Histone demethylase KDM4C is a functional dependency in JAK2-mutated neoplasms

Philipp Ernst1,2, Tina M. Schnöder3, Nicolas Huber3, Florian Perner3, Ashok Kumar Jayavelu4,5, Theresa Eifert3, Chen-Jen Hsu3, Nuria Tubío-Santamaría3, Carl C. Crodel1, Martin Ungelenk6, Christian A. Hübner6, Joachim H. Clement6, Andreas Hochhaus3 and Florian H. Heidel3,8

Mutations of the JAK2 gene are frequent aberrations in the aging hematopoietic system and in myeloid neoplasms. While JAK-inhibitors efficiently reduce hyperinflammation induced by the constitutively active mutated JAK2 kinase, the malignant clone and abundance of mutated cells remains rather unaffected. Here, we sought to assess for genetic vulnerabilities of JAK2-mutated clones. We identified lysine-specific demethylase KDM4C as a selective genetic dependency that persists upon JAK-inhibitor treatment. Genetic inactivation of KDM4C in human and murine JAK2-mutated cells resulted in loss of cell competition and reduced proliferation. These findings led to reduced disease penetrance and improved survival in xenograft models of human JAK2-mutated cells. KDM4C deleted cells showed alterations in target histone residue methylation and target gene expression, resulting in induction of cellular senescence. In summary, these data establish KDM4C as a specific dependency and therapeutic target in JAK2-mutated cells that is essential for oncogenic signaling and prevents induction of senescence.

Int. J. Mol. Sci. 2022, 23, 2776; https://doi.org/10.3390/ijms23062776

INTRODUCTION

JAK2 is frequently mutated in the aging hematopoietic system and in myeloid cancers [1], such as myeloproliferative neoplasms (MPN). Various signaling pathways are constitutively activated by mutated JAK2-kinase. JAK-inhibitors are well tolerated and highly effective in reducing pro-inflammatory cytokine production and inflammation-related symptoms. So far, their use has been restricted to rather symptomatic approaches as meaningful reduction of disease burden is rarely seen [2]. Of note, this finding is in contrast with other small molecules, as tyrosine kinase inhibitors typically induce regression of the mutated clone. Persistence of JAK2-mutated cells under treatment with JAK-inhibitors raises questions about the role of JAK2 as a driver mutation and suggests selective dependencies that may arise from aberrant cell signaling or gene expression. Genome-wide CRISPR-Cas9-based genetic perturbation screens have identified cell type specific dependencies in various cancers in an unbiased manner [3]. Recently, efforts of the Broad Institute have created databases of large-scale functional in vitro screens, identifying genetic vulnerabilities in human cancer cell lines (DepMap; https://depmap.org/; [4]). However, validation of these targets under conditions of targeted therapies remains a necessity.

MATERIAL AND METHODS

Cell lines and culture conditions

Cell lines were purchased from DSMZ (Braunschweig, Germany). Cells were cultured according to standard protocols and tested negative for mycoplasma. For proliferation assays, the number of cells was counted assuming a relevant difference in means of survival. We used a one-sided t-test at \( a = 0.05 \) and a power of \( >80\% \) with an expected difference in means of 1.75 SD (standard deviations) based on previous experience with xenotransplantation of HEL cells. Equal numbers of 8–12 weeks-old male and female mice were used for experiments in all groups.
Primary patient samples
Primary MPN patient samples and healthy donor controls were obtained after informed consent and according to the Helsinki declaration from the Hematology Tumor Bank Jena and Greifswald, approved by the respective local ethics committees (University Hospitals Jena 4753-04/16 or Greifswald BB199/20).

Western blot
Western Blotting was performed according to standard protocols as previously published [5, 6]. Cell lines and whole bone marrow cells were lysed as described previously. All antibodies are indicated in the Supplement.

Quantitative PCR (Q-PCR)
RNA-preparation, reverse transcription and Q-PCR measurement were performed as described before [7, 8]. Q-PCR primers are provided in the Supplement.

