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Selective Phosphorylation of the Dlg1AB Variant Is Critical for TCR-Induced p38 Activation and Induction of Proinflammatory Cytokines in CD8+ T Cells

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CD8+ T cells respond to TCR stimulation by producing proinflammatory cytokines, and destroying infected or malignant cells through the production and release of cytotoxic granules. Scaffold protein Discs large homolog 1 (Dlg1) specifies TCR-dependent functions by channeling proximal signals toward the activation of p38-dependent proinflammatory cytokine gene expression and/or p38-independent cytotoxicity. Two Dlg1 variants are expressed in CD8+ T cells via alternative splicing, Dlg1AB and Dlg1B, which have differing abilities coordinate TCR-dependent functions. Although both variants facilitate p38-independent cytotoxicity, only Dlg1AB coordinates p38-dependent proinflammatory cytokine expression. In this study, we identify TCR-induced Dlg1 tyrosine phosphorylation as a key regulatory step required for Dlg1AB-mediated p38-dependent functions, including proinflammatory cytokine expression. We find that Dlg1AB but not Dlg1B is tyrosine phosphorylated by proximal tyrosine kinase Lck in response to TCR stimulation. Furthermore, we identify Dlg1 tyrosine 222 (Y222) as a major site of Dlg1 phosphorylation required for TCR-triggered p38 activation and NFAT-dependent expression of proinflammatory cytokines, but not for p38-independent cytotoxicity. Taken together, our data support a model where TCR-induced phosphorylation of Dlg1 Y222 is required for TCR-triggered p38 activation and NFAT-dependent expression of proinflammatory cytokines, but not for p38-independent cytotoxicity. We propose blocking Dlg1AB phosphorylation as a novel therapeutic target to specifically block proinflammatory cytokine production but not cytotoxicity. The Journal of Immunology, 2014, 193: 2651–2660.

Proper activation of CD8+ CTLs is essential for maintaining adaptive immunity to intracellular pathogens. In response to TCR stimulation, CTLs produce and release proinflammatory cytokines and destroy infected or transformed cells through targeted release of cytotoxic granules. TCR engagement is coupled to downstream effector function through the recruitment and activation of proximal tyrosine kinases Lck and Zap70. These early activation events initiate a variety of signaling networks including MAPKs ERK, JNK, and p38, which each regulate transcription factors including NFAT and NF-kB. This group of transcription factors controls the expression of genes that dictate TCR-dependent events including: proliferation, differentiation and effector function (1, 2). Recent studies demonstrate that CD8+ T cell activation is not a binary event, but instead represents a spectrum of tightly controlled biological responses, suggesting that TCR-proximal activation events are selectively coupled to specific downstream signaling networks and functions (3). However, the mechanism(s) by which CTL functionality is specified downstream of the TCR remains incompletely understood.

Scaffold proteins have emerged as key points of control that couple extracellular stimuli to intracellular signaling networks and downstream functions, through the formation multicomponent signaling complexes. One such scaffold protein, Discs Large Homolog 1 (Dlg1), localizes to the junction between T cells and APCs known as the immunological synapse, where it regulates Ag-dependent cytoskeletal and signaling events (4–9). The ability of Dlg1 to regulate cellular functions is attributed to its ability to associate with important cytoskeletal regulators and signal transducers through one or more of its modular protein interaction domains (4, 5, 7). These domains include three PSD-95/Dlg/ZO-1 (PDZ) domains, an SH3 domain and a catalytically inactive guanylate kinase (GUK) domain common to all membrane-associated GUK (MAGUK) scaffolds, as well as an L27 oligomerization domain, an N-terminal proline rich region and a C-terminal HOOK domain, which are unique to Dlg1. Dlg1 domain structure can be modified by alternative splicing events that are thought to affect Dlg1 stability, localization and function (10–15).

We and others have demonstrated that Dlg1 specifies TCR signal transduction, by coordinating the activation of p38 through the alternative pathway (5, 7, 16). The alternative p38 pathway is initiated downstream of the TCR, and requires the activity of proximal tyrosine kinase Lck and Zap70 (17). TCR engagement...
triggers activation of Zap70, which phosphorylates p38 at Y323, triggering p38 autophosphorylation at T180 and subsequent up-regulation of p38 kinase activity (17, 18). This pathway is distinct from the canonical p38 pathway, which is triggered by environmental stress and results in direct phosphorylation of p38 at T180 and Y182 by MKK3 or MKK6 (19). A vital downstream target of alternatively activated p38 is the transcription factor NFATc1 (NFAT2), which when phosphorylated at SS4 upregulates numerous CD8+ T cell effector functions including proinflammatory cytokine gene expression (20, 21).

We recently demonstrated that at least two Dlg1 variants are expressed in T cells because of alternative splicing, Dlg1AB and Dlg1IB, that differ in the inclusion or exclusion of the proline-rich i1A region (O. Silva, J. Crocetti, L. Humphries, H. Elaesser, D. Brooks, J. Burkhardt, and M.C. Miceli, submitted for publication). Although both variants can mediate cytoskeletal reorganization and release of cytotoxic granules, only the Dlg1AB variant can facilitate alternative p38 activation leading to induction of NFAT-dependent genes including IFN-γ and TNF-α (O. Silva et al., submitted for publication). The ability of Dlg1AB to activate p38 has been attributed to interaction with proximal tyrosine kinase Lck; however, the mechanism by which Dlg1:Lck association facilitates p38 activation was unclear.

