Mismatch Repair during Homologous and Homeologous Recombination

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Homologous recombination (HR) and mismatch repair (MMR) are inextricably linked. HR pairs homologous chromosomes before meiosis I and is ultimately responsible for generating genetic diversity during sexual reproduction. HR is initiated in meiosis by numerous programmed DNA double-strand breaks (DSBs; several hundred in mammals). A characteristic feature of HR is the exchange of DNA strands, which results in the formation of heteroduplex DNA. Mismatched nucleotides arise in heteroduplex DNA because the participating parental chromosomes contain nonidentical sequences. These mismatched nucleotides may be processed by MMR, resulting in nonreciprocal exchange of genetic information (gene conversion). MMR and HR also play prominent roles in mitotic cells during genome duplication; MMR rectifies polymerase misincorporation errors, whereas HR contributes to replication fork maintenance, as well as the repair of spontaneous DSBs and genotoxic lesions that affect both DNA strands. MMR suppresses HR when the heteroduplex DNA contains excessive mismatched nucleotides, termed homeologous recombination. The regulation of homeologous recombination by MMR ensures the accuracy of DSB repair and significantly contributes to species barriers during sexual reproduction. This review discusses the history, genetics, biochemistry, biophysics, and the current state of studies on the role of MMR in homologous and homeologous recombination from bacteria to humans.

Genetic recombination between unrelated parental DNAs during sexual reproduction appears to solve the problem of Muller’s ratchet, a process in which the accumulation of deleterious spontaneous mutations by asexual organisms eventually leads to extinction (Muller 1964; Felsenstein 1974). Recombination of chromosomes during sexual reproduction (meiosis) is considered to generate competitive genetic hybrids from two potentially noncompetitive mutant parents. The cytological observation of chromosome crossovers (COs) in the early days of Drosophila genetics led to their recognition as important intermediates in the formation of genetic hybrids (Muller 1916). These COs were given a molecular framework...
with the theoretical description of the Holliday junction (Holliday 1964). The fundamental insight of a Holliday junction was that single DNA strands from two parental chromosomes criss-cross to form hybrid DNA duplexes. Resolution of the crossed strands of a Holliday junction may then result in CO of genes on either side of the hybrid DNA duplexes. Sequence differences between the parental DNAs were proposed to result in mismatched nucleotides within the hybrid DNA duplex.

The repair of mismatched nucleotides was proposed nearly simultaneously in 1964 by Evelyn Witkin to account for the processing of brominated nucleotides in bacteria and by Robin Holliday to explain gene conversion following recombination (Holliday 1964; Witkin 1964). Mismatch repair (MMR) was proposed to recognize nucleotide mismatches and perform an excision–resynthesis reaction in which one strand is excised within the hybrid DNA duplex and the remaining strand is used as a template for resynthesis. In 1973, a genetic basis for MMR was discovered when the HexA mutation of Pseudomonas was found to be defective in gene conversion (Tiraby and Fox 1973). The HexA gene turned out to be a homolog of the "Siegel mutator" (MutS), originally described by Eli Siegel in 1967 (Siegel and Bryson 1967). The discovery of MutS added to a growing number of genes with historical roots in the 1954 genetic description of the “Treffers mutator” (MutT) (Treffers et al. 1954). Mutation of these Mut genes substantially elevated spontaneous mutation rates in bacteria (term mutator). Today, most of the Mut genes are known to play a role in the processing of replication misincorporation errors, DNA double-strand breaks (DSBs), and chemical or physical damage to nucleotides (for a review, see Miller 1998; Ciccia and Elledge 2011).

In the early 1980s, a series of clever observations from a number of laboratories showed that a subset of the Escherichia coli mutator genes, MutS, MutL, MutH, and UvrD(MutU), operated in the MMR of polymerase misincorporation errors that occurred frequently during replication (Radman et al. 1980). The recognition of transient undermethylation (hemimethylation) at newly replicated DNA adenine methylation (Dam) GATC sequences was identified as the mechanism for discriminating the error-containing DNA strand (Marinus 1976). Notably, this Dam-directed postreplication MMR mechanism only appears to operate in a subset of γ-proteobacteria, such as E. coli. The mechanism for strand discrimination in Eubacteria, Archa, and Eukaryotes remains speculative. However, as might be expected for important genome maintenance processes, the core MutS homologs (MSHs) and MutL homologs (MLHs)/postmeiotic segregation (PMS) of MMR, have been highly conserved throughout the taxonomic domains (Table 1).

In 1993, a human MSH (HsMSH2) was found to be associated with the common cancer predisposition syndrome hereditary nonpolyposis colorectal cancer (HNPPC; Fishel et al. 1993). Verification of this association (Leach et al. 1993) and the rapid successive discovery of other MSH and MLH/PMS homolog genes linked with HNPPC and sporadic colorectal cancers have solidified a role for MMR defects in tumor development (Bronner et al. 1994; Nicolaides et al. 1994; Papadopoulos et al. 1994, 1995). These observations also underlined the importance of mutators in driving the enormous numbers of mutations required in the microevolutionary selection processes associated with many forms of tumorigenesis (Loeb 1991, 2001; Fishel 2001).

HOMOLOGOUS RECOMBINATION

In addition to meiosis, genetic recombination plays a significant role in somatic cells during the repair of chemical damage to DNA (particularly chemical cross-links), DSBs introduced by physical damage to the DNA, and in the restoration of damaged replication forks (reviewed in Friedberg et al. 2006). If these lesions persist, chromosomal fragmentation, chromosomal loss (aneuploidy), and genetic rearrangements may occur (Kanaar et al. 1998; Rich et al. 2000). DSBs are repaired via homologous recombination (HR), which includes synthesis-dependent strand annealing (SDSA), single-strand annealing (SSA), and nonhomologous end joining.
HR and SDSA faithfully repair DSBs without gain or loss of DNA sequences (Jasin and Rothstein 2013). In contrast, SSA is a mutagenic pathway used by cells when the DSB, committed to homology-directed repair, cannot be timely processed by HR or when the DSB occurs between direct repeats (Weinstock et al. 2006; Moynahan and Jasin 2010). NHEJ mends DNA ends, but almost always results in sequence gain or loss surrounding the site of the DSB. Additionally, the NHEJ process has a high risk of inducing chromosomal translocation because it may fuse virtually any broken end (Lieber 2010; Chiruvella et al. 2013).

This review focuses entirely on HR and SSA because nucleotide mismatches that arise during these processes account for the outcome of the genetic products, as well as regulate the initiation of genetic exchange.

