Tonic LAT-HDAC7 Signals Sustain Nur77 and Irf4 Expression to Tune Naive CD4 T Cells

Graphical Abstract

Highlights

- Loss of the adaptor LAT leads to a Th2-biased lymphoproliferative disease in mice
- LAT transmits a tonic signal through PLCγ that phosphorylates the repressor HDAC7
- Tonic P-HDAC7 is cytoplasmic and does not repress target genes Nur77 and Irf4
- T cell Nur77 and Irf4 levels are critical to curb proliferation and differentiation

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In Brief

Myers et al. find that tonic signals through the adaptor LAT are critical to maintain naive T cell homeostasis. They delineate a tonic LAT-PLCγ-HDAC7 pathway that controls expression of HDAC7 targets. Disruption of the pathway leads to reduced expression of targets Nur77 and Irf4 and aberrant lymphocyte proliferation and differentiation.

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Tonic LAT-HDAC7 Signals Sustain Nur77 and Irf4 Expression to Tune Naive CD4 T Cells

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SUMMARY

CD4+ T cells differentiate into T helper cell subsets in feedforward manners with synergistic signals from the T cell receptor (TCR), cytokines, and lineage-specific transcription factors. Naive CD4+ T cells avoid spontaneous engagement of feedforward mechanisms but retain a prepared state. T cells lacking the adaptor molecule LAT demonstrate impaired TCR-induced signals yet cause a spontaneous lymphoproliferative T helper 2 (TH2) cell syndrome in mice. Thus, LAT constitutes an unexplained maintenance cue. Here, we demonstrate that tonic signals through LAT constitutively export the repressor HDAC7 from the nucleus of CD4+ T cells. Without such tonic signals, HDAC7 target genes Nur77 and Irf4 are repressed. We reveal that Nur77 suppresses CD4+ T cell proliferation and uncover a suppressive role for Irf4 in TH2 polarization; halving Irf4 gene-dosage leads to increases in GATA3+ and IL-4+ cells. Our studies reveal that naive CD4+ T cells are dynamically tuned by tonic LAT-HDAC7 signals.

INTRODUCTION

CD4+ T cells can polarize into T helper cell (Th) subsets, such as Th1 and Th2 (Th helper 2) subsets that produce either interferon-γ (IFN-γ) or interleukin-4 (IL-4), -5, and -13 cytokines (Mosmann et al., 2005). T cell receptor (TCR) signals, cytokine receptor signals, and transcription factors cooperate to establish these lineages via a feedforward loop (Zhu et al., 2010); TCR signals in the presence of IFN-γ result in the activation of STAT4 and induction of the transcription factor T-bet (Tbx21) that induces more IFN-γ expression (reviewed in Ansel et al., 2006; Murphy and Reiner, 2002). Analogously, in a IL-4/STAT6/GATA3/IL-4 induction, reinforcement, and maintenance model for Th2, an initial TCR signal leads to upregulation of IL-4 (termed “early IL-4”) so that subsequent TCR signals with a low-level IL-4/STAT6 signal lead to very robust induction of IL-4, -5, and -13 (Ansel et al., 2006; Paul, 2010). An initial model emerged with T-bet, GATA3, Foxp3, Rorγt, and Bcl6 transcription factors directing T cells to Th1, Th2, T REG (regulatory T cells), Th17, and Th17 (follicular helper T cells) lineages (Figure 1A), though it is clear that additional modes of regulation must exist (Locksley, 2009; O'Shea and Paul, 2010; Zhou et al., 2009; Zhu and Paul, 2010).

It is not completely understood how CD4+ T cells remain in a “resting,” naive state that is permissive to the above-mentioned feedforward mechanisms of Th subset differentiation. Epigenetic control mechanisms of cytokine loci can impact the effect of lineage transcription factors (Hirahara et al., 2011; Kanno et al., 2012), and additional transcription factors with broader expression patterns operate in transcriptional networks with the lineage-specific transcription factors (Li et al., 2014). One of these, Interferon regulatory factor 4 (Irf4), is expressed in different Th subsets (Biswas et al., 2010; Huber and Lohoff, 2014). Irf4 plays a critical role in Th2 differentiation; Irf4 cooperates with NFATc2 to promote IL-4 production and is critical for GATA3 upregulation, and GATA3 overexpression partially rescues IL-4 production by Irf4-deficient Th2 cells (Biswas et al., 2010; Huber and Lohoff, 2014).

