Human MutLγ, the MLH1–MLH3 heterodimer, is an endonuclease that promotes DNA expansion

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MutL proteins are ubiquitous and play important roles in DNA metabolism. MutLγ (MLH1–MLH3 heterodimer) is a poorly understood member of the eukaryotic family of MutL proteins that has been implicated in triplet repeat expansion, but its action in this deleterious process has remained unknown. In humans, triplet repeat expansion is the molecular basis for ~40 neurological disorders. In addition to MutLγ, triplet repeat expansion involves the mismatch recognition factor MutSβ (MSH2–MSH3 heterodimer). We show here that human MutLγ is an endonuclease that nicks DNA. Strikingly, incision of covalently closed, relaxed loop-containing DNA by human MutLγ is promoted by MutSβ and targeted to the strand opposite the loop. The resulting strand breaks license downstream events that lead to a DNA expansion event in human cell extracts. Our data imply that the mammalian MutLγ is a unique endonuclease that can initiate triplet repeat DNA expansions.

Significance

Triplet repeat expansion causes multiple neurological disorders, but the mechanisms of triplet repeat expansion are not well understood. Growing evidence indicates that DNA loops, MutSβ (MSH2–MSH3 heterodimer), and MutLγ (MLH1–MLH3 heterodimer) play important roles in triplet repeat expansion. We demonstrate here that human MutLγ is an endonuclease that nicks DNA in a MutSβ- and loop-dependent manner. Inclusion of loop-containing DNA by MutLγ endonuclease initiates events that lead to DNA expansion. Surprisingly, cleavage of loop-containing DNAs by MutSβ-dependent endonuclease activity of MutLγ is strongly biased to the strand that lacks the loop. These findings document an endonuclease activity and mechanism that may be important for triplet repeat expansion.

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of replication. Genome-wide association studies with human patients have suggested that MMR system genes MSH3, MLH1, MLH3, and PMS2 function as genetic modifiers of the age of onset of symptoms of Huntington’s disease and spinocerebellar ataxias (18, 22, 23, 44–46). Moreover, genetic analyses in mouse models and cultured human cells have demonstrated that the MMR system factors MutSβ, MutLα, and MutLγ are involved in triplet repeat expansion (16, 19, 21, 29, 37, 47–49). Further research has shown that MutSβ promotes triplet repeat expansion as a DNA loop recognition factor and MutLα as an endonuclease (17). However, the action of MutLγ in expansion remains unknown.

The MutLγ homolog MutLα contains the conserved DQHA(X)2E(X)4E motif that is part of its endonuclease active site (11, 12, 50). This motif and three other MutLα endonuclease motifs are present in the MLH3 subunits of yeast and mammalian MutLγ proteins (11, 51). Consistent with the presence of the endonuclease motifs in its MLH3 subunit, yeast MutLγ has been shown to possess an endonuclease activity that nicks DNA (52–54). However, it remains unknown how yeast MutLγ endonuclease activity contributes to the action of this protein in DNA metabolism. Furthermore, it has not been known whether mammalian MutLγ proteins have endonuclease activity. Here we show that human MutLγ has a unique MutSβ-dependent endonuclease activity that incises loop-containing DNAs in the strand that does not have the loop. The resulting nick is used by downstream activities to promote a DNA expansion event.

Results

Human MutLγ Is an Endonuclease. We began this study to advance our understanding of the action of MutLγ in mammalian cells. Because the DQHA(X)2E(X)4E endonuclease motif is preserved in human and several other mammalian MLH3 proteins (11, 51) (SI Appendix, Fig. S1A), we decided to investigate whether human MutLγ had an endonuclease activity. We first purified human MutLγ and its mutant derivative, MutLγ-D1223N, which were produced in insect S9 cells (SI Appendix, Fig. S1 B and C). The MutLγ-D1223N variant contains a D-to-N change in the DQHA(X)2E(X)4E endonuclease motif (SI Appendix, Fig. S1A). The corresponding substitution inactivates the endonuclease function of human MutLα (11). To facilitate purification, a FLAG tag was placed at the N terminus of the MLH3 and MLH3-D1223N subunits. The purity of the proteins obtained at the final purification step was ≥95%. During purification the MutLγ-D1223N variant behaved like wild-type protein, which suggested that the D1223N amino acid substitution did not cause a significant change in protein structure.