Staining and quantification of SA-β-Galactosidase Activity
Staining of SA-β-galactosidase (SA-β-gal) in cells was carried out in triplicates using Histochimical Staining Kit (CS0303-1KT, Sigma-Aldrich) according to manufacturer’s instructions. Quantitative analyses were performed using an Axioskop 2 mot plus provided with a motorized stage (Zeiss, Oberkochen, Germany) and a CX 9000 digital camera (MicroBrightField Europe, Magdeburg, Germany) and the Stereo Investigator 8.1 software (MicroBrightField). Quantification was performed using the dissector method [9]. Cells of different regions of each sample were selected randomly for software supported counting.

Confocal laserscanning microscopy
Cells were fixed and stained as described before [10]. Anti-H3K27me3 mAb (Abcam, # ab194677) was used at 1:800 dilution and a secondary AF488 conjugated anti-Rabbit mAb (ThermoFisher, #A27034) was applied 1:2000. Microscopic evaluation was performed with the Laser Scanning Microscope LSM 980 Airyscan 2 (Carl Zeiss, Jena, Germany) and ZEN 2009 software (Carl Zeiss).

Virus production
Lentiviral particles containing the pooled sgRNA library (see below) or sgRNAs against luciferase, RPA3 and KDM4C, respectively, were generated and virus titer was assessed as described before [9].

Genome editing by CRISPR/Cas9
Genetic editing by CRISPR/Cas9 was performed as previously described [11, 12] unless otherwise stated. Guide RNAs were designed using the Broad GPP tool (Doench, Nat Biotechnology 2014). For cloning of sgRNA sequences, the improved-scaffold-pu6-sgRNA-EF1Alpha-PURO-T2A-RFP (puUSEPR) vector system [13], with puromycin resistance and RFP selection marker was used. Genetic inactivation by CRISPR/Cas9 was performed as published before [10]. sgRNA sequences are provided in the Supplementary Materials. For negative selection competition assays, transduced cells were mixed with non-transduced cells at 9:1 RFP /RFP+ ratio for applying selection pressure. The percentage of RFP+ was monitored by flow cytometry.

For the genome-scale CRISPR/Cas9 screen the human CRISPR Brunello lentiviral pooled library (Addgene, #73178) cloned into the lentiguide-Puro vector backbone (Addgene, #52963) was used. The library includes 76,441 guide RNAs targeting against 19,114 genes and 1000 control guide RNAs. Sequenced confirmed homogeneous representation with a GINI-index 0.011.

Cells were treated with 200 nM ruxolitinib (Selleckchem, Lot# S137813) and DMSO in quadruplicate. Next generation sequencing was performed on an Illumina NextSeq500 platform (75 bp, single reads) aiming for a minimum of 30M reads per sample. Alignment and statistical analysis of the data was performed using MAGIC (https://sourceforge.net/p/mageck/wiki/Home/) [14], version v0.5.9.3: sgRNA counts were retrieved from raw data (fastq files) via “mageck count”. Beta scores and p values were generated from the counts via “mageck mle” using normalization against controls (control sgRNAs) and CNV-normalization (copy number variations in HEL cells).

RNA sequencing
HEL-Cas9 cells were transduced with Luciferase sgRNA and KDM4C sgRNA lentiviruses analogously to the proliferation assay. 48 h after transduction, selection was initiated by adding puromycin 1.5 μg/ml to each sample and the cells were expanded for another 9 days. Transduced cells were then treated with 200 nM ruxolitinib or DMSO as a control in quadruplicates for 48 h. Total RNA was isolated from 2 × 10^7 cells using innuPrep RNA Mini Kit (Analytik Jena AG, Jena, Germany) according to the manufacturer’s instructions. Library preparation and next generation sequencing were performed by GENEWIZ GmbH (Leipzig, Germany). For quantitative analysis raw reads (fastq files, paired end, strand-specific) were trimmed with trimmomatic (v0.39) (parameters: ILLUMINACLIP: TruSeq3-PE.fa:2:30:10, SLIDINGWINDOW:4:20) [15]. Trimmed reads were aligned to the human genome (hg38) using STAR (v2.7.4a) (default parameters) [16]. Counts were generated from bam-files using Subread featureCounts (v2.0.1) (parameters: -p8) [17]. We used DESeq2 (v1.26.0) used to generate log2 fold changes and p values from the counts [18]. Fgsae (v1.12.0) was used to perform gene set enrichment analysis, using pathways from the GSEA website (https://www.gsea-msigdb.org/gsea/index.jsp) [19, 20]. DESeq2 and fgsae were used in R (v3.6.0).

