Involvement of Guanosine Triphosphatases and Phospholipase C-γ2 in Extracellular Signal-regulated Kinase, c-Jun NH2-terminal Kinase, and p38 Mitogen-activated Protein Kinase Activation by the B Cell Antigen Receptor

By Ari Hashimoto,* Hidetaka Okada,* Aimin Jiang,† Mari Kurosaki,* Steven Greenberg,§ Edward A. Clark,‡ and Tomohiro Kurosaki*

From the *Department of Molecular Genetics, Institute for Liver Research, Kansai Medical University, Moriguchi 570-8506, Japan; the ‡Department of Microbiology, University of Washington, Seattle, Washington 98195; and the §Department of Medicine and Pharmacology, Columbia University College of Physicians and Surgeons, New York 10032

Summary

Mitogen-activated protein (MAP) kinase family members, including extracellular signal-regulated kinase (ERK), c-Jun NH2-terminal kinase (JNK), and p38 MAP kinase, have been implicated in coupling the B cell antigen receptor (BCR) to transcriptional responses. However, the mechanisms that lead to the activation of these MAP kinase family members have been poorly elucidated. Here we demonstrate that the BCR-induced ERK activation is reduced by loss of Grb2 or expression of a dominant-negative form of Ras, RasN17, whereas this response is not affected by loss of Shc. The inhibition of the ERK response was also observed in phospholipase C (PLC)-γ2-deficient DT40 B cells, and expression of RasN17 in the PLC-γ2-deficient cells completely abrogated the ERK activation. The PLC-γ2 dependency of ERK activation was most likely due to protein kinase C (PKC) activation rather than calcium mobilization, since loss of inositol 1,4,5-trisphosphate receptors did not affect ERK activation. Similar to cooperation of Ras with PKC activation in ERK response, both PLC-γ2-dependent signal and GTP-ase are required for BCR-induced JNK and p38 responses. JNK response is dependent on Rac1 and calcium mobilization, whereas p38 response requires Rac1 and PKC activation.

Key words: mitogen-activated protein kinase family • Ras • Rac1 • protein kinase C • calcium

The B cell antigen receptor (BCR)1 complex has important functions in the binding and internalization of antigen as well as in transducing signals through the plasma membrane that lead to cell activation, proliferation, and apoptosis. One immediate consequence of triggering the BCR is the activation of protein tyrosine kinases and the resulting induction of phospholipase C (PLC)-γ2-mediated hydrolysis of inositol phospholipids to generate diacylglycerol and inositol 1,4,5-trisphosphate (IP3), which induce protein kinase C (PKC) activation and elevate intracellular calcium ([Ca2+]i), respectively (1–4). Supporting these sequential events, PLC-γ2-deficient DT40 B cells exhibit neither PKC activation nor calcium mobilization after BCR ligation (5), whereas the PKC activation still occurs in IP3 receptor (IP3R)-deficient DT40 cells (6). Thus, combined analyses using PLC-γ2- and IP3R-deficient DT40 cells have allowed us to dissect the requirement of PKC and calcium mobilization in BCR-mediated signaling pathways.

BCR stimulation also leads to extracellular signal-regulated kinase (ERK) activation, which in turn activates transcription factors (7–11). Analogous to receptor tyrosine kinases, the interaction of Grb2, Shc, and Sos, which have been implicated in the activation of Ras, is thought to be a potential pathway for the activation of ERK after BCR ligation (12–14). In addition to this potential pathway, there is evidence that the ERK cascade is activated by treatment with PMA and ionomycin in cultured 3T3 fibroblasts (15). However, it is not known whether BCR-induced ERK activation requires PKC activation and/or [Ca2+]i increase. Moreover, there is no direct evidence for the involvement of Grb2, Shc, and Ras in BCR-induced ERK activation.

In addition to ERK, other mitogen-activated protein

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1Abbreviations used in this paper: BCR, B cell antigen receptor; EGF, epidermal growth factor; ERK, extracellular signal-regulated kinase; GST, glutathione S-transferase; IP3, inositol 1,4,5-trisphosphate; JNK, c-Jun NH2-terminal kinase; MAP, mitogen-activated protein; PKC, protein kinase C; PLC, phospholipase C.

