Expression of a Src Family Kinase in Chronic Myelogenous Leukemia Cells Induces Resistance to Imatinib in a Kinase-dependent Manner

Received for publication, November 28, 2009, and in revised form, May 6, 2010 Published, JBC Papers in Press, May 7, 2010, DOI 10.1074/jbc.M109.090043

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The Bcr-Abl kinase inhibitor imatinib is remarkably effective in chronic myelogenous leukemia (CML), although drug resistance is an emerging problem. Myeloid Src family kinases such as Hck and Lyn are often overexpressed in imatinib-resistant CML cells that lack Bcr-Abl mutations. Here we tested whether Hck overexpression is sufficient to induce imatinib resistance using both wild-type Hck and a mutant (Hck-T338A) that is uniquely sensitive to the pyrazolopyrimidine inhibitor, NaPP1. Expression of either kinase in K562 CML cells caused resistance to imatinib-induced apoptosis and inhibition of soft-agar colony formation. Treatment with NaPP1 restored sensitivity to imatinib in cells expressing T338A but not wild-type Hck, demonstrating that resistance requires Hck kinase activity. NaPP1 also reduced Hck-mediated phosphorylation of Bcr-Abl at sites that may affect imatinib sensitivity exclusively in cells expressing Hck-T338A. These data show that elevated Src family kinase activity is sufficient to induce imatinib resistance through a mechanism that may involve phosphorylation of Bcr-Abl.

Chronic myelogenous leukemia (CML) is a hematopoietic stem cell disease with three clinical phases. The initial chronic phase involves expansion of myeloid cells that retain their ability to undergo terminal differentiation. As the disease progresses, patients enter an accelerated phase followed by a blast crisis characterized by differentiation arrest and accumulation of immature blast cells in the bone marrow and peripheral blood (1).

The cytogenetic hallmark of CML is the Philadelphia chromosome, which fuses the BCR (breakpoint cluster region) locus on chromosome 22 with the c-ABL proto-oncogene on chromosome 9. This translocation is present in greater than 90% of CML patients and leads to the expression of Bcr-Abl, a chimeric and constitutive protein-tyrosine kinase activity (4, 5). Bcr-Abl drives the pathogenesis of CML through the phosphorylation and activation of a broad range of downstream signaling proteins that increase cell survival and promote unregulated cell cycle progression (6). These pathways include but are not limited to the Ras/mitogen-activated protein kinase (MAPK), NF-κB, phosphatidylinositol 3-kinase/Akt, and Stat signaling cascades (7–9).

Bcr-Abl has been shown to activate other non-receptor protein-tyrosine kinases, particularly Src family kinases (SFKs) expressed in myeloid cells such as Hck and Lyn (10). A growing body of evidence supports the relevance of this interaction to CML pathogenesis. For example, expression of a kinase-defective mutant of Hck blocked Bcr-Abl-induced transformation of murine myeloid leukemia cells to cytokine independence (11). In addition, Hck was shown to couple Bcr-Abl to Stat5 signaling and to be required for Bcr-Abl-induced transformation of murine myeloid cells (12). Furthermore, SFK-selective inhibitors block proliferation and induce apoptosis in CML cells without affecting Philadelphia chromosome-negative myeloid leukemia cells (13). More recently, Hck has been shown to play a non-redundant role in Bcr-Abl survival signaling in CML cells (14). Taken together, these findings indicate that SFKs act as essential mediators of Bcr-Abl-induced leukemogenesis.

Because Bcr-Abl plays a critical role in the initiation and maintenance of the CML phenotype, targeting its tyrosine kinase activity is the therapeutic strategy of choice. Imatinib mesylate is a potent small molecule tyrosine kinase inhibitor relatively specific to Bcr-Abl that has become the frontline therapy for patients with CML (6, 15). Most CML patients in the chronic phase achieve and maintain significant hematologic and cytogenetic responses to imatinib treatment (16, 17). However, ~4% of chronic phase patients treated with imatinib develop drug resistance per year. In addition, accelerated or blast crisis patients display higher rates of imatinib resistance (18, 19). Clinical resistance to imatinib can result from mutations in the Abl kinase domain at residues that directly contact imatinib or at positions that allosterically influence imatinib binding (20). Resistance can also arise in the absence of Bcr-Abl mutations. The present study is focused on the role of SFKs in Bcr-Abl mutation-independent imatinib resistance.

Overexpression of the myeloid SFKs Hck and Lyn has been associated with resistance to imatinib in the absence of Bcr-Abl kinase domain mutations. Selection of the CML cell line K562 for resistance to high levels of imatinib increased Lyn protein

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EXPERIMENTAL PROCEDURES

Cell Culture—Sf9 insect cells were cultured in Grace’s medium (Invitrogen) supplemented with 10% fetal bovine serum (FBS) and 50 μg/ml gentamicin (Invitrogen). The K562 myeloid leukemia cell line, derived from a CML patient in erythroid blast crisis (28), was obtained from the ATCC and maintained in RPMI 1640 supplemented with 10% FBS (Atlanta Biologicals) and 0.25 μg/ml Antibiotic-antimycotic (Invitrogen). Rat-2 fibroblasts were purchased from the ATCC and maintained in Dulbecco’s modified Eagle’s medium containing 5% FBS, 100 units/ml penicillin G, 100 μg/ml streptomycin, and 0.25 μg/ml Antibiotic-antimycotic.

