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# Ablation of PI3K blocks BCR-ABL leukemogenesis in mice, and a dual PI3K/mTOR inhibitor prevents expansion of human BCR-ABL+ leukemia cells

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# Research Article

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# Ablation of PI3K blocks BCR-ABL leukemogenesis in mice, and a dual PI3K/mTOR inhibitor prevents expansion of human BCR-ABL+ leukemia cells

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Some cases of pre–B cell acute lymphoblastic leukemia (pre–B-ALL) are caused by the Philadelphia (Ph) chromosome–encoded BCR-ABL oncogene, and these tend to have a poor prognosis. Inhibitors of the PI3K/AKT pathway reduce BCR-ABL—mediated transformation in vitro; however, the specific PI3K isoforms involved are poorly defined. Using a murine model of Ph<sup>+</sup> pre–B-ALL, we found that deletion of both *Pik3r1* and *Pik3r2*, genes encoding class IA PI3K regulatory isoforms, severely impaired transformation. BCR-ABL—dependent pre/pro-B cell lines could be established at low frequency from progenitors that lacked these genes, but the cells were smaller, proliferated more slowly, and failed to cause leukemia in vivo. These cell lines displayed nearly undetectable PI3K signaling function and were resistant to the PI3K inhibitor wortmannin. However, they maintained activation of mammalian target of rapamycin (mTOR) and were more sensitive to rapamycin. Treatment with rapamycin caused feedback activation of AKT in WT cell lines but not PI3K-deficient lines. A dual inhibitor of PI3K and mTOR, PI-103, was more effective than rapamycin at suppressing proliferation of mouse pre–B-ALL and human CD19<sup>+</sup>CD34<sup>+</sup> Ph<sup>+</sup> ALL leukemia cells treated with the ABL kinase inhibitor imatinib. Our findings provide mechanistic insights into PI3K dependency in oncogenic networks and provide a rationale for targeting class IA PI3K, alone or together with mTOR, in the treatment of Ph<sup>+</sup> ALL.

# Introduction

The *BCR-ABL* oncogene causes chronic myelogenous leukemia (CML) and a fraction of pre–B cell acute lymphoblastic leukemia (pre–B-ALL). The genetic lesion encoding the BCR-ABL fusion protein is a translocation, t(9;22), termed the Philadelphia chromosome (Ph) (1). The ABL kinase inhibitor imatinib mesylate produces stable remission for many patients with CML but is less effective in Ph<sup>+</sup> ALL and myeloid or lymphoid blast crisis (BC) phases of CML (2). Clinical resistance to imatinib has prompted investigators to seek novel compounds specific to downstream pathways that are essential for BCR-ABL-mediated cell proliferation and survival (3).

One potential target is PI3K, a family of signaling enzymes of emerging interest for pharmaceutical development (4). PI3Ks are activated by BCR-ABL as well as extracellular signals to produce phosphatidylinositol-3,4,5-trisphosphate (PIP<sub>3</sub>). This lipid is a second messenger that recruits and activates downstream effector proteins such as the serine/threonine kinase AKT (5, 6). Inhibiting the PI3K/AKT pathway reduces BCR-ABL transformation in human and murine myeloid colony-forming assays and murine

Nonstandard abbreviations used: B-ALL, B cell acute lymphoblastic leukemia; BC, blast crisis; CML, chronic myelogenous leukemia; L-CFC, leukemic colony-forming cell; LyBC, lymphoid BC; mTOR, mammalian target of rapamycin; mTORC1, mTOR complex–1; Ph, Philadelphia chromosome; PIP<sub>3</sub>, phosphatidylinositol-3,4,5-trisphosphate; rS6, ribosomal protein S6; S6K, S6 kinase.

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B lymphoid colony-forming assays (7–10). However, the specific PI3K isoforms or subgroups required for BCR-ABL transformation and leukemogenesis are poorly defined.

Class IA PI3Ks, the predominant subgroup triggered by activated tyrosine kinases such as BCR-ABL (5, 6), are composed of a catalytic subunit (p110α, p110β, or p110δ) and a tightly associated regulatory subunit (p85 $\alpha$ , p55 $\alpha$ , p50 $\alpha$ , p85 $\beta$ , or p55 $\gamma$ ). The general functions of the regulatory subunits are to maintain the stability of catalytic subunits and direct the heterodimer to membrane-associated signaling complexes. Broad-spectrum inhibitors of PI3K catalytic subunits, such as LY294002 or wortmannin, block BCR-ABL transformation, but the effects of targeting specific catalytic isoforms have not been reported (7, 9). Among regulatory isoforms, p85α appears to be essential for CML cell survival (8, 11). However, we found that targeted deletion of the mouse gene *Pik3r1*, encoding p85 $\alpha$  as well as p55 $\alpha$ and p50α, caused only partial impairment of leukemic lymphoid colony formation (CFU-pre-B) and leukemogenesis (9). The transformed BM progenitor cells were sensitive to LY294002 and maintained comparable PI3K signaling output. These findings imply that other regulatory isoforms, namely p85  $\beta$  or p55  $\gamma$  , can compensate in this system (9).

To define more clearly the role of class IA PI3K and compensatory pathways in BCR-ABL transformation of B cell progenitors, we used mice with combined deletion of *Pik3r1* and *Pik3r2* (encoding p85β) in the B lineage. Combined deletion of *Pik3r1* and *Pik3r2* caused reduced catalytic subunit expression and abro-



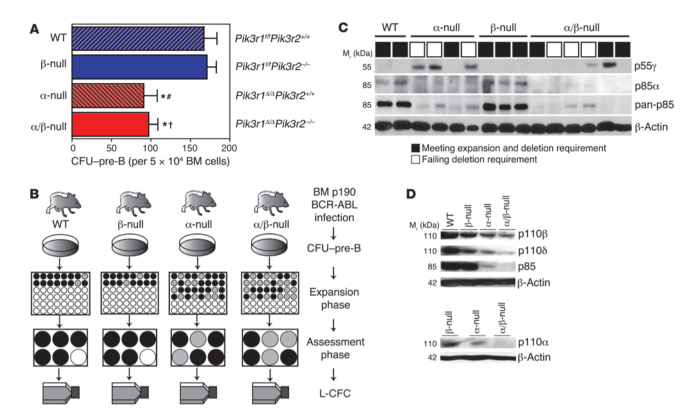


Figure 1

Decreased BCR-ABL-mediated (p190) colony transformation of both  $\alpha$ -null and  $\alpha/\beta$ -null progenitor B cells. (**A**) Plating of p190-infected whole BM (5 × 10<sup>4</sup> cells) from 3- to 6-week-old mice of the indicated genotypes, on M3630 (CFU-pre-B) medium. \*P < 0.05 versus WT; \*P < 0.01 versus  $\beta$ -null; †P < 0.001 versus  $\beta$ -null; 1-way ANOVA, n = 3–7 independent experiments; mean values  $\pm$  SEM are shown. (**B**) Schematic flowchart depicting the clonogenic expansion and assessment of PI3K expression status in L-CFCs. CFU-pre-B colonies were scored at day 7 after infection, and single colonies were selected and transferred to liquid culture. Outgrowth of p190+ L-CFCs was then quantified ("expansion" was defined as reaching  $\ge$ 1 × 10<sup>6</sup> cells), followed by immunoblot assessment of Pik3r1 deletion and p55γ upregulation. Table 1 displays percentages of colonies meeting requirements for expansion, deletion of Pik3r1, and absence of p55γ upregulation. (**C**) Monitoring class IA PI3K subunit expression in expanded L-CFCs. Representative immunoblot shows multiple clones of each genotype with variable loss of p85α and p55γ upregulation. p85α-specific antibody was used to distinguish p85α from p85β. White boxes represent clones that failed to delete p85α or upregulated p55γ. Black boxes represent clones that were used for in vitro and/or in vivo assays. (**D**) Immunoblot for class IA PI3K catalytic subunits in established L-CFCs. Representative blot for 3 independent clones.

gated AKT activation. BCR-ABL transformation of PI3K/AKT-deficient cells was severely impaired, and the few surviving clones showed reduced fitness in vitro and increased sensitivity to imatinib and failed to cause leukemia in vivo. We also identify mammalian target of rapamycin (mTOR) signaling as a vital pathway that remains in cells that lack PI3K activity. A dual PI3K/mTOR inhibitor, PI-103, prevented "rebound" AKT activation caused by rapamycin and was more effective than rapamycin in synergizing with imatinib to suppress survival. PI-103 also augmented the antileukemic effect of imatinib in vivo and reduced the clonogenicity of primary human Ph+ ALLCD19+CD34+ and CML/ALLBCCD19+CD34+ progenitors. These data suggest that combined targeting of PI3K, mTOR, and BCR-ABL would provide an attractive therapeutic strategy in Ph+ leukemias.

