1 Increased apoptotic priming of glioblastoma enables therapeutic targeting by

2 BH3-mimetics

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- 24 Running title: Overcoming apoptotic resistance in GBM
- 25 Keywords: apoptosis/BH3-mimetics/glioblastoma/resistance/stem cells
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28 Abstract

29	IDH wild-type glioblastoma (GBM) is the most prevalent malignant primary brain tumour in
30	adults. GBM typically has a poor prognosis, mainly due to a lack of effective treatment
31	options leading to tumour persistence or recurrence. Tackling this, we investigated the
32	therapeutic potential of targeting anti-apoptotic BCL-2 proteins in GBM. Levels of anti-
33	apoptotic BCL-xL and MCL-1 were consistently increased in GBM compared with non-
34	malignant cells and tissue. Moreover, we found that relative to their differentiated
35	counterparts, patient-derived GBM stem-like cells also displayed higher expression of anti-
36	apoptotic BCL-2 family members. Surprisingly, high anti-apoptotic BCL-xL and MCL-1
37	expression correlated with heightened susceptibility of GBM to BCL-2 family protein-
38	targeting BH3-mimetics. This is indicative of increased apoptotic priming. Indeed, GBM
39	displayed an obligate requirement for MCL-1 expression in both tumour development and
40	maintenance. Investigating this apoptotic sensitivity, we found that sequential inhibition of
41	BCL-xL and MCL-1 led to robust anti-tumour responses in vivo, in the absence of overt
42	toxicity. These data demonstrate that BCL-xL and MCL-1 pro-survival function is a
43	fundamental prerequisite for GBM survival that can be therapeutically exploited by BH3-
44	mimetics.

45 Introduction

46	In adults, IDH wild-type glioblastoma (IDHwt GBM) is the most prevalent and malignant
47	primary brain tumour (1, 2). Despite current multimodal treatment, comprising surgical
48	resection with adjuvant radiotherapy and alkylating chemotherapy, the median survival in
49	newly diagnosed patients remains poor at less than 12 months (3, 4). Resistance to
50	conventional radio- and chemotherapy primarily emerges from persistent cancer stem cells,
51	a tumourigenic subpopulation of GBM cells, consisting of heterogenous subclones and
52	capable of self-renewal (5, 6). Therefore, targeting cells with stem-like capabilities is
53	essential to develop effective treatment options and improve patient survival.
54	
55	Treatment resistance can often be attributed to cells circumventing therapy-induced cell
56	death. Apoptosis is an evolutionarily conserved type of cell death with broad ranging
57	importance in biology (7). The intrinsic (mitochondrial) pathway of apoptosis is controlled by
58	pro- and anti-apoptotic members of the B cell lymphoma 2 (BCL-2) family that regulate
59	mitochondrial outer membrane integrity (8). During apoptosis, pro-apoptotic BCL-2 proteins
60	cause mitochondrial outer membrane permeabilisation or MOMP. This leads to the release
61	of mitochondrial intermembrane space proteins, including cytochrome <i>c</i> , that activate
62	caspase proteases leading to apoptotic cell death (8).
63	

64 Increased anti-apoptotic BCL-2 protein expression has been described in a wide range of 65 solid cancers and is often linked with insensitivity to conventional chemotherapy (9-11). 66 Recently, a new class of chemotherapeutics called BH3-mimetics have been developed that 67 target pro-survival BCL-2 function, sensitising to cell death. BH3-mimetics have proven to be 68 highly effective in haematologic malignancies. For instance, venetoclax (ABT-199), a BCL-2 69 targeted BH3-mimetic (12), is in clinical use for chronic lymphocytic leukaemia (CLL) (13) 70 and acute myelogenous leukaemia (AML) (14, 15). CLL cells typically express high levels of 71 anti-apoptotic BCL-2 protein. Nevertheless, the high intrinsic apoptotic sensitivity - also 72 called apoptotic priming - of CLL renders it sensitive to venetoclax. For solid cancers,

73 venetoclax is currently being tested in combination with conventional chemotherapeutic 74 agents. The combination of venetoclax and tamoxifen has progressed to early phase clinical 75 trials in patients with estrogen receptor positive (ER+), high BCL-2 expressing breast cancer 76 (16). Other BH3-mimetics developed to target BCL-xL and MCL-1 have shown promising 77 pre-clinical results in combination with inhibitors of MEK1/2 for solid cancers harbouring 78 oncogenic mutations in the MAPK pathway (17-19). Approximately 10% of GBM carry a 79 mutation of isocitrate dehydrogenase 1 (IDH1) (1), which has been linked with increased 80 sensitivity to treatment with BH3-mimetics targeting BCL-xL (20). Furthermore, previous 81 studies have proposed BCL-xL as a treatment target in combination with ionising radiation 82 (21) and other chemotherapeutics (22) in GBM.

83

84 Because tumours retain characteristics of their tissue origins, brain derived glial cancers 85 exhibit defined cellular hierarchies found in brain development and homeostasis (23-25). 86 During central nervous system development, anti-apoptotic BCL-2 family proteins play a 87 pivotal role in promoting cell survival (26, 27) while with adulthood the brain becomes 88 refractive to apoptosis (28). Given this important role in cell survival, we hypothesised that 89 GBM, while phenocopying the developing brain, might display similar anti-apoptotic survival 90 dependencies. Indeed, we found increased levels of the major pro-survival proteins in GBM, 91 specifically within the stem-cell enriched population. Surprisingly, high BCL-xL and MCL-1 92 expression correlates with increased apoptotic sensitivity, demonstrating that GBM stem-like 93 cells are primed for apoptosis. Exploiting this, we found that sequential dosing of BCL-xL 94 and MCL-1 targeting BH3-mimetics enables effective treatment responses both, in vitro and 95 in vivo. This could offer a therapeutically tractable approach for patients with IDHwt GBM.

96 Results

97 High anti-apoptotic BCL-2 family protein expression correlates with increased BH3-mimetic 98 sensitivity in GBM 99 Cancer stem cells are proposed to give rise to GBM and contribute to therapeutic resistance 100 (23). We therefore sought to assess the apoptotic sensitivity of GBM stem-like cells (GSC) 101 by treating them with BH3-mimetics with selectivity for BCL-2, BCL-xL and MCL-1. For this 102 purpose, we used a panel of patient-derived IDHwt GSC, cultured under conditions to 103 maintain their tumour specific phenotype and stem cell properties (29, 30). Cell viability was 104 measured using IncuCyte live-cell imaging and SYTOX Green exclusion. Importantly, three 105 cell lines (G1, G7 and R24 GSC) were sensitive to A-1331852, a selective BCL-xL 106 antagonist (31) whereas two cell lines (R9 and R15 GSC) displayed sensitivity to S63845, a 107 potent and selective MCL-1 inhibitor (32) (Figure 1A,B). Moreover, the commonly used 108 GBM cell line U87MG displayed increased sensitivity to BCL-xL inhibition when cultured 109 under stem cell-enriching conditions. One cell line (E2 GSC) was resistant to all single agent 110 treatments. Treatment with venetoclax (ABT-199), a BCL-2 specific inhibitor, induced no 111 more than 26% cell death in any GSC and therefore was comparably inefficient. Collectively, 112 these data show that the majority of tested GSC display survival dependence on anti-113 apoptotic BCL-2 family function. 114 We next used immunoblotting to determine if the individual apoptotic sensitivity of the 115 patient-derived GSC corresponded to anti-apoptotic BCL-2 protein expression. In 116 comparison to human differentiated astrocytes, all GSC exhibited higher expression of BCL-117 xL and MCL-1 and partially higher expression of BCL-2 (Figure 1C). Consistent with their 118 origins, GSC expressed higher levels of neural stem cell marker SOX2 (33), while cell 119 lineage specific GFAP was more abundant in astrocytes (34). Subsequently, we investigated 120 whether anti-apoptotic BCL-2 protein expression also differed in GBM tumours and adjacent 121 brain tissue. Using immunohistochemistry (IHC) we compared BCL-xL and MCL-1 122 expression in matched specimens of three patients diagnosed with GBM. In all cases, MCL-

123 1 and BCL-xL were increased in the tumour cores compared to the predominantly non-

124 tumorous margins (Figure 1D,E). Extending this analysis, we determined MCL-1 and BCL-125 xL mRNA expression in different glioma subtypes and normal brain tissue using the publicly 126 available REMBRANDT database. In line with our IHC analysis, BCL-xL and MCL-1 mRNA 127 levels were more expressed in GBM compared with lower grade gliomas and normal brain 128 tissue (Supplementary Figure 1A,B). These data demonstrate specific sensitivities of 129 patient-derived GSC to individual BH3-mimetics and increased expression of anti-apoptotic 130 BCL-2 proteins in both primary GBM tumour tissues and GSCs. 131 132 Anti-apoptotic MCL-1 is required for the growth and survival of GBM 133 While it has previously been shown that GBM tumoursphere formation is promoted by high 134 BCL-xL expression (35), little is known about the role of MCL-1 on GBM growth and 135 maintenance. To explore the importance of MCL-1 in GBM formation and growth in vivo, we 136 selected a tumourigenic cell line (G7 GSC) and deleted MCL-1 using CRISPR/Cas9 genome 137 editing. Western blot analysis confirmed efficient MCL-1 deletion, while the expression of 138 BCL-xL, SOX2 and GFAP was increased in the MCL-1^{CRISPR} cells (Supplementary Figure **2A**). MCL-1^{CRISPR} GSC were found to proliferate at the same rate and retained a similar 139 capability to form neurospheres as their vector^{CRISPR} counterparts (Figure 2A, 140 141 Supplementary Figure 2B). We next investigated whether MCL-1 was required for tumourigenesis in vivo. iRFP-labelled vector^{CRISPR} and MCL-1^{CRISPR} G7 GSC were 142 143 orthotopically injected in CD-1 nude mice and tumour growth was monitored with cranial 144 magnetic resonance imaging (MRI) and iRFP signal detection (36). We observed a 145 substantial impairment of tumour growth in MCL-1 deleted tumours (Figure 2B,C; 146 Supplementary Figure 2C) that was reflected in the significantly prolonged survival of these 147 mice (Figure 2D). Importantly, IHC analysis of the end-stage tumours revealed an outgrowth of MCL-1 proficient tumour cells in the MCL-1^{CRISPR} xenografts (Supplementary Figure 2D). 148 149 In contrast to the *in vitro* analysis, these data reveal a key role for MCL-1 in initiation and 150 growth of GBM in vivo and identify MCL-1 as a promising therapeutic target. Our results also 151 support an important pro-survival role for anti-apoptotic MCL-1 in GBM.

