INTRODUCTION

Systemic mastocytosis (SM) is characterized by abnormal growth and accumulation of mast cells (MCs) and their precursors in the bone marrow (BM) and in other organs. On the basis of clinical variables and signs of organ damage the WHO classification defines a spectrum of disease variants, including indolent SM (ISM), smoldering SM (SSM), SM with associated hematologic (non-MC) neoplasm (SM-AHN), aggressive SM (ASM) and MC leukemia (MCL). Clinical features and courses in patients with SM vary greatly. In those with ISM and SSM, the clinical course is usually stable and symptoms mostly relate to MC degranulation and mediator release. In contrast, ASM and MCL are characterized by organ damage resulting from malignant MC infiltration. Whereas patients with ISM are considered to have a normal life expectancy, ASM and MCL are severe, fatal forms of SM with short survival times. Therapeutic options are limited in these patients. Somatic mutations leading to ligand-independent activation of the receptor tyrosine kinase KIT (most frequently, KIT D816V) are considered to promote MC differentiation and survival and thus have an important pathogenetic role in SM. However, KIT D816V is detectable both in ASM and in most patients with ISM and several lines of evidence suggest that KIT D816V alone is not a fully transforming oncoprotein. The investigation of a panel of candidate genes involved in myeloid neoplasms has recently shown that additional mutations in genes other than KIT (TET2, NRAS, ASXL1, DNMT3A, SRSF2, CBL, ASXL1) may indeed be found in advanced SM, including SM-AHN and ASM. Patients with additional mutations in ASXL1, RUNX1 and SRS2 were found to have a significantly shorter overall survival. However, only one study using genome-wide approaches to screen for novel molecular aberrations in advanced SM (ASM or MCL) has been published to date. This prompted us to undertake an integrated molecular genetic study of a MCL case who was referred to our center, with the aim to identify novel, potentially druggable lesions involved in the pathogenesis of advanced SM.

ORIgINAL ARTICLE

SETD2 and histone H3 lysine 36 methylation deficiency in advanced systemic mastocytosis

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The molecular basis of advanced systemic mastocytosis (SM) is not fully understood and despite novel therapies the prognosis remains dismal. Exome sequencing of an index-patient with mast cell leukemia (MCL) uncovered biallelic loss-of-function mutations in the SETD2 histone methyltransferase gene. Copy-neutral loss-of-heterozygosity at 3p21.3 (where SETD2 maps) was subsequently found in SM patients and prompted us to undertake an in-depth analysis of SETD2 copy number, mutation status, transcript expression and methylation levels, as well as functional studies in the HMC-1 cell line and in a validation cohort of 57 additional cases with SM, including MCL, aggressive SM and indolent SM. Reduced or no SETD2 protein expression—and consequently, H3K36 trimethylation—was found in all cases and inversely correlated with disease aggressiveness. Proteasome inhibition rescued SETD2 expression and H3K36 trimethylation and resulted in marked accumulation of ubiquitinated SETD2 in SETD2-deficient patients but not in patients with near-normal SETD2 expression. Bortezomib and, to a lesser extent, AZD1775 alone or in combination with midostaurin induced apoptosis and reduced clonogenic growth of HMC-1 cells and of neoplastic mast cells from advanced SM patients. Our findings may have implications for prognostication of SM patients and for the development of improved treatment approaches in advanced SM.

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MATERIALS AND METHODS

Patient samples and cell lines

The index MCL case and 57 SM patients subsequently included in the validation cohort based on sample availability (23 patients with ISM (three with SM-AHN), three with SS, 20 with ASM (three with ASM-AHN) and 11 with MCL (two with MCL-AHN)) were diagnosed and classified according to WHO criteria.2 The clinical characteristics of each patient are shown in Supplementary Table 1. In the index case, CD117+ cells were isolated from BM mononuclear cells (MNCs) collected at the time of diagnosis by immunomagnetic cell separation (Miltenyi Biotech, Cologne, Germany) followed by RNA, DNA and protein extraction with the AllPrep DNA/RNA/Protein Kit (Qiagen, Hilden, Germany). Germline DNA was obtained from saliva. In the validation cohort, DNA, RNA and proteins were obtained from CD117+ cells or MNCs from BM. In this cohort, patients with acute myeloid leukemia-type AHN were excluded. As a result, almost all CD117+ cells isolated from BM samples were CD34-/CD117+ MCs. Investigators performing the experiments were blinded with respect to the WHO subtype, clinical features and outcome of the patients. Sample and data collection were approved by the Institutional Review Boards of the S. Orsola-Malpighi Hospital (protocol 112/2014/U/Tess) and the participating institutions. All patients gave written informed consent. The cell line has two subclones, HMC-1 harboring the V560G mutation and HMC-1Δ harboring both the V560G and the D816V mutation. The control cohort included 95 healthy individuals with age ranging from 20 to 95 years.

