



Published in final edited form as:

Immunity. 2010 April 23; 32(4): 531–540. doi:10.1016/j.immuni.2010.04.005.

Essential role of Ubiquitin and TSG101 in formation and function of the central supramolecular activation cluster

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Abstract

Agonist MHC-peptide complexes in the immunological synapse (IS) signal through T cell receptor (TCR) microclusters (MC) that converge into a central supramolecular activation cluster (cSMAC). The determinants and function of the cSMAC remain unknown. We demonstrate an essential role for ubiquitin and TSG101, but less so for HRS, in signal processing events at the cSMAC. Using siRNA in primary T cells, we show that Ub recognition by TSG101 is required for cSMAC formation, TCR MC signal termination, TCR down-regulation, and segregation of TCR-MHC-peptide from PKC- θ -enriched signaling complexes. Weak agonist MHC-peptide induced CD80-dependent TCR MCs that dissociated in the center of the IS without recruiting TSG101. These results support TSG101-dependent recognition of CD80-independent TCR MCs as a molecular checkpoint for TCR down-regulation.

Introduction

Complex signal integration in cells involves responses to diverse stimuli that often utilize overlapping signal transduction machinery. Appropriate responses to such stimuli may benefit from spatial segregation of ligand-receptor interactions. In T cells, immunological synapses (IS) are specialized cell-cell junctions that combine cell polarization and positional stability with spatial segregation of interacting elements (Dustin, 2002; Dustin et al., 1998; Monks et al., 1998). IS formation is critical for signal integration, as well as coordination of migration, directed secretion, asymmetric cell division, and differentiation (Davis et al., 2007; Dustin, 2008; Huppa and Davis, 2003; Krogsgaard et al., 2003). In CD4⁺ T cells, the IS is composed of three primary subdomains: a central supramolecular activation cluster (cSMAC) rich in proximal signaling components such as TCR-MHC-peptide and PKC- θ , a peripheral supramolecular activation cluster (pSMAC) dominated by ICAM-1-LFA-1 interactions, and a distal supramolecular activation cluster (dSMAC) rich in dynamic actin (Grakoui et al., 1999; Monks et al., 1998). The cSMAC is generated and maintained by microclusters (MCs) of 10-20 TCR that continuously form in an actin-dependent manner in the dSMAC, but become actin-independent as they translocate to the IS center and fuse to form the cSMAC (Krummel et al., 2000; Varma et al., 2006; Yokosuka et al., 2005). TCR MCs are the sites of signal initiation based on recruitment and phosphorylation of Lck, ZAP-70, and LAT as well as SLP-76 and Grb-2 recruitment (Bunnell et al., 2002; Campi et al., 2005; Huse et al., 2007; Yokosuka et al., 2005). In contrast, the cSMAC has 20-fold lower tyrosine phosphorylation than MCs and cannot independently sustain Ca²⁺ signaling

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(Campi et al., 2005; Varma et al., 2006). The molecular basis of signaling differences between MCs and the cSMAC is unknown.

Ligand-mediated TCR downregulation occurs via routing of internalized receptors to lysosomes (Valitutti et al., 1997), and enrichment of the cSMAC in multivesicular body (MVB) markers suggests a role for receptor degradation at the cSMAC (Varma et al., 2006). However, not all ligands induce cSMAC formation and it has recently been proposed that avoidance of cSMAC formation and TCR downregulation may underlie the elevated stimulatory potency of certain weak agonist ligands (Cemerski et al., 2007). TCR degradation may occur via ubiquitin recognition based on involvement of ubiquitin ligases such as Cbl-b (Naramura et al., 2002). Degradation of ubiquitinated substrates via MVBs is a stepwise process coordinated by multiple family members of the endosomal sorting complex required for transport (ESCRT) (Williams and Urbe, 2007). There are 4 ESCRT complexes (0, I, II, and III) with unique roles in signal termination and receptor degradation of epidermal growth factor (EGF) receptor. ESCRT 0 and I directly recognize Ub (Williams and Urbe, 2007). ESCRT-0 can also associate with ubiquitinated cargo through the Ub interaction motif (UIM) of Hrs (Bache et al., 2003b; Hirano et al., 2006) in parallel with inositol-3-phosphate bearing lipids present in endosomes (Raiborg et al., 2001). ESCRT-I recognizes Ub through the ubiquitin E3 variant (UEV) domain of TSG101 and is required for sorting of EGFR into MVBs (Pornillos et al., 2002; Sundquist et al., 2004; Teo et al., 2004). In the absence of TSG101, both MVB formation and sorting of proteins into MVBs is inhibited, resulting instead in persistent EGFR signaling (Bache et al., 2006; Doyotte et al., 2005). The specific role of ESCRT complexes in TCR down-regulation has not been determined.

Unresolved roles for the cSMAC in promoting proximal signaling still exist, particularly with respect to PKC θ , whose enrichment at the cSMAC has been described (Monks et al., 1998; Monks et al., 1997). Activated PKC- θ leads to downstream activation of NF- κ B and AP-1 (Isakov and Altman, 2002; Sun et al., 2000). CD28 is dispensable for PKC- θ recruitment to the IS, but is critical for its concentration in the cSMAC (Huang et al., 2002). Recent reports have suggested discrete sites of CD28/PKC θ and TCR enrichment within the IS (Sims et al., 2007; Tseng et al., 2008; Yokosuka et al., 2008), but the molecular basis for this segregation is as yet unclear. Understanding mechanisms of segregation may provide insight into how cells simultaneously process multiple signals.

To address the role of Ub and ESCRT mediated Ub recognition in cSMAC formation and function we have applied siRNA-mediated suppression and high-resolution fluorescence microscopy of IS formed between primary T cells and supported bilayers. We have determined that Ub is required for cSMAC formation in the IS in response to agonist MHC-peptide complexes. Ub recognition by ESCRT-I component TSG101, but not ESCRT-0 component HRS, is required for cSMAC formation, TCR MC signal termination and segregation of TCR-agonist MHCp complexes from PKC- θ -enriched domains within an outer cSMAC subdomain. We also demonstrate that TSG101 is rapidly recruited to the plasma membrane following T cell activation by fluorescence resonance energy transfer. Finally, we describe signaling MCs formed in response to weak agonist ligands that remain actin-dependent and dissipate at the cSMAC/pSMAC boundary without recruiting ESCRT-I complexes. The IS thus engages in a conditional utilization of ESCRT-I and spatially defined checkpoints to regulate T cell signaling and down-regulation.

