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Disrupting the Intermolecular Self-Association of Itk Enhances T Cell Signaling

Lie Min,*1 Wenfang Wu,†1 Raji E. Joseph,* D. Bruce Fulton,* Leslie Berg, † and Amy H. Andreotti*

The Tec family tyrosine kinase (Itk), is a key component of the TCR signaling pathway. Biochemical studies have shown that Itk activation requires recruitment of Itk to the membrane via its pleckstrin homology domain, phosphorylation of Itk by the Src kinase, Lck, and binding of Itk to the SLP-76/LAT adapter complex. However, the regulation of Itk enzymatic activity by Itk domain interactions is not yet well understood. In this study, we show that full-length Itk self-associates in an intermolecular fashion. Using this information, we have designed an Itk variant that exhibits reduced self-association but maintains normal binding to exogenous ligands via each of its regulatory domains. When expressed in insect cells, the Itk substrate phospholipase Cγ1 is phosphorylated more efficiently by the Itk variant than by wild-type Itk. Furthermore, expression of the Itk variant in primary murine T cells induced higher ERK activation and increased calcium flux following TCR stimulation compared with that of wild-type Itk. Our results indicate that the Tec kinase Itk is negatively regulated by intermolecular clustering and that disruption of this clustering leads to increased Itk kinase activity following TCR stimulation. The Journal of Immunology, 2010, 184: 4228–4235.

Interleukin-2 tyrosine kinase (Itk) is a nonreceptor protein tyrosine kinase of the Tec family that is expressed in T cells, mast cells, and NK cells (1–5). Itk participates in signaling processes following TCR engagement by phosphorylating and activating phospholipase Cγ1 (PLCγ1), leading to production of two second messengers, 1,2-diacylglycerol and inositol 1,4,5-triphosphate (6–9). In addition to Itk, the Tec family includes Btk, Tec, Rlk, and Bmx, each of which shares a similar domain structure with the Src kinase family members (10). Both families contain a Src homology 3 (SH3) domain, a Src homology 2 (SH2) domain, and the catalytic domain. With the exception of Rlk, the Tec kinases also contain a pleckstrin homology (PH) domain and a Tec homology domain at the N terminus (Fig. 1A).

The mechanisms promoting Itk activation following TCR stimulation have been well described. To date, three upstream signals are required for Itk activation, including Itk recruitment to phosphatidylinositol 3,4,5-triphosphate in the membrane via its PH domain, Itk binding to the SLP-76/LAT adapter complex via its SH2 and SH3 domains, and finally Itk phosphorylation by Lck at the activation loop tyrosine in its kinase domain (11). However, the structural changes in Itk that accompany this activation process, as well as the mechanisms by which itk activity is turned off when TCR signaling is terminated, have not been established. One reason for this lack of information is that, despite similarities to the Src kinases in primary structure, there are also significant differences between Itk and Src (Fig. 1A). Most notably, the Tec kinases all lack the C-terminal autoinhibitory sequence that serves to negatively regulate the Src kinases (12–15). Thus, the well-characterized inhibitory interaction in Src between the Src SH2 domain and the phosphorylated Tyr527 in the C-terminal tail cannot occur for the Tec kinase family. The absence of the Src regulatory tail sequence in Itk and the related Tec family members raises questions about the domain interactions and conformational changes that regulate Itk activity during the course of T cell signaling.

We and others have previously reported detailed structural studies for regulatory domain fragments of Itk, Btk, Tec, and Rlk (16–25). An emerging theme for each of these kinases is that the noncatalytic domains form dimeric and higher-order oligomeric structures in solution. For Itk, self-association of the regulatory domains occurs via intermolecular interactions between the SH3 domain and the SH2 domain (18). The structure of the intermolecular Itk SH3/SH2 complex has been solved (16), providing a molecular basis for probing the functional significance of intermolecular association. An intermolecular interaction has also been described for the Itk PH domain (26). The isolated PH domain interacts both with itself and with the PH domain within full-length Itk in coimmunoprecipitation experiments, suggesting that multiple contacts across the regulatory domains stabilize a self-associated form of Itk. Finally, a split yellow fluorescent protein system has previously allowed visualization of intermolecular interactions between full-length Itk molecules in cells (27). Thus, abundant data point to intermolecular clustering of Itk, most likely mediated by multiple Itk regulatory domains, yet to date the functional significance of this self-association has not been explained.

