

Dynamic control of strand excision during human DNA mismatch repair

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Mismatch repair (MMR) is activated by evolutionarily conserved MutS homologs (MSH) and MutL homologs (MLH/PMS). MSH recognizes mismatched nucleotides and form extremely stable sliding clamps that may be bound by MLH/PMS to ultimately authorize strand-specific excision starting at a distant 3'- or 5'-DNA scission. The mechanical processes associated with a complete MMR reaction remain enigmatic. The purified human (*Homo sapien* or Hs) 5'-MMR excision reaction requires the HsMSH2–HsMSH6 heterodimer, the 5' → 3' exonuclease HsEXO1, and the single-stranded binding heterotrimer HsRPA. The HsMLH1–HsPMS2 heterodimer substantially influences 5'-MMR excision in cell extracts but is not required in the purified system. Using real-time single-molecule imaging, we show that HsRPA or *Escherichia coli* EcSSB restricts HsEXO1 excision activity on nicked or gapped DNA. HsMSH2–HsMSH6 activates HsEXO1 by overcoming HsRPA/EcSSB inhibition and exploits multiple dynamic sliding clamps to increase tract length. Conversely, HsMLH1–HsPMS2 regulates tract length by controlling the number of excision complexes, providing a link to 5' MMR.

Lynch syndrome/HNPCC | single molecule | MSH2–MSH6 | MLH1–PMS2 | EXOI

Mismatch repair (MMR) is a highly conserved strand-specific excision-resynthesis process that corrects nucleotide misincorporation errors during replication and nucleotide mismatches arising from recombination between heteroallelic parents or physical damage to the DNA (for review see ref. 1). Mutation of core MMR components results in elevated mutation rates and susceptibility to a variety of cancers (2).

MMR has been reconstituted with purified *Escherichia coli*, *Saccharomyces cerevisiae*, and human proteins (3–6). The core MutS homologs (MSH) and MutL homologs (MLH/PMS) components direct a strand-specific excision reaction, whereas re-synthesis appears to be uniquely performed by the replicative polymerase complex (1). In all organisms the excision process is initiated at a single-strand DNA scission (ssDNA/S) that may be located either 3' or 5' and hundreds to thousands of base pairs distant from the mismatch (4, 7). An ssDNA/S positioned on the newly replicated strand ensures accurate correction of replication misincorporation errors (1).

Excision directionality in γ -proteobacteria (*E. coli*) is linked to the choice of 3' or 5' exonucleases that specifically degrade ssDNA generated by the EcUvrD helicase in concert with EcMutS and EcMutL (1). The lack of a helicase distinguishes yeast and human MMR from γ -proteobacteria. Moreover, the eukaryotic 3'- and 5'-excision reactions require different core MMR components and likely occur by different mechanisms (1). For example, the 3'-MMR excision requires the replicative processivity factor PCNA to activate a cryptic MLH/PMS endonuclease activity (8), whereas 5' MMR uses the only known MMR exonuclease EXOI (3, 5, 6). Unlike the *E. coli* ssDNA exonucleases, EXOI will initiate 5' excision from a ssDNA/S in the absence of a helicase (9). Whereas the purified 5'-MMR reaction does not require MLH/PMS or PCNA, complementation

studies with cellular extracts displayed a substantial requirement for MLH/PMS (10, 11).

A number of models have been proposed to account for the transmission of mismatch recognition to the ssDNA/S (12) as well as the roles of MMR components in the ensuing excision process (1, 13). However, the ensemble functions of the MMR components during excision in all organisms remain largely unknown. We have applied several single-molecule imaging techniques to visualize the complete human 5' MMR strand excision process in real time. Our results suggest that dynamic and stochastic processes ultimately control 5' excision, which may at least partially explain the different factor requirements in crude and purified reactions.

Results

Exonuclease Activity by a Single HsEXO1. Single molecule flow stretching (smFS) was used to introduce a regulated laminar flow drag force (F_D) onto a superparamagnetic (SPM) bead tethered to a flow-cell surface with DNA (*SI Appendix, Fig. S1*) (14, 15). At an applied F_D of 2.5 pN (0.0125 mL/min) we found that a 21.8-kb double-stranded DNA (dsDNA) was almost fully extended across the flow-cell surface ($L_{dsDNA} = 7,204$ nm), whereas an equivalent denatured 21.8-kilo-nucleotide (knt) single-stranded DNA (ssDNA) remained coiled ($L_{ssDNA} = 417$ nm; *SI Appendix, Fig. S1 A and B*). This sizeable force–extension difference is consistent with previous work and was used to monitor the production of ssDNA from dsDNA during MMR excision (16).

A 15.3-kb DNA substrate was constructed that containing a G/T mismatch and an ssDNA/S located 8 kb on the 5' side of the mismatch (Fig. 1A; *SI Appendix, Table S1*). The 5' ends of this linear DNA substrate were effectively blocked, one with 5' biotin bound to the surface and the other with by 5' dig-SPM bead, ensuring that 5' excision was nearly always initiated at the

Significance

The ensemble interactions between the human mismatch repair (MMR) components during mismatch-dependent DNA excision repair remain poorly characterized. We have detailed these interactions in real time to reveal new dynamic collaborations between the four evolutionarily conserved components that ultimately result in a complete MMR event. Our observations demonstrate the stochastic nature of an essential genome maintenance system that ultimately results in robust repair events.