### Results

Decreased leukemic colony formation. We bred  $Pik3r1^{f/P}lik3r2^{-/-}$  (12, 13) with CD19-Cre mice to create mice that have Pik3r1 (p85 $\alpha$ /p55 $\alpha$ /p50 $\alpha$ ) deleted specifically in the B cell lineage and Pik3r2 (p85 $\beta$ ) deleted in all cells. We harvested BM from  $Pik3r1^{f/P}lik3r2^{+/+}$ ,

 $Pik3r1^{1/5}Pik3r2^{-/-}$ ,  $Pik3r1^{\Delta/\Delta}Pik3r2^{+/+}$ , and  $Pik3r1^{\Delta/\Delta}Pik3r2^{-/-}$  mice and assessed transformation efficiency following infection with a retrovirus expressing the p190 isoform of BCR-ABL (p190), together with either GFP or human CD4 lacking the cytoplasmic tail (hCD4). For simplicity, we refer to the B lineage progenitors derived from these mice as WT, β-null, α-null, and α/β-null cells, respectively.

β-null progenitors were equivalent to WT cells in the number of CFUs (CFU–pre-B; Figure 1A). In both α-null and α/β-null progenitors, we observed a significant decrease in transformation efficiency (~50%) relative to that in controls (Figure 1A). We considered the possibility that deletion of Pik3r1 was incomplete and that cells that delete the floxed Pik3r1 allele have a competitive disadvantage. To test this, we selected single colonies (CFU–pre-B), monitored the expansion of these leukemic colony-forming cells (L-CFCs) in liquid culture, and assessed the deletion by immunoblotting for p85 expression (Figure 1, B and C, and Table 1). Only  $33\% \pm 3\%$  (n = 7 experiments; 69 of 215 total colonies selected) of the  $\alpha/\beta$ -null L-CFCs could be expanded versus  $86\% \pm 5\%$  of WT (n = 3; 44 of 52),  $87\% \pm 4\%$  (n = 7; 113 of 134) of  $\beta$ -null, and  $58\% \pm 6\%$  (n = 4; 51 of 90) of  $\alpha$ -null L-CFCs (Table 1). Of the



**Table 1** Decreased expansion of  $\alpha/\beta$ -null L-CFCs, with high frequency of incomplete *Pik3r1* deletion or p55g upregulation

CFU-pre-B transition to L-CFC	WT $(n=3)$	$\beta$ -null ( $n = 7$ )	$\alpha$ -null ( $n = 4$ )	$\alpha/\beta$ -null ( $n=7$ )
L-CFC outgrowth (reaching ≥1 × 10 <sup>6</sup> cells or ≥6 well plate) from CFU-pre-B selected (%)	86 ± 5	87 ± 4	$58 \pm 6^{A,B}$	$33 \pm 3^{B,C}$
L-CFCs deleted of <i>Pik3r1</i> (assessed by immunoblot) (%)	N/A	N/A	$60 \pm 7^{D}$	29 ± 5 <sup>E</sup>
L-CFCs lacking p55γ upregulation (assessed by immunoblot) (%)	100	98 ± 2	$63 \pm 4^{B,D}$	$22 \pm 2^{B,D}$
Establishable L-CFCs meeting expansion and deletion requirements (average %)	86	85	22	2

One-way ANOVA,  $^{A}P$  < 0.01 versus WT;  $^{B}P$  < 0.001 versus  $\beta$ -null;  $^{C}P$  < 0.05 versus WT;  $^{D}P$  < 0.001 versus WT;  $^{E}P$  < 0.01 versus  $\beta$ -null; mean values  $\pm$  SEM are shown. n refers to the number of experiments, for each of which we selected at least 50 colonies for transfer to liquid culture. The total number of colonies for each condition are given in the main text.

expanded L-CFCs from at least 3 experiments, only 29% ± 5% of the  $\alpha/\beta$ -null cells demonstrated deletion by immunoblot analysis compared with 60% ± 7% of the  $\alpha$ -null cells (Table 1). The regulatory isoform encoded by the *Pik3r3* gene, p55γ, was not detected in WT or  $\beta$ -null cells but was upregulated in some  $\alpha$ -null clones and most  $\alpha/\beta$ -null clones (Figure 1C and Table 1). The predominant outgrowth of clones lacking *Pik3r1* deletion and/or p55γ upregulation indicates a strong selective pressure to maintain class IA PI3K for colony survival, particularly when *Pik3r2* is absent.

The outgrowth of nondeleting cells from  $\alpha/\beta$ -null cultures was not due to selection of CD19- progenitors that lack Cre expression, since all clones of each genotype expressed CD19 on the surface (Supplemental Figure 1A; supplemental material available online with this article; doi:10.1172/JCI33337DS1). Most clones of each genotype expressed a mixture of markers of pro-B and pre-B cell progenitors (Supplemental Figure 1B), suggesting that the target cell for transformation is not limiting. Moreover, there was a comparable distribution of B cell progenitor subsets among BM cells 14 hours after infection, at which time the marker-expressing cells were similarly distributed, with mainly pro-B (B220+CD43+IgM-) and fewer pre-B (B220<sup>+</sup>CD43<sup>-</sup>IgM<sup>-</sup>) phenotypes in both β-null and  $\alpha/\beta$ -null samples (Supplemental Figure 1C). The inability of some  $\alpha$ -null and most  $\alpha/\beta$ -null L-CFCs to expand in culture is most likely due to ongoing deletion resulting in loss of proliferation or survival. Consistent with this model, ablating p85 $\alpha$  expression in *Pik3r1*<sup>f/f</sup>*Pik3r2*<sup>-/-</sup> cells by retroviral expression of Cre caused rapid and significant cell cycle arrest and apoptosis (Figure 2). Acute deletion of Pik3r1 in Pik3r2+/+ cells also induced cell cycle arrest and apoptosis; however, the effect on cell cycle was significantly weaker than observed in *Pik3r2*<sup>-/-</sup> cells. Together, the results indicate that B lineage transformation by p190 is almost completely dependent on class IA regulatory isoforms of PI3K, with Pik3r1 and Pik3r2 gene products exhibiting partially redundant functions.

Reduced class IA PI3K catalytic subunit expression and function. In established L-CFCs that showed complete Pik3r1 deletion without p55 $\gamma$  upregulation, we observed a reduction of p110 $\alpha$ , p110 $\beta$ , and p110 $\delta$  expression in cells lacking class IA regulatory isoforms, with greater loss of catalytic isoform expression in  $\alpha/\beta$ -null compared with  $\alpha$ -null or  $\beta$ -null L-CFCs (Figure 1D). Loss of the catalytic subunits is likely due to reduced stability in the absence of regulatory subunit partners (14).

