



Docking and chemoinformatic screens for new ligands and targets

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Computer-based docking screens are now widely used to discover new ligands for targets of known structure; in the last two years alone, the discovery of ligands for more than 20 proteins has been reported. Recently, investigators have also turned to predicting new substrates for enzymes of unknown function, taking docking in a wholly new direction. Increasingly, the hit rates, the true-positives, and the false-positives from the docking screens are being compared to those from empirical, high-throughput screens, revealing the strengths, weaknesses, and complementarities of both techniques. The recent efflorescence of GPCR structures has made these quintessential drug targets available to structure-based approaches. Consistent with their 'druggability', the docking screens have returned high hit rates and potent molecules. Finally, in the last several years, an approach almost exactly opposite to docking has also appeared; this pharmacological network approach begins not with the structure of the target but rather those of drug molecules and asks, given a pattern of chemistry in the ligands, what targets may a particular drug bind to? This method, which returns to an older, pharmacology logic, has been surprisingly successful in predicting new 'off-targets' for established drugs.

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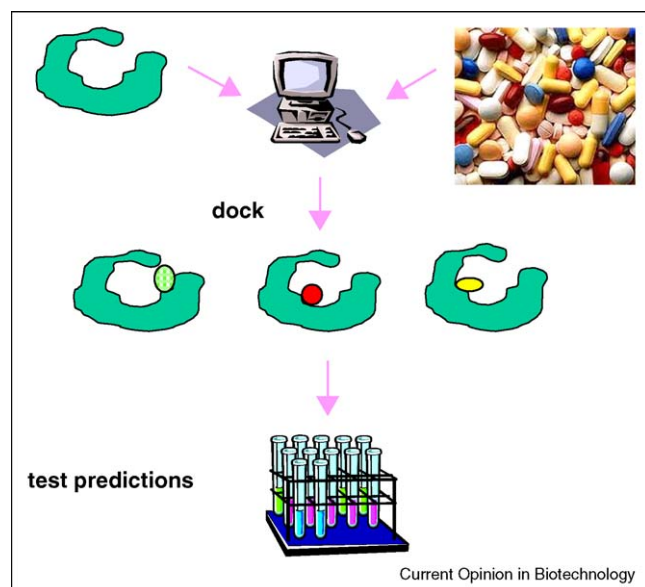
Since the work of Goodford in the mid-1970s [1], protein structures have held the promise of guiding the design of drugs. As is often true, the early potential of the field was largely unmet, and in the early 1990s there was a sense that 'structure-based drug design' was not pragmatic. In the last decade, however, the use of structure-based computational design has made steady, if often unspectacular [2] technical progress, to the point where most pharmaceutical organizations have substantial groups

devoted to its application. If one asks, 'How many drugs have been discovered entirely by structure-based methods?', the answer will be 'none', as these methods largely contribute only to the discovery and early optimization of leads for drug design. If one asks, conversely, 'To the development of how many drugs have structure-based methods been critical?' then the answer will be close to 10. Whereas this number must seem small, it is put into perspective by the number of drugs that owe their origin to empirical high-throughput screening (as of this writing, only two or three [3], though many more are now in trials or awaiting approval).

Perhaps the area where structure-based methods have had the most quantifiable impact is in computational screens of large compound libraries, looking for new chemical matter that will bind to and modulate a protein of known structure. These 'virtual' or 'structure-based' screens typically use molecular docking programs to fit small organic molecules into protein structures, evaluating them for structural and chemical complementarity. Several million molecules may be docked into the structure of the target protein, and those that fit best, according to the docking scoring function, will be tested experimentally (Figure 1). Although these scoring functions retain substantial inaccuracies, the focus on commercially available molecules has made failure cheap — since one can always just purchase and test the next set of compounds — and so pragmatic. In the last two years alone, more than 20 papers have appeared in which docking screens were used to predict a ligand, which was then subsequently confirmed by experiment (Table 1). Most of these studies emerged from academic research groups; many others that have been conducted in industrial groups remain unpublished.

