

Feature Review

RecA: Regulation and Mechanism of a Molecular Search Engine

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Homologous recombination maintains genomic integrity by repairing broken chromosomes. The broken chromosome is partially resected to produce single-stranded DNA (ssDNA) that is used to search for homologous double-stranded DNA (dsDNA). This homology driven ‘search and rescue’ is catalyzed by a class of DNA strand exchange proteins that are defined in relation to *Escherichia coli* RecA, which forms a filament on ssDNA. Here, we review the regulation of RecA filament assembly and the mechanism by which RecA quickly and efficiently searches for and identifies a unique homologous sequence among a vast excess of heterologous DNA. Given that RecA is the prototypic DNA strand exchange protein, its behavior affords insight into the actions of eukaryotic RAD51 orthologs and their regulators, BRCA2 and other tumor suppressors.

RecA: A Molecular Search Engine

Escherichia coli RecA is the defining member of an ancient and ubiquitous clade of DNA strand exchange proteins that are essential for homologous recombination [1]. This clade consists of three distinct families: RecA, RAD51, and the RAD51 paralogs (Figure 1A) [1]. RecA is found in all free-living bacteria, and is the most slowly evolving gene involved in DNA metabolism, with an average sequence conservation of approximately 60–70% across the entire bacterial domain of life [2]. Many bacteriophage also have homologs from phylogenetically defined RecA subfamilies: the Phage SAR1, Phage SAR2, and Phage UvsX, although only phage T4 UvsX has been isolated and studied biochemically (the others were only identified in metagenomic data) [3,4]. The RecA–SAR1 subfamily is likely an ancient paralog of RecA not found in cultured/studied bacteria. Plants – but neither animals nor fungi – have both mitochondrial and chloroplast specific variants of RecA of unknown functional importance, indicating that these genes were acquired through endosymbiotic transfer by an ancestor common to eukaryotes, but then lost during evolutionary divergence [4]. All Archaea and Eukarya share two classes of RecA homologs [1]. The RAD α class is functionally homologous to RecA and includes archaeal RadA, eukaryotic RAD51, and eukaryotic DMC1, the last of which functions specifically during meiosis [5,6]. The RAD β class is a divergent class of proteins that genetically function with RAD51 and are collectively known as the RAD51 paralogs. These paralogs are RAD51B, RAD51C, RAD51D, XRCC2, and XRCC3 proteins in mammals, Rad55 and Rad57 as well as Csm2 and Psy3 in fungi, and RadB in archaea; recently, RadA of *E. coli* (not to be confused with RadA of Archaea), which is a member of this RAD β class, was shown to be a novel branch migration enzyme [7]. Many of the eukaryotic paralogs form various subcomplexes [8]. Although recent biochemical analysis has demonstrated that the yeast Rad55/57 complex promotes homologous recombination by antagonizing an antirecombination helicase (Srs2) [9], the biochemical function(s) of the human paralogs remain unclear.

Trends

Single molecule methods have yielded new insight into how RecA filaments form and find homologous DNA.

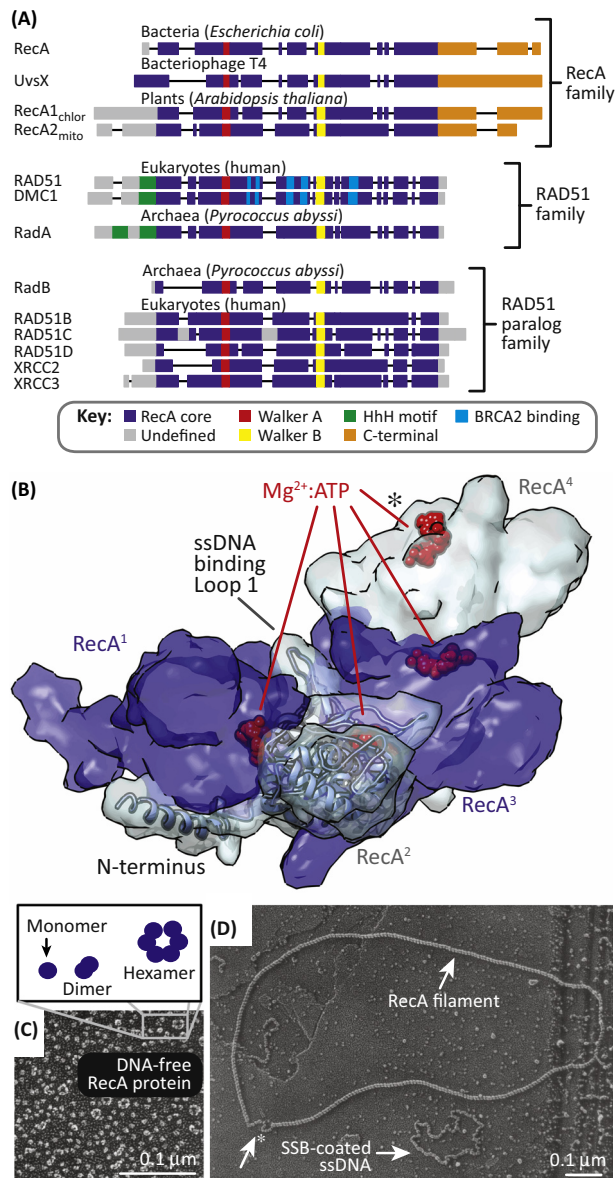
RecA nucleation and growth on SSB-coated ssDNA is kinetically regulated by the RecFOR and RecOR complexes, which both microscopically and macroscopically alter the SSB–ssDNA nucleoprotein fiber.

RecA finds homology through many random, weak, and transient interactions made in parallel and ignores very short regions of microhomology resulting in a reduction of both dimensionality and complexity.

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Figure 1. RecA Conservation and Structure. (A) Conserved domains of the RecA, RAD51, and RAD51 paralog families. (B) Segment of RecA filament showing the Mg²⁺:ATP binding site at the monomer–monomer interface. The asterisk indicates a half-site; for simplicity, only one of ssDNA-binding loops per monomer is shown. Adapted from PDB 3CMW (RecA) [30]. (C) Electron microscopy of DNA-free RecA showing heterogeneous oligomerization states of RecA. (D) Electron microscopy of RecA filament formation on SSB-coated circular ssDNA. Note the dramatic extension of the RecA filament relative to the compaction of the SSB-coated ssDNA. (C, D) Adapted from [39]. Abbreviations: ssDNA, single-stranded DNA; SSB, ssDNA-binding protein.

Failure to properly regulate the assembly of human RAD51 causes an accumulation of mutations that accelerate tumorigenesis in individuals with genetic defects in BRCA2 and its epistasis group, which includes the five RAD51 paralogs (RAD51B, RAD51C, RAD51D, XRCC2, and XRCC3), PALB2, and some proteins of the Fanconi Anemia core complex [8,10]. This accelerated mutation and tumorigenesis is due to the loss of homologous recombination-dependent repair, often in conjunction with – or downstream of – DNA repair pathways that correct specific

types of damage, including crosslinks, alkylations, bulky adducts, single-stranded DNA (ssDNA) gaps, and double-strand breaks [11]. Although this review will focus on *E. coli* RecA, many of the biochemical properties of RecA are fundamentally similar to human RAD51 [5], and studies on RecA continue to inform our understanding of the molecular details of the core homologous recombination machinery across all domains of life [12].

