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Spatial Coupling of JNK Activation to the B Cell Antigen Receptor by Tyrosine-Phosphorylated Ezrin

Neetha Parameswaran, Gospel Enyindah-Asonye, Nayer Bagheri,1 Neilay B. Shah,2 and Neetu Gupta

The ezrin-radixin-moesin proteins regulate B lymphocyte activation via their effect on BCR diffusion and microclustering. This relies on their ability to dynamically tether the plasma membrane with actin filaments that is in turn facilitated by phosphorylation of the conserved threonine residue in the actin-binding domain. In this study, we describe a novel function of ezrin in regulating JNK activation that is mediated by phosphorylation of a tyrosine (Y353) residue that is unconserved with moesin and radixin. BCR, but not CD40, TLR4, or CXCR5 stimulation, induced phosphorylation of ezrin at Y353 in mouse splenic B cells. Ezrin existed in a preformed complex with Syk in unstimulated B cells and underwent Syk-dependent phosphorylation upon anti-IgM stimulation. Y353-phosphorylated ezrin colocalized with the BCR within minutes of stimulation and cotrafficked with the endocytosed BCRs through the early and late endosomes. The T567 residue of ezrin was rephosphorylated in late endosomes and at the plasma membrane at later times of BCR stimulation. Expression of a nonphosphorylatable Y353F mutant of ezrin specifically impaired JNK activation. BCR crosslinking induced the association of Y353-phosphorylated ezrin with JNK and its kinase MAPKK7, as well as spatial colocalization with phosphorylated JNK in the endosomes. The yellow fluorescent protein–tagged Y353F mutant displayed reduced colocalization with the endocytosed BCR as compared with wild-type ezrin-yellow fluorescent protein. Taken together, our data identify a novel role for ezrin as a spatial adaptor that couples JNK signaling components to the BCR signalosome, thus facilitating JNK activation. The Journal of Immunology, 2013, 190: 2017–2026.

Antigen recognition by the BCR in mature B cells triggers a signaling cascade that culminates in transcriptional activation and proliferation (1, 2). At the outset, BCR signaling is accompanied by actin cytoskeletal reorganization that facilitates the formation of BCR microclusters, Ag gathering by the spreading B cell, and the assembly of BCR signalosomes (3, 4). This coordination between intracellular signaling molecules and the cytoskeleton modulates the strength of B cell activation (3, 5, 6). Clustering of the BCR signalosomes is also accompanied by rapid internalization and trafficking of the Ag-bound BCRs to the late endosomes for further processing of the Ag and loading on MHC class II molecules (7). The endocytosed BCRs in turn cosegregate with tyrosine and serine/threonine kinases within the endosomal compartments and continue to support signal transduction (8). It is very likely that cytoskeleton-regulating proteins influence the assembly of intracellular signaling components with the BCR in endosomal signalosomes and play an important role in regulating BCR signaling.

The cortical actin filaments are held underneath the plasma membrane by adaptor proteins that tether transmembrane proteins to actin. Ezrin, a plasma membrane–actin cytoskeleton crosslinking protein of the ezrin-radixin-moesin (ERM) family, contains a conserved threonine residue (T567) in its C-terminal actin-binding domain. Phosphorylation of this threonine is critical for conformational activation and plasma membrane–cytoskeleton crosslinking activity of ERM proteins (9). We previously reported that ezrin is constitutively phosphorylated at T567 in naive B cells, and dephosphorylation of this site upon BCR stimulation results in conformational inactivation facilitating lipid raft coalescence (10). Similarly, chemokine exposure induces T567 dephosphorylation in ezrin in B cells, and the resulting uncoupling of plasma membrane from the actin cytoskeleton is required for the morphological and cytoskeletal changes essential for B cell migration (11). Ezrin-rich networks confine BCR mobility in the absence of Ag (5), but they undergo dynamic remodeling upon Ag stimulation to facilitate Ag-receptor clustering (12). Therefore, Ag-induced conformational inactivation of ezrin is an important regulator of membrane dynamics during BCR signal transduction.

High structural homology between ezrin and moesin and their well-established role as membrane–cytoskeletal crosslinkers has led to the notion that the two proteins have redundant function in lymphocyte activation and migration (9). Indeed, ezrin-deficient mature T cells show defect in TCR-dependent IL-2 production, which is exacerbated upon additional knockdown of moesin expression (13). Interestingly, despite high overall homology between ezrin and moesin, the amino acid sequence of ezrin contains unique phosphorylation sites (S66, Y353, and Y477), and a polyproline stretch at 469–475 (14), suggesting that ezrin may have additional unconserved context-dependent roles. These features in ezrin may
enable protein–protein interactions to facilitate signal transduction and/or localization of interacting proteins. S66-phosphorylated ezrin regulates trafficking of H,K*-ATPase to the apical membrane of gastric parietal cells and is essential for histamine-induced acid secretion (15). Growth factor–dependent phosphorylation of ezrin at Y353 was shown to regulate survival of nonhematopoietic cells (16, 17), whereas Y477 phosphorylation of ezrin regulates growth and invasion of Src-transformed epithelial cells in a three-dimensional environment (18). In human B lymphoma cells, CD81 crosslinking was shown to induce Y353 phosphorylation of ezrin that recruits F-actin and facilitates cytoskeletal reorganization (19). Whether any of these alternate phosphorylation sites contribute toward BCR signaling has not been explored.

