Structural changes in the BH3 domain of SOUL protein upon interaction with the anti-apoptotic protein Bcl-xL

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The SOUL protein is known to induce apoptosis by provoking the mitochondrial permeability transition, and a sequence homologous with the BH3 (Bcl-2 homology 3) domains has recently been identified in the protein, thus making it a potential new member of the BH3-only protein family. In the present study, we provide NMR, SPR (surface plasmon resonance) and crystallographic evidence that a peptide spanning residues 147–172 in SOUL interacts with the anti-apoptotic protein Bcl-xL. We have crystallized SOUL alone and the complex of its BH3 domain peptide with Bcl-xL, and solved their three-dimensional structures. The SOUL monomer is a single domain organized as a distorted β-barrel with eight anti-parallel strands and two α-helices. The BH3 domain extends across 15 residues at the end of the second helix and eight amino acids in the chain following it. There are important structural differences in the BH3 domain in the intact SOUL molecule and the same sequence bound to Bcl-xL.

Key words: apoptosis, Bcl-xL, Bcl-2 homology 3 domain (BH3 domain), crystal structure, NMR, SOUL, surface plasmon resonance.

INTRODUCTION

The Bcl-2 family are a group of evolutionarily conserved proteins that interact to maintain a balance between newly forming and old, damaged or superfluous dying cells. [1]. They play a central role in the regulation of apoptosis, or programmed cell death, a process that in multicellular organisms leads to the controlled death of unneeded or unwanted cells. A crucial event in apoptosis is the MPT (mitochondrial permeability transition), a drastic increase in the permeability of the mitochondrial inner membrane to low-molecular-mass solutes [2]. Through the regulation of apoptosis, the Bcl-2 proteins have an important function in embryogenesis [3], tissue remodelling [4] and the immune response [5]. Their abnormal behaviour is linked to many diseases, such as autoimmunity [6], neurodegenerative disorders [7] and cancer [8].

The effect of the Bcl-2 proteins on the apoptotic process is due to the presence of one or more conserved regions of amino acid sequences, known as BH (Bcl-2 homology) domains, named BH1, BH2, BH3 and BH4 [9,10]. The proteins of this family that contain only the BH3 domain are pro-apoptotic and function as initial sensors of apoptotic signals resulting from various cellular processes, whereas the pro-survival Bcl-2 family members, such as Bcl-2 or Bcl-xL, wield their effect by binding and sequestering their pro-apoptotic counterparts [11]. Peptides spanning the sequence of BH3 domains appear to exert the physiological activity of the intact proteins and their complexes with anti-apoptotic members of the Bcl-2 family have received considerable attention since this interaction is believed to explain the effect at the molecular level. In particular, the complexes of peptides with the sequences of the BH3 domains of Bad, Bim, Bak, Bid and Beclin 1 with Bcl-xL have been examined by X-ray crystallography and NMR, and the conserved crucial interactions between the peptides and the protein have been identified [12,13].

Cancer cells frequently overexpress the anti-apoptotic members of the Bcl-2 family, and small molecules that incorporate the structural features of the BH3 domains necessary for binding to these anti-apoptotic proteins have been synthesized and are being tested as specific cancer cell killers [14].

SOUL was first identified at the transcriptional level by suppression subtractive hybridization in the chicken retina and pineal gland, and its gene was named ckSoul because of the high transcript levels found in the pineal gland, the organ René Descartes hypothesized was the location of the soul [15]. A few years before this report, human SOUL had been isolated and characterized from saline extracts of human term placentas and had been called PP33 (placental protein 23) [16]. More recently, the protein has also been identified in human amniotic fluid [17]. It has subsequently been shown that the gene coding for this protein is very widely distributed in evolution and it has been characterized in many other species, including the popular model organism of plant biology Arabidopsis thaliana. On the basis of its sequence similarity with the mouse gene p22HBP, which codes for HEBP1 (haem-binding protein 1) or p22HBP, SOUL has also been called the alternative name of HEBP2 [18]. Recombinant mouse SOUL was reported to be a dimer in the absence of haem and to hexamerize upon haem binding, with a dissociation constant in the nanomolar range [19].

A very important observation is that SOUL can induce the MPT, a condition that leads to mitochondrial swelling and cell death [20]. More recently, analysis of the human SOUL sequence revealed the presence of a putative BH3 domain of the Bcl-2 family protein whose deletion abolished the apoptotic effects of SOUL [21].

In the present paper, we provide experimental evidence that SOUL is a BH3-only protein as the sequence spanning amino acids 147–172 interacts with the anti-apoptotic protein Bcl-xL. In addition, we have crystallized both intact SOUL and the complex

Abbreviations used: BH, Bcl-2 homology; HEBP, haem-binding protein; HSQC, heteronuclear single-quantum coherence; MPT, mitochondrial permeability transition; rmsd, root mean square deviation; RZPD, Deutsches Ressourcenzentrum für Genomforschung; SPR, surface plasmon resonance.

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The co-ordinates of the models, and the structure factors of SOUL and of the complex of human Bcl-xL with the peptide have been deposited in the PDB under accession codes 3R8S, 3R8J and 3R8K.

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of a peptide that contains its BH3 domain with Bcl-xL, and solved their three-dimensional structures to 1.6 and 2.0 Å (1 Å = 0.1 nm) resolution respectively. The Bcl-xL–SOUL BH3 domain interactions are particularly interesting, since the domain adopts a different structure when bound to Bcl-xL.

