Adaptive immunity depends on V(D)J recombination to assemble antigen receptor genes from their component gene segments during B and T cell development. The recombinase introduces double strand (ds) DNA breaks at recombination signal sequences (RSSs) that flank rearranging gene segments. An RSS consists of conserved heptamer and nonamer sequences separated by a spacer of conserved length (either 12 or 23 bp). The DNA break is made precisely at the junction between the RSS heptamer and the gene segment. These broken DNA molecules are then joined with the assistance of the nonhomologous end joining (NHEJ) family of dsDNA break repair proteins to form coding and signal joints (Fig. 1A) (1). Mutations in genes encoding NHEJ proteins greatly diminish the frequency of productive V(D)J recombination and, when combined with mutations in p53, cause striking genomic instability and cell transformation, resulting in leukemia and lymphoma (2).

The V(D)J recombinase catalyzes DNA transposition and translocation both in vitro and in vivo. Because lymphoid malignancies contain chromosomal translocations involving antigen receptor and protooncogene loci, it is critical to understand the types of “mistakes” made by the recombinase. Using a newly devised assay, we characterized 48 unique TCRβ recombination signal sequence (RSS) end insertions in murine thymocyte and splenocyte genomic DNA samples. Nearly half of these events targeted “cryptic” RSS-like elements. In no instance did we detect target-site duplications, which is a hallmark of recombinase-mediated transposition in vitro. Rather, these insertions were most likely caused by either V(D)J recombination between a bona fide RSS and a cryptic RSS or the insertion of signal circles into chromosomal loci via a V(D)J recombination-like mechanism. Although wild-type, p53, p53 x scid, H2Ax, and ATM mutant thymocytes all showed similar levels of RSS end insertions, core-RAG2 mutant thymocytes showed a sevenfold greater frequency of such events. Thus, the noncore domain of RAG2 serves to limit the extent to which the integrity of the genome is threatened by mistargeting of V(D)J recombination.

Adaptive immunity depends on V(D)J recombination to assemble antigen receptor genes from their component gene segments during B and T cell development. The recombinase introduces double strand (ds) DNA breaks at recombination signal sequences (RSSs) that flank rearranging gene segments. An RSS consists of conserved heptamer and nonamer sequences separated by a spacer of conserved length (either 12 or 23 bp). The DNA break is made precisely at the junction between the RSS heptamer and the gene segment. These broken DNA molecules are then joined with the assistance of the nonhomologous end joining (NHEJ) family of dsDNA break repair proteins to form coding and signal joints (Fig. 1A) (1). Mutations in genes encoding NHEJ proteins greatly diminish the frequency of productive V(D)J recombination and, when combined with mutations in p53, cause striking genomic instability and cell transformation, resulting in leukemia and lymphoma (2).

Strong evidence exists that the lymphoid V(D)J recombinase evolved from an ancient transposase (3). Consistent with this idea, genes encoding the lymphoid-specific components of the recombinase, RAG1 and RAG2, are physically linked in the genome, convergently transcribed, and lack introns in their coding regions (4). Certain aspects of the biochemistry of V(D)J recombination also resemble a transposition reaction. RAG1 contains, and its function depends on, a catalytic triad of amino acids that is common to many transposases, the D-D-E motif (5). RSSs are recognized and cleaved in a pairwise fashion, generating two blunt and two hairpin ends, as is the case in Tn10 and Hermes transposition. In vivo, the two broken RSS ends are most often ligated to one another, generating an episomal signal joint (Fig. 1A); however, infrequently, broken RSSs can rejoin to coding ends, thereby forming what are called hybrid (Fig. 1A, bottom) and open-and-shut joints (6). In vitro, however, RSS ends efficiently transpose, generating a variety of strand-transfer products (7–9). A truncation mutant of RAG2 (core-RAG2, consisting of amino acids 1–383 out of the 527 amino acids of the full-length protein) results in a greatly increased frequency of transposition in vitro (10–12) and hybrid joining in vivo (13).
Understanding how these side reactions of V(D)J recombination are suppressed is of great significance given the frequent observation of chromosomal translocations and deletions in lymphoid malignancy (14).

V(D)J recombination can contribute to genomic instability and malignancy in several ways (15, 16). First, the recombinase can recognize so-called “cryptic” RSSs in nonantigen receptor loci (defined as a DNA sequence identical to a known functional RSS heptamer) and use them together with either bona fide RSSs or other cryptic RSSs to catalyze translocations or deletions. Previous studies revealed that the ability of cryptic RSSs associated with certain protooncogenic loci to undergo V(D)J recombination is within 10–100-fold that of a proper RSS (17). In addition, RAG proteins have the ability to recognize, cleave, and rearrange certain DNA sequences that do not resemble RSSs, but have the tendency to contain non–B-form DNA. A particularly clear example of this was reported in the major breakpoint region (Mbr) of the Bl-2 locus, which is the most frequently re-arranged oncogene in follicular lymphoma (18). The RAGs can also activate the 3′–OH group on broken RSS ends to attack duplex DNA, generating a dsDNA break in the accidental target of this reaction (9). Recently, it was reported that signal joints, previously considered inert, can reinsert into Ig or TCR loci, as well as into cryptic RSSs (19). Finally, as occurs in vitro, the RAGs can catalyze the transpositional insertion of an RSS-ended fragment into genomic DNA (Fig. 1 B). Such RAG-mediated transposition events are characterized by short, target-site sequence duplications. Only three biological examples of this have been published in the literature. In each case, an excised, signal-ended fragment from the TCRα locus was inserted into an intron within the hprt locus (20, 21). Using a retroviral reporter construct in a virally transformed pro–B cell line, Reddy et al. reported that the frequency of RSS-fragment transposition in vivo was ~1 in 44,000 recombination events (22).