Here we focus on research directions in structure-based screening that, even very recently, would have been hard to anticipate. We begin with efforts to combine virtual and high-throughput screens. These campaigns enable the discovery of false-positives and false-negatives for both techniques, and their combination reveals complementary strengths. We then turn to the use of structure-based screens against membrane-bound receptors, the structures for which have only recently become available. Finally, we consider methods that turn the structure-based paradigm on its head, asking not what ligands might be discovered based on the structure of the targets, but rather what targets might be discovered from the identity of the ligands.

Figure 1



Structure-based screens for novel ligands. Large libraries of commercially or otherwise available compounds are fit into target structures by a docking computer program. Each molecule is sampled in thousands to millions of possible configurations and scored based on structural complementarity to the target protein. Of the perhaps millions of molecules in the library, tens to hundreds of top-scoring hits are subsequently tested for activity in an experimental assay.

Virtual versus high-throughput screening

A stringent test of a docking screen is to compare it to a high-throughput screen (HTS), where every single molecule in a library is empirically tested, not just the top-

scoring 50 or several hundred suggested by docking. This could reveal not only false-positives but also false-negatives of docking, which would otherwise always remain opaque, but also whether docking hit rates were high enough to justify the focus on only a relatively small number of candidate ligands. In studies beginning in the early 2000s, this turned out to be the case, with docking hit rates being 10-fold to 1000-fold higher than those returned by HTS against the same target [25–28]. Whereas these early studies were encouraging, the full set of docking predictions was not compared to the HTS results, and sometimes different compound libraries were used.

In recent studies exactly the same molecules have been docked and screened empirically, against exactly the same target. In a quantitative HTS (qHTS), where each compound is tested in 7-point dose response) of more than 70 000 compounds against AmpC β -lactamase, qHTS yielded 1274 initial hits. A detergent counter-screen revealed 95% of these to be colloidal aggregators, which are typically the largest source of false-positives in biochemical assays [29]. Of the 70 ‘hits’ that remained, 25 were β -lactams, which are known to covalently bind to β -lactamase, and so are uninteresting except as controls, 24 were irreproducible on retesting, 9 were aggregators that resisted the low levels of detergent used in the counter-screen, and 9 were promiscuous covalent inhibitors [20*]. No reversible, specific, competitive inhibitors were found by HTS whatsoever.

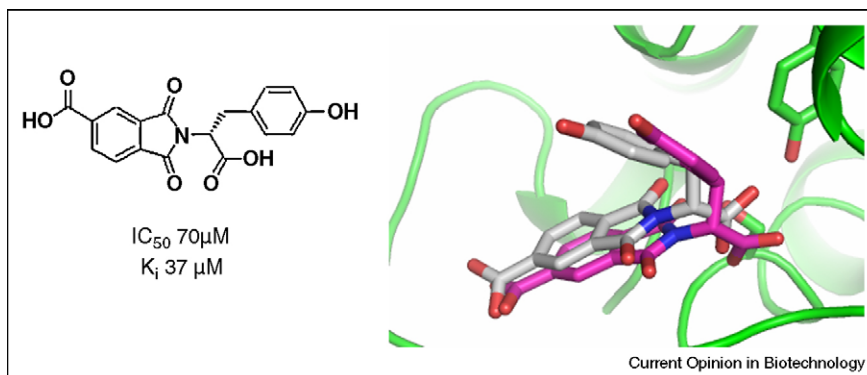
This result was unexpected — a goal of the exercise had been to reveal docking false-negatives, but with no new molecules discovered by the qHTS, this was impossible.

