

REVIEW

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Protein–protein interfaces in molecular glue-induced ternary complexes: classification, characterization, and prediction

Huan Rui,^a Kate S. Ashton,^b Jaeki Min,^c Connie Wang^d and Patrick Ryan Potts^{*,c}

Molecular glues are a class of small molecules that stabilize the interactions between proteins. Naturally occurring molecular glues are present in many areas of biology where they serve as central regulators of signaling pathways. Importantly, several clinical compounds act as molecular glue degraders that stabilize interactions between E3 ubiquitin ligases and target proteins, leading to their degradation. Molecular glues hold promise as a new generation of therapeutic agents, including those molecular glue degraders that can redirect the protein degradation machinery in a precise way. However, rational discovery of molecular glues is difficult in part due to the lack of understanding of the protein–protein interactions they stabilize. In this review, we summarize the structures of known molecular glue-induced ternary complexes and the interface properties. Detailed analysis shows different mechanisms of ternary structure formation. Additionally, we also review computational approaches for predicting protein–protein interfaces and highlight the promises and challenges. This information will ultimately help inform future approaches for rational molecular glue discovery.

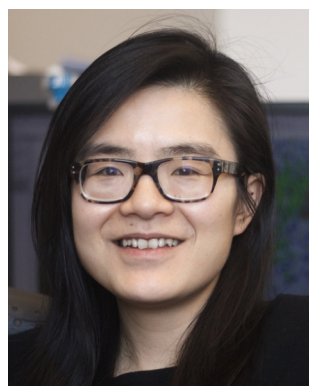
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Introduction

For the past decade, molecular glue degrader discovery has relied on serendipity, *post hoc* rationalization, and phenotypic screening. The most well-known molecular glue degraders are thalidomide and its analogues, lenalidomide and pomalidomide known collectively as immunomodulatory imide drugs (IMiDs). Their discovery as molecular glues was retrospective following FDA approval and subsequent detection of

^a Center for Research Acceleration by Digital Innovation, Amgen Research, Thousand Oaks, CA 91320, USA^b Medicinal Chemistry, Amgen Research, Thousand Oaks, CA 91320, USA^c Induced Proximity Platform, Amgen Research, Thousand Oaks, CA 91320, USA.
E-mail: ryan.potts@amgen.com^d Digital, Technology & Innovation, Amgen, Thousand Oaks, CA 91320, USA

Huan Rui

Huan Rui is a computational chemist at Amgen. She obtained her PhD in computational chemistry from University of Kansas and conducted her postdoctoral research at University of Chicago studying conformational transitions of membrane proteins using molecular dynamics and free energy calculations. Her current research interest is on combining physics and machine learning based methods to predict molecular glue induced protein–protein interfaces.



Kate S. Ashton

Kate Ashton graduated from the University of Cambridge in 2005 with a PhD in organic chemistry before moving to conduct post-doctoral studies at Caltech and Princeton. Since starting at Amgen, she has worked on multiple programs spanning all therapeutic areas and was part of the sotorasib team. She is focused on induced proximity and the challenges it brings to small molecule chemistry.

immunomodulatory and anti-inflammatory activity. The E3 ubiquitin ligase cereblon (CRBN) was identified as the effector protein of thalidomide.¹ Binding of thalidomide to CRBN recruits and degrades neosubstrates, providing an early example of molecular glue-induced targeted protein degradation (TPD).^{1–7} IMiDs are not the only known molecular glue degraders whose mechanisms of action was uncovered retrospectively. Indisulam and its derivatives were also first discovered through a phenotypic screen for small molecules with anti-cancer activity.⁸ Their mechanism of action was uncovered almost 20 years later using mutagenesis and mass spectrometry, identifying RBM39 as their degradation target and DCAF15 as the effector E3 ligase.^{9–11} Even though molecular glue molecules are often discussed in the context of E3 ligases and TPD, they exist in other areas of biology as well. For instance, rapamycin inhibits the mTOR kinase by stabilizing

a protein–protein interface between FKBP12 and the FRB domain in the kinase.¹² Another example is the nuclear receptors. Their specificity is largely modulated by the binding of different ligands, leading to preferential recruitment of either co-repressors or co-activators.^{13,14} There are many other examples of this class of molecules, including FK506,¹⁵ inositol tetraphosphate,¹⁶ and cotylenin A.¹⁷ The discovery of these molecules, which act as protein–protein interaction stabilizers, faces similar challenges as the rest of the molecular glues.

Recently, more systematic screening approaches have been used to identify molecular glue degraders. CR8, a CDK12/CyclinK molecular glue degrader, was identified by a bioinformatics screen, where the cytotoxicity of clinical and preclinical small molecules in different cancer cell lines are correlated with their E3 ligase components mRNA levels.¹⁸ Extensive follow-up testing and a crystal structure revealed that CR8 induces the formation of a complex between CDK12/CyclinK and the E3 ligase DDB1 leading to the degradation of CDK12/CyclinK. Along the same vein, other screening approaches have shown success at discovering novel CyclinK molecular glue degraders as well.^{19,20} These works also highlight the tremendous effort required to perform current screening paradigms, not only due to the nature of the screen itself – typically a cellular assay – but also the follow-up work needed to validate and understand the mechanism of action for any hits. Furthermore, since the hit rates of such screening efforts are usually low, testing a large number of compounds is necessary. Although new technologies such as cellular screens using DNA encoded library technology^{21–24} offer the opportunity to screen millions of compounds at once, hits still need to be remade off DNA and tested again in assays to confirm activity, making the lead generation process slow and resource intensive. Because of this, screening for molecular glues currently falls squarely within the high risk, high reward zone of drug discovery.



Jaeki Min

Chemical Biology Center in the department of chemical biology and therapeutics. Currently he is a group leader of Induced Proximity Platform at Amgen, and his primary research area is targeted protein degradation.

Jaeki Min received his PhD in organic chemistry from Seoul National University in 2004 and then went on to undertake post-doctoral research in chemical genomics at New York University. Afterwards, he joined Emory University's Emory Chemical Biology Discovery to work on the NIH Molecular Library Screening Center Network project. He then moved to St. Jude Children's Research Hospital, where he became the center leader of the



Connie Wang

Connie Wang is a data scientist at Amgen. She obtained her PhD in computational chemistry from Caltech where her research focused on coarse-grained modeling of membrane protein synthesis. Her current work has focused on applying machine learning and data science methods to discover novel therapies and accelerate their delivery to market.



Patrick Ryan Potts

of Cell and Molecular Biology. In 2020 he moved to Amgen, Inc. as Executive Director of Research and Head of the Induced Proximity Platform that is focused on drugging the “undruggable”.

Ryan Potts obtained his PhD in Cell and Molecular Biology from UT Southwestern in 2007. In 2008 he was awarded the Sara and Frank McKnight junior faculty position at UT Southwestern Medical Center and appointed as Assistant Professor in the Departments of Physiology, Pharmacology, and Biochemistry in 2011. In 2016 his lab moved to St. Jude Children's Research Hospital where he was an Associate Member in the Department



Despite these challenges, molecular glue degraders are still highly sought after as a promising new therapeutic modality. Compared to its bi-specific counterpart proteolysis targeting chimeras (PROTACs), molecular glue degraders have the advantage of being small with low molecular weight and the potential for drug-like physiochemical properties, making the late-stage optimization relatively simpler. Many research labs and pharmaceutical companies have started programs aimed at discovering new molecular glues for protein degradation, but without a more efficient way to discover novel degraders and protein targets, the field will face an uphill battle.

Structure-based rational design methods offer a strategy to increase the chance of success if the target and the E3 ubiquitin ligase of the molecular glue are known. In a recent example, the crystal structure of ternary complex between β -TrCP, β -catenin, and a small molecule NRX-1933 led to the discovery of new molecular glues with enhanced mutant selectivity.²⁵ The discovery of the mutant-specific Ikaros degrader, ALV2, was also guided by known crystal structures.^{26,27} However, structural information on the ternary complex is often difficult to obtain and does not always aid in the prediction and discovery of neosubstrates. The crux of this issue lies within the interactions formed by the two proteins and their stabilization by the ligand. Many molecular glues take advantage of weak, fortuitous, pre-existing protein-protein interfaces (PPIs) that can be further strengthened by their binding. To fully understand what drives ternary structure formation and leverage this in the discovery of new molecular glue molecules, a few questions still need to be addressed, including (1) what is the major driving force behind the protein-protein interactions in the ternary complexes, (2) what is the influence of the ligand on the PPI, (3) what are the stability, kinetics, and geometry of the ternary complexes formed, and (4) which of these parameters are most crucial.

A plethora of excellent review articles on molecular glues and the complexes they stabilize have been published in the recent years, each focusing on a slightly different aspect of the topic, with the majority of them focusing on the discovery history, mechanism of action, and design strategy.^{28–37} This review seeks to compile and analyze the current, published molecular glue ternary complex structures with the aim of identifying and quantifying any structural requirements for PPIs that are induced by molecular glues, as well as surveying the best computational methods to predict potential ‘gluable’ interfaces. To gain a general understanding of the molecular glue-induced PPI (MG-PPI), all molecular glue-induced complexes are considered, regardless of whether the molecular glue is a degrader. The review is organized into four sections: (1) a summary of all known MG-PPI and two set of native PPI that do not require molecular glues to form, (2) a comparison of the PPI properties between the sets of interfaces highlighting similarity and differences, (3) an analysis of the ligand-protein interactions occurring in the MG-PPI, showing different mechanisms of MG-PPI formation, and (4) a survey of current computational tools for PPI discovery and their potential utility in the identification of gluable PPI.