We next examined whether the purified human MutLγ possessed an endonuclease activity. Because we previously observed that human and yeast MutLö endonucleases can be gratuitously activated on supercoiled DNA under low-salt conditions in the presence of Mn2+ (11, 12), we tested whether human MutLγ displayed a similar endonuclease activity. The data demonstrated that the purified human MutLγ had a Mn2+-dependent endonuclease activity that nicked supercoiled homoduplex DNA (Fig. 1 A and B). Control experiments revealed that MutLγ-D1223N is defective in supporting this endonuclease reaction (Fig. 1 D). We then compared the levels of the Mn2+-dependent endonuclease activities of human MutLγ and MutLα. The results showed that under the tested conditions, the specific Mn2+-dependent endonuclease activity of human MutLγ was ~3 times higher than that of human MutLα (Fig. 1 C and SI Appendix, Fig. S2). Together these findings demonstrated that human MutLγ is a metal-dependent endonuclease.
The MLH1 and MLH3 subunits of human MutLγ contain conserved motifs that are required for ATP binding and hydrolysis by the members of the GHKL family (55, 56). We analyzed whether the preparations of human MutLγ and MutLγ-D1223N were able to hydrolyze ATP. The data showed that the human MutLγ and MutLγ-D1223N preparations hydrolyzed ATP at similar rates, ~0.27 mol of ATP hydrolyzed per min per mol of the MutL protein at an initial ATP concentration of 0.5 mM (SI Appendix, Fig. S3). This finding suggested that human MutLγ and MutLγ-D1223N had ATPase activities.

Next, we examined the impact of ATP on the endonuclease activity of human MutLγ. We established that the endonuclease activity of human MutLγ was stimulated by 0.25 to 1 mM ATP (Fig. 1D). This implied that ATP is a cofactor for human MutLγ endonuclease. However, higher ATP concentrations were not stimulatory, likely due to chelation of Mn$^{2+}$ by ATP, rendering the cation unavailable for activation of the MutLγ endonuclease. We also established that dATP stimulated the Mn$^{2+}$-dependent endonuclease activity of human MutLγ, but CTP, UTP, and GTP did not (SI Appendix, Fig. S4). This suggests that the enzyme can utilize dATP as a cofactor.

In addition to Mn$^{2+}$, Mg$^{2+}$ and Co$^{2+}$ activate yeast MutLγ endonuclease to nick supercoiled DNA (53). We tested whether human MutLγ endonuclease activity was promoted by Mg$^{2+}$, Co$^{2+}$, Cu$^{2+}$, Ni$^{2+}$, or Zn$^{2+}$. The results showed that human MutLγ endonuclease was activated by Mg$^{2+}$ but not by Co$^{2+}$, Ca$^{2+}$, Ni$^{2+}$, or Zn$^{2+}$ (SI Appendix, Fig. S5). The observation that Co$^{2+}$, a cation that activates yeast MutLγ endonuclease (53), did not promote human MutLγ endonuclease activity suggests that there may be a significant difference between the active sites of human and yeast MutLγ endonuclease.

MutLγ and the mismatch recognition factor MutSβ have been linked to triplet repeat expansion (19–21, 30, 37, 49). We studied whether human MutLγ and MutSβ interacted with each other. We established that the Mn$^{2+}$-dependent endonuclease activity of human MutLγ on supercoiled homoduplex DNA was strongly promoted by human MutSβ (Fig. 1C). These findings demonstrated that human MutSβ directly interacted with human MutLγ endonuclease.

