Depletion of STAT5 blocks TEL–SYK-induced APMF-type leukemia with myelofibrosis and myelodysplasia in mice

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The spleen tyrosine kinase (SYK) was identified as an oncogenic driver in a broad spectrum of hematologic malignancies. The in vivo comparison of three SYK containing oncogenes, SYKwt, TEL–SYK and IL-2-inducible T-cell kinase (ITK)-SYK revealed a general myeloexpansion and the establishment of three different hematologic (pre)diseases. SYKwt enhanced the myeloid and T-cell compartment, without leukemia/lymphoma development. ITK–SYK caused lethal T-cell lymphomas and the cytoplasmic TEL–SYK fusion induced an acute panmyelosis with myelofibrosis-type acute myeloid leukemia (AML) with up to 50% immature megakaryoblasts infiltrating bone marrow, spleen and liver, additional MPN features (myelofibrosis and granulocyte expansion) and MDS stigmata with megakaryocytic and erythroid dysplasia. LKS cells were reduced and all subsets (LT/ST/MPP) showed reduced proliferation rates. SYK inhibitor treatment (R788) of diseased TEL–SYK mice reduced leukocytosis, spleen and liver infiltration, enhanced the hematocrit and prolonged survival time, but could not significantly reduce myelofibrosis. Stat5 was identified as a major downstream mediator of TEL–SYK in vitro as well as in vivo. Consequently, targeted deletion of_stats in vivo completely abrogated TEL–SYK-induced AML and myelofibrosis development, proving Stat5 as a major driver of SYK-induced transformation. Our experiments highlight the important role of SYK in AML and myelofibrosis and prove SYK and STAT5 inhibitors as potent treatment options for those diseases.

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INTRODUCTION

Constitutive activation of tyrosine kinases either by fusion of the kinase domain to dimerizing proteins or by point mutations inducing constitutive activation are a broadly accepted cause for cancer development. One player, the spleen tyrosine kinase (SYK) was shown to be involved in propagation of several hematologic malignancies.

SYKwt is expressed in most hematopoietic cells¹–⁴ and is involved in Fcγ receptor signaling.⁵ B- and T-cell antigen receptor signaling,⁶⁻⁸ immunoglobulin E receptor signaling,⁹ several interleukin receptors¹⁰⁻¹² and integrins like αβ³/β³,⁵,¹³,¹⁴ SYK belongs to the SYK/ZAP-70 family of non-receptor tyrosine kinases.¹⁵,¹⁶ Without receptor stimulation, SYK is autoinhibited and exists in a closed conformation. Upon activation of immunoreceptors, SYK becomes phosphorylated by SRC family kinases and binds to immunoreceptor tyrosine-based activation motifs mediated by its two tandem SH2 domains.⁸,¹⁷,¹⁸ SYK activation induces phosphorylation of SLP65, SLP76, PLCγ1/2 and VAV, resulting in activation of the phosphatidylinositol 3-kinase pathway, calcium ion signaling and mitogen-activated protein kinase signaling.¹⁹⁻二十五

Overexpression and activation of SYKwt was identified in various B-cell lymphoma subtypes.²⁶⁻²⁸ In chronic lymphocytic leukemia,²⁹ SYK functions as a downstream signaling mediator of the autoreactive B-cell receptor³⁰ and propagates microenvironment-driven chemokine receptor signaling like CXCR4.¹³ In acute myeloid leukemia (AML) cells, constitutive activation of SYK occurs independent from the driving oncogene, but depends on tonic activation of the Fc-γR1 and Mac-1 receptors, stimulated by cytokines delivered from the bone marrow (BM) niche.³¹⁻³³ Beside constitutive activation of SYK through upstream signaling events, two fusion oncogenes, interleukin-2 (IL-2)-inducible T-cell kinase (ITK)-SYK and TEL–SYK contain the constitutively activated tyrosine kinase domain of SYK.

ITK–SYK arises from a fusion between SYK and the ITK. It has been identified as a recurrent translocation in 17% of patients with unspecified peripheral T-cell lymphomas.³² The highly aggressive disease is characterized by infiltration of skin, spleen, lymph nodes, BM and other organs with mature T cells. The ITK-part of the ITK–SYK fusion contains a Pleckstrin-homology domain targeting the protein to the plasma membrane, and a Tec-homology domain, which is linked to the tyrosine kinase domain of SYK. Previously, we and others could show that expression of ITK–SYK in murine BM or CD4+ cells leads to T-cell lymphoma development in mice, reflecting all major characteristics from the human disease.³³,³⁴

TEL–SYK was identified in a patient with an atypical myelodysplastic syndrome with leukemic transformation.³⁵ The patient was characterized by refractory anemia, dysplasia of the megakaryocytic and erythroid lineage, as well as myeloid hyperplasia with excess blasts (RAEB-T) with megakaryocytic phenotype. The patient progressed to leukemia with CD41+ megakaryocytic blasts.³⁵

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In TEL–SYK, the E26 transforming-specific translocation variant gene 6 (ETV6) is fused to the C-terminal part of SYK containing the tyrosine kinase domain and the Interdomain B. Although ITK–SYK is mainly localized in lipid rafts at the cell membrane, TEL–SYK has a cytoplasmic localization and is constitutively activated through TEL-mediated oligomerization. Overexpression of the TEL–SYK fusion in murine pre B cells induces a B-ALL like disease in mice. Furthermore, the expression of TEL–SYK in fetal liver cells was shown to induce myelofibrosis and myelodysplasia in transplanted mice.

Given those previously published papers, SYK overexpression as well as constitutive activation by fusions or other activating mechanisms seems to be a general theme for many different hematologic malignancies. On the other hand, the SYK fusion partners, the activation mechanism and the fusion partner-dependent localization of the fusion protein seem to have a pivotal role in selecting the transformed population. Therefore, the aim of the study was to compare the transformation potential of the known oncogenic SYK fusions and SYK isoforms in vivo, to precisely monitor their diverging influence on different hematologic cell subsets and to identify the underlying cell context dependent signaling events with focus on signal transducers and activators of transcription (STAT) signaling.

**MATERIALS AND METHODS**

**Mouse experiments**

All animal experiments were conducted in compliance with the US department of Health and Human Service Guide for Care and Use of Laboratory Animals and in concordance with German law. Experiments were approved by the Regierungspräsidium Freiburg. BALB/c or C57BL/6 males were injected intraperitoneally with 5-fluorouracil (150 mg/kg) and were killed after 4 days. Bone marrow cells (BMCs) were harvested, followed by erythrocyte lysis (Qiagen, Hilden, Germany) and cultivated for 2 days in prestimulation medium with DMEM (Gibco, New York, NY, USA) with 10% FBS (Biochrom, Berlin, Germany), mSCF, mL-6 and mL-3 (Peprotech, Hamburg, Germany) before transplantation. BMCs were transduced twice with retroviral particles generated by HEK293T, which had been transfected with pMSCV/IREC/green fluorescent protein (GFP; pMIG) empty control vector or vector with integrated SYKisoforms, ITK–SYK and TEL–SYK together with an EcoPack packaging vector (BD Bioscience, Heidelberg, Germany). Infection efficiency in different cell subsets was assessed 24 h after the 2nd infection round. BMCs (6 × 10^5) expressing the GFP+ cells represent the cells carrying the respective oncogene or the control vector. (control group obtained 6 × 10^5 normal BMCs and transplanted into irradiated recipient BALB/c females. The control group obtained 6 × 10^5 control BMCs.

