Biofragments: An Approach towards Predicting Protein Function Using Biologically Related Fragments and its Application to Mycobacterium tuberculosis CYP126


We present a novel fragment-based approach that tackles some of the challenges for chemical biology of predicting protein function. The general approach, which we have termed biofragments, comprises two key stages. First, a biologically relevant fragment library (biofragment library) can be designed and constructed from known sets of substrate-like ligands for a protein class of interest. Second, the library can be screened for binding to a novel putative ligand-binding protein from the same or similar class, and the characterization of hits provides insight into the basis of ligand recognition, selectivity, and function at the substrate level. As a proof-of-concept, we applied the biofragments approach to the functionally uncharacterized Mycobacterium tuberculosis (Mtb) cytochrome P450 isoform, CYP126. This led to the development of a tailored CYP biofragment library with notable 3D characteristics and a significantly higher screening hit rate (14%) than standard drug-like fragment libraries screened previously against Mtb CYP121 and 125 (4% and 1%, respectively). Biofragment hits were identified that make both substrate-like type-I and inhibitor-like type-II interactions with CYP126. A chemical-fingerprint-based substrate model was built from the hits and used to search a virtual TB metabolome, which led to the discovery that CYP126 has a strong preference for the recognition of aromatics and substrate-like type-I binding of chlorophenol moieties within the active site near the heme. Future catalytic analyses will be focused on assessing CYP126 for potential substrate oxidative dehalogenation.

Introduction

Predicting the functional roles of proteins remains a major challenge in chemical biology.[1] Advances in high-throughput genome sequencing techniques have seen the rate at which new proteins are identified far exceed that at which they can be biochemically characterized, and it has been reported that less than one percent of proteins have experimentally validated annotations.[1b, 2] The traditional techniques used to assign protein function primarily involve a combination of bioinformatics-based methods, for example, sequence similarity to previously characterized proteins, genomic context, transcriptional patterns, or experimental phenotypes of deletion or knockdown mutants.[1b–d] Newer computational 3D-structure-based methods, involving protein modeling and alignment to find structural similarities (globally or at known or predicted functional sites) and virtual ligand docking, have expanded in recent years but are not yet widely used, and the reliability of docking is also still in question.[1a, 3] For instance, in Mycobacterium tuberculosis (Mtb), the pathogen responsible for tuberculosis (TB) disease, there are 3933 protein-coding genes.[4] Many of these genes are assumed to have essential functions, such as in DNA replication, transcription, translation, and cell-division but this annotation is only on the basis of homologues from other bacteria,[5] and this technique has led to several cases of misassignment.[6] About one third of all Mtb gene products have no functional data at all, assigned as unknown or conserved hypothetical proteins, and without functional classification.[5] For example, in the proline-glutamate (PE)/proline-proline-glutamate (PPE) protein family, which represents about 10% of the Mtb proteome, enzymatic activity has been demonstrated only for LipY (Rv3097c) as a triacylglycerol hydrolase.[7]

Given these challenges, novel approaches for establishing protein function are very much needed. We attempted to...
transfer the technique of fragment-based ligand discovery (FBLD) to this field. FBLD is now an established method for developing small-molecule ligands as chemical tools and leads for drug development.\cite{3b,8} At its heart, this method involves the structure-guided design and synthesis of potent ligands from weak-binding low-molecular-weight fragment molecules (typically < 250 Da).\cite{3b,8} There are two primary advantages to this method: first, because of the low complexity of small fragments, a significantly larger proportion of chemical space can be explored with a relatively small fragment library (usually $10^2 - 10^3$ fragments), compared with the approximately $10^5 - 10^6$ larger molecules ($M_w$ 300–500 Da) typical in a high-throughput screen (HTS).\cite{3b,8} Second, fragment hits must make high-quality interactions with the target to bind with sufficient affinity for detection.\cite{3b}

Herein, we describe a novel fragment-based approach for predicting the function of putative ligand-binding proteins, a method which we have termed “biofragments”. This approach encompasses two main phases: the first phase is the design and construction of more biologically relevant fragment libraries (biofragment libraries) based on known sets of substrate-like ligands for a specific protein class of interest. This step is essential because of the well-recognized disparity between commercial fragment libraries and natural products—commercial fragments have a structural makeup that is largely biased toward readily available small, flat, heterocyclic molecules, whereas natural products have a prevalence of stereogenic centers and even include reactive functional groups.\cite{3b,10}

In the first step of the method, a set of known ligands of both the protein of interest (where available) as well as other related proteins is compiled. Subsequently, a fragment library is assembled that samples the substructural chemical space present in the ligands and is hence expected to increase fragment hit rates. This approach uses the concept of chemogenomics,\cite{10d,e,11} which maps chemical space to biological space in a systematic manner. Herein, we have applied this method to fragments, both in the library design phase and in the step of selecting putative substrate scaffolds.

