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PROTEIN DESIGN

A defined structural unit enables de novo design of small-molecule-binding proteins

Nicholas F. Polizzi* and William F. DeGrado*

The de novo design of proteins that bind highly functionalized small molecules represents a great challenge. To enable computational design of binders, we developed a unit of protein structure—a van der Mer (vdM)—that maps the backbone of each amino acid to statistically preferred positions of interacting chemical groups. Using vdMs, we designed six de novo proteins to bind the drug apixaban; two bound with low and submicromolar affinity. X-ray crystallography and mutagenesis confirmed a structure with a precisely designed cavity that forms favorable interactions in the drug–protein complex. vdMs may enable design of functional proteins for applications in sensing, medicine, and catalysis.

he Anfinsen hypothesis states that a protein's sequence encodes its tertiary structure and underlying function (1). Conversely, a protein's tertiary structure encodes the possible sequences compatible with a particular function. De novo protein design has succeeded in the creation of proteins that fold to various targeted tertiary structures (structure to sequence) (2, 3). Nevertheless, it has been extremely challenging to design proteins that not only fold but also bind to complex small molecules (function and structure to sequence) (2-4). Use of algorithms optimized for packing apolar protein cores leads to difficulty when designing polar cavities required for binding hydrophilic molecules (5). Consequently, design of smallmolecule-binding proteins has generally required recursive experimental screening and large libraries to engender function, mostly starting with natural proteins rather than de novo structures (Fig. 1A) (3, 4, 6-9). Here, we accomplish the reverse of the Anfinsen hypothesis by simultaneously designing structure and binding function from scratch, targeting a small-molecule drug with significant polarity and structural complexity. To do this, we developed a unit of local protein structure that directly links a tertiary structure to key interactions that engender tight and specific binding. These findings illuminate the principles underlying the emergence and evolution of complex function in proteins and provide a methodology for designing useful proteins.

Targeted function and fold

We targeted the factor Xa inhibitor apixaban, an organic compound with five rotatable bonds and eight heteroatoms. Our first objective was to compute a tertiary structure capable of cooperatively binding the polar groups of

Department of Pharmaceutical Chemistry, Cardiovascular Research Institute, University of California, San Francisco, San Francisco, CA 94158, USA. *Corresponding author. Email: nicholas.polizzi@ucsf.edu (N.F.P.); william.degrado@ucsf.edu (W.F.D.) apixaban. Instead of repurposing natural binding proteins or folds that have been shown to bind a similar ligand, in this work, we use de novo four-helix bundles because they are mathematically parameterized (10, 11), designable (12), and share no similarity to the fold of factor Xa. Four-helix bundles generally do not bind small molecules and instead bind metal ions or metalloporphyrins by strong coordinate bonds (10, 13-16). However, fourhelix bundles are tubular and can be designed to have high thermodynamic stability (11, 13) to compensate for the energetically demanding process of building binding cavities replete with buried polar functionality (17). Thus, the design of a de novo helical bundle that binds the drug apixaban critically tests the design method.

The van der Mer structural unit

The design of proteins relies on optimal packing of interior side chains in discrete conformations called rotamers (2, 3, 18-22). However, the design of ligand-binding proteins additionally requires side chains that interact favorably with the target small molecule. Previous design strategies approached this problem by computationally appending the target ligand to rotamers with idealized interaction geometries that-although composed of billions of conformations-sampled only a small fraction of the possible conformational space (6, 8, 23). These strategies rarely deliver submillimolar binders from the initial computational design, so subsequent steps rely on experimental random mutagenesis and screening of libraries.

We wondered how much of the vast, possible conformational space of protein-chemical group interactions is actually sampled in observed protein structures and if sampling interactions directly from this distribution might aid the design of high-affinity binders. Whereas previous analyses have focused on local side chain contacts with chemical groups (24), we sought a structural unit that directly maps backbone coordinates to chemical group loca-

tions, the link between the protein fold and binding function. We developed a unit of protein structure analogous to rotamers-a van der Mer (vdM)-that defines the placement of key chemical groups in the ligand relative to the backbone atoms of the contacting residue (Fig. 1B). vdMs are culled from a nonredundant set of protein structures by (i) identifying all residues of a certain type that interact with a particular chemical group, (ii) performing an all-by-all pairwise superposition of only the backbone and chemical group coordinates (side chains are not considered in the superposition, allowing some variation in their conformation), and (iii) geometric clustering with a tight root mean square deviation (RMSD) cutoff (0.5 Å). The resulting vdMs show backbone φ and ψ dependence (Fig. 1C) and capture compensatory effects of backbone and chemical group placement. Furthermore, single clusters may contain multiple rotamers (Figs. 1D and 6A and fig. S1), given that sidechain coordinates are not explicitly considered in clustering.

