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Involvement of the TCR C β FG Loop in Thymic Selection and T Cell Function

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Abstract

The asymmetric disposition of T cell receptor (TCR) C β and C α ectodomains creates a cavity with a side-wall formed by the rigid C β FG loop. To investigate the significance of this conserved structure, we generated loop deletion ($\beta\Delta$ FG) and β wt transgenic (tg) mice using the TCR β subunit of the N15 CTL. N15 β wt and N15 $\beta\Delta$ FG H-2^b animals have comparable numbers of thymocytes in S phase and manifest developmental progression through the CD4⁻CD8⁻ double-negative (DN) compartment. N15 $\beta\Delta$ FG facilitates transition from DN to CD4⁺CD8⁺ double-positive (DP) thymocytes in recombina-activating gene (RAG)-2^{-/-} mice, showing that pre-TCR function remains. N15 $\beta\Delta$ FG animals possess \sim twofold more CD8⁺ single-positive (SP) thymocytes and lymph node T cells, consistent with enhanced positive selection. As an altered V α repertoire observed in N15 $\beta\Delta$ FG mice may confound the deletion's effect, we crossed N15 $\alpha\beta$ TCR tg RAG-2^{-/-} with N15 $\beta\Delta$ FG tg RAG-2^{-/-} H-2^b mice to generate N15 $\alpha\beta$ RAG-2^{-/-} and N15 $\alpha\beta.\beta\Delta$ FG RAG-2^{-/-} littermates. N15 $\alpha\beta.\beta\Delta$ FG RAG-2^{-/-} mice show an 8–10-fold increase in DP thymocytes due to reduced negative selection, as evidenced by diminished constitutive and cognate peptide-induced apoptosis. Compared with N15 $\alpha\beta$, N15 $\alpha\beta.\beta\Delta$ FG T cells respond poorly to cognate antigens and weak agonists. Thus, the C β FG loop facilitates negative selection of thymocytes and activation of T cells.

Key words: negative selection • thymocyte development • TCR structure • Ig superfamily • transgenic mice

Introduction

The TCR is a multicomponent complex composed of a Ti disulfide-linked heterodimer in noncovalent association with the CD3 subunits (1–5). The majority of T cells express a Ti encoded by α and β TCR genes (Ti $\alpha\beta$) whereas a numerically minor but functionally significant subset expressed Ti encoded by γ and δ TCR genes (Ti $\gamma\delta$; for a review, see reference 6). The Ti $\alpha\beta$ heterodimer whose overall shape and architecture are similar to an antibody Fab fragment represents the peptide–MHC complex (pMHC)* ligand binding unit. Thus, Ti $\alpha\beta$ determines the ligand specificity of an individual T cell. In contrast, the CD3

polypeptides mediate TCR-based signal transduction. The signaling function of the CD3 components involves a conserved motif, termed immunoreceptor tyrosine-based activation motif (ITAM), present at 1–3 copies in the cytoplasmic domain of each CD3 subunit (7–9). The various CD3 subunits exhibit different interactions with intracellular signaling factors and, acting in concert, lead to T cell activation (10–15). However, how pMHC ligand binding to a Ti $\alpha\beta$ or Ti $\gamma\delta$ heterodimer subsequently initiates signaling via the CD3 molecules is currently unknown.

The TCR complex contains one copy of CD3 γ and CD3 δ and two copies of CD3 ϵ and CD3 ζ configured as two CD3 $\epsilon\gamma$ and CD3 $\epsilon\delta$ heterodimers and a disulfide-linked CD3 ζ homodimer (16–21). Recent crystallographic analysis (22) in conjunction with epitope mapping of a class I-restricted TCR (23) have identified a cavity within the Ti $\alpha\beta$ constant (C) module of sufficient size to accommodate one nonglycosylated CD3 ϵ ectodomain. In this regard, the solution structure of a heterodimeric CD3 $\epsilon\gamma$

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*Abbreviations used in this paper: aa, amino acid; APL, altered peptide ligand; BrdU, bromodeoxyuridine; DN, double negative; DP, double positive; PI, propidium iodide; pMHC, peptide–MHC complex; RAG, recombina-activating gene; SP, single positive.

ectodomain complex reveals a unique side-to-side hydrophobic interface between the two C2-set immunoglobulin-like domains and indicates that a modular pair-wise association is shared between CD3 $\epsilon\delta$ as well (24). CD3 ϵ has an elongated shape with length, width and depth being ~ 40 , 25, and 25 Å, respectively.

Herein, we have investigated the biological significance of the unusual structural arrangement of the C β FG loop that forms the side wall of this C-module cavity. Mapping studies using the C β FG loop-specific mAb H57 previously showed that one of two CD3 ϵ subunits resides adjacent to this structural element. To this end, we have created TCR transgenic mice bearing either intact or C β FG loop-deleted β chains. Our results show that the loss of the C β FG loop attenuates negative selection without blocking positive selection within the thymus at the level of double-positive (DP) thymocytes. The sensitivity of mature T cells to cognate peptides as well as altered peptide ligands is also reduced. In contrast, pre-TCR facilitated transition from CD44⁻CD25⁺ to CD44⁻CD25⁻ double-negative (DN) thymocytes and subsequent DP thymocyte generation is not altered. The implications of these findings for TCR and pre-TCR function are discussed.

Materials and Methods

T Cell Receptor Construct. The FG loop-deleted mutant of the N15 TCR β chain constant region (C β) was generated by PCR using N15 β wt cDNA as a template, subcloned into the pCRII vector, and verified by DNA sequencing analysis. The *Stu*I digested DNA fragment containing the FG loop-deleted constant region of the TCR β chain was subcloned into the *Stu*I site in the constant region of an N15 TCR β expression vector, which was previously used for the generation of N15 TCR $\alpha\beta$ tg recombinase activating gene (RAG)-2^{-/-} H-2^b mice (25).

Mice. Mice transgenic for the N15 wt or FG loop-deleted TCR β chain (hereafter for simplicity referred to as N15 β wt or N15 $\beta\Delta$ FG) were produced in the transgenic/gene targeting core facility at Dana-Farber Cancer Institute. The N15 TCR β shuttle constructs with the wt or FG loop-deleted mutant C β were digested with *Pvu*I/*Sal*I, leaving the bacterial vector sequences attached to the TCR gene. These DNAs were injected into (SJL \times C57BL/6) F1 fertilized eggs and implanted into foster mothers. Resulting progeny were screened for integration of the N15 wt or FG loop-deleted TCR β chain DNA by Southern blot analysis of tail DNA and by FACS[®] analysis of N15 TCR β chain expression in peripheral blood nucleated cells. Five transgenic founders were identified, two for N15 β wt, and three for N15 $\beta\Delta$ FG. The mice were backcrossed for 4–5 generations onto the C57BL/6 background before the experiments described below. To create N15 $\beta\Delta$ FG RAG-2^{-/-} mice, the N15 $\beta\Delta$ FG mice were bred with RAG-2^{-/-} C57BL/6 mice and the heterozygous N15 $\beta\Delta$ FG^{+/+} RAG-2^{+/-} F1 mice were intercrossed and offspring which were N15 $\beta\Delta$ FG^{+/+} RAG-2^{-/-} subsequently selected. The expression of the FG loop-deleted N15 β chain and lack of RAG-2 were confirmed based on the Southern blot analysis of tail DNA and FACS[®] analysis of peripheral blood cells. To introduce the N15 α chain into N15 $\beta\Delta$ FG mice and generate Ag-specific T cells with the FG loop-deleted TCR β chain, the N15 $\beta\Delta$ FG RAG-2^{-/-} (N15 $\beta\Delta$ FG^{+/+} RAG-2^{-/-}) mice were bred with N15 TCR $\alpha\beta$ tg RAG-2^{-/-} H-2^b (N15 α wt^{+/+} β wt^{+/+} RAG-2^{-/-}

H-2^b) mice, and the resulting N15 α wt^{+/+} β wt^{+/+} RAG-2^{-/-} H-2^b or N15 α wt^{+/+} β wt^{+/+} $\beta\Delta$ FG^{+/+} RAG-2^{-/-} H-2^b (hereafter for simplicity referred to as N15 $\alpha\beta$ or N15 $\alpha\beta$. β FG mice) were used for the experiments. RAG-2^{-/-} C57BL/6 mice were purchased from Taconic. All lines were maintained and bred under sterile barrier conditions at the animal facility of Dana-Farber Cancer Institute.

Peptide Synthesis. Peptides were synthesized by standard solid phase methods on an Applied Biosystems 430A synthesizer at the Biopolymers Laboratory of the Massachusetts Institute of Technology. All peptides were purified by reverse phase HPLC (Hewlett Packard HPLC 1100) with a C4, 2-mm column. Peptides were analyzed for purity and correct molecular weight by electrospray mass spectrometry, amino acid (aa) analysis, and HPLC. Peptides are named to indicate the substituted aa and the position in the sequence of VSV8 (e.g., L4 denotes replacement of valine by leucine at the fourth residue of the VSV8 peptide). VSV8 is a cognate antigenic peptide agonist of the N15 TCR, which triggers thymic negative selection in N15 TCR $\alpha\beta$ tg RAG-2^{-/-} H-2^b mice, whereas L4 and K1 peptides are altered peptide ligands (APLs) of VSV8 and weak agonists of the N15 TCR, which induce positive selection in N15 TCR $\alpha\beta$ tg RAG-2^{-/-} β_2 M^{-/-} H-2^b mice (26).

