

Oncogenic protein interfaces: small molecules, big challenges

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Abstract | Historically, targeting protein–protein interactions with small molecules was not thought possible because the corresponding interfaces were considered mostly flat and featureless and therefore ‘undruggable’. Instead, such interactions were targeted with larger molecules, such as peptides and antibodies. However, the past decade has seen encouraging breakthroughs through the refinement of existing techniques and the development of new ones, together with the identification and exploitation of unexpected aspects of protein–protein interaction surfaces. In this Review, we describe some of the latest techniques to discover modulators of protein–protein interactions and how current drug discovery approaches have been adapted to successfully target these interfaces.

High-throughput chemical screening

(HTS). A systematic way to measure the modulation of an interaction or a biological function against a given protein target by a large number of individual compounds.

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Protein–protein interactions (PPIs) form the molecular bases of many physiological processes in health and disease. In cancer, PPIs can have important regulatory roles; for example, in the cell division cycle or in cell signalling. The specificity and affinity of these interactions are critically important — changes in them can lead to cellular malfunctions, such as the uncontrolled cell growth that typifies cancer. Thus, it is not surprising that modulators of PPIs, whether as biological agents (such as antibodies) or as small-molecule synthetic compounds, are urgently being sought and developed by the pharmaceutical industry to treat an unmet medical need.

This interest in PPIs as drug targets has historically been tempered by the pervasive view that finding molecules, particularly small synthetic molecules (those <500 Da), that will modulate PPIs is irredeemably hampered by the large, flat, featureless areas of protein surfaces that form the interface — an interface that is thought to be ‘undruggable’ (REFS 1–4). This has often been in spite of compelling evidence from molecular, cellular and animal studies that such targets have an excellent potential for drug development. Nevertheless, major inroads into discovering such molecules have been made in recent years, with more than 12 small-molecule PPI modulators currently in clinical development (TABLE 1) and sales worldwide predicted to be in excess of US\$800 million within 5 years⁵.

As with other areas of drug discovery, a major challenge is to identify and validate physiologically relevant PPIs as cancer-related targets; this task is particularly challenging for many PPIs, as they can be transient in nature.

The most common tools for identifying PPIs have been genetic approaches, such as yeast two-hybrid, synthetic lethal, systematic RNA interference and co-evolution analyses; cell biology approaches, including the localization of proteins with fluorescent markers; proteomic approaches, such as affinity purification and mass spectrometry (MS) of complexes and protein arrays; and direct protein co-purification and immunoprecipitation. The past couple of years have also seen the development of highly sophisticated approaches to genome-wide prediction of interacting proteins, including studies that have identified more than 300,000 potential interaction pairs encoded in the human genome⁶. Of particular relevance to cancer is the genome-wide profiling of tumour samples by initiatives such as [The Cancer Genome Atlas](#) and the [International Cancer Genome Consortium](#).

Validating the physiological relevance of a PPI to cancer facilitates the initiation of a drug discovery programme. The pathways in a typical drug discovery programme are outlined in FIG. 1; the nature of the protein target and the available resources will dictate which pathways are followed. FIGURE 2a illustrates how different approaches are being applied to a single protein target in the hunt for PPI modulators. As well as traditional high-throughput chemical screening (HTS), structure-based discovery approaches are now common. In addition to the challenges described above, these approaches face another challenge that is specific to PPIs. There is a wide gulf between the number of experimentally determined protein structures and the

Key points

- More than 300,000 interaction pairs have already been identified in the human genome; therefore, modulating protein–protein interactions (PPIs) has a huge potential for therapeutic intervention in cancer.
- The refinement of existing bioassay techniques and the development of new ones, together with the identification and exploitation of unexpected aspects of PPI surfaces has led to more than 12 small-molecule PPI modulators making it to the clinic in recent years.
- Advances in the structural and biological understanding of the PPI to be modulated are shaping the compound libraries being used for PPI screening; for example, three-dimensional shape diversity, chirality and pharmacokinetic properties are now included in the library design.
- Improvements in data handling, robotics and miniaturization of assay technology has allowed increasing numbers of compounds to be evaluated against a given protein target using high-throughput screening (HTS) methods.
- The availability of cheap computational power has led to the routine use of virtual screening in the discovery of small-molecule PPI modulators, either as a stand-alone method to identify potential hits or as a prelude to HTS. Active compounds that are identified from the virtual screening process then undergo a traditional medicinal chemistry optimization process.
- Sensitive methods such as protein-based or ligand-based NMR, X-ray diffraction and surface plasmon resonance allow the use of fragments of low affinity and specificity to be used for the discovery and development of small-molecule PPI modulators.
- The therapeutic class of biologicals, which includes antibodies, peptides and aptamers, generally has high target specificity and potency. Biologicals are particularly useful for PPI modulation, as they can be readily tuned to bind to a large variety of protein surfaces and are often used as a starting point for PPI drug discovery programmes.

number of proteins that have been implicated in human cancers. This is especially true for protein–protein complexes, which represent less than 0.5% of the structures deposited in the [Protein Data Bank](#)⁶. Thus, computational methods, such as protein docking and homology modelling, that are combined with low-resolution experimental approaches (for example, mutagenesis, chemical crosslinking, hydrogen–deuterium exchange, cryo-electron microscopy, fluorescence spectroscopy and small-angle X-ray scattering) have proved to be useful when the structure of the protein complex of interest is not available.

Therapeutic proteins, such as monoclonal antibodies, which target PPIs that are accessible to the extracellular environment, have proved to be extremely successful in the clinic during the past decade (see [Supplementary information S1](#) (table)). However, biologicals (protein-based drugs, peptides, aptamers and other macromolecules; see [Supplementary information S2, S3](#) (tables)) can be expensive and are rarely cell-permeable or orally available. Hence, the typically lower cost of treatment, ease of use and potential to target intracellular PPIs all drive the current substantial interest in the discovery of small synthetic molecule PPI modulators. Several recent reviews on the topic of PPI modulators are available, and these cover areas such as PPI characterization⁷, the use of specific techniques for PPI drug discovery^{3,8–11} and a comparison of the properties of PPI modulators with enzyme inhibitors¹². In this Review, we highlight recent technological advances in the field, with a particular emphasis on early stage drug discovery — chiefly focused on small molecules, including peptides and

aptamers — and we illustrate these with examples of cancer-related PPIs being targeted for therapeutic intervention.

PPIs and protein–protein interfaces

The term ‘protein–protein interface’ potentially covers an enormous range of different, physiologically relevant interaction surfaces within biological systems. Interactions between monomers in multimeric proteins, interactions in multi-molecular complexes, recognition of activation signals (such as phosphorylation, proteolysis and receptor–hormone interactions) and many antibody–antigen binding events all involve the formation of protein–protein interfaces with extremely diverse affinities and specificities. The huge diversity of PPIs and corresponding protein–protein interfaces means that it is difficult to generalize about the characteristics of such sites. For most people, the term protein–protein interface brings to mind the large surfaces that are present in globular protein complexes, receptor–hormone complexes and antibody–protein interactions, much like the interleukin-2 (IL-2)–interleukin-2 receptor subunit- α (IL-2R α) interface shown in [FIG. 2b](#), and the term ignores those surfaces that more closely resemble a classic protein–small-molecule interface, such as the BCL-X_L–BAX BH3 domain interface that is shown in [FIG. 2a](#). This perception is unfortunate, as many key interactions between proteins are transient events that occur through small interfaces that are in some ways more akin to ligand binding sites. Some of the aspects of interfaces in PPIs and the characteristics that facilitate their targeting during drug discovery are outlined in [BOX 1](#).

Although many protein interfaces lack clearly defined pockets, there is a bias in the positioning of physiological ligand binding sites to be close to or even directly associated with the edge of protein–protein contacts¹³. Inspection of a set of 1,611 protein–protein complexes showed that in those protein complexes with a known ligand binding site about one-half of the sites contained amino acids from both proteins. In most cases, binding pockets were found to be within 6 Å of the protein–protein interface. Therefore, these sites have the potential to provide convenient anchor points for the development of compounds that interfere with the protein–protein interface.

All modulators of protein activity, including PPI modulators, fall into two broad functional classes. Orthosteric PPI modulators directly target the interacting interface of one of the protein partners and, in doing so, they alter the interaction, usually by preventing the interaction occurring but in some cases by stabilizing the protein complex. By contrast, allosteric PPI modulators bind at a site that is distinct from the interaction interface and, in doing so, they modify their protein target — normally by inducing a conformational change that subsequently alters the formation of the PPI. Thus, approaches to target both types of modulation should be considered. Unless otherwise specified, the small-molecule examples that are discussed in this Review are orthosteric PPI modulators.

Small-angle X-ray scattering

A technique that is used to measure the average size and shape of a protein or a protein complex in solution, in the resolution range of 1–25 nm.

