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The RAG1 N-terminal region regulates the efficiency and pathways of synapsis for V(D)J recombination

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Immunoglobulin and T cell receptor gene assembly depends on V(D)J recombination initiated by the RAG1-RAG2 recombinase. The RAG1 N-terminal region (NTR; aa 1–383) has been implicated in regulatory functions whose influence on V(D)J recombination and lymphocyte development in vivo is poorly understood. We generated mice in which RAG1 lacks ubiquitin ligase activity (P326G), the major site of autoubiquitination (K233R), or its first 215 residues (Δ 215). While few abnormalities were detected in R1.K233R mice, R1.P326G mice exhibit multiple features indicative of reduced recombination efficiency, including an increased Igk⁺:Ig λ ⁺ B cell ratio and decreased recombination of *Igh*, *Igk*, *Ig λ* , and *Tcrb* loci. Previous studies indicate that synapsis of recombining partners during *Igh* recombination occurs through two pathways: long-range scanning and short-range collision. We find that R1 Δ 215 mice exhibit reduced short-range *Igh* and *Tcrb* D-to-J recombination. Our findings indicate that the RAG1 NTR regulates V(D)J recombination and lymphocyte development by multiple pathways, including control of the balance between short- and long-range recombination.

Introduction

The diversity of jawed vertebrate adaptive immune responses depends on programmed assembly and hypermutation of antigen receptor (AgR) genes (Cooper and Alder, 2006). The first AgR gene diversification process to occur in developing lymphocytes is V(D)J recombination, which assembles Ig and TCR genes from arrays of V, in some cases D, and J gene segments (Schatz and Swanson, 2011). V(D)J recombination is initiated by the endonuclease RAG, a heterotetramer made up of RAG1 and RAG2. RAG binds to and cleaves DNA at recombination signal sequences (RSSs) that flank rearranging gene segments. RAG-mediated DNA cleavage requires the synapsis of two RSSs and leads to simultaneous generation of DNA double-strand breaks (DSBs) at the two sites (Schatz and Swanson, 2011). The DNA ends are processed and ligated by nonhomologous end joining repair factors (Rooney et al., 2004).

RAG1 is the primary DNA-binding and cleaving component of RAG, while RAG2 is an essential accessory factor. The minimal portion of RAG1 required for RSS binding and cleavage is called the “core” domain (R1core), consisting of aa 384–1008 of the 1040-aa protein (numbering according to mouse RAG; Fig. 1 A; Sadofsky et al., 1993; Silver et al., 1993). RAG1 also contains two

noncore regions, one at the N-terminus (aa 1–383; hereafter RAG1 N-terminal region [NTR]) and a short C-terminal tail (aa 1009–1040) that modulates RAG cleavage activity in vitro (Grundy et al., 2010; Kim et al., 2015) and is not a focus of this study. Mice expressing R1core exhibit a partial block in early B and T cell development, reduced numbers of mature lymphocytes, an altered V β gene segment repertoire in *Tcrb* recombination events, and an increase in aberrant V(D)J recombination products, demonstrating the importance of RAG1 noncore regions (Dudley et al., 2003; Horowitz and Bassing, 2014; Talukder et al., 2004). The functional significance of the RAG1 NTR is highlighted by the wide array of atypical SCID-like phenotypes found in human patients with mutations in this region (Notarangelo et al., 2016). While multiple regulatory functions have been associated with the RAG1 NTR, largely through in vitro and cell line studies, little is known about its roles in the context of lymphocyte development and AgR repertoire formation in vivo (Jones and Simkus, 2009).

Distinct functions have been attributed to different portions of the RAG1 NTR. RAG1 aa 287–351 constitute a zinc-RING finger domain that binds four zinc atoms and possesses ubiquitin ligase

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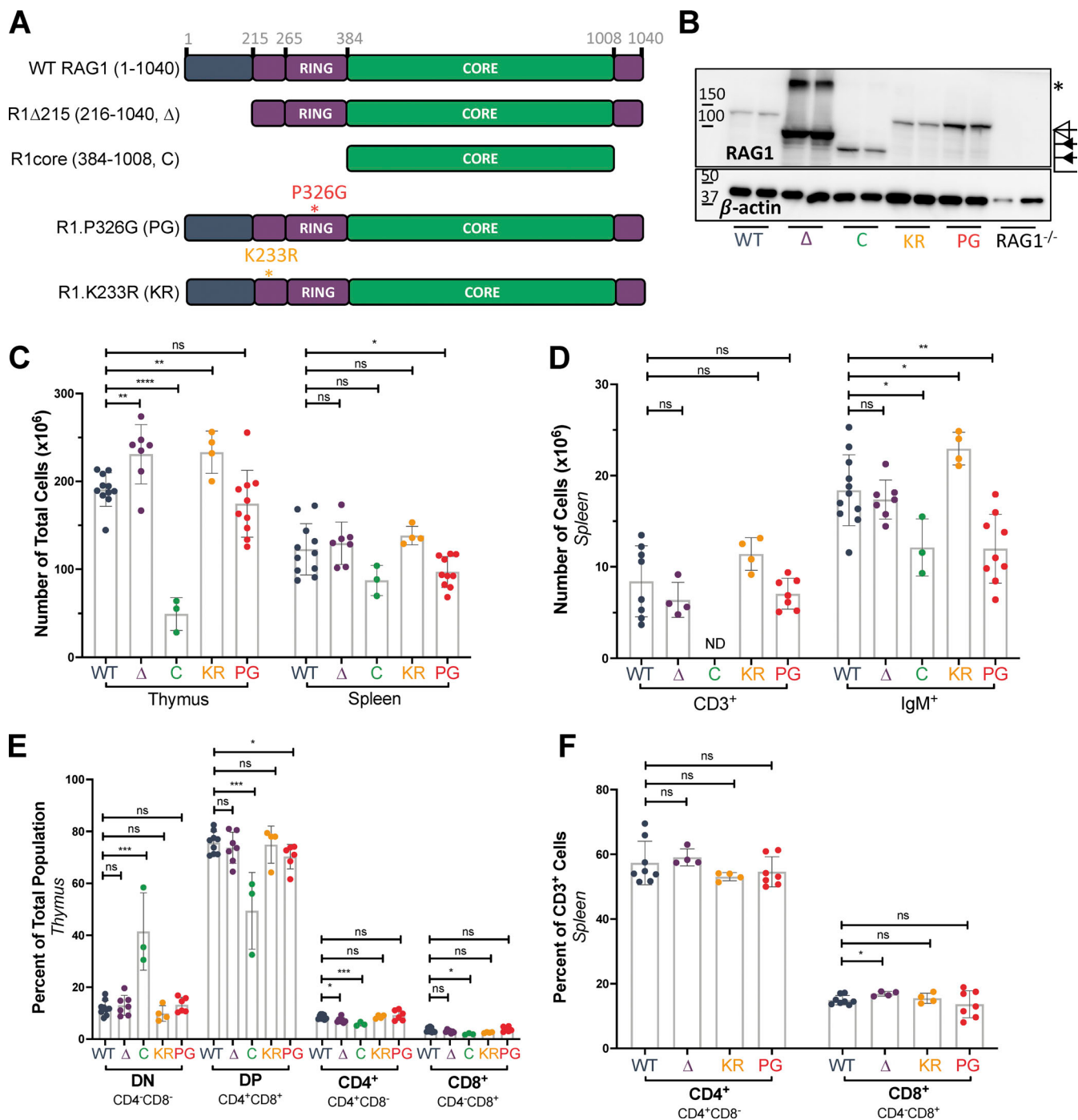


Figure 1. RAG1 NTR regulates RAG1 protein levels and has minimal effect on T cell development. (A) Schematic of RAG1 mutants. Figure not to scale. (B) Western blot of RAG1 in whole thymic lysates from mice 4–6 wk of age. Open triangle, full-length protein; closed triangles, truncated proteins. Asterisk, band approximately twice the molecular weight of R1Δ215 that likely represents a RAG1 dimer resistant to dissociation, as observed previously (Leu and Schatz, 1995). (C) Total numbers of nucleated cells in thymus and spleen. (D) Numbers of live CD3⁺ T cells or IgM⁺ B cells in the spleen. (E) Percentages of DN (CD4⁺CD8⁻), DP (CD4⁺CD8⁺), CD4⁺ (CD4⁺CD8⁻), and CD8⁺ (CD4⁻CD8⁺) thymocytes in whole thymuses. (F) Percentages of CD4⁺CD8⁻ and CD4⁻CD8⁺ T cells among live CD3⁺ cells in the spleen. Data are presented as mean with error bars indicating SEM. Statistical significance determined by two-tailed Welch's *t* test (ns, *P* > 0.05; *, *P* ≤ 0.05; **, *P* ≤ 0.01; ***, *P* ≤ 0.001; ****, *P* ≤ 0.0001).

activity (Jones and Gellert, 2003; Yurchenko et al., 2003). Mutation of the zinc-coordinating residue C325 (C325Y) abrogates ubiquitin ligase activity, dramatically reduces recombination activity, disrupts protein tertiary structure, and almost completely blocks lymphocyte development in mice (Deng et al.,

2015; Simkus et al., 2007). The corresponding mutation in human RAG1 (C328Y) also results in severe B and T cell lymphopenia (Villa et al., 2001). However, because this mutation disrupts RAG1 folding and structure (Simkus et al., 2007), it is difficult to determine the contribution of ubiquitin ligase

activity per se to the phenotypes observed (Deng et al., 2015). Mutation of a proline residue within the RING domain to glycine (P326G) disrupts ubiquitin ligase activity with only minor perturbation to the structural integrity of the domain and modestly reduces recombination of extrachromosomal plasmid substrates (Simkus et al., 2007). Numerous targets of RAG1-mediated ubiquitination have been identified in vitro, including the nuclear import factor KPNA1, histone H3, and histone variant H3.3, with histone ubiquitination implicated in recruitment of post-cleavage repair proteins (Deng et al., 2015; Grazini et al., 2010; Jones et al., 2011; Kassmeier et al., 2012; Simkus et al., 2009). In addition, RAG1 autoubiquitination, primarily at residue K233, stimulates RAG cleavage activity in vitro and has been implicated in increased recombination of extrachromosomal substrates in transfected cells (Singh and Gellert, 2015). A region mapping to the first ~200 aa of RAG1 is involved in the down-regulation of RAG1 protein levels through interactions with VprBP, a subunit of the cullin E3 ubiquitin ligase complex (Kassmeier et al., 2012; Schabla et al., 2018). The RAG1 NTR also regulates RAG1 nucleolar sequestration, with aa 243–249 required for nucleolar import and aa 1–215 required for release from the nucleolus and efficient V(D)J recombination (Brecht et al., 2020).

At AgR loci, RAG accumulates in recombination centers (RCs) encompassing J and J-proximal D gene segments (Schatz and Ji, 2011). RAG is thought to first bind to an RSS within the RC, after which a partner RSS is brought into the RC for synapsis and cleavage, a process long assumed to involve chromatin looping and random collision. Recently however, a model of RAG chromatin scanning has emerged to explain partner RSS capture, based initially on the observation of a strong orientation bias for recombination to cryptic RSSs located >5–10 kb away from a reference RSS (Hu et al., 2015). Subsequent analyses of Ig heavy chain gene (*Igh*) recombination provided strong evidence that V(D)J recombination can occur through a mechanism involving RAG chromatin scanning driven by cohesin-dependent loop extrusion (Ba et al., 2020; Dai et al., 2021; Hill et al., 2020; Jain et al., 2018; Zhang et al., 2019). Short-range recombination events between gene segments within the *Igh* RC were shown to be cohesin independent and governed largely by RSS sequence and not RSS orientation (Ba et al., 2020; Zhang et al., 2019). In contrast, longer-range recombination events were found to require the cohesin component Rad21 and a convergent orientation of the recombining RSSs (Ba et al., 2020; Dai et al., 2021; Jain et al., 2018; Zhang et al., 2019) and to be regulated by the chromatin looping factor CCCTC-binding factor (CTCF; Ba et al., 2020), CTCF-binding elements (CBEs; Ba et al., 2020; Guo et al., 2011; Jain et al., 2018; Lin et al., 2015), and the cohesin loader WAPL (Dai et al., 2021; Hill et al., 2020). Together, these data strongly support the existence of two distinct modes of recombination that differ in the mechanism leading to RSS synapsis—short-range recombination within or close to an RC that involves random RSS collision and long-range scanning driven by cohesin-mediated loop extrusion. It is unknown whether the RAG proteins exert control over the choice between these two modes of recombination.

To study the in vivo roles of the RAG1 NTR in V(D)J recombination and lymphocyte development, we generated and characterized mice in which RAG1 lacks ubiquitin ligase activity

(P326G), the major site of autoubiquitylation (K233R), or its first 215 aa (Δ 215). We found that RAG1 ubiquitin ligase activity, the K233 autoubiquitination site, and the first 215 aa help restrict RAG1 protein levels. We show that the P326G, but not the K233R, mutation results in impaired B cell development and decreased *Igh*, *Igk*, *Igl*, and *Tcrb* recombination, with no evidence that RAG1 or its ubiquitin ligase activity regulates AgR locus accessibility, arguing that RAG1 ubiquitin ligase activity stimulates recombination efficiency. We also demonstrate that the RAG1 NTR regulates the balance between short- and long-range recombination, with the first 215 aa facilitating short-range D-to-J recombination. Overall, our findings identify multiple, distinct roles played by the RAG1 NTR in regulating V(D)J recombination and lymphocyte development.

Results

Post-transcriptional regulation of RAG1 protein levels by the RAG1 NTR

To assess the function of the RAG1 NTR in vivo, we generated and analyzed three mouse lines on the C57BL/6 background with alterations in previously identified functional elements of the NTR. The RAG1 NTR is composed of two subregions, with the first 215 aa contributing to protein stability and intranuclear trafficking and the remainder, aa 216–383, consisting of a basic region harboring the major autoubiquitination site and a zinc-RING finger domain with ubiquitin ligase activity (Fig. 1 A; Brecht et al., 2020; Jones and Gellert, 2003; Jones and Simkus, 2009; Kassmeier et al., 2012; Yurchenko et al., 2003). To distinguish the contributions of these two subregions, a mouse line lacking the first 215 aa was generated (R1 Δ 215; Fig. 1 A and Fig. S1 A). The role of RAG1-mediated ubiquitination was assessed in a mouse line in which RAG1 lacks ubiquitin ligase activity (R1.P326G, hereafter R1.PG), while the function of autoubiquitination was evaluated using a mouse line in which the major site of RAG1 autoubiquitination was mutated (R1.K233R, hereafter R1.KR; Fig. 1 A and Fig. S1 A; Jones and Gellert, 2003; Simkus et al., 2007).

