



Crystal structures of multidrug-resistant HIV-1 protease in complex with two potent anti-malarial compounds

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ABSTRACT

Two potent inhibitors (compounds **1** and **2**) of malarial aspartyl protease, plasmepsin-II, were evaluated against wild type (NL4-3) and multidrug-resistant clinical isolate 769 (MDR) variants of human immunodeficiency virus type-1 (HIV-1) aspartyl protease. Enzyme inhibition assays showed that both **1** and **2** have better potency against NL4-3 than against MDR protease. Crystal structures of MDR protease in complex with **1** and **2** were solved and analyzed. Crystallographic analysis revealed that the MDR protease exhibits a typical wide-open conformation of the flaps (Gly48 to Gly52) causing an overall expansion in the active site cavity, which, in turn caused unstable binding of the inhibitors. Due to the expansion of the active site cavity, both compounds showed loss of direct contacts with the MDR protease compared to the docking models of NL4-3. Multiple water molecules showed a rich network of hydrogen bonds contributing to the stability of the ligand binding in the distorted binding pockets of the MDR protease in both crystal structures. Docking analysis of **1** and **2** showed a decrease in the binding affinity for both compounds against MDR supporting our structure–function studies. Thus, compounds **1** and **2** show promising inhibitory activity against HIV-1 protease variants and hence are good candidates for further development to enhance their potency against NL4-3 as well as MDR HIV-1 protease variants.

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1. Introduction

AIDS (acquired immunodeficiency syndrome) is one of the major causes of global human morbidity [1]. Rapid and error prone replication of HIV-1 (human immunodeficiency virus type-1) leads to mutations that are incorporated into viral proteins resulting in drug resistance [2]. HIV-1 protease [3,4] is a homo-dimeric aspartyl protease with two catalytic aspartic acid residues (Asp25 and Asp125) and has been validated as an important drug target [5] due to its critical role in the viral maturation and virulence [3,4]. In spite of the availability of numerous potent HIV-1 protease inhibitors (PIs), multidrug-resistant (MDR) clinical isolates such as MDR 769 [6] show resistance against most of the PIs [7,8]. Accumulation of mutations can cause changes in the three-dimensional structure of the viral proteins including the protease [9,10]. Such conformational changes were later confirmed by other groups

[11]. Thus, new therapeutic agents are needed to gain an efficient control over the inhibition of such MDR strains.

Previously, HIV-1 protease inhibitors have been shown to exhibit antimalarial activity [12]. In the current study, two highly potent inhibitors, compounds **1** and **2** (Fig. 1), of malarial aspartyl protease, plasmepsin-II (PLM-II) [13,14] were evaluated against wild type (NL4-3) and MDR 769 HIV-1 protease variants. Compounds **1** and **2** were the first examples of fully characterized mechanism-based inhibitors of aspartyl protease, PLM-II [15]. Both compounds were previously shown to have good therapeutic index with low nanomolar potency against PLM-II (Table 1) making them potential new treatments for malaria. Based on the similarities between the catalytic mechanism [16] and the inhibitor design [17] for PLM-II and HIV-1 protease, we hypothesized that **1** and **2** might inhibit the NL4-3 and MDR 769 HIV-1 protease variants. This is a report evaluating anti-malarial compounds against the MDR 769 HIV-1 protease.

Both **1** and **2** feature a transition-state mimic hydroxyl group in a statine based core that is known to enhance the affinity of the compound for the enzyme [18]. Potencies of both compounds were evaluated against NL4-3 and MDR 769 variants of HIV-1 protease. Crystal structures of MDR7 769 A82T mutant HIV-1 protease in complex with **1** and **2** were solved and analyzed. Both compounds