The use of vdMs contrasts with procedures that place ligands at idealized locations relative to the terminal atoms of a side chain (6, 8, 23, 25), which results in vast numbers of ligand-rotamer combinations that might never occur in proteins. Instead, vdMs sample locations of chemical groups relative to the backbone that have been experimentally vetted to achieve favorable interactions. They also implicitly consider interactions with ordered or bulk water, which might influence their interaction geometries. Moreover, unlike ligand-appended and inverse rotamers used in earlier approaches (6, 8, 23, 25), vdMs may be derived from contacts with either main chain, side chain, or both in a multivalent interaction. Finally, the prevalence of a given vdM in the Protein Data Bank (PDB) can be used in scoring functions, similarly to scoring rotamers, which may assist automated selection of binding-site residues for design.

To maximize the number of observed protein-chemical group contacts, we created vdMs using the chemical groups of amino acids that constitute the protein (e.g., $CONH_2$ of Gln and Asn and N-H and C=O of backbone amides). To avoid bias from local structure, we counted only the interactions that were distant in the linear polypeptide chain, as described in the supplementary materials. The set of chemical groups can also be expanded to include those from small-molecule drugs, metal ions, and cofactors, although these are not as pervasive in crystal structures.

We ranked vdMs by their prevalence in the PDB using a log-odds score, C (Fig. 1, D and E; fig. S2; and supplementary text). Although there are hundreds of vdMs associated with a given residue–chemical group combination

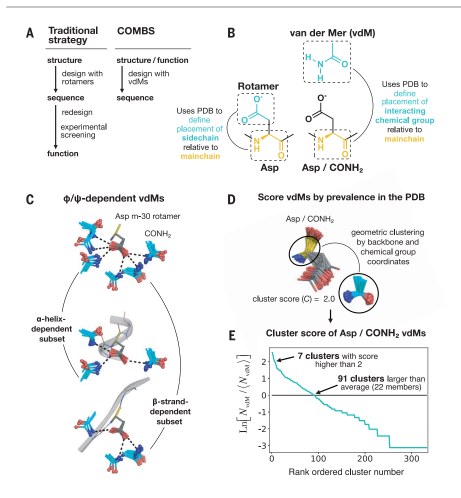


Fig. 1. A vdM is a structural unit relating chemical group position to the protein backbone. (A) Workflow of a traditional protein design strategy versus that of COMBS. **(B)** Definition of a vdM. A chemical group is interacting if it is in van der Waals contact with the protein side chain or main chain. Like rotamers, vdMs are derived from a large set of high-quality protein crystal structures. A vdM of aspartic acid (Asp) and carboxamide (CONH₂, cyan) is shown. **(C)** vdMs are φ , ψ , and rotamer dependent; this is illustrated by the top vdMs of the m-30 rotamer of Asp, clustered by location of CONH₂ after exact superposition of main chain N, C α , and C atoms. **(D** and **E)** We ranked vdMs by prevalence in the PDB, quantified by a cluster score *C* [the natural logarithm of the ratio of the number of members in a cluster (N_{vdM})]. The seventh-largest cluster of Asp/CONH₂ vdMs is shown as an example in (D).

(figs. S3 and S4), only a small fraction of vdMs are highly enriched in protein structures (C > 0). For example, only 91 Asp CONH₂ vdMs have C values >0; these top vdMs map the locations of CONH₂, relative to the backbone of an Asp residue, that are statistically preferred by proteins in the PDB (Fig. 1E and fig. S2D). This is on the order of the number of rotamers used for an amino acid during a typical protein design packing calculation (26). Thus, when combined with an efficient search algorithm, sampling proteinchemical group interactions with vdMs to design ligand-binding sites might be as expedient as sampling rotamers to pack a protein core. Furthermore, functionally relevant lower-probability rotamers may be included if contained in a high-scoring vdM.

