Integrin receptor signals are costimulatory for mitogenesis with the T-cell receptor during T-cell activation. A subset of integrin receptors can link to the adapter protein Shc and provide a mitogenic stimulus. Using a combination of genetic and pharmacological approaches, we show herein that integrin signaling to Shc in T cells requires the receptor tyrosine phosphatase CD45, the Src family kinase member Lck, and protein kinase C. Our results suggest a model in which integrin-dependent serine phosphorylation of Lck is the critical step that determines the efficiency of Shc tyrosine phosphorylation in T cells. Serine phosphorylation of Lck is dependent on PKC and is also linked to CD45 dephosphorylation. Mutants of Lck that cannot be phosphorylated on the critical serine residues do not signal efficiently to Shc and have greatly reduced kinase activity. This signaling from integrins to Lck may be an important step in the costimulation with the T-cell receptor during lymphocyte activation.
with the catalytic domain and surrounding amino acids (Sicheri et al., 1997; Xu et al., 1997, 1999). There are multiple potential ways to disrupt this intramolecular interaction and activate Src kinases, one of which is to dephosphorylate the negative regulatory tyrosine residue located near the carboxy terminus, which is bound by the SH2 domain (Thomas and Brugge, 1997). Because Fyn is activated by integrin ligand binding or antibody cross-linking in adherent cells, we reasoned that integrin ligation could either activate a tyrosine phosphatase or make it accessible to Fyn. This putative tyrosine phosphatase would most likely be membrane bound. In adherent cells, there are data to suggest that the receptor protein-tyrosine phosphatase α and the cytoplasmic tyrosine phosphatase SHP-2 can regulate integrin-dependent Src family kinase activation (Oh et al., 1999; Su et al., 1999).

In hematopoietic cells, one candidate for such a membrane tyrosine phosphatase that can potentially activate Src family members kinase activity is CD45, a leukocyte cell-specific transmembrane glycoprotein with a tandem repeat of protein tyrosine phosphatase domains in its cytoplasmic region (Ralph et al., 1987). The function of CD45 in antigen receptor signaling has been studied extensively. It regulates pre-TCR and TCR complex-mediated signal transduction during T-cell development (Byth et al., 1996). CD45-deficient human T-cell lines are defective in their ability to respond to signals via their TCR-CD3 complex (Koretzky et al., 1991; Shirroz et al., 1992), and this signaling can be restored by reconstitution of functional CD45 (Koretzky et al., 1990, 1992; Desai et al., 1993; Hovis et al., 1993; Volarevic et al., 1993). CD45 has been shown to regulate both the Src family kinases Lck and Fyn. For example, CD45 has been found to be colocalized selectively with Fyn in functional human T lymphocytes. It can dephosphorylate the negative regulatory tyrosine within the carboxy terminus of both Fyn and Lck. In a CD45-deficient T-lymocyte clone (L3), tyrosine phosphorylation of a Fyn C-terminal peptide, which contains the negative regulatory tyrosine, is increased twofold (Cahir McFarland et al., 1993).

Although the role of Fyn in integrin-Shc signaling in adherent fibroblasts is well established, it remains unclear whether Fyn is the only Src family kinase that participates in this signaling process in T cells. In addition to Fyn, T cells express another Src family kinase, Lck. Fyn and Lck share striking overall sequence similarity. Both proteins are myristylated and palmitoylated and are expressed abundantly in mature T cells. Both kinases are important in TCR α/β/CD3-mediated signal transduction in mature T cells, in which Shc tyrosine phosphorylation is also important (Weiss and Littman, 1994; Anderson and Perlmuter, 1995). Furthermore, it has demonstrated that Lck/CD4 cross-linking can result in the phosphorylation of Shc Tyr 317 in a murine T-cell line, which is the same position kinased by Fyn in fibroblasts, and represents a binding site for the SH2 domain of Grb2 (Wary et al., 1996, 1998; Walk et al., 1998). We therefore wondered whether Fyn and Lck could play redundant roles in integrin-Shc signaling in T cells.

