Concomitant targeting of BCL2 with venetoclax and MAPK signaling with cobimetinib in acute myeloid leukemia models

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ABSTRACT

he pathogenesis of acute myeloid leukemia (AML) involves serial acquisition of mutations controlling several cellular processes, requiring combination therapies affecting key downstream survival nodes in order to treat the disease effectively. The BCL2 selective inhibitor venetoclax has potent anti-leukemia efficacy; however, resistance can occur due to its inability to inhibit MCL1, which is stabilized by the MAPK pathway. In this study, we aimed to determine the anti-leukemia efficacy of concomitant targeting of the BCL2 and MAPK pathways by venetoclax and the MEK1/2 inhibitor cobimetinib, respectively. The combination demonstrated synergy in seven of 11 AML cell lines, including those resistant to single agents, and showed growth-inhibitory activity in over 60% of primary samples from patients with diverse genetic alterations. The combination markedly impaired leukemia progenitor functions, while maintaining normal progenitors. Mass cytometry data revealed that BCL2 protein is enriched in leukemia stem/progenitor cells, primarily in venetoclax-sensitive samples, and that cobimetinib suppressed cytokine-induced pERK and pS6 signaling pathways. Through proteomic profiling studies, we identified several pathways inhibited downstream of MAPK that contribute to the synergy of the combination. In OCI-AML3 cells, the combination downregulated MCL1 protein levels and disrupted both BCL2:BIM and MCL1:BIM complexes, releasing BIM to induce cell death. RNA sequencing identified several enriched pathways, including MYC, mTORC1, and p53 in cells sensitive to the drug combination. In vivo, the venetoclax-cobimetinib combination reduced leukemia burden in xenograft models using genetically engineered OCI-AML3 and MOLM13 cells. Our data thus provide a rationale for combinatorial blockade of MEK and BCL2 pathways in AML.

Introduction

Acute myeloid leukemia (AML) is characterized by the uncontrolled proliferation and arrested differentiation of abnormal stem and progenitor cells. Standard induction chemotherapy induces a high rate of complete remission but fails to improve overall survival especially in elderly patients with AML.^{1,2} Genes significantly mutated in AML can be organized into several functional categories that are associated with enhanced proliferation, impaired differentiation, deregulated chromatin modification, and DNA methylation.^{1,3} Therefore, co-targeting downstream pathways that contribute to



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leukemogenesis may deliver the greatest clinical efficacy.

The anti-apoptotic protein BCL2 has been studied extensively for its role in leukemic transformation and chemoresistance. BCL2 is highly expressed in AML leukemia stem cells (LSC) containing low levels of reactive oxygen species which are resistant to chemotherapy.^{4,5} BCL2 inhibitors have been shown to eradicate AML LSC and sensitize chronic myeloid leukemia LSC to tyrosine kinase inhibitors. Oncogenic dependency on BCL2 was found in AML patients carrying mutations in IDH1 and IDH2.67 A recent study used BH3 profiling to discover that co-inhibition of BCL2 with tyrosine kinase inhibitors facilitated eradication of genetically diverse AML in patient-derived xenograft (PDX) models.⁸ We have reported the anti-leukemia potency of venetoclax (ABT-199/GDC-0199), an orally bioavailable BH3 mimetic that selectively binds with high affinity to BCL2, but lacks affinity for BCL-XL and MCL1, in AML models.⁹ In a phase II clinical trial, venetoclax monotherapy had clinical activity in patients with relapsed or refractory AML with a tolerable safety profile.¹⁰ However, the inability of venetoclax to inhibit MCL1 causes resistance in leukemia cells that require MCL1 for survival.^{11,12}

The RAF/MEK/ERK (MAPK) cascade, a major effector pathway activated in 70%-80% of patients with AML, is activated by upstream mutant proteins such as FLT3, KIT, and RAS.¹³⁻¹⁵ The MAPK pathway regulates BCL2 family proteins by stabilizing anti-apoptotic MCL111,16 and inactivating pro-apoptotic BIM (BCL2L11).¹⁷ Monotherapy with MEK inhibitors has had limited clinical efficacy.¹⁵ Recently it was shown that MAPK signaling activation contributed to primary resistance to an IDH2 inhibitor¹⁸ and acquired resistance to venetoclax,¹⁹ suggesting that combination regimens that include MEK inhibitors could be efficacious in these patients. Cobimetinib (GDC-0973) is an allosteric MEK inhibitor with antitumor activity in BRAF- and KRASmutant tumor cells,20 and was recently approved to treat patients with metastatic melanoma. Its anti-AML efficacy, particularly in combination with venetoclax, is unknown.

Furthermore, biomarkers predictive of response to this combination in AML are unknown. Suppression of downstream pERK does not predict sensitivity to MEK inhibition.²¹ In melanoma, it has been demonstrated that mTORC1/2 and pS6 activities are associated with acquired resistance to MEK inhibitors and suppression of pS6 may serve as a biomarker to predict clinical response to MEK inhibitors.^{22,23} The role of pS6 in response to MEK and BCL2 inhibition has not been addressed.