In this study, we report that BCR crosslinking induces phosphorylation of Syk-associated ezrin at Y353, Y353-phosphorylated ezrin associates with MAPKK7 (MKK7) and JNK and couples proximal BCR signaling with the JNK activation machinery in the endosomes, highlighting a novel role for ezrin as a spatial adaptor in the JNK activation pathway.

Materials and Methods

Mice and cells

C57BL/6 (B6) mice were bred in our animal colony, and Lyn−/− mice were obtained from Dr. Anthony DeFranco (University of California, San Francisco). Mice were used at 8–12 wk of age and all experiments were approved by the Institutional Animal Care and Use Committee of the Cleveland Clinic. The murine B lymphoma cell line CH27 was cultured in DMEM medium supplemented with 15% FBS. The DT40 chicken B cell lines were maintained in DMEM supplemented with 10% FBS and 5% DMEM medium supplemented with 15% FBS. The DT40 chicken B cell line was obtained from Dr. Arthur Weiss (University of California, San Francisco). Primary B cells were purified from spleens of B6 and Lyn−/− mice using CD43 microbeads (Miltenyi Biotec).

Plasmids and transfection

The ezrin-pERE52-eGFP and ezrin-yellow fluorescent protein (YFP) constructs have been previously described (11) and were used to mutate the tyrosine 353 residue to phenylalanine, resulting in the Y353F and Y353F-YFP constructs, respectively. Mutagenesis was performed using the QuickChange II site-directed mutagenesis kit (Stratagene). CH27 cells were transfected with 3–6 μg of the appropriate plasmids using Amaxa Nucleofector II (Lonza).

Cell lysis, immunoblotting, and immunoprecipitation

CH27 cells were transfected with vector alone, Y353F, ezrin-YFP, or Y353F-YFP and stimulated with 10 μg/ml anti-IgM at 37˚C for the indicated times. Lysates were prepared as previously described (11) and resolved by SDS-PAGE followed by immunoblotting with appropriate primary and secondary Abs. In some experiments, purified splenic B cells were pretreated with 50 μM PPI (Calbiochem) or indicated concentrations of piceatannol (Calbiochem) for 30 min at 37˚C prior to stimulation with anti-IgM. For immunoprecipitation, CH27 cell lysates were incubated with appropriate Abs and protein G–agarose beads (Invitrogen). Alternatively, lysates were incubated with GST-c-Jun fusion protein–conjugated Sepharose beads (Cell Signaling Technology) to precipitate JNK. Immunoblots were subjected to densitometric analysis using ImageJ software for quantification of ratios.

Flow cytometry assays

Surface IgM expression on the vector and Y353F transfectants of CH27 cells was compared by immunostaining with anti-β-gal-alkaline phosphatase (Becton Dickinson Biosciences) for 30 min at 37˚C. For BCR internalization assay vector and Y353F transfectants of CH27 cells were stained with biotin-conjugated anti-IgM for 30 min on ice, stimulated at 37˚C for the indicated times, fixed, and stained with streptavidin-conjugated PE (Becton Dickinson Biosciences) for 45 min at 4˚C. Samples were acquired using the BD FACSCalibur flow cytometer.

Calcium flux assay

For measurement of intracellular calcium, CH27 cells transfected with vector or Y353F were loaded with 1 μM Indo-1-AM (Invitrogen) and an equal volume of pluronic F-127 (Invitrogen) for 45 min at 37˚C. The cells were washed and resuspended in DMEM medium supplemented with 1% BSA and 20 nM HEPES. The cells were maintained in the dark at room temperature and a ratio of UV450/525 nm was record to establish the baseline fluorescence for unstimulated cells. Cells were stimulated at 30 s with 10 μg/ml anti-IgM, and the UV450/525 ratio was recorded for 200 s. Samples were acquired using a BD LSRII flow cytometer and data were analyzed using FlowJo (Tree Star).

