DOCK8 enforces immunological tolerance by promoting IL-2 signaling and immune synapse formation in Tregs

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Patients deficient in the guanine nucleotide exchange factor DOCK8 have decreased numbers and impaired in vitro function of Tregs and make autoantibodies, but they seldom develop autoimmunity. We show that, similarly, Dock8−/− mice have decreased numbers and impaired in vitro function of Tregs but do not develop autoimmunity. In contrast, mice with selective DOCK8 deficiency in Tregs develop lymphoproliferation, autoantibodies, and gastrointestinal inflammation, despite a normal percentage and in vitro function of Tregs, suggesting that deficient T effector cell function might protect DOCK8-deficient patients from autoimmunity. We demonstrate that DOCK8 associates with STAT5 and is important for IL-2-driven STAT5 phosphorylation in Tregs. DOCK8 localizes within the lamellar actin ring of the Treg immune synapse (IS). Dock8−/− Tregs have abnormal TCR-driven actin dynamics, decreased adhesiveness, an altered gene expression profile, an unstable IS with decreased recruitment of signaling molecules, and impaired transendocytosis of the costimulatory molecule CD86. These data suggest that DOCK8 enforces immunological tolerance by promoting IL-2 signaling, TCR-driven actin dynamics, and the IS in Tregs.

Introduction

T cells play a critical role in fighting pathogens by generating a diverse repertoire of antigen receptors (1). This diversity leads to the generation of T cell clones that recognize self-antigen, potentially causing autoimmunity (2). The majority of autoreactive T cell clones are eliminated in the thymus by negative selection. Autoreactive T cells that escape to the periphery are inactivated by Tregs to maintain peripheral tolerance (3). Interactions between Tregs and DCs are crucial in preventing activation of autoreactive T cells in the periphery (4).

The development of Tregs in the thymus requires intermediate strength interactions between TCRs and self-peptide/MHC ligands and costimulatory signals delivered by CD28 and members of the tumor necrosis factor receptor superfamily (5, 6). These signals lead to the upregulation of the high-affinity IL-2 receptor (IL-2R) α chain (CD25). IL-2 signaling via the high-affinity IL-2R complex containing CD25 activates the transcription factor STAT5, which binds to the promoter of the Foxp3 gene. FOXP3 expression is critical for the survival, maintenance, and function of Tregs (7–9). TCR signaling is also important for the induction of Foxp3 gene transcription and Treg function (10). FOXP3 is dispensable for Treg development but is crucial for Treg function, as evidenced by the development of severe autoimmunity in patients and mice deficient in FOXP3 (11). Thus, both IL-2 signaling and TCR signaling are important for Treg function.

Dedicator of cytokinesis 8 (DOCK8) is a member of the DOCK180 superfamily of DOCK proteins with characteristic DOCK homology region 1 (DHR1) and DHR2 domains (12). The DHR1 domain targets DOCK8 to membranes, through binding phosphatidylinositol (3,4,5)-triphosphate, while the DHR2 domain binds to, and functions as, a guanine nucleotide exchange factor (GEF) for CDC42 (13, 14). The
GEF activity of DOCK8 is critical for its function (15). DOCK8 regulates actin cytoskeleton-dependent functions in T cells, B cells, NK cells, and DCs (14–18). DOCK8 deficiency in humans is caused by biallelic mutations in DOCK8 that abolish protein expression (19). DOCK8 deficiency is associated with atopnic dermatitis, asthma, food allergies, an unusual susceptibility to viral mucocutaneous infections, T cell lymphopenia, reduced proliferative T cell responses, decreased cytokine production, and impaired antibody responses (20, 21).

We previously reported that the number and in vitro–suppressive function of circulating Tregs are significantly reduced in DOCK8-deficient patients (22). However, DOCK8-deficient patients only sporadically develop autoimmune disease (20, 23–26). We report that mice with selective deficiency of DOCK8 in Tregs, but not Dock8–/– mice, develop rampant autoimmunity, suggesting that deficient T effector cells (Teff) function may protect DOCK8-deficient patients from autoimmunity. We show that DOCK8 regulates IL-2–driven STAT5 phosphorylation, TCR-driven actin dynamics, immune synapse (IS) integrity, and transendoctyosis in Tregs, all of which are important for maintaining peripheral tolerance.

**Results**

DOCK8-deficient mice have decreased numbers and impaired in vitro function of Tregs but do not develop autoimmune disease, which could potentially affect their Tregs (27). To circumvent these limitations, we examined mice that carry a homozygous knockin c.C1074T mutation, recapitulating a mutation in a DOCK8null patient (21). These mice, designated Dock8–/– mice, express no detectable DOCK8 protein (16). As previously reported (15, 28), the percentage and number of CD4+ T cells and marginal zone B cells in the spleens of Dock8–/– mice were decreased compared with WT controls (Supplemental Figure 1, A and B; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.94298DS1). Proliferation and IL-2 production and secretion by CD4+CD25– Teffs in response to soluble anti-CD3 and antigen-presenting cells (APCs) was significantly decreased in response to soluble anti-CD3 and antigen-presenting cells (APCs) was significantly decreased compared with WT controls (Figure 1C). When Tregs from Dock8–/– mice were preactivated with IL-2 and anti-CD3+CD28 beads, there was no significant difference in their ability to suppress Teff proliferation (Supplemental Figure 1E). This supports a role for decreased IL-2 secretion from Dock8–/– mice in impairing the in vitro–suppressive ability of Tregs from Dock8–/– mice.

The ability of CD4+CD25+CD39+ Tregs from Dock8–/– mice to suppress the proliferation of WT CD4+CD25+ Teffs in response to soluble anti-CD3 and antigen-presenting cells (APCs) was significantly decreased compared with that of Tregs from WT controls (Figure 1G). Tregs suppress the proliferation of Teffs by several mechanisms, including competition for IL-2 (34) and production of the inhibitory cytokines TGF-β and IL-10 (35, 36). In addition to decreased CD25 surface expression, which reduces the ability to compete for IL-2, sorted CD4+CD25+CD39+ Tregs from Dock8–/– mice had significantly impaired Tgfβ, but not Il10, expression compared with WT controls (Figure 1H). When Tregs from Dock8–/– and WT mice were preactivated with IL-2 and anti-CD3+CD28 beads, there was no significant difference in their ability to suppress Teff proliferation (Supplemental Figure 1E). This supports a role for decreased IL-2 secretion from Dock8–/– Teffs in impairing the in vitro–suppressive ability of Tregs from Dock8–/– mice.