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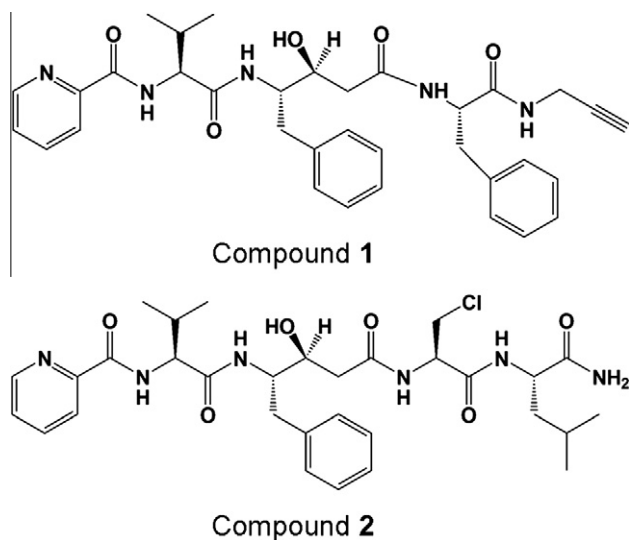


Fig. 1. Structures of compounds **1** and **2**.

Table 1
IC₅₀ (μM) values of compounds **1** and **2**.

Compound	IC ₅₀ (μM) ± standard deviation		
	NL4-3	MDR769	PLM-II
1	0.99 ± 0.13	14.53 ± 1.65	0.02
2	5.7 ± 0.33	30.30 ± 1.97	0.025
Acetyl pepstatin	25.3 ± 1.56	56.75 ± 3.17	7.5

The IC₅₀ values of compounds **1** and **2** against the wild type (NL4-3) and MDR769 HIV-1 protease variants were obtained in the current study and the IC₅₀ values against PLM-II were taken from Gupta et al. [15]. IC₅₀ value for the control, acetyl pepstatin against PLM-II was taken from Parikh et al. [12]. Both compounds show better inhibition profiles against NL4-3 as well as MDR769 HIV-1 protease variants compared to that of the control, acetyl pepstatin.

were further docked into the active site cavity of NL4-3 protease using AutoDock Vina [19]. A detail crystallographic and modeling analysis was performed to understand the binding profiles of both the compounds.

2. Materials and methods

2.1. Compounds and HIV-1 protease

Compounds **1** and **2** were synthesized using peptide coupling method as reported previously [15]. Catalytically active NL4-3 protease was purchased from Bachem (CA, USA) at a concentration of 0.3 mg/ml. Expression and purification of catalytically active (D25/D125) MDR769 and inactive (N25/N125) MDR769 A82T HIV-1 protease variants were performed as described previously [20]. The inactive (D25N/D125N) variant of the MDR769 A82T protease was chosen to avoid the auto proteolysis for crystallographic studies. Briefly, both the active and inactive variants of MDR protease were over expressed and purified from the inclusion bodies by solubilizing them in 6 M urea. Following the ion-exchange chromatographic purification, the protease species were refolded and concentrated. The final yield of the active MDR protease was low (0.5 mg/ml) compared to that of the inactive A82T mutant (1.5 mg/ml) as predicted.

2.2. Enzyme inhibition assay

FRET-based enzyme inhibition assay was performed using the fluorescent HIV-1 substrate (purchased from Molecular probes –

CA, USA) with excitation and emission wavelengths, 340 and 490 nm, respectively. Substrate was dissolved in DMSO at a stock concentration of 0.5 mM. The emission signal was recorded using the Model-M5 plate reader from Molecular Devices (CA, USA). Reaction buffer (0.1 M sodium acetate, 1.0 M sodium chloride, 1.0 mM dithiothreitol (DTT), 1.0 mM ethylenediaminetetraacetic acid (EDTA), 10% DMSO (dimethyl sulfoxide) and 1 mg/ml bovine serum albumin (BSA)) was made fresh each time before the assay. The final pH was adjusted to 4.7. Both NL4-3 and MDR protease variants were diluted by 1600- and 3200-fold, respectively, with fresh reaction buffer before assay in order to achieve proper enzyme activity. Ligand was diluted serially by 2-fold with DMSO. Ninety nine microliters of diluted HIV-1 protease and 1 μl of ligand at various concentrations were incubated at 37 °C for 10 min. One microliter of fluorescent substrate at stock concentration was added right before reading at 37 °C. The fluorescent signal was recorded for 20 min with 1 min intervals. The final IC₅₀ values are average of three experiments.

