Actin polymerization-dependent activation of Cas-L promotes immunological synapse stability

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The immunological synapse formed between a T-cell and an antigen-presenting cell is important for cell–cell communication during T-cell-mediated immune responses. Immunological synapse formation begins with stimulation of the T-cell receptor (TCR). TCR microclusters are assembled and transported to the center of the immunological synapse in an actin polymerization-dependent process. However, the physical link between TCR and actin remains elusive. Here we show that lymphocyte-specific Crk-associated substrate (Cas-L), a member of a force sensing protein family, is required for transport of TCR microclusters and for establishing synapse stability. We found that Cas-L is phosphorylated at TCR microclusters in an actin polymerization-dependent fashion. Furthermore, Cas-L participates in a positive feedback loop leading to amplification of Ca2+ signaling, inside-out integrin activation, and actomyosin contraction. We propose a new role for Cas-L in T-cell activation as a mechanical transducer linking TCR microclusters to the underlying actin network and coordinating multiple actin-dependent structures in the immunological synapse. Our studies highlight the importance of mechanotransduction processes in T-cell-mediated immune responses.

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Most adaptive immune responses require activation of T cells.1–3 The process of T-cell activation involves a multi-step mechanism that begins with weak adhesion and stimulation of the T-cell receptor (TCR) leading to adhesion strengthening and formation of a highly organized immunological synapse.4–7 Spatial organization of the immunological synapse requires f-actin,8–10 myosin IIA,11–13 microtubules and dynein,14 and the endosomal sorting complexes required for transport.15,16 There is growing evidence supporting a physical link between TCR microclusters and the actin cytoskeleton, but this most fundamental connection is the most poorly understood.17–20 TCR and integrin adhesion molecules organize actin polymerization,21–23 which drives transport of distinct TCR and integrin microclusters toward the center of the synapse.24–27 This can be modeled as a ‘frictional’ process as the bulk flow of f-actin is faster than the movement of microclusters, but the molecular basis of the friction-like effect is not known. Furthermore, the TCR and integrins have been implicated in mechanotransduction at the immunological synapse,28–31 but how the TCR participates in mechanotransduction remains unknown.

The temporal and spatial localization of signaling proteins at the immunological synapse correlates with T-cell activation. Proper assembly and localization of signaling complexes is often mediated by scaffold proteins.32 These multidomain adaptors have several binding partners, and by bringing them into close proximity they facilitate protein–protein interactions and signal propagation. Although many scaffold proteins are essential for T-cell activation, how they become activated and how they regulate T-cell signals is largely unknown. We recently described a model for actin-dependent stretch of the mechanosensing protein p130 Crk-associated substrate (p130Cas)33 used by cells in sensing their physical environment, in integrin adhesions and during migration.34–37 p130Cas belongs to a family of adaptor proteins that share a flexible Cas substrate domain that unfolds in response to force exposing Src-family kinase phosphorylation sites.38 The Cas family member most abundant in T cells is Cas-L (also called Hef1 and NEDD9).39,40 Cas-L contains a central substrate domain with 13 repeated motifs each containing a tyrosine residue (YxxP), flanked on one side by an N-terminal SH3 domain, and on the other by a proline-rich four-helix bundle and a Src-family kinase-binding domain with consensus-binding sites YDYVHL and RPLPSP, for SH2 and SH3 domains, respectively. Although Cas-L does not have any enzymatic activity, it has been implicated in a diverse set of physiological and pathological contexts in different cell types.41–47 This functional versatility underscores the importance of Cas-L in mediating receptor-proximal interactions and propagating local stimulatory signals that lead to global changes in cell behavior.32,48

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Seo et al. observed that Cas-L−/− mice have a decreased number of lymphocytes in peripheral lymphoid organs, a thin marginal zone, an impaired TCR-induced response to immunization with a specific peptide, and perturbed integrin-mediated adhesion in comparison with wild-type mice. Previous studies reported that following TCR and integrin stimulation the Cas-L substrate domain tyrosine motifs are transiently phosphorylated. Phospho-Cas-L recruits the adaptor protein CrkL that is constitutively associated with the guanine exchange factor C3G, which in turn is involved in the activation of the integrin regulating Rap1 GTPase. Furthermore, mutational analyses suggest that the substrate domain of Cas-L is required for migration in response to TCR and integrin stimulation. However, the mechanisms leading to assembly of the ternary complex Cas-L-CrkL-C3G are unknown and whether Cas-L regulates T-cell activation is unclear.

To study T-cell activation in the context of the immunological synapse, a system widely used is based on glass-supported planar lipid bilayers that mimic the surface of an antigen-presenting cell. Using bilayers, we and others recently reported that myosin IIa contractility at the immunological synapse is involved in the transport of TCR microclusters, and in sustaining Src-family kinase signals, Ca²⁺ flux, and interleukin-2 (IL-2) secretion. Interestingly, both RNA interference depletion and pharmacological inhibition of myosin IIa activity reduced phosphorylation of Cas-L at the immunological synapse. However, it remains unknown if Cas-L has a role in the formation of the immunological synapse and how Cas-L integrates initial TCR stimulation events with integrin activation and actomyosin contractility.

The interplay between TCR and integrin signaling is complex and it is unclear how it translates into f-actin polymerization dynamics. A widely established paradigm for integrin-signaling studies in T cells is the lymphocyte function activation 1 (LFA-1), a β2 integrin that is critical for immunological synapse formation, and T-cell adhesion and activation. TCR stimulation leads to inside–out activation of the TCR.

