De novo protein fold families expand the designable ligand binding site space

Xingjie Pan1,2*, Tanja Kortemme1,2,3,4*

1 Department of Bioengineering and Therapeutic Sciences, University of California San Francisco, San Francisco, California, United States of America, 2 UC Berkeley–UCSF Graduate Program in Bioengineering, University of California San Francisco, San Francisco, California, United States of America, 3 Quantitative Biosciences Institute (QBI), University of California San Francisco, San Francisco, California, United States of America, 4 Chan Zuckerberg Biohub, San Francisco, California, United States of America

* xingjiepan@gmail.com (XP); tanjakortemme@gmail.com (TK)

Abstract

A major challenge in designing proteins de novo to bind user-defined ligands with high affinity is finding backbones structures into which a new binding site geometry can be engineered with high precision. Recent advances in methods to generate protein fold families de novo have expanded the space of accessible protein structures, but it is not clear to what extent de novo proteins with diverse geometries also expand the space of designable ligand binding functions. We constructed a library of 25,806 high-quality ligand binding sites and developed a fast protocol to place (“match”) these binding sites into both naturally occurring and de novo protein families with two fold topologies: Rossman and NTF2. Each matching step involves engineering new binding site residues into each protein “scaffold”, which is distinct from the problem of comparing already existing binding pockets. 5,896 and 7,475 binding sites could be matched to the Rossmann and NTF2 fold families, respectively. De novo designed Rossman and NTF2 protein families can support 1,791 and 678 binding sites that cannot be matched to naturally existing structures with the same topologies, respectively. While the number of protein residues in ligand binding sites is the major determinant of matching success, ligand size and primary sequence separation of binding site residues also play important roles. The number of matched binding sites are power law functions of the number of members in a fold family. Our results suggest that de novo sampling of geometric variations on diverse fold topologies can significantly expand the space of designable ligand binding sites for a wealth of possible new protein functions.

Author summary

De novo design of proteins that can bind to novel and highly diverse user-defined small molecule ligands could have broad biomedical and synthetic biology applications. Because ligand binding site geometries need to be accommodated by protein backbone scaffolds at high accuracy, the diversity of scaffolds is a major limitation for designing new ligand binding functions. Advances in computational protein structure design methods have significantly increased the number of accessible stable scaffold structures. Understanding
how many new ligand binding sites can be designed into the de novo scaffolds is important for engineering novel ligand binding proteins. To answer this question, we constructed a large library of ligand binding sites from the Protein Data Bank (PDB). We tested the number of ligand binding sites that can be designed into de novo scaffolds and naturally existing scaffolds with the same fold topologies. The results showed that de novo scaffolds significantly expanded the potential ligand binding space of their respective fold topologies. We also identified factors that affect difficulties of binding site accommodation, as well as the relationship between the number of scaffolds and the accessible ligand binding site space. We believe our findings will benefit future method development and applications of ligand binding protein design.

Introduction

Ligand binding is a major class of protein functions, and the ability to design ligand binding de novo has many important applications [1] such as engineering of biosensors and ligand-controlled protein functions [2, 3]. Naturally occurring proteins recognize their cognate ligands with high affinity and specificity using defined three-dimensional geometries of binding sites with high shape complementarity between ligands and proteins. For the formation of favorable hydrophobic and polar interactions, the chemical groups on the protein must be placed at specific spatial positions relative to the ligand [4–6]. Designing new ligand binding proteins therefore requires the ability to build binding sites with defined geometries into stable protein scaffolds that can accommodate the desired interaction geometry with high accuracy. While this approach has led to the successful design of enzymatic activity [7, 8], ligand binding proteins [9, 10], and biosensors [2, 3, 11], it has been limited by both the availability of defined binding site geometries and stable protein scaffolds into which these binding sites can be designed [3].

Several methods have recently been developed to address the first problem, increasing the number of potential ligand binding sites one could generate. The RIF docking method [12] generates ensembles of billions of side chain placements that make defined hydrogen-bonding and non-polar interactions with a target ligand. Other methods [13, 14] use statistics from the protein data bank (PDB) to find three-dimensional placements of amino acid residues that form favorable interactions with fragments of a ligand, which can then be assembled into complete binding site geometries. Protein-ligand interactions defined by these methods have been built successfully into a de novo designed beta barrel [12], and a parametrically designed helical bundle [14].

Naturally occurring proteins solve the second problem, finding a suitable protein backbone to accommodate a specific binding site geometry, not by using a different fold for each function but instead by evolving structural variation in existing protein fold families. This variation allows proteins with the same fold topology (identity and connectivity of secondary structure elements) to tune the precise geometry of binding sites to recognize diverse ligands [15]. This strategy has recently been mimicked by advances in computational protein design methods. These methods have generated de novo designed protein fold families with large numbers of diverse geometries [16, 17], which have significantly expanded the accessible designable protein structure space. The stabilities of designs were validated by computational filters or high-throughput yeast surface display assays, and, for a selected subset, by biophysical assays with purified proteins. In the cases where structures of designs were solved, the experimental structures typically closely agreed with the design models, although the possibility of conformations...
different from the design model cannot be excluded for de novo designed scaffolds without available structures. Nevertheless, the de novo designed proteins could be useful as scaffolds for engineering new functions, but the extent to which de novo fold families could improve binding site design has not been explored. Understanding the relationship between the space occupied by protein structures, and the space available to support different functions, is important for developing methods to design proteins de novo that can bind to novel and highly diverse user-defined ligands.