Whole-exome sequencing (WES) and single-nucleotide polymorphism (SNP)-arrays

For the index case, paired-end DNA libraries were prepared from CD117+ cells and from saliva using the TruSeq DNA Sample preparation kit (Illumina, San Diego, CA, USA). Exon-coding sequences were captured using TruSeq Exome Enrichment Kit (Illumina) and sequenced on an HiSeq 1000 (Illumina). SNP-array analysis was performed in the index case and in 13 additional cases of the validation cohort (8 MCL, 4 ASM, 1 ISM) using the CytoScan HD array (Affymetrix, Santa Clara, CA, USA). Protocols and data analysis are described in the Supplementary Information.

Loss-of-heterozygosity (LOH) analysis at 3p21.3

Genotyping of nine single-nucleotide polymorphisms with minor allele frequency > 40% in the minimal common region of LOH (rs117202139, rs2305634, rs2305635, rs1079276, rs2278963, rs6767907, rs4082155, rs6780013, rs1531875) was performed by direct sequencing as detailed in the Supplementary Information.

SETD2 mutation screening

SETD2 mutations identified by WES in the index case were validated by Sanger sequencing at the DNA and mRNA level using primers shown in the Supplementary Information. Screening of SETD2 coding and promoter sequence in the HMC-1 cell line and in the validation cohort was performed by Sanger sequencing or high throughput sequencing as described in the Supplementary Information.

Quantitative reverse transcription (RT)-PCR for SETD2 expression

In the HMC-1 cell line and in the S7 samples of the validation cohort, SETD2 expression was assessed by quantitative RT-PCR. Total RNA (200 ng) was reverse transcribed to cDNA with the High-Capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific, Waltham, MA, USA). Assays were performed in triplicate on the ABI 7900HT system (Thermo Fisher Scientific) using pre-designed TaqMan Gene Expression Assays (Thermo Fisher Scientific) for SETD2 (Hs01014784_m1) and TBP (Hs00427621_m1) as control gene. SETD2 mRNA levels were quantified using the Comparative Ct method, using a pool of 10 healthy donors of various ages as calibrator.

Methylation analysis of the SETD2 promoter

Methylation status at the SETD2 promoter was assessed in the validation cohort using EpiTect Methyl II PCR Assays (EPHS110216-1A; Qiagen) on an ABI PRISM 7900HT, according to manufacturer’s instructions. A methylation sensitive (EPHS115450-1A) and methylation dependent (EPHS115451-1A) control assay were also run for all the samples analyzed. A pool of 10 healthy donors of various ages served as a negative control.

Drug treatments

After 24 hours of exposure of HMC-1.1 and HMC-1.2 cells to 1 μM AZD1775 (Selleckchem, Houston, TX, USA), 1 μM midostaurin (Selleckchem), 10 nM bortezomib (Selleckchem), 1 μM AZD1775+1 μM midostaurin and 10 nM bortezomib+1 μM midostaurin, apoptotic cell death was assessed by measuring the uptake of fluoresceinated Annexin V and propidium iodide (PI) (Roche Diagnostics, Mannheim, Germany). A FACsCantoll flow cytometer (Beckton Dickinson, Franklin, NJ, USA) set at 488 nm excitation and 530 nm wavelength bandpass filter for fluorescein detection or 580 nm for PI detection, and a dedicated software (DIVA software; Beckton Dickinson) were used. Drug cytotoxicity was evaluated in clonogenic assays. In HMC-1.1 and HMC-1.2, the reduction of colony (generated in 0.9% methylcellulose supplemented with 30% fetal calf serum) number in the presence of increasing sublethal doses of AZD1775 (0.025–0.1 μM), midostaurin (0.025–0.1 μM each), bortezomib (0.25–1 nM), bortezomib+midostaurin (0.25–1 nM)+(0.025–0.1 μM) was assessed after 10 days of incubation at 37 °C in a fully humidified atmosphere and 5% CO2. AZD1775 (0.025–0.1 μM), midostaurin (0.025–0.1 μM), AZD1775+midostaurin (0.025–0.1 μM each), and bortezomib (0.25–1 nM) were also tested in cells from three advanced SM patients (2 with MCL and one with ASM). Cells from two healthy donors were treated for comparison. Nonlinear regression analyses performed using the CompuSyn software19 (CompoSyn, Inc; Paramus, NJ, USA) were used to calculate the efficacy of different drugs alone or in combination in cell lines and in primary patient cells. Combination indexes were calculated using the same software.