In the present study, we have taken the advantage of availability of CD45-deficient and CD45-reconstituted Jurkat cell lines and tested the hypothesis that CD45, or a CD45-like transmembrane tyrosine phosphatase, depending on the cell type, participates in integrin-Shc signaling. We found that CD45 was indeed required for β1 integrin-mediated Shc tyrosine phosphorylation in Jurkat cells. In addition, we have investigated whether Lck takes a part in this signaling event. We found that CD45 regulates Lck kinase activity in Jurkat cells and that Lck kinase activity is required for integrin-mediated Shc phosphorylation. Integrin clustering led to serine phosphorylation of Lck, and this modification is protein kinase C (PKC) dependent. In addition, efficient phosphorylation of Shc in response to integrin clustering in T cells also requires PKC activity. Most strikingly, mutants of Lck with alanine substitution of critical serine residues were profoundly deficient in Shc phosphorylation and kinase activity, showing that these sites are essential for integrin-Shc signaling in T cells.

MATERIALS AND METHODS

Antibodies and Cell Lines

Monoclonal antibodies (mAbs) 4B4 (anti-β1) and 3A5 (anti-Lck) were purchased from Beckman Coulter (Fullerton, CA) and Santa Cruz Biotechnology (Santa Cruz, CA). Cells producing mAb GAP8.3 (anti-CD45) were from American Type Culture Collection (Manassas, VA). Polyclonal anti-human focal adhesion kinase (FAK) and Shc antibodies were from Upstate Biotechnology (Lake Placid, NY). RC-20-H (peroxidase-conjugated recombinant anti-P-Tyr PY 20) was from Transduction Laboratories (Lexington, KY). Polyclonal anti-phospho-Src family Tyr 416 antibody was purchased from Cell Signaling Technology (Beverly, MA).

Leukemic T-cell lines Jurkat, J45.01 (CD45-deficient Jurkat), and Jcam 1.6 (Lck-deficient Jurkat) were obtained from American Type Culture Collection. Dr. G. Koretzky (University of Pennsylvania, Philadelphia, PA) provided J45/CH11 (transfectant of J45.01 that express A2/CD45 chimeric molecule) and J45LB3 (transfectant of J45.01 that express full-length CD45). Jurkat, J45.01, and Jcam 1.6 were routinely maintained in RPMI supplemented with 10% fetal calf serum, penicillin/streptomycin (100 U/ml). J5/CH11 and J45LB3 were maintained in above-mentioned medium supplemented additionally with G418 (2 mg/ml).

A G418-resistant Jcam1 cell line, which expresses a VP16-tetracycline repressor fusion protein, was provided by Dr. D. Straus (Virginia Commonwealth University, Richmond, VA) (Denny et al., 2000). The Lck S42A, S59A, S42A/S59A, S42E, S59E, and S42E/S59E mutants were generated from wild-type (Wt) mouse Lck cDNA by using the QuikChange site-directed mutagenesis kit (Stratagene, La Jolla, CA) and were confirmed by sequencing. These cDNAs were subcloned into the pBP1 vector that contains a cytomegalovirus promoter sequence regulated by tetracycline operators, and a gene conferring resistance to the antibiotic hygromycin (Denny et al., 2000). These vectors were transfected into Jcam1/tetra-VP16 cells by electroporation (250 V and 960 μF) using by a Bio-Rad (Hercules, CA) apparatus. Stable cell lines were selected using hygromycin B (100 μg/ml) and the cytoskeletal associated polystyrene latex beads (2.4 μm in diameter; Interfacial Dynamics, Portland, OR) with 100 μg of 4B4 in 300 μl of conjugation buffer (30 mM Na2CO3, 70 mM NaHCO3, pH 9.5) for 1.5 h at room tempera-
ture. At the end of cross-linking, cells were extracted on ice for 30 min with 1 ml of lysis buffer (1% Triton X-100, 50 mM HEPES, 150 mM NaCl, 1 mM EDTA, 10 mM NaF, 1 mM pyrophosphate, and 2 mM Na3VO4, pH 7.5) supplemented with 10 μl/ml mammalian cell proteinase inhibitor cocktail (Sigma-Aldrich, St. Louis, MO).

