The Tyrosine Kinase PYK-2/RAFTK Regulates Natural Killer (NK) Cell Cytotoxic Response, and Is Translocated and Activated upon Specific Target Cell Recognition and Killing

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Abstract. The compartmentalization of plasma membrane proteins has a key role in regulation of lymphocyte activation and development of immunity. We found that the proline-rich tyrosine kinase-2 (PYK-2/RAFTK) colocalized with the microtubule-organizing center (MTOC) at the trailing edge of migrating natural killer (NK) cells. When polyclonal NK cells bound to K562 targets, PYK-2 translocated to the area of NK–target cell interaction. The specificity of this process was assessed with NK cell clones bearing activatory or inhibitory forms of CD94/NKG2. The translocation of PYK-2, MTOC, and paxillin to the area of NK–target cell contact was regulated upon specific recognition of target cells through NK cell receptors, controlling target cell killing. Furthermore, parallel in vitro kinase assays showed that PYK-2 was activated in response to signals that specifically triggered its translocation and NK cell mediated cytotoxicity. The overexpression of both the wt and a dominant-negative mutant of PYK-2, but not ZAP-70 wt, prevented the specific translocation of the MTOC and paxillin, and blocked the cytotoxic response of NK cells. Our data indicate that subcellular compartmentalization of PYK-2 correlates with effective signal transduction. Furthermore, they also suggest an important role for PYK-2 on the assembly of the signaling complexes that regulate the cytotoxic response.

Key words: CD94/NKG2 • cytotoxicity • microtubule-organizing center • cytoskeletal proteins • HLA-E

Introduction

Cell compartmentalization and rearrangement of membrane and cytosolic proteins seem to be required for lymphocyte activation and the development of immune response (Dustin and Shaw 1999; Lanzavecchia et al., 1999; Sánchez-Madrid and del Pozo, 1999; Serrador et al., 1999). In addition, recent studies on the molecular reorganization that occurs during the specific interaction of lymphocytes with antigen-presenting cells revealed that different molecules involved in T cell stimulation are compartmentalized, favoring the activation of these cells (Monks et al., 1997, 1998; Wülfling and Davis, 1998; Xavier et al., 1998; Viola et al., 1999; reviewed in Dustin and Shaw, 1999).

NK cell cytotoxic activity is modulated by positive and negative signals triggered by different membrane receptors. NK cell receptors specifically interact with molecular histocompatibility complex (MHC) class I on target cells and function either as inhibitory or activatory molecules. Thus, the human killer inhibitory re-

Abbreviations used in this paper: EGFP, enhanced green fluorescent protein; FAK, Focal adhesion kinase; FN, fibronectin; HLA, human leukocyte antigen; IL-2, interleukin-2; ITIM, immunoreceptor tyrosine-based inhibitory motifs; MHC, molecular histocompatibility complex; MTOC, microtubule-organizing center; NK, natural killer; PYK-2, proline-rich tyrosine kinase 2; SHP, Src homology 2 domain containing protein tyrosine phosphatase.

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Receptors from the immunoglobulin superfamily or some members of another family of related Ig-like receptors specifically interact with different human leukocyte antigen (HLA) class I allotypes and suppress NK cell-mediated cytotoxicity (Moreta and Moreta, 1997; Lanier, 1998; Yokoyama, 1998; Long, 1999; López-Botet and Bellón, 1999). In contrast, another set of molecules homologous to the killer inhibitory receptors triggers NK cell activity (Cosman et al., 1997; Samaridis and Colonna, 1997). On the other hand, the CD94 C-type lectin NK cell receptor covalently assembles with distinct members of the NKG2 family, playing different functional roles. CD94/NKG2A constitutes an inhibitory receptor coupled to SHP-1 tyrosine phosphatases through the immunoreceptor tyrosine-based inhibitory motifs (ITIM) bearing NKG2A subunit (Carretero et al., 1997, 1998; Leibson, 1997; Vivier and D’aëron, 1997; Le Drea et al., 1998). In contrast, the association of CD94 with the NKG2C protein, highly homologous to NKG2A but lacking ITIMs, forms a triggering receptor linked to DAP-12 (Houchins et al., 1997; Cantoni et al., 1998; Lanier et al., 1998a,b). Different signaling events such as the rise of intracellular Ca^{2+}, activation of PLC-γ, and release of phosphoinositides, are involved in the activation signaling pathway that ultimately triggers the cytotoxic response, although the precise sequence of these biochemical events has not been fully defined (D’aëron, 1997; Lanier, 1998).