Despite the decreased percentages and impaired in vitro–suppressive activity of their Tregs, Dock8–/– mice aged for 12 months had normal weight gain and did not develop lymphadenopathy, splenomegaly, anemia, thrombocytopenia, or gastrointestinal inflammation. In addition, their sera did not exhibit increased reactivity to HEp-2 cell antigens or to the 128 autoantigens included in the University of Texas Southwestern panel (37) compared with WT controls (data not shown). Intraperitoneal administration of the TLR3 agonist poly(I:C) for 3 months to mimic microbial stimulation did not elicit autoantibody formation against HEp-2 cells and did not provoke autoimmune disease in Dock8–/– mice (data not shown).
Mice with selective DOCK8 deficiency in Tregs have impaired weight gain and develop lymphoproliferation, systemic inflammation, and anemia. Defective Teff function may have protected Dock8−/− mice from developing autoantibodies and autoimmunity. To test this hypothesis, we examined mice with a selective DOCK8 deficiency in Tregs. Dock8flox/flox mice that carryloxP sites flanking exons 44 through 46 (Supplemental Figure 2A) were generated and bred with Foxp3Cre-/+ mice that selectively express a YFP-Cre fusion protein under the control of the Foxp3 promoter. Foxp3Cre-/+ /Dock8floxflox male mice and Foxp3Cre-/+ /Dock8floxflox female mice (both designated as Foxp3Cre-/+ /Dock8floxflox mice) lacked DOCK8 expression in CD4+CD25+ Tregs but retained normal DOCK8 expression in CD4+CD25− Teffs (Supplemental Figure 2B).

Figure 1. Dock8−/− mice have reduced Treg percentages and in vitro Treg-suppressive ability. (A and B) Proliferation measured by Cell Trace Violet dilution (A) and IL-2 secretion in culture supernatants (B) by CD4+CD25− Teffs isolated from the spleens of Dock8−/− and WT mice cultured for 3 days with anti-CD3+anti-CD28-coated beads. (C) Percentage of CD25+FOXP3+ Tregs among CD4+ cells in the thymuses, spleens, and LNs of Dock8−/− and WT mice. n = 17 mice from each group for the thymus, n = 31 mice from each group for the spleen, n = 7 mice from each group for the LN. (D) Percentages of CD44+CD62L+ rTregs and CD44−CD62L− aTregs of total CD4+FOXP3+ cells in the spleens of Dock8−/− and WT mice. n = 5 mice from each group. (E) Representative FACS plots of intracellular FOXP3 and CTLA-4 and surface CD25 expression gating on CD4+FOXP3+ splenocytes (left). Quantitative analysis of surface CD25 expression by splenic CD4+FOXP3+ cells from Dock8−/− mice and WT controls. n = 29 mice from each group. (F) qPCR analysis of Foxp3 and Il2ra mRNA levels in FACS-sorted CD4+CD25+CD39+ Tregs from Dock8−/− and WT mice. Results are expressed as fold increase relative to the WT control ratio of the mRNA of interest/b2microglobulin. (G) Suppression of the proliferation of CD4+CD25+ Teffs by CD4+CD25+CD39+ Tregs from Dock8−/− mice and WT controls. Teff proliferation was measured by FACS analysis of Cell Trace Violet dilution. The left panel is a representative experiment; the right panel shows the pooled results. (H) qPCR analysis of Tgfb and Il10 mRNA expression by sorted CD4+CD25+CD39+ Tregs from Dock8−/− mice and WT controls. Symbols represent individual mice, and error bars represent mean and SEM. Results in A, B, and F–H are representative of 3 independent experiments. t test, NS P > 0.05, * P < 0.05, ** P < 0.01, *** P < 0.001.
Foxp3YFP–Cre/Dock8flox/flox mice had poor weight gain starting at 10 weeks of age, and their weight from 18 weeks of age onward was significantly lower than that of age-matched Foxp3YFP–Cre controls (Figure 2A).

By 28 weeks of age, serum levels of amyloid P, an acute-phase reactant indicative of systemic inflammation (38), were significantly higher in Foxp3YFP–Cre/Dock8flox/flox mice compared with Foxp3YFP–Cre controls (Figure 2B).

Foxp3 YFP–Cre/Dock8flox/flox mice developed microcytic anemia with significantly lower hemoglobin concentration and red blood cell volume compared with Foxp3 YFP–Cre controls (Figure 2C). Platelet counts were not significantly different between Foxp3YFP–Cre/Dock8flox/flox mice and controls (data not shown).

Figure 2. Mice with Treg-specific DOCK8 deficiency develop weight loss, splenomegaly, and lymphadenopathy. (A) Weights between age 10 and 30 weeks in Foxp3YFP–Cre/Dock8flox/flox and Foxp3YFP–Cre controls. n = 11 Foxp3YFP–Cre/Dock8flox/flox and 10 Foxp3YFP–Cre mice. (B) Serum amyloid P levels in Foxp3YFP–Cre/Dock8flox/flox mice and Foxp3YFP–Cre controls at 28 weeks of age. (C) Hemoglobin concentration and mean red blood cell volume (MCV) in Foxp3YFP–Cre/Dock8flox/flox mice and Foxp3YFP–Cre controls at 12 and 28 weeks of age. (D and E) Representative photographs of the spleens (D), cervical LNs (CLN), and inguinal LNs (ILN) (E) from 30-week-old Foxp3YFP–Cre/Dock8flox/flox mice and Foxp3YFP–Cre controls. (F) Nucleated cells counts, CD4+ cell counts, percentage of CD69+ cells among CD4+ cells, and percentage of CD4+CD44 CD62L naive and CD4+CD44+CD62L memory T cells among CD4+FOXP3+ cells in the spleens of 30-week-old Foxp3YFP–Cre/Dock8flox/flox mice and Foxp3YFP–Cre controls. (G) Cytokine mRNA expression in pooled CLNs and ILNs from 30-week-old Foxp3YFP–Cre/Dock8flox/flox mice and Foxp3YFP–Cre controls. Results of the qPCR analysis were expressed as fold increase relative to control of the cytokine mRNA/b2microglobulin mRNA ratio. (H) Representative FACS plots showing intracellular cytokine expression after gating on CD4+ T cells stimulated for 3.5 hours with phorbol 12,13-dibutyrate and ionomycin. (I) Serum IgA and IgE levels in 28-week-old Foxp3YFP–Cre/Dock8flox/flox mice and Foxp3YFP–Cre controls. Symbols represent individual mice, and error bars in A–C, F, G, and I represent mean and SEM. t test, NS P > 0.05, * P < 0.05, ** P < 0.01, *** P < 0.001.
Foxp3YFP–Cre/Dock8flx/flx mice spontaneously developed splenomegaly and peripheral lymphadenopathy by 6–8 weeks of age (Figure 2, D and E). Total cellularity and numbers of CD4+ T cells, CD8+ T cells, and B220+ B cells were significantly higher in the spleens and peripheral LNs of Foxp3YFP–Cre/Dock8flx/flx mice compared with age-matched Foxp3YFP–Cre controls (Figure 2F; Supplemental Figure 2, C and D; and Figure 3). Mice with a Treg-specific DOCK8 deficiency develop gastrointestinal inflammation. (A) Representative photographs of the MLNs from 30-week-old Foxp3YFP–Cre/Dock8flx/flx mice and Foxp3YFP–Cre controls. (B) Total cellularity, number of CD4+ T cells, and percentage of activated CD69+ T cells of CD4+ T cells in MLNs from 30-week-old Foxp3YFP–Cre/Dock8flx/flx mice and Foxp3YFP–Cre controls. (C and D) Representative photographs of the stomachs (C) and representative H&E-stained sections of the forestomach and glandular stomach (original magnification, ×10) (D) from 30-week-old Foxp3YFP–Cre/Dock8flx/flx mice and Foxp3YFP–Cre controls. Scale bar: 100 μm (D). (E) Representative images (original magnification, ×20) of indirect immunofluorescent staining of stomach sections for IgG using dilutions of serum from a 28-week-old Foxp3YFP–Cre/Dock8flx/flx mouse and Foxp3YFP–Cre controls. (F and G) Representative H&E-stained sections (original magnification, ×20) (F) and enteritis pathology score (G) of the small intestine (SI) from 30-week-old Foxp3YFP–Cre/Dock8flx/flx mice and Foxp3YFP–Cre controls. Scale bar: 100 μm. (H) qPCR analysis of cytokine mRNA expression in the SI of 30-week-old Foxp3YFP–Cre/Dock8flx/flx mice and Foxp3YFP–Cre controls. Results are expressed as fold increase of the cytokine mRNA/b2microglobulin mRNA ratio relative to control. (I and J) Representative H&E-stained sections (original magnification, ×20) (I) and colitis pathology score (J) of the mid colon from 30-week-old Foxp3YFP–Cre/Dock8flx/flx mice and Foxp3YFP–Cre controls. (K) Fecal lipocalin-2 concentrations in the stools of 30-week-old Foxp3YFP–Cre/Dock8flx/flx mice and Foxp3YFP–Cre controls. Results are expressed as fold increase of the cytokine mRNA/b2microglobulin mRNA ratio relative to control. Symbols represent individual mice, and error bars in B, G, H, J, and K represent mean and SEM. t test, *P < 0.05, **P < 0.01, ***P < 0.001.
data not shown). The percentage of activated CD4+CD69+ T cells was increased in the peripheral LNs, but not spleens, of Foxp3 YFP–Cre/Dock8flox/flox mice compared with Foxp3 YFP–Cre controls (Supplemental Figure 2D and Figure 2F).

