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Novel Human Lipoxygenase Inhibitors Discovered Using Virtual Screening with Homology Models

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We report the discovery of new, low micromolar, small molecule inhibitors of human platelet-type 12- and reticulocyte 15-lipoxygenase-1 (12-hLO and 15-hLO) using structure-based methods. Specifically, we created homology models of 12-hLO and 15-hLO, based on the structure of rabbit 15-lipoxygenase, for in silico screening of a large compound library followed by in vitro screening of 20 top scoring molecules. Eight of these compounds inhibited either 12- or 15-human lipoxygenase with lower than $100 \,\mu\text{M}$ affinity. Of these, we obtained IC50 values for the three best inhibitors, all of which displayed low micromolar inhibition. One compound showed specificity for 15-hLO versus 12-hLO; however, a selective inhibitor for 12-hLO was not identified. As a control we screened 20 randomly selected compounds, of which none showed low micromolar inhibition. The new low-micromolar inhibitors appear to be suitable as leads for further inhibitor development efforts against 12-hLO and 15-hLO, based on the fact their size and chemical properties are appropriate to classify them as drug-like compounds. The models of these protein—inhibitor complexes suggest strategies for future development of selective lipoxygenase inhibitors.

Introduction

Lipoxygenases are a class of non-heme, iron-containing enzymes that catalyze the incorporation of molecular oxygen into 1,4,-cis,cis-pentadiene-containing fatty acids (e.g. linoleic and arachidonic acids) to form hydroperoxide products. 1-4 Lipoxygenases are the first committed step in a cascade of metabolic pathways that are implicated in the onset of cancers, asthma, and heart disease, making them candidates for inhibitory pharmaceutical therapy.⁵⁻⁷ The human isozymes, 5-, 12-, and 15-lipoxygenase, are associated with different disease states, which suggests that selective inhibition may be important in targeting them for therapeutic purposes.^{8–10} Recent discovery efforts of lipoxygenase inhibitors¹¹⁻¹³ have focused especially on structure-activity relationships¹⁴⁻²² and natural product isolation.^{23–31} In a very recent report, pharmacophore virtual screening methods have yielded novel inhibitors with selectivity for rabbit 15-lipoxygenase.³² One selective human 5-lipoxygenase inhibitor, Zileuton,²¹ has been approved by the FDA for the treatment of asthma.33

We report the discovery of new, low micromolar, small molecule inhibitors of human platelet-type 12-lipoxygenase (12-hLO) and human reticulocyte 15-lipoxygenase-1 (15-hLO) using structure-based, in silico methods and in vitro screening. Prior to this work, structure-based methods have not played a major role in the discovery of lipoxygenase inhibitors, because experimental structures of the human lipoxygenases do not exist. Here, we create homology models of 12-hLO and 15-hLO, based on the structure of rabbit reticulocyte 15-lipoxygenase (15-rLO),³⁴ followed by docking of a large compound library and in vitro screening of the top scoring molecules.

Although docking using comparative models is more challenging and less successful than docking to crystallographic

structures, a recent review by Jacobson and Sali³⁵ describes efforts dating back to at least 1993 in which comparative protein models have been successfully used to aid inhibitor discovery. Some of the proteins successfully targeted using docking against comparative models include cysteine proteases in several parasites,^{36–39} matriptase,⁴⁰ Bcl-2,⁴¹ retinoic acid receptor,⁴² DHFR in T. cruz, 43 human CK2, 44 and CDK4. 45 Despite these successes, however, the role of comparative models in structureguided drug design remains small. A few studies including those by McGovern and Shoichet, 46 Diller and Lee47 and Oshiro et al.48 have attempted to quantify the accuracy of docking to comparative models by evaluating the ability to enrich known inhibitors relative to large databases of "decoy" ligands. The success of docking against homology models varies significantly from case to case in these studies, but in general is somewhat poorer than docking against crystal structures.

