

# In Silico and in Vitro Elucidation of BH3 Binding Specificity toward Bcl-2

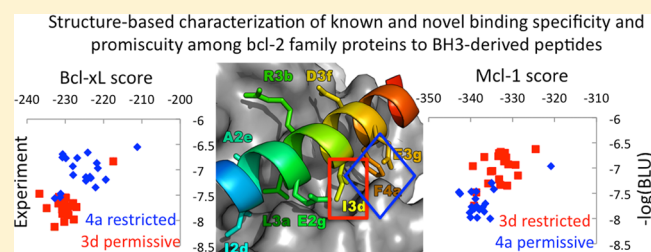
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## S Supporting Information

**ABSTRACT:** Interactions between Bcl-2-like proteins and BH3 domains play a key role in the regulation of apoptosis. Despite the overall structural similarity of their interaction with helical BH3 domains, Bcl-2-like proteins exhibit an intricate spectrum of binding specificities whose underlying basis is not well understood. Here, we characterize these interactions using Rosetta FlexPepBind, a protocol for the prediction of peptide binding specificity that evaluates the binding potential of different peptides based on structural models of the corresponding peptide–receptor complexes. For two prominent players, Bcl-xL and Mcl-1, we obtain good agreement with a large set of experimental SPOT array measurements and recapitulate the binding specificity of peptides derived by yeast display in a previous study. We extend our approach to a third member of this family, Bcl-2: we test our blind prediction of the binding of 180 BIM-derived peptides with a corresponding experimental SPOT array. Both prediction and experiment reveal a Bcl-2 binding specificity pattern that resembles that of Bcl-xL. Finally, we extend this application to accurately predict the specificity pattern of additional human BH3-only derived peptides. This study characterizes the distinct patterns of binding specificity of BH3-only derived peptides for the Bcl-2 like proteins Bcl-xL, Mcl-1, and Bcl-2 and provides insight into the structural basis of determinants of specificity.



The Bcl-2 protein family is a key regulator of programmed cell death, tumorigenesis, and cellular responses to anticancer therapy. Proteins of this family can be grouped into two classes: antiapoptotic proteins that suppress cell death (such as Bcl-2, Bcl-xL, Bcl-w, Mcl-1) and pro-apoptotic proteins that promote cell death. The latter can be further classified based on their conserved domain structure into proteins containing multiple Bcl-2 homology (BH) domains (such as Bak, Bax, Bok) and proteins that contain only the BH3 domain (Bad, Bik, Bid, Bim, Puma).<sup>1,2</sup>

BH3-only proteins promote apoptosis both by the activation of Bak and Bax and by antagonizing pro-survival Bcl-2 family proteins.<sup>3–5</sup> The interaction of the amphipathic helical BH3 domain of a pro-apoptotic protein with the hydrophobic groove created by the conserved BH1, BH2, and BH3 domains of an antiapoptotic family protein is structurally conserved and well characterized.<sup>6–9</sup>

Overexpression of antiapoptotic Bcl-2 family members has been associated with cancers as well as with chemotherapy resistance, and molecules targeting these interactions are consequently pursued for the development of pharmaceuticals.<sup>10–12</sup> As an example, ABT-737 is a selective and potent inhibitor of Bcl-2, Bcl-w, and Bcl-xL,<sup>13</sup> while gx15-070 is a pan-specific Bcl-2 family inhibitor.<sup>14</sup> BH3 binding specificity to different Bcl-2 family members has also been the subject of intensive biophysical and cellular biology research. Structural

studies have shed light on the BH3 binding determinants<sup>6,7,9,15,16</sup> while other studies scanned point mutants<sup>17–22</sup> and random peptide libraries.<sup>21–23</sup> Despite all these efforts, the binding specificity of Bcl-2 family members has still not yet been fully elucidated.

Computational attempts to map peptide-binding specificity are usually restricted to one specific type of peptide-binding domain and can be roughly divided into sequence-based (e.g. refs 24–27) and structure-based approaches. The latter may use a fixed backbone (e.g. refs 28 and 29) or incorporate backbone flexibility (at least on the peptide side; e.g. refs 30–35). Incorporation of backbone conformational flexibility has generally improved computer-aided design of functional protein interactions as well as structure-based prediction of peptide–protein and protein–protein interaction specificity.<sup>36,37</sup> Several recent studies have advanced the computational structural characterization<sup>38–42</sup> and high-resolution modeling<sup>43–48</sup> of peptide–protein interactions. Consequently, these new tools now provide a platform for the prediction of peptide binding specificity.

What determines the ability of peptides to bind to proteins? Compared to regular protein–protein interactions, peptide–

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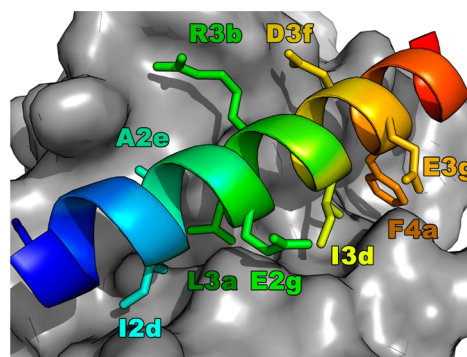
mediated interactions are challenged by the small interface that provides less binding enthalpy as well as by the fact that the peptide loses a considerable amount of entropy upon binding, since in general it lacks a defined structure in its free state. Several strategies are used to address those challenges.<sup>38</sup> Overall entropy loss upon binding is reduced thanks to small conformational changes of the receptor that adopts the bound conformation prior to interacting with the peptide partner. On the peptide side, a considerable fraction of peptide-mediated interactions involve helical peptides that at least partially adopt the helical conformation in the free state. It therefore does not come as a surprise that a predominant part of successful designs of inhibitory peptides involves helical peptides (e.g., refs 49 and 50). The interaction between Bcl-2-like proteins and BH3 domains is an example of a helix–receptor interaction that represents an ideal starting point to characterize high-affinity peptide–protein interactions. Thus, this investigation can lead to helical inhibitory peptides that compete at protein interfaces<sup>51,52</sup> (and can be further optimized, e.g., with staples<sup>53,54</sup>).