**EXPERIMENTAL**

**Protein expression and purification**

The cDNA coding for human SOUL (IMAGE ID 3445763), obtained from RZPD (Deutsches Ressourcenzentrum für Genomforschung), was amplified by PCR using primers designed to introduce restriction sites for BamHI and HindIII endonucleases and a sequence coding for a digestion site for thrombin in the C-terminal end in the amplified fragment. After purification, the fragment and the expression vector pQE50 (Qiagen) were digested with the restriction enzymes designed to introduce restriction sites for BamHI and HindIII. The digested DNA was ligated and the ligation mixture was used to transform E. coli C41 BL21 strain (Novagen, Germany), and clones were selected on agar plates containing ampicillin (50 μg/ml) and IPTG (isopropyl β-D-thiogalactopyranoside). The clones were verified by PCR and sequencing.

**Protein expression in E. coli**

Recombinant human Bcl-xL (IMAGE ID 2823498; RZPD) was expressed in a similar way to the SOUL expression. E. coli BL21 C41 strain Escherichia coli cells were transformed with the resulting vector, grown at 37°C and protein synthesis was induced overnight at 20°C with 0.5 mM IPTG. The cell pellet was collected by centrifugation and washed with 0.1 M NaCl, 0.02% (v/v) Triton X-100, and 20 mM Tris/HCl (pH 7.5). The protein was then purified further by gel filtration on a Superdex 200 column equilibrated with 20 mM Tris/HCl (pH 7.5), 0.15 M NaCl and 0.005% P-20 surfactant at 20°C using a SensiQ Pioneer instrument (ICX Technologies). Data were analysed with the Qdat evaluation analysis software.

**Crystallization, X-ray data collection, structure solution and refinement**

Purified native SOUL was used at approximately 20 mg/ml for the initial screen of crystallization conditions. Molecular Dimensions Structure Screens were used at 20°C with the hanging-drop method, mixing 1 μl of the protein solution with the same volume of the precipitating solution, and equilibrating against a volume of 0.3 ml of the latter in the reservoir. The conditions yielding small crystals were later refined and the sitting-drop method with larger volumes was also tested until crystals that were large enough for data collection were obtained.

The SOUL BH3 peptide spanning amino acids 147–172 was synthesized by TAG Copenhagen. The complex of recombinant human Bcl-xL with the peptide was prepared by mixing the protein at approximately 4 mg/ml with four times the molar ratio of the peptide. The mixture was incubated for about 1 h and then concentrated to approximately 15 mg/ml, and was then used at this concentration for the crystallization experiments.

**NMR measurements**

For the production of 15N-labelled human Bcl-xL lacking only the C-terminal transmembrane domain, host cells were grown in M9 minimal medium using 15NH4Cl as sole nitrogen source. HSQC (heteronuclear single-quantum coherence) NMR spectra were recorded on a Bruker Avance spectrometer operating at 600.13 MHz, equipped with a cryoprobe. The labelled protein, dissolved in 20 mM Tris/HCl (pH 7.5) and 0.15 M NaCl (in 10% 2H2O) and at a concentration of 85 μM, was titered with the SOUL BH3 peptide dissolved in the same buffer at 600 μM. Nine additions were made so that, after correcting for the precise peptide concentration and taking into account dilutions, the BH3 peptide/protein molar ratio was 0.07, 0.17, 0.26, 0.35, 0.52, 0.69, 1.38, 2.77 and 3.83. After each of the additions, the sample was incubated at 20°C for approximately 5 min and a one-dimensional 1H-15N HSQC spectrum was recorded at the same temperature. Standard sequence schemes with pulsed-field gradients were used to achieve the suppression of the solvent signal.

The dissociation constant was calculated from the shifts in two peaks in fast exchange in the methyl region of the decoupled 1H spectrum fitting the data with the equation:

\[
\Delta = \Delta_{\text{max}} \left[ (P + L + K_d) - \sqrt{(P + L + K_d)^2 - 4PL} \right] / 2P
\]

where \(\Delta\) is the chemical shift change, \(P\) is the protein concentration, \(L\) is the total ligand concentration, and \(K_d\) is the dissociation constant. Appropriate corrections for the dilution of protein and ligand were made after the addition of each aliquot during the titration.

**SPR (surface plasmon resonance) studies**

Bcl-xL was immobilized on a COOH1 research-grade sensor chip (Nomadics) by amine-coupling chemistry using the manufacturer’s protocols, and the SOUL BH3 domain peptide was used as the analyte. SPR measurements were carried out in HBS buffer [10 mM Hepes (pH 7.4), 150 mM NaCl, 3 mM EDTA and 0.005% P-20 surfactant] at 20°C using a SensiQ Pioneer instrument (ICX Technologies). Data were analysed with the Qdat evaluation analysis software.
molecules in the asymmetric unit (see Table 1), diffracts to approximately 2.85 Å and appears to be closely related to another crystal form reported in the literature [23]. The crystals grow by adding equal volumes of the protein solution and 0.1 M Tris/HCl (pH 8.5) and 2.0 M ammonium sulfate. The second crystal form is orthorhombic, space group C222₁, and a = 137.7 Å, b = 114.7 Å and c = 67.4 Å, and grows by mixing equal volumes of the protein solution and 0.85 M NaH₂PO₄, KH₂PO₄ and 0.08 M Hepes (pH 7.5). These crystals diffract to a better resolution, approximately 2.3 Å, contain two molecules in the asymmetric unit and are the crystal form that was solved first using the S.I.R. (single isomorphous replacement) method. The hexagonal crystal form was solved later by molecular replacement.