Despite having no inherent enzymatic activity, Dlg1 is dynamically regulated by posttranslational modifications and intramolecular interactions that induce changes in protein stability and binding partners that dictate the scope of Dlg1 function (22–25). Tyrosine phosphorylation of scaffold proteins can lead to conformational changes and creation of new binding sites that play a vital role in scaffold protein function (26, 27). In CD8+ T cells, Dlg1 associates with several tyrosine kinases, including proximal tyrosine kinase Lck and Zap70 (5). However, a role for tyrosine phosphorylation in regulating Dlg1 function in T cells has not been described previously.

In this paper, we report that scaffold protein Dlg1 is phosphorylated on tyrosine residues in response to TCR stimulation. This phosphorylation event occurred specifically on the Dlg1AB splice variant and can be mediated by Dlg1-associated proximal tyrosine kinase Lck. We identified Y222 as a major site of TCR-dependent Dlg1 phosphorylation and demonstrated that loss of Y222 phosphorylation in the Y222F mutant significantly impaired TCR-induced p38 phosphorylation and NFAT-dependent expression of proinflammatory cytokines IFN-γ and TNF-α. Conversely, Dlg1 Y222 phosphorylation was not required for functions mediated by Dlg1B including degranulation. Taken together, our data identify Dlg1 Y222 phosphorylation as a key point of control that selectively impacts Dlg1AB-mediated functions such as proinflammatory cytokine expression. We propose Dlg1 Y222 phosphorylation as a novel therapeutic target to selectively block proinflammatory cytokine production while not impacting other immune functions such as target cell killing.

Materials and Methods

Mice

The generation of Dlg1flox/flox mice has been described previously (28). These mice were crossed with C57BL/6 mice expressing the OT-1 TCR. All mice were maintained and used in accordance with the University of California Los Angeles Chancellor’s Animal Research Committee.

In vitro kinase assays

OT-1 hybridoma cells (4 × 10^6) were lysed using an immunoprecipitation lysis buffer (Pierce) in the presence of protease and phosphatase inhibitors (Pierce). Resulting lysates were cleared by centrifugation and incubated with wild-type (WT) or Y222F GST-Dlg1 fusion proteins bound to glutathione–Sepharose bead slurry and then washed twice with kinase buffer (Cell Signaling Technology). Resulting bead complexes were incubated at 30˚C for 20 min in 50 μl kinase buffer in the presence or absence of ATP (Cell Signaling Technology). Alternatively, WT or Y222F GST-Dlg1 fusion proteins were bound to glutathione Sepharose beads and incubated with recombinant Lck or Zap70 (Active Motif) in 50 μl kinase buffer in the presence or absence of ATP at 30˚C for 20 min.

Dlg1 knockout and expression

Stable Dlg1 knockout lines were created in both OT-1 hybridomas using an mIR-155-based retroviral knockdown vector (29). Sense and antisense sequences specific for the 3′-untranslated region (3′-UTR) of Dlg1 (5′-GTCTCTCCACGTACAGCAGAT-3′) were cloned in to the mIR-155–based knockdown vector. Virus was produced by transfecting 293T cells with 25 μg knockdown vector and 25 μg pcDNA-e vector (Mirus). Resulting virus was used to transfect OT-1 hybridomas for 90 min at 1250 × g on two consecutive days. Cells were allowed to rest for 1 d and then assessed for Dlg1 knockdown. Dlg1 was reintroduced into these cell lines using a murine stem cell virus–based retroviral vector expressing a Dlg1 variant or mutant using the same viral production and transduction methods used for the knockdown virus.

Flow cytometry

For measurement of Dlg1, 1 × 10^6 cells were fixed and permeabilized using FoxP3 staining buffer set (00-5523-00; eBioscience), according to the manufacturer’s instructions. Dlg1 was detected using αDlg1 Ab (610875; BD Biosciences) at 1:1000, followed by Alexa Fluor 647–conjugated donkey anti-mouse IgG F(ab’)2 (715-065-150; Jackson ImmunoResearch Laboratories) at 1:1000. For measurement of p38 phosphorylation, expanded primary CD8+ T cells were harvested, counted, and placed in fresh medium at a concentration of 2 × 10^6 cells/ml. Cells were allowed to rest for 4 h at 37˚C in 6-well dishes at volume of 2 ml/well and were transferred to 12-well plates coated with 5 μg anti-CDS (clone 14-2C11; 553057; BD Biosciences) and incubated for 15 min at 37˚C. Cells were fixed in paraformaldehyde (4% final concentration) for 30 min, followed by permeabilization using the FoxP3 staining buffer set (00-5523-00; eBioscience), according to manufacturer’s instructions. Phosphorylated p38 T180/Y182 was detected using Alexa Fluor 647–conjugated anti-p-p38 Ab (T309/Y320; 560646; BD Biosciences). Cells were surface stained using CD8a-PE and fixed. Events were collected using FACS Calibur (BD Biosciences) and analyzed using FlowJo software.