Here we studied the ability of native and de novo fold families to support a large number of different ligand binding sites. We built a high-quality ligand binding site library from high resolution protein crystal structures. We then matched the binding site library to members of protein folding families using two protocols: a newly developed “fast matching” protocol and the standard method for matching in the Rosetta program for structure modeling and design [5]. “Matching” here refers to the ability to design new binding site residues forming a user-defined binding site geometry into a protein “scaffold”, distinct from the problem of comparing already existing binding pockets. We calculated the number of matched binding sites for four fold families with two different topologies. We studied the effects of binding site sizes (number of binding site residues), ligand sizes and primary sequence separation of binding site residues on the matching success rates and determined the increase of numbers of matches with increasing number of scaffolds in each fold family. Together, we show that de novo fold family design is a promising approach to broaden the scope of designable ligand binding sites.

Results

We first constructed a library of ligand binding sites from native proteins in the PDB. We extracted 25,806 ligands that have at most 100 heavy atoms as well as the ligand binding site residues from 23,238 cluster representative structures from the PDB95 database [18] where chains from the protein data bank are clustered at 95% identity (Methods). The extracted ligands have between 1 and 93 heavy atoms (Fig 1A and 1B). 80.6% percent of the ligands have 13 or fewer heavy atoms, and 7,335 (28.4%) of the ligands have only 1 heavy atom. There are 2,461 unique ligand types in the 25,806 binding sites. The distribution of ligand type frequencies has a long tail (S1 Table). There are 33 frequent ligand types that appear in over a hundred binding sites, while 1,817 ligand types only appear in single binding sites. The frequent ligand types include common crystallographic additives such as glycerol; 1,2-ethanediol; ions such as SO$_4^{2-}$ and Mg$^{2+}$; and cofactors such as heme and flavin adenine dinucleotide (FAD). Ligands that appear in multiple binding sites are seen as vertical stripes in Fig 1B. We also clustered the binding sites by similarities of ligand 3D structures using the extended 3-dimensional fingerprint (E3FP) [19] and Butina clustering [20] (Methods). The ligands form 4232 clusters. Binding sites have between 2 and 41 residues, with 81.2% of the binding sites having 7 or fewer binding site residues. The number of protein residues in binding sites scales linearly with the number of ligand heavy atoms, with a slope 0.35 (Fig 1B). The frequencies of amino acid types in binding sites are different from those for whole proteins reported by UniProtKB/Swiss-Prot (Fig 1C). We defined the enrichment ratios of amino acids as their frequencies in ligand binding sites divided by their frequencies in whole proteins. The large aromatic side chains Trp, Tyr and Phe are the top 1, top 3 and top 6 enriched amino acid residues, respectively. His, characterized by its ability to coordinate metal ions and to catalyze chemical reactions, is the second most enriched amino acid residue. Asp and Arg are the 4th and 5th enriched amino acid residues, which may play important roles in interacting with charged ligands. Binding sites with single heavy atom ligands have different amino acid preference than those binding to ligands with at least two heavy atoms (Fig 1D). For the binding
sites with single heavy atom ligands, the negatively charged residues Asp and Glu, which can form favorable electrostatic interactions with positively charged metal ions, are highly enriched. The enrichment ratios of Asp and Glu are 4.6 and 2.2, respectively. When excluding ligands with only a single heavy atom, the top 5 enriched residues that bind to ligands with at least 2 heavy atoms are Trp, His, Tyr, Phe and Arg (S1 Fig).

The binding site library is useful for testing the ability of protein fold families to support ligand binding sites. A protein scaffold can in principle support a ligand binding site if the binding site residues can be built onto the scaffold such that the key interactions between the ligand and binding site protein residues are preserved. The Rosetta matcher protocol [5] has been shown to be successful in matching ligand binding sites to protein scaffolds [9]. However, the Rosetta matcher is too slow to match tens of thousands of binding sites to thousands of

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scaffolds because it samples all possible side chain rotamers of binding site residues. Therefore, a fast method to test compatibility between binding sites and scaffolds is required. Methods have been developed for comparing binding pocket structures [21, 22]. However, testing the ability of a scaffold to support a ligand binding site for design purposes is different from comparing binding pocket structures. In a ligand binding site design problem, the original side chains on a scaffold will be redesigned, so the matching method should only match the backbone atoms of binding sites at high precision. Additionally, the matching method needs to ensure that there are no steric clashes between binding site side chains newly placed into the scaffold and the scaffold backbone. Given the limitations of existing methods, we developed a new fast match protocol (Fig 2A) to perform all-against-all matching between the library of ligand binding sites and the sets of scaffolds. In the fast match protocol, the binding site is anchored and matched as a rigid body (Methods). This rigid body approximation drastically improved the matching speed. We tested the run time by matching the binding site library to the native NTF2 fold family (CATH superfamily 3.10.450.50) [23]. The mean time to find a successful standard Rosetta match is 706s while the mean time of a successful fast match is 3.1s. As a trade-off, the rigid body approximation of the fast match method may discard binding sites that can be matched by the Rosetta matcher using alternative side chain rotamers. Therefore, in this study we focused on matching ligand binding sites using the side chain rotamers present in the original ligand binding site in the PDB. Using these original rotamers also let us directly compare the backbone geometries in the native binding sites and the backbone geometries in our scaffold libraries.

