EMBO MEMBER'S REVIEW

Replication errors: cha(lle)nging the genome

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Since the discovery of a link between the malfunction of post-replicative mismatch correction and hereditary non-polyposis colon cancer, the study of this complex repair pathway has received a great deal of attention. Our understanding of the mammalian system was facilitated by conservation of the main protagonists of this process from microbes to humans. Thus, biochemical experiments carried out with Escherichia coli extracts helped us to identify functional human homologues of the bacterial mismatch repair proteins, while the genetics of Saccharomyces cerevisiae aided our understanding of the phenotypes of human cells deficient in mismatch correction. Today, mismatch repair is no longer thought of solely as the mechanism responsible for the correction of replication errors, whose failure demonstrates itself in the form of a mutator phenotype and microsatellite instability. Malfunction of this process has been implicated also in mitotic and meiotic recombination, drug and ionizing radiation resistance, transcription-coupled repair and apoptosis. Elucidation of the roles of mismatch repair proteins in these transduction pathways is key to our understanding of the role of mismatch correction in human cancer. However, in order to unravel all the complexities involved in post-replicative mismatch correction, we need to know the cast and the roles of the individual players. This brief treatise provides an overview of our current knowledge of the biochemistry of this process.

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Introduction

Maintenance of genomic stability is one of the key criteria that govern the survival of species. DNA is a reactive molecule and as such is modified continuously by a broad range of agents. Although exogenous sources of DNA damage, such as ionizing and UV radiation or carcinogens contained in foodstuffs and cigarette smoke, have received the greatest share of attention over the past 30 or so years, DNA is damaged primarily from within: through hydrolysis, methylation or active oxygen species (Lindahl, 1996). Given that a mammalian DNA molecule can undergo ~100 000 modifications per day, it is clear that life on Earth would be extinct by now were it not for DNA repair (Friedberg *et al.*, 1995). This term encompasses a multitude of metabolic processes, which can reverse the damage either directly (photolyases, alkyltransferases), or indirectly by removing damaged bases (base excision) or oligonucleotides (nucleotide excision) from DNA, and resynthesizing the removed patch. Moreover, they can repair strand breaks and cross-links, or by-pass non-repairable damage either by specialized mechanisms or by recombination (Friedberg *et al.*, 1995). The main task of DNA repair is to ensure that the DNA molecule is free of modifications or mutations, such that it can be transcribed efficiently and, most importantly, that it can be replicated faithfully and passed on to progeny cells.

DNA replication is a complex process, whose fidelity, estimated to be in the range of one error per 10¹⁰ nucleotides synthesized (Kunkel, 1992), depends on three factors: DNA polymerase(s), exonucleolytic proofreading and mismatch repair (MMR). Replicative DNA polymerases are extremely precise enzymes, which can duplicate the sequence of a given genome within minutes, or at most hours, with an error rate of $\sim 10^{-5}$ (Kunkel, 1992). The high precision with which complementary nucleotides are added to the end of the primer strand is guaranteed by the formation of Watson-Crick base pairs in the active site of the enzyme. This process is controlled by the thermodynamic stability afforded by the formation of two or three hydrogen bonds between the incoming nucleoside triphosphate and the first non-paired base of the template strand, as well as by spacial restrictions within the polymerase active site.

Should a non-complementary nucleotide be incorporated at the end of the primer, extension from such a primer is highly inefficient (Benkovic and Cameron, 1995) and it is in fact likely that the inability of DNA polymerases to go forward in such situations leads to a translocation of the mispaired primer terminus into the active site of the proofreading $3' \rightarrow 5'$ exonuclease activity that is associated with all replicative polymerases and that adds a further two orders of magnitude (Kunkel, 1992) to the fidelity of the replication process by excising the terminal noncomplementary nucleotide, together with two or three additional residues, from the end of the primer terminus.

In some cases, a mispair manages to elude the proofreading process. This happens mostly with the G/T wobble pair, which is stabilized by two hydrogen bonds and brings about only a slight distortion of the double helix (Hunter *et al.*, 1987). As it is the one mispair from which most polymerases are able to extend with the least difficulty (Benkovic and Cameron, 1995), it also happens to escape the translocation into the proofreading site and exits the polymerase complex. A similar situation can arise when the primer and template strands slip with respect to one another (Kunkel, 1990). Such events occur relatively frequently in runs of repeated mono- or dinucleotides,

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where they give rise to loops containing extrahelical nucleotides, the so-called insertion/deletion loops (IDLs). When an IDL is formed by transient dissociation and reassociation of the primer and the template, or by simple slippage, the end of the primer strand will anneal with the template to produce a hydrogen-bonded terminus from which the polymerase can extend (Kunkel, 1993).