Antibodies and Flow Cytometric Analysis. The following mAbs were used: R-phycoerythrin (PE)- or Cychrome-conjugated anti-mouse CD4 (H129.19); FITC- or Cychrome-conjugated anti-mouse CD8 α (53–6.7); Cychrome-conjugated anti-TCR C β (H57–597); PE-conjugated anti-CD25 (PC61); FITC-conjugated anti-CD44 (IM7); FITC-conjugated anti-V β 5.1, 5.2 (MR9.4); biotin-conjugated anti-CD69 (H1.2F3); biotin-conjugated anti-V α 2 (B20.1), anti-V α 3.2 (RR3–16), anti-V α 8.3 (B21.14), anti-V α 11.1, 11.2 (RR8–1), PE-conjugated streptavidin (BD Pharmingen). For flow cytometry, single cell thymocyte or lymph node suspensions were prepared in PBS containing 2% FCS and 0.05% NaN₃. Cells were stained at 5 \times 10⁶ cells per ml in PBS-2% FCS and 0.05% NaN₃ containing the antibodies at saturating concentrations. Phenotypes and proportions of thymocyte and lymph node cell subsets were analyzed by two-color flow cytometry using a FACScan[™] (Becton Dickinson) and the CELLQuest[™] program. Dead cells were excluded from the analysis by forward and side scatter gating.

Bromodeoxyuridine Staining. Bromodeoxyuridine (BrdU) incorporation was examined according to the manufacturer's protocol (BrdU Flow Kit; BD Pharmingen). Mice were administered two intraperitoneal injections of 1 mg of BrdU in PBS at 4 h intervals. After 18 h, thymocytes were fixed/permeabilized and stained with FITC-conjugated anti-BrdU mAb following treatment with 30 μ g/ml DNase. Live cells were gated and analyzed on a FACScan[™].

Apoptosis Assay. Thymocytes from untreated, VSV8 peptide-injected (0.5 μ g intravenously, 18 h previously) or dexamethasone-treated (0.5 mg intraperitoneally, 6 h previously) were examined. Apoptosis of thymocytes was evaluated by cellular DNA content determined by propidium iodide (PI) staining and alterations in plasma membrane permeability examined by Annexin V staining. For PI staining, after fixation with 40% ethanol at 4°C for 30 min, total thymocytes (10⁶ cells) were resuspended in PBS containing 50 μ g/ml RNase A and 2.5 μ g/ml PI and incubated at 37°C for 30 min. PI incorporation was analyzed by flow cytometry. For annexin V staining, after washing with PBS, total thymocytes (10⁶ cells) were resuspended in 1 \times binding buffer containing 0.5 μ g/ml FITC-conjugated annexin V (ApoAlert Annexin V Apoptosis kit; CLONTECH Laboratories,

Inc.) and 2.5 $\mu\text{g/ml}$ PI to gate out dead cells, and incubated in the dark for 10 min at room temperature, followed by the analysis on a FACScan™.

Proliferation Assay. Lymph node cells from N15 $\alpha\beta$ or N15 $\alpha\beta$ - $\beta\Delta\text{FG}$ mice ($10^5/\text{well}$) were incubated at 37°C with 2×10^4 irradiated EL-4 thymoma cells, which had been preloaded for 2 h with the indicated doses of VSV8 or a weak agonist, L4, in AIM-V medium (Life Technologies) containing 50 μM 2-ME. After 48 h incubation, 0.4 μCi per well of [^3H]TdR (ICN Bio-medicals) was added, and after an additional 18 h of culture at 37°C, the cells were harvested and the incorporated radioactivity was measured.

Measurement of IFN- γ . IFN- γ production was induced under the same culture conditions used for proliferation assays (see above). After 48 h of incubation, supernatants were collected and assayed for IFN- γ using a mouse IFN- γ ELISA kit (Mouse IFN- γ OptEIA Set; BD PharMingen). The sensitivity of the assay was

31.3–2,000 pg/ml and results were calculated as the mean of duplicate wells.

Results

The TCR C β FG Loop Structure and Sequence Comparison. One remarkable feature of the TCR C β domain is the striking elongation of the FG loop. Compared with other Ig-like structures (aside from the TCR C γ domain), there is a 13 aa insertion within the FG loop (27). Fig. 1 A shows a view of the crystallographically resolved murine N15 class I MHC-restricted $\alpha\beta$ TCR (V β 5.2 D2J β 2.6 C β 2/V α 8 J α 5C α ; reference 22). The unique protrusion in the C β domain's FG loop is apparent (bracketed in a black box). It was previously noticed that internally the loop is well structured (22). The Trp225 at the tip of the loop

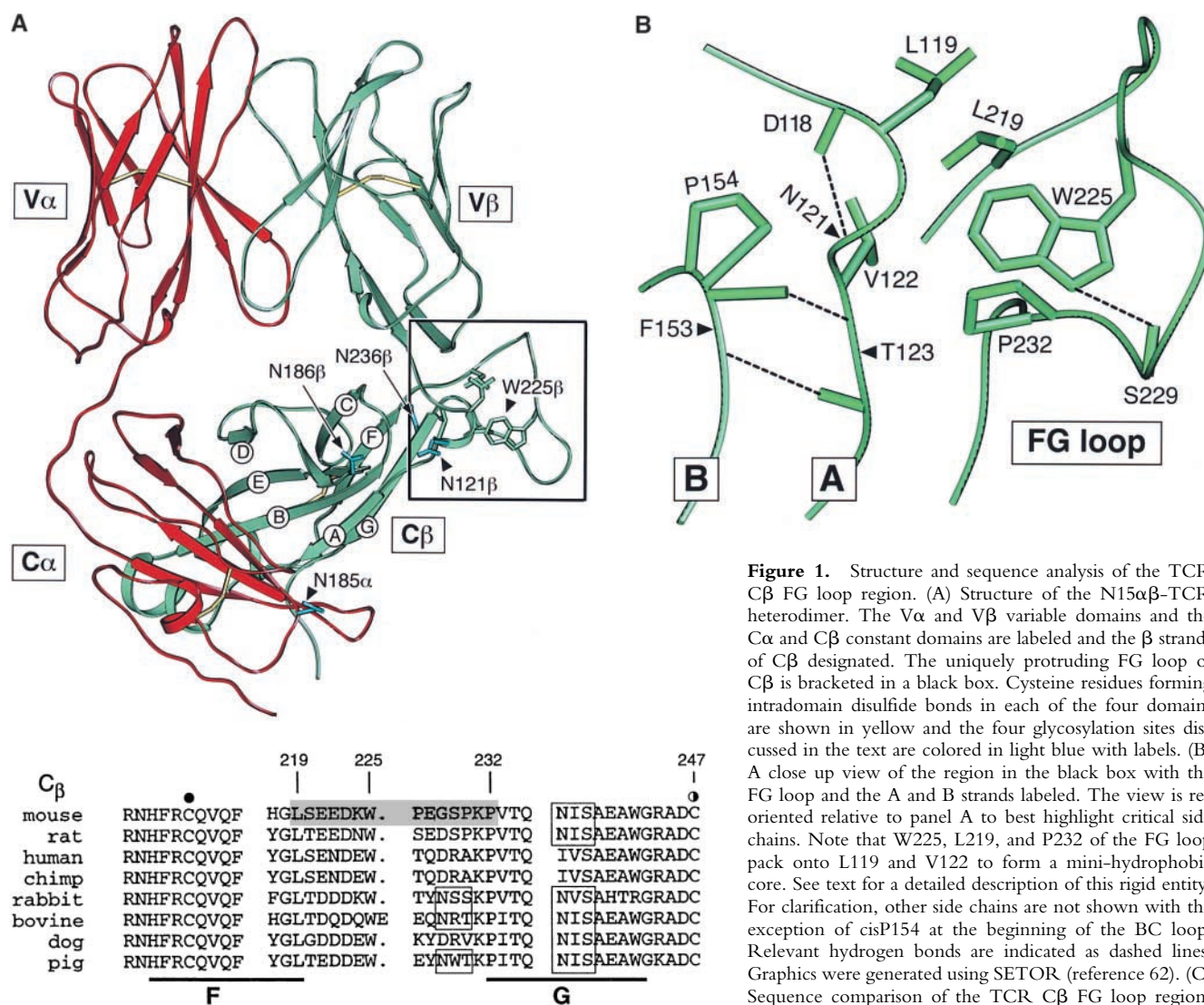


Figure 1. Structure and sequence analysis of the TCR C β FG loop region. (A) Structure of the N15 $\alpha\beta$ -TCR heterodimer. The V α and V β variable domains and the C α and C β constant domains are labeled and the β strands of C β designated. The uniquely protruding FG loop of C β is bracketed in a black box. Cysteine residues forming intradomain disulfide bonds in each of the four domains are shown in yellow and the four glycosylation sites discussed in the text are colored in light blue with labels. (B) A close up view of the region in the black box with the FG loop and the A and B strands labeled. The view is re-oriented relative to panel A to best highlight critical side chains. Note that W225, L219, and P232 of the FG loop pack onto L119 and V122 to form a mini-hydrophobic core. See text for a detailed description of this rigid entity. For clarification, other side chains are not shown with the exception of cisP154 at the beginning of the BC loop. Relevant hydrogen bonds are indicated as dashed lines. Graphics were generated using SETOR (reference 62). (C) Sequence comparison of the TCR C β FG loop regions from various species. The position of the F and G strands is

defined based on the above crystallographic analysis. The shaded region defines the 14 aa residue deleted from the N15 TCR β chain. Boxed residues mark the potential N-linked glycosylation signals. The β chain cysteine (247) which forms an interchain disulfide with the α chain, is indicated by the semi-solid circle and the F strand cysteine (212) which forms the intradomain disulfide with the β strand cysteine (147) of C β (not shown) is indicated by the solid circle.

