An ectopic CTCF-dependent transcriptional insulator influences the choice of Vβ gene segments for VDJ recombination at TCRβ locus

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ABSTRACT
Insulators regulate transcription as they modulate the interactions between enhancers and promoters by organizing the chromatin into distinct domains. To gain better understanding of the nature of chromatin domains defined by insulators, we analyzed the ability of an insulator to interfere in VDJ recombination, a process that is critically dependent on long-range interactions between diverse types of cis-acting DNA elements. A well-established CTCF-dependent transcriptional insulator, H19 imprint control region (H19-ICR), was inserted in the mouse TCRβ locus by genetic manipulation. Analysis of the mutant mice demonstrated that the insulator retains its CTCF and position-dependent enhancer-blocking potential in this heterologous context in vivo. Remarkably, the inserted H19-ICR appears to have the ability to modulate cis-DNA interactions between recombination signal sequence elements of the TCRβ locus leading to a dramatically altered usage of Vβ segments for Vβ-to-DβJβ recombination in the mutant mice. This reveals a novel ability of CTCF to govern long range cis-DNA interactions other than enhancer-promoter interactions and suggests that CTCF-dependent insulators may play a diverse and complex role in genome organization beyond transcriptional control. Our functional analysis of mutated TCRβ locus supports the emerging role of CTCF in governing VDJ recombination.

INTRODUCTION
Insulators prevent interactions between promoters and enhancers in a position-dependent manner due to their ability to influence higher order chromatin structure (1). CTCF (CCCTC-binding factor) appears to be important for their activity as it can facilitate both intrachromosomal and interchromosomal interactions and thus define chromatin domains that may be independently regulated (2). A large number of CTCF binding sites have been identified in mammalian genomes suggesting their extensive involvement in governing cis-DNA interactions among regulatory elements (3–5). Whether CTCF defined domains restrict specifically the enhancer-promoter communication relevant for transcriptional regulation or can influence other types of cis-DNA interactions in the genome is not currently known.

In this context, antigen receptor loci like IgH, TCRξ/δ and TCRβ that code for the immunoglobulin (Ig) and T-cell receptors (TCR), are particularly interesting. By exhibiting long range interactions between different types of elements, they present a useful framework to explore the role of CTCF in defining independently regulated chromatin domains. Enhancer–promoter interactions are necessary for defining the developmental stage specificity of RAG-mediated VDJ recombination at these loci (6). Additionally, recombination requires physical interaction between recombination signal sequence (RSS) elements associated with the V, D and J segments in a combinatorial manner. These segments are located at large distances from each other on the chromosome and higher order chromatin reorganization, manifested as locus contraction, is necessary to bring them together prior to recombination in a lineage specific manner (7,8).

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Recent analysis of the IgH and Igk loci demonstrates that interactions between CTCF binding sites indeed organize the higher order chromatin structure relevant for regulation of transcription and recombination. Multiple CTCF binding sites at these loci were postulated to organize the chromatin in concert with cohesin, Pax5, YY1 etc. (9,10). As predicted, some of the sites are important for enhancer blocking at the IgH locus (11,12). An additional role of CTCF and cohesin in locus contraction seems evident by the multiple interactions between the distantly placed CTCF binding sites at the IgH locus in Pro-B cells (12,13). Knockdown of CTCF led to a partial reduction in these long range interactions thereby supporting the role of CTCF in higher order chromatin organization (14). A conditional knockdown of CTCF in ProB cells demonstrated that CTCF may not be necessary for VDJ recombination per se but plays an important role in restricting Igk enhancer activity and thus influences recombination at this locus (15). While these studies underscore the importance of CTCF for appropriate chromatin organization relevant to VDJ recombination, the manner by which CTCF may impact this process needs further elucidation (16).

To gain further insights into the nature of chromatin domains organized by CTCF and their control on nuclear processes like transcription and VDJ recombination, we adopted a 'gain-of-function' genetic approach. A CTCF binding element [H19 imprint control region (H19-ICR) of the Igf2/H19 locus] was inserted to the murine TCRβ locus by genetic manipulation. We present evidence to show that introduction of ectopic CTCF binding sites from an unrelated locus is sufficient to drastically alter the transcription and recombination patterns at the TCRβ locus.

Organization and regulation of murine TCRβ locus, encoding β chain of α/β T cell receptor (α/β TCR), is useful to delineate the interactions of diverse elements. Unlike the relatively large IgH and Igk loci, TCRβ locus spans only about 700 kb in the genome. It has 31 Vβ gene segments (TRBV1 to TRBV31). Majority of the Vβ segments are located upstream to the two clusters of Dβ and Jβ segments, i.e. Dβ1-Jβ1.1-1.7-Cβ1 (DJCβ1) and Dβ2-Jβ2.1-2.7-Cβ2 (DJCβ2) while the TRBV31 is located downstream to the DJCβ clusters (Figure 1A) (17). A developmentally regulated enhancer (Eβ) and promoters PDβ1 and PDβ2 are the key regulatory elements that interact functionally during early double negative (DN, CD8- CD4-) stage of T cell development (18–21). Their interaction, achieved by formation of a holocomplex comprising enhancer Eβ and promoter PDβ1 and/or PDβ2, leads to generation of chromatin accessibility restricted to about 25 kb region encompassing the DJCβ1 and DJCβ2 clusters (22,23). Consequently, PDβ1 and PDβ2 initiate germline transcripts from DJCβ1 and DJCβ2 clusters, respectively. The altered chromatin also becomes accessible to RAG proteins that orchestrate the RSS-mediated Dβ-to-Jβ recombination at the DJCβ clusters in accordance with the accessibility hypothesis (24). The alleles that have undergone successful Dβ-to-Jβ recombination, act as substrates for RSS-mediated Vβ-to-DβJβ recombination that generates the functional TCRβ gene. Chromatin structure of Vβ segments is also altered in DN T cells but is independent of regulation by Eβ (25). Juxtapositioning of RSS elements associated with Vβ and DJββ segments is achieved by locus contraction prior to recombination (8). Keeping these observations in mind, we argued that the insertion of a CTCF-dependent insulator at the TCRβ locus could potentially interfere with different aspects of VDJ recombination that may be regulated by Eβ and/or be otherwise influenced by chromatin organization.