Protein Purification—Using site-directed mutagenesis, the Thr-338 to Ala (T338A) mutation was introduced into the coding sequence of human p59 Hck-YEEI (QuikChange XL site-directed mutagenesis kit, Stratagene) (29). The Hck-YEEI and Hck-T338A-YEEI constructs also contained N-terminal hexahistidine tags in place of the unique domain (30, 31). The Ncap-c-Abl construct encompassed residues 1–531 of human c-Abl-1b with residues 15–56 deleted and contains a C-terminal cleavage site for the tobacco etch virus (TEV) protease and a hexahistidine tag, both introduced by PCR (32, 33). Hck-YEEI, Hck-T338A-YEEI, and the Ncap-c-Abl constructs were cloned into pVL1392 (BD Biosciences), and each plasmid was used to create high titer recombinant baculovirus in Sf9 insect cells using BaculoGold DNA and the manufacturer’s protocol (BD Biosciences). For protein production, Sf9 cells (1 liter) were cultured to a density of 2 × 10^6 cells/ml and then infected with either the Hck-YEEI or Hck-T338A-YEEI baculoviruses. For the Ncap-c-Abl construct, Sf9 cells were co-infected with Abl and YopH phosphatase baculoviruses at a multiplicity of infection of 10. YopH is a protein-tyrosine phosphatase that promotes a down-regulated conformation of Ncap-c-Abl, permitting high yield purification from Sf9 cells (32). Sf9 cells were grown for 48 h, centrifuged, washed in PBS, and then the pellets were resuspended in buffer A (20 mM Tris-HCl (pH 8.3), 10% glycerol, and 5 mM 2-mercaptoethanol), lyzed by sonication, and centrifuged at 16,000 rpm for 30 min. The recombinant Hck or Ncap-c-Abl proteins were purified from the supernatant using a combination of ion exchange and affinity chromatography as originally described by Schindler et al. for Hck-YEEI (30). Upon purification, the proteins were dialyzed against 20 mM Tris-HCl (pH 8.3) containing 100 mM NaCl and 3 mM dithiothreitol.

In Vitro Kinase Assay—Kinase assays were performed using the fluorescence resonance energy transfer-based Z’-Lyte Src kinase assay kit and Tyr-2 peptide substrate according to the manufacturer’s instructions (Invitrogen). All assays were performed in quadruplicate in low volume, non-binding 384-well plates (Corning). The assay was first optimized to determine the amount of each kinase and the incubation time necessary to phosphorylate 50–60% of the Tyr-2 peptide in the absence of inhibitor. Final ATP and Tyr-2 substrate concentrations were held constant at 50 and 2 μM, respectively. For inhibition experiments, the kinases were preincubated with NaPP1 or imatinib in kinase assay buffer for 30 min followed by incubation with ATP and Tyr-2 peptide for 1 h. Fluorescence was assessed on and activity levels (21). Exposure of these cells to imatinib resulted in an incomplete suppression of Bcr-Abl activity and was accompanied by persistent tyrosine phosphorylation of Bcr-Abl at Tyr-177, a binding site for Grb2 that links Bcr-Abl to the Ras signaling cascade (22, 23). Samples from patients after imatinib failure exhibited increased Hck and/or Lyn activity levels that were not affected by Bcr-Abl inhibition (21). In addition, no Bcr-Abl kinase domain mutations associated with imatinib resistance was present in these patients. However, small interfering RNA-mediated suppression of Lyn expression or treatment with dasatinib, a dual Abl/SFK inhibitor, induced apoptosis in these imatinib-resistant cells (24). Collectively, these findings point to a role for both Hck and Lyn in imatinib resistance in the absence of Bcr-Abl mutations. Although the role of Lyn has been pursued in some detail, the contribution of Hck to this type of imatinib resistance is less well understood.

To address this question, we developed a system in which Hck activity can be inhibited in a highly specific manner. Typically, small molecule kinase inhibitors exhibit limited specificity, especially for individual members of a conserved kinase family. To circumvent this issue, we employed a chemical genetic approach originally described by Shokat and co-workers (for review, see Ref. 25). In this method, pyrazolopyrimidine 1 (PP1), a relatively potent yet non-selective SFK inhibitor, was modified to obtain a more bulky analog, naphthyl-pyrazolopyrimidine 1 (NaPP1), that is far less potent toward wild-type SFKs than the parent compound (Fig. 1A). Using site-directed mutagenesis, the ATP binding sites of SFKs were then modified to accommodate the bulkier naphthyl moiety of the NaPP1 molecule (25, 26). The resulting combination of inhibitor analog and mutant kinase allows for highly selective inhibition of the modified kinase without affecting the corresponding endogenous kinase or related kinases.

In this study we used the chemical genetic principle described above to successfully engineer a Hck mutant (Hck-T338A) that is uniquely sensitive to NaPP1 inhibition. The validity of the mutant was confirmed in vitro using recombinant purified kinases as well as a Rat-2 fibroblast transformation assay. We then used this Hck mutant to address the role of Hck overexpression in imatinib resistance. Wild-type Hck and the T338A mutant were expressed in the CML cell line K562, which was derived from an erythroid blast crisis patient and does not express endogenous Hck (14). Remarkably, expression of either wild-type Hck or the T338A mutant rendered K562 cells resistant to imatinib-induced inhibition of colony formation and apoptosis. This effect correlated with sustained phosphorylation of Bcr-Abl at several regulatory tyrosine residues in the presence of imatinib. These findings suggest that Hck overexpression promotes Bcr-Abl activation, which correlates inversely with inhibition by imatinib (27). Furthermore, NaPP1 treatment restored imatinib sensitivity to cells expressing the Hck-T338A mutant but not the wild-type kinase. This effect correlated with inhibition of Hck-T338A activity and reduction of site-specific Bcr-Abl tyrosine phosphorylation. In contrast, NaPP1 did not affect cells expressing wild-type Hck or control cells. Taken together, these results establish the first causal relationship between Hck kinase activity and Bcr-Abl mutation-independent imatinib resistance.
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...a Gemini XS microplate spectrofluorometer (Molecular Devices). IC₅₀ values were calculated from the means of four wells using a sigmoidal curve fit and Prism software (GraphPad Software, Inc.).