In this study, we evaluated the anti-leukemia effects of concomitant BCL2 and MAPK blockade by venetoclax and cobimetinib in AML cell lines, primary patients' samples, and xenograft murine models. Through the use of reverse-phase protein arrays (RPPA) and RNA sequencing, we identified pharmacodynamic markers that correlated with the efficacy of the combination treatment, in particular activated pS6 (Ser235/236), which discriminated combination-sensitive from -insensitive AML cells. Our data support the rationale for dual inhibition of the BCL2 and MEK pathways.

Methods

Patients' samples, acute myeloid leukemia cell lines, and reagents

Bone marrow and peripheral blood samples were collected from patients with AML or healthy donors after informed con-

sent had been obtained in accordance with the Institutional Review Board of The University of Texas MD Anderson Cancer Center. The cell line culture methodology is described in the *Online Supplementary Methods*.

Assays and other methods

Details of the CellTiter-Glo proliferation assay, colony-forming cell assay, electrochemiluminescent enzyme-linked immunosorbent assay, RPPA, and RNA sequencing are provided in the *Online Supplementary Methods*. Antibody conjugation for mass cytometry staining and the spanning-tree progression analysis of density-normalized events (SPADE) analysis are also explained in detail in the *Online Supplementary Methods*.

Apoptosis in primary acute myeloid leukemia samples

As previously reported,²⁴ 18 primary AML peripheral blood mononuclear cells or AML PDX samples were cultured in LSC medium. Viable AML CD45^{dm} blast cells were enumerated by using CountBright counting beads (Cat. C36950; Invitrogen, Carlsbad, CA, USA) with concurrent annexin V and DAPI detection on a Gallios Flow Cytometer (Beckman Coulter, Indianapolis, IN, USA). Data analysis and additional details are included in the Online Supplementary Methods.

In vivo study of cobimetinib and venetoclax in acute myeloid leukemia xenograft mouse models

The animal studies were performed in accordance with guidelines approved by the Institutional Animal Care and Use Committee at MD Anderson. Nonobese diabetic/severe combined immunodeficient gamma IL3-GM-SF (NSG-SGM3 or NSGS) mice (female, 8-10 weeks old) were purchased from Jackson Laboratory (Bar Harbor, ME, USA). The mice were injected intravenously with OCI-AML3-Luci-GFP (1.0×10⁶) cells, which were lentivirally transduced with firefly luciferase. Leukemia engraftment was confirmed 1 week after injection through a noninvasive in vivo bioluminescence imaging (BLI) system (Xenogen, Alameda, CA, USA) after injection of a Dluciferin (4 mg/mouse) substrate. Mice were distributed into four groups (11 mice/group) with comparable tumor burden and dosed daily for 4 weeks with one of the following oral preparations: vehicle, cobimetinib (10 mg/kg), venetoclax (100 mg/kg), or cobimetinib plus venetoclax. BLI was performed weekly to determine the extent of engraftment. Survival was monitored as an endpoint. A similar MOLM13 model is described in the Online Supplementary Methods.

Statistical analyses

The Student *t*-test was used to analyze the statistical significance of differences between groups, both *in vitro* and *in vivo*. All statistical tests were two-sided, and the results are expressed as the mean \pm standard deviation. A *P* value <0.05 was considered statistically significant. The RPPA and RNA-sequencing data analysis are described in the *Online Supplementary Methods*.

Results

Cobimetinib and venetoclax demonstrate synergistic anti-leukemia efficacy in acute myeloid leukemia cell lines *in vitro*

To assess the anti-leukemia activity of cobimetinib and venetoclax as single agents or in combination, we studied their effects on cell proliferation of 11 AML cell lines (Table 1). The median inhibitory concentration (IC_{50}) values of single agents were determined in a dose-response

manner using CellTiter-Glo (CTG) assays after 72 h of drug treatment. The IC₅₀ values of cobimetinib (range, $0.002 \,\mu\text{M}$ - $3.0 \,\mu\text{M}$) did not correlate with either the status of RAS mutations or the basal levels of p-ERK1/2 determined by flow cytometry (Table 1). To assess pharmacological interactions between cobimetinib and venetoclax, incremental doses were applied based on the IC₅₀ value of each drug. In seven of the 11 cell lines, combination of the agents elicited synergistic growth inhibition based on the Chou-Talalay method of analysis [combination index (CI) <0.8].²⁵ Cell lines with IC₅₀ values below the selected cutoff values (0.3 μ M for cobimetinib²⁰ and 0.1 μ M for venetoclax⁹) were defined as sensitive to the single agent. Patterns of response to single agents and the combination were distinct. Notably, while synergy was observed in both venetoclax-resistant (MOLM14, OCI-AML3, NB4) and cobimetinib-resistant cell lines (KG1, MOLM13), the lowest CI value (0.12) was seen in venetoclaxsensitive/cobimetinib-resistant AML cells (KG1) (Figure 1).