We exploited the CTCF-binding activity of the H19-ICR of the murine Igf2/H19 locus (Figure 1B) for interfering with TCRβ locus regulation. H19-ICR regulates the parent-of-origin specific monoallelic expression of H19 and Igf2 in vivo (26–28). It binds CTCF and organizes an insulator that prevents Igf2 promoter activation by the downstream enhancers on the maternally inherited allele. Loss of CTCF binding due to methylation of CpG residues prevents binding of CTCF and abrogates enhancer blocking on the paternally inherited allele leading to activation of Igf2 promoter by the enhancers (29,30). Loss of CTCF binding due to targeted mutations of CTCF binding sites on the maternal allele activates maternal Igf2 gene emphasizing the crucial link between enhancer blocking and CTCF (31). We chose H19-ICR as the insulator to perturb the TCRβ locus because the enhancer blocking by H19-ICR is clearly dependent on CTCF. Also, when inserted at the Afp locus in mice, H19-ICR was able to interact with enhancer and

**Figure 1.** Schematic diagrams showing organization of the genetic loci used for analysis. (A) Murine TCRβ loci: wild-type TCRβ locus, mutant TCR-ins locus showing insertion of H19-ICR, mutant TCR-mut locus showing insertion of H19-ICR-mut with all four CTCF binding sites lost and mutant TCRβ-del locus showing replacement of region encompassing Jβ1.3 to Cβ2 with neomyrin resistance gene (Neo-r). Grey ovals represent regulatory regions Eβ, PDβ1 and PDβ2 (including 5’PDβ2 and 3’PDβ2 with respect to DJCβ). Each DJCβ cluster has a single Dβ segment (dark vertical line), seven Jβ segments (light vertical lines) and a Cβ region (grey rectangle). The 31 Vβ segments (TRBV1 to TRBV31) are represented as vertical lines (V1 to V31). RSS elements (not shown) are located downstream to each TRBV segment, upstream and downstream to each Dβ segment and upstream to each Jβ segment. (B) Murine Igf2/H19 locus showing the endogenous location of H19-ICR relative to Igf2 and H19 genes and their shared endodermal enhancers. The TCRβ and Igf2/H19 loci span about 700 and 100 kb, respectively, in the mouse genome.
promoter as it does at the endogenous Igf2/H19 locus (32). Although the organization of the regulatory elements at the Afp locus precluded an unambiguous functional readout of enhancer blocking, these observations suggested that the H19-ICR was likely to function as an enhancer blocker in a heterologous context in vivo.

Our analysis demonstrates that the inserted H19-ICR is capable of curtailing functional enhancer–promoter interaction required not only for transcriptional activation but also for DJβ-to-Jβ recombination in the heterologous context of TCRβ.

Most intriguingly, we find that the usage of Vβ gene segments in the TCRβ chains of the mutant mice is altered. This indicates that the inserted H19-ICR can perturb not only enhancer–promoter interactions but can also reorganize the chromatin to influence Vβ-to-DJβ recombination. These results demonstrate the ability of a CTCF-dependent transcriptional insulator to influence long range DNA interactions necessary for VDJ recombination and provide useful insights into the multiple roles of CTCF in light of the topological model of interactions of regulatory elements in the genome.

MATERIALS AND METHODS

All experiments using mice were conducted as approved by Institutional Animal Ethics Committee. Sequences of all primers are given in Supplementary Table S1.

Targeting the H19-ICR and H19-ICR-mut to the TCRβ locus

To generate the TCR-ins (Figure 1A), a targeting vector was constructed (Supplementary Figure S1), that had a 5.7 kb BamHI–HindIII fragment and a 4.7 kb HindIII–BamHI fragment from the TCRβ locus as the 5′ flank and 3′ flank, respectively, which originated from 129S6/SvEvTac gDNA. The 2.4 kb BglII fragment carrying the H19-ICR fragment with all four CTCF binding sites and a ‘neomycin resistance gene’ cassette (Neo-r) were cloned between the flanks along with three Lox P sites in tandem orientation. Additionally, ‘diphtheria toxin-A’ (DTA) gene was also cloned that served as the negative selection marker. The construct was linearized with NotI and electroporated in R1 embryonic stem (ES) cells derived from mouse strain 129X1/SvJ x 129S1 and the clones were selected for G418 resistance. Genomic DNA from G418 resistant clones was digested with BglII and subjected to Southern hybridizations with the 5′ and 3′ probes. The correctly targeted alleles (Tins) generate 6.4 and 7.5 kb bands upon hybridization with the 5′ and 3′ probes, respectively, compared to the 12.6 kb band of the wild-type allele in each case. The Neo-r cassette was subsequently removed by electroporation with a plasmid pBS185 transiently expressing ‘Cre recombinase’. The correct clones (TCR-ins), which had excised Neo-r gene but retained the H19-ICR, were screened using a PCR-based strategy that detected the insulator-TCRβ junctions on either end. Further, the TCR-ins clones having excision of Neo-r gene were confirmed by Southern hybridization of BglII digested gDNA. The BglII fragment size, detected by 3′ probe, in the non-excised Tins clones is 7.5 kb and changes to 6.4 kb in Neo-r excised TCR-ins clones. Correctly manipulated TCR-ins ES cells were micro-injected into C57bl/6 blastocysts to derive founder chimeric mice that were bred to derive TCR-ins mutant mice. The germine transmission of the mutation in the mice was confirmed by Southern hybridization and PCR analysis (Supplementary Figure S1).