The second main phase is the fragment-based screening\cite{3b} of the biofragment library for binding to an uncharacterized protein from the same or similar family from which the biofragments were derived. The characterization of hits (at a pharmacophore level or from determination of their exact structural-binding mode) provides insights into the structural determinants for ligand recognition, the probable structural characteristics of endogenous substrate(s), and hence insight into the functional role of the enzyme. In this regard, it has been shown previously that fragments made by breaking down known endogenous ligands display considerable fidelity of their binding mode and interactions, and can provide an understanding of the substructural energetic contributions for binding, to identify hot spots at protein–ligand binding sites.\cite{12}

As a proof-of-concept, our biofragments approach was applied to the functionally uncharacterized $Mtb$ cytochrome P450 enzyme, CYP126. Cytochrome P450 enzymes have a remarkably diverse repertoire of possible catalytic reactions and substrates,\cite{13} making them an ideal model to test our approach. CYP126 (Rv0778) is located near essential $Mtb$ genes encoding enzymes involved in the de novo biosynthesis of purine, and CYP126 is also part of a putative operon with a probable adenylosuccinate lyase, PurB.\cite{5a} However, CYP126 also shares notable homology (35% identity) with the $Mtb$ cholesterol hydroxylases CYP124 and 125, and it is highly conserved across actinobacteria (including both pathogenic and nonpathogenic...
strains), which suggests that it may participate in an important general function. Further information about the function of CYP126 is, at this stage, not available, which made it a suitable test system for our approach.

To investigate CYP126, we constructed a CYP biofragments library and, through fragment-based screening, biofragment hits were identified that make both substrate-like type-I and inhibitor-like type-II interactions with the enzyme. A chemical-fingerprint-based substrate model was built from the hits and used to search a virtual TB metabolome, this led to the discovery that CYP126 preferentially binds chlorophenol scaffolds in a substrate-like fashion, close to the heme, and suggests that its endogenous substrate may maintain a similar motif upon further interrogation.

Results and Discussion

Design, construction, and analysis of a CYP biofragment library

A CYP-focused biologically relevant fragment library (“CYP biofragment library”) was designed starting from knowledge of CYP ligands found in the protein data bank (PDB; Figure 1). The PDB was used so that structural binding information would be available for comparison to potential biofragment-bound crystal structures. All PDB CYP ligands were extracted and ions, salts, gases, buffer additives, heme groups or other metal complexes, and azoles were removed, along with any ligands labeled as drugs or inhibitors. The in silico filtering process retained 43 bacterial and 24 eukaryotic CYP substrates and substrate-like/mimetic ligands (see Table S1 in the Supporting Information), and these were grouped into six major structural classes (see Figure S1).

Next, a database of commercially available fragment molecules was created by compiling the ChemBridge, ChemDiv, Enamine, Life Chemicals, Ryan Scientific, and Specs supplier datasets (3646719 unique compounds) from the ZINC repository[16] and filtered for those molecules with a predicted solubility in water greater than 1 mM[17] and a molecular weight $M_w < 250$ Da or $M_w < 350$ Da. The two most similar commercial fragments representing each bacteria-

al CYP ligand (henceforth termed CYP biofragments) were then found by passing the fragment database over the ligand database and calculating the Tanimoto similarity coefficient[18] for every combination. CYP biofragments that would not be amenable to 1D $^1$H NMR spectroscopy screening (i.e. no down-field proton NMR signal $>2$ ppm) were manually excluded, and the biofragments were purchased from their respective suppliers. In total, 63 biofragments were compiled in this way for experimental screening.

