Defining the influence of Rad51 and Dmc1 lineage-specific amino acids on genetic recombination

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The vast majority of eukaryotes possess two DNA recombinases: Rad51, which is ubiquitously expressed, and Dmc1, which is meiosis-specific. The evolutionary origins of this two-recombinase system remain poorly understood. Interestingly, Dmc1 can stabilize mismatch-containing base triplets, whereas Rad51 cannot. Here, we demonstrate that this difference can be attributed to three amino acids conserved only within the Dmc1 lineage of the Rad51/RecA family. Chimeric Rad51 mutants harboring Dmc1-specific amino acids gain the ability to stabilize heteroduplex DNA joints with mismatch-containing base triplets, whereas Dmc1 mutants with Rad51-specific amino acids lose this ability. Remarkably, RAD-51 from Caenorhabditis elegans, an organism without Dmc1, has acquired “Dmc1-like” amino acids. Chimeric C. elegans RAD-51 harboring “canonical” Rad51 amino acids gives rise to toxic recombination intermediates, which must be actively dismantled to permit normal meiotic progression. We propose that Dmc1 lineage-specific amino acids involved in the stabilization of heteroduplex DNA joints with mismatch-containing base triplets may contribute to normal meiotic recombination.

[Keywords: DNA repair; Dmc1; homologous recombination; meiosis; Rad51]

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Homologous recombination (HR) enables the exchange of genetic information between DNA molecules and is a major driving force in evolution. HR plays essential roles in double-strand DNA break (DSB) repair (Symington et al. 2014), the rescue of stalled or collapsed replication forks (Cox et al. 2000, Symington et al. 2014), and meiosis (Neale and Keeney 2006, Brown and Bishop 2014). During HR, a presynaptic ssDNA is paired with the complementary strand of a homologous dsDNA, resulting in displacement of the noncomplementary strand (Kowalczykowski 2015; Morrical 2015; Prentiss et al. 2015). Crystal structures of RecA-ssDNA presynaptic and RecA-dsDNA postsynaptic complexes reveal that the DNA is organized into near B-form base triplets separated by ~8 Å between adjacent triplets (Chen et al. 2008; Prentiss et al. 2015). This structural organization likely underpins homology recognition mechanisms and the ability of the Rad51/RecA family of recombinases to promote DNA strand invasion in 3-nucleotide (nt) steps (Ragunathan et al. 2011; Lee et al. 2015, 2017; Prentiss et al. 2015; Qi et al. 2015).

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In mitotic cells, HR is used primarily for the repair of spontaneous DNA breaks, such as those associated with DNA replication errors [Paques and Haber 1999; Cox et al. 2000; Symington et al. 2014]. Meiotic HR is used to repair programmed DSBs generated by the Spo11 complex and is biased toward interhomolog recombination to allow for the formation of crossovers necessary for accurate chromosome segregation in the first meiotic division [Neale and Keeney 2006; Brown and Bishop 2014; Thacker et al. 2014; Zhu and Keeney 2014; Hunter 2015; Marsolier-Kergoat et al. 2018].

Rad51 is the only recombinase in mitotic cells, whereas both Rad51 and Dmc1 are expressed during meiosis in the vast majority of eukaryotes [Neale and Keeney 2006; Brown and Bishop 2014; Hunter 2015]. In the budding yeast Saccharomyces cerevisiae, Dmc1 is responsible for catalyzing interhomolog recombination during meiosis, while Rad51 promotes Dmc1 presynaptic filament assembly and participates in intersister repair that gives rise to noncrossover outcomes [Neale and Keeney 2006; Lao et al. 2013; Brown and Bishop 2014]. Dmc1 mediates the intersister recombination that occurs contemporaneously with interhomolog recombination [Cloud et al. 2012]. Rad51 is inactive during this period and becomes active only after induction of Ndt80 and destruction of the synaptonemal complex [Argunhan et al. 2017; Prugar et al. 2017]. If activated prematurely, Rad51 can mediate interhomolog recombination, but not as efficiently as Dmc1 [Lao et al. 2013; Callender et al. 2016]. It is clear that Rad51 and Dmc1 interact with different subsets of accessory factors [Brown and Bishop 2014]; some examples include the protein complexes Mei5/Sae3 and Hop2/Mnd1, which interact with Dmc1 [Tsubouchi and Roeder 2002; Chen et al. 2004; Hayase et al. 2004; Tsubouchi and Roeder 2004; Petukhova et al. 2005; Ferrari et al. 2009]; Rad54 and Rdh54, which show distinct genetic and biochemical properties in combination with Rad51 and Dmc1 [Klein 1997; Nimonkar et al. 2012]; and Hed1, which is a Rad51-specific binding protein that blocks Rad54 interactions with Rad51 [Tsubouchi and Roeder 2006; Busygina et al. 2008]. In the fission yeast Schizosaccharomyces pombe, the protein Rad22 [an ortholog of S. cerevisiae Rad52] activates Rad51 [also called Rhp51] but inhibits Dmc1, highlighting another example of a recombinase-specific accessory factor [Murayama et al. 2013]. Interestingly, S. pombe Rad51 and Dmc1 promote four-strand exchange reactions, mimicking the formation and branch migration of Holliday junctions, but Rad51 promotes exchange in the 3′-to-5′ direction whereas Dmc1 promotes exchange with the opposite polarity, which could have implications for crossover production in meiosis [Murayama et al. 2008, 2011]. However, there are few other biochemical or biophysical differences between Rad51 and Dmc1 that might help to explain the possible origins or potential evolutionary advantages of using different recombinases during mitosis and meiosis [Neale and Keeney 2006; Sheridan et al. 2008; Brown and Bishop 2014].

Interestingly, biophysical studies have shown that Rad51-ssDNA filaments can bind to dsDNA fragments containing short tracts of sequence microhomology to yield heteroduplex DNA joints, the lifetime of which scales in 3-nt increments consistent with a mechanism involving the stablizing pairing of base triplet interactions [Lee et al. 2015, 2016, 2017; Qi et al. 2015]. However, the introduction of a single nucleotide mismatch within one of the base triplets causes a reduction in binding lifetime of the heteroduplex DNA joint commensurate with the loss of one base triplet pairing interaction [Lee et al. 2015, 2017]. In contrast, Dmc1-ssDNA can tolerate base triplets bearing single, double, or triple mismatches and even abasic sites with no change in the binding lifetimes of the resulting heteroduplex DNA intermediates relative to reactions with fully paired heteroduplex intermediates [Lee et al. 2015, 2017]. These findings suggest that Dmc1 can stabilize mismatched base triplets within heteroduplex DNA joints, whereas Rad51 cannot [Lee et al. 2015, 2017; Borgogno et al. 2016]. Similarly, genetic studies also support the notion that Dmc1 can stabilize mismatch-containing recombination intermediates, whereas Rad51 cannot [Callender et al. 2016]. We and others have hypothesized that the ability of Dmc1 to stabilize imperfectly paired recombination intermediates might reflect an intrinsic difference between the two eukaryotic recombinases [Lee et al. 2015, 2017; Callender et al. 2016]. However, the molecular basis for these differences and their biological implications remained unexplored.

