Waldenstrom macroglobulinemia cells devoid of $BTK^{C481S}$ or $CXCR4^{WHIM-like}$ mutations acquire resistance to ibrutinib through upregulation of Bcl-2 and AKT resulting in vulnerability towards venetoclax or MK2206 treatment

A Paulus$^{1,2}$, S Akhtar$^1$, H Yousaf$^1$, A Manna$^1$, SM Paulus$^2$, Y Bashir$^2$, TR Caulfield$^3$, M Kuranz-Blake$^1$, K Chitta$^1$, X Wang$^4$, Y Asmann$^4$, R Hudec$^5$, W Springer$^3$, S Ailawadhi$^7$ and A Chan-an-Khan$^{1,2}$

Although ibrutinib is highly effective in Waldenstrom macroglobulinemia (WM), no complete remissions in WM patients treated with ibrutinib have been reported to date. Moreover, ibrutinib-resistant disease is being steadily reported and is associated with dismal clinical outcome (overall survival of 2.9–3.1 months). To understand mechanisms of ibrutinib resistance in WM, we established ibrutinib-resistant in vitro models using validated WM cell lines. Characterization of these models revealed the absence of $BTK^{C481S}$ and $CXCR4^{WHIM-like}$ mutations. BTK-mediated signaling was found to be highly attenuated accompanied by a shift in PI3K/AKT and apoptosis regulation-associated genes/proteins. Cytotoxicity studies using the AKT inhibitor, MK2206 and ibrutinib, and the Bcl-2-specific inhibitor, venetoclax and ibrutinib, demonstrated synergistic loss of cell viability when either MK2206 or venetoclax were used in combination with ibrutinib. Our findings demonstrate that induction of ibrutinib resistance in WM cells can arise independent of $BTK^{C481S}$ and $CXCR4^{WHIM-like}$ mutations and sustained pressure from ibrutinib appears to activate compensatory AKT signaling as well as reshuffling of Bcl-2 family proteins for maintenance of cell survival. Combination treatment demonstrated greater (and synergistic) antitumor effect and provides rationale for development of therapeutic strategies encompassing venetoclax+ibrutinib or PI3K/AKT inhibitors+ibrutinib in ibrutinib-resistant WM.

**INTRODUCTION**

Waldenstrom macroglobulinemia (WM), a rare non-Hodgkin lymphoma variant, is characterized by unrestrained clonal proliferation of lymphoplasmacytic cells in the bone marrow and lymphoid tissue (lymph nodes, spleen). Patients usually present with cytopenias, lymphadenopathy and/or hepatosplenomegaly. In addition, WM cells produce and secrete excessive amounts of monoclonal immunoglobulin M (IgM), which can cause hyperviscosity syndrome and its associated complications. Therapeutic strategies have been extrapolated from other low-grade non-Hodgkin lymphoma and until very recently no drug had specifically secured approval in WM. 2

Ibrutinib, a first-in-class Brutons tyrosine kinase (BTK) inhibitor, is the first drug to gain Food and Drug Administration approval for treatment of WM and represents a milestone for patients suffering from this malignancy. In a phase II trial, relapsed or refractory WM patients who received ibrutinib demonstrated an overall response rate of 90.5%, with a major response rate of 70.5%. Estimated progression-free and overall survival (OS) at 24 months of treatment was 69.1% (95% confidence interval (CI): 53.2–80.5) and 95.2% (95% CI: 86.0–98.4), respectively. 3 However, no complete remissions were observed, indicating the WM cell’s ability to maintain their survival under ibrutinib-induced stress.

Despite the clinical benefit derived by patients treated with ibrutinib, undoubtedly the phenomenon of resistance to its effects is increasingly being reported in chronic lymphocytic leukemia (CLL), mantle cell lymphoma (MCL) and also WM (malignancies for which ibrutinib is currently approved). 4–11 Biologically this reflects the malignant tumor clone’s ability to survive sustained BTK inhibition and indicates the lack of curative potential at least with ibrutinib monotherapy. Indeed, ibrutinib-resistant disease is now consistently reported with fatal outcome, with median OS of CLL and MCL patients who relapse on ibrutinib being ~3.1 and 2.9 months, respectively. 12, 13 Although OS data for postibrutinib relapse WM patients is not yet available, it is anticipated that when these patients relapse (or become refractory to ibrutinib), their survival outcome may follow a similar dismal clinical course. Our laboratory efforts preemptively have tried to address this problem through development of unique models to interrogate the biology of ibrutinib resistance in WM in a quest to become prepared for potential salvage approaches. 14–16 Mechanistically, ibrutinib binds the Cys481 residue of the BTK kinase domain-active site and blocks autophosphorylation required for BTK activation. 17 In CLL and MCL patients, it has been reported that a cysteine-to-serine point mutation at residue 481 (C481S) in the allosteric inhibitory segment of BTK diminishes ibrutinib’s antitumor activity. 6, 8, 18 Similar observation has not
yet been confirmed in WM patients, and even in CLL and MCL, BTK\(^{*817}\) is not universally noted in all patients who develop ibrutinib resistance.\(^ {19,20}\) In WM, CXCR4\(^{WHIM+}\) mutations have been suggested as determinants of response to ibrutinib. However, the observation that 38% of WM patients who are MYD88L265P/CXCR4\(^{WHIM+}\) exhibit suboptimal response (i.e. less than major response) vs 62% of MYD88L265P/CXCR4\(^{WHIM+}\) patients who demonstrate major responses suggests that mechanisms other than CXCR4\(^{WHIM+}\) mutation must account for ibrutinib resistance.\(^ {17}\) Considering ibrutinib is the only approved therapeutic for WM, interrogation of the molecular mechanisms of resistance to ibrutinib in WM is of paramount importance to unveil new therapeutic opportunities in patients who have relapsed or become refractory to ibrutinib therapy.\(^ {21}\)