Deletion of Stat5 in vivo was performed with Stat5fl/flMx1Cre BMCs. The original Stat5fl mouse strain was developed by L Hennighausen et al. and Stat5fl/Mx1Cre has previously been described. Stat5fl/Mx1Cre or Stat5flo (as control) BMCs were retrovirally transduced with TEL–SYK as described above and retroorbitally transplanted into twice irradiated recipient BALB/c females (2 × 450 cGy). The Stat5a/b locus was excised by three times intraperitoneal injection of 250 μg Poly (I:C) every 4 days starting day 0 (see Hoelbl et al. 2010).

For SYK inhibitor treatment studies, 2 weeks after transplantation, mice were split into two similar groups (seven to eight mice per group) according to the GFP content in the peripheral blood and body weight. One group was treated via oral gavage twice a day with 40 mg/kg R788 (Shanghai Haoyuan Chemexpress, Shanghai, China), solved in 0.1% carboxymethylcellulose sodium, 0.1% methylparaben, 0.02% propylparaben (Sigma-Aldrich, Munich, Germany) for 21 days and the other group with vehicle.

Plasmids

The pMSCV/IREC/GFP (pMIG) vector, ITK–SYK and TEL–SYK constructs were described previously. The HuSYKisoforms pENTR/mtm221 vector was purchased from Imagenes (Berlin, Germany) and SYKisoforms was cloned into a gateway-adapted pMIG vector via homologous recombination (Invitrogen).

**Cell culture**

293T cells were cultured in DMEM medium (Gibco) supplemented with 10% fetal bovine serum (PAA Laboratories, Pasching, Austria) and 1% penicillin–streptomycin (Gibco). BaF3 and 32D cells were obtained from ATCC and cultivated in RPMI-1640 medium (Gibco) with 1% fetal bovine serum, 1% penicillin–streptomycin and 10% WEHI supernatant. For phosphorylation experiments (phospho-flow or western blots), cells were cultivated without IL-3 for 15 h. Human KG-1 AML (erethroid leukemia, ATCC) and SET-2 cells (JAK2V617F+ megakaryoblastic leukemia, DSMZ) were cultured in RPMI-1640 medium with 10% or 20% FBS, respectively.

**Proliferation and apoptosis assays**

For SYK inhibition, the SYK inhibitor R406 (Axon medchem, Genentech, San Francisco, CA, USA) was dissolved in dimethylsulfoxide and used at a final concentration of 2 and 4 μM. For STAT5-inhibition pimozide (Calbiochem, San Diego, CA, USA) was used in dimethylsulfoxide at a final concentration of 5 and 10 μM. 32D cells supplemented with IL-3, 32D TEL–SYK cells without IL-3, 32D BCR–ABL cells without IL-3, KG-1 cells and

**Figure 1.** TEL–SYK induces a leukemic disease with anemia and thrombocytopenia in mice. (a) Schematic representations of SYKisoforms, ITK–SYK5SYK and TEL–SYK. PH, pleckstrin-homology domain; PNT, pointed N terminal domain; SH2, Src homology 2 domain; TK, tyrosine kinase domain; TH, Tec-homology domain. (b) Kaplan–Meier survival curve representing the survival of Balb/c mice either transplanted with control BM (empty pMSCV/IREC/GFP vector), SYKisoforms, ITK–SYK or TEL–SYK BM (n=8 per group). TEL–SYK mice died after 21–28 weeks, whereas ITK–SYK mice survived 42–76 weeks. Control and SYKisoforms mice died after 200 days without any disease symptoms. (c) Mean spleen weights of control, SYKisoforms, ITK–SYK5SYK and TEL–SYK transplanted Balb/c mice 25 and 42 days post transplantation. ITK–SYK5SYK and TEL–SYK mice develop a splenomegaly with an approximately fourfold increase in spleen weights for TEL–SYK mice after 25 days and a twofold increase for ITK–SYK mice after 42 days (left panel) (**p<0.05; ***p<0.01; ****p<0.001; unpaired t-test). (d) Peripheral blood samples of control, SYKisoforms, ITK–SYK5SYK and TEL–SYK mice were analyzed for red blood cell counts, hematocrit, WBC counts and thrombocytes 25 and 42 days post transplantation with the ADVAS120 blood analyzer (control 25d n=7, SYK n=5, ITK–SYK n=6, TEL–SYK n=7) **p<0.05; ****p<0.001; unpaired t-test. (e) May–Grünewald/Giemsa stain of blood smears of mice expressing control vector, SYKisoforms, ITK–SYK or TEL–SYK 25 days post transplantation. TEL–SYK mice show a strong expansion of differentiated granulocytes and megakaryoblasts in the peripheral blood bars. 50 μm. (f) Differential blood analysis from spleens of the different transfaction groups (n=4 mice per group) 25 days after transplantation. The graph represents the percentage of neutrophils, lymphocytes, bands, monocytes, eosinophils and blasts within total leukocytes and is given as mean±s.e.m. (g) Flow cytometry analysis for CD11b (myeloid cells), B220 (B cells) and CD90/CD3 (T cells) in the peripheral blood of the different transfaction groups analyzed in (d). Bars represent the mean percentage±s.e.m. of CD11b+ cells, B220+ cells and CD90+/CD3+ cells within the GFP+ (black border) and the GFP- (gray border) population, to allow discrimination of oncocogene-driven cell intrinsic and cell extrinsic effects. GFP+ cells represent the cells carrying the respective oncogene or the control vector. (control n=6, SYK 25 days n=6, SYK 42 days n=5). TEL–SYK n=7, ITK–SYK n=7) **p<0.05; ***p<0.01, unpaired t-test. ** above the white bars represents the significance of the difference between the GFP+ and GFP- population within the same mice.

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SET-2 cells were treated as indicated and cell numbers were assessed by Trypan blue staining and counting via Neubauer chamber for 24–72 h. For apoptosis detection after 48 h treatment, cells were stained with AnnexinV-APC/7-AAD (BD Pharmingen, Uppsala, Sweden) according to the manufacturer’s instructions and measured by flow cytometry.