Here, we use structural analysis and bioinformatics to identify Rad51 and Dmc1 lineage-specific amino acid residues that contribute to their unique responses to mismatched base triplets. Based on these analyses, we swapped Rad51 lineage-specific amino acid residues present at the putative DNA-binding interfaces with their lineage-specific counterparts from Dmc1 and vice versa. Single-molecule biophysical analysis of these chimeric recombinases reveals that the differential responses of S. cerevisiae and human Rad51 and Dmc1 to mismatches can be attributed to three lineage-specific amino acid residues within DNA-binding loop L1. Mating type [MAT] switching analysis provides genetic evidence that these L1 residues affect recombination between divergent sequences in vivo. Remarkably, Caenorhabditis elegans RAD-51 L1 amino acid residues more closely resemble Dmc1. Accordingly, wild-type [wt] C. elegans RAD-51 stabilizes mismatch-containing substrates, as is observed for yeast and human Dmc1, whereas mutation of C. elegans L1 residues to their “canonical” Rad51 counterparts abolishes mismatch stabilization. Moreover, worms expressing this chimeric RAD-51 recombinase are proficient for mitotic DNA repair but can accumulate aberrant postsynaptic intermediates during meiosis that must be dismantled by RTEL-1 or HELQ-1 to allow for normal progression through meiosis. Together, our results show that highly conserved lineage-specific amino acid residues in the L1 DNA-binding domain are in part responsible for the differential response of Rad51 and Dmc1 to mismatched sequences and suggest that these amino acid residues may play important roles in mitotic and meiotic recombination.
Results

Identification of Rad51 and Dmc1 lineage-specific amino acid residues

The Rad51 and Dmc1 lineages within the Rad51/RecA family of recombinases arose early in the evolutionary history of eukaryotes (Story et al. 1993; Lin et al. 2006; Chintapalli et al. 2013; Brown and Bishop 2014). These proteins remain closely related, for instance, S. cerevisiae Rad51 (ScRad51) and Dmc1 (ScDmc1) share 56% sequence similarity and 45% sequence identity (Neale and Keeney 2006; Brown and Bishop 2014). For brevity, we use the nomenclature Rad51 and Dmc1 as general designations and ScRad51, ScDmc1, hRAD51 [human RAD51], and hDMC1 [human DMC1] when referring to specific recombinases. Rad51 and Dmc1 form similar filaments on ssDNA, and both promote DNA strand invasion (Neale and Keeney 2006; Brown and Bishop 2014). However, Dmc1 can stabilize imperfectly paired base triplets, whereas Rad51 cannot (Lee et al. 2015, 2017). We speculated that Dmc1-specific amino acid residues might be responsible for this differential response to mismatches. Furthermore, we presumed that residues responsible for this effect might fulfill three criteria: (1) They should be conserved within the Dmc1 lineage of the recombinase family. (2) They should be absent from the Rad51 lineage. (3) They would likely be within one of the two known DNA-binding motifs, DNA-binding loop 1 [L1] or DNA-binding loop 2 [L2], which are present in all recombinase family members (Story et al. 1993; Chen et al. 2008). Given these criteria, we sought to determine whether Rad51 and Dmc1 harbor lineage-specific residues within the L1 and L2 DNA-binding loops.

L1 and L2 were originally identified from examination of the Escherichia coli RecA structure (Story et al. 1993). Therefore, we used E. coli RecA as a model to verify the boundaries of the L1 and L2 motifs [Supplemental Fig. S1; Chen et al. 2008]. E. coli RecA shares 26.5% and 25.6% sequence identity with ScRad51 and ScDmc1, respectively, and the core domain of E. coli RecA cofinals with the core domain of ScRad51 with root mean square deviation of ~1.5 Å (Conway et al. 2004). We then mapped these regions onto a primary structure alignment of Rad51 and Dmc1 from S. cerevisiae, Homo sapiens, Pneumocystis carinii, Oryza sativa, Mus musculus, Entamoeba histolytica, Toxoplasma gondii, and Sus scrofa [Fig. 1A]. From this initial comparison, we identified four amino acid residues within L1 and five amino acid residues within L2 that are conserved within either the Rad51 lineage or the Dmc1 lineage but are divergent between the two recombinases [Fig. 1A]. For ScRad51, these lineage-specific amino acid residues correspond to L1 residues T288, A298, M301, and H302 and L2 residues V328, Q330, V331, D332, and N348. The ScDmc1 lineage-specific amino acids include L1 residues V224, E234, Q237, and K238 and L2 residues Q264, D266, P267, G268, and H285. We validated this initial assignment by analysis of 600 Rad51 protein sequences and 270 Dmc1 sequences [Fig. 1B,C; Supplemental Table S1].

Biochemical characterization of Rad51 and Dmc1 chimeras

If Dmc1 lineage-specific residues present within L1, L2, or both are responsible for mismatch tolerance, then mutation of these residues to those present in Rad51 might abolish this property. Conversely, mutation of the Rad51 lineage-specific residues to their Dmc1 counterparts might enable Rad51 to stabilize mismatched recombination intermediates. To test these hypotheses, we designed chimeric recombinases by swapping the entire L1 and L2 motifs [Supplemental Table S2]. For brevity, we assigned names to the mutants based on the identity of the altered residues (e.g., ScDmc1-RL1 refers to S. cerevisiae Dmc1 harboring the ScRad51 L1, hRAD51-DL12 refers to human RAD51 with the amino acid residues from hDMC1 L1 and L2, etc.) [Supplemental Table S2]. All mutants behaved like their wt counterparts during expression and purification, with the exception of the hDMC1 chimeras, which were significantly less soluble than the wt protein [data not shown]; therefore, we were unable to analyze these hDMC1 mutants.

Each chimeric protein was tested for ATP hydrolysis and DNA strand exchange activity [Supplemental Fig. S2A]. Most of the chimeras retained DNA-dependent ATPase activity, albeit typically at a lower level relative to the wt recombinases, and the single loop swaps exhibited greater ATP hydrolysis activity than the double loop swaps [Supplemental Fig. S2A]. DNA strand exchange assays revealed that the mutant proteins with a single chimeric loop swap exhibited activity comparable with that of their wt counterparts [Supplemental Fig. S2B,C]. However, chimeric recombinases in which both loops were swapped were deficient in strand exchange activity. In particular, ScDmc1-RL12 was significantly compromised for both strand exchange and ATP hydrolysis [Supplemental Fig. S2]. Interestingly, ScDmc1-RL1 exhibited approximately threefold more strand exchange activity compared with wt Dmc1 [Supplemental Fig. S2B,C]. These findings indicate that the identity of the lineage-specific amino acid residues contributes to recombinase interactions with DNA substrates and also show that it is technically possible to swap the Rad51 and Dmc1 L1 or L2 regions without completely abolishing basic protein activities in vitro, although, as specified above, the resulting mutant proteins do show some differences in ATPase and strand exchange activity levels [Supplemental Fig. S2].

Assembly of presynaptic filaments with chimeric recombinases

The chimeric recombinases were tested for the ability to assemble into stable presynaptic complexes using ssDNA curtain assays [Supplemental Fig. S3]. As reported, the addition of wt [unlabeled] Rad51 or Dmc1 results in ATP-dependent displacement of RPA-GFP from the ssDNA, reflecting the assembly of the presynaptic complexes [Supplemental Fig. S3; Ma et al. 2017]. The RPA-GFP reappears when ATP [or both ATP and Ca2+] in the
case of ScDmc1 and hRAD51 is flushed from the sample chamber, reflecting presynaptic complex disassembly (Supplemental Fig. S3; Ma et al. 2017). Most of the chimeras assembled into presynaptic filaments, which remained stable for \( \geq 30 \) min so long as ATP [and Ca\(^{2+}\) when appropriate] was in the reaction buffer (Supplemental Fig. S3; Supplemental Table S3). One exception was ScDmc1-RL12, which failed to assemble into stable filaments [data not shown] and was not characterized further.