Studies with fungi have provided a superb model for studying HR. It should be noted that, like MMR, the genes and mechanical processes of HR are highly conserved up to and including humans. The general purpose and outcomes of HR in *Saccharomyces cerevisiae* are fundamentally different in meiosis and mitosis (reviewed in Andersen and Sekelsky 2010). The main function of HR between paternal and maternal chromosomes in meiosis is to create at least one CO per pair of homologs (reviewed in Page and Hawley 2003; Szekvolgyi and Nicolas 2009). Meiotic COs convert sister chromatid cohesion into homologous chromosome clasps, which ensure accurate segregation in the first meiotic division (meiosis I). The partition of chromosome arms on either side of a CO is the physical basis of Mendelian inheritance and, ultimately, genetic diversity.

The two distinct outcomes of meiotic recombination were first observed following tetrad analysis in *S. cerevisiae* (Lindgren 1955): (1) conserved crossovers, in which genetic information is exchanged reciprocally between chromosomes without genotype gain or loss; and (2) gene conversion, in which the genetic information of one genotype is replaced with an allelic genotype. The gene conversion process may result in CO or noncrossover (NCO) of markers on either side of the gene conversion event. Asymmetric repair of the mismatched nucleotides contained within the hybrid DNA of the Holliday junction was conceived to ac-

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Table 1. Parallels between the proteins involved in MMR and heteroduplex rejection during recombination

<table>
<thead>
<tr>
<th><strong>Escherichia coli</strong></th>
<th><strong>S. cerevisiae</strong></th>
<th><strong>Human</strong></th>
<th><strong>Function</strong></th>
<th><strong>Role</strong></th>
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<tr>
<td>MutS</td>
<td>ScMsh2-ScMsh6</td>
<td>HsMSH2-HsMSH6</td>
<td>Recognition of mismatches and small IDLs; sliding clamp</td>
<td>MMR; ICL repair; gene conversion; heteroduplex rejection</td>
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<td></td>
<td>ScMsh2-ScMsh3</td>
<td>HsMSH2-HsMSH3</td>
<td>Recognition of large IDLs and branched structures; sliding clamp</td>
<td>Postreplicative repair of IDLs; 3'-flap processing; SSA intermediate stabilization</td>
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<tr>
<td>MutL</td>
<td>ScMlh1-ScPms1</td>
<td>HsMLH1-HsPMS2</td>
<td>Downstream mediator; endonuclease</td>
<td>MMR; gene conversion</td>
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<td>?</td>
<td>ScRad1-ScRad10</td>
<td>HsXPF-HsERCC1</td>
<td>Structure-selective nuclease</td>
<td>3'-flap removal</td>
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<td>RecJ</td>
<td>ScExo1</td>
<td>HsEXO1</td>
<td>5' → 3' exonuclease</td>
<td>MMR; HR</td>
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<td>ExoVII</td>
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<td>Bidirectional exonuclease</td>
<td>MMR</td>
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<td>Exol, ExoX</td>
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<td>3' → 5' exonuclease</td>
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<td>UvrD</td>
<td>ScSgs1</td>
<td>HsRECQ1</td>
<td>Structure-selective DNA helicase</td>
<td>MMR; unwinding heteroduplex DNA</td>
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<td>HsBLM</td>
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ICL, Interstrand crosslinks; IDL, insertion–deletion loops; MutS, Siegel mutator; MMR, mismatch repair; SSA, single-strand annealing; MutL, *E. coli* mutator gene; HR, homologous recombination.
count for gene conversion (Holliday 1964). Interestingly, the absence of MMR may lead to PMS of genetic markers contained within the heteroduplex DNA of a Holliday junction, eventually leading to sectored colonies containing both genotypes. This sectored colony phenotype was used in *S. cerevisiae* to isolate the first eukaryotic MMR gene, the MLH/PMS ScPms1 (Williamson et al. 1985), initiating the beginnings of some muddled nomenclature.

To explain the results of plasmid integration (gap-repair) studies in yeast, the double-strand break repair (DSBR) model was proposed (Szostak et al. 1983). The DSBR model and its variants are now thought to engender the major mechanism(s) associated with HR (Fig. 1) (see Mehta and Haber 2014). DSBR in meiotic recombination is initiated by the introduction of a large number of programmed DSBs catalyzed by the ScSpo11 protein (see Keeney et al. 1997; Lam and Keeney 2015). In mouse and human cells, these programmed DSBs number in the several hundred (Kolas et al. 2005; Lenzi et al. 2005). It is the repair of these numerous DSBs, using sequences donated from a homologous parental chromosome, which ultimately links chromosome homologs before meiosis I. At a molecular level, the DSB ends are first resected to produce

![Figure 1](http://cshperspectives.cshlp.org/) Multiple roles of the mismatch repair (MMR) machinery in eukaryotic double-strand break (DSB) repair by homologous recombination (HR). The damaged DNA molecule is shown in blue, the template is shown in black, and the strand synthesized during the double-strand break repair (DSBR) process is shown in red. The heteroduplex rejection steps are represented by the red arrows. Major players involved in every step are indicated. (A) Schematic representation of the DSB repair by single-strand annealing (SSA). The direct repeats on either side of the DSB are shown as arrows. (B) Synthesis-dependent strand annealing (SDSA). See text for details. CO, Crossover; dsDNA, double-stranded DNA; HJ, Holliday junction; NCO, noncrossover.
3’-single-stranded DNA (ssDNA) overhangs (reviewed in Symington and Gautier 2011; see Symington 2014). This process is followed by precisely coordinated DNA transactions, which are orchestrated by highly conserved homologs/orthologs of the prototypical RecA/RAD51 DNA homologous pairing and strand exchange proteins (recombinase) (see Li and Heyer 2008; Zelensky et al. 2014; Morrical 2015). In *S. cerevisiae*, and with the help of recombination mediators, the ScRad51 and/or the RecA/RAD51 paralog ScDmc1 form a nucleoprotein filament (NPF) on the 3’-ssDNA overhangs (Sung 1997; New et al. 1998; Shinohara and Ogawa 1998; Gasior et al. 2001; Liu et al. 2010, 2011; Carreira and Kowalczykowski 2011; Kojic et al. 2011; Murayama et al. 2013; Sasanuma et al. 2013; Daley et al. 2014). One of the RecA/RAD51 NPFs assembled on the 3’-rescissed ends of the DSB then catalyzes the invasion of a homologous parental double-stranded DNA (dsDNA; donor template) forming a displacement loop (D-loop). DNA synthesis, initiated at the 3’ end of the invading strand, then expands the D-loop until the distal end captures the remaining NPF, ultimately forming a double Holliday junction. Varying lengths of heteroduplex DNA may be produced by branch migration of either or both of the individual Holliday junctions (reviewed in Jasin and Rothstein 2013). These Holliday junctions may then be dissolved or resolved to yield either a CO or NCO (see Bizard and Hickson 2014; Wyatt and West 2014). CO and NCO events are readily distinguished in a plasmid gap-repair assay, in which an NCO outcome yields a repaired autonomously replicating plasmid, whereas a CO event integrates the plasmid into genome (Orr-Weaver and Szostak 1983).

