

Linearizable Macrocyclic Peptide Libraries for Affinity Selection-Mass Spectrometry

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1 Abstract

Macrocyclic peptides are attractive for therapeutic development but have been limited in their application to combinatorial library selection from synthetic libraries. Here, we establish a synthetic approach based on split-pool chemistry to produce 100-million membered macrocyclic libraries containing natural and non-natural amino acids. Near-quantitative intramolecular disulfide formation is facilitated by rapid (<10 minute) oxidation by iodine to prepare macrocyclic synthetic libraries in solution. Treatment with dithiothreitol post-affinity selection enables near-quantitative reduction of the library members, rendering the linear analogs amenable to standard tandem mass spectrometry sequencing. We demonstrate the utility of these libraries to discover novel macrocyclic binders to cadherin-2 and the anti-hemagglutinin antibody clone 12ca5. The lead cadherin-binding peptide (**CBP**) is endowed with nanomolar binding affinity measured by biolayer interferometry (BLI, apparent dissociation constant $K_D = 53$ nM). Structure-activity relationship (SAR) studies including alanine and D-amino acid scans reveal the amino acids responsible for driving affinity (hot-spots) and the positions tolerating mutagenesis (cold-spots). Informed by SAR data, two new macrocyclic libraries are designed to derivatize these positions with a variety of abiotic amino acids based on the hot- and cold-spots. Following affinity selection and experimental validation by BLI, 10 high-affinity ligands out of 10 discovered were identified from the library that derivatized the CBP cold-spots, while zero of the two peptide ligands discovered from the hot-spot library were high-affinity binders. Of these noncanonical CBPs (**NCBP**), **NCBP-4** demonstrates improved affinity to cadherin-2 ($K_D = 29$ nM). Overall, we expect that this work will pioneer the use of large-scale macrocyclic libraries to further catalyze therapeutic peptide discovery and development.

2 Introduction

Macrocyclic peptides show significant therapeutic promise with advantages over small molecules to disrupt protein-protein interactions and over proteins to cross biological membranes barriers.^{1–4} Specifically, macrocyclization can impart several potential benefits to linear precursors, including increased proteolytic stability, cell permeability, and oral bioavailability.^{5,6} Proteases often engage and degrade peptides in extended β -strand conformations.^{7,8} Macrocyclization can offer proteolytic resistance by limiting conformational accessibility of the peptide backbone to the enzyme active site, and enable the use of specific engineerable scaffolds (e.g., stapled α -helices).^{9–12} Cyclization is central to the currently applied design principles to achieve passive cell permeability, in addition to strategies that modulate molecular weight, polar surface area, hydrogen bond interactions, and shape.^{13–16} Combining proteolytic stability and passive permeability can impart oral bioavailability for peptide-based drug candidates, which can further be improved by pharmaceutical formulation.^{17,18} For these reasons, macrocyclic libraries are preferred for screening with peptide ligand discovery platforms. In addition, the direct identification of macrocyclic peptide binders from these selections streamlines subsequent development by alleviating the need to optimize suitable cyclization sites. Lastly, the conformational constraint imparted by macrocyclization may improve discovery rates of ligands from libraries against challenging targets.^{19–21}

Genetically-encoded discovery platforms generally access macrocyclic peptide libraries while focusing on high diversity ($>10^8$ members),^{22–28} while synthetic libraries can access the non-natural chemical space at generally lower diversity ($<10^8$ members).^{29–33} While more stable macrocyclization linkages are preferred (e.g., thioether or alkyl chain),^{24,34} the disulfide linkage is suitable at the ligand discovery stage, and does not require any chemical modification or treatment that could compromise genetic amplification in some platforms.^{28,35} The disulfide linkage has been used to create macrocyclic libraries for over two decades in phage display discovery platforms,^{36–38} and is commonly encountered in clinically-approved drugs.^{5,39} Synthetic libraries generally leverage the broader use of non-natural or abiotic functionalities, which have frequently appeared critical to the success of clinical peptide drug candidates including inhibitors to the interleukin-23 receptor (IL-23R), mouse double minute 2 (MDM2), β -catenin, and proprotein convertase subtilisin/kexin type 9 (PCSK9).^{17,29,40–42} Because they cannot be genetically-encoded or amplified, synthetic libraries are screened directly³⁰ as in affinity selection decoded by mass spectrometry (AS-MS).^{43,44} With a key exception of DNA-encoded libraries,⁴⁵ state-of-the-art synthetic macrocyclic libraries generally number below tens of thousands of individual compounds.³⁰

The complexity of decoding macrocyclic peptide sequences in mass spectrometry is a historic limitation for the use of synthetic macrocyclic libraries in affinity selection discovery platforms. Experimental approaches for decoding macrocyclic libraries include computational processing of mass spectra^{46–49} and chemically-triggered linearization.^{50–55} Computational approaches process primary, secondary, and various tertiary mass spectra of cyclic peptide fragments, and have excelled where database matching is possible.^{46,49} For *de novo* sequencing, the complexity of enumerating virtual spectra dramatically increases as the number of monomers and library size increases, and has only been demonstrated up to $\sim 1,000$ -membered libraries.^{47,56} Chemically-triggered linearization adds a synthetic step that must be near-quantitative and high-yielding to enable bottom-up sequencing of non-cyclic peptides, which has been demonstrated at very high diversities.⁵⁷ However, most chemical linearization treatments are harsh and/or rely on the inclusion of non-standard chemical functional groups at fixed positions, limiting library composition.^{50–55} Moreover, these approaches have yet to be demonstrated on high-diversity libraries ($\sim 10^8$ members or more).

We demonstrate here ligand discovery from high-diversity libraries (10^8 members) utilizing AS-MS against anti-hemagglutinin antibody clone 12ca5 (hereinafter abbreviated as 12ca5) and cadherin-2. Cyclization by disulfide bond formation is accomplished using aqueous iodine. We verify the integrity of the library utilizing Ellman's assay and size-exclusion chromatography (SEC) to confirm near-quantitative intramolecular macrocyclization of synthetic combinatorial libraries. Linearization is accomplished by mild reduction with heat

and 1,3-dithiothreitol (DTT) and confirmed by Ellman's assay to enable standard tandem MS sequencing. We apply these new macrocyclic high-diversity libraries containing natural and non-natural (or noncanonical) amino acids in an MS-based affinity selection platform for *de novo* peptide ligand discovery.

We demonstrate successful discovery of nanomolar ligands against 12ca5 and the ectodomain of cadherin-2. The 12ca5 protein binds peptides containing the sequence D**DY(A/S).^{58,59} While 12ca5 has been used to benchmark linear AS-MS libraries,^{33,60} we utilize it here to benchmark and additionally validate the use of the new high-diversity macrocyclic libraries. Cadherin-2 was considered as a second target because of the potential impacts for chemical biology that an affinity reagent could provide, ranging from basic cell adhesion, to neural synapses formation,⁶¹ to the construction of intercalated discs of mammalian heart,⁶² as well as potential drug delivery due to its relative tissue selectivity in the brain and heart.⁶³ These critical roles in biology are generally facilitated by homodimerization in domain 1 and 2.^{61,64,65} Thus, we sought to discover ligands that bind to cadherin-2 domains 4 and 5 as they may not interfere with cadherin-2 function. Outside of domain 1 and 2, there are no ligands to cadherin-2 to our knowledge.^{64,65} Lastly, we demonstrate the incorporation of non-natural amino acids for second-generation ligand discovery in libraries designed with input gained by structure-activity relationship (SAR) data gathered on the initially discovered cadherin-binding peptide (CBP). Taken together, the successful discovery of macrocyclic ligands to both targets from AS-MS demonstrates the potential deployment of ultra-large synthetically-prepared macrocyclic libraries for peptide ligand discovery and development.