plays a key role. A hydrogen bond between the NE1 of Trp225 and the carbonyl group of Ser229 brings the bulky indole ring of Trp225 in position to make hydrophobic contacts with Leu219 and Pro232 at the base of the FG loop (Fig. 1 B, a close up view of the local region shown in the black box of Fig. 1 A). The hydrophobic patch composed of these three FG loop residues packs onto Leu119 and Val122 at the beginning of the C β domain's A strand to form a mini-hydrophobic core. The Leu119 and Val122 themselves lie on a stable local strut created by a tight β turn between Asp118 and Asn121. Moreover, a pair of mainchain hydrogen bonds between Thr123 on the A strand and Phe153 at the end of the B strand further stabilizes the local structure. Collectively, the C β domain FG loop, albeit a long protrusion, is an integrated component of a rigid structural entity that connects the V β and C β domains. Not only are the residues mentioned above all invariant among mouse, rat, human, chimpanzee, rabbit, cow, dog, and pig as shown in the Fig. 1 C, but the three-dimensional structural feature depicted in Fig. 1 B is also conserved in all known TCR crystal structures, including class I MHC-restricted N15 and 2C from mouse (22, 28) and B7 from human (29), as well as the human class II MHC-restricted HA1.7 (30).

T Lineage Development in N15 β Δ FG Mice. To analyze the role of the C β FG loop in thymocyte development and mature T cell function in vivo, we established transgenic mice expressing the wild-type N15 TCR β chain or an FG loop deletion variant lacking aa 219–232. For simplicity, these TCR transgenes will be referred to as N15 β wt and N15 β Δ FG, respectively. The N15 β wt subunit derives from the N15 CD8⁺ cytotoxic T cell clone whose TCR is specific for the vesicular stomatitis virus nucleoprotein aa 52–59 (VSV8) (RGYVYQGL) bound to the MHC class I molecule H-2K^b (31). T lineage cells from the thymus and lymph nodes of N15 β Δ FG mice were then directly compared with those of the N15 β wt mice. Fig. 2 A shows a flow cytometric analysis of CD8⁺ single-positive (SP) lymph node T cells from 1 of 2 representative N15 β wt lines and 1 of 3 representative N15 β Δ FG lines. As expected, the anti-C β FG loop specific mAb H57 (22, 32) reacts with virtually all CD8⁺ SP lymph node T cells in N15 β wt tg mice but almost no CD8⁺ SP lymph node T cells in N15 β Δ FG mice. Those few reactive T cells (6.8%) represent CD8⁺ SP cells that have rearranged endogenous β chains due to incomplete TCR β chain allelic exclusion by the N15 β transgene, as previously reported in other transgenic systems (33). Fig. 2 A also indicates that despite the C β FG loop deletion, N15 β Δ FG tg animals, like N15 β wt tg mice, predominantly express the transgenic V β 5.2 gene as detected by the MR9.4 mAb.

Perhaps more importantly, as shown in Fig. 2 B and Table I, comparison of thymocyte subsets reveals substantial differences between the N15 β wt and N15 β Δ FG tg mice. At 6–12 wk of age, for example, N15 β Δ FG mice show a subtle but significant reduction in the number of total thymocytes (N15 β wt, $12.05 \pm 2.99 \times 10^7$ [$n = 10$]; N15 β Δ FG, $9.51 \pm 2.09 \times 10^7$ [$n = 15$], $P < 0.05$; Table

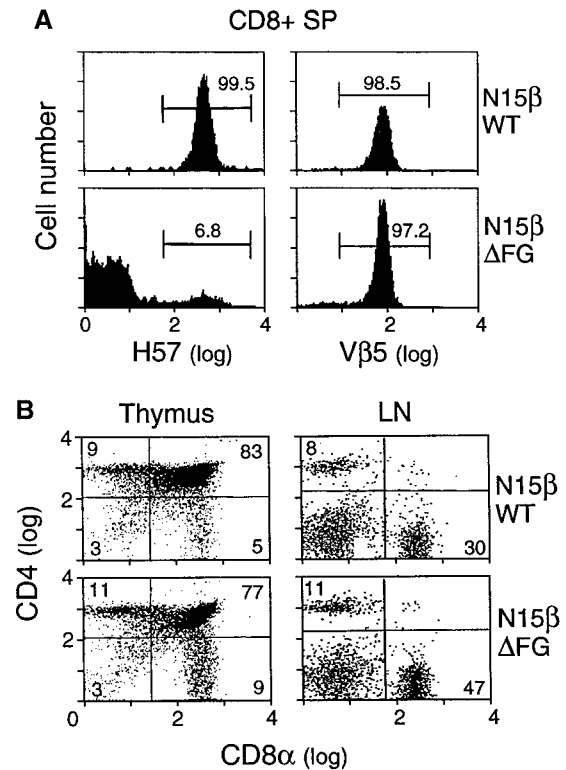


Figure 2. Altered phenotypes of T lineage cells in the N15 β Δ FG mice. (A) TCR expression on CD8⁺ SP lymph node T cells in the N15 β wt and N15 β Δ FG mice. Lymph node cells from 8-wk-old N15 β wt and N15 β Δ FG mice were triple-stained with anti-CD4, anti-CD8, and anti-TCR C β FG loop specific H57 or anti-V β 5 MR9.4 mAbs. The histograms of TCR C β FG loop (left) and V β 5 (right) expression on the gated CD8⁺ SP lymph node T cells are shown. The numbers represent the percentages of H57- or V β 5-positive cells (as indicated). (B) The CD4/CD8 α double-staining profiles in thymocytes and lymph node cells are altered by the lack of the TCR C β FG loop. The thymocytes and lymph node cells from N15 β wt and N15 β Δ FG mice at 8 wk of age were double-stained with PE-anti-CD4 and FITC-anti-CD8 α . The expression of CD4 (y axis) and CD8 α (x axis) in thymocytes and lymph node cells was detected by flow cytometry after gating on 10,000 live cells. The percentages of each subset are indicated.

I). Furthermore, as shown by the representative CD4/CD8 α double-staining profiles in Fig. 2 B, there is a detectable increase in the percentage of the CD8⁺ SP subset and a decrease in that of the DP subset in N15 β Δ FG thymocytes compared with N15 β wt mice. Therefore, N15 β Δ FG mice show a statistically significant increase in the absolute number of CD8⁺ SP thymocytes and a concomitant decrease in DP thymocytes (Table I). This increase in CD8⁺ SP cells is further reflected by the increase in CD8⁺ SP lymph node T cells (N15 β wt, $1.99 \pm 0.77 \times 10^7$ [$n = 10$]; N15 β Δ FG $3.45 \pm 0.82 \times 10^7$ [$n = 15$]). From these findings, we conclude that the absence of the FG loop in the constant region of N15 TCR β chain influences T cell development in the tg mouse model.

There are two major checkpoints at which $\alpha\beta$ thymocyte development is dependent upon the TCR β chain. The earliest occurs during the transition from DN to DP thymocytes and is regulated by the pre-TCR complex

Table 1. Absolute Numbers of Total and Cell Subsets of Thymocytes and LN Cells in N15βwt and N15βΔFG Mice

	Thymus					LN		
	Total ^a	DP ^b	CD8 ⁺ SP ^a	CD4 ⁺ SP	DN	Total	CD8 ⁺ SP ^c	CD4 ⁺ SP
N15βwt	12.05 ± 2.99 ^d	9.95 ± 2.47	0.60 ± 0.17	1.08 ± 0.30	0.41 ± 0.16	6.36 ± 2.11	1.99 ± 0.77	0.49 ± 0.19
N15βΔFG	9.51 ± 2.09	7.31 ± 1.61	0.85 ± 0.29	1.02 ± 0.27	0.33 ± 0.12	7.17 ± 1.83	3.45 ± 0.82	0.71 ± 0.21

^a*P* < 0.05.^b*P* < 0.01.^c*P* < 0.0005.

