

**Progress towards B-Cell Lymphoma 6 BTB Domain Inhibitors for the Treatment of
Diffuse Large B-Cell Lymphoma and Beyond**

Yong Ai¹, Lucia Hwang¹, Alexander D. MacKerell Jr.¹, Ari Melnick² and Fengtian Xue*¹

¹Department of Pharmaceutical Sciences, University of Maryland School of Pharmacy, 20 Penn Street, Baltimore, Maryland 21201, United States

²Department of Hematology and Oncology, Weill Cornell Medical College, New York, New York 10021, United States

²Department of Pharmacology, Weill Cornell Medical College, New York, New York 10021, United States

Abstract

B-cell lymphoma 6 (BCL6) is a master regulator of germinal center formation that produce antibody-secreting plasma cells and memory B-cells for sustained immune responses. The BTB domain of BCL6 (BCL6^{BTB}) forms a homodimer that mediates transcriptional repression by recruiting its co-repressor proteins to form biologically functional transcriptional complex. The protein-protein interaction (PPI) between the BCL6^{BTB} and its co-repressors has emerged as a therapeutic target for the treatment of DLBCL and a number of other human cancers. This perspective provides an overview of recent advances in the development of BCL6^{BTB} inhibitors from reversible inhibitors, irreversible inhibitors, to BCL6 degraders. Inhibitor design and medicinal chemistry strategies for the development of novel compounds will be provided. The binding mode of new inhibitors to BCL6^{BTB} are highlighted. Also, the *in vitro* and *in vivo* assays used for the evaluation of new compounds will be discussed.

Introduction

Diffuse large B-cell lymphoma (DLBCL) is the most common type of non-Hodgkin's lymphomas (NHL) throughout the world with over 18,000 new cases each year. DLBCL is a fast growing cancer arising from germinal center (GC) B-cells.¹ GCs are transient structures formed within lymphoid follicles responding to antigenic stimulation.² The transcriptional repressor protein B-cell lymphoma 6 (BCL6) is needed for B-cells to form GCs.¹ Normally BCL6 is upregulated when B-cells are activated to form GCs and downregulated once the GC reaction is complete.¹ Failure to downregulate BCL6 leads to continuous expression of this transcriptional repressor, which is required to maintain the survival of DLBCL cell lines and primary human DLBCL tumors.³⁻⁷ BCL6 regulates lymphomagenesis by directly repressing DNA damage sensing genes as well as cell cycle and cell death and checkpoint genes.^{3,8-13} In addition, BCL6 also blocks differentiation of DLBCL cells by repressing the *PRDM1* locus that encodes a master regulator for plasma cell differentiation.^{13,14} BCL6 knockdown or inhibition of its repressive activity causes expression of these many checkpoint genes, resulting in proliferation arrest and cell death.^{3-7,15}

BCL6 has been reported as an oncoprotein in many other types of human tumors. Follicular lymphomas (FLs), the second most frequent type of NHL, are documented to be biologically dependent on the repressive function of BCL6 in GC B-cells.¹⁶ In addition, BCL6 is highly upregulated in BCR-ABL positive B-acute lymphoblastic leukemia (B-ALL) cells after treatment by tyrosine kinase inhibitors.¹⁷ Upregulation of BCL6 prevented B-ALL cells from being killed by kinase inhibitors, however, the combination of ABL kinase inhibitor plus BCL6 blockade synergistically eradicated B-ALLs *in vivo*.¹⁷ Moreover, BCL6 is constitutively upregulated in mixed-lineage leukemia (MLL) fusion oncoproteins in B-ALLs with MLL translocation, while suppression of BCL6 kills primary human MLL translocated B-ALL cells.¹⁸ Patients with chronic

myeloid leukemia (CML) are resistant to tyrosine kinase inhibitor imatinib because CML stem cells are also dependent on BCL6,¹⁹ and blockade of BCL6 activity eliminated these critical leukemia-repopulating cells. Furthermore, BCL6 is highly expressed in acute myeloid leukemia (AML) stem cells, and blocking BCL6 kills AML cells and synergizes with chemotherapy.²⁰ BCL6 is also required to maintain the survival of “triple-negative” breast cancer cells.²¹ Recently, BCL6 was characterized as a promising therapeutic target in non-small cell lung cancers, which feature frequent amplification of the BCL6 locus.^{22,23} Taken together, BCL6 has broad oncogenic roles in different cancers and BCL6 targeted therapy can potentially benefit many patients beyond those with lymphomas.

BCL6 has an *N*-terminal broad-complex, tramtrack, and bric-a-brac (BTB) domain (BCL6^{BTB}) that controls transcriptional repression and a *C*-terminal C2H2 zinc finger that binds to a specific DNA consensus sequence.²⁴ The two domains are connected by an unstructured region containing a second repression domain (RD2).²⁵ The BCL6^{BTB} forms a homodimer (Figure 1) that recruits corepressors silencing mediator for retinoid and thyroid receptor (SMRT), nuclear receptor corepressor (NCOR) and BCL6 corepressor (BCOR).^{26,27} The minimal binding domain of these corepressors contains a conserved 18 amino-acid peptide known as BCL6 binding domain (BBD)^{28,29} that binds to a lateral groove (LG) formed at the interface between BCL6^{BTB} monomers (Figure 1).^{28,29} This LG/BBD interaction is essential for the transcriptional repression activity of BCL6^{BTB}.^{28,29} Interestingly, the key BCL6^{BTB} surface residues that contact the BBD are not conserved in any of the other BTB domains.³⁰

Besides GC B-cells, BCL6 also plays essential role in the development of follicular T-helper cells and anti-inflammatory effects in macrophages.³¹ Experimental evidence has shown that BCL6 knockout mice are runted, born at sub-Mendelian ratios and die within weeks of acute

inflammatory diseases.³¹ However, in contrast to BCL6 knockout mice, those with a BCL6^{BTB} mutant that cannot bind to the corepressors, were born at expected rates, and lived normal healthy lives and had no evidence of inflammatory syndrome. The only observed phenotype was failure to form GC B-cells in response to antigenic challenge.³² Therefore, the function of BTB LG is only essential in GC B-cells and in tumors, but is inessential for the function of BCL6 in other lineages. The effect of BCL6 in macrophages, for instance, is mediated by its DNA-binding zinc finger domain, while not affected by targeting the BTB LG.³² Transient suppression of GCs is expected to be non-toxic in humans. Thus, BCL6^{BTB} LG inhibitors represents a safe therapeutic target.

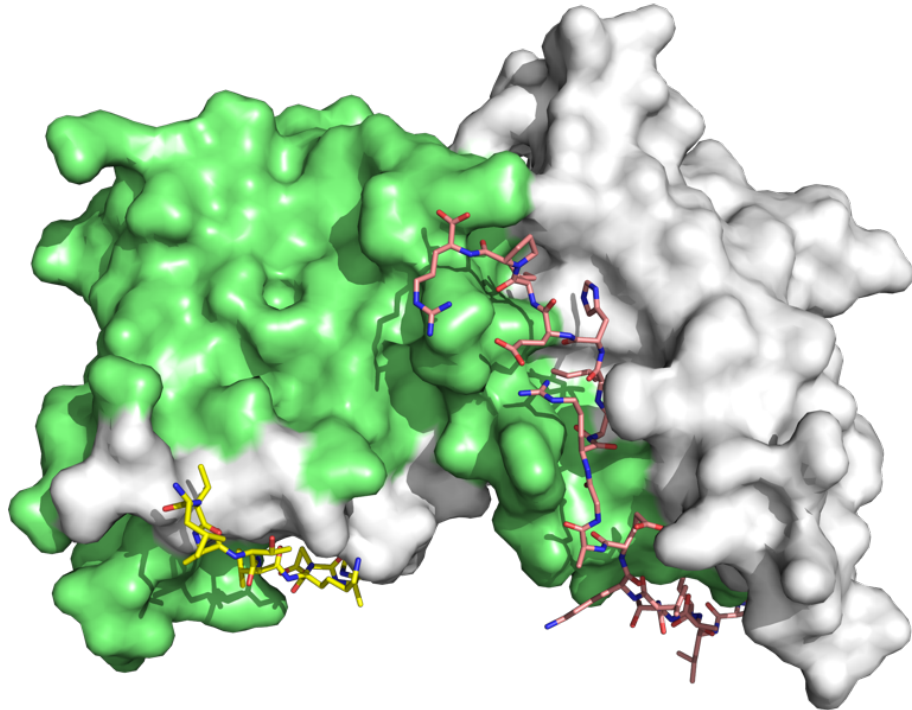


Figure 1. Surface view of the binding mode of the SMRT peptide (LVATVKEAGRSIHEIPR) to BCL6^{BTB} LG (PDB ID: 1R2B). The monomers of BCL6^{BTB} were shown in green and white, respectively. The SMRT peptides were shown in pink and yellow, respectively.

Numerous inhibitors of BCL6^{BTB} have been developed. In pioneering work, Polo and coworkers developed a BCL6 peptide inhibitor (BPI),⁴ which occupied the BTB LG, prevented recruitment of the corepressors by BCL6 and induced expression of BCL6-target genes.⁴ BPI also had potent anti-lymphoma activity, inducing apoptosis and growth arrest in BCL6-dependent DLBCL cell lines but not for BCL6-independent cell lines.⁴ Injections of BPI in mice reproduced the BCL6 null GC phenotype.⁴ Refinement of the BPI yielded a retro-inverso peptidomimetic RI-BPI that bind to BCL6^{BTB} LG and potently killed DLBCL cells *in vitro* and *in vivo* and primary human DLBCL cells *ex vivo*.⁵

A fundamental lesson of these studies was that the duration of occupancy of the BCL6^{BTB} LG was critical for killing of DLBCL cells. Whereas a single dose of short acting decoy peptides with $t_{1/2} < 6$ h could not kill DLBCL cells, the longer acting RI-BPI, which has a nuclear dwell time up to 24 h *in vitro* and *in vivo* did kill DLBCL cells similar to BCL6 siRNA or shRNA. Duration of target engagement is a key parameter for BCL6 targeted therapy.^{33,34} Notably, because BCL6 blockade reactivates checkpoints and cell death, remarkable synergy between RI-BPI and chemotherapy was observed.¹² The lack of toxicity of LG blockade makes BCL6 inhibitors ideally suited to serve as “anchors” for developing novel combination therapy regimens. In sum, BCL6 presents a highly promising therapeutic target that can be effectively and specifically blocked by occluding the BCL6^{BTB} LG.

Early small molecule BCL6^{BTB} inhibitors 1 (79-6)⁶ and 2 (FX1)⁷. The discovery and activity profiles of BCL6^{BTB} inhibitors **1** and **2** have been reviewed previously (Figure 2A).^{35,36} In silico screening hit **1** could disrupt BCL6 repression complexes and kill DLBCL cells.⁶ Subsequent SAR efforts yielded inhibitor **2**,⁷ which bound to BCL6 with affinity similar to endogenous corepressors, exhibited a prolonged dwell time in cells, and mirrored the efficacy and specificity of RI-BPI *in*

vitro by killing BCL6 dependent but not BCL6 independent DLBCL cell lines. Inhibitor **2** was nontoxic and demonstrated promising efficacy *in vivo* in both germinal center B-cell (GCB) and activated B-cell (ABC) type DLBCL xenograft models.⁷ The binding modes of these inhibitors have been characterized by X-ray crystallography and NMR. The indolin-2-one rings of compounds **1** and **2** occupy an aromatic pocket (named the aromatic site) on the top region of the BCL6 LG (red, Figure 2B). The aromatic site is formed by four key residues: R'24, L'25 from one BCL6^{BTB} monomer, and residues M51 and Y58 from the other. The same aromatic site is occupied by corepressor SMRT residues H1426 and I1428,²⁷ or BCOR residues W509 and V511.^{28,29} Also, the side chains of L25' and M51 line a hydrophobic pocket that interacts with the Br atom in inhibitor **1**. The carboxylic acid group of the inhibitor fits snugly into the acid site (cyan, Figure 2B) and forms electrostatic interaction with R'28 (Figure 2C). The NH group of the indolin-2-one ring forms an H-bond (2.5 Å) with the backbone carbonyl O of residue M51. Adjacent to the aromatic site, a shallow hydrophobic pocket (named linker site, blue, Figure 2B), occupied by the side chain of a tryptophan residue in the SMRT protein, is available with the residue E115 located 7.5 Å away from the indolin-2-one ring (Figure 2C). Importantly, the backbone amide NH group, as well as the carboxylate side chain of E115 provide opportunities for interactions such as hydrogen bonds (H-bonds) and electrostatic interactions. Moreover, a large and deep pocket (H'14-D'17-C53-H116), known as the HDCH site (yellow, Figure 2B), is located below residue E115, providing an additional pocket for inhibitor design. The presence of residue C53 in the HDCH site offers a unique opportunity for the development of irreversible inhibitors for BCL6^{BTB}. The specificity of inhibitors can be designed into irreversible inhibitors by exploiting the unique chemical characteristics and spatial orientation of the aromatic and linker sites, as well as the novel HDCH site, to impart specific binding before covalent bond formation.

These early inhibitors **1** and **2** provided proof of principle for targeting the BCL6^{BTB} LG; however, the PAINS classification of the rhodanine moiety³⁷ motivated the design of novel BCL6 inhibitors with improved affinity while lacking undesirable PAINS moieties, which will be the focus of the current review.

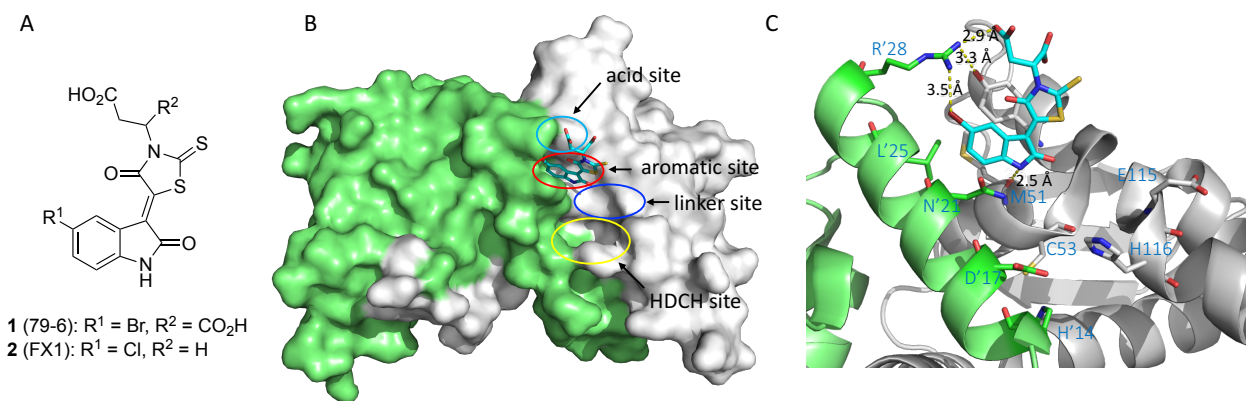


Figure 2. (A) Chemical structures of early BCL6 inhibitors **1** and **2**. (B) Binding mode of compound **1** to BCL6^{BTB} (PDB ID: 3LBZ). Compound **1** was shown in cyan and BCL6^{BTB} monomers are shown in green and gray, respectively. The binding pocket of BCL6^{BTB} LG includes four continuous binding sites: acid site (cyan), aromatic site (red), linker site (blue) and HDCH site (yellow). (C) Detailed binding mode of inhibitor **1** in complex with BCL6^{BTB}. The key residues from BCL6^{BTB} were shown as sticks. The distances of highlighted H-bond interactions were shown in angstrom (Å).

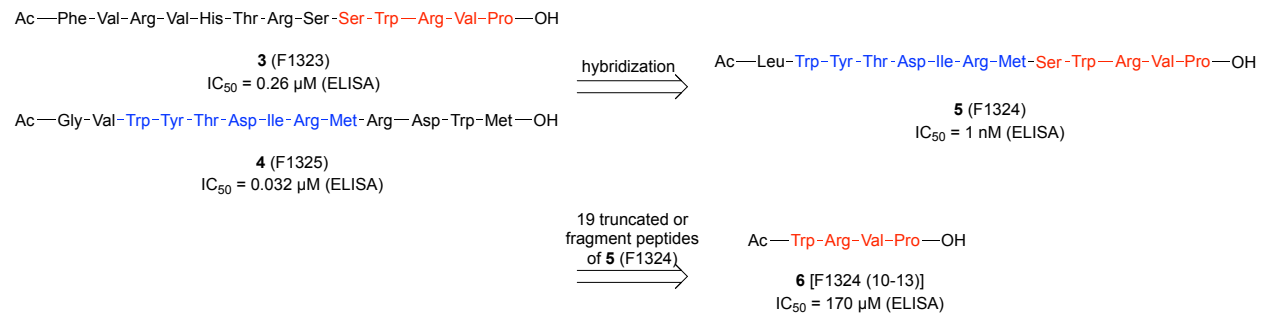
Recent Development of BCL6^{BTB} Inhibitors

Reversible Inhibitors

Sakamoto and coworkers disclosed their efforts using the technology of phage-displayed peptide library to identify novel BPIs (Scheme 1).³⁸ Specifically, the authors panned T7 phage-displayed peptide libraries against the FLAG-BCL6^{BTB} and control proteins (TWEAK and IL17),^{39,40} and determined the binding affinity of resultant polyclonal phage pools using an enzyme-linked

immunosorbent assay (ELISA) on microplates by following the absorbance changes ($\lambda = 450$ nm). After five rounds of panning, two peptides, W-Y/F/I/V/L-T/S-D-I/V/L-R-M and W/F/Y-R/K-V/I-P, were identified. Next, the authors used two representative sequences FVRVHTRSSWRVP (**3**, F1323) and GVWYTDIIRM RDWM (**4**, F1325) to design a hybrid peptide LWYTDIIRMSWRVP (**5**, F1324).

