Src Activation Plays an Important Key Role in Lymphomagenesis Induced by FGFR1 Fusion Kinases

Mingqiang Ren, Haiyan Qin, Ruizhe Ren, Josephine Tidwell, and John K. Cowell

Abstract

Chromosomal translocations and activation of the fibroblast growth factor (FGF) receptor 1 (FGFR1) are a feature of stem cell leukemia–lymphoma syndrome (SCLL), an aggressive malignancy characterized by rapid transformation to acute myeloid leukemia and lymphoblastic lymphoma. It has been suggested that FGFR1 proteins lose their ability to recruit Src kinase, an important mediator of FGFR1 signaling, as a result of the translocations that delete the extended FGFR substrate-2 (FRS2) interacting domain that Src binds. In this study, we report evidence that refutes this hypothesis and reinforces the notion that Src is a critical mediator of signaling from the FGFR1 chimeric fusion genes generated by translocation in SCLL. Src was constitutively active in BaF3 cells expressing exogenous FGFR1 chimeric kinases cultured in vitro as well as in T-cell or B-cell lymphomas they induced in vivo. Residual components of the FRS2-binding site retained in chimeric kinases that were generated by translocation were sufficient to interact with FRS2 and activate Src. The Src kinase inhibitor dasatinib killed transformed BaF3 cells and other established murine leukemia cell lines expressing chimeric FGFR1 kinases, significantly extending the survival of mice with SCLL syndrome. Our results indicated that Src kinase is pathogenically activated in lymphomagenesis induced by FGFR1 fusion genes, implying that Src kinase inhibitors may offer a useful option to treatment of FGFR1-associated myeloproliferative/lymphoma disorders. Cancer Res; 71(23): 1–11. ©2011 AACR.

Introduction

Human stem cell leukemia–lymphoma syndrome (SCLL), also known as 8p11 myeloproliferative syndrome, is a rare atypical myeloproliferative disorder (MPD; ref. 1). SCLL expresses a clinical phenotype with features of both lymphoma and, sometimes, eosinophilic MPDs. SCLL is characterized by a feature of stem cell leukemia (SCLL), an aggressive malignancy characterized by rapid transformation to acute myeloid leukemia and lymphoblastic lymphoma. It has been suggested that FGFR1 proteins lose their ability to recruit Src kinase, an important mediator of FGFR1 signaling, as a result of the translocations that delete the extended FGFR substrate-2 (FRS2) interacting domain that Src binds. In this study, we report evidence that refutes this hypothesis and reinforces the notion that Src is a critical mediator of signaling from the FGFR1 chimeric fusion genes generated by translocation in SCLL. Src was constitutively active in BaF3 cells expressing exogenous FGFR1 chimeric kinases cultured in vitro as well as in T-cell or B-cell lymphomas they induced in vivo. Residual components of the FRS2-binding site retained in chimeric kinases that were generated by translocation were sufficient to interact with FRS2 and activate Src. The Src kinase inhibitor dasatinib killed transformed BaF3 cells and other established murine leukemia cell lines expressing chimeric FGFR1 kinases, significantly extending the survival of mice with SCLL syndrome. Our results indicated that Src kinase is pathogenically activated in lymphomagenesis induced by FGFR1 fusion genes, implying that Src kinase inhibitors may offer a useful option to treatment of FGFR1-associated myeloproliferative/lymphoma disorders. Cancer Res; 71(23): 1–11. ©2011 AACR.

Authors' Affiliation: Georgia Health Sciences University School of Medicine, Augusta, Georgia

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Corresponding Authors: Mingqiang Ren, Georgia Health Sciences University Cancer Center, 1120 15th Street, CN 4121A, Augusta, GA 30912. Phone: 706-721-4381; Fax: 706-721-1671; E-mail: mren@georgiahealth.edu; and John K. Cowell, E-mail: jcowell@georgiahealth.edu

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perhaps, for other human disorders associated with dysregulated FGFR1 activity.