Replication-associated transactions such as mispairs and IDLs that have escaped the proofreading exonuclease become substrates for MMR, whose task is to restore the information contained in the template strand. In this respect, MMR differs from all other DNA repair pathways. Mismatches or IDLs are composed of unmodified nucleotides and exist as detectable moieties in DNA solely in the double-stranded form, inasmuch as separation of the two strands yields two unmodified, intact DNA molecules, neither of which contains recognizable-and therefore repairable—damage. Successful restoration of the original DNA sequence thus requires (i) a factor capable of recognizing base-base mismatches and IDLs, and (ii) a factor that can distinguish between the parent and the daughter strand. It is beyond the scope of this article to review the genetics and the biochemistry of MMR in its entirety. Fortunately, following the discovery of a link between MMR malfunction and a frequent form of human cancer, hereditary non-polyposis colon cancer (HNPCC), this subject has become the topic of intense research and, as a result, interested readers can find a detailed treatise in numerous reviews on MMR that have appeared in the recent literature (Fishel and Kolodner, 1995; Kolodner, 1995, 1996; Radman et al., 1995; Jiricny, 1996; Marra and Boland, 1996; Modrich and Lahue, 1996; Crouse, 1997; Fishel and Wilson, 1997; Modrich, 1997; Fink et al., 1998). The scope of this minireview is to discuss the recent developments in our understanding of the biochemistry of the MMR process and to address some issues that have been little appreciated or those that merit, in my view at least, particular attention.

Mismatch repair in E.coli

The process of post-replicative mismatch correction in *E.coli* represents the best characterized system to date (for reviews see Modrich, 1991; Modrich and Lahue, 1996). The individual factors associated with mismatch recognition, strand discrimination, exonucleolytic degradation of the error-containing strand, resynthesis of the repair patch and ligation have been identified, and there is reason to believe that at least our global concept of the process is correct, as MMR reconstituted from these 10 factors functions *in vitro* (Lahue *et al.*, 1989). Mechanistically, mismatch correction in *E.coli* is thought to proceed as outlined in Figure 1. In the following paragraphs, I shall address some points regarding this model that remain to be clarified.

Criteria for mismatch recognition

There are still numerous details that we do not fully understand. One of these concerns the initial step of the correction cascade, namely mismatch recognition, a role fulfilled by the homodimeric MutS protein (Su and Modrich, 1986). It could be shown in different *in vitro* assays that this factor binds with high affinity to substrates containing most base-base mispairs (Su and Modrich, 1986; Su et al., 1988) and IDLs up to four extrahelical nucleotides (Parker and Marinus, 1992). How these substrates are recognized is not understood. The distortions brought about in the double helix by a G/T mispair as compared with a single extrahelical nucleotide, for example, are dramatically different. The former is hydrogen-bonded, well-stacked and relatively stable (Hunter et al., 1987), whereas the latter either causes a considerable bend and/or perturbation of stacking interactions in the DNA if the extra base is intercalated, or a bulky loop if the unpaired nucleotide is extrahelical. Remarkably, both substrates are bound with a similar affinity in the bandshift assay. More surprisingly still, notable differences in in vitro binding affinity were observed for oligonucleotide substrates containing the G/T or the A/C mispair, where the former was bound strongly and the latter almost undetectably, even though the purines and the pyrimidines were in an identical sequence context. Yet the C/A mispair was bound as well as the G/T and much more strongly than the T/G in these substrates (Jiricny et al., 1988b). These puzzling observations were confounded further by the fact that the efficiency of repair of all these mispairs in vivo was similar. Thus, the take-home message from the binding studies is that affinity of the protein for a particular mispair or a DNA modification in vitro cannot be taken as an indication of repair efficiency in vivo. It would be interesting to examine the fine structures of these particular mismatch combinations and IDLs by highresolution NMR or by X-ray crystallography, such that we might gain an important insight into the types of structural determinants recognized by MutS.

The fact that this mismatch-binding protein has so far eluded crystallization is very frustrating indeed. However, a recent study carried out with the Thermus aquaticus MutS homologue has provided us with the first insight into the interaction of this multifaceted polypeptide with its substrate. Using an iododeoxyuridine-labelled oligonucleotide in UV cross-linking, Malkov et al. (1997) were able to demonstrate that the MutS interaction with DNA was mediated via Phe39, which is situated within a short, but highly conserved motif in the N-terminal domain of the protein. Substitution of this residue for Ala decreased the affinity of the mutant protein for DNA by three orders of magnitude, but the mutation did not affect the ability of the protein to dimerize, nor did it alter its ATPase activity. This suggests that MutS and its homologues interact with DNA via their N-termini, while the C-terminal domains house the dimerization and ATPbinding domains. More studies of this kind would be welcome.

Repair complex assembly

The second mystery concerns the transactions that follow mismatch recognition. Electron microscopic data (Allen *et al.*, 1997) show that the mismatch-bound MutS homodimer undergoes an ATP-driven translocation along the DNA such that a looped structure is formed, with the protein(s) sitting at its base. (This structure is sometimes referred to as the α -loop, although I prefer to think that it resembles the Greek letter Ω rather than α .) This process is accelerated by the addition of the MutL homodimer, which appears to co-localize with MutS. Measurement of