For immunoprecipitation and immunoblotting of Shc, Lck, and FAK, total cell extracts were incubated in lysis buffer with 50 μl of GammaBind G Sepharose (Amersham Biosciences, Piscataway, NJ) and either 10 μg of polyclonal anti-human Shc or FAK or 5 μg of monoclonal anti-human Lck for 2 h at 4°C. Samples were separated by SDS-PAGE (10%) and transferred to nitrocellulose membrane. The blots were blocked with 5% nonfat dry milk (for Shc, Lck, or FAK antibodies) or 5% bovine serum albumin (for RC-20). Nitrocellulose-bound antibodies were detected by chemiluminescence with enhanced chemiluminescence (ECL) (Pierce Chemical, Rockford, IL).

For immune complex autokinase assays, complexes were recovered from extracts prepared with a modified radioimmunoprecipitation assay buffer 1 (1% Triton X-100, 0.5% sodium deoxycholate, 0.1% SDS, 50 mM HEPES pH 7.4, 150 mM NaCl, 1.5 mM MgCl2, 1 mM EGTA, and 10% glycerol). These immune complexes were washed three times with buffer 1, twice with buffer 2 (0.1% Triton X-100, 50 mM HEPES pH 7.4, 150 mM NaCl, and 10% glycerol), and twice with buffer 3 (20 mM Tris pH 7.2, 100 mM NaCl, and 10 mM MgCl2). After the washes, immune complexes were incubated in 30 μl of buffer 3 supplemented with 20 μM cold ATP and 10 μCi of [γ32P]ATP (PerkinElmer Life Sciences, Boston, MA) for 2 min at 30°C. Washing the immune complexes with buffer 3 twice, followed by boiling in SDS-PAGE sample buffer stopped the reaction. After SDS-PAGE, gels were fixed (50% methanol, 10% acetic acid for 30 min, and then 10% methanol, 10% acetic acid for 30 min) and then incubated in 1 M KOH for 2 h at 56°C to hydrolyze phosphates on serine/threonine residues. Finally, gels were rinsed in 10% acetic acid and 10% methanol for 20 min before being dried for autoradiography. Quantification of scanned films was performed using NIH Image software.

For CD45 immune complex phosphatase assays, 107 Jurkat cells were lysed in 1 ml of 1% Triton, 50 mM HEPES pH 7.5, 150 mM NaCl, and 2 mM diethiothreitol for 0.5 h at 4°C. The total cell lysate was then incubated in the lysis buffer with 50 μl of goat anti-mouse Sepharose and 250 μl of supernatant of GAP 8.3 culture medium for 2 h at 4°C. At the end of incubation, Sepharose was washed four times with lysis buffer followed by another four times wash with phosphatase assay buffer (0.1 M sodium acetate, 0.2% Triton X-100, 1 mM EDTA, pH 6.0). The beads were incubated in 20 μl of assay buffer with 5 μl of a phosphotyrosine peptide provided in a tyrosine phosphatase assay kit (Upstate Biotechnology) for 10 min. The assay was stopped by addition of 100 μl of Malachite Green solution and free phosphate was measured by absorbance reading at 620 nm.

**Flow Cytometry**

Fluorescence flow cytometric analysis was performed on a FACScan fluorocytometer (BD Biosciences, San Jose, CA). Harvested cells were washed three times with PCN (phosphate-buffered saline, 0.5% calf serum, and 0.5% NaN3). Cells were stained with primary antibodies on ice for 30 min, followed by three washes with cold PCN. Cells were then resuspended and stained with fluorescein isothiocyanate-conjugated secondary antibodies in PCN for 30 min on ice. After three washes with PCN, cells were fixed in phosphate-buffered saline plus 1% paraformaldehyde and analyzed.