The best crystals of the complex human Bcl-xL–SOUL BH3 peptide grew by mixing equal volumes of the complex solution and 15% PEG [poly(ethylene glycol)] 6000, 0.2 M sodium sulfate and 0.3 M 1-butyl-3-methylimidazol-2-[(2-methoxyethoxy)ethyl] sulfate. They are tetragonal, space group P4₃, with a = b = 66.8 Å and c = 175.2 Å, and diffract to 2.0 Å resolution.

The diffraction data were collected from crystals frozen at 100 K after a brief immersion in a mixture of 80% of the mother liquor and 20% glycerol. The data set for a gold heavy atom derivative used for phasing were obtained using copper Kα radiation from a Rigaku RU-300 rotating anode X-ray generator with a Mar345 imaging plate area detector. The final data sets used for refinement of this and the other crystal forms were collected at the ID14-2 beamline of the ESRF (European Synchrotron Radiation Facility) in Grenoble (λ = 1.001 Å). The data were indexed, integrated and reduced using the programs AUTOMAR, MOSFLM and Scala [24]. The diffraction data statistics of the data sets used for refinement are summarized in Table 1.

Initial phases for the orthorhombic crystals to a 2.3 Å resolution were determined by the single isomorphous replacement method with the derivative data collected at the home source. Two gold sites were located in a difference Patterson map using the program SHELXS [25] and entered as input for the program autoSHARP [26] that was used to locate the minor sites of the derivative, and for density modification and final phasing to 1.8 Å resolution. The electron density map thus produced was of very good quality and could be readily interpreted. The initial model of SOUL was built in this map using the program Coot [27].

Refinement was carried out initially using the program REFMAC [28] and, in a second stage, with the program Phenix. refine [29]. During the process of refinement and model building, the quality of the models was controlled with the program PROCHECK [30]. Solvent molecules were added to the model in the final stages of refinement according to hydrogen-bond criteria and only if their B factors refined to reasonable values and if they improved the Rfree. The model was finally subjected to a final round of TLS refinement.

The structure of the hexagonal crystal form of SOUL was solved using the CCP4 suite of programs for crystallographic computing. The initial phases were calculated by the molecular replacement method.
Figure 1 Overall structure and folding of SOUL

(A) Stereodiagram of the SOUL molecule. The eight-stranded $\beta$-sheet is shown in two colours, red and blue, to emphasize the presence of two repeated units, each with $\beta$-$\beta$-$\alpha$-$\beta$-$\beta$ topology, related by a pseudo-two-fold axis of symmetry. (B) A topological diagram of SOUL. The eight antiparallel $\beta$-strands form a distorted $\beta$-barrel with the two $\alpha$-helices arranged on one face. Strands are labelled in the order of their appearance from the N-terminus to the C-terminus using the letters A–H. The eight strands span the following residues: A, 39–43; B, 46–55; C, 89–94; D, 103–110; E, 127–132; F, 135–142; G, 174–178; H, 190–195. The two helices span residues 58–73 and 148–164. (C) Ribbon representation of the SOUL monomer viewed in a direction rotated approximately 90° with respect to (A). The two helices are in yellow, and the strands of the $\beta$-sheet have been coloured as (A and B) to emphasize the presence of the pseudo-two-fold axis. (D) Ribbon diagram of SOUL with the peptide spanning the BH3 domain represented in light blue for the portion of the chain identified by sequence similarity with known BH3 domains and in yellow for the rest of the second helix of the molecule. (E) Ribbon diagram of the NMR model of BID (lowest energy structure, PDB code 2BID) with the BH3 peptide and domain oriented and colour-coded as in (D). The Figures of the models were prepared using the program PyMOL (http://www.pymol.org).

method as implemented in the program MOLREP [31], with the co-ordinates of the orthorhombic model as the search probe. The automatic search with data up to a resolution of 2.9 Å gave a solution that placed in their correct position three out of the four molecules present in the asymmetric unit. Fixing these co-ordinates, the fourth molecule was found by the same program. The score of this solution was 0.544 and its $R$ factor was 41.6%.

A similar procedure was followed to solve and refine the structure of the complex human Bcl-xL–SOUL BH3 peptide using the co-ordinates of a protomer of Bcl-xL ([32]; PDB code 2YXJ) as the search probe. After the four protomers of Bcl-xL present in the asymmetric unit had been placed in their correct position, it became evident that they dimerized with domain swapping and, at this point, the extra electron density for the four BH3 helices present in the asymmetric unit was also very clear. The final refinement statistics for the models of the three crystal forms are summarized in Table 1.

RESULTS

X-ray structure of human SOUL

The SOUL monomer is a single domain structure organized as an open distorted $\beta$-barrel with eight anti-parallel strands. The barrel is open in the sense that the first and the last strands (A and E) are not in contact in the sheet (Figure 1). Two $\alpha$-helices connect the second to the third strand and the sixth to the seventh strand. They are both located on one face of the molecule and pack against the curved sheet that forms the barrel. A monomer of SOUL fits into a box with the approximate dimensions $56 \times 47 \times 40$ Å. This fold is quite different from that of the canonical member of the BH3-only protein family BID (Figure 1).

Examination of the secondary structure (Figure 1B) reveals that the molecule is made up of two repeated units, each with $\beta$-$\beta$-$\alpha$-$\beta$-$\beta$ topology related by a pseudo-two-fold axis of symmetry. Two SOUL monomers (A and B) are present in the asymmetric unit of the orthorhombic crystal form (Table 1). The secondary structure assignments of monomer A are, for the $\beta$-strands, the following: strand A, residues 39–43, B, residues 46–55, C, residues 89–94, D, residues 103–110, E, residues 127–132, F, residues 135–142, G, residues 174–178 and H residues 190–195. The two $\alpha$-helices span residues 58–73 and 148–164. In addition, residues 25 and 26 extend the $\beta$-sheet and residues 113–116 form an additional helix turn in both monomers. A minor difference between the two monomers was also observed: strand D of molecule B has one residue less at the N-terminal end.