We matched the binding site library to backbone scaffolds of de novo designed Rossmann and NTF2 protein fold families generated by the loop-helix-loop unit combinatorial sampling (LUCS) method [16], as well as the two native fold families with the same topology from the CATH database [23] (Fig 2B, Methods). While some binding sites in the library have the same ligand type, we matched all binding sites because they may represent different ligand or binding site conformations. To determine if a fold family can support a given ligand binding site, we first used fast match to match the ligand binding site to all protein scaffolds in the family. Then we used the Rosetta matcher to match the binding site to the scaffolds that passed the fast match (Methods). If a binding site can be matched to at least one scaffold in a fold family by the fast match method, we call it a fast matched binding site of the fold family. If a binding site can be matched to at least one scaffold in a fold family by both the fast match and the Rosetta matcher, we call it a Rosetta matched binding site of the fold family. We focused on studying whether a binding site can be supported by at least one member from each fold family. Therefore, to limit computational time, once the Rosetta matcher found a match for a given binding site, we skipped matching to further scaffolds in the same family. Since we used stringent matching criteria (Methods), the matched binding sites in the scaffold closely recapitulated the interactions between the ligands and binding site residues in the original protein structures from which the binding sites were derived (Fig 2C).

Between 5896 and 7548 binding sites could be successfully matched by the Rosetta matcher to each fold family when considering all binding sites (Table 1). The number of binding site residues was the major determinant of the matching success rate (Fig 3). For the de novo Rossmann fold family, the success rates for 2, 3 and 4 protein residue binding sites were 93.8%, 33.4% and 6.5%, respectively. Only 13 binding sites with 5 or 6 residues could be matched. There was no match for binding sites with more than 6 protein residues. Similar dependencies on binding site sizes were observed across the 4 different protein fold families (Table 2). Overall, the matching success for the de novo Rossmann family is larger than for the native Rossmann family, for all binding site sizes. However, this is not the case when comparing the de novo NTF2 with native NTF2 fold families. A possible explanation for this finding is that there
are fewer native Rossmann fold scaffolds, so that the possible improvement using de novo scaffolds of this topology is larger.

Because almost all 2-residue binding sites could be matched and the matching success rates were low for binding sites with more than 3 residues, we used 3-residue binding sites to further

Fig 2. Matching ligand binding sites to scaffold libraries. A. Schematic of the matching protocol. The ligand is represented as a yellow triangle. The ligand-binding site as a rigid body (green) is first matched to the scaffold (grey) by anchoring to a scaffold residue shown in the black circle. Then the binding site residues are aligned to the corresponding scaffold residues. Finally, the standard Rosetta matcher is applied to build the binding site side chains (magenta) onto the scaffold. B. The binding sites are matched to native and de novo designed scaffold families with Rossmann or NTF2 fold topologies. C. Examples of matches. The coloring scheme is the same as A.

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study properties of successful matches. We constructed a new library of binding sites that all have 3 protein residues (Methods). We used the Rosetta energy function, which was tested on protein ligand interactions [3, 9, 12, 24], to choose the 3 residues that have strongest predicted interactions with each ligand. Although other residues in a binding site can also contribute to ligand binding, previous studies have shown that it is possible to design binding sites by matching 3 or 4 residues followed by optimizing the surrounding residues with side chain design algorithms [8, 9]. We clustered the ligands in the 3-residue binding site library using the clustering method used above (Methods). The ligands form 4,184 clusters. We matched the 3-residue binding sites to the scaffold libraries using the same protocol as described above. The number of successfully Rosetta-matched 3-residue binding sites ranged from 2,142 to 3,715 (Table 1).

