Negative Regulation of Zap70 by Lck Forms the Mechanistic Basis of Differential Expression in CD4 and CD8 T Cells

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Lck and Zap70, two non-receptor tyrosine kinases, play a crucial role in the regulation of membrane proximal TCR signaling critical for thymic selection, CD4/CD8 lineage choice and mature T cell function. Signal initiation upon TCR/CD3 and peptide/MHC interaction induces Lck-mediated phosphorylation of CD3 ITAMs. This is necessary for Zap70 recruitment and its phosphorylation by Lck leading to full Zap70 activation. In its native state Zap70 maintains a closed conformation creating an auto-inhibitory loop, which is relieved by Lck-mediated phosphorylation of Y315/Y319. Zap70 is differentially expressed in thymic subsets and mature T cells with CD8 T cells expressing the highest amount compared to CD4 T cells. However, the mechanistic basis of differential Zap70 expression in thymic subsets and mature T cells is not well understood. Here, we show that Zap70 is degraded relatively faster in DP and mature CD4 T cells compared to CD8 T cells, and inversely correlated with relative level of activated Zap70. Importantly, we found that Zap70 expression is negatively regulated by Lck activity: augmented Lck activity resulting in severe diminution in total Zap70. Moreover, Lck-mediated phosphorylation of Y315/Y319 was essential for Zap70 degradation. Together, these data shed light on the underlying mechanism of Lck-mediated differential modulation of Zap70 expression in thymic subsets and mature T cells.

Keywords: TCR signaling, Lck kinase, Zap70 kinase, negative regulation, CD4/CD8 T cells

INTRODUCTION

T cells, an indispensable component of adaptive immunity, constantly patrol our body and eliminate pathogen infected or malignant cells via recognition by the T cell receptor (TCR) of antigenic peptide presented in association with the major histocompatibility complex (p/MHC) (1). TCR-induced signal transduction is essential for effective neutralization of such insults by mature T cells. TCR signaling is equally important for development of T cells in the thymus. It is implicated in maturation through the three distinct stages of thymocyte development: CD4 CD8⁻ double negative (DN), CD4⁺CD8⁻ double positive (DP) and CD4⁺ or CD8⁺ single positive (SP) thymocytes (2–4). Thymic selection of DP thymocytes ensures that thymocytes expressing high affinity TCRs are
negatively selected while those with moderate/low affinity for self-p/MHC are positively selected and develop further into MHCII-specific CD4 helper and MHC-I-specific CD8 cytotoxic T cells (5). In the periphery tonic TCR signaling induced by self-peptide/MHC play an important role in T cell homeostasis, while antigenic peptide/MHC complex induce differentiation into effector/memory T cells. Central to initiation and propagation of TCR signaling are two non-receptor tyrosine kinases, Lck, a Src family kinase, and Zap70, a Syk family kinase. The two kinases play an obligatory role in TCR signaling during thymic development and mature T cell function (6, 7). Lck is required for DN to DP transition as well as for thymic selection of DP thymocytes into SP thymocytes (2, 8–10). In contrast, Zap70 is critical for DP to SP transition although it is proposed to play some role in DN to DP transition and survival of DP thymocytes (11, 12).

Studies have shown that the cytoplasmic tails of CD4 and CD8 co-receptors serve as binding sites for Lck (13, 14). Lck binds strongly to CD4 compared to CD8 cytoplasmic tail and engagement of CD4 mediates stronger Lck activation (15). The catalytic activity of Lck is tightly regulated by phosphorylation status of inhibitory Y505 at the C-terminus and activating Y394 located in the kinase domain. Phosphorylation by Csk kinase and dephosphorylation by CD45 phosphatase of the inhibitory Y505 in Lck promotes, respectively, closed inactive and open active conformation (16–23). A signal initiating step following TCR/pMHC engagement involves Lck-mediated phosphorylation of immunoreceptor tyrosine-based activation motifs (ITAM) of TCR-associated invariant chain tails (24, 25). Zap70 binds to phospho-CD3 via two tandem SH2 domains that permits Lck-mediated sequential phosphorylation of Y315/Y319 and Y493 leading to full Zap70 activation triggering downstream signaling cascade culminating in thymocyte or mature T cell activation (26, 27). Y315/Y319 play a critical role in maintaining the auto-inhibitory closed conformation of Zap70. Lck-mediated phosphorylation of Y315/Y319 relieves this auto-inhibition critical for full activation of Zap70 essential for TCR signaling. An essential role for Zap70 kinase is evident from impaired thymic selection in Zap70-deficient mice (28, 29).

