Mechanisms of Pinometostat (EPZ-5676) Treatment Emergent Resistance in MLL Rearranged Leukemia
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Abstract

DOT1L is a protein methyltransferase involved in the development and maintenance of MLL-rearranged (MLL-r) leukemia through its ectopic methylation of histones associated with well characterized leukemic genes. Pinometostat (EPZ-5676), a selective inhibitor of DOT1L, is in clinical development in relapsed/refractory acute leukemia patients harboring rearrangements of the MLL gene. The observation of responses and subsequent relapses in the adult trial treating MLL-r patients motivated preclinical investigations into potential mechanisms of pinometostat treatment emergent resistance (TER) in cell lines confirmed to have MLL-r. TER was achieved in five MLL-r cell lines, KOPN-8, MOLM-13, MV4-11, NOMO-1, and SEM. Two of the cell lines, KOPN-8 and NOMO-1 were thoroughly characterized to understand the mechanisms involved in pinometostat resistance. Unlike many other targeted therapies, resistance does not appear to be achieved through drug-induced selection of mutations of the target itself. Instead, we identified both drug efflux transporter dependent and independent mechanisms of resistance to pinometostat. In KOPN-8 TER cells, increased expression of the drug efflux transporter ABCB1 (P-glycoprotein, MDR1) was the primary mechanism of drug resistance. In contrast, resistance in NOMO-1 cells occurs through a mechanism other than upregulation of a specific efflux pump. RNA-seq analysis performed on both parental and resistant KOPN-8 and NOMO-1 cell lines supported two unique candidate pathway mechanisms that may explain the pinometostat resistance observed in these cell lines. These results are the first demonstration of TER models of the DOT1L inhibitor pinometostat and may provide useful tools for investigating clinical resistance.

Introduction

Rearrangements at the 11q23 locus target the gene MLL (KMT2A), resulting in the oncogenic fusion of the amino terminus of MLL, fused in frame with any of more than 70 fusion partners (1-5). MLL-
rearrangements (MLL-r) occur in 5-10% of acute leukemias of lymphoid, myeloid, or mixed phenotype lineage and are especially common in infant acute leukemias (1, 2, 5). Common MLL oncogenic fusion partners include members of the AF and ENL protein families [1-5]. Translocations of the MLL gene typically involve the loss of the carboxy terminal portion of the MLL protein that contains the SET domain responsible for its methyltransferase activity (1, 2, 5-7). The resulting fusion protein maintains its ability to bind a set of developmentally important genes such as HOXA9 and MEIS1 while gaining a fusion partner capable of recruiting DOT1L to a complex that promotes transcriptional elongation (8-17). DOT1L is a protein methyltransferase (PMT) responsible for catalyzing all three methylation states (mono, di, and tri) of lysine 79 on histone H3 (H3K79) (18, 19). The fusion present in MLL-r patients leads to the aberrant localization of the DOT1L-containing multi-subunit complex within chromatin followed by the ectopic methylation of H3K79 and enhanced gene expression of leukemogenic genes (9-11, 13, 20). Recent studies involving genetic knockdown and pharmacologic inhibition of DOT1L support the hypothesis that DOT1L drives leukemogenesis through overexpression of MLL-r target genes, including HOXA9 and MEIS1 (10, 11, 21-24).

Pinometostat (EPZ-5676) is a small molecule inhibitor of DOT1L currently in clinical development for both adult and pediatric relapsed/refractory acute leukemia patients harboring MLL-r. Preclinical in vitro and in vivo studies conducted with pinometostat using models of MLL-r demonstrated robust single-agent anti-proliferative activity as a direct consequence of pinometostat’s selective inhibition of H3K79 methylation and resultant transcriptional repression of MLL-r target genes HOXA9 and MEIS1 (25). Early reports from the adult phase 1 clinical trial described responses in MLL-r patients, including complete remissions, with subsequent disease progression which motivated preclinical investigation into mechanisms precipitating pinometostat treatment-induced resistance (26).
Despite advances in understanding the molecular mechanisms of disease and implementation of targeted therapies, a common limiting factor to sustained efficacy in cancer patients is the emergence of resistant clones (27, 28). Resistance has been reported to occur through multiple mechanisms such as transformation of the drug target through genetic variation (indels or single base modifications), overexpression, and alternative signal transduction (28). Knowledge of the mechanisms of drug-induced resistance can lead to insights for combinations and rational small molecule drug design to develop compounds for overcoming drug resistance (29-31). Point mutations within the coding region of the target are a frequent cause for resistance to targeted therapies and have been observed clinically in acute myeloid leukemia (AML) with small molecules such as FMS-like tyrosine kinase-3 (FLT3) inhibitors (32, 33). For example, following treatment with FLT3 inhibitors PKC412, SU5614, and sorafenib, acquired resistance mutations have been identified in tyrosine kinase domains 1 and 2 at residues N676, F691, D835, and Y842 (34). Similar to a mechanism found for FLT3 inhibitors, preclinical studies performed with inhibitors of the PMT Enhancer of zeste homolog 2 (EZH2) demonstrated that one mechanism of drug resistance in this enzyme class is through the acquisition of secondary point mutations; whether such EZH2 resistance-causing mutations are relevant to the clinical situation is not clear at present (35, 36). Following up on this work, we set out to examine whether MLL-r cell line models could acquire pinometostat resistance through continuous treatment at concentrations above the pre-determined 14 day proliferation assay IC90 (37). Resistance to pinometostat in all five cell lines tested emerged following three weeks of continued treatment and was defined by increased growth rates in the presence of inhibitor. In depth analysis of two MLL-r models KOPN-8 (MLL-ENL) and NOMO-1 (MLL-AF9) identified common characteristics between the resistant cell lines, but distinct mechanisms by which they became resistant. These mechanisms included examples of increased expression of drug
efflux transporters and evidence of activation of signaling pathways such as PI3K/AKT and RAS/RAF/MEK/ERK.

