KIT Signaling Governs Differential Sensitivity of Mature and Primitive CML Progenitors to Tyrosine Kinase Inhibitors

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Abstract

Imatinib and other BCR-ABL1 inhibitors are effective therapies for chronic myelogenous leukemia (CML), but these inhibitors target additional kinases including KIT, raising the question of whether off-target effects contribute to clinical efficacy. On the basis of its involvement in CML pathogenesis, we hypothesized that KIT may govern responses of CML cells to imatinib. To test this, we assessed the growth of primary CML progenitor cells under conditions of sole BCR-ABL1, sole KIT, and dual BCR-ABL1/KIT inhibition. Sole BCR-ABL1 inhibition suppressed mature CML progenitor cells, but these effects were largely abolished by stem cell factor (SCF) and maximal suppression required dual BCR-ABL1/KIT inhibition. In contrast, KIT inhibition did not add to the effects of BCR-ABL1 inhibition in primitive progenitors, represented by CD34+38 cells. Long-term culture-initiating cell assays on murine stroma revealed profound depletion of primitive CML cells by sole BCR-ABL1 inhibition despite the presence of SCF, suggesting that primitive CML cells are unable to use SCF as a survival factor upon BCR-ABL1 inhibition. In CD34+38 cells, SCF strongly induced pAKTS473 in a phosphoinositide 3-kinase (PI3K)-dependent manner, which was further enhanced by inhibition of BCR-ABL1 and associated with increased colony survival. In contrast, pAKT3473 levels remained low in CD34+38 cells cultured under the same conditions. Consistent with reduced response to SCF, KIT surface expression was significantly lower on CD34+38 compared with CD34+38 CML cells, suggesting a possible mechanism for the differential effects of SCF on mature and primitive CML progenitor cells. Cancer Res; 73(18); 5775–86. ©2013 AACR.

Introduction

The BCR-ABL1 tyrosine kinase inhibitor (TKI), imatinib, induces profound responses in most patients with newly diagnosed chronic phase chronic myelogenous leukemia (CML-CP; ref. 1). Imatinib inhibition of BCR-ABL1 correlates with response, and reactivation of BCR-ABL1 signaling by kinase point mutations with relapse (2). In addition to BCR-ABL1, imatinib targets the tyrosine kinases ABL1, KIT, ARG (ABL2), DDR1/2, PDGFR, CSF-1R, and LCK (2–4). In contrast to BCR-ABL1, we detected no mutations in KIT in platelet-derived growth factor receptor (PDGFR) in patients with imatinib resistance (5).

Imatinib’s capacity to inhibit non-BCR-ABL1 targets has expanded its use to malignancies driven by mutations of KIT or PDGFR (6, 7), but inhibition of physiologic kinase signaling within normal cells may be the cause of side effects such as anemia (8), myelosuppression (9), and fluid retention (10). It is largely unknown whether coinhibition of non-BCR-ABL1 targets within CML cells has therapeutic benefits. KIT has been implicated in CML pathogenesis. BCR-ABL1–expressing progenitors were shown to be hypersensitive to stem cell factor (SCF) due to BCR-ABL1–induced upregulation of its receptor, KIT (11, 12) and SCF was reported to support growth of cytokine-dependent CML but not normal progenitors (13). Furthermore, culture of CML stem and progenitor cells on SCF-deficient stroma favors normal progenitors, suggesting that CML progenitors may be more SCF responsive than their normal counterparts (14). Accordingly, KIT-expressing BCR-ABL1–transduced murine myeloid cells were less sensitive to sole inhibition of either BCR-ABL1 or KIT compared with simultaneous inhibition of both kinases (15). In primary CML CD34+ cells, SCF reduced apoptosis in response to nilotinib (16), but it is unknown which specific pathways are activated by SCF to confer relative TKI resistance, and whether the requirement for KIT inhibition extends to more primitive CML cells. We sought to determine the contribution of KIT inhibition to the effects of TKIs on CML cells at various differentiation stages. We find that dual inhibition of BCR-ABL1 and KIT is required for suppression of mature but not...
primitive CML progenitors. This differential effect is due to the inability of primitive CML cells to activate AKT in response to SCF upon inhibition of BCR-ABL1.

Materials and Methods

Patient samples
Bone marrow or leukapheresis was obtained from patients with newly diagnosed CML-CP. All patients provided informed consent to research protocols approved by the Institutional Review Boards of the participating institutions. Normal bone marrow mononuclear cells (MNC) were from AllCells. Cell selection was as described previously (details in Supplementary Methods; ref. 17).