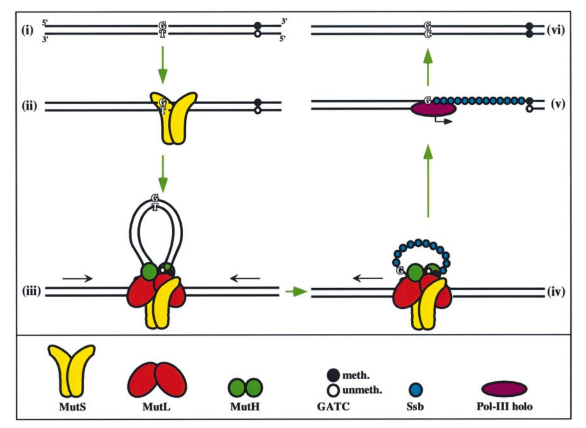


Fig. 1. Suggested mechanism for mismatch correction in *E.coli*. (i) The G/T mispair that arose as an error of DNA polymerase is present in a DNA heteroduplex, which is transiently unmethylated at a GATC sequence in the newly replicated strand. The template GATC site is methylated. (ii) Binding of the mismatch by the homodimeric MutS protein initiates a cascade of events that starts with an ATP-dependent conformational change of MutS, followed by a movement of the bound protein away from the mispair (not shown). (iii) ATP hydrolysis drives the bi-directional movement of the DNA through the bound MutS in the direction of the arrows, as well as the assembly of a multiprotein complex containing the MutS and MutL homodimers and probably also two molecules of the strand discrimination factor MutH, all bound at the base of a looped structure. The assembly of the complex activates the endonucleolytic activity of MutH, which cleaves the newly synthesized DNA 5' from the unmethylated GATC sequence. (iv) The cleaved strand is then degraded from the nick up to and slightly past the mismatch site either by *Exo*VII or *RecJ* (in cases where the unmethylated GATC was situated 5' from the mispair), or by *Exo*I (when the nick was 3' from the mispair). The single-stranded region thus generated is protected by the single strand-binding protein Sb. This step is discussed in more detail in the text. (v) Polymerase III holoenzyme fills the gap and DNA ligase repairs the nick. (vi) The process is completed by methylation of the GATC site by Dam methylase, at which point the substrate becomes refractory to further action by the MMR system.

the loop size and of the mispair-to-terminus distances instigated the authors to suggest that the translocation resulted in a release of the mispair by the protein, such that it is now at the apex of the loop (see Figure 1). This hypothesis has several far-reaching implications. Binding experiments demonstrated that the affinity of MutS for a homoduplex is several fold lower than for a mismatchcontaining heteroduplex (Su et al., 1988). Yet, if the mismatch-bound protein were indeed to release the mispair in the presence of ATP, it would be bound to homoduplex DNA. It is difficult to imagine why the protein should let go of its preferred substrate, unless it were to undergo an ATP-dependent conformational change which would increase its affinity for homoduplex DNA. The electron microscopic data (Allen et al., 1997), moreover, imply that the MutS homodimer sitting at the base of the loop interacts with two double helices, i.e. that each subunit binds to the same homoduplex DNA molecule, albeit several hundred base pairs apart. The only evidence that supports the conclusion that both DNA-binding sites in the MutS homodimer can be occupied comes from studies carried out with the MutS homologue from T.aquaticus, which appears to interact with oligonucleotide substrates with a 1:1 stoichiometry, i.e. two oligonucleotides bound to one MutS homodimer (P.Hsieh, personal communication). It would therefore appear feasible that the MutS homodimer could bind to the homoduplex regions at the base of the loop as depicted in Figure 1, where the ATPdirected conformational change, coupled to the gain in energy through interaction with two DNA homoduplexes (rather than one heteroduplex), would ensure stability of the complex.

It is possible that the conformational change of MutS is catalysed by MutL. In a series of gel-shift experiments, Marinus and colleagues (Drotschmann *et al.*, 1998) have noticed that the presence of MutL in the MutS binding assay increased the efficiency of MutS binding. As the size of the protein–DNA complex did not increase, at least as measured by its mobility through native polyacrylamide gels, it would appear that the MutL protein somehow assisted the loading of MutS onto the mismatched substrate, without being itself a part of the complex. It is interesting to note in this context that the MutL protein has already been assigned a role of a 'molecular matchmaker', in that it is thought to mediate the interaction between MutS and MutH (Modrich, 1991). Moreover, the

role of MutL as a possible chaperone is underscored by the sequence homology of its N-terminus with those of Hsp90 and type II DNA topoisomerases (Bergerat *et al.*, 1997; Grenert *et al.*, 1997). Although no evidence of a DNA–MutS–MutL ternary complex was obtained in gelshifts, the experiments of Allen *et al.* (1997) provide evidence for the fact that the three polypeptides are located together at the base of the loop.

Recent evidence suggests that MutL might play a role also in loading the UvrD (MutU) helicase at the site of the nick (Dao and Modrich, 1998; Yamaguchi *et al.*, 1998). This represents one of the key steps of the repair process, inasmuch as it would help the helicase initiate the unwinding process, due to the fact that most helicases have difficulties with initiation at a nick within a doublestranded DNA molecule. More importantly though, the MutL–UvrD interaction may control the directionality of the unwinding process such that it proceeds towards, rather than away from, the mispair. It is conceivable that the torsional strain of the looped structure might also contribute to the directionality of the process of unwinding and exonucleolytic degradation of the error-containing strand.

