The Interaction of Activated Integrin Lymphocyte Function-associated Antigen 1 with Ligand Intercellular Adhesion Molecule 1 Induces Activation and Redistribution of Focal Adhesion Kinase and Proline-rich Tyrosine Kinase 2 in T Lymphocytes

José Luis Rodríguez-Fernández,* Manuel Gómez,† Alfonso Luque,* Nancy Hogg,‡ Francisco Sánchez-Madrid,† and Carlos Cabañas*§

*Departamento de Bioquímica y Biología Molecular, Facultad de Medicina, Universidad Complutense, 28040 Madrid, Spain; †Servicio de Inmunología, Hospital de la Princesa, 28006 Madrid, Spain; and ‡Leukocyte Adhesion Laboratory, Imperial Cancer Research Fund, London WC2A 3PX, United Kingdom

Submitted August 3, 1998; Accepted March 31, 1999
Monitoring Editor: Joan Brugge

Integrin receptors play a central role in the biology of lymphocytes, mediating crucial functional aspects of these cells, including adhesion, activation, polarization, migration, and signaling. Here we report that induction of activation of the β2-integrin lymphocyte function-associated antigen 1 (LFA-1) in T lymphocytes with divalent cations, phorbol esters, or stimulatory antibodies is followed by a dramatic polarization, resulting in a characteristic elongated morphology of the cells and the arrest of migrating lymphoblasts. This cellular polarization was prevented by treatment of cells with the specific tyrosine kinase inhibitor genistein. Furthermore, the interaction of the activated integrin LFA-1 with its ligand intercellular adhesion molecule 1 induced the activation of the cytoplasmic tyrosine kinases focal adhesion kinase (FAK) and proline-rich tyrosine kinase 2 (PYK-2). FAK activation reached a maximum after 45 min of stimulation; in contrast, PYK-2 activation peaked at 30 min, declining after 60 min. Upon polarization of lymphoblasts, FAK and PYK-2 redistributed from a diffuse localization in the cytoplasm to a region close to the microtubule-organizing center in these cells. FAK and PYK-2 activation was blocked when lymphoblasts were pretreated with actin and tubulin cytoskeleton-interfering agents, indicating its cytoskeletal dependence. Our results demonstrate that interaction of the β2-integrin LFA-1 with its ligand intercellular adhesion molecule 1 induces remodeling of T lymphocyte morphology and activation and redistribution of the cytoplasmic tyrosine kinases FAK and PYK-2.

INTRODUCTION

Integrins are heterodimeric cell surface proteins that function in cell adhesion, cytoskeleton anchorage, and the transduction of cellular stimuli into cytoplasmic signals (Clark and Brugge, 1995; Schwartz et al., 1995). The β2-integrin lymphocyte function-associated antigen 1 (LFA-1; CD11a/CD18) is selectively expressed on leukocytes and mediates important adhesive phenomena of these cells through interaction with its ligands intercellular adhesion molecule 1 (ICAM-1; CD54), ICAM-2 (CD102), and ICAM-3 (CD50). The resting leukocytes, which circulate in the bloodstream, express an inactive form of LFA-1 that is functionally unable to mediate interactions with ICAMs. Induction
of leukocyte activation through different cell surface receptors, including the T cell receptor–CD3 complex and receptors for cytokines, results in the generation of intracellular signals that lead to activation of LFA-1 molecules. This “inside-out” mechanism of LFA-1 activation brings about a change in the conformation of the extracellular region of the integrin, which is functionally reflected by the ability of the cells to adhere to LFA-1 ligands. Activation of LFA-1 molecules can also be induced directly from outside the cells by altering the extracellular divalent cation conditions, i.e. by addition of micromolar concentrations of Mn^{2+} or by the presence of millimolar levels of Mg^{2+} when Ca^{2+} is removed with EGTA (Dransfield et al., 1992a). Direct activation of LFA-1 from outside the cells can also be achieved with specific “activating” or “stimulating” mAbs. These types of antibodies recognize and bind to the α or the β subunits of LFA-1, altering the conformation of the integrin to a state of increased affinity for its ligands. Activating LFA-1 antibodies include NKI-L16, which is specific for the α subunit or CD11a, and KIM-127 and KIM-185, which are directed to the common β2 subunit CD18 (van Kooyk et al., 1991; Ortlepp et al., 1995; Stephens et al., 1995). Phorbol esters such as phorbol-12,13-dibutyrate (FDBu), which are potent and sustained activators of PKC, have also been used extensively as another means of inducing LFA-1–mediated adhesion to ICAMs (Rothlein and Springer, 1986; Stewart et al., 1996).

Upon ligand engagement of LFA-1, a variety of intracellular signals are generated in T lymphocytes, including phosphorylation of phospholipase Cγ, phospholipid hydrolysis, activation of different isoenzymes of PKC, mobilization of intracellular Ca^{2+}, and activation of tyrosine kinases (Wacholtz et al., 1989; Kanner et al., 1993; Arroyo et al., 1994; Lub et al., 1995). The focal adhesion kinase (FAK) is a nonreceptor protein-tyrosine kinase that colocalizes to cellular focal adhesions with integrins (Hanks et al., 1992; Schaller et al., 1992). FAK, one of the major substrates for integrin-dependent tyrosine phosphorylation, is considered an important element in an integrin-regulated pathway (Clark and Brugge, 1995; Richardson and Parsons, 1995; Otey, 1996; Hanks and Polte, 1997). FAK is tyrosine phosphorylated upon stimulation of β1-integrins in fibroblasts, epidermal carcinoma cells, mast cells, and T lymphocytes; upon engagement of β3-integrin in platelets and upon transfection of human fibroblasts with chimeric receptors containing either the β1-, β3-, or β5-integrin intracellular domain (Kornberg et al., 1991; Guan and Shalloway, 1992; Lipfert et al., 1992; Hamawy et al., 1993; Akiyama et al., 1994; Matsumoto et al., 1994; Shattil et al., 1994; Maguire et al., 1995). Concomitant with tyrosine phosphorylation, FAK becomes enzymatically active (Guan and Shalloway, 1992; Lipfert et al., 1992; Otey, 1996). A variety of studies, including the characterization of cells derived from FAK(−/−) embryos, suggests that FAK may be involved in the regulation of cell spreading and motility (Gates et al., 1994; Akasaka et al., 1995; Ilic et al., 1995; Cary et al., 1996; Gilmore and Romer, 1996; Richardson and Parsons, 1996).