While Lck is essential for DN to DP transition and thymic selection, it also modulates CD4 versus CD8 lineage choice of positively selected thymocytes. Augmented (inhibitory Y505F mutation) and diminished (kinase dead K273R mutation) Lck activity promotes CD4 and CD8 lineage choice, respectively, supporting the signal strength/duration model of CD4/CD8 lineage choice (30–32). Interestingly, studies using a mouse model of inducible Zap70 expression showed that sustained Zap70 expression is required for CD8 lineage choice although ablating Zap70 expression after signal transduction did not alter lineage choice (33–35). In WT mice Zap70 is expressed in very low amounts in DN thymocytes and its expression rises in DP and SP thymocytes, CD8 SP thymocytes expressing it particularly strongly (35). This pattern of Zap70 expression is opposite of Syk kinase expression, another member of the Syk kinase family, and the two kinases play unique roles in pre-TCR/TCR signaling during DN to DP transition (12, 36). Compared to mature T cells, DP thymocytes express substantially lower amount of TCR/CD3 complex, which is due to its rapid degradation upon induction of Lck activity following CD4 and MHCII interaction (15, 37, 38). Interestingly, in DP thymocytes Zap70 is associated with phosphorylated CD3ζ (39). Importantly, in DP thymocytes despite significant Lck activity induced by CD4/MHCII interaction TCR-associated Zap70 was shown to be inactive as judged by anti-phosphotyrosine Western blot (39). Zap70 expression is further elevated in SP thymocytes with CD8 T cells expressing highest amounts, yet this does not appear to be due to increased transcription suggesting post-translational regulation of Zap70 although the mechanism of this regulation is not understood.

In the present investigation, we show that under steady state relatively slower degradation explains the higher Zap70 expression in CD8 compared to CD4 SP and DP thymocytes. In addition, differential Zap70 expression in thymic subsets inversely correlated with relative amount of phospho-Zap70 levels in these cells. Augmenting Lck catalytic activity in a heterologous system and T cell line showed significant decrease in Zap70 expression. In transgenic mice expressing constitutively active Lck, we observed a severe diminution in Zap70 expression but a significant increase in relative amount of phosphorylated Zap70 in all thymic subsets and mature T cells. Further experiments showed that phosphorylation of Y315/Y319 of Zap70 by Lck played a critical role in Zap70 degradation. This study provides a strong evidence that Lck activity is a negative regulator of Zap70 expression, a finding with important implications for T cell development and function.

**MATERIALS AND METHODS**

**Mice**

C57BL6 mice were bred in our animal facility. MHC-I-restricted OTI+Rag-/- (chicken ovalbumin 257-264 peptide specific) and P14+Rag-/- (lymphocytic choriomeningitis virus (LCMV) GP33, 41 peptide specific) transgenic mice were obtained from Nathalie Labrecque and Heather Melichar (Centre de Recherche Hopital Maisonneuve-Rosemont (CRHMR)). MHCII-/- mice were obtained from The Jackson Laboratory. All TCR transgenic mice expressing constitutively active LckY505F or Thpok transgene were described previously (40). Mice were genotyped by peripheral blood analysis and/or PCR of genomic DNA isolated from an ear punch. Lymphoid organs from 5–8 week old mice were harvested for all analyses. All mice were housed under specific pathogen free conditions at the CRHMR. Animal experimentation protocols were approved by the CRHMR animal care committee and performed in accordance with the Canadian Committee for Animal Care.

