GS-CA Compounds: First-In-Class HIV-1 Capsid Inhibitors Covering Multiple Grounds

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Recently reported HIV-1 capsid (CA) inhibitors GS-CA1 and GS-6207 (an analog of GS-CA1) are first-in-class compounds with long-acting potential. Reportedly, both compounds have greater potency than currently approved anti-HIV drugs. Due to the limited access to experimental data and the compounds themselves, a detailed mechanism of their inhibition is yet to be delineated. Using crystal structures of capsid-hexamers bound to well-studied capsid inhibitor PF74 and molecular modeling, we predict that GS-CA compounds bind in the pocket that is shared by previously reported CA inhibitors and host factors. Additionally, comparative modeling suggests that GS-CA compounds have unique structural features contributing to interactions with capsid. To test their proposed binding mode, we also report the design of a cyclic peptide combining structural units from GS-CA compounds, host factors, and previously reported capsid inhibitors. This peptide (Pep-1) binds CA-hexamer with a docking score comparable to GS-CA compounds. Affinity determination by MicroScale thermophoresis (MST) assays showed that CA binds Pep-1 with a ~7-fold better affinity than well-studied capsid inhibitor PF74, suggesting that it can be developed as a possible CA inhibitor.

Keywords: human immunodeficiency virus, capsid, assembly, small molecules, inhibitors, disassembly, uncoating

INTRODUCTION

Exceptional developments in combination antiretroviral therapy (cART) have transformed HIV/AIDS from a deadly pandemic to a chronic and manageable disease (Antiretroviral Therapy Cohort, 2017). If administered efficiently, cART significantly reduces morbidity and mortality of HIV-infected individuals, both in resource-rich and in low- and middle-income countries (Quinn, 2008; Sabin, 2013; May et al., 2014; Harries et al., 2016; Teeraananchai et al., 2017). However, emerging drug resistance mutations (DRMs) and the side effects of approved anti-HIV drugs continue to threaten the desired outcome of cART. Hence, current efforts are focused on the discovery of new antivirals acting through novel mechanisms and/or directed to new targets.
HIV-1 CA has two structurally distinct domains, an N-terminal domain (CA-NTD) and a C-terminal domain (CA-CTD), which are connected by a flexible linker of ~5 residues (Figure 1A). The CA-NTD consists of seven α-helices (α1–α7), whereas CA-CTD has four α-helices (α8–α11) and a short 3_10-helix. The structures of CA have revealed interactions between CA protomers in the form of hexameric and pentameric building blocks. The mature capsid core contains ~250 hexamers and 12 pentamers (Pornillos et al., 2009, 2011).

The capsid core is involved at multiple steps of HIV replication. Following fusion of viral and cellular membranes, the capsid core enters the cytosol, where it undergoes controlled disassembly (also known as uncoating). The timing, process, and the extent of uncoating of the capsid core are somewhat controversial. However, published reports implicate that the uncoating is associated with the initiation of reverse transcription (Campbell and Hope, 2015). CA also facilitates nuclear entry and enters the nucleus along with the preintegration complex, suggesting a role of capsid core in integration of viral DNA into the host genome (Schaller et al., 2011; Chen et al., 2016; Francis and Melikyan, 2018). During viral assembly (late stage of viral replication), the Gag polyprotein (precursor of the CA) assembles at the plasma membrane and buds as a spherical, immature, and non-infectious virus. The processing of Gag by the viral protease in several structural proteins and small peptides results in maturation of a conical capsid core.