^dThe numbers of cell subsets in thymocytes and lymph node T cells were calculated by quantifying the total number of thymocytes and lymph node cells from 6–12-wk-old N15β (*n* = 10) and N15βΔFG (*n* = 15) mice and the percentages of each subset as determined by FACS[®] analysis. Results are expressed as mean ± SD (×10⁷ cells). Statistical differences between N15βwt and N15βΔFG numbers in indicated populations are given.

comprised of the invariant pTα subunit disulfide linked to a TCR β chain in noncovalent association with the CD3 subunits (34). The later checkpoint involves the αβ TCR and occurs during the transition from DP to SP thymocyte maturation. At that time, self-MHC-restricted, nonautoreactive T cells are selected for maturation and peripheral exportation (35). To investigate the mechanism of the altered thymocyte phenotype in N15βΔFG mice, we first compared early thymic development in the N15βwt and N15βΔFG mice by examining the fraction of cells synthesizing DNA using BrdU incorporation in vivo. In this way, it is possible to test whether the FG loop deletion in the N15 TCRβ chain affects proliferation and/or turnover of thymocytes. The N15βwt and N15βΔFG tg mice were injected twice intraperitoneally with 1 mg of BrdU in PBS at 4 h intervals, and 18 h later, thymocytes were assayed for the incorporation of BrdU by flow cytometry. As shown in Fig. 3 A, the percentage of BrdU⁺ thymocytes in the N15βΔFG mice was comparable to that observed in control N15βwt mice, indicating that the number of cells that had progressed through S phase is similar. In addition, we compared the developmental stages of immature thymocytes by examining the surface expression of CD25 and CD44 on DN thymocytes of these mice. The percentages of the four CD44/CD25 subsets (CD44⁺CD25⁻, CD44⁺CD25⁺, CD44⁻CD25⁺, and CD44⁻CD25⁻) in the N15βΔFG were equivalent to those of N15βwt mice (Fig. 3 B), suggesting that the deletion of Cβ FG loop from the N15 TCRβ chain does not disrupt the pre-TCR-facilitated transition from CD44⁺CD25⁺ to CD44⁻CD25⁻ DN thymocytes. To further assess the function of the Cβ FG loop-deletion in pre-TCR signaling at the CD44⁻CD25⁺ DN stage, we established RAG-2-deficient mice expressing the Cβ FG loop-deleted N15β chain (N15βΔFG RAG-2^{-/-}). Introduction of a functionally rearranged wild-type TCRβ chain overcomes the developmental arrest at the CD44⁻CD25⁺ DN stage observed in the RAG-2-deficient mice, leading to strong proliferation of DN thymocytes and efficient generation of DP cells (36). As shown in Fig. 3 C, N15βΔFG RAG-2^{-/-} mice generate DP thymocytes with essentially normal cellularity

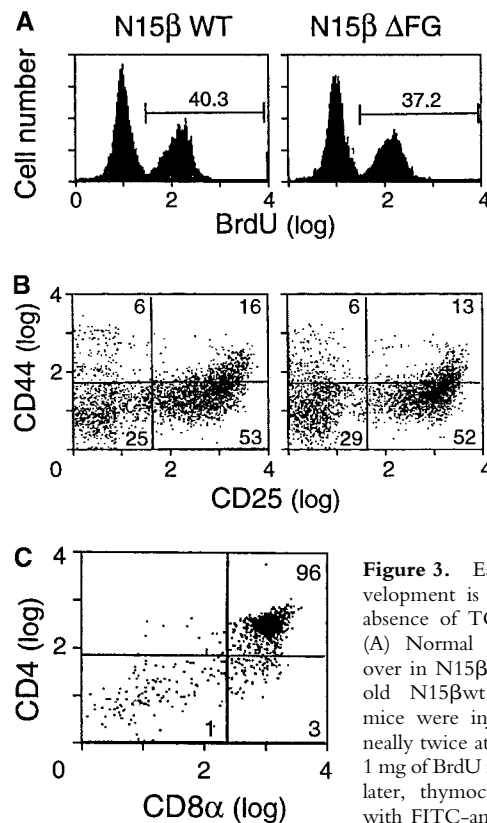


Figure 3. Early thymocyte development is unaffected by the absence of TCR Cβ FG loop. (A) Normal proliferation/turnover in N15βΔFG mice. 9-wk-old N15βwt and N15βΔFG mice were injected intraperitoneally twice at 4 h intervals with 1 mg of BrdU in 0.2 ml PBS. 18 h later, thymocytes were stained with FITC-anti-BrdU mAb and assayed by flow cytometry. The histograms of BrdU incorporation on 10,000 live thymocytes are shown. The numbers represent the percentages of BrdU-positive cells. (B) The transition from CD44⁺CD25⁺ to CD44⁻CD25⁻ DN thymocytes is not blocked in N15βΔFG mice. Total thymocytes were triple-stained with FITC-anti-CD44, PE-anti-CD25, and Cychrome-anti-CD4/anti-CD8α. The profiles of CD44 (y axis) versus CD25 (x axis) expression in Cychrome-negative (CD4⁻CD8⁻ DN) thymocytes are shown. The percentage of each subset is indicated. (C) Efficient generation of DP thymocytes in N15βΔFG RAG-2^{-/-} mice. Thymocytes from 8-wk-old N15βΔFG RAG-2^{-/-} mice were double-stained with PE-anti-CD4 and FITC-anti-CD8α. The expression of CD4 (y axis) and CD8α (x axis) on thymocytes was detected by flow cytometry after gating on 10,000 live cells. The percentages of each subset are indicated.

(6.6×10^7 thymocytes/mouse). These results suggest that signaling through the pre-TCR complex on CD44⁻CD25⁺ thymocytes is not blocked by the deletion of the C β FG loop in the N15 TCR β chain.

Altered V α Usages by Peripheral Mature T Cells in the N15 β Δ FG Mice. Given the nearly twofold increase in the absolute number of CD8⁺ SP lymph node T cells (Table I), we next examined whether deletion of the C β FG loop of the N15 β chain influences thymic repertoire selection. To this end, the TCR V α gene usage of CD8⁺ and CD4⁺ SP lymph node T cells in N15 β wt and N15 β Δ FG mice was assessed by flow cytometry using a panel of anti-V α mAbs. As shown in Table II, there are statistically significant differences in the TCR V α gene usage in peripheral lymph node T cells between N15 β wt and N15 β Δ FG mice. CD8⁺ lymph node T cells from N15 β Δ FG mice display an increased frequency of TCRs incorporating V α 8.3 (mean = 4.37%, SD = 0.58 versus mean = 3.74%, SD = 0.54; $P < 0.02$) and V α 11 (mean = 6.81%, SD = 0.70 versus mean = 5.46%, SD = 0.50; $P < 5 \times 10^{-5}$) gene segments compared with N15 β wt mice (Table II). CD4⁺ lymph node T cells from N15 β Δ FG mice also bear more TCRs incorporating V α 2 (mean = 13.61%, SD = 0.95 versus mean = 11.77%, SD = 1.38; $P < 0.005$) than those from N15 β wt mice. These data suggest that lack of the C β FG loop exerts an influence on selection of the TCR repertoire by affecting usage of functional V α gene segments, leading to the alterations in thymic and peripheral lymph node phenotypes observed in the N15 β Δ FG mice. That V α differences are observed in CD8⁺ SP thymocytes as well as lymph node cells is consistent with this view (data not shown), but does not exclude the possibility that impaired TCR signaling in the peripheral T cells in N15 β Δ FG animals further alters the repertoire.

Quantitative and Qualitative Alterations of Thymocytes and Peripheral T Cells in N15 α β . β Δ FG Mice. The differences between N15 β wt and N15 β Δ FG mice may have been modified by compensatory adaptations of V α gene expression. To analyze more clearly the role of the TCR C β FG loop in thymocyte development and mature T cell function, we next crossed the N15 β Δ FG RAG-2^{-/-} mice

(N15 β Δ FG^{+/-}RAG-2^{-/-} H-2^b) with N15 α β tg RAG-2^{-/-} H-2^b (N15 α wt^{+/+} β wt^{+/+}RAG-2^{-/-} H-2^b) to generate two mouse strains: N15 α wt^{+/-} β wt^{+/-} RAG-2^{-/-} and N15 α wt^{+/-} β wt^{+/-} β Δ FG^{+/-}RAG-2^{-/-} mice. These mice are referred to hereafter for simplicity as N15 α β and N15 α β . β Δ FG, respectively. In these strains, the TCR α chain is fixed and on the RAG-2^{-/-} background, so no confounding endogenous β or α chains can be expressed.

T lineage cells from the thymus and lymph nodes of these animals are directly compared in Fig. 4, A and B. As defined by the anti-V β 5 mAb MR9.4, using quantitative immunofluorescence on FACS[®], the CD8⁺ SP lymph node T cells from N15 α β . β Δ FG mice expressed the same level of N15 β chain as CD8⁺ SP T cells derived from N15 α β mice. However, the fluorescence intensity of anti-TCR C β H57 mAb staining in the CD8⁺ SP T cells from N15 α β . β Δ FG mice is \sim 50% that in N15 α β T cells, indicating that CD8⁺ SP T cells from N15 α β . β Δ FG mice express both wild-type and FG loop-deleted N15 TCR β chains complexed with N15 TCR α chains at almost equal levels on their cell surface (Fig. 4 A). Although not shown, all peripheral T cells are CD8⁺ SP in both N15 α β and N15 α β . β Δ FG mice with equivalent surface CD8 density as revealed by the anti-CD8 α mAb 53–6.7.

