

Critical role for TRIM28 and HP1 β/γ in the epigenetic control of T cell metabolic reprograming and effector differentiation

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Naive CD4⁺ T lymphocytes differentiate into different effector types, including helper and regulatory cells (Th and Treg, respectively). Heritable gene expression programs that define these effector types are established during differentiation, but little is known about the epigenetic mechanisms that install and maintain these programs. Here, we use mice defective for different components of heterochromatin-dependent gene silencing to investigate the epigenetic control of CD4⁺ T cell plasticity. We show that, upon T cell receptor (TCR) engagement, naive and regulatory T cells defective for TRIM28 (an epigenetic adaptor for histone binding modules) or for heterochromatin protein 1 β and γ isoforms (HP1 β/γ , 2 histonebinding factors involved in gene silencing) fail to effectively signal through the PI3K-AKT-mTOR axis and switch to glycolysis. While differentiation of naive TRIM28^{-/-} T cells into cytokine-producing effector T cells is impaired, resulting in reduced induction of autoimmune colitis, TRIM28^{-/-} regulatory T cells also fail to expand in vivo and to suppress autoimmunity effectively. Using a combination of transcriptome and chromatin immunoprecipitation-sequencing (ChIP-seq) analyses for H3K9me3, H3K9Ac, and RNA polymerase II, we show that reduced effector differentiation correlates with impaired transcriptional silencing at distal regulatory regions of a defined set of Treg-associated genes, including, for example, NRP1 or Snai3. We conclude that TRIM28 and HP1 β/γ control metabolic reprograming through epigenetic silencing of a defined set of Treg-characteristic genes, thus allowing effective T cell expansion and differentiation into helper and regulatory phenotypes.

T cells | immunology | epigenetics | autoimmunity | TRIM28

T cells participate and control the vast majority of adaptive immune responses. While $CD8^+$ T cells mediate killing of virus-infected and tumor cells, $CD4^+$ T cells positively (helper T cells, Th) and negatively (regulatory T cells, Tregs) regulate both cytotoxic and antibody responses. Similar to pluripotent stem cells, which retain the ability to differentiate into tissue-specific cell types, naive cells leaving the thymus have the capacity to differentiate into distinct subsets of effector or regulatory cells upon activation that differ in their transcriptional programs and metabolic needs. For example, while all T cells rely on glucose metabolism for clonal expansion, only Th effector functions are linked

to glycolysis, while the suppressive capacity of Tregs has been linked to β -oxidation of fatty acids (1, 2).

Significance

CD4 T cells are major regulators of immune responses against both self and pathogens. Understanding pathways that govern CD4 T cell differentiation and regulation are thus key for the discovery of new immunoregulatory drug targets. Here, we have identified an epigenetic pathway that regulates the expression of a set of proteins that determine T cell responsiveness. By silencing enhancers distal to a set of genes known to be involved in regulatory T cell function, the epigenetic modifiers TRIM28 and HP1 β/γ regulate T cell receptor signaling. This leads to defective metabolic reprograming and inefficient effector differentiation of naive T cells. This mechanism provides an exciting opportunity to regulate T cell responsivity in both autoimmunity and T cell-based immunodeficiencies.

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Competing interest statement: U.G. is currently employed by AstraZeneca AB (Mölndal, Sweden).

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Data deposition: The microarray and sequencing data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, https://www.ncbi.nlm.nih. gov/geo (accession no. GSE140448).

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Typically, these differentiation programs are supported by a series of heritable gene expression programs under the control of specific transcription factors. Epigenetic modifications, including histone acetylation and methylation, or DNA methylation, determine the accessibility of genomic regions to transcription factors. For T cell differentiation into effector T cells, for example, trimethylation of lysine 4 on histone 3 (H3K4me3) is a permissive mark for transcription of lineage specific cytokines IFN γ (Th1), IL4 and IL13 (Th2), and IL17 (Th17), while H3K27me3 is associated with their transcriptional repression (3–7). DNA methylation, in turn, is critical for regulating the expression of Foxp3, the hallmark transcription factor of Tregs (8).

Recently, we showed that H3K9 trimethylation, characteristic for transcriptionally inactive heterochromatin domains, at the IFN γ locus is decisive for Th2 lineage commitment (9). Differentiated Th2 cells from mice lacking the histone methyltransferase Suv39h1 or the heterochromatin protein 1 α (HP1 α), readily converted to IFN γ -expressing Th1-like cells, while wild-type (WT) cells did not. HP1 α and its 2 other isoforms, HP1 β or HP1 γ , are recruited to chromatin by a tripartite motif containing, scaffolding protein, TRIM28 (also known as KAP1 or Tif1 β) (10). TRIM28 interacts with KRAB-zinc finger proteins to target specific genomic regions and modulates transcription through interactions with HP1 isoforms (11, 12). Moreover, recent studies suggest that TRIM28, HP1 β , and HP1 γ regulate initiation and/or elongation of RNA polymerase II (Pol II)-dependent transcription (13–15).

Several studies revealed an important role for TRIM28 in T cell development and activation. Mice with T cell-specific deletion of TRIM28 (TRIM28^{-/-}) lack invariant natural killer T cells due to altered use of T cell receptor α (TCR α) chains and develop autoimmunity due to TGF β 3 overexpression and deregulation of the Th17 pathway (16–18). TRIM28^{-/-} mice also have increased Tregs numbers and a defect in IFN γ production in Th1 cells, but the mechanism for this imbalance remains unclear (17). Although it was shown that TRIM28 could interact with both polycomb and heterochromatin effectors in other cell types, it is still unclear which of these 2 epigenetic effector pathways mediates the effects of TRIM28 in T cells (19).

We now show that TRIM28^{-/-} and HP1 $\beta/\gamma^{-/-}$ T cells fail to differentiate into helper and Tregs due to an early defect in metabolic reprograming to glycolysis. Defective metabolic reprograming results from impaired TCR/CD28 signaling through the PI3K-Akt-mammalian target of rapamycin (mTOR) axis. These defects correlate with impaired epigenetic silencing by TRIM28 and HP1 β/γ at distal regulatory regions to a defined set of Treg-characteristic genes, whose silencing is critical to proper T cell expansion and differentiation into helper and regulatory phenotypes.

Results

TRIM28 Is Necessary for Efficient Teff Differentiation and Function In Vivo. As shown previously by others, TRIM28^{flox/flox} × CD4Cre mice (TRIM28^{-/-}) develop phenotypically normal primary and secondary lymphoid organs and display minor alterations in thymocyte development (17, 18) (*SI Appendix*, Fig. S1 *G–J*). TRIM28 expression is normal in double-negative thymocytes and decreases by about 50% in double-positive thymocytes and by almost 100% in CD4⁺ and CD8⁺ peripheral T cells. Lack of TRIM28 expression in peripheral T cells is also visible by Western blot (*SI Appendix*, Fig. S14).

Bone marrow (BM) from $\text{TRIM28}^{-/-}$ mice contains significantly fewer central and effector memory T cells (*SI Appendix*, Fig. S1*B*) and fewer CD44⁺ memory CD4 T cells expressing the proliferation marker Ki67 in peripheral lymph nodes (*SI Appendix*, Fig. S1*C*). Upon sorting of effector memory T cells from spleen and mesenteric lymph nodes of aged mice, we also detected less effector cytokines (IFN γ , IL13, and IL17) after ex vivo restimulation with PMA/ionomycin (*SI Appendix*, Fig. S1*D*). Adoptively transferred TRIM28-deficient OT-II naive T cells proliferated less and produced less effector cytokines after immunization with ovalbumin (OVA) in complete Freund's adjuvant (CFA) (*SI* *Appendix*, Fig. S1*E*). CFA-OVA immunization of TRIM28^{-/-} resulted in significantly fewer IL4-producing OVA-specific CD4⁺ T cells compared to WT controls (*SI Appendix*, Fig. S1*F*).