Given the role of CA at multiple steps of viral replication, it is an attractive antiviral target, offering novel strategies for therapeutic intervention (Prevelige, 2011; Bocanegra et al., 2012; Li et al., 2013). A handful of compounds that bind HIV-1 CA have been reported (Tang et al., 2003; Sticht et al., 2005; Ternois et al., 2005; Kelly et al., 2007; Blair et al., 2010; Curreli et al., 2011; Bocanegra et al., 2012). These include benzodiazepine (BD) and benzimidazole (BM) compounds (Fader et al., 2011; Lemke et al., 2012; Tremblay et al., 2012), pyrrolopyrazolones (BI-1 and BI-2) (Lamorte et al., 2013), CAP-1 (Kelly et al., 2007; Curreli et al., 2011), and PF-3450074 (PF74) (Figure 1). The crystal structures of PF74-bound CA show that PF74 occupies a pocket between the CA-NTD and CA-CTD (Blair et al., 2010; Bhattacharya et al., 2014; Price et al., 2014). BI-2 and host factors cleavage and polyadenylation specific factor 6 (CPSF6) and nucleoporin 153 (NUP153) (Price et al., 2014) also bind in the same pocket. Therefore, both PF74 and BI-2 are expected to interfere with CA binding to CPSF6 or NUP153 (Price et al., 2012, 2014). An antibiotic Coumermycin-A1 (C-A1) has been shown to inhibit HIV-1 integration, and CA mutation A105S conferred resistance to C-A1 (Chen et al., 2016), suggesting the binding of C-A1 with CA. In addition to above-mentioned compounds, peptides (such as NYAD-1) have been reported to inhibit CA function (Tang et al., 2003; Kortagere et al., 2012; Lamorte et al., 2013; Zhang et al., 2013). Despite many promising leads, none of the compounds or peptides has advanced to clinical trials.

Recently reported CA inhibitors GS-CA1 and GS-6207 (an analog of GS-CA1, also called GS-CA2) (Figure 1) have greater potency than currently approved anti-HIV drugs (Blair et al., 2010; Tse et al., 2017; Zheng et al., 2018). GS-CA1 inhibits HIV-1 replication in T cells and peripheral blood mononuclear cells (PBMCs) at very low concentrations (EC_{50} = 240 and 140 pM, respectively). GS-6207 displays anti-HIV activity in MT-4 cells with an EC_{50} of 100 pM, whereas in PBMCs, it displays a mean EC_{50} of 50 pM (20–160 pM) against 23 HIV-1 clinical isolates from different subtypes (Zheng et al., 2018).
In addition, studies in rats and dogs indicate that a single subcutaneous injection maintains GS-CA1 and GS-6207 plasma concentrations above the plasma-binding-adjusted effective concentration required for 95% HIV-1 replication inhibition for >12 weeks, indicating their potential as long-acting drugs (Jarvis, 2017; Tse et al., 2017; Carnes et al., 2018; Sager et al., 2019). Similar to PF74, GS-CA1 inhibits both early and late stages of virus replication.

The crystal structure of GS-CA1-bound CA hexamer has been reported, but it is not publicly available. Reportedly, GS-CA1 binds CA at the same general site as PF74, CPSF6, and NUP153 (Tse et al., 2017). The crystal structure of CA in complex with GS-6207 is yet to be reported. Although not well understood, GS-CA1 and GS-6207 possibly interact more extensively with CA than does PF74, providing greater binding affinity and thereby, greater efficacy than PF74. Here, using computational approaches and reported inhibitor-bound CA structures, we present the details of interactions between GS-CA compounds and CA. We find that GS-CA compounds contain structural features that are also present in PF74, BI-2, NUP153, and CPSF6. Using the same structural features in our computational modeling, we designed a cyclic peptide (Pep-1), which docked at the GS-CA binding site with comparable docking score. We validated the binding of Pep-1 to CA by determining CA binding affinity of Pep-1 using MicroScale thermophoresis (MST) experiments, which revealed that CA binds Pep-1 with ~7-fold better affinity than PF74, a well-known CA inhibitor.

MATERIALS AND METHODS

HIV-1 CA Structure Preparation

The X-ray crystal structure of native HIV-1 capsid protein bound to PF74 (PDB entry 4XFZ) (Gres et al., 2015) was used to dock GS-CA1 and Coumermycin A1 (C-A1). Initial structures of GS-CA1 and C-A1 were generated with ChemSketch (Advanced Chemistry Development, Inc., Toronto, Ontario, Canada). These structures were subsequently minimized using MacroModel followed by LigPrep (Schrodinger Inc. NY). The PrepWizard (Schrodinger Inc. NY), which adds hydrogens, assigns bond orders, creates heteroatom states, and samples conformations of water molecules, was used to prepare CA-hexamer for docking of GS-CA1 and C-A1.