qPCR analysis revealed significantly increased mRNA expression of Il17a and Il4, but not Il2, Il5, Il13, or Ifng, in peripheral LNs from Foxp3 YFP–Cre/Dock8flox/flox mice compared with Foxp3 YFP–Cre controls (Figure 2G). Intracellular FACS analysis revealed increased percentages of IL-17A+ and IL-4+ CD4+FOXP3– T cells in Foxp3 YFP–Cre/Dock8flox/flox mice compared with Foxp3 YFP–Cre controls (Figure 4A and 4B).

Figure 4. Tregs from mice with a Treg-specific DOCK8 deficiency have impaired fitness. (A and B) Proliferation measured by Cell Trace Violet dilution (A) and IL-2 secretion in culture supernatants (B) by CD4+CD25– Teffs isolated from the spleens of Foxp3 YFP–Cre/Dock8flox/flox and Foxp3 YFP–Cre control mice cultured for 3 days with anti-CD3+anti-CD28–coated beads. (C) Percentage of YFP+ Tregs of CD4+ T cells and percentage of CD44+ FOXP3+ Tregs and CD44+CD62L+ aTregs among total Tregs in spleens of 30-week-old Foxp3 YFP–Cre/Dock8flox/flox mice and Foxp3 YFP–Cre controls. (D) Percentage of YFP+ cells of total CD4+ cells in the intraepithelial lymphocytes (IEL) and LP of the stomachs and colons and in the skin from Foxp3 YFP–Cre/Dock8flox/flox mice and Foxp3 YFP–Cre controls. (E) qPCR analysis of Foxp3, Il2ra, Tgfb, and Il10 mRNA levels in FACSDerived CD4+CD25+YFP+ Tregs from Foxp3 YFP–Cre/Dock8flox/flox mice and Foxp3 YFP–Cre controls. Results are expressed as fold increase relative to the ratio of the mRNA of interest/b2microglobulin in Foxp3 YFP–Cre controls. (F) Suppression of the proliferation of CD4+CD25+ Teffs stimulated with mitomycin C–treated APCs and soluble anti-CD3 mAb at a Teff/Treg ratio of 1:1. Treg proliferation was measured by FACS analysis of Cell Trace Violet dilution. (G) YFP+/YFP– cell ratio in the spleens and MLNs of Foxp3 YFP–Cre/Dock8flox/flox females and Foxp3 YFP–Cre/+ controls normalized to a ratio of 1.0 in the controls. (H) Quantitative analysis of surface CD25 and intracellular FOXP3 protein expression by splenic CD4+YFP+ cells from Foxp3 YFP–Cre/Dock8flox/flox females and Foxp3 YFP–Cre/+ controls. Results in A, B, and F are representative of 3 independent experiments. Symbols represent individual mice, and error bars represent mean and SEM. t test, NS P > 0.05, *P < 0.05, **P < 0.01, ***P < 0.001.
Teffs, but not CD4\(^+\)Foxp3\(^+\) Tregs, in LNs of Foxp3\(^{YFP-Cre}\)/Dock8\(^{flox/flox}\) mice compared with Foxp3\(^{YFP-Cre}\) controls (Figure 2H), indicating that Teffs, but not Tregs, are the major source of increased cytokine production in the LNs of Foxp3\(^{YFP-Cre}\)/Dock8\(^{flox/flox}\) mice. IL-4 drives IgE switching (39). Foxp3\(^{YFP-Cre}\)/Dock8\(^{flox/flox}\) mice had a marked increase in levels of serum IgE (Figure 2I). Serum IgA, but not IgG or IgM, was also significantly elevated (Figure 2I and data not shown). Thus, selective DOCK8 deficiency in Tregs results in the development of lymphoproliferation and T cell activation, with increased expression of IL-17A and IL-4 cytokines in Teffs in the setting of normal numbers of Tregs in the lymphoid tissues.