Summary of protein-protein interfaces

Molecular glue-induced protein-protein interfaces

To establish a baseline, an analysis of currently documented MG-PPI was performed with an emphasis on protein-protein and protein-ligand interfaces, taking into account both their size and nature. Table 1 provides a summary of publicly available molecular glue-induced heterodimeric complex structures from the protein data bank (PDB). The corresponding small molecule structures are shown in Scheme 1. The structures in Table 1 are chosen based on our best knowledge of literature discussing molecular glues and stabilizers. All types of molecular glues are included, including degraders and non-degraders. There are homo-oligomer complex structures available but often times they involve multiple protein-protein interfaces and are excluded from the current review. One exception is BCL6 due to the recent discovery of its degrader BI-3802,³⁸ which causes homo-oligomerization of the BTB domain,³⁹ and the rising interest in its degrader design. Bispecific degrader induced ternary structures are also not included in Table 1 as the ternary structure formation in this case is largely driven by the binding of the warheads to their respective protein targets. The complex structures in Table 1 are organized into two groups based on the interacting mode between the two protein binding partners: Group 1 (domain-domain) features interactions between proteins with well-folded domains, and in Group 2 (sequence motif-domain) one of the binding partners is a stretch of residues that contains a specific pattern for binding (Fig. 1A and B). In all the complexes in Table 1, 45 of them belong to Group 1 and 54 belong to Group 2. Fig. 1B shows two complex structures, DCAF15-indisulam-RBM39 and TIR1-1NAA-IAA7, exemplifying the two groups of interactions respectively. There is a myriad of ways to categorize these ternary complexes, the reason for the current choice is rooted in the nature of these interfaces and the computational discovery strategies that may follow.

Many of the complex structures in Table 1 share the same effector proteins and are clustered together in the table. Fig. 1C shows the breakdown of all the structures into their respective effector proteins. There are a total of 21 effector proteins, 13 of which are from Group 1, the rest belong to Group 2 with a sequence motif from the binding partner protein interacting with the effector protein. A representative structure from each effector protein cluster is used in the following quantitative analyses to show structural characteristics of the MG-PPI. This includes 15 complexes from Group 1 and 14 complexes from Group 2. They are highlighted in bold in Table 1. The selection is based on structure completeness and resolution.

Protein-protein interfaces that are not induced by molecular glues

Two protein-protein docking datasets, Docking Benchmark 5.5 (DB5.5)^{41–45} and Database of Interacting Protein Structures (DIPS),⁴⁶ are used here to represent interfaces that form without the help of molecular glues. DB5.5^{41–45} is a manually curated dataset containing 253 protein-protein complexes



Table 1 Molecular glue-induced ternary complexes from the PDB. Representative structures used for analysis are shown in bold

Binding affinity									
P1	P2	PDB	Glue	Resolution	P1-glue	P1-P2	Ternary	Assay type	
Group 1	RCAR12 (PYL1)	ABI1	3JRQ	Absciscic acid	2.1 Å	K _D = 52 μM (ITC) K _D = 340 μM (SPR)		ITC, SPR ¹⁵⁶	
	RCAR12 (PYL1)	ABI1	3KB3	Absciscic acid	2.0 Å	K _D = 52 μM (ITC) K _D = 340 μM (SPR)			
	RCAR12 (PYL1)	ABI1	3KDJ	Absciscic acid	1.9 Å	K _D = 52 μM (ITC) K _D = 340 μM (SPR)			
	CSPYL1	HAB1	5MN0	Absciscic acid	2.0 Å	N/A			
	RCAR11 (PYR1)	HAB1	3QN1	Absciscic acid	1.8 Å	K _D = 97 μM		NMR ¹⁵⁷	
	RCAR11	HAB1	3ZVU	Absciscic acid	2.1 Å	K _D = 3 μM			
	(PYR1, H60P)							ITC ¹⁵⁸	
	2RCAR14 (PYL2)	ABL2	3UJL	Absciscic acid	2.5 Å	K _D = 59.1 μM			
	RCAR9 (PYL2)	ABL1	4OIC	Absciscic acid	2.0 Å			ITC ¹⁵⁹	
	RCAR13 (PYL3)	HAB1	4DS8	Absciscic acid	2.2 Å				
	RCAR13 (PYL3)	HAB1	5JO2	Absciscic acid	2.4 Å				
	RCAR3	ABL3	5GWO	Absciscic acid	2.8 Å		K _D = 30 nM (at 0.1 mM of ABA) K _D = 2.1 nM (at 0.1 mM of ABA) N/A		
	RCAR3	(S265F/I267M)	ABL3	5GWP	Absciscic acid	2.6 Å			Biolayer interferometry (BLI) ¹⁶⁰
	RCAR3	ABL3	5ZCG	Absciscic acid	2.1 Å				
	RCAR3	(S265L/I267V)	ABL3 (I267W)	5ZCH	Absciscic acid	2.5 Å			
	RCAR3	ABL3 (I267L)	ABL3	5ZCL	Absciscic acid	2.7 Å			
	FKFBP12	FRAP	1FAP	Rapamycin	2.7 Å	K _D = 0.2 nM (P1-Glue) K _D = 26 μM (P2-Glue)	ND		FP assays and SPR ⁵³
	FKFBP12	FRAP	1NSG	Rapamycin	2.2 Å		ND		
	FKFBP12	FRAP (T2098L)	6M4U	Rapamycin	2.2 Å		ND		
	FKFBP12	FRAP	2FAP	C49-methyl Rapamycin	2.2 Å	K _i = 3.5 nM	ND		
FKFBP12	FRAP	3FAP	C15-(R)-methylthienyl Rapamycin	1.9 Å		ND		FKBP12 enzyme inhibition assay ¹⁶¹	
FKFBP12	FRAP	4FAP	C15-(R)-methylthienyl Rapamycin	2.8 Å		ND			
FKFBP12	PPP3CA-PPP3R1	1TCO	FK506	2.5 Å	K _D = 0.4 nM	ND			
FKFBP12	CEP250	6OQA	WDB002	2.2 Å		ND			
DCAF15	RBM39	6QOW	Indisulam	2.9 Å	K _D = 17 μM K _i > 50 μM	K _D = 4.6 μM	K _D = 120 nM	BLI, ⁵⁶ ITC, ¹⁶³ TR-FRET ¹⁶⁴	
DCAF15	RBM39	6UD7	Indisulam	2.3 Å		K _D = 4.6 μM K _D = 4.6 μM			
DCAF15	RBM39	6SJ7	Indisulam	3.5 Å (Cryo-EM)		K _D = 4.6 μM K _D = 4.6 μM			
DCAF15	RBM39	6QOV	Tasisulam	2.9 Å	K _i = 2.9 μM	K _D = 4.6 μM			
DCAF15	RBM39	6PAI	E7820	2.9 Å	K _i = 2.9 μM	K _D = 4.6 μM		K _D = 160 nM	
DCAF15	RBM39	6QOR	E7820	2.9 Å		K _D = 4.6 μM			
DCAF15	RBM39	6UE5	Compound 7	2.6 Å		K _D = 4.6 μM			
ARF1	ARNO	1R8Q	Brefeldin A	1.9 Å	ND	K _D = 4.6 μM			
ARF1	ARNO	1S9D	Brefeldin A	1.8 Å	ND			Fluorescence anisotropy ^{165,166}	
ARF1	GEA	1RE0	Brefeldin A	2.4 Å	ND		K _i = 15 μM K _i = 15 μM EC ₅₀ = 1.61 mM		
Cam	Kcnn2	4G27	Phenylurea	1.7 Å	ND				
									Electrophysiology ¹⁶⁷

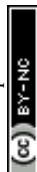
Binding affinity

Binding competition assay¹⁶⁸
 Binding competition assay¹⁶⁸
NMR¹⁶⁹
TR-FRET^{169,170}

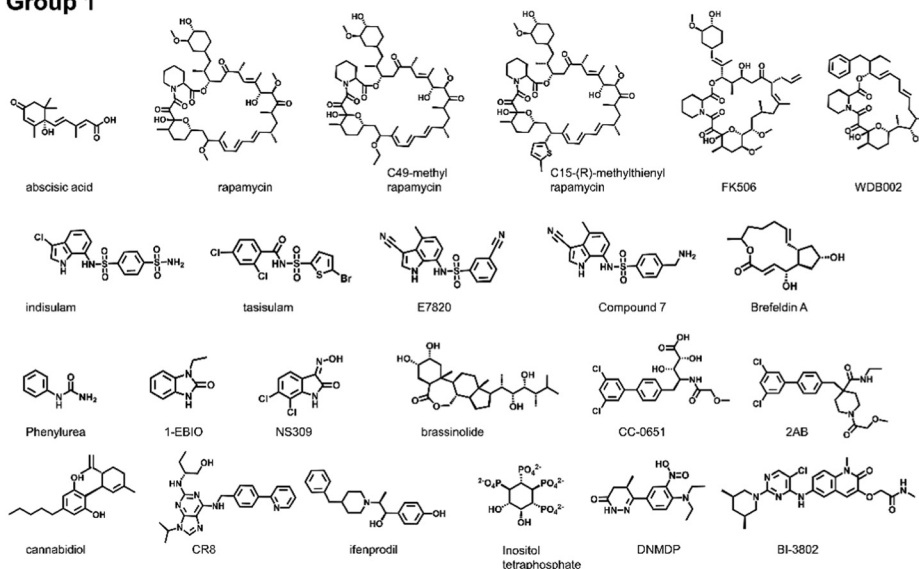
 TR-FRET¹⁷⁰
ITC⁵⁵
BLI⁵⁵
ITC¹⁸
ITC⁵⁴
Analytical ultracentrifugation⁵⁴