Scheme 1. Development and sequence structure of the hybrid BPI **5 (F1324)**



The inhibitory potencies of peptides **3-5** were evaluated using cell-free ELISA assays. Initial peptides **3** and **4** indicated IC_{50} values of $0.26 \mu M$ and $0.032 \mu M$, respectively. The hybrid **5** was a potent inhibitor of BCL6 with an IC_{50} value of 1 nM , which was $\sim 3,000$ -fold more potent than that of the BCL6 corepressor BCOR peptide ($IC_{50} = 3 \mu M$). Note that the BCOR peptide was biotinylated and its sequence is detailed in Table 1. Next, the kinetics for the binding of peptide **5** was assessed using surface plasmon resonance (SPR). The K_d value of peptide **5** was determined to be 0.57 nM , which is $10,000$ -fold more potent than that of BCOR peptide ($K_d = 6,000 \text{ nM}$). The k_{on} and k_{off} values were calculated to be $2.8 \times 10^6 \text{ M}^{-1} \text{ s}^{-1}$ and $1.6 \times 10^{-3} \text{ s}^{-1}$, respectively. The binding half-life ($t_{1/2}$) of peptide **5** was determined to be 441 s , significantly longer than that of the BCOR peptide ($t_{1/2} < 1 \text{ s}$). These results supported that the potent inhibitory activity of peptide **5** against BCL6^{BTB} was likely due to the small k_{off} value.

To identify key amino acid residues responsible for the high inhibitory potency of peptide **5**, the authors synthesized and tested 19 fragment peptides. While most of the *N*-terminal truncated

peptides showed dramatically decreased potencies, the *C*-terminus tetrapeptide WRVP (**6**, Scheme 1) displayed promising inhibition activity ($IC_{50} = 170 \mu\text{M}$), which is even more potent than longer sequences MSWRVP ($IC_{50} = 620 \mu\text{M}$) and SWRVP ($IC_{50} = 1,500 \mu\text{M}$). Besides, replacing the *C*-terminus carboxylate of WRVP, which occupies the acid site of BCL6^{BTB} LG (Figure 4), with an amide group caused a significant decrease in inhibitory potency ($IC_{50} = 170 \mu\text{M}$ to $IC_{50} = 2,100 \mu\text{M}$), highlighting the importance of this carboxylate group for the potency of WRVP.

Although peptide **5** was highly potent in cell-free assays, the authors found this tetrapeptide had no significant inhibitory activity in cellular assays. While various peptide fusion strategies with cell-permeable peptides failed to improve the cellular activity, the authors pursued a cell-based mammalian two-hybrid (M2H) assay²⁷ to deliver peptide **5** into cells. Specifically, the authors built the plasmid coding AcGFP, a nuclear localization sequence DPKKKRKV, and peptide **5**. The expression level of the peptide was confirmed by following the fluorescence signal of AcGFP.⁴¹ The results confirmed that the expressed peptide AcGFP-(PKKKRKV)₃-GGG-**5** localized in the nucleus, causing decreased luminescence from the transcription of the Luc gene by inhibiting the PPI between BCL6^{BTB} and BCOR. In a separate experiment, expression of BCORpep-containing sequence AcGFP-(PKKKRKV)₃-GGG-BCORpep failed to decrease the luminescence signal, indicating that the level of the expressed peptide was significantly less than the K_d value of BCOR peptide.

The co-crystal structure of the complex BCL6^{BTB}-**5** (PDB 5H7G) was solved at 1.85 Å (Figure 3). Peptide **5** binds to BCL6^{BTB} LG in a binding mode similar to those of the SMRT and BCOR peptides. The side chains of amino acids L1, R7, and R11 were completely disordered and those of W2, D5, and P13 were partly disordered. The peptide backbones of W2 and T4 made H-bonds with those of Q10 (2.7 Å) and T12 (2.9 Å) of BCL6^{BTB}, respectively. In addition, the M8,

W10, and V12 backbones formed H-bonds with the sidechains of D'17 (2.8 Å), N'21 (3.0 Å), and R'24 (2.8 Å) of BCL6^{BTB}, respectively (Figure 3). The side-chains of amino acids W10 and P13 formed hydrophobic interactions with residue Y58 of BCL6^{BTB}. The C-terminal -CO₂H of amino acid P13 snugged into the acid site of LG and formed H-bonds with the guanidine group of R'28 (3.3 Å). The *N*-terminus of peptide **5** interacts with a β -strand in the bottom of the BCL6^{BTB} LG. The sidechain of amino acid M8 is deeply buried in the HDCH site of BCL6^{BTB} LG, with a significant conformational change of H116 in BCL6^{BTB}. This methionine residue is substituted by serine residues in both the analog BPI **3** and the BCOR peptide, and an isoleucine (I1425) in the SMRT peptide.

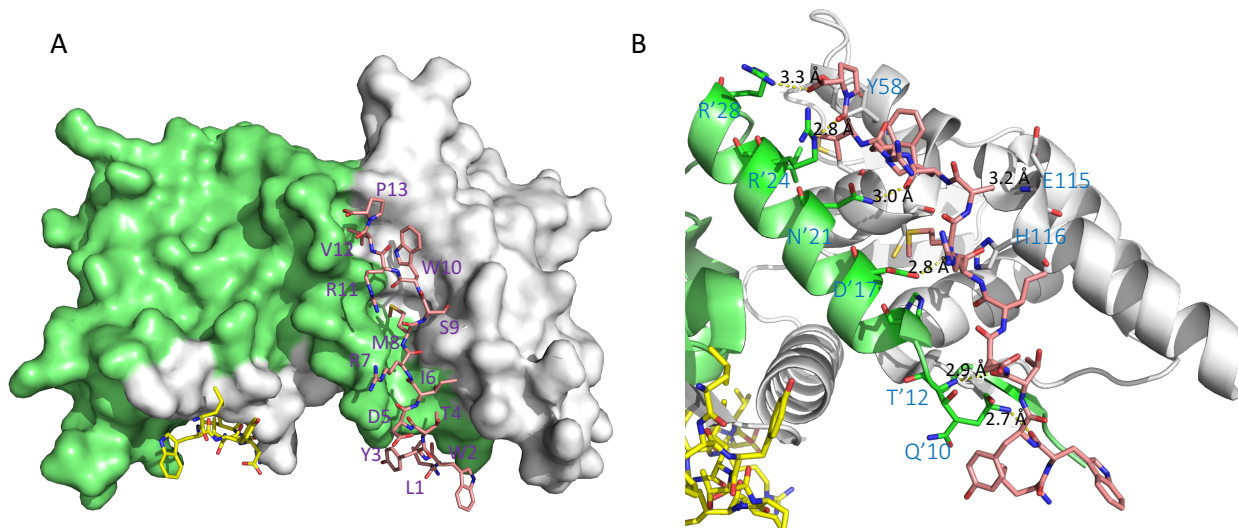


Figure 3. Binding mode of BPI **5** (F1324) to BCL6^{BTB} (PDB ID: 5H7G). (A) Surface view of the binding of BPI **5** to the BCL6^{BTB} LG. Peptide **5** was shown in pink and BCL6^{BTB} monomers are shown in green and gray, respective. (B) Detailed binding mode of inhibitor **5** in complex with BCL6^{BTB}.

The authors also studied the binding mode of the tetrapeptide WRVP (**6**) to BCL6^{BTB} (Figure 4). Interestingly, the truncate peptide **6** occupied the same site as peptide **5** (Figure 4A), with highly similar H-bond patterns to BCL6^{BTB} LG. The carboxylate group of P13 interacted

simultaneously to guanidine groups of R'24 and R'28 in the acid site (Figure 4B). The isopropyl group of V12 fit snugly into the aromatic site while the backbone NH group of V12 and carbonyl O atom of W10 formed two H-bonds to the sidechain of residue N'21. In addition, the guanidine group of R11 formed electrostatic interaction to residue D'17 and the indole NH of W10 formed H-bond to the backbone carbonyl O of residue M51. This unique binding mode of tetrapeptide **6** provided a potential scaffold for effective inhibition of BCL6^{BTB}.

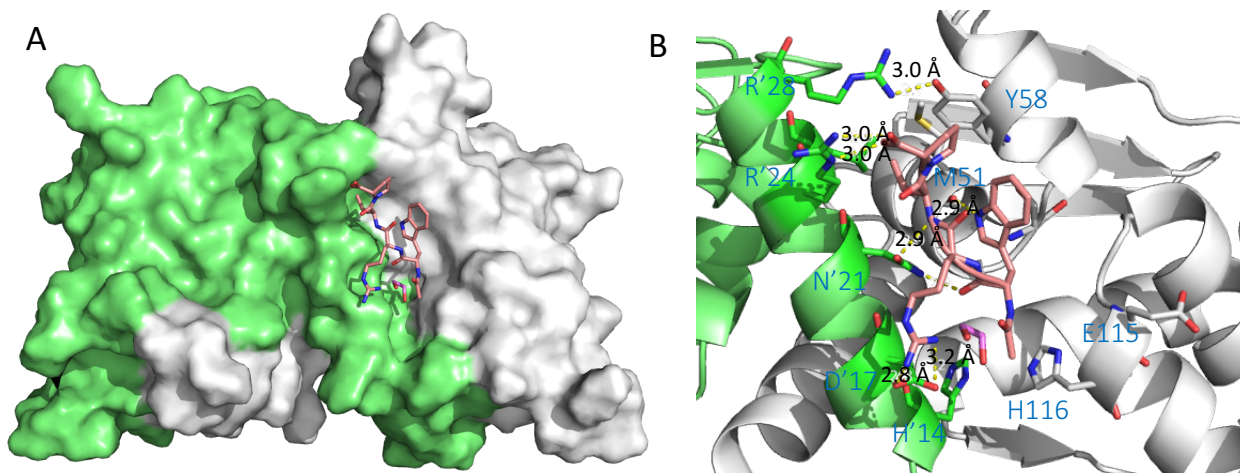


Figure 4. Binding mode of peptide **6** (Ac-WRVP-OH) to BCL6^{BTB} (PDB ID: 5H7H). (A) Surface view of the binding of peptide **6** to the BCL6^{BTB} LG. Peptide **6** was shown in pink and BCL6^{BTB} monomers are shown in green and gray, respective. (B) Detailed binding mode of inhibitor **6** in complex with BCL6^{BTB}.

By screening a library of ~130,000 compounds using ELISA, Yasui and coworkers discovered diphenylamine hits **7** and **8** with IC₅₀ values of 4.5 μ M and 14 μ M, respectively (Scheme 2).⁴² To confirm the binding of these hits to the BCL6^{BTB}, the authors used surface plasmon resonance (SPR) and determined the K_d values of compounds **7** and **8** to be 5.0 μ M and 24 μ M, respectively. The co-crystal structure of BCL6^{BTB} dimer in complex with hit **7** (Figure 5) showed the inhibitor bind to the top part of BCL6^{BTB} LG, the same site as those of compound **1** and tetrapeptide **6**. Specifically, the left ring of inhibitor **7** occupied the aromatic site formed by

sidechains of Y58, N'21 as well as with R'24, and R'28 in the acid site. The Cl atom of the hit was found to fit into the small hydrophobic pocket formed by three residues M51, L'25, and A52. Large substituents (e.g., Br and CF_3) were not tolerated at this position. A key H-bond (3.0 Å) was formed between the diphenylamino NH group and the backbone O atom of residue M51. Another important H-bond (3.1 Å) was detected between the benzimidazolone O atom and the backbone NH group of residue E115.

Scheme 2. Optimization strategy of HTS hits **7** and **8** and identification of compounds **10** and **13**

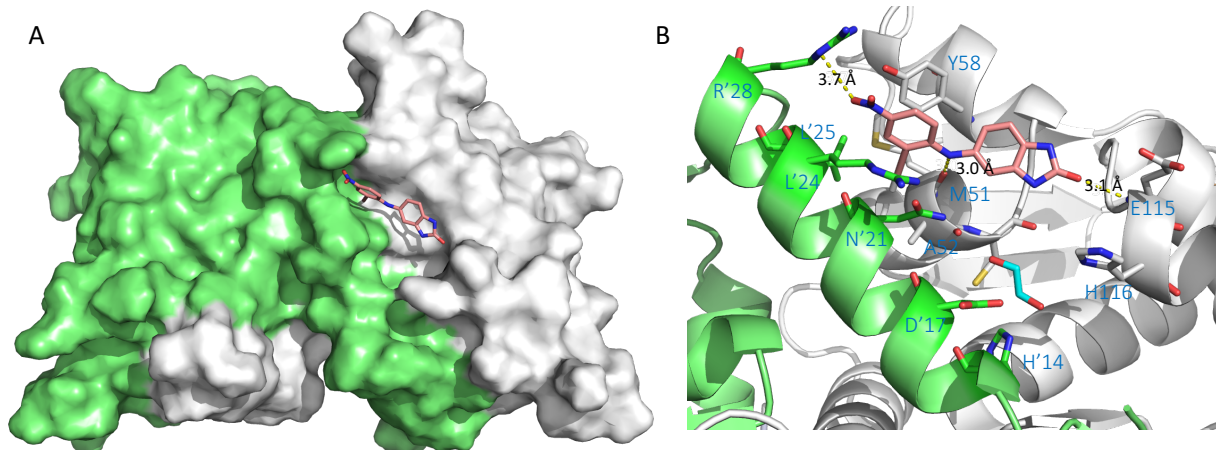
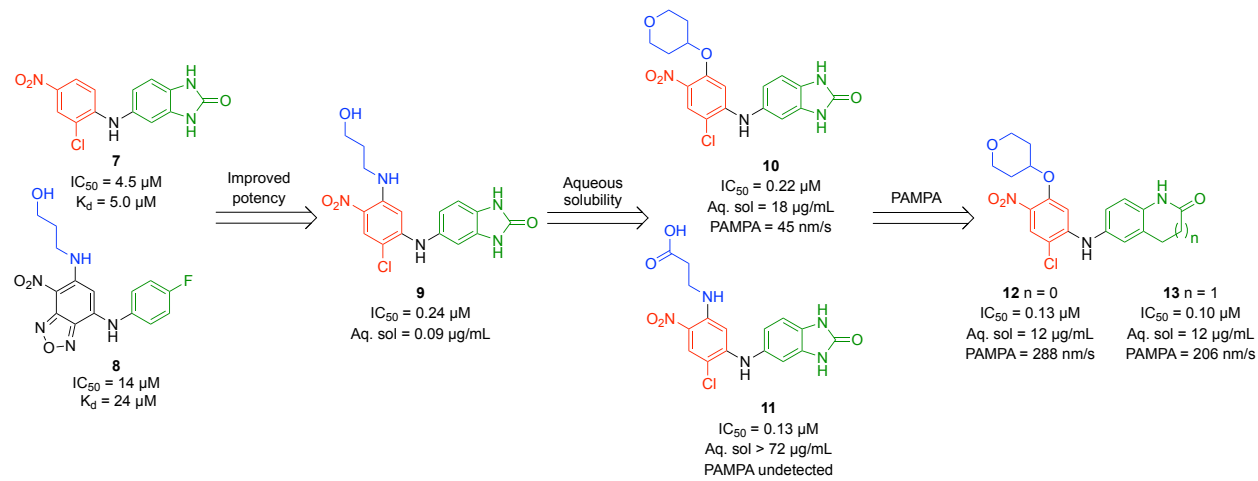


Figure 5. Binding mode of hit **7** to BCL6^{BTB} (PDB ID: 5X9O). (A) Surface view of the binding of hit **7** to the BCL6^{BTB} LG. Compound **7** was shown in pink and BCL6^{BTB} monomers are shown in green and gray, respectively. (B) Detailed binding mode of inhibitor **7** in complex with BCL6^{BTB} .

Suggested by docking results of hit **8** in BCL6^{BTB}, the hydroxypropylamino group of compound **8** was introduced to the left ring of hit **7**, resulting in the hybridized inhibitor **9**, with a 20-fold improvement in potency ($IC_{50} = 0.24 \mu M$) compared to that of hit **7** (Scheme 2). To improve physiochemical properties of hybrid inhibitor **9** (solubility: $0.09 \mu g/mL$, parallel artificial membrane permeability assay (PAMPA): not detected), five additional analogs with different substituents at the 5-position of the left ring were prepared to identify the 4-tetrahydropyranloxy analog **10** and carboxylic acid **11**. Compounds **10** and **11** indicated significantly increased aqueous solubility (**10**: $18 \mu g/mL$, **11**: $> 72 \mu g/mL$)⁴³ without compromising their potencies. The 4-tetrahydropyranloxy analog **10** kept modest PAMPA permeability (45 nm/s) although the carboxylic acid analog **11** failed to show detectable PAMPA permeability, likely due to the presence of the carboxylic acid functionality. To improve membrane permeability of inhibitor **10**, the authors explored the substitution effects on the right ring of the scaffold and reported that the two H-bond donors in the benzimidazolone group contributed to the low PAMPA permeability. Accordingly, oxindole analog **12** and tetrahydroquinolinone analog **13** demonstrated improved PAMPA permeabilities (288 nm/s and 206 nm/s, respectively) compared to that of 4-tetrahydropyranloxy **10**, while maintaining potency and aqueous solubilities (**12**, $IC_{50} = 0.13 \mu M$, aqueous solubility = $12 \mu g/mL$; **13**, $IC_{50} = 0.10 \mu M$, aqueous solubility = $12 \mu g/mL$ vs. **10**, $IC_{50} = 0.22 \mu M$, aqueous solubility = $18 \mu g/mL$). Besides, compounds **12** and **13** exhibited no cytotoxicity at the concentration of $30 \mu M$ in HepG2 cells.

Next, inhibitors **12** and **13** were tested in cellular M2H assays. Both inhibitors indicated good cellular potencies with IC_{50} values of $1.2 \mu M$ and $0.72 \mu M$, respectively. As a control, the unoptimized parent compound **9** demonstrated no cellular activity ($IC_{50} > 100 \mu M$). The PK profile of the more potent inhibitor **13** was studied (0.1 mg/kg i.v.; 1 mg/kg p.o.), and the results showed

that compound **13** had a good PK profile (MRT = 3.3 h, AUC = 1.27 $\mu\text{g}\cdot\text{h}/\text{mL}$, F = 79.9%), making this compound a promising candidate for future *in vitro* and *in vivo* studies.⁴⁴

The binding mode of aqueous soluble analog **11** was determined by X-ray crystallography (PDB 5X9P). As shown in Figure 6, the left ring moiety of compound **11** is positioned in the aromatic site composed of amino acid side chains of Y58, N'21, L'25, and R'28. The NO₂ group of inhibitor **11** interacted with the guanidinium side chain of R'28. The Cl at the 2-position of compound **11** was found to occupy the relatively small hydrophobic pocket formed by three residues M51, L'25, and N'21. Interestingly, the carboxylic acid group of inhibitor **11** was solvent-exposing and largely disordered, while no obvious electrostatic interaction was formed between the carboxylate and the guanidine side chain of R'28 in the acid site. The linker NH group formed a clear hydrogen bond (2.8 Å) with the backbone carbonyl group of the residue M51. Finally, the bicyclic lactam occupied the linker site and the lactam carbonyl O formed a key H-bond (3.1 Å) with the backbone NH of E115.