Tonic or constitutive signals in B lymphocytes rely on surface immunoglobulin M (IgM) and are critical for survival (Lam et al., 1997). In T cells, tonic signaling also occurs (Monroe, 2006), but its physiological role is largely unknown (Hogquist et al., 2003). Survival of naive CD4+ T cells is only modestly impacted following inducible deletion of TCRα chains (Polic et al., 2001). Instead, tonic signals in T cells have been reported as immuno-modulatory, either enhancing (Stefanová et al., 2002) or blunting (Bhandoola et al., 2002; Smith et al., 2001) subsequent TCR responses to foreign antigen. Biochemically, tonic signals such as TCRγ phosphorylation can be detected in T cells rapidly isolated from peripheral lymphoid organs but not when isolated from peripheral blood, and these tonic signals quickly dissipate when cells are cultured in vitro in non-stimulatory conditions (Stefanová et al., 2002; van Oers et al., 1993, 1994). We previously established that the adaptor molecule LAT...
Linker for Activation of T cells (LAT) is critical for sending tonic Ras-ERK kinase signals, which can repress (Roose et al., 2003) or maintain (Markegard et al., 2011) expression of genes. Crippling LAT’s phospho-tyrosine docking site for PLCγ in the mouse germline via mutation of tyrosine 136 into phenylalanine (termed LAT Y136F here) results in a spontaneous TH2 hyperproliferative syndrome (Aguado et al., 2002; Sommers et al., 2002). Sophisticated mouse models with inducible LAT deletion (termed LATNEG here) or inducible switching from wild-type LAT to LAT Y136F demonstrated that the spontaneous TH2 hyperproliferative syndrome also develops when these LAT perturbations occur exclusively in peripheral T cells (Mingueneau et al., 2009; Shen et al., 2009) (Figure 1B). T cells with perturbed LAT function demonstrate impaired TCR-induced PLCγ activation and decreased PLCγ-dependent calcium and ERK signaling (Mingueneau et al., 2009; Shen et al., 2009) yet paradoxically take on a CD44 HIGHCD62LLOW-activated/memory T cell phenotype and produce high levels of intracellular IL-4 (Chuck et al., 2010; Mingueneau et al., 2009; Shen et al., 2009). Whereas it is known that major histocompatibility complex (MHC) class II and CD28 expression are required for the development of the Th2 immune pathology (Mingueneau et al., 2009), no further mechanistic insights relating loss of LAT in naive peripheral T cells to immune abnormalities have been established.

These new genetic mouse models ruled out altered T cell developmental or selection processes as the predominant causes for the Th2 hyperproliferative syndrome, which implies that there is an unappreciated role for LAT in providing some cue to curb CD4 + T cells in their naive state (Brownlie and Zamoyka, 2009). Here, we describe how tonic signals through LAT facilitate constitutive export of the transcriptional repressor histone deacetylase 7 (HDAC7) from the nucleus and thereby promote expression of immune-modulatory genes like Nur77 and Irf4. We reveal repressive roles for Nur77 and Irf4 and demonstrate that these act to tune the naive state of CD4 T cells.

RESULTS

Progressively Altered Gene Expression in Naive CD4 T Cells with LAT Perturbations

CD4 + T cells take on an activated/memory CD44 HIGHCD62LLOW phenotype when LAT is deleted or mutated via tamoxifen-induced Cre recombination (Chuck et al., 2010). In search of a direct molecular mechanism that underlies the hyperproliferative Th2 syndrome when T cells lack the adaptor LAT or express LAT Y136F (Figure 1B), we profiled gene expression in sorted naive CD4 + T cells that express LAT Y136F for 4 weeks demonstrating the most striking changes in gene expression (Figures 1B and S1). These changes could reflect selection of particular T cell clones that occurs over the 4-week period in the LAT Y136F model with a diverse TCR repertoire. LAT protein turnover takes 4 days (Ou-Yang et al., 2012); therefore, 1 week of tamoxifen treatment effectively results in 3 days of LATNEG or LAT Y136F. In search of the immediate and direct effects of LAT perturbation in naive CD4 + T cells, we focused on this short period of LAT perturbation.

HDAC Target Genes Are Repressed 3 Days after LAT Deletion or Y136F Mutation

Examination of expression levels of transcription factors that regulate helper T cell differentiation (Figure 2A) revealed no
conspicuous changes when CD4^{+\text{LOW}}CD4^{+} T cells are LAT^{\text{NEG}} or LAT^{\text{Y136F}} for 3 days. By contrast, expression of a cluster of genes including Egr1, Egr2, and Egr3, as well as Nr4a1 (encoding Nur77), Nr4a2, and Nr4a3 was greatly attenuated in both LAT^{\text{NEG}} and LAT^{\text{Y136F}} T cells (Figures 2B and S2). These genes are immediate-early response genes downstream of mitogen signals in many cell types. Several of these, N-ras in particular, have also been described as target genes of HDAC7 in both thymocytes (Dequiedt et al., 2003) and in DO11.10 T cell hybridoma cells (Kasler and Verdin, 2007).

HDACs deacetylate histone tails, which correlates with gene repression. HDACs are subdivided into four classes (I, IIa, IIb, and IV) (Haberland et al., 2009), and class I and IIa can be inhibited by Trichostatin A (Verdin et al., 2003). We previously reported that gene repression in a Jurkat T cell lymphoma line without LAT could be reversed by Trichostatin A (Markegard et al., 2011) and postulated that there may be a functional connection between LAT, HDACs, and the unexplained Th2 hyperproliferative syndrome. HDAC7 belongs to class IIa, and class IIa members (HDACs 4, 5, 7, and 9) display unique and tissue-specific expression patterns (Haberland et al., 2009; Verdin et al., 2003). Through TaqMan analyses, we established that HDAC7 is the predominant class IIa HDAC expressed in CD4^{+} T cells (Figure 2C) and therefore centered our attention on HDAC7.

We confirmed the reduced expression of Nr4a1 (Nur77) in sorted naive, CD4^{+} T cells by qPCR (Figure 2D). We utilized our previously published gene expression sets from thymocytes that either lack HDAC7 or express HDAC7ΔP, a constitutively nuclear, super-repressor version of HDAC7 with mutated serine phosphorylation sites (Kasler et al., 2011, 2012) to generate a list of potential thymic HDAC7 targets. We applied an arbitrary threshold of 2.0-fold differential expression between loss of HDAC7 and the HDAC7ΔP super-repressor, which resulted in 369 differentially expressed genes (Tables S1 and S2). The Venn diagram in Figure 2E demonstrates that the cluster of immediate early-response genes expressed at lower levels in LAT^{\text{NEG}} or LAT^{\text{Y136F}} naive CD4^{+} T cells overlap with HDAC7 targets in this analysis. In sum, the expression levels of a set immediate early-response genes are maintained by LAT and repressed by HDAC7; we will focus on the functional roles of Nur77 and Ir4 in CD4^{+} T cell later.