Exactly similar strategy was used for targeting of H19-ICR-mut to generate TCR-mut mice. The H19-ICR-mut had all the four CTCF binding sites mutated (31).

Generation of the mouse strain TCR-cas

The congenic mouse strain TCR-cas was generated that carried wild-type TCRβ locus derived from Mus castaneus castaneus in a Mus musculus domesticus genetic background. The female F1 progeny of Mus castaneus castaneus × Mus musculus domesticus was backcrossed to Mus musculus domesticus males for eight generations to derive this strain. At each generation, the presence of castaneus sequences upstream and downstream to TCRβ allele was verified by PCR using oligonucleotide primers DMit274 and D6Nds4.

Flow cytometric analysis

Thymocytes from TCR-ins mutant mice and littermate controls were immunostained with anti-Cd4-PE (clone RM4-5) and anti-Cd8a-Cy5 (clone 53-6.7) and analyzed on BD-LSR Flow Cytometer. For the Vβ choice analysis, these were additionally stained with anti TRBV31-FITC (clone 14-2) or a cocktail of FITC conjugated anti-Vβ antibodies that included anti-TRBV2 (clone KT4), anti-TRBV4 (clone B21.5), anti-TRBV12.1/12.2 (clone MR9-4), anti-TRBV13.1/13.2 (clone MR5-2), anti-TRBV13.3 (clone 1B3.3) and anti-TRBV19 (clone RR4-7). The anti-Vβ antibodies were from the Mouse Vβ TCR screening panel (BD Bioscience cat no. 557004) but Vβ representation is according to IMGT nomenclature. Total thymocytes immunostained with anti-Cd4-PE (clone RM4-5), anti-Cd8a-Cy5 (clone 53-6.7) and anti-Cd25-FITC (clone 7D4) were used to sort DN2/3 thymocytes (CD4+ CD8− CD25+) on BD FACS ARIA III. The purity of the sorted population was more than 95%. All antibodies were from BD Biosciences, USA.

Allele specific transcriptional analysis by SNAPSHOT assay

Allele specific expression of germline transcripts was analyzed in thymocyte RNA of 4–5 weeks old RAG deficient mice which had maternally inherited domesticus allele (wild-type or TCR-ins) and paternally inherited castaneus allele (TCR-cas). Random primers were used for the reverse transcription and gene-specific primers were used to amplify DJβ1, DJβ2, TRBV13.2 and TRBV31. The amplified products were subjected to SNAPSHOT analysis (Snapshot multiplex kit, ABI) that relied on incorporation of fluorescently labelled diodeoxynucleotides by single nucleotide primer extension (SNuPE) based on specific nucleotide differences between
incorporation of dideoxynucleotides by F1 gDNA was domesticus doemsticus before calculating allele specific contribution of the was reverse transcribed using random primers and used Thymocytes from 6–8 weeks old TCR-ins, TCR-mut and Transcriptional analysis by quantitative RT-PCR before calculating allele specific contribution of the doemsticus (TCR-ins or wild-type) and castaneus alleles to total mRNA.

Transcriptional analysis by quantitative RT-PCR
Thymocytes from 6–8 weeks old TCR-ins, TCR-mut and control mice were used to derive total RNA. The RNA was reverse transcribed using random primers and used for quantitative PCR to detect transcripts for DJβ1, DJβ2, Thy1.2, TRBV13-Dβ-Jβ1.1, TRBV31-Dβ-Jβ1.1, TRBV13-Dβ-Jβ2.7 and TRBV31-Dβ-Jβ2.7 using gene specific primers. The forward TRBV13 primer was designed to detect TRBV13.1, TRBV13.2 and TRBV13.3. SYBR-Green chemistry based Real time PCR analysis on ABI PRISM-7000 Sequence Detection System, ABI, USA was used for the quantitative PCR. The amount of transcripts in mutant mice were calculated normalized to the endogenous reference (Thy1.2) and relative to a calibrator (control mice) using the standard curve method for relative quantification.

Dβ-to-Jβ recombination analysis by PCR
Genomic DNA isolated from thymocytes was subjected to PCR and analyzed for the appearance or disappearance of recombination specific bands as described.

DJCβ usage analysis
RNA from thymocytes was reverse transcribed using random primers followed by PCR amplification using gene specific primers to detect rearranged TCRβ genes having TRBV12 (including TRBV12.1 and TRBV12.2), TRBV13 (including TRBV13.1, TRBV13.2 and TRBV13.3) and TRBV31 and any of the two Cβ (Cβ1 or Cβ2). The amplified products were cloned. Sequence analysis from at least 36 clones of each genotype was carried out and fraction of clones using DJCβ1 or DJCβ2 determined.

RESULTS
Genetic manipulation of the TCRβ locus
We used a ‘knock-in’ approach to insert the H19-ICR at the TCRβ locus in mouse ES cells such that the 2.4 kb H19-ICR element, carrying all four CTCF binding sites and known to be sufficient for enhancer blocking in vitro (30,33), was introduced immediately downstream to the PDβ1-regulated transcription unit comprising Dβ1-Jβ1.1-1.7-Cβ1 (DJCβ1; Figure 1A). This positioned the H19-ICR between the promoter PDβ1 and the enhancer Eβ. PDβ2 dual promoters, responsible for the expression of the second transcription unit Dβ2-Jβ2.1-2.7-Cβ2 (DJCβ2), are located downstream to the H19-ICR insertion site. Correctly targeted ES cells (TCR-ins) were used to derive mutant mice that carried the H19-ICR inserted at the TCRβ locus. To unambiguously ascertain the importance of CTCF in the H19-ICR function in the heterologous context, we also generated mutant mice (TCR-mut) that carried a mutant version of H19-ICR (H19-ICR mut) wherein all the four CTCF binding sites were mutated (Figure 1A). The four site-specific mutations are known to abrogate CTCF binding by H19-ICR in vitro as well as in vivo (31). All heterozygous and homozygous mutants were viable, fertile and indistinguishable from the wild-type mice in physical appearance.