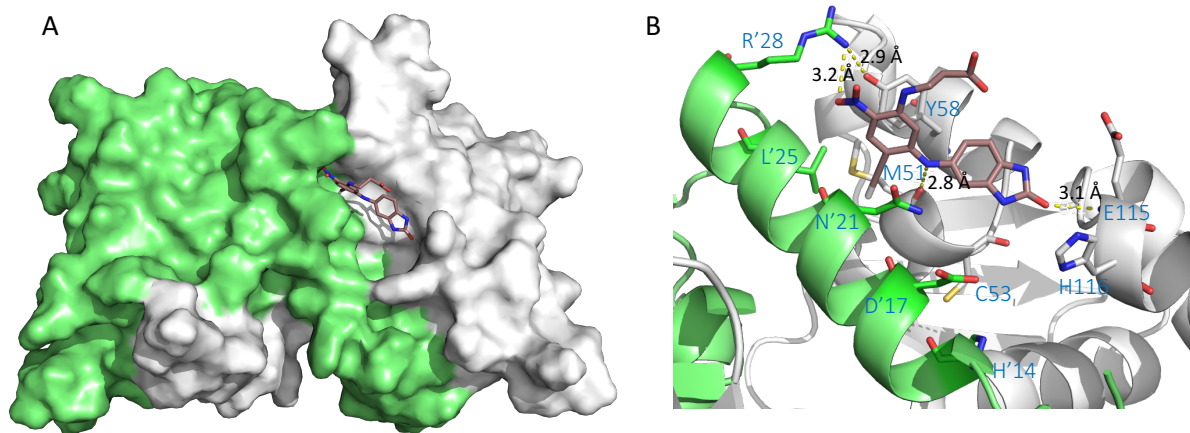
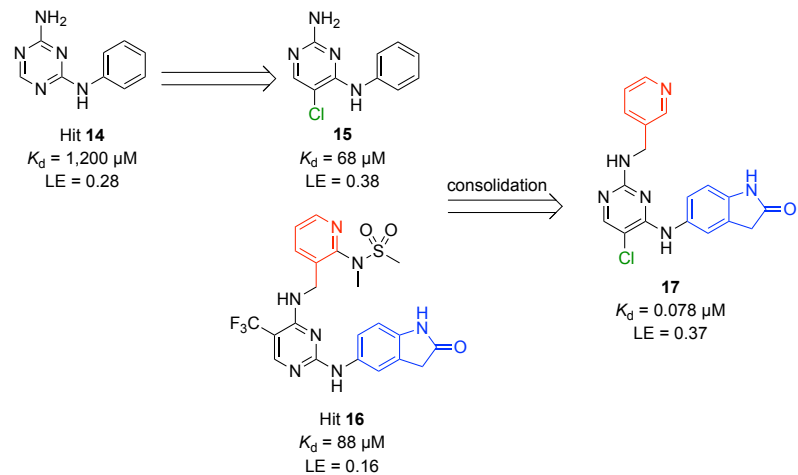


Figure 6. Binding mode of compound **11** to BCL6^{BTB} (PDB ID: 5X9P). (A) Surface view of the binding of inhibitor **11** (pink) to the BCL6^{BTB} LG. Compound **11** was shown in pink and BCL6^{BTB} monomers are shown in green and gray, respectively. (B) Detailed binding mode of inhibitor **11** in complex with BCL6^{BTB}.

Using fragment-based drug discovery (FBDD),⁴⁵⁻⁴⁹ Kamada and coauthors conducted screens of 1494 fragment-sized molecules against four BCL6^{BTB} proteins:⁵⁰ captured wt BCL6^{BTB}, captured mt BCL6^{BTB}, coupled wt BCL6^{BTB} and NeutrAvidin, using SPR.⁵¹⁻⁵⁴ From the identified hits, the authors chose 64 pan-active candidates and conducted full dose-responses. The binding of selected hits was confirmed by STD-NMR.⁵⁵ Based on these efforts, a triazine hit **14** (Scheme 3) was identified with a K_d value to 1,200 μM (SPR⁵⁶⁻⁵⁸) and ligand efficiency (LE⁵⁹) of 0.28 to BCL6^{BTB}.

Scheme 3. Identification of hit **14** and optimization to the generation of inhibitors **17**



Hit **14** binds to BCL6^{BTB} LG (PDB 5X4M) with a similar binding mode to that of inhibitor **1**. The linker NH group formed a key H-bond with the backbone carbonyl O atom of M51. Next, the authors tested the K_d values of four additional pyrimidine analogs of triazine **14**, and found the chloropyrimidine analog **15** with a significantly improved potency ($K_d = 68 \mu\text{M}$) and LE (0.38).

Parallelly, another hit **16** ($K_d = 88 \mu\text{M}$, LE = 0.16) was obtained from an ELISA-based HTS campaign by measuring the PPI between the BCL6^{BTB} dimer and BCOR peptide. The co-crystals revealed that inhibitors **15** and **16** bind to BCL6^{BTB} dimer with a similar orientation. The NH linkers bridging the two aromatic rings overlapped. The Cl atom on the pyrimidine ring in compound **15** fit into a small hydrophobic pocket formed by residues M51, L'25, and A52) (PDB

5X4N). The carbonyl O atom of the cyclic amide in hit **16** occupied the linker site and interacts with the backbone NH group of residue E115 (PDB 5X4O).

The structural similarities of compounds **15** and **16** inspired the authors to develop a hybrid compound **17**, which showed significantly improved potency ($K_d = 78$ nM, >15,000-fold more potent than hit **14**). Co-crystal structure of inhibitor **17** in complex with BCL6^{BTB} (Figure 7) showed that its pyridine ring was solvent exposed and mostly disordered. The two key H-bonds remained between the bridging NH group and the backbone carbonyl oxygen atom of M51 (2.8 Å), and between the cyclic amide oxygen with the backbone NH of residue E115 (3.0 Å). Interestingly, one molecule of ethylene glycol was observed in the HDCH site. One of the OH groups of the ethylene glycol formed an H-bond with the sidechain of H'14, while the other OH located 3.6 Å away from the cyclic amide fragment of compound **17**. These results highlighted the potential of further expansion of this inhibitor series toward the HDCH site.

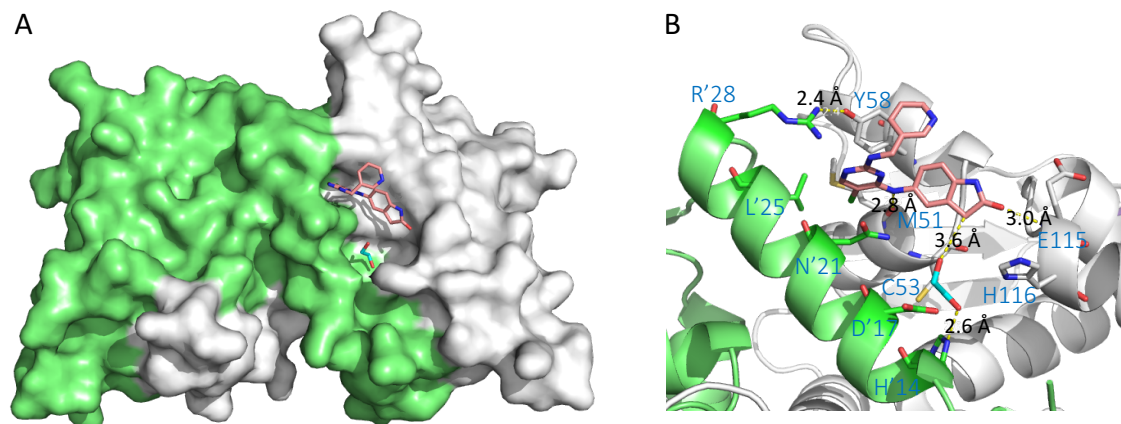
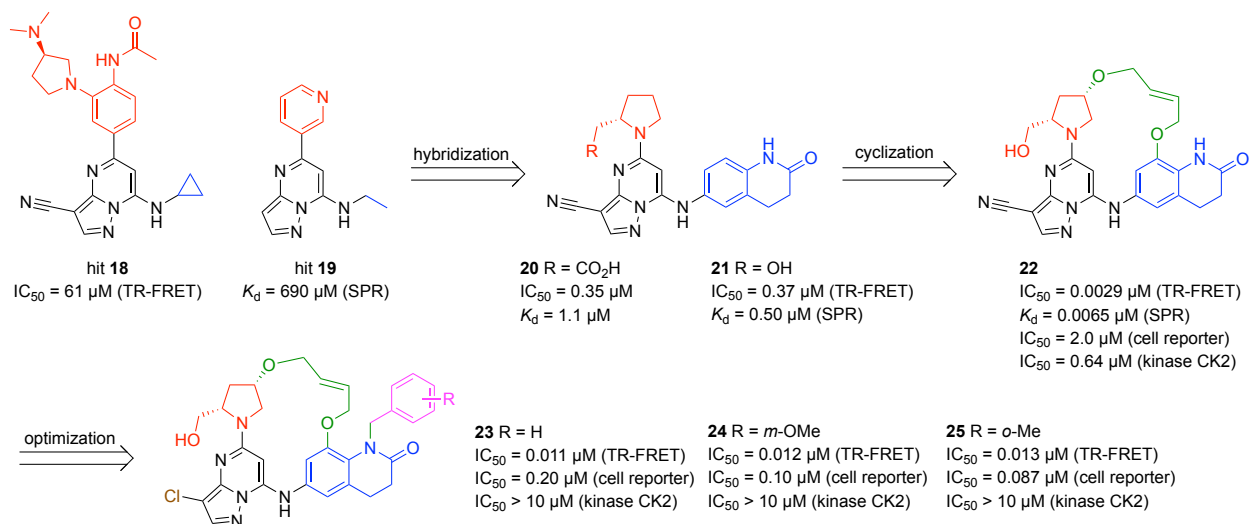


Figure 7. Binding mode of compound **17** to BCL6^{BTB} (PDB ID: 5X4Q). (A) Surface view of the binding of inhibitor **17** (pink) to the BCL6^{BTB} LG. Compound **17** was shown in pink and BCL6^{BTB} monomers are shown in green and gray, respective. (B) Detailed binding mode of inhibitor **17** in complex with BCL6^{BTB}.

Based on the encouraging binding affinity and ligand efficiency of inhibitor **17**, the authors further evaluated the activity of this compound in additional assays. In a competitive binding ELISA assay, compound **17** potently inhibited the PPI between BCL6^{B_TB} and BCOR peptide with the IC₅₀ value of 0.48 μ M. However, the cellular activity of this inhibitor was moderate (IC₅₀ = 8.6 μ M in M2H assay). Overall, compound **17** represented a promising BCL6^{B_TB} inhibitor for future development.

Using a time-resolved fluorescence resonance energy transfer (TR-FRET) assay, McCoull and coauthors screened a library of ~8,000 compounds from the AstraZeneca collection, and identified a pyrazolo[1,5-*a*]pyrimidine-based hit **18** with the IC₅₀ value of 61 μ M (Scheme 4).⁶⁰ Parallelly, the authors also used SPR to screen ~3,500 fragments and identified another hit **19** employing a same pyrazolo[1,5-*a*]pyrimidine core with the K_d value of 690 μ M. The binding of hits **18** and **19** to BCL6^{B_TB} was confirmed using 2D ¹H, ¹⁵N-TROSY NMR experiments in which, competitive displacement of the SMRT peptide by testing compounds was studied.

Scheme 4. Optimization strategy of hits **18** and **19** and generation of pyrazolo[1,5-*a*]pyrimidine inhibitors using TR-FRET, SPR, and cellular gene reporter assays



Based on the co-crystal structures of **18** (PDB 5N20) and **19** (PDB 5N1X) in complex with BCL6^{BTB}, the authors found that the pyrazolo[1,5-*a*]pyrimidine of both hits bind to the aromatic site of BCL6^{BTB} LG. The acid site residue R'24 formed a cation- π interaction with the pyridine ring of hit **19** while the disubstituted phenyl group of hit **18** extruded from the protein surface. Next, the authors conducted a solvent analysis (CCG, MOE) using 3D-RISM⁶¹, and identified a number of unstable H₂O molecules that are present in the linker site. Specifically, one of the H₂O molecules, which indicated high occupancy and low stability, interacted with the backbone NH of residue E115 through an H-bond. Then, a new analog **20** was synthesized employing a bicyclic lactam group designed to replace the E115-interacting H₂O molecule. Indeed, the co-crystal structure of inhibitor **20** (PDB 5N21) in complex with BCL6^{BTB} confirmed that the bicyclic lactam group occupied the linker site with the lactam carbonyl interacting with the E115 backbone NH via an H-bond. Interestingly, the carboxylate of inhibitor **20** did not interact directly to R'24. Instead, these two groups interacted through a structural H₂O molecule. As a confirmation of this result, the hydroxy analog **21** also indicated an excellent potency (IC₅₀ = 370 nM). Based on the structural data, a macrocyclic compound **22** was synthesized by tethering the two ring systems of compound **21** (Scheme 4). Remarkably, the new macrocyclic inhibitor **22** indicated an IC₅₀ value of 2.9 nM, over 120-fold more potent than its parent **21**. To explain the increase of affinity from acyclic inhibitor **21** to macrocycle **22**, the authors conducted conformational analysis using NMR in solution.^{62,63} Two major conformations were observed for both compounds, with an active/inactive ratio of 79:18 for macrocycle **22** and 60:40 for acyclic **21**, indicating that the flexible rings limited the inhibitor from adopting the bioactive configuration.

The authors further studied the binding kinetics of inhibitors **20-22** using SPR,⁶⁴⁻⁶⁶ and found that the K_d values correlated well with the IC₅₀ values obtained from the TR-FRET ($R^2 =$

0.99), such as compounds **20** (SPR $K_d = 1.1 \mu\text{M}$ vs. TR-FRET $\text{IC}_{50} = 0.35 \mu\text{M}$), **21** (SPR $K_d = 0.50 \mu\text{M}$ vs. TR-FRET $\text{IC}_{50} = 0.37 \mu\text{M}$), and **22** (SPR $K_d = 6.5 \text{ nM}$ vs. TR-FRET $\text{IC}_{50} = 2.9 \text{ nM}$). The authors then reasoned the affinity increase from acyclic **21** to macrocycle **22** as a combination of an increased associate rate constant k_{on} due to enhanced structural rigidity, and decreased dissociate rate constant k_{off} achieved from additional H-bond interactions due to the favorable orientation of the (5S)-hydroxymethyl group.

A co-crystal structure of macrocycle **22** (PDB 5N1Z) in complex with BCL6^{BTB} dimer was solved, in which, compound **22** occupied both the aromatic and linker sites similarly to that of acyclic analog **20** (Figure 8). The hydroxy group formed an H-bond to R'28 in the acid site. The linker region of the macrocycle was solvent-exposing and formed non-polar interactions with the protein. The NH group bridging the two ring systems formed a key H-bond (2.8 Å) to the backbone carbonyl O atom of M51. The bicyclic lactam of compound **22** occupied the linker site, with the lactam carbonyl O atom forming an H-bond (3.4 Å) to the side chain of E115. As the HDCH site was not occupied by the inhibitor, a chloride ion was detected in the co-crystal structure.

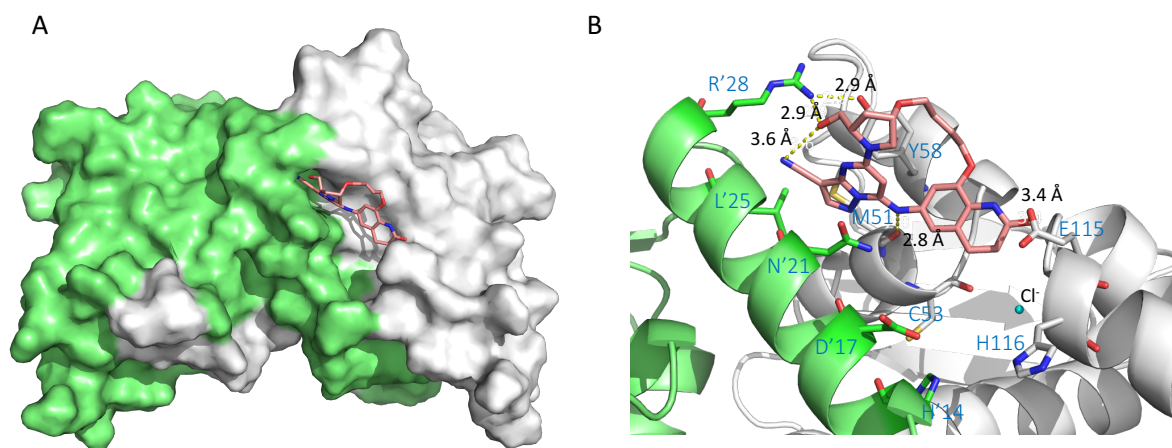


Figure 8. Binding mode of compound **22** to BCL6^{BTB} (PDB ID: 5N1Z). (A) Surface view of the binding of inhibitor **22** to the BCL6^{BTB} LG. Compound **22** was shown in pink and BCL6^{BTB}

monomers are shown in green and gray, respective. (B) Detailed binding mode of inhibitor **22** in complex with BCL6^{BTB}.

Considering macrocyclic inhibitor **22** originated from a kinase CK2-targeting scaffold,⁶⁷ a kinase screen was performed employing a panel of 126 kinases. The results indicated that across the panel only CK2 indicated > 60% inhibition at a relatively high concentration (1 μ M). More detailed evaluation revealed that compound **22** inhibited CK2 with an IC₅₀ value of 0.64 μ M, which was 220-fold weaker compared to that for BCL6^{BTB} (IC₅₀ = 2.9 nM) and 50-fold weaker compared to that of the original hit **18** (CK2 IC₅₀ = 12 nM). Interestingly, co-crystal structures of compound **22** bound to these two proteins (BCL6^{BTB}, PDB 5N1Z; CK2, PDB 5N1V) showed highly similar binding modes: the diarylamino group formed a key H-bond to both proteins. While in CK2, the NH group interacted with the kinase hinge domain via residue V116, in BCL6, it formed an H-bond to the carbonyl group of M51 backbone. The binding pockets of BCL6^{BTB} dimer and CK2 provided similar interactions and compound **22** showed similar bound conformations in both proteins, underlining the potential challenge in achieving selectivity for BCL6^{BTB} over CK2.