The space within the $\beta$-barrel is filled by the side chains of both hydrophobic and hydrophilic residues: Trp$^{39}$, Met$^{135}$, Leu$^{137}$,
Leu<sup>130</sup> and Trp<sup>193</sup>, but also Arg<sup>41</sup> and Arg<sup>132</sup>, Asp<sup>130</sup>, Asp<sup>191</sup>, Thr<sup>90</sup> and Tyr<sup>179</sup>. Packing of the first helix takes place through hydrophobic residues and the following specific contacts: Trp<sup>98</sup> (NE1)–Ser<sup>91</sup> (OG), Lys<sup>68</sup> (O)–Glu<sup>124</sup> (OE), Tyr<sup>72</sup> (OH)–Pro<sup>120</sup> (O), Tyr<sup>72</sup> (O)–Gln<sup>77</sup> (OE) and Ile<sup>3</sup> (O)–Ile<sup>83</sup> (N). Packing of the second helix involves the following specific contacts: Gln<sup>154</sup> (OE)–Arg<sup>140</sup> (NE) and Leu<sup>162</sup> (O)–Lys<sup>167</sup> (N).

Two NMR structures of murine p22HBP have been published [33,34]. The protein has approximately 28% sequence identity with murine SOUL. Comparison of the two SOUL molecules present in the crystallographic asymmetric unit of the orthorhombic crystal form with the two NMR models of murine p22HBP revealed that the four models differ substantially only in rather limited areas (Supplementary Figure S1 at http://www.BiochemJ.org/bj/438/bj4380291add.htm). The zone where the two SOUL molecules in the asymmetric unit differ more from one another are the region before the first strand of the β-sheet and the loop connecting strands C and D. The chains before the first strand are totally exposed to the solvent, whereas in the case of the connection of strands C and D the loop in chain A is in close contact with a symmetry-related molecule, and the equivalent area in molecule B is in contact with the solvent. These differences are thus probably simply a consequence of molecular packing in the crystal. Although there is more variability in the two NMR structures of murine p22HBP, the two models are very similar in the region connecting strands C and D, which is also the region where both are most different from SOUL. This particular region of the SOUL molecule thus appears to be more variable than the rest of the molecule.

The BH3 domain predicted by sequence alignment to be present in SOUL spans residues 158–172, i.e. the last seven residues of the second α-helix of the molecule and eight amino acids in the loop connecting it to strand G. The domain is represented in light blue in Figure 1(D). Additional details on this structure have been included as Supplementary information (at http://www.BiochemJ.org/bj/438/bj4380291add.htm).

**Does SOUL bind haemin?**

All our attempts to prepare co-crystals of SOUL and haemin failed. When crystallizations were set up, with haemin/SOUL molar ratios of even up to five, the crystals obtained, after screening many different conditions, were invariably those of the apo protein. Soaking the pre-formed crystals did not reveal any electron density other than that of the apo protein. It was thus suspected that the interaction of the two molecules, if present, was not as strong as expected. For this reason, the UV–visible spectrum of haemin was examined alone and in the presence of ten times the molar ratio of SOUL, with BSA used as a control (Supplementary Figure S2A at http://www.BiochemJ.org/bj/438/bj4380291add.htm). Although the sample containing BSA had, due to the ligand–protein interaction, its peak higher and shifted from 390 to 401 nm, as expected for haemin bound with no axial co-ordination to a hydrophobic cavity [35], the sample containing SOUL showed only negligible differences that can be explained by the absorption of the protein present in the sample and cannot be considered as evidence of a SOUL–haemin interaction.

A second control was carried out recording <sup>15</sup>N-<sup>1</sup>H HSQC NMR spectra of <sup>15</sup>N-labelled SOUL in the presence of increasing amounts of haemin dissolved in Tris/HCl buffer. Haemin aliquots corresponding to 0.5, 1, 2, 3 and 4 equivalents of SOUL were added to the protein sample and the spectra were recorded at 25°C (Supplementary Figure S2B). After the addition of up to four equivalents of haemin per molecule protein, the spectrum remained unaltered. Given the time involved to record the different spectra, a kinetic effect can be excluded in this case or at least if there was such an effect it has to be proposed that the reaction is so slow that, even after more than 1 day of observation, no change in the spectrum is detectable at all. These observations thus lead to the conclusion that, with the methods described in the present study, no interaction between haemin and SOUL, that may be considered of physiological relevance, can be observed.

**Interaction of the SOUL BH3 peptide with human Bcl-xL**

The BH3 domain predicted by sequence alignment to be present in SOUL spans residues 158–172, i.e. the last seven residues of the second α-helix of the molecule and eight residues in the loop connecting it to strand G. BH3 domains are known to be helical and, since we knew that in the structure of SOUL the helix began before, we decided to examine the interaction of a 26-amino-acid-long peptide spanning residues 147–172 with the anti-apoptotic protein Bcl-xL, i.e. covering the entire helix and the region predicted to be part of the BH3 domain by sequence homology. The sequence of the peptide studied is: SAQKNEQQLTLASILREDGKVFDK. Figure 1(D) shows the SOUL monomer with the BH3 peptide coloured light blue for the chain predicted to be part of the domain by sequence alignment and yellow for the rest of the peptide corresponding to the N-terminal portion of the second α-helix of SOUL.