Materials and Methods

Cell Culture and Analysis of Cell Proliferation and Viability

Human leukemia cell lines KOPN-8 (ACC 552), NOMO-1 (ACC 542), MOLM-13 (ACC 554), and SEM (ACC 546) were obtained from DSMZ, while MV4-11 (CRL-9591) cells were attained from ATCC, between the years 2009 to 2010. All cell lines were authenticated by STR profiling (KOPN-8 and NOMO-1 cells were authenticated using the ATCC fee for service; MOLM-13, MV4-11 and SEM were authenticated by the vendors and used within 6 months of resuscitation), and grown in vendor recommended media (all cell culture reagents were supplied by ThermoFisher Scientific) in a humidified incubator set to 37°C and 5% CO2. Pinometostat resistant cells were cultured in media supplemented with 1 μM (MV4-11, MOLM-13) or 4.5 μM (KOPN-8, NOMO-1, SEM) of the inhibitor. Every 3-4 days cell number and viability was assessed by trypan blue exclusion staining (Vi-CELL, Beckman Coulter) and growth media and pinometostat replaced, splitting cells back to a density of 2x10^5 cells/mL. Cell splitting ratios were recorded and used to express the total number of viable cells per milliliter. Pinometostat was made by Epizyme (38). Proliferation assays with combination agents valsparse (PSC 833, R&D Systems), trametinib (Selleck Chem), GDC-0994 (Selleck Chem), and dabrafenib (Selleck Chem) were completed with the above conditions (39-42).

ELISA Analysis on Inhibition of H3K79me2

The H3K79me2 ELISA was performed as previously described (25). Briefly, 1-2x10^6 KOPN-8 and NOMO-1 cells were collected after 10 and 28 days of dosing with 4.5 μM pinometostat or DMSO. Histones were extracted as previously described (21). Histones were run in matched H3K79me2 and
total H3 ELISAs. The optical density of the H3K79me2 ELISA was normalized to the total H3 ELISA of each sample.

**Sanger Sequencing**

Cell pellets from the resistant and parental KOPN-8 and NOMO-1 cell lines were provided to Genewiz (South Plainfield, NJ) for endogenous cDNA verification of DOT1L by Sanger sequencing. RNA extraction and purification was performed with the RNeasy Plus Mini Kit (Qiagen, 74134) followed by cDNA library generation with the SuperScript III First-Strand Synthesis Supermix Kit (Invitrogen, 18080-400). A 1% agarose gel was used to resolve PCR products, followed by purification with the Qiaquick Gel Extraction Kit (Qiagen, 28706). The procedures followed for the RNA extraction, cDNA library generation and gel extraction were completed as outlined in the manufacturer’s protocols. BigDye Terminator Cycle Sequencing was used for sequencing and DNASTAR Lasergene12 software for data analysis. The threshold for SNP detection was set to 10%. Sequencing results were compared to the reference sequence NM_032482.2. PCR and sequencing primers can be found in the supplemental methods. DOT1L sequences were deposited in GenBank for KOPN-8 (accession number KX792913) and NOMO-1 (accession number KX792914).

**Quantitative Real-time PCR**

Total RNA was isolated from approximately 3x10^6 cells using the RNeasy Total RNA isolation kit (Qiagen, 74106) according to manufacturer’s instructions. ABCB1 (Hs00184500_m1, ThermoFisher Scientific) mRNA levels were assessed and normalized to 18S (Hs99999901_s1, ThermoFisher Scientific), HOXA9 (Hs00365956_m1, ThermoFisher Scientific) and MEIS1 (Hs00180020_m1, ThermoFisher Scientific) were assessed and normalized to beta-2-microglobulin (B2M) (4333766F, ThermoFisher Scientific) by qRT-PCR as previously described (21).
RNA-seq

Samples were collected after 28 days (KOPN-8) or 53 days (NOMO-1) of 4.5 μM pinometostat treatment for whole transcriptome analysis through RNA-seq. In order to distinguish changes in gene expression due to DOT1L inhibition from acquired resistance, naïve cells were dosed with 4.5 μM pinometostat for 10 days and included in the RNA sequencing analysis as a reference for sensitivity to pinometostat. All samples were then evaluated relative to the vehicle control. Q² Solutions (Morrisville, NC) performed the RNA extraction, amplification, and microarray processing on the samples detailed above. RNA-seq data was deposited in GEO, accession number GSE94849. Detailed library preparation and sequencing methods can be found in the supplemental methods.