Retroviral Transduction of Rat-2 Fibroblasts and K562 Cells—Using site-directed mutagenesis, an active form of Hck was constructed by replacing the negative regulatory tail tyrosine residue with phenylalanine (Hck-YF) (34). The wild-type Hck, Hck-T338A, Hck-YF, and Hck-T338A-YF constructs were subcloned into the retroviral vector pSRoMSVtkneo (35) and used to generate high titer retroviral stocks by co-transfection of 293T cells with an ecotropic packaging vector (29, 34, 36, 37). Control retroviruses were prepared using the empty parent vector. Rat-2 fibroblasts were infected as follows: 2.5 × 10⁴ Rat-2 cells were plated per well in 6-well plates and incubated with viral stocks in a final volume of 5 ml in the presence of Polybrene (4 μg/ml final). To enhance infection efficiency, the plates were centrifuged at 3000 rpm for 4 h at 18 °C. After infection, the virus was replaced with fresh medium. G418 selection (800 μg/ml) was started 48 h after infection and continued for 14 days. At the end of the selection period, the G418 concentration was decreased to 400 μg/ml.

For K562 cells, wild-type Hck and the Hck-T338A mutant were subcloned into the retroviral expression vector pMSCV-IRES-neo (Clontech) between the MSCV promoter and internal ribosome entry site-neo sequence. Retroviral stocks were produced from the resulting constructs in 293T cells using an amphotropic packaging vector as described above for Rat-2 fibroblasts. K562 cells were plated in 6-well plates at 1 × 10⁶ cells/well in 5 ml of undiluted viral supernatant in the presence of 4 μg/ml Polybrene and centrifuged at 3000 rpm for 3 h at room temperature. After infection, cells were washed, returned to regular medium for 48 h, and then placed under G418 selection (800 μg/ml) for 14 days. At the end of the selection period, cells were maintained in medium with 400 μg/ml G418.

Soft Agar Colony Assay—Rat-2 fibroblast or K562 cell colony-forming activity was assayed in triplicate 35-mm Petri dishes (Falcon) using Seaplaque-agarose (FMC Bioproducts). One ml of 0.5% bottom agarose layer in complete culture medium was poured in the presence of either vehicle (0.5% DMSO) or NaPP1/imatinib at twice the final desired concentration. After hardening of the bottom layer, 1 × 10⁴ Rat-2 cells or 2 × 10⁵ K562 cells were mixed in culture medium containing 0.3% agarose, and 1 ml was added to the plates. 7–10 days later, the colonies were stained with 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide and counted from scanned images using colony counting software (Bio-Rad QuantityOne).

Apoptosis Assay—Apoptosis was determined by measuring cell-surface phosphatidylserine using an Alexa Fluor 488-conjugated anti-phosphatidylserine antibody (Upstate Biotechnology) and flow cytometry. Cells (1 × 10⁷/ml) were treated with vehicle alone (0.5% DMSO), imatinib, or combinations of NaPP1 and imatinib for 72 h at 37 °C. After incubation, cells were centrifuged at 1000 rpm for 5 min, washed 3 times with ice-cold PBS, and resuspended to 4 × 10⁶ cells/ml in staining buffer (1% PBS in PBS). Aliquots (50 μl) were transferred to 96-well round-bottom tissue culture plates, mixed with the anti-phosphatidylserine antibody (0.21 μg/well), and incubated on ice for 1 h. Cells were washed three times in ice-cold PBS and analyzed using a FACSCalibur flow cytometer (BD Biosciences) and CellQuest software.

Antibodies—The following antibodies were used in this study: anti-actin (MAB1501; Chemicon), anti-Hck (N-30 polyclonal, Santa Cruz Biotechnology; monoclonal, Transduction Laboratories), anti-Lyn (Santa Cruz), anti-c-Abl (Calbiochem), anti-Src phosphospecific (Src phospho-Tyr-416; Upstate Biotechnology), anti-Bcr phospho-Tyr-177 (Cell Signaling), anti-c-Abl phospho-Tyr-89 (Cell Signaling), anti-c-Abl phospho-Tyr-245 (Cell Signaling), and anti-phosphotyrosine (PY99; Santa Cruz).

Immunoprecipitation and Immunoblotting—K562 cells (5 × 10⁶) were incubated overnight in 0.5% FBS and then treated for 5 h with vehicle control (DMSO, 0.5%) or with imatinib and/or NaPP1. At the end of the incubation period, the cells were collected by centrifugation, washed twice with PBS, and lysed in ice-cold radioimmune precipitation assay buffer (34). For Hck or Lyn immunoprecipitation, total protein concentrations were first normalized with lysis buffer followed by the addition of 1 μg of anti-Hck or anti-Lyn antibody and 25 μl of protein G-Sepharose (50% slurry; GE Healthcare). After incubation for 2 h at 4 °C, immunoprecipitates were washed 3 times with 1.0 ml of radioimmune precipitation assay buffer and heated in SDS sample buffer. After SDS-PAGE, proteins were transferred to polyvinylidene difluoride or nitrocellulose membranes and incubated with the indicated primary antibodies. Immunoreactive proteins were visualized with appropriate secondary antibody-alkaline phosphatase conjugates and nitro blue tetrazolium/5-bromo-4-chloro-3-indolyl phosphate colorimetric substrate (Sigma). For quantitative immunoblotting, the membranes were incubated with IRdye680- or IRdye800CW-conjugated secondary antibodies (LI-COR, Lincoln, NE) and scanned with the Odyssey infrared imaging system (LI-COR). Protein band intensities were quantitated using the Odyssey software.

Statistical Analysis—Data obtained from multiple independent experiments are reported as the mean ± S.D. values. Statistical tests were conducted using Student’s t test or analysis of variance (ANOVA) where indicated. Statistical tests were performed using GraphPad Prism, Version 3.03.

