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RIBP, a Novel Rlk/Txk- and Itk-binding Adaptor Protein That Regulates T Cell Activation

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Summary

A novel T cell–specific adaptor protein, RIBP, was identified based on its ability to bind Rlk/Txk in a yeast two-hybrid screen of a mouse T cell lymphoma library. RIBP was also found to interact with a related member of the Tec family of tyrosine kinases, Itk. Expression of RIBP is restricted to T and natural killer cells and is upregulated substantially after T cell activation. RIBP-disrupted knockout mice displayed apparently normal T cell development. However, proliferation of RIBP-deficient T cells in response to T cell receptor (TCR)-mediated activation was significantly impaired. Furthermore, these activated T cells were defective in the production of interleukin (IL)-2 and interferon γ, but not IL-4. These data suggest that RIBP plays an important role in TCR-mediated signal transduction pathways and that its binding to Itk and Rlk/Txk may regulate T cell differentiation.

Key words: T cell activation • signal transduction • adaptor protein • Tec tyrosine kinases • T helper type 1/T helper type 2 cells

Abbreviations used in this paper: ES, embryonic stem; KO, knockout; PRR, proline-rich region; PTB, phosphotyrosine-binding; PTK, protein tyrosine kinase; RAG, recombine-activating gene; RIBP, Rlk/Itk-binding protein; SH, Src homology; TBST, Tris-buffered saline/Tween.

It is well established that T cell activation critically depends on signal transduction pathways mediated by protein tyrosine kinases (PTKs) of the Src (Fyn and Lck) and Syk (Syk and ZAP-70) families (1–4). In contrast, the biological functions of members of the Tec family of tyrosine kinases are not well understood, nor have the biochemical mechanisms by which they operate been elucidated. These kinases contain a Src homology (SH)3 domain capable of binding proline-rich sequences, an SH2 domain capable of binding phosphotyrosine-containing sequences, and a tyrosine kinase domain (5, 6). However, unlike other Src family kinases, members of the Tec family characteristically lack a negative regulatory tyrosine residue at the COOH terminus (7–9), whose dephosphorylation is normally required for kinase activity.

Multiple members of the Tec subfamily have been identified in T cells and B cells, including Itk, Rlk/Txk, Btk, and Tec. Itk, a 72-kD IL-2-inducible PTK expressed in T cells and NK cells (9, 10), has been reported to be involved in T cell activation. Mice deficient in Itk display deficits in both T cell development and peripheral T cell activation (11); these T cells exhibit a diminished intracellular Ca2+ flux in response to TCR/CD3 signaling, and consistent with this, reduced phospholipase C-γ1 phosphorylation and inositol trisphosphate production (12). Additionally, Itk has been implicated in CD28 signal transduction, although whether it functions to positively or negatively regulate this pathway is unclear (13–15). Tec is expressed in multiple hematopoietic and nonhematopoietic cell types, including T cells (16, 17). Although a recent report suggests that it may function in antigen receptor signal transduction in T cells (18), other evidence suggests that Tec is involved in signaling through receptors for the cytokines IL-3, IL-6, and GM-CSF (19–21, respectively). Finally, we identified previously a novel member of the Tec subfamily of Src-related PTKs, Rlk (also called Txk), that is expressed primarily in T cells, mast cells, and testes (22, 23). Unlike other Tec subfamily members, Rlk (a 62-kD protein) lacks an NH2-terminal pleckstrin homology domain, and its expression is downregulated in naïve T cells after activation. Moreover, there is substantially more Rlk expressed in Th1-
type than T h2-type cells (22), suggesting that it may be involved in T cell differentiation. Although one recent study implicates Rlk in the binding of the cytoplasmic tail of the counter-costimulatory receptor CTLA-4 (24), little is known about the biochemical targets of this PTK or its possible functions in T cell activation.

To begin to elucidate the function(s) of Rlk and related Tec kinases, we sought to identify associated proteins using the yeast two-hybrid system for analyzing interprotein interactions. A unique cDNA clone was identified that interacted with both Rlk and Itk. This cDNA encodes a putative adaptor protein, termed RIBP (for R lk/Itk-binding protein). RIBP mRNA is expressed in T cells and N K cells and is significantly upregulated after T cell activation. Finally, targeted disruption of the RIBP gene via homologous recombination was shown to inhibit TCR/CD3-induced T cell proliferation, and IL-2 and IFN-γ production, but not IL-4 production. Together these data suggest that this novel molecule may provide an important adaptor function for Tec subfamily kinases critically involved in T cell activation and differentiation.