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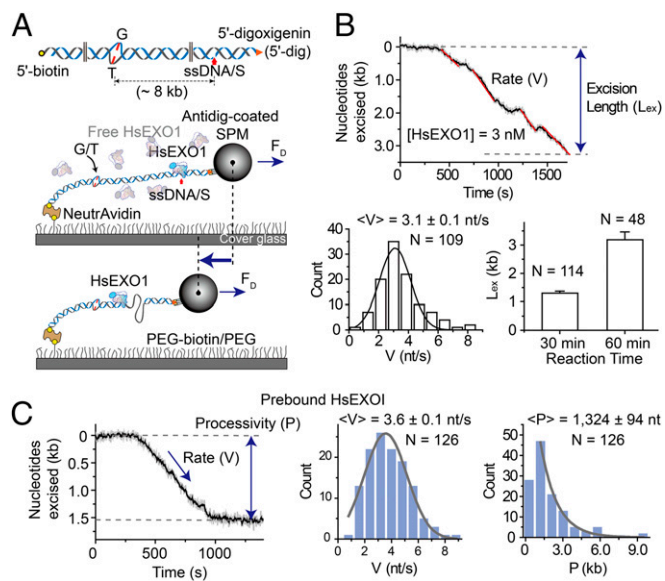


Fig. 1. The rate and processivity of HsEXO1 exonuclease activity. (A, Top) Illustration of the 15.3-kb DNA substrate containing a G/T mismatch. A single-stranded DNA scission (ssDNA/S) that is located 8 kb 5' of the mismatch was introduced using Nb.BbvC1 (New England Biolabs). (Middle and Bottom) Schematic illustration of the smFS system to visualize real-time DNA excision by the HsEXO1 exonuclease. (B, Top) Representative time trajectory of the SPM bead tethered to the 15.3-kb mismatched DNA during a steady-state HsEXO1 excision reaction. The rate (V , red) and excision length (L_{ex}) are shown. (Middle Left) Histogram of binned excision rates that were fit to a Gaussian curve to derive the average rate ($\langle V \rangle$). (Middle Right) The average excision length following a 30- and 60-min reaction. (C, Left) Representative time trajectory of the SPM bead tethered to the 15.3-kb mismatched DNA during a single HsEXO1 excision event (see text). The rate (V) and processivity (P) are shown. (Middle) Histogram of binned excision rates that were fit to a Gaussian curve to derive the average rate ($\langle V \rangle$). (Right) Histogram of binned excision processivity that were fit to a single exponential decay to derive the average processivity ($\langle P \rangle$). All error bars and N indicate SE and the number of molecules, respectively.

internal ssDNA/S. A real-time decrease in the dsDNA extension was observed following the addition of *Homo sapiens* (Hs)EXO1 (Fig. 1B) (14, 15). The rate appeared relatively constant and corresponds to steady-state 5'-exonuclease activity (3.1 ± 0.1 nt/s; Fig. 1B) that ultimately resulted in an average excision of $1,301 \pm 71$ nt after 30 min and $3,188 \pm 277$ nt after 60-min incubations (Fig. 1B; for conversion of nanometers to base pairs, see *SI Appendix*, Fig. S1). Whereas an examination of steady-state excision scans appeared to reveal pauses (Fig. 1B), both the time and spatial resolution of the smFS system were insufficient to accurately establish any connections to HsEXO1 turnover kinetics or excision intermediates.

Previous studies have demonstrated that in presence of Ca^{2+} cations, HsEXO1 remains stably bound to a single-strand scission without catalyzing DNA excision (9, 17). This property was used to examine the excision kinetics of a single HsEXO1 exonuclease event by prebinding the protein in the presence of Ca^{2+} (5 mM) and then initiating the excision reaction by introducing Mg^{2+} (5 mM) into the flow cell (Fig. 1C). Whereas the catalytic excision rate from single HsEXO1 events was not significantly different from the steady-state rate (3.6 ± 0.1 nt/s; Fig. 1C), the average processivity (excision tract length) was $1,324 \pm 94$ nt (Fig. 1C).

HsRPA Inhibits HsEXO1 by Binding to a Nascent Excision Gap. The eukaryotic ssDNA binding heterotrimeric protein RPA is required for MMR (18). To examine the role of HsRPA during 5'-MMR excision, we first needed to establish whether the smFS system could resolve dsDNA from HsRPA-bound ssDNA (ssDNA_{HsRPA}). We found that saturating HsRPA (>10 nM) extended the

21.8-knt ssDNA to 60% of the length of the corresponding dsDNA (4,200 nm; *SI Appendix*, Figs. S1 and S2), which was insufficient to clearly resolve exonuclease activity in real time.

To overcome this technical issue, a static single-molecule total internal reflection fluorescence (smTIRF) analysis was developed with the identical DNA substrate used in the smFS system (Fig. 2A, Top). In this system, the production of ssDNA was visualized by localizing bound HsRPA with Alexa647-conjugated anti-HsRPA70 antibody (Alexa647-HsRPA) and the duplex region was imaged using the dsDNA specific Sytox Orange fluorescent dye. Although not real time, the formation of ssDNA by HsEXO1 (alone) was easily observed as a shortening of the Sytox Orange-stained material (green) associated with intense foci that denote coiled ssDNA excision tracts bound by Alexa647-HsRPA added after the excision reaction was terminated (orange; Fig. 2A, Bottom Left). When HsRPA was added together with HsEXO1, we observed small low-intensity Alexa647 foci on numerous DNA molecules (Fig. 2A, Bottom Middle and *SI Appendix*, Fig. S3, lane 5). We interpret these foci to represent brief ssDNA excision events that by nature are bound by fewer Alexa647-HsRPA. To quantify the frequency of sustained excision events that resulted in long ssDNA tracts with intense Alexa647-HsRPA foci, we first determined the average Alexa647-HsRPA staining intensity of the brief ssDNA excision events produced when HsEXO1 and HsRPA were included together (*SI Appendix*, Fig. S3, lane 5). The number of events that equaled or exceeded one SD above this average intensity (*SI Appendix*, Fig. S3, red

