Impaired Integrin-mediated Adhesion and Signaling in Fibroblasts Expressing a Dominant-negative Mutant PTP1B

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Abstract. To investigate the role of nonreceptor protein tyrosine phosphatase 1B (PTP1B) in β1-integrin–mediated adhesion and signaling, we transfected mouse L cells with normal and catalytically inactive forms of the phosphatase. Parental cells and cells expressing the wild-type or mutant PTP1B were assayed for (a) adhesion, (b) spreading, (c) presence of focal adhesions and stress fibers, and (d) tyrosine phosphorylation. Parental cells and cells expressing wild-type PTP1B show similar morphology, are able to attach and spread on fibronectin, and form focal adhesions and stress fibers. In contrast, cells expressing the inactive PTP1B have a spindle-shaped morphology, reduced adhesion and spreading on fibronectin, and almost a complete absence of focal adhesions and stress fibers. Attachment to fibronectin induces tyrosine phosphorylation of focal adhesion kinase (FAK) and paxillin in parental cells and cells transfected with the wild-type PTP1B, while in cells transfected with the mutant PTP1B, such induction is not observed. Additionally, in cells expressing the mutant PTP1B, tyrosine phosphorylation of Src is enhanced and activity is reduced. Lysophosphatidic acid temporarily reverses the effects of the mutant PTP1B, suggesting the existence of a signaling pathway triggering focal adhesion assembly that bypasses the need for active PTP1B. PTP1B coimmunoprecipitates with β1-integrin from nonionic detergent extracts and colocalizes with vinculin and the ends of actin stress fibers in focal adhesions. Our data suggest that PTP1B is a critical regulatory component of integrin signaling pathways, which is essential for adhesion, spreading, and formation of focal adhesions.

Key words: protein tyrosine phosphatase • integrins • cell-substrate adhesion • cell signaling • tyrosine phosphorylation

An increasing body of evidence indicates that the function of adhesion receptors is controlled by signaling pathways regulating their association with the cytoskeleton (for reviews see Luna and Hitt, 1992; Gumbiner, 1993; Lilien et al., 1997; Yamada and Geiger, 1997). A major feature of these signaling pathways is tyrosine phosphorylation/dephosphorylation of proteins that link or modulate the interaction of adhesion receptors with the actin-containing cytoskeleton. Among cadherins, this linkage is mediated by α- and β- or γ-catenin (for reviews see Kemler, 1993; Aberle et al., 1996). Hyperphosphorylation of β-catenin on tyrosine residues is correlated with the loss of cadherin function (Behrens et al., 1993; Hamaguchi et al., 1993; Balsamo et al., 1996; Daniel and Reynolds, 1997; Hazan and Norton, 1998; Ozawa and Kemler, 1998) and destabilization or loss of the actin–cadherin linkage (Balsamo et al., 1996; Fujii et al., 1996; Daniel and Reynold, 1997; Hazan and Norton, 1998; Ozawa and Kemler, 1998). Our laboratory has shown that, in retinal cells, dephosphorylation of β-catenin by the nonreceptor protein tyrosine phosphatase PTP1B is essential for N-cadherin function (Balsamo et al., 1996). Moreover, in fibroblasts constitutively expressing N-cadherin that are transfected with a dominant-negative mutant PTP1B, β-catenin retains phosphate on tyrosine residues, N-cadherin is uncoupled from the cytoskeleton, and N-cadherin–mediated adhesion is lost (Balsamo et al., 1998).

Tyrosine phosphorylation/dephosphorylation is also critical to integrin-mediated adhesion and focal contact formation (for reviews see Clark and Brugge, 1995; Schwartz et al., 1995; Jockusch et al., 1995; Parsons, 1996; Yamada, 1997). Vinculin, talin, and α-actinin, which are prominent structural components of focal contacts, mediate the anchorage of actin stress fibers at these sites (for reviews see Burridge et al., 1988; Jockusch et al., 1995; Burridge and Chrzanowska-Wodnicka, 1996). Other proteins of focal adhesions, such as paxillin and p130Cas, appear to function as adaptors, while focal adhesion kinase...
(FAK) and Src, among others, are important intracellular signaling components (for reviews see Jockusch et al., 1995; Schwartz et al., 1995; Burridge and Chrzanowska-Wodnicka, 1996). Recent mutational studies have outlined a sequence of tyrosine phosphorylation events after integrin engagement. Early in this sequence, tyrosine 397 on FAK is autophosphorylated, creating a high-affinity binding site for the SH2 domain of the Src family kinases, recruiting Src and/or Fyn into integrin complexes (Lipfert et al., 1992; Schaller et al., 1994; Xing et al., 1994; Eide et al., 1995). Recruitment and activation of Src leads to phosphorylation of FAK on additional tyrosine residues, contributing to its full activation and to the recruitment of additional proteins to integrin complexes (Calab et al., 1995; Brown and Cooper, 1996; Hanks and Polte, 1997; Parsons and Parsons, 1997). The formation of this complex appears to be important for tyrosine phosphorylation of downstream targets, including paxillin (Schaller and Parsons, 1995). A late manifestation of this chain of tyrosine phosphorylations, in which FAK and Src are primary actors, is the assembly of focal adhesions. Consistent with a role for tyrosine kinases in focal contact assembly, the tyrosine kinase inhibitor herbimicin A inhibits the phosphorylation of FAK and paxillin and the formation of focal adhesions and stress fibers among fibroblasts plated on fibronectin (Burridge et al., 1992). The balance between tyrosine phosphorylation and dephosphorylation appears to be critical to the normal assembly of focal adhesions. Expression of v-Src (Rohrschneider, 1980; Nigg et al., 1982; Krueger et al., 1983; Nermut et al., 1991) or deregulated c-Src (Howel and Cooper, 1994; Thomas et al., 1995) leads to hyperphosphorylation of focal adhesion components, aberrant formation of focal adhesion sites, and alteration of the actin cytoskeleton.

