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Shifts in targeting of class switch recombination sites in mice that lack μ switch region tandem repeats or Msh2

Irene M. Min, Lisa R. Rothlein, Carol E. Schrader, Janet Stavnezer, and Erik Selsing

The mechanisms that target class switch recombination (CSR) to antibody gene switch (S) regions are unknown. Analyses of switch site locations in wild-type mice and in mice that lack the Sμ tandem repeats show shifts indicating that a 4–5-kb DNA domain (bounded upstream by the Iμ promoter) is accessible for switching independent of Sμ sequences. This CSR-accessible domain is reminiscent of the promoter-defined domains that target somatic hypermutation. Within the 4–5-kb CSR domain, the targeting of S site locations also depends on the Msh2 mismatch repair protein because Msh2-deficient mice show an increased focus of sites to the Sμ tandem repeat region. We propose that Msh2 affects S site location because sequences with few activation-induced cytidine deaminase targets generate mostly switch DNA cleavages that require Msh2-directed processing to allow CSR joining.

Antibody gene class switch recombination (CSR) occurs between switch (S) regions that are located upstream of each CH gene and contain tandem repeat sequences that are often G-rich on the nontranscribed DNA strand (1, 2). The unusual nature of S region tandem repeat (Sμ.TR) sequences suggests that they play a role in CSR. However, mice lacking Sμ tandem repeats (Sμ.TR−/−, previously designated as ΔSμ; reference 3) show only two- to threefold reductions in isotype switching, which indicates that sequences outside of the Sμ.TR can target the switching mechanism (3). Larger deletions of Sμ sequences have greater defects in switching (4), and an extensive deletion of Sy1 shows almost complete abrogation of γ1 switching (5), indicating that S region sequences are critical for CSR. The mechanisms that target CSR to S regions are unknown.

Activation-induced cytidine deaminase (AID) and proteins involved in nonhomologous end joining are critical for CSR activity. AID deamination of deoxycytidine residues appears to initiate events that result in S region DNA cleavages (6, 7), whereas nonhomologous end joining proteins may be involved in the religation of broken S region DNA ends (8–10). CSR also involves the uracil DNA glycosylase protein, possibly for the generation of abasic sites from AID-generated dU–dG DNA mismatches (11). DNA mismatch repair (MMR) proteins are also important for CSR (12, 13), but their roles are unclear. Studies of S sites located within a portion of Sμ showed that the absence of the Msh2 MMR protein caused CSR to focus on GAGCT/GGGGT sequences (12). Furthermore, the presence of the Sμ.TR region is critical for class switching in Msh2-deficient mice, suggesting that Msh2 affects the sequences that are targeted or selected for CSR (14). Msh2 is also important for the low levels of CSR found in uracil DNA glycosylase–deficient mice, suggesting an Msh2 role in some DNA cleavages that initiate CSR (15).

We have analyzed the locations of S sites in two mutant mouse strains that lack either the Sμ.TR region or the Msh2 protein to determine how these mutations affect CSR targeting. S site locations in wild-type and Sμ.TR−/− mice show that CSR events occur within a 4–5-kb DNA region located downstream of the Iμ promoter and provide the first indication that CSR is targeted to a fixed-length “domain” that is linked to the location of a transcriptional promoter. Comparing S site locations in wild-type with Msh2−/− mice shows that the absence of Msh2 focuses CSR sites on the Sμ.TR region. This finding indicates that...
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RESULTS AND DISCUSSION

Upstream S sites are not affected by SpμTR deletion

We used several digestion-circularization (DC)-PCR assays to analyze CSR sites in stimulated splenic B cells. To quantify Spμ/Sy1 CSR sites located upstream of the SpμTR region in wild-type and SpμTR−/− mice, a BsrGI digestion was added to the standard EcoRI DC-PCR assay (16) before DNA circularization (Fig. 1 a). Only Spμ/Sy1 junctions upstream of the Spμ BsrGI site will lack the BsrGI site and generate EcoRI fragments that can be self-ligated and PCR amplified. Comparing band intensities in EcoRI DC-PCR with those in EcoRIBsrGI DC-PCR assays allows the percentages of junctions located upstream or downstream of the Spμ BsrGI site to be calculated.

