Structure of a Myeloid cell leukemia-1 (Mcl-1) inhibitor bound to drug site 3 of Human Serum Albumin

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Abstract

Amplification of the gene encoding Myeloid cell leukemia-1 (Mcl-1) is one of the most common genetic aberrations in human cancer and is associated with high tumor grade and poor survival. Recently, we reported on the discovery of high affinity Mcl-1 inhibitors that elicit mechanism-based cell activity. These inhibitors are lipophilic and contain an acidic functionality which is a common chemical profile for compounds that bind to albumin in plasma. Indeed, these Mcl-1 inhibitors exhibited reduced \textit{in vitro} cell activity in the presence of serum. Here we describe the structure of a lead Mcl-1 inhibitor when bound to Human Serum Albumin (HSA). Unlike many acidic lipophilic compounds that bind to drug site 1 or 2, we found that this Mcl-1 inhibitor binds predominantly to drug site 3. Site 3 of HSA may be able to accommodate larger, more rigid compounds that do not fit into the smaller drug site 1 or 2. Structural studies of molecules that bind to this third site may provide insight into how some higher molecular weight compounds bind to albumin and could be used to aid in the design of compounds with reduced albumin binding.

1. Introduction

Defects in apoptotic signaling are central to tumor initiation and maintenance of cancer cells, particularly when associated with unregulated tumor cell proliferation.\textsuperscript{1-4} Amplification of the gene encoding the anti-apoptotic protein Myeloid cell leukemia-1 (Mcl-1) is one of the most common genetic aberrations in human cancer.\textsuperscript{5} Mcl-1 over expression in human cancers is associated with high tumor grade and poor survival.\textsuperscript{6} This suggests that inhibition of Mcl-1 could cause tumor cell death in Mcl-1-dependent cancers and cause an increase in sensitivity to many standard chemotherapeutics by releasing apoptotic restrictions.

Mcl-1 mediates its effects primarily through a protein-protein interaction and is difficult to target with small molecules.\textsuperscript{7} Recently, we reported on ligands that bind tightly to Mcl-1 that are lipophilic and contain an acidic functionality.\textsuperscript{8-11} Although these molecules bind tightly to Mcl-1, they also bind to albumin as suggested by their lower \textit{in vitro} cell activity in the presence of 10% FBS and no measurable \textit{in vitro} free fraction. To improve our series’ \textit{in vitro} cell activity, we wanted to design Mcl-1 inhibitors with reduced binding to albumin. This was accomplished in the past when targeting the Bcl-2 family members Bcl-xl/Bcl-2 using a structure-based approach.\textsuperscript{12} By comparing ligand binding to HSA and Bcl-xl, Bcl-2 family inhibitors were designed with a significant decrease in albumin binding. A similar approach was used for inhibitors of Methionine aminopeptidase-2 (MetAP2) which led to compounds with good cellular activity in both proliferation and methionine processing assays.\textsuperscript{13} Many other studies have monitored ligand binding to plasma proteins to discover small molecules with reduced albumin binding.\textsuperscript{14} For example, in the discovery of a series of DPP-IV inhibitors, a correlation between \textit{in vitro} potency, plasma concentration and free fraction was observed\textsuperscript{15} that was used to guide inhibitor design.

Human Serum Albumin (HSA) binds multiple natural ligands in different sites. The endogenous ligands are typically acidic lipophilic compounds like fatty acids or bilirubin. Fatty acids have nine binding sites on albumin, and each are potential binding sites for drugs.\textsuperscript{16,17} Drugs have three major reported binding sites on albumin. Smaller drugs (molecular weight less than 500 Daltons) mostly bind to drug sites 1 or 2 with a few also binding to a third
site near domain IB of albumin.\textsuperscript{17} Larger, rigid, and extended, higher molecular weight ligands typically bind to a third site in domain IB. All three sites share common physical characteristics; a hydrophobic pocket near multiple basic residues, which helps explain the protein's broad tolerance for a diverse group of lipophilic ligands which contain an acidic moiety.