RESULTS
KDM4C is highly expressed and a specific dependency in JAK2-mutated cells
Most recently, we have used in depth phospho-proteome profiling to define the global signaling landscape downstream of mutated JAK2-kinase and its perturbation by stimulation with its physiologic ligand erythropoietin or JAK-inhibitor treatment. Using RNAi-screens to explore relevant JAK2-targets, we have defined JAK2-selective vulnerabilities related to perturbation of the splicing machinery through oncogenic cell signaling [10]. To identify further vulnerabilities specifically related to mutated JAK2-kinase, we analyzed DepMap CRISPR-Cas9 datasets and identified genes that represent dependencies in JAK2-mutated hematopoietic cell lines compared with non-JAK2-mutated cells (Fig. 1A). We identified a total of 22 genes mainly related to cell signaling, apoptosis and epigenetic regulation. Of note, only 4/22 candidates had been identified as targets of mutated JAK2-kinase in phospho-proteome analysis: BAK1, CNTLN, KDM4C and UHRF2 (Fig. 1B). KDM4C was highly expressed in JAK2-mutated cell lines and was a selective dependency as compared to non-JAK2-mutated cell lines (Fig. 1C). In contrast, other members of the KDM-family could not be identified as specific vulnerabilities for JAK2-mutated cells (Fig. 1D). KDM4C (JMJD2C) is a jumonji domain-containing protein and acts as a trimethylation-specific demethylase that contributes to epigenetic regulation of both oncogene and tumor suppressor genes and is frequently overexpressed in human cancers [21, 22]. In more detail, KDM4C specifically demethylates H3K9me3/me2, H1.4K26me3, and H3K36me3/me2 via a 2-oxoglutarate-dependent dioxygenase reaction. KDM4C activity is required for development of acute myeloid leukemia [21, 22]. However, while genetic inactivation of multiple KDM4 enzymes is detrimental to normal HSCs, selective deletion of KDM4C appears to be dispensable for HSC function [23], which indicates a potential therapeutic window.
Regarding the high dependency of JAK2-mutated cells, we sought to investigate the functional role of KDM4C in vitro and in vivo. KDM4C was highly expressed in peripheral blood granulocytes derived from MPN patients when compared to age-matched healthy donor controls (Fig. 2A). Increased mRNA expression was detectable across all phenotypic subtypes of MPN, including polycythemia vera (PV), essential thrombocythemia (ET), myelofibrosis (MF) or unclassifiable MPN (MPN-U) (Fig. 2B).

As indicated above, Janus kinase inhibitors are well tolerated and highly effective in reducing pro-inflammatory cytokine production and inflammation-related clinical symptoms. However, reduction of disease burden is rarely seen, which highlights the necessity to identify targets that are accessible and effective in combination with JAK-inhibitor treatment. In order to confirm the functional dependency on KDM4C under treatment conditions with the JAK-inhibitor ruxolitinib (RUX), we applied a genome-wide CRISPR-Cas9 screen in the human JAK2-mutated cell line HEL. This cell line was selected for its sensitivity to RUX and ability to be transduced among the JAK2-mutated human cell lines. For the screen, HEL cells were infected with the human CRISPR Brunello lentiviral pooled library including 76,441 guide RNAs targeting against 19,114 genes and 1000 control guide RNAs and treated with 200 nM RUX (or DMSO, as control) for 12 days (Fig. 3A). Alignment and statistical analysis of the data was performed using the MAGeCK and MAGeCK-Flute algorithms as described before [24]. Specifically, MAGeCK was used to align guide sequences from FASTQ files based on the guide-matrix. Subsequently, the MAGeCK-MLE algorithm was used to statistically compare dropout and enrichment of guides between day 0 and day 12 separately for treated (RUX) versus untreated (DMSO) conditions. Here, 749 genes were identified as functional dependencies upon ruxolitinib treatment (dependency score < −0.5). Of note, KDM4C was detected among the ruxolitinib “persistent” dependencies (HEL dependency score −0.92) (Fig. 3B).