Table 1

Docking predictions subsequently confirmed by experiments: 2007 to mid-2009

Target	Docking program	Lead inhibitor IC ₅₀ (μ M)
AdoMetDC [4]	Glide	12
AHAS [5]	DOCK 4/AutoDock	15.2
Aldose reductase [6]	N/A	0.53
CDC25 phosphatase [7]	FRED/Surflex/LigandFit	13
DNA gyrase [8]	DOCK 5	50
EphB4 [9]	DAIM-SEED-FFLD	1.5
FFAR1 [10]	Glide	3.6
Histamine H4 [11]	FlexX	95.8
Human pregnane X [12]	Surflex	0.049
MCH-R1 [13]	ICM	7.5
Pim-1 kinase [14]	Glide	0.091
PNP [15]	GOLD	18.9
PPAR- γ [16]	Glide/IFD	2.9
Tm0936 [17]	DOCK 3.5	$10^5 \text{ M}^{-1} \text{ S}^{-1}$ ($K_{\text{cat}}/K_{\text{m}}$)
TRH-R1/TRH-R2 [18]	FlexX	0.29
β_2 -Adrenergic receptor [19*]	DOCK 3.5.54	0.009
β -Lactamase [20*]	DOCK 3.5.54	140
SHP2 [21]	DOCK	100
AI-2 quorum sensing [22]	DOCK 5	35
Anthrax edema factor [23]	HINT/AutoDock	1.7
hPRMT1 [24]	GOLD	12

Figure 2



A docking-derived novel β -lactamase inhibitor. Structure of the new inhibitor and overlay of the crystal structure (white carbons) to the initial pose proposed by DOCK (magenta carbons). The RMS deviation of the heavy atoms between the two poses was 0.9 Å.

To explore whether the empirical screen suffered from false-negatives, 16 high-ranking docking hits were retested in low throughput at concentrations higher than used in the qHTS campaign (where the maximum concentration of compounds was 30 μ M). Two mid-micromolar competitive inhibitors were found, one that had ranked 80th and one that had ranked 200th of the 66 000 library molecules docked. These compounds had K_i values of 37 and 55 μ M, and IC_{50} values higher still, which explains why they were missed by the qHTS. Subsequently, a crystal structure of AmpC in complex with one of these inhibitors showed close correspondence between the X-ray structure and that predicted by docking (Figure 2) [20*].

In a search for formylpeptide receptor ligands, approximately the top 1% of the library ranked by virtual screening was tested by HTS, and nine chemical families of hits were confirmed [30]. In another study, a combination of 2D and 3D similarity approaches was used to screen 10 000 compounds against the estrogen receptor GPR30. Biochemical screens of the top 100 ranked compounds identified a potent ligand ($K_i = 5.7$ nM) [31]. These studies illustrate how prioritization of compounds to be tested can lead to the identification of true hits. However, they do not answer the question of how many scaffolds were missed by virtual screening. To do so one needs to screen exactly the same library by both methods and compare the hits. A good example of such a study was reported in a search for GSK-3 β in which four out of six scaffolds found by HTS were found in the top 1% by docking [28], indicating the potential for using virtual screening as a guide in the follow up of HTS hits. Unfortunately, few direct comparisons between these methods are currently reported, and a bigger number of similar studies would help to draw conclusions on how frequently such success is attained by virtual screening.