**RESULTS**

**A Requirement for CD45 in Integrin Shc Signaling in Jurkat Cells**

To address the role of CD45 in β1 integrin-mediated Shc tyrosine phosphorylation, we compared the integrin signal-
cells (J45.01). Cells (10^7) for each cell line were either clustered with Wild-type Jurkat cells were compared with CD45-dependent Jurkat deficient cell line J45.01. (B) FAK phosphorylation in Jurkat cells. dependent phosphorylation of Shc in Jurkat, but in the CD45-deficient cell line J45.01. Cells (10^7) for each cell line were either clustered with anti-β1 antibody beads (+) or left unclustered (−) and incubated for 15 min at 37°C followed by extraction. Extracts were immunoprecipitated using rabbit anti-Shc antibodies and separated by SDS-PAGE. After transfer to nitrocellulose, blots were stained for phosphotyrosine by using peroxidase-conjugated anti-P-Tyr (RC-20) and visualized by ECL. Blots were then stripped and reprobed with rabbit anti-FAK antibodies and peroxidase-conjugated protein A. The band below, which is present in all lanes, is related to the polyclonal anti-Shc, because it remains present even in the absence of cell extracts (our unpublished data). Note the cross-linking-dependent phosphorylation of Shc in Jurkat, but in the CD45-deficient cell line J45.01. (B) FAK phosphorylation in Jurkat cells. Wild-type Jurkat cells were compared with CD45-dependent Jurkat cells (J45.01). Cells (10^7) for each cell line were either clustered with anti-β1 antibody beads (+) or left unclustered (−) and incubated for 15 min at 37°C followed by extraction. Extracts were immunoprecipitated using rabbit anti-Shc antibodies and separated by SDS-PAGE. After transfer to nitrocellulose, blots were stained for phosphotyrosine by using peroxidase-conjugated anti-P-Tyr (RC-20) and visualized by ECL. Blots were then stripped and reprobed with rabbit anti-FAK and peroxidase-conjugated protein A. Note the cross-linking-dependent phosphorylation of FAK in both cell lines, demonstrating that FAK phosphorylation does not require CD45 and that β1 cross-linking in J45.01 is effective. (C) Integrin-dependent Shc phosphorylation: requirement of the presence of CD45 phosphatase. CD45 cDNA (J45LB3) or a chimera consisting of the extracellular domain of HLA A2 connected to the intracellular domain of CD45 (J45CH11) was expressed in previously CD45-deficient J45.01 cells. Cells (10^7) for each line were either clustered with anti-β1 antibody beads (+) or left unclustered (−) and incubated for 15 min at 37°C followed by extraction. Extracts were immunoprecipitated using rabbit anti-Shc antibodies and separated by SDS-PAGE. After transfer to nitrocellulose, blots were stained for phosphotyrosine by using peroxidase-conjugated anti-P-Tyr (RC-20) and visualized by ECL. Blots were then stripped and reprobed with rabbit anti-Shc and peroxidase-conjugated protein A. Note the cross-linking–dependent phosphorylation of Shc is restored with both CD45 (J45LB3) and with the chimera molecule (J45CH11).

**Figure 1.** Integrin-dependent Shc phosphorylation in Jurkat cells: requirement for CD45. (A) Wild-type Jurkat cells were compared with CD45-deficient Jurkat cells (J45.01). Cells (10^7) for each cell line were either clustered with anti-β1 antibody beads (+) or left unclustered (−) and incubated for 15 min at 37°C followed by extraction. Extracts were immunoprecipitated using rabbit anti-Shc antibodies and separated by SDS-PAGE. After transfer to nitrocellulose, blots were stained for phosphotyrosine by using peroxidase-conjugated anti-P-Tyr (RC-20) and visualized by ECL. Blots were then stripped and reprobed with rabbit anti-FAK and peroxidase-conjugated protein A. Note the cross-linking–dependent phosphorylation of FAK in both cell lines, demonstrating that FAK phosphorylation does not require CD45 and that β1 cross-linking in J45.01 is effective. (C) Integrin-dependent Shc phosphorylation: requirement of the presence of CD45 phosphatase. CD45 cDNA (J45LB3) or a chimera consisting of the extracellular domain of HLA A2 connected to the intracellular domain of CD45 (J45CH11) was expressed in previously CD45-deficient J45.01 cells. Cells (10^7) for each line were either clustered with anti-β1 antibody beads (+) or left unclustered (−) and incubated for 15 min at 37°C followed by extraction. Extracts were immunoprecipitated using rabbit anti-Shc antibodies and separated by SDS-PAGE. After transfer to nitrocellulose, blots were stained for phosphotyrosine by using peroxidase-conjugated anti-P-Tyr (RC-20) and visualized by ECL. Blots were then stripped and reprobed with rabbit anti-Shc and peroxidase-conjugated protein A. Note the cross-linking–dependent phosphorylation of Shc is restored with both CD45 (J45LB3) and with the chimera molecule (J45CH11).
observed (our unpublished data). Furthermore, when we using immune complex assays. No obvious difference was integrin cross-linked and noncross-linked Jurkat cells by

strates the requirement of Lck for

show that Shc phosphorylation is indeed defective in Jcam 1.6 cells after antibody-mediated integrin cross-linking.