Figure 1 TCR microcluster movement is impaired in the absence of Cas-L. (a) Western blot of total cell lysates from CD8⁺ T cells isolated by negative selection from spleen of Cas-L-null (Cas-L−/−) mice, and from the control Cas-L heterozygous littermates (WT). A monoclonal antibody for Cas-L was used to assess the presence of Cas-L. (b) Cells were fixed in 2% paraformaldehyde 15 min after seeding, and images of synapses formed in both cell types were acquired in both channels. Red = TCR; blue = ICAM-1; yellow = synapse outline. Scale bar = 2 μm. (c) Radial profiles of TCR and ICAM-1 fluorescence intensities at the synapses of WT and Cas-L−/− cells (radial sweep averaged from at least 50 cells in each cell type and two independent experiments). Solid and dashed lines represent WT and Cas-L−/− cells, respectively. Red = TCR, blue = ICAM-1. P-value < 0.05 (d) Time-lapse images of TCR microcluster formation and translocation at the immunological synapse. Yellow arrowheads highlight individual TCR microcluster position at the synapse at different time points. Freshly isolated CD8⁺ T cells from spleen of WT or Cas-L−/− mice were seeded on a bilayer embedded with fluorescently labeled antibody to TCR (red) and ICAM-1 (blue), and imaged in a TIRF microscope to follow immunological synapse formation. Simultaneous imaging of four different fields of the bilayer was performed with a × 100 objective, with an acquisition rate of 13 s between frames. Scale bar = 5 μm. (e) Average speed of translocation of individual TCR microclusters during synapse formation. Mean ± s.e.m. represent two independent experiments. At least five individual microclusters per cell were analyzed in at least three cells. One asterisk indicates P-value < 0.05. (f) Change in mean fluorescence intensity of TCR microclusters during synapse formation in WT and Cas-L−/− cells. Mean values of four cells of each type, pooled from two independent experiments. P-value < 0.0005. (g) Change in area occupied by TCR microclusters at the synapse of WT and Cas-L−/− cells analyzed in D (mean values of four cells of each type, pooled from two independent experiments).
LFA-1, which undergoes a conformational change into a higher affinity state to bind its ligand intercellular adhesion molecule 1 (ICAM-1). Interestingly, the strength of the LFA-1/ICAM-1 interaction is regulated by a mechanical feedback loop, but whether there is a mechanical link between TCR and LFA-1 remains unknown. However, it is known that activation of Rap1 is critical for LFA-1-mediated T-cell adhesion, indicating that f-actin-based contractility and Cas-L might provide a mechanical connection between TCR and LFA-1.

Among the vast number of molecules that regulate actin polymerization in T cells is the Wiskott-Aldrich Syndrome protein (WASp). WASp is a multidomain scaffold protein containing binding sites for actin and the Arp2/3 complex, which is an actin nucleation factor responsible for branching of actin filaments. WASp is recruited to TCR microclusters and can become activated via several pathways. Activated WASp then recruits Arp2/3 complex and initiates actin polymerization from existing filaments. Furthermore, WASp favors stability of the immunological synapse, and treatment of WASp−/− T cells with an inhibitor of protein kinase C theta (PKCθ) restored immunological synapse stability, suggesting opposing effects within a single pathway for the activity of those two proteins at the immunological synapse.

Here, we wished to determine whether Cas-L has a role in T-cell activation in the context of the immunological synapse using T cells from Cas-L−/− mice. We report a role for actin polymerization in Cas-L activation at the immunological synapse. In analogy to p130Cas, we hypothesized that a mechanical feedback loop between TCR-mediated Cas-L activation, actin polymerization and LFA-1 activation might regulate the processes of adhesion, migration and activation of T cells.

RESULTS

Cas-L promotes TCR microcluster translocation

To investigate a potential role for Cas-L in T-cell activation we measured the accumulation of TCR microclusters and ICAM-1 at the immunological synapse of Cas-L−/− T cells compared with control cells. No Cas-L protein was detected in T cells from Cas-L−/− mice (Figure 1a). We seeded cells on supported lipid bilayers embedded with laterally mobile fluorescently labeled anti-CD3ε and ICAM-1, fixed them after 15 min, imaged them and plotted the average radial profile of intensities of TCR and ICAM-1 over the entire synapse of both cell types (Figures 1b and c). In control cells, ICAM-1 stays distributed in a pericentral ring at the synapse, but in contrast, ICAM-1 accumulation in Cas-L−/− cells was severely compromised and its pericentral ring organization was lost (Figure 1b blue channel, and Figure 1c blue lines). Further, in Cas-L−/− T cells, accumulation of TCR at the center of the synapse was significantly decreased compared with control cells (Figure 1b red channel, and Figure 1c red lines).

Interestingly, a fraction of the TCR microclusters remained arrested at the synapse periphery in Cas-L−/− T cells (Figure 1b red channel). Thus, we asked whether the decreased accumulation of TCR at the immunological synapse of Cas-L−/− T cells was a consequence of deficient transport of TCR microclusters from the periphery to the central domain where extracellular vesicles are formed. In control cells, TCR assembled into microclusters at the periphery of the synapse within seconds of interaction with the bilayer and rapidly moved toward the center, with TCR accumulating in one bright focus at the center of the synapse (Figure 1d top panel; Movie 1 top panel). In contrast, in Cas-L−/− T cells, movement of TCR microclusters toward the center of the synapse was impaired, and ~30% of TCR microclusters remained dispersed in the periphery and did not move to the center (Figure 1d bottom panel; Movie 1 bottom panel).

Importantly, in Cas-L−/− T cells, TCR microclusters exhibited a lower speed of translocation (1.92 ± 0.20 μm min⁻¹) and a lower mean square displacement (path length 1.78 ± 0.16 μm) than in control cells, where microclusters moved at an average speed of 2.78 ± 0.37 μm min⁻¹, with a path length of 2.82 ± 0.38 μm (Figure 1e). Furthermore, we tracked individual TCR microclusters, measured their fluorescence intensities and areas and plotted the corresponding averages throughout the course of synapse formation (Figures 1f and g). Individual TCR microclusters from Cas-L−/− cells exhibited, on average, decreased intensity and area compared with TCR microclusters from control cells, suggesting that in the absence of Cas-L TCR microclusters have a lower rate of maturation compared with control cells.

Activated Cas-L colocalizes with TCR microclusters at the periphery of the immunological synapse

As localization of TCR-signaling proteins to the immunological synapse correlates with T-cell activation, we examined if activation of Cas-L was spatially and temporally coupled to TCR microcluster assembly at the immunological synapse as previously suggested. To assess the level of activation of Cas-L we used an antibody that recognizes phosphorylated tyrosine motifs located in the substrate domain of Cas-L. We seeded cells on anti-CD3ε/ICAM-1-embedded bilayers, fixed them after 2 min and stained them for phospho-Cas-L (Figure 2a). Importantly, phospho-Cas-L colocalized with nascent TCR microclusters at the periphery of newly formed synapses (Figure 2a yellow arrows on the bottom-right panel), indicating that Cas-L phosphorylation is spatially coupled to TCR microcluster assembly at an early stage of immunological synapse formation.

