Wiskott-Aldrich syndrome (WAS) is associated with mutations in the WAS protein (WASp), which plays a critical role in the initiation of T cell receptor–driven (TCR-driven) actin polymerization. The clinical phenotype of WAS includes susceptibility to infection, allergy, autoimmunity, and malignancy and overlaps with the symptoms of dedicator of cytokinesis 8 (DOCK8) deficiency, suggesting that the 2 syndromes share common pathogenic mechanisms. Here, we demonstrated that the WASp-interacting protein (WIP) bridges DOCK8 to WASp and actin in T cells. We determined that the guanine nucleotide exchange factor activity of DOCK8 is essential for the integrity of the subcortical actin cytoskeleton as well as for TCR-driven WASp activation, F-actin assembly, immune synapse formation, actin foci formation, mechanotransduction, T cell transendothelial migration, and homing to lymph nodes, all of which also depend on WASp. These results indicate that DOCK8 and WASp are in the same signaling pathway that links TCRs to the actin cytoskeleton in TCR-driven actin assembly. Further, they provide an explanation for similarities in the clinical phenotypes of WAS and DOCK8 deficiency.

Introduction

The integrity of the actin cytoskeleton is important for T cell migration into tissues, defense against pathogens, and immunosurveillance (1). The Wiskott-Aldrich syndrome (WAS) protein WASp plays an important role in the organization and function of the actin cytoskeleton in hematopoietic cells (2). WASp and its homolog neuronal WASp (N-WASp) are important activators of the actin-related protein 2/3 (ARP2/3) complex, which initiates actin polymerization (2). In resting cells, WASp exists in a closed, inactive conformation due to intramolecular interactions that prevent the C-terminal acidic domain from interacting with the ARP2/3 complex. The small Rho GTPase cell division cycle 42 (CDC42) is an important activator of WASp (3). Binding of CDC42-GTP to the GTPase-binding domain (GBD) of WASp causes a conformational change, which allows WASp to interact with and activate the ARP2/3 complex (4, 5). T cells deficient in WASp have low baseline F-actin levels, impaired T cell receptor-driven (TCR-driven) actin assembly, immune synapse (IS) formation, actin foci formation and mechanotransduction, and defective migration into tissues (2, 6). WASp is complexed with the WASp-interacting protein (WIP), which is essential for WASp stability (2). WIP also binds to actin (2). T cells from WIP-deficient mice and knockin mice expressing a WIP mutant that fails to bind actin exhibit cytoskeletal and functional defects similar to, but more severe than, those of WASp-deficient T cells (7), suggesting that WIP can regulate the T cell actin cytoskeleton independently of WASp.

Dedicator of cytokinesis 8 (DOCK8) is one of the eleven members of the DOCK180 superfamily (8). DOCK proteins have characteristic DOCK homology region 1 and 2 (DHR1 and DHR2) domains. The DHR1 domain is important for targeting DOCK proteins to membranes, through its binding of phosphatidylinositol (3, 4, 5)-triphosphate (PIP3). The DHR2 domain binds to and functions as a GEF for CDC42 (9). Several observations suggest that DOCK8 regulates actin cytoskeleton–dependent functions. DOCK8-deficient B cells, T cells, and NK cells have defective IS formation (10–12), and DOCK8-deficient DCs have defective migration in vivo and in vitro in a 3D collagen gel matrix (13). WASp deficiency, which results in the X-linked WAS, and DOCK8 deficiency, which causes autosomal recessive hyper IgE syndrome, share clinical and laboratory findings that include recurrent sinopulmonary infections, viral skin infections, eczema, elevated serum IgE levels, food allergies, and an increased risk for malignancy and autoimmune diseases (14–16). Moreover, DOCK8 is a GEF for the WASp activator CDC42 (17), and DOCK8 and WASp exist in a complex in a human NK cell line (18). We show here that
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precipitates (Figure 1, A and B). These results indicate that DOCK8
constitutively associates with WASp and WIP in primary T cells. TCR ligation induces tyrosine phosphorylation of DOCK8, WASp,
and WIP (19, 20). Ligation of the TCR complex in human
T cells with anti-CD3 mAb had no detectable effect on the associ-
ation of DOCK8 with WASp or WIP (Figure 1C) but caused robust
ERK phosphorylation, demonstrating effective TCR signaling.
Similar results were obtained with mouse splenic T cells (data not
shown). These results indicate that TCR signaling does not alter
the constitutive association of DOCK8 with WASp and WIP.

The subcellular localization of DOCK8, WASp, and WIP in T cells
was examined in the human T cell acute lymphoblastic leukemia
(T-ALL) lymphoblast cell line DND41, which has a relatively
abundant cytoplasm compared with that of primary resting T cells.
As in primary T cells, WASp and WIP coimmunoprecipitated with
DOCK8 from DND41 cell lysates (Figure 1D). Intracellular immuno-
fluorescence revealed that DOCK8 colocalized with both WASp and
WIP bridges DOCK8 to WASp and actin and that DOCK8, and in
particular DOCK8 GEF activity, is essential for TCR-driven WASp
activation and F-actin assembly, the integrity of the subcortical
actin cytoskeleton, TCR-driven actin foci formation and mechno-
transduction, and transendothelial migration (TEM) of T cells, all
of which also depend on WASp. These findings provide a molecular
explanation for the shared features of DOCK8 deficiency and WAS.

Results

DOCK8 binds to and colocalizes constitutively with WASp and WIP in T cells. We examined whether DOCK8 interacts with WASp and WIP
in primary T cells. Lysates from CD3+ T cells purified from human
blood and mouse spleens were immunoprecipitated with rabbit
anti-DOCK8 polyclonal Ab, or rabbit anti-MALT1 polyclonal Ab
as a negative control, and the immunoprecipitates were probed for
DOCK8, WASp, and WIP. Immunoblot analysis demonstrated the
presence of WASp and WIP in DOCK8, but not MALT1, immuno-
precipitates (Figure 1, A and B). These results indicate that DOCK8
constitutively associates with WASp and WIP in primary T cells.

TCR ligation induces tyrosine phosphorylation of DOCK8, WASp,
and WIP (19, 20). Ligation of the TCR complex in human
T cells with anti-CD3 mAb had no detectable effect on the associ-
ation of DOCK8 with WASp or WIP (Figure 1C) but caused robust
ERK phosphorylation, demonstrating effective TCR signaling.
Similar results were obtained with mouse splenic T cells (data not
shown). These results indicate that TCR signaling does not alter
the constitutive association of DOCK8 with WASp and WIP.