**Transient Transfection**

WT Lck and mutant expressing Y505F or K273R were cloned into pCCL lentiviral vector (kind gift of Jonathan Bramson, McMaster University). We also cloned Lck and mutants fused to mCherry into pCCL lentiviral vector (WT Lck-mCherry fusion kind gift of N. Gascoigne; National University of Singapore) (41).
The Lck-mCherry fusion molecule expression is driven by an EF1α promoter. The plasmid also expresses truncated human NGFR under the control of CMV promoter. Lentivirus plasmid encoding full-length mouse Zap70 was obtained from Abm (pLenti-GIII-CMV-RFP-2A-Puro; cat # LV434826). In this plasmid Zap70 and RFP reporter expression were driven by CMV and SV40 promoter, respectively. RFP coding sequence was replaced by ZsGreen coding sequence (pLenti-Zap70-ZsGreen). Control lentiviral plasmid was generated by deleting Zap70 coding sequence (pLenti-ZsGreen). Y315F/Y319F (YYFF) or Y319F mutated in pLenti-Zap70-ZsGreen were generated by overlap PCR. Mutations were confirmed by DNA sequencing. One million 293T cells were co-transfected with plasmids encoding mouse Zap70 and pCCL plasmids expressing control or Lck variants in the presence of polyethylenimine (PEI). Ninety-six hours post transfection cells were fixed, permeabilized and stained for Zap70 followed by flow cytometry analysis. Mouse Zap70 expression was determined in ZsGreen+NGFR+ cells and normalized to Zap70 expression ZsGreen+ cells transfected with control plasmid for Lck expression. Plasmid encoding human Zap70-GFP fusion molecule linked to EGFR by self-cleaving T2A peptide sequence was obtained from Johannes Huppa (Medical University of Vienna) (42) and a hZap70-GFP expressing 293T stable cell line was generated. This stable cell line was transfected with various Lck-mCherry fusion lentivirus constructs. For human Zap70-GFP fusion protein expression analysis, GFP expression in mCherry expressing or not cells was analyzed by flow cytometry and expressed as ratio of mCherry-mCherry+ cells. In some experiments Zap70-GFP expressing 293T cells were transfected in triplicate with NGFR reporter expressing control, LckWT or LckY505F plasmids and stained intracellularly with PE-conjugated anti-pY319 antibody. Ratio of MFI for pY319 over GFP in NGFR+ve and NGFR-ve cells was determined and normalized to control transfectants.

**Lentivirus Production and Transduction**

For lentivirus production 293T cells were plated in 10cm tissue culture dish and transfected with LckWT-mCherry, LckY505F-mCherry or LckK273R-mCherry lentiviral plasmid and pMD2.G and psPAX2 packaging plasmids (Addgene) in the presence of polyethylenimine. Culture supernatant containing lentiviral particles was harvested at 48 and 72 hours post-transfection and used to transduce the RLM mouse CD4 SP thymocyte cell line. Zap70 expression in transduced cells (mCherry+) was determined and normalized to control transfectants.

**Flow Cytometry**

A total of 1 x 10⁶ thymocytes or RBC-depleted spleen cells were incubated with a combination of fluorescently labeled Abs to CD4 (GLK5), CD8 (53-6.7), TCRβ (H57-957), CD5 (53-7.3), CD69 (H1.2F3), CD24 (M1/69), CD44 (IM7), CD62L (MEL-14), Vα2 (B2O.1), Vβ5 (MR9-4), and analyzed using flow cytometry (LSRFortessa X-20 or LSR II, BD Biosciences). For intracellular labeling thymocytes or spleen cells were surface stained, fixed and permeabilized using commercial kit (Ebiscience, FoxP3 fixation/permeabilization buffer) and stained for Lck, Zap70, pY319-Zap70, and pY493-Zap70. Appropriate isotype control antibody staining was included in all experiments involving intracellular staining. Antibodies were obtained from ebiscience, BioLegend, or Cell Signaling Technology. Data were analyzed using FlowJo software (Tree Star).

**Quantitative RT-PCR**

Various thymic or splenic T cell subsets were FACS purified and total RNA isolated using TRizol (Invitrogen). Complementary DNA was synthesized using a commercial kit as per manufacturer’s protocol (Bio-Rad Laboratories). Quantitative PCR (QPCR) for target gene was performed in triplicate using SYBER green dye (Bio- Rad Laboratories) or EvaGreen dye (Abcam). Amplification of housekeeping gene Hprt served as an internal control. QPCR data were analyzed by Applied Biosystems software ABI 7500 v2.0.5. Data were normalized to Hprt expression in each population. Relative expression values were calculated using ΔΔ cycle threshold method. Ratio of gene-specific values to housekeeping gene for each subset was determined and normalized to CD4 T cells for splenic T cell analysis and to DN thymocytes for thymic subset analysis. Data are presented as an average of triplicate values with standard deviation (SD). QPCR primers for Hprt, Lck and Zap70 were obtained from Integrated DNA Technologies.

