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Mitochondrial dynamics regulate genome stability via control of caspase-dependent DNA damage

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Summary

Mitochondrial dysfunction is interconnected with cancer. Nevertheless, how defective mitochondria promote cancer is poorly understood. We find that mitochondrial dysfunction promotes DNA damage under conditions of increased apoptotic priming. Underlying this process, we reveal a key role for mitochondrial dynamics in the regulation of DNA damage and genome instability. The ability of mitochondrial dynamics to regulate oncogenic DNA damage centres upon the control of minority MOMP, a process that enables non-lethal caspase activation leading to DNA damage. Mitochondrial fusion suppresses minority MOMP, and its associated DNA damage, by enabling homogenous mitochondrial expression of anti-apoptotic BCL-2 proteins. Finally, we find that mitochondrial dysfunction inhibits pro-apoptotic BAX retrotranslocation, causing BAX mitochondrial localization thereby promoting minority MOMP. Unexpectedly, these data reveal oncogenic effects of mitochondrial dysfunction that are mediated via mitochondrial dynamics and caspase-dependent DNA damage.
Introduction

Mitochondrial dysfunction has pleiotropic impact on cancer (Giampazolias and Tait, 2016). For instance, mitochondrial respiratory complex proteins and TCA enzymes bearing tumour associated mutations, generate oncometabolites (Isaacs et al., 2005; Pollard et al., 2007; Sciacovelli et al., 2016; Selak et al., 2005). Moreover, loss of function mutations in mitochondrial DNA (mtDNA) are common in cancer and have been shown to accelerate tumorigenesis (Gorelick et al., 2021; Smith et al., 2020). Nonetheless, how dysfunctional mitochondria promote cancer largely remains an open question.

While inhibition of mitochondrial apoptosis has well established oncogenic effects, through increased apoptotic priming, tumour cells are often sensitized to cell killing cancer therapies (Certo et al., 2006; Singh et al., 2019). Mitochondria regulate apoptosis via mitochondrial outer membrane permeabilization or MOMP (Bock and Tait, 2020). This key event releases soluble mitochondrial intermembrane space proteins into the cytoplasm, notably cytochrome c, that activate caspases proteases causing rapid cellular demise. Because it dictates cell fate, mitochondrial outer membrane integrity is tightly regulated by BCL-2 protein family members (Campbell and Tait, 2018).

MOMP is usually considered a lethal point-of-no-return due to its extensive nature, often occurring in all mitochondria, coupled to an invariable loss of mitochondrial function (Goldstein et al., 2000; Lartigue et al., 2009; Rehm et al., 2003). However, we have previously described conditions whereby MOMP can be heterogenous permitting cell survival (Ichim et al., 2015; Tait et al., 2010). Following a sub-lethal stress, a limited mitochondrial cohort selectively permeabilizes, which we termed minority MOMP (Ichim et al., 2015). Strikingly, minority MOMP can engage sub-lethal caspase activity promoting DNA damage that is dependent upon caspase-activated DNAse (CAD) (Ichim et al., 2015). By causing DNA damage, minority MOMP may contribute to the paradoxical oncogenic effects of apoptotic signaling reported in different studies (Ichim and Tait, 2016). Moreover, minority MOMP has been recently implicated in an expanding array of functions including increased cancer aggressiveness, innate immunity and inflammation triggered by mtDNA double-strand breaks (Berthenet et al., 2020; Brokatzky et al., 2019; Tigano et al., 2021).

Here, we investigated the relationship between mitochondrial dysfunction and DNA damage. Surprisingly, we uncovered a key role for mitochondrial dynamics in the regulation of DNA damage. Mitochondrial fission, a consequence of mitochondrial dysfunction, promotes minority MOMP causing caspase-dependent DNA damage and genome instability. Secondly, we find reduced retrotranslocation of pro-apoptotic BAX on dysfunctional mitochondria, thus
facilitating minority MOMP. These data reveal an unanticipated link between mitochondrial dysfunction and oncogenic DNA damage that is mediated through minority MOMP and caspase activity.

Results

Mitochondrial dynamics regulate DNA damage

We aimed to understand how mitochondrial dysfunction can be oncogenic. Given the tumor promoting roles of DNA damage, we initially investigated its interconnection with mitochondrial function. To cause mitochondrial dysfunction, U2OS and HeLa cells were treated with the uncoupler carbonyl cyanide m-chlorophenyl hydrazone (CCCP). In order to phenocopy increased apoptotic priming that is found in pre-malignant and tumour cells, we co-treated cells with ABT-737, a BH3-mimetic compound that selectively neutralises anti-apoptotic BCL-2, BCL-xL and BCL-w. The response to DNA damage was measured by γH2AX staining and flow cytometry. In both HeLa and U2OS cells, BH3-mimetic treatment led to an increase in γH2AX positive cells that was significantly enhanced by combined treatment with CCCP, consistent with mitochondrial dysfunction promoting DNA damage (Figure 1A). Given the low level of γH2AX-positive cells observed by flow cytometry, we visualised the degree of DNA damage in U2OS and HeLa cells treated with ABT-737 by indirect immunofluorescence and western blot. In agreement with the flow cytometry data, we observed a low, but detectable, level of γH2AX and pATM foci in cells treated with ABT-737 (Figure 1B, Supplementary Figure 1A, B). Mitochondrial dynamics and function are tightly interconnected such that mitochondrial dysfunction causes mitochondrial fission (Oltersdorf et al., 2005). We therefore investigated whether mitochondrial dynamics affected DNA damage triggered by BH3-mimetic treatment. To disrupt mitochondrial fusion, we used Mfn1/2−/− murine embryonic fibroblasts (Mfn1/2−/− MEF) and, as control, reconstituted these cells with MFN2 (Mfn1/2−/− + Mfn2 MEF). As expected, Mfn1/2−/− MEF displayed a hyper-fragmented mitochondrial network whereas MFN2 reconstitution of these cells (Mfn1/2−/− + Mfn2) restored mitochondrial fusion, resulting in a filamentous mitochondrial network, but did not affect growth following treatment with ABT-737 (Figure 1C, D, Supplementary Figure 3A, B). Of note, although Mfn2 protein expression is higher in Mfn1/2−/− MEF reconstituted with Mfn2 compared to Wt MEF, mitochondrial fusion was not enhanced in Mfn1/2−/− + Mfn2 MEF and mitochondrial fragmentation occurred with the same kinetics when exposed to CCCP, relative to Wt MEF (Supplementary Figure 1C, D). Furthermore, expression of mitochondrial fusion proteins Mfn1 and Mfn2 and the mitochondrial fission protein Drp1 were not changed following ABT-737 treatment (Supplementary Figure 1E). Mfn1/2−/− and Mfn1/2−/− + Mfn2 MEF were treated with ABT-737 (10 µM, 3 hours) and the DNA damage response was assessed by analyzing γH2AX levels
by western blot or by flow cytometry (Figures 1E, F). Mfn1/2−/− MEF exhibited increased γH2AX, consistent with mitochondrial fission promoting DNA damage. Because DNA damage can be oncogenic, we investigated if cells with extensive mitochondrial fission were more prone to transformation. Mfn1/2−/− and Mfn1/2−/−+ Mfn2 MEF were passaged repeatedly in ABT-737. Following treatment, cells were assayed for transformation in vitro by determining anchorage-independent growth in soft agar. Specifically following culture in ABT-737, Mfn1/2−/− MEF formed colonies more readily than Mfn1−/− + Mfn2 MEF (Figure 1G). MFN2 can have functions independent of mitochondrial fusion, for instance mitochondrial-endoplasmic reticulum (ER) tethering (de Brito and Scorrano, 2008). Therefore, we investigated whether mitochondrial dynamics regulate DNA-damage through an alternative approach by inhibiting mitochondrial fission. DRP1 plays a central role in mitochondrial fission (Ishihara et al., 2009; Wakabayashi et al., 2009). To inhibit mitochondrial fission we used Drp1fl/fl MEF, which when infected with adenoviral Cre efficiently delete Drp1, causing a hyper-fused mitochondrial network (Figure 1I, Supplementary Figure 1F, 3A). Infection with adenoviral Cre did not cause DNA damage, nor impact ABT-737-induced DNA damage or growth following ABT-737 treatment (Supplementary Figure 1G, 3B). Drp1fl/fl and Drp1−/− MEF were treated with ABT-737 and γH2AX was measured by flow cytometry, as before. MEF expressing Drp1 have elevated levels of γH2AX after exposure to ABT-737, but this was completely abolished in Drp1-deficient cells (Figure 1J). These data suggest that mitochondrial dysfunction and fission promote oncogenic DNA damage and transformation.