To test whether this defect in effector T cell function translates to decreased T cell-mediated immunopathology, we studied a T cell transfer colitis model (20). Rag2^{-/-} mice transferred with TRIM28^{-/-} × RorcGFP⁺ naive CD4⁺ T cells survived longer and developed significantly less colitis, as evidenced by the delay in weight loss, decreased T cell infiltration in colonic mucosa, increased colon length, and lower immunohistopathology scores of colonic tissue (Fig. 1 *A*–*E*). Time course analysis of CD4⁺ T cells in the blood revealed a strong defect in T cell expansion and development into Roryt-expressing cells that translated into fewer IFNγ- and IL17-producing splenic CD4⁺ T cells (Fig. 1 *F*–*I*). We conclude that TRIM28 is necessary for efficient effector

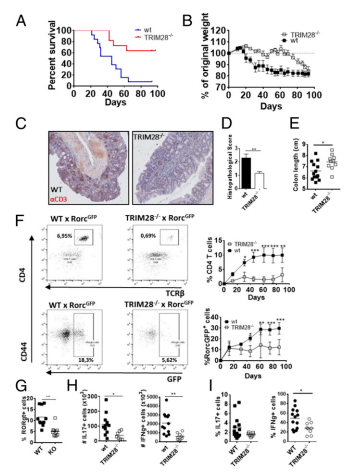


Fig. 1. T cell-mediated immunopathology depends on TRIM28. (A and B) Survival and weight loss (percentage of starting weight) of Rag2⁻ mice after transfer of either TRIM28^{-/-} × RorcGFP or littermate control naive CD4⁺ T cells. (C) Immunohistochemistry of colonic mucosa of Rag2^{-/-} mice injected with either TRIM28^{-/-} or littermate control naive CD4⁺ T cells using α CD3 (brown) to stain for infiltrated T cells. (D and E) Colon length and histopathologic analysis of colonic sections from mice killed in A. (F) Representative dot plots (6w) and quantification of total (upper row) or RorcGFP⁺ (lower row) CD4 T cells in blood of Rag2^{-/-} mice, at different time points after adoptive transfer. (G) Percentage of RorcGFP⁺ CD4 T cells in spleen of $Rag2^{-/-}$ mice at termination. (H and I) Numbers (H) and percentages (I) of IL17- or IFN_γ-producing splenic CD4⁺ T cells after ex vivo restimulation with PMA/ionomycin/brefeldin A. (B and F) Symbols (A and F) or bars (D, E, G, H, and I) show mean \pm SEM. The dots represent individual mice (E, G, H, and I). Student's t test (E, G, H, and I) or 2-way ANOVA (B and F) was used to calculate statistical significance. Data are from 2 independent experiments; **P* < 0.05; ***P* < 0.01; ****P* < 0.001; and *****P* < 0.0001.

T cell development and expansion in vivo and that its absence prevents the development of T cell-mediated pathology.

TRIM28^{-/-} Chimeras Develop Severe Autoimmunity due to Deficient Treg Expansion. The aforementioned results seem inconsistent with previous studies showing that $TRIM28^{-/-}$ mice develop autoimmunity (18). While $TRIM28^{-/-}$ mice from our animal facility only develop mild symptoms of autoimmunity upon aging (SI Appendix, Fig. S1 G-P), TRIM28^{-/-} BM chimeras suffered from wasting disease and displayed significant weight loss starting 5 wk after reconstitution (Fig. 24), confirming the development of autoim-munity. TRIM28^{-/-} chimeras generated from T cell-depleted BM or isolated BM T cells developed similar symptoms, indicating that autoimmunity is not solely due to expansion of preexisting, autoreactive BM T cells (SI Appendix, Fig. S2A). Weight loss was not observed in mixed BM chimeras that were reconstituted with as little as 25% of WT BM (Fig. 2*B*), suggesting that the pheno-type in the TRIM28^{-/-} chimeras could be due to defective expansion or differentiation of TRIM28^{-/-} Tregs. Indeed, in these mixed chimeras, regulatory T cells derived mainly from WT BM precursors, while effector T cells developed in similar proportions from WT and TRIM28^{-/-} precursors (Fig. 2 C and D). In addition, TRIM28^{-/-} chimeras reconstituted with T cell-depleted BM or \dot{T} cells isolated from BM, harbored significantly fewer splenic Tregs, while the effector compartments developed similarly or were slightly increased, compared to WT chimeras (SI Appendix, Fig. S2B). Also, fewer Tregs from TRIM28^{-/-} mice expressed the proliferation marker Ki67 compared to WT controls (SI Appendix, Fig. S2C).

To test whether the observed autoimmunity is a consequence of defective Treg expansion or loss of Treg stability, we transferred CD45.2⁺ WT or TRIM28^{-/-} Tregs alongside CD45.1⁺ naive T cells into Rag2^{-/-} mice. TRIM28^{-/-} Treg numbers were significantly reduced following adoptive transfer (SI Appendix, Fig. S2 D and E). This was not due to loss of Treg phenotypic stability as ex vivo isolated Tregs maintained higher Foxp3 expression upon redifferentiation in Th17 effector cytokine conditions (SI Appendix, Fig. S2F). Thus, failure of TRIM28^{-/-} Treg to reconstitute the Treg compartment was likely due to an expansion defect. Finally, while WT and TRIM28-/ Tregs performed similarly in an in vitro suppression assay (SI Appendix, Fig. S2G), only WT, but not TRIM $28^{-/-}$, Tregs suppressed autoimmunity when transferred to TRIM $28^{-/-}$ BM chimeras at the time of reconstitution (Fig. 2E and SI Appendix, Fig. S2H). We conclude that TRIM28 is required for both effector and regulatory T cell expansion and that defective Treg expansion, and not their functional differentiation, is likely responsible for the development of autoimmunity in TRIM28^{-/-} mice.

TRIM28 Is Required for Metabolic Reprogramming Early during Teff Differentiation. In order to delineate the underlying mechanisms for the observed phenotypes, we differentiated naive CD4⁺ T cells into Teff in vitro. TRIM28 deficiency impairs effector cytokine production in differentiated Th1, Th2, Th17 cells after PMA/ ionomycin restimulation (Fig. 3 A and B and SI Appendix, Fig. S3 *A–C*). Deficient IFN γ , IL13, and IL17 production was not due to impaired proliferation, as TRIM28^{-/-} cells having undergone the same number of cell cycles still produced less effector cytokines (SI Appendix, Fig. S3D). We also observed a significant increase in Foxp3-expressing Th1 cells in vitro (Fig. 3A and B) and increased percentages of Foxp3-expressing cells in vivo upon injection of a 1:1 ratio of CD45.1 WT and CD45.2 TRIM28^{-/-} T cells into Rag2^{-/-} mice (SI Appendix, Fig. S3E). Defective cytokine production and Foxp3 overexpression were both cell intrinsic since TRIM28⁻ - naive T cells from mixed BM chimeras showed the same proliferation and priming defects (SI Appendix, Fig. S3F). Addition of TGF_{β3} to WT T cells, which was previously shown to be overexpressed in TRIM $28^{-/-}$ T cells, did not reproduce the TRIM28 phenotype (*SI Appendix*, Fig. S3G and ref. 18). Consistently, blocking aTGFB antibodies did not rescue cytokine production in knockout TRIM28^{-/-} cells (SI Appendix, Fig. 3H).

Therefore, TRIM28^{-/-} T cells bear an intrinsic defect that impairs effector differentiation.