For the successfully matched 3-residue binding sites, we first analyzed the positions of matches relative to the surface of the scaffolds. For each scaffold, we used the Rosetta Layer residue selector [25] to assign layers to all of its residues in a side chain independent manner because the side chains surrounding the matched binding sites are usually redesigned after matching in actual ligand binding protein design problems (Methods). Residues on the surfaces of scaffolds were assigned to the surface layer; residues with many neighbors (defined as weighted neighbor counts greater than 5.2, see Methods) were assigned to the core layer; and the rest of residues were assigned to the boundary layer (Fig 4A). Core layer residues are often not completely buried, i.e. are accessible to ligands. In all of the fold families, surface layer residues were most abundant, which accounted for 47%-63% of all residues. 29%-39% residues were in the boundary layer and 6%-20% residues were in the core layer (Fig 4B). NTF2 fold proteins had more surface layer residues which was likely due to the pocket of this fold. We defined the layer of each residue in a matched binding site as the layer of its matched scaffold residue position. The frequencies of matched residue layers are similar to the frequencies of scaffold layers (Fig 4C). To evaluate the positions of matches at the binding site level, we defined a depth score for each matched binding site. The depth score of a matched binding site is the number of boundary residues plus two times the number of core residues. The depth scores for binding sites matched to different fold families had similar distributions (Fig 4D). 20%-27% binding sites were entirely matched to protein surfaces and had depth scores of 0. The remainder of matched binding sites were buried to some extent. The majority of binding sites were in shallow pockets with depth scores ranging from 2 to 4. Only 8%-12% binding sites were matched to core layer positions with depth scores of 5 or 6. For comparison, ligand-

| Binding site library | Match type | Native Rossmann \(N^a = 20\) | Native NTF2 \(N^b = 103\) | De novo Rossmann \(N^b = 1,000\) | De novo NTF2 \(N^b = 1,000\) | De novo Rossmann \(n^c = 20\) | De novo NTF2 \(n^c = 103\) |
|----------------------|------------|-------------------------------|-----------------------------|-----------------------------|-----------------------------|-------------------------------|-------------------------------|
| All binding sites \(N^a = 25,806\) | fast       | 6860 (248)\(^e\)             | 8761 (795)                  | 9034 (2442)\(^d\)           | 8909 (943)                  |                                |                                |
|                      | Rosetta    | 5896 (212)                    | 7450 (580)                  | 7475 (1791)                 | 7548 (678)                  |                                |                                |
| 3 protein residue binding sites \(N^a = 20,102\) | fast       | 3556 (324)                    | 5714 (1306)                 | 6537 (3305)                 | 6128 (1720)                 | 2544\(^f\) (2744)\(^f\)     | 3864\(^f\) (3977)\(^f\)      |
|                      | Rosetta    | 2142 (199)                    | 3541 (807)                  | 3715 (1772)                 | 3686 (952)                  | 1500\(^f\) (1677)           | 2395\(^f\) (2482)           |

\(^a\) Total number of binding sites in the library.  
\(^b\) Total number of scaffolds in the fold family.  
\(^c\) Numbers in parentheses are binding sites that cannot be matched to de novo scaffolds with the same topology.  
\(^d\) Numbers in square brackets are binding sites that cannot be matched to native scaffolds with the same topology.  
\(^e\) Numbers of scaffolds in randomly resampled subsets of scaffolds.  
\(^f\) Average numbers of matches to 100 randomly resampled subsets of scaffolds.  
\(^g\) Numbers in curly braces are from the best subset of scaffolds in the 100 randomly resampled subsets.
binding proteins that were successfully designed computationally to date can have binding site residues in any of the three layers [8, 9, 14], including cases where residues that mediate designed ligand interactions all belong to the surface layer [26] or the core layer [27]. Proteins

![Histograms of numbers of matches vs binding site sizes (number of protein residues in the binding site). Binding sites that cannot be matched to any scaffold are shown in blue. Binding sites that can be matched to at least one scaffold by the fast match method but cannot be matched by the standard Rosetta matcher are shown in orange. Binding sites that can be matched to at least one scaffold by the standard Rosetta matcher are in green. A-D. Results for 4 scaffold libraries; scaffold sets are indicated in each panel title.](https://doi.org/10.1371/journal.pcbi.1009620.g003)

**Table 2. Dependency of matching success on binding site size (number of protein residues).**

| Binding site size | Native Rossmann | Native NTF2 | De novo Rossmann | De novo NTF2 |
|-------------------|-----------------|-------------|------------------|--------------|
|                   | success count   | success rate| success count    | success rate |
| 2                 | 4590            | 80.9%       | 5340             | 94.2%        |
| 3                 | 1182            | 21.4%       | 1792             | 32.5%        |
| 4                 | 118             | 2.7%        | 272              | 6.3%         |
| 5                 | 6               | 0.2%        | 38               | 1.4%         |
| 6                 | 0               | 0.4%        | 1                | 0.06%        |
| 7                 | 0               | 0.2%        | 0                | 0%           |

![Table 2](https://doi.org/10.1371/journal.pcbi.1009620.t002)
with binding sites that are largely buried in the core can be destabilized in the absence of the ligand; this property has been exploited to engineer protein systems capable of ligand sensing \[2, 28, 29\]. We also did an analysis on the secondary structures of the matched binding sites, which showed that alpha helices were enriched in the matched binding sites (S2 Fig).