Mitochondrial dynamics regulate DNA damage and genome-instability in a caspase and CAD dependent manner

We next sought to understand how mitochondrial dynamics regulate DNA damage. Because we had found that pro-apoptotic BH3-mimetic treatment potentiated DNA damage, we investigated a role for apoptotic caspase function. Wild type MEF or MEF overexpressing DRP1 were treated with the pan-caspase inhibitor qVD-OPh and γH2AX was measured by flow cytometry and western blot. MEF cells overexpressing DRP1 displayed a more fragmented mitochondrial network and had higher levels of γH2AX compared to their empty vector counterparts, consistent with our earlier data, but showed similar proliferation rates following treatment with ABT-737 (Figure 2A, B, Supplementary Figure 2A, B, 3A, B). Crucially, γH2AX was prevented by treatment with the pan-caspase inhibitor qVD-OPh, demonstrating a key role for caspase activation in DNA damage (Figure 2B). We next set out to establish whether loss of Drp1 impacted oncogenic transformation following repeated culture in ABT-737; however, loss of Drp1 alone was sufficient to render cells resistant to Myc/Hras-induced transformation (Supplementary Figure 2C), consistent with the results of
Serasinghe and colleagues (Serasinghe et al., 2015). Given these findings, we investigated a possible correlation between expression of the mitochondrial fission protein DRP1 and mutational burden in cancer. TCGA PanCancer Atlas studies were investigated through cBioportal. Of these, a significant association between increased mutational count in DNM1L mRNA high quartile versus DNM1L mRNA low quartile was found in invasive breast carcinoma and lung adenocarcinoma (out of 22 studies) with the inverse relationship not observed in any cancer type (Figures 2C, 2D, Supplemental Table 1 and data not shown). In both invasive breast cancer and lung adenocarcinoma, DNA damage response pathways were enriched in the DNM1L mRNA high quartile consistent with engagement of DNA damage (Supplementary Figures 2D, E, F). Moreover, high DNM1L mRNA expression correlates with poorer survival in a cohort of lung adenocarcinoma (Supplementary Figure 2G). To further investigate the role of caspase activity, we investigated the impact of mitochondrial dynamics upon genome instability. To this end, we used the PALA assay, in which gene amplification of CAD (carbamyl phosphate synthetase/aspartate transcarbamylase/dihydro-orotase, note that this is distinct from caspase-activated Dnase) described later) enables resistance to PALA (N-phosphonoacetyl-L-aspartate) (Wahl et al., 1979). To determine if alterations in mitochondrial dynamics, also affect genome instability dependent upon caspase activity, we passaged Mfn1/2−/− and Mfn1/2−/− + Mfn2 MEF with sub-lethal doses of ABT-737 in the presence or absence of QVD-OPh. Following treatment, cells were grown in the presence of PALA and clonogenic survival was measured (Figure 2E). Importantly, ABT-737 treated Mfn1/2−/− MEF gave significantly more colonies than Mfn1/2−/− + Mfn2 following PALA treatment, in a caspase-dependent manner (Figure 2E, F). In line with increased survival following PALA treatment, qPCR revealed amplification of the Cad locus only in Mfn1/2−/− MEF repeatedly treated with ABT-737 (Figure 2G). We and others have previously found that non-lethal caspase activity can cause DNA damage and genome instability dependent upon caspase-activated DNAse (CAD) (Ichim et al., 2015; Lovric and Hawkins, 2010). To examine the role of CAD in genomic instability we used the Mfn1/2−/− and Mfn1/2−/− + Mfn2 MEF in which we deleted the Dff40 gene (encoding CAD) using CRISPR-Cas9 genome editing (Supplementary Figure 3C). As before, Mfn1/2−/− cells resisted PALA treatment and efficiently grew as colonies following ABT-737 treatment, whereas Mfn1/2−/− + Mfn2 cells did not (Figure 2H, I). However, deletion of CAD completely abrogated clonogenic potential. Cad DNA expression and anchorage-independent growth were also diminished following ABT-737 treatment in CAD/Dff40 deleted cells as compared to their controls (Figure 2J, K, Supplementary Figure 3D). Together, these data show that mitochondrial fission promotes genome instability in a caspase and CAD dependent manner.
Minority MOMP occurs on fragmented mitochondria and is regulated by mitochondrial dynamics

