Isoform-selective phosphoinositide 3'-kinase inhibitors inhibit CXCR4 signaling and overcome stromal cell–mediated drug resistance in chronic lymphocytic leukemia: a novel therapeutic approach

Matthias Niedermeier,1 Bryan T. Hennessy,2 Zachary A. Knight,3 Marina Henneberg,1 Jianhua Hu,4 Antonina V. Kurtova,1 William G. Wierda,1 Michael J. Keating,1 Kevan M. Shokat,3 and Jan A. Burger1

Departments of 1Leukemia and 2Systems Biology and Gynecologic Medical Oncology, University of Texas M. D. Anderson Cancer Center, Houston; 3Department of Cellular and Molecular Pharmacology, Howard Hughes Medical Institute, University of California, San Francisco; and 4Department of Biostatistics, University of Texas M. D. Anderson Cancer Center, Houston

Phosphoinositide 3-kinases (PI3Ks) are among the most frequently activated signaling pathways in cancer. In chronic lymphocytic leukemia (CLL), signals from the microenvironment are critical for expansion of the malignant B cells, and cause constitutive activation of PI3Ks. CXCR4 is a key receptor for CLL cell migration and adhesion to marrow stromal cells (MSCs). Because of the importance of CXCR4 and PI3Ks for CLL-microenvironment cross-talk, we investigated the activity of novel, isoform-selective PI3K inhibitors that target different isoforms of the p110-kDa subunit. Inhibition with p110α inhibitors (PIK-90 and PI-103) resulted in a significant reduction of chemotaxis and actin polymerization to CXCL12 and reduced migration beneath MSC (pseudoepipodesis). Western blot and reverse phase protein array analyses consistently demonstrated that PIK-90 and PI-103 inhibited phosphorylation of Akt and S6, whereas p110δ or p110δ/p110δ inhibitors were less effective. In suspension and MSC cocultures, PI-103 and PIK-90 were potent inducers of CLL cell apoptosis. Moreover, these p110α inhibitors enhanced the cytotoxicity of fludarabine and reversed the protective effect of MSC on fludarabine-induced apoptosis. Collectively, our data demonstrate that p110α inhibitors antagonize stromal cell–derived migration, survival, and drug-resistance signals and therefore provide a rationale to explore the therapeutic activity of these promising agents in CLL. (Blood. 2009;113:5549-5557)

Introduction

Chronic lymphocytic leukemia (CLL), the most prevalent form of adult leukemia in Western countries, is characterized by the progressive accumulation of phenotypically mature, monoclonal B lymphocytes in the peripheral blood, lymph nodes, and bone marrow. These long-lived CLL B cells are mostly arrested in the G0/G1 phase of the cell cycle and display features consistent with a defect in programmed cell death (apoptosis), such as overexpression of Bcl-2-family proteins.1,2 Despite their apparent longevity in vivo, CLL cells undergo spontaneous apoptosis in vitro, once removed from their in vivo microenvironment and placed into suspension culture without supportive stromal cells.3,4 Spontaneous apoptosis can be prevented by coculture with various stromal cells, such as marrow stromal cells (MSCs), follicular dendritic cells, or nurse-like cells.5-8 This prosurvival effect of stromal cells is largely dependent on direct cell contact between CLL and stromal cells.4,5,9 Chemokine secretion by stromal cells and expression of corresponding chemokine receptors on leukemia cells play a critical role in directional migration (chemotaxis) and adhesion of leukemia cells to MSCs, both in vitro10 and in vivo.11

CXCL12, previously called stromal cell–derived factor–1, is a chemokine constitutively secreted by MSCs that attracts and confines CLL cells to stromal cells via its cognate receptor CXCR4 expressed at high levels on CLL cells.10,12 This mechanism is shared with normal hematopoietic stem cells that require this receptor for homing to stromal niches in the marrow.13,14 Besides its activity on adhesion and migration of CLL cells,10 which is partially dependent on PI3K activation,15 CXCL12 also has a direct prosurvival effect on CLL cells.8,10 Once they engage in adhesion to stromal cells, CLL cells become resistant to the cytotoxic effects of drugs commonly used to treat CLL patients, such as fludarabine17 or corticosteroids.4 This primary drug resistance mechanism, also called cell adhesion–mediated drug resistance,18 may account for minimal residual disease in tissue compartments such as the marrow and relapses commonly seen in treatment of CLL patients.19-21

We previously demonstrated that CXCR4 antagonists can partially resensitize CLL cells to cytotoxic drugs in cocultures with MSCs,17 a finding that is currently pursued in clinical trials in leukemia patients,22 using the small molecule CXCR4 antagonist AMD3100 (now called Plerixafor). However, from our previous work and other studies,23,24 it is also apparent that targeting of CXCR4 not only partially overcomes stromal cell–mediated drug resistance; therefore, other CLL-microenvironment interactions may represent alternative therapeutic targets.

Phosphoinositide 3-kinases (PI3Ks) are among the most commonly activated signaling pathways in human cancers.25-27 In freshly isolated CLL cells, PI3Ks are constitutive activated,28 and CLL patients with unmutated immunoglobulin variable heavy chain genes, which generally display a more aggressive clinical
course than variable heavy chain-mutated patients, show overexpression of PI3K by real-time quantitative polymerase chain reaction.\textsuperscript{29} Furthermore, growth and survival signals from the microenvironment, such as adhesion to MSCs,\textsuperscript{9} CXCR4 activation,\textsuperscript{15} and B-cell receptor (BCR) activation,\textsuperscript{30} cause PI3K activation in CLL cells. Therefore, we investigated the activity of isoform-selective PI3K inhibitors using a panel of novel isoform-selective PI3K inhibitors that target different isoforms of the p110 subunit.