To further evaluate the selectivity of the macrocyclic inhibitors, compounds were parallelly tested in cellular gene reporter assays employing either BCL6^{BTB} or another BTB domain PLZF. All tested compounds indicated no inhibitory activity against PLZF at the concentration of 30 μ M (IC₅₀ range against BCL6^{BTB}: 0.1-6.1 μ M). One of the active macrocycle **23** was further screened (1 μ M) against a panel of 398 kinases and over 30 diverse pharmacological targets.⁶⁸ Inhibitor **23** indicated high selectivity⁶⁸ by hitting only two targets (GABA receptor, K_i = 0.25 μ M; 5-HT1B receptor, EC₅₀ = 0.57 μ M).

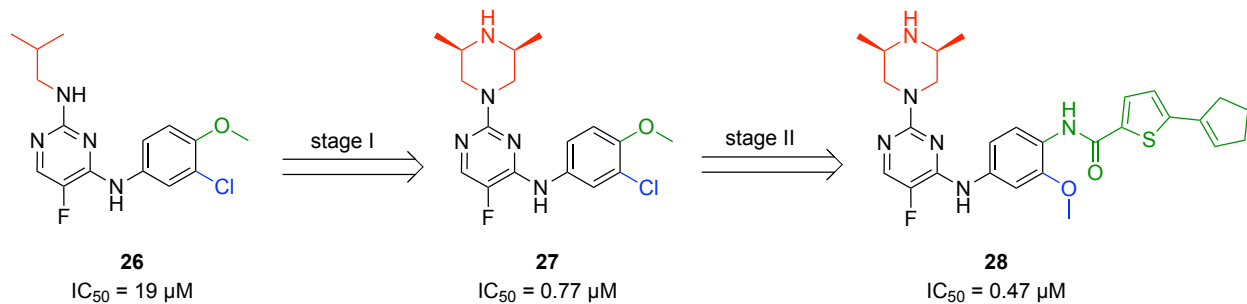
Finally, to examine the anticancer activity of the macrocyclic inhibitors, the authors performed cell proliferation assays using a panel of BCL6-dependent⁷ (e.g., OCI-Ly1, SUDHL-4,

OCI-Ly3, SUDHL-2, U2932, OCI-Ly10, TMD8) and BCL6-independent⁷ (e.g., Karpas422, OCI-Ly19) DLBCL cell lines, along with a multiple myeloma (MM) cell line AMO-1 with undetectable levels of BCL6. Note that Karpas422 has also been reported by Kerres and coworkers as a BCL6-dependent cell line.⁶⁹ Unfortunately for all macrocyclic inhibitors and the control compound FX1, the authors only observed no distinguishable features with low micromolar antiproliferative activity for all tested cell lines. While the authors didn't expand the discussion on these results, the lack of the antiproliferative effects of these macrocycles might due to the evaluation platform used in the study. Prior to the antiproliferation studies, the cellular activity of the macrocycles was only evaluated using a luciferase reporter assay employing engineered BCL6^{BTB} and a corepressor peptide but not the corresponding full-length proteins. Thus, one cannot exclude the possibility that the compounds could not inhibit the PPI between full length BCL6 protein and its corepressors. It is also worth noting that although good correlation between the results of TR-FRET and SPR were obtained, the authors didn't observe reasonable correlation between cell-free assays and cellular reporter assays. Furthermore, in the antiproliferation study, only a short 3-day proliferation study was pursued, which might not provide sufficient time for tested compounds to show BCL6-mediated effects.

Guo and coauthors screened an in-house collection of 230 compounds using a TR-FRET assay at three concentrations (100, 50, and 25 μ M) and identified a diaminopyrimidine compound **26** that inhibited the PPI between BCL6^{BTB} and SMRT with an IC₅₀ value of 19 μ M (Scheme 5).⁷⁰ Structural optimization of hit **26** focused on the amino-substitution of the diaminopyrimidine core and the *meta*- and *para*-substitutions of the phenyl ring. In the first stage, the isobutyl group of hit **26** was replaced by 14 different functional groups, which led to the generation of the (2*S*,6*R*)-2,6-dimethylpiperazinyl **27** with an IC₅₀ value of 0.77 μ M. Next, based on the structure of compound

27, the substitutions of the phenyl ring were investigated by 18 additional analogs to yield compound **28** with an IC_{50} value of $0.47 \mu\text{M}$. Overall, the SAR study employed a total number of 32 new compounds with the IC_{50} range of $>10 \mu\text{M}$ to $0.47 \mu\text{M}$.

Scheme 5. Optimization strategy of HTS hit **26** and generation of diaminopyrimidine inhibitor **28**



Next, the activity of selected inhibitors in reactivating BCL6 target genes was evaluated in the BCL6-dependent SUDHL-4 cells using real-time quantitative PCR (RT-PCR). All tested compounds, at the concentration of $5 \mu\text{M}$, reactivated ATR, CD69 and CXCR4⁷¹ compared to the vehicle. Then, compound **28** was chosen for further evaluation in two BCL6-dependent DLBCL cell lines SUDHL-4 and Farage. At the concentration of $5 \mu\text{M}$, compound **28** caused more significant reactivation effects of BCL6 target genes (p53, ATR, CD69, CXCR4, and CDKN1A)⁷¹ compared to those induced by positive controls **17** ($5 \mu\text{M}$, Takeda)⁵⁰ and **2** (FX1, $50 \mu\text{M}$).⁷

To explore the selectivity of inhibitor **28**, the authors followed the mRNA levels of SUDHL-4 and Farage along with a BCL6-independent DLBCL cell line Toledo. The results indicated that treatment of **28** caused a dose-dependent increase in the mRNA levels of six tested BCL6 target genes (p53, ATR, CD69, CXCR4, CDKN1A, and CD80) in both BCL6 dependent lines.⁷¹ Specifically, the mRNA level of CXCR4 in SUDHL-4 cells was ~ 12 times greater with exposure to compound **28** compared to the control **2**. Interestingly, compound **28** had no obvious effect on tested genes in the BCL6-independent Toledo cells.

The kinetics of inhibitor **28** binding to BCL6^{BTB} was studied using bio-layer interferometry (BLI). Compound **28** binds to BCL6^{BTB} with a K_d value of 0.37 μM , 20-fold and 3-fold greater than those of controls **2** (K_d 7.9 μM) and **17** (K_d 1.2 μM). To study its antiproliferative effects, compound **28** was tested in four BCL6 dependent DLBCL cell lines (SUDHL-4, Farage, DOHH-2, and OCI-Ly7) and a BCL6 independent Toledo cell line using the Cell Counting Kit 8 (CCK8) assay.⁷ Compound **28** killed all four BCL6 dependent cell lines with IC_{50} values \sim 1 μM , which was $>$ 15-fold more potent than positive controls **2** and **17**. On the other hand, compound **28** indicated an IC_{50} value of 3.3 μM against Toledo cells, a BCL6 independent GCB-DLBCL cell line that is less sensitive than the tested GCB-DLBCL cell lines. While the results are encouraging, the observed antiproliferative effects of compound **28** was in disagreement with other reports where significant antiproliferation could be obtained after a long study usually ranging from 14-17 days. Subsequently, selectivity of **28** over human normal cell lines (L02, HAF, NCM460, and PNT1A) were determined, and the IC_{50} values of compound **28** for normal cells were \sim 20-fold higher than that of DLBCL cells, confirming that compound **28** was selective against DLBCL cells.

The inhibitory effects of compound **28** in SUDHL-4 cell were evaluated using the EdU flow cytometry assay.⁷ Compound **28** significantly inhibited DLBCL cell growth. Treatment of compound **28**, at the concentrations of 0.625 μM , 1.25 μM , and 2.50 μM significantly decreased the number of living cells (from 48%, to 27%, 13%, and 6.6%, respectively) in 48 h. The results were more dramatic than those of controls **2** and **17**. The effect of compound **28** on DLBCL apoptosis was studied using flow cytometry, and the results revealed that compound **28** dose-dependently induced apoptosis (from 4.0%, to 6.8%, 16%, 73%), while at the same concentration, control compounds **2** and **17** indicated no obvious activity in inducing apoptosis.

To assess whether compound **28** could diminish the number of GC B cells, C57BL/6 mice were immunized with 4-hydroxy-3-nitrophenyl)acetyl-chicken gamma globulin (NP-CGG) and then treated with the compound (i.v., 10 mg/kg/3d for 12 days, vehicle: 20% (2-hydroxypropyl)- β -cyclodextrin). Flow cytometry was conducted to detect splenic GC B cells. Compared to the untreated group (frequency of GC B cells, ~2.0%), the treated group indicated a significantly decreased GC formation (frequency = 0.45%). The group treated with control **2** (50 mg/kg, i.p. daily injection) indicated a frequency of 0.73%. As BCL6 was a key regulator for the differentiation and maturation of CD4+ Tfh cell, the author further studied the development of Tfh cells upon treatment of compound **28** using flow cytometry,⁷² and found that the proportion of Tfh cells was significantly lower in the **28** treated group (0.99%) compared to in the control group (2.3%). Moreover, using immunofluorescent staining, the author also studied the splenic architecture, and found that both size and number of GCs were significantly decreased (~80%) upon treatment of compound **28**, which was consistent with the results of the flow cytometry analysis. It was noted that control **2** indicated modest inhibitory effects (~50%) in GC B cell development and GC formation.

Next, the authors used ELISA to determine the effects of compound **28** in impairing immunoglobulin affinity maturation.³² Compared to the controls, mice treated with compound **28** indicated ~50-fold lower titers of high-affinity immunoglobulin G1 measured by NP5-BSA. A similar decrease was also observed in total NP-specific immunoglobulin G1 production determined by NP23-BSA.

To evaluate the *in vivo* efficacy of compound **28**, an SCID DLBCL xenograft model with SUDHL-4 cells was used.⁷ At the dose level of 5 mg/kg/3d (i.v., vehicle: 20% (2-hydroxypropyl)- β -cyclodextrin), compound **28** suppressed the tumor weight (~70%) and volume (~60%) compared

to the vehicle-control. Treatment of compound **28** caused no apparent adverse events as little difference was observed in body weight and no obvious damage was detected to major organs (heart, liver, spleen, lung and kidney) determined by H&E staining. In addition, the Ki67 staining was used to assess tumor aggressiveness, and the results indicated that the expression level of Ki67 was decreased by half in **28** treated mice, confirming the *in vivo* effects of compound **28** in inducing DLBCL cell growth arrest. Moreover, using RT-PCR, the authors also showed that compared to controls, the mRNA levels of the BCL6 target genes CD69, CXCR4 and ATR were increased by >150%, >60% and >200%, respectively when compound **28** was administered (5 mg/kg/3d).

Finally, the *in vitro* ADME profile of compound **28** was assessed. The results indicated that this lead compound had low microsomal clearance CL_{int} ($CL_{int(mic)} < 9.6 \mu\text{l}/\text{min}/\text{mg}$, $CL_{int(liver)} < 38.0 \text{ ml}/\text{min}/\text{kg}$), acceptable metabolic stabilities ($t_{1/2} > 60 \text{ min}$) in human liver microsomes, and modest binding to plasma proteins (7.6% free fraction).

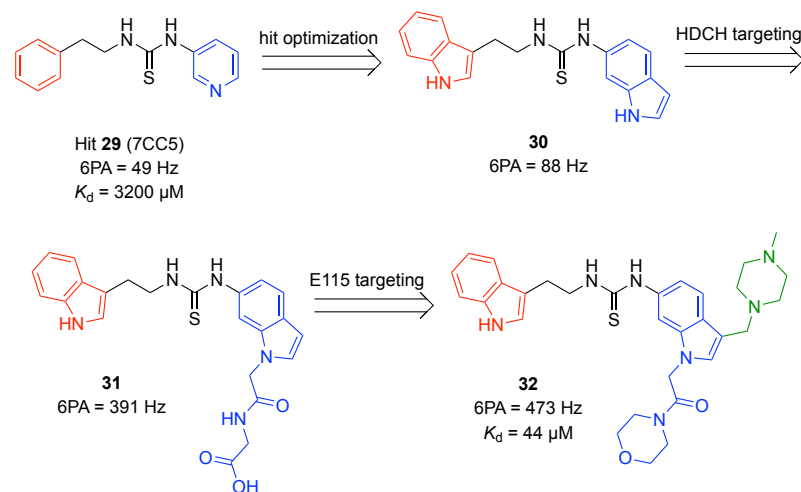
Cheng and coworkers reported a series of inhibitors for the BCL6^{BTB} dimer based on a novel thiourea scaffold.⁷³ Screening of a library of ~1,500 fragment-sized molecules by protein-observed NMR spectroscopy led to the identification of a thiourea hit **29** with a K_d value 3.2 mM (Scheme 6). The co-crystal structure of BCL6^{BTB}-**29** (PDB 6C3N) revealed that hit **29** binds in the aromatic site of the BCL6^{BTB} LG. The S atom of this hit fit into a hydrophobic pocket formed by N'21, L'25, M51, A52, and Y58. One of the thioamide hydrogens formed an H-bond (2.6 Å) with the backbone carbonyl O atom of M51.

Based on this result, new inhibitors were designed using a computer-aided drug design (CADD) method called site identification by ligand competitive saturation (SILCS).⁷⁴⁻⁷⁷

Optimization of hit **29** involved replacing the 3-substituted pyridine with an indole ring and expanding the chemical structure of inhibitors into the HDCH site via the indole nitrogen. Moreover, a piperazine group was introduced to the indole group to interact with the acidic sidechain of residue E115. Finally, the phenethyl group of hit **29** was replaced by a second indole ring.

The authors characterized the binding affinity of early weak inhibitors using ^1H - ^{15}N HSQC spectra NMR. Specifically, the sum of chemical shift perturbations of six selected amide (T62, T48, F61, N23, R28, and V18) resonances (6PA) were used to rank the relative affinity of new compounds to BCL6^{BTB} . Compared to hit **29**, the di-indole analog **30** showed an increased 6PA value (88 Hz vs 49 Hz for hit **29**). Substitution of the right-arm indole nitrogen of compound **30** using eight additional analogs yielded compound **31** with a significantly increased 6PA value of 391 Hz. Next, a methyl piperazine was introduced into the right-arm indole, followed by a further investigation of substitutions on the right-arm indole nitrogen by nine additional analogs. These efforts resulted in compound **32** with a 6PA value of 473 Hz. Overall, the SAR employed a total number of 19 new compounds with the 6PA range of 49 Hz to 473 Hz.

Scheme 6. Identification and optimization of thiourea-based inhibitors of BCL6^{BTB} .



Using NMR-titration experiments, the K_d value of inhibitor **32** to BCL6^{BTB} dimer was determined to be 44 μM , which was 70-fold more potent than hit **29** ($K_d = 3,200 \mu\text{M}$). Isothermal-titration calorimetry (ITC) was also performed to validate the binding of **32** to BCL6^{BTB} dimer in which a very similar affinity ($K_d = 36 \mu\text{M}$) was reported. The crystal structure of the BCL6^{BTB}-**32** complex (Figure 9) showed **32** occupying the aromatic, linker and HDCH sites of the BCL6^{BTB} LG. Specifically, the NH group of the thiourea functionality formed an H-bond to M51 (2.8 \AA). In addition, the tertiary amino group of the piperazine ring interacted to the carboxylate sidechain of E115 via an electrostatic interaction (3.3 \AA). Moreover, the morpholine group of amide tail binds into the HDCH site formed by the backbones of A52 and C53, and side chains of H'14, A'17, V'18 and N'21.

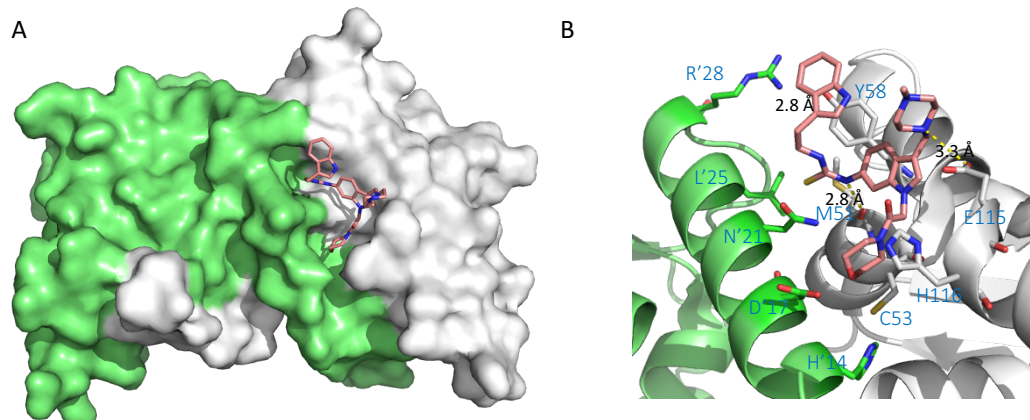


Figure 9. Binding mode of compound **32** to BCL6^{BTB} (PDB ID: 6C3L). (A) Surface view of the binding of inhibitor **32** (pink) to the BCL6^{BTB} LG. Compound **32** was shown in pink and BCL6^{BTB} monomers are shown in green and gray, respective. (B) Detailed binding mode of inhibitor **32** in complex with BCL6^{BTB}.

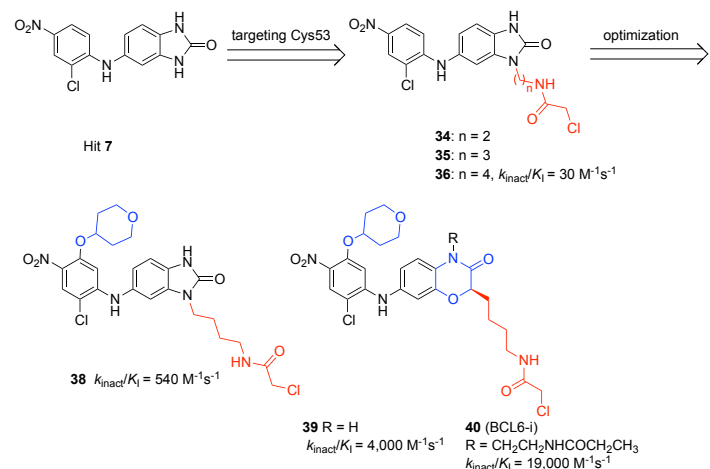
To test the potencies of compounds, an AlphaLisa assay was established involving labeled BCOR peptide binding to BCL6^{BTB} dimer. As the most potent candidate, compound **32** showed

an IC_{50} value of 27 μ M. Next, differential cell-killing assays were conducted for selected inhibitors using a panel of four BCL6-dependent lymphoma cell lines (OCI-Ly1, OCI-Ly7, SUDHL-4, and SUDHL-6) and three BCL6-independent lymphoma cell lines (Karpas422, Toledo, and OCI-Ly1-B50). Note that Karpas422 has reported by Kerres and coworkers as a BCL6-dependent cell line.⁶⁹ Unfortunately, the tested inhibitors only showed relatively weak cellular activity likely due to limited cell permeability.