**MATERIALS AND METHODS**

**Cell lines, cell culture and reagents**

WM cell lines and their corresponding ibrutinib-resistant clones, developed in our laboratory, were used in experiments. All cell lines were cultured in RPMI-1640 containing 10% fetal bovine serum, penicillin (100 U/ml) and streptomycin (100 µg/ml). Cell viability was always maintained at >90% and was measured by trypan blue exclusion assay using ViCell-XR viability counter (Beckman-Coulter, Indianapolis, IN, USA). RPMI, penicillin, streptomycin, tetracycline, methyl ester (TMRM) and fetal bovine serum were purchased from Life Technologies (Carlsbad, CA, USA). Ibrutinib, MK2206 and ABT-199 (venetoclax) were purchased from Selleckchem (Houston, TX, USA). Annexin-V/Propidium Iodide Staining Kit was purchased from BD Biosciences (San Jose, CA, USA).

**Cell proliferation, viability and apoptosis assays**

MTS assay (Molecular Probes, Thermofisher Scientific, Rockford, IL, USA) or the CellTiter Glo Luminescent Cell Assay (Promega, Madison, WI, USA) were used to establish proliferation and viability of tumor cells, under previously reported conditions.\(^ {15,22}\) Apoptosis was measured by annexin-V and propidium iodide staining of cells followed by flow cytometry analysis. The conditions and method for these assays have been previously described by us.\(^ {15}\) Drug combination synergy, additivity or antagonism were determined using the CompuSyn Software (CompuSyn Inc., Paramus, NJ, USA), which uses the Chou-Talalay principle.\(^ {23}\) Details for this analysis are presented in the Supplementary Material and methods.

**Determination of MOMP**

Cells were treated with venetoclax for 12 h and assessed for mitochondrial outer membrane permeability (MOMP) using TMRM in a manner similar to that previously reported by us.\(^ {15,22,24,25}\)

**Cell respirometry assay**

Mitochondrial oxygen consumption rate was measured using the Seahorse XF96 Flux Analyzer (Seahorse Bioscience, Billerica, MA, USA). BCWM.1 and BCWM.1/IR cells were collected by centrifugation (room temperature (RT), 10 min, 200 g) washed twice with custom-made no phenol red containing low-buffered non-bicarbonated Seahorse medium (pH 7.35): powder Dulbecco’s modified Eagle’s media supplemented with 10 mm glucose, 10 mm pyruvate and 2 mm l-glutamine (all from Sigma-Aldrich, St Louis, MO, USA). Cell counts were adjusted and 2.5 x 10\(^{4}\) cells per well were seeded on Seahorse 96-well plate previously coated with poly-lysine—in pentaplicates per each condition. Pelleted cells (room temperature, 1 min, 600 g) were further capped in CO\(_2\)-free incubator for 1 h prior to the experiment. The Seahorse program included a 20 min incubation step followed by a 3 min mix, and 3 min measurement, which was repeated three times before and after each injection. Drugs were injected sequentially in final concentration as follows: 2 µM oligomycin, 1.5 µM carbonyl cyanide-4-phenylhydrazone and combination of rotenone and antimycin A each 0.5 µM (all from Sigma-Aldrich). After completion of the assay, the obtained oxygen consumption rate values were normalized to total protein content. Cells in each well were lysed in RIPA buffer: 50 µM Tris-Cl (pH 8.0), 150 mM NaCl, 0.1% sodium dodecyl sulfate, 0.5% deoxycholate and 1% NP-40 (all from Sigma-Aldrich) and protein content was determined using the BCA Protein Assay (ThermoFisher Scientific) according to the manufacturer’s recommendation. Normalized data were analyzed using Seahorse Wave software and Excel macro file XF Mito Stress Test Report Generator (Agilent Technologies, Santa Clara, CA, USA). Displayed data show mean values and standard error of the mean.

**Immunoblot analysis**

Western blot was performed for specific cellular protein estimations as reported previously.\(^ {21}\) Total protein extracts were prepared using RIPA lysis buffer alongside 0.2% protease and a phosphatase inhibitor cocktail (Sigma-Aldrich). Supernatant was collected for western blot analyses after centrifugation, and the protein content was measured by BCA protein assay reagent. Twenty micrograms of total protein were then boiled in Laemmli sample buffer, loaded onto 10% sodium dodecyl sulfate-polyacrylamide gel electrophoresis gels and transferred onto a nitrocellulose membrane. Blots were developed using chemiluminescence (ThermoFisher Scientific).