While tyrosine kinases have been a major focus of attention in the analysis of focal adhesion formation and in the tyrosine phosphorylation occurring after integrin–matrix interaction, a potential role for tyrosine phosphatases in focal adhesion assembly is beginning to emerge. Several lines of evidence suggest that PTP1B may play a role. Engagement of integrins in platelets and T lymphocytes is accompanied by cleavage of PTP1B from a 50- to a 42-kD form and by its translocation to the cytoskeleton (Franconi et al., 1983; Ezumi et al., 1995) or deregulated c-Src (Howel and Cooper, 1994; Thomas et al., 1995) leads to hyperphosphorylation of focal adhesion components, aberrant formation of focal adhesion sites, and alteration of the actin cytoskeleton.

Reverse Transcription PCR

Expression of transfected chick PTP1B was assessed by reverse transcription PCR (RT-PCR). In brief, total RNA was isolated from cultured cells using a Qiagen kit (Chatsworth, CA) and reverse transcribed with Superscript II and oligo dT primers (GIBCO BRL). The resulting cDNA was used to amplify PTP1B by two rounds of PCR. Primers used for the first round were specific to the e-myc tag and the chicken PTP1B sequence.
The first primer set was 5' GAGCAAAAGCTCATTTGTGAG 3' from the e-mytag and 5' TTCAACAGGTCCTCCCTCC 3' from the chck PTPIB. The second primer set was 5' GCCCTAACCTGAGTGGAT- TATGGACCTG 3' (nucleotides 718–727) and 5' AGCAGTGGCTGACTGGACATCTTATC 3' (nucleotides 1154–1130), both from the chck PTPIB sequence (data sequence available from GenBank/EMBL/DDBJ under accession number U86040; Balsamo et al., 1998). For amplification of the mouse β-actin cDNA (accession number X03672), the primers used correspond to nucleotides 225–278 at the 5' end and 1023–103 at the 3' end.

**Flow Cytometry**

Surface expression of β1-integrin was analyzed in parental (LP) cells and cells expressing the wild-type (LWT) and mutant (LMU) PTPIB. The cells were grown in culture for 1–2 d and detached by a brief incubation with 0.06% trypsin. The trypsin was inactivated by addition of 1 mM (4-[2-aminoethyl]benzenesulfonfluoride) (AEBSF; Calbiochem). Cells were collected by centrifugation, resuspended, and incubated in DME/0.1% BSA for 30 min at 4°C. Primary antibodies (20 μg/ml of either rat anti-mouse α5β1-integrin or rat anti–mouse β1-integrin subunit) were then added, and the cells were incubated for an additional 30 min at 4°C. After three washes with DME, the cells were incubated with FITC-conjugated goat anti-rat IgG for 30 min at 4°C, washed three times with DME and once with PBS (137 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4, pH 7.4), and fixed in 3% paraformaldehyde. Fluorescence was analyzed on a Becton Dickinson FACScan (Bedford, MA). Background fluorescence was assessed in cells immunostained with normal rat IgG (20 μg/ml) as primary antibody, followed by FITC-conjugated goat anti–rat IgG. 5,000 cells per sample were analyzed.

**Immunofluorescence Microscopy**

LP, LWT, and LMU cells were grown on acid-washed round coverslips (Fisher Scientific) coated with 20 μg/ml fibronectin (FNRG, overnight, 4°C). At the indicated times, cells were fixed with 3% paraformaldehyde in PBS for 20 min at 4°C, permeabilized with 0.5% Triton X-100 in PBS for 10 min at room temperature, and blocked with 3% BSA in PBS for 30 min. Cells were then incubated with either of the following monoclonal antibodies: mouse anti-FAK (2.5 μg/ml), mouse antipaxillin (2.5 μg/ml), rat anti-β1-integrin (20 μg/ml), or mouse antivinculin (5 μg/ml). Incubations were carried out in a humid chamber for 1 h at 37°C. Cells were washed three times with PBS and incubated for a similar period with a combination of 0.5 mg/ml TRITC-phalloidin (Sigma Chemical Co.) and 1/60 dilution of either FITC-conjugated goat anti-mouse IgG or FITC-conjugated goat anti-rat IgG. After washing thoroughly, samples were mounted in 0.1 M Tris-HCl, pH 8.5, 8% glycerol (1:4, v/v), containing 1 μM phallolidin and 0.1% nynelidamine. Stained cells were observed using a universal microscope (Carl Zeiss, Inc., Thornwood, NY) equipped for epifluorescence. Images were captured with a SenSys CCD camera (Photometrics, Tucson, AZ) and analyzed with a Metamorph image analysis system, version 2 (Universal Imaging Corp., West Chester, PA).