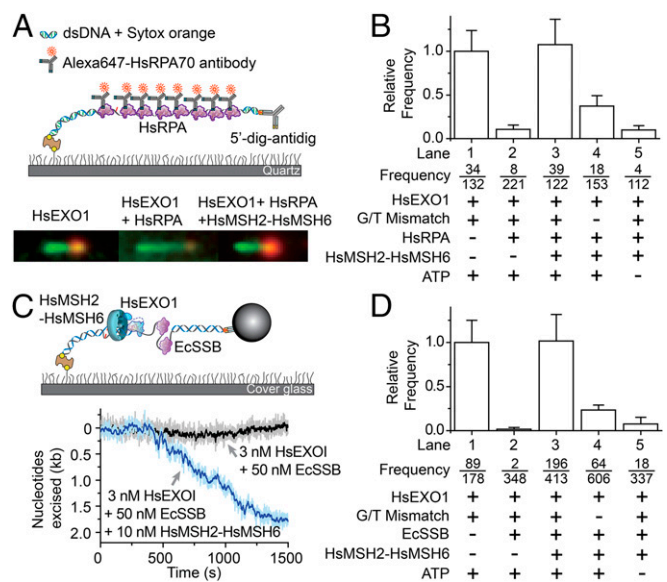


Fig. 2. Reconstitution of 5'-mismatch repair excision on single DNA molecules. (A, Top) Schematic illustration of the static single molecule total internal reflection fluorescence (smTIRF) system used to visualize the 5'-MMR excision process (Methods and *SI Appendix*, *SI Methods*). (Bottom Left) Representative molecule following incubation with HsEXO1 alone. (Bottom Middle) Representative molecule following incubation with HsEXO1 in the presence of HsRPA. (Bottom Right) Representative molecule following incubation with HsEXO1, HsRPA, and HsMSH2-HsMSH6 in the presence of ATP. (B) The relative frequency (\pm SE) of HsEXO1 excision events in the presence of various MMR components determined by smTIRF (3 nM HsEXO1, 10 nM HsMSH2-HsMSH6, 10 nM HsRPA). Frequency is the ratio of the number of corrected excision events (see text and *SI Appendix*, *SI Methods*) to the total number of Sytox Orange-stained dsDNA molecules. (C, Top) Schematic illustration of the real-time smFS system to examine the complete 5'-MMR reaction. (Bottom) Representative time trajectories of SPM beads tethered to the 15.3-kb mismatched DNA. (D) The relative frequency (\pm SE) of HsEXO1 excision events in the presence of various MMR components determined by smFS (Methods). Frequency is the ratio of the number of excision events to the total number of bead-tethered DNA molecules.

dotted line) relative to the total number of Sytox Orange-stained DNA molecules was then determined for each excision reaction condition (Fig. 2*B*, see “frequency”). The frequency of sustained ssDNA excision events under these reaction conditions was then normalized to the frequency of sustained ssDNA excision events when HsEXO1 was added alone (frequency in lane 1) to obtain the relative frequency (Fig. 2*B* and *SI Appendix, SI Methods*). The results clearly indicate that HsRPA dramatically inhibits sustained excision by HsEXO1 (Fig. 2*B*). These observations are similar to bulk studies of human 5'-MMR excision (5), but appear to contrast recombination end-processing studies with the *S. cerevisiae* homologs (19).

There are at least two mechanisms that HsRPA might inhibit HsEXO1 excision activity. HsRPA may interfere with HsEXO1 binding to the strand scission or HsRPA might bind to a short HsEXO1 excision gap ultimately inhibiting the exonuclease. To test these hypotheses, we used single-molecule Förster resonance energy transfer (smFRET) (20, 21). A 73-bp DNA containing a G/T mismatch and a ssDNA/S was attached to a flow-cell surface via a 5'-biotin-NeutrAvidin linkage (*SI Appendix, Fig. S4A*). The ssDNA/S was located between a Cy3 donor and Cy5 acceptor pair (*SI Appendix, Fig. S4A*). HsEXO1 5' → 3' exonuclease activity beginning at the ssDNA/S will form an ssDNA gap that spontaneously coils placing the Cy3 in proximity for FRET with Cy5 (*SI Appendix, Fig. S4 A and B*). We observed no difference in the relative frequency of HsEXO1 excision-induced FRET in the absence of HsRPA or following preincubation with HsRPA (*SI Appendix, Fig. S4C*). These results suggest that HsRPA does not inhibit HsEXO1 entry or exonuclease activity initiated at a 5'-strand scission.

We examined HsEXO1 activity on a substrate DNA containing a mismatch with a recessed 5' end and a 3'-ssDNA tail (30 nt) that would be a model for a nascent excision gap (*SI Appendix, Fig. S4D*). A Cy3 donor was located within the dsDNA 15 bp from the recessed 5' end and a Cy5 acceptor was located on the 3'-ssDNA tail 5 nt from the ssDNA/dsDNA junction. To prevent release of the internal Cy3 by HsEXO1 exonuclease activity, two sequential phosphorothioate linkages were introduced into the phosphate backbone (*SI Appendix, Fig. S4D*). In the absence of HsRPA, we observed ~17% of the DNA molecules transitioned to a steady high FRET state ($E_{\text{low}} = 0.3 \rightarrow E_{\text{high}} = 0.5$; *SI Appendix, Fig. S4 D and E*), indicating HsEXO1-mediated excision of the 15 nt from the recessed 5' end to the phosphorothioate linkages. In contrast, when HsRPA was preassembled onto the ssDNA tail, we observed only 0.5% of the DNA molecules transitioned to high FRET (relative frequency = 0.03; *SI Appendix, Fig. S4 E and F*). These results suggest that HsEXO1 is substantially inhibited when HsRPA binds to the ssDNA gap adjacent to the 5' end that would serve as an entry point for exonuclease activity. Taken together with the small low intensity Alexa647-HsRPA foci in the smTIRF studies, these results suggest that HsEXO1 may begin excision from any 5'-strand scission, but is halted when HsRPA binds to the resulting nascent excision gap.

