Synergism of FAK and tyrosine kinase inhibition in Ph⁺ B-ALL

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Introduction

Acute lymphoblastic leukemia (ALL) comprises a number of subtypes defined by constellations of founding chromosomal alterations, structural alterations, and mutations that influence leukemogenesis and treatment outcome. BCR-ABL1⁺ (Ph⁺) B progenitor ALL (B-ALL) constitutes 3% to 5% of childhood and up to one-third of adult ALL cases. It is a highly aggressive disease that is often refractory to currently available therapies. Tyrosine kinase inhibitors (TKIs) that target the constitutively ABL1 kinase have revolutionized the treatment of Ph⁺ chronic myeloid leukemia but have only partly improved the poor prognosis of Ph⁺ ALL. Genomic profiling studies have provided important insights into the genetic basis of these differences in outcome. In contrast to CML, which at chronic phase harbors few if any additional genomic alterations, Ph⁺ ALL is characterized by alterations of the lymphoid transcription factor gene, \textit{IKZF1} (1, 2). In both Ph⁺ and Ph⁻ ALL, \textit{IKZF1} alterations are associated with inferior treatment outcome (3). Additional genetic alterations, including deletion of the \textit{CDKN2A/CDKN2B} (INK4/ARF) tumor suppressor locus and deletions of \textit{PAX5}, are also common in Ph⁺ ALL.

In mouse models of Ph⁺ ALL, perturbations of \textit{Ikaros}, including loss-of-function deletions and expression of the dominant-negative isoform IK6 (deletion of exons 4–7), result in the acquisition of stem cell–like features, enhanced self-renewal, overexpression of adhesion molecules, and upregulation of focal adhesion kinase (FAK), in part due to the derepression of stem cell and adhesion genes normally silenced by IKZF1 (4, 5). This results in leukemic cell adhesion to other tumor cells as well as stromal cells of the bone marrow microenvironment, aberrant localization of leukemic cells to the periarteriolar bone marrow niche, and reduced sensitivity to therapy (4). A detailed understanding of the interactions between leukemic cells and the bone marrow microenvironment is important to fully understand leukemogenesis, tumor maintenance, and resistance to therapy. These recent insights indicate that aberrant interactions of leukemic cells with the microenvironment are dictated by genetic alterations in leukemic cells and that targeting pathways that influence the tumor cell microenvironment interactions may prove indispensable in improving responses to standard therapies and reducing the incidence of relapse for high-risk, \textit{IKZF1}-mutated Ph⁺ B-ALL.
RESULTS

FAK pathway upregulation in Ph+ B-ALL, with further activation in Ikzf1-mutated Ph+ B-ALL cells. Gene expression profiling of primary mouse Arf−/− pre-B cells expressing empty vector, BCR-ABL1, or EBF1-PDGFRB revealed that Fak1 is overexpressed in Ph+ cells (Figure 1A). Expression of EBF1-PDGFRB, a fusion identified in Ph-like B-ALL that results in constitutive kinase activity, cytokine-independent cell line proliferation, and leukemogenesis (12), did not result in upregulation of Fak1 (Figure 1A). The expression of the dominant-negative Ikaros isoform, IK6, in combination with BCR-ABL1 resulted in further upregulation of Fak1, whereas IK6 did not increase Fak1 expression in empty vector or EBF1-PDGFRB-expressing pre-B cells (Figure 1A). Interestingly, IK6 expression upregulated Fak2 (also known as Ptk2B or Pyk2) in non-Ph+ cells but not in BCR-ABL1–expressing cells (Figure 1, A and B). RNA (mRNA) and protein sequencing confirmed that FAK1 was overexpressed in IK6-expressing Ph+ pre-B cells, and FAK2 expression was relatively high in all BCR-ABL1–expressing cells (Figure 1, B and C). Although a broad range of FAK1 expression was observed by mRNA sequencing of 19 Ph+ and 109 Ph-like patient samples regardless of IKZF1 status (Figure 1D), activation of the pathway was revealed by reverse-phase protein array in human Ph+ IK6+ B-ALL cells compared with Ph+ cells with intact IKZF1 (Figure 1E).