Figure 1. DOCK8 interacts constitutively and colocalizes with WASp and WIP in T cells. (A and B) Co-IP of WIP and WASp with DOCK8, but not MALT1, in
human peripheral blood T cells (A) and mouse splenic T cells (B). Aliquots from the lysates used for IP were probed with DOCK8, WASp, WIP, and MALT1 to
to ensure equal loading. (C) Lack of a detectable effect of TCR ligation with anti-CD3 mAb on the association of DOCK8 with WIP and WASp in human blood
T cells. Lysates were probed with a phospho-specific ERK1/ERK2 (p-ERK1/2) Ab to verify TCR signaling and with GAPDH as a loading control. IP with an
IgG isotype Ab was used as a negative control. (D) Co-IP of WIP and WASp with DOCK8, but not MALT1, in the human DND41 T cell line. (E) Colocalization
of WIP and WASp with DOCK8. Representative DIC images (left), confocal immunofluorescence microscopic images (middle), and merged images (right)
of DND41 T cells permeabilized and stained with Abs against DOCK8, WIP, and WASp, followed by fluorochrome-labeled secondary Abs. Original magnifi-
cation, ×63. Scale bars: 2 μm. Data in A-D represent 3 independent experiments each. Images in E are representative of 50 cells that were examined in 2
independent experiments.
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controls (Figure 2B). These results indicate that DOCK8 binds to WIP independently of WASp.

WIP-deficient T cells express virtually no WASp (21) and are not suitable for determining whether DOCK8 associates with WASp independently of WIP. To determine whether DOCK8 directly binds to WIP or WASp, we examined whether recombinant DOCK8 (rDOCK8) associates with rWIP and rWASp in vitro. DOCK8-Myc, WIP-EGFP, and Flag-tagged WASp fusion proteins were generated using an in vitro wheat germ extract transcription translation system. EGFP and MAL T1-Flag recombinant proteins served as negative controls, and lysate aliquots were probed for Myc and EGFP to ensure equal loading. The * denotes a non-specific band. (F) Co-IP of DOCK8 and WASp with WIP in human T cells, using the 3D10 and D12CS anti-WIP mAbs directed against distinct epitopes in the N-terminus and C-terminus of WIP, respectively. Data in C, E, and F represent 4 independent experiments.
WASp, but not DOCK8 (Figure 2F). These results suggest that DOCK8 binds to the N-terminus of WIP, while WASp binds to the C-terminus of WIP, and are consistent with the finding of WIP bridging DOCK8 and WASp.

WIP bridges DOCK8 to actin. We previously showed that a WIP mutant that lacks the actin-binding domain (aa 43-54), designated WIPΔABD (Figure 3A), fails to bind actin (7). We examined the ability of EGFP-WIPΔABD to bind DOCK8-Myc from lysates of 293T cotransfectants. EGFP-WIPΔABD coprecipitated with DOCK8-Myc to an extent comparable to that seen with EGFP-WIP (Figure 3B), indicating that the WIP-binding sites for DOCK8 and actin are distinct.

To examine whether WIP bridges DOCK8 and actin in T cells, we examined the association of DOCK8 and actin in T cells from WT mice and mice homozygous for a knocked-in mutant Wipf1 gene that encodes for WIPΔABD and designated WIPΔABD mice (7). We had previously shown that actin coprecipitates with WIP in WT T cells but that this association is virtually abolished in WIPΔABD T cells (7). Actin was readily detected in DOCK8 immunoprecipitates from WT T cells, but was virtually absent from DOCK8 immunoprecipitates from WIPΔABD T cells (Figure 3, C and D). There was less WIP, as well as less WASp, in lysates and DOCK8 immunoprecipitates from WIPΔABD T cells compared with WT T cells (Figure 3C); however, the WIP/DOCK8 and WASp/DOCK8 ratios in DOCK8 immunoprecipitates relative to lysates and normalizing the values to those obtained in WT T cells by setting the WT ratio to 1. Ctrl, control. Data are representative of 3 independent experiments in B and 2 independent experiments in C and D. Error bars in D represent the mean ± SEM.
immunoblotted with the anti–WASp F-8 mAb, which recognizes both activated and nonactivated WASp. We performed quantitative analysis of activated WASp by calculating the ratio of the intensity of the WASp band in the CSAb immunoprecipitates to that in total lysates. Activated WASp was weakly detectable in unstimulated T cells (Figure 4A). TCR ligation caused a significant increase in activated WASp in T cells from HCs, but not in T cells from DOCK8null patients (Figure 4A). TCR-driven phosphorylation of ERK was comparable in DOCK8-deficient and normal T cells. These results indicate that DOCK8 is required for TCR-driven WASp activation.

Figure 4. DOCK8 GEF activity mediates TCR-driven WASp activation. (A) Representative (left) and pooled (right) results of the generation of active WASp following TCR ligation in human T cells from an HC and a DOCK8null patient as determined by immunoblotting WASp and N-WASp CSAb immunoprecipitates with anti–WASp F-8 mAb. Aliquots from total lysates were probed for WASp to ensure equal loading and for p-ERK1/2 to verify TCR signaling. Results are representative of 2 experiments involving 2 patients and 2 controls. Quantification was performed by calculating the ratio of activated WASp in the IP to total WASp in the lysates, relative to controls. (B) Ribbon diagram of the DOCK8–CDC42 complex. Residue S1827 is in red, and the region affected by the S1827P mutation is in magenta. (C) Interaction of DOCK8 residue S1827 with CDC42 (top) and its disruption by the S1827P DOCK8mut mutation (bottom). (D) GEF activity for CDC42 of WT DOCK8 and DOCK8mut DHR2 domains. Relative fluorescence units (RFU) of MANT-GTP over time are shown. Results are representative of 3 experiments. (E) Representative immunoblot and quantitation of DOCK8 expression in Dock8mut/mut and WT mice. Quantification was performed by calculating the DOCK8/GAPDH ratio relative to that in WT controls. (F and G) Representative immunoblot (F) and quantitation (G) of the association of DOCK8 with WIP and WASp in T cells from DOCK8-mutant mice. Quantification was performed by calculating the WIP/DOCK8 ratio in DOCK8 immunoprecipitates relative to that in WT controls. (H and I) Representative (H) and pooled (I) results of the generation of active WASp following TCR ligation in splenic T cells from Dock8–/–, Dock8mut/mut, and WT mice. The experiment was performed and the results expressed as described in A. Results in E–I are representative of 3 experiments using 3 mice from each strain. Error bars in A, E, G, and I represent the means SEM. *P < 0.05, by Student’s t test.