**Sanger sequencing of BTK and CXCR4**

RNA was isolated from all WM cells, including ibrutinib-resistant clones, using the miCURY RNA Isolation Kit (300110; Eixion, Woburn, MA, USA) and quantified using NanoDrop 3000 (Thermofisher Scientific). Using specific primer sequence for the whole gene, RNA was converted to cDNA by using SuperScript III One Step RT-PCR (12574030; Thermofisher Scientific). Primers for BTK and CXCR4 are included in the Supplementary Materials and methods.

**Cell surface antigen analysis**

For staining of cell surface markers, WM cells were washed two times with cold phosphate-buffered saline (PBS) and suspended in 300 µl of binding buffer (PBS solution with 2% fetal bovine serum). Cells were divided into three tubes: unstained, isotype control and those with antibody against CD19 (MACS Miltenyi Biotech, San Diego CA, USA), k-light-chain and \(\lambda\)-light-chain (Invitrogen, Life Technologies, Carlsbad, CA, USA). Five microliters of the respective antibody was added and cells were incubated for 30 min at room temperature. Tumor cells were washed two times with cold PBS and suspended in 100 µl of 4% paraformaldehyde in PBS solution (ThermoFisher Scientific), followed by analysis using an BD Accuri C6 Flow Cytometer (Agilent Technologies, Santa Clara, CA, USA). FCS Express 4 (De Novo Softwares, Glendale, CA, USA) was used to analyze the data.

**Cell cycle analysis**

Cells were serum starved for 4 h and treated with the drug for 12 h under normal culture conditions (10% fetal bovine serum). One million cells per sample were washed two times with cold PBS (SH30256; GE Healthcare Life Sciences, Logan, UT, USA). Cells were later resuspended in 10% fetal bovine serum. Cells were then resuspended in 1 ml of PBS containing 100 µg/ml and propidium iodide stain (50 µg/ml) solution (BD Biosciences) and incubated in water bath (37 °C) for 3 min. With a pipette, cells were resuspended and read on an Accuri C6 Flow Cytometer (BD Biosciences) at slow speed. Samples were analyzed using FCS Express 4 (De Novo Softwares).

**Human IgM measurement**

To quantify human IgM secreted in the sera of WM cells, we used the E-80M Human IgM ELISA Kit (Immunology Consultants Laboratory, Portland, OR, USA) according to the manufacturer’s instructions.

**Digital mRNA expression profiling (nCounter NanoString platform)**

Focused mRNA expression profiling was carried out using the NanoString nCounter Assay (NanoString, Seattle, WA, USA) to determine the expression of select cancer genes in WM cells treated with dimethyl sulfoxide (DMSO) or ibrutinib, as described previously.\(^ {22,25,26}\)

**Whole-exome sequencing**

DNA samples from BCWM.1, MWCL-1 and RPCI-WM1 and their respective ibrutinib-resistant derivatives (BCWM.1/IR, MWCL-1/IR, RPCI-WM1/IR) were captured using SureSelect Human All Exon V5+UTRs from Agilent Technologies. Additional details are described in the Supplementary Materials and methods.
RESULTS

Induction of ibrutinib resistance, independent of BTK<sup>C481S</sup> or CXCR4<sup>WHIM-like</sup> mutations.

To understand the molecular mechanisms that underlie induction of ibrutinib resistance, we exposed established human WM cell lines (n = 3) to sequentially escalating concentrations of ibrutinib over a period of 6 months. Compared with parent WM cells (wild-type, WT), ibrutinib-resistant derivatives (denoted by IR) maintained viability and proliferative potential when exposed to lethal concentrations of ibrutinib. Median half-maximal inhibitory concentration (IC<sub>50</sub>) of ibrutinib-resistant clones was 22 μM (range, 20–28 μM), whereas those of WT parent cells was 3.5 μM (range, 1–6.5 μM) (Figure 1a). Resistance was also examined by means of determining apoptotic potential of ibrutinib in ibrutinib-resistant and WT WM cells. While cells experienced ~30–50% apoptotic cell death at concentrations ranging between 5 and 10 μM, ibrutinib-resistant clones displayed significant resistance with almost no apoptosis occurring in these cells at concentrations as high as 30 μM (Figure 1b). As BTK<sup>C481S</sup> and CXCR4<sup>WHIM-like</sup> mutations have been reported to contribute towards ibrutinib resistance, we sequenced the corresponding DNA regions and found neither BTK<sup>C481S</sup> nor CXCR4<sup>WHIM-like</sup> mutation in any of the WT or ibrutinib-resistant WM cells by either whole-exome or Sanger sequencing (Figures 1c–f and Supplementary Figure S1). Altogether, these findings suggested that ibrutinib resistance in WM cells can emerge without the presence of BTK<sup>C481S</sup> or CXCR4<sup>WHIM-like</sup> mutations, indicating other potential mechanisms for WM cell survival in the presence of sustained BTK inhibition.