Results

Ubiquitination is required for cSMAC formation

Prior observations that MVBs are enriched at the cSMAC led us to investigate whether ubiquitination is involved in IS patterning (Varma et al., 2006). Treatment of cells with MG132 inhibits the proteasome resulting in accumulation of poly-ubiquitinated substrates and depletion of the free ubiquitin pool (Melikova et al., 2006). We therefore incubated MG132-treated AND TCR Tg T cells on supported planar bilayers presenting I-E^k-MCC + ICAM-1. Control treated T cells formed TCR MCs, which accumulated in the cSMAC in response to bilayers presenting agonist MHC-peptide (Fig. 1A, Fig. S1A and Movie S1). In contrast, we found that MG132 treated T cells were able to efficiently form TCR MCs, but were unable to translocate MCs from the site of formation to the IS center. The amount of TCR accumulating at the interface was increased significantly in the MG132-treated cells (Fig. 1B). Furthermore, MG132 treatment significantly impeded termination of TCR signals, as phosphorylation of peripherally accumulated TCR MCs was 5-10 fold higher in MG132 treated T cell IS as compared to control T cell IS (Fig. 1A, B). While MG132 treatment may have a number of additional effects, these results suggest that ubiquitination is required for cSMAC formation and encouraged us to further pursue this line of experimentation.

TSG101 is required for cSMAC formation and MC dephosphorylation

The suggested role for ubiquitination in cSMAC formation led us to investigate the role of ubiquitin recognizing ESCRT complex components in regulating TCR signaling within the IS. We chose to target HRS and TSG101, the respective critical components of the ESCRT-0 and ESCRT-I complexes with established ubiquitin binding (Fig. 1C) (Williams and Urbe, 2007). We developed a procedure to efficiently suppress expression of HRS or TSG101 in 100% of activated AND TCR transgenic (Tg) T cells, by nucleofection of either specific siRNA duplexes or scrambled versions of the same sequence as controls. We were able to reduce HRS and TSG101 expression more than 10-fold during primary expansion of AND TCR Tg T cells (Fig. 1D).

In EGFR systems, ESCRT-0 is initially recruited to ubiquitinated cargo within clathrin-coated pits and is required for efficient receptor degradation (Bache et al., 2003a; Bilodeau et al., 2003; Katzmann et al., 2003). However, HRS knockdown (KD) T cells formed a cSMAC containing dephosphorylated TCR (Fig. 1E-F). In addition, T cells transfected with GFP-tagged clathrin light chain showed no colocalization of clathrin-coated pits with TCR MCs in the IS (Fig. S1B and Movie S2). However, clathrin-coated pits were present in distinct puncta emerging in the IS periphery, and accumulated in the IS center (Fig. S1B and Movie S2). Furthermore, HRS KD resulted in accumulation of phosphotyrosine in the IS periphery that did not co-localize with TCR (Fig. 1E-F). This leaves open the possibility that clathrin/HRS might be involved in regulation of downstream signaling complexes that are known to dissociate from TCR MCs and signal independently (Bunnell et al., 2002), however, because this phenotype is distinct from that of MG132 treatment we did not pursue this further here.

In contrast to HRS, TSG101 KD generated a phenotype more similar to that of MG132 treatment. Control siRNA treated cells formed TCR MCs, which accumulated in the cSMAC in response to bilayers presenting agonist MHCp. In contrast, TCR MCs failed to translocate to the cSMAC and instead accumulated more peripherally in TSG101 KD T cells (Fig. 1G, S2A,B and Movie S3). TSG101 KD resulted in accumulation of phosphorylated TCR MCs in quantities and distribution that were similar to MG132 treated T cell IS (Fig. 1G-H, S2A, B), suggesting that TSG101 translocates ubiquitinated TCR MCs to the cSMAC for dephosphorylation and degradation. The elevated anti-phosphotyrosine intensity in

TSG101 KD ISs did not correlate directly with the TCR intensity in MCs, but all visible TCR MCs were intensely phosphotyrosine positive, except in rare cases where TCR MCs penetrated into the IS center. A proportion of the elevated phosphotyrosine signal in TSG101 KD ISs did not colocalize with TCR. This may be due to phosphorylation of small clusters of TCR that are below the limit of detection. Alternatively, they may represent other surface proteins, such as LAT and associated proteins that are known to cluster independently following TCR signaling (Balagopalan et al., 2007). The accumulation of large TCR MCs in the periphery of TSG101 KD continued to engage agonist MHCp (Fig. S2C) and resulted in slightly reduced LFA-1-ICAM-1 accumulation in the pSMAC, presumably due to competition with the TCR MCs for space (Fig S2D). The similarity in TCR location and co-localized phosphorylation between the MG132 treatment and TSG101 KD led us to further characterize TSG101's role in IS formation and signal processing.

In order to establish whether TSG101 was positioned to play direct roles in IS patterning, we investigated the distribution of TSG101 within T cells forming IS. We found that both endogenous and recombinant fluorescent TSG101 were strongly recruited to the cSMAC in response to agonist pMHC engagement on supported planar bilayers (Fig. S2E).

TSG101 is required for cSMAC formation and function in T-B conjugates

It remained possible, although unlikely, that our findings using supported planar bilayers did not faithfully recapitulate IS formation and hence TSG101 function in cell-cell interfaces. To rule out this possibility, we imaged control and TSG101 KD T cells forming ISs with primary B by confocal microscopy. Reconstructed T-B cell interfaces displayed comparable peripheral TCR accumulation and elevated phosphotyrosine following TSG101 KD to those observed on bilayers (Fig. 2A-C). Consistent with our imaging results, flow cytometry analysis of intact or permeabilized T-B conjugates demonstrated a significant defect in antigen-induced TCR surface downregulation and degradation, respectively, in the absence of TSG101 (Fig. 2D).

