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Dynamic template-assisted strategies in fragment-based drug discovery

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Fragment-based methods for drug discovery are increasingly popular because they provide drug leads with greater ligand efficiency than conventional high-throughput screening. However, established methods for fragment detection do not address the central question in fragment-based ligand discovery: how can a primary ligand be optimally extended by a secondary fragment? Dynamic screening methods solve this issue by using a protein target as a template for ligand assembly, thus yielding high-affinity binders from low-affinity fragments. This review summarizes recent work on dynamic screening methodology, which resulted in the development of several high-affinity binders for various targets. Strengths and limitations of the published approaches are discussed and possible contributions of dynamic screening methodology to the drug discovery process are highlighted.

Introduction

Since an ever-increasing number of proteins have been discovered or postulated as potential new drug targets [1], the demand of the pharmaceutical industry for novel drug candidates and of the chemical biology field for chemical tools has been strongly stimulated. However, the actual output of research and development (R&D) has been described as insufficient in proportion to the investment made. For example, in 2006 the overall spending on biopharmaceutical R&D reached a record of $55.2 billion in North America. In the same year the US Food and Drug Administration (FDA) approved only 22 new molecular entities (NMEs) and biologics. In contrast, in 1996 some 53 NMEs were approved for clinical use, although R&D expenditure in 1996 was less than half the amount spent in 2006.¹ This discrepancy has been termed a ‘compound crisis’ and it has been postulated that it results from inefficient methods in drug discovery summarized as an ‘innovation deficit’ [2–5]. Thus, it is apparent that the pharmaceutical industry must improve its R&D productivity by using new technologies for the development of NMEs.

The concept of fragment-based drug discovery: small is beautiful

Fragment-based drug discovery concepts have been developed since the mid-1990s as an alternative to conventional combinatorial chemistry and high-throughput screening (HTS) approaches (Figure 1a). As opposed to combinatorial chemistry and HTS, which use large chemical libraries (>10⁶) requiring considerable effort, expensive library storage, quality control and data handling, fragment-based approaches require only small libraries (often <1000 members) as a starting point.

Conceptually, fragment-based drug discovery is based on the consideration that the free binding energy of a protein ligand results from the contributions of its molecular components. Therefore, small contributions from molecular fragments can add up to yield a high-affinity protein ligand (Figure 1b) [6]. First, a small molecule fragment that binds to the protein pocket of interest is identified. The starting fragment is then chemically modified to generate a binder of higher affinity, which is subsequently further optimized to a lead structure (Figure 1b). The concept has become very popular for two main reasons. First, there are much fewer fragments than drug-sized molecules. Rough estimates indicate that approximately 10⁷ fragments with up to 12 heavy atoms do exist (excluding 3- and 4-ring-containing structures) [7], whereas there are 10⁶³ possible small drug-like molecules with up to 30 heavy atoms [8]. For comparison, only approximately 10⁶ molecules have been synthesized to date [9]. Thus, initial screening of fragment libraries is expected to sample the chemical space much more efficiently than traditional approaches ever could.

The second reason is that fragment-derived lead structures have significantly higher ligand efficiency (free binding energy per non-hydrogen atom of the ligand) than molecules discovered by screening of large compound libraries. An investigation of 150 known natural and synthetic ligands revealed that the free binding energy increased in proportion to ligand size up to a maximum of 15 atoms [10,11]. The maximum average free-energy contribution per heavy atom was ~1.5 kcal/mol. For molecules larger than this, no further increase in ligand efficiency was observed [12]. These observations confirm how crucial limiting the molecular size is for the efficiency of protein ligands, thereby supporting the preference of fragment hits (<12 heavy atoms per molecule) over typical HTS hits. These results also indicate easier optimization and hit-to-lead development of fragment hits relative to that of HTS hits.

Dynamic template-assisted strategies in fragment-based drug discovery

The major challenges of fragment-based drug discovery are the identification of low-affinity fragments and efficient and biologically active linkage of the fragments identified.
Weakly binding ligands are difficult to detect. In general, four main biophysical techniques have been used for this task: NMR spectroscopy [14,15], X-ray crystallography [16,17], surface plasmon resonance [18] and isothermal titration calorimetry [19,20]. Major bottlenecks of these biophysical techniques are their high protein consumption, the need for expensive detection equipment and their limited throughput. Moreover, the methods do not solve the other challenging question in fragment-based discovery, i.e. how can two low-affinity binders be linked optimally to yield a high-affinity binder [21]? Biophysical methods do not provide information regarding the optimal bioactive combination of fragments.