Immunofluorescence microscopy

To visualize the BCR, CH27 cells were stimulated with biotin-conjugated anti-IgM and fixed with 4% PFA, blocked with PBS containing 15% goat serum for 30 min, followed by permeabilization and staining with Alexa Fluor 488/633–conjugated streptavidin (Molecular Probes) for 45 min at 4˚C. Next, the cells were stained with the pY353 Ab overnight at 4˚C and Alexa Fluor 568–conjugated anti-rabbit IgG (Molecular Probes) for 30 min. Puriﬁed spleen B cells from B6 mice were stimulated with anti-IgM, ﬁxed, permeabilized, and stained with phosphorylated threonine in ERK (pThrERK) Ab overnight and Alexa Fluor 568–conjugated anti-rabbit IgG for 30 min. Ezrin-YFP– or Y353F–YFP–transfected CH27 cells were stimulated and stained for the BCR as described above. For simultaneous imaging of pJNK and Y353-phosphorylated ezrin, cells were stimulated, ﬁxed, and stained with pY353 Ab as described above, followed by staining with mAb to pJNK and Alexa Fluor 488/647–conjugated anti-mouse IgG (Molecular Probes) for 30 min at 4˚C. To mark the endosomal aggregates, CH27 or splenic B cells were stained with Abs to early endosome Ag 1 (EEA1; Invitrogen) or lysosomal-associated membrane protein 1 (LAMP1; BD Pharamingen) for 1 h at 4˚C followed by Alexa Fluor 488–conjugated anti-chicken IgG (Molecular Probes) or Alexa Fluor 633–conjugated anti-rat IgG (Molecular Probes), respectively, for 45 min. The cells were stained with DAPI (Invitrogen), washed, and image slices were acquired through the z-axis with an interval of 0.2 μm. Images were acquired using a Leica AM TIRF microscope DMi6000 (Leica Microsystems) with an attached Hamamatsu EM-CCD camera, using HCX PL APO ×100 oil objective at an additional ×1.6 magnification with a numerical aperture of 1.47 and appropriate filter cubes. Imaging was performed using the Leica acquisition software LAS AP version 2.2.0. For Fig. 3, CH27 cells were stained with Abs to pY353, BCR, and DAPI as described above, imaged by confocal microscopy, and image slices were acquired through the z-axis with an interval of 0.1 μm. Images were acquired using a PerkinElmer UltraVIEW VoX confocal imaging system and a Leica-AM microscope DMi6000 SD (Leica Microsystems) with an attached Hamamatsu EM-CCD C9100-50 camera, using HCX PL APO ×100 oil objective with a numerical aperture of 1.47 and appropriate filter cubes. Images were acquired using the Velocity 5.5 software.

Image analysis

The images across the z-series were combined using the Velocity version 6.0.1 for three-dimensional volumetric rendition of z-stacks. The z-stacks of images were digitally deconvolved using the “iterative restoration” function in Velocity. Briefly, point spread functions were calculated for each fluorochrome channel and restoration was performed using 85–95% confidence limits and 25–30 iterations. For colocalization analysis, the deconvolved images were thresholded and Pearson’s correlation coefficients calculated for the specified pairs of channels across the stack of images using Velocity. For Fig. 7E, Metamorph image analysis software was used for digital “nearest-neighbors” deconvolution of the z-stacks. ImagePro 6.1 software was used to quantify pixel intensity in Fig. 7F. Briefly, z-stack images were analyzed using an algorithm written to quantify the number of green (pJNK) pixels in the region of interest. The images were filtered to enhance/ equalize their appearance, thresholded, and the total number of green pixels in each region of interest was calculated by summing the segmented pixels in the region of interest across the stack of images.

Statistical analyses

Exact p values were calculated using the nonparametric two-tailed Mann–Whitney U test except in Fig. 1A and Fig. 11B, where an unpaired t test was used. A p value <0.05 was considered statistically significant. The tests were performed with 95% conﬁdence interval (α level 0.05) using Prism (GraphPad Software). Means ± SD are reported along with p values.

Results

BCR-specific tyrosine phosphorylation of ezrin

We first tested whether B cell stimulation would lead to phosphorylation of ezrin at Y353. Puriﬁed spleen B cells from B6 mice were treated with anti-mouse IgM, anti-CD40, LPS, or CXCL13 at...
experiments. The data in (A) show the ratio of pY353 to actin (mean ± SD) from three independent experiments. *p < 0.05, ***p < 0.001.

37°C for indicated time periods. Out of the four stimuli tested only BCR ligation induced phosphorylation of Y353 in ezrin (Fig. 1A). In contrast, all the ligands induced dephosphorylation of the conserved T567 (Fig. 1B). Phosphorylation of Y353 upon BCR crosslinking indicated that ezrin might have a second role in BCR signaling in addition to its membrane–cytoskeleton tethering function that is known to regulate BCR diffusion and micro-clustering.

**Phosphorylation of ezrin at Y353 requires Syk tyrosine kinase activity**

As Lyn is the predominant Src family kinase (SFK) in B cells, we used B cells from B6 or Lyn−/− mice to determine whether it is required for Y353 phosphorylation in response to BCR cross-linking. Absence of Lyn kinase reduced the Ag-dependent tyrosine phosphorylation of ezrin (Fig. 2A). In contrast, pretreatment of splenic B cells with the SFK inhibitor PP1 abolished global tyrosine phosphorylation as well as phosphorylation of Y353 in ezrin (Supplemental Fig. 1A). These data suggest that SFK activity initiates the signaling events leading up to Y353 phosphorylation and that Lyn contributes to it but is not essential. SFK-mediated phosphorylation of ITAM in Igα and Igβ further initiates sequential activation of downstream tyrosine kinases such as Syk and Btk (2). As Lyn deficiency did not eliminate Y353 phosphorylation of ezrin, we tested whether Syk or Btk played a role in phosphorylation of ezrin at Y353. Treatment of mouse splenic B cells with a Syk kinase inhibitor, piceatannol, also resulted in a dose-dependent inhibition of Y353 phosphorylation (Supplemental Fig. 1B). As phosphorylation of Syk was reduced and not completely abrogated in the absence of Lyn (Supplemental Fig. 1C), the remaining Y353 phosphorylation of ezrin in Lyn-deficient B cells may be due to the residual Syk activity.