As BTK signaling is relevant to WM cell proliferation, we evaluated what effect continuous BTK suppression has on the cell cycle and proliferative potential of ibrutinib-resistant clones. Although ibrutinib-resistant WM cells continued to survive under chronic stress of ibrutinib (as these cells are kept in ibrutinib-containing media), their proliferation rate was much lower than parental WT cells. By 48 h, ibrutinib-resistant WM (BCWM.1/IR and RPCI-WM/IR) cell numbers were 1.46 and 1.73 million/ml, respectively, vs 2.73 and 3.69 million/ml in their respective WT counterparts (Figure 2a). Focused transcriptome analysis of ibrutinib-resistant cells revealed downregulation in the cell cycle genes, CDKN1A, CDKN2A in RPCI-WM1/IR vs RPCI-WM1 cells (Supplementary Figure S2 and Supplementary Table S1) and CDKN2A in BCWM.1/IR vs BCWM.1 cells (Supplementary Table S2). Cell cycle analysis by flow cytometry at 12 h showed that while ibrutinib treatment of WT WM cells retarded cell growth/division (as is consistent with the drugs known cytostatic effect), ibrutinib-
resistant cells (after 48 h washout) remained in a growth-arrested state where > 90% of cells were in the G1/G2 phase of the cell cycle (Figure 2b). Light microscopy showed that ibrutinib-resistant cells tended to cluster, and phenotypically were larger in size and less uniform in shape, confirming the inability of these cells to complete the G1 phase (Figure 2c).

Figure 2. Ibrutinib-resistant WM clones show altered growth kinetics, reduced IgM secretion and Ig-light-chain expression. (a) WT and ibrutinib-resistant WM clones (two representative isogenic pairs shown) were seeded at 0.5 million/ml, followed by measurement of total cell numbers and viability at 24 and 48 h. (b) WM cells and ibrutinib-resistant derivatives (two representative isogenic pairs shown) were treated with DMSO or increasing concentrations of ibrutinib (1 or 5 μM for 24 h), followed by flow cytometry and cell cycle analysis. Ibrutinib treatment resulted in accumulation of WM tumor cells in the G1 phase of the cell cycle. By comparison, ibrutinib-resistant WM clones (BCWM.1/IR and RPCI-WM1/IR) were noted to be G1 arrested. (c) Light microscopy showed ibrutinib-resistant WM cells to be more irregular in shape and larger in size as compared with their WT counterparts (one isogenic pair shown as representative). (d) Mitochondrial respiration was measured side by side in parental cells (BCWM.1) and ibrutinib-resistant WM cells (BCWM.1/IR) at baseline condition, followed by inhibition of mitochondrial ATP production (introduction of oligomycin), uncoupling oxidative phosphorylation (OP) and ATP production causing maximal mitochondrial respiration (introduction of carbonyl cyanide-4-phenylhydrazone (FCCP)) and finally complete inhibition of the OP by the combination of rotenone and antimycin A. oxygen consumption rate (OCR) is displayed in values normalized to total protein content. (e) Compared with parental cells, BCWM.1/IR cells showed increased values in all analyzed parameters: basal respiration (left), ATP production (middle left) and maximal respiration (middle right) and non-mitochondrial respiration (right). (f) IgM secretion was quantified in the supernatant of WT and ibrutinib-resistant WM clones (± ibrutinib 1 μM, 24 h exposure). In WT WM cells, secretion of IgM decreased after exposure to ibrutinib. In ibrutinib-resistant clones, IgM secretion was measured in cells after a washout period (cells cultured in ibrutinib-free media for 1 week) as well as after ibrutinib treatment and was noted to be markedly reduced in washout cells compared to WT counterparts, regardless of exposure to ibrutinib thereafter. (g) In a similar manner, all WM cell lines were incubated with anti-λ-light-chain or anti-k-light-chain antibody and analyzed by flow cytometry, which demonstrated decreased surface expression of native Ig-light-chain on ibrutinib-resistant WM cells.
CD19- and this was also noted in RPCI-WM1/IR cells. Compared to WT parental cells, BCWM.1/IR and MWCL-1/IR cells showed reduced surface expression of CD19. RPCI-WM1 is known to be clones- before or after washout, was not impacted. (4) Immunobot analysis showed very low pBTK, pSYK or pPLC\(\gamma\) protein levels in these cells. Moreover, carbonyl cyanide-4-phenyl-

The G1 phase of the cell cycle has been shown to be associated with changes in the mitochondria and increased respiration. 28 This led us to examine mitochondrial welfare and bioenergetics, comparing mitochondrial respiration in parental and resistant cells. Using the Seahorse XF96 Flux Analyzer and examining BCWM.1 and its corresponding ibrutinib-resistant subclone (BCWM.1/IR), we found the resistant cells to have an overall increased oxygen consumption rate compared with parental cells (Figure 2d). Basal respiration in the resistant cells was increased by 30%. A similar level of increase was found in mitochondrial ATP production supporting the observation of increased mitochondrial utilization in these cells. Moreover, carbonyl cyanide-4-phenyl-hydrazone treatment revealed elevation in maximal respiration that reflects higher spare respiratory capacity of mitochondria in BCWM.1/IR cells as compared with parental BCWM.1 cell (Figure 2e). These findings suggest higher capability of ibrutinib-resistant cells to maintain and sustain increased mitochondrial demand under stress conditions. Moreover, we noted more than 50% increase in non-mitochondrial oxygen consumption/respiration, suggesting the ability of mitochondria of ibrutinib-resistant WM cells to adapt in a pro-oxidative environment, thus making them more resilient toward stress-associated cytotoxicity.