Proline-rich tyrosine kinase 2 (PYK-2), also called RAFTK (related adhesion focal tyrosine kinase), CAK-β (cell adhesion kinase β), CADTK (cell adhesion–dependent tyrosine kinase), or focal adhesion kinase-2 (FAK2), is a member of the FAK non-receptor tyrosine kinase family that is expressed by different cell types, including neural and hematopoietic cells (A vraham et al., 1995; E arp et al., 1995; Lev et al., 1995; Sasaki et al., 1995; Herzog et al., 1996). PYK-2 shares significant sequence homology with FAK (60% identity in the central catalytic domain and 40% identity in both the COOH and NH₂ termini) and does not contain SH2 or SH3 domains, but bears several binding sites for SH2/SH3-containing signaling proteins (A vraham et al., 1995; Lev et al., 1995). PYK-2 may be activated by a variety of stimuli that induce a rise in intracellular calcium concentrations (Lev et al., 1995). In addition, phorbol esters and agonists of G-protein–coupled receptors as well as signals triggered through integrins, may lead to tyrosine phosphorylation of PYK-2 (Lev et al., 1995; D ikić et al., 1996; Li et al., 1996). It has been reported that NK cells express PYK-2, but not FAK, and that β1 and β2 integrin outside-in signaling stimulates PYK-2 activation (G ismondi et al., 1997; Rodríguez-Fernández et al., 1999). Furthermore, it has been shown that PYK-2, together with Src, functions as a link between heterotrimeric G-protein–coupled receptors and the mitogen-activated protein (MAP) kinase signaling pathway (D ikić et al., 1996). Recent studies indicate that PYK-2 may be also involved in regulation of vesicular transport through its interaction with a new GTPase activating protein designated Pap (A ndreev et al., 1999).

In this report, we found that changes in the subcellular localization of the tyrosine kinase PYK-2 occur in an NK cell receptor specifically regulated manner during the effector–target cell conjugate formation. The translocation of PYK-2 from the trailing edge of NK cells to the area of NK-target cell interaction correlated with its activation, suggesting that the compartmentalization of PYK-2 significantly contributes to both the signal transduction in NK cells and the microtubular rearrangements that occur during the cytotoxic process. In addition, overexpression of both a wt and a dominant-negative mutant of PYK-2, but not ZAP-70 wt, blocked the NK cell–mediated cytotoxic response, suggesting an additional role for PYK-2 in the assembly of the signaling complexes that participate in the cytotoxic response.

Materials and Methods

Cells

Interleukin-2 (IL-2)–cultured polyclonal NLK cells were obtained essentially as described (Abramov et al., 1999). In brief, peripheral blood lymphocytes were cultured with irradiated (5 Gy) RPMI 1640 lymphoblastoid cells for 6–9 d in RPMI 1640 supplemented with 10% FCS (complete medium), followed by a negative selection step using an anti-CD3 mAb and rabbit complement (Behring). The CD3+ cells (<5% CD3⁰) were cultured with 50 U/ml of rhIL-2 until use. We routinely obtained a cell population with >95% of CD5⁶ and CD16⁶ cells and <5% of CD3⁰, CD19⁰, or CD14⁰ cells. NK cell clones were established as described (Pérez-Villar et al., 1995). The inhibitory clones were selected by their ability to kill .221 target cells, that was inhibited by HLA-E in .221 A-EH cells. The selected activatory clones showed a low basal lysis of .221 target cells, thus allowing the detection of the lysis activation by HLA-E on .221-A-EH target cells. A cell culture systems were systematically phenotyped by flow cytometry analysis with a panel of NK receptor-specific mAbs, as described (Pérez-Villar et al., 1997). The NKL cell line, kindly provided by M. Roberston (D ana-Farber, Boston, MA; Roberston et al., 1996), was maintained in RPMI 1640 medium supplemented with 10% (vol/vol) heat-inactivated human AB serum, 2 mM glutamine, 1 mM sodium pyruvate, and 100 U/ml IL-2.

The HLA class I–deficient EBV-transformed B lymphoblastoid cell line 721.221 (.221) was grown in complete medium. The 721.221 cell line transfected with the A EH chimeric construct (.221-A EH), in which the HLA-E leader sequence is replaced by that of HLA-A2 was kindly provided by Dr. G. Schäffer (Lee et al., 1998).

Antibodies and Reagents

The anti-CD56 mAb (C218) was kindly provided by Dr. A. Moreta (Istituto Nazionale per la Ricerca sul Cancro e Centro Biotecnologie A varnale, University of Genova, Genova, Italy). The anti-CD94 (H-P 38B1) and the anti-HLA-class I (H-P 37F7) mAbs have been described (Pérez-Villar et al., 1995). PYK-2 affinity-purified anti-peptide polyclonal antibodies C-19 and N-19, as well as the C-19 antibody cognate blocking peptide were from Santa Cruz Biotechnology, Inc. The anti-y-tubulin and antitalin were purchased from Sigma-Aldrich. The anti-paxillin mAb was from Transduction Laboratories, and the anti-Tyr(p) 4G10 was from Santa Cruz. The anti-moesin 3B8/7 was kindly provided by Dr. R. Schwartz-A Kalbe (German Cancer Research Center, Heidelberg, Germany) and has been described (Lankes et al., 1988). The anti-p37 15B6 mAb was kindly provided by Dr. R. Blasco (Instituto Nacional de Investigaciones A garías, Madrid). The anti-ZAP-70 701 polyclonal antibody was kindly provided by Dr. R. Blasco (Instituto Nacional de Investigaciones A garías, Madrid). The anti-ZAP-70 701 polyclonal antibody was kindly provided by Dr. R. Blasco (Instituto Nacional de Investigaciones A garías, Madrid).