Irreversible $BCL6^{BTB}$ Inhibitors

By studying the binding mode of the HTS hit **7**⁴² to the $BCL6^{BTB}$ protein, Sameshima and coauthors learned that compound **7** binds into the HDCH site in proximity to the nucleophilic residue C53 (Scheme 7). Based on this observation, the authors synthesized chloroacetamides **34**-**36** as potential C53-targeting irreversible inhibitors for $BCL6^{BTB}$.⁷⁸ The covalent bond formation between $BCL6^{BTB}$ and these chloroacetamide analogs were assessed by mass spectrometry, and the results confirmed a 1:1 covalent modification by chloroacetamide **36** and its homologated analogs **34** and **35**. Interestingly, the acrylamide analogs of compound **36** failed to modify $BCL6^{BTB}$.

Scheme 7. Design of irreversible $BCL6$ inhibitors by targeting C53 residue from compound **7**



Next, TR-FRET assays were used to evaluate the inhibitory activity of compounds **34**–**36** for the PPI between BCL6^{BTB} dimer and the TAMRA peptide, a fluorescent tetramethylrhodamine-conjugated BCL6^{BTB} binding sequence (TAMRA-Abu(4)-VWYTDI MRDWM-OH) derived from peptide **4**.³⁸ Although none of the inhibitors gave a full dose response curve, the results confirmed that compound **36** was more potent than its homologated analog **35** at the concentration of 6.3 μ M, while the other analog **34** indicated no inhibitory activity at the same concentration. To identify the specific BCL6^{BTB} amino acid residue modified by inhibitor **36**, the authors conducted MS studies in the presence of compound **5** (50 μ M),³⁸ a highly potent peptide BCL6^{BTB} inhibitor covering the C53-containing HDCH site. The results indicated that compound **36**, at concentrations as high as 20 μ M, failed to modify BCL6^{BTB}, implying that inhibitor **36** likely achieved his inhibitory activity by modifying C53 of BCL6^{BTB}. Next, irreversible modification of BCL6^{BTB} by inhibitor **36** was confirmed using jump dilution assays in which compound **36** (6.3 μ M) or its non-covalent analog (100 μ M) was incubated with BCL6^{BTB} for 10 h, followed by a 100-fold dilution using a buffer containing excess TAMRA peptide (20-fold K_d). Different to the noncovalent analog whose inhibitory activity disappeared, the inhibitory effect of compound **36** maintained in 24 h after dilution. These results again, confirmed irreversible inhibitory mechanism of compound **36**.

Next, optimization of compound **36** was pursued by following the k_{inact}/K_I value of new inhibitors in an irreversible probe competition (IPC) assay using a covalent fluorescent probe **37** (BCL6-FP) (Figure 10).^{79,80} The k_{inact}/K_I value of inhibitor **36** was determined to be $3.0 \times 10^1 \text{ M}^{-1} \text{ s}^{-1}$. Introduction of a 4-tetrahydropyranloxy group to compound **36** resulted in a new inhibitor **38** with an 18-fold increased k_{inact}/K_I value ($5.4 \times 10^2 \text{ M}^{-1} \text{ s}^{-1}$). Expansion of the five-membered benzo[*d*]imidazol-2-one ring to a six-membered benzoxazine ring yielded compound **39**, which

indicated an another 7.4-fold improvement of $k_{\text{inact}}/K_{\text{I}}$ value ($4.0 \times 10^3 \text{ M}^{-1} \text{ s}^{-1}$). Finally, introduction of a 2-propanoylaminoethyl group to the chemical scaffold gave inhibitor **40** (BCL6-i) that was highlighted by an excellent $k_{\text{inact}}/K_{\text{I}}$ value of $1.9 \times 10^4 \text{ M}^{-1} \text{ s}^{-1}$, over 670-fold higher than that of the parent **36**.

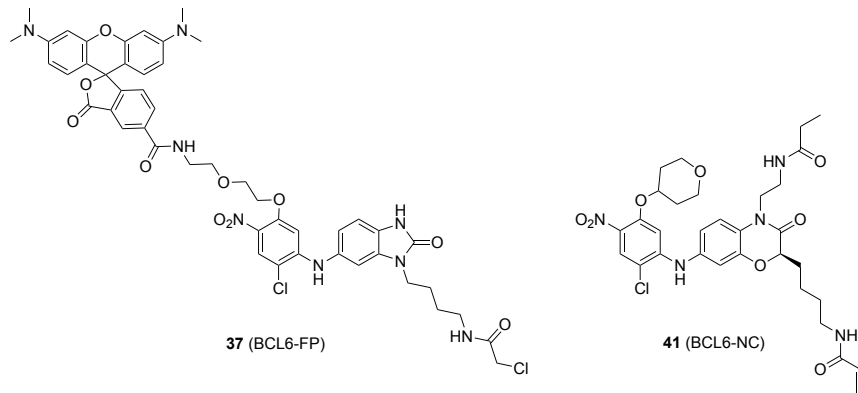


Figure 10. Chemical structures of **37** (BCL6-FP) and **41** (BCL6-NC).

MS analysis confirmed the covalent binding of inhibitor **40** to BCL6^{BTB} with 1:1 stoichiometry. In addition, compound **40** indicated identical IC₅₀ values after several hours of incubation in the IPC assay, confirming the irreversible binding of the compound to BCL6^{BTB}. The authors further confirmed that inhibitor **40** covalently modified residue C53 in the HDCH site of BCL6^{BTB} via time-dependent inhibition experiments. Compound **40** effectively inhibited BCL6 mutants containing C53 time-dependently, while the inhibitory activity disappeared for BCL6 mutants without C53. These results together supported that the optimized electrophilic chloroacetamide inhibitor **40** modified C53 residue in the HDCH site of BCL6^{BTB} LG.

To address the potential toxicity of these irreversible inhibitors, the intrinsic chemical reactivity (k_{chem}) values^{81,82} of compound **40** and its analogs was compared to an acrylamide-based irreversible pan-erbB kinase inhibitor CI-1033 that is under Phase II clinical trials. Compound **40** showed a 23-fold smaller k_{chem} value ($1.6 \times 10^{-3} \text{ M}^{-1} \text{ s}^{-1}$) than that of compound CI-1033 ($3.7 \times$

10^{-2} M $^{-1}$ s $^{-1}$), indicating an acceptable k_{chem} value of compound **40** for further cellular or *in vivo* evaluations.

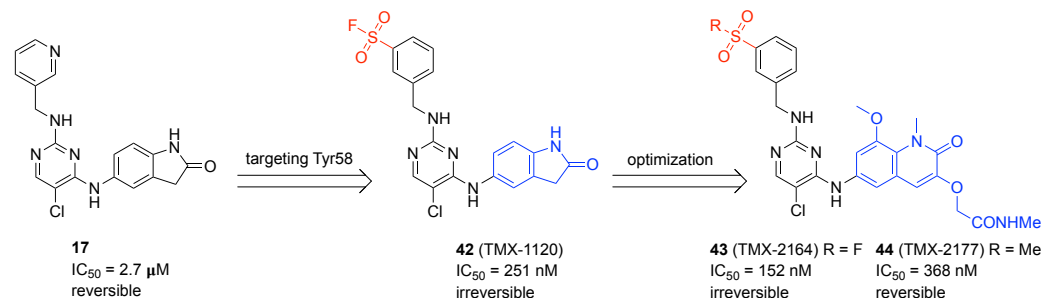
To assess the cellular activity of inhibitor **40**, the authors used a fluorescence imaging assay to follow colocalization of BCL6 with the BCL6-FP **37**. Cells expressing full length FLAG-BCL6 were treated with DMSO, **40** (1 μ M), or reversible control **41** (BCL6-NC, Figure 10) (1 μ M) for 1 h, then allowed to react with BCL6-FP **37** (1 μ M) for another 1 h. Colocalization of BCL6 protein with BCL6-FP **37** was detected, indicating that BCL6-FP **37** covalently interacts with the BCL6 protein in cellular environment. On the other hand, pre-treatment of inhibitor **40** prohibited the fluorescence signal from BCL6-FP **37**, whereas the treatment of a reversible compound **41** did not. These results together confirmed that inhibitor **40** irreversibly modifies the protein BCL6 in cell. Next, using an M2H assay, the authors confirmed that compound **40** inhibited the PPI between intracellular BCL6 and BCOR. Specifically, after a 24 h treatment, compound **40** (1 μ M) inhibited the M2H signal by \sim 80%, whereas the negative control compound **41** (1 μ M) did not (< 10% inhibition).

Finally, the authors tested the antiproliferation activity of inhibitor **40** and the negative control **41** against two BCL6-dependent GCB-DLBCL cell lines SUDHL-4 and OCI-Ly3. The results revealed that treatment of inhibitor **40**, at the concentration of 3 μ M after 72 h, significantly suppressed growth of SUDHL-4 (\sim 80%) and OCI-Ly3 (\sim 70%) compared to negative control **41** (\sim 15% and \sim 25%, respectively).

Teng and coauthors reported a series of irreversible BCL6^{BTB} inhibitors by targeting the side chain of amino acid residue Y58 in the aromatic site using a well-known sulfonyl fluoride functionality.⁸³ The initial inhibitor **42** (TMX-1120) was designed based on the chemical scaffold of Takeda compound **17**⁵⁰ (Scheme 8). Based on the co-crystal structure, the authors noticed that

the Y58-OH group and *meta*-carbon of the pyridine are located in reasonable distance (4.2 Å) for the insertion of an additional functional group. In addition, the guanidinium group of R'28 was in close proximity (2.4 Å) to Y58-OH, which could help deprotonation of the Y58-OH group. Furthermore, the solvent-exposed and disordered pyridine moiety suggested no steric hindrance to reach the Y58-OH. Therefore, the authors introduced a sulfonyl fluoride,⁸⁴⁻⁸⁷ a widely used tyrosine-targeting electrophilic warhead, to target Y58.

Scheme 8. Development of irreversible inhibitor 43 (TMX-2164) targeting Y58 of BCL6^{BTB}



The potency of new compounds was determined using a TR-FRET assay employing a bodipyFL-labeled BCOR peptide in 30 min. Compound **42** showed an IC₅₀ value of 251 nM, > 10-fold more potent than parent **17** (IC₅₀ = 2.7 μM). To verify the formation of a covalent bond of the inhibitor to Y58, the authors used protein mass spectrometry by analyzing the changes of molecular weights of the recombinant BCL6^{BTB} after incubating with 10 equivalent compound **42** for 2 h. A mass shift was observed corresponding to 1:1 stoichiometric modification of the protein by compound **42**. The site of covalent modification was further confirmed by digesting the labeled protein with trypsin followed by analyzing the resulting peptides using capillary electrophoresis-mass spectrometry (CE-MS).

Next, the authors synthesized new hybrid **43** (TMX-2164) by merging the sulfonyl fluoride warhead with the chemical scaffold of another BCL6 inhibitor **62** (BI-3812)⁶⁹. Compound **43**

displayed an IC_{50} value of 152 nM and its reversible sulfone analog **44** (TMX-2177) revealed a comparable IC_{50} of 368 nM. Covalent modification of BCL6 by inhibitor **43** was also confirmed by mass spectrometry. A mass shift of 572 Da, corresponding to the addition of compound **43** with the loss of an HF, was detected.

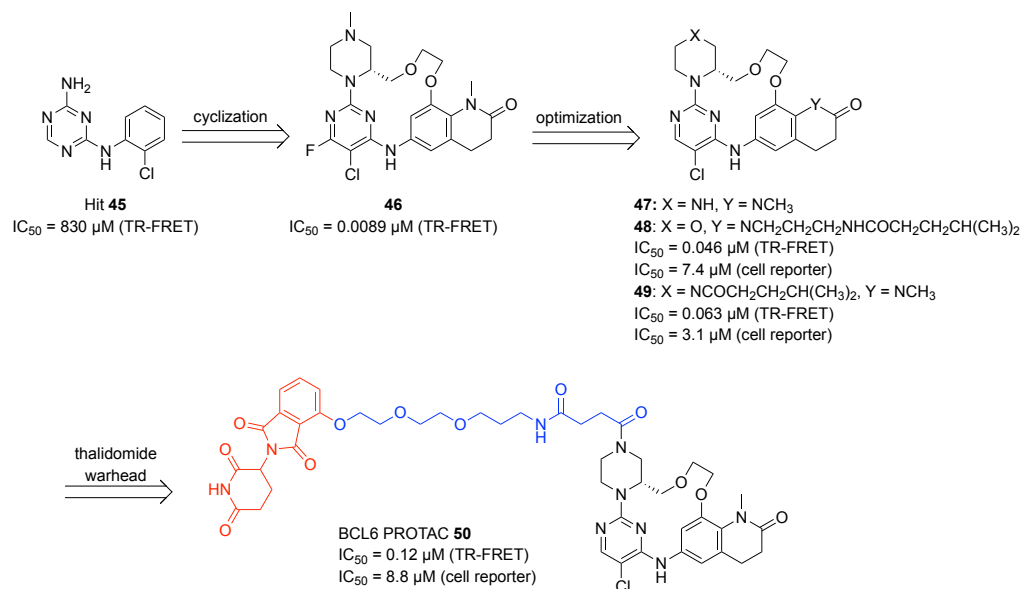
To examine the cellular activity of the compounds, the authors developed a fluorescence-activated cell sorting (FACS) assay that enabled quantification of BCL6 levels by following the ratio of eGFP/mCherry.⁸⁸ Cells were treated with irreversible inhibitors **42** and **43** or reversible controls **17** and **44** at 5 μ M for 30 h, followed by a BCL6 degrader **61** (BI-3802)⁶⁹ at different concentrations. While treatments of covalent inhibitors **42** and **43** protected BCL6 from **61**-induced degradation, treatment of reversible controls **17** and **44** did not indicate the same protection, demonstrating permanent modification of BCL6 protein by covalent inhibitors. Finally, the anti-DLBCL activity of new covalent inhibitors were evaluated using SUDHL-4 cells. After 5 days of treatment, the irreversible inhibitor **43**, compared to both reversible inhibitors **17** and **44**, indicated the most potent antiproliferation effect with the GI_{50} value in the single digit micromolar range.

BCL6 Protein Degraders

Based on the small molecule reversible inhibitors developed by the team in AstraZeneca (Scheme 4),⁶⁰ McCoull and coauthors reported their efforts in the development of cell permeable BCL6 inhibitors and a subsequent PROTAC,^{89,90} which effectively degraded the BCL6 protein.⁹¹ A triazine hit **45** (IC_{50} = 830 μ M, TR-FRET, Scheme 9) in this study was identified from a fragment screen and optimized by 2D-NMR-guided effective conformational analysis, to yield a macrocycle **46** with a remarkable ~93,000-fold improvement in inhibitory potency (IC_{50} = 8.9 nM). The co-

crystal structure of compound **45** in complex with BCL6^{BTB} homodimer (PDB 6EW6) confirmed the binding site of the ligand in the BCL6^{BTB} LG.

Scheme 9. Development of BCL6 PROTAC **50**.



The authors assessed BCL6 target engagement in OCI-Ly1 cells using a cellular thermal shift assay (CETSA),³⁴ in which, ligand binding could lead to increased BCL6 stability and therefore a higher protein unfolding temperature.³⁴ The EC₅₀ value of a testing ligand could be calculated based on the isothermal dose response curve. Using this assay, compound **46** was determined to have an EC₅₀ value of 0.48 μM. The authors further demonstrated a good correlation between the CETSA results and an additional cellular luciferase gene reporter assay established in HEK293T/17 cells, in which, the compound **46** indicated an IC₅₀ value of 0.71 μM. Using a similar gene reporter assay, the authors demonstrated the specificity of compound **46** to BCL6^{BTB} over another BTB domain protein PLZF (IC₅₀ >32 μM). Specificity of compound **46** was further evaluated via a kinase screen against a panel of 399 kinases, and the result revealed that compound **46** had no obvious inhibitory activity to the tested kinases (< 25% inhibition at 1 μM).

Analysis of the co-crystal structure of compound **47** (Figure 11), a close analog of compound **46**, indicated that vectors from both the lactam and piperazine N atoms were directed toward solvent accessible space that could allow PROTAC installation. Substitution at these two N atoms were subsequently pursued yielding compounds **48** and **49**, and the result indicated that compound **49** maintained higher cellular activity than compound **48** (IC_{50} 3.1 μ M vs. 7.4 μ M). Accordingly, compound **49** was attached to a thalidomide warhead,⁹²⁻⁹⁴ through a two-unit PEG hydroxypropylamine butanediamide linker, to generate the BCL6 PROTAC **50** (Scheme 9). Compared with parent compound **49**, PROTAC **50** displayed improved BCL6 inhibitory activity with purified protein (IC_{50} = 0.12 μ M, RT-FRET), although it only indicated modest inhibitory potency in cell (IC_{50} = 8.8 μ M, luciferase gene reporter assay). Besides, PROTAC **50** had some specificity for BCL6^{BTB} over PLZF (IC₅₀ > 32 μ M, luciferase gene reporter assay). However, PROTAC **50** indicated decreased lipophilicity ($\log D$ = 1.6 vs 4.2 for compound **49**) and Caco2 permeability (0.081×10^{-6} cm/s vs 31×10^{-6} cm/s for compound **49**).