Therapeutic targeting of PI3K has been decelerated until recently because of the lack of specific inhibitors that possess sufficient activity, specificity, and bioavailability. The prototype PI3K inhibitors wortmannin and LY294002 are pan-specific PI3K inhibitors that sensitize human cancer cells to chemotherapy and radiation in vitro and in vivo\textsuperscript{31} but lack substrate specificity and show toxicity in animal studies,\textsuperscript{32} precluding their clinical development. However, over the past few years, we have witnessed a rapid expansion of information about new small molecules that target the PI3K family.\textsuperscript{33-35}

In response to cell stimulation by various growth factors and chemokines, PI3Ks phosphorylate phosphatidylinositol lipids at the D-3 position of the inositol ring, catalyzing the production of phosphatidylinositol-3,4,5-triphosphate (PIP\textsubscript{3}), which then sets in motion a coordinated set of events leading to cell growth, migration, and survival.\textsuperscript{25,36} The PI3Ks have been classified into 3 groups according to their structure and substrate specificity. Class IA isoforms couple to tyrosine kinases and consist of a p110 catalytic subunit (p110\textsubscript{α}, p110\textsubscript{β}, or p110\textsubscript{γ}), which is bound to one of 5 distinct p85 regulatory subunits, linking PI3K activity to the receptor tyrosine kinases (class Ia) or G protein–coupled receptors (class Ib).\textsuperscript{37} These heterodimers can be recruited either directly to cell-surface receptors, for instance, growth factor receptors, or indirectly by adaptor molecules, such as Shc, Grb2, or IRS-1.\textsuperscript{38} The p110\textsubscript{α} and p110\textsubscript{β} isoforms are ubiquitous, whereas p110\textsubscript{γ} is predominantly expressed in leukocytes.\textsuperscript{39,38} P110\textsubscript{β} was originally identified in leukocytes but is also expressed in other cell types, including breast tissue and melanocytes.\textsuperscript{40}

The serine-threonine protein kinase Akt (also protein kinase B) is activated as a result of PI3K activity and thus mediates many of the downstream effects of PI3K and consequently plays an important role in both normal and pathologic signaling through the PI3K pathway. Activation of Akt involves membrane binding as well as phosphorylation. Akt requires PIP\textsubscript{3} to be translocated to the cell membrane.\textsuperscript{41} After PIP\textsubscript{3} binding and positioning in the cell membrane, full activation of Akt requires phosphorylation on Ser\textsuperscript{473} in the hydrophobic motif by the mTOR complex 2 (mTORC2) and on Thr\textsuperscript{308} in the activation loop by the phosphoinositide-dependent kinase 1.\textsuperscript{38,42} Further downstream, Akt phosphorylates a variety of substrates involved in the regulation of key cellular functions. Because of the importance of PI3Ks in transducing a variety of extracellular, microenvironment-derived migration, growth, and survival signals, we tested new, isoform-selective PI3K inhibitors for activity on CLL cell migration to CXCL12, stromal cell interactions, and stromal cell-mediated drug resistance.

### Methods

#### Reagents and Antibodies

All reagents were purchased from Sigma-Aldrich (St Louis, MO) unless otherwise stated. Trypsin/ethylenediaminetetraacetic acid solution was obtained from Invitrogen (Carlsbad, CA), synthetic human CXCL12 (stromal cell–derived factor–1α) from Upstate Biotechnology (Chattanooga, VA), and Ly294002 (phosphatidylinositol 3-kinase inhibitor) from Calbiochem (San Diego, CA). The isoform-selective PI3K inhibitors PIK-90, PI-103, TGX115, ZK75, and IC87114 were synthesized after published patent specifications as described,\textsuperscript{43} prepared as 20-mM stocks in dimethyl sulfoxide and stored in aliquots at −20°C before use. The structures and key characteristics of these inhibitors have previously been detailed.\textsuperscript{33,35} The isoform-selective PI3K inhibitors were used at 1- and 10-μM concentrations for the various assays because we previously established that these inhibitors display activity in cellular systems at concentrations between 0.1 and 10 μM.\textsuperscript{43,44} Moreover, pharmacokinetic analysis revealed that concentrations of greater than 1-μM concentrations of these drugs (PI103) can be achieved in vivo.\textsuperscript{45}

If not otherwise stated, all primary antibodies were bought from Cell Signaling Technology (Danvers, MA). Secondary anti–mouse IgM (μ)-peroxidase antibody was purchased from Roche Diagnostics (Mannheim, Germany) and secondary enhanced chemiluminescence peroxidase-labeled anti–rabbit antibody from Jackson Immunoresearch Laboratories (West Grove, PA). The dephosphorylated fludarabine nucleoside arabinosyl-2-fluoroadenine (F-ara-A) was purchased from Sigma-Aldrich.

#### Cells and cell-culture conditions

After informed consent, acquired in accordance with the Declaration of Helsinki and Institutional Review Board approval, peripheral blood samples were obtained either from patients diagnosed with CLL according to clinical and immunophenotypic criteria at the Leukemia Department, University of Texas M. D. Anderson Cancer Center, or from healthy volunteers. Patients’ characteristics are summarized in Table S1 (available on the Blood website; see the Supplemental Materials link at the top of the online article). Peripheral blood mononuclear cells (PBMCs) were isolated via density gradient centrifugation over Ficoll Paque (GE Healthcare, Little Chalfont, United Kingdom). Cells were used fresh or viably frozen in fetal calf serum (FCS) plus 10% dimethyl sulfoxide (DMSO) and stored in aliquots at −80°C. CLL B cells were resuspended at 3 to 6 × 10\textsuperscript{6} cells/mL in RPMI 1640 (HyClone Laboratories, Logan, UT) with 10% FCS (heat-inactivated) and 1% penicillin-streptomycin (Invitrogen) at 37°C in a humidified atmosphere containing 5% carbon dioxide. The murine marrow stromal cell line M2-10B4 was purchased from ATCC (Manassas, VA).

