Structure—Activity Relationships of Cbx7 Inhibitors, Including Selectivity Studies against Other Cbx Proteins

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Supporting Information

ABSTRACT: The five human polycomb (Pc) paralog proteins, chromobox homolog (Cbx) 2/4/6/7/8, are a family of chromodomain containing methyllysine reader proteins that are canonical readers of trimethyllysine 27 on histone 3 (H3K27me3). The aberrant expression of the Cbx7 gene is implicated in several cancers including prostate, gastric, thyroid, pancreas, and colon cancer. Previous reports on antagonizing the molecular recognition of Cbx7–H3K27me3 with chemical inhibitors showed an impact on prostate cancer cell lines. We report here on the design, synthesis, and structure—activity relationships of a series of potent peptidomimetic antagonists that were optimized on a trimethyllysine-containing scaffold to target Cbx7. The ligands were characterized using fluorescence polarization (FP) for their binding efficiency and selectivity against the Pc paralog Cbx proteins. The most selective ligand 9, as indicated by the FP data analysis, was further characterized using the isothermal titration calorimetry (ITC). Compound 9 exhibits a 220 nM potency for Cbx7 and exhibits 3.3, 1.8, 7.3 times selective for Cbx7 over Cbx2/4/8 and 28-fold selective over the HP1 family member Cbx1. Our research provides several potent and partially selective inhibitors for Cbx2/4/7 that do not contain trimethyllysine. Our models and binding data suggest that the aromatic cages of Cbx7/Cbx4 can accommodate larger alkyl groups such as diisobutyl substitution on the lysine nitrogen.

INTRODUCTION

Proteins that control the dynamic state of chromatin are critical in the epigenetic regulation of gene expression.1 One of the key mechanisms of the epigenetic control of chromatin is post-translational modifications (PTMs) on histones. Methylation is among the most common PTMs on the histone side chains along with acetylation, phosphorylation, ubiquitylation, and glycosylation.2 Side chains of the lysine residues can be mono-, di-, or trimethylated.2c,3 Methylation of histone residues serves as a chemical switch to recruit effector protein complexes through methyl reader domains. The recruited proteins and multiprotein complexes influence the accessibility of DNA to transcriptional factors, which in turn regulate gene expression. The degree and the site of methylation determine which reader protein is recruited and ultimately the biological outcome, such as the activation or repression of gene expression.1

Methylysine reader proteins have diverse roles in the development, cell-cycle regulation, and oncogenesis.4 Chromodomains are a family of methyllysine reader proteins that include the polycomb (Pc) paralog proteins, chromobox homolog (Cbx) 2/4/6/7/8, that recognize trimethyllysine 27 on histone 3 (H3K27me3).5 A closely related family, the homolog (Cbx) 2/4/6/7/8, that recognize trimethyllysine include the polycomb (Pc) paralog proteins, chromobox domains are a family of methyllysine reader proteins that are canonical readers of trimethyllysine 27 on histone 3 (H3K27me3). The degree and the site of methylation determine which reader

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at the Ink4a/Arf locus. In prostate cancer cell lines, Cbx7 downregulates p16 expression, leading to tumorigenesis. Given the established role of Cbx7 in prostate cancer cells, chemical agents that selectively target the ChD Cbx7–H3K27me3 interaction are proposed to be therapeutically beneficial.

Selectively targeting a single Pc paralog is challenging because of the high degree of sequence homology within the family (Figure 1a). The ligands are bound as the central strand in a three-stranded beta sheet. The binding pockets include an aromatic cage that binds the Kme3 residue, a small hydrophobic pocket that binds the residue, that is, the N-terminal of two residues to the Kme3 (−2) pocket, and a shallow, extended β-groove that accommodates the ligand residues from the (−3) position onward.

The structural similarities and differences among chromodomains vary according to the binding subsites. The aromatic cage residues, Phe 11, Trp 32, and Trp 35, are highly conserved (Figure 1b). The Trp 35 in Cbx2/4/6/7/8 is replaced with the less-electron-rich phenylalanine residue in the HP1 family (not shown). This tryptophan/phenylalanine swap is a possible determinant for selectivity between the two families, in addition to the other structural differences distal to the aromatic cage that have been described. Efforts to make Cbx-selective inhibitors have therefore mainly depended on the (−2) pocket and on the differences in the extended β-groove of the protein. The (−2) pocket is buried under the side-chain-clasping bridge of the hydrophobic residues Val 10/Leu 49. These hydrophobic clasp residues (Val 10/Leu 49) of the Pc group are replaced with polar residues in HP1 (glutamic acid/aspartic acid). This switch influences the selectivity of the side chains of the residues at the (−1) and (−3) positions of the ligand. The floor of the (−2) pocket is defined by Val 13 in Cbx7/Cbx4, which is replaced by an Ala 13 residue that creates a larger (−2) pocket in Cbx2/6/8 (Figure 1c). The HP1 paralogs Cbx1/3/5 also have a Val 13 residue in the corresponding position (not shown). Substitution of a larger group in the (−2) pocket was recently shown to confer good selectivity for Cbx6 over all other Pc and HP1 paralogs. Although significant structural differences exist in the extended β-groove portion of the Cbx proteins, little is known of how this region affects the selectivity among Cbx proteins.