T cells compared with WT T cells (Figure 3D). These results indicate that WIP bridges DOCK8 to actin in T cells and suggest that WIP acts as a scaffold on which DOCK8, WASp, and actin assemble.

DOCK8 GEF activity mediates TCR-driven WASp activation. A conformation-sensitive Ab (CSAb), directed against a conserved peptide close to the GBD of N-WASp, selectively recognizes the activated, but not the nonactivated, form of WASp (23). To determine whether WASp activation following TCR ligation depends on DOCK8, WASp CSAb immunoprecipitates from anti-CD3–stimulated human T cells from 2 DOCK8null patients and 3 healthy controls (HCs) were
CDC42 has been shown to play an important role in WASp activation and actin reorganization (24–27), along with subsequent T cell activation, cytokine secretion, polarization, and chemotaxis (28–31). To determine the role of DOCK8 GEF activity for CDC42 in TCR-mediated WASp activation, we compared T cells from WT, Dock8−/−, and Dock8pri/pri mice. Dock8−/− mice were generated as detailed in Supplemental Figure 1A; supplemental material available online with this article; doi:10.1172/JCI85774DS1) and have no detectable DOCK8 protein expression (Supplemental Figure 1B). Dock8pri/pri mice carry a homozygous S1827P mutation in the DHR2 domain of DOCK8 postulated to interfere with GEF activity (11). Examination of the crystal structure of the DOCK8 DHR2-CDC42 complex (17) indicates that S1827 is located in the middle of a long helix (residues 1823-1838) of DOCK8, whose N-terminal region interacts with CDC42 (Figure 4B) and engages in water-mediated interactions with CDC42 residues T161 and F28 (Figure 4C). Modeling the S1827P mutation suggests that the Pro residue probably fails to form similar interactions, which may reduce the affinity to CDC42 (Figure 4C). Furthermore, structural dynamics simulation suggests that the S1827P mutation diminishes the interaction with CDC42 indirectly by inducing local conformational changes (data not shown). Using an in vitro exchange activity assay, we directly demonstrated that the DHR2 domain of the S1827P mutant had virtually no detectable GEF activity toward CDC42 (Figure 4D). We observed a modest, but not significant, decrease in DOCK8 expression in Dock8pri/pri T cells compared with that in WT T cells (Figure 4E). Importantly, the association of the S1827P mutant with WASp and WIP was preserved.

Figure 5. Defective actin cytoskeleton structure and function in DOCK8-deficient T cells. (A) Representative EM images of the apical membrane of T cells stimulated with anti-CD3 mAb. Shown are the cytoskeletal actin fibers associated with the cytoplasmic side of the adherent plasma membranes. Results are representative of 2 experiments with T cells from 1 mouse of each strain. More than 50 cells were examined in each experiment. Scale bar: 200 nm. (B) Actin filament length in T cell membranes from the Dock8−/−, Dock8pri/pri, and WT mice represented in A; 114 WT T cells, 139 Dock8−/− T cells, and 120 Dock8pri/pri T cells were measured in 2 independent experiments. (C) Representative FACS analysis of FITC-phalloidin staining for F-actin in resting T cells from Dock8−/−, Dock8pri/pri, and WT mice and quantitative analysis of the results as a percentage of F-actin content in T cells from WT controls. Results are representative of 3 independent experiments using 3 mice from each strain. (D) Increase in the F-actin content of T cells from Dock8−/−, Dock8pri/pri, and WT mice following stimulation with anti-CD3 mAb. Results are expressed as the increase in the mean fluorescence intensity (MFI) of F-actin from the baseline (time 0). Results are representative of 3 independent experiments using 3 mice from each strain. Error bars in B and C and symbols and bars in D represent the mean ± SEM. ***P < 0.001, **P < 0.01, and *P < 0.05, by Student’s t test.
as evidenced by comparable WIP/DOCK8 and WASp/DOCK8 ratios in DOCK8 immunoprecipitates from T cells of Dock8<sup>−/−</sup> and WT mice (Figure 4, F and G). TCR ligation caused a significant increase in activated WASp in T cells from WT mice, but not from Dock8<sup>−/−</sup> or Dock8<sup>pri/pri</sup> mice (Figure 4, H and I). TCR-driven ERK phosphorylation was comparable in DOCK8-mutant and WT T cells. These results demonstrate that DOCK8 GEF activity is critical for TCR-driven WASp activation.

**DOCK8 GEF activity is essential for the integrity of the subcortical actin cytoskeleton and TCR-driven actin polymerization.** Given that DOCK8 is important for the activation of WASp, which regulates F-actin assembly in T cells (32), we examined the actin cytoskeleton and actin dynamics in DOCK8-deficient T cells. To evaluate the ultrastructure of the actin cytoskeleton, purified T cells were spread on anti-CD3–coated glass chambers, their apical surface removed, and the attached cell membranes processed for electron microscopy (EM). EM images revealed that the amount of F-actin associated with the inner side of the plasma membrane was greatly diminished in Dock8<sup>−/−</sup> and Dock8<sup>pri/pri</sup> T cells compared with the dense, branched actin network observed in WT T cells (Figure 5A). The residual filaments in Dock8<sup>−/−</sup> and Dock8<sup>pri/pri</sup> T cells had significantly longer F-actin filaments than did WT T cells (Figure 5B). These findings indicate that DOCK8 GEF activity is important for the integrity of the T cell subcortical actin cytoskeleton.

T cells from Dock8<sup>−/−</sup> and Dock8<sup>pri/pri</sup> mice had significantly reduced basal F-actin levels (Figure 5C), consistent with their sparse subcortical F-actin network. Moreover, the increase in F-actin content following TCR cross-linking was significantly impaired in T cells from Dock8<sup>−/−</sup> and Dock8<sup>pri/pri</sup> mice compared with those from WT controls (Figure 5D). These findings suggest that DOCK8 GEF activity for CDC42 is essential for maintaining the basal level of F-actin in T cells and for TCR-driven actin assembly.