Co-immunoprecipitation/immunoblotting and western blotting analyses

Co-immunoprecipitation/immunoblotting and western blotting were performed as described in the Supplementary Information using the following antibodies: anti-SETD2 (Abnova, Taipei, Taiwan), anti-H3K36Me3 (Merck Millipore, Darmstadt, Germany), anti-RNAPol II (Merck Millipore), anti-hnRNP (Cell Signaling Technology, Danvers, MA, USA) and anti-ubiquitin (Bethyl Laboratories, Montgomery, TX, USA). Beta-actin (Santa Cruz Biotechnology, Dallas, TX, USA) and anti-hnRNP (Merck Millipore, Darmstadt, Germany), anti-RNAPol II (Merck Millipore), anti-RNAPol I (Merck Millipore), and anti-SUMO (Cell Signaling Technology, Danvers, MA, USA) were used as loading controls. Immunoreactive proteins were visualized by probing with horseradish peroxidase-conjugated secondary antibodies and then by enhanced chemiluminescence (ECL; Thermo Fisher Scientific). Immunoblots were scanned and band intensities were quantified with the ImageJ software. Signal intensities in single blots obtained from three individual experiments were averaged, normalized to those of the loading controls and expressed as a fraction of the intensities, set to 1, obtained in a pool of 10 healthy donors of various ages used as a control.

Immunochemistry (IHC)

IHC was performed on sections from paraffin-embedded trephine 8 BM biopsies of selected 5 cases. IHC Staining with anti-SETD2 antibody (HPA042451, Sigma-Aldrich, Milan, Italy) was performed using the Ventana BenchMark Ultra automated IHC slide staining system (Ventana Medical Systems, Tucson, AZ, USA). Tissues from normal testis and from a testicular seminoma were used as external controls and to determine the antibody dilution giving the best staining results. Antibody reactivity was evaluated with diaminobenzidine. Slides were counterstained with hematoxylin.

Statistical analysis

Analysis of variance followed by Tukey post-hoc test was applied in order to detect differences in SETD2 and H3K36Me3 expression between groups. Spearman correlation index was calculated between SETD2 and H3K36Me3 level and disease subtype. Differences in the distribution of continuous and categorical variables between SETD2 > 0.5 and < 0.5 subgroups were assessed by means of non-parametric tests—the Mann–Whitney U-test and the Fisher’s exact test, respectively. There was no adjustment for multiple testing as all analyses were explorative. Survival analysis was considered from the date of diagnosis to the date of death or last contact. Survival curves were prepared by the Kaplan and Meier method and statistical significance of differences in survival was calculated by the log-rank test.
rank test. Analyses were carried out by SPSS (IBM Corp, Chicago, IL, USA) and R. All P-values were two-sided and those less than 0.05 were considered statistically significant.

RESULTS
WES of the index MCL case identifies loss of function mutations in SETD2
Among the mutated genes found in neoplastic MCs but not in saliva, SETD2 stood out among others because of its known role as a tumor suppressor and its involvement in various solid tumors and in acute leukemias. Notably, two distinct putatively inactivating heterozygous mutations were identified in the patient, a nonsense mutation in exon 15 (NM_014159:c.G6753T:p.Glu2234Ter) and a frameshift insertion in exon 20 (NM_014159:c.7595_7596insC:p.Gly2515ArgfsTer5) (Figures 1a and b). Cloning of a cDNA fragment encompassing both mutant residues showed that the mutations were biallelic. The first mutation was predicted to result in a truncated protein lacking 330 amino acids at the C-terminal, whereas the second mutation was predicted to result in a truncated protein lacking 46 amino acids at the C-terminal. Western blotting, however, showed that only the SETD2-truncated isoform resulting from the nonsense mutation was expressed in neoplastic MCs (Figure 1c). This truncated protein (tSETD2) lacked the highly conserved SRI domain (Figure 1b). The normal SETD2 protein is responsible for trimethylation of lysine 36 of histone H3 (H3K36Me3). Co-immunoprecipitation showed that tSETD2 lost the ability to bind the H3 histone and, most importantly, western blotting showed that H3K36Me3 was abrogated (Figure 1c). The SRI domain is also known to mediate SETD2 interaction with the phosphorylated C-terminal domain of the RNA polymerase II large subunit and with the heterogeneous nuclear ribonucleoprotein-L, thus coupling H3K36Me3 with transcription and splicing. Co-immunoprecipitation confirmed that tSETD2 did not bind either protein (Figure 1c), further underpinning the loss of function nature of the SETD2 mutations found in the index case.