**Human MutLγ Has a MutSβ-Dependent Endonuclease Activity That Cleaves a Loop-Containing DNA in the Strand That Lacks the Loop.** We next investigated whether in the presence of ATP and Mg$^{2+}$ human MutLγ and MutSβ formed a two-protein system that incised a relaxed covalently closed DNA (ccDNA) containing a 3-nt (5′-AGC) loop in the top strand (Fig. 2). These experiments were done at physiological salt concentration to suppress nonspecific incision, and products were visualized by Southern hybridization after BanI cleavage and separation on a denaturing agarose gel. The data showed that the two-protein system incised the loop-lacking strand, but not the loop-containing strand, of the heteroduplex ccDNA (Fig. 2A and B, lane 3). Inspection of the cleavage pattern of the heteroduplex ccDNA (Fig. 2C, lane 3) indicated that MutLγ endonuclease cleaved the loop-lacking strand at multiple sites. The omission of MutLγ or MutSβ or replacement of MutLγ with MutLγ-D1223N abolished the

![](image1)

**Fig. 2.** Human MutLγ has a MutSβ-dependent endonuclease activity that incises a 3-nt loop-containing DNA in the strand that does not carry the loop. The defined reaction mixtures contained Mg$^{2+}$ and the indicated proteins and DNAs. Reactions were carried out and analyzed as detailed in Materials and Methods. The products of the indicated defined reactions were cleaved with BanI, separated in 1.4% denaturing agarose gels, and visualized by Southern hybridizations with 32P-labeled oligonucleotides 5′-GACAGTACCAATCATTAGCTG-3′ (A) and 5′-CAGTGTTAAACATTGG-TAATGTG-3′ (B). (A and B) Incision of the loop-lacking (A) and 3-nt loop-containing (B) strands of the heteroduplex DNA in the indicated reactions. The arrows indicate locations of DNA products that were formed by cleavage of the ccDNA at or near the loop site. The diagrams on the left illustrate the 3-nt loop-containing ccDNA substrate and show the relative positions of the 3-nt loop, the BanI site, and the 32P-labeled probes. (C) Quantification of the incision of the loop-lacking and 3-nt loop-containing strands of the heteroduplex DNA in the indicated reactions. The incision values were determined from phosphorimager data and are presented as averages ±1 SD (n ≥ 3).
incision of the loop-lacking strand of the heteroduplex ccDNA (Fig. 2, lanes 2 and 4, and Fig. 2C). Importantly, at the physiological salt concentration used, the two-protein system did not cause significant cleavage of either strand of the control homoduplex ccDNA (Fig. 2A and B, lane 6, and Fig. 2C). These experiments revealed that human MutLα has a MutSβ-dependent endonuclease activity that incises a 3-nt loop-containing ccDNA at multiple sites that are located on the loop-lacking strand.

**Human MutLα Endonuclease Promotes DNA Expansions.** Small loops are formed in triplet repeat DNA in vivo and in vitro and are likely to be the structures that initiate triplet repeat expansion (17, 30, 41). We determined whether human MutLα endonuclease and MutSβ promoted DNA expansion in the 3-nt loop-containing ccDNA in a reconstituted cell extract system that included ATP, the four dNTPs, and Mg2+ (Fig. 3). In these experiments, we utilized a cell-free extract that was prepared from human MLH1+/−/− MSH3+/−/− H6 cells (8, 11, 57). If the loop-containing bottom strand of this relaxed ccDNA is incised and then subject to repair DNA synthesis using the top strand as a template, a 3-bp expansion takes place (Fig. 3A). However, if the top strand is incised and then repaired using the bottom strand as a template, the 3-nt loop is removed. The two events, the 3-bp DNA expansion and 3-nt loop removal, can be differentiated from each other by diagnostic cleavages of the reaction products with the restriction endonucleases BmtI and HindIII (Fig. 3A). Supplementation of the MLH1+/−/− MSH3+/−/− H6 cell extract-containing reaction mixture with purified human MutLα and MutSβ triggered repair to a 3-bp DNA expansion (Fig. 3B and SI Appendix, Fig. S6). Although the fraction of DNA subject to expansion in these experiments was small (~3%), it is similar to the yield of expanded DNA in a MutLα endonuclease-dependent reaction in a human nuclear extract system (17), as well as the extent of cyclobutane thymine dimer repair that occurs in a CHO cell extract system (58).

