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Early changes in rpS6 phosphorylation and BH3 profiling predict response to chemotherapy in AML cells

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Abstract

Blasts from different patients with acute myeloid leukemia (AML) vary in the agent(s) to which they are most responsive. With a myriad of novel agents to evaluate, there is a lack of predictive biomarkers to precisely assign targeted therapies to individual patients. Primary AML cells often survive poorly in vitro, thus confounding conventional cytotoxicity assays. The purpose of this work was to assess the potential of two same-day functional predictive assays in AML cell lines to predict long-term response to chemotherapy. (i) Ribosomal protein S6 (rpS6) is a downstream substrate of PI3K/akt/mTOR/kinase and MAPK kinase pathways and its dephosphorylation is also triggered by DNA double strand breaks. Phospho-rpS6 is reliably measurable by flow cytometry and thus has the potential to function as a biomarker of responsiveness to several therapeutic agents. (ii) A cell's propensity for apoptosis can be interrogated via a functional assay termed “Dynamic BH3 Profiling” in which mitochondrial outer membrane permeabilization in drug-treated cells can be driven by pro-apoptotic BH3 domain peptides such as PUMA-BH3. The extent to which a particular cell is primed for apoptosis by the drug can be determined by measuring the amount of cytochrome C released on addition of BH3 peptide. We demonstrate that phospho-rpS6 expression and PUMA-BH3 peptide-induced cytochrome C release after 4 hours both predict long term chemoresponsiveness to tyrosine kinase inhibitors and DNA double strand break inducers in AML cell lines. We also describe changes in expression levels of the prosurvival BCL-2 family member Mcl-1 and the pro-apoptotic protein BIM after short term drug culture.

Introduction

AML is a heterogeneous clonal disorder of haemopoietic progenitor cells where both failure to differentiate and over proliferation results in accumulation of non-functional cells termed myeloblasts.[1] While nearly 80% of younger AML patients may initially achieve complete remission with current therapy, most will relapse with resistant disease.[2] Clinical outcomes
in the elderly are even more modest as these patients do not tend to tolerate intensive chemotherapy regimens and frequently have adverse cytogenetics.[1]

There are many obstacles a chemotherapeutic drug has to circumvent before it can kill a leukaemia cell. Briefly, the drug has to reach its target, initiate the appropriate pro-apoptotic signals and overcome the cell’s anti-apoptotic defences. Despite the abundance of novel agents that have the potential to improve patient outcome, we are still lacking assays that clinicians can be offered to demonstrate which drugs the individual patient will best respond to. Most efforts in assigning therapy involve predicting a patient’s response to an agent based on their cytogenetic profile. The obvious solution might be chemosensitivity assays, but these have been tried, tested and found wanting.[3, 4] A confounding issue is that cells can be fragile ex vivo, and most AML cells will die spontaneously in culture fairly quickly (median survival at 48 hours = 38% of baseline).[5] Assays have been developed to maintain leukaemia cells in vitro, but AML samples are so heterogeneous that there is no “one-size-fits all” methodology for keeping them alive once they have been isolated from the patient. By focusing on same-day assays, with intact cells, we hope to overcome this obstacle.

Ribosomal protein S6 is a downstream substrate of PI3K/akt/mTOR/ p70S6 kinase and MAPK/p90S6 kinase pathways and is also dephosphorylated following DNA double strand breaks.[6–10] RpS6 is a constituent of the 40S ribosomal subunit that is phosphorylated at several sites including serines 235/236 and 240/244 upon activation by p70S6 and p90S6 kinases. [6] Phosphorylation of rpS6 controls mRNA translation in dividing cells,[11] and its recruitment to the 7-methylguanosine cap structure suggests a role in regulating assembly of the translation preinitiation complex.[8] The AKT and/or MAPK signalling pathways are constitutively active in the majority of AML cases.[12, 13] We have recently determined that rpS6 is hyperphosphorylated in AML patient samples, with phosphorylation being over 20 fold higher than in normal mobilised CD34+ cells.[14] Moreover, rpS6 phosphorylation in patient samples can be abrogated by AKT and/or ERK inhibitors.[15] Some types of AML therapy, such as DNA damaging agents and receptor kinase inhibitors, might be expected to converge to dephosphorylate rpS6 through their actions on the akt/mTOR, ATM/AMPK/mTOR and/or ERK pathways.[9, 16] Phosphorylated rpS6 is therefore a potential biomarker of responsiveness to several therapeutic agents. Antibodies to rpS6 phosphorylated at serine 235/236 have been optimised for flow cytometry, where they are well-established as biomarkers for mTORC1 activity.[17, 18]

The B-Cell Lymphoma-2 (Bcl-2) family of proteins act at the mitochondria and regulate the internal apoptotic pathway. For apoptosis to occur, the proapoptotic effector molecules BAK and BAX must oligomerise and form pores that cause mitochondrial outer membrane permeabilisation (MOMP) resulting in cytochrome c release. Effector molecule activation can be triggered by BH3-only proapoptotic BCL-2 family members such as PUMA, BIM and BID.[19] These pro-apoptotic family members are usually sequestered by BCL-2 family prosurvival members such as MCL-1, BCL-2, and BCL-XL which serves to hold apoptosis in check.[20] A cell’s propensity for apoptosis can be interrogated via a functional assay termed “Dynamic BH3 Profiling”. [21] Dynamic BH3 profiling can predict cellular responses to therapy based on measuring the capacity of drugs to prime mitochondria for apoptosis. The technique involves the addition of permeable pro-apoptotic BH3 peptides to drug primed cells to induce rapid mitochondrial outer membrane permeabilisation. The extent to which a particular cell is primed for apoptosis can be determined by measuring the amount of cytochrome c released. [22] Dynamic BH3 profiling using imatinib (with a 16 hour incubation) was found to predict clinical responsiveness in CML patients.[21]

In this study we assess whether short term (4 hour) rpS6 de-phosphorylation and/or PUMA-BH3 peptide-driven cytochrome c release can predict long-term (48 hour) response to
chemotherapeutic drugs. As primary AML cells are unstable in vitro we utilize a panel of AML cell lines in order to obtain robust 48 hour IC$_{50}$ values for reliable comparison with the short term functional assays. We also investigate whether drug exposure induces rapid changes in expression levels of Bcl-2 protein family members.

