Supplementary Information

MATERIALS AND METHODS

Generation of CD3 multicistronic vectors

CD3 multicistronic constructs were generated as described previously (Holst et al., 2006a; Holst et al., 2006b; Holst et al., 2008). Briefly, 2A peptide-linked CD3 constructs were generated by recombinant PCR and cloned into pMIG, an MSCV-based retroviral vector containing an IRES-GFP cassette. The lysines on 2A peptides were mutated to arginines without altering ‘self-cleaving’ capability (data not shown). The lysines on the cytoplasmic domain of each CD3 chain were mutated to arginines by recombinant PCR and subsequent CD3 mutant constructs were then generated by appropriate subcloning.

Generation of retroviral producer cells

Retroviral producer cell lines were generated as described previously (Holst et al., 2006a; Holst et al., 2006b; Holst et al., 2008). Briefly, HEK-293T cells were transiently transfected with CD3 multicistronic vectors (4 μg), together with packaging and envelope vectors using TransIT LT1 transfection reagent (Mirus). The supernatant containing virus was collected and used to transduce GP+E86 cells in the presence of polybrene (6 μg/ml) every 12 h for 3-4 days until a viral titer greater than 10^5/ml after 24 h was obtained.

Flow cytometric analysis, intracellular staining and cell sorting

Thymocytes were purified by FACS using mAbs against CD4 and CD8, with or without CD69 mAb (BD-PharMingen). Splenic T cells and B cells from C57BL/6J mice were purified by FACS using TCRβ and B220 mAbs. Peripheral T cells from the spleens and lymph nodes of C57BL/6J or retrogenic mice
were purified by negative MACS (AutoMACS; Miltenyi Biotec). Following red blood cell lysis with Gey’s solution, cells were stained with biotinylated mAbs against Mac1, Ter119, Gr1, B220, NK1.1 (BD-PharMingen), followed by streptavidin coupled magnetic beads. For flow cytometric analysis, mAbs against the following molecules were used: CD4 (RM4-5), CD5 (53-7.3), CD8 (53-6.7), CD25 (PC61), CD69 (H1.2F3), B220 (RA3-6B2), TCRβ (H57), and Foxp3 (FJK-16s) (All BD-PharMingen). Flow cytometric analysis was performed using a FACSCalibur (Becton Dickinson).

**CD3ε crosslinking and intracellular staining**

Bulk thymocytes were plated at 2 × 10^6 cells per well in 96 well flat bottom plate and rested for 1 hour at 37°C in 0.5% FCS RPMI. After resting, MEK inhibitor was added at a final concentration of 10μM U0126 (Sigma) in 0.1% DMSO to medium or thymocytes were treated with 0.1% DMSO only (Vehicle). Cells were fixed in 4% final Formaldehyde (Polysciences, Inc.) and permeabilized in 95% final ice cold Methanol. Permeabilized cells were stained with a pERK Ab (P-p44/42 MAPK, Cell Signaling) and detected with Alexa-647–conjugated Ab against rabbit IgG.

**Lipid bilayers and microscopic analyses**

Lipid bilayers were constructed essentially as described (Sumen et al., 2004), with some modifications (Kaizuka et al., 2007). Briefly, liposomes comprised of 90% dioleoylphosphocholine, 10% DOGS (1,2-dioleoyl-SN-Glycero-3-{[N(5-amino-1-carboxypentyl) iminodiacetic acid]succinyl}), and 0.2% biotin-CAP-PE were deposited on glass coverslips cleaned with piranha solution (50:50 mixture of 30% H₂O₂ and 96% H₂SO₄). Alexa-555-conjugated streptavidin and biotinylated TCRβ mAb were sequentially loaded onto the bilayer, while unlabeled poly-his-tagged ICAM1 produced in a baculovirus system was also loaded to facilitate T cell adhesion. Sorted DP thymocytes were added to the bilayer in RPMI medium containing HEPES and 2% FBS. Cells were fixed with 4% formaldehyde following 15 min
stimulation and immediately imaged using a Zeiss 200M spinning disk confocal microscope and analyzed using Slidebook software.