2.3. X-ray crystallography

A 20-fold molar excess of either **1** or **2** was added to the catalytically inactive A82T variant of MDR769 HIV-1 protease and incubated on ice for 1 h before crystallization. Hanging drop, vapor diffusion method was used for crystallization in the pH range of 2.0–12.0. For each condition, 1 μl of enzyme-drug complex was mixed with 1 μl of well solution. Diffraction quality crystals were obtained in the range of pH 7.0–8.5 at a sodium chloride concentration of 0.3–0.5 M. Diffraction datasets were collected at the LS-CAT beam line 21 at the APS (advanced photon source), IL. Diffraction data was processed using HKL2000 [21]. Structure solutions were obtained using MOLREP [22] through CCP4 [23] suite of programs with MDR769 HIV-1 protease (PDB ID: 1TW7) as a search model. The models were then refined further using REFMAC5 [24]. Electron density maps were calculated using XtalView/Xfit [25] program. The difference electron density (calculated using XtalView/Xfit program) was contoured between 1.8 and 2.0 σ while fitting the ligand. REFMAC libraries for the ligands were obtained from PRODRG server [26]. Refined coordinates for the ligand conformation with highest occupancy and reasonable thermal factor values were included in the final coordinate file. Solvent molecules were then built in using ARP/wARP [27,28] through CCP4. Validation of the final structures was performed by PROCHECK [29,30]. Coordinates for crystal structures of MDR769 A82T HIV-1 protease in complex with compounds **1** and **2** were deposited in RCSB (Regional Collaboratory for Structural Bioinformatics) PDB (protein data bank) with accession codes 3R0Y and 3R0W, respectively. Analysis of the protease-inhibitor contacts was performed using CONTACT/ACT [31] module from CCP4 suite of programs.

2.4. Docking

Crystal structures of wild type HIV-1 protease in complex with darunavir (PDB ID: 2IEN) and PLM-II in complex with TIT-1330 (PDB ID: 1W6H) were chosen as starting structures. The coordinates for ligand, water and other small molecules were deleted from both the structures and apo-enzyme coordinates were then used as receptors to build the docking grid. All the polar hydrogen atoms were added to the receptors before generating the docking grid. The docking grid was generated in and around the active site cavity for all the receptors using AutoDock tools [32] GUI (graphic-user interface). The ligands were prepared in the same way using AutoDock tool GUI. Initially, darunavir taken from the crystal structure was re-docked back into the active site of the HIV-1 protease taken from 2IEN as a positive control using AutoDock Vina.

Similar positive control was performed for TIT-1330 against PLM-II taken from 2W6H. For the MDR769 HIV-1 protease, crystal structures 3R0Y and 3R0W were used. Mutations N25D and N125D were introduced into the MDR protease (to represent catalytically active protease) model before receptor grid generation. All the graphics in this article were prepared using the open source molecular graphics program, PyMol (v0.99rc6) (www.pymol.org).

3. Results and discussion

3.1. Compounds **1** and **2** show better potency against wild type HIV-1 protease than the MDR 769 HIV-1 protease

Fluorescence resonance energy transfer (FRET)-based enzyme inhibition assays revealed that both **1** and **2** show inhibitory activity with better potency against NL4-3 than MDR 769 HIV-1 protease. As shown in Table 1, the IC₅₀ values show that **1** is 14-fold and **2** is 5-fold more potent against NL4-3 than MDR 769 HIV-1 protease. Compound **1** is at least 25-fold (against NL4-3) and 4-fold (against MDR 769) more potent compared to the control, acetyl pepstatin (a general aspartyl protease inhibitor). Similarly, **2** is at least 4-fold (against NL4-3) and almost 2-fold (against MDR 769) more potent compared to acetyl pepstatin.