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(MAP) kinase family members c-Jun N \( \text{H}_2 \)-terminal kinase (JNK) and p38 are known to be stimulated by BCR cross-linking, although less effectively than is ERK (11, 16, 17). In a fibroblast cotransfection system, Rac1 has been shown to stimulate JNK and p38 (18–20). Furthermore, recent experiments have demonstrated that Vav, upon tyrosine phosphorylation, can activate members of the Rho family of GTPases (21–23). Although these observations provide the intriguing possibility that the activated Rac1 by Vav leads to activation of JNK and p38, any link between BCR-regulated Rac1 and the activation of JNK and p38 has not yet been explored.

Different activation patterns of these MAP kinase family members (ERK, JNK, p38) may lead to differential expression of genes and contrasting cellular phenotypes, such as growth or apoptosis in B cells. This concept is supported by the evidence obtained from the cultured B cell system and the anti-IgM-induced apoptosis (16). Moreover, in transgenic mice, both ERK and JNK are activated by foreign antigens in naive B cells, whereas ERK, but not JNK, is activated by self-antigens in tolerant B cells (24).

In this study, we focus upon how BCR activates ERK, JNK, and p38. Rac1 inhibited the BCR-induced ERK activation without affecting JNK and p38 responses. Conversely, the JNK and p38 responses were abrogated by expression of a dominant-negative form of Rac1 and Rac1N17, but not of Rac1N17. In addition to demonstrating requirement of these GTPases, our results reveal the importance of the PLC–γ2 pathway in activation of these MAP kinase family members.

Materials and Methods

Cells, Expression Constructs, and Antibodies. Wild-type and various mutant DT 40 cells were cultured in RPMI 1640 supplemented with 10% FCS, 1% chicken serum, 50 \( \mu \)g/ml 2-ME, 2 mM L-glutamine, and antibiotics. The cDNA of Rac1N17 (provided by T. Ueno, Department of Biochemistry, Nagoya University School of Medicine, Nagoya, Japan) and Myc-tagged Rac1N17 (reference 25; provided by G. Bokoch, Department of Immunology and Cell Biology, Scripps Research Institute, La Jolla, CA) were cloned into expression vector pA3uro (26). These cDNAs were transfected by electroporation at 550 V, 25 \( \mu \)F, and selected in the presence of 0.5 \( \mu \)g/ml puromycin (Sigma Chemical Co., St. Louis, MO). Expression of transfected cDNA was confirmed by Western blot analysis. The mAb used for the stimulation of BCR was M4, which recognizes chicken IgM (27). Anti-Rac2 Ab was obtained by immunizing rabbits with bacterially expressed glutathione S-transferase (GST) fusion protein containing chicken BCR SH2 domain. The following Abs were purchased: anti-ERK2 and anti-p38 from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA); anti-JNK1 and anti-Myc from Pharmingen (San Diego, CA); anti-Grb2 Ab from Transduction Labs, Inc. (Lexington, KY); anti-Shc from Upstate Biotechnology, Inc. (Lake Placid, NY); and anti-Ras from Oncogene (Cambridge, MA).