A recent review has described the various structural and sequence features that influence and determine binding specificity.<sup>55</sup> A wide variety of different protein–protein interactions have been observed. Of particular importance to the complexity of the interaction network in a cell are interaction types that have been duplicated and are multiply reused. A range of different specificities can be observed for homologous protein pairs: certain proteins will bind only to one specific protein partner, while others show less specificity and bind promiscuously to a range of different homologues. The wide variability and conservation observed in different systems have revealed both robustness and sensitivity: binding can be obtained in different ways for the same interface, while single mutations can strongly affect binding. These differences will have profound functional consequences on the cell. The interaction between Bcl-2-like proteins and BH3-derived peptides represents a model case to elucidate the structural basis of such intricate specificity effects.

In this study we have applied our previously developed FlexPepBind protocol that predicts peptide-binding specificity (e.g., ref 35) to recapitulate the BH3 peptide binding preferences to Bcl-2 protein family members. After an initial calibration and application of FlexPepBind to Mcl-1 and Bcl-xL based on *in vitro* SPOT array experiments<sup>56</sup> for a set of BIM-derived peptides,<sup>21</sup> we used this protocol to blindly predict the corresponding specificity profile of Bcl-2. Our prediction was shown to agree well with an additional, independently performed *in vitro* SPOT array experiment: Both prediction and experiment indicate that Bcl-2 displays a BH3 binding profile that is highly similar to the profile previously reported for Bcl-xL. Finally, we used this approach to recapitulate the distinct patterns of binding specificity of a set of BH3-like peptides to Bcl-xL and Mcl-1 and the binding of several native human BH3 peptides to all three receptors. The successful performance of our protocol on several receptors of this system bodes well for future application and suggests that, after proper calibration, it is general enough for a large variety of peptide–protein interactions.

## MATERIALS AND METHODS

**Data Sets.** All data sets of peptide binding assays Bcl-xL and Mcl-1 were taken from Dutta et al.,<sup>21</sup> and the numbering of the residues is according to the definition therein (see also Figure



**Figure 1.** BIM peptide in complex with Mcl1. The amphiphilic BIM BH3-only helical peptide (cartoon representation, rainbow coloring) binds Mcl1 (surface representation, gray) in a hydrophobic groove. Ten peptide interface positions that were mutated in the different data sets used in this study are indicated in stick representation, accompanied by their position nomenclature.

1) *SPOT arrays*: (a) TRAIN1: a SPOT array displaying all possible combinations of selected Bim-BH3 mutants at the following positions: 2d: Ile/Ala/Phe; 3a: Leu/Ile/Phe/Ala; 3b: Arg/Asp; 3d: Ile/Phe/Asp/Asn/Ala; 4a: Phe/Val/Asn ( $n = 360$  sequences); (b) TRAIN2: a SPOT array displaying Bim-BH3 single mutant variants, in which 10 interface positions of the peptide (2d, 2e, 2g, 3a, 3b, 3d, 3e, 3f, 3g, and 4a) were substituted to all amino acids (excluding Met and Cys; the native Bim-BH3 sequence was probed 20 times but only used once for calculation of the correlation) ( $n = 181$ ). This set was probed by either 1  $\mu$ M or 100 nM of the receptor. (2) *Yeast surface display experiments*: (a) YEAST-X: sequences from a yeast surface display experiment that were shown to bind to Bcl-xL in preference to Mcl-1 ( $n = 40$ ). (b) YEAST-M: sequences from a yeast surface display experiment that were shown to bind to Mcl-1 in preference to Bcl-xL ( $n = 33$ ). (c) YEAST-BOTH: sequences from a yeast surface display experiment that were shown to bind both Bcl-xL and Mcl-1 ( $n = 17$ ).

**Modeling the Peptide–Receptor Complex. General Modeling Scheme.** Our scheme is based on the previously reported peptide-docking protocol Rosetta FlexPepDock.<sup>43</sup> For a given peptide, we thread the desired sequence onto the peptide in the native structure, while keeping the side chains of the receptor fixed. We then use Rosetta FlexPepDock in full-atom mode (i.e., without centroid preoptimization) to refine the structure of the complex with the threaded target peptide. (All of the peptide's side chains as well as the receptor's interface side chains are flexible.) We create 1000 models and score the sequence as the average score of the top 50 models (see below) to evaluate the binding ability of the specific peptide sequence.

**Modeling of Specific Protein–Peptide Complexes.** For each of the Bcl-xL, Mcl-1, and Bcl-2 receptors, we evaluated different available PDB structures to serve as templates to discriminate binders from nonbinders in the TRAIN1 set, using a quick protocol that involves only simple minimization of the template structure. (1) Bcl-xL: Two crystal structures of the Bcl-xL/Bim-BH3 peptide interaction are available (PDB id: 3io8, 3fdl). 3io8 (chains C and D) provided the best discrimination (see Supporting Information Table S1), and we therefore refer to 3io8:CD as the “native structure”. (2) Mcl-1: We evaluated a set of different PDB structures as templates (2nl9, 2pqk, 3d7v,

3io9, 3kj0, 3kj1, 3kj2, 3kz0). Among those, 2pqk provided the best discrimination of the TRAIN1 set. (3) Bcl-2: The receptor structure was taken from PDB 2xa0, and the BH3 BIM peptide template from PDB 2pqk.

**Score Function.** We use the standard Rosetta score function (score12) for the simulations, with two minor adjustments: (1) The penalty for the burial of a carboxyl oxygen atom is increased (the  $\Delta G^{\text{free}}$  parameter of the Lazaridis–Karplus solvation potential<sup>57</sup> was modified from  $-10$  to  $-13.5$  for this atom type). This was added to better account for a group of sequences that received poor experimental binding values in the TRAIN set but good peptide scores. Inspection of the structural models detected a buried Asp that explains the discrepancy. (2) A weak short-range Coulombic electrostatic energy term was added with weight 0.5 (as described in ref 58).

We assessed each model by its “weighted score”, which is the average of the Total score, Interface score, and Peptide score (where Total score is the overall Rosetta energy score for the complex, Interface score is the energy of pairwise interactions across the peptide–protein interface, and Peptide score is the sum of the Rosetta energy function over the peptide residues). This score was proven useful in previous modeling studies using FlexPepDock (e.g., ref 44).

**Estimation of Difference in SPOT Arrays.** To find the point mutations that differ the most between the Bcl-2 SPOT array experiment and prediction, the 200 scores (or  $-\log(\text{BLU})$ ) were normalized to the average score of 10 wild-type scores. The differences between the normalized scores were rank-ordered: the largest delta values indicate positions that differ most between the arrays.

