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Orientation-specific RAG activity in chromosomal loop domains contributes to *Tcrd* V(D)J recombination during T cell development

Lijuan Zhao,^{1*} Richard L. Frock,^{1*} Zhou Du,¹ Jiazhi Hu,¹ Liang Chen,² Michael S. Krangel,² and Frederick W. Alt¹

¹Howard Hughes Medical Institute, Program in Cellular and Molecular Medicine Children's Hospital Boston, Department of Genetics, Harvard Medical School, Boston, MA 02115

²Department of Immunology, Duke University Medical Center, Durham, NC 27710

T cell antigen receptor δ (*Tcrd*) variable region exons are assembled by RAG-initiated V(D)J recombination events in developing $\gamma\delta$ thymocytes. Here, we use linear amplification-mediated high-throughput genome-wide translocation sequencing (LAM-HTGTS) to map hundreds of thousands of RAG-initiated *Tcrd* D segment (*Trdd1* and *Trdd2*) rearrangements in CD4⁺CD8⁺ double-negative thymocyte progenitors differentiated in vitro from bone marrow-derived hematopoietic stem cells. We find that *Trdd2* joins directly to *Trdv*, *Trdd1*, and *Trdj* segments, whereas *Trdd1* joining is ordered with joining to *Trdd2*, a prerequisite for further rearrangement. We also find frequent, previously unappreciated, *Trdd1* and *Trdd2* rearrangements that inactivate *Tcrd*, including sequential rearrangements from V(D)J recombination signal sequence fusions. Moreover, we find dozens of RAG off-target sequences that are generated via RAG tracking both upstream and downstream from the *Trdd2* recombination center across the *Tcrd* loop domain that is bounded by the upstream INT1-2 and downstream TEA elements. Disruption of the upstream INT1-2 boundary of this loop domain allows spreading of RAG on- and off-target activity to the proximal *Trdv* domain and, correspondingly, shifts the *Tcrd* V(D)J recombination landscape by leading to predominant V(D)J joining to a proximal *Trdv3* pseudogene that lies just upstream of the normal boundary.

INTRODUCTION

The RAG endonuclease (RAG) initiates V(D)J recombination by introducing DSBs between a pair of variable (V), diversity (D), and joining (J) gene segments and flanking recombination signal sequences (RSSs) to generate a pair of blunt signal ends (SEs) and a pair of hairpin-sealed coding ends (CEs; Schatz and Swanson, 2011; Alt et al., 2013). Bona fide RSSs are composed of a conserved heptamer (consensus: 5'-CAC AGTG) and an AT-rich nonamer separated by nonconserved 12- (12RSS) or 23-bp (23RSS) spacers (Schatz and Swanson, 2011). Normal RAG targeting and cleavage occurs only at pairs of coding segments flanked, respectively, by 12RSSs and 23RSSs (Alt et al., 2013). After binding to a Y-shaped RAG heterodimer (Kim et al., 2015; Ru et al., 2015) and subsequent cleavage, the CEs and SEs are held in a postcleavage synaptic complex from which SEs are directly joined to each other and hairpin CEs are opened, processed, and joined to each

other (Schatz and Swanson, 2011). The joining steps occur via classical nonhomologous end joining (Alt et al., 2013).

V(D)J recombination occurs in early B and T lymphocyte development and is tightly regulated by modulating accessibility of V, D, and J RSSs to RAG (Alt et al., 2013). Prior studies have shown that V(D)J recombination is initiated from a recombination center (RC) where RAG is recruited by epigenetic modifications and other factors (Matthews and Oettinger, 2009; Desiderio, 2010; Ji et al., 2010). In the *immunoglobulin heavy chain* (*IgH*) locus, the initial RC appears to form over the proximal D and J_H region (Teng et al., 2015). The *IgH* locus contains a key V(D)J recombination regulatory element, termed intergenic control region 1 (IGCR1), which lies between the *IgH* V_H and D_H gene segments (Guo et al., 2011). IGCR1 regulates *IgH* V(D)J recombination in the context of lineage specificity, order, and proximal V_H feedback regulation (Guo et al., 2011; Lin et al., 2015). IGCR1 function in these contexts relies on a pair of divergently oriented CTCF-binding elements (termed CBE1 and CBE2). Mutation of both CBE1 and CBE2 abrogates all of these forms of regulation and results in strongly increased

*L. Zhao and R.L. Frock contributed equally to this paper.

Correspondence to Michael S. Krangel: michael.krangel@duke.edu; or Frederick W. Alt: alt@enders.tch.harvard.edu

Abbreviations used: BE, broken end; CBE, CTCF-binding element; CE, coding end; CIL, chromatin interaction loop; CJ, coding join; δ REC, δ -deleting element; DN, double negative; IGCR1, intergenic control region 1; LAM-HTGTS, linear amplification-mediated high-throughput genome-wide translocation sequencing; RAG, RAG endonuclease; RC, recombination center; RSS, recombination signal sequence; SE, signal end; SJ, signal join; TEA, T early α .

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utilization of the most proximal V_H (V_H81x) coupled with a major reduction in distal V_H utilization (Guo et al., 2011). The two IGCR1 CBEs have been suggested to cooperatively regulate *IgH* V(D)J recombination by limiting the activity of the DJ_H RC to a chromosomal loop domain containing the D and J_H segments (Lin et al., 2015) and insulating the activity from the V_H8 portion of the locus (Hu et al., 2015; Lin et al., 2015). Such a mechanism would ensure ordered rearrangement of D-to- J_H segments before appendage of a V_H segment (Lin et al., 2015).

A potentially new mechanistic aspect of RAG function involving directional, linear tracking was recently implicated through the study that used the linear amplification-mediated high-throughput genome-wide translocation sequencing (LAM-HTGTS) approach to follow RAG cleavage and joining events (Hu et al., 2015). This study showed that pairs of bona fide RSSs integrated into a variety of chromosomal loop domains at various sites across the genome promote robust RAG off-target activity at flanking cryptic target sites, with cleavage occurring between convergent CAC motifs and associated surrogate CEs. Such joining is directionally oriented with respect to CAC motifs used and confined within the specific convergently oriented CTCF-anchored loop domains containing the bona fide RSSs (Hu et al., 2015). The mechanism that drives such RAG off-target directional- and orientation-specific joining biases has been proposed to involve unidirectional RAG tracking over great linear distances after being activated in the context of paired bona fide RSSs within a RC. In progenitor (pro)-B cells harboring a DJ_H rearrangement, such RAG tracking is robust within a V(D)J recombination domain that extended from IGCR1 to just downstream of the DJ_H RC (Hu et al., 2015). Moreover, deletion of IGCR1 allowed this off-target activity to directionally extend from the DJ_H to the proximal V_H81x , resulting in dramatically increased overutilization of V_H81x in joining to the downstream DJ_H (Hu et al., 2015).

The *TCR δ* gene (*Tcrd*) lies within the locus encoding *TCR α* (*Tcra*) in a contiguous 1.5-Mb region of the 129 mouse strain (Carico and Krangel, 2015). The 3' portion of *Tcrd* consists of two Ds (*Trdd1* and *Trdd2*) upstream of two Js (*Trdj1* and *Trdj2*), followed by C δ . There are 16 V δ s, five of which lie in a proximal unique region upstream of the D δ s, and one of which is in an inverted orientation downstream of C δ (Carico and Krangel, 2015). Other V δ s, also used by *Tcra*, lie at greater distances from D δ s (Carico and Krangel, 2015). Unlike ordered *IgH* rearrangement, which generates D-to-J rearrangements before V-to-DJ rearrangements (Alt et al., 2013), *Tcrd* rearrangements in mice have been concluded to be disordered, with intermediate V-to-D, D-to-D, D-to-J, V-to-D(D), and D(D)-to-J joins (Chien et al., 1987; Migone et al., 1995; Carico and Krangel, 2015). Based on RAG binding, the *Trdd2-Trdj1* region has been indicated to contain the initiating *Tcrd* RC (Ji et al., 2010; Teng et al., 2015). *Tcrd* recombination relies on the E δ enhancer (Monroe et al., 1999) and occurs early in T cell development in

CD4⁻CD8⁻ double-negative (DN) thymocytes at the DN2/DN3 stage (CD44⁺CD25⁺/CD44⁻CD25⁺; Capone et al., 1998; Livák et al., 1999; Carico and Krangel, 2015). *Tcra* recombination is dependent on the E α enhancer downstream of C α and on the T early α (TEA) promoter region upstream of the *Traj* cluster and occurs in CD4⁺CD8⁺ double-positive thymocytes (Sleckman et al., 1997; Carico and Krangel, 2015). Like *IgH*, *Tcrd* also contains a pair of intergenic CBEs (INT1 and INT2) between the V δ s and D δ s, and INT2 makes a loop (termed a chromatin interaction loop [CIL]) with a downstream convergently oriented CBE within the TEA region (Chen et al., 2015). The INT1-2 CBEs have been implicated in regulating *Tcrd* and *Tcra*V(D)J recombination (Chen et al., 2015).

To gain additional insight into how RAG orchestrates V(D)J recombination within the *Tcra-Tcrd* locus, we now use the LAM-HTGTS approach to map genome-wide junctions from RSSs flanking *Trdd2* and *Trdd1* gene segments during early T cell development.

RESULTS

LAM-HTGTS detection of joining events involving RAG-initiated DSBs at *Trdd2*

To study mechanisms of RAG activity and V(D)J recombination control in the *Tcrd* locus, we performed LAM-HTGTS studies with primary DN2/DN3 T cell precursors that represent the developmental stage at which *Tcrd*V(D)J recombination occurs (Capone et al., 1998; Livák et al., 1999; Carico and Krangel, 2015). To gain further potential insights into the initiating *Tcrd* RC (Teng et al., 2015), we examined the published RAG ChIP-seq profiles at higher resolution and noted that the major peak of RAG binding lies over *Trdd2* (Fig. S1). Therefore, for initial application of LAM-HTGTS for *Tcrd*V(D)J recombination studies, we performed LAM-HTGTS studies using RAG-initiated DSBs at RSSs flanking one or the other sides of *Trdd2* as bait (Fig. 1, A–C; and Table S1). To generate large numbers of DN2/DN3 T cell precursors for individual libraries, we cultured bone marrow-derived WT hematopoietic stem cells on OP9-DL1 cells (Schmitt et al., 2004; Holmes and Zúñiga-Pflücker, 2009) in the presence of IL-7 and Flt3-L for 14 d (Huang et al., 2005; Zakrzewski et al., 2006).