A second member of the FAK nonreceptor tyrosine kinase family, known as proline-rich tyrosine kinase 2 (PYK-2), and also called related adhesion focal tyrosine kinase, cell adhesion kinase β, calcium-dependent tyrosine kinase, or FAK2, has been described (Avraham et al., 1995; Earp et al., 1995; Lev et al., 1995; Sasaki et al., 1995; Herzog et al., 1996). PYK-2 is less evenly expressed in tissues and restricted to a lower number of cell types compared with FAK; however, the expression is high in cells of neural, epithelial, or hematopoetic origin (Avraham et al., 1995; Lev et al., 1995; Yu et al., 1996). PYK-2 shares significant homology with FAK (60% identity in the central catalytic domain and 40% identity in both C and N termini) and, like FAK, does not contain Src homology 2 (SH2) or SH3 domains but contains several sites for binding of SH2- and SH3-containing signaling proteins. Regulation of PYK-2 localization and tyrosine phosphorylation seems to be cell type and integrin specific. PYK-2 localizes to focal contacts and displays a β1-integrin–dependent phosphorylation in natural killer cells, B lymphocytes, megakaryocytes and transfected COS cells and exhibits a β3-dependent tyrosine phosphorylation in T lymphocytes (Li et al., 1996; Astier et al., 1997; Gismondi et al., 1997; Ma et al., 1997). However, PYK-2 was localized to intercellular junctions and was not tyrosine phosphorylated upon adhesion of transfected 3Y1 fibroblasts on fibronectin or integrin ligation during platelet aggregation (Sasaki et al., 1995; Raja et al., 1997). It has been suggested that the differences in localization and tyrosine phosphorylation of FAK and PYK-2 may be due to differences in the sequences of the C termini of FAK and PYK-2 (Zheng et al., 1998). Several studies suggest that PYK-2 may be involved in the control of apoptosis and the regulation of ion channels in neuronal cells (Lev et al., 1995; Xiong and Parsons, 1997).

After activation and autophosphorylation of FAK and PYK-2, these enzymes may serve as recruiting molecules for various signaling and cytoskeletal proteins, including the tyrosine kinases Src and Fyn, and the focal contact proteins paxillin and p130Cas, which are also potential targets for FAK and PYK-2 phosphorylation (Richardson and Parsons, 1995; Dikic et al., 1996; Astier et al., 1997; Ganju et al., 1997; Gismondi et al., 1997; Hanks and Polte, 1997; Hiregowdara et al., 1997; Ohba et al., 1998; Schlaepfer and Hunter, 1998). Therefore, integrin signaling, involving the activation of FAK and PYK-2, has also been suggested to modulate the reorganization of the actin cytoskeleton and to regulate changes in cell morphology. Polarization is a key feature in the biology of T cells (Wilkinson and
Higgins, 1987; del Pozo et al., 1996; Sánchez-Madrid and del Pozo, 1999). Lymphocytes acquire a polarized phenotype after activation, upon interaction with antigen-presenting cells (APCs) and during chemokine-directed transendothelial migration. The leukocyte integrin LFA-1 plays a key role in these crucial events of immune and inflammatory responses (Springer, 1990; Dransfield et al., 1992a). Furthermore, cell polarization in lymphocytes involves a reorganization of the cytoskeleton, including polymerization and redistribution of actin and reorientation of the microtubule-organizing center (MTOC) toward the opposing cell (Singer, 1992; Dustin et al., 1997; Lowin-Kropf et al., 1998).

In this study we have analyzed some of the morphological and functional changes that occur in T lymphocytes after induction of activation of the integrin LFA-1 and its interaction with its ligand ICAM-1. We demonstrate that functional activation of LFA-1 brings about dramatic changes in the morphology and polarization of these cells, converting them into a non-motile phenotype. We also demonstrate, for the first time, that concomitantly with these changes, the two homologous cytoplasmic tyrosine kinases FAK and PYK-2 become activated and redistributed intracellularly, colocalizing in the polarized T cell with the MTOC.

MATERIALS AND METHODS

Cells
Human T lymphoblasts were prepared from peripheral blood mononuclear cells by treatment with 0.5% phytohemmagglutinin for 48 h. Cells were washed and cultured in RPMI 1640 (Flow Laboratories, Irvine, Scotland) containing 10% FCS (Flow Laboratories) and interleukin 2 (20 U/ml). T lymphoblasts cultured by 7–12 d were typically used in all experiments (Dransfield et al., 1992b; Luque et al., 1996; Stewart et al., 1996).

Antibodies and Reagents
A dimeric form of an ICAM-1-Fc chimeric protein consisting of the five domains of ICAM-1 fused to the Fc fragment of human IgG1 was prepared as previously described (Berendt et al., 1992). The LFA-1-stimulatory mAb KIM-127, which is specific for the β2-integrin subunit (Ortlepp et al., 1995; Stephens et al., 1995), was a generous gift of Dr. Martyn Robinson (Celltech, Slough, United Kingdom). FAK antipeptide polyclonal antibodies C-20 and A-17 and PYK-2 antipeptide polyclonal antibodies C-19 and N-19 were from Santa Cruz Biotechnology (Santa Cruz, CA). The anti-FAK C-20 and A-17 polyclonal antibodies were raised against peptides corresponding to amino acids 1033–1052 mapping at the C terminus and amino acids 2–18 mapping at the N termini of human FAK, respectively. The anti-PYK-2 polyclonal antibodies C-19 and N-19 were raised against peptides corresponding to amino acids 990–1009, mapping at the C terminus, and amino acids 2–20, mapping at the amino terminus, of human PYK-2, respectively. The anti-FY(p) PY20 and the anti-FAK a-FAK mAbs were from Transduction Laboratories (Lexington, KY). The mAb a-FAK recognizes an epitope in the kinase domain of FAK, corresponding to amino acids 354–533 of the chicken protein. The anti-FAK mAb 2A7 was from Upstate Biotechnology (Lake Placid, NY). The 2A7 mAb recognizes an epitope in the C domain of FAK (Schaller et al., 1993). The anti-FY(p) mAb clone PY72 was obtained from the Hybridoma Development Unit, Imperial Cancer Research Fund. [γ-32P]ATP (400 Ci/mmol) and 1-[35S]methionine (400 Ci/mmol) were from ICN Biochemicals (Costa Mesa, CA). Phytomemagglutinin, BSA, poly Glu-Tyr (4:1), poly-l-lysine, cytochalasin D, colchicine, the anti-α-tubulin mAb, and the FITC- and TRITC-conjugated secondary antibodies were all obtained from Sigma (St. Louis, MO). Protein A-agarose and Protein G-agarose were from Boehringer Mannheim (Mannheim, Germany). ECL reagents were from Amersham (Buckinghamshire, United Kingdom). Tyrosine kinases inhibitor genistein was from Calbiochem-Novabiochem (Nottingham, United Kingdom). Interleukin 2 was from Eurocetus (Amsterdam, The Netherlands). Taxol was a kind gift from Dr. J.M. Andreu (Centro de Investigaciones Biológicas, Consejo Superior de Investigaciones Científicas, Madrid, Spain). All other reagents used were of the purest grade available.