H3K79me2 Chromatin Immunoprecipitation Followed by Sequencing (ChIP-seq)

Samples were collected from KOPN-8 cells treated with 4.5 μM pinometostat or DMSO for 10 and 28 days, and from NOMO-1 cells treated for 14 and 53 days. Cells were fixed for ChIP-seq according to the protocol provided by Active Motif as follows. 1/10 volume of formaldehyde solution (11% formaldehyde, 0.1 M NaCl, 1 mM EDTA pH 8, 50 mM HEPES pH 7.9) was added to the media of each flask of cells. Following agitation at room temperature for 15 minutes, fixation was halted with the addition of 1/20 volume 2.5 M glycine solution for 5 minutes. Cells were spun down in a 4° C cooled microcentrifuge at 800 x g for 10 minutes and the supernatant removed. Following centrifugation, cells were washed with 10 mL ice cold PBS-Igepal (1X PBS, 0.5% Igepal CA-630) and supernatant removed. Cells were washed again with 10 mL ice cold PBS-Igepal with the addition of 100 μl PMSF (100 mM in ethanol, final concentration 1 mM). Cell pellets were flash frozen and stored at -80° C.
Fixed samples were provided to Active Motif for chromatin immunoprecipitation with an antibody against H3K79me2 (Abcam, ab3594) and next generation sequencing. Detailed methods can be found in the supplemental methods.

**ABCB1 Transporter Study**

ABCB1 transporter studies were performed at Optivia Biotechnology (Menlo Park, CA) as described in the supplemental methods.

**Western Blot Analysis**

Please see supplemental methods

**Bioinformatics Analysis**

Utilizing different R packages, PCA, hierarchical clustering, differentiation and pathway analyses were performed on the RNA-seq data as described in the supplemental methods.

**Results**

**Development of Pinometostat Resistance in KOPN-8 and NOMO-1 Cell Line Models of MLL-r**

*MLL*-r cell lines KOPN-8 (*MLL-ENL*) and NOMO-1 (*MLL-AF9*) are examples of cellular models that have demonstrated anti-proliferative phenotypes, with 14 day IC_{50} values of 71 and 658 nM respectively, when treated with pinometostat as a single agent (25). We tested the potential for identifying resistance mechanisms using these two sensitive *MLL*-r models by treating each cell line continuously in the presence of 4.5 µM pinometostat until resistance was acquired. The process to achieve pinometostat resistance for both cell lines was completed within approximately three weeks from the initiation of treatment (Fig. 1). We have previously demonstrated approximately seven days of
treatment with pinometostat is required to realize clear anti-proliferative activity in vitro (25). Anti-proliferative activity in the form of cell stasis continues from day 7 through day 20, after which a resistant phenotype emerges. Resistance was defined as when the growth rate of the cells cultured in the presence of pinometostat was similar to the naïve control cell line. Stability of the resistant cell line pools was ascertained by allowing the cells to grow in the presence of pinometostat for an additional three weeks following the establishment of resistance.

Cellular responses to pinometostat in resistant and control pools were examined by a number of pharmacological measures (Table 1) including proliferation rates and the ability to inhibit the DOT1L pharmacodynamic marker H3K79me2. In both the KOPN-8 and NOMO-1 cells tested, there was a hundreds-fold increase in proliferation IC₅₀ values between the control and resistant cell lines. Interestingly, despite similar shifts in proliferation IC₅₀ values, the degree of H3K79me2 inhibition four days post treatment was discordant between the KOPN-8 and NOMO-1 resistant cells. In KOPN-8 cells, much like the proliferation shift, there was an approximate 100-fold increase in IC₅₀ values for inhibition of the H3K79me2 mark between the control and resistant cell lines, from 15 nM to 2 μM, suggesting a significant decrease in target inhibition. In contrast, the NOMO-1 cell lines exhibited a muted shift of a less than 3-fold increase in the H3K79me2 mark IC₅₀ values between the control and resistant cell lines, from 27 nM to 95 nM. The rate at which both of these cell lines became refractory to pinometostat is suggestive of the acquisition of a mutation in the SAM binding domain of DOT1L or increased activity of a drug efflux transporter. The pharmacological measurements in Table 1 potentially support either hypothesis as the mechanism for KOPN-8 resistance, as the shift in cell growth inhibition and H3K79me2 is suggestive of the drug’s decreased ability to interact with DOT1L. The sustained inhibition of H3K79me2 in pinometostat resistant NOMO-1 cells is in contrast to the KOPN-8 results and could indicate alternative mechanisms of pinometostat resistance.
Measures of Pharmacodynamic Markers over Time