Materials and Methods

Cell culture and proliferation assays

All cell lines were cultured in RPMI (Invitrogen) with 10% FBS (Hyclone), at 37°C in 10% CO₂. For drug treatments, 40,000 cells per well were seeded in 96-well plates and incubated overnight, then treated with the either dimethyl sulfoxide (DMSO; control) or the drugs indicated in the results section at concentrations defined by the experiments. Cell viability was determined using Cell TiterGlo luminescence cell viability kits (Promega) and a SpectraMax M5e (Molecular Probes) luminescence plate reader. CellTrace Violet (Invitrogen) or PKH26 (Promega) and a SpectraMax M5e (Molecular Probes) luminescence reader was used to track cell division. In these approaches, cells are initially labeled with specific fluorochromes. As the cells divide, the fluorochrome is distributed to the daughter cells and so the intensity of fluorescence in the population declines at a rate proportional to the rate of cell proliferation.

Transduction and infection

The dominant negative mouse Src K295R/Y527F construct (Addgene) in pCMV5 was subcloned into the YFP pMIYII vector (a kind gift from Dr. Darío Vignali, St. Jude Children’s Research Hospital, Memphis, TN) and designed as pMIY-DNSrc. The presence of mutations was confirmed by sequencing. The procedure for transduction and infection was as described previously (17). Supernatants containing retroviral particles of the empty pMIYII vector and pMIY-DNSrc were generated using Phoenix package cells as described previously (17). Cells were infected with the respective supernatant 4 times. The infected cell pools were used for apoptosis assays 1 week after infection.

Cell apoptosis assay, cell cycle, and phospho-specific flow cytometry

For analysis of apoptosis, cells were stained with Annexin V and 7-AAD (BD Biosciences) following the manufacturer’s protocol. Appearance of Annexin V and 7-AAD in flow cytometric analysis indicated onset of apoptosis. Cell cycle was analyzed by flow cytometry using standard procedures following propidium iodide staining. For intracellular staining of phospho-Src following drug treatment, cells were fixed and permeabilized before reacting with a phospho-Src antibody (BD Biosciences) according to standard procedures. Cells were analyzed using a LSRII flow cytometer. All flow cytometry data was analyzed using FlowJo software (Tree Star).

Immunoprecipitation and Western blot analyses

Proteins were isolated using standard TRIzol procedures. For immunoprecipitations (IP), 1,500 µg of cell lysates were incubated with the FRS2 antibody (Sigma) at 4°C over night, followed by addition of protein A/G-agarose beads (Pierce). Immunoprecipitates, or whole-cell lysates (50 µg), were separated using SDS-PAGE and immunoblotted with the following specific antibodies: anti-FGFR1 and GAPDH (Santa Cruz), anti-Src and anti-pSrc (Cell signaling) and beta-actin (Sigma) using standard protocols.

Anchorage-independent growth

Cells were suspended in Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal calf serum and 0.3% agar and then seeded into 6-well plates precoated with 0.6% to 1.0% agar in DMEM and 10% FBS. Twenty thousand cells were seeded in triplicate wells. Medium was changed twice a week. Colonies were counted at 5× magnification 3 weeks postseeding.

Animals and treatment schedule

Female Balb/c mice (Taconic, 6–8 weeks old) were injected with ZNF112 or CEP2A cells at 2×10⁶ per mouse through the tail vein. One week after injection, mice were randomized to treatment with vehicle control or dasatinib (ChemieTek). ZNF112 transplanted mice were divided into 3 groups: Control (n = 10), dasatinib 20 mg/kg (n = 10), and dasatinib 40 mg/kg (n = 10). CEP2A mice were divided into 2 groups: Control (n = 13) or dasatinib 40 mg/kg (n = 10). Dasatinib was dissolved in 80 mmol/L citrate buffer (pH 3.1; ref. 18) and given orally using a gavage needle twice per day. The control group of mice was given an equal volume of the citrate buffer by gavage. All treatments were done 6 days per week for 4 weeks. All animal experiments were carried out under protocols approved by the Institutional Animal Care and Use Committee of the Medical College of Georgia.

Human CD34+ cord blood cell culture and infection

Human cord blood cells were obtained from the Cord Blood Bank of the Medical College of Georgia under approved Institutional Review Board protocols. CD34+ cells were isolated using the EasySep Cord Blood CD34 Positive Selection Kit (StemCell Technologies) following the manufacturer’s protocol and expanded in StemSpan SFEM medium (StemCell) supplemented with recombinant human cytokines: LDL 10 µg/mL, Flt-3 100 ng/mL, SCF 100 ng/mL, TPO 50 ng/mL, interleukin (IL)-3 20 ng/mL, and IL-6 20 ng/mL (R&D Systems). After 24 hours prestimulation, CD34+ cells were infected with either BCR-FGFR1 or the control MIEG3 vector as described previously (17).