An additional challenge in targeting the Pc proteins is the induced-fit mechanism of the substrate binding. Ligand-free structures for Cbx proteins are very different than the ligand-bound structures for Cbx proteins (not shown). Whereas the H3K27me3-bound structures overlay very well (see Figure 1), the structures of Cbx7 cocrystallized with different inhibitors are more structurally divergent—especially in the ligand-binding aromatic cage region—demonstrating that the proteins can adopt multiple competent binding conformations. Probably for this reason, virtual screens have not yet yielded any chromodomain inhibitors.

Successful approaches to Cbx7 inhibition have been driven by peptide-driven approaches or by high-throughput screening. We first reported on the 200 nM peptidomimetic antagonists of Cbx7, which was followed by the development of a potent (97 nM) cell-active inhibitor UNC3866 by the Frye group. Frye and James recently reported a one-bead-one-compound combinatorial approach that retargeted peptidic compounds toward the CDY family of chromodomains. The group of Zhou published an HTS-derived small-molecule inhibitor for Cbx7 with a 29 μM affinity, and 3-fold and >10-fold selectivity over Cbx4 and Cbx2/6/8, respectively. The authors showed transcriptional derepression of the p16 gene in PC3 cancer cells upon their treatment with the inhibitor MS37452 at 500 μM. The weaker binding of MS37452 to Cbx7 is due to MS37452 occupying only two binding pocket subsites, the aromatic cage and the partly solvent-exposed (−1) pocket of Cbx7. A follow-up report on an extended analog, engaging the hydrophobic (−2) pocket, showed improved affinity.

We report here our own efforts to obtain an improved potency and selectivity for Cbx7. We used the existing peptidic scaffold for inhibitor development because it addresses all binding subpockets and because it has been demonstrated as the most potent cell-active tool against Cbx7. The ligand’s alamine, which engages the (−2) pocket, was maintained because we already showed separately that inhibitors with larger elements in this position lose affinity for Cbx7 and are reprogrammed for the selective inhibition of Cbx6. We focused on two key binding regions that have limited a structure–activity relationship (SAR) established for potency and Cbx paralog selectivity: the aromatic cage and the extended β-groove. On the basis of the binding profiles and structural features of Cbx proteins, we hypothesized that targeting these
regions would create agents with strong selectivity for Cbx7. In the end, we generated new SAR for agents that lean toward different subsets of members within the whole Pc family and that provided good selectivity over its closest relative Cbx4 but did not provide exclusive selectivity for Cbx7.

**RESULTS AND DISCUSSION**

We have established a fluorescence polarization (FP) assay using a panel of Cbx Pc proteins that include Cbx 2/4/6/7/8 and the HP1 paralog Cbx1. These assays were initially developed with the native H3K27me3 ligand, but the low inherent potency of this complex (\(K_d\) values between 6 and 36 \(\mu M\) for different Cbx proteins: Table 1) made studies with

| Chromodomain | \(K_d\) for FITC–H3K27me3 probe (\(\mu M\)) | \(K_d\) for second-generation probe 1 (\(\mu M\)) |
|--------------|-------------------------------------------|-----------------------------------------------|
| Cbx1         | 23 ± 3.6                                  | 5.5 ± 0.54                                    |
| Cbx2         | 5.8 ± 0.2                                 | 0.31 ± 0.03                                   |
| Cbx4         | 19 ± 1.6                                  | 0.21 ± 0.01                                   |
| Cbx6         | 36 ± 4                                    | 0.32 ± 0.03                                   |
| Cbx7         | 8.2 ± 1.1                                 | 0.09 ± 0.10                                   |
| Cbx8         | 30 ± 4                                    | 1.02 ± 0.08                                   |

“Binding data are obtained by titrating varied concentrations of Cbx proteins into a fixed concentration of the FITC-labeled probe. Data are the averages of two or three independent experimental trials and expressed as \(K_d\) ± standard error in \(\mu M\). Solutions contain 20 mM Tris, 250 mM NaCl, 1 mM PMSF, 1 mM benzamidine, and 0.01% Tween, at pH = 8.0. [FITC–H3K27me3] = 500 nM. Stock solution concentrations of Cbx1, 2, 4, 6, 7, and 8 are 450, 280, 800, 450, 175, and 760 \(\mu M\), respectively. [Compound 1] = 100 nM. Stock solution concentrations of Cbx1, 2, 4, 6, 7, and 8 are 500, 500, 390, 500, 450, and 500 \(\mu M\), respectively.

More potent compounds run against the tight-binding limit for the assay for some Cbx proteins, but not for other Cbx proteins.17 To enable competitive FP against the entire panel of proteins, we developed and used a probe (1), that is, a promiscuous binder across the whole test set of proteins (Figures 2 and 3).10

![Figure 2. Structure of probe used for the second-generation competitive FP assays.](image)

Ligand-optimization efforts were initiated based on the core structure (2) (Figure 4). Compound 2 binds to Cbx7 with 0.22 \(\mu M\) potency and exhibited 2.1/3.9/9.1/30-fold selectivity against Cbx2/4/6/8, respectively, and achieved a 36-fold selectivity against HP1 homolog, Cbx1.