The model as shown in Figure 1 therefore appears to fit most of the experimental data, with one notable exception: repair-patch size. As mentioned above, the EM results of Allen et al. (1997) show that the mismatchdependent, ATP-driven translocation of the MutS-MutL complex involves the bidirectional threading of the DNA through the proteins such that the released mispair finds itself roughly equidistant from the two extremities of the loop. However, measurement of the repair patch size in the same system showed that the degradation began at the nick and proceeded up to and just past the mismatch (Su et al., 1989). This implies that the exonucleolytic degradation of the newly synthesized strand stops shortly after the non-complementary nucleotide had been removed. In the repair of damaged bases or nucleotides, such a possibility would not be questioned. However, as mentioned above, mismatches do not contain modified nucleotides and are undetectable when the two DNA strands have been separated by, for example, a helicase. The fact that the degradation mostly stops only a short distance past the site where the mismatch had been suggests that the site is either tagged, or that, following the MutH-mediated binding and nicking of the nearest GATC site, the threading direction is reversed, such that the mismatch returns to the MutS fold. This would now be an asymmetric process, because only one half of the loop is free to migrate-the other being bound by the MutH protein. This model is purely speculative; we have no evidence at present in support of either mechanism. Further experiments aimed at refining it are clearly needed.

Mismatch repair in human cells

The discovery of sequence similarities between the PMS1 protein of *Saccharomyces cerevisiae* and the MutL protein of *E.coli* (Kramer *et al.*, 1989) represented an important milestone in the study of eukaryotic MMR, inasmuch as it provided the first evidence of a close conservation of the pathway between prokaryotes and higher organisms. Moreover, it became clear that the post-meiotic segregation

(PMS) phenotype, described by Fogel and colleagues (Williamson et al., 1985), was closely linked with MMR and thus implied an important role for this process in meiotic recombination. The availability of yeast MMR mutants helped us to understand the phenotypes of human cells with deficiencies in MMR. Thus, while a strong mutator phenotype was widely expected, MMR deficiency appeared to accentuate an instability of microsatellite repeats (Strand et al., 1993), a trait that was to prove of immense importance in the identification of human tumours with MMR gene defects (Boyer et al., 1995). Moreover, the characterization of the S.cerevisiae MutS and MutL homologues (MSH and MLH, respectively) made possible the identification of human MMR genes by degenerate PCR (Leach et al., 1993; Fishel et al., 1994c; Nicolaides et al., 1994; Papadopoulos et al., 1994; Kolodner et al., 1995), thanks to the conservation of the amino acid sequences from bacteria to yeast to man. Due to this conservation, the MMR process in humans appears at first glance very similar to that shown in Figure 1. However, upon closer examination, several key differences emerge, the most notable of which are the heterodimeric nature of both MutS and MutL homologues and the lack of a strand discrimination mechanism resembling the Dam methylase-MutH endonuclease interplay. Moreover, the system has an in-built redundancy.

MutS homologues and mismatch recognition

A factor binding with high affinity to oligonucleotide substrates containing G/T mispairs was first described in 1988 (Jiricny et al., 1988a). It was later purified to near homogeneity (Hughes and Jiricny, 1992) and, due to its preference for G/T mispair-containing substrates, was named GTBP (G/T-binding protein). The purest fraction contained two polypeptides of apparent molecular mass of 100 and 160 kDa, which were at the time thought to be related, in that the smaller protein was thought to be a proteolytic fragment of the 160 kDa protein. However, sequencing of tryptic peptides derived from the two proteins revealed that they were encoded by distinct genes (Palombo et al., 1995). The 100 kDa protein was shown to be the product of the hMSH2 gene (Palombo et al., 1994), which was identified a few months previously (Leach et al., 1993; Fishel et al., 1994c) as the first locus of HNPCC, on chromosome 2p15-16 (Peltomaki et al., 1993). hMSH2, which stands for human MutS homologue 2, was so named because it is closely related to the S.cerevisiae MSH2 protein, first described in 1992 (Reenan and Kolodner, 1992). The gene encoding the 160 kDa polypeptide co-localizes with hMSH2 (Papadopoulos et al., 1995). It originally was named GTBP (Hughes and Jiricny, 1992), but because of the close relatedness of its product to the S.cerevisiae MSH6 protein described some months later (Marsischky et al., 1996), the name hMSH6 has now been adopted.

hMSH2 and hMSH6 are tightly associated and copurify through a series of chromatographic steps (Hughes and Jiricny, 1992; Drummond *et al.*, 1995). The purified heterodimer, often referred to as hMutS α , has been shown to complement extracts of cells mutated in *hMSH2* or *hMSH6* in an *in vitro* MMR assay (Drummond *et al.*, 1995). Attempts at expression of the individual proteins in the baculovirus system met with only partial success. hMSH2 was expressed in high yields and could be purified easily to homogeneity (Fishel *et al.*, 1994a,b; Iaccarino *et al.*, 1998). Although we found the protein to lack specific mismatch-binding activity in our gel-shift assays (Iaccarino *et al.*, 1998), several reports using recombinant hMSH2 described its ability to bind to substrates containing base–base mispairs, as well as IDLs of varying sizes (Fishel *et al.*, 1994a,b). Similar results were reported also for the yeast MSH2 protein (Alani *et al.*, 1995), with the sole difference that the binding of the human protein to heteroduplex DNA was augmented in the presence of ATP (Fishel *et al.*, 1994b), whereas the *S. cerevisiae* protein binding was unaltered in the presence of the nucleotide (Alani *et al.*, 1995). We currently are attempting to identify the cause underlying these differences.