**Kinetics of Zap70 Expression**

To determine the kinetics of Zap70 degradation splenic T cells were isolated using negative selection beads (Stem Cell Technology) followed by FACS sorting to obtain purified CD4 and CD8 T cells. One million CD4 and CD8 T cells were cultured in the presence of cyclohexamide, an inhibitor of protein translation, for different time points followed by lysis in Laemmeli buffer. Lysates were resolved on polyacrylamide gel under reducing condition, transferred onto PVDF membrane, blocked in 5% milk in TBST buffer (10mM Tris pH7.6, 150mM NaCl and 0.1% Tween20) for 1 hour at room temperature followed by anti-Zap70 or anti-actin antibody overnight (O/N) at 4°C. Membranes were washed four times with 0.1%TBST and further incubated with HRP coupled secondary antibody, washed four times with 0.1%TBST, and developed using chemiluminescent reagent. Bands were visualized using Azure c600 imaging system and their intensity determined using ImageJ software. Ratio of band intensity of Zap70 over actin was determined and normalized to that at 0 minute as 100%. To study the effect of activation on Zap70 degradation one million purified CD4 and CD8 T cells were cultured with biotinylated anti-CD3 antibody (10μg/ml) for 30 minutes on ice, washed with ice-cold media and stimulated by adding pre-warmed media containing streptavidin (20μg/ml) for 5, 10, 60 and 120 minutes. Cell were processed for Western blot as described above. Incubation, wash and stimulation media contained cyclohexamide. For the analysis of Zap70 degradation in thymocytes, cells were cultured in the presence of cyclohexamide, surface stained for CD4/CD8/TCR followed by Zap70 intracellular staining at 30 min, 1, 2, and 4 hr post-incubation. To account for variability in staining we determined and compared the ratio of Zap70 expression in target subset over DN thymocytes (which express low amount of
Zap70); the ratio for DN thymocytes was treated as one. To determine Zap70 ubiquitination purified T cells from OTI mice expressing or not Lck<sup>K273R</sup> transgene were lysed and anti-Zap70 immunoprecipitate was Western blotted with anti-Ubiquitin or anti-Zap70 antibody, and the ratio of anti-Ub to anti-Zap70 band intensity was compared.

**Statistical Analyses**

Statistical analyses were performed using GraphPad or Microsoft Office. Excel software. Data are displayed as a mean with SD error bar. Unpaired two-tailed Student t test was used to determine statistical significance when thymic and splenic T cell subsets from different mice. For experiments involving a comparison of T cell subsets isolated from the same mouse, a paired Student t test was used. A p value < 0.05 was considered statistically significant (*p < 0.05, **p < 0.005, ***p < 0.0005 and ****p < 0.00005).

**RESULTS**

**Zap70 Expression Is Inversely Correlated With Its Activation Status**

To dissect the mechanism of differential Zap70 expression in CD4 and CD8 T cells or thymic subsets, we first evaluated Zap70 expression in cells from wild type (WT) mice. To this end, we first surface stained splenocytes for CD4, CD8 and TCR followed by fixation, permeabilization and intracellular staining for Zap70. This highly sensitive approach allowed us to accurately evaluate Zap70 expression at the single cell level. Here, we observed a significantly higher level of Zap70 protein in mature CD8 than CD4 splenic T cells despite higher Zap70 mRNA expression in the latter subset (Figures 1A,B; Figure S1), which is in agreement with published report (35). Similar analysis of WT thymocytes showed a direct correlation between thymocyte maturation and Zap70 expression with CD8 SP and DP thymocytes expressing the highest and lowest Zap70 amount, respectively (Figure 1C) (12, 35). Further, CD4<sup>+</sup>CD8<sup>+</sup> and CD4<sup>+</sup> SP thymocytes expressed significantly more Zap70 than DP thymocytes but less than CD8 SP thymocytes. Interestingly, at the transcriptional level, higher Zap70 mRNA levels were observed in DP compared to more mature CD4<sup>+</sup>CD8<sup>+</sup> or CD4<sup>+</sup> and CD8<sup>+</sup> SP thymocytes (Figure 1D). These data suggest differential Zap70 stability in DP thymocytes and in CD4 mature thymocytes compared to CD8 thymocytes or mature T cells. To test this notion, we evaluated the kinetics of Zap70 degradation in thymocytes and mature T cells in the presence of cyclohexamide, a protein translation inhibitor. Indeed, flow cytometric analysis of Zap70 expression kinetics of WT thymocytes showed faster degradation in DP and CD4 SP thymocytes compared to CD8 SP thymocytes (Figure 1E). Similarly, anti-Zap70 Western blot analysis of mature CD4 and CD8 T cells showed faster Zap70 degradation in CD4 T cells compared to CD8 T cells; at 180 minutes Zap70 expression was reduced by 60% in CD4 T cells compared to about 35% in CD8 T cells (Figure 1F). We then determined that T cell activation resulted in accelerated Zap70 degradation with CD4 T cells showing higher degradation compared with CD8 T cells (Figure 1G) and is in agreement with a previous report (43). These data support the notion that Zap70 is relatively less stable in DP thymocytes and CD4 SP thymocytes compared to CD8 SP thymocytes or T cells.

