The Recognition of Unrelated Ligands by Identical Proteins

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Supporting Information

ABSTRACT: Unrelated ligands, often found in drug discovery campaigns, can bind to the same receptor, even with the same protein residues. To investigate how this might occur, and whether it might be typically possible to find unrelated ligands for the same drug target, we sought examples of topologically unrelated ligands that bound to the same protein in the same site. Seventy-six pairs of ligands, each bound to the same protein (152 complexes total), were considered, classified into three groups. In the first (31 pairs of complexes), unrelated ligands interacted largely with the same pocket residues through different functional groups. In the second group (39 pairs), the unrelated ligand in each pair engaged different residues, though still within the same pocket. The smallest group (6 pairs) contained ligands with different scaffolds but with shared functional groups interacting with the same residues. We found that there are multiple chemically unrelated but physically similar functional groups that can complement any given local protein pocket; when these functional group substitutions are combined within a single molecule, they lead to topologically unrelated ligands that can each well-complement a site. It may be that many active and orthosteric sites can recognize topologically unrelated ligands.

INTRODUCTION

Chemical biology and drug discovery often seek new chemical scaffolds for receptor targets, unrelated to established ligand series. Such new chemotypes can illuminate previously unknown signaling, engage new regulatory networks, overcome key bioavailability limits for a target, or confer new selectivity among target family members. Whereas one might expect a genuinely novel ligand to occupy a new binding site, such as an allosteric site, it is often the main enzyme active or receptor orthosteric sites that are the most preorganized for binding and to which the novel ligands bind. By classical structure–activity or “similar properties principle” medicinal chemistry, this can be surprising.

Biophysically, of course, there is little to preclude the favorable binding of unrelated chemotypes in the same protein site. For any given protein subsite, several functional groups can be found to complement it well, and these may be combined in multiple unrelated ligand topologies to satisfy overall receptor complementarity. This was shown in a study of protein−ligand pockets by Gao and Skolnick, who found that many pockets bound unrelated ligands, characterizing four examples in detail. Also, many of the structurally distinct ligands from library screening bind to the main active or orthosteric sites of the target proteins. Our interest here is how unrelated ligands are recognized in the same site, and whether one might expect it to be common or rare. We will make no distinction here between endogenous and synthetic ligands, nor will we consider the evolution of protein−ligand recognition or its promiscuity, although this is a rich topic to which the biophysics of recognition relates.

Here, we investigate 76 pairs of unrelated ligands binding to the same target site for 71 proteins—over 150 complexes overall. While we do not pretend that this is comprehensive, this set covers most major drug target families, including G-protein coupled receptors (GPCRs), kinases, enzymes, nuclear hormone receptors (NHRs), and ion channels. We will consider whether these complexes, in the context of the biophysical considerations sketched above, support the likelihood that many proteins may recognize, and therefore may be targeted for, diverse, unrelated ligands, even in the same binding site. Said another way, if it is theoretically reasonable that a binding site can recognize unrelated ligand chemotypes, and if there are multiple convincing examples of such recognition among extant protein−ligand complexes, then we might expect such recognition to be plausible for many drug targets.

RESULTS AND DISCUSSION

The 76 proteins pairs were taken from the Protein Data Bank (PDB) (see Tables SI-1–SI-3 in the Supporting Information). In each pair, the same protein was crystallized with two different unrelated ligand complexes, then we considered whether these complexes, in the context of the biophysical considerations sketched above, support the likelihood that many proteins may recognize, and therefore may be targeted for, diverse, unrelated ligands, even in the same binding site. Said another way, if it is theoretically reasonable that a binding site can recognize unrelated ligand chemotypes, and if there are multiple convincing examples of such recognition among extant protein−ligand complexes, then we might expect such recognition to be plausible for many drug targets.
and if the poses of the two ligands overlapped. The ligand pairs were chosen for each target based on their pairwise dissimilarity; each ligand pair was compared by the widely used extended connectivity fingerprint (ECFP4) and considered if the Tanimoto coefficient (Tc) was below 0.15. Briefly, in ECFP4 fingerprints ligands are converted to bitstrings that represent the chemical environments within a four-bond diameter of each atom. The Tc represents the ratio of matching features to all possible features between two ligands and ranges between a value of 1 (identical molecules) and 0 (no features in common). For ECFP4 fingerprints, random pairs of molecules drawn from ChEMBL have average Tc values of ~0.20. In a few cases, the stringent Tc filter of 0.15 was relaxed to include protein families that are underrepresented in the PDB or in our set (all Tc values are reported in Tables SI-1–SI-3). The 76 pairs of protein–ligand complexes were selected to represent most drug-relevant protein families, with at least three examples of each class (Figure 1).