Phosphorylation of ezrin at Y353 and its subcellular localization in B cells

Protein kinases are known to interact with their substrates during the phosphorylation event. Therefore, we tested whether Syk interacts with ezrin to phosphorylate it at Y353. Ezrin coimmunoprecipitated with Syk from lysates of CH27 cells regardless of whether the cells were stimulated with anti-IgM or not (Fig. 2C). However, the association of Y353-phosphorylated ezrin with Syk was maximal at 1 min (Fig. 2C), which coincides with the peak of Y353 phosphorylation observed in B cell lysates (Fig. 1A). These results indicate that ezrin and Syk exist in a preformed complex that facilitates Syk-mediated phosphorylation at Y353 upon BCR stimulation.

**Tyrosine-phosphorylated ezrin colocalizes with the BCR**

The interaction between ezrin and Syk suggested that Y353-phosphorylated ezrin may be part of the BCR signalosome that assembles at the membrane and supports Ag-induced early signaling events. We employed confocal microscopy to determine the subcellular localization of Y353-phosphorylated ezrin with respect to the BCR in CH27 cells using Abs to pY353 and BCR. Unstimulated B cells showed punctate distribution of the BCRs at the plasma membrane and negligible signal for Y353-phosphorylated ezrin (Fig. 3A). Anti-IgM stimulation induced tyrosine phosphorylation of ezrin at 1 min, showing a scattered, punctate distribution with patches of colocalization with the BCR. At 10 min of anti-IgM stimulation the BCRs appeared focused and Y353-phosphorylated ezrin colocalized with the BCR. The BCR and Y353-phosphorylated ezrin localized to the perinuclear space and remained colocalized up to 30 min of stimulation. The perinuclear colocalization at later times of anti-IgM stimulation was observed in >80% of the B cells imaged (Fig. 3B).
BCR crosslinking induces receptor oligomerization and generation of signaling microclusters or “microsignalosomes.” The endocytosed Ag-bound BCRs traffic through early and late endosomes to deliver the Ag for proteolytic processing and transfer onto the MHC class II molecules (20). As Y353-phosphorylated ezrin colocalized with the BCR in the perinuclear region, we examined whether it traffics through the endosomal compartments together with the BCR. We imaged Y353-phosphorylated ezrin and BCR along with EEA1 and LAMP1, membrane proteins that mark early and late endosomes, respectively. CH27 cells were stimulated with anti-IgM and immunostained with Abs to pY353, BCR, and EEA1 (Fig. 4) or LAMP1 (Fig. 5). To clearly visualize the subcellular localization of Y353-phosphorylated ezrin and BCR we generated volumetric reconstructions of each cell using z-stacks of the images. Resting B cells showed punctate distribution of the BCRs, negligible tyrosine-phosphorylated ezrin, and scattered distribution of EEA1-positive early endosomes (Fig. 4A). Anti-IgM–induced Y353-phosphorylated ezrin appeared scattered at 5 min of stimulation and was colocalized in patches with the BCR-containing EEA1-positive early endosomes (Fig. 4). At 10 min of anti-IgM stimulation, BCR and Y353-phosphorylated ezrin were colocalized, with a few patches still associating with EEA1-positive endosomes (Fig. 4). BCR and Y353-phosphorylated ezrin remained colocalized at 15 min and 30 min of stimulation as seen in Fig. 3A but they were mostly outside the EEA1-positive endosomes at these times (Fig. 4). Imaging of the BCR, Y353-phosphorylated ezrin and the late endosomal protein, LAMP1, showed negligible association of BCR and Y353-phosphorylated ezrin with LAMP1 at 5 min of anti-IgM stimulation (Fig. 5A). Significant colocalization of the BCR and Y353-phosphorylated ezrin with LAMP1 was observed starting at 10 min that further increased at 30 min (Fig. 5). Taken together, these results show that within minutes of BCR crosslinking, Y353-phosphorylated ezrin and the BCR colocalized in EEA1-positive endosomes and remained colocalized in LAMP1-positive endosomes at later times of stimulation.