In this paper, we extend the earlier studies of Itk domain fragments and demonstrate that full-length Itk self-associates in an intermolecular fashion in vitro. To evaluate the functional significance of the observed self-association, we then designed a mutant Itk molecule that retains all of the structural features of the wild-type...
enzyme yet exhibits diminished self-association. We next examined substrate phosphorylation levels by the wild-type and mutant Itk molecules as well as the signaling properties of wild-type and mutant Itk following expression in primary CD4+ T cells. The results of these experiments indicate that the Itk mutant exhibiting diminished self-association has increased activity and signaling capacity both in vitro and upon TCR engagement in primary T cells. On the basis of these findings, we discuss a mechanistic explanation for this observation and propose a model for the control of Itk activity during TCR signaling.

Materials and Methods

Constructs and baculovirus production

V5- or myc-tagged proteins were cloned into the pcDNA3.1/D/V5-HisTOPO vector (Invitrogen, Carlsbad, CA). Itk(BtkSH3) was generated by replacing the Itk SH3 domain sequence from Pro171 to Asn232 with the human Btk sequence spanning Ser214 to Ser275 by PCR. Flag-tagged full-length, wild-type (mouse Itk) or Itk(BtkSH3) were cloned into the pENTR/DTOPO vector (Invitrogen) by TOPO cloning. Point mutations (Y180F) were introduced using the QuickChange Site-Directed Mutagenesis Kit (Stratagene, La Jolla, CA). The pENTR vectors with various inserts were recombined in vitro with BaculoDirect C-Term Linear DNA (Invitrogen) according to the manufacturer’s instructions (Invitrogen) for virus production. The PLCγ1 baculovirus has been described previously (9).

Immunoprecipitation and Western blot

NIH 3T3 cells were transfected with V5- or myc-tagged DNA using Effectene Transfection Reagent from Qiagen (Valencia, CA). Twenty-four hours post-transfection, cells were lysed in buffer containing 0.5% NP-40, 50 mM Tris (pH 7.4), 150 mM NaCl, 10 mM MgCl2, and protease inhibitors (Roche Applied Science, Indianapolis, IN). Immunoprecipitation, Western transfer, and Western blotting were performed using standard techniques. Abs used throughout are as follows: anti-Erk (Cell Signaling Technology, Beverly, MA), anti–phospho-Erk (Cell Signaling Technology), anti-Ik (Upstate Biotechnology), Lake Placid, NY), anti-PLCγ1 (Upstate Biotechnology), anti-pY783 (Upstate Biotechnology), anti-myc (Invitrogen), and anti-V5 (Invitrogen). Anti-Btk pY223 (used to probe the analogous site in Itk, pY180) and anti-Btk pY551 (used to probe the analogous site in Itk, pY551) are from Dr. Owen Witte.

Protein expression and purification

The purification method for bacterially expressed protein has been described previously (18). For production of Itk enzyme for in vitro kinase assays, baculovirus encoding either full-length, wild-type Itk or Itk(BtkSH3) along with Lck virus (if indicated) were used to infect Sf9 cells in a 1:1 ratio. Proteins were purified using methods reported previously (28–30). A kinase inactive mutant of Itk is used to ensure that activity measurements reflect Itk activity alone and not copurifying Lck. Coexpression of Itk [or Itk(BtkSH3)] with PLCγ1 in Sf9 cells did not include Lck expression to avoid spurious phosphorylation of the substrate by Lck. It has been demonstrated previously that Itk produced in this manner is active (31).

Nuclear magnetic resonance spectroscopy

Nuclear magnetic resonance (NMR) spectra were recorded at 298 K on a DRX500 spectrometer (Bruker, Billerica, MA) operating at a 1H frequency of 499.867 MHz. Protein concentrations were adjusted to 1 mM. Global reactivity of 499.867 MHz. Protein concentrations were adjusted to 1 mM. Global reactivity of molecules as well as the signaling properties of wild-type and mutant Itk following expression in primary CD4+ T cells. The results of these experiments indicate that the Itk mutant exhibiting diminished self-association has increased activity and signaling capacity both in vitro and upon TCR engagement in primary T cells. On the basis of these findings, we discuss a mechanistic explanation for this observation and propose a model for the control of Itk activity during TCR signaling.

Purification of primary CD4+ T cells

Spleen and lymph node cells were isolated from wild-type or Itk−/− (6) C57BL/10 (The Jackson Laboratory, Bar Harbor, ME) mice, and CD4+ T cells were purified by positive selection using anti-CD4 Ab-coated magnetic microbeads (Miltenyi Biotec, Auburn, CA).