Expression of hMSH6 was less successful. The fulllength protein appeared proteolytically labile and could be isolated only in poor yields. It displayed no mismatchbinding activity (Iaccarino *et al.*, 1998). It was, however, considerably stabilized when co-expressed with hMSH2 (Palombo *et al.*, 1996; Gradia *et al.*, 1997; Iaccarino *et al.*, 1998), which permitted the isolation of soluble hMutS α in high yields. This heterodimeric factor bound G/Tcontaining substrates with a low nanomolar affinity (Gradia *et al.*, 1997; Iaccarino *et al.*, 1998).

All MutS homologues characterized to date are highly conserved at their C-terminal regions, which contain the consensus ATP-binding site. Addition of ATP to gel-shift assays results in an apparent dissociation of the proteins from the oligonucleotide substrates (Hughes and Jiricny, 1992; Gradia et al., 1997; Iaccarino et al., 1998), which is most likely to be the result of their running off the end of the short duplex following the ATP-dependent translocation described above for MutS (see also below). This ATP-dependent process requires only ATP binding, not ATP hydrolysis (Hughes and Jiricny, 1992; Gradia et al., 1997; Iaccarino et al., 1998). Mutations within the conserved GxxxxGKS motif of the Salmonella typhimurium MutS (Haber and Walker, 1991), the S.cerevisiae MSH2 and MSH6 (Alani et al., 1997; Studamire et al., 1998) or the human hMSH2 and hMSH6 proteins (Iaccarino et al., 1998) had little influence on mismatch binding, but brought about a considerable decrease in the sensitivity of the protein–DNA complex to ATP. Assuming that, following binding at the mismatch site, hMutS α undergoes an ATP-dependent conformational change and translocates along the double helix in a manner similar to the MutS homodimer, thereby generating a looped structure, it would be predicted that ATPase-defective mutants would be MMR deficient. This is indeed the case, although both hMSH2 and hMSH6 sites needed to be mutated before MMR activity was lost in an in vitro assay (Iaccarino et al., 1998).

Recently, Fishel and colleagues (Gradia *et al.*, 1997) suggested that the hMSH2/6 heterodimer acts as a molecular switch homologous in some way to G proteins, being ON for mismatch binding in an ADP-bound form and OFF when complexed with ATP. This suggestion stems from the finding that, as already mentioned, the heterodimer forms stable complexes with mismatch-containing oligonucleotide substrates, which dissociate upon the addition of ATP. This interesting hypothesis needs to be substantiated further. Previous experiments have shown

that the heterodimer does not require any cofactors for binding to heteroduplex DNA; it binds oligonucleotide substrates with a similar affinity in the presence or absence of ADP (Hughes and Jiricny, 1992), and it can exchange ADP for ATP without the need for additional exchange factors (I.Iaccarino and J.Jiricny, unpublished). However, it is possible that the presence of other proteins, such as the MutL homologues, might accelerate the exchange. It seems likely that this ATP-dependent switch is in fact a conformational change of hMutS α that is required for the binding factor to leave the mismatch site and thus initiate the downstream repair events as proposed (Gradia *et al.*, 1997). There is evidence that a conformational change does indeed take place upon ATP binding by the *S.cerevisiae* MSH2/6 heterodimer (Studamire *et al.*, 1998).

Although the in vitro binding experiments are interesting, they do not provide us with much information regarding the sequence of events at the mismatch following the binding of hMutS α . It is unlikely that the factor will leave DNA upon ATP binding (as suggested by the gelshift experiments), without first either marking the site or inducing the assembly of the repairosome. As the oligonucleotide substrates are apparently intact and unmodified following ATP-driven hMutS α dissociation, we can only assume that mismatch binding catalyses the assembly of the repair complex in a manner similar to that described for *E.coli* (see Figure 1). This would imply that the complex remains bound to the DNA, but not at the mismatch, i.e. that it translocates following ATP binding. Although EM evidence of this process in the human system is not yet available, ATP fails to mediate the dissociation of hMutS α from a 200 bp mismatchcontaining duplex when both ends of the molecule are blocked with streptavidin (L.Blackwell and P.Modrich, personal communication). This suggests that the protein slides along the molecule as proposed for MutS.

Although hMSH2 and hMSH6 function as a heterodimer in MMR, the phenotype of cells mutated in the hMSH2 gene is dramatically different from hMSH6 mutants. The former cells are deficient for the repair of base-base mismatches and small IDLs, and correspondingly display instability in microsatellite sequences consisting of runs of mono-, di-, tri- and tetranucleotides. In contrast, hMSH6 mutants are deficient for the repair of base-base mismatches and IDLs of one nucleotide, but dinucleotide repeat instability was not noted in these cells (Papadopoulos et al., 1995) and their extracts are partially proficient for the repair of IDLs of two nucleotides or more (Drummond et al., 1995). These finding led to the proposal (Karran, 1995) that loop repair is dependent on another mismatch-binding factor that contains hMSH2, but not hMSH6. This hypothesis could soon be substantiated. Genetic experiments in S.cerevisiae demonstrated that the mutator phenotype of the msh2 strain was similar to that of a double mutant in msh3 and msh6 (Marsischky et al., 1996), and it was suggested that the MSH3 and MSH6 proteins can both interact with MSH2, whereby the MSH2/3 heterodimer displays affinity for IDLs, while the MSH2/6 factor might preferentially bind base-base mismatches and loops of one nucleotide. Indeed, using recombinant hMSH2/6 (hMutS α) and hMSH2/3 (hMutS β) heterodimers expressed in the baculovirus system (Acharya et al., 1996; Palombo et al., 1996), it could be shown that the former