To address this problem, alternative so-called dynamic and template-assisted strategies have been proposed for fragment-based drug discovery and are discussed here. All these methods use a target protein as a template for selection and/or assembly of optimal fragment combinations. All dynamic template-assisted approaches covered in this review have in common that a chemical reaction, which can be reversible or irreversible and enzymatic or non-enzymatic, is exploited for detection of the best fragment combination.

The first step in this direction was to shift chemical equilibria by introducing proteins, as realized in dynamic combinatorial chemistry (DCC) (Figure 2) [22–24], which has been conceptually extended in tethering [25–28], dynamic combinatorial resolution (DCR) [29–31], and pseudo-dynamic combinatorial chemistry (pDCC) [32–34] approaches. A second subcategory of fragment detection comprises template-assisted strategies such as target-guided synthesis (TGS), in which the chemical reaction used for detection is catalyzed by the protein [31,35–40]. Substrate activity screening (SAS) [41–43], which is discussed below, can be used for the transfer of fragment information from substrates to non-substrate protein ligands. Finally, dynamic ligation screening (DLS) [44,45] uses classical bioassays, such as fluorogenic substrate competition or fluorescence polarization [43], for fragment detection and can thus be used for HTS fragment-based drug discovery.

There are significant differences in dynamic template-assisted strategies, which largely determine their practical use in drug discovery. One major criterion is the amount of protein required for fragment detection. Some of the strategies, such as DCC [23,24], pDCC [32], and TGS [40], require near-stoichiometric or stoichiometric protein quantities, whereas more recently developed strategies, such as DCR [29], SAS [41] and DLS [44], use the protein template in only catalytic amounts. A second relevant criterion is the detection technology required. In DCC, tethering, DCR, pDCC, and TGS, fragment identification relies on NMR spectroscopy [29], chromatography [22,25,32,40,46], X-ray crystallography [47] or mass
spectrometry [25–28]. Recently, fragment detection consistent with established high-throughput methodologies that use standardized microtiter plates and detection by fluorescence, absorption, or fluorescence polarization in DLS have been developed [43]. Finally, the methods described here differ significantly in their scope. Whereas some can be used for detection of a wide range of protein binders, others are more specific for one protein or for a class of proteins. Thus, the aim of our review is to provide an orientation guide for potential users of dynamic template-assisted fragment-based methodology.

**Dynamic combinatorial chemistry (DCC)**

If a mixture of compounds formed in a reversible chemical equilibrium interacts with a protein, a shift in equilibrium towards the best binding components can be expected in accordance with the law of mass action. The first application of this principle for the discovery of protein binders was reported by Huc and Lehn in 1997 and termed DCC [22].

A DCC application was demonstrated by Hochgurtel et al. in 2002 [46,48]. They created an imine library by condensing a diamine (Figure 2b) that was derived from Tamiflu — a known inhibitor of neuraminidase, a key influenza virus enzyme — with 41 different ketones. For detection of enzyme activity, the imines formed were reduced to the corresponding amines by addition of NaCNBH3 and the resulting set of amines was analyzed by HPLC. High-affinity binders of neuraminidase were identified by comparing HPLC chromatograms for reactions containing the template with controls. However, in this particular application of the DCC concept the relative amplification of fragment combinations did not correspond to the binding affinities of the compounds identified (Figure 2b) [46,48]. For example, the compound exhibiting the greatest amplification was not a potent inhibitor, whereas amplification of the strongest inhibitor obtained from this screen was three-fold less. There are a number of possible explanations for this observation. In particular, the final reduction reaction can distort the results in two ways. Amines, which are derived from chemically more stable imines, are expected to be preferentially formed, irrespective of the protein template. Second, reduction of a strongly binding imine can yield an amine product with poorer binding properties.