**RNA, cDNA and quantitative real-time PCR**

RNA from sorted thymocytes, T cells or B cells was isolated using the Qiagen microRNA extraction kit following the manufacturer’s instructions. RNA was quantitated spectrophotometrically and cDNA was reverse transcribed using the cDNA archival kit (Applied Biosystems) following the manufacturer’s guidelines. TaqMan primers and probes were designed with PrimerExpress software and synthesized in the St. Jude Hartwell Center for Biotechnology and Bioinformatics:

**CD3\(\delta\) 5’ primer, AAAGGTGGTGTCTTCTGTGCAA;**

**CD3\(\delta\) 3’ primer, CCCGAGTCTAGCTCCACACAGT;**

**CD3\(\delta\) probe, FAM-TCCATTACCGAATGTGCCA.**

**CD3\(\gamma\) 5’ primer, GCGGGACAGGATGGAGTTC;**

**CD3\(\gamma\) 3’ primer, TTCATTTTGCAACAGAGTCTGCTT;**

**CD3\(\gamma\) probe, FAM-CCAGTCAAGAGCTTCAGA.**

**CD3\(\epsilon\) 5’ primer, CAGCCTCAAATAAAAAACACGTACTTG;**

**CD3\(\epsilon\) 3’ primer, TCAGGTCCACCTCCACACAGT;**

**CD3\(\epsilon\) probe: FAM-CTGAAAGCTCGAGTGTGTGA.**

**CD3\(\zeta\) 5’ primer, GAGGAGGAACCCCCAGGAA;**

**CD3\(\zeta\) 3’ primer: TGTAGGCTTCTGCCATCTTGTGTC;**
For quantitative analysis, an ABI Prism 7900 Sequence Detection System was used for real-time RT-PCR. Expression was normalized relative to the expression of endogenous β actin.

**Transient transfection of HEK-293T cells**

Transient transfection of HEK-293T cells was performed as previously described (Liu et al., 2000; Szymczak et al., 2004) with some modifications. HEK-293T cells were incubated in 6 well plates at 2 × 10^5/plate overnight at 37°C. TCRαβ (2A-linked) plasmid (1 μg) and indicated CD3 plasmid (1 μg) with or without a construct containing WT or a dominant negative dynamin (1 μg) were transfected to HEK-293T cells using 6 μl TransIT LT1 transfection reagent (Mirus). Cells were harvested 40 h after transfection and stained with CD3ε and TCRβ mAbs for flow cytometry analysis.

**Thymic Organ Culture**

Thymic organ culture was performed as previously described (D'Oro et al., 1997; Hogquist, 2001; Holst et al., 2008) with some modifications. The culture medium is RPMI (Gibco) buffer supplemented with 10% FBS, 50 μM 2-mercatoethanol, 2 mM L-glutamine, 50 U/mL penicillin, 50 mg/mL streptomycin, 50 mg/mL gentamycin sulfate, and 5 mM HEPES. Thymic lobes from newborn mice (P1) were cultured on a culture plate insert (Millipore) floating in a prepared well for 20 h at 37°C. Cultures were carried out in 1 ml of medium containing indicated reagents or vehicle control in 6-well plates. The plate was placed in a tupperware dish which has a water soaked paper towel in its bottom. Individual thymic lobes were harvested and single-cell suspensions were analyzed for surface TCR expression by flow cytometry. Following reagents were used in thymic organ culture: MG132, Lactocytin, NH₄Cl (Sigma),
Concanamycin A (Sigma), MiTMAB (EMD Chemicals Inc.), PP2 (Calbiochem), Syk inhibitor (Calbiochem), and PP1 (Invitrogen).

**Histological examination**

Histological examination was performed as previously described (Holst et al., 2008). Hematoxylin and eosin stained, formalin-fixed (10% neutral buffered), paraffin-embedded tissue sections (4 μm) of liver, lung and intestinal tract were evaluated in a blinded fashion by an experienced veterinary pathologist (Kelli Boyd, Department of Pathology, St. Jude Children’s Research Hospital). Inflammation in each organ was assigned a score of 0 to 3 based on a semi-quantitative grading scheme established for this model (Supplementary Table 1).
REFERENCES

D'Oro U, Vacchio MS, Weissman AM, Ashwell JD (1997) Activation of the Lck tyrosine kinase targets cell surface T cell antigen receptors for lysosomal degradation. *Immunity* 7: 619-628