Homology models of lipoxygenase enzymes have been used in several previous studies for purposes other than inhibitor discovery. Specifically, these studies have used homology models, sometimes in combination with docking methods, to propose models for substrate binding and specificity in various lipoxygenases.

In previous work, we have used docking methods to propose a binding model where linoleic acid binds "carboxylic end first" in the active site of soybean lipoxygenase, which was consistent with experimental mutagenesis studies.⁵² In this work, we performed virtual screening of 50 000 drug-like molecules against the active sites of 12-hLO and 15-hLO homology models. The top 10 compounds from each virtual screen were then experimentally screened against each enzyme; no filtering of the computational results was performed. Eight of these 20 compounds inhibited either 12-hLO or 15-hLO with lower than $100 \,\mu\text{M}$ affinity. Of these, we obtained IC₅₀ values for the three best inhibitors, all of which displayed low micromolar inhibition. One compound shows specificity for 15-hLO versus 12-hLO; however, a selective inhibitor for 12-hLO was not identified. As a control we also screened 20 randomly selected compounds, of which none showed low micromolar inhibition. The new low-

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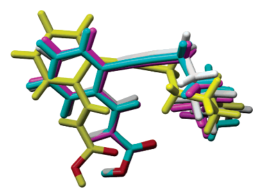


Figure 1. Superimposed images of the cocrystallized inhibitor RS7 in the 15-rLO crystal structure (grey), as well as the same inhibitor docked into 15-rLO structure (purple) and the 12-hLO (yellow) and 15-hLO (teal) homology models. The carboxylic acid group on the ligand is protonated because it forms a hydrogen bond with the backbone carbonyl group of Leu597 in 15-rLO (2.6 Å O-O distance). The lipoxygenase proteins are not shown, but are structurally superimposed on the 15-rLO crystal structure. Note that the binding pose of RS7 in 12-hLO and 15-hLO is not known. We assume that the inhibitor will bind similarly in 15-hLO as in 15-rLO, as suggested by the docking poses, but the pose of RS7 in 12-hLO may not be meaningful.

micromolar inhibitors generally adhere to Lipinski's rules of five, with average molecular weights of 350, less than five rotatable bonds and an average log P of four. Moreover, these inhibitors are chemically dissimilar from known lipoxygenase inhibitors and provide new scaffolds that may be useful for further development of selective lipoxygenase inhibitors.

Results

As a first test, we confirmed the ability of the docking algorithm to reproduce the cocrystallized pose of the RS7 inhibitor in the 15-rLO crystal structure (Figure 1). In these tests, the carboxylic acid group on the ligand is protonated because it forms a hydrogen bond with the backbone carbonyl group of Leu597 in 15-rLO (2.6 Å O-O distance). The RMS deviation of the RS7 inhibitor from its cocrystallized position in 15-rLO is 0.67 Å (docking pose accuracy of better than 2.0 Å RMSD is generally considered acceptable). We also docked the RS7 inhibitor against the 12-hLO and 15-hLO homology models, although we know of no published results showing that it inhibits these enzymes. We assume that the inhibitor is likely to bind similarly in 15-hLO as in 15-rLO, as suggested by the docking poses, but the pose of RS7 in 12-hLO may not be meaningful. The RMS deviations of the RS7 inhibitor from its cocrystallized position in 15-rLO (by structurally superimposing the proteins) are 0.90 Å and 1.84 Å when docked against 15hLO and 12-hLO, respectively.

Figure 2 depicts surface representations^{65,66} of the substrate binding sites in the 15-hLO and 12-hLO homology models as well as the 15-rLO template. Selected residues that determine the size of the active site, and thus determine the chemistry performed by the respective enzymes, are depicted. The active sites of the 15-hLO and 15-rLO are roughly the same size but the 12-hLO active site is larger and extends further toward the back of the active site. This is consistent with experimental mutagenesis studies that demonstrated that the 12-hLO active site is larger than that of 15-hLO.⁶⁷

Subsequently, a set of 50 000 drug-like compounds, commercially available from ChemBridge ("diversity set"), was docked against the 12-hLO and 15-hLO homology models, using Glide's SP ("standard precision") mode. The top 1000 hits from this initial screen (ranked according to GlideScore) for both the