152

153 GSC display increased apoptotic priming and can be effectively killed by dual BCL-xL, MCL-154 1 inhibition 155 Currently, no treatment regimen is able to achieve long-time remission of GBM, with tumours 156 inevitably developing resistance to treatment and recurring, eventually leading to patient 157 death (37). Anti-apoptotic BCL-2 family members have overlapping binding affinities for 158 several pro-apoptotic BH3-only proteins (8). We asked whether GBM might circumvent 159 single inhibitor treatment by compensatory upregulation of untargeted anti-apoptotic 160 proteins. Indeed, upon treatment with BCL-xL inhibitor A-1331852 or navitoclax (ABT-737), 161 an inhibitor of BCL-xL, BCL-2 and BCL-w, we found that levels of MCL-1 protein were 162 increased in surviving GSC (Figure 3A, Supplementary Figure 3A). We reasoned that 163 GBM might counteract drug-induced neutralisation of BCL-xL function via increased MCL-1 expression. To address this, we treated MCL-1^{CRISPR}G7 and R24 GSC with the BCL-xL 164 165 inhibitors A-1331852 or ABT-263. Cell viability was measured by live-cell IncuCyte imaging 166 with Sytox Green exclusion or in a clonogenic survival assay. In all cases, MCL-1 deletion 167 significantly increased cellular sensitivity to the BCL-xL specific antagonist and navitoclax 168 (Figure 3B, Supplementary Figure 3B). Similarly, dual inhibition of BCL-xL and MCL-1 with 169 A-1331852 and S63845 displayed a substantial combinatorial effect resulting in up to 100% 170 cell death across a range of GSC, determined in both short term cell viability assays and 171 long-term clonogenic survival assays (Figure 3C,D; Supplementary Figure 3C,D). This 172 effect was observed even at 10-fold decreased doses compared to effective single 173 treatment. Verifying on-target engagement of mitochondrial apoptosis, combined MCL-1 and 174 BCL-xL inhibition led to Caspase 3 and PARP-1 cleavage as well as cell death in a BAK, 175 BAX and caspase-dependent manner (Figure 3E, Supplementary Figures 3E-F). 176 Sensitivity of GBM cells to chemotherapy and ionising radiation inversely correlates with 177 tumour cell stemness (6, 38, 39). We therefore hypothesised that the differentiated 178 counterparts (DIFF) of the patient-derived GBM stem-like cells may be more sensitive 179 towards BH3-mimetic treatment. To ensure comparable culture conditions for GSC and

180 DIFF, we conducted these experiments using 1% FCS containing Ad-DMEM medium during 181 the experimental procedure. Cells were treated with A-1331852 and S63845 to inhibit BCL-182 xL and MCL-1 respectively and cell viability measured by live-cell IncuCyte imaging and 183 Sytox Green exclusion. Following treatment with single BH3-mimetic, we found that cell 184 viability was largely comparable for DIFF and GSC (Supplementary Figure 3G). 185 Surprisingly, following dual MCL-1 and BCL-xL inhibition, E2 and R15 GSC were more 186 sensitive than DIFF cells; while >50% cell death was observed within about 5 hours in GSC, 187 it was not observed in DIFF cells until 16 hours (Figure 3F). Moreover, 100% cell death was 188 not achieved in either of the DIFF cell lines. Together, these data suggest that GSCs are 189 more primed for apoptotic cell death than DIFF cells. To investigate this further, we 190 compared expression of pro- and anti-apoptotic BCL-2 proteins in the paired cell lines. 191 Although more sensitive to apoptosis, GSCs displayed higher levels of anti-apoptotic BCL-2 192 proteins, BCL2, BCL-xL and MCL-1 than their differentiated counterparts (**Supplementary** 193 Figure 3H). In summary, these data indicate that GSC can display increased apoptotic 194 priming and reveal potent cytotoxic effects of dual-targeting BCL-xL and MCL-1. 195 196 *TrkB* signalling regulates sensitivity of GSC to anti-apoptotic treatment 197 We next sought to explore the differential priming between GSC and their isogenic 198 differentiated counterparts. To this end, bulk RNA sequencing data from E2, G7 GSC and 199 their DIFF counterparts was analysed (30). Consistent with enrichment of GSC, RNAseq 200 analysis revealed increased levels of CD34, a surface glycoprotein, first described as marker 201 for haematopoietic progenitor cells (40). Interestingly, high expression of NTRK2 mRNA was 202 detected in both GSC (Figure 4A). This finding was validated in E2 and G7 GSC as well as 203 R15 and R24 GSC via immunoblotting (Figure 4B, Supplementary Figure 4A). NTRK2 the

- 204 gene coding for the tropomycin receptor kinase B (TrkB) is primarily known for its function in
- 205 neurodevelopment inducing downstream signalling upon binding of brain-derived
- 206 neurotrophic factor (BDNF) (41). Recently, Wang and colleagues have reported a role for
- 207 TrkB-expressing cancer stem cells in GBM progression in response to BDNF stimulation by

208 differentiated tumour cells (42). As TrkB-mediated activation of MAPK and PI3K-AKT 209 signalling is generally associated with cell survival (43), we hypothesised that BDNF-210 mediated TrkB stimulation might enable GSC to evade cell death. Unexpectedly, following stimulation of GSC with BDNF or 7,8-dihydroxyflavone (7,8-DHF), a specific TrkB agonist, 211 212 we found that GSC were further sensitised to cell death following treatment with BH3-213 mimetics targeting BCL-xL and or MCL-1 (Figure 4C, Supplementary Figure 4B). This 214 sensitising effect was not observed in the DIFF cells (Figure 4D). DIFF cells express 215 significantly lower levels of the TrkB receptor and therefore prove to be comparably 216 unresponsive to BDNF stimulation (Supplementary Figure 4C). BDNF-induced TrkB 217 phosphorylation also led to increased BCL-xL protein expression, alongside stabilisation of 218 the BIM protein downstream of MAPK signalling, independently of BCL-xL and MCL-1 219 inhibition (Figure 4E). These data demonstrate a key role for BDNF-TrkB signalling in the 220 increased apoptotic priming of GSCs.

221

222 Combined BCL-xL and MCL-1 inhibition causes apoptosis in human GBM ex vivo 223 Current in vitro methodologies fail to recapitulate important aspects of the brain 224 microenvironment and tissue context. Given this, we sought to use a more physiologically 225 relevant model to investigate functional responses to BCL-xL and MCL-1 inhibition in GBM. 226 For this purpose, we developed an assay tailored to the use of freshly resected human GBM 227 to be cultured ex vivo as tissue slices that could be readily exposed to candidate drugs 228 (experimental setup illustrated in Supplementary Figure 5A). All three patients included in 229 the study were diagnosed with *IDHwt* GBM. Tissue slices were treated for 72 hours in total. 230 In all cases, we found that a combined therapy with A-1331852 and S63845 (BCL-xL and 231 MCL-1 inhibition) significantly reduced tumour cell count compared with single drug 232 treatment or control (Figure 5A-D, Supplementary Figure 5B-D). Moreover, dual treatment 233 induced a significant reduction of cell proliferation (Ki67 IHC) and amount of SOX2 positive 234 tumour cells (Figure 5E,F, Supplementary Figure 5E,F), while Caspase 3 cleavage was 235 increased. This data indicates that the dual treatment efficiently targets GBM stem-like cells

236 ex vivo. Importantly, the integrity of the brain tissue and vasculature was maintained

237 (Supplementary Figure 5G).

238 In recent years, selective MCL-1 and BCL-xL inhibitors have been developed that show 239 effective *in vivo* potency (31, 32). However, systemic exposure to both inhibitors is limited 240 due to its combined toxicity (44). Because the blood-brain-barrier is only permissible to 241 certain drugs, local or intrathecal drug application (45) might allow to circumvent systemic 242 side effects in a clinical setting. To explore potential toxicities to resident brain cells, we 243 obtained brain slices from 11-week old adult mice and exposed them to dual treatment with 244 the indicated BH3-mimetics. In regions of the subventricular zone, an important neural stem 245 cell niche, we detected only a moderate increase of cleaved Caspase 3 in a fraction of glial 246 and neural progenitor cells (GFAP and NESTIN IF stain) (Figure 5G,H). In summary, these 247 data suggest that dual targeting of BCL-xL and MCL-1 may provide a novel therapeutic 248 approach to treat GBM.

249

250 Priming with BCL-xL inhibition renders GSC vulnerable to MCL-1 inhibition, promoting

251 tumour regression in vivo

Given the potency of joint BCL-xL/MCL-1 inhibition in our *in vitro* and *ex vivo* findings, we sought to maximise this combinatorial effect whilst mitigating possible systemic toxicity. To address this, we investigated which pro-apoptotic proteins are involved in regulating intrinsic apoptosis in GBM. Upon single agent treatment with A-1331852, we observed upregulation of the BH3-only protein BIM as well as anti-apoptotic MCL-1. This was seen in both control and BAX/BAK deficient GSC (**Figure 6A**). BIM is an important BH3-only protein in the

258 canonical apoptotic pathway where it functions by regulating both BCL-xL and MCL-1

259 mediated cell death responses (46). We hypothesised that MCL-1 might bind and neutralise

260 BIM that is released by the A-1331852 complexing to BCL-xL. Accordingly,

261 immunoprecipitation of MCL-1 following treatment with A-1331852 revealed increased

262 binding of BIM to MCL-1 (Figure 6B). To explore whether this mechanism could be

therapeutically exploited, we questioned whether BCL-xL inhibition would render GSC more

264	sensitive to subsequent MCL-1 inhibition. GSC were treated with a BH3-mimetic targeting
265	either BCL-xL or MCL-1 for up to 48 hours followed by a washout and 24 hours treatment
266	pause. Subsequently the complementary inhibitor was applied for up to 48 hours. Whereas
267	prior inhibition with MCL-1 inhibitor failed to sensitise the cells to BCL-xL inhibition, pre-
268	treatment with the BCL-xL inhibitor substantially increased the susceptibility of GSC to
269	subsequent MCL-1 inhibition (Figure 6 C,D). To further investigate the relevance of BIM in
270	mediating apoptosis following BCL-xL and MCL-1 inhibition, we deleted BIM by
271	CRISPR/Cas9 genome editing (Supplementary Figure 6A). Using IncuCyte live cell
272	imaging and Sytox Green exclusion to detect cell death under treatment we observed that
273	knockout of BIM did not impede the sensitivity of G7 GSC to concurrent dual BCL-xL and
274	MCL-1 inhibition (Supplementary Figure 6B). However, after pre-treatment with BCL-xL
275	inhibitor A-1331852 G7 BIM ^{CRISPR} GSC were less primed for following MCL-1 inhibition
276	compared to their vector ^{CRISPR} counterparts (Figure 6E). These results indicate that Bcl-xL-
277	inhibition mediates sensitisation of GSC to subsequent MCL-1 neutralisation via pro-
278	apoptotic BIM.
279	Finally, we aimed to investigate the potency of alternating BH3-mimetic treatments in GBM
280	in vivo. Due to the poor blood-brain-barrier penetrance of ABT-263, we chose a
281	subcutaneous model. As the GSC used in our study do not grow as subcutaneous
282	xenografts we explored whether human U87MG respond to combined and alternating
283	inhibition of BCL-xL and MCL-1 in like manner to GSC. Indeed, we found that dual inhibition
284	of BCL-xL and MCL-1 induced substantial cell death in U87MG neurospheres
285	(Supplementary Figure 6C). Testing different treatment regimens using a clonogenic
286	survival assay, we found a profound decrease in colony formation upon alternating treatment
287	with ABT-263 and S63845 (Supplementary Figure 6D). In our in vivo cohort, mice were
288	treated with vehicle or alternating therapy with ABT-263 and S63845 every 48 hours for two
289	weeks upon tumour establishment (treatment schematic illustrated in Supplementary
290	Figure 6E). Compared to mice receiving vehicle control, mice treated with the sequential
291	therapy showed significant attenuation and/or regression of tumours (Figure 6F). Most

- importantly, we observed significantly improved survival in mice treated with ABT-263,
- followed by S63845 (Figure 6G). One mouse had a complete tumour regression after
- sequential treatment with no reoccurrence of the subcutaneous tumour over an 8-week
- follow up period. With this treatment schedule no significant weight loss (Supplementary
- 296 Figure 6F) or signs of neurological deficits were detected in mice. Histopathological analysis
- 297 of tumours reaching clinical end point showed a higher prevalence of a large central necrotic
- areas in within the treatment cohort compared to vehicle control (83% vs. 33%)
- 299 (Supplementary Figure 6G). Collectively, these data demonstrate the therapeutic potential
- 300 of sequential BCL-xL and MCL-1 inhibition in GBM.