In order to understand how Cas-L becomes activated at TCR microclusters we treated T cells with a small molecule inhibitor of Src-family kinases, PP2. We found that inhibition of Src-family kinases by PP2 led to a significant decrease in the level of phospho-Cas-L at the immunological synapse (Figures 2b and c). As shown in Figure 2b, in the presence of PP2, the LFA-1-dependent adhesion was nearly completely eliminated, as previously reported, but TCR clustering still occurred, suggesting that cell adhesion and Cas-L phosphorylation might be related. The level of phospho-Cas-L at TCR clusters was greatly decreased with PP2 treatment, but as revealed by Pearson’s correlation coefficient analysis (Supplementary Figure 1) the phospho-Cas-L signal was more strongly co-localized with TCR clusters in PP2-treated cells than in the absence of PP2, as TCR clusters were relatively more resistant to inhibition by PP2 compared with LFA-1/ICAM-1 interaction.

We further pursued the relationship between phospho-Cas-L and TCR microclusters. It is known that TCR ligation leads to the recruitment and activation of Src-family kinase Lck, which in turn phosphorylates the immunoreceptor tyrosine-based activation motifs located on the CD3 chains of the TCR complex. Then, the zeta-chain-associated protein kinase 70 kDa (ZAP70) binds to the doubly phosphorylated zeta-chain immunoreceptor tyrosine-based activation motifs, exposing phosphorylation sites on its kinase domain activation loop, which in turn are phosphorylated by Lck. Thus, the localization of ZAP70 at TCR microclusters and the phosphorylation status of ZAP70 correlate with TCR activation and Lck activity, respectively. To determine whether different baseline levels of TCR activation could account for our observations in Cas-L−/− T cells, we compared the levels of ZAP70 localized with TCR microclusters in...
both cell types and detected no significant changes (Supplementary Figure 1A). We also compared the baseline activity levels of the Src-family kinase Lck by probing for the phosphorylated form of ZAP70 at TCR microclusters and detected no differences between Cas-L−/− T cells and control cells (Supplementary Figure 1B). As the baseline levels of TCR activation and Lck activity were not altered in Cas-L−/− T cells compared with control cells, the defects observed in Cas-L−/− T cells were not due to differences in the TCR proximal tyrosine kinase cascade up to ZAP70 recruitment and activation.

Noteworthy, treatment with a selective Lck inhibitor76 led to a significantly greater decrease in mean intensity levels of phospho-Cas-L than treatment with the selective ZAP70 inhibitor piceatannol,74 suggesting that Cas-L phosphorylation depends on Lck activity (Figures 2d and e).

These results demonstrate that Lck phosphorylates tyrosine residues in the Cas-L substrate domain immediately downstream of TCR stimulation. Thus, although Cas-L is dispensable for the assembly of TCR microclusters, it is required for the efficient transport of newly formed TCR microclusters to the center of the immunological synapse. As the centripetal transport of microclusters correlates with sustained TCR signaling,5,8 these findings suggest that Cas-L might regulate signals downstream of the TCR proximal tyrosine kinase cascade.

Ca2+ release from intracellular stores is impaired in Cas-L−/− T cells
As myosin Ila-dependent transport of TCR microclusters is required to sustain TCR signals in the bilayer model,11,12,77 in particular Ca2+

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**Figure 2** Phosphorylated Cas-L colocalizes with new TCR microclusters at the periphery of the immunological synapse. (a) TIRF images of synapses from mouse spleen CD8+ T cells fixed at 2 min after seeding on bilayers embedded with fluorescently labeled antibody to TCR (red) and ICAM-1 (blue), and stained with an antibody specific for phosphotyrosine repeats in the substrate domain of Cas-L (pCas-L, green). Yellow boxes are expanded on the right panel. Arrowheads highlight colocalization of TCR microclusters and phospho-Cas-L. Scale bar = 2 μm. (b) Cas-L phosphorylation at the synapse depends on Src-family kinase activity. Scale bar = 2 μm. (c) Cas-L recruitment to TCR microclusters depends on Src-family kinases activity. Dot plots display the median (central line), ± 1.5 × interquartile range (whiskers). One asterisk represents P-value ≤ 0.05; two asterisks represent P-value ≤ 0.005; three asterisks mean P-value ≤ 0.0005; data represent at least 30 cells from two independent experiments. Graphs display 30 cells from two experiments pooled in one graph. (d and e) Cas-L phosphorylation at the immunological synapse is significantly decreased after inhibition of Lck by treatment with piceatannol or an Lck selective inhibitor. Mean values ± s.e.m. are representative of at least 30 cells (for each condition) from two independent experiments. Scale bar = 2 μm. One asterisk indicates P-value ≤ 0.05; three asterisks P-value ≤ 0.0005.
signaling, we wanted to determine whether the defective TCR microcluster transport observed in Cas-L−/− T cells was correlated with any defects in Ca^{2+} signaling. To visualize changes in the concentration of free intracellular Ca^{2+}, we loaded cells with the fluorescent Ca^{2+} probe Fluo-4 as described in Methods. We then seeded cells on anti-CD3ɛ/anti-CD28/ICAM-1-coated coverslips and imaged Fluo-4 fluorescence intensity changes (Movie 2). As shown in Figure 3a and plotted in Figure 3b, when cells touched the antibody-coated coverslip they exhibited a sharp increase of Fluo-4 fluorescence intensity corresponding to Ca^{2+} released from intracellular stores, reaching a maximum Fluo-4 fluorescence value (imax1), followed by a decrease of fluorescence intensity down to a sustained lower level, which is dependent upon influx of extracellular Ca^{2+} (Movie 2). The cells were then treated with ionomycin to determine the maximum Fluo-4 fluorescence value (imax2) for scaling, which is important as Fluo-4 loading differs between cells. For each cell the pre-stimulation baseline was set to 0 and the imax2 value was set to 1 and other values normalized accordingly. We observed that Cas-L−/− cells exhibit a lower value of imax1 compared with control cells, but the normalized value of sustained Ca^{2+} was similar in both cell types (Figures 3c and d). Raw values of imax2 in wild-type and Cas-L−/− cells were similar suggesting no significant differences in dye loading or Ca^{2+} homeostasis. We also observed that ionomycin quickly halted Cas-L−/− T-cell migration, suggesting that the machinery linking Ca^{2+} to migration arrest is intact in the absence of Cas-L (Movie 2 bottom panel). Together these observations suggest that the defect in imax1 Ca^{2+} release observed in Cas-L−/− T cells was due to a TCR-signaling defect.
Cas-L regulates immunological synapse stability via integrin-mediated adhesion