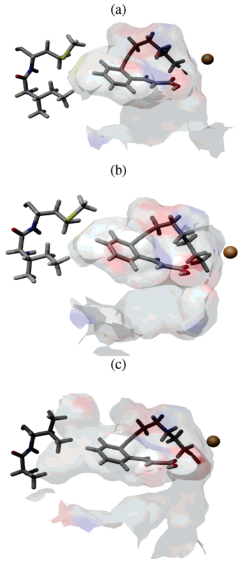


Figure 2. Surface representation of the 15-rLO (a), 15-hLO (b), and 12-hLO (c) active sites with the cocrystallized pose of the RS7 inhibitor shown for reference. The residues shown in wireframe are those that are primarily responsible for determining the size/depth of the active site among the three isozymes (Met and Ile in 15-rLO and 15-hLO; Val and Ala in 12-hLO). The iron is shown in a space-filling representation. The colors on the surface reflect the identity of the heavy atoms lining the cavity: red for oxygen, blue for nitrogen, and gray for carbon. These figures were prepared using Chimera,65 which employs the MSMS algorithm for surface generation.⁶⁶

12-hLO and 15-hLO homology models were then refined by redocking them using Glide's XP ("extra precision") mode.

The top 10 compounds from this docking protocol against both 15-hLO (compounds 1-10) and 12-hLO (compounds 11-20) were purchased and experimentally screened for their ability to inhibit lipoxygenase activity. All 20 compounds taken from the docking calculations were visually inspected to ensure reasonableness, but none were omitted from experimental testing. All 20 of the top hits formed putative hydrogen bonds to Gln546 and Glu355 in the active site and formed extensive hydrophobic interactions in the rear of the active site.

The results obtained from the experimental screen are shown in Table 1. Of the top 10 scoring compounds from the 15-hLO docking calculations, one compound (9) showed nonselective inhibition against both 15- and 12-hLO with IC₅₀ values of 9.2 \pm 1.4 and 12.3 \pm 0.9 μ M, respectively. Of the top 10 scoring compounds taken from the 12-hLO docking calculation, two

Table 1. Chemical Structures, Docking Ranks, and IC_{50} Values for Top Scoring Compounds in the Docking Screens against the Models of 12-hLO and 15-hLO a

	ID	_	SP Rank		XP I	Rank	IC ₅₀	
Chembridge ID Number	Number From	Structure		15- 12-		15- 12-		12-
6485150	Text 1	S S S S S S CI	708	hLO 2949	hLO 1	hLO –	>50	hLO >100
5245411	2		107	-	2	_	>100	>200
7523797	3	O NH	6	ı	3	ı	> 100	> 500
6738761	4		194	31905	4	ı	>100	>200
6712889	5	ST O O O O O O O O O O O O O O O O O O O	67	24753	5		>50	>100
6671738	6	HN CI	969	3277	6	=	>100	>100
7510319	7		298	ı	7	ı	>100	>200
6704363	8	S HN O	40	12917	8	ı	>50	>100
6640337	9	HO OHO N-NH	569	4781	9	-	9.2 ± 1.4	12.3 ± 0.9
6615087	10	O NH H O NH H O NH H	150	32049	10	-	>100	>200

	ID Number	Structure		SP Rank		XP Rank		IC ₅₀	
Chembridge ID Number	From Text			12- hLO	15- hLO	12- hLO	15- hLO	12- hLO	
6942880	11		ı	713	_	1	>100	>200	
5680672	12	S NH ₂ NH O NH Cl	1892	180	ı	2	18.8 ± 4.7	30.7 ± 6.8	
6862629	13		ı	491	ı	3	>50	>100	
6945303	14	TO HE COLOR	ı	609	_	4	>200	>200	
6635967	15	$\begin{array}{c} \begin{array}{c} \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ \begin{array}{c} \\ \\ \end{array} \\ \begin{array}{c} \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \end{array} \\ \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \end{array} \\ \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\$	1409	301	-	5	6.8 ± 1.2	>200	
7402594	16		ı	128	-	6	>50	>100	
7282756	17	HN O CI	I	832	-	7	>100	>200	
7383862	18	O NH	_	255	_	8	>100	>100	
6558319	19		_	709	_	9	>100	>100	
7232391	20		-	240	_	10	>100	>100	