FAK inhibition decreases survival, clonogenicity, and adhesion of Ikzf1-mutated Ph+ B-ALL cells. As a single agent, VS-4718 is cytotoxic to primary human Ph+ IK6-expressing B-ALL cells and mouse Arf−/− or ikzf1−/− pre-B cells expressing BCR-ABL1 with and without IK6 or a dominant-negative IKZF1 point mutation, D186A, at concentrations above 0.1 μM (Figure 2A). Ikzf1 haploinsufficiency and IK6 expression resulted in less sensitivity to FAK inhibition compared with cells with intact Ikzf1, whereas cells harboring IKZF1 D186A were as sensitive as cells expressing empty GFP vector (MSCV-IRES-GFP [MIG]; Figure 2A). Human Ph+ cells underwent apoptosis in response to exposure to VS-4718 ex vivo (Figure 2B).

Ikef1 alterations confer aberrant hematopoietic stem cell–like properties to Ph+ pre-B cells, including increased self-renewal and adhesion to ECMs in vitro and stromal cells in the bone marrow niche (4). VS-4718 attenuated the capability of cells with or without Ikef1 alterations to form colonies from single cells at concentrations that do not affect cell viability (Figure 2C). FAK inhibition also blocked adhesion to fibronectin fragment monolayers regardless of Ikef1 status; although the greatest effects were observed in cells harboring Ikef1 alterations (Figure 2D). VS-4718 treatment inhibited FAK pathway signaling, as revealed by reverse-phase protein array analysis of key targets in human Ph+ IK6+ B-ALL cells (Figure 2E).

In the bone marrow niche, Arf−/− Ph+ IK6+ leukemic cells adopt a spindle-like adherent phenotype, in stark contrast to the typical spherical shape of pre-B cells (4). To visualize the effects of VS-4718 on adherence of cells in the bone marrow niche, we transplanted GFP-labeled Arf−/− Ph+ IK6+ cells into recipient mice that were randomized and treated daily with vehicle or VS-4718 for 3 days, starting 24 hours after transplant. Calvaria were harvested and immediately imaged by multiphoton microscopy. Vehicle-treated mice displayed the adherent, irregularly shaped spindle-like morphology, whereas VS-4718 treatment abrogated the adhesive phenotype of the cells, without directly affecting engraftment or viability of cells (Figure...
Quantification of the spindle-like or typical round morphology of cells revealed a reduction in the number of cells with an adherent, irregular appearance in the bone marrow niche after VS-4718 treatment (Figure 3C).

VS-4718 synergizes with dasatinib in affecting cell survival, adhesion, and inhibition of downstream targets of FAK. Dasatinib is a TKI that targets BCR-ABL1; it is widely used in frontline treatment of Ph⁺ B-ALL. To assess the combinatorial effects of dasatinib and FAK inhibition, we performed cell viability assays using Arf–/– Ph+ pre-B cells expressing either empty vector (MIG) or IK6 at fixed drug ratios to determine combination index (CI) values. Dasatinib synergized with VS-4718 in decreasing the survival of cells at increasing concentrations of drug (Figure 4A). CI values ranged from 0.68 to 0.91 for cells expressing empty vector (MIG), suggesting synergistic to additive effects, whereas IK6-expressing cells displayed CI values indicative of strong synergistic effects of combination therapy (Figure 4B). Four primary human Ph⁺ B-ALL lines harboring IKZF1 alterations (ALL-4 [IK6], ALL-55 [deletion of exons 2–7], ALL-56 [deletion of exons 2–7], and h9407 [IK6] cells) displayed highly variable responses to dasatinib and VS-4718 as single agents ex vivo; however, exposure to both drugs decreased the IC₅₀ values compared with values after exposure to either alone for all lines except ALL-55, which was refractory to both drugs (Figure 4, C and D).