Immunofluorescence Analysis

NK cells (5 × 10⁶) were added to coverslips coated with 30 μg/ml fibronectin (FN) in flat-bottomed, 24-well plates (Costar Corp.) in a final volume of 500 μl complete medium and allowed to settle for 30 min at 37°C in a 5% CO₂ atmosphere. Then, cells were either fixed and permeabilized with 0.5% Triton, 2% formaldehyde, PBS or fixed in 2% formaldehyde, and then permeabilized in 0.1% saponin, 1% BSA, 0.005% NaN₃, PBS, and
stained with appropriate antibodies as previously described (Nieto et al., 1998).

For studies of cell conjugates, 1 × 10^5 target cells were added to cover-slips coated with 30 μg/ml FN in flat-bottomed, 24-well plates (Costar Corp.) in a final volume of 500 μl complete medium and allowed to settle 30 min at 37°C in a 5% CO₂ atmosphere. Nonadherent cells were aspirated and 1 × 10^5 NK cells in 500 μl complete medium were then added and incubated for 1 h at 37°C in a 5% CO₂ atmosphere. Cells were then fixed and permeabilized as previously indicated, stained with the indicated antibodies, and analyzed using a Nikon Labophot-2 photomicroscope with a 100× oil immersion objective. Confocal microscopy was performed as described (Rodríguez-Fernández et al., 1999) using a MRC-1024 Confocal Laser Scanning System (Bio-Rad Labs.).

**Cytotoxic Assays**

Microcytotoxicity chromium (⁵¹Cr) release assays were performed as described (Pérez-Villar et al., 1997), using 721.221 and 221-AEH target cells. The cytotoxic effect of anti-CD94 mAb (for effectors) or anti-HLA mAb (for targets) was tested by preincubating cells for 10 min before performing the cytotoxicity assay. The percent of specific lysis was calculated (Aramburu et al., 1990) and only those experiments with a spontaneous release under 20% of the maximum ⁵¹Cr release were considered.

**Cell Lysates and Immunoprecipitation Assays**

The NKL cell line and the target cell lines (.221 and .221-A EH) were washed twice with RPMI 1640. Experiments were initiated by mixing 10^5 effector cells and 2 × 10^5 of each target cells (.221 or .221-A EH) in a final volume of 200 μl. When indicated, target cells were previously fixed in 1% paraformaldehyde and washed extensively before incubating with NK cells. Cells were spun, incubated at 37°C for the indicated times, and lysed in 1 ml of ice-cold lysis buffer (10 mM Tris-HCl, pH 7.65, 5 mM EDTA, 150 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 phenylmethylsulfonyl fluoride). Cell lysates were clarified by centrifugation at 14,000 rpm for 10 min, and supernatants were immunoprecipitated at 4°C overnight with protein G–agarose coupled to goat polyclonal anti-PYK-2 (C-19) antibodies. Immunoprecipitates were washed three times with lysis buffer, and either used for in vitro kinase assays or Western blot (see below).

**In Vitro Kinase Assay**

This assay was performed as described (R. Rodriguez-Fernández and R. Renzengurt, 1998). Briefly, immunoprecipitates were washed three times in lysis buffer and twice with kinase buffer (20 mM Heps, 3 mM MnCl₂, pH 7.35), pelleted dissolved in 40 μl of kinase buffer, and reactions initiated by adding 10 μCi of γ⁻³²P]-ATP. The reactions were carried out at 30°C for 15 min, and stopped by transferring to ice and adding 10 mM EDTA. Pellets were then 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. A utoradiograms were processed using an AGFA Studio Scan IIsi scanner and bands were quantified using the Biorad Multi-Analyset Software.

**Western Blot**

After SDS-PAGE, immunoprecipitated proteins were transferred to Immobilon membranes using a Bio-Rad SD Transblot. Membranes were then blocked with 3% nonfat dried milk in PBS, pH 7.2, and incubated for 2 h RT with anti–PYK-2 (C-19 or N-19) polyclonal antibodies at a 1:500 dilution, anti PYTyr 4G10 mAb at 1:1000, and anti-p37 15B6 polyclonal antibody at a 1:50 dilution in PBS containing 3% nonfat dried milk. Bound antibodies were detected with horseradish peroxidase conjugate secondary antibodies, followed by visualization with ECL reagents.