The occurrence of mismatches during meiotic recombination has been documented in genetic experiments via the manifestation of sectored PMS colonies, as well as through physical methods (White et al. 1985; Lichten et al. 1990). Correction of mismatched meiotic recombination intermediates by the MMR machinery results in gene conversion (Alani et al. 1994). Although the precise mechanism(s) associated with the processing of meiotic mismatched (heteroduplex) DNA remain obscure, it is likely that there is significant similarity with postreplication MMR (reviewed in Kolodner and Marsischky 1999; Kolodner et al. 2007).

In contrast to meiosis, mitotic recombination is critical for the repair of spontaneous DSBs and collapsed replication forks. It is significantly biased toward NCOs because the unidirectional transfer of information, which consequently results in the removal of any impediments to chromosome segregation, appears essential for high-fidelity repair. Only a small fraction of mitotic events proceed through the classic DSBR pathway (Bzymek et al. 2010). The majority of DSB repair occurs by an SDSA HR mechanism (Fig. 1) (Andersen and Sekelsky 2010). SDSA begins with DNA strand invasion and D-loop formation similar to classical DSBR. The 3’ end of the invading ScRad51 NPF is then extended by polymerase δ-catalyzed DNA synthesis (Li et al. 2009), which is essential to restore any DNA sequences that might be degraded during the exonucleolytic processing of the DSB. The major difference between SDSA and the DSBR model is that the D-loop is disassembled and the proximal polymerase-extended ssDNA strand is then used to capture the distal ScRad51 NPF (Fig. 1). Any DNA gaps and/or unannealed DNA flaps may then be processed and the remaining strand scissions are ligated to produce an intact duplex DNA. DSB repair by SDSA inevitably results in NCO events.

Three helicases in *S. cerevisiae* have been implicated in regulating HR events. These include the slow growth suppressor of a topoisomerase 3 (Top3) mutation ScSgs1, the suppressor of Rad6 sensitivity ScSrs2, and the mutator phenotype ScMph1 (Ira et al. 2003; Prakash et al. 2009; Heyer et al. 2010). All three helicases have been either directly or genetically shown to dismantle and/or prevent the formation of D-loop structures. The most recent genetic evidence implicates ScMph1 in the disassembly of D-loops, whereas ScSgs1 dissolves intact double Holliday junctions and ScSrs2 dismantles single and double Holliday junctions containing strand scissions (Mitchel et al. 2013). In all cases, dissolution of these recombination intermediates results in NCO events. Additional bias toward NCO events during mitotic recombina-
tion, as well as the CO/NCO balance during meiotic recombination, may also be achieved by linking tight regulation of structure-selective nucleases to cell-cycle progression (Matos et al. 2011; Matos and West 2014).

In both meiosis and mitosis, the numerous steps associated with the recombination event must discriminate between identical and non-identical template sequences and then either repair, process, or reject the ensuing heteroduplex DNA. In contrast to meiosis, which tolerates some degree of sequence heterology, mitotic recombination is extremely sensitive to the presence of mismatched nucleotides. This exquisite sensitivity is generally not caused by the selectivity of the RecA/RAD51 recombinase, but rather depends on the MMR machinery (Datta et al. 1997).

HOMEOLOGOUS RECOMBINATION

In 1989, Radman and colleagues suggested that, in addition to resolving recombination intermediates during gene conversion, MMR introduces a barrier to recombination between species containing excessive sequence heterology (Rayssiguier et al. 1989). The experimental system was relatively simple. *E. coli* and *Salmonella typhimurium* are 97% identical at the DNA sequence level. Yet, recombination between well-defined genetic markers contained in their largely identical chromosomes is quite rare; occurring at a frequency of $10^{-6}$ – $10^{-8}$ depending on the genetic marker (Rayssiguier et al. 1989; Stambuk and Radman 1998). This is in spite of the fact that F-pillus formation and DNA transfer between these organisms occurs quite readily (Clark and Adelberg 1962; Curtiss 1969). Remarkably, when the recipient bacteria are MMR deficient, the frequency of recombination between several genetic markers increased over 1000-fold (Rayssiguier et al. 1989; Stambuk and Radman 1998). These results were interpreted to suggest that MMR either aborts the formation of recombination intermediates or resolves nascent recombination intermediates containing excess heterology such that genetic recombination rarely occurs. Genetic exchange between partially homologous sequences was termed “homeologous” recombination to distinguish it from HR between fully homologous chromosomal DNA.

THE GENETIC REQUIREMENTS FOR THE SUPPRESSION OF HOMEOLOGOUS RECOMBINATION IN E. COLI

The suppression of homeologous recombination in *E. coli* required the *EcMutS*, *EcMutL*, *EcMutH*, and *EcUvrD* genes (Table 1) (Rayssiguier et al. 1989; Stambuk and Radman 1998). However, the relative suppression of homeologous recombination was quite different between these MMR genes. A broad pathway analysis determined that the *mutS* mutation increased homeologous recombination 735-fold (Stambuk and Radman 1998). Although the requirement for *MutL* was not examined in this pathway analysis, the initial study suggested that *MutS* and *MutL* affected the magnitude of homeologous recombination similarly. In contrast, a *mutH* mutation only increased homeologous recombination 22-fold, whereas mutation of *uvrD* increased homeologous recombination fivefold. These observations were qualitatively similar to the initial study (Rayssiguier et al. 1989). Interestingly, the combined *mutS uvrD* mutation increased homeologous recombination approximately two times over *mutS* alone, whereas the combined *mutH uvrD* mutation increased homeologous recombination almost tenfold over *mutH* alone. Mutation of genes involved in recombination initiation, Holliday junction branch migration, or the up-regulation of damage-inducible genes had little or no effect on homeologous recombination (Rayssiguier et al. 1989). The additive nature of *uvrD* mutations with MMR mutations suggested that there were likely two redundant pathways capable of aborting homeologous recombination in *E. coli*: one involving the core MMR genes MutHLS and a second involving the UvrD helicase. The separation of these pathways appeared most convincing in the absence of *mutH* than *mutS*. The biochemical properties of these proteins provide a foundation to help explain these distinctions.
BIOCHEMICAL ACTIVITIES OF THE MMR PROTEINS