could bind base-base mismatches and IDLs of one and two nucleotides, while the latter failed to bind to oligonucleotide substrates containing base-base mismatches, but displayed considerable affinity for IDL substrates. In other studies, using proteins purified from human HeLa cells, it could be shown that hMutSa could complement hMSH2deficient extracts for the repair of base-base mismatches and small as well as large loops, while only loop repair was restored to these extracts by the addition of hMutS β (Genschel et al., 1998; Marra et al., 1998). It was proposed, therefore, that hMutS α most likely initiates the repair of both base-base mispairs and IDLs under most circumstances. Support for this hypothesis came on one hand from the examination of extracts of human cell lines, which showed that hMutS α is significantly more abundant than hMutS β (Drummond *et al.*, 1997) and, on the other hand, from the finding that cells lacking hMSH3 do not display microsatellite instability (Inokuchi et al., 1995). In agreement with these findings is also the fact that *msh3* mutants in S.cerevisiae have only a very weak mutator and PMS phenotype (Strand et al., 1995). hMutSa and hMutS β are therefore functionally redundant for the repair of IDLs, although it seems probable that the latter fulfils only a back-up function in MMR.

The existence of hMutS α and hMutS β implies that hMSH3 and hMSH6 have to compete for hMSH2. As already mentioned above, the outcome of this competition in normal cells is strongly biased in favour of hMutSa (Drummond et al., 1997; Marra et al., 1998), with the result that these cells are MMR proficient. However, this equilibrium could easily change if hMSH3 were overexpressed, due to the fact that hMutS β acts in the repair of IDLs and not base-base mispairs. Such a situation was shown to exist in methotrexate-resistant cell lines, where amplification of the DHFR locus resulted in a coamplification of the hMSH3 gene (Drummond et al., 1997; Marra *et al.*, 1998). It could be demonstrated that extracts of these cells contained normal levels of hMSH2, but that hMSH6 levels were significantly decreased, most probably due to the proteolytic degradation of the partnerless protein. In contrast, hMSH3 was abundant in these extracts. Correspondingly, these cells have a strong mutator phenotype (Caligo et al., 1990; Drummond et al., 1997) and are defective in the repair of base-base mismatches, but are proficient for IDL repair in vitro (Drummond et al., 1997; Marra et al., 1998). The clinical relevance of this phenomenon in vivo is not known at this time; however, its importance lies in the demonstration that MMR deficiency can be attained also without mutation, through a simple dysregulation of expression of one of its components.

Role of MutL homologues

It could be shown that, similarly to the MutS homologues, the human counterpart of MutL is a heterodimeric factor, hMutL α , consisting of hMLH1 and hPMS2 (Li and Modrich, 1995). As in the case of the bacterial MutL, the precise biochemical role of the eukaryotic MutL homologues in MMR is unclear. It was anticipated that hMutL α will interact with a mismatch-bound hMutS α in a fashion similar to MutS and MutL. Indeed, in a gelshift experiment (Prolla *et al.*, 1994), the *S.cerevisiae* MSH2 complex with a mismatch-containing oligonucleo-

tide was supershifted in the presence of MLH1 and PMS1 (note that PMS1 is the *S.cerevisiae* homologue of hPMS2). The caveat of this experiment is that MSH2 has since been shown to exist in a complex with MSH6 (see above), the latter protein being absent from this experiment. The relevance of this finding is therefore uncertain at this time. However, the S.cerevisiae MLH1-PMS1 complex was shown to enhance the binding of the MSH2/3 heterodimer to an IDL substrate (Habraken et al., 1997), and interaction of human MutS and MutL homologues on a DNA substrate was demonstrated recently by co-immunoprecipitation studies, where the anti-hMLH1 antibody also succeeded in bringing down hPMS2 and hMSH2 (the authors did not test for the presence of hMSH6, but it can be assumed that it too was present in the precipitate). This reaction was ATP-dependent (Gu et al., 1998). Interestingly, the co-immunoprecipitated proteins also contained proliferating cell nuclear antigen (PCNA), whose possible role in MMR is discussed below.