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Figure 4

A

Gating

DP thymocytes

SP thymocytes

% of Max

TCRβ

Neg. Control  Medium only  Dynamin Inhibitor II

B

Surface TCR expression (% of medium only)

Medium only  Dynamin Inhibitor II

DP  CD4 SP

C

Surface TCR expression (% of medium only)

Medium only  10 μM MG132  20 μM Lacto-cystin  500 nM Concana-mycin A

% of Max
Figure 5

A

| CD3δ | HETGRPSGAEEVQALLKEQGLYQPLREDTQYSRLGNWPRNKKSB |
| CD3γ | QDGVRQSRASTDKQTLLQNEQLYQPLKREDYQYSHLQGNQLRKKE |
| CD3ε | SKNRKAKAXPVRCTGAGSRFRQNLQERPPVPNPDYEPIRKQQRDLGSLNQRAV |
| CD3ζ | RAKFSRSEQTALNDPNQLYNELNLGRREEYDVEKKRARDPEMGKDQRRNRPOEGVYNALQDFDSEAAYSEIGTKGERGRKGHDGLYQGLSTAKDQYDALHMQTLAPR |

B

Activated by anti-CD3ε

CD3WT CD3KR

Activated by anti-CD3ζ

IP: α-CD3ζ Blot: α-Ubiquitin

IP: α-CD3ζ Blot: α-CD3ζ

C

Gating DP thymocytes SP thymocytes

Surface TCR level on DP thymocytes (Normalized by CD3WT)

| Surface TCR level on DP thymocytes (%) of Max | CD3δ ε | M M | M M | M M |
| Surface TCR level on DP thymocytes (%) of Max | CD3γ | M M | γ M |
| Surface TCR level on DP thymocytes (%) of Max | CD3ε | ε M |
| Surface TCR level on DP thymocytes (%) of Max | CD3ζ | M M |
| Surface TCR level on DP thymocytes (%) of Max | # Lysine | 37 0 3 4 12 18 |
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**Figure 7**

A. Surface TCR level on DP thymocytes (normalized by CD3<sup>WT</sup>)

B. GFP+ Thymocytes Number (x10<sup>7</sup>)

C. % GFP+ Thymocytes

D. Gated on GFP+ thymocytes

E. Gating on GFP+ CD4+ Splenocytes

F. CD4<sup>+</sup>foxp3% of Splenic CD4<sup>+</sup> T cells

G. Thymic Foxp3+ cell Number (10<sup>4</sup>)

CD3<sup>WT</sup>, CD3<sup>KR</sup>, CD3<sup>KR-monoUb</sup>

CD4+CD8+, CD4+CD8−, CD4−CD8−, CD4-CD8+
Figure 8

(A) Flow cytometry plots showing CD8 Vβ8.3 expression on GFP+ blood cells gated on CD3WT, CD3KR, and CD3KR-monoUb in males and females. The dot plots display cell counts and the histograms show the mean fluorescence intensity (MFI) for CD5 expression on DP thymocytes.

(B) Bar graph depicting thymic cellularity (x10^8) in females and males for CD3WT, CD3KR, and CD3KR-monoUb.

(C) Bar graph showing CD5 expression on DP thymocytes (MFI) for CD3WT, CD3KR, and CD3KR-monoUb in females and males.
<table>
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<th>Score</th>
<th>Lung and Liver</th>
<th>Small and Large Intestine</th>
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<tr>
<td>0</td>
<td>Rare inflammatory cells in the perivascular space</td>
<td>Rare inflammatory cells in the perivascular space</td>
</tr>
<tr>
<td>(Negative)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Well delineated cuffs of inflammatory cells around vessels</td>
<td>Lymphocytes and plasma cells with few neutrophils and macrophages in the lamina propria, without necrosis of crypts</td>
</tr>
<tr>
<td>(Mild)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Perivascular cuffing and vasculitis</td>
<td>Lamina propria inflammation multifocal crypt necrosis, multifocal enterocyte apoptosis, moderate dilation of lymphatics</td>
</tr>
<tr>
<td>(Moderate)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Perivascular cuffing, vasculitis and parenchymal necrosis</td>
<td>Mucosal ulceration, inflammatory infiltrate in lamina propria and mucosa, wide spread crypt necrosis, transmural edema</td>
</tr>
<tr>
<td>(Severe)</td>
<td></td>
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**Supplementary Table 1 | Pathology Score Description**