To assess PI3K activity, we visualized PIP $_3$  in fixed cells using a specific antibody. Control L-CFCs exhibited PIP $_3$  immunostaining with a concentration in the plasma membrane and/or intracellular organelles (Figure 3A and Supplemental Figure 2). Pretreatment

with wortmannin blocked the staining in most cells, supporting the specificity of detection. Residual PIP<sub>3</sub> production was abolished by pretreatment with wortmannin for longer times (Supplemental Figure 2C). The  $\alpha/\beta$ -null L-CFCs displayed significantly reduced PIP<sub>3</sub> staining, resembling wortmannin-treated control cells. Thus, the few viable L-CFCs lacking Pik3r1 and Pik3r2 do not sustain PIP<sub>3</sub> production by residual class IA catalytic isoforms or the class IB enzyme.

Most  $\alpha/\beta$ -null clones showed no detectable AKT phosphorylation (p-AKT), and clones with residual p-AKT also showed upregulation of p55 $\gamma$  (Figure 3B). Acute *Pik3r1* deletion also abrogated AKT phosphorylation (Figure 2A). Cells that compensated for the loss of *Pik3r1* and *Pik3r2* by upregulating p55 $\gamma$  were eliminated from our further biochemical and leukemogenic investigations.

Lack of PI3K/AKT functional output. Transcription factors of the forkhead box subgroup O (FOXO) family are important AKT substrates in p190 B-ALL cells (9). In agreement with the lack of p-AKT,  $\alpha/\beta$ -null cells maintained a very low level of phosphorylated FOXO1 and FOXO3a (Figure 2A and Figure 3, B and C), which could not be reduced by pharmacological inhibition of PI3K (Figure 3C). This implies that a PI3K-independent kinase accounts for the residual FOXO phosphorylation. We investigated the sensitivity of  $\alpha/\beta$ -null cells to FOXO-mediated death via overexpression of FOXO3a or a mutant (FOXO3a.A3) that lacks the 3 consensus AKT phosphorylation sites (Figure 3D). The control cells that maintained normal levels of PI3K activity were resistant to WT FOXO3a while remaining sensitive to the PI3K-independent form FOXO3a.A3, as described previously (9). In contrast, the  $\alpha/\beta$ -null cells were equally sensitive to death caused by WT and AKT-resistant mutant FOXO3a. This indicates that in the presence of diminished PI3K output, the residual FOXO phosphorylation by an alternative kinase is unable to inactivate overexpressed FOXO proteins.

Proliferation and cell cycle deficits. We labeled cells with PKH26 to assess proliferation of established  $\alpha/\beta$ -null cells. We found that  $\alpha/\beta$ -null cells partitioned the PKH26 dye at a lower rate in comparison to control cells (Figure 4A). Cell cycle analysis supported these results, showing an increase in the percentage of cells in  $G_1$  and a dramatic reduction of cells in S phase.  $\alpha/\beta$ -null clones were also smaller, as assessed by forward light scatter (Figure 4A).

Reduced in vivo leukemogenic potential. To assess the in vivo consequences of diminished PI3K/AKT signaling, we examined the tumorigenic capacity of multiple L-CFC clones. In this assay, we used only  $\alpha/\beta$ -null L-CFCs lacking detectable p85 proteins or p55 $\gamma$ . WT (1 clone, n = 4 mice),  $\beta$ -null (5 clones, n = 16),  $\alpha$ -null



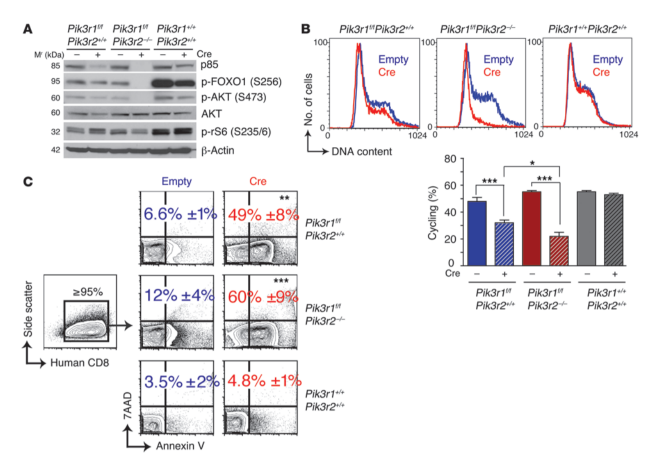


Figure 2 Dependence on continued PI3K signaling for maintenance of p190 L-CFCs. (A) L-CFCs were transduced with MSCV-IRES-human CD8 (Empty) or MSCV-Cre-IRES-human CD8 (Cre) and monitored for deletion of Pik3r1 and PI3K/mTOR signaling by immunoblotting 48 hours after infection. The blot is representative of 2 independent experiments using 3 separate clones. (B) L-CFCs were stained for human CD8 48 hours after infection and subsequently fixed and stained with propidium iodide for DNA content analysis of live gated human CD8+ cells by FACS. Graph shows the percentage of cells cycling (S/G<sub>2</sub>/M); mean  $\pm$  SEM; n = 3 clones. \*P < 0.05, \*\*\*P < 0.001; 1-way ANOVA. (C) Human CD8+ L-CFCs were assayed for apoptosis by annexin V/7AAD staining 48 hours after infection. Cell death is expressed as the percentage ( $\pm$ SEM) of annexin V+ cells to both 7AAD+ and 7AAD- cells; n = 3 clones. \*P < 0.01, \*\*\*P < 0.001, empty versus Cre-infected; 1-way ANOVA.

(1 clone, n = 4), and  $\alpha/\beta$ -null L-CFCs (5 clones, n = 23) were injected into sublethally irradiated NOD/SCID hosts. Mice with emerging EGFP<sup>+</sup> (or hCD4<sup>+</sup>) leukemic cells developed lymphadenopathy and splenomegaly, with lymphoblasts exhibiting pro/pre-B-ALL morphology and surface phenotype but lacking myeloid lineage markers (Figure 4C and Supplemental Table 1). Mice receiving WT or β-null cells died from pro/pre-B-ALL after an average of 15 and 13 days, respectively (Figure 4B), whereas mice receiving  $\alpha$ -null cells succumbed to disease after an average of 35 days (P = 0.052 vs. WT).  $\alpha/\beta$ -null cells had a significantly reduced capacity to initiate leukemia in vivo (Figure 4B). Among all  $\alpha/\beta$ -null transplanted mice, only 1 (of 23) developed a leukemic disease in which cells retained an  $\alpha/\beta$ -null status by immunoblot analysis. This mouse developed leukemic blasts in the BM, lymph nodes, and spleen with some mature B cell markers: CD19+, IgM+, B220+, CD43-, CD25-, CD24+, BP-1+, IgD-, CD11b-. These blasts displayed detectable p-AKT and prominent upregulation of p55y (Supplemental Figure 3). The striking block in leukemogenesis and the upregulation of p55y in the 1 instance of emerging leukemia from  $\alpha/\beta$ -null cells strongly suggest that p190 drives B-ALL in a manner completely dependent on class IA PI3K function. However, the residual proliferation in vitro prompted us to examine the drug sensitivity of control and PI3K-deficient L-CFCs.