Given that VS-4718 and dasatinib synergize in decreasing cell viability, we explored their effects on cell adhesion by exposing Arf⁻/⁻ Ph⁺ pre-B cells expressing either empty vector (MIG) or IK6 to RetroNectin
monolayers after single and combination treatments. As expected, IK6-expressing cells were more adherent after a short (30-minute) exposure to RetroNectin than their empty vector–expressing counterparts (ref. 4 and Figure 4E). Alone, FAK inhibition with VS-4718 or ABL inhibition with dasatinib significantly reduced the ability of IK6-expressing cells to adhere to RetroNectin, whereas the number of adherent MIG-expressing cells was slightly, yet not significantly reduced after VS-4718 or dasatinib treatment alone.
In contrast, the ability of both MIG- and IK6-expressing cells to adhere to Retronectin was significantly attenuated following combination dasatinib and VS-4718 treatment, and the reduction was greater than that with either drug alone (Figure 4E). Enhanced inhibition of p-P130Cas, a downstream target of FAK, was observed in the adherent fraction of cells, chosen due to maximal activation of the pathway by ECM exposure at 15 minutes, 30 minutes, and 2 hours after treatment with a combination of VS-4718 and dasatinib on Retronectin (Figure 4F).

Combination treatment with VS-4718 and dasatinib improves therapeutic outcome of Ph+ B-ALL. For in vivo efficacy studies, BCR-ABL1–transduced pre-B cells expressing luciferase were derived from Arf−/− donor mice and subsequently transduced with either empty GFP vector (MIG) or IK6-GFP, sorted, and injected into sublethally irradiated wild-type recipients. Mice were monitored for leukemic burden by detecting luciferase activity in vivo by live animal bioluminescence imaging (Figure 5A). By day 7, mice had measurable tumor burden and treatment was commenced. Untreated animals from both empty vector and IK6 groups succumbed to disease by day 14 (4), and VS-4718 treatment alone did not affect leukemic burden or survival (data not shown). Although inefficacious alone, VS-4718 in combination with dasatinib dramatically decreased leukemic burden and extended the lives of both MIG and IK6 cohorts, with one long-term survivor from the IK6 group that underwent a complete remission, despite the cessation of treatment (Figure 5, B and C). At the time of sacrifice for moribund animals that succumbed to disease, spleen weights were slightly reduced in combination-treated mice (Figure 5D).

To further test the efficacy of VS-4718 in vivo, human Ph+ IK6+ (h9407) cells were xenografted into NOD/SCID γ-null (NOD.Cg-Prkdcscid Il2rgtm1Wjl/SzJ [NSG]) mice, harvested, marked with luciferase ex vivo, retransplanted into mice for expansion, and then harvested, sorted, and retransplanted for in vivo efficacy studies tracing leukemic burden by monitoring luciferase activity. Preliminary studies revealed that VS-4718 was not effective as a single agent (Figure 6A); however, in combination with dasatinib, FAK
inhibition significantly reduced leukemic burden (Figure 6B). Survival was not a reliable readout for drug efficacy using h9407 cells, as mice were able to harbor high leukemic burden without manifesting symptoms of illness; hence, luciferase quantifications were employed as a measure of efficacy, and animals were sacrificed at the end of the study. Activity against a second human Ph⁺ IK6⁺ B-ALL line, ALL-4, was evaluated by monitoring human CD45 (hCD45) in peripheral blood, and survival was commensurate with disease burden. Again, VS-4718 provided no benefit as a single agent; however, it displayed remarkable efficacy in combination with dasatinib, whereby disease progression was markedly slowed and survival was prolonged on combination therapy (Figure 6, C and D). The number of circulating and