Taken together, KDM4C was identified as a selective vulnerability of JAK2-mutated cells also under exposure to JAK-inhibitor treatment.
were (partially) outcompeted by non-transduced cells in JAK2-mutated cell lines (HEL, $p = 0.0017$) compared to non-targeting (sg-Luc) controls (Fig. 4B). Therefore, these in vitro results suggest that KDM4C is relevant under conditions of cell competition in JAK2-mutated cell lines. To confirm these results, we aimed to investigate cell proliferation in different murine and human JAK2-mutated cells following genetic inactivation of KDM4C in vitro (Fig. 4C). Here, loss of cell competition could be attributed to impaired proliferative capacity of KDM4C deficient murine (Ba/F3-V617F-Cas9-Blast; $p = 0.0012$) and human (HEL, $p = 0.002$; SET-2, $p = 0.0081$) JAK2-mutated cells. To validate the functional impact of KDM4C deletion in human JAK2-mutated cells in vivo, we performed CRISPR-Cas9-mediated deletion of KDM4C in HEL cells and assessed disease dynamics after transplantation into humanized NKG mice at different dilutions (Fig. 4D). Inactivation of KDM4C reduced disease activity as indicated by reduced spleen size of recipient mice ($p = 0.0188$ and $p = 0.0227$, respectively; Fig. 4E). Moreover, deletion of KDM4C delayed disease progression at different dilutions. When injecting 30,000 transduced cells, disease penetrance was reduced in KDM4C-depleted cells (gLuc: 40%; KDM4Cg4: 20%; KDM4Cg5: 0%; Fig. 4F, upper panel). At higher counts of 60,000 transduced HEL cells per animal, overall survival was significantly improved (median survival of gLuc: 48 days; sgKDM4C: not reached; $p = 0.0025$) (Fig. 4F, lower panel).

**Loss of KDM4C results in altered methylation of target histones and induction of senescence**

Genetic inactivation of the lysine-specific demethylase KDM4C resulted in increased methylation of H3K36, which is the specific residue targeted by KDM4C. This effect was detected by Western blotting (Fig. 5A) and immunofluorescence staining (Fig. 5B) in murine (Ba/F3-VF) and human (SET-2, HEL) JAK-mutated cells.
H3K36me3 can mediate multiple transcriptional-related events, such as the regulation of transcriptional activity, is associated with both facultative and constitutive heterochromatin and has been linked to cellular processes involved in senescence [25]. When examining the transcriptome of KDM4C-depleted JAK2-mutated HEL cells, we found increased expression of CDKN1A (p21), IL1beta, BCL2, and downregulation of THS81 or MTOR. Consistently, gene set enrichment analysis (GSEA) revealed a strong induction of gene sets related to cellular senescence (NES = 1.55; p = 0.0327; Fig. 5C, D). Furthermore, gene set enrichment analysis (GSEA) demonstrated enrichment of PI3K/AKT/MTOR pathway and NF-κB signaling, whereas genes encoding transcription initiation and genotoxicity pathways appeared repressed.

For functional validation, genetic deletion of KDM4C by CRISPR-Cas9 was induced and compared to daunorubicin treatment as a positive control to induce cellular senescence. Here, we found induction of H3K36me, H3K9me3 and p21 (data not shown) along with significant increase in SA-βGal staining in JAK2-mutated HEL cells in vitro (Fig. 5E, F). These findings indicate that inactivation of KDM4C reduces cell competition and proliferative capacity and induces cellular senescence in JAK2-mutated cells.