Reconstitution of the Human MMR 5'-Excision Reaction on Single DNA Molecules. The addition of HsMSH2-HsMSH6 with HsRPA and HsEXO1 resulted in significant shortening of the dsDNA (Fig. 2*A*, *Bottom Right*) and nearly equivalent frequency of excision events as HsEXO1 alone (Fig. 2*A*, *Bottom Right*; Fig. 2*B*, lane 3; and *SI Appendix, Fig. S3*, lane 4). HsMSH2-HsMSH6 stimulation of HsEXO1 excision in the presence of HsRPA was dependent on a mismatch (Fig. 2*B*, lane 4 and *SI Appendix, Fig. S3*, lane 6) and ATP (Fig. 2*B*, lane 5 and *SI Appendix, Fig. S3*, lane 7). The observation that HsMSH2-HsMSH6, HsEXO1, and HsRPA catalyze a mismatch-dependent 5'-excision reaction is similar to previous studies (3, 5, 6) and suggests that we have fully reconstituted 5' MMR on single DNA molecules.

Previous studies have suggested that EcSSB may substitute for HsRPA in the human 5'-MMR excision reaction (5). We could not directly compare EcSSB and HsRPA in the smTIRF system

because a specific EcSSB antibody does not exist. However, we determined that the 21.8-knt ssDNA was extended 2,013 nm in the presence of saturating EcSSB (50 nM), resulting in a conversion factor of 4.2 nt/nm for the dsDNA → ssDNA_{EcSSB} transition that was well within the resolution of smFS (*SI Appendix, Fig. S5*). The significant differences in extension when ssDNA is bound by EcSSB compared with HsRPA likely reflects distinct binding mechanisms; whereas the EcSSB tetramer fully wraps ~65 nt of ssDNA in a “baseball seam” structure (22), HsRPA merely bends ~30 nt of ssDNA (23). Including EcSSB with HsEXO1 completely impeded the contraction of mismatched DNA in the smFS system (Fig. 2*C* and *D*, compare lanes 1 and 2), confirming that EcSSB inhibits HsEXO1 exonuclease activity similar to HsRPA (Fig. 2*B* and *D*) (5). The addition of HsMSH2-HsMSH6 overcomes the EcSSB inhibition of HsEXO1, resulting in the real-time observation of 5' MMR (Fig. 2*C* and *D*, lane 3). The reaction was dependent on a mismatch and ATP (Fig. 2*D*, lanes 4 and 5) underlining the strong functional similarity between EcSSB and HsRPA in 5' MMR (compare Fig. 2*B* and 2*D*).

HsEXO1 Excision Is Dynamic and Controlled by Multiple HsMSH2-HsMSH6 Sliding Clamps. To examine the MMR mechanics, we first determined the concentration-dependent number of HsMSH2-HsMSH6 sliding clamps bound to a single mismatched DNA using mEos3.2-HsMSH2-HsMSH6 that displayed wild-type activity (*SI Appendix, Table S2 and Fig. S6*). The results suggested that at 2 nM, nearly 90% of the DNA molecules contained a single HsMSH2-HsMSH6 (*SI Appendix, Fig. S6*). In contrast, at 60 nM, nearly 70% of the DNA molecules contained two to four HsMSH2-HsMSH6 (*SI Appendix, Fig. S6*). For comparison, the cellular concentration of HsMSH2-HsMSH6 has been estimated to be ~250 nM (24). All of the HsMSH2-HsMSH6 proteins appeared to diffuse randomly on the mismatched DNA consistent with ATP-bound sliding clamps as described previously (20, 25–27).

We examined the interplay between HsMSH2-HsMSH6 and HsEXO1 during 5'-MMR excision (Fig. 3*A*) and found that the excision length did not vary over a range of HsEXO1 concentrations in the presence of 2 nM preassembled HsMSH2-HsMSH6 (Fig. 3*B*). However, in the presence of 10 nM or 60 nM preassembled HsMSH2-HsMSH6 the excision length significantly increased with increasing HsEXO1 (Fig. 3*B*). These results suggest that increasing the number of HsMSH2-HsMSH6 sliding clamps is correlated with increased total excision length. The rate of 5'-MMR excision at a concentration of HsMSH2-HsMSH6 (2 nM) where most events would be due to a single sliding clamp appeared nearly identical to mismatch-independent HsEXO1 excision (3.5 ± 0.2 nt/s; Fig. 3*C*). This result suggests that HsMSH2-HsMSH6 does not influence the fundamental HsEXO1 catalytic process. Interestingly, the excision processivity of the HsMSH2-HsMSH6/HsEXO1 complex appeared nearly twofold shorter than HsEXO1 alone (830 ± 33 nt; Fig. 3*C*).