We have previously found that permeabilization of a small number of mitochondria in a cell — called minority MOMP — can engage non-lethal caspase activity causing CAD activation and DNA damage (Ichim et al., 2015). This knowledge, coupled to our previous data, led us to investigate a role for mitochondrial dynamics in the regulation of minority MOMP. To address this, we combined super-resolution Airyscan confocal microscopy together with our fluorescent reporter that allows detection of minority MOMP (Ichim et al., 2015). This reporter comprises cytosolic FKBP-GFP (cytoGFP) and mitochondrial inner membrane targeted FRB-mCherry (mito-mCherry). Upon loss of mitochondrial outer membrane integrity, and in the presence of chemical heterodimeriser (AP21967), these two proteins bind one another, recruiting cytoGFP to the permeabilized mitochondria (Figure 3A). HeLa or U2OS were treated with a non-lethal dose of BH3-mimetic ABT-737 (10 µM) for 3 hours. Consistent with our previous data, this treatment was sufficient to engage minority MOMP, as evidenced by localization of cytoGFP to specific mitochondria (Figure 3B). Super-resolution analysis of these mitochondria revealed that selectively permeabilised mitochondria were separate from the mitochondria network, suggesting that minority MOMP preferentially occurs on fragmented mitochondria (Figure 3B, C). Extensive mitochondrial fission is a well-established consequence of MOMP (Bhola et al., 2009; Frank et al., 2001). Therefore, to approach whether mitochondria fragmentation was a cause or consequence of minority MOMP, U2OS cells expressing cytoGFP and mito-mCherry were imaged by live-cell microscopy. Treatment with ABT-737 (10 µM) led to minority MOMP, apparent by the translocation of cytoGFP into mitochondria after 124 minutes. Importantly, these mitochondria were fragmented from the mitochondrial network prior to cytoGFP translocation at 120 minutes (Figure 3D, Movie 1). This suggests that minority MOMP preferentially occurs on fragmented mitochondria. We next used these cells to investigate a role for mitochondrial fusion in regulating minority MOMP. Mfn1/2−/− and Mfn1/2−/+ Mfn2 MEF expressing the MOMP reporter, were treated with a sub-lethal dose of ABT-737. Strikingly, increased levels of minority MOMP were observed in Mfn1/2−/− MEF when compared to Mfn1/2−/+ Mfn2 MEF (Figure 3E, Supplementary Figure 4A). This is consistent with minority MOMP occurring primarily on fragmented mitochondria, with mitochondrial fusion having an inhibitory effect. To further address this, we investigated the impact of inhibiting mitochondrial fission upon minority MOMP following treatment of Drp1fl/fl and Drp1−/− MEF with ABT-737. MEF expressing Drp1 undergo minority MOMP after exposure to ABT-737, but this was completely abolished in Drp1-deleted cells (Figure 3F, Supplementary Figure 4B). Together, these data demonstrate that mitochondrial dynamics regulate minority MOMP; mitochondrial fusion appears to be inhibitory whereas fission promotes minority MOMP.
Pro-survival BCL-2 proteins display inter-mitochondrial heterogeneity in localisation

Our data demonstrate that mitochondrial fission promotes minority MOMP enabling caspase-dependent DNA damage. Nevertheless, how mitochondrial dynamics regulate minority MOMP is not known. Mitochondrial outer membrane integrity is regulated by the balance of pro- and anti-apoptotic BCL-2 family proteins (Campbell and Tait, 2018). We hypothesised that inter-mitochondrial variation in BCL-2 family localisation may underlie minority MOMP. To investigate this hypothesis, we set out to visualise endogenous levels of BCL-2 family proteins on individual mitochondria. CRISPR-Cas9 genome editing can be used to knock-in fluorescent proteins at defined genomic loci to enable endogenous tagging of proteins (Bukhari and Muller, 2019). Using this approach, we generated clonal knock-in HeLa cell lines where the red fluorescent protein Scarlet was fused to the N-termini of BCL-2, BCL-xL and MCL-1. As verification, western blotting using antibodies specific BCL-2, BCL-xL, MCL-1 and Scarlet confirmed that these cell lines expressed these fusion proteins at similar levels to their endogenous counterparts (Figure 4A, B). Secondly, Airyscan super-resolution microscopy demonstrated mitochondrial localisation of Scarlet-BCL-2, BCL-xL and MCL-1, as expected (Figure 4C) and is comparable to mitochondrial localisation of untagged endogenous BCL-2 proteins in parental cells (Supplementary Figure 5A, B). Finally, we monitored cell viability using SYTOX Green exclusion and IncuCyte real-time imaging in response to BH3-mimetic treatment (ABT-737 and S63845). This demonstrated that all knock-in cell lines underwent cell death in response to BH3-mimetic treatment (Supplementary Figure 5C). Using these knock-in cells, we next acquired super-resolution microscopy images of Scarlet-tagged BCL-2, BCL-xL and MCL-1 then applied a colour grading lookup table (LUT) such that the brighter the Scarlet signal, the more bright the image. This revealed heterogeneity of Scarlet BCL-2, BCL-xL and MCL-1 across the mitochondrial network, which is also evident in parental cells stained with BCL-2 family antibodies (Figure 4D, Supplementary Figure 5A). Given our previous data, we hypothesised that BCL-2 family protein heterogeneity is regulated by mitochondrial dynamics. To test this, we inhibited mitochondrial fission through CRISPR-Cas9 deletion of DRP1. Western blot confirmed DRP1 deletion, resulting in extensive mitochondrial hyperfusion (Supplementary Figure 5D, E). Strikingly, cells with hyperfused mitochondria displayed much reduced inter-mitochondrial heterogeneity of MCL-1, BCL-2 or BCL-xL (Figure 4E, F, Supplementary Figure 5F). Combined, these data show that within a cell extensive inter-mitochondrial heterogeneity in BCL-2 localisation exists that is impacted by mitochondrial dynamics.

