BH3 mimetics targeting BCL-XL impact the senescent compartment of pilocytic astrocytoma

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Running title
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Funding
Brain Tumor Foundation and the PLGA Program at Dana-Farber Cancer Institute. T.M. was supported by the Bilger family.

**Conflict of interest**

T.M. received research funding from Biomed Valley and Day One Therapeutics. C.M.T. participated in advisory boards of Novartis and Bayer. S.M.P. receives funding from Bayer, Pfizer, Eli-Lilly, Roche, Amgen, AstraZeneca, PharmaMar, Sanofi and Servier in the context of an IMI-2-funded EU project entitled ITCC-P4 (www.itccp4.eu). O.W. participated in advisory boards of Novartis, BMS, Janssen and receives research grants from BVD, Day One Therapeutics. J.D. and Y.Z. are employees of Ascentage Pharma. J.M. is co-inventor of dynamic BH3 profiling, was paid consultant for Oncoheroes Biosciences and Vivid Biosciences, is unpaid board member for The Society for Functional Precision Medicine, is collaborating with AstraZeneca.

**Authorship**


**Total word count**

6982 words
Abstract

Background: Pilocytic astrocytoma (PA) is the most common pediatric brain tumor and a mitogen-activated protein kinase (MAPK)-driven disease. Oncogenic MAPK-signaling drives the majority of cells into oncogene-induced senescence (OIS). While OIS induces resistance to anti-proliferative therapies, it represents a potential vulnerability exploitable by senolytic agents.

Methods: We established new patient-derived PA cell lines that preserve molecular features of the primary tumors and can be studied in OIS and proliferation depending on expression or repression of the SV40 large T antigen. We determined expression of anti-apoptotic BCL-2 members in these models and primary PA. Dependence of senescent PA cells on anti-apoptotic BCL-2 members was investigated using a comprehensive set of BH3-mimetics.

Results: Senescent PA cells upregulate BCL-XL upon senescence induction and show dependency on BCL-XL for survival. BH3 mimetics with high affinity for BCL-XL (BCL-XLi) reduce metabolic activity and induce mitochondrial apoptosis in senescent PA cells at nanomolar concentrations. In contrast, BH3 mimetics without BCL-XLi activity, conventional chemotherapy and MEK inhibitors show no effect.

Conclusions: Our data demonstrates that BCL-XL is critical for survival of senescent PA tumor cells and provides proof-of-principle for the use of clinically available BCL-XL-dependent senolytics.

Key words  pilocytic astrocytoma, BH3 mimetics, BCL-XL, oncogene-induced senescence

Key Points

- New patient-derived PA models for studies in oncogene-induced senescence and proliferation
• BH3 mimetics targeting BCL-XL induce apoptosis in senescent PA cells
• BCL-XL is the first reported target for senolytic treatment of PA

Importance of the Study

PAs are the most common pediatric brain tumors. Incompletely resected tumors frequently progress after cessation of anti-proliferative treatments. Relapses and multiple lines of salvage-therapy cause substantial long-term morbidity and toxicity. Because most PA cells are in OIS, they are not sensitive to anti-proliferative treatments and can therefore serve as source for relapse and progression. OIS may thus be an unexploited vulnerability of PA cells targetable by senolytic drugs. We here report the first evidence supporting the use of BCL-XLi to target the so far un-targeted senescent compartment of PA. Senolysis induced by clinically available BCL-XLi could open a new avenue to improve long-term outcomes of PA patients. Based on the data presented, conceptualization of a clinical trial investigating BCL-XLi in relapsed and progressive PAs is currently being discussed.
Introduction

Pilocytic astrocytomas (PA) are the most frequent pediatric brain tumors and the largest subgroup of pediatric low-grade gliomas (pLGG). In contrast to the excellent 10-year overall survival of over 90%, the event-free survival of pLGG patients is only ca. 50%. More than half of the incompletely resected patients treated with conventional chemotherapy progress and require one or more lines of salvage therapy. PAs are characterized by activation of the mitogen-activated protein kinase (MAPK) pathway. Although the mitogen-activated protein kinase kinase (MEK) inhibitors (MEKi) selumetinib and trametinib were shown to be well tolerated and effective in progressive or relapsing pLGGs in phase I/II clinical trials and retrospective case studies, tumors may relapse or progress shortly after discontinuation of treatment. Therefore, development of complementary new treatment strategies is needed to improve long-term outcome of PA patients.