Comparison of the thymocyte cell number and subset distribution among N15 α β or N15 α β . β Δ FG mice reveal striking differences. At 8–12 wk of age, N15 α β . β Δ FG mice have a markedly larger thymus relative to that of N15 α β animals with an \sim sevenfold increase in the number of total thymocytes (Fig. 4 C). Furthermore, as shown by the CD4/CD8 α double-staining profiles in Fig. 4 B, there is a substantial increase in the DP subset percentage and a decrease in the DN and CD8⁺ SP subset percentages in N15 α β . β Δ FG thymocytes. As the total number of N15 α β . β Δ FG thymocytes is \sim 7 times more than N15 α β thymocytes, N15 α β . β Δ FG mice show no significant differences in the absolute numbers of DN and CD8⁺ SP thymocytes, but \sim 10-fold more DP thymocytes than N15 α β mice (N15 α β , 1.48×10^7 DP cells versus N15 α β . β Δ FG, 13.2×10^7 DP cells; Fig. 4 C). These results indicate that the absence of the TCR C β FG loop influences intrathy-

Table II. V α Usages by CD8⁺ SP and CD4⁺ SP Lymph Node Cells in N15 β wt and N15 β Δ FG

	CD8 ⁺ SP				CD4 ⁺ SP			
	V α 2	V α 3.2	V α 8.3 ^a	V α 11 ^b	V α 2 ^c	V α 3.2	V α 8.3	V α 11
N15 β wt	12.21 \pm 0.95 ^d	1.13 \pm 0.43	3.74 \pm 0.54	5.46 \pm 0.50	11.77 \pm 1.38	1.90 \pm 0.46	2.00 \pm 0.59	5.74 \pm 1.54
N15 β Δ FG	12.19 \pm 0.75	0.99 \pm 0.38	4.37 \pm 0.58	6.81 \pm 0.70	13.61 \pm 0.95	1.77 \pm 0.58	2.35 \pm 0.48	4.74 \pm 1.04

^a $P < 0.02$.

^b $P < 0.00005$.

^c $P < 0.005$.

^dLymph node cells from 6–12-wk-old N15 β ($n = 10$) and N15 β Δ FG ($n = 15$) mice were stained with FITC anti-CD4, CyChrome anti-CD8, biotin anti-V α mAb, followed by PE-conjugated streptavidin. Values represent the mean and SD of percentages in CD8⁺ and CD4⁺ SP lymph node T cell subsets as determined by FACS[®] analysis.

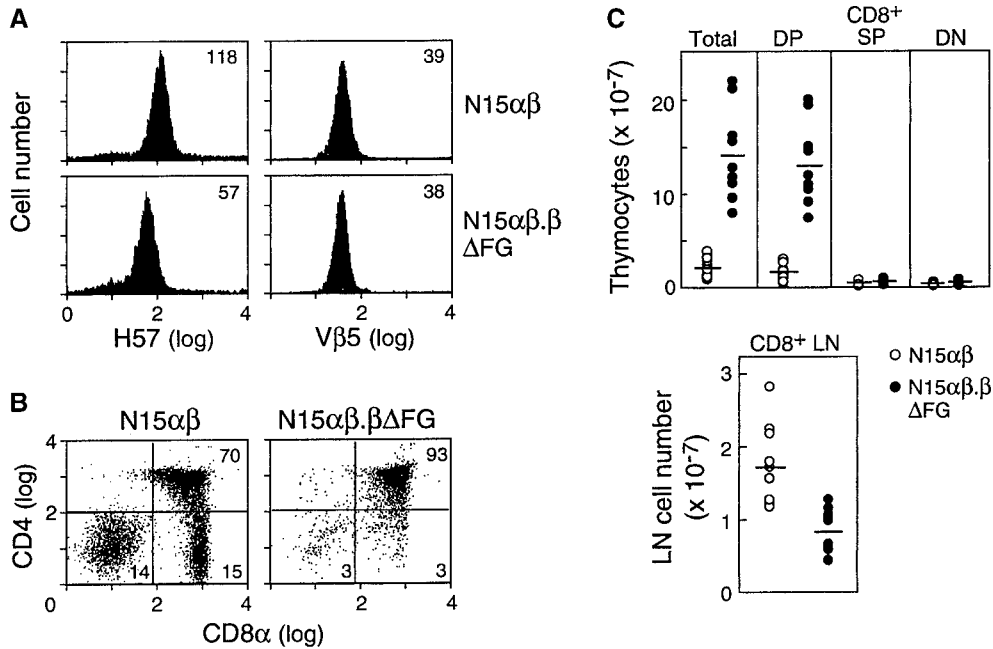


Figure 4. Quantitative and qualitative T lineage abnormalities in the N15αβ.βΔFG mice. (A) TCR expression on lymph node T cells in the N15αβ and N15αβ.βΔFG mice. Lymph node cells from 12-wk-old N15αβ and N15αβ.βΔFG mice were triple-stained with anti-CD4, anti-CD8, and anti-TCR Cβ FG loop-specific H57 or anti-Vβ5 MR9.4 mAbs. The histograms of TCR Cβ FG loop (left) and Vβ5 (right) expression on the gated CD8⁺ SP lymph node T cells are shown. The numbers represent the values of mean fluorescence intensity. (B) The CD4/CD8α double-staining profile in thymocytes is altered in N15αβ.βΔFG mice. The thymocytes from N15αβ and N15αβ.βΔFG mice at 12 wk of age were double-stained with PE-anti-CD4 and FITC-anti-CD8α. The expression of CD4 (y axis) and CD8α (x axis)

on thymocytes was detected by flow cytometry after gating on 10,000 live cells. The percentages of each subset are indicated. (C) The N15αβ.βΔFG mice show a drastic increase in the total and DP thymocytes, but a decrease in the CD8⁺ SP lymph node T cells. The numbers of total, DP, CD8⁺ SP, and DN thymocytes and CD8⁺ SP lymph node T cells were calculated by quantifying the total numbers of thymocytes and lymph node cells from 8–12-wk-old N15αβ and N15αβ.βΔFG mice and the percentages of each subset as determined by FACS[®] analysis. Open circles (N15αβ mice [*n* = 13]) and closed circles (N15αβ.βΔFG mice [*n* = 9]) represent values of individual mice and bars represent average values for a given group. The total and DP thymocyte numbers are significantly higher, and CD8⁺ SP lymph node cell number is significantly lower in the N15αβ.βΔFG mice, compared with the N15αβ mice (total and DP thymocytes, *P* < 0.0001; CD8⁺ SP lymph node cells, *P* < 0.0005).

mic T cell development in the N15 TCR αβ tg mouse system. In contrast to the difference in thymocyte number, the number of peripheral CD8⁺ SP lymph node T cells in N15αβ mice is ~2 times more than that in the N15αβ.βΔFG mice (N15αβ, 1.71 × 10⁷ cells versus N15αβ.βΔFG, 0.87 × 10⁷ cells). This difference in peripheral CD8⁺ SP T cell number may result either from faulty homeostatic expansion of T cells in the peripheral lymphoid system of N15αβ.βΔFG mice as a consequence of impaired T cell activation (37) or represent a qualitative difference in CD8⁺ SP thymocyte maturation and/or release into the peripheral T cell compartment. CFSE dye tracking of the CD8⁺ SP transgenic T cells injected into B6 RAG-2^{-/-} mice or irradiated B6 mice excludes detectable alteration in cell divisions, however (data not shown).

Decreased Thymic Negative Selection in the N15αβ.βΔFG Mice. The T cell repertoire is shaped by both negative and positive thymic selection processes acting primarily at the level of the immature DP thymocytes (for reviews, see references 35 and 38). Previous studies showed that DP thymocytes are deleted during negative selection via a caspase-dependent apoptotic mechanism (39). To investigate the basis for the dramatic increase in DP thymocytes in N15αβ.βΔFG, we evaluated the efficiency of the negative selection process in the N15αβ versus N15αβ.βΔFG animals. Apoptotic cells were enumerated through measurement of DNA content in total thymocytes using propidium iodide and flow cytometric analysis. As shown in Fig. 5 A,

the percentage of thymocytes with subdiploid DNA content, a typical feature of apoptotic cell death, is significantly higher in the N15αβ mice than in the N15αβ.βΔFG mice (N15αβ, 12.5% versus N15αβ.βΔFG, 1.1%). Consistent with this result, alterations in plasma membrane permeability, one of the earliest changes in cells undergoing apoptosis, is also altered differentially in N15αβ versus N15αβ.βΔFG mice. As shown in Fig. 5 B, about half (48.5%) of the thymocytes in the N15αβ mice are positive for annexin V staining, suggesting that a large number of DP thymocytes are being continuously deleted by apoptosis. In contrast, only a small percentage (10.0%) of thymocytes are annexin V positive in N15αβ.βΔFG mice. This finding suggests that the nature of constitutive ongoing negative selection in DP thymocytes is different between N15αβ and N15αβ.βΔFG mice.