MMR is a mismatch or nucleotide lesion-dependent DNA excision-resynthesis DNA repair process (Modrich 1989, 1997; Kolodner and Marsischky 1999; Fishel et al. 2000). Strand excision may be initiated by several hundred to several thousand base pairs on either the 3'- or 5'-side of the mismatch, and the excision tract extends from this distant site to just past the mismatch (Lahue et al. 1987). A number of models have been proposed to account for the biochemical properties of the individual MMR proteins and the ability to initiate excision bidirectionally at a site significantly distant from the mismatch/lesion (Fig. 2) (Acharya et al. 2003; Kolodner et al. 2007). The mechanism that is most consistent with the data and the Brownian nature of molecular biology appears to be the molecular switch model (Gradia et al. 1997, 1999; Fishel 1998, 2000; Acharya et al. 2003; Jeong et al. 2011; Cho et al. 2012; Gorman et al. 2012; Qiu et al. 2012; Spies 2013). Most, if not all, biochemical discrepancy can be traced to differences in the experimental conditions, an issue that persists today (Hall et al. 2001; Drotschmann et al. 2002; Tessmer et al. 2008; Sass et al. 2010; Tham et al. 2013).


MSH proteins are members of the AAA⁺ ATPase family and contain a highly conserved Walker A/B nucleotide-binding motif (Walker et al. 1982; Iyer et al. 2004). Mismatch nucleotide recognition was first demonstrated in 1986 with the EcMutS (Su and Modrich 1986). Remarkably identical structures of numerous MSH proteins bound to mismatched DNA have appeared in the literature in succeeding years (Lamers et al. 2000; Obmolova et al. 2000; Warren et al. 2007; Gupta et al. 2011). These structures showed that bacterial and human MSHs form a clamp-like configuration around the mismatched DNA with a Phe-residue within a highly conserved Phe-X-Glu motif interrogating the DNA 3' of the mismatch and inducing a 45°–60° bend. Additional nonspecific electrostatic contacts with the DNA by protein residues surrounding the Phe-X-Glu motif further stabilize the mismatch-bound structure. Only nucleotide-free or ADP-bound structures have been crystallized. Infusion of ATP or ATPγS destroys the crystals (Obmolova et al. 2000), suggesting significant differences in the mismatch-bound and postmismatch conformations.

Multiple functionally distinct forms of the MSH clamp on DNA have been detected using single-molecule biophysical imaging analysis (reviewed in Spies 2013). While searching for a mismatch on duplex DNA, the Thermus aquaticus TₐMutS was revealed to form an incipient clamp that underwent facilitated 1D rotational diffusion while in continuous contact with the helical backbone (Fig. 2A) (Cho et al. 2012). A similar mismatch search mechanism has been observed for the ScMsh2-ScMsh6 heterodimer (Gorman et al. 2012) and is likely conserved in all MSH proteins (Heinen et al. 2011). On encountering a mismatch, MutS pauses (presumably forming the clamp structure shown in structural studies), releases bound ADP, and binds ATP (ADP → ATP exchange) (Acharya et al. 2003; Jeong et al. 2011). DNA flexibility surrounding the mismatch has been proposed to distinguish it from the normally smooth DNA backbone triggering the pause in MSH diffusion (Mazurek et al. 2009). Detection of DNA contour alterations, and not the mismatch itself, would explain the wide range of mismatch/lesion recognition properties shown by MSH proteins (Mazurek et al. 2009).

Mismatch, lesion, and/or structure provoked ADP → ATP exchange is a central function of all MSH proteins examined to date.
A TP binding by the MSH dimer/heterodimer instigates a significant conformational transition (Gradia et al. 1997; Wilson et al. 1999; Acharya et al. 2003; Snowden et al. 2004), which results in an extremely stable (5- to 10-min lifetime) sliding clamp that freely diffuses along the DNA without A TP hydrolysis and in the absence of any stable backbone interactions (Fig. 2A) (Cho et al. 2012). A TP binding and release of the MutS from the mismatch allows the loading of multiple MutS A TP-bound sliding clamps (Fig. 2B) (Gradia et al. 1997; Acharya et al. 2003; Jeong et al. 2011; Cho et al. 2012). 1D diffusion by the A TP-bound MSH sliding clamp is iterative with multiple MSH sliding clamps and MLH/PMS interactions followed by dynamic, likely small, excision events that ultimately release the entire MSH-MLH/PMS complex. Although not entirely known, the strand to be excised and repaired may be distinguished by endonuclease discontinuities and/or a 3′ or 5′ end. The bacterial β-clamp and eukaryotic proliferating cell nuclear antigen (PCNA), along with their respective clamp loading factors γ-complex and replication factor C (RFC), as well as the single-strand binding protein (SSB) and replication protein A (RPA), respectively, play important but not fully defined roles in the processes outlined in A and B. (C,D) The bidirectional nature of MSH-MLH/PMS diffusion ensures that excision may take place bidirectionally on either side of the mismatch in which an appropriate end occurs. (E) The multiple nucleolytic events remove the DNA region from the initial strand scission to just past the mismatch. Once the mismatch is removed, the loading of MSH sliding clamps is halted and the excision reaction ultimately ceases. Mismatch excision requires strand-specific exonucleases (3′ → 5′ ExoI, 3′ → 5′ ExoX, bidirectional ExoVII, or 5′ → 3′ RecJ in Escherichia coli and 5′ → 3′ EXO1 in eukaryotes). Resynthesis of the excision gap is thought to be accomplished by the replicative polymerase Pol I in bacteria and POL-δ in eukaryotes.