H19-ICR functions as an enhancer blocker specifically upon maternal inheritance at the endogenous Igf2/H19 locus. Hence, the transcription and recombination analysis was carried out in an allele specific manner. Thymocytes for the analysis were derived from mutant mice that maternally inherited TCR-ins (M, TCR-ins/TCR-cas) or TCR-mut (Mx, TCR-mut/TCRβ-del) or wild-type TCRβ allele (C, +/TCRβ-del). In each case, the presence of TCRβ-del (Figure 1A), carrying 15 kb deletion (encompassing DJCβ1 and DJCβ2) as the paternal allele, ensured that it would not contribute to any analysis as it could neither generate transcripts in the deleted region nor undergo recombination (34,35). Alternatively, using a congenic mouse strain, TCR-cas (see ‘Materials and Methods’ section for details), mutant mice were obtained that maternally inherited domesticus TCR-ins (M*, TCR-ins/TCR-cas) or domesticus wild-type TCRβ allele (C*, +/TCR-cas). In each case, the paternally inherited allele was castaneus wild-type TCRβ (TCR-cas). Nucleotide differences between the domesticus and castaneus alleles afforded allele-specific transcriptional analysis of thymocytes in these mice.

H19-ICR insertion at the TCRβ locus does not arrest overall thymocyte development
Thymocyte development proceeds sequentially via double negative (DN, CD4–CD8–), double positive (DP, CD4+CD8+ and single positive (SP, CD4+CD8– and CD4–CD8+) stages characterized by the presence of cell surface markers CD4 and CD8. The development beyond DN stage depends on the synthesis of a functional TCRβ chain consequent to successful VDJ recombination of TCRβ locus (35). Since the inserted H19-ICR could potentially interfere in the regulation of transcription and/or recombination at TCRβ locus, we first examined the effect of our TCRβ manipulation on overall thymocyte development.

The cellularity of the thymus in TCR-ins (M), TCR-mut (Mx) and control (C) mice as well as their thymocyte developmental profiles were comparable (Figure 2). As described above, the TCRβ-del background allele is not capable of supporting recombination and contributing to T-cell development beyond DN stage (35). Therefore, T-cell development beyond DN stage to generate DP and eventually SP thymocytes in the mutant mice demonstrated the ability of the TCR-ins and TCR-mut
alleles to undergo VDJ recombination and generate functional TCRb chains. Thus, the inserted H19-ICR neither completely abolished VDJ recombination nor arrested thymocyte development. The ability of TCR-ins and TCR-mut mutant alleles to support VDJ recombination cannot be used to simplistically infer that regulation of VDJ recombination was not affected. Multiple Vb, Db, and Jb segments are used in a combinatorial manner during VDJ recombination. Hence, it was plausible that the inserted position-dependent H19-ICR insulator influenced transcription and recombination in a gene-segment-specific manner. Thymocytes isolated from the mutant mice were, therefore, utilized to unambiguously investigate gene-segment-specific transcription as well as Db-to-Jb and Vb-to-DbJb recombination events at the manipulated alleles.

**Effective CTCF-dependent enhancer blocking by inserted H19-ICR**

We hypothesized that if the H19-ICR organized a functional insulator between promoter PDβ1 and enhancer Eβ in the TCR-ins allele, it would reduce the generation of DJβ1 germline transcripts that depend on activation of PDβ1 by Eβ. An assay was designed wherein RT-PCR was followed by SNAPSHOT analysis. Based on this, single nucleotide differences between the *domesticus* and *castaneus* alleles were utilized to determine the contribution of the mutant (TCR-ins) or the wild-type (+) allele to the germline transcripts originating from various promoters of the TCRβ locus in thymocytes of TCR-ins mutants (M*) and control mice (C*). The analysis was performed in RAG deficient mice (Rag−/−) to prevent VDJ recombination and avoid heterogeneity in DNA templates that could interfere in accurate detection of germline transcripts.

A drastic reduction in the DJβ1 transcription from the mutant TCR-ins allele was evident as it contributed to less than 5% of the total DJβ1 transcripts in the mutant mice (M*; Figure 3A). The control mice (C*) exhibited nearly equal contribution of the *domesticus* and *castaneus* alleles (biallelic expression) for DJβ1 as expected. We also observed that the expression of DJβ2 transcripts continued to be biallelic in mutants as in the control mice (Figure 3A). Further, the Vβ germline transcripts (TRBV13.2 and TRBV31) also continued to be expressed from both alleles in mutants as in control mice (Figure 3A).

**Figure 2.** Effect of H19-ICR insertion on T-cell development analyzed in thymocytes isolated from mutant mice. (A) Mutants that maternally inherited the TCR-ins (TCR-ins/TCRβ-del) or wild-type TCRβ allele (+/TCRβ-del). (B) Mutants that maternally inherited TCR-mut (TCR-mut/TCRβ-del) or wild-type TCRβ allele (+/TCRβ-del). Thymocytes were immunostained to detect the presence of cell surface markers CD4 and CD8.
The significant impairment in DJβ1 transcripts indicated that the inserted *H19-ICR* abrogates Eβ and PDβ1 interaction necessary for transcriptional activation. Insulators curtail enhancer-based promoter activation only when located between the enhancer and the promoter. Our finding of abrogation of Eβ-PDβ1 interaction taken together with no alteration in Eβ-PDβ2 interaction unambiguously demonstrated that the inserted *H19-ICR* acts as an effective position-dependent enhancer blocking insulator when inserted at the TCRβ locus. The Vβ promoters are known not to be affected by Eβ (25). An unaltered transcriptional profile of Vβ germline transcripts indicated that *H19-ICR* insertion specifically abrogated the Eβ-PDβ1 interaction rather than lead to generalized alterations in germline transcription at the TCRβ locus.