To identify the transcriptional pathways deregulated in TRIM28^{-/-} Th1 cells, we analyzed the transcriptomes of in vitro differentiated Th1 cells using Affymetrix microarrays. Genes involved in cell cycle progression and in metabolism were significantly downregulated in TRIM28^{-/-} T cells, compared to WT littermates (Fig. 3C). Anabolic pathways, such as "Cholesterol Biosynthesis" or "Glucose Transport and Metabolism," were down-regulated in TRIM28^{-/-} Th1 cells, compared to control littermates (*SI Apppendix*, Fig. S3*I*). The only metabolic RNA category up-regulated in TRIM28^{-/-} over WT Th1 cells was mitochondrial β -oxidation (*SI Appendix*, Fig. S3*I*). The switch to glucose metabolism is a hallmark of effector T cell differentiation, while naive T cells and Tregs generate their energy mainly through β -oxidation of fatty acids (1). These results suggest that the defect in effector function observed in TRIM28^{-/-} T cells results from an impaired switch from oxidation to glycolysis.

To investigate whether TRIM28 interferes with effector function in already differentiated effector T cells or early during T cell differentiation, we used naive T cells from TRIM28^{flox/flox} or TRIM28^{wt/M} mice crossed with tamoxifen-inducible RosaCreERT2 mice. Addition of tamoxifen in vitro, at the time of naive T cell activation, resulted in strong TRIM28 depletion in effector T cells (after 5 d of differentiation). TRIM28-depleted effector T cells, however, showed no defect in cytokine production, while Foxp3 expression was still increased in Th1 cells (Fig. 3*E*). Therefore, impaired effector cytokine production in T cells requires early TRIM28 depletion, and cannot be obtained by delayed depletion, while the effect on Foxp3 follows TRIM28 depletion, even if it occurs at later stages (suggesting a more direct effect of TRIM28 on silencing of FoxP3 during effector differentiation).

Loss of TRIM28 Leads to Defective mTOR Signaling and Glycolysis in Naive and Regulatory T Cells. To investigate whether the defect in glycolytic switch is due to defective T cell activation after TCR engagement, we first measured proliferation and blasting after stimulation of naive CD4⁺ T cells with α CD3/ α CD28. TRIM28⁻ cells displayed a significant delay in proliferation and blasting starting 24 h after stimulation (as defined by forward-scatter acquisition) (Fig. 4A and B). Transcriptomic analysis and ¹³C-glucose tracing experiments revealed deregulation of glucose (and lipid) metabolisms only in activated, not naive TRIM28^{-/-} CD4 T cells (Fig. 4C and *SI Appendix*, Fig. S4 A–C). In activated TRIM28^{-/-} T cells, ¹³C-labeled carbons accounted for a significantly smaller fraction of glycolysis-dependent metabolites such as lactate, serine, methionine, citrate, and succinate, compared to WT T cells (Fig. 4C). Bioenergetics measurements revealed a significant defect in glycolysis in TRIM28^{-/-} T cells after 24 h (but not 4 or 0 h) of activation, compared to cells from control littermates, while mitochondrial respiration remained unchanged (Fig. 4D and SI Appendix, Fig. S4 \hat{D} and E). Defective glycolysis at 24 h was not due to increased presence of Foxp3⁺ cells (SI Appendix, Fig. S4F). Since glucose metabolism is mainly regulated via the mTOR pathway, we measured S6 and 4-EBP1 phosphorylation as surrogate markers of mTOR activ-ity. As expected, activated TRIM28^{-/-} T cells exhibited lower levels of 4-EBP1 and S6 phosphorylation, both in Foxp3⁺ and Foxp3⁻ T cells (Fig. 4 E and F and SI Appendix, Fig. S4 Gand H).

In recent years, mTOR has emerged as a central signaling hub that distinguishes effector from regulatory T cell differentiation by regulating key metabolic pathways, such as glycolysis (21). Importantly, while the absence of mTOR activity is important for Foxp3 expression and iTreg differentiation, its activity is necessary for the expansion of Tregs and maintenance of immune homeostasis in vivo. Tregs from TRIM28^{-/-} mice exhibited reduced glycolysis and lactate production 24 h after α CD3/ α CD28 activation, coinciding with a decrease in α CD3/ α CD28 and IL2-dependent S6 phosphorylation and proliferation (Fig. 4 *G* and *H*). Together, these results suggest that the absence of TRIM28 impairs

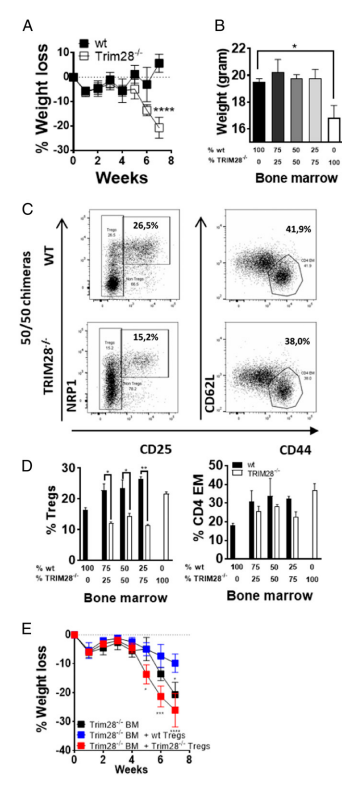


Fig. 2. TRIM28 is required for Treg expansion and suppression in vivo. (A) Weight (as percent starting weight) of $Rag2^{-/-}$ mice irradiated and reconstituted with BM from WT or TRIM28^{-/-} mice. (B) $Rag2^{-/-}$ BM chimeras were reconstituted with the indicated percentages of WT and TRIM28^{-/-} BM and weighed 6 wk after reconstitution. (C and D) Representative dot plots (C) and quantification (D) of splenic Tregs and CD4 EM cells of mice from (B). Data are presented as percentage of CD45.1 (WT) or CD45.2 (TRIM28^{-/-}) CD4⁺ T cells. (E) Weight (as percent starting weight) of Rag2^{-/-} mice. (A-E) Data are from 2 independent experiments with 4 to 7 mice per group. Symbols (A and E) or bars (B and D) show

mTOR signaling in naive and regulatory CD4 T cells and thereby interferes with effector T cell differentiation and Treg expansion.

Defective Signaling through CD28 in TRIM28^{-/-} **T Cells.** To explain the decreased mTOR activity in TRIM28^{-/-} T cells, we investigated signaling events proximal to TCR/CD28 stimulation. Metabolic switch to glycolysis in effector T cells is mainly driven by CD28-PI3K-AKT-mTOR signaling axis (22). Expression of CD28, but not CD3, was slightly, but significantly, reduced in TRIM28^{-/-}, compared to control T cells (*SI Appendix*, Fig. S5A). To test whether defective signaling through CD28 may explain the mTOR defect, we measured the response of WT and TRIM28^{-/-} CD4⁺ T cells to increasing concentrations of aCD28 (leaving the concentration of aCD3 unchanged). While S6 phosphorylation, T cell blasting (FSC), and IFNy production were similar in WT and TRIM28 cells stimulated with α CD3 alone, increasing concentrations of aCD28 led to a significant increase in pS6, FSC, and IFNy production in activated CD4⁺ T cells from WT, but not from TRIM28^{-/-} mice (Fig. 5 *A*–*C*). In contrast, increased CD28 en-gagement in TRIM28^{-/-}, but not in WT Th1 cells, resulted in up-regulation of Foxp3 expression (Fig. 5 *D*, *Left*). Foxp3 expression was further increased by inhibition of PI3K by Ly294002 (Fig. 5 D, Right), suggesting that CD28-dependent Foxp3 induction is achieved through an alternative, PI3K-independent pathway. In contrast, we did not observe any differences in Ca^{2+} signaling between WT and TRIM28^{-/-} T cells following activation with $\alpha CD3/\alpha CD28$ or ionomycin (*SI Appendix*, Fig. S5*B*). The signaling defect was limited to stimulation through CD28, as T cell activation, S6 phosphorylation, and blasting were rescued by stimulation with PMA/ionomycin (SI Appendix, Fig. S5C). Western blot analysis of downstream signaling molecules revealed that phosphorylation of the Akt substrate Foxo1/3a, but not that of ERK1/2, downstream of CD3 ϵ signaling, was impaired in TRIM28^{-/-}, compared to WT T cells, also consistent with a defect in signaling through the CD28-PI3K-AKT-mTOR axis (Fig. 5 E and F). PKC θ recruitment is a hallmark difference in CD28 signaling between Tregs and Tconv (23). Strikingly, α CD3/ α CD28 stimulation led to decreased recruitment of PKC θ to the immune synapse in TRIM28^{-/-} compared to WT cells (SI Appendix, Fig. S5 D and E).

