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*J. Bacteriol.* 2006, 188(16):5812. DOI: 10.1128/JB.00358-06.

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## Distinguishing Characteristics of Hyperrecombinogenic RecA Protein from *Pseudomonas aeruginosa* Acting in *Escherichia coli*

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Received 13 March 2006/Accepted 9 June 2006

**In *Escherichia coli*, a relatively low frequency of recombination exchanges (FRE) is predetermined by the activity of RecA protein, as modulated by a complex regulatory program involving both autoregulation and other factors. The RecA protein of *Pseudomonas aeruginosa* (RecA<sub>Pa</sub>) exhibits a more robust recombinase activity than its *E. coli* counterpart (RecA<sub>Ec</sub>). Low-level expression of RecA<sub>Pa</sub> in *E. coli* cells results in hyperrecombination (an increase of FRE) even in the presence of RecA<sub>Ec</sub>. This genetic effect is supported by the biochemical finding that the RecA<sub>Pa</sub> protein is more efficient in filament formation than RecA K72R, a mutant protein with RecA<sub>Ec</sub>-like DNA-binding ability. Expression of RecA<sub>Pa</sub> also partially suppresses the effects of *recF*, *recO*, and *recR* mutations. In concordance with the latter, RecA<sub>Pa</sub> filaments initiate recombination equally from both the 5' and 3' ends. Besides, these filaments exhibit more resistance to disassembly from the 5' ends that makes the ends potentially appropriate for initiation of strand exchange. These comparative genetic and biochemical characteristics reveal that multiple levels are used by bacteria for a programmed regulation of their recombination activities.**

RecA protein, a central enzyme of homologous recombination and recombinational DNA repair in bacteria, plays a pivotal role in genome reproduction and the maintenance of genome integrity (13, 20–22, 29, 44). The RecA protein of *Escherichia coli* (RecA<sub>Ec</sub>) was the first member found of a larger family that includes the nearly ubiquitous RecA proteins in bacterial species, the Rad51 and Dmc1 proteins of eukaryotes, and the archaeal RadA proteins (8, 44).

The in vivo activity of bacterial RecA recombinase is most readily monitored during conjugation. When a fragment of donor DNA is injected into a recipient cell during bacterial conjugation, it is integrated either in its entirety (that is without internal recombination exchanges) or in parts (with additional exchanges inside the fragment). The former integration is stimulated by the two outer ends of the donor fragment (“ends-out” events). The integration of donor fragments accompanied by additional exchanges is stimulated by inner single-stranded (ss) or double-stranded (ds) ends (“ends-in” events) and is classically represented as that proceeding through single-strand gap repair (SSGR) and double-strand break repair (DSBR) mechanisms, respectively (for a review, see reference 14). The ends-out events are quantitatively characterized by genetic parameters, such as the yield of recombinants, which is usually normalized per number of donors or transconjugants. The ends-in events can be quantitatively described by the linkage frequency between multiple donor markers. This can be expressed as the frequency of recombi-

nation exchanges (FRE) per DNA unit length (for reviews, see references 24 and 25).

*E. coli* possesses two major recombination systems called the RecBC (later called the RecBCD) and RecF pathways (11), which appear to act on different types of lesions (2, 14). The RecBCD pathway predominates in the DSBR mechanism, while SSG appear to be repaired by the RecF pathway. During conjugation, the RecF pathway is readily observed only in a *recBC sbcB sbcC* background (11, 12). This involves inactivation, respectively, of the RecBCD helicase-ss endonuclease (3), the (3'→5')-directed ss exonuclease I (39, 40), and the ATP-dependent dsDNA exonuclease SbcCD (12).

As previously suggested (2, 14, 24), ends-in exchanges are stimulated by daughter strand gaps. These can occur during replication and range from 240 to 730 bases per replication fork in vivo (17). The gaps are constantly presented during conjugation, both during replication of the donor DNA, when the single-stranded DNA (ssDNA) of the incoming donor fragment is converted to a duplex form via the synthesis of Okazaki fragments (28), and in recipient DNA during normal chromosome duplication. In vivo, the ssDNA is presumably complexed with ssDNA-binding protein (SSB). Consequently, to use ssDNA gaps as substrates for recombination, RecA protein has to displace SSB and polymerize on ssDNA. Under normal conditions, such events are very infrequent and recombination events are initiated mainly by proximal and distal ends of the donor DNA fragment (54). In fact, during conjugational recombination, the FRE value is only one exchange per 20 min of the *E. coli* chromosome (9, 25, 60). However, FRE can be elevated as much as 26-fold under special conditions of hyperrecombination that include SOS derepression, MutS inactivation, and RecA mutational activation (25). Note that the hyperrecombination does not increase, at least noticeably, the

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yield of conjugal recombinants, because all events with low or increased FRE proceed on the same set of chromosomes inside the same number of recipient cells (5, 25).

The RecF, RecO, and RecR proteins assist RecA in competition with SSB for binding to ssDNA and in some specific recombination functions as well. First, RecO physically interacts with both RecR and SSB (49, 56). The RecOR complex binds to SSB-covered ssDNA and facilitates formation of a RecA filament proficient in strand exchange (57). Second, the RecO protein promotes the annealing of complementary ssDNAs complexed with SSB. The RecR protein inactivates the annealing function of RecO but stimulates its mediator function in RecA-promoted strand exchange (18). Third, in the presence of ATP, RecF and RecR proteins physically interact and bind to double-stranded DNA (dsDNA). With enough protein, RecFR can coat duplex DNA uniformly (59). Fourth, RecA filament assembles on linear ssDNA by extension in the 5'-to-3' direction (41, 49). The filament also disassembles with the same polarity (6), but the RecOR proteins stabilize RecA filaments even at the 5' ends of linear ssDNA (49) and thus facilitate RecA-mediated D-loop formation at these ends (7). Fifth, recombination-directed replication (a break copy mechanism) is stimulated *in vitro* by the RecO and RecR proteins and inhibited by the presence of RecF (7, 61). Sixth, the RecF protein physically interacts with the RecX protein to protect RecA from the inhibitory activity of RecX during RecA filament extension (30).

The yield of conjugal recombinants can be greatly affected in cells lacking the *recF*, *recO*, and *recR* genes (16, 19, 32). Interestingly, each of the two parameters characterizing the ends-out and ends-in events, the yield of recombinants and FRE, has been found to be similar for both the RecBCD and RecF pathways (9, 16).