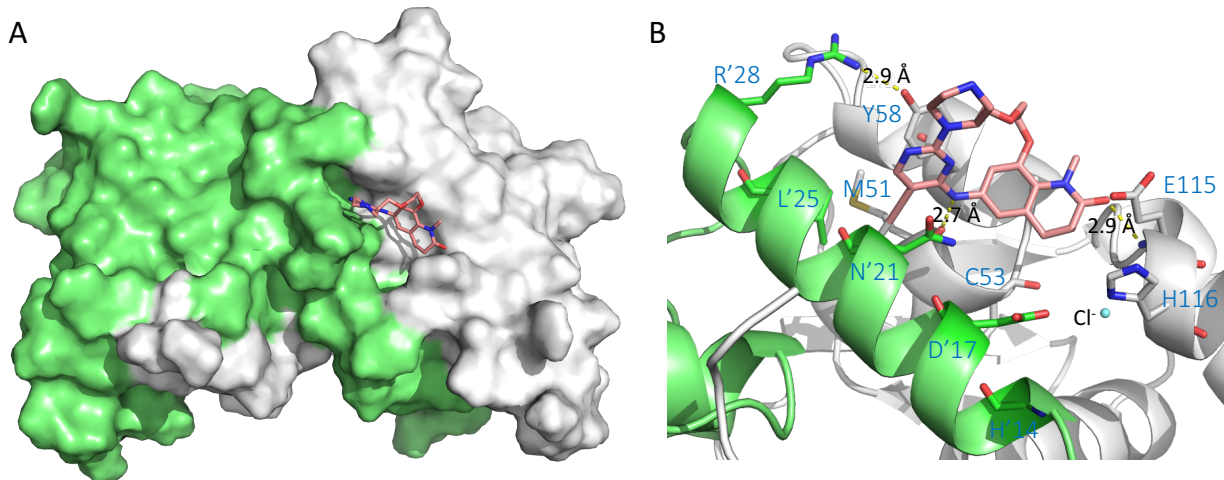


Figure 11. Binding mode of compound **47** to BCL6^{BTB} (PDB ID: 6EW8). (A) Surface view of the binding of inhibitor **47** (pink) to the BCL6^{BTB} LG. Compound **47** was shown in pink and

BCL6^{BTB} monomers are shown in green and gray, respective. (B) Detailed binding mode of inhibitor **47** in complex with BCL6^{BTB}.

The degradation ability of PROTAC **50** was evaluated by following the BCL6 protein levels after treatment with the compound in OCI-Ly1 cells. PROTAC **50** dose-dependently degraded BCL6 in OCI-Ly1 cells and the effect was specific to BCL6 as it caused no degradation on another protein involved in the BCL6-corepressor complex, TBLR1. It was noted however, the degradation obtained from the treatment of PROTAC **50** was not complete at the highest concentration tested (1 μ M, 82% degradation). It was also shown that significant BCL6 protein degradation was observed as early as 1 h and the effect lasted for at least 72 h. In contrast, upon treatment of reversible inhibitor **49** caused no obvious change in BCL6 levels over time, suggesting that the thalidomide group was responsible for BCL6 protein degradation.

To elucidate the mechanism of action of PROTAC **50**, three control experiments were performed by including: i) proteasome inhibitor MG132, ii) BCL6 inhibitor **49**, and iii) a potent thalidomide derivative pomalidomide to the tests. Reduced levels of degradation were observed for all three experiments. More complete protection from protein degradation was achieved when a combination of BCL6 inhibitor **49** and pomalidomide were added. Altogether, these results supported the mechanism of PROTAC-based BCL6 protein degradation. Furthermore, PROTAC **50** at the concentration of 1 μ M, also caused significant BCL6 degradation effects (59% - 84%) for other DLBCL cells SUDHL-4, Karpas422 and ULA, and Burkitt's lymphoma cell line Ramos.

To assess the antiproliferative activity of PROTAC **50**, a three-day cell proliferation study was performed using DLBCL cell lines of both GCB (OCI-Ly1, Karpas422, SUDHL-4, WILL-2 and OCI-Ly19) and ABC (OCI-Ly10, RI-1, SUDHL-2, OCI-Ly3 and TMD8) subtypes with varying levels of BCL6, along with the MM cell line AMO-1. Unfortunately, PROTAC **50** failed

to show significant selectivity with the similar low micromolar pGI₅₀ values for all tested cell lines. Furthermore, PROTAC **50** did not indicate improved antiproliferative potency compared to the BCL6 inhibitor **49** in BCL6-dependent DLBCL cells. Finally, a long 16-day proliferation study was pursued. However, similarly modest antiproliferative effects of PROTAC **50** were obtained for OCI-Ly1, SUDHL-4, and OCI-Ly19 cell lines.

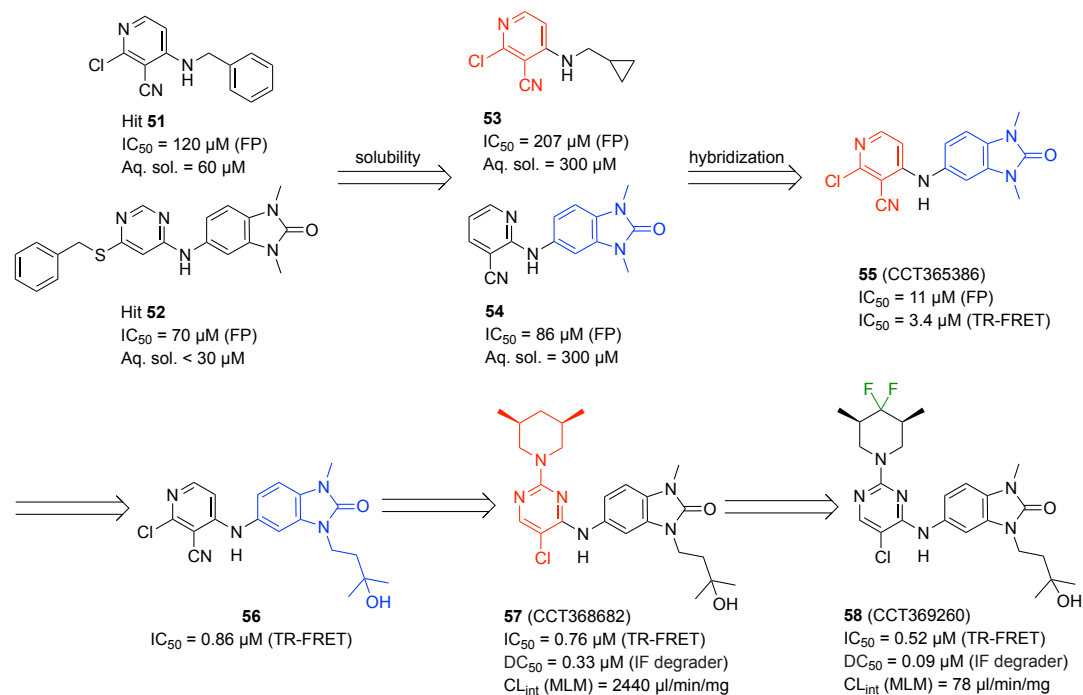
To explain the relatively weak antiproliferative effect of the compound, the authors followed the subcellular BCL6 levels after the treatment the PROTAC (10 μ M, 24 h) using an immunofluorescence (IF) staining assay in OCI-Ly1 cells. Although reduced levels of diffuse and punctate staining of BCL6 were detected, significant punctate staining of BCL6 remained. Considering the incomplete degradation of BCL6 by PROTAC **50**, quantitative mass spectrometry was used to estimate specific intracellular concentrations of compound **50**. After treatment with PROTAC **50**, the concentrations of PROTAC **50** in cytoplasm was 0.76 μ M, over 10-fold below the IC₅₀ (8.8 μ M) of the compound in BCL6 cell reporter assay. Importantly, the concentration of PROTAC **50** in the nucleus, where the target BCL6 locates, was only detected as 0.05 μ M. The extremely low levels of compound concentration in the nucleus might explain the relatively weak cellular potency of the PROTAC.

The authors further showed that after treatment of PROTAC **50**, BCL6 levels were similar in different subcellular compartments including cytoplasmic (DMSO:**50** = 7.3:2.3), soluble nuclear (DMSO:**50** = 86.0:20.2), and chromatin bound (DMSO:**50** = 6.7:1.9) fractions. Finally, the intracellular levels of BCL6 were measured after treatment of PROTAC **50** (1 μ M, 4 h). PROTAC **50** could sufficiently access the chromatin bound (1.9 μ M) and soluble nuclear (0.46 μ M) fractions, to cause BCL6 degradation within cytoplasm, nucleus, and chromatin bound fractions.

Bellenie and coworkers at the Institute of Cancer Research (ICR) reported benzimidazolone-based BCL6 inhibitors.⁹⁵ Interestingly a subset of compounds triggered rapid degradation of BCL6, of which, compound **58** (CCT369260) was highlighted as a probe unlike known PROTAC and non-PROTAC degraders (Scheme 10).

By screening an in-house compound library using a fluorescence polarization (FP) assay, the authors identified two hits **51** and **52** with the IC_{50} values of 120 μ M and 70 μ M, respectively (Scheme 10). To improve aqueous solubility, their analogs **53** and **54** were synthesized, which indicated \sim 5-fold increased solubility without compromising the inhibitory potencies. Next, hybridization of compounds **53** and **54** yielded the benzimidazolone **55** (CCT365386) with an IC_{50} value of 11 μ M. The co-crystal structure of compound **55** bound to BCL6^{BTB} (PDB: 6TOH) was determined. Results from additional assays revealed that compound **55** had a favorable drug-like profile including low microsomal clearance, high permeability, and good ligand efficiency.

Scheme 10. Optimization strategy of hits **51 and **52** and generation of benzimidazolone **58****



To enable the accurate measurement of inhibitors with nanomolar potency, the authors developed a TR-FRET assay with which the IC₅₀ value of inhibitor **55** was determined to be 3.4 μ M. Structural refinement of compound **55** involved the expansion of inhibitors into the HDCH site via substitution of the N³-position of the benzimidazolone, and optimization of the interactions to the aromatic site around residue Y58. The authors found that introduction of a 3-hydroxyl-*n*-butyl group at the N³-position of the benzimidazolone yielded an achiral tertiary alcohol **56** with a 5-fold improvement of inhibitory potency. The co-crystal structure of compound **56** bound to BCL6^{BTB} (PDB: 6TOJ) showed that the pyridine group formed hydrophobic interactions with aromatic sidechain of Y58, and the pyridine nitrogen formed an H-bond with R'28 through a water molecule. The authors also found that a pyrimidine ring demonstrated the same effects. Based on these results, the 2-position on the pyrimidine was further explored and demonstrated that substitution was tolerated. Next, modification of the substituted pyridine was investigated by 24 additional analogs, and the (3R,5S)-3,5-dimethylpiperidine compound **57** (CCT368682) was identified as a potent BCL6 degrader with an IC₅₀ value of 0.76 μ M and DC₅₀ value of 0.33 μ M (100%) in an immunofluorescence degrader assay. To further investigate the degradation ability of compound **57**, SUDHL-4 and OCI-Ly1 cells were treated with compound **57** at 37 °C for 4 h, and the protein levels of BCL6 was monitored by Western blots. Compound **57** reduced the protein levels of BCL6 dose dependently, with complete degradation at 10 μ M of the compound. This result was consistent with the mechanism of compound-mediated degradation of BCL6.

Finally, to improve stability and reduce metabolic clearance of the compound, substitutions on the pyrimidine ring were investigated. These efforts yielded the difluoromethylene degrader **58** (CCT369260), which indicated an IC₅₀ value of 0.52 μ M (TR-FRET) and excellent DC₅₀ value

of 0.09 μM in SUDHL4 cells. Further evaluation indicated that compound **58** had a CL_{int} mouse liver microsomes (MLM) value of 78 $\mu\text{l}/\text{min}/\text{mg}$.

To confirm the observed degradation was not cell line specific, the authors used meso scale discovery (MSD) assays to follow BCL6 protein degradation in different cells. Compound **58** caused >85% degradation of BCL6 in OCI-Ly1 ($\text{DC}_{50} = 0.049 \mu\text{M}$) and Karpas422 ($\text{DC}_{50} = 0.062 \mu\text{M}$) cells. Then, the antiproliferative activity of compound **58** and four analogs were examined using a 14-day proliferation assay in BCL6-dependent (SUDHL-4 and OCI-Ly1) and BCL6-independent (OCI-Ly3) cell lines. Compound **58** indicated differential cell killing effects for BCL6-dependent cells over the BCL6-independent OCI-Ly3 cells. Compound **58** was further evaluated in a panel of BCL6-independent cell lines (Toledo, OCI-AML3, PLB-985, and MM.1S) using a 17-day proliferation assay. The results indicated that compound **58** had no antiproliferative activity for these cell lines at the concentration of 1 μM .

PK studies of compound **58** were performed using female Balb/C mice (1 mg/kg, i.v.; or 5 mg/kg, p.o.). Compound **58** indicated a reasonably good profile with modest clearance ($\text{CL} = 20 \text{ mL min}^{-1} \text{ kg}^{-1}$), good oral bioavailability (54%), and high plasma protein binding (0.07% free in SCID mouse plasma, $n = 6$). To determine if sufficient exposure could be achieved in the SCID mouse used for xenograft studies, additional PK studies were conducted at three different dose levels (5, 15, and 50 mg/kg, p.o.). At the medium dose (15 mg/kg), the mean total blood concentration of compound **58** was above the desired concentration for $\sim 10 \text{ h}$.

Next, to determine if, and at what concentration BCL6 depletion could be achieved, a PK/PD study was conducted in mice using a single dose of compound **58** at 15 mg/kg (p.o.) in an OCI-Ly1 DLBCL xenograft model, with dosing of compound commencing 20 days after injection. Tumor BCL6 levels were quantified using capillary electrophoresis and normalized to a GADPH

loading control. A clear decrease of BCL6 levels in the tumor was observed up to 10 h after dosing, with a maximal effect at ~4 h with free drug concentrations of > 1 nM. This result supported that small molecule-triggered BCL6 degradation is a promising therapeutic approach to develop anticancer agents.

Binding mode of BCL6 degrader **58** was studied (Figure 12, PDB 6TOM) along with a closely related non-degrader (PDB: 6TOL). Both compounds bind in the BCL6^{BTB} LG with highly similar binding modes, suggesting that the mechanism of action of the compounds, either as an inhibitor or a degrader, depended more on the nature of the substituents than a particular binding mode. Both molecules occupied the aromatic, linker, and part of HDCH sites (Figure 12A). The Cl atom fit into the deep hydrophobic group formed by M51, L'25 and N'21. Clear H-bonds were observed between the backbone carbonyl O atom of M51 and the bridging NH group (2.7 Å) as between the backbone NH of E115 and the urea O atom (3.1 Å).

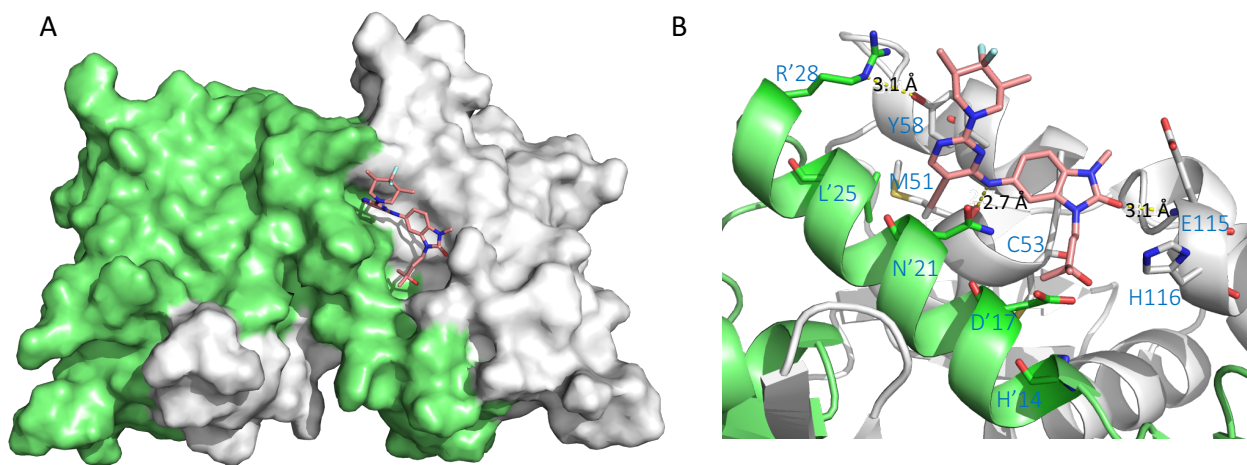
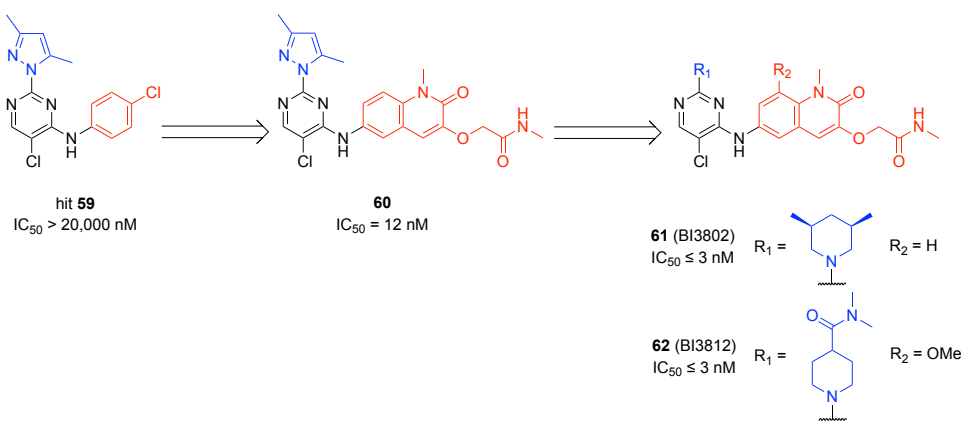


Figure 12. Binding mode of degrader **58** to BCL6^{BTB} (PDB ID: 6TOM). (A) Surface view of the binding of degrader **58** (pink) to the BCL6^{BTB} LG. Degrader **58** was shown in pink and BCL6^{BTB} monomers are shown in green and gray, respective. (B) Detailed binding mode of degrader **58** in complex with BCL6^{BTB}.