To determine which of the biochemical activities of the V(D)J recombinase contribute to genomic instability, and to understand the mechanisms that limit RSS fragment transposition and other RSS end insertion events in developing lymphocytes, we developed a methodology to detect and characterize TCRβ locus RSS end insertions, including transposition, in genomic DNA purified from mouse T lineage cells. We applied this assay to wild-type and various mutant murine DNA samples. Our results reveal that the genomes of developing thymocytes are littered with TCR signal end integration events that may contribute to genomic instability, that a high fraction of such events target cryptic RSSs, and that the noncore domain of RAG2 serves to limit these potentially deleterious events.

RESULTS

LM-TECA assay for RSS fragment insertion

We designed the ligation-mediated transferred-end capture assay (LM-TECA) to detect the insertion of a RSS-ended DNA fragment generated during V(D)J recombination into random sites around the genome (Fig. 1 C). High molecular weight genomic DNA is restriction digested, denatured, and subjected to primer extension using a biotinylated primer homologous to a region adjacent to either the Dβ1 RSS (D-side) or the JB1.1 RSS (J-side). Primer extension products are captured using streptavidin-coated paramagnetic beads and ligated to an oligonucleotide linker. The linker-ligated DNA is analyzed by PCR using a linker primer and a RSS fragment primer. Amplified products are purified by gel electrophoresis (Fig. 2) and characterized by DNA sequence analysis (Table I). Once the genomic location of an inserted RSS end has been determined, direct PCR can be used in an attempt to identify the sequence at the other end of the RSS fragment (Fig. 1 D, top) or the other joining product of a balanced translocation (Fig. 1 D, bottom).
LM-TECA amplifies several classes of background events. Most of these are mechanical breaks in purified genomic DNA that are ligated and captured by the assay after initial primer extension. A second source of background events are signal end breaks, which exist in cells undergoing D-to-Jβ rearrangement. Captured signal end PCR products are of known size and can thus be identified on gels and ignored. For assays on the Dβ1-side, a third source of background events result from direct Vβ-to-Dβ1 rearrangements that occur rarely in wild-type animals and 10 to 1,000 time more frequently (unpublished data) in core-RAG2 mutant mice (23). We spent the bulk of our sequencing efforts on capturing Jβ1-sided events to avoid the confounding effects of these direct V-to-D coding joints.

As a negative control for LM-TECA, we used a RAG1−/− mouse expressing a TCRβ chain transgene. Thymocytes from these mice contain large numbers of preT cells lacking RAG-mediated breaks or rearrangements. DNA sequence analysis of 91 uniquely sized Dβ1-side or Jβ1-side PCR products revealed random DNA breaks across this region of the TCRβ locus (unpublished data). A second negative control was designed to address the concern that PCR artifacts might generate presumed RSS insertion fragments, particularly from genomic DNA samples that contain considerable amounts of broken RSS ends. We mixed mouse thymocytes with equal numbers of human HeLa cells and then purified genomic DNA. If LM-TECA results in the generation of artifacts caused by RSS-ended DNA fragments joining with random DNA fragments, we would expect to detect “insertions” of TCRβ RSSs into human DNA. We performed numerous LM-TECA assays on this source of material and never found any mouse–human DNA recombinants; however, we did detect four insertions of TCRβ RSS fragments into mouse DNA (Table I).

Fig. 2 shows a typical set of PCR reactions in which multiple 0.2-μl aliquots from a single Jβ-side LM-TECA of wild-type thymus DNA were assayed. 9 out of the 10 reactions show a 655-nt product that corresponds to the germline TCRβ locus cleaved at the restriction site (filled arrow). This amplified fragment serves as an internal control for the assay. Individual fragments both smaller and larger than the germline fragment can be seen. From these 10 PCR reactions, 2 of the sequenced fragments proved to be signal end insertion events (open arrows). Depending on the sample, between 5 and 10% of sequenced products were RSS insertions (unpublished data), and the remainder were “background” events as noted in the previous paragraph.

Using this strategy, we were able to isolate and characterize a series of Dβ1-side or Jβ1-side RSS insertion events from primary tissues (Table I). Most of the events we detected were found in WT-thymic DNA samples. We obtained one event from total splenic DNA and five others from sorted splenic T cells. We also analyzed thymic DNA from core-RAG2 mutant mice (23) and from several DNA repair–deficient mouse strains (p53, p53 x sad, 53bp1, H2Ax, and ATM) (24–27).