Mice with selective DOCK8 deficiency in Tregs develop severe gastrointestinal inflammation. Treg defects are often associated with gastrointestinal inflammation (40). Foxp3\(^{YFP-Cre}\)/Dock8\(^{flox/flox}\) mice spontaneously developed enlarged mesenteric LNs (MLNs) by 6–8 weeks of age (Figure 3A). Total cellularity, numbers of CD4\(^+\) T cells, and the percentage of activated CD4\(^+\)CD69\(^+\) T cells were all significantly higher in the MLNs from Foxp3\(^{YFP-Cre}\)/Dock8\(^{flox/flox}\) mice compared with Foxp3\(^{YFP-Cre}\) controls (Figure 3B).
Figure 6. DOCK8-deficient Tregs have abnormal shape and F-actin kinetics and an unstable immune synapse. (A) Baseline F-actin content of CD4^+CD25^+ Tregs from Dock8^{−/−} and WT mice, and effect of CD3 crosslinking on the F-actin content of Tregs from Dock8^{−/−} and WT mice. Results are expressed as the change in the MFI of F-actin from the baseline (time 0). (B) Representative images of CD4^+CD25^+ Tregs from Dock8^{−/−} mice and WT controls plated on anti-CD3- and ICAM-1-coated glass chambers and stained for F-actin by phalloidin at 5 and 45 minutes (original magnification, ×100). (C) Relative number of adherent cells per unit area at 5 minutes. (D) Quantitative analysis of F-actin staining of Tregs after 5 and 45 minutes of stimulation. (E) Localization of pTYR, F-actin, and DOCK8 in Dock8^{−/−} and WT Tregs at 10 minutes (original magnification, ×100). (F) A representative TIRF image of DOCK8 and actin distribution across the synapse after 10 minutes of synapse formation. The graph shows line scan profiles of DOCK8 and ACTIN across the middle of the cell; the green trace represents ACTIN intensity distribution, and the red trace represents DOCK8 intensity distribution. Note that DOCK8 and ACTIN intensities coincide in the cell periphery, represented by the peaks at the beginning and end of the line scan profiles. (G) Side view of 2 representative T cells, showing the whole cell ACTIN, pTYR, and DOCK8 distribution. Cells stained with DOCK8, pTYR, and ACTIN were imaged using confocal microscopy, and the images show a side view of maximum intensity projection of the confocal images. (H) Quantification of cell edge roughness determined by “shape descriptors” utility of ImageJ at 5 and 45 minutes. (I) Measurement of immune synapse instability, as denoted by the kinapse index (transient interactions index > 1) at 5 and 45 minutes. (J) Relative staining intensity of pTYR, phospho-ZAP70 (pZAP70), and TALIN at 10 minutes. (K) Transendocytosis of CD86-GFP by Tregs from Dock8^{−/−} and WT mice cocultured with CD86-GFP-expressing CHO cells. The percentage of GFP^+ Tregs of total Tregs is shown. Results in A and K are representative of 3 independent experiments. Results in B–J are representative of 2 independent experiments. Symbols represent individual measurements, and error bars represent mean and SEM. Significance was determined by unpaired t test. ns P > 0.05, *P < 0.05, **P < 0.01, ***P < 0.001.
Figure 7. DOCK8-deficient Tregs have decreased expression of cytoskeleton- and Treg-associated genes. (A and B) CD4⁺CD25⁺CD39⁺YFP⁻ (DOCK8-deficient) and YFP⁺ (DOCK8-sufficient) Tregs were sorted from Foxp3 YFP⁻Cre/+ Dock8 flox/flox female mice. Tregs were cultured overnight in media alone (A) or with anti-CD3+CD28 beads (B). Heatmaps of selected genes differentially expressed in YFP⁻ and YFP⁺ Tregs from Foxp3 YFP⁻Cre/+ Dock8 flox/flox female mice. The cutoff for significance was P < 0.05. P values were calculated using the Wald test for differential expression analysis. P values were corrected afterward for multiple testing. Expression of genes is centered and scaled by row to highlight differences in each gene sample. Each column represents an individual mouse. (C and D) CD4⁺CD25⁺CD39⁺ Tregs were FACS sorted from Dock8⁻/- and WT mice. RNA was prepared from Tregs directly after isolation (C) or after 24-hour culture with anti-CD3+CD28 beads (C and D). qPCR results are expressed as fold increase of mRNA of interest/b2microglobulin ratio relative to the unstimulated WT Tregs. (E) MFI of surface marker staining on YFP⁺ DOCK8-sufficient and YFP⁺ DOCK8-deficient CD4⁺CD25⁺CD39⁺ Tregs from Foxp3 YFP⁻Cre/+ Dock8 flox/flox female mice. Symbols represent individual mice. Bars in C-E represent the mean and SEM. t test, NS P > 0.05, *P < 0.05, ***P < 0.001.
The stomachs of Foxp3^{YFP-Cre}/Dock8^{−/−} mice were pale and enlarged compared with those from age-matched Foxp3^{YFP-Cre} controls (Figure 3C). Histologic examination revealed hyperkeratosis of the forestomach in Foxp3^{YFP-Cre}/Dock8^{−/−} mice. In addition, the glandular stomach was hyperplastic with cystic crypts and mononuclear infiltration of the lamina propria (LP) (Figure 3D). Gastric-specific antibodies have long been associated with Treg deficiency in neonatally thymectomized mice (41, 42). Gastric autoantibodies were detected in the sera of 6 of 8 (75%) Foxp3^{YFP-Cre}/Dock8^{−/−} mice at dilutions up to 1:320, but none were detected in the sera from 6 age-matched Foxp3^{YFP-Cre} controls (Figure 3E). Histologic examination of the small intestine showed thickened villi with lymphocytic infiltration of the LP in Foxp3^{YFP-Cre}/Dock8^{−/−} mice compared with Foxp3^{YFP-Cre} control (Figure 3F). Foxp3^{YFP-Cre}/Dock8^{−/−} mice had significantly elevated enteritis histopathology scores and an increase in Ifng mRNA expression in the small intestine compared with age-matched Foxp3^{YFP-Cre} controls (Figure 3, G and H). The colons from Foxp3^{YFP-Cre}/Dock8^{−/−} mice had LP densely infiltrated with lymphocytes and plasma cells. Goblet cells were reduced, and the mucosa was markedly thickened (Figure 3I). The colitis histopathology score was significantly elevated in Foxp3^{YFP-Cre}/Dock8^{−/−} mice compared with age-matched Foxp3^{YFP-Cre} controls (Figure 3J). Lipocalin-2 is secreted by neutrophils, and its levels are increased in the feces in murine models of colitis (43). The fecal content of lipocalin-2 was significantly higher in Tregs from Foxp3^{YFP-Cre}/Dock8^{−/−} mice compared with Foxp3^{YFP-Cre} controls (Figure 3L). These data demonstrate that selective DOCK8 deficiency in Tregs results in the development of severe gastrointestinal inflammation.

Tregs from mice with selective DOCK8 deficiency in Tregs have normal in vitro function but impaired fitness. As expected, proliferation and IL-2 secretion by CD4^+CD25^− Teffs following stimulation with anti-CD3+anti-CD28–coated beads were not significantly different between Foxp3^{YFP-Cre}/Dock8^{−/−} mice and Foxp3^{YFP-Cre} controls (Figure 4, A and B). There was an increased percentage of total and activated YFP^+ Tregs in the spleens and LNs of Foxp3^{YFP-Cre}/Dock8^{−/−} mice compared with Foxp3^{YFP-Cre} controls (Figure 4C, Supplemental Figure 3A, and data not shown). However, there was a significant decrease in the percentage of YFP^+ Tregs among CD4^+ T cells in the LP of the inflamed stomachs and colons and among intraepithelial lymphocytes in the stomachs, small intestines, and colons of Foxp3^{YFP-Cre}/Dock8^{−/−} mice compared with Foxp3^{YFP-Cre} controls (Figure 4D). In contrast, the percentage of Tregs in the noninflamed skin of Foxp3^{YFP-Cre}/Dock8^{−/−} mice was comparable to that in the skin of Foxp3^{YFP-Cre} controls (Figure 4D). These results suggest that DOCK8 is important for the homeostasis of Tregs in the inflamed gut.

Expression of CD25, FOXP3, CTLA-4, CD39, and HELIOS was comparable in CD4^+YFP^+ Tregs from Foxp3^{YFP-Cre}/Dock8^{−/−} mice and Foxp3^{YFP-Cre} controls (Supplemental Figure 3B). Only ICOS expression was significantly higher in Tregs from Foxp3^{YFP-Cre}/Dock8^{−/−} mice compared with Foxp3^{YFP-Cre} controls (Supplemental Figure 3B), suggestive of activation-induced Treg instability (44). Foxp3, Il2ra, and Tgfb, but not Il10, mRNA levels were significantly lower in sorted CD4^+YFP^+ Tregs from Foxp3^{YFP-Cre}/Dock8^{−/−} mice compared with Foxp3^{YFP-Cre} controls (Figure 4E). The ability of CD4^+YFP^+ Tregs from Foxp3^{YFP-Cre}/Dock8^{−/−} mice to suppress the proliferation of WT Teffs in response to stimulation with anti-CD3 beads and APCs was comparable to that of CD4^+YFP^+ Tregs from Foxp3^{YFP-Cre} controls (Figure 4F), suggesting that cell-extrinsic factors affected the decreased in vitro–suppressive activity observed in Tregs from Dock8^{−/−} mice (Figure 1G).