Western blots of whole thymic lysates revealed elevated RAG1 protein levels in all RAG1 mutant strains compared with WT, with levels dramatically elevated in R1 Δ 215 (Fig. 1 B and Fig. S1 B). The increase in protein levels was not due to changes in RAG1 mRNA levels (Fig. S1 C). Elevated levels of RAG1 protein in R1.KR and R1.PG thymocytes are consistent with the hypothesis that autoubiquitination serves to down-regulate RAG1 protein levels (Singh and Gellert, 2015), while increased levels of the R1 Δ 215 protein are consistent with loss of VprBP-cullin-mediated protein degradation (Kassmeier et al., 2012; Schabla et al., 2018). R1 Δ 215, R1.PG, and R1.KR mice could be maintained to at least 18 mo of age without overt evidence of pathologies (minimum of five mice per genotype), indicating that their elevated RAG1 protein levels do not severely compromise life span or physiology.

RAG1 ubiquitin ligase activity is required for efficient *Igk* and *Igl* recombination and production of normal numbers of *Ig*⁺ B cells

To assess the role of the RAG1 NTR in T and B cell development, lymphocyte subpopulations were analyzed by flow cytometry in

thymus, bone marrow, and spleen in WT and RAG1 mutant mouse strains. Gross $\alpha\beta$ T cell development was largely unperturbed in R1 Δ 215, R1.PG, and R1.KR mice: thymocyte and splenic T cell numbers and percentages were either normal or slightly elevated (Fig. 1, C, D, and F; and Fig. S2, A and B), and percentages of CD4⁺/CD8⁺ double-positive (DP) thymocytes were not different from those in WT mice, with the exception of a small (~5%) decrease in DP thymocytes in R1.PG mice (Fig. 1 E; Burn et al., 2021 Preprint). As expected, R1core mice displayed reduced thymocyte numbers and a relative accumulation of DN thymocytes and reduction in DP thymocytes (Fig. 1, C and E; Dudley et al., 2003).

B cell development was largely unperturbed in R1 Δ 215 and R1.KR mice, with normal numbers of pro-, pre-, and immature B cells in the bone marrow (although the percentages of preB and immature B cells were slightly reduced in R1.KR mice) and normal or somewhat elevated numbers of IgM⁺ cells in the spleen (Fig. 1 D; and Fig. 2, A and B). In contrast, numbers of bone marrow preB and immature B cells and of mature splenic B cells were reduced in R1.PG mice (Fig. 1 D and Fig. 2 A), with the reduction most evident in the immature B cell population, which also exhibited a relative decline (Fig. 2, A and B; and Fig. S2 C). Furthermore, the Ig κ /Ig λ ratio on IgM⁺ bone marrow and splenic B cells was significantly elevated in R1.PG and RAG1core, but not in R1 Δ 215 or R1.KR, mice relative to WT mice (Fig. 2 D). This elevated Ig κ /Ig λ ratio is due to decreased genesis of Ig λ B cells in the bone marrow, which is also reflected in the spleen (Fig. 2, C–F). Given the normal parameters of B cell development in R1.KR mice, the RAG1 ubiquitin ligase domain appears to perturb B cell development by a mechanism that is independent of RAG1 autoubiquitination.

Ig λ B cells develop from preB cells that fail to generate a nonautoreactive B cell receptor from Ig κ recombination and proceed to Ig λ recombination (Gorman and Alt, 1998). In preB cells, RAG-mediated DSBs up-regulate expression of the Pim2 kinase (Bednarski et al., 2012), whose activity enhances preB cell survival, thereby facilitating Ig λ recombination and development of Ig λ B cells (Derudder et al., 2009). The decreased production of Ig λ B cells and reduced numbers of developing and mature B cells in R1.PG mice are consistent with decreased efficiency of V(D)J recombination, a specific impairment in Ig λ rearrangement, and/or diminished survival of preB cells. To investigate V(D)J recombination, we conducted Taqman PCR to quantify Ig κ and Ig λ rearrangements in sorted preB cells of WT and R1.PG mice. Recombination of both loci is reduced in R1.PG mice relative to WT (Fig. 3 A). The decrease in Ig λ recombination is somewhat more pronounced than at Ig κ , which might reflect the ordered nature of Ig κ and Ig λ rearrangements rather than a specific defect in Ig λ recombination.

We considered that RAG1 ubiquitin ligase activity might stimulate recombination by driving transcription-associated chromatin accessibility. To investigate this possibility, we analyzed *Igh*, *Ig λ* , and *Tcrb* locus transcriptional accessibility in developing lymphocytes lacking RAG1 protein or expressing a DNA cleavage-incompetent R1.D708A mutant (Ji et al., 2010). We observed equivalent levels of λ 1 germline transcripts and genome accessibility (Fig. S2, D and E) in cells of each genotype.

Similarly, we observed equivalent levels of germline transcripts in RAG1^{−/−} and R1.D708A mice in the *Igh* RC in proB cells and in the two *Tcrb* RCs in DN thymocytes (Fig. S2, F and G). These data demonstrate that parameters of accessibility are not altered by the absence of RAG1 and argue that R1.PG and the other RAG1 mutants under study are unlikely to affect recombination by altering locus accessibility.

To determine whether R1.PG preB cells exhibit decreased induction of Pim2 mRNA and protein, we used an approach that isolates preB cell responses to RAG cleavage of *Ig κ* loci (Bednarski et al., 2012). The approach involves culturing preB cells from mice lacking the Artemis DSB repair factor and containing the anti-apoptotic *E μ BCL2* transgene and an *IgH* transgene that drives development of preB cells. Withdrawal of the IL-7 cytokine initiates *Ig κ* recombination, which leads to accumulation of RAG DSBs at *Ig κ* loci and signals from these lesions. We generated *E μ BCL2:IgH:Artemis^{−/−}* mice on both the WT RAG1 (BIA) and R1.PG (BIAPG) backgrounds and analyzed preB cells cultured from each genotype. Consistent with our recombination TaqMan PCR data, we detected a modest increase in germline λ 1, indicating a reduction of RAG *Ig κ* cleavage, in BIAPG cells (Fig. 3 B). Despite this decreased RAG cleavage, BIAPG cells exhibited no difference from BIA cells in up-regulation of Pim2 transcripts or protein (Fig. 3, C–E). Together, our data indicate that RAG1 ubiquitin ligase activity is required for normal efficiency of *Ig κ* and *Ig λ* recombination and that loss of ubiquitin ligase activity impairs the progression from *Ig κ* to *Ig λ* recombination in preB cells and the development of B cells through subsequent stages.

Alterations in *Igh* repertoire in R1 Δ 215 and R1.PG but not R1.KR mice

The murine *Igh* locus spans 2.7 Mb, with more than 100 V_H gene segments located ~100 kb upstream of 13 D_H and 4 J_H gene segments (Fig. 4 A). *Igh* recombination is initiated in proB cells by recruitment of RAG to the *Igh* RC, which encompasses the J_H segments and the most downstream D_H gene segment, DQ52 (Fig. 4 A; Ji et al., 2010; Teng et al., 2015). DQ52-to-J_H recombination occurs through a topologically unconstrained collision-based mechanism (Zhang et al., 2019), whereas recombination of other D_H gene segments to J_H involves a distinct mechanism in which a J_H-bound RAG complex scans chromatin, driven by cohesin-mediated loop extrusion (Ba et al., 2020; Zhang et al., 2019). J_H-bound RAG chromatin scanning is arrested by the CBE-containing element IGCR1, which lies immediately upstream of DFL16.1 (Fig. 4 A), explaining why DFL16.1 is the most frequently used D_H gene segment (Ba et al., 2020; Jain et al., 2018; Zhang et al., 2019).

To investigate the possibility that the RAG1 NTR mutations alter gene segment usage, we used high-throughput genome-wide translocation sequencing-adapted repertoire sequencing (HTGTS-Rep-seq) to characterize the *Igh* repertoire in B220⁺IgM[−] bone marrow B-lineage cells (predominantly preB cells) and B220⁺ spleen cells (predominantly naive, mature B cells) using J_H1 and J_H3 as baits (Lin et al., 2016). This approach reveals the repertoire of V_H and D_H gene segments that become joined to these J_H gene segments. We did not observe major alterations in V_H gene segment usage in the RAG1 NTR mutant mice generated for this study. Thus, we focus

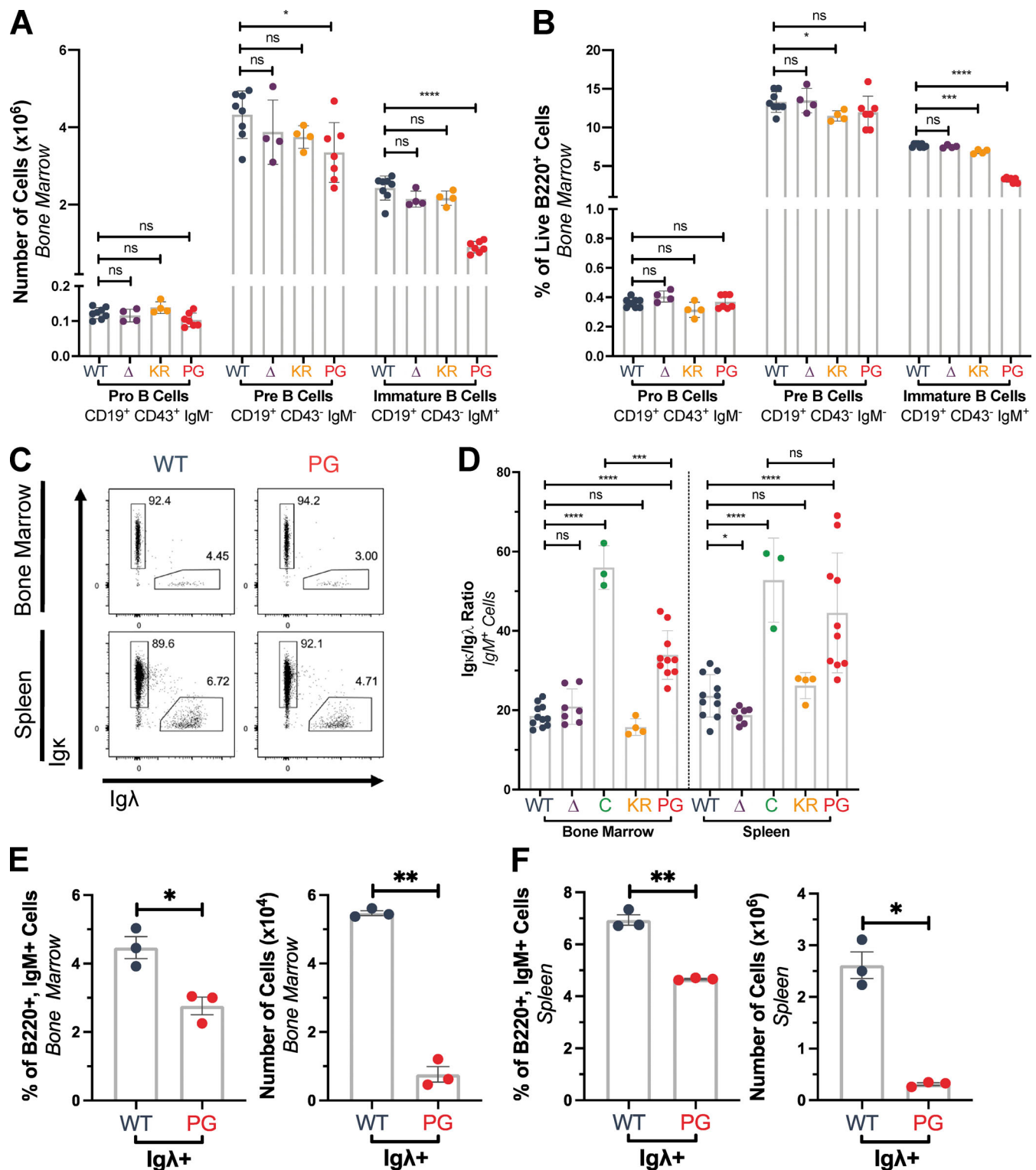


Figure 2. RAG1 ubiquitin ligase promotes Ig λ ⁺ B cell development. (A) Numbers of pro-, pre-, and immature B cells in bone marrow from mice 4–6 wk of age. (B) Percentages of pro-, pre-, and immature B cells among live B220⁺ cells in the bone marrow. (C) Representative flow cytometry data showing the populations of B220⁺ IgM⁺ B cells of the Ig κ or Ig λ lineage in bone marrow or spleen of WT and R1.PG mice. (D) Ig κ :Ig λ ratio in bone marrow and spleen. Calculated by taking exact number of IgM⁺ Ig λ ⁺ divided by exact number of IgM⁺ Ig κ ⁺ cells. All analyses were done from organs isolated from mice aged 4–6 wk. Each dot in B and D indicates data from an independent mouse, with data presented as mean $\pm$ SEM. (E and F) Graphed flow cytometry data depicting the frequency or number of B220⁺ IgM⁺ bone marrow (E) or spleen (F) cells that express Ig λ on their surface. Shown are individual data points collected from three mice from each genotype, with the average $\pm$ SEM. Statistical significance determined by two-tailed Welch's *t* test (ns, *P* > 0.05; *, *P* $\leq$ 0.05; **, *P* $\leq$ 0.01; ***, *P* $\leq$ 0.001; ****, *P* $\leq$ 0.0001).