We categorized the pairs into three classes (Table 1): in class I, the same protein residues, or a subset of them, engage different ligand functional groups, sometimes reflecting intentional replacement of one functional group with a bioisostere (31 pairs, Table SI-1); in class II, different protein residues within the same binding site engage different ligand functional groups (39 pairs, Table SI-2); and in class III, the same protein residues engage the same ligand functional groups, but these groups emerge from different core ligand scaffolds (6 pairs, Table SI-3). With each complex pair superposed, the average root-mean-square deviation (RMSD) for all binding site atoms was of 0.41 Å from the 76 proteins, confirming that unrelated ligand binding did not reflect large conformational changes on the part of the protein (Figure 2A). Ligands in classes I, II, and III had average volume overlaps of 56.1%, 52.0%, and 64.9% with the complexes superposed, confirming they bound in the same site (Figure 2B). In only one pair of complexes, the class II complex of α-amylase with myricetin and with D-gluconhydroximo-1,5-lactam, was the overlap of <30% of each ligand’s volume. In >70% of the complex pairs in classes I and III, 60% of at least one ligand’s respective volume was overlapped (~54% of the pairs in class II meet this criterion). Ligand volumes ranged from 65 Å³ to 825 Å³, and the average volume difference in the ligand pairs ranges between 80 Å³ and 100 Å³ in each class (see Tables SI-1–SI-3). Overall, each of the 76 pairs of ligands occupies the same space in almost identical binding sites.

**Ligand Dissimilarity.** The average Tc across all 76 pairs is 0.10 (Figure 2C), which is a level of dissimilarity far greater than that characteristic of most scaffold hops. Since the shape and structural dissimilarity can be uncorrelated with topological dissimilarity, we compared the pairs of ligands by Rapid Overlay of Chemical Structures (ROCS) shape similarity and by TanimotoCombo, which incorporates the ROCS color score. The color score is akin to aligning pharmacophores, measuring the overlap of hydrogen bonds donors and acceptors, cations and anions, and hydrophobic groups. The average ROCS shape similarity was 0.52; only seven pairs were above the threshold of 0.75, where ligands are typically considered shape-similar. Meanwhile, the TanimotoCombo average was 0.68 and 75 pairs were below 1.15, which is substantially lower than the threshold of 1.4 for shape and chemical similarity. These levels of dissimilarity support the idea that the ligand pairs represent distant chemotypes.

**Class I Complexes.** We identified 31 pairs of complexes where the same protein residues, or a shared subset of them, interacted with unrelated functional groups from the paired ligands (see Tables SI-1 and SI-4 in the Supporting Information). We illustrate three examples here.

Retinol-binding protein 4 (RBP4) binds a benzoic urea (PDB: 3FMZ) and a pyrimido-thiophene tricycle (PDB: 4O9S) that share a Tc of 0.24 (Figure 3A). The ligand volumes overlap by 74% and 65%, respectively, in binding sites whose residues have an RMSD of 0.56 Å. Aromatic groups of each ligand occupy a deep hydrophobic pocket; a trifluorotoluene of compound 6 makes fewer contacts than the larger
two complexes, the two ligands have IC50 values of 90 and 280 

combination of electrostatic and nonpolar interactions in the π-to-face with Phe96, complemented by polar contacts with Tyr90 and the urea of

The benzoic acid of 3UON) by the M2 muscarinic G-protein coupled receptor

Leu37, and its benzoic acid forms a salt bridge with Arg121. 

accepts a hydrogen bond from the backbone nitrogen of 6 with Phe77 and His104. The urea linker of compound 7 is positioned similarly to 10—now part of a succinimide moiety—in contrast to an aromatic nitrogen in 11—now part of a triazole ring. The other carbonyl oxygen of the succinimide in 10 points directly at the face of Tyr1203, whereas this residue makes a π-contact with a methoxyphenyl in 11. The methoxyphenyl also makes an edge-to-face π-interaction with Phe1188. Although Tyr1203 rearranges to accommodate the methoxyphenyl of 11, the binding sites superpose to an RMSD of 0.88 Å. These chemically different ligands are similarly situated, contact similar residues, and inhibit TNKS with similar IC50 values of 240 nM and 33 nM, respectively.30,31