Genomic DNAs isolated from SpμTR−/− or wild-type B cells were analyzed using the two assays (Fig. 1 b). Based on the EcoRI DC-PCR results, total levels of Spμ/Sy1 CSR in SpμTR−/− B cells were reduced about twofold relative to the wild type, which was consistent with previous findings (3, 14). However, analyses of Spμ/Sy1 CSR upstream of the BsrGI site (Fig. 1 b) showed that these were the same in SpμTR−/− and wild-type B cells (Fig. 1 c). These results indicate that, in this upstream region, CSR events are not influenced by the SpμTR deletion in nearby downstream regions.

Switch recombinations in SpμTR−/− mice shift to downstream JH-Cμ regions

To further locate S sites within the JH-Cμ intron, downstream CSR events measured in HindIII and XbaI DC-PCR assays were compared with the total number of switch events measured by the EcoRI DC-PCR assay (Fig. 2, a and b). Because the three DC-PCRs can have different efficiencies, we used a control plasmid (pSpμ/Sy3) that contained Spμ and Sy3 sequences together with the three enzyme sites to normalize the DC-PCR assays. The frequencies of CSR sites within different segments of the JH-Cμ region were determined by comparison with standard curves generated using pSpμ/Sy3.

We first assessed total Spμ/Sy3 recombinations by EcoRI DC-PCR assays (Fig. 2 a) and used DC-PCR quantitation of the Rag-1 gene to normalize the content of genomic DNA from different samples. The total frequency of Spμ/Sy3 switch events in wild-type B cells was estimated to be ~7.3% (Fig. 2 b). This frequency from real-time DC-PCR matches the levels of in vitro γ3 switching as assessed by flow cytometry (14). Spμ/Sy3 CSR in SpμTR−/− mice was reduced about threefold relative to the wild type (Fig. 2 b). Of the total Spμ/Sy3 junctions in SpμTR−/− mice, ~67% were located downstream of the HindIII site located at the 3′ end of the SpμTR region (Fig. 2 a). This high frequency of junctions downstream of H1 in SpμTR−/− mice is in striking contrast to the negligible number of Spμ/Sy3 sites located within the same sequence in wild-type mice (<1% of all switch events; Fig. 2 b).

Of the Spμ/Sy3 recombinations in SpμTR−/− mice located downstream of the H1 site, about half occurred downstream of the XbaI site located near the 5′ end of the Cμ exons (Fig. 2 b). Comparisons of EcoRI DC-PCR with XbaI DC-PCR assays indicated that ~30% of Spμ/Sy3 recombinations in SpμTR−/− mice occurred at sequences downstream of the XbaI site.

For wild-type and SpμTR−/− mice, we compared the locations of CSR sites determined by DC-PCR with the locations of CSR events in hybridoma panels (3). For wild-type mice, 23 hybridomas show five (22%) CSR sites upstream of the SpμTR, 17 (78%) sites within the SpμTR, and no sites downstream of the SpμTR. DC-PCR analyses of wild-type CSR events show 17% upstream, 82% SpμTR, and 1% downstream. For CSR sites in SpμTR−/− mice, 10 hybridomas show three (30%) upstream of the H1 site and seven (70%) downstream of H1, whereas DC-PCR shows 33% upstream and 67% downstream. Thus, although the hybridoma panels are small and will miss CSR events that result in nonfunctional Ig proteins,
the comparisons show a good correlation between the distributions of CSR sites as determined by the two methods.

The CSR site frequency between the HindIII and XbaI sites might be slightly underestimated because of the H2 HindIII site in the Cμ segment (Fig. 2 a). For SpμTR−/− mice, CSR events occurring downstream of H2 were estimated using the HindIII-circularized B cell DNA samples. Among SpμTR−/− mice, CSR sites located downstream of H1, only ~7% were located downstream of H2 (Fig. 2 c), assuming that both HindIII DC-PCR assays have equal efficiencies. An examination of the threshold cycles of the two HindIII PCR assays indicated that the efficiencies differ by less than a factor of two (not depicted). Thus, although the majority of switch recombinations in SpμTR−/− mice are skewed toward downstream JH-Cμ intron sequences, there are few junction sites downstream of the Cμ exons.

Lack of the MMR Msh2 protein shifts the targeting of CSR to focus on the SpμTR region
The role of Msh2 in the CSR process is unclear. Previous analyses of SpμTR−/−:Msh2−/− mice have shown that the SpμTR region is critical for isotype switching when Msh2 is lacking (14). These studies suggested that Msh2 might be more important for switching in the sequences flanking the SpμTR region than within the SpμTR itself. To directly test this model, S site locations were determined in Msh2−/− mice that retain the SpμTR region.