Figure 2. Molecular switch model for eukaryotic mismatch repair (MMR). (A) An MutS homolog (MSH) scanning clamp binds dsDNA and searches for rare nucleotide misincorporation replication errors by one-dimensional (1D) rotational diffusion while maintaining continuous contact with the DNA duplex. (B) An encounter with a mismatch and subsequent ADP → A TP exchange converts the MSH into a sliding clamp, which dissociates from the mismatch, but still encircles the DNA duplex and moves by 1D diffusion while in discontinuous contact with the DNA backbone. The MutL homolog (MLH)/postmeiotic segregation (PMS) heterodimer associates with the A TP-bound MSH sliding clamp. The interaction between MSH with MLH/PMS, and likely some DNA structure, promotes A TP binding by the MLH/PMS, which activates its intrinsic endonuclease activity. This process is iterative with multiple MSH sliding clamps and MLH/PMS interactions followed by dynamic, likely small, excision events that ultimately release the entire MSH-MLH/PMS complex. Although not entirely known, the strand to be excised and repaired may be distinguished by endonuclease discontinuities and/or a 3′ or 5′ end. The bacterial β-clamp and eukaryotic proliferating cell nuclear antigen (PCNA), along with their respective clamp loading factors γ-complex and replication factor C (RFC), as well as the single-strand binding protein (SSB) and replication protein A (RPA), respectively, play important but not fully defined roles in the processes outlined in A and B. (C,D) The bidirectional nature of MSH-MLH/PMS diffusion ensures that excision may take place bidirectionally on either side of the mismatch in which an appropriate end occurs. (E) The multiple nucleolytic events remove the DNA region from the initial strand scission to just past the mismatch. Once the mismatch is removed, the loading of MSH sliding clamps is halted and the excision reaction ultimately ceases. Mismatch excision requires strand-specific exonucleases (3′ → 5′ ExoI, 3′ → 5′ ExoX, bidirectional ExoVII, or 5′ → 3′ RecJ in Escherichia coli and 5′ → 3′ EXO1 in eukaryotes). Resynthesis of the excision gap is thought to be accomplished by the replicative polymerase Pol I in bacteria and POL-δ in eukaryotes.
clamp(s) has been calculated to easily cover the several thousand nucleotides between the mismatch and the DNA strand scission where excision begins.

Long-lived ATP-bound hydrolysis-independent MSH sliding clamps are the single most critical intermediates in initiating MMR and suppression of homeologous recombination. This conclusion is supported by a number of genetic and biochemical observations. First, ATP-binding or hydrolysis-deficient MSH mutations, located in the conserved Walker ATP-binding motif, retain mismatch-binding activity, but are deficient in both MMR and homeologous recombination (Haber and Walker 1991; Wu and Marinus 1994; Iaccarino et al. 1998; Hess et al. 2002; Junop et al. 2003; Lin et al. 2004; Mendillo et al. 2005; Ollila et al. 2008). Second, the ability to form a sliding clamp strictly correlates with biological function, whereas mismatch/lesion/structure binding is necessary, but not sufficient for biological function (Sia et al. 1997; Genschel et al. 1998; Wilson et al. 1999; Snowden et al. 2004; Lenzi et al. 2005; Mendillo et al. 2005; Hargreaves et al. 2010). Finally, titration studies suggest that 4–8 MSH molecules are required for a single repair event in vitro (Zhang et al. 2005; Goeliner et al. 2014). Taken together, these observations clearly suggest that for biochemical or biophysical studies to fully reflect the cellular function(s) of MSH proteins, it is important to select conditions in which sliding clamp formation is robust and not dramatically reduced or absent. Besides the obvious prerequisite for ATP or the poorly hydrolysable analog ATPγS (MSH proteins do not readily bind AMP-PNP or AMP-PCP), ionic conditions between 90 mM and 150 mM are essential for the formation of stable mismatch-dependent ATP-bound MSH sliding clamps (Gradia et al. 1999; Wilson et al. 1999; Hess et al. 2002; Acharya et al. 2003; Mendillo et al. 2005; Mazur et al. 2006; Jeong et al. 2011). A requirement for specific ionic conditions is not surprising considering the nonspecific electrostatic contacts observed with MSH-DNA complexes outside the conserved Phe-X-Glu motif.

The EcMutH protein uniquely recognizes a hemimethylated Dam (GATC/GAmεTC) site and introduces a single-strand scission between the guanine and unmethylated adenine nucleotides (Welsh et al. 1987). It is this distant strand scission where the MMR excision reaction starts in γ-proteobacteria, such as E. coli. No other biochemical activity for EcMutH has been identified. Moreover, a preexisting strand scission appears to completely eliminate the EcMutH requirement for MMR in vitro (Lahue et al. 1987). EcUvrD is a 3′ → 5′ helicase (Matson 1986; Matson and George 1987), which is activated by EcMutL (Dao and Modrich 1998; Hall et al. 1998). Its likely role in MMR is to unwind the excised DNA strand before and/or in concert with exonuclease digestion (Lee and Yang 2006). One could envisage a number of different mechanistic strategies for the rejection of homeologous recombination intermediates to account for the known genetic requirements. For example, the EcMutH hemimethylated strand scission function might activate MMR strand excision following D-loop-dependent DNA synthesis. Alternatively, EcUvrD helicase might disassemble a homeologous D-loop intermediate. In both of these hypothetical mechanisms, it is likely that EcMutS-mediated mismatch detection would be required to target the homeologous recombination intermediate.

The detailed function of MLH/PMS proteins in MMR remains enigmatic. MLH/PMS proteins contain a GKL ATP-binding domain (Dutta and Inouye 2000), which displays extremely weak ATPase activity (Ban et al. 1999; Spampinato and Modrich 2000; Acharya et al. 2003). Moreover, the yeast MLH/PMS, ScMlh1-ScPms1, appears to undergo conformation contractions between the carboxy-terminal heterodimer interaction domain and the amino-terminal ATP-binding and adenosine nucleotide-dependent dimerization domain (Sacho et al. 2008). The function, if any, of these conformational transitions is unknown. ATP binding by EcMutL significantly enhances the EcMutH endonuclease activity (Acharya et al. 2003). Importantly, EcMutL and MLH/PMS heterodimer proteins form a stable complex with ATP-bound EcMutS and MSH sliding clamps (Fig. 2B) (Acharya et al. 2003; Mendillo et al. 2005). The diffusion characteristics of the
MSH-MLH/PMS complex do not appear to be different from the MutS/MSH sliding clamp alone (Gorman et al. 2012). These observations strongly suggest that the ATP-bound MutS/MSH sliding clamp recruits and transports the MutL/MLH/PMS for downstream MMR functions. Potential functions for EcMutL in bacterial MMR may be to stabilize the EcMutH endonuclease at the hemimethylated GATC site and/or enhance EcUvrD helicase activity during excision. Interestingly, EcMutL and ScMlh1-ScPms1 have been shown to bind ssDNA (Drotschmann et al. 2002; Park et al. 2010). However, EcMutL ssDNA-binding activity is nearly undetectable at physiological ionic strength and unlikely to be significant during MMR (Park et al. 2010).