**DISCUSSION**

In summary, we have identified lysine-specific demethylase KDM4C as a downstream effector of mutated and constitutively active JAK2 kinase. KDM4C represents a functional vulnerability in JAK2-mutated cells and its inactivation remains detrimental upon co-treatment with JAK1/2-inhibitors. Most recently, KDM4C has been described as a regulator of stemness [26–29], cancer cell resistance [30, 31] and cancer progression [32, 33] in various cancer models and KDM4C germline variants may increase multi-cancer vulnerability through dysregulation of target histone methylation [34]. In murine models of acute myeloid leukemia (AML) driven by MLL-rearrangements, genetic inactivation of all Kdm4 family members blunted leukemia development while inactivation of Kdm4c alone showed minor effects regarding proliferation of leukemic cells [21]. In contrast, KDM4C was shown to regulate ALKBH5 expression in human AML cells by increasing chromatin accessibility to MYC and Pol II and to maintain KDM4C-ALKBH5-AXL signaling [29]. Here, inactivation of the KDM4C-ALKBH5 axis resulted in reduced proliferation, impaired colony formation and loss of leukemia stem cells. The role of KDM4-family proteins has been also investigated in detail in hematopoietic stem cells and models of acute myeloid leukemia (AML). Deletion of Kdm4 family members (Kdm4a/b/c) in murine hematopoietic stem cells resulted in accumulation of H3K9me3 on transcription start sites, transcriptional silencing and loss of stem cell self-renewal [23]. In contrast, inactivation of Kdm4c alone did not reveal deleterious effects in normal HSCs, indicating a potential therapeutic window for defining it as a target in myeloid neoplasms. Our data provides first evidence that JAK2-mutated cells may show an even greater dependency than other myeloid leukemia cell lines. Moreover, this vulnerability is

---

**Fig. 5** Loss of KDM4C leads to differential methylation of target histones and induction of cellular senescence. A Western blot analysis in BaF3/JAK2-V617F_Cas9_Blast, SET-2_Cas9_Blast and HEL_Cas9_Blast cells following CRISPR-Cas9 knockout using KDM4C specific sgRNAs (KDM4C-sg2, -sg3) or non-targeting control (LUC). B Immunofluorescence analysis of H3K36me3 and Hoechst-staining in SET-2_Cas9_Blast and HEL_Cas9_Blast cells following CRISPR-Cas9 knockout using KDM4C specific sgRNAs (KDM4C-sg2, -sg3) or non-targeting control (LUC). C GSEA showing enrichment of genes related to senescence and autophagy in cancer. Plotted are normalized enrichment scores (NES). D Heat map of differentially expressed genes in KDM4C deleted HEL cells versus non-targeting control (LUC); n = 4. Red zones represent higher gene expression (upregulation), blue zones represent lower gene expression (downregulation). E Representative cytopsins of SA-beta-Gal-staining of KDM4C depleted HEL cells compared to non-targeting control (sgLUC). Cells exposed to daunorubicin (Dauno) serve as positive control for SA-beta-Gal staining. F Bar graph depicting quantification of SA-beta-Gal staining as depicted in E. n = 3 independent replicates, two-tailed t-test.
uncoupled from JAK-inhibitor treatment, which is the standard-therapy for relevant subgroups of JAK2-mutated cancers. Given the fact that KDM4C was dispensable for normal hematopoietic stem- and progenitor cells, targeting this enzyme may represent a strategy to selectively target JAK2-mutated cells in myeloproliferative neoplasms.

Induction of cellular senescence has been linked to epigenetic modulator genes and specifically lysine-specific demethylases such as LSD1 and KDM4C. In melanoma cells, enforced expression of KDM4C promoted melanomagenesis through altered methylation of relevant target histone residues [35]. Targeting relevant demethylases such as LSD1 or KDM4C had deleterious effects on melanoma growth by inducing senescence. In JAK2-mutated cells, inactivation of KDM4C resulted in reduced proliferation in vitro, increased methylation of target histone residues (H3K36, H3K9) and induction of senescence. This is of major interest, as the interplay between inflammation and induction of senescence in MPN is so far not well understood. Aging of the microenvironment through MPN induced chronic inflammation may foster induction of senescence in normal, non-mutated HSPCs or niche cells such as mesenchymal stroma cells (MSC). Conversely, senescent cells may in turn contribute to the maintenance of chronic inflammation through the senescence-associated secretory phenotype (SASP). Further studies exploring the long-term effects of senescent JAK2-mutated cells in this therapeutic context are clearly needed. Finally, development of specific pharmacologic KDM4C inhibitors will allow pre-clinical validation of demethylases as relevant clinical targets in JAK2-mutated cancers.