The composition of the new biofragment library was compared to a conventional commercial drug-like fragment library in regards to physicochemical properties and chemical diversity, using analytical procedures reported previously[10] (Figure 2). Percentage frequency distributions of the libraries by $M_w$ and $c\log P$ (calculated logarithm of partition coefficient) show similarities in size and hydrophilicity based on the fragments rule-of-three constraints used in the construction of both libraries (Figure 2 A,B). A bimodal distribution is observed for the $M_w$ of the biofragments (Figure 2 A), and this is probably reflective of the two commercially available fragment databases that were combined during the process of biofragment design ($M_w < 250$ or $< 350$ Da). Significant contrast was observed when plotting the library distributions of the ratio of...
sp²- to sp³-hybridized carbons in each molecule (Figure 2C). The traditional commercial fragments contain a high proportion of sp² carbons, primarily from aromatics, but the CYP biofragments are richer in sp³ centers. A principal moment-of-inertia (PMI) plot\cite{10f,19} corroborates this finding (Figure 2D), showing the conventional fragments lying heavily in 2D space, whereas the biofragments have more uniform coverage of molecular shape and structural diversity (by visual inspection). This is a direct consequence of designing the library to resemble nonaromatic natural CYP ligands (see Figure 1 and Figure S1). This theme also appears in the context of drug discovery, where more challenging drug targets (e.g., protein–protein interactions) have pushed efforts towards improving the structural diversity of inhouse fragment libraries and also towards increasing the three-dimensional structures available\cite{3b,10c,11}. Overall, the above analysis affirms the differing properties of biologically relevant fragment libraries that originate from the first phase in the biofragments approach; the importance of this will be shown later by significantly increased hit rates.

### Biofragment screening by NMR spectroscopy

The 63 CYP biofragments were screened for binding to CYP126 by CPMG,\cite{20} STD,\cite{21} and WaterLOGSY\cite{22} ligand-detect ed 1D ¹H NMR spectroscopy. The antifungal agent, ketoconazole, is known to bind to CYP126 in a typical azole–heme type-II coordination mode by using heme absorbance shift assays ($K_d = 1.4 \mu M$; K. J. McLean et al., unpublished results), and ketoconazole was used here in displacement experiments to indicate hits that might bind within the CYP126 active site (displacement in CPMG only). Nine hits were identified in total by NMR that were displaced by ketoconazole from the active site (Figure 3 A,B). This represents a hit rate for the screen of 14%, which is significantly higher than for a standard commercial drug-like fragment library that was screened previously against Mtb CYP121 and 125 (4 and 1%, respectively\cite{23}). This confirms that biologically related small molecules would be more likely to bind and be recognized by biomacromolecular targets.\cite{184} The hits identified by ketoconazole displacement were all aromatic compounds, suggesting that CYP126 might preferentially recognize aromatic moieties within its catalytic site.

### Biofragment screening by heme absorbance shift

To give further information on the type of interaction the biofragments make within the CYP126 active site and whether they bind in close proximity to the heme, a spectrophotometric heme-absorbance-shift assay\cite{33a,32a} was performed for all nine hits. Two biofragments (BIO-A7 and BIO-B10) were found to induce a type-II red shift in the CYP126 heme Soret absorbance peak (likely from heme-coordination; see Figure S2), and a single hit (BIO-B6) gave a substrate-like type-I blue shift (Figure 3 C). Looking at the structures of these hits, we envisaged that BIO-A7 could coordinate the heme iron atom through its arylamine and BIO-B10 possibly through the lone pair of the thioether sulfur atom.\cite{24} It is intriguing that CYP126 could support a thioether coordination mode because this has been noted as a rather unusual CYP binding interaction.\cite{24} Of greatest interest, however, is the type-I shift induced by the chlorophenol BIO-B6, which is typically associated with substrate-dependent displacement of the weakly
bound resting water molecule from the distal position of the CYP heme iron atom for the first step of the CYP catalytic cycle.[34] There are only four chlorophenols in our traditional drug-like fragment library (0.30% of the total library size), which illustrates the difficulty of reaching the same conclusion from a conventional library. Subsequent efforts were focused on exploring the basis of this BIO-B6–CYP126 interaction. In particular, we attempted to experimentally characterize the BIO-B6–CYP126 structural binding interaction by crystallography, but we were only able to obtain native CYP126 crystals with the enzymatic active site occluded by a dimer interface (K. J. McLean et al., unpublished results).

Construction, search, and selection of a virtual TB metabolome and screening by using a heme-absorbance-shift assay