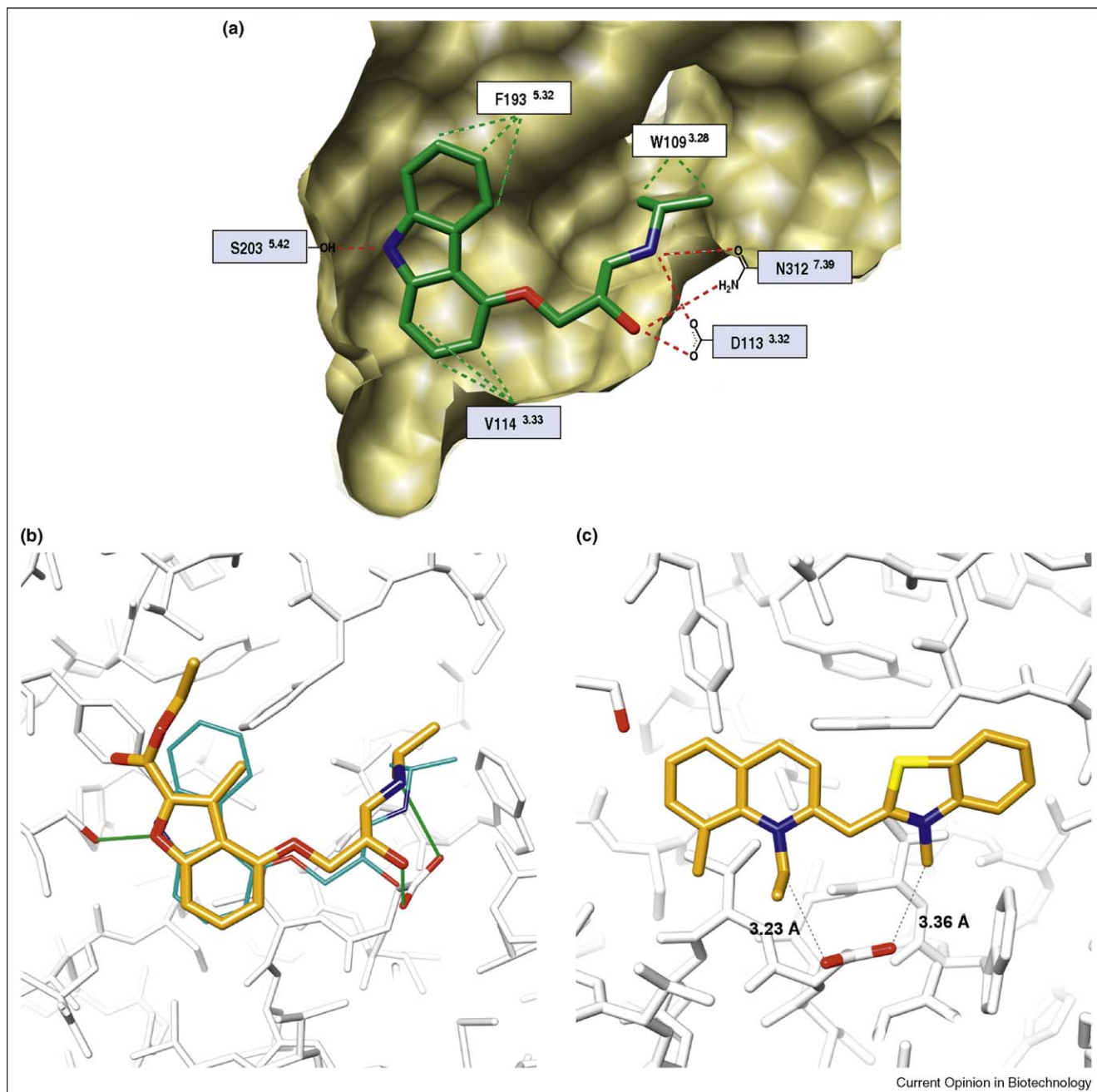
At this early point, three tentative lessons may be drawn: first, both docking and HTS suffer from false-positives and false-negatives; second, that an intense amount of work is required to follow up and confirm screening hits; and third, that the two techniques may complement each other. These conclusions are supported by a new docking and qHTS campaign of now 198 000 molecules against the enzyme cruzain, with the proviso that here, finally, screening revealed docking false-negatives (unpublished results). How reliable these conclusions are, and whether we may expect docking and HTS to be combined as a standard procedure, must await the outcome of further studies now underway.

Docking to membrane proteins

Structure-based approaches to ligand design for membrane proteins have been hampered by a lack of X-ray structures. This was especially grave in the case of guanine nucleotide-binding protein (G-protein)-coupled receptors (GPCRs), where, until recently, only the structure of rhodopsin was known [32]. This absence was keenly felt, as 25% of drug targets are GPCRs, and 40% of drugs bind to these targets [33]. To overcome this gap, protein structure modeling was widely used to generate 3D structures as scaffolds to which to dock. In the past two years, modeling was mainly concerned with the refinement of the 'raw' modeled structures. These structures were then successfully used in several docking screens, although the hit rates remained low [13,18,34*].