RESULTS

Design of an Inhibitor Analog-sensitive Hck Allele—Crystal structures of various protein-tyrosine kinases have shown that a small hydrophobic pocket lies adjacent to the ATP binding site, access to which is controlled by a “gatekeeper residue.” The side chain of the gatekeeper (typically a bulky residue such as threonine, methionine, isoleucine, or phenylalanine) sterically hinders the access of many small molecule inhibitors to this pocket (38, 39). Based on these observations, several groups have successfully applied a kinase inhibitor sensitization approach to a number of protein kinase families by replacing the gatekeeper residue with a much smaller residue such as glycine or alanine (25, 40, 41). In particular, replacement of the v-Src gatekeeper (Ile-338; numbering is based on the crystal structure of human c-Src (42)) with either glycine or alanine rendered it susceptible to inhibition by NaPP1 (43), suggesting that this
to adopt the down-regulated conformation, enabling high yield purification from Sf9 cells without the need for Csk co-expression (30, 31, 45). NaPP1 sensitivity of recombinant purified Hck-YEEI and Hck-T338A-YEEI protein kinases was then assayed using an in vitro kinase assay (Z’-Lyte) and a fluorescence resonance energy transfer-peptide substrate (a detailed explanation of the Z’-Lyte assay and the calculation of percent inhibition is provided in supplemental Fig. S1.) As shown in Fig. 1C, NaPP1 potently inhibited the Hck-T338A-YEEI mutant with an IC$_{50}$ of 26.6 ± 1.26 nM. In contrast, the NaPP1 IC$_{50}$ for Hck-YEEI was almost 50-fold higher (1.3 ± 0.39 μM). Importantly, NaPP1 did not inhibit a recombinant purified form of the c-Abl tyrosine kinase core consisting of the N-terminal cap (Ncap) region, the SH3 domain, the SH2 domain, and the kinase domain (referred to hereafter as “Ncap-c-Abl”) (33).

One possible issue concerning the introduction of the gatekeeper mutation in the ATP binding site of Hck is cross-sensitization to imatinib. To rule out this possibility, we also tested Hck-T338A sensitivity to imatinib in the in vitro kinase assay using the same set of purified recombinant kinases. As shown in Fig. 1D, imatinib did not inhibit wild-type Hck or the T338A mutant, further supporting the validity of this approach. In contrast, imatinib inhibited Ncap-c-Abl with an IC$_{50}$ of 350 ± 53 nM, consistent with previous work (46).

**NaPP1 Potently and Selectively Inhibits Hck-T338A in a Fibroblast Transformation Assay**—To determine whether the Hck-T338A mutant is sensitive to NaPP1 in a cell-based assay, we next tested the ability of NaPP1 to revert the transformed phenotype of fibroblasts expressing an active form of Hck-T338A. For these experiments, we replaced the C-terminal negative regulatory tail tyrosine residue (Tyr-527) with phenylalanine (“YF” mutation) in both wild-type Hck and the T338A mutant. This mutation was previously shown to prevent tail Tyr-527 phosphorylation by Csk and SH2 domain engagement, resulting in a marked increase in kinase activity and oncogenic transformation of fibroblasts (29, 34, 36, 37, 48). The resulting Hck-YF and Hck-T338A-YF mutants were expressed in Rat-2 fibroblasts, and their transforming potential and sensitivity to NaPP1 were compared in a soft-agar colony formation assay. Fibroblasts expressing wild-type Hck, Hck-T338A, or the G418 selection marker were used as negative controls. As shown in
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Fig. 2A, neither wild-type Hck nor the T338A mutant induced fibroblast transformation, demonstrating that alanine substitution at the gatekeeper position does not result in kinase activation. In contrast, both Hck-YF and Hck-T338A-YF had robust and comparable transforming activity, demonstrating that the T338A mutation does not interfere with the biological functions of the kinase in this system. Hck activity in each of the Rat-2 cell lines was then assessed by anti-phosphotyrosine immunoblotting, which reveals the endogenous Hck substrate protein, pp40 (Fig. 2B). Both Hck-YF and Hck-T338A-YF showed strong constitutive activity in this assay. In contrast, wild-type Hck and Hck-T338A showed very little kinase activity, consistent with their lack of transforming function.

We next assessed the specificity of NaPP1 for Hck-T338A-YF in this system. As shown in Fig. 2A, NaPP1 signifi- cantly blocked colony formation by cells transformed with Hck-T338A-YF with an IC_{50} of ~0.1 μM. This effect correlated with inhibition of Hck-T338A-YF activity as assessed by anti-phosphotyrosine immunoblotting, which reveals the endogenous Hck substrate protein, pp40 (Fig. 2B). Conversely, neither colony formation nor kinase activity was affected by NaPP1 in cells transformed by Hck-YF, even at concentrations as high as 1 μM. These results establish that the T338A mutation is functionally silent yet confers NaPP1 sensitivity on Hck in a cell-based assay.

Overexpression of Hck and Hck-T338A Protects K562 CML Cells from Imatinib-induced Apoptosis and Inhibition of Colony Formation—Previous studies have shown that a subset of imatinib-resistant blast crisis CML patients display increased Hck kinase protein expression and/or activity in the absence Bcr-Abl mutations (21, 22, 24). To determine whether Hck overexpression in CML cells is sufficient to induce imatinib resistance, wild-type Hck and Hck-T338A were expressed in K562 cells using recombinant retroviruses. Cells infected with a retrovirus carrying only the neomycin resistance marker served as a negative control. After selection with G418, the K562-Neo, K562-Hck, and K562-Hck-T338A cell populations were treated over a range of imatinib concentrations for 72 h. The fraction of apoptotic cells was then determined by anti-phosphatidylinerine antibody staining and flow cytometry. As shown in Fig. 3, imatinib induced apoptosis in a dose-dependent manner in the K562-neo cell population, with apoptotic cells evident with as little as 0.3 μM imatinib. Strikingly, overexpression of wild-type Hck or Hck-T338A in K562 cells was sufficient to reduce the apoptotic effects of imatinib at both 0.3 and 1 μM (p < 0.05, one-way ANOVA).