Together, these results indicate that, while CD28 is active in both WT and TRIM28^{-/-} T cells, CD28 signals through alternative pathways in TRIM28^{-/-} T cells, leading to Foxp3 expression rather than mTOR activation. The results also show that TRIM28^{-/-} naive T cells present differences in precocious events of activation, minutes after TCR engagement, when transcriptional or epigenetic regulation events are unlikely to have occurred. We therefore explored the possibility that the observed phenotype is due to epigenetic deregulation in naive T cells before they are activated.

TRIM28 Deficiency Reactivates Silent Regulatory Elements in Naive T Cells through H3K9 Histone Modifications. To investigate possible differences in gene expression between naive WT and TRIM28^{-/-} cells, we analyzed their respective transcriptomes by means of Affymetrix microarrays. In total, 222 RNA species were significantly up-regulated, and 76 are down-regulated in naive TRIM28^{-/-} T cells, compared to WT naive T cells (Fig. 6.4). Differences in gene expression were statistically significant, but small (ranging between 1.2- and 10-fold). Three times more genes were up- than down-regulated, consistent with the known role of TRIM28 as a transcriptional corepressor (11).

TRIM28 functions as a molecular adaptor for gene silencing through both the polycomb and the HP1 pathways. HP1-mediated control of gene expression depends on H3K9 trimethylated (H3K9me3) and acetylated (H3K9Ac) epigenetic marks, which are associated with stable silencing and active

mean \pm SEM. Two-way ANOVA (*A* and *E*), one-way ANOVA (*C*), or Student's *t* test (*D*) was used to calculate statistical significance. **P* < 0.05; ***P* < 0.01; ****P* < 0.001; and *****P* < 0.0001.

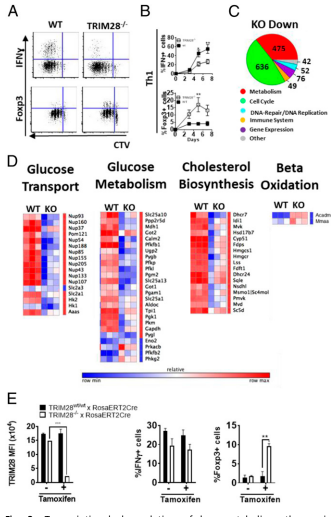


Fig. 3. Transcriptional deregulation of key metabolic pathways in TRIM28^{-/-} T cells. (A and B) Dot plots and quantification showing Cell Trace Violet (CTV) dilution (A), IFN_Y (A and B) and Foxp3 (A and B) expression in differentiated TRIM28-/- or littermate control Th1 at day 5 of culture after restimulation. Data are from 6 experiments and 12 mice per time point. (C) Pie chart of pathways enriched in genes with decreased expression levels in Th1 TRIM28^{-/-} cells compared to littermate control cells. Numbers of deregulated genes are indicated in the corresponding plot section. (D) Heatmaps and hierarchical clustering of metabolic pathways showing genes differentially regulated in Th1 cells from TRIM28^{-/-} or littermate control mice. Coloring corresponds to relative expression values, normalized for each gene. Data are generated from 3 TRIM28-/- and matching littermate control mice. (E) Quantification of TRIM28 expression, IFNy production, and Foxp3 expression after Th1 differentiation of naive T cells from TRIM28^{+/+} or TRIM28^{-/-} \times Rosa26ERTCre mice in the presence or absence of tamoxifen, at day 5, following restimulation. Data are from 2 independent experiments. (A and E) Symbols (A) or bars (E) show mean \pm SEM, and 2-way ANOVA was used to determine statistical significance. *P < 0.05; **P < 0.01; ***P < 0.001.

transcription, respectively (24). To explore whether the observed transcriptional changes would be correlated to changes in H3K9 histone modifications, we performed chromatin immunoprecipitation sequencing (ChIP-seq) for H3K9me3, H3K9Ac on naive CD4⁺ T cells from TRIM28^{-/-} and WT mice. We observed a significant correlation between increased acetylation and decreased trimethylation 20 to 50 kB around differentially regulated genes (Fig. 6 *B* and *C*). Increase in acetylation and decrease in trimethylation positively correlated with changes in gene expression in TRIM28^{-/-} T cells compared to WT littermates. H3K9Ac was significantly increased on 51 promoters

and 347 distal regulatory elements (defined based on a classical H3K4me1/H3K27ac signature) around the differentially regulated genes (*SI Appendix*, Fig. S6A), indicating a more pronounced effect of TRIM28 depletion on distal regions and thus on putative active enhancers, compared to promoters (Fig. 6*C*; $P < 10^{-5}$, Fisher test).

RNA-seq and ChIP-seq for RNA Pol II revealed an increased transcription at H3K9-hyperacetylated distal regions in TRIM287 compared to WT CD4⁺ T cells (SI Appendix, Fig. S6B). This is exemplified at the TNFRII, Snai3/Rnf166, and NRP1 loci, which showed increased H3K9ac signals at a distal, regulatory upstream regions (E) that also correlated with decrease of the H3K9me3 signal (Fig. 6D and SI Appendix, Fig. S6C). Reanalysis of a published ChIP-seq dataset using TRIM28-specific antibodies on thymocytes revealed significant enrichment of TRIM28 peaks around transcriptionally deregulated genes (SI Appendix, Fig. S6D). We further confirmed the presence of TRIM28 at transcriptionally deregulated genes using TRIM28-specific ChIP-PCR and primers designed for TRIM28 peaks identified in SI6C, negative and published positive controls (SI Appendix, Fig. S6E and refs. 17 and 18). Taken together, these results suggest that TRIM28 regulates the levels of acetylation vs. trimethylation of H3K9 at a selective set of distal regulatory elements (and promoters) of genes that are up-regulated in TRIM28-defective cells.

To investigate the nature of the link between this set of deregulated genes and the phenotype observed in TRIM28^{-/} T cells, we used pathway and transcription factor binding site analysis. While we did not detect any significant gene enrichment for published pathways and GO terms among differentially deregulated genes, we did detect a significant enrichment of binding sites for Foxo1, a transcription factor strongly associated with Tregs and metabolic regulation (25-28) (Fig. 6E and SI Appendix, Fig. S6 F and G). Gene set enrichment analysis (GSEA) revealed that a significant proportion of differentially expressed genes overlap with a published Treg signature (Fig. 6 \dot{E} and F and ref. 29). Increased RNA levels of Tnfrsf1b, Folr4, and Nrp1 coincided with increased surface expression levels for TNFR II, FR4, or NRP1 on naive CD4⁺ T cells (SI Appendix, Fig. S6H). Also, a similar increase in FR4 and NRP1 was observed on naive TRIM28^{-/-} T cells and CD4⁺ single- and double-positive thymocytes from mixed BM chimeras, indicating a direct, cellintrinsic effect of TRIM28 on these genes during thymic development (SI Appendix, Fig. S6I). These results suggest that a defect in epigenetic silencing of a set of Treg-associated genes alters early T cell activation, causing defective downstream mTOR signaling and effector differentiation.