**Experimental Measurement of Bcl-2 Binding Specificity with SPOT Arrays.** Human Bcl-2 protein and SPOT arrays were prepared and probed as previously described.<sup>21</sup> Briefly, Bcl-2 (residues 1–172) was expressed as a TEV-cleavable C-terminus fusion construct with His-tagged maltose binding protein (MBP). Purification was achieved using a Ni-NTA column followed by cleavage with TEV protease to yield N-terminally c-myc-tagged Bcl-2 product. A second Ni-NTA affinity step was used to capture His-tagged MBP and His-tagged TEV, and the Bcl-2 fraction was further purified using size exclusion chromatography. SPOT arrays were synthesized on activated nitrocellulose support using Fmoc protection/deprotection chemistry by an Intavis AutoSpot robot in the MIT Biopolymers Laboratory. The 26-residue peptides were synthesized with PEG3 (three ethylene glycol units) linkers at the carboxy terminus. The array consisted of single site substitutions of 10 interfacial positions of the wild-type BIM sequence (MRPEIWIAQELRRIGDEFNAYYARRV) to all amino acids with the exception of cysteine and methionine. Resulting BIM substitution arrays were probed with either 100 nM or 1  $\mu\text{M}$  Bcl-2 followed by detection with anti-c-myc-Cy3 antibody (Sigma-Aldrich C6594) as previously described.<sup>21</sup> Fluorescence intensity for the probed SPOT arrays was measured on a Typhoon 9400 (GE Healthcare), and the resulting images were analyzed with ImageQuant (GE Healthcare) without background correction. Data from the 100 nM experiments were used in this work. Raw signals were used without background correction, measured in BLU.

## RESULTS

We have previously developed a general framework for the prediction of binding specificity of flexible peptides to protein

receptors. In general, we model the structure of peptides with variable sequences to the target receptor using our high-resolution peptide docking protocol (Rosetta FlexPepDock<sup>43,44</sup>) and use the energy scores of the models to assess binding ability. We have applied this protocol to determine which peptides can bind to the enzyme farnesyltransferase (FTase) and consequently undergo farnesylation.<sup>35</sup> This protocol can potentially be applied to any new system after simple calibration on a set of sequences for which binding properties have experimentally been determined. Here, we first calibrate and test our protocol for the binding of BH3 derived helical peptides to Bcl-xL and Mcl-1 on different sets of experimental data reported by Dutta et al.<sup>21</sup> We then apply the protocol to the homologue Bcl-2 and validate our predictions with a SPOT array experiment. Finally, we successfully reproduce binding specificity profiles for these three Bcl-2-derived proteins for a range of different human BH3-derived peptides.

**1. Calibration of the Protocol.** Dutta et al. performed two different SPOT peptide array experiments to determine the binding of different peptides to Bcl-xL and Mcl-1.<sup>21</sup> TRAIN1 ( $n = 360$ ) is a combinatorial library of mutations at five positions of the BIM BH3 peptide. TRAIN2 ( $n = 181$ ) is a library of all possible point mutations at 10 different positions of the BIM BH3 peptide (see Figure 1 and Materials and Methods for detailed description of the sets; TRAIN2 was probed by either 1  $\mu\text{M}$  or 100 nM of the receptor). We modeled each of the sequences in these peptide libraries in complex with Bcl-xL and with Mcl-1. The binding energy of each of these models was then used to define their relative binding abilities to Bcl-xL and Mcl-1. We describe in the following how these sets were used to calibrate the sampling and scoring schemes of our protocol.

Several solved structures are available for both Bcl-xL and Mcl-1. To select the most suitable template for our protocol, we modeled the TRAIN1 set onto each of the different templates and minimized the threaded peptides in the context of the complex structure. To assess the performance of each template, we examined the Pearson correlation between five different scoring schemes (see Materials and Methods) and the experimental estimation of binding (represented as the logarithm of the SPOT binding intensity, BLU). The weighted score (“Wsc”), a scoring scheme including the total score of the complex while emphasizing the interactions across the interface and the internal energy of the peptide (see Materials and Methods), was shown useful in previous peptide modeling studies<sup>44</sup> and is indeed also adequate for both receptors in this case. In the following we report results obtained using this score. For Bcl-xL, the score of the minimized peptides with the template structure 3io8<sup>16</sup> in the Protein Data Bank (PDB<sup>59</sup>) showed better correlation ( $r = 0.54$ ) than the other available template, while for Mcl-1 among eight tested templates PDB 2pqk<sup>9</sup> resulted in the best correlation ( $r = 0.46$ ).

We next applied the full FlexPepBind protocol on these templates: As opposed to the fast and simple minimization-based protocol, we created for each peptide sequence 200 models using our flexible peptide docking protocol FlexPepDock and selected the top-scoring models as representatives for that sequence. This exhaustive protocol samples extensively the peptide flexibility within the binding pocket. Indeed, the top-scoring model of each peptide sequence improved the correlation ( $r = 0.62$  for both receptors). We note that results based on minimization of different templates correlate with the performance of these templates using the full FlexPepDock



protocol (see Supporting Information Table S1) and conclude that simple minimization is a good indicator for the initial selection of the best structural template.

Manual inspection of several false positive peptide sequences (i.e., sequences that in the experiment did not bind but were nevertheless predicted to bind well by our protocol) revealed a common I3dD mutation in which the aspartic acid was buried in a hydrophobic pocket (numbering of peptide positions according the scheme used in Dutta et al.;<sup>21</sup> see Figure 1). This indicates that the burial of charges and polar atoms may not be penalized strongly enough in the current default Rosetta energy function. We therefore modified the Rosetta solvation model parameters (that are based on the Lazaridis–Karplus solvent exclusion model<sup>57</sup>) to further penalize buried carboxyl groups (see Materials and Methods). Application of our protocol with this adapted energy function slightly improved the correlation to experiment (Bcl-xL  $r = 0.66$ ; Mcl-1  $r = 0.68$ ). Addition of a simple Coulombic electrostatic energy term<sup>58</sup> further improved performance for Bcl-xL ( $r = 0.76$ ), without any influence on the Mcl-1 predictions.