The recent determinations of the structures of the β_2 -adrenergic (β_2 AR) [35,36*], the β_1 -adrenergic [37*], and the adenosine A_{2A} [38*] GPCRs reveal why these receptors are wonderful targets for small molecules. The binding sites almost entirely enclose the ligands, ensuring close complementarity. Each site combines a mixture of polar groups, allowing for specificity, and non-polar ones, for affinity (Figure 3). The use of an X-ray structure for

Figure 3



The GPCR β 2AR as a target for structure-based screening. **(a)** Side view of the orthosteric site, the proximal portion of the protein has been cut away for clarity. Several key interacting residues are marked. Red and green dashed lines indicate polar and hydrophobic contacts, respectively. Residues in a light-blue box are essential for agonist and antagonist binding. **(b)** Comparison of the docked pose of a docking-derived, 9 nm inverse agonist [19^{*}] (gold carbons), with the X-ray structure of carazolol (cyan). Hydrogen bonds are shown as green sticks, and residues Asp-113 and Ser-203 are emphasized with red oxygens. **(c)** Predicted binding mode of a docking-derived novel ligand [19^{*}] (gold carbons), chemically distinct from previously known β AR ligands. The distances between the alkyl substituents and the respective closest oxygen of Asp-113 are shown as dashed lines.

docking should be a substantial advantage over modeled structures, not least because of how the structure reveals the precise layouts of the receptor binding sites and their interactions with the ligand. With these new structures, can we anticipate an efflorescence of ligand discovery,

will they be better templates for discovery than the homology models that preceded them, and how will the new ligands compare to those discovered, over the last 50 years, by traditional ligand-based methods, which progressed without the advantage of a crystal structure?

In a screen against the β_2 AR, about a million ‘lead-like’ molecules from the ZINC database (<http://zinc.docking.org>) were screened against the structure of the target. Twenty-five were chosen from the top 400 docking-ranked molecules and tested in ligand displacement assays. Six modulated β_2 AR with affinities between 9 nM and 4 μ M, a hit rate of 24% [19^{*}] (Figure 3). Intriguingly, five of these were inverse agonists, as was the ligand bound in the X-ray structure, carazolol, against which the screen occurred. In a similar study, an in-house database of 400 000 ligands was screened against β_2 AR [39]. 36% of the molecules tested were active, with the best having a 0.114 nM K_i . An innovative experiment was to dock for pharmaceutical chaperones of misfolded rhodopsin [40]. About 24 000 compounds were docked from the NCI database against the X-ray structure of the P23H mutation of opsin, which is associated with retinitis pigmentosa. Five docking prioritized compounds were tested, one of which was weakly active as an inhibitor of opsin regeneration.

The relatively high hit rates and high potencies of the ligands to emerge against the X-ray structures support the idea that these are better templates for discovery and design, compared to the earlier homology models. This is borne out in a community-wide, blind assessment (GPCR Dock 2008 [41]) of the prediction of the structure of the human adenosine A_{2A} receptor in complex with the ligand ZM241385 [38^{*}]. Twenty-nine groups submitted a total of 206 structural models before the actual publication of the X-ray structure. The best model had a ligand rmsd of 2.8 Å and a C α rmsd of 3.0 Å, but ranked only second among the models submitted by that participant. Slightly more discouragingly, the average rmsd of the ligand predictions was 9.5 Å, despite an average C α rmsd of only 4.2 Å between protein model and X-ray structure. This indicates that docking did not work well and is sensitive to changes in protein structure, a point that has been made before.

By docking standards — where a 5% hit rate is considered substantial, and where a ‘hit’ might have a mid-micromolar affinity, the results in the GPCR docking screens seem extraordinary. This is partly explained by the substantial bias in even our commercial libraries toward GPCR-like ligands, the product of 50 years of intense medicinal chemistry in this area. On the other hand, the β_2 AR and opsin docking screens suggest that despite the attention lavished on these targets, novel chemotypes, with arguably new biology, may yet be found. Both observations support the idea that further structure-based screens against GPCRs, both those now determined and the new structures that are eagerly anticipated, will merit the effort. A challenge for the future will be leveraging the often antagonist, or inverse agonist, bound complexes to discover agonists; recent work from the Rognan and Abagyan labs suggest that this will be possible [42,43].