**Materials and methods**

**Materials**

Drugs and suppliers used in the study were as follows: 17-AAG, rapamycin, sorafenib, U0126 and torin 1 from LC labs (www.lclabs.com); AC220 and vosaroxin from Selleck (supplied by Stratech UK); etoposide from Tocris; gemtuzumab ozogamicin (GO) was a gift from Wyeth, Pearl River USA. C2 ceramide and Calyculin A were from Santa Cruz Biotechnology, Santa Cruz, CA, USA. Ly294006 was from Millipore, Watford, UK. Other drugs and reagents were from Sigma (Poole, Dorset, UK) unless specified.

**Cells**

OCI-AML3, MOLM-13 and M-07e myeloid leukaemia cell lines were obtained from the German Collection of Microorganisms and Cell Cultures (DSMZ, Braunschweig, Germany). U937 and KG1a cell lines were from the European Collection of Animal Cell Cultures (Salisbury, UK). MV4-11 and TF-1a cells were obtained from the American Type Culture Collection (Manassas, VA, USA). HL-60 cells were a gift from Dawn Bradbury (Nottingham University Hospitals, UK), OCI-AML6.2 cells were a gift from Dr. Jo Mountford (University of Glasgow, UK), M0-91 cells were a gift from Joseph Scandura (Cornell Medical College, USA). OCI-AML DNR cells were developed in our laboratory.[23] HL-60, U937, OCI-AML3, OCI-AML DNR, OCI-AML6.2, MOLM-13, TF-1a, M0-91 and MV4-11 cell lines were maintained in RPMI 1640 medium with 10% foetal calf serum (FCS; First Link, Birmingham, UK), 2mM L-glutamine, 100 U/ml penicillin and 10μg/ml streptomycin. The KG1a and M-07e cell lines were maintained as above with 20% FCS and the M-07e having the addition of 10ng/ml GM-CSF (Novartis, Basel, Switzerland). All cultures were kept at 37°C in 5% CO$_2$ and all experiments were performed with cell lines in log phase. Regular testing to authenticate these cell lines was performed using multiplex short tandem repeat analysis (Powerplex 16; Promega, Southampton, UK). Mycoplasma testing was carried out routinely using the Mycoalert mycoplasma detection kit (Lanza, Rockland, USA) and following the manufacturer’s instructions.

**Chemosensitivity assay**

Cells were plated in triplicate at 2.5x10$^5$/ml with drug or untreated controls in 96 well plates. Plates were incubated for 48 hours at 37°C in 5% CO$_2$ with the addition of alamar blue (Sero-tec, BUF012A) for the final 4 hours. Fluorescence was recorded using a POLARStar optima plate reader (BMG technologies, UK). Cell lines were deemed sensitive or resistant to each agent using the following criteria ($<5$ X 10$^{th}$ centile IC$_{50}$ = sensitive; $>5$ X 10$^{th}$ centile IC$_{50}$ = resistant).

**Phospho-S6 ribosomal protein expression**

Cells were incubated at 5x10$^5$/ml in culture medium for four hours with the indicated drugs. Phospho-S6 ribosomal protein expression (using Alexa-647-conjugated rpS6 p-ser235/236 antibody, CST #4851) was measured following fixation in 2% paraformaldehyde and permeabilisation with 0.1% saponin as described.[14] Baseline rpS6 phosphorylation was determined by culturing with the mTOR inhibitors rapamycin (100 nM) and torin1 (1 μM) and the ERK
inhibitor U0126 (3 μM). Adjustments for baseline rpS6 phosphorylation and expression in untreated cells were made using the calculation \( \% \text{rpS6 dephosphorylation} = 100 - 100 \times (\text{MFI with agent–baseline MFI})/(\text{untreated MFI–baseline MFI}) \), where MFI = mean fluorescence intensity.

**Dynamic BH3 profiling**

Cells were incubated at 5x10^5/ml in culture medium for four hours with the indicated drugs. Cytochrome c release (using Alexa-647-conjugated cytochrome c antibody, Becton Dickinson #558709) was measured after a further 60 minute incubation of digitonin-permeabilised cells with PUMA-BH3 peptide as described.[22, 24] In preliminary assays, the PUMA-BH3 was optimised to 3 μM in all cells except M-091, as this was the concentration of the peptide found to be sufficient to induce mitochondrial outer membrane permeabilisation in drug-primed cells, but not so high that it induced a high degree of mitochondrial outer membrane permeabilisation without drugs. In M-091, the peptide was used at 0.5 μM. Adjustments for peptide-induced cytochrome c release in untreated cells were made in order to establish agent-specific release, using the formula 100X (percent cytochrome c positive with peptide–percent cytochrome c positive with drug plus peptide)/(percent cytochrome c positive with peptide). A mutated PUMA-BH3 peptide (PUMA2A) [22] at 100 μM and BIM-BH3 peptide (10 μM) were used as controls in all experiments. Data were collected on a FACSCanto II flow cytometer (Becton Dickinson) and analysed with FACS Diva software (Becton Dickinson).

**Western blot analysis**

MV4-11 cells were treated for four hours with 1 μM etoposide, 10 nM AC220 or 1 μM torin1. Cell lysates were prepared, separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis, and transferred to nitrocellulose membranes. Detection antibodies included anti-β-Actin (N-21, sc-130656), anti-MCL1 (S-19, sc-819) and anti-Bcl-2 (C-2, sc-7382) from Santa Cruz Biotechnology, Santa Cruz, CA, USA, anti-BID (#2002, Cell Signalling Technology, Leiden, Netherlands), anti-BIM (Y36, ab32158), anti-BAD (phospho S136, ab28824) and anti-PUMA (EP512Y, ab33906) from Abcam, Cambridge, UK.

**Calculations and statistics**

Statistical analysis was carried out using the Statistical Package for Social Sciences, version 23 (SPSS, Chicago, IL, USA). P values of ≤0.05 were considered to represent significance. Area under curve (AUC) receiver-operating characteristic (ROC) predictive analysis was used to evaluate the relationship between sensitivity of cell lines to an agent and rpS6 phosphorylation or cytochrome c release.[25]

**Results**

**Multiple signalling pathways converge on rpS6 phosphorylation**

The MAPK and/or AKT signalling pathways are constitutively active in the majority of AML cases.[12, 13] RpS6 is a downstream mediator of both pathways,[7, 8] and is constitutively phosphorylated in patient samples.[14] P-rpS6 has been used previously as a sensitive marker for mTORC1 activation in flow cytometric studies.[15, 17, 18] To examine basal phosphorylation and inhibitor-induced dephosphorylation of rpS6 in our assay system, we cultured MV4-11s for 4 hours with the mTORC1 inhibitor torin1, the ERK inhibitor U0126 and the akt inhibitor LY294002. All these agents were able to greatly reduce rpS6 phosphorylation (Fig 1). The
Fig 1. RpS6 dephosphorylation by signalling inhibitors. (A) MV4-11 cells were cultured for 4 hours with 1 μM torin1, 5 μM U0126, 30 μM LY294002, 30 μM C2-ceramide or a combination of all 4 inhibitors. (B) Values are a percent of untreated cell phospo-rpS6 as described in the methods. (Mean +/- SD for n = 3–5). The phosphatase inhibitor Calyculin A (3 nM) was used as positive control.