Retrovirus production

Itk and Itk(BtkSH3) were cloned into the retroviral mouse stem cell virus vector MSCV2.2-internal ribosomal entry site (IRES)-GFP (32). Phoenix-E retroviral packaging cells (Orbigen, San Diego, CA) were transfected with each retrovirus construct plus the pCL-Eco retrovirus packaging vector (Imegenex, San Diego, CA) as described previously (9), and virus particles were harvested and stored at −80°C. For each infection, 2 × 10⁶ primary CD4+ T cells were stimulated with 5 ng/ml PMA (Sigma-Aldrich, St. Louis, MO) and 375 ng/ml ionomycin (Calbiochem, La Jolla, CA) for 24 h and then incubated with 2 μl viral supernatant plus IL-2 (30 ng/ml) and Lipofectamine (Invitrogen). After 3–4 d, CD4+ GFP+ infected cells were sorted by flow cytometry and cultured a further 2 wk in IL-2. For analysis, 3 × 10⁶ cells were restimulated with biotinylated-anti-CD3 Ab (25 μg/ml; eBioscience, San Diego, CA) for 10 min, followed by streptavidin (50 μg/ml; Pierce Protein Research Products, Rockford, IL) crosslinking for 5 min. Cell lysates were analyzed for phospho-ERK by immunoblot; alternatively, cells were permeabilized and stained with anti-phospho-ERK Ab followed by flow cytometry.

Ca2+ flux assay

Wild-type Itk and Itk(BtkSH3) were cloned into the retroviral vector pMX-IRES-hCD8 (33). Four days postinfection, cells were sensitized with 10 μg/ml biotin–oCD3 for 10 min on ice in OPTI-MEM, washed, and incubated with 16 μM Fura-Red (Invitrogen) and 16 μM Fluo-3 (Invitrogen) in RPMI 1640/5% FBS at 37°C for 45 min. Cells were washed twice with RPMI 1640/5% FBS, incubated at room temperature in Tyrode’s buffer/BSA for 30 min. Cells were analyzed by flow cytometry with the addition of streptavidin at 30 s. Tyrode’s buffer/BSA for Ca2+ influx assay: contained 137 mM NaCl, 1.2 mM HEPES, 2.7 mM KCl, 0.04 mM NaH2PO4· H2O, 5.6 mM glucose, 0.5 mM CaCl2· H2O, 1 mM MgCl2·6H2O, and 2% FBS.

Statistical analysis

Statistical analysis was performed using the Student t test.

Results

The Itk SH3 domain interacts specifically with the Itk SH2 domain in a nonclassical manner

SH2 and SH3 domains are well-studied signal transduction modules that mediate protein–protein interactions to alter protein localization and to promote proximity for enzyme–substrate interactions. SH2 and SH3 binding domains typically mediate interactions with target proteins that contain phosphotyrosine motifs or proline-rich sequences, respectively. For instance, the Itk SH2 domain binds to phosphorylated tyrosine residues on SLP-76, and the Itk SH3 domain binds to proline-rich regions in SLP-76 and Vav1 (11). However, the regulatory SH2 and SH3 domains of Itk interact with each other via an intermolecular association that does not fit these established paradigms for SH2 and SH3 recognition (18). The Itk SH3/SH2 interaction is neither phosphotyrosine-dependent nor mediated by a polyproline motif (16). Thus, the Itk amino acid residues mediating this self-association are distinct from those involved in binding to other signaling proteins, such as SLP-76. Given this distinction, Itk provides us with an ideal opportunity to specifically disrupt self-association by altering the SH3 and/or SH2 domains in a manner that does not affect binding to canonical proline-rich or phosphophoryl tyrosine ligands. These alterations then would permit functional assays, allowing us to compare full-length, wild-type Itk to a full-length Itk variant in which self-association is disfavored.

Molecular determinants of the intermolecular SH3/SH2 interaction

The three-dimensional structure of the binary Itk SH3/SH2 complex is shown in Fig. 1B. Using this structure, we aimed to identify sequence changes that would abolish the interaction between the Itk...
SH3 and SH2 domains but maintain the classical ligand binding functions of the SH3 and SH2 domains, proline-rich peptide recognition and phosphotyrosine binding, respectively. A simplistic approach to this goal would be mutation of the central residue in the SH3 binding cleft, Trp208, which has been previously shown to disrupt the interaction between the SH2 domain and the SH3 domain of Itk (18). However, this conserved tryptophan also makes extensive contacts to cognate proline-rich ligands (34, 35) and is therefore not an appropriate target for mutation in our current study. In fact, the SH2/SH3 interface in this complex either fully or partially overlaps with the binding sites on these domains that contact the classical phospholigand and proline-rich ligand sequences. Because our goal is to selectively disrupt just the SH2/SH3 interaction without affecting the other binding functions of the Itk SH3 and SH2 domains, we turned our attention to an approach that would disrupt the SH3/SH2 interaction but maintain the canonical ligand binding characteristics of each of these regulatory domains.