Immunoprecipitation and blotting

For immunoprecipitation, T cell lysates were incubated with 20 μl 50% (v/v) protein G–Sepharose beads (2625; Cell Signaling Technology), GST (2625; Cell Signaling Technology), or phosphorylated tyrosine kinase Lck (clone mAb2C11; 553057; BD Biosciences). Cells were washed and boiled with Laemmli buffer. Blotting was performed according to manufacturer’s instructions. Dlg1 was detected using anti-Dlg1 Ab (mAb1Lck) in 1:1000. For measurement of p38 phosphorylation, expanded primary CD8+ T cells were harvested, counted, and incubated for 5 or 15 min at 37˚C to stimulate. Cells were fixed in paraformaldehyde (4% final concentration) for 30 min, followed by permeabilization using the FoxP3 staining buffer set (00-5523-00; eBioscience), according to manufacturer’s instructions. Phosphorylated p38 T180/Y182 was detected using Alexa Fluor 647–conjugated anti-p-p38 Ab (T309/Y320; 560646; BD Biosciences). Cells were surface stained using CD8a-PE and fixed. Events were collected using FACS Calibur (BD Biosciences) and analyzed using FlowJo software.

RNA isolation, Reverse transcription, and quantitative PCR

RNA was isolated using TRIzol reagent as described previously (5). Two micrograms of RNA was reverse transcribed using Superscript II reverse transcriptase (Invitrogen), according to the manufacturer’s instructions, using random hexamer and oligo(dT)12-18 as primers. The iCycler thermocycler was used for quantitative PCR analysis according to the manufacturer’s instructions (Bio-Rad). A final volume of 25 μl was used for each quantitative PCR as described previously (5). The following gene-specific primers were used for amplification: L32(F), 5′-AAG CGAAGCCGCGGAAGAAAC-3′, and L32(R), 5′-TAAAACCAGTTTGGCCATCAG-3′; NFATc1(F), 5′-GGCTGTCTGATTGCTGGAAG-3′; NFATc1(R), 5′-GGAAAGCAGTTGATTTGAACAAA-3′; IκBα(F), 5′-CTGGAGGCCACACATCAACA-3′, and IκBα(R), 5′-CAGCACCCAAAATGGCAGATG-3′; IFN-γ(F), 5′-GTCACAGAACCCCAAGGGGCAG-3′, and IFN-γ(R), 5′-CCCTTTCTCGCTCTCTTGAGG-3′; TNF-α(F), 5′-CCTCCCGGGGCTGTCACCA-3′.
**Results**

Dlg1AB is tyrosine phosphorylated by a Dlg1-associated kinase in response to TCR stimulation

Dlg1 is rich in tyrosine residues and is known to associate with several tyrosine kinases; therefore, we considered the possibility that the signaling potential of Dlg1 may be regulated by phosphorylation of specific tyrosines (5). To determine whether Dlg1 was phosphorylated in T cells, we performed in vitro kinase assays using Dlg1 complexes formed using T cell lysates. Specifically, GST fusion proteins expressing full-length Dlg1AB (GST-Dlg1AB) or full-length Dlg1B (GST-Dlg1B) were incubated with T cell lysates to form Dlg1 complexes with T cell–specific ligands. Upon addition of ATP to these complexes, robust tyrosine phosphorylation (pY-Dlg1) of Dlg1AB but not Dlg1B was observed (Fig. 1A, 1B). These data indicate that Dlg1AB but not Dlg1B associates with and is phosphorylated by at least one kinase found in T cell lysates. To determine whether Dlg1 phosphorylation was TCR induced, Dlg1 complexes were immunoprecipitated from resting or TCR-stimulated T cells, and the level of Dlg1 tyrosine phosphorylation was assessed. We observed an increase in tyrosine phosphorylation of the top band of Dlg1 in TCR-stimulated samples compared with resting samples, which we attributed to Src family kinase activity in response to TCR stimulation.

Tyrosine 222 (Y222) is a major site of TCR-induced Dlg1 phosphorylation

Dlg1AB and Dlg1B differ in the inclusion or exclusion of the proline-rich i1A region, which permits association of Dlg1 with proximal tyrosine kinase Lck (Fig. 2A; O. Silva et al., submitted for publication). The Dlg1AB protein variant contains 927 aa, 29 of which are tyrosines (GenBank accession number AAH57118.1) (Fig. 2A). To narrow our search for specific sites of Dlg1 tyrosine phosphorylation, we used the ScanSite algorithm to determine the most likely candidate tyrosine residues (30). Tyrosine 222 (Y222) was identified as the most likely site of tyrosine phosphorylation using high-stringency search criteria. Y222 is located in the N-terminal region of Dlg1 in the flexible linker region between the i1B and PDZ1 domains. In addition, Y222 is part of a highly conserved Tyr-Glu-Glu-Iso (YEEI) motif that is a predicted target of known Dlg1-associated tyrosine kinases Lck and Zap70 (5, 31, 32) (http://ppsp.biocuckoo.org) (Fig. 2A).

To determine whether this candidate tyrosine was a site of phosphorylation, we created GST-Dlg1AB fusion proteins in which Y222 was mutated to phenylalanine (Y222F) to disrupt phosphorylation at this position. GST-Dlg1AB (WT) or GST-Dlg1AB Y222F (Y222F) fusion proteins were used to form complexes with T cell ligands and subsequently in vitro phosphorylated. Upon the addition of ATP, GST-Dlg1AB Y222F consistently showed a 40% decrease in tyrosine phosphorylation signal relative to GST-Dlg1AB demonstrating that Y222 was a major site of Dlg1 phosphorylation (Fig. 2B). Although the region surrounding Y222 is conserved in both variants (Fig. 2A), we found that Dlg1B is not tyrosine phosphorylated, suggesting that the tyrosine kinase responsible for Y222 phosphorylation may not be present in Dlg1B complexes (Fig. 1).