Proteins use the same set of 20 amino acids to fold as well as to recognize a vast array of highly functionalized ligands. We therefore hypothesized that the interaction modes used by amino acids to stabilize their tertiary structures would also be used to achieve tight binding of ligands, even those containing structurally distinct heterocyclic chemical groups. To test this hypothesis, we examined the streptavidin-biotin complex (Fig. 2). Using the natural sequence of streptavidin, we examined the positions of vdMs of N-H, C=O, and COO⁻, where these groups were derived from protein main chains and side chains. In each case, we observed that the side-chain interactions with biotin's polar groups involved highly favorable vdMs, with enrichment scores of ~8-fold or greater (C > 2). The streptavidin sequence-fold pairing cooperatively positions highly favorable vdMs to cover each polar chemical group of biotin simultaneously.

Our analysis of the streptavidin-biotin complex suggests that binding sites can be designed by considering folds that position vdMs to collectively bind the distinct chemical groups found in a target small-molecule ligand. Moreover, the vdMs of the binding site should be maximally prevalent in the PDB. We developed a search algorithm, called Convergent Motifs for Binding Sites (COMBS), to discover favorable poses of a ligand that satisfy these criteria.

De novo design strategy

Our design strategy consists of several hierarchic steps, which prioritize the most essential and difficult features to avoid sampling regions in sequence and structure space with little chance of success (fig. S5). First, we define the chemical groups within the small molecule that will be targeted. We initially focus on polar chemical groups, which are the most challenging to dehydrate but must be satisfied with H-bonds to achieve high affinity and specificity (27). Second, we choose a designable protein fold and create an ensemble of backbones with geometries that are consistent with the known plasticity of the fold. Next, for each backbone we use COMBS to identify members of the backbone ensemble that can position vdMs to collectively engage each of the targeted chemical groups of the small molecule. In this way, the binding of the desired ligand dictates the precise backbone geometry. Having discovered candidate backbones and binding sites, the design is completed by engineering a tightly packed folding core that supports the vdMderived keystone interactions in the binding site (13). In this step, we constrain the keystone interactions and use flexible backbone design (13, 26) to pack additional residues within the binding site while simultaneously packing the protein core.

We focused on apixaban's carboxamide (both the C=O and -NH₂), as well as two additional carbonyls (Fig. 3A). (Other groups that were internally H-bonded or easily dehydrated were not initially targeted.) We created a set of vdMs of carboxamide (CONH2 from Asn and Gln side chains) and carbonyl [C=O from the protein backbone (supplementary text)] and used these vdMs to discover preferred CONH₂ and C=O binding locations within a set of 32 mathematically generated de novo polyglycine backbones (10, 28) (Fig. 3, B and C; fig. S2; and table S1). For each of the mathematically generated backbones, we placed apixaban in the protein interior by using a separate set of vdMs with apixaban superimposed onto the chemical group of the vdM. For example, the CONH₂ of apixaban can be superimposed

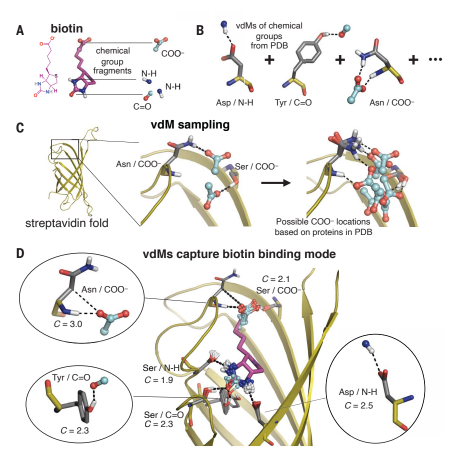


Fig. 2. Prevalent vdMs describe the binding site of biotin in streptavidin. (A and B) We constructed vdMs of the polar chemical groups of biotin by searching the PDB for protein interactions with the (i) backbone amide nitrogen (N-H), (ii) backbone carbonyl or carbonyl from Asn or Gln side chains (C=O), and (iii) carboxylate of Asp or Glu side chains (COO⁻). (**C**) Using the native sequence of streptavidin, vdMs were sampled on the streptavidin backbone to generate possible locations for productive interactions with the chemical groups. Here, Asn and Ser vdMs of COO⁻ are sampled at two positions of the backbone. (D) vdMs with chemical groups (cyan) that are nearest neighbors (0.6 Å RMSD) to those of biotin in its binding site are overlaid on top of biotin (purple).