Materials and Methods

Bait cDNA Cloning and Yeast Two-Hybrid cDNA Library Screen. The Y153 Saccharomyces cerevisiae strain (auxotrophic for adenine, uracil, leucine, and tryptophan) as well as the bait and trap expression vectors, pAS1 and pACT, respectively, have been described (25). The Rlk and Itk chimeric bait constructs were generated by cloning full-length cDNA, PCR-amplified using primers encoding Ncol and BamH I sites, into the Ncol and BamHI sites in the multiple cloning site of pAS1. The Itk cDNA in pcdNA1 (Invitrogen) was a gift of Dan Littman (New York University, New York, N.Y.). A mouse T cell lymphoma MATCHMAKER cDNA library in pACT (Clontech) was screened using the Rlk bait as per the manufacturer’s protocol. Full-length clones were identified by screening a custom-made Uni-ZAP mouse day 16 fetal thymic cDNA library (Stratagene) using a cDNA probe derived from the RIBP coding region.

For Yeast Two-Hybrid cDNA Library Screen, total RNA was isolated from the indicated tissues and cell types using TRIzol (GIBCO BRL) according to the manufacturer’s protocol or as described previously (23). Northern blots were probed for the presence of RIBP mRNA using 32P-labeled cDNA probes derived from either full-length RIBP or RIBP 1–208 (encoding RIBP from the NH2 terminus to the end of the SH2 domain) by random priming, using a Primer-It II kit (Stratagene). The RIBP transcript was detected as a band ~1.7 kb in size, migrating faster than the 18S rRNA band.

For Southern blot analysis, genomic DNA was prepared from embryonic stem (ES) cell clones by standard methods, and digested with the indicated restriction endonucleases. The DNA was dephosphorylated and, after gel electrophoresis and membrane transfer, hybridized to either a 5’ or 3’ 32P-labeled RIBP cDNA probe (as indicated in the text, and see Fig. 4 B). Bands representing either the endogenous or targeted RIBP allele were detected based on size, as described in the text and in the legend to Fig. 4 B. Targeted deletion of the RIBP gene and generation of RIBP-deficient Mice. The targeting construct was electroporated into the R1 ES cell line, derived from the 129/Sv mouse strain, using standard techniques (29). Homologous recombination at the RIBP locus was detected by Southern blot analysis using 5’ and 3’ probes binding outside the RIBP regions used in the targeting construct (see above, and Fig. 4 B). Targeted ES cell clones were used for blastocyst injection, and the resulting chimeric animals were backcrossed to C57BL/6 mice. Agouti offspring carrying the targeted RIBP allele were interbred to generate progeny homozygous for the targeted RIBP allele. Progeny were typed by Southern blot analysis of tail genomic DNA (see Fig. 4 B), and subsequently, by PCR analysis, using primer pairs which either amplified endogenous sequence between exons 2 and 3 or sequences in the neo cassette.

Flow Cytofluorimetric Analysis. Single cell suspensions from thymi, lymph nodes, and spleens were prepared according to standard methods. Cells were then blocked with an anti-F,R mAb, 2.4G2, and stained with the Abs indicated in the text. Abs used in these experiments were conjugated to biotin or one of the following: Texas Red-goat anti-mouse IgG (Southern Biotechnology Associates), Texas Red-PEA goat anti-rat IgG (BD PharMingen, San Diego, CA), PE-anti-mouse IgG2b (BD PharMingen), and FITC-anti-mouse IgM (BD PharMingen).
following fluorochromes FITC, PE, or Cy-Chrome. Ab-stained cells were analyzed on a FACScan™ (Becton Dickinson) flow cytometer.

Proliferation Assays. Whole lymph node cells, splenocytes, or a 1:1 mixture of lymph node or splenic T cells cultured with T cell–depleted, irradiated syngeneic splenic APCs, from wild-type (either progeny from heterozygous intercrosses or [C57BL/6 × 129] F2 mice; The Jackson Laboratory), heterozygous, or homozygous knockout (KO) mice were used in these experiments. Soluble anti-CD3e mAb (145-2C11) with or without soluble anti-CD28 mAb (PV-1) was added to wells of round-bottomed 96-well plates (Costar) at the concentrations indicated in the text. When used, final concentrations of PM A and ionomycin were 10 ng/ml and 0.5 μM, respectively. Cells (2 × 10^6) were added to each well at a final concentration of 1 × 10^6/ml and cultured at 37°C in DMEM supplemented with 10% FCS, penicillin/streptomycin (100 μg/ml) and 2-ME. After 40 h, individual wells were pulsed with [3H]thymidine (1 μCi/well) for 8 h, for a total duration of 48 h of stimulation. Cell contents were then harvested onto fiberglass filters, scintillation fluid was added (25 μl/well), and counts were determined using a TopCount microplate scintillation counter (Packard).