A major difference between *E. coli* MutL and virtually all MLH/PMS proteins outside γ-proteobacteria is the presence of an intrinsic endonuclease activity (Kadyrov et al. 2006, 2007; Pillon et al. 2010). It has been suggested that this MLH/PMS intrinsic endonuclease substitutes for the MutH endonuclease activity found only in γ-proteobacteria like *E. coli* (Kadyrov et al. 2006). This parallel seems, at the very least, incomplete because the MLH/PMS endonuclease appears to introduce multiple strand scissions during the MMR reaction (Kadyrov et al. 2006). The MLH/PMS endonuclease appears more efficient in the presence of manganese-divalent cation and can also be stimulated by zinc (Kadyrov et al. 2006, 2007; Pillon et al. 2010). Importantly, the *Thermus thermophilus* homologs were used to show that the TtMutL ATP-dependent endonuclease is activated only on its association with ATP-bound TtMutS sliding clamps (Shimada et al. 2013).

The molecular switch model, as applied to γ-proteobacteria, suggests that multiple MutS/MutL complexes may diffuse along the DNA to activate MutH incision, UvrD helicase, and exonuclease excision (Fig. 2A,B) (Acharya et al. 2003). The process is proposed to be dynamic such that successive MutS-MutL-UvrD-exonuclease complexes produce redundant small excision tracts, which ultimately discharge the mismatch. For organisms that do not contain a MutH and do not use a helicase during MMR (Eubacteria, Archea, and Eukaryotes), the MLH/PMS ATP-dependent endonuclease presumably associated with ATP-bound MSH sliding clamps, and an exonuclease could conceivably produce small excision tracts by a similar dynamic and redundant mechanism (Fig. 2C,D) (Fishel et al. 2000). Once the excision tract releases the mismatch, successive loading of MutS sliding clamps ceases and the DNA degradation reaction grinds to a halt. A free 3′ end would allow the polymerase machinery to assemble and complete the resynthesis reaction (Fig. 2E).

**THE SUPPRESSION OF HOMELOGOUS RECOMBINATION IN YEAST**

DSBs and the formation of early recombination intermediates in meiosis are required to establish stable pairing interactions between homologs (Peoples et al. 2002). When formed in unique nonrepetitive sequences, meiotic DSBs initiate recombination between identical allelic positions on homologous chromosomes. DSBs can also form within or near repetitive elements that are found in abundance in all eukaryotes (Lander et al. 2001). Recombination between such repeats carries a risk for deleterious genome rearrangements in the germ line. As with bacteria, in eukaryotes, the MMR machinery is largely responsible for the rejection of heteroallelic homeologous DNA during recombination (Datta et al. 1996). Both the precise mechanism of homeologous recombination rejection and molecular triggers, which signal to either repair or reject these partially homologous sequences, remain unknown.

How many mismatches are required to distinguish homologous from homeologous recombination? A log-linear relationship between sequence divergence and sexual isolation (frequency of recombination) was observed for soil isolates of *Bacillus subtilis* (Roberts and Cohan 1993). In these studies, as little as 0.3% sequence divergence reduced recombination, thus, increasing sexual isolation. Although a complete answer to this question may vary between organisms, in *S. cerevisiae*, one mismatch in a recombination region of ~300 bp (0.3% sequence divergence) reduced the recombination rate al-
most threefold, and that rate decreased logarithmically until ~1% sequence divergence was reached (Datta et al. 1997). The rate decrease in recombination then becomes linear over three orders of magnitude in sequence divergence (Datta et al. 1997). Remarkably, the impact of MMR in suppressing recombination increased steadily until ~10% sequence divergence (Datta et al. 1997). The recombination rate between sequences with >10% sequence divergence decreased dramatically and appeared to be largely unaffected by MMR, suggesting that genetic exchange was most likely aborted before any actions by the core MMR machinery. Similar results have been reported for the effect of sequence divergence and MMR on Arabidopsis thaliana recombination (Li et al. 2006).

The MMR machinery is required for both recognizing and processing mismatch-containing heteroduplex during homologous recombination in yeast (Fig. 1). Genetically, msh2, msh3, and msh6 play critical roles in the heteroduplex rejection, suggesting the roles for both ScMsh2-ScMsh3 and ScMsh2-ScMsh6 heterodimers (Fig. 1). The antirecombination activity of ScMsh2 is severalfold greater than that of the ScPms1 or ScMlh1 (Datta et al. 1996). These observations appear qualitatively similar to E. coli and suggest redundancy or bifurcation into subpathways in which both rejection mechanisms depend on ScMsh2, but only one requires both MSH and MLH/PMS. The ScSgs1 and ScSrs2 helicases play additional roles in antagonizing homologous recombination, but have no known roles in postreplicative MMR (Welz-Voegele and Jinks-Robertson 2008). Notably, deletion of either msh2 or sgs1 increases homologous recombination to the level of HR (Welz-Voegele and Jinks-Robertson 2008). These observations appear similar to bacteria and suggest the existence of at least two heteroduplex rejection pathways associated with HR (Fig. 1).

HETERODUPEX REJECTION DURING SSA BETWEEN DIRECT REPEAT SEQUENCES

SSA represents the major mechanism for the repair of DSBs that occur between direct repeats (Fig. 3) (Fishman-Lobell et al. 1992). As with the classical DSBR and the SDSA pathways, the ends of the DSB are resected in SSA (Fig. 3, left). If resection covers two complementary regions on either side of the DSB, the exposed ssDNAs are annealed with the help of the ScRad52 protein (Fig. 3, left). Although the SSA pathway is quite robust, it is also utterly mutagenic as it results in the loss of coding sequence between the repeats. SSA has been extensively used to investigate heteroduplex rejection during yeast mitosis because it provides a straightforward process for generating heteroduplex in vivo. As with DSBR and SDSA, SSA is sensitive to

Figure 3. Roles of the mismatch repair (MMR) proteins in single-strand annealing (SSA). Schematic representation of the double-strand break (DSB) repair by SSA. The direct repeats on either side of the DSB are shown as arrows. See the text for details of the rejection (left) and the mismatch correction (right) mechanisms.
the presence of heterology (Sugawara et al. 1997). Heteroduplex rejection in SSA depends on the ScMsh2-ScMsh6 protein, which recognizes mismatches within the heteroduplex, and the Sgs1 helicase, which presumably unwinds heteroduplex DNA strands (Fig. 1B, left) (Sugawara et al. 2004). The SSA heteroduplex rejection process is also significantly reduced in \textit{mlh1}Δ and \textit{pms1}Δ, \textit{mlh2}Δ, \textit{mlh3}Δ triple mutant cells, albeit to a much lesser extent than that observed in the \textit{msh6}Δ strain (Sugawara et al. 2004). These results are consistent with the conclusion that SSA heteroduplex rejection requires ScMsh2-ScMsh6 and one of the three known \textit{S. cerevisiae} MLH/PMS heterodimers (ScMlh1-ScPms1, ScMlh1-ScMlh2, ScMlh1-ScMlh3) (Sugawara et al. 2004).