Kerres and coauthors screened a ~1.7 million-compound library using an FP assay and identified a 4-amino-5-chloro-pyrimidine hit **59** that inhibited the PPI between a co-repressor peptide and the BCL6^{BTB} dimer with an IC₅₀ value of 18 μ M (Scheme 11).⁶⁹ The binding affinity of the selected hit **59** to BCL6^{BTB} of hit was further validated by SPR ($K_d = 20 \mu$ M) and a biochemical ULight corepressor binding assay (IC₅₀ > 20 μ M). Optimization of hit **59** was conducted through a two-stage process focusing on the chlorophenyl ring and amino-substitution of the pyrimidine core, using a biochemical ULight corepressor binding assay. In the first stage, the chlorophenyl group of hit **59** was replaced by four different functional groups designed to interact with the backbone NH of E115. These efforts led to the identification of a new inhibitor **60** with a dramatically increased IC₅₀ value of 12 nM. Based on compound **60**, 10 additional compounds with various substitutions at the 2-position of the pyrimidine core were tested to yield two optimal inhibitors, **61** (BI-3802) and **62** (BI-3812) with IC₅₀ values \leq 3 nM. Overall, the SAR study included a total number of 14 new compounds with the IC₅₀ range of 20 μ M to \leq 3 nM.

Scheme 11. Optimization of hit **59 and generation of compounds **61** (BI-3802) and **62** (BI-3812)**



During biophysical characterization of the new compounds using co-crystallization and protein NMR experiments, the authors noticed that some of the new compounds caused protein precipitation. By confirming these atypical compounds inhibited the PPI of BCL6^{BTB} and its

corepressor while not disrupting BCL6^{BTB} dimerization using a cellular luminescence-based mammalian interactome (LUMIER) assays,⁹⁶ the authors ruled out the possibility that these compounds could denature the protein. In addition, the thermal stability of BCL6^{BTB} dimer was increased after treatment with both compounds **61** and **62** in differential scanning fluorimetry measurements, also supporting the maintenance of protein folding upon treatment of either type of compound. Moreover, the authors explored the binding regions of either compound on BCL6^{BTB} using a hydrogen deuterium exchange mass spectrometry (HDX-MS).⁹⁷ The atypical compound **61** showed more protection from HDX compared to the typical compound **62** in both close and remote regions of the inhibitor binding site. The authors attributed the extended protection to the induced dimer-dimer interaction by the atypical compounds.

To investigate whether this atypical behavior can translate into cellular effects, the authors performed Western blotting where they assessed BCL6 levels after treatment of compound **61** (1 μ M, 60 min). Compound **61** showed similar effects on BCL6 levels in ten lymphoma cells. Importantly, the authors noticed that the effect of compound **61** could be reversed by proteasome inhibitor MG-132,⁹⁸ indicating a mechanism of proteasome-dependent protein degradation. Moreover, treatment of compound **61** in combination with MG-132 led to the generation of a BCL6 form with decrease migration rate, in agreement with the formation of multi-ubiquitylated protein. The presence of multi-ubiquitin chains on BCL6 protein, after the treatment of degradation-inducing compound **61**, was further confirmed by probing a BCL6 immune precipitate with tandem ubiquitin-binding entities (TUBEs). Together, these results revealed that compound **61** led to cellular BCL6 degradation via the mechanism of ubiquitylation followed by proteasome-dependent degradation.

Compound **61** caused degradation of BCL6 by 50% in just ~5 min, and by over 90% in 2 h. This rapid degradation was also dose dependent with a DC₅₀ (half degradation concentration) value of 20 nM in SUDHL-4 cells. The correlation between the DC₅₀ values and IC₅₀ values derived from the inhibition of the PPI between BCL6^{BTB} and co-repressor peptides are high ($r^2 = 0.82$), supporting direct binding of the degradation-inducing compounds to BCL6^{BTB}. In addition, the authors found no correlation between the degradation efficiency and degradation potency for more potent compounds did not cause a higher percentage of protein degradation.

Next, degrader **61** was studied for its cellular effects, along with non-degrader inhibitor **62** (Scheme 11). To assess the selectivity of degrader **61** in cell, an affinity pull-down experiment was conducted using immobilized inhibitor **61** from cell lysate of Farage. The results confirmed that BCL6 was the major target of compound **61**. Importantly, no other BTB/POZ domain proteins were identified in the same pull-down experiment.

The authors found that the effective degradation of BCL6 by compound **61** required a functional ZF DNA-binding domain because corresponding construct with either deletion or mutation of the ZF domain could not get degraded when treated by the degrader **61**. To study the domains required for BCL6 degradation, the authors used HEK293 cells that lack endogenous BCL6. The levels of BCL6 after treatment of compound **61** (500 nM, 90 min) were determined by Western blotting. When a group of DNA-binding domains from different transcription factors were introduced to BCL6^{BTB}, the DNA-binding domains from ESRRA and basic helix-loop-helix domain of TCF4 could restore **61**-induced BCL6 degradation. On the other hand, the isolated BTB domain lacking the DNA-binding domain wound not get degraded. These results indicated that to achieve efficient degradation, the presence of the BCL6 ZF domain is not required, however, the localization of BCL6 protein to DNA, via a DNA binding domain, is required.

The effects of compounds **61** and **62** on levels of BCL6 target genes were studied by qPCR. Upon treatment of degrader **61** and inhibitor **62** (500 nM, 20 h) in SUDHL-4 cells, the number of genes that were increased and decreased were 87 and 17, respectively. Specifically, degrader **61** significantly induced BCL6 target genes including ATM, PRDM1, PTPN6, CD69, IRF4, and DUSP5. Importantly, many induced genes also indicated proximal (95% within 5kb of the transcription start site) BCL6 binding sites confirmed by ChIP experiments. Additionally, the authors took three of the robustly induced genes (PTPN6, RAPGEF1, and CHST2) and showed that degrader **61** (50 nM and 500 nM) induced these BCL6 target genes time-dependently (4, 6, 8, 16, 24 h). In comparison, the authors found in parallel experiments that the potency of inhibitor **62** in inducing BCL6 target genes (PTPN6 and RAPGEF1) was weaker than that of degrader **61**.

Finally, the antiproliferative effects of degrader **61** were evaluated in DLBCL cells using long-term proliferation assays with a concentration range of 0.01 to 3 μ M. Degrader **61** indicated antiproliferative activities (GI_{50} in the low micromolar range) in tested BCL6-dependent DLBCL cell lines including Farage, SUDHL-4, OCI-Ly7 and OCI-Ly1, and no obvious antiproliferative activities in BCL6-independent cells such as OCI-Ly19, MV-4-11 and Toledo. The authors also found that inhibitor **62** only showed antiproliferative activity at concentrations from 0.3 to 3 μ M, at least 100-fold higher than its IC_{50} value (≤ 3 nM). When a collection of compounds was tested, the observed antiproliferative potencies were well correlated with both the half degradation concentration (DC_{50}) ($r^2 = 0.68$) and co-repressor binding (IC_{50} , LUMIER assay) with the r^2 value of 0.68 and 0.89, respectively. These results highlighted that the effect of degrader **61** was through a BCL6-mediated mechanism. In summary, **61**-mediated BCL6 degradation caused antiproliferative effects in DLBCL cells. However, no significant cell death was observed upon treatment of degrader **61**. Subsequently, PK studies in SCID mice were performed after oral

dosing of compound **61** at 10 mg/kg (mean plasma AUC 1860 nM·h and Cmax 193 nM) and 100 mg/kg (AUC, 4650 nM·h; Cmax, 599 nM). Due to the relatively poor bioavailability of the compound, the authors didn't further explore the effect of compound **61** in animal models.

The co-crystal structure of hit **59** (PDB 5MW6) revealed the ligand binding to the BCL6^{BTB} LG. The key interaction between hit **59** and BCL6^{BTB} dimer was the anilinic HBD interacting with the backbone carbonyl O atom of M51, which brought the chloropyrimidine ring to fit snuggly into the aromatic site and placed the pyrazole ring in an approximate distance (3.6 Å) to R'24 for a potential hydrogen bonding interaction. The chlorophenyl moiety was found resting in the linker site, offering a handle for further optimization of binding affinity. As shown in Figure 13, the co-crystal structures of compound **61** (PDB 5MW2) revealed that the lactam carbonyl of this compound occupied the linker site and the O atom formed a key hydrogen bonding interaction (2.8 Å) to the backbone NH of E115. Furthermore, the acetamide tail of compound **61** added an H-bond interaction to the backbone NH of V117 (3.6 Å), which could also contribute to the potency improvement. Notably, the (3*S*,5*R*)-3,5-dimethylpiperidine ring on the pyrimidine ring was solvent-exposed without obvious interaction with the protein surface.

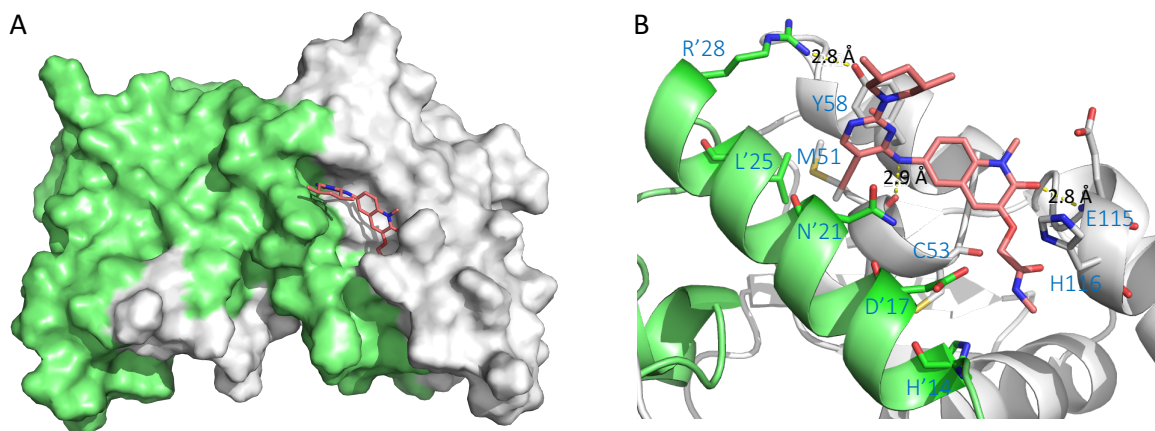


Figure 13. Binding mode of compound **61** to BCL6^{BTB} (PDB ID: 5MW2). (A) Surface view of the binding of inhibitor **61** (pink) to the BCL6^{BTB} LG. Compound **61** was shown in pink and BCL6^{BTB} monomers are shown in green and gray, respective. (B) Detailed binding mode of inhibitor **61** in complex with BCL6^{BTB}.

Summary and Perspective

As an oncogenic transcription factor, BCL6 is a promising target for the treatment of DLBCL and other important human cancer such as FLs, various forms of leukemia, breast cancer, and lung cancer. Currently there are no clinically approved drugs available targeting BCL6. Although BCL6 knockout is lethal, experimental evidence indicated that disrupting the PPI between the BCL6^{BTB} LG and its corepressors provides a safe therapeutic strategy.

First generation of BCL6^{BTB} inhibitors include peptide-based BPI, RI-BPI and small molecules **1** and **2**. These early inhibitors, albeit weak BCL6^{BTB} binders, provided good supporting evidence that targeting BCL6^{BTB} blocks the oncogenic activity of BCL6 and kills cancer cells. Also, the binding mode studies of these compounds have defined the binding pocket of the BCL6^{BTB} LG, which includes the acid, aromatic, linker, and HDCH sites. On the other hand, these inhibitors suffered from major drawbacks. For instance, BPI and RI-BPI involved challenging synthesis and poor oral bioavailability, and rhodanine-based compounds **1** and **2** were limited with respect to their modest binding affinity and PAINS-related pharmacologic properties.

Recent discovery of novel BCL6^{BTB} inhibitors is commonly driven by different high throughput assays (Table 1). Interestingly all the hits identified from different screen campaigns were small aromatic molecules that reversibly bind to the BCL6^{BTB} LG into the aromatic site formed by residues R'24, L'25, M51 and Y58. This aromatic site is occupied by SMRT residues H1426 and I1428,²⁸ or BCOR residues W509 and V511.²⁹ Another common feature of the

identified hits was the conserved HBD that interacted with the backbone carbonyl of residue M51. Note that the same backbone carbonyl O atom of M51 forms strong H-bonds with the imidazole sidechain of SMRT residue H1426 or indole sidechain of BCOR residue W509.^{28,29} Optimization of the chemical structure of the hits included two often parallel strategies: 1) expanding the molecule to occupy the linker and HDCH sites of the binding pocket, and 2) replacing the substitution on the aromatic core to the aid site for additional interactions to basic residues (R'28 and R'24) or favorable physiochemical properties (e.g., polarity and aqueous solubility).

Covalent inhibitors of the BCL6^{BTB} have also been pursued by employing reactive warheads to the chemical scaffolds of different reversible inhibitors, to target either the residue C53 in the HDCH site or the phenolic sidechain of Y58 on the edge of the aromatic site. To selectively target the thiol sidechain of residue C53, chloroacetamides, which showed reasonable intrinsic chemical reactivity, were introduced to aliphatic groups that could fit into the HDCH site. On the other hand, the phenol sidechain the solvent-exposed residue Y58 was activated by its neighboring residue R'28. This unique networking around the aromatic site offered an opportunity for covalent interaction by introducing a phenol-reactive group such as sulfonyl fluoride. Interestingly, both C53- and Y58-targeting irreversible inhibitors could effectively modify intracellular BCL6. However, these compounds only showed modest antiproliferative effects against BCL6-dependent DLBCL cells.

BCL6 protein PROTACs have also been pursued by installing PROTAC warhead (e.g., thalidomide) onto solvent accessible positions of optimized inhibitors. These PROTACs were cell permeable and caused fast and dramatic degradation of the BCL6 protein in cell. Interestingly, introduction of a dimethylpiperidine (or difluoromethylene analog) group directly to BCL6^{BTB} inhibitors without a spacer also caused sufficient degradation of the BCL6 protein in cell. The

antiproliferative effects was in consistence with the observed DC₅₀ values of these compounds. The rapid turnover of BCL6, which is maintained by continuous rapid translation through eIF4E,⁹⁹ makes it challenging to sustain low levels of the protein. Moreover, the fact that BCL6 knockout is lethal also raises concerns about full elimination of BCL6.³¹ On the other hand, the fact that degrader **58** was well-tolerated in mouse may ameliorate these concerns.

Several cell-free high-throughput competitive binding assays have been developed to measure the inhibitory potency and enable structure-activity relationship analyses. ELISA commonly includes a FLAG-tagged BCL6^{BTB} protein along with a biotinylated BCOR peptide immobilized onto streptavidin-coated microplates.^{38,42,50} Despite its success, ELISA requires wash steps to separate unbound labeled compound from bound ones in order to detect specific binding event. The involvement of the washing steps also limited this method in detecting low-affinity ligands. Compared to ELISA, the no-wash TR-FRET^{60,69,70,78,83,91,95} assay found their application in multiple studies, in which a long-lifetime fluorescent lanthanide (Er or Tb) labeled BCL6^{BTB} (energy donor) and a fluorescence-tagged corepressor peptide (energy acceptor) were employed in solution, which can be used to provide highly sensitive and reproducible measurement of competitive displacement of the fluorescence-tagged corepressor peptide by ligands from the BCL6^{BTB}-corepressor interface. In addition, FP assay represents another homogenous method to measure the inhibitory potency of compounds against the PPI between BCL6^{BTB} dimer and fluorescence-labeled BCOR repressor peptide.^{69,95} The potential utility of un-labeled protein is a big advantage of FP assays. However, FP can experience significant limitation from autofluorescence of testing compounds and light scattering.^{100,101} Moreover, AlphaLISA assay, including a pre-coated BCOR peptide (donor bead) and a BCL6^{BTB}-tagged acceptor bead, has also been developed to test potential inhibitors. AlphaLISA requires no wash steps, however, metal

chelators can cause false positives in the assays.⁷³ Besides, to evaluate the potency of nonequilibrium binding of irreversible inhibitors for BCL6^{BTB} protein, IPC assays have been developed. IPC offers a rapid determination of the specific binding constant k_{inact}/K_I for irreversible ligands,⁷⁹ although a control covalent ligand (e.g. irreversible fluorescence probe **37** for BCL6^{BTB} dimer⁷⁸) must be developed before using the assay.

Table 1. Competitive Binding Assays and Conditions

Assay	BCL6 ^{BTB} ^a	Co-repressor	Reference
ELISA	FLAG-BCL6 ^{BTB}	Biotinylated BCOR peptide ^b	38, 42, 50
TR-FRET	His tagged BCL6 ^{BTBc}	Hilyte647 SMRT peptide ^d	60, 91
	Biotinylated BCL6 ^{BTB}	TAMRA-peptide 4 ^e	72
	Biotinylated BCL6 ^{BTB}	FITC-BCOR peptide ^f	83
	Trx-His tagged BCL6 ^{BTBg}	AF633-BCOR peptide ^h	95
	Avi-tag biotinylated BCL6 ^{BTBi}	BCOR ULight peptide ⁱ	69
	GST-BCL6 ^{BTBk}	His tagged SMRT peptide ^j	70
FP	Avi-tag biotinylated BCL6 ^{BTBi}	FP-labeled BCOR peptide ^m	69
	BCL6 ^{BTB}	AF633-BCOR peptide ^h	95
AlphaLisa	His tagged BCL6 ^{BTB} ^c	Biotinylated BCOR peptide ^b	73
IPC	Avi-tag biotinylated BCL6 ^{BTBi}	37 (BCL6-FP)	78

^aBCL6^{BTB} (5-129); ^bBiotinylated BCOR peptide = Ac-RSEIISTAPSSWVVPGP-Lys-(ε-Biotin-(AC₅)₂)-OH; ^cHis tagged BCL6^{BTB} = Histidine (His) tagged BCL6^{BTB} (5-129) triple mutant (C8Q, C67R, and C84N); ^dHilyte™ Fluor 647 SMRT peptide H1426W = Hilyte™ Fluor 647-LVATVKEAGRSIWEIPR; ^eTAMRA peptide **4** = TAMRA-Abu(4)-VWYTDIIMRDWM-OH; ^fFITC, fluorescein isothiocyanate (Note: In the paper, BodipyFL-labeled BCOR peptide, but in SI FITC-BCOR peptide); ^gTrx, Thioredoxin; ^hAlexa Fluor 633-BCOR peptide = Ac-RSEIISTAPSSWVVPGP-Cys-Alexa Fluor 633-CONH₂; ⁱAvi-tag, amino acid sequence GLNDIFEAQKIEWHE; ^jBCOR ULight peptide = Ac-RSEIISTAPSSWVVPGP-Cys-ULight; ^kGST, Glutathione S-transferase; ^lHis tagged SMRT peptide = H6-LVATVKEAGRSIHEIPR; ^mFP-labeled BCOR peptide = 5-TAMRA-RSEIISTAPSSWVVPGP;

Direct binding assays including SPR, BLI, MST and ITC, have been utilized to characterize the binding event of small molecules to labeled or label-free BCL6^{BTB} protein (Table 2). SPR is a widely used optical biosensor technology employing a tagged BCL6^{BTB} protein for surface immobilization.^{38,42,50,60,69,91} It provides real-time measurement of the binding affinity (K_d) as well as kinetic parameters such as k_{on} and k_{off} with high sensitivity. By measuring the heat change

during ligand binding to GST-tagged BCL6^{BTB} protein,⁷⁰ BLI represents another biosensor technology for real-time detecting and quantifying the binding affinity and kinetics. Compared to SPR, BLI has advantages including increased throughput and wider sample compatibility. Because BLI is a fluidics-free system, it requires lower instrumentation cost as well. Moreover, by measuring directed movement of molecules in a temperature gradient, MST has also been used to quantify binding event between GST-labelled BCL6^{BTB} protein and its ligands.¹⁰² Compared to SPR and ITC, MST is a surface immobilization-free approach. Recent advances in the MST technology by following intrinsic fluorescence of tryptophan residue of a protein would make it a completely label-free method to study molecular interactions.¹⁰³ Besides, MST experiment can be done in versatile environment even in plasma and cell lysate.¹⁰³ ITC has been used to define the binding affinity, stoichiometry, and the enthalpy and entropy changes of the ligand binding to the BCL6^{BTB} protein.⁷³ In addition to binding affinity, results from ITC can provide information about structural features. However, ITC has limitations such as low throughput and relatively low resolution, it also requires relatively large amount of the protein.¹⁰⁴ Therefore, it is usually used as a secondary confirmation assay instead of primary screening.