Seo et al. observed that T cells from Cas-L−/− mice exhibit perturbed integrin-mediated adhesion. Integrin-mediated adhesion, namely mediated by LFA-1 ligation by ICAM-1, is critical for immunological synapse formation and enhances sensitivity of T-cell activation to MHC-peptide complexes. We showed in Figure 1c that recruitment of ICAM-1 to the immunological synapse was impaired in Cas-L−/− T cells, relative to control cells. In addition, we noticed that although the number of cells seeded on the bilayers was the same for both cell types (1 × 10^6 cells ml^−1), the population of Cas-L−/− T cells that adhered to the bilayers was significantly decreased in relation to control cells (results not shown). Thus, we wanted to determine whether Cas-L had a role in LFA-1-mediated T-cell adhesion by imaging and comparing synapse stability in Cas-L−/− T cells and control cells seeded on bilayers with anti-CD3ε and ICAM-1. We saw that the population of T cells that adhered to the bilayers followed a bimodal distribution, whereby one fraction of the cell population formed stable persistent synapses that remained in the same initial position during the course of the experiment (15 min), and another fraction was unstable, and polarized and migrated after contacting the bilayers (Figure 4a; Movie 3). The majority of control cells formed stable synapses, whereas most Cas-L−/− T cells failed to form a stable synapse, tended to polarize and exhibited uncoordinated migration (Figures 4b and c; Movie 3). Although more Cas-L−/− T cells migrated than control T cells, the migration speed of Cas-L−/− T cells (3.5 ± 0.51 μm min^−1) was significantly lower than that of control cells (5.3 ± 0.91 μm min^−1) (Figure 4c). Furthermore, in contrast to the directionally persistent migration of control cells, Cas-L−/− cells were less persistent, with frequent turns (Movie 3). As a result of their slower migration speed and more frequent turning, Cas-L−/− cells showed smaller displacement, that is, Cas-L−/− cells traveled shorter distances than control cells in a given time period (Figure 4c). Further, Cas-L−/− T cells revealed defects in early spreading, had significantly smaller synapse area than control cells (Figure 4d), and a larger circularity index (Figure 4e), reflecting their inability to fully spread over the bilayers.

To further understand the relationship between Cas-L and the actin cytoskeleton, we seeded T cells on bilayers with anti-CD3ε and ICAM-1 and compared their Cas-L phosphorylation levels (Figure 5c), suggesting that Cas-L has a role in the substrate domain of Cas-L−/− T cells (Figure 5c). Although control cells were able to form synapses on bilayers with as low as 30 molecules of ICAM-1 per μm^2, Cas-L−/− cells failed to form stable synapses even over threefold the typical level of ICAM-1 (650 molecules of ICAM-1 per μm^2) (Figure 4g). This adhesion defect was not due to lower expression of LFA-1 as the expression levels of LFA-1 were similar in wild-type and Cas-L−/− T cells, as analyzed by flow cytometry (Supplementary Figure 1D).

Altogether, these results show that Cas-L has a role in establishing and maintaining stability of the immunological synapse by promoting inside-out integrin activation leading to T-cell adhesion.

Actin polymerization regulates Cas-L activation

In vivo, T cells undergo an extensive search for antigen in lymphoid tissues by using ameboid migration, a process that requires actin polymerization. The actin cytoskeleton has, in addition, a critical role in immunological synapse formation and regulation of T-cell function. Previous studies using mutational analysis indicated that the substrate domain of Cas-L was required for T-cell migration, but the mechanism of Cas-L activation was not known. Here, we hypothesize that in a similar way to its paralp p130Cas34 Cas-L activation depends on actin polymerization. Thus, we looked at how disruption of actin polymerization by cytochalasin D treatment affected phosphorylation levels of Cas-L substrate domain at the immunological synapse. We saw that cytochalasin D induced a strong decrease in the levels of phospho-Cas-L at the synapse compared with dimethyl sulfoxide (DMSO) carrier, and its effect was stronger at higher doses (Figure 5a), indicating that phosphorylation of the Cas-L substrate domain depends on actin polymerization.

To further understand the relationship between Cas-L and the actin cytoskeleton, we seeded T cells on bilayers with anti-CD3ε and ICAM-1 and compared f-actin organization in Cas-L−/− cells and control cells. Confocal imaging of fluorescently labeled phalloidin staining in Cas-L−/− cells revealed a disorganized cortical f-actin network, smaller lamellipodial area and undefined lamella/lamellipodia boundaries (Figure 5b). The radial profile of the average f-actin intensity at the synapse also revealed a defective f-actin network in Cas-L−/− cells (Figure 5c), suggesting that Cas-L has a role in the organization of the structure of the f-actin network. Furthermore, inhibition of the actin polymerization factor Arp2/3 with a small-molecular inhibitor (CK666) caused a significant decrease in phospho-Cas-L levels at the immunological synapse compared with DMSO carrier treatment (Figure 5d). As Arp2/3 also colocalizes with nascent TCR microclusters at the periphery of the synapse, these data indicate that Cas-L activation at the TCR microclusters is dependent on the actin polymerization-promoting factor Arp2/3.

To further investigate how actin polymerization regulates Cas-L activation, we asked if WASp had a role in Cas-L activation. We seeded WASp−/− T cells and control cells on bilayers with anti-CD3ε and ICAM-1 and compared their Cas-L phosphorylation levels (Figure 5e). We saw significantly lower levels of phospho-Cas-L at the immunological synapse compared with DMSO carrier treatment (Figure 5d). As Arp2/3 also colocalizes with nascent TCR microclusters at the periphery of the synapse, these data indicate that Cas-L activation at the TCR microclusters is dependent on the actin polymerization-promoting factor Arp2/3.
migration observed in Cas-L−/− cells was due to PKCθ driven symmetry breaking. In fact, when we treated Cas-L−/− T cells with an inhibitor of PKCθ activity, we saw that Cas-L−/− cells formed significantly more stable synapses compared with DMSO-treated Cas-L−/− cells (P-value ≤ 0.0005). Values of averages ± s.e.m. are representative of at least 30 cells of each type and three independent experiments. (c) Quantification of adhesion and migration parameters of WT and Cas-L−/− cells (values represent mean ± s.e.m. of at least 30 cells from three independent experiments). (d and e) Cas-L−/− cells exhibit deficient spreading, having smaller contact areas, and more circular contact areas, in comparison with control cells. (f) ICAM-1 recruitment to the synapse is impaired in Cas-L−/− cells, especially at early time points of synapse formation. (g) Providing bilayers with increasing amounts of ICAM-1 is not sufficient to rescue synapse instability exhibited by Cas-L−/− cells. One asterisk: P-value ≤ 0.05; 'n.s.': non-significant; data represent three independent experiments.