Replacements for the N-terminal p-bromobenzamide group of 2 involved a selection of substituted benzoic acids. Tertiary amine replacements of the Kme3 group involved a variety of dialkyl substituents on the lysine nitrogen (Figure 4).

We envisioned that replacements to the N-terminal group of the ligand that engages the extended \(\beta\)-groove would improve the potency and selectivity for Cbx7. We first explored changing the position of bromine on the N-terminal phenyl group followed by substitution with different electron-donating, electron-withdrawing, and hydrophobic groups. Different choices of substituents were made to expand on the existing results by systematically exploring the substituent size, geometry, and chemistry. Synthesis was achieved on the solid phase, by treatment with different benzamides as the end-capping group (Scheme 1a). Selected replacements and binding affinities of the N-terminal group are presented in Table 2. The Supporting Information contains the synthesis and characterization data for all compounds.

The compounds were all tested by a competitive FP assay using probe 1 and recombinant, purified Cbx chromodomains (see the Supporting Information). \(K_{i,0}\) values for the disruption of the Cbx–probe complexes were determined using XLFit and are reported in Tables 2, 3, and 4.

Changing the bromo to the meta/ortho position of the N-terminal phenyl group of 2 reduced the selectivity for Cbx7 and lowered the binding affinity to all Cbx proteins tested. The meta bromo substitution (3) lowered the binding affinity by 21-fold for Cbx7, 5.1-fold for Cbx2, and ~2-fold for Cbx4/6. The ortho bromo substitution (4) was detrimental for Cbx7/Cbx2. Meta or ortho substitutions did not significantly influence binding to Cbx1/8. General potency trends observed for the bromo-substituted compounds to Cbx’s is para > meta > ortho. Replacement of bromine with a methoxy group reduced the binding affinity to all Cbx proteins. All analogs with a methoxy substitution (5, 6, and 7) showed a decreased binding affinity to each Cbx protein, exhibiting the same trend as the bromo compounds (para > meta > ortho). These results indicate that the bromine in the para position is making interactions with Cbx7 more favorably than it does with the other paralogs tested.

We hypothesized that the replacement of the p-bromophenyl group with the p-bromopyridinium group (11) would improve potency either by accepting a hydrogen bond from nearby donors or by tuning the electronics of the halogen.19 In doing so, compound 11 resulted in a 7.3-fold decrease in the potency for Cbx7, 6.6/5.0/4.4/3.5-fold decrease in the potency for Cbx2/4/6/8, respectively, and 1.5-fold decrease in the potency for Cbx1.

Replacement of the para bromo group with an iodo or trifluoromethyl group showed reduced or abolished binding affinity to all Cbx’s. Replacement with iodine (8) showed no binding to Cbx7. Replacement with CF3 (9) resulted in a small loss of potency up to 2-fold for each Cbx protein but showed a slightly improved selectivity for Cbx7 overall. Compound 9 showed a 2.7-fold/9-fold selectivity for Cbx7 over Cbx2/6 and showed 36-fold selectivity over each of Cbx1/8. Compound 9 showed a 5.3-fold selectivity for Cbx7 over Cbx4, which is the best selectivity achieved for any Cbx7 peptide-based antagonist against Cbx4 (the chromodomain that is most structurally similar to Cbx7).11,13c

Substitution with a hydrophobic tert-butyl group (10), as in UNC3866, improved the potency by ~1.2 fold to all Pc group Cbx and showed a 3.1-fold loss to Cbx1. Compound 10 is the most potent compound for Cbx7, is 2.2-, 4.3-, 8.5-, and 28-fold
Figure 3. Binding data of compound 2 against a panel of Cbx proteins. Overlay of FP binding curves of the inhibitor 2 against a panel of Cbx proteins alongside the binding data presented as a bar graph. Gray = Cbx1, purple = Cbx2, yellow = Cbx4, green = Cbx6, blue = Cbx7, and red = Cbx8. Data expressed as IC\textsubscript{50} ± standard errors.

Figure 4. Design of potent and selective Cbx7 inhibitors with N-terminus and Kme3 substitutions on the parent scaffold (2).

Scheme 1. Synthetic Routes to Prepare Peptidomimetics\textsuperscript{a}

\textsuperscript{a}(a) Synthesis of N-terminal modifications of the scaffold and (b) synthesis of N-alkylated peptidomimetics.
selective for Cbx7 over Cbx2/4/6/8, respectively, and is 140-fold selective for Cbx7 over Cbx1.

A variety of quaternary methylammonium group replacements were designed with varied size, hydrophobicity, and basicity to explore binding interactions to the aromatic cages of Cbx proteins (Figure 4). Synthesis of peptides 12−16 was achieved on the solid phase by N-alkylation following the Mtt deprotection of the lysine side chain without cleavage from the resin. A reductive amination was carried out with varying aldehydes added to the free lysine-ε-NH₂ group (see Scheme 1b), and then the synthesis was completed by cleavage and purification as normal. The results for tertiary ammonium side chain replacements are shown in Table 3.