**Recombinant Vaccinia Viruses and Vaccinia Virus Infection of NKL Cell Line and NK Clones**

To generate recombinant vaccinia viruses encoding the wt and the kinase dead form of PYK-2, the full-length PYK-2 and PYK-2-K457A (PYK-2-K-M) cDNA's (Li et al., 1996) were subcloned as a HindIII-XbaI fragment into the pRB21 vector (kindly provided by Dr. J. A. Molerio, Instituto de Biología Fundamental, Instituto de Salud Carlos III, Madrid). The full-length ZA P-70 wt cDNA in EcoRI site of pSV10.1 (kindly provided by Dr. B. A. larcón) was subcloned in EcoRI site of pRB21. Then, the recombinant pRB21 vector and the constructs containing ZA P-70 wt, PY K-2 wt and PY K-2 (K-M) in the correct orientation were inserted into the defective vR B12 strain of vaccinia virus (Blasco and Moss, 1995) by homologous recombination. Selection of recombinants was carried out by plaque formation as described (Blasco and Moss, 1995). Viruses were then cloned, purified, and standard viral plaque assays on CV1 cells were used to determine the titer of the constructs. To generate PY K-2-EGFP fusion proteins, PY K-2 cDNA's were subcloned as a HindIII-XbaI fragment into the pEGFP-C3 vector (CLONTECH Laboratories, Inc.), and then the N hel-XbaI fragment subcloned in the pRB21 vector. ZA P-70-EGFP (kindly provided by Dr. B. A. larcón) was subcloned as an EcoRI-XbaI fragment into the pRB21 vector. As described before, viruses coding EGFP, EGF ZA P-70 wt, EGFPP K 2 wt, and EGFPP KY K-2 (K-M) fusion proteins were generated.

NKL cell line and NK clones were infected by recombinant vaccinia viruses essentially as described previously (Wagmann et al., 1999). In brief, NKL cells and NK clones were washed three times with infection medium (1% serums, 0.5% BSA, 1× nonessential amino acids, 2 mM l-glutamine) to room temperature and resuspended at 10⁶ cells/ml in infection medium. Cell suspensions were given either no virus or recombinant vaccinia virus (control pRB21, ZA P-70 wt, PY K-2 wt, KY K-2 (K-M), or the EGFPP fusion proteins) at 20 pfu/cell and incubated 2 h at 37°C with occasional stirring. After one wash in complete RPMI, the vaccinia-infected or uninfected control cells were simultaneously plated for standard ⁵¹Cr-release killing assays, for Western blot, and for immunofluorescence analysis as described. The level of infection was monitored by measuring the expression of the viral envelope p37 protein in the infected cells by Western blot.

**Results**

**PYK-2 Colocalizes with the MTOC in Migrating NK Cells**

IL-2–activated NK lymphocytes show a high degree of cell polarization, in accordance with their high locomotive capability (Somersalo, 1996–1997; Nieto et al., 1998). Immunofluorescence studies were performed to determine the cytoplasmic distribution of PYK-2 in IL-2–activated NK cells adhered and migrating onto FN. Confocal microscopy analysis revealed that PYK-2 is highly concentrated in a specific location in the cytoplasm of both rounded and polarized NK cells adhered to FN (Fig. 1 a, A and B). In polarized NK cells, the majority of PYK-2 was concentrated at the uropod, the cytoplasmic projection developed at the rear of polarized cells. Double staining of cells for PY K-2 and moesin, a member of the ezrin-radixin-moesin family of proteins that connects the cell membrane to the actin-cytoskeletal network, and that marks the cytoplasmic membrane of the uropod (Serrador et al., 1997), confirmed that PY K-2 localizes in the protrusion of migrating, polarized NK cells. Confocal microscopy of migrating, polarized NK cells. Confocal sections of cells double-stained for γ-tubulin, a microtubule protein that is a highly conserved component of the MTOC (Sterns et al., 1991; Zheng et al., 1991), and for PY K-2, demonstrated that this tyrosine kinase colocalizes with γ-tubulin, and thus with the MTOC, at the uropod of polarized NK cells adhered to FN (Fig. 1 b).
Figure 1. PYK-2 is localized at the uropod projection of NK cells adhered to fibronectin. (a) Confocal section of PYK-2 immune staining of NK cells migrating on FN (A). The same section under brightfield illumination is shown in B. Epifluorescence images of NK cells double-stained for PYK-2 (green) and moesin (red, C), and PYK-2 (green) and talin (red, D). Control cells stained with goat serum or secondary antibody alone did not show any specific staining, and preincubation of the anti-PYK-2 antisera with its specific blocking peptide prevented staining of NK cells (not shown). (b, A) Confocal serial sections of polarized NK cells adhered to FN and double-stained for γ-tubulin (red, upper panels, γ-TUB) and PYK-2 (green, lower panels, PYK-2). Sections were taken every 0.8 μm from the substratum (0) in the z-axis. B shows a merged image of double-labeled cells demonstrating the colocalization of γ-tubulin and PYK-2 at the uropod of NK cells.
PYK-2 Is Translocated in NK Cells to the Area of Cell-to-Cell Contact during Specific Recognition of Target Cells