#### Chemotaxis assay

The migration of PI3K inhibitor–pretreated CLL B cells in response to CXCL12 was evaluated using 24-well Corning chemotaxis chambers (Corning Life Sciences, Acton, MA) as described.\textsuperscript{10} In brief, transfected chambers with polycarbonate inserts with a 5-μm pore size were used, and CLL B cells (5 × 10\textsuperscript{6} in 100 μL) were added to the top chamber, whereas CXCL12 was added to the lower wells. The filter inserts were then placed into the wells and incubated at 37°C. Assay medium (RPMI with 0.5% bovine serum albumin [BSA]) without CXCL12 was used as baseline control. After 3 hours, cells that transmigrated into the lower chamber were collected and counted by flow cytometry (FACScan) at high flow for 20 seconds. A 1:20 dilution of input cells was counted under the same conditions. Results are shown as migration index (mean ± SEM), which represents the ratio between cells that migrated to the lower chamber in the presence of CXCL12 and cells that migrated in response to medium alone.

Pretreated CLL cells were incubated with the PI3K inhibitors or with vehicle (untreated controls) for 2 hours at 37°C.

#### Pseudoeperipolesis assay (in vitro migration of CLL B cells beneath stromal cells)

The migration of CLL cells beneath MSCs was observed and quantified as previously described.\textsuperscript{10} Briefly, the day before performing the assay, M2-10B4 stromal cells in RPMI supplemented with 10% FCS and antibiotics were seeded in collagen-coated 24-well plates at a concentration of 1.5 × 10\textsuperscript{5} cells per well. After overnight culture, the confluence of the stromal cell layer was controlled by phase-contrast microscopy, and then untreated and pretreated CLL B cells were added into each well to a final concentration of 5 × 10\textsuperscript{4} CLL cells per well. Pretreatment of the CLL cells...
was performed as described in “Reagents and antibodies.” The plates were incubated at 37°C for 5 hours, and after that nonmigrated cells were removed by vigorously washing of each well for 5 times with RPMI medium. Before detaching the migrated cells containing stromal cell layer for 1 minute with prewarmed (37°C) trypsin/ethylenediaminetetraacetic acid solution, the complete removal of nonmigrated cells and the integrity of the stromal cell layer were checked via phase-contrast microscopy. The detached cells were immediately suspended in 200-μL RPMI with 10% FCS for counting by flow cytometry. To exclude stromal cells from the counts, a lymphocyte gate was set using the different relative size and granularity (forward scatter, side scatter), and cell counting then was performed at high flow for 20 seconds.

**Actin polymerization assay**

Actin polymerization was performed as described.10 Briefly, 1.5 × 10^6 CLL cells, pretreated with PI3K inhibitors as described in “Reagents and antibodies,” were suspended in RPMI medium with 0.5% BSA and incubated with 200 ng/mL CXCL12 at 37°C for various amounts of time. At the indicated time points, 400 μL of the suspension was added to 100-μL staining solution (4 × 10^-7 mol/L fluorescein isothiocyanate–labeled phalloidin, 0.5 μg/mL 1-α-l-lyso phosphatidylcholin, and 8% formaldehyde, all Sigma-Aldrich) in phosphate-buffered saline. The analyses of the fixed cells were performed by flow cytometry on a FACScalibur. All time points are plotted relative to the mean fluorescence intensity of the same sample before the chemokine was added.

**Measurement of cell viability**

To determine the viability of CLL B cells, 200-μL cells were removed from the wells of a 24-well plate at the indicated time points and incubated for 15 minutes in fluorescence-activated cell sorter buffer (RPMI + 0.5% BSA) containing 40 NM 3,3’-dihexylxocarbocyanine iodide (DiOC<sub>6</sub>) and 10 μg/mL propidium iodide (PI), as described.5 Within 30 minutes, the cells were then analyzed by flow cytometry on a FACScalibur. BD Biosciences, San Jose, CA). Viable cells show high DiOC<sub>6</sub> fluorescence, whereas apoptotic cells have low DiOC<sub>6</sub> and PI fluorescence; necrotic cells are characterized by low DiOC<sub>6</sub> and high PI fluorescence (Figure 7A). We also cultured normal PBMCs under the same conditions, with or without the various PI3K inhibitors, fludarabine, and with or without stromal cell support, and their viability was also determined by staining with DiOC<sub>6</sub> and PI.

**Western blot analysis of Akt, pAkt, and β-actin**

A total of 3 × 10^6 serum-starved CLL B cells were pretreated with the PI3K inhibitors for 2 hours and then stimulated with 200 ng/mL CXCL12 for 1 minute. After that, the cells were lysed in ice-cold lysis buffer (Upstate Biotechnology) containing COMPLETE protease inhibitor cocktail tablets and PhosSTOP tablets (both Roche Diagnostics), 1 mM phenylmethylsulfonyl fluoride, 1 mM sodium orthovanadate, and the lysates were kept on ice for 30 minutes after a centrifugation at 14 000g for 30 minutes. The protein concentration of the supernatant was determined using Protein Assay solution (Bio-Rad, Hercules, CA) according to the manufacturer’s protocol. A total of 30 μg total protein of each sample was denatured by boiling for 5 minutes in Laemmli sample buffer, separated on a 4% to 15% Tris-HCL gel (Bio-Rad), and finally transferred to nitrocellulose membranes (Millipore, Billerica, MA). The membranes were blocked with 5% nonfat dry milk in TBST (10 mM Tris-HCL, 150 mM NaCl, 0.5% Tween) and incubated with the indicated primary antibodies overnight at 4°C. After incubation with the horseradish peroxidase–conjugated secondary antibody, membranes were treated with a chemiluminescence reagent, enhanced chemiluminescence (GE Healthcare), and subsequently exposed to a radiographic film (Eastman Kodak, Rochester, NY).