We also examined IgM secretion from the ibrutinib-resistant cells and found it to be significantly decreased in all resistant clones (after 48 h washout and reintroduction to ibrutinib) compared with WT cells (before and after treatment with ibrutinib) (Figure 2f). Consistent with this effect, we also observed reduced cell surface expression of cognate Ig-light-chain receptors on ibrutinib-resistant WM cells (Figure 2g). Altogether, these data indicate the altered proliferative capacity yet tenacity of ibrutinib-resistant cells to maintain survival under selective pressure from ibrutinib, through mechanisms independent of \(BTK^{C481S}\) and \(CXCR4\) mutations.

BCR signaling is attenuated in ibrutinib-resistant WM cells without consequence to tumor cell viability and growth

As ibrutinib abrogates BTK-mediated signaling that is propagated through the B-cell receptor (BCR) hub, 5,29 we examined this signaling pathway in all WM cell lines. Ibrutinib treatment (1 \(\mu\)M) of WT WM cell lines markedly decreased phospho (p)BTK, pPLC\(\gamma\)2 and pSYK levels, demonstrating drug–target engagement and inhibition of BCR signaling as would be anticipated (Figure 3a).

We also examined IgM secretion from the ibrutinib-resistant cells and found it to be significantly decreased in all resistant clones (after 48 h washout and reintroduction to ibrutinib) compared with WT cells (before and after treatment with ibrutinib) (Figure 2f). Consistent with this effect, we also observed reduced cell surface expression of cognate Ig-light-chain receptors on ibrutinib-resistant WM cells (Figure 2g). Altogether, these data indicate the altered proliferative capacity yet tenacity of ibrutinib-resistant cells to maintain survival under selective pressure from ibrutinib, through mechanisms independent of \(BTK^{C481S}\) and \(CXCR4\) mutations.

Contrastingly, in ibrutinib-resistant derivatives, we noted constitutively decreased pBTK, pPLC\(\gamma\)2 and pSYK protein levels—with the cells retaining >90% viability. Interestingly, after 48 h washout and culture in ibrutinib-free media, we observed marked upregulation of pBTK, pPLC\(\gamma\)2 and pSYK, but this did not impact ibrutinib-resistant cell viability (Figure 3b). As CD19 can complex with the BCR receptor to enhance signaling, 30 we examined CD19 surface receptor expression by flow cytometry and observed significant decrease in ibrutinib-resistant cells compared with WT cells (Figure 3c). Taken together, these data suggested to us that ibrutinib-resistant WM cells shift survival and growth dependence away from BCR/BTK-mediated signaling and engage alternative pathway(s) to maintain viability.

Phospho-AKT expression in ibrutinib-resistant cells is upregulated and its pharmacologic inhibition results in decreased tumor cell survival and apoptosis

Dysfunctional AKT signaling has been reported to contribute to ibrutinib insensitivity in \(CXCR4^{MM}\)-positive WM cells. 9 As such, we investigated the relevance of AKT in our \(CXCR4^{MM}\)-negative ibrutinib-resistant WM cells. Targeted transcriptome analysis showed expression of genes associated with AKT signaling (such as MET and WNT10B) to be increased in RPCI-WM1/IR cells (Supplementary Table S1). Indeed, we had noted that after treatment of WT BCWM.1 cells with ibrutinib, phosphorylation of AKT (at Thr308 and Ser473) was markedly increased (Figure 4a, left panel set). And, this effect was prominent at baseline in BCWM.1/IR cells with a slight increase in pAKT (Ser473) and its pharmacologic inhibition results in decreased tumor cell survival and apoptosis.
WM1/IR cells showed constitutive expression of pAKT (more prominent at the Ser473 residue), which did not change following washout and incubation with ibrutinib (Figure 4a, right panel set). To examine functional relevance of increased pAKT in CXCR4 WHIM-negative ibrutinib-resistant WM cells, we treated these cells (BCWM.1 and BCWM.1/IR; RPCI-WM1 and RPCI-WM1/IR) with the clinical-grade allosteric pan-AKT inhibitor, MK2206, and observed marked reduction of pAKT as well as evidence of apoptosis as indicated by PARP-1 cleavage (Figure 4a). However, when cells were treated concurrently with ibrutinib and MK2206, we noted marked reduction in pBTK and nearly complete loss of pAKT, which was associated with more robust cleavage of PARP-1 (Figure 4a). Overall, it is important to note that the minor differences noted in protein levels of different signaling molecules between the various cell line models is expected given our prior report/analysis on the variable state of generation of these cell lines (i.e. relapse/refractory vs untreated). Observing that concurrent BTK and AKT inhibition could result in potentially more programmed cell death, we treated BCWM.1 and RPCI-WM1 (and their respective ibrutinib-resistant derivatives) with different concentrations of MK2206, ibrutinib or MK2206+ibrutinib for 48 h and assessed the viability of the CellTiter Glo Luminescent Cell Assay to determine pharmacologic synergy. A combination index (CI) value of 1 signifies additivity, whereas a CI < 1 signifies synergism and a CI > 1 signifies antagonism between the drugs at particular concentrations. Bar graph shows representative results of combination experiments where drug combination synergy was noted. In BCWM.1 cells, ibrutinib (2 μM)+MK2206 (2 μM) reduced cell viability to 19% and in BCWM.1/IR cells, a similar trend was noted at similar concentrations, where the combination of both agents caused significant reduction in cell viability (40% viable cells) compared with either agent alone. (c) Representative isobologram analysis of BCWM.1/IR cells shows synergistic cytotoxic activity of ibrutinib+MK2206 in 6/6 drug combinations tested (represented by blue circles). (d) In RPCI-WM1 and RPCI-WM1/IR cells, ibrutinib or MK2206 alone did not significantly reduce cell viability; however, the combination of the two agents (ibrutinib 4 μM+MK2206 4 μM in RPCI-WM1 cells; ibrutinib 16 μM+MK2206 8 μM in RPCI-WM1/IR cells) significantly reduced cell viability to 16 and 11% in RPCI-WM1 and RPCI-WM1/IR cells, respectively. (e) Representative isobologram analysis of RPCI-WM1/IR cells showed synergistic cytotoxic activity in 2/6 combinations (represented by blue circles).