MAPK pathway activation leads to oncogene-induced senescence (OIS) in PA. Less than 5% of primary PA cells express the proliferation marker Ki67, indicating that the majority of cells are not cycling. While cells in OIS are not responsive to anti-proliferative treatments and may constitute a reservoir of tumor cells that can lead to tumor relapse upon cessation of anti-proliferative therapies, OIS might represent a yet unexploited complementary vulnerability of PA targetable by senolytic drugs.

In 2015 Zhu et al identified the activation of pro-survival networks in senescent cells and silenced expression of the key nodes of these networks to target senescent cells. Subsequently, pharmacological clearance of senescent cells by compounds of mechanistically diverse classes including BH3 mimetics was described. BH3 mimetics are inhibitors of the anti-apoptotic B-cell lymphoma 2 (BCL-2) family members, which comprise BCL-2, BCL-XL, Bcl-2-like protein 2 (BCL-W), induced myeloid leukemia cell differentiation protein (MCL-1) and BCL2 related protein A1 (BFL-1). Within the tightly balanced process of intrinsic apoptosis, the main function of these anti-apoptotic BCL-2 members is to prevent the induction of mitochondrial membrane potential breakdown and apoptosis.
Previous data from our group indicated the senolytic activity of the BH3 mimetic navitoclax, an inhibitor of BCL-2, BCL-XL and BCL-W, in PA.\textsuperscript{20} The lack of more PA models, that would allow for validation and comprehensive testing of senolytic compounds in OIS, is a major obstacle. While short term cultures can only be incompletely characterized and do not yield enough material for comprehensive studies, the few existing long-term pLGG-derived \textit{in vitro} models (BT40\textsuperscript{21}, JHH-NF1-PA1\textsuperscript{22}, Res186, Res259\textsuperscript{23}) are not suited for testing in senescence because they continuously proliferate and do not recapitulate the PA senescence biology.

The aim of the present study was to evaluate the senolytic properties of BH3 mimetics in PA and to decipher their translational potential. We made use of our PA cell line DKFZ-BT66\textsuperscript{24} and a set of three completely newly established PA cell lines suitable for testing in proliferation and OIS depending on expression or repression of SV40 large T antigen. The aim was to identify the BCL-2 members essential for survival of senescent PA cells, which need to be targeted by BH3 mimetics to induce senolysis.
Materials and methods

Processing of primary patient samples

Primary PA tumor material (DKFZ-BT308, DKFZ-BT314 and DKFZ-BT317) were collected during therapeutic intervention. Informed consent for sample collection and use was obtained within the study S-304/2014, which was approved by the institutional review board of the University of Heidelberg. Clinical, histopathological and molecular data of the samples are summarized in Supplementary Table 1. For processing and culture of primary samples see Supplementary Methods.

Inducible expression of simian vacuolating virus large T antigen (SV40-TAg)

For generation of pCW57.1 GFP-TAg, allowing for doxycycline-inducible co-expression of destabilized GFP (dsGFP) and SV40 TAg, see Supplementary Methods. Primary PA cell cultures were transduced after 2 to 7 days in culture with 3 ml pCW57.1 GFP-TAg supernatant in 6-well plates (Corning). ABM medium and 1µg/ml doxycycline were added 6 hours after infection. Work with lentiviruses was approved by the authorities (internal project number 80935) and performed in a biosafety level two laboratory.

Cell culture and drugs

DKFZ-BT66, DKFZ-BT308, DKFZ-BT314 and DKFZ-BT317 cell lines were grown as described before 24. For details see Supplementary Methods.

Metabolic activity, DSS calculation and drug combination profiling

For measurement of metabolic activity see Supplementary Methods. Drug sensitivity scores were calculated based on dose-response data using the R package DSS (v 1.2)25. DSS3 mode was used, with output ranging from 0 (insensitive) to 100 (highly sensitive).
Cell viability assessment and cell counting

Five days prior to treatment 0.8 x 10^6 cells/well were seeded in 6-well plates (Corning) without doxycycline. Medium changes were performed every second to third day. Cells were then incubated with different BH3 mimetics at concentrations indicated or DMSO. After 72 hours the remaining attached cells were collected and cell number as well as viability was assessed using trypan-blue based automated cell counting (Vi-CELL XR automated cell counter; Beckmann Coulter).