The final calibration of our protocol involved the investigation of two additional features that in previous applications improved predictions, namely the number of models created by the docking protocol and the number of top-scoring models used to evaluate the binding energies. Overall, marginal improvement could be observed when the number of models was increased (from  $n = 200$  to  $n = 1000$ ). Scoring based on averaging the top 10- or top 50-scoring models did not significantly improve performance.

We note that significant improvement was observed for the Bcl-xL predictions for some of the tested parameter sets, but at the expense of reduced performance on Mcl-1. Since we aim at one general protocol that can be applied to this whole family, we focused on a protocol that performs best for both Bcl-xL and Mcl-1 (see Table S1).

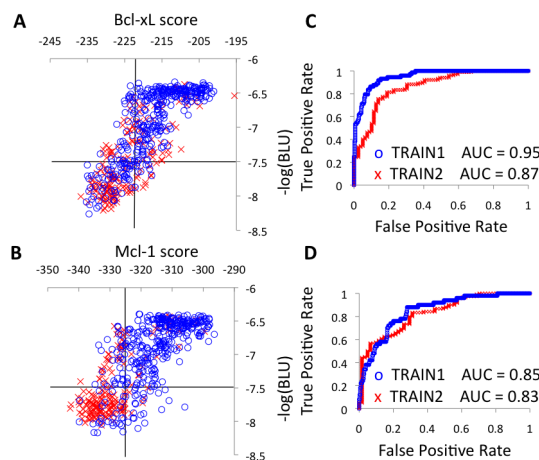
**Table 1. Pearson Correlation between Predicted and Experimental Binding Values of BIM-Derived Peptides to Bcl-xL and Mcl-1<sup>a</sup>**

	BIM BH3-derived peptide sets <sup>b</sup>		
	TRAIN1 ( $n = 360$ ; 100 nM)	TRAIN2 ( $n = 181$ ; 1 $\mu$ M)	TRAIN2 ( $n = 181$ ; 100 nM)
Bcl-xL	$r = 0.80$	$r = 0.73$	$r = 0.69$
Mcl-1	$r = 0.67$	$r = 0.71$	$r = 0.65$

<sup>a</sup>Experimental values were taken from SPOT array experiments.<sup>21</sup> Binding predictions were performed using the weighted score of the peptide (Wsc), according to the calibrated protocol described in the text (see Materials and Methods and Results for more details and Table S1 for alternative scoring schemes). <sup>b</sup>TRAIN1: combinatorial library of mutations at five positions. TRAIN2: 180 point mutations at 10 positions.<sup>21</sup> TRAIN2 array was probed by either 1  $\mu$ M or 100 nM of the receptor.

Overall, Table 1 and Figure 2A,B show a good agreement between our predicted values and the logarithm of the measured SPOT intensities, both for the TRAIN1 set and for the subsequently evaluated TRAIN2 set.

**2. Prediction of Binding for the TRAIN 1 and TRAIN 2 SPOT Arrays.** Examination of the TRAIN correlation plots revealed two distinct regions: a group of nonbinding sequences, which can be detected by high energy scores, and a group of



**Figure 2.** Predicted binding values for BIM-derived peptides to Bcl-xL and Mcl-1 are in good agreement with experimental results from SPOT arrays. (A, B) Correlation of calculated score (X-axis) to the experimental peptide binding signal (Y-axis;  $-\log(\text{SPOT intensity})$  measured in BLU) for (A) Bcl-xL and (B) Mcl-1, for TRAIN1 (blue circles) and TRAIN2 probed with 1  $\mu$ M receptor (red crosses) (see Materials and Methods for description; the experimental data for TRAIN2 are also shown in Figure S5A). The  $-7.5$   $-\log(\text{BLU})$  threshold (see Figure S1) and the selected score thresholds are indicated with lines. (C, D) ROC curves for the discrimination of binders and non binders (threshold for binding defined as  $-\log(\text{BLU}) = -7.5$ ) for (C) Bcl-xL and (D) Mcl-1. The high area under the curve (AUC) values demonstrate good discrimination between BH3 Bim-derived peptides that bind/do not bind to different Bcl-2 binding proteins.

binding sequences that correlate with our predicted score. This observation led us to inspect the performance of the protocol in a binary discrimination task. That is, we defined a threshold for the experimental binding value, below which a peptide is considered a “binder” and above a “nonbinder”. If one wishes to utilize this protocol to detect high-affinity binders, this threshold should be higher (strict), whereas if the objective is to detect any type of binding, this threshold should be lower (permissive). Given this binary labeling of the data points, we created receiver operating characteristic (ROC) curves and calculated the area under the curve (AUC), which indicates the discrimination ability of our scoring. For both receptors, and on both training sets, we obtained AUC values above 0.83 (see Figure 2C,D and Table 2). We report the results using an arbitrary threshold of  $-7.5$  (logarithm of the normalized fluorescence intensity in the SPOT experiment) to define “binders”. However, we note that the predictions would improve with increasing threshold stringency, at the expense of detecting fewer binders (see Figure S1). This indicates that our protocol is particularly suitable for the selection of high-affinity binding peptides.

We selected a score threshold of  $-223$  for Bcl-xL and  $-325$  for Mcl-1, corresponding to a false positive rate (FPR) of 10% on the TRAIN1 data set. These thresholds achieve a true positive rate (TPR) of 86% and 54% for Bcl-xL and Mcl-1, respectively, on the TRAIN1 set. On the TRAIN2 sets, the same threshold gives a much higher TPR for Mcl-1 (Bcl-xL 88%; Mcl-1 96%) together with a corresponding increase in FPR (Bcl-xL 32%; Mcl-1 59%). This could be attributed to different backgrounds of the SPOT arrays. It is also reasonable that the TRAIN1 and TRAIN2 sets contain different numbers of binders. TRAIN2 contains only single point mutations,

**Table 2. Identification of Binders and Nonbinders in the Training Sets<sup>a</sup>**

	threshold <sup>b</sup>	TRAIN1 (100 nM)					TRAIN2 (1 $\mu$ M)				
		AUC <sup>c</sup>	bind <sup>d</sup>	NON <sup>e</sup>	TPR (%)	FPR (%)	AUC	bind	NON	TPR (%)	FPR (%)
Bcl-xL	−223	0.95	73	287	86	10	0.87	102	79	88	32
Mcl-1	−325	0.85	50	310	54	10	0.83	120	61	96	59

<sup>a</sup>Binders and nonbinders were defined using a threshold of  $-\log(\text{BLU}) = -7.5$ . <sup>b</sup>Score threshold set to achieve 10% FPR rate on TRAIN1 (see Figure 2). <sup>c</sup>Area under the ROC plot. <sup>d</sup>Number of “binding” peptides in the set (according to a  $-7.5 - \log(\text{BLU})$  threshold). <sup>e</sup>Number of “nonbinding” peptides in the set (according to a  $-7.5 - \log(\text{BLU})$  threshold).