Inhibition of BCR-ABL1, KIT, MEK, and PI3K

Sole BCR-ABL1 inhibition was achieved with PPY-A (a gift of ARIAD Pharmaceuticals; ref. 18). Sole KIT inhibition was achieved by three methods: (i) use of a SCF-blocking antibody K44.2 (SCF-block; Sigma-Aldrich), a human-specific antibody that binds extracellularly to KIT and prevents SCF-induced dimerization; (ii) BAW667, a small molecule that targets KIT but not BCR-ABL1, the chemical structure of which is still proprietary. The activity profile of BAW667 was determined as previously described (19, 20) and is provided in Table S4. Requests to obtain BAW667 should be directed to Paul Manley, Novartis; (iii) downregulation of KIT using a shRNA. Mitogen-activated protein/extracellular signal-regulated kinase (MEK) inhibition was achieved with PD98059 (Cell Signaling Technology) and phosphoinositol 3-kinase (PI3K) inhibition with LY294002 (Cell Signaling Technology). For details on vector construction, see Supplementary Methods.

Immunoblot analysis of cell lines and patient samples

Analysis of primary cells and cell lines was as previously published (17, 21). For details and a list of antibodies, see Supplementary Methods.

Immunofluorescence

Fluorescence-activated cell sorting (FACS)-sorted CD34+ or CD34+38+ and CD3438+ cells were labeled with CD117/KIT-PerCP-Cy5.5, Lin-FITC, CD34-APC, and CD38-PE antibodies (BD Biosciences) and mean fluorescence intensity of CD117 in Lin-CD34+38+ and Lin-CD34+38+ cells was measured.

Quantitative real-time PCR for SCF

Details are described in Supplementary Methods.

Results

Imatinib inhibits both BCR-ABL1 and KIT, PPY-A inhibits BCR-ABL1 but not KIT, and BAW667 inhibits KIT but not BCR-ABL1

We initially determined the specificity of imatinib, dasatinib, PPY-A (18), BAW667 (a compound with activity against KIT but not BCR-ABL1), and a SCF-block. SCF-stimulated Mo7e or Mo7ep210BCR-ABL1 cells were treated with inhibitors at concentrations reported to effectively inhibit BCR-ABL1 (if applicable; refs. 18, 22) and cell lysates were immunoblotted for pKITY725 or pBCR-ABL1. SCF-block was titrated against KIT to determine an appropriate working concentration (not shown). Imatinib (2 μmol/L) and dasatinib (50 μmol/L) inhibited both BCR-ABL1 and KIT, whereas PPY-A (1 μmol/L) only inhibited pBCR-ABL1 and BAW667 (1 μmol/L) only inhibited pKITY725 (Fig. 1A). No KIT inhibition was seen with 10 μmol/L PPY-A, whereas 10 μmol/L BAW667 slightly reduced pBCR-ABL1 (Supplementary Fig. S1). We concluded that PPY-A and
BAW667 at 1 μmol/L selectively inhibit BCR-ABL1 or KIT, respectively. SCF-block at 200 ng/mL suppressed KIT phosphorylation without affecting BCR-ABL1 activity. Similar results were obtained in CD34⁺ CML cells, using CRKL as a marker for BCR-ABL1 activity (Fig. 1B). KIT was phosphorylated in CML CD34⁺ cells in the absence of SCF and this phosphorylation was reduced by imatinib or BAW667, but not PPY-A, suggesting that some KIT activation occurs without SCF, independent of BCR-ABL1 kinase activity. The band corresponding to pKIT[721] was not completely suppressed in CML CD34⁺ cells under any conditions, including imatinib and BAW667 treatment, suggesting that a kinase other than BCR-ABL1 or KIT may maintain a low level of KIT phosphorylation in primary CML cells.

Figure 1. BCR-ABL1 and KIT inhibitor profile. A, Mo7ep210[BCR-ABL1] or Mo7e cells stimulated with SCF were treated overnight with 2 μmol/L imatinib (IM), 50 nmol/L dasatinib (DA), 1 μmol/L BAW667, 1 μmol/L PPY-A, or 200 ng/mL SCF-block. Lysates were immunoblotted for phosphotyrosine or pKITY721, respectively. Total BCR-ABL1 and total KIT are shown as loading controls. Mean relative signal intensity in the presence of inhibiting agents, determined by densitometric quantitation of band intensity, is shown for n = 3 replicates. Error bars represent SEM; *, P < 0.05; **, P < 0.01; ***, P < 0.001 (Student t test). B, CD34⁺ cells from patients with newly diagnosed CML-CP (n = 3) were treated for 4 hours with 2 μmol/L imatinib, 1 μmol/L BAW667, 1 μmol/L PPY-A, or 200 ng/mL SCF-block/SCF stimulation as indicated. Lysates were immunoblotted for pCRKL or pKITY721. Total CRKL and total KIT are shown as loading controls. One representative experiment is shown. Dasatinib was not tested because of cell number limitations.