We then investigated whether the recruitment of TSG101 to the IS seen in T cells incubated on supported planar bilayers was recapitulated in a cell-cell system. In particular, we wondered whether T cell activation resulted in recruitment of TSG101 to membrane compartments, as would be required in order to execute sorting processes within the IS. We therefore evaluated activation-induced recruitment of TSG101 to T cell membranes by fluorescence resonance energy transfer (FRET) (Wouters et al., 1998). Efficient FRET between membranes and associated signaling proteins has been reported by donor dequenching after acceptor photo-bleaching, using GFP tagged proteins as donor and the membrane dye DiI as acceptor (McLean et al., 2000). In order to use this method to probe membrane interactions of TSG101, we stably expressed a murine TSG101-GFP chimera in AND T cells by retroviral transduction, and sorted them for uniform expression. We then briefly labeled transduced T cells with DiI before allowing them to form conjugates with MCC-loaded murine B cells. Changes in donor fluorescence following photobleaching of DiI revealed a low and diffuse FRET signal throughout the T cell, indicating loose coupling of TSG101 with plasma membrane and endomembranes in the steady-state (Fig. 2E-F) as previously reported (Welsch et al., 2006). Upon IS formation with antigen-pulsed B cells, TSG101 underwent a marked redistribution, polarizing strongly to the IS, but also outside the IS (Fig. 2E-F). The high apparent FRET efficiencies observed is consistent with direct apposition of membranes and tagged TSG101. Notably, the FRET signal was most prominent at the cSMAC in the T-B cell contacts (Fig. 2E). Therefore, T cell activation results in redistribution of TSG101 to membrane compartments concentrated in the IS.

Sorting of TCR MCs into the IS center depends on Ub recognition by TSG101

We then sought to confirm the specificity of TSG101 in inducing cSMAC formation and signal termination. We observed similar defects in cSMAC formation and dephosphorylation when TSG101 was suppressed with 3 different siRNA oligomers (Fig. S3). Restoration of wild-type IS patterning by reintroduction of wild-type human TSG101 following KD of endogenous mouse TSG101 indicated that our observed defects in TCR dephosphorylation and cSMAC formation were specifically attributable to TSG101 function (Fig. S3). However, we found that a mutant TSG101 with substitutions in the UEV domain that abrogate Ub binding (Pornillos et al., 2002) was completely unable to restore cSMAC formation or dephosphorylation following TSG101 KD (Fig. 3). Cell lysates from TSG101 KD T cells demonstrated a marked accumulation of ubiquitinated species as compared to control siRNA treated T cells (Fig. S4A), and ubiquitination was further increased by agonist MHCp stimulation. These higher molecular weight species were also identifiable by antibodies to CD3 ζ (Fig. S4B). This suggests that CD3 ζ may be a ubiquitinated protein recognized by TSG101. In combination with the equivalent defects observed in TSG101 KD and MG132 treated T cells forming IS, these data all support the model that TSG101 specifically interacts with ubiquitinated components of TCR MCs to induce cSMAC formation and TCR signal termination.

Differential sorting of receptors by TSG101

Numerous receptors are engaged within the IS that are regulated by diverse mechanisms. A recent example of sorting of such receptors in the IS is the observation that CD28 and TCR traffic together in peripheral MCs upon engagement and are then segregated from each other in the cSMAC (Yokosuka et al., 2008). In light of our finding that Ub-mediated TSG101 recognition is necessary for its incorporation of TCR into a cSMAC, we wondered whether TSG101 might also be involved in segregating TCR from CD28. We evaluated these sorting events by observing control and TSG101 KD T cells interacting with bilayers containing MHCp, ICAM-1, and CD80 and following PKC- θ enriched compartments. Consistent with prior reports, we found that in control cells, PKC θ and TCR colocalized in peripheral MCs but became segregated in the cSMAC (Fig 4A and S5A)(Yokosuka et al., 2008). In TSG101 KD T cells, however, PKC- θ and TCR failed to segregate and remained colocalized in large MCs in the pSMAC region, with a significantly elevated Pearson's correlation coefficient (Fig. 4A, Fig S4A). Therefore, TSG101 functions at the cSMAC-pSMAC interface to generate distinct TCR and PKC- θ enriched domains of the cSMAC.

We then investigated the mechanism by which TSG101 selectively terminates TCR MC signals at the IS center. Normally, TCR MCs become relatively immobile as they accumulate at the IS center (Varma et al., 2006). The association of immobile central TCR with LBPA staining is consistent with incorporation of TCR into MVBs (Varma et al., 2006). In control T cells, TCR MCs became immobilized as they became co-localized with LBPA⁺ structures at the cSMAC (Fig 4B-C). In contrast, we observed continued jostling movements of large TCR MCs in the periphery of TSG101 KD IS (Fig. 4B and Movie S3). This jostling movement is represented in Fig. 4B by overlaying 3 sequential images at 2 minute intervals with red, green, and blue coding such that immobile structures appear white and mobile structures appear colored. The continued jostling of TCR in TSG101 KD IS correlated with a striking segregation of TCR from LBPA⁺ structures (Fig. 4C). This segregation was confirmed by a significant decrease in the Pearson's correlation coefficient between TCR and LBPA in TSG101 KD IS (Fig. S5B). Thus, TSG101 is required for TCR co-localization with LBPA⁺ compartments.

TSG101 suppression leads to chronic TCR signaling

The striking persistence of phosphorylated TCR MCs following TSG101 KD (Fig. 1G-H) suggested to us that the duration of signaling through engaged TCR might be extended in TSG101 KD T cells. To test this, we allowed control and TSG101 KD T cells to be activated with agonist MHCp ligand on bilayers for 10 minutes, sufficient to induce stable TCR engagements, prior to treatment with blocking anti-I-E^k-MCC antibodies (D4) to prevent new TCR MC formation. Signaling was monitored using intracellular Ca²⁺. In control siRNA treated T cells, the Ca²⁺ signal decayed over 2 minutes (Fig. 5A,B, Fig. S6), as previously described (Varma et al., 2006). In contrast, Ca²⁺ elevation decayed only partially and was then sustained at an intermediate level in TSG101 KD T cells after D4 treatment (Fig. 5A,B, Fig. S6). The net effect of increased TCR signaling in TSG101 KD T cells was increased IL-2 production compared to control siRNA treated T cells (Fig. 5C). It was not possible to examine proliferation of T cells in the TSG101 KD because TSG101 is required for the abscission step of cellular cytokinesis (Morita et al., 2007); as such, we noted that TSG101 KD consistently inhibited T cell proliferation. Thus, TSG101 plays an important role in limiting downstream T cell signals and effector responses following productive proximal signaling in response to TCR engagement.