Effect of β1 Integrins Clustering on Fyn and Lck Kinase Activity

We next decided to investigate whether the clustering of integrins could regulate the activity of the Src family kinases Fyn or Lck in Jurkat cells. Others (Wary et al., 1998) have shown that kinase activity of Fyn was increased in fibroblasts after antibody-mediated β1 integrin cross-linking. Therefore, we determined the kinase activity of Fyn from β1 integrin cross-linked and noncross-linked Jurkat cells by using immune complex assays. No obvious difference was observed (our unpublished data). Furthermore, when we tested the kinase activity of Lck from β1 integrin-clustered and -unclustered Jurkat cells, again, we did not see any significant change (Figure 4A). This result may be due to the high level of Lck kinase activity observed before cross-linking. In fibroblasts, using anti-β1 antibody beads, we did see a significant activation of Fyn kinase activity by using the same assay conditions as shown above (Figure 4C). However, these cells have a much lower level of kinase activity before clustering. As a control for loading the same amount of enzyme in the autokinase assays, we split the immunoprecipitates and probed Western blots with Lck antibody.

Change in Gel Mobility of Lck Induced by β1 Integrin Cross-Linking

It has been reported that serine phosphorylation of the N-terminal unique region of Lck and the accompanying gel retardation (from 56–63 kDa) can occur upon treatment of T cells with phorbol ester, interleukin 2, and CD4/TCR cross-linking agents (Casnellie and Lamberts, 1986; Veillette et al., 1988; Einspahr et al., 1990; Horak et al., 1991). Ser 42 and Ser 59 in the N-terminal unique region of Lck have been identified as the major phorbol ester-induced phosphorylation sites. Phosphorylation of Ser 59 is responsible for the shift from 56 to 61 kDa, whereas phosphorylation of Ser 42 is required for the shift from 61 to 63 kDa. It has been found in vitro that mitogen-activated kinase and PKC or protein kinase A (PKA) could be responsible for the phosphorylation of Ser 59 and Ser 42, respectively (Winkler et al., 1993).
We decided to investigate this phosphorylation and gel retardation of Lck, induced upon integrin clustering, to determine whether this process played a role in the integrin-Shc signaling pathway. Our first step was to determine whether this shift occurs both with antibody-induced clustering and with authentic integrin ligands. Because flow cytometry measurements (our unpublished data) had determined that one of the β1 integrin heterodimers on the surface of Jurkat cells is α1β1, we decided to use antibodies and ligand for this integrin. To this end, we compared the electrophoretic mobility of Lck in untreated Jurkat cells with those treated with beads coated with TS2/7, an anti-human integrin α1 antibody, and with cells treated with beads coated with CB3, the cell binding fragment of collagen IV, a ligand for α1β1 (Kern et al., 1994). As shown in Figure 5, A and B, both treatments resulted in a mobility shift of a significant portion of the Lck. We next hypothesized that these phosphorylations may be relevant to Lck regulation if they were dependent on the presence of active CD45, because the carboxyl-terminal phosphotyrosine intramolecularly to the SH2 domain might prevent access to the amino terminus, where the modifications of Lck are likely to be occurring. Therefore, we clustered integrins on CD45 null J45.01 cells or on Jurkat cells. Strikingly, as shown in Figure 5C, Jurkat cells are able to generate the electrophoretic mobility shift upon integrin clustering, whereas the CD45 null cells cannot, which indicates that this shift requires CD45. These results suggest that the cleavage of the carboxyl-terminal P-Tyr is linked to the serine phosphorylation of the amino terminus.

We next determined which protein kinase inhibitors could block this process. Figure 6, A–C, shows that neither mitogen-activated protein kinase kinase (MEK) nor PKA inhibitors could prevent the change in electrophoretic mobility despite the fact that these two classes of kinases had been implicated in vitro in this mobility shift (Winkler et al., 1993). In contrast, PKC inhibitors, such as bisindolylmaleimide (Figure 6C) or staurosporine (our unpublished data), completely abolished the shift induced by integrin clustering. Thus, integrin-dependent modification of Lck requires PKC activity.