IL-2 ELISAs. IL-2 ELISAs were performed on culture supernatants of activated cell cultures (described above). Nunc-immuno plates (N gel N unc) were used for these assays. Plates were coated with an anti-IL-2-coating Ab overnight at 4°C, washed, and blocked in 2% BSA/PBS for 1 h at room temperature. Supernatants diluted in 2% BSA/PBS, were incubated on the coated plates overnight at 4°C. The next day, plates were washed, then incubated with an anti-IL-2 detecting Ab for 1 h at room temperature. Plates were washed and incubated with a 1:5,000 dilution of streptavidin-horseradish peroxidase (Zymed), after which they were developed with TMB One-Step Substrate (Dako). Abs and IL-2 standards were all purchased from Endogen.

R N ase Protection Assays. Splenocytes from wild-type (either [C57BL/6 × 129] F2 described above, or when indicated, C57BL/6 mice), heterozygous, and homozygous KO mice were activated with the indicated concentrations of soluble anti-CD3 mAb with or without anti-CD28. At the indicated times, cells were harvested, and R N A was prepared as described above. Cyto- kine R N ase protection assays were performed using the RibopQuant M ulti-Probe R N ase Protection Ass Say System (Pharmingen) according to the manufacturer’s protocol. In brief, a 32P-labeled R N A probe was synthesized via in vitro transcription from the mCK-1 plasmid template. Aliquots of 2.5 μg of total RNA per sample were hybridized overnight at 56°C to the probe. Samples were then digested with RNase A, and after which they were treated with Proteinase K and phenol/chloroform extracted. Samples were then precipitated, resuspended in 1× loading buffer, heated briefly at 90°C, and subjected to electrophoresis on a 5% denaturing polyacrylamide gel. Gels were dried, and autoradiography was performed. Densitometry on the cytokine and housekeeping gene bands was performed using a Molecular Dynamics ImageQuant® densitometer.

Results

R I B P is a Novel Adaptor Protein That Interacts with R Lk and I t k. The yeast two-hybrid system of detecting interprotein interactions was used to identify potential R Lk-binding proteins that might mediate or regulate Tec family kinase function. A cDNA encoding full-length R Lk was cloned into a yeast vector encoding the GAL4 DNA-binding domain, thus yielding a “bait” fusion protein consisting of R Lk physically linked to this DNA-binding protein. A mouse T cell lymphoma cDNA library was screened using the R Lk bait. This library consisted of “trap” cDNAs cloned into a yeast vector encoding the GAL4 transactivation domain. The T cell cDNA library was chosen as R Lk is highly expressed both in the thymus and in peripheral T cells. Two specificity criteria were used as measures of GAL4-mediated transprokine activation: (a) histidine synthesis by the yeast, and (b) β-galactosidase synthesis. A total of 750,000 yeast colonies were screened. 86 of these colonies were identified as positive for bait-trap interaction by the first pass criterion of growth on histidine-deficient (His-) medium supplemented with 3-aminotriazole, an inhibitor of weak endogenous histidine synthesis. Among the 86 clones, 56 were positive for interaction with the R Lk bait fusion protein based on the second criterion of β-galactosidase synthesis (blue/white selection). The cDNA library plasmids isolated from the β-galactosidase-positive yeast colonies were subsequently transformed into bacteria to distinguish multiple trap cDNAs present within the same yeast colony during the original screening. A total of 65 unique cDNA clones were identified and subsequently retransformed into yeast. One clone, 2.3.2, was found to interact with R Lk with high specificity, and was chosen for further study.

Initial sequence analysis demonstrated that clone 2.3.2 did not represent a full-length cDNA; consequently, a mouse fetal thymic cDNA library was screened using a probe derived from the positive yeast cDNA clone. This screen yielded a full-length cDNA that was used to examine interactions with R Lk and I t k. Yeast transformed with the full-length cDNA, termed R I B P, interacted specifically with both R Lk and I t k (Fig. 1 A, bottom; nos. 2 and 6, respectively). We further examined whether the interaction of R I B P with R Lk depended on the kinase function of R Lk. An altered R Lk that contained a point mutation in the kinase domain (K309→R, denoted K309R) previously shown to abolish kinase activity (our unpublished observations) was incapable of binding R I B P (Fig. 1 A, bottom; no. 4). This suggests that the bait–trap interaction is critically dependent on the functionality and activity of the tyrosine kinase domain of R Lk.