Subcellular localization of GFP and GFP–PTPIB fusion proteins was examined with a laser scanning microscope (Carl Zeiss, Inc.) using a 488-nm argon laser. GFP expression was analyzed after paraformaldehyde fixation. For colocalization analysis, cells were fixed and permeabilized as indicated above. In this case, GFP and GFP fusion proteins were detected with a polyclonal anti-GFP antibody followed by an FITC-conjugated donkey anti-rabbit IgG. Images were printed on a color printer (model 8650 PS; Eastman Kodak Corp., Rochester, NY).

Staining of the endoplasmic reticulum was performed using rhodamine B hydroxyl ester. Living cells were incubated 1 min with 1 μM of the dye, washed three times with PBS, and fixed. Observations were carried out in a confocal microscope using a 543-nm helium-neon laser.

**Electrophoresis, Immunoprecipitation, and Western Blots**

Cultured cells were rinsed quickly with cold TBS (20 mM Tris-HCl, pH 7.4, 137 mM NaCl) containing 0.5 mM NaVO3 and removed from the culture dish in the same buffer containing 1% Triton X-100, 1 mM PMSF, 10 μg/ml leupeptin, 2.5 mM NaVO3, and 10 mM NaF. Cell lysates were incubated at 4°C for 15 min, with rotation, and then centrifuged for 2 min in a microcentrifuge (Eppendorf, Madison, WI) at maximum speed. Approximately 1 mg of supernatant protein was mixed with 2 μg/ml of either mouse anti-FAK or mouse antipaxillin and incubated with rotation for 3 h at 4°C. Goat anti–mouse IgG conjugated to magnetic beads (PerSeptive Biosystems, Inc., Framingham, MA) was then added, and the suspension was incubated for 1.5 h at 4°C. For β1-integrin immunoprecipitation, ~2 μg of supernatant protein was first diluted with 1 vol of TBS and incubated with 3 μg/ml of rat anti–mouse β1 antibody followed by goat anti-rat IgG conjugated to magnetic beads. Immunocomplexes were collected and washed three times with TBS/0.5% NP-40 and once with TBS, both containing the protease and phosphatase inhibitors at the concentrations indicated above. Final pellets were resuspended in sample buffer (Laemmli, 1970) and boiled for 5 min, and the beads were pulled down by centrifugation. The resulting supernatant was subjected to SDS-PAGE (8% acrylamide gels) and transferred to polyvinyl difluoride (PVDF) membranes (Bio-Rad Labs, Hercules, CA). After blocking in TBS containing 3% BSA and 0.2% Tween 20 for 1 h at room temperature, membranes were washed briefly with TBS and incubated for 1 h at 37°C with the primary antibody: 1 μg/ml antiphosphotyrosine antibody, 1/600 dilution of anti–β1-integrin antisera (Giancotti and Ruoslahti, 1990), or 2 μg/ml anti-PTPIB polyclonal IgG. Blots were washed three times with TBS/ Tween 20 and once with TBS, and incubated 1 h with the appropriate HRP-conjugated anti-IgG antibody. For determining the levels of phosphotyrosine on Src, cells were serum starved for 2 h, plated on FN-coated dishes, and harvested in RIPA buffer (0.1% SDS, 0.1% deoxycholate, 1% NP-40, 10 mM Tris, pH 7.4, 150 mM NaCl, 2 mM EDTA, 5 mM NaF) with 2.5 mM o-vanadate, 1 mM PMSF, 5 μg/ml leupeptin, and 10 μg/ml DNase. The lysates were centrifuged as above, and the supernatant was incubated with 2 μg/ml rabbit IgG or anti-Src antibody for 4 h at 4°C. The immunocomplexes were collected with goat anti-rabbit magnetic beads, washed and processed as above. Membranes were developed using the ECL system (Amersham Corp., Arlington Heights, IL). To reuse the membranes, they were first incubated in TBS containing 5% 2-mercaptoethanol and 2% SDS for 30 min at 55°C and then thoroughly washed with TBS, blocked again, and probed with a different antibody.

The anti-Src immunoprecipitates were also used to estimate the relative activity of Src from each of the cell lines. The immunoprecipitates were separated by SDS-PAGE and transferred to nitrocellulose. The membrane was incubated for 30 min in two changes of TBS containing 10 mM MgCl2, and the Src-containing band (~0.5 cm) was removed and incubated with 8 μg recombinant mutant GST-chkPTPIB in 250 μl of TBS containing 10 mM MgCl2, 0.5 mM DTT, 25 μM ATP, and 0.4 mM o-vanadate for 30 min at 37°C. The reaction was stopped by addition of 50 μl of 5× SDS sample buffer, and the samples were boiled for 2 min. The samples were separated by SDS-PAGE, transferred to PVDF membranes and immunoblotted with anti-PY antibody 4G10. To determine the amount of substrate in each lane, the immunoblot was stripped and rebotted with anti-PTPIB antibody.