Importantly, we determined that the addition of the endonuclease mutant MutLα-D1223N and MutSβ to the H6 extract-containing mixture did not affect the level of the DNA expansions. Strikingly, the addition of MutLα and MutSβ to the H6 extract-containing reaction mixture caused no significant increase in the level of 3-nt loop removal. In contrast, the addition of MutLα and MutSβ to the H6 extract-containing reaction mixture caused significant increase in the level of 3-nt loop removal. A similar result was obtained in a recent study of human MutLα and MutSβ (17). The above findings demonstrated that MutLα endonuclease and MutSβ act in the same pathway that specifically promotes DNA expansions.

**Cleavage of (CTG)γ/(CAG)β and (CTG)β/(CAG)γ Heteroduplex DNAs by Activated Human MutLα Endonuclease.** Expansion of CTG/CAG repeats in the human DMPK gene is an essential step in the process that causes myotonic dystrophy (59–61). In the next series of experiments, we studied whether the two-protein system cleaved a relaxed (CTG)α/(CAG)β heteroduplex ccDNA in which a 6-nt loop was within the sequence context of the human DMPK gene (Fig. 4) (17). (Due to the surrounding sequence the 6-nt loop sequence in a (CTG)α/(CAG)β heteroduplex molecule may be CTGCTG, GCTGCT, or TGCTGC.) The data demonstrated that in the presence of Mg2+ the two-protein system cleaved the loop-lacking strand of the (CTG)β/(CAG)α heteroduplex ccDNA in a reaction that required the presence of both proteins and MutLα endonuclease function (Fig. 4A, lanes 2 to 5, and Fig. 4C).

Strikingly, the activated MutLα preferred to incise the loop-lacking strand of the heteroduplex DNA at two regions located to either side of the lesion, each of which was ~300 bp away from the loop (Fig. 4A, lane 3). We also found that the presence of PCNA and RFC did not increase the level of incision of the loop-lacking strand of the heteroduplex DNA. Consistent with the results shown in Fig. 2, we established that neither the control (CTG)β/(CAG)α homoduplex DNA nor the loop-containing strand of the (CTG)β/(CAG)α heteroduplex DNA was incised by the MutSβ-dependent endonuclease activity of MutLα (Fig. 4A and B, lanes 8 and 9; Fig. 4B, lanes 3 and 6; and Fig. 4C). The experiments summarized in Fig. 4 utilized a relaxed ccDNA that contained a 6-nt loop in the top strand. We also studied how human MutLα endonuclease acted on a similar relaxed ccDNA, a (CTG)γ/(CAG)β heteroduplex, that carried a 6-nt loop in the bottom strand. The results showed that the endonuclease activity of human MutLα incised the loop-lacking strand of (CTG)γ/(CAG)β heteroduplex ccDNA in a MutSβ- (SI Appendix, Fig. S7) and ATP-dependent reaction (SI Appendix, Fig. S8). Incision of the loop-lacking strand often took place in a region that encompassed the 6-nt loop but also occurred at several more distant minor sites (SI Appendix, Fig. S7B, lane 3, and SI Appendix, Fig. S8, lane 2). Collectively, the results of the above experiments (Figs. 2 and 4 and SI Appendix, Figs. S7 and S8) demonstrated that MutLα has a unique MutSβ-dependent endonuclease activity that incises loop-containing ccDNAs in the strand that lacks the loop.

In our previous analysis of the defined two-protein system (Figs. 2 and 4 and SI Appendix, Figs. S7 and S8), we carried out the endonuclease reactions in the presence of Mg2+. We have also found that MutSβ- and loop-dependent endonuclease activity of human MutLα on the (CTG)γ/(CAG)β heteroduplex ccDNA is also evident in the presence of Mn2+ provided that the salt...
concentration is elevated to suppress nonspecific MutLγ nuclease activity. As observed in the presence of Mg\(^{2+}\), the MutSβ-dependent endonuclease activity of human MutLγ incised the loop-lacking strand of the loop-containing ccDNA in the presence of 0.05 to 1 mM of Mn\(^{2+}\), but it was silent in the presence of 0.01 to 0.03 mM of Mn\(^{2+}\) (SI Appendix, Fig. S9), concentrations similar to those present in the mammalian cell (62).