The Btk SH3 domain is 51% identical to that of Itk. Of the Itk SH3 domain surface residues that directly contact the Itk SH2 domain in the SH3/SH2 complex, there are eight residues that differ between Itk and Btk SH3 domains (Fig. 2A). These sequence differences are enough to eliminate the intermolecular SH3/SH2 interaction. Using NMR spectroscopy, we measured binding between the Itk SH2 domain and the Btk SH3 domain. Changes in NMR peak position (chemical shift perturbations) indicate specific binding between proteins. Unlike the Itk SH3/Itk SH2 interaction that is readily detected by NMR, we find that no chemical shift perturbations are observed in NMR spectra upon mixing the Btk SH3 domain with the Itk SH2 domain (Fig. 2B). Thus, the Itk SH2 domain and the Btk SH3 domain do not show detectable binding to each other.

To ensure that the Btk SH3 sequence binds the canonical proline-rich ligand in a manner similar to the Itk SH3 domain, we next compared the affinities of the Itk and Btk SH3 domains for a proline-rich peptide derived from Slp-76 (Q184QPPVPPQRPMA195), previously shown to bind the SH3 domain of Itk (36). Separate titrations of the QPPVPPQRPMA peptide into 15N-labeled Itk or Btk SH3 domains induced chemical shift perturbations that were analyzed to determine their respective dissociation constants. The binding curves generated upon addition of increasing concentrations of the Slp-76 peptide into the Btk and Itk SH3 domains are indistinguishable (Fig. 2C). Thus, the Btk SH3 domain does not interact with the Itk SH2 domain yet maintains normal proline binding characteristics. These data provide a strategy for altering the full-length Itk sequence to disfavor the observed self-association of wild-type Itk while maintaining classical proline ligand binding characteristics. A full-length Itk protein [designated Itk(BtkSH3)] that contains the Btk sequence within the region that normally spans the Itk SH3 domain (Fig. 2D) was constructed. All of the other regions of Itk(BtkSH3) are identical to those of full-length, wild-type Itk.

Using the newly constructed Itk(BtkSH3) protein, we examined the phosphorylation status of both Tyr511 in the Itk activation loop and Tyr180 within the Itk SH3 domain. Tyr180 is strictly conserved among SH3 domains, and the corresponding residue in the Btk SH3 domain is Tyr223. Phosphorylation at both tyrosine sites (Tyr511 and Tyr180) accompanies activation of Itk in T cells and if disrupted would interfere with Itk function in T cells (9). Full-length, wild-type Itk and Itk(BtkSH3) were separately expressed as flag-tagged fusion proteins and were coexpressed with Lck in insect cells. The proteins were purified, separated by SDS-PAGE, and then probed by Western blotting with a Btk pY223-specific Ab [previously shown to recognize specific phosphorylation on both Itk Tyr511 and Btk Tyr223 (9)] or an Itk pY511-specific Ab. Both wild-type Itk and Itk(BtkSH3) were phosphorylated to the same degree at the expected sites: Tyr511 and Tyr511 (Fig. 2E). Thus, sequence changes in the full-length Itk(BtkSH3) diminish intermolecular self-association but do not alter the normal phosphorylation patterns associated with wild-type Itk activation.

To examine the extent that the intermolecular Itk SH3/SH2 interaction mediates self-association of the full-length Itk kinase, we generated differentially tagged versions of full-length, wild-type Itk and Itk(BtkSH3) for coinmunoprecipitation experiments (Fig. 3A). First, the V5- and myc-tagged wild-type Itk were transiently cotransfected into NIH 3T3 cells and subjected to immunoprecipitation using anti-myc Ab. The immunoprecipitated fraction was then resolved by SDS-PAGE and probed with an Ab to the V5 tag. As a control, NIH 3T3 cells were also transiently transfected with V5-tagged Itk and/or the empty vector alone. The V5-tagged full-length Itk is readily detected in anti-myc immunoprecipitates from cotransfected NIH 3T3 cells, indicating that full-length, wild-type Itk self-associates in an intermolecular fashion (Fig. 3A). These data are consistent with previous in vivo data that also indicate intermolecular association of full-length Itk molecules (27).