Table 3. IC₅₀ Data of p-Bromobenzamide Replacements of the Scaffold 2 for the Disruption of Cbx7−Compound 1 Complex

| Entry | R = | Cbx1 | Cbx2 | Cbx4 | Cbx6 | Cbx7 | Cbx8 |
|-------|-----|------|------|------|------|------|------|
| 2     | Br  | 7.9 ± 2 | 0.45 ± 0.09 | 0.86 ± 0.2 | 2 ± 0.3 | 0.22 ± 0.7 | 6.8 ± 1 |
| 3     | Br  | 8.3 ± 2 | 2.4 ± 0.03 | 2.1 ± 0.3 | 4.2 ± 0.4 | 4.6 ± 0.6 | 12.4 ± 4 |
| 4     | Br  | NA | >25 | NA | NA | >25 | 11 ± 2 |
| 5     | NA  | 0.83 ± 0.2 | 0.3 | 0.3 | 0.4 | 0.62 ± 0.1 | 9.6 ± 3 |
| 6     | NA  | 11 ± 2 | 1.3 ± 0.2 | 1.9 ± 0.3 | 3.9 ± 0.4 | 0.56 ± 0.1 | 11 ± 2 |
| 7     | NA  | NA | >25 | NA | NA | 14 ± 1 | >50 |
| 8     | NA  | NA | NA | NA | NA | >100 | NA |
| 9     | NA  | 11 ± 2 | 0.80 ± 0.2 | 1.6 ± 0.3 | 2.7 ± 0.3 | 0.3 ± 0.04 | 11 ± 2 |
| 10    | F   | 25 ± 5.6 | 0.39 ± 0.2 | 0.77 ± 0.4 | 1.53 ± 0.4 | 0.18 ± 0.06 | 5.1 ± 1.2 |
| 11    | Me  | 12 ± 3 | 3 ± 0.3 | 4.3 ± 0.4 | 8.8 ± 0.6 | 1.6 ± 0.1 | 24 ± 3 |

“Binding data determined from the inhibitor concentration needed to half-dissociate the Cbx7−compound 1 complex. Data are the averages of two or three independent experimental trials and expressed as IC₅₀ ± standard error in μM. FP data obtained upon titrating varied concentrations of inhibitors into a fixed concentration of Cbx protein and compound 1. Buffer used in the FP experiments is the same as reported in Table 1. The concentration of compound 1 is 100 nM. The concentrations used for Cbx1, 2, 4, 6, 7, and 8 are 10, 1, 1, 1, 0.4, and 7 μM respectively. NA = not applicable, that is, compounds were not tested.

selective for Cbx7 over Cbx2/4/6/8, respectively, and is 140-fold selective for Cbx7 over Cbx1.

A variety of quaternary methylammonium group replacements were designed with varied size, hydrophobicity, and basicity to explore binding interactions to the aromatic cages of Cbx proteins (Figure 4). Synthesis of peptides 12−16 was achieved on the solid phase by N-alkylation following the Mtt deprotection of the lysine side chain without cleavage from the resin. A reductive amination was carried out with varying aldehydes added to the free lysine-ε-NH₂ group (see Scheme 1b), and then the synthesis was completed by cleavage and purification as normal. The results for tertiary ammonium side chain replacements are shown in Table 3.

The dipropyl- and dibenzyl-lysine-containing compounds did not show binding to Cbx7. Compound 15 showed a very weak binding affinity, >1000 μM, to Cbx7. No change in the FP signal was observed during the titration of 14 into Cbx7, which implies that the compound does not bind to Cbx7. Protein−probe aggregation was a problem in these studies because of the poor solubility of the ligands, and this was noted particularly in titrations with Cbx1/6/8.

Of all replacements tested, the diethyllysine compound 12 was the best-tolerated replacement for the trimethyllysine (see Tables 3 and 4). Its binding affinity is less than that of the parent Kme3 compound by 1.5−3.6-fold across the panel. Compound 12 exhibited the strongest binding affinity to Cbx7, but when compared with the parent compound 2, it was less selective for Cbx7 against most of the other Cbx proteins in the panel. The disobutyl compound 13 showed a small decrease in the binding affinity to Cbx7 and a small increase in the binding affinity to Cbx4. Aggregation problems with Cbx1/6/8 prevented the determination of binding affinities and suggested that the solubility was not ideal. Compound 13 was the most potent Cbx4 inhibitor within the series but was not selective over Cbx7.

Combining the most selective Cbx7 CF₃ group with the best-tolerated Kme3 replacement (compound 16) resulted in a potent and partially selective Cbx7 inhibitor. Table 4 shows the
data for compound 16, along with those of its three most closely related parent compounds. Compound 16 registered generally lower potencies when compared with the trimethyllysine-containing parent compound (9) and also with the closely related diethyllysine analog (12). Compound 16 showed 1.5/3.1/5.7/12-fold selectivity for Cbx7 over Cbx2/4/6/8, respectively, and 19-fold selective over Cbx1.