To further evaluate the mechanistic differences between pinometostat resistance in the KOPN-8 and NOMO-1 cells, we collected cells at various time points over the course of developing pinometostat resistance. Samples from these time points underwent detailed analyses of global and locus specific H3K79me2 in addition to whole transcriptome analysis through RNA-seq. Assessment of global H3K79me2 from resistant samples continuously maintained in 4.5 µM pinometostat, a dose higher than the IC50 for global reduction of H3K79me2, demonstrated that relative to their DMSO (vehicle) controls, both the resistant KOPN-8 and NOMO-1 cells exhibited maximal and sustained inhibition of the H3K79me2 mark across both time points evaluated, at approximately 10% of the vehicle control (Fig. 2A). RNA-seq analysis was performed in an attempt to identify signaling pathways altered by the development of pinometostat TER. Using these data to assess MLL-r target genes HOXA9 and MEIS1 in the KOPN-8 and NOMO-1 resistant cell lines, we noted a divergence in response between the two cell lines (Fig. 2B). The effects of DOT1L inhibition through either small molecule inhibition or genetic knockdown have established that loss of DOT1L enzymatic activity will lead to decreased transcription of MLL-r target genes HOXA9 and MEIS1 (11, 16, 21, 25). As shown in Figure 2B, resistant KOPN-8 cells maintained significantly higher levels of HOXA9 and MEIS1 in the presence of pinometostat than their sensitive counterparts. In contrast, both resistant and sensitive NOMO-1 cells showed similar depletion of HOXA9 and MEIS1, suggesting that two different resistance mechanisms may be at work. Moreover, these data also suggest that KOPN-8 cells are able to maintain HOXA9 and MEIS1 levels even when global H3K79me2 is reduced. To understand the discrepancy between the persistent global inhibition of H3K79me2 and recovery of target gene expression in the resistant KOPN-8 cells, we performed ChIP-seq using an antibody specific for the dimethylation state of H3K79 (Fig. 2C). Results from the ChIP-seq provided a potential explanation for how target gene expression recovered in the
KOPN-8 resistant cells, despite sustained global methyl mark inhibition. In the sensitive KOPN-8 cells, reduced methylation was observed across the entire genome while the resistant cells showed significantly increased methylation at many MLL-r target gene loci, including HOXA9 and MEIS1. Thus, despite global methylation inhibition in the resistant KOPN-8 line, we propose that the recovery of H3K79me2 levels at key target genes reactivated transcription, as observed with HOXA9 and MEIS1. An evaluation of all genes with recovered H3K79me2 in the resistance setting reveals a significant overlap with genes harboring high baseline H3K79me2 (Fig. 2D). Unlike the KOPN-8 line, gene specific H3K79me2 inhibition was maintained across the whole genome in both the resistant and sensitive NOMO-1 cells.

**Overexpression of Drug Efflux Transporter ABCB1 Correlates with Primary and Secondary Resistance to Pinometostat**

Increased expression and function of drug transporters is a common mechanism for both primary and secondary resistance to anticancer drugs (43-46). Drug metabolism studies performed with pinometostat identified it as a substrate for the common drug transporter ATP-binding cassette subfamily B member 1 (ABCB1, P-gp, P-glycoprotein, MDR1) (Supplemental Table S1). Based on this result, we reviewed our RNA-seq data to determine if development of pinometostat resistance influenced the expression of a set of membrane transporters commonly assessed in small molecule drug development. Examination of expression data from this panel of transporters revealed that resistant KOPN-8 cells had a greater than 100-fold increase in ABCB1 expression over the vehicle control cells (Fig. 3A). In contrast, and consistent with sustained HOXA9 and MEIS1 inhibition, little to no change in ABCB1 expression was observed in the resistant NOMO-1 cell line (Fig. 3B). Based on these data it is reasonable to suspect activity of a drug efflux transporter is responsible for the recovery of H3K79me2 and MLL-r target genes in our KOPN-8 pinometostat resistant cell line model.
Expression of $ABCB1$ has been described as a poor prognostic factor for elderly patients with AML and has been implicated with relapsed AML, leukemia arising from myelodysplastic syndrome, and treatment with certain anti-cancer therapies such as daunorubicin (47-49). We sought to determine if a correlation existed between $ABCB1$ expression and the inhibitory effect of pinometostat on H3K79me2 levels. We measured $ABCB1$ expression levels by qPCR in a panel of leukemia cell lines and observed a wide range of expression, from undetectable to up to nearly 250 times the expression of the lowest detectable expression (Fig. 4A, Supplemental Table S2). We then determined the H3K79me2 IC$_{50}$ after four day treatment with pinometostat in a subset of the leukemia cell lines. In the cell lines with detectable expression of $ABCB1$, the relative expression of $ABCB1$ correlated with the H3K79me2 IC$_{50}$ for that line (Fig. 4B). Of the cell lines analyzed, the non $MLL$-r AML line KG-1a had the highest expression of $ABCB1$ and correspondingly greatest H3K79me2 IC$_{50}$. To determine if high $ABCB1$ expression was causing the high H3K79me2 IC$_{50}$ in the KG-1a cell line, we co-treated KG-1a with pinometostat and the ABCB1 inhibitor, valsodar (PSC-833) (Fig. 4C). Valspodar is a non-immune suppressant analogue of cyclosporin A and has been tested clinically in combination with a variety of anti-cancer treatments, including paclitaxel, doxorubicin, and daunorubicin (50-54). When the KG-1a cell line was treated with pinometostat and valsodar, the H3K79me2 IC$_{50}$ was significantly reduced.