on the CONH₂ of a vdM, uniquely defining the position of apixaban in the binding site. Apixaban's conformation in this step was fixed in a low-energy conformer found in its cocrystal structure with factor Xa (PDB code 2p16) (Fig. 3A and fig. S6; extension to multiple conformers is discussed in the supplementary text and illustrated in fig. S7). vdMs that cover the remaining C=O groups of the placed ligand, as well as additional vdMs to the carboxamide, were then queried in the nearby space (Fig. 3D). We chose binding poses by maximizing the PDB prevalence of sterically compatible vdMs ($\sum C$) (Fig. 3E).

Side chains from vdMs in six selected binding poses were fixed, and their H-bonding interactions with apixaban were constrained in all subsequent steps of sequence design performed within the Rosetta modeling suite. After insertion of interhelical loops, we used a flexible backbone design protocol (13) (Fig. 3F) to compute the hydrophobic core while simultaneously completing the packing of the binding site. For some designs, new polar interactions were recruited during this step, as were Gly residues, which are known to interact favorably with aromatic groups (27). The use of small residues to make hydrophobic contacts minimizes the number of large, apolar side chains that might lead to nonspecific binding or hydrophobic collapse in the absence of ligand.

Description of designs and biophysical characterization

We designed six proteins of varying length, topology, ligand position, ligand burial, and keystone interactions (fig. S8). By contrast to factor Xa, which engages polar groups of apixaban through main-chain amides in loops (fig. S6), the designs interact with apixaban using predominantly side chains in helices. The six designs were well-expressed in bacteria, and each was helical based on far UV circular dichroism spectroscopy (fig. S9). Proton nuclear magnetic resonance (NMR) showed that two designs, ABLE (apixaban-binding helical bundle) and LABLE (longer ABLE), bound apixaban (fig. S10). These two designs had the same orientation of apixaban within the bundle and shared the same vdM-derived keystone interactions (Fig. 3E and fig. S8). For example, they shared a buried, high-scoring His/C=O vdM (8-fold enrichment, C = 2.1) (Fig. 3E). However, ABLE and LABLE differed in length (125 vs. 165 residues), topology, and loop geometry and shared only 22% sequence homology.

Binding of apixaban to ABLE restricts the drug's conformation, resulting in a red shift of

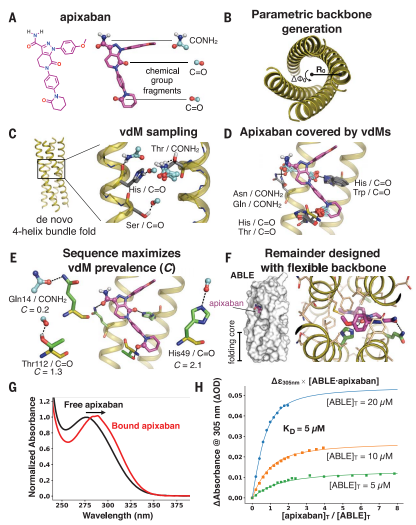
its electronic absorbance spectrum (Fig. 3G). Spectral titrations and fluorescence polarization competition experiments showed that ABLE and LABLE bind apixaban with a dissociation constant (K_D) of 5 (± 1 [SEM]) μ M and 0.6 (\pm 0.1) μ M, respectively (Figs. 3H and 4D; figs. S11 and S12). Although LABLE showed a dispersed two-dimensional ¹H-¹⁵N heteronuclear single-quantum coherence spectrum by NMR (fig. S13), indicative of a wellstructured protein, it failed to crystallize in a sparse matrix screen, and so we focused our attention on characterization of ABLE. ABLE is monomeric in solution (fig. S14) and highly stable to heat denaturation (melting temperature of >95°C), despite the inclusion of three Glv and a polar His within its core (fig. S15).