It was important to confirm the abrogation of Eβ-PDβ1 transcription in RAG sufficient (Rag<sup>+/+</sup>) mice since normal RAG expression in the developing T cells was necessary for the analysis of interference of inserted *H19-ICR* with VDJ recombination. To minimize the influence of recombination on detection of germline transcription, DN thymocytes at DN2/3 stage of development were isolated. Such cells have a high proportion of alleles that have not undergone recombination and hence are in the germline configuration. A quantitative RT-PCR assay was designed to specifically estimate the abundance of DJβ1 germline transcripts arising due to Eβ-PDβ1 interaction prior to recombination (Figure 3B). Further, the RT-PCR primer design ensured that the DJβ1 transcripts arising from the paternal TCRβ-del allele were not detected. In this assay also, the DJβ1 transcripts in
mutant mice (M) were observed to be drastically reduced compared to controls (C; Figure 3B, left graph). Further, the Eβ–PDβ2 interaction-dependent DJβ2 germline transcripts continued to be expressed at high levels in TCR-ins mutants (M) like the control mice (C) in the sorted DN2/3 thymocytes (Supplementary Figure S2). It was evident, therefore, that the inserted H19-ICR functions as a position-dependent enhancer blocker in RAG sufficient mice also. All subsequent experiments were performed on mice with RAG sufficient background.

Finally, to ascertain the requirement of CTCF for the insulator organized by H19-ICR, we analyzed the abundance of DJβ1 germline transcripts in TCR-mut mice (Mx). Clearly, in these mice, the Eβ–PDβ1 interaction-dependent DJβ1 transcription was not affected (Figure 3B, right graph). This demonstrated that the enhancer blocking organized by the inserted H19-ICR at the TCRβ locus is CTCF dependent.

Severe abrogation of Eβ–PDβ1 interaction-mediated recombination

Deletion analysis has indicated that the interaction of Eβ with the promoters PDβ1 and PDβ2 leads to generation of chromatin accessibility both for transcription and recombination (22). Since the inserted H19-ICR was able to prevent Eβ–PDβ1 interaction-mediated transcriptional activation, we examined its ability to interfere in DJβ-to-Jββ recombination by PCR based on genomic DNA from thymocytes (Figure 4A).

Amplification products of 450 and 318 bp arising due to the recombination of DJβ1 with Jββ1.1 and Jββ1.2, respectively, were evident in control mice (C) but were absent in TCR-ins mutants (M). Further, the 1.1 kb band representing the non-rearranged germline configuration was robustly amplified in mutants (Figure 4A, left panels). This indicated a severe abrogation of recombination at the DJCβ1 cluster. The block in recombination was CTCF dependent as TCR-mut mice (Mx) did not exhibit such an abrogation. In accordance with our observation that H19-ICR did not influence Eβ–PDβ2 interaction for transcription, the PDβ2-regulated recombination of DJβ2 with Jβ2 was also not significantly altered in TCR-ins mutants. Both mutants (M) and control littermates (C) exhibited the recombination of DJβ2 with Jβ2.1 and Jβ2.2, evidenced by appearance of 411 and 208 bp band and reduction in amplification of 991 bp band (Figure 4A, right panels). These results suggest that enhancer–promoter interaction-dependent recombination can also be curtailed by an insulator in a position-dependent manner.

**Figure 4.** Effect of H19-ICR insertion on DJβ-to-Jββ recombination and consequent choice of DJCβ1 cluster for Vβ-to-DJββ recombination. (A) Recombination at DJCβ1 and DJCβ2 gene clusters. Schematic diagram shows the regulatory regions Eβ, PDβ1 and PDβ2 (grey ovals) and PCR primers (grey and black triangles) that detect DJβ-to-Jββ recombination. Recombination of DJβ1 to Jββ1.1 and Jββ1.2 (left panels) at DJCβ1 and DJβ2 to Jββ2.1 and Jββ2.2 (right panels) at DJCβ2 was analyzed in thymocyte gDNA of TCR-ins mutants (M), TCR-mut mutants (Mx) and control mice (C). Genomic DNA derived from kidney (kid) and thymocytes (Thy) of wild-type mice (+/+; Mx  (TCR-mut / TCR-ins)) and TCR-mut (Mx, TCR-mut/TCR-ins) was used for RT-PCR using forward primers specific to TRBV12 or TRBV13 or TRBV31 (black arrows on the locus diagram) and a reverse primer common to exon 1 of both Cβ1 and Cβ2 (grey arrows on the locus diagram). The amplification products were cloned and sequenced to determine the fraction of clones with DJCβ1 or DJCβ2 as depicted in the graph. At least 36 clones were sequenced for each genotype. For PCR and RT-PCR analysis, thymocytes were isolated from mice that maternally inherited the wild-type TCRβ allele (C, +/TCRβ-del) or TCR-ins (M, TCR-ins/TCRβ-del) or TCR-mut (Mx, TCR-mut/TCRβ-del).
Preferential usage of DJCβ2 cluster for VDJ recombination