Docking of GS-CA1, GS-6207, and C-A1

All docking simulations were conducted by the Induced-Fit Docking (IFD) module of Schrodinger Suite (Schrodinger Inc., NY). The IFD used Glide (Schrodinger Inc., NY) and the Refinement module in Prime (Schrodinger Inc., NY) to accurately predict ligand binding modes and concomitant structural changes in the receptor. A grid of 36 Å × 36 Å × 36 Å centered on the PF74 in the crystal structure of the native form of CA-hexamer (PDB file 4XFZ) for the docking of GS-CA1, GS-6207, and C-A1 was generated by the Receptor Grid Generation utility of Glide. The IFD optimized the side chain conformation to best determine the docking poses. The pose with the best IFD score was selected for comparison purposes.

Docking of Designed Peptide Pep-1

The structure of peptide Pep-1 was generated by Prime and subjected to energy minimization using the MM/GBSA (Molecular Mechanics–Generalized Born Surface Area) method (Genheden and Ryde, 2010). The docking of the peptide into the crystal structure of CA-hexamer was conducted by IFD (Schrodinger Inc., NY). The best scoring complex of CA/Pep-1 peptide was selected for analyses. We also used PatchDock (Schneidman-Duhovny et al., 2005), though the PatchDock web server1 to assess if the two softwares predicted different docking conformation of Pep-1.

MicroScale Thermophoresis Assays

The binding affinities of CA with Pep-1 and PF74 were determined by measuring thermophoresis of fluorescently labeled CA-hexamers in the presence of increasing Pep-1 or PF74 concentrations. Peptide Pep-1 was synthesized in the Molecular Interaction Core (University of Missouri) and PF74 was purchased from Sigma-Aldrich (St. Louis, MO, USA). Fluorescent labeling of CA with Alexa Fluor 647 analog NT647 was performed according to the manufacturer’s instructions (MO-L004 Monolith Protein Labeling Kit; NanoTemper Technologies GmbH, Munich, Germany). Briefly, 20 μM protein was incubated overnight with 3 M excess of dye at room temperature in a conjugation buffer provided with the labeling kit. The unreacted dye was removed by filtration through a gravity flow column provided with the kit. The elution fractions were collected in 2× MST buffer (40 mM MOPS, pH 7.2, 200 mM NaCl, and 0.2% pluronic F-127). Fluorescence intensity of each fraction was evaluated by MST (Monolith NT.115, NanoTemper Technologies GmbH, Munich, Germany), and fractions containing labeled protein were pooled. Protein concentration was determined by NanoDrop (Thermo Scientific, Waltham, MA) spectrophotometer. Aliquots were stored at –80°C until use. The reaction mixtures containing 200 nM labeled CA-hexamer and increasing concentrations of Pep-1 (1–2,000 nM) were loaded in the capillaries and the thermophoresis was monitored at 20% LED power, high MST power with 20 s MST-on time. The data were analyzed using MO. Affinity software (version 2.3) (NanoTemper Technologies, CA) by fitting the data point to a quadratic equation (Eq. 1) and plotting by Prism (Version 6.0) (GraphPad Inc., La Jolla, CA).

\[ F_{\text{norm}} = \frac{\left( K_d + [CA_0] + [P_0] \right)}{\sqrt{\left( -4[P_0][CA_0] \right)^2 + \left( K_d + [CA_0] + [P_0] \right)^2}} \]

where \( A \) is an arbitrary parameter, \( K_d = [P][CA]/[P - CA] \), \( [P] \) is the concentration of free Pep-1 or PF74, \([CA] \) is the concentration of free CA, \([CA_0] \) is the concentration of added CA, and \( P_0 \) is the concentration of added Pep-1 or PF74.