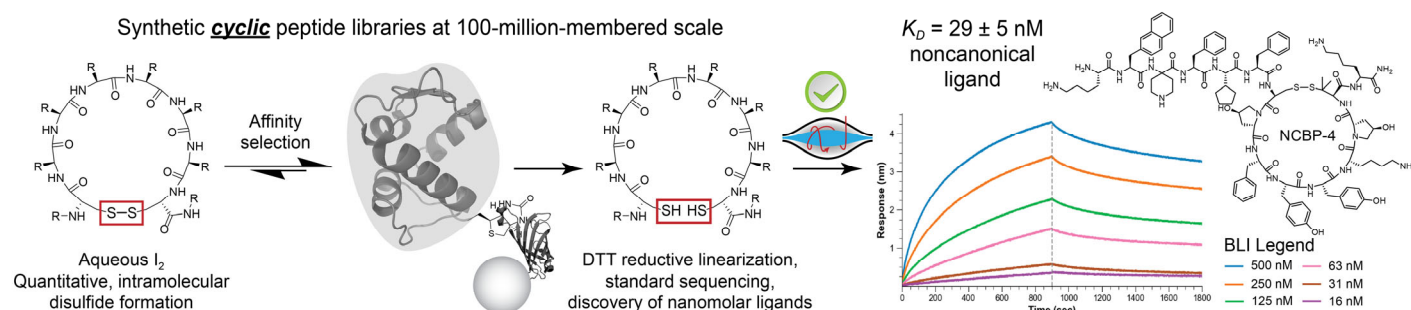


Figure 1. Disulfide linkages allows for high-number diversity libraries compatible with decoding by tandem MS/MS. Libraries of macrocyclic peptides are prepared for affinity selection by oxidation of cysteine analogs using aqueous iodine, providing a near-quantitative conversion to intramolecular macrocyclic peptides. After affinity selection, peptides can be quickly linearized using dithiothreitol (DTT) for sequencing using MS/MS.

3 Results and Discussion

Due to its demonstrated utility in genetically-encoded libraries, disulfide-induced macrocyclization of peptides has become a routine approach with a variety of existing methods to facilitate the oxidation step. Several different methods exist to form the disulfide linkage on single peptides, including oxidation using dimethyl sulfoxide,⁶⁶ a gentle stream of air, or aqueous iodine. Ideally, the macrocyclization step can be introduced during standard peptide library synthesis without incurring production delays or yield losses. The isolation of peptides in DMSO-containing solutions could be challenging as the solvent cannot be easily evaporated or lyophilized and solid-phase extraction could incur sample loss due to the DMSO content without further aqueous dilution.⁶⁷ In comparison, oxidation utilizing iodine presents itself as a rapid method compatible with mixtures of aqueous or organic solvents, and can even facilitate the formation of disulfide bonds on resin during solid-phase peptide synthesis.^{43,68} However, longer reaction times on resin promote iodine-based side reactions, therefore rapid in-solution oxidation is preferred (< 15 min).⁶⁹

Iodine facilitated formation of macrocyclic peptide libraries at 200-million membered scale. We synthesized macrocyclic libraries by split-and-pool solid-phase peptide synthesis using mono-sized 20 μm resin (8.33 g of resin, 2.00 mmol scale total), with each bead providing ~ 1 pmol of peptide. Two billion-membered libraries were prepared with the designs of $\underline{\text{C}}\text{X}_{12}\underline{\text{C}}\text{K}$ and $\text{X}_6\underline{\text{C}}\text{X}_6\underline{\text{C}}\text{K}$, where X = all canonical amino acids except Cys, to control disulfide formation, and Ile because it is isobaric in mass with Leu (18 amino acids) and $\underline{\text{C}}$ = cysteine, homocysteine, and penicillamine (Figure 3A). The libraries were split into five separate 200-million-membered aliquots and cleaved from the solid phase resin using a cleavage cocktail. After ether trituration and lyophilization, peptide libraries were cyclized in 5% acetonitrile in water (with 0.1% trifluoroacetic acid) at ~ 2 mg/mL (~ 1 mM) by dropwise addition of ~ 1 eq. iodine in methanol until a yellow-brown color persisted. After 5-10 minutes at room temperature in the dark, the reaction was quenched with aqueous ascorbic acid to provide a colorless solution again (3.5 eq.). These libraries were then characterized to verify the efficiency of the oxidation and linearization reactions as well as their structure (intramolecular vs intermolecular disulfide formation).

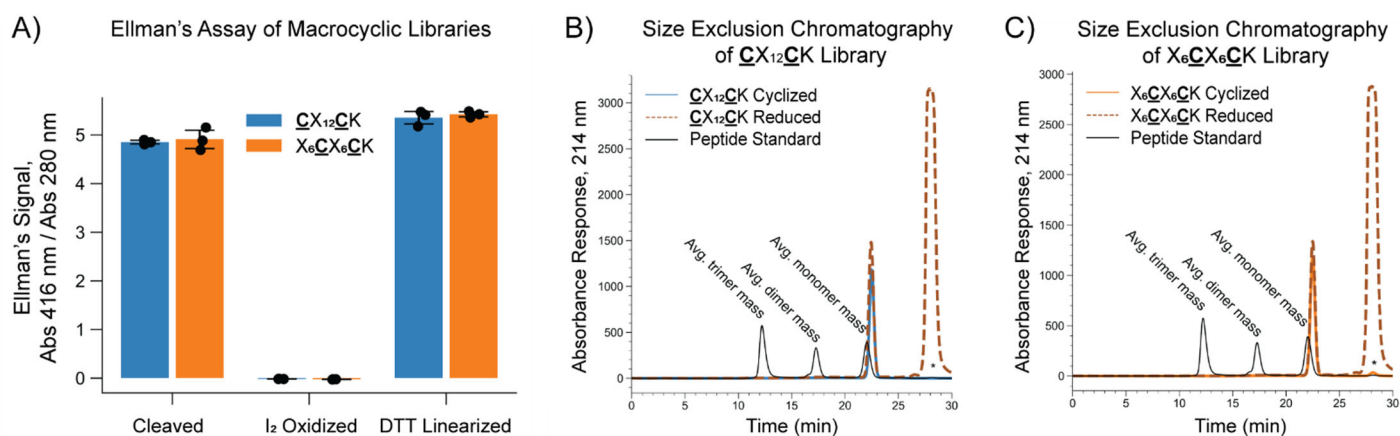


Figure 2. Characterization of macrocyclic libraries based on size and thiol concentration showed near-quantitative formation of intramolecular disulfide bonds. (A) Ellman's assay showed expected changes in total thiol concentration of the peptides directly after cleavage from the solid phase resin, after oxidation by dropwise addition of ~ 1 eq. of 60 mM iodine in methanol to facilitate disulfide formation (room temperature, 5-10 minutes in the dark, subsequent quench with 3.5 eq. aqueous ascorbic acid), and after reduction using DTT (50 mg/mL, ~ 1000 eq at 60 $^{\circ}\text{C}$ for 15 minutes). Free thiol was quantitatively consumed during the oxidation process and was restored after linearization to concentrations comparable to those determined directly after cleavage. (B,C) Size exclusion chromatograms of absorbance at 214 nm of two macrocyclic libraries compared to molecular weight standards corresponding to the average mass of monomeric, dimeric, and trimeric species. Library samples were ran using the cyclized form (later used in affinity selection experiments) and the DTT linearized form, demonstrating the formation of intramolecular disulfide bonds. Peaks marked with an asterisk (*) were residual elements from the sample buffer. $\underline{\text{C}}$ = cysteine, homocysteine, and penicillamine.