Although mismatch recognition and unwinding of the heteroduplex DNA produced during DSBR and SSA likely share many mechanistic features, the two processes should not be completely equated. Mismatch recognition within a D-loop produced by a ScRad51 NPF may present a unique set of challenges distinct from those associated with the SSA structure annealed by the ScRad52 protein. For example, the three-stranded structure of a D-loop is clearly distinct from the simple heteroduplex produced by ScRad52-dependent annealing of resected stands. Moreover, the heteroduplex region of the D-loop likely remains coated with ScRad51 after the strand exchange step, perhaps until it is removed by ScRad54 (Li and Heyer 2009). Both ScRad52 (Sugiyama et al. 1998) and HsRad52 after it is activated by phosphorylation (Honda et al. 2011) are exquisitely selective for ssDNA and the homology search mechanism depends on diffusion (Rothenberg et al. 2008). These observations suggest a largely protein-free heteroduplex region following SSA.

**THE MMR SYSTEM IN HETEROLOGOUS FLAP PROCESSING**

When free homologous ends are available, the RAD51 recombinase NPF forms plectonemic joints in which the 3’ end of the invading strand is paired to the template DNA duplex. This structure then serves as a template for DNA synthesis, as discussed above for SDSA. When fully complementary ends are not available, the exchange of the DNA strands may produce less stable paranemic joints in which the two complementary strands are not topologically interwound. Removal of the protruding heterologous flap by a structure-specific nuclease, such as ScRad1-ScRad10, both converts a paranemic joint into a more stable plectonemic joint and produces a substrate for DNA synthesis (Fig. 3, right). Similarly, nonhomologous flaps must be removed from the ends of the heteroduplex formed during SSA. Because it is likely that the ends of the resected DSB will not be complementary, virtually every SSA event will almost certainly require a flap-processing step. The 3’-protruding end removal in D-loop and SSA intermediates has been extensively studied (Schiestl and Prakash 1990; Fishman-Lobell et al. 1992; Tomkinson et al. 1993). The process appears to depend primarily on the ScMsh2-ScMsh3 heterodimer, which both recognizes and stabilizes flap-containing heteroduplexes (Paques and Haber 1997; Sugawara et al. 1997; Kearney et al. 2001). Stabilization of the flap-containing heteroduplex by ScMsh2-ScMsh3 increases SSA when the DSB occurs in a region flanked by direct repeats with complementary region <1 kb and nonhomologous tails >30 nucleotides (Paques and Haber 1997; Sugawara et al. 1997). Separation-of-function \textit{msh2} mutations defective in ScMsh2-ScMsh3-dependent nonhomologous end removal, but functional in ScMsh2-ScMsh6 postreplicative MMR suggest that the 3’ protruding-end removal may involve a distinct DNA-binding mode (Goldfarb and Alani 2005). On ScMsh2-ScMsh3 recognition, flaps appear to be processed by ScRad1-ScRad10 (Fig. 1B, right) (Tomkinson et al. 1993). Moreover, the flap-removal function of ScMsh2-ScMsh3 is independent of ScMsh6, ScMlh1, and ScPms1 (Paques and Haber 1997; Sugawara et al. 1997; Kearney et al. 2001).

**PARALLELS BETWEEN YEAST AND MAMMALIAN SYSTEMS**

Mammalian MMR proteins participate in processing HR intermediates and suppressing ho-
meologous recombination (Elliott and Jasin 2001). With exception of HsPMS2, which is the ScPms1 homolog, the core MMR machinery that controls recombination retains both the nomenclature and functionalities of their yeast and bacterial counterparts (Fig. 2; Table 1).

Mammalian cells carrying mutations in XPF or ERCC1 are sensitive to DSB-inducing agents (Ahmad et al. 2008). Thus, it is likely that the mammalian homolog of ScRad1-ScRad10, HsXPF-HsERCC1, plays a similar role in processing heterologous flaps during SDSA and SSA in human cells (Fig. 4; Table 1) (Sargent et al. 2000). In addition, HsERCC1 physically interacts with HsMSH2 and HsRAD52 (Lan et al. 2004). Moreover, the interaction with HsRAD52 stimulates HsXPF-HsERCC1 flap-endonuclease activity on 3′ ends (Fig. 4; Motycka et al. 2004). It is less clear which human helicase homologs play the same role as ScSgs1 (Fig. 4; Table 1). Among the five known human RecQ family helicases, three, HsRECQ1 (Doherty et al. 2005), HsBLM (Pedrazzi et al. 2003), and HsWRN (Saydam et al. 2007), have been shown to interact with the HsMSH2-HsMSH6 heterodimer. HsBLM and HsWRN also interact with the HsMSH2-HsMSH3 heterodimer (Pedrazzi et al. 2003; Saydam et al. 2007). In addition, HsBLM interacts with HsMLH1 and HsExol and, in conjunction with TopIIIα and both RMI1 and RM2, has been shown to dissolve double Holliday junctions (Langland et al. 2001; Pedrazzi et al. 2003; Wu and Hickson 2003; Nimmonkar et al. 2011). HsRECQ1 (Doherty et al. 2005) and HsWRN (Saydam et al. 2007) also interact with HsMLH1, and HsRECQ1 as well as with HsExol (Cui et al. 2004). Furthermore, purified HsRECQ1 forms a subnanomolar affinity complex with HsMSH2-HsMSH6, and HsMSH2-HsMSH6 stimulates HsRECQ1 helicase activity on the 3′-flap structures (Doherty et al. 2005). Finally, both HsMSH2-HsMSH6 and HsMSH2-HsMSH3 stimulate HsWRN helicase activity on forked DNA substrates containing 3′-ssDNA tails (Saydam et al. 2007). Interestingly, the substrate specificity of HsRECQ1, HsBLM, and HsWRN only partially overlaps. Thus, HsRECQ1 readily unwinds immobile Holliday junctions, but does not support the Holliday junction branch migration activity characteristic of HsBLM and HsWRN (Sharma et al. 2005; Popuri et al. 2008).