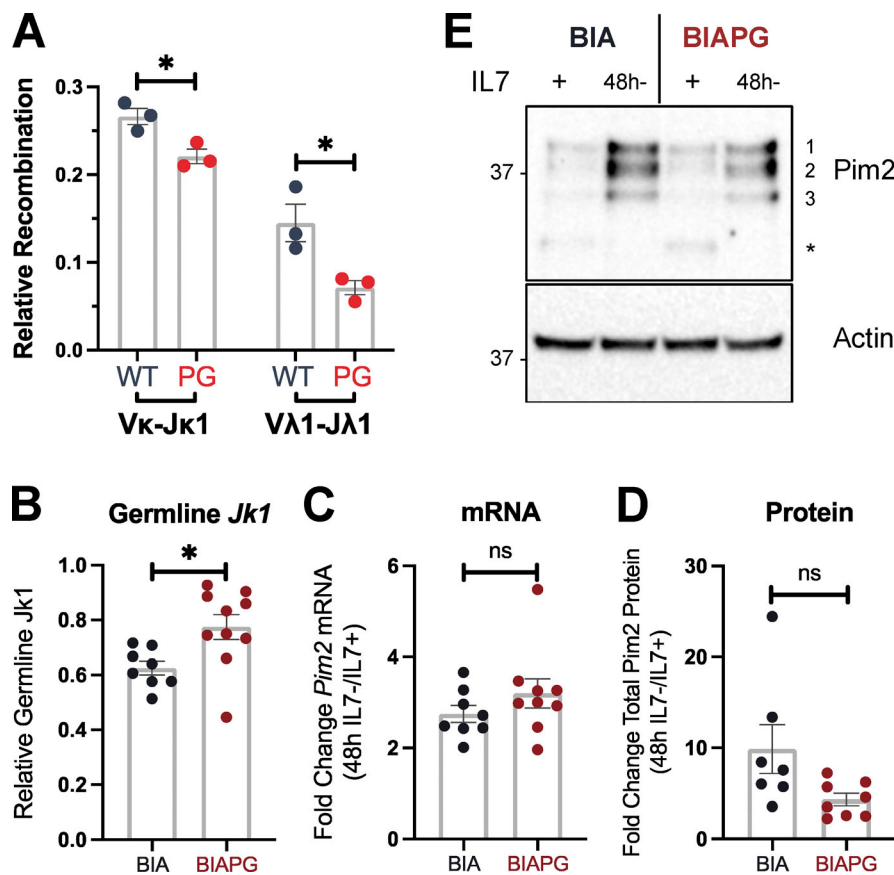


Figure 3. RAG1 ubiquitin ligase stimulates *Igκ* and *Igλ* gene recombination. (A) Taqman qPCR quantification of *Vκ-Jκ1* or *Vλ1-Jλ1* recombination conducted on genomic DNA from sorted preB cells. Signals from each sample were normalized to values from an assay for the invariant CD19 gene. Shown are the average values $\pm$ SEM from three independent isolations of preB cells. Statistical significance determined by two-tailed unpaired t test. (B–E) Characterization of the cellular response to RAG cleavage of *Igκ* alleles in primary preB cell cultures 48 h after IL-7 withdrawal. (B) Taqman qPCR quantification of intact germline *Jκ1* segment calculated by dividing germline *Jκ1* signals after IL-7 withdrawal by signals before IL-7 withdrawal. CD19 was used to normalize the input DNA. (C) Quantitative real-time-PCR quantification of the induction of *Pim2* transcripts calculated by dividing mRNA signals after IL-7 withdrawal by signals before IL-7 withdrawal. CD19 was used to normalize input cDNA. (D) Western quantification of the induction of *Pim2* protein calculated by dividing combined signals of all three *Pim2* isoforms after IL-7 withdrawal by signals before IL-7 withdrawal. (E) Representative Western blot with the locations of each *Pim2* isoform and a nonspecific band (asterisk) indicated. Statistical significance determined by two-tailed unpaired t tests (ns, $P > 0.05$; *, $P \leq 0.05$).

here on the D_H gene segment repertoire, with usage frequency presented for DQ52, DFL16.1, and the intervening D_H gene segments considered as a group (DSP; Fig. 1 A). The full dataset of V_H and D_H gene segment usage is provided in Table S1, Table S2, Table S3, Table S4, Table S5, and Table S6.

R1Δ215 mice exhibited a striking decrease of up to ~10-fold in the frequency of usage of DQ52 in rearrangements involving J_H1 and J_H3 and in both the bone marrow and spleen relative to WT mice (data for bone marrow and spleen are shown in Fig. 4, B–D; and Fig. S3, A–C, respectively). This was true in total VDJ rearrangements, nonproductive (VDJ[−]) rearrangements, and DJ rearrangements. Data for productive (VDJ⁺) rearrangements closely resemble those for total VDJ rearrangements and are not shown. In contrast, usage of DFL16.1 was not substantially or consistently altered in R1Δ215 relative to WT mice (Fig. 4, B–D; and Fig. S3, A–C). DFL16.1 was the most frequently used D_H gene segment in all genotypes analyzed, with usage higher in J_H1 than J_H3 rearrangements, as reported previously (Lin et al., 2016; Fig. S4). We conclude that deletion of the first 215 aa of RAG1 strongly reduces DQ52 usage and that this phenotype arises at the step of D_H -to- J_H rearrangement and is not due to selection for productive rearrangements.

R1.PG mice also exhibited perturbations in *Igh* gene segment repertoire. DQ52 usage was increased and DFL16.1 usage was decreased relative to WT in total VDJ, VDJ[−], and DJ rearrangements in bone marrow (Fig. 4, B–D). A similar trend was observed in spleen rearrangements, although the magnitudes of the changes were smaller and were not statistically significant

(Fig. S3, A–C). In contrast, D_H segment usage in R1.KR mice closely resembled that of WT in DJ_H and VDJ_H joints in bone marrow and spleen (Figs. 4, B–D; and Fig. S3, A–C). We conclude that loss of RAG1 ubiquitin ligase activity leads to increased representation of DQ52 and decreased representation of DFL16.1 in the *Igh* repertoire, most strikingly in preB cells, and that this phenotype is not due to loss of RAG1 autoubiquitination at K233. AgR repertoire alterations in R1core mice are considered together at the end of the Results.

Reduced secondary D-to- J_H recombination contributes to the R1.PG *Igh* repertoire phenotype

An attractive explanation for increased DQ52 and decreased DFL16.1 usage in R1.PG mice, suggested by our finding of reduced *Igκ* and *Igλ* recombination in these mice, is decreased recombinase activity leading to a decrease in secondary D-to- J_H recombination, which should preserve DQ52- J_H alleles at the expense of DFL16.1. If this explanation is correct, then R1.PG mice should exhibit increased usage of 5' and decreased usage of 3' J_H gene segments relative to WT. To test this prediction, we assessed *Igh* repertoires using ImmunoSEQ (Adaptive Biotechnologies), an approach that provides the repertoire of V_H , D_H , and J_H segments in VDJ_H rearrangements. The analysis was performed on bone marrow B220⁺IgM[−] cells from WT, R1Δ215, R1.PG, and R1core mice (two mice of each genotype).

The ImmunoSEQ results (Fig. 5) recapitulate the HTGTS-Rep-seq findings regarding D_H segment usage in R1Δ215 and R1.PG mice: in R1Δ215, DQ52 usage was reduced ~10 fold

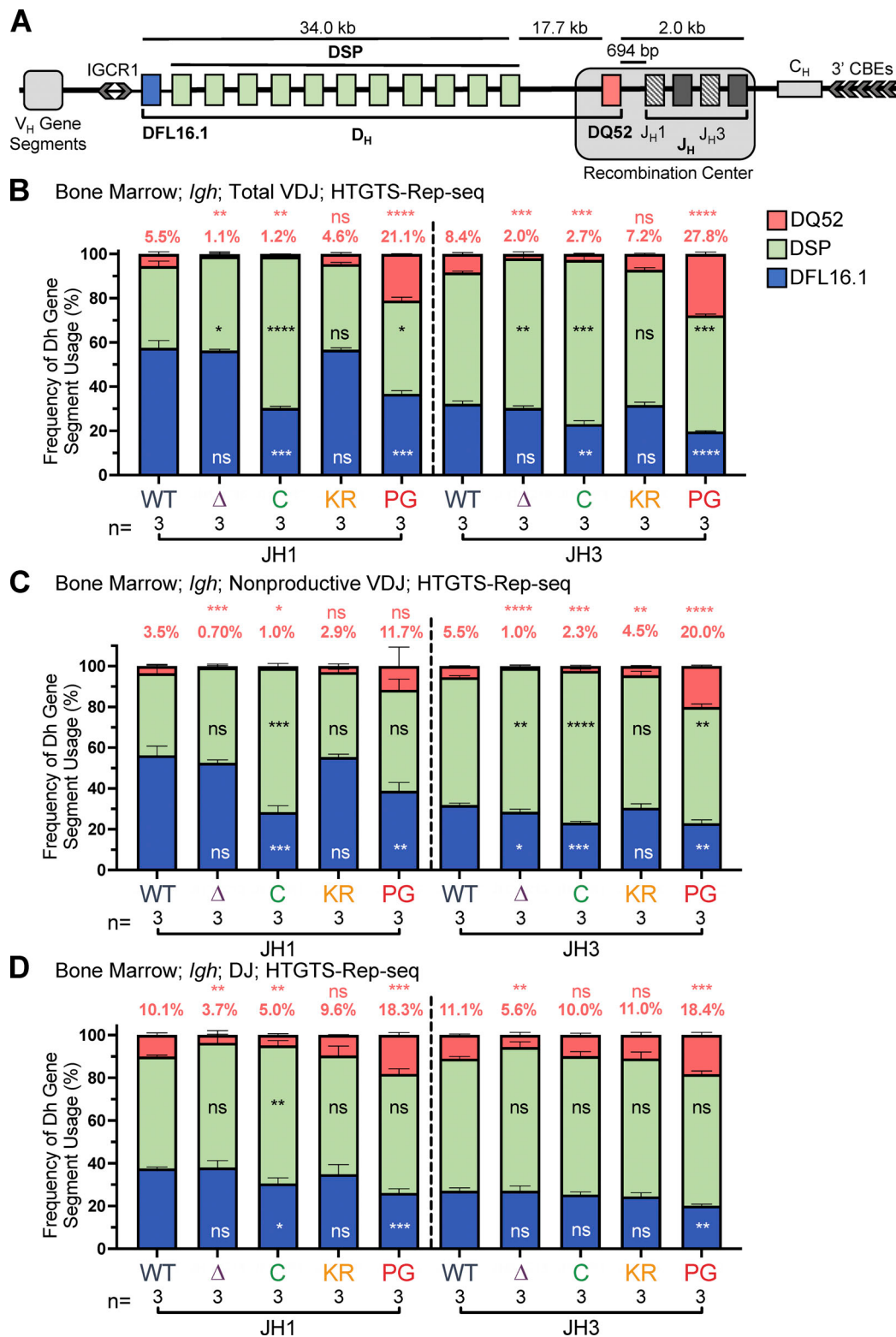


Figure 4. **D_H gene segment usage in *Igh* rearrangements in bone marrow assessed by HTGTS-Rep-seq.** (A) Schematic of murine *Igh* with the RC containing the J_H-proximal DQ52 gene segment and four J_H segments highlighted. Figure not to scale. Chevrons indicate the orientation of CBEs. CBEs in V_H portion of locus not depicted. (B–D) *Igh* HTGTS-Rep-seq data from mice 5–7 wk of age were analyzed for frequency of DFL16.1 (blue), DQ52 (salmon), and intervening 11 D_H gene segments (green) in total (B), nonproductive VDJ_H (C), or DJ recombination events (D) with J_H1 or J_H3 in bone marrow. Percentage of DQ52 usage is indicated above each bar. n, number of mice analyzed per genotype. Number of recombination events analyzed can be found in Table S4, Table S5, and Table S6. Data are presented as mean with error bars indicating SEM. Statistical significance relative to WT determined by two-tailed unpaired *t* tests (ns, *P* > 0.05; *, *P* ≤ 0.05; **, *P* ≤ 0.01; ***, *P* ≤ 0.001; ****, *P* ≤ 0.0001). DSP, intervening D_H gene segments considered as a group.

compared with WT, with little change observed for DFL16.1, while in R1.PG, DQ52 usage was increased and DFL16.1 usage was decreased compared with WT (Fig. 5 A). These differences were seen in total and nonproductive VDJ_H rearrangements (Fig. 5 A) and were recapitulated at each of the four J_H segments (Fig. 5 B). J_H repertoire analysis revealed increased J_{H1} and J_{H2} and decreased J_{H3} and J_{H4} usage in R1.PG mice relative to WT in total VDJ and VDJ_H rearrangements, as predicted by the model that R1.PG mice perform secondary D-to-J_H recombination inefficiently (Fig. 5 C). The J_H repertoire of R1Δ215 mice exhibited only minor differences from that of WT, with small increases in J_{H3} and J_{H4}, decreased J_{H2}, and slightly decreased J_{H1} usage, consistent with a normal or slightly elevated efficiency of secondary D-to-J_H recombination (Fig. 5 B). Together, our data argue strongly that the altered D_H segment repertoire in R1.PG mice is due in part or entirely to reduced secondary D-to-J_H recombination and that altered secondary D-to-J_H recombination makes at most a minor contribution to the R1Δ215 phenotype.

Reduced nonproductive VDJ_H alleles further support inefficient recombination in R1.PG mice

Reduced recombinase activity during *Igh* assembly in proB cells might reduce the frequency of VDJ_H[−] alleles because such alleles can only be carried forward to subsequent stages of B cell development in cells able to perform V-to-DJ recombination on both alleles. In support of reduced recombinase activity in proB cells due to the P326G mutation, the frequency of VDJ_H[−] alleles in R1.PG mice was consistently lower than in WT mice in both the HTGTS-Rep-seq and ImmunoSEQ analyses (Fig. 6, A and B; and Fig. S3 D). Indeed, the frequency of VDJ_H[−] alleles was as low or lower than in R1core mice, which are known to perform *Igh* gene assembly inefficiently (Dudley et al., 2003). In both R1.KR and R1Δ215 mice, the frequency of VDJ_H[−] alleles was similar to that in WT (Fig. 6, A and B; and Fig. S3 D), suggesting a normal efficiency of *Igh* gene assembly and consistent with the normal parameters of B cell development observed in these mice.

Analysis of Tcrb repertoires reveals reduced D_{β1}-to-J_{β1} recombination in R1Δ215 mice

Tcrb is made up of 21 functional V_β gene segments upstream and 1 V_β gene segment downstream of two D_β-J_β-C_β clusters (Fig. 7 A). Recombination occurs within two RCs, each containing a single D_β gene segment followed by seven J_β gene segments. The relative contributions of the RAG chromatin scanning versus collisional modes of recombination have not yet been established at *Tcrb*. If *Tcrb* follows the precedent set by *Igh*, then intra-RC D_{β1}-to-J_{β1} and D_{β2}-to-J_{β2} recombination events would be predicted to occur through a short-range collisional mechanism, whereas inter-RC D_{β1}-to-J_{β2} recombination events, which span at least 9 kb, might occur by RAG chromatin scanning. These assumptions, together with our finding that the R1Δ215 protein is selectively defective in short-range recombination, lead to the prediction that recombination events involving D_{β1} should be biased toward J_{β2} gene segments over J_{β1} gene segments in R1Δ215 mice relative to WT mice.

To test this prediction, we characterized *Tcrb* repertoires in sorted CD3e^{lo} DP thymocytes using ImmunoSEQ. The data

revealed that in R1Δ215 mice, VDJ_β rearrangements involving D_{β1} were indeed biased away from J_{β1} and toward J_{β2} gene segments, with usage of J_{β1} dropping from 55% in WT to 35% in R1Δ215 mice (Fig. 7 B). This bias in J cluster rearrangements involving D_{β1} was seen in nonproductive rearrangements (Fig. 7 B) and at all individual J_{β1} and J_{β2} gene segments with the curious exception of J_{β1.1} (Fig. S5 A; see Discussion). No such bias was observed in R1.KR and R1.PG mice, indicating that disrupting RAG1 ubiquitin ligase activity or autoubiquitination does not perturb D_{β1}'s preferences for the J_{β1} versus J_{β2} cluster. These findings argue that deletion of the first 215 aa of RAG1 disfavors intra-RC D_{β1}-to-J_{β1} recombination.