In vivo, RecA preferentially binds to ssDNA that is generated through the resection of double-stranded DNA (dsDNA) breaks, or the formation of ssDNA gaps, which arise when replication forks stall or collapse. To initiate homologous recombination, helicases and/or nucleases process a broken chromosome or stalled replication fork to produce ssDNA, which is rapidly coated with – and continuously sequestered by – ssDNA-binding proteins (SSB, in bacteria). An essential step during homologous recombination is the assembly of a presynaptic filament of RecA on the ssDNA of the broken chromosome, which in turn uses the ssDNA sequence to search for a homologous region in the dsDNA genome. Importantly, bacteria are either haploid or merodiploid (partially diploid) organisms, and therefore use the sister chromatid as the repair template during replication but before the cell has divided. Formation of the RecA filament triggers the SOS response through the autocatalytic proteolysis of the LexA repressor via a direct LexA–RecA–ssDNA complex. In addition to inducing the expression of DNA repair proteins and activating translesion polymerases, the SOS response delays cell division, temporarily increasing the copy number of the chromosome and causing physiological filamentation of *E. coli*. Recent *in vivo* super-resolution imaging of RecA filaments demonstrates that the RecA filament (or bundles of filaments) transverses the mother and daughter cells in response to DNA damage [13].

RecA filament assembly in response to a dsDNA break is not spontaneous, but rather tightly regulated by RecBCD, a helicase/nuclease that processes dsDNA breaks and directly loads RecA onto ssDNA upon encountering a bacterial self-recognition sequence called χ (*crossover hotspot instigator*, Chi, 5'-GCTGGTGG-3'), which is spaced approximately every 4–5 kb throughout the *E. coli* genome [14]. When the χ sequence is not present, RecBCD will rapidly and processively degrade linear dsDNA (up to 30 kb during an average processing event *in vitro*, but ~10 kb *in vivo* [15]) to defend against bacteriophage infection. Interestingly, the degradation products generated by RecBCD can be used by the CRISPR system to identify foreign DNA sequences during 'adaptation', when protospacer sequences are acquired [16]. During DNA replication, RecA filaments are assembled on ssDNA gaps and breaks that form when replication forks stall or collapse upon encountering either chemical damage (nicks, crosslinks, adducts, etc.), or in some cases, transcriptional impediments (e.g., collision with RNA polymerases) [17]. These so-called daughter strand gaps are processed not by RecBCD (which requires a nearly blunt DNA end to initiate) but rather by the concerted action of RecQ helicase and RecJ nuclease, which produces long regions (i.e., several thousand nucleotides) of SSB-coated ssDNA tails or gaps (i.e., an ssDNA region flanked by either one or two segments of dsDNA) [18,19]. Unlike RecBCD, neither RecQ nor RecJ are reported to directly interact with RecA during processing; rather, RecA is loaded onto SSB–ssDNA by the RecF, RecO, and RecR proteins [20–22]. In this context, both the RecFOR and RecOR complexes regulate RecA filament assembly by enhancing nucleation and growth of the filament through structural perturbation of the SSB-coated ssDNA nucleoprotein (i.e., protein–DNA) complex and/or altering the kinetics of SSB sliding/wrapping on ssDNA [23,24]. Here, we aim to provide a synthesis of our current understanding of the mechanism of RecA, which is illuminated by both a rich history of genetic and biochemical research, as well as recent single molecule experiments.

The Search for *rec* Genes: Identification and Biochemical Isolation of RecA

Clark and Margulies first identified *recA* in 1965 in a genetic screen where they mutagenized F^- bacteria and screened for strains unable to recombine with an Hfr donor strain [25].

Subsequent genetic analysis revealed that *recA* is essential for the RecBCD and RecF pathways of homologous recombination and that *recA* mutants are extraordinarily pleiotropic, affecting DNA repair, SOS mutagenesis, induction of λ prophage, cell division, and chromosomal segregation [26]. A decade after the discovery of the *recA* gene, the protein was first purified and a wide range of biochemical activities were reported, including DNA binding, transactivation of autoproteolysis of the LexA and λ phage repressors, ATP hydrolysis, the ability to form filaments on DNA, and the capacity to exchange homologous DNA strands [26–28]. Owing to a rich history of elegant genetic and biochemical investigation of RecA, many of the biochemical and biophysical properties of RecA can be summarized in relation to the physiology of *E. coli* (Table 1).

RecA is a 38-kDa protein that is basally expressed at approximately 1000 monomers per cell and forms right-handed nucleoprotein filaments on ssDNA and dsDNA, although substantially more slowly on the latter. The *in vivo* concentration of RecA is $\sim 1 \mu\text{M}$ in the absence of DNA damage; however, the concentration is increased approximately 10-fold during the SOS response [28]. RecA has a potent ATP hydrolysis activity in the presence of DNA, a result of a complex and dynamic cycle of filament assembly and disassembly on DNA [29]. The nucleoside triphosphate-binding site lies between adjacent monomers in a filament (Figure 1B), and the hydrolyzed state of the nucleotide strongly affects the stability of the RecA–DNA complex [30]. The ATP-bound form has a high affinity for DNA, while the ADP-bound form has a lower affinity, and, therefore, is both thermodynamically and kinetically less stable. The ATP-bound nucleoprotein filament serves as a surface catalyst for the search and capture of a homologous sequence of DNA, a process known as synapsis. Once a region of homology is found, the ssDNA strands on the homologous chromosomes are exchanged, producing heteroduplex DNA (i.e., each single strand originated from a different chromosome). Although RecA must bind ATP to form an active filament, hydrolysis is not required to catalyze synapsis or exchange the DNA strands to produce heteroduplex DNA; however, hydrolysis is required for filament disassembly [29,31]. The progression of these steps – synapsis and strand exchange – produces a complex, metastable intermediate of recombination known as a joint molecule that is then repaired through a combination of DNA synthesis, ligation, and resolution [11].

The Search for Structural Clues: Insight from Electron Microscopy and X-Ray Crystallography

Although the first crystal structure of RecA was solved in 1992 by Story *et al.* [32], detailed structural information of the protein in complex with ATP and either ssDNA or dsDNA was remarkably elusive. For nearly three decades, most of the structural information on the RecA nucleoprotein filament came from high-resolution electron microscopy, which yielded tremendous insight into the morphological changes in the nucleoprotein structures during recombination (Figure 1C,D) [33–35]. Under physiological conditions, ssDNA folds into a multitude of structures that are stabilized by localized annealing of short regions of complementarity, called secondary structure, which inhibit RecA filament formation [23]. DNA secondary structure is largely overcome by the SSB protein, which binds tightly to ssDNA, effectively denaturing the secondary structure [36]. SSB-coated ssDNA is compacted relative to dsDNA and is morphologically distinct from RecA filaments because each tetramer of SSB wraps a segment of 30–70 nucleotides around itself (Figures 1D and 2A) [35,37]. When a molecule of ssDNA is completely and contiguously coated with RecA in the presence of ATP γ S [adenosine 5'-(gamma-thio)triphosphate, a nonhydrolyzable ATP analog], the ssDNA is extended by 150–160% relative to a molecule of dsDNA of the same length, has one RecA monomer bound for approximately three nucleotides, and has a helical pitch of 9–10 nm (~ 6 monomers per turn) with a diameter of ~ 10 nm [38]. When the filament is strictly in the ADP-bound form, the pitch is reduced to 6–7 nm and the diameter expands to ~ 12 nm [39]. When RecA filaments are formed