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cytotoxic drug-activated signalling lipid ceramide [26–28] was also inhibitory. In contrast, the phosphatase inhibitor calyculin A enhanced S6 phosphorylation.

**Rps6 dephosphorylation at 4 hours predicts 48 hour drug sensitivity**

Data from Fig 1 suggests early changes in rps6 phosphorylation may be used as a predictor of chemotherapeutic response in AML cells. To test this hypothesis, 11 AML cell lines were subjected to 48 hour dose response assays with the following agents: FLT3 inhibitors (AC220 and sorafenib),[29, 30] DNA double strand break (DSB) inducers (etoposide, GO and vosaroxin),[31–33] the standard-of-care drug cytarabine and the hsp90 inhibitor 17-AAG.[34]

Using ROC analysis (predictive accuracy test), rpS6 dephosphorylation at 4 hours predicted the 48 hour response to the DSB inducing drugs etoposide and GO, the FLT3 inhibitors sorafenib and ACC220 as well as to cytarabine (Fig 2 and S1 Table). ROC analysis of 4 hour rpS6 dephosphorylation compared to 48 hour drug sensitivity confirmed that the assay was highly sensitive and specific with AUC values of 1.0 for etoposide, GO, sorafenib, AC220 and cytarabine (S1 Fig shows individual ROC curves). An AUC value close to 1 indicates excellent predictive capabilities providing evidence that this laboratory test performed after 4 hours does have an ability to predict chemosensitivity at 48 hours. For vosaroxin, the AUC value was 0.82. HL-60 cells, which were sensitive to vosaroxin at 48 hours, did not dephosphorylate rpS6.

17-AAG, which had been included because of clinical interest in novel hsp90 inhibitors,[35] was expected to be particularly effective because hsp90 plays a direct role in maintaining the stability of rpS6.[36] The predictive value for 17-AAG (AUC 0.9) was statistically significant, albeit with U937 cells showing confoundingly high dephosphorylation.

**PUMA-BH3 peptide-induced cytochrome c release after 4 hours drug treatment predicts 48 hour drug sensitivity**

Dynamic BH3 profiling involves exposing mitochondria to BH3 domain derived peptides following short term drug exposure to prime mitochondria for changes in MOMP as measured by cytochrome c release. It has been demonstrated that this technique can predict cytotoxicity.[21] PUMA-BH3 peptide can sensitisise all the anti-apoptotic BCL-2 family of proteins[37] and was used for our assay. To obtain optimal sensitivity with this technique it is important to establish suitable drug and peptide concentrations so that agent or peptides do not overwhelm the system by inducing too much cytochrome c release individually. As the technique demands outer membrane permeabilisation whilst maintaining mitochondrial integrity, we set up appropriate assay controls and defined their interpretation as >90% induced cytochrome c release in the presence of a high concentration (10 μM) of the activator peptide BIM-BH3 but less than 10% when incubated with the non-specific peptide PUMA2A-BH3. PUMA-BH3 peptide-induced cytochrome c release closely predicted the 48 hour IC₅₀ for DSB inducing drugs and FLT3 inhibitors, with AUC values of 1.0 for etoposide, GO, AC220 and sorafenib (Fig 3 and S1 Table). For vosaroxin, the AUC value was 0.95. As with the rpS6 assay, HL-60 cells, which were sensitive to vosaroxin at 48 hours, were insensitive in the short-term assay. BH3 profiling was not predictive for responses to 17-AAG, (AUC value 0.63), or cytarabine (AUC value 0.77). See S2 Fig for individual ROC curves.

Summarising the ROC analysis across the 11 cell lines and 7 drugs (Fig 4), both rpS6 dephosphorylation and dynamic BH3 profiling assays showed highly significant predictive ability.
Fig 2. RpS6 dephosphorylation at 4 hours predicts 48 hour drug sensitivity. (A) MOLM-13 cells were cultured for 4 hours with 100 nM Rapamycin, 1 μM torin1 and 3 μM U0126 to determine p-rpS6 baseline. RpS6 dephosphorylation after 4 hours culture with 500 nM 17-AAG is also shown. Example histograms are representative of 3 individual experiments. (B) Based on the 48 hour IC50 values a drug sensitive/resistant cut off for cell lines was determined. Cell lines were cultured for 4 hours with 1 μM etoposide, 50nM sorafenib, 600ng/ml GO, 10nM AC220, 1 μM vosaroxin, 500nM 17-AAG and 2 μM cytarabine. Values are a percent of untreated cell p-rpS6 as described in the methods. Each point represents a cell line and is the product of three individual experiments.

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Changes in expression of apoptotic modulator proteins after four hours drug exposure

For apoptosis to occur, the effector molecules BAK and BAX oligomerise and form pores that cause MOMP, resulting in cytochrome c release. Effector molecule activation can be triggered by the BH3-only proapoptotic BCL-2 family members BIM, BID and PUMA. Prosurvival members such as MCL-1, BCL-2 and BCL-X_L inhibit the BH3-only proteins by sequestration and hold apoptosis in check. Our dynamic BH3 profiling results clearly demonstrate that drugs are priming cells to PUMA-BH3 peptide after only 4 hours exposure. We investigated what effect the drugs were having on BCL-2 apoptotic protein family members during this time period that might underpin this rapid priming. MV4-11 cells were used as these cells were the most sensitive cell line to the majority of drugs in our panel (S1 Table). These cells do not over-express the anti-apoptotic protein BCL-X_L.\(^2\) Of the other BCL-2 family prosurvival members MCL-1 has the shorter half-life (approximately 1 hour) and can be rapidly...

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**Fig 3.** PUMA-BH3 peptide-induced cytochrome c release after 4 hours drug treatment predicts 48 hour drug sensitivity.