Results

Transformation BaF3 cells by ZMYM2–FGFR1, BCR–FGFR1, or CEP110–FGFR1 is associated with Src activation

It was previously suggested that chimeric FGFR1 proteins had lost their ability to recruit Src kinase due to loss of the FRS2-binding site in the FGFR1 fusion proteins (19). However, it has also been shown that 8/9 members of the Src family kinases were highly phosphorylated in the human KG-1 myeloid cell line, which express the FGFR1OP2–FGFR1 fusion kinase (20). Activation of Src kinase plays a crucial role in the normal process of FGFR1 signaling dynamics (14, 15). This observation suggests that either the chimeric FGFR1 proteins...
can still recruit FSR2 or that Src family kinases are activated indirectly in the presence of the FGFR1 fusion kinases. On the basis of recently updated sequence of the FGFR1 gene (NCBI: NM_023110.2), the break point in all FGFR1-associated translocations occurs within intron 12 (previously described as intron 8). As a result, all chimeric FGFR1 proteins contain amino acids 429–822 (based on FGFR1 transcript variant 1) which retains part of the FRS2-binding site (Fig. 1A; refs. 21, 22). The extended FRS2-binding site involves amino acids 420 to 433 (21, 22), although a 5’-truncated FRS2-binding site retained is in the chimeric FGFR1 proteins (Fig. 1A), which raised the possibility that the FGFR1 fusion proteins may, in fact, be able to recruit the FRS2 adaptor protein. If this is the case for the FGFR1 fusion kinases, then constitutive activation of Src

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**Figure 1.** Activation of Src is associated with FGFR1 chimeric kinase expression. A, schematic outline of the FGFR1 break point junction in the chimeric FGFR1 fusion kinases shows conserved amino acids within the FRS2 consensus binding domain (amino acids 420–433; refs. 21, 22). B, cell proliferation rates in BaF3 cells transformed by ZMYM2–FGFR1 (ZMYM2-F), BCR–FGFR1 (BCR-F), or CEP110–FGFR1 (CEP110-F) show both IL-3 independence and an increased growth rate relative to cells carrying the empty vector (MIEGS). C, transduction of BaF3 cells (left) with the chimeric FGFR1 kinases identifies constitutive tyrosine phosphorylation in total cellular proteins using the anti-PY20 antibody compared with cells carrying the empty vector. These cells also show phosphoactivation of Src kinase (right). D, ectopic expression of ZMYM2–FGFR1 in adherent 293T cells leads to increased levels of phospho-Src. E, IP using anti-FRS2 antibodies coprecipitates the CEP110–FGFR1 fusion protein in CEP110–FGFR1 (CEP110-F)-transformed BaF3 cells, but not in cells transduced with the empty vector. F, when BaF3 cells expressing either ZMYM2–FGFR1 or CEP110–FGFR1 are treated with the TKI258 (400 nmol/L) FGFR1 inhibitor, activation of both the FGFR1 fusion kinases and Src is inhibited. In contrast, TKI258 does not affect activation of either BCR–FGFR1 or Src in BCR–FGFR1-transformed cells. 

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kinases may play an important role in SCLL-associated leukemogenesis. To investigate this possibility, we first determined whether Src activation could be detected in cells expressing exogenous FGFR1 fusion kinases.