3.2. Crystal structures of MDR 769 HIV-1 protease in complex with compounds **1** and **2** show altered binding orientations of the compounds

Crystal structures of MDR 769 HIV-1 protease variant in complex with **1** and **2** were solved in the space group P4₁ with one protease dimer per asymmetric unit. Diffraction data and refinement statistics are given in Table 2. Crystallographic analysis suggested that both compounds bind the MDR protease in two possible orientations each. The major binding orientation for each compound with higher occupancy and lower thermal factors was determined to be the final binding orientation and were refined to obtain the final structures. The MDR protease in both structures showed the typical wide-open conformation of the flaps contributing to the expansion in the volume of the active site cavity. Due to this wide-open conformation of the flaps, the overall binding orientation of either compound looks altered when compared to other HIV-1 protease inhibitors. Analysis of protease-inhibitor contacts suggests that both **1** and **2** are involved in three direct polar contacts each with MDR protease and multiple polar contacts with active site water molecules. As shown in Fig. 2, the transition-state mimic hydroxyl moiety from either compound is involved in hydrogen bonding with Asn25 and Asn125. Details of polar contacts are given in Table 3. As shown in Table 4, compound **1** is involved in a total of 13 hydrophobic contacts while compound **2** is involved in a total of 15 hydrophobic contacts.

3.3. Crystallographic water molecules compensate for the loss of direct contacts in the expanded active site cavity of the MDR 769 HIV-1 protease

Multiple water molecules were observed in both crystal structures surrounding the ligand in the expanded active site cavity of the MDR protease. Both structures show a rich network of hydrogen bonding among the waters in and around the active site area contributing to the stability of the compounds in the expanded active site cavity. Out of the three water molecules that showed direct polar contacts with compound **1**, two were observed to be bridging between the protease (one with N125 and the other with G127) and the compound (Fig. 2A). Similarly, out of the four water molecules that showed direct polar contacts with compound **2**, three were observed to be bridging between the protease (one with

Table 2
X-ray diffraction data and refinement statistics.

Parameters	Compound 1	Compound 2
PDB entry	3R0Y	3R0W
<i>Crystal parameters</i>		
Resolution (Å)	18.95–1.65	18.98–1.7
Unit cell (Å)	$a = b = 45.64$, $c = 102.00$, $\alpha = \beta = \gamma = 90$	$a = b = 45.73$, $c = 102.15$, $\alpha = \beta = \gamma = 90$
Space group	P4 ₁	P4 ₁
Solvent content (%)	47.0	47.0
<i>Data processing</i>		
No. of unique reflections	25092 (2505)	22795 (2281)
Mean $I/\sigma(I)$	30.81 (5.0)	40.62 (6.9)
R_{merge} (%) ^b	5.7 (45.6)	4.0 (26.6)
Data redundancy	7.6 (7.5)	4.0 (4.0)
Completeness (%)	100 (100)	99.3 (100)
<i>Refinement statistics</i>		
No. of reflections used	23751	21607
R_{cryst} (%) ^a	19.78	19.78
R_{free} (%)	23.06	24.54
No. of protein atoms	1512	1512
No. of ligand atoms	44	44
No. of water molecules	193	208
<i>Mean temperature factors (Å²)</i>		
Protein	21.34	21.25
Main chains	20.08	20.20
Side chains	22.72	22.41
Ligand	50.52	59.66
Waters	37.63	35.44
R.M.S.D. bond lengths (Å)	0.019	0.013
R.M.S.D. bond angles (°)	1.67	1.50
<i>Ramachandran plot</i>		
Most favored (%)	92.9	93.6
Additional allowed (%)	7.1	6.4
Generously allowed (%)	0	0
Disallowed (%)	0	0

Values in parentheses are for the highest resolution shell.