The mTOR pathway is a vital signaling node for  $\alpha/\beta$ -null cells. To assess the importance of different signaling mechanisms in the absence of PI3K signaling output, we measured cell cycle progression after treatment with pathway-selective inhibitors. Wortmannin, used at a concentration (50 nM) that inhibits all classes of PI3K but not mTOR (15, 16), caused a modest increase in the G<sub>1</sub> fraction and a significant decrease in cycling ability (percentage of cells in S/G<sub>2</sub>/M phase) of control cells (β-null) (Figure 5A, right panel, P < 0.001). However, wortmannin had no significant effect on the cycling ability of  $\alpha/\beta\text{-null}$  cells, indicating that other PI3K subclasses do not compensate appreciably for class IA PI3K in cell cycle control. LY294002, a global PI3K inhibitor that also inhibits mTOR directly (15, 17), induced  $G_1$  arrest in both control and  $\alpha/\beta$ -null cells; moreover, the  $\alpha/\beta$ -null cells were hypersensitive, as shown by the greater decrease in the number of cycling  $\alpha/\beta$ -null compared with control cells (P < 0.05). Consistent with the interpretation that the effects of LY294002 were mainly due to direct effects on the rapamycin-



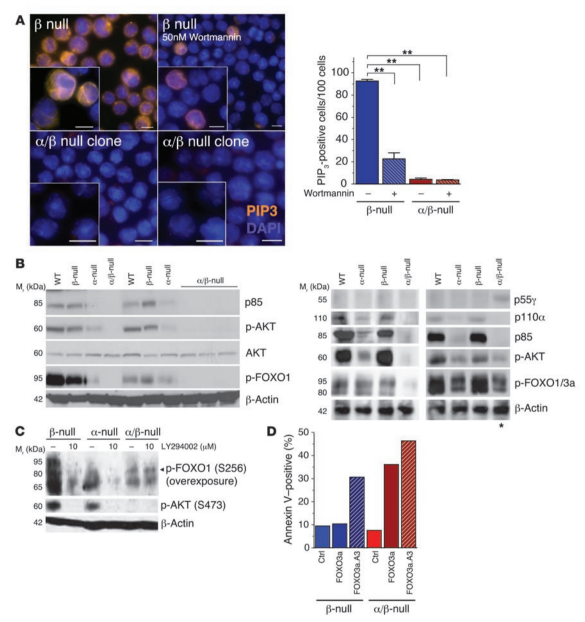


Figure 3

Lack of PI3K/AKT/FOXO functional signaling output in  $\alpha/\beta$ -null L-CFCs. (**A**)  $\beta$ -null and  $\alpha/\beta$ -null cells were fixed, permeabilized, and immunolabeled with anti-PIP $_3$  antibodies (orange). Nuclei were stained by DAPI (blue). Indicated cells were pretreated with 50 nM wortmannin 30 minutes prior to fixation. Immunoreactive cells were quantitated by counting 100 cells in images acquired with a 40× objective (represented in the graph) and visualized with a 100× objective (represented in the images). Typical examples of n=4 clones per genotype, 2 independent experiments (see also Supplemental Figure 2). Scale bars: 10.0 μm. \*\*P<0.001 versus untreated  $\beta$ -null cells; 1-way ANOVA. (**B**) Multiple clones of L-CFCs of the indicated genotypes were immunoblotted for class IA PI3K isoforms and assessed for phosphorylation of AKT (p-AKT S473) and its substrates FOXO1 and FOXO3a (p-FOXO1 T24/p-FOXO3a T32, right panel; or p-FOXO1 S256, left panel. Note: Anti-p-FOXO1 (S256) can detect p-FOXO4 (S193), but antibodies are not available to confirm mouse FOXO4 expression. Asterisk indicates representative  $\alpha/\beta$ -null L-CFC clone that upregulated p55γ, with a concomitant increase in AKT activity. (**C**) p190 L-CFCs of the indicated genotypes were treated for 15 minutes with LY294002 (10 μM) and subsequently immunoblotted for p-AKT (S473) and p-FOXO1/4 (S256/S193; note the overexposure with p-FOXO for detection of low signal). n=3 experiments using 2 different clones. (**D**) p190 L-CFCs were infected with retroviruses MSCV-IRES-Thy1.1 (Ctrl), MSCV-FOXO3a-IRES-Thy1.1, or MSCV-FOXO3a.A3-IRES-Thy1.1. Apoptosis was assessed in Thy1.1+ cells by annexin V/7AAD staining 48 hours after infection. Data are representative of 3 independent experiments using 2 different clones.

sensitive mTOR complex–1 (mTORC1), addition of rapamycin also caused a greater decrease in cycling  $\alpha/\beta$ -null cells compared with control cells (P < 0.01). These findings indicate that residual mTORC1 signaling in  $\alpha/\beta$ -null cells is required for proliferation in

the absence of class IA PI3K activity. The finding that rapamycin induced greater apoptosis in the  $\alpha/\beta$ -null cells, and that LY294002 caused apoptosis in cells of both genotypes, indicates that only the combined loss of the PI3K/AKT and mTORC1 pathways can lead to



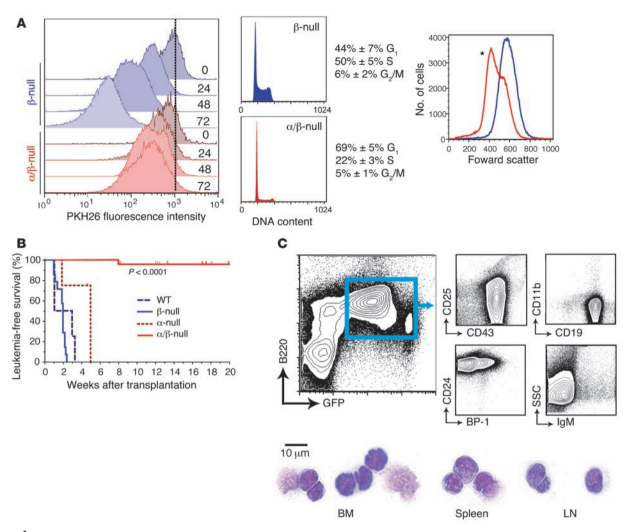


Figure 4 Severe loss of leukemogenic potential of  $\alpha/\beta$ -null L-CFCs. (A) Proliferation and cell cycle in  $\alpha/\beta$ -null L-CFCs were assessed in vitro, where  $\beta$ -null (blue) and  $\alpha/\beta$ -null (red) L-CFCs were labeled with 2 μM PKH26 and subsequently analyzed for dilution of PKH26 every 24 hours (left panel, representative of 2 independent experiments using 2 clones) and cellular size analyzed (right panel; \*P < 0.05, 2-tailed, paired Student's t test) by forward scatter; n = 3 clones. (B) Survival analysis of sublethally irradiated NOD/SCID mice transplanted with p190 L-CFCs (1 × 10 $^6$  cells). EGFP+ (or human CD4+) cells (>98% purity) from BM and spleen were collected to assess class IA PI3K deletion status (immunoblot analysis and leukemia-free survival were adjusted as described in Methods and Supplemental Figure 3 and Supplemental Table 1) (C) Representative immunophenotype and cytospins of  $\beta$ -null leukemic blasts in BM, spleen, and lymph nodes.

a substantial level of cell death in this system. Loss of PI3K/AKT signaling output also rendered p190 cells more sensitive to apoptosis caused by imatinib (Figure 5, A and B).

Considering that  $\alpha/\beta$ -null cells were hypersensitive to rapamycin, we examined the status and drug sensitivity of signaling events downstream of mTORC1, including potential feedback regulation of AKT. The serine/threonine kinase mTOR is regulated by both PI3K/AKT and other inputs (18, 19). We found that genetic or pharmacological inhibition of PI3K only partially reduced mTORC1 function as assessed by phosphorylation of ribosomal protein S6 (p-rS6) (Figure 6, A–D; also see Figure 2A for acute *Pik3r1* deletion). The residual p-rS6 was comparable to  $\beta$ -null cells treated with the pan-PI3K inhibitor wortmannin. Rapamycin suppressed p-rS6 almost completely in control and  $\alpha/\beta$ -null cells, confirming the presence of PI3K-independent inputs to mTORC1 activity in these cells. Rapamycin also greatly reduced the fraction

of phosphorylated 4EBP-1, a direct mTORC1 substrate (Figure 6, B and C), whereas 4EBP-1 phosphorylation was partly maintained in  $\alpha/\beta$ -null cells or wortmannin-treated controls.

The  $\alpha/\beta$ -null cells lacked p-AKT, while the control cells showed p-AKT that was sensitive to wortmannin, LY294002, and imatinib (Figure 6, A and B). Rapamycin caused an increase in AKT phosphorylation in control cells, with an accompanying increase in AKT activity as judged by FOXO phosphorylation (Figure 6, A–C). This "AKT rebound" phenomenon has been described in many cell contexts and is partly attributed to feedback inhibition of PI3K/AKT signaling by S6 kinases (S6Ks), which is prevented by rapamycin (18–20). A significant consequence of deleting both Pik3r1 and Pik3r2 was the complete loss of rapamycin-induced enhancement of AKT phosphorylation. This might explain the greater sensitivity of  $\alpha/\beta$ -null cells to cell cycle arrest and death following rapamycin treatment (Figure 5A).