DATA AVAILABILITY
RNA-seq data have been deposited in the Gene expression Omnibus database with accession number GSE203060. CRISPR screen data has been deposited to the Gene expression Omnibus database with the accession code GSE203059.

REFERENCES
1. Perner F, Perner C, Ernst T, Heidel FH. Roles of JAK2 in aging, inflammation, hematopoiesis and malignant transformation. Cells. 2019;8:854.
2. Neuringer M, Radich J, Burn TC, Huber R, Paranganama D, Verstovsek S. The effect of long-term ruxolitinib treatment on JAK2p.V617F allele burden in patients with myelofibrosis. Blood. 2015;126:1551–4.
3. Hart T, Chandrahshekar M, Aregger M, Steinhart Z, Brown KR, MacLeod G, et al. High-resolution CRISPR screens reveal fitness genes and genotype-specific cancer liabilities. Cell. 2015;163:1515–26.
4. Meyer RM, Bryan JG, McFarland JM, Zaleski C, Jha S, et al. STAR: ultrafast and memory-efficient general purpose program for RNA-seq data with DESeq2. Genome Biol. 2014;15:554.
5. Bolger AM, Lohse M, Usadel B. Trimmmomatic: a flexible trimmer for Illumina sequence data. Bioinformatics. 2014;30:2114–20.
6. Zielinski D, Davis CA, Schlegl F, Drenkow J, Zaleski C, Jha S, et al. STAR: ultrafast universal RNA-seq aligner. Bioinformatics. 2013;29:15–21.
7. Liao Y, Smyth GK, Shi W, featureCounts: an efficient general purpose program for assigning sequence reads to genomic features. Bioinformatics. 2014;30:923–30.
8. Love MI, Huber W, Anders S. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol. 2014;15:550.
9. Subramanian A, Tamayo P, Mootha VK, Mukherjee S, Ebert BL, Gillette MA, et al. Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. Proc Natl Acad Sci USA. 2005;102:15454–50.
10. Mootha VK, Lindgren CM, Eriksson KF, Subramanian A, Sihag S, Lehár J, et al. PGC-1alpha-responsive genes involved in oxidative phosphorylation are coordinately downregulated in human diabetes. Nat Genet. 2003;34:267–73.
11. Agger K, Miyagi S, Pedersen KT, Kooistra SM, Johansen JV, Helin K. Jmjd2/Kdm4 demethylases are required for expression of β3ra and survival of acute myeloid leukaemia cells. Genes Dev. 2016;30:1278–88.
12. Cheung N, Fung TK, Zeisig BB, Holmes K, Rane JK, Mowen KA, et al. Targeting aberrant epigenetic networks mediated by PRMT1 and KDM4C in acute myeloid leukemia. Cancer Cell. 2016;29:32–48.
13. Agger K, Nishimura K, Miyagi S, Messing JE, Rasmussen KD, Helin K. The KDM4/JMJD2 histone demethylases are required for hematopoietic stem cell maintenance. Blood. 2019;134:1154–8.
14. Wang B, Wang M, Zhang W, Xiao T, Chen CH, Wu A, et al. Integrative analysis of pooled CRISPR genetic screens using MAGeCKFlute. Nat Protoc. 2019;14:756–80.
15. Ernst P, Heidel FH. Molecular mechanisms of senescence and implications for the treatment of myeloid malignancies. Cancers. 2021;13:612.
16. Chen GQ, Ye P, Ling RS, Feng Z, Zhu XS, Chen L, et al. Histone demethylase KDM4C is required for ovarian cancer stem cell maintenance. Stem Cells Int. 2020;2020:8860185.
17. Lang T, Xu J, Zhou L, Zhang Z, Ma X, Gu J, et al. Disruption of KDM4C-ALDH1A3 feed-forward loop inhibits stemness, tumorigenesis and chromoresistance of gastric cancer stem cells. Signal Transduct Target Ther. 2021;6:336.
18. Massett ME, Monaghan L, Patterson S, Mannion N, Bunschoten RP, Hoose A, et al. A KDM4A-PAF1-mediated epigenomic network is essential for acute myeloid leukaemia cell senescence and survival. Cell Death Dis. 2021;12:573.
19. Wang J, Li Y, Wang P, Han G, Zhang T, Chang J, et al. Leukemogenic chromatin alterations promote AML leukaemia stem cells via a KDM4C-ALKBH5-AXL signaling axis. Cell Stem Cell. 2020;27:81–97.e88
20. Gao Y, Liu J, Liu Y, Peng Y, Tuan B, Xin Y, et al. UHRF1 promotes androgen receptor-regulated CD6 transcription and anti-androgen receptor drug resistance in prostate cancer through KDM4C-Mediated chromatin modifications. Cancer Lett. 2021;520:172–83.
21. Jie X, Fong WP, Zhou R, Zhao Y, Zhao Y, Meng R, et al. USP9X-mediated KDM4C deubiquitination promotes lung cancer radiosensitivity by epigenetically inducing TGF-beta2 transcription. Cell Death Differ. 2021;28:2099–111.
22. Lee DH, Kim GW, Yoo J, Lee SW, Jeon YH, Kim SY, et al. Histone demethylase KDM4C controls tumorigenesis of glioblastoma by epigenetically regulating p53 and c-Myc. Cell Death Dis. 2021;12:89.
23. Shao N, Cheng J, Huang H, Gong X, Lu Y, Idris M, et al. GASC1 promotes hepato-cellular carcinoma progression by inhibiting the degradation of ROCK2. Cell Death Dis. 2021;12:253.
24. Katanen R, Donner I, Raisanen M, Berta D, Kuusmanen A, Kaasinen E, et al. Novel germline variant in the histone demethylase and transcription regulator KDM4C induces a multi-cancer phenotype. J Med Genet. 2021. https://doi.org/10.1136/jmedgenet-2021-107747. Online ahead of print.
25. Wu Y, Schleich K, Yue B, JS, Lohnies P, Kemper K, et al. Targeting the Senescence-Overriding Cooperative Activity of Structurally Unrelated H3K9 Demethylases in Melanoma. Cancer Cell. 2018;33:322–36.e328.