^aOnly the top 1000 compounds from the Glide SP results were subjected to rescoring with Glide XP. Compounds 1-10 and 11-20 represent the top 10 hits from the Glide XP docking against 15-hLO and 12-hLO, respectively. Some compounds did not successfully dock into one or both of the models, generally due to a steric clash or other highly unfavorable interaction. The IC50 values for the three low micromolar inhibitors were determined by fitting the data presented in Supporting Information. The IC₅₀ values for the remaining molecules were estimated using single-point screens, and categorized as follows: $>50 (50-100 \mu M)$, $>100 (100-200 \mu M)$, $>200 (200-1000 \mu M)$, and $>1000 \mu M$.

exhibited low micromolar inhibition. Compound 12 inhibited both 12-hLO and 15-hLO with IC50 values of 30.7 \pm 6.8 μM and 18.8 \pm 4.7 μ M, respectively. However, compound 15 was selective against 15-hLO (IC₅₀ = 6.8 \pm 1.2 μ M) and not 12hLO (IC₅₀ > 200 μ M). We note that compound 15 has a basic nitrogen on the amidinium group (p $K_a \sim 6-7$) in the central region of the molecule that, if protonated, could form an ion pair with Glu356, as suggested by the docking pose.

Figure 3 shows compound 15 docked into both 12-hLO and 15-hLO. This compound makes the same hydrogen bond

Figure 3. Surface representations of 15-hLO selective compound **15** docked to 15-hLO (left) and 12-hLO (right). Glu356 from the receptor is shown in wireframe, because it forms hydrogen bonds to the inhibitor. The iron is shown in a space-filling representation. The colors on the surface reflect the identity of the heavy atoms lining the cavity: red for oxygen, blue for nitrogen, and gray for carbon. Compound **15** fills the binding pocket in 15-hLO much better than in 12-hLO, possibly providing an explanation for its selectivity against 15-hLO. These figures were prepared using Chimera, ⁶⁵ which employs the MSMS algorithm for surface generation. ⁶⁶

contacts in both enzymes, but fills out the hydrophobic end of the binding pocket to a much greater extent in 15-hLO relative to 12-hLO. We believe that the tighter binding in 15-hLO may be due to the favorable van der Waals contacts that form at the bottom of the binding pocket, as well as, potentially, an unfavorable free energy contribution in 12-hLO arising from *not* filling the binding pocket. That is, we believe that the pose of this ligand docked in the homology models provides a post facto but reasonable explanation for the observed selectivity of compound 15.

More broadly, 15 of the 20 selected compounds, including the 3 for which IC₅₀s were determined, showed somewhat greater inhibition of 15-hLO relative to 12-hLO, although the differences were small in some cases. There are several possible explanations, which are all admittedly speculative in the absence of further investigation. The first is that 15-hLO is more closely related structurally to the template 15-rLO. It should be noted that the docking results against 12-hLO also showed a bias toward 15-hLO selective inhibitors, including compound 15. However, this could still represent a "bias" inherent to creating a homology model based on 15-rLO. A second possibility is that the binding site of 15-hLO is simply more easily targeted due to the smaller size of the binding pocket.

As a control, we also purchased 20 randomly chosen compounds from the ChemBridge library (using a simple computer script employing a random number generator). None of these randomly selected compounds were among the top 1000 compounds from the 12-hLO and 15-hLO docking calculations. No low micromolar inhibitors were identified among the controls, although a few relatively weak inhibitors were identified (Supporting Information Table 1).