Analysis of the relative use of the D_{β1} and D_{β2} gene segments revealed that R1Δ215, R1.PG, and R1core mice exhibited significant increases in usage of D_{β1} over D_{β2} relative to WT in VDJ_β recombination events (Fig. S5 C). Notably, these increases parallel the reductions in the proportions of nonproductive VDJ_β rearrangement alleles observed in these three genotypes relative to WT (Fig. 6 C). R1.KR mice displayed a normal proportion of nonproductive VDJ_β rearrangements (Fig. 6 C). These data argue that RAG1 aa 1–215 and ubiquitin ligase activity, but not autoubiquitination, promote efficient recombination of *Tcrb*, and they raise the possibility that inefficient recombination contributes to preferential usage of D_{β1} over D_{β2}. They also indicate that inefficient *Tcrb* recombination is unlikely to explain the bias toward J_{β2} over J_{β1} recombination observed in R1Δ215 mice since recombination is at least as inefficient in R1.PG as in R1Δ215 (Fig. 6 C), but no J_{β2} > J_{β1} bias is seen in R1.PG (Fig. 7 B).

Igh and Tcrb repertoire changes in R1core mice

Igh and *Tcrb* repertoire alterations in R1core mice exhibited features resembling those in R1Δ215 and R1.PG mice, as expected given the complete absence of the RAG1 NTR. DFL16.1 usage was reduced in R1core relative to WT in a pattern that resembled that in R1.PG, whereas DQ52 usage in R1core was reduced relative to WT in a pattern that resembled that in R1Δ215 (Fig. 4, B–D; Fig. 5 A; and Fig. S3, A–C). As in R1Δ215 mice, R1core *Tcrb* rearrangements involving D_{β1} were biased away from J_{β1} and toward J_{β2} gene segments (Fig. 7 B).

We observed a substantially altered repertoire of V_β usage in R1core thymocytes (Fig. S5 B), consistent with a prior study (Horowitz and Bassing, 2014). A similar pattern of changes in V_β repertoire was observed in R1Δ215 thymocytes, although they are more modest in magnitude than in R1core (Fig. S5 B). The V_β repertoires in R1.KR and R1.PG thymocytes closely resemble that of WT thymocytes (Fig. S5 B; Burn et al., 2021 Preprint).

Discussion

RAG1 ubiquitin ligase activity contributes to efficient recombination and lymphocyte development

Our data indicate that the RAG1 P326G mutation decreases V(D)J recombinase activity during both B and T cell development. This reduced activity manifests in vivo as reduced secondary D-to-J_H recombination, reductions in nonproductive VDJ_H and VDJ_β alleles, decreased *Igκ* and *Igλ* recombination, and diminished development of Igλ⁺ B cells. Consistent with our observations,

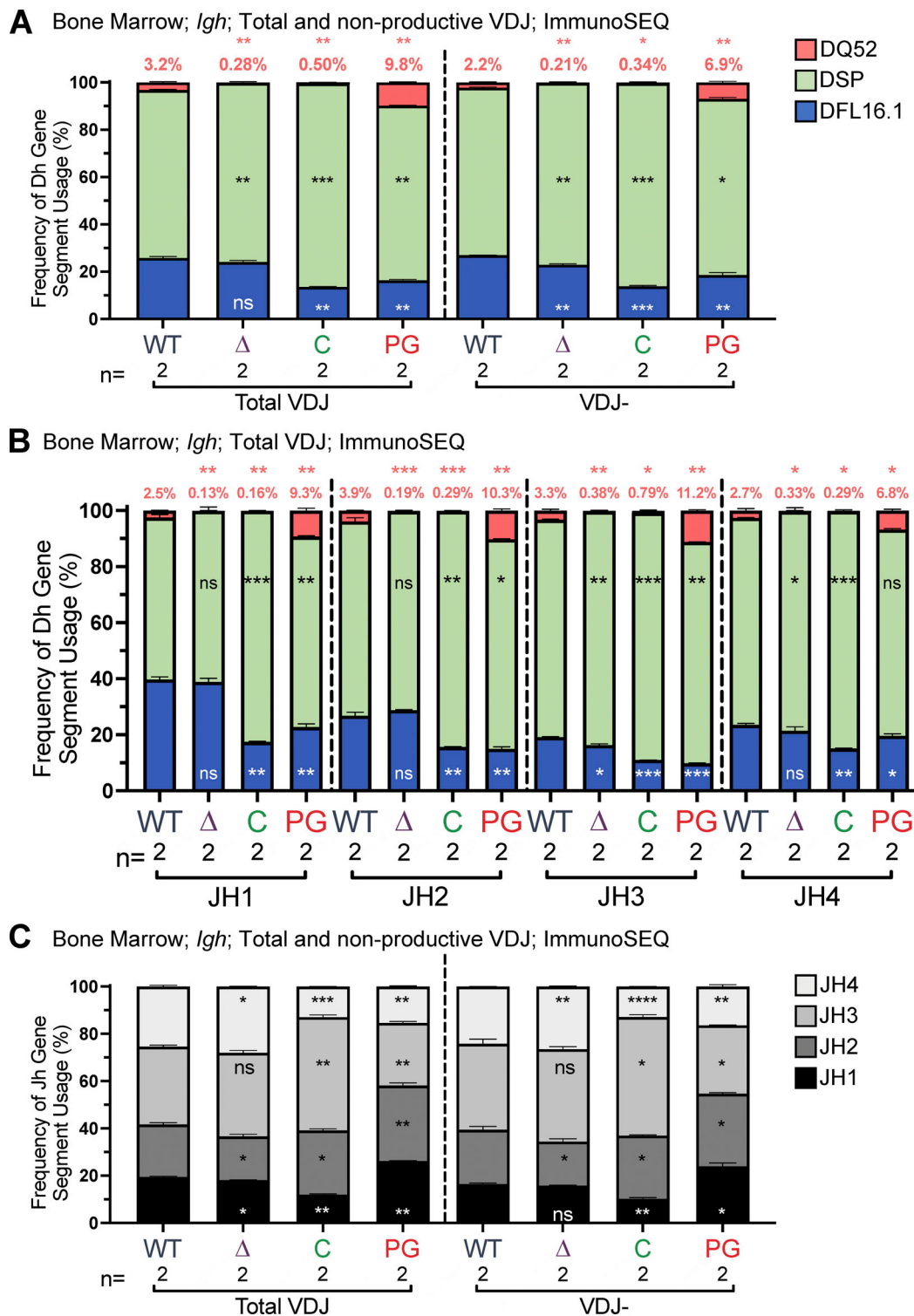


Figure 5. **D_H** and **J_H** gene segment usage in **VDJ_H** rearrangements in bone marrow assessed by ImmunoSEQ. (A–C) ImmunoSEQ data from mice 5–7 wk of age were analyzed for frequency of DFL16.1 (blue), DQ52 (salmon), and intervening 11 D_H gene segments (green) in total or nonproductive VDJ_H rearrangements (A) or VDJ_H rearrangements to individual J_H gene segments (B) and for J_H gene segments in total and nonproductive VDJ_H recombination events (C). Percentage of DQ52 usage is indicated above each bar in A and B. n, number of mice analyzed per genotype. Number of recombination events analyzed can be found in Table S7. Data are presented as mean with error bars indicating SEM. Statistical significance relative to WT determined by two-tailed unpaired *t* tests (ns, *P* > 0.05; *, *P* ≤ 0.05; **, *P* ≤ 0.01; ***, *P* ≤ 0.001; ****, *P* ≤ 0.0001).

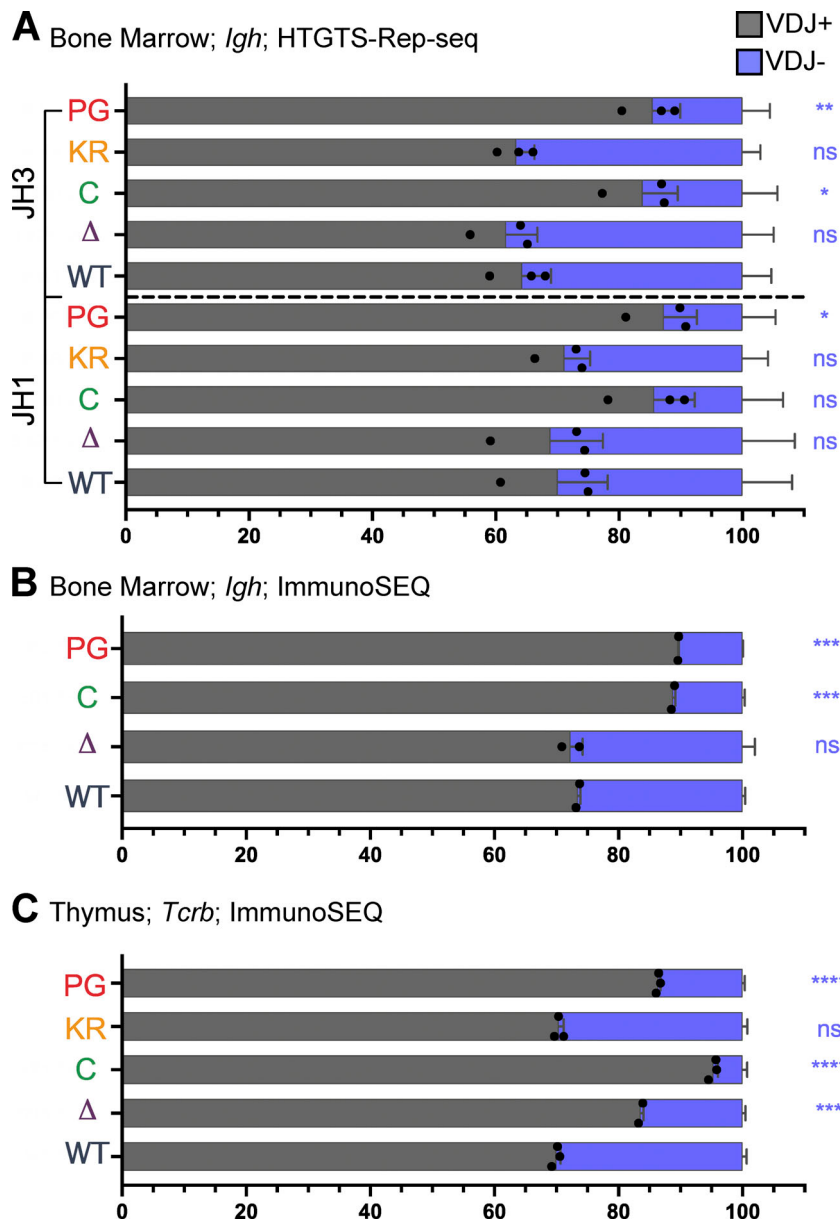


Figure 6. Frequency of productive and nonproductive VDJ_H and VDJ_B rearrangements. (A and B) Frequency of productive (VDJ⁺) and nonproductive (VDJ⁻) recombination events at *Igh* in bone marrow with J_H1 or J_H3 using HTGTS-Rep-seq data (A) and using ImmunoSEQ data (B). Number of recombination events analyzed can be found in Table S4, Table S5, Table S6, and Table S7. The frequency of VDJ⁺ alleles in mature B cells is predicted to be ~71% and was measured to be 73% in splenic B cells (Lin et al., 2016). **(C)** Frequency of VDJ⁺ and VDJ⁻ recombination events in thymus using ImmunoSEQ data. Data are presented as mean with error bars indicating SEM. Number of recombination events analyzed can be found in Table S8. Statistical significance relative to WT determined by two-tailed unpaired *t* tests (ns, *P* > 0.05; *, *P* ≤ 0.05; **, *P* ≤ 0.01; ***, *P* ≤ 0.001; ****, *P* ≤ 0.0001).

Burn et al. reported lower than normal levels of V_β and V_α rearrangements in DN and DP thymocytes, respectively, of R1.PG mice (Burn et al., 2021 Preprint). The decreased cleavage of *Igκ* loci that we detected in R1.PG preB cells argues that the RAG1 P326G mutation reduces RAG endonuclease activity. Impaired repair of RAG DSBs in the absence of RAG1-mediated ubiquitylation of H3 histones might also contribute to reduced levels of completed V(D)J rearrangements (Grazini et al., 2010). Reduced endonuclease activity could result from altered folding of the RAG1 N-terminus, abrogation of RAG1 ubiquitin ligase activity, or both. A change in structure of the RING domain could negatively impact the RAG1 endonuclease catalytic site or destabilize RAG1 dimerization and thereby stability of the heterotetrameric RAG enzyme. The normal lymphocyte development and V(D)J rearrangements observed in R1.KR mice argue that the RAG1 ubiquitin ligase does not stimulate RAG endonuclease activity through RAG1 autoubiquitination, at least at K233. We

cannot rule out that elevated R1.KR protein levels mask the reduction in recombination activity that would have been predicted from biochemical and cell line studies (Singh and Gellert, 2015). Notably, our finding of substantial levels of lymphocyte development in R1.PG mice argues strongly that the severe lymphocyte developmental defects reported previously in R1.C325Y mice and in a human bearing a RAG1 C328Y mutation were a consequence of perturbations of RAG1 function that extend beyond abrogation of ubiquitin ligase activity (Deng et al., 2015; Villa et al., 2001). Finally, considering that Burn et al. observed decreased expression of signaling molecules in R1.PG thymocytes, including the Syk protein that mediates B cell receptor signaling (Burn et al., 2021 Preprint), altered proB and preB cell proteomes could contribute to impaired development of Igκ⁺ and Igλ⁺ B cells in R1.PG mice. In this regard, while DSBs up-regulate Pim2 expression normally in R1.PG preB cells, we cannot rule out a contribution of decreased cell survival to the R1.PG phenotype.