Generation of Various Deficient DT 40 Cells. Based on published sequence of Grap (28) and Shc (29), each chicken cDNA was cloned by reverse transcriptase-PCR method using RNA from DT 40 B cells. Chicken Grb2 cDNA sequence has already been published (30). Genomic clones of Grb2, Grap, and Shc were obtained by PCR using oligonucleotides designed from each cDNA sequence and genomic DNA as a template. The targeting vectors, pGrb2-neo and pGrb2-hisD, were constructed by replacing the genomic fragment containing exons that correspond to chicken Grb2 SH2 domain, amino acid residues 59–166 (30), with neo and hisD cassettes. These cassettes were flanked by 2.3 and 1.8 kb of Grb2 sequence on the 5′ and 3′ sides, respectively. Selection was done in the presence of G418 (2 mg/ml) and clones were screened by Southern blot analysis. The pGrb2-hisD was again transfected into the neo-targeted clone and selected with both G418 (2 mg/ml) and histidinol (1 mg/ml). Introduction of a single copy of each transfecting vector was verified by reproping the blots with internal neo or hisD probe. The bsr and bleo targeting constructs of Grap were made by replacing the genomic fragment containing exons corresponding to human Grap SH2 domain, amino acid residues 59–160 (28), with bsr and bleo cassettes. These constructs spanned 0.8 (5′ side) and 1.2 kb (3′ side) of Grap sequence. They were sequentially transfected into DT 40 cells by electroporation to obtain a null mutant. Selection for drug-resistant clones was carried out by using blasticidin S (50 \( \mu \)g/ml) and phleomycin (0.3 mg/ml). The constructs (pShc-neo and pShc-hisD) were designed for neo and hisD cassettes to replace exons that correspond to human Shc amino acid residues 101–140 (29). The neo and hisD cassettes were flanked by 0.9 and 1.3 kb of Shc sequence on the 5′ and 3′ sides, respectively. Selection was performed as described above for Grb2. In this study, we also used DT 40 cells deficient in PLC–γ2 and all three isoforms of IP3R, all of which have been described previously (5, 6).

Northern Blot Analysis. RNA was prepared from wild-type and mutant DT 40 cells using the guanidium thiocyanate method. Total RNA (20 \( \mu \)g) was separated in a 1.2% formaldehyde gel, transferred to Hybond-N (Amer sham, Arlington Heights, IL), and probed with 32P-labeled cDNAs.

Western Blot Analysis. Wild-type, Grb2-, Grap-, and Shc-deficient cell lysates were separated by SDS-PAGE gel, transferred to nitrocellulose membrane, and detected by appropriate Abs and the ECL system (Amer sham, Arlington Heights, IL).

In vitro Kinase Assay. After incubation with the indicated stimuli, 2.5 × 106 cells were lysed in 500 \( \mu \)l of lysis buffer (50 mM Tris-Cl, pH 7.5, 150 mM NaCl, 5 mM EDTA, 2% Triton X-100, 100 \( \mu \)M sodium vanadate, 10 mM sodium pyrophosphate, 2 mM PM SF, 10 \( \mu \)g/ml leupeptin, and 2 \( \mu \)g/ml aprotinin) for 30 min on ice. Cell debris were removed by centrifugation at 16,000 g for 15 min at 4°C. Precleared lysates were immunoprecipitated by 1 \( \mu \)g anti-ERK2 Ab, 1 \( \mu \)g anti-JNK1 Ab, or 1 \( \mu \)g anti-p38 Ab with 40 \( \mu \)l protein G-Sepharose (Amersham Pharmacia Biotech, Piscataway, N J). The beads were pelleted by centrifugation and then washed three times with lysis buffer, two times with washing buffer (50 mM Hepes, pH 7.4, 10 mM MgCl2, 100 \( \mu \)M sodium vanadate, and 2 mM dithiothreitol). Immunoprecipitates were divided and half of them were used for Western blot analysis. The remaining half were washed once with kinase assay buffer (20 mM Hepes, pH 7.4, 10 mM MgCl2, 10 mM MnCl2, 2 mM dithiothreitol), and 10 \( \mu \)M sodium vanadate). Immune complexes were suspended in 30 \( \mu \)l kinase assay buffer containing γ-\( ^{32} \)PATP (>3,000 Ci/mmol; NEN Life Science Products, Boston, MA) and 5 \( \mu \)M cold ATP. A 5 \( \mu \)g of GST–Elk, GST–c-Jun (provided by M. Hibi, Biomedical Research Center, Osaka University Medical School, Suita, Japan) and GST–ATF2 fusion protein were added as substrates for ERK2, JNK, and p38, respectively. After 20 min

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of incubation at 30°C, the reaction was terminated by the addition of SDS sample buffer followed by boiling for 5 min. The samples were separated by SDS-PAGE gel, dried, and subjected to autoradiography.