Cell Attachment Assays
Cell adhesion assays were essentially performed as described elsewhere (Dransfield et al., 1992b; Luque et al., 1996). Briefly, 96-well flat-bottom plates were precoated overnight at 4°C with 6 μg/ml ICAM-1-Fc in adhesion buffer (20 mM Tris-HCl and 150 mM NaCl, pH 8.2), blocked with 1% BSA in adhesion buffer for 1 h at room temperature, and then washed twice with RPMI 1640. In control nonspecific adhesion experiments, wells were precoated overnight at 4°C with 20 μg/ml poly-l-lysine in water. T lymphoblasts (3 × 10⁵ cells per well) were then added to the wells containing or not the relevant stimuli for LFA-1 activation. When Mg²⁺-EGTA was used to stimulate LFA-1, cells were washed in HEPES-NaCl buffer (20 mM HEPES, 150 mM NaCl, and 2 mg/ml glucose, pH 7.4) instead of RPMI 1640 medium. After 20 min on ice, incubation of plates continued for 60 min at 37°C. Plates were then washed gently four times with prewarmed RPMI 1640, and bound cells were quantitated by measuring the absorbance of wells at 540 nm after fixation and staining with 0.5% crystal violet in 20% methanol.

Digital Confocal Microscopy
For immunofluorescence analysis, round glass coverslips (13 mm diameter) were precoated with recombinant ICAM-1-Fc protein and blocked with 1% BSA as indicated above. T lymphoblasts were washed twice in HEPES-NaCl buffer and allowed to adhere to the ICAM-1-Fc–coated coverslips for 60 min at 37°C. After two washes in PBS, adherent cells were fixed in 3.7% formaldehyde in PBS for 10 min at room temperature and then washed twice in HEPES-NaCl buffer and allowed to adhere to the ICAM-1-Fc–coated coverslips for 60 min at 37°C. After two washes in PBS, adherent cells were fixed in 3.7% formaldehyde in PBS for 10 min at room temperature and then permeabilized in 0.1% Triton X-100 in PBS and 0.5% BSA. Cells were then incubated for 45 min with rabbit anti-FAK (C-20 or A-17) or goat anti-PYK-2 (C-19 or N-19) polyclonal antibodies. Cells were then extensively washed in PBS and 0.5% BSA and incubated for 60 min with the corresponding anti-goat or anti-rabbit FITC-conjugated secondary antibody. In some experiments, the samples were washed again with PBS, double stained with the anti-α-tubulin mAb for 60 min, and washed in PBS and 0.5% BSA, followed by an incubation for 60 min with an anti-mouse TRITC-conjugated secondary antibody. In the experiments in which actin was stained, cells were fixed as above and then treated with TRITC-conjugated phalloidin in PBS and 0.5% BSA for 20 min and washed extensively in PBS. Before mounting the samples for fluorescence microscopy, they were washed again with PBS and distilled water. Confocal microscopy was performed using a MRC-1000 confocal laser scanning system (Bio-Rad, Watford, United Kingdom) connected to a Nikon (Tokyo, Japan) Diaphot 200 inverted microscope. Images of 20 serial vertical cellular sections were acquired every 0.5 μm with the Bio-Rad COMOS graphical user interface and software.
Quantitative Time-Lapse Video Microscopy of T Lymphoblast Motility

Plastic dishes (35 mm) were precoated with recombinant ICAM-1Fc protein and blocked with BSA as indicated above. T lymphoblasts were plated immediately before video recording in RPMI 1640 medium supplemented with 1% FCS in the presence or absence of the relevant stimuli for LFA-1 activation. Time-lapse video films of cells were generated as a sequence of individual digital images (“frames”) that were obtained every 10 s for 2.30 h in an Zeiss (Thornwood, NY) Axiosvert 135 video microscope using the IP-Lab Spectrum software (Signal Analytics, Vienna, VA). The cellular random migration tracks, distances, and average speeds of individual cells for each experimental condition were obtained using the Cell Tracking software extension for IP-Lab Spectrum developed by Tim Hutton (Confocal Microscopy and Digital Image Unit, Imperial Cancer Research Fund).

Immunoprecipitation

T lymphoblasts (100 × 10^6 or 2.5 × 10^6 cells to immunoprecipitate FAK or PYK-2, respectively, unless otherwise stated) were washed twice with RPMI 1640, plated on dishes coated with either BSA or ICAM-1, and, after 15 min on ice, stimulated with 10 μg/ml mAb KIM-127 for 15 min. The stimulation was terminated by solubilizing the cells in 1 ml of ice-cold lysis buffer (10 mM Tris-HCl, pH 7.65, 5 mM EDTA, 50 mM NaCl, 30 mM sodium pyrophosphate, 50 mM NaF, 2 mM sodium orthovanadate, 1% Triton X-100, 50 μg/ml aprotinin, 50 μg/ml leupeptin, 5 μg/ml pepstatin, and 1 mM PMSF). Lysates were clarified by centrifugation at 14,000 rpm for 10 min, and the pellets were discarded. After centrifugation, supernatants were transferred to fresh tubes, and proteins were immunoprecipitated at 4°C overnight with either protein A-agarose–linked rabbit polyclonal anti-FAK antibodies (C-20 or A-17) or protein G-agarose–linked mAbs directed against FAK (2A7 or a-FAK mAbs) or against Tyr(P) proteins (PY20 and PY72 mAbs) or protein G-agarose–linked goat polyclonal anti-PYK-2 antibody (C-19). Immunoprecipitates were washed three times with lysis buffer and either used for in vitro kinase reactions (see below) or extracted in 2× SDS-PAGE sample buffer (200 mM Tris-HCl, pH 6.8, 0.1 mM sodium orthovanadate, 1 mM EDTA, 6% SDS, 2 mM EDTA, 4% 2-mercaptoethanol, and 10% glycerol), by boiling 5 min, fractionated by one-dimensional SDS PAGE, and further analyzed as described in RESULTS and figure legends.