Table 1 (continued)

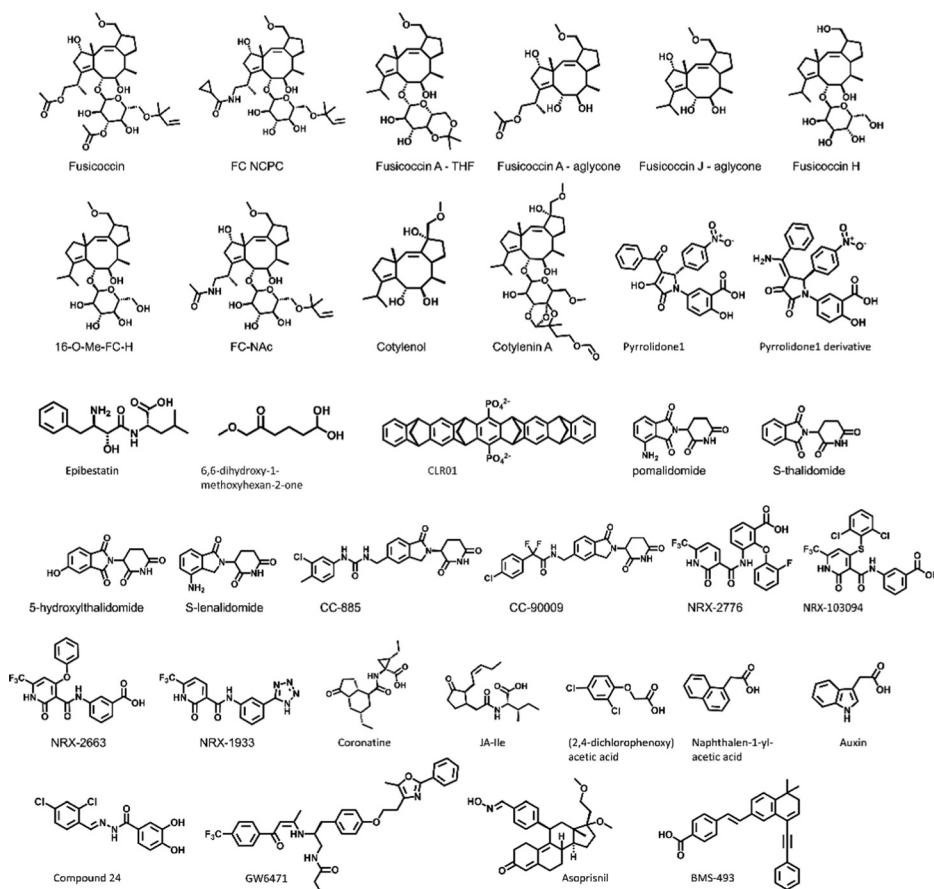
P1	P2	PDB	Glue	Binding affinity			Assay type
				Resolution	P1-glue	Ternary	
14-3-3	TASK-3	3UX0	Fusicoccin H	1.8 Å	$K_D = 4.1 \mu\text{M}$ (ITC)	$K_{D,\text{app}} = 0.27 \mu\text{M}$ (ITC)	FP ¹⁸⁵
14-3-3	TASK-3	4FR3	16-O-Me-FC-H	1.9 Å	$K_D = 1.46 \mu\text{M}$	$K_{D,\text{app}} = 0.55 \mu\text{M}$	FP ¹⁸⁵
14-3-3	TASK-3	6GHP	FC-Nac	2.0 Å	$K_D = 0.45 \mu\text{M}$	$K_{D,\text{app}} = 0.07 \mu\text{M}$	FP ¹⁸⁶
CRBN	IKZF1(ZF2)	6H0F	Pomalidomide	3.3 Å	$K_D = 157 \text{ nM}$	$K_i = 2314 \text{ nM}$ (CRBN-ZF2)	FP assay ⁶
					$K_D = 250 \text{ nM}$ (CRBN-ZF2-3)	(CRBN-ZF2)	TR-FRET assay ¹⁸⁷
CRBN	ZNF692(ZF4)	6H0G	Pomalidomide	4.3 Å	$K_D = 157 \text{ nM}$	$K_D = 53 \text{ nM}$ (CRBN-ZF2-3)	BLI ¹⁸⁵
CRBN	SALL4	6UML	Pomalidomide	3.6 Å	$K_D = 157 \text{ nM}$	$K_i = 2314 \text{ nM}$	Alphascreen ^{5,5}
CRBN	SALL4	7BQU	S-Thalidomide	1.9 Å	$K_D = 4 \mu\text{M}$		TR-FRET ¹⁸⁷
CRBN	SALL4	7BQV	5-Hydroxythalidomide	1.8 Å	$K_D = 0.76 \mu\text{M}$		ITC ¹⁸⁸
CRBN	GSP1	5HXB	CC-885	3.6 Å	$K_D = 71 \text{ nM}$	$K_D = 1.8 \text{ nM}$	ITC ¹⁸⁸
CRBN	GSP1	6XK9	CC-90009	3.6 Å	$\text{IC}_{50} = 18 \text{ nM}$		MST ¹⁸⁹
CRBN	CK1α	5FQD	S-Lenalidomide	2.5 Å	$\text{IC}_{50} = 824 \text{ nM}$		FP assay ¹⁹⁰
					$K_D = 180 \text{ nM}$		FP assay ¹⁹⁰
β-TRCP	β-catenin (pS33/S37)	6M90	NRX-2776	2.1 Å	$K_D = 689 \text{ nM}$	$K_D = 75 \text{ nM}$	BLI ¹⁹¹
β-TRCP	β-catenin (pS33/S37A)	6M91	NRX-103094	2.4 Å	$K_D = 272 \text{ nM}$		FP assay ⁶
β-TRCP	β-catenin (pS33/S37)	6M92	NRX-2663	2.4 Å	$K_D = 689 \text{ nM}$	$K_D = 0.6 \text{ nM}$ (at 40 μM NRX-103094)	BLI ¹⁹¹
β-TRCP	β-catenin (pS33/S37)	6M93	NRX-1933	2.5 Å	$K_D = 689 \text{ nM}$	$K_D = 54.8 \text{ nM}$ (at 250 μM NRX-2663)	TR-FRET ²⁵
COI1	JAZ1	3OGK	Coronatine	2.8 Å			TR-FRET ²⁵
COI1	JAZ1	3OGM	Coronatine	3.3 Å		$K_D = 68 \text{ nM}$	Radio-ligand binding assay ⁴⁷
COI1	JAZ1	3OGL	JAI-isoleucine (JAI-Ile)	3.2 Å		$K_D = 68 \text{ nM}$	Radio-ligand binding assay ⁴⁷
TIR1	IAA7	2PIN	(2,4-Dichlorophenoxy) acetic acid	2.5 Å	$K_D = 1.4 \mu\text{M}$	$K_i = 1.8 \mu\text{M}$	Competitive binding ⁴⁷
TIR1	IAA7	2PIO	Naphthalen-1-yl-acetic acid	1.9 Å	$K_D = 1.3 \mu\text{M}$		Competitive binding ¹⁹¹
TIR1	IAA7	2PIQ	Auxin	1.9 Å	$\text{IC}_{50} = 84 \text{ nM}$	Exists but not quantified	Competitive binding ¹⁹¹
Aldolase	TRAP	4TR9	Compound 24	2.1 Å			BLI ¹⁹¹
PPARα	SMRT	1KKQ	GW6471	3.0 Å	$\text{IC}_{50} = 8 \mu\text{M}$	$K_D = 1.6 \mu\text{M}$	Competitive binding, SPR, NMR ^{191,192}
PR	NCoR	2OVM	Asoprisnil	2.6 Å	$K_D = 0.5 \mu\text{M}$		Fluorescence energy transfer assays and FP ¹⁹³
RAR	N-CoNR1	3KMZ	BMS-493	2.1 Å	$K_D = 1.8 \mu\text{M}$	$K_{D,\text{app}} = 2.4 \mu\text{M}$ $K_D = 0.2 \mu\text{M}$	Fluorescence anisotropy ¹⁴



Group 1



Group 2



Scheme 1 All molecular glues in Table 1.

spanning eight categories based on protein functions. The binding affinities of these interfaces span a range between picomolar as seen in some of the antibody-antigen complexes to single digit micromolar. It is commonly used to benchmark

protein-protein docking algorithms. DIPS is a much larger nonredundant protein-protein complex dataset assembled by Townshend and coworkers.⁴⁶ It is constructed by filtering all protein-protein interfaces in the PDB based on structure