**DOCK8 is essential for TCR-driven actin foci formation and mechanotransduction.** TCR ligation triggers foci formation at the IS, which leads to invadopodia-like T cell membrane protrusions that push into the target cell (6). In addition, TCR-driven actin remodeling induces mechanical strain at the IS, reflected by the phosphorylation of the mechanosensory molecule CasL (the p130Cas isoform expressed in lymphocytes) at the Y165 residue (6). Both TCR-driven actin foci formation and CasL phosphorylation are severely impaired in the absence of WASp (6). To examine the role of DOCK8 in actin foci formation and mechanical stress at the IS, T cells were plated on a glass cover-

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**Figure 6. DOCK8 is essential for TCR-driven actin foci formation and mechanotransduction.** (A and B) Representative images of splenic CD4<sup>+</sup> T cells from WT, Dock8<sup>−/−</sup>, and Dock8<sup>pri/pri</sup> mice plated on anti-CD3– and ICAM-1–coated glass chambers (A) or a superantigen-treated endothelial cell monolayer (B) and stained for F-actin (phalloidin, pseudocolored green) and pCasL (pseudocolored red). Orange indicates merging. The extracellular striated pattern observed in B (phalloidin panel on the left) is due to the F-actin fluorescence contributed by the endothelial cell cytoskeleton. Original magnification, ×100. Scale bar: 5 μm. (C-G) Quantitative analysis of actin foci (C and E), pCasL intensity (D and F), and synaptic area (G) in T cells. Each data point represents a value obtained for a single cell. Similar results (A-G) were obtained in 2 independent experiments. Error bars in C-G represent the mean ± SEM. ***P < 0.001, **P < 0.01, and *P < 0.05, by Student’s t test.
slip coated with anti-CD3 mAb and ICAM-1, and actin foci formation and phosphorylated CasL (p-Casl) levels were assessed using total internal reflection microscopy. F-actin foci formation and p-Casl were significantly reduced in Dock8–/– and Dock8pri/pri T cells compared with that detected in WT T cells (Figure 6, A, C, and D). To better mimic the physiologic interactions between T cells and vascular endothelium, T cells were plated on a superantigen-loaded endothelial cell monolayer, and cell-cell contacts were examined by spinning-disc confocal microscopy. WT T cells plated on the endothelial monolayer exhibited robust F-actin foci formation and CasL phosphorylation; both were significantly reduced in Dock8–/– and Dock8pri/pri T cells (Figure 6, B, E and F).
integrin-mediated firm adhesion and arrest, spreading, and TEM (35). In vitro Th1-polarized CD4+ T cells show high TEM activity and are routinely used to study this process in vitro (36). We generated Th1 T cells in vitro from Dock8–/–, Dock8pri/pri, and WT mice by stimulating splenic CD4+ T cells with anti-CD3 and anti-CD28 under Th1-polarizing conditions. The percentage of generated Th1 cells determined by intracellular staining for IFN-γ was comparable between the 3 groups (76.8% ± 13.3% for Dock8–/–, 77.8% ± 13.5% for Dock8pri/pri, and 84.3% ± 3.6% for WT T cells). Th1 cells from all 3 groups had comparable surface expression of the selectin ligands CD44 and P-selectin glycoprotein ligand 1 (PSGL-1) and the integrins LFA-1 and VLA-4, which mediate adhesion to ICAM-1 and VCAM-1, respectively (Figure 7A). The accumulation of Th1 cells on immobilized E-selectin-Fc and VCAM-1-Fc chimeric proteins under a range of physiological flow conditions (0.5–1.0 dynes/cm2) was comparable among Dock8–/–, Dock8pri/pri, and WT mice (Figure 7B).

In addition, the area of the IS was significantly reduced in T cells from Dock8–/– and Dock8pri/pri mice compared with that of T cells from WT controls (Figure 6G). These results show that DOCK8 GEF activity for CDC42 is important for TCR-driven formation of actin foci and generation of mechanical force at the IS.

DOCK8 is essential for T cell spreading and in vitro TEM. Dynamic rearrangements of the actin cytoskeleton, including actin foci formation, are important for T cell motility and transcellular diapedesis (33). Splenic CD4+ T cells from Dock8–/– and Dock8pri/pri mice migrated normally toward SDF1α under static conditions in a 2D Transwell migration assay (data not shown), like T cells from Dock8cpm/cpm-mutant mice (34). Migration of T cells from blood vessels into tissues occurs under physiological shear flow conditions and involves sequential interactions between adhesion molecules expressed on T cells and endothelial cells, including selectin-mediated rolling and tethering, followed by chemokine-induced integrin-mediated firm adhesion and arrest, spreading, and TEM (35). In vitro Th1-polarized CD4+ T cells show high TEM activity and are routinely used to study this process in vitro (36). We generated Th1 T cells in vitro from Dock8–/–, Dock8pri/pri, and WT mice by stimulating splenic CD4+ T cells with anti-CD3 and anti-CD28 under Th1-polarizing conditions. The percentage of generated Th1 cells determined by intracellular staining for IFN-γ was comparable between the 3 groups (76.8% ± 13.3% for Dock8–/–, 77.8% ± 13.5% for Dock8pri/pri, and 84.3% ± 3.6% for WT T cells). Th1 cells from all 3 groups had comparable surface expression of the selectin ligands CD44 and P-selectin glycoprotein ligand 1 (PSGL-1) and the integrins LFA-1 and VLA-4, which mediate adhesion to ICAM-1 and VCAM-1, respectively (Figure 7A). The accumulation of Th1 cells on immobilized E-selectin-Fc and VCAM-1-Fc chimeric proteins under a range of physiological flow conditions (0.5–1.0 dynes/cm2) was comparable among Dock8–/–, Dock8pri/pri, and WT mice (Figure 7B).

Figure 8. Defective in vivo homing of DOCK8-deficient T cells. (A) Representative FACS analysis of a mixture of equal numbers of Alexa Fluor 555-labeled WT T cells (designated by red lettering) and Alexa Fluor 488-labeled WT, Dock8–/–, and Dock8pri/pri T cells (designated by green lettering) used for injection into genetically matched WT recipients. (B) Representative FACS plots from the blood and LNs of WT recipients 1 hour after i.v. administration of 1:1 mixtures of Alexa Fluor 555-labeled WT T cells and Alexa Fluor 488-labeled WT, Dock8–/–, or Dock8pri/pri T cells. (C) Quantitative analysis of cells from the blood and LNs of WT recipients obtained 1 hour after i.v. administration of 1:1 mixtures of equal numbers of Alexa Fluor 555-labeled WT T cells and Alexa Fluor 488-labeled WT, Dock8–/–, or Dock8pri/pri T cells. Graph shows the homing index of T cells from Dock8–/–, Dock8pri/pri, and WT mice relative to the mean homing index of WT T cells set at 1.0. (D) Percentages of annexin V+ eFluor-780+ apoptotic cells and eFluor-780+ dead cells among Alexa Fluor 555-labeled WT, Dock8–/–, and Dock8pri/pri donor T cells in pooled LN cells from WT recipients. Results in A–D are representative of 2 independent experiments each using 5 recipients per donor strain. Error bars represent the mean ± SEM. ***P < 0.001, **P < 0.01, and *P < 0.05, by Student’s t test.
However, a significant decrease of 57% and 36% in the adhesion of Dock8−/− Th1 cells to ICAM-1-Fc (coimmobilized with SDF1α) was observed at 0.8 and 0.5 dynes/cm², respectively. There was an 18% and 29% decrease in the adhesion of Dock8−/− Th1 cells to ICAM-1-Fc at 0.8 and 0.5 dynes/cm², respectively, but these decreases were not statistically significant. These findings suggest that Dock8 functions, beyond its GEF activity, may contribute to optimal LFA-1-dependent Th1 cell adherence to ICAM-1 under shear flow conditions. Next, DIC microscopic recordings of adherent Th1 cells were analyzed to assess their ability to undergo spreading after arrest on ICAM-1 and SDF1-α, which revealed a significant reduction in the spreading of Th1 cells from Dock8−/− and Dock8−/+ mice compared with that seen in Th1 cells from WT mice (Figure 7C).