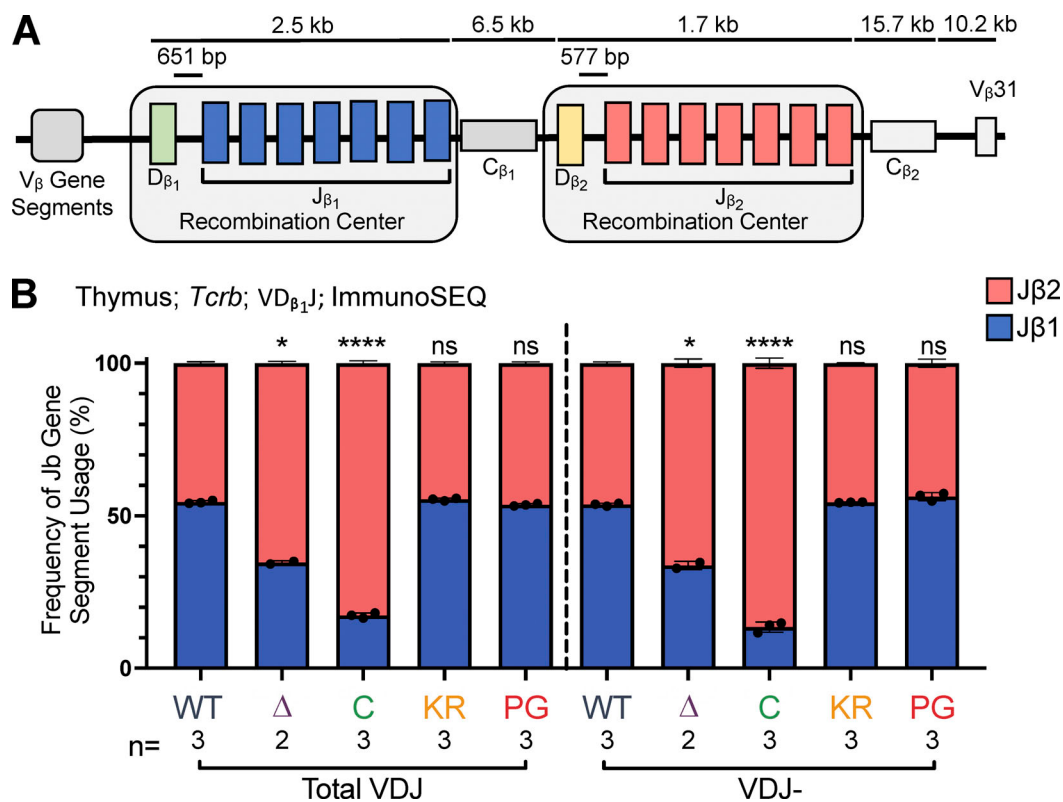


Figure 7. **J β gene segment usage in VDJ β rearrangements.** (A) Schematic of murine *Tcrb* with the two RCs containing D β 1 and J β 1 or D β 2 and J β 2 gene segments highlighted. Figure not to scale. (B) *Tcrb* ImmunoSEQ data from mice 5–7 wk of age in CD4⁺CD8⁺CD3 ϵ ^{lo} thymic T cells were analyzed for frequency of J β 1 and J β 2 gene cluster usage in VDJ recombination events using the D β 1 gene segment. Number of recombination events analyzed can be found in Table S8. Data are presented as mean with error bars indicating SEM. Statistical significance determined by two-tailed unpaired t tests (ns, $P > 0.05$; *, $P \leq 0.05$; ****, $P \leq 0.0001$).

The most striking alterations in AgR gene segment repertoire observed in R1.PG mice are increased usage of DQ52 and decreased usage of DFL16.1, accompanied by increased usage of J H 1 and J H 2 and decreased usage of J H 4. These findings are fully consistent with, and plausibly explained by, reduced secondary D-to-J H recombination in R1.PG mice relative to WT, although other mechanisms might also contribute.

Modes of recombination and the RAG1 NTR

Deletion of the first 215 aa of RAG1 results in strongly decreased usage of DQ52 in *Igh* rearrangements and reduced usage of J β 1 gene segments in *Tcrb* rearrangements involving D β 1 without grossly perturbing B or $\alpha\beta$ T cell development. Our data argue strongly that reduced usage of DQ52 is not due to increased secondary D-to-J H recombination. In particular, R1Δ215 mice exhibit levels of DFL16.1 usage that are, if anything, lower than in WT and levels of J H 1 usage that are only slightly reduced relative to WT (Fig. 4 and Fig. 5), neither of which is consistent with a large increase in secondary D-to-J H recombination. The possibility that reduced DQ52 usage in R1Δ215 mice is due to a small increase in secondary D-to-J H recombination that selectively targets DQ52-J H alleles is ruled out by the finding that DQ52 usage is strongly reduced in rearrangements involving all four J H segments, including J H 4 (Fig. 5 B); D-J H 4 alleles cannot be targets of secondary recombination. We conclude that decreased

usage of DQ52 in R1Δ215 mice is due primarily to mechanism(s) distinct from altered secondary D-to-J H recombination.

The high levels of expression of the R1Δ215 protein might have been predicted to lead to a hyper-recombination phenotype, but we see little evidence for this. Indeed, the decreased frequency of nonproductive *Tcrb* alleles observed in R1Δ215 mice suggests reduced recombination activity in thymocytes, despite the high levels of expression of the R1Δ215 protein. We propose that potential hyper-recombination effects of elevated R1Δ215 protein levels are counterbalanced by the inability of this protein to efficiently escape the repressive environment of the nucleolus. In our previous study, R1Δ215 was ~10-fold less active than WT RAG1 for recombination of a chromosomal substrate even though expressed at higher levels (Brecht et al., 2020). In this context, we emphasize that decreased recombination per se does not result in reduced usage of the J β 1 gene segment cluster, since J β 1 gene segment usage is normal in P326G thymocytes. While we cannot rule out the possibility that deletion of the first 215 aa perturbs folding of other portions of the RAG1 protein, aa 215 corresponds closely to an NTR domain boundary (Arbuckle et al., 2011), and robust RAG1 protein accumulation and normal lymphocyte development in R1Δ215 mice argue against major structural perturbations.

There are multiple possible explanations for the defect in short-range D-to-J recombination observed for the R1Δ215

protein. The first 215 aa of RAG1 have been implicated in regulation of RAG1 protein stability (Schabla et al., 2018), interaction with the splicing factor SF3A2 (Maitra and Sadofsky, 2009), zinc binding involving three conserved pairs of cysteine residues (Arbuckle et al., 2011), and release of RAG1 from the nucleolus (Brecht et al., 2020). It is unknown if these functions contribute to the phenotypes we observe in R1Δ215 mice. RAG1 noncore regions influence the targeting of RAG1 in the genome through chromatin interactions (Maman et al., 2016; Teng et al., 2015), and the NTR has been implicated in histone binding (Deng et al., 2015; Grazini et al., 2010; Jones et al., 2011). It will be important to determine whether aa 1–215 contribute to RAG1–chromatin interactions.

We propose that the first 215 aa of RAG1 are required to maintain the proper balance between short-range, collisional recombination and long-range recombination mediated by RAG chromatin scanning. How this balance is established is not known, and to our knowledge, no prior evidence has implicated RAG in its control. Collisional and chromatin scanning-mediated recombinations differ in the constraints placed on partner RSS capture by a RAG–RSS complex in the RC (Fig. 8). While collisional recombination is largely unconstrained topologically (Fig. 8 C), in RAG chromatin scanning the cohesin ring is envisioned to bring potential partner RSSs into close proximity of RSS-bound RAG in the RC in a topologically constrained manner (Fig. 8 B; Zhang et al., 2019). For RSSs separated by a short distance from one another in the RC (2–3 kb for the *Igh* and *Tcrb* RCs), loop extrusion (measured to occur at 0.5–2 kb/s; Davidson et al., 2019; Kim et al., 2019) would be predicted to move a potential partner RSS to the opposite side of the cohesin ring from the RAG-bound RSS very rapidly, likely sequestering it from the RC (as depicted for DQ52 in Fig. 8 B, panel i). Substantial collision-mediated recombination of gene segments that have been extruded through the cohesin ring adjacent to the RAG–RSS complex would be inconsistent with the powerful orientation bias observed for recombination of V_H , most D_H , and cryptic RSS sequences (Ba et al., 2020; Dai et al., 2021; Hill et al., 2020; Hu et al., 2015; Jain et al., 2018; Zhang et al., 2019). Cohesin-mediated loop extrusion might thereby compete with and inhibit short-range collisional recombination. RAG1 residues 1–215 might act to facilitate collisional recombination, perhaps by stabilizing the 12RSS–23RSS synaptic complex. We propose that such stabilization would be less important for scanning-mediated recombination due to the presence of the cohesin ring. Alternatively, or in addition, residues 1–215 might inhibit loop extrusion, allowing more time for collisional recombination to occur. We note that in the current model of RAG chromatin scanning (Zhang et al., 2019), the cohesin ring encounters the RAG-bound RSS at the nonamer end of the RSS, where the nonamer binding domain and the RAG1 NTR would be located. Our data argue that ubiquitin ligase activity is not required for the ability of the RAG1 NTR to facilitate short-range recombination. Further mechanistic insights might require reconstitution of RAG chromatin scanning in vitro.

We note that, because all V_H recombination involves RAG chromatin scanning (Dai et al., 2021; Hill et al., 2020), an altered balance between collisional and scanning modes of recombination

might not be expected to perturb V_H repertoire. Consistent with this, we did not observe substantial alterations in V_H repertoire in R1Δ215 mice.

Support for our model regarding the role of RAG1 aa 1–215 in establishing the balance between short- and long-range recombination derives primarily from our findings with the *Igh* locus, where collisional and scanning modes of recombination have been firmly established and mapped to particular classes of rearrangements. Because a similar mechanistic understanding is not yet available for *Tcrb* recombination, findings from this locus should be interpreted cautiously. Our observation that R1Δ215 thymocytes exhibit a skewing of $D_{\beta 1}$ recombination toward the $J_{\beta 2}$ cluster and away from the $J_{\beta 1}$ cluster should be considered only preliminary corroboration of our findings at *Igh*. $J_{\beta 1.1}$ is the only J_{β} segment that does not show this skewing, suggesting that its recombination is influenced by additional mechanisms. One speculative possibility is that Fos-driven RAG loading onto the $D_{\beta 1}$ 3'RSS (Wang et al., 2008), combined with the close proximity of $J_{\beta 1.1}$ to $D_{\beta 1}$, compensates for the R1Δ215 short-range recombination defect.

Insights into the phenotype of R1core mice

While the B cell developmental defects observed in R1core mice are mirrored in R1.PG mice (albeit in a milder form), the substantial defects in T cell development were not observed in any of the NTR mutant mice analyzed here and likely reflect the combined loss of multiple noncore domains/activities. *Igh* repertoire alterations in R1core B cells appear to represent a combination of loss of the first 215 aa, abrogation of ubiquitin ligase activity, and likely other deficits as well. *Tcrb* repertoire perturbations in R1core mice, such as reduced usage of $J_{\beta 1}$ gene segments and V_{β} repertoire changes, are mirrored to a substantial extent in R1Δ215 mice, and loss of ubiquitin ligase activity appears to make less of a contribution. The severe reduction in nonfunctional VDJ_{β} rearrangements observed in R1core thymocytes appears to be due to a combination of the defects observed in R1Δ215 and R1.PG thymocytes.

The generally more severe phenotypes observed in R1core mice compared with the RAG1 NTR mutants created for this study could be due to one or more additional domains/activities missing in the R1core protein, including a positively charged region between residues 216 and 238 that enhances recombination activity (McMahan et al., 1997), a signal for nucleolar localization between aa 243 and 249 (Brecht et al., 2020), the RAG1 C-terminal tail, loss of the strong dimerization activity provided by the zinc–RING finger, and the ability to interact with Ku (Raval et al., 2008) and chromatin (Maman et al., 2016), which have yet to be mapped within the RAG1 noncore regions.

Conclusions

Our findings implicate different portions/activities of the RAG1 NTR in regulating distinct functions: (i) maintenance of proper RAG1 protein levels (residues 1–215, ubiquitin ligase activity, major site of autoubiquitination), (ii) efficient V(D)J recombination and lymphocyte development (ubiquitin ligase activity), and (iii) efficient short-range recombination within RCs (residues 1–215). The RAG1 NTR is proposed to be derived from a

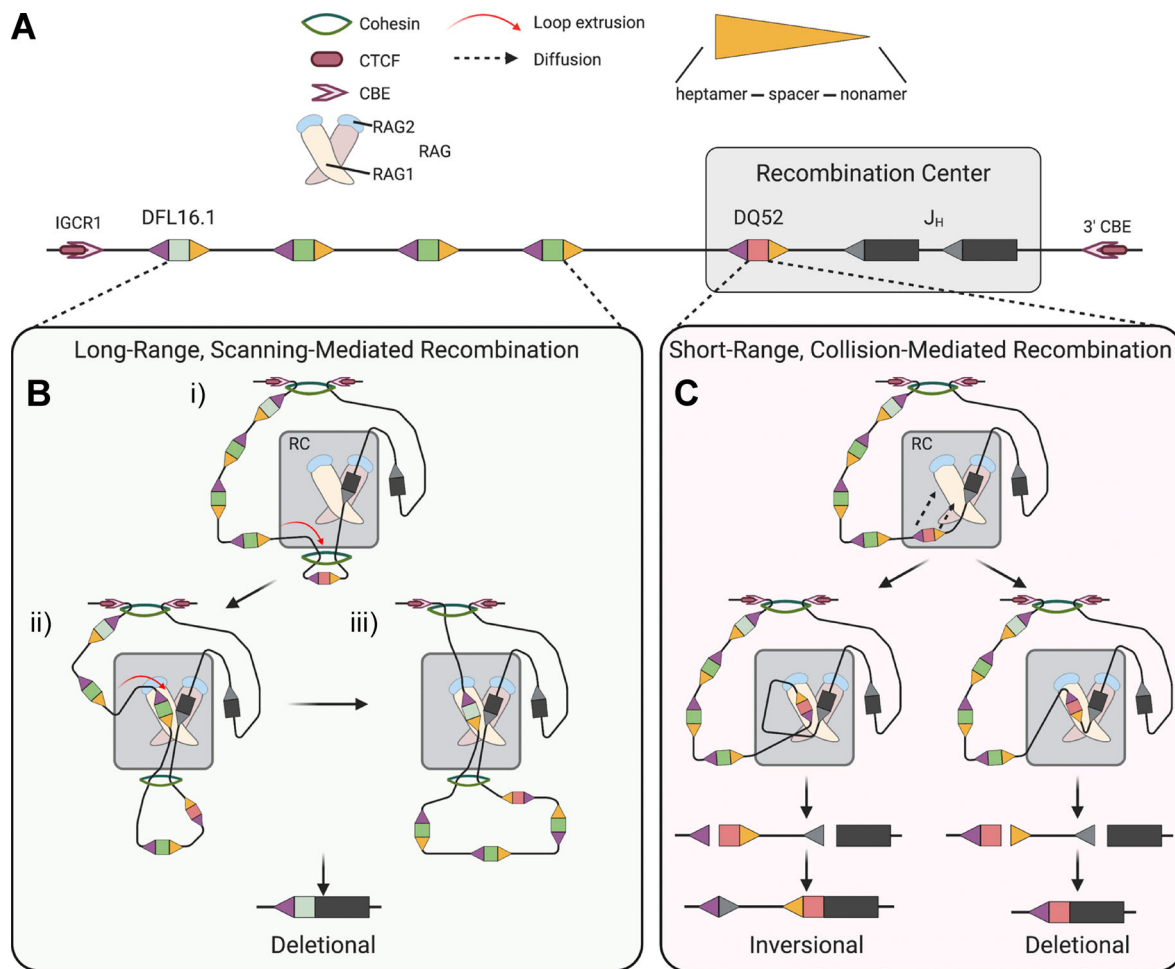


Figure 8. Schematic diagrams depicting models for short-range collision- and long-range scanning-mediated D-to-J recombination. (A) Schematic of murine *Igh* depicting some D_H and J_H gene segments. Gray box represents the RC, encompassing J_H gene segments and DQ52. Gene segments are depicted as rectangles, 23RSSs as gray triangles, and 3' and 5' D_H 12RSSs as orange and purple triangles, respectively. **(B)** Long-range recombination is thought to occur through RAG chromatin scanning in which the cohesin ring brings RSSs into the RC in close proximity of RSS-bound RAG in a topologically constrained manner. Panel i: DQ52, which resides very close to the J_H gene segments, has already been extruded through the cohesin ring and now is separated from J_H by cohesin. Long-range recombination (panels ii and iii) occurs predominantly by deletion due to the topological constraint imposed by cohesin. **(C)** Short-range recombination occurs through collision-mediated recombination, largely topologically unconstrained. Short-range recombination can result in either inversion or deletion; however, DQ52-to-J_H recombination occurs almost exclusively by deletion due to the different sequences of its 5' and 3' RSSs (Zhang et al., 2019). Models in B and C adapted from Zhang et al. (2019). This figure was created with BioRender.com.