**HDAC7 Effects, Expression, and Phosphorylation**

Early support for a role for HDACs in T_{\text{H}} cell function came from studies where T cells were treated with HDAC inhibitors, which enhanced the expression of both IFN-γ and T_{\text{H}2} cytokines (Bird et al., 1998; Valapour et al., 2002). To test whether HDAC7 may, in principle, impact T_{\text{H}} cell function, we isolated T cells from mice with HDAC7 deletion or expression of HDAC7ΔP, the super-repressor version of HDAC7 (Kasler and Verdin, 2007; Kasler et al., 2011). Direct ex vivo stimulation followed by intracellular fluorescence-activated cell sorting (FACS) staining for cytokines revealed that HDAC7 deficiency led to increased percentages of CD4^{+} T cells producing IFN-γ (Figure 3A and S3A). Fewer HDAC7ΔP-expressing T cells expressed IFN-γ compared to wild-type (WT) (Figures 3B and S3B). HDAC7ΔP did not cause spontaneous increases in IL-4-producing cells under short-term stimulatory conditions (Figure 3B). Transduced purified WT T cells with a retroviral construct for HDAC7ΔP-GFP also revealed suppression of IFN-γ production (Figure S3C).

We next explored the possibility that HDAC7 levels could impact Th2 differentiation. Microarray expression data revealed no statistically significant differences in HDAC7 mRNA levels between WT, LAT^{\text{NEG}}, and LAT^{\text{Y136F}} CD4^{+}CD4^{+} T cells (Figure 3C). Relative HDAC7 protein levels fell when thymocytes developed into naive T cells and rose when these were stimulated (T_{\text{H}0}) or stimulated and polarized into different effector populations (Figure 3D). Comparisons of HDAC7 levels between in vitro-generated T_{\text{H}2} versus T_{\text{H}1}, T_{\text{H}2}, T_{\text{H}17}, and T_{\text{REG}} or T_{\text{H}0}; populations did not show striking differences (Figure 3D). These results argue that altered expression levels of HDAC7 are not the source of the LAT^{\text{NEG}} and LAT^{\text{Y136F}} T cell abnormalities.

Both HDAC-5 and -7 are regulated via phosphorylation of N-terminally located serine residues; phosphorylation leads to nuclear export and cytoplasmic retention of HDAC-5 and -7 and counteracts their repressive effects on gene expression (Dequiedt et al., 2003; Parra et al., 2005; Vega et al., 2004; Verdin et al., 2003; Zhang et al., 2002). B cell receptor (BCR)-induced phosphorylation of HDAC7-5 and -7 had previously been reported in avian- and murine-B cell lines and primary B cells (Matthews et al., 2006). We observed that TCR stimulation could induce phosphorylation of HDAC-5 or -7 transfected into a Jurkat T cell leukemia cell line (Figures 3E and 3F). However, we noted a substantial amount of tonic HDAC phosphorylation in the unstimulated Jurkat samples, which had also been observed in a B cell line system (Matthews et al., 2006). Examination of primary lymph node cells revealed that TCR-induced HDAC7 phosphorylation was very modest and contrasted with the robust induction of ERK phosphorylation. Instead, tonic HDAC-7 phosphorylation was very pronounced in these primary cells (Figure 3G). Given our previous studies on LAT’s role in tonic signals in non-stimulated cells and gene regulation (Markegard et al., 2011; Roose et al., 2003), we next investigated whether the adaptor LAT regulates HDAC7 phosphorylation, localization, and function in a tonic fashion.

**LAT-Dependent, Tonic Nucleo-cytoplasmic Shuttling of HDAC7**

HDAC7 has nuclear (N) localization in CD4^{+}CD8^{+} double positive thymocytes but becomes predominantly cytoplasmic (C) in more developed CD4^{+} or CD8^{+} single positive thymocytes (Kasler et al., 2011). Unstimulated, naive CD4^{+} T cells maintain the predominant cytoplasmic distribution of HDAC7 seen in their predecessor CD4-single positive thymocytes (Figure 4A), which agrees with the evident tonic phosphorylation of HDAC7 in primary lymph node cells (Figure 3G). These data show that tonic signals in CD4^{+} T cells constitutively keep HDAC7 phosphorylated and exported from the nucleus. Uniform cytoplasmic localization of HDAC7 in T_{\text{H}0}, T_{\text{H}1}, T_{\text{H}2}, T_{\text{H}17}, and T_{\text{REG}} cells demonstrated that nucleo-cytoplasmic trafficking occurs efficiently in all T_{\text{H}} subsets (Figure 4A).

Primary cells rapidly lose tonic signals in vitro (Stefanová et al., 2002; van Oers et al., 1993) making biochemical studies on tonic signaling and pathway mapping challenging; we therefore used a model cell line to investigate biochemical details of the tonic
A 1 week Tamoxifen