Senescence-associated β-galactosidase staining

Senescence-associated (SA) β-galactosidase staining was performed as described. 1 x 10^5 cells were seeded per well of a 6-well plate (Corning) 5 days prior to fixation without doxycycline. Images were taken with a Nikon Eclipse Ts2 microscope and Nikon Elements software (v 5.20).

Gene expression analysis

Gene expression analysis of PA cell lines was performed using Human Genome U133 Plus 2.0 chips (Affymetrix GeneChips®) with n=3 independent biological replicates per condition. Affymetrix U133 Plus 2.0 expression array data of primary PA and normal cerebella was from R2 (Tumor Pilocytic astrocytoma (DKFZ) - Kool; R2 internal identifier: ps_mkheidel_mkd kfz209_u133p2). Array data was MAS5.0 normalized. Gene expression data of cell lines were downloaded from GDSC database (GDSC2). Gene set enrichment analysis (GSEA) was done using GSEA software (v 4.1.0). Single sample GSEA (ssGSEA) was performed using the public server of GenePattern (www.genepattern.org). BCL2 and BCL2L1 expression was analyzed in previously published single cell RNA sequencing data of primary PA.
Western blot and immunoprecipitation

For information about antibodies used see Supplementary Methods. Western blot analysis was performed as described before. Immunoprecipitation (IP) was done using Dynabeads™ Protein G Immunoprecipitation Kit (Invitrogen, 10007D) following the manufacturer’s instructions. Antibodies were crosslinked to Dynabeads™ using freshly dissolved 20 mM dimethyl-pimelimidate (DMP) in 0.2 M triethanolamine buffer.

Tissue micro array and immunohistochemical staining

A tissue micro array of 83 PAs (CNS WHO grade I), part of the LOGGIC Core study, was generated and stained, for details see Supplementary Methods. Inconspicuous CNS tissue adjacent to low-grade gliomas was used as normal controls (n=16).

shRNA mediated gene silencing

The lentiviral vector pGIPZ (Horizon Discovery Biosciences Limited) was used for shRNA knockdowns of BCL-XL. A mix of four shRNAs with the following clone IDs (all from Horizon Discovery Biosciences Limited) and mature antisense sequences were used: V3LHS_641297 (GCATCTCCTTGTCTACGCT); V3LHS_413478 (AAATTCTAGAAAACTAGCT); V2LHS_269487 (TTTCCGACTGAAGAGTGAG); V3LHS_393128 (CACTAAACTGACTCCAGCT). For details on lentivirus production and infection see Supplementary Methods. Protein samples for determination of knockdown efficiency were harvested 96 h after transduction. The amount of remaining attached cells was counted 14 days after transduction using Vi-CELL XR automated cell counter (Beckmann Coulter).

Real time quantitative reverse transcription real-time PCR (RTqPCR)

RNA extraction, cDNA synthesis and RTqPCR were performed as described before. For information about primers used see Supplementary Methods.
Gene panel sequencing and DNA methylation analysis

Capture-based next-generation DNA sequencing was performed on a NextSeq 500 instrument (Illumina) and DNA methylation analysis done as described. For details see Supplementary Methods.

Droplet digital PCR (ddPCR)

All ddPCR experiments were conducted on the QX200 Droplet Digital PCR System (Bio-Rad) and analysis was performed using QuantaSoft Analysis Pro software (Bio-Rad), for details see Supplementary Methods.

BH3 profiling and caspase-3 activity

BH3 profiling experiments were performed as described before, for details as well as on caspase-3 activity measurement see Supplementary Methods.

Measurement of mitochondrial membrane potential (MMP)

Loss of MMP was measured using a tetramethyl rhodamine ethylester (TMRE) mitochondrial membrane potential assay kit (Cayman chemicals, Item No. 701310) following the manufacturer’s instructions. For details see Supplementary Methods.