Dual inhibition of BCR-ABL1 and KIT is required for maximal suppression of CFU-GM colony formation by CML CD34⁺ cells

We initially compared colony-forming unit, granulocyte-macrophage (CFU-GM) colony formation upon sole BCR-ABL1 inhibition (PPY-A), sole KIT inhibition (SCF-block), or dual BCR-ABL1/KIT inhibition (imatinib or PPY-A+SCF-block). Cells were plated in interleukin (IL)-3, granulocyte macrophage colony-stimulating factor (GM-CSF), and SCF. Unlike imatinib, which reduced colonies by approximately 80%, PPY-A suppressed CFU-GM colony formation by only approximately 30% and SCF-block by approximately 50% (Fig. 2A). Dual BCR-ABL1 and KIT inhibition by PPY-A and SCF-block, however, reduced colony numbers by approximately 80%, suggesting that both
BCR-ABL1 and SCF/KIT contribute independently to colony growth. Normal CFU-GM colony formation was unaffected by sole BCR-ABL1 inhibition (PPY-A), but suppressed by imatinib or SCF-block, consistent with dependence on KIT signaling. The lack of efficacy of PPY-A was not due to drug instability, as pCRKL was inhibited in PPY-A–treated colonies harvested following the culture period (Fig. 2B). We also tested inhibitor effects on blast-forming unit-erythroid (BFU-E) colony formation. PPY-A had no effect, whereas sole SCF-block reduced BFU-E colony numbers to those observed with imatinib (Supplementary Fig. S2A). Thus, CML erythroid colony growth is independent of BCR-ABL1 and its suppression by imatinib is due entirely to KIT inhibition. Responses of normal BFU-E were identical, confirming that growth inhibition was cell-type rather than CML-specific. The insensitivity of CML BFU-E to PPY-A is not due to autocrine SCF production, as SCF is not expressed by CML CD34⁺ cells and not induced by PPY-A (Supplementary Fig. S2B).

In a second independent series of experiments, we included BAW667 and shKIT as alternative means of suppressing KIT activity. CD34⁺ cells from patients with CML or normal controls were plated with or without SCF and BAW667, PPY-A, BAW667+PPY-A, or imatinib were added (Fig. 3A). SCF increased CML and normal CFU-GM colonies by approximately 2.1-fold and approximately 1.7-fold, respectively. BAW667 abrogated the colony increase imparted by SCF in CML and normal cells. In addition, BAW667 reduced CML colony formation in the absence of SCF by approximately 50%, whereas effects on normal colonies were minimal, suggesting that KIT is constitutively active in CML but not normal progenitor cells and contributes to their growth. PPY-A inhibited colony formation by CML progenitor cells, and this was partially rescued by SCF, but had no effect on normal progenitor cells. Combination of PPY-A and BAW667 had effects similar to imatinib. To specifically inhibit KIT without concerns about possible off-target effects of biochemical inhibitors, we used a lentiviral vector for simultaneous expression of shKIT and GFP in human cells (Supplementary Fig. S3). Without SCF, shKIT had little effect on normal cells, but reduced colony formation of CML CD34⁺ cells by approximately 45% (Fig. 3B), similar to BAW667 alone (Fig. 3A). shKIT also abrogated the increase in colony formation caused by SCF in both normal and CML CD34⁺ cells. Finally, combining shKIT and PPY-A had similar effects as PPY-A+BAW667 or imatinib. Altogether these data show that CML CD34⁺ progenitor cells are slightly more responsive to SCF than normal CD34⁺ cells and that KIT is intrinsically active in CML but not normal cells. As a result, KIT inhibition differentiates between normal and CML CD34⁺ progenitor cells in the presence and absence of SCF, and this differential is further increased by inhibition of BCR-ABL1.

For additional validation, we analyzed CML CFU-GM colony growth following removal of the individual cytokines SCF, GM-CSF, or IL-3. We found that removal of SCF had the most pronounced effect; combining SCF removal with PPY-A had effects comparable with imatinib, suggesting that the differential sensitivity of CML CFU-GM to imatinib and PPY-A is due exclusively to their differential effects on KIT (Supplementary Fig. S4A and S4B). It should be noted that some colony growth is due to IL-3 or GM-CSF, evidenced by an approximately 15% reduction of colony growth upon removal of IL-3 or GM-CSF in the presence of PPY-A (Supplementary Fig. S4A and S4B). Removal of individual cytokines did not enhance PPY-A effects against BFU-E colony formation (Supplementary Fig. S4C), unsurprisingly given our finding that these cells are dependent on KIT but not BCR-ABL1.

