The use of a neutral peptide aptamer scaffold to anchor BH3 peptides constitutes a viable approach to studying their function

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The B-cell CLL/lymphoma-2 (Bcl-2) family of proteins are important regulators of the intrinsic pathway of apoptosis, and their interactions, driven by Bcl-2 homology (BH) domains, are of great interest in cancer research. Particularly, the BH3 domain is of clinical relevance, as it promotes apoptosis through activation of Bcl-2-associated x protein (Bax) and Bcl-2 antagonist killer (Bak), as well as by antagonising the anti-apoptotic Bcl-2 family members. Although investigated extensively in vitro, the study of the BH3 domain alone inside cells is more problematic because of diminished secondary structure of the unconstrained peptide and a lack of stability. In this study, we report the successful use of a novel peptide aptamer scaffold – Stefin A quadruple mutant – to anchor and present the BH3 domains from Bcl-2-interacting mediator of cell death (Bim), p53 upregulated modulator of apoptosis (Puma), Bcl-2-associated death promoter (Bad) and Noxa, and demonstrate its usefulness in the study of the BH3 domains in vivo. When expressed intracellularly, anchored BH3 peptides exhibit much the same binding specificities previously established in vitro, however, we find that, at endogenous expression levels, Bcl-2 does not bind to any of the anchored BH3 domains tested. Nonetheless, when expressed inside cells the anchored PUMA and Bim BH3 α-helices powerfully induce cell death in the absence of efficient targeting to the mitochondrial membrane, whereas the Noxa helix requires a membrane insertion domain in order to kill Mcl-1-dependent myeloma cells. Finally, the binding of the Bim BH3 peptide to Bak was the only interaction with a pro-apoptotic effector protein observed in this study.

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Programmed cell death is a tightly controlled process encompassing a wide range of regulatory mechanisms. Permeabilisation of the mitochondrial membrane and the resulting release of cytochrome c into the cytoplasm is a central event in the induction of the intrinsic apoptotic pathway. The B-cell CLL/lymphoma-2 (Bcl-2) family of proteins is chiefly involved in the regulation of apoptosis at the mitochondrion.¹–⁵ As well as apoptosis, more recent evidence has implicated the Bcl-2 family in the regulation of mitochondrial morphology and autoptaphagy.⁶,⁷ Structurally, the Bcl-2 proteins are characterised by four alpha helical Bcl-2 homology domains (BH1–4), at least one of which is present in each family member. In functional terms, the protein family is divided into three groups: the apoptotic effector proteins Bcl-2-associated x protein (Bax) and Bcl-2 antagonist killer (Bak; and possibly Bok), the anti-apoptotic proteins Bcl-2, Bcl-2-related gene long isoform (Bcl-xl), Bcl-w, Mcl-1 and Bcl-2-related gene A1 (A1), and finally the pro-apoptotic BH3-only proteins. In the last group, one can further distinguish between activators (Bcl-2-interacting mediator of cell death (Bim) and Bcl-2-interacting domain death agonist (Bid)) and sensitisers (e.g., p53 upregulated modulator of apoptosis (Puma), Bcl-2-associated death promoter (Bad), Noxa, Bcl-2-interacting killer (Bik) and more). In the event of cytotoxic stress, Bak and/or Bax oligomerise and produce pores in the mitochondrial outer membrane (MOM) thus initiating MOM permeabilisation (MOMP) and cytochrome c release.³ The mechanism of Bcl-2 family-regulated apoptosis is predominantly described by either the direct or indirect activation models, or a combination thereof (reviewed in Chipuk et al.³ and Shamas-Din et al.⁵).