Discovering that inhibition of ABCB1 led to increased target engagement in a model with inherent overexpression, we sought to confirm the extent to which the mechanism of resistance in the KOPN-8 resistant cells was mediated by overexpression of ABCB1. We dosed both the resistant KOPN-8 and NOMO-1 cell lines with a combination of pinometostat and valsodar (Fig. 5A). Inhibition of ABCB1 through valsodar fully resensitized the resistant KOPN-8 cells to the action of pinometostat, as evidenced by their reduced growth rate from day four through the end of the experiment at day 21. No effects on growth were observed in the KOPN-8 control line with single agent valsodar (Fig. 5A). After
four days of co-treatment with valspodar in the KOPN-8 and NOMO-1 cell lines, we also assessed H3K79me2 at several concentrations (Fig. 5B). Treating the resistant KOPN-8 cells with valspodar and pinometostat shifted H3K79me2 back to a similar level as the naïve cells treated with pinometostat alone. To verify drug transporters were not playing a role in the NOMO-1 resistance, we also dosed these cells in combination with valspodar and pinometostat. Valspodar had no single agent activity with the NOMO-1 cells, nor did it resensitize the resistant cells to pinometostat, consistent with an ABCB1 independent mechanism of pinometostat resistance in NOMO-1 (Fig. 5A). Samples were also collected for analysis of H3K79me2 and the results were consistent with the proliferation results, as the combination with valspodar had no effect (Fig. 5B).

**Additional MLL-r Models Support Overexpression of ABCB1 as the Primary TER Mechanism**

Because our studies identified two distinct resistance mechanisms we wanted to test additional MLL-r models to identify if there was a common mechanism among MLL-r cell lines. Additional MLL-r cell lines, MV4-11 (MLL-AF4), MOLM-13 (MLL-AF9), and SEM (MLL-AF4) were treated continuously with pinometostat, and similar to the KOPN-8 and NOMO-1 cells, acquired resistance within three weeks from the initiation of treatment. All three of these additional models demonstrated that the phenomenon of ABCB1 overexpression was not unique to the KOPN-8 cells, with fold increases of ABCB1 ranging from 10 to 2,000 times the vehicle control (Supplemental Fig. S1A). We then sought to determine if the mechanism of resistance was mediated by overexpression of ABCB1. Utilizing the MV4-11 and SEM resistant cell lines, we treated with a combination of pinometostat and valspodar. Inhibition of ABCB1 with valspodar fully resensitized both the MV4-11 and SEM cell lines to the effect of pinometostat (Supplemental Fig. S1B). Results from these additional cell lines identify ABCB1 overexpression as a common mechanism by which MLL-r cell lines can bypass the effect of pinometostat.
Hypothesis Generation and Testing of NOMO-1 Mechanism of Resistance