**DISCUSSION**

Building an immunological synapse provides a mechanism to assess sensitivity and specificity of the interaction between antigen and TCR. The intrinsically transient TCR-MHC-peptide interaction is organized into clusters that are subjected to physical forces associated with transport as a critical determinant of T-cell signaling. These dynamic processes take place in stable immunological synapse to allow for signal integration and effector functions. Generation of effective recall response may be particularly reliant on stable immunological synapses.

Our results here show that Cas-L is involved in stabilizing the immunological synapse and establishing synapse symmetry. Furthermore, we show that Cas-L is necessary for efficient transport of TCR microclusters from the periphery to the center of the synapse, and without Cas-L, microcluster translocation slows down or is completely arrested. Our findings reveal that both activation of phospho-Cas-L localizes to early TCR microclusters where tyrosine phosphorylation and WASp-dependent actin polymerization occur. We found that disrupting actin polymerization with cytochalasin D or inhibiting actin nucleation factor Arp2/3 led to a significant decrease in phospho-Cas-L levels at TCR microclusters. Finally, we propose a new role for Cas-L as a mechanical link bridging f-actin foci at TCR microclusters, as well as at the lamellipodial f-actin ring facilitated by LFA-1/ICAM-1-dependent adhesion.

Stable immunological synapses are not required for T-cell proliferation, but appear to be critical for asymmetric cell division and functional memory. In vivo analyses with tracking of single T cells by bar-coding have challenged the need for asymmetric division as a drive, but still demonstrate striking heterogeneity in the behavior of individual T-cell clones, which may rely on a spectrum...
of interactions including stable immunological synapses.\(^82\) It has also been proposed that synapse stabilization in vivo may help T cells of lower affinity for an antigen decide whether or not to participate in a response.\(^83\) In particular for a T-cell effector response, the initial free intracellular Ca\(^{2+}\) spike (\(i_{\text{max}1}\)) is critical for rapid arrest of migrating cells and direct cell–cell communication that establishes that response. Here, we saw that Cas-L\(^{-/-}\) CD8\(^{+}\) T cells release only approximately half of their total Ca\(^{2+}\) reserves, which amounts to a decrease of approximately 30% compared with wild-type cells. Remarkably, the proportion of Cas-L\(^{-/-}\) T cells that adhered to anti-CD3\(\varepsilon\)/ICAM-1-embedded supported planar bilayers was only half of that of control cells, and they had a smaller contact area compared with control cells. Furthermore, Cas-L\(^{-/-}\) cells formed unstable synapses that failed to remain symmetric, exhibiting irregular lamellipodial ruffling, and uncoordinated migration. This phenotype might be explained by the defects we saw in the actin cytoskeleton of Cas-L\(^{-/-}\) T cells, which exhibited a disordered cortical f-actin network, and lost the defined border between lamella and lamellipodia structures. In addition, studies have shown that the size of TCR microclusters correlates with T-cell activation.\(^6\) Here, we saw that immunological synapses formed by Cas-L\(^{-/-}\) T cells showed a decreased accumulation of TCR in the central region and the size of microclusters was smaller compared with control cells. Altogether, these data suggest that Cas-L might be important for T-cell survival and proliferation\(^80,81,83\) by establishing immunological synapse stability.

In an effort to understand the mechanism by which Cas-L establishes immunological synapse stability, we looked into a model we previously reported\(^72\) describing how T cells regulate synapse symmetry and migration. There we demonstrated that PKC\(\theta\) activity promotes synapse symmetry breaking leading to migration, whereas

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**Figure 5** Cas-L phosphorylation is dependent on actin polymerization. (a) Actin disruption affects the radial profiles of phospho-Cas-L (pCasL) intensity at the synapse. T cells treated for 30 min with DMSO, or 60 nM and 200 nM Cytochalasin D (CytoD) were seeded on anti-CD3\(\varepsilon\)/ICAM-1-embedded bilayers, allowed to interact with the bilayers for 2 min, fixed with 2% paraformaldehyde and stained with an antibody specific for phosphotyrosine residue repeats in the substrate domain of Cas-L. The synapses were imaged with TIRFM, the levels of mean fluorescence intensity of pCasL were measured, processed and the average radial profiles and standard error of the mean of at least 30 cells per condition were plotted (data represent two independent experiments). (b) Confocal slices from the synapses of WT and Cas-L\(^{-/-}\) cells, and corresponding vertical section profiles. The border between lamella and lamellipodia structures is lost in synapses of Cas-L\(^{-/-}\) cells. Scale bars=2 \(\mu\)m. (c) Radial profiles of f-actin in synapses of WT and Cas-L\(^{-/-}\) T cells indicated lower f-actin accumulation at the synapses of Cas-L\(^{-/-}\) T cells. Cells were seeded on bilayers, fixed at 2 min and stained with fluorescently labeled phalloidin (green). Curves represent the mean±s.e.m. of at least 30 cells from three independent experiments. (d) Phospho-Cas-L levels at the synapse are dependent on Arp2/3 activity. T cells were treated with a small molecular inhibitor of Arp2/3 (CK666), or DMSO as control. Three asterisks: \(P\)-value\(\leq\)0.0005. (e) Phospho-Cas-L levels at the synapse are dependent on WASp activity. WASp\(^{-/-}\) T cells or WT cells were seeded on bilayers, fixed at 2 min, and stained for phospho-Cas-L. (f) Cas-L and PKC\(\theta\) play opposing roles in regulating synapse stability. T cells from Cas-L\(^{-/-}\) mice were seeded on bilayers, and 20 min after seeding a small molecule inhibitor for PKC\(\theta\) (C20) was added to the medium (at 10 \(\mu\)M or 33 \(\mu\)M) and its effect was followed for another 20 min. Graph reflects the change in the number of new synapses formed after C20 treatment relative to the DMSO control. Values represent mean±s.e.m. of at least 30 cells from three independent experiments.
WASp activity favors synapse symmetry and promotes synapse stability. In line with those studies, here we saw that inhibition of PKCθ activity in Cas-L^{−/−} T cells impeded migration and restored synapse stability. Furthermore, activated phospho-Cas-L at TCR microclusters was decreased in WASp^{−/−} T cells, suggesting that Cas-L activation at TCR microclusters is dependent on WASp activity. Altogether, these data further corroborate a role for Cas-L in promoting synapse symmetry and stability.