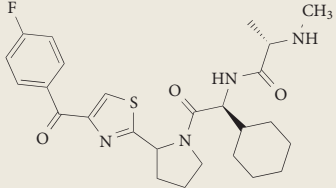
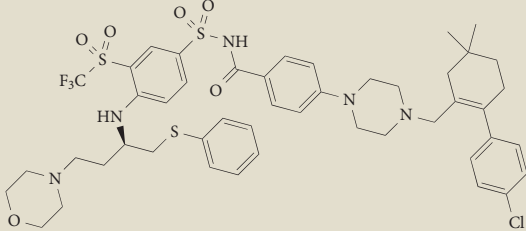
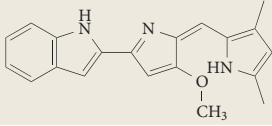
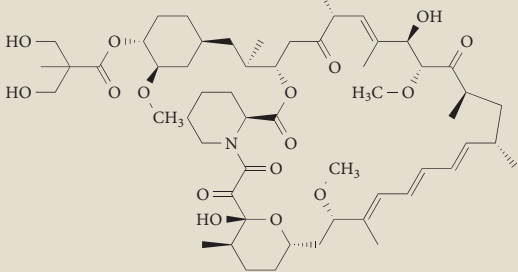
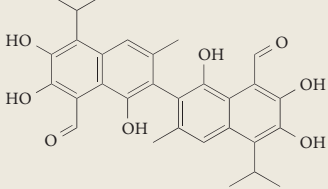
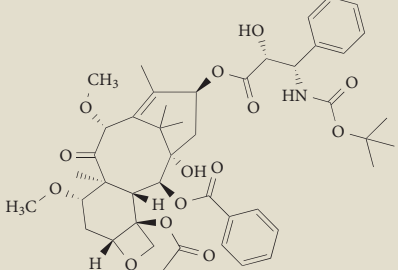
Orthosteric PPI modulators

A site on the protein–protein interaction surface of one of the protein partners is targeted by an orthosteric PPI modulator to alter the interaction, usually by preventing the interaction occurring but in some cases by stabilizing the protein complex.

Table 1 | Recent examples of small-molecule modulators of cancer-related protein–protein interactions in the clinic

Cancer type	Target PPI	Chemical structure	Molecular mass (cLogP)	Generic name (type of modulator)	Clinical phase	Refs
AML, CML, sarcoma and solid tumours	MDM2–p53		728 Da (10.6)	RG7112 or RO5045337 (orthosteric inhibitor)	Phase I	29
AML, CML, pancreas and solid tumours	β-catenin–CREB-binding protein		549 Da (6.1)	PRI-724 or ICG-001 (orthosteric inhibitor)	Phase I/II	143, 144
AML, lymphoma and solid tumours	IAP family–SMAC		807 Da (2.8)	TL32711 (orthosteric inhibitor)	Phase I/II	145
Bladder	Tubulin-α–tubulin-β		817 Da (5.4) free base; 1117 Da, tartrate salt	Vinflunine ditartrate or PM391 (allosteric inhibitor)	In the clinic	146, 147
Breast	Tubulin-α–tubulin-β		730 Da (1.2) free base; 826 Da, mesylate salt	Eribulin mesylate or E7389 (allosteric inhibitor)	In the clinic	148, 149
Breast	Tubulin-α–tubulin-β		507 Da (3.1)	Ixabepilone (allosteric stabiliser)	In the clinic	150, 151

Table 1 (cont.) | Recent examples of small-molecule modulators of cancer-related protein–protein interactions in the clinic

Cancer type	Target PPI	Chemical structure	Molecular mass (cLogP)	Generic name (type of modulator)	Clinical phase	Refs
Breast, pancreas and solid tumours	IAP family–SMAC		501 Da (3.6)	LCL161 (orthosteric inhibitor)	Phase I/II	152, 153
Breast, colorectal, GIST, lung, multiple myeloma and solid tumours	BCL-2–BAX and BAK BH3 domain		975 Da (12.4)	ABT-263 (orthosteric inhibitor)	Phase I/II	154
GIST and solid tumours	BCL-2 family–BH3 domain		317 Da (4.0)	GX15-070 (orthosteric inhibitor)	Phase II	155–157
Haematological malignancies and solid tumours	mTOR–FKBP12		1030 Da (7.5)	Temsirolimus or CCI-779 (allosteric inhibitor)	In the clinic	158, 159
Lung	BCL-2 family–BH3 domain; BCL-2–beclin 1; and BCL-X _L –beclin 1		519 Da (6.1)	(-)-gossypol (orthosteric inhibitor)	Phase I/II	160, 161
Lung and prostate	Tubulin- α -tubulin- β		836 Da (5.4)	Cabazitaxel (allosteric inhibitor)	In the clinic	162

AML, acute myelogenous leukaemia; cLogP, calculated LogP; CML, chronic myelogenous leukaemia; GIST, gastrointestinal stromal tumour; IAP, inhibitor of apoptosis; PPI, protein–protein interaction; SMAC, second mitochondria-derived activator of caspases.

Methods of monitoring PPI disruption

Monitoring the effect of PPI modulators can require quite different experimental approaches, depending on the specific PPI. Clearly, the most important aspect of PPI modulation is whether there is a physiologically

relevant effect on the system that is being targeted. Although the direct binding of compound to protein or the impact of compounds on the stability of the protein complex are important, an assay of biological activity is ultimately going to be the defining result:

Isothermal calorimetry (ITC). A way of directly measuring the heat that is released or absorbed in a sample upon complex formation (that is, protein–protein or protein–ligand complex formation).

Surface plasmon resonance (SPR). A technique that measures the interaction of molecules through changes in the refractive index that occur when ligands that are bound to a surface interact with another molecule in solution.

Microscale thermophoresis A technique that measures changes in the mobility of a protein in solution along an induced temperature gradient upon ligand binding or complex formation.

Fluorescence resonance energy transfer (FRET; also known as Förster resonance energy transfer). An assay technique that is used to measure interactions such as protein–protein interactions. A measurable signal is generated when an acceptor chromophore and a donor chromophore that are attached to two components of a system are stabilized in close proximity and are able to transfer excitation energy from the acceptor to the donor.

Amplified luminescent proximity homogeneous assay screen (AlphaScreen; PerkinElmer). A non-radioactive assay that measures the interactions between two components attached to luminescent donor and acceptor beads. A measurable signal is generated when an acceptor bead and a donor bead are held in close proximity through the attached molecules.

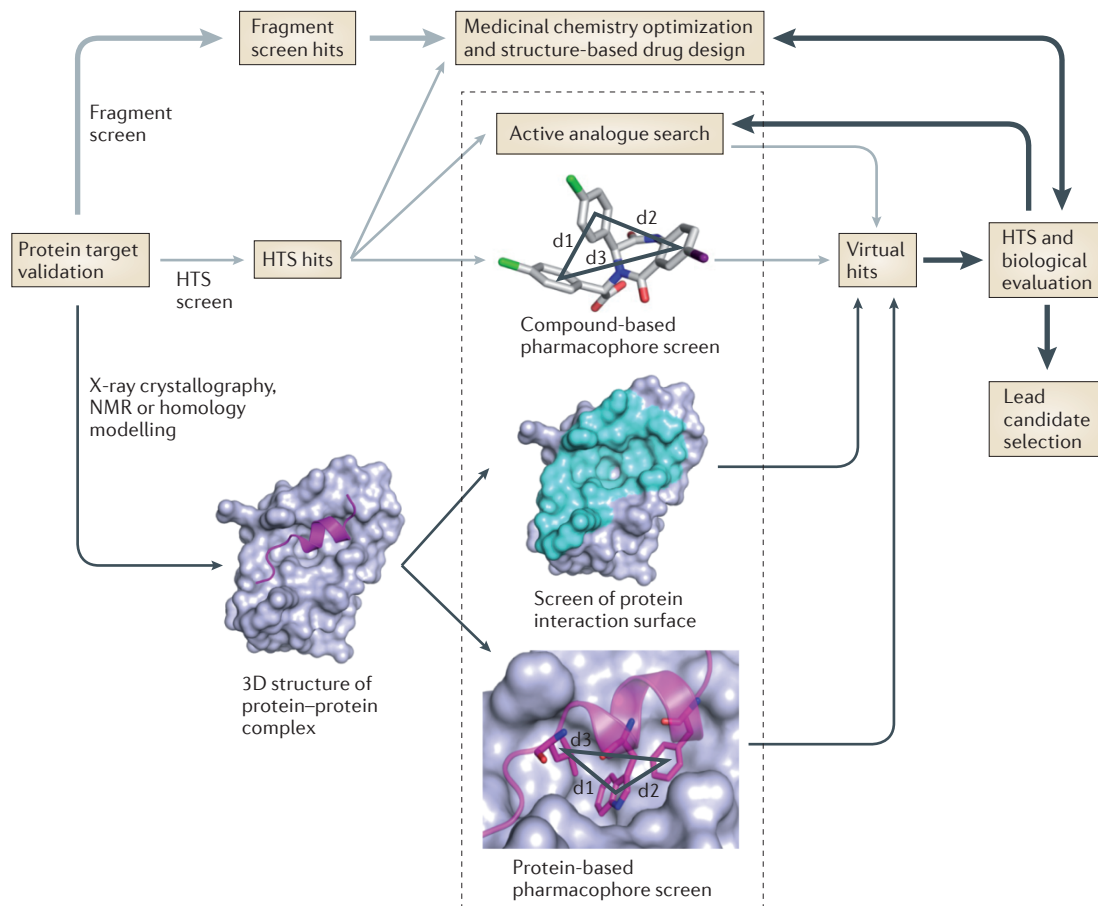


Figure 1 | The drug discovery process and protein–protein interactions. Once the protein target has been validated, screening of fragment or small-molecule compound libraries is undertaken to identify ‘hits’ that bind to the protein and modulate the protein–protein interaction. The virtual screening (VS) methodologies are enclosed in the dashed box. It is common for combinations of fragment screening, high-throughput screening (HTS) and VS to be done in parallel, with the data being fed into the structure-based drug design and medicinal chemistry optimization process. 3D, three-dimensional.