^a $R_{\text{cryst}} = \sum ||F_{\text{obs}}| - |F_{\text{calc}}|| / \sum |F_{\text{obs}}|$.

^b $R_{\text{merge}} = \sum |I - \langle I \rangle| / \sum I$.

G148 and the other two with D129 backbone amide nitrogen and side chain oxygen atoms) and the compound (Fig. 2B). Typically one conserved water molecule is observed in most of the HIV-1 protease crystal structures that bridges the inhibitor to the protease flaps but in the current study, due to the wide-open conformation of the flaps, the conserved water molecule was lost. Both structures showed a conserved trend of inter-flap water triad consisting of three water molecules bridging the wide-open flaps. The tips of the flaps (Gly48 to Gly52) show a curly conformation with one water molecule bound in the concavity of either flap (intra-flap water). The intra-flap water molecules showed a conserved trend of hydrogen bonding with the backbone of residues Gly49, Ile50, Gly51 and Gly52 in each monomer from both structures.

3.4. Docking solutions suggest loss of binding affinity for either compound against the MDR 769 HIV-1 protease

Both compounds **1** and **2** were individually docked into the active site of NL4-3. Docking solutions were analyzed for contacts between the protease and compounds. As shown in Fig. 2, compound **2** showed more contacts with the NL4-3 receptor in the docked model compared to compound **1**. Both compounds show a stretched-out binding orientation in the docked models of NL4-