We then tested factors that affect the “matchability” of 3-residue binding sites. We calculated the number of binding sites that can be matched to two of the fold families, which are termed as overlapping binding sites. (Fig 5A). If matching to one fold family is independent from matching to another fold family, the probability of overlapping binding sites should be the product of the probabilities of matching to each fold family. We compared 4 pairs of scaffold libraries (Fig 5A): de novo designed Rossmann folds versus de novo designed NTF2 folds (top left) or versus native Rossmann folds (top right), and native NTF2 folds versus de novo designed NTF2 folds (bottom left) or versus native Rossmann folds (bottom right). For all 4 pairs of scaffold libraries, the observed number of overlapping binding sites was significantly
higher than the number of expected overlapping binding sites (chi-squared test p-value < 10^{-300}). This result indicates that some binding sites had higher matchabilities (probabilities to be matched to multiple scaffold libraries). Similar results were observed at the ligand cluster level (S3 Fig). We investigated the contribution of ligand sizes to binding site matchabilities. As expected, the matching success rates for 3-residue binding sites decreased with an increase of the number of ligand heavy atoms (Fig 5B), likely because larger ligands are more likely to clash with the scaffold backbones. We also hypothesized that binding sites whose residues have larger separations in primary sequences are more difficult to match. To confirm that non-local binding sites are harder to match, we calculated the mean inter-residue primary sequence distances for each 3-residue binding site and plotted the mean distances against the matching success rates (Fig 5C). When the 3-residues in a binding site were consecutive in primary sequence, the mean primary sequence distance was 1.33, and the matching success rates were higher than 80%. The success rates dropped rapidly with the increase of mean distance and reached a plateau at low match success rates when the mean distance reached 70.
Next, we studied how the number of matched 3-residue binding sites grew with an increase of the number of scaffolds in fold families (Methods). The log of the number of matched binding sites scaled linearly with the log of the number of scaffolds (Fig 6A–6D). This power law relationship was valid for both the number of fast matches and Rosetta matches across the 4 different fold families. The powers of the power law functions (slopes of the log-log plots) ranged from 0.184 to 0.298. Since the powers were small, the increase of matches progressively diminished as the number of scaffolds got large. Because there is a limited number of designable structures for each fold family, the power law relationship cannot continue indefinitely, but it can still provide a reasonable estimation of the upper bound of the number of matches. Extrapolating the de novo fold family power law relationships to the number of representative structures from the PDB95 database, i.e., 23,238 structures, the numbers of expected Rosetta matches for the Rossmann fold family and the NTF2 fold family would be 7,346 and 6,640. Based on this analysis, the extrapolated numbers of matchable binding sites are still much smaller than the number of total binding sites, highlighting the importance of having diverse fold topologies for different functions.

Since there are many more de novo designed than native scaffolds in each fold family, we also analyzed the number of matches to subsets of de novo fold families that have same numbers of scaffolds as their corresponding native fold families (20 and 103 for the Rossmann and NTF2 fold families, respectively). We used fast match and the Rosetta matcher to match the 3-residue binding site library to all scaffolds in the de novo fold families (in contrast to previous simulations, we did not skip any matching jobs even when the Rosetta matcher already found a match). For the de novo Rossmann fold family, we randomly picked 20 scaffolds and calculated the number of matches. We resampled 100 times and calculated the mean and maximum of the numbers of matches. For the de novo NTF2 fold family, we did a similar analysis, but picked 103 scaffolds for each round of resampling. In this analysis, the de novo fold sub-families supported fewer binding sites than the native fold families (Table 1, right two columns).

This observation can be partly explained by the fact that the structural variation in native fold families is larger than the variation of the de novo fold families that had only a part of the fold (two helices) varied (S4 Fig). However, larger regions of de novo proteins could be varied; when considering only the LHL regions in the de novo scaffolds that were varied between designs, the structural variations were large [16] (S4 Fig).

Finally, to understand how de novo scaffolds expand protein function space, we studied the binding sites that can be matched to de novo scaffolds but not to the native scaffolds of the same topology. We plotted the number of binding sites that were matched to only de novo fold families versus the number of de novo scaffolds (Fig 6E and 6F). For each topology, there are more than 1,000 binding sites that are exclusively matched to de novo scaffolds. These relationships also follow power law functions. The slopes are larger than the slopes of the total matches (Fig 6A–6D), indicating that binding sites that can match to both native and de novo fold families saturate quickly.

**Discussion**

Advances in computational protein structure sampling methods [30] have expanded the accessible structure space of de novo designed proteins. In particular, two recently developed computational methods [16, 17] are capable of engineering de novo protein families that contain defined variations in geometry of proteins that share the same overall fold topology. We probed the functional implications of de novo protein fold families generated by the LUCS method [16] by matching known ligand binding sites to both native and de novo fold families. The de novo Rossmann fold family can support many more binding sites than the native
Fig 6. Numbers of matches scale as power-law functions of numbers of scaffolds in fold families. A-D. Log-log plots of the number of 3-residue matches vs the number of scaffolds. E-F. Log-log plots of the number of 3-residue binding sites that can only be matched to de novo scaffolds of specific topologies vs the number of scaffolds. The black lines represent linear fits.