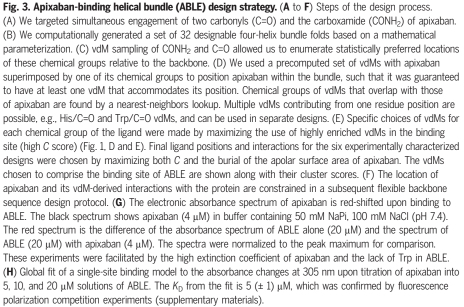
Structures of apixaban-bound and drug-free ABLE

ABLE readily crystalized with apixaban and diffracted to 1.3-Å resolution. Two very closely related monomers were observed in the asymmetric unit (fig. S16); apixaban is bound to both monomers, as expected for a specific, high-affinity complex. The structure of the drug-bound protein is in excellent agreement with the design (C α RMSD of 0.7 Å) (Fig. 4). The rotamers of the core residues of ABLE, including the binding-site residues, overwhelmingly agree with the design model.

Superimposing by all heavy atoms of core amino acids, including apixaban, gives an RMSD of 0.98 Å. ABLE buries almost all available apolar surface area (504 Å²) of apixaban,

and it also forms most polar interactions included in the design (Fig. 4, B and C). Apixaban's conformation is close to that used in the design (0.6 Å heavy atom RMSD), with small devia-





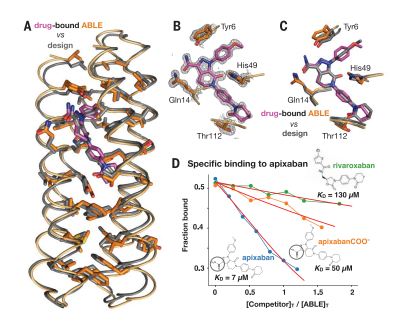
tions that bring it closer to a quantum mechanically optimized geometry (fig. S17). The rigid body translation between apixaban's center of mass in the designed versus that in observed structures is only 0.2 Å, with a rigid body rotation of 6°. The bespoke binding site is specific for apixaban, as shown by fluorescence polarization competition experiments (Fig. 4D), which indicated that ABLE binds apixaban 20-fold more tightly than a similar factor Xa inhibitor, rivaroxaban.

To assess the extent of preorganization of the protein, we also solved the drug-free structure to 1.3-Å resolution (Fig. 5). The structure shows an open, preorganized binding pocket, with an overall C α RMSD of 0.65 Å to the apixaban-ABLE complex. The unoccupied binding site is solvated by nine ordered water molecules plus an acetate from the buffer (Fig. 5D). Binding of apixaban displaces ordered solvent from this site, suggesting a release of local frustration upon binding. The pocket has a 480-Å² solvent-exposed surface area, which expands by 40% to accommodate the drug (680 Å^2). The drug-free protein has nearly identical rotamers to that of the drug-bound protein throughout the core and binding site (Fig. 5, G to I). Unliganded ABLE shows two alternate rotamers for several of the residues that form H-bonds to apixaban (e.g., Tyr⁴⁶ and His⁴⁹); binding of apixaban selects one each of these alternate rotamers. Thus, like many natural proteins (29), ABLE has a limited degree of flexibility, which is reduced upon ligand binding, and the binding event appears to trade configurational entropy for enthalpically favorable interactions.

Insights from the structure and function of ABLE

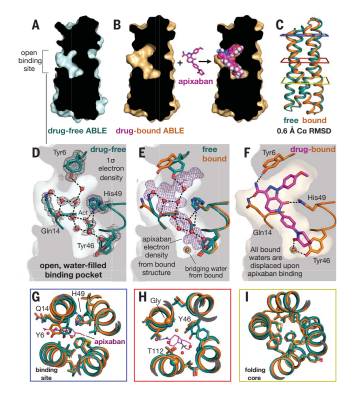
Two of the three keystone interactions identified by COMBS contribute appreciably to binding affinity. Substitution of His⁴⁹ or Gln¹⁴ with alanine individually decreases affinity by approximately 1 kcal/mol (~3-fold) (Fig. 6D and fig. S18). Gln¹⁴ was observed in its intended rotamer, whereas His49 occupied an alternate rotamer that nevertheless maintained the intended position of apixaban's carbonyl relative to the main chain (Fig. 6A). Indeed, the cluster describing this His/C=O vdM contains multiple His rotamers, each capable of achieving identical placements of C=O relative to the main chain. Thus, we observed vdM convergence, even amidst rotamer divergence.