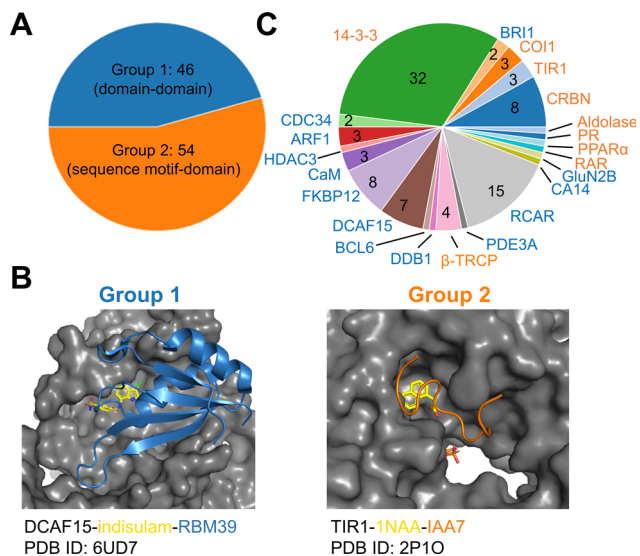


Fig. 1 Summary of all molecular glue-induced ternary complexes. (A) Breakdown of complexes in Table 1 into two groups based on the interaction mode. (B) Two example structures showing Group 1 (left, DCAF15-indisulam-RBM39, PDB ID: 6UD7) and Group 2 (right, TIR1-1NAA-IAA7, PDB ID: 2P1O) interaction modes. The effector proteins are shown in surface presentation and the binding partner proteins are shown in cartoon representation. The bound molecular glue molecules are shown in yellow. The graphics are generated using PyMOL.⁴⁰ (C) Breakdown of complexes by effector proteins. The names of the effector proteins are shown in the outer perimeter of the pie chart with blue (Group 1) and orange (Group 2) indicating the mode of interactions.

resolution, PPI size, and sequence homology to the DB5 complexes. It contains more than 42 000 interfaces. The DIPS dataset was further trimmed down by removing complexes in Table 1 to exclude MG-PPI. Interface properties are computed for complexes from both datasets and compared with those extracted from representative MG-PPI in Table 1.

Comparison between different types of ppis

Protein–protein buried surface area

The average buried surface area (BSA) of the PPI of the representative complexes in Group 1 and 2, as well as in the DB5.5 and DIPS datasets, are shown in Fig. 2A. The average BSA of Group 1 MG-PPI and the DB5.5 dataset are similar, but smaller than that seen in the DIPS dataset (Fig. 2A). Complexes in Group 2 in general have smaller BSA than those in all three other groups (*i.e.*, Group 1, DB5.5, and DIPS), which is unsurprising due to the significantly shorter amino acid sequence of one protein partner in complexes from this dataset. The PPI in the other three datasets (Group1, DB5.5, and DIPS) are largely biased towards complexes formed between folded protein partners. Intrinsically disordered proteins are not included when building DB5.5 dataset.^{41–45} The DIPS dataset contains complexes with large PPIs such as the membrane protein complex phosphatidylcholine flippase Dnf2-Lem3 (PDB ID: 7KY8) with a PPI BSA at 4995 Å² and the Myosin II complete

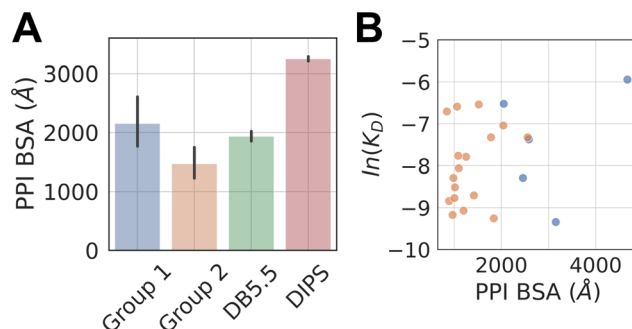


Fig. 2 PPI BSA of complexes in different datasets and their relationship with experimentally measured intrinsic protein–protein interaction K_D . (A) Average PPI BSA in four datasets. The PPI BSA for each complex is computed by taking the difference between two solvent accessible surface areas (SASA). One of them is the sum of the SASA of the two proteins as monomers. The other is the SASA of the two proteins as a complex. The difference is the reported PPI BSA. Molecular glues are excluded when computing the PPI BSA for Group 1 and 2 complexes. Standard deviation is reported for Group1, 2, and DB5.5. A two-tailed student *t*-test is performed on the BSA from Group 1 and Group 2. A *p*-value of 0.019 is achieved, indicating the observed difference is statistically significant. DIPS dataset is randomly sampled 20 times and each time with 500 samples to produce a standard error, which is reported. All properties reported below for DIPS are sampled using this scheme. (B) Shows the scatter plots of natural log of K_D against the PPI BSA for Group 1 (blue) and 2 (orange) complexes with known protein–protein binding affinities (K_D).

coiled-coil domain (PDB ID: 7KOG) with an extensive PPI BSA at 25 746 Å². Furthermore, the DIPS dataset contains a filter to remove complexes with buried surface area smaller than 500 Å².⁴⁶ This shifts the PPI BSA distribution to the higher end and as a result the average PPI BSA is much larger in the DIPS dataset than any of the others.

The PPI BSA values of Group 1 complexes span a wide range from the smallest value of 423 Å² seen in the FKBP12-rapamycin-FRAP complex to the largest value of 2336 Å² seen in the DDB1-CR8-CDK12 complex. Five of these complexes have reported protein–protein binding affinity data in the absence of the molecular glue (Table 1). Fig. 2B plots the log values of the protein–protein dissociation constants without the molecular glue, $\ln(K_D)$, against the PPI BSA for these five complexes. There is a rough correlation between the two, with interfaces that have larger PPI BSA also registering better protein–protein binding, even though the data is extremely limited. There is also one outlier, namely the DDB1-CR8-CDK12 complex (PDB ID: 6TD3). It has a large BSA PPI at 2,336 Å², but the binding affinity between DDB1 and CDK12 without CR8 is low ($K_D \approx 50 \mu\text{M}$).¹⁸ This is because when CR8 is not present, the energy cost for CDK12 to adopt the bound conformation is likely high, leading to the observed low affinity.

The protein–protein binding affinities of Group 2 complexes in the absence of their corresponding molecular glues span a similar range compared to those in Group 1 (Table 1). The weakest of them is the TIR-IAA7 complex with a K_D at 18.5 μM and the strongest is the complex formed between 14-3-3 and the Gab2 peptide with a K_D of 0.5 μM . Additionally, as many



interacting sequence motifs in Group 2 have key, specific interactions with the effector protein, such as buried hydrogen bonds and salt bridge interactions, these interfaces are perhaps more efficient than those largely non-specific interactions present in complexes from the other datasets. For the eight complexes where protein–protein dissociation K_D exist, plotting the $\ln(K_D)$ against PPI BSA shows little correlation and almost the opposite trend as that seen in Group 1 complexes (Fig. 2B). The complex formed between β -TrCP and its substrate pSer33 β -catenin peptide (PDB ID: 6M90) only has a BSA PPI of 509 Å² but has one of the highest affinities in all the complexes with known protein–protein binding K_D (K_D = 530 nM).²⁵ The efficiency of this interface with such a small PPI BSA likely stems from the partially buried charge–charge interactions between the phosphorylated Ser33 and the surrounding residues on the effector protein. On the other hand, the complex formed between the plant hormone receptor COI1 and its JAZ1 degron (PDB ID: 3OGM) has the largest PPI BSA at 1279 Å² in this set but only a moderate protein–protein K_D of 5 μ M is observed.⁴⁷ This shows that the nature of the interactions formed in complexes belonging to Group 2 is likely different than those formed between folded domains.

Amino acid preference at the interface

Amino acid preference profiles show which residues are more likely to be observed in a class of PPI. Fig. 3 plots these for all 20 amino acids at the PPI from complexes in four datasets described above. While Group 1 complexes exhibit similar residue preference profiles as those in datasets DB5.5 and DIPS, complexes in Group 2 show some differences for a few amino acids: Gln, Thr, His, Met, Trp, Phe, and Pro. Based on two-tail student *t*-tests, some of these differences are statistically significant, including those for Gln, His, Phe, and Pro. Although the sample size is still quite small, some of these trends may be meaningful. For example, Thr and Pro only exist in some sequence motifs but not others likely because phosphorylated Thr serves as a chemical signal for protein recognition and Pro is known to introduce kinks in protein structures that may provide the basis for shape recognition. A similar analysis done by Hou and coworkers found that by clustering and analyzing sequences of eukaryotic linear motifs,⁴⁸ which form reversible interactions with protein partners to illicit downstream effects, they could identify residue preferences in different types of sequence motifs.⁴⁹ Such preferences should be harnessed to discover neo-substrates and potential glues for



Fig. 3 Interface amino acid preference profiles. The protocol below describes how the preference profile is computed. First, an interface residue list is generated for a complex. Each residue is counted once even if there are multiple occurrences at an interface. The lists of all complexes in a dataset are added up based on residue types and normalized by the total number of complexes in the dataset. The value of the preference is between zero and one. With a zero indicating a residue that is missing from all PPI in the dataset and one indicating that this residue is ubiquitous in all PPI in the dataset at hand. Interface residues are defined as residues that are within 4.5 Å of any non-hydrogen atoms from the interacting protein. Residues with modifications like phosphorylation and protonation are counted as their original form. The 20 residues are color by their properties: negatively charged (red), positively charged (blue), polar (magenta), hydrophobic (green), and others (yellow). Phosphorylated residues (Ser, Thr, and Tyr) are represented with a darker shade of color. The residues with statistically significant difference in residue preference between Group 1 and 2 are marked with a star. These include Gln, His, Phe, and Pro with *p*-values at 0.0292, 0.0003, 0.0292, and 0.0008, respectively.