With the inability of valsapodar to resensitize NOMO-1 cells to pinometostat treatment and minimal increase in ABCB1 expression observed in these cells, we set out to determine what other mechanisms were playing a role in the NOMO-1 resistance. Published results with another PMT, EZH2, demonstrated that one potential mechanism of resistance in this enzyme family is through acquisition of a secondary mutation leading to decreased binding of small molecule inhibitors (35, 36). To explore if NOMO-1 cells acquired resistance through a secondary mutation, DNA was obtained from both the parental and TER NOMO-1 cells and subjected to Sanger sequencing across the full coding region of DOT1L. In addition to sequencing the NOMO-1 cells, we also obtained DNA from KOPN-8 parental and TER cells, as a representative model from the group demonstrating ABCB1 overexpression as the primary mechanism of resistance. In both cell line models the DOT1L sequence remained unchanged between parental and resistant lines. Therefore, we were unable to detect any evidence of a resistance mechanism for pinometostat being driven through the acquisition of mutations within DOT1L (Supplemental Fig. S2). Given the apparent lack of pinometostat selected mutations in DOT1L we investigated possible alternative mechanisms which could potentially lead to pinometostat resistance. Detailed gene expression and pathway analysis was performed on the ChIP-seq and RNA-seq data sets generated from the naïve and resistant NOMO-1 cell lines. Principal component analysis of the RNA-seq data demonstrated that NOMO-1 cell line gene expression signatures could be distinguished by whether they were treated with pinometostat or vehicle (PC1) and their pinometostat sensitivity status and treatment time, i.e. resistant or naïve cell line status (PC2), indicating that altered gene expression occurs following pinometostat exposure and likely plays a role in the development of resistance (Supplemental Fig. S3). Combined analysis of the ChIP-seq and RNA-seq data identified many potential mechanisms involved in the development of resistance, including the PI3K/AKT and
RAS/RAF/MEK/ERK signaling pathways, the latter being of greater interest to us due to the availability of a number of specific inhibitors targeting a variety of nodes in the pathway (Supplemental Fig. S4A-C, Supplemental Table S3, Supplemental Table S4) (55). We tested combinations of pinometostat with MEK, ERK, and BRAF inhibitors in the resistant NOMO-1 cells. We first determined IC_{50} values in naïve NOMO-1 cells then used those doses to treat in combination to determine whether the inhibitors could resensitize the resistant NOMO-1 cells. None of the three compounds tested (trametinib, GDC-0994, and dabrafenib) were effective in resensitizing the resistant NOMO-1 cells to pinometostat (Supplemental Fig. S5)(40, 41). Of note, both the control and resistant NOMO-1 lines displayed exquisite single agent sensitivity to the MEK1/2 inhibitor trametinib with three day IC_{50} values less than 1 nM (40). To reconcile the lack of resensitization of the resistant NOMO-1 line with inhibitors of the RAS/RAF/MEK/ERK signaling pathway with the finding of pathway activation, we measured levels of phospho-proteins in the RAS/RAF/MEK/ERK pathway by immunoblot. Consistent with the proliferation results, we did not observe any changes in levels of protein phosphorylation between the control and resistant NOMO-1 lines (Supplemental Fig. S6). The lack of pathway activation as determined by phospho-protein suggests that although the gene expression data indicated key pathway signaling in the resistant cell lines, this change in transcription did not appear to translate functionally into altered pathway activation as measured by phosphorylation status of the pathway components. Thus, while RAS/RAF/MEK/ERK signaling may be altered transcriptionally in the resistant NOMO-1 cell line, this alteration does not appear to be driving resistance to pinometostat. To further interrogate the RNA-seq data set, pathway analysis was performed using the KEGG, GO – Biological Process, and MSigDB databases comparing the resistant cell line with the pinometostat treated (sensitive) parental cell line. From the KEGG and GO analysis, 838 and 313 significant pathways were identified, respectively. In the MSigDB analysis, of 1036 gene sets, none were significantly (FDR < 0.05)
upregulated while 26 were significantly downregulated (Supplemental Table S5, Supplemental Table S6). The most highly significant pathways were largely unrelated to established oncogenic pathways, but these results provide future avenues for potential investigation into the molecular understanding of NOMO-1 TER.

Discussion

DOT1L is known to play a role in MLL-r leukemia through its recruitment by MLL fusion proteins and subsequent hypermethylation of a characteristic set of genes known to drive leukemogenesis (9, 10, 16). The strong scientific rationale supporting the role of DOT1L in a subtype of acute leukemias makes it a target of interest for cancer treatment. With clinical development of pinometostat ongoing we sought to identify potential mechanisms leading to TER.

Small molecule based targeted therapies have greatly expanded the options available for oncology patients, and resistance to those drugs can emerge. A common resistance mechanism found with many of the tyrosine kinase inhibitors is the acquisition of molecular variants ranging from single nucleotide changes to intragenic deletions (27). Specifically in the PMT family, which includes DOT1L and EZH2, publications by Gibaja et al and Baker et al highlighted secondary mutations in both wild-type and mutant EZH2 alleles leading to resistance of specific small molecule PMT inhibitors in preclinical in vitro models (35, 36). In this study, we developed five MLL-r cell line models of acquired pinometostat resistance. Sanger sequencing of DOT1L in the KOPN-8 and NOMO-1 cell lines determined that our resistant pools had not acquired a secondary mutation. However, given the limitations of Sanger sequencing for detecting low variant allelic frequencies we cannot exclude the possibility of the existence of a sub-clonal DOT1L mutation. Given that the sub-clone would not likely be a driver of
resistance, we believe we uncovered PMT resistance mechanisms beyond what was previously described in the literature.