Figure 4. VS-4718 synergizes with dasatinib to decrease cell survival and adhesion of BCR-ABL1 B progenitor acute lymphoblastic leukemia cells. (A) VS-4718 synergizes with dasatinib to lower the IC₅₀ of Arf⁻/⁻ BCR-ABL1 (Ph) pre-B cells expressing MIG or IK6 in cell viability assays. (B) Combination index (CI) values from viability assays shown in A indicate synergistic to additive effects of VS-4718 and dasatinib on MIG cells and strong synergy for IK6 cells (CI ≤ 0.3 = strong synergy; 0.3 < CI ≤ 0.85 = synergy; 0.85 < CI ≤ 1.2 = additive; 1.2 < CI ≤ 3.3 = antagonism; CI > 3.3 = strong antagonism). (C) Cell viability assays with human ALL-4 and ALL-56 cells displayed synergism between VS-4718 and dasatinib; however, the survival of ALL-55 cells was not decreased with combination treatment. (D) Human h9407 B progenitor acute lymphoblastic leukemia cells are less sensitive to increasing concentrations of dasatinib alone compared with combination treatment with VS-4718 in cell viability assays ex vivo. (E) Adhesion assays were used to detect the effects of VS-4718 and dasatinib on the ability of Arf⁻/⁻ Ph pre-B cells expressing MIG or IK6 to adhere to RetroNectin monolayers in vitro. Combination treatment with VS-4718 and dasatinib synergistically abolished cellular adhesion to RetroNectin. (F) Downstream FAK signaling is inhibited synergistically by VS-4718 and dasatinib in the adherent fraction of cells from the RetroNectin assays shown in E, as detected by Western blotting for p-P130Cas expression at 15 minutes, 30 minutes, and 2 hours of exposure to drug. Data represent averages ± SD of 3 technical replicates for each group in A–E; *P ≤ 0.05, **P ≤ 0.005, ***P ≤ 0.0005, 2-tailed Student’s t test. MIG, MSCV-IRES-GFP (empty GFP vector); Das, dasatinib.
Figure 5. Combination VS-4718 and dasatinib therapy decreases leukemic burden and increases survival of murine BCR-ABL1 B progenitor acute lymphoblastic leukemia. (A) Live whole-animal imaging of representative mice transplanted with Arf−/− BCR-ABL1 (Ph) pre-B cells expressing luciferase, with and without IK6, used for testing the in vivo efficacy of VS-4718 and dasatinib against murine Ph+ B progenitor acute lymphoblastic leukemia. (B) Quantification of luciferase activity to monitor the progression of disease during treatment. (C) Kaplan-Meier survival curve displaying increased survival on combination VS-4718 and dasatinib therapy, with one long-term remission achieved until the surviving mouse was sacrificed with no indication of leukemia at day 200. (D) Combination treatment slightly reduced spleen weight, as observed at the time of sacrifice when mice were moribund. n = 10 mice per group; *P ≤ 0.05 Student’s t test. ***P ≤ 0.0005, Mantel-Cox test. MIG, MSCV-IRES-GFP (empty GFP vector); Das, dasatinib
organ-infiltrating hCD45+ blasts was reduced to nearly undetectable levels at day 28 after treatment with dasatinib and VS-4718 (Figure 6, C and E). Although a durable remission was achieved on combination therapy, mice relapsed between 1 and 3 weeks after completion of treatment on day 28 and ultimately succumbed to florid disease (Figure 6, C–E). Analyses of spleen samples by Western blot and ELISA confirmed that phosphorylation of FAK at residue Y397 (pFAK-Y397), which is indicative of FAK activation, was inhibited by in vivo treatment with VS-4718 (Figure 6, F and G).

**Discussion**

Here, we have shown that FAK inhibition attenuates the aberrant adhesive and stem cell–like phenotypes induced by IKZF1 alterations in Ph+ ALL and show that combined therapy with FAK inhibitors and the ABL inhibitor dasatinib is remarkably effective in abrogating leukemic cell growth in vivo. Indeed, this combinatorial approach induced durable remission without recurrence following cessation of therapy.

Genetic alterations resulting in loss of function or dominant-negative inhibition of B lineage transcription factors are a hallmark of B progenitor ALL (13), with significant differences observed in the nature of alteration between B-ALL subtypes. Notably, favorable risk subtypes such as *ETV6-RUNX1* ALL harbor deletions of *PAX5*, which is required for B lineage commitment and maintenance, in one-third of cases but rarely have alterations of *IKZF1*. In contrast, Ph+ ALL, Ph-like ALL, and *ERG*-dysregulated ALL have a high frequency of *IKZF1* alterations (1, 2). In contrast to *IKZF1*, *PAX5* alterations, which are associated with poor prognosis in Ph+ and Ph-like ALL, are not consistently associated with poor outcome, suggesting distinct roles of each genetic alteration in leukemogenesis and responsiveness to therapy.

Multiple studies have shown that loss of function of *PAX5*, *EBF1*, and *IKZF1* increases the penetrance of mouse B-ALL either in Ph-driven spontaneous or mutagenesis-primed models (14–16). In Ph+ ALL, loss of *IKZF1* activity shifts the phenotype from a myeloid, CML-like disease to an aggressive, transplantable lymphoid leukemia, without expanding the pool of lymphoid precursors (4). Alterations of *IKZF1*, including loss-of-function deletions, dominant-negative alleles, and dominant-negative point mutations, also result in resistance to TKI-based therapy independent of inhibition of the ABL1 kinase (4).