Recovery of threonine phosphorylation of ezrin occurs at the plasma membrane and in late endosomes

Anti-IgM–induced dephosphorylation of ezrin and moesin at the regulatory threonine residue is transient, as rephosphorylation is observed at 30 min of stimulation (Fig. 1B). At this time, Y353-phosphorylated ezrin is observed within late endosomes (Figs. 4, 5). To identify the intracellular compartment where ezrin becomes rephosphorylated at the threonine residue, we imaged pThrERM together with LAMP1 in B cells stimulated with anti-IgM over a longer time period. Rapid loss of pThrERM from the plasma membrane was observed at 3 and 10 min of stimulation (Fig. 6A, 6B). Most of the recovery of pThrERM
occurred at the plasma membrane starting at 20 min of anti-IgM stimulation (Fig 6A, 6B); however, significant colocalization of pThrERM was also detected with LAMP1 at 30 min of stimulation (Fig. 6A, 6C), indicating that a pool of ezrin located within late endosomes also undergoes threonine phosphorylation.

Expression of the Y353F mutant of ezrin specifically inhibits JNK activation

To investigate the function of Y353 phosphorylation of ezrin in BCR signaling we mutated Y353 to phenylalanine (Y353F) in the plasmid vector pIRES-2–enhanced GFP (pI2E). Bicistronic expression of enhanced GFP was used as a reporter of transfection efficiency. The expression of IgM was equivalent in CH27 cells transfected with vector or the Y353F mutant (Supplemental Fig. 2A). Because Y353-phosphorylated ezrin colocalized with the BCR in endosomes within 5 min of stimulation, we first tested whether expression of the Y353F mutant altered stimulation-induced BCR internalization in CH27 cells transiently transfected with the vector pI2E (referred to as vector) or pI2E-Y353F (referred to as Y353F). Anti-IgM–induced BCR internalization was unaltered in the presence of the Y353F mutant of ezrin (Supplemental Fig. 2B). Expression of the Y353F mutant also did not alter anti-IgM–induced calcium flux (Fig. 7A). Next, CH27 cells expressing the vector or Y353F were stimulated with anti-IgM for indicated times and lysates were probed with Abs to phosphorylated forms of key signaling intermediates in BCR activation. Because the Y353F mutant of ezrin is untagged, expression of the Y353F mutant was assessed by increase in intensity of the protein band corresponding to endogenous ezrin in the immunoblot (Fig. 7B, bottom panel). The phosphorylation of BLNK and PLCγ2, two proteins that participate early in the BCR signaling cascade, was unaltered in the presence of the Y353F mutant of ezrin (Fig. 7B). ERK1/2 phosphorylation also remained intact in CH27 cells expressing the Y353F mutant of ezrin (Fig. 7C), whereas anti-IgM–induced phosphorylation of both the p54 and p46 isoforms of JNK was markedly inhibited (Fig. 7D). A prominent nonspecific band of 43 kDa was observed below the p46 isoform of phosphorylated JNK (Fig. 7D), but its

FIGURE 5. Y353-phosphorylated ezrin and BCR continue to colocalize in late endosomes. (A) CH27 cells were stimulated with 10 μg/ml anti-IgM for the indicated time and stained for BCR (green), pY353 (red), LAMP1 (cyan), and nucleus (blue). Volumetric renditions of representative cells at each time of stimulation are shown, with arrows pointing to overall and pairwise colocalization between Y353-phosphorylated ezrin, BCR, and LAMP1. Scale bars, 10 μm. (B) Pearson’s colocalization coefficients for the indicated pairs. Each dot represents an individual cell and horizontal lines indicate the mean (n = 15–20 cells/group). Statistical significance is shown for the indicated stimulated groups in comparison with the unstimulated group. ***p < 0.001.

FIGURE 6. T567 rephosphorylation of ezrin occurs at the plasma membrane and in late endosomes. (A) CH27 cells were stimulated with 10 μg/ml anti-IgM for the indicated time and stained for pThrERM (red) and LAMP1 (green). Middle plane (left column) and volumetric renditions (right column) of representative cells at each time of stimulation are shown, with arrows pointing to colocalization between pThrERM and LAMP1. Scale bar, 1 μm. Data are representative of two independent experiments. (B) Pixel intensity of pThrERM at the plasma membrane. (C) Pearson’s colocalization coefficients for pThrERM and LAMP1. Each symbol in (B) and (C) represents an individual cell and horizontal lines indicate the mean (n = 15–20 cells/group). *p < 0.05, **p < 0.01, ***p < 0.001.
Intensity was similar between the vector and Y353F-expressing cells (Supplemental Fig. 2C). To confirm the effect of Y353F mutant on JNK activation, we employed immunofluorescence microscopy to image the active, phosphorylated form of JNK (pJNK) in vector and Y353F-expressing CH27 cells stimulated with anti-IgM. Consistent with the data in Fig. 7D, anti-IgM stimulation of CH27 cells transfected with vector alone or the Y353F mutant were stimulated with 10 μg/ml anti-IgM for the indicated time and lysates were probed with (B) pBLNK, BLNK, pPLCγ2, PLCγ2, and ezrin or (C) pERK and ERK, or (D) pJNK and JNK Abs. *, Nonspecific protein cross-reacting with the pJNK Ab in lysates from CH27 cells. The bar graphs show ratios of phosphoprotein to the respective total protein. Data in (B)–(D) are averaged from three independent experiments (mean ± SD). (E) CH27 cells transfected with vector or Y353F were stimulated with 10 μg/ml anti-IgM for the indicated time, stained for pJNK, and a z-stack of images was acquired. Scale bars, 5 μm. Representative cells from each stimulation time are shown. (F) Quantification of total green (pJNK) pixels; the horizontal bar indicates the mean (n = 42–64 cells/group). *p < 0.05, **p < 0.01, ***p < 0.001.