Finally, we determined whether PKC activity was required for integrin-Shc signaling in T cells. This hypothesis was tested by clustering Jurkat cells with anti-β1 integrin antibody either in the absence or presence of the PKC inhibitor bisindolylmaleimide. Strikingly, as shown in Figure 7A, inhibition of PKC activity caused a dramatic reduction in the phosphorylation of Shc. Quantitation of this experiment showed that this was a fourfold reduction. We confirmed...
this result by down-regulating PKC in Jurkat cells with a high-dose overnight treatment with phorbol ester, followed by clustering Jurkat cells with anti-β1 integrin antibody and assaying Shc phosphorylation (Figure 7B). This treatment also led to a dramatic reduction in integrin-dependent Shc phosphorylation. Thus, PKC activity is required for both the electrophoretic mobility shift of Lck and for the efficient tyrosine phosphorylation of Shc in T cells.

**Mutants of Lck**

To test directly the role of phosphorylation of Ser 42 and Ser 59 of the Lck unique amino-terminal domain, we have made mutants of Lck and expressed them in Jcam1.6 cells, which lack Lck activity. Single S→A and S→E mutants were made along with double mutants. These mutants and the wild-type Lck were then expressed in the Lck-deficient cells. Multiple clones were isolated and the analysis was done...
with lines expressing similar levels of Lck protein (Figure 8A). Mutation of S42A, S42E, S59A, or S59E had no effect on integrin-induced Shc phosphorylation. Strikingly, the double mutant S42A, S59A is profoundly defective in its ability to mediate integrin-mediated Shc signaling (Figure 8B). These results demonstrate that these two serine residues are required for integrin-mediated Shc phosphorylation. In contrast, when these residues were both mutated to glutamic acid, integrin-dependent Shc phosphorylation was normal (Figure 8B).

Because we have shown that PKC signals are required for Shc phosphorylation in these cells, one would predict that the presence of negative charges at these two serines should mimic phosphorylation and may render this form of Lck resistant to PKC inhibition. Therefore, we compared the effect of the PKC inhibitor on Shc phosphorylation induced in cells expressing wild-type Lck vs. S42E S59E Lck. The results (Figure 8C) show that the presence of negative charges at these two sites does indeed provide substantial resistance (>90%) to PKC inhibitors.

Because the double mutant S42A, S59A is defective in its ability to mediate integrin-mediated Shc signaling, we decided to determine its kinase activity. Using both autokinase assays and Western blotting for the presence of the activation loop phospho-tyrosine, we find that this mutant has dramatically reduced kinase activity in these cells (Figure 9). Thus, the amino terminal unique domain of Lck seems to be involved in the regulation of kinase activity.

**DISCUSSION**

The results shown in this article demonstrate a requirement for the receptor tyrosine phosphatase CD45 in the signaling of integrin to Shc. We have shown that the lack of CD45 leads to decreased Lck kinase activity, and restoration of the phosphatase activity of CD45 leads to increased Lck kinase activity. CD45 is known to be essential for TCR antigen-dependent signaling, both in cell lines and in vivo (Neel, 1997). Others have shown that the kinase activity of Lck is increased after expression of CD45 in L3 M93 T-cell clone (Cahir McFarland et al., 1993), in peripheral blood lymphocytes (Mustelin et al., 1989), and HPB.MLT, a human CD4+8+ leukemic T-cell line (Mustelin and Altman, 1990). In Jurkat cells, CD45 expression was shown to modulate the binding of Lck to an 11-amino acid tyrosine-phosphorylated peptide containing the carboxy terminus of Lck, suggesting that CD45 could positively regulate Lck kinase activity (Sieh et al., 1993).

Our data demonstrate a requirement of CD45 for integrin signaling in T cells. In macrophages, CD45 has been shown to colocalize with integrins in focal adhesions, along with Src family kinases, and to be required for maximal adhesive activity (Roach et al., 1997). However, in this cell type, the lack of CD45 did not increase tyrosine kinase activity of Hck and Lyn, due to persistent hyperphosphorylation of the regulatory tyrosines located within the activation loop of the kinase domain. Thus, the exact role of CD45 in integrin receptor function depends on which Src family kinase members are required for these pathways in a given cell type, and how they may be activated.