The interaction of the peptide and the protein was examined in solution by one- and two-dimensional NMR and by SPR. Figure 2(A) shows the amidic region of the <sup>15</sup>N-<sup>1</sup>H HSQC NMR spectrum of <sup>15</sup>N-labelled human Bcl-xL (lacking the transmembrane domain after amino acid 209) titrated with increasing amounts of the peptide. Note the significant changes in some of the peaks of the protein as the interacting BH3 peptide was added to the solution. Figure 2(B) shows the chemical shift displacements in the two peaks in fast exchange in the methyl region of the <sup>1</sup>H spectrum that were used to calculate an approximate dissociation constant of the interaction, and Figure 2(C) shows the fitting of these displacements as a function of equivalents of the BH3 peptide added. The dissociation constant values calculated using the two peaks are in reasonable agreement with one another and are 47.8 and 41.3 μM (see Figure 2C).

The SOUL BH3 peptide–Bcl-xL interaction was also studied by SPR. Both intact Bcl-xL and a form lacking amino acids 27–82 that is known to bind BH3 domains like the intact protein [36] were immobilized on a sensor chip and the SOUL BH3 domain peptide was used as the analyte. Similar results confirming the interaction were observed for both variants of Bcl-xL. Figure 2(D) shows a sensorgram of the Δ27–82 form of the protein. The dissociation constant estimated with this method is approximately five times the value observed in the NMR experiments. The discrepancy is probably due to errors in the estimate of the peptide concentration used in the experiments. However, we did not detect significant binding between intact SOUL and Bcl-xL (results not shown), which may reflect either a lack of interaction under the conditions tested or more probably the requirement of drastic structural changes in SOUL.

**X-ray structure of the SOUL BH3 domain peptide complexed with Bcl-xL**

The data collection and refinement statistics of the co-crystals of human Bcl-xL (Δ27–82) with the SOUL BH3 peptide are summarized in Table 1. The crystals are tetragonal, space group

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**Interaction of SOUL protein BH3 domain with Bcl-xL**

The BH3 domain predicted by sequence alignment to be present in SOUL spans residues 158–172, i.e. the last seven residues of the second α-helix of the molecule and eight residues in the loop connecting it to strand G. BH3 domains are known to be helical and, since we knew that in the structure of SOUL the helix began before, we decided to examine the interaction of a 26-amino-acid-long peptide spanning residues 147–172 with the anti-apoptotic protein Bcl-xL, i.e. covering the entire helix and the region predicted to be part of the BH3 domain by sequence homology. The sequence of the peptide studied is: SAQKNEQQLTLASILREDGKVFDK. Figure 1(D) shows the SOUL monomer with the BH3 peptide coloured light blue for the chain predicted to be part of the domain by sequence alignment and yellow for the rest of the peptide corresponding to the N-terminal portion of the second α-helix of SOUL.

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Figure 2 Interaction of the SOUL BH3 peptide with human Bcl-xL

(A) 15N-1H NMR correlation spectra of 15N-labelled human Bcl-xL titrated with increasing amounts of a 26-amino-acid-long peptide spanning residues 147–172 of human SOUL. The peptide sequence is SAQKNQEQLLTLASILREDGKVDEK. Arrows indicate the direction of peak shifts. Only four spectra are represented in the Figure; the black one is before any addition, the green after the addition of 0.52 equivalents of the peptide, the blue after the addition of 1.38 equivalents and the red after adding 3.83 equivalents of the BH3 domain peptide. (B) Decoupled 1H spectrum showing the shifts in two peaks in fast exchange in the methyl region used to calculate an approximate dissociation constant. The colour of the spectra is the same as in (A) and corresponds to the addition of the same amounts of peptide. (C) Magnitude of the change in the 1H chemical shift of the two selected peaks plotted as a function of the total number of equivalents of BH3 domain peptide added. The best fit curve is shown along with the coefficient of determination ($R^2$) and the calculated dissociation constants. The values determined using the two peaks are 47.8 and 41.3 μM. The equation used to fit the data is given in the Experimental section. (D) SPR studies of the same interaction. The sensorgram shows the binding of the BH3 peptide to truncated immobilized Bcl-xL. Relative units (RU) are plotted as a function of time (in s). (E) A plot of the response as a function of the peptide concentration used to estimate the dissociation constant. The diagram is the result of several experiments and higher peptide concentrations could not be used because the BH3 peptide had a tendency to aggregate.

The model was refined to give the statistics listed in Table 1. The asymmetric unit contains four Bcl-xL protomers organized as dimers that exhibit domain swapping, exchanging their N-terminal α1-helix (Figure 3A). This kind of domain swapping has been observed in several co-crystals of BH3 domains and human Bcl-xL (Δ27–82) [13,37–39]. It is considered to be an artifact
The BH3-only members of the Bcl-2 protein family play a central role in the process that leads to programmed cell death or apoptosis. Their effect is due to inhibition through binding of their BH3 domain in the hydrophobic cleft of the anti-apoptotic members of the Bcl-2 family, such as Bcl-2, Bcl-xL or MCL-1.
Indeed, one of the criteria to include a protein in this group is a demonstration of its interaction with one of the pro-survival proteins of the Bcl-2 family, in addition to showing that they have a cellular death-inducing activity. On the basis of these criteria, since the discovery of the first member of the family, BIK [44], many other members have been added to the list [9,10]. In many cases, the proteins seem to have additional functions besides their role in inducing cellular death. Although a very large number of complexes of BH3 peptides with pro-survival proteins of the Bcl-2 family are available [12,45], only one NMR structure of an intact BH3-only protein is known, BID [46,47].