**Adhesion and Spreading Assays**

96-well Falcon polystyrene tissue culture plates (Becton Dickinson) were coated overnight at 4°C with 20 μg/ml bovine fibronectin (Collaborative Biomedical Products, Bedford, MA), 250 μg/ml polylysine (PL), or 1% BSA and washed with PBS, FN, and PL-coated wells were blocked for 1 h at 37°C with 1% BSA before use. LP, LWT, and LMU cells grown overnight in serum-free medium were detached with trypsin as indicated above. 20,000 cells were added per well in HBSGK buffer (20 mM Hepes, pH 7.2, 150 mM NaCl, 3 mM KCl, and 2 mM glucose). To ensure rapid and even distribution of the plated cells onto the bottom of the wells, plates were briefly centrifuged (model GH3.8 rotor; Beckman Instruments, Fullerton, CA). Cells were allowed to attach for 45 min at 37°C. Nonadherent cells were removed by washing with HBSGK and the number of attached cells was quantitated by crystal violet staining (Bonnekoh et al., 1989). In brief, attached cells were fixed for 10 min with 3% paraformaldehyde, washed with PBS, and stained with a crystal violet solution (0.2%, wt/vol in 10% ethanol, vol/vol) for 10 min. Stained cells were rinsed thoroughly with 0.5 M Tris-HCl, pH 7.8, and resuspended in 50 ml of the same buffer containing 0.5% SDS and 50% ethanol. After 30 min, optical density was quantitated at 585 nm using a microplate reader (Bio-Rad Laboratories). When indicated, cells were preincubated for 20 min on ice with the GRO/SG or GRAD/SP peptide (10 μM) and plated in the presence of the peptides.

For spreading assays, cells were prepared as above, except that they were plated on fibronectin-coated coverslips. After 30 min, cells were fixed with 3% paraformaldehyde, washed with PBS, and incubated with DII (10 μM) in the dark for 30 min. Excess dye was washed with PBS, and the cells were mounted in 80% glycerol/PBS. The area of fluorescent cells was measured using a Metamorph image analysis system (Universal Imaging Corp.).
Results

Cell–Substrate Adhesion and Spreading Is Impaired in L Cells Expressing Catalytically Inactive PTP1B

To investigate the role of PTP1B in integrin-mediated adhesion, we transfected L cells with a vector containing the full-length cDNA for chicken PTP1B (wild-type: LWT) or a catalytically inactive form of the phosphatase (C215S: LMU). Both cDNAs carry a c-myc tag in frame at the 5′ end to facilitate subsequent analysis. Stable cell lines were selected with zeocin, and expression of the c-myc–tagged PTP1B was determined by RT-PCR, using a combination of primers specific to c-myc and the chick enzyme, and by immunoblot. A unique band of 437 bp corresponding to the chick PTP1B message is readily detected in LWT and LMU cells (Fig. 1A). As expected, chick PTP1B message is not detected in LP cells (Fig 1A). Actin message was used to normalize the amount of starting material from each cell preparation (Fig. 1A). PTP1B is readily detected in both control and transfected populations (Fig. 1B). Because we used a pan-specific anti-PTP1B, we were able to estimate the level of overexpression to be about twofold (Fig. 1B).

To assess the effect of transfected wild-type and inactive PTP1B on integrin-mediated adhesion, several clones of LWT and LMU cells as well as LP cells were plated on fibronectin-coated, 96-well plates in serum-free medium, and the attached cells were quantified after 45 min incubation. Only 30% of the input LMU cells adhered, as compared with more than 90% of the LP and LWT cells (Fig. 2A). Attachment of control LP and LWT cells to fibronectin is integrin-dependent: over 90% attachment is inhibited by an RGD-containing peptide (Fig. 2A). Several clones expressing wild type and mutant were analyzed by phase contrast microscopy after culture in the presence of serum. All clones expressing the mutant PTP1B (LMU) are similar and morphologically different from clones expressing the wild-type protein (LWT) or the parental cell line (LP) (Fig. 2B). LMU cells show a bipolar, spindle shape and do not flatten onto the substrate. In contrast, LP and LWT cells form extensive lamellae and are flattened on the matrix (Fig. 2B).

The poor attachment of LMU cells to fibronectin is also reflected in their inability to flatten and spread. 30 min after plating on fibronectin, most LMU cells remain round...
and very loosely attached to the substrate, with only a few cells showing limited spreading. In contrast, most of the LP and LWT cells flatten and form large lamella (Fig. 3A). The surface areas of LP and LWT cells are similar and approximately three times that of LMU cells, as determined by quantitative image analysis (Fig. 3B). The reduced attachment and spreading of LMU cells on fibronectin is not due to a decreased expression of integrins at the cell surface since flow cytometry profiles of surface-labeled $\alpha_5\beta_1$-integrin and the major fibronectin receptor, $\alpha_5\beta_1$, are equivalent in all three cell types (Fig. 4). In addition, immunoblots of cell lysates with anti–mouse $\beta_1$-integrin antibody show equal levels of expression (not shown).

**Focal Adhesions and Actin Stress Fibers Are Altered in L Cells Expressing Catalytically Inactive PTP1B**

In normal fibroblasts, integrin-mediated adhesion to fibronectin leads to the formation of focal adhesions and assembly of actin stress fibers (Burrage et al., 1988). The impaired attachment and spreading of LMU cells on fibronectin suggests that these cells may be unable to assemble normal focal contacts and actin stress fibers. To investigate this possibility, LMU cells were plated on fibronectin for 30 min and then fixed and processed for double immunofluorescence to detect F-actin and $\beta_1$-integrin. Actin appears diffusely distributed throughout the cell body but accumulates in the submembranal cytoskeleton (Fig. 5A). $\beta_1$-integrin is distributed uniformly in a diffuse pattern all over the cell, with occasional small aggregates (Fig. 5B). In contrast, among LWT cells, actin is organized in stress fibers (Fig. 5C), and $\beta_1$-integrin accumulates in focal contacts (Fig. 5D).