One possible explanation for the reduced excision processivity is a decreased dynamic lifetime of the ATP-bound HsMSH2-HsMSH6 sliding clamps. We used single molecule protein-induced fluorescence enhancement (smPIFE) to examine the lifetime of HsMSH2-HsMSH6 sliding clamps on mismatched DNA (Fig. 3*D*) (28). A 40-bp Cy3-labeled DNA containing a G/T mismatch was attached at the proximal end to the flow-cell surface via a 5'-biotin-NeutrAvidin linkage, whereas the distal end was blocked with a 5' dig-antidig that traps freely diffusing ATP-bound MSH sliding clamps (Fig. 3*D* and *SI Appendix, Table S3*) (20). Time-averaged diffusion of trapped HsMSH2-HsMSH6 sliding clamps on this relatively short DNA produced a PIFE signal that was used to determine dwell time (τ_{on}). We found that the lifetime of HsMSH2-HsMSH6 on tightly blocked-end mismatched DNA ($\tau_{\text{on}} = 509.1 \pm 14.3$ s; Fig. 3*D*) was at least twofold longer than the time required for an average excision tract ($830 \text{ nt} \div 3.5 \text{ nt/s} = 237 \text{ s}$; Fig. 3*C* and *D*). In contrast, HsMSH2-HsMSH6 sliding clamps rapidly dissociated from either unblocked mismatched DNA with blunt end or unblocked mismatched DNA containing an oligo-dT₁₀ ssDNA tail ($\tau_{\text{on}} = 2.3 \pm 0.1$ s;

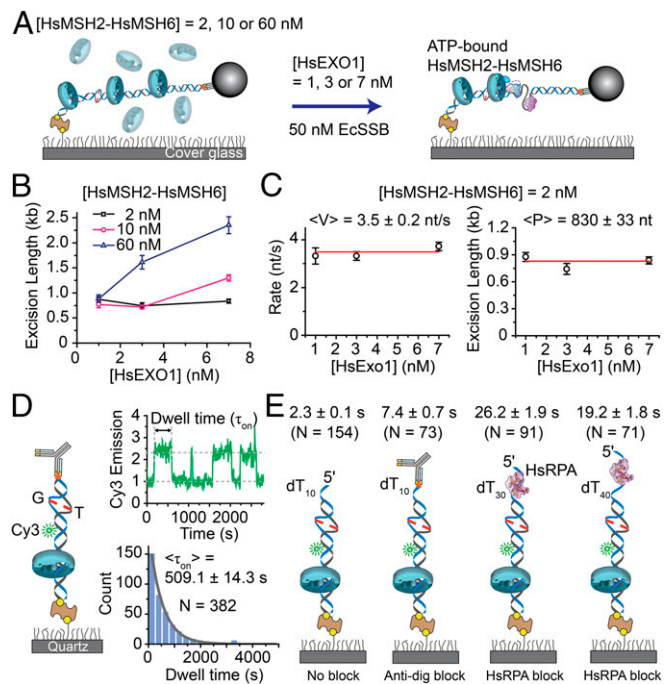


Fig. 3. Multiple HsMSH2-HsMSH6 sliding clamps enhance HsEXO1 excision. (A) Schematic illustration of the single molecule analysis of HsMSH2-HsMSH6 preassembled on mismatched DNA, followed by infusion of HsEXO1 in the presence EcSSB. The HsMSH2-HsMSH6 concentration determines the number of sliding clamps loaded on the mismatched DNA (SI Appendix, Fig. S6). (B) The dependence of excision length (\pm SE) on HsEXO1 concentration in the presence of different HsMSH2-HsMSH6 concentrations. The excision lengths were obtained from the analysis of \sim 32–160 molecules. (C) The dependence of 5'-MMR excision rate ($\langle V \rangle \pm$ SE) and processivity ($\langle P \rangle \pm$ SE) on HsEXO1 concentration in the presence of HsMSH2-HsMSH6 (2 nM) where 88% of the single mismatched DNA molecules contain a single sliding clamp. $n = 32, 34,$ and 126 for 1 nM, 3 nM, and 7 nM HsEXO1. (D, Left) Schematic illustration of the smPIFE 40-bp mismatched DNA substrate (SI Appendix, Table S2). (Right Top) Representative time trace of HsMSH2-HsMSH6 lifetime (τ_{on}) analysis. (Right Bottom) Binned histogram fit with a single exponential decay to determine the lifetime ($\langle \tau_{on} \rangle \pm$ SE) of HsMSH2-HsMSH6 on tightly blocked-end mismatch DNA. (E) Average smPIFE lifetime ($\langle \tau_{on} \rangle$) for HsMSH2-HsMSH6 on various illustrated 40-bp mismatched DNA substrates (SI Appendix, Table S2). The DNA substrates used are (left to right): oligo-dT₁₀ tail; oligo-dT₁₀ tail containing 5' dig-antidig blocked end; oligo-dT₃₀ tail bound with HsRPA (20 nM); and oligo-dT₄₀ tail bound with HsRPA (20 nM). Dwell times are shown above each illustration (\pm SE).

Fig. 3E). The dissociation kinetics was only modestly reduced when the oligo-dT₁₀ ssDNA tail was blocked with dig-antidig ($\tau_{on} = 7.4 \pm 0.7$ s; Fig. 3E). These observations appear similar to *Thermus aquaticus* TaMutS (20) and suggest that HsMSH2-HsMSH6 rapidly dissociates from mismatched DNA when it encounters a ssDNA/dsDNA junction.

We speculated that HsMSH2-HsMSH6 sliding clamps might be destabilized in the absence of a fully protected ssDNA gap. HsRPA displays a high affinity 30 nt ssDNA binding mode (29), although structural analysis suggests that only 25 nt are actually bound (23). We found that mismatched DNA containing an oligo-dT₃₀ ssDNA tail prebound by HsRPA significantly increased the lifetime of HsMSH2-HsMSH6 ($\tau_{on} = 26.2 \pm 1.9$ s) compared with unblocked dT₁₀ or dig-antidig blocked dT₁₀ (Fig. 3E). Increasing the ssDNA tail with oligo-dT₄₀ that is substantially larger than the HsRPA footprint further decreased the lifetime of HsMSH2-HsMSH6 ($\tau_{on} = 19.2 \pm 1.8$ s; Fig. 3E). Taken as a whole, these observations are consistent with the conclusion that HsMSH2-HsMSH6 stabilizes HsEXO1 on mismatched DNA allowing it to catalyze 5' excision. However, any

encounter with an exposed ssDNA gap reduces the lifetime of the HsMSH2-HsMSH6 sliding clamp.