Flow cytometry
Surface staining of BMC and spleen cells previously treated with erythrocyte lysis buffer (Qiagen) was performed with antibodies from BD Pharmingen against CD3-PE-Cy7, CD4-APC, CD8-PB, B220-APC, Ter119-PE-Cy7, CD11b-PE-Cy7, CD19-PE-Cy7, Gr-1-PE-Cy7, Gr-1-PB and...
Sca1-PB; cKit-APC from BioLegend (San Diego, CA, USA), and CD90.2-PE-Cy7 from eBioscience (San Diego, CA, USA) and measured using the CyanADP flow cytometer (Beckmann Coulter). Flow cytometry data were analyzed using the FlowJo 7.6 software (Tree Star, Inc., Ashland, OR, USA).

Intracellular phospho flow cytometry
For intracellular phospho flow cytometry, spleens of SYK wt, TEL–SYK, ITK–SYK or control mice were harvested, passed through a 100-μm filter (Millipore, Billerica, MA, USA) and directly fixed in 2% formaldehyde.
followed by 30 min permeabilization with 90% methanol. Then cells were stained with APC-labeled antibodies against phospho-STAT5 pY694, phospho-STAT6 pY641, phospho-PLCγ1 pY783, phospho-PLCγ2 pY759 and phospho-SLP76 pY128 (all BD Phosflow) for 1 h and measured using the CyanADP flow cytometer (Beckmann Coulter).

Phosphorylation analysis after R406 (Axon medchem) treatment of 32D TEL-SYK cells (depleted of IL-3)− Kit+Sca1+CD135− cells, ST as Lin−Kit+Sca1+CD135−CD150+FLT3− cells, MPP as Lin−Kit+Sca1+CD135−CD150+ and myeloid progenitors (MPs) (Lin−Kit+Sca1−) were sorted with the ARIA III cell sorter (BD Biosciences). Samples were prepared according to the manufacturer’s instructions and were analyzed for EdU+ cells by flow cytometry (Beckmann Coulter). Cell cycle analysis

TEL−SYK mice and GFP-control mice were injected with 100μl 20g body weight of 10μl EdU (Click-it Edu Pacific Blue Flow Cytometry Assay Kit, Molecular Probes, Eugene, OR, USA). BMCS were harvested 15 h later and LT-HSCs (Lin−CD48−CD11b−CD100−CD150−CD44hi), ST-HSCs (Lin−Kit+Sca1+CD135−CD150−), MPs (Lin−Kit+Sca1+CD135−CD150−) and myeloid progenitors (MPs) (Lin−Kit+Sca1−) were sorted with the ARIA III cell sorter (BD Biosciences). Samples were prepared according to the manufacturer’s instructions and were analyzed for EdU+ cells by flow cytometry (Beckmann Coulter).

Gene arrays

TEL−SYK and GFP-control LT/ST-HSCs were pooled from three to four mice and sorted as described above. RNA from LT and ST-HSCs was extracted with the RNeasy Micro Kit (Qiagen). RNA integrity was analyzed by capillary electrophoresis using a Fragment Analyser (Advanced Analytical Technology) were used at 1:2500. Membranes were incubated with ECL (Amersham, London, UK) for 2 min and chemiluminescence was detected using a western blot detection system (Cotix 60, AGFA).

Immunohistochemistry staining

Paraffin-embedded spleen, liver and BM sections were acquired from the pathology department of the University of Freiburg and stained for hematoyxlin/eosin, periodic acid−Schiff stain and reticulin following standard procedures. Blood smears were stained with May−Grünüwald−Giemsa. Histology slides were analyzed by a pathologist from the Medical University Freiburg. Images were taken with the LCmicro from Olympus.

RESULTS

ITK−SYK and TEL−SYK, but not SYKwt cause a lethal hematologic disease in mice

To investigate differences of SYKwt, ITK−SYK and TEL−SYK on hematopoiesis development and transformation, the oncogenes were cloned into a retroviral vector (pMSCV/IRES/GFP = pMIG; Figure 1a, composition of SYKwt and SYK fusion domains), expressed in 5-Fluorouracil-pretreated BM and BM cells were then infected and sublethally irradiated Balb/c recipients. BM infection rates measured by GFP content 48 h after infection were in between 25 and 50% for all constructs and infection of different cell subsets (B cells, T cells, myeloid cells, HSCs) was equal in between the different groups and as previously described.

The three SYK oncogenes caused remarkable differences in disease development. Three weeks after transplantation, TEL−SYK
mice developed general disease symptoms (weight loss of more than 20%, reduced mobility, rough fur) and all mice in that group (n=8) had to be killed due to final disease within 28 days after transplantation (Figure 1b). ITK-SYK-transplanted mice also developed a lethal disease, but had nearly twice as long survival time of 41–76 days. SYK wt, as well as GFP-control mice did not show any signs of disease until 200 days after transplantation (Figure 1b).

Analysis of spleens and livers at the time of death revealed a significant fourfold increase in spleen weights for TEL–SYK mice compared with controls (mean 506 mg versus 118 mg for control; Figure 1c, Supplementary Figure S1). Spleen weights of ITK–SYK mice were twofold increased (mean 255 versus 121 mg for control; Figure 1c), whereas spleens of SYK wt mice were normal up to 200 days after transplantation. Western blot analysis of spleens or BM of diseased mice showed expression of the respective oncogene (Supplementary Figure S2). Liver weights were normal for all transplantation groups (Supplementary Figure S3).

Blood analysis of SYKwt mice showed no lasting alterations in red blood cell counts, white blood cell (WBC) counts,
Figure 4. Hierarchical clustering and Gene set enrichment analysis (GSEA) of differentially expressed genes in TEL-SYK or GFP LKS subpopulations. (a) Hierarchical clustering was used to visualize genes that were differentially expressed between the four analyzed subpopulations compared to the median expression levels among the groups with a false discovery rate (FDR) < 0.005. Blue shades indicate a lower gene expression level than the median, whereas red shades indicate higher gene expression levels. Three clusters can be identified which are specific for TEL-SYK-expressing LT-HSCs and/or ST-HSCs. The gene names are listed for each cluster. (b) Enrichment plots and heat maps for core enrichment genes were generated by GSEA using the KEGG gene sets. NES, nominal P-value and FDR are shown for each gene set. Comparison of TEL-SYK versus GFP ST-HSCs identifies downregulation of the ‘Cell Cycle’ gene set and upregulation of the ‘Cytokine Cytokine Receptor Interaction’ gene set. (c) Comparison of TEL-SYK versus GFP ST- and LT-HSCs identifies downregulation of the ‘Antigen Processing and Presentation’ and ‘Primary Immunodeficiency’ gene sets and upregulation of the ‘Amino Sugar and Nucleotide Sugar Metabolism’ gene set.
thrombocytes or GFP content over a time period of 200 days (Figure 1d, Supplementary Figure S4). In contrast, TEL–SYK mice developed an anemia with a decrease of the red blood cell count to half of that observed in the control group (TEL–SYK mean 4.4 × 10^6 cells/μl versus control 9.3 × 10^6 cells/μl) and a reduced hematocrit of 29.4% (control 45.0%; Figure 1d). Also thrombocytes