Given the low selectivities (<10-fold within the Pc group) of ligands by FP and variations in the concentration of each Cbx protein tested against the ligands, we sought to validate our FP findings using isothermal titration calorimetry (ITC) for one representative compound from each series, using the same recombinant Cbx constructs. Cbx6 poorly behaved at ITC concentrations and was omitted. We targeted compounds 9 and 16 for ITC titrations, each of which have a hydrophobic CF3 group that leads to strong binding. We could not perform ITC with 16 because at the working concentrations of ligand, we encountered solubility problems in either dissolving the compound or precipitation during the titration of inhibitor into protein. ITC data of compound 9 presented in Table 5 show that compound 9 binds to Cbx7 with a $K_d$ value of 0.22 μM and 3.3-, 1.8-, 7.3-fold selective for Cbx7 over Cbx2/4/8 and 28-fold over Cbx1. Ligand-binding events with each Cbx protein registered favorable negative enthalpies and unfavorable entropies. There are strong signs of enthalpy−entropy compensation when comparing the binding of compound 9 to one protein or another, but we have insufficient structure-function data to comment in detail on the thermodynamics of...
binding across the series. Overall, the binding affinity trends of Cbx paralogs determined using FP and ITC are similar. There is a significant deviation between the IC$_{50}$ and $K_d$ values for Cbx8 and Cbx1. This is because of the weaker binding affinity of these proteins to probe peptide 1, which ultimately leads to differences in the binding affinity determined using FP. Although probe peptide 1 is an improvement over earlier probes, with higher affinities that are relatively similar across the panel of Cbx proteins, these results suggest that FP data comparison in these studies is quantitative only for Cbx proteins.

Table 5. ITC Binding Data for Compound 9 with Different Cbx Proteins$^a$

| protein | $K_d$ ($\mu$M) | $\Delta H$ (kJ mol$^{-1}$) | $-\Delta S$ (kJ mol$^{-1}$) | selectivity |
|---------|----------------|-----------------------------|-----------------------------|-------------|
| Cbx1    | 6.21 ± 0.08    | $-28.4$ ± 0.28              | $-1.32$ ± 0.54              | 28          |
| Cbx2    | 0.73 ± 0.32    | $-21.3$ ± 0.97              | $-14.0$ ± 2.12              | 3.3         |
| Cbx4    | 0.39 ± 0.15    | $-13.4$ ± 0.72              | $-23.4$ ± 1.75              | 1.8         |
| Cbx7    | 0.22 ± 0.02    | $-22.4$ ± 0.01              | $-15.7$ ± 0.62              | 1.8         |
| Cbx8    | 1.61 ± 0.01    | $-6.42$ ± 0.002             | $-26.7$ ± 0.01              | 7.3         |

$^a$Average values are determined by ITC titrations in duplicate at 298 K in buffer, 20 mM Tris, and 250 mM NaCl at pH = 8.

![Table 5. ITC Binding Data for Compound 9 with Different Cbx Proteins](image)

Figure 5. Understanding the binding affinities of inhibitors using molecular models. (a) Chemical structure of UNC3866. Key differences compared with compound 12 are highlighted in pink. (b) Overlay of compound 12 with UNC3866 in Cbx7. Cbx7 is shown in pink, UNC3866 in cyan, and compound 12 in slate blue (pdb code used for the model is 5EPJ). (c) Hydrogen bonding network between UNC3866 and Cbx7. (d) Overlay of para/meta/ortho bromo compounds in Cbx7. (Compounds 2, 3, and 4 are shown in gray white, green, and purple, respectively.) (e,f) Close-up views of the tert-butylbenzamide and diethyllysine portions of the ligands shown in panel d. (g) Two possible meta orientations of the ligand 3. (h) Diisobutyl side chain of the ligand 13 within the aromatic cage of Cbx7 (compound 13 is shown in violet purple). The figure was created using the PyMOL visualization software.
proteins and ligands in which the affinities are not near the tight-binding limit.\textsuperscript{17} The Frye group recently reported the Cbx7 inhibitor UNC3866, which is an analog of compound 2 and is similar to compounds 12 and 16 (Figure 5a). The key differences are methyl ester as the C-terminus, leucine side chain at the \((-1)\) position, and para \(t\)-butyl group at the N-terminus. To fully understand the SARs of our inhibitors, we carried out molecular modeling studies based on the X-ray cocrystal structures of the peptidic ligand Ac-FAYKme3S-NH\(_2\) with Cbx7\textsuperscript{11} as well as UNC3866 with Cbx2/4/7/8.\textsuperscript{13} The all-atom MAB force field as implemented in MOLOC was used for energy minimizations and qualitative comparisons.\textsuperscript{19} The cocrystal structures revealed that UNC3866 makes similar interactions with the chromodomains of Cbx2/4/7/8 at the N-terminus, diethyllysine side chain, and the C-terminus portions of the ligand. The energy-minimized MOLOC model of the diethyllysine-containing compound, 12, was overlaid with UNC3866 in Cbx7, as shown in Figure 5b, and the model suggests that the backbones are highly aligned. Similar to that of Ac-FAYKme3S-NH\(_2\), the backbone of UNC3866 adopts a beta-strand conformation and is participating in an extensive backbone-to-backbone anti-parallel \(\beta\)-sheet-like hydrogen-bonding network with the strands of Cbx7 on either side (Figure 5c). The N-terminal tert-butylbenzamide portion of the ligand is partly embedded in a solvent-exposed cleft, which is formed by the Asp 50, Arg 52, and Leu 53 residues of the protein (Figure 5d). The shallow cleft is defined by one wall that is formed by a flat salt bridge between the side chains of Asp 50 and Arg 52. We conclude from the increasing potency in going from MeO to Br to \(t\)-Bu (5 to 2 to 10), and also from the comparison of benzene versus pyridine (2 vs 11), that the hydrophobic interactions are of main importance for this binding element. The charged Arg–Asp salt bridge would seem an unlikely platform for the hydrophobic interactions, but in fact the faces of such salt bridges have been demonstrated to be key hydrophobic-binding elements in model systems\textsuperscript{20} and in protein structural surveys.\textsuperscript{21} The motif D-L-R in the extended \(\beta\)-groove of Pc proteins interacts with the tert-butylbenzamide group and is replaced by C-E-D-L in the HP1 group. This swap abolishes the salt bridge present in the Pc proteins that contributes to the selectivity of inhibitors for Pc proteins against HP1 proteins.