Modulation of Bcl-2 expression as a function of resistance induction resulting in sensitivity to venetoclax treatment

In addition to compensatory AKT signaling, we investigated other potential mechanisms for ibrutinib resistance in WM cells. Deferred apoptosis is a hallmark of many B-cell malignancies associated with drug resistance; as such, we examined expression of apoptotic regulators in our transcriptome expression data set. This analysis revealed modulation of several genes known to regulate apoptotic machinery such as BMI1, GNAS, IL8

Figure 4. Phospho-AKT is increased in ibrutinib-resistant WM cells and its pharmacologic inhibition results in decreased tumor cell survival and apoptosis. (a) WM cell lines were treated with DMSO, ibrutinib or MK2206 (allosteric pan-AKT inhibitor) at indicated concentrations for 24 h and immunoblot analysis was performed probing for BTK, pBTK, AKT (Thr308), pAKT (Ser473) and PARP-1 protein expression. pAKT was noted to be decreased in all WM cells (alone or in combination with ibrutinib). PARP-1 cleavage in WM cells was most markedly observed after combination ibrutinib and MK2206 treatment. (b) Two isogenic cell line pairs (BCWM.1 and BCWM.1/IR; RPCI-WM1 and RPCI-WM1/IR) were treated with DMSO or various concentrations of MK2206, ibrutinib or ibrutinib+MK2206 for 48 h and assessed for viability by the CellTiter Glo Luminescent Cell Assay to determine pharmacologic synergy. A combination index (CI) value of 1 signifies additivity, whereas a CI < 1 signifies synergism and a CI > 1 signifies antagonism between the drugs at particular concentrations. Bar graph shows representative results of combination experiments where drug combination synergy was noted. In BCWM.1 cells, ibrutinib (2 μM)+MK2206 (2 μM) reduced cell viability to 19% and in BCWM.1/IR cells, a similar trend was noted at similar concentrations, where the combination of both agents caused significant reduction in cell viability (40% viable cells) compared with either agent alone. (c) Representative isobologram analysis of BCWM.1/IR cells shows synergistic cytotoxic activity of ibrutinib+MK2206 in 6/6 drug combinations tested (represented by blue circles). (d) In RPCI-WM1 and RPCI-WM1/IR cells, ibrutinib or MK2206 alone did not significantly reduce cell viability; however, the combination of the two agents (ibrutinib 4 μM+MK2206 4 μM in RPCI-WM1 cells; ibrutinib 16 μM+MK2206 8 μM in RPCI-WM1/IR cells) significantly reduced cell viability to 16 and 11% in RPCI-WM1 and RPCI-WM1/IR cells, respectively. (e) Representative isobologram analysis of RPCI-WM1/IR cells showed synergistic cytotoxic activity in 2/6 combinations (represented by blue circles).
and PIM1 in BCWM.1/IR and RPCI-WM1/IR cells (Supplementary Tables S1 and 2). Thus, we immunoblotted directly for apoptosis effectors; anti-Bcl-2, Mcl-1, A1) and proapoptotic (Bax, Bak, Bim and PUMA) Bcl-2 family members in WT and ibrutinib-resistant clones and observed increased Bcl-2 and Mcl-1 in the latter, whereas proapoptotic proteins displayed more variable expression patterns (Figure 5a). This demonstrated that dysfunctional Bcl-2 interactions may have a role in maintaining ibrutinib-resistant cell survival. Next, we investigated anti-WM activity of the Bcl-2-BTK inhibitor venetoclax (ABT-199, recently Food and Drug Administration-approved for treatment of 17p deletion-positive relapsed/refractory CLL),37–39 in WT and ibrutinib-resistant derivatives. Modeling studies of venetoclax revealed its occupancy at the P2 and P4 pockets in the BH3 groove of the Bcl-2 protein. (c) A 48 h MTS assay was conducted to assess sensitivity of all WM cell lines towards venetoclax, which demonstrated comparable activity in WT and ibrutinib-resistant cells alike, with the exception of MWCL-1/IR cells, which showed greater growth inhibition compared with MWCL-1 parental cells, with the exception of MWCL-1/IR cells, which showed greater growth inhibition compared with MWCL-1 parental cells. (d) WM cell lines were treated with DMSO or venetoclax (5 μM) for 24 h and stained with annexin-V and propidium iodide, followed by flow cytometry to examine apoptosis. Venetoclax induced significant apoptosis in all WM cell lines tested; heat density plots from a representative isogenic pair model, BCWM.1 and BCWM.1/IR, show 60% and 63% annexin-V positivity, respectively, after treatment with the drug. (f) Mitochondrial-mediated apoptosis was validated in WM cells treated with DMSO or venetoclax by immunoblotting for PARP-1, caspase-9 and caspase-3 protein products, which demonstrated cleavage of these proteins in all cells treated with the drug.