**Characteristics of RSS fragment insertions**

The 48 RSS end insertion events we recovered mapped to 18 different chromosomes (Table I). 36 were located within or near annotated genes, including 6 into the TCRγ locus and 4 into non–RSS-containing regions of the TCRβ locus. 35 of the events contained N regions (nontemplated nucleotides) ranging from 1 to 10 nt in length. Interestingly, 4 of 12 Dβ1 RSS fragment insertions occurred near, but not exactly at, the RSS heptamer, deleting 1, 2, 9, or 12 nt from the RSS end. There was even more alteration of sequence among the Jβ1 RSS end insertions, with only 14 of 36 sequences containing the full heptamer, whereas in the other 22, the break deleted between 2 and 12 nt of the RSS.

Many of the chromosomal targets of RSS-end insertion were at or near RSSs or cryptic heptamers (Table I, underlined sequences). 9 of the 12 Dβ1 RSS end insertions were of this type. Two of these involved TCR Jγ RSSs, whereas the other seven breakpoints occurred within 2 nt of a consensus heptamer. The situation was somewhat different in respect to Jβ1 RSS end targets, with only 13 of 36 sequences targeted near cryptic heptamers and 4 involving bona fide RSSs (TCRβ and γ loci). Cryptic heptamers at target loci were found on either side of the RSS insertion site, suggesting that ligation events could resemble signal, hybrid, or coding joint formation.

**Detecting the second end of an RSS insertion**

For the majority of the Dβ-side or Jβ-side RSS insertion events, we designed a specific direct PCR assay to detect possible insertion of the other end of the RSS-ended fragment into the same target locus (Fig. 1 D, Fig. 3, and Fig. 4). These assays involved nested PCRs with TCRβ-sided and target locus-sided primers. Using this strategy, we were able to identify the “other” end of a total of five events (Fig. S1, available at http://www.jem.org/cgi/content/full/jem.20070583/DC1).
### Table I. Partial DNA sequences of Dβ1/H9252 and Jβ1/H9252 signal end insertions into various chromosomal locations

| Event Name | Geno | Chromosome breakpoint | N and P | Dβ1 Side | Gene | UCSC chr. position |
|------------|------|------------------------|---------|----------|------|------------------|
| **A**      |      |                        |         |          |      |                  |
| E1-4       | WT   | CCCTATGG CAGTTGTCtcactgctc | AATGGCC | CACGGTG ATTCATT | TCR Jy1 | chr:13:18678545 |
| E2-2       | WT   | CCCTATGG CAGTTGTCtcctaaaga | ACC | CACGGTG ATTCATT | TCR Jy1 | chr:16:55739889 |
| E3-4       | WT   | HPCCTATGG CAGTTGTCtcgcagcag | CCG | CACGGTG ATTCATT | Scarb1 | chr:7:69460414 |
| EMAT2      | WT   | ACATCTAGAGGtaagaagcaaga | CACGGTG ATTCATT | BO42698 | chr:1:138847427 |
| EMAT5-1    | WT   | AGCTATGG CAGTTGTCcctcc | CACGGTG ATTCATT | C80913 (NNX3) | chr:7:34900516 |
| E2b-6      | cR2  | GAGCTATGG CAGTTGTCagctgctggc | CACGGTG ATTCATT | lama3 | chr:18:12680543 |
| b196       | WT   | TCCTATGG CACTGTTgcagcagtgg | see note | CACGGTG ATTCATT | hotspot | chr:14:3968210 |
| b154       | WT   | TCCTATGG CACTGTTgcagcagctgg | CACGGTG ATTCATT | Scarb1 | chr:5:124540600 |
| EMAT5-2    | WT   | GAGCTATGG CAGTTGTCgctacagcg | ACCTATGG CAGTTGTC | DEAD box pept 21 | TCR Jy2 | chr:13:18624358 |
| E3-1       | WT   | TCCTATGG CACTGTTgcagcagctgg | CACGGTG ATTCATT | TCR Jy2 | chr:13:18624358 |
| b220       | cR2  | TAGCTATGG CAGTTGTCgctacagcg | CACGGTG ATTCATT | BC096449, unknown mRNA | chr:1:152252917 |
| EL3-1      | WT   | AGCTATGG CAGTTGTCcctcc | CACGGTG ATTCATT | Vh J558 | chr:12:112181483 |
| **B**      |      |                        |         |          |      |                  |
| Jβ1.1.1    |      | TCCTATGG CAGTTGTC | ??????GACCCCGTC | too short to align |
| EL2-4      | WT   | TCCTATGG CAGTTGTC | aatcagtgaatagaATCG | Vh J558 | chr:12:112181482 |
| S3         | WT   | TCCTATGG CACTGTTGC | GACCTGGTATCGTTCTCT | TCR Vx16 | chr:14:48296888 |
| b175       | WT   | TCCTATGG CACTGTTGC | gcgtctcgctagGAGCCAA | histone deac. 7A | chr:15:97821023 |
| S16        | WT   | TCCTATGG CACTGTTGC | gaaaggtgtAAGAAGACTCAAC | AK133119, mRNA | chr:2:6346533 |
| B120       | WT   | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| S31        | WT   | TCCTATGG CACTGTTGC | gcaatgtgatAGCTCGGGCT | TCR yJ1 Coding | chr:13:18678643 |
| b196       | WT   | TCCTATGG CACTGTTGC | cccctgccctggtgc | chr:14:13968211 |
| Jβ1.2.1    |      | TCCTATGG CACTGTTGC | ??????GACCCCGTC | too short to align |
| b108       | WT   | TCCTATGG CACTGTTGC | aatcagtgaatagaATCG | Vh J558 | chr:12:112181482 |
| b183       | WT   | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| b177       | WT   | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| S1         | WT   | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| b14        | WT   | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| S2         | WT   | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| b35        | WT   | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| Jβ1.3.1    |      | TCCTATGG CACTGTTGC | ??????GACCCCGTC | too short to align |
| b154       | WT   | TCCTATGG CACTGTTGC | aatcagtgaatagaATCG | Vh J558 | chr:12:112181482 |
| E1-6       | WT   | TCCTATGG CACTGTTGC | aatcagtgaatagaATCG | Vh J558 | chr:12:112181482 |
| b114       | WT   | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| b126       | WT   | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| c (Events recovered from mutants) |      |                        |         |          |      |                  |
| Jβ1.1.1    |      | TCCTATGG CACTGTTGC | ??????GACCCCGTC | too short to align |
| b75        | H2AX | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| b235       | cR2  | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| E2b-6      | cR2  | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| Jβ1.2.1    |      | TCCTATGG CACTGTTGC | ??????GACCCCGTC | too short to align |
| b220       | cR2  | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| E2a-6      | cR2  | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| b243       | p53  | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| b314       | p53  | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| b346       | p53  | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
| b340       | p53  | TCCTATGG CACTGTTGC | gtaatgtgactgtgCAGTCATG | Plut in Intron | chr:1:134452052 |
Event EL3–1 (Table I), for example, was an insertion of the DB1-to-JB1.1 RSS-ended fragment into a region of the IgHC locus 5′ to a V558-family gene segment (Fig. 3 and Fig. S1). This is somewhat surprising, given the inaccessibility of the IgHC V region to the V(D)J recombinase in developing T cells. In this insertion, the RSS-end identified by LM-TECA was intact (the JB1.1 RSS), whereas the other end, detected by direct PCR in 6 out of 6 DNA aliquots from the same thymus, had undergone a 12-nt deletion (Fig. 3 A). We were unable to detect this DB1-side RSS insertion in splenic DNA from the same animal or in thymic DNA from any of four additional wild-type mice (Fig. 3 B, top). Dilution analysis (Fig. 3 B, bottom) led us to estimate the frequency of this RSS fragment insertion as at least 1 per ~17,000 cells based on our ability to detect the amplicon in a sample containing as little as 100 ng of genomic DNA. Both end insertions contained N regions. Using this sequence information, we designed a PCR assay that would amplify across the entire insertion, identifying the predicted molecule in multiple separate DNA aliquots from the same thymus, but not in splenic DNA from that same animal (Fig. 3 C). The identity of the fragment was confirmed by DNA sequencing (unpublished data). The fact that we were unable to detect the EL3–1 RSS fragment insertion in splenocytes suggests the cell containing the initial transposition event had undergone multiple rounds of cell division within the thymus, but failed to be positively selected into the peripheral repertoire.