Cobimetinib and venetoclax demonstrate on-target suppression of cell proliferation and clonogenic potential of leukemia progenitors in a subset of primary acute myeloid leukemia cells *ex vivo*

The anti-leukemia activities of cobimetinib and venetoclax were examined in 18 primary samples with diverse genetic alterations, collected from patients with AML or spleen from PDX models (Table 2). Primary AML blasts were treated with cobimetinib and venetoclax alone or in combination at 0.1 μ M for 5 days in LSC medium to maintain the immature state of the leukemia cells.²⁴ Cobimetinib alone induced minimal cell death (specific apoptosis, $6.7 \pm 5.9\%$), which was significantly enhanced when the drug was given in combination with venetoclax $(27.7 \pm 20.2\%, P=0.001)$ (Figure 2A, left). Cobimetinib inhibited cell proliferation in the majority of cases (34.2 \pm 23.7%), and this suppression was more pronounced when the drug was combined with venetoclax ($60.2 \pm 28.8\%$, P < 0.001) (Figure 2A, right). Venetoclax as a single agent reduced viable cell numbers by more than 50% in six cases (33.3%). Three of the four AML samples demonstrating over 50% growth inhibition by the cobimetinib

treatment carried the FLT3-ITD and/or D835 point mutation (AML 12, 13, and 17). As previously reported, IDHmutant AML samples were highly sensitive to venetoclax as a single agent (AML 2 and 15). Over 60% (11 of 18) of the patients' samples responded to the combination treatment, including those insensitive to either compound alone (AML 1, 8 and 11). Importantly, induction of apoptosis in AML stem/progenitor CD34⁺CD38⁻CD123⁺ population following the combination treatment was observed in two out of four AML samples tested (Online Supplementary Figure S1). The clonogenic potential of myeloid progenitors was significantly suppressed by the combination (82.5 \pm 20.0%), as compared to cobimetinib $(38.3 \pm 14.6\%, P=0.01)$ or venetoclax $(41.9 \pm 18.6\%,$ P < 0.05) alone. Normal progenitor function was minimally affected (Figure 2B and Online Supplementary Figure S2).

To test the on-target efficacy of both agents, we developed a 28-parameter mass cytometry [time-of-flight mass spectrometry (CyTOF)] panel comprising antibodies against surface antigens to define AML stem/progenitor fractions and intracellular proteins of the BCL2 family and various signaling pathways²⁶ (Online Supplementary Table S1). The CyTOF study was performed in AML13 (sensitive to the combination) and AML14 (resistant to the combination) samples (Figure 2C). SPADE trees were built and annotated using all cell surface markers (Online Supplementary Table S1); the positive markers were included in the heat maps (Online Supplementary Figure S3). BCL2 protein levels were significantly enriched in CD34⁺ stem/progenitor cells compared to CD34- cells and BCL2 was expressed at a higher level in the venetoclax-sensitive sample (AML13) than in the venetoclax-resistant sample (AML14), consistent with our published data9 (Figure 2D). These results support the notion that venetoclax preferentially target LSC in AML. As previously reported, the cancer signaling network relies on the manner in which cancer cells respond to external stimuli rather than their basal phosphorylation state.²⁷ Therefore, following cobimetinib treatment, we stimulated primary AML cells with granulocyte colony-stimulating factor (G-CSF) or stem cell factor (SCF) to study MEK downstream signaling pathways under conditions mimicking a cytokine-rich bone marrow

fable 1. Cytotoxicity of	of cobimetinib and	venetoclax in acute	myeloid leukemia cell lines.
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Cell line	Mutations	Cobimetinib IC₅₀ (µM)	Venetoclax IC₅₀ (µM)	CI value	p-ERK (R-MFI)
MOLM13	<i>FLT3</i> -ITD	0.46	0.01	0.49	6.60
MOLM14	<i>FLT3-</i> ITD	0.16	1.88	0.36	9.68
MV4;11	<i>FLT3</i> -ITD	0.29	0.005	0.99	11.7
TF-1*	NRAS, TP53	0.51	10.3	1.51	2.56
OCI-AML3	NPM1, DNMT3A, NRAS	0.17	2.90	0.29	8.67
OCI-AML2	DNMT3A	0.002	0.04	0.78	3.72
THP1*	NRAS, TP53	0.56	39.1	0.54	4.48
KG1*	ITGB8, SMC2	3.06	0.03	0.12	3.73
NB4	PML-RARA	0.04	0.73	0.30	2.34
U937*	PTPN11, WT1	3.00	9.75	0.88	3.30
HL-60	CDKN2A, TP53, NRAS	0.45	0.004	0.89	3.83