We directly examined the capacity of Th1 cells to arrest, crawl over, and transmigrate across a TNF-α–activated murine endothelial cell monolayer under physiological low-flow conditions (0.8 dynes/cm²). Th1 cells from Dock8−/− and Dock8−/+ mice adhered normally to the monolayer; however, the fraction of adherent cells that transmigrated was significantly lower for Dock8−/− and Dock8−/+ T cells than for WT T cells (Figure 7D). These results suggest that Dock8 GEF activity is important for T cell spreading and TEM under shear flow conditions.

Dock8 is essential for T cell homing to lymph nodes. Circulating T cells enter lymph node (LTs) by migrating across high endothelial venules (37). To examine the role of Dock8 in the in vivo T cell homing to LNs, equivalent numbers of Alexa Fluor 488–labeled Dock8−/−, Dock8−/+, or WT T cells were mixed with Alexa Fluor 555–labeled WT T cells and administered i.v. to WT recipients. One hour later, the percentages of transferred WT and Dock8−/− or Dock8−/+ T cells were analyzed in the blood and cervical, axillary, mesenteric, and inguinal LNs by flow cytometry (Figure 8, A and B). The homing index was calculated as the ratio of Alexa Fluor 488– to Alexa Fluor 555–labeled cells and normalized to the ratio of WT controls. The homing index of Dock8−/− and Dock8−/+ T cells was significantly decreased in all LNs examined (Figure 8C). There was a significantly higher retention of Dock8−/− and Dock8−/+ T cells in the blood (Figure 8C). Similar results were obtained when the cell-labeling dyes were reversed (data not shown). Migration of Dock8−/− deficient T cells through 3D spaces that constrain cell movement results in catastrophic cell death that becomes evident after 12 hours (13). Analysis of LNs from WT recipients 1 hour after adoptive transfer revealed no significant differences in the percentages of apoptotic cells or dead cells among Dock8−/−, Dock8−/+, and WT donor T cells (Figure 8D).

These results suggest that Dock8 GEF activity is important for the homing of naive T cells to LNs.

Discussion

We demonstrate that Dock8 and WASp constitutively associate in T cells in a complex in which WIP bridges Dock8 to WASp and actin and that Dock8 GEF activity is essential for TCR-driven WASp activation and actin cytoskeleton reorganization. We also show that Dock8 GEF activity is important for the integrity of the subcortical actin cytoskeleton in T cells and for a number of actin cytoskeleton–mediated T cell functions that also depend on WASp. These findings establish a mechanistic basis for the phenotypic convergence of Dock8 deficiency and WASp deficiency.

IP studies demonstrated that Dock8 constitutively associates with WASp and its partner WIP in human and murine T cells. Consistent with this finding, Dock8 colocalized with WASp and WIP in the T cell line DND41. The association of Dock8 with WASp and WIP was not altered by TCR ligation, indicating that TCR-driven posttranslational modifications of these proteins have no detectable effect on the Dock8-WIP-WASp complex. The association of Dock8 with WIP was direct and independent of WASp. In contrast, WIP was essential for the association of Dock8 and WASp. WIP was also found to mediate the association between Dock8 and actin, and the WIP-binding sites for Dock8, WASp, and actin were found to be distinct. This suggests that WIP serves as a scaffold on which Dock8, WASp, and actin assemble.

Using cells from Dock8−/− patients and mice and an Ab that selectively recognizes the open, active conformation of WASp, we established that Dock8 is critical for TCR-driven WASp activation in T cells. More importantly, we demonstrated that Dock8 GEF...
activity was essential for WASp activation, because TCR ligation failed to activate WASp in T cells from Dock8–/– and Dock8<sup>gmo/gmo</sup> mice, which express a mutant DOCK8 protein with a missense mutation in the DHR2 domain that we show abolishes CDC42-directed GEF activity. These findings place DOCK8 upstream of WASp and suggest that the spatial proximity of DOCK8 and WASp may facilitate WASp activation following TCR ligation. The mechanism by which TCR ligation triggers DOCK8 GEF activity is not known. CCK7 signaling in T cells induces MST1 and MST2 kinase-dependent phosphorylation of MOB1, causing it to associate with DOCK8 and promote its GEF activity (38). TCR ligation activates MST1 (39), thus, MST1 and MOB1 may link the TCR to DOCK8 in T cells.

DOCK8, and in particular DOCK8 GEF activity, were found to be essential for the integrity and function of the actin cytoskeleton in T cells. The subcortical actin network was severely attenuated in T cells from Dock8<sup>–/–</sup> and Dock8<sup>gmo/gmo</sup> mice. Disruption of the subcortical actin network is also observed in WASp-deficient T cells, albeit to a lesser extent than in DOCK8-deficient and WIP-deficient T cells (40, 41), suggesting that DOCK8 and WIP may regulate actin dynamics independently of WASp. In this regard, WIP binds to cortactin, which can drive F-actin assembly (42), and DOCK8 may have GEF activity for RAC1, which activates WASp family verprolin-homologous protein (WAVE) to drive actin polymerization (38). WASp facilitates the branching of filamentous actin in T cells by activating the ARP2/3 complex (43). The residual F-actin filaments in Dock8<sup>–/–</sup> and Dock8<sup>gmo/gmo</sup> T cells were significantly elongated, possibly because of impaired branching due to deficient WASp-mediated ARP2/3 activation in these cells.