**Discussion**

Despite their significant impact on DNA metabolism (19–21, 24), mammalian MutLγ proteins and the mechanisms of their action have been poorly understood. Since the discovery of the DOHA(X)\(_2\)E(X)\(_4\)E endonuclease motif in PMS2 and its homologs (11, 12), a key question has been whether a mammalian MutLγ possesses an endonuclease activity. Here we have shown that human MutLγ is a metal-dependent endonuclease. Our study has discovered that human MutLγ displays a distinct MutSβ-dependent endonuclease activity that incises loop-containing DNAs in the strand that does not carry the loop.

A recent report was the first to implicate endonuclease activity of a MutL protein in triplet repeat expansion (17). That study showed that human MutLα endonuclease promotes triplet repeat expansion by incising ccDNA in a loop-dependent manner. The incision of triplet repeat extrusion-containing ccDNA by human MutLα occurs in either strand and requires the presence of MutSβ, PCNA, and RFC (17). The current work has provided evidence for the involvement of endonuclease activity of another human MutL protein in loop-directed DNA expansion (Figs. 2–4 and SI Appendix, Figs. S6 and S7). Unlike MutLα, MutLγ incises triplet repeat expansion-containing ccDNA in a reaction that does not depend on PCNA and RFC (Fig. 4).

Human MutLγ alone incises a supercoiled DNA in the presence of Mn\(^{2+}\) and 70 mM KCl + NaCl (Fig. 1C). However, human MutLγ alone does not cleave the (CTG)\(_3\)/(CAG)\(_1\) heteroduplex ccDNA in the presence of Mn\(^{2+}\) at an ionic strength that approximates physiological conditions (140 mM KCl + NaCl) (SI Appendix, Fig. S9). We attribute this difference to suppression of Mn\(^{2+}\)-dependent endonuclease activity at the higher salt concentration (SI Appendix, Fig. S10). Both MutLα and MutLγ endonucleases promote DNA expansion in a manner that requires the integrity of the DOHA(X)\(_2\)E(X)\(_4\)E motif (17) (Fig. 3). Nevertheless, there is a fundamental difference between the behaviors of the two proteins: whereas the activated MutLα endonuclease incises either strand of loop-containing ccDNA (17), the activated MutLγ only incises the loop-lacking strand (Figs. 2 and 4 and SI Appendix, Fig. S7). This difference in the behaviors of the two nucleases provides a simple explanation for the fact that in the H6 extract system the activated MutLγ exclusively promotes DNA expansion, whereas the activated MutLα endonuclease promotes both DNA expansion and loop removal (17) (Fig. 3).

The activated MutLγ endonuclease cleaves the 3-nt loop-containing ccDNA at numerous sites on the loop-lacking strand (Fig. 2A, lane 3). However, the pattern of incision of the (CTG)\(_3\)/(CAG)\(_3\) and (CTG)\(_3\)/(CAG)\(_3\) heteroduplex ccDNAs by the activated MutLγ endonuclease is different (Fig. A4 and SI Appendix, Fig. S7B). In the case of the (CTG)\(_3\)/(CAG)\(_3\) heteroduplex ccDNA, activated MutLγ endonuclease often cleaves the loop-lacking strand within two regions on either side of the loop (Fig. A4). For the (CTG)\(_3\)/(CAG)\(_3\) heteroduplex ccDNA, the activated protein frequently cuts the loop-lacking strand at the region that encompasses the loop (SI Appendix, Fig. S7B), although cleavage does occur at several other regions as well. Similar to the activated MutLγ (Fig. 4), the activated MutLα often cleaves the (CTG)\(_3\)/(CAG)\(_3\) heteroduplex at several preferred regions that are near the 6-nt loop (17). These different modes of endonuclease action on the several DNAs presumably reflect differences in the structural nature and/or lifetimes of the protein-DNA complexes involved.

**Materials and Methods**

Annealing of DNA strands was carried out and quantified with a phosphorimager data and are presented as averages ±1 SD (n = 4).