Half maximal inhibitory concentration (IC_{st}) values were calculated on the basis of the number of viable cells quantified by CTG assay. CI: combination index; R-MFI: relative median fluorescence intensity determined by the ratio of the signal in the antibody-stained cells/autofluorescence of unstained cells; ITD: internal tandem duplication. *Data on gene mutations are from Cancer Cell Line Encyclopedia: <u>http://www.broadinstitute.org/ccle/home</u>

microenvironment. The SPADE trees were colored based on expression levels of CD34. In AML13, annotation 4 (6.5% of total viable cells) represented the leukemia stem/progenitor cell population by phenotypically positive expression of CD34, CD123, CD25, CD135, and CD64. Annotation 8 accounted for 53.4% of total viable cells in AML14, and was highly positive for expression of CD34, CD123, CD117, CD135, and CD64 (Online Supplementary Figure S3). In both samples, we observed low basal levels of pERK, which increased following G-CSF stimulation (3.9-fold in AML13 and 5.7-fold increase in AML14). G-CSF-stimulated pERK in both patients' samples was largely inhibited by cobimetinib despite differential responses in proliferation assays (Figure 2C, D), indicating that suppression of pERK does not predict sensitivity to MEK inhibition and is consistent with previous reports.²¹ Several studies have shown that suppression of mTORC1 and downstream pathways (especially S6) predicted sensitivity to MEK inhibition.^{21,22} We found that pS6 was highly activated by SCF and effectively suppressed by cobimetinib in the cobimetinib-sensitive AML sample, whereas the cobimetinib-resistant AML sample did not demonstrate a response to SCF. As in this study, we treated cells overnight and transiently stimulated then for 10 min to look into activation of signal transduction pathways (Figure 2). Due to limited exposure to the inhibitors (2 h), we were unable to detect changes in frequencies of AML stem/progenitor cells (*Online Supplementary Figure S4*).

Transcriptomic and proteomic profiles identify pharmacodynamic markers underlying responses to targeted agents

To identify the pharmacodynamic markers underlying the observed drug responses, we treated the 11 AML cell lines (Table 1) with cobimetinib and venetoclax as single



Figure 1. Anti-leukemia efficacy of cobimetinib and venetoclax in acute myeloid leukemia cell lines. Eleven acute myeloid leukemia cell lines were left untreated or treated with cobimetinib or venetoclax as single agents at 0.001, 0.01, or 1.0 μ M for 72 h. Calcusyn software was used to calculate the median inhibitory concentration (IC₅₀) values. Combinations of the two drugs were then tested on the same cell lines at dose ranges of 0.25, 0.5, 1, 2, and 4 times the IC₅₀ value of each combination index of each combination in each cell line was calculated on the basis on the luminescent intensity that correlated with number of viable cells determined by the CellTiter-Glo assay. Responses to treatment were categorized into four patterns: (i) sensitive to both drugs; (ii) sensitive only to venetoclax and showing synergy for the combination; (iv) resistant to both drugs. AML: acute myeloid leukemia; Cobi: cobimetinib; Ven: venetoclax; Cl: combination index

agents or in combination for 24 h at doses that were 0.5, 1 or 2 times their IC₅₀ values, followed by RPPA and RNA sequencing analysis. As already described, cells with IC₅₀ values <0.3 μ M for cobimetinib or <0.1 μ M for venetoclax were categorized as sensitive and those with IC₅₀ values above these cutoffs were considered resistant. For the combination groups, CI values <0.8 were considered synergistic.

Quantification of 90 proteins by RPPA identified several biomarkers that correlated with *in vitro* drug responses. For example, S6 phosphorylation at Ser235/236 was significantly reduced in both cobimetinib-sensitive and -resistant cell lines compared to untreated cells, with sensitive cells displaying higher basal phosphorylation at Ser235/236. Significant pMEK induction was observed in cobimetinib-resistant cell lines (Figure 3A). Several signaling pathways were highly activated under basal condi-

tions in cobimetinib-sensitive cells compared to resistant cells, including pS6 (Ser235/236), pRSK, pERK, p38MAPK and pPTEN (Online Supplementary Figure S5A). Proteins indicating responses to venetoclax treatment were largely limited to the caspase-dependent apoptotic cascade (data not shown). Higher levels of BAX and BCL2 and lower levels of BIM and pS6 (Ser240/244) correlated with sensitivity to venetoclax (Online Supplementary Figure S5B). In cell lines in which cobimetinib and venetoclax had a synergistic effect, several MEK downstream pathways were significantly downregulated and cleaved poly (ADP-ribose) polymerase (PARP) was detected, indicating induction of apoptosis (Figure 3B). These changes were not identified in cell lines in which a synergistic effect did not occur. The heat maps of the complete RPPA datasets are shown in Online Supplementary Figure S6.

Western blotting was performed to validate the RPPA

 Table 2. Clinical information for primary acute myeloid leukemia patients' samples.