(A) MOLM-13 cells were cultured for 4 hours with 1 μM vosaroxin or 50 nM sorafenib followed by 1 hour treatment with PUMA-BH3 peptide or PUMA2A control. Example dot plots are representative of 3 individual experiments. (B) Based on the 48 hour IC\(_{50}\) values a drug sensitive/resistant cut off for cell lines was determined as described in the methods. Cell lines were cultured for 4 hours with 1 μM etoposide, 50nM sorafenib, 600ng/ml GO, 10nM AC220, 1 μM vosaroxin, 500nM 17-AAG or 2 μM cytarabine followed by 1 hour incubation with PUMA-BH3 peptide. Values are corrected for cytochrome c release with PUMA2A control peptide as described in the methods. Each point represents a cell line and is the product of three individual experiments.

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**Fig 4.** ROC curve analysis confirms highly significant overall sensitivity and specificity for the ability of both rpS6 dephosphorylation and PUMA-BH3 induced cytochrome c release after 4 hours drug treatment to predict 48 hour sensitivity to drugs. Summary ROC curves for percent change in rpS6 phosphorylation and PUMA induced cytochrome c release after 4 hours treatment with 1 μM etoposide, 50nM sorafenib, 600ng/ml GO, 10nM AC220, 1 μM vosaroxin, 500nM 17-AAG or 2 μM cytarabine in 11 AML cells lines. Cell lines were classified as sensitive or resistant according to 48 hours drug response (The standardised definition of sensitivity is described in the methods section). Each data point used to generate the analysis is the mean of three individual experiments.

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downregulated [38] whilst BCL-2 is a much more stable protein. [37, 39] We used etoposide and AC220, as DSB-inducing agents and FLT3 inhibitors are of particular clinical interest in AML: both rpS6 dephosphorylation and dynamic BH3 profiling had been able to predict response to these drugs (S1 Table). We found that etoposide and AC220 significantly depletes MCL-1 in the MV4.11 cells without affecting the expression of BCL-2 (Fig 5). The mTORC1 antagonist and translation inhibitor torin1 [40] was included as a control because inhibition of translation depletes MCL-1. [41] Activation of effector molecules BAK and BAX using conformation-specific antibodies as previously described [24] was not observed (data not shown). Of the activator BCL-2 family members BID, BIM and PUMA, BIM expression was significantly increased after 4 hours treatment with AC220 only whilst BID and PUMA were unaffected by either etoposide or AC220 (Fig 5). See S3 Fig for uncropped blots.

Discussion

This work is a first step towards ascertaining whether short term functional predictive assays might have clinical application in AML. A history of failure for chemosensitivity assays in clinical practice [3] has been discouraging, but the need for the development of such assays is still being strongly asserted, [4, 42, 43] with the rationale that molecular assays in unstimulated cells cannot recapitulate complex behaviour.

We have demonstrated the potential of two same-day functional flow cytometric assays to predict 48-hour response to chemotherapy in AML cell lines. FLT3 inhibitors are documented to induce rapid changes in signalling pathways [44–46] and we have shown here that rpS6 dephosphorylation is a suitable indicator of this activity. Early (i.e. less than four hours) pro-apoptotic changes invoked by DNA damaging drugs are rarely documented. However, in an exception to the general focus on late changes, Nijhawan and colleagues documented the early loss of MCL-1 after genotoxic stress, and they included etoposide in their analysis. [47] The rapidity of ceramide induction following DNA damage [26, 27] and the involvement of ceramide in phosphatase activation [48] also encouraged us to postulate that we might be able to document responses to DNA-damaging agents after four hours. The cell lines were appropriately classified as sensitive or resistant to etoposide and GO. Vosaroxin sensitivity was also predicted by both assays in 10/11 cell lines.

The main obstacle for further development of this work in clinical samples is that primary samples are unstable over time in vitro [5] and therefore cannot be used for proof of principle. Bolt-on studies in the context of clinical trials will be needed to properly evaluate the methodologies. Since AML samples can be hypocellular at presentation, the availability of cells can be a problem for diagnostic assays. Phospho-rpS6 can be performed on very small samples with minimal cell loss during processing. However, some primary samples have low basal rpS6 phosphorylation. [14] BH3 profiling requires permeabilisation of the outer cell membrane, while maintaining viable mitochondria: this is technically challenging [22] and the cell requirement includes positive and negative controls for permeabilisation as well as no peptide controls for each drug. BIM-BH3 has been used by others for profiling without drugs. [49, 50] Our choice of PUMA-BH3 rather than BIM-BH3 was made because PUMA gave a more consistent low baseline level of cytochrome release at a single concentration across several cell lines in preliminary experiments. In performing the assay in 11 cell lines for this study, we found that one (M0-91) was hypersensitive to PUMA-BH3 alone (see methods) and the whole assay had to be repeated with a lower peptide concentration. In summary, BH3 profiling requires more cells and may also require sample-specific optimisation of baseline conditions. The area under the curve in Fig 4 was lower for BH3 profiling than for rpS6 phosphorylation. However, dynamic BH3 profiling is the more logical assay for early chemosensitivity testing as it takes
into account apoptosis resistance that might occur separately from or downstream of pRS6 inhibition.

Perturbations of signalling pathways can induce drug resistance mechanisms,[51, 52] but the finding that rpS6 dephosphorylation at 4 hours was predictive of cell death at 48 hours implies early, irreversible changes. In the final part of the study we probed for early drug-induced changes in BCL-2 family proteins that might indicate commitment to apoptosis. Apoptosis is effected by oligomerisation of BAX and BAK and activated by BCL-2 family members such as BIM, BID and PUMA.[19] The only change we noted in these pro-apoptotic molecules was induction of BIM by AC220 in sensitive MV4-11 cells (Fig 5). We found significant early inhibition of MCL-1 by AC220 in these cells (Fig 5). This might be predicated on early down-regulation of STAT5 signalling: [53] activation of this pathway is documented to maintain MCL-1 expression.[54] However, MCL-1 was only 20% decreased in etoposide-treated cells. Given that elimination of MCL1 is reported to be required for the initiation of apoptosis following double strand breaks[47] the delineation of early irreversible pre-apoptotic changes will likely require attention to post-translational changes and binding partners as well as expression levels of MCL-1. MCL-1 is sequentially translocated to mitochondria, phosphorylated and ubiquitinylated prior to degradation.[55] A possible explanation for the clear-cut predictive value of the assays in the face of resistance mechanisms is the homogeneity of cell lines. A future application of the system might be to elucidate whether sensitive and resistant subsets within patient samples can be defined.