Results

Requirement for Grb2, not Shc, in BCR-mediated ERK Activation. To determine the necessity or redundancy of Grb2 and Shc in the BCR-induced ERK activation, we generated DT40 B cells deficient in these molecules by the gene targeting method. Since Grap has recently been shown to possess similar structural characteristics to Grb2 and this protein is prominently expressed in lymphoid cells (28, 31), we also established Grap-deficient DT40 cells. Lack of Grb2, Grap, and Shc was verified by Northern and Western blot analyses (Fig. 1, A and B). The level of cell surface expression of BCR on these targeted DT40 cells was the same as that of parental DT40 cells (data not shown).

RNA analysis by reverse transcriptase-PCR method and Western blot analysis (32) indicated that ERK2 is the predominant isoform expressed in DT40 B cells. After BCR stimulation, DT40 cells were lysed and immunoprecipitated with anti-ERK2 Ab. The immunoprecipitates were assayed for its in vitro kinase activity by the ability to phosphorylate the substrate Elk-1. This anti-ERK2 Ab does not recognize JNK or p38 (Fig. 2). Consistent with previous reports (9, 10, 33, 34), activation of ERK2 in DT40 cells was maximal at 1 and 3 min, after which activity declined to a lower level (Figs. 3 and 4). Compared with wild-type cells, the BCR-induced ERK2 response was reduced approximately twofold by loss of Grb2. However, the ERK2 response was not reduced in Shc-deficient DT40 cells (Fig. 3), demonstrating that Grb2, but not Shc, is required for the BCR-induced ERK2 activation. Moreover, despite the structural similarity between Grb2 and Grap, Grap-deficient DT40 B cells exhibited no decrease in BCR-mediated ERK2 activation (Fig. 3), implicating that Grb2 and Grap have distinct function(s) in BCR signaling.

Since Grb2 is essential for Ras activation in receptor-type tyrosine kinase signaling, we decided to test whether Grb2 is involved in the ERK2 response along a Ras-dependent pathway in BCR signaling. As shown in Fig. 5, expression of RasN17 (Fig. 1 C) inhibited ERK2 activation by threefold compared with wild-type cells, suggesting that Ras as activation, perhaps via Grb2, is required for BCR-induced ERK2 activation. The residual ERK2 response by loss of Grb2 or expression of RasN17 implicates that an additional signal(s) is required for the maximum ERK2 activation and that this additional signal is able to stimulate ERK2 to some extent even in the absence of Ras as activation.

Both PKC and Ras Pathways Are Required for ERK Activation. Recent work has implicated PKC and/or [Ca^{2+}]_{i} increase as a cofactor for ERK activation (7, 15). To determine whether PKC and [Ca^{2+}]_{i} increase are required for BCR-induced ERK activation, we used PLC-γ2- and IP_{3}R-deficient DT40 cells. We have previously shown that the BCR-induced [Ca^{2+}]_{i} increase is completely abrogated in both PLC-γ2- and IP_{3}R-deficient DT40 cells, whereas PKC activation is abolished only in PLC-γ2-deficient cells (6). BCR ligation stimulated ERK2 activation even in the IP_{3}R-deficient cells, whereas this activation was markedly inhibited by loss of PLC-γ2 (Fig. 3). These data suggest that the PKC pathway rather than the calcium pathway is required for the BCR-induced ERK2 activation. Furthermore, the extent of the inhibition by loss of PLC-γ2 was more than that in DT40 cells expressing RasN17 (Fig. 5), suggesting that the PKC pathway may play a more dominant role than Ras as in BCR-mediated ERK2 activation.

The residual ERK2 activation in PLC-γ2-deficient DT40 cells was completely abolished by expression of
RasN17 in this mutant DT40 cell (Fig. 5). Together, these data suggest that Ras activation and the PKC-dependent pathway may work synergistically for the maximum BCR-induced ERK2 activation.