Chimeric recombinases exhibit base triplet stepping

We developed a ssDNA curtain assay for visualizing DNA strand exchange intermediates at the single-molecule level (Lee et al. 2015, 2016, 2017; Qi et al. 2015; Qi and Greene 2016; Ma et al. 2017). In brief, a series of Atto565-labeled dsDNA substrates (70 bp) harboring 8- to 15-nt tracts of microhomology targeted toward unique sequences in the M13 ssDNA are incubated with the presynaptic complexes, and unbound dsDNA is flushed away (Fig. 2A,B). The resulting intermediates are visualized by total internal reflection fluorescence microscopy (TIRFM), and dsDNA dissociation rates are obtained from the survival probabilities of the bound dsDNA fragments (Lee et al. 2015, 2017; Qi et al. 2015). Importantly, presynaptic complexes prepared with the chimeric recombinases could bind the Atto565-labeled dsDNA, the resulting dsDNA dissociation rates were comparable with those measured for the wt proteins, and the dissociation rates also varied in 3-nt increments (Fig. 2C–J; Supplemental Fig. S4, S5). We conclude that the chimeric Rad51 and Dmc1 recombinases possess dsDNA-binding and base triplet stepping attributes similar to those determined for their wt counterparts.
Figure 2. Chimeric recombinases undergo base triplet stepping. (A) Schematic of dsDNA capture assay. (B) Illustration of dsDNA substrates used for base triplet stepping assays. The 8-nt tract of microhomology highlighted in green is required for efficient binding, and the nucleotides highlighted in blue represent incremental increases in the microhomology length. Color-coded designations here and in all subsequent figure panels indicate the length of homology at which steps 1–3 are detected (Lee et al. 2015, 2016, 2017). (C,D) Survival probabilities $[\text{C}]$ and dissociation rate data $[\text{D}]$ obtained from the survival probability plots for wt ScRad51. (E,F) Survival probabilities $[\text{E}]$ and dissociation rate data $[\text{F}]$ for ScRad51-DL1. (G,H) Survival probabilities $[\text{G}]$ and dissociation rate data $[\text{H}]$ for ScRad51-DL2. (I,J) Survival probabilities $[\text{I}]$ and dissociation rate data $[\text{J}]$ for ScRad51-DL12. For all dissociation rate graphs, arrows indicate stepwise reductions in dissociation rates coincident with recognition of the third base of each triplet, dashed lines report the mean rate for each step, and the free energy changes $[\Delta\Delta G^\circ]$ associated with each triplet step are indicated. Here and throughout, error bars for survival probability plots represent 70% confidence intervals (CIs) obtained through bootstrap analysis, error bars for the dissociation rate data represent mean ± standard deviation (SD), the number of events used to calculate these values are shown in the survival probability panels, and the color-coded designations indicating steps 1–3 are indicated.

Dmc1 L1 lineage-specific amino acid residues regulate mismatch stabilization

We next asked how the chimeric recombinases responded to DNA mismatches. These assays are conducted by calculating the $\Delta\Delta G^\circ$ values for a given recombinase in reactions with fully paired dsDNA oligonucleotide substrates [as described above] compared with reactions with the same recombinase using dsDNA oligonucleotide substrates that have single nucleotide mismatches introduced at defined locations [Fig. 3A, Lee et al. 2015, 2017]. Rad51, RecA, and Dmc1 require perfect Watson-Crick pairing interactions to stabilize base triplets located at the terminal positions of tracts of microhomology tracts embedded within the dsDNA; in the absence of perfect pairing, they fail to take a step [Fig. 3A, panel i; Lee et al. 2015, 2017]. Rad51 and RecA also require perfect Watson-Crick base-pairing interactions to stabilize base triplets located at internal positions, and the presence of a mismatch at these internal positions is revealed as the loss of a base triplet step [Fig. 3A, panel ii; Lee et al. 2015, 2017]. In contrast, Dmc1 can stabilize mismatches at internal positions [Fig. 3A, panel iii; Lee et al. 2015, 2017]. Indeed, Dmc1 can stabilize internal base triplets containing single, double, and triple mismatches and even abasic sites so long as these imperfect triplets are flanked by homologous sequences [Lee et al. 2017].

Consistent with previous results, none of the recombinases was capable of stabilizing a base triplet located at the terminal position of a 12-nt tract of microhomology [Fig. 3B], and the resulting substrates exhibited dissociation rates similar to those measured for a substrate with only 9 nt of microhomology [Fig. 3D–F]. We next tested the chimeric recombinases with mismatch-containing base triplets that were juxtaposed to a single perfectly paired triplet with a 15-nt tract of microhomology [Fig. 3C]. As shown previously, ScRad51 and hRAD51 could step over internal mismatches but could not stabilize the internal mismatched triplet, instead yielding dissociation rates comparable with a substrate bearing only 12 nt of microhomology [Fig. 3G,H; Lee et al. 2015, 2017]. In contrast, ScDmc1 and hDMC1 yielded dissociation rates comparable with the corresponding substrate bearing 15 nt of perfect microhomology [Fig. 3I]. These findings demonstrate that Dmc1 L1 lineage-specific amino acid residues regulate mismatch stabilization.

Effects of Rad51 and Dmc1 specific amino acids

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played filament assembly and disassembly kinetics that Dmc1 lineages (Fig. 1A,B). The resulting mutants dis- these residues were most conserved across the Rad51 or mismatches. We focused on the ScRad51 chimeric point set of these L1 residues could confer the ability to stabilize ScRad51, we sought to determine whether a smaller sub-

resemble the general biochemical characteristics of wt

effort to identify a mutant protein that might more closely

ing and ATPase activity relative to ScRad51 (Supplemen-

ScRad51-DL1 swap mutant has compromised DNA-bind-

to DNA mismatch stabilization

Three Dmc1 L1 amino acid residues contribute
to DNA mismatch stabilization

ScRad51-DL1 has a total of six amino acid residues from ScDmc1 (Supplemental Tables S1, S2), and the complete ScRad51-DL1 swap mutant has compromised DNA-binding and ATPase activity relative to ScRad51 (Supplemental Fig. S2A, S3; Supplemental Table S1). Therefore, in an effort to identify a mutant protein that might more closely resemble the general biochemical characteristics of wt ScRad51, we sought to determine whether a smaller subset of these L1 residues could confer the ability to stabilize mismatches. We focused on the ScRad51 chimeric point mutants T288V, A298E, M301Q, and H302K because these residues were most conserved across the Rad51 or Dmc1 lineages [Fig. 1A,B]. The resulting mutants displayed filament assembly and disassembly kinetics that closely resembled the wt protein [Supplemental Table S3]. Each mutant exhibited dissociation rates for the Atto655-dsDNA fragments harboring the 9-, 12-, and 15-nt tracts of microhomology, which were essentially indistinguishable from wt ScRad51 [Supplemental Table S4]. As with all recombinases, the point mutants were unable to stabilize mismatches present at the terminal position of an embedded tract of microhomology [Fig. 4A,C]. ScRad51-T288V behaved like wt ScRad51, as it was able to step over the internal mismatches but did not stabilize the mismatch [Fig. 4B,D]. However, ScRad51-M301Q stabilized mismatches regardless of the relative position of the mismatch within the base triplet, exhibiting behavior comparable with wt ScDmc1 [Fig. 4D]. Interestingly, both ScRad51-A298E and ScRad51-H302K mutants were able to stabilize an internal mismatch located at the edge of a base triplet [nucleotide position 12 in Fig. 4B], but neither mutant could stabilize an internal mismatch located at the center of the triplet [nucleotide position 11 in Fig. 4B,D], these properties were confirmed using an alternative set of dsDNA substrates targeted to a different region.