In Vitro Kinase Reactions

Reactions were performed as described (Rodriguez-Fernández and Rozenburg, 1996, 1998). Briefly, immunoprecipitates were washed and pelleted (2500 rpm 10 min in the cold) three times in lysis buffer and twice with kinase buffer (20 mM HEPES and 3 mM MnCl2, pH 7.3). Pellets were dissolved in 40 μl of kinase buffer, and reactions were started by adding 10 μCi of [γ-32P]ATP. The reactions were carried out at 30°C for 15 min and were stopped by ice by adding 10 mM EDTA. After the in vitro kinase reactions, the pellet were washed in lysis buffer containing 10 mM EDTA, extracted for 5 min at 95°C in 2× SDS-PAGE sample buffer, and analyzed by SDS-PAGE. In some experiments poly-Glu-Tyr (4:1; 40 μg) was added to the immunocomplexes. The incorporation of 32P label into poly-Glu-Tyr (4:1) was stopped by removing the supernatant from the agarose beads and adding 2× SDS-PAGE sample buffer. Samples were then analyzed by SDS-PAGE and autoradiography. After fixing and drying of the gels, autoradiography was performed at ~80°C. Autoradiograms were analyzed using an Agfa (Mortsel, Belgium) Studio Scan IIsi scanner, and bands were quantified using the Bio-Rad Molecular Analyst software.

Western Blotting

Cell lysis and immunoprecipitations were performed as described above. After SDS-PAGE, proteins were transferred to Immobilon membranes (Millipore, Bedford, MA) using a Bio-Rad SD Transblot. Membranes were blocked using 3% nonfat dried milk in PBS, pH 7.2, and incubated for 2 h at 22°C with the polyclonal antibodies anti-PYK-2 (C-19 or N-19), diluted 1:500 in PBS containing 3% nonfat dried milk. After incubating membranes with HRP-conjugated secondary antibodies, immunoreactive bands were visualized using ECL reagents.

Metabolic Labeling and Autoradiography

Cells were labeled overnight with [35S]methionine at 100 μCi/ml in methionine-free RPMI 1640 containing 5% FCS, as described previously (Rodriguez-Fernández et al., 1992). Labeled proteins were used to immunoprecipitate FAK with the C-20 antibody. After immunoprecipitation and SDS-PAGE, the gels were soaked in DMSO twice, for 30 min periods, followed by treatment with 19% 2,5-diphenyloxazole (PPO) in DMSO for 2 h. The DMSO-PPO was removed, and the gels were thoroughly washed in water. Gels were dried in a heated gel drier under vacuum and exposed to x-ray films.

RESULTS

Morphological Changes in T Lymphoblasts after Induction of Activation of Integrin LFA-1

In the absence of stimulation the majority of T lymphoblasts grow in suspension and remain as individual cells displaying a rounded appearance. Despite the high expression of both the integrin LFA-1 and its ligand ICAM-1 on the surface of these activated T cells, they do not form large intercellular aggregates, indicating that LFA-1 is in an inactive state unable to mediate ligand interactions. When unstimulated T lymphoblasts are allowed to adhere to immobilized ligand ICAM-1, only a relatively low percentage of these cells are able to attach (Figure 1A, CONTROL). Induction of activation of LFA-1 molecules on T cells using activating mAb KIM-127, Mg2+-EGTA, or the phorbol ester PDBu results in an important increase in the percentage of cells adhering to ICAM-1 (Figure 1A). Interestingly, this augmented adhesion is accompanied by profound changes in the morphology of the adherent T cells. The lymphoblasts changed from their round and moderately spread morphology to an elongated and highly spread phenotype, in which a cell body and a long, and generally unique, cell projection were clearly distinguishable (Figure 1B, ICAM-1). Staining of polymerized actin revealed that after 1 h of adhesion to ICAM-1, actin was redistributed and accompanied by profound changes in the morphology of the adherent T cells. The lymphoblasts changed from their round and moderately spread morphology to an elongated and highly spread phenotype, in which a cell body and a long, and generally unique, cell projection were clearly distinguishable (Figure 1B, ICAM-1). Staining of polymerized actin revealed that after 1 h of adhesion to ICAM-1, actin was redistributed and organized in a thin rim around the nucleus of the cell and, more importantly, in the long cytoplasmic process (Figure 1B, PLL). These adhesion experiments under static conditions, together with the staining of actin in fixed cells, demonstrate that activation of LFA-1 induces important changes in the adhesion and morphology of T lymphocytes.
Figure 1. Agents that activate LFA-1 integrin cause an increment in adhesion and polarization of lymphoblasts. (A) T lymphoblasts plated on ICAM-1-Fc-coated plastic dishes were allowed to adhere for 60 min in the absence (CONTROL) or the presence of different of LFA-1-activating agents: 10 \( \mu \text{g/ml} \) mAb KIM-127 (KIM-127), 10 mM \( \text{Mg}^{2+} \) plus 1 mM EGTA (\( \text{Mg}^{2+}/\text{EGTA} \)), and 50 nM PDBu. Cell adhesion assays were carried out as specified in MATERIALS AND METHODS. The results are the mean \( \pm \) SEM of four independent experiments. (B) T lymphoblasts were plated either on ICAM-Fc-coated dishes (ICAM-1) or on poly-l-lysine-coated dishes (PLL) and stimulated as indicated above. The cells were fixed, permeabilized, and stained with Texas Red-conjugated phalloidin. Immunofluorescence was analyzed by confocal laser microscopy as described in MATERIALS AND METHODS. Arrows indicate the position of the cellular extensions in representative polarized cells.
Kinetics of Morphological Changes and Effects on T Cell Motility of the Activation of LFA-1 Integrin

We used time-lapse digital video microscopy to investigate in detail the kinetics of the morphological changes and the effects that these changes had on the motility of lymphocytes after induction of activation of the LFA-1 molecules. The kinetics of induction of morphological changes was very similar for the three different protocols of activation of LFA-1 used (Mg2+ -EGTA, activating mAb KIM127, and PDBu). The changes in cell morphology were already evident after 20 min, and after 60 min of induction of LFA-1 activation, >90% of T lymphoblasts displayed the characteristic morphological features shown in Figure 1B.