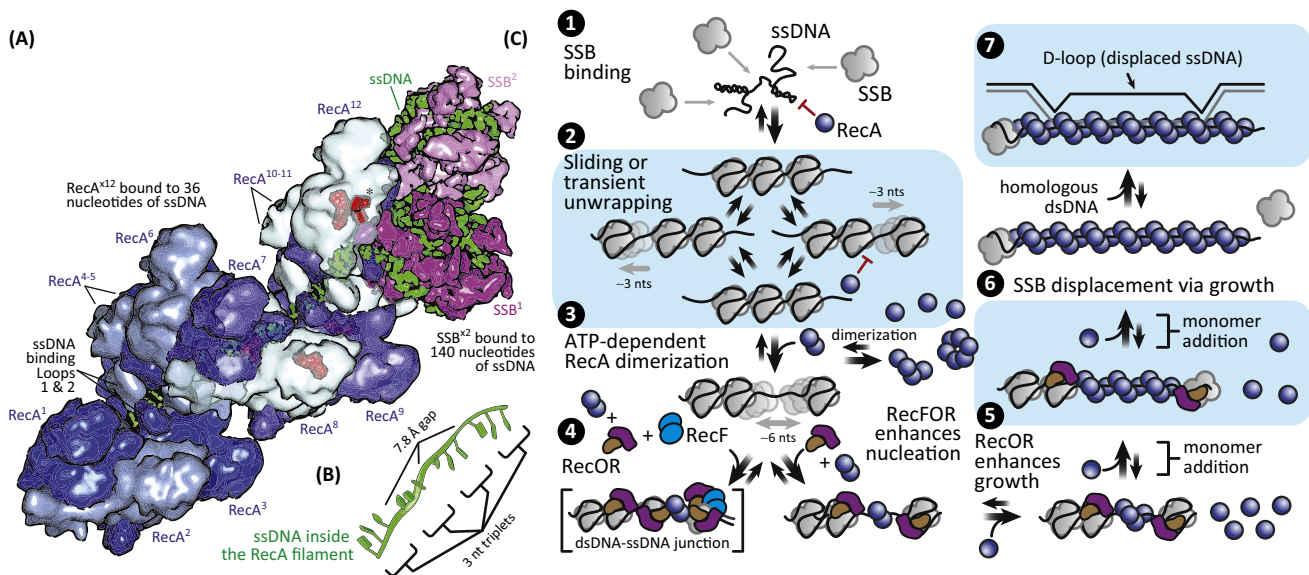
Table 1. RecA Filament Assembly and Synapsis during Replication and Homologous Recombination

Rates and Relevant Physical and Physiological Measurements		Refs
Volume of an <i>E. coli</i> cell	~1 f l (1×10^{-15} l)	[103]
<i>E. coli</i> genome size	4.7 Mb (4.7×10^6 bp)	[104]
Replication rate	650–800 bp/s	[105]
ssDNA generated at replication fork (average Okazaki fragment size)	1000–2000 nucleotides	[106]
Average size of daughter strand gaps	100–800 nucleotides	[107,108]
dsDNA breaks per division (average)	0.1–1	[28,109]
(maximum tolerated)	< 3	[110]
DNA crosslinks per division (maximum tolerated)	50–70	[111]
Oxidative lesions per division	~2000	[28]
Rate of RecBCD resection	1000–2000 bp/s	[14]
Average χ (Chi) frequency	1 per 4500 bp	[14]
Average length of dsDNA resection by RecBCD (<i>in vitro</i>)	30 000 bp	[14]
(<i>in vivo</i>)	10 000 bp	[15]
SSB site size per tetramer	30–70 nucleotides	[37,112]
RecA site size per monomer	3 nucleotides	[29]
Persistence length of dsDNA	~50 nm	[80]
Persistence length of ssDNA	~1 nm	[80,82]
Persistence length of RecA–ssDNA	~900 nm	[44]
Radius of gyration (R_g) for λ dsDNA (48.5 kb)	~900 nm	
RecA nucleation time (rate) ^a , spontaneous	10–60 min (1–6 nuclei/h)	[24]
RecOR-mediated	5–30 min (2–12 nuclei/h)	
RecFOR-mediated	2–10 min (6–30 nuclei/h)	
RecA growth rate ^a , spontaneous	0.3–1.3 RecA monomers/s	[24]
RecOR-mediated	2–6 RecA monomers/s	
RecFOR-mediated	2–6 RecA monomers/s	
RecA K_d for ATP	~15 μ M	[113]
RecA K_d for ATP (+ssDNA)	~2.5 μ M (ssDNA)	
RecA K_m for ATP (+ssDNA)	~20 μ M (ssDNA)	[114]
(+dsDNA)	~100 μ M (dsDNA)	
RecA k_{cat} for ATP (+ssDNA)	~21 per min per RecA	[56]
for dATP (+ssDNA)	~33 per min per RecA	
RecA–ssDNA complex salt titration midpoint ^b , 0.1 mM nucleotide cofactor	255 mM NaCl ~400 mM NaCl (+ATP) 165 mM NaCl (+ADP)	[115]
RecA–dsDNA complex salt titration midpoint ^c , 1 mM nucleotide cofactor	~300 mM NaCl (+ATP) 190 mM NaCl (+ADP)	[116]

^aNucleation and growth rates reported were measured in the presence of ATP γ S (1 μ M RecA, 2 mM ATP γ S, pH 7.5, 37°C, ~8000 nt substrate) [24]. The numbers in Table 1 represent our best estimate for physiologically relevant nucleation and growth rates in the presence of ATP (instead of ATP γ S), which is ten times slower for nucleation and two-thirds slower for growth. However, these rates are affected by temperature, pH, excluded volume, and concentrations of proteins, and monovalent, divalent, and trivalent salts in nonlinear ways. Nonetheless, these estimates are consistent with ensemble experiments with RecFOR [22] and *in vivo* imaging of RecA bundle appearance and growth [13].

^b20 mM Tris acetate, pH 7.5, 4 mM Mg(OAc)₂, 25°C.

^c20 mM MES, pH 6.2, 10 mM MgCl₂, 25°C.



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Figure 2. Mechanism of RecA Nucleation and Growth on SSB-Coated ssDNA. (A) Structural model of 12 monomers of RecA bound to 36 nucleotides of ssDNA adjacent to two tetramers of SSB bound to 140 nucleotides of ssDNA demonstrating the vast differences in site size and accessibility of the ssDNA bound to each protein. Mg: ATP (red) is visible at the monomer-monomer interface for RecA⁷–RecA¹². Adapted from PDB 3CMW (RecA) and PDB 1EYG (SSB) [30,37]. (B) ssDNA within the filament is stretched into nucleotide triplets that can maintain Watson–Crick interactions during homologous pairing. (C) (1) SSB binds rapidly to ssDNA, removing secondary structure that impedes RecA-mediated DNA strand exchange. (2) SSB kinetically blocks RecA filament formation. (3) Nucleation of RecA onto rare and transient microscopic gaps requires ATP-dependent dimerization, making nucleation infrequent. (4) RecOR binds to the C-terminal tails of SSB, microscopically altering the SSB–ssDNA complex, but not displacing it. This ‘stoichiometric remodeling’ creates microscopic gaps that are large enough and long-lived enough for RecA to stably bind, enhancing nucleation. Similarly, RecF further, in coordination with RecOR, enhances RecA nucleation at dsDNA–ssDNA junctions. (5) RecA filament growth through monomer addition is impeded by SSB, although less so than nucleation; however, in the presence of RecOR, RecA filament growth is stimulated ~3-fold. (6) The RecA filament grows monotonically and displaces SSB from ssDNA. Adapted from [24]. (7) The RecA–ssDNA filament catalyzes pairing and strand exchange with a homologous dsDNA molecule, resulting in an intermediate, three-stranded molecule called a D-loop (or ‘displacement loop’); SSB binds to the displaced strand to stabilize the D-loop (not shown). Abbreviations: dsDNA, double-stranded DNA; ssDNA, single-stranded DNA; SSB, ssDNA-binding protein.