TRIM28 Regulates T Cell Activation and Differentiation through Treg-Associated Genes and Akt. NRP1 has been shown to interfere with T cell activation by recruiting the phosphatase PTEN to the immunological synapse (30). To test whether thymic epigenetic silencing of NRP1 is required for activation and proliferation of peripheral T cells, we overexpressed NRP1 alongside Ametrine, a fluorescent reporter gene in an OVA-specific T cell hybridoma and activated the transduced cells via OVA-pulsed dendritic cells. NRP1-overexpressing cells proliferated less in response to dendritic cell-mediated activation compared to the Ametrine reporter transfection control, as evidenced both by less CFSE dilution and lower cell numbers up to 12 d posttransfection, relative to the Ametrine-only control (Fig. 7 A-C). To further explore the contribution of overexpressed Treg genes to TRIM28-associated defects, we first sought to block Nrp1 function in TRIM28^{-/} naive T cells. Using blocking antibodies against Nrp1 and its ligand semaphorin4a or CRISPR/Cas9-mediated gene ablation, we found IL17, but not IFNy production was partially restored by blocking the Nrp1-signaling axis (Fig. 7 D and \tilde{E}). In contrast, CRISPR/Cas9 targeting of Snai3 increased both IFNy and IL17 production. CRISPR/Cas9-mediated knockdown led to a 40 to 50% decrease in Nrp1 surface levels as assessed by flow cytometry (SI Appendix, Fig. S7A) and 30 to 50% Snai3 frameshift mutations (SI Appendix,

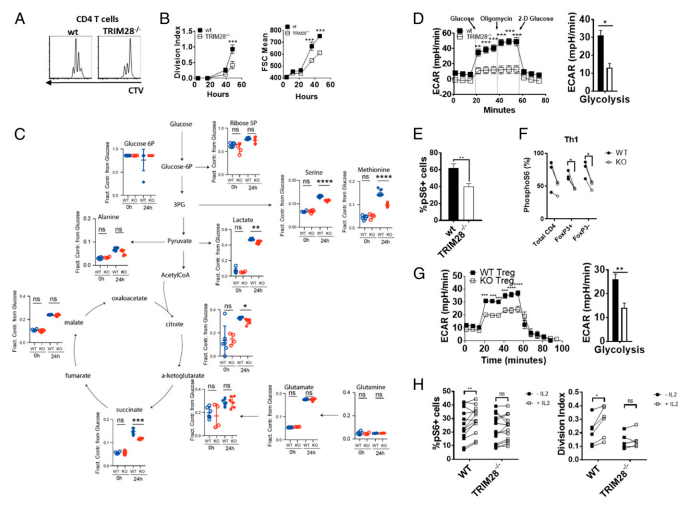


Fig. 4. TRIM28 regulates the metabolic switch to glycolysis in naive and regulatory CD4⁺ T cells following TCR activation. (*A* and *B*) Histogram plots of Cell Trace Violet (CTV) dilution (*A*), division index (*B*, *Left*), or FSC (*B*, *Right*; used as a measure of T cell blasting), 48 h (*A*) or at different time points (*B*) after stimulating TRIM28^{-/-} or littermate control naive CD4⁺ T cells with α CD3/ α CD28. (*C*) Graphic representation of glucose-dependent metabolic pathways and scatter plots of ¹³C incorporation in indicated metabolites in TRIM28^{-/-} or littermate control CD4⁺ activated for 24 h using α CD3/ α CD28. Data are from biological replicates of 1 to 2 mice per group. (*D* and G) Quantification of the extracellular acidification rate (ECAR) of TRIM28^{-/-} or littermate control naive CD4⁺ (*D*) or Tregs (*G*) activated for 24 h using α CD3/ α CD28 (*D*), in presence or absence of IL2 (*G*) as measured by the Seahorse XFe96 Analyzer. (*E* and *F*) Quantification of ribosomal S6 phosphorylation 24 h after activation of naive CD4⁺ T cell from TRIM28^{-/-} or littermate control mice with α CD3/ α CD28 in total cells (*E*) or Foxp3⁻ vs. Foxp3⁺ cells from cells stimulated in Th1 conditions (*F*). (*H*) Quantification of ribosomal S6 phosphorylation at 24 h and division index 48 h after activation of CD4⁺ α CD3/ α CD28 in presence of absence of IL2. (*A*, *B*, *D*, and *H*) Data are from 2 independent experiments. Two-way ANOVA (*B*, *D*, and *G*), Student's *t* test (*C* and *E*), or paired *t* test (*F* and *H*) was used to determine statistical significance. **P* < 0.05; ***P* < 0.01; ****P* < 0.001.

Fig. S7B) as assessed by tracking of indels by decomposition, as described previously (31).

Since NRP1 has been shown to recruit PTEN, an antagonist of the PI3K-Akt pathway, we asked whether overexpression of constitutively active Akt in TRIM28-/- CD4 T cells would rescue effector T cell differentiation. Following transduction of activated naive CD4 T cells with a construct coding for myristoylated Akt and subsequent differentiation into Th1, Th2, or Th17 effector cells, TRIM28^{-/-} Th1 cells produced significantly more IFNy (Th1) and IL17 (Th17) than the respective transduction control (Fig. 7 F and G and SI Appendix, Fig. S7C). Cytokine levels reached those of control vector transduced WT Th1 and Th17 cells (Fig. 7 F and G and SI Appendix, Fig. S7C). Similar findings were made for Th2 cells, although differences did not reach statistical significance (SI Appendix, Fig. S7D). Increased cytokine responses in Th1 and Th17 cells were dependent on glucose concentrations in the medium suggesting that myr-Akt overexpression rescues effector T cell differentiation by restoring glycolysis (SI Appendix, Fig. S7E). We conclude that TRIM28 permits effector T cell differentiation due to epigenetic silencing of genes that interfere with the Akt signaling pathway. In the absence of TRIM28, restoring the Akt pathway can overcome the TRIM28-induced hyporesponsiveness in a glucose-dependent manner.

TRIM28 Requirement for Effector T Cell Differentiation Is Dependent on HP1. Since transcriptional changes in TRIM28^{-/-} T cells correlated with changes in H3K9 histone modifications, we investigated whether TRIM28 needs to interact with the H3K9me³-binding HP1 isoforms (α, β, and γ) to control the induction of glycolysis and effector/regulatory fates. We first analyzed T cell effector differentiation in mice harboring a mutated version of TRIM28 (V488L490/AA), which impairs interactions with all HP1 isoforms α, β, or γ (32). TRIM28^{ΔHP1box/flox} × CD4Cre (TRIM28^{ΔHP1box/-}) mice express normal levels of the mutated TRIM28 protein (*SI Appendix*, Fig. S8*A*). While TRIM28^{ΔHP1box/-} mice had phenotypically normal spleens, thymus, and T cell compartments (*SI Appendix*, Fig. S8 *B–D*, *Left*), CD4⁺ T cells from TRIM28^{ΔHP1box/-} mice

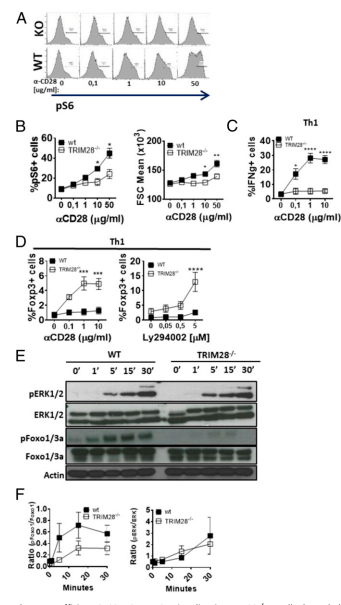


Fig. 5. Inefficient CD28–PI3K–mTOR signaling in TRIM28^{-/-} T cells. (A and B) Representative profiles (A) and guantification of ribosomal S6 phosphorylation (B) and blasting (B) 24 h after activation of TRIM28^{-/-} or littermate control naive CD4⁺ T cells with α CD3 and increasing concentrations of α CD28. (C) IFN γ expression on day 5 of Th1 cell cultures from TRIM28^{-/-} or littermate naive CD4⁺ T cells in the presence of increasing doses of aCD28. (D) Foxp3 expression on day 5 of Th1 cell cultures from TRIM28^{-/-} or littermate control mice in the presence of increasing doses of aCD28 or PI3K inhibitor Ly294002. (E and F) Representative Western blots (E) and quantification of band intensities (F) using cell lysates from TRIM28^{-/-} or littermate control naive CD4⁺ T cells activated for up to 30 min using aCD3 and aCD28. (A-D) Data are from 2 independent experiments, and 2-way ANOVA was used to calculate statistical significance. *P < 0.05; **P < 0.01; ***P < 0.001; and ****P < 0.0001.

displayed the same defects in proliferation, blasting, and IFNy and Foxp3 as TRIM28^{-/-} cells (Fig. 8 A, Left, and SI Appendix, Fig. S8 E and F). These results indicate that TRIM28–HP1 interactions are critical to TRIM28 functions in T cells.