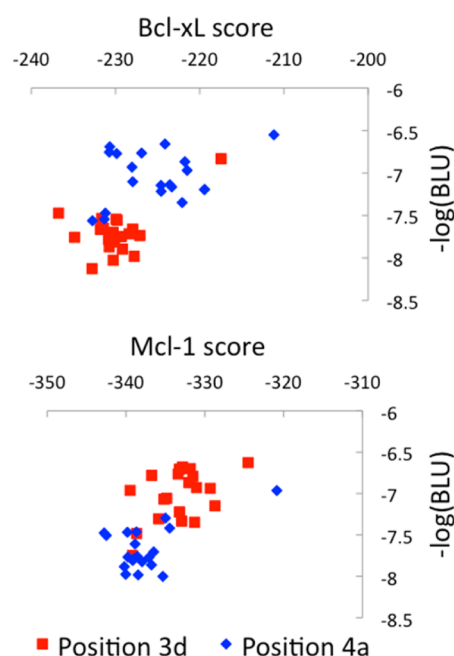
which are expected to perturb binding less than the combinations of mutations explored in TRAIN1 (see Materials and Methods for descriptions of the data sets).

Although we sometimes refer below to “binding” and “nonbinding” peptides, this is a system- and assay-specific designation, and the interpretation should be that these are higher-affinity/lower-affinity binding peptides. Further, numerical score thresholds should not be applied to predictions of peptides from different sequence regimes. Rather, the relative ranking of scores within data sets serves as a better indicator for comparison.

**3d and 4a Positions.** Of particular interest in the TRAIN2 set are point mutations in BIM positions 3d and 4a. Mcl-1 binds almost singularly to peptides with a native isoleucine at position 3d, and any point mutation abolished binding on the SPOT arrays (at least at 100 nM concentration). In turn, Bcl-xL is promiscuous at the 3d position and allows almost any amino acid point mutation. Position 4a shows the reverse behavior; i.e., many mutations can be introduced without impairing binding to Mcl-1, but binding to Bcl-xL is very specific. A plot of predicted scores for these point mutants largely recapitulates this behavior (see Figure 3).

**3. Characterization of Binding Specificity of Peptides Isolated from a Yeast Surface Display.** In addition to the SPOT arrays, Dutta et al. also report a third set of sequences of peptides that were recovered from a yeast surface display library.<sup>21</sup> This set contains peptides that bind specifically to Bcl-xL (YEAST-X;  $n = 40$ ), to Mcl-1 (YEAST-M;  $n = 33$ ), or to both (YEAST-BOTH;  $n = 17$ ). We used this set to test the ability of our computational protocol to characterize binding specificity. For each of the peptides, we calculated binding scores for both Bcl-xL and Mcl-1. Figure 4 shows that the Bcl-xL and Mcl-1 specific peptides are clearly separable using these two values. Our Bcl-xL score distinguishes Bcl-xL binders from nonbinders and detects most sequences that bind both proteins. However, FlexPepBind fails to differentiate Mcl-1 binders vs nonbinders using the Mcl-1 score. Mcl-1 specific peptides obtain a similar range of scores as the peptides that bind solely to Bcl-xL. This behavior was also observed by Dutta et al., who used a SPOT-derived PSSM based on point mutation data (TRAIN2 in this study) to predict the yeast data. The PSSM was subsequently improved by assessing an additional set of mutants (TRAIN1 in this study) to create a modified model with better discrimination (see Figure 6E in ref 21).

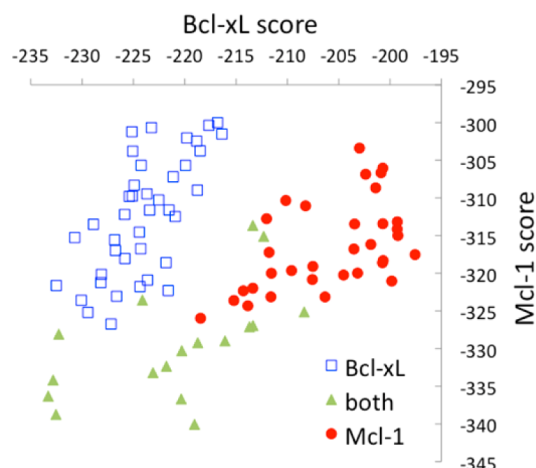
Examination of our structural models suggests that the problem can be traced back to a set of peptides that contain a mutation at position 3b. According to the TRAIN1 set, most peptides containing an R3bD mutation are nonbinders, but those that do bind are rescued by a F4aV mutation. Indeed, many of the YEAST-M sequences contain such an R3b to D/E mutation (13/33) or R3b to G mutation (9/33). Fourteen of these 22 are combined with the F4aV mutation (actually, most



**Figure 3.** Binding specificity and promiscuity at positions 3d and 4a. Binding score estimation for peptide binding to Bcl-xL (top) and Mcl-1 (bottom) recapitulates distinct specificity-determining residues identified using SPOT experiments.<sup>21</sup> Bcl-xL binding peptides show a restricted range of amino acids at position 4a (blue diamonds), while many residues are tolerated at position 3d (red squares). Mcl-1 binding peptides in contrast show specific preferences for position 3d, but little discrimination at position 4a.

YEAST-M sequences, 19/33, contain valine at position 4a). Structurally, these positions are not spatially adjacent, and coupling between these positions is thus not expected (see Figure S2). It seems that our protocol fails to properly weight the contribution of the 4a valine and therefore predicts the wrong outcome for these sequences. In contrast, the experiment-based PSSM approach overcomes this failure by using data from TRAIN1, which captures the valine preference (see Figure 6F in ref 21).