Figure 3. The Dmc1 L1 DNA-binding loop allows for mismatch stabilization. [A] Schematic illustration of a dsDNA substrate (shown in blue; the noncomplementary strand is omitted for clarity) bound to a presynaptic ssDNA (in red). A single mismatch is indicated with a star [★] and is positioned in either the terminal base triplet [panel I], an internal base triplet that is not stabilized [as with Rad51; panel II], or an internal triplet that is stabilized [as with Dmc1; panel III] [Lee et al. 2015]. The binding steps relative to triplet length are highlighted, whereas step 1 corresponds to the initial binding interaction, whereas steps 2 and 3 reflect the changes in dissociation rates that occur for each 3-nt increase in length. [B] Schematic of dsDNA substrates with mismatches positioned within the terminal base triplet. The locations and identities of the mismatches are highlighted as an underlined magenta “X.” (C) Schematic of dsDNA substrates with mismatches positioned within the internal triplet. Assays with terminal mismatch substrates for wt and chimeric versions of ScRad51 (D), hRAD51 (E), and ScDmc1 (F). Assays with internal mismatch substrates for wt and chimeric versions of ScRad51 (G), hRAD51 (H), and ScDmc1 (I). In D–I, the location and identity of the mismatched nucleotide are highlighted in magenta and underlined. Error bars represent mean ± SD. All ΔΔG° values for the mismatched substrates for each protein were calculated to relative reactions performed with the same protein and the fully paired substrates. In A–C, color-coded dashed lines indicate the length of homology at which steps 1–3 are detected for fully homologous substrates (see Fig. 2B,D), and in D–H, the corresponding free energy changes associated with each step (for fully homologous substrates) are indicated with color-coded
of the presynaptic ssDNA (data not shown, Lee et al. 2015). However, a ScRad51-A298E, H302K double mutant (Supplemental Table S3) could stabilize internal mismatched base triplets regardless of whether the mismatch was located at the center or edge of the triplet (Fig. 4D). These results suggest that ScRad51 L1 amino acids A298, M301, and H302, when mutated to the corresponding Dmc1 L1 residues, can all contribute to mismatch stabilization.

Genetic characteristics of ScRad51 chimeras

A major challenge in understanding why eukaryotes have two recombinases is that Rad51 and Dmc1 cannot simply be replaced for one another because of the many mitotic- and meiotic-specific cofactors necessary for each of their in vivo functions, respectively (Neale and Keeney 2006; Brown and Bishop 2014). However, the chimeric proteins described here offer the opportunity to examine potential benefits of the dual recombinase system within the context of mutants that have a well-defined biochemical characteristic; namely, the ability or inability to stabilize mismatched recombination intermediates.

For genetic testing of the chimeric ScRad51 proteins, we constructed S. cerevisiae strains in which the chromosomal rad51 gene was replaced with rad51 mutants containing Dmc1 lineage-specific amino acids. Western blot analysis confirmed that all proteins were expressed (Supplemental Fig. S6). In addition, wt ScRad51, ScRad51-DL1, ScRad51-A298E, and ScRad51-M301Q all supported some level of YFP-Rad54 DNA repair focus formation upon exposure to ionizing radiation (Supplemental Fig. S7A,B). Since the assembly of DNA repair-specific Rad54 foci is known to be dependent on Rad51-ssDNA filaments

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**Figure 4.** L1 amino acid identity influences recombination between divergent DNA sequences. Schematic illustrations of dsDNA sequences used to analyze the effects of terminal (A) and internal (B) mismatched triplets. (C) Terminal mismatch assays for the ScRad51 point mutants. (D) Internal mismatch assays for the ScRad51 point mutants. In C and D, data for wt ScRad51 are reproduced from Figure 3, F and I, for comparison. (E) Schematic illustration of the MAT switching assay highlighting the products that arise from nonhomologous end joining (NHEJ) and HR. (F) Example of a genomic blot analysis of StyI-digested DNA after galactose induction of HO endonuclease. (G) Bar graph showing the fraction of repair products attributed to HR for strains expressing either wt ScRad51 or ScRad51-M301Q for templates with varying levels of sequence divergence. Each bar represents the mean ± SD from n independent experiments, as indicated. The statistical significance for the observed differences between wt and mutant for the 20% and 25% sequence divergence data sets are indicated. P-values from an unpaired two-tailed Student’s t-test are indicated. (H) Bar graph illustrating the percent difference for HR-mediated MAT switching between ScRad51-M301Q and wt Rad51. Each bar represents the mean ± SEM calculated from data in F. In A and B, color-coded designations indicate the length of homology at which steps 1–3 are detected for fully homologous substrates, and in C and D, the corresponding free energy changes associated with each step (for fully homologous substrates) are indicated with color-coded dashed lines.
(Lisby et al. 2004), these results provide evidence that the mutant Rad51 proteins could form presynaptic filaments in vivo, although the A298E mutant was compromised in focus formation compared with wt ScRad51. In contrast, ScRad51-H302K and the ScRad51-A298E, H302K double mutant were unable to support YFP-Rad54 focus formation, suggesting that these mutants were defective in presynaptic filament assembly in cells [Supplemental Fig. S7B]. We next asked whether ScRad51 chimeras could support cell growth on media containing the DNA-damaging agents methyl methanesulfonate (MMS) or zeocin [Supplemental Fig. S7C]. Importantly, strains expressing either ScRad51-A298E or ScRad51-M301Q exhibited near wt levels of cell growth in the presence of MMS or zeocin [Supplemental Fig. S7B], demonstrating that these point mutants retain function in vivo. As expected, ScRad51-H302K and ScRad51-A298E, H302K were compromised for growth on plates with MMS or zeocin [Supplemental Fig. S7C]. Interestingly, ScRad51-DL1 was also compromised for growth on MMS or zeocin plates even though this chimera was functional in vitro and supported Rad54 focus formation [Supplemental Fig. S2, S3, S7B,C]. We speculate that swapping the entire Rad51 L1 motif may hinder some downstream step in the HR pathway.

**Lineage-specific amino acid residues contribute to recombination in yeast**

We used a modified version of the mating type switching assay to determine whether lineage-specific L1 residues might contribute to recombination fidelity. In this assay, the MAT locus is cleaved by the HO endonuclease, and Rad51-mediated recombination takes place between the cleaved MAT locus and either the HMRa or HMRa donor locus [Haber 2012, Mehta et al. 2017]. Strand invasion initiates from the Z-box within the MAT locus, which is homologous to sequences in the donor loci [Fig. 4E, Haber 2012, Mehta et al. 2017]. The cleaved intermediates can be repaired by HR or nonhomologous end joining (NHEJ), and the identity of the resulting products can be defined by genomic blot analysis [Fig. 4F]. To examine the effects of mismatches on recombination, we introduced point mutations at every eighth, seventh, sixth, fifth, or fourth position within the Z-box, corresponding to 12.5%, 14.2%, 16.7%, 20%, or 25% sequence divergence between donor and acceptor loci, respectively [Fig. 4E, Supplemental Table S5].

For the strain expressing wt ScRad51, MAT switching was remarkably tolerant of mismatches [Fig. 4G]. For instance, with the wt templates [0% divergence] 75.1% ± 1.33% of the repair products could be attributed to HR-mediated repair, while 66.1% ± 4.75% of the repair products could be attributed to HR for templates with 16.7% sequence divergence [Fig. 4G]. However, HR efficiency drops markedly for templates with mismatches at every fifth [20% divergence] and fourth [25% divergence] position, yielding values of 35.0% ± 4.75% and 5.52% ± 1.46%, respectively [Fig. 4G]. These observations are in good agreement with previous reports indicating that wt ScRad51 supports efficient break-induced replication (BIR) for templates with similar levels of sequence divergence [Anand et al. 2017].