A second such event, E2b–6 (Table I, Fig. S1, and Fig. S2, available at http://www.jem.org/cgi/content/full/jem.20070583/DC1), was detected in thymus from a core-RAG2 mutant mouse. We identified the JB1.1-side event by LM-TECA and used target locus sequence to directly amplify the corresponding DB1-side event (Fig. S2 A). The DB1 RSS formed a precise signal joint with a cryptic heptamer in an exon of a gene called lama3 on chromosome 18, whereas the JB1.1 RSS had suffered a 4-nt deletion. The JB1.1 insertion also contained a 4-nt N region. Thymocytes with this RSS fragment insertion were rarer than those with the aforementioned EL3–1 transposition because only occasional aliquots of thymic DNA from one animal gave the expected amplification product corresponding to the complete inserted RSS-ended fragment (Fig. S2 B, lane 3; identity was confirmed by DNA sequence analysis). It, too, was undetectable in splenic DNA.

A third event led us to define a “hotspot” for RSS end insertion in developing T cells. Event b196, detected by LM-TECA, consisted of a partial deletion of the JB1.1 RSS ligated to a region of chromosome 1 that lacks nearby annotated genes (Fig. 4 and Fig. S1). Direct PCR assays allowed us to identify the other end of this putative RSS fragment insertion, which consists of the precise fusion of an intact DB1 RSS to a cryptic heptamer at the chromosome 1 site. Remarkably, we recovered multiple different sequences for this insertion junction from the same and from different thymuses, thus identifying this region of chromosome 1 as a hotspot for RSS end insertion (Fig. 4, B and C). The sequences all contained full-length DB1 heptamers, but differed in the nature of N regions inserted within the joined fragment. We did not detect the inserted DB1 RSS in a limited number of 1 μg DNA aliquots from spleen or bone marrow, suggesting that this event might have a negative effect on T cell development or viability (Fig. 4 C). We were unable to detect by PCR a single amplicon containing both RSS ends inserted into the b196 hotspot (unpublished data). Given the apparent frequency of DB1 RSS insertion at this cryptic heptamer, we went on to examine this site for reciprocal translocation events between the DB1 coding end and the coding end generated by cryptic heptamer cleavage. We were able to detect this recombination event, as well as the formation of a hybrid joint between the cryptic RSS and the JB1.1 and JB1.2 coding elements, by direct PCR in multiple independent aliquots of thymus DNA (Fig. S3, available at http://www.jem.org/cgi/content/full/jem.20070583/DC1). The presence