**Fig. 4.** Human MutLγ incises the loop-lacking strand of a (CTG)\(_3\)/(CAG)\(_1\) heteroduplex ccDNA in a reaction that displays a significant site specificity. Defined reactions that occurred in the presence of Mg\(^{2+}\) were carried out and analyzed as described under Materials and Methods. The products of the indicated reactions were cleaved with Scal, separated in 1% denaturing agarose gels, and visualized by Southern hybridizations with \(^{32}\)P-labeled oligonucleotides S-GTGTATCCGCGGACCGGAGTTGCTCTTG-3' (A) and S-CAGAGGCAATCGGTCGCCGATACACA-3' (B). (A) Incision of the loop-lacking strand of the (CTG)\(_3\)/(CAG)\(_1\) heteroduplex ccDNA by the activated MutLγ endonuclease. (B) Lack of incision of the loop-containing strand of the (CTG)\(_3\)/(CAG)\(_1\) heteroduplex ccDNA by the activated MutLγ endonuclease. The arrows mark locations of DNA products that were formed by cleavage of the ccDNA at the loop site. The diagrams on the left depict the (CTG)\(_3\)/(CAG)\(_1\) heteroduplex ccDNA and show the relative positions of the 6-nt loop, the Scal site, and the \(^{32}\)P-labeled probes. (C) Quantification of incision of the two strands of (CTG)\(_3\)/(CAG)\(_1\) heteroduplex ccDNA by the activated MutLγ endonuclease. The incision values were determined from phosphorimager data and are presented as averages ±1 SD (n = 4).
Like its homolog MutLα (11, 12), human MutLγ has Mg2+- and Mn2+-dependent endonuclease activities that nick DNA (Figs. 1, 2, and 4 and SI Appendix, Figs. S5 and S7–S9). The Mn2+-dependent endonuclease activities of human MutLα and MutLγ are promoted by ATP (11, 12) (Fig. 1D). This supports the view that the two human MutL proteins act as ATP-dependent endonucleases. In contrast, the endonuclease activity of yeast MutLγ is not influenced by ATP (52, 53). The comparison of the specific Mn2+-dependent endonuclease activities of human MutLγ and MutLα revealed that they are in the same range (Fig. 1C), which indicates that the mechanisms of DNA nicking in the active sites of human MutLα and MutLγ endonucleases may be similar. An important question is, does the Mn2+-dependent endonuclease activity of MutLγ contribute to DNA expansion? Given that the intracellular concentration of Mn2+ is likely to be low (~30 μM with only 0.7 μM free) (62), the Mn2+-dependent endonuclease activity of human MutLγ probably does not play a significant role in DNA expansion when the intracellular Mn2+ concentration is in the normal range (SI Appendix, Fig. S9).

MutSβ recognizes 1- to 12-nt loops (63, 64) and plays an important role in triplet repeat expansion (5, 30). Consistent with this, we have determined that human MutSβ activated the Mg2+-dependent endonuclease activity of human MutLγ to incise relaxed ccDNA in a loop-dependent manner (Figs. 2 and 4 and SI Appendix, Fig. S7). Furthermore, we have determined that human MutSβ promoted the Mn2+-dependent endonuclease activities of human MutLγ and MutLα (Fig. 1C and SI Appendix, Figs. S9 and S11). This is in agreement with a previous study that showed that yeast MutSβ stimulated the Mn2+-dependent endonuclease activity of yeast MutLγ on supercoiled homoduplex DNA (53). Thus, it is likely that the ability of MutSβ to interact with MutLγ and MutLα endonucleases has been conserved throughout evolution of eukaryotes. Because both MutLγ and MutLα contain MLH1 as a subunit, it would be important to determine whether the PCNA-binding motif, which is located near the N terminus of its MSH3 subunit (65). It would be important to study whether the PCNA-binding motif of MutSβ is also required for the interaction of this mismatch recognition factor with MutLγ endonuclease. Although MutSβ activates the Mn2+-dependent endonuclease of both yeast and human MutLγ on a supercoiled homoduplex, the basis of this activation is unclear, but it may involve recognition of non-B DNA structures, the formation of which is driven by superhelical free energy.