Two different mutually non-excluding functions have been attributed to SOUL: haem transport [19] and a role in apoptosis as a BH3-only protein [20,21]. We have used NMR, SPR, UV spectroscopy and crystallography to explore both functions and, in addition, we have determined the three-dimensional structure of the protein. Human SOUL is a monomer with a fold which is quite different from that of the other BH3-only protein whose three-dimensional structure is known, BID, which is similar to Bcl-xL. BID contains eight α-helices: two central hydrophobic helices surrounded by six amphipathic helices with their hydrophilic face exposed to the solvent. The SOUL molecule is similar to murine p22HBP [33,34].

The interaction of human SOUL with haemin was explored by titrating the latter with the former and following the UV spectrum, using BSA as a control. In our experiments, we made sure that the protein used did not contain any traces of the histidine tag used for purification and that no imidazole, residual from the purification
Asp170 (shown in Figure 4) are indicated with dotted lines. The aspartate residue is conserved of Arg139 of Bcl-xL with an aspartate residue in Beclin 1 (conserved in all BH3 domain–Bcl-xL in Beclin 1 and the equivalent of the arginine residue is a lysine. Note that the important contact present study and work on the related p22HBP proteins. It is also reported to be a dimer in the absence of haem and a hexamer is missing in the SOUL peptide.

We do not have an explanation as to why our results appear to be in the bound state, which is different from both the results of the other studies and control, revealed that SOUL is not a substrate of this enzyme but up to now there has been no information at the molecular level on the mechanism through which this function is accomplished.

However, an experiment with caspase 8, using human BID as a control, revealed that SOUL is not a substrate of this enzyme. This cleavage site is not present in SOUL, although the similar sequence LESDV spans residues 123–127 and is exposed to the solvent in the molecule, in the loop connecting strand D to E. This sequence LESDV spans residues 123–127 and is exposed to the solvent in the molecule, in the loop connecting strand D to E. However, an experiment with caspase 8, using human BID as a control, revealed that SOUL is not a substrate of this enzyme.

The role of SOUL in inducing apoptosis has been documented, but up to now there has been no information at the molecular level on the mechanism through which this function is accomplished. We have shown that its BH3 domain interacts with a pro-survival member of the Bcl-2 family and thus we have provided new evidence that SOUL behaves like a novel member of the survival Bcl-2 targets. No important changes were found in Bcl-xL, but the changes in the BH3 domain are remarkable and are sufficient to explain why no interaction is observed with the intact SOUL molecule. These drastic modifications might require conditions that have not yet been found, but that should be explored further given the importance of this interaction in the functionality of the two proteins.

The mechanism of activation of BID, the prototype of BH3-only proteins, involves cleavage by caspase 8 in a region with the sequence LQTDG. The second amino acid in the sequence can be any amino acid other than phenylalanine, glutamate, glutamine, lysine or arginine. This cleavage site is not present in SOUL, although the similar sequence LESDV spans residues 123–127 and is exposed to the solvent in the molecule, in the loop connecting strand D to E. However, an experiment with caspase 8, using human BID as a control, revealed that SOUL is not a substrate of this enzyme (results not shown).

Figure 5 Structural changes in the BH3 peptide of SOUL upon binding to Bcl-xL

(A) Comparison of the peptide bound to Bcl-xL (Δ27–82) (blue) and the same sequence in the intact SOUL molecule (red). Eight amino acids change their conformation to extend the BH3 helix towards its C-terminus. The co-ordinates were superimposed by using the CCP4 suit of programs. Three charged amino acids that give specificity to the interaction are represented as stick models in the two conformations. (B) Superposition of the SOUL BH3 peptide Bcl-xL (Δ27–82) complex (the peptide is in yellow and Bcl-xL is in orange) and the same complex of Bcl-xL (in light blue) and the Beclin 1 BH3 peptide (in green). The PDB code of the model of the Beclin 1 peptide complex is 2P1L [13]. The interactions of the SOUL amino acids Arg163 and Asp170 (shown in Figure 4) are indicated with dotted lines. The aspartate residue is conserved in Beclin 1 and the equivalent of the arginine residue is a lysine. Note that the important contact of Arg163 of Bcl-xL with an aspartate residue in Beclin 1 (conserved in all BH3 domain–Bcl-xL complexes) is missing in the SOUL peptide.

in the affinity columns, was present in the samples. We did not find any evidence of an interaction between haem and SOUL. This result was confirmed by the 15N-1H HSQC NMR spectra of 15N-labelled SOUL in the presence of increasing amounts of haem.

We do not have an explanation as to why our results appear to be at variance with those reported for mouse SOUL [19]. The latter is reported to be a dimer in the absence of haem and a hexamer in the bound state, which is different from both the results of the present study and work on the related p22HBP proteins. It is also worth noting that the histidine residue that Sato et al. [19] found to be essential for haem binding does not seem to be involved in haem binding to p22HBP, which appears to bind this moiety through hydrophobic interactions and not metal co-ordination.