Heterogeneity in apoptotic priming underpins minority MOMP
We next investigated whether there was a relationship between localisation of anti-apoptotic BCL-2 proteins and minority MOMP. To determine this, we acquired super-resolution images of HeLa cells expressing endogenous Scarlet-BCL-2, BCL-xL or MCL-1, together with Omi-GFP and MitoTracker Deep Red. During MOMP, soluble intermembrane space proteins, including Omi, are released from mitochondria (Bock and Tait, 2020). Mitochondria retain MitoTracker Deep Red even after loss of mitochondrial integrity; thus, mitochondria that have undergone MOMP are identifiable by loss of Omi and MitoTracker retention. Surprisingly, live-cell imaging of BCL-2 family protein knock-in cells following treatment with ABT-737 revealed that mitochondria (determined by MitoTracker positivity) that release Omi-GFP have higher levels of BCL-2, BCL-xL or MCL-1 local mitochondrial levels prior to MOMP (Figure 5A-C). Computational segmentation allowed us to distinguish BCL-2 family protein mitochondrial localisation, on MitoTracker positive structures, which lack Omi expression, confirming that these mitochondria have indeed undergone minority MOMP. Quantification across a number of cells shows that mitochondria which undergo minority MOMP have increased BCL-2 family protein mitochondrial localisation (Figure 5D-F). Furthermore, line scans revealed regions of the mitochondrial network with high BCL-2 family residency, but low Omi expression, that is mitochondria which have likely undergone minority MOMP (Supplementary Figure 6A, B). Unexpectedly, these data reveal a correlation between increased anti-apoptotic BCL-2 mitochondrial localisation and selective mitochondrial permeabilisation. We reasoned that this may be analogous to increased apoptotic priming at the cellular level, where high anti-apoptotic BCL-2 mitochondrial localisation can correlate with apoptotic sensitivity in some cell types. Mitochondrial association of pro-apoptotic BAX is indicative of increasing apoptotic priming (Edlich et al., 2011; Reichenbach et al., 2017; Schellenberg et al., 2013). To investigate whether mitochondria with high BCL-2 local mitochondrial localisation may also display high BAX localisation (indicative of selective, increased apoptotic priming), we generated GFP-BAX expressing BCL-2 family knock-in HeLa cells and imaged them by super-resolution microscopy. In line with the notion that mitochondria with higher BCL-2 family localisation also have elevated BAX localisation, we observed BAX co-localising with high BCL-2 mitochondria, indicative of increased apoptotic priming (Figure 5G-I). These data demonstrate that inter-mitochondrial heterogeneity in anti-apoptotic BCL-2 mitochondrial localisation and apoptotic priming underlies minority MOMP.

Mitochondrial dysfunction promotes BAX accumulation promoting minority MOMP

We aimed to define the underlying mechanism of mitochondrial intrinsic apoptotic priming. In healthy cells, BAX undergoes mitochondrial retrotranslocation and inhibiting this process causes BAX mitochondrial accumulation, sensitising to MOMP (Edlich et al., 2011;
Schellenberg et al., 2013). HeLa cells expressing GFP-BAX and iRFP-Omp25 were treated with the uncoupler carbonyl cyanide m-chlorophenyl hydrazone (CCCP) to induce mitochondrial dysfunction. To facilitate visualisation of mitochondrial localised GFP-BAX, cells were treated with digitonin to selectively permeabilise the plasma membrane, as described previously (Bender et al., 2012). Inducing mitochondrial dysfunction by CCCP treatment led to robust mitochondrial recruitment of GFP-BAX (Figure 6A, Supplemental Figure 7A, Movies 2-5). Immunostaining of HeLa cells with the activation specific BAX antibody 6A7 revealed BAX activation, as expected, under conditions of apoptosis (combined BH3-mimetic treatment) but not following CCCP treatment (Supplementary Figure 7B). Since mitochondrial fusion promotes efficient oxidative phosphorylation, reducing heterogeneity in mitochondrial function (Chen et al., 2003), we hypothesised that by impacting mitochondrial function, mitochondrial fission may promote BAX recruitment thereby facilitating minority MOMP. We imaged Mfn1/2−/− and Mfn1/2−/− + Mfn2 MEF with MitoTracker Red, a potentiometric dye, that reports mitochondrial Δψm as a measure of mitochondrial function. Consistent with defective mitochondrial function, mitochondria in fusion defective cells (Mfn1/2−/−) displayed heterogenous MitoTracker Red signal and lower total signal than fusion competent Mfn1/2−/− + Mfn2 MEF (Figures 6B-D). We next analysed GFP-BAX localisation in Mfn1/2−/− and Mfn1/2−/− + Mfn2 MEF, using fluorescence loss in photobleaching (FLIP) to help visualise mitochondrial localised GFP-BAX. FLIP analysis of GFP-BAX revealed slower mitochondrial retrotranslocation in Mfn1/2−/− cells compared to Mfn1/2−/− + Mfn2 cells (Figure 6E-H, Movies 6, 7). This suggests that mitochondrial dysfunction, a consequence of defective mitochondrial dynamics, can promote GFP-BAX mitochondrial accumulation, serving as an mitochondrial intrinsic priming mechanism that facilitates minority MOMP.

Discussion

We describe that mitochondrial dysfunction, inducing mitochondrial fission, promotes DNA damage and genome instability. This process requires caspase activity, that is engaged by minority MOMP, in order to trigger DNA damage. Investigating the underlying mechanism, we find that mitochondrial dynamics affect inter-mitochondrial heterogeneity of anti-apoptotic BCL-2, permitting increased apoptotic priming of fragmented mitochondria. Mitochondrial dysfunction acts as a mitochondrial intrinsic priming signal by inhibiting pro-apoptotic BAX retrotranslocation, promoting minority MOMP. Unexpectedly, by affecting mitochondrial BCL-2 heterogeneity and apoptotic priming, our data reveal crucial roles for mitochondrial dysfunction and dynamics in the regulation of minority MOMP leading to caspase dependent DNA damage and genome instability.
Our study highlights that mitochondrial dynamics are integral to minority MOMP, whereby mitochondrial fusion inhibits, and fission promotes this process. Consistent with this finding, the ability of sub-lethal apoptotic stress to engage oncogenic caspase-dependent DNA damage and genome instability was regulated in a similar manner. Moreover, we found in some cancer types, a correlation between the expression of the mitochondrial fission protein DRP1, DNA damage and mutational burden. These data support an oncogenic role for mitochondrial fission, through its capacity to promote minority MOMP and associated sub-lethal caspase activity. This also suggests that the multitude of cellular signalling pathways and stresses that impact mitochondrial dynamics, for instance as hypoxia or high glycolytic rates, might facilitate minority MOMP induced transformation (Chen and Chan, 2017; Wu et al., 2016). Indeed, we found that enforced mitochondrial fission (through MFN1/2 deletion), promoted minority MOMP induced transformation. Supporting previous findings, we also report that DRP1 contributes to oncogene induced transformation - potentially, minority contributory role Serasinghe et al. Our study underscores extensive interplay exists between mitochondrial dynamics and cancer (Chen and Chan, 2017; Gao et al., 2017; Kashatus et al., 2015; Zhao et al., 2013).