To identify which HP1 isoforms are recruited by TRIM28, we used mice lacking either one or a combination of 2 HP1 isoforms $(\alpha, \beta, \text{ and } \gamma)$. Spleens and thymi of HP1-deficient mice were phenotypically normal (SI Appendix, Fig. S8 B-D and ref. 9). Neither single HP1 isoform defects, nor combined deletion of HP1 α and β , affected T cell activation, blasting, or differentiation

(Fig. 8 A, Right, and SI Appendix, Fig. S8 E and F). Simultaneous disruption of HP1 β and HP1 γ , however, reproduced all of the observed phenotypes of TRIM28^{-/-} or TRIM28^{Δ HP1box/-} T cells (Fig. 8 A, Right). An impaired glycolytic switch, reduced blasting, and decreased mTOR activity were all observed in HP1 $\beta/\gamma^{-/-}$, but not HP1α-KO T cells (Fig. 8B and SI Appendix, Fig. S8 E and F). We also detected a similar increase in Treg signature genes, in both TRIM28^{Δ HP1box/-} and HP1 $\beta/\gamma^{-/-}$ but not HP1 $\alpha^{-/-}$ T cells, both at the RNA and protein levels (Fig. 8 *C* and *D* and *SI Appendix*, Fig. 88 *G* and *H*). Finally, HP1 $\beta/\gamma^{-/-}$ or TRIM28^{Δ HP1box/-} BM chimeras developed autoimmune symptoms (similar to TRIM28^{-/-} chimeras), as indicated by severe weight loss and a lack of splenic Tregs (Fig. 8 E and F). We conclude that effector T cell differentiation and Treg expansion both require the interaction of TRIM28 with HP1 β/γ isoforms, establishing a direct link between the observed phenotypes and heterochromatin dynamics.

Discussion

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Over the last few years, it has become clear that T cell functions are linked to specific modes of cell metabolism. While naive and memory T cells, similar to Tregs, rely mainly on oxidative respiration, differentiation to effector helper T cells, capable of active expansion and secreting high levels of cytokines, requires a "switch" to glycolysis. This glycolytic switch depends on gene expression reprograming that occurs after TCR/CD28-mediated activation of the mTOR pathway. In this study, we unravel an unexpected level of epigenetic control of metabolic reprograming by heterochromatin-dependent gene silencing. TRIM28 or $HP1\beta/\gamma$ expression in early differentiating T cells represents an absolute requirement for effective switch of CD4⁺ T cells to glycolysis. In the absence of the heterochromatin regulators, the balance between H3K9me3 and H3K9ac is modified at regulatory enhancers at a series of genes, causing their overexpression and inhibiting glycolytic reprograming.

It is unclear at this point which genes are direct targets of TRIM28-HP1 β/γ regulation, and which genes are overexpressed as a consequence of indirect effects, as TRIM28 ChIP-seq proved to be irreproducible (in our hands) and unconvincing (in published datasets). It is also unclear whether the defective glycolytic switch is due to deregulation of a complex combination of genes, or to one individual gene among this list. While decreased H3K9me3 levels at some gene regulatory regions were decreased in TRIM28^{-/-} cells, neither TRIM28 nor HP1 possess H3K9 trimethylation activity, suggesting that the TRIM28-HP1 β/γ complex recruits a histone-methyltransferase (HMT) of H3K9. HP1 proteins can recruit the HMTs Suv39h1 or Suv39h2, while TRIM28 can recruit another H3K9 HMT, SETDB1 (33-35). Our preliminary results indicate that neither Suv39h1- nor Suv39h2-defective T cells phenocopy the observed defect in effector T cell differentiation or the increased expression of the Treg signature. It is therefore most likely that other H3K9 trimethylases, such as SETDB1, are involved.

If TRIM28 is necessary for the full development of effector T cell responses, how then can TRIM28-deficient mice develop autoimmunity [as reported by others (18)]? The authors suggested a defect in the suppressive activity of TRIM28-defective Tregs, even though they did not detect differences in Treg-suppressive activity in vitro. Our results indicate that Tregs from WT and TRIM28^{-/-} mice were equally suppressive in vitro (*SI Appendix*, Fig. S2F). Tregs from TRIM28^{-/-} mice, however, did not respond to IL2 and failed to expand in vivo to suppress colitis as well as in pure and mixed BM chimeras (Fig. 2C and SI Appendix, Figs. S2 B, D, and E and S4F). Our data further suggest that this expansion defect is due to a defect in mTOR signaling and glycolytic switch (Fig. 4 E and F), which was recently suggested to be necessary for Treg expansion and thereby for their suppressive capacity in vivo (36). Thus, in addition to its role in regulating effector T cell responses, TRIM28 also regulates the expansion of regulatory T cells. A partial defect in Teff development, associated to a strong defect in Treg expansion, most likely accounts for the autoimmune

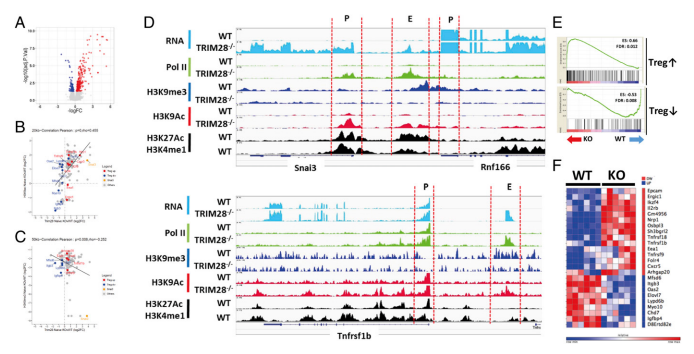


Fig. 6. Naive TRIM28^{-/-} T cells display a Treg-like signature. (*A*) Volcano plot of gene expression in naive TRIM28^{-/-} or littermate control CD4⁺ T cells. Differentially expressed genes (P < 0.01) are colored blue (down in KO) and red (up in KO). (*B*) Correlation dot plot using log₂ FC values for gene expression in TRIM28^{-/-} over WT naive CD4 T cells (*x* axis) and H3K9 acetylation in a 20-kb window around the transcription start site (TSS) of differentially expressed genes (*y* axis). Correlation was calculated using the Pearson method, and trend line is indicated in black. (*C*) Correlation dot plot using log₂ FC values for gene expression in TRIM28^{-/-} over WT naive CD4 T cells (*x* axis) and H3K9 trimethylation in a 50-kb window around the TSS of differentially expressed genes (*y* axis). Correlation was calculated using the Pearson method, and trend line is indicated in black. (*C*) Expressed in TRIM28^{-/-} over WT naive CD4 T cells (*x* axis) and H3K9 trimethylation in a 50-kb window around the TSS of differentially expressed genes (*y* axis). Correlation was calculated using the Pearson method, and trend line is indicated in black. (*D*) Examples of epigenetic and transcriptional perturbations associated with presence or absence of Trim28 at promoters and enhancers at the Trim24 and Snai3 loci. The labels of ChIP-seq tracks are indicated on the *Left*. P, P1, and P2 represent promoters, and E, putative enhancers, in which gain of H3K9ac active mark and loss of H3K9me3 are observed. (*E*) Gene set enrichment analysis (GSEA) using the transcriptome of naive TRIM28^{-/-} or littermate control CD4⁺ T cells in comparison to published Treg† and Treg‡ signatures. (*F*) Heatmap of Treg signature genes differentially expressed in KO T cells.

phenotype observed by others in TRIM28-deficient mice, and by us in BM chimeras (17, 18).