Weak agonist MHCp do not involve TSG101

Typical recognition of agonist pMHC results in both tyrosine phosphorylation-mediated T cell activation and Ub-mediated TCR down-regulation (Naramura et al., 2002; Samelson et al., 1986). However, T cell activation in the absence of TCR down-regulation has recently been reported. The altered peptide ligand (APL) MCC-K99A, a variant of MCC88-103 with a single amino acid substitution that greatly accelerates I-E^k-based tetramer dissociation compared to the native sequence, effectively stimulates proliferation of AND Tg T cells when presented by splenic APCs despite inducing diminished TCR downregulation and central TCR accumulation (Cemerski et al., 2007). We began investigating IS formation by AND T cells in response to I-E^k-MCC-K99A in a planar bilayer system to understand the properties of this ligand-receptor interaction. Supported planar bilayers presenting I-E^k-MCC-K99A and ICAM-1 arrested T cell migration and induced sustained Ca²⁺ elevation in AND T cells only in the presence of CD80 (Fig. S7A, B). This is consistent with reports in which CD28 engagement is required for response to weak agonist ligands (Bachmann et al., 1997). Real-time imaging of TCR and LFA-1/ICAM-1 interactions demonstrated that, in the presence of CD80, T cells responding to I-E^k-MCC-K99A formed IS with many structural similarities to I-E^k-MCC-induced IS. T cells spread symmetrically on K99A-presenting surfaces, and LFA-1-ICAM-1 interactions quickly organized into a pSMAC (Fig. S7C). Tyrosine phosphorylation and proximal TCR signaling (indicated by recruitment and phosphorylation of LAT) were concentrated in small MCs in the periphery of the IS with a similar intensity and frequency to IS formed in response to I-E^k-MCC (Fig. S7C, D). However, in striking contrast to I-E^k-MCC-induced IS, TCR did not accumulate in the IS center even in response to 10 molecules/μm² of I-E^k-MCC-K99A (Fig. 6A, B, p < 0.0001; Fig. S7C, D.). Time-lapse TIRFM imaging of IS formed in response to I-E^k-MCC-K99A revealed co-stimulation dependent formation and centripetal translocation of TCR MCs (Fig 6A and Movie S4); however, these MCs dissipated as they reached the center of the IS rather than accumulating in a cSMAC as they do in response to I-E^k-MCC.

In order to resolve this difference between I-E^k-MCC-K99A and I-E^k-MCC-induced IS, we investigated the contribution of TSG101 to ISs formed in response to these respective ligands. Notably, TSG101 was recruited to the cSMAC of I-E^k-MCC-induced IS, but not to I-E^k-MCC-K99A-induced IS (Fig. 6B). Furthermore, TSG101 KD had no impact on TCR MC formation, size, transport, or phosphotyrosine levels within MCC-K99A induced ISs (Fig. 6C and Fig S8A). These results demonstrate that TSG101 is not required for early TCR

signaling, TCR MC transport, pSMAC formation or a radially symmetric IS in response to the MCC-K99A weak agonist MHCp. It has previously been demonstrated that TCR MCs form in an F-actin dependent manner, but then become rapidly F-actin independent based on their stability in the presence of latrunculin A, which sequesters G-actin (Varma et al., 2006). Weak agonist MCC-K99A induced TCR MCs remained F-actin dependent, in contrast to many F-actin independent MCs that accumulate in response to I-E^k-MCC (Fig. S8B). The cSMAC is generally free or depleted of F-actin (Kaizuka et al., 2007). Analysis of the F-actin distribution in the IS formed in response to MCC and MCC-K99A MHCp confirmed that I-E^k-MCC-K99A does not induce an F-actin free central zone (Fig. 8C).

Discussion

We previously demonstrated that the cSMAC shows a striking enrichment of LBPA (Varma et al., 2006), a lipid associated with MVBs. This suggested that the cSMAC might be a site for degradation of TCR through an ESCRT dependent pathway (Williams and Urbe, 2007). Other results in the field were consistent with this notion including earlier observations that CD2AP, an adapter linking actin and Cbl family ubiquitin E3-ligases is required for normal TCR degradation and signal termination in the cSMAC (Lee et al., 2003). There was also ample evidence for a role of Cbl family E3 ligases in TCR signal termination, and a role for lysosomes in TCR degradation (Liu et al., 2000; Naramura et al., 2002; Valitutti et al., 1997). It had also been demonstrated that TCR down-regulation correlates with MHCp potency (Itoh et al., 1999) and that a subset of weak agonist ligands do not trigger TCR down regulation at all (Cemerski et al., 2007). Furthermore, the TCR ζ and δ chains are directly ubiquitinated upon strong TCR engagement (Cenciarelli et al., 1992). Based on these observations we anticipated that ubiquitination and ESCRT machinery might be important for TCR signal termination and eventual degradation after engagement with agonist MHCp. We were surprised by our finding in this study that depletion of free ubiquitin with MG132 or knockdown of TSG101 not only blocked TCR signal termination and degradation, but also blocked cSMAC formation in response to agonist MHCp. In contrast, a weak agonist MHCp could induce robust TCR MCs, Ca²⁺ elevation and pSMAC formation without recruiting TSG101 or inducing a cSMAC in normal T cells. These results have important implications for signal integration in response to agonist MHCp and how this differs from weak agonist MHCp.