The pro-apoptotic BH3-only proteins are diverse in structure and largely unrelated except for their BH3 domain, which represents the minimal unit of their function and is necessary

Abbreviations: A1, Bcl-2-related gene A1; Bad, Bcl-2-associated death promoter; Bak, Bcl-2 antagonist killer; Bax, Bcl-2-associated x protein; Bcl-2, B-cell CLL/lymphoma-2; Bcl-xl, Bcl-2-related gene long isoform; BH domain, Bcl-2 homology domain; Bid, Bcl-2-interacting domain death agonist; Bik, Bcl-2-interacting killer; Bim, Bcl-2-interacting mediator of cell death; CD, circular dichroism; DAPI, 4',6-diamidino-2-phenylindole; GFP, green fluorescent protein; MOM(P), mitochondrial outer membrane (permeabilisation); PCR, polymerase chain reaction; Puma, p53 upregulated modulator of apoptosis; SQT, Stefin A quadruple mutant Tracy

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for target protein binding and induction of apoptosis.8–11 Binding of the BH3-only Bcl-2 family members to each of their anti-apoptotic counterparts occurs in a selective manner, which closely correlates with their apoptotic potency.12,13 Bim, Puma and Bid have been found to bind to all five anti-apoptotic Bcl-2 proteins; when overexpressed in cells they potently induce cell death. Bad and Noxa, on the other hand, bind to complementary subsets of anti-apoptotic factors, with Bad interacting strongly with Bcl-2, Bcl-xI and Bcl-w and Noxa binding to Mcl-1 and A1.11,14 Overexpression of Bad and Noxa individually failed to cause apoptosis in mouse embryonic fibroblasts.12 Furthermore, Bim and Bid (and maybe Puma) directly bind and activate Bax and Bak.8,15–17

As aberrant regulation of apoptosis by the Bcl-2 family often lies at the heart of many diseases, the development of molecular tools, which help dissect the pathways involved, is of paramount importance. In particular, the role of the BH3 domain in isolation (i.e., outside the context of the full-length protein) as the mediator of the death signal is of significance to clinical research, as this interaction site has become an attractive drug target in the treatment of cancer.18,19 The interactions of the BH3 peptides with the anti-apoptotic and pro-apoptotic-effector Bcl-2 proteins have been studied thoroughly in vitro, however, the BH3-only proteins are not easily express recombinantly,12 making it difficult to derive reliable conclusions from peptide-based interaction studies. Moreover, introducing peptides into cells is challenging and once inside peptides are susceptible to proteolytic degradation.20,21 Consequently, rather less emphasis has been placed on the study of the BH3 peptides in vivo. A further caveat to the use of the BH3 peptides in isolation is their lack of secondary structure, which – in the absence of a structural constraint – precludes high-affinity interactions with their targets.16,22

We have recently reported the design and validation of a series of new peptide aptamer scaffolds, culminating in Stefin A quadruple mutant Tracy (SQT), derived from the mammalian protease inhibitor Stefin A.23 Peptide aptamers represent a relatively new class of alternative binding proteins, and are defined as binding peptides constrained within a protein scaffold.24 As binding molecules they are superior to free peptides, which are susceptible to proteolytic cleavage in cells, and which – because of their flexible nature – tend to bind to protein surfaces non-specifically. In this study, we explore the use of a novel peptide aptamer scaffold to anchor and present four different BH3 domains as a means of investigating their functions when expressed in cells. We propose that such tools may in time allow a direct correlation between biochemical and intracellular studies of BH3 helix function.

**Results**

**Structural analysis of the scaffold-constrained BH3 domains.** Based on previous studies,12,25 26-amino-acid stretches covering the BH3 region of Puma, Bim, Noxa and Bad, respectively (Figure 1a), were introduced into the N-terminal insertion site of the SQT peptide aptamer scaffold (Figure 1b), thus creating pepPuma, pepBim, pepNoxa and pepBad (collectively referred to as ‘pepBH3’). Circular dichroism (CD) spectroscopy was used to assess the secondary structure of the pepBH3 proteins (Figure 1c).