To determine if LMU cells have a delayed ability to assemble stress fibers and focal contacts, cells were plated on fibronectin and cultured for 2 d in the presence of serum before staining with TRITC-phalloidin and anti-$\beta_1$ antibody. After this time, most LMU cells have an elongated shape but still lack actin stress fibers and focal adhesions. Actin is distributed in short filaments within the cell body and in the submembranal cytoskeleton (Fig. 6A). Actin filaments are also seen aligned along the main axis of the cells and following a radial pattern within the lamellipodia (Fig. 6, B and D). As expected, LWT and LP cells actin accumulates in prominent stress fibers and $\beta_1$-integrin in focal adhesions (Fig. 6, LWT: E and F; LP: G and H).

The distributions of other typical focal adhesion components, such as FAK, paxillin, and vinculin, are also altered in LMU cells. In all cases, the distribution is dotted, with occasional formation of small aggregates (Fig. 7).
These proteins are distributed in large and elongated focal adhesions in LWT (Fig. 7) and LP cells (not shown).

**Tyrosine Phosphorylation of FAK and Paxillin Is Reduced in L Cells Expressing Catalytically Inactive PTP1B**

Tyrosine phosphorylation of FAK is an early event occurring after integrin binding to the ligand (Guan and Shalloway, 1992; Lipfert et al., 1992; Kornberg et al., 1992; Clark and Brugge, 1995). To examine whether phosphorylation of FAK on tyrosine residues is affected in LMU cells, non-ionic detergent extracts of cells plated on polylysine or fibronectin for 30 min were immunoprecipitated with an anti-FAK antibody and immunoblotted with an antiphosphotyrosine antibody. On polylysine, phosphotyrosine levels on FAK are barely detectable in the three cell lines (not shown). On fibronectin, phosphotyrosine levels in FAK are significantly reduced in LMU cells as compared with those in LP and LWT cells (Fig. 8 A, top), suggesting that phosphorylation of FAK on tyrosine residues in response to the attachment to fibronectin is indeed impaired in LMU cells.

Similarly, paxillin is localized in focal adhesions and becomes tyrosine phosphorylated after integrin-dependent adhesion (Burridge et al., 1992). To determine whether phosphorylation of paxillin on tyrosine is also affected in LMU cells, lysates from cells plated on fibronectin were immunoprecipitated with a specific antipaxillin antibody, and phosphotyrosine residues were detected as described above. Indeed, in LMU cells, paxillin shows reduced levels of phosphotyrosine when compared with LP and LWT cells (Fig. 8 B, top). The reduced phosphorylation of FAK and paxillin in LMU cells cannot be attributed to a lower level of protein expression or to differential immunoprecipitation since equivalent amounts of protein were used for all immunoprecipitations, and the amount of immunoprecipitated FAK and paxillin is similar in the three cell lines, as determined by stripping the membranes and blotted with anti-FAK (Fig. 8 A, bottom) or antipaxillin (Fig. 8 B, bottom) antibodies.

To eliminate the possibility that a general dephosphorylation of proteins occurs in cells expressing the mutant PTP1B, lysates from cells plated on fibronectin were fractionated by SDS-PAGE, transferred to PVDF membrane, and immunoblotted with antiphosphotyrosine antibody. Tyrosine phosphate residues at the appropriate molecular mass for FAK and paxillin are reduced in LMU cells (Fig. 8 C). There are changes in the phosphorylation levels of other proteins in LMU cells; however, there are no wholesale changes in the pattern. It is of interest that phosphorylation of high (~200 kD) and low (~30 kD) molecular mass bands are reduced in LWT cells, indicating that in-

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**Figure 5.** Distribution of β1-integrin and actin in cells plated for 30 min on fibronectin. Serum-starved LMU (A and B) and LWT (C and D) cells in DME were plated on fibronectin-coated coverslips and fixed 30 min later. The cells were processed for double immunofluorescence staining with rat anti-mouse β1 mAb/FITC-goat anti-rat IgG (B and D) and TRITC-phalloidin (A and C). Note the lack of spreading, focal contacts, and actin stress fibers in LMU cells. Bar, 15 µm.

**Figure 6.** Distribution of β1-integrin and actin in cells cultured for 2 d. LMU (A–D), LWT (E and F), and LP (G and H) cells in DME/FBS were plated on fibronectin-coated coverslips. After 2 d, cells were fixed, permeabilized, and double stained with rat anti-β1 mAb/FITC-goat anti-rat IgG (B, D, F, and H) and TRITC-phalloidin (A, C, E, and G). Bar, 10 µm.
creased levels of active PTP1B does alter the pattern of phosphorylation.

**Tyrosine Phosphorylation of Src Is Increased and Activity Decreased in L Cells Expressing Catalytically Inactive PTP1B**

The decrease in tyrosine phosphorylation of FAK and paxillin in cells expressing the catalytically inactive PTP1B indicates that PTP1B plays a role in regulating the tyrosine phosphorylation of components upstream from these phosphorylation events. Src is an obvious candidate since FAK and paxillin are downstream targets of Src (Clark and Brugge, 1993; Howel and Cooper, 1994; Kaplan et al., 1994; Calalb et al., 1995; Schaller and Parsons, 1995; Hanks and Polte, 1997). Tyrosine 527 must be dephosphorylated to activate Src (Courtneidge, 1985; Cooper and

![Figure 7](image7.png)

*Figure 7.* Distribution of FAK, paxillin, and vinculin in LMU and LWT cells plated on fibronectin-coated coverslips. LWT (A, C, and E) and LMU (B, D, and F) cells cultured for 2 d were fixed and processed for immunofluorescence using monoclonal antibodies against paxillin (A and B), FAK (C and D), or vinculin (E and F). Bar, 20 µm.