Our results suggest distinct roles for major Ub recognizing ESCRT components HRS and TSG101 in IS formation and TCR down-regulation. HRS is part of ESCRT-0 and associates with ubiquitinated cargo through its ubiquitin interaction motif (UIM) (Bache et al., 2003b; Hirano et al., 2006), in parallel with inositol-3-phosphate bearing lipids present in endosomes (Raiborg et al., 2001). In EGF-R systems, HRS promotes ubiquitin recognition by TSG101 by directing TSG101 to early endosomes containing ubiquitinated cargo (Katzmann et al., 2003). This process is dependent on the recruitment of HRS recruitment to receptors within clathrin lattices (Raiborg et al., 2002), and clathrin clustering of ubiquitinated cargo is required for efficient HRS recognition (Raiborg et al., 2006). Indeed, in EGF-R systems involving both clathrin-dependent and clathrin-independent endocytic mechanisms, HRS is involved only in degradation of clathrin-endocytosed receptors (Myromslien et al., 2006). The specificity of HRS for clathrin-coated pits and the failure of TCR MCs to intersect with these compartments may explain the finding that a cSMAC is formed in the absence of HRS (Razi and Futter, 2006). Furthermore, in non-EGFR systems, the association of HRS and TSG101 is not so clear. Indeed, TSG101-mediated endosomal trafficking of Tyrp1 is independent of HRS function in melanocytes (Truschel et al., 2009). Furthermore, HRS is involved in alternate internal trafficking pathways, such as sequence-dependent recycling of the beta-2 adrenergic receptor (Hanyaloglu et al., 2005). Thus, prior models suggesting a strictly sequential transfer of ubiquitinated cargo from ESCRT-0 to

ESCRT-I may not apply in all situations, as has been recently demonstrated (Shields et al., 2009). Downregulation of engaged TCR complexes appears to be clathrin independent (Monjas et al., 2004), while internalization of the proximal signaling molecule LAT has been shown to be clathrin dependent (Brignatz et al., 2005). We detected a weak, but significant effect of HRS KD on non-TCR MC associated phosphotyrosine. This suggests that HRS may act on distinct substrates from the TCR during T cell activation and this is worthy of further study.

Our results provide insight into the distinct mechanisms of sustained signaling by agonist and weak agonist MHCp. TSG101 prevents chronic signaling by agonist MHCp induced TCR MCs. TCR-agonist MHCp and CD28-CD80 interactions are initially co-localized in MCs, but then are segregated within the cSMAC (Yokosuka et al., 2008). We show that this segregation requires TSG101. There are a number of possibilities for how TSG101 organizes this process. Two possibilities are polymerization of ESCRT III components to generate MVBs that are then translocated toward the center by microtubules (Wollert et al., 2009) or linkage to a final actin dependent translocation step through Alix before later ESCRT steps are initiated (Pan et al., 2006; Strack et al., 2003). This sorting process, however mediated, led to formation of the PKC- θ -rich compartment, which is a hallmark of the IS formed with professional APCs (Monks et al., 1997; Tseng et al., 2008; Yokosuka et al., 2008). Weak agonist MHCp required CD80 in the bilayer to form TCR MCs and sustain Ca^{2+} signals, which is not the case for agonist MHCp. This result is consistent with earlier studies showing that responses to weak agonist MHCp are CD28 dependent (Bachmann et al., 1997). Weak agonist MCC-K99A induced TCR MCs that remained F-actin dependent and disappeared in the center of the IS without a requirement for an F-actin free central zone or TSG101. The mechanism by which weak agonist MHCp signals are terminated is not clear, but we speculate that the requirement for a dynamic contribution from CD80 and the continued requirement for F-actin may enable physical disengagement and recycling of TCR at the point of convergence of centripetal F-actin flow (Kaizuka et al., 2007). More stable MCs formed by agonist MHCp might persist in response to the same physical process, necessitating the use of TSG101 to mark irreversibly clustered TCR-MHCp complex for elimination. The other distinguishing feature of signaling mediated by agonist and weak agonist MHCp is the generation of the consolidated PKC- θ -rich compartment by agonist MHCp. Since the cSMAC size is linearly related to agonist MHCp density (Varma et al., 2006) the formation of the PKC- θ -rich outer cSMAC may provide a mechanism to measure integrated TCR avidity in an IS as long as CD80 or CD86 are present on the APC.

These data lead us to a model in which ESCRT-I is required for sorting of TCR into MVB-like structures within the cSMAC and for signal termination in response to agonist MHCp (Fig 7). Our findings clarify the mechanism of cSMAC formation and the origin of recently described segregation of inner TCR and outer, PKC- θ enriched, cSMAC regions (Kaizuka et al., 2007; Varma et al., 2006; Yokosuka et al., 2008). Integrating TSG101 into our current picture of IS structure clarifies the inhibitory role of the cSMAC and the sorting function of TSG101 reconciles the co-existence of signaling and degradation compartments within this central structure in T cell activation (Cemerski et al., 2008; Lee et al., 2003; Varma et al., 2006).

Experimental Procedures

Please refer to supplementary information for complete methods.

Mice, Cells and Cell Culture

AND TCR Tg spleen cells were activated in OK-DMEM (Invitrogen) with 10% FBS (Hyclone) for 48 h using 2 μ M MCC peptide (MCC88-103). Cells were replenished with

fresh media and IL-2 every 24 h for 4 days following activation (Campi et al., 2005). B cells were obtained from B10.Br mice by negative selection of splenocyte suspensions using anti-CD43 and -CD4 magnetic beads (Miltenyi Biotech) following the manufacturer's instructions. B cells were activated with 100 µg/ml LPS (Sigma) for 24 hrs in RPMI-1640 media, washed, and loaded with 50 µM MCC peptide for 5 hrs.

Suppression of Protein Expression by siRNA electroporation

AND T cells were activated as previously described. 96h after activation, proliferating CD4 T cells were transfected with 4ug of siRNA duplexes either against the desired target or a negative control sequence. Electroporation reagents were purchased from Amaxa Biosystems. Custom siRNA duplexes were purchased from Dharmacon, Inc. After electroporation, the cells were rested for 4h before switching the media to supplemented T cell media containing IL2. 48h later, the cells were once again electroporated with either specific or control siRNA. Cells were used for experiments 48h after the second electroporation.

Bilayers

Glass-supported DOPC bilayers incorporating GPI-anchored forms of I-E^k, CD80, and ICAM-1 were applied to positively charged glass coverslips treated with 70% H₂SO₄/30% H₂O₂. This treatment allows lipid bilayers to adhere to coverslips while permitting lateral mobility. These cover slips were then incorporated into flow cells (Bioptechs) as previously described (Grakoui et al., 1999; Varma et al., 2006).