**4. Characterization of Bcl-2 Binding Specificity: SPOT Array for Bcl-2.** In order to evaluate the general ability of our protocol to characterize BH3 binding proteins, we extended our applications to another member of this family: Bcl-2. We applied the same protocol to a crystal structure of Bcl-2 in complex with a BH3 peptide (PDB: 2xa0<sup>15</sup>) and constructed an *in silico* SPOT array analogue to TRAIN2. Independently of the blind prediction, a SPOT array displaying 26-residue Bim-BH3 variants representing the TRAIN2 set was constructed using solid-phase synthesis in the same way as described by Dutta et al.<sup>21</sup> and probed with 100 nM Bcl-2. Figure 5 displays the predicted Bcl-2 spot array side by side with the experimental results. Overall, there is good agreement between the predicted

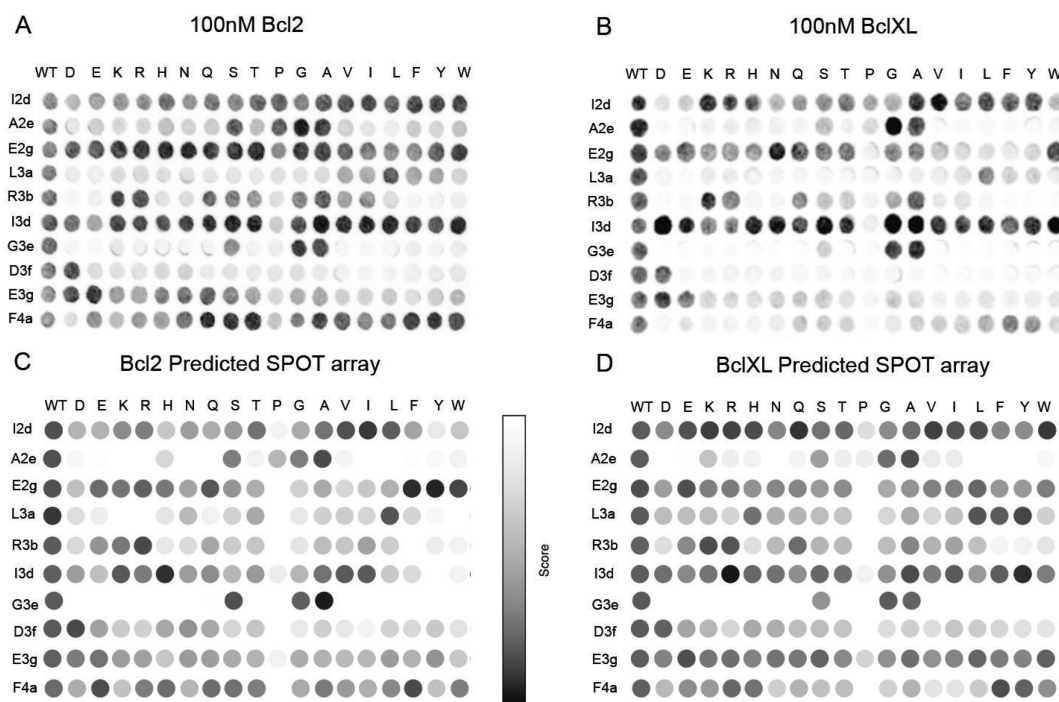


**Figure 4.** Specificity prediction for yeast display-derived peptides. FlexPepBind is able to discriminate between Bcl-xL binders/nonbinders (Bcl-xL specific binders all obtain good scores for binding to Bcl-xL but worse scores for binding to Mcl-1; blue squares in upper left quadrant) as well as detect sequences that bind both proteins (peptides that bind to both proteins mostly obtain good scores for both; green triangles in lower left quadrant). However, it performs badly in detecting the Mcl-1 specific peptides, which all obtain unfavorable scores both for Bcl-xL and Mcl-1 binding (red circles in upper right quadrant—should be located in lower right quadrant).

binding and experimental results ( $r = 0.65$ , which is similar to the values obtained for both Mcl-1 and Bcl-xL for this data set; see Figure S3), indicating that indeed this approach might be extensible and general for additional receptors.

Both computational and experimental assessments indicated that the overall specificity profile emerging from this set for Bcl-2/BIM interaction is similar to that of Bcl-xL ( $r = 0.88$  between the two experiments). Nevertheless, some differences can be identified. Most notable is position 4a that previously showed divergent behavior for Mcl-1 and Bcl-xL.<sup>17,21</sup> Similar to Mcl-1, Bcl-2 also shows promiscuity with regard to this position and is able to bind a range of different substitutions that do not bind Bcl-xL (compare last row of Figure 5A,B). This trend was to some extent predicted *a priori* (compare the last row of Figure 5C,D). An additional position that shows better binding to Bcl-2 over Bcl-xL across the array is position 2g. Substitutions at this site are more promiscuous with regards to Bcl-2 in comparison to Bcl-xL, which prefers mostly polar residues at this position. Lastly, Bcl-2 is a little more permissive for substitutions at position 2e. While this position still prefers Ala (native) or Gly, other small amino acids (Ser,Pro) bind equally well, and even bulkier residues are better accommodated than for Bcl-xL.

The similarity between the binding profiles of Bcl-2 and Bcl-xL can be identified using the predicted Bcl-2 array: Comparison of this array to the published experimental Bcl-xL array performed by Dutta et al. (Figure 5C,B) reveals considerable similarities: e.g., the “wild type” column is among the darkest in both arrays, proline can hardly be incorporated into any position, and position 3e permits only A, G, S for both. Thus, even though quantitatively the Pearson correlation between the predicted Bcl-2 array and the experimental Bcl-xL array is lower than for the two experimental arrays ( $r = 0.67$  when compared to the  $1 \mu\text{M}$  Bcl-xL array), the overall agreement is surprisingly good.



**Figure 5.** Computational and experimental characterization of Bcl-xL and Bcl-2 BIM-derived peptide-binding specificities. Sequences of 180 point mutants of the BIM BH3 peptide (TRAIN2) were evaluated for binding to Bcl-2 (left panels) and Bcl-xL (right panels). Experimental and predicted SPOT arrays are shown in the upper and lower panels, respectively. The predicted SPOT intensities are similar to the experimental SPOT arrays ( $r = 0.72$  for Bcl-xL and  $0.67$  for Bcl-2). The experimental specificity profiles of Bcl-xL and Bcl-2 are very similar ( $r = 0.88$ ; see text for comparison). The array image in (B) was taken from ref 21.