Table 2. Direct Binding Assays and Conditions

Assay	BCL6 ^{BTB}	Reference
SPR	Biotinylated BCL6 ^{BTB}	38, 69
	Avi-tagged BCL6 ^{BTB}	42, 50
	H6-BCL6 ^{BTB}	60, 91
BLI	GST-BCL6 ^{BTB}	70
MST	GST-BCL6 ^{BTB}	102
ITC	BCL6 ^{BTB}	73

Three types of cellular assays have been developed (Table 3). The luciferase reporter assay was first established based on a GAL4-DBD-BCL6^{BTB} fusion construct. This assay was valuable not only in testing potential ligand interaction to BCL6^{BTB} in cellular environment, but also providing a flexible system to study the specificity of small molecules for BCL6^{BTB} over other related BTB domain proteins such as such as PLZF, Kaiso and HIC1.^{6,7,60,91} To follow the PPI of BCL6^{BTB} and its corepressors in cell, an M2H assay, based on activation of the luciferase reporter upon interaction of genetically fused GAL4-DBD-BCL6^{BTB} (“bait”) protein and GAL4-VP16-corepressor (“prey”) peptide, has also been developed.^{38,42,50,78} This assay platform was used by multiple groups to study the potency of reversible inhibitors in cellular environment. Besides, the bioluminescence resonance energy transfer-based assay system NanoBRET has been utilized to study the PPI between BCL6^{BTB} and its compressors in live cells. This assay uses the NanoLuc luciferase to provide the donor signal and the HaloTag ligand (HL) as the fluorescence energy acceptor. Compared to the luciferase reporter and M2H assays, the NanoBRET assay has a significant advantage for it offers cellular evaluation of a potential inhibitor in blocking the PPI between full length BCL6 and corepressor proteins. Recently, by following the fluorescence signal of eGFP using a translational fusion eGFP-BCL6^{BTB}, a FACS-based flow cytometry assay was developed that can be used to study the binding of irreversible inhibitors to in living cells.⁸³

Table 3. Cellular Assays and Conditions

Assay	Protein	Readout	Reference
Luc reporter	GAL4-BCL6 ^{BTB}	Luciferase catalysis	60, 91
M2H	^a GAL4-BCL6 ^{BTB} , VP16 ^b	Luciferase catalysis	38, 42, 50, 78
NanoBRET	full length BCL6 and corepressor	HL Fluorescence	95
FACS	eGFP-BCL6 ^{BTBc}	eGFP / mCherry	83

^aGAL4-BCL6^{BTB} = GAL4-BCL6^{BTB}(5-129); ^bVP16 = VP16-BCOR(112-753); ^ceGFP, enhanced green fluorescent protein.

Co-crystal structures of various compounds in complex with BCL6^{BTB} have been solved and successfully used in the design and development of new inhibitors. These compounds, although belonged to different families, bind to a common binding pocket in the BCL6^{BTB} LG employing four continuous binding sites: acid site, aromatic site, linker site and the HDCH site. It is worth noting that numerous inhibitors included a fragment that extruded from the acid site to the solvent-exposing space. These fragments could play an important role in crystallization of the inhibitor/BCL6^{BTB} complexes.^{60,69,73,91} As one example, the indole group of inhibitor **47**⁹¹ was sandwiched between two BCL6^{BTB} dimers (Figure 14A). The piperazine ring of compound **47** extended beyond the acid site and fit into a polar pocket formed by five residues E81, N84, I85, D88 and N96 (Figure 14B). Notably, this interaction mode involves the presence of crystal contacts of the ligand with the adjacent BCL6^{BTB} dimer. Such interactions may impact details of the interaction orientation of the ligand with the primary BCL6^{BTB} dimer to which it is bound, as previously discussed.⁷¹

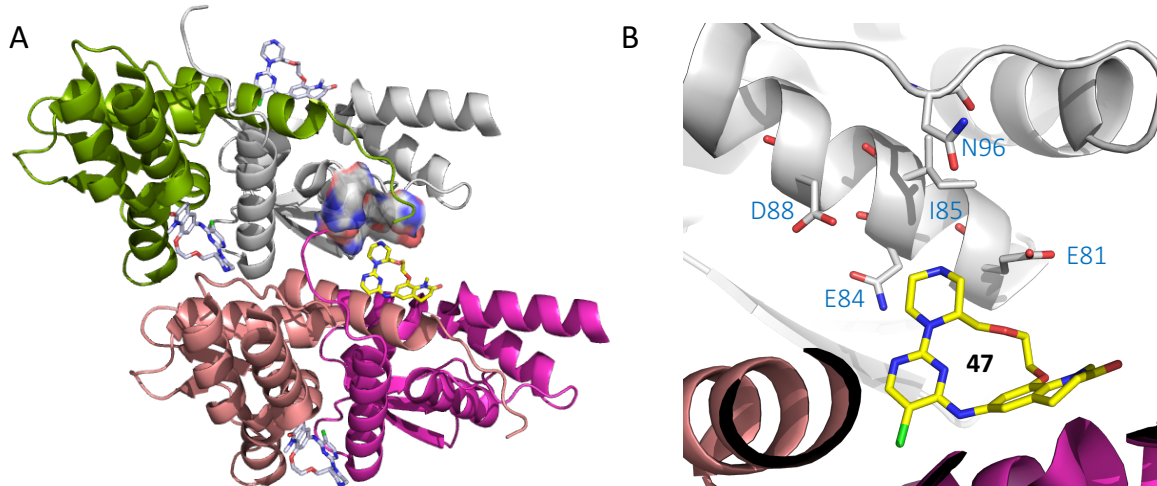


Figure 14. Binding mode of compound **47** to BCL6^{BTB} (PDB ID: 6EW8). (A) Overall view of the packing of one BCL6^{BTB} dimer (green and white) on top of a second BCL6^{BTB} dimer (orange and purple) via inhibitor **47**. The surface area of the neighboring BCL6^{BTB} dimer that was interact

with inhibitor **47** (yellow) was highlighted. (B) Detailed binding mode of inhibitor **47** to the neighboring BCL6^{BTB} dimer. Key residues involved in the interactions were shown in sticks.

Using a combination of various biological and biophysical methods, numerous BCL6^{BTB}-targeting hits have been identified, which were subsequently developed into lead compounds through medicinal chemistry efforts. These leads could disrupt BCL6 interaction with its partner corepressor-derived peptide in various binding assays with an affinity range of low-nanomolar to low-micromolar. They usually also demonstrated mid-nanomolar to micromolar activities in engineered cellular reporter system. Their binding modes to BCL6^{BTB} LG were confirmed by X-ray co-crystal structures and 2D NMR (e.g., ¹H,¹⁵N-HSQC). Of these new compounds, reversible inhibitors (e.g., compound **28**) by Guo and coworkers showed strong antiproliferative effects.⁷⁰ Irreversible inhibitors (e.g., compound **40**) from Takeda indicated antiproliferative effects although selectivity for BCL6-dependent DLBCL cells was not determined.⁷⁸ Moreover, recent BCL6 degraders from both ICR (e.g., compound **58**)⁹⁵ and Boehringer (e.g., compound **61**)⁶⁹ showed selective antiproliferative effects for BCL6-dependent over BCL6-independent DLBCL cells. It has also been shown that the antiproliferative effects of BCL6 degraders were stronger than those of the corresponding inhibitors. On the other hand, several optimized leads indicated either no effect on DLBCL proliferation,^{38,50,73} or non-selective antiproliferative effects for both BCL6-dependent and BCL6-independent cells.^{60,91} The failure of these compounds is likely due to the limited cellular permeability^{38,50,73} or likely the lack of adequate target engagement. Considering no detailed target engagement studies were performed in these studies, it is not known whether the compounds could actually block BCL6 mediated formation of repression complexes nor for how long the inhibitory event might occur.

The mechanism of action of potent BCL6 PROTACs and degraders was evaluated to confirm the involvement the BCL6 protein.^{69,91,95} For example, PROTAC **50** caused selective degradation of the BCL6 protein in cell,⁹¹ although only incomplete degradation of the protein has been observed. The degrader **58** caused rapid but incomplete degradation of the BCL6 protein,⁹⁵ another degrader **61**, on the other hand, caused degradation of BCL6 in a DNA-binding dependent manner.⁶⁹ Although the degrader **61** caused gradual proliferation arrest of DLBCL cells without apoptosis, which was quite different than what was observed with BCL6 shRNA, siRNA, CRSPR, or BPI, similar effects by the degrader **61** was recently observed using inducible BCL6 knockout model for DLBCL.¹⁰⁵

In summary, BCL6^{BTB} holds excellent potential as a therapeutic target that has only recently started to be systematically studied. Numerous BCL6 inhibitors and degraders have been characterized using various *in vitro* and *in vivo* assays, with a few examples indicating promising effects in killing BCL6-dependent cancer cells. In general, the BCL6 degraders have demonstrated more favorable antiproliferative effects than those of BCL6 inhibitors. One of the major impediments to currently available studies was the challenge of transforming inhibitors with high binding potency to BCL6^{BTB} into anticancer efficacies in cell and *in vivo*. This may be due to the fact that none of the new compounds have been evaluated through the proper target engagement experiments such as QChIP or target occupancy over time assays,⁷ which are considered critical to understand the likely impact of those compounds. Specifically, because BCL6 must recruit its repression complex in order to perform its transcriptional repression function,^{28,29} it is critical to test the compounds' effects in disrupting the repression complex by compounds. Without these results, one cannot know if the compounds had on-target efficacy. For example, the new compounds from Takeda, AstraZeneca, and Boehringer have not been tested for the effects of

disrupting the BCL6 repression complex on chromatin of BCL6, as occurred with the early BPI,⁴ small molecule inhibitor **2**,⁷ or by CRISPR knockdown.¹⁰⁶ BCL6 CRISPR is lethal to lymphoma cells.¹⁰⁶ Up to now, there have been no compounds able to sufficiently down-regulate BCL6 so as to mimic the effect of genetic ablation by CRISPR. Therefore, none of the new compounds (except for inhibitor **28**) to date has been shown to kill lymphoma cells in short period of time (48-96 h) and has indicated anticancer effects in DLBCL animal models. Continuous efforts are needed in the development of potent, selective, and cellularly active BCL6^{BTB} inhibitors to facilitate studies on the therapeutic potential of the oncogenic transcription repressor BCL6 in drug discovery. We expect that the fundamental insights gained both published and from future studies will ultimately enhance the ability of medicinal chemists to design inhibitors for BCL6^{BTB} as well as a wide range of disease-relevant proteins that contain BTB domains. Such novel compounds, in addition to their potential therapeutic utility, are expected to prove useful in investigating the pharmacology of other protein members of the BTB/POZ family such as Kaiso and LRF.

AUTHOR INFORMATION

Corresponding Author

Fengtian Xue – Department of Pharmaceutical Sciences, University of Maryland School of Pharmacy, 20 Penn Street, Baltimore, Maryland 21201, United States; Email: fxue@rx.umaryland.edu; orcid.org/0000-0002-4132-9887

Authors

Yong Ai – Department of Pharmaceutical Sciences, University of Maryland School of Pharmacy, 20 Penn Street, Baltimore, Maryland 21201, United States; orcid.org/0000-0002-7598-7992

Lucia Hwang – Department of Pharmaceutical Sciences, University of Maryland School of Pharmacy, 20 Penn Street, Baltimore, Maryland 21201, United States

Alexander D. MacKerell – Department of Pharmaceutical Sciences, University of Maryland School of Pharmacy, 20 Penn Street, Baltimore, Maryland 21201, United States

Ari Melnick – Department of Medicine Weill Cornell Medicine, New York, New York 10021, United States; Department of Pharmacology, Weill Cornell Medicine, New York, New York 10021, United States; orcid.org/0000-0002-8074-2287

Notes

The authors declare no competing financial interest.

Biographies

Yong Ai earned his Ph.D. in Medicinal Chemistry from China Pharmaceutical University under the supervision of Professors Yihua Zhang and Sixun Peng. He is currently a Postdoctoral Fellow of the Department of Pharmaceutical Sciences, School of Pharmacy, University of Maryland,

Baltimore. His research interests mainly focus on inhibitors of the Wnt signaling pathway and BCL6 BTB domain.

Lucia Hwang is a second-year PharmD candidate at the University of Maryland School of Pharmacy. Her research focuses on developing small molecules to inhibit the BCL6^{BTB} and heme uptake in *Pseudomonas aeruginosa*.

Alexander D. MacKerell Jr. received a Ph.D. in Biochemistry in 1985 from Rutgers University, which was followed by postdoctoral fellowships in the Department of Medical Biophysics, Karolinska Institutet, Stockholm, Sweden and the Department of Chemistry, Harvard University. In 1992 he assumed his faculty position in the School of Pharmacy, University of Maryland where he is currently the Grollman-Glick Professor of Pharmaceutical Sciences and the Director of University of Maryland Computer-Aided Drug Design Center. MacKerell is also Co-founder and CSO of SilcsBio LLC, a biotech company in the area of computer-aided drug design and development.

Ari Melnick received his doctorate in Medicine from the University of Buenos Aires in Argentina. He completed a residency in internal medicine and a fellowship in hematology/oncology at the Mount Sinai School of Medicine in New York City. Then, he did a post-doctoral fellowship with Dr. Jonathan Licht at Mount Sinai on the topic of transcriptional regulation in leukemia. Dr. Melnick started his independent laboratory at the Albert Einstein College of Medicine, and then moved to the Weill Cornell Medical College in New York City, where he is currently the Gebroe Family Professor of Hematology/Oncology and Professor of Pharmacology and Immunology.

Fengtian Xue received his BS in chemistry from the University of Science and Technology of China in 2001. He completed his PhD in chemistry under the direction of Professor Christopher Seto at Brown University in 2007, and later pursued a post-doctoral fellowship in Medicinal

Chemistry under the supervision of Professor Richard Silverman at Northwestern University from 2007 to 2009. Dr. Xue is currently an Associate Professor in the Department of Pharmaceutical Sciences at University of Maryland Baltimore. His research focuses in using experimental methods from Organic Chemistry and Medicinal Chemistry to develop small molecule therapeutic agents for various diseases including various cancers, bacterial/parasitic infections, and metabolic diseases.

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ABBREVIATIONS USED

ABL, abelson proto-oncogene; AML, acute myeloid leukemia; B-ALL, B-acute lymphoblastic leukemia; BBD, BCL6 binding domain; BCL6, B-cell Lymphoma 6; BCOR, BCL6 corepressor; BPI, BCL6 peptide inhibitor; BTB, Broad-Complex, tramtrack and bric a brac; GCB, germinal center B-cell; ABC, activated B-cell; CADD, computer-aided drug design; CETSA, cellular thermal shift assay; CML, chronic myeloid leukemia; CCK8, Cell Counting Kit 8; DC₅₀, the concentration at which the target is degraded by fifty percent (50%); DLBCL, diffuse large B-cell lymphomas; DMSO, dimethyl sulfoxide; DTT, dithiothreitol; ELISA, enzyme-linked immunosorbent assay; FBDD, fragment-based drug discovery; FLs, follicular lymphomas; FP, fluorescence polarization; GC, germinal center; HDX-MS, hydrogen deuterium exchange mass spectrometry; HTS, high-throughput screening; IC₅₀, the concentration of drug at which 50% of

the target is inhibited; IPC, irreversible probe competition; ITC, isothermal-titration calorimetry; LE, ligand efficiency; LG, lateral groove; LUMIER, luminescence-based mammalian interactome; M2H, mammalian two-hybrid; MLL, mixed-lineage leukemia; MSD, Meso Scale Discovery; MST, microscale thermophoresis; NCOR, nuclear receptor corepressor; NHL, non-Hodgkin's lymphomas; NMR, nuclear magnetic resonance; PAMPA, parallel artificial membrane permeability assay; PDB, RSCB Protein Data Bank; PPI, protein-protein interaction; PRDM1, positive regulatory domain zinc finger protein 1; PROTAC, proteolysis targeting chimera; RD2, second repression domain; RTHS, reverse two-hybrid system; SILCS, site identification by ligand competitive saturation; SMRT, silencing mediator for retinoid or thyroid hormone receptors; SPR, surface plasmon resonance; STD-NMR, saturation transfer difference-nuclear magnetic resonance; TR-FRET, time-resolved fluorescence resonance energy transfer; NP-CGG, 4-hydroxy-3-nitrophenyl)acetyl-chicken gamma globulin; MLM, mouse liver microsomes; BLI, biolayer interferometry; ICR, Institute of Cancer Research.

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