**Reverse-phase protein arrays**

Reverse-phase protein arrays (RPPAs) were performed as described.46 In brief, CLL B cells were pretreated with PI3K inhibitors and then stimulated with CXCL12 for 5 and for 30 minutes. The cells then were lysed in ice-cold lysis buffer as described for Western blot analysis. Cell lysates were normalized to a concentration of 4 μg/mL using bicinechinonic acid and boiled with 1% sodium dodecyl sulfate, and the supernatants were manually diluted in 5 2-fold serial dilutions with lysis buffer. An Aushon Biosystems arrayer (Aushon Biosystems, Billerica, MA) created 6336 spot arrays on nitrocellulose-coated FAST slides (Whatman Schleicher and Schuell, Dassel, Germany) using the serial dilutions. Each slide was probed with a validated primary antibody, and the signal was amplified using a Dako North America-catalyzed system (Dako North America, Carpenteria, CA). A secondary antibody (anti–mouse or anti–rabbit) was used as an amplification starting point. The slides were scanned, analyzed, and quantified using MicroGenie software (VirgeneTech, Carlisle, MA) and the Supercurve software package, as described previously. To generate heat maps, Treeview (University of Glasgow) and X-cluster software were used.

**Data analyses, statistics**

Unless otherwise stated, results are shown as mean plus or minus SEM of at least 3 experiments each. For statistical comparison between groups, the Student paired t test or Bonferroni t test was used. Analyses were performed using Prism 5 for Mac (GraphPad Software, San Diego, CA) and Primer of Biostatistics software (McGraw Hill, Columbus, OH). Data collected via flow cytometry were analyzed using FlowJo (TreeStar, Ashland, OR). To determine whether the combination of F-ara-A and PIK90 or the combination of F-ara-A and PI103 had an additive or more than additive cytotoxic effect in CLL suspension cultures and/or MSC cocultures, we performed Wilcoxon rank sum tests. Briefly, we computed the differences in mean relative viabilities between the combination treatment and the 2 individual treatments (eg, difference = viability for F-ara-A in combination with PIK90 at 1 μM − [viability for F-ara-A alone + viability for PIK90 at 1 μM]/2). The hypothesis was that, if there were an additive effect, the expected difference would be 0. If there were synergism or antagonism, the difference would be significantly different from 0. There was synergism if the difference were significantly smaller than 0.

**Results**

CXCL12-induced CLL B-cell chemotaxis, actin polymerization, and pseudoeoiperiplois keep PI3K activation

To determine the effects of new, isofom-selective PI3K inhibitors on CXCR4 responses in CLL cells, we examined the nonspecific PI3K inhibitor Ly294002 and the isofom-selective PI3K inhibitors PIK-90, PI-103, IC87114, and TGX115 in chemotaxis, pseudoeoiperiplois, and actin polymerization assays, as these processes are well known to be regulated by activation of CXCR4 by its ligand CXCL12 in CLL B cells.15 As shown in Figure 1, the rather
multitargeted p110α/p110γ inhibitors PI-103 and PIK-90 inhibited chemotaxis to levels that were 57.8% plus or minus 8.9% (PIK-90) or 51.4% plus or minus 1.7% (PI-103) of controls at 1 μM (mean ± SEM; n = 5) and 56.8% plus or minus 10.6% (PIK-90) or 53% plus or minus 2.9% (PI-103) of controls at 10 μM (mean ± SEM; n = 5, P < .05). IC87114, which is highly selective for p110δ, and TGX115, also highly selective for p110β/p110δ, did not affect chemotaxis. Ly294002 displayed similar activity as the p110α inhibitors, whereas pertussis toxin, which interferes with the Gi component of adenylate cyclase, almost completely abrogated CLL cell chemotaxis. Consistent with the chemotaxis data, we found that PIK-90 and PI-103 inhibited pseudopodiploisis to levels that were 74.2% plus or minus 12.9% (PIK-90) or 69.2% plus or minus 19.4% (PI-103) of controls (mean ± SEM; n = 4) at 1 μM and 57.9% plus or minus 13.7% (PIK-90) or 62.9% plus or minus 17% (PI-103) of controls at 10 μM (mean ± SEM; n = 4). These inhibitions were significant with P < .05, except for the 1-μM concentrations. The phase-contrast photomicrographs displayed in Figure 2 illustrate the effect of pretreatment of CLL cells with 10 μM PIK-90, demonstrating a marked reduction of CLL cell migration into the stromal cell layer compared with untreated controls (Figure 2A vs Figure 2B). Cell migration in response to chemokines, such as CXCL12, requires actin polymerization, and we therefore analyzed the effects of PI3K inhibitors on this cellular response. As shown in Figure 3, CXCL12-induced actin polymerization was decreased to levels that peaked at 170.7% plus or minus 5.1% at 15 seconds with 10 μM PIK-90, compared with 188.5% plus or minus 3.6% (t = 15 seconds) for 10 μM PI-103 and 195.1% plus or minus 4.1% (t = 15 seconds) for the untreated controls (mean ± SEM; n = 4 for each condition). This was a significant inhibition for PIK-90 (P = .01), but not for PI-103 (P = .27).

**Figure 2. Reduction of spontaneous migration of CLL cells beneath marrow stromal cells (pseudopodiploisis) by PIK-90 and PI-103.** CLL B cells were preincubated with PIK-90 or PI-103 and seeded onto confluent marrow stromal cell layers. After overnight incubation, the CLL cells that had not migrated into the stromal cell layer were vigorously washed off. Then, CLL B cells that migrated beneath the stromal cells were microphotographed (A) and quantified by flow cytometry (B). In phase-contrast, pseudopodiploisis is characterized by the dark appearance of CLL cells that have migrated into the same focal plane as the stromal cells. Cells were imaged in medium using a phase-contrast microscope (Model ELWD 0.3; Nikon, Garden City, NY) with a 0.25 NA objective lens. Images were captured with a Nikon D40 digital camera (Nikon, Tokyo, Japan) using Camera Control Pro software (Nikon Japan); when necessary, Adobe Photoshop 9.0 (Adobe Systems, San Jose, CA) was used for image processing. (A) Representative phase-contrast photomicrographs of untreated CLL cells, labeled “control,” and, in comparison, reduced pseudopodiploisis of the same CLL sample pretreated with 10 μM PIK-90 (bottom row). White bars in the left panels represent 100 μm (100× magnification); bars in the right panels, 50 μm (400× magnification); white filled arrow, a nonmigrated CLL cell; gray filled arrow, a migrated CLL cell; black filled arrow, a narrow stromal cell (framed). In panel B, each bar represents the mean (+SEM) relative number of migrated CLL cells from four different patients after pretreatment with the agents displayed on the horizontal axis, compared with the untreated controls. *Significant inhibition with P values less than .05.