301 Discussion

302	Largely due to a dearth of effective treatment options, GBM patients have a dismal
303	prognosis (37). Addressing this, we investigated the therapeutic potential of targeting anti-
304	apoptotic BCL-2 proteins in GBM. Our analysis revealed high expression of anti-apoptotic
305	BCL-xL and MCL-1 in GBM. Moreover, we also observed increased expression of BCL-xL
306	and MCL-1 in GBM stem-like cells - a population of cells that are key for GBM development
307	and treatment resistance in vivo (5, 6). Rather than promoting apoptotic resistance, elevated
308	anti-apoptotic BCL-xL and MCL-1 expression in GSC compared to isogenic DIFF correlated
309	with increased susceptibility to targeted inhibition using BH3-mimetics. This indicates that
310	GSC are inherently primed for apoptosis. Exploiting this, we found that GBM were sensitive
311	to BH3-mimetics targeting either MCL-1 or BCL-xL. Crucially, alternating dosing with BCL-xL
312	followed by MCL-1 specific BH3-mimetics, led to durable treatment responses with
313	preceding BCL-xL inhibition sensitising to MCL-1 inhibition in vivo. These data highlight the
314	therapeutic potential of targeting BCL-xL and MCL-1 in GBM.

315

316 Recently, highly specific and potent BH3-mimetics have been developed to specifically 317 target BCL-2, BCL-xL and MCL-1 (12, 31, 32). We used these to probe the individual 318 dependencies of GBM in a panel of patient-derived GSC. Importantly, we found that GBM 319 cells are largely dependent on BCL-xL or MCL-1 for survival, whereas BCL-2 plays a lesser 320 role. Genetic deletion of MCL-1 corroborated its key role in both promotion and maintenance 321 of GBM. Consistent with our findings, indirect targeting of MCL-1 through CDK7 inhibition, 322 causing transcriptional repression, sensitises GBM cell lines to ABT-263 (47). Further, we 323 could demonstrate that both BCL-xL and MCL-1 are highly expressed, not only in GBM 324 tumour cores, but also in GBM stem-like cells compared to their isogenic differentiated 325 counterparts and astrocytes. The high expression and dependence of GBM on anti-apoptotic 326 BCL-2 function is consistent with an increased state of apoptotic priming. As Sarosiek and 327 colleagues have demonstrated, the tissue of origin plays a major role in determining the 328 apoptotic sensitivity of a cell (28), therefore the high dependency of stem-like cells on BCL-

329 xL and MCL-1 might relate to their resemblance of cerebral precursor cells. Both anti-

apoptotic proteins play a major role in neurogenesis, while with brain maturation neurons

331 become refractory to apoptotic cell death (26, 28).

332

333 Unexpectedly, we observed that in comparison to their differentiated counterparts, stem-like 334 cells were more susceptible to BH3-mimetic treatment. Investigating the basis of the 335 differential apoptotic priming we identified NTRK2, a stem-cell specific surface receptor (42) 336 as a key component. NTRK2 signalling, mediated by the soluble brain derived neurotrophic 337 factor BDNF, plays a major role in cell survival promotion of growth in glial tumours (42, 48). 338 Surprisingly, we found that BDNF stimulation led to increased apoptotic priming. This result 339 reinforces the notion that stem-like cells are especially dependent on the tight regulation of 340 apoptotic sensitivity. More profound understanding of the tumour-environmental context 341 should shed light on how these interactions can be therapeutically exploited to maximise 342 treatment efficacy of BH3-mimetics.

343

344 To facilitate translation to the clinic, we developed an ex vivo assay to investigate 345 chemosensitivity of fresh patient derived GBM tissue to BCL-2 targeting BH3-mimetics. 346 Across different freshly resected IDHwt GBM samples, we found that targeting both MCL-1 347 and BCL-xL led to an extensive induction of apoptosis and sustained reduction in tumour cell 348 viability ex vivo, without compromising tissue and vessel integrity. Following inhibition of 349 BCL-xL we found an increased amount of BIM bound to MCL-1, leading to a sensitisation of 350 the GBM cells to MCL-1 antagonists. This mirrors studies in haematologic malignancies 351 where the susceptibility to BH3-mimetics was dependent on BCL-2 complexed to BIM (49, 352 50) and increased BIM levels sensitised to BCL-2 inhibition (51).

353

To circumvent reported systemic toxicity, we developed a sequential treatment schedule. Our analysis of bioavailability in an orthotopic patient-derived xenograft revealed that the blood-brain-barrier is barely penetrable for ABT-263. Using the GBM cell line U87MG in a

357	subcutaneous xenog	raft model, the	e observed effect	of sequential	priming could also be	è
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- 358 recapitulated *in vivo* with a profound regression of tumour size and significant survival
- 359 benefit. Importantly, the brain, due to its blood-brain-barrier, provides unique opportunities
- 360 for drug delivery. For instance, local drug delivery or intrathecal chemotherapy can be
- 361 exerted to use the blood-brain-barrier and in turn circumvent systemic side effects (45). In
- 362 summary, these data provide a rationale for further investigating alternating inhibition of
- 363 BCL-xL and MCL-1 pro-survival function in GBM to maximise the therapeutic effect.
- 364

365 Materials and methods

- 366 Patient-derived GBM cell lines and cell culture reagents
- 367 Patient-derived GBM stem-like cells (E2, G1, G7, R9, R15, R24 GSC), obtained from
- 368 surgical resection specimens of anonymised patients as described (52, 53), were kindly
- 369 provided by Prof. Colin Watts. GSC and U87MG were cultured in serum-free Advanced
- 370 Dulbecco's modified Eagle's medium F12 (Thermo Fisher Scientific), supplemented with
- 2mM glutamine, 4µg/ml heparin (Sigma), 1% B27 (Thermo Fisher Scientific), 0.5% N2
- 372 (Thermo Fisher Scientific), 20ng/ml EGF and 10ng/ml FGF (Thermo Fisher Scientific). DIFF
- 373 cells were cultured in 10% foetal calf serum (FCS) containing high-glucose DMEM (Thermo
- 374 Fisher Scientific) complemented with 2mM glutamine. Human astrocytes were originally
- 375 provided by Prof. Steven Pollard as human foetal neural stem cells and previously
- 376 differentiated to astrocytes by 7 days culture in differentiation inducing medium as described
- before (54). All cells were kept in 37°C incubator at 5% CO₂ and, when grown as
- 378 monolayers on Matrigel (Corning) pre-coated plates or as spheres in uncoated plates. For all
- 379 experiments, cells were used up to ten passages after thawing. All cell lines used were
- 380 routinely tested for mycoplasma.
- 381 For our in vitro studies the following drugs and chemicals were used: ABT-199 (AdooQ
- 382 BioScience, #A12500-50), ABT-263 (ApexBio, #A3007), ABT-737 (ApexBio, #A8193),
- 383 A1331852 and A1155463 (ApexBio, #B6164 and #B6163), S63845 (Chemgood, #C-1370),
- 384 Actinomycin D (Calbiochem, #114666), q-VD-OPh (QVD, AdooQ BioScience, #A14915-25),
- 385 Sytox Green (Thermo Fisher Scientific, #S7020), Brain-derived neurotrophic factor (BDNF;
- 386 Peprotech, #450-02), 7,8-Dihydroxyflavone hydrate (7,8-DHF; Merck, #D5446).
- 387
- 388 Lentiviral transduction
- 389 GSC transduction was performed using CRISPR/Cas9 genome editing with the following
- 390 guide sequences:
- 391 hBAX: 5'-AGTAGAAAAGGGCGACAACC-3'
- 392 hBAK: 5'-GCCATGCTGGTAGACGTGTA-3'

393 hMCL-1.1: 5'-GGGTAGTGACCCGTCCGTAC-3'

394 hMCL-1.2: 5'-GTATCACAGACGTTCTCGTA-3'

395 hBIM: 5'- TACCCATTGCACTGAGATAG-3'

396 For stable cell line generation HEK293-FT cells (4x10⁶ in a 10 cm dish) were transfected

397 using 4µg polyethylenimine (PEI, Polysciences) per µg plasmid DNA with the

398 LentiCRISPRv2-puro (Addgene #52961) or LentiCRISPRv2-blasti (55) backbone, lentiviral

399 transfer vector plasmid, packaging plasmid (Addgene #14887) and envelope plasmid

400 pUVSVG (Addgene #8454), mixed in a 4:2:1 ratio. DNA/PEI mixtures were incubated at

room temperature for 10 to 15 minutes, prior to application on HEK293-FTs. 24 and 48

402 hours later, virus containing supernatant was harvested and filtered (0.45µM). Virus was

403 extracted using Lenti-X concentrator (Clontech Takara) according to the manufacturer's

404 instructions. The virus containing pellet was resuspended in serum-free stem-cell medium

and target cells were infected in the presence of $1\mu g/ml$ polybrene (Sigma Aldrich). Two

406 days following infection, cells were selected by growth in puromycin (E2: 1µg/ml, G7:

407 0.5μg/ml; Sigma Aldrich) or blasticidin (G7, R15 and R24: 10 μg/ml; InvivoGen) containing

408 medium. As described previously, plasmids encoding iRFP IRES puro have been inserted
409 into a pBABE vector (36).

410

411 Cell proliferation and live-cell viability assay

412 Cell death and cell confluence were determined using live-cell imaging in the IncuCyte Zoom

413 and S3 (Sartorius). For cell confluence 50x10³ cells were seeded in Matrigel-coated 6-well

414 plates. Cell area per well was measured using IncuCyte imaging analysis software

415 (Sartorius). For cell death assays, 6x10³ or 12x10³ GSC per well were seeded in Matrigel-

416 coated 96-well plates and treated with the indicated drugs in the presence of 30nM SYTOX

417 Green. Plates were applied to the IncuCyte imager at 37°C in a humidified 95% air/ 5% CO₂

418 incubator. Every hour, two images per well were taken over a period of 24 to 48 hours.