Iodine-promoted cyclization was highly efficient and provided near-quantitative oxidation to disulfide by thiol quantification using Ellman's assay. We quantitated thiol oxidation by performing an Ellman's assay, normalized by the absorbance of the library at 280 nm (Figure 2A). The thiol content of the library was quantified by Ellman's reagent after cleavage, cold ether trituration, and solid-phase extraction (SPE), to remove any remaining reducing scavengers. Upon aqueous resuspension of the library, a strong thiol signal was observed.

This signal was eliminated completely by the treatment of the library with iodine, ascorbic acid quench, SPE purification, and aqueous resuspension, consistent with the near-quantitative formation of disulfide bonds.

With macrocyclic libraries in hand, a MS-friendly protocol to reduce and linearize the peptides was devised to enable standard tandem sequencing approaches. Both DL-dithiothreitol (DTT) and tris(2-carboxyethyl)phosphine (TCEP) were considered for disulfide reduction. Compared to TCEP, DTT is smaller and more hydrophilic and less likely to be retained on reverse phase columns. Because of the concern for its retention on column, TCEP was utilized in a bead-immobilized form, whereas DTT was directly added to each sample just before mass spectrometry, reducing handling steps and potential sample loss. Due to its high solubility, DTT was utilized at 50 mg/mL, (~1000 eq) at 60 °C for 15 minutes, whereas immobilized TCEP was utilized at 20 eq at room temperature for 25 minutes per manufacturer protocol.⁷⁰ For DTT-treated samples, an SPE purification was performed to remove excess DTT reagent, whereas samples treated with immobilized TCEP were isolated by centrifugation. While TCEP only provided incomplete reduction (40% for X₆CX₆CK and 85% for CX₁₂CK), DTT provided near-quantitative disulfide reduction (~100% of original Ellman's signal across all libraries, Figure S1). Additionally, the reduction efficiency by DTT was found to be similar at pH 3 and pH 8 (Figure S1). The direct reduction at pH 3 was performed to mimic a prepared sample in 0.1% formic acid in water, which could then directly be injected in the mass spectrometer for tandem sequencing. Overall, these data support the near-quantitative formation and reduction of disulfide bonds.

Size exclusion chromatography (SEC) confirmed disulfide bonds correspond to formation of intramolecular species, producing an almost-exclusively monomeric macrocyclic peptide library. Utilizing SEC to separate the library by its apparent molecular weight, we assessed if the iodine-facilitated disulfide bond formation was intramolecular or intermolecular. Library samples were injected on a SuperDex[®] 30 10/300 GL column, which can distinguish the molecular weight range of 100 to 7000 Da, to analyze the presence of monomeric, dimeric, and oligomeric peptides induced by iodine oxidation. Aliquots of 25 µg of library were injected after oxidation by iodine and purification by SPE, as well as in the reduced form after treatment with DTT. A custom low-molecular weight peptide standard was also prepared using peptides that correspond to the average molecular weights of monomeric, dimeric, and trimeric species along with the exclusion limit of the column; the components of the custom standard are given in Table S1. As shown in Figures 2B and 2C, only monomeric library species were observed when compared to the peptide standard, indicating that the disulfide bonds formed were intramolecular rather than intermolecular. This result, in tandem with the data from the Ellman's assay, asserted a nearly quantitative conversion of peptide thiols to intramolecular disulfide bonds and confirmed this technique provides a facile method for preparing high-diversity macrocyclic peptide libraries.

The high-diversity macrocyclic libraries verified to be compatible with standard tandem MS/MS sequencing protocols and DTT facilitated the expected recovery of high-confidence peptide sequencing. Small aliquots of library (~1000 beads equivalent to ~1000 sequences) were taken at various points in the affinity selection workflow from cyclization to linearization, including directly after cleavage from resin, after cyclization with iodine, and after linearization with DTT. About 8 µg of peptide library was purified via a C₁₈ STAGE tip⁷¹ from three each steps in the protocol: linear from cleavage, iodine-macrocyclized, and DTT-linearized. As expected, the library samples taken directly after cleavage and after linearization showed high sequencing confidence, described by *de novo* sequence IDs using PEAKS Studio 8.5 with an assigned local confidence (ALC) greater than 85% and an absolute mass error <5 ppm (see Figure 3B and Figure S2).⁷² Conversely, the macrocyclized library sample showed poor sequencing confidence, consistent with unproductive fragmentation caused by the disulfide macrocycle. Lastly, the linearization by DTT in the mass spectrometry sample provided a significant recovery of the peptides discovered in standard tandem sequencing methods (Figure 3B).

Analysis of the MS/MS sequencing data demonstrated a balanced distribution of amino acid monomers throughout the library, as well as the incorporation of non-canonical cysteine analogs. Histograms showing the monomer distribution among high-confidence sequence assignments are given in Figure 3C. Although penicillamine and methionine are isobaric, the fixed positions of penicillamine allowed for effective filtering of

sequences to prevent inaccurate assignments. Interestingly, the X₆CX₆CK library design sequenced with higher confidence overall compared to the CX₁₂CK library design, suggesting a benefit of the intermediate cysteine position during fragmentation events in sequence assignment. Overall, these results corroborate the successful split-and-pool synthesis, oxidation, reduction, and tandem sequencing decoding of the macrocyclic libraries.