The table describes the scale and the types of inflammation observed in those mice shown in Supplementary Fig. 7.
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Figure S1

A) Thymi were isolated from 6 week old C57BL/6J mice. Surface expression of the TCR:CD3 complex on DN, DP, SP and total thymocytes was measured by staining with a CD3ε antibody and was analyzed by flow cytometry. (B) Thymocytes were stained with antibodies to CD4 and CD8, and DP and SP thymocytes were purified by FACS. Total, DP and SP thymocytes were lysed and an equal amount of total protein was loaded in each sample. The protein levels of CD3δ, CD3γ, CD3ε, CD3ζ and tubulin (loading control) were measured by SDS-PAGE and Western blot. (C) DP and SP thymocytes were purified by FACS as above. DN thymocytes were purified using Lin- (CD4-CD8-B220-Mac1-Ter119-panNK-TCRβ-TCRδγ) and Thy1.2* antibodies. B cells were purified from splenocytes on B220. RNA was extracted and reverse transcribed to cDNA. By using β-actin gene as an internal control, the CD3δ, CD3γ, CD3ε and CD3ζ mRNA levels were measured by qPCR. All data were normalized by the CD3 mRNA levels in DP thymocytes.
Figure S2 | SLAP is involved in the regulation of TCR expression in DP thymocytes

(A) Thymi were isolated from either C57BL/6J mice or Slap<sup>−/−</sup> mice. Surface expression of the TCR:CD3 complex was measured by staining anti-TCRβ and analyzed by flow cytometry. (B) Thymocytes were stained with antibodies to CD4 and CD8, and DP and SP thymocytes were purified by FACS. Purified thymocytes were fixed, permeabilized, and stained with a CD3ζ antibody conjugated with Alexa-647. The localization of CD3ζ (red) and DAPI stained nucleus (blue) are shown. (C) The distribution of CD3ζ in the thymocyte populations indicated is shown as a bar chart.

Figure S3 | Colocalization of CD3ζ with LAPTM5 in DP thymocytes

Thymocytes were stained with antibodies to CD4 and CD8. DP and SP thymocytes were purified by FACS. Thymocytes were fixed, permeabilized, and stained with a CD3ζ antibody conjugated with Alexa-647 and a LAPMT5 antibody, which was subsequently detected with an Alexa-488-tagged rabbit IgG secondary antibody. Nuclei were stained with DAPI. The scale bar is 5 μM.
Figure S4 | Constitutive multi-mono ubiquitylation of CD3ζ in DP thymocytes.

DP (240 × 10⁶) and SP (30 × 10⁶) thymocytes were purified by FACS and lysed in 1 ml lysis buffer. The lysates were immunoprecipitated with a CD3ζ antibody (551-ζ), separated by SDS-PAGE and Western blots probed with a mAb to either polyUb (FK1, left), K63Ub (HWA4C4, middle), or Ub (P4D1, right). Blots were stripped and re-probed with anti-CD3ζ (H146) to show that an equal amount of CD3ζ was analyzed in each sample.

Figure S5 | Comparable retroviral transduction efficiency

(A) Schematic representation of the TCRαβ:CD3γδεζ complex. (B) Schematic representation of the CD3-2A constructs in the MSCV-ires-GFP (pMIG) vector as shown previously. The lysines (K) in 2A peptide are highlighted in red and are mutated to arginines to prevent ubiquitylation. (C) Retrogenic mice were generated by retroviral-mediated stem cell gene transfer using CD3εζ–/– as bone marrow donor mice and sublethally irradiated Rag1–/– as recipient mice. Thymi were isolated 5-8 weeks post bone marrow transplant. The GFP mean fluorescence intensity in GFP+ thymocytes was measured by flow cytometry. Bars represent the mean ± s.e.m.
Figure S6 | No other detectable sumoylation, neddylation, or methylation occurring on CD3ζ