**Dlg1-associated Lck can mediate Dlg1 phosphorylation**

Dlg1 Y222 is part of a YEEI motif predicted to be a target of Src and Syk family kinases (Fig. 2A). To determine a possible role for Src kinases in Dlg1 phosphorylation, we examined Dlg1 phosphorylation in vitro phosphorylated GST-Dlg1AB complexes in the presence or absence of Src kinase family inhibitor PP2. Dlg1 tyrosine phosphorylation was observed upon the addition of ATP, which was completely blocked by the addition of PP2, demonstrating that Src family kinase activity was required for Dlg1 phosphorylation (Fig. 2C). However, because Lck kinase activity regulates Zap70 activation, these data cannot rule out a role for Zap70 in Dlg1 phosphorylation.

To assess the role of these kinases in phosphorylating Dlg1, we investigated whether the kinase activity of each kinase was directed to phosphorylate Dlg1. We found that Lck but not Zap70 directly phosphorylated Dlg1.
Furthermore, phosphorylation of GST-Dlg1AB Y222F by rLck was significantly reduced compared with GST-Dlg1AB, demonstrating that Lck is likely responsible for Y222 phosphorylation (Fig. 2D). Although rZap70 did not phosphorylate Dlg1, it did phosphorylate our positive control (GST-LAT), demonstrating that the kinase was active in our system (Fig. 2D). These results rule out a role for Syk family member Zap70 in Dlg1 phosphorylation. Taken together, our data demonstrate that Src family kinases play an important role in Dlg1 phosphorylation and identify Lck as one possible kinase capable of phosphorylating Dlg1. Although we cannot rule out a role for Src family kinase Fyn, previous studies demonstrating that Lck but not Fyn binds Dlg1 suggest that Lck is likely the preferred Src family member responsible for Dlg1 tyrosine phosphorylation (4).

Dlg1 Y222 phosphorylation is required for TCR-induced alternative p38 activation in CD8+ T cells

Dlg1AB coordinates the activation of p38 in response to TCR stimulation via the alternative activation pathway (5). To assess the requirement of Dlg1 Y222 in coordinating alternative p38 activation, we formed Dlg1 complexes by incubating GST-Dlg1AB or GST-Dlg1AB Y222F with T cell lysates, centrifuged, washed, and incubated with ATP. Protein complexes subjected SDS-PAGE and blotted for using anti-4G10 (pY-Dlg1) and anti-GST (GST-Dlg1). In vitro kinase assay using GST-Dlg1AB fusion proteins bound to glutathione-Sepharose beads and incubated with T cell lysates, centrifuged, washed, and incubated with ATP in the presence or absence of 10 μM PP2 inhibitor. Protein complexes were subjected SDS-PAGE and blotted for using anti-4G10 (pY-Dlg1) and anti-GST (GST-Dlg1). GST-Dlg1AB WT or Y222F fusion proteins were bound to glutathione-Sepharose beads and incubated with ATP and rLck, rZap70, or buffer only (none). Proteins were subjected to SDS-PAGE and blotted for using anti-4G10 (pY-GST-Dlg1 and pY-GST-LAT) or anti-GST (GST-Dlg1). Blots were quantitated using Li-Cor Odyssey software. pY-Dlg1 was normalized to total Dlg1 for each lane, and WT phosphorylation was set to 1.0; error bars represent the SD of four independent experiments. **p < 0.05.

**FIGURE 2.** Dlg1-associated Lck phosphorylates Dlg1 at position tyrosine 222. (A) Domain structure of Dlg1AB and Dlg1B. Asterisks indicate the location of the 29 tyrosine residues along the Dlg1 sequence. The bold asterisk indicates the location of Y222. (A, bottom inset) Amino acid sequence alignment of the region of Dlg1 surrounding Y222 from various all species listed in the National Center for Biotechnology Information Database. Shaded region indicates evolutionarily conserved linker region of Dlg1. (B) In vitro kinase assays using GST-Dlg1AB WT or GST-Dlg1 Y222F (Tyr to Phe mutation) fusion proteins bound to glutathione–Sepharose beads and incubated with T cell lysates, centrifuged, washed, and incubated with ATP. Protein complexes subjected SDS-PAGE and blotted for using anti-4G10 (pY-Dlg1) and anti-GST (GST-Dlg1). (C) In vitro kinase assay using GST-Dlg1AB fusion proteins bound to glutathione–Sepharose beads and incubated with T cell lysates, centrifuged, washed, and incubated with ATP in the presence or absence of 10 μM PP2 inhibitor. Protein complexes were subjected SDS-PAGE and blotted for using anti-4G10 (pY-Dlg1) and anti-GST (GST-Dlg1). (D) GST-Dlg1AB WT or Y222F fusion proteins were bound to glutathione–Sepharose beads and incubated with ATP and rLck, rZap70, or buffer only (none). Proteins were subjected to SDS-PAGE and blotted for using anti-4G10 (pY-GST-Dlg1 and pY-GST-LAT) or anti-GST (GST-Dlg1). (B, right; D, right) Blots were quantitated using Li-Cor Odyssey software. pY-Dlg1 was normalized to total Dlg1 for each lane, and WT phosphorylation was set to 1.0; error bars represent the SD of four independent experiments. **p < 0.05.
FIGURE 3. Dlg1 Y222 phosphorylation is required for TCR-induced alternative p38 activation. (A) In vitro kinase assay were GST-Dlg1AB WT, or GST-Dlg1 Y222F fusion proteins were bound to glutathione–Sepharose beads and incubated with T cell lysates, centrifuged, washed, and incubated with 10 mM ATP for 20 min at 30°C. Protein complexes were boiled off beads and subjected SDS-PAGE, and blotted using anti-4G10 (pY-p38) and anti-p38. Blots were quantitated using Li-Cor Odyssey software. pY-p38 was normalized to total p38 for each lane, and WT phosphorylation was set to 1.0; error bars represent the SD of three independent experiments. **p < 0.05. (B and C) Control, Dlg1-deficient OT-I hybridomas (3′-UTR) re-expressing vector only (3′-UTR+Vector), Dlg1AB (3′-UTR+Dlg1AB), or Dlg1AB Y222F (3′-UTR+Dlg1Y222F) were lysed and subjected to SDS-PAGE and blotted using anti-Dlg1 or anti-p38 (B) or stimulated using 5 μM anti-CD3 and 5 μg/ml CD28 (C), followed by cross-linking with donkey anti-Armenian hamster Ab for 15 min at 37°C, followed by lysis. Dlg1 was immunoprecipitated from resulting lysates and subjected to SDS-PAGE then blotted using anti-p-p38 180/182 and anti-p38. (D and E) CD8+ T cells were isolated from the spleens of OT-I mice and expanded on plate-bound anti-CD3 and anti-CD28 Ab for 72 h, followed by overexpression of Dlg1AB or Dlg1Y222F. Resulting cells were permeabilized and stained for intracellular Dlg1 (D) or stimulated with plate-bound anti-CD3 and CD28 (E) for 15 min, followed by fixation, permeabilization, and staining for intracellular p-p38 180/182. Data presented are representative of three independent experiments.