**Dual inhibition of BCR-ABL1 and KIT is required to suppress CML progenitor cell growth, whereas sole BCR-ABL1 inhibition is sufficient to suppress CML stem cell growth**

Because BCR-ABL1 and KIT signaling both contribute to survival of CML progenitor cells, we determined whether this

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**Figure 2.** Suppression of CML and normal CFU-GM colony formation by PPY-A, imatinib, SCF-block, or PPY-A+SCF-block. A, CFU-GM were assessed in samples from newly diagnosed CML-CP patients (left) or normal MNCs (right) cultured for 14 days in semisolid medium containing IL-3, GM-CSF, and SCF. Imatinib, PPY-A, SCF-block, or PPY-A+SCF-block were added as indicated. Mean colony number of triplicate plates is shown normalized relative to untreated for n = 4 samples. Error bars represent SEM. B, CFU-GM colonies from untreated and PPY-A–treated plates were pooled and lysates were immunoblotted for pCRKL to assess BCR-ABL1 activity (n = 3). Total CRKL is shown as a loading control. A representative immunoblot is shown.
was also the case for more primitive CML cells, using immunophenotype or functional capacity to distinguish between mature and primitive CML progenitor cells. Lin<sup>−</sup>CD34<sup>−</sup>38<sup>−</sup> (representing relatively mature progenitor cells) and Lin<sup>−</sup>CD34<sup>−</sup>38<sup>−</sup> cells (representing a primitive population, including stem cells; ref. 17) from newly diagnosed patients with CML-CP were plated in semisolid medium supplemented with 1 μmol/L PPY-A, 1 μmol/L BAW667, or a combination thereof (Fig. 4A). KIT inhibition with BAW667 reduced colony formation by approximately 49% in progenitor cells and approximately 42% in primitive cells, respectively. In contrast, isolated BCR-ABL1 inhibition (PPY-A) had a more significant effect on primitive cells (~74% reduction) than on mature progenitor cells (~58% reduction). Combining KIT and BCR-ABL1 inhibition increased inhibition of progenitor cells by 27% to approximately 76%, whereas inhibition of primitive cells was only mildly increased by about 8% to approximately 95%.

We also plated CML progenitors on murine stroma for 1, 3, or 6 weeks in the presence of PPY-A, SCF-block or both. Although precise assignment of these populations to a specific differentiation stage is difficult, the progressively longer duration of culture permits expansion and maturation of increasingly primitive cells that are quantified by CFC assays as a composite readout for survival, expansion, and maturation (23). Resulting colonies were genotyped for BCR-ABL1 by FISH. Neither sole BCR-ABL1 (PPY-A) nor sole inhibition of KIT (SCF-block) achieved the CFC suppression seen with dual inhibition by imatinib or PPY-A + SCF-block in week 1 and 3 colonies (Fig. 4B and C). In 6-week LTC-IC, representative of primitive CML progenitors and stem cells, sole BCR-ABL1 inhibition achieved more than 95% suppression of Ph<sup>+</sup> CFC (Fig. 4D) and was comparable with imatinib (P = 0.8), whereas minimal growth suppression of 6-week LTC-IC was observed with SCF-block (Fig. 4E). Dependence on BCR-ABL1 and KIT progressively increased or decreased, respectively, with duration of growth on murine stroma (Fig. 4E). FISH revealed a mix of CML and LTC-IC, typical of early CML-CP (24). Although growth suppression was variable between patients and in some