We next investigated whether expression of Hck in K562 cells reverses imatinib-induced inhibition of colony formation. To address this issue, all three K562 cell populations were plated in soft agar over a range of imatinib concentrations. As shown in Fig. 4, K562 cells expressing wild-type Hck or Hck-T338A yielded a comparable number of colonies as the vector control cells when plated in soft-agar in the absence of imatinib. This finding suggests that expression of wild-type Hck or Hck-T338A alone does not enhance the basal level of K562 cell colony-forming activity. Imatinib induced a dose-dependent inhibition of colony formation in all three cell lines. However, cells expressing wild-type Hck or Hck-T338A displayed substantial resistance to imatinib-induced inhibition of colony formation when compared with control cells. More specifically, although 0.03 μM imatinib reduced the number of K562-neo colonies by nearly 50%, it did not have a significant impact on cells expressing either form of Hck (p = 0.0003, one-way ANOVA). Furthermore, K562-Hck and K562-Hck-T338A cells were three times more resistant to 0.1 μM imatinib than K562-neo control cells in this assay (p < 0.0001, one-way ANOVA). Taken together, results from both the apoptosis and colony assays demonstrate that overexpression of Hck is sufficient to induce resistance to imatinib.

Resistance to Imatinib-induced Apoptosis and Inhibition of Colony Growth Requires Hck Kinase Activity—To determine whether resistance to imatinib requires Hck kinase activity, we tested whether specific inhibition of Hck-T338A with NaPP1
Expression of wild-type Hck or Hck-T338A protects K562 cells from imatinib-induced apoptosis. K562-Neo, K562-Hck, and K562-Hck-T338A cell populations were incubated for 72 h in the absence or presence of the indicated concentrations of imatinib. Apoptotic cells were stained with an anti-phosphatidylserine-Alexa Fluor 488-conjugated antibody and the percentage of apoptotic cells was determined by flow cytometry. A, shown are histograms from a representative flow cytometry experiment with the percentage of apoptotic cells shown above each plot. PS, phosphatidylserine. B, bar graph showing the average percentage of apoptotic cells from three independent experiments ± S.D. Two-way ANOVA showed significant effects of Hck or Hck-T338A expression (p < 0.0001) and of imatinib treatment (p < 0.0001). One-way ANOVA was performed separately for each concentration of imatinib across the three groups. The percent apoptosis among the three cell lines was statistically significant at 0.3 µM imatinib and 1 µM imatinib (*, p = 0.0003 and **, p = 0.006). Bonferroni’s post test for multiple comparisons showed a p < 0.001 for Neo versus Hck and Neo versus Hck-T338A at 0.3 µM imatinib and a p < 0.05 for Neo versus Hck and Neo versus Hck-T338A at 1 µM imatinib.

FIGURE 3. Expression of wild-type Hck or Hck-T338A protects K562 cells from imatinib-induced apoptosis. K562-Neo, K562-Hck, and K562-Hck-T338A cell populations were plated in soft agar with various concentrations of imatinib and NaPP1 and incubated for 7 days to allow colony formation. Imatinib-induced inhibition of colony formation was determined for each concentration of imatinib shown. A, colonies were stained 7–10 days later, and representative images of the plates are shown. B, the bar graph shows the average number of colonies from three independent experiments ± S.D. Two-way ANOVA showed significant effects of Hck or Hck-T338A expression (p < 0.0001) and of imatinib treatment (p < 0.0001). One-way ANOVA was performed separately for each concentration of imatinib across the three groups. The effect of imatinib on colony formation by the three cell lines was statistically significant at 0.03, 0.1, and 0.3 µM (*, p = 0.0003; **, p < 0.0001; ***p = 0.0015, respectively). Bonferroni’s post test for multiple comparisons showed a p < 0.001 for Neo versus Hck and Neo versus Hck-T338A at 0.3 µM imatinib and a p < 0.01 for Neo versus Hck and Neo versus Hck-T338A at 0.3 µM imatinib.

Next, we determined whether NaPP1 resensitizes K562-Hck-T338A cells to imatinib in the soft-agar colony assay. Each K562 cell population was plated in soft agar with various concentrations of imatinib and NaPP1 and incubated for 7 days to allow colony formation. Imatinib-induced inhibition of colony formation was determined for each NaPP1 concentration, and the results are plotted in Fig. 6. In addition, colony formation IC50 values for imatinib were determined by curve-fitting. Imatinib inhibited K562-Neo colony formation with an IC50 of 35.9 ± 6.8 nm in the absence of NaPP1, and this value was unaffected by NaPP1 treatment. Overexpression of wild-type Hck or Hck-T338A in K562 cells increased the IC50 value for the inhibition of colony formation by imatinib to about 4-fold, to 135 ± 11 and 146 ± 17.7 nm, respectively. Similar to the K562-Neo control cells, NaPP1 did not affect the inhibition of colony formation by imatinib in K562 cells expressing wild-type Hck.

level similar to K562-neo control cells, demonstrating that resistance to imatinib-induced apoptosis requires the kinase activity of Hck.

FIGURE 4. Expression of wild-type Hck or Hck-T338A protects K562 cells from imatinib-induced inhibition of soft-agar colony formation. K562-Neo, K562-Hck, and K562-Hck-T338A cell populations were plated in soft agar colony assays in the presence of the concentrations of imatinib shown. A, colonies were stained 7–10 days later, and representative images of the plates are shown. B, the bar graph shows the average number of colonies from three independent experiments ± S.D. Two-way ANOVA showed significant effects of Hck or Hck-T338A expression (p < 0.0001) and of imatinib treatment (p < 0.0001). One-way ANOVA was performed separately for each concentration of imatinib across the three groups. The effect of imatinib on colony formation by the three cell lines was statistically significant at 0.03, 0.1, and 0.3 µM (*, p = 0.0003; **, p < 0.0001; *** p = 0.0015, respectively). Bonferroni’s post test for multiple comparisons showed p < 0.001 for Neo versus Hck and Neo versus Hck-T338A at 0.3 µM imatinib and p < 0.01 for Neo versus Hck and Neo versus Hck-T338A at 0.3 µM imatinib.
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Hck, consistent with the lack of NaPP1 activity against the wild-type kinase. In contrast, the addition of NaPP1 to K562-Hck-T338A cells induced a dose-dependent reversal of the inhibition of colony formation by imatinib, with the IC_{50} value dropping from 146 ± 17.7 to 32 ± 6.6 nM in the presence of 3 μM NaPP1. These results demonstrate that selective inhibition of Hck-T338A kinase activity completely restores sensitivity to imatinib in terms of colony-forming activity.