To capture joins involving the 12RSS SE (*Trdd2*-12RSS-SE) broken ends (BEs), we used a bait primer 75 bp upstream of *Trdd2* (5'-Primer; Fig. 1 B). This primer also captures joins to downstream 23RSS CE (*Trdd2*-23RSS-CE) BEs, but only those on alleles that had not undergone a prior joining event to the upstream *Trdd2*-12RSS-CE (Fig. 1 B). We used a bait primer 64 bp downstream of *Trdd2* (3'-Primer) to capture joins of the 23RSS SE (*Trdd2*-23RSS-SE) BEs, and also upstream *Trdd2*-12RSS-CE BEs that occur on alleles that had not undergone a prior joining event to the downstream *Trdd2*-23RSS-CE (Fig. 1 C). For both the 5' and 3' *Trdd2* priming strategies, prey junctions with bait sequence lengths that corresponded to either SEs or CEs of the *Trdd2*

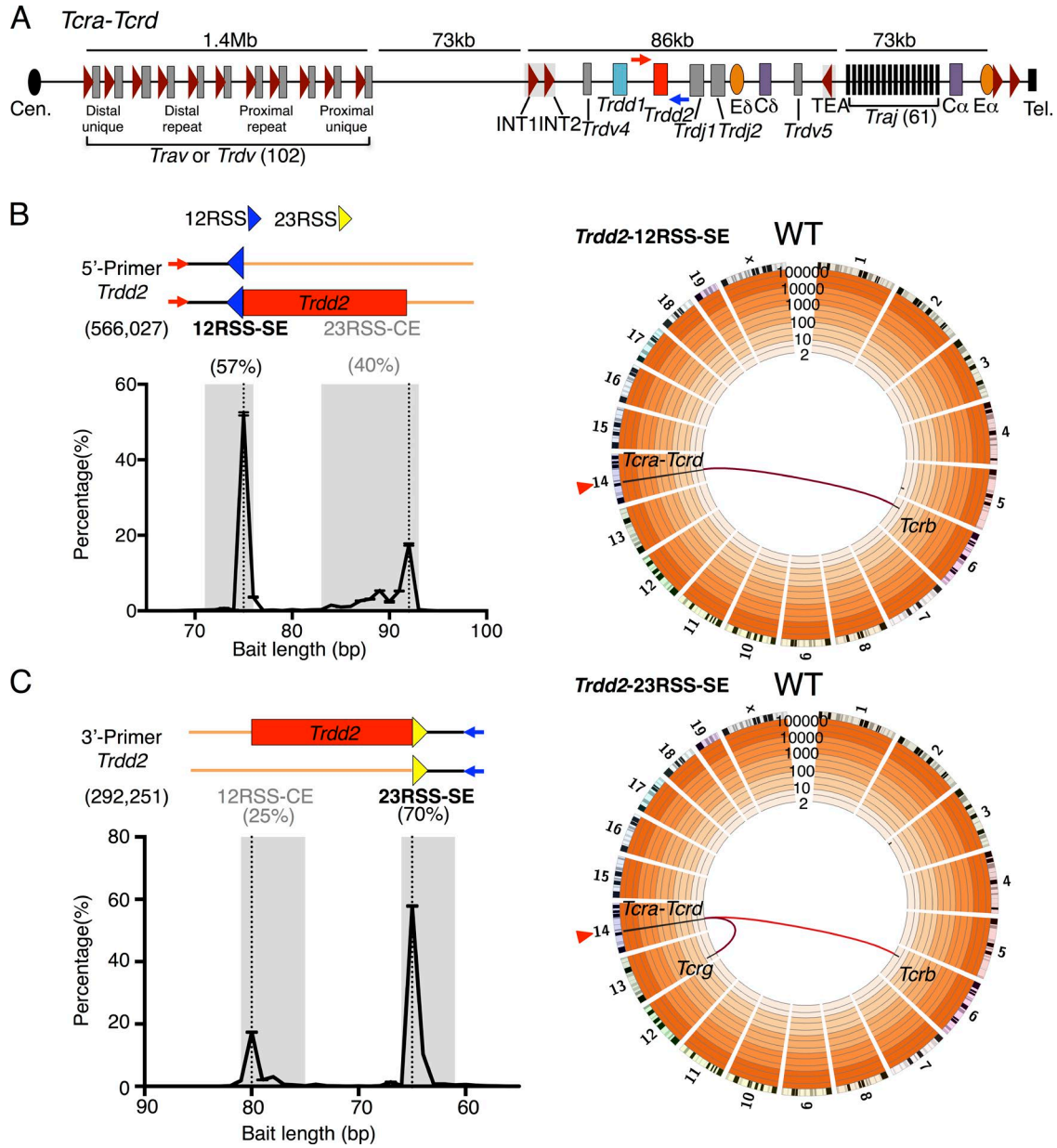


Figure 1. **Genome-wide translocation landscape of DN2/DN3 T cell progenitors.** (A) Schematic of murine *Tcra-Tcrd* locus. Dark red triangles represent position and orientation of CBEs. INT1-2 and TEA CBEs are labeled. Cen., centromere; Tel., telomere. 5'- and 3'-Primers (red and blue arrows, respectively) for generation of *Trdd2* LAM-HTGTS libraries are indicated. (B and C, top left) Junction reads are composed of the bait primer (red or blue arrows) and the bait sequence (black lines) leading up to the junction (encompassing bait and prey BE sequences), followed by the prey sequence (orange lines). Relative position of 12RSS (blue triangle), 23RSS (yellow triangle), and *Trdd2* gene segment (red rectangle) are indicated. (bottom left) Distribution of bait sequence length, plotted as the percentage of total junctions, corresponds to the relative position of the predicted bait break-sites (dotted lines). Numbers in parentheses denote total junctions analyzed. Ranges of bait sequence lengths of signal end (SE) and coding end (CE) bait libraries are shaded in gray (see Materials and methods for details). Junction percentages of total junctions in indicated ranges are shown above. 5'-Primer ($n = 4$; B); 3'-Primer ($n = 3$; C). (right) Circos plots displaying genome-wide prey junction distribution from either *Trdd2*-12RSS-SE ($n = 4$; B) or *Trdd2*-23RSS-SE ($n = 3$; C) libraries. Bin size is 5 Mb (black bars). *Trdd2* bait site (red triangle) and translocations from the bait to *Tcr* loci hotspots (red links) are indicated. *Trdd2*-12RSS-SE and *Trdd2*-12RSS-CE (Fig. S2) are normalized to 76,966 junctions. *Trdd2*-23RSS-SE and *Trdd2*-23RSS-CE (Fig. S2) are normalized to 106,193 junctions.

12 or 23 bait RSSs were identified (Fig. 1, B and C; Fig. S2, A and B; and Table S2). As anticipated, more junctions were recovered from primer-proximal SE baits than from

primer-distal CE baits (57 vs. 40% for 12RSS and 70 vs. 25% for 23RSS, respectively; Fig. 1, B and C) due to disordered V-to-D, D-to-D, and D-to-J joining of *Trdd2* (Chien et al.,

1987; Migone et al., 1995; Carico and Krangel, 2015). Given the expected underrepresentation of *Trdd2* CE bait junctions, we limit quantitative analyses to SE bait junctions. However, we observed reciprocal junction patterns for *Trdd2*-12RSS SE and CE baits and also for *Trdd2*-23RSS SE and CE baits, consistent with normal V(D)J joining (e.g., Figs. 2 and 3).

The highly precise joining of RAG-initiated SE junctions and limited diversity of CE junctions during normal V(D)J recombination tremendously limits junction diversity relative to that recovered, for example, with designer endonucleases. Therefore, we include duplicate junctions in our analysis of LAM-HTGTS libraries of RAG-initiated baits and obtain overall significance by analyzing at least three biological repeats of each experiment with a given bait or genetic background (Hu et al., 2015). We analyzed 323,910 *Trdd2*-12RSS-SE junctions (four libraries) and 76,966 *Trdd2*-12RSS-CE junctions (three libraries), as well as 203,485 *Trdd2*-23RSS-SE junctions (three libraries) and 229,666 *Trdd2*-23RSS-CE junctions (four libraries; Table S2). For both upstream and downstream *Trdd2* baits, >99% of recovered junctions were within the *Tcra-Tcrd* locus, with very low, but clear-cut, translocation junctions to *Tcrb* and *Tcrg* (Fig. 1, B and C; Fig. S2, A and B; and Table S3). Strikingly, though, in contrast to studies performed with designer nuclease bait DSBs in various cell types including DN2/DN3 T cell precursors (Hu et al., 2014; Frock et al., 2015; unpublished data), there were virtually no junctions recovered along the break-site chromosome or from other nonantigen receptor locus genomic sites in *Trdd2* bait libraries from WT T cell precursors (Fig. 1, B and C; Fig. S2, A and B; and Table S3).

RAG-initiated *Trdd2* SE and CE junctions in WT T cell precursors

The 12RSS of *Trdd2* can pair with bona fide 23RSSs of the *Trdd1* or *Trdv* gene segments (which all lie upstream of *Trdd2*) resulting in excision circle signal joins (Fig. 2 A) and deletional *Trdd1*-to-*Trdd2* or *Trdv*-to-*Trdd2* coding joins (Fig. 2 B). Both types of coding joins commonly contribute to the normal *Tcrd* repertoire in mice (Carico and Krangel, 2015). The 12RSS of *Trdd2* also can pair with the 23RSS of the inverted *Trdv5* that lies downstream of *Trdd2*, resulting in inversional signal joins and inversional *Trdv5*-to-*Trdd2* coding joins, respectively, which also inverts the intervening sequence between *Trdd2* and *Trdv5* that contains *Trdj1*, *Trdj2*, and *Cδ* (Fig. 2, A and B). We visualized overall RAG on-target patterns across *Tcra-Tcrd* via IGV plots (Fig. 2 C; Robinson et al., 2011; Hu et al., 2015). Junctions are displayed in + orientation if the prey sequence aligns in a centromere to telomere direction and in - orientation if prey sequence aligns in the opposite direction (Chiarle et al., 2011; Hu et al., 2015). *Trdd2*-12RSS-SE and *Trdd2*-12RSS-CE libraries identified all of the same bona fide RSS sites throughout the *Tcrd* locus, but, as expected for normal V(D)J recombination, occurred in + and - orientations, respectively (Fig. 2 C).