1https://bioinfo3d.cs.tau.ac.il/PatchDock/
RESULT AND DISCUSSION

Interactions of GS-CA1 and GS-6207 With CA in Modeled CA/GS-CA Complexes

To gain insights into the interactions between the CA hexamer and GS-CA1, we used Induced-Fit Docking (IFD) interfaced with Maestro of Schrodinger Suite (Schrodinger LLC, NY) as detailed in section “Materials and Methods.” A docked pose of GS-CA1 (with best Glide score) in the crystal structure of the native form of CA-hexamer (PDB entry 4XFZ) (Gres et al., 2015) is shown in Figure 2. This figure shows that GS-CA1 binds in the close proximity to residues L56, M66, Q67, N74, and A105 (colored orange in Figures 2B,C). In vitro selection studies have identified GS-CA1 resistance mutations L56I, M66I, Q67H, N74D, and A105E (Perrier et al., 2017), suggesting that these mutations may affect GS-CA1 binding to the CA hexamer. Notably, IFD docking was conducted without any bias toward L56, M66, Q67, L74, or A105. In addition, in our model of the CA/GS-CA1 complex, CA-NTD residues I37, P38, S41, N53, T54, N57, Q63, L69, K70, T106, T107, Y130, Y169, L172, R173, and Q179 also directly interact with GS-CA1 (Figure 2D). Many of these residues are critical to bind small molecules or peptides derived from host factors CPSF6 and NUP153 (Price et al., 2014).

In a limited size cohort (n = 15), the antiviral activity of GS-CA1 was reported to be comparable among clinical isolates from different subtypes (Tse et al., 2017), suggesting a strong conservation of amino acid residues in the GS-CA1 binding pocket. To assess whether the GS-CA1 binding pocket is conserved among subtypes, we generated a consensus sequence of CA from HIV-1 subtype C (HIV-1C), which accounts for more than 50% of all HIV-1 infections, using the Los Alamos HIV sequence database2. The results showed that the GS-CA1 binding site in HIV-1C was highly conserved. We noted only one substitution in HIV-1C (F169) compared to HIV-1B (Y169). The nearest (Ca) atom of Y169 (or F169 in HIV-1C) is within interacting distance of GS-CA1 (< 3.8 Å), suggesting a weak interaction with GS-CA1. The effect of the change from tyrosine to phenylalanine remains to be investigated.

GS-6207 differs from GS-CA1 by three modifications: (1) a cyclopropane moiety on sulfonamide group was replaced by a methyl group, (2) difluoroethyl groups on indazole ring was replaced by a trifluoroethyl group, and (3) difluoromethyl group on cyclopenta-pyrazole ring was replaced by a trifluoromethyl moiety. At present, the specific rationale for these replacements is not known. We docked GS-6207 in the crystal structure of native form of CA (Gres et al., 2015). The results showed that GS-6207 binds in the same binding pocket as GS-CA1 and with a slightly better Glide score (−14.362 for GS-6207 versus −11.271 for GS-CA1), suggesting a better binding affinity. We also noted

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2https://www.hiv.lanl.gov
that the orientation of cyclopenta-pyrazole ring in docked GS-6207 was switched by ~180° compared to that in GS-CA1, leading to the exposure of trifluoromethyl moiety to the solvent (Figure 3A). Another remarkable difference between docked complexes of CA/GS-CA1 and CA/GS-6207 is the conformation of K70 and R173 side chains. In CA/GS-6207 complex, K70 side chain moves around 5 Å from the position in CA/GS-CA1 complex (Figure 3B, solid arrow) toward the binding pocket and forms a hydrogen bond with C=O of amide group in GS-6207 (Figure 3B, dotted line). An additional H-bond may be one of the reasons that GS-6207 has better Glide score than GS-CA1. While the side chain conformation of R173 is also altered (Figure 3B), it does not appear to be significant.