The mechanistic basis of *IKZF1*-mediated resistance to therapy has now been elucidated. *IKZF1* is a zinc finger transcription factor that mediates transcriptional activation and repression, in part through interactions and recruitment of histone-modifying complexes, such as the nucleosome remodeling and deacetylase complex (17, 18), transcriptional elongation complex positive-transcription elongation factor b (19), and polycomb repressor complex 2 (20). The dominant-negative deletions and mutations of *IKZF1* remove or perturb function of the N-terminal zinc fingers, resulting in loss of DNA-binding activity but retention of dimerization, resulting in cellular mis localization (4, 21). The resulting loss or inhibition of activity dysregulates and activates multiple cellular pathways, including reactivation of hematopoietic stem cell and adhesion pathways (4). Key adhesion molecules expressed include CD90 (THY-1) and integrins, particularly ITGA5, which, upon engagement of ligand, stimulate intracellular signaling pathways (4, 5). FAK expression is elevated in *IKZF1*-mutated leukemias, and phosphorylation of FAK itself, as well as downstream effectors, is likely augmented by the direct action of activated ABL1 itself.

We previously showed that this phenotype was accompanied by aberrant self-renewal and markedly enhanced intracellular and cell-stromal adhesion, resulting in extravasation and relocalization of leukemia cells to a periarteriolar location in the bone marrow niche (4), which in part may account for the aggressive phenotype of *IKZF1*-mutated ALL observed in mouse models characterized by extramedullary invasion and aggressive central nervous system infiltration. Using high-content screening, we showed that retinoids, specifically retinoid X receptor agonists, potently reversed this phenotype, abrogating self-renewal and inducing differentiation. In part, retinoids exerted this effect by inducing expression of wild-type IKZF1, relocalizing the protein to the nucleus, suppressing expression of stem cell and adhesion molecules, and abrogating adhesion. This also suggested that approaches directly targeting these adhesion pathways may also be efficacious.

Our results confirm this, showing that FAK inhibition blocks phosphorylation of FAK and downstream effectors. With the exception of FLT3, VS-4718 is specific for FAK at low concentrations in vitro (0.1 μM), with partial inhibition of additional kinases at higher concentrations, including ACK1 (activated Cdc42-associated tyrosine kinase 1), aurora-A, CDK2 (cyclin-dependent kinase 2), insulin receptor, LCK (lymphocyte-specific protein tyrosine kinase), and TRKA (tropomyosin-related kinase A) (22). Importantly, little single-agent activity was observed, whereas the combination of dasatinib and FAK inhibition
was potently effective in killing leukemic cells and reversing adhesion in vitro and in vivo. A significant, although less striking, effect was observed in IKZF1 wild-type, Ph+ ALL, supporting the notion that BCR-ABL1 may directly activate FAK signaling and that FAK inhibition may be efficacious in all cases of ALL, regardless of IKZF1 status.

These results provide a compelling rationale for clinical evaluation of combined ABL and FAK inhibition in Ph+ ALL. Single-agent ABL1 inhibition has transformed the outcome of Ph+ CML, resulting in deep and prolonged remissions in the majority of patients (23). However, Ph+ CML is a distinct disease from ALL, in which Ph alone is not only required but is sufficient to induce a myeloproliferative neoplasm,
and additional recurring genetic lesions are observed at transition to blast crisis, including \textit{IKZF1} at progression to ALL (2, 24). Ph+ ALL exhibits only transient responses to single-agent TKI therapy, and disease progression is frequent with chemotherapy alone (25–27). The combination of TKI with chemotherapy has improved outcomes (28), yet these remain inferior to non-IKZF1-mutated ALL (29, 30). Recent preclinical trials using the same \textit{Arf}^{−/−} BCR-ABL1 pre-B model of B-ALL used in these studies suggest that pairing dasatinib with the Janus kinase inhibitor ruxolitinib and chemotherapy (dexamethasone) is a promising therapeutic approach in Ph+ B-ALL, although the 2-drug combination of dasatinib and ruxolitinib was only modestly more effective than dasatinib alone (31). These experimental findings and clinical observations strongly suggest that directly disrupting the perturbations induced by IKZF1 alterations, either by restoring expression with retinoids or by disrupting the effects of integrin and FAK signaling, represent a powerful approach to improve outcome in this disease.