The binding affinity results show that the \(para\) position is in the best orientation to be embedded in the extended \(\beta\)-groove of Cbx proteins. The models suggest that this simply minimizes steric repulsions that are introduced by substitutions at the \(ortho/meta\) positions (Figure 5d–f). This explains higher potencies to all Cbx5 for \(para\)-substituted compounds compared with those of the \(ortho/meta\)-substituted analogs. The only exception to this rule in the whole data set is the \(meta\)-substituted compound 6, which binds 1.6-fold stronger to Cbx2 than its \(para\)-substituted counterpart 5. The models show that one binding orientation for the \(meta\)-substituent in compound 3 could be directed away from the steric clashes with the cleft (Figure 5g), and we propose that this mode is more accessible for Cbx2 than for the other Cbx proteins.

The diethyllysine side chain is tolerated similarly in the aromatic cages of all Cbx proteins. The diethyllysine side chain of UNC3866 was first modeled and found to be favoring \(gauche\) conformation about the \(\text{C}_\gamma-C_\delta\) bond when the ligand is bound to the Cbx7/Cbx8 proteins. The high-resolution cocrystal structures\textsuperscript{3c} confirmed the same conformation as observed with our ligands, as illustrated in Figure 5b. In UNC3866, one of the ethyl groups on the lysine \(\epsilon\)-NH\(_2\) group interacts favorably with the aromatic pocket residues of Cbx, whereas the other ethyl group is coming out of the pocket. In comparison to adapting \(\text{C}_\gamma-C_\delta\) bond in anti conformation, adapting \(gauche\) conformation about the \(\text{C}_\gamma-C_\delta\) bond shortens the side chain, which helps to minimize steric clashes in the aromatic cage. Accordingly, all diethyllysine peptide analogs in Table 3 maintain moderate potency to all Cbx proteins. Larger alkyl chains such as propyl (15), isobutyl (13), and benzyl groups (14) on lysine have comparatively more steric hindrance, explaining the reduction in binding affinity. The basicity of lysine nitrogen decreases in the order diethyllysine < dipropyllysine < disobutyllysine < dibenzyllysine, which also negatively influences binding affinity in addition to the steric hindrance factor. Unexpectedly, the disobutyl lysine containing compound 13 still holds some potency against Cbx7 and became relatively more potent for Cbx4. This can be explained by an energy-minimized model, which indicates that the binding affinity mostly comes from the hydrophobic interactions with one of the disobutyl side chains of compound 13, and the loss in binding affinity is explained by the ammonium ion coming out of the aromatic cage because of the change in torsion angle of the lysine C-alpha (Figure 5h). The overall conclusion for this series of compounds is that the aromatic cage cannot be used as a source of selectivity among the Cbx paralogs.

## CONCLUSIONS

We report on the inhibitors built from a lead scaffold (2) and selectivity studies against the five Pc paralog proteins. With a goal of achieving potency and selectivity for Cbx7 over Cbx2/4/6/8 and Cbx1, we synthesized two families of inhibitors by altering the N-terminal and Kme3 groups of the scaffold. Potent submicromolar Cbx7 inhibitors with partial selectivity were synthesized. ITC data of compound 9 reveal that the compound binds to Cbx7 with a \(K_d\) value of 220 nM, with a \(\sim 2–7\)-fold of selectivity for Cbx7 over other Pc paralogs and >25-fold selectivity over HP1 paralog, Cbx1. The ITC data of compound 9 also reveal binding affinity trends that are in line with FP findings but also suggest some caution in interpreting selectivities obtained from the FP data quantitatively, especially where the probe peptide binds more weakly to its target (e.g., Cbx1 and Cbx8).