**Figure 1d:** Graphs showing the effects of R788 treatment on TEL–SYK Balb/c mice. **a** Percent survival over 200 days after treatment initiation. **b** Comparison of spleen and liver weights between Vehicle and R788 groups. **c** Hemoglobin (HGB) and hematocrit (HCT) levels with statistical significance marked. **d** GFP+ and GFP- cells in the bone marrow (BM). **e** Histological comparison of liver sections stained with H&E and BM sections stained with reticulin for R788 and Vehicle groups. **f** Reticulin score graphs.
were strongly reduced (TEL–SYK mean 45.7 × 10^3 cells/μl versus control 583.3 × 10^3 cells/μl). Furthermore, TEL–SYK mice developed a leukocytosis with a 10-fold increase in WBCs compared with controls (TEL–SYK mean 35.0 × 10^3 cells/μl versus control 3.1 × 10^3 cells/μl). Blood smears stained with May-Grünwald/Giemsa showed accumulation of mature granulocytes in the peripheral blood (Figures 1e and f), but also the occurrence of 9–15% immature blasts as signs of a leukemic disease (Figure 1f and Supplementary Table 1). Flow cytometry analysis revealed that the GFP+ population in TEL–SYK mice mainly consisted of CD11b+/Gr-1+–positive myeloid cells (TEL–SYK 85.56% versus control 38.9%), whereas CD90+/CD3+ T cells (both CD4 and CD8) and B220+ B cells were suppressed (Figure 1g, Supplementary Table 1, Supplementary Figure S5). The percentage of CD11b+ cells within the GFP-negative population in TEL–SYK mice was only 36.67%, indicating a cell intrinsic effect of TEL–SYK driving the accumulation of myeloid cells in the peripheral blood. Despite the occurrence of immature blasts in the blood smears, we could not detect an increase in cKit+ (CD117) immature myeloid cells in the peripheral blood, pointing towards an atypical phenotype of the visualized blasts (Supplementary Figure S5).

In contrast to TEL–SYK, SYKwt and ITK–SYK showed no difference in total leukocytes compared with controls and mice did not develop a myeloproliferative or leukemic disease (Figures 1d and e). Nevertheless, both, SYKwt and ITK–SYK, induced a relative expansion of the myeloid compartment (Figures 1f and g). Hereby, SYKwt, similar to TEL–SYK, induced an expansion of myeloid cells only within the SYK/GFP+ population. In contrast, ITK–SYK enhanced myeloid cells and reduced B220+ cells within the GFP+ and GFP− population, pointing towards a cell intrinsic effect as driver for myeloid expansion and B-cell suppression in those mice (Figure 1g).

SYK oncogenes drive myeloid and T-cell expansion and alter the HSC pool

To further characterize the disease in TEL–SYK mice, to compare it with the other oncogenes and to discriminate between cell intrinsic and cell extrinsic effects, hematopoietic cell subsets within the GFP+ (oncogene carrying) and GFP− population in spleens and BM were analyzed by flow cytometry. Transplantation groups were compared at 25 days (final disease for TEL–SYK) and 42 days (final disease for ITK–SYK) after transplantation.

Interestingly and consistent with the findings from the peripheral blood, the GFP+ cells in the BM and spleens of TEL–SYK mice consisted of more than 90% myeloid cells (CD11b+), whereas B cells (B220+), erythroid progenitors (Ter119+) and T cells (CD3+/CD90+; spleen) were suppressed (Figure 2a, Supplementary Table 2). Furthermore, we detected a significant decrease in Lin-cKit+ myeloid progenitors in the BM and also a reduction of Lin-Sca1+ lymphoid progenitors within the GFP+ population (Figure 2b). Relative amounts of LKS cells (percentage of Lin-cKit+Sca+ cells within complete BM) in the BM and spleens were also strongly reduced (Figures 2b and c). Further analysis of LKS subpopulations showed a dramatic reduction of total LT-HSCs and ST-HSCs and to a lower extent also of the MPP population in TEL–SYK mice (Figure 2d). Cell cycle analysis of TEL–SYK+ LKS cells by DNA staining showed reduced cells in S/G2 phase compared with controls (Figure 2e). Further analysis of the cell cycle status of all LKS subpopulations (LT/ST/MPP cells) and myeloid progenitors in the BM was performed by EdU in vivo staining for 15 h and confirmed the reduction of cycling cells mainly within the LT and ST-HSC compartment (Figure 2f).

In concordance with the loss of nearly all stem- and progenitor populations in BM and spleens, total cell counts in femurs and spleens of TEL–SYK mice were strongly reduced compared with control mice (femur control 7.3 × 10^9 total cells versus femur TEL–SYK 0.75 × 10^9 total cells), resulting in a dramatic loss of all cell types including myeloid cells and all progenitor subtypes in both organs (Figure 2g).

Retransplantation of 2 million cells isolated from the BM of TEL–SYK mice into secondary recipient mice did not induce engraftment of the disease, pointing towards the MDS phenotype as a major event caused by the fusion.

In contrast to TEL–SYK, total cell numbers in femurs and spleens of ITK–SYK mice were equal (or slightly increased in spleen) compared with GFP controls (Figure 2d). Analysis of the GFP+ population of ITK–SYK mice revealed a significant increase in myeloid cells in BM and spleens, although this increase within the GFP+ population was much lower than seen in TEL–SYK mice (Figure 2a). In concordance with results from the peripheral blood, the GFP− population also showed an increase in myeloid BMCs as sign of a cell extrinsic effect driving the myeloid expansion. Although B cells and erythroid cells (Ter119) were suppressed, we detected a significant increase of CD4+ and CD8+ T cells within the GFP+ population in the BM reflecting the previously described T-cell lymphoproliferative/lymphoma disease induced by ITK–SYK (Figure 2a, Supplementary Figure S6, Supplementary Table 2).

Interestingly, and in contrast to TEL–SYK, ITK–SYK also strongly enhanced the Lin-Kit-Sca+ population (Figures 2b and c) and the Kit-Sca+B220+ lymphoid progenitor (CLP-II) compartment within the GFP+ population (Supplementary Table 1) and induced a 10-fold expansion of LKS cells in BM and spleens within only 45 days (Figures 2b and c; Supplementary Table 2).

SYKwt mice also showed normal total cell numbers in femurs and spleens compared with GFP controls (Figure 2d). Relative cell distribution showed features of both, the TEL–SYK and ITK–SYK phenotype, with a cell intrinsic expansion of the myeloid compartment in the BM and the T-cell compartment in the spleen (Figure 2a). B cells and Ter119+ erythroid progenitors were reduced (Figures 2a–d; Supplementary Table 2).