We examined the fitness of DOCK8-deficient Tregs by comparing the ratio of DOCK8-deficient (YFP^+) to DOCK8-sufficient (YFP^−) Tregs in Foxp3^{YFP-Cre}/Dock8^{−/−} females. DOCK8-deficient Tregs demonstrated a competitive disadvantage compared with DOCK8-sufficient Tregs in populating the spleen and LNs, as evidenced by a significantly lower ratio of CD4^+FOXP3^+YFP^+ cells to CD4^+FOXP3^+YFP^− cells in Foxp3^{YFP-Cre+/−}/Dock8^{−/−} females compared with Foxp3^{YFP-Cre+/+} controls (Figure 4G). This is despite comparable expression of CD25 and FOXP3 by CD4^+YFP^+ Tregs from Foxp3^{YFP-Cre+/−}/Dock8^{−/−} females and Foxp3^{YFP-Cre+/+} female control mice (Figure 4H).

DOCK8-deficient Tregs have defective IL-2–driven STAT5 phosphorylation, and DOCK8 constitutively associates with STAT5. STAT5 phosphorylation in Tregs is important for the ability of Tregs to suppress autoreactive Teffs (45). IL-2 stimulation of splenic Tregs resulted in a significantly lower percentage of phosphorylated STAT5^+ (pSTAT5^+) cells in Foxp3^{YFP-Cre}/Dock8^{−/−} mice compared with Foxp3^{YFP-Cre} controls (Figure 5A). As expected, IL-2–driven STAT5 phosphorylation was comparable in CD4^+YFP^+ DOCK8-sufficient Teffs from the two strains (Supplemental Figure 4A). pSTAT5 content following IL-2 stimulation was substantially lower in YFP^+ Tregs compared with YFP^− Tregs in Foxp3^{YFP-Cre+/−}/Dock8^{−/−} female mice, which had no evidence of inflammation, but was not significantly different between YFP^+ and YFP^− Tregs in Foxp3^{YFP-Cre+/+} controls.
Dock8–/– WT control Tregs (Supplemental Figure 5B). However, F-actin content between Dock8–/– dependent on TCR/CD3-driven F-actin reorganization (48). There was no significant difference in baseline does not have a detectable effect on the gross localization of Tregs in LNs.

primarily in the T cell–rich zone of LNs (Supplemental Figure 5A), demonstrating that DOCK8 deficiency or to the onset of lymphadenopathy. Similar to DOCK8-sufficient Tregs, DOCK8-deficient Tregs were found

adhered to the coverslip and formed a synapse (Figure 6, B and C).

anti-CD3 mAb and ICAM-1 to mimic the interaction between Tregs and APCs. Total internal reflection

phosphorylation in Tregs. IL-2–induced STAT5 phosphorylation was impaired in CD4+FOXP3+ Tregs as

"kinapse" (52).

Dock8 –/– actions with other DCs (51). This sampling behavior with rapidly dissolving and reforming IS is termed
costimulatory molecules on the DC, Tregs become increasingly motile and move on to form transient inter-
decreasing the costimulatory function of DCs, rendering them tolerogenic. Following the decrease in

CTLA-4–expressing Tregs acquire the costimulatory molecules CD80 and CD86 from DCs, thereby

maintenance of total pTYR proteins, pZAP70, and pCASL was reduced in the IS of Tregs from

accumulation of TALIN in the IS was significantly reduced in Tregs from Dock8–/– mice compared with control

incidence of kinapses, characterized by an elongated, polarized morphology (Figure 6, B and I), a reflec-
tion of a motile synapse. These data suggest that DOCK8-deficient Tregs are impaired in their ability to

maintain a stable IS with APCs.

TCR-driven IS formation is characterized by the accumulation of tyrosine phosphorylated proteins (pTYR) that include the TCR signal transducer ZAP70 and the mechanosensor CASL (53, 54). Accumulation of total pTYR proteins, pZAP70, and pCASL was reduced in the IS of Tregs from Dock8–/– mice compared with control WT Tregs (Figure 6I). TCR/CD3 ligation causes conformational changes in LFA-1 via inside-out signaling that promote adhesiveness to ICAM-1 (55). TALIN, a protein that binds to integrins, as well as DOCK8, accumulates at the IS of T cells (56, 57). Consistent with reduced adhesion, the accumulation of TALIN in the IS was significantly reduced in Tregs from Dock8–/– mice compared with control
expression of the IL-2–regulated genes Foxp3, Tgfb2, Tbx21 (refs. 58–60 and Figure 7, A and B). Treg transcription factors with differential expression included which encode proteins that localize with F-actin and either bind F-actin directly and/or regulate its assembly Rgs1 (64–66), and genes that encode markers of Tregs and Treg subsets or enable tissue homing, including Nrp1 all N-GAL, which is present in high levels in the stools of patients with colitis (43, 76). While colitis developed in CTLA-4 expression, which was comparable in anti-CD3+anti-C28–stimulated Tregs from er than that of Tregs from WT controls (Figure 6K). This difference was not due to a difference in surface CTLA-4 expression by Tregs. Tregs were sorted from Foxp3YFP–Cre/+/Dock8flox/flox and WT controls (Supplemental Figure 5C).

In addition, reduced surface expression of TIGIT, PD1, and NRP1 was observed on YFP+ DOCK8-deficient Tregs compared with YFP– DOCK8-sufficient Tregs from (DOCK8-deficient) and CD4+CD39+CD25+YFP– (DOCK8-sufficient) Tregs were 123 before stimulation (112 genes were downregulated) and 162 (128 genes were downregulated) after stimulation. We focused our analysis on actin cytoskeleton– and Treg-related genes. Several of these genes were underexpressed, whereas none were overexpressed in DOCK8-deficient Tregs. Actin cytoskeleton–related genes included with decreased expression included Vill, Cagg, Zyx, and Cadl1, which encode proteins that localize with F-actin and either bind F-actin directly and/or regulate its assembly (refs. 58–60 and Figure 7, A and B). Treg transcription factors with differential expression included Foxp3, Stat1, Tbx21, Id2, and Gata3, which are involved in the polarization and stabilization of the Treg phenotype (61–63). Other Treg-related genes included Tgfb2, Tigit, Tnfsf18, and Il2ra, which are involved in Treg-suppressive activity (64–66), and genes that encode markers of Tregs and Treg subsets or enable tissue homing, including Nrp1, Rgs1, Baf1, Fdx1, Itgae, Socs1, Cxcr3, Ccr4, and Ccl24 (Figure 7, A and B, and refs. 61, 67–75). Decreased expression of the IL-2–regulated genes Foxp3, Il2ra, Tnfsf18, Socs1, and Cxcr3 in Dock8–/– Tregs is consistent with impaired IL-2–driven STAT5 phosphorylation in these cells. The decreased levels of Cagg, Foxp3, Il2ra, Stat1, Tigit, and Cxcr3 in Dock8–/– Tregs was confirmed by qPCR analysis of mRNA (Figure 1F and Figure 7, C and D). In addition, reduced surface expression of TIGIT, PD1, and NRPI was observed on YFP– DOCK8-deficient Tregs compared with YFP– DOCK8-sufficient Tregs from Foxp3YFP–Cre+/Dock8flox/flox female mice (Figure 7E).