*Pseudomonas aeruginosa* is an aerobic gram-negative bacterium commonly found free living in moist environments. It is also an opportunistic pathogen of plants, animals, and humans. The RecA protein of *P. aeruginosa* (RecA<sub>Pa</sub>) is similar to RecA<sub>Ec</sub>, with amino acid changes at 101 positions (overall 71% identity and 86% similarity). A distinguishing characteristic of the RecA<sub>Pa</sub> protein is its constitutive hyperrecombinogenic (hyper-rec) activity during conjugation when introduced into *E. coli* cells. This enhancement of recombination is SOS independent (5). In *E. coli*, expression of RecA<sub>Pa</sub> increases FRE by six- to eightfold (37). RecA<sub>Pa</sub> displaces both the *E. coli* SSB and *P. aeruginosa* SSB from ssDNA more efficiently than RecA<sub>Ec</sub> does (4). RecA<sub>Pa</sub> forms more salt-stable (4, 37) and temperature-stable (10) presynaptic structures. It possesses a greater affinity for ssDNA than RecA<sub>Ec</sub> does (4). It promotes faster joint molecule formation in DNA strand exchange reactions but is less effective in generating the final products of DNA strand exchange (37). The propensity of RecA<sub>Pa</sub> to initiate but not complete strand exchange is a characteristic of recombinase nucleoprotein filaments with enhanced DNA pairing and/or duplex DNA binding activities (21, 27). Once a strand exchange reaction is initiated, additional interactions between the recombinase filament and the same or different duplex DNAs can impede the DNA rotation needed to extend the heteroduplex DNA in the nascent joint molecule (43, 52, 53).

Here, we continue the analysis of RecA<sub>Pa</sub> function by focus-

ing on two additional characteristics of RecA<sub>Pa</sub> that are thought to be important for the molecular mechanism of recombination in general and hyperrecombination in particular. These include the RecA<sub>Pa</sub> ability (i) to form hyperreactive presynaptic filament in *E. coli* in a high background of native RecA<sub>Ec</sub> protein and (ii) to suppress, at least partially, defects of RecF, RecO, and RecR proteins during presynaptic complex formation.

## MATERIALS AND METHODS

***E. coli* strains and plasmids.** Donor strain KL227 (HfrP4x *metB*) and recipient strains AB1157 (*thr-1 leuB6 proA2 hisG4 argE3 thi-1 supE44 rpsL31*) of the RecBCD recombination pathway, JC7623 (as AB1157 but *recBC sbcCD*) of the RecF pathway, and recombination-deficient JC10289 (as AB1157 but  $\Delta[\textit{recA-srlR306}::\textit{Tn10} = \Delta\textit{recA306}]$ ) were from A. J. Clark's collection. Recombination-deficient strains JC7623-F, JC7623-O, and JC7623-R were constructed by P1 transduction to transfer *recO1504::Tn5* (36), *recR252::Tn10-9* (33), and *recF349(del)* (47) mutant genes, respectively, into strain JC7623. Plasmids pRecA<sub>Ec</sub> (pUC19-*recA1.1*) and pRecA<sub>Pa</sub> (pUC19-PA189.1) contain *recA<sub>Ec</sub>* and *recA<sub>Pa</sub>* genes, respectively, together with their native operator-promoter regions as described previously (37). Plasmid P200 F'-*lac* was used to standardize conjugation abilities of recipient strains.

**DNA.** Circular M13mp8 ssDNA was from New England Biolabs. Its nucleotide molar concentration was determined by absorbance at 260 nm using an extinction coefficient of  $6.5 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ . The etheno-modified calf thymus ssDNA was as described previously (1). Its nucleotide molar concentration was determined by absorbance at 260 nm using an extinction coefficient of  $8.325 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ . M13mp8.1037(+) circular ssDNA and supercoiled M13mp8 dsDNA were from M. Cox's lab.

**Other reagents.** ATP was from Sigma. All other reagents used in the study were research grade and were commercially available.

**Proteins.** The RecA<sub>Ec</sub> and RecA<sub>Pa</sub> proteins were purified as described previously (37). RecA K72R protein was from M. Cox's lab. Concentrations of RecA<sub>Ec</sub> and RecA K72R were determined by absorbance at 280 nm, using an extinction coefficient of  $2.23 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$ . The RecA<sub>Pa</sub> protein concentration was determined with the help of a protein Coomassie Plus protein assay reagent kit (Pierce) using the RecA<sub>Ec</sub> protein as a standard. *E. coli* SSB was from Sigma, USB. The concentration was determined by absorbance at 280 nm, using an extinction coefficient of  $2.83 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$ . Lactate dehydrogenase, phosphoenolpyruvate, and pyruvate kinase were from Sigma.

**Conjugation.** Conjugation was carried out essentially as described previously (25). Both Hfr and F<sup>-</sup> strains were grown, crossed, and selected for recombinants at 37°C in mineral salts 56/2 medium supplied with all necessary growth factors at pH 7.5. The ratio between donors and recipients in mating mixtures was 1:10,  $2 \times 10^7$  to  $4 \times 10^7$  donors and  $2 \times 10^8$  to  $4 \times 10^8$  recipients per 1 ml. The yield of Thr<sup>+</sup> Str<sup>r</sup> recombinants in all independent crosses (5 to 7% relative to donors) was normalized by the mating ability of each recipient used. The latter was determined by the yield of transconjugants F'-*lac*<sup>+</sup> in crosses between the recipients and donor P200 F'-*lac*.

**FRE value calculations.** FRE value calculations were carried out as described previously (5, 25). Quantitative estimations of FRE alterations ( $\Delta\text{FRE}$ ) promoted by the *P. aeruginosa recA* (*recA<sub>Pa</sub>*) gene relative to the FRE value promoted by the *E. coli recA* (*recA<sub>Ec</sub>*) gene were done by use of the following formula:  $\Delta\text{FRE} = \ln(2\mu_1 - 1)/(2\mu_2 - 1)$ , where  $\mu_2$  is the linkage of selected *thr*<sup>+</sup> and unselected *leu*<sup>+</sup> markers in a cross with strain AB1157 and  $\mu_1$  is similar linkage in the cross analyzed. Donor KL227 transfers *leu*<sup>+</sup> and *thr*<sup>+</sup> as a proximal and distal marker, respectively. Calculations of uncertainty in determinations of relative values of FRE were done as deviations from the average values by making use of the Excel 97 program with formula ( $= 2 \times \text{standard deviation}$ ) and by inputting the values from independent repeats of three experiments.

**Determination of intracellular RecA<sub>Ec</sub> and RecA<sub>Pa</sub> amounts.** The intracellular amounts of RecA<sub>Ec</sub> and RecA<sub>Pa</sub> were determined in all recipients used for FRE analyses and presented relative to the amount of RecA<sub>Ec</sub> in strain AB1157. *E. coli* cells were grown up to mid-log phase in LB medium at 37°C.