Requirement for Rac1 in BCR-induced JNK and p38 Activation. After BCR stimulation, the activities of JNK and p38 were determined by their ability to phosphorylate the substrates c-Jun and ATF-2, respectively. The kinetics of JNK activation were distinct from that of ERK2, being marked at 10 min and being declined by 30 min. The activation of p38 was maximal at 3 and 10 min, and more sustained than that of JNK (Figs. 4 and 6). Contrary to ERK2 activation, expression of RasN17 did not significantly affect the JNK and p38 responses upon BCR engagement (Fig. 6). Consistent with these observations, Grb2-deficient DT40 cells also showed normal JNK and p38 responses (data not shown). These findings prompted us to examine the involvement of Rac1 in BCR-induced JNK and p38 activation, since recent experiments in fibroblasts have shown that Rac1, rather than Ras, is an efficient activator of a cascade leading to JNK and p38 activation (18-20). Expression of Rac1N17 in DT40 cells (Fig. 1C) markedly inhibited both JNK and p38 activation after BCR cross-linking (Fig. 6), whereas ERK2 activation was not affected (data not shown). These results indicate that Rac1, rather than Ras, is required for JNK and p38 activation after BCR ligation.

PLC-γ2-dependent Signaling Is Required for JNK and p38 Activation. A calcium ionophore has been shown to activate JNK in B cells (16, 35). This suggests that other activators of JNK, such as ligated BCR, may use [Ca²⁺]i increase for maximum JNK activation. Likewise, p38 response also might be dependent on the [Ca²⁺]i increase in B cells. To test these possibilities, we examined the BCR-induced activation of JNK and p38 in PLC-γ2- and IP3R-deficient DT40 cells. As shown in Fig. 7, both PLC-γ2- and IP3R-deficient DT40 cells exhibited impaired JNK responses, suggesting the dependency of the JNK response on the calcium pathway rather than the PKC pathway. However, a residual JNK response was still observed even in the IP3R-deficient DT40 cells. Activation of p38 upon BCR engagement was completely abolished by loss of PLC-γ2, whereas this response was still observed in IP3R-deficient DT40 cells (Fig. 7). These results suggest that the BCR-induced p38 response requires a dominantly PKC pathway.

Discussion

In this study, we have addressed three questions regarding the regulation of ERK, JNK, and p38 by BCR, potentially leading to altered transcription of genes. First, are dis-
distinct GTPases required for the BCR-induced ERK, JNK, and p38 responses? Second, particularly in the ERK response, is either Grb2 or Shc, or both of them, essential for this response? Third, are GTPases the sole regulators of BCR-induced activation of these MAP kinase family members?

Since the two SH3 domains of Grb2 bind to proline-rich residues near the COOH terminus of Sos, Grb2 is thought to mediate the translocation of Sos to the plasma membrane, allowing Sos to activate membrane-bound Ras. In the case of the epidermal growth factor (EGF) receptor, translocation of Grb2 to membrane can occur upon binding of its SH2 domain to the autophosphorylated tail of the EGF receptor (36–40). However, because the BCR complex does not possess Grb2 binding sites despite activating Ras, an alternate model for Ras activation has been proposed. According to this model, BCR ligation leads to the recruitment of Shc to the plasma membrane, whereupon it undergoes tyrosine phosphorylation of the Grb2 binding site, thereby recruiting Grb2 to the plasma membrane (38). Indeed, BCR stimulation leads to tyrosine phosphorylation of Shc and to the assembly of an Shc–Grb2–Sos complex (8, 41–43). However, the data presented here indicate that Grb2 but not Shc is required for BCR-induced ERK response. These findings indicate the dispensability of Shc in Ras activation in BCR signaling, and raise the previous question of how Grb2 is recruited to the plasma membrane after BCR ligation. One candidate adaptor molecule might be LAT, an integral membrane protein that is phosphorylated and bound to Grb2 (44). Although LAT is expressed predominantly in T cells, B cells might express a similar molecule that is required for Grb2 recruitment to the membrane. This candidate molecule is tyrosine phosphorylated presumably by Syk, since the accompanying study has shown that Syk, but not Lyn, is a responsible kinase for BCR-induced ERK activation (32).