**Figure 3. PI3K inhibitors PIK-90 and PI-103 reduce actin polymerization in response to CXCL12 stimulation.** Intracellular F-actin was measured in CLL cells using fluorescein isothiocyanate–labeled phalloidin after adding 200 ng/mL CXCL12 at time 0. Results at the time points indicated on the horizontal axis are shown as percentage of intracellular F-actin relative to the value of F-actin before addition of CXCL12. Each data point represents the mean (±SEM) of four different patients. Pretreatment of CLL cells with Ly294002 and PIK-90 reduces intracellular F-actin to approximately 70% of controls, whereas PI-103 had only a minor effect with an approximate 10% reduction.

**Figure 4. PI3K inhibitors induce apoptosis in CLL B cells.** To determine the effects of the isoform-selective PI3K inhibitors on CLL cell viability, we incubated CLL cells from different patients with various concentrations of the inhibitors for 24, 48, and 72 hours. As shown in Figure 4, we found that the PI3K inhibitors PI-103 and PIK-90 revealed the strongest apoptosis-inducing effects at both concentrations and at all different time points. Using a concentration of 10 μM, PIK-90 reduced the viability of CLL cells to 51.1% plus or minus 6.6% at 24 hours, whereas 1 μM PIK-90 reduced the viability to 77.8% plus or minus 6.4%; 10 μM PI-103 and PIK-90 induce apoptosis in CLL B cells

**Figure 4. PI3K inhibitors induce apoptosis in CLL B cells.** CLL cell viability was determined 24, 48, and 72 hours after addition of the PI3K inhibitors that are displayed on the vertical axis by staining with DiOC6 and PI. The results represent the mean (±SEM; n = 7) relative viability of PI3K inhibitor-treated CLL B cells compared with untreated controls (100%). The strongest cytotoxic effects at 1 μM and 10 μM were obtained with PIK-90 and PI-103, whereas TGX115 and IC87114 were found to be less potent. *Significant decrease in viability (P < .05).
Western blotting. CXCL12 stimulation resulted in the phosphorylation of Akt, especially at Ser473, whereas Thr308 revealed a significantly weaker phosphorylation (data not shown). A total of 1 and 10 μM PIK-90 reduced the phospho-Akt(Ser473) to 63.8% and 65%, and PI-103 reduced the phospho-Akt(Ser473) to 56% or 78.5% of controls, as shown in Figure 5.

PIK-90 and PI-103 inhibit signaling through AKT and S6 ribosomal proteins and up-regulate apoptosis-related proteins

We next used a sensitive high-throughput RPPA approach to assay PI3K inhibitor-induced changes in the phosphorylation status of proteins that are involved in PI3K-AKT signaling and to analyze the expression of apoptosis-related proteins. The advantage of RPPA is that it allows a more comprehensive interrogation of signaling events than does Western blotting. As shown in Figure 6A, pretreatment of CLL cells with PIK-90 and PI-103 resulted in decreased levels of phosphorylated Akt (especially at Ser473), of phosphorylated serine/threonine protein kinase glycogen synthase kinase 3 (at Ser21/9-PIK90 only), and of phosphorylated S6 ribosomal protein (at Ser235/236 and Ser240/244). A total of 1 μM PIK-90 significantly reduced phosphorylation of Akt at Ser473 to 80% plus or minus 2% and phosphorylation of S6 at Ser235/236 to 77% plus or minus 8% of the levels in untreated controls after stimulation with CXCL12. PIK-90 at 1 μM also produced a trend to decreased phosphorylation of glycogen synthase kinase 3 at Ser21/9 (to 85.2% ± 10% of the level in untreated controls) and to increased cleavage of caspase 7 (to 114% ± 12% of the level in untreated controls).

PIK-90 and PI-103 inhibit Akt(Ser473) phosphorylation

CXCL-12 induced activation of the PI3-K effector Akt, and the effects of the isoform-specific PI3K inhibitors were assayed by Figure 5. Activation of Akt after stimulation with CXCL12 is inhibited by PIK-90 and PI-103. After preincubation with the PI3K inhibitors that are indicated above the bands, CLL B cells were stimulated with CXCL12 for 1 minute and then lysed. The phosphorylated forms of Akt (p473; p308 not shown) were visualized by Western blot, and the band intensities were analyzed using the Software Labimage 1D. The relative band intensities compared with the band intensity of the control sample (+ CXCL12, no PI3K inhibitor, 100%) are indicated below each band. PIK-90 and PI-103 reduced the p473 phosphorylation of Akt by approximately 50% compared with the control. In contrast, the inhibitory effects of TGX115 and Ly294002 on CXCL12-induced Akt phosphorylation were minor. The figure shows data from 1 representative patient sample of 3 patient samples with similar results.

PI-103 reduced the viability of CLL cells to 77.5% plus or minus 3.4% and to 80.5% plus or minus 2.8% at 24 hours with 1 μM PI-103 (mean ± SEM; n = 7 for each condition). The asterisks in Figure 4 indicate which inhibitor and at what concentrations induced significant reduction in CLL viabilities with P less than .05. Using normal PBMCs from healthy donors, we found that the cytotoxicity of PI3K inhibitors was not restricted to CLL cells, but also evident in normal PBMCs. Generally, this cytotoxic effect was less pronounced compared with CLL cells (Table S3).