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**Figure 1** Structural analysis of the SQT scaffold presenting four different pro-apoptotic BH3 domains. (a) Amino-acid sequence of the four BH3 peptides used in this study. In the Puma sequence, the underlined amino acids are deleted in pepPumaDel. (b) Nuclear magnetic resonance (NMR) structure of Stefin A (adapted from pdb ref: 1DVC). The three sites for peptide presentation in the SQT peptide aptamer scaffold are highlighted in red. (c) The CD spectrum of SQT alone and with N-terminal insertions of the HA tag (a negative control not expected to adopt an alpha helical configuration) or with each BH3 peptide. The single inflection point around 218 nm indicates predominantly β-strands in the SQT and SQT-HA spectra. A shift in the inflection point to 220–222 nm, the deeper inflection of the spectrum and an additional shoulder signal the alpha helix formed by the BH3 peptides. SQT with Noxa BH3 and the HA peptide have been reported previously23.
The empty SQT scaffold contains predominantly β-strands, which is clearly reflected in its CD spectrum. Insertion of the BH3 domains altered the CD spectrum in a manner that is indicative of α-helices. When introducing the HA epitope tag into the same insertion site, the CD spectrum of SQT remains largely unchanged, suggesting that anchoring the BH3 peptides within a larger open reading frame (ORF) allows them to adopt a stable α-helical conformation.

**Functional analysis of pepBH3.** To study these newly created pepBH3 proteins intracellularly, a retroviral system (ProteoTuner) was used to stably transduce A375 melanoma cells with the pepBH3-encoding ORFs. In this expression system, proteins are constitutively targeted to the proteasome until addition of a chemical stabiliser (Shield). To confirm the suitability of the ProteoTuner system, expression of SQT after addition of Shield (5 μM) at numerous time points was examined (Figure 2a). SQT protein was detectable after 0.5 h and increased during the time course. Next, co-immunoprecipitation (co-IP) was used to investigate the ability of each pepBH3 protein to interact with other Bcl-2 protein family members inside cells (Figure 2b). PepPuma and pepBim co-IP with the anti-apoptotic proteins Bcl-xL and Mcl-1, whereas pepBad only co-IP with Bcl-xL and pepNoxa with Mcl-1, the latter as previously reported. The only interaction detectable between a pepBH3 and a pro-apoptotic effector protein was between pepBin and Bax. No in vivo binding to Bak was observed. A functionally inactive version of pepPuma (pepPumaDel), with a mutated BH3 domain where three amino acids, which are crucial to the protein–protein interaction, have been deleted (Figure 1a) did not bind to any of the Bcl-2 family members assayed. Surprisingly, none of the pepBH3 proteins showed intracellular interaction with Bcl-2, which was nonetheless present. We performed a co-IP experiment using an anti-Bad antibody, to test whether endogenous Bad was bound to endogenous Bcl-2 in staurosporine-treated A375 cells (Figure 2c). Although Bad successfully co-IP with Bcl-xL, no interaction with Bcl-2 was observed, consistent with the pepBad result (Figure 2b). In order to ascertain whether this surprising observation was specific to A375 cells, two separate experiments were carried out. First, Hela cells, transfected with pepBim and the negative control pepPumaDel, respectively, were submitted to a co-IP experiment (Figure 2b). Again no interaction between pepBim and endogenous Bcl-2 was detected. Second, the binding of endogenous full-length Bad to endogenous Bcl-2 was tested in the non-cancerous cell line human embryonic kidney 293T cells (Figure 2c). As for the A375 cells, an anti-Bad antibody column was used to establish whether Bcl-2 IP with Bad in apoptosing cells and again the result was negative.

As all the experiments so far had looked at endogenous levels of Bcl-2, we decided to test the interaction with pepBim when Bcl-2 was overexpressed. To that end, a mammalian expression vector carrying Flag-tagged Bcl-2 was transiently transfected into A375 cells carrying pepBim and pepPumaDel, respectively. Following transfection with Bcl-2 and induction of peptide aptamer expression, the cells were lysed (4 h after addition of 5 μM Shield) and co-IP carried out using an anti-Stefin A antibody (Figure 2b, bottom panel). This time efficient binding of pepBim to Bcl-2 could be detected, while no interaction with pepPumaDel occurred.