![Figure 3. Effects of genetic or biochemical KIT inhibition on PPY-A sensitivity in normal and CML CD34<sup>+</sup> cells. A, CD34<sup>+</sup> cells from newly diagnosed patients with CML-CP (top; n = 4) or healthy controls (bottom; n = 4) were cultured for 14 days in semisolid medium containing IL-3 and GM-CSF. BAW667, PPY-A, BAW667 + PPY-A, or imatinib were added as indicated. CFU-GM colonies were scored on day 14. Untreated controls (without SCF) were set to 1. Error bars represent SEM; **, P < 0.01; ***, P < 0.001 (Student t test). B, CD34<sup>+</sup> cells from the same patients (top) and controls (bottom) as in A were infected with lentivirus for simultaneous expression of GFP and either shKIT or shSCR. Cells were plated in semisolid medium containing IL-3 and GM-CSF, PPY-A, SCF, or both were added as indicated. GFP<sup>+</sup> colonies were scored after 14 days. shSCR controls cultured with IL-3 and GM-CSF only were set to 1. Error bars represent SEM; **, P < 0.01; ***, P < 0.001 (Student t test).]
Figure 4. Suppression of mature versus primitive CML progenitors and stem cells by sole inhibition of BCR-ABL1, sole inhibition of KIT, or combined inhibition of KIT and BCR-ABL1. A, Lin−CD34−38− and Lin−CD34−38+/C0 cells were isolated from viably frozen MNCs from patients with CML-CP (n = 3) and plated in semisolid media containing IL-3 and GM-CSF with or without 1 μmol/L PPY-A, 1 μmol/L BAW667, or their combination. CFU-GM colonies were counted after 14 days. Untreated controls were set to 1. Error bars represent SEM. ***, P < 0.001. B–E, a separate series of experiments was carried out on CD34+ cells from newly diagnosed patients with CML (n = 3). Cells were cultured in LTC-IC assays on murine stromal cells (M2-10B4) in the presence of 2 μmol/L imatinib, 1 μmol/L PPY-A, 200 ng/mL SCF-block, or PPY-A+SCF-block for a total duration of 6 weeks. At 1 week (B), 3 weeks (C), and 6 weeks (D), triplicate cultures were assessed for CFC growth. Total CFC derived from 10,000 input cells are shown for each sample as well as normalized mean values for each time point. Error bars represent SEM. Differences in LTC-IC numbers were evaluated by Student t test. The frequency of Ph+ colonies was determined by FISH analysis of individual colonies from all treatment conditions. Data presented in B and C include only Ph+ colonies. E, suppression of mature and primitive CML cell outgrowth in the presence of sole BCR-ABL1 or sole KIT inhibition was compared for the three LTC-IC assays combined. F, frequency of normal (Ph−) colonies in 6-week CML LTC-IC. Mean values for the three samples are shown. Error bars represent SEM.
instances the absolute number of surviving colonies was small, Ph+ colonies were observed in all treatment conditions. Either imatinib or PPY-A, but not SCF-block increased the proportion of Ph/C0 6-week LTC-IC relative to untreated (Fig. 4F). Altogether these data suggest that primitive cells are less dependent on KIT and more dependent on BCR-ABL1 and that the capacity of KIT signaling to rescue CML cells in the presence of sole BCR-ABL1 inhibition is largely restricted to mature CML progenitors.

Differences in sensitivity to KIT inhibition may depend on KIT expression

KIT has been implicated in CML pathogenesis, but it is not precisely known how KIT and BCR-ABL1 signaling interact and whether any such interaction may depend on the differentiation stage of the cells (12). Figures 1B and 3A and B show that KIT is constitutively active in CML CD34+ cells in the absence of SCF, and that inhibition of KIT activity alone reduces colony growth, showing that KIT contributes to CML progenitor cell growth independently of SCF. To confirm this, we infected bone marrow from 5-fluorouracil (5-FU)–treated mice with retrovirus for simultaneous expression of BCR-ABL1, GFP, and shKIT or scrambled shRNA. Equal numbers of GFP-positive (GFP+) cells were plated in colony assays with or without SCF. KIT shRNA significantly reduced colony formation in the absence and presence of SCF, confirming that KIT is involved in BCR-ABL1 transformation irrespective of receptor engagement by ligand (Fig. 5A–D).

To further characterize interactions of KIT and BCR-ABL1 signaling, we used Mo7ep210<sup>BCR-ABL1</sup> cells. SCF rescued Mo7ep210<sup>BCR-ABL1</sup> cells treated with PPY-A (Fig. 6A), similar to colony assays of CML CD34+ cells (Fig. 3A). Because KIT signals through several canonical pathways including PI3K and MEK (reviewed in ref. 25), we tested how BCR-ABL1 and KIT signaling influence activation of these pathways. Mo7ep210<sup>BCR-ABL1</sup> cells were treated with SCF prior PPY-A inhibition of BCR-ABL1 signaling. In the presence of active BCR-ABL1, KIT activation of AKT (downstream of PI3K) and ERK1/2 (downstream of MEK) was weak and short-lived. In contrast, strong and sustained activation of AKT and ERK1/2 occurred when BCR-ABL1 was inhibited (Fig. 6B). Activation of AKT was associated with a reduction of Foxo3A (Supplementary Fig. S5). Cotreatment with a PI3K inhibitor (LY294002) completely and cotreatment with a MEK inhibitor (PD98059) partially inhibited SCF-mediated proliferation, suggesting that the SCF signal is transmitted mainly via PI3K/AKT and enhanced by BCR-ABL1 inhibition (Fig. 6C). To validate this finding in primary CML cells, we conducted