To determine whether the reversal of imatinib sensitivity was associated with inhibition of Hck-T338A tyrosine kinase activity, Hck was immunoprecipitated from cells after exposure to the same imatinib concentrations used in the apoptosis and colony assays in combination with 3 μM NaPP1 (Fig. 7). Quantitative immunoblot analysis using an antibody specific to phospho-Tyr-416 in the activation loop of the active form of Hck showed that NaPP1 completely inhibited the T338A mutant of the kinase, consistent with the complete reversal of imatinib sensitivity observed at this concentration in both the apoptosis and colony assays. In contrast, NaPP1 did not inhibit the activity of wild-type Hck. To confirm the specificity of NaPP1 further, we also immunoprecipitated endogenous Lyn and performed similar immunoblot analysis with the anti-phospho-Tyr-416 antibody. NaPP1 did not affect Lyn activation loop phosphorylation in the presence or absence of imatinib (data not shown).

Overexpression of Hck in K562 Cells Increases Tyrosine Phosphorylation of Multiple Proteins in the Presence of Imatinib—To explore the mechanism of Hck-induced imatinib resistance, we performed quantitative anti-phosphotyrosine immunoblotting on lysates from K562-Neo, K562-Hck, and K562-Hck-T338A cells after treatment with DMSO carrier solvent (control), 1 μM imatinib, 3 μM NaPP1, or a combination of both compounds. As shown in Fig. 8, expression of wild-type Hck and the Hck-T338A mutant enhanced the tyrosine phosphorylation of multiple proteins in the presence of imatinib.
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FIGURE 7. NaPP1 selectively inhibits Hck-T338A in K562 cells. K562-Neo, K562-Hck, and K562-Hck-T338A cell populations were grown in 0.5% FBS overnight and treated with the indicated concentrations of imatinib and NaPP1 (3 μM) for 5 h. Hck was immunoprecipitated from clarified cell lysates and immunoblotted with a phosphospecific antibody that recognizes the tyrosine-phosphorylated activation loop of Hck (pHck). Duplicate membranes were blotted for Hck as a loading control. Western blots were analyzed using the Odyssey Infrared Imaging System. The Hck phosphotyrosine signal intensities were normalized to the levels of Hck protein from the blots of two independent experiments, and the average intensity ratios ± S.D. are presented in the bar graph. The entire experiment was repeated twice from independently derived cell populations with comparable results; representative blots are shown at the top. Two-way ANOVA was performed to determine statistical significance of the effect of imatinib and NaPP1 on the phosphorylation of Hck within each cell line. For K562-Hck, neither imatinib nor NaPP1 had a significant effect (p > 0.3), whereas for K562-Hck-T338A, NaPP1 treatment produced a significant reduction in Hck phosphorylation (p < 0.0001).

FIGURE 8. Wild-type Hck or Hck-T338A overexpression enhances tyrosine phosphorylation of several proteins in K562 cells without changing the overall tyrosine phosphoprotein banding pattern. K562 cells were infected with recombinant retroviruses carrying a neomycin selection marker (Neo), wild-type Hck or Hck-T338A and selected with G418. The resulting cell populations were plated in 0.5% FBS overnight and treated with the DMSO carrier solvent, imatinib (1 μM), NaPP1 (3 μM), or imatinib plus NaPP1 for 5 h. Cell lysates were resolved by SDS-PAGE, transferred to nitrocellulose membranes, and probed with anti-phosphotyrosine (top) and anti-Hck antibodies. Immunoreactive proteins were simultaneously detected using the Odyssey Infrared Imaging System (LI-COR). As a loading control, replicate blots were also probed with an anti-actin antibody. Using the Odyssey 3.0 software, the position of each molecular weight marker was used to estimate the molecular weights of each tyrosine phosphoprotein. The positions of the three bands showing the most prominent changes in phosphotyrosine content after Hck expression are indicated (p210, p72, and p59). This entire experiment was repeated three times from independently derived cell populations and produced comparable results in each case.

Overexpression of Hck Is Associated with Enhanced and Sustained Phosphorylation of Bcr-Abl at Multiple Regulatory Sites in the Presence of Imatinib—Previous studies have shown that Hck and other SFKs transphosphorylate c-Abl and Bcr-Abl (52–54). This may in turn stabilize the active conformation of the Abl kinase domain and, thus, decrease sensitivity to imatinib as well as increase Bcr-Abl-dependent signaling. For example, both c-Src and Hck phosphorylate c-Abl on its activation loop at Tyr-412 (53), which promotes kinase activation and induces imatinib resistance (Abl numbering is based on the crystal structure of the Ncap-c-Abl core; PDB code 2FO0) (32). Moreover, Hck and other SFKs phosphorylate Bcr-Abl at Tyr-89 in the SH3 domain and Tyr-245 in the SH2 kinase linker region both in vitro and in CML cells (54, 55). Phosphorylation at either of these residues may disrupt the negative regulatory interaction between the SH3 domain and the SH2 kinase linker (56). Furthermore, Hck phosphorylates Bcr-Abl in the Bcr-derived portion of the protein at Tyr-177 (Bcr numbering), a residue that links Bcr-Abl to Ras activation and is required for leukemogenesis (57–60).