E3 ubiquitin ligases. The implication of the notable differences between the interfaces formed in Groups 1 and 2 is that different computational schemes may need to be applied for their prediction.

Residue-residue pair preference at the interface

To understand whether the specific interacting residues in the PPI of the four different types of complexes show similarity or differences, we analyzed the frequency of interacting residue pairs. Fig. 4 plots the residue-residue pair frequencies seen at the PPI. As expected, interactions between hydrophobic residues are seen ubiquitously in all four groups of interfaces, especially between Leu, Phe, and the other hydrophobic residues. Another residue that is seen often in all PPIs is Arg. The salt bridge interactions between Arg and the negatively charged residues are observed in nearly all interfaces. This highlights the seminal role of hydrophobic and electrostatic interactions in the formations of PPIs, and this is common in all PPI including those induced by molecular glues. Interactions between polar residues (Asn, Gln, Ser, Thr, Tyr, His) are slightly less frequent in PPI from Group 1 and 2 than those in the DB5.5 and DIPS datasets but given the small sample size in the MG-PPI, the significance of such difference is unclear. Nevertheless, this analysis illustrates that the main driving force of all PPI formation, regardless of whether a small molecule is needed to stabilize the interface, is hydrophobic and electrostatic.

This implies that prediction algorithms that rely on physics- or statistics-based modeling to predict PPIs without molecular glues may still be valuable.

Ternary structure formation mechanism imparted by ligand–protein interactions

Aside from protein–protein interactions, molecular glue-induced complex structures offer an opportunity to examine the interactions between the small molecule and the two protein partners. Fig. 5 plots the two ligand–protein BSA in the representative structures taken from Table 1. The diagonal line in Fig. 5 indicates a region where the two ligand–protein BSA in a complex are equal. As the complex structures deviate from this diagonal region, increasing asymmetry is seen between the interfaces formed between the molecular glue and the two binding partner proteins. The asymmetry of interactions between the molecular glue and its protein binding partners can guide the understanding of their ternary structure formation.

As may be expected for molecular glues, the majority of compounds show asymmetric binding where the ligand BSA is larger for one protein in the PPI compared to the other. This is more prominent in complexes from Group 2, likely due to two factors: (1) unresolved structure of disordered regions in one

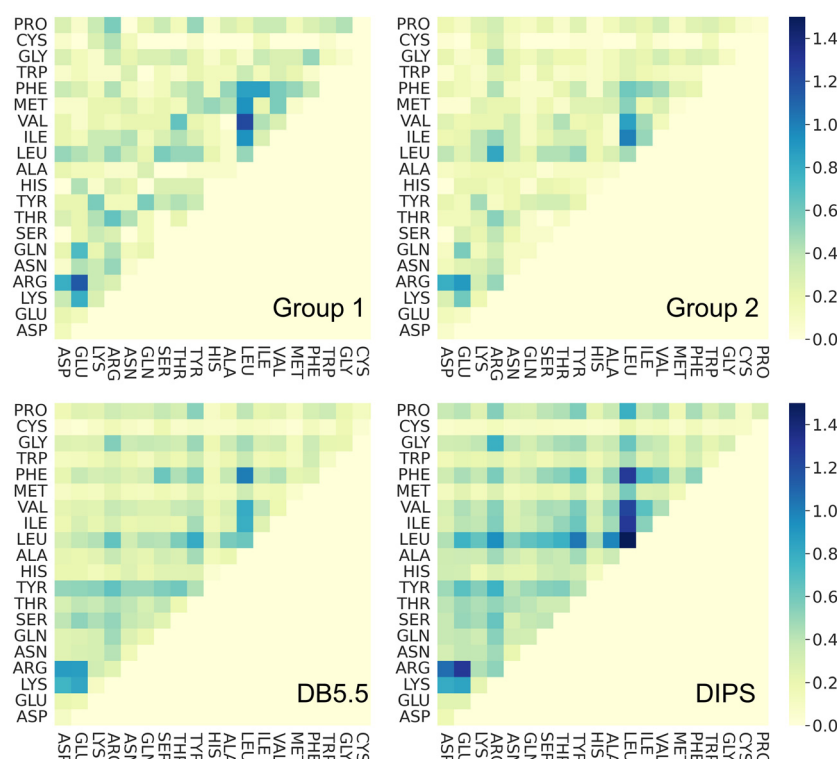


Fig. 4 Residue pair preference profiles. The residue pair preference profile is calculated using the procedure below. All pairs of residue–residue interaction counts are recorded for an interface, resulting in a 20 by 20 matrix. The interaction matrices are then averaged over all complexes in the dataset without normalization. The higher the value, the more prevalent such pair of interaction is observed in a dataset. Interacting residue pairs are defined as those with any non-hydrogen atoms within 4.5 Å of each other and the two residues must be from two different proteins. Residues with modifications like phosphorylation and protonation are counted as their original form.



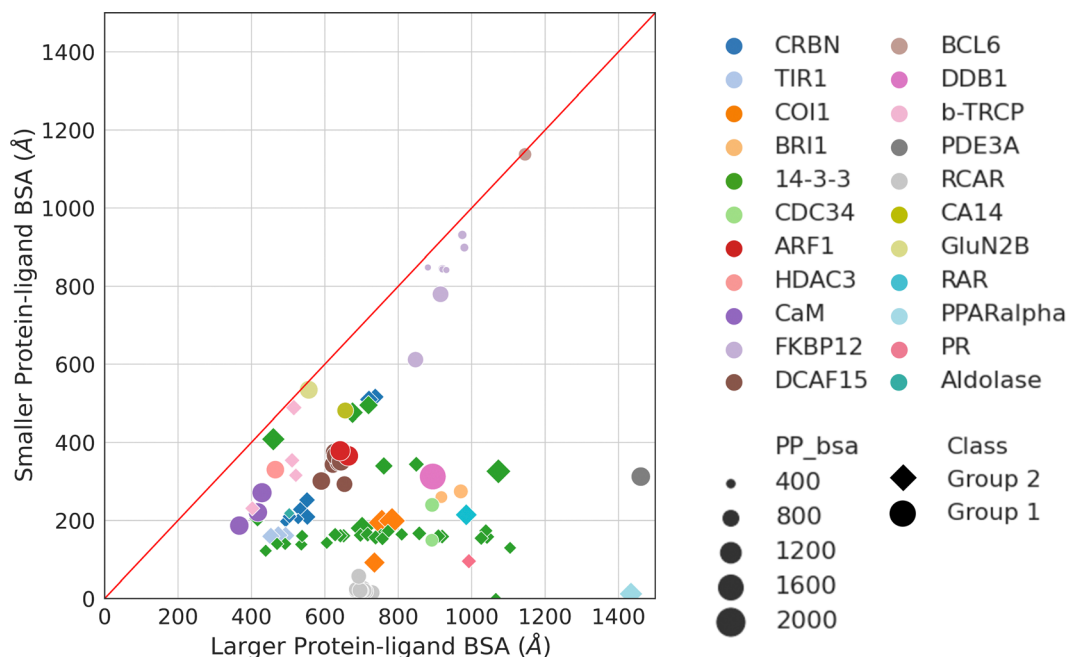


Fig. 5 Ligand–protein BSA of all structures in Table 1. For those effector proteins with multiple binding partners (e.g., FKBP12 and 14-3-3), all of them are included in this plot with the same color and shape indicating the same effector protein. Ligand–protein BSA is computed in a similar fashion as the PPI BSA. In a molecular glue induced ternary complex, the ligand (*i.e.*, molecular glue) interacts with both protein partners. The BSA of the two interfaces are calculated separately. For each of the ligand–protein interface, the SASA is computed for the two components alone and for the complex. The difference is then calculated and reported as the final ligand–protein BSA. The two ligand–protein BSA for a complex are then ranked by their magnitudes, resulting in a smaller and a larger BSA. Plotting the smaller BSA against larger BSA resulted in this plot. The complexes are colored by their effector proteins, with the shape indicating which group of interaction mode they belong to (Group 1: circle and Group 2: diamond). The size of the symbols indicates the size of the PPI BSA of the complex. The PDB IDs of these complexes can be found in Table 1.

protein partner, usually the one containing the sequence motif, and (2) intrinsic binding between the glue and one of the proteins even without the second protein partner, which will be discussed further below. The most asymmetric cases lie in the bottom right corner, for example the complex structure formed between the peroxisome proliferator-activated receptor- α (PPAR α) and the silencing mediator for retinoid and thyroid hormone receptors, SMRT (PDBID: 1KKQ⁵⁰). There is an extensive interface between the glue GW6471 and PPAR α while little interaction is seen between GW6471 and SMRT. This is supported by the already existing binding affinity between GW6471 and PPAR α .⁵⁰ Without GW6471, the binding between PPAR α and SMRT is weak, with a K_D of $8.0 \pm 3.4 \mu\text{M}$ measured by fluorescence polarization assay.⁵⁰ Adding the molecular glue enhances the K_D by 5-fold. By comparing the receptor structures bound with antagonist GW6471 and agonist GW409544, it is clear that the observed binding cooperativity lies in the induced conformational change of the AF-2 helix upon GW6471 binding, which moves and exposes a surface that is favorable for SMRT binding (Fig. 6A). Another example is the CR8-induced ternary complex between DDB1 and CDK12-CyclinK (PDBID:6TD3).¹⁸ CR8 is a known CDK inhibitor⁵¹ and the crystal structure of the ternary complex shows that it has extensive interactions with CDK12. Without CR8 the binding between CDK12-CyclinK and DDB1 is minimal with a K_D of $\sim 50 \mu\text{M}$ measured by isothermal titration calorimetry (ITC).¹⁸

Binding of CR8 likely shifts the distribution of CDK12 conformations to one that favors binding to DDB1 ($K_D < 100 \text{ nM}$, measured by ITC¹⁸). Fig. 6B shows the C-terminal structural difference between an inactive CDK12 in complex with ADP and the CR8 bound DDB1-CDK12/CyclinK complex. When CR8 is bound to CDK12, it disrupts the interactions between the C-terminal short helix and the C-lobe of the kinase as seen in the inactive state, causing the C-terminus to adopt a conformation that can be recognized by DDB1, leading to the degradation of CyclinK. In both complexes which show pronounced asymmetry, the ternary structure formation is driven by ligand–protein binding, which induces allosteric conformational change of the protein surface, making it recognizable by the binding partner.