Figure 5. SYK inhibitor treatment prolongs survival of TEL–SYK mice, but cannot reduce myelofibrosis. (a) Kaplan–Meier survival curve of R788 treated (n = 8) versus vehicle-treated (n = 7) Balbc/c mice transplanted with the TEL–SYK oncogene. Treatment started 2 weeks after transplantation. Vehicle mice died within 16 days after treatment initiation, whereas R788 treated mice were killed after 21 days, when significance was reached (log-rank test) and to compare organ infiltration. (b) Splenic and liver weights of both treatment groups compared at the survival end point. R788 treated mice had a 1.5-fold decrease in spleen weight and 1.3-fold decrease in liver weight. **P < 0.05; ***P < 0.01; ****P < 0.001, unpaired t-test. (c) Peripheral blood analysis of R788 treated (n = 7) versus vehicle-treated (n = 6) Balbc/c mice 14 days after treatment initiation shows hemoglobin (HGB), hematocrit (HCT), WBC and percentages of granulocytes and lymphocytes. *P < 0.05; **P < 0.01; ***P < 0.001, unpaired t-test and Mann–Whitney test for WBC. (d) Flow cytometry analysis of GFP+ and GFP− cells in the BM as well as GFP+ CD11b+ and GFP−/Ter119+ BM cells 16 days (vehicle treated) and 21 days (R788 treated) after treatment initiation. *P < 0.05; **P < 0.01; ***P < 0.001, unpaired t-test. (e) Hematoxylin and eosin-stained liver slides and reticulin stained BM slides are shown from one R788 treated mouse versus control mouse. Arrows show liver infiltrates in the control group, which are absent in the treatment group. (f) Grading of fibrosis in BM slides for vehicle treated versus R788 treated Balbc/c mice (n = 7). Grade 1 shows a slight expansion of reticulin fibers, grade 2 represents a dense network of reticulin fibers, whereas grade 3 shows a strong expansion of collagen fibers with osteoneogenesis. Grading was done by two researchers blinded to the group attribution by considering reticulin staining in three fields of vision at ×100 magnification.
Figure 6. TEL–SYK exclusively induces factor independency and STAT5/6 phosphorylation in myeloid and B-lymphoid cells. (a) Expression of SYKwt, ITK–SYK and TEL–SYK after retroviral infection and GFP-sorting of the IL-3 dependent cell lines BaF3 and 32D. For detection of SYKwt and ITK–SYK by western blot, a primary SYK antibody was used. For detection of TEL–SYK, we used a TEL antibody. (b) Growth curves for IL-3-depleted BaF3 and 32D cells expressing the different oncogenes over 5 days were performed by counting Trypan blue negative cells via Neubauer chamber. (c) Intracellular phospho flow cytometry for phosphorylated STAT5/6, PLCγ2 and SLP76 in BaF3 cells (upper panel) and 32D cells (lower panel) expressing the different constructs. (d) Western blot of BaF3 and 32D cells expressing either control vector or the different SYK oncogenes with antibodies against phosphorylated STATs, PLCγ, JAK2, SLP76, ERK and AKT. Left panel shows the phosphorylated protein, right panel the total protein.
TEL–SYK induces an AML with features of human acute panmyelosis with myelofibrosis in mice

To further characterize the disease in TEL–SYK mice, hematoyxin and eosin stainings, periodic acid–Schiff staining and reticulin staining were performed from BM, spleens and livers of the diseased mice and all slides were examined by a mouse pathologist.

BM from TEL–SYK mice showed signs of multilineage dysplasia as seen in myelodysplastic syndrome, with hypoplasia of erythrocytes showing nuclear fragmentation, irregular nuclear contours and abnormal mitotic figures. Myelopoesis showed abnormal nuclear maturation with open chromatin, lobulated nuclei and hypogranulation. Immature megakaryocytes/megakaryoblasts were strongly increased reaching up to 50% of total cell counts and partially displayed multiple separated nuclei or unlobulated nuclei as well as micromegakaryocytes (Figure 3a, Supplementary Figure S7). Megakaryocytes were mainly periodic acid–Schiff stain positive (Figure 2b) and were organized in clusters. The immature megakaryoblasts in the BM were phenotypically identical with blasts seen in the peripheral blood. Reticulin staining showed a grade 2–3 myelofibrosis, at least partially explaining the reduced total cell counts in TEL–SYK mice (Figure 3c, Supplementary Figure S7). TEL–SYK spleens had lost their normal follicular architecture, were strongly fibrotic and were infiltrated with myeloid cells, predominantly consisting of mature granulocytes, dysplastic erythrocytes and clusters of dysplastic and immature megakaryocytes (Figure 3c). The same cell types also infiltrated the liver (Figure 3a). Taken together and according to the Bethesda classification of nonlymphoid hematopoietic neoplasms in mice34 (myeloid disorders), the TEL–SYK phenotype in the majority of mice can be classified as AML with features of human acute panmyelosis with myelofibrosis (APMF). Due to the frequent MDS features, we also have to consider the diagnosis of a myelodysplastic syndrome with features of human refractory anemia with excess blasts and fibrosis (RAEB-F) and at least one mouse, with more than 50% megakaryoblasts in the BM can be diagnosed as megakaryocytic leukemia.

In contrast to TEL–SYK, ITK–SYK mice showed neither signs of dysplasia nor myelofibrosis in the BM nor displayed any accumulation of immature megakaryocytes (Figures 3a–c). Overlapping with TEL–SYK mice, granulopoiesis was enhanced in BM and spleens and even infiltrated the liver. Additionally and consistent with our previous publication, we also found infiltration of T cells into the splenic lymph follicles, the BM and the liver, reflecting the previously described lymphoproliferative disease caused by ITK–SYK mice. SYK wt mice had a normal splenic and BM architecture and no liver infiltration (Supplementary Figure S7). Compared with GFP controls and consistent with our flow cytometry data, SYK wt mice had enhanced relative numbers of mature granulocytes in BM and spleens.

TEL–SYK alters the gene expression profile of LKS subpopulations and downregulates genes important for B-cell development and cell cycle regulation

To further characterize the effect of TEL–SYK on the stem cell compartment, we sorted TEL–SYK+ LT-HSCs (Lin−Kit+Sca+CD150 +FLT3−) and ST-HSCs (Lin−Kit+Sca+CD150−FLT3+) and compared them with control counterparts by gene expression profiling. Hundred and eight genes with a P-value < 0.005 discriminated between the different groups and are shown by gene cluster analysis (Figure 4a and Supplementary Table 3). Gene cluster 1 shows genes exclusively overexpressed in TEL–SYK LT-HSCs. Gene cluster 2 shows genes overexpressed in TEL–SYK LT and ST-HSCs compared with GFP controls and cluster 3 shows genes downregulated in TEL–SYK+ LT and ST-HSCs. Genes identified in cluster 1 and elevated in TEL–SYK LT-HSCs are Spr2a1 (2x), Kcnd1 (potassium voltage gated channel) and Stxbp6 (syntaxin binding protein 6). In cluster 2, we found Gfz2f2 (general transcription factor 2 F), Enah (Mena, associated with epithelial-mesenchymal transition in solid tumors) and Ak2. Ak2 was described as being mutated in reticular dysgenesis leading to loss of neutrophils and lymphocytes35 and might therefore be involved in the MPN phenotype seen in our mice.