Next, cleavage of PARP was monitored to test whether pepBH3 proteins are capable of inducing apoptosis (Figure 3a). Expression of either pepPuma or pepBim, but not of pepBad or pepNoxa, results in efficient PARP cleavage. This is consistent with the ability of pepPuma and pepBim to bind to Bcl-xL and Mcl-1 (Figure 2b) and with the previously

**Figure 2** PepBH3 expression and identification of their binding partners. (a) Following induction, pepBH3 expression is detectable after 30 min. (b) Co-IP of pro- and anti-apoptotic Bcl-2 proteins with pepBH3 following their expression in A375 (top panel) and Hela cells (middle panel). There is no interaction between Bcl-2 and pepBim in A375 or Hela cells, however, when Bcl-2 is overexpressed (in A375) binding to pepBim is significant (bottom panel, underneath dashed line). (c) IP of endogenous full-length Bad from A375 cell lysate reveals binding of Bcl-xL, but not Bcl-2 (top panel). Equally in 293T cells, the Bad:Bcl-2 interaction does not occur (bottom panel). PARP cleavage indicates that apoptosis was induced successfully in these cells.
downregulation of anti-apoptotic proteins starts after 2–4 h.

Cleavage of PARP is detectable within 30 min of induction with Shield in cells expressing pepPuma or pepBim, and a significant decrease in the levels of the anti-apoptotic proteins Mcl-1, Bcl-2 and Bcl-xL are observed at 4 h.

**PepBim/Puma weakly localise to the mitochondria, but strongly induce apoptosis.** A cell viability assay following induction of pepBim/Puma expression in A375 cells with increasing amounts of Shield reveals that these peptide aptamers efficiently cause cell death. Induction of protein expression with 5 μM shield leads to a reduction in cell viability by 80% (Figure 4a). PepPumaDel does not affect cell viability (Figure 4b). To study the cellular localisation of the peptide aptamers immunocytochemistry (ICC) was used to probe for pepBim, pepPuma and empty SQT in Hela cells expressing these peptide aptamers (Figure 4c). SQT is expressed at high levels in the cells and displays a diffuse staining pattern, distributed evenly across all cellular compartments. PepPuma and pepBim show increased localisation to areas rich in mitochondria, especially surrounding the nucleus, however, staining remains rather diffuse and exhibits variability across different cells. These observations are consistent with the fact that pepBim/Puma lack the C-terminal hydrophobic domains of the full-length proteins, which have previously been shown to be essential for mitochondrial localisation. We conclude that the BH3 helix is sufficient for induction of apoptosis even in the absence of a mitochondrial-targeting sequence.

In order to verify that the BH3 presenting peptide aptamers cause programmed cell death via initiation of cytochrome c release from the mitochondrion, a Hela cell line, which stably expresses a green fluorescent protein (GFP)–cytochrome c fusion protein, was used (Figure 5). In a healthy cell, the mitochondria are visible as distinct features under the confocal microscope (Figure 5a). Upon expression of pepPuma, cytochrome c is released efficiently starting in some cells from 30 to 60 min following induction. Visually this can be registered by the dissolution of the distinct mitochondrial illumination pattern, which leads to a diffuse staining in the cytoplasm of these cells (Figures 5a and b).

**PepNoxa and pepBad do not individually induce apoptosis.** Expression of pepNoxa or pepBad did not lead to cleavage of PARP in A375 cells (Figure 3), indicating that apoptosis was not induced by these anchored BH3 helices. A similar observation has been made with the full-length proteins, and was explained by the fact that Noxa and Bad neutralise only a subset of anti-apoptotic Bcl-2 proteins. To test pepNoxa’s apoptotic potential, a U266 myeloma cell line was used, whose survival is dependent on Mcl-1. However, expression of pepNoxa in this cell line was not sufficient to drive apoptosis, as viability was unaffected and PARP was not cleaved (Figure 6a). It has previously been shown that the full-length Noxa protein possesses a C-terminal mitochondrial-targeting domain, whose presence is required for induction of apoptosis. In order to test whether targeting pepNoxa to the mitochondrion might result in induction of cell death, Bim’s membrane-targeting domain (Bim hydrophobic region, BHR) was introduced into the C-terminus of pepNoxa. Expression of the pepNoxa-BHR protein in U266 cells led to a clear reduction in cell numbers compared with SQT or pepNoxa alone (Figure 6a). Fluorescent microscopy in Hela cells confirmed that pepNoxa-BHR localises to the mitochondria, in contrast to pepNoxa, which exhibits a diffuse staining pattern (Figure 6b). Co-IP using an anti-Stefin A antibody to capture pepNoxa revealed that inclusion of the BHR significantly increases the formation of the pepNoxa:Mcl-1 complex (Figure 6c). Finally, as pepBad is predicted to mimic the effects of the chemical Bcl-2 inhibitor ABT-737 and synergise with the proteasomal inhibitor MG-132, the ability of pepBad to promote apoptosis in A375 cells treated with MG-132 was tested (Figure 6d). The strongest effect was measured at a concentration of 100 nM of MG-132, where expression of pepBad leads to approximately 50% reduced cell viability compared with the empty SQT scaffold. As with full-length Bad, which has no known membrane-targeting sequence, pepBad’s localisation pattern in Hela cells is diffused (Figure 6b).