Figure 6A, pretreatment of CLL cells with PIK-90 and PI-103 resulted in decreased levels of phosphorylated Akt (especially at Ser473), of phosphorylated S6 ribosomal protein (at Ser235/236 and Ser240/244). A total of 1 μM PIK-90 at 1 μM PI-103 reduced phosphorylation of Akt at Ser473 to 80% plus or minus 2% and phosphorylation of S6 at Ser235/236 to 77% plus or minus 8% of the levels in untreated controls after stimulation with CXCL12. PIK-90 at 1 μM also produced a trend to decreased phosphorylation of glycogen synthase kinase 3 at Ser21/9 (to 85.2% ± 10% of the level in untreated controls) and to increased cleavage of caspase 7 (to 114% ± 12% of the level in untreated controls). A total of 1 μM PI-103 reduced phosphorylation of Akt at Ser473 to 72% plus or minus 3% and of S6 ribosomal protein at Ser236/236 to 83% plus or minus 8% along with a trend to increased cleavage of PARP to 114% plus or minus 16% of the level in untreated controls (mean ± SEM; n = 5 for all data). The heat map in Figure 6B shows the individual results from 5 different PI3K-AKT signaling and up-regulation of apoptosis-related proteins by PIK-90 and PI-103. (A) PIK-90 and PI-103 reduce CXCL12-induced phosphorylation of Akt (especially at Ser473) and of its downstream effector S6 ribosomal protein (at Ser235/236 and Ser240/244) and produce a trend to increased cleavage of the apoptosis-related proteins caspase 7 and its target poly-(ADP-ribose) polymerase (PARP). To generate the bar graphs, (phospho)protein expression of B CLL cells from 5 different patients was quantified with RPPA, and the quantification data were then corrected for loading and normalized to controls that were stimulated with CXCL12 but not preincubated with PI3K inhibitors. Each bar represents the mean (± SEM; n = 5) relative quantity of the (phospho)proteins that are displayed on the right vertical axis after stimulation of Akt, especially at Ser473, whereas Thr308 revealed a significantly weaker phosphorylation (data not shown). A total of 1 and 10 μM PIK-90 reduced the phospho-Akt(Ser473) to 63.8% and 65%, and PI-103 reduced the phospho-Akt(Ser473) to 56% or 78.5% of controls, as shown in Figure 5.

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We next used a sensitive high-throughput RPPA approach to assay PI3K inhibitor-induced changes in the phosphorylation status of proteins that are involved in PI3K-AKT signaling and to analyze the expression of apoptosis-related proteins. The advantage of RPPA is that it allows a more comprehensive interrogation of signaling events than does Western blotting. As shown in Figure 6A, pretreatment of CLL cells with PIK-90 and PI-103 resulted in decreased levels of phosphorylated Akt (especially at Ser473), of phosphorylated serine/threonine protein kinase glycogen synthase kinase 3 (at Ser21/9-PIK90 only), and of phosphorylated S6 ribosomal protein (at Ser235/236 and Ser240/244). A total of 1 μM PIK-90 significantly reduced phosphorylation of Akt at Ser473 to 80% plus or minus 2% and phosphorylation of S6 at Ser235/236 to 77% plus or minus 8% of the levels in untreated controls after stimulation with CXCL12. PIK-90 at 1 μM also produced a trend to decreased phosphorylation of glycogen synthase kinase 3 at Ser21/9 (to 85.2% ± 10% of the level in untreated controls) and to increased cleavage of caspase 7 (to 114% ± 12% of the level in untreated controls). A total of 1 μM PI-103 reduced phosphorylation of Akt at Ser473 to 72% plus or minus 3% and of S6 ribosomal protein at Ser236/236 to 83% plus or minus 8% along with a trend to increased cleavage of PARP to 114% plus or minus 16% of the level in untreated controls (mean ± SEM; n = 5 for all data). The heat map in Figure 6B shows the individual results from 5 different PI3K-AKT signaling and up-regulation of apoptosis-related proteins by PIK-90 and PI-103. (A) PIK-90 and PI-103 reduce CXCL12-induced phosphorylation of Akt (especially at Ser473) and of its downstream effector S6 ribosomal protein (at Ser235/236 and Ser240/244) and produce a trend to increased cleavage of the apoptosis-related proteins caspase 7 and its target poly-(ADP-ribose) polymerase (PARP). To generate the bar graphs, (phospho)protein expression of B CLL cells from 5 different patients was quantified with RPPA, and the quantification data were then corrected for loading and normalized to controls that were stimulated with CXCL12 but not preincubated with PI3K inhibitors. Each bar represents the mean (± SEM; n = 5) relative quantity of the (phospho)proteins that are displayed on the right vertical axis after stimulation of Akt, especially at Ser473, whereas Thr308 revealed a significantly weaker phosphorylation (data not shown). A total of 1 and 10 μM PIK-90 reduced the phospho-Akt(Ser473) to 63.8% and 65%, and PI-103 reduced the phospho-Akt(Ser473) to 56% or 78.5% of controls, as shown in Figure 5.

PIK-90 and PI-103 inhibit signaling through AKT and S6 ribosomal proteins and up-regulate apoptosis-related proteins