A cell pellet containing  $5 \times 10^7$  cells was lysed by boiling with sodium dodecyl sulfate, electrophoresed through sodium dodecyl sulfate-10% polyacrylamide gels. The RecA<sub>Ec</sub> and RecA<sub>Pa</sub> amounts were detected by immunoblotting using polyclonal chicken antibodies to these proteins (Genetel Lab) in a standard procedure (46). Primary antibody binding was visualized with secondary antibodies coupled to horseradish peroxidase (Genetel Lab). The blots were then

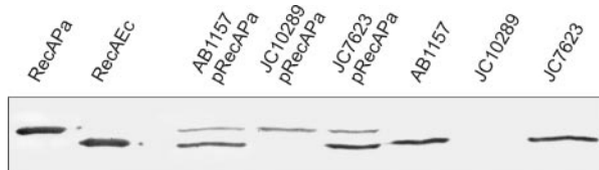


FIG. 1. Visualization and quantitation of the RecA<sub>Ec</sub> and RecA<sub>Pa</sub> proteins by immunoblotting with a mixture of polyclonal antibodies raised against RecA<sub>Ec</sub> and RecA<sub>Pa</sub>. For controls, lanes RecAPa and RecAEc contain equal amounts of pure RecA proteins. Other lanes show the data from a typical experiment comparing the RecA<sub>Pa</sub> and RecA<sub>Ec</sub> amounts in different strains when protein was extracted from the same number of cells (for details, see Materials and Methods).

stained with diaminobenzidine (Sigma) and scanned, and the amounts of proteins were documented by the use of the TotalLab program. The data of two independent experiments were averaged. Polyclonal antibodies raised against RecA<sub>Ec</sub> and RecA<sub>Pa</sub> showed a strong cross-reactivity though, in principle, their specificities were observed with the amount of RecA proteins lower than 0.03  $\mu\text{g}$  (data not shown). Although the molecular weights of RecA<sub>Ec</sub> (37,842) and RecA<sub>Pa</sub> (36,877) are close enough, they form well-separated bands in the 10% polyacrylamide gels used in the analysis. This fact allowed us to use an equimolar mixture of antibodies against RecA<sub>Ec</sub> and RecA<sub>Pa</sub> to reveal the intracellular amounts of RecA<sub>Ec</sub> and RecA<sub>Pa</sub> in recipients analyzed relative to the amount of RecA<sub>Ec</sub> in strain AB1157. Figure 1 shows that this equimolar mixture of antibodies visualizes in equal manner both RecA<sub>Ec</sub> and RecA<sub>Pa</sub> proteins presented in equal amount (0.0068  $\mu\text{g}$ ). This control means that this mixture can be successfully used for quantitative determination of the intracellular amounts of RecA<sub>Ec</sub> and RecA<sub>Pa</sub> in different strains.

**ssDNA-dependent ATP hydrolysis.** ssDNA-dependent ATP hydrolysis reactions were carried out at 37°C in TMD buffer (25 mM Tris-HCl [pH 7.5], 10 mM MgCl<sub>2</sub> and 1 mM dithiothreitol) containing 2  $\mu\text{M}$  RecA, 2 mM ATP with its regenerating system (5 mM phosphoenolpyruvate and 30 units ml<sup>-1</sup> pyruvate kinase), coupling system (3 mM NADH and 30 units ml<sup>-1</sup> lactate dehydrogenase), and circular M13mp8 ssDNA and SSB as indicated.

**RecA-mediated D-loop formation.** RecA-mediated D-loop formation was carried out between circular supercoiled dsDNA and linear ssDNA with homologous regions at the 3' or 5' end. The agarose gel assay was used to visualize joint molecule formation. The reactions were carried out in TMD buffer containing 3  $\mu\text{M}$  RecA, 6  $\mu\text{M}$  linear ssDNA [either M13mp8.1037(+)*Pst*I or M13mp8.1037(+)*Eco*RI], 6  $\mu\text{M}$  supercoiled M13mp8 dsDNA, 2 mM ATP with its regenerating system, and 1.4  $\mu\text{M}$  SSB. The reaction mixture was prepared at 0°C, and the reaction was initiated by the temperature shift to 37°C for the time indicated.

**Fluorescence assay of RecA/ATP/ $\epsilon$ DNA complex formation under conditions of poly(dT) challenge.** Experiments were performed at 37°C in TMD buffer containing 2.5  $\mu\text{M}$  RecA, 2 mM ATP with its regenerating system, and 3  $\mu\text{M}$   $\epsilon$ DNA. After RecA<sub>Pa</sub> and RecA<sub>Ec</sub> complexes were preformed and the increase

of fluorescence was established, 50  $\mu\text{M}$  poly(dT) was added to initiate the quenching of fluorescent signal (time zero) as a result of poly(dT) challenge.

## RESULTS

**Low levels of RecA<sub>Pa</sub> significantly increase the frequency of recombinational exchanges even in the presence of wild-type RecA<sub>Ec</sub>.** As mentioned above, FRE is a genetic parameter that calculates ends-in events and thus characterizes the aggressiveness of RecA protein in the initiation of recombination. As expected (25), in the conjugational cross KL227  $\times$  AB1157, the absolute value of FRE was about one exchange per each 20-min region of the *E. coli* chromosome, as measured by *thr-leu* linkage (Table 1).

Plasmid pRecAPa confers ampicillin resistance on its host and contains the *recA<sub>Pa</sub>* gene under the control of its own promoter. This gene is weakly expressed in *E. coli*. In fact, immunoblotting using an equimolar mixture of polyclonal antibodies to the RecA<sub>Ec</sub> and RecA<sub>Pa</sub> proteins (see Materials and Methods) showed that the level of RecA<sub>Pa</sub> in JC10289/pRecAPa cells is only 30% of the RecA<sub>Ec</sub> concentration found in strain AB1157 (compare lanes AB1157, JC10289, and JC10289/pRecAPa in Fig. 1 and their quantitation presented in Table 1). However, this relatively small amount of RecA<sub>Pa</sub> provided a 7.9-fold increase in recombination frequency as measured by  $\Delta$ FRE (Table 1). Moreover, the same low-level expression of RecA<sub>Pa</sub> in a background including wild-type RecA<sub>Ec</sub> resulted in a fivefold enhancement of recombination when measured separately for both the RecBCD and RecF pathways (see data for AB1157/pRecAPa and JC7623/pRecAPa, respectively, in Fig. 1 and Table 1).

In principle, recombination may have occurred between the *recA<sub>Ec</sub>* and *recA<sub>Pa</sub>* genes (located in the chromosome and plasmid, respectively) in these strains, resulting in exchange of location of the genes or production of chimeric *recA<sub>Ec</sub>/recA<sub>Pa</sub>* genes with novel properties. In order to be sure that we used conjugation recipients with the reported configuration of the *recA<sub>Ec</sub>* and *recA<sub>Pa</sub>* genes, the following control experiments were carried out. Two recipient strains used in mating, AB1157/pRecAPa and JC7623/pRecAPa, were diluted 1 to 10 in 56/2 growth medium without ampicillin (which served as a plasmid selection factor) in order to allow multiplication of

TABLE 1. Intracellular amounts of RecA<sub>Ec</sub> or RecA<sub>Pa</sub> protein in different recipients and relative change in FRE values measured in crosses between donor KL227 (HfrR4) and recipients