Summary of Class I Complexes. The ligands in these complexes are unrelated topologically and use different functional groups to interact with the same protein residues (for all 31 complexes, see Table SI-1 and Figure SI-3 in the Supporting Information); most pairs of ligands nevertheless bind with similar affinities. Binding site lysines and arginines may complement anions in one ligand and neutral hydrogen bond acceptors in another (as with retinol binding protein, above, or with the demethylase KDM4a; see Table SI-1 and Figure SI-3). The same aromatic residues of a binding site may complement anions in one ligand and neutral hydrogen bond acceptors in another (as with tankyrase. Not discussed in our three examples are the many examples of nonpolar protein residues that complement a wide range of topologically unrelated hydrophobic ligand side chains, as in the dithiufuroromethyl and steroid groups bound to the glucocorticoid receptor (NR3C1, Table SI-1 and Figure SI-3). In class I complexes, no single type of interaction or driving force can explain the binding of the unrelated ligands to their shared binding sites.

Figure 2. Statistics for binding site residue overlap, ligand volume overlap, and ligand similarity for the 76 complex pairs. (A) Binding site residue overlap (all heavy atoms) of complex pairs in each of the three classes and totaled overall. (B) Ligand volume overlap in each complex pair, separated by class and summed overall. Most pairs contain one ligand with 50% of its volume overlapped, indicating ligands are in the same binding pocket. (C) Dissimilarity of ligands in complex pairs for all three classes, by three different metrics: ECFP4 fingerprints, ROCS shape, and ROCS TanimotoCombo. Most are below common values used to distinguish chemically dissimilar scaffolds (dashed lines with corresponding colors).

pyrimido-thiophene tricycle of 7, which makes π-interactions with Phe77 and His104. The urea linker of compound 6 accepts a hydrogen bond from the backbone nitrogen of Leu37, and its benzoic acid forms a salt bridge with Arg121. Although the amide of compound 7 is positioned similarly to the urea of 6, it instead rotates to hydrogen-bond with Arg121. The benzoic acid of 6 also makes an edge-to-face π-interaction with Phe96, complemented by polar contacts with Tyr90 and GlN98, whereas a terminal trifluorotoluene of 7 makes a face-to-face π-interaction with Phe96. Despite the different combination of electrostatic and nonpolar interactions in the two complexes, the two ligands have IC50 values of 90 and 280 nM, respectively.26

A second example is the recognition of the agonist Iperoxo (8)27 (PDB: 4MQS) and the antagonist QNB (9)28 (PDB: 3UON) by the M2 muscarinic G-protein coupled receptor (GPCR) (Figure 3B). In both ligands, a cationic tertiary or quaternary amine ion pairs with Asp103. However, this charged amine is the only shared feature between these ligands, and they have a Tc of merely 0.05. A terminal dihydroisoxazole of Iperoxo accepts a hydrogen bond from Asn404, whereas an α-hydroxy ester of QNB both accepts and donates a hydrogen bond to the same Asn. The dihydroisoxazole of Iperoxo also superposes with a phenyl ring of QNB, making more-distant π-interactions with surrounding Trp155 and Trp400. The π-system of the alkyne linker of Iperoxo is sandwiched by Tyr104 and Trp400, whereas these residues T-stack with the phenyl rings of QNB that extend opposite the cationic nitrogen; the terminal Trp400 torsion angle is rotated nearly 60° to optimize these interactions. More generally, the binding sites shift somewhat more than is typical for this set, superposing with a 1.54 Å RMSD. The position of the ligands also overlaps slightly less, at 57% and 35% of their respective volumes. Notwithstanding their chemical, functional, and spatial differences, both ligands bind in the mid-pM range.27,29