In summary we have described two assays which detect changes occurring in sensitive cells within four hours of drug application and which predict sensitivity and resistance to DNA damaging agents and FLT3 inhibitors in a panel of AML cell lines.

Supporting information
S1 Table. Cell line IC50, rpS6 and cytochrome C data. 11 cell lines were treated with 7 different drugs or untreated controls for 48 hours to determine an IC50 (nM) (shown in bold;
superscript R = resistant, superscript S = sensitive). Percent rpS6 dephosphorylation (regular font) and percent PUMA induced cytochrome C release (bold italic) was determined in the same cell lines after 4 hours drug treatment. Each value is the product of three individual experiments.

**S1 Fig. Individual ROC curves for rpS6 dephosphorylation after 4 hours drug treatment.** ROC curves for percent change in rpS6 phosphorylation after 4 hours treatment with 1 μM etoposide, 50nM sorafenib, 600ng/ml GO, 10nM AC220, 1 μM vosaroxin, 500nM 17-AAG or 2 μM cytarabine in 11 AML cells lines. Each data point used to generate the analysis is the mean of three individual experiments.

(TIF)

**S2 Fig. Individual ROC curves for PUMA induced cytochrome c release after 4 hours drug treatment.** ROC curves for PUMA induced cytochrome c release after 4 hours treatment with 1 μM etoposide, 50nM sorafenib, 600ng/ml GO, 10nM AC220, 1 μM vosaroxin, 500nM 17-AAG or 2 μM cytarabine in 11 AML cells lines. Each data point used to generate the analysis is the mean of three individual experiments.

(TIF)

**S3 Fig. Original uncropped western blots.** MV4-11 cells were treated for four hours with 1 μM etoposide, 10 nM AC220 or 1 μM torin1 before probing for the apoptotic modulator proteins Mcl-1, Bcl-2, BIM, PUMA and BID.

(TIF)

**Author Contributions**

**Conceptualization:** Monica Pallis.

**Data curation:** Martin Grundy, Thomas Jones, Liban Elmi, Michael Hall, Adam Graham, Monica Pallis.

**Formal analysis:** Martin Grundy.

**Funding acquisition:** Martin Grundy, Nigel Russell.

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**Writing – original draft:** Martin Grundy, Monica Pallis.

**Writing – review & editing:** Martin Grundy, Thomas Jones, Liban Elmi, Michael Hall, Adam Graham, Monica Pallis.

**References**

1. Estey E. Acute myeloid leukemia: 2016 Update on risk-stratification and management. American journal of hematology. 2016; 91(8):824–46. Epub 2016/07/16. https://doi.org/10.1002/ajh.24439 PMID: 27417880.

2. Burnett A, Wetzler M, Lowenberg B. Therapeutic advances in acute myeloid leukemia. Journal of clinical oncology : official journal of the American Society of Clinical Oncology. 2011; 29(5):487–94. Epub 2011/01/12. https://doi.org/10.1200/jco.2010.30.1820 PMID: 21220605.
3. Burstein HJ, Mangu PB, Somerfield MR, Schrag D, Samson D, Holt L, et al. American Society of Clinical Oncology clinical practice guideline update on the use of chemotherapy sensitivity and resistance assays. Journal of clinical oncology : official journal of the American Society of Clinical Oncology. 2011; 29(24):3328–30. Epub 2011/07/27. https://doi.org/10.1200/jco.2011.36.0354 PMID: 21788567.

4. Friedman AA, Letai A, Fisher DE, Flaherty KT. Precision medicine for cancer with next-generation functional diagnostics. Nature reviews Cancer. 2015; 15(12):747–56. Epub 2015/11/06. https://doi.org/10.1038/nrc4015 PMID: 26536825; PubMed Central PMCID: PMCPMC4970460.

5. Pallis M, Turzanski J, Grundy M, Seedhouse C, Russell N. Resistance to spontaneous apoptosis in acute myeloid leukaemia blasts is associated with p-glycoprotein expression and function, but not with the presence of FLT3 internal tandem duplications. British journal of haematology. 2003; 120(6):1009–16. Epub 2003/03/22. PMID: 12648071.

6. Holz MK, Blenis J. Identification of S6 kinase 1 as a novel mammalian target of rapamycin (mTOR)-phosphorylating kinase. The Journal of biological chemistry. 2005; 280(28):26089–93. Epub 2005/05/21. https://doi.org/10.1074/jbc.M504045200 PMID: 15905173.

7. Ma L, Chen Z, Erdjument-Bromage H, Tempst P, Pandolfi PP. Phosphorylation and functional inactivation of TSC2 by Erk implications for tuberous sclerosis and cancer pathogenesis. Cell. 2005; 121(2):179–93. Epub 2005/04/27. https://doi.org/10.1016/j.cell.2005.02.031 PMID: 15851026.

8. Roux PP, Shahbazian D, Vu H, Holz MK, Cohen MS, Taunton J, et al. RAS/ERK signaling promotes site-specific ribosomal protein S6 phosphorylation via RSK and stimulates cap-dependent translation. The Journal of biological chemistry. 2007; 282(19):14056–64. Epub 2007/03/16. https://doi.org/10.1074/jbc.M700906200 PMID: 17360704; PubMed Central PMCID: PMCPMC3618456.

9. Cam M, Bid HK, Xiao L, Zambetti GP, Houghton PJ, Cam H. p53/TAp63 and AKT regulate mammalian target of rapamycin complex 1 (mTORC1) signaling through two independent parallel pathways in the presence of DNA damage. The Journal of biological chemistry. 2014; 289(7):4083–94. Epub 2013/12/25. https://doi.org/10.1074/jbc.M113.530303 PMID: 24366874; PubMed Central PMCID: PMCPMC3924274.

10. Li Y, Mitsuhashi S, Ikejo M, Miura N, Kawamura T, Hamakubo T, et al. Relationship between ATM and ribosomal protein S6 revealed by the chemical inhibition of Ser/Thr protein phosphatase type 1. Bioscience, biotechnology, and biochemistry. 2012; 76(3):486–94. Epub 2012/03/28. https://doi.org/10.1271/bbb.110774 PMID: 22451389.