### 5. Binding Specificity of Additional BH3 Peptides.

Encouraged by the performance of FlexPepBind in this blind test for an additional Bcl-2 like protein, we wanted to assess its predictions for additional BH3 peptides. While until now all modeled peptide sequences were derived from the BIM BH3 peptide, the receptors encounter in the cell other BH3 sequences, such as those in proteins Bid, Bad, Bik, Noxa, Puma, Hrk, and Bmf. We modeled these BH3 sequences and predicted their binding profiles to Bcl-xL, Bcl2, and Mcl-1. Table 3 compares the predicted binding to experimental IC50

**Table 3. Computational Recapitulation of Distinct Binding Preferences of Bcl-2 Family Proteins to Different BH3 Substrates**

	peptide binding affinity		
	<100 nM <sup>a</sup>	<10 $\mu$ M	>100 $\mu$ M
Bcl-xL	Bim, Bmf, Bad, Puma, Hrk, Bid, Bik		Noxa
predicted rank <sup>b</sup>	1, 2, 3, 4, 5, 6, 8		7
Bcl-2	Bim, Bmf, Puma, Bad	Hrk, Bid, Bik	Noxa
predicted rank	1, 2, 4, 6	3, 5, 7	8
Mcl-1	Bim, Noxa, Puma	Hrk, Bmf, Bik, Bid	Bad
predicted rank	1, 2, 3	4, 5, 6, 7	8

<sup>a</sup>IC50 values of Bcl-2-like protein with BH3-only derived peptide (as reported in ref 60). <sup>b</sup>Predicted rank of binding by FlexPepBind among the biological substrates Bim, Bid, Bad, Bik, Noxa, Puma, Hrk, and Bmf.

values determined by Chen et al.<sup>60</sup> Even though no linear correlation is observed, there is a good correspondence between the experimental and predicted values. All three receptors bind tightest to the BIM peptide, and BIM indeed obtains for all three receptors the best score among the BH3 peptides. Mcl-1 is the only receptor to bind NOXA, and indeed NOXA obtains a good binding value for Mcl-1 but very poor values for Bcl-xL and Bcl-2. The reverse is seen for BAD, which binds Bcl-xL and Bcl-2, but not Mcl-1, and again the prediction results in intermediate binding values for the first two and a very poor value for the latter. This is an example of the generalization ability of our protocol, which can be calibrated based on significantly less experimental data compared to corresponding sequence based prediction methods.

## DISCUSSION

The range of interactions mediated by the binding of Bcl-2-like proteins to BH3 domains in their partners plays an important role in the regulation of apoptosis.<sup>5</sup> This interaction has been intensively studied in the past using a plethora of different approaches.<sup>18,21</sup> In addition, the detailed analysis of different solved crystal structures and drug design into the BH3 binding pocket have probed important structural characteristics of this interaction.<sup>6–9</sup> This well-characterized system therefore represents an ideal platform for the validation and further improvement of approaches that attempt prediction of binding specificity in peptide-mediated interactions, such as the structure-based approach presented in this study. In turn, insights from successful modeling of peptide binding specificity can also further improve our understanding of the important but complex system of regulation of apoptosis.

### Computational Structural Specificity Prediction for Flexible Peptides.

We have previously used structure-based prediction of binding specificity to successfully identify both known and novel protein farnesyltransferase (FTase) substrate peptides.<sup>35</sup> The structure-based characterization of BH3 peptides binding to Bcl-2-like proteins in this study demonstrates that this approach can indeed be generalized and applied to additional biological peptide–protein interactions. The two systems are markedly different: While for FTase, the binding motif is very short (four residues) and constrained by three specific interactions (conserved hydrogen bonds and Zn coordination), BH3 peptides are much longer (~20 residues), and although constrained by their helical conformation, they display a much larger overall degree of flexibility. It is probably for this reason that the latter required more intensive modeling (in comparison to a minimization-only protocol that provided good results for FTase<sup>35</sup>).

Modifications in the protocol during calibration that improved its performance for the system at hand highlight general problems mainly in the scoring function used for the modeling of the peptide–receptor complex structure and the ranking of the different sequences. The main improvements in the performance of the protocol (measured as Pearson correlation to the experimental SPOT array binding values for the training sets) were the modification of the solvation model to further penalize desolvation of Asp and Glu and the addition of an explicit electrostatic term. Both modifications make sense in light of the amphiphilic nature of the peptide–protein interface and should now be assessed on a larger scale to see how they impact the overall modeling accuracy of a variety of Rosetta protocols. Current discrepancies between predictions and experiment, such as the possible underestimation of the binding energy of valine at position 4a, emphasize future directions to explore for improvements.

The analysis of binding specificity of peptides from yeast surface display highlights the advantages of a sequence-based approach rooted in experiments, where the experiments can be fine-tuned for a specific system by assessment of additional sequences for correction of background effects. This has been nicely demonstrated also in a study just published that has significantly extended the amount of sequenced peptides that bind to different bcl-2 proteins and thus could report improved characterization of bcl-2 binding specificities.<sup>22</sup> Our structure-based approach in contrast uses experimental binding data merely to define a threshold. Despite its limitation in this specific case, this approach is expected to be more general in comparison to the PSSM-based methodology that will always be biased toward similar sequences. (Indeed, we showed in our previous work on FTase substrates that our approach can be used to identify novel substrates not detectable based on sequence similarity.<sup>35</sup>)

The conformational flexibility of the protein receptor might also play a major role in determining binding specificity. Structural studies have previously shown that Mcl-1 is able to accommodate surprising point mutations in the BIM BH3 sequence, without significant effect on its low nanomolar binding affinity. This is facilitated in part by side chain repacking (e.g., I2dA mutation<sup>9</sup>) as well as backbone movement of both the peptide and the Mcl-1 binding pocket (mainly the  $\alpha$ 3– $\alpha$ 4 helices, which are 1.5 Å RMSD apart<sup>9</sup>). While our protocol is able to capture side chain movements and the peptide flexibility (which for different mutants can differ up to 1 Å RMSD from the native conformation<sup>9</sup>), we do not

model the receptor backbone flexibility, which is critical for accommodation of some mutants (e.g., the binding of the BIM I2dY mutant to Mcl-1<sup>9</sup>). Indeed, our predictions correctly identify I2dA and F4aE as binders, while the mutation I2dY that involves backbone changes achieves a score that does not pass the defined threshold. Targeted incorporation of receptor flexibility in the future will therefore improve performance, provided that the new false positive conformations accessible thanks to increased flexibility can efficiently be screened.