We hypothesized that fingerprint-based characterization of the biofragment hits (particularly BIO-B6) could be used to provide insights into possible classes of substrate for CYP126 based on previously known TB metabolites. A virtual TB metabolome was constructed from the KEGG PATHWAY database[25] and searched by Tanimoto similarity to BIO-B6. An additional Naïve Bayes substrate model based on Molprint2D fingerprints of all nine NMR hits was constructed[26] and used as a secondary search. Twenty-three TB metabolite matches (0.60% of the total metabolome) were then screened against CYP126 by heme-absorbance-shift assay as described above for the original biofragment hits (Figure 4). This screen identified a substrate-like type-I hit, pentachlorophenol TB23, and a chloroaniline type-II hit, TB8 (Figure 4B and Figure S3). A complete titration heme-absorbance-shift assay was performed for TB23, successfully confirming it has high binding affinity and ligand efficiency (Figure 4B; $K_d = 150 \, \mu \text{M}$, $LE = 0.43 \, \text{kcal mol}^{-1} \text{NHA}^{-1}$). Both of these metabolites are aromatic chlorobenzenes, like biofragments BIO-B6 and BIO-A7, which further supports an apparent preference by CYP126 for binding this type of motif close to the heme and implies that this scaffold could be consistent in the endogenous substrate. BIO-B6 and TB23, which induce a substrate-like type-I shift, are both chlorophenols. The selection of the negatively charged pentachlorophenol TB23 over the other similarly assayed non-heme-coordinating chlorophenols and benzyl alcohols is also intriguing (Figure 4A), which indicates that the additional electron-withdrawing chlorine atoms promote CYP126 active-site recognition. While an exact pentachlorophenol metabolite pathway has not been explicitly reported for Mtb, the pathway appears as part of the Mtb H37Rv KEGG dataset, based on extension of the well-known chlorocyclohexane and chlorobenzene biodegradation processes found in microorganisms from diverse environments.[27] In this pathway, pentachlorophenol is dehalogenated oxidatively to tetrachlorohydroquinone.[27b, d, 28] This metabolic process has recently been shown to be CYP-mediated in the white rot fungus, *Phanerochaete chrysosporium*,[28a] by human cytochrome P450 3A4,[28b] and CYP involvement is also suspected in *Mucor ramosissimus*[29] and *Mycobacterium chlorophenolicum.[30]*

Figure 4. TB metabolite screening against *Mtb* CYP126 by heme-absorbance-shift assay. A) Commercially available TB metabolites matched from the fingerprints represented by the CYP biofragment NMR hits. Only those matches that were viable for screening by heme absorbance shift (see the Experimental Section) are shown. Chemical formulas are drawn in their predominant protonation state at physiological pH. B) Absorbance difference spectra for CYP126 (5 \, \mu \text{M}) titrated with various concentrations of TB23. The substrate-like type-I blue shift in the Soret absorbance band of the heme induced by TB23 is shown as a $\Delta A_{409} \text{nm}$ and $\Delta A_{427} \text{nm}$ in the difference spectra. Inset: the shift in the absorbance band of the heme (quantified as $\Delta A_{409} \text{nm} - \Delta A_{427} \text{nm}$) as a function of TB23 concentration (squares) with the one-site binding equilibrium model fitted for calculating the $K_d$ (line).
Conclusions

Herein, we presented a novel fragment-based approach, biofragments, to help address the fundamental problem in biology of assigning function to proteins. The approach has two main stages: first, a biologically relevant fragment library (biofragment library) is constructed based on known sets of substrate-like ligands for the protein class of interest. Second, the biofragment library is screened (fragment-based screening cascade) for binding to a novel putative ligand-binding protein from that class. The characterization of hits provides insights into the basis of ligand recognition, selectivity, and function.

We applied this approach to the functionally uncharacterized Mtb CYP isoform, CYP126. The designed CYP biofragment library had notable 3D characteristics and its screening hit rate against CYP126 (of 14%) was significantly higher than for the conventional drug-like fragment libraries versus Mtb CYP121 and 125 (which were 4% and 1%, respectively[23]). Overall, the biofragment hits and their follow-up TB metabolites (found by a chemical fingerprint-based substrate model search of a virtual TB metabolome), indicate a strong preference for the recognition of aromatics by CYP126 and the substrate-like type-I binding of chlorophenol moieties within the active site near the heme. Future studies will assess CYP126 for potential substrate oxidative dehalogenation or similar CYP-mediated reactions.[13, 27b, 28a, 31] These findings confirm our hypothesis that biologically related fragments are more likely to bind to biomacromolecules, and suggest that biofragments could also be used to improve the hit rates for screens against traditionally difficult target classes, such as protein–protein interactions or structured nucleic acids.[30] In summary, we conclude that the biofragments approach is a novel method to deconvolute protein function, adding to the previously established value of biofragments approach is a novel method to deconvolute protein function, adding to the previously established value of biofragments approach is a novel method to deconvolute protein function, adding to the previously established value of biofragments approach is a novel method to deconvolute protein function.

Experimental Section

All biological and computational experimental methods are given in the Supporting Information.

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