Finally, two relatively rigid, aromatic ligands 1030 (PDB: 4OA7) and 1131 (PDB: 3UDD) inhibit tankyrase-1 (TNKS) by targeting the adenosine subsite of the NAD binding pocket (Figure 3C). Both ligands share ~66% volume overlap, but have a Tc of only 0.11. At one end, a quinolone of compound 10 and a methoxyphenyl of compound 11 both π-stack with His1201. Both ligands also hydrogen-bond with the backbone nitrogen of Asp1198; as a hydrogen bond acceptor, the amide oxygen of 10 mimics the 2-nitrogen of the 1,2,4-oxadiazole in 11. Similarly, the backbone nitrogen of Tyr1213 donates a hydrogen bond to each ligand; again, the ligand acceptors are a carbonyl oxygen in 10—now part of a succinimide moiety—in contrast to an aromatic nitrogen in 11—now part of a triazole ring. The other carbonyl oxygen of the succinimide in 10 points directly at the face of Tyr1203, whereas this residue makes a π-contact with a methoxyphenyl in 11. The methoxyphenyl also makes an edge-to-face π-interaction with Phe1188. Although Tyr1203 rearranges to accommodate the methoxyphenyl of 11, the binding sites superpose to an RMSD of 0.88 Å. These chemically different ligands are similarly situated, contact similar residues, and inhibit TNKS with similar IC50 values of 240 nM and 33 nM, respectively.30,31
Figure 3. Examples of class I complexes, where the same protein residues interact with different ligand functional groups (see Table SI-1 and Figure SI-4 in the Supporting Information). The structure overlap, binding site of each complex, 2D outline of protein–ligand interactions (blue dots are water, red dashed lines hydrogen bonds, orange lines represent hydrophobic contact; blue residue labels are common to both), and similarity metrics are shown. (A) Retinol-binding protein 4 (RBP4) bound to compound 6 and to compound 7. (B) M2 muscarinic acetylcholine receptor (CHRM2) bound to Iperoxo (8) and to QNB (9). (C) Tankyrase-1 bound to compounds 10 and 11.
Figure 4. Examples of class II complexes where different protein residues engage different ligand functional groups (see Table SI-2 and Figure SI-5 in the Supporting Information). The 3D crystal structure overlap, enlarged binding site of each complex, 2D outline of protein–ligand interactions (colored as in Figure 3), and similarity metrics are shown. (A) Coagulation factor Xa bound to DX-9056a (12) and to compound 13. (B) Progesterone receptor bound to ulipristal acetate (14) and to compound 15. (C) Glucose transporter bound to cytochalasin B (16) and to compound 17.
Figure 5. Examples of class III complexes where the same protein residues interact with similar functional groups on the ligands (also see Table SI-3 and SI-6). The structure overlap, enlarged binding site of each complex, 2D outline of protein–ligand interactions (colored as in Figure 3), and similarity metrics are shown: (A) β2-adrenergic receptor bound to timolol (18) and to BI167107 (19); (B) FABP4 bound to compound 20 and to ibuprofen (21); and (C) adenosine receptor A2a bound to adenosine (22) and to compound 23.
Class II Complexes. Class II are those complex pairs where two chemically different ligands interact with mostly different protein residues, yet still overlap in the same binding site. We categorized 39 pairs as class II, and again describe three examples, now emphasizing protein–ligand interactions unique to each ligand (a full list is given in Tables SI-2 and SI-5 in the Supporting Information).

Coagulation factor Xa recognizes DX-9056a (12) (PDB: IFAX), which has long been a characteristic inhibitor, and the novel chloroindole pyridine compound 13 (PDB: 1WU1) (Figure 4A). Whereas the IC_{50} values of the ligands are within one logarithmic unit, at 85 nM and 18 nM, respectively, they share little chemical similarity with a Tc of 0.16, especially in their warheads. Still, at a gross level, each adopts an L-shape and overlap 63% of their volumes in binding sites that superpose with an RMSD of 0.45 Å. The S1 pocket, lined by Asp189, Ser195 and Tyr228, was historically targeted with a basic amine, such as the amidinium in DX-9056a. It was thus astonishing that the chloroindole in 13 could target the same pocket by making a halogen−π interaction with Tyr228; the ability to make such halogen-bonds in coagulation-cascade proteases has enabled drugs such as rivaroxaban to overcome the pharmacokinetic problems of the cationic inhibitors. A tertiary carbon with a carboxylic acid substituent in DX-9056a makes the L-turn, whereas the shape is defined by a sulfonlipiperazene in 13. The S4 aryl-binding pocket, made up by Tyr99, Phe174, and Trp215 is the other focal point for ligand design. DX-9056a features a phenyl ring and a terminal acetimidoylpiprrolidine; the ligand’s second amidinium simultaneously hydrogen-bonds to the backbone carbonyl of Glu97 and forms a salt bridge with its side-chain carboxylic acid. In contrast, compound 13 features a 5-pyridylpyrimidine that makes no polar or ionic interactions in S4; the two linked aromatic rings π−π or T-stack with Tyr99, Phe174, and Trp215.