Recipient	Relevant genotype		Relative intracellular amt (mean $\pm$ SD) of <sup>b</sup> :		Recombination parameter	
	<i>recA</i>	Other <i>rec</i> gene <sup>a</sup>	RecA <sub>Ec</sub>	RecA <sub>Pa</sub>	Linkage ( <i>thr<sup>+</sup>-leu<sup>+</sup></i> ) (mean $\pm$ SD)	$\Delta$ FRE
AB1157	<i>recA<sub>Ec</sub><sup>+</sup></i>	wt	1.0	0.0	0.924 $\pm$ 0.018 (800) <sup>c</sup>	1
JC10289	$\Delta$ <i>recA<sub>Ec</sub></i>	wt	0.0	0.0	ND <sup>d</sup>	ND
JC10289/pRecAPa	$\Delta$ <i>recA<sub>Ec</sub></i> + <i>recA<sub>Pa</sub><sup>+</sup></i>	wt	0.0	0.27 $\pm$ 0.05	0.636 $\pm$ 0.028 (1,000)	7.9
AB1157/pRecAPa	<i>recA<sub>Ec</sub><sup>+</sup></i> + <i>recA<sub>Pa</sub><sup>+</sup></i>	wt	0.67 $\pm$ 0.09	0.20 $\pm$ 0.02	0.718 $\pm$ 0.015 (1,200)	5.0
JC7623	<i>recA<sub>Ec</sub><sup>+</sup></i>	<i>recBC sbcBC</i>	0.90 $\pm$ 0.20	0.0	0.919 $\pm$ 0.014 (1,200)	1.1
JC7623/pRecAPa	<i>recA<sub>Ec</sub><sup>+</sup></i> + <i>recA<sub>Pa</sub><sup>+</sup></i>	<i>recBC sbcBC</i>	0.84 $\pm$ 0.05	0.32 $\pm$ 0.05	0.715 $\pm$ 0.018 (600)	5.1

<sup>a</sup> wt means that other *rec* genes are of wild type or that the recipient has the RecBCD pathway of recombination; *recBC sbcBC* recipients have the RecF pathway.

<sup>b</sup> Intracellular RecA<sub>Ec</sub> and RecA<sub>Pa</sub> amounts were determined relative to the amount of RecA<sub>Ec</sub> in strain AB1157. The data of two independent experiments were averaged. The data are given after subtraction of the background (0.003  $\pm$  0.002).

<sup>c</sup> Numbers in parentheses are the numbers of clones analyzed (for details, see Materials and Methods).

<sup>d</sup> ND, not determined.

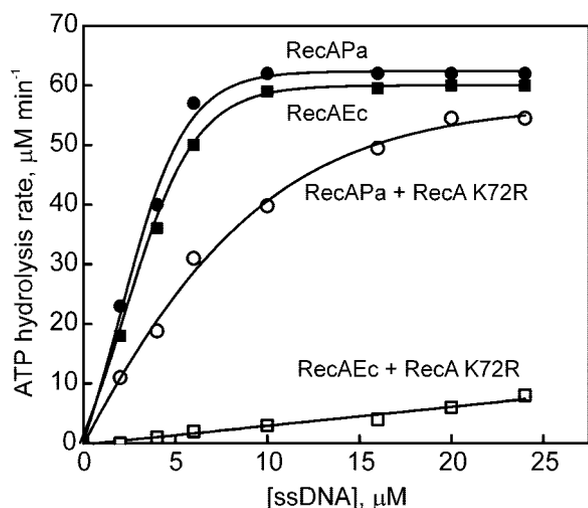


FIG. 2. ssDNA-dependent ATP hydrolysis of four presynaptic complexes formed by RecA<sub>Pa</sub>, RecA<sub>Ec</sub>, RecA<sub>Pa</sub> plus RecA K72R, and RecA<sub>Ec</sub> plus RecA K72R at increasing concentrations of circular ssDNA. SSB was used at a concentration providing one SSB monomer per 10 nucleotides of ssDNA. Each point on the curves represents an individual sample and shows a steady-state rate measured over a period of about 15 min.

cells lacking the pRecAPa plasmid. The recombination parameters of these novel populations of AB1157 and JC7623 recipients were determined in crosses with donor KL227 and compared with those presented in Table 1. After 6 to 8 generations in the medium without ampicillin, the recombination frequency (FRE) was restored to near wild-type levels in both AB1157 and JC7623. The ΔFRE declined from 5.0 to 3.4 after 4 generations for strain AB1157 and from 5.1 to 2.2 after 3 generations for strain JC7623. The ΔFRE changed to 1.1 after an additional 3 or 4 generations for both AB1157 and JC7623 when 45 to 47 clones from 50 analyzed for each culture had lost the Amp<sup>r</sup> phenotype because of the loss of pRecAPa plasmids. These nearly complete restorations of FRE values after the loss of pRecAPa plasmids by recipients show that both AB1157/pRecAPa and JC7623/pRecAPa cultures used in the experiments were not contaminated to any significant degree by recombinant variants present in the *E. coli* chromosome.

The data presented in this section indicate the following. (i) The RecA<sub>Pa</sub> protein exhibits a more robust recombinase activity, as measured by decreased genetic linkage *in vivo*, than the RecA<sub>Ec</sub> protein does. (ii) This property does not depend on the recombination pathway used. (iii) The RecA<sub>Pa</sub> protein is much more efficient in initiation of additional recombination exchanges even in the presence of RecA<sub>Ec</sub>. The latter statement is supported by the following biochemical observation.

**Unlike RecA<sub>Ec</sub> protein, RecA<sub>Pa</sub> effectively competes with RecA K72R in presynaptic complex formation.** Figure 2 shows the ssDNA concentration dependence of the ATPase activity produced by different presynaptic complexes of the type RecA/ATP/circular M13 ssDNA, all formed in the presence of sufficient SSB to remove the ssDNA secondary structures. Two control and two test complexes were compared. The controls included presynaptic complexes with pure RecA<sub>Pa</sub> and RecA<sub>Ec</sub>; the test complexes contained protein mixtures of either RecA<sub>Ec</sub>

and RecA K72R or RecA<sub>Pa</sub> and RecA K72R taken in equal proportion. The Lys-to-Arg substitution in the RecA K72R protein occurs in a well-conserved nucleotide binding region. The protein binds but does not hydrolyze ATP, though it is still able to promote the fundamental DNA pairing reaction (42). Moreover, this mutant protein forms mixed presynaptic filaments with RecA<sub>Ec</sub>, competing on an equal basis with wild-type RecA<sub>Ec</sub> for binding sites in ssDNA (50, 51).

Each point of the curves presented in Fig. 2 is the rate of ATP hydrolysis obtained 15 min after initiation of the reaction by mixing RecA(s), ATP with its regenerating system, and closed circular ssDNA of phage M13mp8 in buffer (pH 7.5) at 37°C. For pure RecA<sub>Ec</sub> and RecA<sub>Pa</sub>, the ATP hydrolysis rate increased with ssDNA concentration to a maximum of 60 to 63 μM min<sup>-1</sup> with an apparent stoichiometry of about three bases per RecA subunit. As expected (50), RecA K72R did not hydrolyze ATP, and the hydrolysis rate observed for an equal mixture of RecA<sub>Ec</sub> and RecA K72R proteins increased only slightly with increasing ssDNA concentration and had an approximately 10-fold-lower acceleration. This resulted from formation of mixed RecA filaments and inhibition of ATP hydrolysis because of the dimension of cooperative interactions between RecA<sub>Ec</sub> protomers in filaments when RecA K72R monomers are included (50). Quite another situation was found for the mixture of RecA<sub>Pa</sub> plus RecA K72R (equal amounts of proteins). The ATP hydrolysis rate increased with ssDNA concentration, reaching 87% of the maximal value described for pure RecA<sub>Pa</sub>. Thus, the excess ssDNA in the reaction mixture resulted in formation of presynaptic complexes with ATPase activity that was closer to that of pure RecA<sub>Pa</sub> filaments.