transposable element known as *N-RAG-TP* (Panchin and Moroz, 2008) and to have become incorporated into a RAG1-like protein early in the evolution of RAG-like transposons (transposons that encode RAG1-like and RAG2-like proteins; Carmona and Schatz, 2017). The NTR of extant RAG1-like proteins shows considerable variation in the domains that have been retained (Fugmann et al., 2006; Huang et al., 2016; Martin et al., 2020), suggesting that the NTR has been a flexible substrate for the evolution of new functions. Given that known RAG-like transposons are <10 kb in size (Martin et al., 2020), an activity to ensure efficient short-range recombination might have arisen in a RAG-like transposon to facilitate synapsis and cleavage of the transposon terminal inverted repeats. In this scenario, the domain would have subsequently been repurposed for short-range recombination in jawed vertebrates. Alternatively, such an activity might have evolved only after RAG had begun to function as a recombinase in jawed vertebrates, to

help address the complexities of assembling large AgR loci containing many gene segments. It will now be important to determine which residues in the first 215 aa of RAG1 contribute to this activity and whether this activity can be detected in RAG-like transposases.

Materials and methods

Mouse strains

WT C57BL/6 mice were purchased from The Jackson Laboratory. R1core mice have been described previously (Dudley et al., 2003). R1Δ215 and R1.KR mice were generated by the Yale Genome Editing Center, as previously described, with CRISPR-Cas9 technology into C57BL/6N embryos (Wang et al., 2013). To generate R1Δ215 mice, two guide RNAs (Table S9) targeting the far N-terminus-encoding region of *Rag1* were coinjected into fertilized WT C57BL/6N eggs with a single-strand DNA Ultramer

(IDT) donor template (Table S9). The knock-in cassette replaces the sequence encoding the initial 215 aa of RAG1, such that upon homology-directed repair-mediated genomic integration, the protein will begin at aa 216. To generate R1.KR mice, a guide RNA (Table S9) and single-strand DNA Ultramer (IDT; Table S9) were coinjected into fertilized WT C57BL/6N eggs. This sequence introduces the K233R point mutation into *Rag1* and introduces a HindIII site that does not affect the amino acid sequence of the protein to facilitate genotyping. For both lines, single heterozygous males were generated and backcrossed to C57BL/6 females for eight generations. Heterozygote-by-heterozygote breeding was then conducted to establish homozygote lines of mice. R1Δ215 and R1.KR lines were confirmed to have only the designated mutations by sequencing of the *Rag1* locus. R1.PG mice were generated as described (Burn et al., 2021 Preprint). Sequencing revealed that R1.PG mice harbor an additional V238A mutation in *Rag1* (Fig. S1 A). This mutation is conservative, and V238 is not well conserved in vertebrates. These mice are deposited in the Mutant Mouse Resource & Research Centers Repository (strain RAG1-tml; #37105) and are cryo-archived.

All animal studies were conducted in accordance with national guidelines and were approved by the Institutional Animal Care and Use Committee of the Children's Hospital of Philadelphia or by the Yale Institutional Animal Care and Use Committee. Female and male mice were used for all experiments.

Western blotting

For Western blots for Fig. 1 B and Fig. S1 B, whole thymuses were lysed in RIPA buffer containing 1× cComplete protease inhibitor cocktail (Sigma-Aldrich; 4693159001) and Pierce Universal Nuclease for Cell Lysis (Thermo Fisher Scientific; 88700) and incubated on ice for 10 min. 25 µg of lysates was resolved on 8% SDS-PAGE gels. Immunoblotting was performed according to standard procedures with mAbs for RAG1 (mAb 23; Coster et al., 2012) and β-actin (Sigma-Aldrich; A5441).

For Western blots for Fig. 3 E, 6–10 million preB cells were harvested at indicated time points, washed with PBS, and resuspended in ice-cold lysis RIPA buffer with protease and phosphatase inhibitor cocktail (Cell Signaling) and benzonase (Sigma-Aldrich). Laemmli buffer was added to the lysate; then samples were boiled for 5 min. 20 µg of protein was loaded and run on ExpressPlus PAGE 4–12% gels (Genscript). Electrophoresed proteins were transferred to Immobilon-FL polyvinylidene fluoride membrane (EMD Millipore). Membranes were blocked with a blocking buffer (Tri base 1X, Tween 0.1%, and BSA 5%) and incubated overnight with a primary antibody. Primary antibodies were monoclonal mouse anti-Pim2 antibody (Santa Cruz Biotechnology; clone 1D12) and monoclonal mouse anti-βactin antibody (Sigma-Aldrich; clone AC-15). Blots were washed and incubated with appropriate HRP-conjugated secondary antibodies. After washing, proteins were revealed using WesternBright ECL HRP Substrate Kits (Advansta). All blots were acquired with the G:box Chemi-XRQ. Quantification was performed using ImageJ software (Wayne Rasband, National Institutes of Health, version 1.52q).

RAG1 mRNA analysis

RNA was isolated from whole thymuses with TRIzol reagent (Invitrogen) and was converted to cDNA with SuperScript III

and random hexamers according to the manufacturer's instructions (Life Technologies). Transcripts were quantified by iTaq Universal SYBR Green (Bio-Rad) with primers RAG1 Forward and RAG1 Reverse. Analysis was normalized using *Hprt* expression (primers *Hprt* Forward 1 and *Hprt* Reverse 1). Primer sequences are listed in Table S9.

Flow cytometry and cell sorting

Lymphocyte development was analyzed in age-matched mice aged 4–6 wk. Both males and females were analyzed, with no differences noted. The analyses for Fig. 1, C–F; Fig. 2, A, B, and D; and Fig. S2, A–C were performed as follows. Total thymocytes were harvested from whole thymuses, total splenocytes were harvested from whole spleens, and total bone marrow was harvested from whole bone marrow. Total cells were calculated based on number of nucleated cells after erythrocytes were lysed using ZAP-OGLOBIN II Lytic Reagent (Beckman Coulter; 7546138). Lymphocyte development analyses were conducted using flow cytometry. Thymic and splenic T cells were analyzed using PE anti-mouse CD3 (BioLegend clone 17A2), APC anti-mouse CD4 (BioLegend; clone GK1.5), and FITC anti-mouse CD8a (BioLegend; clone 53–6.7) antibodies and DAPI (BioLegend; 422801). B cell development in the bone marrow was analyzed using FITC anti-mouse CD43 (BD Biosciences; clone S7), PE anti-mouse CD19 (BioLegend; clone 6D5), PerCP anti-mouse B220 (BioLegend; clone RA3-6B2), and APC anti-mouse IgM (BioLegend; clone RMM-1) antibodies and DAPI. Igκ and Igλ staining was conducted in the bone marrow and spleen using PerCP anti-mouse B220, APC anti-mouse IgM, PE anti-mouse Igκ (BD Biosciences; 559940), and FITC anti-mouse Igλ (BD Biosciences; clone R26-46) antibodies and DAPI. Samples were run on BD LSRII or CytoFlex LX machines, and data were analyzed using FlowJo software (Tree Star).

The analyses for Fig. 2, C, E, and F were performed as follows. Cells isolated from bone marrow and spleens were depleted of red blood cells with NH₄Cl lysis buffer, and FC receptors were blocked using anti-CD16/CD32 (BD Biosciences; 2.4G2). Single-cell suspensions from bone marrow cells or splenocytes were stained in PBS containing 2% BSA, 0.5 mM EDTA with BUV395-conjugated anti-B220 (clone RA3-6B2), APC- or PE-conjugated anti-IgM (clone 11/41), BV421- or APC-conjugated anti-CD43 (clone S7), PE-conjugated anti-Igκ (clone 187.1), and FITC-conjugated anti-Igλ (clone R26-46) antibodies. Dead cells were stained with LIVE/DEAD Fixable Aqua Dead Cell Stain Kit (Invitrogen). Data were acquired on an LSRFortessa (BD Biosciences) using Diva software (BD Biosciences) and analyzed using FlowJo software. Cell sorting was conducted using a MoFlo Astrios (Beckman Coulter, Inc.) using PE-Cy7 anti-mouse CD19 and FITC anti-mouse CD43 to identify preB cells.

Taqman PCR quantification of Ig light chain recombination

Genomic DNA was isolated from sorted preB cells using the QiaAMP DNA mini kit (Qiagen). Taqman quantitative PCR (qPCR) assays were performed using conditions previously described (Gopalakrishnan et al., 2013) and the following reagents: Vk forward primer, Jk1 reverse primer, and Jk1 probe or Vλ1 forward primer, Jλ1 reverse primer, and Jλ1 probe (Integrated

DNA Technologies; Table S9). *Vk-Jkl* and *Vλ1-Jλ1* rearrangements were normalized to an unrearranged region of the genome (CD19; using primers CD19 Forward, CD19 Reverse, and CD19 probe).

Quantification of germline *Jλ1* transcripts

Total RNA was isolated from sorted preB cells using the Trizol Reagent (Ambion). RNA was incubated with DNase (Promega) to destroy contaminating DNA. cDNA was synthesized using Pro-toScript First Strand cDNA synthesis Kit (New England BioLabs) as described (Brady et al., 2010). Primers used to detect *Jλ1* germline transcripts were the *Jλ1* Forward 1 and *Jλ1* Reverse 1 primers (Table S9). Levels of *Jλ1* transcripts were calculated using $\Delta\Delta C_t$ analysis with values normalized to levels of *HPRT*.

Quantification of germline *Jλ1* accessibility

Nuclei were isolated from the sorted preB cells and resuspended into 1× restriction digestion buffer (Mandal et al., 2015). Samples were divided into two equal parts and either treated as undigested control or digested with 50 U of *Bse*YI restriction enzyme (New England BioLabs) in 100 μ l volume at 37°C for 1 h. Genomic DNA was extracted using QiaAMP DNA mini kit (Qiagen). The isolated DNA was subjected to quantitative analysis by SYBR Green PCR using the primer pairs flanking the *Bse*YI restriction site. *CD19* was used to normalize the input DNA. *ICAM2* served as a control reference. Primers used were *CD19* Forward, *CD19* Reverse, *ICAM2* Forward 2, *ICAM2* Reverse 2, *Jλ1* Forward 2, and *Jλ1* Reverse 2 (Table S9).

Primary preB cell culture

Primary bone marrow cells were harvested by flushing bone marrow from all leg bones of at least three mice of the appropriate genotype for each experiment. These bone marrow cells were cultured for 4 d in RPMI 1640 supplemented with 10% FBS, 10 mM Hepes, 13 nonessential aa, 1 mM L-glutamine, 1 mM sodium pyruvate, 100 U/ml penicillin-streptomycin, 50 mM 2-ME, and 5 ng/ml IL-7. Cells were plated at a density of 5 million cells per milliliter of media. Each 2 d, cells were harvested and put back into culture in fresh media at a density of 5 million cells per milliliter. After 4 d of culture, B cells were sorted by depletion using EasySep mouse B cell isolation kit according to the manufacturer's instructions (StemCell Technologies, Inc.). PreB cell purity was assessed by flow cytometry and was above 90% (live B220⁺CD43⁺IgM⁺ cells). To induce G1 arrest and activate transcription of *Rag1* and *Rag2* by IL-7 withdrawal, we pelleted cells by centrifugation, resuspended them in the same media lacking IL-7 at a density of 2 million cells per milliliter. Cells were harvested after 48 h.

Real-time PCR quantification of *Pim2* mRNA and *DQ52*, *D β 1*, and *D β 2* transcripts

For *Pim2* (Fig. 3 B), 3 million cultured preB cells were harvested at indicated time points and immediately lysed in TRIzol (Life Technologies). For *DQ52* transcripts (Fig. S2 F), CD19⁺ cells were purified from either *RAG1*^{-/-} or R1.D708A total bone marrow using the EasySep Mouse CD19 Positive Selection Kit II (cat no. 18954) according to the manufacturer's directions and

immediately lysed in TRIzol. For *D β 1* and *D β 2* transcripts (Fig. S2 G), total thymocytes from either *RAG1*^{-/-} or R1.D708A mice were isolated and lysed in TRIzol. Total RNA was isolated using RNeasy mini kit (Qiagen), treated with DNase (RNase-Free DNase Set; Qiagen), and reverse transcribed to generate cDNA with High-Capacity RNA-to-cDNA Kit (Thermo-Fisher Scientific) according to the manufacturer's directions. The cDNAs were then used as a template for real-time PCRs performed with SYBR Green Mastermix (Applied Biosystems) and run on a Quant Studio Flex 7 machine using the following corresponding primers: *Pim2* Forward, *Pim2* Reverse, *DQ52* Forward, *DQ52* Reverse, *D β 1* Forward, *D β 1* Reverse, *D β 2* Forward, and *D β 2* Reverse primers (Table S9). *Pim2* and *DQ52* values were normalized to *CD19* using *CD19* forward and *CD19* reverse primers. *D β 1* and *D β 2* values were normalized to *Lck* using *Lck* Forward and *Lck* Reverse primers. Fold change was determined by $\Delta\Delta C_t$ analysis.