Consistent with the MMS and zeocin resistance assays, ScRad51-DL1, ScRad51-H302K, and the ScRad51-A298E, H302K double mutant were all compromised for MAT switching even at 0% divergence, and these defects were exacerbated at higher levels of sequence divergence [Supplemental Fig. S7D,F,G]. In contrast, the ScRad51-M301Q point mutant supported levels of HR-mediated repair comparable with wt ScRad51 for the substrates with up to 16.7% divergence. Remarkably, at 20% sequence divergence, representing the midpoint of the curve for wt ScRad51 [Fig. 4G], there was a 26.7% ± 5.00% [Student’s t-test, \( P < 0.001 \)] increase in HR-mediated repair by ScRad51-M301Q relative to wt ScRad51, and there was a 22.2% ± 10.2% [Student’s t-test, \( P < 0.05 \)] increase in HR for the template with 25% sequence divergence [Fig. 4G, H]. These findings, together with our biophysical data, demonstrate that ScRad51-M301Q, which can stabilize mismatched HR intermediates in vitro, also supports a higher recombination frequency for mismatched substrates in vivo. Interestingly, although ScRad51-A298E was functional for MAT switching, this mutant showed no differences in HR efficiency compared with wt ScRad51 for templates with increasing sequence divergence [Supplemental Fig. S7E]. Given this outcome, it is notable that ScRad51-A298E could only stabilize mismatches located at the edge of a base triplet [Fig. 4B,D], and the divergent Z-box sequences will always have mismatches at both the center and edge positions of the mismatched base triplets [irrespective of the frame of reference] [Supplemental Table S5]. The inability of ScRad51-A298E to stabilize mismatches located in the center of a base triplet could explain why this particular mutant does not behave like ScRad51-M301Q in the MAT switching assays.

**C. elegans RAD-51 behaves like Dmc1 in vitro**

Some eukaryotes, such as *Caenorhabditis sp.*, have lost the *DMC1* gene, although the reasons for this loss remain uncertain [Brown and Bishop 2014]. Surprisingly, inspection of RAD-51 from *Caenorhabditis sp.* revealed that the lineage-specific residues present in L1 were not the same as “canonical” Rad51 [we use the term “canonical” to identify Rad51 from species that have both recombination pathways] but instead more closely resembled Dmc1 [Fig. 5A]. If our hypothesis regarding the role of L1 in mismatch stabilization is correct, then *C. elegans* RAD-51 (CeRAD-51) may stabilize mismatches, whereas a RAD-51 mutant in which the “Dmc1-like” amino acids were converted to the Rad51 lineage-specific residues might lose the ability to stabilize mismatches.

To test this hypothesis, we made a CeRAD-51 N246S, E256A, K260H triple mutant protein (corresponding to ScRad51 amino acids T288, A298, and H302, respectively) [Fig. 5A], which we refer to as CeRAD-51-TM for brevity. Both CeRAD-51 [Taylor et al. 2016] and CeRAD-51-TM retain similar biochemical activity, although the ssDNA-
and dsDNA-binding activity of CeRAD-51-TM is modestly reduced (Supplemental Fig. S8A,B). CeRAD-51 and CeRAD-51-TM also form presynaptic complexes in the ssDNA curtains with similar assembly and disassembly kinetics, although CeRAD-51-TM binds $\sim 21\%$ more slowly and dissociates $\sim 26\%$ more quickly than CeRAD-51 (Supplemental Fig. S8C,D; Supplemental Table S3). Assays with Atto565-dsDNA fragments confirmed that both CeRAD-51 proteins exhibited base triplet stepping (Fig. 5B,C; Supplemental Fig. S8E,F), and, as with the other recombinases, wt and triple mutant RAD-51 were unable to stabilize mismatches present at the end of an embedded tract of microhomology (Fig. 5D). However, in contrast to ScRad51 and hRAD51, wt CeRAD-51 could stabilize mismatches located at an internal position within the embedded tract of microhomology (Fig. 5E). This finding demonstrates that wt CeRAD-51 does not behave like “canonical” Rad51 when presented with a mismatched substrate; rather, it responds similarly to Dmc1. However, the ability to stabilize mismatched base triplets was abolished for the CeRAD-51-TM protein (Fig. 5E). Remarkably, CeRAD-51 could also promote D-loop formation with mismatched substrates (32\% sequence divergence), albeit at low efficiency, whereas CeRAD-51-TM lacks this activity (Supplemental Fig. S8G,H). Together, these findings provide additional support for the premise that lineage-specific L1 residues in Dmc1 confer the ability to stabilize mismatched base triplets within the context of heteroduplex strand exchange intermediates.

Chimeric CeRAD-51 gives rise to toxic recombination intermediates

To investigate the importance of the L1 lineage-specific amino acids in vivo, we performed genome editing by CRISPR–Cas9 to introduce three substitution mutations [N246S, E256A, and K260H] into the C. elegans rad-51 gene. The resulting rad-51(knu529) strain (Supplemental Fig. S9A,B) was assessed for evidence of meiotic dysfunction, genome instability, and loss of fecundity. Deletion of rad-51 results in defective meiotic DSB repair and embryonic lethality in worms [Alpi et al. 2003; Martin et al. 2005]. In contrast, the rad-51(knu529) strain exhibited brood sizes, embryonic viability, and meiotic chromosome nondisjunction rates (as assessed by the frequency of males) comparable with N2(Wt) control strains (Fig. 6A,B). The lack of an apparent phenotype associated with the rad-51(knu529) allele indicates that CeRAD-51-TM is a functional recombinase in vivo.

Figure 5. C. elegans RAD-51 has Dmc1-like amino acids and dsDNA-binding properties. (A) Comparison of Caenorhabditis sp. RAD-51 sequences (C. elegans, Caenorhabditis remanei, and Caenorhabditis brenneri) with Rad51 and Dmc1 sequences from organisms with both recombinases (S. cerevisiae, H. sapiens, and O. sativa). Color-coding is the same as shown in Figure 1A. Base triplet stepping data for wt CeRAD-51 [B] and the CeRAD-51 triple mutant [C]. (D) Terminal mismatch triplet assays for CeRAD-51 and CeRAD-51-TM (CeRAD-51 N246S, E256A, K260H triple mutant protein). Data for wt ScRad51 and ScDmc1 are shown for comparison and are reproduced from Figure 3, D and F. [E] Internal mismatch triplet assays for CeRAD-51 and CeRAD-51-TM. Data for wt ScRad51 and ScDmc1 are shown for comparison and are reproduced from Figure 3, D, F, G, and I. In D and E, the free energy changes associated with each step (for fully homologous substrates) are indicated with color-coded dashed lines.
and rol-1 markers, while the noninverted chromosome carries the dpy-25 allele. Rare heterologous recombination [het-rec] events occurring between this synapsed 8-Mb inverted sequence in meiosis can be observed by scoring recombination between the visible markers [Fig. 6A]. In wt and rad-51(knu529) strains, het-rec events are extremely rare. However, it has been shown that depletion of RTEL-1 or BRC-1 in the N2(wt) strain leads to a significant increase in the frequency of het-rec events, which are mismatch-dependent [León-Ortiz et al. 2018]. Given the latter, this system can be used to assess the mismatch tolerance of C. elegans RAD-51. When compared with N2 [wt], depletion of RTEL-1 or BRC-1 yielded a significantly reduced frequency of het-rec events in the rad-51(knu529) strain. Given that RAD-51-TM is intolerant to mismatch events seen in the rad-51(knu529) strain likely reflects an inability to promote pairing reactions between mismatch-containing sequences within the mln-1 inversion. Importantly, scoring the visible markers dpy-17/unc-36, carried on a fully homologous genetic interval on chromosome III, did not reveal any significant difference in recombination frequencies between N2(Wt) and rad-51(knu529) strains [Supplemental Fig. S9C]. These observations suggest that the reduced het-rec frequencies in rad-51(knu529) background arise due to lower mismatch tolerance of CeRAD-51-TM rather than an overall impairment of its recombinase activity. Furthermore, since deletion of the mismatch repair gene msh-2 suppresses het-rec events occurring in the retel-1 background [León-Ortiz et al. 2018], our data further support the importance of mismatch tolerance and recognition in this process.