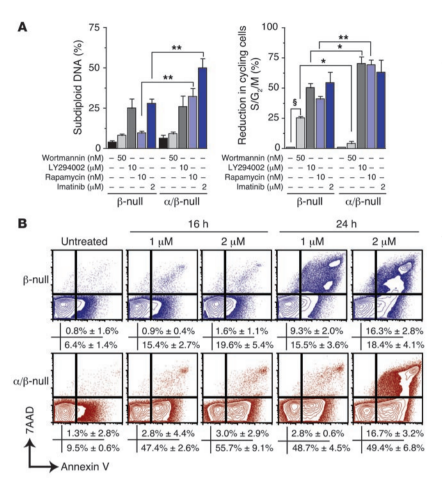


Figure 5

 $\alpha/\beta$ -null L-CFCs show altered drug sensitivity. (A) L-CFCs were treated with the indicated inhibitors for 24 hours and assessed for apoptosis (subdiploid DNA, left panel) and cycling cells (% S/G<sub>2</sub>/M phase, right panel) by flow cytometry. Differences in treatment effects between  $\beta$ -null and  $\alpha/\beta$ -null were analyzed by 2-way ANOVA (\*P < 0.05, \*\*P < 0.01) and 1-way ANOVA for wortmannin effects compared with untreated controls ( ${}^{\S}P$  < 0.001). Mean values ± SEM are shown; n = 4, using 4 different clones. (B) L-CFCs (blue,  $\beta$ -null; red,  $\alpha/\beta$ -null) were treated with or without 1  $\mu$ M or 2  $\mu$ M imatinib for 16 hours or 24 hours and analyzed for apoptosis. Cells that stained positive for annexin V and negative or positive for 7-AAD are reported as mean percentage ± SEM; n = 3 independent experiments.

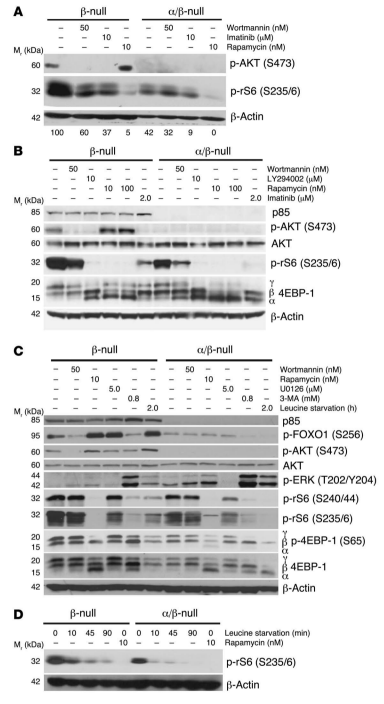
Class IA-independent mechanisms for mTOR pathway activation. To explore the mechanism of PI3K/AKT-independent mTOR activity, we assessed the role of 2 potential mTOR regulatory pathways: ERK activity and amino acid sensing. Basal ERK phosphorylation was consistently elevated in  $\alpha/\beta$ -null L-CFCs and blocked by treatment of cells with the MEK inhibitor U0126 (Figure 6C). However, MEK inhibition did not alter mTOR activity as judged by phosphorylation of 4EBP-1 (Figure 6C). p-rS6 expression was modestly reduced in both control and  $\alpha/\beta$ -null L-CFCs, likely due to stimulatory effects of ERK on S6K (21). We assessed the contribution of amino acid sensing by withdrawal of leucine from the culture media. Under these conditions, mTOR activity was rapidly extinguished in  $\alpha/\beta$ -null L-CFCs and to a lesser degree in control cells (Figure 6, C and D), as in other cell systems (16, 22). Amino acid sensing by mTOR is promoted by class III PI3K (hVPS34), an enzyme whose activity is sensitive to wortmannin (22). This might explain the partial inhibition of mTOR signaling by wortmannin in  $\alpha/\beta$ -null L-CFCs that lack class IA PI3K (Figure 6, A-C). Moreover, the class III PI3K inhibitor 3-methyladenine (3-MA) reduced mTOR activity (Figure 6C). In summary, residual mTORC1 activity in  $\alpha/\beta$ -null L-CFCs is ERK independent and sustained by amino acid sensing and other pathways that remain to be defined.

Multitargeted inhibition of pre–B-ALL proliferation with BCR-ABL kinase inhibitor and dual PI3K/mTOR kinase inhibitor. The residual proliferation and mTORC1 signaling in  $\alpha/\beta$ -null cells, together with the feedback activation of AKT in rapamycin-treated control cells, suggested that dual targeting of PI3K and mTOR might

be more effective than single targeting of either signaling node. A small molecule inhibitor, PI-103, selectively blocks the kinase activity of class I PI3K catalytic isoforms and both mTOR complexes at nanomolar concentrations (15, 23, 24). We treated p190 L-CFCs (from BALB/c BM) for 48 hours with multiple concentrations of imatinib, rapamycin, or PI-103, alone and in combination (Figure 7A). The percentage of viable cells was determined by MTS assay, and drug synergy was calculated using CalcuSyn software by the method of Chou (25). Both rapamycin and PI-103 suppressed survival with a maximal effect of approximately 50%, though rapamycin was approximately 70-fold more potent on a molar basis (Figure 7A). As previously reported (26), rapamycin showed slight to strong synergy with imatinib, except at lower concentrations of rapamycin (<40% IC<sub>50</sub>) where the combination produced antagonism (<20% IC<sub>50</sub>) to near additivity (<40% IC<sub>50</sub>). PI-103 combined with imatinib exhibited near additivity (<10% IC<sub>50</sub>) to strong synergy that was consistently greater than observed with rapamycin and imatinib.

To define the mechanism of action of drug combinations, we immunoblotted for p-AKT and mTOR activity over a time course of inhibitor treatment (Figure 7B). Rapamycin enhanced p-AKT during the first 6 hours of treatment, though this effect was lost at 24 and 48 hours. Simultaneous inhibition of PI3K and mTOR with PI-103 prevented this feedback rebound, such that p-AKT was effectively suppressed during the first 6 hours. Cells treated with imatinib plus rapamycin showed residual p-AKT that was eliminated in cells treated with imatinib plus PI-103. Also, imatinib





plus PI-103 suppressed mTORC1 signaling to a similar extent as imatinib plus rapamycin. Collectively, these results suggest that suppression of rebound AKT activity might explain the greater functional effects of imatinib plus PI-103.