These data suggest that tyrosine phosphorylation of ezrin positively regulates JNK activation.

Y353-phosphorylated ezrin colocalizes with phosphorylated JNK within endosomal compartments

Intracellular MAPK signaling depends on their compartmentalization with the BCR within endosomes (8). Because Y353-phosphorylated ezrin and the BCR comigrated in endosomes, we considered the possibility that tyrosine-phosphorylated ezrin spatially localizes JNK in proximity to the BCR. To investigate
whether Y353-phosphorylated ezrin colocalizes with the active, phosphorylated form of JNK, we imaged Y353-phosphorylated ezrin and pJNK using immunofluorescence microscopy. CH27 cells were stimulated with anti-IgM and immunostained with Abs to pY353 and pJNK and the endosomal proteins EEA1 and LAMP1. Unstimulated B cells showed minimal signal for pJNK and Y353-phosphorylated ezrin (Fig. 8A). Anti-IgM stimulation resulted in Y353 phosphorylation of ezrin and JNK phosphorylation, and both showed a scattered distribution at 1 min of stimulation (Fig. 8A). The colocalization of Y353-phosphorylated ezrin and pJNK increased significantly at 10 and 30 min of anti-IgM stimulation (Fig. 8B), and the Y353-phosphorylated ezrin/pJNK complex began to appear more focused at 30 min (Fig. 8A). The colocalization of Y353-phosphorylated ezrin/pJNK complex with EEA1-positive early endosomes peaked at 10 min of stimulation and decreased by 30 min (Fig. 8). The Y353-phosphorylated ezrin/pJNK complexes started associating with the LAMP1-positive late endosomes at 10 min of stimulation and were maximally colocalized at 30 min (Fig. 9).

Ezrin physically associates with JNK and MKK7

JNK-interacting proteins (JIPs) are scaffolds that facilitate JNK signal transduction by physically interacting with JNK and its upstream kinases such as MLK and MKK7 to bring them into close proximity for serial phosphorylation (21, 22). JIPs are poorly expressed in B cells (23) and their contribution toward BCR-dependent JNK activation has not been explored. We considered the possibility that Y353-phosphorylated ezrin regulates JNK activation in a JIP-like manner. Therefore, we examined the interaction between Y353-phosphorylated ezrin and JNK in anti-IgM-stimulated B cells by precipitating JNK from CH27 cells stimulated with anti-IgM using GST-c-Jun fusion protein–conjugated Sepharose beads and by looking for the presence of coprecipitated Y353-phosphorylated ezrin. Y353-phosphorylated ezrin associated with JNK at 5 min of anti-IgM stimulation and the association peaked at 10 min (Fig. 10A). Next, we examined the association of ezrin with MKK7, one of the upstream kinases that directly phosphorylates JNK. Maximal association between tyrosine-phosphorylated ezrin and MKK7 was detected at 5 min of BCR stimulation (Fig. 10B).

Y353F mutant of ezrin shows reduced colocalization with the internalized BCR

To address the possibility that the nonphosphorylatable Y353F mutant of ezrin may display altered colocalization or cotrafficking with the BCR or phosphorylated JNK, we created a tagged version of the Y353F mutant in which YFP was attached to its C terminus (Y353F-YFP). This tagged fusion protein allowed us to distinguish between the ectopic Y353F mutant ezrin and the endogenous wild-type ezrin. As expected, the Y353F-YFP mutant did not undergo Y353 phosphorylation upon anti-IgM stimulation (Supplemental Fig. 2D). In view of the fact that phosphorylated JNK is signifi-
of intracellular signaling molecules and require morphological and cytoskeletal reorganization (24–27). Dephosphorylation of the regulatory T567 in ezrin by all the ligands tested in our study suggests a common and conserved mechanism of conformational inactivation that may facilitate membrane–cytoskeletal remodeling during proliferation and migration of B cells. Exclusive induction of Y353 phosphorylation of ezrin by BCR ligation indicated that Y353-phosphorylated ezrin has a specific function in Ag-dependent B cell activation. Previously, CD81 but not BCR or CD19 crosslinking was reported to induce Y353 phosphorylation in human B cell lines (19). BCR-mediated Y353 phosphorylation was not observed in the OCI-Ly8 DLBCL cell line in that study, possibly due to the lower dose of stimulus and longer stimulation time (30 min) (19). In our study, Y353 phosphorylation of ezrin was still detectable at 30 min of anti-IgM stimulation, but it was much lower than that observed at 1 min. We have previously shown that anti-IgM stimulation induces rapid but transient loss of T567 phosphorylation of ezrin that is recovered later (10). The repolymerization of ezrin at T567 in late endosomes indicates that it may reassociate with the actin cytoskeleton in this compartment as well and recycle back to the plasma membrane. Ezrin is known to facilitate recycling of the α1-adrenergic receptor (28) in an actin-dependent manner.