Genes downregulated in TEL–SYK LT and ST-HSCs (gene cluster 3) are mainly involved in early B-cell development (Blnk, IL7r, Ifng, Rag1 and so on), which fits to our observation that nearly all B cells are lost in TEL–SYK+ mice. Furthermore, we found genes downregulated that are involved in interferon-alpha signaling (Tlrp1,6,7 und 9; Figure 4a).

Gene set enrichment analysis using the KEGG gene set database from TEL–SYK+ versus control ST-HSCs showed downregulation of genes involved in cell cycle regulation (Figure 4b) supporting our in vivo observation of reduced proliferation rates and cell cycle alterations in TEL–SYK+ HSCs. Furthermore, within ST-HSCs, we found the upregulation of the ‘cytokine cytokine receptor interactions’ gene set with increased levels for IL-6, IL-13, IL-1B, LIF and many others recapitulating the cytokine storm shown in those mice (Figure 4b).36 Pathways involved in immune regulation, B-cell receptor signaling and antigen processing were reduced, which matches the lymphopenic phenotype of the mice, whereas metabolic pathways were increased (Figure 4c, Supplementary Table 4).

SYK inhibition can partially revert the TEL–SYK-induced phenotype

To investigate if the TEL–SYK-induced AML with myelofibrosis and myelodysplasia can be reverted by SYK inhibitor treatment, we treated mice with the R406 prodrug R788 starting 2 weeks after transplantation. SYK inhibitor treatment could significantly enhance the survival of the mice (Figure 5a), reduced the spleen and liver size (Figure 5b) and reduced the WBC count (Figure 5c). Relative lymphocyte counts increased, whereas granulocyte counts decreased, bringing relations of both blood components closer to normal state. The hemoglobin levels, the hematocrit and total erythrocyte counts after 14 days of treatment were significantly enhanced (Figure 5c). Comparison of cell subsets in the BM reflected the results from the peripheral blood with reduced neutrophils (CD11b+ cells) and enhanced Ter119+ erythroid progenitors in GFP+ and GFP− populations (Figure 5d). The GFP+ population in the BM was strongly reduced after treatment with R788, whereas total GFP− cells were not affected (Figure 5d). IHC staining also showed repopulation of the BM with erythroid progenitors and a reduction of megakaryoblast infiltration (Figure 5e). Liver infiltration by granulocytes and megakaryoblasts was nearly absent in the R788 treatment group (Figure 5e).

Interestingly, and in contrast to our expectations, myelofibrosis was not significantly reduced in the treatment group according to our reticulin staining (Figure 5f), which might indicate that this process once initiated, is not reversible by SYK inhibition in late stages of the disease.

TEL–SYK exclusively induces IL-3 independent growth and STAT5/6 phosphorylation in myeloid and B-lymphoid cells

To further examine the differences in signal transduction between TEL–SYK, SYK wt and ITK-SYK dependent on the background of the transformed cell, we overexpressed the oncogenes in myeloid (32D cells) and B-lymphoid (BaF3 cells) cell lines (Figure 6a).

Interestingly, only the overexpression of TEL–SYK, but not ITK–SYK or SYK wt, induced growth factor independence of the otherwise IL-3-dependent 32D- and BaF3 cells (Figure 6b). Therefore, we aimed to identify signaling pathways specifically activated by TEL–SYK and performed western blots with phospho-specific antibodies and additionally phospho-flow cytometry (Figures 6c and d, respectively).
Supplementary Figure S8. Myeloid 32D cells carrying TEL–SYK showed an exclusive activation of STAT5, SLP76 and PLCγ2 (p-PLCγ2 by flow cytometry), ERK and AKT. In B-lymphoid BaF3 TEL–SYK cells, we found a similar activation pattern with phosphorylation of STAT5, SLP76, PLCγ2 and ERK, no AKT phosphorylation, but additionally activation of STAT6.

In contrast, SYKwt and ITK–SYK did not activate any of those pathways in 32D cells (Figures 6c and d), but SYKmut was able to...
activate PLCγ2 and SLP76 in BaF3 cells as assessed by flow cytometry (Figure 6c).

STAT5 is an essential downstream target of TEL–SYK in vivo and is generally regulatable by SYK inhibitors. To verify if any of those pathways activated by the SYK oncopgenes in vitro is also relevant in vivo, we analyzed BM and spleen cells of SYK oncogene carrying mice using phospho-flow cytometry of GFP+ cells. TEL–SYK positive murine myeloid cells showed a strongly increased phosphorylation of STAT5 and STAT6 in vivo, supporting the in vitro results (Figure 7a). In contrast, other phosphorylation events like PLCγ2 activation could not be confirmed in vivo and SLP76 phosphorylation was only moderately increased. The phosphorylation pattern of SYK wt cells was completely different, with a strongly increased phosphorylation of PLCγ2 and SLP76, but not STAT5 and STAT6, which is in line with the in vitro results. According to those findings, we hypothesized that STAT5 and STAT6 phosphorylation, but not SLP76 and PLCγ2, make the difference in between malignant transformation caused by TEL–SYK or only myeloid expansion as caused by SYKwt. STAT5 is known to be a major player in oncogenic transformation of myeloid cells and is activated in myeloproliferative diseases and different types of AML. To verify that STAT5 is directly regulated by the oncogenic TEL–SYK kinase fusion in myeloid cells, we treated TEL–SYK 32D cells with the SYK inhibitor R406 for up to 2 h. SYK inhibition in TEL–SYK 32D cells resulted in a fast dephosphorylation of STAT5 within 10 min and induction of apoptosis and reduced proliferation in the absence of IL-3 (Figures 7b–d). SYK inhibition did not reduce STAT6 phosphorylation nor phosphorylation of SLP76 or PLCγ2 within these 2 h, pointing towards those genes as secondary phosphorylation events.