LOH at the SETD2 locus is detectable in SM patients
The SETD2 gene maps to chromosome 3p21, where allelic loss due to deletions or copy neutral (CN)-LOH has frequently been observed in kidney, lung, breast and many other cancer samples and cell lines. We thus examined the copy number status and searched for LOH at 3p21 in 13 cases of the validation cohort (eight MCL, four ASM, one ISM) for whom SNP-array data were available. CN-LOH of a small region ranging from 613 kb to 2.02 Mb at 3p21.3 was found in 10/13 cases (7 MCL, 2 ASM, 1 ISM). The minimal common region (MCR) of LOH included SETD2 and nine additional genes (CCDC12, NBEAL2, NRADDP, KIF9-AS1, KIF9, KLH18, PTNP23, SCAP, ELP6) (Supplementary Figure 1). We thus extended the screen for LOH specifically at 3p21.3 in the remaining patients of the validation cohort. Genotyping of nine SNPs with minor allele frequency >40% in the MCR was suggestive of LOH in 17 additional cases. Thus, overall, LOH was

![Figure 1](image_url)

**Figure 1.** SETD2 loss of function mutations in the index MCL case (MCL1). (a) Sanger sequencing chromatograms with the frameshift (top) and nonsense (bottom) mutations identified by whole-exome sequencing. (b) Localization of the mutations with respect to the key functional domains of the SETD2 protein. The SRI domain is necessary for histone H3 lysine 36 trimethylation (H3K36Me3) and mediates SETD2 interaction with the phosphorylated C-terminal domain of the RNA polymerase II large subunit (RNA pol II) and with the heterogeneous nuclear ribonucleoprotein-L (HnRNP), thus coupling H3K36Me3 with transcription elongation and splicing. (c) from top to bottom: western blotting (WB) showing the truncated SETD2 (tSETD2) protein as compared to full-length SETD2 detectable in a pool of proteins from mononuclear cells of healthy donors; co-immunoprecipitation experiments performed by using: an anti-RNA pol II, an anti-HnRNP and an anti-histone H3, respectively, to isolate the proteins of interest and then an anti-SETD2 as primary antibody to label the PVDF membrane on which the immunoprecipitates were transferred; WB for H3K36Me3. Histone H3 and actin were used as loading controls.
SETD2 and H3K36Me3 deficiency are recurrent events in SM and cluster in advanced disease.

To investigate the frequency of SETD2 inactivation in the validation cohort, we used western blotting to assess H3K36Me3 levels as a surrogate marker of loss of SETD2 function and to screen for the expression of SETD2 full-length vs truncated isoforms. No evidence of abnormal isoforms was found. However, the SETD2 protein turned out to be reduced or not at all expressed in the great majority of patients examined (Figure 2a). H3K36Me3 paralleled SETD2 expression and was reduced or absent accordingly (Figure 2a). In one MCL, one ASM and two ISM patients, SETD2 and H3K36Me3 levels were assessed in total MNC fractions obtained from BM and in purified CD117+ cells in parallel and no differences were observed. To confirm western blotting results, IHC with an anti-SETD2 antibody was performed on BM trephine biopsies of patients displaying various SETD2 protein levels. The intensity of nuclear staining and the percentage of immunostained atypical MCs varied among the cases and were consistent with SETD2 protein expression as assessed by western blotting (Supplementary Figure 2). Internal positive controls (BM myeloid genes (for example, SF3B1, SRSF2, TET2, ASXL1 and so on; Supplementary Table 4). In an attempt to explain the nonrandom accumulation of LOH at 3p21.3 and the observed SETD2 protein deficiency, we sequenced the entire SETD2 gene, including the promoter region, in all the patients of the validation cohort. Only one ASM patient (who had no evidence of 3p21.3 LOH) had a missense mutation significantly correlated with each other (Spearman R = 0.91, P < 0.001) and were lower in advanced SM (R = 0.57, P < 0.001) (Figures 2b and c). In particular, patients with MCL and ASM displayed significantly lower levels of SETD2 and H3K36Me3 as compared to ISM (SETD2: P = < 0.001 and P = 0.002, respectively; H3K36Me3: P < 0.001 and P = 0.004, respectively). There seemed to be no differences in SETD2 expression between advanced SM patients with (n = 8) or without (n = 26) AHN. When the distribution of SETD2 protein levels within the ASM and ISM subgroups was examined, no significant differences regarding baseline clinical variables or survival emerged between cases with SETD2 < 0.5 and > 0.5 (data not shown). However, when patients with SETD2 > 0.5 and < 0.5 were compared regardless of disease subtype, lower SETD2 expression was found to be associated with older age (P = 0.02), higher serum tryptase levels (P = 0.03), lower hemoglobin (P = 0.03) and platelet count (P = 0.005). In addition, patients with lower SETD2 levels were more likely to have splenomegaly (P = 0.04) and less likely to have skin lesions (P = 0.03). There was also a trend towards shorter overall survival (P = 0.06) (data not shown). There seemed to be no preferential association between SETD2 loss of function and the presence or absence of specific additional mutations or mutation patterns in myeloid genes (for example, SF3B1, SRSF2, TET2, ASXL1 and so on; Supplementary Table 4).