Given the suppression of the retel-1 phenotype in the het-rec assay, we next assessed whether the rad-51(knu529) allele could also suppress other retel-1-associated phenotypes, including decreased brood size, reduced fecundity, and DNA damage sensitivity [Barber et al. 2008]. Contrary to expectation, we instead observed a synthetic phenotype when retel-1 and rad-51(knu529) were combined, including a further decrease in brood size [Fig. 6B], elevated levels of embryonic lethality [Fig. 6C], and increased sensitivity to genotoxins [Supplemental Fig. S9D]. We also observed a modest increase in chromosomal aberrations in diakinesis [Supplemental Fig. S9E,F].
RECLQ5 was implicated previously in suppressing HR by dismantling RAD51 filaments (Hu et al. 2007), while RTEL1 has been shown to disrupt D-loop HR intermediates to alter the outcome of meiotic DSB repair (Barber et al. 2008; Youds et al. 2010). *C. elegans* RECLQ5 does not seem to play a role in suppressing additional meiotic crossovers or het-rec events (Barber et al. 2008; Youds et al. 2010). Importantly, no discernable genetic interaction was observed when *rad-51(knu529)* was combined with a deletion in *rcq-5* [Fig. 6B,C].

Next, we analyzed the *rtel-1; rad-51(knu529)* strain for alterations in RAD-51 focus formation and/or resolution in the *C. elegans* germline. Premiotic S phase at the distal end of the germ line is followed by progressive stages of meiosis I, which are defined by specific chromosome alterations associated with homologous chromosome alignment, pairing, synapsis, and chiasma formation [Fig. 6D]. Recombination intermediates formed by RAD-51 at SPO-11-induced meiotic DSBs are repaired to produce crossovers or noncrossovers and can be monitored by the appearance and timely resolution of RAD-51 foci in late pachyten/early diplotene. Quantification of RAD-51 foci in each of these zones revealed that both the timing and levels of RAD-51 focal accumulation at meiotic DSBs were comparable between N2(Wt) and the *rtel-1; rad-51(knu529)* strain, further supporting the conclusion that RAD-51-TM is a functional recombinase in vivo. However, in contrast to the N2(Wt) and *rad-51(knu529)* strain alone, RAD-51 foci persisted into late stages of meiotic prophase in the *rtel-1; rad-51(knu529)* strain [Fig. 6D,E]. This phenotype is reminiscent of the phenotype observed in *helq-1; rtel-1-1* double mutants that also present with reduced brood sizes, embryonic lethality, and persistent RAD-51 foci [Ward et al. 2010]. Prompted by this similarity, we crossed *rad-51(knu529)* into the *helq-1* background. Intriguingly, the *helq-1; rad-51(knu529)* strain also displayed persistence of RAD-51 foci in late pachyten and diplotene phases of meiosis [Fig. 6D,E]. The *helq-1; rad-51(knu529)* strain also showed a significant brood size reduction [Fig. 6B], increased numbers of unhatched eggs [Fig. 6C], and a modest effect on chromosomal aberrations in diakinesis [Supplemental Fig. S9E,F].

Taken together, our data reveal that CeRAD-51-TM is a functional recombinase proficient for meiotic HR in vivo but is unable to tolerate mismatch-containing heterologous DNA sequences duringstrand invasion. Since RTEL-1 and HELQ-1 are believed to act to dismantle post-synaptic recombination intermediates, the synthetic genetic interactions with *rtel-1* and *helq-1* also reveal a dependence on postsynaptic recombination regulators for the survival and effective repair of meiotic DSBs in the *rad-51(knu529)* strain. This is suggestive of the presence of toxic recombination intermediates arising in the *rad-51(knu529)* strain, which must be effectively removed by RTEL-1 or HELQ-1 to allow meiotic progression.

**Discussion**

Here we explored the structural elements that contribute to the differential response of Rad51 and Dmc1 to mismatch-containing base triplets. We propose that these structural and functional differences represent a fundamental distinction between the Rad51 and Dmc1 lineages of the Rad51/RecA family of DNA recombinases. Here, we discuss possible implications of these findings with respect to recombinase structures, recombination mechanisms, and the evolution of Rad51/RecA family members.

**Lineage-specific amino acid residues help determine recombinase behaviors**

L1 and L2 DNA-binding loops of the Rad51/RecA family of DNA recombinases contain amino acids that are specifically conserved within either the Rad51 lineage or the Dmc1 lineage but not both. Our data demonstrate that three amino acid residues within the L1 DNA-binding loop regulate the response of Rad51 and Dmc1 to mismatched base triplets. Mutations that swap the identities of these residues alter the response of the respective recombinase to match that of its paralog, and this behavior appears to be conserved between recombinases from both *S. cerevisiae* and humans. We speculate that within the Rad51 lineage, these residues may help enhance recombination fidelity, whereas the identity of these residues within the Dmc1 lineage may contribute to meiosis, perhaps by allowing for more favorable recombination between polymorphic parental alleles. Indeed, a single point mutation in Rad51 (M301Q) is sufficient to recapitulate Dmc1-like mismatch stabilization in vitro and also enhances recombination between divergent sequences in vivo. However, we also note that the precise spatial geometry of the L1 DNA-binding loop is likely to be influenced by its native context, in particular its connectivity to the rest of the recombinase core domain. Thus, it is possible that the mismatch tolerance observed for Rad51-M301Q may not exactly match that of Dmc1. Indeed, Dmc1 itself may prove to be even more mismatch-tolerant in vivo than the Rad51 chimeras harboring Dmc1 residues. Although the natural prevalence of sequence polymorphisms between parental alleles would not approach the high levels of sequence divergence in our MAT switching assays, we anticipate that more subtle differences in recombination efficiency involving templates with lower divergence may be important when considered over longer evolutionary time scales. Moreover, MAT switching is a highly efficient HR-dependent process that is greatly facilitated by local chromosomal architecture (Haber 2012), which may make MAT switching inherently more tolerant to mismatches. It is possible that less efficient HR-mediated repair events could exhibit even more pronounced effects at lower levels of sequence divergence.

Interestingly, ScRad51-DL2 retains basic biochemical functions but does not support Rad54 focus formation or allow for growth on MMS or zeocin plates (data not shown). The disparity between the in vivo and in vitro activities of this chimera suggest that lineage-specific amino acids in L2 may be important for assembly of the native presynaptic complex. Finally, in addition to the L1 and L2 amino acids described in this study, we also identified...
~19 lineage-specific amino acid residues present in other regions of the two recombinases (data not shown). We anticipate that many of these residues may mediate Rad51- and Dmc1-specific protein–protein contacts. Exploring the roles of these lineage-specific amino acids may yield further insights into the differential properties of the two eukaryotic recombinases.

Recombination between divergent sequences

Rad51/RecA family recombinases require at least consecutive 8 nt of microhomology for efficient recognition of short dsDNA substrates in vitro [Hsieh et al. 1992; Raganathan et al. 2011; Danilowicz et al. 2015; Qi et al. 2015]. However, the efficiency of BIR [Anand et al. 2017] and MAT switching [this study] indicate that the in vivo requirements for donor DNA recognition and strand invasion are significantly less stringent. Two factors may account for these differences. First, in vivo recombination requires many other proteins, a number of which may influence the activities of Rad51 [Symington et al. 2014; Kowalczykowski 2015]. Second, long substrates likely allow for multiple simultaneous interactions, the cumulative effects of which may circumvent the kinetic benefits observed in vitro for shorter substrates bearing a single contiguous tract of homology. For example, the Z-box with mismatches at every eighth position has 40 adjacent 5-nt tracts of microhomology separated from one another by single mismatches, and the Z-box with mismatches at every sixth position has 52 adjacent 5-nt tracts of microhomology [Supplemental Table S5]. At present, the lengths and complex sequence compositions of these substrates preclude detailed biophysical analysis in ssDNA curtain assays, and we cannot yet recapitulate the protein composition of a native presynaptic complex in vitro. Future work will be necessary to understand how these parameters influence the efficiency and kinetics of HR reaction mechanisms.