BaF3 murine, pro-B lymphoma cells normally require IL-3 supplements for growth, and oncogenic transformation results in IL-3-independent growth. Following forced expression of either ZMYM2–FGFR1, BCR–FGFR1, or CEP110–FGFR1 [from retroviral constructs coexpressing green fluorescent protein (GFP)], BaF3 cells became IL-3 independent in all cases (Fig. 1B), compared with BaF3 control cells transfected with the empty vector (MIEG3) which died within 24 hours in the absence of IL-3 (data not shown). These observations indicated that these 3 FGFR1 fusion genes can successfully transform BaF3 cells. We then compared the growth rate of transformed BaF3 cells in the absence of IL-3 with that of MIEG3–BaF3 (GFP) sorted cells in the presence of IL-3. BaF3 cells transformed with all 3 fusion kinases proliferated significantly faster than MIEG3–BaF3 cells (Fig. 1B). Western blot analysis confirmed high expression levels of ZMYM2–FGFR1, BCR–FGFR1, and CEP110–FGFR1 in the transformed BaF3 cells (Fig. 1C). As expected, constitutive activation of the chimeric kinases was associated with significantly higher global tyrosine phosphorylation levels compared with MIEG3–BaF3 cells (Fig. 1C). High levels of activated Src were also observed in these cells using Western blot analysis with an antibody that recognizes Y416-activated Src (Fig. 1C). In addition, ectopic expression of the ZMYM2–FGFR1 chimeric kinase in adherent 293T cells, which have been used extensively to study chimeric FGFR1 signaling (23, 24), also induced Src activation (Fig. 1D). Next, we used representative BaF3–CEP110–FGFR1 and BaF3–MIEG3 cells to determine whether FRS2 interacts with the FGFR1 fusion protein. Following IP with an anti-FRS2 antibody, Western blot analysis clearly showed that FRS2 can co-IP with the CEP110 protein. Following IP with an anti-FRS2 antibody, Western blot analysis with an antibody that recognizes Y416-tyrosine phosphorylated Src showed that 72 hours exposure to dasatinib reduced cell proliferation in cells expressing ZMYM2–FGFR1 but not cells carrying the empty MIEG3 vector or treated with DMSO (vehicle).

Inhibition of Src activation in FGFR1 fusion transduced BaF3 cells inhibits cell proliferation

To determine whether inhibiting Src activation affects cell growth in FGFR1 chimeric kinase expressing cells, we treated BaF3 cells transformed with the 3 chimeric kinases with dasatinib for 24 hours at their GI50 doses. Western blot analysis shows that dasatinib can demonstrably reduce phospho-Src levels (Fig. 2A). Using CellTrace Violet Fluorescence to monitor cell proliferation, we then showed that growth of the IL-3-independent cells was reduced by 50% following treatment with 10 nmol/L dasatinib for 72 hours (Fig. 2B). Growth rate was even slower using 100 nmol/L dasatinib (Fig. 2B). The same treatments did not significantly affect growth of BaF3 cells carrying the empty vector (Fig. 2B).

Src activation occurs in primary lymphomas induced by FGFR1 fusion genes in vivo

We previously described a murine model for ZMYM2–FGFR1 SCLL using a bone marrow transduction and transplantation approach (17). These mice developed T-lymphoblastic lymphoma with a double positive, CD4+/CD8+, immunophenotype. We have since also developed murine models for BCR–FGFR1 and CEP110–FGFR1 SCLL. BCR–FGFR1 induces leukemias with a CD19+/B220+/CD45– immunophenotype that is consistent with pro-B lymphoma (Tidwell and colleagues, submitted manuscript). In the CEP110–FGFR1 SCLL model, mice developed both T-cell and B-cell lymphomas (Ren and colleagues, manuscript in preparation). Mice in all 3 FGFR1-associated SCLL models concomitantly develop myeloid lineage disease. Here we focused on the T and B lymphomas from these mouse models. Western blot analysis using an

Figure 2. Downregulation of activated Src correlates with growth inhibition. A, dasatinib treatment (100 nmol/L) leads to suppression of Src kinase autophosphorylation in cells expressing ZMYM2–FGFR1, CEP110–FGFR1, and BCR–FGFR1. B, flow cytometric tracking of cell division with CellTrace Violet shows that 72 hours exposure to dasatinib reduces cell proliferation in cells expressing ZMYM2–FGFR1 but not cells carrying the empty MIEG3 vector or treated with DMSO (vehicle).
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Figure 3. Src is activated in primary lymphomas. Western blot analyses of phospho-activated and total Src levels in normal thymocytes (Thy) and normal lymph nodes (LN) from normal Balb/c mice compared with cells derived from lymphoma in lymph nodes (Lym) from mice with leukemia/lymphoma from the 3 different fusion kinase models. In these experiments, Src activation is seen in the majority of primary tumors from the 3 different models.

anti–phospho-Src (Y416) antibody clearly shows high levels of Src activation in the majority of lymphomas from the 3 in vivo FGFR1-associated SCLL models (Fig. 3), suggesting that constitutive activation of Src could be involved in the etiology of the lymphomagenesis induced by the chimeric FGFR1 genes.