The most frequent class of joins recovered from the *Trdd2*-12RSS-SE libraries involved excision circle joining

between the *Trdd2*-12RSS-SEs and the 23RSS of *Trdd1* (56%; Fig. 2, A [2-SJ] and C; and Table S4). *Trdd2*-12RSS-SE libraries contained substantial numbers of excision circle signal joins to 23RSSs of upstream *Trdv* segments (25%; Fig. 2 A [1-SJ] and C; and Table S4) and also inversional signal joins to the downstream *Trdv5* 23RSS (8%; Fig. 2, A [3-SJ] and C; and Table S4). The most frequent upstream *Trdv*-to-*Trdd2* joins involved *Trdv2-2* (9%) and the *Trdv3* pseudogene (6%), which lie at the D-proximal end of the proximal unique region (Fig. 2 C). *Trdv2-2* has also been found to be one of the most frequently used *Trdvs* by repertoire sequencing (Passoni et al., 1997; Weber-Arden et al., 2000). Despite its location in the CIL and close proximity to *Trdd2*, *Trdv4* junctions occurred very rarely (<1%), possibly consistent with its preferential usage in fetal repertoires (Hao and Krangel, 2011). The *Trdd2*-12RSS-SE also revealed joining to five additional bona fide RSSs not associated with a known coding segment that lie upstream of *Trdd2* and therefore qualify as δ -deleting elements (δ RECs; Fig. 2 C), termed δ REC1, δ REC2, δ REC3, δ REC4, and δ REC5 (positioned from centromere to telomere). The strongest of these sites is δ REC3, which represents the previously described δ REC (de Villartay et al., 1988; Hockett et al., 1989), whereas the other four have not been previously described. δ REC2 and δ REC3 are conserved across multiple species, whereas δ REC1, δ REC4, and δ REC5 are conserved between mouse and rat (unpublished data). Notably, junctions involving the *Trdd2*-12RSS-CEs gave the same pattern of joins to all of these *Trdvs*, *Trdd1*, and δ RECs, but, consistent with normal V(D)J joining, in the opposite orientation (Fig. 2 C).

The 23RSS downstream of *Trdd2* can pair with a bona fide 12RSS from the upstream *Trdd1* to form deletional signal joins (Fig. 3 A [1-SJ]) and excision circle *Trdd2*-to-*Trdd1* coding joins (Fig. 3 B [1-CJ]). The *Trdd2* 23RSS can also pair with a 12RSS from the downstream *Trdjs* leading to excision circle signal joins and normal deletional DJ coding joins (Fig. 3, A [2-SJ] and B [2-CJ]). The *Trdd2*-23RSS-SE and *Trdd2*-23RSS-CE libraries again identified all of the same major bona fide RSS sites throughout the *Tcrd* locus, which, as expected for normal V(D)J recombination, occurred in - and + orientations, respectively (Fig. 3 C). The most frequent class of joins recovered were the excision circle signal joins between the *Trdd2*-23RSS-SEs and the 12RSS of *Trdj1* (78%; Fig. 3, A [2-SJ] and C) and the corresponding normal intra-chromosomal coding joins (Fig. 3, B [2-CJ] and C). However, there were few *Trdd2*-23RSS-SE joins to the 12RSS of *Trdj2* (<1%) consistent with earlier findings (Chien et al., 1987; Fig. 3 C). We also found a high frequency of *Trdd2*-23RSS-SE joins to the 12RSS of *Trdd1* (19%; Fig. 3, A [1-SJ] and C) and a high frequency of the reciprocal coding joining of *Trdd2*-to-*Trdd1* (26%) that results in the chromosomal deletion of both *Trdd1* and *Trdd2* coding gene segments within excision circles (Fig. 3, B [1-CJ] and C). This finding is striking and suggests this class of *Trdd* recombination could severely diminish repertoire diversity and,

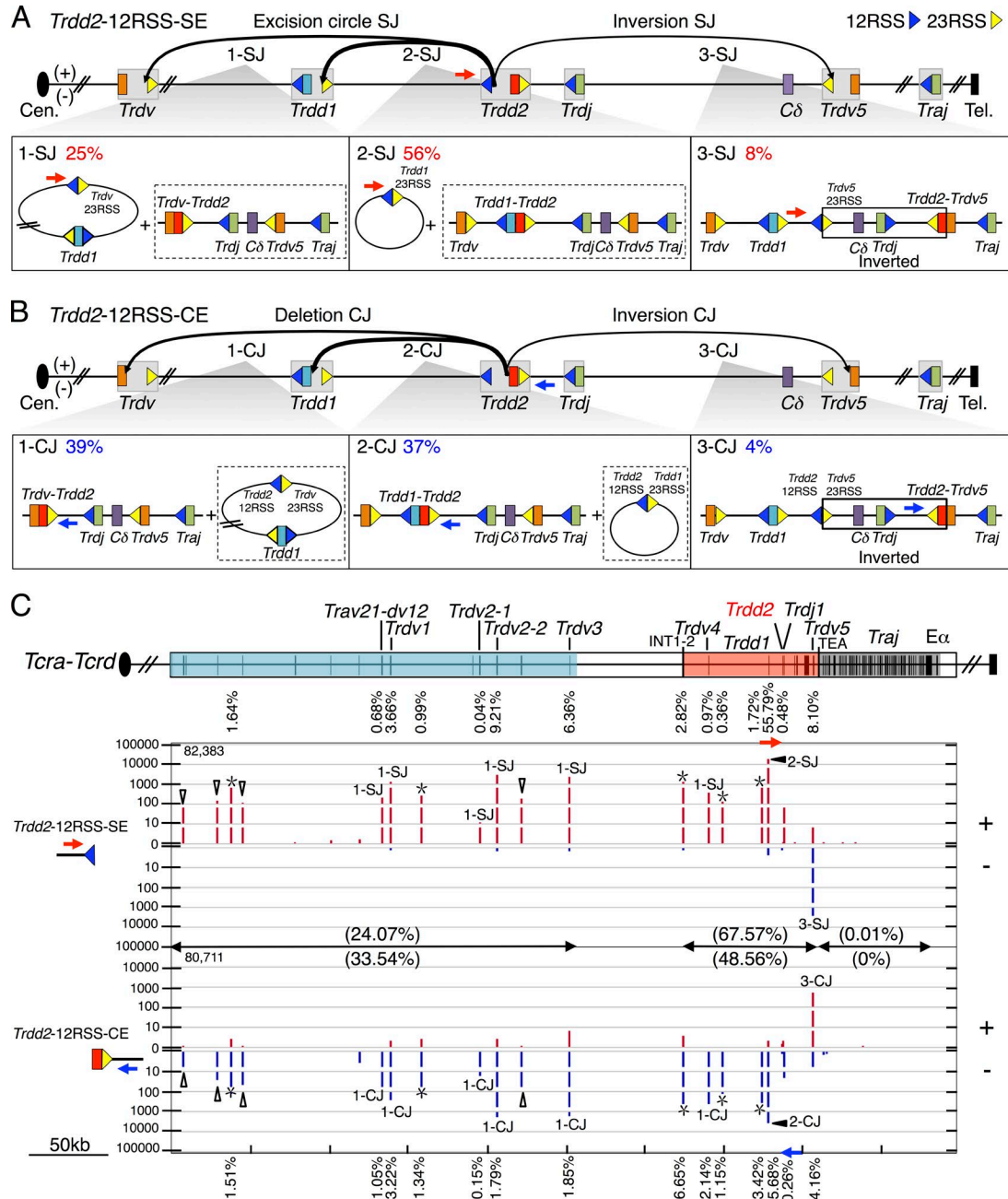


Figure 2. **Joining distribution of *Trdd2*-12RSS-SE and *Trdd2*-12RSS-CE libraries across the *Tcra*-*Tcrd* locus.** (A and B, top) Predicted joining outcomes of *Trdd2*-12RSS-SE (A) and *Trdd2*-12RSS-CE (B) baits. Gray boxes indicate gene segments with their flanking RSS(s): 12RSS and 23RSS are represented by blue and yellow triangles, respectively. The used primers are labeled with blue or red arrows. (bottom) Diagram of joining outcomes in excision circles plus deletional joins (1-SJ, 2-SJ, 1-CJ, and 2-CJ); and inversional joins (3-SJ and 3-CJ). The junction percentage of such joining outcomes as of total RAG on-targets are listed in the top left corner. Associated joining outcomes not detected by the primer listed are indicated in dashed box. Inverted regions in 3-SJ and 3-CJ are boxed. (C, top) Gene segment organization of the *Tcra*-*Tcrd* locus, organized in a centromere to telomere (+) chromosomal orientation. Proximal unique *Travs* and *Trdvs* (blue), the CIL (red), and the downstream *Traj* cluster (dark gray) are shown. Black vertical bars in the colored regions indicate the position of gene segments. (bottom) IGV plots displaying distribution of RAG on-targets (red in + and blue in - orientations) from pooled *Trdd2*-12RSS-SE ($n = 4$) and -CE libraries ($n = 3$). Junctions are displayed as stacked tracks (i.e., log scale between tracks, linear scale within each track) with the total number of RAG on-targets for the bait used are listed in the top left corner. Junction percentages for each indicated gene segment are labeled above or below their corresponding junction peaks. Open triangles mark *Trav* junctions and asterisks indicate δ REC (δ REC1, δ REC2, δ REC3, δ REC4, and δ REC5 from centromere to telomere). Junction percentages for each region are denoted in parentheses. *Trdd2*-12RSS-SE and -CE libraries are normalized to 83,572 total junctions.