### Comparison With PF74/CA and BI-2/CA Crystal Structures

Five mutations (Q67H, K70R, T107N, L111I, and H87P) confer resistance to PF74 (Blair et al., 2010; Shi et al., 2011, 2015; Zhou et al., 2015). Residues Q67, K70, T107, and L111 reside on helices 4 and 5, whereas H87 is part of the CypA binding loop (residues 85–93) (Gamble et al., 1996; Ambrose and Aiken, 2014). The only common resistance mutation between GS-CA1 and PF74 is Q67H (Perrier et al., 2017), although other GS-CA1 resistance residues (L56, M66, L74, and A105) are also within interacting distance of PF74. A superposition of the CA/PF74 crystal structure (Gres et al., 2015) and the CA/GS-CA1 model is shown in Figure 3C. It is clear from the figure that all three rings of PF74 (two phenyl rings and one indole ring) superpose extremely well on three different rings GS-CA1 (dotted circles 1, 2, and 3 in Figure 3D). The PF74 indole ring superposes on the cyclopenta-pyrazole ring of GS-CA1 (circle 1). One of the two phenyl rings of PF74 superposes on the difluorobenzene ring of GS-CA1 (circle 2), whereas the other PF74 phenyl ring is at a topologically similar position to the indazole ring of GS-CA1 (circle 3). Additionally, the polar moieties of PF74 match topologically with the polar moieties of GS-CA1. Thus, the acetamide moiety of GS-CA1 superposes well on the corresponding moiety of PF74. These data suggest that certain structural features and interactions are common between GS-CA1 and PF74.

During IFD of GS-CA1 into the CA-hexamer, the conformations of most of the side chains in the GS-CA1/PF74 binding pocket did not change significantly as compared to the CA/PF74 crystal structure, with the exception of the side chain of K70 (Figure 3C). The position of the K70 NZ atom was shifted by ~4.7 Å from its position in the CA/PF74 complex (cyan versus gray carbons in Figure 3C), suggesting an absence of interactions between K70 and GS-CA1, in contrast to K70 interactions with PF74. The absence of this interaction is a possible reason that mutation at K70 did not emerge during GS-CA1 in vitro resistance selection studies (Perrier et al., 2017). As mentioned above, the interaction of K70 is restored in the CA/GS-6207 complex. At present, the resistance mutation profile of GS-6207 is not known. Hence, the significance of this interaction awaits virological studies.

BI-2 is one of the two 4, 5-dihydro-1H-pyrrolo[3,4-c]pyrazol-6-one series compounds shown to bind the CA hexamer. BI-2 was shown to stabilize CA hexamers and inhibit HIV-1 at early stages of infection (Lamorte et al., 2013). Selection of viruses resistant to BI-2 identified mutations at residues A105 and
and T107 of CA-NTD (Lamorte et al., 2013). The high resolution structure of CA in complex with BI-2 showed that it binds at the PF74 binding site. The superposition of the three compounds (GS-CA1, PF74, and BI-2) obtained from the superposition of Ca-atoms of CA-NTD showed that the three compounds have a common binding mode with CA-hexamer (Figure 3E). Our docking results of GS-CA1 showed that CA residue A105 is within interacting distance of GS-CA1, and the common resistance mutation A105T between GS-CA1 and BI-2 further confirms that the two compounds share part of the binding site.

**Comparison With CPSF6/CA and NUP153/CA Crystal Structures**

The crystal structures of CA in complex with short peptides derived from CPSF6 and NUP153 showed that both peptides share the binding pocket occupied by PF74 and BI-2 (Price et al., 2014), although the bound peptides had additional interactions. To determine whether common structural features among GS-CA1, CPSF6, and NUP153 exist upon binding to CA, we superimposed the crystal structures of CA/CPSF6 and CA/NUP153 on our modeled CA/GS-CA1 complex. The superposition is shown in Figure 4, demonstrating that the conformation of GS-CA1 docked into the CA-hexamer follows the folding of the CPSF6 peptide (Figure 4A). Remarkably, the side chain of F321 of CPSF6 perfectly superposed on the difluorobenzyl moiety of GS-CA1.

Similar to CPSF6, the NUP153 backbone follows the conformation of GS-CA1, and F1417 of NUP153 perfectly superposes on the difluorobenzyl moiety of GS-CA1 (Figure 4B). In addition, there exists a hydrophobic interaction between the methylsulfonyl moiety of GS-CA1 and P38 of CA (atoms of P38 are not shown). A similar interaction is noted between F1415 of NUP153 and P38 (Figure 4B).