We next used a sensitive high-throughput RPPA approach to assay PI3K inhibitor-induced changes in the phosphorylation status of proteins that are involved in PI3K-AKT signaling and to analyze the expression of apoptosis-related proteins. The advantage of RPPA is that it allows a more comprehensive interrogation of signaling events than does Western blotting. As shown in Figure 6A, pretreatment of CLL cells with PIK-90 and PI-103 resulted in decreased levels of phosphorylated Akt (especially at Ser473), of phosphorylated serine/threonine protein kinase glycogen synthase kinase 3 (at Ser21/9- PIK90 only), and of phosphorylated S6 ribosomal protein (at Ser235/236 and Ser240/244). A total of 1 μM PIK-90 significantly reduced phosphorylation of Akt at Ser473 to 80% plus or minus 2% and phosphorylation of S6 at Ser235/236 to 77% plus or minus 8% of the levels in untreated controls after stimulation with CXCL12. PIK-90 at 1 μM also produced a trend to decreased phosphorylation of glycogen synthase kinase 3 at Ser21/9 (to 85.2% ± 10% of the level in untreated controls) and to increased cleavage of caspase 7 (to 114% ± 12% of the level in untreated controls). A total of 1 μM PI-103 reduced phosphorylation of Akt at Ser473 to 72% plus or minus 3% and of S6 ribosomal protein at Ser236/236 to 83% plus or minus 8% along with a trend to increased cleavage of PARP to 114% plus or minus 16% of the level in untreated controls (mean ± SEM; n = 5 for all data). The heat map in Figure 6B shows the individual results from 5 different
patients in terms of phosphorylation of the aforementioned PI3K-AKT signaling proteins and of cleavage of the apoptosis-related proteins caspase 7 and PARP after treatment with 2 different concentrations of the PI3K inhibitors PIK-90 and PI-103. The heat map demonstrates a general increase in cleavage of caspase 7 and of PARP after treatment with the PI3K inhibitors shown along with a general decrease in phosphorylation of Akt (at Thr308 and Ser473) and of S6 (at Ser235/236 and Ser240/244) in comparison with untreated controls. S6 phosphorylation was most consistently down-regulated in the 5 patients after treatment with PIK90.

**PIK-90 and PI-103 display a more than additive cytotoxic effect in combination with fludarabine and reverse the protective effect of stromal cells**

Inhibitors of the PI3K-AKT pathway combined with conventional cytotoxic drugs may provide a more effective strategy to enhance treatment responses compared with either agent by itself. Because PIK-90 and PI-103 were the most potent PI3K inhibitors in our migration and stromal cell–adhesion assays and induced significant apoptosis as single agents in suspension cultures (Figure 4), we subsequently tested these PI3K inhibitors alone and in combination with fludarabine in CLL-MSC cocultures. Figure 7A shows a representative analysis of CLL cells cultured with M2-10B4 cells and treated with 10 μM F-ara-A and different concentrations of PIK-90 for 24 hours. Although in this case the addition of 10 μM F-ara-A alone did not lead to a significant decrease in viability (87.8% for controls, and 88% after treatment with 10 μM F-ara-A), 1 μM PIK-90 caused a decreased in viability to approximately 75.6%, and the addition of both compounds together decreased the viability of the B CLL cells to only 23%. A total of 10 μM PIK-90 alone also reduced the viability to 29.4%, whereas the combination of both compounds resulted in an additional decrease in viability to 20.7%. Figure 7B compares the viability of CLL B cells from 10 different patients cultured with or without M2-10B4 cells and treated with 10 μM F-ara-A or PI3K inhibitors alone, or a combination of both agents at different time points (24, 48, 72 hours). Pretreatment of CLL cells with the PI3-K inhibitors Ly 294002, PI-103, and PIK-90 enhanced the cytotoxicity of fludarabine (compare right column to left column) and also partially reversed the protective effect of stromal cells on fludarabine-induced apoptosis (compare top row with bottom row). For example, the relative viability of CLL cells cultured on MSC treated with 10 μM F-ara-A was 93.8% plus or minus 2.4% at 24 hours, 66.3% plus or minus 11% at 48 hours, and 68.9% plus or minus 7.6% at 72 compared with untreated controls. For comparison, treatment with 1 μM PIK-90 decreased the viability to 68.3% plus or minus 3.7% at 24 hours, 70.6% plus or minus 5.9% at 48 hours, and 79.8% plus or minus 6% at 72 hours. However, the combination of both 10 μM F-ara-A and 1 μM PIK-90 decreased the viability to 57.7% plus or minus 5.4% at 24 hours, 21% plus or minus 6.3% at 48 hours, and 26.5% plus or minus 5.3% at 72 hours (mean ± SEM; n = 10 for all data points; Figure 7B bottom right diagram). We used the Wilcoxon rank sum test to determine that there was an additive or more than additive effect for the combination treatment of PI3K inhibitors and F-ara-A. We found that both PIK90 and PI103 displayed a more than additive cytotoxic effect when combined with F-ara-A, both in suspension and CLL cell cocultures with MSCs. Moreover, this synergistic
effect was noticed for both concentrations of the PI3K inhibitors (10 and 1 μM). The data analysis for this experiment is shown in Tables S2A and S2B, which list the difference in viability for each of the combination treatments at 1 μM and at 10 μM for the PI3K inhibitors, as well as the result of Wilcoxon rank sum test, for the groups with and without M2, respectively.

### Table 1. Specificity profiles of the PI3K inhibitors used in this study, based on IC50 values (μM)

<table>
<thead>
<tr>
<th>Name</th>
<th>p110α</th>
<th>p110β</th>
<th>p110γ</th>
<th>p110δ</th>
<th>p110ε</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ly294002</td>
<td>9.3</td>
<td>2.9</td>
<td>38</td>
<td>6.0</td>
<td></td>
<td>Pan-PI3K inhibitor</td>
</tr>
<tr>
<td>PIK-90</td>
<td>0.011</td>
<td>0.35</td>
<td>0.018</td>
<td>0.058</td>
<td></td>
<td>Low mTOR activity, cell permeable</td>
</tr>
<tr>
<td>PI-103</td>
<td>0.008</td>
<td>0.088</td>
<td>0.15</td>
<td>0.048</td>
<td></td>
<td>Targets PI3K(22p, mTORC2, DNA-PK, p235)</td>
</tr>
<tr>
<td>TGX115</td>
<td>61</td>
<td>0.13</td>
<td>100</td>
<td>0.63</td>
<td></td>
<td>p110β- and p1-selective</td>
</tr>
<tr>
<td>ICB7114</td>
<td>200</td>
<td>16</td>
<td>61</td>
<td>0.13</td>
<td></td>
<td>Selective p110β inhibitor, cell permeable</td>
</tr>
</tbody>
</table>

The IC50 values and the key characteristics of the inhibitors are based on the data published by Knight et al33 and Marone et al.56