The TCRβ alleles that have successfully recombined DJβ segments show a preference for usage of DJCβ regions as a substrate for RSS-mediated recombination between Vβ and DJβ segments. The severe reduction in recombinating at DJCβ1 clusters upon maternal inheritance of TCR-ins allele (Figure 4A), was therefore expected to substantially reduce the usage of DJCβ1 region in TCRβ chains in these mutants. To verify this, RNA was isolated from thymocytes of TCR-ins mutant and control mice. TRBV12 and TRBV13 together contribute to TCRβ chains on about 30% of thymocytes (36). Primers for RT-PCR were designed such that the forward primer could bind to specific Vβ regions (TRBV12, TRBV13 or TRBV31) while the reverse primer was common to exon I of Cβ1 and Cβ2 and avoided any inherent bias for amplification of Cβ1 or Cβ2 (Figure 4B). Amplified products after RT-PCR were cloned and sequenced to estimate the usage of DJCβ1 and DJCβ2 in transcripts encoding TCRβ chain. Sequence analysis revealed a very strong bias in favor of DJCβ2 usage in mice that had maternally inherited TCR-ins (M) and exhibited severely compromised DJβ1-to-Jβ1 recombination compared to the mice that had a wild-type allele (C; Figure 4B). Again, CTCF dependence was clearly evident as the DJCβ2 usage in TCR-mut mice (Mx) was comparable to control mice.

The enhanced usage of DJCβ2 cluster for Vβ-to-DJββ recombination was a logical consequence of efficient enhancer blocking by the inserted H19-ICR that curtailed Eβ-PDβ1 interaction based events, i.e. DJβ1 transcription and DJβ1-to-Jβ1 recombination and hence reduced the usage of DJCβ1 for Vβ-to-DJββ recombination.

Altered usage of Vβ segments for VDJ recombination

VDJ recombination at TCRβ locus involves long range cis-DNA interactions of RSS elements between the recombined DJβ segments and the Vβ segments located a large distance away. In the TCR-ins allele, in addition to the two DJCβ clusters, Vβ regions also get partitioned by the insulator such that TRBV13-TRBV30 are located upstream to the DJCβ1 cluster and inserted H19-ICR while TRBV31 is located downstream to the H19-ICR and DJCβ2 cluster (Figure 5A). Having established that the inserted H19-ICR organizes a CTCF-dependent functional insulator at TCRβ locus, we analyzed the choice of Vβ gene segments used for Vβ-to-DJββ recombination in TCR-ins mutant mice.

Thymocytes were analyzed for surface expression of TCRβ that use Vβ gene segments present on either side of the H19-ICR. We first analyzed the TRBV31 specific TCRβ by immunostaining thymocytes from TCR-ins mice that organize a functional insulator. We observed an enormous increase (~8- to 10-fold), in the number of CD4-SP thymocytes (CD4+CD8−) that had TRBV31 in the TCRβ chain in TCR-ins mutants (M; Figure 5B) compared to control littermates (C). Also, in the same thymocyte samples, there was a drastic reduction (about 5-fold) in the usage of several other Vβ tested (TRBV2, TRBV4, TRBV12, TRBV13 and TRBV19). A similarly altered Vβ usage was observed in CD8-SP cells (data not shown). Importantly, such a drastic alteration was not observed for TCR-mut mice (Mx; Figure 5B) that are unable to organize an insulator due to mutated CTCF binding sites.

This suggested that the presence of the ectopic H19-ICR insulator is responsible for a decrease in usage of Vβ segments located upstream to the inserted H19-ICR and a concomitant increase in the usage of TRBV31 located downstream to the inserted H19-ICR. Thus, Vβ choice for Vβ-to-DJββ recombination was influenced by H19-ICR in a CTCF-dependent manner. In conjunction with the increased DJCβ2 usage (Figure 4B), these results suggest a strong bias in favor of TRBV31 recombination specifically with DJCβ1 cluster in the TCR-ins allele. In sharp contrast, recombination of all other Vβ segments with DJCβ2 cluster, i.e. interactions across H19-ICR, appear to be hindered significantly.

To confirm our finding, we directly tested the choice of recombination of DJCβ1 and DJCβ2 clusters with TRBV13 and TRBV31. An RT-PCR assay was designed such that the forward primers, in independent PCR reactions, recognize TRBV13 or TRBV31 while the reverse primer recognizes Jβ1.1 in both these reactions. Jβ1.1 was taken as a representative Jβ for the use of DJCβ1 cluster as we observed it to be used maximally in the sequence based analysis of Vβ-to-DJββ recombination described earlier. In the mice that inherited the TCR-ins maternally (M), there was a drastic reduction in TRBV13-DJβ-Jβ1.1 as well as TRBV31-DJβ-Jβ1.1 transcripts (Figure 6A) consistent with the observation of a reduced ability of DJCβ1 to support Vβ-to-DJββ recombination (Figure 4A and B).

In a similar assay, by designing the reverse primer in Jβ2.7, we estimated the choice of DJCβ2 for recombination with TRBV13 and TRBV31. In TCR-ins mutants (M), the TRBV31-DJβ-Jβ2.7 transcripts were increased by more than 10-fold while the TRBV13-DJβ-Jβ2.7 transcripts were about 50% less abundant compared to control mice (C; Figure 6B). Thus, effectively, the usage of TRBV31 increased more than 20-fold relative to usage of TRBV13 for recombination to DJCβ2 cluster in the presence of a functional insulator (Figure 6C, upper graph). Unlike TCR-ins mutants (M), TCR-mut mice (Mx) did not exhibit any enhancement in the TRBV31-DJβ-Jβ2.7 to TRBV13-DJβ-Jβ2.7 transcript ratio compared to control mice (C; Figure 6C, lower graph).