Ratio Gene Expression

LAT^Y136F
LAT^NEG
LAT^WT
LAT^WT

Gene expression in CD4^+CD44^low T cells

B

Gene expression in CD4^+CD44^low T cells

wt control
1 week
LatNEG
1 week
LatY136F
1 week

C

Relative Gene Expression

Heart
Thymus
CD4^+ T cells

Hdac4
Hdac5
Hdac7
Hdac9

D

Relative Nur77 expression

Control
LAT^NEG
LAT^Y136F
model
mouse

+ tamoxifen

E

Down in LAT Mutant
19

Down in LAT K0
15

Thymic HDAC7 Targets
12

Nr4a1 (Nur77)
Irft
Egr1
Egr2
Gpr83
Egr3
Nr4a3
Dusp2
Nr4a2
Fkbp5

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signals. To examine LAT, we used a LAT-deficient Jurkat T cell line, J.CaM2. We determined that tonic phosphorylation of HDAC-5 and -7 is decreased when cells lack LAT and is partially restored when J.CaM2 cells are stably reconstituted with a WT LAT cDNA construct (Figures 4B and 4C). Our results with the LATY136F mouse model point to a role for PLCγ1 in the tonic signaling pathway, since PLCγ1 normally docks at phosphorylated Y136. We previously utilized overexpression of DGKζ to reduce tonic PLCγ1-diacylglycerol (DAG) signals as DGKζ converts DAG to phosphatidic acid (Markegard et al., 2011). Overexpression of FLAG-tagged DGKζ expression reduced tonic phosphorylation of HDAC-5 in unstimulated Jurkat T cells (Figure 4D), indicating that a LAT-PLCγ1-DAG pathway is an important component of the tonic signaling pathway. Concomitant with decreased tonic HDAC phosphorylation in LAT-deficient J.CaM2 cells, its nuclear/cytoplasmic ratio increased 4.8-fold when cells do not express the adaptor LAT (Figure 4E).

**Tonic Regulation of HDAC7 Impacts Nur77 Expression and Proliferation of CD4⁺ T Cells**

To substantiate our findings in Figure 4, we now exploited the fact that tonic signals rapidly dissipate when primary cells are rested ex vivo in non-stimulatory medium (Stefanová et al., 2002; van Oers et al., 1993). Resting lymph node cells for 30 or 60 min in PBS at 37°C resulted in a 70%–80% decrease of tonic HDAC7 phosphorylation as well as in previously reported decreases of tyrosine-phosphorylated proteins detected by 4G10 immunoblotting (Figure 5A). Analysis of HDAC7 with CD4-costaining on cytopsins of cells fixed immediately after isolation revealed a predominantly cytoplasmic HDAC7 distribution in CD4⁺ T cells (Figure 5B) with the highest pixel intensity for HDAC7 at the cell perimeter (Figure 5C). Resting cells for 30 min in PBS before fixation resulted in a translocation to predominantly nuclear HDAC7 localization (Figures 5D and 5E), as expected based on our phosphorylation data (Figure 5A).

CD5 expression is a sensitive reporter of TCR affinity (Azzam et al., 1998) and can be used as a marker of tonic TCR signaling (Mandl et al., 2013). Naïve CD4⁺ T cells display a range of CD5 expression in which CD5HIGH cells received most tonic signal input (Mandl et al., 2013). We used CD44 and CD5 markers to sort CD44LOW naïve CD4⁺ T cells into the most bright and most dim for CD5 while keeping cells ice cold and established that CD5HIGH naïve CD4⁺ T cells have substantially more tonic HDAC7 phosphorylation than CD5LOW naïve CD4⁺ T cells (Figure 5F). Furthermore, CD5HIGH naïve CD4⁺ T cells expressed significantly higher mRNA levels of the HDAC target Nur77 compared to their CD5LOW counterparts (Figure 5G). Single deficiency for Nur77 leaves T cell development largely intact (Lee et al., 1995). More recently it has been reported that loss of Nur77 results in increased proliferation of stimulated CD8⁺ T cells (Nowyhed et al., 2015), implying that Nur77 has suppressive functions. In support of this notion, we found that stimulation of sorted naïve CD4⁺ T cells resulted in more proliferation measured by CTV dilution when these cells lack Nur77 (Figure 5H), indicating that Nur77 not only is simply under tonic signal control, but provides a negative feedback loop to suppress T cell proliferation. In sum, by utilizing three systems—induced LAT perturbation, PBS rest, and CD5-based sorted populations—we demonstrate that naïve CD4⁺ T cells constitutively export HDAC7 from the nucleus in a dynamic manner that relies on tonic LAT-PLCγ1-DAG signals. Loss of this tonic control signal results in reduced expression of immediate-early response genes; one of these, Nur77, functions to suppress T cell proliferation (Figure 5).

**Tonic Regulation of Irf4 Expression Limits Th2 Polarization of CD4⁺ T Cells**

Our gene expression analyses also indicated that tonic LAT-HDAC7 signals maintain Irf4 expression in naïve T cells (Figure 2). Irf4 is a member of the Irf (interferon regulatory factor) family of transcription factors (Lohoff and Mak, 2005; Tamura et al., 2008) and plays a critical role in the function and homeostasis of mature but not developing T cells (Mittrücker et al., 1997). Irf4 function in T cells is complex. First, specific ablation of Irf4 in Foxp3-positive regulatory T cells yielded the surprising phenotype of a spontaneous Th2 immune disorder (Zheng et al., 2009); thus, Irf4 fulfills a Th2-intrinsic role to prevent effector T cells from polarizing toward Th2 (Figure 6A). Multiple studies identified Irf4 as a critical factor promoting differentiation of Th2 cells and production of Th2 cytokines under Th2 polarizing conditions in vitro and demonstrated that Irf4 is required for Th2 responses in vivo (reviewed in Biswas et al., 2010; Huber and Lohoff, 2014). However, it has also been observed that Irf4 inhibits IL-4 production in naïve CD4⁺ T cells from BALB/c mice (Honma et al., 2008). Thus, Irf4 may have both Th2-stimulatory and Th2-suppressing functions in CD4⁺ T cells depending on their state (Figure 6B).