Data availability and statistics

Data were generated by the authors and are available on request. For details on statistics see Supplementary Methods.
Results

Establishment of new patient-derived PA in vitro models

Short-term cultures from native tissue of three fresh PA tumors (Supplementary Table 1) were transduced with pCW57.1 GFP-TAg, allowing for doxycycline inducible expression of SV40-TAg to circumvent OIS. Culture of transduced primary cells in the presence of doxycycline allowed for establishment of long-term cell lines (>15 passages). DNA-methylation patterns, established for molecular classification of brain tumors,33 revealed a stable methylome of all patient-derived cell lines at passage 7 and 15 clustering close to the DNA-methylation group of the corresponding primary samples (Figure 1A). Copy number plots (CNPs) derived from these DNA-methylation analyses were flat as expected for PA (Supplementary Figure 1A).35 CNPs indicated the presence of a BRAF-fusion in DKFZ-BT308 and DKFZ-BT317 (Supplementary Figure 1A), confirmed by ddPCR and RTqPCR on DNA and mRNA level (Supplementary Figure 1B-E). Gene panel sequencing detected a BRAF V600E mutation in DKFZ-BT314 (Supplementary Table 2), confirmed by ddPCR (Supplementary Figure 1F). ddPCR experiments indicated a nearly 100% tumor cell purity of all new models (Supplementary Figure 1B, C and F) explaining the close proximity but distinct difference from primary PA bulk tumors (containing also microenvironmental cells) on DNA-methylation level (Figure 1A).

The MAPK Pathway Activity Score (MPAS)36 (Figure 1B) as well as phosphorylation status of ERK1/2 (Thr202/Tyr204) (Supplementary Figure 1 G) revealed activation of the MAPK-pathway in all cell lines. Robust proliferation of the new cell lines was observed only in the presence of doxycycline (proliferation; +DOX), and withdrawal of doxycycline (OIS; -DOX) led to growth arrest in all cell lines (Figure 1C) following repression of SV40 TAg (Figure 1D). All PA cell lines showed expression of the senescence marker p21Cip1 on protein level (Figure 1E) and marked SA-beta-galactosidase positivity (Supplementary Figure 1H) in OIS mode. Gene expression analysis revealed upregulation of gene sets known to be upregulated in senescence (BUHL SASP20, FRIDMAN SENESCENCE UP37 and BT66 OIS UP24) and downregulation of gene sets known to be downregulated in senescence (BT66 OIS DOWN24).
in all three new cell lines in the absence of doxycycline (Figure 1E). In summary, we established three new patient-derived PA cell lines that preserved molecular characteristics of their primary tumors and that are uniquely suitable for drug studies in OIS conditionally induced by repression of SV40 TAg after expansion.

BH3 mimetics with high binding affinity for BCL-XL preferentially decrease metabolic activity of PA cells in oncogene-induced senescence

Navitoclax, a BH3 mimic targeting BCL-2, BCL-XL and BCL-W, decreased metabolic activity of DKFZ-BT66, -BT314 and -BT317 at nano-molar concentrations, validating the preferential susceptibility in OIS previously described in DKFZ-BT66 in two more models (Figure 2A). DKFZ-BT308 was relatively resistant to navitoclax in OIS and proliferation.

To identify the precise inhibitory profile needed for the induction of senolysis in PA, we tested a comprehensive set of BH3 mimetics with different inhibitory profiles (Table 1), compared to chemotherapeutics and MEK inhibitors (Figure 2B). Vincristine and vinblastine showed high DSS (indicating high sensitivity) only in proliferation (DSS>30 for all cell lines) but not in OIS (DSS = 0), as expected. The DSS for venetoclax, a BCL-2 selective inhibitor, MCL-1 inhibitors (A-1210477, S63645, AZD5991) and MEKis (trametinib, selumetinib, binimetinib) were overall low (<7), independent of OIS and proliferation. In contrast, the DSS for all inhibitors with strong affinity to BCL-XL indicated sensitivity of DKFZ-BT66, -BT314 and -BT317 in OIS (DSS >33). IC\textsubscript{50} values in OIS for all tested BCL-XLi were in the nanomolar range (Figure 2C) and the cell lines were among the most navitoclax-sensitive compared to 751 cell lines from the GDSC2 database (ranks: DKFZ-BT66:10/755; DKFZ-BT314: 39/755; DKFZ-BT317: 43/755) (Supplementary Figure 2A). DKFZ-BT308 was relatively resistant to all BCL-XLi, in OIS and proliferation. The differential DSS (dDSS; DSS\textsuperscript{OIS} minus DSS\textsuperscript{proliferation}) for all BCL-XLi were positive in the BCL-XLi sensitive cell lines, indicating a preferential susceptibility in senescence and a senolytic mechanism of action (Figure 2D). No positive dDSS was observed for BH3 mimetics without BCL-XL affinity or MEK-inhibitors. The dDSS of vinca alkaloids were
negative, indicating sensitivity only in proliferation. The IC$_{50}$s of the BCL-XLi navitoclax, A-1331852 and A-1155463 were at least 8.5-fold higher in non-neoplastic brain-derived control NHA TAg cells compared to the BCL-XLi sensitive PA cell lines (Figure 2E and Supplementary Figure 2B). Taken together, we identified senolytic activity of BCL-XLi on metabolic activity level in senescent PA cell line models.