We sought to define how mitochondrial dynamics might control minority MOMP. Surprisingly, we found inter-mitochondrial heterogeneity in anti-apoptotic BCL-2 localisation. This heterogeneity was suppressed by mitochondrial fusion, most likely because mitochondrial fusion enables homogenous distribution of BCL-2 proteins across the mitochondrial network. As we further discuss, heterogeneity in anti-apoptotic BCL-2 localisation enables differences in apoptotic priming of specific mitochondria. Interestingly, during cell death, mitochondrial variation in pro-apoptotic BAK levels have previously been found to influence the kinetics of MOMP (Weaver et al., 2014). Though myriad interconnections between mitochondrial dynamics and apoptosis exist, mitochondrial fission is largely considered a consequence of cell death. For instance, during apoptosis, extensive mitochondrial fragmentation occurs subsequent to MOMP (Bhola et al., 2009). By promoting homogenous BCL-2 localisation across the mitochondrial network, our data reveal an indirect role for mitochondrial fusion in preventing minority MOMP.

We have previously found that ectopic expression of BCL-2 can lead to incomplete MOMP, consistent with BCL-2 anti-apoptotic function (Tait et al., 2010). In the current study, we find that increased local mitochondrial levels of anti-apoptotic BCL-2 family proteins correlates with selective mitochondrial permeabilisation. While this may seem initially counter-intuitive, precedence for increased apoptotic priming, correlating with high anti-apoptotic BCL-2 levels is evident in various cancers (Certo et al., 2006; Singh et al., 2019). This is perhaps
best demonstrated in high-BCL-2 expressing chronic lymphocytic leukaemia (CLL) that is often highly sensitive to the BCL-2 selective BH3-mimetic, venetoclax (Roberts et al., 2016). In healthy cells, BAX mitochondrial localisation is indicative of increased apoptotic priming (Edlich et al., 2011; Kuwana et al., 2020; Reichenbach et al., 2017; Schellenberg et al., 2013). Indeed, further investigation revealed that high pro-apoptotic BAX localisation correlated with high-BCL-2 localisation on mitochondria. Our data argue that heterogeneity in apoptotic priming exists not only between cell types, but also intracellularly, at the level of individual mitochondria.

Finally, we sought to understand how inter-mitochondrial heterogeneity in apoptotic priming might occur. Pro-apoptotic BAX is subject to constant mitochondrial retrotranslocation; inhibition of BAX retrotranslocation leads to mitochondrial accumulation, sensitising to apoptosis (Edlich et al., 2011; Schellenberg et al., 2013). We find that reduction of mitochondrial inner membrane potential ($\Delta \psi_m$) promotes BAX mitochondrial localisation. Importantly, reduction of $\Delta \psi_m$, provides a mitochondrial-intrinsic signal to increase apoptotic priming. Under conditions of mitochondrial dysfunction imposed by loss of mitochondrial fusion, we find decreased rates of BAX retro translocation, enabling its mitochondrial accumulation. In essence, BAX retrotranslocation may serve as a barometer of cellular metabolic health. Because loss of mitochondrial function causes mitochondrial fission, it promotes minority MOMP in a two-fold manner, segregating dysfunctional mitochondria and promoting BAX accumulation (Twig et al., 2008). Further investigation will be required to mechanistically delineate how mitochondrial function regulates BAX retrotranslocation. We consider it likely that additional mechanisms of mitochondrial-intrinsic priming also exist, for instance levels of anti-apoptotic BCL-2 mitochondrial localisation will dictate a given cell’s propensity to engage minority MOMP. Moreover, degradation of dysfunctional mitochondria through mitophagy, may also impact the occurrence of minority MOMP. Indeed, others have reported that permeabilised mitochondria are targeted for mitophagy - this likely affects the steady-state detection of minority MOMP (Lindqvist et al., 2018).

Our study investigates the impact of mitochondrial dynamics and dysfunction upon BH3-mimetic induced minority MOMP - when might this occur physiologically? One possibility is during intracellular bacterial infection (Salmonella Typhimurium and Chlamydia trachomatis) where minority MOMP has been shown to cause inflammation as an innate immune response (Brokatzky et al., 2019). Interestingly, while C. trachomatis actively promotes mitochondrial fusion during early infection, late-stage infection is associated with mitochondrial fragmentation, providing an ideal setting to engage minority MOMP induced inflammation.
More recently, mitochondrial DNA (mtDNA) damage has been shown to promote minority MOMP, enabling mitochondrial-nuclear retrograde signalling (Tigano et al., 2021). In this setting, mitochondrial fragmentation most likely enables some mitochondria to selectively permeabilise in response to mtDNA-damage. A final example may relate to the minority MOMP that we have previously observed under homeostatic conditions (Ichim et al., 2015). Indeed, others have found basal levels of caspase/CAD-dependent DNA-damage in cycling cells (Liu et al., 2017). Given extensive mitochondrial fission occurs during mitosis, this may serve to underpin basal levels of minority MOMP and caspase-dependent DNA-damage.

In summary, our findings that reveal that mitochondrial dynamics regulate DNA damage and genome instability via minority MOMP induced caspase-activity. This provides a mechanism linking mitochondrial dysfunction to pro-oncogenic DNA damage. Beyond pro-tumourigenic effects, minority MOMP has also been shown to have roles in innate immunity and inflammation, as such, our findings suggest new approaches to modulate minority MOMP and its downstream functions.

Methods

Cell Lines

HeLa and U2OS cells were purchased from ATCC (LGC Standards). 293FT cells were purchased from Thermo Fisher Scientific.

Mfn1/2−/− MEF were provided by David Chan, Caltech and reconstituted with LZRS-MFN2 in our laboratory. Drp1−/− MEF were provided by Hiromi Sesaki, Johns Hopkins University School of Medicine. MEF Wt and MEF loxP-STOP-loxP EYFP were a kind gift from Douglas Green, St Jude’s Research Children’s Hospital. All cell lines were cultured in DMEM high-glucose medium supplemented with 10% FCS, 2 mM glutamine, 1 mM sodium pyruvate, penicillin (10,000 units/ml) and streptomycin (10,000 units/ml).