**Methods**

**Expression vectors and retroviral production.** cDNAs encoding BCR-ABL1 were cloned into mouse stem cell virus–internal ribosome entry site–GFP (MSCV-IRES-GFP [MIG]) (32) or luciferase (MSCV-IRES-luc) vectors (33). cDNAs encoding wild-type dominant-negative IK6 were PCR amplified from human leukemic cell RNA (2) and cloned into MIG. Vectors were packaged into replication-incompetent ecotropic retroviral particles by transient transfection of 293T cells with a triple plasmid (pMD old gag pol and pCAG4-Eco) system.

**Gene transduction and transplantation.** MSCV vectors expressing human BCR-ABL1 and luciferase were used for transduction of whole bone marrow from \textit{Arf}^{−/−} or \textit{Ikzf1}^{+/−} (34, 35) mice and generation of in vitro pre-B cell cultures (32, 36). Upon establishment of BCR-ABL1–transformed cultures after 7 to 8 days, cells were transduced with MSCV retroviral supernatants expressing IK6 and GFP or empty vector (MIG). Cells were GFP sorted prior to subsequent culture and/or transplantation into mice. All pre-B cell cultures were maintained in RPMI supplemented with 10% fetal calf serum, penicillin, streptomycin, glutamine, and 55 \( \mu \)M \( \beta \)-mercaptoethanol (BCM10). For all pre-B cell transplantation experiments, 200,000 BCR-ABL1–expressing pre-B cells were transplanted by tail vein injection into sublethally (5 Gy) irradiated 8- to 10-week-old C57BL/6 recipients. Animals were monitored daily and sacrificed when moribund or upon clinically manifest central nervous system involvement.

**Generation of xenografted mice.** A human Ph+ IK6 B-ALL sample (h9407) was provided by John Dick from the University Health Network, Toronto, Ontario, Canada (37); ALL-4 samples were generated as previously reported (38). h9407 cells were transduced for 3 hours with pCL20-MSCV-Luciferase-ires-YFP viral supernatant, washed 3 times in PBS, and transplanted by tail vein injection into unirradiated immunodeficient NSG mice. After recipients were moribund with disease, YFP+ cells were sorted and used for in vivo treatment studies. Disease progression was monitored either by flow cytometric analysis of peripheral blood cells stained for hCD45 or whole body xenogen imaging following injection of D-luciferin to detect luciferase activity.

**In vivo drug treatment studies.** Dasatinib (LC Laboratories) was administered by oral gavage at 10 mg/kg (C57BL/6), 2 mg/kg (NSG h9407), or 15 mg/kg (NOD/SCID ALL-4) in 80 mM citric acid (pH 3.1). VS-4718 in 0.5% carboxymethyl cellulose (Sigma-Aldrich) and 0.1% Tween 80 (Sigma-Aldrich) in sterile water was administered by oral gavage at 50 mg/kg. Mice were treated once daily with dasatinib (C57BL/6 and NSG h9407) or 5 days per week (NSG ALL-4).

**Calvarial imaging.** Calvaria were harvested in triplicate at 72 hours after transplant, placed in PBS, and immediately imaged using a 40× 1.1 NA LD C-Apochromat water immersion objective on a Zeiss LSM 780 microscope (Carl Zeiss Microscopy). Internal detectors were used, and the GFP signal was spectrally unmixed from the td-Tomato signal. Excitation was with a Coherent Chameleon Ti:Sapphire laser tuned to 900 nm. Three 200 × 200 micron fields of view within bone marrow cavities were analyzed per calvarium. Bone marrow cavities were identified by the absence of a second harmonic signal from the surrounding bone. For enumeration of round versus spindle-like cells, the investigator was blinded to the treatment group of calvarial images.