**BCL-XL is expressed in primary PA, PA tumor cell lines and upregulated in oncogene-induced senescence**

Following the observation of exclusive sensitivity to BCL-XLi in the responsive PA cell lines we investigated the expression of anti-apoptotic BCL-2 members in PA with a specific focus on BCL-XL. Expression of $BCL2L1$ (encoding BCL-XL) mRNA (Figure 3A) and abundance of BCL-XL protein was confirmed in in primary PA samples (Figure 3B and C; Supplementary Table 3). Of note, $BCL2L1$ mRNA expression was significantly higher in normal CNS tissue but BCL-XL protein was significantly higher expressed in PA. The four PA cell lines expressed $BCL2L1$/BCL-XL and showed upregulation in OIS compared to proliferation (Figure 3D and E).

Upregulation in OIS was not detected for the remaining anti-apoptotic Bcl-2 members BCL-2, BCL-W and MCL-1 (Supplementary Figure 3A and B). $BCL2$ transcription was significantly lower in the PA cell lines compared to bulk PA and no BCL-2 protein was detected by Western blot in the PA cell lines, consistent with the observation that BCL-2, in contrast to BCL-XL, is predominantly expressed in the tumor infiltrating immune cells (Supplementary Figure 3C). In summary, we confirmed BCL-XL target expression in primary PA and cell lines, and showed upregulation of BCL-XL in senescence in PA cell lines.

**BCL-XL inhibition reduces viable cell number and induces mitochondrial apoptosis in senescent PA cells**

Navitoclax treatment led to displacement of anti-apoptotic BCL-XL from pro-apoptotic BAK in all PA cell lines demonstrating target engagement (Figure 4A). The BCL-XLi A-1331852 and
navitoclax significantly reduced the viable cell numbers at a concentration of 40 nM and higher in DKFZ-BT314 and -BT317 (Figure 4B). In contrast, the BCL-2 inhibitor venetoclax and the MCL-1 inhibitor S63845 did not significantly impact the number of viable cells in DKFZ-BT314 and only to a limited extent in DKFZ-BT317 (Figure 4B). This observation was in line with the respective metabolic activity data: metabolic activity IC\(_{50}\)s for venetoclax and S63845 were >3µM in both models. A-1331852 and navitoclax led to a significant loss of MMP in the senescent PA cell lines DKFZ-BT314 and DKFZ-BT317 (Figure 4C) and an activation of caspase-3 (Figure 4D) indicating induction of mitochondrial apoptosis. Caspase 3 activation upon navitoclax treatment was significantly lower in proliferating DKFZ-BT314 and DKFZ-BT317 indicating a preferential induction of apoptosis in senescent PA cells (Supplementary Figure 4 A). Taken together, this data confirmed the on-target effect and an in-class effect of BCL-XL on viable cell number, mediated by mitochondrial apoptosis in senescent PA cells.

**Senescent PA cell lines depend on BCL-XL to maintain viability and to prevent mitochondrial outer membrane permeabilization (MOMP)**

DKFZ-BT308 showed relative resistance to BCL-XL. We therefore tested the dependence on BCL-XL protein in all four PA models in OIS. The number of viable cells relative to non-silencing control shRNA was reduced in all cell lines after BCL-XL knockdown with comparable efficiency (Supplementary Figure 4B and C), indicating a dependence on the protein for cell survival (Figure 4E), however without conclusive difference between the BCL-XL resistant line and the sensitive lines (Figure 4E). Treatment with the two specific synthetic sensitizer BH3 peptides, BCL-2-antagonist of cell death (BAD; specific for dependence on BCL-2/BCL-XL) and activator of apoptosis harakiri (HRK; specific for dependence on BCL-XL) induced mitochondrial outer membrane permeabilization (MOMP) as measured by cytochrome c release to a similar extent in BCL-XL sensitive (DKFZ-BT66) and resistant (DKFZ-BT308) cells (Figure 4F and G). In summary, target dependence and BCL-XL dependent priming were validated, however without differences to explain the low BCL-XL sensitivity of DKFZ-BT308.
A xenobiotics metabolism gene set upregulated in DKFZ-BT308 predicts navitoclax resistance