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Rossmann fold family. The de novo and native NTF2 fold families have comparable total numbers of matched binding sites, but there are hundreds of binding sites that can only be matched to the de novo NTF2 fold family. These results show that LUCS generated structures expand both the accessible protein structure and the accessible protein function space for potential protein engineering applications. The number of matched binding sites increased as a power law function of the number of scaffolds. This relationship allowed us to estimate the upper bound of matches as the number of scaffolds grew and shows that, in addition to geometric variation, different fold topologies are necessary to support diverse functions.

Computational design studies have shown that ligand binding proteins can be designed by matching binding sites with 3 or 4 residues followed by designing second shells of residues to stabilize the binding sites [9]. However, the possible conformational space for even only 3 or 4 binding site residues is large. Consider the relative conformation between a ligand and the backbones of its binding site residues. Each residue backbone can be considered as a rigid body and has 6 degrees of freedom (3 translational and 3 rotational). Therefore, the conformational spaces of 3 or 4 residues have 18 or 24 dimensions. On the other hand, relative conformations between ligands and their binding site residues must be placed with high accuracy to allow favorable interactions, especially polar interactions. Together, due to the high dimensionality of the possible space and the requirement of high accuracy, binding site matching often requires extensive sampling of protein backbones; this sampling is possible with de novo designed proteins. Our analysis on naturally existing and de novo fold families supports this rationale. The de novo scaffolds tested in this work sample conformational variability in local regions only. As a consequence, the de novo fold families cover a smaller fraction of overall conformational space than native fold families when considering the same number of structures (S4 Fig). As a consequence, when we used subsets of de novo fold families that have the same numbers of scaffolds as their corresponding native fold families, we found that the de novo fold sub-families support fewer binding sites than the native fold families. However, it is possible to generate many more de novo scaffolds than the number of existing native scaffolds with available structures. Moreover, the conformational variability that can be sampled in local regions is larger with de novo proteins (S4 Fig). As a result of this more extensive sampling, we show that the total numbers of matched binding sites to de novo fold families are higher than the numbers of matches to native fold families.

Previous studies have shown that computationally generated artificial (ART) compact homopolypeptide structures can match virtually every native ligand binding pocket [31, 32]. In contrast, the native and de novo fold families we studied here can only be matched to a limited fraction of native binding sites. A likely reason is that we only used structures with two topologies while the ART structures are generated using secondary structure preferences from thousands of random PDB structures with many different fold topologies. These two behaviors together support that the diversity of topologies is important for the repertoire of native ligand binding functions. Additionally, the de novo designed structures we used were subjected to filters for a set of physical properties such as core packing, hydrogen bonding and surface exposed hydrophobic patches [16]. These filters are designed to eliminate structures that are not likely to fold, whereas the ART structures are model polyleucine homopolypeptides. Requirements for folding places diverse additional constraints on the accessible conformational space of protein structures.

Using the new fast match protocol introduced here as well as the Rosetta matcher, we were able to match a library of high-quality binding sites to de novo protein fold families. To engineer new ligand binding proteins, the matching step is typically followed by sequence design [3, 9] to optimize the binding site protein environment. Ligand binding site design is a challenging problem because the designed sequence must simultaneously be compatible with the protein fold and precisely place binding site residues in their desired geometries for favorable
interactions with the ligand. Given the typically high stability of de novo designed protein families [16, 33], matches generated by the protocol described here could be good model systems for testing binding site design algorithms.

Another advantage of using de novo fold families for ligand binding site design is that the systematic sampling of diverse geometries could provide an ensemble of negative states. Using negative states in design has been shown to improve accuracy in protein stability prediction [34]. Thus, a de novo ensemble of negative states may increase success rates of ligand design where high accuracy in both sampling and scoring designs is required. Ensembles of different conformational states in de novo fold families also pave the way to engineer ligand binding-induced conformational changes. Small molecule-induced switches could be designed by building a ligand binding site in one of the structures in the de novo fold family and tuning the free energy gaps between the ligand binding state and the other states. We envision that de novo designed protein fold families will play an important role in designing functions such as ligand binding and protein switches.

Methods

Binding site library construction

Ligand binding sites were extracted from the PDB95 [18] database. The representative pdb structures for each cluster, which were listed in the pdb_95.cod file, were used for binding site extraction. The representative structures were filtered by resolution. Only crystal structures whose resolutions were better than 2 Å were kept. Ligand residues were identified by built-in functions in PyRosetta [35]. In this study, we focused on ligands that had at most 100 heavy atoms. Ligands that had average heavy atom B-factors greater than 60 Å² were filtered out. Ligands that did not have protein residues within 5 Å were also excluded from subsequent processing. We calculated the Rosetta 2-body energy scores [36, 37] between ligands and protein residues that have at least one heavy atom within 5 Å from any ligand heavy atom. Ligand binding site residues were defined as protein residues that had favorable van der Waals, electrostatic or hydrogen bond interactions with the ligand. A residue was included in a binding site if the sum of its Rosetta energy [37] terms fa_atr, fa_elec, hbond_bb_sc and hbond_sc was less than -1 Rosetta energy units (REU). We excluded protein residues from consideration that had total Rosetta scores greater than 50 to avoid poorly modeled residues, such as those who have severe clashes with the protein environment. We also excluded all residues with missing heavy atoms in the PDB file. We only kept ligand binding sites that have at least two protein residues. To prevent overcounting ligands in structures which had multiple chains of the same protein in their asymmetric units, only one binding site was extracted for the same ligand in a given structure.