Human epithelial kidney (HEK293) cells were transfected transiently with constructs encoding HA-tagged R I B P and/or I t k to directly examine protein–protein interactions in mammalian cells. Cell lysates were immunoprecipitated with an anti-HA Ab, resolved by SDS-PAGE, and subjected to immunoblotting using an anti-Itk antiserum. As shown in Figure 1 B, a specific interaction between R I B P and I t k was detected (top panel, lane 5). Coexpression of the Src family kinase, Lck, increased the amount of I t k coprecipitated with R I B P (Fig. 1 B, top; compare lane 6 with lane 5). This Lck-dependent augmentation of R I B P–I t k complex formation was not due to increased levels of I t k in the cell lysates or R I B P-HA in the immunoprecipitates (Fig. 1 B, top, lane 2 vs. lane 1; and bottom, lane 6 vs. lane 5, respectively). The results suggested that the interaction
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of RIBP with Itk is regulated, at least in part, by Src family kinase-specific phosphorylation of one or both proteins.

Structural Analysis and Chromosomal Location of RIBP. Sequence and genomic analyses were performed on the RIBP gene product. RIBP contains an SH2 domain (highly homologous to the SH2 domain of Shc, 47% [data not shown]), a proline-rich region (PRR) capable of binding SH3 domains, and an NPIY substrate sequence for binding by phosphotyrosine-binding (PTB) domains (30) present within the PRR (Fig. 1 C). Further analysis of the cDNA library identified a second RIBP transcript that represented an alternatively spliced form with an 8 amino acid sequence insert (VWASQQKA). Other potentially significant amino acid sequences within RIBP include two YXXP motifs, implicated in binding the SH2 domains of rasGAP, Abl, and Crk (31), and one YXXV motif, potentially capable of binding the SH2 domain of the tyrosine phosphatase, SHP-2 (32). Additionally, RIBP contains three potential sites for N-myristoylation (Fig. 1 C). These structural features and the absence of a catalytic domain suggest that RIBP functions as an adaptor protein. One adaptor protein, TSAd (33), showed extensive sequence homology to RIBP. The human TSAd protein exhibits a 68% sequence identity and a 76% sequence similarity with RIBP. The sequence ho-
mology between RIBP and TSAd was greatest within the SH2 domain, where the two proteins share an 87% sequence identity (80/92 amino acids; Fig. 1 C). Furthermore, TSAd exists in two alternatively spliced forms, similar to RIBP, with a 10 amino acid insert present in the longer form. Although these results suggested that RIBP and TSAd may be species homologues, the sequence insertions in RIBP and TSAd are in different locations within the molecules. The 10 amino acid insertion in TSAd is located within the SH2 domain, whereas the 8 amino acid insertion in RIBP is located within one of the exons in the region between the SH2 domain and the PRR (Fig. 1 C).

Genomic linkage studies of the RIBP gene were performed by Southern blot analyses of two sets of multilocus crosses (described in Materials and Methods). Digestion of DNA samples from parental mice with HindIII produced RIBP fragments of 8.6 and 2.9 kb in M. spretus and 11.0 and 10.8 kb in NFS/N. BamHI produced fragments of 12.4 and 4.0 kb in NFS/N and 16.5 kb in M. m. musculus. Inheritance of the variant fragments was compared with that of other markers previously typed in these crosses. The RIBP locus was mapped to mouse chromosome 3 (Fig. 2). Closest linkage was detected with Gba, for which no recombinants were detected in 171 mice, indicating that the genes are within 1.7 cM of one another at the upper limit of the 95% confidence interval. Similarly, no recombination was detected with Lma in 103 mice, placing RIBP within 2.9 cM of Lma. The RIBP gene maps to a region of mouse chromosome 3 with conserved synteny to human chromosome 1q21 (34, 35), where both human GBA (36) and LMNA (37), as well as TSAd (33) and Shc (38), are located. Thus, the RIBP gene has a genomic localization congruent with that of the TSAd and Shc genes, supporting the possibility of gene duplication events in this region.