Predicting the activities for enzymes of unknown function

In the early years of docking several investigators entertained the idea of predicting not only what would inhibit an enzyme or a receptor, but also their true physiological substrates and agonists. For two reasons the idea was put aside. First, we could not then (early 1990s) imagine many proteins whose structures had been determined but whose functions remained unknown — why would anyone go to such trouble? Second, predicting activity seemed much harder than predicting inhibitors of activity, notwithstanding the relevance of both for drug discovery and for biological understanding. This idea thus remained little more than idle conversation for two decades.

With the advent of the Structural Genomics projects, however, an increasing number of proteins of unknown function have had their structures determined, more than 50 such are cataloged by the Protein Databank (PDB) alone. This has inspired several groups to return to the challenge of predicting the enzyme activity. Whereas the original technical concerns remain germane, not least because of the difficulties in modeling conformational changes to which activity is often coupled, there are preliminary signs of progress. Thornton and colleagues have investigated docking a limited set of metabolites against the structures of short-chain dehydrogenases/reductases, exploring different docking protocols in retrospective studies [44]. An interesting aspect of this work is the use of representative metabolites to represent a class of molecules, reducing computational costs. In more prospective work, Hermann and co-workers and Xiang and co-workers have predicted and tested the activities of two targets from structural genomics — the amidohydrolases Tm0936 [17] and Dr0930 [45]. Docking the high-energy intermediate, transition-state-like forms of the KEGG metabolites, Tm0936 was correctly predicted to act as a deaminase of adenosine and δ -adenosine homocysteine, while Dr0930 was correctly predicted to act as a lactonase. Whereas both of these predictions were confirmed experimentally, close inspection of the results reveals the strengths and weaknesses of the approach. For Tm0936 the docking was almost entirely correct — not only was the right substrate predicted, but also was its docked geometry compared to a crystal structure of the product of the reaction that was subsequently determined [17]. With Dr0930, conversely, whereas the docking identified lactones as a general class of substrate, the precise preferences among lactones was *not* captured, as δ -lactones were predicted to be as good as, often better than γ -lactones, when in fact the reverse was found to be true. Thus docking the high-energy intermediate forms of the metabolites was grossly successful, but missed important particulars.

Thus it is too early to tell whether a structure-based approach to function prediction has yet overcome the

concerns that kept it on the sidelines for the last two decades. What can be said is that it is no longer so sidelined, and is being pursued actively by several groups. Hopefully, two years from now when the next version of this review becomes timely, we will have a better sense if this approach has advanced to the point of reliability — for now, it remains intriguing enough, with enough preliminary successes, to justify a focused research effort.

Polypharmacology and the chemical view of biology

The logic used by classical pharmacology inverts the target-oriented approach of molecular biology — in pharmacology, biology is characterized by the actions of organic small molecules. Even now, most receptors are characterized by more or less specific ligands that, for instance, distinguish α -adrenergic from β -adrenergic receptors, and β_1 subtypes from β_2 . Recently, computational chemists and biologists have returned to this older chemical view, quantifying the relationships among receptors based not on their structures or sequences, but on chemical patterns among the ligands that bind to them. This has revealed connections among targets that to a traditional pharmacologist — now largely extinct — might seem entirely reasonable, but to the now dominant molecular view will be surprising.

A seminal paper in this area, by Hopkins and colleagues when at Pfizer, mapped the links among targets as articulated by shared ligands. Many targets that were unrelated by sequence or structure nevertheless had common ligands [46]. Similarly, Vidal and colleagues adapted network techniques developed for protein–protein association to drug–target associations [47]. Many drugs shared multiple targets, and unsurprisingly the converse was also true. Older drugs were often distinguished from newer, sometimes investigational drugs by the greater specificity of the latter, which the authors attributed to the influence of target-based or ‘rational’ design. A comparison of drug targets to those directly involved in disease etiology suggested that most drugs bound to proteins distant from the etiological targets, implying that most drugs did not work on the cause of the disease, but were rather palliative.