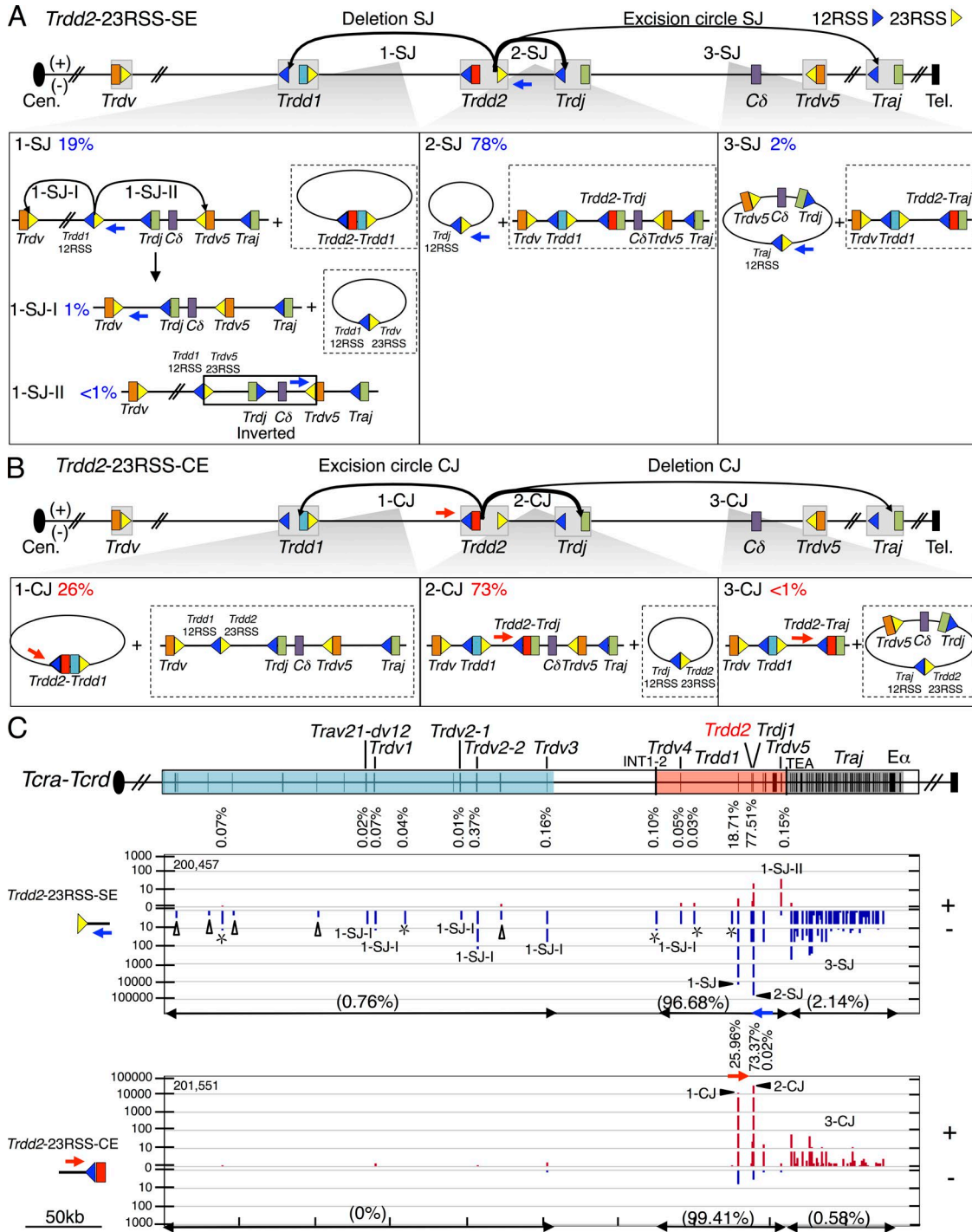


Figure 3. **Joining distribution of *Trdd2*-23RSS-SE and *Trdd2*-23RSS-CE libraries across the *Tcra-Tcrd* locus.** (A and B) Schematic of predicted joining outcomes for *Trdd2*-23RSS-SE (A) and *Trdd2*-23RSS-CE (B) bait libraries. Each class of joining events links to a lower box illustrating the corresponding joining outcomes in excision circles plus deletional joins (see Fig. 2 legend for other details). 1-SJ panel depicts secondary events from the *Trdd1*-12RSS-SE/*Trdd2*-23RSS-SE fusion (i.e., 1-SJ-I and 1-SJ-II). Inverted region in 1-SJ-II is boxed. (C) Gene segment organization of the *Tcra-Tcrd* locus is displayed in the top panel. IGV plot panels below show junction distribution of pooled *Trdd2*-23RSS-SE ($n = 3$) and -CE ($n = 4$) libraries (see Fig. 2 legend). *Trdd2*-23RSS-SE and -CE libraries are normalized to 202,196 total junctions.

given that direct *Trdv* to *Trdj* joining appears very infrequent (Table S4), potentially functionally inactivate the *Tcrd* locus. We also found a small fraction (2%) of excision circle *Trdd2*-23RSS-SE and *Trdd2*-23RSS-CE junctions to *Traj* 12RSS SEs and CE, respectively (Fig. 3, A [3-SJ], B [3-CJ], and C). Finally, we found very low levels of *Trdd2*23RSS-SE joins to *Trdv*-23RSS-CE (Fig. 3, A [1-SJ-I and 1-SJ-II] and C), which appear to arise via a 12/23 restricted intermediate (see the following section).

Profiles of RAG-initiated *Trdd1* SE and CE junctions in T cell precursors

We investigated joining patterns of DSBs generated at *Trdd1* bona fide RSSs by using 5' and 3' primers flanking *Trdd1*, as described above for *Trdd2* (Fig. S3 A and B; and Table S1; see Materials and methods). We analyzed 36,956 *Trdd1*-12RSS-SE junctions (three libraries) and 43,699 *Trdd1*-12RSS-CE junctions (three libraries), as well as 119,344 *Trdd1*-23RSS-SE junctions (three libraries) and 30,322 *Trdd1*-23RSS-CE junctions (three libraries; Table S2). Genome-wide joining patterns for all *Trdd1* baits used mostly were similar to those described above for *Trdd2* baits (Fig. S3, A and B; and Table S3).

Within the *Tcra-Tcrd* locus, we found a substantial level (15%) of excision circle *Trdd1*-12RSS-SE joins to 23RSSs associated with upstream proximal *Trdv* segments, most notably to *Trdv2-2* and *Trdv3* (Fig. 4, A [1-SJ] and C), with δ RECs (4%; Fig. 4 C), and also with the downstream inverted *Trdv5* (6%; Fig. 4, A [3-SJ] and C). However, recovery of *Trdd1*-12RSS-CE joins to *Trdv* segments were surprisingly rare (4%; Fig. 4 B [1-CJ and 3-CJ]), which in combination with the total *Trdd1*-12RSS-SE junctions involving *Trdv* 23RSSs (>20% of junctions; Fig. 4 A [1-SJ and 3-SJ]) raises the possibility that *Trdd1* joining is, in fact, ordered. Consistent with this notion, the vast majority of joins recovered from *Trdd1*-23RSS-SE libraries involved excision circle joining between *Trdd1*-23RSS-SE and the 12RSS of *Trdd2* (99%; Fig. 5, A [1-SJ] and C) with surprisingly few *Trdd1*-23RSS SE joins to the 12RSS of *Trdj* (<1%; Fig. 5, A [2-SJ] and C). Although we observe apparent coding joining to *Trdj1* from the *Trdd1*-23RSS-CE, this joining is not direct, as these junctions contain intervening *Trdd2* coding sequence (Fig. 5 B [1-CJ-I]). We also addressed this finding by analyzing *Trdj1*-12RSS-SE bait libraries (404,591 junctions; three libraries; Tables S1 and S2). These studies confirmed that <1% of *Trdj1*-12RSS-SE junctions joined directly to *Trdd1*, whereas 99% joined directly to *Trdd2* (Table S4). As expected, *Trdd1-Trdd2-Trdj1* sequential joining was also detected from *Trdj1*-12RSS-CE libraries (40,274 junctions; three libraries; Tables S1, S2, and S4). Overall, this set of findings indicates that for *Trdd1* segments with an intact 23RSS, *Trdd1* 12RSS rarely joins to *Trdvs*; moreover, the *Trdd1* 23RSS rarely joins to *Trdj*. Therefore, in contrast to prior expectations, it appears that most mature VDJ junctions involving *Trdd1* arise from a *Trdd1*-to-*Trdd2* intermediate via an ordered, as opposed to disordered, joining process.

We also found several *Trdd1* joining patterns, some of which are major, that had not been previously recognized. First, we found that the 12RSS of *Trdd1* also joins to the 23RSS of *Trdd2* to generate a substantial level (23%) of deletional signal joins, which would leave the two fused 12RSS/23RSS SEs in the chromosome (Fig. 4, A [2-SJ] and C). Remarkably, most (94%) of the recovered *Trdd1*-12RSS-CE joins are to the 23RSS CE of *Trdd2*, which resulted in deletion of the fused *Trdd1* and *Trdd2* segments within excision circles, and thus would severely reduce repertoire diversity (Fig. 4, B [2-CJ] and C). Although these 12RSS SE and CE joins would be expected to be reciprocal products, their differential recovery (23 vs. 94%) is striking. An explanation, however, comes from our finding that 50% of recovered *Trdd1*-12RSS-SE junctions occur at the CE of *Trdj1* (Fig. 4, A [2-SJ-I] and C), and appear to represent secondary rearrangements of the fused *Trdd1*-12RSS-SE/*Trdd2*-23RSS-SE junctions that explain the lower than expected recovery of the latter junctions. Although these junctions might first suggest apparent hybrid joins that seem to break the 12/23 rule, they appear to arise as a secondary recombination event of the fused *Trdd1*-12RSS-SE/*Trdd2*-23RSS-SE (Fig. 4 A [2-SJ]; and Fig. S4 A) in which the *Trdd2*-23RSS SE subsequently paired with the *Trdj1*-12RSS SE and the *Trdd1*-12RSS SE is used as a surrogate CE to join to *Trdj1* (Fig. 4 A [2-SJ-I]; and Fig. S4 A). Consistent with this interpretation, the 12RSS side of most of these junctions shows end-processing expected for CE joining, which would then inactivate the RSS as a further RAG target (Fig. S4 A). We also observe evidence for this type of secondary V(D)J joining activity for *Trdd1*-23RSS-CEs joining to 23RSSs of *Trdv* segments (Fig. 5 B, X-I), which most likely occurs within excision circles harboring *Trdd1* that are generated via *Trdd2*-12RSS and *Trdv*-23RSS joins (Fig. 2 A [1-SJ]). The reciprocal joining outcome of excision circles harboring *Trdd1*-23RSS/*Trdd2*-12RSS fusions (Fig. 5 B, X-I) is readily detected from *Trdd2*-12RSS-SE and *Trdd1*-23RSS-SE libraries and explains why the observed frequency of *Trdd1*-to-*Trdd2* joining events is slightly higher in the SE library than in the corresponding CE library (Figs. 2, A [2-SJ] and B [2-CJ]; and 5, A [1-SJ] and B [1-CJ and 1-CJ-I]).