Previously published results point toward SYK as a major driver of human AML and SYK inhibition or deletion resulted in differentiation and reduced proliferation of AML cell lines.31 In concordance with those previous reports, SYK inhibition in the human AML cell line KG-1 reduced proliferation of those cells while not inducing apoptosis (Figures 7c and d). Furthermore, SYK inhibition resulted in the dephosphorylation of STAT5, but also PLCγ2 and SLP76. As the murine TEL–SYK model reflects features from a classical AML, but also characteristics of a megakaryocytic leukemia with accumulation of megakaryoblasts in the BM and myelofibrosis, we chose the human cell line SET-2 for further experiments. SET-2 cells were obtained from a patient with leukemia with accumulation of megakaryoblasts in the BM and mainly consisting of mature granulocytes, whereas TEL–SYK/Stat5−/− mice had normal WBC counts in between 1 and 3 × 10^3 cells/μl (Figures 8e and f) without enhancement of the myeloid population. Further analysis of BMcs and spleen cells showed, that the Stat5−/− did not only prevent the expansion of TEL–SYK+ myeloid cells, but blocked the expansion of the complete TEL–SYK+ population (GFP+ cells within total BMcs; Figure 8f). Histological evaluation revealed that not only the AML phenotype (liver and spleen infiltration with immature blasts, leukocytosis) was suppressed by the depletion of Stat5, but that also myelofibrosis was abrogated (Figure 8h), pointing toward Stat5 as the major mediator of TEL–SYK-induced disease development.

Taken together, SYK inhibition in different cell lines seems to merge in STAT5 dephosphorylation and can affect proliferation and viability of AML cells and megakaryocytic leukemia cells, despite their dependence on other oncogenes.

To further validate the role of STAT5 in TEL–SYK-induced transformation, 32D TEL–SYK cells were treated with the STAT5 inhibitor Pimozide. Pimozide treatment resulted in significant apoptosis induction and reduced proliferation, similar to that seen with the SYK inhibitor R406, supporting the role of STAT5 downstream of oncogenic SYK (Figures 7e and f).

Depletion of Stat5 blocks TEL–SYK-induced leukemia and myelofibrosis

To further validate the role of STAT5 downstream of oncogenic SYK, we overexpressed TEL–SYK in BMCs from conditional Stat5fl/fl Mx1Cre mice versus control Stat5fl/fl mice.64 This experimental system was previously used to demonstrate the addition of BCR–ABL-transformed leukemic cells to Stat5 signaling, whereas normal adult hematopoiesis was rarely affected (mild reduction of hematocrit and thrombocytes within the first 8 weeks of Stat5 depletion).41 TEL–SYK/Stat5fl/fl/Mx1Cre BMCs and TEL–SYK/Stat5fl/fl control BMCs were transplanted into irradiated recipients and treated with Poly-IC 7 days after transplantation to allow full engraftment of normal and leukemic cells before depletion of Stat5. Stat5 depletion could be detected by western blot analysis of total Stat5 protein levels and by intracellular phospho-Stat5 staining of hematopoietic cells derived from the BM at the end of the experiment (Figures 8a and b). Thirty-two days after transplantation, the control group developed a lethal disease with leukocytosis and increase in myeloid cells in the peripheral blood and the experiment was terminated. Comparison of the spleens showed a dramatic difference with TEL–SYK control spleens being enlarged to a medium weight of 630 ± 79 mg, whereas TEL–SYK/Stat5−/− spleens were normal in size and weight (mean 121 ± 9 mg; Figures 8c and d). Analysis of the peripheral blood at the time of death revealed a leukocytosis in the TEL–SYK control group with an average of 40 × 10^3 cells/μl mainly consisting of mature granulocytes, whereas TEL–SYK/Stat5−/− mice had normal WBC counts in between 1 and 3 × 10^3 cells/μl (Figures 8e and f) without enhancement of the myeloid population. Further analysis of BMcs and spleen cells showed, that the Stat5−/− did not only prevent the expansion of TEL–SYK+ myeloid cells, but blocked the expansion of the complete TEL–SYK+ population (GFP+ cells within total BMcs; Figure 8f). Histological evaluation revealed that not only the AML phenotype (liver and spleen infiltration with immature blasts, leukocytosis) was suppressed by the depletion of Stat5, but that also myelofibrosis was abrogated (Figure 8h), pointing toward Stat5 as the major mediator of TEL–SYK-induced disease development.
Figure 8. Depletion of Stat5 blocks TEL-SYK-induced leukemia and myelofibrosis. (a) Western blot showing depletion of Stat5 in Stat5<sup>fl/fl</sup>Mx1Cre BMCs versus control Stat5<sup>fl/fl</sup>BMCs from mice treated with three times injection of Poly (I:C) and 35 days after transplantation. (b) Intracellular phospho-Stat5 staining was assessed by flow cytometry from BMCs from mice treated as in A (n = 4 per group). (c) Photographs showing spleens of three TEL–SYK control and three TEL–SYK Stat5<sup>−/−</sup> mice in comparison after 35 days. (d) Spleen and liver weights from TEL–SYK control and TEL–SYK Stat5<sup>−/−</sup> mice. (e) Peripheral blood of TEL–SYK control and TEL–SYK Stat5<sup>−/−</sup> mice (n = 4) was analyzed for red blood cell count, hematocrit, WBC count, thrombocytes and granulocytes 35 days post transplantation with the ADVIA120 blood analyzer. Graphs show results for total WBCs and the percentage of granulocytes. (f) Photographs from blood smears of control TEL–SYK and TEL–SYK Stat5<sup>−/−</sup> mice, which were stained with May-Grünwald/Giemsa. Bars, 50 μm. (g) GFP contents, CD11b+/GFP+ expression, CD90+3+/GFP+ expression and Ter119+/GFP+ expression within total live cells was assessed by flow cytometry analysis in BM and spleen cells from TEL–SYK control and TEL–SYK Stat5<sup>−/−</sup> mice 35 days after transplantation (n = 4). *P < 0.05; **P < 0.01; ***P < 0.001, unpaired t-test. (h) Representative images showing hematoxylin/eosin and reticulin (left image) stained BM, spleen and liver slides from TEL–SYK control and TEL–SYK Stat5<sup>−/−</sup> mice. Bars, 50 μm.
DISCUSSION

Oncogenic SYK was found to be an important driver of different hematologic malignancies.\textsuperscript{26,27,29,46} In our study comparing the different SYK oncogenes in mice, we could confirm the diverse picture of SYK involvement in human disease, with ITK–SYK driving T-cell neoplasias\textsuperscript{52} and TEL–SYK inducing an AML phenotype. Interestingly, myeloid expansion was seen with all SYK oncogenes, even with SYK\textsuperscript{wt}, pointing toward this cell type as being most susceptible to alterations in expression and activation levels of the SYK kinase. Interestingly, myeloid expansion was not always driven by a cell intrinsic effect as seen with the TEL–SYK oncogene, but was partially driven by cell extrinsic effects at least in the ITK–SYK mouse model. ITK–SYK caused a cell intrinsic T-cell expansion in the BM, which induced a strong cell extrinsic (GFP\textsuperscript{−} and GFP\textsuperscript{+} cells) expansion of the myeloid compartment and even a 10-fold increase in the LKS compartment. Several cytokines, like interferon-\textgamma, were reported to be highly upregulated in the ITK–SYK mouse model\textsuperscript{23} and might cause this phenomenon.