Several compounds have been found that bind to site 3 of albumin.\textsuperscript{10} However, the number of molecules that have been structurally characterized is much lower than to site 1 or 2 making further structural characterization of ligand binding to this site of broad interest. Additional structural data for this site would provide additional data to optimize QSAR models and for docking approaches to predict albumin binding of compounds.\textsuperscript{18-20}

Here we report on the X-ray structure of an Mcl-1 inhibitor when bound to Human Serum Albumin in an effort to understand how to reduce HSA interference for this class of compounds. We were surprised to find that this Mcl-1 inhibitor binds predominantly to HSA site 3. When we compared our structure to other ligands that bind to the same pocket, we observed common binding hot spots and interactions that stabilize complex formation. The structure of this ligand when bound to HSA is consistent with the SAR we obtained on the plasma binding of other members of this chemical series. These data may thus be used to help design new analogs with reduced HSA binding without affecting binding to Mcl-1.

2. Experimental

2.1. Chemistry

Preparation of compound 1 (Fig. 1) was described previously.\textsuperscript{11}

2.2. Purification of Human Serum Albumin

Albumin from human serum lyophilized powder and albumin human recombinant were purchased from Sigma-Aldrich. The protein was further purified by gel-filtration using S75 column chromatography.

HSA ligand free crystals were grown by sitting-drop vapor diffusion using a protein concentration of 100 mg/ml in 50 mM potassium phosphate (pH 7). The HSA complex was generated by soaking the ligand free crystals in mother liquor containing 2 mM compound 1 with 5% DMSO overnight.

2.3. X-ray crystallography

Data were collected on the Life Sciences Collaborative Access Team (LS-CAT) 21-ID-D beamline at the Advanced Photon Source (APS), Argonne National Laboratory. The data were indexed, integrated and scaled with HKL2000. Using a previously determined HSA ligand free structure (PDB: 4K2C) as a starting model, phasing was done by molecular replacement\textsuperscript{21} with Phaser\textsuperscript{22} as implemented in CCP4. Refinement of the structural models were performed with Phenix, and included rounds of manual model building in COOT.\textsuperscript{23} The complex was refined to resolution of 2.7 Å. The final model has Rwork/Rfree 0.17/0.21 and good stereochemistry (Table 2 Supplementary).

2.4. Graphics software

All figures were produced using the PyMOL Molecular Graphics System, version 1.7 (Schrödinger, LLC: New York, 2010).

2.5. Expression and purification of Mcl-1

Mcl-1 (172-318), corresponding to the N-terminal portion of Mcl-1, with an MBP tag was used for the TR-FRET biochemical assays. The design, characterization and sequence of this construct have been described earlier.\textsuperscript{8} Soluble protein was purified by Ni affinity chromatography.

2.6. Caspase activation assay

NCI-H929 cells were washed twice in RPMI 1640 without Fetal Bovine Serum (FBS), then dispensed in 96 well plates at 10,000 cells per ml in RPMI 1640 containing 0.05% Pluronic F-68 non-ionic surfactant (Thermo Fisher Scientific, Waltham MA) and 0, 1 or 10% FBS in white 96 well plates. Compound dose response curves were generated and added to the cells in their respective media. Plates are incubated with compound for 3 h then Caspase-Glo (Promega, USA) reagent is added and incubated at room temperature in the dark, for 30 min. Luminescence is measured on a BioTek Cytation 3 (Winooski VT, USA) and the I_{50} was determined using Graphpad Prism software (San Diego, CA).

2.7. TR-FRET binding experiments

Time resolved fluorescence resonance energy transfer (TR-FRET) based competitive binding assays were carried out in a 20 µL assay mixture in OptiPlate-384, White opaque 384 well MicroPlates (Perkin Elmer, Waltham, MA). The assay mixture was composed of 1 nM recombinant, Mcl-1 fused to maltose binding protein (MBP), 100 nM fluorescein isothiocyanate (FITC) labeled Bak BH3 peptide (FITC-Ahx-GQVGRQLAIIGDDINR-NH2); (GenScript USA Inc, Piscataway, NJ), and 1 nM Terbium (Tb) conjugated antibody directed against MBP (CisBio Bioassays, Bedford, MA) in an assay buffer composed of 4.52 mM monobasic potassium phosphate, 15.48 mM dibasic potassium phosphate, 1 mM sodium EDTA, 0.05% Pluronic F-68 detergent, 50 mM sodium chloride, and 1 mM DTT (pH 7.5). Compounds were added to the reaction mixture in DMSO. The final DMSO concentration was 0.25%. TR-FRET measurements were determined using a Biotek 3 multimode plate reader (Winooski VT, USA) at an excitation wavelength of 340 ± 30 nm, and dual excitation wavelengths of 520 ± 10 nm and 620 ± 10 nm. Delta F was determined by dividing the fluorescence at 520 nm by the fluorescence at 620 nm.