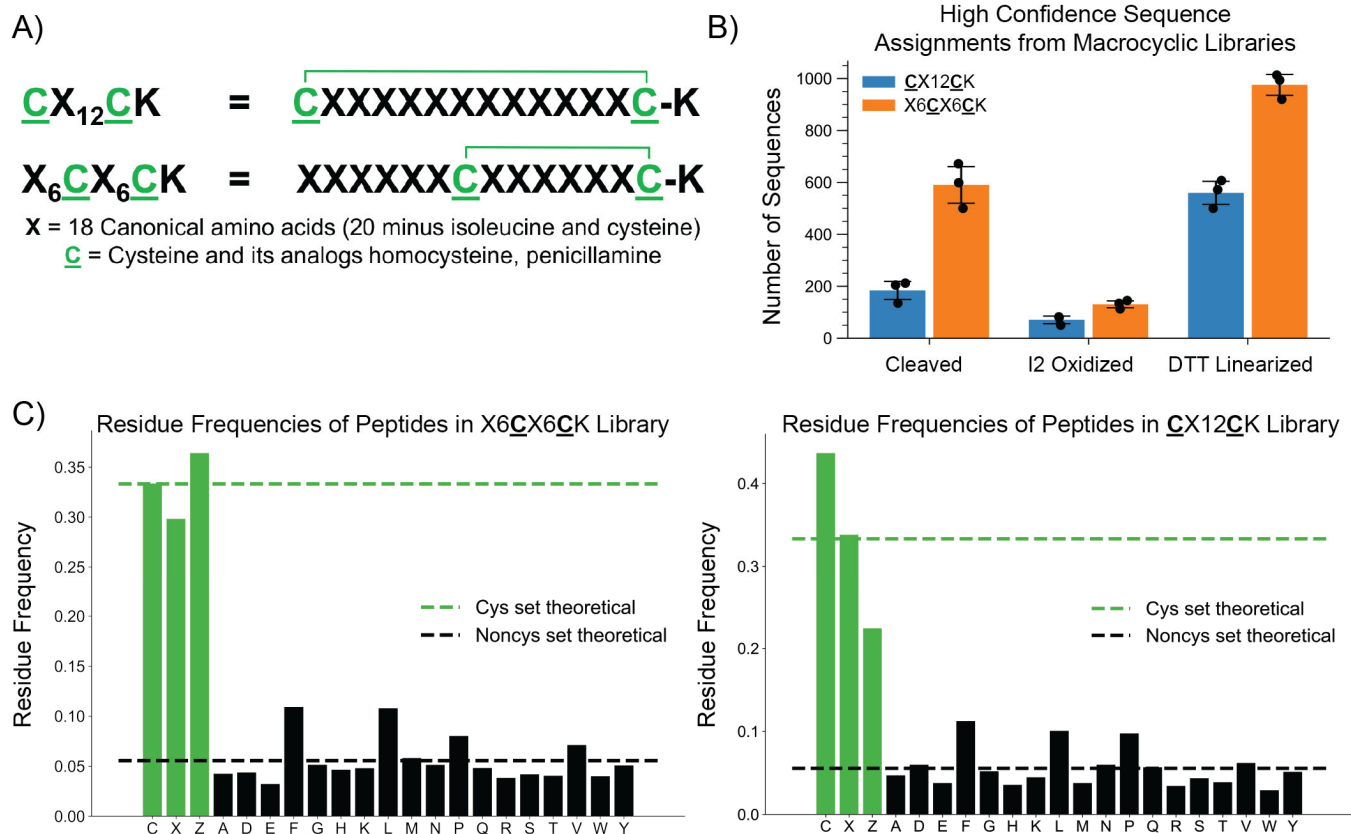


Figure 3. Characterization of the macrocyclic peptide libraries by tandem MS/MS sequencing shows successful split-and-pool synthesis of 15-residue libraries. (A) The macrocyclic peptide libraries were synthesized according to two designs, with a large 12-member macrocycle or a smaller 6-member macrocycle. Additionally, cysteine analogs including homocysteine and penicillamine were used to increase the diversity around the resulting disulfide linkage. (B) High-confidence sequence assignments of ~1000-member library samples directly after cleavage, after oxidation by I₂, and after subsequence reduction using DTT show a loss and gain of sequencing capabilities with the macrocyclization process as expected (n = 3). High-confidence sequences were determined as having a calculated average local confidences (ALC) in PEAKS Studio 8.5 for the *de novo* sequence assignment by PEAKS Studio 8.5 greater than 85% and an absolute mass error <5 ppm. (C) Normalized residue frequencies assignments (as a fraction) show a balanced incorporation of amino acids in the variable positions (black) and cysteine analog positions (green, X is homocysteine, Z is penicillamine).

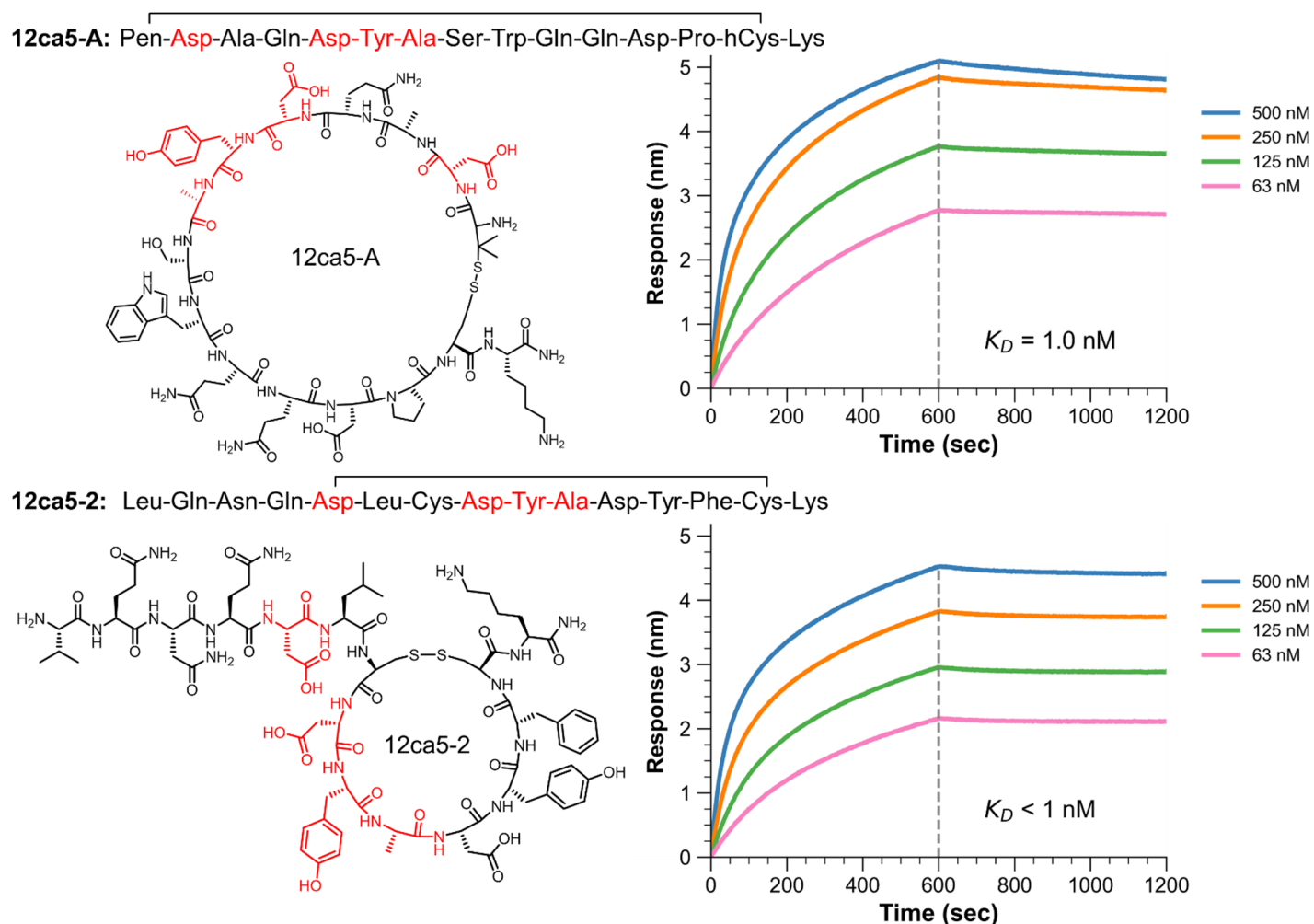


Figure 4. High-affinity macrocyclic peptide ligands to 12ca5 were enriched and identified via affinity selection-mass spectrometry (AS-MS), and the binding affinity was confirmed and measured using biolayer interferometry (BLI). All peptides were prepared with Lys(Biotin)-aminohexanoic acid (Ahx) attached to the N-terminus of the shown sequence and immobilized onto the BLI tip. The BLI tip was then dipped into solutions containing varying concentrations of cadherin-2 to record the concentration-dependent association and dissociation events. The characteristic 12ca5-binding motif D**DY(A/S) is highlighted in red and appeared exclusively at a single position in the X₆CX₆CK library (7 discovered peptides). In comparison, only one motif-containing peptide was discovered from the CX₁₂CK library.