Table I. Partial DNA sequences of DB1 and JB1 signal end insertions into various chromosomal locations (Continued)

| Event | DB1 RSS | JB1 RSS |
|-------|---------|---------|
| b356  | p53scid TCTATGG CACTG | GTTA | ggtttaggttAAGGTAGGG | AK148393, REPEAT | chr18:48387988 |
| b338  | p53scid TCTATG TGATG | ATGAG | aagtacgcaCTACGCCCA | 114668bps |(chr B) 5′ of D1 |
| b256  | p53scid CTGATCG TCGATG | CC | acaaggacgCAAGCCAGT | AF397014, unknown mRNA | chr13:17054115 |
| b292  | p53scid CTGATCG TCGATG | GGTT | acltacgtagCAGTTTCTC | TCR J1 | chr13:18575789 |
| b327  | p53scid CTGATCG TCGATG | aatgctgctgATATTTTTCT | chr12:110724915 |
| b361  | p53scid CTGATCG TCGATG | aaccttgggGAGGGGGGA | 4531bps 5′ of D2 | chr6:41486702 |
| b347  | p53scid CTGATCG TCGATG | T | tgtgtgtngGTGTTGTTGA | chr9:120505807 |
| b350  | p53scid CTGATCG TCGATG | aaattttgagCTTITCAAGG | 432bps 5′ of D1 | chr6:41463838 |
| b328  | p53scid CTGATCG TCGATG | TCGT | tgtgtctctgGCTTATCGGT | 467bps 3′ of D1 | chr6:41467144 |

Sequence of the DB1 (part A) or JB1 (parts B and C) signal end is given in upper case, with sequences aligned to reveal partial RSS deletions. N or P nucleotides are shown in lower case. Underlined target sequence indicates the position of a cryptic heptamer, as tested according to Cowell et al. (44). Potential stem-loop structures are denoted with the superscript HP and shown in boldface and italic. Events in which both RSS ends of a putative RSS fragment insertion were amplified from the same DNA sample are indicated in boldface and shown diagrammatically in Fig. S1. Nucleotide positions and target gene annotation refer to the August 2005 release of the University of California Santa Cruz mouse genome browser (http://genome.ucsc.edu/cgi-bin/hgGateway?org=mouse; cfr2, core-RAG2; T-cell, sorted splenic T cell; m/h, a mixture of murine thymus and Hela cell DNA; H2Ax, H2Ax null thymus; p53scid, p53 x scid thymus.
that such mice might exhibit more readily detectable RSS insertion events because of the inefficiency of proper joint formation. Using LM-TECA, we found multiple J-sided events in p53 x scid mice (Table I). In 7 of 13 instances, the Jβ1 RSS underwent modest (2 – 11 nt) deletion before insertion, as compared with 12 of 18 instances of deletion in wild-type mice (2 – 12 nt). Interestingly, only 3 of the 13 target loci in scid x p53 mice, compared with 7 of 18 in wild-type mice, contained authentic or cryptic RSS sequences. Also, 7 of the 13 targets in p53 x scid thymus were loci that undergo V(D)J recombination in that tissue, compared with only 3 out of 18 such targets in wild-type.

To more precisely compare the frequency of RSS fragment insertion among thymocytes from various mutant and wild-type strains, we used the second end of a transposed RSS fragment (either RSS fragment insertions or chromosomal translocations) to this same location in multiple mice allowed us to use this event as a measure of RSS end insertion frequency (see the following section).

The frequency of RSS insertion events in thymocyte DNA from wild-type and mutant mice

Scid mice contain a mutation in the gene encoding DNA-PK, which is a kinase involved in dsDNA break repair and V(D)J recombination. They are deficient in coding and, to a lesser extent, signal joint formation and produce few mature lymphocytes (28, 29). Coding joints that are formed often contain large deletions. Breeding the scid mutation onto a p53−/− genetic background results in the partial rescue of T cell development to the double-positive stage and allows accumulation of TCRβ locus coding joints (25). We reasoned that such mice might exhibit more readily detectable RSS insertion events because of the inefficiency of proper joint formation. Using LM-TECA, we found multiple J-sided events in p53 x scid mice (Table I). In 7 of 13 instances, the Jβ1 RSS underwent modest (2 – 11 nt) deletion before insertion, as compared with 12 of 18 instances of deletion in wild-type mice (2 – 12 nt). Interestingly, only 3 of the 13 target loci in scid x p53 mice, compared with 7 of 18 in wild-type mice, contained authentic or cryptic RSS sequences. Also, 7 of the 13 targets in p53 x scid thymus were loci that undergo V(D)J recombination in that tissue, compared with only 3 out of 18 such targets in wild-type.