In a second class II pair, the progesterone receptor binds both the antagonistic steroid-analogue ulipristal acetate (14) (PDB: 4AAR) and the agonist isoxazole sulfonamide 15 (PDB: 4APU) (Figure 4B). The two molecules are unrelated topologically, with a Tc of 0.10, but overlap in the binding site by 52% and 67% of their respective volumes. Ulipristal acetate hydrogen-bonds to Thr894 through its terminal ester and forms a salt bridge with its side-chain carboxylic acid. In contrast, compound 15 only hydrogen bonds once to the tertiary carbon with a carboxylic acid substituent in DX-9056a and forms a salt bridge and hydrogen bonds with Arg126, Tyr128, and an ordered water molecule. Conversely, compound 15 only hydrogen bonds once to the receptor, and to a different residue, Asn719. The isoxazole of 15 does make a polar interaction with the same ordered water molecule as ulipristal acetate, bridging to Gln725 and Arg766. While both ligands make nonpolar interactions with the receptor, the specific residues with which they interact differ. For instance, ulipristal acetate makes π−π contacts with Phe778 and Trp755, whereas compound 15 does so with Phe778 and Tyr890. Both ligands are potent, with an IC_{50} value of 0.2 nM (14) and an EC_{50} value of 8.6 nM (15). Aside from a conformational change in Met909, the binding site residues superpose with an RMSD of 0.58 Å.

A final class II example is the glucose transporter (SLC2A1), which binds the multicyclic cytochalasin B (16) (PDB: SEQ), and a more linear phenylalanine derivative (17) (PDB: SEQQ) (see Figure 4C). Again, the IC_{50} values of each ligand are similar at 110 nM and 267 nM, respectively, yet the compounds share little topologically with a Tc of 0.15. Nevertheless, both ligands overlap 53% and 60% of their respective volumes in structures that superpose with an RMSD of 0.36 Å. Cytochalasin B hydrogen-bonds to Thr137 and Trp388, and otherwise makes apolar contacts with Asn411 and Trp412, which are characteristic of inhibitors of this transporter. Compound 17 similarly interacts with Asn411 and Trp412, but instead of hydrogen-bonding with Trp388, it forms π−π interactions with this residue, and in addition with His160, and Phe379.

Summary of Class II Complexes. The ligands in these complexes, like those in class I, are unrelated topologically, but here they more often engage with different protein residues, and less so with shared ones. Nevertheless, they bind in the same protein site, often with similar affinities (all 39 complexes are summarized in Tables SI-2 and SI-5). A mix of different driving interactions may be seen between the different ligand pairs in any given complex, such as the recognition of an amidinium or a halogen bond—by different residues—by the factor Xa, or the swapping of a π−π interaction in the SLC2A1 transporter for a tryptophan-donated hydrogen bond. In this sense, the class II complexes resemble the diverse forms of alternating interactions observed in the class I complexes. Still, the class II complexes more often swap nonpolar interactions among different hydrophobic protein residues than seemed to be true for the class I complexes.

Class III Complexes. In six pairs of complexes that we analyzed, the protein recognizes functional groups that are shared between the paired ligands (i.e., the ligand “warheads”), while the core ligand scaffolds that present these shared side chains differ. We consider three examples (a full list is presented in Tables SI-3 and SI-6 in the Supporting Information).

The β2-adrenergic receptor binds the inverse agonist, timolol (18) (PDB: 3D4S) and the agonist BI167107 (19) (PDB: 4LDE) (Figure 5A). Both ligands make conserved interactions through a cationic nitrogen and an alcohol to Asp113, Asn312, and Tyr316. The common anchor causes substantial volume overlap of the ligands: 75% and 62%, respectively. Substituted morpholines in both scaffolds make a crucial polar interaction with Asn293 deep in the orthosteric pocket and aromatic ligand groups T-stack with Phe290. Apart from the morpholine, the molecular scaffolds differ, and the overall Tc of the two molecules is 0.17. An additional tolyl on BI167107, which interacts with Trp109, and different polar groups are responsible for the topological dissimilarity between the two ligands. A thiadiazole of timolol hydrogen bonds to Thr118, while BI167107 makes two additional hydrogen bonds to Ser203 and Ser207. The binding site residues of the two complexes overlap with an RMSD of 0.64 Å; both ligands bind the β2-adrenergic receptor with subnanomolar affinity.