TCR stimulation results in strong upregulation of Irf4 expression (Biswas et al., 2010; Huber and Lohoff, 2014). In CD8⁺ T cells, Irf4 expression levels increase in a graded manner directly proportional to the strength of incoming TCR receptor signal (Nayar et al., 2012, 2014; Yao et al., 2013). TaqMan analysis revealed that Irf4 levels are reduced by roughly 60% in both
Figure 3. HDAC7 Represses IFN-γ and Is Constitutively Phosphorylated in CD4+ T Cells

(A and B) Bar graph representation of percentages of cytokine-producing cells isolated from WT mice compared to HDAC7-deficient (KO in A) and HDAC7 ΔP-transgenic mice (ΔP in B) that were acutely stimulated with PMA and ionomycin for 4 hr ex vivo in the presence of Brefeldin A. Data are representative of three to six (A) or five (B) independent experiments with one to two mice/group, and error bars represent the SEM. See also Figure S3.

(C) Gene expression levels of HDAC7 in LAT WT, LATY136F, and LAT NEG mice. Gene expression levels (mean ± SEM) were obtained from the three data points from the microarray with 1-week tamoxifen treatment. ns, not significant.

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LAT\textsuperscript{NEG} and LAT\textsuperscript{136F} CD44\textsuperscript{LOW}CD4\textsuperscript{+} T cells (Figure 6C). Using the CD5-sorting strategy, we confirmed that baseline Irf4 expression levels are impacted by tonic signaling; CD5\textsuperscript{LOW} naive CD4\textsuperscript{+} T cells with the lowest level of tonic signaling reproducibly expressed roughly half of the Irf4 mRNA and Irf4 protein compared to CD5\textsuperscript{HIGH} naive counterparts (Figures 6D and 6E).

To specifically address how decreases in Irf4 expression levels impact CD4\textsuperscript{+} T cells, we generated an Irf4 allelic series of two WT copies, one WT copy, or zero WT copies of Irf4 by crossing floxed Irf4 mice to CD4-Cre mice (Figure 6F). Irf4 TaqMan revealed a 40% reduction in Irf4 expression in heterozygous Irf4\textsuperscript{+/−} CD4 T cells (Figure 6G). CD3/CD28 stimulation of sorted Irf4\textsuperscript{+/−} naive CD4\textsuperscript{+} T cells (CD44\textsuperscript{LOW} with exclusion of CD25\textsuperscript{+} cells) induced upregulation of Irf4 expression, in agreement with published work (Biswas et al., 2010; Huber and Lohoff, 2014), but Irf4 levels lagged behind in Irf4\textsuperscript{−/−} cells (HET), measured after 10 hr (Figure 6H). Functionally, an increased percentage of Irf4\textsuperscript{−/−} cells expressed GATA3 compared to WT cells stimulated for 10 hr in T\textsubscript{i}2-polarizing culture conditions (Figure 6i). Thus, Irf4 plays a suppressive role during the T\textsubscript{i}2 initiation phase, when cells are still more naive. This suppressive effect of Irf4 was still detectable in 5-day polarization assays: while full Irf4 deletion resulted in the complete absence of IL-4 induction, we observed an increase in percentage of IL-4-producing cells when the gene dose of Irf4 is halved (Irf4\textsuperscript{+/−}), compared to cells with WT levels of Irf4 (Figure 6j). Using CD4-Cre, these results unequivocally demonstrate that some (expression level) of Irf4 is critical for any polarization toward T\textsubscript{i}2 cells but also revealed that normal expression levels of Irf4 that are sustained by tonic signals function to curb T\textsubscript{i}2 polarization.

**DISCUSSION**

Mouse models with T cell-specific LAT perturbation develop a T\textsubscript{i}2 hyperproliferative syndrome through unknown mechanisms (Chuck et al., 2010; Mingueneau et al., 2009; Shen et al., 2009). Utilizing three distinct experimental systems, we demonstrate the existence of dynamic maintenance cues in the form of tonic signals in naive CD4\textsuperscript{+} T cells that rely on the presence of an intact LAT adaptor molecule. We reveal that these tonic effects are critical for constitutive phosphorylation and nuclear export of the repressor HDAC7 in naive CD44\textsuperscript{LOW}CD4\textsuperscript{+} T cells; without tonic HDAC7 regulation, expression of target genes becomes repressed. We particularly focused on the LAT-HDAC7 target genes Nur77 and Irf4 and revealed that these are immunosuppressive; these data offer mechanistic insights into how tonic signals tune the naive state of CD4\textsuperscript{+} T cells.

We show that Nur77 fulfills a negative feedback function to suppress proliferation of CD4\textsuperscript{+} T cells. In addition, one of Nur77’s reported targets is Fasl (Fas ligand) (Rajpal et al., 2003), which regulates T cell homeostasis via induction of apoptosis of activated T cells. Fasl is also a target of Egr transcription factors (Rengarajan et al., 2000), and we found Egr-1, Egr-2, and Egr-3 expressed at reduced levels in T cells with perturbed LAT function (Figure 2B). LAT\textsuperscript{136F} T cells fail to upregulate Fasl in response to TCR engagement (Ou-Yang et al., 2012; Sommers et al., 2002). The transcription factor Irf4 has received a lot of interest as a T cell fate-determining factor (Biswas et al., 2010; Huber and Lohoff, 2014). We were inspired by the reported linear correlation between strength of TCR signal and Irf4 expression levels (Nayar et al., 2012, 2014; Yao et al., 2013), and our studies revealed that Irf4 expression in naive CD4\textsuperscript{+} T cells is under tonic control. Sustained and optimal Irf4 expression functions to curb T\textsubscript{i}2 differentiation, as a larger portion of Irf4\textsuperscript{−/−} T cells, which have a 40% reduction in Irf4 mRNA, express GATA3 and IL-4. Recent work uncovered that Irf4 interacts with Fos/Jun heterodimers of the activator protein-1 (AP1) complex to regulate genes that contain AICE elements (AP-1/IRF composite elements) (Glasmacher et al., 2012; Li et al., 2012; Murphy et al., 2013). It is possible that reduced Irf4 levels result in altered composition of transcription factor complexes and/or altered regulation of AICE elements, which is an area for future investigation. Together, our studies with emphasis on Nur77 and Irf4 offer an explanation how perturbation of a tonic signal via LAT can cause a T\textsubscript{i}2 hyperproliferative syndrome.