A previous report showed, by Western blot, that non-phosphorylated Zap70 is associated with a substantial fraction of surface TCR in the preselection DP thymocytes via phospho-CD3<sup>ζ</sup> (39). As our data and published report show that DP thymocytes express the lowest amount of Zap70, and since recruitment of Zap70 to phospho-CD3<sup>ζ</sup> should promote its phosphorylation we asked whether Zap70 levels correlated with its phospho status. To this end, we used flow cytometry to determine relative amounts of phospho Y319 (pY319) and pY493 forms of Zap70, which represent partially and fully active kinase, respectively. As Zap70 is differentially expressed in the thymic and splenic subsets, we determined relative amounts of phophos-Zap70 by calculating the ratios of the two phospho forms of Zap70 to the total amount of Zap70 in each thymic or splenic T cell subset and normalized it to the average of ratio in CD8 SP thymocytes or mature T cells. Surprisingly, we observed that the relative amount of pY319 and pY493 Zap70 expression correlated inversely with the total Zap70 expression. Thus, DP thymocytes expressed significantly higher relative amount of pY319 and pY493 Zap70 than post-selection mature thymocyte subsets (Figure 2A). Further, CD4 SP thymocytes showed higher relative pY493 levels than CD8 SP thymocytes. Similar analysis of splenocytes showed higher relative amount of pY319 and pY493 Zap70 in CD4 compared to CD8 T cells although the difference was not significant (Figure 2B). Together, the inverse correlation between total Zap70 level and its phosphorylation suggest that partial and/or full Zap70 activation may negatively affect its stability.

**Zap70 Expression Is Negatively Regulated by Lck Activity**

It is well established that Lck is more active in CD4 than CD8 T cells (15, 44). As Lck phosphorylates Y319 and Y493 in Zap70 (45, 46) and data described above suggest an inverse correlation between phospho status of Zap70 and its total amount, we asked if differential Lck activity was the cause of differential Zap70 expression in CD4 and CD8 T cells. To test this prediction, we generated constitutively active Lck<sup>K273R</sup> and kinase inactive Lck<sup>K273F</sup> mutants. In these constructs Lck was fused to mCherry allowing us to monitor Lck expression by analyzing mCherry expression (41). We also generated 293T cells stably expressing human Zap70-fused to GFP (hZap70-GFP). This system allowed us to determine Zap70 expression by monitoring GFP levels (42). We transfected hZap70-GFP expressing 293T cells with Lck-mCherry fusion constructs and monitored GFP expression in mCherry expressing cells to determine the impact of Lck activity on Zap70 expression. Indeed, we observed significant decrease in Zap70 expression in the presence of Lck<sup>K273F</sup> (30% decrease in GFP) compared to control or WT Lck, while it was unaltered in the presence of...
kinase inactive Lck$^{K273R}$ mutant (Figure 3A). A similar decrease in mouse Zap70 expression was observed in the presence of Lck$^{Y505F}$ compared to WT Lck in 293T cells (data not shown). Importantly, we observed an opposite pattern for pY319 Zap70 staining; WT Lck expression resulted in a significant increase in pY319 Zap70 level compared with basal level in control transfectants, and this amount increased by 3-fold in the presence of Lck$^{Y505F}$ (Figure 3B). To extend this observation to T cells we transduced RLM, a CD4 SP thymocyte cell line, with lentivirus expressing WT or Y505F or K273R mutated Lck fused to mCherry and analyzed endogenous Zap70 expression by flow cytometry in mCherry+ve and mCherry-ve RLM cells. As expected, CD4 cell line expressing constitutively active Lck$^{Y505F}$ showed significant decrease in Zap70 levels compared to those expressing WT Lck or kinase inactive Lck$^{K273R}$ (Figure 3C). Zap70 expression was comparable in untransduced mCherry-ve RLM cells in all conditions. These data strongly suggest an inverse correlation between Zap70 amount and its activation state.