CSR sites throughout the JH-Cμ region were analyzed by DC-PCR in Msh2−/− mice. CSR sites upstream of the SpμTR in Msh2−/− mice were decreased about sixfold relative to the wild type, whereas switching within the SpμTR was only reduced by a factor of 1.4 (Fig. 3, a and b). The fraction of CSR events located upstream in Msh2−/− mice was significantly different from the wild type (two-sided P < 0.0001 by χ² test; Fig. 3 b). Very few CSR events occurred downstream of the SpμTR in both Msh2−/− and wild-type mice (Fig. 3 C), and we cannot assess whether any specific reductions occur. Within the regions that exhibited the majority of CSR activity, however, our results show a shift of CSR site focus into the SpμTR region in Msh2−/− mice (Fig. 4) and support the model that Msh2 is more important for CSR in non-SpμTR sequences and less important for CSR within the SpμTR.

Figure 2. SpμTR−/− mice have increased levels of CSR events targeted to downstream sequences. (a) DC-PCR measuring CSR junction sites downstream of the SpμTR region. HindIII and XbaI sites downstream of the SpμTR region were used in DC-PCR assays to measure CSR junctions in sequences downstream of each site. Total CSR sites were determined by EcoRI DC-PCR (E, EcoRI; H1 and H2, HindIII; X, XbaI). Primer pairs used for the different DC-PCR assays are indicated by small arrows of different types. (b) Wild-type and SpμTR−/− CSR events measured by EcoRI, HindIII, and XbaI real-time DC-PCR. Copy numbers for CSR events were derived from pSpμSy3 standard curves. Rag-1 PCR normalized CSR events relative to Rag-1 copy number. Mean values and SEM are from seven independent experiments with four mice per group. (c) Percentage of SpμTR−/− switch events downstream of the H2 relative to events downstream of H1 (set to 100%). Mean values and SEM are from three independent experiments per group.

Figure 3. CSR within sequences upstream of the SpμTR are reduced in Msh2−/− mice. (a) Twofold dilutions of genomic DNAs were assayed by SpμSy1 EcoRI DC-PCR and EcoRI–BsrGI DC-PCR. Samples were normalized using nAChR DC-PCR. (b) CSR sites in wild-type or Msh2−/− mice measured relative to total wild-type CSR (set to 100%). Mean values and SEM of at least four independent experiments are presented. (c) CSR events in Msh2−/− mice measured by EcoRI, HindIII, and XbaI real-time DC-PCR and compared with wild-type mice. Mean values and SEM are from six independent experiments with two Msh2−/− mice and seven independent experiments with four wild-type mice.
Figure 4. CSR junction sites and AID target sites within different segments of the JH-Cμ intron. Histograms of CSR site, WRC motif, and WGCW motif frequencies are shown for different segments of the JH-Cμ region. (a) Comparisons of wild-type with Msh2−/− mice that both have wild-type Sμ regions. (b) Comparisons of SμTR−/− mice with a shortened Sμ region. DNA segments are drawn in proportion on the x axis. Areas of boxes in the histogram are proportional to frequencies in each segment, resulting in box heights that indicate relative recombinational frequencies per nucleotide. Boxes for mutant mice are drawn to scale relative to wild-type mice. Switching reductions in SμTR−/− varied slightly in different DC-PCR assays and were averaged. Junction site distributions in each segment were assumed to be uniform. Wild-type CSR frequencies are set to 100%; other CSR frequencies are normalized to this total. Total WRC and WGCW motif frequencies were set to 100%; each mouse strain. The SμTR is only partially sequenced (available from GenBank/EMBL/DDBJ under accession no. AC073553). Densities of motifs are assumed conserved throughout the SμTR. The total numbers of each motif in the Cμ region are indicated in the top right of each graph.

Patterns of S sites in wild-type and SμTR−/− mice suggest a 4–5-kb switching domain

SμTR−/− mice exhibit shifts of S site locations into downstream JH-Cμ intron sequences that show almost no CSR in wild-type mice. These shifts indicate that, in wild-type mice, there is a mechanism that restricts switching in these downstream regions because the same sequences can clearly support numerous switch events when the SμTR region is deleted. This mechanism does not appear to reflect competition between different sequences for the recruitment of the CSR machinery because the sequences upstream of the SμTR, which can clearly support CSR even in wild-type mice, show no increases in CSR when the SμTR is absent. Based on S site locations in wild-type mice, it appears that a 4–5-kb-long region downstream of the Cμ promoter is accessible for CSR in stimulated B cells. The pattern of S sites in SμTR−/− B cells suggests that a similarly sized region is also accessible for CSR, but, because of the SμTR deletion, this region now extends into Cμ sequences (Fig. 5).