Venetoclax synergistically enhances the antiproliferative effects of ibrutinib

Drug combination therapy produces enhanced antitumor effect and counters drug resistance.41 We addressed the question of whether concurrently inhibiting Bcl-2 (with venetoclax) and BTK (with ibrutinib) could induce cytotoxicity in WM cells and overcome ibrutinib-resistant WM cell insensitivity to single agent ibrutinib. Two isogenic WM cell line pairs were treated with sub-IC50 concentrations of venetoclax (V), ibrutinib (I) or venetoclax +ibrutinib (V+I) for 48 h and viability was assessed thereafter. In BCWM.1 and BCWM.1/IR cells, the combination of V+I synergistically reduced tumor cell viability; significantly more so than either V or I alone at a combination index of 0.03 and 0.21, respectively (Figure 6a). Isobologram analysis of BCWM.1/IR cells formation and induction of apoptosis.40 Thus, we examined MOMP in WT and ibrutinib-resistant WM cells before and after venetoclax treatment. As anticipated, venetoclax compromised mitochondrial function in both WT and ibrutinib-resistant WM cells by increasing MOMP (Figure 5d), which was associated with induction of apoptosis as determined by annexin-V/PI staining (Figure 5e). Furthermore, (mitochondrial-mediated) programmed cell death in venetoclax-treated cells was confirmed by the presence of PARP-1 cleavage and cleaved caspase-3 (Figure 5f).
demonstrated synergistic cytotoxic activity of V+I in 6/6 combinations (represented by blue circles) (Figure 6b). In the RPCI-WM1 cell line and its ibrutinib-resistant derivative, the combination of V (1.5 μM)+I (15 μM) synergistically reduced tumor cell viability (CI of 0.48 and 0.80, respectively), an effect more notable than when cells were treated with V or I alone. (d) Representative isobologram analysis of RPCI-WM1/IR cells showed synergistic cytotoxic activity in 2/6 combinations (represented by blue circles).

DISCUSSION
Ibrutinib-resistant Waldenstrom macroglobulinemia A Paulus et al

Ibrutinib resistance represents a unique challenge due to its association with a poor clinical outcome. Overall, biological understanding of how ibrutinib resistance occurs in WM remains limited. Mechanisms of resistance in other B-cell cancers are reported to be associated with ibrutinib-binding site mutations in BTK, whereas in WM, CXCR4WHIM-like mutations have been reported to correlate with ibrutinib insensitivity in a limited number of cases. As such, the optimal therapeutic strategies to extend clinical remission in patients who have become refractory to ibrutinib are unclear and remain an active area of investigation. To this end, we dedicated our efforts towards establishment of preclinical WM models to study the phenomenon of ibrutinib resistance as is observed clinically—where chronic exposure to ibrutinib over an extended time leads to outgrowth of tumor clones that can maintain proliferative potential under ibrutinib-induced stress.

Characterization of ibrutinib-resistant clones revealed the absence of BTK or CXCR4WHIM-like mutations. A C481S mutation has been shown to disrupt covalent (irreversible) binding of ibrutinib to BTK and instead permits for a reversible binding interaction, limiting the drugs effect by nearly 25-fold. The absence of a C481S anomaly in our resistant models implied that...
Ibrutinib should optimally bind and functionally inhibit BTK, irrespective of impact on cell viability. The latter is important to note, as our assessment of phosphorylation status of BCR signalosome components in resistant cells showed that tyrosine phosphorylation of not only BTK (at Y551 and Y223 amino acids) but effectors upstream such as SYK (Y322 and Y525/526) and downstream, PLCγ2 (Y759 and Y1275), were inhibited by ibrutinib, indicating that drug binding is intact and BCR signaling is abrogated. And, this effect (BCR signalosome phosphorylation status) was reversed when resistant cells were washed out and grown in ibrutinib-free media. In either condition, ibrutinib-resistant cells remained viable with stable growth kinetics, indicating to us that in these cells BTK-mediated signaling was not a primary driver of disease. We also noted decreased lgh secretion in the sera of ibrutinib-resistant cells. It would appear that in this drug-resistant state, the emphasis of the tumor cells is more towards survivability and transitioning to an ‘oligosecretory phase’. Clinically this is noted also in patients with plasma cell malignancies, whereby with increasing resistance to therapeutics the cell can eventually evolve into a non-secretory/oligosecretory phase.