control cells whereas Dlg1AB Y222F did not (Fig. 3D, 3E). Taken together, our results highlight an essential role for Dlg1 Y222 phosphorylation in the regulation of TCR-induced alternative p38 activation in CD8+ T cells.

Loss of Dlg1 Y222 phosphorylation results in decreased NFAT-, but not NF-kB-dependent expression

Previous studies from our group and others have demonstrated that Dlg1-mediated alternative p38 activation induces NFAT- but not NF-kB-dependent gene expression (5, 7, 16). Recently, we have demonstrated that Dlg1AB but not Dlg1B can facilitate p38-dependent NFAT activation (O. Silva et al., submitted for publication). To determine whether Dlg1 Y222 plays a role in TCR-induced NFAT or NF-kB activation, CD8+ T cells re-expression Dlg1AB or Dlg1AB Y222F were stimulated through their TCR, and the level of inducible *Nfatc1* (NFAT-dependent gene) and *Ikba* (NF-kB–dependent gene) gene expression was measured via quantitative PCR. In agreement with published results, CD8+ T cells lacking Dlg1 demonstrated impaired induction of *Nfatc1* upon TCR stimulation (Fig. 4A). Expression of Dlg1AB in Dlg1 knockout cells was able to rescue *Nfatc1* levels, whereas expression of Dlg1AB Y222F or Dlg1B only further attenuated *Nfatc1* expression (Fig. 4A). Alterations in Dlg1 expression had no significant effect on the level of *Ikba* induction upon TCR stimulation (Fig. 4B). We further interrogated these findings in primary CD8+ T cells where overexpression of Dlg1AB but not Dlg1AB Y222F or Dlg1B was able to enhance TCR-dependent *Nfatc1* transcription, whereas Dlg1 expression levels had no significant effect on *Ikba* transcription (Fig. 4F, 4G). These findings demonstrate that Dlg1 Y222 phosphorylation is required for the specific activation of NFAT- but not NF-kB–dependent transcription in response to TCR stimulation.

*Dlg1 Y222 is required for optimal NFAT-dependent IFN-γ and TNF-α but not IL-2 production*

Dlg1AB-mediated alternative p38 activation has been implicated in selective upregulation of proinflammatory cytokines IFN-γ and TNF-α but not IL-2 in response to TCR stimulation (5, 20, 21, 33, 34) (O. Silva et al., submitted for publication). To examine the requirement of Dlg1 Y222 in TCR-induced cytokine production, CD8+ T cells re-expressing Dlg1AB, Dlg1B, or Dlg1AB Y222F were TCR-stimulated, and their ability to induce cytokine gene expression was measured. Dlg1-deficient CD8+ T cells had impaired induction of IFN-γ and TNF-α but not IL-2 transcription compared with control cells (Fig. 4C–E). Expression of Dlg1AB was able to rescue the defect in IFN-γ and TNF-α whereas Dlg1AB Y222F or Dlg1B was not. In keeping with previous results, expression of Dlg1 had no significant effect on the level of IL-2 transcription (Fig. 4E; O. Silva et al., submitted for publication). In addition, overexpression of Dlg1AB in primary CD8+ CTLs was able to enhance IFN-γ and TNF-α gene expression, whereas expression of Dlg1B or Dlg1AB Y222F led to decreased IFN-γ and TNF-α expression compared with control cells, suggesting that these variants may have a dominant negative effect. Dlg1 expression had no significant effect on IL-2 gene expression (Fig. 4H–J). Taken together, our results support a model where phosphorylation of Dlg1 at Y222 is a key point of control that facilitates selective upregulation of NFAT-dependent gene transcription, including the production of proinflammatory cytokines IFN-γ and TNF-α but not IL-2.