![Figure 8](image8.png)

*Figure 8.* Phosphotyrosine levels of FAK and paxillin in LP, LWT, and LMU cells. LP, LWT, and LMU cells were plated on fibronectin for 30 min, lysed (see Materials and Methods), and immunoprecipitated (IP) with (A) anti-FAK mAb or (B) anti-paxillin mAb and immunoblotted (IB) with antiphosphotyrosine (PY) antibody. Lower panels show stripped membranes reblotted with the same antibodies used for immunoprecipitation. (C) Phosphotyrosine staining of total proteins in Triton X-100 lysates. LP, LWT, and LMU cells plated on fibronectin for 30 min were lysed as described in Materials and Methods. Numbers at the left indicate the position of molecular mass markers in kilodaltons.

King, 1986; Brown and Cooper, 1996) and mutant PTP1B may block this event. An increase in tyrosine phosphate content of Src has been correlated with decreased activity (Soltész et al., 1997). To assess the tyrosine phosphate content of Src, extracts of cells plated on fibronectin for 1 h were immunoprecipitated with an anti-Src antibody and immunoblotted with an antiphosphotyrosine antibody. The phosphotyrosine content of Src is increased in the LMU cells as compared with LP cells (Fig. 9A). It is interesting to note that overexpression of wild-type PTP1B decreases the phosphotyrosine content of Src, without an apparent alteration of the phenotype (Fig. 9A). The amount of Src in each lane is also shown (Fig. 9A, bottom).

To directly estimate the relative activity of Src in each of the cell types, Src was immunoprecipitated and assayed in vitro using mutant chk-PTP1B as the substrate (Jung et al., 1998). Transfer of phosphate to the substrate was determined by SDS-PAGE and immunoblot with anti-PY antibody (Fig. 9B). Src isolated from LMU cells is clearly less active than that isolated from either LP or LWT cells.

**Subcellular Distribution of PTP1B**

It has been shown that PTP1B is predominantly localized in the endoplasmic reticulum (ER) (Frangioni et al., 1992;
Woodford-Thomas et al., 1992). To investigate whether a fraction of PTP1B is associated with integrin complexes, nonionic detergent extracts of LWT and LMU cells plated for 30 min on fibronectin were immunoprecipitated with a rat anti–β1-integrin antibody, and the precipitates were analyzed for the presence of PTP1B after SDS-PAGE and immunoblot. Rabbit anti-PTP1B antibody reveals a band at 50 kD, the approximate molecular mass for the intact chicken PTP1B protein (Fig. 10A). This band is not present in control immunoprecipitates. To confirm the immunoprecipitation of β1-integrin, blots were probed with a polyclonal anti-β1 antibody (Fig. 10B).

In an alternative approach, L cells were transiently transfected with wild-type and catalytically inactive (C215S) forms of chick PTP1B tagged with the green fluorescent protein at the NH2 terminus. The cells were analyzed for expression of the GFP–PTP1B construct by immunoblot with anti-PTP1B antibody and anti-GFP antibody (Fig. 11). Parental PTP1B migrates at ~50 kD, while the GFP fusion migrates at ~77 kD (27 kD + 50 kD). The anti-GFP antibody recognizes the band at ~77 kD as expected. Among cells that express high levels of GFP–wtPTP1B and GFP–mutPTP1B, both proteins are seen in a pattern typical for the ER as revealed by staining with the ER marker rhodamine B hexyl ester (Terasaki and Reese, 1992) (Fig. 12, compare A and B with D). In contrast, cells transfected with the GFP alone show a uniform distribution of the GFP (Fig. 12C). These results indicate that the GFP tag does not significantly affect the targeting of PTP1B to the ER, a subcellular localization previously reported for endogenous PTP1B (Frangioni et al., 1992; Woodford-Thomas et al., 1992).
Among cells that express moderate or low levels of GFP–wtPTP1B, localization at focal adhesions is observed. To confirm this, double immunofluorescence analyses were performed comparing the distribution of the GFP–wtPTP1B with that of vinculin and actin, two prominent components of focal adhesions. Analysis of 1-μm optical sections taken at the cell–substrate level shows that GFP–wtPTP1B colocalizes with actin at the ends of the stress fibers (Fig. 13, arrows in C, and compare with A). GFP–wtPTP1B also shows a high degree of colocalization with vinculin (Fig. 13, arrows in D, and compare with B).

In addition, GFP–wtPTP1B staining was compared with images taken by reflection interference microscopy, where the pattern of focal adhesions is revealed. The overlap between the immunofluorescence of GFP–wtPTP1B and the dark regions at the margins of cellular lamella seen by reflection interference optics indicates that wtPTP1B localizes in areas of focal adhesions (Fig. 13, arrowheads). Similar results are seen using anti-PTP1B antibody (not shown).