AML#	Status	WBC (x10°/L)	Blasts,%	Cytogenetics	Molecular mutations	
Samples for 5-	lay culture					
1	NA	NA	NA	NA	IKZF1, NOTCH1, BCOR	
2	Relapsed	34.3	95	Complex	DNMT3A, IDH2, TP53, FLT3-N841	
3	Relapsed	21	96	Complex	FLT3-ITD, NPM1, WT1, DNMT3A	
4	Relapsed	14.9	98	Complex	JAK2, MPL, WT1	
5	Relapsed	6.5	94	46,XY,t(9;11)(p22;q23)	CEBPA, ATM	
6	Relapsed	18.3	57	47,XY,+21	RUNX1, TET2	
7	Relapsed	19.9	69	NA	EVI1	
8	De novo	13.5	18	46,XX	EZH2, MPL	
9	De novo	20.6	74	NA	NA	
10	De novo	5.9	21	Complex	TP53	
11	NA	40	24	Complex	FLT3-D835	
12	Relapsed	19	94	47,XY,+8	FLT3-D835, NOTCH1, ASXL1, KIT, TET2	
13	Relapsed	45.8	94	45,XY,der(17;18)	<i>FLT3</i> -ITD and <i>D835</i>	
14	Relapsed	6.4	72	Complex	EGFR, PTPN11, WT1	
15	Relapsed	5.4	25	47,XY,+8	RUNX1, ASXL1, IDH1, KRAS, NRAS, TET2	
16	De novo	12.7	31	Complex	RUNX1, TET2, TP53	
17	Relapsed	4.8	89	46,t(X;X)(q22;q26)	<i>FLT3</i> -ITD and <i>D835</i>	
18	Relapsed	4.6	48	Complex	FLT3-ITD, JAK2, RUNX1	
Samples for CFG	C assays					
19	De novo	85.5	51	46,XX	DNMT3A, IDH2m NPM1, ASXL1	
20	Relapsed	2.4	50	47,XX,+8	RUNX1, ASXL1, IDH1, TET2, NRAS, KRAS	
21	Relapsed	1.7	82	Complex	ASXL1, EZH2, IDH1, TET2, RUNX1	
22	Relapsed	5.8	32	Complex	DNMT3A, IDH1	
Samples for Cy	OF (only) study					
23	Relapsed	104.4	3	Complex	IDH2	
24	Relapsed	163.5	98	Complex	<i>FLT3</i> -ITD	
25	Relapsed	10.9	10	Complex	IDH2	
26	Relapsed	5.1	88	Complex	TP53, ATM	
27	Relapsed	80.1	72	46,XX	No mutations	
28	Relapsed	13.1	63	46,XX	TP53, IDH2	

AML: acute myeloid leukemia; WBC: white blood cell count; NA: not available; ITD: internal duplication; CFC: colony-forming cells; CyTOF: time-of-flight mass spectrometry. For 5-day culture assays, all samples were collected from peripheral blood, except AML #15, which was from bone marrow, and AML #1, #5, and #9, which were from patient-derived xenograft mouse spleens. All the samples for CFC assays were bone marrow.

L. Han et al.



SCF G-CSF Unstim Unstim G-CSF Unstim SCF Unstim Figure 2. Treatment with cobimetinib and venetoclax causes on-target suppression of cell proliferation and impairs leukemia progenitor function in a subset of primary acute myeloid leukemia cases. (A) Primary acute myeloid leukemia (AML) peripheral blood mononuclear or bone marrow cells from AML cases were cultured in serum-free expansion medium supplemented with BIT 9500 Serum Substitute and cytokines, including stem cell factor (SCF; 100 ng/mL), Flt3 ligand (50 ng/mL), interleukin 3 (IL3; 20 ng/mL), and granulocyte colony-stimulating factor (G-CSF; 20 ng/mL) as well as StemRegenin 1 (SR1; 1 µM). Cells were left untreated or treated with cobimetinib (Cobi) or venetoclax (Van), both at 0.1 μ M, as single agents or in combination (Combo). After culture for 5 days, cells were stained with CD45-PE, Annexin-V-APC, and DAPI. Apoptotic leukemia blasts (CD45dimAnnexin-V⁺) were isolated by flow cytometry. Results are expressed as percentage of specific apoptosis calculated by the formula: $100 \times (\%$ apoptosis of treated cells – % apoptosis of control cells)/(100 - % apoptosis of control cells). Percentage of growth inhibition was calculated on the basis of the number of control viable cells (Annexin-V⁻/DAPI⁻). **P<0.01; ***P<0.001. (B) Mononuclear cells collected from AML patients (100,000 cells) or healthy donors (50,000 cells; NBM) were plated in methylcellulose, then treated with venetoclax or cobimetinib (both at 0.1 μ M) as single agents or in combination. Colonies were scored on day 14. Data are presented as percentage inhibition compared to control groups. (C) The absolute cell counts of AML13 and AML14 samples as determined in (A) are shown in comparison with those of untreated controls (Contr), with median inhibitory concentration (IC₅₀) values indicated for each sample. (D) AML13 and AML14 samples were treated with cobimetinib 1.0 uM overnight followed by 10 min with or without (Unstim) stimulation with SCF or G-CSF (100 ng/mL). Cells were fixed, permeabilized, and processed for time-of-lfight mass spectrometry. Spanning-tree progression analysis of density-normalized events (SPADE) trees were generated by using markers shown in Online Supplementary Figure S2. The leukemia stem/progenitor populations were manually annotated and highlighted by analysis of all surface markers. The median intensities of pERK and pS6 in gated populations are shown. BCL2 expression in CD34 and CD34⁻ fractions in both samples is shown. DMSO: dimethylsulfoxide.