Direction- and orientation-specific joining of RAG off-target DSBs at the *Tcra-Tcrd* locus

LAM-HTGTS studies with *Trdd2*-12RSS and *Trdd2*-23RSS bait BEs also revealed thousands of lower level, but reproducible, off-target junctions that amounted to ~1% of the total junctions (Table S2). These off-target junctions were largely generated at convergent CAC motifs (Fig. 6, A-E; Fig. S5, A-D; and Table S5). In this regard, 85% of *Trdd2*-12RSS-SE junctions were joined perfectly to upstream convergent CAC motifs resulting in excision circle junctions (+ orientation; Fig. 6, A and E). Likewise, ~93% of the *Trdd2*-12RSS-CE junctions occur upstream and involve the surrogate CEs associated with convergent CAC motifs, resulting in end-processed deletional junctions (Fig. 6, B and E). Both types of

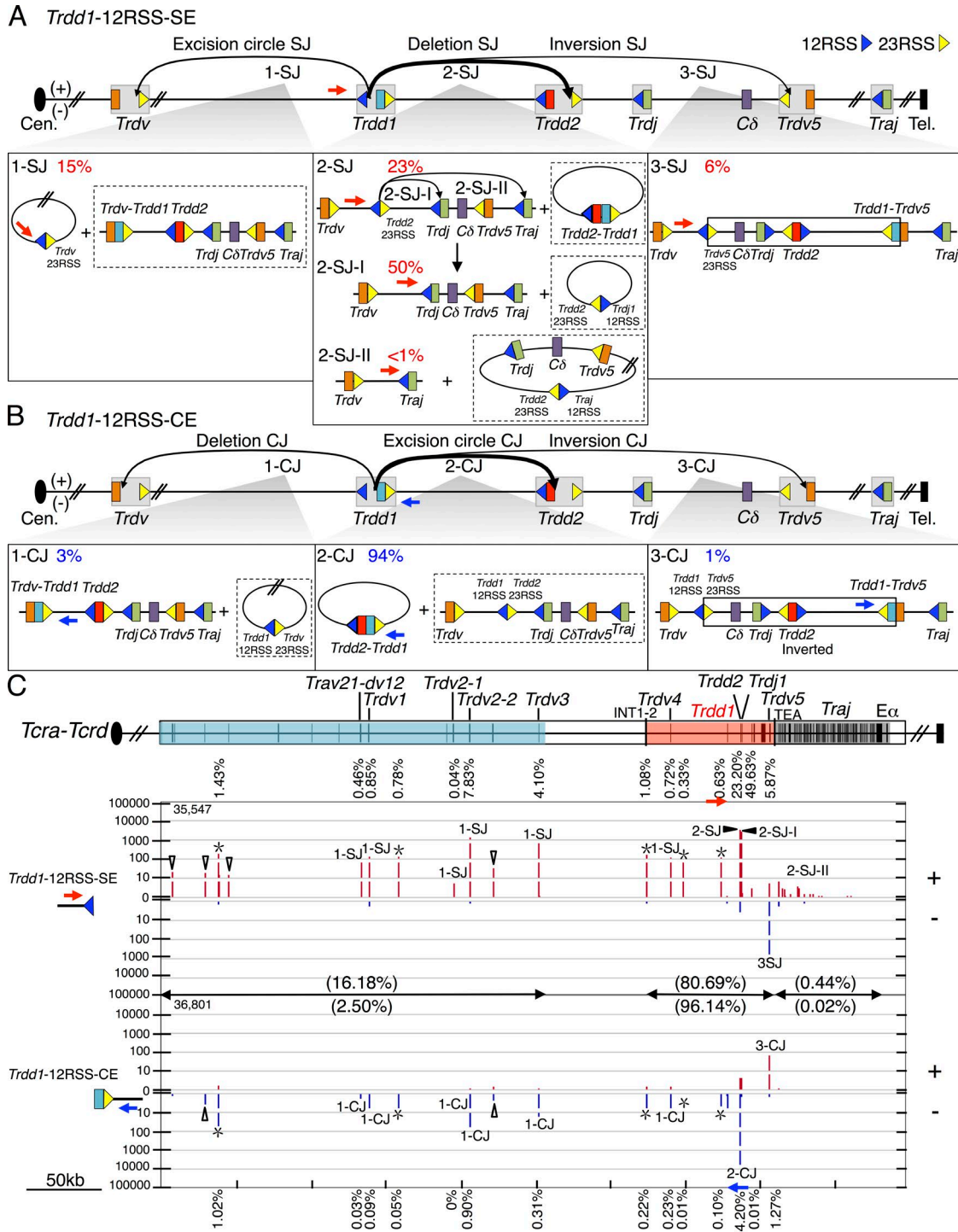


Figure 4. **Joining distribution of *Trdd1*-12RSS-SE and *Trdd1*-12RSS-CE libraries across the *Tcra-Tcrd* locus (A and B) Predicted joining outcomes for *Trdd1*-12RSS-SE (A) and *Trdd1*-12RSS-CE (B) bait libraries.** Labels (e.g., 1-SJ) for each class of joining events correspond to a lower box displaying joining outcomes and their corresponding percentages as of total RAG on-target joining events (see Fig. 2 legend). 2-SJ panel includes secondary events arising from *Trdd1*-12RSS-SE/*Trdd2*-23RSS-SE fusions (i.e., 2-SJ-I and 2-SJ-II). (C, top) *Tcra-Tcrd* gene segment organization. (C, bottom) Profile of RAG on-targets displayed by IGV from pooled *Trdd1*-12RSS-SE ($n = 3$) and -CE libraries ($n = 3$; see Fig. 2 legend). *Trdd1*-12RSS-SE and -CE libraries are normalized to 36,596 total junctions.

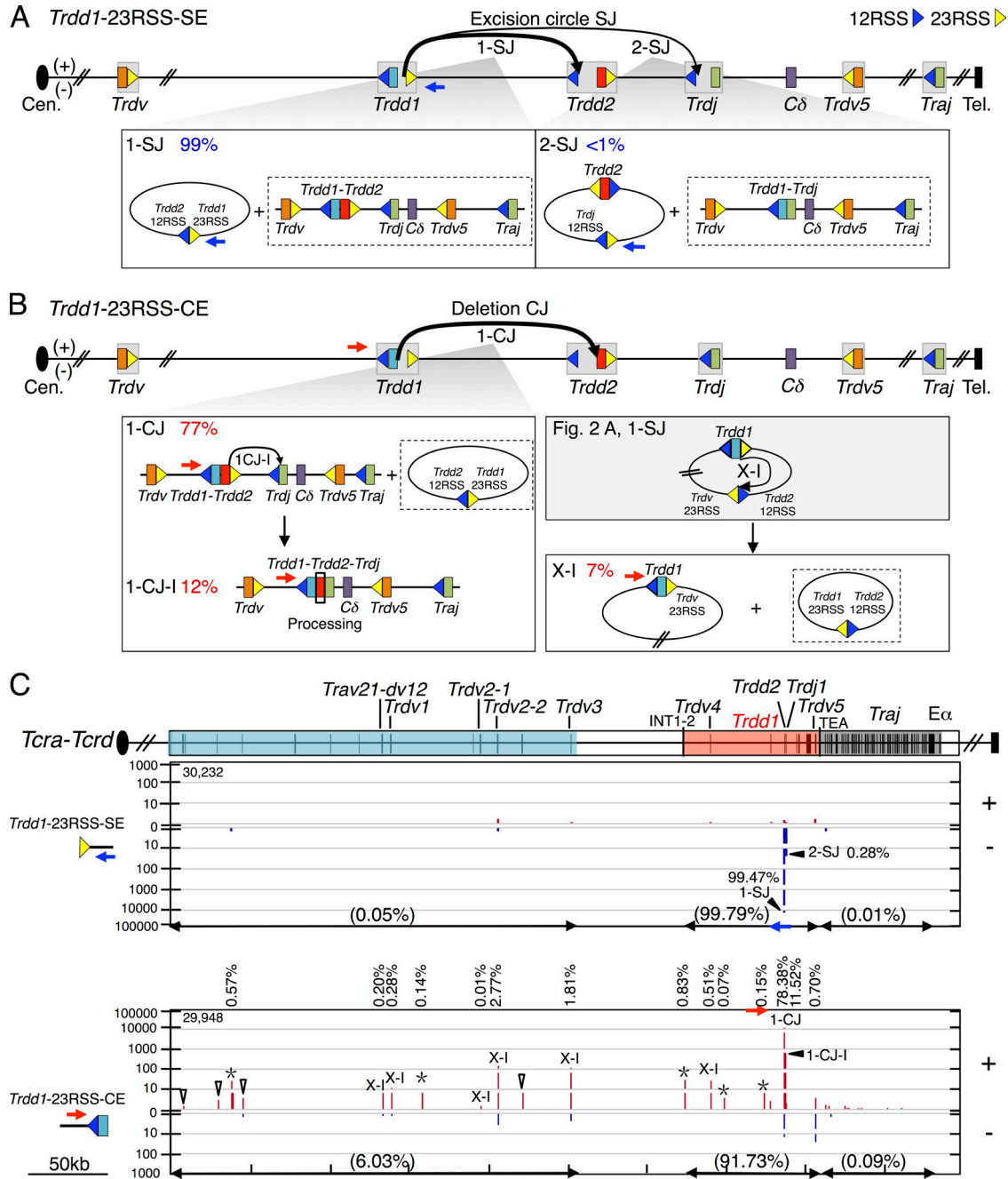


Figure 5. **Joining distribution of *Trdd1*-23RSS-SE and *Trdd1*-23RSS-CE libraries across the *Tcra-Tcrd* locus.** (A and B) Diagram of expected joining outcomes for *Trdd1*-23RSS-SE (A) and *Trdd1*-23RSS-CE (B) libraries. Each event links to a lower box showing joining outcomes resulting in excision circles plus deletional joins. (B) Event 1-CJ-I indicates hairpin processing on both sides of *Trdd2*. Event X-I comes from the excision circle harboring *Trdv*-23RSS and *Trdd2*-12RSS, described in Fig. 2 A. (C, top) Organization of gene segments in the *Tcra-Tcrd* locus. (bottom) IGV plots displaying distribution of RAG on-target junctions (red in + and blue in - orientations) from pooled *Trdd1*-23RSS-SE ($n = 3$) and -CE libraries ($n = 3$; see Fig. 2 legend). *Trdd1*-23RSS-SE and -CE libraries are normalized to 30,313 total junctions.

these upstream RAG off-target junctions stop abruptly at the CBE-containing INT1-2 elements (Fig. 6 E). A large fraction (69%) of *Trdd2*-23RSS SE junctions were perfectly joined to downstream convergent CAC motifs, resulting in excision

circle (- orientation) junctions; whereas the majority (90%) of *Trdd2*-23RSS-CEs joined downstream to surrogate CEs associated with these CACs to form deletional, end-processed junctions (Fig. 6, C-E). The downstream junctions also largely

terminate at the CBE-containing TEA element, which forms a convergent loop with INT2 (Fig. 6 E).