The mechanism that limits the CSR domain is not known. The Cμ promoter appears to be the upstream boundary of most CSR events (4, 17) and may localize the upstream limit of the CSR domain (Fig. 5). The CSR domain might be explained by a transcriptional-tracking model similar to models for the targeting of somatic hypermutation (18) in which switch cleavage factors are carried with RNA polymerase during the transcription process, but dissociate after a distance. AID interactions with RNA polymerase (19) provide some support for such a transcription-based model.

The CSR-accessible DNA domain could also involve a chromatin structure that regulates the switching machinery, similar to the ability of chromatin structure to regulate V(D)J recombination (20, 21). Studies of B cells undergoing somatic hypermutation indicate that mutating V(D)J segments are found in chromatin where the histones H3 and H4 are hyperacetylated relative to the C-region of the same gene (22). For switching, however, relationships between CSR and S region H3/H4 hyperacetylation are complex, suggesting that other modifications or protein factors might be needed for CSR targeting (23). In pro-B cells, dimethylation of histone 3 at lysine 4 in the JH-Cμ region peaks around Cμ with a rapid drop off within 2 kb of this central peak (21). Thus, the dimethylation of histone 3 at lysine 4 modification rises rapidly in the region just downstream of the SμTR that forms the 3′ boundary of the switching domain and could be involved in limiting CSR accessibility.

In wild-type mice, CSR sites correlate with the distribution of WRC motifs (Fig. 4 a) that are known sites for AID deamination on single-stranded DNAs (24, 25). In SμTR−/− mice, however, CSR sites do not correlate with WRC motifs (Fig. 4 b). The HindIII–XbaI focus of SμTR−/− CSR sites could reflect the high concentration of nontranscribed G residues in this region (3). Such G-rich sequences promote RNA–DNA R-loops in S regions and might provide single-stranded DNA targets for AID activity (5, 26).

Mice with a larger Sμ-region deletion show more drastic CSR reductions and have Sμ sites mostly constrained to the Cμ exon region (4). Sequence elements important for the
4–5-kb CSR domain proposed in Fig. 5 may be disrupted by the larger Sµ deletion. Identifying sequences that establish the CSR domain might distinguish transcriptional and chromatin structure targeting models and lead to the identification of targeting factors. S regions clearly contain sequences important for CSR because replacing S region sequences with a random DNA sequence does not provide for CSR (27).

Msh2 may be important for processing “frayed” switch DNA cleavages

Our results show that S sites in Msh2−/− mice are strongly focused into the Sµ TR region, indicating that CSR events within Sµ TR sequences are much less dependent on Msh2 than are events within non-Sµ TR sequences. The Msh2 activity that leads to this sequence-specific effect is not clear. Msh2 is known to recognize base pair mismatches, and recent studies suggest that Msh2 recognition of G–U might lead to S region DNA breaks (15). However, Msh2 G–U recognition appears to play a minor role in switch cleavage (15), and there is no apparent reason for this G–U recognition activity to affect switching outside of Sµ TR sequences to a greater extent than within the Sµ TR.

The shifts of CSR sites in Msh2−/− mice are consistent with a model that we have previously proposed in which Msh2 processes DNA “flaps” found predominantly on cleavage ends generated from sequences outside of the Sµ TR (14). This model suggests a postcleavage end-processing function for Msh2 that converts staggered single-stranded breaks to the double-stranded breaks required for CSR, in addition to its role in initiating some switch cleavages.