HsMLH1-HsPMS2 Regulates HsMSH2-HsMSH6/HsEXO1 Excision. To examine the relationship between the excision tract length and the mismatch location, we constructed a series of single molecule substrates containing a 31-nt ssDNA gap (ssDNA/G) at variable distances (D) from the mismatch (Fig. 4A, Top). Excision by HsEXO1 (3 nM) alone appeared relatively continuous over 60 min regardless of a mismatch or its position relative to the ssDNA/G (Fig. 4A, Bottom Left; D = 0.5 kb). The addition of EcSSB dramatically reduced the frequency of excision events nearly 70-fold relative to HsEXO1 alone (relative frequency = 1.5%). HsMSH2-HsMSH6 overcomes EcSSB HsEXO1 inhibition and the SPM bead approaches an asymptotic maximum excision length (L_{ex} ; Fig. 4A, Bottom Right, D = 0.5 kb).

Using a box plot, we displayed the L_{ex} median (constriction between quartiles), the mean (+), 1.5 times the interquartile range from the first and third quartiles (filled colors) and all L_{ex} observations as whiskers. Importantly, the L_{ex} was dependent on HsMSH2-HsMSH6 concentration but not the distance between the ssDNA/G and the mismatch (Fig. 4B). For example, slightly more than 50% with 10 nM HsMSH2-HsMSH6, and nearly all of excision tracts with 60 nM HsMSH2-HsMSH6, terminated past the mismatch on the DNA substrate containing D = 0.5 kb (Fig. 4B). A similar trend was observed when the ssDNA/G was 1.0 or 2.0 kb from the mismatch, except that in most cases, the excision tract terminated before the mismatch (Fig. 4B, D = 1.0 and 2.0 kb). Additional HsEXO1 (7 nM) resulted in more than 50% of the excision tracts terminating past the mismatch with 60 nM HsMSH2-HsMSH6, which could be improved to 90% with

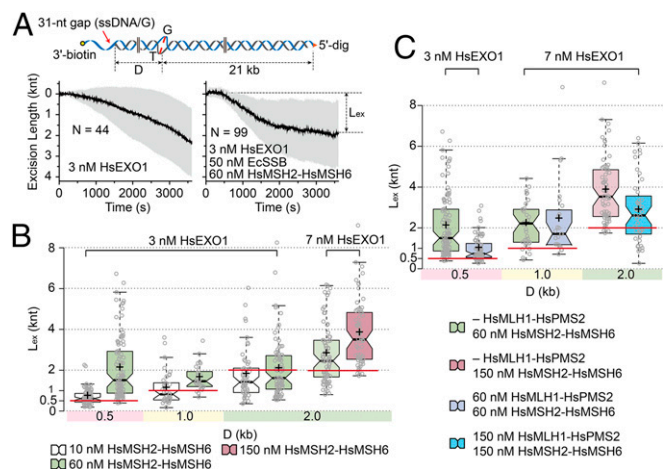


Fig. 4. The 5'-MMR excision is enhanced by HsMSH2-HsMSH6 and down-regulated by HsMLH1-HsPMS2. (A, Top) Schematic illustration of 21 kb mismatched DNA substrate containing a DNA extension that places a ssDNA 5' gap at a distance (D) of 0.5, 1.0, and 2.0 kb from the mismatch. (Bottom Left) Time trajectories of the average (black) and SD (gray) during 5' excision by HsEXO1 (3 nM) alone on the 21 kb mismatched DNA with D = 0.5 kb. (Bottom Right) Time trajectories of the average (black) and SD (gray) during 5' excision on the 21 kb mismatched DNA with D = 0.5 kb. (C) Box plots of the excision length (L_{ex} ; knt) vs. D (kb) in the presence of 50 nM EcSSB (from left $n = 78, 96, 38, 28, 49, 81, 74,$ and 64). Concentrations of HsEXO1 and box plot colors indicating HsMSH2-HsMSH6 concentration are shown above. (C) Box plots of the excision length (L_{ex} ; knt) vs. D (kb) in the presence of 50 nM EcSSB (from left $n = 96, 67, 34, 23, 64,$ and 43). Concentrations of HsEXO1 are indicated above plots. Box plot colors indicating HsMSH2-HsMSH6 and HsMLH1-HsPMS2 concentration are shown below. The cross mark and the constriction line indicates the mean and the median, respectively. Solid extensions above and below the median indicate upper and lower quartiles, respectively. Red lines indicate the distance between the mismatch and ssDNA/G.

150 nM HsMSH2–HsMSH6. Under nearly all conditions, the excision length could exceed 2–3 times the distance past the mismatch as the distance between the 5′ ssDNA/G and the mismatch.

Introducing HsMLH1–HsPMS2 into the HsMSH2–HsMSH6/HsEXO1 5′-MMR excision reaction reduced the median length of the excision tracts under all conditions examined (Fig. 4C). In most cases, the excision tract terminated just past the mismatch (Fig. 4C). As a control, we found that HsMLH1–HsPMS2 did not activate HsEXO1 in the absence of HsMSH2–HsMSH6 nor does a threefold excess of HsMLH1–HsPMS2 decrease the HsEXO1 excision tract length activated by a single preassembled HsMSH2–HsMSH6 sliding clamp (*SI Appendix, Fig. S7* and see Fig. 3C and *SI Appendix, Fig. S6B*). Another mechanism that is consistent with previous studies suggests that MLH/PMS might alter the association of MSH with the mismatched DNA (26, 30). We found that the number of mEos3.2–HsMSH2–HsMSH6 sliding clamps was significantly reduced in the presence of HsMLH1–HsPMS2 (*SI Appendix, Fig. S8*). Combined with studies of the yeast homologs that suggest that the stability of the ScMsh2–ScMsh6/ScMlh1–ScPms1 complex is not significantly different from ScMsh2–ScMsh6 sliding clamps alone (26), these observations are consistent with the hypothesis that HsMLH1–HsPMS2 controls runaway HsMSH2–HsMSH6/HsEXO1 excision by stochastically regulating the number of HsMSH2–HsMSH6 sliding clamps associated with the mismatched DNA.