11. Bandi HR, Ferrari S, Krieg J, Meyer HE, Thomas G. Identification of 40 S ribosomal protein S6 phosphorylation sites in Swiss mouse 3T3 fibroblasts stimulated with serum. The Journal of biological chemistry. 1993; 268(6):4530–3. Epub 1993/02/25. PMID: 8440735.

12. Fransecky L, Mochmann LH, Baldus CD. Outlook on PI3K/AKT/mTOR inhibition in acute leukemia. Molecular and cellular therapeutics. 2015; 3:2. Epub 2015/06/10. https://doi.org/10.1186/s40591-015-0040-8 PMID: 26056603; PubMed Central PMCID: PMCPMC4452048.

13. Milella M, Kornblau SM, Estrov Z, Carter BZ, Lapillonne H, Harris D, et al. Therapeutic targeting of the MEK/ERK signal transduction module in acute myeloid leukemia. The Journal of clinical investigation. 2001; 108(6):851–9. Epub 2001/09/19. https://doi.org/10.1172/JCI12807 PMID: 11560954; PubMed Central PMCID: PMCPMC2009930.

14. Pallis M, Harvey T, Russell N. Phenotypically Dormant and Immature Leukaemia Cells Display Increased Ribosomal Protein S6 Phosphorylation. PloS one. 2016; 11(3):e0151480. Epub 2016/03/18. https://doi.org/10.1371/journal.pone.0151480 PMID: 26985829; PubMed Central PMCID: PMCPMC4795744.

15. Chow S, Minden MD, Hedley DW. Constitutive phosphorylation of the S6 ribosomal protein via mTOR and ERK signaling in the peripheral blasts of acute leukemia patients. Experimental hematology. 2006; 34(9):1183–91. Epub 2006/08/31. https://doi.org/10.1016/j.exphem.2006.05.002 PMID: 16939811.

16. Gunawardane RN, Nenomuceno RR, Rooks AM, Hunt JP, Ricono JM, Belli B, et al. Transient exposure to quizzartinib mediates sustained inhibition of FLT3 signaling while specifically inducing apoptosis in FLT3-activated leukemia cells. Molecular cancer therapeutics. 2013; 12(4):438–47. Epub 2013/02/16. https://doi.org/10.1158/1078-0432.ccr-12-0305 PMID: 23412931.

17. Dietertlen MT, Bittner HB, Klein S, von Salisch S, Mittag A, Tarnok A, et al. Assay validation of phosphorylated S6 ribosomal protein for a pharmacodynamic monitoring of mTOR-inhibitors in peripheral human blood. Cytometry Part B, Clinical cytometry. 2012; 82(3):151–7. Epub 2012/01/04. https://doi.org/10.1002/cyto.b.21005 PMID: 22213594.

18. Perl AE, Kasner MT, Shank D, Luger SM, Carroll M. Single-cell pharmacodynamic monitoring of S6 ribosomal protein phosphorylation in AML blasts during a clinical trial combining the mTOR inhibitor sirolimus and intensive chemotherapy. Clinical cancer research : an official journal of the American Association for Cancer Research. 2012; 18(6):1716–25. Epub 2011/12/15. https://doi.org/10.1158/1078-0432.ccr-11-2346 PMID: 22167413; PubMed Central PMCID: PMCPMC3306511.
19. Czabotar PE, Lessene G, Strasser A, Adams JM. Control of apoptosis by the BCL-2 protein family: implications for physiology and therapy. Nature reviews Molecular cell biology. 2014; 15(1):49–63. Epub 2013/12/21. https://doi.org/10.1038/nrm3722 PMID: 24355989.

20. Chipuk JE, Green DR. How do BCL-2 proteins induce mitochondrial outer membrane permeabilization? Trends in cell biology. 2008; 18(4):157–64. Epub 2008/03/04. https://doi.org/10.1016/j.tcb.2008.01.007 PMID: 18314333; PubMed Central PMCID: PMC3242477.

21. Montero J, Sarosiek KA, DeAngelo JD, Maertens O, Ryan J, Ercan D, et al. Drug-induced death signaling strategy rapidly predicts cancer response to chemotherapy. Cell. 2015; 160(5):977–89. Epub 2015/02/28. https://doi.org/10.1016/j.cell.2015.01.042 PMID: 25723171; PubMed Central PMCID: PMC4391197.

22. Ryan J, Letal A. BH3 profiling in whole cells by fluorimeter or FACS. Methods (San Diego, Calif). 2013; 61(2):156–64. Epub 2013/04/24. https://doi.org/10.1016/j.jytemth.2013.04.006 PMID: 23607990; PubMed Central PMCID: PMC3686919.

23. Grundy M, Seedhouse C, Russell NH, Pallis M. P-glycoprotein and breast cancer resistance protein in acute myeloid leukaemia cells treated with the aurora-B kinase inhibitor barasertib-hQPA. BMC cancer. 2011; 11:254. Epub 2011/06/18. https://doi.org/10.1186/1471-2407-11-254 PMID: 21679421; PubMed Central PMCID: PMC3146447.

24. Pallis M, Burrows F, Ryan J, Grundy M, Seedhouse C, Abdul-Aziz A, et al. Complementary dynamic BH3 profiles predict co-operativity between the multi-kinase inhibitor TG02 and the BH3 mimetic ABT-199 in acute myeloid leukaemia cells. Oncotarget. 2017; 8(10):16220–32. Epub 2016/04/20. https://doi.org/10.18632/oncotarget.8742 PMID: 27092880; PubMed Central PMCID: PMC5369858.

25. Zweig MH, Campbell G. Receiver-operating characteristic (ROC) plots: a fundamental evaluation tool in clinical medicine. Clinical chemistry. 1993; 39(4):561–77. Epub 1993/04/01. PMID: 8472349.

26. Perry DK, Carton J, Shah AK, Meredith F, Uhlinger DJ, Hannun YA. Serine palmitoyltransferase regulates de novo ceramide generation during etoposide-induced apoptosis. The Journal of biological chemistry. 2000; 275(12):9078–84. Epub 2000/03/18. PMID: 10722793.

27. Sawada M, Nakashima S, Banno Y, Yamakawa H, Hayashi K, Takenaka K, et al. Ordering of ceramide formation, caspase activation, and Bax/Bcl-2 expression during etoposide-induced apoptosis in C6 glioma cells. Cell death and differentiation. 2000; 7(9):761–72. Epub 2000/10/24. https://doi.org/10.1038/sj.cdd.4400711 PMID: 11042671.