RIBP Is Restricted in Expression to T Lymphocytes and N K Cells. Tec family members have been shown to have restricted tissue expression. In the cases of Rlk and Itk, expression is largely restricted to T cells with Itk also expressed in N K cells and Rlk also expressed in mast cells. Northern blot analyses were performed on RNA samples isolated from various tissues in order to examine the tissue expression of RIBP. A 1.7-kb transcript representing RIBP was expressed in the thymus, lymph node, and to a lesser extent, in the spleen and bone marrow (Fig. 3 A, left panel, lane 5; right panel, lanes 1 and 2; and left panel, lanes 4 and 3, respectively). In addition, analysis of multiple hematopoietic cell lines revealed that RIBP was expressed in T cell lines (e.g., BW.P and C57), but not expressed in a B cell lymphoma, A20, or a mastocytoma, P815 (data not shown). Northern blot analyses of thymocyte subpopulations demonstrated that RIBP mRNA was expressed in CD4+CD8+ thymocytes (Fig. 3 A; right panel, lane 5), and in both CD4+ and CD8+ peripheral T cells (Fig. 3 A, right panel, lane 6), little expression was detected in recombinase-activating gene (RAG)-2–deficient thymus (Fig. 3 A, right panel, lane 7). These data suggest that RIBP expression may be dependent on signal transduction through the pre-TCR, and consequently, that its expression levels vary during the course of T cell development.

RIBP mRNA was found to be rapidly induced after T cell activation (Fig. 3 B, top panel). RIBP expression was upregulated in the Th1 T cell clone, A.E7, in as little as 2 h after exposure to anti-CD3 (Fig. 3 B, top left panel). Similarly, RIBP mRNA was induced in another Th1 T cell clone, pSL10, when activated with anti-CD3 (Fig. 3 B, top right panel) or ovalbumin in the presence of appropriate APCs (data not shown). The addition of exogenous IL-2 had no effect on RIBP expression, although it significantly increased [3H]thymidine incorporation (Fig. 3 B, top right panel, and data not shown). In addition to thymocytes, N K cells stimulated with IL-2 were observed to express high levels of RIBP mRNA (Fig. 3 B, bottom panel). Furthermore, RIBP was also expressed in the Th2 clones PL104 and PL3, as also shown in Fig. 3 B (bottom panel).

Generation of RIBP-deficient Mice. RIBP KO mice were generated by disrupting the RIBP gene via homologous recombination. A targeting construct was generated that replaced exons 3 and 4 (encoding the SH2 domain) with a DNA segment encoding a neo gene. The targeting construct, depicted in Fig. 4 A, was introduced into the R1 ES cell line derived from the 129/Sv mouse strain. Southern blot analyses of drug-resistant ES cells demonstrated successful targeted disruption at the RIBP locus (Fig. 4 B, top). Genomic DNA samples from control and targeted ES cells were digested with EcoRI and XhoI, resolved by gel electrophoresis, and hybridized to a 5′ RIBP cDNA probe. As seen in Fig. 4 B (top left panel), probing of control digested ES cell DNA revealed bands of the predicted size (3.8 kb).

**Figure 2.** Chromosomal localization of mouse RIBP. To the right of the map are recombination fractions between adjacent loci, with the first fraction representing data from the M. m. musculus crosses and the second from the M. spretus crosses. Numbers in parentheses are recombinational distances ± SE. The M. m. musculus crosses were not typed for D3Mit22.
In contrast, DNA from targeted ES cell clones revealed bands migrating at 5.5 kb, consistent with the replacement of the third and fourth exons with the neo cassette. Similar results were observed using a 3" RIBP cDNA probe of control versus targeted ES cell genomic DNA samples digested with Asp718 (an isoschizomer of KpnI) and XhoI (Fig. 4B, top right panel). The neomycin-resistant ES cells (clone 2) were injected into C57BL/6 blastocysts using standard techniques (29). Chimeric mice were bred to C57BL/6 mice to establish germline transmission, and heterozygous mice were interbred to obtain homozygous KO progeny. Southern blot analyses performed on tail genomic DNA samples from the progeny demonstrated successful targeted disruption at the RIBP locus (Fig. 4B, bottom panel). Initial studies revealed that homozygous KO mice lacked RIBP mRNA expression based on Northern blot analyses (data not shown). The lack of RIBP expression in the RIBP KO offspring was confirmed by immunoblot analyses using an RIBP-specific polyclonal antiserum. As seen in Fig. 4C, lysates from wild-type C57BL/6 and heterozygous splenocytes activated with anti-CD3 contained significant amounts of RIBP protein migrating at ~46 kD. In contrast, no RIBP protein was detected in activated T cells from homozygous RIBP-deficient animals. Analyses of Thymic and Peripheral T Cell Populations in RIBP-deficient Mice. RIBP KO mice did not display any gross developmental abnormalities, and a normal Mendelian distribution of RIBP KO progeny was observed. The percentages of CD4+CD8−, CD4+CD8+, CD4+CD8−, and CD4−CD8+ thymocytes did not differ between RIBP KO and wild-type (Fig. 5A) or heterozygous (data not shown) control mice, and no significant qualitative or quantitative differences in lymph node and splenic T cell populations were evident (Fig. 5B, and data not shown, respectively). In addition, the CD4+/CD8+ ratio in both lymph node (Fig. 5B) and spleen (data not shown) was similar among the various groups. Moreover, no obvious differences in the sizes or cell yields of thymus, spleen, and lymph nodes isolated from RIBP KO mice were apparent compared with wild-type control mice (data not shown). These data suggest that RIBP does not have an essential role in T cell development or homeostasis of peripheral T cell populations.