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mice, we took advantage of the b196 RSS end insertion hotspot on chromosome 1 described in the previous section. We performed sets of 85 identical direct PCR assays for b196 Db-sided RSS insertion on 1-µg aliquots of DNA and used the number of negative samples to calculate the frequency of this insertion based on the Poisson distribution (see Materials and methods). The frequencies of b196 RSS insertion in thymus samples from the DNA damage detection mutants p53, ATM, and H2Ax were within twofold of the wild-type frequency (Fig. 5). Remarkably, the b196 RSS insertion was undetectable in 53bp1 and p53 x sid1 mutant mice (Fig. 5; frequency was at least 15-fold lower than wild type), although the region of DNA surrounding the hotspot had the identical sequence in all strains of mice in this study (not depicted).

As noted in the Introduction, there is biochemical evidence that suggests that the noncore domain of RAG2 suppresses RAG-mediated RSS transposition (10–12). This led us to examine core-RAG2 mutant mouse thymus for examples of TCRβ fragment RSS insertion (23). Using the b196 hotspot assay, we found that TCRβ RSS insertion was sevenfold greater in core-RAG2 compared with wild-type thymus (Fig. 5).

Of the four novel Jβ1 RSS end insertions we identified by LM-TECA in core-RAG2 thymus DNA, we were able to detect the corresponding Dβ1 RSS end insertion twice (Table I, events E2b-6 and b220). In both of these instances, the Dβ1 and Jβ1 RSS ends were inserted at or very near cryptic heptamers in the target locus. There was no evidence of target site duplication in either case.

**Signal circles as a potential source of DNA ends for RSS insertion**

More than half (26 of 48) of the RSS insertion events detected by LM-TECA displayed partial deletion of the TCRβ RSS, and 15 others contained N regions adjacent to full-length RSS ends (Table I). These events cannot be accounted for by a simple transposition mechanism in which the RAGs remain bound to a broken RSS end and catalyze attack by the 3’ hydroxyl group on a DNA duplex (Fig. 1 B) because such a mechanism would not result in deletion of RSS sequences or N region addition. Therefore, we went on to test the hypothesis that a signal circle might serve as donor for RAG-catalyzed RSS fragment insertion.

To test this potential mechanism, we inserted a PGK-neo cassette into plasmid clones containing (a) a precise Dβ1-Jβ1 signal joint; (b) a signal joint with a partially deleted RSS-23 heptamer; and (c) a signal joint with intact RSS ends separated by a 6-nucleotide N region (Fig. 6 A). Each intact circular plasmid was then transfected into a core-RAG2 mutant AMuLV-transformed preB cell line (23) that had been treated with the Abl-kinase inhibitor STI-571 to induce high levels of RAG expression and endogenous Vk-to-Jκ gene rearrangement (30). Stable transfectants were selected with G418, and purified genomic DNA from each culture was assayed by PCR for plasmid RSS insertion at Vk or Jκ RSSs associated with gene segments undergoing recombination in these cells. We detected insertion of the wild-type (a) and the N region-containing (c) signal joint plasmids at Vk RSS-12 sequences (Fig. 6, B and C). These insertions most often had a precise signal joint (Fig. 6 C) and diverse “coding” joints (Fig. 6 B and not depicted). Furthermore, each of 10 sequenced insertions of construct c contained portions of the donor plasmid’s N region in a joint with the Vk coding segment, indicating that RAG cleavage occurred at the RSS-23 heptamer within the signal joint construct. Thus, a subset of imprecise signal circles can serve as a source genomic instability. Interestingly, we failed to detect any insertions into Jκ RSS-23 sequences.

**DISCUSSION**

The V(D)J recombinase generates dsDNA breaks at the junctions between RSSs and Ig or TCR gene segments. It then relies on the NHEJ pathway to assemble these broken molecules to form the variable exons of antigen receptor genes and to safely discard RSS-ended DNA fragments. Most often, RSS-ended fragments are joined to form extrachromosomal signal circles. Previous biochemical and transfection experiments have shown that the V(D)J recombinase catalyzes various other reactions that might result in chromosomal translocations and RSS fragment transpositions or insertions (1). We designed the LM-TECA assay to allow us to detect and characterize in an unbiased fashion such events involving the D-J region of the TCRβ locus in developing thymocytes. Using this approach, we found that rearranged genomes of the lymphoid compartment are littered with remnants of V(D)J recombination reaction intermediates.

LM-TECA permits the recovery and sequence analysis of single chromosomally integrated RSS ends. We then used public database genome sequence information to design direct PCR strategies in efforts to recover the “other end” of a potential RSS fragment insertion or balanced chromosomal translocation. Using this strategy, we were able to demonstrate...
None of the events in which we recovered both RSS ends corresponded to a balanced translocation. In only one case, the b196 hotspot, did we detect a second fragment insertion. In three other cases, we could detect DJ1 RSS integration events involving the other side of the target locus break, but were unable to amplify the full contiguous event. Many other RSS end integration events could not be paired with corresponding partner ends. This is probably because of the inefficiency of our assays and the likelihood that not all of the original events occurred in cells that underwent subsequent clonal expansion. Small clonal populations bearing paired DJ1-J1 RSS insertion events might still be missed simply because of their infrequency in the entire thymic population. In only one case, the b196 hotspot, did we detect a second fragment corresponding to a balanced translocation.