The data indicate that RecA<sub>Pa</sub> protein is much more active in competition with RecA K72R for presynaptic filament formation. Because the capacity of the mutant RecA K72R protein to bind ssDNA is not changed significantly relative to that of RecA<sub>Ec</sub> (42), the data reveal the RecA<sub>Pa</sub> advantages in recombination initiation in comparison with RecA<sub>Ec</sub>.

**RecA<sub>Pa</sub> protein partially suppresses the genetic consequences of *recF*, *recO*, and *recR* mutations.** Inactivation of the *recF*, *recO*, or *recR* gene results in a substantial decrease in the yield of recombinants during conjugation in *E. coli* cells relying on the RecF pathway. On the other hand, as shown in Table 1, relatively low levels of RecA<sub>Pa</sub> protein promote hyperrecombination in *E. coli* cells utilizing the RecF pathway. We decided to determine whether the presence of RecA<sub>Pa</sub> could compensate for the loss of *recF*, *recO*, or *recR* function.

Table 2 presents the relative yield of Thr<sup>+</sup> Str<sup>r</sup> recombinants (RYR) obtained in three sets of crosses (each repeated twice) between donor strain KL227 or KL227 *recF349*, as indicated, and four different recipient strains with different RecF pathway genotypes including (i) classical strain JC7623 *recBC sbcCD* (a basal cross); (ii) JC7623/pRecAPa, in which hyperrecombination by RecA<sub>Pa</sub> protein is promoted in the RecA<sub>Ec</sub> background (the first testing cross); (iii) JC7623-0, JC7623-R, or JC7623-F recipients deficient in general recombination because of defects in the *recO*, *recR*, or *recF* gene, respectively (the second test); and (iv) JC7623-0/pRecAPa, JC7623-R/pRecAPa, or JC7623-F/pRecAPa recipients deficient in RecA<sub>Ec</sub>-dependent recombination while potentially proficient in recombination promoted by RecA<sub>Pa</sub> (the third test). The yield of recombi-

TABLE 2. Partial suppression of recombination deficiency that resulted from *recO*, *recR*, or *recF* mutations in the RecF pathway of recombination by RecA<sub>Pa</sub> protein as observed via the relative yield of Thr<sup>+</sup> Str<sup>r</sup> recombinants in crosses between donor KL227 (HfrR4-type) and recipients originating from JC7623 *recBC sbcCD*

Recipient or recombination	Relevant genotype		R <sub>YR</sub> <sup>a</sup>	
	<i>recA</i> <sup>+</sup>	Other <i>rec</i> gene <sup>b</sup>	Expt 1	Expt 2
JC7623	<i>recA</i> <sub>Ec</sub> <sup>+</sup>		1.0 (7.1) <sup>c</sup>	1.0 (3.4)
JC7623/pRecAPa	<i>recA</i> <sub>Ec</sub> <sup>+</sup> + <i>recA</i> <sub>Pa</sub> <sup>+</sup>		0.96	1.0
JC7623-O	<i>recA</i> <sub>Ec</sub> <sup>+</sup>	<i>recO1504</i>	0.024	0.020
JC7623-O/pRecAPa	<i>recA</i> <sub>Ec</sub> <sup>+</sup> + <i>recA</i> <sub>Pa</sub> <sup>+</sup>	<i>recO1504</i>	0.30	0.35
Deficiency <sup>d</sup> ( <i>recO</i> <sup>+</sup> / <i>recO1504</i> )			41.6	50.0
Suppression <sup>e</sup> ( <i>recA</i> <sub>Ec</sub> <sup>+</sup> <i>recA</i> <sub>Pa</sub> <sup>+</sup> <i>recO1504</i> / <i>recA</i> <sub>Ec</sub> <sup>+</sup> <i>recO1504</i> )			12.5	17.5
JC7623	<i>recA</i> <sub>Ec</sub> <sup>+</sup>		1.0 (5.2)	1.0 (5.0)
JC7623/pRecAPa	<i>recA</i> <sub>Ec</sub> <sup>+</sup> + <i>recA</i> <sub>Pa</sub> <sup>+</sup>		0.85	1.0
JC7623-R	<i>recA</i> <sub>Ec</sub> <sup>+</sup>	<i>recR252</i>	0.0067	0.0056
JC7623-R/pRecAPa	<i>recA</i> <sub>Ec</sub> <sup>+</sup> + <i>recA</i> <sub>Pa</sub> <sup>+</sup>	<i>recR252</i>	0.108	0.05
Deficiency ( <i>recR</i> <sup>+</sup> / <i>recR252</i> )			145	178
Suppression ( <i>recA</i> <sub>Ec</sub> <sup>+</sup> <i>recA</i> <sub>Pa</sub> <sup>+</sup> <i>recR252</i> / <i>recA</i> <sub>Ec</sub> <sup>+</sup> <i>recR252</i> )			16.1	8.9
JC7623	<i>recA</i> <sub>Ec</sub> <sup>+</sup>		1.0 (4.3)	1.0 (4.4)
JC7623/pRecAPa	<i>recA</i> <sub>Ec</sub> <sup>+</sup> + <i>recA</i> <sub>Pa</sub> <sup>+</sup>		0.95	1.05
JC7623-F <sup>f</sup>	<i>recA</i> <sub>Ec</sub> <sup>+</sup>	<i>recF349</i>	0.0033	0.0049
JC7623-F/pRecAPa <sup>f</sup>	<i>recA</i> <sub>Ec</sub> <sup>+</sup> + <i>recA</i> <sub>Pa</sub> <sup>+</sup>	<i>recF349</i>	0.029	0.03
Deficiency ( <i>recF</i> <sup>+</sup> / <i>recF349</i> )			300	204
Suppression ( <i>recA</i> <sub>Ec</sub> <sup>+</sup> <i>recA</i> <sub>Pa</sub> <sup>+</sup> <i>recF349</i> / <i>recA</i> <sub>Ec</sub> <sup>+</sup> <i>recF349</i> )			8.8	6.1

<sup>a</sup> R<sub>YR</sub> means the yield of Thr<sup>+</sup> Str<sup>r</sup> recombinants relative to that found in crosses with JC7623, which is identical to the yield of Thr<sup>+</sup> Str<sup>r</sup> recombinants for strain AB1157 (cells with the RecBCD pathway). The yield of recombinants in all independent crosses was normalized by the mating ability of each recipient used.

<sup>b</sup> The full names of mutations used were as follows: *recO1504*::Tn5, *recR252*::Tn10-9, and *recF349*(del).

<sup>c</sup> Numbers in parentheses show the yield of Thr<sup>+</sup> Str<sup>r</sup> recombinants as a percentage of the donors in the conjugation mixture.

<sup>d</sup> Deficiency of recombination is a decrease in the R<sub>YR</sub> that resulted from the *recO*, *recR*, or *recF* insertion or deletion mutations used in the study.

<sup>e</sup> Suppression of the deficiency mentioned above resulted from introduction into recipients the pRecAPa plasmid, which expressed the RecA<sub>Pa</sub> protein.