We performed GSEA to discover potential differences between BCL-XL sensitive and resistant cell lines at mRNA level. 50 hallmark gene sets covering a wide range of biological processes were investigated in DKFZ-BT308 versus the BCL-XLi sensitive lines in OIS (Figure 5A). Among all gene sets, only the drug metabolism gene set “HALLMARK_XENOBIOTIC_METABOLISM” was significantly enriched in DKFZ-BT308 compared to the remaining cell lines (Figure 5B). 38 genes of the gene set contributed to the core enrichment and were used to derive a new BT308_UP signature (Supplementary Table 4). To test a potential relation of this signature to BCL-XLi resistance, we used an independent data set of 751 cell lines (GDSC2) (Figure 5C). GSEA and ssGSEA confirmed a significant enrichment of the BT308_UP signature in the group of navitoclax resistant cell lines (Figure 5D and E), as well as ABT-737 (another BCL-XLi) resistant cell lines (Supplementary Figure 5A). Binary logistic regression analysis revealed that expression of BT308_UP (independent continuous predictor) was able to predict “navitoclax resistance” at an optimal ssGSEA threshold score between -217 and -197 with $p = 3.37 \times 10^{-9}$, a sensitivity of 85.53% and a specificity of 75.00% (Supplementary Figure 5B). Primary PAs showed a wide variability of BT308_UP signature expression, but all primary samples tested expressed the signature to a lower extent compared to the resistant cell line DKFZ-BT308 (Figure 5F). Taken together, we identified upregulation of a drug metabolism gene set in DKFZ-BT308 that could explain the phenotypical differences in BCL-XLi sensitivity.
Discussion

OIS is a well-known feature of PA but currently not therapeutically exploited. The lack of preclinical data supporting senolytic drugs can be explained by the lack of PA OIS models. In this respect the three new cell lines (DKFZ-BT308, DKFZ-BT314 and DKFZ-BT317) described in the present study are outstanding because they can be analyzed in OIS and complement the previously published DKFZ-BT66 model.

Using our four PA models we discovered that BCL-XL is essential for PA cells in OIS and BH3 mimetics induce senolysis via inhibition of BCL-XL. Several recent studies have highlighted the potential role of senolysis in glioma therapy including BCL-XL inhibition. However, these studies are fundamentally different from the pediatric low-grade glioma/OIS background of the present work since they investigated senolytics in another glioma type (adult high-grade glioma) and in the context of a different form of senescence (therapy-induced senescence).

Beyond the first proof of BCL-XL as a target in PA, our study provides several translational implications. We confirmed abundance of the target BCL-XL in primary PA and higher expression of BCL-XL protein compared to normal CNS tissue. Of note, a broad variability in BCL2L1/BCL-XL expression among all primary samples was observed. However, this may not necessarily indicate differences in BCL-XLi susceptibility, as BCL2L1 expression levels per se were not found to be predictive for dependence on BCL-XL in a recent study using different cancer cell lines. Several BCL-XLi are already in clinical testing and therefore potentially available for studies in PA. Thrombocytopenia was a major dose-limiting toxicity observed in navitoclax phase I trials. However, adjusted treatment schedules led to improved tolerability in adult patients and two other BCL-Xli, AZD0466 and pelcitoclax, have shown lower platelet toxicity while maintaining anti-tumor activity. The BCL-Xli responsive PA models were among the most navitoclax sensitive cell lines compared to cell lines from the GDSC2 database. The IC50 values of lower than 300 nM seem clinically relevant in the context of a navitoclax Cmax of 6607 ± 3262 nM at RP2D. No data is available on blood brain barrier (BBB) penetration of clinically available BCL-XLi. The relatively high molecular weight of navitoclax (974.61 g/mol), AZD4320 (954.5 g/mol) and pelcitoclax (1159.78 g/mol) might...
militate against an effective BBB penetrance per se. However, many PAs have a disrupted BBB, which is evident from the uptake of MR contrast agent in these tumors\textsuperscript{49,50}, indicating that high molecular weights do not a priori preclude effective tumor concentrations of these compounds.