with ATP and hydrolysis results in ADP accumulation, RecA filaments are heterogeneous in pitch even along a single filament, consistent with a model of localized, cooperative ATP hydrolysis [38]. The failure of mutant RecA proteins (exemplified best by RecA 142 [40]) to stretch ssDNA to the canonical filament length, despite their ability to bind to ssDNA and hydrolyze ATP, is one of the strongest morphological factors that correlates with an inability to catalyze synapsis and DNA strand exchange, showing that stretching DNA is an essential component of DNA strand exchange [29,41].

In 2008, Chen *et al.* solved the crystal structure of both the presynaptic (ssDNA-bound) and postsynaptic (dsDNA-bound) RecA filaments, revealing surprising mechanistic details (Figure 1B and Figure 2A) [30]. While thought to be relatively uniform (or isotropic) throughout the complex, the DNA stretched inside the RecA filament is in fact segmented into nucleotide triplets that maintain approximately normal B-form dimensions (~3.2–3.5 Å for dsDNA and 3.5–4.2 Å for ssDNA) but are separated by an internucleotide extension of ~7–8 Å (Figure 2B) [30]. Biochemical analysis had previously demonstrated that ATP binding is necessary and sufficient to promote DNA strand exchange, whereas hydrolysis of ATP is dispensable [31]. This observation substantiated the hypothesis that synapsis is ‘simply’ a bimolecular collisional process between the RecA–ssDNA nucleoprotein filament and the homologous genomic locus (see subsequent section on ‘Search optimization through parallel processing: RecA finds homology by simultaneously sampling many transient and weak contacts’), rather than a process that is coupled to some putative ATP-dependent directed motion. The presynaptic and postsynaptic crystal

structures demonstrated how this is possible [30]. By using the free energy of ATP binding to stretch and untwist the ssDNA within the RecA filament, binding of the incoming dsDNA to the secondary site within the filament is stabilized through base pairing interactions only when a sufficient amount of homology is attained (as few as 8 bases). By maintaining these B-form triplet states (Figure 2B), the RecA filament creates a series of genetic ‘words’ that are held in register to one another throughout a contiguous filament allowing the nucleotide triplets to test for localized homology through normal, although transient, base pairing [30]. Each RecA monomer interacts with the entire triplet closest to itself in the structure, as well as with two more nucleotides, one from each of the preceding and following triplets. As a result, each nucleotide triplet is bound by three monomers, and each monomer interacts with five nucleotides, but because the first and fifth nucleotides are obligatorily ‘shared’ with adjacent monomers, the net stoichiometric ratio (or binding site size) is three nucleotides per RecA. In a similar manner, the dsDNA in the postsynaptic filament is also stretched into nearly perfect Watson–Crick base-paired triplets [30].

The structural elucidation of the RecA–ssDNA complex was a substantial breakthrough that advanced our understanding of how DNA is recognized at the base-pair level during genetic recombination. However, how RecA filament assembly is regulated and whether the collisional, diffusion-driven model for homology search was a correct or feasible model remained open questions that traditional, ensemble biochemical approaches could only partially address.

The Search for Dynamics: Single Molecule Microscopy Brings RecA Assembly and Search Mechanisms into Focus

Single molecule techniques are a diverse and powerful toolbox with which to probe biological function with high spatiotemporal resolution. The first reported experiment to use a single molecule assay to probe the dynamics of RecA was performed in 1998 by John Marko’s research group using a glass fiber attached to one end of a single dsDNA molecule and a bead held by a micropipette at fixed tension at the other end [42]. By pulling on the molecule with the micropipette, the authors could accurately measure the force required to stretch the DNA in the presence and absence of RecA by observing the displacement (i.e., bending) of the glass fiber, which acted as a force transducer. Polymerization of RecA onto the dsDNA resulted in extension of the DNA molecule, which was observed in real time. Other groups reported similar experiments shortly thereafter, where a single DNA molecule was attached to the surface of a piezo-driven stage at one end and a bead at the other [43,44]. The bead was held in place by an optical trap that was calibrated to measure either the change in force upon displacement, or the displacement under constant force. Like the previous study, the authors were able to measure extension of a single RecA–DNA molecule in real time as well as the force–extension relationship of filaments; however, they expanded upon the previous work by investigating the role of various nucleotide cofactors in filament assembly and by monitoring filament disassembly. Despite the advance presented by single molecule manipulation, the ability to distinguish between the nucleation and growth phase of filament assembly remained elusive – in part because the DNA molecule used was very long and multiple filaments could simultaneously nucleate and grow on a single molecule of DNA.

In 2006, two independent groups used very different single molecule assays to differentiate between these kinetic parameters of nucleation and growth. The first method used total internal reflection fluorescence (TIRF) microscopy to detect Förster resonance energy transfer (FRET) between a donor (Cy3) fluorophore that had been attached to the 5′-end of a dsDNA–ssDNA junction and an acceptor (Cy5) fluorophore that had been incorporated internally into an ssDNA overhang 13 nucleotides away (although the total length of the overhang was greater). In the absence of RecA, the short persistence length (and therefore high flexibility) of ssDNA results in high FRET efficiency, whereas formation of the filament results in an increased persistence length

that diminishes the FRET efficiency approximately 4-fold or greater by physically separating the donor and acceptor molecules [45]. When RecA filaments were at steady-state in this assay, approximately four or five quantized FRET states of varying efficiency were observed depending on the length of the ssDNA substrate. Each quantized state was dependent on the number of RecA monomers bound between the donor and acceptor molecules and the authors were able to extract kinetic parameters from thousands of transitions between states, on hundreds of individual molecules, counting the frequency of addition and subtraction of monomers in time.

At essentially the same time, a complementary direct visualization assay was developed [46]. Individual clusters of RecA filaments could be observed directly while forming and growing on dsDNA, which was attached at one end to a polystyrene bead immobilized in an optical trap within a multichannel, laminar flow cell [47]. An epifluorescence microscope was used to visualize fluorescent RecA bound to dsDNA tethered to the bead, which could be transferred between channels through the movement of the microscope stage. This approach is remarkably powerful as a result of the relative ease with which basic biochemical parameters such as concentration, cofactor, or salt can be varied, while simultaneously maintaining tight control of the incubation time for each individual molecule. These experiments provided three major insights into the mechanism of RecA filament assembly. First, that the nucleation kinetics of individual clusters forming on dsDNA follow a power dependence, where the rate increases proportionally to the fourth to fifth power of RecA concentration [$k_{\text{nuc-dsDNA}} \propto (\text{RecA})^{\sim 4-5}$], indicating that approximately four to five monomers of RecA are the nucleation unit when binding to dsDNA. Second, that RecA filament growth is bidirectional on dsDNA, where the rate of filament growth at internal locations on the DNA is approximately 2-fold faster than filaments growing at the distal end of the DNA molecule. Third, that RecA filaments could be formed in the presence of ATP and stabilized by either ATP γ S or ATP:Ca²⁺, providing direct evidence that the nucleotide cofactor can be exchanged within the filament without disassembly.