Previous research (17, 33) and our findings suggest a model for MMR system-dependent triplet repeat expansion events that may occur in nonreplicating DNA (Fig. 5). In this model, expansion is a net result of multiple events of two different classes. One class of event depends on MutSβ, MutLα, PCNA, and RFC, and the other class of event requires MutSβ and MutLγ. RFC contributes to the MutLα-dependent event by loading PCNA onto the loop-containing DNA (17). Both the MutLα-dependent and MutLγ-dependent events are initiated by recognition of a small loop in the triplet repeat DNA by MutSβ. After recognizing a small loop, MutSβ cooperates with loaded PCNA to activate MutLα or it acts alone to activate MutLγ. The activated MutLα or MutLγ incises the loop-containing DNA. The incision of the loop-containing DNA by the activated MutLα endonuclease occurs in the loop-containing or loop-lacking strand, whereas the cleavage of the loop-containing DNA by the activated MutLγ endonuclease only takes place in the loop-lacking strand. Pol β (66, 67) or another DNA polymerase utilizes the generated strand break to perform a DNA synthesis reaction that leads to triplet repeat expansion or contraction in the event that involves MutLα endonuclease and to triplet repeat expansion in the event that involves MutLγ endonuclease. After the DNA synthesis step, the nick is sealed by DNA ligase I or III (68). If this model is correct, it would be important to determine the relative contributions of the MutLα- and MutLγ-dependent events to triplet repeat expansion.

A consensus that has emerged from studies of triplet repeat instability indicates that triplet repeats can be expanded via MMR system-dependent and independent mechanisms (29, 33, 69, 70). Importantly, genetic studies in mice have linked the MMR system to both germ line expansion and somatic expansion/contraction events that involve a relatively small number of repeat units (16, 21, 30, 37, 71). Thus, our model for MMR system-dependent triplet repeat expansion (Fig. 5) might account for conversion of normal alleles, especially long normal alleles, into premutation alleles and for small-scale somatic expansions that take place in expanded alleles in Huntington’s disease. However, MMR-dependent events may be not necessary for production of large intergenerational expansions that have been observed in some repeat expansion diseases.

Materials and Methods

Proteins, H6 Cell-Free Extract, and DNAs. Human MutLα, MutSβ, PCNA, and RFC were isolated in near homogeneous forms as previously described (11, 14, 65). Human MutLβ and MutLγ-D1223N proteins containing a FLAG tag at the N terminus of their MSH3 subunits were produced in insect Sf9 cells and then purified by chromatography on M2 anti-Flag beads (Sigma) and a MonoQ column (GE HealthCare). Human MLH1−/− MSH2−/− H6 cells were grown, and their cytosolic extracts were prepared as previously described (72).

The relaxed 3-nt loop-containing ccDNA and the relaxed control homoduplex ccDNA were prepared using the gapped form of the plasmid pAH1A (73) according to previously described procedures (17, 74). To prepare the 3-nt loop-containing ccDNA a phosphorylated 39-mer oligonucleotide with the sequence 5′-GCTACGTTCCAGAATCTCCGATCGGAGTCCGACG-3′ (the 3-nt loop sequence is underlined) was utilized. The control homoduplex ccDNA was prepared using a phosphorylated 36-mer oligonucleotide 5′-GCTACGTTCCAGAATCTCCGATCGGAGTCCGACG-3′. The relaxed (CTG)1/(CAG)1 and (CTG)1/(CAG)3 heteroduplex and (CTG)γ/(CAG)3 homoduplex cDNAs that carry a part of 3 untranslated region of the human DMPK gene were prepared as previously described (17). The DNA sequence of the human DMPK gene 3′ untranslated region in the homoduplex DNA was 5′-GCTACGTTCCAGAATCTCCGATCGGAGTCCGACG-3′, and the DNA sequences of the human DMPK gene 3′ untranslated region in the loop-containing strands of the relaxed (CTG)γ/(CAG), and (CTG)γ/(CAG)
Endonuclease, ATPase, and DNA Expansion Assays. Endonuclease, ATPase, and DNA expansion assays were based on previously developed methods (11, 12). Details of these assays are available in SI Appendix.

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