For biochemistry experiments, bilayers were incubated with 5 µM silica beads (Bangs Laboratories) for 10 minutes at room temperature. Blocking and loading of His-tagged proteins was then performed at equivalent concentrations used in preparation of flow cells. AND T cells were made to form conjugates with beads by gentle co-sedimentation with bilayer-coated beads and incubation at 37°C in HBS/HSA.

TIRFM Imaging

TIRF alignment and imaging was performed as previously described (Varma et al., 2006). AND T cells were suspended in HBS/HSA, and were labeled with 5 µg/ml Alexa568 conjugated H57 Fab. The labeled cells were made to interact with the bilayers at 37°C. For TCR tracking purposes, images were collected at an interval of 5 s in the constant presence of labeled anti-TCR Fab. In wide-field or TIRFM mode, images were exposed for 500-800 msec.

Image Analysis and Statistics

Images were acquired using IPLab software. Images were background subtracted using ImageJ. All data was analyzed with the Metamorph Software, with the exception of calcium measurements and tracking, which was analyzed with Volocity. Statistical analyses were done using GraphPad software. Mann-Whitney and Fisher's Exact tests were used for continuous and categorical variables, respectively.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

We thank W. Sundquist for human TSG101 expression plasmids, U. Ungewickell for the clathrin light chain-GFP fusion expression plasmid, and J. Gruenberg for the anti-LBPA monoclonal antibody. This work was supported by

NIH grant T32 GM07308 (S.V.), a Cancer Research Institute Fellowship (K.C.), and NIH grants R01 AI043542 and PN2 EY016586 (M.L.D.).

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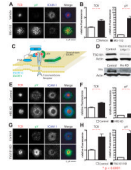


Figure 1. TSG101 is required for central TCR accumulation and dephosphorylation

a-b) AND T cells were pretreated with DMSO (vehicle) or the proteasomal inhibitor MG132 to deplete cells of free ubiquitin. They were then incubated on glass-supported planar bilayers containing 10 molec/ μm^2 I-E^k-MCC and 200 molec/ μm^2 ICAM-1 for 30 minutes at 37°C, fixed, and stained for TCR and phosphotyrosine (pY). Bar graphs (b) depict quantification in arbitrary units (AU) of mean fluorescence at cell contact interfaces, and represent >100 cells across 2 or more experiments. Activated AND Tg T cells were transduced with either control or TSG101-specific siRNA. c) A model for ESCRT recruitment to ubiquitinated substrates. HRS, the critical component of ESCRT-0, binds ubiquitin through its UIM domain but requires recruitment to PI-3-P-rich endosomes through its FYVE domain. TSG101, the critical component of ESCRT-I, binds ubiquitin through its UEV domain. d) Cell lysates of 10⁶ control TSG101 knockdown, and HRS knockdown (KD) T cells were separated by SDS-PAGE and blotted for TSG101 and HRS. Anti-alpha actin was used as a loading control. e-f) Control and HRS KD T cells were incubated on glass-supported planar bilayers containing 10 molec/ μm^2 I-E^k-MCC and 200 molec/ μm^2 ICAM-1 for 30 minutes at 37°C, fixed, and stained for TCR and pY. g-h) Control and TSG101 KD T cells were incubated on glass-supported planar bilayers containing 2 molec/ μm^2 I-E^k-MCC and 200 molec/ μm^2 ICAM-1 for 30 minutes at 37°C, fixed, and stained for TCR and pY.

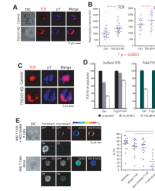


Figure 2. TSG101 controls central TCR accumulation and dephosphorylation in T cell-B cell conjugates

Control and TSG101 KD T cells mixed with MCC-pulsed activated B cells at a 1:2 ratio, and brought into contact by brief centrifugation. Following incubation at 37°C for 30 minutes, cells were fixed, stained for TCR and pY, and placed on clean coverslips for confocal microscopy. Representative images (a) and dot plots (b) demonstrate distribution and quantification of TCR and phosphotyrosine (pY) in Control and TSG101 KD synapses. For en face synapse reconstructions (c), z-stacks of appropriately oriented cell conjugates were collected at 200 nm steps to a distance of ~8 μm from the coverslip surface. d) Control or TSG101 KD T cells were incubated with A20 B cells expressing I-E^k and loaded with either 10 μM or 100 μM MCC. After 30 minutes, they were cooled to 4°C and either stained directly for surface TCR levels or fixed with 2% PFA, permeabilized with 90% methanol, and stained for total TCR levels using antibodies against the AND TCR Vβ3 region. TCR levels were assessed by flow cytometry. Values are reported as a percent of TCR on T cells in culture with unloaded A20 cells. e) TSG101-GFP expressing AND T cells were labeled briefly with CM-DiI and incubated either alone or mixed with MCC-pulsed activated B cells at a 1:2 ratio, and brought into contact by brief centrifugation. Following incubation at 37°C for 30 minutes, cells were fixed and placed on clean coverslips for confocal microscopy. Cells were imaged for GFP and DiI fluorescence prior to and following photobleaching of DiI (Pre and post bleach). D1 and D2 denote donor (GFP) fluorescence pre and post bleach of DiI. D2-D1 is a difference image, and D2-D1/D2 is the FRET image. Both are depicted in pseudo-color (see color key). White represents highest value. Graph (right) represents apparent FRET ($E_a = D2-D1/D2 \times 100\%$) for different regions of the T cell. Lines indicate comparisons, and numbers are P values for comparisons of sample means by ANOVA.

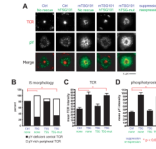


Figure 3. Sorting of TCR into the IS center depends on ubiquitin recognition by TSG101

Control or TSG101 KD T cells were rescued by infection with retroviral siRNA-resistant constructs containing either wild-type human TSG101 or a construct containing mutations in the UEV domain that abrogates ubiquitin recognition (N45A D46A) (Pornillos 2002). Human TSG101 shares 94.6% amino-acid sequence homology with its murine orthologue.

a) Shown are T cells transfected with the indicated siRNA (blue) and re-expression vectors forming IS on glass-supported planar bilayers containing 2 molec/ μm^2 I-E^k-MCC and 200 molec/ μm^2 ICAM-1. Cells were incubated on bilayers for 30 minutes, fixed and stained for TCR and pY. Rescue vectors are indicated as wild-type (TSG101) or ubiquitin-binding mutant (TSG-mut). b) Synapses were categorized by cSMAC morphology based on central accumulation and phosphotyrosine enrichment. Bar graphs (c-d) depict quantification in arbitrary units (AU) of mean fluorescence at cell contact interfaces, and represent >100 cells across 2 or more experiments.