Searching for Control in the Face of Competition: Regulation of RecA Filament Assembly on SSB-Coated ssDNA

Within its cellular context, RecA filaments must nucleate and grow on ssDNA in direct competition with SSB, which rapidly and contiguously sequesters ssDNA generated during replication and nucleolytic processing of damaged DNA. This competition serves two basic functions: first, it suppresses unwanted and potentially detrimental recombination. Second, SSB denatures localized DNA secondary structure that impedes contiguous RecA filament formation, which in turn reduces the efficiency of DNA strand exchange. Although methods for measuring the nucleation and growth of other filament forming proteins – actin and tubulin – have existed for several decades, the complexity of forming a filament on a linear template in the presence of a contiguous kinetic competitor made quantitative measurements of RecA assembly difficult. To accomplish this, long ssDNA molecules were attached to a surface within a flow cell and the nucleation and growth of filaments in the presence of SSB was directly imaged using TIRF microscopy, which enabled the unexpectedly long time-lapse experiments required to observe filament formation [24].

Several features of RecA filament assembly and regulation were revealed by this approach. First, the nucleation rate of RecA increased in proportion to the second power of the RecA concentration [$k_{\text{nuc-ssDNA}} \propto (\text{RecA})^{\sim 2}$], indicating that a dimer of RecA – the minimal oligomer that can bind ATP – is the critical nucleus required for stable filament formation [24,30]. This smaller critical nucleus required for binding to ssDNA (dimers) relative to dsDNA (tetramers to hexamers) is presumably due to the higher affinity of RecA for ssDNA, an interpretation that is supported by the observation that nucleation on dsDNA was extremely sensitive to low concentrations of salt, while nucleation on SSB-coated ssDNA was largely insensitive over this same range [24,46,48]. Second, RecA filament growth on ssDNA is rapid and bidirectional, although 50–60% faster in

the 5'→3' direction, and proceeds through monomer addition to the ends of the filaments coupled to a conformational change that is slower than the diffusion limit. Third, nucleoside triphosphate binding activates a high-affinity state, acting as a conformational effector that increases the success of a nucleation event by stabilizing capture of the ssDNA transiently released from SSB during sliding or unwrapping. This is supported by the observation that RecA nucleation is faster when ATP is replaced with either ATP γ S or dATP, which is hydrolyzed ~30% faster than ATP. The observed increase in nucleation rate is energetically linked to the increased affinity of RecA for ssDNA when bound to the nucleotide or analog. Therefore, both the ability to hydrolyze ATP and the rate at which it is hydrolyzed are dispensable properties with respect to assembly, although they are essential properties for disassembly during DNA strand exchange. Fourth, both nucleation and growth of RecA are severely depressed at physiological pH (i.e., ~7.5). If RecA filament assembly were intrinsically at its maximum, then the cell would have little means by which to regulate its activity. This finding therefore rationalizes the necessity of mediator proteins – RecFOR and RecOR – to potentiate filament assembly by binding to SSB via RecO to trap ssDNA transiently released during SSB sliding or unwrapping (Figure 2C) [24]. Although RecOR and RecF had long been known to stimulate RecA filament formation, the mechanism of this stimulation remained unclear. In the single molecule experiments described earlier, the addition of RecOR reduced the lag time for RecA filament nucleation and stimulated the rate of growth. The addition of RecF to the RecOR-stimulated reaction further reduced the lag time, but had no effect on growth. Hence these experiments were able to clearly distinguish, for the first time, the relative contributions of the so-called 'recombination mediator proteins' in the kinetic regulation and stimulation of RecA filament formation [24].

Although the intracellular pH is highly regulated in *E. coli*, it can be altered [49,50]. For example, exposure to lipophilic acids such as propionic acid or sodium benzoate induce a stable reduction in the intracellular pH to ~6.5–6.8 that is toxic to cells if prolonged [49,51,52]. The expression profile of many bacterial genes, including the SOS response genes, is altered by environmental pH [53,54]. In fact, lowering the intracellular pH was used to reactivate a pH-sensitive mutant of RecA (*RecA142*) that is inactive at normal physiological pH, but active at lower pH [53,54]. Therefore, it is interesting to speculate that the alteration of filament assembly by solution conditions could be one of the many mechanisms by which bacteria respond to environmental stress, wherein the chemical environment (i.e., pH, osmolality, metabolites, etc.) might directly modulate the DNA damage response. Recent single molecule FRET experiments have further demonstrated that the pH-induced rate changes to filament growth and disassembly are moderate; however, the effect is amplified owing to both an acceleration of binding and a reduction in dissociation as pH decreases [55]. Similarly, it was proposed that the enhanced activity of RecA in the presence of dATP may be a reflection of an *in vivo* mechanism of biochemical regulation [56]. This interpretation is supported by the observation that the intracellular pool of dATP is increased during the bacterial DNA damage-induced SOS response due to a shift in ribonucleotide and deoxyribonucleotide concentrations mediated by the upregulation of ribonucleotide reductase (RNR) [57]. Nonetheless, the primary biological mechanism for filament regulation in *E. coli* is through the concerted action of RecFOR and RecOR, the latter of which binds to the autoregulatory C-terminal tail of SSB [20,21,23,24].

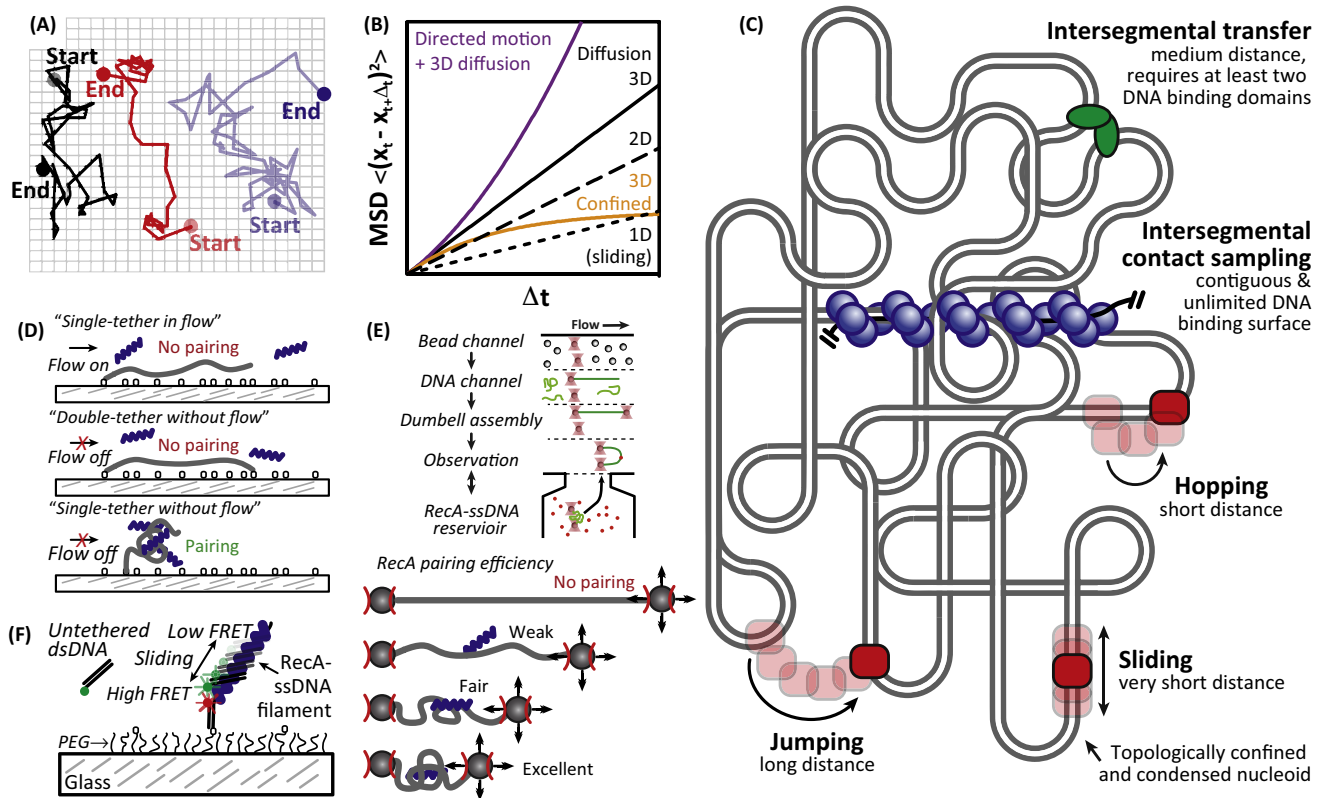
Search Optimization through Parallel Processing: RecA Finds Homology by Simultaneously Sampling Many Transient and Weak Contacts