Mechanisms of RSS fragment insertion

Perhaps the most surprising aspect of this study is that we did not observe RAG-mediated RSS fragment transposition. None of the events in which we recovered both RSS ends contained the short target site duplications characteristic of RAG-mediated RSS transposition. The RSS fragment insertions we did detect can be placed into two broad categories—one in which the recombination break point in the target locus is at or near a “cryptic” heptamer and one in which there is no such target sequence present (Table I) (15). Of the DJ1 RSS-23 events, nearly all the targeted chromosomes (9 of 12) contain complete RSS heptamers. In contrast, only 13 of 36 J1 RSS-12 insertions involve chromosomal targets with potential cryptic heptamers. This observation raises the possibility that RSS-12 (J1) sequences may differ from RSS-23 sequences in their potential for chromosomal insertion or the nature of the events in which they participate. Perhaps, as suggested previously, RSS-12 RAG complexes can attack phosphoester bonds in duplex DNA regardless of DNA sequence (9). Subsequent transesterification can result in dsDNA breakage with formation of a hairpin end that can be joined to the TCR gene-segment hairpin. However, almost every J1 RSS insertion was associated with the addition of nontemplated nucleotides, an observation that may not be consistent with this mechanism. Alternatively, it is possible that RSS-12 ends can pair with non-RSS sequences that can serve as recombination targets because of their unique structural features. Precedent for such a situation involving the breakage cluster region within the Bcl-2 protooncogene was noted in the Introduction (18).

The observation that 4 of the DJ1 and 20 of the J1 RSS end insertions had lost between 1 and 12 nt of RSS sequence raises the possibility that the reaction that forms these joints may not proceed directly from a free signal end intermediate. One interesting possibility is suggested by the work of Vanura et al., who recently showed that plasmids containing signal joints could undergo V(D)J recombination with plasmids containing proper RSSs in cotransfected cell lines and excised signal circles could integrate back into chromosomal RSSs in vivo (19). Neiditch et al. reported that signal joints undergo continuous RAG-mediated cleavage and rejoining until the recombination is inactivated during lymphoid development (31). Signal joint cleavage would cease, however, if rejoining were accompanied by nucleotide deletion on one or the other RSS. In this case, an “imprecise” signal joint might undergo synapsis, followed by V(D)J recombination in trans with a cryptic RSS, resulting in the insertion of the signal circle into the chromosome. In this case, the mutant RSS would function as a coding end. We showed that just this type of RAG-mediated insertion reaction could occur in cells performing V(D)J recombination (Fig. 6). Our observation that a majority of the J1 RSS-end insertions contain multiple-nucleotide RSS deletions is also consistent with this possibility.

Our failure to observe the target site duplications characteristic of RAG-mediated RSS transposition (7, 8) in any of the five events in which we recovered both RSS ends, as well as the high frequency of events targeting cryptic heptamers, are in potential conflict with a recent study from Reddy et al. These investigators used a chromosomally integrated transposition reporter construct to estimate the frequency of RSS fragment transposition as 1 per ~44,000 V(D)J recombination events (32). In that study, however, only ~1/3 of the recovered events were of this type. Most other events consisted of integrations at or near RSSs in the Ig κ locus that was actively rearranging in the transfected cells. These events were similar to what we observed (Fig. 6). It is possible that constraints imposed by native Ig or TCR locus chromatin structure effectively suppress true transposition events. Alternatively,
such events might fall below the limits of detection of the LM-TECA assay.

The frequency of RSS fragment insertion

Although the design of LM-TECA is qualitative, one can nonetheless make inferences regarding the frequency of RSS end insertion based on how the assay is performed. Each assay starts with 40 µg of genomic DNA containing a total of \(13.3 \times 10^6\) TCR\(\beta\) alleles. If each step was fully efficient (which is unlikely), then each microliter of the final recovered fraction would contain events from 133,000 TCR\(\beta\) alleles. From 4 µL of this fraction, we were able to recover, on average, two RSS end insertion events (unpublished data).

Thus, the frequency of TCR\(\beta\) D–J interval RSS fragment insertion in vivo is at least 1 in 266,000 genomes. We have measured the recovery of TCR\(\beta\) locus sequences after biotinylated primer extension to be \(\approx 6\%\) (unpublished data), and previous studies allow us to estimate the efficiency of linker ligation as \(\approx 5\%\) (33), thus allowing us to adjust our estimated transposition frequency to at least 1 event per 800 genomes or \(\approx 125,000\) events per young mouse thymus. This is likely an underestimate because some RSS end insertion events may be deleterious or may diminish the chance of functional TCR\(\beta\) variable exon assembly.

RSS end insertion in mutants

We used direct PCR assays of D\(\beta\) or J\(\beta\) RSS end insertion into the b196 hotspot (Fig. 4) to compare insertion frequencies among a panel of mutant mice (Fig. 5). To our initial surprise, ATM mutant mice, which, based on a recent reporter construct study, might be expected to show an increased frequency of transposition (34), showed a frequency of RSS end insertion, in this case translocation, similar to that of wild-type. The recent ATM study, however, showed defects in the processing of coding, but not signal ends. This study focuses on signal end behavior that appeared unaffected in that study (34). We also failed to detect increased hotspot RSS translocation in p53, H2Ax, and \(\gamma\delta\) mutant thymocytes (Fig. 5).