Despite functioning downstream of WASp and actin polymerization at TCR clusters, Cas-L also contributes to the proper organization of the peripheral f-actin ring in defining the lamella/lamellipodia regions, which suggests that Cas-L is part of a feedback loop mechanism favoring TCR signal amplification and immunological synapse stability. Our colocalization analysis (Supplementary Figure 1F) suggests that both the TCR microcluster actin foci and the lamellipodial f-actin ring that is facilitated by LFA-1-ICAM-1 interaction contribute to phospho-Cas-L activation. It is known that ligation of TCR causes recruitment of active Lck, and recruitment and activation of WASp via the WIP-CrkL pathway, which in turn recruits Arp2/3, thus triggering actin polymerization.65–68 As phospho-Cas-L associates strongly with CrkL,69,70 it is plausible to hypothesize that Cas-L is part of a multiprotein complex comprising Cas-Crk-C3G-Rap1, thus leading to activation of LFA-1-dependent adhesion through effectors including Rapl and RIAM.63,64–67 In addition, previous studies show that the proline-rich tyrosine kinase 2 associates with LFA-1 and it is essential for CD8γ T-cell adhesion and migration. Proline-rich tyrosine kinase 2 has been reported to phosphorylate Cas-L,71 and unpublished data from Sheetz lab suggest that Cas-L might form a complex with Proline-rich tyrosine kinase 2 and LFA-1 that could work as a physical link connecting TCR clusters, LFA-1, and the actin cytoskeleton. Moreover, Cas-L might also be involved in Rap1 activation, integrin-mediated adhesion and f-actin polymerization by associating with the adaptor protein Chat-H, thus forming a complex that has previously been implicated in integrin-mediated adhesion and naive T-cell migration in response to TCR stimulation.69,70 In those studies, RNA interference-mediated knockdown of Chat-H resulted in impaired Rap1 activation, integrin-mediated adhesion and T-cell migration.69 Those results are in line with our model and support a role for the Cas-L/Chat-H complex as an early regulator of T-cell adhesion and migration upstream of Rap1 in response to TCR stimulation.

It is known that upon TCR stimulation, ZAP70, a kinase important for TCR-mediated signaling leading to Ca^2+ flux and IL-2 production, is recruited to the phosphorylated immunoreceptor tyrosine-based activation motifs on the zeta chains of the CD3ε subunit of TCR. Then, ZAP70 phosphorylates adaptor proteins LAT and SLP76, which in turn recruit PLCγ1.72,80,82 Subsequently, PLCγ1 cleaves PIP2, generating IP3 and DAG, which lead to Ca^2+ release, and PKCθ activation and Ras activation, respectively.92 PLCγ-1 activation is a critical step in TCR signaling that requires the function of the TCR proximal tyrosine kinase cascade and links it to rapid release of Ca^2+ from intracellular stores. Here, we saw that Cas-L^{−/−} CD8γ T cells have a significant defect in Ca^2+ release in response to TCR stimulation. This defect was accompanied by decreased levels of phospho-PLCγ-1 at the immunological synapses of Cas-L^{−/−} T cells relative to the control, supporting a role for Cas-L in the amplification of TCR signals that ultimately lead to T-cell activation.

In order to see whether the absence of Cas-L had an effect on signaling steps further downstream of TCR stimulation, we compared the levels of IL-2 production in Cas-L^{−/−} and wild-type T cells and saw no significant difference in IL-2 production. Consistent with this observation, we showed that pharmacological inhibition of ZAP70 had a minor effect on phospho-Cas-L levels. In line with these results, previous studies have also shown that disruption of f-actin polymerization reduced levels of phospho-PLCγ-1 within TCR microclusters without affecting ZAP70 activity. These observations suggest that alternative pathways downstream of ZAP70 take place independently of Cas-L activation.

In addition, the molecular events leading to Cas-L activation take place at a different stage and time scale than the events leading to IL-2 expression and secretion. In fact, whereas Cas-L recruitment and phosphorylation at the TCR microclusters is transient, occurring in the first 2–5 min of synapse formation and then dropping, IL-2 production takes place at later stages, 24–48 h after synapse formation. Thus, Cas-L seems to have a particularly important role at early stages, whereas WASp has a broader role such that IL-2 production is lower in WASp^{−/−} T cells,71 perhaps owing to structurally distinct TCR-mediated signaling events that are not as dependent upon Cas-L. Nevertheless, we provide clear evidence that Cas-L^{−/−} T cells show impaired early Ca^2+ signaling compared with wild-type cells, demonstrating a role for Cas-L in TCR-mediated signaling and in regulating immunological synapse stability.

Recent studies in WASp^{−/−} T cells reveal that TCR microcluster-associated f-actin foci have an important role of PLCγ-1 activation without having any effect on the TCR proximal tyrosine kinase cascade.95 Whereas WASP-dependent f-actin foci were closely associated with TCR signaling, phosphorylation of Cas-L was only weakly impacted by inhibition of ZAP70 catalytic activity, which is also a characteristic of LFA-1 regulation downstream of the TCR.95 Altogether these data suggest that WASp-dependent activation of Cas-L by TCR microcluster-associated f-actin likely has an important role in the initiation and maintenance of TCR signaling.

The guanine nucleotide exchange factor DOCK2 is found to be associated with CrkL in leukemia cell lines,95 which in turn binds phospho-Cas-L via its SH2 domain downstream of TCR activation.50,52 Thus, we speculate that a Cas-L-CrkL-DOCK2-Rac multiprotein complex can that directly activate the small GTPase Rac downstream of TCR activation, thus promoting f-actin polymerization through activation of the WAVE2 complex.96–98 Furthermore, a recent study reported that early TCR microclusters are able to activate T-cell cytokine expression, centralization of TCR microclusters is more important for Notch-ADAM10-Vav1-dependent T-cell proliferation. It is possible that the role of Cas-L in the centripetal movement of TCR microclusters at the immunological synapse might be more relevant for the signaling pathway leading to T-cell proliferation than to cytokine production. Hence, Cas-L may have distinct roles in different signaling pathways leading to cytokine production and T-cell proliferation, thus providing alternative regulation mechanisms for the TCR-signaling events that ultimately lead to IL-2 production and T-cell-mediated immune responses.