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**Figure 5.** Effects of KIT knockdown on colony formation by murine bone marrow cells in the presence and absence of SCF. A, murine bone marrow MNCs were infected with MIG-p185-SCR or MIG-p185-shmKIT-2 lentivirus. GFP+ cells were sorted and p185-SCR or p185-shmKIT-2 and KIT expression were detected by immunoblot. B–D, bone marrow from 5-FU–treated mice was infected with MIG-p185-SCR or MIG-p185-shmKIT-2 lentivirus and colony assays were set up in the presence or absence of 50 ng/mL rmSCF without any other cytokines. After 15 days, plates were photographed (B), GFP+ colonies were identified and counted under a fluorescence microscope (C), and colony numbers were compared between conditions (D). The experiment was carried out twice, with similar results. Error bars represent SEM. *<i>P</i> < 0.05 (Student t test).
Figure 6. Effects of SCF in combination with inhibition of BCR-ABL1, KIT, PI3K, and/or MEK on Mo7ep210^{BCR-ABL1}, Mo7e cells and primary CD34^{+} CML-CP cells. A, Mo7ep210^{BCR-ABL1} and Mo7e cells were treated with 25 ng/mL SCF, 1 \mu mol/L PPY-A, 1 \mu mol/L BAW667 or their combination(s) as indicated. Viable cell numbers were measured after 72 hours. Results at 24 hours and 48 hours were comparable (not shown). Error bars represent SEM. ***P < 0.001 (Student t test). B, total cellular lysates were harvested after 30 and 60 minutes and subjected to immunoblot analysis for pKIT^{T721}, pABL^{Y402}, pERK^{Y202/204}, pAKT^{S473}, pSTAT^{S727/729}, and \alpha-tubulin (loading control). C, Mo7ep210^{BCR-ABL1} cells were treated with PPY-A in combination with 20 \mu mol/L PD98059 (MEK inhibitor) or 20 \mu mol/L LY294002 (PI3K inhibitor) in the presence or absence of 25 ng/mL SCF. Viable cells were measured by MTS assay at 72 hours. Error bars represent SEM. *P < 0.05; ***P < 0.001 (Student t test). D and E, CD34^{+} cells from newly diagnosed patients with CML-CP (n = 3) were incubated with or without PPY-A (1 \mu mol/L) for 2 hours, followed by SCF (25 ng/mL) for 30 minutes. Aliquots of cells were analyzed for pAKT^{S473} and pERK^{Y202/204} by immunofluorescence (D) or cultured in semisolid medium using identical conditions, with CFU-GM colonies assessed after 15 days (E). Error bars represent SEM. *P < 0.05 (Student t test).
immuno

fluorescence for pAKTS473 and pERK1/2Y202/204 on CD34

þ cells treated with SCF

/C6 prior PPY-A treatment. PPY-A alone or SCF alone had moderate impact on pAKTS473 and pERK1/2Y202/204 (Fig. 6D). However, both were strongly induced by simultaneous treatment with PPY-A and SCF, confirming the results in Mo7ep210BCR-ABL1 cells (Fig. 6B).

Inhibition of PI3K (50 µmol/L LY294002) completely and inhibition of MEK (20 µmol/L PD98059) partially rescued the SCF effect (Fig. 6E), validating the data on Mo7ep210BCR-ABL1 cells (Fig. 6C). Altogether these results show that SCF rescues CML CD34

þ progenitor cells upon inhibition of BCR-ABL1, mainly via PI3K/AKT, which explains why dual inhibition of KIT and BCR-ABL1 is required to effectively target these cells.

We next examined possible differences in signaling response to SCF between mature (CD34+38+) and primitive (CD34+38−) progenitor cells. CD34+ cells were cultured overnight in B1 medium without cytokines and sorted, treated with SCF, PPY-A, or both, and then analyzed by immunofluorescence for pAKTS473 (Fig. 7A). pAKTS473 was lower in CD34+38− cells than CD34+38+ cells, as previously reported (17). PPY-A alone minimally reduced pAKTS473 in CD34+38+ cells. Strikingly, SCF strongly induced pAKTS473 in PPY-A–treated CD34+38+ cells, analogous to Mo7ep210BCR-ABL1 cells (Fig. 6B), but had little effect in CD34+38− cells, suggesting that upon BCR-ABL1 inhibition CD34+38− CML cells fail to launch a robust pAKTS473 response to SCF stimulation. To identify the underlying mechanism, we measured CD117 (KIT) expression on Lin−/C0 CD34+38+ versus Lin−/C0 CD34+38− cells and found significantly lower expression in the primitive Lin− CD34+38− cells, which may explain their decreased response to SCF and greater vulnerability to sole BCR-ABL1 inhibition (Fig. 7B).

Discussion

Previous studies have implicated KIT in CML pathogenesis, suggesting that the efficacy of TKIs such as imatinib may be
due to their combined activity against BCR-ABL1 and KIT (13, 15, 26). For example, KIT– BCR-ABL1–transduced murine progenitor cells are more sensitive to dual inhibition of KIT and BCR-ABL1 than to inhibition of either kinase individually (15). In another study, SCF rescued CML CD34+ cells from nilotinib but not imatinib effects and ascribed the difference to nilotinib’s relatively weaker anti-KIT activity (16). However, it remained unknown which specific pathways are activated by SCF to confer relative TKI resistance and whether the requirement for KIT inhibition depends on the stage of differentiation. Here, we use TKIs, shRNA and blocking antibodies to examine whether the complimentary activity of imatinib against both BCR-ABL1 and KIT contributes to its efficacy in mature and primitive primary CML cells.