Discussion

Our findings establish that DOCK8 expression in Tregs is critical for peripheral self-tolerance. Mice with selective deficiency of DOCK8 in Tregs develop spontaneous lymphoproliferation and autoimmune disease. DOCK8-deficient Tregs had impaired IL-2–driven STAT5 phosphorylation, abnormalities in TCR actin dynamics, an unstable IS, and diminished transendocytosing, all of which are important for the ability of Tregs to maintain tolerance.

Foxp3YFP–Cre+/Dock8flox/flox mice with a selective DOCK8 deficiency in Tregs, but intact Teff function, developed spontaneous inflammation and autoimmunity at a young age. This is evidenced by their failure to gain weight and the development of anemia, progressive lymphoproliferation, elevated serum amyloid P levels, increased numbers of activated CD4+T cells, and increased expression of Il17a and Il4 mRNA in their LNs. Foxp3YFP–Cre+/Dock8flox/flox mice also developed autoantibodies against gastric antigens and an autoimmune gastroenteropathy characterized by a dense lymphocytic infiltrate in the LP throughout the gut. The small intestines of Foxp3YFP–Cre+/Dock8flox/flox mice demonstrated significantly increased expression of Ifng, suggesting a role for Th1-mediated inflammation in driving enteritis, while colons demonstrated significantly increased expression Il2, Ifng, Il17a, and Il4 mRNA, suggesting that multiple cytokines were involved in driving colitis in these mice. Foxp3YFP–Cre+/Dock8flox/flox mice had an increased fecal content of lipocalin-2, the homolog of human N-GAL, which is present in high levels in the stools of patients with colitis (43, 76). While colitis developed in all Foxp3YFP–Cre+/Dock8flox/flox mice, dermatitis did not. The gut microbiota may be important in driving colitis in these mice, as in other models of autoimmune enteropathy (77–79). Together, these findings suggest that Treg dysfunction may play a role in the colitis that develops in patients with DOCK8 deficiency (80).

In contrast to Foxp3YFP–Cre+/Dock8flox/flox mice, Dock8–/– mice did not develop autoimmunity even with aging or chronic stimulation with poly(I:C) to mimic microbial infection. We similarly found no signs of spontaneous autoimmunity in CD4-Cre/Dock8flox/flox mice, which lack DOCK8 in both Teffs and Tregs (E. Janssen and R.S.
Geha, unpublished observations). These findings support the hypothesis that DOCK8 function in Teffs is crucial for the establishment of autoimmunity in the context of DOCK8-deficient Tregs and may explain the low prevalence of autoimmunity in DOCK8-deficient patients. Decreased availability of IL-2 from Teffs together with impaired IL-2–driven STAT5 activation may have contributed to the decreased percentage of Tregs and impaired in vitro–suppressive ability of Tregs from Dock8−/− mice. In contrast, Foxp3YFP-Cre/Dock8flox/flox mice, in which IL-2 production by Teffs was intact, had normal or elevated percentages of Tregs in lymphoid organs and normal Treg in vitro–suppressive activity.

A role for DOCK8 in the maintenance of Tregs is supported by the competitive disadvantage of YFP+ DOCK8-deficient Tregs compared with YFP+ DOCK8-sufficient Tregs in populating the lymphoid organs of Foxp3YFP-Cre+/+/Dock8flox/flox females. A function for DOCK8 in Treg homeostasis is also suggested by the significant decrease in the percentage of Tregs in the inflamed stomach and colon, but not the uninflamed skin, of Foxp3YFP-Cre/Dock8flox/flox mice. Both TCR and IL2-driven signals, which were impaired in DOCK8-deficient Tregs, are important for the competitiveness and fitness of Tregs (7, 8, 10, 47).

There was a significant decrease in baseline and IL-2–induced STAT5 phosphorylation in Tregs from Foxp3YFP-Cre/Dock8flox/flox mice. This defect was cell intrinsic and not secondary to inflammation, since it was found in YFP+ DOCK8-deficient Tregs from Foxp3YFP-Cre+/+/Dock8flox/flox female mice, which showed no evidence of inflammation. The IL-2 signaling defect in Tregs may have contributed to the development of autoimmune disease in Foxp3YFP-Cre/Dock8flox/flox mice. We demonstrated that DOCK8 associates constitutively with STAT5 in T cells. This association is not increased by IL-2 stimulation. DOCK8 may bring STAT5 in proximity to the IL-2R facilitating STAT5 phosphorylation by receptor-associated JAK kinases. DOCK8 exists in T cells in a high-molecular-weight complex that contains proteins that include WASP, WIP, and TALIN (16, 57). The association of DOCK8 and STAT5 in T cells may be direct or may be mediated by one of the other proteins in this complex.

DOCK8-deficient Tregs had impaired ability to increase their F-actin content in response to TCR/CD3 ligation, exhibited defective adhesion to anti-CD3– and ICAM-1–coated coverslips, and displayed shape abnormalities over time compared with WT Tregs. Using an anti-CD3 and ICAM-1 system designed to mimic the Treg and DC interaction, we found that DOCK8 localizes to the lamellar actin ring in Tregs. DOCK8-deficient Tregs formed a less stable IS and were more prone to transient kinapse formation. This was accompanied by decreased recruitment to the IS of phosphotyrosine proteins, including phospho-ZAP70. In addition, accumulation at the IS of the mechanosensor pCASL, which is essential for the establishment of IS stability (81), and of TALIN, a known DOCK8 interactor crucial for IS stability, was reduced in Dock8−/− Tregs. DOCK8-deficient Tregs were significantly less efficient at transendocytosis of CD86 from CHO cells, a system that relies on Treg interactions with CD86 exclusively. These findings suggest that DOCK8 has a critical role in the formation of a stable IS between Tregs and DCs and in transendocytosis.

Gene expression analysis by RNAseq suggested that DOCK8 plays a role in the expression of several cytoskeletal proteins important for actin assembly, IS stability, adhesion, and motility (58–60). DOCK8-deficient Tregs demonstrated decreased expression of several IL-2–regulated genes that included Foxp3 and Il2ra, consistent with impaired IL-2–driven STAT5 phosphorylation. There was also decreased expression of several transcription factors that affect Treg stability and subset development (62, 67, 68, 72, 74, 82, 83). This may be relevant to the rampant Th1/Th17 intestinal inflammation observed in Foxp3YFP-Cre/Dock8flox/flox mice. We did not observe a change in the expression of genes previously shown to be differentially expressed in Tregs lacking TCR expression (47), suggesting that TCR signals affecting gene expression remain intact in the absence of DOCK8.