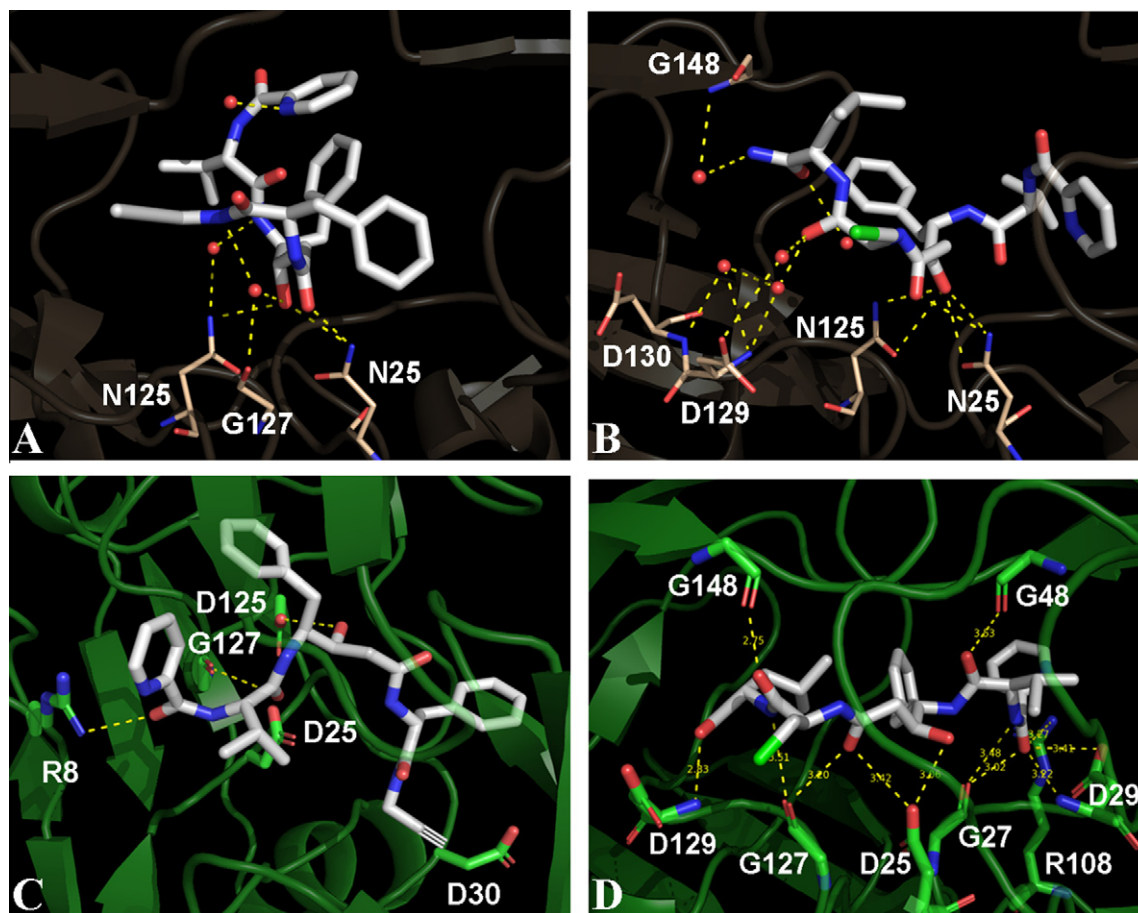


Fig. 2. Three-dimensional view of contacts for compounds **1** and **2**. (A and B) Show contacts made by compounds **1** and **2**, respectively with the MDR protease in their corresponding crystal structures. (C and D) Show docked models of wild type HIV-1 protease with compounds **1** and **2**, respectively. The compounds are represented as white sticks in all panels while the protease residues that are involved in polar contacts are highlighted as sticks. The protease residues are colored in beige and green for the MDR and wild type protease, respectively. All the polar contacts are represented as yellow dashed lines and the active site water molecules are represented as small red spheres. Three waters that have direct contacts with compound **1** are shown in (A). Out of the three water molecules in (A), two bridge the compound to N125 and G127 of the MDR protease. Five waters are shown in (B) out of which, four waters are directly in contact with compound **2**. Three waters bridge the compound to D129 and G148 in (B). The binding orientation of compound **2** is more stretched-out, with more contacts compared to that of the compound **1**. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Table 3
Protease inhibitor polar contacts.

Enzyme	Compound 1	Total	Compound 2	Total
MDR769	N25 (2), N125 (1), H ₂ O (3)	6	N25 (3), N125 (2), H ₂ O (4)	9
NL4-3	R8 (1), D125 (1), G127 (1)	3	D25 (2), G27(2), D29(2), G48 (1), R108 (1), G127 (2), D129 (1), G148 (1)	12
PLM-II	D34 (2), G36 (1), S79 (2), Y192 (1), G216 (1), T217 (1)	8	D34 (3), G36 (1), V78 (1), S79 (3), Y192 (2), D214 (1), T217 (1), S218 (2)	14

Polar contacts made by compounds **1** and **2**. Amino acid residues (represented as single letter amino acid codes) involved in polar contacts with compounds **1** and **2** are listed with their corresponding number of contacts in parenthesis. In the case of NL4-3 and MDR769, the protease residues are numbered 1–99 for monomer-A and 101–199 for monomer-B. Active site water molecules that are involved in polar contacts with either compound are listed along with the protease residues. Compound **2** shows more contacts than that of compound **1** in all three cases. Both compounds show highest number of polar contacts with PLM-II.

3 compared to their corresponding crystal structures with MDR protease. As shown in Fig. 3, both compounds show better binding affinity values against NL4-3 and PLM-II compared to MDR protease. Compounds **1** and **2** show a loss of 2.8 and 2.5 kcal/mol, respectively in their binding affinities against the MDR protease.

Table 4
Protease inhibitor hydrophobic contacts.