419 Images were presented in green phase contrast at 10x magnification. For image

420 quantification IncuCyte imaging analysis software was used. Percentage cell death was

421	calculated by normalising against maximal cell death control upon 24 to 48 hours treatment
422	(1µM Actinomycin D, 10µM ABT-737 and 1µM S63845). Alternatively, 100% cell death
423	control was verified by visual inspection of IncuCyte images, where 100% Sytox Green
424	positive cells = total cell count.
425	
426	Clonogenic survival assay
427	GSC were seeded at a density of 250 cells per well in Matrigel-coated, 6-well plates with
428	three technical repeats per experiment and left to adhere overnight. After 16 hours cells
429	were treated as indicated for 24 hours, followed by replacement of fresh media. Cells were
430	left to form colonies for 2 to 3 weeks prior to methanol fixation and crystal violet staining.
431	Visible colonies consisting of minimum 50 cells were counted manually.
432	
433	Neurosphere formation assay
434	G7 GSC vector ^{CRISPR} and MCL-1 ^{CRISPR} were seeded at a density of 10 cells per well in
435	uncoated 96-well plates. Serum-free stem-cell medium was refreshed every week. Spheres
436	were left to grow for 14 days before manual scoring of the 60 inner wells.
437	
438	Immunoblotting, immunoprecipitation and antibodies
439	GSC were lysed and collected in RIPA buffer (50mM Tris-HCl pH 7.5, 150 mM NaCl, 1 mM
440	EDTA, 1% NP-40), supplemented with complete protease inhibitor (Roche) and PhosSTOP
441	(Roche). Protein concentration was determined using Pierce BCA protein assay kit (Thermo
442	Fisher Scientific) and protein lysates were subjected to electrophoresis through SDS–PAGE
443	or 4-12% NuPage Bis-Tris protein gels (Thermo Fisher Scientific) followed by blotting onto
444	nitrocellulose membranes. After blocking in 5% non-fat, dry milk or 2% BSA (Roche),
445	membranes were probed with primary antibody (dilution 1:1000) BAK (Cell Signaling
446	#12105), BAX (Cell Signaling #2772), BCL-2 (Cell Signaling #2762), BCL-xL (Cell Signaling
447	#2762), MCL-1 (Cell Signaling #5453), BIM (Cell Signaling #2933), TrkB (Cell Signaling
448	#4603), pTrkA (Tyr674/675)/pTrkB (Tyr706/707) (Cell Signaling #4621), ERK1/2 (Cell

449 Signaling #4695), pERK1/2 (Cell Signaling #4370), AKT (Cell Signaling #9272), pAKT 450 (Ser473; Cell Signaling #4066), Caspase 3 (Cell Signaling #9662), cleaved Caspase 3 (Cell 451 Signaling #9664), PARP1 (Cell Signaling #9532) and SOX2 (Abcam #ab92494), NESTIN 452 (Abcam #ab22035), GFAP (Santa Cruz #SC-6170) at 4°C overnight in blocking buffer. α-453 tubulin (Sigma #T5168, 1:5000), β -tubulin (Cell Signaling #2146, 1:5000), HSP60 (Cell 454 Signaling #4870, 1:1000), or actin (Sigma #A4700, 1:5000) served as loading controls. Each 455 blot was probed with primary antibodies and a loading control. Representative loading 456 controls are shown in figures. Membranes were incubated with Li-Cor secondary antibodies 457 (IRDye 680RD donkey anti-mouse, IRDye 800CW donkey anti-rabbit, IRDye 800CW donkey 458 anti-goat) for 1 hour at room temperature. 459 For immunoprecipitation (IP), rabbit antibodies were coupled to magnetic beads conjugated 460 to anti-rabbit IgG (Dynabeads Sheep anti-rabbit IgG, Invitrogen, #11203D). The buffer 461 containing 200 mM NaCl, 75 mM Tris-HCl pH 7, 15 mM NaF, 1.5 mM Na3VO4, 7.5 mM 462 EDTA, 7.5 mM EGTA, 0.15% (v/v) Tween-20 and protein inhibitors (Thermo Fisher) were 463 used to prepare cell lysates. Lysates were passed several times through a 26-gauge needle 464 followed by centrifugation at 10,000g for 5 min at 4 °C. Lysates were added to the beads 465 and rotated for 2 hours at 4 °C. After washes in Tween-20-containing buffer, lysates were 466 analysed by immunoblotting. 467 Blots were imaged using Li-Cor Odyssey CLx (Li-Cor), acquired and processed using 468 Image-Studio software (Li-Cor) and subsequently arranged using Adobe Illustrator. 469

470 Orthotopic intracranial and subcutaneous xenografts

471 All mouse experiments were carried out in accordance with the Animals Act 1986 (Scientific

472 Procedures on living animals) and the regulatory guidelines of the EU Directive 2010 under

- 473 project licences PPL P4A277133 and PP6345023 and ethical review (University of
- 474 Glasgow). For intracranial xenograft 7-week old female CD1-nude mice (Charles River, UK)

475 were orthotopically injected with 1 x 10⁵ iRFP-labelled vector^{CRISPR} and MCL-1^{CRISPR} G7 GSC

into the right striatum. Mice were monitored for the duration of the experiment and humanely

477 sacrificed when they showed neurological (hemiparesis, paraplegia) or general symptoms

478 (hunched posture, reduced mobility, and/or weight loss >20%).

479 For subcutaneous xenograft 1 x 10⁶ U87MG cells, previously cultured in stem-cell medium,

480 were diluted in PBS and 50% growth factor reduced Matrigel and injected in the right flank of

481 8-week old female CD1-nude mice (Charles River, UK). For in vivo dosing, ABT-263

482 (ChemieTek #A263) was dissolved in 10% ethanol, 30% PEG glycol 400 and 60% Phosal

483 50 PG at 20 mg/kg and administered via oral gavage. S63845 (ChemieTek #S63845) was

484 prepared in 2% vitamin E/d-α-tocopheryl polyethylene glycol 1000 succinate (Sigma)

immediately prior to IV administration by tail vein injection at 25 mg/kg. Mice were treated

486 with a 48-hours pause between drug administrations over a 14 days period. Tumour growth

487 was monitored by caliper measurement three times per week and volume calculated using

488 the equation ([length x width²]/2). Clinical end point, at which mice were euthanised, was

489 15mm diameter or ulceration of the tumour.

490

491 Intravital cranial iRFP imaging and magnetic resonance imaging (MRI)

492 To examine intravital, intracranial tumour growth in animals bearing iRFP-positive G7 GSC,

493 mice were monitored by PEARL imaging (Li-Cor) as previously described (36). MRI scans

494 performed on brain tumour bearing mice using a nanoScan PET/MRI scanner (Mediso

495 Medical Imaging Systems, Hungary). Mice were maintained under inhaled isoflurane

496 anaesthesia (induction 5% v/v; maintenance 1.5 - 2.0% v/v) in the medical air during imaging

497 procedure duration. Whole brain T2 Fast Spin Echo (FSE) 3D Axial Sequences (slice

thickness 1.0 mm, repetition time (TR) 2000 msec, echo time (TE) 83.7 msec, Flip Angle 90

499 degrees) were used to acquire MRI scans. For assessments of scans, volume-of-interest

500 (VOI) was manually drawn around the tumour region on MRI scans by visual inspection.

501 Separate VOI were drawn for each scan to adjust for the position of the mice on the scanner

502 and tumour size.

503 Patient-derived GBM specimens and tissue culture

GBM specimens were obtained from surplus tumour tissue resected from patients treated
within the OPARATIC study (NCT01390571). Patients had consented for use of surplus
tissue for future research projects.

Fresh GBM tissues were obtained from surplus surgical resection tissue from patients at the 507 508 Queen Elizabeth University Hospital (QEUH) in Glasgow after review by neuropathology 509 with appropriate consent and in accordance with the NHS GG&C ethical committee review 510 (Biorepository Application No. 432). The patient study was conducted in accordance with the 511 Declaration of Helsinki. Neuropathological diagnosis and selected patient information are 512 displayed in the figures. Further, details of these patients are restricted by institutional 513 requirements. All experiments were performed conform to relevant regulatory standards of 514 the CRUK Beatson Institute. Fresh samples were attenuated in 2% low gelling temperature 515 agarose (Merck) and cut into 350µm thick slices using the McIlwain tissue slicer (Campden 516 Instruments). Tissue slices were dissected under the microscope in ice cold PBS before they 517 were transferred on top of hydrophilic Millicell cell culture inserts (Merck Millipore) into 518 serum-free Advanced Dulbecco's modified Eagle's medium F12 supplemented with 0.5% 519 N2, 1% B27, 1% glutamine and 1% penicillin-streptomycin and left to equilibrate for 24 hours 520 at 37°C in a humidified 95% air/ 5% CO₂ incubator, before treatment with indicated drugs for 521 72 hours. Following PBS washes brains were fixed in 4% paraformaldehyde (PFA) 522 overnight.

523

524 Organotypic adult mouse brain slice culture

525 Extracted brains from three 11-week old C57BL/6J mice were transferred to sterile PBS on

526 ice, divided into both hemispheres and cut into coronal, 100µm thick slices using a

527 vibratome (Campden Instruments 5100mz, advance speed 1mm/sec, oscillation amplitude

528 1.5mm, 80Hz). Up to 5 slices per hemisphere were cut around the subventricular zone

529 (SVZ). Slices were cultured on top of cell culture inserts in neurobasal medium as described

530 in the previous section and left to equilibrate for 1 hour at 37°C in a humidified 95% air/ 5%

531 CO₂ incubator before treatment with indicated drugs for 24 hours. After PBS washes slices

532 were fixed in 4% PFA overnight.

533

534 Immunohistochemistry (IHC) and immunofluorescence (IF)

535 H&E staining and IHC was performed on $4\mu m$ formalin fixed paraffin embedded (FFPE) 536 sections. For BCL-xL (Cell Signalling #2764), cleaved Caspase 3 (Cell Signalling #9661) 537 and MCL-1 (Abcam # ab32087) IHC staining the Leica Bond Rx Autostainer was used. All 538 FFPE sections underwent on-board antigen retrieval for 20 minutes using ER2 retrieval 539 buffer (Leica, UK) before staining at a previously optimised dilution (BCL-xL 1:500; cleaved 540 Caspase 3 1:500; MCL-1 1:200) and visualised with Liquid DAB (Agilent, UK). Ki67 (Agilent 541 #M7240) staining was performed on a Dako Autostainer Link 48 using high pH TRS retrieval 542 buffer performed in a PT module (20 mins at 97°C). Ki67 was applied at 1:100 dilution 543 before visualising using Liquid DAB. For SOX2, IHC epitope retrieval was achieved by 544 heating to 98°C in pH6 citrate buffer for 60 minutes before proceeding as per the 545 manufacturer's instructions with SOX2 antibody used at a dilution of 1:500. Scanning and 546 image analysis was conducted using Halo (Indica Labs). Algorithms were optimised for each 547 stain individually and automated, quantitative analysis undertaken with Halo software (Indica 548 Labs). 549 For IF staining tissue slices were permeabilised and blocked in PBS with 10% NGS, 1% 550 BSA, 0.3% TX-100 and 0.05% Azide for 1 hour at room temperature. After washes with 10% 551 NGS, 1% BSA, 0.1% TX-100 and 0.05% Azide containing buffer slices were incubated with 552 primary antibodies (NESTIN 1:300, GFAP 1:400, cl. Caspase 3 1:400) in washing buffer for 553 72 hours at 4°C. After washes slices were incubated in secondary antibodies (1:200, Alexa 554 Fluor 568 goat anti-mouse (#A11004), Alexa Fluor 488 goat anti-rabbit (#A11034), Life 555 Technologies) in washing buffer for 24 hours. Following washes in PBS tissues were 556 counterstained with DAPI (VECTASHIELD, LSBio) and mounted in gaskets (BioRad Seal 557 Frame Incubation Chambers) on glass cover slips. Images were acquired using a Zeiss 710

558 laser scanning microscope with an EC Plan-Neofluar 40x/1.30 Oil DIC M27 objective and

559 Zen 2.3 SP1 FP3 (black edition) software. 70µm Z-stacks were acquired at 2.5µm intervals

- and Maximum Intensity Projections (MIPs) were generated using Zen 2.1 (blue edition).
- 561 Image processing was performed using Fiji (ImageJ 1.53c). Cleaved Caspase 3 positive
- 562 cells were counted manually and nuclei were counted automatically using CellProfiler

563 (Version 4.0.7).