In summary, our findings suggest that DOCK8 is important for IL-2 signaling, TCR-driven actin remodeling, IS stability, and transendocytosis in Tregs and, thereby, Treg function in maintaining of peripheral immunological tolerance.

Methods
Mice. The generation of Dock8−/− mice is described in ref. 16. To generate Dock8 conditional knockout mice, intron 43–44 to 46–47 were amplified by PCR from the genomic DNA of 129 ES cells. The 3.15-kb 5′ arm, 6.5-kb 3′ arm, and a 5.4-kb sequence containing exons 44–46 were cloned into the pEZ-Frt-Lox-DT vector (Supplemental Figure 2A). Mice were generated using standard techniques and mated with mice expressing the Flp recombinase under the Rosa promoter to remove the neo gene and then backcrossed with C57BL/6 mice for 10 generations. Homozygous Dock8flox/flox mice were then mated with B6.129(Cg)-Foxp3tm4(YFP/cre)Ayr/J.
mice (Jackson Laboratories) to generate mice with a Treg-specific deficiency in DOCK8. Both female and male mice were studied, and sex matching was done between comparison groups. All mice were kept in a specific pathogen–free environment. Unless otherwise indicated 3 mice were in all comparison groups, and individual experiments were repeated at least 3 times. No randomization was performed.

Autoantibody detection. Prepared slides with HEp-2 cells and mouse stomach tissues were obtained from Trinity Biotech. Serum was diluted in PBS and applied to each slide for 30 minutes. Slides were washed with PBS+1% Tween, and bound IgG was detected with anti-mouse IgG conjugated to Alexa Fluor 488 (Biolegend). Samples were scored by 3 blinded individuals.

Flow cytometry. Single-cell suspensions were made of the thymuses, spleens, and LNs; red blood cells were lysed using ACK lysis buffer. For staining of surface molecules, cells were incubated on ice with fluorochrome-conjugated mAbs against CD4 (clone GK1.5), CD8 (clone 53-6.7), CD45.2 (clone 104), CD25 (clone PC61), CD44 (clone IM7), CD62L (clone MEL-14), CD39 (clone DaVa59), CD69 (clone H1.2F3), and ICOS (clone C398.4A) (Biolegend). For intracellular staining with antibodies against HELIOS (2F6), FOXP3 (FJK-16s), and CTLA-4 (UC10-4B9), cells were fixed, permeabilized, and stained according to the manufacturer’s directions (Affymetrix).

For intracellular cytokine staining, cells were either stimulated with anti-CD3+anti-CD28 dynabeads (ThermoFisher) for 2 days followed by the addition of Golgistop and Golgiplug (BD Biosciences) for 16 hours or were stimulated with phorbol 12,13-dibutyrate and ionomycin (Sigma-Aldrich) in the presence of Golgistop and Golgiplug for 3.5 hours. They were then fixed and permeabilized using the BD Bioscience Cytofix/Cytoperm kit and incubated with fluorochrome-conjugated antibodies against IL-2 (JES6-5H4), IL-17A (eBio17B7), and IL-4 (11B11) (Affymetrix).

All data were acquired on a BD LSRFortessa cell analyzer using FACS Diva software (BD Biosciences). Analyses were performed using FlowJo software (Tree Star Inc.).

Preparation of intraepithelial lymphocytes and LP cells. The stomach and colon were isolated and flushed with PBS+1% FBS on ice. After mincing with scissors, the epithelial layer was stripped by agitation in 10 mM EDTA at 37°C. This was followed by collagenase VIII digestion (Sigma-Aldrich) and mechanical disruption to free the LP cells. Specimens were enriched for lymphocytes with a 40% Percoll gradient, before analysis by flow cytometry as detailed above.

Treg suppression assays. CD4+ T cells were isolated by negative selection using the CD4+ T cell isolation kit (Miltenyi Biotec) according to the manufacturer’s directions. CD4+CD25+ Teffs were isolated by depleting CD25+ cells using anti-CD25-biotin and anti-biotin magnetic beads (Miltenyi Biotec). Teffs were labeled with Cell Trace Violet (Life Technologies) to monitor cell divisions by flow cytometry. Tregs were isolated by fluorescent cell sorting for CD4+CD25+CD39+ Tregs or CD4+CD25+CD39+YFP+ Tregs. For preactivation of Tregs, cells were cultured for 48 hours with anti-CD3+anti-CD28 dynabeads (ThermoFisher) at a ratio of 1 bead/1 Treg in the presence of 20 ng/ml IL-2 (R&D Systems). Tregs were then washed several times, the dynabeads were removed with a magnet, and Tregs were recounted before their addition to Teff cultures. Teffs were stimulated with anti-CD3 mAb (1 μg/ml, 2C11, Affymetrix) in the presence of T cell–depleted splenocytes treated with 25 μg/ml mitomycin C (Santa Cruz Biotechnology). On day 4, Teff division was evaluated by flow cytometry.

Cellular proliferation and IL-2 secretion. CD4+CD25+ Teffs were isolated using magnetic beads (Miltenyi) and labeled with Cell Trace Violet as described above. Prepared Teffs were stimulated with anti-CD3+anti-CD28 beads (Invitrogen) at a ratio of 1 bead/1 cell in round bottom wells. After 72 hours in culture, supernatants were harvested and IL-2 content was measured by ELISA (Affymetrix). An equal volume of complete media was added back to each well, and proliferation was evaluated by flow cytometry on day 4 of culture.

Equivalent numbers of CD4+CD25+ Tregs were stimulated with IL-2 at the indicated concentrations and/or anti-CD3+anti-CD28 beads (Invitrogen). After 6 days in culture, T cells were analyzed by FACS for viability, and cell numbers were normalized using Precision Count beads (Biolegend).

Immunoglobulins and protein assays. Serum IgM, IgG, IgA, and IgE levels were measured by ELISA using capture and detection antibodies from Southern Biotech. Serum amyloid P and haptoglobin (Genway) and stool lipocalin-2 (R&D Systems) levels were measured by ELISA. For the lipocalin-2 ELISA, fresh fecal samples were vortexed with PBS+0.1% Tween for 20 minutes before being spun down at 4°C, and the lipocalin-2 concentration of the supernatant was measured.

Histologic analysis. Enteritis and colitis scoring was performed by a blinded trained pathologist based on the scoring criteria in ref. 84.
Immunofluorescence analysis of Tregs in LNs. LNs were fixed overnight in 4% periodate-lysine-paraformaldehyde solution at 4°C; serially passed through 10%, 20%, and 30% sucrose solutions for 24 hours each; embedded in OCT compound; and rapidly frozen in ethanol cooled on dry ice. Thawed 20-μm cryotome sections adhered onto superfrost glass slides were washed 3 times for 5 minutes each with PBS without Ca²⁺/Mg²⁺, permeabilized with precooled 90% methanol for 10 minutes at –20°C, washed for 30 minutes, treated with the Endogenous Biotin-Blocking Kit (Thermo Fisher), and blocked in Fc block (1:200, Biolegend) with 1% goat serum and 0.25% BSA in PBS for 1 hour. Sections were then incubated overnight at 4°C with biotinylated anti-FOXP3 antibody (1:50, clone FJK-16s, eBioscience) in blocking solution and, after thorough washing, stained with Streptavidin–Alexa Fluor 647 (1:500, Thermo Fisher) for 8 hours at 4°C, washed, and stained with Pacific Blue–conjugated anti-B220 antibody (1:200, Biolegend) for 8 hours at 4°C. After several thorough washings, sections were mounted on coverslips in Prolong Gold (Thermo Fisher). In order to visualize entire LN sections, tile imaging was performed on a LSM 780 AxioObserver confocal microscope (Carl Zeiss) equipped with a ×40 lens (Apochromat, 1.1 W Korr M27). Pacific blue was excited by 405-nm laser line, and emission was collected with MBS 405 (424–509 nm). Alexa Fluor 647 was excited by 633-nm laser line, and emission was collected with MBS488/568/633 (638–755 nm). Images were processed in FIJI (open-source).