Macrocyclic low-nanomolar peptide ligands were discovered by affinity selection-mass spectrometry (AS-MS) performed against 12ca5 as a model protein target. The anti-hemagglutinin protein 12ca5 binds peptides containing the sequence D**DY(A/S) and has been used to benchmark AS-MS libraries.^{33,58–60} Seven high affinity peptide ligands were pulled down from the X₆CX₆CK library design, while only one peptide was from the CX₁₂CK library (see Table S2 for all identified sequences). A select number of these sequences were synthesized and validated (see Figure S3) for their binding affinity using biolayer interferometry (BLI). All identified binders exhibited apparent dissociation constants (K_D) in the single-digit nanomolar to estimated high picomolar range, nearing the lower limit of detection for the instrument (see Figures 4 and S4). The binding motif was present in the same position or frameshift in all sequences found from the X₆CX₆CK library. Specifically, the cysteine analog in the middle of the library design was located inside of the 12ca5 motif at the

third position (i.e., D*ΨDY(A/S), where Ψ was discovered to be cysteine, homocysteine, or penicillamine). This outcome could potentially be explained by macrocyclization mimicking the geometry of the proline present at the same position within the natural substrate (HA peptide, YPYDVDPYA), leading to a stronger affinity. Moreover, this trend also suggests the X₆CX₆CK library design is more amenable to enrich high affinity binders against 12ca5 relative to the CX₁₂CK library.

Nanomolar affinity binders to cadherin-2 (CDH2) were discovered by AS-MS with the macrocyclic libraries. Due to the critical roles of CDH2 in adhesion in neural and cardiac junctions, the discovery of ligands outside of the protein homodimerization site could be of importance toward its study without affecting biological function. The homodimerization of cadherin-2 is largely driven by molecular interactions involving domains 1 and 2 of the ectodomain.^{61,62,64,65} Based on these considerations, a fusion protein construct comprised of domains 4 and 5 of the CDH2 ectodomain (residues 498 to 724, Uniprot P15116) was used as the target for AS-MS selections. The fusion protein was cloned and expressed in mammalian cells, purified using SEC and Anti-Protein-C (clone HPC4) affinity tag purification, and verified by analytical SEC, reducing and non-reducing SDS-PAGE gels, as well as Western blotting (Figure S5). To our knowledge, no peptide or small molecular ligands have been reported to these CDH2 domains outside the homodimerization site.^{64,65}

Nanomolar peptide ligands were discovered by AS-MS with both X₆CX₆CK and CX₁₂CK libraries against the CDH2[498-724] fusion protein. Only one peptide with high sequencing confidence was enriched against CDH2 (KMTFLFCNFTYKDZK, called cadherin-binding peptide, **CBP**, where Z is penicillamine disulfide bonded to C). **CBP** was synthesized and tested for its binding affinity to CDH2 by BLI, yielding a 53 nM K_D value (Figure 5). To verify sequence binding specificity, a scramble sequence that preserved the size of the macrocycle was synthesized and shown to have negligible binding response to CDH2 by BLI (Figure S6). The linearized form of **CBP** was also tested and demonstrated a greatly-reduced binding response (0.2 nm versus 1.6 nm response for the macrocycle), with an affinity of K_D of 150 nM (Figures S7 and S8). The more rigid structure of the macrocyclic **CBP** thus appears to favor higher affinity.

Structure-activity relationships (SAR) were delineated to characterize **CBP** using single residue replacement studies (alanine and D-amino acid scans) and truncations. In all SAR studies the Cys7-Pen14 disulfide bond was maintained to provide a consistent macrocycle structure to optimize from. First, an alanine scan was performed by synthesis of 13 variants featuring individual alanine mutations, which were assayed by BLI against CDH2 (Table S3, Figure S9). This alanine scan revealed multiple residues to be important for binding (hot-spots) including Lys4, Phe6, Lys12, and Lys15, due to the complete ablation of binding to CDH2 observed in BLI assays (Figure S10). Other residues including Met2, Thr3, Leu5, Thr10, Tyr11, and Asp13 had no effect on binding (cold-spots), suggesting they are not drivers of **CBP** binding to CDH2. Second, a truncation study focused at shortening **CBP** from the N-terminus, producing five additional peptides for BLI testing (Table S4 and Figure S11). BLI assays with these peptides confirmed the impact of the N-terminal residues on binding affinity, especially Phe6 as well as Leu5, while Met2 and Thr3 contributed minimally to CDH2 binding (Figure S12). Third, a D-amino acid scan of **CBP** was performed by iteratively replacing L-amino acids with D-amino acids to determine the impact of stereochemistry on the ligand interactions.^{73,74} Notable hot-spots identified from the D-amino acid scan were Phe6 and Tyr11, further reinforcing the importance of the aromatic residues for binding (see Table S5 and Figures S13-S14). In summary, these initial SAR studies outline the hot-spot residues that appear to drive the high affinity binding of **CBP** to CDH2, including Lys4, Phe6, Lys12, and Lys15, while Met2, Thr3, Leu5, Thr10, and Asp13 are designated as cold-spots with a minimal effect on binding.