**Comparison With CA/CAP-1, CA/BD, and CA/BM Complexes**

CAP-1 1-(3-chloro-4-methylphenyl)-3-(2-(((5-((dimethylamino)methyl)furan-2-yl)methyl)thio)ethyl)urea is an assembly inhibitor for which the resistance mutation profile has not been reported (Kelly et al., 2007). The structure of CAP-1 bound CA-NTD has been solved by NMR and X-ray crystallography (Kelly et al., 2007). A comparison of the crystal and NMR structures demonstrated that CA undergoes significant conformational change upon CAP-1 binding. The superposition of the crystal structure of the CA/CAP-1 complex on the model structure of the CA/GS-CA1 complex showed that the two inhibitors did not bind at a common site (Figure 5). However, two residues (M66 and L69) interacted with both GS-CA1 and CAP-1. The positions of M66 in the CA/GS-CA1 and CA/CAP-1 complexes are shown in Figure 5. The compounds of the benzodiazepine (BD1–BD4) and benzimidazole (BM1–BM5) series bind to CA at a site that is close to the CAP-1 binding site (Lemke et al., 2012). While compounds from both series have been shown to bind at the same pocket, they have distinct resistance mutation profiles. Mutations V36T and G61E were selected with BD inhibitors, whereas K30R and S33G were selected with BM inhibitors. Both V36 and G61 are part of BM3 binding pocket (PDB entry 4E91) (Sticht et al., 2005). K30 is not within the interacting distance of BM4, and the backbone carbonyl group of S33 only forms a Van der Waals interaction with BM4 (PDB entry 4E92) (Lemke et al., 2012). Hence, the resistance mechanism of BM4 does not seem to operate through direct interactions. The crystal structures of BD3 and BM4 bound to CA-NTD showed that both compounds are within interacting distance of M66, similar to CAP-1. Hence, BD and BM series compounds do not share a binding site with GS-CA1, but they all have a common interaction with M66.

**Coumermycin A1 Binding to CA-Hexamers**

Coumermycin A1 (C-A1) is a gyrase B inhibitor that also inhibits HSP90 (Vozzolo et al., 2010) (reviewed in Carnes et al., 2018). A crystal structure of CA/C-A1 has not been solved. However, docking studies predict the binding of C-A1 in a pocket formed by two adjacent capsid monomers (Chen et al., 2016). This predicted binding site may be relevant, as mutations N74D and A105S conferred resistance to C-A1, and both residues (N74 and A105) are at the interface of two capsid monomers.

![Figure 4](image-url)

**Figure 4** | (A) Superposition of CA/CPSF6 crystal structure on the molecular model of CA/GS-CA1. For clarity, residues P313-P316 and backbone atoms of CPSF6 have been omitted. Peptide CPSF6 is shown in yellow carbons, and the backbone of CPSF6 peptide is shown as a yellow tube. This figure shows the superposition of the difluorobenzyl ring of GS-CA1 on F321 of CPSF6. (B) Superposition of CA/NUP153 crystal structure complex on the molecular model of CA/GS-CA1. For clarity, only residues F1415-G1418 are shown. The carbon atoms and backbone (tube) of NUP153 are shown in faded red-orange color. This figure shows the superposition of the difluorobenzyl ring of GS-CA1 on F1417 of NUP153. In addition, F1415 of NUP153 and the methylsulfonyl group of GS-CA1 have a common interaction with P38 of CA-hexamer. The atoms of P38 are not shown for clarity.
Ebselen is a small molecule that was discovered in a search for inhibitors of CA dimerization. Electrospray ionization mass spectrometry experiments revealed that ebselen covalently binds CA-CTD, most likely through a selenylsulfide linkage involving C198 and C218 (Thenin-Houssier et al., 2016). Both of these residues are part of the CA-CTD, and they are not within interacting distance of the GS-CA1 in our modeled CA/GS-CA1 complex. Therefore, we predict that ebselen and GS-CA1 binding sites do not overlap.