**Btk SH3 sequence limits Itk self-association**

In a manner identical to the wild-type Itk sequence, V5- and myc-tagged Itk(BtkSH3) were transiently cotransfected into NIH 3T3 cells and subjected to coimmunoprecipitation. Consistent with the absence of an intermolecular interaction between the isolated Itk SH2 domain and the Btk SH3 domains, coimmunoprecipitation data indicate that full-length Itk(BtkSH3) does not self-associate to a level that is detected in the coimmunoprecipitation experiments for wild-type Itk (Fig. 3B). The observed differences in the coimmunoprecipitation data for Itk and Itk(BtkSH3) suggest that
disruption of the noncanonical Itk SH3/SH2 interaction reduces intermolecular self-association of full-length Itk compared with that of wild-type protein. The functions of these Itk proteins were next examined both in vitro and in primary T cells. Recombinant flag-tagged, wild-type Itk and Itk(BtkSH3) were separately purified from baculovirus-infected insect cells and used for in vitro kinase assays. With the in vitro kinase assay that we have described previously for wild-type Itk (28), a biotinylated peptide substrate (30) was used to measure the initial velocity of wild-type Itk and Itk(BtkSH3) at different concentrations of enzyme (Fig. 4).

**FIGURE 2.** Specific disruption of the intermolecular SH3/SH2 interaction. A, The Itk SH3 and Btk SH3 domain structures are superimposed (only labeled side chains from the Btk SH3 domain are shown for clarity). The Itk SH3/SH2 interaction surface is indicated in orange on the SH3 structure (18). Within this binding surface, only eight residues are not conserved between Itk and Btk. Each of the nonconserved pairs of residues are labeled and indicated on the structure (orange are Itk side chains and red are Btk side chains). B, Select region of heteronuclear single quantum coherence spectra of 15N-labeled SH3 domains from either Itk (left panel) or Btk (right panel), indicating the extent of chemical shift perturbations associated with binding to an unlabeled Itk SH2 domain. For both, the black spectrum is 15N-labeled SH3 protein alone, and the red spectrum has been acquired following addition of equimolar unlabeled Itk SH2 domain in each case. The arrow indicates the change in resonance frequency for a representative peak upon addition of Itk SH2 to 15N-labeled Itk SH3 domain. Addition of unlabeled Itk SH2 domain to 15N-labeled Btk SH3 domain results in no change in chemical shift values. This result confirms the absence of an interaction between Itk SH2 and Btk SH3. C, Concentration dependence of the normalized 1H chemical shift (Δδ/Δδmax) for representative residues of 15N-labeled Btk SH3 domain (top panel) and 15N-labeled Itk SH3 domain (bottom panel) upon titration of the SLP-76–derived proline-rich peptide. The titration data were acquired and analyzed as previously described (21). D, Design of the Itk(BtkSH3) protein. The Itk SH3 sequence within full-length Itk is replaced with the SH3 sequence of Btk. The sequence of the Btk SH3 domain is shown in bold below the domain map of Itk, and the surrounding Itk residues within the resulting Itk(BtkSH3) are lightface and underlined. E, Western blot analysis using phosphotyrosine-specific Abs to the autophosphorylation site in the SH3 domain (Tyr180 for Itk and Tyr223 for Btk) and the activation loop tyrosine (Tyr511). (i) Lane 1, full-length, wild-type Itk; lane 2, full-length, mutant Itk (Y180F); and lane 3, Itk(BtkSH3) were expressed as fusions with the flag peptide and purified from insect cells, resolved by SDS-PAGE, transferred to polyvinylidene difluoride membrane, and blotted with the Btk Ab to pY223 (Btk numbering; this tyrosine corresponds to Tyr180 in Itk) to detect the autophosphorylation status in the SH3 domain (top panel). Cross reactivity of the Btk pY223 Ab for Itk pY180 has been previously demonstrated (9). (ii) Lane 1, full-length, wild-type Itk; and lane 2, Itk(BtkSH3) were prepared as described above and blotted with the anti-pY551 Ab to detect the activation loop phosphorylation status (top panel). Itk protein levels are indicated in the bottom panel for both i and ii.
not due to intrinsic activity differences between the wild-type Itk and Itk(BtkSH3) constructs and 2) the increase in activity is a concentration-dependent phenomenon, consistent with a role for the SH3/SH2 intermolecular interaction in regulating Itk kinase activity. We next compared the activities of both wild-type Itk and Itk(BtkSH3), by coexpressing S9 insect cells with the Itk substrate, PLCγ1, and either wild-type Itk or Itk(BtkSH3). Tyrosine phosphorylation of PLCγ1 on Tyr783 was more robustly induced by Itk(BtkSH3), than by wild-type Itk, suggesting enhanced kinase activity for Itk(BtkSH3) with that for wild-type Itk on its native substrate (Fig. 4B).