One of the most striking differences between WT and TRIM28 KO T cells is signaling downstream of CD28. While stimulation with aCD28 induced efficient mTOR phosphorylation and Th1 differentiation in WT T cells, it resulted in increased Foxp3 induction in TRIM28^{-/-} T cells, even under Th1 priming conditions (Fig. 4A). Foxp3 induction was antagonized by PI3K, as pharmacological inhibition of this kinase further increased Foxp3 expression in KO T cells, while inhibiting IFNy expression in WT Th1 cells (Fig. 4F). Therefore, CD28 activation can lead to at least 2 different signaling pathways: one inducing PI3K-Akt-mTOR activity and favoring effector T cell differentiation, the second one leading to Foxp3 expression independently of PI3K. It is interesting to notice that TRIM28 and even HP1 have been implicated as targets for phosphorylation after TCR or IL2 activation (16, 18, 37, 38). TRIM28 can also be phosphorylated downstream of Grb2, an adaptor for the intracellular domain of CD28 (39). Thus, TRIM28 acts proximal to TCR signaling, possibly through chro-matin remodeling. Interestingly, Sauer et al. (40) described a TGFβindependent pathway of Foxp3 induction that involved chromatin remodeling. The authors demonstrated that premature termination of CD3/CD28 signaling leads to Foxp3 expression in around 10% of activated cells and could further be augmented by inhibition of mTOR and PI3K. Premature termination of signaling also led to increased levels of H3K4me2/3 marks and chromatin accessibility at the promoter and 5'-untranslated region of Foxp3 and other Tregassociated genes (40). Altogether, the results suggest that, in addition to controlling the expression of critical genes in naive T cells, TRIM28 is also necessary for chromatin remodeling at enhancers of Treg-associated genes upon TCR engagement.

Previous studies suggested that hyperresponsivity to TGFβderived signals causes the bias toward Treg cell differentiation (17). While we confirm the overexpression of TGF β 3 mRNA, especially in Th17 cells, we found that deficient effector cell differentiation is a T cell-intrinsic defect and could not be restored by blocking TGF β present in the culture (*SI Appendix*, Fig. S1 *D* and *E*). Furthermore, the transcriptomic Treg signature was enriched in TCR signaling- or Foxp3-dependent genes, but not in genes related to TGF β signaling. Interestingly, we could not detect any changes in H3K9me3, H3K9Ac, or Pol II proximal to the Foxp3 gene (*SI Appendix*, Fig. S7*C*), and its expression was not significantly changed between WT and KO naive T cells up to 24 h postactivation (*SI Appendix*, Fig. S4*F*).

These results also suggest that TRIM28 is an important epigenetic regulator during thymic development. Zhou et al. (16) showed that TRIM28 intervenes in TCR rearrangement, and Chikuma et al. (18) found severe defects in thymus development when TRIM28 was deleted at the CD4⁻/CD8⁻ stage. Disruption of TRIM28 at the CD4⁺/CD8⁺ stage of thymocyte differentiation (in our case) does not lead to abnormal T cell development. However, silencing of Treg-associated genes, such as NRP1, already occurs in the thymus and might be a prerequisite for the development of naive CD4⁺ T cells with the full potential to differentiate into Treg or Teff in the periphery. Supporting this notion, overexpression of NRP1 in a T cell hybridoma strongly reduced the proliferative response to antigenic stimulation by dendritic cells (Fig. 7A-C), while Nrp1 inactivation in naive TRIM28^{-/-} T cells partially restored cytokine secretion. Interestingly, genetic ablation of the Tregassociated genes Nrp1 or Snai3 had different outcomes. Indeed, Snai3 deletion partially restores cytokine secretion both in Th1 and Th17 conditions, while Nrp1 inactivation only had an effect on IL17 production. This suggests that the different Treg-associated genes overexpressed in TRIM28^{-/-} T cells contribute by various mechanisms to the observed defects. Full phenotype restoration is thus likely to depend on targeting multiple genes simultaneously.

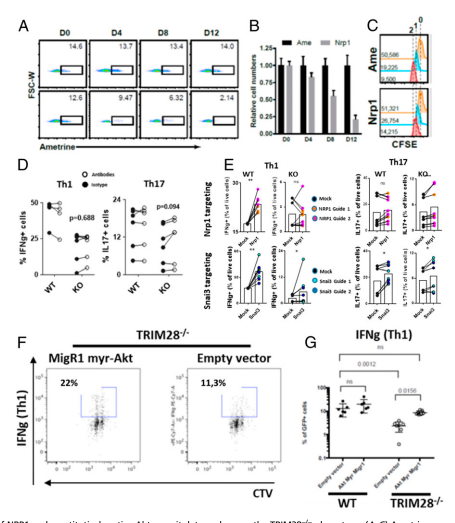


Fig. 7. Overexpression of NRP1 and constitutively active Akt recapitulate and rescue the TRIM28^{-/-} phenotype. (A–C) Ametrine expression (A) and proliferation (B and C) of Ametrine-reporter gene⁺ B097 (hybridoma) T cells transduced to express Ametrine (fluorescent reporter) or Nrp1-2A-Ametrine. (A) Ametrine expression from day 2 to day 12 posttransduction. (B) Relative cell numbers based on Ametrine-positive cells presented in A. Data are normalized based on Ametrine-positive cells in the Ametrine-only condition. (C) CFSE dilution over the course of the first 48 h. Days following CFSE labeling and CFSE mean fluorescence intensity are indicated. (D) Percentage of IFN₇⁺ TRIM28^{-/-} or littermate control Th1 cells following differentiation in presence of NRP1 and Sma4a or isotype control antibiodies. Data are from 2 independent experiments; dots represent individual mice, and paired Wilcoxon test was used to test for statistical significance. (E) Percentage of IFN₇⁺ or IL17⁺ TRIM28^{-/-} or littermate control cells following CRISPR/Cas9-mediated genetic ablation of Nrp1 (upper row) or Snai3 (lower row). Data are from 3 to 4 independent experiments. The colors indicate 2 different single-guide RNAs used for gene targeting. The dots represent individual mice, and paired Wilcoxon test was used to test for statistical significance. (F and G) Representative dot plots (F) and quantification (G) of IFN₇⁺ in WT and TRIM28^{-/-} CD4 T cells following overexpression of myristoylated Akt and differentiation in Th1 conditions. Wilcoxon (to compare mock to Akt-Myr for each sample) or Mann–Whitney U test (to compare WT to KO) test were used to test for statistical significance. The dots represent individual mice. Data are from at least 2 independent experiments. *P < 0.05; **P < 0.01.

Since the overexpression of a constitutively active form of Akt in TRIM28^{-/-} CD4 T cells rescued their ability to differentiate into cytokine-expressing effectors, and since NRP1 has been shown to interfere with the PI3K–Akt pathway by recruiting the phosphatase PTEN (30), we believe that TRIM28 regulates peripheral T cell activation by epigenetically silencing antagonists of the PI3K–Akt–mTOR pathway during thymic development.

Altogether, this work unravels an unexpected, mechanism of control of effector T cell responses through epigenetic imprinting in thymocytes and/or naive T cells. Whether this process of "imprinting" is physiologically controlled during infections, chronic inflammation or aging will be a fascinating subject for future studies.

Methods

For detailed information and protocols, see SI Appendix.