Figure 2. SETD2 protein and H3K36Me3 deficiency in SM. (a) representative western blot results for SETD2 protein and H3K36Me3 levels in SM patients as compared to a pool of healthy donors (HDs). One of three independent experiments is shown. (b and c) Box and whiskers plots of SETD2 and H3K36Me3 level estimates obtained by densitometric analysis of western blots. Median, interquartile range, minimum, maximum and outliers are indicated. SETD2 and H3K36Me3 signal intensities in single blots obtained from three individual experiments were normalized to those of beta-actin and H3 histone, respectively, and averaged. Normalized SETD2 and H3K36Me3 levels calculated in SM patients were then expressed in comparison to normalized SETD2 and H3K36Me3 levels detected in a pool of HDs, conventionally set to 1. The asterisks indicate that MCL and ASM had significantly lower levels of SETD2 protein (P < 0.001 and P = 0.002, respectively) and H3K36Me3 (P < 0.001 and P = 0.004, respectively) as compared to ISM.
(c.1351A>G; p.Thr451Ala) not reported as a SNP (Supplementary Table 5). The mutation was not found either in the Catalogue of Somatic Mutations in Cancer or in ClinVar databases and was predicted to be benign/tolerated by Mutation Taster and Polyphen-2. Thus, despite LOH, the classical two-hit hypothesis predicted to be benign/tolerated by Mutation Taster and Somatic Mutations in Cancer or in ClinVar databases and was not found either in the Catalogue of Somatic Mutations in Cancer or in ClinVar databases.

To rule out the possibility that SETD2 loss of function might be another age-associated alteration in an epigenetic regulator, 80 healthy older adults with an age ranging from 65 to 95 years were screened for SETD2 mutations and for alterations in SETD2 protein expression or H3K36Me3. No sequence variants other than the Pro1962Leu and the Asn1155Asn SNPs were detected. All the older adults did express SETD2; and densitometric analysis of western blots (done using a pool of 10 younger adults with an age ranging from 20 to 30 years as a calibrator, after having excluded individual variations of expression) showed comparable SETD2 and H3K36Me3 levels among individual older adults as well as between older and younger adults (Supplementary Table 6).

We next wondered whether the HMC-1 cell line, established from a patient with MCL, also exhibits SETD2 and H3K36Me3 deficiency. Indeed, western blotting revealed no SETD2 protein and no H3K36Me3 in either subclone (Figure 3a). Sequencing ruled out the presence of SETD2 gene mutations. Similarly to the patients of the validation cohort, both HMC-1.1 and HMC-1.2 cells were found to have no significant reduction in SETD2 transcript levels as compared to SETD2 wild-type cell lines and the gene promoter was found to have very low methylation levels (4.08% and 1.08% in HMC-1.1 and HMC-1.2, respectively). This suggested that similar mechanism(s) acting at the translational or post-translational level underlie SETD2 deficiency in advanced SM patients and in HMC-1 cells.

SETD2 deficiency results from enhanced proteasomal degradation

In HMC-1 cells, incubation with the proteasome inhibitor bortezomib (10 nM for 24 h) rescued SETD2 protein expression and restored H3K36Me3 levels (Figure 3b), indicating that a functional SETD2 protein is regularly translated in neoplastic MCs, and that failure to detect it by western blotting is rather due to altered turnover and proteasomal degradation. Co-immunoprecipitation with an anti-SETD2 antibody after bortezomib treatment showed that blockage of proteasome-mediated degradation results in accumulation of ubiquitinated and SUMOylated SETD2 (Figure 4a). We therefore speculated that reduced or undetectable SETD2 expression in advanced SM patients might be accounted for by hyper-ubiquitination and -SUMOylation leading to proteasomal degradation. We thus compared the effects of proteasome inhibition in malignant MCs from advanced SM patients displaying no SETD2 protein and from ISM patients with near-normal SETD2 levels. In contrast to SETD2-deficient patients, where results were superimposable to those obtained in HMC-1 cells, patients with near-normal SETD2 expression had similar levels of ubiquitinated and SUMOylated SETD2 protein (and similar H3K36Me3 levels) before and after the inhibition of proteasome-mediated degradation (Figure 4b).