This may be caused by the fact that the RSS end insertion events we observed resemble mistargeted V(D)J recombination more than they do aberrant NHEJ.

In contrast, thymocytes from core-RAG2 mutant mice showed an approximately sevenfold increase in RSS end translocation frequency. This is interesting in light of previous reports that recombinase containing core-RAG2 catalyzes transposition in vitro with much greater frequency than recombinase containing wild-type RAG2 (10–12). In contrast, TCR\(\beta\) RSS end insertions lacked the target site duplications characteristic of the in vitro RAG-mediated transposition reaction (Table I). The C-terminal region of RAG2, which is missing in core-RAG2 mice, interacts with nucleosomal histones (35). Perhaps this interaction contributes to the stability of the postcleavage RSS complex and serves to limit RSS fragment insertions. This function of RAG2 might be considered analogous to the newly appreciated role of ATM in enhancing the stability of the postcleavage coding end complex (34). Alternatively, the core-RAG2 mutant recombinase may be more permissive for RSS synopsis with cryptic sites in trans.

Our data regarding RSS end insertion frequency must be interpreted with caution because the hotspot assay measures rearrangement of a small number of TCR\(\beta\) RSS ends with a single cryptic RSS on a different chromosome. This event may or may not be representative of the entire class of RSS end translocations involving native and cryptic RSSs. Of the 48 RSS insertion events characterized in this study, b196 was the only site targeted by more than one independent event, and the only one that displayed evidence of a translocation. Also, although we were unable to detect hotspot RSS end insertion in \(p53 \times \gamma\delta\) thymus, LM-TECA identified an array of other RSS insertion events in this same material (Table I).

Protecting the genome

Lymphoid leukemias and lymphomas are invariably associated with chromosomal translocations or deletions that affect protooncogenes, and many of these have the characteristics of aberrant V(D)J recombination (14, 16, 36). The mouse provides an excellent model system with which to study genes that limit and mutations that predispose to this type of developmentally regulated genomic instability. Young mice generate between 10 and 30 million thymocytes each day. During their ontogeny, each of these cells must rearrange at least one allele at two different antigen receptor loci (\(\alpha\beta\) or \(\gamma\delta\)). Thus, the lifetime total number of dsDNA breaks introduced by the V(D)J recombinase in mouse thymus is truly enormous. Given the ability of the RAG proteins to catalyze various strand-invasion reactions in vitro, including RSS fragment transposition, it is perhaps surprising how infrequent such events appear to be in vivo. This may be caused by several constraints on the recombinase in developing lymphocytes. First, RAG-mediated dsDNA cleavage in vivo requires synapsis between an RSS-12 and -23 sequence (37). Recent data from several groups has led to the suggestion that synapsis might be regulated by RAG-independent chromosome contraction (38). In addition, the frequency of inter-allelic V(D)J recombination is far lower than recombination in cis. Thus, chromatin structure and the compartmentalization of various chromosomal regions within the nucleus might serve to limit opportunities for transposition and translocation (39). Second, the same DNA–protein and protein–protein interactions that allow the RAGs to bind to RSSs and contribute to their synapsis are likely involved in holding RSS ends together after pairwise cleavage. The involvement of the RAG proteins themselves in protection against transposition may be of clinical significance in light of a recent epidemiological study that associated human lymphoma with a genetic polymorphism in RAG1 (40). Third, unbalanced translocations and some RSS fragment insertions are likely to disrupt essential cellular functions and result in lethality. In fact, the vast majority of developing thymocytes fail to assemble a self-tolerant, positively selectable TCR and undergo apoptosis, reducing the frequency of surviving cells...
with aberrant recombination events. To our surprise, null mutations in the genes encoding p53, ATM, H2AX, and DNA-PK did not seem to result in an increase in cells carrying RSS transposition events. We are continuing to survey mutations in NHEJ pathway proteins, as well as genes encoding “guardians of the genome” for effects on aberrant V(D)J recombination.

MATERIALS AND METHODS

Animals and cell lines. 63–12 and cR2-25, A-MuLV-transformed pro-B cell lines from RAG2−/− (41), and core-RAG2 mutant (23) bone marrow, respectively, were cultured in RPMI supplemented with 10% fetal calf serum and antibiotics. Human embryonic kidney 293T cells were cultured in DME within 8 gel plugs) were digested overnight with the restriction enzymes MaeIII or RsaI (NEB) at 37 °C. Plugs were rinsed with TE (10 mM Tris-

Full/jem.20070583/DC1), and 20 U of a hot-start Taq polymerase (Jump-

Approximately 40 μg of DNA (8 × 10⁶ cells embedded within 8 gel plugs) were digested overnight with the restriction enzymes MaeIII or RsaI (NEB) at 37 °C. Plugs were rinsed with TE (10 mM Tris-

DNA-PK with aberrant recombination events. To our surprise, null mutations in the genes encoding p53, ATM, H2AX, and DNA-PK did not seem to result in an increase in cells carrying RSS transposition events. We are continuing to survey mutations in NHEJ pathway proteins, as well as genes encoding “guardians of the genome” for effects on aberrant V(D)J recombination.

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