On each plate, wells containing the fluorescent peptide, Mcl-1-MBP protein, Terbium conjugated antibody, and vehicle (DMSO) served as a negative control. Wells containing peptide and antibody, in the absence of Mcl-1 protein served as a positive control for total displacement of peptide. For each compound, a dose response curve was generated by potting delta F values against compound concentration and the I_{50} was determined using a four parameter logistic curve fit with Graphpad prism software. The dissociation constant (K_{i} value) was derived from:\textsuperscript{24}

\[ K_{i} = \frac{\left[ I \right]_{50}/\left[ I \right]_{50}/K_{d} + [P]_{0}/K_{d} + 1) \]

where [I]_{50} is the concentration of the free inhibitor at 50% inhibition, [I]_{50} is the concentration of the free labeled ligand at 50% inhibition, [P]_{0} is the concentration of the free protein at 0% inhibition, K_{d} represents the dissociation constant of the FITC peptide probe. Experiments which determined the effect of albumin or bovine serum were performed in the same manner in assay buffer containing FBS, BSA, or HSA at the indicated concentration.
3. Results and discussion

3.1. Shift in activity due to albumin/serum binding

Compound binding to serum proteins can cause shifts in biochemical and in vitro cell assays making interpretation of structure activity relationships (SAR) complicated. To test the effect of adding Fetal Bovine Serum (FBS) or albumin on the apparent compound activity of compound 1, we performed a plasma shift assay. The plasma shift assay is a common method to ascertain interference from FBS by adding serum to an in vitro assay and then observing a change in the apparent affinity to the target due to compound sequestration by plasma proteins. The plasma shift assay can be quantified to help monitor changes to compound binding to plasma proteins such as albumin. Quantification can be useful (Supplementary Materials) but in most cases the fold shift in the presence and absence of albumin is all that is used to compare compounds in the same chemical series.

For compound 1 in the TR-FRET competition assay, we observed a shift in IC50 and apparent Ki for binding to the target, Mcl-1, of over 3-fold between no FBS and 1% FBS (Fig. 2A). When we repeated this experiment using delipidated HSA or BSA at the approximate concentration of albumin found in FBS (i.e. from FBS suppliers analysis 1% FBS corresponds to 180 µg/ml BSA), and we found a nearly identical shift in apparent Ki (Table 1). This result is consistent with albumin being the major component in FBS that sequesters compound 1 and causes the right shift in the IC50 in the binding assay. We next tested what effect different amounts of FBS had on compound 1 in an in vitro cell based activity assay.

To test the effect of FBS on in vitro cell based activity, we used a caspase activation assay. Rapid caspase activation is a characteristic of on target Bcl-2 family inhibitors and for compound 1, we see robust caspase activation in less than 3 h after dosing. We found that cells are more tolerant to different amounts of FBS in the caspase assay with a readout at 3 h, than in a 24 or 72 h proliferation assay. However, we still carefully normalized all of the data using DMSO as a control to reduce differences in caspase activity due to changes in cell viability and changing serum conditions. As shown in Fig. 2B when we dosed cells with different amount of FBS, we observed more than a fifty fold shift in EC50 in the caspase activation assay between 0% and 10% FBS. The pronounced shift in EC50 in this cell based assay is similar to the effect we observed in the biochemical binding assay (Fig. 2A) and is consistent with the hypothesis that compound binding to serum proteins is what causes an increase in the EC50 needed to see a response in the cell based caspase activation assay. This experimental data demonstrates that serum protein binding is an important factor.

![Fig. 1. Structures of compound 1, Bilirubin (Pdb ID: 2VUE), and Camptothecin (Pdb ID: 4L9K) that bind to drug site 3 in domain 1B of albumin.](image1)

![Fig. 2. In vitro biochemical assays of compound 1 showing titration curves with different amounts of FBS (0,1,10%). (A) TR-FRET binding and (B) Caspase activation in NCI-H929 cancer cells.](image2)

| Compound Characterization in Biochemical and Cell-based Assays in the Presence and Absence of FBS or Albumin. |
|-----------------|-----------------|-----------------|-----------------|
| **K<sub>i</sub> (nM)** TR-FRET Assay | Mcl-1 + 0% FBS | 0.5 | 1% FBS | 1.4 |
| | Mcl-1 + 2% FBS | 1.5 | Mcl-1 + 200 µg/ml Bovine Serum Albumin | 1.5 |
| | Mcl-1 + 200 µg/ml Human Serum Albumin | 1.4 |
| **Predicted Albumin K<sub>a</sub> (µM)** | Eq. 3 | 1.8 |
| **Caspase 3/7 IC<sub>50</sub> (µM)** | H929 0% FBS | 0.1 | H929 1% FBS | 1.4 |
| | H929 10% FBS | 6.0 |

Based on the manufacturers analysis, the concentration of albumin in the lot used for these studies is 180 µg/ml at 1%. 