Src activation in stable cell lines from leukemia mice and inhibition of Src activation induce cell death

Recently, we generated several cell lines from the 3 different SCLL mouse models. ZNF112 carries the ZMYM2–FGFR1, BBC1 carries the BCR–FGFR1 rearrangement, and CEP2A and CEP5A carry CEP110–FGFR1, ZNF112, CEP2A, and CEP5A cells are derived from T-cell lymphomas. BBC1 cells have pro-B cell immunophenotypes. Both Western blot and phospho-flow analyses showed high levels of activated Src in these cell lines (Fig. 4A). When ZNF112, CEP2A, CEP5A, and BBC1 cells were treated with dasatinib in the 300 to 1,000 nmol/L range for 24 hours, phospho-flow and Western blot analyses showed an almost complete loss of phospho-Src (Fig. 4A). Consequently, these cells undergo apoptosis and cell death (Fig. 4B) with a concomitant reduction in the passage of cells through the cell cycle, showing that, for these cells, activated Src is a critical survival factor. Similar to the chimeric FGFR1-transformed BaF3 cells (Fig. 1E), IP with the anti-FRS2 antibody clearly shows FRS2 can co-IP with both the phosphorylated FGFR1 fusion protein and Src kinase (Fig. 4C) in ZNF112 and CEP2A cells, whereas IP with the rabbit IgG isotype did not. To further exclude the nonspecific effects of dasatinib, we infected ZNF112, CEP2A, and CEP5A cells with retrovirus constructs carrying either a dominant negative Src (pDNSrc; ref. 26) or control vector (pMIYII). One week postinfection, flow cytometry analysis showed 2 to 4 times greater cell death (double positive of Annexin V and 7-AAD) in the pDNSrc cells compared with the pMIYII cells (Fig. 4D). Taken together, these observations show that the chimeric FGFR1 may drive tumorigenesis through activation of Src kinase.

Knockout of Src impairs transformation of 3T3 cells by ZMYM2–FGFR1

Although dasatinib inhibits both BCR/ABL and Src family tyrosine kinases, it can also inhibit other tyrosine kinases, such as PDGFRα/b and KIT (27, 28). To show a specific association between FGFR1 chimeric kinase expression and Src activation, we introduced either the ZMYM2–FGFR1, or MIEG3, retroviral constructs into the Src kinase family null Src-Yes-Fyn (SYF) mouse embryo fibroblasts (MEF; ref. 29). Nontumorigenic NIH3T3 cells were used as controls. Stably infected, GFP-positive MEFs were isolated by fluorescence-activated cell sorting (Supplementary Fig. S1A). Western blot analysis showed that expression of ZMYM2–FGFR1 results in high levels of Src activation in normal NIH3T3 cells but not in the SYF−/− MEFs (Fig. 5A). Red fluorescent PKH26 was used to analyse cell-cycle progression which showed that ZMYM2–FGFR1 can significantly promote cell proliferation in NIH3T3 cells but not in SYF−/− MEFs (Supplementary Fig. S1B), which is consistent with activation of Src. To further support this observation, cell proliferation assays clearly show that ZMYM2–FGFR1–transformed 3T3 cell grow faster than cells transfected with the empty vector after 4 days. No difference was observed between ZMYM2–FGFR1 and MIEG3-transduced SYF cells (Fig. 5B). Furthermore, Soft-agar colony formation showed the development of larger colonies by the ZMYM2–FGFR1–infected NIH3T3 cells, compared with cells carrying the empty MIEG3 vector (Fig. 5C), although no difference was observed between ZMYM2–FGFR1- and MIEG3-infected MEF-SYF−/− cells (Fig. 5C). These results show that abrogation of Src activity can significantly reduce in vitro phenotypes of transformation in ZMYM2–FGFR1 expressing cells.