Activation of the Src family tyrosine kinases is the earliest event in the BCR crosslinking–mediated signaling cascade, and accordingly lack of Lyn lowered Y353 phosphorylation of ezrin. Furthermore, an absolute requirement of Syk kinase activity for Y353 phosphorylation of ezrin was observed in the Syk-deficient DT40 B cells. The residual Y353 phosphorylation observed in Lyn-deficient mouse B cells (Fig. 2A) and DT40 chicken B cell line (Fig. 2B) suggests that Pyn and/or Blk cooperate with Lyn to transduce the signal to Syk. Notably, CD81-dependent Y353 phosphorylation of ezrin was also shown to be Syk-dependent (19). Although CD40 and CXCR5 engagement synergize with BCR crosslinking to enhance Syk kinase activation, independently these stimuli induce negligible activation of Syk (27, 29). This correlates well with our observations of insignificant Y353 phosphorylation of ezrin in B cells stimulated with their ligands, anti-CD40 and CXCL13. The form of ezrin that associates with Syk during P-selectin glycoprotein ligand-1–induced transcriptional activation in T cells (30). The mode of association of ezrin with Syk in B cells remains to be tested. As the Y353 residue is not conserved in moesin, the ability of ezrin to regulate JNK activation may not be redundant with moesin.

Pervanadate stimulation of murine CD4+ T cells, but not TCR engagement alone, was reported to induce Y353 phosphorylation of ezrin but its function remains unidentified (13). Ezrin was shown to associate with and recruit the tyrosine kinase ZAP-70 to the immunological synapse in T cells (31). Both ZAP-70 and Syk belong to the Syk family of cytoplasmic nonreceptor tyrosine kinases that are involved in Ag receptor–mediated signaling in T and B cells, respectively. Given the functional similarities between ZAP-70 and Syk, and their role in Ag-mediated signaling, ZAP-70 may catalyze Y353 phosphorylation of ezrin at the immunological synapse in T cells. In a separate study, TCR ligation in human T cell lines was shown to induce tyrosine phosphorylation of ezrin in an Lck- and ZAP-70–dependent manner (32), but the site of tyrosine phosphorylation in ezrin was not identified.

Discussion

In this study we expand the role of ezrin in B cells beyond its established role as a linker protein between the plasma membrane and actin cytoskeleton. In addition to dephosphorylation of T567, ezrin undergoes Syk-dependent phosphorylation at Y353 upon BCR crosslinking. BCR stimulation–induced increases in association of Y353-phosphorylated ezrin with JNK and its kinase MKK7, as well as its increased colocalization with the BCR and phosphorylated JNK within endosomal compartments, point toward an adaptor function for Y353-phosphorylated ezrin in JNK activation.

Stimulation of naive B cells with ligands that engage different receptors on B cells induces various functional outcomes, including proliferation, survival, and migration. It is known that B cell proliferation downstream of BCR, TLR4, or CD40 ligation, as well as B cell migration, induced by CXCL13 involve polarization

![FIGURE 10. Ezrin physically associates with JNK activation components. CH27 cells were stimulated with 10 μg/ml anti-IgM for the indicated time, and lysates were subjected to precipitation with (A) GST-c-Jun fusion protein–Sepharose beads or (B) MKK7 Ab and protein G-agarose beads. The precipitates were probed with Abs to pY353 and (A) JNK or (B) MKK7 Ab. Data in each panel are representative of two independent experiments.](http://www.jimmunol.org/content/204/7/3455/F10.large.jpg)
BCR “microsignalosomes” assemble at the surface of B cells and include intracellular signaling molecules such as Syk, PLCγ2, and Vav (33). As BCR signaling proceeds, intracellular proteins sequentially enter and exit the BCR signaling complex. Continual signal transduction by the internalized BCRs within endosomes underscores the importance of compartmentalization of signaling components in optimal regulation of the MAPK pathways (8). Preventing BCR endocytosis and its subsequent sequential colocalization with kinases within the endosomes was shown to alter the extent and outcome of MAPK activation (8). Recent studies have identified MAPK scaffold proteins (e.g., MP1 and β-arrestin) that modulate stimulus-dependent activation of ERK and JNK signaling pathways by spatially localizing signaling modules to endosomes (34, 35). Interestingly, the prominent perinuclear colocalization of Y353-phosphorylated ezrin with the BCR and pJNK at later times of B cell stimulation resembles the localization reported for JNK scaffolds in fibroblasts and neuronal cells (36). The major groups of JNK scaffolds are the JIP proteins, of which JIP1 and JIP2 are almost exclusively expressed in neuronal cells (23, 37). Follicular B cells express very low levels of JIP3 and JIP4, of which JIP4 was shown to facilitate the activation of p38 instead of JNK (23, 38). Our data show association of ezrin with JNK and MKK7, indicating that ezrin may substitute for JIPs and act as a JNK scaffold in B cells. Our immunoprecipitation data showed that Y353-phosphorylated ezrin and JNK associated maximally at 10 min and dissociated at later times of stimulation. However, we continued to observe the colocalization of Y353-phosphorylated ezrin and pJNK within late endosomes at 30 min, suggesting that the physical interaction between Y353-phosphorylated ezrin and JNK or MKK7 may be a transient requirement and serve to spatially localize these kinases within the BCR-occupied signaling endosomes. Y353-phosphorylated ezrin may either associate directly with MKK7 and JNK, and function as a JIP-like adaptor, or facilitate the function of another as yet unidentified JNK scaffold protein. Our data using the YFP-tagged fusion proteins indicate that BCR internalization was unaffected in the presence of Y353F-YFP. Interestingly, the region around the internalized BCR was found to be rich in ezrin-YFP but devoid of Y353F-YFP, which indicates the importance of Y353 phosphorylation in recruiting ezrin to the endosomal compartment. Our biochemical and microscopy analyses show impaired JNK activation upon expression of the Y353F mutant of ezrin confirming the requirement of Y353 phosphorylation of ezrin. Our data also suggest that the Y353F mutant acts in a dominant negative manner, and further characterization of its protein–protein interactions will reveal the mechanism by which it interferes with the ability of endogenous ezrin to facilitate JNK activation.