To delete Drp1 from Drp1−/− MEF, 2 x 10⁶ cells were seeded and infected with 200 MOI Ad5CMVCre (Viral Vector Core, University of Iowa) for 8 h, after which the media was replaced. Cells were used for experiments from the following day.

METHOD DETAILS

Generation of Scarlet-BCL-2 knock-in cell lines
We used a modified version of the knock-in strategy described in (Stewart-Ornstein and Lahav, 2016). Two vectors were used: the first vector comprises 500bp homology arm before and after the start codon of BCL-2, in between which is the Scarlet coding sequence, cloned into pUC-SP. The second vector, pSpCas9(BB)-2A-Puro (Addgene #48139) comprises Cas9 and the sgRNA targeting sequence. The following sgRNA sequences were used:

- **Human BCL-2**: 5'- ATGGCGCACGCTGGGAGAAC -3'
- **Human BCL-xL**: 5'- AAAAAATGTCTCAGACCAACC -3'
- **Human MCL-1**: 5'- CGGCGGCGACTGGCAATGTT -3'

To generate the knock-in cells, HeLa cells were transfected with 1 μg of homology arm vector and 1 μg of pSpCas9(BB)-2A-Puro with Lipofectamine 2000, according to the manufacturer’s instructions. Media was removed 5 h later, and replaced with media containing 1 μM SCR7 for 2 days. Cells were selected with 1 μg/mL puromycin for a further two days before selecting Scarlet positive clones by FACS. Cells which expressed Scarlet signal which co-localised with mitochondria we used for further experiments.

**Generation of stable overexpressing cell lines**

For retroviral transduction, 293FT cells were transfected with 5 μg of plasmid, together with 1.2 μg gag/pol (Addgene #14887) and 2.4 μg VSVG (Addgene #8454) using Lipofectamine 2000. Media was changed after 6 hours and collected, filtered and used to infect cells 24 and 48 h post-transfection in the presence of 1 μg/ml Polybrene. 24 h following infection, cells were allowed to recover in fresh medium and incubated with selection antibiotic 24 h after. Cells were selected with appropriate antibiotic or FACS sorted to isolate a high-expressing population. Concentrations used for antibiotic selection were 200 μg/ml zeocin (Invivogen) or 1 μg/ml puromycin (Sigma).

For lentiviral transduction, the procedure was the same as for retroviral transduction, except 5 μg plasmid was transfected into 293FT along with 1.86 μg psPAX2 (Addgene #12260) and 1 μg VSVG (Addgene #8454) using Lipofectamine 2000.

**Generation of CRISPR knock-out cell lines**

Human Drp1 and mouse Dff40 knock-out cell lines were generated by CRISPR-Cas9 gene deletion, using the lentiviral transduction protocol above. The following sequences were cloned into LentiCRISPRv2-puro (Addgene #52961):

- **Human Drp1**: 5'- AAATCAGAGAGCTCATTCTT - 3'
- **Mouse Dff40**: 5'- ACATGGAGCCAAGGACTCG -3'
Plasmids

LZRS-Drp1 was generated by cloning the Drp1 coding sequence from pcDNA3.1(+) Drp1 (Addgene #34706) into LZRS backbone using Gibson Assembly. pBABE iRFP-Omp25 was cloned by Gibson Assembly using fragments derived from pLJM2 SNAP-Omp25 (Addgene #69599) and pMito-iRFP670 (Addgene #45462). Omi-GFP (in eGFPN2) was a kind gift from Douglas Green, St Jude’s Children’s Research Hospital.

Western Blotting

Cells were collected and lysed in NP-40 lysis buffer (1% NP-40, 1 mM EDTA, 150 mM NaCl, 50 mM Tris-Cl pH 7.4) supplemented with complete protease inhibitor (Roche). Protein concentration of cleared lysates was determined by Bradford assay (Bio-Rad). Equal amounts of protein lysates were subjected to electrophoresis through 10 or 12% SDS-PAGE gels and transferred onto nitrocellulose membranes, which were blocked for 1 h in 5% milk/PBS-Tween at room temperature. Membranes were incubated with primary antibody overnight at 4°C overnight. After washing, membranes were incubated with either goat-anti-rabbit Alexa Fluor 800, goat-anti-mouse Alexa Fluor 680 or goat-anti-rat DyLight 800 for 1 h at room temperature before detection using a Li-Cor Odyssey CLx (Li-Cor).

Flow Cytometry

For measuring levels of γH2AX, cells were trypsinised and washed once with PBS and fixed in 4% PFA for 15 minutes at room temperature. After washing once in PBS, cells were resuspended in 300 µL and 700 µL cold ethanol added dropwise while slowly vortexing. Samples were frozen at -20°C overnight. The following day, samples were washed with PBS and blocked in 2% BSA in PBS for 1 h at room temperature and incubated with anti-γH2AX antibody conjugated to Alexa Fluor 647 (BioLegend) for 30 minutes protected from light. Samples were analysed on the BD LSRFortessa flow cytometer (BD Biosciences) using standard protocols.

To measure mitochondrial potential in Mfn1/2−/− and Mfn1/2−/− + Mfn2 MEF, cells were incubated with 50nM MitoTracker CMXRos (Thermo Fisher Scientific) for 15 mins before collection. Cells were analysed on a Attune NxT flow cytometer (Thermo Fisher Scientific) using standard protocols, and analysed in FlowJo (BD).

PALA Assay and Cad Genomic Amplification

Cells were seeded in triplicate in 6 well plates at a density of 2500 cells per well and cultured in nucleoside-free α-MEM medium supplemented with 10% dialysed FBS. PALA was added
at the LD<sub>50</sub> dose and cells maintained until visible colonies formed. Colonies were fixed and stained in methylene blue (1% methylene blue in 50:50 methanol:water).

To assay Cad genomic amplification, DNA was extracted from PALA resistant colonies, or, in the case of control treated cells where no colonies were viable, DNA was extracted from cells passaged twenty times in DMSO, but not subjected to PALA treatment.

**Anchorage-independent growth assay**

A 1% base low melting temperature agarose solution (Sigma-Aldrich) was added to 6 well plates and allowed to set. 7,500 cells were suspended in 0.6% agarose in a 1:1: ratio to achieve a final concentration of 0.3% agarose., which was added on top of base agarose. When set, the cell/agarose mix was overlaid with complete DMEM media and colonies counted 14 days later from 15 fields of view per cell line.