DMSO

Cobi

Median intensity

DMSO

Cobi

DMSO

Cobi

Median intensity

DMSO

Cobi



Figure 3. Pharmacodynamic markers of drug response identified through reverse-phase protein arrays and RNA sequencing. Acute myeloid leukemia (AML) cell lines were left untreated or treated with cobimetinib or venetoclax as single agents or in combination at 0.5, 1, or 2 times the median inhibitory concentration (IC_{so}) value of each compound in each cell line for 24 h. Cell pellets were harvested after treatment and subjected to reverse-phase protein array (RPPA) analysis as previously reported. (A) Plots depict proteins differentially expressed between cobimetinib-sensitive and cobimetinib-resistant cells. (B) The mean values of corresponding proteins in cell lines showing synergy to the combination treatment (CI-0.8 as presented in Table 1) are shown in the heatmap. Only the proteins showing significant differences (P<0.05) between control and treated groups are shown. C: untreated control; T: treated; S: sensitive; R: resistant; Syn: synergy. (C) Cells were treated with cobimetinib (Cobi), venetoclax (Ven), or a combination (Combo) at 10, 100 and 1000 nM for 4 h and subjected to lysis; proteins were separated and probed with the antibodies indicated. (D) AML cell lines were left untreated (control) or treated with cobimetinib or venetoclax as single agents or in combination at 10 times the median inhibitory concentration (IC_{so}) value of each compound in each cell line for 4 h. Cell pellets were harvested after treatment and subjected to electrochemiluminescent enzyme-linked immunosorbent assay. The levels of BCL2, MCL1, BCL2:BIM and MCL1:BIM complexes were plotted based on percentages of the levels in the control group. (E) AML cells were treated and processed as described above for the RPPA assay. RNA was isolated using the RNeasy kit and sent for mRNA sequencing. The enriched pathways in cell types showing synergy in response to the combination are shown.

data in four cell lines representing different response patterns (Table 1, Figure 3C and Online Supplementary Figure *S5C*). Suppression of pERK by cobimetinib was observed in both sensitive (OCI-AML3 and MV4;11) and resistant (MOLM13 and U937) cells at 0.01 µM, irrespective of response patterns. pS6 (Ser235/236 or Ser240/244) was inhibited by a low dose $(0.1 \ \mu M)$ of cobimetinib in sensitive OCI-AML3 cells but not in resistant U937 cells (Figure 3C), consistent with our RPPA findings noted above. These data also indicate direct suppression of mTORC1 signaling by cobimetinib as the Ser240/244 site is regulated exclusively by mTORC1. MYC was downregulated by cobimetinib alone or in combination with venetoclax in OCI-AML3, but not in U937 cells. Cell death characterized by elevated levels of cleaved PARP was observed in the combination group in OCI-AML3 cells, consistent with pro-apoptotic synergy (Figure 3C). To capture the dynamic interactions of pro- and anti-apoptotic BCL2 family members, a Meso Scale Discovery assay was performed in OCI-AML3 cells (Figure 3D). High basal levels of BCL2 and BCL2:BIM complexes were identified. After venetoclax treatment, BCL2:BIM complexes were disrupted and MCL1 protein levels were upregulated, resulting in increased MCL1:BIM complexes. The combination of cobimetinib with venetoclax suppressed both BCL2:BIM and MCL1:BIM complexes, enabling release of free BIM to induce cell death (Figure 3D). In addition, cobimetinib treatment induced total BIM protein levels in MV4;11 cells, thereby priming the cells for death (Online Supplementary Figure S7).

To refine our search for potential biomarkers correlating with response to the venetoclax-cobimetinib combination, we performed RNA sequencing and evaluated differential gene expression after exposure to the drugs. The aim was to identify hallmark cancer pathways significantly altered specifically in cells sensitive to the drug combination (Figure 3E and Online Supplementary Table S2). Our analysis demonstrated that several downstream pathways, including MYC, E2F and their target genes, were significantly altered after treatment in cells that responded synergistically to the combination. Consistent with western blot data, mTORC1 signaling was also altered in cells showing synergistic responses. Hypoxia and unfolded protein response (UPR) pathways were also significantly enriched, possibly downstream of mTOR/4EBP1/eIF4E signaling, which directs protein synthesis of HIF-1 α ,²⁸ and can trigger the UPR.²⁹ Glycolysis, another enriched pathway, is regulated by the ERK signaling pathway through RNK126-mediated ubiquitination of pyruvate dehydrogenase kinase, which may account for resistance to apoptosis.³⁰