To further probe differences in the negative selection threshold of DP thymocytes from N15αβ and N15αβ.βΔFG mice, we injected the cognate peptide VSV8 intravenously and compared the phenotypic changes in thymocytes. 18 h after intravenous injection with 0.5 μg of VSV8 peptide, N15αβ mice showed an ~90% reduction in total thymocyte number (without VSV8 injection, 2.07 × 10⁷ cells; after VSV8 injection, 0.26 × 10⁷ cells), consistent with earlier observations (25). By contrast, there is no peptide-induced change in the total thymic cellularity in N15αβ.βΔFG mice (without VSV8 injection, 10.4 × 10⁷ cells; after VSV8 injection, 10.6 × 10⁷ cells). These re-

sults suggest that deletion of the C β FG loop alters the sensitivity of immature DP thymocytes to negative selection induced by the cognate peptide in N15 TCRtg mice. Consistent with the thymic cellularity data, DNA staining with propidium iodide shows a significant increase in the percentage of subdiploid DNA-containing cells after VSV8 injection in the N15 $\alpha\beta$ (35.8%) but not in the N15 $\alpha\beta$. $\beta\Delta$ FG mice (1.3%; Fig. 5 A). Furthermore, as shown in Fig. 5 B, the N15 $\alpha\beta$ mice display a prominent increase in the annexin V⁺ thymocytes after 18 h of VSV8 (80.4%), compared with N15 $\alpha\beta$. $\beta\Delta$ FG mice (14.7%).

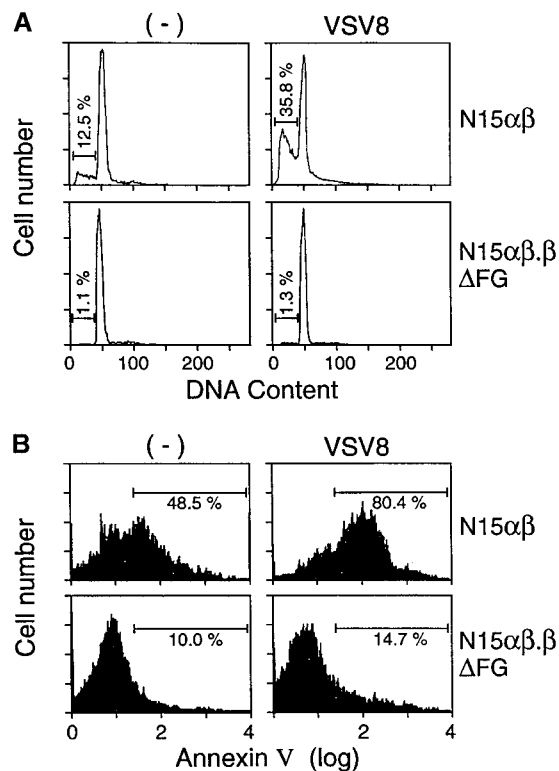


Figure 5. Decreased apoptotic cell death in thymocytes of the N15 $\alpha\beta$. $\beta\Delta$ FG mice. 7-wk-old N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG mice were injected intravenously with 0.5 μ g of VSV8 cognate peptide in 0.1 ml PBS. 18 h later, thymocytes were counted and analyzed by flow cytometry for DNA content and annexin V staining to detect apoptotic cells. Treatment of the N15 $\alpha\beta$ mouse with VSV8 resulted in a dramatic reduction of thymocyte number (without VSV8, 2.07×10^7 cells/mouse; with VSV8, 0.26×10^7 cells/mouse). By contrast, the N15 $\alpha\beta$. $\beta\Delta$ FG mouse showed no significant changes in the absolute number of thymocytes (without VSV8, 10.4×10^7 cells/mouse; with VSV8, 10.6×10^7 cells/mouse) or their subset distribution after VSV8 injection. (A) Subdiploid DNA-containing thymocytes are decreased in the N15 $\alpha\beta$. $\beta\Delta$ FG mice. After fixation and permeabilization, total thymocytes from N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG mice with or without VSV8 injection were stained with PI and assayed by flow cytometry. The histograms of DNA content on the total thymocytes are shown. The numbers represent the percentages of subdiploid DNA-containing apoptotic thymocytes. (B) Annexin V⁺ thymocytes are decreased in the N15 $\alpha\beta$. $\beta\Delta$ FG mice. Total thymocytes from N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG mice with or without VSV8 injection were stained with FITC-conjugated annexin V and assayed by flow cytometry. Dead cells were gated out using PI to reveal the proportion of live thymocytes undergoing early stages of apoptosis. The histograms of annexin V staining on the PI-negative live thymocytes are shown. The numbers represent the percentages of annexin V⁺ apoptotic thymocytes.

Collectively, these results suggest that the increase in DP thymocytes in the N15 $\alpha\beta$. $\beta\Delta$ FG mice can be explained, at least in part, by the reduced sensitivity of those DP thymocytes to undergo peptide-triggered negative selection. Although not shown, higher doses of VSV8 peptide induce apoptotic cell death in both N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG thymocytes to a similar extent, suggesting that the N15 $\alpha\beta$. $\beta\Delta$ FG mice require a larger number of pMHC-ligated TCRs to trigger negative selection. That dexamethasone-induced DNA fragmentation of thymocytes is equivalent in both mouse strains implies no general apoptotic defect in N15 $\alpha\beta$. $\beta\Delta$ FG animals (data not shown). Note that the observed apoptosis of thymocytes is not secondary to peripheral cytokine production observed in other systems (40) as shown by the rapid kinetics of the N15 TCR tg system and prior parallel FTOC analysis (25, 39).

Impaired Proliferation and Cytokine Production upon Antigen-triggered Stimulation of Naive T Cells in the N15 $\alpha\beta$. $\beta\Delta$ FG Mice. To examine the role of the C β FG loop in mature T cell function, we next tested the antigen responsiveness of N15 $\alpha\beta$. $\beta\Delta$ FG T cells by culturing lymph node T cells with VSV8 cognate peptide or variant APLs in vitro. To first address whether the C β FG loop deletion affects early phases of naive T cell activation, we assessed surface expression of the cell activation “markers,” IL-2R α (CD25) and CD69, which represents a reliable method for determining the number of T cells that are activated in culture (41, 42). The lymph node T cells from N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG mice were stimulated in vitro for 18 h with varying molar concentrations of VSV8 or the APL variants, L4 and K1 (26), using irradiated EL-4 cells as H-2K^b-bearing APCs, and then were assayed for the expression of CD25 and CD69 by flow cytometry. As shown in Fig. 6 A, VSV8 peptide was capable of activating virtually almost the same percentages of T cells at the peptide concentration of 10^{-4} – 10^{-8} M in both N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG mice. However, a very subtle difference is observed in the activation responses of the N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG lymph node T cells after stimulation with a lower concentration of VSV8 (10^{-10} M) and more convincingly with the APLs L4 and K1, as assessed by CD69 (Fig. 6 A) and CD25 (data not shown) expression. At peptide concentrations of 10^{-5} – 10^{-7} M for L4 and 10^{-4} – 10^{-5} M for K1, the percentages of activated T cells from N15 $\alpha\beta$ mice are slightly higher than those from N15 $\alpha\beta$. $\beta\Delta$ FG mice; the dose–response curve from N15 $\alpha\beta$. $\beta\Delta$ FG lymph node cells after treatment with L4 or K1 peptide is shifted to the right by a factor of ~ 10 . These findings indicate that in the mature lymphoid compartment, the early T cell activation by weak agonist ligation is modestly affected by the C β FG loop deletion as measured by this assay.

We next examined and compared Ag-induced IFN- γ production and proliferative responses of unprimed naive lymph node T cells from N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG mice. As shown in Fig. 6 B, when incubated for 48 h with VSV8 prepulsed EL-4 cells, both the N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG lymph node T cells secrete IFN- γ into the culture supernatant at each peptide concentration tested (10^{-5} – 10^{-9} M).

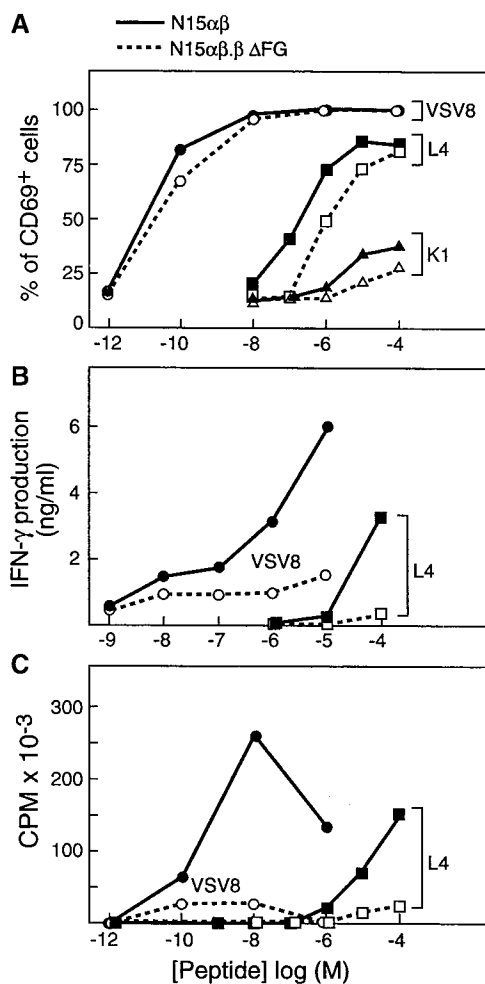


Figure 6. Impaired IFN- γ production and proliferation upon Ag-triggered stimulation of naive T cells in the N15 $\alpha\beta$. $\beta\Delta$ FG mice. (A) Lymph node T cells (1×10^5 cells) from 8–11-wk-old N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG mice were stimulated in vitro with varying molar concentrations of VSV8, L4, or K1 peptide, using irradiated EL-4 cells (2×10^4 cells) as H-2K^b-bearing APCs. After 18 h of culture, cells were stained with Cychrome-anti-CD8, PE-anti-CD25, and biotin-anti-CD69 followed by FITC-streptavidin. Gated CD8⁺ cells were analyzed and the percentages of CD25- or CD69-positive cells were determined by flow cytometry. The data on CD69 expression only are shown but staining with anti-CD25 mAb showed similar results. (B) Lymph node T cells (1×10^5 cells) from 8–11-wk-old N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG mice were incubated with the indicated concentrations of VSV8 and L4 and irradiated K^b-bearing EL-4 cells (2×10^4 cells). 48 h later, supernatants were assayed for IFN- γ production. Mean of duplicate cultures is shown. (C) Lymph node T cells (1×10^5 cells) from 8–11-wk-old N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG mice were incubated in vitro with the indicated concentrations of VSV8 and L4, using irradiated EL-4 cells (2×10^4 cells) as H-2K^b-bearing APC. 48 h later, [³H]Tdr was pulsed for 18 h and T cell proliferation was judged by [³H]Tdr incorporation. Mean of duplicate cultures is shown. Results are representative of four independent experiments.