Sole BCR-ABL1 inhibition with PPY-A modestly suppressed CML CFU-GM colony formation in the presence of cytokines, whereas SCF-block caused a slightly more pronounced reduction (Fig. 2A). This reduction was not exclusively dependent on active SCF signaling, as SCF-block (Fig. 2A) or removal of SCF (Supplementary Fig. S4A) had quantitatively similar effects to BAW667 (Fig. 3A) or shKIT knockdown in SCF-free cultures (Fig. 3B). These data show that KIT contributes to the proliferation of mature CML progenitors in the absence of ligand, in accord with previous reports in BCR-ABL1–expressing cell lines (12). Consistent with this, we detected low levels of pKIT Y722 in serum-starved CML CD34+ cells that was reduced by imatinib and BAW667 (Fig. 1B). The effects of BAW667 inhibition of KIT were maximal in cultures of CML CD34+ cells that were supplemented with SCF (Fig. 3A, left: compare dark bars in control vs. BAW667). Expression of BCR-ABL1 with or without simultaneous KIT knockdown in murine bone marrow (Fig. 5) reproduced the data on primary human CML cells, indicating that both SCF-induced and -independent KIT activation contribute to CML progenitor cell growth. Surprisingly, shKIT significantly enhanced the effects of PPY-A on CML CD34+ colony formation in the absence of SCF (Fig. 3B, top). This could reflect persistence of a low level of BCR-ABL1 kinase activity not detected by pCRKL immunobLOTS (27) or constitutive KIT activation that is independent of BCR-ABL1 kinase activity. In addition, the band corresponding to pKIt Y722 was not completely abolished by BAW667 or imatinib (Fig. 1B), suggesting that a kinase other than BCR-ABL1 or KIT may phosphorylate KIT on tyrosine 722, although this residue is generally regarded as an autophosphorylation site. For example, SRC family kinases have been shown to phosphorylate tyrosine 900 of KIT (28). Additional studies will be required to distinguish between these two possibilities.

In contrast to the limited effects of targeting either BCR-ABL1 or KIT in isolation, simultaneous inhibition of both kinases dramatically reduced CML CFU-GM growth. Disruption of KIT signaling was achieved with four independent approaches (BAW667, SCF-block, SCF removal, and KIT knockdown), all of which produced similar effects when combined with the BCR-ABL1 inhibitor PPY-A. Dual dependence of primary CML progenitor cells on BCR-ABL1 and KIT signaling is restricted to granulocyte precursors. Although erythroid progenitors express both BCR-ABL1 and KIT (29), erythroid colony formation was maximally suppressed by inhibition of KIT alone and independent of BCR-ABL1 activity, identical to normal BFU-E (Supplementary Fig. S2A). Thus, imatinib suppression of leukemic BFU-E is due entirely to KIT inhibition and BCR-ABL1 expression in erythroid lineage cells is not synonymous with dependence on BCR-ABL1 (30). Accordingly, erythrocytosis is not a feature of CML.

Unlike the balanced contribution of BCR-ABL1 and KIT inhibition to suppression of CFU-GM colonies, effects on primitive CML cells, defined either by a CD34+ 38− phenotype (Fig. 4A) or LTC-IC functionality (Fig. 4B) were mostly due to BCR-ABL1 inhibition. In particular, in 6-week LTC-IC assays, which select primitive CML progenitor cells (23), both imatinib and PPY-A reduced Ph− LTC-IC colonies by more than 95%, consistent with an effect that requires inhibition of BCR-ABL1, but not KIT. On the surface, the capacity of sole BCR-ABL1 inhibition to suppress primitive CML cells seems to contradict reports by us and others that that CML stem cells are insensitive to BCR-ABL1 inhibitors (17, 31). Furthermore, previous studies reported only modest imatinib effects on CML LTC-IC (32, 33). The differences are readily explained by the fact that prior studies evaluated the effects of short-term (72–96 hours) drug treatment of CML progenitors followed by 6-week culture on stroma without TKIs. These assays show the inability of TKIs to effectively induce apoptosis in primitive cells, but do not reflect conditions of long-term imatinib treatment. In contrast, we examined how continuous suppression of BCR-ABL1, KIT, or their combination throughout the 6-week culture period would affect LTC-IC outgrowth. Importantly, to generate an environment devoid of human cytokines, we conducted the LTC-IC assays using unmanipulated murine (M210B4) stromal cells (i.e., not engineered to express human cytokines). Because most cytokines and chemokines are not cross-reactive between species (34), these conditions minimize extrinsic factors that might support CML stem cells despite BCR-ABL1 inhibition. In these conditions, imatinib and PPY-A resulted in profound suppression of the most primitive cells. Notably, the differential effects of sole BCR-ABL1 versus sole KIT inhibition on mature versus primitive CML progenitor cells were consistent irrespective of whether the cell populations were defined by immunophenotype (Fig. 4A) or functionality (Fig. 4B–F). Given the overall profound effect of sole BCR-ABL1 inhibition on primitive CML progenitor cells, it is impossible to exclude a small contribution of KIT inhibition to the suppression of this population. Despite small numbers of colonies, in all samples Ph+ LTC-IC survived in the presence of BCR-ABL1 inhibitors, consistent with reports of residual BCR-ABL1 LTC-IC and CD34− 38− cells in patients with sustained molecular response to imatinib (35, 36).