Dasatinib significantly reduces leukemogenesis

Src activation clearly plays a role in the in vitro tumorigenic potential of FGFR1 fusion kinases. To determine whether the same was true in vivo, we transplanted ZNF112 and CEP2A cell lines into normal female Balb/c mice and treated them with
dasatinib. Leukemogenesis was then monitored over several months. Our pilot studies showed that both of these cell lines predominantly induce T-cell leukemia/lymphoma in vivo. Although injection of 100,000 cells from either cell line typically results in SCLL within 3 weeks, for the drug treatment experiments, we injected 2 million cells into each mouse to ensure a large initial tumor burden. Dasatinib treatments were initiated 8 days after transplantation, to provide sufficient time for the injected cells to home to the bone marrow. Initially, we treated ZNF112 carrying mice with dasatinib at 20 mg per kg body

Figure 4. Inhibition of Src activation results in apoptotic cell death. A, flow cytometric and Western blot analysis of activated Src shows reduced phospho-Src levels in BaF3 cells stably expressing the 3 chimeric FGFR1 kinases following dasatinib treatment. ZNF112 expresses ZMYM2–FGFR1, CEP2A, and CEP5A cells express CEP110–FGFR1 and BBC1 expresses BCR–FGFR1. B, apoptosis and cell-cycle analyses show that dasatinib treatment (300 nmol/L for ZNF112, CEP2A, and CEP5A; 1,000 nmol/L for BBC, treated for 48 hours) remarkably increased cell apoptosis rate and decreased the percentage of cells in the S+G2 phase in ZMYM2–FGFR1 and CEP110–FGFR1 expressing cells. C, immunoprecipitation with anti-FRS2 in CEP2A and ZNF112 cells shows that both the phopho-FGFR1 fusion kinase and phospho-Src are present in the same immunocomplex. D, flow cytometry analysis shows that infection with a dominant negative K295/R/Y527F Src (pDNSrc) retroviral construct induces cell death in the cell lines expressing the chimeric FGFR1 kinases compared with the empty vector alone.
weight, according to the regimen described by Hu and colleagues (18). In these experiments, dasatinib prolonged the median survival time by 2 weeks (P < 0.05, Fig. 6A). We did not observe any remarkable side effects because of dasatinib treatment in these mice, such as diarrhea or reduced body weight. Postmortem analyses did not reveal evidence of gastrointestinal hemorrhage, which is found in approximately 8% of human patients treated with dasatinib (30). When we increased the dasatinib dose to 40 mg/kg body weight in subsequent experiments, leukemogenesis was significantly inhibited for both ZNF112 and CEP2A-injected mice compared with animals treated with the vehicle control (Fig. 6A and D). Two of 10 mice (20%) injected with ZNF112 cells and 30% (3/10) of mice injected with CEP2A show no GFP+ cells in their peripheral blood 4 months after transplantation (data not shown). Consistent with the survival analysis, the spleen weights in the dasatinib-treated groups were also significantly decreased compared with control groups (Fig. 6B and E). Flow cytometric analysis of these dasatinib-treated animals shows a remarkable decrease in GFP+ CD4+ CD8+ cells in the peripheral blood after only 2 weeks of treatment (Fig. 6C and F). Together, these data suggest that inhibition of Src activation or possibly other non-Src targets of dasatinib can inhibit leukemogenesis induced by FGFR1 fusion kinases.

Activation of Src is also seen in BCR–FGFR1–transduced human CD34+ cells and the KG-1 cell line

To determine whether Src is activated in human hematopoietic cells, we transduced BCR–FGFR1 into CD34+ progenitor cells isolated from human cord blood samples. Intracellular phospho-flow cytometry analysis showed that phospho-Src levels were increased 62% in the GFP+ BCR–FGFR1 subpopulation compared with that in CD34+ cells transduced with the control vector (Fig. 7B). Furthermore, when KG-1 cells were treated with either dasatinib or the FGFR1 inhibitors PD17073 or TKI258, phospho-Src levels were reduced 1.5- to 2-fold compared with the vehicle-treated cells (Fig. 7B), suggesting activation of Src in human hematopoietic progenitor cells. In addition, higher activated Src levels were also found in human KG-1 cells, expressing the FGFR1OP2–FGFR1 variant fusion gene (20), compared with the normal human mononuclear cells (Fig. 7B). Furthermore, when KG-1 cells were treated with either dasatinib or the FGFR1 inhibitors PD17073 or TKI258, phospho-Src levels were reduced 1.5- to 2-fold compared with the vehicle-treated cells (Fig. 7B), suggesting activation of Src in KG-1 cells could be associated with constitutive activation of FGFR1 fusion kinase. Treatment with dasatinib at low nanomolar concentrations also induces growth inhibition in KG-1 cells (Fig. 7C), indicating Src activation plays an important role in sustaining KG-1 survival and proliferation. Together, these observations show that activation of Src is one of the mechanisms that drive chimeric FGFR1–associated tumorigenesis and progression. Consequently, molecularly targeting Src kinase may provide a new therapeutic option for SCLL patients.