- 564
- 565 In silico and transcriptomic analysis
- 566 REMBRANDT microarray data was obtained from gliovis.bioinfo.cnio.es. Data was filtered 567 for histology and tumour grading.
- 568 RNA sequencing data was obtained from a previously published GBM database (30). In
- order to determine the most differentially expressed genes, calculation of expression rank
- 570 product was employed to assess relative gene expression in paired GSC and DIFF cell lines
- 571 (56). Only results > 10 reads were incorporated.
- 572
- 573 Statistical analyses
- 574 For comparison of two experimental groups two tailed, unpaired *t* test with Welch's
- 575 correction (Welch's test) or Mann-Whitney test were used. For tumour related Kaplan–Meier
- 576 survival curves Mantel-Cox (Log-rank) was plotted. All statistical analyses were executed
- 577 with Prism software version 9 (GraphPad, La Jolla, CA, USA).
- 578
- 579

580 Acknowledgements

581 First and foremost, our special gratitude goes to the GBM patients who agreed to make their

tumour specimens available for research. Special thanks to Mary Fraser, Dr. Alexandru

- 583 Stan, Dr. Zoltan Hanzely and the Neuropathology Department of the QUEH in Glasgow, who
- obtained patient consent and provided support with GBM tissue sampling. Further, we would
- 585 like to thank the Core Services at the Cancer Research UK Beatson Institute with particular
- thanks to Gemma Thomson, David Strachan, the Biological Services Unit and Histology. We

587 thank Prof. Colin Watts for generous sharing of the patient-derived GBM cell lines, Prof. 588 Steven Pollard and Rodrigo Gutierrez Quintana for providing the human astrocytes. Many 589 thanks to Dr. Jayanthi Anand and Dr. Dimitris Athineos for their invaluable support with the 590 animal work. Many thanks to members of the Tait laboratory for discussion and input, and 591 Catherine Winchester for critical reading and assistance in the preparation of this 592 manuscript. Figures were created with BioRender.com. Funding: This work was supported 593 by CRUK core funding to the Beatson Institute (A17196) to C.N., (A17196/A31287) to 594 N.R.P., (A22903) to K.M.R. and J.O'P., and (A18277) to J.C.N., a CRUK Programme 595 Foundation Award (C40872/A20145) to S.W.T., CRUK Clinical Research Fellowship 596 (A23220) to A.L.K., funding by the University of Glasgow to D.K. and A.J.C. and funding by 597 the Beatson Cancer Charity and Cancer Research UK RadNet Centre Glasgow (A28803) to 598 K.S., L.M.-E. was funded by the by Erasmus+ Program and Short Research stay fellowship 599 for trainees by Universitat Autònoma de Barcelona.

600

601 Author Contributions

602 A.L.K. and S.W.G.T. conceived the study, designed the work plan and wrote the manuscript.

603 A.L.K. performed the majority of the experiments. D.K., L.M.-E., J.O'P. and F.J.B. acquired

data and provided technical support. N.R.P. performed IF imaging and analysis. D.K., K.S.,

605 C.C., C.N. and G.M. provided assistance with the *in vivo* models and related imaging. K.K.

and W.S. supervised and supported patient specimen collection. A.J.C. provided the

607 RNAseq dataset. Y.E., D.K. and M.R.J. performed and supported data analysis. Expertise

and critical input as well as review of the manuscript was given by K.J.C., G.I., K.B., A.J.C.,

609 K.M.R. and J.C.N.

610

611 Competing interests

K.J.C. is a recipient of a share in royalty payments paid to the Walter and Eliza Hall Instituteof Medical Research in relation to venetoclax.

614

615

616 References

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774

775 Figure legends

776 Figure 1. High anti-apoptotic BCL-2 family protein expression correlates with

777 increased BH3-mimetic sensitivity in GBM

(A) Schematic overview of BH3-mimetic drugs used and their respective targets (B) Panel of
 six GSC cell lines and the human primary GBM cell line U87MG were treated with indicated

780 drugs for 24 to 48 hours and analysed for cell viability using an IncuCyte imager and SYTOX

- 781 Green exclusion. Results are presented as heatmap. Percentage cell death was calculated
- by normalising against maximal cell death (treatment with 1µM Actinomycin D, 10µM ABT-
- 783 737 and 1µM S6384), n=3 independent experiments per cell line. (C) Immunoblot of BCL-2
- family proteins, cell-line specific neural stem cell marker SOX2 and astrocyte lineage
- differentiation marker GFAP in human astrocytes and patient-derived GSC. α-tubulin served
- as loading control. Representative image from n=3 independent experiments. (D,E) Matched
- tumour and margin specimens were obtained from three patients diagnosed with GBM and
- stained for haematoxylin and eosin (H&E), BCL-xL and MCL-1 IHC (representative images
- of one case shown). H-score (analysis of intracellular BCL-xL and MCL-1 expression) was
- 790 determined using automated analysis with Halo. Error bars represent mean +/-SD
- 791 (***p=0.0002, *p=0.0404) Welch's test. Further representative images in lower

magnifications are shown in Supplementary Figure 1C.

793

Figure 2. Anti-apoptotic MCL-1 is required for the growth and survival of GBM

- 795 (A) Representative images of neurosphere growth from G7 GSC vector^{CRISPR} (upper panel),
- 796 MCL-1.1^{CRISPR} (middle panel) MCL-1.2^{CRISPR} (lower panel), respectively. Quantification of

797 neurosphere formation capacity by G7 GSC vector^{CRISPR} vs. MCL-1.1^{CRISPR} or MCL-1.2^{CRISPR}.

- Firor bars represent mean +/-SD from n=4 independent experiments (p=0.8146, ns,
- 799 nonsignificant) Welch's test. (B) Representative images of brain MRI scans (tumour
- 800 indicated by red dashed line) next to corresponding pseudocolour representations of iRFP
- signal of mice bearing iRFP tagged G7 GSC vector^{CRISPR} (upper panel) and MCL-1.2^{CRISPR}
- 802 (lower panel) xenografts, respectively. iRFP signal was detected by PEARL scans (700nm

803 channel) (1) at week 8 and (2) at week 20 (vector^{CRISPR}) or week 36 (MCL1^{CRISPR}) post

- injection. (**C**) Quantification of time to 20% iRFP signal increase of G7 vector^{CRISPR} n=13 vs.
- 805 G7 MCL-1^{CRISPR} tumours n=19, compared to 4 weeks post injection (baseline signal). Error
- bars represent mean +/-SD (**p=0.0013) Mann-Whitney test. (D) Kaplan-Meier survival
- graph of mice with orthotopic xenografts of G7 GSC iRFP vector^{CRISPR} n=9 (median survival
- 808 87 days) vs. MCL-1^{CRISPR} tumours n=10 (median survival undefined) post tumour cell
- 809 implantation (p<0.0001) Log-rank (Mantel-Cox) test.
- 810

811 Figure 3. GSC display increased apoptotic priming and can be effectively killed by

812 dual BCL-xL, MCL-1 inhibition

- 813 (A) G7 and E2 GSC were treated with DMSO (ctrl), A-1331852 or ABT-737 as indicated for
- 814 16 or 24 hours, respectively, harvested and protein expression was analysed by
- 815 immunoblot. α -tubulin served as loading control. Representative image from n=3
- 816 independent experiments shown. (B) G7 or R24 GSC vector^{CRISPR} vs. MCL1.1^{CRISPR} and
- 817 MCL1.2^{CRISPR} were treated with A-1331852 for 24 hours and analysed for cell viability using
- 818 an IncuCyte imager and SYTOX Green exclusion. Percentage cell death was calculated by
- 819 normalising against maximal cell death verified by visual inspection. Error bars represent
- 820 mean +/-SEM from n=3 independent experiments. (G7: *p=0.0218, **p=0.002, ***p=0.0006)
- 821 (R24: **p=0.0028, ****p<0.0001) Welch's test. (C) E2, G1, G7, R24 GSC were treated with a
- 822 combination of A-1331852 and S63845 in indicated concentrations for 24 hours and
- analysed for cell viability using an IncuCyte imager and SYTOX Green exclusion.
- 824 Percentage cell death was calculated by normalising against maximal cell death as
- 825 described in Figure 1B. Error bars represent mean +/-SEM from n=3 independent
- 826 experiments. (E2 **p=0.0021, G1 **p=0.0022, G7 **p=0.0016, R24 *p=0.0123) Welch's test.
- 827 (D) Clonogenic survival assay of G7 GSC iRFP treated with indicated drugs 16 hours after
- 828 plating 250 cells per well. Colonies counted manually after 14 days. Error bars represent
- 829 mean +/-SD from n=3 independent experiments (**p=0.0038) Welch's test. Representative
- 830 images of a replicate in one independent repeat scanned on LICOR imager. (E) E2 GSC

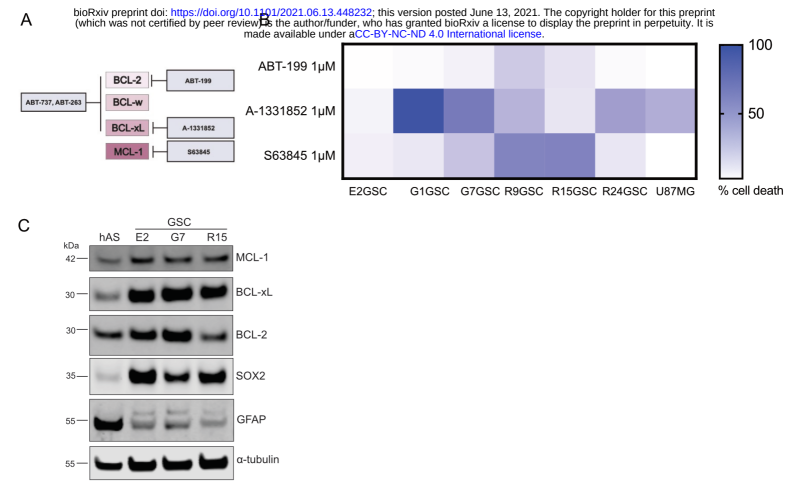
831 vector^{CRISPR} and BAK/BAX^{CRISPR} were treated as indicated for 2 hours, harvested and protein 832 expression was analysed by immunoblot. HSP60 served as loading control. Representative 833 image from n=3 independent experiments. (F) E2 and R15 GSC and paired DIFF cells were 834 treated either with DMSO (grey) or a combination of A-1331852 and S63845 (both 0.1 μ M) 835 for 24 hours and analysed for cell viability using an IncuCyte imager and SYTOX Green 836 exclusion. Error bars represent mean +/-SEM from n=3 independent experiments. 837 Representative IncuCyte images 24 hours after treatment are shown. 838 839 Figure 4. TrkB signalling regulates sensitivity of GSC to anti-apoptotic treatment 840 (A) Most differentially expressed genes in RNAseg analysis of E2 and G7 GSC vs. E2 and 841 G7 DIFF. NTRK2 codes for TrkB. (B) Immunoblot of TrkB in E2 and G7 GSC compared with 842 paired DIFF cells. α -tubulin served as loading control. Representative image from n=2 843 independent experiments. (C) E2 and G7 and GSC treated with BDNF (100ng/mL) or 7,8-844 DHF (20µg/mL) +/- A-1331852 1µM and S63845 1µM for 24 hours (after a 24-hour 845 starvation period in 1% glutamine containing DMEM/F12) and analysed for cell viability using 846 an IncuCyte imager and SYTOX Green exclusion. Error bars represent mean +/-SEM from 847 one of n=3 independent experiments (E2 *p=0.0166, G7 ***p=0.008) Welch's test. (D) G7 848 DIFF treated as described in Figure 4C and analysed for cell viability using an IncuCyte 849 imager and SYTOX Green exclusion. Error bars represent mean +/-SEM from n=3 850 independent experiments (ns, p=0.9284). (E) G7 GSC were treated with 7,8-DHF (20µg/mL) 851 +/- A-1331852 1µM and S63845 1µM for 1 hour, harvested and protein expression was 852 analysed by immunoblot. α -tubulin served as loading control. Representative image from 853 n=3 independent experiments. 854 855 Figure 5. Combined BCL-xL and MCL-1 inhibition causes apoptosis in human GBM ex 856 vivo

