The Liaison of Isotype Class Switch and Mismatch Repair: An Illegitimate Affair

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The Ig class switch allows the expression of a V region with new C\text{H} regions associated with various effector functions (for reviews, see references 1 and 2). Switch recombination (SR) occurs by an intrachromosomal deletion process in which the intervening genetic material between the switch (S) regions is excised as a circle. The recombination breakpoints are located within S regions but not at consensus recombination signal sequences, suggesting that SR is not site-specific recombination (3). Switch junctions do not contain long stretches of homology, arguing that homologous recombination does not contribute to this process. Thus, SR is unique in that it does not fall neatly into one of the well-defined categories of recombination.

Results indicating an important role for mismatch repair (MMR) proteins in Ig class switch, published in a recent issue of EMBO (European Molecular Biology Organization) Journal (4) and in this issue of The Journal of Experimental Medicine (5), serve to significantly augment our understanding of the long-elusive molecular mechanism of Ig SR.

A model for the mechanism of SR was previously proposed in which heteroduplex DNA was located at DNA ends in recombination intermediates (6). Successful completion of the switch reaction required removal of the nonhomologous ends. In a perspicacious search for activities that mediate processing of heteroduplex DNA ends in SR, Schrader et al. (5) analyzed mice deficient for the MMR proteins M.sh2, Pms2, Mlh1, and the compound knockout of Pms2/Mlh1 for their ability to support isotype switching in vitro. Extensive analysis of proliferation by triitated thymidine uptake and of cell cycle by propidium iodide staining and flow cytometry demonstrated no difference between MMR-deficient and normal splenic B cells. MMR-deficient splenic B cells stimulated with LPS and the appropriate lymphokines displayed a 35–75% reduction in isotype switching to IgG3, IgG1, IgG2b, and IgA. This evaluation was based on cell surface staining for switched isotypes and was confirmed by digestion-circularization PCR. It is striking that the degree of isotype switch reduction is partial and somewhat variable among the isotypes.

In a screen for the effects of the MMR protein M.sh2 on somatic hypermutation, it was noted that mice deficient in M.sh2 could make good IgM but poor IgG responses to T cell-dependent antigen, suggesting that SR might be selectively affected (7). Ehrenstein and Nueuberger (4) have now systematically examined this issue in M.sh2-deficient mice to find that in both the antigen-specific T cell-dependent response to (4-hydroxy-3-nitrophenyl)acetyl-chicken \(\gamma\)-globulin (NP-CG) and the T cell-independent response to NP-Ficoll, the IgG response was diminished while the IgM response was intact. Analysis of the LPS or the LPS plus IL-4 response in B cells derived from mice deficient for M.sh2 showed reduced switching to IgG3 or to IgG1, respectively, whereas proliferative responses were normal (4). Similar to the study by Schrader et al. (5), these findings reveal a partial reduction of isotype switching. The diminished response to T-independent antigen strongly suggests that this effect is intrinsic to B cells. In another recently published study of the immune response in M.sh2-deficient mice, it was found that IgG expression was diminished but not abolished in antigen-specific T-dependent and -independent responses, whereas the LPS response was intact (8). Detection of an LPS response in M.sh2-deficient B cells does not agree with the results of Schrader et al. (5) and Ehrenstein and Nueuberger (4), and may be related to the method used to prepare splenic B cells for in vitro culture.

Classically, MMR maintains genetic stability by correcting mismatched nucleotides that arise from DNA replication errors, DNA damage, and genetic recombination (9). In Escherichia coli, the MutS, MutL, and MutH proteins provide the unique functions of mismatch recognition and DNA incision that are required to initiate bidirectional strand-specific repair (10). Homologues of MutS have been shown to function similarly in Saccharomyces cerevisiae and in humans. Both mice and humans have at least five homologues of MutS, termed Mut2-6. MMR proteins, most notably M.sh2, bind to mismatched basepairs (11), heteroduplex loops (11), Holliday junctions (12), and other branched intermediates (13). What possible relationship can there be between isotype switching and MMR pathways? To address this question, we must digress for a moment into consideration of the mechanism of isotype class switch.