Accumulation of ICAM-1 at the immunological synapse is rapidly initiated and then sustained for many minutes even when TCR signaling is selectively blocked.99 However, we saw that accumulation of ICAM-1 at the synapses of Cas-L^{−/−} T cells was impaired, suggesting a potential role for Cas-L-mediated integrin activation. Surprisingly, using bilayers with increasing amounts of ICAM-1 (and a fixed amount of anti-CD3ε antibody) did not rescue spreading defects in Cas-L^{−/−} T cells. These were particularly interesting results as in control cells as few as 30 molecules of ICAM-1 per μm^2 were sufficient to allow synapse formation, whereas increasing to 650 molecules of ICAM-1 per μm^2 was insufficient to rescue this spreading defect in Cas-L^{−/−} T cells. The ability of Cas-L-deficient T cells to form a few
synapses, rather than none, may be a result of TCR and Cas-L-independent activation of integrins.\(^{100,101}\)

Our observations here are in line with recent studies we\(^{11,13}\) and others\(^{15}\) reported on the role of myosin IIa activity in immunological synapse formation and Cas-L activation. In those studies, inhibition of myosin IIa activity or myosin IIa knockdown activity led to a decrease, but not complete elimination, in phospho-Cas-L levels. Thus, those results supported a two-step model in which initial TCR ligation, activation of Lck, assembly of TCR microclusters, initial activation of Cas-L and recruitment of myosin IIa are all independent of myosin IIa activity, whereas amplification of Cas-L phosphorylation and microcluster transport are dependent on myosin IIa activity.

We can now develop a working model for the role for Cas-L as a link between TCR, actin and integrins (Figure 6). We previously described a model for actin polymerization-dependent stretch of p130Cas leading to TCR signal amplification (8–9) thus stabilizing the immunological synapse.

Figure 6 Working model illustrating the critical steps that lead to cytoskeletal stretch-dependent Cas-L activation and TCR signaling. Based on our observations, we propose a model where the initial steps of TCR ligation and clustering (1–2), assembly of actin nucleating factors (3) and initial f-actin polymerization at TCR microclusters (4) take place independently of Cas-L, and the subsequent steps leading to integrin activation and stabilization of lamellipodia structures (6), and TCR transport (7), are regulated by actin-dependent Cas-L activation at TCR microclusters. In turn, this mechanical activation of Cas-L acts via a potential feedback loop that leads to TCR signal amplification (8–9) thus stabilizing the immunological synapse.

Despite the striking morphological and functional differences between fibroblasts and T cells, here we demonstrate through the Cas model that the fundamental principles of mechanotransduction are evolutionarily conserved across different cell types.\(^{34,102}\)

METHODS

Cells and culture conditions

Cas-L\(^{-/-}\) mice on the C57BL6 background\(^{49}\) were maintained in a specific pathogen-free facility at Mount Sinai School of Medicine under the supervision of the institutional animal care and use committee. Naive CD8\(^+\) T cells were isolated from the spleens of Cas-L\(^{-/-}\) mice and their wild-type littermates using the Dynal negative isolation (Invitrogen, Waltham, MA, USA) following the manufacturer’s protocol. In brief, splenocytes were incubated with a mixture of antibodies against the CD8-negative T cells, and then incubated with super-paramagnetic polystyrene beads coated with a polyclonal sheep anti-rat immunoglobulin G antibody. Then, the bead-bound cells were separated by a magnet, and the remaining untouched CD8\(^+\) T cells were used for the experiments described here.

All experiments were performed in primary murine lymphocytes from the spleen, freshly isolated from mice about the same age (6–10 weeks), and enriched for CD8\(^+\) T cells using a standard negative selection procedure, yielding a population purity of 95–97%, as measured by flow cytometry (data not shown). Quiescent T-cell populations comprised a mixture of naive, memory and effector T cells, as measured by the levels of CD62L\(^{high}\) and CD127\(^^{+}\) (data not shown).

Cell culture medium was RPMI 1640 (Sigma, St Louis, MO, USA or GIBCO, Carlsbad, CA, USA) with 10% heat-inactivated fetal calf serum (Life Technologies, Carlsbad, CA, USA). Cells were kept in a 37°C, 5% CO\(_2\) humidified incubator.

Glass-supported lipid bilayer assembly

To recreate early events at the immunological synapse between a T-cell with an antigen-presenting cell in real-time we prepared planar lipid bilayers embedded with fluorescently labeled ligands to engage key T-cell surface receptors: 2C11 monoclonal antibodies against TCR/CD3\(\varepsilon\) and ICAM-1 at 35 molecules per \(\mu\)m\(^2\) and 150 molecules per \(\mu\)m\(^2\), respectively. Planar lipid bilayers provide a valid model for studying immunological synapse formation dynamics by high-resolution microscopy, faithfully reconstituting the supramolecular organization of the T-cell/APC interface previously described.\(^{5,58}\) The bilayer-presented TCR and LFA-1 ligands can move freely throughout the
plane of the bilayers, and we labeled them with different fluorophores to allow high-resolution spatio-temporal characterization of immunological synapse formation. To validate the quality of the bilayers, we tested the lateral mobility of the ligands by recording their fluorescence recovery after photobleaching kinetics before seeding cells. In all experiments, we use fresh primary CD8+ T cells purified by negative selection from spleen of Cas-L−/− mice or their wild-type littermates.

Supported planar bilayers were assembled in parallel plate flow cells (Biotecs, Butler, PA, USA or Ibidi, Madison, WI, USA) from unilamellar vesicles containing 12% mol% 1,2-dioleoyl-sn-glycero-3-[(N-(5-amino-1-carboxypentyl)-iminodiacetic acid)-succinyl] (nickel salt), 0.01 mol% 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine-N-(capriobutyl), and 87.99% mol% 1-oleoyl-2-palmitoyl-phosphatidylycholine (Avanti Polar Lipids, Alabaster, AL, USA). Bilayers were loaded with monobiotinylated 564-2C11 mouse antibody pre-polymerized with 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine-N-cyanoethylmaleimide-dodecanoylphosphatidylethanolamine (Avanti Polar Lipids, Alabaster, AL, USA). Non-specific binding was reduced by blocking with 5% casein in phosphate-buffered saline. Cells were allowed to settle and form contacts with the bilayer during imaging.

The estimate of the number of molecules of ICAM-1 on the bilayers was performed on an independent assay using bilayers on glass beads as described elsewhere.