### Docking of a Designed Cyclic Peptide Inhibitor (Pep-1)

Using the crystal structures of PF74, NUP153, CPSF6, and BI-2-bound CA as well as the modeled structure of the CA/GS-CA1 (CA/GS-6207) complex, we designed a cyclic peptide, Pep-1, containing common structural components/groups among CA-bound small molecules or peptides derived from CPSF6 and NUP153. The docking of Pep-1 showed that it binds in a pocket that is shared by PF74, NUP153, CPSF6, GS-CA1, and GS-6207. The structural components that superposed in different complexes are listed in Table 1. It appears that the designed peptide shares binding site and chemical moieties that may inhibit CA function. The structural and chemical details of Pep-1 will be reported elsewhere. However, we determined the binding affinity of CA with Pep-1 and compared with the PF74 binding affinity (presented below).

### Binding Affinity of CA With Pep-1 and PF74

We used the MicroScale thermophoresis (MST) assays to determine the binding affinity of CA to Pep-1 and PF74. MST is based on the thermophoresis, a directed movement of molecules in a temperature gradient, which depends on a variety of molecular properties including size, charge, hydration shell, and conformation. Thus, it is highly sensitive to virtually any change in molecular properties, allowing for precise quantification of molecular events independent of the size or nature of the investigated sample (Jerabek-Willemsen et al., 2014). During the MST experiment, a temperature gradient is induced by an infrared laser. The directed movement of molecules through the temperature gradient is detected and quantified using a covalently attached fluorophore. The binding isotherms obtained by plotting the difference in normalized fluorescence against increasing Pep-1 and PF74 concentration are shown in Figures 6A,B, respectively. The binding affinities of Pep-1 ($K_{d,\text{Pep-1}}$) and PF74 ($K_{d,\text{PF74}}$) with CA were extrapolated by fitting the data points to a quadratic
The binding affinity of CA-hexamers with PF74 was previously determined by isothermal calorimetry (ITC) to be 262 nM (Bhattacharya et al., 2014), which is in good agreement with the $K_d$ determined here using MST. These data suggest that Pep-1 binds CA with ~7-fold greater affinity.

CONCLUSION

The results obtained from flexible docking and comparative structural analyses presented above show that the compounds GS-CA1 and GS-6207 contain structural features that are present in several previously discovered small molecule inhibitors of CA, as well as in CA-interacting host factors. Most importantly, the phenyl moieties of PF74 and BI-2 perfectly superpose on the difluorobenzyl group of GS-CA1 and GS-6207, as do the phenylalanine residues of CPSF6 and NUP153 peptides. The position of the phenyl ring is critical in binding of these compounds, as it forms direct contacts with critical CA residues L56, N57, and M66. A significantly larger size of GS-CA1 and GS-6207 compared to PF74 likely ensures additional contacts with CA. These contacts are either present in CA/CPSF6, CA/NUP153, CA/BD (or CA/BM), or CA/CAP-1 crystal structures. Collectively, it appears that GS-CA compounds have been designed to include structural features of many CA-binding molecules to ensure that they mimic the interactions of many ligands.

Furthermore, we suggest that a cyclic peptide designed based on the structures of small molecules inhibitors and host factors bound CA has strong binding affinity to CA.

DATA AVAILABILITY

The raw data supporting the conclusions of this manuscript will be made available by the authors, without undue reservation, to any qualified researcher.

AUTHOR CONTRIBUTIONS

KS and AS conceived and designed the study. UN conducted sequence analyses, and KS conducted computational analyses. KS, FG, TQ, and UN analyzed the results. KS wrote the original manuscript. KS, KH, ML, TQ, DB, UN, and AS edited the manuscript.

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FIGURE 6 | Pep-1 (A) and PF74 (B) binding affinities of CA as determined by MicroScale thermophoresis. This figure shows the change in fluorescence due to thermophoresis at the increasing concentrations of Pep-1 and PF74 (1–2,000 nM) in the presence of 200 nM CA-hexamers.
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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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