857 (A-C) Representative images of H&E, Ki67 and cleaved Caspase 3 IHC of three patients 858 diagnosed with IDHwt GBM (case 1-3; ctrl=DMSO, dual treatment=A-1331852 1µM + 859 S63845 1µM) (D-F) Quantification of cellularity, percentage Ki67 and SOX2 positive 860 cells/total cell count in all three cases treated with the drugs described in (A-C) for 72 hours. 861 Error bars represent mean +/-SEM (**p=0.0021) Welch's test. (G) Representative images of 862 IF staining of GFAP, NESTIN (both red) and cleaved Caspase 3 (green) in the 863 subventricular zone (SVZ) of mouse brain slices, cultured and treated with DMSO or a 864 combination of ABT-263 5µM and S63845 2µM for 24 hours, counterstaining with DAPI 865 (blue). (H) Quantification of cleaved Caspase 3 positive cells normalised to nuclear count in 866 single mice. Single dots represent analysed images. Error bars represent median (mouse 1: 867 p=0.9198, p=0.3086; mouse 2: p=0.149, p=0.2974; mouse 3: **p=0.0044, p=0.0593; ns, 868 nonsignificant) Welch's test. Scale bars = 50µm. 869 870 Figure 6. Priming with BCL-xL inhibition renders GSC vulnerable to MCL-1 inhibition, 871 promoting tumour regression in vivo (A) E2 GSC vector^{CRISPR} and BAK/BAX^{CRISPR} were treated with DMSO (-), A-1331852 and 872 873 S63845 as indicated for 24 hours, harvested and protein expression was analysed by 874 immunoblot. Representative image from n=2 independent experiments. β -tubulin served as 875 loading control. (B) Direct binding interactions between MCL-1 and BIM were 876 immunoprecipitated and interacting proteins were detected by western blot in G7 GSC 877 treated for 16 hours with A-1331852 (Input=total cell lysate, IP=immunoprecipitated fraction). 878 Representative image from n=2 independent experiments. (C) G7 GSC were treated with 879 DMSO, A-1331852 or S63852 1µM for 48 hours, followed by 24 hours drug washout with 880 exchange to fresh medium and treatment with indicated drugs for 24 hours. For cell viability 881 analysis, IncuCyte imager and SYTOX Green exclusion was used. Error bars represent 882 mean +/-SEM from n=3 independent experiments, (**p=0.003) Welch's test. (D) E2 GSC

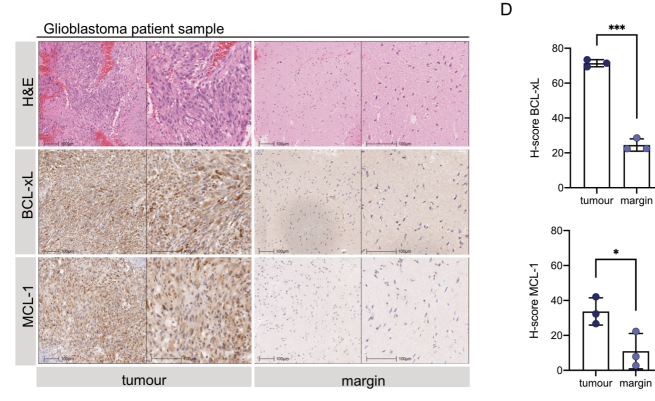
883 were pre-treated with DMSO or A-1331852 1µM (#) for 24 hours, followed by 24 hours drug

884 washout with exchange to fresh medium and treatment with indicated drugs for 48 hours. 885 For cell viability analysis, IncuCyte imager and SYTOX Green exclusion was used. Error 886 bars represent mean +/-SEM from n=3 independent experiments, (***p=0.0004) Welch's test. (E) G7 GSC vector^{CRISPR} and BIM^{CRISPR} were treated for 48 hours with A-1331852 1µM, 887 888 followed by 24 hours drug washout with exchange to fresh medium and treatment with 889 S63845 1µM for 24 hours. For cell viability analysis, IncuCyte imager and SYTOX Green 890 exclusion was used. Error bars represent mean +/-SEM from n=3 independent experiments, 891 (*p=0.0132) Welch's test. (F) Percent U87MG subcutaneous tumour volume change at the 892 end of 2 weeks alternating treatment with ABT-263, followed by S63845, relative to tumour 893 size at start. Treatment commenced when tumours were >5mm diameter. n=6 vehicle 894 treated (grey dots) and n=7 drug treated (blue dots). Bars are mean +/- SD (*p=0.0104) 895 Welch's test. (G) Kaplan-Meier survival analysis of U87 vehicle treated (grey line, n=6, 896 median survival 22 days) vs. U87 drug treated (blue line, n=7, median survival 34 days) 897 since treatment start (**p=0.0056) Log-rank (Mantel-Cox) test.

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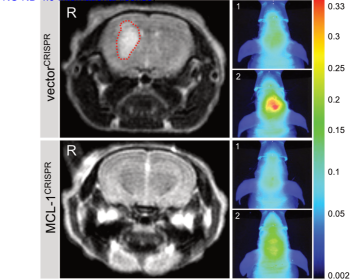


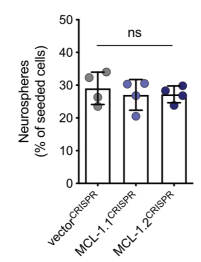
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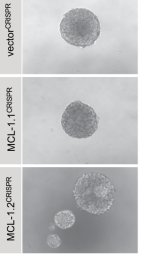


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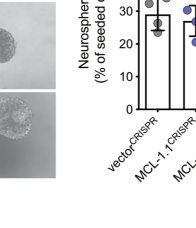


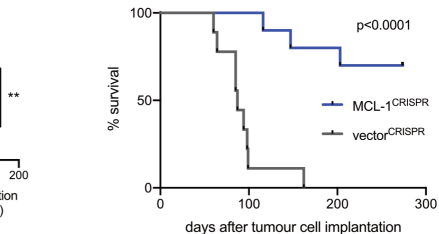


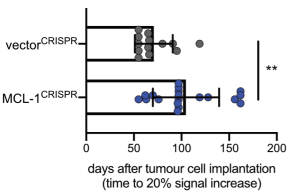


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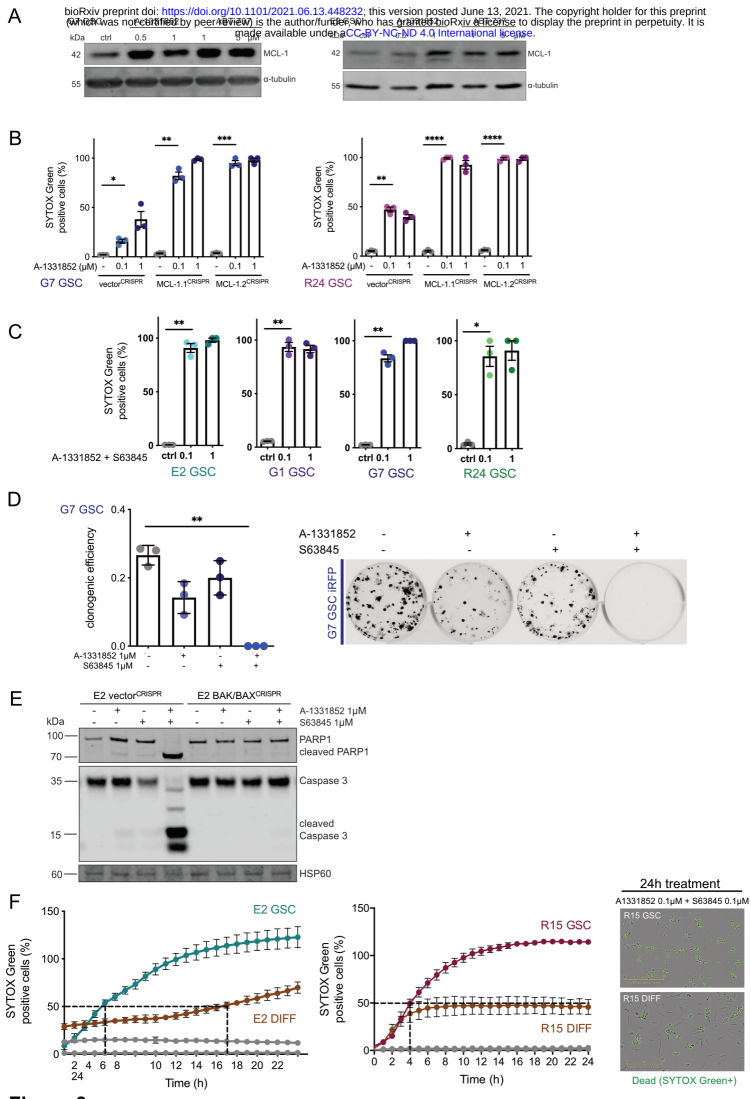