Our results indicate that the H19-ICR significantly alters the regulation of transcription as well as recombination at the TCRβ locus and this effect is not merely due to the presence of an inserted 2.2 kb DNA fragment. Ability of inserted H19-ICR to bind CTCF was essential for its influence on these processes. Just as it prevents the enhancer–promoter communication when present between them, the H19-ICR insulator also prevented RSS elements from interacting with each other in a CTCF-and-position-dependent manner, i.e. when located between them and led to an alteration in the choice of Vβ segments for recombination (Figure 6D).
DISCUSSION

In this investigation, we took advantage of the precisely regulated transcription and recombination at the TCRβ locus to understand the ability of a CTCF-dependent insulator to interfere in the interactions between several cis-acting regulatory elements. By organizing an ectopic insulator at TCRβ locus, we demonstrate a novel ability of CTCF to modulate cis-DNA interactions between RSS elements during VDJ recombination.

Our analysis of the TCR-ins and TCR-mut alleles indicated that H19-ICR acts as an efficient CTCF-dependent enhancer blocker when inserted to the TCRβ locus and prevents Eβ-regulated transcription and recombination. H19-ICR interacts with several CTFC and cohesin binding sites at the Igf2/H19 locus (37) for organizing insulator function. However, it also interacts with enhancer and promoter elements at the Igf2/H19 locus and when inserted at the Afp locus suggesting context independence (32). Our analysis confirms the context independence functionally and clearly establishes the ability of the H19-ICR to act as a position-dependent enhancer blocker in the absence of any specific cis-acting elements of the Igf2/H19 locus in vivo.

Effective organization of an insulator by the inserted H19-ICR at the TCRβ locus also led to a marked reduction in the Eβ-PDβ1-dependent recombination between Dβ1 and Jβ1 segments. As predicted by the accessibility hypothesis (24), deletion of accessibility control elements (ACE), i.e. either Eβ or PDβ1, has been shown to lead to loss of accessibility generation for transcriptional activation and subsequent recombination (22). Our results provide a strong support for the accessibility hypothesis by relying on a novel approach of altering the ACE activity rather than deletion. Modified histones acetyl-H3K9 and trimethyl-H3K4 have been observed to be associated with chromatin which is accessible for RAG mediated recombination at the antigen receptor loci (38,39). Predictably, an altered epigenetic landscape of TCRβ locus due to the ectopic insulator is the basis for the observed curtailment of Eβ-dependent transcription.

Figure 5. Altered usage of Vβ segments for V to DJ recombination at the TCRβ locus due to H19-ICR insertion. (A) Schematic diagram of the TCRβ locus showing relevant gene segments and regulatory elements along with the H19-ICR insertion. (B) Usage of Vβ segments for Vβ-to-DJββ recombination analyzed by flow cytometry. Thymocytes were immunostained to detect the presence of CD4, CD8 and TRBV31 or CD4, CD8 and pooled Vβ (TRBV2, TRBV4, TRBV12, TRBV13 and TRBV19) on the cell surface. Results from the gated CD4-SP thymocyte population (CD4+CD8+) are shown in the bar graph as relative numbers of cells that express TRBV31 or any of other Vβ that was included in the pooled Vβ (TRBV2, TRBV4, TRBV12, TRBV13 and TRBV19) in mice that maternally inherited wild-type TCRβ allele (C, +/TCRβ-del) or TCR-ins (M, TCR-ins/TCRβ-del) or TCR-mut (Mx, TCR-mut/TCRβ-del). Error bars represent ± SD between independent experiments (n = 6 for M and C and n = 3 for Mx and C). Histograms depict representative results from individual FACS experiments as described above. Numbers within the histograms denote the percentage of CD4-SP thymocytes positive for the Vβ under investigation (TRBV31 or pooled Vβ).
and recombination in TCR-ins mutant alleles. Hence, an extensive comparison of the epigenetic landscape in wild-type TCR\(b\) and TCR-ins alleles is likely to provide useful insights into the nature of ACE interactions and their ability to alter the chromatin accessibility necessary for transcription and RAG-mediated recombination. Further, the promoters PD\(b\)1, 5PD\(b\)2 and 3PD\(b\)2 are all regulated by E\(b\) but exhibit subtle differences in temporal regulation (21). Due to the drastic reduction in E\(b\)-PD\(b\)1 interaction, the TCR-ins allele provides a useful model to understand the basis for the possible competition between the promoters for the shared enhancer as well as their temporal regulation.

The most remarkable feature that emerged in this study was an alteration in the V\(b\) usage during VDJ recombination due to introduction of the four ectopic CTCF binding sites. An important role of CTCF in VDJ recombination by defining functional domains was recently demonstrated at IgH and Igk loci. At these loci, CTCF-defined chromatin loops seem to insulate the proximal V regions from the influence of enhancers E\(m\) and iE\(k\). Consequently, the usage of V segments was altered at IgH locus upon deletion of CTCF binding sites (12,40) and at Igk locus upon loss of CTCF (15) or deletion of the CTCF and Ikaros binding element Sis (41). In the present investigation, we show that the normal chromatin interactions for ordered VDJ recombination can be altered simply by the ectopic insertion of non-locus specific CTCF binding sites. Specifically, we found that TRBV31 usage was drastically enhanced in TCR-ins allele while the usage of V\(b\) segments located upstream was concomitantly reduced for recombination with DJC\(b\)2. Unlike the V regions of IgH and Igk, the V\(b\) segments of TCR\(b\) are not regulated by E\(b\) (25). Accordingly, the germline transcription of upstream V\(b\) segment (TRBV13 to DJC\(b\)2 recombination).