In contrast to the trends toward specificity inferred by Vidal, Roth has argued that polypharmacology is not only common but that, especially in the CNS, it is often essential for efficacy [48] (polypharmacology is taken to mean the action of one drug on multiple targets at physiologically relevant concentrations). Effective CNS drugs typically target at least two targets, such as serotonin and dopamine subtypes, and often more; genuine specificity correlates with lack of efficacy. Roth has established an NIH Roadmap Psychoactive Drug Screening Program (PDSP, <http://pdsp.med.unc.edu/indexR.html>) to screen pharmacologically active agents against a broad

spectrum of targets, including more than 300 GPCRs and, increasingly, ion channels, transporters, and kinases. Recently, systematic screening has revealed that the histamine H1 receptor can be responsible for weight gain under antipsychotic treatment, an enormous problem for drugs such as Zyprexa, that the kappa opioid receptor is the target of the potent hallucinogen salvinorin A, and that the clozapine metabolite desmethylclozapine activates M1 muscarinic receptors, which may contribute to the clinical efficacy of the parent antipsychotic [49].

In collaboration with Roth’s group, we have tried to predict and test new off-target effects of established drugs. Drawing on a systematic map relating drug targets by their ligands, using BLAST-derived algorithms to normalize for random chemical similarity, we found that targets related by ligands quantitatively differed from those related by sequence [50•]. For instance, the serotonergic 5HT-3 receptor was closely related to the 5HT-4 receptor, even though the former is an ion channel and the second a GPCR; both bind congeners of serotonin. Similar relationships among the ligands for multiple ionotropic (ion channels) and metabotropic (GPCRs) receptors, and even transporters, were observed. On the basis of these observations, new off-targets were predicted, including the antagonistic effects of the μ -opioid methadone on M3 muscarinic receptors, the activity of the antiparasitic emetine on α -adrenergic receptors, and the activity of the gut μ -opioid loperamide on NK2 receptors. When tested in the Roth lab, all three had low or submicromolar activities on their predicted off-targets. Similarly, Bork and colleagues at the EMBL have analyzed the shared side effect profiles of 746 drugs using a text-based analysis of drug inserts, correlating these with chemoinformatic similarities. A network of 1018 drug–drug relations was revealed, 261 of which were formed among dissimilar drugs from different indications. Twenty specific drug off-targets were tested, and 13 were confirmed experimentally [51••]. Other studies, which return to a more structure-based method for predicting off-target effects, including docking, have also revealed new off-targets, though these efforts remain at an earlier stage than the chemoinformatics approaches [52].

Drug polypharmacology is common [53], occasionally essential, often has unwanted side effects [54], and can cross classic molecular categorizations. The most successful methods to characterize and, occasionally, predict [50•,51••] such off-target effects have been those that are strictly ligand-based, but is there a way to understand these effects from a molecular target view? In a fascinating recent paper, Klebe and colleagues compared the structural similarities of binding sites among often unrelated targets, and find that the similarity among these is often unrelated to the sequence identities of the proteins that harbor them [55]. It may be that as the ligand-bound

structures of more and more targets become available, we will be able to understand these cross target activities. For now, the older, pharmacological organization is, paradoxically, the more generative for predicting and classifying polypharmacology than the structure-based approaches which have been our primary focus here [56*].

Conclusions

With all their weaknesses, docking screens are now common in molecular discovery. The ongoing growth of molecular structures, not least among membrane targets, will continue to widen the remit of this technique. In upcoming research one can look not only for the discovery of new molecular entities, but also for their genuine application to pragmatic problems in drug discovery and chemical biology. In the short term, the highest impact studies may well come from the chemoinformatics approaches that return to an older, classical pharmacology view of drug action, as these work with drugs themselves, whose impact on human health and biology is already clear.

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