does the PPI modulator change the biology? However, bioassays have the disadvantage that they are often relatively insensitive: they need compounds with low micromolar activity or higher before an effect on the assay is observed. Although this is suitable for later stage development, identifying initial ‘hit’ compounds of relatively low potency requires more sensitive techniques than the typical bioassay can provide.

Screening for PPI modulators is mostly carried out in the same way as screening for any inhibitor of protein function. Various biophysical techniques can be used to directly measure the interaction between a drug and the target protein, such as MS, NMR, isothermal calorimetry (ITC), surface plasmon resonance (SPR), microscale thermophoresis, fluorescence anisotropy or X-ray crystallography. Examples of these approaches being used to characterize the effects of molecules that regulate cancer-related PPIs have been published^{8,14–16}, and some are described below: in many cases, identical PPI targets were explored using completely different techniques by independent research groups.

HTS approaches to modulate PPIs

HTS is a well-established strategy that has been used for several decades by industry and, more recently, by academic groups to identify new small molecules that can modulate the activity of a protein target^{14,17,18}. Continued improvements in data handling, robotics and miniaturization of assay technology has allowed increasing numbers of compounds from in-house libraries and commercial suppliers to be tested efficiently, reliably and with a relatively low cost per assay^{14,17}. Despite this impressive history in classical drug discovery, HTS has had mixed results in identifying new leads for modulating PPIs¹, partly owing to the difficulties with developing suitable high-throughput assays to assess the PPI to be modulated *in vitro*. Despite the potential of robust assay methods such as fluorescence resonance energy transfer (FRET; also known as Förster resonance energy transfer), amplified luminescent proximity homogeneous assay screen (AlphaScreen; PerkinElmer), enzyme-linked immunosorbent assay (ELISA) and fluorescence polarization (FP), which involve a measurable change in fluorescence or absorbance upon formation of a PPI, or the use of highly

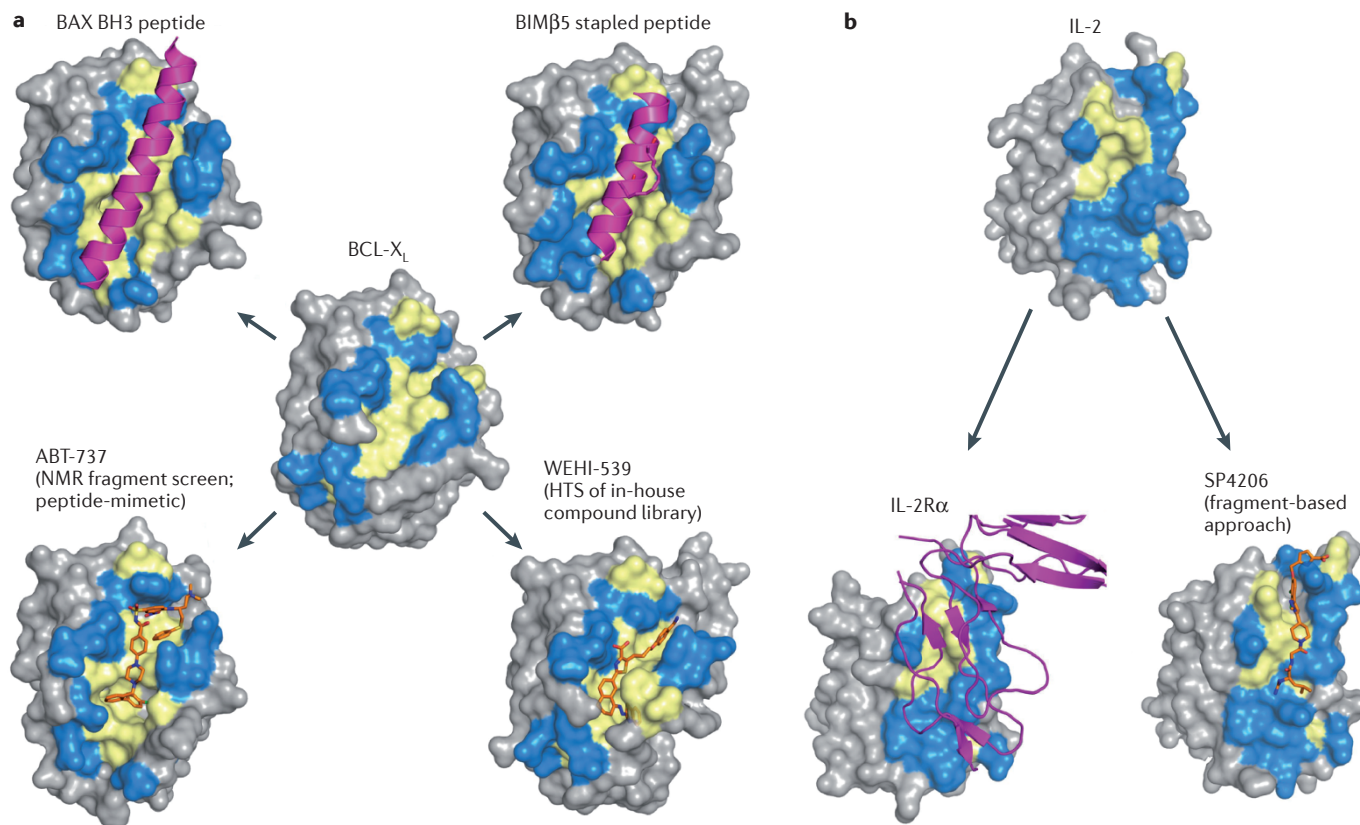


Figure 2 | Cancer-related protein–protein interaction targets BCL-X_L and IL-2. BCL-X_L–BAX BH3 domain is an example of a ‘helix in groove’ interaction, and interleukin-2 (IL-2)–interleukin-2 receptor subunit-α (IL-2Rα) is an example of a flat interaction surface. In the figure, the protein to be inhibited is shown as a grey surface and the residues that interact with the natural protein partner are shown in blue (polar residues) and yellow (hydrophobic residues). The same view and colour scheme has been used to highlight the conformational change in the protein interaction surface when it is bound to the interacting partner protein, peptide or small molecule. **a** | The BCL-X_L crystal structure (middle; [Protein Data Bank](#) (PDB) ID: [1R2D](#)¹³⁵); the complex of BCL-X_L with the BAX BH3 peptide (magenta; top left; PDB ID: [3PL7](#) (REF. 130); half-maximal inhibitory concentration (IC₅₀) = 184 nM¹³⁶); the complex of BCL-X_L with BIMβ5, which is a stapled BIM BH3 peptide (magenta, with the linker (‘staple’) shown as sticks; top right; PDB ID: [2YQ6](#) (REF. 137); dissociation constant (K_D) = 45 nM¹³⁷); the complex of BCL-X_L with ABT-737 (shown as sticks, coloured by atom type; bottom left; PDB ID: [2YXJ](#)¹³⁸; K_D = 0.4 nM⁹¹; IC₅₀ = 5 nM⁹¹); and the complex of BCL-X_L with WEHI-539 (shown as sticks, coloured by atom type; bottom right; PDB ID: [3BZR](#)⁹¹; K_D = 0.6 nM⁹¹; IC₅₀ = 1.1 nM⁹¹). **b** | The IL-2 crystal structure (top; PDB ID: [1M47](#) (REF. 94)); the complex of IL-2 with IL-2Rα (magenta; bottom left; PDB ID: [1Z92](#) (REF. 139); inhibition constant (K_i) = 10.5 nM¹⁴⁰); and the complex of IL-2 with the inhibitor SP4206 (shown as sticks, coloured by atom type; bottom right; PDB ID: [1PY2](#) (REF. 141); K_i = 68.8 nM¹⁴⁰). HTS, high-throughput chemical screening.