In spite of these limitations, some clear conclusions can be drawn from the FP data. Compounds 10 and 9 are \(>4\)-fold selective for Cbx7 over its closest relative Cbx4. The same two compounds show 2–5-fold selectivity across the series of human Pc paralogs and are very selective for Cbx7 over the HP1 paralog Cbx1 (>30-fold and >100-fold, respectively). In addition to tuning the selectivity toward Cbx7, this series of studies showed some substitutions that tip the selectivity away from Cbx7 and toward other Cbx proteins. Compound 10 possesses a \(t\)-butyl group that achieves the highest potency (390 nM) for Cbx2. Compound 13 contains a disobutyl-substituted lysine side chain and was the most potent Cbx4 inhibitor (490 nM).

FP is a powerful and convenient method for high-throughput screening and for detailed studies on the binding affinity across a panel of proteins. The tight-binding limit for FP probes is not a problem for the weaker-binding hits, but the advent of this entire class of submicromolar Cbx inhibitors requires the creation of even more potent FP probe ligands that are highly potent across all Cbx proteins. Because the tight-binding limit
also applies to many other binding assays, such potent, promiscuous probes would be useful for a variety of Cbx-targeted screening techniques.

This study did not produce a single compound that can exclusively bind Cbx7, but it does provide a set of compounds with different patterns of affinities in this important but scarcely explored family of targets. The inhibition of methyllysine reader proteins is in its infancy, but this can be considered analogous to the development of kinase inhibitors—almost none of them are exclusively selective for a single target, and yet their powerful biological activities arise from their patterns of selectivity in interesting ways. We will report on the cell-based studies of these inhibitors in due course.

■ EXPERIMENTAL SECTION

Materials. All N-Fmoc-protected natural and non-natural amino acids and Rink amide resin were purchased from Chem-Impex International. Fmoc-l-Lys(Me)2-Oh was purchased from GL Biochem. Substituted benzoic acids used for making N-terminal modifications were purchased from Sigma-Aldrich or Matrix Scientific. Aldehydes used for the synthesis of N-alkylated lysine peptides were purchased from Sigma-Aldrich.

Peptide Synthesis. Peptide synthesis was carried out using Fmoc-solid-phase peptide synthesis protocols on Rink amide resin on a CEM Liberty 1 microwave peptide synthesizer. Fmoc-deprotection, addition of Fmoc-amino acid, coupling and cleavage and purification of the peptides on a CEM Liberty 1 microwave peptide synthesizer. Compounds 12–16 were purified using liquid chromatography–mass spectrometry (LC–MS).

Synthesis of N-Terminal Modification Peptides. Substituted benzoic acids were added to the N-terminal of the peptides on a CEM Liberty 1 microwave peptide synthesizer. Cleavage and purification were carried out as outlined in the above-mentioned general procedure for the peptide synthesis. Compounds 2–11 were synthesized in this fashion.

Synthesis of N-Alkylated Lysine Peptides. Syntheses of compounds 12–16 were achieved using the general procedure mentioned above. Lysine side-chain functionalization was carried out using a side-chain MtT-protected lysine. Selective N-deprotection of lysine Mt group on resin was achieved by stirring with a solution of 2% TFA/2% triisopropylsilane in dichloromethane (DCM) (10 mL, 30 min × 2), followed by stirring with a solution of 3% TFA/3% triisopropylsilane in DCM (10 mL, 25 min), and finally by stirring with a solution of 5% TFA/5% triisopropylsilane in DCM (10 mL, 15 min × 2). The resin was washed with CH2Cl2 (10 mL × 3) after each treatment. R–CHO (10–100 equiv), sodium cyanoborohydride (NaBH3CN) (10 equiv), glacial acetic acid (1 drop), and DMF (10 mL) were added to the peptide on resin and stirred at room temperature overnight. The resin was flushed and washed with DMF (10 mL, 3 times). Again, R–CHO (10–100 equiv), NaBH3CN (10 equiv), glacial acetic acid (1 drop), and DMF (10 mL) were added to the peptide on resin and stirred at room temperature overnight. The resin was washed with DMF (10 mL × 3) and then with CH2Cl2 (10 mL × 3). Peptide cleavage from resin, purification, and characterization were performed as outlined above.

Protein Expression. BL21 CodonPlus RIL Escherichia coli cells (Stratagene) were transformed using Addgene plasmids 25245 (Cbx1), 25158 (Cbx2), 25237 (Cbx4), 25296 (Cbx6), 25241 (Cbx7), and a Cbx8 plasmid donated by Arrowsmith (Structural Genomics Consortium, Toronto, Canada). E. coli was grown at 37 °C in 2XYT media to OD ≈ 1.8. Temperature was then dropped to 15 °C for 1 h and then induced with 1 mM IPTG (Life Technologies). Cultures were left shaking overnight and then pelleted, resuspended in binding buffer, and frozen the next day. The chromodomains of Cbx1 (20–73), Cbx2 (8–62), Cbx4 (8–65), Cbx6 (8–65), Cbx7 (8–62), and Cbx8 (8–61) were overexpressed as N-terminal HisX6-tagged proteins.