Baseline F-actin levels were significantly decreased, and TCR-driven F-actin assembly as well as IS formation were impaired in DOCK8-mutant T cells, as in WASp-deficient and WIP-deficient T cells (2, 40, 44, 45); this further suggests that DOCK8, WIP, and WASp form a functional unit. Defective accumulation of F-actin at the IS has been demonstrated in T cells from Dock8<sup>–/–</sup> and Dock8<sup>gmo/gmo</sup> mice (10). The residual increase in TCR-driven F-actin assembly in DOCK8-deficient T cells may be explained by CDC42-independent activation of WASp (2, 46), as well as by WASp-independent pathways of actin polymerization, e.g., via WAVE (47). Residual activation of WASp via CDC42-independent pathways may also explain the less severe effect of DOCK8 deficiency on F-actin foci formation compared with that of WASp deficiency (6).

T cell extravasation across the endothelial cell lining is essential for the recruitment of effector T cells to sites of infection, immunosurveillance, and tolerance (48). Dock8<sup>–/–</sup> and Dock8<sup>gmo/gmo</sup> Th1 effector cells expressed selectin ligands normally and adhered normally to immobilized VCAM-1 and E-selectin. It has been established that circulating T cells constitutively express VLA-4 integrins in conformations that recognize VCAM-1 under flow conditions (49). The comparable adhesion of Dock8<sup>–/–</sup>, Dock8<sup>gmo/gmo</sup>, and WT Th1 cells to VCAM-1 suggests that expression of these active conformations by T cells is independent of DOCK8 activity. In contrast, Dock8<sup>–/–</sup> Th1 cells adhered significantly less to ICAM-1 compared with WT Th1 cells under shear flow conditions. Although Dock8<sup>gmo/gmo</sup> Th1 cell adhesion was lower than that of WT Th1 cells, the decrease did not reach statistical significance. These results suggest that DOCK8 functions, beyond its GEF activity, contribute to optimal LFA-1-dependent Th1 cell adherence to ICAM-1 under shear flow conditions. In this respect, DOCK8 is known to interact with talin and protein tyrosine kinase 2 (PYK2), both of which play a role in integrin-mediated cell adhesion (19, 50–52). Post-arrest spreading of both Dock8<sup>–/–</sup> and Dock8<sup>gmo/gmo</sup> effector cells on ICAM-1 coimmobilized with SDF-1α was reduced compared with WT cells. A similar spreading defect has been observed in WASp- and WIP-deficient T cells (40, 53). Importantly, Th1 cells that lacked DOCK8 or expressed a DOCK8-mutant deficient in GEF activity showed a striking reduction in their ability to transmigrate across a TNF-α-activated murine endothelial cell monolayer, a scenario that mimics the egress of effector T cells from blood vessels into an inflamed tissue. Moreover, both DOCK8-deficient and DOCK8-mutant T cells were defective in their ability to home to LNs. Defects in TEM have been documented in WASp- and WIP-deficient T cells (54, 55) as well as in LFA-1 and ICAM-1–deficient T cells (56). The reported normal homing of mature thymocytes from Dock8<sup>gmo/gmo</sup> mice to LNs (34) may reflect differences between mouse strains and between thymocytes and splenic T cells. Decreased contact with the substrate due to defective spreading, defective formation of actin foci, podosome equivalents important for pushing through endothelial cells, and reduced mecanotransduction may all contribute to the impaired TEM and defective homing to LNs of DOCK8-deficient T cells.

In summary, our data demonstrate that a DOCK8-WIP-WASP complex links the TCR to the actin cytoskeleton. The spatial proximity of DOCK8, WASp, and actin in this complex ensures that CDC42 activated by DOCK8 following TCR ligation drives WASp-mediated actin polymerization, as illustrated in Figure 9. Moreover, we show that DOCK8-deficient T cells share with WASp-deficient T cells a number of actin cytoskeletal defects that include an attenuated subcortical actin cytoskeleton, reduced F-actin content, defective TCR-driven actin assembly, actin foci formation, and mecanotransduction, and impaired TEM and homing to LNs. These defects probably contribute to the susceptibility of DOCK8-deficient and WASp-deficient patients to infections, allergy, autoimmunity, and malignancy. Collectively, our findings provide a mechanistic explanation for the shared features of DOCK8 and WASP deficiencies.

Methods

Mice. War<sup>+/+</sup> mice, a gift of Scott Snapper (Boston Children’s Hospital, Boston, Massachusetts, USA) have been previously described (57). Homozygous knockin WIPAABB mice that express a WIP mutant lacking the actin-binding domain (ABD) (aa 43–54) are described in ref. (7). The generation of Dock8<sup>–/–</sup> mice, which carry a homozygous knockin c.C1074T mutation that introduces a stop codon in exon 9 of the endogenous Dock8 gene is described in Supplemental Figure 1. This mutation recapitulates a mutation in a DOCK8<sup>–/–</sup> patient (16). Mice were generated using standard techniques and mated with mice expressing the Flp recombinase under the control of the Rosa promoter to remove the neo gene. The homozygous mutant mice, which express no detectable DOCK8 protein and are designated Dock8<sup>–/–</sup> mice, were then bred for 10 generations on the C57BL/6 background. Dock8<sup>gmo/gmo</sup> mice, generated by N-ethyl-N-nitrosourea mutagenesis from a pure...
C57BL/6 background, carry a homozygous S1827P mutation in the DOCK8 DHR2 domain as previously described (11). WT C57BL/6 mice (Charles River Laboratories) were bred in our facility. Mice were sacrificed at 7 to 12 weeks of age for harvesting of LN and spleen and at 8 to 12 days of age for harvesting of heart endothelial cells. All mice were housed under specific pathogen-free conditions.

Patients. Blood samples were obtained from healthy volunteers, 3 DOCK8-deficient patients with homozygous mutations in DOCK8 (c. T181A p.Ser135X, deletion of exons 1-3, and deletion of exons 10-12) that abolish DOCK8 expression, and 2 WAS patients with mutations in WAS (c.G291A p.R86H, and c.559+5G>A donor splice-site mutation in intron 6) that abolish WASP expression.

Cell isolation. Human peripheral blood mononuclear cells (PBMCs) were isolated from whole blood by Ficoll-Paque PLUS density gradient centrifugation (GE Healthcare). T cells were purified from PBMCs by negative selection using a Pan T Cell Isolation Kit II (Miltenyi Biotech). Mouse LN and spleens were harvested and processed into single-cell suspensions. Splenocytes were pelleted by centrifugation. Mouse T cells were purified by negative selection using a mouse Pan T Cell Isolation Kit II and CD4+ T Cell Isolation Kit, respectively (Miltenyi Biotech).