Enzyme	Amino acid residues involved in hydrophobic contacts	
	Compound 1	Compound 2
MDR769	G27, I50, P81, T82, V84, I147, P181, T182	A28, V32, V84, G148, I150, P181, T182, V184
NL4-3	R8, A28, D30, V32, I47, I50, I84, A128, D129, D130, V132, I147, G149, I150, L176, P181, V182, I184	L23, A28, V82, I84, A128, D129, I147, G149, I150, V182, I184
PLM-II	I14, M15, Y17, I32, D34, G36, S37, M75, N76, Y77, V78, S79, F111, T114, S118, F120, I123, L131, Y192, D214, G216, T217, S218, A219, T221, F244, Q275, N288, I290, L292, F294, I300	I14, M15, Y17, I32, D34, G36, S37, A38, M75, N76, Y77, V78, S79, F111, T114, A117, S118, F120, I123, L131, Y192, I212, D214, G216, T217, S218, A219, T221, F244, I290, L292, F294, I300

Hydrophobic contacts made by compounds **1** and **2**. Amino acid residues (represented as single letter amino acid codes) involved in hydrophobic contacts with compounds **1** and **2** are listed for MDR769, NL4-3 and PLM-II. In the case of NL4-3 and MDR769, the protease residues are numbered 1–99 for monomer-A and 101–199 for monomer-B. Both compounds show almost equal number of hydrophobic contacts with the MDR769 HIV-1 protease. Compound **1** shows more hydrophobic contacts than **2** in the docked models of NL4-3 HIV-1 protease. Both compounds show highest number of hydrophobic contacts with PLM-II.

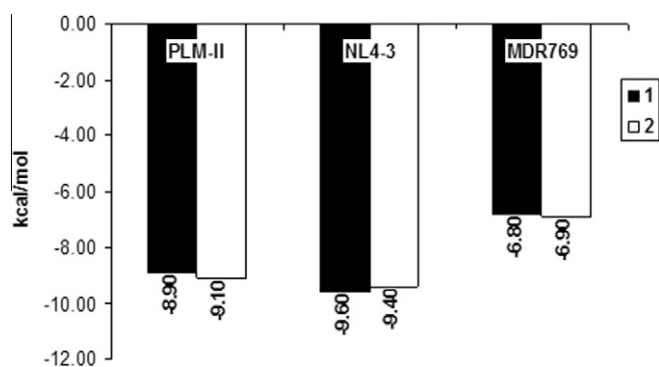


Fig. 3. Binding affinities of compounds **1** and **2**. Binding affinity values in kcal/mol (y-axis) are plotted for compounds **1** (black bars) and **2** (white bars) against Plasmepsin-II (PLM-II), wild type (NL4-3) and MDR 769 HIV-1 protease receptors. Compounds **1** and **2** show equal or better binding affinity values against the NL4-3 receptor compared to their corresponding PLM-II models. The MDR 769 HIV-1 protease models show loss (>2 kcal/mol) of binding affinity for both compounds.

Availability of larger chemical space in the expanded active site cavity of the MDR protease causes multiple possible binding orientations of the inhibitor until stable binding is achieved. This increase in the chemical space in part explains the loss of binding affinity for compounds **1** and **2** docked against the MDR 769 HIV-1 protease receptor.

3.5. Compound **2** shows less fold-resistance against MDR 769 HIV-1 protease compared to compound **1**

Compounds **1** and **2** showed promising inhibition profiles against the NL4-3 protease. Previously, MDR clinical isolates such as MDR 769 HIV-1 protease were shown to be 14- to more than 100-fold resistant against FDA approved and experimental PIs [6]. Recently, it has been shown that MDR 769 HIV-1 protease exhibits 2- to almost 600-fold resistance against FDA approved PIs [8]. In the current study, the MDR 769 showed 14- and 5-fold resistance against **1** and **2**, respectively suggesting that both compounds are within the range of the regular PIs with respect to the fold-change in their potency. Crystal structures and docking models showed that compound **2** is bound in a stretched-out fashion with relatively more contacts with the protease than compound **1** suggesting that compound **2** is less prone to loss of potency due to mutations in the protease compared to compound **1**.

In summary, the crystals structures and docking models support the inhibitory activity of **1** and **2** against the NL4-3 as well as against MDR 769 HIV-1 protease variants suggesting that further development of these compounds would enhance their potency. We are currently in the process of evaluating new compounds designed based on the crystal structures reported here.

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