Discussion

SCLL is a distinct disease with bilineage involvement in the same patient. To date, a wide variety of therapeutic regimens have been used for patients with SCLL, including protocols for acute lymphoblastic leukemia, AML, chronic myelogenous leukemia, and for other myeloproliferative neoplasms or
myelodysplastic syndromes (6). Overall, the success of targeted therapy has not been achieved in patients with SCLL. In this study, we have shown that activation of Src kinase is consistently seen in vitro in cells transformed with different chimeric FGFR1 kinases, as well as in vivo in mice transduced and transplanted with FGFR1 fusion genes. Pharmacologic inhibition of Src kinase, using dasatinib treatment or genetic inactivation of Src using dominant/negative

Figure 6. Dasatinib reduces lymphomagenesis of ZNF112 and CEP2A cells. A, ZNF112 xenografted into Balb/c mice (n = 10) and then treated with DMSO (vehicle) rapidly succumb. Xenografted mice treated with 20 mg/kg (n = 10) dasatinib prolongs survival and at 40 mg/kg (n = 10) significantly prolongs survival. In these animals (B), spleen enlargement in the 40 mg/kg group are significantly reduced. **P < 0.05. Two weeks after treatment, using both doses shows reduced numbers of GFP+ cells in the peripheral blood of dasatinib-treated mice (C). In the same way, (D) treatment of CEP2A xenografts (n = 10) with 40 mg/kg dasatinib significantly prolongs survival and reduces spleen weight (E) in these animals. **P < 0.05. Reduced numbers of GFP+ cells is also seen in this model (F).

Figure 7. Constitutive Src activation in BCR–FGFR1–transduced human CD34+ and KG-1 cells. A, phospho-flow cytometry analysis showed that phospho-Src levels in BCR–FGFR1–transduced normal human CD34+ progenitor cells (GFP+ BCR–FGFR1) is increased 62% compared with MIEG3-transduced CD34+ cells (GFP+ MIEG3) based on the mean of fluorescent intensity (MFI) of the pSrc-PE channel. This experiment was repeated independently in triplicate and consistent results were obtained. B, KG-1 cells, expressing FGFR1OP2-FGFR1, contain higher levels of phospho-Src compared with normal human mononuclear cells (hPBMN). Treatment with dasatinib (100 nmol/L), or FGFR1 inhibitors PD17073 (50 nmol/L) or TKI258 (200 nmol/L), inhibited Src activation in KG-1 cells. C, treatment with dasatinib induces growth inhibition in KG-1 cells.
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Dasatinib can also significantly delay leukemogenesis in vivo and in 20% to 30% of cases can completely prevent tumorigenesis. These results provide proof-of-principle evidence that molecular targeting of Src kinase could be a potential strategy to treat SCLL patients, either alone or in combination with other drugs.

Src overexpression and activation is found in a large number of human malignancies, including lung, breast, pancreatic, colon, and prostate (see review refs. 31, 32). High-level activation of Src has also been linked to the development of Philadelphia chromosome-positive chronic myeloid leukemia (CML) and acute lymphoblastic leukemia (33). In normal cells, Src plays a critical role in mediating signal transduction via interactions with multiple proteins and protein complexes, including the normal FGFR1 signaling pathway (15). Wild-type FGFR1 is tethered in the cell membrane, whereas the ZMYM2–FGFR1 fusion protein, and other FGFR1 fusion molecules, are found in the cytoplasm due to loss of putative nuclear localization signals (34, 35). Previously, it was assumed that the chimeric FGFR1 proteins had lost their ability to recruit Src kinase due to loss of the juxtamembrane FRS2-binding site in the chimeric protein (6, 19). However, we have now shown that all FGFR1 fusion kinases retain part of the FRS2 (Fig. 1) which in IP experiments seem to be sufficient to bind FRS2 and lead to phosphoryactivation of Src in both ZNF112 (harboring ZMYM2–FGFR1) and CEP2A (containing CEP110–FGFR1) cells. Consistent with these observations, Gu and colleagues showed that 8/9 members of the Src family kinases were highly phosphorylated in the human KG-1 myeloid cell line, which harbors the rare variant FGFR1OP2–FGFR1 fusion kinase (20). These data indicate that activation of Src kinase may be a common event during initiation of FGFR1-related SCLL.