A portion of molecular glue induced complexes have roughly similar buried surface area between the molecular glue and the two protein binding partners. These are presented as data-points close to the diagonal line in Fig. 5. Some complexes belong to this category do not show any protein–protein binding without the molecular glue. For example, the complex formed by FKBP12, rapamycin, and FRAP (PDB ID: 1FAP) has no detectable interactions between FKBP12 and FRAP. However, rapamycin is known to have high affinity binding to FKBP12 with a K_D of 0.2 nM .^{52,53} It also binds to FRAP but weakly ($K_D = 26 \mu\text{M}$ ⁵³). This can be seen from Fig. 6C, where the PPI BSA is rather small but there is a deep, concaved binding



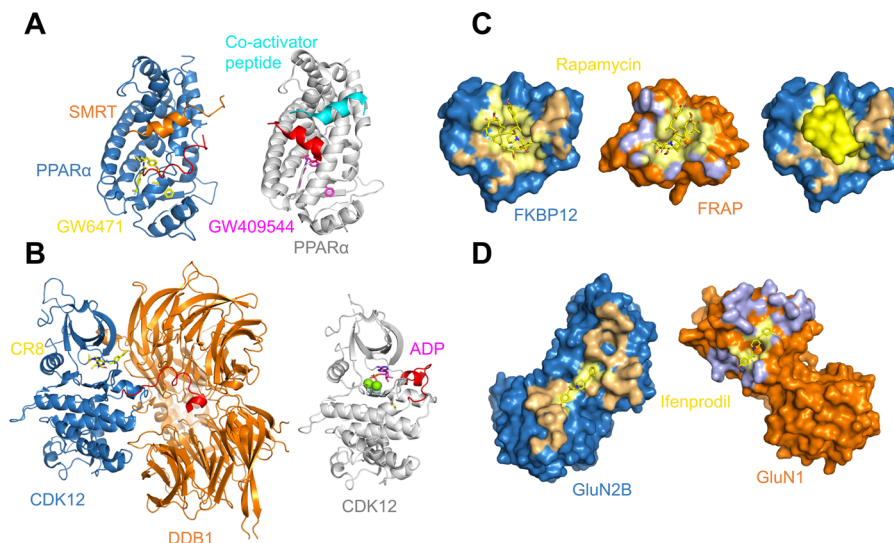


Fig. 6 (A) Binding of antagonist GW6471 (yellow) to the PPAR α receptor (blue) alters the AF-2 helix (red) conformation (left; PDB ID: 1KKQ) compared to when PPAR α is bound to an agonist GW409544 (magenta) (right; PDB ID: 1K7L). The conformational difference in the AF-2 helix dictates the binding partner protein whether it is the co-repressor peptide SMRT (left; orange) or a co-activator peptide (right; cyan). (B) Binding of CR8 (yellow) to CDK12 (blue) alters the conformation of its C-terminal region (red) (left; PDB ID: 6TD3) compared to when CDK12 is inactive and bound to ADP (magenta) (right; PDB ID: 4NST). The new conformation of the CDK12 C-terminus resulted from CR8 binding is well-suited to bind to DDB1 (left; orange). (C) The protein binding area (light orange) and the ligand binding pocket (light yellow) in FKBP12 (blue) shown in surface presentation (left). The protein binding area (light blue) and the ligand binding surface (light yellow) in FRAP (orange) shown in surface presentation (middle). Binding of rapamycin (yellow) changes the surface of FKBP12 (blue), making it favorable for FRAP binding (right). The structures are taken from the FKBP12-rapamycin-FRAP complex (PDB ID: 1FAP). (D) The protein binding area (light orange) and the ligand binding surface (light yellow) in GluN2B (blue) shown in surface presentation (left). The protein binding area (light blue) and the ligand binding surface (light yellow) in GluN1 (orange) shown in surface presentation. The structures are taken from the GluN2B-ifenprodil-GluN1 complex (PDB ID: 3QEL).

site on FKBP12 used to bind rapamycin. The rapamycin binding surface on FRAP is shallower in line with the weaker K_D between two. Even though rapamycin only binds weakly to FRAP, the FKBP12-rapamycin complex binds to FRAP in much higher affinity (ternary $K_D = 12$ nM) revealed by SPR measurements.⁵³ This is likely due to the binding of rapamycin to FKBP12 creating a new composite surface, driving the recruitment of FRAP.

In the diagonal region of Fig. 5, there are also complexes with intrinsic protein-protein binding affinities even without the presence of molecular glues. Adding molecular glues to the mix further strengthens the PPIs, leading to stable ternary complexes. This is exemplified by the complex formed between the NMDA receptor amino-terminal domains GluN1, GluN2B, and ifenprodil (PDB ID: 3QEL).⁵⁴ In the ternary structure, the interface areas between ifenprodil and the two amino-terminal domains are almost identical, and there are no obvious pockets on either of the receptor subunits that indicate ifenprodil-subunit binding (Fig. 6D). This coincides with the observation that no binding could be detected between ifenprodil with either GluN1 or GluN2B.⁵⁴ Without ifenprodil, GluN1 and GluN2B do interact and form a heterodimer with a K_D of 0.7–1 μ M when mixed. In fact, there is extensive PPI between the two proteins in the ternary complex with a PPI BSA at 1,229 \AA^2 (Fig. 6D). When ifenprodil is introduced, a 20-fold dimer stabilization effect is seen. The ternary structure formation in this case likely starts with the dimerization of GluN1

and GluN2B, forming a binding pocket for ifenprodil. Once ifenprodil is bound, it further stabilizes the complex.

Analysis of molecular glue-induced ternary structures indicate that there are likely two main pathways of how the ternary structure is formed (Fig. 7); either a protein-protein binary interface is formed first and then the complex incorporates a small molecule ligand at the interface, which further enhances binding (Path 1 in Fig. 7), or a small molecule binds to a protein partner first, which leads to altered protein surface properties, much like the effects of post-translational modification,³⁴ making this surface available for association with another protein (Path 2 in Fig. 7). A similar scheme of molecular glue-induced ternary structure formation has been proposed by Cao and coworkers.⁵⁵ The two pathways for ternary structure formation are not independent of each other, even though in some complexes one may dominate over the other as is the case in GluN2B-ifenprodil-GluN1 and FKBP12-rapamycin-FRAP (Fig. 7). There are cases where both paths are exploited to form a ternary complex. One example is CRBN-lenalidomide-CK1 α , where relatively weak binary binding exists between the two proteins, as well as between CRBN and lenalidomide. Even though it is tempting to categorize all known molecular glue-induced complexes into these three categories, it remains a difficult task until experimental affinity for the binary components in the ternary systems are measured and reported, especially for those that are weak. Fig. 7 also reveals that the ternary structure formation process is a closed reaction cycle