28. Tepper AD, de Vries E, van Blitterswijk WJ, Borst J. Ordering of ceramide formation, caspase activation, and mitochondrial changes during CD95- and DNA damage-induced apoptosis. The Journal of clinical investigation. 1999; 103(7):971–8. Epub 1999/04/09. https://doi.org/10.1172/JCI5457 PMID: 10194469; PubMed Central PMCID: PMC408258.

29. Lyons JF, Wilhelm S, Hibner B, Bollag G. Discovery of a novel Raf kinase inhibitor. Endocrine-related cancer. 2001; 8(3):219–25. Epub 2001/09/22. PMID: 11566613.

30. Zarrinkar PP, Gunawardane RN, Cramer MD, Gardner MF, Brigham D, Belli B, et al. AC220 is a uniquely potent and selective inhibitor of FLT3 for the treatment of acute myeloid leukemia (AML). Blood. 2009; 114(14):2984–92. Epub 2009/08/06. https://doi.org/10.1182/blood-2009-05-222034 PMID: 19654408; PubMed Central PMCID: PMC2756206.

31. Howlin RE, Stockett DE, Wong OK, Lundin C, Helleday T, Fox JA. Homologous recombination repair is essential for repair of vosaroxin-induced DNA double-strand breaks. Oncotarget. 2010; 1(7):606–19. Epub 2011/02/15. https://doi.org/10.18632/oncotarget.101106 PMID: 21317456; PubMed Central PMCID: PMC3248135.

32. Vock EH, Lutz WK, Hormes P, Hoffmann HD, Varnavas S. Discrimination between genotoxicity and cytotoxicity in the induction of DNA double-strand breaks in cells treated with etoposide, melphalan, cisplatin, potassium cyanide, Triton X-100, and gamma-irradiation. Mutation research. 1998; 413(1):83–94. Epub 1998/02/29. PMID: 9602861.

33. Yamauchi T, Matsuda Y, Tasaki T, Negoro E, Ikegaya S, Takagi K, et al. Ordering of ceramide generation, caspase activation, and mitochondrial changes during CD95- and DNA damage-induced apoptosis. The Journal of biological chemistry. 2001; 276(25):27072696; PubMed Central PMCID: PMC4995156.

34. Kelland LR, Sharp SY, Rogers TM, Myers TG, Workman P. DT-Diaphorase expression and tumor cell sensitivity to 17-allylamino, 17-demethoxygeldanamycin, an inhibitor of heat shock protein 90. Journal of the National Cancer Institute. 1999; 91(22):1940–9. Epub 1999/11/24. PMID: 10564678.

35. Shrestha L, Bolaender A, Patel HJ, Taldone T. Heat Shock Protein (HSP) Drug Discovery and Development: Targeting Heat Shock Proteins in Disease. Current topics in medicinal chemistry. 2016; 16(25):2753–64. Epub 2016/04/14. PMID: 27072696; PubMed Central PMCID: PMC4995156.