*Dlg1 Y222 is not required for Dlg1-mediated target cell killing by CD8+ T cells*

Dlg1 is known to regulate cytoskeletal events in epithelial, neuronal, and immunologic systems (4, 7, 10, 35). In particular, Dlg1 association with WASp and ezrin is implicated in TCR-dependent positioning of the microtubule-organizing complex in response to antigenic stimulation (7). Recently, we have demonstrated that both Dlg1AB and Dlg1B can facilitate cytotoxic granule and cytokine release through association with and activation of WASp (O. Silva et al., submitted for publication). To test the requirement of Dlg1 Y222 phosphorylation in release of cytotoxic granules (degranulation), we incubated CD8+ T cells re-expressing Dlg1AB or Dlg1AB Y222F with APCs and tracked the surface expression of degranulation marker CD107a. Total Dlg1 knockdown resulted...
in significantly impaired Ag-dependent degranulation. Surprisingly, both Dlg1AB and Dlg1Y222F were able to rescue and enhance Ag-dependent degranulation to a similar level (Fig. 5A). Overexpression of Dlg1AB or Dlg1AB Y222F in primary CD8+ CTLs enhanced Ag-dependent degranulation to a similar level compared with control cells (Fig. 5B). Examination of perforin production in primary CD8+ T cells overexpressing Dlg1AB or Dlg1AB Y222F show no significant differences, suggesting that cytotoxic granule components are not affected by loss of Dlg1 Y222 phosphorylation. In the same experiment, Dlg1AB but not Dlg1Y222F was able to enhance NFAT-dependent transcription. Taken together, these data suggest that phosphorylation of Dlg1 Y222 does not control Dlg1-mediated Ag-dependent degranulation but not production of cytotoxic granule components.

Acute Dlg1 knockout block TCR-dependent p38 activation, NFAT-dependent transcription and target cell killing in CD8+ T cells

We and others have demonstrated that Dlg1 plays a vital role in T cell development, activation and effector responses utilizing Dlg1 knockdown and overexpression systems (4, 5, 7, 9, 10, 16). However, initial knockout studies that ablate Dlg1 early in T cell development do not show striking functional differences, likely because of compensation from other Dlg1 family members (36, 37). Therefore, we developed an acute Dlg1 knockout system to examine the requirement for Dlg1 in TCR-dependent effector functions. CD8+ T cells isolated from the spleens of OT-1 mice and expanded on plate-bound anti-CD3 and anti-CD28 Ab for 72 h, followed by overexpression of Dlg1AB, Dlg1B, Dlg1Y222F, or empty vector (control). (A–J) Cells were stimulated with anti-CD3 and anti-CD28 Ab for 2 or 6 h, followed by mRNA isolation, reverse transcription, and qPCR analysis using primers for NFATc1 (A and F), IκBα (B and G), IFN-γ (C and H), TNF-α (D and I), or IL-2 (E and J). All values were normalized to L32 and unstimulated values set to 1.0. Error bars represent SD of triplicates; data are representative of three independent experiments. **p < 0.01.

Discussion

CD8+ T cells have the ability to develop a broad range of functionality, with differing capacities to produce and release cytokines and lytic factors in response to TCR stimulation (3). Scaffold protein Dlg1 specifies signaling and function downstream of the TCR by facilitating p38 and NFAT activation leading to the production of proinflammatory cytokines. In ad-
FIGURE 5. Dlg1 Y222 phosphorylation is not required for Ag-dependent CTL degranulation. (A) Control, Dlg1-deficient OT-1 hybridomas (3'-UTR) reexpressing vector only (3'-UTR+Vector), Dlg1AB (3'-UTR+Dlg1AB), or Dlg1AB Y222F (3'-UTR+Dlg1AB Y222F) (A) or CD8+ T cells were isolated from the spleens of OT-1 mice and expanded on plate-bound anti-CD3 and anti-CD28 Ab for 72 h, followed by overexpression of Dlg1AB, Dlg1B, Dlg1Y222F, or empty vector (control) cells were incubated with EG.7 thymoma target cells at indicated E:T ratios for 2 h in the presence of anti-CD107a Ab. CD107a was assessed by flow cytometry where control cells at E:T of 1:0 was set to 5%. Data are representative of three experiments. (B and C) CD8+ T cells were isolated from the spleens of OT-1 mice and expanded on plate-bound anti-CD3 and anti-CD28 Ab for 72 h, followed by overexpression of Dlg1AB, Dlg1Y222F, or empty vector (control). (B) CTLs were incubated with EG.7 thymoma target cells at indicated E:T ratios for 2 h in the presence of anti-CD107a Ab. CD107a was assessed by flow cytometry where control cells at E:T of 1:0 was set to 5%. (C) CTLs were stimulated with anti-CD3 and anti-CD28 Ab for 6 h, followed by mRNA isolation, reverse transcription, and qPCR analysis using primers for perforin. **p < 0.05.

dition. Dlg1 coordinates TCR-dependent activation of WASp leading to cytoskeletal reorganization and release of cytotoxic granules (4, 5) (O. Silva et al. submitted for publication). These functions can be differentially regulated by the two Dlg1 splice variants expressed in CD8+ T cells: Dlg1AB and Dlg1B (O. Silva et al., submitted for publication). Both Dlg1AB and Dlg1B facilitate cytotoxicity, whereas only Dlg1AB can coordinate p38 activation and proinflammatory cytokine production. The ability of Dlg1AB to activate p38 has been attributed to its interaction with proximal tyrosine kinase Lck; however, the mechanism by which Dlg1AB but not Dlg1B coordinates p38 activation was unclear. In this study, we identify TCR-induced phosphorylation of Dlg1AB at Y222 as a key point of control required for TCR-induced, Dlg1-mediated p38 activation and expression of proinflammatory cytokines but not cytotoxic granule release.