**RNA and protein sequencing.** Leukemic cell mRNA and protein sequencing was performed as previously described (4). In brief, RNA was extracted using TRIzol reagent and quantitated by fluorimetry (Qubit), and integrity was assessed using a TapeStation 2200 (Agilent Technologies). RNA sequencing was performed using Truseq poly-A selected library construction (Illumina) and paired end 100-bp sequencing.
using HiSeq 2000 and 2500 sequencers. Proteins were extracted from pre-B cells and digested in solution by sequential addition of Lys-C and trypsin proteases. The resulting peptides were individually desalted and labeled with 6-plex Tandem Mass Tag reagents (Thermo Scientific) for sequencing by tandem mass tag-based mass spectrometry.

Reverse-phase protein array analysis. Freshly harvested patient-derived xenografts, h9407 (IK6) and TB-08-3679 (IKZF1 wild type), were harvested and treated with 0.1 μM VS-4718 or DMSO for 24 hours at 37°C, 8% CO₂. Cell lysates were prepared according to the specifications of Theranostic Health, which performed the arrays.

In vitro apoptosis, viability, and colony-formation assays. To analyze the effects of VS-4718 on apoptosis, one million cells were seeded in triplicate wells of a 6-well dish and treated with 0, 1, or 3 μM of drug for 24 hours prior to propidium iodide staining to determine DNA content and annexin V–APC staining to assess cell viability. Cells were analyzed on a BD FACSCalibur (BD Biosciences). For cell viability assays, murine pre-B cells or human xenograft cells from freshly harvested bone marrow were plated in 96-well plates and treated with VS-4718 and dasatinib dissolved in DMSO for 48 hours at the indicated concentrations. Cells were incubated for 4 hours with resazurin and read on a Synergy HT (Biotek). For assays testing the effects of VS-4718 on colony formation, quadruplicate 96-well plates containing 0.1 μM of drug in 200 μl media/well were seeded with 1 pre-B cell per well for each of the indicated genotypes via a BD FACSARia II cell sorter (BD Biosciences). Plates were incubated at 37°C, 8% CO₂, for 7 days before colonies were counted. To test the effects of drugs on colony formation, drug was added to the media in each plate prior to sorting.

Adhesion assays. Arf⁻/⁻ BCR-ABL1 empty vector (MIG) and Arf⁻/⁻ BCR-ABL1 IK6-IRES-GFP pre-B cells were pretreated with continuous rotation for 3 hours with 1 μM VS-4718 and plated in triplicate at 6 × 10⁶ cells per well in nontissue culture treated 6-well plates coated with RetroNectin (Takara Bio). Plates were centrifuge at 450 g for 5 seconds, and cells were allowed to adhere for 30 minutes, followed by mild agitation of the plate and aspiration of unbound cells in the supernatant. For combination assays, dasatinib was added at 10 nM for the 30-minute incubation. Wells were lightly washed with PBS prior to dissociation and collection of adherent cells for counting on a Bio-Rad TC10 automated cell counter.

Immunoblotting and ELISA. Spleen fragments from control or VS-4718–treated mice (50 mg/kg) were frozen in RNALater and lysed in mPER buffer (ThermoFisher Scientific) with 3× protease/phosphatase inhibitors (Halt Protease and Phosphatase Inhibitor Cocktail). For lysates from RetroNectin assays, pre-B cells adhering to RetroNectin monolayers were lysed by addition of mPER buffer directly to the well. Western blotting was performed using anti- p-P130Cas and anti-pFAK-Y397 antibodies (Cell Signaling Technologies, 4015 and 3283, respectively). ELISA was done using the FAK [pY397] Phospho-ELISA Kit (ThermoFisher Scientific, KHO0441).

Accession numbers. RNA-sequencing data have been deposited in the Gene Expression Omnibus (accession number GSE68391).

Statistics. All graphs and Kaplan-Meier survival curves were generated and analyzed using GraphPad Prism Version 6.0 (GraphPad). All data are presented as mean ± SD. Significance was determined using 2-tailed Student’s t test, 2-way ANOVA, or Mantel-Cox log-rank test as appropriate. A P value of less than 0.05 was considered significant.

Study approval. Mice were housed in an American Association of Laboratory Animal Care–accredited facility and were treated using Institutional Animal Care and Use Committee–approved protocols in accordance with NIH guidelines.

Author contributions
MLC, IMS, JAP, DTW, RBL, PJH, MAS, and CGM designed experiments. MLC, KE, JR, AR, LJ, and IMS performed experiments. MLC and IMS analyzed data. MLC and CGM wrote the manuscript.

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