The differential sensitivity of mature and primitive CML progenitors to sole BCR-ABL1 or KIT inhibition suggested cell type–specific differences in the response to SCF. We initially studied Mo7ep210 (CR-ABL1) cells and found that SCF rescued these cells from the effects of PPY-A inhibition of BCR-ABL1 (Fig. 6A). Although active BCR-ABL1 blunted SCF activation of AKT and MEK, key pathways downstream of KIT (25), inhibition of BCR-ABL1 sensitized cells to SCF. SCF rescue was completely blocked by PI3K inhibition, but only partially by MEK inhibition, implicating PI3K/AKT as
the critical pathway downstream of KIT (Fig. 6C). Similar results were obtained in primary CML CD34+ cells whereas PPY-A or SCF alone had little effect on pAKT$^{S473}$, their combination greatly increased pAKT$^{S473}$ (Fig. 6D). LY294002 abrogated the SCF-induced increase in colony formation (Fig. 6E). In contrast, basal pAKT$^{S473}$ in CD34+38 cells was low, as previously reported in BCR-ABL1-expressing murine stem cells and CD34+38 CML cells (17, 37), unchanged upon PPY-A treatment and minimally increased by SCF alone or in combination with PPY-A (Fig. 7A). This is in stark contrast to CD34+38 CML cells (Fig. 6D) and suggests that the PPY-A sensitivity of CD34+38 CML cells is due to their inability to strongly activate SCF signaling upon BCR-ABL1 inhibition, possibly reflecting the lower CD117 expression on CD34+38 compared with CD34+38 CML cells (Fig. 7B). As the most primitive CML cells are not strongly KIT dependent, this may explain why no KIT mutations have been observed in patients with imatinib resistance (5). Additional mechanisms may be involved in blunting the SCF response of primitive CML progenitor cells in vivo. For example, Naka and colleagues reported that TGF-β blocks BCR-ABL1-induced AKT activation in BCR-ABL1-transduced murine stem cells (37). Because TGF-β is reported to prevent SCF rescue of mast cells after IL-3 withdrawal (38), it is possible that TGF-β in the microenvironment may further reduce SCF-induced rescue of CD34+38 CML cells.

From the drug development perspective, it is important to consider whether there is benefit to inhibiting nononcogenic targets, or whether “surgical” inhibitors with the narrowest possible target spectrum are preferred. Among the second- and third-generation BCR-ABL1 inhibitors, dasatinib, nilotinib, and ponatinib directly inhibit KIT, whereas bosutinib has no activity against KIT (39). The clinical activity of bosutinib (40, 41) can be explained by its inhibitory activity toward SRC kinases, which play a critical role in KIT signaling (42, 43). Altogether our data suggest that, in vivo, CML stem cells may survive BCR-ABL1 inhibition through a pathway other than SCF/KIT that is not activated by M210B4 stromal cells. Therefore, the TKI resistance of primitive CML cells may simply reflect the fact that these inhibitors fail to target a second critical pathway, in contrast to the fortuitous situation in mature progenitor cells. Identifying pathways that support CML stem cells in the presence of BCR-ABL1 TKIs should open new therapeutic options.

Disclosure of Potential Conflicts of Interest
P.W. Manley is employed as Research Investigator in Novartis Pharma. J.E. Cortes has commercial research grant from Bristol-Myers Squibb, Novartis, Pfizer, ARIAD Pharmaceuticals, and Teva and is a consultant/advisory board member of ARIAD Pharmaceuticals, Pfizer, and Teva. B.J. Drucker has commercial research grant from Novartis, Bristol-Myers Squibb, and ARIAD Pharmaceuticals. M.W. Deininger has commercial research grant from Novartis and Bristol-Myers Squibb and is a consultant/advisory board member of Bristol-Myers Squibb, Novartis, ARIAD Pharmaceuticals, and Incyte. No potential conflicts of interest were disclosed by the other authors.

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References
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