The BCL-XL\textsubscript{i} resistant cell line DKFZ-BT308 differed from the sensitive cell lines in expression of a xenobiotics and drug metabolism related gene set. This gene set was able to predict navitoclax resistance in an independent dataset, indicating that the genes differentially regulated in the resistant PA cell line are related to and involved in the observed resistance phenotype. Moreover, the BT308\textunderscore{UP} signature was also enriched in cells resistant to the BH3 mimetic ABT-737, indicating an in-class relevance of this gene set. While resistance to BH3 mimetics has been linked to upregulation of non-inhibited BCL-2 members\textsuperscript{51}, innate resistance caused by upregulation of drug metabolism genes has not been described in the context of BH3 mimetics so far and represents a new and possibly general observation that might be important also for many other cancer entities. A potential limitation of our observation is the lack of validation of this signature for the other BH3 mimetics used, as independent sensitivity data for these BH3 mimetics was not available. Expression of BT308\textunderscore{UP} is variable within primary PA samples according to the presented data. Although the larger proportion of primary tumors show an expression of the signature similar or lower compared to our sensitive cell lines, a smaller proportion showed higher expression of the BT308\textunderscore{UP} signature, possibly indicative of relative resistance to BH3 mimetics. The clinical relevance and predictive validity of the signature described here needs to be prospectively validated in a clinical BCL-XL\textsubscript{i} trial.

In summary, our study provides the first reported target for senolytic treatment of PA cells. In contrast to chemotherapeutics and MEKi currently used in the treatment of PA, BCL-XL\textsubscript{i} induce apoptosis in PA cells in OIS. A main limitation of this study, as for most preclinical pLGG studies, is the lack of \textit{in vivo} data. To our knowledge, there are no \textit{in vivo} models of true molecular PA available to date that could be used for testing of senolytic drugs in the context of OIS to surmount this obstacle. Based on the \textit{in vitro} data presented here, BCL-XL inhibition
is a promising treatment approach and the first attempt to target the so far un-targetable senescent compartment of PA tumors. The translation of our findings into clinical trials exploring the safety and efficacy of clinically available BCL-XL in PA patients is urgently needed.
Acknowledgements

We thank the patients and families for consent to collect and use tumor material. We thank Daniela Kuhn, Isabel Büdenbender, Carina Müller and Anna Laudat for excellent technical support. J.P.M.B. is grateful to the Great Ormond Street Hospital Charity and NIHR Biomedical Research Centre at University College London Hospitals NHS Foundation Trust and University College London. We thank the Microarray Unit of the Genomics and Proteomics Core Facility, DKFZ, for providing excellent Expression Profiling services.
References


Figure Caption

**Figure 1:** Characterization of new patient-derived PA *in vitro* models DKFZ-BT308, DKFZ-BT314 and DKFZ-BT317. A t-SNE analysis of DNA methylation profiles of pediatric brain tumors (selected tumor types from mnp V12.3 reference set; www.molecularneuropathology.org), midline PA (PA_MID) and posterior fossa PA (PA_PF) enlarged on the right. Dashed circles: PA cell lines from p7 and p15 (passage 7 and 15), closed circles: corresponding primary samples. B MPAS signature ssGSEA scores. normal: normal cerebellum, PA: primary PA. *** Tukey multiple comparisons of means adjusted p-value: 2.7 x10^-6. Expression data: ps_mkheideli_mkdkfz209_u133p2. C Cell counts of PA cell lines in proliferation or OIS. DOX: doxycycline. DKFZ-BT66: historical data. D Western blot of SV40 TAg and p21^Cip1^ OIS: 5 days of doxycycline (DOX) withdrawal. E ssGSEA z-scores of senescence gene sets. n=3 independent expression samples per condition. DOX: doxycycline.