![Table 1](table1)
to consider in interpreting our compound SAR in cell based assays containing serum.

3.2. Structures of compounds binding to site 3 of Human Serum Albumin

To help understand the structural basis for binding to albumin, we determined the structure of compound 1 bound to HSA. The three dimensional structure of compound 1 bound to Human Serum Albumin was obtained by X-ray crystallography. From the structure of compound 1 complexed to HSA (Fig. 3A), we observed that this compound binds to site 3 in domain IB (Fig. 3B). Some additional, but small volumes of unaccounted-for electron density (at the 3 sigma level) are present in site 2. The residual density was however poorly defined and a model for the ligand could not be built in this density. There are two HSA molecules in the asymmetric unit, and the electron density of the inhibitor in site 3 is present in both copies with a ligand occupancy in each copy of about 0.8.

The site 3 binding pocket in domain IB of albumin is formed by four alpha helices and is covered by an extended loop (Fig. 3AB). The pocket consists of a mix of hydrophobic and polar residues, which enables it to recognize fatty acids as well as bilirubin and other larger acid containing lipophilic molecules. Compound 1 contains three parts (Fig. 1) with different physical characteristics. The propyl linker and dimethyl chloro phenyl (region A in Fig. 1) is the most hydrophobic and flexible part of the ligand and it packs against complementary hydrophobic residues of albumin near to residue I142. The top of the ligand containing the trimethyl pyrazole (region B) is a balance of polar hydrophilic and hydrophobic moieties. This region of the ligand is covered by loop L1 (115–118) of albumin. The hydrophobic loop residues, L115 and P118 sit above the ligand in the pocket, and the guanidine of R117 makes both polar and hydrophobic contacts with compound 1. Most of the pyrazole at the 7 position of the indole packs tightly against albumin with the exception of the 1 position at the top of the pyrazole which is partially exposed to solvent. The 6-Cl indole core sits in a more polar region of the pocket and is near to K137, Y161, and Y138. Finally, the indole acid headpiece (region C) is the most exposed part of the molecule when bound to albumin. The acid of the indole headpiece sits near R186 and R117 with most of the indole headpiece sitting at the edge of the cleft between domains IB and IIIB. The Arg residues near the ligand in the structure suggests that removal of the acid or changing its position could change the albumin binding affinity of the ligand.

In Fig. 3 (CD), we compare the binding of our ligand to two other compounds that bind to site 3. Bilirubin (Fig. 3C, PDB ID: 2VUE) and Camptothecin (Fig. 3D, Pdb ID: 4LSK). Although very different in their physical characteristics, all of the ligands share common binding interactions. In the bilirubin (cyan) structure, the 3-methyl-4-vinyl-dihydro-pyrrol-2-one, binds into the same hydrophobic pocket near I142 as the dimethyl chloro phenyl of our Mcl-1 inhibitor. Bilirubin’s other 3-methyl-4-vinyl-dihydropyrrol-2-one overlaps with the 7-pyrrole of our ligand. The two acids of bilirubin interact with R186 and R117. Similar contacts are made by camptothecin. In camptothecin, the quinolone of the pentacyclic ring sits near residue I142 making a hydrophobic contact analogous to the dimethyl chloro phenyl of compound 1 and the 3-methyl-4-vinyl-dihydropyrrol-2-one of bilirubin. The hydroxyl and acid of camptothecin makes polar contacts to both R186 and R117 in a similar fashion as bilirubin and compound 1.

The binding motif of these three ligands share 2 common interactions with albumin. The first is hydrophobic burial of the ligand in the pocket around I142, which may serve as the hydrophobic hotspot that anchors ligands to Site 3. The second interaction, involves a polar contact to R186 and/or R117 which is also likely to be a common contact exploited by ligands that bind to site 3.