Discussion

We have examined the human 5′-MMR excision process in real time on single DNA molecules containing a G/T mismatch and a single-strand scission or gap. To our knowledge, this is the first reconstitution of a multicomponent DNA repair reaction on single DNA molecules. Several imaging techniques were applied to visualize the mechanics. Our observations expand previous bulk studies with yeast and human MMR proteins (3, 5, 6) by resolving the complex biophysical assemblies between HsRPA, HsEXO1, HsMSH2–HsMSH6, and HsMLH1–HsPMS2.

HsEXO1 appeared active on any DNA substrate containing a 5′ single-stranded DNA scission or gap (5, 9, 17). HsRPA dramatically inhibited HsEXO1 exonuclease activity (Fig. 2A and B) (5). However, inhibition only occurred after a ssDNA gap was formed that could be bound by HsRPA (Fig. 2A, *Bottom Middle* and *SI Appendix, Fig. S4*). These results suggest a previously unknown very early role for HsRPA in regulating unfettered HsEXO1 5′ excision. In the presence of a mismatch and ATP, HsMSH2–HsMSH6 overcomes the HsRPA inhibition of HsEXO1 (3, 5, 6). A surprising result was that EcSSB could fully substitute for HsRPA in all HsEXO1-mediated excision reactions (Figs. 2C and D, 3, and 4). This observation strongly suggests that there are no explicit interactions between HsMSH2–HsMSH6 and HsEXO1 with either HsRPA or EcSSB, and that the singular role of HsRPA during 5′ MMR is as an ssDNA binding protein. These studies appear to clearly explain why no MMR-deficient RPA mutations were identified in yeast mutator genetic screens (31).

The rate of excision by HsEXO1 alone was virtually identical to the rate of excision by the HsMSH2–HsMSH6/HsEXO1 complex (compare Fig. 1C and Fig. 3C). An attractive hypothesis is that ATP-bound MSH sliding clamps (20, 25, 26) provide a freely diffusible platform that stabilizes EXO1 on the 5′ end of an ssDNA scission or gap via well-known protein–protein interaction domains (32). We envision a model where MSH sliding clamps function similarly to PCNA sliding clamps that tether the polymerase machinery to the replication fork (33). By counting the number of HsMSH2–HsMSH6 sliding clamps loaded onto a mismatched DNA molecule, we determined that a single HsMSH2–HsMSH6/HsEXO1 complex processively degrades 830 nt during a single 5′-MMR excision event (Fig. 3C). This implies that the lifetime of the HsMSH2–HsMSH6/HsEXO1 complex is ~2 min shorter than HsEXO1 alone. It is likely that this reduced lifetime is linked to transient exposure of the ssDNA gap during 5′ excision, which the PIFE studies showed decreased the lifetime of HsMSH2–HsMSH6 sliding clamps 10- to 100-fold depending on the extent of

HsRPA occlusion (Fig. 3D and E). We also note that a significantly decreased HsMSH2–HsMSH6 lifetime on a mismatched DNA containing an exposed ssDNA gap is an alternate explanation for the dramatically decreased MMR efficiency in the absence of HsRPA observed by Genschel et al. (5).

Whereas there is strong biochemical evidence that MSH proteins form multiple extremely stable sliding clamps on mismatch DNA in the presence of ATP (20, 25, 27), one could argue that these are anomalous structures that develop in the absence of additional MMR components. However, we found a near direct correlation between the HsEXO1-dependent 5′-MMR excision tract length (Figs. 3B and 4B) and numbers of HsMSH2–HsMSH6 sliding clamps observed on the mismatched DNA (*SI Appendix, Fig. S6*). Moreover, at any fixed concentration of HsEXO1, the average excision tract length appeared to be uniquely dependent on HsMSH2–HsMSH6 concentration and was largely independent of the ssDNA/G location relative to the mismatch (Fig. 4B). Together, these observations suggest a concentration-dependent loading of multiple HsMSH2–HsMSH6 sliding clamps that dynamically activate HsEXO1 5′ excision, and argue against different configurations, complexes, or functions of HsMSH2–HsMSH6 during its interaction with HsEXO1.

Previous studies have demonstrated that HsMLH1–HsPMS2 is not required for the 5′-MMR excision reaction (3, 5), but when added, appeared to alter excision termination past the mismatch (6). We found that the HsMSH2–HsMSH6/HsEXO1 excision endpoints were shorter in the presence of equivalent HsMLH1–HsPMS2 physiological stoichiometries (Fig. 4C). These observations are consistent with the conclusion that HsMLH1–HsPMS2 inhibits runaway excision by the HsMSH2–HsMSH6/HsEXO1 complex. Interestingly, when the mismatch and the ssDNA/G were separated by 2.0 kb, the addition of HsMLH1–HsPMS2 resulted in a significant fraction (40%) of excision tracts that did not reach the mismatch (Fig. 4B, D = 2.0 kb). This was despite relatively high concentrations of HsMSH2–HsMSH6 (150 nM) that should have loaded sufficient sliding clamps capable of activating HsEXO1 to easily cover that distance. These results suggest that HsMLH1–HsPMS2 is unlikely to detect the mismatch as part of its excision-regulation role. We also found that the excision tract length of a single HsMSH2–HsMSH6/HsEXO1 event was not reduced by HsMLH1–HsPMS2 (*SI Appendix, Fig. S7*). Instead, we showed that HsMLH1–HsPMS2 appears to reduce the number of dynamic excision events by stochastically modulating the number of HsMSH2–HsMSH6 sliding clamps capable of activating HsEXO1 5′ excision (*SI Appendix, Fig. S8*). Our data cannot rule out the possibilities that HsMLH1–HsPMS2 might regulate HsMSH2–HsMSH6 loading at the mismatch or titrate an excision component such as HsEXO1 independent of the mismatch (32).