Lysophosphatidic Acid Stimulates Assembly of Focal Adhesions in LMU Cells

Growth factors present in serum, such as lysophosphatidic acid, induce the assembly of focal contacts and actin stress fibers, as well as increase the phosphorylation of FAK and paxillin on tyrosine residues (Ridley and Hall, 1992; Zachary et al., 1993; Seufferlein and Rozengurt, 1994). To determine whether expression of the mutant PTP1B also af-
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fects the LPA-dependent signaling pathway, LMU cells were plated on fibronectin in the presence or absence of LPA. In the presence of LPA, LMU cells rapidly flatten and produce extensive lamella (Fig. 14, compare A with C). The area of spreading among LMU cells in the presence of LPA is equivalent to that of LWT cells (Fig. 14, compare B with C) or LP cells (not shown) as determined by quantitative image analysis. Moreover, in LMU cells treated with LPA, β1-integrin and F-actin show their typical distribution in focal contacts and stress fibers (Fig. 14, C and D). The distribution of vinculin and FAK is similar to that of β1-integrin (not shown).

To determine whether there is an increase in phosphorylation of FAK on tyrosine residues as a response to LPA, nonionic detergent lysates of LMU cells were immunoprecipitated with anti-FAK antibody, and the precipitates were separated by SDS-PAGE and immunoblotted with antiphosphotyrosine antibody. 15 min after plating LMU cells on fibronectin, in the presence of LPA, the levels of phosphorylated tyrosine residues on FAK are similar to those in LP cells (Fig. 14). The effects of LPA in inducing spreading, focal contact and stress fiber formation, and FAK phosphorylation require integrin ligation since they are not observed in cells grown on polylysine (not shown). The rapid response of LMU cells to LPA treatment rules out the possibility that the machinery essential for phosphorylation of FAK and paxillin and for assembly of focal contacts and stress fibers is defective in LMU cells.

Discussion

In this paper, we present data that suggest a regulatory role for PTP1B in integrin-mediated adhesion and spreading. L cells expressing a catalytically inactive form of PTP1B display reduced attachment and are unable to spread on fibronectin. They assume an elongated spindle shape after prolonged culture but fail to form the typical flattened phenotype. Consistent with this altered morphology, LMU cells also show a dramatic decrease in focal adhesions and actin stress fibers. They are able, however, to assemble actin into long filaments parallel to the long axis of the cell and short filaments in the submembrane region of cells and lamellipodia. Additionally, the proportion of polymeric actin in LMU cells is similar to that in LWT or LP cells (Arregui, C.O., unpublished observation). The level of expression of β1-integrin at the cell surface is unaltered in LMU cells; however, its distribution changes to a dotted pattern that contrasts with that of morphologically differentiated focal adhesion complexes in control cells.

Since activated rho is required for the assembly of focal adhesions and actin stress fibers (Ridley and Hall, 1992; Barry and Critchley, 1994; Nobes and Hall, 1995), we considered the possibility that rho-dependent signaling could be impaired in LMU cells. However, when incubated with serum or lysophosphatidic acid, both of which activate the rho pathway (Ridley and Hall, 1992), LMU cells rapidly acquire a phenotype similar to parental or LWT cells, indicating that the machinery to assemble actin stress fibers and focal adhesions is intact. The effect of LPA or serum is dramatic but transient, since cells cultured for longer periods do not retain the spread shape and do not develop focal adhesions and stress fibers. Thus, the normal interac-

![Figure 14](image-url)

Figure 14. Effect of lysophosphatidic acid in LMU cells. Serum-starved cells were plated in DME on fibronectin-coated coverslips in the absence or presence of 200 ng/ml lysophosphatidic acid and fixed after 15 min. Cells were permeabilized and stained with rat anti–β1 antibody (D) and TRITC-phalloidin labeled (A–C). Anti–β1-integrin staining was visualized by a FITC-conjugated goat anti–rat IgG. (A) LMU cells plated in the absence of LPA. (B) LWT cells plated in the absence of LPA. (C and D) LMU cells exposed to LPA and stained for actin (C) or β1-integrin (D). Note the formation of actin stress fiber (C) and induction of focal contacts (D) in LMU cells exposed to LPA. Tyrosine phosphorylation of FAK is shown in the immunoblot below. LP, LWT, and LMU cells were plated on fibronectin, lysed after 15 min under denaturing conditions, and immunoprecipitated with anti-FAK antibody. The immunoprecipitates were fractionated by SDS-PAGE and analyzed by immunoblotting with the 4G10 monoclonal antiphosphotyrosine antibody. Note the difference in the phosphorylation of FAK when LMU cells were plated in the absence or presence of LPA. The bottom panel shows the same blot stripped and probed with anti-FAK antibody. Numbers at the left indicate the position of molecular mass markers in kilodaltons. Bar, 20 μm.
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**PTP1B as Modulator of Integrin-mediated Adhesion**

Focal adhesion complexes contain many components, some of which serve structural roles, forming the actual adhesion and links to the actin-based cytoskeleton. Other components of the complex serve to propagate signals initiated outside the cell by ligation of integrins, or those initiated by growth factors and activation of integrins from inside the cell (for reviews see Burridge and Chrzanowska-Wodnicka, 1996; Machesky and Hall, 1996). Furthermore, it is clear that there is redundancy of function among components of focal adhesion complexes (Soriano et al., 1991; Ilic et al., 1995; Thomas et al., 1995). It is important to emphasize that, given this complexity, interruption of the signaling pathway(s) leading to loss of focal contact formation may result in several different phenotypes with respect to the phosphorylation of focal contact components, or even the array of components associated with integrin adhesion complexes. These phenotypes will depend on the point at which the signal cascade is interrupted.