Bortezomib and the combination of Midostaurin+AZD1775 reduce colony growth and induce apoptosis in HMC-1 cells

Given the ability of bortezomib to rescue SETD2 expression and H3K36Me3, we tested whether proteasome inhibition might be effective in vitro in HMC-1 cells. Bortezomib as single agent strikingly inhibited colony formation (LD50 in HMC-1.1 = 0.173 nM; LD50 in HMC-1.2 = 0.226 nM) and induced apoptosis in these cells (Figures 5f and g). Combination with D816V KIT tyrosine kinase inhibition by midostaurin did not increase significantly bortezomib efficacy (Supplementary Figure 3).

Recently, a synthetic lethality interaction between H3K36Me3 deficiency and enhancement of replicative stress via Wee1 G2 checkpoint Serine/Threonine protein kinase inhibition has been uncovered. As a consequence, inhibition of Wee1 with AZD1775 was found to kill SETD2/H3K36Me3-deficient cells. We thus decided to explore also the in vitro efficacy of a strategy using AZD1775 alone or in combination with midostaurin in HMC-1 cells. AZD1775+midostaurin combination suppressed colony growth with a LD50 of 0.036 μM and 0.038 μM for HMC-1.1 and -1.2, respectively (Figure 5b; results of AZD1775 and midostaurin as single agents shown in Figures 5c and d; summary of LD50 values in Figure 5e). When compared to either agent alone, AZD1775 +midostaurin combination displayed a strong synergistic effect in apoptotic cell death induction in HMC-1.1 cells, whereas only an additive effect was observed in HMC-1.2 cells (Supplementary Table 7).

Bortezomib and the combination of AZD1775+midostaurin reduce the clonogenic potential of SETD2/H3K36Me3-deficient neoplastic MCs

We next assessed the effects of bortezomib, AZD1775, midostaurin and AZD1775+midostaurin combination on the clonogenic potential of neoplastic MCs from two patients with MCL with no SETD2 expression (MCL3 and MCL7) and one patient with ASM (ASM18) with very low SETD2 expression (20% as compared to a pool of healthy donors). Growth inhibition curves and LD50 values
are shown in Figure 6. Bortezomib induced a marked dose-dependent reduction in colony formation with LD50 ranging from 0.105 to 0.295 nM (Figure 6a). The combination of AZD1775 + midostaurin appeared to be slightly superior to the single agent strategy in MCL3 and ASM18, with LD 50 values of 0.054 and 0.058 μM, respectively, but not in MCL7 where AZD1775 as single agent showed the lowest LD50 (0.026 μM for AZD1775 alone as against 0.048 μM for the combination; Figures 6b–e).

DISCUSSION

In recent years, deregulation of epigenetic modifiers by chromosomal translocations, genomic loss and/or point mutations has turned out to be a recurrent event in cancer evolution. Loss of function of the SETD2 tumor suppressor gene, which encodes the only histone methyltransferase that can catalyze H3K36Me3, was first identified in clear cell renal cell carcinomas, and subsequently, albeit at lower frequency, in several other solid tumors including lung, breast, skin and brain cancers. More recently, SETD2 loss of function and global loss of H3K36Me3 have been observed in acute lymphoid and myeloid leukemias. In renal cell carcinomas, SETD2 loss of function mainly results from large monoallelic deletions or CN-LOH at chromosome 3p, followed by mutation of the remaining allele. In acute leukemias, it rather derives from biallelic missense or truncating mutations. Here, we report for the first time that SETD2 and H3K36Me3 deficiency are recurrent events in patients with advanced SM. Of note, our findings demonstrate that SETD2 loss of function may also occur at the post-translational level, in the absence of mutations or structural aberrations. Biallelic inactivating mutations were indeed found only in the index case. LOH of a relatively small region at 3p21.3 was detected in almost 50% of the patients of the validation cohort, with higher frequency in advanced SM as compared to ISM. However, LOH was neither associated with coding, splicing or promoter mutations, nor with epigenetic silencing, and no significant differences in SETD2 mRNA levels were found when comparing SM patients with or without LOH and healthy donors. Nonetheless, the great majority of the patients had reduced SETD2 protein expression, albeit to a varying degree, as compared to a pool of healthy donors as assessed by western blotting. As expected, this was mirrored by similarly reduced, or totally abrogated, H3K36Me3 (Figure 2a). IHC in BM biopsies of the same patients further confirmed this observation and showed weak or no nuclear staining for SETD2 in neoplastic MCs (Supplementary Figure 2).