There has been a sneaking suspicion for the last several years that nonhomologous end joining (NHEJ), also called illegitimate recombination, mediates Ig class switching. Illegitimate recombination in mitotic cells is stimulated by DNA damage, leading to the formation of double strand breaks (DSBs). DSB repair is a process characterized by recombination junctions with distinct features (14, 15) and the involvement of specific proteins in both yeast and mammalian cells (16, 17). In some cases, DSBs repaired by...
NHEJ undergo blunt end ligation, which uses no homology between the ligated ends. More frequently, the recombination junctions contain microhomologies of one to five nucleotides and short deletions of genetic material. In mammalian cells, 97% of the end-joining events are within 15 nucleotides of the original broken end (15). Repair of a site-specific DSB is associated with introduction of mutations in the flanking DNA (18). In murine cells, DSBs are frequently subjected to extensive single strand 5'→3' or 3'→5' exonuclease digestion during NHEJ (19). The gap is thought to be resynthesized, resulting in little net loss of genetic material (19). The resynthesis of DNA by an error-prone DNA polymerase could introduce nucleotide substitutions which would be detected as mutations in the regions flanking the DSB.

The recombination junctions in composite S/S regions conform to the classical features associated with DSB repair by NHEJ. There is either no homology or microhomologies of one to eight nucleotides at the recombination breakpoints for S/S composite genes (3). Mutations flanking the switch junctions have been frequently observed in recombinant S regions (3, 20, 21). It has been difficult to assess the degree of deletion at broken DNA ends since the precise locations of most DSBs in S regions are not known. However, if DSBs occur at specific sites in the S region tandem repeat, the process that leads to short deletions at the DNA ends is predicted to result in clustered recombination breakpoints for switch junctions. Clustered recombination breakpoints in switch junctions of Sμ/S3, Sμ/S1e, or Sμ/S2b have been reported (21, 22). Moreover, a mitogen-inducible DSB found in normal splenic B cells in Sμ3 DNA was located in the region of the clustered breakpoints (23). Taken together, these findings indicate that switch junctions contain the characteristic features associated with DSB repair by NHEJ.

NHEJ is a well-regulated process mediated by DNA-dependent protein kinase (DNA-PK) (16, 17). DNA-PK is composed of the DNA-PK catalytic subunit (DNA-PKcs) and the Ku70/Ku80 heterodimer, which has DNA end-binding activity. The finding of DSBs in Sμ3 DNA and the dependency of SR on the DSB repair proteins Ku80 (24), Ku70 (25), and DNA-PKcs (26) strongly suggest that isotype switching occurs through an NHEJ mechanism. Alternatively, it may be that DSBs in S DNA are not related to the switching process but are normally repaired in a Ku-dependent pathway. Thus, viability of Ku-deficient B cells is poor in the presence of DSBs at S regions, making it appear as though SR is dependent on DNA-PK. Nonetheless, both the distinctive DNA sequence at switch junctions and the identification of a role for DSB repair proteins in SR are consistent with the tentative conclusion that isotype switching is mediated by NHEJ.

The key to understanding a possible role for MMR in SR requires consideration of the physical structure of the DNA ends that are involved in an NHEJ transaction. I begin with the assumption that blunt end DSBs, observed in S DNA (23), are indeed intermediates in the SR reaction (Fig. 1 A). There are two pathways in which a blunt DSB may be processed for use in NHEJ. First, the broken end may be directly ligated (Fig. 1 B). This is presumed to be an inefficient process, since these types of recombination products are relatively infrequent in populations of NHEJ events (15). Second, the broken end may be chewed back along one strand in a 5'→3' (as shown in Fig. 1 C) or in the 3'→5' direction (not shown). In some cases, microhomology between the recombinogenic ends will occur precisely at the DNA termini (Fig. 1 D). These events use an error-prone polymerase to fill in the residual gaps (Fig. 1 E). In other cases, short homologies will be located at more internal locations, leaving heteroduplex DNA at the ends (Fig. 1 F). The removal of the heteroduplex ends is postu-
lated by both Schrader et al. (5) and Ehrenstein and Neuberger (4) to be MMR dependent (Fig. 1 G), and must be followed by gap fill-in. Precedent has been established for involvement of the MMR proteins Msh2 and Msh3 in the removal of heteroduplex ends during DSB repair by homologous recombination (13).

The NHEJ model predicts that isotype switching will be reduced but not abolished by mutations in the MMR pathways, since only those recombination intermediates containing heteroduplex DNA require MMR activity for resolution. Moreover, the frequency at which microhomologies are found at DSB termini may be a function of S sequence. The observations of Schrader et al. (5) and Ehrenstein and Neuberger (4) demonstrating a partial reduction of SR in the absence of MMR and variation in the severity of this reduction as a function of isotype fit remarkably well with the NHEJ model for SR.

The NHEJ model for SR makes a second provocative prediction. Initial DSB formation may be intrinsically biased to consensus sites, as observed previously (23). In MMR deficient mice, heteroduplex DNA ends cannot be processed used and significantly informs our understanding of the molecular mechanism of the Ig class switch.

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