We compared the efficacy of different drug combinations on BCR-ABL-driven leukemic cell expansion in vivo over a 2-day treatment course. BALB/c mice were lethally irradiated and reconstituted with a syngeneic radioprotective graft of BM cells and syngeneic p190 cells (marked by hCD4). After 11 days, when hCD4\* cells could be detected in peripheral blood (Supplemental Figure 4), mice were treated for 2 consecutive days with vehicle alone or sub-

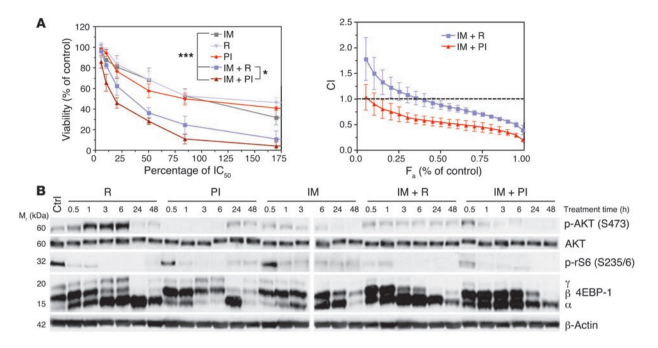
### Figure 6

 $\alpha/\beta$ -null L-CFCs maintain mTOR signaling via class IA PI3K-independent mechanisms. (A) p190 L-CFCs were treated for 2 hours with the indicated inhibitors, and the percentage of p-rS6 was quantified by ImageJ (upper p-rS6 band was normalized to β-actin; representative of 4 independent experiments and multiple clones). (B and C) L-CFCs were treated with the indicated inhibitors and/or leucine starved for 2 hours and immunoblotted with the indicated antibodies. n=3, using 3 clones. 3-MA, 3-methyladenine. (D) Cells of the indicated genotypes were starved in leucine-deficient medium (containing 10% dialyzed FCS) for the indicated times or treated with rapamycin (1.5-hour treatment). Control samples were incubated with leucine-sufficient medium (with matched dialyzed serum). n=3, using 2 clones.

optimal imatinib (70 mg/kg, twice per day), rapamycin (7 mg/kg, once per day), and PI-103 (40 mg/kg, twice per day), alone or in combination. Mice were then sacrificed, and several aspects of leukemic disease were assessed. Treatment with imatinib alone, compared with vehicle, significantly reduced leukemic disease as measured by spleen weight and percentage of hCD4+ cells in the peripheral blood (Figure 8 and Supplemental Figure 4). Imatinib alone also reduced the number of cycling hCD4+ cells in BM and spleen but caused no significant increase in apoptosis of hCD4+ cells. Rapamycin or PI-103 alone did not significantly reduce spleen weight but did reduce leukemic cell cycling, with variable effects on apoptosis. Imatinib plus PI-103 produced a significantly stronger antileukemic effect than imatinib alone, as measured by spleen weight. In accordance with this, imatinib plus PI-103 caused significantly greater cell cycle arrest and death in leukemia cells compared with imatinib alone. As previously reported (26), imatinib plus rapamycin was also effective and caused a significant increase in apoptosis compared with imatinib alone. However, the reduction in spleen weight and leukemic cell cycling was not statistically greater for imatinib plus rapamycin compared with imatinib alone. Thus, PI-103 but not rapamycin enhanced the effects of imatinib on spleen weight and BM leukemic cell cycling. Note that the PI-103 dose (80 mg/kg/d) was only 11-fold greater than the rapamycin dose (7 mg/kg/d), whereas the IC<sub>50</sub> of PI-103 in vitro is nearly 70-fold higher. Overall, these results support the in vitro data in showing that PI-103 is more effective than rapamycin in combining with imatinib to suppress leukemic cell expansion.

To extend these findings from the murine system, we compared the effects of PI-103 and rapamycin on primary human Ph<sup>+</sup> leukemias. CD19<sup>+</sup>CD34<sup>+</sup> cells were purified from peripheral blood of patients with Ph<sup>+</sup> ALL, PH<sup>+</sup> CML with lymphoid BC, or Ph<sup>-</sup> B-ALL, and clonogenic potential was assessed in the presence of various drug combinations. Ph<sup>-</sup> B-ALL samples were resistant to imatinib, as expected. In each case of Ph<sup>+</sup> disease, the presence of PI-103 combined with imatinib suppressed colony formation to a greater extent than imatinib alone or imatinib in combination with rapamycin; overall, these differences were statistically significant (Figure 9 and Supplemental Figure 5). Notably, the results from samples B and G also indicate that PI-103 can potentiate the





Attenuating PI3K and mTOR signaling together with BCR-ABL inactivation provides a greater synergistic response. (**A**) p190 L-CFCs were treated for 48 hours with the indicated inhibitors alone or in combination with imatinib (IM; left panel). Cell viability and IC<sub>50</sub> were determined using the MTS assay. \*P < 0.05, imatinib plus PI-103 (PI) versus imatinib plus rapamycin (R); \*\*\*P < 0.001, imatinib versus drug combination; repeated-measures 1-way ANOVA. The viability of cells treated with drug combinations at fixed ratios was plotted in the same graph (left panel), and drugs were assessed for synergy by calculating the combination index (CI) using CalcuSyn software (right panel). The CI is plotted as a function of the fractional growth inhibition (CI-F<sub>a</sub> plot using median-effect equation; ref. 25) by Monte Carlo simulation of 3 compiled independent experiments. CIs of less than 1, of 1, and of greater than 1 indicate synergism, additive effect, and antagonism, respectively. Mean  $\pm$  SD is shown; n = 3 independent experiments, using 2 separate clones. (**B**) p190 L-CFCs were treated for the indicated times with drug combinations, then cell lysates were subjected to immunoblot analysis using 2 gels run and developed concurrently with the same sample set. n = 2. Here we used concentrations of rapamycin (10 nM; ~120% IC<sub>50</sub>) and PI-103 (250 nM; ~45% IC<sub>50</sub>) that are achievable in vivo without excessive toxicity

(24, 26). Compounds were tested alone or in combination with a clinically achievable concentration of imatinib (2 µM; ~355% IC<sub>50</sub>).

antileukemic effect of imatinib in Ph<sup>+</sup> samples that are clinically resistant to this ABL inhibitor.

### **Discussion**

Pediatric and adult Ph<sup>+</sup> pre-B-ALL represents a subtype of leukemia with poor prognosis and rapidly acquires resistance to imatinib treatment during therapy. Induction regimens that target signaling pathways downstream of BCR-ABL have been hypothesized to be effective therapeutic strategies in patients that develop resistance to imatinib (3, 27). Here we have presented evidence that: (a) targeting the class IA PI3K subgroup strongly suppresses transformation and leukemogenesis in a mouse model of Ph<sup>+</sup> pre-B-ALL; and (b) dual inhibition of PI3K and mTOR is a particularly effective approach to suppressing survival of mouse pre-B-ALL cells and clonogenic expansion of human Ph<sup>+</sup> CD19<sup>+</sup>CD34<sup>+</sup> cells.

Increased PI3K/AKT signaling through activation of tyrosine kinases, loss of PTEN, or activating mutations in the p110 $\alpha$  catalytic subunit of PI3K occurs in a wide variety of malignancies (28). PI3K pathway activation drives proliferation, apoptotic resistance, and changes in metabolism. Cells expressing BCR-ABL show constitutive production of PIP3 and phosphorylation of AKT and its substrates (5) (Figure 10). A requirement for class IA PI3K activation, or "PI3K addiction," seems very likely in CML (7, 8) but has not been established in Ph $^+$  ALL. Previous work has suggested a

requirement for class IA PI3K in the murine pre-B-ALL model but has not distinguished the roles of different isoforms (5, 10, 29). Genetic ablation of *Pik3r1* encoding p85 $\alpha$ /p55 $\alpha$ /p50 $\alpha$  causes only partial inhibition of B lineage transformation by p190, and deletion of Pik3r2 (p85 $\beta$ ) has no effect (9). Here we show that loss of p85 $\beta$  in addition to p85 $\alpha$ /p55 $\alpha$ /p50 $\alpha$  reduces PI3K signaling output to an undetectable level. BM-derived progenitor B cells lacking both Pik3r1 and Pik3r2 showed greatly reduced transformation by p190, and outgrowth of nondeleting cells in this conditional Cre-loxP system indicates a strong selective pressure for PI3K signaling. The appearance of clones that upregulate the p55γ regulatory isoform supports the model of selective pressure for class IA PI3K expression and function. Acute deletion of *Pik3r1* also caused rapid cell cycle arrest and death, with greater effects in cells also lacking Pik3r2. Importantly, Pik3r1/Pik3r2 ablation essentially abolished leukemogenesis in an in vivo model, with a single mouse displaying a late-onset malignancy with p55γ upregulation. Thus, the additional signals received by leukemia cells in the living animal are not sufficient to bypass the requirement for class IA PI3K signaling.

Using the few  $\alpha/\beta$ -null L-CFCs that delete the floxed allele, we examined the status of signaling nodes downstream of PI3K/AKT. Most growth factors and many oncoproteins activate the mTOR signaling pathway in a PI3K/AKT-dependent manner (18, 19, 30).