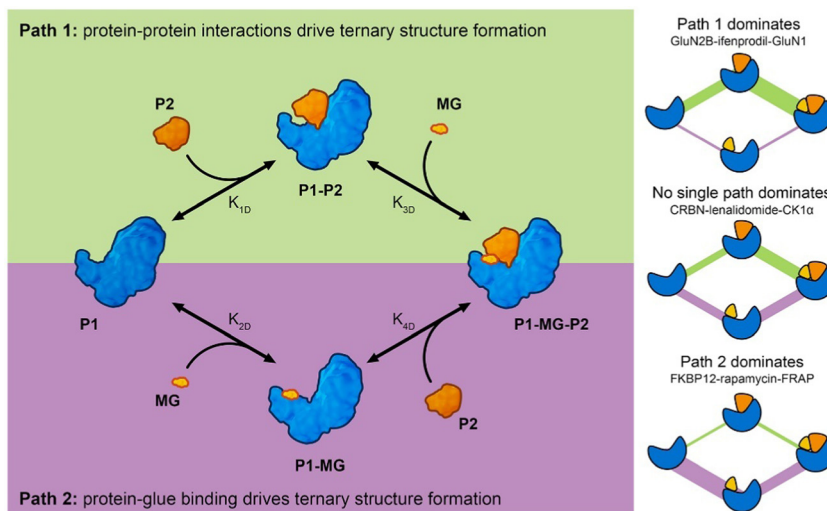


Fig. 7 Molecular glue-induced ternary structure formation mechanism. (left) The elementary reactions leading to ternary complexes. There are two paths for ternary structure formation. Path 1 starts from protein–protein binding while Path 2 starts from protein and molecular glue binding. Both paths meet at the final product – the formed ternary complex. (right) Three examples showing complexes with different ternary structure formation mechanisms. In the complex formed by GluN2B, ifenprodil, and GluN1, Path 1 dominates with $K_{1D, \text{GluN2B-GluN1}} = 1 \mu\text{M}$, $K_{2D, \text{GluN2B-ifenprodil}} = \text{N.D.}$, and $K_{3D, \text{GluN2B-ifenprodil-GluN1}} = 50 \text{ nM}$. In the complex formed by CRBN, lenalidomide, and CK1 α , no single path dominates the ternary structure formation ($K_{1D, \text{CRBN-CK1}\alpha} = 2 \mu\text{M}$, $K_{2D, \text{CRBN-lenalidomide}} = 180 \text{ nM}$, $K_{4D, \text{CRBN-lenalidomide-CK1}\alpha} = 12 \text{ nM}$). In the complex formed by FKBP12-*rapamycin*-FRAP, Path 2 dominates with $K_{1D, \text{FKBP12-FRAP}} = \text{N.D.}$, $K_{2D, \text{FKBP12-*rapamycin*}} = 0.2 \text{ nM}$, $K_{4D, \text{FKBP12-*rapamycin*-FRAP}} = 12 \text{ nM}$. The color schemes of the components follow those in the graph on the left. These components include P1, P1–P2, P1–MG–P2, and P1–MG starting from the left, going clockwise. The edges connecting the symbols represent the reaction fluxes between different states. The thickness is a rough indication of the amount of flux with thicker lines representing higher flux and route dominance.

with every elementary step reversible. Because of this feature, the K_D of all four elementary reactions are related by the cooperativity (α) of the system, where $\alpha = K_{1D}/K_{4D} = K_{2D}/K_{3D}$. This provides the basis for further kinetic modeling of these systems. Such a framework is useful in the rational discovery of molecular glues and neosubstrates. The implication of this analysis is that different ternary structure formation mechanisms should be taken into consideration as one searches for molecular glues and neosubstrates of E3 ubiquitin ligases. Strategies for molecular glue discovery where ternary complexes are driven primarily by protein–protein binding *versus* ligand–protein binding need to be considered differently.

Computational approaches for identifying weak protein–protein interfaces

Discovering weak protein–protein interactions that can be further stabilized is key to develop molecular glues. Rational design of molecular glues has been difficult mainly due to a lack of understanding and predictability of weak interactions, for example between the protein of interest and the E3 ubiquitin ligase. As outlined above, the molecular glue can simply enhance an already existing PPI or it can induce one *via* tight binding to a single protein first. Here, we focus on the first scenario, identifying existing, weak, fortuitous interactions between two proteins could be the first step in rational design and discovery of new molecular glues. However, this approach

remains challenging. For known MG–PPI, the binding affinity between the two proteins without the molecular glue can be in the μM range,^{18,25,47,55,56} too weak for structure determination approaches such as X-ray crystallography. NMR spectroscopy can be used to probe weak protein–protein interactions,⁵⁷ but the throughput is slow and therefore is ill-suited for the task of screening many potential protein pairs. Computational methods of PPI prediction could be leveraged to fill the gap of identifying PPIs with weak intrinsic binding affinity.

Computational approaches for predicting protein–protein complex structures typically involve two steps, complex generation and ranking. The two tasks can be achieved *via* either physics-based or machine learning (ML)-based and deep learning (DL)-based algorithms. Physics-based approaches rely on the underlying physics of the system and require rather little *a priori* knowledge, while ML/DL models leverage existing protein–protein complex data to learn the correct patterns for prediction.

Physics-based protein–protein docking methods can further be divided into *ab initio* approaches, or homology-based approaches to reduce the search space. The homology-based approach searches for homologous complexes with known structures (*e.g.*, templates), which are used to guide docking pose generation.^{58–62} The *ab initio* approach uses a variety of methods to search the conformational space of the PPIs, including fast Fourier transform^{63–65} and geometric hashing,^{66,67} and spherical polar Fourier correlations.^{68,69} Both approaches provide candidate docked poses called decoys. *Ab initio* docking is especially helpful when no homologous



complex structures exist of the queries, as is largely the case for molecular glue induced complexes. This makes *ab initio* docking an attractive tool for generating candidate structures of gluable PPI. Recently, DL techniques have also been used to generate protein–protein complex decoys and have provided results that are on par or better than those generated using traditional physics-based docking approaches and with much less computing time.^{70,71}

After docking poses are generated, a scoring scheme is employed to rank them so that the most plausible structure can be selected. There are two different types of scoring functions, physics-based and knowledge-based. Physics-based scoring functions usually include a weighted, linear combination of energy terms such as van der Waals, electrostatics, and desolvation energies.^{72,73} They can also include empirical terms such as shape and chemical complementarity of the interacting surfaces and the size of buried interfaces.^{74,75} The optimal weights of these terms need to be determined using known protein–protein complex systems. Knowledge-based scoring functions typically take the shape of a statistical potential that is constructed using known complex structures.^{76–79} The frequencies of interactions between atoms or residues are recorded for a set of known, diverse complexes. By applying Boltzmann's equation to the ratio of the observed and expected frequencies for an atom/residue pair, one can estimate its potential of mean force. A total score can then be computed for a given interface as a contribution from all interacting atom/residue pairs and is used in ranking.

Another avenue for developing these special scoring functions is to use a ML model trained on appropriate datasets. There are already several ML-based decoy generation and ranking algorithms^{46,70,80–83} publicly available. One example is the MaSIF framework⁷⁰ which uses protein surface patches with precomputed geometric and physicochemical features as inputs. It then applies a geometric neural network to the inputs to generate fingerprint representations. The fingerprints of potential binding partners are scanned and those that show complementarity were used to generate decoys, followed by reranking using another trained neural network. The performance on a test set of 100 complexes is on par with commonly used physics-based docking programs PatchDock⁸⁴ and ZDOCK^{85,86} but with a fraction of the computational cost. A recent development of the framework, dMaSIF,⁸⁷ is a differentiable end-to-end DL model built on the MaSIF framework. It does not require precomputed features and uses a point cloud to represent the protein surface. The change introduced by dMaSIF led to a significant improvement in run time and memory use while maintaining a similar level of performance as MaSIF.

Inspired by the MaSIF and dMaSIF protein–protein interaction prediction workflow, Orasch and coworkers⁸⁸ devised an DL model that learns the most suitable embeddings for predicting protein–protein interactions using point cloud representation of protein surfaces together with precomputed features. They trained this model using the MaSIF dataset and a more diverse orthogonal dataset and then applied it to predict

protein–protein interfaces formed upon PROTAC binding. For a set of 16 complexes, an average area under the receiver operating characteristic curve (ROC AUC) of 0.87 is achieved, much better than a random model which would give an ROC AUC of 0.5. This indicates that the features present in the training set of known protein–protein complexes are likely important in the PROTAC-induced protein–protein interfaces as well. One caveat of the study is that no precision or recall values were reported for the PROTAC-induced complex set. These are important measures of model performance and should be part of model performance evaluation. Still, this study highlights the potential of using ML and DL in predicting weak interfaces, like the ones induced by molecular glues and PROTACs, especially when appropriate datasets are used in training.

A more rigorous approach for ranking the quality of protein–protein poses is to compute the free energy of association. This can be achieved using free energy simulations coupled with enhanced sampling techniques. There are many flavors of free energy calculations. The most commonly used ones are free energy perturbation (FEP),^{89–91} thermodynamic integration (TI),⁹² and potential of mean force (PMF) calculations.⁹³ The first two are routinely used to predict binding free energies between proteins and ligands in the drug discovery settings and have shown promise in terms of accuracy and efficiency.^{94,95} FEP calculations are also starting to gain popularity in mutation effect prediction at protein–protein binding interfaces as encouraging results emerge.^{96–98} PMF calculations can be used to compute the absolute binding free energy of protein complexes. For example, umbrella sampling simulations^{99,100} impose a series of biases along a reaction coordinate, in this case the distance between the two binding partners, guiding the dissociation or association of the complex. Methods like the weighted-histogram analysis method^{101,102} or multistate Bennett acceptance ratio¹⁰³ can then be used to obtain the unbiased the population distribution along the reaction coordinate, from which the PMF along the reaction coordinate can be computed. The binding free energy is the PMF difference between the associated and fully dissociated states. This approach has been used to study the association of well-characterized barstar–barnase¹⁰⁴ protein complex and the predicted binding free energy is within 2 kcal/mol of experimental value.¹⁰⁵ Other techniques like metadynamics¹⁰⁶ and adaptive bias force^{107–109} simulations can also be used to sample the conformational space upon association or dissociation and obtain the PMF, from which the binding free energy is extracted. A recent study by Wang *et al.*, showcased the utility of metadynamics in protein–protein binding free energy predictions.¹¹⁰ The dissociation PMF profiles and the binding free energies of 19 distinctive protein–protein complexes were computed and compared with the corresponding experimental measurements. A remarkable R^2 of 0.74 was observed between the predicted and the experimental binding free energy values, indicating a high degree of correlation. The main advantage of applying the PMF approach to compute binding free energy is that it uses explicit representation for both the solvent



(e.g., water) and the solute atoms and employs molecular dynamics simulations to incorporate conformational flexibility. Even though it produces accurate binding free energies, the computational cost is also considerable. Adequate sampling is the key for success in PMF-based free energy calculations. In the metadynamics study,¹¹⁰ 50 replicas and an accumulative simulation time of 2 μ s are required to obtain the converged binding free energy for each system tested. However, given the low throughput this approach is restricted for use when a small subset of decoys has already been chosen for analysis or when no other alternatives are available.