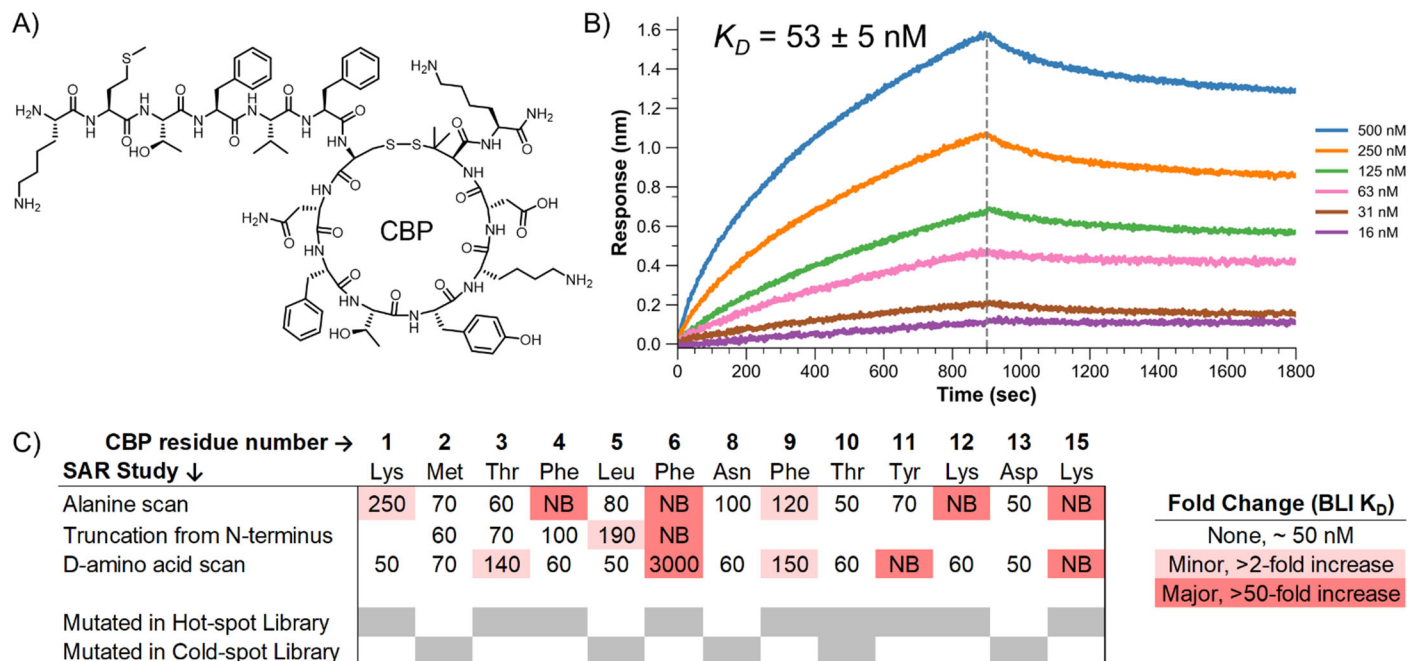


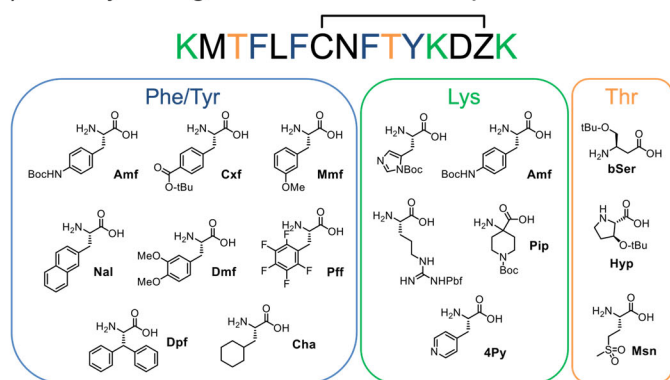
Figure 5. Macrocyclic peptide libraries enabled discovery of a 53 nM peptide ligand to a portion of the ectodomain of cadherin-2. (A) Structure of CBP. (B) BLI experiment reports the affinity of CBP to CDH2 with $K_D = 53$ nM binding affinity. All peptides were prepared with Lys(Biotin)-Ahx attached to the N-terminus in addition to the sequence shown and immobilized onto the BLI tip. The BLI tip was then dipped into solutions containing varying concentrations of CDH2 to record the concentration-dependent association and dissociation events. (C) Summary of experimental (alanine scan, D-amino acid scan, and N-terminal truncation study) SAR data. “NB” denotes non-binding. This SAR information was used to inform the designation of CBP “hot-spots” and “cold-spots,” which do or do not drive high affinity binding, respectively.

SAR data informed the design of two focused libraries based on **CBP**: one to derivatize the high-affinity hot-spot residues, and the other to derivatize the cold-spot residues non-essential for binding. A common approach to design noncanonical libraries is to diversify the hot-spots.^{75,76} However, peptide development and optimization often also considers the cold-spot residues that do not drive high affinity binding to the target. These cold-spot residues can be non-intuitively critical to improving binding affinity, solubility, or proteolytic stability.^{42,43} Thus, we chose to compare the results from both maturation strategies. The set of non-canonical amino acids for incorporation in both the hot- and cold-spot focused libraries were selected based on the consensus data provided by the docking, alanine scan, D-amino acid scan, and truncation studies (Figure 6A and 6B). These libraries were synthesized and subjected to validation by SEC as shown in Figure S15 to demonstrate the lack of apparent oligomerization after disulfide bond formation.

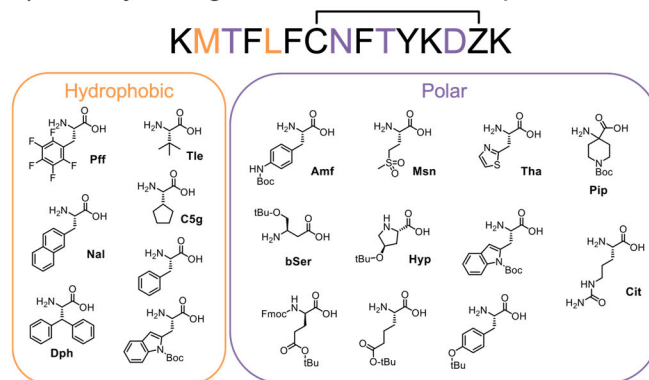
No high-affinity ligands were discovered from the hot-spot focused library by AS-MS, while the cold-spot library provided ten new high-affinity noncanonical macrocyclic cadherin-2 peptide binders (**NCBPs**). From the hot-spot library, only two candidates were identified with high sequencing confidence, which featured multiple mutations from the original **CBP** sequence and shared replacements including Thr10Msn, Tyr11Dph, and Lys15Arg. However, the two hot-spot candidates (**NCBP-1** and **NCBP-2**) were synthesized and tested by BLI, revealing that they were non-binders to CDH2 under these conditions (Figures S16-S17). From the cold-spot library, ten noncanonical putative binders with high sequencing confidence were discovered and synthesized (Table S6, Figure S16). All discovered sequences from the cold-spot library (**NCBP-3** to **NCBP-12**) were high-affinity binders to CDH2 in the BLI experiments, with determined K_D values between 20 and 50 nM (Figure S17).

The significant improvement in binding affinities observed by BLI upon derivatization of the cold-spot residues support this strategy as an efficient avenue for further optimization.