The majority (74%) of off-target *Trdd1*-12RSS-SE junctions occurred to upstream convergent CAC motifs (Fig. S5 E), resulting in excision circles and terminated at INT1-2 (Fig. S6, A and E). A smaller portion of *Trdd1*-12RSS-SE junctions (20%) occurred to downstream surrogate CEs resulting in deletions (Fig. S6, A and E); characteristics of these junctions, including *Trdd1*-12RSS end processing, indicate that they occur from the fused *Trdd1*-12RSS-SE/*Trdd2*-23RSS-SE junctions with the *Trdd1*-12RSS-SE acting as a surrogate CE (Fig. S4 A). *Trdd1*-12RSS CE, *Trdd1*-23RSS-SE, and *Trdd1*-23RSS-CE off-target joins were found much less frequently (Fig. S6, B–E), and apparently can be explained by ordered *Trdd1* joining to *Trdd2*.

These analyses also identified certain unanticipated types of off-target junctions. In this regard, we identified a surprisingly high frequency (25%) of deletional (– orientation) *Trdd2*-23RSS-SE junctions to surrogate CEs associated with upstream CAC motifs (Fig. 6, C and E). Notably, the 23RSS SEs involved in such joins frequently showed end processing (Fig. S4 C), consistent with functioning as surrogate CEs from fused *Trdd1*-12RSS/*Trdd2*-23RSS junctions (see above). We also found a low, but reproducible, level of *Trdd1*-23RSS-CE junctions to upstream CACs that are used as surrogate CEs (Fig. S5 F; and Fig. S6, D and E); these junctions likely occur within *Trdd2*-12RSS/CAC excision circles harboring unrearranged *Trdd1* segments (Fig. S6 D).

INT1-2 blocks RAG-mediated joining to upstream neighboring domain

To assess the functional role of the INT1-2 CBEs in regulating RAG on- and off-target joining patterns, we performed LAM-HTGTS with *Trdd2*-12RSS and *Trdd2*-23RSS bait BEs on DNA from cultured INT1-2-deficient T cell precursors generated from homozygous INT1-2-deficient mice (Chen et al., 2015). The frequency of *Trdd2*-12RSS-SE joins to bona fide 23RSSs within the *Trdd2* to INT1-2 interval decreased in INT1-2-deficient T cell precursors versus those of WT, with the greatest decreases for *Trdd1* (threefold; Fig. 2 C; Fig. 7 A; and Table S4). We also observed a large decrease in the frequency of *Trdd2*-12RSS-SE joins to the downstream bona fide inverted 23RSS of *Trdv5* (sixfold; Fig. 2 C; Fig. 7 A; and Table S4). In contrast, the frequency of *Trdd2*-12RSS-SE junctions with the *Trdv3* 23RSS, which lies 73 kb upstream of the INT1-2 locale, markedly increased (10-fold) in INT1-2-deficient T cell precursors relative to those of WT (Fig. 2 C; Fig. 7 A; and Table S4). We also found low level, but reproducible, *Trdd2*-12RSS-SE junctions to an apparently bona fide 23RSS ~1 kb downstream of *Trdv3* in INT1-2-deficient, versus WT, T cell precursors (Fig. 7 A and Table S4). *Trdd2*-12RSS-SE RAG off-target joins within the *Trdd2* to INT1-2 locale interval also decreased in INT1-2-deficient versus WT T cell precursors; however, off-targets then spread 73 kb upstream of the INT1-2 locale to *Trdv3*

(Fig. 6 E; Fig. 7 A; and Table S4). We also observed similar differences in RAG on- and off-target joining to the 12RSS CEs in INT1-2-deficient versus WT T cell precursors (Table S4). However, *Trdd2*-23RSS-SE and *Trdd2*-23RSS-CE RAG on- and off-target joining patterns were similar between INT1-2-deficient and WT, consistent with these joins mainly occurring downstream of *Trdd2* and, therefore, not being impacted by the upstream INT1-2 deletion (Table S4).

We also performed LAM-HTGTS with *Trdd1*-12RSS and *Trdd1*-23RSS bait BEs on genomic DNA from INT1-2-deficient T cell precursors. In the INT1-2-deficient T cell precursors, we find decreased joining to bona fide 23RSSs in the *Trdd1* to the INT1-2 locale interval and a corresponding increase in joining to the *Trdv3* 23RSS (12-fold) and to the bona fide 23RSS downstream of *Trdv3* (Fig. 4 C; Fig. 7 B; and Table S4). Likewise, RAG off-target joining was decreased within the *Trdd1* to INT1-2 locale interval in the INT1-2-deficient T cell precursors with off-target activity spreading to *Trdv3* (Fig. S6 E; Fig. 7 B; and Table S4). Although we rarely detected *Trdd1*-12RSS-CE joining to *Trdvs* in WT precursors, INT1-2 deficiency led to substantial joining of the *Trdd1*-12RSS-CE to *Trdv3* (12.49% of total bona fide joining), indicating increased disordered joining (Table S4). Again, *Trdd1*-23RSS-SE and CE joining patterns were not substantially altered in INT1-2-deficient T cell precursors (Table S4).

DISCUSSION

Our LAM-HTGTS studies provide strong confirmation of disordered *Trdd2* gene segment joining, which comes specifically from the detection of *Trdd2*-to-*Trdj1*, *Trdd2*-to-*Trdd1*, and *Trdd2*-to-*Trdv* as major V(D)J recombination intermediates from *Trdd2* CE baits. However, in contrast to *Trdd2* disordered joining, primary downstream joining of *Trdd1* occurred almost exclusively (99% of junctions) to the upstream CE of *Trdd2*, either to the germline *Trdd2* segment or to a *Trdd2*-*Trdj1* intermediate. These findings indicate that *Trdd1* joining is ordered and occurs to *Trdd2* before *Trdd1*-*Trdd2* joining to *Trdvs* via *Trdd1*-12RSS/*Trdv*-23RSS pairing. Ordered rearrangement of *Trdd1*, as opposed to the disordered rearrangement of *Trdd2*, can be mechanistically understood based on the observation that *Trdd2*, as opposed to *Trdd1*, is highly accessible in DN thymocytes (Carabana et al., 2005; Hao and Krangel, 2011) and appears to be the initiating RC (Teng et al., 2015). Thus, RAG bound to the *Trdd2*-12RSS may capture either *Trdd1* or *Trdv* gene segments, whereas the RAG-poor and weakly accessible *Trdd1*-12RSS would be much less likely to capture *Trdv* gene segments. A corollary to this would be that the upstream *Trdd1*-12RSS must be activated as a RAG substrate once brought into the RC by *Trdd1*-to-*Trdd2* rearrangement. Previous studies have identified germline *Tcrd* transcripts initiating within *Trdd2* and upstream of and within *Trdj1*, as well as a strong promoter associated with *Trdd2* (Carabana et al., 2005). This promoter is likely responsible for *Trdd2* accessibility and formation of