Proliferative Responses of RIBP-deficient T Cells in Response to TCR Ligation Are Impaired. Given the findings that (a) RIBP expression is augmented by T cell activation, and (b) its binding partners Itk and Rlk have been shown to be requisite for T cell responses to activation stimuli (11, 39), studies to evaluate the functional status of RIBP KO T cells were undertaken. In response to TCR/CD3-mediated signals delivered by an anti-CD3 mAb, proliferation of RIBP KO T cells was considerably reduced relative to control T cell proliferation (Fig. 6A). This proliferative impairment was observed across the range of anti-CD3 concentrations tested, and ranged from an 80% reduction in proliferation at the lowest (anti-CD3 concentration) to a 50% reduction in proliferation at the highest (anti-CD3 concentrations). Although RIBP KO T cells displayed an apparent defect in TCR-mediated T cell activation, they were capable of receiving CD28-mediated co-stimulation (Fig. 6A). These data indicate that RIBP KO T cells are impaired in their ability to respond to TCR-mediated signals, whereas responsiveness to CD28 signals is less affected.

To determine whether the RIBP KO T cells were defective in either proximal or distal components of the TCR signaling pathway, RIBP KO T cells were treated with either anti-CD3 or PMA and ionomycin, and proliferation
was compared with that of wild-type T cells activated with the same stimuli. As shown in Fig. 6 B, RIBP KO T cell proliferation in response to anti-CD3 was 50% reduced compared with wild-type T cells. However, wild-type and RIBP KO T cells stimulated with PMA and ionomycin proliferated equivalently, suggesting that the inadequacy in TCR-mediated signaling in RIBP KO T cells lies upstream of pathways triggered by protein kinase C activation or intracellular Ca\(^{2+}\) flux. These findings are consistent with those reported for T cells deficient in either Itk (11) or both Itk and Rlk (39).

Reduced Cytokine Production by Activated RIBP KO T Cells. RIBP KO splenocytes were stimulated with anti-CD3 and examined for cytokine production. Consistent with the reduced proliferative response (Fig. 6, A and B), a significant reduction in the amount of IL-2 in the culture supernatant was observed (Fig. 6 C). Furthermore, although CD28-mediated costimulation increased IL-2 production by both wild-type and KO T cells, the relative differences in IL-2 levels were maintained. These results were confirmed using an RT-PCR protection assay to determine relative levels of IL-2 transcripts in activated RIBP KO versus wild-type T cells. As shown in Fig. 6 D, IL-2 mRNA was detected at 6 h in (C57BL/6 \times 129) F2 wild-type T cells stimulated with anti-CD3. In contrast, little IL-2 mRNA was apparent in activated RIBP KO T cells at this time point (Fig. 6 D, lane 5 versus lane 4). Denstometry analyses revealed that IL-2 mRNA levels in RIBP KO T cells were reduced by \( \approx 80\% \) relative to wild-type cells (data not shown). Also, consistent with the reduction in supernatant levels of IL-2 (Fig. 6 C), the relative amounts of IL-2 mRNA from RIBP KO T cells activated with anti-CD3 and anti-CD28 were still significantly reduced compared with control T cells (Fig. 6 D, lane 7 versus lane 6).

In addition to a reduction in IL-2 levels, a pronounced decrease in IFN-\(\gamma\) production was observed in RIBP KO T cells (Fig. 7 A). As was the case for IL-2, this reduction in IFN-\(\gamma\) production correlated with decreased IFN-\(\gamma\) mRNA levels (Fig. 7 B), suggesting that RIBP affects lymphokine production at the level of transcription and/or mRNA stabilization. Little IFN-\(\gamma\)-mRNA was observed in anti-CD3-activated RIBP KO T cells relative to wild-type T cells; denstometry revealed that this was a \( >90\% \) reduction (Fig. 7 B; densitometry data not shown). However, anti-CD28 costimulation did restore the IFN-\(\gamma\) decrease observed in RIBP KO T cells, suggesting that this defect may be an indirect consequence of the reduced activation and IL-2 production. Finally, despite the reductions in IL-2...
and IFN-γ production by RIBP KO T cells, no significant reduction in IL-4 production was observed (Fig. 7 C). In fact, in some experiments, activated RIBP KO splenocytes produced more IL-4 than did control splenocytes. Thus, the lack of RIBP appears to more profoundly affect the ability of T cells to produce Th1-type cytokines than Th2-type cytokines.