Cells cultures. The T cell acute lymphocytic leukemia (T-ALL) cell line DND41 (58) was obtained from the ATCC and maintained in RPMI 1640 supplemented with 10% FBS, 100 U/ml penicillin G, and 100 μg/ml streptomycin. Murine CD4+ T cells were derived from naive T cells by anti-CD3ε mAb (mAb 145-2C11) and anti-CD28 (mAb E18) stimulation (both from BioLegend) in the presence of anti-IL-14 mAb (clone 11B11; BioLegend) and recombinant IL-12 and IL-2 (R&D Systems) as previously described (36, 59). Murine heart endothelial cells (MHECs) were prepared from 8- to 12-day-old WT C57BL/6 mice as previously described (36, 59).

Cell stimulation. Human T cells were incubated with 10 μg/ml mouse anti-human CD3 mAb OKT3 (eBioscience) for 10 minutes at 37°C, followed by cross-linking with 20 μg/ml F(ab′)2 goat anti-mouse IgG second antibody (Jackson ImmunoResearch Laboratories) for the times indicated in the figures and figure legends at 37°C. Mouse T cells, rat anti-mouse CD3 mAb KT3 (Serotec) and goat F(ab′)2 anti-rat IgG second antibody (Jackson ImmunoResearch Laboratories) were used.

Cell transfection. HEK293 cells were cotransfected using the lipofectamine LTX and PLUS reagents from Invitrogen with plasmids (vectors) as previously described (36, 59). Murine heart endothelial cells (MHECs) were prepared from 8- to 12-day-old WT C57BL/6 mice as previously described (36, 59).

Flow cytometry. Standard flow cytometric methods were used to evaluate cell-surface marker expression using the following Abs (clones): PE-Cy7 anti-mouse CD44/L-selectin ligand IM7 mAb and PE-Cy7 anti-mouse CD49d/L-selectin ligand IM7 mAb from Biolegend; PE mouse anti-CD11a/LFA-1a M17/4 mAb from eBioscience; and PSGL-1/α1-selectin ligand 2PH1 mAb from BD Biosciences. T cells activated under Th1-polarizing conditions were processed for intracellular staining using a BD Cytofix/Cytoperm Fixation/Permeabilization Solution Kit (BD Biosciences) and stained with eFluor 450 anti-mouse IFN-γ mAb XMGl.2 (eBioscience) to corroborate the differentiation of Th1 samples. Cells were acquired on a BD LSRFortessa cell analyzer using FACSDiva software (BD Biosciences). Analyses were performed using FlowJo software (Tree Star).

Immunofluorescence and confocal microscopy. DND41 cells (5 × 10^5) were plated on fibronectin-coated coverslips (20 μg/ml in PBS) overnight and then fixed with 4% paraformaldehyde for 10 minutes at room temperature. Fixed cells were permeabilized with 0.2% saponin–0.2% BSA in PBS for 20 minutes at room temperature and incubated overnight at 4°C with the following primary Abs: rabbit polyclonal anti-DOCK8 H-159 (Santa Cruz Biotechnology Inc.) paired with mouse anti-WASP mAb F-9 (Santa Cruz Biotechnology Inc.) or mouse anti-WIPI mAb D12C5 mAb; mouse anti-Myc mAb 9E10 (05-419/9E10; EMD Millipore); mouse anti-gamma IFN mAb XMG1.2 (eBioscience) to corroborate the differentiation of Th1 samples. Cells were acquired on a BD LSRFortessa cell analyzer using FACSDiva software (BD Biosciences). Analyses were performed using FlowJo software (Tree Star).
Structural modeling of the SI1827P mutation. The structural flexibility of WT DOCK8 and the SI1827P mutant were analyzed by the CABS-FLEX server (http://biocomp.chem.uw.edu.pl/CABSFlex/) (62). DOCK8 fragment 1792-1879 from the 3vh structure was used in the simulation. For each simulation, 12 clustered structure models were obtained and analyzed through residue fluctuation and dynamic movements. Coot (63) and Pymol (64) were used in structural visualization and presentation.

DOCK8 guanine nucleotide exchange activity assay. The DHR2 containing C-terminal regions of WT or mutant DOCK8 (aa 1639-2071, WT, or SI1827P) fused to Gaussia luciferase secretory peptide were cloned into the pCDNA6/V5-His (Invitrogen, Thermo Fisher Scientific) mammalian expression vector and expressed in HEK293 cells. The Histagged recombinant proteins were purified using a TALON Gravity Column Purification Kit (Clontech). Nucleotide exchange was determined using a fluorescence-based RhogEF exchange activity assay kit according to the manufacturer’s instructions (Cytoskeleton). Briefly, CDC42-GDP was mixed with MANT-GTP [2'-3'-O-(N'-methylanthraniloyl)] guanosine-5'-O-triphosphate, trisodium salt] in the exchange buffer and brought to room temperature. The DHR2 domains of WT or SI1827P mouse DOCK8 (5 μM) were added to the mixture, and the fluorescence (excitation [ex] = 360 nm and emission [em] = 440 nm) was measured over time using a spectrofluorometer (LS50B; PerkinElmer).

EM. Glass coverslips were coated with 1 mg/ml poly-L-lysine (Sigma-Aldrich) for 1 hour at room temperature, followed by 10 μg/ml rat anti-mouse CD3 mAb OKT3 (Serotec) in a humidified chamber overnight at 4°C. T cells were adhered to anti-CD3-coated coverslips by centrifugation at 280 × g for 5 minutes at 4°C, stimulated for 30 minutes at 37°C, and mechanically unroofed with poly-L-lysine-coated coverslips in PHEM buffer (60 mM PIPES, 25 mM HEPES, 10 mM EGTA, 2 mM MgCl2, pH 6.9) containing 1 μM phallacidin and protease inhibitors. Unroofed cells were briefly washed in PHEM buffer and fixed with 1% glutaraldehyde in PHEM buffer for 10 minutes. The coverslips were washed extensively in distilled water, rapidly frozen on a helium-cooled copper block, freeze dried in a Cressington CFE-50 apparatus (Cressington), and rotary coated with 1.5 nm tantalum/tungsten at a 25° angle and 4 nm carbon at 90° without rotation. Specimens were examined and photographed in a JEOL 1200 EX tantalum/tungsten at a 25° angle and 4 nm carbon at 90° without rotation. Coot (63) and Pymol (64) were used in structural visualization and presentation.

Determination of cellular F-actin content. Purified splenic CD3+ T cells were rested in RPMI-1640 for 2 hours on ice. Cells were then incubated with 10 μg/ml rat anti-mouse CD3 mAb KT3 (Serotec) for 30 minutes on ice and stimulated by cross-linking with 20 μg/ml F(ab′)2 goat anti-rat IgG secondary Ab (Jackson ImmunoResearch Laboratories) for the times indicated in the figure 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 FITC-labeled phalloidin (Sigma-Aldrich), and F-actin content was measured with a BD LSRFortessa cell analyzer.