Discussion

The primary conclusion of this study is that structure-based virtual screening represents a viable approach to the development of new lipoxygenase inhibitors, despite the fact that no experimental structures are available for human lipoxygenases. Although this is admittedly a relatively small-scale study, with only 20 compounds tested based on the virtual screening results, 3 new low micromolar inhibitors were discovered. These compounds appear to be suitable as leads for further inhibitor development efforts against 12-hLO and 15-hLO, based on the fact their size and chemical properties are generally appropriate to classify them as drug-like compounds (Table 2). This study

Table 2. Chemical Properties of the Low Micromolar Inhibitors

Chembridge ID	ID no. from text	mol weight	$\log P$	rotatable bonds	H bond donors	H bond acceptors
6635967	15	302.75	5.42	4	1	3
5680672	12	306.81	4.64	2	2	1
6640337	9	296.32	3.93	3	3	4

thus joins a relatively small but growing literature in which homology models have been used in conjunction with virtual screening to successfully identify new enzyme inhibitors.

We wish to underscore two methodological strengths of this study. First, in contrast to many reported applications of virtual screening methods, we did not employ human judgment in selecting compounds from the docking "hit list" for experimental testing. Thus, the successful identification of new inhibitors directly reflects the quality of the homology models and the docking algorithms. Second, as a control study, 20 randomly chosen compounds, from the same library used for the virtual screening, were also tested for inhibition of 12-hLO and 15hLO. Although this again is clearly a small-scale study, it is nonetheless reassuring that no low-micromolar inhibitors were identified among the randomly selected compounds. We note also that detergent was used in the assay to help disrupt small molecule aggregates that can lead to false positives. Altogether, our results suggest that virtual screening against the homology models is capable of identifying low-micromolar inhibitors at a rate that, in all likelihood, greatly exceeds random compound selection.

Our ongoing goal is to identify inhibitors that are both potent and *selective* for specific lipoxygenase enzymes. In the work reported here, we can claim, at best, to be only partially successful in this endeavor. One of the low micromolar inhibitors showed selectivity against 15-hLO over 12-hLO. However, this compound was selected from the virtual screen against the 12-hLO homology model. Nonetheless, we were able to use our models to retrospectively rationalize why compound 15 showed selectivity for 15-hLO. Finally, the models of the docked low-micromolar inhibitors suggest clear strategies for exploiting the differences in binding site volume between 12-hLO and 15-hLO and we remain optimistic that our structure-based approach will ultimately be capable of aiding the discovery of new selective inhibitors, including compounds with selectivity for 12-hLO.

Experimental Section

Homology Model Construction. The 12-hLO and 15-hLO homology models were built using the 2.4 Å resolution 15-rLO structure (PDB ID: 1LOX), with the cocrystallized Roche RS7 inhibitor, as a template.³⁴ This is the only publicly available crystal structure of a mammalian lipoxygenase. It has 80% overall sequence identity to 15-hLO and 61% sequence identity to 12-hLO. The template structure was first refined by the addition of hydrogen atoms not present in the crystal structure, as well as the construction of atoms and residues not resolved in the crystal structure. Residues 201-211 and 601-602 lacked electron density and could not be resolved. In addition, the electron densities of residues 177-188 were ambiguous due to the presence of a 2-fold crystallographic symmetry operator. Construction of the missing segments was carried out in the Jacobson group using their in-house software, Protein Local Optimization Program (PLOP, distributed commercially under the name Prime), which uses loop prediction, ⁵³ side chain prediction, 54,55 and energy minimization to construct the missing atoms and residues.

The homology modeling of 12-hLO and 15-hLO was also carried out using PLOP, based on alignments obtained from NCBI-BLAST (blastp).⁵⁶ In brief, the primary structural modifications that must be made during the initial model construction are the closing of chain breaks associated with gaps in the sequence alignment, and side chain optimization (for all residues that are not identical between target and template in the sequence alignment). Chain break closure is accomplished using iterative application of a loop prediction algorithm, which is described in detail elsewhere. 53 After all chain breaks have been closed, side chain optimization^{54,55} and complete energy minimization are performed on all portions of the protein whose coordinates were either not taken from the template at all, or were modified during the model building procedure. The homology modeling procedure uses the OPLS all-atom force field^{54,57,58} and a Generalized Born solvent model^{59,60} for choosing low-energy structures.