In Vitro T Cell Activation and Differentiation. To isolate naive T cells for in vitro assays, secondary lymphoid organs were collected, pooled, and homogenized into single-cell suspensions. Naive T cells were enriched using magnetic and fluorescence-activated cell sorting (MACS and FACS) of naive CD4⁺, CD62L⁺, CD25⁻, CD44⁻ and CD8⁺, CD62L⁺, CD44⁻ T cells using a FACSAria II (BD Biosciences).

Sorted, naive T cells were stained with Cell Trace Violet (Thermo Fisher Scientific) according to the manufacturer's instructions, stimulated on α CD3 (BD Biosciences)-coated flat-bottomed 96-well plates (Corning) with soluble α CD28 (BD Biosciences) in the absence or presence of T cell differentiation mixtures for Th1, Th2, Th17, or Treg.

In Vivo Differentiation Experiments. Naive WT or TRIM28-deficient CD4⁺ OT-II-GFP cells were isolated using a CD4⁺ T cell enrichment kits (Thermo Fisher Scientific) as described above, followed by FACS sorting and staining with Cell Trace Violet as described above. On day -1, 5×10^5 sorted naive T cells were injected i.v. into C57BL/6 recipient mice followed by s.c. immunization with 50 µg of OVA in CFA (Thermo Fisher Scientific) into the left flank on day 0. Four and 8 d later, mice were killed and draining lymph nodes and spleen were removed for analysis of OT-II proliferation (division index based on Cell Trace Violet dilution) and cytokine production. Animal care and use for this study were performed in accordance with the recommendations of the

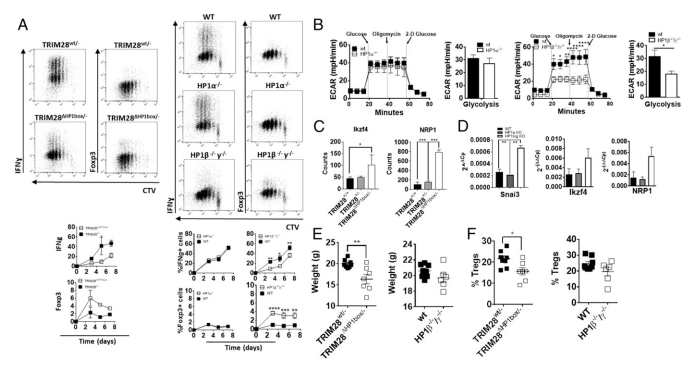


Fig. 8. TRIM28 recruits HP1 β and HP1 γ to silence Treg signature genes and regulate the metabolic switch. (A) Representative dot plots and quantification of IFN γ and Foxp3 expression in differentiated Th1 cells from TRIM28^{ΔHP1box/-}, HP1 $\alpha^{-/-}$, and HP1 $\beta^{-/-}/\gamma^{-/-}$ or littermate control mice. (*B*) Quantification of the extracellular acidification rate (ECAR) of CD4⁺ T cells from HP1 $\alpha^{-/-}$ and HP1 $\beta^{-/-}/\gamma^{-/-}$ or littermate control T cells activated for 24 h using α CD3/ α CD28 as measured by the Seahorse XFe96 Analyzer. The bars or dots indicate mean + SEM. Data are from at least 4 independent experiments. (*C*) mRNA quantification using NanoString probes and RNA isolated from naive CD4⁺ T cells from TRIM28^{ΔHP1box/-}, TRIM28^{MH/-}, or WT mice. (*D*) mRNA quantification using qPCR and RNA isolated from naive CD4⁺ T cells from TRIM28^{ΔHP1box/-}, TRIM28^{MH/-}, or WT mice. (*D*) mRNA quantification using qPCR and RNA isolated from naive CD4⁺ T cells from TRIM28^{ΔHP1box/-}, TRIM28^{MH/-}, or WT mice. (*D*) mRNA quantification. (*E*) Weight of Rag2^{-/-} mice reconstituted with BM from TRIM28^{ΔHP1box/-}, HP1 $\beta^{-/-}/\gamma^{-/-}$, or littermate control mice 6 wk after reconstitution. The dots represent individual mice. (*F*) Percentage of Tregs among CD4⁺ T cells from BM chimeras from *E*. (*A*, *C*, and *E*) Data are from at least 2 independent experiments. Two-way ANOVA (*A* and *B*, kinetics) or Student's *t* test (*B*, average; *C*, *D*, *E*, and *F*) was used to test for statistical significance. **P* < 0.05; ***P* < 0.001; ****P* < 0.001;

European Community (2010/63/UE) for the care and use of laboratory animals. Experimental procedures were specifically approved by the ethics committee of the Institut Curie CEEA-IC #118 (CEEA-IC 2014-03 and CEEA-IC 2017-006) in compliance with the international guidelines.

T Cell Transfer Colitis. Transfer colitis was performed similar as reported by Powrie et al. (20).

BM Chimeras. BM was prepared from tibia and femur of TRIM28^{-/-}, littermate control, or CD45.1 congenic mice, as described previously (41).

Seahorse Assay. Sorted, naive CD4⁺ and CD8⁺ T cells were activated in vitro using plate-bound α CD3 and soluble α CD28 for 24 h as described above. Activated T cells were harvested, counted, and washed, and resuspended in assay medium (Seahorse Biosciences). Assay plates (Seahorse Biosciences) were coated with Cell-Tak (Corning) at 22.4 μ g/mL in PBS for 1 h before cells were plated at a density of 2 \times 10⁵ cells per well according to the manufacturer's instructions. Mitochondrial respiration and glycolysis were Biosciences) according to the manufacturer's instructions.

Immunohistochemistry and Microscopy. Immunohistochemistry of colon samples. Swiss roll samples were fixed in 4% formaldehyde, and 4-µm-thick slices were cut and embedded in paraffin for further immunohistochemical analysis. CD3 (clone 145-2C11) staining was performed at pH9 (Dako 52367) using a 1:100 dilution for 1 h at room temperature. Staining was revealed using an Elite ABC kit (Vector Laboratories) using diaminobenzidine for 10 min. Nuclei were stained using diluted Harris hematoxylin for 1 min. All immunostainings were performed on a LabVision Autostainer (Thermo Fisher Scientific).

RNA Isolation and Analysis by qPCR or NanoString nCounter. RNA was isolated from naive, activated, or differentiated effector or regulatory cells using the

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miRNEasy isolation kit (Qiagen) according to the manufacturer's instructions. For qPCR, 200 ng of RNA was transcribed into cDNA using reverse transcriptase (Thermo Fisher Scientific) according to the manufacturer's instructions. qPCR was performed using 5 ng of template cDNA, H₂O, Sybr Green (Roche), and specific primers on a LightCycler 480 (Roche) (45 cycles, 60 °C annealing temperature). The following primer sequences were used (9, 42, 43).

Chromatin Immunoprecipitation. Chromatin immunoprecipitation was performed similar as previously published (9). Data is deposited in the GEO database (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE140448).

CRISPR/Cas9 Deletion in Naive CD4 T Cells. CRISPR/Cas9-based deletion of NRP1 and Snai3 in naive CD4 T cells was performed as described in ref. 44.

Transduction and Protein Overexpression in CD4 T Cells. T cell transduction protocol was performed similar to Singh et al. (45). Cytokine production was measured 2 d after transduction. See *SI Appendix* for detailed information.

Statistical Analysis. Student's *t* test and one-way ANOVA were used for normally distributed data, while Mann–Whitney *U* or Kruskal–Wallis tests were used for not normally distributed data. For paired samples (e.g., data from mixed BM chimera), paired *t* test or Wilcoxon test was used, depending on the type of distribution. Data from kinetic experiments were analyzed using 2-way ANOVA using Bonferroni's multiple-test correction. **P* < 0.05, ***P* < 0.01, ****P* < 0.001, and *****P* < 0.001. Statistical analysis was performed using GraphPad software).

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