The experiments of video microscopy also revealed that induction of activation of LFA-1 on T lymphoblasts adhering to ICAM-1 results in a switch from a highly motile to a nonmotile phenotype (Figure 2). In the arrested lymphoblasts, the cellular projection extending from the body of the cell seems to be firmly anchored to the substrate, and only the cell body of the lymphoblasts exhibited some lateral motility. Furthermore, quantitation of the distances traveled by the cell bodies of individual lymphoblasts and their average speeds in micrometers per hour confirmed that induction of activation of LFA-1 with PDBu, mAb KIM127, or Mg2+-EGTA results in a striking 13-fold reduction in the motility of these cells, decreasing from almost 1700 μm/h (nonactivated) to 130 μm/h (activated LFA-1) (Table 1).

Inhibition of Morphological Changes Induced by Stimulation of LFA-1 by the Tyrosine Kinase Inhibitor Genistein

To investigate whether tyrosine phosphorylation is required for the observed changes in morphology and level of adhesion of T lymphoblasts induced through activation of LFA-1, we tested the effect of the specific tyrosine kinase inhibitor genistein (Akiyama et al., 1987). As observed in Figure 3A, the addition of genistein at 30 μM to cells plated on dishes coated with ICAM-1 and stimulated with mAb KIM127 inhibited almost completely the morphological changes induced by LFA-1, and the majority of cells remained round shaped (Figure 3A). In contrast with its dra-
matic effects on lymphoblast morphology and polarization, genistein did not significantly affect the adhesion of these cells to ICAM-1 (Figure 3B). These results suggest that tyrosine kinase activity is required for LFA-1–induced changes in the morphology of lymphoblasts.

**LFA-1 Stimulates FAK and PYK-2 Tyrosine Kinase Activity in Human Lymphoblasts**

Because a tyrosine kinase seemed to be involved in the signaling from LFA-1, we examined the effect of stimulation of LFA-1 on the tyrosine kinase activities present in lymphoblasts. Cells plated on dishes coated with ICAM-1 were treated with the activating mAb KIM-127 for 30 min and lysed. Tyr(P) proteins were immunoprecipitated from the cell lysates with the mAb PY20, and the resulting immune complexes were incubated with \[^{32}\text{P}]\text{ATP, analyzed by SDS-PAGE, and subjected to autoradiography. In the immunoprecipitates obtained from KIM-127–treated cells, we observed strong phosphorylation of several bands in the } M_r 110,000–180,000 and 70,000–80,000 ranges (Figure 4, left panel). Results identical to those shown in Figure 4, left panel, were obtained when immunoprecipitates were prepared using a different anti-Tyr(P) mAb (PY72).

To determine whether the tyrosine kinases FAK and PYK-2 were among the \(M_r 110,000–180,000\) bands induced after activation of LFA-1. At the percentage of SDS-PAGE used in this experiment (8%), bands corresponding to FAK and PYK-2 showed very similar electrophoretic mobilities, but at lower SDS-PAGE percentages, PYK-2 migrated at a slightly lower \(M_r \) compared with FAK (Earp et al., 1995).

To confirm directly that LFA-1 stimulates FAK activity, lysates of KIM-127–treated lymphoblasts were incubated with the anti-FAK antibody C-20, and the C-20 immunoprecipitates were incubated with \[^{32}\text{P}]\text{ATP and analyzed by SDS-PAGE. As observed in Figure 5A (IP: C-20), FAK activity increases upon KIM-127 stimulation. No kinase activity was detected when the lysates were immunoprecipitated with nonimmune rabbit serum (Figure 5A, IP: NIS). To further confirm that the radiolabeled 125-kDa band was FAK, we eluted this band by denaturation and reprecipitated the eluted proteins with the mAb a-FAK, an antibody that recognizes an epitope in the kinase domain of FAK (Figure 5A, IP: C-20 and r-IP: a-FAK). The reprecipitation of the 125-kDa radiolabeled band in these experiments confirmed that it is phosphorylated FAK (Figure 5A, IP: C-20 and r-IP: a-FAK). No band was detected when the C-20 immunoprecipitate was reprecipitated with nonimmune rabbit serum (Figure 5A, IP: C-20 and r-IP: NIS). A similar increase in FAK activity after in vitro kinase assays was also observed when lysates of lymphoblasts stimulated with KIM-127 were immunoprecipitated with A-17, a polyclonal antibody that recognizes an epitope in the N terminus of FAK (Figure 5A, IP: C-20 and r-IP: A-17) or with mAb 2A7, an antibody reactive with the C-terminal sequence of FAK. SDS-PAGE and fluorography of FAK immunoprecipitates obtained from \[^{35}\text{S}]\text{methionine metabolically labeled cells prepared in parallel with those used for the kinase assays verified that similar amounts of FAK were recovered before and after KIM-127 treatment (Figure 5A, ML: \(^{35}\text{S}-\)Met and IP: C-20).
Quantitative densitometric scanning showed that activation of LFA-1 induced by treatment of the T lymphoblasts with mAb KIM-127 induced a 2.5-± 0.4-fold (n = 3) and 2.4-± 0.4-fold (n = 4) increase in the phosphorylation of FAK in immunoprecipitates obtained with the C-20 and A-17 polyclonal antibodies, respectively (Figure 5B). An increase in the activity of FAK after KIM-127 treatment of lymphoblasts was demonstrated when lysates from control and KIM-127–stimulated cells were immunoprecipitated with the C-20 antibody, and kinase activity was determined by the ability of the immunoprecipitates to phosphorylate exogenously added substrate poly-Glu-Tyr (4:1) (Figure 5C).