The in vitro data shown in Fig. 4 prompted us to introduce wild-type Itk and Itk(BtkSH3) into Itk−/− primary CD4+ T cells by retrovirus transduction to compare the signaling properties of the two Itk constructs. With this retrovirus vector, GFP is translated from the same mRNA transcript as Itk; thus Itk and Itk(BtkSH3) were expressed at comparable levels in Itk−/− cells, as indicated by GFP fluorescence (Fig. 5A). Previous studies have shown that optimal activation and phosphorylation of the p42/44 ERK MAPKs following T cell activation are dependent on Itk (8, 37). To compare wild-type Itk and Itk(BtkSH3) function in primary T cells, transduced T cells were stimulated by TCR engagement, and ERK phosphorylation was assessed. Itk−/− T cells expressing wild-type Itk were ~20% positive for ERK phosphorylation, in contrast to Itk−/− T cells transduced with the retrovirus vector alone (~10% phospho-ERK*). Expression of Itk(BtkSH3), further enhanced ERK activation, leading to ~30% phospho-ERK* cells. Average data from three experiments are shown in Fig. 5B. Of note, the levels of wild-type Itk and Itk(BtkSH3) in retrovirus-infected Itk−/− T cells are substantially lower than the normal level of endogenous Itk in wild-type T cells (Fig. 5C). Nonetheless, even this low level of Itk(BtkSH3) induces ERK phosphorylation that is similar if not greater than that seen in wild-type T cells following TCR stimulation (Fig. 5C). These data indicate that Itk(BtkSH3) is able to function in the TCR signaling pathway of primary T cells, and compared with wild-type Itk expressed at the same level, the nonclustering Itk mutant exhibits enhanced TCR signaling.

We also expressed wild-type Itk or Itk(BtkSH3) in Itk−/+ primary CD4+ T cells. According to our model, the active pool of Itk is the monomeric fraction, and further addition of wild-type Itk to the endogenous Itk supply should not significantly increase the pool of free monomer due to its self-association. In contrast, we expect that addition of Itk(BtkSH3) should increase the population of monomer supply because the Itk(BtkSH3) molecules are less likely to interact with themselves or with wild-type Itk. Indeed, expression of Itk(BtkSH3), in Itk−/+ T cells enhanced ERK phosphorylation following TCR stimulation compared with that in T cells transduced with the control retrovirus, whereas transduction of Itk−/+ T cells with wild-type Itk had no effect (Fig. 5D, 5E). These results indicate that Itk(BtkSH3) enhances Itk functional activity in the presence of wild-type Itk, suggesting that coexpression of the two forms of Itk leads to an increase in Itk monomers in T cells.

Given the increased activity of Itk(BtkSH3), measured in vitro (Fig. 4A), the higher levels of PLCγ1 phosphorylation induced by Itk(BtkSH3) compared with those induced by wild-type Itk (Fig. 4B), and the increase in phosphorylated Erk in primary T cells expressing Itk(BtkSH3) (Fig. 5A–E), we next monitored calcium flux in stimulated Itk−/− T cells expressing either wild-type Itk, Itk(BtkSH3), or vector alone (Fig. 5F). As can be seen, Itk−/− cells transduced with vector alone have a markedly impaired calcium response compared with that of Itk-sufficient wild-type CD4+ T cells; expression of wild-type Itk in Itk−/− cells partially restores this response. Importantly, Itk−/− T cells expressing Itk(BtkSH3) showed enhanced calcium flux compared with that of cells.
expressing wild-type Itk, consistent with the earlier observations of increased activity for the nonclustering Itk mutant. Efforts to assess IL-2 responses by the retrovirus-transduced Itk$^{+/+}$ T cells indicated a trend toward higher activity in cells expressing Itk (BtkSH3) versus that in cells expressing wild-type Itk, but the differences seen were not statistically significant. Nonetheless, these data overall provide further support for a model in which disruption of intermolecular Itk clustering leads to an increase in Itk-mediated signaling following TCR stimulation.