The observation that marginal recovery of H3K79me2 at specific MLL-r target loci led to reactivation of HOXA9 and MEIS1 transcription strongly suggests a threshold increase in H3K79me2 levels is required to support a resistant state. However, once this threshold is reached, the loss of H3K79me2 at remaining genetic loci does not impact the cell’s viability or typical functionality. A well-known mechanism of drug resistance is the overexpression of drug efflux transporters (43-46). We mined our RNA-seq data comparing naïve and resistant pools looking specifically at transcripts of transporters highlighted by the FDA as being clinically significant to drug-drug interactions (56). Several transporters published in the FDA guidance also play a role in drug resistance such as ABCB1 and ABCG2 (57). Analysis of the RNA-seq data identified greater than 100-fold upregulation of ABCB1 in KOPN-8 resistant cells. Overexpression of ABCB1 is consistent with the focused recovery of H3K79me2 at genes critical to driving proliferation and maintaining a cell differentiation block. Following this observation, we treated resistant KOPN-8 cells with an inhibitor of ABCB1 and were able to resensitize the cells to pinometostat. These results suggest the mechanism of resistance is likely due to decreased intracellular levels of pinometostat in KOPN-8 resistant cells, due to rapid efflux via the ABCB1 transporter. Considering inhibition of ABCB1 is not currently a viable clinical strategy, the question remains as to how this mechanism of resistance might be handled clinically. One potential strategy would be to develop a new generation of DOT1L inhibitors with a drug design emphasis on removing characteristics which lead to the molecule being a substrate for drug efflux. Lead candidates from a new series of inhibitors could be screened in models of resistance and/or cellular biochemical assays run in cell lines that have high baseline expression of ABCB1 such as KG-1a. Alternatively, signaling pathways responsible for the modulation of ABCB1 expression could also be investigated. However, these
signaling pathways are known to be complex and varied and to date have yet to be validated as a clinically relevant mechanism of pinometostat resistance (45, 46).

The NOMO-1 cell line generated with resistance to pinometostat was established to have a mechanism distinct from drug efflux. This observation led us to explore pathways identified through transcript analysis as being altered between the naïve and resistant cells. Unlike ABCB1, several of the pathways identified in the resistant NOMO-1 cells are druggable with clinically relevant inhibitors and could easily be tested to determine if pathway inhibition resensitized the cells to pinometostat. Treatment of resistant NOMO-1 cells with representative inhibitors of the RAS/RAF/MEK/ERK signaling pathway (e.g. dabrafenib and trametinib) failed to resensitize the cells to pinometostat. These data suggest that mechanisms and pathways identified through our analysis of transcripts did not represent cellular dependence on increased signal transduction of this pathway. Indeed, when we probed protein lysates harvested from naïve and resistant NOMO-1 cells we saw no evidence of increased phosphorylation of the RAS/RAK/MEK/ERK pathway. Although not all pathways shown to be differentially expressed by RNA-seq were exhaustively pursued, we believe that given the lack of active signaling, further exploration would have a low probability of success. AML is a complex disease with multiple molecular variants capable of driving leukemogenesis (58-60). Considering NOMO-1 cells harbor a KRAS G13D mutation, it is possible that MLL-r may not be the main oncogenic driver (61). Unsurprisingly, NOMO-1 cells were exquisitely sensitive to single agent treatment with the MEK inhibitor trametinib. Given the timeframe to development of pinometostat resistance, along with the maintenance of H3K79me2 inhibition both globally and at MLL-r target gene loci, it is conceivable that the development of resistance indicated a shift from reliance on programming caused by the MLL-AF9 fusion to RAS pathway addiction.
While our investigations in this manuscript utilized established cell line models, the authors believe corroboration of pinometostat resistance in primary patient samples would provide more clinical relevance. However, these studies cannot currently be performed given the limited life span of primary patient samples and the extended timeframe required to induce resistance. Furthermore, the potential for investigation of TER in vivo is limited by the pharmacokinetic properties and exposure levels of pinometostat needed to perform xenograft studies, necessitating delivery through a continuous IV infusion in nude rats. Delivery of pinometostat through continuous IV infusion presents additional challenges due to the extended dosing period required and the need to maintain pump and IV line integrity for several weeks. Given the practical limitations of alternative models and the need for long term maintenance of pinometostat exposure to drive resistance, apart from a clinical trial, cell lines are currently the only human model system in which it is feasible to explore pinometostat resistance.

In summary, we developed several novel models of resistance to the DOT1L inhibitor pinometostat. The evidence provided suggests that resistance was not induced via a direct target mutation-based mechanism. Instead, investigations into the mechanisms of TER identified that two independent mechanisms were operating in our cell line models. In four of the five resistant models we observed a clear association of pinometostat activity with ABCB1 overexpression, and ABCB1 was identified as the likely primary mechanism of acquired pinometostat resistance. In contrast the NOMO-1 mechanism of resistance remains unknown but is independent of ABCB1. Development of these cell line models and understanding the mechanisms of resistance may have utility in exploring clinical mechanisms of resistance to pinometostat.

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References


Tables

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Table 1. Inhibition of H3K79 methylation and cellular proliferation by pinometostat in control and resistant cell lines. Pinometostat 4 day H3K79me2 IC_{50} (µM) and 14 day long term proliferation IC_{50} (µM) measured in the control and resistant cell lines. ¹ ± error represents the 95% confidence interval ² Fold-change calculated as [(Resistant – Control)/Control] values.