In our case, we have shown that Lck activity was regulated by CD45. Furthermore, Lck-deficient cells also do not phosphorylate Shc in response to integrin clustering. Thus, we have shown that Lck is required, and at least in Jurkat and Jcam cells, Fyn does not substitute. The expression level of Fyn in these cells is very low (Denny et al., 2000), and this low expression level likely accounts for the inability of Fyn to compensate. Further experiments will be required to dissect the relative roles of these two Src family kinase members in integrin-Shc signaling in vivo. The requirement of Lck in Jurkat cells strongly suggests that many Src family members could mediate integrin-Shc signaling in different cell types. Perhaps the most prominent exception is c-Src itself, which would not substitute for Fyn in fibroblasts (Wary et al., 1998). This result may be due to differences in lipid modification, because Src associates to the membrane via a myristyl and a basic motif, whereas both Lck and Fyn share a similar lipid modification; they are myristylated and
palmitylated (Resh, 1994). However, differences within cell types may also include the role of caveolin 1 vs. lymphocyte-specific proteins within lipid microdomains of the membrane, because both Lck and Fyn are associated with such domains (Brown and London, 1998).

A model has been proposed for the regulation of receptor tyrosine phosphatases, which suggests that oligomerization of these molecules is the latent state, whereas dissociation leads to activation (Weiss and Schlessinger, 1998). Thus, it is possible that integrins, for example, could bind CD45 and cause dissociation of dimers, which would activate the phosphatase domain, leading to Lck activation. We attempted to test this idea by measuring CD45 phosphatase activity in vitro after immunoprecipitation from extracts of cells untreated or treated with integrin clustering and did not find an increase in phosphatase activity. It is possible that immunoprecipitation of CD45 leads to its activation, so that we cannot measure potential increases in activity via this assay. Thus, we cannot definitively rule out a role for integrins in the regulation of CD45 activity. However, at this point, we

Figure 8. Serine 42 and serine 59 of the Lck N-terminal unique region are critical for integrin-dependent Shc phosphorylation. (A) Determination of the expression levels of Wt Lck and single and double mutants of serine 42 and serine 59 to alanine (S42A, S59A and S42A/S59A) or glutamic acid (S42E, S59E and S42E/S59E) of Lck in Jcam 1.6 cells. Equal amounts of protein from the extracts of these stably transfected Jcam 1.6 cell lines were immunoprecipitated with monoclonal anti-Lck antibody. The resulting immune complexes were separated by SDS-PAGE. After transfer to nitrocellulose, the blot was stained with monoclonal Lck antibodies and peroxidase-conjugated anti-light chain antibodies, and visualized by ECL. Similar levels of Lck are expressed in all of the lines used. (B) Integrin-dependent Shc phosphorylation requires both the Ser 42 or Ser 59 sites in the Lck N-terminal unique region. The cell lines shown above were either left unclustered (−) or clustered with antibody beads (+) and incubated for 15 min at 37°C followed by extraction. Equal amounts of protein from the extracts were immunoprecipitated with rabbit anti-Shc antibody and separated by SDS-PAGE. After transfer to nitrocellulose, the blot was stained for phosphotyrosine by using peroxidase-conjugated anti-p-Tyr (RC-20) and visualized by ECL. Blots were then stripped and reprobed with rabbit anti-Shc antibodies and peroxidase-conjugated protein A. Note the markedly reduced Shc phosphorylation in S42A/S59A double mutant Lck Jcam 1.6 cell line upon integrin activation, whereas the S42E/S59E mutant had normal amounts of Shc phosphorylation. (C) S42E/S59E Lck is resistant to inhibition of PKC. Wt Lck and S42E/S59E Lck-transfected Jcam1.6 cell lines were incubated with 5 μM bisindolylmaleimide, a specific inhibitor of PKCa, βI, βII, and γ subtypes, for 1 h at 37°C or without bisindolylmaleimide. The cells were then clustered with antibody beads (+) and incubated for 15 min at 37°C followed by extraction. Equal amounts of protein from the extracts were immunoprecipitated with rabbit anti-Shc antibody and separated by SDS-PAGE. After transfer to nitrocellulose, the blot was stained for phosphotyrosine by using peroxidase-conjugated anti-p-Tyr (RC-20) and visualized by ECL. Blots were then stripped and reprobed with rabbit anti-Shc antibody and peroxidase-conjugated protein A. Note that the S42E S59E form of Lck produced high levels (>90% of untreated) of Shc despite pretreatment with a PKC inhibitor, whereas the Wt shows a fivefold reduction (as determined by NIH Image software) in Shc phosphorylation with the same treatment.
have no evidence that CD45 activity is regulated by integrins.