To further investigate the stimulation of PYK-2 activity through LFA-1, lysates of KIM-127–treated lymphoblasts were incubated with the C-19 anti-PYK-2 polyclonal antibody. PYK-2 immunoprecipitates were incubated with [γ-32P]ATP and analyzed by SDS-PAGE. As observed in Figure 6A, in contrast...
to FAK, PYK-2 shows relatively high basal activity in unstimulated cells that is further increased upon LFA-1 activation. No kinase activity was detected when the lysates were immunoprecipitated with nonimmune goat serum (Figure 6A, NIS). Immunoblotting with N-19 antibody or C-19 antibody of PYK-2 immunoprecipitates prepared in parallel with those used for the kinase assays verified that similar amounts of PYK-2 were recovered after KIM-127 treatment (Figure 6A, IP: C-19 and WB: C-19). Densitometric scanning showed that KIM-127 induced a 7.5-6.14-fold (n = 5) increase in the phosphorylation of PYK-2 (Figure 6B). Accordingly, we also observed a strong increase in the ability of PYK-2 immunoprecipitates to phosphorylate the exogenous substrate poly-Glu-Tyr (4:1) when cells were stimulated with KIM-127 (Figure 6C). No increase in FAK or PYK-2 phosphorylation was observed when lymphoblasts plated on BSA or poly-L-lysine were stimulated with 10 μg/ml mAb KIM-127 or when the cells plated on ICAM-1 were stimulated with 10 μg/ml control antibody. Taken together, these results show that both FAK and PYK-2 increased their tyrosine kinase activities upon LFA-1 activation.

**Time Course of LFA-1–stimulated FAK and PYK-2 Tyrosine Kinase Activities**

To study in detail the kinetics of activation of FAK and PYK-2, lymphoblasts plated on dishes coated with ICAM-1 were treated with the activating mAb KIM-127 for different times and lysed. The lysates were incubated with C-20 and C-19 antibodies to immunoprecipitate FAK and PYK-2, respectively. The resulting immune complexes were incubated with [γ-32P]ATP, analyzed by SDS-PAGE, and subjected to autoradiography. As observed in Figure 7, A and B (upper panels), KIM-127 stimulates the activity of FAK with a kinetics different from that observed for PYK-2. The increase in the activity of FAK reached a maximum within 45 min and remained sustainedly at this maximal level for at least 60 min (Figure 7A, upper panel). In contrast, the increase in PYK-2 activity reached a maximum 30 min after stimulation with KIM-127, decreasing at longer times (Figure 7B, upper panel). SDS-PAGE and fluorography of anti-FAK immunoprecipitates of [35S]methionine-labeled cells (Figure 7A, lower panel) and immunoblotting with C-19 antibody of anti-PYK-2 immunoprecipitates (Figure 7B, lower panel), prepared in parallel with those used for the...
kinase assays, confirmed that similar amounts of FAK and PYK-2 kinases were recovered after different times of KIM-127 treatment.

**FAK and PYK-2 Are Redistributed after Activation of LFA-1 and Colocalize with the MTOC**

To study further the possible role of FAK and PYK-2 in the morphological changes observed in lymphoblasts, we examined the localization of these tyrosine kinases in resting and stimulated cells. Lymphoblasts plated on ICAM-1, unstimulated or treated with Mg\(^{2+}\)-EGTA, were stained by double-labeling immunofluorescence with either C-20 rabbit polyclonal anti-FAK antibody or C-19 goat polyclonal anti-PYK-2 antibody and with an anti-α-tubulin mAb, as described in MATERIALS AND METHODS. In unstimulated cells, FAK (Figure 8A; unstimulated, FAK) and PYK-2 (Figure 8B; unstimulated, PYK-2) exhibited a diffuse distribution. Interestingly, in these cells, PYK-2 also localized to the cell–cell contact area (Figure 8B, unstimulated, PYK-2), as reported by others (Sasaki et al., 1995). In cells in which LFA-1 was activated, FAK (Figure 8A; stimulated, FAK) and PYK-2 (Figure 8B; stimulated, PYK-2) are redistributed mainly to the region between the cell body and the stemming projection. Similar staining was observed with the polyclonal antibodies A-17 and N-19, which recognize sequences in the N-termini of FAK and PYK-2, respectively. This characteristic staining was specific, because it was not observed in cells labeled with control nonimmune rabbit [Figure 8A, stimulated, NIS (rabbit)] or goat [Figure 8B, stimulated, NIS (goat)] sera. Interestingly, costaining with an anti-α-tubulin antibody showed that the location of FAK and PYK-2 in stimulated cells coincides with the position of the MTOC.
MTOC in polarized lymphoblasts (Figure 8, A and B, stimulated, α-tubulin). Similar results were obtained when single immunofluorescences for FAK, PYK-2, or α-tubulin were performed, ruling out cross-detection of fluorescence.

**Actin and Microtubular Cytoskeletal Networks Are Involved in the Morphological Changes Induced by Stimulation of LFA-1**

To evaluate the importance of cytoskeletal integrity in the observed changes in cell morphology that take place in lymphoblasts upon stimulation of activation of LFA-1, we studied the effect of different cytoskeletal perturbing agents in this process. Lymphoblasts were pretreated with cytochalasin D (2.5 μM), a drug which disrupts the actin cytoskeleton, colchicine (10 μM), or taxol (1 μM), drugs that disrupt and stabilize the tubulin cytoskeleton, respectively, and the cultures were subsequently stimulated with the mAb KIM-127. At the concentrations used, all the agents exerted only a slight inhibitory effect on the adhesion of lymphoblasts to ICAM-1. However, as shown in Figure 9A, treatment with all these agents blocked the morphological changes in response to the stimulation with KIM-127.

The changes in morphology observed in lymphoblasts upon stimulation of activation of LFA-1 take place concomitantly with a redistribution of FAK and PYK-2 to the MTOC (Figure 8, A and B). We performed immunofluorescence experiments to study whether treatment with cytochalasin D or colchicine had any effect on the distribution of FAK and PYK-2.