We analyzed the JH-Cµ sequence to determine whether motifs associated with CSR might account for the S site distribution in Msh2−/− mice. WRC motifs are preferred sites for AID (24, 25), and the WGCW motif is a site where AID activity on opposing strands could provide a double-stranded cleavage with a single base overhang (24). WRC motifs within the JH-Cµ intron correlate well with distributions of CSR sites in the 4–5-kb Sµ TR domain for wild-type mice (Fig. 4 a). For Msh2−/− mice, on the other hand, CSR sites correlate better with WGCW motifs (Fig. 4 a). CSR downstream of the Sµ TR is rare in both wild-type and Msh2−/− mice, so that Msh2 effects are not discernable. However, Sµ TR−/− mice show numerous S sites in this downstream region and previous analyses have shown that Sµ TR−/−: Msh2−/− mice show large decreases in all CSR events (14). Thus, Msh2 is also important when CSR occurs downstream of the Sµ TR, consistent with the paucity of WGCW sites within these sequences (Fig. 4 a). CSR sites in Sµ TR−/−:Msh2−/− mice also show a significant focus on WGCW motifs (two-tailed P = 0.0043 by Fisher’s exact test) relative to CSR sites in Sµ TR−/− mice (14), emphasizing the correlation between WGCW motifs and CSR sites when Msh2 is not present. Furthermore, the number of WGCW motifs in the wild-type Sµ region is about half the number of WRC motifs in the same region (Fig. 4 a). Perhaps the twofold reduction in CSR observed between wild-type and Msh2−/− mice correlates roughly with the fraction of cleavage sites that cannot complete CSR because of frayed ends.

The involvement of Msh2 in processing S region breaks would suggest that Msh2 alters switch targeting by affecting the selection of switch breaks that can undergo CSR. Thus, Msh2 has an important role to broaden the types of DNA sequences that can successfully participate in CSR. Analyses of Sµ TR−/− mice crossed with mice deficient in other MMR proteins, such as Exo1, Mlh1, and Pms2, should indicate whether these proteins are involved in the same processes of the CSR mechanism that are affected by Msh2.

MATERIALS AND METHODS

Mice and cell culture. Sµ TR−/− and Msh2−/− mice were used in these studies. Animal experiments were approved by Tufts University Institutional Animal Care and Use Committee. Splenic B cells were cultured with 50 μg/ml LPS (Sigma-Aldrich) with or without 0.3 ng/ml anti-ß-dextran, or with LPS and IL-4 (800 U/ml) as described previously (14).

DC-PCR. DC-PCR was performed as previously described (3, 14). Circularized DNAs were diluted twofold, and PCR was amplified as follows. For switch events located upstream of the Sµ TR, DNAs were digested with BglI and EcoRI before circularization. Sµ/Sy1 DC-PCR products were amplified and phosphorimager quantification was normalized using the nicotinic acetylcholine receptor (nAChR) gene (16). BglI digestion was monitored by PCR of the BglI site using primers Sµ out (14) and BglI_bk (5′-TCATTACTGTGGCTGGAGAG-3′). Switching upstream was quantitated by comparing EcoRI-BsrGI DC-PCR with EcoRI DC-PCR products. For switch events located downstream of the Sµ TR, genomic DNAs were digested with EcoRI, HindIII, or Xbal, and circularized for real-time PCR using fluorescent SYBR green dye (Applied Biosystems). The pSµ/Sy3 control plasmid provided standard curves for each real-time DC-PCR. Rag-1 copy numbers were measured by real-time PCR using Rag-1 plasmid standards (Rag-1 plasmid provided by E. Marchlik, Tufts University, Boston, MA). Gel electrophoresis of PCR products and melting temperature analyses were used to confirm the specificity of real-time PCR amplifications. Total Sµ/Sy3 junctions were measured in an EcoRI DC-PCR assay using published primer pairs DC-µ2 and DC-γ3 (28).

Switch events downstream of the H1 site (Fig. 2) were measured by HindIII DC-PCR using primers H3-IM (5′-CTTTACACCATGATCACAT-3′) and H3-G3F (5′-GATAGGACAGATGGAGCAGTTACAGA-3′). HindIII DC-PCR measured switch events downstream of the Xbal site located 5′ of the Cµ exon (Fig. 2 a) using primer pairs XBA-IM (5′-GAACGACTTGGGAGCCCTCA-3′) and XI-G3F (5′-GCTGAGGAGGACCTGGTGT-3′). HindIII DC-PCR measured S junctions downstream of the H2 site (Fig. 2 c) using primers CMH3-R (5′-GGGGAGGACCTGAGTCAAG-3′) and H3-G3F.
Statistics. χ² tests were used to assess the significance of Msh2⁻/⁻ upstream CSR differences. Fisher’s exact test was used to assess the significance of differences in CSR at WGCM motifs between SpμTR⁻/⁻:Msh2⁻/⁻ and SpμTR⁻/⁻ mice.

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