**Figure 2:** Impact of BH3 mimetics on metabolic activity. (OIS (-DOX): oncogene-induced senescence, 5 days doxycycline withdrawal; proliferation (+DOX): +1μg/ml doxycycline. D: DMSO; BCL-Xli: BCL-XL inhibitors) A Relative metabolic activity after treatment with navitoclax (ATP-based measurement of viable cells; mean +/- SD of at least n=3 biological replicates). B DSS (DSS3) of BH3 mimetics, chemotherapeutics and MEKi in OIS and proliferation (n=3 biological replicates for each drug, cell line and mode). C Plot of log10 of absolute IC$_{50}$ values of indicated drugs in OIS (n=3 biological replicates +/-SD). 10µM (highest concentration applied) was assumed whenever insufficient drug effects precluded IC$_{50}$ calculation. D Mean differential DSS (dDSS) of indicated drugs across three BCL-XLi sensitive PA cell lines (+/-SD). dDDS = DSS of proliferating cells - DSS of cells in OIS. E Relative metabolic activity (ATP-based measurement of viable cells) after treatment withnavitoclax in normal human astrocytes NHA TAg compared to PA cell lines in OIS (mean +/- SD; at least n=3). Numbers in dose-response plots are absolute IC$_{50}$.
Figure 3: **BCL-XL expression in PA.** (OIS (-DOX): oncogene-induced senescence, 5 days doxycycline withdrawal; proliferation (+DOX): +1µg/ml doxycycline) A **BCL2L1** mRNA expression in primary PA compared to normal cerebellum. Expression data: ps_mkheidel_mkdkfz209_u133p2. Unpaired t-test: * p<0.05. B H-scores of BCL-XL protein staining intensity in 75 primary PA samples compared to 16 inconspicuous CNS tissues adjacent to low-grade gliomas. Unpaired t-test: * p<0.05. C Exemplary microscopic images of BCL-XL immunohistochemistry in PA showing strong (PA 15), medium (PA 23) and weak (PA 4) staining. neg. ctrl.: negative control, muscle tissue. Scale bars indicate a distance of 100 µm. D **BCL2L1** mRNA expression in four PA cell lines in proliferation vs. OIS compared to normal cerebellum (n=18) and primary PA (n=191) (ps_mkheidel_mkdkfz209_u133p2). Unpaired t-test: * p<0.05, ** p<0.01, *** p<0.001. E Western blot of BCL-XL protein in proliferation vs. OIS mode.

Figure 4: **On-target activity and cell death induction upon BCL-XLi treatment.** (OIS (-DOX): oncogene-induced senescence; 5 days doxycycline withdrawal) A Western blot of BCL-XL and BAK after immunoprecipitation (IP) of BCL-XL after 4h treatment of PA cells in OIS with 1 µM navitoclax. B Relative viable attached cells after 72h of drug treatment (mean +/- SD, n=3 biological replicates). Grey boxes indicate drugs’ inhibitory profiles (K_i <1nM). Unpaired t-test: * p<0.05, ** p<0.01, *** p<0.001. ****p<0.0001 (comparison to DMSO). C Loss of mitochondrial membrane potential (MMP) measured by TMRE incorporation after 24h treatment (mean +/- SD, n=3). Unpaired t-test. * p<0.05, ** p<0.01, *** p<0.001 (comparison to DMSO). D Relative caspase 3 activity after 24h of treatment with 100 nM navitoclax or 100 nM A-1331852, control: DMSO (mean +/- SD, n=3). E Relative viable cells 14 days after BCL-XL knock-down (100%: control shRNA) (mean +/- SD, n=3). * Tukey multiple comparisons of means adjusted p-value <0.01. F, G Relative cytochrome c released into the cytosol after 1h of treatment with 10µM BAD peptide (F) or 100µM HRK peptide (G), respectively, relative to DMSO (mean +/- SD, n= at least 3). rel.: relative; c: concentration.
Figure 5: Identification of a gene signature differentiating the BCL-XL\textit{i} resistant cell line DKFZ-BT308 from BCL-XL\textit{i} sensitive cell lines. \textbf{A} Flowchart depicting the deduction of BT308\textsubscript{UP} signature. \textbf{B} GSEA of the HALLMARK\_XENOBIOTIC\_METABOLISM gene set in BT308 vs. BCL-Xli sensitive PA cell lines. \textbf{C} Work-flow of BT308\textsubscript{UP} signature validation in the independent GDSC2 dataset. \textbf{D} Comparison of BT308\textsubscript{UP} signature expression in the GDSC group “navitoclax resistant” vs. the GDSC group “navitoclax sensitive”. \textbf{E} ssGSEA scores of BT308\textsubscript{UP} signature in the two GDSC groups “navitoclax resistant” and “navitoclax sensitive”. \textbf{F} BT308\textsubscript{UP} signature ssGSEA z-scores in the four PA cell lines in OIS mode (n=3 samples per cell line) and in primary PA (ps\_mkheidel\_mkdkfz209\_u133p2). GSEA; geneset enrichment analysis; ssGSEA; single sample GSEA. NES: normalized enrichment score; norm.: normalized; FDR: false discovery rate.