<sup>f</sup> In crosses with JC7623 *recF349*, Hfr KL227 *recF349* was used. It was necessary to exclude the possibility of transferring the *recF*<sup>+</sup> allele by the donor during a 60-min mating used in the study.

nants in all three testing crosses was normalized to that in the basal cross. The result of the first test showed that addition of a small portion of RecA<sub>Pa</sub> protein (30%) to the RecA<sub>Ec</sub> protein in the cells, which rendered them hyper-rec as measured by FRE (Table 1), did not change R<sub>YR</sub> significantly, if any (Table 2). The second test showed that both insertion mutations, *recO1504* and *recR252*, and the deletion mutation *recF349* lowered the yield of recombinants by 40 to 50, 140 to 180, and 200 to 300 times, respectively, a result that is in satisfactory agreement with the data in the literature (33, 36, 47). The third test cross revealed average increases of 15-, 12.5-, and 7.5-fold in recombination frequency when RecA<sub>Pa</sub> was provided in backgrounds deficient in the *recO*, *recR*, and *recF* genes, respectively.

These in vivo data indicate that relative to RecA<sub>Ec</sub> protein, the hyper-rec RecA<sub>Pa</sub> protein is less dependent on the *E. coli* RecO, RecR, and RecF proteins in initiation of recombination events in the RecF pathway.

As known (7), the RecOR protein complex facilitates RecA-mediated D-loop formation at the 5' ends of linear ssDNA. If RecA<sub>Pa</sub> were able to compensate at least partially for the inactivity of the RecOR complex, one could expect the RecA<sub>Pa</sub> function to be more active at 5' ends of ssDNA. This was the case.

**RecA<sub>Pa</sub> protein promotes D-loop formation with equal efficiency at both the 3' and 5' ends of linear ssDNA, even in the presence of SSB.** To determine the extent of any end bias in the D-loop formation promoted by the RecA<sub>Pa</sub> and RecA<sub>Ec</sub> presynaptic complexes, an in vitro joint molecule assay was used. The assay monitors homologous pairing between linear M13mp8 ssDNA and supercoiled dsDNA of the same phage (7). A 7-kb region of homology between these substrates was restricted to either the 3' or 5' end of the linear ssDNA by insertion at its 5' or 3' end, respectively, of a 1,037-nucleotide segment heterologous to the M13mp8 DNA (see the scheme at the top of Fig. 3).

D-loop formation between the linear ssDNA and supercoiled dsDNA was carried out in the presence of SSB at 37°C. The time course of appearance of the 3'- or 5'-end-directed joint molecules promoted by RecA<sub>Ec</sub> or RecA<sub>Pa</sub> proteins is presented in Fig. 3A. Quantitation of these events is presented in Fig. 3B. Consistent with previous observations (37), joint molecule formation proceeded faster for RecA<sub>Pa</sub> than for RecA<sub>Ec</sub>, though both reached a maximum within 4 min and sustained that level for some time thereafter.

The values presented in Fig. 3C are the quantitative estimations of maximum joint molecule formation observed in Fig. 3A and analyzed in Fig. 3B (reactions after 4, 6, and 8 min),