Although in principle DCC should be applicable to different types of proteins or enzymes, the method is limited by the near-stoichiometric amounts of protein
required. Second, the reaction time is long, ranging from 1 day to 1 week. Third, larger libraries are difficult to screen because HPLC cannot easily be used to separate considerably larger libraries [48]. In addition, the detection of preferentially formed library members via reduction does not correlate with the affinity of the originally binder. Thus, differences in amplification and the affinity of the reduction products are likely to occur.

Tethering

Tethering constitutes a specific form of DCC and was been introduced by the Sunesis company (www.sunesis.com). Their approach exploits a reversible disulfide exchange reaction on the surface of a protein [25,49]. For disulfide ligation, native cysteine residues can be used; alternatively, these can be engineered in proximity to the site of interest [26,28] or added by a cross-linking reaction [27]. Thiol fragments that bind tightly in the screened pocket will form a stable disulfide with the cysteine residue of the protein or the thiol-containing cross-linker. The key advantage of tethering is the ability to focus on a particular region of the template. For example, in the case of caspase-3, the cross-linker, a derivative of aspartate, was bound covalently to the enzyme and the cross-linked caspase-3 was incubated with 7000 disulfides. The thiol group of the cross-linker interacted with the disulfides by exchange reaction. Disulfides that fit to caspase-3 are preferentially formed. The most stable disulfide–enzyme complex could be detected by high-resolution mass spectrometry, from which a potent caspase-3 inhibitor could be obtained (Figure 3).

Disadvantages of the tethering approach are the requirement for near-stoichiometric amounts of template, the need for a specialized disulfide library, which is not commercially available, and fragment detection by high-resolution mass spectrometry. In addition, because tethering products are template-linked disulfides, the fragments need to be converted into a chemically stable ligand, which needs to retain the affinity of the disulfide, before further information can be obtained. Nevertheless, the tethering approach enables site-directed screening via cross-linking and, in contrast to DCR, pDCC and SAS, is broadly applicable to every type of protein and enzyme.

Dynamic combinatorial resolution (DCR) and pseudo-dynamic combinatorial chemistry (pDCC)

Both DCR and pDCC rely on the concept of DCC. A dynamic combinatorial library \( A_n \ast B_m \) is generated and incubated with an enzyme that catalyzes an irreversible reaction of the library members \( A_n \ast B_m \) to \( C_{nm} \) with a certain degree of selectivity (Figure 4a). The selection process can be performed using enzymes that either catalyze bond formation (in DCR) (Figure 4b) or cleavage (in pDCC) (Figure 4c).

Inspired by dynamic resolution, a method for separation of a racemic mixture [41], Ramström’s group introduced
the concept of DCR [29–31]. The proof of principle for this approach was provided by a nitroaldol reaction (the so-called Henry reaction), in which β-nitro alcohols were reversibly formed by addition of nitroalkanes to carbonyl compounds. On addition of the selector, in this case the lipase PS-C I from Pseudomonas cepacia, library members that could fit into the active site of the lipase were acylated (Figure 4b) and subsequently identified by $^1$H-NMR [29].
pDCC was developed by the Kazlauskas group [32,34]. They replaced the equilibrium used for selection in DCC by combining an irreversible reaction used for library formation with enzymatic cleavage of library members. Here, the selection protein template protects better-binding fragment combinations from enzymatic cleavage, whereas unbound compounds are rapidly removed from the mixture. The concept was successfully demonstrated for a carbonic anhydrase template using a synthetic dipeptide library (Figure 4c). After incubation with the protein template, the non-specific protease pronase from Streptomyces griseus was added to hydrolyze all unbound dipeptides. As expected, the cleavage reaction was strongly inhibited for the best binder compared to weaker binders, allowing detection by NMR.

In both approaches, amplification correlated directly with the binding affinity of the compounds identified. However, as above, one major disadvantage is the difficulty in detecting the best fragment combinations by NMR, which limits the possible library size. Another limitation is the restricted applicability, which is confined to chemical libraries for which a cleavage reaction can be established. For example, in the case of DCR, the choice of selector templates is confined to relatively few enzymes such as lipases, preventing broader applicability of this particular approach for drug discovery. However, only catalytic amounts of the selector protein are required, which constitutes a clear advantage over DCC. Similar to DCR, pDCC is also confined to specific chemicals. Dipeptides, as demonstrated here, are ideal for establishing the concept; however, they are not very likely drug leads. Thus, in summary these approaches, although conceptually interesting, are less likely to be widely used in drug discovery.