To provide in vivo evidence for this inverse correlation between Lck activity and Zap70 expression, we took advantage of mice expressing constitutively active Lck transgene (Lck$^{Y505F}$), which promotes redirection of MHCI-signaled thymocytes into CD4 lineage (10, 31). In addition, we recently showed that expression of Lck$^{Y505F}$ transgene augments Thpok-induced CD4 lineage choice of MHCI-signaled thymocytes (40). We
MHCII-/- mice also showed significant decrease in Zap70 expression compared to CD8 T cells from OTI+LckY505F mice. Interestingly, CD4 T cells consistently showed lower Zap70 expression relative to control mice compared to CD8 T cells (Figures 4A,B) (both restricted to MHCI).

Therefore asked if Zap70 expression was altered in T cells from MHCI-specific OTI-TCR transgenic mice expressing LckY505F. In agreement with our ex vivo data, we observed a dramatic reduction in Zap70 expression, as much as 70%, in the thymic subsets (Figure 4A) as well as mature CD4 and CD8 splenic T cells from OTI+LckY505F mice compared to OTI control mice (Figure 4B). This was confirmed by anti-Zap70 Western blot of splenic T cells from OTI+LckY505F and OTI control mice (Figure 4C). Anti-ubiquitin Western blot of anti-Zap70 immunoprecipitates of purified splenic T cells showed higher relative amounts of ubiquitinated Zap70 in purified splenic T cells from OTI+LckY505F compared to OTI control (Figure 4C). Interestingly, CD4 T cells consistently showed lower Zap70 expression compared to CD8 T cells from OTI+LckY505F mice although it was not significant (Figures 4A,B). Despite the decrease in total Zap70 the relative amounts of pY493 and pY319 were significantly increased in mature T cells from OTI+LckY505F mice compared to those in CD8 T cells from OTI mice (Figure 4D) supporting the idea that Zap70 expression inversely correlates with its activation status. Decrease in Zap70 expression thymocytes and mature T cells from MHCI-/- (Figure 5) and P14 (Figure S2) mice expressing LckY505F compared to control mice confirmed these data. Interestingly, we found that a small number of CD4 T cells that develop in MHCI-/- mice also showed significantly lower Zap70 expression compared to CD8 T cells (Figure 5) (both restricted to MHCI) providing further evidence for a role of the co-receptor associated Lck in regulating Zap70 expression.

To ensure the decrease in Zap70 expression was due to LckY505F expression rather than lineage redirection we performed Zap70 staining of thymocytes and mature T cells from OTI mice expressing the Thpok transgene. We previously showed that enforced Thpok expression promotes CD8 to CD4 lineage redirection in dose dependent manner and is augmented by LckY505F expression (40). Thus, we analyzed Zap70 expression in mice expressing two different Thpok transgenes, Thpok-163 and Thpok-H. We detected a small decrease in Zap70 levels in the redirected CD4 T cells while it remained unchanged in CD8 T cells from OTI mice expressing Thpok-163 (Figure 6) and Thpok-H transgene (Figure S3). Together, these data strongly suggest that Lck activity, while it augments Zap70 phosphorylation, negatively impacts the stability of Zap70 in SP thymocytes and mature T cells, and explain the lower Zap70 expression in CD4 compared to CD8 T cells in wild type mice.

**Phosphorylation of Zap70 Y315/Y319 Contribute to Lck-Induced Zap70 Degradation**

In its inactive form Zap70 is auto-inhibited by intra-molecular interactions between interdomain A (linking the two SH2 domains) and kinase domain mediated by Y315 and Y319 located in the interdomain B (linking C-SH2 to kinase...
domain). Lck phosphorylates Y315/Y319 in the interdomain B relieving auto-inhibition and leading to an open conformation of Zap70 that is partially active. Doubly phosphorylated pY315/pY319 Zap70 serves as a docking site for Lck via its SH2 domain causing phosphorylation of Y493 in the kinase domain and full activation of Zap70. Data described above strongly suggest an inverse correlation between Lck activity and Zap70 expression. Interestingly, Y315A/Y319A (YYAA) knockin mutations, which result in an open conformation, generate a hypomorphic Zap70 allele and impair thymic selection and mature T cell function in mice (47). Lck cannot bind to YYAA Zap70 mutant although the mutations promote auto-phosphorylation and LAT phosphorylation (48). This suggests that relieving auto-inhibitory constrain by phosphorylation of Y315/Y319 in Zap70 may induce its degradation. To test this notion we generated and co-transfected WT, Y319F or YYFF mutated Zap70 (also expressing ZsGreen reporter) with WT or constitutively active LckY505F (expressing NGFR reporter). YYFF mutated Zap70 cannot be phosphorylated by Lck, thus resulting in a constitutively closed auto-inhibitory structure that impairs TCR signaling and affects T cell development and function with Y319F, compared to Y315F, mutated Zap70 having more profound impact on T cell function (46, 48–50).