**Discussion**

The study of BH3 helices has value in basic research to help further elucidate the role of these domains in apoptosis and the regulation of mitochondrial fission, and in translational research to inform drug design. However, in some cases the BH3 helix alone is not sufficient to mediate biological function,
and further tools are required. Here the use of the SQT peptide aptamer scaffold to present BH3 peptides has been successfully validated.

**At endogenous levels Bcl-2 does not interact with the BH3-only proteins.** The binding of the BH3 helices to members of the Bcl-2 family has been thoroughly investigated over the past 10 years. Chen *et al.* have systematically tested the binding of several BH3 peptides to the five anti-apoptotic Bcl-2 proteins in vitro, as well as the interaction between full-length BH3-only proteins and a subset of the anti-apoptotic proteins following their overexpression in cells. Their findings match our observations to a large extent, with Puma and Bim binding to both endogenous Bcl-xL and Mcl-1 and Bad binding strongly to Bcl-xL but not Mcl-1 whereas Noxa was selective for Mcl-1. Interestingly, however, endogenous Bcl-2 did not co-IP with any of the pepBH3 (in A375 or Hela cells), or with endogenous, full-length Bad (in A375 or 293T cells). In the scientific literature there are, to our knowledge, only a few instances of endogenous Bcl-2 co-IP, and in those studies the cell types used are almost exclusively either mature or immature T cells or B cells; all cell types which were not investigated in this study. The vast bulk of the

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**Figure 4** Cell viability and localisation of pepPuma/Bim in cells. (a) PepPuma/Bim expression in A375 cells reduces cell viability by approximately 80%. The western blot shows levels of SQT with increasing amounts of Shield (error bars are ± S.D.; n = 3). (b) PepPumaDel does not cause cell death (error bars are ± S.D.; n = 3). (c) Immunofluorescence of Hela cells expressing SQT reveals no specific localisation pattern. PepBim or pepPuma (green) colocalise with mitotracker (red) in some but not all cells. DAPI-stained nuclei are shown in blue.
literature has only reported positive BH3:Bcl-2 interactions in vitro or with overexpressed Bcl-2 inside cells. The latter observation could be successfully reproduced in this study. Notable in this context is the fact that several haematological malignancies are driven by translocation of the bcl-2 gene and consequent overexpression of the protein. Furthermore, there is some evidence that Bcl-2 is a much weaker pro-survival factor than, for instance, Bcl-xL and work with mouse embryonic fibroblasts has revealed that sequestration of Bcl-xL and Mcl-1 is sufficient to induce apoptosis. Thus, we hypothesise that, in most cell types, Bcl-2 might only be a relevant anti-apoptotic player and BH3-only protein inhibitor in the overexpressed state. Evidence corroborating our findings comes from Deng et al. who tested Bcl-2:Bim interaction via IP in 18 different diffuse large B-cell lymphoma cell lines. Importantly, they find that significant Bcl-2:Bim complex formation is generally only detectable in those cell lines carrying the t(14;18) translocation, where Bcl-2 levels are high. Recently, Plötz et al. co-IP BimL with endogenous Bcl-2 in Mel-2a melanoma cells, a cell line known to overexpress Bcl-2, particularly compared with A375, the melanoma cell line used in this study.