The extent of inhibition by RasN17 of BCR-mediated ERK2 activation was almost the same level as that observed in Grb2-deficient DT40 cells, suggesting that the Grb2-deficient phenotype in BCR signaling can be accounted for by loss of Ras as activation. In contrast to the almost complete inhibition of EGF receptor-induced ERK response by expression of RasN17 (45), we observed residual BCR-induced ERK2 activation despite loss of Grb2 or expression of RasN17. These observations provide important implications. First, the sensitivity to requirement of Ras as for ERK activation might vary depending on receptor systems. In this regard, TCR-induced ERK response appears to be more strictly dependent on Ras as activation than is BCR (46). Second, the BCR uses an additional signaling pathway to stimulate ERK even in the absence of Ras as, and probably this additional signaling pathway, in addition to Ras as, is required for maximum BCR-induced ERK activation. This additional signal is provided by PKC activation, since PLC-γ2-deficient DT40 cells, but not IP3R-deficient, exhibited profound loss in ERK2 activation by BCR ligation. Furthermore, expression of RasN17 in PLC-γ2-deficient cells led to the complete abrogation of BCR-induced ERK2 activation, suggesting the synergistic action of Ras as and the PKC pathway for BCR-induced ERK acti-
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In contrast to the ERK response, the JNK and p38 responses are not affected by expression of RasN17. Conversely, expression of RacN17 abolished the JNK and p38 activation without affecting the ERK response in B cells. This difference presumably reflects that distinct exchange factors, Sos and Vav, participate in ERK and JNK/p38 responses through activating Ras and Rac1, respectively, in BCR signaling. Supporting this notion, recent experiments have shown that Vav, upon tyrosine phosphorylation, stimulates Rac1 (21–23). Our results reveal that not only Rac1 but also PLC-γ2-dependent signal is required for BCR-induced JNK/p38 response; JNK is dependent on calcium mobilization rather than PKC activation, whereas p38 requires the PKC pathway. Requirement of calcium for JNK activation agrees with the data that JNK response, but not p38 response, is inhibited by treatment of BAPTA-AM, a chelator of both intracellular and extracellular calcium (32). The requirement of both Rac1 and PLC-γ for JNK activation is well conserved between B and T cells. Indeed, the data presented in this study and the accompanying paper (32) are consistent with the findings that JNK activation in T cells requires calcium-dependent calcineurin and Syk, in addition to Ras (49, 50).

Although this study has not addressed the mechanism by which PKC and calcium regulate p38 and JNK activation by BCR, modulation of the signaling pathway upstream of Rac1 by PKC and calcium seems to be unlikely, as Vav behaves as a tyrosine phosphorylation-dependent exchange factor for Rac1 (21–23). Rather, distinct requirement for PKC and calcium for p38 and JNK, respectively, might suggest that downstream molecules of Rac1 are potential targets of PKC and calcium. Analogous to Rac, which is recruited by Ras (51), and which is a substrate of PKC (52), another kinase participating in the cascade leading to p38 and JNK response might be a substrate of PKC and a calcium-dependent kinase, respectively.

Regulation of ERK, JNK, and p38 by PKC and calcium might provide insights into the mechanism by which CD19 costimulates the activation of these MAP kinase family members by BCR. Although coclustering CD19 and BCR leads to enhanced activation of ERK, the CD19-mediated enhancement of JNK and p38 responses is more dramatic (53). Since CD19 associates with Vav and phosphatidylinositol 3 kinase (54–57), coligation of CD19 with BCR brings these molecules into the close proximity of the BCR signaling complex. This may lead to further enhancement of tyrosine phosphorylation of Vav by Lyn and Syk, leading to its enhanced activation towards Rac1. Recently, phosphatidylinositol 3 kinase has been reported to participate in the enhanced calcium mobilization (58–60), which potentially could lead to augmented JNK activation. Conversely, inhibitory receptors such as PIR-B and FcγRIIB on B cells are predicted to inhibit these MAP kinase family members, as these receptors downregulate the BCR-induced PLC-γ2 activation and calcium mobilization (61–63).

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Figure 7. BCR-induced JNK and p38 responses in PLC-γ2− and IP3R− deficient DT40 cells. Cells were stimulated with M4 (4 μg/ml) for indicated times and cell lysates were immunoprecipitated with anti-JNK1 and anti-p38 Ab. The kinase assays were done as in Fig. 6.
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