To assess whether PY K-2 is translocated during the killing process, we investigated the cellular localization of PY K-2 in NK–target cell conjugates. In a first set of experiments, K 562 cells were used as targets since these cells do not express classical MHC class I molecules and are efficiently killed by IL-2–activated polyclonal NK cells. As it has been described (Nieto et al., 1998), asymmetrical binding of NK cells to target cells through the NK cell leading edge was observed, with the uropod projecting away from the area of cell–cell contact (Fig. 2). When NK–K 562 cell conjugates were stained with anti–PY K-2 polyclonal antibodies, it was found that this kinase concentrated in a specific domain of the effecter cells that faces the site of adhesion with the target cell (Fig. 2). In addition, in some effector cells, PY K-2 localized to cell-to-cell contact areas. In contrast, no staining for PY K-2 was detected at the cell uropod, developed at the opposite site of the contacting region. Therefore, PY K-2 is translocated from the rear of the cell to the area of cell contact during effector–target cell conjugate formation.

The triggering receptors for lysis of K 562 cells have not been defined yet. To ascertain whether the translocation of PY K-2 in NK cells was related to the recognition of target cells through specific receptors, the positioning of PY K-2 was determined during the interaction of distinct NK cell clones with target cells that show a different sensitivity to be killed. To this end, we have used two different types of NK cell clones that express CD 94/NKG2C or CD 94/NKG2A heterodimers that interact with HLA-E. Fig. 3 shows the cytotoxic activity of NKG2A+ and NKG2C+ NK cell clones against target cells expressing (.221-A EH) or not (.221) HLA-E. NK G2A+ cell clones showed an efficient cytotoxicity against .221 cells, but not to .221-A EH. The protection conferred by HLA-E was reversed by blocking mAbs against both CD 94 and MHC I molecules, demonstrating that the inhibitory signal of cytotoxicity was induced through the specific interaction of these molecules. In contrast, the lytic ability of the NK clones that express the CD 94/NKG2C is enhanced by .221-A EH HLA-E-bearing cells (Borrego et al., 1998; Braud et al., 1998; Lee et al., 1998; Llano et al., 1998). Hence, we selected some activatory clones whose basal lysis against .221 was low, thus allowing the detection of the enhancing effect by HLA-E (Fig. 3). In this case, mAbs against CD 94 and MHC I molecules blocked the activation signal induced by HLA-E (Fig. 3).

NKG2A+ and NKG2C+ cell clones were allowed to interact with .221 or .221-A EH target cells and then PY K-2 was stained in effector–target cell conjugates, and analyzed by confocal microscopy. In conjugates of inhibitory NKG2A+ cell clones with HLA-E–bearing target cells (.221-A EH), PY K-2 was clearly located at the cell uropod. In contrast, PY K-2 was found facing the area of cell–cell interaction in NK cells when .221 target cells did not express HLA-E (Fig. 4 a, left panels, and Table I). The correlation between the triggering of cytotoxic activity and...
Figure 4. PYK-2 and the MTOC are translocated to the area of cell–cell contact of NK cells with target cells during specific recognition and cytotoxicity. (a) Confocal sections showing PYK-2 localization on NK cell clones (E) which are inhibited (CD94/NKG2A), or activated (CD94/NKG2C) by HLA-E, and that were interacting with HLA-E⁻ (221) or HLA-E⁺ (AEH) target cells (T). Cell–cell conjugates were stained for PYK-2 (green) and CD94 (red) and analyzed using confocal microscopy. Confocal sections under brightfield illumination are shown for each conjugate. Arrowheads show the localization of PYK-2 on effector cells. (b) MTOC localization on NK cell clones that are inhibited (CD94/NKG2A), or activated (CD94/NKG2C) by HLA-E, and that were interacting with HLA-E⁻ (221) or HLA-E⁺ (AEH) target cells. Cell conjugates stained for γ-tubulin were photographed under epifluorescence (red) and brightfield conditions using a Nomarski 60× objective. T and E indicate target and effector cells, respectively, while arrowheads point to the MTOC on effector cells.
translocation of PYK-2 was further confirmed in reciprocal experiments using activatory NKG2C-expressing clones. In this case, PYK-2 was mainly localized at the NK cell uropod in cell conjugates formed with 221 cells, whereas it was translocated to the cell-to-cell contact area when cytotoxicity was triggered by recognition of HLA-E on 221-AEH cells (Fig. 4, right panels, and Table I).