Only the anchored Bim BH3 helix interacts with Bax in A375 cells. The interaction of the BH3-only proteins with the anti-apoptotic Bcl-2 members serves to antagonise their function. There is, however, mounting evidence that Bax and Bak are directly activated by some of the BH3-only proteins. In this study, only the anchored Bim α-helix was found to bind to Bax in cells. This observation is consistent with that by Mérino et al. who report binding of Bax to the Bim isoform BimS. Replacement of the Bim BH3 peptide within the Bim backbone with that of Bad, Noxa or Puma led to no interaction. A different study, investigating binding to both Bax and Bak, found that BimS IP with Bax but found no interaction between Bak and the BH3-only proteins, consistent with our results. Both studies have also looked at the binding affinity between Bax and the Bim BH3 peptide biophysically, and only low (μM) affinities were measured. Here we show that an anchored Bim BH3 peptide is sufficient to interact with Bax in cells. The efficiency of the co-IP, which is comparable to that of pepBim and Bcl-xL, suggests that the interaction in vivo is robust. Furthermore, Puma has previously been shown to promote oligomerisation and activation of Bax and Bak followed by cytochrome c release from the mitochondria and a bacterial two-hybrid assay has identified Puma as well as its BH3 peptide alone as interactors of Bax. In this study, however, no binding of the constrained Puma helix to either Bax or Bak was detected. Others using IP assays have also found that full-length Puma does not bind to the pro-apoptotic effector proteins in lysed cells.

The BH3 domain of Puma and Bim is sufficient to induce apoptosis; a membrane-targeting sequence is not required. One of the advantages of the anchored peptide system described here is that it allows the study of the BH3 peptides in a neutral and structurally stable scaffold outside the context of their native proteins. Previously, Weber et al. have shown that a truncated BimS lacking the C-terminal hydrophobic region was incapable of localising to the mitochondria and could not induce apoptosis. Equally a truncated Puma, with an intact BH3 domain but no C-terminus failed to target the mitochondria and kill cells.

Figure 5 PepPuma expression results in cytochrome c release from the mitochondria. (a) Shown are three time points in the process of apoptosis. In the left-hand panel, both cells in the main field of view have their mitochondria intact. In the central panel, the right-hand cell is starting to undergo apoptosis with mitochondrial membrane permeabilisation leading to the GFP signal becoming more diffuse. In the right-hand panel, the right-hand cell has completely lost mitochondrial integrity and GFP–cytochrome c has translocated to the nucleus. (b) White arrows indicate cells before (left panel) and after (right panel) mitochondrial permeabilisation.
Here we show that the anchored BH3 helices of both Puma and Bim exhibit a strong apoptotic effect in A375 cells. In Hela cells, pepPuma powerfully induces cytochrome c release (pepBim was not tested). Although pepBim and pepPuma localise only poorly to the mitochondria, probably because the hydrophobic C-terminus is not present, these proteins do show increased accumulation around mitochondria compared with SQT alone. These findings thus prove that the Puma/Bim BH3 helix alone is enough to cause apoptosis even in the absence of efficient membrane targeting. In support of these results, several studies have successfully induced apoptosis in cancer cells by using a cell-permeable stapled BH3 peptides. Furthermore, previous studies have shown that the Bim helix colocalises very specifically with mitotracker (red). pepNoxa-BHR is expressed at lower levels, but purifies with a larger amount of Mcl-1 than pepNoxa lacking a BHR. pepBad and MG-132 synergise to reduce viability in A375 cells (most prominently at 100 nM MG-132; error bars are ± S.D.; n = 3).

**Conclusion.** The anchored helix approach to studying the BH3 peptides constitutes a way of informing the growing body of knowledge about the roles of these proteins in apoptosis and cancer. The results suggest that the potential implications for the design of Mcl-1 (Noxa’s target) inhibitors to treat malignancies such as chronic lymphocytic leukaemia and multiple myeloma, which have been shown to be dependent on Mcl-1 for survival, are promising. PepBad, on the other hand, showed apoptotic potential in the absence of a membrane-targeting domain consistent with the fact that the natural full-length protein lacks such a feature. Consequently, the results support the idea that the potency of the BH3-only Bcl-2 proteins is not only determined by their affinity for the anti-apoptotic proteins, but is also regulated by their intracellular localisation.