We also examined the structural consequences of substituting His⁴⁹ with Ala by solving the crystal structure of the unliganded His⁴⁹ \rightarrow Ala (H49A) mutant protein (fig. S19). Although the structures of drug-free ABLE and drug-free H49A are similar (C α RMSD = 1.2 Å), the residues that surround His⁴⁹ show rotameric differences in the absence of this side chain; released from the restraints of tight Fig. 4. The structure of apixaban-bound ABLE agrees with the design. (A) Superposition of backbone $C\alpha$ atoms of structure (protein in orange, apixaban in purple) and design (gray; 0.7 Å RMSD), showing side chains of amino acids in the protein core. (B) ABLE's binding site from the structure (1.3-Å resolution), showing vdM-derived interactions with apixaban (purple). The 2mFo-DFc composite omit map is contoured at 1.5 σ . The map was generated from a model that omitted coordinates of apixaban. The protein backbone of these residues is shown in cartoon format. (C) Overlay of designed interactions (gray), after the designed model was superimposed onto the Ca atoms of the structure (protein in orange, apixaban in purple). (D) Fluorescence anisotropy competition experiments (485-nm excitation, 528-nm emission) showed that ABLE binds apixaban specifically. The bound fluorophore apixabanpolyethylene glycol-fluorescein isothiocyanate (apixaban-PEG-FITC) (supplementary text and fig. S9) is dislodged by addition of competing ligand. Anisotropy was converted to the fraction bound by use of a one-site binding model (supplementary text). The ABLE concentration was 20 µM, and the apixaban-PEG-FITC concentration was 25 nM in buffer containing 50 mM NaPi, 100 mM NaCl (pH 7.4). Apixaban COO⁻ is identical to apixaban except that it contains a



carboxylate instead of a carboxamide (circled). Rivaroxaban is another inhibitor that also binds tightly to factor Xa by using the same binding mode as apixaban but shows only very weak binding to ABLE. Fits to a competitive binding model are shown in red. K_D values: rivaroxaban, 130 (± 10) μ M; apixaban COO⁻, 50 (± 5) μ M; apixaban, 7 (± 2) μ M.

Fig. 5. Drug-free ABLE has a preorganized structure with an open binding site competent for binding. (A) A slice through a surface representation of the 1.3-Å resolution structure of unliganded ABLE shows an open binding cavity. (B) Same slice, shown for the structure of apixaban-bound ABLE. (C) The C α atom backbone superposition of unliganded and liganded ABLE. Colored squares surrounding the structure correspond to panels in (G), (H), and (I), looking down from the top. (D) The binding site of drug-free ABLE shows nine buried, crystallographic waters (red spheres, occupancy > 0.9) involved in an extensive H-bonded network with binding-site residues Tyr⁶, Gln¹⁴, Tyr⁴⁶, and His⁴⁹. The 2mFo-DFc electron density map of drug-free ABLE is contoured at 1 σ . An acetate (Act) group from the crystallization condition H-bonds with His⁴⁹. His⁴⁹ and Tyr⁴⁶ are observed with alternate rotamers. (E) Same view as in (D) but with the addition of the corresponding residues from the apixaban-bound structure, after an all-C α -atom backbone superposition. The 1- σ 2mFo-DFc electron density (purple) of apixaban from the drug-bound structure shows where the crystallographic waters bind in the ligand-free structure relative to the bound structure. A water (shown as an orange sphere) mediates the H-bond between Tyr⁴⁶ and apixaban. This water is not observed in the unliganded structure. (F) Binding of apixaban in the drug-bound structure displaces all of the nine buried waters in the drug-free structure. Stick renderings, as well as the surface background, show the binding site of the ABLE-apixaban complex. (G and H) Binding-site overlay of liganded (orange, apixaban purple) and unliganded (cyan) ABLE shows preorganized rotamers. (I) The remote folding core contains identical rotamers in drug-free and drug-bound ABLE, predisposing the drug-free protein for binding.