Strand discrimination in human MMR

As mentioned earlier, the process of correction of replication errors has to be directed to the newly synthesized strand. This means that the MMR system must possess a strand discrimination function. In E.coli and S.typhimurium, strand discrimination makes use of a transient undermethylation of the newly synthesized strand, and employs an endonuclease, the MutH protein, capable of specifically cleaving the unmethylated strand of the hemimethylated DNA molecule (Figure 1). This mechanism is very elegant, but it should be remembered that this Dam-directed system represents the exception rather than the rule. Thus, in Streptococcus pneumoniae, no such system appears to operate (Balganesh and Lacks, 1985). In lower eukaryotes such as S.cerevisiae, there is no or only very little DNA methylation, and a similar situation applies in Drosophila melanogaster. Vertebrates and plants do methylate their DNA, mostly at the 5-position of cytosine in CpG or CpXpG motifs, but the methylation patterns are irregular, inasmuch as there are stretches of DNA up to several kilobases long, the so-called CpG islands, which remain unmethylated. MutH-like, methylation-sensitive endonuclease involved in the correction of a mismatch in such a sequence stretch could nick either strand. Such an event would lead in the best case to undirected repair and in the worst scenario to a doublestrand break. Thus, contrary to one published report (Hare and Taylor, 1985), methylation is unlikely to direct MMR in higher organisms. Yet, all these systems possess a postreplicative MMR system capable of strand discrimination. So how does the MMR system manage to distinguish between the template and the primer strand? In vitro studies of the MMR process demonstrated that strand discontinuities, such as nicks, can direct the mismatchdependent exonucleolysis to the nicked strand (Holmes et al., 1990; Thomas et al., 1991). This is true for all the above-mentioned systems, including *E.coli* where a single nick can substitute for MutH (Au et al., 1992). Lacks and colleagues proposed a long time ago that the MMR system of S.pneumoniae is nick-directed (Balganesh and Lacks, 1985). Indeed, taking as a model the lagging strand, we know that the Okazaki fragments are perhaps 2 kb long and that a mismatch anywhere within such a piece of

DNA is well within the reach of the MMR system, which could use either end of the fragment for the initiation of the repair process. The problem was always presented by the leading strand, where DNA synthesis is continuous. Where is the discrimination signal in this strand? Is it possible that the template DNA contains an as yet unidentified epigenetic modification that could be used to distinguish between it and the newly synthesized strand? We have as yet no answers to these questions. However, recent evidence coming from several different experimental systems has begun to shed light on this problem. As briefly noted above, immunoprecipitation experiments with antihMLH1 antibodies also co-precipitated hMSH2, hPMS2 and PCNA (Gu et al., 1998). The reason why these authors looked for PCNA was most likely thanks to seminal work coming from Kunkel's laboratory, where a two-hybrid screen search for MLH-interacting proteins picked out a full-length cDNA clone encoding the *S.cerevisiae* PCNA (Umar et al., 1996). This was the first experimental evidence which hinted at an interaction between DNA replication and MMR, especially as in vitro MMR experiments implied that PCNA was involved not only in the expected resynthesis of the excised repair tract (Gu et al., 1998), but also at a step prior to this (Umar et al., 1996). Based on these results, the authors (Umar et al., 1996) suggested that PCNA bound to the primer termini at the replication fork might itself act as the strand discrimination signal. Extending this prediction a little further, I would offer the following model as to how strand discrimination in eukaryotes might work. Let us assume that the replicative polymerase has just incorporated a non-complementary nucleotide at the end of the primer strand, which then escaped the proofreading activity of the replication complex. If the MMR proteins were able to physically interact with the replication enzymes via PCNA, detection and binding of the mispair by the MSH2/6 heterodimer, followed by a threading process that would eventually contact PCNA in the polymerase complex, would arrest the forward movement of the replication fork. This may well result in the dissociation of the polymerase from the primer terminus, an event which would leave the 3' nucleotide of the newly synthesized strand free and therefore accessible to any eventual 3'-5' exonucleases, which could then initiate the degradation of the primer strand towards the mispair (Figure 2). An equally feasible alternative is that the MMR proteins are linked directly to PCNA throughout the replication process. Although we have at present no experimental evidence in favour of this model, its beauty lies in the fact that it requires no MutH homologue for strand discrimination. The fact that no open reading frame which could be considered related to MutH (other than bacterial restriction endonucleases) has been detected in an ever-increasing number of organisms whose genomes have been completely (or almost completely) sequenced should come as no surprise, because MutH is a sequence-specific, methylation-sensitive endonuclease, with a preference for an epigenetically modified motif absent from the DNA of other organisms. However, it would appear probable that MMR in most species dispenses with a MutH-like function. Circular, mismatchcontaining heteroduplex molecules lacking a nick are generally refractory to MMR in vitro (Holmes et al., 1990; Thomas et al., 1991) and in vivo (Brown and Jiricny,

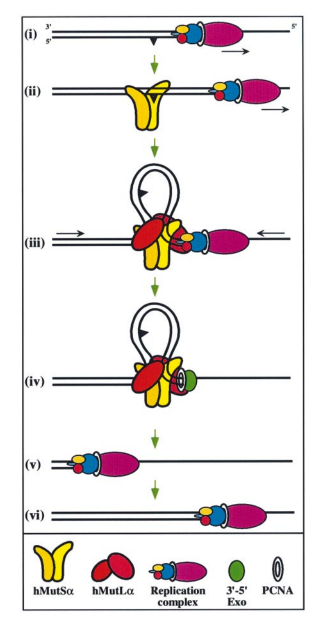


Fig. 2. Putative model of strand discrimination of human mismatch repair in the leading strand. (i) The mismatch, introduced into the newly synthesized strand by the replication complex consisting of pol- δ , PCNA, RFC and other replication factors, has escaped the proofreading exonuclease. (ii) The mispair is recognized by hMutSa. (iii) ATP drives the assembly of the MMR complex and the bi-directional threading of DNA through the hMutS α heterodimer until the replication complex is contacted via an interaction with PCNA. This event leads to replication arrest and dissociation of pol- δ . (iv) Exonucleolytic degradation of the error-containing primer strand can commence from its exposed 3' terminus. (v) Following the dissociation of the MMR complex, the PCNA, still bound at the end of the primer, recruits the replication complex and (vi) the DNA synthesis is re-initiated. Replication protein A (RPA), which probably fulfils the role of Ssb in this process, is covering the single-stranded DNA throughout the process and is not shown here. Another alternative is that the replication complex is physically linked with the MMR proteins already in steps (i) and (ii), and that it acts as a break which halts the polymerase complex upon mismatch binding (see text for details).