Once a RecA filament has formed on a resected chromosome, it must use the sequence information encoded within the single strand of DNA within the nucleoprotein filament to find a homologous dsDNA region and then exchange the individual strands [29]. This biochemical process is unique to the RAD α subclass, but elements of this process are now recognized as being employed by CRISPR systems to uniquely target any sequence in dsDNA using a protein-bound ssRNA, rather than ssDNA, sequence. Although the search mechanism might seem

simple and intuitive, the task of finding a homologous locus within the short time window required for DNA repair is challenging. This difficulty arises due to the vast excess of heterologous DNA within the genome – a challenge that scales exponentially with genome size, but may be overcome through organizing principles that establish chromatin territories within the eukaryotic nucleus [58,59]. To access this information, RecA uses two DNA-binding sites: a primary site that stretches the ssDNA within the nucleoprotein filament and a weaker affinity secondary DNA-binding site that lies proximal to the ssDNA within the interior of the filament groove [60]. When homologous dsDNA comes in close contact with the stretched ssDNA within the groove of the filament, it can randomly sample homology through either canonical or noncanonical base pairing until a sufficient threshold (8 bases) stabilizes the interaction and permits DNA strand exchange products to be long-lived (~26 base pairs) [61]. DNA strand exchange is favored by virtue of the higher affinity displayed by the secondary site for the product of DNA strand exchange (the displaced ssDNA) rather than the substrate (the complementary dsDNA) [60]. This was clarified by the structural studies described earlier in this review [30]; however, the mechanism by which RecA kinetically searches for its homologous target was, until recently, largely a black box [29,62–65].

By contrast, how a site-specific DNA-binding protein can find a particular DNA sequence is relatively well understood, as many DNA-binding proteins (e.g., transcription factors and restriction enzymes) and specific structural motifs (zinc fingers, TALENs, etc.) that enable a protein to bind to a specific sequence have been extensively characterized both kinetically and structurally [66–69]. These processes are largely driven by a diffusive, random walk (shown schematically in Figure 3A) until a collision between the protein and its target site results in a successful binding event [70–74]. How quickly a target can be found is dependent on the degrees of freedom of the random walk (i.e., 1D sliding along a lattice, 2D diffusion on a plane, or 3D diffusion in space) and whether the diffusion is facilitated and/or directed (i.e., coupled to an accelerating factor such as an electrostatic potential) or confined (Figure 3B) [71,72,75].

More than three decades ago, Otto Berg, Robert Winter, and Peter von Hippel proposed three major diffusion-driven mechanisms that could explain the partitioning kinetics between specific, high-affinity binding to a target site and nonspecific, low-affinity binding to random DNA (Figure 3C) [71–75]. These general search mechanisms can be thought of as a model for all protein–DNA interactions, with only a handful of special case exceptions. First, in 1D diffusion, a protein can slide along a DNA lattice (i.e., a DNA polymer composed of repetitive and identical units) by randomly stepping away from – and back towards – its initial binding site. Sliding is theoretically limited to sampling only a small segment of DNA because – on average – the protein would randomly step backwards as frequently as it steps forward, requiring long lifetimes for large mean squared displacements. Second, proteins can ‘hop’ along DNA by iteratively binding and dissociating. Each release will be followed by 3D diffusion; however, because DNA behaves as a rod over short distances, rather than a point source, the probability of rebinding to an adjacent site close to the point of dissociation is statistically high. Therefore, this correlation between unbinding and rebinding results in a statistically probable ‘hop’ within ~10–20 bp [70]. Alternatively, *in vivo*, because the genome is tightly compacted within a confined space [76–78], there is a high probability of a new binding event being far away when measured along the linear molecule (e.g., tens of kilobase pairs to several megabase pairs), but which is very close in 3D space. Such an event is simply normal dissociation followed by rebinding (and therefore microscopically indistinguishable from ‘hopping’), but because of the dense coiled nature of DNA, the term ‘jumping’ has been attributed to events associated with rebinding events that are very far away from the point of dissociation on the linear chromosome [79]. Finally, if a protein has at least two or more DNA-binding domains, then it can iteratively bind and release via one domain while remaining tethered to the DNA through another – a phenomenon called ‘inter-segmental transfer’ or ‘looping’. This transfer can occur between segments that are close in the



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Figure 3. Diffusion-Driven Mechanism of RecA-Mediated Homology Search. (A) Diagram of three particles demonstrating random walk diffusion represented on a 2D plane. Adapted from [117]. (B) Plot showing the mean squared displacement as a function of time for 1D, 2D, and 3D diffusion compared with 3D diffusion in a confined space without or with directed motion. Adapted from [99,118]. (C) Cartoon depicting different modes by which proteins find their targets by sliding, hopping, jumping, intersegmental transfer, and intersegmental contact sampling [70–72,75]. Single molecule methods used to measure RecA-mediated homology search. (D) Total internal reflection fluorescence (TIRF) microscopy used to visualize ssDNA–RecA filaments pairing with λ phage DNA [86]. (E) DNA micromanipulation experiments demonstrating RecA pairing efficiency increases as the DNA is allowed to adopt 3D, random-coil configurations [86]. (F) Single molecule Förster resonance energy transfer (FRET) experiments used to demonstrate microscale sliding of RecA filaments [89]. Abbreviation: ssDNA, single-stranded DNA.

linear arrangement of DNA or between segments that are widely separated in the genome. In each of these models, the target site is short and rare, but is bound tightly and specifically when it is found and is often linked to a regulatory function such as binding to a regulatory metabolite or protein partner [71].