Discussion

In this study, a yeast two-hybrid T cell cDNA library screen was used to search for proteins that could associate with the T cell–specific members of the Tec family of tyrosine kinases, Rlk and Itk. A novel adaptor molecule, termed RIBP, was identified that interacts with a site in Rlk and Itk regulated by tyrosine phosphorylation. RIBP is expressed in thymocytes, and at low levels in resting peripheral T cells. Expression of RIBP is upregulated after T cell activation, coinciding with the expression levels of one of its binding partners, Itk, and high expression levels are observed in activated NK cells. The restriction of RIBP expression to T cells and NK cells mirrors the expression patterns of several signaling molecules, including Itk (9), linker for activation of T cells (LAT [40]), which is also expressed in mast cells, and ZAP-70 (41). These results are consistent with the proposed common lineage of these cells (42). Finally, genetic disruption of RIBP was found to significantly impair TCR/CD3-mediated proliferation, and IL-2 and IFN-γ production, although IL-4 production was unaffected. Furthermore, the TCR-mediated signal transduction defects observed in RIBP KO T cells were bypassed using the downstream mitogens, PMA and ionomycin. These results suggest that RIBP positively regulates TCR signaling by acting as a molecular link between Itk/Rlk and other components of the TCR signaling pathway. The consequences of this interaction include the regulation of TCR-mediated signal transduction, and consequently, lymphokine production.

RIBP is a complex adaptor protein. The gene encodes a PRR that may bind SH3 domain–containing proteins such as other Src family kinases and other adaptor proteins. In addition, a tyrosine-phosphorylated NPIY sequence present in RIBP may bind PTB domain–containing proteins such as Shc. Proteins such as Shc have been implicated in TCR signal transduction via interactions with ZAP-70 (43) and Cbl, an adaptor which is thought to negatively regulate ZAP-70 activation (44, 45) and binds to the SH3 domain of Itk. The tyrosine-based protein binding motifs present in RIBP (YXXP and YXXV) may bind the SH2 domains of rasGAP, Abl, Crk, and SHP-2, molecules previously shown to regulate T cell signal transduction. Finally, RIBP contains an SH2 domain that may be involved in binding to other phosphotyrosine-containing proteins. In this regard, preliminary results indicate that the SH2 domain of RIBP is required for interaction with Itk, whereas the PRR and tyrosine-based protein binding motifs are unnecessary (our unpublished observations).

The molecular mechanisms that link RIBP, Rlk, and Itk, to the proximal event of TCR ligation are unclear. The finding that Lck coexpression augments the association of RIBP with Itk is consistent with a role for RIBP in the regulation of TCR signal transduction. Tyrosine phosphorylation of Itk, catalyzed by Lck or Fyn, is required for optimal Itk kinase activity (46, 47), and consequently, T cell activation. Phosphorylation of Itk occurs at Y511, a residue in the activation loop of the kinase domain of Itk (48). Thus, RIBP could be functionally coupled to TCR engagement, as its binding to Itk is enhanced by the activity of this CD4/CD8 coreceptor–associated Src family kinase in vitro.

RIBP may bind both Itk and Rlk, raising the question as to its physiological partner and the relationship of its binding specificity to the functional deficit evident in RIBP KO T cells. The expression pattern of RIBP suggests that its physiological binding partner may be Itk in the periphery. Previous studies have shown that Rlk is highly expressed in resting T cells, but downregulated after T cell activation. In
mice were stimulated with the indicated concentrations of soluble anti-CD3 alone (triangles), with or without soluble anti-CD28 added at 1.0 μg/ml (circles). Proliferation was assessed at 48 h, with a [3H]thymidine pulse added after 40 h, for an 8-h pulse duration. Results are representative of four independent experiments performed using splenocytes, and three independent experiments using purified T cells and irradiated, T cell–depleted APCs. (B) Proliferation of T cells from wild-type (WT) and RIBP KO mice in response to either anti-CD3 or PMA plus ionomycin (P + I). Proliferation was assessed as above. (C) Splenocytes from wild-type and RIBP KO mice were stimulated with 1 μg/ml of soluble anti-CD3, with or without 1 μg/ml of soluble anti-CD28. Supernatants were collected at either 6, 24, or 48 h, and the concentration of IL-2 was determined by ELISA; values listed above were obtained at 24 h. Results are representative of 11 independent experiments. (D) Splenocytes from RIBP KO or wild-type mice were activated for the indicated times with anti-CD3 with or without anti-CD28. RNA prepared from cells was subsequently subjected to RNase protection assays, as described in Materials and Methods. 32P autoradiogram of RNase protection assay polyacrylamide gel. Top: IL-2 mRNA levels in wild-type and RIBP KO T cells activated for the indicated times with the stimuli listed. EL4 mRNA is a positive control for cytokine gene expression, and yeast tRNA is a negative control. Bottom: mRNA levels for the L32 housekeeping gene. Results are representative of three independent experiments.