Y353-phosphorylated ezrin has been reported to regulate the survival of nonhematopoietic cells (39). Expression of the Y353F mutant of ezrin in kidney epithelial cells led to increased apoptosis that was attributed to decrease in association of ezrin with the p85 subunit of PI3K and impaired activation of Akt downstream of PI3K (39). Ezrin is highly expressed in HS2 leukemic stage cell lines of proerythroblastic cells, and overexpression of the Y353F mutant of ezrin leads to increased cell death, supporting a role for Y353-phosphorylated ezrin in enhancing cell survival (40). These studies and ours suggest that ezrin can partner with different signaling proteins depending on the cellular context and stimulus.

Phosphorylation of ezrin recovers within late endosomes at 30 min and at the plasma membrane between 20 and 45 min of BCR stimulation. *p < 0.05. **p < 0.01.
Finally, our data suggest a model wherein BCR crosslinking induces activation of Syk leading to Y353 phosphorylation of ezrin (Fig. 11C). Y353-phosphorylated ezrin accesses and associates with JNK and MKK7 and couples proximal BCR signaling with the JNK signaling module within endosomes (Fig. 11C). As the BCR continues to transduce signals in the endosomal compartment it supports JNK activation. Moreover, ezrin undergoes rhesphorylation at TS67 in both the LAMP compartment and at the plasma membrane. Therefore, we propose that Y353-phosphorylated ezrin functions as an adaptor to recruit JNK and MKK7 spatially close to the active BCR within endosomes. In conclusion, we have identified Y353-phosphorylated ezrin as a novel regulator of JNK signaling in Ag-activated B cells.

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Disclosures
The authors have no financial conflicts of interest.

References
SUPPLEMENTARY FIGURE LEGENDS

Supplementary Figure S1. Effect of Src family and Syk kinase inhibitor on Y353 phosphorylation. (A) Splenic B cells were pretreated with 50 µM of the SFK inhibitor PP1, stimulated with 10 µg/ml of anti-IgM for 1 min and lysates were probed with antibodies to pY353, ezrin and total phosphotyrosine. (B) Splenic B cells were pretreated with indicated doses of the Syk inhibitor piceatannol, stimulated with 10 µg/ml of anti-IgM for 1 min and lysates were probed with pY353 and ezrin antibody. The bar graph shows the ratio of pY353 to ezrin. (C) Purified B cells from B6 or Lyn-deficient mice were stimulated with 10 µg/ml of anti-IgM for the indicated time and lysates probed with pSyk and Syk antibody.

Supplementary Figure S2. Surface BCR levels in resting and stimulated Y353F expressing CH27 cells, and tyrosine phosphorylation of YFP fusion constructs of ezrin. (A) CH27 cells were transfected with the vector pI2E or the Y353F mutant of ezrin and analyzed by flow cytometry for surface IgM levels on GFP+ cells. (B) CH27 cells transfected with the vector or the Y353F mutant were stimulated with biotin-conjugated anti-IgM, fixed and stained with phycoerythrin-conjugated streptavidin. Surface BCR levels on GFP+ cells are shown. (C) Ratio of the non-specific band (*) in the pJNK blot shown in Fig. 7D to the total p46 isoform JNK. (D) CH27 cells transfected with the Ezrin-YFP or Y353F-YFP fusion constructs of ezrin were stimulated with 10 µg/ml of anti-IgM for the indicated time and lysates were probed with pY353 and ezrin antibodies.