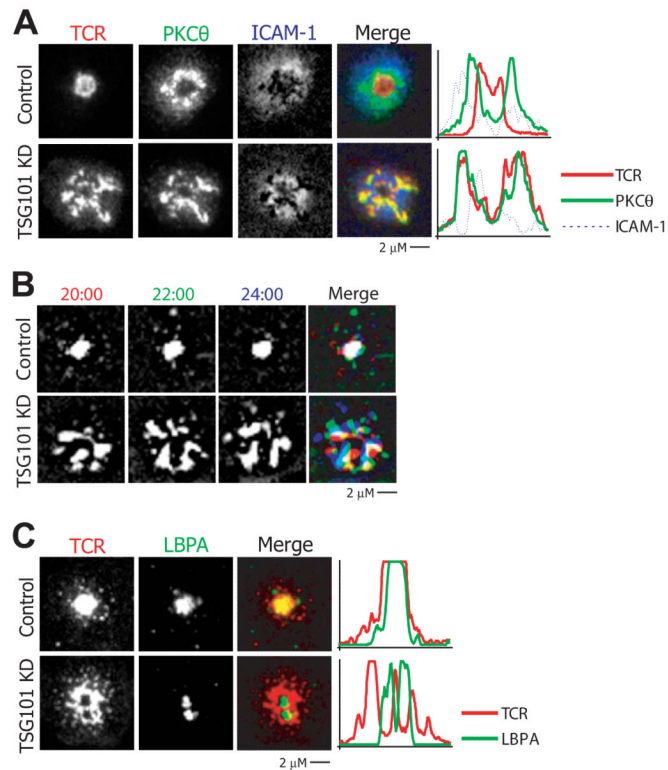


Figure 4. TSG101 controls sorting within plasma membrane and endosomal compartments
 a) Control and TSG101 KD T cells were incubated on glass-supported planar bilayers containing 10 molec/ μm^2 I-E^k-MCC, 100 molec/ μm^2 CD80, and 200 molec/ μm^2 ICAM-1 for 30 minutes, fixed, and stained for TCR and PKC θ . Images display representative distribution of TCR and PKC θ in Control and TSG101 KD synapses. b) Control and TSG101 KD T cells were incubated on glass-supported planar bilayers containing 2 molec/ μm^2 I-E^k-MCC and 200 molec/ μm^2 ICAM-1 and allowed to form contacts for 20 minutes at 37°C. They were then imaged by TIRFM in the presence of fluorescent anti-TCR Fab. White regions indicate immobilization (overlap of red, green, blue), while any individual color (red green, or blue) indicates movement. c) Control and TSG101 KD T cells were incubated on glass-supported planar bilayers containing I-E^k-MCC and ICAM-1 for 30 minutes at 37°C, fixed, and stained for TCR and LBPA. Images show representative distribution of TCR and LBPA in Control and TSG101 KD synapses.

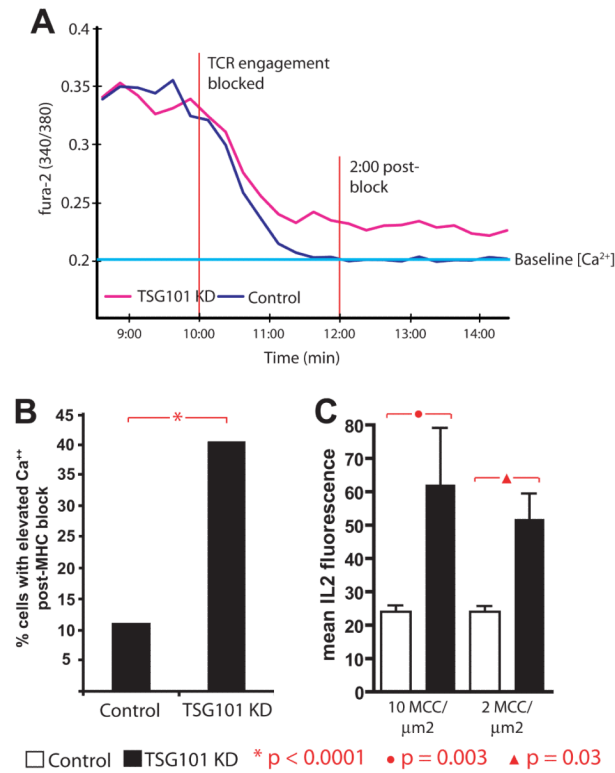


Figure 5. TSG101 suppression leads to chronic TCR signaling

Single-cell intracellular calcium measurements were performed on Fura-2 loaded (a) control (n= 91) and (b) TSG101 KD (n= 57) T cells following incubation on glass-supported planar bilayers containing 2 molec/ μm^2 I-E^k-MCC and 200 molec/ μm^2 ICAM-1 for 10 minutes at 37°C. Cells were then treated with saturating concentrations (30 $\mu g/mL$) of anti-MHC blocking antibodies. Each track depicts individual cell intracellular calcium signal presented as a ratio of fluorescence at 510 nm when excited at 340/380 nm. c) Persistent calcium elevation following anti-MHC block was determined as follows: Normally, anti-MHC treatment eliminates elevated Ca^{2+} levels within 2 minutes. Cells were scored as positive for persistent signaling if intracellular Ca^{2+} levels were 50% above baseline calcium levels for at least 3 minutes after treatment. Baseline Ca^{2+} values were determined by imaging cells on bilayers containing ICAM-1 alone. d) Control and TSG101 KD T cells were loaded with an IL2 capture reagent (Miltenyi) and then incubated on glass-supported planar bilayers containing 2 molec/ μm^2 I-E^k-MCC and 200 molec/ μm^2 ICAM-1 for 6 hours. Cells were fixed, stained with fluorescent anti-IL2-PE antibodies, and imaged for total PE fluorescence. Bar graphs depict quantification of total IL2 fluorescence per cell in arbitrary units (AU), and represent >100 cells across 2 or more experiments.