Clustering ligand 3D structures

We clustered the ligands in the binding site libraries based on their 3-dimensional similarities quantified by their extended 3-dimensional fingerprints (E3FP) [19]. A 1024-bit fingerprint was calculated for each ligand by the e3fp python package. The fingerprints were clustered using the Butina clustering method [20] implemented in the RDKit open source cheminformatics package. During the clustering, the distance between two fingerprints was defined as one minus their Tanimoto Coefficient. The distance cutoff for the Butina clustering was 0.65.

Fast match protocol

We developed a new fast match protocol to rapidly match the library of binding sites to the sets of protein scaffolds. During the fast match, a ligand binding site is treated as a rigid body.
When the fast matcher matches a ligand binding site to a scaffold, it first iterates through all pairs of binding site protein residues and scaffold residues. For each pair of residues, the protocol superimposes the N, Ca and C atoms of the binding site residue to the corresponding atoms in the scaffold residue. The remainder of the binding site is transformed as a rigid body. Then the matcher finds the closest scaffold residues to each binding site protein residue. The distances between residues are defined as the Ca-Ca distances. If all distances between binding site protein residues and their closest scaffold residues are within 2 Å, the backbone N, Ca and C atoms of the binding site protein residues are superimposed to the N, Ca and C atoms of their closest scaffold residues. The superimposition minimizes the root mean squared deviation (RMSD) between the corresponding atoms. If the RMSD is within 1 Å, the cosine of angles between the vectors pointing from Ca to Cb of corresponding residues are calculated. If all the cosine values are greater than 0.7, clashes between the matched binding site and the scaffold backbone are checked. Two atoms are defined to clash when the distance between them is less than the sum of their Lennard-Jones radii times a scale factor of 0.6. The match is accepted if the ligand and protein side chains from the binding site do not clash with the scaffold backbone atoms that are not matched to binding site residues.

**Standard Rosetta matcher**

For each binding site successfully matched to a scaffold using fast match, we ran the standard Rosetta matcher [5]. We made mol2 files for ligands using Open Babel [38] and generated ligand parameter files with the molfile_to_params.py script distributed with Rosetta. The relative positions of a ligand and a binding site protein residue are defined by 6 heavy atoms. On the ligand side, the heavy atom closest to the protein residue and the two ligand heavy atoms closest to the first ligand heavy atom are defined as the anchor atoms. On the protein residue side, the heavy atom closest to the ligand and two protein atoms closest to the first protein heavy atom are defined as the anchor atoms. For each binding site, we generated a constraint file where the relative positions between the ligand anchor atoms and protein residue anchor atoms were constrained. We used stringent matching criteria similar to those used in previous work [9, 13]. The relative distances between ligands and binding site residues are sampled at ideal values; the relative angles and torsions are sampled at the ideal values and ±10° from the ideal values. The binding sites were matched using the standard Rosetta matcher with the following command:

```
match.linuxgccrelease -match:output_format PDB -match:match_grouper SameSequenceGroup -match:consolidate_matches -match:output_matches_per_group 1 -use_input_sc -in:ignore_unrecognized_res -ex1 -ex2 -enumerate_ligand_rotamers false -match::lig_name LIG_NAME -match:geometric_constraint_file CST_FILE -s SCAFFOLD_PDB -match::scaffold_active_site_residues POS_FILE
```

where LIG_NAME is the 3-letter name of the ligand, CST_FILE is the constraint file, SCAFFOLD_PDB is the pdb file of the scaffold structure and POS_FILE is the file that stores the matchable residues. In this study, all residues on a scaffold are matchable.

**Construction of scaffold libraries**

The de novo Rossmann and NTF2 fold families were reported in ref. [16]. The scaffolds in these fold families were generated by the LUCS method and filtered by a set of designability filters [16]. We randomly selected 1,000 scaffolds from each de novo fold family as the scaffold set for ligand binding site matching. The native fold families of Rossmann and NTF2 folds were obtained from the CATH database [23]. The native Rossmann fold scaffolds were extracted from the CATH 3.40.50.1980 superfamily and the native NTF2 family structures...
were from the CATH 3.10.450.50 superfamily. Because the automatic classification algorithm of the CATH database did not force all structures in a CATH superfamily to have the same topology, we manually excluded the CATH structures that have different topologies from the de novo designed scaffolds. As a result, the native Rossmann fold scaffold set had 20 structures and the native NTF2 fold scaffold set had 103 structures. The C-terminal helices in de novo NTF2 scaffolds occluded the ligand binding pocket. In contrast, only 35 out of 103 native NTF2 scaffolds had C-terminal helices. Among these native C-terminal helices, 31 helices pointed away from pocket entrances, and thus, did not affect the accessibility of ligand binding sites, leaving only 4 scaffolds with pocket occluding C-terminal helices. We therefore trimmed the C-terminal helices in de novo NTF2 proteins to expose the ligand binding pocket.