**Target-guided synthesis (TGS)**

A logical and straightforward extension of template-assisted shifts in chemical equilibria is the direct use of protein targets as either templates or catalysts in irreversible chemical reactions, a concept termed TGS [31,50]. In TGS, the protein template binds two reagents that are in close proximity, thus accelerating their irreversible chemical reactions, a concept termed TGS [31,50]. In TGS, the protein template binds two reagents that are in close proximity, thus accelerating their irreversible chemical reactions, a concept termed TGS.

For example, in the case of the wild-type HIV1 protease (IC50 6 nM)[40]. Because of the growing importance of triazoles in drug discovery, the most widely used reaction for TGS is the so-called click chemistry [39,40,51]. Click reactions of azides with alkynes regioselectively yield 1,4-disubstituted triazoles by copper(I) catalysis [52]. In the presence of a suitable protein template, however, a well-bound azide and alkyn can react in the absence of the copper salt. For example, in the case of HIV protease, an alkyn (IC50 >100 µM) and an azide (IC50 4.2 µM) were incubated in the presence of the enzyme for 24 h. HPLC analysis revealed that the triazole was formed, which is an inhibitor of the wild-type HIV1 protease (IC50 6 nM) [40]. Because of the growing importance of triazoles in drug discovery [52], in situ click chemistry could become a powerful tool in lead discovery.

In principle, TGS is suitable for the development of highly active molecules. For broader applicability, however, some issues have to be addressed. For template-assisted reaction and detection of the designed products by HPLC, NMR, or mass spectrometry, they need to be present in relatively high concentrations (≥100 µM). Second, the need for a near-stoichiometric amounts of the template protein limits the feasibility of TGS. We anticipate that the set of reactions for which TGS is particularly suited will not be fully explored until further research has been carried out.

**Substrate activity screening (SAS)**

SAS was introduced by the Ellman group and uses substrate libraries that are screened in standard enzyme assays to determine the best binding fragments [41]. Initially, this method was demonstrated for proteases. In this particular application, a fluorophore was coupled to a variety of low-molecular-weight fragments, resulting in a so-called fluorogenic substrate library of ~100 members that was screened using several proteases (such as cathepsin S shown in Figure 5a) to identify the best protease substrates via cleavage of the amide bond, which released the fluorescent dye. Here, high substrate turnover indicated high affinity of the respective substrate fragments. In the case of cathepsin S, a non-peptidic substrate could be identified. The best substrate identified in this reaction was then converted into an inhibitor by introducing an aldehyde at the C-terminus, providing a non-peptidic nanomolar inhibitor of cathepsin S [42].

SAS was also used to identify protein tyrosine phosphatase inhibitors (Figure 5b). A substrate library of O-aryl phosphates was prepared and screened with Mycobacterium tuberculosis protein tyrosine phosphatase B (MptpB), a secreted virulence factor and potential target against tuberculosis (TB). A biphenyl scaffold could be identified as a potential lead fragment. The phosphate group in the molecule was then replaced by a phosphate mimetic and, following optimization, a potent and selective MptpB inhibitor was obtained (Figure 5b) [53].

Several molecules acting on therapeutically relevant enzymes have already been identified by SAS [42,53]. The strength of this method is its ability to efficiently identify substrate structures that bind to the active site. The use of protein in only catalytic amounts contributes to its efficiency. However, SAS requires the catalytic function of enzymatic drug targets, such as proteases or protein tyrosine phosphatases, and therefore cannot be applied more generally to other types of drug targets. The approach is further restricted by the limited availability of substrate libraries, which are not commercially available and have to be synthesized on a case-by-case basis. Moreover, the classical rationale of fragment-based drug discovery, i.e. to connect several low-affinity binders for specific pockets, is not applied in SAS. Another drawback is that the substrates discovered have to be converted into useful inhibitors. For example, in the case of cathepsin S, the substrate fluorophore was replaced with an aliphatic aldehyde, which is less suitable in a drug. Similarly, for phosphatase inhibitors, replacement of the phosphate by a mimetic such as isothiazolidine or isoxazole carboxylic acid [53] resulted in adverse effects, such as lower selectivity and cell permeability.
Dynamic ligation screening (DLS)