Our studies here focused on tonic LAT-HDAC signals in naive CD4\textsuperscript{+} T cells, but it is possible that such signals are important for regulatory T cells as well. We previously demonstrated that perturbations in LAT lead to impaired T\textsubscript{REG} function (Chuck et al., 2010; Shen et al., 2010), and recent studies support the idea of functional constitutive signals in T\textsubscript{REG}—deletion of the TCR in T\textsubscript{REG} resulted in impaired homeostasis of T\textsubscript{REG} (Levine et al., 2014; Vahi et al., 2014). Notably, TCR deletion in T\textsubscript{REG} did not impact expression of FoxP3 or other T\textsubscript{REG} signature genes, but, instead, expression of Egr1, Egr2, Nr4a1 (Nur77), Irf4, and Ctia4 was reduced in the TCR-ablated T\textsubscript{REG} (Levine et al., 2014; Vahi et al., 2014). Since we find that these same genes are under tonic LAT-HDAC7 control in naive CD4\textsuperscript{+} T cells, this suggests that tonic LAT-HDAC7 signals may also regulate this gene set in T\textsubscript{REG}. Furthermore, Nur77 and the two other Nr4a family members are critical for the generation of T\textsubscript{REG} (Sekiya et al., 2013) and suppression of T\textsubscript{i}2 by T\textsubscript{REG} (Sekiya et al., 2015), and Irf4 was shown to have a T\textsubscript{REG}-intrinsic role to suppress T\textsubscript{i}2 differentiation as well (Zheng et al., 2009).

Together, our studies presented here combined with the above-cited work put forth a model in which loss of tonic LAT-HDAC7 signals alters the tuning of naive CD4\textsuperscript{+} T cells (as well
as the homeostasis of T_{REG} and that tonic regulation of Nur77 and Irf4 levels are important mediators and feedback regulators for maintaining the naive, undifferentiated state. We propose that dynamic maintenance cues in T cells should be considered in the context of cancer immunotherapy and immune diseases, where these tonic signals may either be influenced by checkpoint blockade therapy or be altered through SNPs. In support of the latter idea, it is of interest to note that three Nur77 SNPs have been associated with severity of bronchial hyperresponsiveness in asthma patients (Kurakula et al., 2015) and that a SNP in the 3' UTR region of Irf4 has been associated with recurrent bronchitis in children (Pinto et al., 2013).
Figure A: Lymph node cells were treated with different concentrations of HDAC7 inhibitors for 0, 30, or 60 minutes in PBS. The inhibitors were P-HDAC7 and HDAC7.

Figure B: Direct fixation of CD4 and HDAC7 expression in cells.

Figure C: HDAC7 pixel intensity across cell diameter.

Figure D: CD4 and HDAC7 expression in cells after 30 minutes of PBS rest.

Figure E: HDAC7 pixel intensity across cell diameter.

Figure F: CD4+CD44^LOW T cells were sorted into CD5^LOW and CD5^HIGH populations.

Figure G: Relative Nur77 expression in CD5^HIGH and CD5^LOW CD4^+ T cells.

Figure H: WT cells treated with Nur77^−/− and CTV expression at 72 and 96 hours.

Figure I: LAT Perturbation with Reduced Nr4a1 & Irf4 expression.
**Experimental Procedures**

**Mice**

WT C57BL/6 mice were bred in house at UCSF, LAT-/- and LATY136F mice crossed to ER-Cre (Chuck et al., 2010; Markegard et al., 2011) and HDAC7 KO mice and HDAC7-ΔP mice (Kasler et al., 2011, 2012) have been described previously. Nur77+/− mice were obtained from The Jackson Laboratory, Irr34th (The Jackson Laboratory) and CD4-Cre (Taconic Farms) mice were purchased and crossed. All mice used in experiments were between 8 and 14 weeks of age, and both males and females were analyzed. Mice were housed and treated in accordance with the guidelines of the Institutional Animal Care and Use Committee (IACUC) guidelines of the University of California, San Francisco (AN098375-03B for J.P.R. and AN111072-01 for E.V.) and Duke University (A160-14-06 for W.Z.). All animal experimentation occurred with institutional oversight and IACUC approval (protocol AN145252-01A, approval date: October 4, 2016).

**Microarrays**

Microarray analysis of CD4+ T cells was carried out using Agilent gene arrays using protocols optimized in our group and the UCSF Functional Genomics Core Facility. For details, see the Supplemental Information.