Immune synapse formation, actin foci formation, and CasL phosphorylation. CD4+ T cells freshly isolated from WT, Dock8−/−, and Dock8Δp130 mice were incubated in glass chambers (MatTek) precoated with 10 μg/ml anti-CD3 (2C11) and 1 μg/ml ICAM-1 (extracellular fragment) for 5 minutes at 37°C (6). 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, Thermo Fisher Scientific) and anti-p130Cas (pY165) phospho-specific Ab (Cell Signaling Technology) overnight, followed by incubation with Alexa Fluoro 647- donkey anti-rabbit secondary Ab for 1 hour. Cells were then imaged using total internal reflection fluorescence (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. For the assessment of “foci” actin features within the overall polymerized F-actin network, phalloidin images were processed further: raw phalloidin (actin) images were subjected to a 1.6-×1.6-micron rolling box high-pass filter, and the resultant high-pass filter image was subtracted from the original raw image, as described previously (6). The final subtracted images were used for actin foci measurements. The intensities of actin foci and p-CasL as well as the synapse areas were measured using ROI tools in ImageJ; the measured values were then analyzed for statistical significance as well as plotted using GraphPad Prism (GraphPad Software).

For T cell-endothelial monolayer assays, the bEnd3 cell line (ATCC), which constitutively expresses ICAM-1 and VCAM-1, was loaded with TSST-1 and SEB (2 μg/ml each) for 1 hour, washed, and incubated with CD4+ T cells derived from WT, Dock8−/−, and Dock8Δp130 mice for 15 minutes at 37°C. The cells were fixed using PHEM buffer and processed for immunofluorescence, as described above. Subsequently, cells were imaged using a spinning-disc confocal microscope, and a maximum intensity projection encompassing 1 micron at the cell-cell conjugate interface was analyzed for F-actin foci and pCasL levels.

T cell adhesion and transmigration under defined laminar shear flow conditions in vitro. Live-cell imaging of leukocyte adhesion was recorded by a video camera coupled to a Nikon TE2000 inverted microscope equipped with a 20×/0.75 NA phase-contrast objective and VideoLab software (Mitov) (36, 59). Briefly, in vitro polarized WT, Dock8−/−, and Dock8Δp130 Th1 cells (5 × 105 in 100 μl) were suspended in Dulbecco's PBS containing 0.1% (w/v) BSA and 20 mM HEPES, pH 7.4, and were drawn across immobilized murine E-selectin-Fc (20 μg/ml), ICAM-1-Fc (20 μg/ml), and VCAM-1-Fc (5 μg/ml) chimeric proteins (R&D Systems) initially at a shear stress level of 1.0 dynes/cm², and subsequently, the flow rate was reduced at 3 minutes to 0.8 dynes/cm² and finally to 0.5 dynes/cm² 3 minutes later (59). Th1 cell adhesion was determined in 5 separate fields at the last minute of each flow rate and reflects the accumulation of adherent cells over the range of shear stress examined. At the end of the assay, each adherent T cell on ICAM-1 and 250 ng/ml SDF-1α was assessed for its ability to spread. Fully spread cells were scored as 2 points, partially spread cells as 1 point, and round, unspread cells scored as 0 points. The sum of scores was divided by the total number of adherent cells to calculate average scores for each condition.

Polarized Th1 cells were drawn across confluent TNF-α-activated (100 ng/ml, 5 hours) MHECs grown on glass coverslips at 37°C in an in vitro flow chamber at an estimated shear stress level of 0.8 dynes/cm² (59). Live-cell imaging of Th1 cell TEM was performed using a digital imaging system coupled to a Nikon Eclipse Ti inverted microscope (Nikon) equipped with a 20×/0.75 NA DIC objective. Time-lapse videos were acquired using MetaMorph software (Molecular Devices). The total number of accumulated Th1 cells was determined by counting total adherent and transmigrated cells per coverslip. The percentage of TEM = total transmigrated leukocytes/(total adhered + transmigrated leukocytes) × 100.

In vivo homing of T cells to LNs. In vivo homing of T cells was performed as previously described (7). T cells were purified from spleen by negative selection using a Pan T Cell Isolation Kit II (Miltenyi Bio-
WT T cells were labeled with 20 μg/ml Alexa Fluor 555 carboxylic acid succinimidyl ester (Life Technologies, Thermo Fisher Scientific) or Alexa Fluor 488 carboxylic acid succinimidyl ester (Life Technologies, Thermo Fisher Scientific) for 15 minutes at 37°C. Dock8<sup>−/−</sup> or Dock8<sup>−/−</sup> T cells were labeled with Alexa Fluor 488. Labeled cells were then centrifuged, washed, and suspended in RPMI-1640. A total of 10<sup>5</sup> Alexa Fluor 555-labeled WT T cells were mixed with an equal number of either Alexa Fluor 488-labeled WT, Dock8<sup>−/−</sup>, or Dock8<sup>−/−</sup> T cells. Cells were injected i.v. into WT recipient mice. After 1 hour, the mice were sacrificed, and LNs were harvested. Single-cell suspensions were analyzed by flow cytometry, and the percentages of Alexa Fluor 488- and Alexa Fluor 555-labeled cells were determined. The T cell homing index was calculated as the ratio of Alexa Fluor 488-labeled cells to Alexa Fluor 555-labeled cells and divided by the ratio of WT to WT cells. To stain for cell death markers, cells were incubated with fixable viability dye eFluor 780 (ebioscience) for 30 minutes at 4°C, followed by APC-conjugated annexin V (ebioscience) for 15 minutes at room temperature. The percentage of annexin V<sup>+</sup> (apoptotic), eFluor 780<sup>+</sup> (dead), and eFluor 780<sup>−</sup> live cells was calculated using FlowJo software.

**Statistics.** Statistical analysis of the data using 2-tailed Student’s t test or 2-way ANOVA for multiple groups was performed with GraphPad Prism (GraphPad Software version 7.01). A P value of less than 0.05 was considered significant.

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**Study approval.** All mouse studies were approved by IACUC of Boston Children’s Hospital. All human participants gave written informed consent to participate in a research protocol approved by the Committee on Clinical Investigation at Boston Children’s Hospital.

**Author contributions**

EJ, MH, MT, ML, SK, NR, SU, VA, and JH performed the experiments and analyzed the data. QQ and HW did the molecular modeling. EJ, MJM, KLR, and CCG generated the mutated mice. WAH provided patients’ samples. DC generated the CSab. JH, DJI, FWL, and RSG designed and supervised the research. EJ, MH, and RSG wrote the manuscript.

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