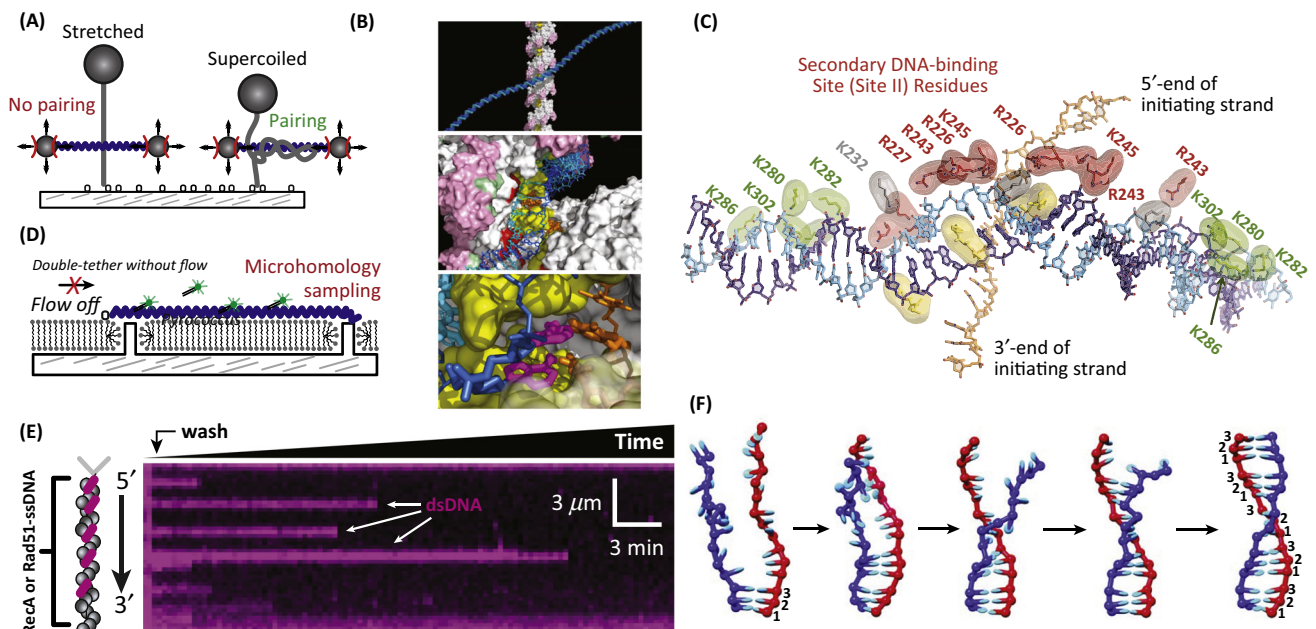
The RecA–DNA complex is not a simple, compact entity; rather, it is a long partially flexible filament that has a persistence length (i.e., a measure of polymer stiffness) of ~ 900 nm for the static ATP γ S filament [44], which is approximately 20 times greater than dsDNA (45–50 nm [80,81]) and several hundred times greater than ssDNA (0.6–3 nm [44,82,83]) (Table 1). So if diffusion of a RecA filament is severely restricted through physical confinement, how then does RecA find its homologous partner? Although it was over three decades ago that RecA was found to assimilate ssDNA into homologous duplex DNA [84,85], the details of the kinetic intermediates formed and the transient physical mechanics of the RecA–DNA complexes remained elusive until several groups recently used modern single molecule assays to study the process. Historically, this was because – despite the existing familiarity with the concepts of facilitated diffusion – under the extant ensemble reaction conditions, the rate of homologous pairing is not limited by the rate of the homology search [29,62]; therefore, it was not possible to elucidate the mechanism of the search process itself, despite clever attempts [63–65].

To detect DNA pairing in real time using direct visualization, Forget and Kowalczykowski incubated fluorescent RecA–ssDNA nucleoprotein filaments with λ phage DNA (48.5 kb). In the initial attempts, the dsDNA was attached at one end to the surface of a flow cell, extended by solution flow to just under contour length, and then the opposite DNA end was attached to the surface [86]. When imaged using TIRF microscopy, the vast majority of the stretched molecules did not form stable complexes, which was perplexing because when the molecules were incubated together under identical conditions and subsequently attached to the surface, pairing efficiency was high. Interestingly, a subset of molecules that were either tethered only at one end, or proximally tethered in such a way that the ends of the DNA molecule were close to each other, demonstrated stable pairing when the reaction was incubated without flow (Figure 3D). These observations suggested that the 3D conformation of the dsDNA target was instrumental in the homology search. To further test this hypothesis, a dual beam optical trap was used to micromanipulate a single molecule of dsDNA between two beads so as to vary the end-to-end distance in a controlled manner [86,87]. When a λ dsDNA molecule was extended to beyond $\sim 16 \mu\text{m}$ (the length at which the DNA was at its theoretical B-form length), homologous pairing was not observed, in agreement with the TIRF experiments; however, when the ends of the molecule were brought together so that the dsDNA could adopt a more randomly coiled polymer, the efficiency of RecA pairing increased monotonically as the end-to-end distance was decreased (Figure 3E). Furthermore, transient interactions at off-target, non-homologous sites were observed to induce DNA looping events that were short-lived (<10 s) and were released as the DNA molecule was extended through micromanipulation. These observations provided direct evidence that a single RecA–ssDNA nucleoprotein filament simultaneously samples different sites within the same dsDNA while the filament is searching for homology, and also that efficient searching occurred when the target dsDNA adopted a 3D, random coil that allowed multiple contacts with the filament. This process was described as a ‘intersegmental contact sampling’, wherein a single RecA filament of a theoretically unlimited length, makes many transient and weak interactions with the randomly coiled dsDNA containing the target site; both the polyvalent binding and the behavior of coiled DNA ensure that segments of the dsDNA target are statistically confined to, and quickly sampled within, the 3D space around the filament (Figure 3C) [86]. This intersegmental contact sampling is analogous to parallel processing, where only the local concentration of the dsDNA target and the net length of the RecA nucleoprotein filament limit the number of interactions that can be simultaneously tested.

But are random 3D collisions sufficient for the homology search? Ragunathan *et al.* used single molecule FRET to probe the dynamic fluctuations of RecA–nucleoprotein filaments on short, oligonucleotide length filaments and dsDNA targets (Figure 3F) [88,89]. They observed rapid fluctuations consistent with 1D sliding of the filament along the dsDNA, with an average interaction time of 0.5–10 s and a sampling site size of 60–300 base pairs [89]. It is very likely that this rapid, 1D sliding is another important aspect of homology sampling at each contact point along the RecA filament; classic analysis by Berg and von Hippel showed that the optimal search strategy employs a combination of 3D searching and 1D sliding [66,70,71]. Ultimately, it will be essential to understand these reactions in the context of the cellular process. Live imaging of *E. coli* showed that RecA filaments span the length of dividing cells and that they can laterally aggregate into bundles [13]. *In vivo*, the homology search took approximately 50 min [13].