Taken together, these studies strongly support a dynamic model for MMR (1, 30) and effectively eliminate static or single-complex models (13, 34). A surprising observation was that even in the absence of a mismatch, HsEXO1 efficiently converts a 5′-strand scission into a gap bound by HsRPA before its exonuclease activity is fully inhibited (Fig. 2A, *Bottom Middle* and *SI Appendix, Fig. S4*). There is compelling evidence that a combination of RNaseH, the flap endonuclease FEN1, and Polδ polymerase strand-displacement activity ultimately remove the RNA primer from the 5′ end of an Okazaki fragment (35). EXO1 has been proposed to play at least a minor role in Okazaki fragment maturation because the rad27Δ(ScFen1) exo1Δ double mutant is lethal in yeast (36). An appealing model would envision HsRPA associated with a transient gap exposed during the removal of the RNA primer that functionally inhibits any potential HsEXO1 excision (*SI Appendix, Fig. S9A*).

The molecular switch/sliding clamp model remains the most consistent mechanism for MMR excision (*SI Appendix, Fig. S9*) (1, 30). In the presence of a mismatch, multiple ATP-bound HsMSH2–HsMSH6 sliding clamps may be loaded onto the DNA that may then interact with HsMLH1–HsPMS2 (*SI Appendix, Fig. S9B*). The HsMSH2–HsMSH6 sliding clamp that is closest to the 5′ end initiates 5′ → 3′ excision by stabilizing HsEXO1

(SI Appendix, Fig. S9B). A single HsMSH2–HsMSH6/HsEXO1 complex generates an ~800-nt excision tract, which is successively bound by HsRPA before the complex spontaneously dissociates (SI Appendix, Fig. S9C). The next closest HsMSH2–HsMSH6 sliding clamp then stabilizes another HsEXO1 at the newly located 5' end, reinitiating excision (SI Appendix, Fig. S9C). This animated process is proposed to be iterative until the mismatch is released and no additional HsMSH2–HsMSH6 sliding clamps may be loaded onto the DNA (SI Appendix, Fig. S9D). The regulatory role of HsMLH1–HsPMS2 in 5'-MMR excision is likely to be significantly different from its catalytic role on 3'-MMR excision where HsPCNA may activate its intrinsic endonuclease (8). However, the control of HsEXO1 by HsMSH2–HsMSH6 and HsMLH1–HsPMS2 appears to provide a dynamic yin and yang for efficient 5'-MMR excision.

Methods

See SI Appendix, SI Methods for comprehensive methods.

Preparation of Proteins and DNA. HsMSH2–HsMSH6 and HsRPA were purified as previously described (37, 38). HsEXO1 and HsMLH1–HsPMS2 were purified with modifications to published methods (SI Appendix, SI Methods) (9, 11).

Single-Molecule Flow Stretching, Total Internal Reflection Fluorescence, and Protein-Induced Fluorescence Enhancement Microscopy. A 15.3-kb DNA substrate containing a G/T mismatch was constructed as previously described (25). The construction of DNA substrates containing an ssDNA/G at various distances from the G/T mismatched is described in the SI Appendix, SI Methods.

For smF5 microscopy, a custom flow chamber was constructed with mismatched DNA attached to the surface and the SPM bead as previously

described (39). The SPM bead was imaged with a 10× objective (N.A. = 0.40, Olympus) containing ~300-tethered DNA molecules in a field of view, recorded with a high-resolution CCD (RETIGA 2000R, Qimaging) using MetaVue (Molecular Devices) imaging software at 1-s time resolution, the bead position determined with high accuracy (~20 nm) by DiaTrack 3.03 (40), and the data were analyzed using OriginPro8 (OriginLab) and Matlab 2013b (Mathworks).

The smTIRF microscopy analysis was performed as previously described (20, 25). Alexa647-labeled anti-HsRPA70 antibody (Abcam) was infused into the flow chamber following HsRPA (10 nM) incubation in reactions performed in the absence of HsRPA or directly after a 60-min reaction that contained HsRPA. Alexa647 emission was imaged after 10 min and then colocalized with dsDNA following Sytox Orange (Thermo Fisher Scientific) staining.

For smPIFE microscopy, 40 bp of G/T mismatched DNA molecules with various ssDNA tails (0, 10, 30, and 40 oligo-dT) were prepared by annealing paired DNAs (SI Appendix, Table S2 and SI Methods). All of the DNA substrates were labeled with Cy3 (Monofunctional NHS-ester, GE Healthcare), and attached to the flow cell surface via 5'-biotin–NeutrAvidin as previously described (20). DNAs with an oligo-dT₀ or oligo-dT₁₀ tail contained a 5' dig. Anti-dig antibody (50 nM; Roche) or HsRPA (20 nM) was introduced into the flow chamber followed by HsMSH2–HsMSH6. Emission of Cy3 was imaged by smTIRF as previously described (20, 25). The time resolution was adjusted ranging from 0.2 s to 2 s according to the lifetime of HsMSH2–HsMSH6 using a time-lapse method (SI Appendix, SI Methods). PIFE data were analyzed using IDL 6.4 (EXELIS VIS) and Matlab 2013b (Mathworks).

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