A) Library Design around **CBP** Hotspots:



B) Library Design around **CBP** Coldspots:



C) **NCBP-4**: Lys-Nal-Pip-Phe-C5g-Phe-Cys-Hyp-Phe-Tyr-Tyr-Lys-Asp-Pen-Lys

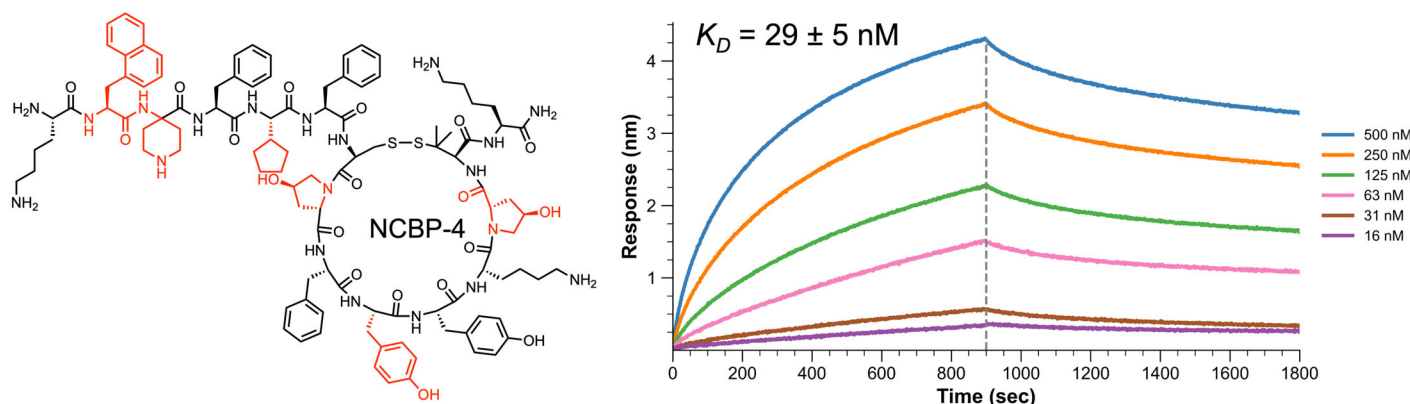


Figure 6. Single-peptide SAR information informs combinatorial library design and affinity maturation with noncanonical amino acids. (A and B) From the single peptide SAR studies summarized in Figure 5C, two libraries were designed to perform affinity maturation. The first library focused on minimally derivatizing the hot-spots by matching the original natural amino acids properties (e.g., hydrophobicity or positive charge). The second library focused around diversifying the cold-spots to examine the possibility that the cold-spots could be mutated to improve the overall binding of the CBP peptide, either by pre-arranging the conformation of the peptide or facilitating new binding interactions with CDH2. Both libraries were prepared using split-pool synthesis, except the entirety of the theoretical sequence space was sampled by the library due to its smaller focused design. Specifically, the number of beads used in split-pool synthesis approximately matched the theoretical diversity: Hot-spot library total number of beads: 2.5×10^6 with theoretical sequence space diversity: 2.7×10^6 and cold-spot library total number of beads: 7.0×10^5 with theoretical sequence space diversity: 7.2×10^5 . (C) Sequence and structure of **NCBP-4** discovered from affinity selection and its BLI binding response, which exhibited high-affinity binding ($K_D = 29 \pm 5$ nM). Nal = 3-(2-naphthyl)-L-alanine, Pip = 4-aminopiperidine-4-carboxylic acid, C5g = cyclopentylglycine, and Hyp = L-trans-4-hydroxyproline.

The **NCBP-4** noncanonical binder exhibits nanomolar binding affinity to CDH2 ($K_D = 29 \pm 5$ nM). The results from AS-MS of the cold-spot macrocyclic library show most cold-spot amino acids were replaced in the identified sequences (Table S6). In all candidates, Met2 was replaced by a hydrophobic amino acid, with 3-(2-

naphthyl)-L-alanine (Nal) appearing in six of the ten. Similarly, Leu5 was replaced by Phe, Nal, cyclopentylglycine (C5g), or pentafluoro-L-phenylalanine (Pff). For the polar subset of amino acids in the cold-spot library, the replacements made to **CBP** were more mixed. Asn8 and Thr10 were replaced with a diverse set of amino acids, possibly indicating their lack of contribution to the binding interaction. Thr3 was replaced by cationic 4-aminopiperidine-4-carboxylic acid (Pip) and polar L- β -Homoserine (bSer). And lastly, Asp13 demonstrated a preferred replacement to trans-4-hydroxyproline (Hyp), appearing in five of ten candidates. With these results in mind, **NCBP-4** was chosen for detailed investigation by BLI as it featured a consensus of amino acid replacements including Met2Nph, Thr3Pip, Leu5C5g, and Asp13Hyp (Figure S18). **NCBP-4** exhibited clear concentration-dependent binding to CDH2 and a resulting binding affinity of $K_D = 29 \pm 5$ nM determined by BLI (Figure 6). Moreover, **NCBP-4** demonstrated a stronger response illustrated by a higher BLI signal (~ 4.3 nm), more than double the response seen with **CBP** (~ 1.6 nm, Figure S19).

A specific mutant of **NCBP-4** was constructed with serine mutated at the hot-spots to specifically examine if the cold-spot residues were effective in creating new binding interactions with CDH2 (sequence: Ser-Nal-Pip-Ser-C5g-Ser-Cys-Hyp-Ser-Tyr-Ser-Ser-Hyp-Pen-Ser-NH₂). Interestingly, this mutant retained moderate binding to CDH2 with an apparent K_D of ~ 150 nM (Figure S20). This serine-substituted mutant was also tested against 12ca5 to assess non-specific binding. The mutant showed no affinity towards 12ca5 (Figure S19). This result suggests the amino acid replacements made in the cold-spots of **CBP** to deliver the **NCBP** compound series resulted in peptide ligands that maintain target specificity despite the presence of additional hydrophobic residues, which are commonly associated with non-specific interactions.^{77,78}

4 Conclusions

We established a protocol for the split-and-pool synthesis of high-diversity macrocyclic peptide libraries and demonstrated their use in the discovery of nanomolar ligands against two protein targets of different structure. Formation of the macrocycle is performed using a simple disulfide bond, which is rapidly installed in aqueous solution using iodine. Quantification of free thiol content via Ellman's assay in both the cyclized and linearized forms confirmed the complete conversion of thiols to disulfides in the library, while SEC revealed that the disulfide bonds were formed exclusively intramolecularly without oligomerization. While the disulfide bond is not the most robust linkage for cyclization, there are several approaches available to replace it when needed to improve stability toward therapeutic development.^{11,79,80}

Affinity selection was able to identify protein-specific ligands from these synthetic macrocyclic libraries for both a model protein (12ca5) and a novel target, cadherin-2, which participates in basic biological adhesion of cells in neural and cardiometabolic function. Both the canonical **CBP** and non-canonical **NCBP-4** peptides demonstrated concentration-dependent binding to CDH2, binding specificity, and high affinity with K_D values of 50 and 29 nM, respectively. After examining the SAR of **CBP**, several hot-spot residues were revealed to be critical for binding to CDH2, featuring several hydrophobic and cationic residues.

From the SAR studies, two additional macrocyclic libraries containing a diverse set of noncanonical amino acids were synthesized focusing on the affinity-driving hot-spots and the non-essential cold-spots, respectively. The subsequent affinity selection experiments investigated the hypotheses of whether the hot-spots can be further refined or if the cold-spots can become meaningful contributors to the binding affinity upon maturation with focused library designs. Overall, AS-MS utilizing the hot-spot CDH2-focused library did not provide any binders or improvement to the original **CBP** peptide. However, AS-MS experiments utilizing the cold-spot library were able to provide several candidates individually validated to be high-affinity binders. Of these, **NCBP-4** was examined more closely for its high affinity ($K_D \sim 29$ nM), specificity, and specific side-chain contributions demonstrated by the amino acids that replaced the original **CBP** cold-spot residues.