Protein Purification. E. coli was lysed by incubating on a rocker for 30 min with a CHAPS/dH2O solution (Biobasic), followed by four short (10–15 s) rounds of sonication using a probe-tip sonicator. Centrifugation was subsequently performed to remove cell debris. The remaining supernatant was run through a nickel–nitrilotriacetic acid chelating column (Qiagen) and then through a HiloLoad 16/600 Superdex 75 pg size exclusion column (GE). The FP–LC step performed a buffer exchange into our “minimal” FP buffer consisting of 20 mM Tris HCl, 250 mM NaCl, and 0.01% Tween-20. After concentrating protein to the desired concentration using Amicon centrifugal filter units, phenylmethane sulfonyl fluoride (PMSF), dithiothreitol (DTT), and benzamidine were added to a 1 mM final concentration. Protein purity was assessed using SDS-PAGE gel. Cbx7 and Cbx8 histidine tags were removed after the nickel column step. Cleavage was performed overnight with our in-house TEV protease.

Direct Fluorescence Polarization Assay. Cbx1, 2, 4, 6, 7, and 8 proteins, at maximum concentrations ranging from 120–800 μM, were titrated, in duplicate, into FITC-labeled H3K27me3 or compound 1 at fixed concentrations of 500 and 100 nM, respectively. Both labeled probes were dissolved in distilled water and stored at −20 °C until use. Cbx proteins with a low affinity to the particular probe had to be run at higher concentrations to acquire an accurate Kd value. The assay was performed on 384-well black NUNC plates with optical bottoms. Each well contained a final volume of 50 μL at the time of the read. The buffer used for the assay contained 20
mM Tris—HCl pH 8.0, 250 mM NaCl, 1 mM DTT, 1 mM benzamidine, 1 mM PMSF, and 0.01% Tween. After performing a serial dilution of Cbx proteins from right to left on a plate and leaving one buffer well (column 1 of plate) and one well with no Cbx (column 2 of plate), the plate was incubated in the dark for 15 min before being read. Plates were read with a SpectraMax M5 plate reader (molecular devices) with λ_max 450 nm, λ_exc 530 nm, and an instrument cutoff of 515 nm. The FP read yielded values for parallel (S) and perpendicular (P) light emission.

**Competitive Fluorescence Polarization Assay.** The competitive assay was performed in duplicate using 96-well black plates (NUNC). Compound 1 was dissolved in water and stored at −20 °C until use. The buffer used for the assay contained 20 mM Tris—HCl pH 8.0, 250 mM NaCl, 1 mM DTT, 1 mM benzamidine, 1 mM PMSF, and 0.01% Tween.

Various concentrations of inhibitors dissolved in water (N-alkylated lysine compounds needed up to maximum 10% DMSO to completely dissolve) were titrated into a set concentration of compound 1 (100 nM) + Cbx protein of interest (Cbx 1, 2, 4, 6, 7, and 8). The maximum concentration that the inhibitor was titrated at varied from 1.2 mM to 10 μM. The set concentrations for Cbx proteins were 10 μM (Cbx1), 1 μM (Cbx2, 4, and 6), 0.4 μM (Cbx7), and 7 μM (Cbx8). All wells had a final volume of 100 μL. Plates were incubated in the dark for 15 min after the dilutions. The reads and data analysis were performed in the same manner as for direct FP. Many of the inhibitors required up to 10% DMSO to properly dissolve. Separate competitive FP assays were performed with a known medium strength binder of Cbx7 titrated into compound 1 + Cbx7, with the inhibitor dissolved in various percentages of DMSO. This assay showed that the 10% DMSO would be tolerated without significantly affecting the IC_{50} values (data not shown).

**Fluorescence Polarization Data Analysis.** Microsoft Excel was used to determine the averaged P and S values, minus values for buffer control wells, for each data point. Millipolarization (mP) units were calculated using the formula [(S − P)/(S + P)] × 1000, and the resulting data were plotted with a curve-fitting add-in for Excel (XLFit) using the equation \((C + \beta n_{max} x)/(K_{x} + x))\). In some instances, the data points for the highest inhibitor concentrations had to be excluded where the points had abnormally high mP due to aggregation.

**Isothermal Titration Calorimetry.** Cbx proteins were dialyzed into ITC buffer (20 mM Tris, 250 mM NaCl, pH = 8.0), and the concentrations of proteins were measured at 298 K using MicroCal VP-ITC. The ligand solution (40–60 mM) in syringe was titrated into the Cbx protein (16–48 μM) in a sample cell. Each titration contains 30 injections, with the first injection measuring 2 μL and remaining injections measuring 10 μL each. The duration between two consecutive injections was 240 s. Control titration of heat from ligand dilution was measured from a titration of the ligand solution into the buffer with the above-mentioned instrument settings. Data treatment of titrations to obtain thermodynamic parameters was performed as follows: baseline corrections were performed for both the main titration and control titration data, control data were subtracted from the main data, and data fitting was carried out with a OneSites binding model using the Origin data-fitting software. Thermodynamic parameters presented in Table 5 are the mean values from two individual runs.

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