sensitive radiometric assays such as scintillation proximity assays and FlashPlate (PerkinElmer)¹⁷, the problem often lies not in the availability of an assay method but in the recreation of the PPI in an *in vitro* setting. Components can be difficult to isolate in a fully functional form on the scale that is required for HTS, owing to various biological complexities, such as essential cofactors, membrane requirements, multiprotein complexes and so on. Often, one or more protein partners are present as a fragment (for example, as a peptide or a specific domain) in assays, and inhibition of the PPI *in vitro* might not be recapitulated with the full-length proteins. A greater understanding of the quality of the compound libraries that are used during HTS is also developing as groups merge experimental results from different screening campaigns and investigate common trends. This has led to a better understanding of the chemistries of compounds that frequently lead to false

positives and ultimately lead to resources being wasted; in particular, this is exemplified by Baell and Holloway’s ‘pan assay interference compounds’ (PAINS; see below)¹⁹. Moreover, researchers at F. Hoffmann–La Roche identified zinc-contaminated compounds as giving a false-positive signal in their screen directed against RAS²⁰. On a more positive note for the development of RAS pathway inhibitors, one group has identified a series of benzimidazole compounds through an AlphaScreen assay that selectively bind to the prenyl-binding pocket of phosphodiesterase 6D (PDEδ; also known as GMPPDEδ) with nanomolar affinity, and these disrupt the KRAS–PDEδ PPI²¹.

Helix–groove binders. The greatest success for HTS so far has been with PPIs in which a helix of one protein binds into a groove of the interacting partner; for example, the BAX BH3 domain interacting with BCL-X_L (REF. 22)

Box 1 | Hot spots and O-rings

Globular protein interfaces typically have 1,200–2,000 Å² of buried surface, which is similar in composition to the available surface of typical monomeric proteins. The distribution of polar and lipophilic groups is comparable to a normal protein surface⁷. Packing of residues in the protein interface is usually tight, with a similar density to that of atoms in the hydrophobic core of a protein¹²⁹. This is increased by interfacial water molecules that form polar interactions between the proteins. There is, however, a high degree of variation in the 'flatness' of the interface, with many interfaces having quite well-defined binding sites, such as the helix-binding groove on BCL-X_L (FIG. 2a). Others have no immediately obvious site of interaction (such as the interleukin-2 interface (FIG. 2b)). However, painstaking experimental studies using mutagenesis and affinity measurement have shown that many of these apparently featureless interfaces contain 'hot spots', which are small clusters of residues that contribute a significant proportion of the overall binding energy in the interaction¹³⁰. The residues in hot spots also show a higher degree of flexibility and motion than the rest of the interfacial residues, which increases their potential for use as drug target sites¹³¹. This adaptivity provides a mechanism for optimizing the fit of the protein–protein interaction while at the same time giving a greater opportunity for small-molecule inhibitors to bind to the target protein, thereby potentially inducing the formation of a new pocket with more typical druggable features. Hot spots are usually ringed by a layer of residues (the 'O-ring' (REF. 132)) that almost completely exclude solvent, which means that the hot-spot interactions are driven solely by the residues themselves, without any bridging solvent molecules. Around the O-ring are the 'rim' residues, which form the periphery of the interface and are only partially buried from solvent. FIGURE 4 shows these key features on the structure of the human growth hormone binding protein (GHBP; also known as the extracellular GHR domain)–growth hormone (GH) complex¹³³. Hot-spot residues were defined through mutagenesis and binding studies as providing approximately 85% of the total binding energy of the interaction¹³⁴.

(FIG. 2a) and p53 interacting with MDM2 (REF. 23). The α -helix-binding grooves are surface pockets that are more akin to a classic enzyme–substrate site because of their well-defined and pronounced clefts compared to other types of globular protein interfaces (BOX 1).

The BCL-2 family of proteins are necessary for the regulation of apoptosis or programmed cell death. The family is composed of two related groups — the pro-survival proteins, such as BCL-2, BCL-X_L, BCL-W, BCL-B, MCL1 and A1 (also known as BCL-2A1), and the pro-apoptotic proteins, such as BAX, BAK, BOK, BAD, BIM, PUMA and NOXA²². Inhibitors of this family of apoptosis regulators were identified by NMR²⁴, MS and FP²⁵, and computational screening²⁶. The Nutlins — imidazoline-based compounds that inhibit the interaction between the p53 tumour suppressor and MDM2, which is one of the key repressors of p53 activity — were discovered and developed through screening the interaction of the mouse MDM2 homologue with p53 by SPR²⁷. The proof-of-concept for this PPI target came from screening 12-mer and 15-mer phage display libraries²⁸. The minimal length of the p53-derived peptide that was required to retain micromolar inhibition of MDM2 was determined. In addition, modification of the minimal length peptide to include a 6-chloro substituent on the tryptophan residue increased inhibition. These drugs were successfully optimized into mid-nanomolar inhibitors (half-maximal inhibitory concentration (IC₅₀) values in the 100–300 nM range) using structure-based drug design (SBDD)²⁸. Further investigation around this series led to the F. Hoffmann-La Roche candidates RG7112 (REFS 23,29) and RG7338 (REF. 30). These and two other MDM2–p53 inhibitors are in clinical trials:

RG7112 (REF. 29) (two active Phase I trials and five completed trials under the name RO5045337; targeting various leukaemias and liposarcoma), RG7338 (REF. 30) (currently in three Phase I studies under the name RO5503781; targeting solid tumours), SAR299155 (REFS 28,31) (currently undergoing Phase I clinical trials; targeting acute myelogenous leukaemia and solid tumours) and MK-8242 (REF. 28) (also known as SCH900242; currently in two Phase I studies; targeting acute myelogenous leukaemia and solid tumours) — see ClinicalTrials.gov. A different HTS approach was taken by Johnson & Johnson, in which they used differential scanning fluorimetry (DSF) to screen 338,000 compounds; from these, 116 benzodiazepinedione compounds were selected for further analysis using a fluorescent peptide displacement assay that was designed to detect specific inhibitors of the MDM2–p53 interaction³². They used an MDM2 peptide consisting of residues 17 to 125, which included the minimal p53 binding domain³³. A hit-to-lead optimization programme established extensive structure–activity relationships (SARs) and the importance of the absolute stereochemistry for their lead series, but these compounds showed weak cellular activity and were not developed further.

Another interesting example of this class of PPI being addressed by HTS is provided by the 14-3-3 proteins³⁴. These eukaryotic adaptor proteins, which are involved in the regulation of cell cycle control, signal transduction, protein trafficking and apoptosis, act as functional dimers, with each monomer containing an amphipathic groove that accommodates interaction motifs on their partner proteins. In one study, a surface-based fluorescence assay of 14-3-3 interacting with a plant (*Nicotiana plumbaginifolia*) plasma membrane H⁺-ATPase-2 (PMA2) was used to screen a 37,000-compound library³⁵ for compounds that stabilized the 14-3-3 dimer. By contrast, competitive inhibitors that block the amphipathic groove and prevent formation of the active 14-3-3 dimer have been identified through the screening of a hybrid peptide and small-molecule library³⁶. This demonstration that the same PPI can be successfully addressed by modulators that block or increase the protein interaction makes it clear that assay development for PPI HTS programmes requires extremely careful planning.

Non-helical-groove binders. There are now excellent examples of small-molecule inhibitors that have been developed to compete with non-helical-groove binders. For example, the X-linked inhibitor of apoptosis (XIAP) has a downstream role in the apoptosis pathway and regulates the expression of the 'executor' cysteine–aspartic acid proteases (caspases)³⁷. Antagonists of XIAP were identified by an enzymatic derepression assay of caspase 3 function³⁸. They were confirmed to be specific for inhibiting the binding to caspase 3 by ELISA assays of the interaction of XIAP with endogenous XIAP inhibitors, such as second mitochondria-derived activator of caspases (SMAC). SMAC is known to interact with a deep groove in XIAP in an extended conformation³⁹. Other XIAP antagonists were identified by NMR studies of XIAP–peptide complexes, with binding affinities of compounds measured by FP⁴⁰.

Structure-based drug design

(SBDD). An approach that uses the three-dimensional structure of a protein or a protein complex to guide the development of ligands.

Differential scanning fluorimetry

(DSF). An assay that is used to measure the thermal stability of proteins through the binding of a hydrophobic, fluorescent dye to partially unfolded proteins. Changes in the temperature of unfolding in the presence of compounds reflect the stability of a protein–ligand complex.

Structure–activity relationships

(SARs). Relationships between the chemical or three-dimensional structure of molecules and their biological activity.

Medicinal chemistry optimization

The process in which chemists synthetically modify lead compounds to improve therapeutic properties such as efficacy and specificity for the target protein, as well as pharmacological properties such as bioavailability and absorption, distribution, metabolism and excretion (ADME) or toxicity.

In silico combinatorial chemistry

The virtual synthesis of a large number of chemical or biological entities using a small set of reagents together in all combinations. The resultant libraries are used for virtual or *in silico* screening.