IL-2–driven STAT5 phosphorylation. The percentage of pSTAT5⁺ T cells and cellular content of pSTAT5 was determined before and after stimulation for 15 minutes with concentrations of mouse IL-2 (R&D Systems) as indicated in the figure legends. Cells were fixed using BD Cytofix fixation buffer and permeabilized with BD Phosflow Perm buffer III. After permeabilization, cells were stained with anti-STAT5 pY694 (clone 47, BD Biosciences) and anti-STAT5 (44-386G, ThermoFish).

Immunoprecipitation and immunoblotting assays. Cells were lysed in 0.5% NP-40 buffer (150 mM NaCl, 25 mM Tris-CI [pH 7.5], 5 mM EDTA) containing complete protease inhibitors (Roche). Cell lysates were incubated with anti-DOCK8 (rabbit polyclonal, Sigma-Aldrich) or anti-STAT5 (C-17, Santa Cruz). Immune complexes were captured with protein G-Sepharose (EMD Millipore), washed, denatured by boiling in sample buffer, separated on acrylamide gels, and analyzed by immunoblotting with anti-DOCK8 (H-159, Santa Cruz) and anti-STAT5 (C-17, Santa Cruz). Protein band intensities were quantified by densitometry using the ImageJ software (NIH).

Determination of cellular F-actin content. Splenocytes were rested in RPMI-1640 for 3 hours on ice. Cells were then incubated with 10 μg/ml rat anti-mouse CD3 mAb KT3 (Serotec) for 20 minutes on ice and stimulated by cross-linking with 20 μg/ml F(ab')₂ goat anti-rat IgG secondary antibody (Jackson Immunoresearch Laboritories Inc.) for the indicated times at 37°C. Cells were fixed in 4% formaldehyde, washed, and permeabilized with the BD Cytofix/Cytoperm kit (BD Biosciences). F-actin was stained with fluorescein isothiocyanate-labeled phallidin (Sigma-Aldrich) and F-actin content was measured with a BD LSRRFortessa cell analyzer.

ILS visualization. CD4⁺CD25⁺ T cells freshly isolated from Dock8⁻/⁻ and WT mice were incubated in glass chambers (Mat-tek) precoated with 10 μg/ml anti-CD3 (2C11, Affymetrix) and 1 μg/ml ICAM-1 (extracellular fragment) (85), for 5, 10, or 45 minutes at 37°C (53). Cells were then fixed using PHEM buffer at 37°C, permeabilized (0.1% Triton-X 100, 5 minutes), and stained with Alexa 568-phalloidin (Life Technologies), p130CASL (pY165), anti-phosphotyrosine (multi-mAb rabbit antibody), anti-pZAP70 (pTyr319)/pSyk(Tyr352) (Cell Signaling Technologies), and anti-TALIN (C20, goat polyclonal) (Santa Cruz Biotechnology) antibodies overnight, followed by incubation with fluorescently labeled anti-rabbit or anti-goat antibodies for 1 hour. Cells were then imaged using TIRF microscopy using a Nikon Eclipse Ti microscope equipped with a ×100 1.49 NA TIRF objective, and an AndorDU897 back illuminated EMCCD camera. The acquired images were analyzed using ImageJ software as described previously (53). The cell boundaries were identified using the “particle analysis” plug-in of ImageJ, and the ROI obtained were further utilized extract morphological and intensity information associated with the raw images.

Transendocytosis. CHO-K1 cells (ATCC) were transfected with a vector encoding CD86-GFP (Origene), and stable transfectants were selected for with G418 (Sigma-Aldrich). CD86-GFP expressing CHO cells were then cocultured with sorted CD4⁺CD25⁺CD39⁺ Tregs at a 1:1 ratio in the presence of 10 nM bafilomycin A (Invigene). Uptake of GFP by Tregs was measured by FACS after 24 hours of coculture.

Quantitative analysis of gene expression and code availability. Tregs were isolated by fluorescent sorting for CD4⁺CD25⁺CD39⁺, CD4⁺CD25⁺CD39⁺YFP⁺, or CD4⁺CD25⁺CD39⁺YFP⁻ Tregs. RNA was isolated using a RNAeasy Plus micro kit (Qiagen) according to the manufacturer's instructions. Total small intestine and colonic RNA were isolated by homogenization in lysis/binding solution followed by RNA isolation using an RNAxesous phenol-free total RNA isolation kit (Ambion) according to the manufacturer's instructions.
Complementary DNA was reverse transcribed using an iScript cDNA synthesis kit (Biorad) according to the manufacturer’s instructions. Analysis of transcripts was performed with commercial TaqMan primer/probe sets against mouse b2m, Il2ra, Foxp3, Tgfb1, Capg, Tigit, Stat1, Il10, Il2, Ilng, Il17a, Il4, Il5, and Il13 (Thermo Fisher).

For RNAseq, CD4+CD25+CD39+YFP+ and CD4+CD25+CD39+YFP- Tregs were isolated from the spleens and LNs of Foxp3\textsuperscript{Cre+/-}Dock8\textsuperscript{flox/flox} mice. RNA was isolated as described above. Smart-Seq2 libraries were generated and sequenced using a Broad Genomics Platform (86–88). The quality of the sequencing files was initially assessed with the FastQC tool. They were then trimmed to remove contaminant sequences (like polyA tails), adapters, and low-quality sequences with cutadapt. These trimmed reads were aligned to the mouse genome (UCSC build mm10) using STAR aligner. Alignments were checked using a combination of FastQC, Qualimap, MultiQC, and custom tools. Counts of reads aligning to known genes were generated by featureCounts. In parallel, transcripts per million measurements per isof orm were generated using Sailfish. Differential expression at the gene level was called with DESeq2. We used the counts per gene estimated from Sailfish by tximport as input to DESeq2, as quantifying at the isof orm level has been shown to produce more accurate results at the gene level. The GEO accession is GSE102577.

Statistics. Comparisons were analyzed for statistical significance using unpaired 2-tailed Student’s \textit{t} test and 2-way ANOVA to determine \(P\) values using GraphPad Software. A \(P\) value of less than 0.05 was considered significant.

Study approval. All mouse studies were approved and performed in accordance with Boston Children’s Hospital Institutional Animal Research and Care Committee.

Author contributions
EJ, SK, MT, SU, JMJT, MCR, RTB, VB, and SMU performed the experiments and analyzed the data. EJ generated the mutated mice. TRM, DJI, and RSG designed and supervised the research. EJ, SK, and RSG wrote the manuscript.

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