**qPCR**

Genomic DNA was isolated from cells using the GeneJET DNA Extraction Kit (Thermo Fisher Scientific). PCR was performed on a Bio-Rad C1000 Thermal Cycler using the following conditions: 3 min at 95°C, 40 cycles of 20 s at 95°C, 30 s at 57°C, 30 s at 72°C and a final 5 min at 72°C using Brilliant III Ultra-Fast SYBR Green qPCR Master Mix (Agilent Technologies). Relative DNA quantification was analysed by the 2<sup>ΔΔCt</sup> method. Primer sequences used are as follows:

- Mouse CAD-F: AAGCTCAGATCCTAGTGCTAACG
- Mouse CAD-R: CCGTAGTTGCGATGAGG
- Mouse 18S-F: ATGGTAGTCCCGGTGCCTAC
- Mouse 18S-R: CCGGAATCGAACCCTGATT

**Microscopy**

**Fixed cell imaging**

Cells were grown on coverslips and fixed in 4% PFA/PBS for 10 min, followed by permeabilization in 0.2% Triton-X-100/PBS for 15 min. Cells were blocked for 1 h in 2% BSA/PBS and incubated with primary antibodies overnight at 4°C in a humidified chamber. The following day, cells were washed in PBS and secondary antibodies added for 1 h at room temperature, before final wash steps and mounting in Vectashield antifade mounting media.

**MOMP assay**

Cells were transfected with 250ng CytoGFP and 250ng mito-mCherry for 16 h with either Lipofectamine 2000 or GeneJuice before treatment in combination with 50 nM A/C
heterodimer (Clontech). Minority MOMP was scored based on co-localisation of CytoGFP with mito-mCherry. A minimum of 100 cells were analysed per condition, and we defined minority MOMP as a cell which has 1 or more instances of cytoGFP/mito-mCherry co-localisation.

**Airyscan super-resolution imaging**

Super-resolution Airyscan images were acquired on a Zeiss LSM 880 with Airyscan microscope (Carl Zeiss). Data were collected using a 63 x 1.4 NA objective for the majority of experiments, although some were acquired using a 40 x 1.3 NA objective. 405nm, 561nm and 640 nm laser lines were used, in addition to a multi-line argon laser (488nm) and images acquired sequentially using the optimal resolution determined by the Zeiss ZEN software. When acquiring z-stacks, the software-recommend slice size was used. Live-cell experiments were performed in an environmental chamber at 37°C and 5% CO₂. Airyscan processing was performed using the Airyscan processing function in the ZEN software, and to maintain clarity some images have been pseudocoloured and brightness and contrast altered in FIJI (ImageJ v2.0.0).

**Nikon A1R imaging**

Confocal images were acquired on a Nikon A1R microscope (Nikon). Data were collected using a 60 x Plan Apo VC Oil DIC N2 objective. 405nm, 561nm, 638nm laser lines were used, in addition to a multi-line argon laser (488nm). Images were acquired sequentially to avoid bleedthrough. For live-cell imaging, cells were imaged in a humidified environmental chamber at 37°C and 5% CO₂. Images were minimally processed in FIJI (ImageJ v2.0.0) to adjust brightness and contrast.

**3D rendering and image analysis**

Z-stacks acquired on the Zeiss LSM 880 with Airyscan microscope were imported into Imaris (Bitplane, Switzerland). To segment Omi and BCL-2, a surface was created using the Omi-GFP pixel information. Masks were applied to differentiate between BCL-2 inside and outside the Omi surface. From these masks, spots were created from the BCL-2 channel and quantified based on intensity of BCL-2 on mitochondria undergoing minority MOMP.

**Fluorescence Loss in Photobleaching**

The time lapse images were taken at one frame per second on a Zeiss LSM880 microscope in a 37°C chamber, with a Fluor 40x/1.30 NA oil immersion lens. Cells were in HEPES containing media. Fluorescence was excited using the 488nm line of an argon ion laser at 5.4% power through the AOTF and the emitted signal was captured with an Airyscan detector.
in Resolution vs. Signal mode using a combination of LP500 and BP485-550 filters with a zoom of 3.6. A letterbox bleach region covering approximately a third of the cell area was achieved using combination of 405 and 488nm wavelengths of light at 100% laser power and 100 iterations which took approximately 14 seconds.

**Digitionin permeabilisation**

Prior to digitonin permeabilisation, cells were incubated in FluoroBrite DMEM without FBS. To permeabilise the plasma membrane, 20 µM digitonin (Sigma) was added and cells imaged immediately.

**Mitochondrial analysis**

Cells stained with TOM20 and imaged on the Zeiss LSM 880 with Airyscan. These images were analysed by dividing mitochondrial length by width (mitochondria aspect ratio) in ImageJ. Heterogeneity of BCL-2 localisation was measured by calculating the standard deviation of Scarlet and mitochondrial signals in mitochondrial regions in FIJI (ImageJ v2.0.0).

**Live-cell viability assays**

Cell viability was assayed using either an IncuCyte ZOOM or IncuCyte S3 imaging system (Sartorius). Cells were seeded overnight and drugged in the presence of 30 nM SYTOX Green (Thermo Fisher Scientific), which is a non-cell-permeable nuclear stain. Data were analysed in the IncuCyte software, and where different cell lines are compared the data are normalised to starting confluency.

**Bioinformatic Analysis**

Relationship between DRP1 (*DNM1L*) expression and mutational count were investigated in TCGA PanCancer Atlas studies through cBioportal (Cerami et al., 2012; Gao et al., 2013). Studies with greater than 100 samples were analysed and samples divided into quartiles of *DNM1L*: mRNA expression z-scores relative to diploid samples (RNA Seq V2 RSEM). Of these, a significant association between increased mutational count in *DNM1L* mRNA highest quartile versus *DNM1L* mRNA lowest quartile was found in 2 out of 22 studies with the inverse relationship not observed in any cancer type. Mutation count in *DNM1L* quartiles was viewed in the Clinical Tab, statistical analysis of mutation count was performed by cBioportal, Wilcoxon test, q-value <0.05 was considered significant. As the relationship between *DNM1L* and mutational count was highly significant in Invasive Breast Carcinoma and Non-Small Cell Lung Cancer, we used these studies for further interrogation with cases of Lung adenocarcinoma selected from Non-Small Cell Lung Cancer dataset (not Lung squamous cell carcinoma). Data were downloaded from cBioportal and mutational count in *DNM1L* mRNA
highest quartile versus DNM1L mRNA lowest quartile (mRNA expression z-scores relative to diploid samples (RNA Seq V2 RSEM)) plotted in GraphPad Prism Version 9.0.0 and statistical significance between groups calculated by Mann-Whitney test. Data points represent individual patient samples, bar is mean (SD). DNM1L quartiles each contain 128 samples (Lung Adenocarcinoma TCGA PanCancer Atlas dataset) or 271 samples (Breast Invasive Carcinoma TCGA PanCancer Atlas dataset). Differentially expressed proteins in DNM1L highest versus lowest quartiles were also determined in cBioportal (measured by reverse-phase protein array, Z-scores) where significant differences are determined by Student’s t-test (p value) and Benjamini-Hochberg procedure (q value). Pathway analysis was performed using gene names of proteins identified with significantly higher expression in DNM1L high versus DNM1L low quartiles (excluding phospho-specific proteins, see lists in Supplementary Table 1) in GO Biological Process 2018 through Enrichr (Chen et al., 2013; Kuleshov et al., 2016).