Calculated LogP

(cLogP). The calculated base-10 logarithm of the relative partition of a compound in an organic phase (octanol) versus an aqueous phase (water), which is usually reported at 25 °C. cLogP values are generally reliable, and the parameter is used as an indicator of the solubility of the compound.

Polar interactions

The interactions between two chemical groups that both contain an electric dipole or multipole moment.

Hydrophobic interactions

Interactions of nonpolar (unionized and uncharged) regions of molecules with each other. The interactions have contributions from both dispersion forces and van der Waal's effects; they are entropy-driven through solvent exclusion.

An interesting recent example of the combined use of HTS and structural biology to find potent PPI inhibitors is that of the disruption of the complexes that are formed by mixed lineage leukaemia (MLL) fusion proteins and the tumour suppressor protein menin, which are complexes found in the development of human acute leukaemias⁴¹. A peptide fragment of MLL was shown to interact with menin in a deep groove, where it adopts a U-shaped conformation with a single β -turn. A collection of 49,000 small molecules was screened using an FP assay to target the complex⁴². The most potent compound ($IC_{50} = 1.9 \mu\text{M}$) that was identified belonged to the thienopyrimidine chemical class. Initial medicinal chemistry optimization led to a more potent compound, MI-2, which inhibited the interaction with an IC_{50} of 446 nM. The crystal structure of menin that was complexed with MI-2 showed that the inhibitor mimicked many of the key interactions that MLL has with menin, and it provided the basis for the development of a potent nanomolar inhibitor (dissociation constant (K_d) = 22 nM) that showed strong inhibition of cell proliferation and differentiation in leukaemia cells in which *MLL* is translocated.

Computational approaches to modulating PPIs

Virtual screening (VS; also known as *in silico* screening) is the process of searching a computer-based collection of chemical entities that have the greatest potential to interact with the protein target⁴³ and, in the context of this Review, to modify a specific PPI. VS can be used as a stand-alone method to identify potential hits or as a prelude to HTS, which is still a relatively expensive process (FIG. 1). Using VS before HTS is beneficial because it allows a more focused and enriched compound set to be evaluated in the bioassay. The *in silico* compound library can be derived from several sources, such as proprietary in-house compound collections, commercial and non-commercial supplier databases (see [Supplementary information S4](#) (table) and Further information for examples) and even sets of 'virtual' compounds that arise from *in silico* combinatorial chemistry. It is now common for *in silico* compound libraries to comprise millions of compounds. Compounds that contain undesirable physicochemical properties are typically removed from the *in silico* compound library by applying filters; for example, excluding all metal-containing compounds, known promiscuous compounds (PAINS)¹⁹ and those with a molecular mass above 1,000 Da or a calculated LogP (cLogP) >6. The nature of the search query for VS depends on whether the structure of the target protein is known (FIG. 1). When the structure of the target protein is available (from X-ray crystallography, NMR or homology modelling) the interaction surface can be searched (screened) using a computational algorithm that assesses the ability of each compound in the *in silico* library to interact with the experimentally determined protein surface 'hot spots' (BOX 1 describes hot spots and the painstaking experimental methods that are used to identify them). This process is also known as 'protein-compound docking'. Given the plastic nature of protein-protein interfaces and the availability of

cheap computational power, the current trend is to screen ensembles of potential protein conformations rather than a single protein model. These ensembles can be derived from multiple crystal structures, from snapshots of molecular dynamics simulations or from NMR ensembles^{44–47}. The fit of the compound to the protein interaction surface is evaluated or 'scored' on the basis of criteria that include shape complementarity, polar interactions, hydrophobic interactions and van der Waal's interactions. The compound score is used to rank the quality of the interaction of each compound with the target site. An excellent review of the methodology of protein-compound docking, docking software and scoring algorithms has recently been published⁴³. The ranked compounds from the VS process must then be assayed for their ability to modulate the target PPI before undergoing a traditional medicinal chemistry optimization process. The number of compounds that are selected for biological screening depends on the resources (for example, funds and facilities) of the investigators (for example, those in academic research or working in industry) and the throughput of the bioassay.

Numerous early drug discovery examples of VS for modulators that targeted cancer-related PPIs have been published, including for XIAP-caspase 9 (REF. 48), the BCL-2-BAK BH3 domain⁴⁹ and the urokinase plasminogen activator receptor (uPAR)-urokinase-type plasminogen activator (uPA) interaction⁴⁵.

XIAP is upregulated in many cancers, including acute myeloid leukaemia, as well as breast, pancreas, prostate and solid tumours^{11,50} (TABLE 1). In the case of XIAP, the compound embelin (see [Supplementary information S4](#) (table)) was identified as a cell-permeable, low molecular mass inhibitor of the XIAP-caspase 9 interaction⁴⁸. The program DOCK⁵¹ was used to virtually screen a library that comprised 8,000 structurally diverse small molecules against a model of the SMAC protein binding site that was extracted from a crystal structure of the XIAP BIR3 domain-SMAC complex. The authors used their in-house scoring function, 'X-score', to rank the compounds and the top 200 were considered to be potential inhibitors of the XIAP BIR3 domain. From the top 200 ranked compounds, 36 were obtained and their affinities were determined in an FP binding assay. The authors did not disclose how these 36 compounds were chosen over the other 164. Although the authors stated that five of the 36 compounds screened in the FP binding assay showed affinity for the XIAP BIR3 domain, data were reported for only one compound. Embelin was reported to be the strongest binder, with an IC_{50} of 4.1 μM (see [Supplementary information S4](#) (table)), which was comparable to the binding affinity of the natural 9-mer SMAC peptide ($IC_{50} = 2.8 \mu\text{M}$)⁴⁸. Having established that embelin bound to the target XIAP protein, the ability of the compound to inhibit cell growth in prostate cancer cells (PC-3 and LNCaP) and to induce apoptosis through the activation of caspase 9 in PC-3 cells was evaluated⁴⁸. Embelin inhibited PC-3 and LNCaP cell growth ($IC_{50} = 3.7 \mu\text{M}$ and 5.7 μM , respectively) and induced apoptosis in PC-3 cells via the activation of caspase 9 (a 10-fold and 20-fold increase in

Pharmacophore

A description of molecular features for a binding interaction and their relative position in three-dimensional space. Pharmacophores can be derived from either the protein receptor site (protein-based pharmacophore) or from a superimposition of known compounds (compound-based pharmacophore).

the level of activated caspase 9 was induced by a 42-hour treatment with 20 μM and 40 μM embelin, respectively) in a dose-dependent manner. Identifying embelin as an XIAP BIR3 domain PPI inhibitor means that the compound could function as the starting point for a medicinal chemistry optimization programme to improve activity and pharmacokinetic properties.

Inhibitors of the anti-apoptotic protein BCL-2 are currently being used to treat various cancers, including breast cancer, chronic lymphocytic leukaemia, lymphoma, prostate cancer, small-cell lung cancer and solid tumours⁵² (TABLE 1). Small-molecule inhibitors of the BCL-2–BAK BH3 domain interaction were identified from the US National Cancer Institute DIS 3D database of 206,000 compounds⁴⁹. The BH3 α -helix binding cavity in a homology model of the BCL-2 protein was screened using DOCK⁵¹. The energy scoring function in DOCK was used to rank the compounds and the top 500 were considered to be potential BCL-2 inhibitors. Chemical samples of 80 of the top 500 compounds were requested from the US National Cancer Institute; how these 80 compounds were selected was not disclosed, but only 35 were available. These compounds were tested in an FP binding assay and seven were shown to have affinity for BCL-2; the strongest binder had an IC_{50} of 1.6 μM (see Supplementary information S4 (table))⁴⁹. The seven compounds were then investigated for HL-60 cell viability and proliferation (HL-60 is a human myeloid leukaemia cell line that expresses high levels of BCL-2 protein); for the inhibition of cell viability, six of the compounds had an IC_{50} < 50 μM and one compound inhibited cell growth with an IC_{50} of 4 μM ⁴⁹. BCL-2 is overexpressed in breast, prostate and other forms of cancer, and further optimization of these compounds through SBDD would be required to produce potential clinical candidates.

uPAR is highly expressed in most cancers⁵³ and the uPAR–uPA interaction has been implicated in tumour formation and progression⁴⁵. In a search for small molecules that are capable of inhibiting the uPAR–uPA interaction, the binding cavities from two uPAR crystal structures were screened against a library of nearly 5 million commercially available compounds⁴⁵. Seven different scoring functions were used to select the top 10,000 compounds. These 10,000 compounds were further virtually screened using a multiple protein conformer strategy, whereby an ensemble of 50 uPAR conformations that were generated by molecular dynamics simulations were searched using the programs AutoDock4 (REF. 54) and Glide (version 5.5; Schrödinger). The docked complexes were finally scored and ranked using Glide SP; the 250 highest ranked compounds were clustered by similarity and the highest scoring compound from the top 50 clusters was purchased and screened in an FP binding assay. Of the 50 compounds screened, three showed affinity for uPAR, with the strongest binder (IPR-456) having an inhibition constant (K_i) of 140 nM (see Supplementary information S4 (table))⁴⁵. The authors went on to show that IPR-456 inhibited both the uPAR–uPA PPI interaction and cell invasion in the breast cancer cell line MDA-MB-231, which overexpresses uPAR (IC_{50} for

inhibiting invasion estimated to be 30 μM). Although IPR-456 is a useful chemical tool compound to probe the role of the uPAR–uPA PPI interaction in blocking metastatic processes such as invasion, further medicinal chemistry optimization will be required to develop clinical candidates.