To test whether the imatinib resistance observed in response to Hck overexpression correlates with enhanced phosphorylation of Bcr-Abl at these key regulatory tyrosines, we performed quantitative immunoblotting with phosphospecific antibodies on lysates from K562-Neo, K562-Hck, and K562-Hck-T338A cells after treatment with imatinib and NaPP1. Fig. 10 shows that expression of wild-type or T338A Hck causes enhanced phosphorylation of the Bcr-Abl SH3 domain at Tyr-89 and that phosphorylation of this site is sustained in the presence of 0.03–0.3 μM imatinib. The addition of NaPP1 completely reversed this Hck-dependent phosphorylation of Tyr-89 in cells expressing Hck-T338A but not wild-type Hck. This effect of NaPP1 is
particularly marked at 0.1 and 0.3 μM imatinib and suggests that Hck directly phosphorylates this key regulatory site in the Bcr-Abl SH3 domain. Tyr-89 phosphorylation is markedly reduced in all three cell lines at 1.0 μM imatinib in the presence or absence of NaPP1, suggesting that Bcr-Abl kinase activity may be required to stimulate Hck or endogenous SFK activation, which in turn leads to phosphorylation of this site. Bcr-Abl autophosphorylation may occur at Tyr-89 as well. Expression of wild-type or T338A Hck also increased phosphorylation of the Bcr-Abl activation loop at Tyr-412 (Fig. 10). Similar to Tyr-89, phosphorylation at this site was resistant to inhibition by lower concentrations of imatinib and dramatically reversed by NaPP1 exclusively in K562-Hck-T338A cells. Hck-mediated phosphorylation of Tyr-89 and Tyr-412 may induce an active conformation of Bcr-Abl due to the disruption of the intramolecular negative regulatory interaction between the SH3 domain and the linker or via direct phosphorylation of the activation loop (54, 56). In either case such changes favor an active Bcr-Abl kinase domain conformation less compatible with imatinib binding (27).

Phosphorylation of Bcr-Abl in the SH2 kinase linker region on Tyr-245 was also enhanced after expression of either wild-type Hck or Hck-T338A, and this effect was sustained in the presence of imatinib except at the highest concentration tested (1 μM; supplemental Fig. S3). In this case, however, the addition of NaPP1 did not significantly inhibit Tyr-245 phosphorylation in cells expressing Hck-T338A, suggesting that the enhancement of phosphorylation observed is independent of Hck kinase activity. One possible kinase-independent mechanism may involve binding of Hck to the regulatory SH2 and SH3 domains of Bcr-Abl (11).

Finally, we investigated the effect of Hck expression on Tyr-177 phosphorylation in the Bcr-derived portion of the oncoprotein (supplemental Fig. S3). Bcr-Abl from K562-Hck and K562-Hck-T338A cells demonstrated enhanced phosphorylation of this site that persisted even in the presence of 1 μM imatinib, whereas phosphorylation of Tyr-177 in control cells was inhibited by about 50% at this imatinib concentration. The addition of NaPP1 partially reduced Bcr-Abl Tyr-177 phosphorylation in K562-Hck-T338A but not K562-Hck cells. Because phosphorylation of Tyr-177 creates a binding site for Grb2 (22, 23) and other adaptors, it is possible that Hck may increase Ras-dependent signaling via Tyr-177 that contributes to imatinib resistance. The inability of imatinib to completely reverse Tyr-177 phosphorylation in control cells suggests that other SFKs coupled to Bcr-Abl or possibly other non-receptor tyrosine kinase families contribute to phosphorylation of this site.

DISCUSSION

Resistance to imatinib, the first-line therapy for CML, is typically associated with reactivation of Bcr-Abl signaling due to selection of drug-resistant Bcr-Abl mutants. However, emerging evidence shows that other mechanisms of clinical resistance can occur in the absence of Bcr-Abl mutations. Consistent with the role of SFKs in CML pathogenesis, recent evidence has linked Hck and Lyn overexpression or enhanced activity to imatinib resistance in the absence of Bcr-Abl mutations (21, 22, 24, 61). In this report, we establish a direct cause and effect relationship between Hck overexpression and resistance to imatinib in a CML cell line with wild-type Bcr-Abl and demonstrate that drug resistance requires Hck kinase activity.

To determine whether imatinib resistance requires Hck activity, we generated a Hck allele (Hck-T338A) that is uniquely targeted by NaPP1, a bulky analog of the non-selective SFK inhibitor PP1. The advantage of this chemical genetic approach is that it allows for specific inhibition of Hck or endogenous SFK activation, which in turn leads to phosphorylation of this site. Bcr-Abl autophosphorylation may occur at Tyr-89 as well. Expression of wild-type or T338A Hck also increased phosphorylation of the Bcr-Abl activation loop at Tyr-412 (Fig. 10). Similar to Tyr-89, phosphorylation at this site was resistant to inhibition by lower concentrations of imatinib and dramatically reversed by NaPP1 exclusively in K562-Hck-T338A cells. Hck-mediated phosphorylation of Tyr-89 and Tyr-412 may induce an active conformation of Bcr-Abl due to the disruption of the intramolecular negative regulatory interaction between the SH3 domain and the linker or via direct phosphorylation of the activation loop (54, 56). In either case such changes favor an active Bcr-Abl kinase domain conformation less compatible with imatinib binding (27).

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As part of the Hck-T338A validation strategy, we first needed to establish that this mutant did not exhibit a loss or gain of function and that it was selectively sensitive to inhibition by NaPP1 but not affected by imatinib. Our results both in vitro and in cell-based assays show that the T338A mutation does not affect Hck catalytic or biological activities. Furthermore, we found that the T338A mutation made Hck almost 50 times more sensitive to NaPP1 than the wild-type kinase in vitro, without inducing sensitivity to imatinib. Importantly, NaPP1 did not inhibit a recombinant Abl kinase protein establishing the specificity of this compound for the Hck-T338A mutant.