Taking advantage of YYFF or Y319F mutated Zap70, we analyzed and compared Zap70 expression in NGFR+ZsGreen+ cells. As predicted, constitutively active LckY505F had no effect on Zap70YYFF expression but resulted in severe reduction in Zap70 WT expression. Interestingly, Lck Y505F induced significantly less degradation of Zap70 Y319F compared to Zap70 WT but more compared to Zap70YYFF mutant (Figure 7).
These data provide substantial evidence that relieving auto-inhibition via Lck-mediated phosphorylation of Y315/Y319 renders Zap70 considerably unstable.

**DISCUSSION**

Lck and Zap70 kinases are crucial for proximal TCR signaling that forms the basis of thymic selection and CD4/CD8 lineage choice in the thymus and mature T cell homeostasis and response to antigenic challenge. While Lck is more active in CD4 than CD8 T cells (15, 44, 51), higher Zap70 levels are found in CD8 than CD4 T cells (35). Here, we have studied the mechanistic basis of differential Zap70 expression in correlation to Lck activity in thymic subsets and mature T cells employing flow cytometry staining that allows analysis at single cell level. Our data show that Lck-mediated phosphorylation of Y315 and Y319 in the interdomain B, which results in an open active conformation, is sufficient to induce rapid Zap70 degradation. Apart from its role in regulating TCR expression via CD3 ITAM phosphorylation and serving as a bridge between Zap70 and its substrate LAT (15, 52, 53), our finding that Lck kinase function negatively regulates Zap70 expression identifies a novel function for Lck and may explain several aspects of T cell biology (Figure 8). Given that the two kinases are essential for proximal events in signal transduction in all T cells (αβ, γδ, NKT cells) this simple mechanism of regulating Zap70 expression by Lck may regulate TCR signaling in developing thymocytes in response to selecting ligands critical for production of mature T cells devoid of autoimmunity and in mature T cells in response to self or foreign peptide.

In the thymus Zap70 expression inversely correlates with thymocyte maturation with expression in DP < CD4 < CD8 thymocytes with DP thymocytes expressing about 2.5 to 3-fold less Zap70 than CD8 SP thymocytes, and lower Zap70 expression in CD4 than CD8 SP thymocytes was retained in mature T cells with CD4 expressing lower Zap70 than CD8 T cells. Analysis of relative pY493 and pY319 specific staining showed exactly opposite pattern with higher relative expression in DP thymocytes compared to SP thymocytes. Previous study showed that non-phosphorylated Zap70 is associated with phospho-CD3 chain in the preselection DP thymocytes (39, 54). It is possible that lower amount of Zap70 in DP thymocytes combined to anti-pY Western blot technique likely resulted in lack of detection of pY-Zap70 in DP thymocytes in these studies. This opposite pattern of total and relative phospho-Zap70 levels prompted us to evaluate role of Lck-mediated phosphorylation of Zap70 in regulating Zap70 expression. If non-phosphorylated
Zap70 remains associated with phosho-CD3 chains it may act as a dominant negative (55) leading to impaired signaling in the preselection DP thymocytes. Lck-mediated phosphorylation of Y315/Y319 in Zap70 in DP thymocytes may serve two functions; in the absence of optimal receptor/ligand interaction it may lead to dissociation of Zap70 from TCR complex (56) and its subsequent degradation thereby preventing its potential dominant negative effect on TCR signaling. Such a mechanism may help explain sensitive nature of DP thymocytes to selecting self-peptide ligands (57); the “primed” Zap70 readily available for full activation upon engagement of DP thymocytes with a selecting ligand. However, we do not rule out the possibility that association of non-phosphorylated Zap70 to phospho-CD3 may protect it from degradation and that partially or fully open but free Zap70 is targeted for degradation in DP thymocytes thereby preventing undesired signal transduction in unsignaled DP thymocytes. Equally, fully Zap70 activation upon signal transduction in DP thymocytes, while necessary for efficient signaling, may cause over activation if its levels are not controlled; degradation would then regulate the amount of fully active Zap70 available for signaling (Figure 8).