The interaction of a peptide spanning residues 147–172 of human SOUL with human Bcl-xL was also studied with 15N-1H HSQC NMR spectroscopy using two different forms of human Bcl-xL, the entire molecule lacking only the hydrophobic transmembrane domain after amino acid 209 and another truncated form lacking also amino acids 27–82 (Δ27–82). The results were comparable and indicated in both cases that there is an interaction of the 26-amino-acid SOUL peptide with Bcl-xL with a dissociation constant estimated to be 40–50 μM. These results were confirmed by SPR measurements and by preparing co-crystals of the complex and solving their three-dimensional structure. The new crystal structure revealed swapping of the first helix of the Bcl-xL dimer, a phenomenon associated with the presence of the Δ27–82 truncation. The amino acids participating in the interaction of the BH3 domain and protein were identified. The interactions are mostly hydrophobic, but a significant number of specific charged residue contacts were also found to be present.

When the SPR experiments performed to detect the interaction of the SOUL BH3 domain peptide were repeated using the entire SOUL molecule instead, no interaction was detected between ligand and analyte. This result might be explained by the fact that our findings predict a very drastic conformational change in the protein molecule to allow the portio of polypeptide chain involved in the contacts to adopt the conformation required for the contacts to be established. The last eight amino acids of the BH3 domain are not in a helical conformation in intact SOUL and, in addition, side chains that are important for the interaction point towards the interior of the molecule and are not available on the protein surface. Conformational changes in the anti-apoptotic members of the Bcl-2 family upon interaction with BH3 domains have been described [38], as well as changes in the interacting BH3 domains [41,46]. In addition, it has been shown that Bim, Bad and Bmf have intrinsically unstructured BH3 domains that undergo a localized conformational change upon binding to pro-survival Bcl-2 targets [48]. No important changes were found in Bcl-xL, but the changes in the BH3 domain are remarkable and are sufficient to explain why no interaction is observed with the intact SOUL molecule. These drastic modifications might require conditions that have not yet been found, but that should be explored further given the importance of this interaction in the functionality of the two proteins.

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for the interaction to take place, and (ii) the precise specificity of the interaction of the SOUL BH3 domain with different members of the Bcl-2 family.

AUTHOR CONTRIBUTION
Emmanuele Ambrosi produced the SOUL protein, prepared the crystals and solved the structure, and performed the related data analysis; Stefano Capaldi prepared Bcl-xL, crystallized the complex with the peptide and performed the NMR studies; Michele Bovi performed the SPR studies; Gianmaria Saccomani helped in the initial phases of SOUL protein expression, purification and crystallization; Massimiliano Perduca collected and processed the X-ray diffraction data; and Hugo Monaco directed the research and wrote the paper. All authors contributed to revising and improving the final version of the paper prior to publication.

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Interaction of SOUL protein BH3 domain with Bcl-xL


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SUPPLEMENTARY ONLINE DATA

Structural changes in the BH3 domain of SOUL protein upon interaction with the anti-apoptotic protein Bcl-xL

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SUPPLEMENTARY INFORMATION

X-ray structure of human SOUL

For the sake of simplicity, only the structure of the orthorhombic crystal form which diffracts to 1.6 Å resolution was described (in the main paper). At the end of this section, we will discuss a difference that was found in the hexagonal crystal form solved to 2.85 Å resolution. The final model of the orthorhombic crystal form of SOUL comprises 180 amino acid residues of both chains A and B, the two monomers present in the asymmetric unit of this crystal form. The maps do not show electron density for the first 18 residues and the last seven residues of both chains. The model contains 2974 non-hydrogen protein atoms, one phosphate and 395 water molecules. The conventional R factor is 17.5% and the free R factor is 19.7% (see Table 1 in the main paper). The R factors and rmsd values of Table 1 in the main paper were calculated with the program Phenix.refine [1]. The stereochemical quality of the protein model was assessed with the program PROCHECK [2]; 93.5% of the residues are in the most favourable region of the Ramachandran plot and the remaining 6.5% in the additionally allowed region. The two molecules in the asymmetric unit are not related by a non-crystallographic dyad and there are some minor differences between the two chains that can be explained by the packing of the molecules in the crystal (see below). The rmsd between the two main chains is 0.72 Å calculated over 180 Cα pairs of equivalent residues.

The resolution of the orthorhombic crystal form is quite adequate for the analysis of the structure of the solvent molecules within the β-barrel cavity of the protein (see Table 1 of the main paper), and we have identified the water molecules that bind in the interior of barrel in the same position in the two SOUL monomers of the asymmetric unit. Trp46 binds a water molecule at its Nε1 atom, and another water molecule, in proximity of the first, binds the NH1 atom of Arg132. Two other water molecules, in the proximity of the first two, bind the carbonyls of Ala133 and Gly44, the latter is external to the cavity. The four water molecules are close enough so that the group should be described as a cluster organized in a net of hydrogen-bonded solvent molecules. There is another cluster that involves three conserved water molecules positioned in the interior of the barrel (bound to the OG1 of Thr176 and the carbonyl of Val68), at the interface (bound to the carbonyl of Ser181) and outside (bound to the water molecule at the interface). Two other internal water molecules bridge the OG1 of Thr186 with the carbonyl of Lys84 and the N of Tyr110, and another pair is bound to the carboxy moiety of Asp130. The solvent molecules in the interior of the cavity are in contact with others at the interface which in turn are in the proximity of water molecules in conserved positions on the external surface of SOUL.

An interesting structural difference was observed in the hexagonal crystal form that diffracts to 2.85 Å and contains four SOUL monomers in the asymmetric unit. Two of these monomers (A and B) swap the first portion of their polypeptide chains, the portion running from amino acids 19 to 32, i.e. before the first strand of β-sheet. This structure swapping is not confirmed in the other two monomers, since the portion where it is expected to take place is disordered in the maps. This disorder is a clear indication of high flexibility of that loop and we have not attributed any particular significance to our observation that we think is a consequence of the molecular packing in the crystals.