The notion that Lck participates in integrin signaling raises the question of how this kinase is regulated by integrins. Our findings that cross-linking could result in a slower migration of Lck from 56 kDa to either 61 or 63 kDa may have shed some light on this problem. Based on studies on various T cells, this mobility shifting is due to phosphorylation of serine residues in the N-terminal unique region of Lck. Because these serine residues lie very close to the SH2 domain of Lck in the three-dimensional structure, the phosphorylation state of these residues is believed to be critical in determining the binding affinity and specificity of the SH2 domain of the protein (Joung et al., 1995; Park et al., 1995).

For example, it has been found that a novel 62-kDa protein competes with phosphotyrosine for binding to SH2 domain of Lck when the serine 59 in the N-terminal region of Lck is phosphorylated (Park et al., 1995). The phosphorylation of these serine residues could also have an impact on the binding of potential inhibitors to Lck, or in the binding of the SH2 domain to the carboxy-terminal phosphotyrosine. In vitro, several different protein kinases, PKC/PKA, or mitogen-activated protein (MAP) kinase has been implicated in the shifting from 59 to 61 kDa and from 59 to 63 kDa, respectively (Winkler et al., 1993). Although our data show that the phosphorylation is dependent on PKC, we do not know whether this is the only protein kinase involved. Serine 59 is most likely phosphorylated by a proline-dependent kinase, consistent with the surrounding sequence within Lck (Winkler et al., 1993). In vitro, MAP kinase can mediate this phosphorylation; however, the lack of an effect by the MEK inhibitor, shown in Figure 6, suggests that either another proline-dependent kinase is involved or that a MEK-independent MAP kinase activation is involved. Because another group has shown that a PKA-dependent serine phosphorylation of Src is critical in its regulation (Schmitt and Stork, 2002), it is possible that these modifications of the unique amino-terminal domain of Src family kinases are a common, general mechanism for regulation. Further experiments will be necessary to determine the mechanism of this profound effect (8- to 10-fold reduction in kinase activity of the S42AS59A mutant).

Our results suggest a working model for integrin-dependent phosphorylation of Shc in T cells (Figure 10). Upon integrin clustering, there is a PKC-dependent phosphorylation of Lck, which may alter its binding to a number of partners, allowing for a portion of it to bind and phosphorylate Shc. This phosphorylation is dependent on the previous cleavage of the COOH terminal pTyr by CD45 (Figure 5C). Although PKC activation is clearly required, it is likely that clustering is a critical feature, because the phospho-mimic mutant (S42ES59E) does not phosphorylate Shc in the absence of cross-linking (Figure 8B). Most likely, Lck would bind to Shc through its SH3 domain, as has been shown for Fyn and Shc (Wary et al., 1998). Lck, like Fyn, then phosphorylates Shc on tyrosine 317, which is a potent Grb2 binding site. Interestingly, others (Miranti et al., 1999) have shown that PKC is required for optimal integrin-Shc signaling in Cos cells, where Fyn is the most likely kinase. The experiments performed in this study were done using fibronectin adhesion to trigger signaling, and perturbation of PKC affected cell spreading as well as Shc signals, although FAK signaling was spared. Thus, the exact basis for PKC requirement in Cos cells could be similar as we propose for Lck in Jurkat cells, but more experiments will be necessary to determine whether this is the case.

Future directions for this work will hopefully answer a number of important questions that remain open. The mechanism of integrin-dependent PKC activation remains unclear but may involve PLCγ isoforms binding to FAK. In addition, it is unclear which PKC isoforms are involved; however, it is intriguing to speculate that PKCζ, which is a central player in T-cell signaling at the immunological synapse (Isakov and Altman, 2002), may be involved.
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REFERENCES