However, the values of IFN- γ secreted from N15 $\alpha\beta$ lymph node T cells are higher than those from N15 $\alpha\beta$. $\beta\Delta$ FG T cells at 10^{-5} – 10^{-8} M VSV8. In fact, compared with N15 $\alpha\beta$ T cells, the dose–response curve from N15 $\alpha\beta$. $\beta\Delta$ FG lymph node T cells is shifted to the right by a factor of 100–1,000. Furthermore, when incubated with

EL-4 cells prepulsed with the weak agonist L4, the N15 $\alpha\beta$ T cells can produce readily detectable amounts of IFN- γ at a peptide concentration of 10^{-4} M. In contrast, N15 $\alpha\beta$. $\beta\Delta$ FG lymph node T cells cannot induce IFN- γ production after L4 stimulation at any peptide concentration tested (Fig. 6 B). Similarly, as shown in Fig. 6 C, significant differences are also observed in the IL-2–dependent proliferative responses of the N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG lymph node T cells after stimulation with antigenic peptides as assessed by [³H]thymidine incorporation. When N15 $\alpha\beta$ and N15 $\alpha\beta$. $\beta\Delta$ FG lymph node T cells are cocultured with irradiated EL-4 cells prepulsed with VSV8 for 48 h, the amounts of incorporated [³H]thymidine in N15 $\alpha\beta$. $\beta\Delta$ FG lymph node T cells are clearly less than those in N15 $\alpha\beta$ T cells at peptide concentrations from 10^{-6} to 10^{-10} M (Fig. 6 C). In addition, compared with N15 $\alpha\beta$ T cells, N15 $\alpha\beta$. $\beta\Delta$ FG lymph node T cells show significantly decreased proliferative responses to L4 stimulation at peptide concentrations from 10^{-4} to 10^{-6} M, with the dose–response curves from N15 $\alpha\beta$. $\beta\Delta$ FG lymph node T cells shifted to the right by a factor of 10–100 (Fig. 6 C). Collectively, these results clearly demonstrate that responsiveness to antigenic peptides is impaired in the N15 $\alpha\beta$. $\beta\Delta$ FG T cells as determined by peptide antigen-specific induction of proliferation and cytokine production.

Discussion

In this study, we generated a TCR β chain mutant (N15 $\beta\Delta$ FG) in which the entire C β FG loop was deleted. The N15 $\beta\Delta$ FG construct or the normal N15 β wt cDNA was then introduced as a transgene into mice and differentiation/functional analysis of thymocytes conducted on the H-2^b background. While N15 $\beta\Delta$ FG facilitated pre-TCR activity (Fig. 3, and Table I), our analysis reveals a distinct phenotype with respect to effects of the $\beta\Delta$ FG mutation on further thymocyte maturation. N15 $\beta\Delta$ FG tg mice show a statistically significant increase in CD8⁺ SP thymocytes and lymph node T cells (Table I) in association with alteration of endogenous V α repertoires (Table II). These results imply that, in balance, enhanced positive selection is favored over negative selection. Given the potential confounding effects of repertoire alteration, we subsequently restricted V α expression to that of the N15 α subunit on a RAG-2^{-/-} background in a second set of animals. Thus, N15 $\alpha\beta$ TCR tg RAG-2^{-/-} and N15 $\beta\Delta$ FG tg RAG-2^{-/-} mice were crossed to generate N15 $\alpha\beta$ RAG-2^{-/-} and N15 $\alpha\beta$. $\beta\Delta$ FG RAG-2^{-/-} littermates. N15 $\alpha\beta$. $\beta\Delta$ FG tg RAG-2^{-/-} mice were striking with regard to an 8–10-fold increase in thymocyte cellularity as a consequence of an increase in DP thymocytes without alteration in cell numbers of DN or CD8⁺ SP compartments (Fig. 4). This increase was due to reduced negative selection with decreased hypodiploid cellular DNA content constitutively as well as following in vivo VSV8 cognate peptide injection in N15 $\alpha\beta$. $\beta\Delta$ FG tg RAG-2^{-/-} compared with N15 $\alpha\beta$ tg RAG-2^{-/-} H-2^b mice (Fig. 5). Annexin V staining further confirmed the re-

duced apoptosis in N15 $\alpha\beta$. $\beta\Delta$ FG RAG-2^{-/-} versus N15 $\alpha\beta$ RAG-2^{-/-} mice.

At first glance, these data may appear in conflict with earlier studies by Degermann et al. (43), whose findings indicated that a TCR β chain lacking the identical solvent-exposed C β FG loop supported normal $\alpha\beta$ T cell development in tg mice as well as normal function in cell transfectants. Their β chain was derived from the 14.3.d TCR specific for HA₁₁₀₋₁₁₉ PR8 influenza hemagglutinin peptide complexed with I-E^d and comprised of V β 8.2 and J β 2.1 segments, unlike the N15 $\beta\Delta$ FG chain derived from the VSV8/K^b-specific N15 CD8⁺ SP CTL encoded by V β 5.2 J β 2.6 segments. In 14.3.d $\beta\Delta$ FG tg mice, superantigen-induced proliferation, anti-H-2^d CTL effector function, or helper T cell responses for anti-hapten antibody production were essentially normal. However, given that in the 14.3.d $\beta\Delta$ FG tg mice, endogenous TCR α subunits pair with the introduced β transgene product, the biologic system can readily overcome any deficiency mediated by the $\beta\Delta$ FG deletion. Hence, polyclonal T cell responses are not a sensitive indicator of the functional consequences of the C β FG loop deletion. Consistent with this notion, additional studies by the same authors in 14.3.d $\beta\Delta$ FG mice show that $\alpha\beta$ T cells coexpressing NK1.1 are severely depleted (44). This is particularly noteworthy, as ~80% of NK1.1 T cells express the invariant (V α 14-J α 281) segment required for their development (45). Thus, even in the RAG-2^{+/+} background, a near single TCR specificity is generated with regard to NK1.1 T cells, permitting the effects of the C β FG loop deletion to be recognized. Our results, in conjunction with those of Degermann, imply that the C β FG loop is functionally significant for immune recognition-based signaling by both conventional $\alpha\beta$ T cells and NK1.1 T cells of pMHC and CD1 specificities, respectively.

Prior mAb epitope mapping analysis showed that one of the two CD3 ϵ subunits within a TCR complex is in close proximity to the TCR C β chain FG loop whereas the second CD3 ϵ subunit lies elsewhere. Crystallographic analysis of the N15 TCR $\alpha\beta$ heterodimer reveals a striking asymmetry within the constant domain module such that virtually half of the C β domain's ABED surface is exposed, yet uninvolved in C α domain contacts. This asymmetry creates a cavity beneath the β chain with the C β ABED surface as a ceiling and the plasma membrane as the floor. The two protruding C α loops (CD loop and EF loop) plus one C α (N185) and two C β (N121, N186) glycans form one side-wall while the C β FG loop and one C β (N236) glycan form the second wall of this cave. In structural terms, the role of this extensive FG loop is to further rigidify the interface between V β and C β domains, thereby limiting the lateral movement of V β . In addition, careful analysis of the C β FG loop in several TCR structures (22, 28, 30, 46) reveals conservation of the three-dimensional structure of the loop, the adjacent C β domain and the cave of both pMHCI- and pMHCII-restricted TCRs in human (29, 30) and mouse (22, 28). Such conservation implies an important role for this structural element on both CD4⁺ and CD8⁺ SP T cells. By forming a side wall of the cavity into

which a CD3 ϵ subunit of one CD3 heterodimer can insert, the C β FG loop presumably stabilizes the interaction between CD3 ϵ and the T $\alpha\beta$ heterodimer. Not surprisingly, therefore, immunoprecipitation studies using the CD3 $\epsilon\gamma$ dimer-specific mAb 7D6 with N15 $\alpha\beta$ and N15 $\alpha\beta\Delta$ FG transfectants reveals a markedly weakened association of CD3 $\epsilon\gamma$ with the TCR $\alpha\beta$ heterodimer and CD3 ζ in N15 $\alpha\beta\Delta$ FG T cells (unpublished data). It is further noteworthy that the loop is negatively charged, perhaps influencing the electrostatics of the TCR surface in a manner similar to an N-linked glycan. Antedating the evolution of the FG loop in higher vertebrates, such a glycan localizes to the same region in primitive species (shark, chicken and axo; data not shown). Although the mouse has an adjacent N-linked glycan as well as the elongated FG loop in its TCR structure, both are not essential since the human TCR lacks an N-linked attachment site.