Figure 3

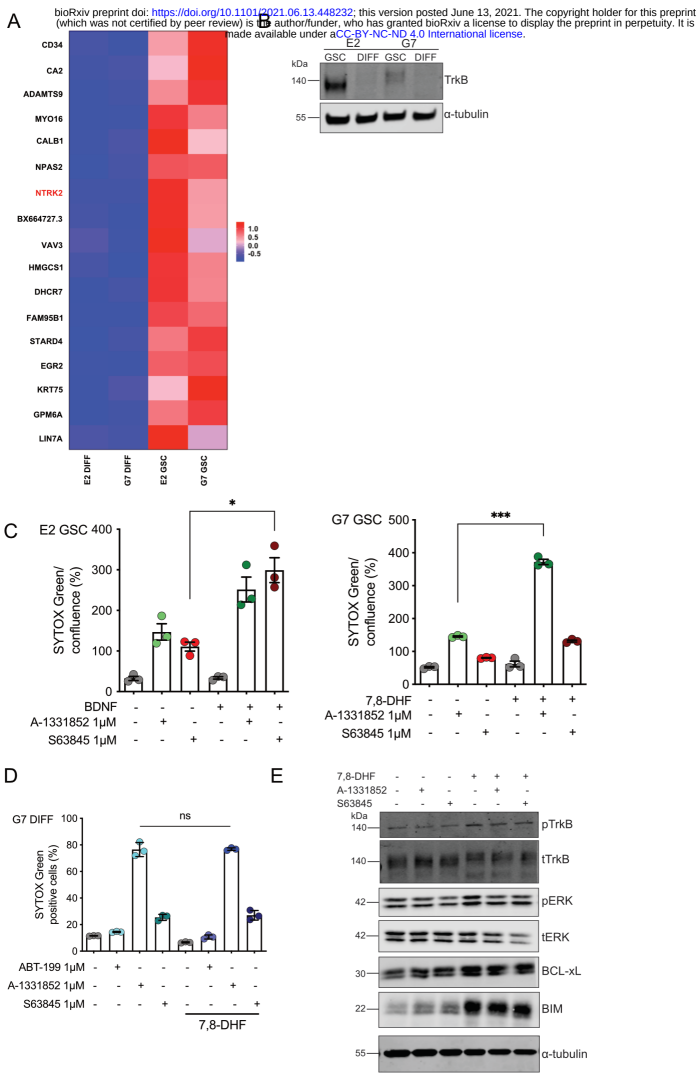
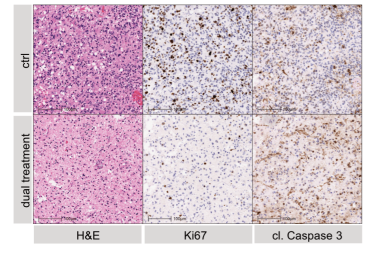
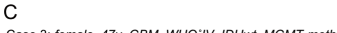
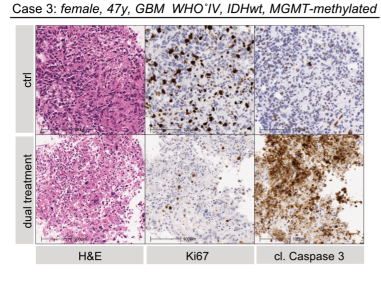


Figure 4

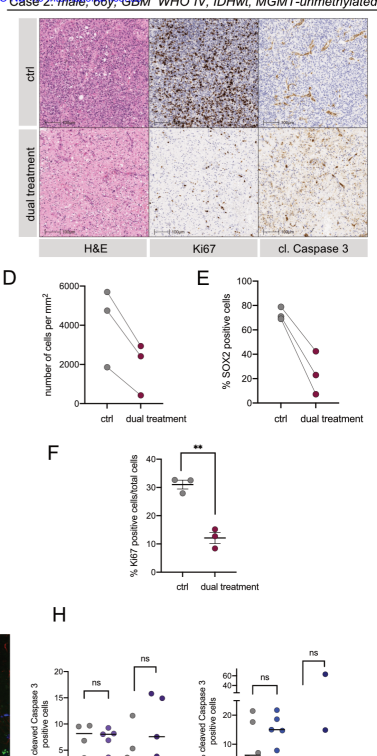
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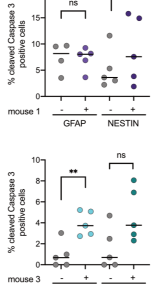






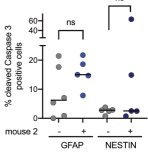
G DMSO + ABT-263 and S63845 cleaved Caspase 3 **GFAP** Nuclei cleaved Caspase 3 **Nestin Nuclei**