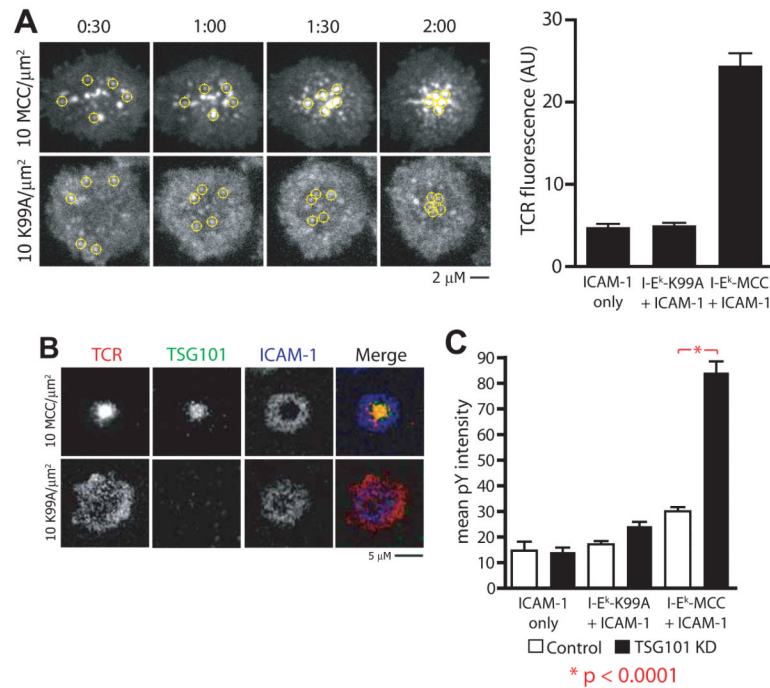


Figure 6. Low avidity ligands signal through microclusters but avoid ESCRT-mediated TCR downregulation

a) AND T cells were incubated on glass-supported planar bilayers containing 200 molec/ μm^2 ICAM-1, 100 molec/ μm^2 CD80, and 10 molec/ μm^2 I-E^k-MCC or I-E^k-K99A as indicated, and immediately imaged by TIRFM at 37°C in the presence of fluorescent anti-TCR Fab. Yellow circles indicate tracks of individual TCR MCs. Bar graph (right) depicts quantification in arbitrary units (AU) of mean TCR fluorescence at cell contact interfaces in IS formed by AND T cells in response to ICAM-1, ICAM-1 + I-E^k-K99A, or ICAM-1 + I-E^k-MCC, and represent >100 cells across 2 or more experiments. b) AND T cells were incubated on glass-supported planar bilayers containing 200 molec/ μm^2 ICAM-1, 100 molec/ μm^2 CD80, and 10 molec/ μm^2 I-E^k-MCC or I-E^k-K99A as indicated for 30 minutes at 37°C, fixed, and stained for TCR and TSG101. Images show representative distribution of TCR and TSG101 in IS formed in response to I-E^k-MCC or I-E^k-K99A. c) Control and TSG101 KD T cells were incubated on glass-supported planar bilayers containing 200 molec/ μm^2 ICAM-1, 100 molec/ μm^2 CD80, and 10 molec/ μm^2 I-E^k-MCC or I-E^k-K99A as indicated for 30 minutes at 37°C, fixed, and stained for TCR and pY. Bar graphs depict quantification in arbitrary units (AU) of mean pY fluorescence at cell contact interfaces, and represent >100 cells across 2 or more experiments.

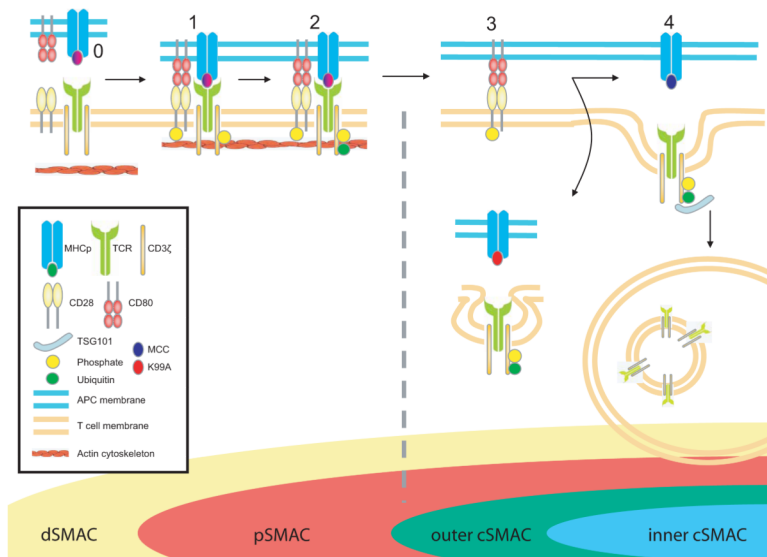


Figure 7. A model for IS patterning

TCR and CD28 engagements occur primarily in the IS dSMAC, resulting in receptor phosphorylation (1), actin-driven centripetal movement within microclusters (depicted as single receptors for simplicity), and ubiquitination of TCR, but not CD28 complexes (2). Upon reaching the pSMAC/cSMAC border, TSG101 specifically sorts strong agonist-engaged ubiquitinated TCR into central MVB/lysosomal compartments (4), leaving CD28 along with associated PKC θ (not depicted) to accumulate and continue signaling from an annular cluster (3). Weak agonist-engaged TCR clusters are not subject to TSG101-mediated sorting, and are thus not incorporated into the cSMAC. The broken grey line indicates receptor signaling state and stage at which TSG101 suppression affects signaling downregulation and IS patterning. For simplicity only a single CD3 ζ chain (of dimeric signaling subunits $\zeta\zeta$, $\epsilon\delta$ and $\epsilon\gamma$) is shown associated with the TCR, and associated proximal signaling molecules (such as Lck, ZAP-70 and LAT) are not depicted.