With these details in mind, we note that the TCR is a largely rigid structure: one component consists of the FG loop-“reinforced” V β -C β extracellular segment associated via an extensive (~2400 Å²) buried surface interface with the C α domain of the TCR α subunit, a second component. The C β -C α interface contains multiple rigidifying salt bridges. Hence, the V β -C β -C α domains form a single conformationally rigid module. This V β -C β -C α module in turn is noncovalently associated with the CD3 $\epsilon\gamma$ and/or CD3 $\epsilon\delta$ heterodimers such that the glycosylated CD3 γ and CD3 δ subunits extend away from the TCR heterodimer while the two nonglycosylated CD3 ϵ subunits are proximal. NMR structural studies show that CD3 ϵ and CD3 γ (and by extension CD3 $\epsilon\delta$) adopt C2-set IgSF folds, forming a stable heterodimeric complex by parallel strand pairing (G strand) and hydrophobic interaction (24). The CD3 ϵ -CD3 γ G strand pairing presumably creates a rod-like connector, permitting displacement of transmembrane helices of CD3 $\epsilon\gamma$ and CD3 $\epsilon\delta$ upon TCR triggering, leading to intracellular kinase-dependent events and resulting in T lineage activation. Removal of the C β FG loop, by eliminating one wall of the cave in the C β domain would reduce one key component of the rigidifying architecture such that signaling might be impaired. Hence the altered negative selection in DP thymocytes of the N15 $\alpha\beta$. $\beta\Delta$ FG animals and the reduced capacity of their CD8⁺ SP lymph node T cells to proliferate to cognate antigens and APL are anticipated sequelae. That polyclonal thymocytes and $\alpha\beta$ T cells in $\beta\Delta$ FG tg mice (i.e. RAG-2^{+/+}) can overcome this block, implies either that compensatory adaptations in the TCR α subunit or elsewhere readily occur or that additional protein interactions among TCR subunits (for example, within their transmembrane segments) maintain TCR-based signaling even in the absence of FG loop constraint. These possibilities are not mutually exclusive.

The prominent effect of the $\beta\Delta$ FG transgene in N15 $\alpha\beta$. $\beta\Delta$ FG RAG-2^{-/-} mice is the marked inefficiency of negative selection processes compared with that of N15 $\alpha\beta$ RAG-2^{-/-} mice. Interestingly, the C β FG loop deletion impairs negative selection without diminishing the capacity of DP thymocytes to differentiate to CD8 SP thy-

mocytes; hence, the number of CD8⁺ SP thymocytes in N15αβ.βΔFG RAG-2^{-/-} H-2^b and N15αβ RAG-2^{-/-} H-2^b littermates is identical. In general, pMHC ligands of stronger affinity are negatively selecting while those with weaker affinity for a given TCR are positively selecting (for reviews, see references 47 and 48). Given that the Cβ FG loop deletion will reduce rigidification and likely attenuate signaling from negatively selecting ligands, this result is not surprising. In contrast, positive selection must not be attenuated below a threshold required for initiating effective survival signaling. As other studies have demonstrated that positive and negative selection are under the control of different signaling pathways (for reviews, see references 38 and 49), this differential concept is entirely tenable. Nevertheless, in N15αβ.βΔFG RAG-2^{-/-} mice, the numbers of CD8⁺ SP lymph node T cells are somewhat reduced compared with N15αβ RAG-2^{-/-} mice on the same H-2 background. This observation in conjunction with increased HSA and PNA staining of CD8⁺ SP thymocytes in N15αβ.βΔFG RAG-2^{-/-} mice (data not shown) suggests that positive selection may not be entirely unaffected by the CβΔ FG mutation.

The pre-TCR consists of a TCRβ chain disulfide-bonded to the thymus-restricted pTα subunit expressed on DN thymocytes in association with CD3ε heterodimers and CD3ζ homodimers (34, 50). Although the single Ig-like ectodomain of pTα bears only 12% homology to TCR Cα, key residues participating in the formation of the rigid interface between Cα and Cβ are preserved (22). This structural conservation suggests that the heterodimeric interface of pTα-Cβ is similar to the interface of Cα-Cβ. From this conserved feature, we have inferred that a CD3ε heterodimer slots into a pre-TCR cave comparable to that created within the TCR. That a CD3ε knockout mutation disrupts pTα function (51) is consistent with this view. Furthermore, given that a CD3γ knockout also disrupts pre-TCR function (52), it is tempting to speculate that CD3εγ is the cave-associated CD3ε heterodimer. Limited glycosylation of CD3γ relative to CD3δ allows for ready modeled docking of the CD3εγ but not CD3εδ heterodimer at this cavity while still accommodating H57 mAb binding (unpublished results). Three additional facts also support the presumptive CD3εγ localization. First, it is known from chain association studies that CD3εδ pairs with TCRα whereas the CD3εγ heterodimer binds to TCRβ (53). Second, these experiments show that CD3ε and TCRβ interact via their respective ectodomains whereas the associations of the CD3γ, δ, and ζ components with TCR α and β are mediated via their transmembrane regions. Third, chemical cross-linking experiments in man and mouse imply proximity of TCRβ and CD3γ (18, 54). The above results notwithstanding, verification of these predictions still requires definitive structural elucidation.

The inability of the Cβ FG loop deletion to influence the function of the pre-TCR is consistent with recent transgenic experiments in the pTα^{-/-} background: pTα constructs lacking both the ectodomain and much of the cytoplasmic tail still support pre-TCR function (55). These

data imply that pTα stabilizes the pre-TCR predominantly via transmembrane interactions, with CD3ε heterodimer association not singularly dependent on the cave for pre-TCR function. Further support for this notion comes from three other observations: (a) mice transgenic for a TCRβ chain lacking the Vβ domain support DN to DP transition (56); (b) pTα and TCRβ chains lacking their respective Ig-like domains still facilitate thymic differentiation in RAG-deficient mice (57); and (c) mutational analysis of the two tyrosines in the TCRβ transmembrane region show that even conservative tyrosine to phenylalanine changes disrupt thymocyte survival, proliferation, and differentiation (58), further underscoring the importance of the transmembrane segment for pre-TCR function. Alterations within the TCR ectodomains are less well tolerated with regard to αβTCR function, significantly impinging on TCR activity. For example, Cβ FG loop deletion reduces the efficiency of negative selection in N15αβ.βΔFG RAG-2^{-/-} mice. Mutations of the TCRα chain connecting peptide profoundly affect positive selection. The latter occurs through attenuation of ERK activation and a reduction of activated lck, Zap-70, phospho-CD3ζ, and phospho-LAT within lipid rafts (59). A similar phenotype with equivalent biochemical signaling aberrations is observed in CD3δ^{-/-} mice (60). If CD3εγ maps to the TCRβ side and CD3εδ to the TCRα side of the αβ TCR complex, these findings can be rationalized.

In N15αβ.βΔFG RAG-2^{-/-} H-2^b mice, the level of N15 TCR as judged by both MR9.4 (anti-Vβ5) or R53 (anti-Vβ clonotype) is virtually identical to that found on T cells of N15αβ RAG-2^{-/-} H-2^b mice. However, in the former case, ~50% of TCRs are wild-type and 50% incorporate the mutant βΔFG subunit. Yet mature CD8⁺ SP T cell function is significantly reduced in N15αβ.βΔFG RAG-2^{-/-} mice. Thus, despite half of the TCRs being wild-type in structure, functional activation of these T cells vis-a-vis cytokine production or proliferation is dramatically reduced. In contrast, tetracycline regulatable TCR transgene expression shows that cells expressing as few as 1/20 of the normal levels of TCRs preserve their functional activity (61). Thus, paradoxically, the presence of TCRs with the βΔFG subunit is more detrimental to TCR-triggered activation than the complete absence of wild-type TCRs. Although the basis for this disparity remains to be elucidated, N15αβ.βΔFG TCRs may dysregulate wt TCRs by activating inhibitory pathways (phosphatases or others) in the absence of productive stimulation.

These data indicate an important functional role for the Cβ FG loop in both thymic negative selection and mature T cell activation, T lineage activities requiring threshold stimulation to achieve a stringent set point. On the other hand, positive selection is only modestly altered. In view of evidence that positive selection requirements vis-a-vis pMHC interaction are less arduous with regard to affinity and avidity than those for negative selection (47, 48), this result is perhaps not surprising. Thus, attendant reduction in signaling strength through TCRs incorporating mutant βΔFG subunits may still be sufficient to achieve the more

“relaxed” set point threshold for successful positive selection. Apparently, structural features of the $\alpha\beta$ TCR have evolved to facilitate the “multi-tasking” requirements of TCR signaling function during development and mature T cell function. In sum, TCR β -linked transduction events are more critically involved in negative selection while it appears that TCR α -linked transduction processes are key for positive selection.

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