Figure Legends

Figure 1: Development of Pinometostat Resistance in Cell Line Models of MLL-r. Cellular MLL-r models sensitive to single agent treatment of pinometostat were continuously treated with 4.5 µM inhibitor to determine if resistance would develop. The cell lines KOPN-8 and NOMO-1 regained growth rates similar to the vehicle control after approximately 21 days of dosing. Following the transition to a pinometostat refractive state, both cell lines were maintained in growth media containing pinometostat to assure the resistant pool was stable. Viable cells were counted and split every 3 to 4 days in the presence of pinometostat or DMSO vehicle control and split adjusted results plotted on a logarithmic scale.
Figure 2: Measures of Pharmacodynamic Markers over Time. Analysis of pharmacodynamic markers over the course of development of resistance [A] Global H3K79me2 analysis by ELISA in KOPN-8 and NOMO-1 cells following treatment with 4.5 μM pinometostat at indicated time points. H3K79me2 levels are plotted as a percentage of vehicle control. [B] RNA-seq analysis of DOT1L target genes in KOPN-8 and NOMO-1 vehicle control, pinometostat sensitive and pinometostat resistant cells. Relative mRNA expression levels are plotted as a percentage of those in vehicle-treated control cells. HOXA9 and MEIS1 gene expression begins to recover in resistant KOPN-8 cells while no recovery of target gene expression is observed in resistant NOMO-1 cells. [C] Assessment of H3K79me2 at HOXA9 and MEIS1 gene loci using ChIP-seq. The difference between the sensitive and resistant cell lines is shown as the FDR corrected p-value (q-value). Significant recovery (q ≤ 0.05) of H3K79me2 was observed at both the HOXA9 and MEIS1 gene locus as the KOPN-8 cells developed resistance to pinometostat. No significant recovery of H3K79me2 was observed in NOMO-1 cells at any locus. [D] KOPN-8 ChIP-seq data plotted as the significance (-log(FDR)) of the change in H3K79me2 methylation levels between the day 28 vs day 10 treatment vs the baseline methylation status. The majority of the significant changes in methylation status (q ≤ 0.05) occur in genes with high baseline methylation signal. The histogram above the plot demonstrates the distribution of genes across the methylation signal spectrum. Genes with a high baseline methylation signal represent only a small portion of all genes assessed.

Figure 3: Membrane Transporter mRNA Expression in Resistant Cell lines. RNA-seq analysis of common drug efflux transporters in [A] KOPN-8 resistant cells at day 28 of treatment and [B] NOMO-1 resistant cells at day 53 of treatment, plotted as the fold change in RNA-seq counts in the resistant versus control cell line.
Figure 4: Overexpression of \textit{ABCB1} Positively Correlates with Primary Resistance to Pinometostat. Correlation of \textit{ABCB1} expression with primary resistance to pinometostat in a panel of leukemia cell lines [A] Quantitative real-time PCR analysis of \textit{ABCB1} expression and H3K79me2 IC$_{50}$ values in a panel of leukemia cell lines. Relative mRNA expression levels are plotted relative to KOPN-8. Table of H3K79me2 IC$_{50}$ values available in Supplemental Table S2 [B] Correlation of \textit{ABCB1} expression with H3K79me2 IC$_{50}$ values [C] \textit{ABCB1}-high cell line KG-1a was treated with pinometostat single agent or in combination with 1 µM valsopodar for 4 days, and inhibition of H3K79me2 mark assessed by ELISA.

Figure 5: Pinometostat Resensitization of KOPN-8 but not NOMO-1 Resistant Cell Lines Achieved through Dosing in Combination with Valspodar. Resensitization of ABCB1-mediated resistance with ABCB1 inhibitor. [A] Growth of KOPN-8 and NOMO-1 resistant and DMSO vehicle control cells when treated with 1 µM valsopodar for 21 days. Resistant cells were cultured in media containing 4.5 µM pinometostat. Viable cells were counted and split every 3 to 4 days and split adjusted results plotted on a logarithmic scale. [B] Global H3K79me2 analysis by ELISA in resistant KOPN-8 and resistant NOMO-1 cells following combination treatment with 1 µM valsopodar and pinometostat at indicated concentrations for 4 days. H3K79me2 levels are plotted as a percentage of vehicle control.
Figure 1

KOPN-8
(*MLL-ENL*)

Viable cells/mL

Viable cells/mL

NOMO-1
(*MLL-AF9*)

Time (Days)

Time (Days)

Vehicle
Pinometostat

Vehicle
Pinometostat

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**Figure 2**

**A.**

Methylation signal (vehicle) log(FDR) for differential methylation (Day 28 vs Day 10 pinometostat).

**B.**

Relative Expression of Vehicle Control

**C.**

Relative Expression of Vehicle Control

**D.**

-log(FDR) for differential methylation (Day 28 vs Day 10 pinometostat) vs Methylation signal (vehicle).
Figure 3

A. Relative Expression Fold Increase Over Control

KOPN-8

B. Relative Expression Fold Increase Over Control

NOMO-1
Figure 4

A. ABCB1 Expression and H3K79me2 IC$_{50}$ in a Leukemia Panel

B. ABCB1 Expression Relative to H3K79me2 IC$_{50}$

C. KG-1a

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Mechanisms of Pinometostat (EPZ-5676) Treatment Emergent Resistance in MLL Rearranged Leukemia

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