Interestingly, SYK\textsuperscript{wt}, expressing mice showed an intermediate phenotype with a cell intrinsic myeloid, but also T-lymphoid expansion. In contrast to the other two oncogenes, SYK\textsuperscript{wt} is not bound to a certain cell compartment, but can still alter its localization and despite its overexpression is still controllable by cell intrinsic regulatory processes like phosphorylation events or protein degradation. In contrast, TEL–SYK is constitutively activated by the TEL oligomerization domain and is fixed in the cytoplasm, which might be causative for the exclusive myeloid, but not T-lymphoid expansion. Despite our expectations, we could not detect an expansion of the B-cell population, although SYK\textsuperscript{wt} is thought to be a target in a broad spectrum of B-cell lymphoma subtypes.\textsuperscript{26,27,29,46} This might on one hand argue for a minor role of SYK in B-cell malignancies, but could also be caused by our experimental design of using 5-Fluorouracil-pretreated BM, which has lost most of the target B cells for the retroviral infections.

The TEL–SYK phenotype seen in the mice strongly resembles the disease from the patient, where the fusion kinase was originally identified.\textsuperscript{35} Both mice and men show an MDS phenotype in the BM with signs of dysplasia in the erythroid and megakaryocytic lineage. In both cases, there was a myeloid hyperplasia and an increase in immature blasts with a megakaryocytic phenotype in the BM. The patient later progressed to AML with megakaryocytic blasts. According to the new WHO classification of 2008 and the Bethesda classification for murine leukemias, the disease can be characterized as AML with features of human APMF. The fact that the disease is not transplantable does not interfere with this diagnosis, as the other main criteria for AML (more than 20% blasts in the BM and lethal disease within 4 weeks) are fulfilled. A recently published paper, where the investigators overexpressed TEL–SYK in fetal liver cells, shows part of the phenotype that we observe in our mice.\textsuperscript{39} The authors describe the same MDS and MPN features (granulocytosis, myelofibrosis), but the disease is not in all cases lethal and mice succumb to disease at a much later time point at 60 days after transplantation. Furthermore, the mice do not accumulate megakaryoblasts in the BM (as described here and in the patient) and also have no blasts in the peripheral blood and can therefore not be diagnosed as leukemia. The major difference in our experimental procedures is the use of adult BM instead of fetal liver cells. To further support our diagnosis, we overexpressed the TEL–SYK oncogene not only in Balb/c mice, but also in C57BL/6 mice, and here we find the APMF phenotype with infiltration of the BM and liver with megakaryoblasts (data not shown).

One mouse with more than 50% megakaryocytic blasts in the BM has even reached the criteria for a megakaryocytic leukemia bringing megakaryocytic disorders into the focus of our research. SYK is highly expressed within the megakaryocytic lineage\textsuperscript{35} and was previously shown to be essential for migration of megakaryocytes from the BM niche towards the vascular niche.\textsuperscript{48} In megakaryocytes and other hematopoietic cells, the SYK kinase has a critical role in the inside-out activation of integrins by glycoprotein receptors such as the collagen receptor GPVI\textsuperscript{49,51}. In addition, SYK has an important role in outside-in signals from integrins that mediate changes in cytoskeletal organization leading to cell spreading and motility.\textsuperscript{32} Megakaryocytic maturation is guided by microenvironmental factors provided by different niche cell subsets and platelet release only occurs in the vascular niche. Therefore, alterations in migration as induced by alterations in SYK activity\textsuperscript{46} could strongly affect the maturation processes and also the functionality of megakaryocytes and argues for a tight regulation of SYK expression as necessary for this process. In TEL–SYK mice, the SYK kinase is completely deregulated and constitutively activated within the cytoplasm, and its expression results in accumulation of immature and dysplastic megakaryoblasts in clusters in the BM. The mice develop a severe thrombocytopenia, showing the dysfunctionality of those cells. Dysplastic megakaryocytes are known to produce factors like platelet-derived growth factor\textsuperscript{53,54} or transforming growth factor-\textbeta,\textsuperscript{55,57} which are both potent stimulators of fibroblast proliferation and can contribute to myelofibrosis development, which is also seen in TEL–SYK mice. The fact that only 3 weeks are sufficient to cause a grade II–III myelofibrosis makes the TEL–SYK mouse model extremely interesting to identify further factors and cell types inducing or contributing to myelofibrosis and to test treatment strategies.

Previous publications found constitutive activation of SYK in AML induced by B2 integrins stimulated by factors coming from the BM niche and not by cell intrinsic oncogenes.\textsuperscript{5} In line with those findings, SYK inhibition in AML/MPN cell lines effectively induced apoptosis and reduced proliferation, even in the presence of other oncogenic driver mutations. For example, the cell line SET-2 is derived from a patient with megakaryocytic leukemia after essential thrombocytosis and has an activated Jak2V617F mutation. Despite the known dependence of this cell line on oncogenic JAK2, the SYK inhibitor R406 was highly effective in inducing apoptosis, reduced proliferation and even blocked STAT5 signaling in those cells, pointing towards megakaryocytic leukemias as susceptible towards SYK inhibition even in the presence of Jak2V617F.

STAT5 is known to be a major driver of myeloproliferative diseases and is regulated by JAK kinases.\textsuperscript{58,59} Previous publications have shown that SYK can directly interact with STAT5 and STAT3 in AML cell lines and can activate both proteins.\textsuperscript{5} In our hands, we could not detect STAT3 activation by our SYK oncogenes, but in all our in vivo and in vitro models, we found activation of STAT5 by SYK kinases as the converging event. Furthermore, in SET-2 cells, STAT5 dephosphorylation was the sole phosphorylation event influenced by the SYK inhibitor treatment. Furthermore, the targeted deletion of STAT5 could completely block leukemia development and also myelofibrosis otherwise caused by TEL–SYK, strongly supporting the role of STAT5 as the major mediator downstream of the SYK kinase. As STAT5 is known to be deregulated in many myeloid malignancies, SYK inhibitors might be a highly interesting target in influencing STAT5 activity downstream of the classical oncogenes like JAK2V617F, BCR–ABL or various AML oncogenes, especially in situations where resistance occurs against the primary target. Furthermore, our results strongly support the use of SYK inhibitors in megakaryocytic leukemias and also in myelofibrosis patients and can pave the way towards a clinical trial in this patient cohort.

CONFLICT OF INTEREST

The authors declare no conflict of interest.
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AUTHOR CONTRIBUTIONS
CS, HM, DB, TAM, JH, SK and JA performed the experiments; CK established the research; CD and CS wrote the manuscript. KA analyzed the histology slides.

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