A second characteristic class III pair involves fatty acid binding protein 4 (FABP4), which binds both indole compound 20 (PDB: 5D4S) and ibuprofen (21) (PDB: 3P6H) (Figure 5B). The carboxylic acid of each ligand makes a salt bridge and hydrogen bonds with Arg126, Tyr128, and an ordered water molecule bridging to Arg106. The rest of the recognition elements in the site appear to be hydrophobic and it is perhaps unsurprising that ligands have found several topologies to complement these residues, resulting in a pairwise Tc of 0.14. Compound 20 is much larger, with two additional phenyl rings that displace members of a water network around ibuprofen. The size difference between the two ligands is reflected in only 44% of the larger compound 20 being overlapped, compared to 80% of ibuprofen. The binding
sites of each complex superpose with an RMSD of 0.29 Å and both ligands bind with $K_i$ values of $\sim 100$ nM.\textsuperscript{39,40}

A final example of class III complexes is shown with the $\alpha_2A$ adenosine receptor, a GPCR, which binds adenosine (22)\textsuperscript{41} (PDB: 2YDO) and a triazine- and chlorophenol-containing antagonist 23 (PDB: 3UZC) (Figure 5C). Both ligands hydrogen bond to Asn253 and His278 through an extracyclic nitrogen on nitrogen-rich heterocycles and hydroxyl groups, respectively. Although the agonist and antagonist share these polar contacts and overlap 65% and 54% of their respective volumes in the binding site, they differ chemically, with a Tc of only 0.09. Whereas adenosine makes several hydrogen bonds with Glu169, Ser277, and a bridging water molecule to His250, compound 23 makes a halogen–π interaction with His278 and T-stacks with Trp246 and His250 through two phenyl rings. Both ligands bind well, with $K_i$ values of 30.9 nM and 1.41 nM, respectively.\textsuperscript{41,42} The binding site residues superpose with an RMSD of 0.98 Å.

**Summary of Class III Complexes.** As in the class I and class II complexes, the class III ligand pairs are unrelated topologically. Unlike class I and II, the differences in the class III ligands occur in the core scaffolds, not the ligand side chains most involved in interacting with the proteins. Accordingly, there can be more similarity in the interactions between the unrelated ligands. Still, even among these complexes, one observes the same switching of driving interactions that prevails in the class I and, to a lesser extent, the class II complexes: the atom types of donors or acceptors can change, hydrogen bonds themselves may be swapped for dipole–quadrupole interactions, and there is great plasticity in the nonpolar interactions. A theme that emerges in all three classes is that there are multiple topologically unrelated ligand

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**Figure 6.** Complex pairs where water mediates interactions with one ligand whereas the second ligand makes these interactions directly. The structure overlap, binding site of each complex, 2D outline of protein–ligand interactions (colored as in Figure 3), and similarity metrics are shown. (A) Pteridine reductase co-crystallized with compound 24 and with compound 25. (B) Renin co-crystallized with compound 26 and with compound 27.
groups that can recognize most receptor side chains and binding site environments.

**Water Molecules Mediating Ligand Binding.** Water molecules can mediate interactions between ligand and protein\(^3,4,44\) to enable more diverse ligands to bind in the same site. Of the 76 pairs studied, 45 had at least one water molecule; 13 of them bridged one ligand to a residue that was directly engaged by the other ligand in the pair. We illustrate two examples.

Pteridine reductase binds both the benzimidazole 24\(^{35}\) and the flavanol 25\(^{46}\) (Figure 6A). The chemically unrelated ligands have a Tc of 0.05 and overlap well in the biotin pocket (80% and 39% of each ligand's volume). Compound 25 hydrogen bonds directly to Asp161 through its 3' phenolic oxygen, which also engages two ordered water molecules, forming a hydrogen bond network with Cys168 and Asn175. This phenolic oxygen is 1 Å from a superposed water molecule from the cocystal of 24. In this second complex, the imidazole nitrogen of compound 24 hydrogen-bonds to this water molecule, which connects to the water network with the same residues, Asp161, Cys168, and Asn175. The 7-OH of 25 hydrogen-bonds to Ser95 and to the NADP cofactor, whereas a water molecule bridges the exocyclic nitrogen of 24 to the NADP cofactor; the water molecule is again <1 Å from the superposed 7-OH of 25. The two binding sites superpose with an RMSD of 0.18 Å; the \(K_i\) of 24 is 288 μM, while the IC\(_{50}\) of 25 is 104 μM.

A second example of the role of water is shown in the complexes of renin with the flexible polyhydroxy 26\(^{77}\) (PDB: 1HRN) and with the more-rigid dianimopyrimidine benzoxazine 27\(^{48}\) (PDB: 2G1Y) (Figure 6B). Here, too, the ligands are chemically unrelated with a pairwise Tc of 0.06. Nevertheless, 43% and 58% of their respective volumes overlap in the site, and they each interact with several of the same residues, which superpose with an RMSD of 0.26 Å. This is partly explained by bridging water molecules, one of which bridges the terminal dianimopyrimidine of 27 and Ser76, while a hydroxyl of 26 displaces this water and hydrogen-bonds directly to the same Ser76. Similarly, in the S3 subpocket, a water network bridges a hydroxyl in 26 to the backbone of Tyr14, while compound 27 hydrogen-bonds directly to the same Tyr14. Meanwhile, the IC\(_{50}\) values of the two compounds are within one logarithmic order at 9 and 90 nM, respectively.