36. Kim TS, Jang CY, Kim HD, Lee JY, Ahn BY, Kim J. Interaction of Hsp90 with ribosomal proteins protects from ubiquitination and proteasome-dependent degradation. Molecular biology of the cell. 2006; 17
Kapoor S, Natarajan K, Baldwin PR, Doshi KA, Lapidus RG, Mathias TJ, et al. Concurrent inhibition of
53.
52.
Bruner JK, Ma HS, Li L, Qin ACR, Rudek MA, Jones RJ, et al. Adaptation to TKI Treatment Reactivates
45.
Spiekermann K, Bagrintseva K, Schwab R, Schmieja K, Hiddemann W. Overexpression and constitu-
Nijhawan D, Fang M, Traer E, Zhong Q, Gao W, Du F, et al. Elimination of Mcl-1 is required for the initia-
47.
Grundy M, Seedhouse C, Shang S, Richardson J, Russell N, Pallis M. The FLT3 internal tandem dupli-
46.
Montero J, Letai A. Dynamic BH3 profiling—poking cancer cells with a stick. Molecular & cellular oncol-
45.
Kitada S, Miyashita T, Tanaka S, Reed JC. Investigations of antisense oligonucleotides targeted
44.
Levis M, Tse KF, Smith BD, Garrett E, Small D. A FLT3 tyrosine kinase inhibitor is selectively cytotoxic
to acute myeloid leukemia blasts harboring FLT3 internal tandem duplication mutations. Blood. 2001;
98(3):885–7. Epub 2001/07/27. PMID: 11468194.
43.
Spiekermann K, Bagrintseva K, Schwab R, Schmieja K, Hiddemann W. Overexpression and constitutive
activation of FLT3 induces STAT5 activation in primary acute myeloid leukemia blast cells. Clinical
cancer research : an official journal of the American Association for Cancer Research. 2003; 9(6):2140–
50. Epub 2003/06/11. PMID: 12796379.
42.
Grundy M, Seedhouse C, Shang S, Richardson J, Russell N, Pallis M. The FLT3 internal tandem dupli-
cation mutation is a secondary target of the aurora B kinase inhibitor AZD1152-HQPA in acute myeloge-
nous leukemia cells. Molecular cancer therapeutics. 2010; 9(3):661–72. Epub 2010/02/18. https://doi.
or/10.1158/1535-7163.MCT-09-1144 PMID: 20599992.
41.
Nijhawan D, Fang M, Traer E, Zhong Q, Gao W, Du F, et al. Elimination of Mcl-1 is required for the initia-
tion of apoptosis following ultraviolet irradiation. Genes & development. 2003; 17(12):1475–86. Epub
1995/03/01. PMID: 8796880; PubMed Central PMCID: PMCPMC2120408.
39.
Kitada S, Miyashita T, Tanaka S, Reed JC. Investigations of antisense oligonucleotides targeted against
bcl-2 RNAs. Antisense research and development. 1993; 3(2):157–69. Epub 1993/01/01. PMID:
8400801.
38.
Thoreen CC, Kang SA, Chang JW, Liu Q, Zhang J, Gao Y, et al. An ATP-competitive mammalian target
of rapamycin inhibitor reveals rapamycin-resistant functions of mTORC1. The Journal of biological
chemistry. 2009; 284(12):8023–32. Epub 2009/01/20. https://doi.org/10.1074/jbc.M900301200 PMID:
19150980; PubMed Central PMCID: PMCPMC2658096.
37.
Rooswinkel RW, van de Kooij B, de Vries E, Pauwe M, Braster R, Verheij M, et al. Antiapoptotic
potency of Bcl-2 proteins primarily relies on their stability, not binding selectivity. Blood. 2012; 123
(18):2806–15. Epub 2014/03/14. https://doi.org/10.1182/blood-2013-08-519470 PMID: 24622325.
36.
Yang T, Kozopas KM, Craig RW. The intracellular distribution and pattern of expression of Mcl-1 over-
association for Cancer Research. 2017. Epub 2017/10/28. https://doi.org/10.1101/gad.1093903
PMCID: PMC5901262.
35.
Oaks J, Ogretmen B. Regulation of PP2A by Sphingolipid Metabolism and Signaling. Frontiers in oncol-
chemistry. 2009; 284(12):8023–32. Epub 2009/01/20. https://doi.org/10.1074/jbc.M900301200 PMID:
19150980; PubMed Central PMCID: PMCPMC2658096.
34.
Pan R, Hogdal LJ, Benito JM, Bucci D, Han L, Borthakur G, et al. Selective BCL-2 inhibition by ABT-199
causes on-target cell death in acute myeloid leukemia. Cancer discovery. 2014; 4(3):362–75. Epub
https://doi.org/10.1158/1940-6207.crd-16-0305. Epub 2016/11/24. PMID:
27314085; PubMed Central PMCID: PMCPMC4909438.
33.
Yaffe MB. The scientific drunk and the lamppost: massive sequencing efforts in cancer discovery and
treatment. Science signaling. 2013; 6(269):pe13. Epub 2013/04/04. https://doi.org/10.1126/scisignal.
2003684 Epub 2015/02/03.
32.
Levis M, Tse KF, Smith BD, Garrett E, Small D. A FLT3 tyrosine kinase inhibitor is selectively cytotoxic
to acute myeloid leukemia blasts harboring FLT3 internal tandem duplication mutations. Blood. 2001;
98(3):885–7. Epub 2001/07/27. PMID: 11468194.
31.
Spiekermann K, Bagrintseva K, Schwab R, Schmieja K, Hiddemann W. Overexpression and constitutive
activation of FLT3 induces STAT5 activation in primary acute myeloid leukemia blast cells. Clinical
cancer research : an official journal of the American Association for Cancer Research. 2003; 9(6):2140–
50. Epub 2003/06/11. PMID: 12796379.
30.
Grundy M, Seedhouse C, Shang S, Richardson J, Russell N, Pallis M. The FLT3 internal tandem dupli-
cation mutation is a secondary target of the aurora B kinase inhibitor AZD1152-HQPA in acute myeloge-
nous leukemia cells. Molecular cancer therapeutics. 2010; 9(3):661–72. Epub 2010/02/18. https://doi.
or/10.1158/1535-7163.MCT-09-1144 PMID: 20599992.
29.
Nijhawan D, Fang M, Traer E, Zhong Q, Gao W, Du F, et al. Elimination of Mcl-1 is required for the initia-
tion of apoptosis following ultraviolet irradiation. Genes & development. 2003; 17(12):1475–86. Epub
2003/06/05. https://doi.org/10.1101/gad.1093903 PMID: 12783855; PubMed Central PMCID:
PMCPMC196078.
28.
Oaks J, Ogretmen B. Regulation of PP2A by Sphingolipid Metabolism and Signaling. Frontiers in oncol-
chemistry. 2014; 4:388. Epub 2015/02/03. https://doi.org/10.3389/fonc.2014.00388 PMID:
25642418; PubMed Central PMCID: PMCPMC4295541.
27.
Pan R, Hogdal LJ, Benito JM, Bucci D, Han L, Borthakur G, et al. Selective BCL-2 inhibition by ABT-199
causes on-target cell death in acute myeloid leukemia. Cancer discovery. 2014; 4(3):362–75. Epub
https://doi.org/10.1158/1940-6207.crd-16-0305. Epub 2016/11/24. PMID:
27314085; PubMed Central PMCID: PMCPMC3975047.
26.
Vo TT, Ryan J, Carrasco R, Neuberg D, Rossi DJ, Stone RM, et al. Relative mitochondrial priming of
myeloblasts and normal HSCs determines chemotherapeutic success in AML. Cell. 2012; 151(2):344–
55. Epub 2012/10/16. https://doi.org/10.1016/j.cell.2012.08.038 PMID:
23063124; PubMed Central PMCID: PMCPMC3534747.
25.
Lito P, Rosen N, Solit DB. Tumor adaptation and resistance to RAF inhibitors. Nature medicine. 2013;
19(11):1401–9. Epub 2013/11/10. https://doi.org/10.1038/nm.3392 PMID: 24202393.
24.
Bruner JK, Ma HS, Li L, Qin ACR, Rudek MA, Jones RJ, et al. Adaptation to TKI Treatment Reactivates
ERK Signaling in Tyrosine Kinase-Driven Leukemias and Other Malignancies. Cancer research. 2017;
77(20):5554–63. Epub 2017/09/20. https://doi.org/10.1158/0008-5472.CAN-16-2593 PMID:
28923853.
23.
Kapoor S, Natarajan K, Baldwin PR, Doshi KA, Lapidus RG, Mathias TJ, et al. Concurrent inhibition of
Pim and FLT3 kinases enhances apoptosis of FLT3-ITD acute myeloid leukemia cells through
increased Mcl-1 proteasomal degradation. Clinical cancer research : an official journal of the American
Association for Cancer Research. 2017. Epub 2017/10/28. https://doi.org/10.1158/0008-5472.ccr-17-
1629 PMID: 29074603.
54. Nogami A, Oshikawa G, Okada K, Fukutake S, Umezawa Y, Nagao T, et al. FLT3-ITD confers resistance to the PI3K/Akt pathway inhibitors by protecting the mTOR/4EBP1/Mcl-1 pathway through STAT5 activation in acute myeloid leukemia. Oncotarget. 2015; 6(11):9189–205. Epub 2015/04/01. https://doi.org/10.18632/oncotarget.3279 PMID: 25826077; PubMed Central PMCID: PMCPMC4496211.

55. Nakajima W, Hicks MA, Tanaka N, Krystal GW, Harada H. Noxa determines localization and stability of MCL-1 and consequently ABT-737 sensitivity in small cell lung cancer. Cell death & disease. 2014; 5: e1052. Epub 2014/02/15. https://doi.org/10.1038/cddis.2014.6 PMID: 24525728; PubMed Central PMCID: PMCPMC3944277.