ACKNOWLEDGEMENTS
The authors thank Nicole Schröder for technical assistance.

Leukemia (2022) 36:1843 – 1849

1848

P. Ernst et al.
AUTHOR CONTRIBUTIONS
PE, TMS, NH, FP, AKJ, TE, CJH, and NTS performed research. FP, MU, CH, JC, and AH provided material, analyzed data and contributed to the writing of the manuscript. FHH designed research and analyzed data. PE and FHH wrote the manuscript.

FUNDING
This work was supported by funding from the “Else Kröner-Fresenius-Stiftung, EKFS”, within the Research Program “Else Kröner-Forschungskolleg AntiAge”, Jena (to PE) and in part by grants of the German Research Council (DFG; HE6233/4-2 to FHH) and the Thuringian state program ProExzellenz (RegenerAging–FSU-I-03/14) of the Thuringian Ministry for Economics, Science and Digital Society (TMWWDG; to FHH). Open Access funding enabled and organized by Projekt DEAL.

COMPETING INTERESTS
FHH received fees for consulting services and research funding from Novartis, Celgene, CTI. AH received research support from Novartis, BMS, Pfizer and Incyte. PE received fees for consulting services from Pfizer. The remaining authors declare no competing interests.

ADDITIONAL INFORMATION
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41375-022-01611-3.