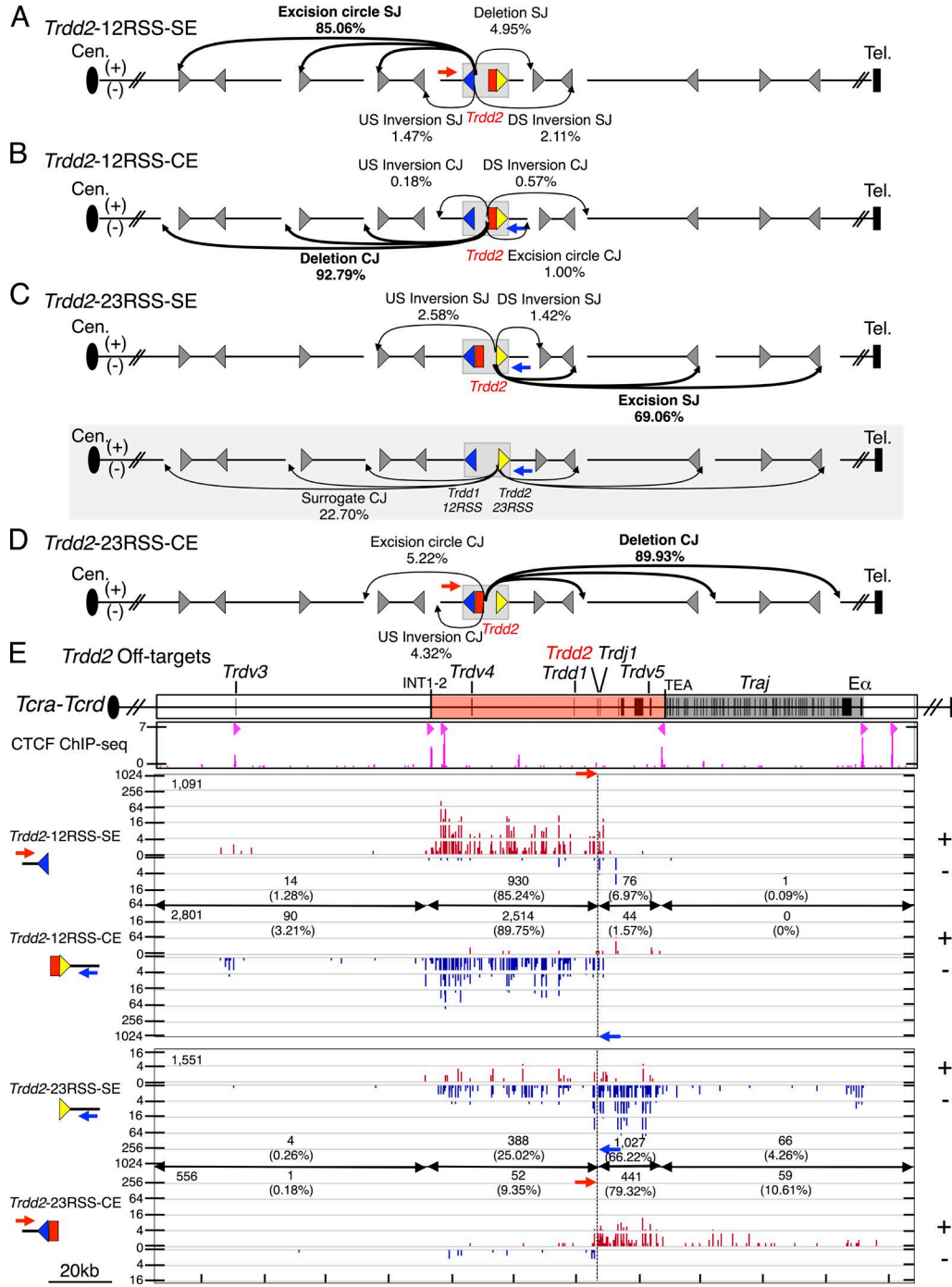


Figure 6. Direction- and orientation-specific off-target joining of RAG-initiated *Trdd2* rearrangements. (A–D) Joining outcomes to cRSS (CAC) sites (gray triangle) from *Trdd2*-12RSS-SE ($n = 4$; A), -CE ($n = 3$; B), *Trdd2*-23RSS-SE ($n = 3$; C), and -CE ($n = 4$; D) libraries. Diagram of joining outcomes from *Trdd2*-23RSS-SE bait in the fused *Trdd1*-12RSS/*Trdd2*-23RSS configuration to cRSSs is shaded in gray (C). Junction percentages as of total RAG off-targets in excision circle, deletion, upstream (US) inversion, and downstream (DS) inversion joining events are shown. Red and blue arrows indicate the position and orientation of 5'- and 3'-Primers, respectively. Dotted line indicates the position of bait BEs. (E, top) Gene segment organization of the region between *Trdv3* and *E α* (3' portion of the *Tcra-Tcrd* locus). CIL (red) and *Traj* cluster (dark gray) are indicated. (middle) ChIP-seq profile of CTCF. CBE orientation is marked by light red triangles. (bottom) IGV plots of RAG off-target joining patterns of *Trdd2*-12RSS-SE and -CE and *Trdd2*-23RSS-SE and -CE libraries. The total number of RAG off-targets is displayed on the top left corner of each plot. Junction numbers and percentages as of total RAG off-targets in indicated regions are shown.

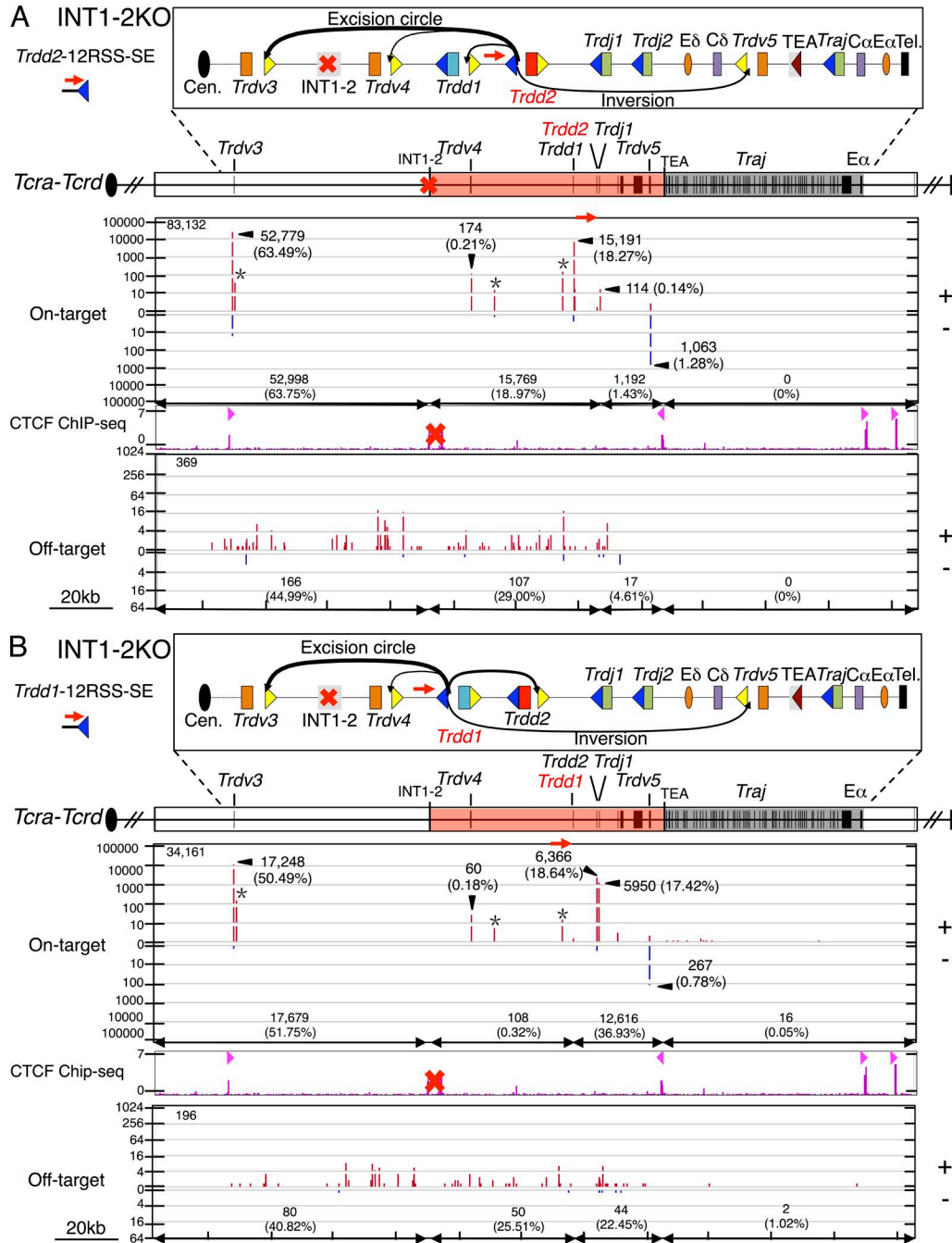


Figure 7. **INT1-2 CBEs block RAG tracking across the insulated neighboring domain.** (A and B, top) Gene segment organization of the 3' portion of the *Tcra-Tcrd* locus. Diagram of predicted RAG on-target joining outcomes of INT1-2-deficient *Trdd2*-12RSS-SE (A) or *Trdd1*-12RSS-SE (B) libraries is displayed in the box above. (bottom) IGV plots of RAG on- and off-target joining patterns for INT1-2 deficient *Trdd2*-12RSS-SE ($n = 3$; A) and *Trdd1*-12RSS-SE ($n = 3$; B) bait libraries. Profile of CTCF ChIP-seq is shown and CBE orientation is marked by light red triangles. Red "X" represents the INT1-2 deletion. INT1-2-deficient *Trdd2*-12RSS-SE and *Trdd1*-12RSS-SE libraries are normalized to the respective WT libraries described in Figs. 2 and 4. Total RAG on- and off-target numbers of each bait library are listed in the top left corner. RAG on- and off-target numbers and percentages as of the total number of each category in indicated regions are shown.

the *Tcrd* RC, but it would be disrupted by *Trdd1*-to-*Trdd2* rearrangement. Although analysis of unrearranged alleles identified no germline transcripts mapping to *Trdd1*, and no

substantial promoter activity associated with *Trdj1* (Carabana et al., 2005), it is possible that promoter activity associated with one or the other is enhanced by *Trdd1*-to-*Trdd2* re-

arrangement, allowing RAG-loading at the *Trdd1*-12RSS to stimulate *Trdv*-to-*Trdd1*-*Trdd2* rearrangement.

Beyond the well-defined δ REC 120 kb upstream of the *Trdd* gene segments (Janowski et al., 1997; Krangel et al., 1998), we also discovered additional, previously uncharacterized δ RECs that between them provide nearly 70% of the deletional activity of δ REC and, as a result of their location in the INT1-2-TEA loop or in the proximal unique region, generate variably sized truncations of the *Tcrd* locus upon joining to *Trdd* segments. We find that nearly 10% of all such *Tcrd*V(D)J rearrangements to the upstream 12RSS of *Trdd2* occur to this overall set of δ RECs, recombination events that would functionally diminish repertoire diversity and potentially severely impair *Tcrd* V(D)J recombination. Moreover, we also found an additional, even more frequent, mechanism that could lead to *Tcrd* inactivation. Joins between *Trdd1*-12RSSs and *Trdd2*-23RSSs contribute \sim 20% of all *Trdd2* 23RSS rearrangements. This joining event deletes both *Trdd1* and *Trdd2* segments, leaving in the chromosome the perfectly fused *Trdd1*-12RSS-SE/*Trdd2*-23RSS-SE. Such fused 12 and 23 RSSs can be recut and joined as surrogate CEs (Hu et al., 2015). In this regard, we find joining of the *Trdd1* 12RSSs to *Trdj1* and joining *Trdd2* 23RSS to *Trdvs*. However, in most of these junctions the *Trdd2* or *Trdd1* SEs serving as surrogate CEs are processed via normal CE junctional diversification mechanisms and, thus, are inactivated. Together, our findings directly support the notion that *Tcrd*V(D)J recombination evolved deletional mechanisms to developmentally delete *Tcrd* and, thereby, to promote appropriate V(D)J recombinational activation of the greater *Tcra* locus in which *Tcrd* is embedded (Chen et al., 2015).