HTGTS-Rep-seq B cell repertoire library construction

B cells were isolated from mice aged 5–7 wk. Splenic B cells were isolated using B220 magnetic beads (Miltenyi Biotec) or by FACS sorting. B220⁺IgM⁺ cells were purified via FACS sorting from bone marrow. Genomic DNA samples were obtained using phenol-chloroform extraction of whole-cell lysates. HTGTS-Rep-seq libraries were prepared as previously described (Hu et al., 2016; Lin et al., 2016). Briefly, genomic DNA was sonicated and subjected to linear amplification-mediated PCR using biotinylated *J β 1* or *J β 3* bait primers (Lin et al., 2016). Linear amplification-mediated PCR products were purified using Dynabeads MyONE C1 streptavidin beads (Life Technologies; 65002) and ligated to bridge adaptors. Adaptor-ligated products were amplified by nested PCR with indexed *J β 1*/*J β 3* primers (8-bp indexes were unique to each sample; Table S9) and primer annealed to the adaptor. PCR products were further tagged with Illumina sequencing adaptor sequences and size-selected via gel extraction. Libraries were sequenced by paired-end 300-bp sequencing on an Illumina MiSeq by the Yale Center for Genome Analysis. Primers were previously described and are listed in Table S9 (Hu et al., 2016; Lin et al., 2016).

HTGTS-Rep-seq B cell repertoire analysis

To quantify *Igh* repertoires, an HTGTS pipeline was implemented that provides the all-inclusive steps for the analysis of HTGTS-Rep-seq data (Lin et al., 2016). In brief, the pipeline involves fastq-multx tool-based (<https://expressionanalysis.github.io/ea-utils/>) demultiplexing followed by the implementation of cutadapt (<https://cutadapt.readthedocs.io/en/stable/>) to trim the adaptors in the metadata file for each sample. The high-quality (Phred score >20) adaptor-trimmed reads were then joined using fastq-join tool if, and only if, the reads from both sides had at least a 10-bp overlap with mismatch rate $\leq 8\%$. Both joined and unjoined reads were mapped against the mouse (mm10) IGH V(D)J gene database (IMGT) using default parameters of the IgBLAST. The aligned reads (now assigned with V, D, and J genes) were further filtered to have high IgBLAST score >150 with total alignment length >100, comprising overall mismatch ratio <0.1. The usage of V and D genes was computed based on the processed IgBLAST results

as provided by the pipeline. To quantitate DJ_H rearrangements, we implemented the transloc_pipeline (https://github.com/robinmeyers/transloc_pipeline) that involves D_H-to-J_H junction detection based on the Optimal Query Coverage algorithm, as previously described (Hu et al., 2016). The identified junctions were then intersected with expanded RSS locations (±40 bp) to annotate the region with their respective D and J gene segments. After gene segments were called, all quantitative analysis was conducted in R.

ImmunoSEQ B and T cell repertoire library building and repertoire analysis

DNA was extracted from FACS-sorted bone marrow B220⁺IgM⁺ cells and thymic CD4⁺CD8⁺CD3e^{lo} cells using the Qiagen DNeasy Blood & Tissue Kit (Qiagen; 69506). High-throughput DNA sequencing of rearranged *Igh* and *Tcrb* genes was performed using ImmunoSEQ and was performed by Adaptive Biotechnologies. Gene segment usage was analyzed by ImmunoSEQ Analyzer software (Adaptive Biotechnologies) and R.

Online supplemental material

Fig. S1 shows RAG1 mutant mouse sequencing tracks and protein and mRNA expression. Fig. S2 shows FACS gating strategies and germline transcription and genome accessibility in D708A mice. Fig. S3 presents HTGTS-Rep-seq data from splenic B cells. Fig. S4 shows individual D_H gene segment usage from HTGTS-Rep-seq experiments. Fig. S5 shows individual J_β and V_β gene segment usage from thymic ImmunoSEQ experiments. Table S1, Table S2, Table S3, Table S4, Table S5, and Table S6 detail data for *Igh* HTGTS-Rep-seq experiments from spleen and bone marrow. Table S7 and Table S8 show detailed data for *Igh* and *Tcrb* ImmunoSEQ experiments from bone marrow and thymus, respectively. Table S9 lists oligonucleotides used in these studies and their sequences.

Data availability

The accession no. for the HTGTS-Rep-seq datasets reported in this paper is Gene Expression Omnibus no. GSE180734. The ImmunoSEQ datasets reported can be found at <https://doi.org/10.21417/HAB2021JEM> and <https://clients.adaptivebiotech.com/pub/beilinson-2021-jem>.

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Author contributions: H.A. Beilinson, C.H. Bassing, and D.G. Schatz designed experiments, which were performed by H.A. Beilinson, R.A. Glynn, J. Xiao, E. Corbett, H. Saribasak, R. Arya, and C. Miot. H.A. Beilinson, A.D. Yadavalli, and D.G. Schatz designed computational analyses, which were performed by H.A. Beilinson and A.D. Yadavalli. J.M. Jones generously provided the R1.PG mouse generated by A. Bhattacharyya. J.M.R. Pongubala provided mentorship to A.D. Yadavalli. H.A. Beilinson, R.A. Glynn, C.H. Bassing, and D.G. Schatz assembled figures and wrote the paper with input from other authors.

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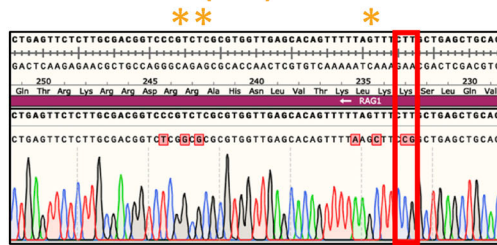
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Supplemental material

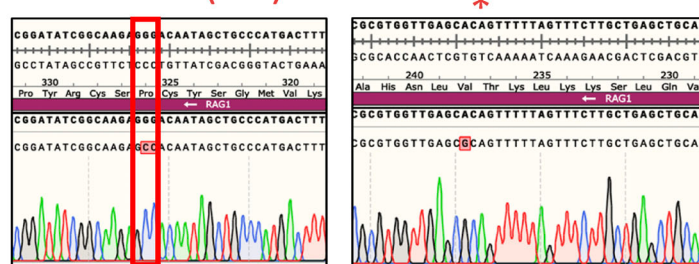
A R1Δ215



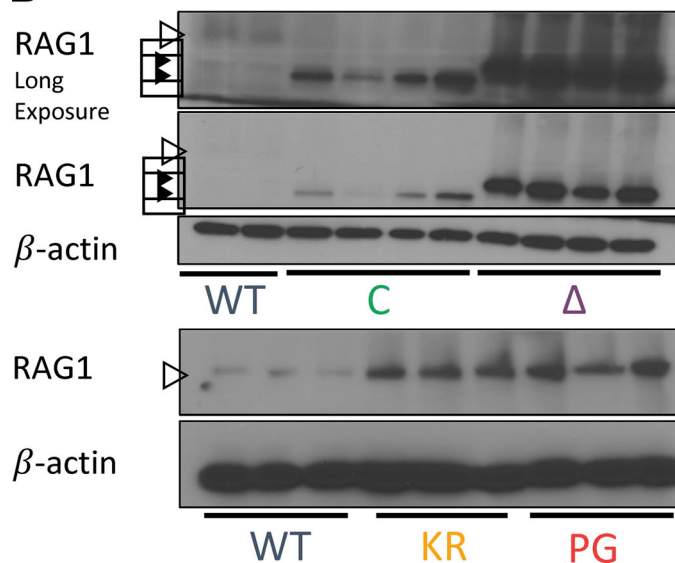
R1.K233R (KR)



R1.P326G (PG)



B



C

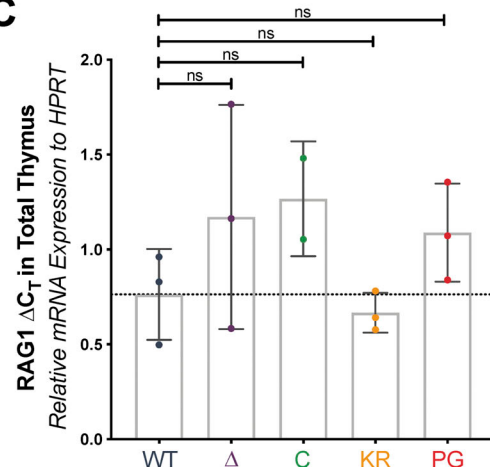


Figure S1. **RAG1 protein and mRNA expression in WT and RAG1 mutant mice.** (A) Sequencing tracks of PCR products amplified from the genomes of RAG1 NTR mutant mice. Red boxes indicate site of indicated mutation. Single asterisk indicates K234 and L235, where silent mutations were introduced to disrupt PAM site in R1.KR mice. Double asterisks indicate R243 and R244, where silent mutations were introduced to add a site for HindIII for genotyping purposes. The red asterisk indicates a passenger V238A mutation in R1.PG mice. This mutation is conservative, and V238 is not highly conserved and is replaced by alanine or isoleucine in several jawed vertebrates (Fig. S4 B from Brecht et al., 2020). It is not predicted to have an impact on RAG activity. (B) Western blot of RAG1 in whole thymic lysates from additional independent WT and RAG1 mutant mice. Open triangles, full-length protein; closed triangles, truncated proteins. (C) RAG1 mRNA expression in whole thymic lysates from mice of indicated genotypes, calculated relative to that of HPRT. Each dot indicates data from an independent mouse, with data presented as mean ± SEM. Statistical significance determined by two-tailed unpaired *t* tests (ns, *P* > 0.05; *, *P* ≤ 0.05; **, *P* ≤ 0.01; ***, *P* ≤ 0.001; ****, *P* ≤ 0.0001).

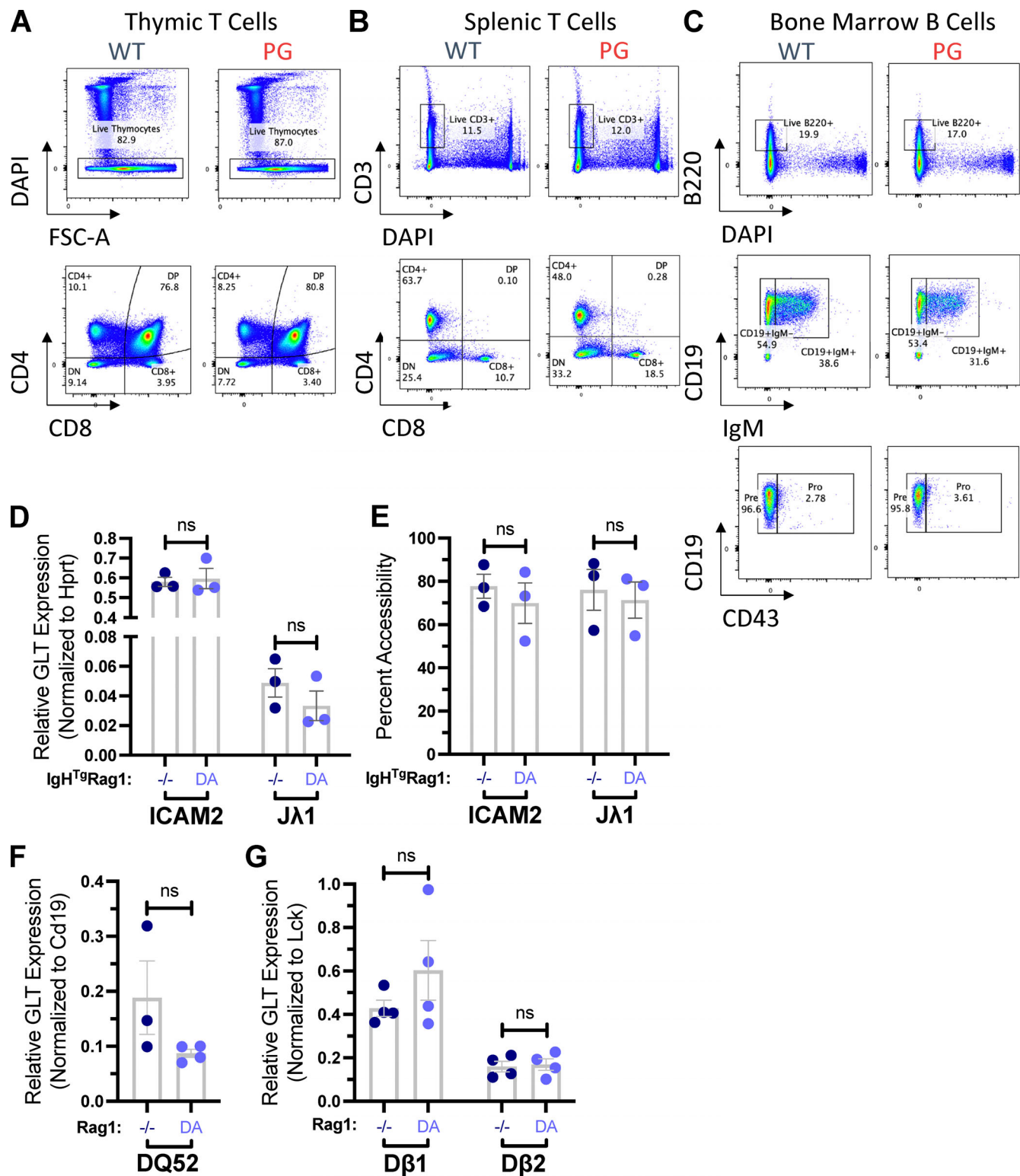


Figure S2. Flow cytometry analyses of B and T cell development in WT and R1.PG mice and characterization of germline transcripts in developing lymphocytes in WT and R1.D708A mice. (A–C) Representative flow cytometry data of WT and R1.PG mice showing the populations from thymic T cells (A), splenic T cells (B), and bone marrow B cells (C). (D–G) Quantification of RNA transcripts (D, F, and G) or DNA accessibility (E) of *ICAM2* or *Jλ1* (D and E), *DQ52* (F), or *Dβ1* and *Dβ2* (G) conducted on samples from three or four independent samples of sorted preB cells (D and E), CD19⁺ cells from total bone marrow (F), or total thymocytes (G) from *RAG1*^{-/-} and R1.D708A (DA) mice with (D and E) or without (F and G) an IgH transgene. Signals from each assay were normalized to values from assay of the HPRT (D), CD19 (F), or Lck (G) gene. Shown are mean values ± SEM. Statistical significance determined by two-tailed unpaired *t* tests (ns, *P* > 0.05; *, *P* ≤ 0.05; **, *P* ≤ 0.01; ***, *P* ≤ 0.001; ****, *P* ≤ 0.0001). FSC-A, forward scatter-A.