Our data reveal that the chimeric ScRad51-M301Q mutant, which stabilizes mismatches in biophysical assays, can also support more efficient HR-mediated repair during MAT switching for a Z-box bearing ≥20% sequence divergence. It is interesting to note that for wt ScRad51, we observed the largest decline in HR efficiency only for templates with ≥20% divergence [Fig. 4G,H]. Inspection of these DNA sequences reveals an interesting feature that coincides with this dramatic change in HR efficiency; namely, for templates ranging up to 16.7% sequence divergence, the mismatched base triplets are never adjacent to one another, but this is not true for the templates with ≥20% sequence divergence, which instead are comprised of repeating patterns in which there are always two or three mismatched triplets adjacent to one another [Supplemental Table S5]. One possible explanation for the marked decline in HR efficiency at ≥20% sequence divergence is that Rad51 may have particular difficulties in promoting recombination when two or more adjacent base triplets contain mismatches, whereas Dmc1 [or ScRad51 harboring Dmc1 L1 amino acids] may be more tolerant of these substrates.

What are the mechanistic impacts of mismatches on recombination?

We can envision at least two general mechanisms by which DNA mismatches could affect recombination: [1] by altering the intrinsic stability of the heteroduplex DNA intermediates or [2] by altering the structure of the resulting nucleoprotein complex, perhaps making these intermediates more susceptible to disruption by regulatory enzymes. We do not favor the hypothesis that a small number of mismatches greatly alters the intrinsic stability of the heteroduplex intermediates. Although a single mismatch impacts the binding lifetimes of dsDNA fragments in our biophysical assays, on the whole, these are all still relatively long-lived intermediates [e.g., for ScRad51, a 70-bp dsDNA fragment with 12 nt of microhomology, has a lifetime of ~33 min; increasing the microhomology length to 15 nt yields a lifetime of ~47 min, and introduction of a single mismatch within the 15-nt tract of microhomology reduces the lifetime to ~35 min]. This conclusion is also consistent with studies demonstrating that bacterial RecA is surprisingly tolerant of mismatches in vitro [Danilowicz et al. 2015]. We cannot yet predict how our in vitro observations will scale for the longer in vivo substrates or currently measure the equivalent biophysical parameters in vitro with longer substrates. Nevertheless, the available biophysical data imply that mismatched HR intermediates are not intrinsically unstable. Thus, we favor the hypothesis that mismatches may somehow alter the structure of the nucleoprotein complexes, thus rendering them more susceptible to disruption by regulatory enzymes. Several proteins are known to dissociate HR intermediates, including the S. cerevisiae helicases Srs2 and Sgs1 [BLM in humans], and the postreplicative mismatch repair machinery also plays a role in minimizing HR between divergent sequences [Spell and Jinks-Robertson 2004; Sugawara et al. 2004; Spies and Fishel 2015; Branzei and Szakal 2017; Lorenz 2017]. One possibility is that these or other enzymes may recognize some distinct mismatch-dependent structural feature that enables them to more readily act on Rad51-bound intermediates, whereas Dmc1 may shield mismatched intermediates from these enzymes [Fig. 7A]. This hypothesis is consistent with the observed synthetic lethality between the CeRAD-51-TM mutant and RTEL-1 and HELQ-1 [see below].

Potential mechanisms of DNA mismatch stabilization

Three lineage-specific residues clustered together within L1 DNA-binding loop contribute mismatch stabilization by Dmc1. We proposed that these amino acids may allow Dmc1 to make compensatory DNA contacts that are independent of Watson-Crick pairing interactions. Consistent with this hypothesis, molecular dynamics (MD) simulations of wt hRAD51 and an hRAD51 chimeric mutant bearing three hDMC1 residues in the L1 domain—corresponding to the mutations A241E, M244K and H245K [numbering based on hRAD51] [Fig. 1A]—reveal that the introduction of these Dmc1-specific residues...
results in more intimate contacts between L1 and the phosphate backbone of the complementary strand of the postsynaptic complex (Fig. 7B,C). The notion that Dmc1 residues contact the phosphate backbone of the nascent DNA joint is also consistent with the observation that Dmc1 can stabilize base triplets bearing abasic sites (Lee et al. 2017). At present, these simulations must be interpreted with caution, given that they are based on structural data that lack sufficient information to fully describe the relevant protein–nucleic acid interfaces. In the case of hDMC1, the crystal structures lack DNA (Kinebuchi et al. 2004). For hRAD51, existing cryo-electron microscopy structures lack sufficient resolution to accurately define the L1 protein–DNA interface (Short et al. 2016; Xu et al. 2017). Similarly, although there is a crystal structure of ScRad51 bound to DNA, the L1 and L2 contacts are not visible (Conway et al. 2004). Finally, there is as yet no structural information regarding how recombinases interact with a DNA joint that harbors mismatches, and our simulations do not account for the potential influence of mismatches on recombinase-DNA interactions. Regardless, the MD simulations, together the observation that Dmc1 can stabilize abasic sites [Lee et al. 2017], are consistent with the general notion that the lineage-specific amino acid residues responsible for the differential responses of Rad51 and Dmc1 to mismatches are positioned to interact with the phosphate backbone of the complementary DNA strand within the nascent DNA joint.

C. elegans RAD-51 requires Dmc1-like amino acids

Ecdysozoans, such as Caenorhabditis sp. and Drosophila sp., possess RAD51 but have lost DMC1 as well as genes encoding core meiotic proteins necessary for Dmc1 activity (e.g., Hop2/Mnd1) [Villeneuve and Hillers 2001; Ramesh et al. 2005; Lin et al. 2006; Chintapalli et al. 2013; Hunter 2015]. It remains unknown why these organisms have lost DMC1, and the evolutionary implications of this loss remain unexplored. However, Caenorhabditis sp. and Drosophila sp. have among the highest rates of evolution for the RAD51 genes [Lin et al. 2006]. Interestingly, we show that the lineage-specific amino acid residues within the L1 DNA-binding loop of RAD-51 from Caenorhabditis sp. have adapted to more closely match Dmc1 and that CeRAD-51 can stabilize mismatched DNA.

Figure 7. Potential mechanism of mismatch stabilization and L1 conservation among different Rad51/RecA family members. (A) Model for differences between Rad51 and Dmc1 interactions involving imperfectly paired HR intermediates. (B, C) Snapshots taken from MD simulations of hRAD51 (B) and hRAD51 harboring three Dmc1 lineage-specific amino acids substitutions (A240E M243Q H244K) (C), suggesting that the Dmc1 amino acids are better positioned to contact the incoming complementary DNA strand. Insets highlight potential protein contacts (red dashed lines) with the ribose ring of the phosphate backbone. (D) Comparison of L1 and L2 sequences from E. coli RecA, “canonical” Rad51 and Dmc1 from organisms harboring both recombinases, Rad51 from four Drosophilia sp. [Drosophilia melanogaster, Drosophilia virilis, Drosophilia mojavenis, and Drosophilia simulans], Rad51 sequences designated as “other” from Ustilago maydis, Sodaria macrospora, and Neurospora crassa, and RadA from Halofexar volcanii, Pyrococcus furiosus, Methanooccus maripaludis, Methanooccus voltae, and Sulfolobus solfataricus. Asterisks denote the amino acids that contribute to mismatch stabilization for Dmc1.
base triplets in vitro. Thus, the loss of DMC1 may have
curred, or allowed for, adaptation of CeRAD-51 to become
more “Dmc1-like” with respect to the L1 DNA-binding
loop. In accordance with the importance of these
“Dmc1-like” residues, we found that the chimeric
CeRad51-TM mutant, which bears canonical Rad51-like
amino acids in the L1 loop, is a functional recombinase
in vivo but exhibits an inability to promote recombina-
ion through highly divergent sequences in vivo and a syn-
thetic phenotype with antirecombinases [RTEL1 and
HELQ-1]. These data indicate that the identity of the na-
tive “Dmc1-like” amino acids present in the L1 loops of
wt CeRAD-51 cannot be converted to amino acids found
in the canonical Rad51 lineage without affecting the fidelity
of meiotic HR and increasing the frequency of toxic re-
combination intermediates that places a dependence on
motor proteins such as RTEL-1 and HELQ-1. We specu-
late that mismatches that would normally be masked by
CeRAD-51 are not properly masked by the CeRAD-51-
TM, yielding a dead-end D-loop intermediate that must
disrupted by RTEL-1 or HELQ-1. Thus, the most im-
portant effect of the canonical Dmc1 amino acid residues
may be with respect to differences in their ability to mask
mismatched heteroduplexes from aberrant processing
(Fig. 7A). Interestingly, Drosophila sp. Rad51 L1 amino ac-
cids closely match those found in canonical Rad51 [Fig.
7D]. However, N301 (numbering based on S. cerevisiae
Rad51 for comparison) in Drosophila Rad51 does not
match the methionine found in canonical Rad51 but is in-
stead more similar to the glutamine from Dmc1 [Fig. 7D].
Thus, Drosophila Rad51 may also have a similar require-
ment for “Dmc1-like” amino acids in the L1 DNA-bind-
ing loop.