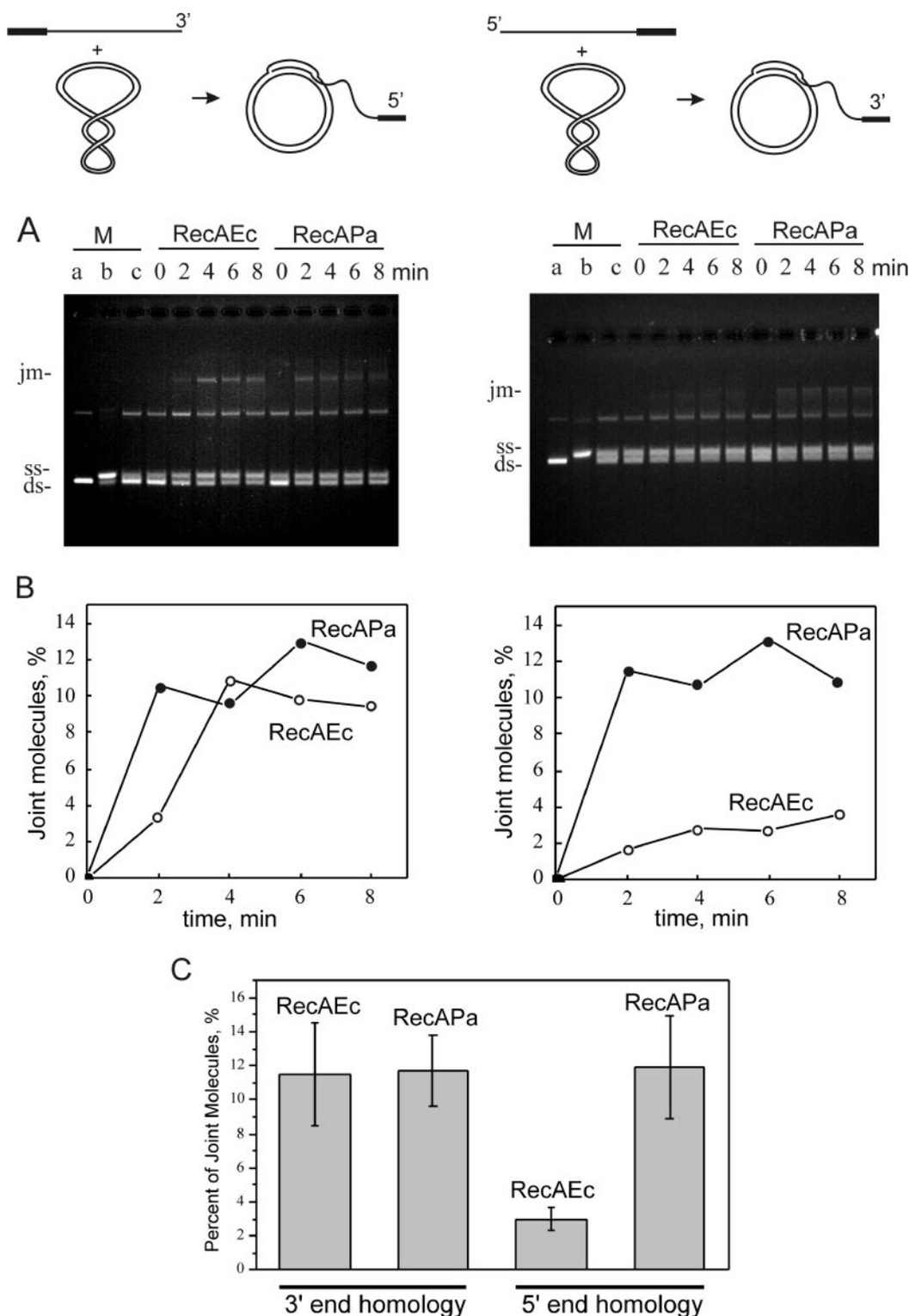


FIG. 3. Time course of initiation of D-loop formation between circular supercoiled dsDNA and linear ssDNA with a region of homology at either the 3' or 5' end in a comparison of the characteristics of RecA<sub>Pa</sub> and RecA<sub>Ec</sub> in promotion of the reaction. The DNA substrates and the reactions are illustrated by schemes presented at the top of the figure. M13mp8.1037(+) circular ssDNA was linearized with either EcoRI or PstI to place the 1,037-nucleotide insert (filled rectangle) at the 5' or 3' end, respectively. The supercoiled M13mp8 dsDNA has 7,229 bp of homology to the 3' end of EcoRI-digested M13mp8.1037(+) linear ssDNA or to the 5' end of PstI-digested M13mp8.1037(+) linear ssDNA. The agarose gel assay was used to visualize joint molecule formation as described in Materials and Methods. (A) Agarose gel presentation of joint molecule formation. Lanes a, b, and c contain markers (M) of dsDNA, ssDNA, and dsDNA plus ssDNA, respectively. The positions of joint molecules (jm), linear ssDNA (ss), and closed circular dsDNA (ds) are shown to the left of the gels. (B) Quantitation of the data presented in panel A by use of the Kodak DS1D image analysis software. (C) Values calculated from the data presented in panel B and two other independent experiments. The bands obtained in the D-loop reactions proceeding 4, 6, and 8 min after initiation were summarized because they looked similar.

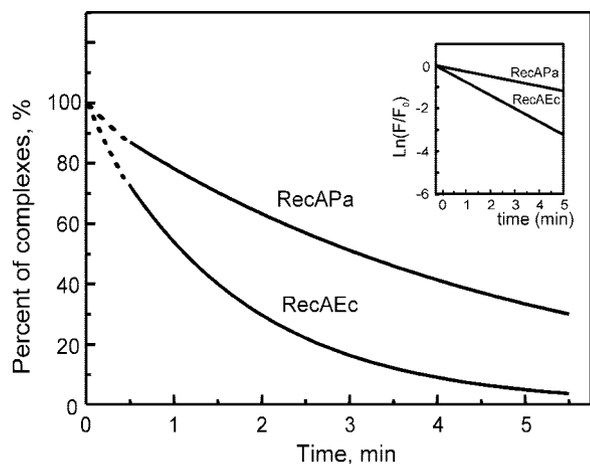


FIG. 4. Time-course decrease in fluorescence after the addition of excess poly(dT) to the RecA/ATP/ $\epsilon$ DNA complex in a comparison of disassembly of RecA<sub>Pa</sub> and RecA<sub>Ec</sub> complexes. The presynaptic complex RecA/ATP/ $\epsilon$ DNA was formed through the primary DNA binding site of RecA (excess RecA compared to  $\epsilon$ DNA). Experiments were performed as described in Materials and Methods. After RecA<sub>Pa</sub> and RecA<sub>Ec</sub> complexes were preformed, the increase of fluorescence was established. Excess poly(dT) was then added to initiate the quenching of fluorescent signal (time zero) as result of the poly(dT) challenge. The measurements were started 0.5 min later. The broken lines are the extrapolation of data to the moment of poly(dT) addition to the reaction mixture. (Inset) The same data but presented in semilogarithmic scale.  $F/F_0$  is a portion of fluorescence  $F$  of the RecA<sub>Ec</sub> or RecA<sub>Pa</sub> presynaptic complexes still existing to the time indicated;  $F_0$  is the fluorescence at time zero.

averaged from three independent experiments as the percentage of the total DNA (ssDNA plus dsDNA plus joint molecules) participating in the reaction. When the linear ssDNA was homologous at its 3' end with its supercoiled partner, joint molecules were formed with equal efficiency in the reactions promoted by the RecA<sub>Ec</sub> and RecA<sub>Pa</sub> proteins. However, when the homology was limited to the 5' ends, the efficiency of reactions promoted by RecA<sub>Ec</sub> was reduced about fourfold, in good agreement with observations described earlier (7, 15). In contrast, the RecA<sub>Pa</sub> protein promoted the formation of joint molecules with equal proficiency at both the 5' and 3' ends of the linear ssDNA.

This finding is in good accordance with the capacity of RecA<sub>Pa</sub> protein to form relatively stable presynaptic filaments on linear ssDNA which might persist at the 5' ends of the ssDNA (6, 7, 40).

**RecA<sub>Pa</sub> protein filaments disassemble at only half the rate of RecA<sub>Ec</sub> protein filaments.** As documented previously (6), the RecA<sub>Ec</sub> presynaptic complex disassembles on linear ssDNA in the 5'-to-3' direction. To compare the rates of disassembly of the RecA<sub>Pa</sub> and RecA<sub>Ec</sub> filaments physically and measure the rate quantitatively, the change of fluorescence of RecA protein complexed with etheno-DNA ( $\epsilon$ DNA) was used. RecA binding to  $\epsilon$ DNA results in a proportional increase of fluorescence upon complex formation (34) and vice versa in a fluorescence decrease when the complex disassembles. To form the presynaptic RecA/ATP/ $\epsilon$ DNA complex so that no more than one ssDNA was bound per RecA filament, we limited the amount of  $\epsilon$ DNA in the reaction mixture. The

binding of calf thymus  $\epsilon$ DNA used in the study to RecA<sub>Ec</sub> or RecA<sub>Pa</sub> proteins resulted in an increase of the specific fluorescence by a factor of 3.25 (data not shown). These complexes were formed and then challenged with excess poly(dT) to bind free RecA and any dissociated RecA molecules, and the resulting decline in fluorescence was monitored.

As shown in Fig. 4, the time-dependent drop in fluorescence proceeds exponentially for both the RecA<sub>Ec</sub> and RecA<sub>Pa</sub> presynaptic complexes, as expected for a first-order disassembly process. When plotted as  $\ln(F/F_0)$  versus time (see inset to Fig. 4), where  $F$  is the fluorescence at a given time and  $F_0$  is the original fluorescence (at time zero), both the RecA<sub>Pa</sub> and RecA<sub>Ec</sub> are converted into linear functions. The ratio of the half-time period of the drop in fluorescence, RecA<sub>Ec</sub>/RecA<sub>Pa</sub>, is 1:2.7. This means that the RecA<sub>Pa</sub> filaments are more than twice as resistant to poly(dT) challenge as the RecA<sub>Ec</sub> filaments are.

The data indicate that the rate of RecA<sub>Pa</sub> disassembly from the 5' end of ssDNA proceeds much slower than that of RecA<sub>Ec</sub>.

## DISCUSSION

Genetic analysis of the dependence of FRE values on RecA protein (either RecA<sub>Ec</sub> or RecA<sub>Pa</sub> or both proteins) mediating homologous recombination in *E. coli* supported a previous observation (37) that the RecA<sub>Pa</sub> protein promotes the ends-in type of recombination events more frequently (eight times) during conjugation. Note that ends-in but not ends-out events appear to be involved in the FRE increase because otherwise a significant increase of recombinant yield could also be observed, but this was not the case. Also, the analysis showed that even a small portion of intracellular RecA presented in *E. coli* by RecA<sub>Pa</sub>, which forms only one third of normal RecA<sub>Ec</sub> concentration (Fig. 1) (that is usually about 10,000 molecules per cell [35, 48]) gave a fivefold increase of FRE, which was reproduced in both RecBCD and RecF genetic pathways of recombination (Table 1). It means that RecA<sub>Pa</sub>, being in *E. coli*, successfully predominates over RecA<sub>Ec</sub> in formation of such presynaptic complexes which are able to initiate additional ends-in recombination events that result in hyperrecombination. Two questions arise. (i) What is the RecA composition of such a presynaptic complex? (ii) What events of DNA metabolism stimulate the ends-in events?