Changes in the Distribution of Cytoskeletal Components during NK Cell Receptor–mediated Effector–Target Cell Interaction

It has been described that the MTOC of cytotoxic T lymphocytes is translocated to face the area of cell adhesion with the target cell during the cytotoxicity process (Kupper and Singer, 1989). Therefore, it was of interest to analyze the effect of NK cell–specific cytotoxic triggering in the cellular positioning of some cytoskeletal components. We then characterized the MTOC translocation triggered by CD94/NKG2-specific recognition. In accordance with their cytotoxic responses, the translocation of the MTOC observed when NKG2A− clones interacted with 221 targets was inhibited when killing was prevented by HLA-E in 221-AEH cells (Fig. 4, left panels, and Table I). Conversely, an efficient translocation of the MTOC was observed on CD94/NKG2C NK cell clones interacting with 221-AEH target cells. The frequency of MTOC translocation was very low when these clones interacted with 221 target cells (Fig. 4, right panels, and Table I). Thus, the triggering of cytotoxic activity in these clones correlated with the translocation of MTOC to the area of interaction with sensitive target cells.

It has been recently reported that PYK-2 is constitutively associated with paxillin in NK cells (Gismondi et al., 1997). Accordingly, we found that paxillin colocalized with PYK-2 at the site of MTOC, and that both paxillin and PYK-2 were translocated to the cell–cell contact area when NK cells were bound to sensitive, but not to resistant target cells (Fig. 5). In contrast, no changes in the subcellular localization of other cytoskeletal components such as moesin, localized at the NK cell uropod, or talin, which participates in integrin-mediated cell adhesion and localizes at cell-to-cell boundaries areas, were observed upon triggering of killing (Fig. 5). These results demonstrate that PYK-2 and paxillin are specifically translocated, correlating with microtubular rearrangements, to the area of NK cell–target cell interaction in response to NK cell receptor–mediated specific signals that modulate cytotoxicity.

PYK-2 Activation during Cytotoxic Activity of NK Cells

The selective translocation of PYK-2 during target cell killing suggests that this tyrosine kinase is activated as a consequence of the signals involved in the triggering of cytotoxicity. To assess this possibility, we analyzed the activation of PYK-2 during killing by an in vitro kinase assay using as effector the CD94/NKG2A expressing NKL cell line. The cytotoxic response and the translocation of the MT OC and PYK-2 in this cell line are similar to the CD94/NKG2A clones described above, and they are also inhibited upon recognition of HLA-E (data not shown). To analyze the kinase activity of PYK-2 during cytotoxicity, this kinase was immunoprecipitated from cell lysates of mixtures of NKL cells with either 221 wild-type cells or cells expressing HLA-E (.221-AEH). Immunoprecipitates were incubated with γ[32P]ATP and analyzed by SDS-PAGE. The results obtained revealed a statistically significant difference between the activation state of PYK-2 in the two distinct effector–target cell conjugates (Fig. 6, a and b). Correlating with the triggering of cytotoxic activity (65.7 ± 5.6% specific lysis), PYK-2 was clearly activated when NKL cells interacted with wild-type–sensitive 221 cells. In contrast, a significant lower change in the activation state of PYK-2 was observed when NKL cells interacted with 221-AEH cells, which trigger an inhibitory signal resulting in a lower cytotoxic activity (27.5 ± 3.8% specific lysis). Immunoblotting with anti–PYK-2 C-19 antibody immunoprecipitates carried out in parallel verified that similar amounts of PYK-2 were recovered in each case (Fig. 6, a and b). Kinase activity was not detected when the PYK-2 antibody was preincubated with the immunizing peptide before the addition to the lysates or when the lysates were immunoprecipitated with a nonspecific goat serum (results not shown). Kinetics studies showed that the increase in the kinase activity of PYK-2 during cytotoxicity reached a maximum from 25 to 45 min (Fig. 6c). The possible contribution of PYK-2 protein from target cells was ruled out when the same experiments performed with fixed target cells rendered similar results (data not shown). Fixation of target cells did not affect their capability to induce translocation of PYK-2 or MT OC on NK cells (data not shown). These results indicate that PYK-2 is activated in response to NK cell receptor–driven specific signals that lead to NK cell–mediated cytotoxicity.

Overexpression of PYK-2 in NK Cells Inhibits Cytotoxicity and Translocation of MT OC and Paxillin

To further assess the role of PYK-2 in the cytotoxic response, we used a vaccinia virus system to overexpress the wt and a kinase dead mutant of PYK-2 (PYK-2 K M) in the NKL cell line. Immunoblotting analyses with anti–PYK-2 confirmed the overexpression of the wt and the kinase dead mutant of PYK-2 (Fig. 7a). As expected, overexpression of PYK-2 wt resulted in a substantial increase in the PYK-2 autokinase activity in immunoprecipitates from lysates of NKL-infected cells. In contrast, PYK-2 immunoprecipitates from cells infected with the kinase dead mutant PYK-2 K-M did not show any kinase activity and.