The combination of cobimetinib and venetoclax reduces leukemia burden in acute myeloid leukemia models *in vivo*

To test the efficacy of cobimetinib and venetoclax *in vivo*, we induced leukemia in NSGS mice by injecting the animals with genetically engineered OCI-AML3/Luc/GFP cells. Leukemia engraftment was confirmed 1 week after injection using BLI. Mice were randomly distributed into four arms and dosed orally with vehicle, cobimetinib (10 mg/kg), venetoclax (100 mg/kg), or cobimetnib plus venetoclax daily for 28 days. BLI demonstrated that the leukemia burden was significantly reduced in treated groups compared to controls over time (Figure 4A). At

week 5, the tumor reduction was significantly greater in the groups that received single-agent cobimetinib (P<0.001) or cobimetnib plus venetoclax (P<0.001) than in the control group (Figure 4B). The tumor reduction was greater following combination treatment than following venetoclax (P<0.05) or cobimetinib (P<0.05) alone. All drug treatments, including combinations, were tolerated *in vivo* based on minimal changes in body weights (*data not shown*).

As a second AML cell line-derived xenograft model, we introduced genetically engineered MOLM3/Luc/GFP cells into NSGS mice and initiated treatment as for the OCI-AML3/Luc/GFP model. Again, BLI demonstrated significantly reduced leukemia burden in the treated groups compared to controls and the reduction was more pronounced in the groups treated with single-agent venetoclax or cobimetinib plus venetoclax (Figure 4C, D). Additionally, human CD45 engraftment and cell counts in both bone marrow and spleen demonstrated a trend toward decreased tumor burden in mice treated with the drug combination compared to that in mice treated with either agent alone (Online Supplementary Figure S8A, B). As in the OCI-AML3/Luc xenograft model, all drug treatments were tolerated based on minimal changes in body weights. We performed additional PDX studies in NSG mice using an AML PDX generated from primary sample AML11 (Table 2). The mice were treated with the same doses of drugs as those used in the cell line models. As shown in Online Supplementary Figure S9, the combination therapy extended survival in the AML11 PDX model. These data demonstrate that the combination of cobimetinib plus venetoclax potently suppresses leukemia burden in tumor-bearing mice *in vivo* at tolerable doses.

Discussion

Although gain-of-function mutations often represent secondary events in the pathogenesis of AML,^{31,32} they are required for AML maintenance and are therefore attractive therapeutic targets.³³ While MEK inhibitors have demonstrated limited activity in AML as single agents,^{15,34} preclinical studies with first generation MEK and BCL2 inhibitors demonstrated synergistic induction of apoptosis by suppression of MCL1 following MEK inhibition.^{11,35}

In this study, five of the 11 AML cell lines tested were sensitive to cobimetinib, including two that harbored a *FLT3*-ITD mutation (MOLM14 and MV4;11) and one with an *NRAS* mutation (OCI-AML3). Consistent with previous reports, the baseline levels of ERK phosphorylation did not correlate with response to cobimetinib.^{21,36} Venetoclax as a single agent had activity in five of the cell lines tested while the combination with cobimetinib was synergistic in seven of the cell lines, including those that were resistant to each agent alone.

To extend our preliminary findings in cell lines, we studied a selection of genetically diverse primary AML patients' samples. Venetoclax induced pronounced apoptosis (>50%) in only three of the samples (16.7%), a rate similar to that of clinical responses to venetoclax monotherapy (19%),¹⁰ possibly reflecting protective properties of the tumor microenvironment, as our culture conditions were cytokine-rich. Cobimetinib induced very limited cell death in all AML samples, consistent with previous reports that MEK inhibitors preferentially suppress proliferation and promote differentiation, rather than induce death.^{33,37,38} Remarkably, over 60% of patients' samples responded to the combination therapy, notably including samples that were insensitive to both agents on their own. Moreover, these responders carried diverse genetic alterations that affect leukemia cell proliferation (*FLT3, RAS*), differentiation (*RUNX1*), genomic stability (*NPM1*), and epigenetic modifications (*TET2, IDH1* and *IDH2*). Clonogenic assays demonstrated that the combination markedly impaired the colony-forming functions





of AML progenitors, while normal progenitors were only minimally affected.

CyTOF has proven to be a powerful approach for identifying functional proteins in diverse cell populations at single-cell levels.³⁹ Several groups, including ours, have studied the feasibility of CyTOF in AML.^{26,40,41} In this study, we utilized CyTOF combined with SPADE software⁴² to investigate the efficacy of cobimetinib and venetoclax in two primary patients' samples: one responder and one non-responder. BCL2 was highly expressed in CD34⁺ stem/progenitor cells compared to the CD34⁻ cells, underlying the critical need for BCL2 inhibition to eliminate LSC. The venetoclax-sensitive sample displayed a higher level of BCL2 protein than the resistant sample. Cobimetinib inhibited G-CSF-induced pERK irrespective of response status. In line with several studies reporting that suppression of mTORC1 and its downstream pathways (specifically S6) predicted sensitivity to MEK inhibition,^{21,22} our data also demonstrated that the pS6 signaling pathway was suppressed in the cobimetinib-responding sample, suggesting that S6 phosphorylation may be a more predictive pharmacodynamic marker for MEK inhibition. However, the latter requires validation in a larger cohort of samples from patients.