Very low or absent SETD2 protein expression was especially found in cases with advanced SM (Figure 2b), with some exceptions. The most striking outliers were three ISM patients with no SETD2 expression by western blotting and IHC: one of these patients was a severely symptomatic ISM case, with histologically documented gastrointestinal involvement; one had diffuse osteoporosis; the remaining one seemed to have no peculiar clinical features. The small number of patients belonging to each disease subtype limited the possibility to perform statistical comparisons to identify peculiar clinical features or outcome differences between patients with high vs low SETD2 expression within the ASM and ISM groups. When patients with high and low SETD2 levels were compared irrespective of the WHO category, lower SETD2 levels correlated with older age, higher serum tryptase levels, lower hemoglobin and platelet count, splenomegaly and absence of skin lesions. There was also a not statistically significant trend towards shorter overall survival. In
This regard it is noteworthy that in renal cell carcinomas lower SETD2/H3K36Me3 levels, assessed by IHC, correlate with larger tumor burden, more advanced stage of disease, and a worse prognosis. Similarly, in chronic lymphocytic leukemia recent data suggest that SETD2 monoallelic deletions or mutations are associated with inferior progression-free and overall survival. Analysis of a larger series of patients is warranted to fully understand the clinical and prognostic meaning of SETD2 loss of function in SM.

Proteasomal degradation was found to have a major role in the observed lack of SETD2 protein. In the HMC-1 cell line and in neoplastic MCs from advanced SM patients, incubation with the proteasome inhibitor bortezomib rescued SETD2 expression and H3K36Me3. Immunoprecipitation and immunoblotting showed that blockage of proteasomal degradation results in accumulation of ubiquitinated and SUMOylated SETD2 in HMC-1 cells and in primary MCs in advanced SM patients with no SETD2 protein expression but not in ISM patients with near-normal SETD2. This evidence of ‘non-genomic’ loss of function suggests that measuring protein and H3K36Me3 levels by western blotting or IHC is the most informative screening approach and that the frequency of SETD2 involvement in solid tumors or hematologic malignancies might be higher than sequencing and copy number data alone would suggest.

Figure 5. Effects of bortezomib and AZD1775+midostaurin combination in HMC-1 cells. (a and b) reduction of clonogenic growth of HMC-1.1 (red curve) and -1.2 (blue curve) cells, in the presence of increasing doses of bortezomib (0.25–1 nM) and AZD1775+midostaurin (0.025–0.1 μM), respectively. All the clonogenic survival rates are expressed as mean ± standard deviation of counts from three independent experiments. Curves with AZD1775 (0.025–0.1 μM) and midostaurin (0.025–0.1 μM) as single agents are shown in c and d. (e) Comparison of LD50 values for the different strategies. (f and g) Induction of apoptosis in HMC-1.1 and -1.2, respectively. Columns represent the mean of three independent experiments and the bars represent the standard error. ctrl, control; mido, midostaurin.

| DRUGS                  | HMC 1.1   | HMC 1.2   |
|------------------------|-----------|-----------|
| BORTEZOMIB             | 0.173 nM  | 0.226 nM  |
| AZD1775+MIDOSTAURIN    | 0.036 μM  | 0.038 μM  |
| AZD1775                | 0.058 μM  | 0.046 μM  |
| MIDOSTAURIN            | 0.057 μM  | 0.049 μM  |

Proteasomal degradation was found to have a major role in the observed lack of SETD2 protein. In the HMC-1 cell line and in neoplastic MCs from advanced SM patients, incubation with the proteasome inhibitor bortezomib rescued SETD2 expression and H3K36Me3. Immunoprecipitation and immunoblotting showed that blockage of proteasomal degradation results in accumulation of ubiquitinated and SUMOylated SETD2 in HMC-1 cells and in primary MCs in advanced SM patients with no SETD2 protein expression but not in ISM patients with near-normal SETD2. This evidence of ‘non-genomic’ loss of function suggests that measuring protein and H3K36Me3 levels by western blotting or IHC is the most informative screening approach and that the frequency of SETD2 involvement in solid tumors or hematologic malignancies might be higher than sequencing and copy number data alone would suggest.
Patients with evidence of LOH at 3p21.3 did not always harbor lower SETD2 protein levels as compared to patients with no LOH. Thus, the biological significance of the recurrent 3p21.3 LOH events that can be observed in SM remains, at present, unclear. Ligand-independent activation of the KIT pathway, often resulting from the D816V KIT mutation, has a central role in SM pathogenesis. In acute myeloid leukemias, SETD2 has been found to be a critical cooperating event for a variety of major chromosomal or genetic aberrations that are known to be driver mutations in leukemogenesis. In these studies, SETD2 knockdown contributed to leukemia initiation, accelerated progression, enhanced the number and the self-renewal potential of leukemic stem cells and was associated with increased expression of mTOR and JAK-STAT pathway components—which are also important downstream effectors of KIT. It can be hypothesized that reduced SETD2 expression/absence and H3K36Me3 deficiency can cooperate with, and potentiate the effects of KIT constitutive activation to determine the phenotype of advanced SM. Increased DNA damage, reported to occur in SETD2-deficient cells, would then be responsible for the accumulation of additional molecular/cellular defects accounting for the phenotypic and clinical heterogeneity of the disease.