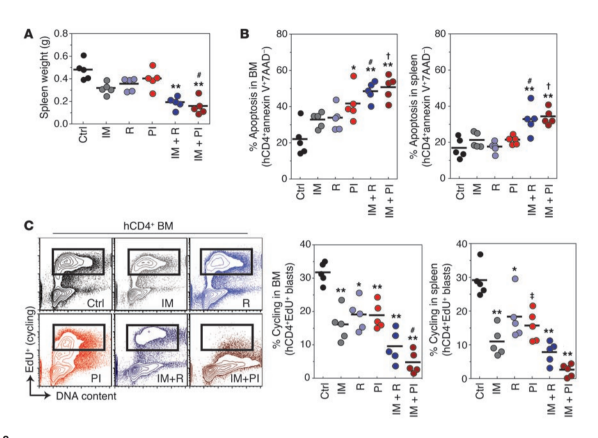


Figure 8

Short-term in vivo administration of Pl-103 combined with imatinib suppresses leukemic cell expansion more effectively than imatinib alone. (**A**) Lethally irradiated syngeneic recipients were transplanted with p190 L-CFCs (hCD4+) together with normal BM, then treated on days 11–13 after injection (i.p.; 2 days of treatment) with single and combination doses of imatinib (70 mg/kg, b.i.d.) and rapamycin (7 mg/kg, once per day) or imatinib and Pl-103 (40 mg/kg, b.i.d.) or double placebo control (diH<sub>2</sub>O and 75%/25% DMSO/saline mixture, b.i.d.) prior to sacrifice. Spleen weights after treatment regimen are reported. (**B**) Percent apoptosis in the BM and spleen was determined by annexin V/7AAD staining of hCD4+ blasts after 2-day treatment regimen. (**C**) Two hours prior to sacrifice, mice were injected with 5-ethynyl-2'-deoxyridine (EdU; i.p., 1.125 mg) and assayed for cycling cells that incorporated EdU in the BM and spleen with Click-iT EdU AF647 azide and propidium iodide staining (left and right panels). Representative depiction of cycling cells in the BM (left panel). \*P < 0.05, †P < 0.01, \*P < 0.001 versus control; \*P < 0.05, †P < 0.05, \*P < 0.001 versus imatinib; 1-way ANOVA; mean values  $\pm$  SEM are shown; P < 0.05 mice per treatment group.

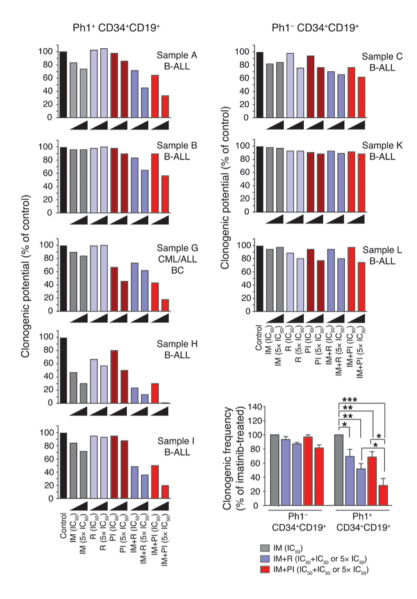
The mTOR serine/threonine kinase exists in 2 complexes. Complex 1 (mTORC1) is the form that is inhibited by rapamycin (Figure 10) and integrates signals from growth factors and nutrients to promote ribosome biogenesis, translation, and cell growth and proliferation. It is well established that BCR-ABL-expressing cells exhibit active mTORC1 signaling (26, 31), which is likely important to support the increased cell cycling and metabolic demands imposed by leukemic transformation. Surprisingly, we found that PI3K-deficient cells retain residual mTORC1 signaling, though at lower levels that correlated with smaller cell size. PI3K-deficient cells remained sensitive to rapamycin, establishing that mTORC1 has a required function even in the absence of AKT activity. The nutrient-sensing pathway independent of BCR-ABL appears to be essential to maintain mTORC1 activity in cells with low PI3K/AKT signaling. Further work is necessary to determine whether nutrient sensing drives residual mTORC1 activity or is merely permissive for another activating signal.

Although compounds that inhibit PI3K catalysis are considered promising therapeutics, our work suggests likely benefits of developing agents that can interfere with the function of class IA regulatory subunits. In this regard, it is important to consider that

class IA regulatory isoforms might have scaffolding functions in oncogenic signaling that are independent of their role in localizing and activating the catalytic subunits (32). We consider this unlikely in the murine pre–B-ALL system, as clones upregulating p55 $\gamma$  displayed normal growth characteristics, which correlated with maintenance of AKT phosphorylation. p55 $\gamma$  lacks the N-terminal modules (SH3, RhoGAP-homology) that are required for known scaffolding functions of p85 $\alpha$  while retaining conserved SH2 domains required for recruitment to tyrosine-phosphorylated BCR-ABL substrates. These observations imply that potential Ph<sup>+</sup> leukemia therapies targeting class IA regulatory subunits might be most effective if designed to interfere with conserved features of the different isoforms.

Rapamycin and related mTORC1 inhibitors are clinically important immunosuppressive agents that are currently in trials for anticancer activity (30, 33). In Ph+ leukemia, rapamycin analogs are being tested in combination with imatinib. A key concern with this approach is that many tumors treated with mTORC1 inhibitors show elevated AKT activity, suggesting a mechanism for resistance to therapy. Blocking mTORC1 can augment mTORC2 activity, recently shown to mediate phosphorylation of AKT on serine 473





(34). mTORC1 inhibition also reduces S6K1 activity, disrupting the negative feedback phosphorylation of signaling adaptor proteins. Either of these mechanisms can lead to rebound enhancement of AKT function and induce a recovery of prosurvival signals (Figure 10). High AKT activity is associated with a poor prognosis as well as resistance to chemotherapy (35, 36).

To achieve blockade of PI3K, AKT, and mTOR simultaneously, we used the multitargeted inhibitor PI-103 (Figure 10). This compound has similar potency against class IA PI3K catalytic subunits and both mTOR complexes and blocks proliferation of many human cancer cell lines (15, 24). We found that PI-103 is less potent than rapamycin on a molar basis but synergizes significantly better with imatinib to suppress the proliferation and survival of p190-transformed L-CFCs. Supporting the potential of this compound for cancer therapy, PI-103 suppresses solid tumor growth in xenograft models more potently than agents targeting PI3K or mTOR alone (23, 24). Although the doses of PI-103 used in these mouse studies are well tolerated, the compound has unfavorable pharmacokinetics (24). Hence, development of similar compounds with greater potency and better drug properties should be a prior-

### Figure 9

Anticlonogenic effects of PI-103 combined with imatinib in Ph+ leukemia progenitors. Primary Ph+ leukemia samples (Ph+ B-ALLCD34+/CD19+, n = 5; Ph+ CML/ALL-BC<sup>CD34+/CD19+</sup>, n = 1; and Ph<sup>-</sup> ALL<sup>CD34+/CD19+</sup>, n = 3) were assessed for colony formation potential with or without single and combination treatments of imatinib, rapamycin, or PI-103. 100% clonogenic potential was normalized to vehicle-treated samples. Clonogenic frequencies of combination treatments were compared with that of imatinib alone (expressed as percent of imatinib-treated). \*P < 0.05, \*\*P < 0.01, \*\*\*P < 0.001; 2-way ANOVA; mean values  $\pm$  SEM are shown; Ph<sup>+</sup>, n = 5 and Ph<sup>-</sup>, n = 3, comparing the source of variation from Ph<sup>+/-</sup> and treatment. Note that in the combination treatments, the imatinib concentration was fixed at its IC<sub>50</sub>. Cytogeneics, cytospins, and flow cytometric markers of human specimens are given in Supplemental Table 2.

ity. It will also be important to distinguish whether the potency of PI-103 relative to rapamycin arises from more sustained suppression of AKT activation or from other mechanisms.