Utilizing noncanonical amino acids in combinatorically prepared macrocyclic libraries, we demonstrated the rapid affinity maturation of **CBP**. This process was successful through the replacement of cold-spot residues

with noncanonical monomers. Overall, due to the improvements that macrocyclization often offer over linear peptide scaffolds, we expect this work to be fundamental to the impactful deployment of macrocyclic synthetic libraries for the advancement of peptide therapeutic discovery and development.

5 Data Availability Statement

The data supporting the findings of this study are available in the main text and Supporting Information.

6 Author Contributions

All authors designed research. Michael A. Lee, Joseph S. Brown and Charlotte E. Farquhar performed experimental work. All authors performed data analysis. Michael A. Lee and Joseph S. Brown wrote the manuscript with input from all authors. Bradley L. Pentelute and Andrei Loas supervised the work.

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8 Conflict of Interest Statement

The authors declare the following competing interests: Bradley L. Pentelute is a co-founder and/or member of the scientific advisory board of several companies focusing on the development of protein and peptide therapeutics.

9 Methods

9.1 Rapid oxidation of peptide thiols for intramolecular cyclization using iodine

After cleavage and lyophilization, each split-pool prepared peptide library (e.g., 10 mg, 4.2 μmol , average molecular weight ~ 2400 g/mol) was resuspended at 2 mg/mL in 5% AcN in Water (0.1% TFA) and treated with 10 μL portions of freshly prepared 60 mM I_2 in MeOH until the solution remained yellow. For the example 10 mg scale, approximately 70 μL I_2 stock solution total was used, resulting in ~ 1 eq of I_2 with respect to the library. The iodine-treated library was incubated for 10 minutes in the dark at rt, upon which 15 μL of freshly prepared 1 M ascorbic acid was added (~ 3.6 eq of ascorbic acid with respect to the library example). This solution was immediately loaded onto a pre-equilibrated SPE column, SPE-purified to remove any remaining iodine and ascorbic acid, and lyophilized. The lyophilized powder was then resuspended at approximately 0.1 mM and its thiol concentration as quantified by Ellman's assay.

9.2 Reductive linearization of cyclized library

Peptide libraries were resuspended at 0.62 mg/mL to mimic the maximum concentration possibly isolated at the end of AS-MS, due to the maximum capacity of the STAGE tip (8 μg for a double plug, using 13 μL). As described in the Main Text, reduction after resuspension at pH 3 from 0.1% formic acid in ultrapure water was successful using 1,4-DL-dithiothreitol (DTT, Chem-Impex Cat: 00127). The reduction was also tested in 200 mM sodium phosphate, 5 mM EDTA, pH 8 with DTT, and immobilized tris(2-carboxyethyl)phosphine (TCEP, Thermo Fisher Scientific, 77712). DTT was freshly prepared in a stock solution of 500 mg/mL and added to samples to

provide 50 mg/mL final, 1000 eq and incubated at 60 °C for 15 minutes. Samples were then SPE-purified, lyophilized, and Ellman's quantified upon resuspension. Immobilized TCEP beads (8 mM stock suspension) were washed three times before use with the assay buffer using centrifugation at 1000 rcf for 1 minutes. Treatment of the library peptides with immobilized TCEP used 20 eq for 25 minutes at room temperature rocking on a nutating mixer. The supernatant was isolated from centrifugation, lyophilized, and Ellman's' quantified.

9.3 Sequencing validation of reduced and oxidized libraries

A small portion of library resin was measured and made into a 1 mg/mL stock solution in DMF. Several aliquots of ~1000 beads (for 20 µm resin, this will be about ~10 µL of 1 mg/mL stock) were taken, centrifuged, and aspirated of DMF. Each aliquot was cleaved from the solid phase support using 60 µL of 95% trifluoroacetic acid, 2.5% water and 2.5% triisopropylsilane at 60 °C for 15 minutes. Half of the liquid was then evaporated under a gentle stream of N₂, followed by dilution to a total volume of 240 µL using water (0.1% TFA). A third of the solution was aliquoted to represent the library before cyclization. The remaining solution was cyclized according to the Section 3.5. The cyclized peptide library and aliquoted linear library were both prepared for nLC/MS-MS analysis according to Section 3.4. The dried library samples were then reconstituted in water (0.1% TFA) at a concentration of 100 pg/µL/peptide (for example, prepare 8 µg of an aliquot of 1000 peptides in 80 µL). Half of the cyclized peptide sample was then linearized using DTT as described in Section 3.6. All three types of samples, peptide post-cleavage, cyclized, and cyclized then reduced, were subjected to nLC-MS/MS analysis as described in Section 3.11.

9.4 Ellman's thiol quantification assay

The thiol concentration of suspended peptides was completed using Ellman's reagent (Millipore-Sigma, 5,5'-dithiobis(2-nitrobenzoic acid), D8130, ≥98%, BioReagent) using the following conditions. Ellman's stock solution was prepared at 10.0 mM and assay buffer was 1x PBS at pH 8.0 with 1 mM of EDTA. Nonsterile Greiner 96-well polystyrene plates (Millipore-Sigma, M2936) were used. Using the assay buffer to have a final 200 µL well volume, 3.6 µL of Ellman's stock solution was combined with the peptide solution to give a final peptide intended concentration of 0.1 mM, which was determined to be within the linear regime in which signal could be observed from a standard curve constructed using cysteine (Millipore-Sigma, C7352, ≥98%, BioReagent). After combining, all materials were incubated for 7 minutes, and then read at 416 nm using a TECAN Spark Plate Reader. The concentration of the library was inferred by measuring its absorbance at 280 nm (NanoQuant Plate) and was used to normalize the Ellman's thiol concentration to account for slight variations in the intended resuspended concentrations.

Sample preparation to measure thiol signal from cleavage and SPE of library (reduced library): Approximately 50 mg of peptide library (peptide + resin) was globally deprotected and cleaved from resin with 95% (v/v) TFA, 2.5% (v/v) water, and 2.5% (v/v) triisopropylsilane, for 15 minutes at 60 °C (~20 mL/g of resin). Precipitated peptide was triturated (3 x 100 mL/g resin) with cold diethyl ether, resuspended in 5% acetonitrile in water (0.1% TFA) and solid-phase extracted. After lyophilization, this sample was resuspended in assay buffer at 0.1 mM and measured for its thiol concentration by Ellman's assay.

9.5 Size exclusion chromatography (SEC) of libraries

Size exclusion chromatography (SEC) was performed using an Agilent 1260 Infinity II LC System with a Superdex 30 Increase 10/300 GL column (10 x 300 mm, 9 µm particle size from Cytiva Life Sciences, separation MW range 100 to 7000 Da). 25 µg of library was injected in 200 µL of total solution. Cyclized peptide samples were aliquoted from the main stock prepared according to Sections 3.5 and 3.3. Linearized samples were prepared by adding 1000 eq from a 50 mg/mL DTT stock solution and heating the sample to 60 °C for 10 minutes before dilution to 200 µL using 1x PBS. Column conditions: isocratic 1x PBS for 1.5 column volumes

at 0.8 mL/min. Buffer blanks were prepared for both cyclized and linearized samples and were subtracted from the library samples. A custom mass standard was prepared by adding 10 µg of a mixture of peptides corresponding to the following molecular weights: 1807, 3750, 5312, 8305 for average monomer mass, average dimer mass, average trimer mass, and the exclusion limit of the SEC column.

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