---

**Figure 8.** FAK and PYK-2 colocalize with the MTOC in polarized lymphoblasts. Lymphoblasts were allowed to adhere onto ICAM-1-Fc–coated dishes for 60 min either in HEPES-NaCl buffer containing 1 mM Mg2+ and 1 mM Ca2+ (unstimulated) or in HEPES-NaCl containing 10 mM Mg2+ and 1 mM EGTA (stimulated). (A) The cells were fixed, permeabilized, and double stained with C-20 rabbit polyclonal antibody against FAK (FAK) or with a control rabbit nonimmune serum (NIS rabbit) and with an anti-α-tubulin mAb (α-tubulin) as specified in MATERIALS AND METHODS. (B) The cells were fixed, permeabilized, and double stained with C-19 goat polyclonal antibody against PYK-2 (C-19) or with a control goat nonimmune serum (NIS goat) and with an anti-α-tubulin mAb (α-tubulin) as specified in MATERIALS AND METHODS. Arrowheads indicate the positions of the MTOC in representative cells.
Figure 9. Cytoskeletal interfering agents block LFA-1–dependent polarization of lymphoblasts and reduce translocation of FAK and PYK-2 to the MTOC. (A) T Lymphoblasts were washed in RPMI 1640 medium and then pretreated for 4 h with 0.3% DMSO (KIM-127), with 2.5 μM cytochalasin D dissolved in DMSO (KIM-127 + CYT D), with 10 μM colchicine dissolved in DMSO (KIM-127 + COLCH), or with 1 μM taxol dissolved in DMSO (KIM-127 + TAXOL). Cells were subsequently plated on ICAM-Fc–coated dishes and stimulated for 60 min with 10 μg/ml mAb KIM-127. Attached cells were fixed and stained with 0.5% crystal violet in 20% methanol before photography. A representative experiment is shown. (B and C) T Lymphoblasts were washed in RPMI medium and then pretreated for 4 h with 0.3% DMSO (stimulated), with 2.5 μM cytochalasin D (stimulated + CYT D), or with 10 μM colchicine (stimulated + COLCH). After this pretreatment, cells were washed in HEPES-NaCl buffer, plated on ICAM-1-Fc–coated dishes, and stimulated for 60 min with 10 mM Mg^{2+} plus 1 mM EGTA. The cells were fixed, permeabilized, and double stained with C-20 rabbit polyclonal antibody against FAK (B, FAK) or with C-19 goat polyclonal antibody against PYK-2 (C, PYK-2) and with an anti-α-tubulin mAb (B and C, α-tubulin). Immunofluorescence samples were analyzed by confocal laser microscopy as specified in MATERIALS AND METHODS. Arrowheads indicate the position of the MTOC. A representative experiment is shown.
LFA-1–mediated Activation of FAK and PYK-2 Requires Cytoskeletal Integrity

Because cytoskeletal disrupting agents affect the relocation of FAK and PYK-2 to the MTOC in lymphoblasts, we also studied the effect of these agents on the activation of FAK and PYK-2. For this purpose, lymphoblasts were pretreated with cytochalasin D (2.5 μM), colchicine (10 μM), or taxol (1 μM). The cells were subsequently plated on dishes coated with ICAM-1 and treated with the activating mAb KIM-127. As shown in Figure 10, A and B (upper panels), all the agents blocked the increase in FAK and PYK-2 activity in response to the stimulation of LFA-1. Taken together, these results suggest that the reorganization of the actin and tubulin networks that takes place after stimulation of LFA-1 is required for FAK and PYK-2 activation.

DISCUSSION

Polarization of T lymphocytes upon interaction with APCs or during chemokine directed transendothelial migration is a relevant phenomenon that is thought to contribute importantly to the specificity and effectiveness of immune responses (Geiger et al., 1982; del Pozo et al., 1996; Sánchez-Madrid and del Pozo, 1999). Cell polarization involves the conversion from a spherical to an elongated morphology, a process in which important changes in the organization of the cytoskeleton take place. In the present study we report that upon induction of activation of the β2-integrin LFA-1, T lymphocytes undergo a dramatic polarization, characterized by the conversion from a spherical morphology, characteristic of the nonadherent cells, to an elongated morphology, characterized by the presence of a cell body and a long cellular extension. These morphological changes take place as a result of the interaction between activated LFA-1 integrin molecules with ligand ICAM-1, because they are not observed when cells are plated on poly-L-lysine. The induction of activation of the LFA-1 integrin on T lymphoblasts adhering to ICAM-1 also involves the conversion of the cells from a highly motile to a nonmotile phenotype. Furthermore, we demonstrate that binding of T lymphoblasts to the LFA-1 ligand ICAM-1 induces the activation and intracellular redistribution of the tyrosine kinases FAK and PYK-2.

The described changes in the adhesive capacity, morphology, and locomotive behavior in T lymphoblasts upon induction of activation of LFA-1 molecules take place using three different protocols for
induction of activation of LFA-1 molecules, i.e., 1) the presence in the medium of divalent cation Mg$^{2+}$, 2) incubation of cells with the $\beta$-specific stimulatory mAb KIM-127, and 3) the addition of the phorbol ester PDBu. Interestingly, the first two protocols of induction of LFA-1 activation have been reported to affect mainly the affinity of this integrin, whereas the treatment with phorbol esters induces integrin adhesive capacity primarily through changes in avidity (Stewart et al., 1996). In parallel with the polarized morphology, we observed that the lymphocytes acquire a nonmotile phenotype. The cell polarization and the arrest of the migrating T lymphocytes seem to be related phenomena, which contribute to the proper function of T lymphocytes. Thus, after T cell antigen receptor engagement during interaction with APCs, a stop signal is delivered through LFA-1 that converts the migrating T lymphocytes into nonmotile cells. This process appears to be an important requisite for an adequate immune response (Dustin et al., 1997; Lowinsky, Kropf et al., 1998).

To study whether a tyrosine kinase activity was required for the observed LFA-1–mediated increase in T cell adhesion and polarization, we used the specific tyrosine kinase inhibitor genistain. Pretreatment with genistain blocked the polarization of the lymphoblasts with mAb KIM-127, suggesting that the morphological changes induced through the integrin LFA-1 require a tyrosine kinase activity. Genistain, however, had no significant effect on the adhesion of these cells to ICAM-1. Because in these experiments we used the mAb KIM-127 to activate the integrin LFA-1, these results suggest either that adhesion mediated by LFA-1 in lymphoblasts does not require a genistein-sensitive tyrosine kinase activity and/or that the affinity of LFA-1 for ligand ICAM-1 that is induced by KIM-127 treatment is high enough to maintain the cells strongly adhered, even when tyrosine kinases are inhibited.