Finally, the structural features of the unbound BH3 peptides might also contribute to their binding ability. The unstructured regions in BH3 containing proteins play an important regulatory role, and the length of these regions is regulated by alternative splicing. The BH3 peptides themselves are unstructured in solution and adopt a helical conformation upon binding to Bcl-2-like family proteins.<sup>61</sup> Helical propensity itself might play a major role in the binding process: Indeed, studies have shown that mutations that mainly affect helical propensity (either by replacement with helix-favoring amino acids or by elongation/shortening of the helix) can dramatically affect binding.<sup>62</sup> By itself, however, helical propensity might not be the main determinant of binding, as assessment of helical propensity of the different mutants analyzed in the present study (using AGADIR<sup>63</sup>) did not show any clear and consistent correlation between the ease of helix formation and the ability of BIM-derived peptides to bind to Bcl-2-like proteins (the correlation coefficients of AGADIR scores with TRAIN1/TRAIN2 binding data were Bcl-xL  $r = -0.6/0.12$  and Mcl-1  $r = -0.23/0.33$ , for the 100 nM binding assays).

**Large-Scale Identification of BH3 Domains in Humans and Pathogens.** We have shown that our approach is not limited to a specific receptor, nor to sequences derived from a specific BH3 peptide. We therefore anticipate that FlexPepBind can be used for the identification of novel BH3 binding sequences in the human genome and in other genomes, similar to its application to the identification of potential novel human FTase targets in our previous study.<sup>35</sup> BH3-only proteins are not limited to the Bcl-2 like protein family. Such functional sequences were found for example e.g. in an E3 ubiquitin ligase that utilizes a BH3 domain to bind specifically to its target Mcl-1 and mark it for degradation.<sup>64</sup> Application of our protocol to the putative BH3 domain of this protein (mule) indeed predicted a high-affinity interaction with Mcl-1 (data not shown). Another validated BH3 sequence can be found in nuclear clusterin (CLU), which is a multifunctional glycoprotein that is overexpressed in prostate and breast cancers.<sup>65,66</sup> Because of the poor sequence identity of known BH3 domains, sequence-based predictors are not suitable for exact determination of the binding properties of BH3-like sequences but can rather serve as an initial fast screen to offer candidate sequences later to be filtered by FlexPepBind.

Apoptosis is part of the cell's combat arsenal against invading viruses. Consequently, many viruses have evolved strategies to undermine cell suicide. One such mechanism is the expression of Bcl-2-like antiapoptotic proteins: For example, Epstein–Barr and Kaposi's sarcoma viruses both encode Bcl-2 homologues (BHRF-1 and KSHV Bcl-2, respectively). It was shown that these proteins are not only able to bind BH3 domain peptides found in the cell but also bear unique specificity profiles: BHRF-1 binds Bim, Bid and Puma, but not other BH3 domains,<sup>67</sup> while KSHV Bcl-2 shows a specificity profile similar to Mcl-1,<sup>68</sup> binding most BH3 domains except for Bad.<sup>69</sup> Application of FlexPepBind on a solved crystal structure of

BHRF1 bound to Bim (PDB: 2wh6<sup>67</sup>) recovers the main binding preferences of BHRF1 to cellular BH3 domains: Bim and Bid are identified as tight binders (ranked 1 and 3), while Puma is ranked less favorably (rank 6 out of 8 BH3 peptides). The structure of KSHV Bcl-2 in complex with a ligand has not been solved yet, but an obvious extension of our protocol would be to predict binding using the free structure or a homology model. This would however require further calibration and testing. These preliminary results indicate that FlexPepBind might be a valuable tool for the detection and characterization of additional viral receptors.

**Specificity Profiles of Bcl-2 and Bcl-xL.** Our results demonstrate that Bcl-2 and Bcl-xL share a very similar specificity profile for BIM-BH3 derived point mutants. This is surprising in light of the fact that the two proteins are not highly conserved: only 14 of the 21 interface residues in Bcl-xL are conserved in Bcl-2 (i.e., 66% sequence identity at the interface; overall sequence identity of 40%). On the other hand, previous studies have shown that the binding preferences for biological BH3 sequences of these two proteins are indeed similar. What then is the biological role of this apparent redundancy? One possibility is robustness: owing to their cardinal role in determining cell fate, they act as a backup for each other (this motif is found in many signal transduction pathways). Another possibility is that the small differences in binding preferences (such as a ~100-fold preference for Bcl-xL to bind Bik or Bid) might lead to an overall different outcome when integrating the cellular network of interactions. Indeed, in a large-scale assessment of squamous cell carcinoma, most cell lines upregulated either Bcl-2 or Bcl-xL, but not both. Moreover, the survival profiles of the different subgroups differed significantly.<sup>70</sup> This and other studies indicate that these two proteins play similar but not identical roles.<sup>71</sup> Further investigation of these differences with tools such as the one presented in this study may help to obtain a more comprehensive view of the underlying features of the cellular network of interactions and their contribution to cellular behavior.

## ■ ASSOCIATED CONTENT

### § Supporting Information

Calibration of the protocol: correlations between predicted and experimental BIM-derived peptides to Bcl-xL and Mcl-1 are indicated for different sets of parameters tested (Table S1); performance of the protocol as a function of the threshold used to define binding peptide sequences in the SPOT arrays (Figure S1); details of positions R3b and F4a on the structure of the Mcl1–BIM complex (Figure S2). This material is available free of charge via the Internet at <http://pubs.acs.org>.

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## Notes

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## ABBREVIATIONS

AUC, area under roc curve; Bcl2, B cell lymphoma 2; BH, Bcl2-homology domain; BIM, Bcl-2-interacting mediator of cell death; BLU, Boehringer light unit; FPR, false positive rate; FTase, farnesyltransferase; Mcl1, myeloid leukemia cell differentiation protein; PDB, Protein Data Bank; PSSM, position specific scoring matrix; RMSD, root mean square deviation; ROC, receiving operating characteristic; TPR, true positive rate; Wsc, weighted score.

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