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### Discussion

Interactions between CLL cells and the microenvironment are critical for maintenance and expansion of the neoplastic B-cell clone.6,19,48 These interactions occur in tissue compartments, such as the marrow and secondary lymphatic tissues, where CLL cells interact with a variety of nonmalignant accessory cells collectively referred to as stromal cells. These interactions provide growth and survival signals and also confer drug resistance and therefore may account for residual disease and relapses after conventional therapies.10,19,44 Therefore, increasing emphasis is being placed on therapeutically targeting the leukemia microenvironment. Targeting stroma-derived signaling pathways (such as the PI3K pathways) that control CLL cell survival and drug resistance is an attractive, new approach. Our data show that PI3K inhibitors can effectively antagonize migratory, survival, and drug-resistance signals derived from CXCL12 and CLL-MSC interactions. More specifically, we found that the class I PI3K inhibitors PIK-90 and PI-103 effectively inhibited CLL cell migration and signaling responses to CXCL12. Furthermore, PIK-90 and PI-103 inhibited CLL cell migration beneath stromal cells (Figure 2), S6, and Akt phosphorylation on Ser473 (Figures 5, 6), and induced CLL cell apoptosis in suspension (Figure 4) and MSC cocultures (Figure 7B). In addition, PIK-90 and PI-103 displayed a more than additive effect when used in combination with F-ara-A for induction of CLL cell apoptosis, and PIK-90 and PI-103 inhibited CLL cell migration and signaling responses to CXCL12. However, chemical disruption of CXCL12 sensing and cell polarization. However, chemical disruption of PI3K signaling by p110α inhibitors did not completely abolish chemotaxis (Figure 1), suggesting that PIP2-independent pathways are also involved in the regulation of chemotaxis, as suggested earlier.53,54

In cancer cells, constitutive PI3K activation is a common feature that can be caused through different mechanism. In several solid tumors, such as breast, colon, or endometrial cancers, PIK3CA, the gene encoding the PI3K catalytic subunit (p110α), is mutated and encodes mutant p110α proteins that constitutively activate its kinase activity and induce malignant growth in normal cells.57 Other mechanisms of PI3K activation in malignant cells are related to PTEN inactivation or to mutations in the p85 regulatory subunit.59 In contrast to solid tumors, where activating PI3K mutations are found in approximately 30% of the cases,58 PI3K mutations appear to be absent or extremely rare in cancers of the hematopoietic system, such as leukemias.57 No activating mutation in the PIK3CA or PI3KCD genes has been identified in acute myeloid leukemia and lymphoma patients.60,62
from activating signals from the microenvironment, in particular contact-dependent signals from MSCs. On the molecular level, stromal cells in the microenvironment can activate PI3Ks in CLL cells via the CXCL12-CXCR4 axis, as demonstrated in this study, or by activating BCR signaling. Collectively, these data suggest that constitutive PI3K activation in CLL and other leukemias is induced by external signals from the microenvironment, rather than being intrinsic to the leukemia cells, and stromal cell–derived signals, such as activation of the CXCL12-CXCR4 axis, play a pivotal role in the stroma-CLL cross talk leading to PI3K activation.

One may argue that signaling pathways activated by stromal cells could be redundant; therefore, blocking a single pathway, such as the PI3Ks, could be bypassed by alternative signaling routes. However, PI3Ks are involved in CXCR4-, BCR-, and other stromal cell–related signaling pathways, and are found to be overexpressed in the unmutated subgroup of CLL patients. These findings suggest that PI3Ks integrate multiple relevant signaling pathways in CLL cells. Moreover, our finding that the PIK inhibitors PIK-90 and PI-103 induce CLL cell apoptosis at low micromolar concentrations, even in the presence of MSCs, suggests that these drugs can turn off a critical survival pathway for CLL cells in stromal cell contact cultures. This is again highly relevant in the clinical context, where new drugs for CLL patients are needed to overcome stromal cell–mediated drug resistance in the marrow microenvironment to eradicate minimal residual disease. Another important concern regarding the therapeutic implications of our findings is the question of therapeutic specificity and safety of PIK inhibitors. Clearly, given the importance of PI3Ks in various tissues and biologic functions, PI3K inhibitors and particularly their long-term application could cause unwanted toxicities. On the other hand, pan-PI3K inhibitors are currently developed for cancer therapy, such as PI-103, and show encouraging preclinical toxicity data. Our finding that the PI3K inhibitors PIK-90 and PI-103 also induce cytotoxicity in normal blood mononuclear cells (Table S3) indicates that hematotoxicity and/or lymphocyte toxicity needs to be characterized and closely monitored in these initial clinical trials.

Collectively, our data demonstrate that PI3Ks play an important role in regulating interactions between CLL cells and MSCs via the CXCR4 receptor. The isoform-selective PI3K inhibitors PIK-90 and PI-103 induced CLL cell apoptosis in MSC cocultures, synergized with fludarabine in induction of CLL cell apoptosis, and reversed stromal cell–mediated resistance to fludarabine, providing a rationale to evaluate this class of PI3K inhibitors in CLL patients.

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Authorship

Contribution: M.N. performed experiments and data analysis and wrote the paper; M.H. and A.V.K. performed experiments and data analysis; B.T.H. performed the RPP arrays and RPPA analysis and reviewed the manuscript; Z.A.K. and K.M.S. helped with the experimental design and data analysis related to PI3K inhibitors and reviewed the manuscript; J.H. analyzed data regarding synergism between PI3K inhibitors and F-ara-A; W.G.W. and M.J.K. provided patient samples, analyzed data, and reviewed the manuscript; and J.A.B. designed the research, supervised the study, analyzed the data, and wrote the paper.

Conflict-of-interest disclosure: The authors declare no competing financial interests.

Correspondence: Jan A. Burger, Department of Leukemia, Unit 428, University of Texas M. D. Anderson Cancer Center, PO Box 310402, Houston, TX 77230-1402; e-mail: jaburger@mdanderson.org.

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