contrast, Itk gene expression is upregulated after T cell activation, similar to RIBP. Furthermore, the phenotype observed in RIBP KO T cells more closely resembles the phenotype of Itk-deficient T cells, which display diminished proliferation and IL-2 production in response to TCR ligation. In contrast, Rlk KO T cells have only minor deficits in TCR-mediated T cell activation (39). However, there are circumstances when the binding of RIBP to Rlk may be physiologically relevant. For instance, both RIBP and Rlk are expressed in thymocytes. Although we have not identified any profound changes in T cell development in the RIBP KO or Rlk KO mice (our unpublished observations, and reference 39, respectively), there may be subtle, as yet unidentified effects of this interaction in the thymus. Moreover, both Rlk and RIBP are expressed in NK cells, and may play a role in this lymphocyte subset. In fact, recent evidence indicates that Itk and Rlk have overlapping functions in T cell activation, as mice deficient in both kinases have more severely diminished peripheral T cell function than mice deficient in either kinase alone (39). Thus, further biochemical and breeding studies to generate RIBP/Itk and RIBP/Rlk double-KO mice will be needed in order to determine the differential effects of RIBP on the functions of Itk versus Rlk. It is important to emphasize that the functional impairment present in RIBP KO T cells was not as severe as that observed in Rlk/Itk double-KO T cells. There are two possible reasons for these findings. First, there may be functional homologues of RIBP that can partially compensate for the absence of RIBP in KO T cells. Second, not all of the intracellular effects on
TCR signaling that Rlk and Itk mediate may require RIBP or any putative homologues. The fact that CD28-mediated costimulation partially compensates for the severely diminished proliferative response to TCR/CD3-mediated T cell activation in RIBP KO mice (Fig. 6 A) suggests that the RIBP KO T cells are capable of responding to the CD28-mediated costimulatory signal. Interestingly, Itk has been reported to be both a positive and negative regulator of CD28-mediated T cell activation. On the one hand, Itk is physically associated with CD28 after T cell activation (49), and Jurkat T cells expressing CD28 cytoplasmic tail mutants incapable of recruiting Itk are suppressed in their ability to produce IL-2. However, Littman and colleagues demonstrated that T cells from Itk-deficient mice were hyperresponsive to CD28 costimulation (15). Therefore, it remains to be determined in what context, if any, RIBP is involved in CD28 signal transduction.

Finally, the results of this study support a role for RIBP in the regulation of Th cell subset differentiation. IL-2 and IFN-γ production were both significantly reduced in activated RIBP KO T cells (Figs. 6 and 7), whereas IL-4 production was unaffected (Fig. 7 C). Such a selective impairment in IFN-γ and IL-2 production suggests that RIBP may act in signaling pathways that promote Th1 differentiation. The lack of RIBP may reduce TCR-mediated signal strength via impairment of Itk and/or Rlk function, resulting in a selective loss of Th1 differentiation, as suggested by Bottomly and colleagues (50). This may occur by generating decreased TCR signals that drive Th2 skewing. In fact, both Itk and Rlk have been implicated in T cell differentiation (Schaeffer, E., personal communication). Also, Rlk is selectively expressed in Th1-type T cells. These results may explain, at least in part, the finding that RIBP selectively affects Th1-type cytokine production.

In conclusion, we have identified and functionally characterized a novel T cell adaptor molecule involved in the regulation of responses to T cell activation stimuli, specifically proliferation and lymphokine production. This adaptor, RIBP, appears to function in TCR/CD3-mediated signal transduction, consistent with previously reported data regarding its binding partners, the Tec tyrosine kinases Rlk and Itk. The binding of RIBP to Itk and Rlk may provide important biochemical links of these two important kinases with other components in the T cell activation machinery. Further molecular studies will be needed to determine the specific sites of action of RIBP within various T cell signal transduction pathways.

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