Replacement of the corresponding gatekeeper residue in other protein-tyrosine kinases such as c-Abl, c-Src, or the platelet-derived growth factor receptor was recently reported to have kinase-activating effects as shown by the increased tyrosine phosphorylation of cellular proteins in transfected human 293T cells (62). However, these mutants were not able to transform murine BaF3 myeloid cells to interleukin-3 independence (62). In the case of Hck, evidence presented here shows that the replacement of the gatekeeper threonine with alanine does not induce a gain of function. Expression of Hck-T338A in Rat-2 fibroblasts did not induce cellular transformation or phosphorylation of endogenous substrates. Although these data indicate that the T338A mutation does not up-regulate Hck kinase activity, we cannot rule out the possibility that this gatekeeper mutation affects other biological functions.

Having validated the NaPP1-sensitive mutant, we next showed that both wild-type Hck and Hck-T338A overexpression are sufficient to restore K562 cell survival and colony-forming activity in the presence of imatinib. Colony formation in soft agar was strongly affected, with the IC_{50} value for inhibition of colony growth increased by 4-fold upon overexpression of either form of Hck. The resistance to imatinib induced by Hck-T338A overexpression was completely reversed by NaPP1 in a concentration-dependent manner in cells expressing Hck-T338A but not in cells expressing wild-type Hck. The reversal of imatinib sensitivity in terms of survival and colony growth correlated with a complete inhibition of Hck-T338A activity by NaPP1 treatment, whereas wild-type Hck activity was unaffected. These results show that imatinib resistance induced by Hck overexpression requires Hck kinase activity.

Hck-induced resistance to imatinib may occur through a feedback mechanism that sustains an active conformation of the Bcr-Abl tyrosine kinase domain and thereby promotes imatinib resistance. Using phosphospecific antibodies, we present evidence that both wild-type and T338A Hck expression enhance phosphorylation of Bcr-Abl Tyr-89 (SH3 domain) and Tyr-412 (activation loop) that is sustained in the presence of imatinib. Phosphorylation of both sites is markedly sensitive to NaPP1 solely in K562-Hck-T338A cells, strongly suggesting that Hck is responsible for phosphorylation of these sites in vivo. Recent work has shown that phosphorylation of Tyr-89, which localizes to the binding surface of the SH3 domain, disrupts its intramolecular interaction with the SH2 kinase linker and is predicted to promote an active conformation of the Abl core (56). Data presented here also show a Hck kinase-dependent increase in phosphorylation of the Abl activation loop tyrosine residue (Tyr-412), which may also increase Bcr-Abl activity and imatinib resistance. However, it is not clear whether this effect is due to direct Hck-mediated phosphorylation of Tyr-412 or disruption of intramolecular regulatory interactions within Bcr-Abl (e.g. via phosphorylation of Tyr-89 in the SH3 domain).

Our results also show an increase in Tyr-177 phosphorylation in the Bcr-derived part of Bcr-Abl that is partially sensitive to NaPP1 in K562-Hck-T338A cells. Because phosphorylation of Bcr-Abl Tyr-177 facilitates Ras signaling (22, 23), enhanced phosphorylation of this site may stimulate mitogenic and anti-
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apoptotic pathways downstream of Ras that contribute to the increased growth and survival of Hck-expressing K562 cells observed in the presence of imatinib. This finding may also help to explain the remarkable enhancement of K562 colony formation observed in the presence of imatinib after expression of either form of Hck in K562 cells.

Although our data suggest that Hck may directly influence Bcr-Abl activity or signaling to account for imatinib resistance, Bcr-Abl-independent mechanisms may play a role as well. For example, recent evidence shows that Bcr-Abl activates autocrine insulin-like growth factor-1 signaling through Hck and Stat5b (63). In other experimental systems, insulin-like growth factor-1 was shown to have anti-apoptotic effects due to β-catenin stabilization (64). Therefore, it is possible that Hck-induced imatinib resistance observed here may be mediated in part through insulin-like growth factor-1 release. In addition, several SFKs have been shown to phosphorylate the cytoplasmic domain of Muc-1, a transmembrane glycoprotein overexpressed in CML. Phosphorylation of this Muc-1 region promotes the binding, stabilization, and increased nuclear targeting of β-catenin (65, 66). Whether or not Hck-mediated stabilization of β-catenin through either of these pathways contributes to imatinib resistance will require further investigation.

Recently, several second-generation Bcr-Abl-tyrosine kinase inhibitors, such as nilotinib, have been developed to override the problem of imatinib resistance. This has led in turn to studies of nilotinib resistance mechanisms. One report describes the generation of a nilotinib-resistant K562 cell line by exposure to gradually increasing concentrations of the drug (47). In this cell line nilotinib resistance was linked to Lyn up-regulation. In addition, an increase in Lyn mRNA expression was found in two of seven patients that developed resistance to nilotinib (47). This report suggests that SFK up-regulation may represent a common mechanism of resistance to Bcr-Abl inhibitors in addition to selection of drug-resistant Bcr-Abl kinase domain mutants. This observation also suggests that patients developing imatinib resistance without Bcr-Abl mutations may also be cross-resistant to other Bcr-Abl inhibitors. In this context, it would be interesting to determine whether Hck overexpression also induces resistance to nilotinib.

In summary, imatinib is arguably the best known and most successful kinase inhibitor in clinical use, although resistance to this drug is a growing problem in CML. Mutations in the Bcr-Abl kinase domain account for many but not all cases of resistance. More vexing are a growing number of imatinib-resistant CML patients who do not exhibit mutations in the Abl kinase domain but instead show a correlation with overexpression of SFKs, which are not direct targets for imatinib action. Results presented in this report demonstrate for the first time that overexpression of the myeloid SFK Hck is sufficient to confer imatinib resistance to blast crisis CML cells that express wild-type Bcr-Abl and that resistance requires Hck kinase activity. Furthermore, our data suggest that the resistance mechanism may involve direct phosphorylation of Bcr-Abl by Hck, although enhanced phosphorylation of substrate proteins shared by these kinases is also a possibility. Inhibitors selective for Hck or other SFKs may be of particular clinical benefit in this type of imatinib-resistant CML.

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