BIOCHEMICAL RECONSTITUTIONS OF HETERO DupLEX REJEcTION

There have been two approaches toward the biochemical and biophysical analysis of heterodu-
plex rejection. Both approaches reconstitute the initial steps and/or intermediates in homeologous recombination. One approach has been to determine any inhibitory role for the E. coli MMR machinery on DNA strand exchange and D-loop formation catalyzed by EcRecA (Worth et al. 1994; Tham et al. 2013). Two different recombinase assays have been used to observe homologous pairing and strand exchange in vitro: (1) a three-strand system in which RecA/RAD51 catalyzes the exchange of homologous strands between a double-stranded linear DNA and a large circular ssDNA, and (2) RecA/RAD51-catalyzed formation of a D-loop between a relatively short linear ssDNA and a supercoiled circular DNA (Kowalczykowski and Eggleston 1994). The three-strand reaction generally results in extensive branch migration that progresses through partial strand exchange intermediates into the formation of a terminal circular dsDNA product, whereas the D-loop reaction does not result in extensive branch migration (Kowalczykowski et al. 1994).

Purified EcMutS, EcMutL, EcUvrD, EcSSB, and EcRecA were combined in a recent study to examine the effect of MMR proteins on EcRecA-catalyzed three-strand exchange and D-loop formation with and without homeologous recombination substrates (Tham et al. 2013). The investigators show an apparent mismatch-dependent kinetic effect of MMR proteins in the rejection of heteroduplex recombination intermediates. This was evident from the accumulation of three-stranded intermediates and a reduction in circular dsDNA product, whereas the D-loop reaction does not result in extensive branch migration (Kowalczykowski et al. 1994).

An important caveat in these studies is that EcMutS is incapable of forming stable ATP-bound sliding clamps in the experimental ionic conditions (Acharya et al. 2003). Recall that both genetic and biochemical studies have clearly indicated that ATP-bound MSH sliding clamps are essential for both MMR and heteroduplex rejection (Wu and Marinus 1994; Hess et al. 2002; Acharya et al. 2003; Mendillo et al. 2005; Hargreaves et al. 2010). Moreover, persistent mismatch-bound EcMutS has been shown to normally alter MMR functions (Hess et al. 2002). An additional limitation in these studies is that both EcMutS and EcMutL display unusual ssDNA-binding activities at these low experimental ionic strength conditions (Su and Modrich 1986; Gradia et al. 2000; Acharya et al. 2003; Park et al. 2010). It is also well known that ssDNA-binding proteins influence both the DNA interaction(s) and kinetic processes of RecA recombinase activity (Kowalczykowski et al. 1987). Atypical ssDNA-binding activity could also confuse studies that concluded secondary structures in the displaced strand of the three-strand reaction-targeted heteroduplex rejection by EcUvrD (Tham et al. 2013). These issues underscore the complexity of incorporating multiple proteins containing numerous independent and overlapping biochemical activities. Expanding biochemical reconstitutions to include a broader range of experimental conditions compatible with well-known MMR protein functions will likely reveal the nuanced impacts of these DNA repair components in the multifaceted heteroduplex rejection reaction.

In a second approach, the HsMSH2-HsMSH6 heterodimer was shown to bind and form ATP-bound sliding clamps on D-loop intermediates containing a single mismatch in the heteroduplex region (Honda et al. 2014). These studies also showed that HsMSH2-HsMSH6 was capable of recognizing a mismatch within the D-loop heteroduplex when it was bound by the human ssDNA-binding heterotrimer HsRPA and/or the HsRAD51 recombinase (Honda et al. 2014). The results revealed a surprising affinity of MSH proteins for mismatched nucleotides regardless of its surrounding protein and DNA environment. Additional
single molecule analysis established that the lifetime of the HsMSH2-HsMSH6 ATP-bound sliding clamp was almost threefold shorter than on a duplex DNA containing a mismatch (Honda et al. 2014). The reduced stability was traced to interactions between HsMSH2-HsMSH6 with the ssDNA–dsDNA junction on either side of the mismatch-containing heteroduplex DNA within the D-loop. This observation appeared similar to studies with the TaMutS in which reduced sliding clamp lifetime was observed when an ssDNA tail was introduced adjacent to the duplex DNA containing a mismatch (Jeong et al. 2011). The instability associated with ssDNA–dsDNA junctions appears contrary to the observations with the E. coli MMR proteins performed under very low ionic strength conditions (Tham et al. 2013). If correct, the shorter lifetime of the MSH sliding clamps on a D-loop would likely reduce the efficiency of MMR and/or heteroduplex rejection. Such a reduced heteroduplex rejection activity during HR would seem important to preserve a small number of gene conversion events as is observed during Mendelian genetics. Clearly, these initial observations will require confirmation along with more complicated biochemical reconstitution containing the remaining MMR machinery, the recombination initiation machinery, relevant recombination substrates, and a wide range of biochemical conditions.

CONCLUDING REMARKS

Despite the importance of cellular processes that balance homologous and homeologous recombination, our understanding of the heteroduplex rejection mechanism, which is at the heart of this balancing act, remains rudimentary. Most of our current knowledge comes from microbial genetic and cellular studies. There is, as yet, only an elementary understanding of the basic biochemical activities of the proteins and enzymes orchestrating different steps in the heteroduplex rejection, as only a few studies have attempted to tackle this important process, producing tantalizing, but still incomplete conclusions (Worth et al. 1994; Tham et al. 2013; Honda et al. 2014). Complete biochemical reconstitution of the heteroduplex rejection reactions occurring during HR and SSA will be necessary to produce a coherent mechanistic description of the intricate molecular choreography of these processes. More thorough and complete biochemical, biophysical, and single-molecule studies performed under a wider range of experimental conditions would seem necessary for a comprehensive understanding of the MMR-dependent heteroduplex rejection process. Finally, all the eukaryotic players of HR, SSA, and MMR appear to be controlled by post-translational modifications. The effect of these modifications on the heteroduplex rejection and the choice between rejection, processing, or repairing the heteroduplex remains entirely unexplored.

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Mismatch Repair during Homologous and Homeologous Recombination

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