**CONCLUSION**

A key observation from this study is that many proteins can bind unrelated ligands in the same binding site. This is true for all major target families, including GPCRs, transporters, ion channels, nuclear hormone receptors, and enzymes. Often, the more deeply that a target is studied, the more chemotypes one can find for it. Consider the long-studied \(\mu\)-opioid\(^{49}\) and estrogen receptors, for which at least 22 and 21 unrelated chemotypes, respectively, may be distinguished from the known ligands with <1 nM affinity in the ChEMBL database\(^18\) (using an ECFP4 Tc ≤ 0.35, suggesting scaffold hops\(^21\)). The recognition of different ligands that overlap in the same site can occur in several ways: wholly unrelated ligands can interact with largely the same residues (our class I), they can bind in the same site but engage different residues within it (our class II), or unrelated scaffolds might project similar functional groups to interact with the same protein residues (our class III). No single type of driving force seems to explain this plasticity, with multiple types of functional group recognition occurring in many of the pairs of ligands (salt-bridges exchanged for ion-dipole interactions, one type of hydrogen-bond donor or acceptors swapped for another, hydrogen bonds replaced with dipole quadrupole interactions, and great plasticity among nonpolar side-chain interactions). An encouraging inference of this study is that structural complementarity allows for the discovery of entirely new chemotypes, even for heavily studied targets.\(^{13,14}\)

At first glance, the recognition of diverse ligands by the same site may seem unsurprising. For many targets, multiple chemotypes will be well-known to specialists; for instance, opioid researchers are conscious of morphine, fentanyl, oliceridine,\(^{50}\) and others,\(^{14}\) where pairwise ECFP4 Tc values range from 0.12 to 0.19. The ability of unrelated proteins to bind the same ligands—the logical complement of this study—has also been shown both for substrates—leading to enzyme moonlighting and perhaps evolution\(^3\)—metabolites\(^52\) and for synthetic molecules.\(^{11,33,34}\) Meanwhile, medicinal chemists have long used bioisosteres to exchange topologically unrelated functional groups. Well-known examples include the replacement of carboxylates with tetrazoles, hydroxamic acids, and isoxazoles,\(^{56}\) the replacement of benzene with thiophene,\(^{17}\) the replacement of catechols with benzimidazoles,\(^{56}\) and the replacement of phosphates with salicylates or isothiazolidinones.\(^{20}\) Of the 76 pairs of protein–ligand complexes studied (152 total complexes), 32 reflected such bioisosteric swaps. More surprising, but just as common in this set, are cases where the same protein residue interacts with pairs of ligand functionalities that are not classically bioisosteric, including the hydrogen bonding or π-stacking of a tryptophan with either a ligand lactam or ary group (as in 16 and 17 interacting Trp388 of the glucose transporter, Figure 4C) or the interaction of a cation or a halogen in the S1 pocket of factor Xa (Figure 4A). From a biophysics standpoint, we are taught to consider energies of interacting groups, not their topologies. From this perspective, two ligands that both present an anion to interact with a threonine will likely capture similar interaction energies, after correcting for \(pK_a\) differences, regardless if one is an isoxazole phenolate and the other is a carboxylate, as in glutamate and AMPA binding to the ionotropic glutamate receptor (see Tables SI-1 and SI-4).

Even nodding to this biophysical understanding, what emerge from this study are the wide range of targets for which unrelated ligands can bind, and the profound nature of the dissimilarity. Thus, classic bioisosteric swaps are a minority of our cases, and even where they can be detected they rarely represent the largest change. Especially in classes I and II, no single chemical switch links most of the ligand pairs. None of this breaks the laws of biophysics, of course, but that is one of the key points of this study.