**Discussion**

Strength and duration of TCR signaling are important in regulating T cell development and lineage commitment. As a key component of the TCR signaling pathway, Itk has been found to modulate several aspects of thymocyte maturation, including positive selection, negative selection, and conventional versus innate CD8+ T cell development (11, 38). These observations suggest that Itk activity is involved in fine-tuning TCR signaling and that this, in turn, regulates T cell fate decisions. Further investigation of this process would be greatly aided by a mouse model in which Itk kinase activity is increased relative to that of wild-type Itk. Previous efforts to increase Itk activity by overexpression have uniformly been unsuccessful (36). The data presented in this report provide a biochemical explanation for these failures. Quite unlike the Src family kinases, there are to date no known mutations that activate Itk signaling. To our knowledge, Itk(BtkSH3) is the first Itk variant to show activity greater than that of the wild-type enzyme in T cells. Thus, in the future, a knock-in mouse expressing Itk(BtkSH3) in place of wild-type Itk should provide a valuable tool for studying the role of Itk in T cell development and function.

**FIGURE 5.** Itk(BtkSH3) exhibits enhanced signaling in T cells. A–C, Itk$^{-/-}$ primary CD4$^+$ T cells were infected with the retrovirus vector alone [mouse stem cell virus (MSCV), solid line], a retrovirus expressing wild-type Itk (Itk, long dashed line), or a retrovirus expressing Itk(BtkSH3) (short dashed line). GFP$^+$ cells were sorted on day 4 postinfection. A, Histograms of GFP fluorescence at the time of analysis are shown compared with noninfected control cells (filled histogram). Cells were stimulated with anti-CD3 Ab for 5 min, fixed, permeabilized, and stained with anti–phospho-ERK Ab. The percentages of phospho-ERK$^+$ cells are indicated on each histogram. The nonstimulated control shown is MSCV-infected cells. Nonstimulated controls for all cell lines are provided in Supplemental Fig. 1. B, The graph shows data from three experiments described in A. Bars indicate mean ± SD of the percentage of phospho-ERK$^+$ T cells. *p < 0.0005; **p < 0.0006; ***p < 0.004 (using a paired t test). C, Itk$^{-/-}$ T cells infected with vector alone (lane 3), retrovirus expressing wild-type Itk (lane 4), or retrovirus expressing Itk(BtkSH3) (lane 5) were stimulated with crosslinking anti-CD3 Ab for 5 min, lysed, and immunoblotted for Itk (top panel), phospho-ERK (middle panel), and total ERK (bottom panel). Lanes 1 and 2 represent primary Itk$^{+/+}$ CD4$^+$ T cells stimulated with anti-CD3 Ab for 0 and 5 min, respectively. D and E, Itk$^{+/+}$ primary CD4$^+$ T cells were infected with the retrovirus vector alone (MSCV, solid line), a retrovirus expressing wild-type Itk (Itk, long dashed line), or a retrovirus expressing Itk(BtkSH3) (short dashed line) and treated in a manner identical to the data shown in A and B. *p < 0.059; **p < 0.006; ***p < 0.048. F, Wild-type or Itk$^{-/-}$ CD4$^+$ T cells were infected with pMX-IRES-hCD8 vector alone or vector containing Itk constructs. Four days postinfection, human CD8$^+$ T cells were gated on and analyzed for calcium mobilization in response to anti-CD3 stimulation. Levels of wild-type Itk and Itk(BtkSH3) protein in the transduced Itk$^{-/-}$ T cells were comparable, as assessed by human CD8 staining (see Supplemental Fig. 1). Data are representative of three independent experiments with similar results.
of wild-type Itk could provide a system for examining T cell development and lineage commitment in cells with increased Itk kinase activity.

The quantitative effects of modulating Itk self-association on TCR signaling events are similar to those seen in studies of the Src family kinase, Lck. Src kinases, including Lck, are negatively-regulated by an intramolecular association between the SH2 domain and a phosphorylated tyrosine in the C-terminal tail that is absent in the Tec kinases. A comprehensive examination of Lck autoinhibition was recently reported in which this intramolecular interaction is either disrupted or strengthened by sequence changes close to the phosphotyrosine (39). Lck-deficient Jurkat T cells were reconstituted with wild-type Lck or Lck mutants, and TCR signaling leading to Erk activation was examined. Similar to our findings for wild-type Itk and Itk(BuSH3S), modulation of the Lck autoinhibitory interaction led to detectable but modest changes in Erk activation. Together these studies indicate that shifting the equilibrium between active and inactive kinase conformation alters the strength of TCR-mediated signaling but does so within a limited range of outcomes. This is in contrast to mutations located in active sites that have pronounced, all or nothing, effects on downstream signaling.