Our findings may have important therapeutic implications. Polychemotherapy, interferon, cladribine or midostaurin are currently the main options in advanced SM, and hematopoietic stem cell transplant is also considered in younger patients. However, treatment results are often unsatisfactory and many patients relapse on therapy or have resistant disease. In addition, in many cases with advanced SM-intensive therapy cannot be applied because of age or poor performance status. As a result, the prognosis of patients with advanced SM is poor and the median survival of patients with MCL is measurable in months. Our study points to bortezomib as a promising novel approach worth exploring clinically for the treatment of advanced SM. Bortezomib had already been reported to counteract the growth of cell lines and primary neoplastic MCs, an effect that had been attributed to the bortezomib-mediated re-expression of the pro-apoptotic factor BIM (suppressed in SM as a result of constitutive activation of the KIT/SCF pathway). However, the concentrations of bortezomib required to suppress proliferation and induce apoptosis were slightly lower than that required to re-express BIM, which had led to the hypothesis that bortezomib might act in part via additional mechanisms. We here show that bortezomib treatment at subnanomolar concentrations promotes SETD2 protein re-expression, reduces clonogenic potential and promotes apoptosis. Whether concomitant KIT D816V inhibition by midostaurin might exert an additive or synergistic effect was also tested—but the striking efficacy of bortezomib as single agent appeared not to be significantly enhanced further (Supplementary Figure 3). In renal cell carcinomas, it has been reported that SETD2-negative cells display elevated DNA damage and that SETD2 depletion abrogates p53 activation.
hypothesized that apoptosis triggered by bortezomib may at least in part result from restoration of a functional p53 checkpoint in a context of DNA damage.

We were also intrigued by the recent report suggesting that H3K36Me3-deficient tumors can be targeted in vitro and in vivo by AZD1775 (a Wee1 kinase inhibitor) through a mechanism of synthetic lethality ultimately resulting in dNTP starvation.34 AZD1775 alone or in combination with other targeted drugs or chemotherapy is currently being assessed in phase 2 studies for several types of solid tumors as well as for acute leukemias and myelodysplasias—which would greatly facilitate its repurposing. This prompted us to explore whether the elimination of H3K36Me3-deficient cells by AZD1775, alone or in combination with KiT D816V inhibition by midostaurin, might prove effective in vitro in our cell line model as well as in neoplastic MCs from patients with advanced SM. AZD1775 and midostaurin showed strong synergy in inducing apoptosis in HMC-1 cells, but only an additive effect in HMC-1.2 cells. Clonogenic growth of both cell line subclones was affected in a dose-dependent manner by AZD1775+midostaurin combination, that was slightly more effective than the single agents alone. In neoplastic MCs from patients with advanced SM, the combination of AZD1775 +midostaurin appeared to be slightly superior to the single agent strategy in two out of three cases, whereas in the remaining case AZD1775 as single agent showed the lowest LD50. Additional testing in a greater number of patient samples will thus be required to better investigate the efficacy of a strategy using AZD1775 alone or in combination.

In conclusion, we show that non-genomic loss-of-function of SETD2 is a recurrent event in advanced SM. SETD2 and H3K36Me3 deficiency might cooperate with the ligand-independent activation of the KiT pathway to enhance leukemic transformation and progression and determine the phenotype of ASM and MCL.

Targeting SETD2 down-modulation itself (with bortezomib or second-generation proteasome inhibitors) and/or SETD2/KiT cooperation are promising therapeutic strategies to improve the outcome of ASM and MCL patients.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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AUTHOR CONTRIBUTIONS

Contribution: SS and GM designed the study, supervised the study, analyzed and interpreted results. MMancini, CDB, MR, MManfrini, MMeggendorfer, RC, VG, MCF, AP, LB, EZ, MD, LR, FM and TH performed experiments, analyzed and interpreted results. CP, LP, RZ, LS, GS, FA, SM, CE, PS, DG, PT, FC and GV provided patient samples and clinical data. MC coordinated the clinical and research activities. All authors contributed to drafting the manuscript and gave final approval for submission.

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