Virtual Screening Procedure. Flexible ligand docking was performed using the Glide (Schrödinger, Inc.) program, ^{61,62} which uses a modified version of the Chemscore energy function to score the protein—ligand interactions. ⁶³ This docking algorithm has recently been shown to outperform other docking algorithms in its ability to reproduce cocrystallized poses from a set of 69 ligand—receptor complexes from the PDB. ⁶⁴ Molecules were docked using either the standard precision mode (SP) or the extra precision mode (XP), which uses a more optimized scoring function as well as a more extensive search of ligand confirmations than the SP mode. The resulting poses of the docked compounds were ranked according to their GlideScore.

The modeled structures for 12-hLO and 15-hLO and the 15-rLO crystal structure were prepared for docking by minimizing the energy of each structure with the RS7 inhibitor bound in the active site.

The homology models were built with the active site iron in the active ferric state (Fe^{3+}) with no bound water. The ferrous (Fe^{2+}) form of the enzyme is dominant at equilibrium and is activated by oxidation of 1 equiv of the hydroperoxide product [(9Z-11E)-13-hydroperoxy-9,11-octadecadienoic acid (HPOD)] to the ferric state (Fe^{3+}) . The interaction with the substrate, linoleic or arachidonic acid, reduces the iron from the ferric to the ferrous state. We assumed that the putative inhibitors would act as competitive inhibitors, thereby disallowing the entry of the substrate and subsequently not allowing the iron to be reduced.

The databases of small molecules were prepared for docking using the LigPrep (Schrödinger, Inc) ligand preparation software, which generates a minimized conformation of each ligand, and multiple protonation/tautomerization states when appropriate.

Experimental Determination of IC_{50}s. The experimental screening was performed as previously described. Briefly, the human lipoxygenases were expressed and purified via our standard histagged protein protocol. ²⁷ 12-hLO and 15-hLO were then screened for inhibition by monitoring the rate of formation of the conjugated

diene products at 234 nm via UV spectroscopy. One-point screens were preformed with 20 μ L of a 1 mg/mL solution of inhibitor in DMSO aliquoted into a 2 mL cuvette containing 4 µM of substrate in the appropriate buffer (HEPES pH 8.0 for 12-hLO and HEPES pH 7.5 for 15-hLO) containing 0.01 wt % Triton X-100 to disrupt small molecule aggregates that can lead to false positives.⁴⁶ The percent inhibition (%inh) is given as %inh = $(1 - R_I/R_c)$, where $R_{\rm I}$ is the enzyme rate with the inhibitor present and $R_{\rm c}$ is the control rate of the enzyme. The control is performed as described above in the absence of inhibitor but with the same volume of DMSO. Initially, one-point screens were performed with 0.01% concentration of inhibitor, in duplicate, to assess the effectiveness of inhibitor. If the inhibitor did not display potent inhibition, the IC₅₀ was estimated with the one-point screen, utilizing the hyperbolic equation, $%\inf = [I]/([I] + IC_{50})$, where [I] is the concentration of the inhibitor and %inh is the percent inhibition, assuming 100% inhibition at infinite [I]. Compounds that displayed potent inhibition were screened at multiple inhibitor concentrations (2-60 μ L of a 0.1 to 1 mg/mL solution in DMSO, depending on inhibitor strength) and fit with a standard hyperbolic equation to determine IC₅₀ values and their corresponding error.

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Supporting Information Available: Data on the 20 randomly selected compounds used as a control is provided in supplementary Table 1. The inhibition curves and least-squares fits used to determine IC_{50} s are provided in supplementary Figures 1–3. The coordinates of the homology models are provided as PDB files. This material is available free of charge via the Internet at http://pubs.acs.org.

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