Substitution of the third keystone residue, Thr¹¹², with Ala resulted in little change in affinity (Fig. 6D). In the complex, its side chain did not form the intended H-bond to apixaban but instead formed an intrahelical H-bond to a backbone carbonyl (Fig. 6C). The intended Thr/C=O vdM is favored in the backbone-independent vdM library used in the design of ABLE, but it is disfavored in a backbone-dependent vdM library. The lack of engagement with apixaban's carbonyl resulted in some disorder of the terminal oxopiperidine,

packing, they instead adopt their preferred

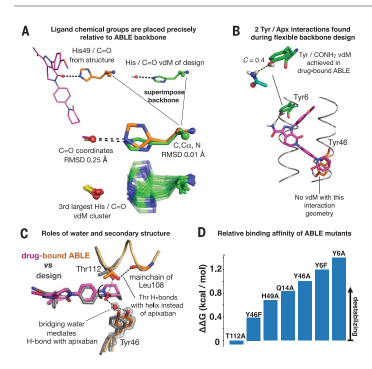
rotamers. The structure illustrates that glo-

bal packing of core residues supports the

positioning of a key functional group, even

when this requires local frustration at indi-

vidual sites.



which has higher b-factors and two alternate conformations (related by a 180° ring flip) in the structure (fig. S16). Thus, backbonedependent vdM libraries should be used in future applications.

Flexible backbone sequence design of ABLE recruited two Tyr residues that interact with apixaban (Fig. 6B). One of these interactions was represented in the vdM database (Tyr⁶/ CONH_2 , C = 0.4), but the other $(\text{Tyr}^{46}/\text{C}=\text{O})$ was not. The structure of drug-bound ABLE confirmed the H-bond of $Tyr^6/CONH_2$ (Fig. 4C), but an unanticipated water enters the binding site to mediate an H-bond between apixaban and Tyr⁴⁶ (Fig. 6C). Furthermore, substitution of Tyr⁶ with Phe or Ala was more destabilizing than the same substitutions for Tyr⁴⁶, tracking with prevalence in the PDB. Thus, vdMs can be used to filter and rank interactions obtained using a variety of computational methods (30).

Finally, we wondered if ab initio folding predictions (26) might distinguish between successful versus unsuccessful designs. Of the six designs, only two-ABLE and LABLE-were predicted by folding simulations to maintain uncollapsed binding sites (fig. S20). Moreover, the lowest-energy models predicted from ab initio folding simulations of ABLE's sequence largely agreed with the crystallographic structure (fig. S20A). Thus, ab initio folding may be useful as a screen to ensure that designs maintain an open, preorganized site. These results emphasize the degree to which the folding and binding problems are intimately coupled.

Conclusion

Previously, the design of de novo proteins that bind in a shape-selective manner to rigid, flat, hydrophobic dyes or lipidic metabolites had been possible, but binding flexible molecules replete with polar atoms has been more challenging (4, 8, 31-33). Natural proteins bind highly functionalized ligands by first accruing the ability to weakly bind fragments within the context of a particular fold (34-36). To mimic this process, we developed the vdM structural unit to directly link the protein fold to statistically preferred binding modes of chemical groups. We sampled vdMs on the backbone of a designable four-helix bundle to create constellations of chemical groups that, when matched with the shape of apixaban, defined the binding site. This contrasts with previous approaches that search for positional matching of whole ligands, sampled using idealized interaction geometries. Such approaches are highly sensitive to small changes in the interaction geometries and thus require an enormous amount of sampling to discover possible binding solutions, many of which may contain interactions not observed in the PDB.

vdMs sample from the experimentally vetted distribution of observed protein structures. vdMs are surprisingly sparse and discrete (Fig. 1E and figs. S3 and S4), and they enable facile sampling of sequence space to discover convergent combinations of keystone interactions (supplementary text and fig. S2). We consider only the backbone and the orientation of the pendant chemical group, which obviates the need to enumerate a large ensemble of ligandappended rotamers for each amino acid type at each position of the sequence. We focused here on simple, fully de novo scaffolds rather than redesigning the specificity of natural ligand-binding proteins, because we wished to address the challenge of designing function entirely from scratch. Indeed, ABLE shares