Considering the limitations of different dynamic fragment-based methods for drug discovery, DLS was developed as an approach that combines dynamic target-assisted formation of inhibitory species with detection via a biochemical assay [44]. For example, an enzyme reaction can be used for amplified detection of the fragment, thus drastically reducing the amount of protein required and allowing HTS (Figure 6). In addition, the use of chemically reactive protein ligands that bind to a defined pocket on the protein surface makes it possible to test for inhibitors fragments acting at the pocket of interest.

The DLS approach was first demonstrated for the main protease of SARS coronavirus (SARS-CoV Mpro) as the protein target. For site-directed identification of inhibitory fragments, an activity assay for SARS-CoV Mpro was first developed using a peptidic substrate (Figure 6). Enzymatic cleavage of the peptidic substrate released a fluorophore, which served as an indicator for protease activity. The chemically reactive protein ligand, a peptide aldehyde [45,54], was then incubated with an excess of a nucleophilic fragment in the presence of the enzyme. Following addition of the fluorogenic substrate, rate differences in substrate turnover were quantified to identify active inhibitory fragments. For SARS-CoV Mpro, one fragment was identified that showed considerably stronger inhibition in presence of the peptide aldehyde than the peptide aldehyde inhibitor alone. The fragment alone showed no significant activity against the SARS enzyme. To verify the result, the fragment identified was converted by chemical synthesis into an electrophile, which was shown to be an active inhibitor of SARS-CoV Mpro and was used as a new chemically reactive protein ligand. Thus, the DLS approach was repeated with the newly identified electrophile to identify a new nucleophile fragment. Covalent linkage of the new electrophile to the fragment identified yielded an active,
Figure 6. Concept of dynamic ligation screening (DLS) illustrated by the development of a non-peptidic SARS-CoV Mpro inhibitor. The substrate 1 (shown in green) competes with a peptide aldehyde inhibitor 2 (orange) targeting the S1 pocket of the SARS-CoV main protease (gray). An active amine fragment 3 (red) binds to the S1’ pocket, leading to increased inhibition via dynamic ligation in the active protein site [44]. Based on the active amine fragment 3 (red), an analogous electrophilic aldehyde fragment 4 (red) that also binds to S1’ was synthesized and used for DLS of the S1 pocket, yielding active fragment 5 as a hit. Finally, both hit fragments were chemically linked to yield the non-peptide inhibitor 6 with a $K_I$ value of 2.9 $\mu$M [44].
non-peptidic SARS-CoV M<sup>pro</sup> inhibitor with a <i>K<sub>I</sub></i> value of 2.9 µM (Figure 6).

In conclusion, DLS provides site-directed detection of low-affinity fragments. The sensitivity of the method is therefore higher than for conventional and other dynamic fragment-based approaches. The method can be applied in a high-throughput format. No additional equipment besides a standard microtiter plate reader is needed. Most importantly, DLS can be operated iteratively in an evolutionary process and has been shown to succeed in transforming a moderately active peptidic inhibitor into an entirely non-peptidic inhibitor [44]. Although successful DLS has only been demonstrated for the development of protease inhibitors to date [43,44], DLS assays could easily be extended to other proteases and to other enzyme classes, as well as to any type of protein–protein interaction, so that standard biochemical assays, including binding assays, can easily be adapted to the particular strategy required [43].

**Outlook**

Methodological innovations in the drug discovery process are required to alleviate the dwindling success of current drug development efforts in the pharmaceutical industry. Recent advances in dynamic fragment-based screening methods have the potential to contribute significantly to this process because they combine the power of high-throughput screening with the advantages of fragment-derived hits. With the development of dynamic ligation screening approaches, fragment-based drug discovery has become more competitive with traditional methods because only catalytic amounts of the target protein are required and standard HTS methods can be adapted. Considering recent breakthroughs in the field, dynamic template-assisted strategies are now facing the next challenge: it must be demonstrated that they can significantly support the development of an approved drug. It is our firm belief that successful application of these approaches will yield positive results in the near future.

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