Our data also explains paradox that sustained Zap70 expression is required for CD8 lineage choice, while augmented Lck activity promotes CD4 lineage choice (6, 10, 35, 40). It is likely that stronger association of Lck with CD4 ensures greater availability of relative amount of fully activated Zap70 in MHCII-signalized thymocytes leading to stronger signal and consequent CD4 lineage choice. This also ensures rapid degradation of partially activated Zap70 that could interfere in signaling and promote CD8 lineage choice in MHCII-signalized cells. In contrast, weaker Lck association with CD8 (15, 44) would result in relatively fewer phospho-Zap70 available for signaling leading to signal disruption and CD8 lineage choice in MHCII-signalized thymocytes. A recent report describing FlipFlop mouse strongly support the signal duration model of CD4 and CD8 lineage choice (58). We predict that in FlipFlop mouse even though CD8 coreceptor encoded by the Cd4 locus is weakly associated with Lck, its higher and continuous expression may result in accumulation of a sufficient amount of activated Lck, higher
Zap70 activation and its rapid degradation leading to helper lineage choice, while the converse may happen with lower and disrupted CD4 co-receptor expression from the Cd8 locus. Thus, cytotoxic lineage choice may be more susceptible to Zap70 amount, while helper lineage choice may be more affected by level of Lck activity (6). It may also explain why CD8 T cell, compared CD4 T cell, production is more severely affected by deficiency or hypomorphic expression of Zap70 in human patients (59–65).

Why would Lck, which promotes Zap70 activation critical for TCR signaling, negatively regulate Zap70 amount? Binding of Zap70 to phospho-CD3 promotes Zap70 activation due to partially destabilized auto-inhibition due to disruption of I-A linker and kinase domain interaction. This promotes Lck-mediated phosphorylation of Y315/Y319 and Y493 results in partially or fully active Zap70 essential for signal transduction during thymic maturation and mature T cell activation. However, partially or fully active Zap70 must be rapidly degraded to prevent potentially dominant negative effect of partially active Zap70 or over-activation during thymic development or mature T cell response to ligand. Partially or fully open but free (not associated with phospho-CD3) Zap70 may be rapidly degraded as well thereby preventing undesired antigen independent signaling during thymic development. It is quite possible that association of partially open Zap70 with phospho-CD3 may protect it from degradation.

![Model of Lck-mediated regulation of Zap70 expression/activation in T cell activation. In DP thymocytes higher amount of Zap70 is associated with surface TCR complex compared with mature T cells (ref 39) likely due to higher Lck activity resulting from CD4 and MHCII interaction (ref 15). We propose that the closed auto-inhibited conformation of Zap70 is stable and upon its recruitment to phosphor-CD3 chains Lck-mediated phosphorylation of Y315/Y319 and Y493 results in partially or fully active Zap70 essential for signal transduction during thymic maturation and mature T cell activation. However, partially or fully active Zap70 must be rapidly degraded to prevent potentially dominant negative effect of partially active Zap70 or over-activation during thymic development or mature T cell response to ligand. Partially or fully open but free (not associated with phospho-CD3) Zap70 may be rapidly degraded as well thereby preventing undesired antigen independent signaling during thymic development. It is quite possible that association of partially open Zap70 with phospho-CD3 may protect it from degradation.](image-url)
DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding author.

ETHICS STATEMENT

Animal experimentation protocols were approved by the CRHMR animal care committee and performed in accordance with the Canadian Committee for Animal Care.

AUTHOR CONTRIBUTIONS

HD, CT, and MV performed experiments. VD and HD analyzed data. D-CR provided reagents and resources. VD conceptualized the study, and VD and D-CR supervised the study. All authors discussed and commented on manuscript. All authors contributed to the article and approved the submitted version.

ACKNOWLEDGMENTS

We thank Jean-Philippe Bastien and Annabelle Minguy for comments; Martin Dupuis and Fred Duval for help with flow cytometry; Marie-Leur Durand and Marie-Philippe Boucher for animal help. We thank Johannes Huppa for human Zap70-GFP fusion plasmid (Medical University of Vienna) and Nicholas Gascoigne for Lck-mCherry fusion plasmid (National University of Singapore). This work was supported by the Hopital Maisonneuve-Rosemont Foundation-Isabelle Marcoux/Transcontinental Fund. All authors are members of the Quebec Cell, Tissue and Gene Therapy Network –TheCell (a thematic network supported by the FRQS).

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2022.935367/full#supplementary-material
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