Dlg1 is rich in tyrosine residues of uncharacterized function and is known to associate with several tyrosine kinases; therefore, we hypothesized that tyrosine phosphorylation may play an important role in regulating Dlg1 function. We found that Dlg1 was tyrosine phosphorylated in response to TCR stimulation in a splice variant dependent manner, where Dlg1AB but not Dlg1B was phosphorylated. We determined that Y222 in the N-terminal linker region of Dlg1 is a phosphorylation site likely targeted by Dlg1-associated Lck. The loss of Y222 phosphorylation impaired TCR-induced p38 activation and NFAT-dependent gene expression of proinflammatory cytokines IFN-γ and TNF-α but not IL-2. Surprisingly, we found that Dlg1 phosphorylation is not required for TCR-mediated degranulation. These data highlight the role of Dlg1 Y222 phosphorylation as a key point of control required for the activation of a subset of effector functions downstream of the TCR. In support of the importance of Dlg1 in regulating T cell function, we demonstrated that acute knockout of Dlg1 in CD8+ CTLs resulted in selective impairment of TCR-induced p38 activation, NFAT-dependent transcription, proinflammatory cytokine gene expression, and Ag-dependent degranulation. Our results indicate that Dlg1 plays an important role in specifying CD8+ T cell function and that Lck-mediated phosphorylation at Y222 is a molecular switch required to authorize a subset of Dlg1-mediated functions.

At least two Dlg1 protein variants, Dlg1AB and Dlg1B, are expressed in CD8+ T cells through alternative splicing that differ in the inclusion or exclusion of the proline-rich i1A region (O. Silva et al., submitted for publication). The inclusion of the i1A region allows Dlg1AB to associate with tyrosine kinase Lck and facilitate p38 activation and NFAT-dependent transcription of proinflammatory cytokine genes in response to TCR stimulation. In contrast, Dlg1AB and Dlg1B are both capable of mediating Ag-dependent degranulation, suggesting that direct association with Lck is not required for coordinating these functions (O. Silva et al., submitted for publication). In this study, we showed that Dlg1AB but not Dlg1B is phosphorylated in response to TCR stimulation and that this phosphorylation is dependent on Dlg1-associated Lck. We also identify Y222 as an Lck-dependent phosphorylation site required for TCR-induced p38 activation and downstream transcriptional activation but not Ag-dependent degranulation. Therefore, we propose that the differential functionality of the two known Dlg1 splice variants is regulated, at least in part, by differential ability to be phosphorylated on tyrosine residues.

Scaffold protein structure, and therefore function, is often regulated through a series of phosphorylation and/or ligand binding events that lead to the gradual unfolding of the scaffold into an active conformation (39–41). These phosphorylation events are hypothesized to provide molecular memory by creating a primed scaffold capable of binding new ligands, endowing the scaffold with novel and/or heightened functionally upon secondary stimulation. Dlg1 forms intramolecular interactions yielding at least two distinct closed conformations hypothesized to impact Dlg1 function by masking ligand binding sites or affecting the juxtaposition of bound ligands (22, 23). However, a role for these intramolecular interactions has not been characterized in the context of Dlg1 ligands in T cells. In this study, we showed that tyrosine phosphorylation of Dlg1 in the N-terminal linker region is required for TCR-induced p38 activation. We predict that phosphorylation at Dlg1 Y222 causes a conformational change required for proper association of Dlg1 with ligands such as Zap70. Alternatively, structural changes could affect the positioning of Dlg1-associated proteins relative to each other. Therefore, the closed conformation could disrupt signal propagation on the scaffold even in the presence of all required ligands. Allosteric regulation has been described in other MAGUK family members including CARMA1, which facilitates activation of NF-κB in T and B cells through formation of a signaling complex with Bcl10 and MALT1 (42). Inducible phosphorylation of serine residues in the linker region of CARMA1 regulates its structure, binding partners and function.
FIGURE 6. Acute knockout of Dlg1 impairs p38 activation and NFAT-dependent proinflammatory cytokines gene expression and target cell lysis in response to TCR stimulation. (A) Genomic organization of Dlg1flox/flox mice as described previously (28). F and R refer to the location of the forward and reverse primers for gDNA analysis. (B–L) CD8+ T cells were isolated from spleens of OT-1 Dlg1flox/flox mice and expanded on plate-bound anti-CD3 and anti-CD28 Ab for 48–72 h, followed by infection with retroviral cre recombinase or vector control. (B) Genomic DNA was isolated, and PCR analysis was performed. (C) Whole-cell lysates were subjected to SDS-PAGE and blotted with anti-Dlg1 or anti-p38. (D) Cells were stimulated with plate-bound anti-CD3 and anti-CD28 Ab for 15 min, followed by fixation, permeabilization, and staining for p-p38 180/182. Error bars represent SD of three independent experiments. (E–J) Cells were left unstimulated or stimulated with plate-bound anti-CD3 and anti-CD28 for 2 h (E and F) or 6 h (G–J), followed by mRNA isolation, reverse transcription, and qPCR analysis using primers specific for NFATc1(E), IκBα (F), IFN-γ (G), TNF-α (H), IL-2 (I), or granzyme B (J). All values were normalized to L32 and unstimulated values set to 1.0. Error bars represent standard deviation of triplicates; data are representative of four independent experiments. (K and L) Cells were incubated with EG.7 thymoma cells at indicated ratios for 2 h, followed by surface staining for CD107a (K) or lactate dehydrogenase cytotoxicity assay (L). **p < 0.01, ***p < 0.001.