| Table I. Frequency of Translocation of PYK-2 and MT OC in NK Cells Interacting with Target Cells |
|--------------------------------------------------|
|                     | CD94/NKG2A (INH) | CD94/NKG2C (ACT) | CD94/NKG2A (INH) | CD94/NKG2C (ACT) |
| .221                 | 78.7 ± 13.5      | 32.6 ± 8.9       | 75.5 ± 12.8      | 30.1 ± 7.5       |
| AEH                  | 23.1 ± 8.3       | 81.2 ± 15.4      | 19.2 ± 7.4       | 82.7 ± 17.9      |

NK cells bearing the activatory (CD94/NKG2C), or inhibitory (CD94/NKG2A) receptors for HLA-E were allowed to interact with .221 (HLA-E−) or .221-AEH (HLA-E−) target cells on FN-coated plates. After 1 h, cell conjugates were fixed, and stained for PYK-2 or γ-tubulin. Translocation of PYK-2 or MT OC was measured in more than 100 conjugates in three independent experiments. Results correspond to the arithmetic mean ± SD.
instead, it caused a significant inhibition of the basal kinase activity of endogenous PYK-2 (Fig. 7 a), as previously described (Sieg et al., 1998; Kumar et al., 1999). No significant effect on the kinase activity of PYK-2 was observed when cells were infected with the control VV pRB21 at the same pfu/cell. Equivalence of the level of infection between the vaccinia viruses containing different constructs was assessed by analysis of the expression of the viral p37 protein (Fig. 7 a).

To assess the specificity of the effect of PYK-2 overexpression on NK cell function, we generated ZAP-70 wt viruses overexpressing an equivalent level of this tyrosine kinase, compared to the overexpression of PYK-2 wt (Fig. 7 b). The ZAP-70 kinase has been previously described not to affect cytotoxicity (Brumbaugh et al., 1997). We then analyzed the cytotoxic activity of the NKL infected with the different vaccinia viruses against the .221-sensitive target. Interestingly, the killing was inhibited by the overexpression of both the wt and the kinase dead PYK-2, as compared to the slight effect of the infection with control pRB21 vaccinia virus and the tyrosine-kinase ZAP-70 (Fig. 7 c). In addition, the overexpression of another kinase, the serine-threonine kinase PAK-1, did not affect cytotoxicity, further confirming the specificity of the effect of PYK-2 on NK cell-mediated killing (data not shown).

To explore the mechanism by which overexpression of PYK-2 inhibits the cytotoxic activity, we studied the subcellular localization of the different forms of PYK-2-EGFP fusion proteins. Vaccinia viruses were used to overexpress PYK-2 wt-EGFP, PYK-2 (K-M)-EGFP, ZAP-70-EGFP, and EGFP fusion proteins. Inhibitory clones (CD94/NKG2A) were infected and allowed to interact with .221 target cells, subcellular localization of MTOC, and paxillin studied by immunofluorescence analysis (Fig. 8 a). PYK-2 wt-EGFP and (K-M)-EGFP showed a broad distribution throughout

Figure 5. Subcellular localization of different cytoskeletal proteins in NK/target cell conjugates. Cytoplasmic localization of PYK-2 (green) and the cytoskeletal proteins moesin, talin, and paxillin (red) in cell conjugates formed by NK cell clones (E) which are inhibited by HLA-E<sup>+</sup> (INH-AEH) but not by HLA-E<sup>-</sup> (INH-.221) target cells (T). Arrowheads indicate staining of the protein indicated at each panel.
the cell cytoplasm, with a higher concentration colocalizing with the MTOC and paxillin, whereas EGFP and ZAP-70-EGFP were found uniformly distributed in the cell cytoplasm (Fig. 8a, and not shown). As expected, efficient translocation of the MTOC to the area of contact with target cells was observed on cells overexpressing EGFP or ZAP-70-EGFP, and only a slight reduction on the frequency of MTOC translocation was observed as compared to uninfected cells (Fig. 8a and b). Interestingly, overexpression of both the fusion proteins PYK-2 wt-EGFP and PYK-2 (K-M)-EGFP inhibited the translocation of MTOC and paxillin in response to the sensitive .221 target cell (Fig. 8a and b). Given the well-known interactions of PYK-2 with several regulatory molecules...
These results suggest that PYK-2 has an important additional role as a scaffolding protein, whose overexpression deregulates the balance of signaling complexes that control the killing response, probably by interfering with the microtubular rearrangements involved in the translocation and orientation of the secretory apparatus.