The distinct response patterns in AML cell lines and patients' samples led us to search for additional pharmacodynamic markers correlating with drug responses using proteomic and transcriptomic profiling. In line with the findings of an extensive study of MEK inhibition,²⁰ we observed bypass induction of pMEK signaling upon MEK inhibition, which was more pronounced in cobimetinibresistant cell lines. Several signaling pathways were highly activated in cobimetinib-sensitive cell lines, including pS6, pERK, p38MAPK, and pPTEN. Lauchle and colleagues demonstrated that leukemia clones with pre-existing resistance to MEK inhibition displayed reduced p38 kinase activity and increased RasGRP1 levels.¹³ It was also previously reported that the RSK signaling pathway, which is downstream of MAPK, regulates an mTOR-independent pathway to induce S6 phosphorylation.⁴³ Western blotting analysis performed to validate the RPPA data showed that S6 phosphorylation at both Ser235/236 and Ser240/244 sites was markedly suppressed in cobimetinib-sensitive OCI-AML3 and MV4-11 cells. In OCI-AML3 cells, the combination treatment resulted in significant cell death characterized by elevated levels of cleaved PARP, which could be attributed to disruption of BCL2:BIM complexes, releasing BIM to trigger apoptosis. We also observed BIM induction in MV4-11 cells, underscoring its critical role in the efficacy of the combination of BCL2 and MEK inhibitors.^{37,38,44} Although RPPA data showed no modulation of MCL1 after cobimetinib treatment, both western blot and Meso Scale Discovery assays showed downregulation of MCL1 in OCL-AML3 cells and upregulation of MCL1 after venetoclax treatment. These data suggest that

increased MCL1 levels induced by venetoclax favor the formation of MCL1:BIM complexes were disrupted, freeing BIM to initiate apoptosis. Consistent with these findings, we recently showed that MCL-1 degradation associated with MDM2 inhibition occurs through MEK/ERK suppression and GSK3 activation.⁴⁵ The downregulation of MYC levels by cobimetinib also suggests a MEK/ERK-GSK3 β link, as ubiquitination and degradation of MYC requires phosphorylation at T58 by GSK3β.⁴⁶ Furthermore, RNA sequencing analyses revealed enhanced expression of MYC and E2F target genes in cells demonstrating a synergistic response to the cobimetinib-venetoclax combination. This finding is consistent with a previous report that MEK inhibition sensitized cells to ABT-263-induced apoptosis by promoting a G1 cell cycle arrest.³⁷ Glycolysis and oxidative phosphorylation (OXPHOS) are known to be regulated by ERK signaling through RNK126-mediated ubiquitination of pyruvate dehydrogenase kinase.³⁰ Potent antitumor efficacy has been demonstrated in melanoma cells through combined inhibition of BCL2, OXPHOS and MAPK signaling.⁴⁷ Alterations in p53 and UPR pathways identified by transcriptome analysis may also account for synergy between MEK and BCL2 inhibition.48 These proposed mechanisms of actions are summarized in Online Supplementary Figure S10. However these models require further validation in controlled mechanistic studies.

The potency of the cobimetinib and venetoclax combination was further demonstrated in vivo using models established with OCI-AML3 (resistant to venetoclax) and MOLM13 (resistant to cobimetinib) leukemia cells. Although we observed strong synergistic effects in both cell lines in vitro, the combination did not confer significant survival benefits in the *in vivo* models. This may be due to protection against cell death provided by the microenvironment, as we have observed in patients' samples cultured in cytokine-rich medium. Similar to our in vitro observations, the OCI-AML3 xenograft model is hypersensitive to cobimetinib, and we found no significant survival differences between animals that received single-agent cobimetinib and those that received the combination. In the very aggressive MOLM13 model, in which untreated mice die 3 weeks after cell injection, the combination reduced but did not eliminate leukemia burden markedly on day 17.

In summary, combinatorial blockade of the MAPK and BCL2 pathways promotes cell death and suppresses proliferation in the majority of primary AML cells. This antileukemia efficacy is associated with the simultaneous inhibition of BCL2 by venetoclax and the downregulation of MCL1 mediated by cobimetinib, which together enable the release of the pro-death protein BIM. These preclinical data provided a strong mechanistic rationale for evaluating the combination of cobimetinib with venetoclax in a phase I trial now enrolling elderly patients with relapsed/refractory AML (NCT02670044), and initial data have included objective clinical responses.⁴⁹

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