Various pharmaceutical companies have drug discovery efforts to develop compounds targeting specific PI3K isoforms or subgroups, especially within the class IA and IB enzyme subfamilies. Our data establish that class IA PI3K signaling contributes quantitatively to BCR-ABL transformation and is essential for in vivo leukemogenesis. Moreover, even in the rare event when cells escape PI3K addiction, they proliferate slowly and remain dependent on mTORC1. Our genetic and pharmacological data in murine and human pre-B-ALL cells suggest that targeting the PI3K pathway at multiple nodes will provide greater therapeutic efficacy.

### **Methods**

Antibodies and other reagents. A list of antibodies is provided in Supplemental Methods. PI-103 was synthesized as described previously (15, 23). Other pharmacological

inhibitors were obtained from commercial sources: LY294002, U0126, 3-MA (Sigma-Aldrich), rapamycin (LC Laboratories), imatinib (Novartis), and wortmannin (Calbiochem).

Retroviral vectors and generation of virus stocks. Bicistronic retroviral vectors were used to express exogenous genes and a marker gene from the murine stem cell virus (MSCV) long terminal repeat. Details of these vectors and preparation of virus stocks are provided in Supplemental Methods.

*Mice.* The generation of *Pik3r1*-flox and mice lacking *Pik3r2* is described in detail elsewhere (12, 13, 37). These strains were kindly provided by Lewis Cantley (Harvard University, Boston, Massachusetts, USA) and crossed to CD19-Cre knock-in mice (gift of Robert Rickert, Burnham Institute, La Jolla, California, USA). Mice were maintained in a mixed background (C57BL/6 × 129SvEv) for the experiments, and BM was harvested when animals were 3–6 weeks of age. NOD.CB17-*Prkdcscid*/J (NOD/SCID) and BALB/c mice were purchased from The Jackson Laboratory. All animal protocols were approved by the Institutional Animal Care and Use Committee of University of California, Irvine.

BM progenitor cell transduction and establishment of p190 BCR-ABL L-CFCs. BM cells were flushed from the long bones (tibias and femurs) of 3- to 4-week-old mice. Cells were spinoculated with retroviral supernatant in



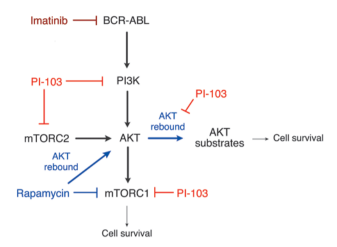


Figure 10

Schematic diagram of PI3K and mTOR signaling and inhibitor action in BCR-ABL—transformed pre—B-ALL. Rapamycin inhibits mTORC1 but leads to rebound AKT activation through either (a) increased mTORC2 activity or (b) loss of S6K-mediated feedback inhibition of PI3K recruitment to signaling adaptors. Not shown: ablation of PI3K prevents activation of AKT but not mTORC1. Dual inhibition with PI-103 overcomes PI3K-independent mTORC1 activity and prevents rebound activation of AKT.

the presence of 5 µg/ml polybrene for 45 minutes at 450 g and 37°C with RPMI-20 culture medium (see Supplemental Methods) supplemented with recombinant mouse IL-7 (10 ng/ml; R&D Systems) to promote cell cycle entry. Following spinoculation, cells were incubated overnight at 37°C, 5% CO<sub>2</sub> and plated (5 × 10<sup>4</sup> cells) in M3630 methylcellulose cultures (Stem-Cell Technologies) as described previously (9). Cells were then selected, expanded, and infected as described further in Supplemental Methods.

Immunoblotting. Immunoblotting was performed using standard procedures (see Supplemental Methods). Where indicated, cells were subjected to amino acid starvation as described in Supplemental Methods. In Figure 6A, ImageJ (http://rsbweb.nih.gov/ij/) was used to quantitate band intensity.

Flow cytometry and immunofluorescence microscopy. FACS acquisition was performed on a BD FACSCalibur or LSRII, and surface markers were detected with fluorescent mouse- or human-specific antibodies as detailed in Supplemental Methods.

In vitro drug combination studies. p190 L-CFCs (generated from M3630 clonal cultures of 3-week-old BALB/c BM;  $2 \times 10^4$  cells) were incubated in triplicate wells of 96-well culture plates for 48 hours in the presence of increasing concentrations of imatinib, rapamycin, and PI-103. Cell viability and IC<sub>50</sub> were determined using the MTS assay (CellTiter 96 AQ<sub>ueous</sub> One Solution Cell Proliferation Assay kit; Promega). Absorbance values were normalized to vehicle control-treated (0.2% DMSO) cells and expressed as percent cell viability. A range of fixed ratios of drug combinations was employed to assess synergy using the combination index (CI) with Calcu-Syn software (Biosoft) using the median-effect method of Chou (25). Potency (IC<sub>50</sub>), shape (m, coefficient of the slope equation), and the linear correlation coefficient (r) parameters of dose-effect curves for imatinib (IC<sub>50</sub>= 563 ± 17 nM, m = 1.178, r = 0.98, p = 3), rapamycin (IC<sub>50</sub>= 8.5 ± 2.5 nM, m = 1.083, r = 0.97, p = 3), and PI-103 (IC<sub>50</sub>= 592 ± 38 nM, m = 1.030, p = 0.98, p = 3) were calculated from 6 dose ranges (5%–350% of IC<sub>50</sub>).

Transplantation and in vivo drug delivery studies. For studies involving transplantation of p190 L-CFCs generated from Pik3r1-floxed mice, single colonies from 7-day M3630 methylcellulose cultures were expanded in RPMI-20

and assessed for immunophenotype, and for expression of p85 $\alpha$ , p85 $\beta$ , and p55 $\gamma$  and p-AKT/FOXO status by immunoblotting before in vivo transplantation. Following a 2-week in vitro expansion,  $1\times 10^6$  cells were injected into the retroorbital venous sinus of 6- to 10-week-old NOD/SCID mice that had been sublethally irradiated with 240 cGy of total body  $\gamma$ -irradiation. For studies involving in vivo drug treatment of animals, the BM was flushed from 3-week-old BALB/c mice and spinoculated with retroviral supernatants (MSCV-BCR-ABL-IRES-hCD4) as plated in colony assays, expanded, and injected into to 8-week-old syngeneic BALB/c mice. Further details are provided in Supplemental Methods.

Primary human leukemia samples. All human leukemia specimens were obtained by one of the authors (M.B. Lilly) while treating the subjects at Loma Linda Medical Center, Loma Linda, California, USA. Heparinized peripheral blood was collected during the course of routine clinical care or as part of a Loma Linda Medical Center Institutional Review Board-approved specimen bank protocol. Mononuclear cells were then isolated by density gradient centrifugation, suspended in 90% FCS/10% DMSO, and stored in liquid nitrogen. Their use for these studies was approved by the University of California Irvine Institutional Review Board, with the stipulation that the samples lacked personal identifiers and that the donors were either deceased or had initially signed a specimen banking consent. Cryopreserved samples (90% FCS/10% DMSO) of mononuclear cells from the peripheral blood of patients with Ph<sup>+</sup> ALL, Ph<sup>+</sup> CML with lymphoid BC, or Ph<sup>-</sup> B-ALL were thawed, cultured overnight, and plated in methylcellulose as detailed in Supplemental Methods.

Statistics. GraphPad Prism (version 4.0c) software was used for all statistical analyses. Homogeneity of variances was calculated by Bartlett's test. Random continuous variables were analyzed using 2-tailed Student's ttest, 1-way ANOVA, and 2-way ANOVA, where indicated. Bonferroni post-hoc analysis was used throughout, and the Tukey post-hoc test was used in Figure 9. Survival curves were compared using the log-rank test. Data are reported as the mean ± SEM, unless otherwise indicated. A P value less than 0.05 was considered significant.

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# research article



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