Discussion

We report herein the changes in the subcellular localization of the tyrosine kinase PYK-2 in NK cells and its activation during the effector–target cell conjugate formation and the triggering of cytotoxic activity. In addition, we provide evidence on the role of PYK-2 in the regulation of cytoskeletal rearrangements, required for NK cell-mediated killing activity. It is now becoming evident that in addition to other regulatory elements such as phosphorylation or aggregation of molecules, protein compartmentalization provides an additional modulatory mechanism in the cell (Mochly-Rosen, 1995; Faux and Scott, 1996; Pawson and Scott, 1997). Subcellular positioning of signaling molecules can be accomplished by their association with cytoskeletal scaffolds or, as it has been recently demonstrated, by their preferential localization into specific cell membrane domains (Dustin and Shaw, 1999). The selective confining of proteins into specific cell domains has been shown to be of particular importance during the immune response (reviewed in Dustin and Shaw, 1999; Lanzavecchia et al., 1999; Penninger and Crabtree, 1999; Serrador et al., 1999). During the T cell response, the concentration of several proteins, including the T cell receptor molecules, into distinct segregated domains, is necessary to accomplish T cell activation. Interestingly, the protein kinase C-β and Src-
kinase family members Lck and Fyn colocalize with the T cell receptor complex at the area of cell–cell contact during antigen presentation (Montes et al., 1997, 1998). Additionally, these observations suggest that protein compartmentalization, via recruitment and inclusion of activating signaling molecules and exclusion of potential negative regulators, may provide a unique restricted subcellular microenvironment that promotes optimal signal transduction. By this way, the cytoskeleton can directly regulate, both temporally and spatially, the molecular dynamics of signaling pathways.

The localization of PYK-2 at the site of cell contact during conjugate formation could be explained by its association with the MTOC, but such phenomenon would also imply the topological restricted regulation of signaling events at the site of intercellular contacts such as the aforementioned vesicle secretion or cell adhesion. In this regard, it has been shown that PYK-2 is involved in the outside-in signaling mediated by integrins, which are involved in NK cell adhesion to their targets, and act as co-stimulatory receptors during NK cell function (Aster et al., 1997; Gismondi et al., 1997; Ma et al., 1997). In addition, it is worth mentioning that a small fraction of PYK-2 is located at the advancing front of some polarized NK cells, the site that nucleates cell adhesion. Furthermore, PYK-2 was observed predominantly concentrated in the area of cell adhesion in polygonal NK-target cell conjugates. The translocation of PYK-2 to the site of contact with target cell upon the triggering of cytotoxic activity suggests that this enzyme participates in the signaling mediated by integrins during NK cell activation. In agreement with this possibility, we found that PYK-2 colocalizes with paxillin at the site of MTOC and that both proteins are translocated to the cell–cell contact area of NK–target cell–specific conjugates during the cytotoxic process. In this regard, it has been recently reported the constitutive association of PYK-2 with paxillin, which has been largely implicated in the formation of adhesion structures (Burrage et al., 1997; Gismondi et al., 1997).

However, several pieces of evidence indicate that integrin signaling does not seem to account by itself for the activation and translocation of PYK-2 observed after the triggering of cytotoxic activity by sensitive target cells. First, the interaction of NK cells with FN through β1 integrins does not induce the translocation of PYK-2 and the MTOC. Second, we found adhesion of NK cells, likely mediated by β2 integrins, to both sensitive and resistant target cells, but the translocation and activation of PYK-2 was only induced by targets that elicited NK cell cytotoxicity. Furthermore, PYK-2 translocation was induced upon its overexpression, thus resulting in nonfunctional signaling complexes. This possible role of PYK-2 as an adapter molecule, independently of its kinase activity has also been suggested for FA K to explain the effects of its overexpression on the motility of CHO cells (Cary et al., 1996). Our finding on the prevention of the rearrangement of microtubules and the translocation of MTOC upon the overexpression of PYK-2 supports this view. However, the issue of the possible function of the kinase activity of PYK-2 during cytotoxicity, which may play a role since PYK-2 becomes activated as a consequence of cytolytic signals, remains uncertain. In this regard, a recent report described that overexpression of wild-type SHP-1 blocks tyrosine phosphorylation of PYK-2 (Kumar et al., 1999). This tyrosine phosphatase has been involved in KIR-mediated inhibition of cytotoxicity (Binstadt et al., 1996), and it would be feasible to inhibit this inhibitory effect via SHP-1 phosphatases on cytotoxicity could be mediated by PYK-2 inhibition. In addition, the kinase activity of PYK-2 could be important for its adapter role, since the association of important kinases which would be involved in cytotoxicity, such as Src, is dependent on the autophosphorylation of the Tyr402 of PYK-2 (Schlaepfer et al., 1999). The precise mechanism of action of PYK-2 in NK cell cytotoxicity deserves further research.

In summary, our data indicate that the specific recognition of target cell by NK cell receptors directs the translocation and activation of PYK-2, paxillin, and MTOC from
the uropod to the cell-cell contact area. In addition, PYK-2 regulates NK cell cytotoxic activity. These findings further support the important role of cell compartmentalization in the key functions of immune cells.

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Note Added in Proof. In a recent paper, another group has also shown a functional role for PYK-2 in NK cell cytotoxicity (Gismondi, A., J. Jacobson, et al. Science 283:484-486).

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