We demonstrate for the first time that upon activation of LFA-1 and binding of T lymphocytes to ligand ICAM-1, the two related nonreceptor tyrosine kinases FAK and PYK-2 become activated. FAK and PYK-2 immunoprecipitates possess intrinsic autokinase activities and are able to phosphorylate exogenous substrates, such as the tyrosine kinase substrate poly-Glu-Tyr (4:1). Because several studies have demonstrated that the major phosphorylation sites on FAK and PYK-2 (i.e., Tyr-397 and Tyr-402, respectively) are high-affinity binding sites for the SH2 domain of members of the Src kinase family (Richardson and Parsons, 1995), and Src members are able to phosphorylate FAK, it has been suggested that FAK and PYK-2 kinase activities could also reflect the presence of Src in the immunoprecipitates, rather than an intrinsic increase in their activities. Nevertheless, several lines of evidence indicate that this is not the case in our assays.

First, in the lysis buffer in which our experiments were carried out, which contains the nonionic detergent Triton X-100 (see MATERIALS AND METHODS), the interaction between Src and FAK is minimal (Rodríguez-Fernández and Rozengurt, 1998; Schlaepfer et al., 1998). Second, when we used in our in vitro phosphorylation assays pyrazolopyrimidine, a novel specific inhibitor of Src kinases, we did not observe any inhibition in FAK activity at concentrations that virtually abolished Src kinase activity in parallel Src immunoprecipitates (Rodríguez-Fernández and Rozengurt, 1998). Therefore, the previous results strongly suggest that the increases in FAK and PYK-2 tyrosine kinases induced upon activation of integrin LFA-1 cannot be attributed to the presence of Src in the immunoprecipitates.

Several groups have shown that in leukocytes and other cells FAK and PYK-2 become tyrosine phosphorylated and activated upon stimulation of members of the $\beta_1$, $\beta_3$, and $\beta_5$ family of integrins (Kornberg et al., 1991; Guan and Shalloway, 1992; Hanks et al., 1992; Lipfert et al., 1992; Schaller et al., 1992; Shattil et al., 1994; Maguire et al., 1995; Li et al., 1996; Astier et al., 1997; Gismondi et al., 1997; Ma et al., 1997). Furthermore, it has been shown that fakB, a putative homologue of FAK present in lymphocytes, is regulated by the $\beta_2$-integrin LFA-1 (Kanner et al., 1994; Kanner, 1996). However, to our knowledge there are no reports demonstrating activation of FAK and/or PYK-2 upon engagement of a member of the $\beta_2$ family of integrins. Although the precise mechanism by which integrins are able to activate FAK or PYK-2 is not known, the N-terminal region of FAK has been shown to associate in vitro with peptides corresponding to cytoplasmic segments of $\beta_1$-, $\beta_2$-, and $\beta_3$-integrin subunits (Schaller et al., 1995). It has been speculated that the direct interaction with the cytoplasmic domain of the integrins could contribute to the activation of FAK (Schaller et al., 1995).

In polarized T lymphoblasts the activation of FAK and PYK-2 occurs in parallel with a redistribution of these kinases from a relatively diffuse localization in the cytoplasm to a location close to or in association with the MTOC of the lymphocytes. The MTOC is a critical organizing point within the cell, which becomes reoriented during cell polarization, and is thought to participate in the redistribution and concentration of surface molecules in membrane caps, including different adhesion receptors (Dustin et al., 1997; Sánchez-Madrid and del Pozo, 1999). Furthermore, several studies have suggested that the MTOC helps position the T cell secretory apparatus. For example, it was observed that reorientation of the MTOC was followed by the polarized concentration of cytokines and cytotoxic mediators at the T cell–APC surface (Kupfer et al., 1991).
It has been reported that engagement of T cell receptor by physiological ligands delivers a stop signal to lymphocytes migrating on purified ligand ICAM-1 (Dustin et al., 1997). In these experiments, it was observed that the MTOC was repositioned to a location adjacent to the site of stable lymphocyte anchorage, suggesting that the MTOC in the arrested cells could have a role in maintaining the nonmotile phenotype of these cells. We observe in our immunofluorescence and video microscopy experiments that the MTOC in polarized lymphocytes is consistently found at a specific cytoplasmic location, in the region between the cell body and the long cellular extension, which is characterized by strong anchorage to the substrate ICAM-1. These results concur with two recent reports describing that the MTOC localizes to the distal portion of the posterior appendage known as uropod in polarized migrating lymphocytes (Ratner et al., 1997; Serrador et al., 1997). Furthermore, it has been reported that activation of human T lymphocytes induces tyrosine phosphorylation of α-tubulin (Ley et al., 1994; Marie-Cardine et al., 1995). Interestingly, LFA-1-induced activation of PYK-2 was blocked by pretreatment of the cells with agents that interfere with actin and tubulin components of cytoskeleton, suggesting that PYK-2 activation requires cytoskeletal integrity to allow the activation of this enzyme. It is noteworthy that in other systems FAK and PYK-2 activation is blocked with agents that act on actin, but not tubulin, polymerization (Sinnett-Smith et al., 1993; Astier et al., 1997; Rodriguez-Fernández et al., 1998).

In conclusion, in this study we demonstrate that the interaction between activated integrin LFA-1 and its ligand ICAM-1 leads to changes in cell morphology and to the acquisition of a nonmotile phenotype of T lymphocytes. We show, for the first time, that concomitant with these changes, LFA-1 induces activation of the cytoplasmic tyrosine kinases FAK and PYK-2 and redistribution of both kinases to a region close to the MTOC.

ACKNOWLEDGMENTS

We greatly appreciate the generous supply of mAb KIM-127 provided by Dr. Martyn Robinson (Celltech, Slough, United Kingdom) and the generous gift of taxol provided by Dr. J.M. Andreu. We thank Tim Hutton, Peter Jordan, Alex Stokes, and Reiner Pepperkok (Confocal Microscopy and Digital Image Unit, Imperial Cancer Research Fund) for help with the video and confocal microscopy experiments, Reyes Tejedor for help with the preparation of recombinant ICAM-1-Fc, and Dr. Joaquín Teixido for carefully reviewing the manuscript. This work was supported by Dirección General de Investigación Científica y Técnica Promoción General del Conocimiento grant PB94-0231 (to C.C.), Comunidad de Madrid Acciones Coordinadas en Ciencias de la Salud grants 07/044/96 (to C.C. and F.S.M.) and 08/1/0013/97 (to C.C.), a grant from Fundación Científica de la Asociación Española contra el Cáncer (to F.S.M. and C.C.) and Comisión Interministerial de Ciencia y Tecnología grants SAF 96/0039 (to F.S.M.) and SAF 98/0080 (to C.C.). C.C. was also partially supported by the European Molecular Biology Labo-

REFERENCES