FIGURE 7. Model of Dlg1-dependent signaling downstream of the TCR. In response to TCR engagement, Dlg1-bound Lck phosphorylates Dlg1AB at several tyrosine residues including Y222. This phosphorylation event empowers Dlg1 to enhance p38 activation and NFAT-dependent transcription of IFN-γ and TNF-α. Dlg1AB or Dlg1B is able to facilitate Ag-induced degranulation and target cell killing in a phosphorylation-independent manner.

Using knockdown methodologies several groups have demonstrated a role for Dlg1 in regulating functionality and differentiation in both CD4+ and CD8+ T cells (4, 5, 7, 16) (O. Silva et al., submitted for publication). Despite growing evidence for Dlg1 in specifying signal transduction downstream of the TCR, recent attempts by numerous groups to extend these studies to knockout models have been inconsistent and largely unavailing (36, 43, 44). The mild phenotype seen in these models is attributed to compensation of other Dlg1 family members; such as PSD-95, which is known to have functional redundancy with Dlg1 in neurons (37). In this study, we have described an acute Dlg1 knockout system where cre recombinase is overexpressed in CD8+ CTLs from Dlg1flox/flox mice, and TCR-triggered functions are assessed 72 hr later. In this system, we demonstrate that TCR-triggered p38 activation, NFAT activation, induction of IFN-γ and TNF-α gene expression, and degranulation are impaired in the Dlg1flox mice compared with controls.

The spectrum of effector functions developed by T cells during infection is predicted to optimize T cell responsiveness to specific pathogens (3). Indeed, development of polyfunctional CD8+ T cells is correlated with improved clinical outcomes against HIV and tumors, whereas dysregulation of CD8+ T cell function is implicated in persistence of chronic viral infections and autoimmune disorders (45–49). Therefore, it is essential to understand the molecular mechanisms that regulate TCR-dependent signaling in CD8+ T cells, which together dictate the T cell function. We have recently demonstrated that Dlg1 plays a vital role in CD8+ T cell proinflammatory cytokine production and target cell killing in response to acute viral infection (O. Silva et al., submitted for publication). In this study, we show that TCR-induced Dlg1 phosphorylation is a key point of control capable of uncoupling cytokine production and cytotoxicity in CD8+ T cells. Given these data, it is interesting to imagine that TCR-induced phosphorylation of Dlg1AB acts as a molecular switch turning on p38 dependent functions, including proinflammatory cytokine production, in response to certain Ags, whereas other Ag signals may be sufficient to trigger degranulation.

Our data place Dlg1 Y222 phosphorylation as a vital regulatory step required for activation of p38 and NFAT. In the CD8+ T cells examined in this study, these pathways control the expression of proinflammatory cytokines IFN-γ and TNF-α. Dlg1-mediated p38 and NFAT activation also is thought to play an important role in the development of antibody responses by T cells.
role in regulating CD4+ T cell development and function. In CD4+ regulatory T cells, Dlg1 expression and localization to the immunocyte synapse correlates with p38 phosphorylation, NFAT-dependent transcription, and regulatory T cell-suppressive activity (16). In CD4+ effector cells, Dlg1 is thought to play a role in skewing toward expression of Th1 cytokines (36). In addition, Dlg1 also has been shown to influence the development of CD4+ T cell memory (44). Although both Dlg1AB and Dlg1B are expressed in CD4+ T cell subsets, the relative abundance of each isoform and the role of phosphorylation have yet to be explored (O. Silva et al., submitted for publication). Future studies into these cell types are likely to yield important insights into mechanisms of CD4+ T cell development and function.

In summary, Dlg1 is a key signaling and cytoskeletal specifier downstream of the TCR. Our study identifies Lck-mediated Dlg1 phosphorylation as a previously uncharacterized point of control regulating Dlg1-mediated signaling downstream of the TCR. In response to TCR stimulation, Lck phosphorylates the Dlg1 splice variant at several tyrosine residues, including Y222. Loss of Dlg1 phosphorylation at Y222 through Y222F mutation led to a significant and specific loss of p38 activation and up regulation of IFN-γ and TNF-α gene expression but did not impair cytotoxic granule release. A better understanding of the mechanisms that specifically regulate proinflammatory cytokines should ultimately lead to the discovery and design of more efficient and safe anti-inflammatory treatments for a variety of inflammatory conditions, including autoimmune disorders and severe influenza infection.

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Disclosures
The authors have no financial conflicts of interest.

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2. Rutishauser, R. L., and S. M. Kaech. 2010. Generating diversity: transcriptional activity (16). In CD4+ effector cells, Dlg1 is thought to play a role in regulating CD4+ T cell development and function. In CD4+ regulatory T cells, Dlg1 expression and localization to the immunocyte synapse correlates with p38 phosphorylation, NFAT-dependent transcription, and regulatory T cell-suppressive activity (16). In CD4+ effector cells, Dlg1 is thought to play a role in skewing toward expression of Th1 cytokines (36). In addition, Dlg1 also has been shown to influence the development of CD4+ T cell memory (44). Although both Dlg1AB and Dlg1B are expressed in CD4+ T cell subsets, the relative abundance of each isoform and the role of phosphorylation have yet to be explored (O. Silva et al., submitted for publication). Future studies into these cell types are likely to yield important insights into mechanisms of CD4+ T cell development and function.

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