Relationship between DRP1 (DNM1L) expression and patient survival were investigated in TCGA PanCancer Atlas studies through cBioportal (Cerami et al., 2012; Gao et al., 2013). Lung Adenocarcinoma TCGA PanCancer Atlas and Breast Invasive Carcinoma TCGA PanCancer Atlas datasets were downloaded from cBioportal, samples divided into quartiles of DNM1L: mRNA expression (z-scores relative to normal samples log RNA Seq V2 RSEM), with survival of highest and lowest quartiles plotted in GraphPad Prism Version 9.0.0 and statistical significance between groups calculated by Log-rank (Mantel-Cox) test.

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Author contributions
KC, JSR and SWGT conceived the study and designed the workplan. Experimental work: KC, JSR, RH, AEM-G, EV, KB, CC, YE, DGS, GI, KJC, APG. Development and contribution of reagents: KC, JSR, CC, Gl. Data analysis: KC, JSR, YE, KJC, APG, SWGT. Intellectual input: KC, JSR, KJC, APG, SWGT. Manuscript writing: JSR and SWGT.
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A) DMSO + Dimeriser  ABT-737 + Dimeriser

B) % of minority MOMP

C) U2OS ABT-737 + Dimeriser

Figure 1
Figure 2

A. Images showing cellular morphology of Mfn1/2−/− and Mfn1−/− cells.

B. Western blot analysis of MFN2 and α-Tubulin expression levels in Mfn1/2−/− and Mfn1−/− cells.

C. Graph showing percentage of cells with minority MOMP in Mfn1/2−/− and Mfn1−/− cells treated with DMSO or ABT-737.

D. Graph showing γH2AX+ cell counts in Mfn1/2−/− and Mfn1−/− cells treated with DMSO or ABT-737.

E. Western blot analysis of γH2AX and β-Tubulin expression levels in Mfn1/2−/− and Mfn1−/− cells treated with DMSO or ABT-737.

F. Images showing γH2AX localization in Wt and Drp1−/− cells.

G. Graph showing γH2AX+ cell counts in Mfn1/2−/− and Mfn1−/− cells treated with DMSO or ABT-737.

H. Graph showing γH2AX+ cell counts in Mfn1/2−/− and Mfn1−/− cells treated with DMSO or ABT-737/QVD.

I. Images showing γH2AX localization in LZRS-EV and LZRS-DRP1 cells.

J. Graph showing γH2AX+ cell counts in LZRS-EV and LZRS-DRP1 cells treated with DMSO, ABT-737, or ABT-737/QVD.
Figure 3
Figure 4
Figure 5

**Figure 5**

A. HeLa Scarlet-BCL-2  
B. HeLa Scarlet-BCL-xL  
C. HeLa Scarlet-MCL-1

D. Average Scarlet-BCL-2 Intensity  
E. Average Scarlet-BCL-xL Intensity  
F. Average Scarlet-MCL-1 Intensity

G. HeLa Scarlet-BCL-2  
H. HeLa Scarlet-BCL-xL  
I. HeLa Scarlet-MCL-1
**Supplementary Figure 2 - related to Figure 3**

### A

- **Dff40**
- **CRISPR**
- **Mfn1/2**
- **-/-**
- **Mfn1**
- **-/-**
- **Mfn1/2**
- **-/-**
- **DMSO**
- **EMPTY**
- **ABT-737**
- **Digestion**

### B

- **EMPT-CRISPR**
- **DMSO**
- **ABT-737**
- **100 µm**

### C

**Lung Adenocarcinoma**

- GO Biological Processes Enriched in DNMM1L High

### D

**Breast Invasive Carcinoma**

- GO Biological Processes Enriched in DNMM1L High

### E

**Breast Invasive Carcinoma and Lung Adenocarcinoma**

- GO Biological Processes Enriched in DNMM1L High

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*Supplementary Figure 2 - related to Figure 3*
Supplementary Figure 3

A. Graph showing SYTOX Green^+ (normalised to starting confluence) over time for HeLa cells and HeLa cells expressing Scarlet-BCL-2, Scarlet-BCL-xL, and Scarlet-MCL-1.

B. Western blot analysis showing DRP1 and α-Tubulin expression for EMPTY and Drp1 CRISPR conditions.

C. Microscopy images of HeLa Scarlet-BCL-2 cells under EMPTY and Drp1 CRISPR conditions.

D. Scatter plots showing the Scarlet/MitoTracker StDev Ratio for BCL-2, BCL-xL, and MCL-1 under EMPTY and Drp1 CRISPR conditions.

F. Graph showing time (h) versus SYTOX Green^+ (normalised to starting confluence) for HeLa Scarlet-BCL-2, HeLa Scarlet-MCL-1, and HeLa Scarlet-BCL-xL cells.
Supplementary Figure 5 - related to Figure 6

A: Digitonin permeabilisation

| Time (s) | Untreated | CCCP |
|----------|-----------|------|
| 0        | GFP-BAX   | BCL-2|
| 75       |           |      |
| 150      |           |      |
| 225      |           |      |
| 300      |           |      |
| 375      |           |      |

B: ABT-737/ S63845/ QVD CCCP

|          | Untreated | ABT-737/ S63845/ QVD | CCCP |
|----------|-----------|-----------------------|------|
| TOM20    |           |                       |      |
| GFP-BAX  |           |                       |      |
| BAX (6A7)|           |                       |      |
| Merge    |           |                       |      |