Pharmacophore searching. An alternative VS approach is to identify key features from one of the proteins involved in the interaction and represent this motif as a pharmacophore (FIG. 1). The protein-based pharmacophore can then be used to search the *in silico* compound library and identify compounds that satisfy the pharmacophoric elements. Each compound is scored against how well it matches these elements and ranked. The highest ranked compounds can then be assessed in an appropriate bioassay^{47,55–59}. An example of the protein-based pharmacophore approach is the identification of substituted 1,2,4-triazoles as inhibitors of the S100A10–annexin A2 protein interaction (see Supplementary information S4 (table))⁵⁵. This PPI was shown to be essential for the recruitment of macrophages to tumour sites and is implicated in the process of neo-angiogenesis⁶⁰. S100A10 mediates the recruitment of macrophages to inflammatory stimuli, while annexin A2 is an auto-crine factor that is involved in osteoclast formation and bone resorption⁶¹. S100A10 has recently been identified as a potential therapeutic target in colorectal cancer⁶². A pharmacophore model that represented the amino acid interactions made by the amino-terminal residues of annexin A2 to the S100A10 protein interface was generated from the crystal structure of the S100A10–annexin A2 complex. A library of 704,000 commercially available compounds was screened against the pharmacophore (see Supplementary information S4 (table)) and 586 compounds that fitted the pharmacophore were identified. These compounds were then docked into the defined annexin A2-binding site on the S100A10 protein surface using the program GOLD⁶³. One hundred and ninety compounds were selected on the basis of both the ranked pharmacophore fit and GOLD docking scores for screening in a competitive FRET assay. Three compounds were confirmed as inhibitors of the S100A10–annexin A2 interaction (IC_{50} values of 24 μM , 66 μM and 90 μM) and provided the basis for a medicinal chemistry optimization programme.

Even in the absence of structural information about a PPI, it is still possible to perform VS by using the chemical structures of known active compounds that have been identified using other approaches, such as HTS (FIG. 1). Key chemical features in the active compounds can be used to generate a compound-based pharmacophore (FIG. 1), which in turn can be used to search *in silico* compound libraries^{64–66} in a similar manner to that described above for protein-based pharmacophores. An example of the compound-based pharmacophore searching approach is the identification of tubulin polymerization inhibitors (see Supplementary information S4 (table))⁶⁵. Compounds that inhibit tubulin polymerization are allosteric PPI inhibitors, in that they do not bind at the

α -tubulin- β -tubulin heterodimer interface but they bind to different regions on the α -tubulin- β -tubulin heterodimer, and this binding in turn regulates tubulin oligomerization. Tubulin inhibitors have been used for many years¹¹ to treat cancers such as bladder cancer, breast cancer and prostate cancer, and some recent examples are given in TABLE 1. In the above computational approach example, a compound-based pharmacophore was generated from a training set of 21 compounds with IC_{50} values ranging from 1.2 nM to 6,000 nM for the inhibition of human oral squamous carcinoma KB cells using CATALYST software (version 4.10; *Accelrys*). The pharmacophore was used to screen a compound database of 130,000 compounds, and 1,000 compounds with the best fit to the pharmacophore features were examined further. Compounds were visually examined for any structural features that might sterically clash with the tubulin binding site, and these compounds were eliminated from the compound list. The remaining 142 compounds were screened against the KB cell line for their anti-proliferative activities; four compounds inhibited the KB cell line with $IC_{50} < 6 \mu M$ ⁶⁵. The most active compound (see Supplementary information S4 (table)) was also shown to be a tubulin inhibitor in an *in vitro* tubulin polymerization assay

with an IC_{50} of $4.4 \mu M$. The four compounds with anti-proliferative activity would provide an ideal starting point for a medicinal chemistry optimization programme.

The VS approaches described here are typically used in the early drug discovery phase to identify compounds with some activity for the protein target. The same computational methods have been applied for many years to the traditional cancer-related protein targets, such as G protein-coupled receptors, protein kinases and enzymes⁴³. VS should essentially be considered as an early lead generator, and compounds with activity for the protein target should necessarily undergo substantial medicinal chemistry optimization, such as scaffold hopping and analogue exploration.

Fragment-based design for modulating PPIs

Fragment-based screening uses a different but complementary approach to HTS. Fragments are low molecular mass compounds (<300 Da) that typically comprise one or two fused or linked ring systems with one or two functional groups^{67–76}. The rationale behind fragment screening is that the molecules in fragment libraries explore chemical space more efficiently and are more likely to form an optimal set of interactions with a target protein than the larger and more complex HTS compounds (FIG. 3). The downside of the simpler interactions is that the initial hits typically have a very low affinity ($K_D = 100 \mu M$ to 10 mM) and therefore require specialized and sensitive techniques to measure binding interactions⁷⁷. Fragments also tend to be more promiscuous than compounds that are identified through HTS, as the simpler chemistry is likely to form a match with multiple protein binding sites. Ideally, a structure-guided approach is used to develop a fragment hit to counter the issue of promiscuity. An emerging example of this is the development of inhibitors of replication protein A (RPA), which is involved in the DNA-damage response^{78,79}. Researchers at Vanderbilt University, Nashville, Tennessee, USA, determined the crystal structures for several fragment hits, which they then linked and optimized to develop a lead compound with a sub-micromolar K_D (REF. 79) (FIG. 3). Optimization of detection techniques and increased knowledge of methods for fragment growth and elaboration have meant that fragment-based screening is now a routine technique in the pharmaceutical industry and is increasingly being used in academia.

The small size, low affinity and binding promiscuity of fragments means that sensitive methods are needed to detect and measure interactions. Methods including protein-based or ligand-based NMR⁸⁰, X-ray diffraction^{81,82} and SPR^{83–85} are well-established for the detection of fragment binding⁷⁵. DSF, AlphaScreen, FRET, FP and microscale thermophoresis are among techniques that are being explored for measuring fragment binding^{14,86,87}. Traditional biochemical assays have also been adapted for fragment screening in an approach that has been dubbed ‘high concentration screening’. In this approach, fragments are tested in established biochemical assays for a given protein target but at a much higher concentration than normal assays (typically $250 \mu M$ to 5 mM)^{87,88}. Although activity in a bioassay is the gold standard for

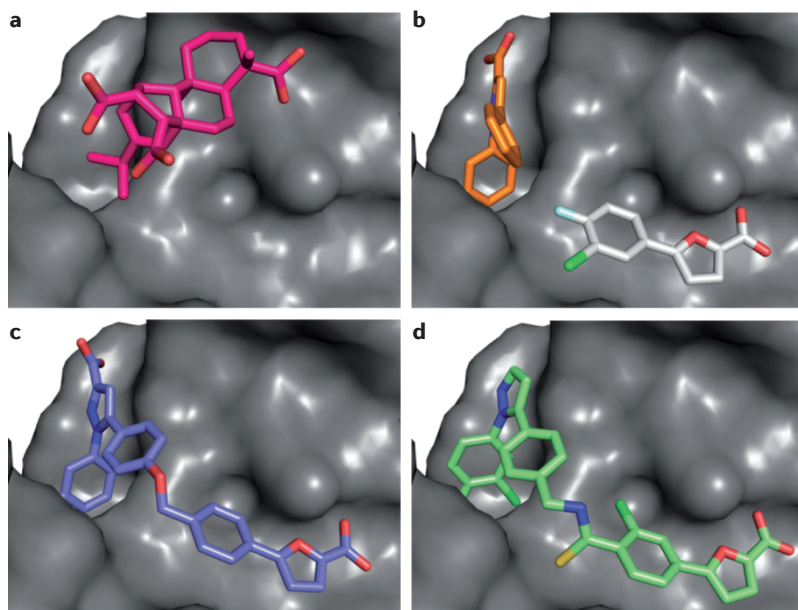


Figure 3 | Optimizing protein–ligand interactions using fragment-based screening.