**Figure 6** Apoptotic potential and cellular localisation of pepNoxa and pepBad. (a) Expression of pepNoxa in U266 cells has no effect on cell viability and does not cause PARP cleavage. PepNoxa with the BHR domain efficiently induces cell death and leads to PARP deavage (error bars are ± S.D.; n = 3). The inducible expression system is leaky in this cell type and thus pepNoxa-BHR expression occurs even in the absence of shield. The set of columns labelled ‘U266’ denotes viability measurements of untransfected cells (which are not 100% viable). PepBH3 degradation products are indicated by an asterisk (*). (b) Immunofluorescence analysis of Hela cells expressing pepNoxa or pepBad (green) reveals no mitochondrial (red) localisation. PepNoxa-BHR colocalises very specifically with mitotracker (red). (c) Co-IP experiment using an anti-Stefin A antibody and lysate from cells expressing pepNoxa or pepNoxa-BHR. PepNoxa-BHR is expressed at lower levels, but purifies with a larger amount of Mcl-1 than pepNoxa lacking a BHR. (d) PepBad and MG-132 synergise to reduce viability in A375 cells (most prominently at 100 nM MG-132; error bars are ± S.D.; n = 3).
field of potential therapeutics that are the BH3 helix mimetics \(^{18,19,52–54}\) and stapled helices, \(^{37,55,56}\) as it presents a neutral and robust platform to study the intracellular effects of the BH3 domains outside the context of the native protein backbone. Other methods, such as using Bim\(_m\) as a common backbone to study the BH3 helices by replacing the Bim BH3 sequence with that of the BH3-only protein to be studied have been used successfully in the past.\(^{12,35}\) This approach, however, might not always reflect the true character of the BH3 domain under investigation as the BHR targets the protein to intracellular membranes very efficiently. Another important finding reported here is the apparent inability of Bcl-2 to interact with the pro-apoptotic BH3-only proteins, when the former is only present at endogenous levels in the cell. Conversely, when overexpressed, binding occurs readily. This suggests that Bcl-2’s role in counteracting apoptosis might only be significant in specific pathological situations where the protein’s expression levels are unnaturally high (e.g., following genetic translocation). Future studies should prioritise the verification of this hypothesis.

### Materials and Methods

**DNA manipulation and cloning.** Double-stranded DNA cassettes encoding each BH3 peptide (Figure 1a) or the BHR flanked by AvrII or XhoI restriction enzyme site, respectively, were made by annealing oligonucleotides (Supplementary Table S1). For bacterial expression, restriction enzyme digested dsDNA cassettes were ligated into the appropriate restriction sites of the SQT scaffold-encoding ORF in pET30a(+)\(^{35}\). For viral transfection, each construct was amplified by polymerase chain reaction (PCR) from pET30a(+)\(^{35}\) using SQT-specific primers (F 5′-GGGCGGCGCATCTCTAGATCAGGCCG-3′, R 5′-GAGAGGG GGCGCCATGCTAAAAAGCCGCGACGCTG-3′). PCR products were cloned into the Retro-X ProtoTuner vector using In-Fusion cloning and sequenced, as per manufacturer’s instructions (Clontech, St Germain-en-Laye, France).

**PepBH3 expression in E. coli and purification.** pET30a(+)\(^{35}\) expression vectors containing the pepBH3 ORF were transformed into BL21 (DE3) pLyS competent cells (Stratagene, Santa Clara, CA, USA) and expressed under induction with IPTG for 3 h at 37 °C. Cells were harvested, lysed using a sonicator, and the recombinant protein purified using Ni-NTA sepharose beads (Novagen, Billerica, MA, USA), as described previously.\(^{37}\)

**Production of retrovirus and transduction of cells.** Constructs were transfected into Phoenix A packaging cells (ATCC, Manassas, VA, USA), using siPORT XP-1 transfection agent (Life Technologies, Carlsbad, CA, USA). After 48 h, the medium was harvested, 0.4 μm filtered and mixed in equal amounts with fresh medium containing 8 μg/ml of polybrene (Sigma, Gillingham, UK). Cells were incubated with retroviral supernatants for 8 h. At 48 h after transfection, cells were transferred into selection medium containing puromycin.