**Real-Time qPCR**

RNA was extracted from harvested cells using Trizol, the RNeasy kit (Qiagen), and treated with DNase I. Random primers (Invitrogen) and M-MLV Reverse Transcriptase (Invitrogen) were used to generate cDNA. For mRNA gene expression assays, TaqMan primers/probe were purchased from Life Technologies: HDAC7: Mm00469527_m1, Nr4a1 (Nur77): Mm01300401_m1, Irf4; Mm00516431_m1, and Actb: Mm00230941_g1. TaqMan Real-Time PCR was performed using SensiMix II Probe Kit (Bioline). TaqMan reactions were run on a Mastercycler EP Reaplex system (Eppendorf) in triplicate. Values was performed using SensiMix II Probe Kit (Bioline). TaqMan reactions were run on a Mastercycler EP Reaplex system (Eppendorf) in triplicate. Values were normalized to β-actin on each sample.

**Cell Lines, Transfections, Cell Isolation, Stainulations, and Fractionations**

Jurkat T cells and derived J.CaM2 cells were grown, transfected, and stimulated as described before (Das et al., 2009; Markegard et al., 2011). In short, cells were transfected with 10 μg of HDAC7-GFP, HDAC5-GFP, or GFP, rested overnight, and stimulated with 30 ng/mL antibody to trigger TCR signaling. For biochemical analysis, lymph node CD4+ T cells were purified through magnetic-activated cell sorting (MACS) depletion (Miltenyi) using negative isolation and, where indicated, were subsequently sorted to >98% purity on a MoFlo XDP (Beckman Coulter) or a FACs Aria (BD Biosciences). Cells were cultured for 72 and 96 hr on CD3/CD28-coated plates (0.5 μg/mL each) at which point viable CD4 + cells were analyzed for CTV dilution by flow cytometry. Cells were cultured for 72 and 96 hr on CD3/CD28-coated plates (0.5 μg/mL each) at which point viable CD4+ cells were analyzed for CTV dilution by flow cytometry. Cells were stimulated in vitro with α-CD3 (2C11, UCSF mAb core) and α-CD28 (clone GK1.5, UCFS Hybridoma Core) followed by secondary antibodies; goat-anti-rabbit antibody conjugated to Alexa Fluor 568 (Thermo Fischer Scientific), and subsequently sorted to >98% purity. Cells were stimulated in vitro with α-CD3 (2C11, UCSF mAb core) and α-CD28 (clone GK1.5, UCFS mAb Core) before (Das et al., 2009) and lysed with SDS lysis buffer. Nuclear and cytoplasmic fractions were prepared from indicated T cell subsets using NE-PER Nuclear and Cytoplasmic Extraction Kit (Thermo Fisher Scientific, catalog no. 78838).

**Immunoblotting**

Western blot analyses were performed and quantitated using a LAS3000 Imaging System (Fuji) and MultiGauge software as described previously (Das et al., 2009; Markegard et al., 2011). Antibodies to phosphorylated and total ERK, Irf4, and actin were from Cell Signaling Technology; antibodies to HDAC7 and Grb2 were from Santa Cruz Biotechnology; antibody to Gfp was from Clontech; antibody to lamin b was from Abcam, and antibody to histone 3 was from EMD Millipore. Rabbit antiserum to phosphorylated HDAC7 (S259) was a gift from Dr. Timothy McKinsey.

**Microscopy**

For microscopic analysis of HDAC7 distribution in cytopsins, single-cell suspensions were prepared from lymph nodes, washed with PBS, and either immediately fixed in 2% paraformaldehyde or were rested in PBS for 30 min prior to fixation. Cells were then washed and spun onto slides using a Cytospin (700 rpm, 4 min) and stained with primary antibodies to HDAC7 (Santa Cruz Biotechnology) and CD4 (clone GK1.5, UCFS Hybridoma Core) followed by secondary antibodies; goat-anti-rabbit antibody conjugated to Alexa Fluor 568 (Invitrogen, A-11036) for HDAC7 and goat anti-actin antibody conjugated to Alexa Fluor 488 (A-11006) for CD4. Cells were counterstained with DAPI. Images were acquired on a Zeiss spinning disk confocal microscope. Images and pixel intensity were analyzed using ImageJ software.

**Flow Cytometry Assays on Primary T Cells**

CD4+ T cells used in polarization assays were purified from lymph nodes using MACS negative isolation and, where indicated, were subsequently sorted to >98% purity on a MoFlo XDP (Beckman Coulter) or a FACS Aria (BD Biosciences). Cells were stimulated in the UCSF flow cytometry core. Cells were stimulated in α-CD3 (2C11) and α-CD28 (37.51)-coated plates in the presence of Th2-polarizing media (50 ng/mL rIL-4 [PeproTech] and 10 μg/mL α-IFN-γ [XMG1.2, UCFS mAb core]). After 5 days, cells were restimulated for 4 hr with phorbol-12-myristate-13-acetate (PMA) (5 ng/mL) and ionomycin (0.67 mM) in the presence of monensin (BD GolgiStop). Following restimulation, cells were labeled with Live/Dead fixable viability dye (Life Technologies) and antibodies to CD4 (GK1.5), IFN-γ (XMG1.2), and IL-4 (11B11). For GATA3 assays, cells were stimulated on coated plates for up to 48 hr in Th2 media before fixation and permeabilization in Foxp3/Transcription Factor Buffer set (eBioscience) and staining with GATA3 (TWAJ, eBioscience).

For proliferation assays, CD4+ T cells from Nur77+/− and WT B6 lymph nodes were enriched using MACS negative isolation, labeled with Cell Trace Violet (Thermo Fischer Scientific), and subsequently sorted to >98% purity. Sorted cells were cultured on CD3/CD28-coated plates for up to 96 hr prior to FACS analysis.

**Figure 5. Tonic Signals in CD4+ T Cells Dynamically Regulate HDAC7 and Its Target Nur77 to Limit Proliferation**

(A) Immunoblotting for phosphorylated HDAC7 (top) and total phosphorytrosine (4G10, bottom) in lymph node cells isolated from mice that were lysed directly or first restimulated in PBS for the indicated time points.