1987, 1988). (The small amount of repair in the latter experiments could be due to the introduction of sporadic strand breaks into the substrate DNA during the transfection procedure.) The notable exception to the rule might be *Xenopus laevis* extracts, which appear to be proficient for MMR on circular substrates (Brooks *et al.*, 1989; Varlet *et al.*, 1996). However, as the repair patches were short and centred predominantly around the mismatch, the possibility must be considered that the system addressing these mispairs was distinct from the post-replicative mismatch correction pathway.

Other proteins involved in MMR

The proteins participating in MMR can be divided into two categories. Those that are dedicated exclusively to mismatch correction and those that participate in other DNA metabolic pathways. In the case of *E.coli*, the former class is represented by the MutSLH polypeptides, while the helicase, exonucleases, polymerase and ligase belong to the second class. In human cells, we have already listed the members of the MMR-specific group: $hMutS\alpha$ (hMSH2/6) or hMutS β (hMSH2/3) and hMutL α (hMLH1/ hPMS2). A third MutL homologue, hPMS1, has also been identified (Nicolaides et al., 1994), but has to-date remained without a known biochemical function. As to the second group, the only activity known to participate in human MMR, at least in the *in vitro* assay, is polymerase- δ (Longley et al., 1997). A 5'-3' exonuclease, EXO1 (Tishkoff et al., 1997), has also been implicated in the process, largely based on the phenotype of S.cerevisiae and Schizosaccharomyces pombe mutant strains, but here it is difficult to say with any certainty what, if any, role these enzymes might play in the MMR process, due to their functional redundancy, as well as to the fact that they also participate in a number of other DNA transactions. RPA, the functional homologue of the bacterial single strand-binding protein Ssb, has also been implicated in the process (Lin et al., 1998; Umezu et al., 1998). The confirmation of the role(s) of these enzymes must await the results of ongoing biochemical studies. No evidence is as yet available regarding the helicase function. Of the four known human ligases, DNA ligase I is probably the one involved in MMR, but this is only an assumption based on its frequent association with pol- δ - and PCNAdependent processes (Montecucco et al., 1998).

Concluding remarks

Since 1994, there have been close to 300 publications appearing each year that mention the term mismatch repair. Correspondingly, this field is becoming very difficult to review objectively and I would like to give this as a reason for not having cited a great many interesting papers. In the preceding paragraphs, I have tried to discuss those experiments that address some of the key issues in the MMR field that have stimulated me personally into thinking about the molecular mechanism involved in mismatch correction and that have helped me to define where the shortcomings in our knowledge of the process lie. We have yet to identify several of the missing players in the MMR game, before we can reconstitute the system from its individual components such as was achieved in E.coli (Lahue et al., 1989). We need to know more about proteinprotein and protein-DNA interactions; crystallization efforts should be stepped up, but much can be learned also from missense mutations identified in HNPCC families and from mutagenesis studies. By learning more about MMR individual proteins in the process. The involvement of MMR in drug resistance is of great clinical relevance and we need to understand how these mechanisms operate, with the hope of finding agents that kill MMR-deficient cells more effectively. Mismatch repair proteins have been reported to be involved also in other DNA metabolic pathways such as transcription-coupled nucleotide excision repair (NER) (Mellon and Champe, 1996; Mellon et al., 1996; Leadon and Avrutskaya, 1998), where we currently have no mechanistic insights as to how the MMR and NER pathways might interact. It is of further clinical interest to identify the cause(s) of the inactivation of the wild-type allele of the MMR gene mutated in HNPCC kindreds and why this happens predominantly in the colonic epithelium. Epigenetic inactivation has been shown to play a role here (Kane et al., 1997; Veigl et al., 1998), but there must be also other mechanisms at work. Of interest here are experiments carried out by Meuth and collaborators (Richards et al., 1997), who showed that some MMR-deficient cells acquire a stronger mutator phenotype under conditions of restrained growth, such as might be found in the context of a tumour. This phenotype is reminiscent of the bacterial 'adaptive mutation' phenomenon described several years ago (for a review see Foster, 1998). Another phenomenon that requires further study is the role of the immune system in HNPCC. Frameshift mutations give rise to many aberrant proteins, which are displayed on MHC class I receptors as nonself-peptides. It appears that MMR-deficient cells avoid elimination by the immune system through abolishing expression of the MHC receptors (Bicknell et al., 1996). How do these cells escape elimination by the natural killer cells?

malfunction, we can learn more about the function of the

These and many more questions require answers before we can say that we understand MMR. It is my hope that better knowledge of this important DNA repair pathway will help us devise better ways of dealing with cancers associated with MMR deficiencies, or better still, for preventing these malignancies altogether.

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