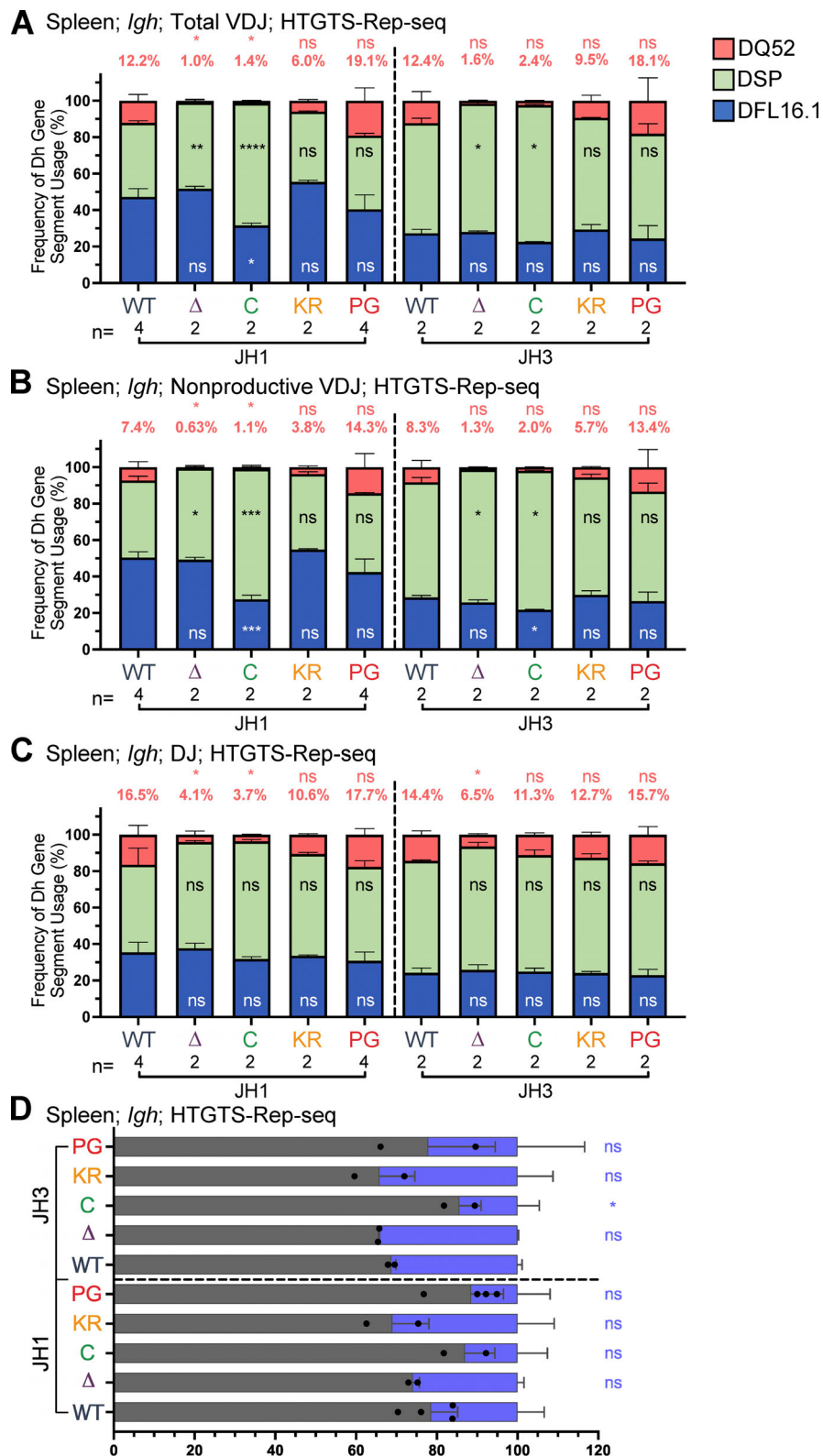


Figure S3. **D_H gene segment usage and analysis of productive and nonproductive rearrangements in VDJ_H rearrangements in spleen assessed by HTGTS-Rep-seq.** (A–C) *Igh* HTGTS-Rep-seq data from mice 5–7 wk of age were analyzed for frequency of DFL16.1 (blue), DQ52 (salmon), and intervening 11 D_H gene segments (green) in total (A) or nonproductive (B) VDJ_H or DJ recombination events (C) with J_{H1} or J_{H3} in spleen. Percentage of DQ52 usage is indicated above each bar. n, number of mice analyzed per genotype. (D) Frequency of productive (VDJ⁺) and nonproductive (VDJ⁻) recombination events at *Igh* in spleen with J_{H1} or J_{H3} using HTGTS-Rep-seq data. Number of recombination events analyzed can be found in Table S1, Table S2, and Table S3. Data are presented as mean with error bars indicating SEM. Statistical significance determined by two-tailed unpaired *t* tests (ns, *P* > 0.05; *, *P* ≤ 0.05; **, *P* ≤ 0.01; ***, *P* ≤ 0.001; ****, *P* ≤ 0.0001).

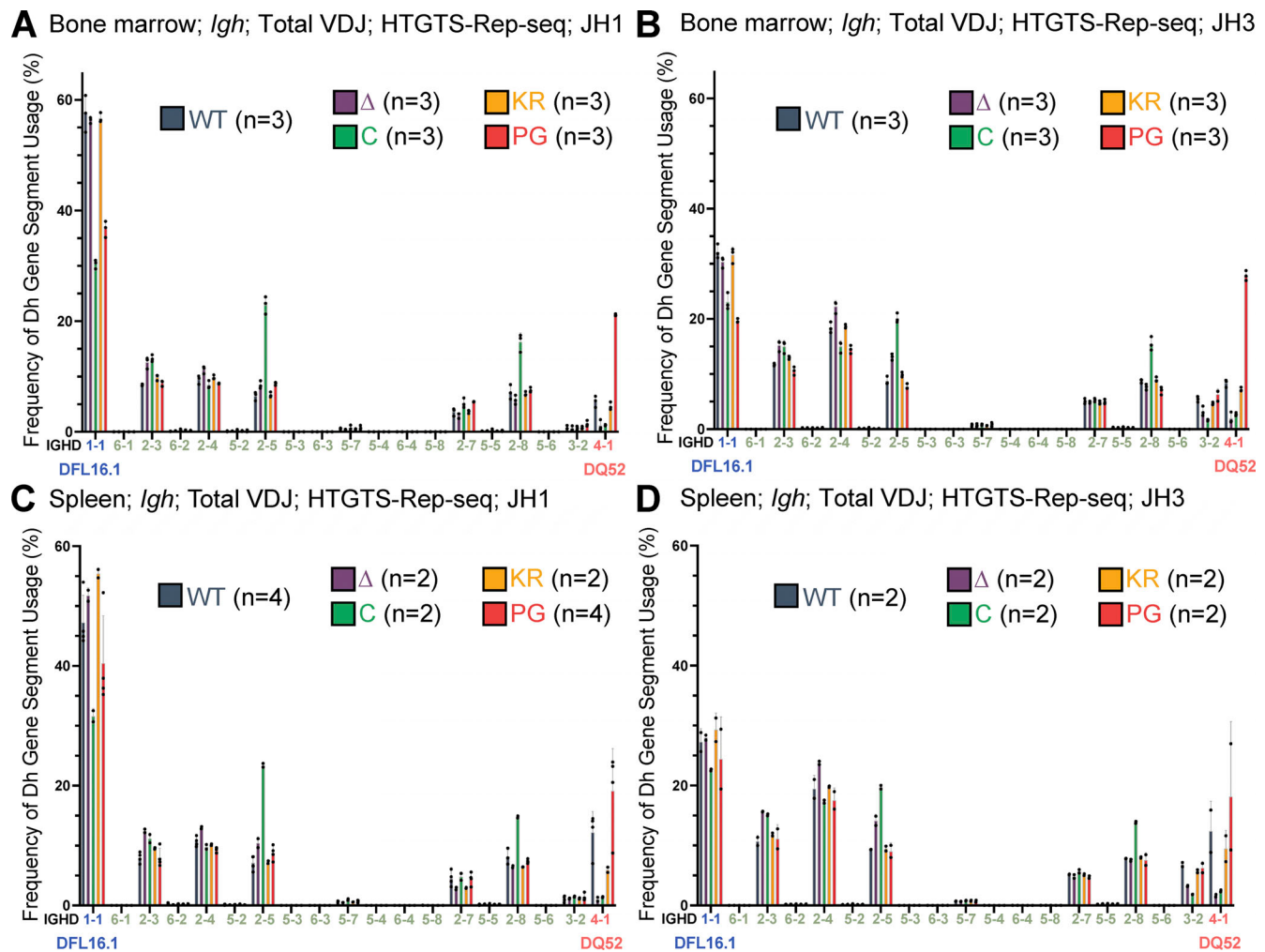


Figure S4. **D_H gene segment usage in total VDJ_H recombination events assessed by HTGTS-Rep-seq.** (A–D) Frequency of D_H gene segment usage in total VDJ recombination events in mice 5–7 wk of age with J_H1 (A and C) or J_H3 (B and D) in bone marrow (A and B) and spleen (C and D) using HTGTS-Rep-seq. n represents number of mice analyzed per genotype. Each dot indicates data from an independent mouse, with data presented as mean ± SEM. Number of recombination events analyzed can be found in Table S1, Table S2, Table S3, Table S4, Table S5, and Table S6.

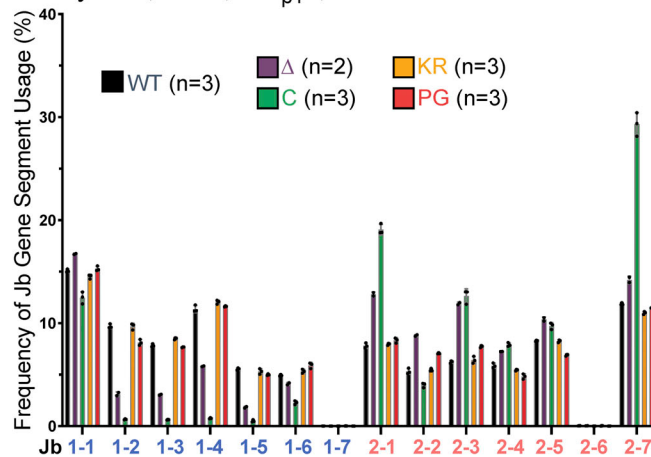
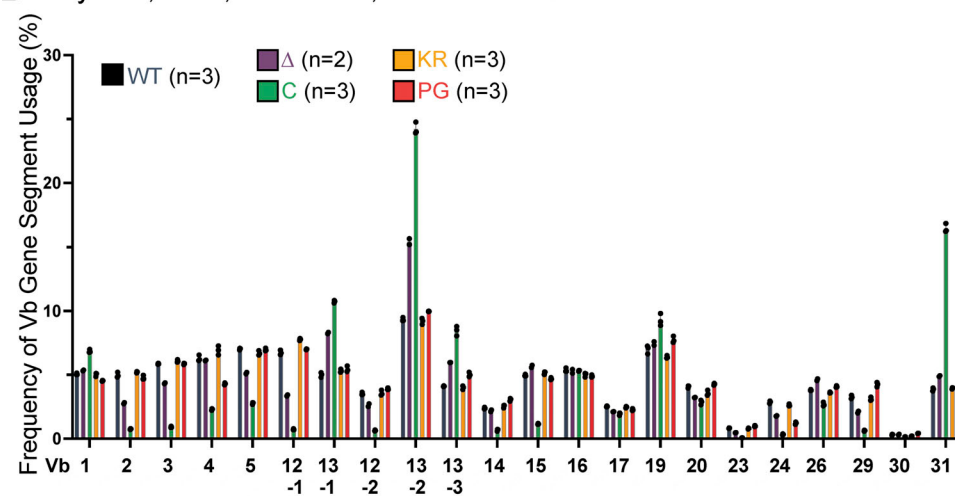
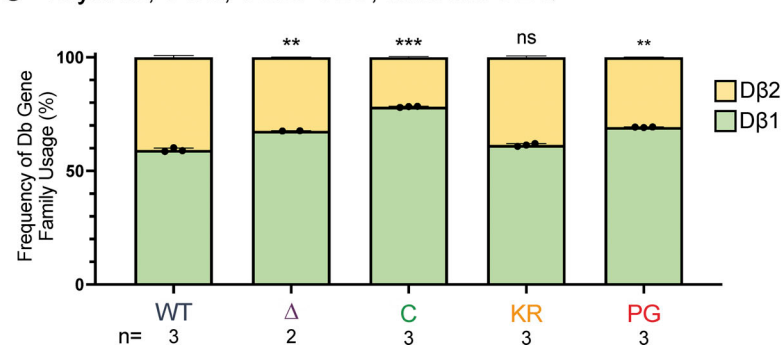
A Thymus; *Tcrb*; VD_{β1}J; ImmunoSEQ**B** Thymus; *Tcrb*; Total VDJ; ImmunoSEQ**C** Thymus; *Tcrb*; Total VDJ; ImmunoSEQ

Figure S5. **J_β, V_β, and D_β gene segment usage in VDJ_β recombination events in thymocytes assessed using ImmunoSEQ.** (A) Frequency of J_β gene segment usage in VDJ recombination events containing D_β1 in CD4⁺CD8⁺CD3^ε^{lo} thymic T cells. (B) Frequency of V_β gene segment usage in total VDJ recombination events in CD4⁺CD8⁺CD3^ε^{lo} thymic T cells. Repertoires were analyzed in mice of 5–7 wk of age. n represents number of mice analyzed per genotype. (C) Frequency of D_β1 and D_β2 gene segment usage in total VDJ_β recombination events. Each dot indicates data from an independent mouse, with data presented as mean ± SEM. Number of recombination events analyzed can be found in Table S8. Statistical significance determined by two-tailed unpaired *t* tests (ns, *P* > 0.05; **, *P* ≤ 0.01; ***, *P* ≤ 0.001).

Provided online are nine Excel tables. Table S1 shows utilization of *Igh* gene segments in primary B220⁺ splenocytes (HTGTS-Rep-seq, experiment 1). Table S2 shows utilization of *Igh* gene segments in primary B220⁺ splenocytes (HTGTS-Rep-seq, experiment 2). Table S3 shows utilization of *Igh* gene segments in primary B220⁺ splenocytes (HTGTS-Rep-seq, experiment 3). Table S4 shows

utilization of Igh gene segments in primary B220⁺IgM⁻ bone marrow cells (HTGTS-Rep-seq, experiment 1). Table S5 shows utilization of Igh gene segments in primary B220⁺IgM⁻ bone marrow cells (HTGTS-Rep-seq, experiment 2). Table S6 shows utilization of Igh gene segments in primary B220⁺IgM⁻ bone marrow cells (HTGTS-Rep-seq, experiment 3). Table S7 shows utilization of Igh gene segments in primary B220⁺IgM⁻ bone marrow cells (ImmunoSEQ). Table S8 shows utilization of Tcrb gene segments in VDJ rearrangements in primary CD8⁺CD4⁺CD3ε^{lo} thymocytes (ImmunoSEQ). Table S9 lists oligos used and their sequences.