Recognition of DNA Sequence Homology: Testing Involves Concerted Sampling of Short Duplet or Triplet Matches

Recognition of DNA sequence homology by the RecA filament was detected in an elegant set of experiments where one end of a dsDNA molecule was tethered to the surface of a flow cell with the opposite end attached to a magnetic bead, while a second transverse molecule of a RecA–ssDNA nucleoprotein filament was held in a dual-beam optical trap (Figure 4A) [90]. The RecA filament could be physically micromanipulated, where it could be pushed against or dragged



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Figure 4. Reduction in Complexity through Microhomology Sampling. (A) 'Dual molecule' method simultaneously using magnetic and optical traps to measure filament DNA contacts formed on stretched or supercoiled DNA [90]. (B) Molecular modeling and simulation of dsDNA interactions with the RecA-ssDNA nucleoprotein filament during initial contact, binding to the secondary DNA-binding site (site II) and homology sampling. Adapted from [91]. (C) Molecular model of a RecA pairing intermediate [91,92]. Labeled amino acids indicate positively charged residues that constitute the secondary DNA-binding site that binds to and stabilizes the incoming homologous dsDNA. (D) Schematic and (E) representative data from a single molecule 'DNA curtain' experiment measuring RecA-mediated DNA 'microhomology sampling' between a long RecA-ssDNA filament and short, fluorescently labeled dsDNA molecules [94]. (F) Molecular modeling (for simplicity, only two strands are shown) of the transient kinetic intermediates formed during the initial microhomology sampling (eight base pairs or less) and subsequent DNA pairing reaction (nine base pairs or greater), resulting in the postsynaptic, Rad51/RecA-Stretched DNA intermediate that is stretched into segments of three base pairs containing normal Watson-Crick spacing, interspersed with 7–8 Å gaps formed by intercalating hydrophobic residues (e.g., Met164 and Ile99). Adapted from [95]. Abbreviations: dsDNA, double-stranded DNA; ssDNA, single-stranded DNA.

across the target dsDNA molecule. Transient homologous recognition events could be measured due to the high sensitivity and time resolution of the experiment; these pairing events were only observed when the dsDNA molecule was supercoiled through the rotation of the paramagnetic trap. Supercoiling was long known to facilitate RecA pairing in D-loop assays (an assay wherein an ssDNA oligonucleotide coated with RecA is paired with supercoiled DNA), by stabilizing the paired heteroduplex product [85].

Although pairing intermediates of the RecA-ssDNA filament bound to the incoming dsDNA have not been crystallized, integration of RecA biochemistry, structural analysis, and molecular dynamics was used to generate a molecular model of the pairing process (Figure 4B). Elements of the model included both the position of positively charged residues within the nucleoprotein filament that are thought to comprise the second strand DNA-binding site, also called site II (Figure 4C), the energetic costs of dsDNA extension upon binding to the filament, and the energy 'payback' when new Watson-Crick base pairs formed in a homologous product [91,92]. A numerical simulation of the process offers a gratifying molecular perspective of the DNA pairing and recognition process (see supplemental Video S1 in the supplemental information online).

To measure the transient kinetic interactions between the RecA filament and dsDNA molecules containing varying degrees of heterologous sequences, a TIRF-based approach was used by the Greene lab. 'Curtains' of either Rad51- or RecA-coated ssDNA were tethered between nanofabricated barriers, and short fluorescently labeled duplex oligonucleotides were incubated

with the filaments (Figure 4D,E). This approach allowed the authors to measure the kinetic lifetimes of the pairing intermediates, and thereby test the model for homology recognition described by Prentiss and colleagues [93,94]. Interestingly, when dsDNA substrates containing seven nucleotides or less of microhomology were used, the authors saw only transiently paired complexes; however, adding an eighth homologous nucleotide resulted in a dramatic increase in both pairing efficiency and binding energy. Adding an additional ninth nucleotide of homology contributed to another dramatic increase in binding energy, constituting a transition from homology sampling to pairing (Figure 4F) [94]. Interestingly, the stability of paired intermediates increases in three nucleotide increments, and this energetic profile is conserved between RecA, Rad51, and Dmc1 [95]. Therefore, RecA also discriminates against heterologous sequences by effectively ignoring homologous sequences shorter than seven nucleotides, using a fundamental eight nucleotide homology test during the molecular search process [94]. Together, both a reduction in dimensionality (through intersegment contact sampling) and a reduction in complexity (through microhomology sampling) provide an elegant and satisfying solution to a classical and fundamental problem in molecular search theory.

Whether these processes are sufficient to explain the *in vivo* homology search in larger eukaryotic cells remains unclear. Although only 12, 13, or 17 nucleotides are required to define a unique sequence in the *E. coli*, *Saccharomyces cerevisiae*, and human genomes, respectively [94], mammalian cells have nuclear volumes approximately 1000 times greater than a yeast nucleus or a bacterial cell [58,59,96]. Complicating the problem of an expanded search volume and larger genome is that eukaryotic DNA is densely packaged into chromatin that is tightly regulated and organized into stable ‘territories’ within a single cell’s nucleus. In yeast, *in vivo* labeling experiments have demonstrated that centromeres and telomeres only explore approximately 3–5% of the nuclear volume, indicating that once the chromosomes are packaged into the nucleus, they are relatively well confined [96]. This in itself would not be a problem, but FISH and chromatin capture experiments have demonstrated that homologous chromosomes can occupy territories that are far apart from one another, suggesting that the homology search process could be greatly impeded by physical distance [77,78]. This problem is mitigated after DNA replication due to sister chromatid cohesion. When exposed to DNA damage such as ionizing radiation or inducible endonucleolytic dsDNA breaks, the motility of eukaryotic chromosomes increases; however, the extent of motility seems to vary greatly between organisms and cell cycle, with some loci inherently more mobile than others [96–102]. These *in vivo* chromosome mobility assays are dependent on ploidy, where and how the loci is labeled, and length of observation; nonetheless, it is clear that cells exposed to DNA damage exhibit an increase in chromosome mobility, in principle, enabling loci to explore larger volumes. The precise factors that contribute to this ‘nuclear jiggling’ and how these dynamics contribute to chromosome pairing remain controversial and are still being defined. Defining and characterizing the factors that are required to accelerate and facilitate the eukaryotic homology search, either through conformational changes in global or local chromatin structure or directed, motor-dependent motion will be important steps in understanding how the recombination machinery finds its homologous target (see Outstanding Questions). We expect that advances in super-resolution microscopy and single molecule manipulation and detection methods will be essential in future studies probing the dynamics of the search process in living cells, and that those findings will impact not only our understanding of recombination and DNA repair but will also directly inform the development of template-directed gene therapies using RNA-guided CRISPRs.

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Outstanding Questions

How do other mediator proteins and regulatory motor proteins alter the dynamic behavior (assembly or disassembly) of the RecA/RAD51 filament?

What is the role of SMC (Structural Maintenance of Chromosomes) proteins in recombination-dependent homology searches?

How is the homology search affected by the length of the ssDNA within the filament: at some characteristic length, will the search be hindered by more off-target sampling?

How does RAD51 search for homology in the context of chromatin and heterochromatin?

How does the physical organization of the genome influence search and recombination bias?

How are repair outcomes influenced by differential euchromatin and heterochromatin states?

To what extent do histone chaperones and chromatin remodeling enzymes affect DNA pairing and strand exchange by RAD51?

What other chromatin modifications and remodelers are required for recombination?

How does chromatid cohesion impose sister bias and is this merely a consequence of physical proximity, or are other biochemical mechanisms at play?

How do chromatin-associated RNAs (e.g., R-loops, noncoding RNA, etc.) affect recombination?

Supplemental Information

Supplemental information associated with this article can be found, in the online version, at <http://dx.doi.org/10.1016/j.tibs.2016.04.002>.

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