Among organisms that have Dmc1, chromosome
pairing is initiated through recombination-dependent
mechanisms [Neale and Keeney 2006; Brown and Bishop
2014; Hunter 2015]. In contrast, both C. elegans and
D. melanogaster have evolved alternative, recombin-
ion-independent mechanisms for initiating chromosome
pairing during meiosis [Villeneuve and Hillers 2001;
Gerton and Hawley 2005]. However, use of alternative
pairing mechanisms is not necessarily the defining attribute
of organisms lacking Dmc1. For instance, Ustilago
maydis, Sodaria macrospora, and Neurospora crassa
have all lost the DMC1 gene but require recombination
to initiate chromosome pairing during meiosis [Storlazzi
et al. 2008]. Interestingly, U. maydis, S. macrospora,
and N. crassa all possess canonical Rad51 lineage-specific
amino acids in the L1 DNA-binding loop [Fig. 7D]. Finally,
it should be noted that the L1 [and L2] amino acids from
bacterial RecA are highly divergent from the eukaryotic
recombinases [Figs. 1A, 7D]. Therefore, the mechanisms
by which RecA interacts with the DNA intermediates
differ from those that define the action of the eukary-
ocic recombinases.

Potential origins of the dual-recombinase paradigm
Rad51 and Dmc1 arose early in eukaryotic evolution from
a gene duplication event involving an ancestral archaeal
RadA recombinase, and this gene duplication event may
have coincided with or allowed for the emergence of mei-
osis and sexual reproduction [Ramesh et al. 2005; Lin et al.
2006; Chintapalli et al. 2013]. Some extant archaea, such
as Haloferax volcanii, undergo a conjugation process in-
volving the exchange of highly divergent genetic informa-
tion via HR, and this conjugation process bears some
resemblance to the eukaryotic meiotic program [Mevar-
ech and Werezberger 1985; Rosenshine et al. 1989; Papke
et al. 2004; Cohan and Aracena 2012; Naor et al. 2012;
Naor and Gophna 2013]. Most intriguingly, the RadA L1
DNA-binding loop amino acid residues located at the po-
positions involved in mismatch stabilization are all identical
to those found in the Dmc1 lineage of the Rad51/RecA
family [Fig. 7D]. One speculative possibility is that a “low-
er-fidelity” recombinase (i.e., more “Dmc1-like”) may
have predated the “higher-fidelity” Rad51 present in mod-
ern eukaryotic lineages, and the emergence of Rad51 may
have allowed eukaryotes to take better advantage of HR as
a high-fidelity DNA repair pathway.

Materials and methods

Sequence and crystal structure alignments

Crystal structures of E. coli RecA [1CMX] [Chen et al. 2008],
ScRad51 [1SZP] [Conway et al. 2004], hDMC1 [1V5W] [Kinebuchi
et al. 2004], and hRAD51 [5H1C] [Xu et al. 2017] were obtained
from the RCSB Protein Data Bank and aligned in MacPyMOL
(PyMOL molecular graphics system, version 1.8). All protein se-
quences were obtained from the NCBI database, and sequence
alignments were performed using National Institutes of Health
COBALT [Papadopoulos and Agarwala 2007]. Most of the se-
quenced RAD51 and DMC1 genes have not been experimentally
validated; therefore, to help ensure the alignments reflected in in-
formation from bona fide Rad51 and Dmc1 proteins, we restrict our
analysis to proteins that were ≥200 amino acids in length and
also contained the highly conserved L1 motif GRGEL (or GRGDL,
corresponding to amino acids 294–298 in ScRad51). Sequences
that did not fulfill these two criteria were excluded from the analy-
sis. Aligned sequences were further analyzed for common fea-
tures and annotated using ESPript 3.0 [Robert and Gouet 2014].

Single-molecule dsDNA-binding assays

All single-molecule dsDNA-binding experiments were per-
formed as described previously [Lee et al. 2015, 2017; Qi et al.
2015]. In brief, presynaptic filaments were assembled, as de-
scribed above, by injecting the indicated recombinase at a con-
centration of 2 µM in the presence of 2 mM ATP followed by a
20-min incubation at 30°C. Free protein was then flushed from
the sample chamber, followed by an injection of HR buffer (as in-
dicated above for each recombinase), and the reactions were incu-
bated for an additional 10 min. Unbound dsDNA was then
quickly flushed from the sample chamber using a 30-sec wash
at 1 mL/min, flow was then reduced to 0.2 mL/min, and images
(90-msec integration) were collected at 30- to 60-sec intervals for
1–2 h, as described previously [Lee et al. 2015, 2017; Qi et al.
2015]. The data collection intervals were optimized relative to
the overall lifetime of each dsDNA substrate, and the laser was
shuttered between acquired images to minimize photobleaching.
Kymographs were then generated from the resulting movies
using Fiji. Survival probabilities were determined from analysis.
of the resulting kymographs by measuring the time [dwell time] that each molecule of Atto565-dsDNA remained bound to the presynaptic complexes after flushing the unbound DNA from the sample chamber. Error bars for the survival probability measurements and binding distributions represent 70% confidence intervals obtained through bootstrap analysis, providing a close approximation of expectations for one standard deviation from the mean (Lee et al. 2015; Qi et al. 2015). All reported free energy calculations were performed using the following equation, as described previously (Lee et al. 2015, 2016, 2017; Qi et al. 2015; Qi and Greene 2016; Ma et al. 2017):

\[ \Delta \Delta G^\ddagger = \Delta G^\ddagger_2 - \Delta G^\ddagger_1 = k_b T \ln \frac{k_3^\ddagger}{k_2^\ddagger}. \]

where \( k_b \) is the Boltzmann constant, \( T \) is temperature, and \( \Delta \Delta G^\ddagger \) is the free energy difference between two escape processes described by the dissociation rates \( k_2^\ddagger \) and \( k_3^\ddagger \). The experimentally measured data used to calculate \( \Delta \Delta G^\ddagger \) were the dissociation rate data for the different oligonucleotide substrates [see Supplemental Table S4], which were obtained from the survival probability analysis for each different recombinase with the different dsDNA oligonucleotide substrates. The reported free energy values for analysis of base triplet stepping are indicated by double-ended arrows, and the step 1, step 2, and step 3 designations are also highlighted by color-coded schematics and arrowheads. For analysis of mismatch-bearing oligonucleotides, the \( \Delta \Delta G^\ddagger \) assigned for the first step was set to zero (“step 1” in figure diagrams in Figs. 2B, 3A–B), and the step 1–3 designations reflect the values obtained for fully paired substrates for each specific protein (as indicated by dashed color-coded lines).

Additional Materials and Methods are in the Supplemental Material.

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Effects of Rad51 and Dmc1 specific amino acids


Defining the influence of Rad51 and Dmc1 lineage-specific amino acids on genetic recombination

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