(B) Immunofluorescence of HDAC7 localization in CD4+ T cells in a cytospin preparation co-stained for HDAC7 (red) and CD4 (green). Lymph node cells were immediately fixed and spun onto slides prior to preparation for imaging.

(C) Quantification of pixel signal intensity across the diameter of the 10 CD4+ T cells was obtained with Fiji image analysis software.

(D and E) Analysis of HDAC7 localization exactly as in (B) and (C), except that lymph node cells were restimulated in PBS for 30 min prior to fixation and cytokin analysis. Note the switch to predominant nuclear HDAC7 localization following PBS rest.

(F) Schematic of CD5 sorting scheme of naive CD4+ T cells from lymph node and immunoblot analysis of phospho-HDAC7 and -ERK levels in sorted populations. An arbitrary 20% gap was chosen to segregate CD5−high from CD5−low cells yet obtain sufficient cells for biochemistry. Total HDAC7 and ERK serve as loading controls.

(G) TaqMan qPCR analysis of Nur77 mRNA levels in CD5-sorted cells. TaqMan samples were run in triplicate. Nur77 mRNA levels were arbitrarily set at 1.0 in one CD5−low sample. Unpaired t test: p = 0.018 (indicated by *). (A)−(G) are representative examples of three independent experiments. Error bars represent SEM.

(H) Flow-cytometric analysis of Cell Trace Violet (CTV) dilution in sorted (TCR+) CD4+ CD25− CD44low naive CD4+ T cells from two WT and two Nur77−/− mice. Cells were cultured for 72 and 96 hr on CD3/CD28-coated plates (0.5 μg/mL each) at which point viable CD4+ cells were analyzed for CTV dilution by flow cytometry. Data are representative of two independent experiments.

(i) Cartoon summarizing how tonic signals through LAT influence phosphorylation and nucleo-cytoplasmic shuttling of HDAC7 and expression of HDAC7 target genes.
Figure 6. Irf4 Expression Is under Tonic Signal Regulation and Curbs T_{H2} Polarization of CD4+ T Cells

(A) Schematic of the role of Irf4 in regulatory T cells (T_{REG}) and suppressive effects on T_{H2} differentiation.

(B) Schematic of the role of Irf4 in effector (top) and naive (bottom) CD4+ T cells in controlling T_{H2} differentiation and IL-4 production.

(C) TaqMan qPCR analysis of Irf4 levels in T cells isolated from 4-week tamoxifen-treated WT, LAT KO, and LAT Y136F mice. Representative example of three independent assays is shown on these three samples.

(D and E) (D) TaqMan qPCR analysis (n = 3) and (E) immunoblot analysis (n = 2) of Irf4 levels in CD5-sorted cells. TaqMan samples were run in triplicate. Irf4 mRNA levels were arbitrarily set at 1.0 in one CD5 LOW sample. Unpaired t test: p = 0.034 (indicated by *). In (D), error bars represent SEM.

(F) Cartoon of Irf4 allelic series.

(legend continued on next page)
All antibodies used were from eBioscience, BD, or Tonbo Biosciences. All flow cytometry data were acquired on LSRII or Fortessa instruments (BD Biosciences) and analyzed using FlowJo (Tree Star). Statistical significance was determined using unpaired t tests.

ACCESSION NUMBERS

The accession number for the microarray data reported in this paper is GEO: GSE76897.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, three figures, and two tables and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2017.04.078.

AUTHOR CONTRIBUTIONS

M.Z. provided material form LAT mouse models. A.B. performed the array in Figure 1C. H.K. performed the analyses in Figure 2E. H.W.L. executed Figures 3A, 3B, 3D, and 4A. J.P.H. performed experiments in Figure 5H. D.R.M., T.L., and E.M. performed all other experiments and interpreted data. J.Z., D.J.E., E.V., and W.Z. provided reagents and technology and edited the manuscript. J.P.R. conceived the study, oversaw all work and the collaboration, and wrote the manuscript with D.R.M. and T.L.

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(B) TaqMan qPCR analysis of ifr4 levels in T cells isolated from ifr4 WT and ifr4 KO CD4+ Cre mice. Representative example of two independent experiments performed in triplicate. Unpaired t test; p = 0.0005 (indicated by **). Error bars represent SEM.

(G) Flow cytometry for ifr4 expression in sorted CD4+ CD25+ CD44low T cells from ifr4 WT and HET mice cultured for 10 hr in T2,2-polarizing conditions on α-CD3/CD28-coated plates (0.5 μg/mL each). Data are representative of two independent experiments with two mice/group.

(H) Flow cytometry for ifr4 expression in sorted CD4+ CD25+ CD44low T cells from ifr4 WT and HET mice cultured for 10 hr in T2,2-polarizing conditions on α-CD3/CD28-coated plates (0.5 μg/mL each). Data are representative of two independent experiments with two mice/group.

(J) Flow-cytometric analysis of IL-4 production in viable T2,2-polarized naive CD4+ CD25+ CD44low T cells sorted from ifr4 WT and ifr4 KO CD4+ Cre mice (WT), ifr4 KO CD4+ Cre mice (KO) and ifr4 WT and ifr4 KO mice stimulated with 0.5 μg/mL α-CD3 and α-CD28. Percentage of cytokine-positive cells was calculated, and data were normalized and compiled from three independent experiments with two mice per group. Error bars represent SEM, and significance was determined using an unpaired t test, with Th2 WT versus HET p = 0.0446 (indicated by *) and WT versus KO p < 0.0001.