Fig. 6. Design inferences from the structure and function of

ABLE. (A) Exact sidechain positioning is not necessary for precise placement of ligand chemical groups relative to the mainchain. The placement of the C=O chemical group of apixaban relative to the backbone of residue 49 is exact (0.25 Å RMSD). The His⁴⁹/C=O vdM from the design (green) (Fig. 3E) was superimposed onto His⁴⁹ (orange) of the drug-bound ABLE structure through use of backbone atoms (N, $C\alpha$, C atoms, spheres). This backbone superposition places the C=O group of the original vdM precisely (0.25 Å RMSD) onto that of apixaban (purple) in the structure. The cluster describing the His C=O vdM, shown beneath, contains multiple rotamers of His that achieve the same placement of C=O relative to the position of the backbone. The rotamers of His⁴⁹ in the structure and His from the original vdM are both observed in the cluster. (B) Flexible backbone sequence design (Fig. 3F) resulted in recruitment of two additional polar interactions with apixaban from Tyr⁶ and Tyr⁴⁶. A Tyr⁶/CONH₂ vdM is prevalent in the PDB, whereas the Tvr⁴⁶/C=O interaction is not found in the database. (C) A water mediates an H-bond between Tyr⁴⁶ and the C=O group of apixaban. Thr¹²² H-bonds the C=O of the helix backbone at residue 108. (D) Relative binding affinities of ABLE mutants with apixaban-PEG-FITC fluorophore by fluorescence anisotropy experiments (supplementary text and fig. S18).

> no sequence homology to any known proteins (BLAST E value of <0.42 against the nonredundant protein sequence database nr). We used only prevalence to rank vdMs and choose binding sites, but we suspect the true power of vdMs may lie in higher-order correlations of the interactions.

> COMBS and vdMs can now be used for a variety of protein engineering applications and in full partnership with experimental optimization strategies for exploring sequence space. We anticipate that vdMs can also be used to predict chemical group hot spots of proteins with fixed sequence. vdMs may also enable design of protein-protein interfaces in a selfconsistent manner. Finally, because vdMs sample from the distribution of evolved interaction geometries observed in protein structures, it is tempting to view the chemical group constellations constructed by vdMs as a structural hypothesis of the evolutionary path to acquire binding within the context of a given fold.

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W.F.D. are inventors on a provisional patent application submitted by the University of California, San Francisco, for the design, composition, and function of the proteins in this study. **Data and materials availability:** Computational code and design scripts are available in the supplementary materials and at Zenodo (37). Coordinates and data files of ABLE structures have been deposited to the PDB with accession codes 6W6X (drug-free ABLE), 6W70 (apixaban-bound ABLE), 6X8N (H49A ABLE mutant). Materials are available from the authors on request. The plasmid of ABLE is available from Addgene (no. 158627).

SUPPLEMENTARY MATERIALS

science.sciencemag.org/content/369/6508/1227/suppl/DC1 Materials and Methods Supplementary Text Figs. S1 to S20 Tables S1 to S4 References (38-56) MDAR Reproducibility Checklist Data S1 View/request a protocol for this paper from *Bio-protocol*.

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A defined structural unit enables de novo design of small-molecule-binding proteins

Nicholas F. Polizzi and William F. DeGrado

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A new tool in the protein design toolbox

Protein design compute protein folds from first principles. However, designing new proteins that are functional remains challenging, in part because designing binding interactions requires simultaneous optimization of protein sequence and protein-ligand conformation. Polizzi and DeGrado designed proteins from scratch that bind a small-molecule drug (see the Perspective by Peacock). They introduced a new structural element called a van der Mer (vdM), which tracks the orientation of a chemical group relative to the backbone of a contacting residue. Assuming proteins bind ligands using interactions similar to intraprotein packing, they determined statistically preferred vdMs from a large set of structures in the Protein Data Bank. By including weighted vdMs in their computations, they designed model. *Science*, this issue p. 1227; see also p. 1166

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