Calculation of pairwise RMSD distributions

Pairwise RMSDs between scaffold structures were calculated using the mTM-align software [39] with its default settings. For the native Rossmann fold family, pairwise RMSDs between all structures were calculated. For the de novo Rossmann, native NTF2 and de novo NTF2 fold families, 100 random structures were selected for the respective pairwise RMSD calculation. The pairwise RMSD distribution between native and de novo Rossmann fold families was calculated using all 20 native Rossmann fold structures and 20 random de novo Rossmann fold structures. The pairwise RMSD distribution between native and de novo NTF2 fold families was calculated using 50 random native NTF2 fold structures and 50 random de novo NTF2 fold structures.

Construction of a library of 3-residue binding sites

The 3-residue binding site library was constructed from the library of all binding sites. We eliminated binding sites with fewer than 3 residues. The binding sites with 3 protein residues were kept unchanged. For binding sites with more than 3 protein residues, we scored the total Rosetta two-body energy [36] between the ligand and each protein residue. We kept the 3 protein residues with lowest total two-body energies and removed the remainder of the binding site residues.

Assignment of layers to scaffold residues

The Rosetta Layer selector [25] with the default settings was applied to assign layers to each scaffold residue. The layer of a residue was determined by a weighted count of the number of neighbor amino acid residues in a cone extending along its Ca-Cb vector. The weight of a neighbor residue is defined as a distance factor multiplied by an angle factor. The distance factor is defined as

$$1/(1 + \exp((d - 9)))$$

where d is the distance of the neighbor from the residue CA in angstrom. The angle factor equals to

$$\left((\cos(\theta) + 0.5)/(1 + 0.5)\right)^2$$

where $\theta$ is the angle between the CA-CB vector and the CA-neighbor vector. More details about the Layer selector are explained in the Rosetta online documentation (https://new.rosettacommons.org/docs/latest/scripting_documentation/RosettaScripts/ResidueSelectors/ResidueSelectors#residueselectors_conformation-dependent-residue-selectors_layerselector). A residue is assigned to the surface layer if the weighted count is less than 2; a residue is
assigned to the core layer if the weighted count is greater than 5.2; all other residues are assigned to the boundary layer.

**Assignment of secondary structures**

Secondary structures were assigned to the scaffolds by the DSSP algorithm [40] implemented in the PyRosetta package [35]. The secondary structure of each residue in a matched binding site was defined as the secondary structure of its matched scaffold residue position.

**Calculation of the numbers of matches for subsets of fold families**

During the process of matching a binding site to a fold family, we recorded the number of scaffolds in the fold family that we tested to find the first successful fast match and called this number the first-fast-match-encounter-number. The number of fast matches for a subset of a fold family with N scaffolds was defined as the number of binding sites with first-fast-match-encounter-numbers smaller than or equal to N. The number of Rosetta matches for subsets of fold families were calculated in the same way.

**Supporting information**

S1 Fig. Amino acid (AA) frequencies and enrichment ratios in binding sites that bind to ligands with at least 2 heavy atoms.

(TIFF)

S2 Fig. Secondary structure distributions in fold families and matched 3-residue binding sites. A. Distributions of secondary structures of scaffolds in different fold families. B. Distributions of secondary structures of matched 3-residue binding sites.

(TIFF)

S3 Fig. Venn diagrams of numbers of Rosetta-matched 3-residue binding site clusters between pairs of scaffold sets. The 3-residue binding sites were clustered by 3D similarities of their ligands. A cluster is defined as matched if it has at least one matched binding site. For each Venn diagram, the number in the overlapping region is the observed number of clusters that can be matched to both scaffold sets, with the expected number in parentheses. The number in the non-overlapping region within a circle denotes the clusters that can only be matched to this scaffold set. The number outside the circles denotes the clusters that cannot be matched to either of the two scaffold sets.

(TIFF)

S4 Fig. Structural variation in protein fold families. A. Distributions of pairwise RMSDs between structures in native fold families (native), de novo fold families (de novo), cross comparison between native and de novo fold families (cross), or pairwise RMSDs between the de novo designed LHL units when aligned the remainder of the protein (de novo LHL, calculated using the method in ref(16)). B. Examples of scaffold structures from each fold family.

(TIFF)

S1 Table. Ligand type frequencies in the binding site library.

(TSV)

S1 File. Summary tables of matching results to all fold families.

(GZ)
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Author Contributions
Conceptualization: Xingjie Pan, Tanja Kortemme.
Data curation: Xingjie Pan.
Formal analysis: Xingjie Pan.
Funding acquisition: Tanja Kortemme.
Investigation: Xingjie Pan.
Methodology: Xingjie Pan.
Project administration: Tanja Kortemme.
Resources: Xingjie Pan.
Software: Xingjie Pan.
Supervision: Tanja Kortemme.
Validation: Xingjie Pan.
Visualization: Xingjie Pan.
Writing – original draft: Xingjie Pan.
Writing – review & editing: Xingjie Pan, Tanja Kortemme.

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