Dasatinib is a novel, potent, oral, multitargeting inhibitor of Src, Abi1/2, KIT, and PDGFRα/β at nanomolar levels (27, 28). In the clinic, dasatinib has shown efficacy in patients with imatinib-resistant CML, and, currently, dasatinib is also the most effective treatment for patients with accelerated-phase CML (30). To date, dasatinib has not apparently been used to treat SCLL patients. Here we show that dasatinib treatment can substantially prolong the survival of mice with chimeric FGFR1-induced tumors. Consistent with these observations, we also showed that the human KG-1 myeloid leukemia cells, which carry the FGFR1OP2–FGFR1 chimeric kinase, are also highly sensitive to dasatinib treatment. These results provide preclinical evidence that Src inhibitors might be the basis for clinic trials involving SCLL patients. Our previous studies suggested that multiple genetic and, possibly, epigenetic changes are required for ZMYM2–FGFR1 to fully drive leukemogenesis and progression (17). We have recently identified activating deletions in the 5′ region of Notch1 in all T-ALL samples from ZMYM2–FGFR1 leukemia/lymphoma mice (36) which provides convincing evidence that ZMYM2–FGFR1 can drive the development of T-lineage disease through nonrandom constitutive activation of Notch1. Interestingly, aberrant activation of Notch1 was also seen in cells both from primary leukemic cells from a patient with ZMYM2–FGFR1 SCLL, as well as KG-1 cells. Although these observations show that targeting either Src or Notch1 alone in SCLL patients may provide some benefit, targeting FGFR1, SRC, and Notch1 in combination might be more effective.

Treatment of cells expressing ZMYM2–FGFR1 and CEP110–FGFR1 kinases with TK1258 reduced FGFR1 activation, with a concomitant reduction in Src activation. However, TK258 did not significantly inhibit Src activation in cells expressing the BCR–FGFR1 kinase (Fig. 1F). Furthermore, targeting Src with dasatinib only induced apoptosis and death in vitro at high concentrations. Consistent with these observations, in vivo treatment of BCR–FGFR1 expressing cells with the same dosage of dasatinib that prolonged survival in mice transplanted with ZMYM2–FGFR1 and CEP110–FGFR1 expressing cells did not prolong survival in these BCR–FGFR1 mice (data not shown). SCLL patients expressing the BCR–FGFR1 kinase differ from SCLL patients expressing the other kinases, in that they present with an atypical CML, without the Philadelphia chromosome (3, 37). In mice, BCR–FGFR1 rapidly induces CML-like MPD and B-cell lymphoma, with significantly shortened survival times compared with ZMYM2–FGFR1 (ref. 19 and Tidwell and colleagues, submitted manuscript), as well as in CEP110–FGFR1–induced disease (Ren, in preparation). Unlike the other fusion partners, BCR itself also has kinase activity (38) and so potentially serves not only as an oligomerization domain to activate FGFR1 but can also trigger downstream signaling pathways in its own right. Mutation of the critical Y177 tyrosine in BCR–FGFR1, which is required to bind the GRB2 adaptor protein, significantly reduces murine SCLL development in vivo (19) and those mice that develop disease show T-cell lymphomas typical of the other fusion kinases. Taken together, these observations suggest that the BCR component of the fusion kinase may be driving the disease to the myeloid and B-cell lineage and that targeting FGFR1 alone in this disease is not as efficient a therapeutic approach as it is in the other chimeric kinase-induced diseases. From our data, it seems that this may be due in some part to the inability of FGFR1-targeting drugs to specifically suppress Src activation.

In summary, our present study shows that ZMYM2–FGFR1, BCR–FGFR1, and CEP110–FGFR1 chimeric proteins can constitutively activate Src kinase both in vitro and in vivo. Pharmacologic and genetic inhibition of Src function induces apoptosis and cell death in cells harboring these chimeric FGFR1 genes, suggesting that that Src activation is an important oncogenic event for FGFR1 fusion gene–driven disease development and progression. As such, molecular targeting of Src kinase should be considered as part of a combination treatment for SCLL patients.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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Mingqiang Ren, Haiyan Qin, Ruizhe Ren, et al.

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