Src is an obvious target for the effect of the dominant-negative PTP1B. FAK and paxillin are downstream targets of Src (Clark and Brugge, 1993; Howel and Cooper, 1994; Kaplan et al., 1994; Calalb et al., 1995; Schaller and Parsons, 1995; Hanks and Polte, 1997), and the enzymatic activity of Src is regulated by the balance of phosphorylation/dephosphorylation of tyrosine 527. The COOH-terminal Src kinase (Csk) phosphorylates Y527 and represses Src activity. Recruitment of Src to focal adhesions is increased in cells isolated from Csk-minus mice (Kaplan et al., 1995). We are suggesting that PTP1B may also play such a role. Our results are consistent with this hypothesis; transfection with the dominant-negative, catalytically inactive PTP1B results in increased phosphorytosine content of Src and decreased activity, presumably by preventing the dephosphorylation of Tyr 527 by endogenous PTP1B or other phosphatases, keeping Src in a repressed state (see Fig. 15).

Several additional lines of evidence are also consistent with the hypothesis that the reduced levels of tyrosine phosphorylation of FAK and paxillin in LMC cells are due to repression of Src. Recruitment of Src to focal adhesions follows integrin engagement (Clark and Brugge, 1993; Kaplan et al., 1995) and is accompanied by a transient increase in Src activity concomitant with a decrease in the phosphorylation of Tyr 527 (Clark and Brugge, 1993; Kaplan et al., 1995). FAK and paxillin tyrosine phosphorylation is increased in cells isolated from Csk-minus mice (Thomas et al., 1995), and overexpression of Csk in HeLa cells results in impaired cell-substrate adhesion and spreading and a redistribution of integrins from focal contacts to point contacts (Bergman et al., 1995), a phenotype similar to our L cells expressing mutant PTP1B. Finally, fibroblasts from Src knock out mice also show reduced spreading (Kaplan et al., 1995).

In seeming contrast to our results, Liu et al. (1998) have reported that introduction of wild-type PTP1B into 3Y1 fibroblasts reduces the rate of cell spreading and ultimately results in the formation of abnormally dense bundles of stress fibers and enlarged, elongated focal contacts. This was accompanied by reduced levels of phosphorylation of FAK (Liu et al., 1998). We do not see an effect of wild-type PTP1B on adhesion or focal contacts in L cells. In agreement with our observations, spreading and cytoskeletal organization are not altered by expression of wild-type PTP1B in human foreskin fibroblasts or W-87MG glioblastoma cells (Tamura et al., 1998). Interestingly, we do see a significant enhancement of neurite outgrowth by PC12 cells expressing wild-type PTP1B (Pathre, P., C. Arregui, J. Lilien, and J. Balsamo, manuscript in preparation). This may reflect effects similar to those seen by Liu et al. (1998), a possibility we are investigating. The different phenotypes resulting from perturbation of PTP1B may therefore be due to differences in host cell type and/or expression levels of PTP1B.

**Subcellular Distribution of PTP1B**

We find that PTP1B coimmunoprecipitates with β1-inte...
This suggests possibilities for cross regulation of these two complexes (Balsamo et al., 1996; Balsamo et al., 1998). It is interesting that other tyrosine phosphatases also interact with p130Cas (Black and Blisika, 1997; Garton et al., 1997; Shen et al., 1998).

Like targeting to p130Cas, the proline-rich region of PTP1B may also interact with the SH3 domain of Src itself. The selective dephosphorylation of Tyr 527 over 416 may be due to sequence recognition by PTP1B—tyrosines 527 and 416 of Src are presented in entirely different sequence motifs (Takeya and Hanafusa, 1983)—and/or accessibility based on targeting. It is worth emphasizing that targeting mechanisms can impose limits on the available protein substrates and specific tyrosine residues.

The Many Faces of PTP1B

Full-length PTP1B was initially described as localized to the endoplasmic reticulum (Frangioni et al., 1992; Woodford-Thomas et al., 1992). PTP1B has also been reported to interact with both the insulin and IGF-1 receptors regulating the ligand-induced signaling via these receptors (Kenner et al., 1996; Seely et al., 1996; Bandyopadhyay et al., 1997). It also interacts with and is phosphorylated by the EGF receptor (Milarski et al., 1993; Flint et al., 1997; Liu et al., 1997). Here we report that PTP1B is functionally and physically associated with focal contacts. We have also reported that, in chick neural retina cells and in L cells stably expressing N-cadherin, a fraction of PTP1B binds to N-cadherin and regulates its function through dephosphorylation of β-catenin (Balsamo et al., 1996; Balsamo et al., 1998). PTP1B thus plays multiple roles depending on specific interactions, whether via substrate recognition or other specific targeting regions of the molecule.

Finally, it is interesting to note that PTP1B plays an important role in regulating both integrin adhesion complexes (Liu et al., 1998: this paper) and cadherin adhesion complexes (Balsamo et al., 1996; Balsamo et al., 1998). This suggests possibilities for cross regulation of these two developmentally important cell adhesion systems.

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