Limitations still exist for computational prediction of complex structures. One major obstacle is the lack of ability to incorporate structural rearrangement upon complex formation. Most physics-based and ML-based methods work well on complexes formed between rigid components, but the performance drops when one or both monomers undergo conformational change upon complex formation.^{59,111,112} Techniques such as soft docking,^{113–115} ensemble docking,^{116,117} sidechain optimization,^{72,118,119} and adaptive conformer generation¹²⁰ can be applied in *ab initio* docking to overcome some of the flexibility issues, especially those involving small and local changes (e.g., sidechain rearrangement). Predicting complexes involving large conformational change upon binding is still challenging for *ab initio* docking. With ML-based methods, by choosing the appropriate protein representation and model architecture, one may be well-positioned when faced with complex prediction challenges where flexibility is involved. For example, a coarse-grained representation of protein surfaces using residue-level instead of atom-level features will likely be less sensitive to sidechain conformational changes and produce reasonable results for those with small induced conformational changes upon binding.^{112,121} DL methods like AlphaFold^{71,122} and RoseTTAFold¹²³ use multiple sequence alignment and residue pairwise relations to represent protein structures. These representations are processed by multiple neural networks to predict the relative distances between residues, which are then used to guide the folding of a single chain protein or the binding of two proteins. Because of these representations, such methods are able to succeed in predicting complex structures that undergo large conformational changes upon binding and are traditionally deemed as hard targets.¹²⁴

Another factor preventing the application of these methods in predicting molecular glue-induced PPI is accuracy. Many of the current scoring functions in docking programs and ML/DL methods are developed using complex datasets with little or no presence of MG-PPI and thus fail. For example, AlphaFold-multimer⁷¹ relies on co-evolutionary information encoded in the multiple sequence alignments to guide the complex prediction, but most of the MG-PPI do not have this intrinsic property, except for those involving disease causing mutations on the native substrates of E3 ubiquitin ligases, such as those pS33/pS37 degron mutants and β -TRCP.²⁵ This review serves as a first attempt to understand the interactions governing MG-PPI and how they differ from other protein–protein complexes.

With the development of special scoring functions and new, appropriate training datasets we can overcome this important problem.

Concluding discussion

Molecular glue-induced ternary complex is an umbrella term used to describe all protein complexes that require a third entity, usually a small molecule, to assemble or to gain stability. As there is appreciable difference in all known molecular glues,³⁶ there is heterogeneity in the interfaces they induce, as well as the mechanisms of how the ternary complexes are formed. In this review, we divided the PPIs stabilized by molecular glues into two categories, those that involve interfaces from well-folded protein domains and those that feature specific sequence motif and folded domain interactions. The interaction patterns differ for these two types of interfaces with the first showing similarity to the rest of protein–protein interfaces seen in the PDB. The second category of interfaces involve sequence motifs spanning three to thirteen amino acids, many of which have signature interaction motifs such as phosphorylated Ser/Thr and leucine zipper. The different nature of these two types of PPIs demands separate discovery strategies.

Based on the currently known molecular glue-induced ternary complexes, the mechanism of ternary structure formation varies for different complexes. On one extreme of the spectrum, ternary structure formation starts from a small molecule binding to one of the protein partners, either altering its surface properties or its conformational distribution, making it more attractive to the other protein partner. On the other end of the spectrum, the two proteins come together first with some binding affinity. The newly formed complex then offers a binding pocket for a molecular glue to bind and stabilize the ternary complex. Most of the ternary complexes likely form using a mixture of these two mechanisms. In essence, the formation of ternary complexes involving molecular glues is similar to that induced by bi-specific degraders.¹²⁵ From monomers, there are two pathways to form a ternary complex. Each path contains two reactions, binary and ternary formation (Fig. 7). The K_D of these reactions are constrained by cooperativity. Three-component systems like these have been studied extensively.^{125,126} This means that one can interrogate these ternary systems using analytical and numerical simulations. For example, ordinary differential equations can be set up to describe the system of reactions. For a given initial condition, with the K_D known, one may compute the amount of ternary complexes formed. On the other hand, for a desired amount of ternary structure formation, one may back calculate the K_D that produce such an amount and use this information to guide the identification of molecular glues and neosubstrates.

In the context of E3 ubiquitin ligases, the rational discovery of neosubstrates and molecular glues is difficult because one needs to consider both variables simultaneously. To simplify this problem, it is useful to narrow down the search space by



reducing the number of candidate neosubstrates or molecular glues. In the effort to reduce the number of neosubstrates, being able to predict which target protein and E3 ubiquitin ligase pairs are likely to form weak protein–protein interactions that can be further stabilized by small molecules is key. Direct prediction of whether two proteins could come together with the help of a small molecule using cutting edge ML algorithms is challenging due to the lack of appropriate datasets since the number and diversity of known molecular glue-induced complexes is limited. Predicting the binding affinities between two protein partners in a high throughput fashion is still an open challenge. However, protein–protein decoy generation and ranking algorithms, both physics-based and ML-based, show promise in finding the most likely complexes formed between two proteins.^{70,84–88} These complexes then need to be evaluated by their ability to bind and to be stabilized by small molecules. Such a problem is similar to those in traditional small molecule drug discovery where one needs to identify effective binders for the targets of interest. Experimentally, this is achieved by performing high throughput screens and hit validation followed by lead optimization. Computational approaches such as binding pocket identification,^{127–132} protein–ligand docking,^{133–138} virtual screening,^{139–141} and ligand binding free energy calculations^{94,95,142,143} can also be applied and have been shown to be helpful in this area. With a combination of computational and experimental approaches, rational discovery of molecular glues may indeed be feasible, even though the best and the most effective solution to this task is yet to be determined.

Recent development of DL models trained on large datasets to predict protein structures^{122,123} and protein–protein complex structures^{71,144,145} puts ML/DL methods in the spotlight in all fields of biology. Methods inspired by these ML/DL programs can be extremely helpful in predicting potential complexes between two proteins of interest. Recently, Tsaban *et al.*¹⁴⁶ revealed that AlphaFold2 can be used to predict peptide–protein complexes with reasonable accuracy without using multiple sequence alignment information for the peptide portion based on the hypothesis that binding of a peptide to its well-folded partner is similar to the final step of protein folding,¹⁴⁷ where the last piece of unstructured fragment folds into the folded portion. DL methods like AlphaFold and RosettaFold also hold potential to incorporate conformational changes into complex structure prediction. This is especially helpful for predicting complexes like DDB1–CR8–CDK12, where binding of CR8 to apo CDK12 leads to large conformational change of the latter, revealing a new surface for DDB1 association. By applying attention mechanism to key interactions and folding the complex from scratch, one may finally overcome the challenge of flexible docking. The research in this area is still in early stage but it has been shown in a recent NeurIPS conference paper that using an iterative transformer network, the DL approach, GeoDock¹⁴⁸ is able to predict backbone conformational changes upon protein–protein complex formation. Another example of using DL in protein–protein complex prediction is the protein surface embedding developed in the

MaSIF framework⁷⁰ trained using a large structural dataset. This embedding could be used in different kinds of predictions tasks. Language models like BERT (Bidirectional Encoder Representations from Transformers)¹⁴⁹ have also been shown to be able to learn protein representations that encode their fundamental properties such as secondary structure, binding site hotspots, and sites for post-translational modification.^{150,151} Using such a model trained on degron data from the ELM database, Hou and coworkers⁴⁹ show that they can predict new degrons and assign them to E3 ubiquitin ligases. In conclusion, the outlook for molecular glue discovery is bright and ML/DL methods hold tremendous potential to aid in this process. Framing the question correctly and constructing appropriate datasets are crucial elements to success.

Lastly, being able to predict whether and how ternary structures would form is only solving part of the puzzle. Ultimately, one needs to design small molecule glues that can induce ternary structure formation and degradation of disease relevant neo-substrates. Ternary structure formation does not always guarantee degradation. This has been thoroughly discussed in the review on enzymology of degraders.¹⁵² A few recent studies have used physics modeling^{153,154} and ML techniques¹⁵⁵ to predict or rationalize degradability of PROTAC molecules. These studies all point to lysines, especially those that are E2-accessible ubiquitination sites, are indicative of degradation potential. It is difficult to say how generalizable these approaches are as they are mostly based on a handful of E3 ligases. The majority of E3 ligases are underexplored in terms of their structures and their ability to engage in molecular glue-induced degradation. The field is in its early days, but it is quickly evolving. Many exciting new studies are well under way. Concerted efforts from both computational and experimental groups are required to fully understand and harness the E3 ligase-mediated degradation system.

Conflicts of interest

All authors are employees and stockholders for Amgen, Inc.

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