An illustration of a fragment linking approach to develop inhibitors of the replication protein A (RPA; grey molecular surface) protein–protein interaction with partner proteins that are involved in the DNA-damage response, such as p53, RAD9, ATR-interacting protein (ATRIP) and MRE11 (REF. 79). **a** | A docked model of a high-throughput chemical screening (HTS) ‘hit’, fumaropimaric acid (pink), bound to RPA (fluorescence polarization (FP)-determined dissociation constant (K_D) = $18.3 \mu M$ ¹⁴²). **b** | Crystal structures of fragment hits bound to RPA (Protein Data Bank (PDB) ID: 4LUV⁷⁹; NMR-determined K_D = $580 \mu M$ (orange) and K_D = $1,400 \mu M$ (white)⁷⁹. Each fragment has fewer but more efficient and optimized interactions with the protein target compared to the HTS hit. **c** | By linking these fragments, the binding affinity was improved (the crystal structure shown is of PDB ID: 4LUZ; purple; NMR-determined K_D = $26 \mu M$; FP-determined K_D = $20 \mu M$ ⁷⁹). **d** | Further optimization gave the crystal structure of a compound (green) with an FP-determined K_D of $0.19 \mu M$ (PDB ID: 4LWC⁷⁹).

identifying a PPI modulator, using this approach for fragments can lead to a high rate of false positives owing to compound aggregation that interferes with the assay, and can be confounded by the lack of adequate solubility for fragment compounds in suitable assay buffers. The development of the BCL-2 and BCL-X_L inhibitor ABT-263 (Navitoclax)⁸⁹, which is currently in Phase II clinical trials for the treatment of various lymphomas and leukaemias, is among the first success stories to originate from fragment screening (TABLE 1). Researchers at Abbott laboratories²⁴ used a technique called 'SAR by NMR' (REF. 71) to identify two fragments that bound adjacent to one another in the BH3 α -helix-binding groove of BCL-X_L. These compounds were developed into the pre-clinical candidate ABT-737 (FIG. 2a) before being further optimized for oral bioavailability to ABT-263 (REF. 89). Efforts are now focused on introducing selectivity for individual members of the BCL-2 protein family by using a combination of directed fragment libraries⁹⁰ and peptide-mimetic scaffold libraries⁹¹. Researchers are also using NMR techniques to probe conformational changes of the BCL-X_L protein during fragment and inhibitor binding to increase specificity for particular BCL-2 protein family members⁹².

An alternative fragment-based approach for PPI modulator development is molecular tethering⁹³. Tethering uses a specific anchor point, such as a cysteine, that is adjacent to the binding site of interest, as well as a reactive group (a sulphhydryl group in the case of a cysteine anchor point) on the fragment to form a covalent link. Binding can be confirmed using MS and visualized using X-ray crystallography. Tethering was used to identify inhibitors of the cytokine signalling molecule IL-2 (REFS 94,95). Arkin and colleagues^{94,95} used tethering to screen for novel compounds with tractable chemistry and that bound to IL-2 in a manner distinct from a known inhibitor. From a modest lead compound (active between 500–600 μ M in an ELISA assay) an inhibitor with an IC₅₀ of 60 nM was developed — guided by SBDD. The ligand-free structure of IL-2 is flat and featureless (FIG. 2b), and it was only once a crystal structure of IL-2 in complex with a known inhibitor was obtained that the PPI hot spot became apparent^{93–95}.

Library design. Common to VS, HTS and fragment screening is the assembly of a library of molecules to be used as potential ligands. Compound libraries have seen considerable evolution in the past 30 years, from libraries of natural product extracts through to streamlined and characterized small-molecule libraries that contain the products of combinatorial chemistry¹⁸. There has more recently been a move to again incorporate a high degree of three-dimensional shape diversity and chirality in an attempt to capture some of the complexity that has been lost in the move away from natural products^{96,97}. The distinct nature of PPIs as targets for drug development has also begun to shape the libraries that are being used for PPI screening^{12,98}. Indeed, library design targeting PPIs that involve α -helix-binding epitopes has been described^{57,91,99}. As more screens for PPI targets are carried out, compound library design will continue

to evolve to exploit the insights that are gained from the success or failure of compound classes and from the physicochemical profile of the hit series of compounds that emerge.

Biological approaches to modulating PPIs

Biologicals are medicinal products that include protein-based drugs, peptides, aptamers and other macromolecules. The main advantages of biologicals as therapeutics are their high target specificity and potency. They also often share a metabolic pathway with endogenous macromolecules, which, along with their high specificity, results in a low level of systemic toxicity. Biologicals are particularly useful for PPI modulation, as they can be readily tuned to bind to a large variety of protein surfaces, and they are often used as a starting point for PPI drug discovery programmes.

Protein-based drugs. Examples of protein-based drugs include antibodies and other entities such as enzymes, growth factors, hormones, interferons, interleukins and engineered protein scaffolds^{100,101}. Antibodies have become the fastest growing class of biological therapeutics, particularly in oncology (see Supplementary information S1 (table)). Antibodies that bind with high affinity and specificity to their targets elicit an anti-tumour effect via immune stimulation, the focused delivery of conjugated toxins or by modulating PPIs on the surface of target tissues. The limitations of protein-based drugs include a tendency to elicit a systemic immune response before reaching the drug target (particularly with extended dosing) and an inability to access intracellular targets^{102,103}. Difficulties in modifying pharmacokinetic characteristics after production, formulation issues and excessive production or purification steps complicate the manufacture of protein-based drugs and add substantial expense^{104–107}.

Peptide-based modulators. Compared with protein-based drugs, peptides are less likely to prompt any serious immune response and are generally cheaper and quicker to produce. Indeed, more than 100 peptide-based drugs are currently on the market, with annual sales in excess of \$40 billion per year, which represents 10% of the annual sales for all available drugs¹⁰⁸ (see Supplementary information S2 (table)). In spite of this success, limitations to accessing intracellular targets and unfavourable pharmacokinetic profiles mean that peptides are often used as a starting point to develop mimetics and non-peptide drugs¹⁰⁹. Peptide modulators of PPIs have been identified through various approaches, including rational design, as well as screening random and biased peptide libraries¹¹⁰.

The rational design of peptides as PPI modulators stemmed from the observation that many PPIs involve a continuous epitope of one partner and a well-defined groove or series of specific small pockets on the target protein surface (BOX 1; FIG. 4). Perhaps the most well-known example of this is the development of the stapled BIM β 5 peptides (FIG. 2a) as rationally designed inhibitors of BCL-X_L (REF. 111). These peptides have since been

Libraries of natural product extracts

Compound libraries that are prepared from extracts of natural products such as plants, animals or parts thereof. Compounds in a natural product library may be present as crude extracts, fractionations or purified products.

Stapled BIM β 5 peptides

Stapled BIM β 5 peptides have been chemically locked into a stable conformation by the introduction of an intramolecular linker between the residues. This modification results in a peptide that is resistant to proteases, has increased cell penetration, improved pharmacokinetics and, usually, a high affinity to the target protein surface.

used to develop peptide mimetics and have aided in the development of small-molecule BCL-2-selective inhibitors^{91,112}. The pro-apoptotic BH3-only protein NOXA provides a more recent example of a case in which rational design has been successfully used to increase the potency, selectivity, proteolytic stability and cell permeability of a peptide¹¹³. The α -helix of NOXA binds to the BCL-2 family member MCL1 with high affinity and selectivity. MCL1 has been implicated in multiple cancers and has specific clinical implications for chronic myelogenous leukaemia and multiple myeloma. Guided by X-ray crystallography, a cysteine-mediated crosslink was used to stabilize the NOXA α -helical peptide. However, the stabilized peptide was still not cell-permeable, and further structure-based refinement of the peptide was carried out. By replacing solvent-exposed residues and methylating the peptide backbone,

a peptide mimetic with a 65-fold increase in inhibitory activity and a >200-fold selectivity for MCL1 over other BCL-2 protein family members was obtained. Examples such as this illustrate the subtlety that is involved in the rational design of peptides into potential therapeutics that target intracellular PPIs.

Peptide modulators have also been discovered through screening techniques, including alanine scanning mutagenesis and phage screening^{109,110}. Phage screening identifies linear peptide epitopes that are expressed on the surface of filamentous bacteriophage viruses. Using a random bacteriophage library with the target of choice, phages that have weak peptide–protein target interactions are selected and used to infect colonies of bacteria. During the infection, random substitution occurs in the surface epitopes, and then additional cycles of more stringent selection–replication of the phage progeny are carried out. The end result is a set of phage-carrying peptide sequences that have been tuned by evolution to bind strongly to the target protein¹¹⁴. The identified sequence (or sequences) can be directly used as a peptide ligand or further developed by subsequent modification and adaptation. For example, as mentioned above, phage screening was used to create the pharmacophore model behind the current Phase I MDM2–p53 inhibitors²⁸. These inhibitors are being sought as tumour-selective drugs for multiple cancer types¹¹⁵. A screen of phage display peptide libraries identified a sequence with a 28-fold greater inhibition of the MDM2–p53 interaction than the wild-type p53-derived peptide. A crystal structure of the N-terminal domain of MDM2 bound to a wild-type p53-derived peptide showed important structural motifs, such as a type 1 β -turn, that were contained within the peptide. The phage-derived peptide was modified through cysteine-mediated crosslinking chemistry and *N*-methylation of solvent-exposed residues to adopt these motifs, which resulted in a peptide with a 63-fold greater potency (IC_{50} of 5 nM) than the wild-type-derived peptide¹¹⁶. These peptides did not reach the clinic owing to poor membrane permeability and physiological stability, but they contributed to the design of non-peptidic inhibitors that are in early phase clinical trials.