In an attempt to answer the first question, we compared the ATPase activities of presynaptic complexes formed by two protein mixtures: RecA<sub>Pa</sub> plus RecA K72R and RecA<sub>Ec</sub> plus RecA K72R, where all three constituents were presented in equimolar concentrations. Because the RecA K72R protein can only bind but not hydrolyze ATP (42, 52), its participation in presynaptic complex formation might result in a decrease of the ATPase activity of the latter. This expectation was found for RecA<sub>Ec</sub>/RecA K72R complexes, while the decrease was much lower for RecA<sub>Pa</sub>/RecA K72R complexes (Fig. 2). In these experiments, RecA K72R mimics the behavior of RecA<sub>Ec</sub>; thus, this in vitro test allows the conclusion that in full concordance with in vivo analysis, RecA<sub>Pa</sub> also predominates over RecA<sub>Ec</sub> in presynaptic complex formation.

In principle, RecA<sub>Pa</sub> can form mixed filaments with RecA<sub>Ec</sub> though the interfaces of their protomers differ in nine positions

(5, 55). This statement follows from a high recombination activity of the RecAX45 chimera containing one interface from RecA<sub>Ec</sub> and the other complementary interface from RecA<sub>Pa</sub> (5, 38). Since RecA<sub>Pa</sub> has more affinity for ssDNA (4, 37) and forms more stable filaments (Fig. 4) than RecA<sub>Ec</sub> does, the mixed RecA<sub>Pa</sub>/RecA<sub>Ec</sub> presynaptic complexes appeared to be strong enough to realize hyperrecombination with  $\Delta$ FRE of 5 (Table 1) though less efficiently than pure RecA<sub>Pa</sub> filaments ( $\Delta$ FRE = 7.9 [Table 1]).

To answer the second question, we should consider two types of ends-in events: ss- or dsDNA breaks and take into account that (i) the former can be converted into the latter in a special enzymatic reaction (23) and (ii) the latter form the basis of recombination-directed replication (13, 20). However, for hyperrecombination when the number of ends-in events should increase 5, 8, or 26 times (5, 25), it seems reasonable to suggest SSGR (14) as the main mechanism in realization of hyper-rec ends-in events. In fact, RecA<sub>Pa</sub> effectively displaces SSB from ssDNA (4, 37) and thus from ssDNA gaps in donor and recipient DNA during conjugation, which results in the formation of presynaptic complexes initiating the SSGR process.

According to a common view, the SSGR mechanism is directed by the RecF pathway (2, 14). However, in the case of RecA<sub>Pa</sub>, the full activity of this pathway appeared to be unnecessary. Genetic analysis of the RecF pathway activity, measured through the yield of conjugal recombinants, on RecA<sub>Pa</sub>- or RecA<sub>Ec</sub>-mediated recombination showed that even deficiency of this pathway can be compensated, at least partially, by recombinogenic activity of the RecA<sub>Pa</sub> protein. This *in vivo* finding is in good agreement with the role of the RecOR protein complex in stabilization of RecA<sub>Ec</sub> filament recombination activity at its 5' end (7) and with the abilities of the RecA<sub>Pa</sub> filament (i) to initiate recombination from the 5' and 3' ends in an equal manner (Fig. 3) and (ii) to be more stable at the 5' end (Fig. 4). These observations are in concordance with the SSGR model predicted by Cromie and Leach (14) and the *in vivo* evidence for the abilities of both 3' and 5' ssDNA ends to stimulate recombination (40).

Recombinational characteristics of RecA803 (31) and RecA730 (26) (= RecA1211 [58]) proteins bearing amino acid substitutions V37M and E38K, respectively, were described earlier. Like RecA<sub>Pa</sub>, these point mutant RecA<sub>Ec</sub> proteins can partially complement the deficiency of RecF protein. Moreover, genetic analysis has revealed twofold and eightfold increases of FRE values for *recA803* and *recA730* (= *recA1211*) mutants, respectively (25; V. A. Lanzov and A. J. Clark, unpublished results). Interestingly, RecA<sub>Pa</sub> protein also has amino acid residue substitutions in the same positions (V37I and E38P). In fact, all residues in positions from 31 to 39 in RecA<sub>Pa</sub> are changed relative to those in RecA<sub>Ec</sub>. On the whole, this observation provides indirect evidence that loops 29 and 44 of the RecA protein structure elucidated by Story et al. (55) might be responsible for regulation of RecA interactions with the RecFOR proteins.

Qualitatively, some biochemical characteristics of RecA803 and especially RecA730, which are thought to be responsible for recombination, look like those of RecA<sub>Pa</sub> that are improved or enhanced relative to those of RecA<sub>Ec</sub>. The important difference between RecA<sub>Pa</sub> and RecA730 proteins in-

volves their relation to SOS induction. RecA730 promotes hyperrecombination mainly (but not fully) in an SOS-dependent manner (25), while RecA<sub>Pa</sub> generally does this independently of the SOS response. Nevertheless, the existence of these point mutant RecA<sub>Ec</sub> proteins help us to understand that the weak interconnections between RecA<sub>Pa</sub> and RecFOR do not reflect only the different origins of the proteins (the former is from *P. aeruginosa*, while the latter is from *E. coli*) but rather are based in the stronger recombinogenic activity of the RecA<sub>Pa</sub> protein. In fact, the RecFOR system of proteins exists in *P. aeruginosa* and in many other bacteria (45). However, to date, little is known about the interchangeability of RecFOR systems from different sources. It seems reasonable to suggest that the inherent recombinase function of RecA<sub>Pa</sub> has been more completely unmasked by evolution to provide efficient DNA repair as appropriate for a respiratory pathogen subjected to high levels of oxidative damage.

Given its close structural relationship to RecA<sub>Ec</sub>, RecA<sub>Pa</sub> provides an excellent system with which to investigate the structural basis of recombinase function. We anticipate that continued efforts to compare the two proteins, focusing on key amino acid residues already defined as important to the enhanced function of RecA<sub>Pa</sub>, will yield new insights.

#### ACKNOWLEDGMENTS

This work was supported by a Fogarty International Research Collaboration Award (2 R03 TW001318-04; to M.M.C.) and by grants from the Russian Foundation for Basic Research (05-04-48266; to V.A.L.), the Russian Ministry of Education and Science (RNP 2.2.1.1.4663), and the National Institutes of Health (GM32335; to M.M.C.).

We thank Leonid M. Firsov (PNPI) for fruitful discussions of some experiments. D.M.B., I.V.B., and Y.V.K. acknowledge with gratitude M. Cox's research group for providing assistance with the portion of the work carried out in the United States.

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