**Transient transfection of A375 cells.** A pCMV-Tag2B vector carrying Bcl-2 with an N-terminal Flag-tag was ordered from Addgene (Cambridge, MA, USA). The transfection mix was assembled as follows: 3:3 μg of high-quality plasmid DNA (midprep) per six-well plate well is diluted in sterile water to a concentration of 20 ng/μl. Then FuGenDE transfection reagent (Promega, Southampton, UK), which has been warmed to room temperature (RT), is added directly to the solution at 10 μl/3.3 μg DNA. This is followed by brief vortexing and incubation at RT for 5–10 min. The mix is then added to cells, which are approximately 80% confluent, and are grown in complete DMEM Glutamax medium (Life Technologies, Paisley, UK), with 3 ml of the medium per six-well plate well. The transfected cells were harvested after 24 h for downstream applications.

**Far-UV CD spectroscopy.** CD spectra were measured on a Jasco J715 spectropolarimeter (Jasco, Easton, MD, USA) as described previously.\(^{37}\) Briefly, pepBH3 proteins were transferred into 50 mM sodium phosphate buffer (pH 7.4) and their CD spectra recorded from 200 to 260 nm. The spectrum for buffer alone was subtracted from the measurements and the data analysed and visualised using Microsoft Excel.

**IP experiments.** IP experiments were carried out using the Crosslinking Immunoprecipitation kit (Pierce, Rockford, IL, USA) according to the manufacturer’s instructions. Briefly, cells expressing the desired protein were lysed using the kit’s IP lysis/wash buffer (containing a non-ionic detergent) and the soluble fraction applied to agarose beads carrying the appropriate antibody (α-Bad, Abcam, Cambridge, UK; α-Stefin A, R&D Systems (Minneapolis, MN, USA); α-Flag-Tag, Sigma). Lysates were incubated overnight at 4 °C and the beads were then washed four times and then stored at ~80 °C until analysis by western blotting.

**Western blotting.** Western blotting was preceded by sodium dodecyl sulphate polyacrylamide gel electrophoresis and transfer onto nitrocellulose or PVDF membrane. The membrane was blocked in 3% milk for 1 h at RT followed by incubation with antibody at 4 °C overnight. Antibodies used were α-Stefin A (1:3000; R&D Systems), α-Bad (1:10 000; AbD Serotec, Kidlington, UK), α-Mcl-1, α-Bcl-2, α-Bcl-xL, α-Bad, α-Bak and α-Bax (all 1:1000; all from Cell Signalling Technology, Hitchin, UK) and α-PARP (1:1000; BD Biosciences, Oxford, UK). HRP-conjugated secondary antibodies were α-mouse, α-rabbit, α-goat (all 1:12 000; all from Jackson ImmunoResearch, West Grove, PA, USA) and were applied for 1 h at RT. For detection, the membrane was incubated with ECL reagent (Pierce) and exposed to photographic film.

**Immunocytochemistry.** For ICC, Hela cells were cultured on glass cover slips. Mitotracker red CMXRos (25 nM; Life Technologies, Carlsbad, CA, USA) was added to the culture medium 30 min before fixing in 4% paraformaldehyde. Cells were permeabilised in 2% Triton-X100 for 5 min, washed in PBS and blocked in 2% milk for 30 min. Subsequently, the anti-Stefin A antibody was applied (1:100; R&D Systems) for 2 h at RT followed by incubation with Alexa Fluor 488 labelled secondary antibody (1:200; Life Technologies, Carlsbad, CA, USA) and 4,6-diamidino-2-phenylindole (DAPI) for 1 h. Before fluorescence microscopy, the cover slips were mounted onto glass slides using ProLong Gold antifade reagent (Life Technologies, Carlsbad, CA, USA).

**Monitoring cell viability and cytochrome c release.** Viability and apoptosis assays were performed using the Guava EasyCyte System (Millipore, Billerica, MA, USA), according to the manufacturer’s instructions. Hela cells, which stably express GFP–cytochrome c (a kind gift from Douglas Green, St. Jude Children’s Research Hospital, Memphis, TN, USA), were transfected with pepPuma as described above. One hour after addition of Shield, the cells were monitored under green fluorescence and micrographs taken every 5 min for 2 h.

### Conflict of Interest

The authors declare no conflict of interest.

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