In the current study, the sequence changes that we have introduced into full-length Itk have been limited to the SH3/SH2 interface. As already mentioned, the isolated PH domain of Itk also forms intermolecular self-associated complexes (26) and might therefore play a significant role in Itk clustering in T cells. Itk membrane association occurs via PH domain interactions with phosphatidylinositol 3,4,5-trisphosphate in a T cell stimulation-dependent fashion (40–43). Localization of Itk at the membrane could favor PH/PH intermolecular interactions resulting in further stabilization of self-associated Itk complexes. Indeed, the contribution of the PH domain to Itk self-association may be minimal in systems, such as the NIH 3T3 cells used in this study for coimmunoprecipitation of self-associated Itk complexes. Indeed, the contribution of the PH domain to Itk self-association may be minimal in systems, such as the NIH 3T3 cells used in this study for coimmunoprecipitation experiments, but significantly more pronounced in the context of the T cell membrane environment. Hence, the modest functional effects observed upon altering the SH3/SH2 interface could be due to a dominant role for the PH domain that has not been affected in the Itk(BuSH3S) mutant. When more detailed structural insights into the PH/PH domain interface become available, this region of the Itk regulatory structure can be probed for specific sequence changes that disrupt PH domain self-association but are silent with respect to the membrane binding function of the PH domain. It is likely that such mutations, by themselves or in combination with mutations in the SH3/SH2 interface, may shift the equilibrium further away from the self-associated form of Itk, leading to further enhancement of Itk signaling in T cells.

Specific intermolecular self-association has been characterized for a number of protein systems (44). Within the kinase superfamily, this mechanism activates receptor kinases by promoting trans autophosphorylation both within and outside of the protein kinase domain (45). In another example of activation by intermolecular association, the antiviral protein kinase PKR dimerizes via phosphotyrosine-dependent binding to dsRNA (46, 47). Alternatively, inhibition by dimerization occurs for the receptor-like protein tyrosine phosphatase-α where an inhibitory “wedge” on one molecule inserts into the catalytic site of another molecule (48). A crystal structure of the kinase domain from yeast Snf1 also reveals a dimeric arrangement that putatively impedes catalytic activity by steric means (49). Likewise, a crystal structure of Ca<sup>2+</sup>/calmodulin-dependent protein kinase II reveals a regulatory segment that sterically blocks substrate binding to the catalytic site (50). The Ca<sup>2+</sup>/calmodulin-dependent protein kinase II kinase domain itself is intrinsically active, and dimer formation brings the regulatory segment into position to inhibit activity. 

In light of these examples, a simple mechanistic model for Itk autoinhibition by self-association would invoke steric blockade of the catalytic site. However, it has been well documented that the isolated Itk kinase domain exhibits little or no catalytic activity (28–30), suggesting the possibility of an alternative autoinhibition mechanism. We have previously reported a kinetic analysis of a series of Itk fragments and have shown that the SH2 domain and linker between SH2 and kinase domains are required to achieve wild-type levels of activity (28). Like the Csk kinase (51), the SH2 and SH2–kinase linker region make direct contact with the kinase domain to stabilize the active conformation of Itk. Thus, inhibition of Itk catalytic activity by SH3/SH2-mediated self-association could be explained if the intermolecular interactions between Itk regulatory domains (particularly those involving the SH2 domain) compete with activating interactions between the SH2 and kinase domains.

T cell activation itself produces signals that compete with the self-association equilibrium and would shift the population of Itk in the cell toward a monomeric state. Specifically, exogenous binding partners, such as transiently produced phospholipids, would compete directly with Itk self-association by interfering with the SH2/SH3 interaction interface (9, 18, 22, 36, 52, 53). If self-association is disfavored, then the regulatory domains could then adopt the catalytically competent conformation (28). Conversely, in the absence of such activating factors, Itk might remain self-associated and autoinhibited.

It is also possible to envision a role for Itk autoinhibition following activation of Itk by TCR engagement; in this case, self-association would be one mechanism that could terminate Itk signaling (in addition to the activity of phosphatases). In binding studies of the Itk SH3 and SH2 domains, we found that Tyr<sup>180</sup> phosphorylation in the SH3 domain enhances affinity for the Itk SH2 domain (54). Together with the observations that autophosphorylation at Tyr<sup>180</sup> in the SH3 domain has no effect on Itk kinase activity and occurs in cis (54), it is reasonable to suggest that autophosphorylation may be a first step toward turning off Itk-mediated signaling by promoting intermolecular self-association leading to a drop in kinase activity. Indeed, a previous report suggests that Itk clustering in T cells occurs following membrane association (27).

Although the precise mechanistic details of when and how intermolecular association of Itk modulates its activity remain a question, this mode of autoinhibition for a nonreceptor tyrosine kinase suggests that reagents (or mutations) that shift the equilibrium toward the self-associated state could dampen T cell activation while disfavoring self-association will increase Itk activity and concomitant T cell activation.

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References