**Microscopy**

TIRF imaging was carried out using a Nikon Eclipse Ti with ×60 NA 1.49 objective and an Andor DU987 back illuminated electron multiplying charge coupled device camera. Solid-state lasers (Coherent, Santa Clara, CA, USA) provided illumination at 488, 561 and 641 nm and narrow pass filters (Chroma Technology, Bellows Falls, VT, USA) were used for detection. Widefield epifluorescence imaging was performed with a 150 W Xe lamp on an Olympus IX70 with a Zeiss ×40 1.3 NA Fluar objective, Ludl filters (Chroma Technology) and a CoolSNAP HQ camera (Roper Scientific, Acton, MA, USA). Illumination for Ca2+ imaging was provided by a 150 W Xe lamp on an Olympus IX70 with a Zeiss ×40 1.3 NA Fluar objective, Ludl filters (Chroma Technology) and a CoolSNAP HQ camera (Roper Scientific, Acton, MA, USA). Second order polynomials were used to fit the measured data for Ca2+ imaging.

**Image analysis**

The cell boundaries were defined in ImageJ using the ‘Magic wand’ tool after making the images binary with the ‘Threshold’ function. The ‘Magic wand’ tool uses an algorithm that selects all connected pixels above the lower and below the upper threshold values (or value 1 in the case of a binary image), which were set based on the image background and pixel saturation, respectively. To obtain a coherent mask of the cells, the functions ‘Close’, ‘Fill Holes’ and ‘Remove Outliers’ were also used when necessary. Each cell was added to a list using ‘ROI Manager’, and then measurements of area and mean intensity were compiled. Our method to define cell boundaries involved using the program CellProfiler,104 setting a threshold intensity mask in one of the channels to identify the whole synapse area, and setting another mask on the other channel to identify TCR microclusters. Thus, we were able to measure total or mean fluorescence intensity of proteins over the whole synapse area, or just over the TCR clusters only. Using the masks defined in CellProfiler we computed the Pearson’s correlation coefficient to quantify the overlap between phospho-Cas-L and TCR over the whole synapse area or just over the TCR clusters only. The Pearson’s coefficient was then plotted to compare signal colocalization.

**Intracellular Ca2+ detection**

We tracked spreading cells by internal reflection microscopy and used a Ca2+-specific fluorescent probe (Fluo-4-AM, Molecular Probes, Eugene, OR, USA, F14217) to detect variations of Ca2+ levels with epifluorescence. The Fluor-4-AM Ca2+ dye was dissolved in DMSO at 5 mM (×1000) and Phoronic (Molecular Probes P3000MP) was added at 50 mg mL−1 to aid the solubilization of the dye into aqueous buffers. Cells were pelleted by centrifuging at 200 g (1250 r.p.m.) for 5 min at 4 °C, and the pellet was resuspended in 1 ml Hepes buffered saline supplemented with 5 mM glucose, 2 mM MgCl2, 1 mM CaCl2, and 1% human serum albumin (HBS/HSA). The freshly made Fluo-4/DMSO/Phoronic solution was added to the cell suspension to achieve a 5 μM final Fluo-4-AM concentration, and incubated 30 min at room temperature, to allow efficient uptake and de-esterification of the AM esters from the dye. Finally, cells were washed with phosphate-buffered saline, pelleted and resuspended in phenol free HBS/1%HSA. As a measurement of the relative changes in free intracellular Ca2+, raw values of total fluorescence intensity for each cell were background-corrected and normalized to the maximum intracellular Ca2+ release measured after addition to the medium of ionomycin (1 μM), an ionophore that releases Ca2+ from intracellular stores. The acquisition rate was 16 frames min−1 for the long term acquisition.

**IL-2 production measurement**

Freshly isolated Cas-L−/− or control mouse splenic CD8+ T cells (1×106 cells) were cultured in 200 μl Rosewell Park Memorial Institute medium with 10% fetal calf serum for 48 h on glass chambers coated with anti-CD3ε antibody and ICAM-1 (both at 5 μg ml−1), or uncoated chambers as unstimulated (negative) control. Cell culture supernatant was collected at 6 h, 12 h, 24 h or 48 h of culture, centrifuged to remove cells and debris and stored at −20 °C until analysis. IL-2 concentration in the supernatant was determined using a standard ELISA kit for mouse IL-2 (Invitrogen). A plate reader was used to measure 450 nm absorbance of each well, and a standard curve was plotted with a second order polynomial fit.

**Pharmacological perturbations**

The Src-family kinase inhibitor PP275 was added to 1×106 cells in 100 μl at a final concentration of 10 μM, incubated for 15 min at 37 °C, and added to the bilayers. Treatments with Lck inhibitor (10 μM), Lck- and Syk-family inhibitor Picatannol (10 μM), PKCθ inhibitor C20 (10 μM or 33 μM), or PKCδ inhibitor C20 (10 μM), and actin polymerization inhibitor cytochalasin D (60 μM or 200 μM) followed similar protocol described for PP2. All inhibitors were dissolved in DMSO, and all pharmacological experiments performed included a DMSO only incubation control.

**Immunocytochemistry**

After incubating cells for 3 min at 37 °C, cells were fixed with 2% paraformaldehyde, and permeabilized with 0.01% Triton-X in phosphate-buffered saline for 2 min, followed by blocking with 5% casein for 30 min. Primary antibody incubation was performed overnight at 4 °C, followed by fluorescently labeled secondary antibody incubation (with or without Phalloidin) for 1 h at room temperature. Primary antibodies used: Cas-L (clone 2G9; Novus Biologicals, Littleton, CO, USA, #NB100-1699); phospho-Cas-L (Cell Signaling, Danvers, MA, USA, #4015); ZAP70 (clone 99F2; Cell Signaling #2705); phospho-ZAP70 (Tyr319; Cell Signaling #2701). Fluorescently labeled secondary antibodies were obtained from Molecular Probes.

**Sodium dodecylsulfate polyacrylamide gel electrophoresis and immunoblotting**

Cells (1×106 in 100 μl of RPMI 1640 complete medium) were incubated with pharmacological agents as indicated for 15 min prior to seeding on anti-CD3ε/anti-CD28/ICAM-1–coated glass coverslip. After 5 min of interaction with the coverslip cells were lysed with radioimmunoprecipitation assay buffer, solubilized in Laemmli sample buffer supplemented with 5% beta-mercaptoethanol, boiled and stored at −70 °C. Samples were run on a gradient (4–20%) sodium dodecylsulfate polyacrylamide gel electrophoresis and transferred to a nitrocellulose membrane, blocked, incubated overnight with primary antibody, washed, incubated for 1 h with horseradish peroxidase-labeled secondary antibodies and immunoreactivity was visualized by chemiluminescence detection with film. Relative quantification of protein levels was performed from intensity measurements of the bands on the membrane images using ImageJ ‘Measurements/Intensity’ tool after inversion of the lookup table to generate positive values.

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Cas-L coordinates T-cell actin cytoskeleton

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

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