In addition to BTK, we also probed for CXCR4 (WHIM-like) mutations, which in WM have been shown to decrease CXCR4 receptor internalization, thereby allowing continuous signaling and downstream activation of ERK and AKT, leading to sustained WM cell proliferation/survival. However, the role of AKT and its relevance in WT CXCR4 ibrutinib-resistant WM is unclear. While pAKT was effectively downregulated with MK2206 treatment alone in all WM cells tested herein, this did not translate to a significant loss of viability in WT or ibrutinib-resistant RPCI-WM1/IR cells—and could potentially just be a function of increased basal AKT activity in these cells. However, combined BTK and AKT inhibition induced a significant decrease in RPCI-WM1 and RPCI-WM1/IR cell viability, suggesting that the protective role of AKT could be overcome with concurrent targeting of BTK with ibrutinib (at sub-IC50 concentrations). Further studies are needed to explicate the role of AKT in CXCR4 (WHIM-like) negative ibrutinib-resistant WM and novel, clinically relevant inhibitors that target upstream of AKT, that is, PI3K isoforms (idelalisib, duvelesib) may hold greater anti-WM potential when combined with ibrutinib.

In conjunction with growth-stimulating signaling pathways such as those propagated by AKT/ERK/NFkB activation, WM cell survival is intricately supported by malfunctions in initiation of apoptosis through dysregulated expression of Bcl-2 family proteins. Interestingly, AKT and Bcl-2 are directly linked, where studies by Pugazhenthi et al. have shown that AKT activates the transcription factor CREB (CAMP response element-binding protein), which binds to the Bcl-2 promoter region and positively regulates its expression. Whether upregulation of Bcl-2 in ibrutinib-resistant WM is related to CREB and AKT is currently being examined by us. Nevertheless, we have previously shown the utility of disrupting proapoptotic Bcl-2 protein interactions in CLL, WM and multiple myeloma and that this functional interference results in malignant cell death or resensitization (or heightened sensitivity) of tumor cells towards other therapeutics. Indeed, this observation also held true for CXCR4 (WHIM-like) negative ibrutinib-resistant WM cells, where the Bcl-2 specific BH3 mimetic, venetoclax, decreased ibrutinib-resistant WM cell viability and induced apoptosis as a single agent. Our investigations further highlighted that concomitant disruption of BTK and Bcl-2 with sub-IC50 doses of ibrutinib and venetoclax, respectively, synergistically induce tumor cell lethality in WT and ibrutinib-resistant cells. These data are in line with observations by Cao et al. in WM, whereas we are the first to demonstrate the utility of combined Bcl-2 and BTK inhibition in ibrutinib-resistant WM cells in which neither BTK (Kh73) nor CXCR4 (WHIM-like) mutations are responsible for drug resistance. The rationale for Bcl-2 plus BTK inhibition is further supported by preclinical studies in CLL and MCL, the latter where the combination of venetoclax and ibrutinib is being investigated in two separate phase clinical trials (NCT02419560, phase I study; NCT02471091, phase II study).

In conclusion, we report on the development of novel ibrutinib-resistant WM cell lines, which exhibit decreased survival dependency on BTK-mediated signaling. These models all lack ibrutinib-binding site (BTK Cys481) mutations as well as CXCR4 (WHIM-like) mutations, providing new insight into mechanism underpinning ibrutinib resistance independent of these aberrations. Focused transcriptome analysis combined with protein profiling revealed AKT and Bcl-2-associated pathways may have a role in maintaining survival of ibrutinib-resistant WM cells. In vitro efficacy and mechanistic analysis confirmed reliance on both pathways and suggests that drug combination strategies encompassing BTK/AKT/PI3K inhibition or BTK+Bcl-2 inhibition can potentially overcome ibrutinib resistance in WM. Finally, our observations are unique as they feature previously unknown mechanisms directing resistance to ibrutinib in WM cells and provide rationale for potential combination therapeutic strategies to address this clinical problem in patients with WM.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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AUTHOR CONTRIBUTIONS
AP designed and performed the research, analyzed and interpreted the data, contributed to writing of the manuscript and approved the final draft. SA performed the research, collected and analyzed the data, contributed to writing of the manuscript and approved the final draft. HY performed the research, analyzed the data and approved the final draft. AM performed the research and provided critical revision for important intellectual content and approved the final draft. SP performed the research, performed critical revision for important intellectual content and approved the final draft. YB performed the research, performed critical revision for important intellectual content and approved the final draft. TRC performed the research, collected and analyzed the data, contributed to writing of the manuscript and approved the final draft. KC designed and performed the research, collected and analyzed the data, contributed to writing of the manuscript and approved the final draft. KW performed the research, collected and analyzed the data, contributed to writing of the manuscript and approved the final draft. RH performed the research, collected and analyzed the data, contributed to writing of the manuscript and approved the final draft. WA analyzed the data, performed critical revision for important intellectual content contributed to writing of the manuscript and approved the final draft. HY performed the research, collected and analyzed the data, contributed to writing of the manuscript and approved the final draft. WS performed the research, performed critical revision for important intellectual content and approved the final draft. SA analyzed the data, performed critical revision for important intellectual content and approved the final draft. AC-K designed the research, analyzed the data provided critical scientific and intellectual oversight, wrote the manuscript and approved the final draft.
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