Comparison of SOUL with HEBP1

The p22HBPs have been studied more extensively than the SOUL family. They are ubiquitously expressed but are extremely abundant in liver, have a cytoplasmic location and there is solid evidence that they bind haem and several porphyrins with micromolar Kd values [3]. The p22HBPs are highly homologous, monomeric and soluble, and bind metalloporphyrins, free porphyrins and N-methylprotoporphyrin with similar affinities. Two NMR structures of the same protein, murine p22HBP, have been published [4,5]. In both models, the 22-kDa monomer is described to fold as a distorted β-barrel flanked by two long α-helices arranged on one face. Using a 15N-1H HSQC titration experiment, the porphyrin-binding site of murine p22HBP was mapped and found to comprise a number of loops and one of the two α-helices with all of the residues participating in ligand binding located on a single face of the molecule [5].

Murine p22HBP has approximately 28% sequence identity with murine SOUL, which was reported to be a dimer in its apo form and to hexamerize upon haem binding with a dissociation constant in the nanomolar range [6]. It was reported further that His42, the only histidine residue present in the sequence, plays a crucial role and that its mutation abolishes haem binding. Comparison of ligand binding of murine p22HBP and murine SOUL reveals that binding has to take place in very different sites and therefore one has to confront the intriguing situation of two highly similar proteins binding the same ligand in a very different way.

The percentage of sequence identity between human SOUL and murine p22HBP is approximately 29 %, which supports the prediction that the two folds should be very similar to one another. In spite of this, all our attempts to solve the crystal structure of SOUL by molecular replacement, using the two available NMR structures as search probes, failed and the SOUL structure had to be solved using the alternative single isomorphous replacement method.

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The co-ordinates of the models, and the structure factors of SOUL and of the complex of human Bcl-xL with the peptide have been deposited in the PDB under accession codes 3R85, 3R8J and 3R8K.
After the final co-ordinates of SOUL became available, the model of chain A of SOUL was superimposed to that of chain B and the two lowest energy NMR structures available of murine p22HBP (PDB codes 2GOV and 2HV A; [4,5]), and the distances between equivalent α carbons were calculated. The results are represented in Supplementary Figure S1(A) as a function of the amino acid number. In the Figure the blue trace represents the differences between the two SOUL molecule models in the orthorhombic crystal asymmetric unit (A and B) and the black trace indicates the difference between the two murine p22HBP NMR structures. The area where the two SOUL molecules in the asymmetric unit differ more are the region before the first strand of β-sheet and the loop connecting strands C and D. The chains before the first strand are totally exposed to the solvent, whereas in the case of the connection of strands C and D the loop in chain A is in close contact with a symmetry related molecule and the equivalent area is in contact with the solvent in molecule B. We thus believe that these differences are simply a consequence of molecular packing in the crystal. Although there is more variability in the two NMR structures of murine p22HBP, the two structures are very similar in the region connecting strands C and D, which is also the region where both are most different from SOUL. This particular region of the SOUL molecule thus appears to be more variable than the rest of the molecule.

Supplementary Figure S1(B) is a stereo pair in which the model of SOUL A is superimposed to the two NMR models of murine p22HBP. In the Figure, SOUL is represented in blue, whereas the two models of murine p22HBP are in red and green. As the Figure shows there are areas of larger variability in the conformation of the proteins. The N-terminal region of the two NMR structures is very different and both differ substantially from the X-ray structure, which is due to the fact that this part of the chain is very flexible and becomes more structured in the crystals. A more significant difference is observed in the loop connecting strands C and D between the A chain of SOUL and the two NMR structures. This area of the molecule is, however, in different conformations in the two SOUL molecules of the asymmetric unit and so the differences observed with murine p22HBP probably simply reflect the high degree of flexibility of this area.

Overall the three structures are quite similar, although, as expected, the differences between the two NMR structures are somewhat smaller than those observed between them and the crystal structure of SOUL.

### Table S1  Main contacts between the SOUL BH3 domain and human Bcl-xL

Selected distances between the closest human Bcl-xL residues of molecule A and the SOUL BH3 domain in contact with it (labelled molecule E). The hydrophilic contacts are highlighted in bold.

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Figure S1  Comparison of the models of human SOUL and murine p22HBP

(A) rMsd between $\alpha$-carbon atoms of the SOUL models and the models of murine p22HBP: A chain compared with B chain of the orthorhombic form (blue); lowest energy NMR murine structure (PDB code 2GOV) compared with the other equivalent structure available of the same protein (PDB code 2HVA) (black). The green and red traces compare chain A of SOUL with the two NMR structures of murine p22HBP (PDB codes 2GOV (green) and 2HVA (red)). The strip at the bottom of the Figure represents the elements of secondary structure of SOUL. The blue colour identifies the buried residues, whereas white is used to indicate the exposed amino acids. (B) Stereoimage with the superposition of the three models. SOUL is in blue and the two NMR models of murine p22HBP are in red and green. The Figures of the models were prepared using the program PyMOL. (http://www.pymol.org).
Figure S2 Interaction of SOUL with haemin

(A) UV–visible spectra of haemin alone (broken line) and with ten times a molar excess of SOUL (dotted line) and BSA (solid line). (B) HSQC titration of 15N-labelled SOUL with haemin. The black spectrum corresponds to protein SOUL before the titration, whereas the red spectrum was recorded after the addition of four equivalents of haemin. No differences were detected between the two spectra.

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