Aptamers as PPI modulators. Aptamers are synthetic oligonucleotides that are usually identified through multiple rounds of selection and amplification in a similar way to phage display libraries. Aptamers can be developed to bind to their protein targets with extremely high affinity and have the capacity to modulate PPIs in a similar manner to peptides and antibodies¹¹⁷. Like antibodies, aptamers are sought not only as therapeutics but also as drug conjugates for targeted drug delivery and biomarker identification¹¹⁸. The key advantages that aptamers have over antibodies are that they are relatively small, comparatively inexpensive, free from cell-culture-derived contaminants and essentially non-immunogenic. Aptamers can also be developed against almost any protein target, including toxins, molecules that do not elicit an immune response

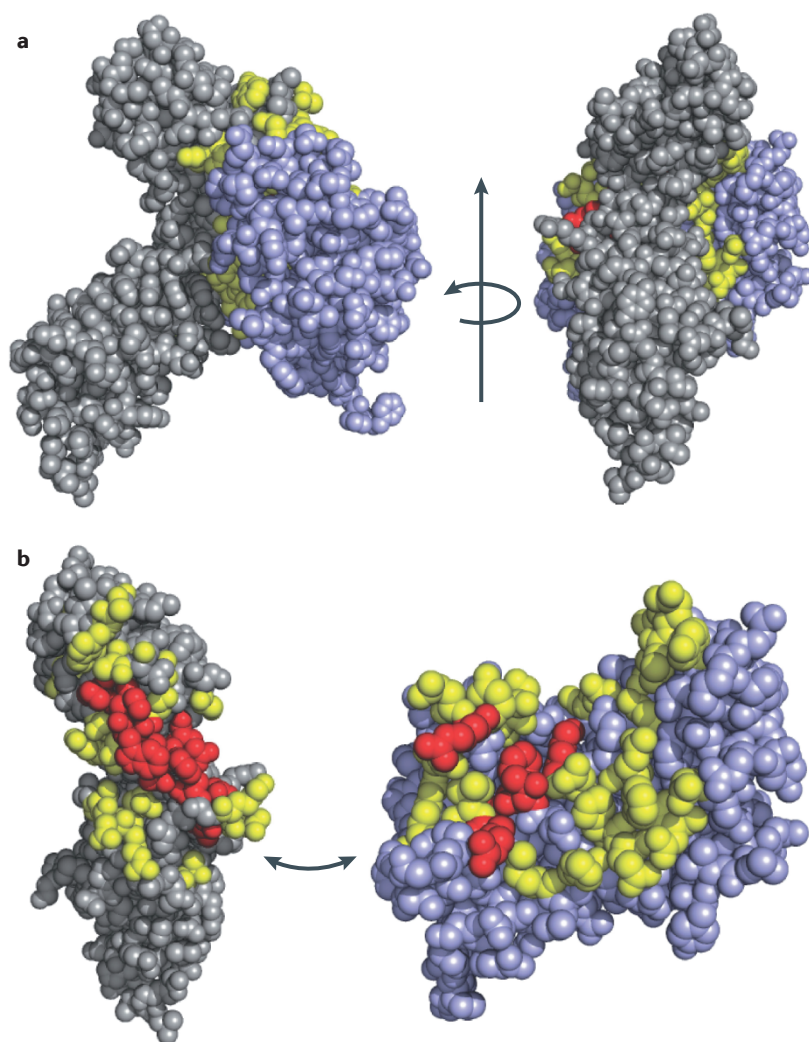


Figure 4 | Hot spots and O-rings at the protein–protein interface. a | Orthogonal views of the human growth hormone binding protein (GHBP)–growth hormone (GH) complex (Protein Data Bank (PDB) ID: 3HHR¹³³), with the proteins depicted as atom spheres coloured either purple (GH) or grey (GHBP) except for the ‘hot spot’ (red) and ‘rim’ (yellow) residues. Hot spots are usually ringed by a layer of solvent-excluding residues called the ‘O-ring’, and around the O-ring are the rim residues. **b** | The complex is shown opening to expose the interacting surface.

Allosteric binding sites

Sites on a protein surface that influence the biological function of a protein upon binding of a regulatory substance but that are not considered to be the protein–protein interaction site or the enzymatic or receptor active site.

and small epitopes for which antibodies cannot be raised. Several aptamers for use in cancer patients have been successfully taken to the market or are currently undergoing clinical trials (see Supplementary information S3 (table)). For example, ERBB2-specific anticancer aptamers have been recently developed^{119–123} to deliver small interfering RNAs that target BCL-2 to breast cancers¹²⁴, and aptamers that target platelet-derived growth factor (PDGF) have been developed to be used in conjunction with traditional kinase inhibitors, such as imatinib mesylate¹²⁵, potentially for the treatment of patients with gastrointestinal tumours¹²⁶. These studies highlight the versatile selection process, low production cost and low batch-to-batch variability that make aptamers an attractive enterprise for clinical application.

Conclusions and future directions

Approaches to identify and target PPIs for therapeutic development have gathered pace over the past few years. This development is likely to continue in light of pharmaceutical industry analyst predictions that worldwide sales of small-molecule PPI modulators are set to exceed

\$800 million per annum by 2018 (REF. 5). There is cause for much optimism, particularly for cancer in which progress has been perhaps the most dramatic. Although only a few small-molecule drugs that modulate PPIs have reached human clinical trials to date, there has been exciting progress in developing the appropriate methodology — for example, in fragment screening, compound library design and in peptide mimetics — that should in future lead to the development of better cancer-targeted drugs. Furthermore, evidence is emerging that PPIs can be indirectly disrupted through the binding of small molecules at allosteric binding sites¹²⁷, some of which engage ‘cryptic’ pockets, which are not present in crystal structures of the uncomplexed protein target¹²⁸. The fragment screening approach is proving a powerful means to discover these conformationally adaptable pockets. As a class, PPI modulators tend to be larger than typical orally available drugs⁸. A challenge for the future will be finding ways to optimize the pharmacokinetic properties, such as absorption, distribution, metabolism and excretion, of PPI modulators early in the discovery process to ensure that the compounds are suitable for the clinic.

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Competing interests statement

The authors declare competing interests: see Web version for details.

DATABASES

The Cancer Genome Atlas: <http://cancergenome.nih.gov/>
International Cancer Genome Consortium: <http://icgc.org/>
ClinicalTrials.gov: <http://ClinicalTrials.gov>
National Cancer Institute DIS 3D database: http://dtp.nci.nih.gov/docs/3d_database/dis3d.html
Protein Data Bank: <http://www.rcsb.org/pdb/home/home.do>

FURTHER INFORMATION

2P2ldb: <http://2p2ldb.cnrs-mrs.fr/>
Accelrys: www.accelrys.com
ANCHOR: <http://structure.pitt.edu/anchor/>
Click2Drug: <http://www.click2drug.org/>
Developmental Therapeutics Program, NCI/NIH: <http://dtp.nci.nih.gov/index.html>
Dictionary of Marine Products: <http://dmnp.chemnetbase.com>
Dictionary of Natural Products: <http://dnp.chemnetbase.com>
DrugBank: <http://www.drugbank.ca/>
FAF-Drugs2 server: <http://bioservrpbis.univ-paris-diderot.fr/FAF-Drugs/>
FDA Drug-related Databases: <http://www.fda.gov/Drugs/default.htm>
Human Cancer Protein Interaction Network (HCPIN): <http://www.nesg.org/9090/HCPIN/index.jsp>
iPPI-DB: <http://www.ippidb.cdithem.fr/>
National Cancer Institute (NIH, USA): <http://cancer.gov/>
NCGC Pharmaceutical Collection (NPC): <http://tripod.nih.gov/npc/>
PPI Network: <http://ppi-net.org/>
PrePPI: <http://bhapp.c2b2.columbia.edu/PrePPI/>
Protein-Protein Interactions: <http://bip.weizmann.ac.il/toolbox/structure/protein-protein.htm#db2>
Schrödinger: <http://www.schrodinger.com>
The MIPS Mammalian Protein-Protein Interaction Database: <http://mips.helmholtz-muenchen.de/proj/ppi/>
The Protein Interaction Network Analysis (PINA): <http://cbg.garvan.unsw.edu.au/pina/>
TIMBAL: <http://mordred.bioc.cam.ac.uk/timbal/>
Virtual Ligand Screening in 3D.com: <http://www.vls3d.com/links.html>
Wiki-Pi: <http://severus.dbmi.pitt.edu/wiki/pi/>

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