Certain caveats merit airing. We do not pretend to be comprehensive in this study: there are likely many other pairs of dissimilar ligands binding to the same site. Conversely, and more generally, unrelated ligands binding to the same site remain exemplary exceptions, not the rule; molecules falling into related families dominate most ligand–protein complexes, as this reflects the history of the medicinal chemistry undertaken against them. Consequently, the ratios of our classes I, II, and III are not meaningful in themselves, nor is the ratio of complexes with unrelated ligands meaningful to those with related ligands. Nor do we pretend that even our classes are definitive—others might easily be used. Even among our classes, choosing the class into which a pair of complexes fell
was often a judgment call; we had no fully quantitative metric for assigning a pair of complexes to one class or the other, although often the choice seemed clear. Although we sought examples from multiple protein families, we cannot confidently write that all proteins will recognize unrelated chemotypes. Although the binding sites of each complex pair overlap by crystallography, they may differ sufficiently to change the energy landscape. Finally, using any single metric, such as topology-based Tanimoto coefficients, to characterize chemical similarity can mislead, especially when one molecule is much larger than the other.

These caveats notwithstanding, it seems clear that many proteins can bind—and many others should be able to bind—unrelated chemotypes in the same site. While the study was not comprehensive, it was extensive, represented many families of drug targets, and it was unbiased in the selection of complexes. Crucially, biophysical considerations, such as the equivalence of topologically unrelated groups, and the size and diversity of the chemical space from which such groups might be drawn, bolster the empirical observations described here. Thus, while we make no statistical argument as to the relative diversity of the chemical space from which such groups might be drawn, bolster the empirical observations described here. The ability of most binding sites to recognize diverse and even unrelated chemotypes supports the community’s longstanding program of structure-based novel ligand discovery, and the new biology that they promise.

■ METHODS

Protein–ligand complexes were selected in two ways. First, a list of PDB structures containing ligands, solved by X-ray crystallography to a resolution of better than 2.5 Å, was compiled using the “Advanced Search” interface in RCSB.org. All entries with Uniprot identifiers were processed using the KNIME analytics platform (https://www.knime.com). Of 70879 PDB structures with 17892 ligands, we excluded the ~11% that were crystallized with only one ligand, and another ~6% with frequent ligands such as buffer components and ions. Only entries with at least 20 unique ligands were selected, thereby reducing this sizable list to 188 Uniprot IDs with up to 389 ligands for a single target. Using the RDKit Fingerprint module within KNIME, we calculated pairwise ECFP4-based Tcs, and generated similarity matrices between all ligands for each Uniprot ID. Using these matrices, we prioritized complexes with maximally dissimilar ligands; these were visually inspected in PyMOL and in DiscoveryStudio 4.1 (Biovia, Inc.).

Second, to account for under-represented protein classes, the list of complexes was complemented by 20 unique ligands. After removing buffers, ions and other crystallizing agents, this left 15120 complexes, corresponding to 1306 unique Uniprot IDs. Again, a pairwise ECFP4-based Tc was calculated for all ligands within each Uniprot ID. The lowest Tc pair from randomly selected protein structures were visualized until each protein major class was represented. If the lowest Tc-pair seemed inappropriate—for instance, if there were large domain shifts in the protein, or if the ligands were not in the same binding pocket—then other pairs of ligands were visualized.

Initially, 365 unique PDB pairs were visualized in PyMOL, using a script that was optimized to superpose the structures based on their fold. A maximum of 10 ligand–ligand receptor pairs were visualized for each protein; these were the lowest pairs for that protein, as determined by pairwise Tc similarity of their ligands. For each protein, we selected the first appropriate complex that we visualized, based on the evaluation of hydrogen bond networks, hydrophobic stacking interactions, water-mediated interactions (where applicable), interaction with individual amino acids, and superposition of ligand volumes. Ultimately, 76 protein–ligand complex pairs were selected for 71 unique proteins and 151 unique ligands; five proteins are duplicated, as are two ligands.

To calculate the ligand volume overlap for each complex pair, the two PDB structures were superposed using the MatchMaker function in Chimera, and the overlapping ligands saved. The volume of each ligand was calculated with Python scripts that computes van der Waals spheres for each atom. The two ligands are virtually combined and their joint volume is calculated. The volume overlap of two ligands is found by adding the individual volumes and subtracting the joint volume. ROCS shape-based overlap values were computed using ROCS Release 3.2.0.4 from OpenEye. The same ligand input was used for the volume ROCS calculations.

Three-dimensional structural images were generated with PyMOL. Two-dimensional ligand–receptor interactions were generated using LigPlot and Pose view (a part of the ProteinsPlus web service), the interactions were manually merged in ChemDraw Prime 15.1, to ensure same ligand site orientation.

■ ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acschembio.8b00443.

Full list of complex pairs from classes I, II, and III, including binding site RMSD, ligand similarity metrics, volume overlap, and 3D comparisons (PDF)

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Notes
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■ REFERENCES