3.3. Species differences in binding site 3 of albumin

Rodent PK studies are generally used to help predict the PK in humans. Differences in albumin from different species can however change drug binding in plasma and hence the PK in rodents versus humans. We used a sequence alignment (Supplementary Material Table 3) between rodent and human albumin to compare the differences in the albumin binding sites. It can be seen from this alignment that the ligand contact residues in drug binding site 1 and 2 are mostly conserved. However, site 3 has two residues that are different in rodents and humans, R186K and I142V, both of which contact compound 1. These are conservative differences and when we tested our compound using albumin from rat serum and human albumin both at 200 μg/ml, we saw no significant difference in the plasma shift assay (data not shown).

3.4. Comparison to ligand binding to Mcl-1

Many of the features of compound 1 that contribute to binding to albumin are also important for binding to Mcl-1. When compound 1 binds to Mcl-1 (Fig. 4A PDB ID: 5IF4), the 6-CI indole core of 1 binds to the top of the pocket in the BH3 binding groove of Mcl-1. The di-methyl chloro phenyl binds deeply into a lower pocket, which is formed by a ligand induced bend in Helix 4 (H4) which helps accommodate this group into the deep hydrophobic pocket. The top of H4 protrudes slightly into the BH3 binding cleft and causes the loop connecting H4–H5 to be pulled toward the ligand. The 7 indole di-methyl pyrazole substitution is oriented out of the indole plane with one of the methyls pointing to the P3 pocket. Much of the 7 position pyrazole substitution is exposed and could tolerate polar or larger groups. The lactam of 1 locks the peptide bond conformation and is used to attach the final indole acid headpiece of the molecule. The acid of the head piece points towards the backbone of a loop containing G257-V258 with the indole headpiece sitting over R263.
If the ligand structure of 4A is used to superimpose the Mcl-1 complex onto the albumin complex by using a pairwise fit of all ligand and no protein atoms, the superposition shown in Fig. 4B is obtained. In both complexes, the di-methyl chloro phenyl of the ligand pushes deep into a hydrophobic pocket. In Mcl-1, this pocket is formed by helix 4 and 5 while in HSA the pocket is formed by helix 9 and 10 of site 3 domain IB. The propyl-di-methyl chloro phenyl of the ligand is flexible and adapts different binding poses in the hydrophobic pocket of Mcl-1 and HSA Site 3. The indole acid is also rotated in the HSA structure when compared to its pose in Mcl-1 (Fig. 4B). This suggests that reducing the rotation of the headpiece could also decrease ligand binding to HSA.

In Fig. 4C, a surface representation of Mcl-1 with the binding poses of the ligand when bound to Mcl-1 and albumin site 3 is displayed. This can be compared to the surface representation of albumin with the binding poses of the ligand when bound to Mcl-1 and site 3 in 4D. Compound 1 is more buried in albumin (4D) than in Mcl-1 (4C), suggesting that adding substituents to the ligand would reduce binding to albumin and if done correctly still enhance or maintain binding to Mcl-1. From the top down view of the protein’s surfaces, shown in Fig. 4C,D, three areas of the ligand are evident that could potentially be used to create a steric hindrance to albumin binding. These are substitutions in the headpiece, lactam bridge or on the pyrazole of the ligand. We show in the Supplementary Material, earlier data on ligands with larger substitutions on the pyrazole. The improvement in plasma shift observed in these earlier compounds is consistent with our structural analysis of compound 1 binding to albumin.

4. Conclusion

We observed that the cell efficacy of compound 1 was reduced in in vitro binding and cell based assays in the presence of FBS or albumin. In order to help understand the structural basis of this interference, we determined the structure of compound 1 when bound to albumin and found that it binds to site 3 in domain IB. Site 3 is likely better suited to accommodate our compound because it is larger than the pockets in drug sites 1 and 2. The structure of compound 1 with HSA and with Mcl-1 was compared to identify the common and different binding features of the ligand to each individual protein. Even though HSA can adapt to many different ligands, a structural analysis can generally identify regions of the ligand that can be modified to reduce albumin binding and still maintain or improve binding to target.12,13 For our compound, we observed three potential regions that could be modified to reduce albumin binding while maintaining binding to Mcl-1.

Ligands binding to drug site 3 of HSA may be more common for larger more extended molecules. The structural data presented here adds to the knowledge base of preferences for this site and could be useful for the development of QSAR models and docking approaches to predict albumin binding of compounds to this site.18,19

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A. Supplementary data

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References