Splenic T cells were purified by MACS (A-C, 30 × 10^6 cells alone or activated by crosslinking with CD3ζ antibodies for 2 min) and SP thymocytes (D-F, 30 × 10^6) or DP thymocytes (D-F, 240 × 10^6) were purified by FACS, and lysed in 1 ml lysis buffer. The lysates were immunoprecipitated with a CD3ζ antibody (551-ζ), separated by SDS-PAGE and Western blots probed with Sumo-1 (A and D), Nedd8 (B and E), or pan-Methyl (C and F) antibodies. Purified Sumo-1, Nedd8 and Histone proteins were loaded as positive controls. The predicted size of modified CD3ζ is indicated by arrows. Blots were stripped and re-probed with anti-CD3ζ (H146) to show that an equal amount of CD3ζ was analyzed in each sample.
Figure S7 | Attaching mono-ubiquitin induces TCR downregulation
(A) HEK-293T cells were transiently transfected with TCRαβ (2A-linked) and CD3δγεζ (wild type or mutants as indicated). Transfected cells were stained with a TCRβ antibody, and analyzed by flow cytometry. Bar chart shows the mean ± s.e.m. of the TCRβ mean fluorescence intensity for the GFP+ gated cell population. (B) HEK-293T cells were transfected as for (A), along with either wild type or a dominant negative dynamin. Transfected cells were stained with antibodies to CD3ε and TCRβ, and analyzed by flow cytometry. Dot plots show the level of TCRβ and CD3ε surface expression for the GFP+ gated cell population.

Figure S8 | Activation induced phosphorylation of ERK
Thymocytes from Sla−/− and littermate controls were activated by crosslinking using anti-CD3ε in the presence (+U0126) or absence (+Veh.) of the MEK inhibitor U0126. Thymocytes were then fixed at the indicated time, permeabilized, and stained with antibodies to CD4, CD8 and pERK. (A) CD4+CD8+ DP thymocytes were gated and pERK expression measured. (B) Representative flow cytometry dot plots are shown 10 minutes post-stimulation. Statistical significance was determined using an unpaired t test in Prism software.
**Figure S9 & S10**

**Figure S9 | Ratio of SP thymocyte number to DP thymocytes number**
Retrogenic mice were generated by transducing CD3εζ bone marrow (A) or CD3εζ⁺/-Rag1⁻⁻ bone marrow (B), transplanting into sub-lethally irradiated Rag1⁻⁻ recipients, and analyzing 6-8 weeks post-transfer. Thymocytes were stained with antibodies to CD4, CD8, and TCRβ, and analyzed by flow cytometry. (A) The ratio of absolute numbers of SP to numbers of TCRβ⁺DP thymocytes is shown. (B) The ratio of absolute numbers of CD8 SP to numbers of TCRβ⁺ DP thymocytes in female MataHari mice is shown.

**Figure S10 | Pathology of CD3KR-monoUb mice**
Retrogenic mice were generated reconstituting sub-lethally irradiated Rag1⁻⁻ recipients with transduced CD3εζ⁻⁻ bone marrow. Mice were sacrificed 8-12 weeks post bone marrow transfer, and the lung, liver, small and large intestine were removed for histological examination. The pathologist was blinded, and scored the level of inflammation as described in Supplementary Table 1. (A) Hematoxylin and eosin staining of formalin-fixed, paraffin-embedded sections of organs are shown. (B) Bar chart shows average pathology score of each mice group.
Figure S11 | Generation of DP thymocytes is blocked in absence of TCRβ.
Retrogenic mice were generated by transducing CD3εζ bone marrow (upper panel) or CD3εζ–/–Rag1–/– bone marrow (lower panel), transplanting into sub-lethally irradiated Rag1–/– recipients, and analyzing 6-8 weeks post-transfer. Thymocytes were stained with antibodies to CD4 and CD8, and analyzed by flow cytometry.

Figure S12 | Thymus size of MataHari retrogenic mice
Retrogenic mice were generated by transducing CD3εζ–/–Rag1–/– bone marrow, transplanting into sub-lethally irradiated Rag1–/– recipients, and analyzing 5-7 weeks post-transfer. Thymi were isolated and photographed to compare size of thymus.
Figure S13| Tonic ubiquitylation and T cell development
Schematic representation of the model: different TCR levels in the indicated retrogenic mice (CD3^{KR}>CD3^{WT}>CD3^{KR monoUb}) alter the signal strength threshold for negative/positive selection and generation of natural T_{reg}. 

Figure S13