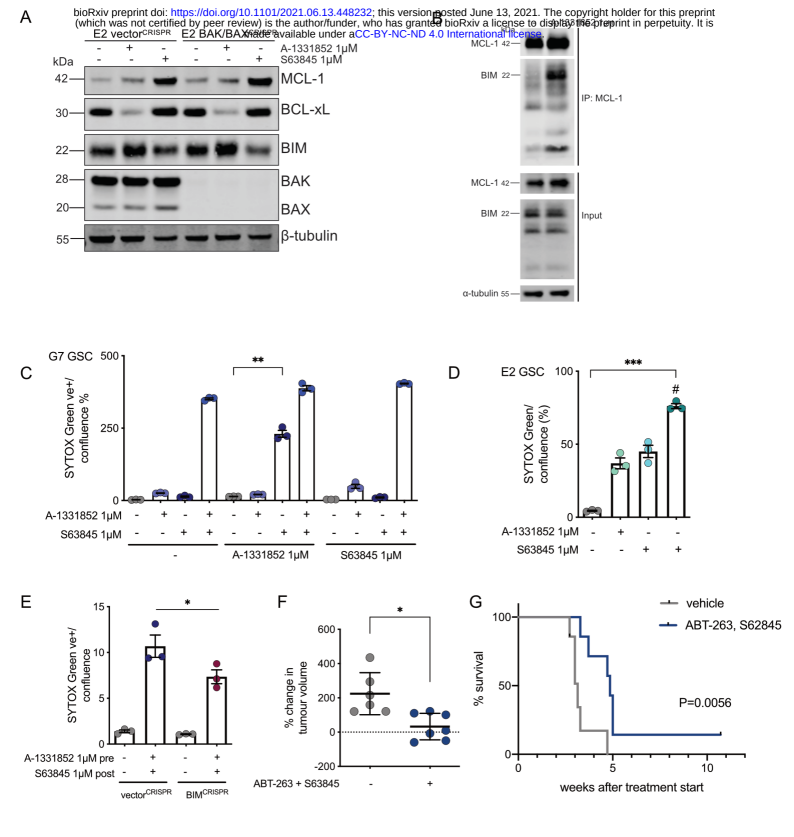


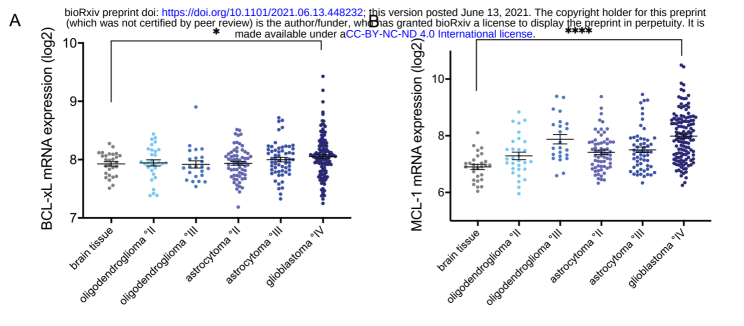


GFAP

NESTIN

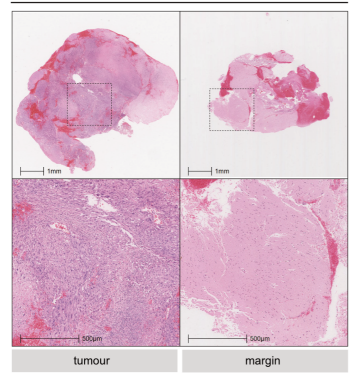




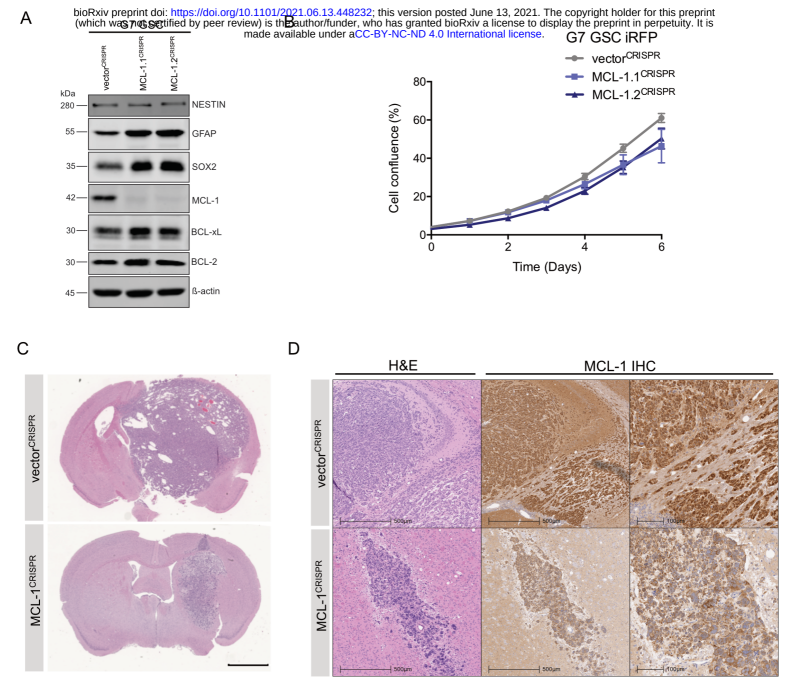


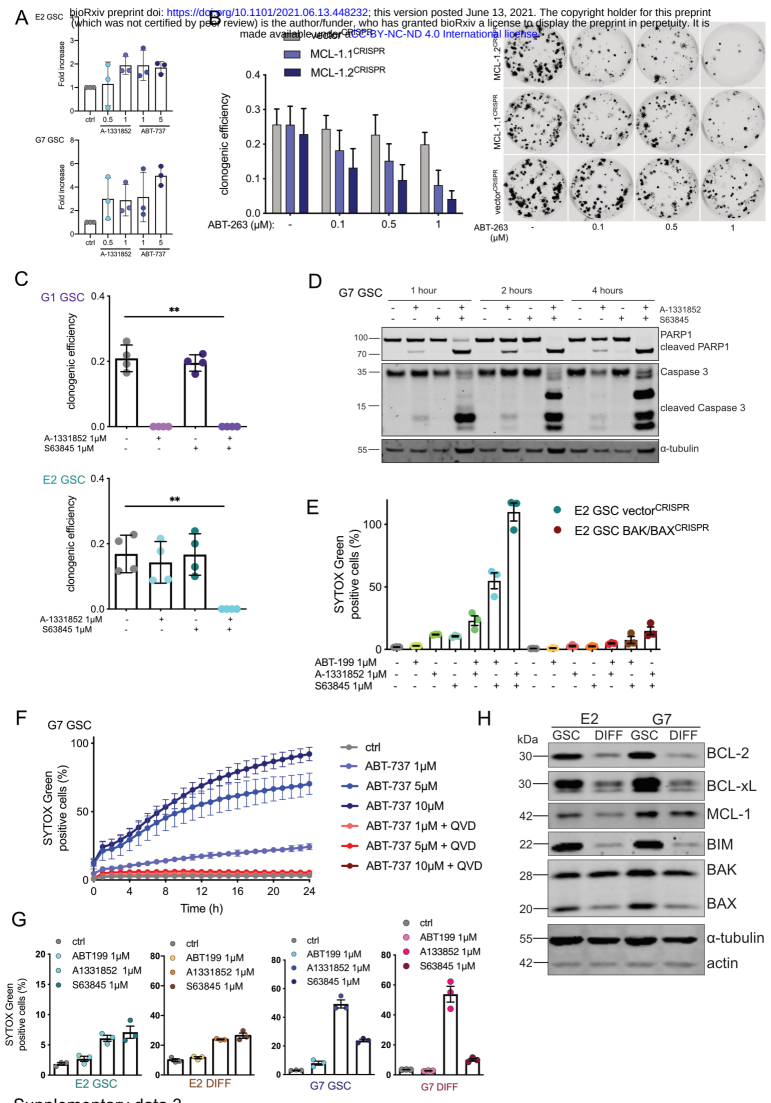
Glioblastoma patient sample

D

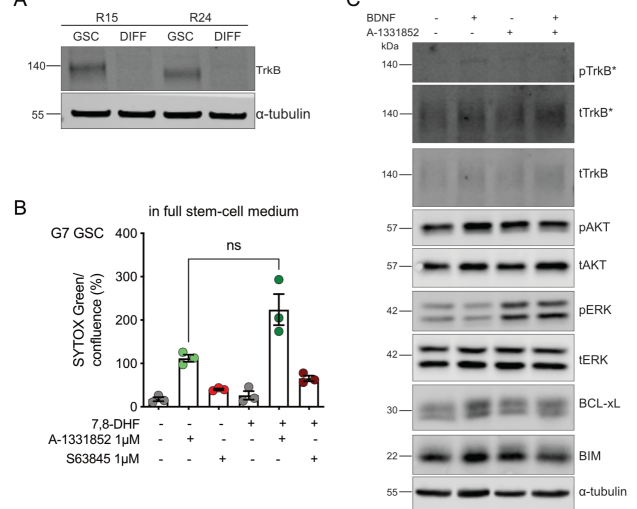


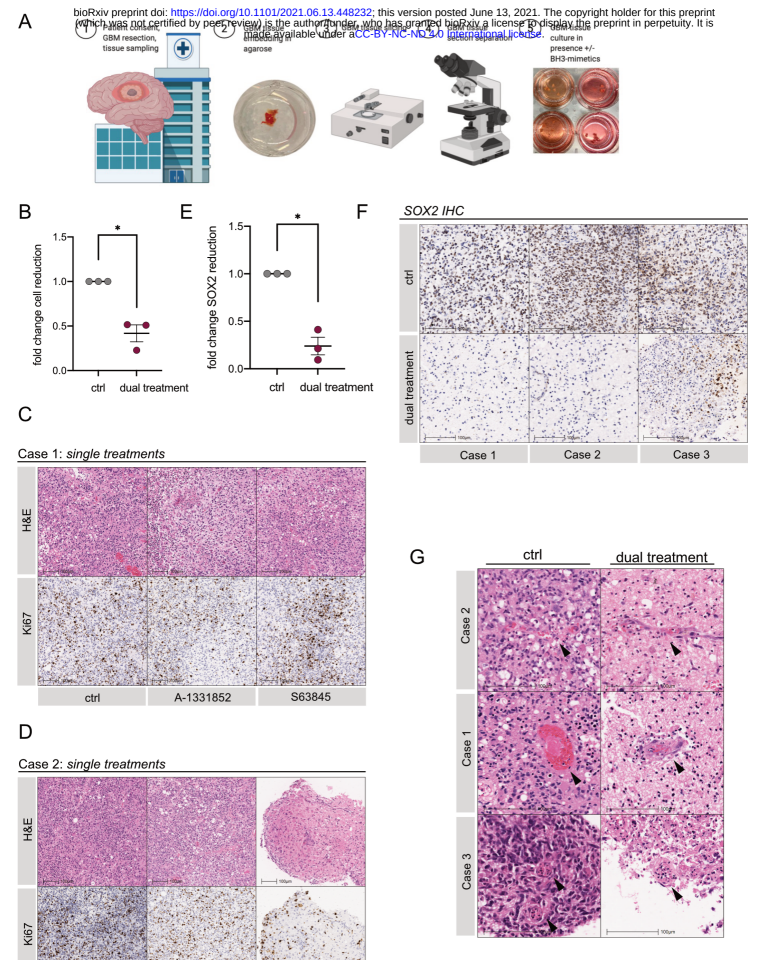
Supplementary data 1





Supplementary data 3





Supplementary data 5

ctrl

S63845

A-1331852

1 Supplementary Figure legends

2 Supplementary Figure 1. Relevant to Figure 1.

3 (A,B) BCL-xL and MCL-1 mRNA expression from the publicly available REMBRANDT GBM

4 microarray dataset. Data plotted for different glioma subtypes and normal brain tissue. Error

5 bars represent mean +/-SEM (*p=0.0118, ****p<0.0001) Welch's test. (C) Corresponding

6 representative H&E images of GBM tumour and margin samples shown in lower

- 7 magnification than in Figure 1D.
- 8

9 Supplementary Figure 2. Relevant to Figure 2.

(A) Immunoblotting of G7 GSC vector^{CRISPR}, MCL1.1^{CRISPR} and MCL1.2^{CRISPR} for BCL-2 10 11 family proteins, cell-line specific neural stem cell marker (SOX2, NESTIN) and astrocyte 12 lineage differentiation marker GFAP. Actin served as loading control. (B) Proliferation assay 13 of indicated cell lines using IncuCyte Imager and – percentage cell density over 6 days. 14 Error bars represent mean +/-SD from n=3 independent experiments. (C) Representative 15 H&E images of G7 GSC iRFP vector^{CRISPR} and MCL1^{CRISPR} orthotopic xenografts at end 16 point (corresponding to Figure 2B). Scale bar = 100µm. (D) Representative images of H&E and MCL-1 IHC of G7 GSC iRFP vector^{CRISPR} and MCL1^{CRISPR} orthotopic xenografts at end 17 18 point.

19

20 Supplementary Figure 3. Relevant to Figure 3.

21 (A) Quantification of immunoblots shown in Figure 3A. (B) Clonogenic survival assay of G7 GSC iRFP vector^{CRISPR} vs. MCL1.1^{CRISPR} and MCL1.2^{CRISPR} treated with indicated drugs 16 22 23 hours after plating 250 cells per well. Colonies counted manually after 14 days. Error bars 24 represent mean +/-SD from n=4 independent experiments. Representative images scanned 25 on LICOR imager. (C) Clonogenic survival assay of E2 and G1 GSC iRFP treated with 26 indicated drugs 16 hours after plating 250 cells per well. Colonies counted manually after 14 27 days. Error bars represent mean +/-SEM from n=4 independent experiments (E2**p=0.0099, 28 G1**p=0.002) Welch's test. (D) G7 GSC were treated with DMSO (-), A-1331852 and

29 S63845 for indicated times, harvested and protein expression was analysed by immunoblot. 30 α -tubulin served as loading control. Representative image from three independent experiments. (E) E2 GSC vector^{CRISPR} and BAK/BAX^{CRISPR} treated with indicated drugs for 48 31 32 hours and analysed for cell viability using an IncuCyte imager and SYTOX Green exclusion. 33 Percentage cell death was calculated by normalising against maximal cell death as 34 described in Figure 1B. Error bars represent mean +/-SEM from n=3 independent 35 experiments. (F) G7 GSC treated with indicated drugs (+/-QVD 10µM) for 24 hours and 36 analysed for cell viability using an IncuCyte imager and SYTOX Green exclusion. 37 Percentage cell death was calculated by normalising against maximal cell death as 38 described in Figure 1B. Error bars represent mean +/-SEM from n=3 independent 39 experiments. (G) E2 and G7 GSC with paired DIFF cells treated with indicated drugs for 24 40 hours and analysed for cell viability using an IncuCyte imager and SYTOX Green exclusion. 41 Percentage cell death was calculated by normalising against maximal cell death as 42 described in Figure 1B. Error bars represent mean +/-SEM from n=3 independent 43 experiments. (H) Immunoblot of E2 and G7 GSC with paired DIFF cells for BCL-2 family 44 proteins. α -tubulin served as loading control. Representative image from n=2 independent 45 experiments.

46

47 Supplementary Figure 4. Relevant to Figure 4.

(A) Immunoblot of TrkB in R15 and R24 GSC compared with paired DIFF cells. α-tubulin
served as loading control. Representative image from n=2 independent experiments. (B) G7
GSC treated with indicated drugs for 24h hours in full stem-cell medium (including EGF and
FGF) and analysed for cell viability using an IncuCyte imager and SYTOX Green exclusion.
Error bars represent mean +/-SEM from n=3 independent experiments. (ns, p=0.0817)
Welch's test. (C) G7 DIFF were treated with A-1331852 2µM +/- BDNF (100ng/mL) for 1
hour, harvested and protein expression was analysed by immunoblot (*indicates high

55 exposure). α-tubulin served as loading control. Representative image from n=3 independent

56 experiments.

57

58 Supplementary Figure 5. Relevant to Figure 5.

59 (A) Schematic model of fresh GBM patient tissue collection and processing. (B)

60 Quantification of cell reduction in GBM case 1-3 normalised to DMSO-treated control. Error

61 bars represent mean +/-SEM (*p=0.0257) Welch's test. (C,D) Representative H&E and Ki67

- 62 IHC images of GBM patients case 1 and 2 treated with single inhibitors A-1331852 2μM or
- 63 S63845 2µM for 72 hours. In both cases H&E control images were used from the same area
- 64 as shown in Figure 5A,B in different magnification. (E) Quantification of SOX2 positive cell
- reduction in GBM case 1-3 normalised to DMSO-treated control. Error bars represent mean
- 66 +/-SEM (*p=0.0145) Welch's test. (**F**) Representative images of SOX2 IHC in GBM case 1-3.
- 67 (G) Representative H&E images of GBM case 1-3, arrows indicate intratumoural vessels.
- 68

69 Supplementary Figure 6. Relevant to Figure 6.

(A) Immunoblot of E2 and G7 GSC vector^{CRISPR} and BIM^{CRISPR} for BCL-xL, MCL-1 and BIM. 70 71 Representative images from n=2 independent experiments. α -tubulin served as loading 72 control. (**B**) G7 GSC vector^{CRISPR} and BIM^{CRISPR} treated with DMSO or indicated combination 73 for 24 hours and analysed for cell viability using an IncuCyte imager and SYTOX Green 74 exclusion. Error bars represent mean +/-SD from n=3 independent experiments. (C) 75 Representative images of U87MG neurospheres in stem-cell medium under treatment with 76 indicated drugs over a total of 60 hours. Quantification of neurosphere treatment for cell 77 viability using an IncuCyte imager and SYTOX Green exclusion given as mean signal 78 intensity (GCU). One representative of n=3 independent experiments shown. Error bars 79 represent mean +/-SD. (D) Clonogenic survival assay of U87MG. Treatment commenced 16 80 hours after plating 250 cells/well with either DMSO or ABT-263 5 μ M for 24 hours followed by 81 drug washout and treatment pause for 24 hours and S63845 2µM for 24 hours. Alternating

- treatment was continued over the experimental period of 14 days. Colonies were counted
- 83 manually. Quantification of one representative of n=2 independent experiments shown.
- 84 Representative images of a replicate in one independent repeat scanned on LICOR imager.
- 85 (E) Schematic of treatment schedule for in vivo study. (F) Percent weight change of mice
- 86 (vehicle n=6, ABT-236, S63845 n=7) during the drug treatment period of the experiment in
- 87 Figure 1G,H. (G) Representative H&E and Ki67 IHC images of U87MG xenografts treated
- 88 with vehicle or alternating ABT-263 and S63845 therapy at end point.
- 89