**Figure 6.** RSS-mediated interactions between chromatin domains defined by H19-ICR at TCR\(b\) locus as assessed by relative abundance of transcripts arising due to RSS-mediated recombination in mutants maternally inheriting wild-type TCR\(b\) allele (C, +/TCR\(b\)-del), TCR-ins (M, TCR-ins/TCR\(b\)-del) or TCR-mut (Mx, TCR-mut/TCR\(b\)-del). (A) Abundance of transcripts arising due to recombination TRBV13-D\(b\)-J\(b\)1.1 (denoted V13-DJ1) and TRBV31-D\(b\)-J\(b\)1.1 (denoted V31-DJ1). (B) Abundance of transcripts arising due to recombination TRBV13-D\(b\)-J\(b\)2.7 (denoted V13-DJ2) and TRBV31-D\(b\)-J\(b\)2.7 (denoted V31-DJ2). (C) Abundance of recombined transcripts V31-DJ2 relative to V13-DJ2. In each case, relative abundance was estimated by real-time quantitative RT-PCR analysis. Error bars denote ± SEM. (D) Schematic of TCR\(b\) locus in wild-type (upper) and TCR-ins (lower) mice. Arrows indicate the long range interactions necessary for promoter activation (black arrows) and RSS-mediated recombination (grey arrows) between V\(b\) and DJC\(b\) regions. H19-ICR presence in TCR-ins drastically reduces E\(b\) based PD\(b\)1 activation. It also defines chromatin domains such that ‘intradomain’ interaction (TRBV31 to DJC\(b\)2 recombination) is significantly increased at the expense of ‘interdomain’ interaction (TRBV13 to DJC\(b\)2 recombination).
insertion of an enhancer blocker. Also, the transcriptional status of TRBV31, located relatively close to Εβ, was not altered in the TCR-ins allele. Yet the Vβ-to-Dβ junction recombination profile was completely altered. The most plausible explanation for the altered Vβ usage in TCR-ins allele appears to be the CTCF-directed chromatin domain organization such that the RSS-mediated V-to-DJ recombination is efficient only when V and DJ segments share the same domain defined by CTCF (Figure 6). This suggests that insulators may have the potential to organize specific functional domains with respect to recombination at the antigen receptor loci in addition to their well known ability to define transcriptionally distinct domains. Taken together with the recent evidence from IgH and Igx loci, these data present an emerging picture of CTCF and insulator elements governing multiple chromatin interactions at the recombining loci.

Chromatin loop organization is mediated by multiple CTCF and cohesin binding sites at the recombining loci (16,42). Based on analysis of IgH locus, it has been proposed that CTCF organizes dynamic rosette structures to facilitate access to all Vh segments for recombining with DβJh segments (43). CTCF manifests its influence by organizing chromatin loops (2). Consequently, the interactions between DNA elements can be promoted as well as hindered depending on their positions relative to the CTCF-binding insulator (2,44). Keeping this in mind, we designed the TCR-ins allele such that the H19-ICR insulator effectively partitioned the TCRβ locus. We observed increased intradomain and reduced interdomain functional interactions between promoter–enhancer regulatory elements as well as between RSS elements across the H19-ICR insertion in the TCR-ins allele. Thus, CTCF appears to have three activities that impact VDJ recombination: (i) enhancer blocking that regulates ordered assembly of genes (12), (ii) contraction of loci prior to recombination (13,14,16) and (iii) organization of chromatin domains that define the choice of recombining gene segments as observed during this investigation. These activities complement each other and emphasize the importance of CTCF for VDJ recombination. More detailed investigations of the interactions between the ectopic and endogenous CTCF binding sites at the TCRβ locus as well as at other antigen receptor loci will be meaningful to elucidate these roles in future.

Finally, our results also provide useful insights into the nature of interactions between distantly placed cis-acting regulatory elements which have, so far, been considered only in the context of transcriptional regulation using chromosome conformation capture (3C) analysis (32,45–47). Enhancers can activate their cognate promoters in a contact-dependent and contact-independent manner according to the topological loop domain and tracking models, respectively. Accordingly, the intervening insulator may either organize the chromatin domains to abrogate enhancer–promoter contact or prevent the tracking signal from reaching the cognate promoter (48). A fundamental assumption of the topological model is that ‘intradomain’ contacts are favored over ‘interdomain’ contacts when an insulator defines the distinct domains. This has been difficult to test functionally as reduction in interdomain promoter–enhancer contacts by the intervening insulator cannot be unambiguously discriminated from the block in tracking by an intervening insulator; both the situations would lead to a loss in transcriptional activation.

In this context, it was intriguing to note that the functional H19-ICR insulator was able to hinder RSS contact mediated Vβ-to-Dβ recombination across itself just as it prevents enhancer–promoter functional interaction. VDJ recombination unambiguously relies on a contact based RSS interaction and ‘captures’ the physical juxtaposition of the interacting RSS elements in vivo reporting it as a specific recombination event. When H19-ICR defined the chromatin domains at the TCRβ locus, the contact between RSS elements and consequent Vβ-to-Dβ recombination, was strongly favored between RSS elements sharing the same chromatin domain (intradomain) rather than those separated by the insulator (interdomain; Figure 6D). Mechanistically, RSS-mediated recombination and enhancer-mediated promoter activation are entirely distinct. The only common feature between them is that both require interaction of specific cis-acting DNA elements. The ability of the insulator to interfere in a position-dependent manner with each of these distinct processes provides functional evidence to support the topological model of interaction of cis acting regulatory elements.

In conclusion, our analysis clearly demonstrates a previously unreported ability of a CTCF-dependent insulator to modulate interactions between RSS elements during VDJ recombination. This emphasizes that the CTCF binding sites may have a complex role in influencing functional interactions between regulatory elements beyond modulating enhancer–promoter interactions and defining transcriptionally independent domains in the genome.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online: Supplementary Table 1 and Supplementary Figures 1 and 2.

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