Analyses of INT1-2-deficient mice the DN thymocytes revealed partial inhibition of $\gamma\delta$ T cell development and a twofold increase in usage of *Trdv2-2* (V δ 4), the most proximal functional *Trdv* in the unique region upstream of INT1-2 (Chen et al., 2015). Our current LAM-HTGTS analysis of in vitro differentiated DN2/DN3 thymocytes, the stage at which *Tcrd* rearrangement occurs (Capone et al., 1998; Livák et al., 1999; Carico and Krangel, 2015), confirmed a modest increase in *Trdv2-2* usage. However, our current studies further revealed a major 10-fold increase in usage of *Trdv3*, a pseudogene that is the most proximal upstream *Trdv* to INT1-2. The markedly increased usage of this *Trdv* pseudogene in primary rearrangements in the absence of INT1-2 offers a plausible explanation for the $\gamma\delta$ T cell developmental defect in INT1-2-deficient thymocytes. Notably, we also find directional joining consistent with RAG tracking upstream from the *Trdd2* RC 12RSS to the INT1-2 loop domain boundary in differentiating DN2/DN3 thymocytes, as revealed by junctions to \sim 70 different sets of cryptic RSSs and associated surrogate CEs across this domain. This upstream tracking abruptly ends at INT1-2. Correspondingly, INT1-2 deletion, which disrupts the upstream boundary of the INT2-to-TEA loop domain (Chen et al., 2015), allows apparent RAG tracking to continue upstream of the INT1-2

locale into the proximal *Trdv* domain. Thus, INT1-2 function in sequestering the *Trdd2* RC appears similar to that of IGCR1 in sequestering the *IgH* DJ_H RC (Guo et al., 2011; Hu et al., 2015), and suggests a potential contribution of RAG tracking to increased *Trdv3* utilization. Finally, RAG also potentially tracks downstream from *Trdd2* 23RSS to TEA, as revealed by directional junctions to 12 convergent cryptic RSSs in this 24-kb region.

Overall, our findings demonstrate that the INT2-to-TEA loop domain largely confines directional RAG joining activity from the initial *Trdd2* RC. Correspondingly, major RAG off-target activity is also confined within this domain, helping to suppress RAG activity during *Tcrd*V(D)J recombination at the huge number of potential off-target sites genome-wide. We do find low-level RAG-initiated *Tcrd* DSB translocations to DSBs in other TCR loci, consistent with the latter loci undergoing higher levels of DSBs, because they are also RAG targets, than other genomic loci. We also found that the level of such translocations was substantially increased in the absence of ATM (unpublished data), consistent with our prior finding of increased RAG-initiated translocations to Ig loci in ATM-deficient pro-B cells (Hu et al., 2015). Thus, our current findings support the proposal that, in contrast to other types of DSBs, RAG-generated DSBs to two target sites within a chromosomal antigen receptor locus loop are fused in association with the initiating RAG post-cleavage complex by being directly channeled into nonhomologous DNA end-joining (Bredemeyer et al., 2006; Deriano et al., 2011; Hu et al., 2015).

MATERIALS AND METHODS

Mice

The INT1-2KO mice have been previously described (Chen et al., 2015). Mouse work was performed under protocols approved by the Boston Children's Hospital (Boston, MA) and the Duke University (Durham, NC) Institutional Animal Care and Use Committees.

OP9-DL1 co-culture

The generation of DN2/DN3 T lymphocytes from adult bone marrow cells have been previously described (Huang et al., 2005; Holmes and Zúñiga-Pflücker, 2009). In brief, OP9-DL1 stromal cell lines were maintained in α -MEM (12571-071; Invitrogen), supplemented with 20% FBS (10438-026; Invitrogen). Flt3-L (427FL; R&D Systems) and IL-7 (407-ML; R&D Systems) were used at a concentration of 5 ng/ml each during co-culturing. After 14 d of differentiation, stromal cells and B cells were excluded by MACS-negative selection using CD140a (130-101-547; Miltenyi Biotec) and B220 (130-049-501; Miltenyi Biotec) MicroBeads, respectively. Genomic DNA was collected for LAM-HTGTS library preparation.

LAM-HTGTS and junction mapping

Primers used to generate libraries of *Trdd2* and *Trdd1* RAG-initiated bait-ends are listed in Table S6. No restriction enzyme blocking was performed for *Trdd2* and *Trdd1* librar-

ies. All LAM-HTGTS libraries were sequenced by Illumina Miseq and have been described previously (Hu et al., 2016). Junctions were mapped genome-wide using custom circos plots (Krzywinski et al., 2009; Frock et al., 2015) and mapped locally using IGV plots (Robinson et al., 2011; Hu et al., 2015).

Data analysis and normalization

Preprocessing of Miseq reads has been described previously (Frock et al., 2015). Processed reads were aligned using Bowtie2 (Langmead and Salzberg, 2012) to a modified-mm9 genome, in which *Tcra-Tcrd* locus at positions chr14: 52971974–54848751 was replaced with a 1666365nt-long segment from the 129S1/SvImJ strain (NT_039614.1). *Tcra-Tcrd* locus annotation was modified accordingly. For libraries from RAG-initiated bait-ends, we included duplicate junctions for analysis (Hu et al., 2015). We compared libraries, including duplicates, with those containing only unique junctions and observed similar patterns between them. We isolated the 12RSS-SE and 23RSS-CE libraries from 5-Primer libraries, as well as 23RSS-SE and 12RSS-CE libraries from 3-Primer libraries according to their coordinated bait sequence length (see the following paragraph). RAG-initiated bait-end libraries prepared from WT and INT1–2-deficient cells were normalized to the isolated junction numbers for comparison.

Isolating SE and CE bait junctions from the library generated by one common primer

We used specific criteria to separate SE and CE bait junctions from the same library. From the predicted SE and CE, we included an additional nucleotide beyond the predicted break site with respect to the primer used, to take into account the small fraction of junctions that coincidentally align beyond the predicted position as a result of nucleotide addition activities. We included several nucleotides for each predicted SE and CE, which represents the end processing of the bait BE sequences. Therefore, junctions with 71–76-bp bait sequence lengths were isolated from 5′-*Trdd2* LAM-HTGTS libraries as 12RSS-SE bait-end libraries, and 84–92-bp bait sequence lengths were similarly isolated for 23RSS-CE bait-end libraries, and for 3′-*Trdd2* LAM-HTGTS libraries, 61–66-bp bait sequence lengths were separated for 23RSS-SE and 75–81-bp for 12RSS-CE. 5′-*Trdd1* LAM-HTGTS libraries were similarly isolated, with 91–103-bp and 106–112-bp representing 12RSS-SE and 23RSS-CE bait sequence lengths, respectively, and for 3′-*Trdd1* LAM-HTGTS libraries, 91–98- and 101–107-bp bait sequence lengths for 23RSS-SE and 12RSS-CE, respectively were isolated. Junctions with 46–52-bp bait sequence lengths were isolated from 5′-*Trdj1* libraries for *Trdj1*-12RSS-SE and 45–52-bp bait sequence lengths were isolated from 3′-*Trdj1* libraries for *Trdj1*-12RSS-CE.

Adjusting the sequential joins containing *Trdd2* sequence

For *Trdd1*-23RSS-CE and *Trdj1*-12RSS-CE libraries, insertion sequences of junctions were screened for *Trdd2* consen-

sus sequence by using a sliding window approach with a 1-bp step size and a window size of 5 bp across *Trdd2* (12 iterations), and such junctions were manually adjusted back to *Trdd2*.

Junction hotspots and RAG on- and off-target identification

Junction-enriched regions were identified by MACS2 (Zhang et al., 2008) with custom parameters (extsize, 20 bp; FDR cut-off, 10^{-9}). Enriched regions identified in at least three individual libraries were considered as recurrent hotspots and used in the following study. The bona fide RSS sequence information flanking *Tcr* gene segments was collected from the IMGT/GENE-DB database (Giudicelli et al., 2005). Hotspots that overlapped with bona fide RSS were recognized as RAG on-targets. Hotspots positioned <10 bp away from the simple CAC motif were defined as associated with RAG off-targets.

ChIP-Seq

We obtained CTCF ChIP-seq data from DN thymocytes from (Shih et al., 2012; available from GEO under accession no. GSM1023416), RAG1, RAG2, and H3K4me3 ChIP-seq data from (Teng et al., 2015; available from GEO under accession nos. GSM1701786, GSM1701790, and GSM30317). Data were reanalyzed with our mm9-modified genome using the ChIP-seq data analysis pipeline Chilin.

Accession code

The GEO accession no. for the datasets reported in this paper is GSE79892.

Online supplemental material

Fig. S1 shows the high-level enrichment of RAG binding and H3K4me3 at *Trdd2*. Fig. S2 displays genome-wide distribution of prey junctions from *Trdd2* 12RSS and 23RSS CE libraries. Fig. S3 shows bait length and genome-wide prey junction distributions baited from RAG-initiated *Trdd1* DSBs at *Trdd1*. Fig. S4 includes examples of precise signal joins and processed surrogate coding joins. Fig. S5 shows junctions not corresponding to bona fide RSS sites are associated with RAG off-targets. Fig. S6 displays the profiles of off-targets baited from RAG-initiated DSBs at *Trdd1*. Table S1 is the summary of all LAM-HTGTS libraries from various bait-ends. Table S2 is the summary of total translocations to RAG on- or off-targets identified by RAG-initiated bait-ends flanking *Trdd2*, *Trdd1*, and *Trdj1*. Table S3 lists the percentage of relative junction distributions of all LAM-HTGTS libraries from RAG-initiated bait-ends flanking *Trdd2*, *Trdd1* and *Trdj1*. Table S4 lists *Tcra-Tcrd* gene segments and joining percentages from RAG-initiated bait-ends flanking *Trdd2*, *Trdd1*, and *Trdj1* in T cell precursors. Table S5 lists of δ RECs and RAG off-target sites identified at *Tcra-Tcrd* via LAM-HTGTS of WT T cell precursors. Table S6 lists LAM-HTGTS oligos used to clone *Trdd2*, *Trdd1*, and *Trdj1* bait-end junctions. Tables S1–S6 are available as Excel files. Online

supplemental material is available at <http://www.jem.org/cgi/content/full/jem.20160670/DC1>.

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Author contributions: L. Zhao, R.L. Frock, and F.W. Alt designed the study. L. Zhao performed all the experiments. L. Chen and M.S. Krangel provided the INT1-2-deficient mice. Z. Du helped with statistical analyses. J. Hu provided suggestions during manuscript preparation. L. Zhao, R.L. Frock, M.S. Krangel, and F.W. Alt interpreted the data, designed the figures, and wrote the manuscript.

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