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Improving success rates for lead generation using affinity binding technologies
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Affinity technologies have been applied at several stages of the drug discovery process, ranging from target identification and purification to the identification of preclinical candidates. The detection of ligand–macromolecule interactions in lead discovery is the best studied and most powerful of these techniques. Although affinity methods have been in widespread use for about a decade, only recently have many reports emerged on their utility. Primary affinity screens of large libraries of small molecules or fragments have begun to produce results for challenging targets. Furthermore, in secondary assays affinity methods are opening new avenues to tackle important medicinal chemistry tasks.

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Introduction
Lead generation is a critical first step in the drug discovery process. Over the past decade, high-throughput screening (HTS) of corporate compound decks has emerged as the primary paradigm for hit or lead discovery. Typically, targets are interrogated with 1–3 million discrete compounds in parallel. The HTS era has thus far delivered mixed results. Hit rates for targets tend to be extremely low or very high. Two key factors are likely to be responsible for this. First, most compound decks in the industry are skewed in compound distribution favoring specific, well-studied protein subfamilies. Second, and most importantly for novel targets, sampling of drug-like chemistry space by a few million compounds is inadequate to consistently deliver several diverse lead classes [1]. Challenges facing HTS technologies include high false-positive and false-negative rates, the need for reporter assays, and the limitation in throughput imposed by testing compounds individually [2]. To address these issues, affinity or bio-affinity screening approaches have emerged as orthogonal methods for early lead discovery. For affinity-based techniques, the readout is typically a qualitative or quantitative signal based on the physical interaction between macromolecule (RNA, DNA or protein) and the small-molecule partner. This approach has several advantages over biochemical assays for both drug-like and fragment-like methods (see Glossary). First, it has the capability to sample mixtures (including natural products) rather than discrete entities, thus enabling the exploration of larger chemical spaces without a concomitant increase in the number of samples. Second, it has the ability to investigate multiple forms of macromolecules, such as various ternary complexes, activated or inactivated forms and forms with or without cofactor. Third, it is possible to carry out the assays at small-molecule concentrations much higher or lower than that of the corresponding biochemical assays. Last, this approach eliminates many assay artifacts that arise from nonspecific aggregation, fluorescence absorption or quenching. On the downside, affinity-based technologies do not deliver a functional readout. Thus, affinity hits require biochemical, cellular or in vivo validation. Nevertheless, the advantages of affinity techniques have extended their use into follow-up or secondary screens, especially in the field of ATP-dependent enzymes.

In this article, we review a subset of affinity techniques that has been shown to be valuable for lead discovery (Table 1). We provide several examples of demonstrated success, considering achievements from the perspectives of the technology and drug discovery.

Affinity selection in screens of drug-like libraries
Affinity selection techniques can be classified as homogeneous or heterogeneous. A heterogeneous screening environment is created by immobilization of either the macromolecule of interest or of the small molecule(s) on a reactive surface followed by a static or flow-based analysis of binding. By contrast, a homogeneous environment leaves both macromolecular and small-molecule species to interact in their native states. Following the binding event, a qualitative or quantitative assessment of the interaction can be made using one of many available detectors that have been used for affinity screening, such as nuclear magnetic resonance (NMR), mass spectrometry (MS), X-ray crystallography or surface plasmon resonance (SPR). A subset of these techniques employs a filtration step before detection. Specific details of some affinity
Fragment-based drug or lead discovery: A lead generation technique that calls for primary screens of weak-binding (IC\textsubscript{50} 5 mM–50 μM) fragment-like molecules followed by optimization towards leads, rather than for primary screens of more potent (IC\textsubscript{50} 25 μM–50 nM) drug-like molecules via HTS. The advantage of the former is the better sampling of chemistry space, because the number of compounds relevant for drug discovery grows exponentially with molecular weight.

Fragment evolution: A fragment optimization technique that typically involves the systematic exploration of structure-activity relationships along various substitution vectors of the fragment hits.

Site-directed fragment screening: A fragment optimization technique that typically involves screening fragment libraries in the presence of another fragment added at saturation concentrations. The binding mode of the latter fragment is typically known from previous studies. Any new hits from the second screen would target new binding sites and such hits could later be the subject of ‘fragment linking’ with the previously identified fragment.

**Table 1**

| Affinity-based technologies in primary screening applications. |
|-------------------|-----------------|-----------------|-----------------|-----------------|
| Detection     | Kinetics\(^a\) | Binding assay   | Main applications                                      | Pros                                                                                     | Cons                                                                                   |
| MS            |                | Homogeneous     | Ultra-HTS of large libraries                           | Improved sampling of drug-like chemistry space (higher throughput and ability to analyze mixtures), preference for slower K\textsubscript{off} | K\textsubscript{off}-dependent signal                                                   |
|               | K\textsubscript{d}-dependent | Homogeneous reversible | Ultra-HTS of large libraries                           | Improved sampling of drug-like chemistry space (higher throughput and ability to analyze mixtures), suitable for weak compounds | Harder to automate and miniaturize                                                     |
|               |                | Homogeneous, covalent | Fragment-based lead discovery, Ultra-HTS of large libraries, mixture capability | Smaller space to sample, lead-likeness                                                    | Mutated proteins might be required                                                     |
|               |                | Heterogeneous: immobilized protein | Fragment-based lead discovery, Ultra-HTS of large libraries, mixture capability | Improved sampling of drug-like chemistry space, enabling natural product screens         | Immobilization of target                                                              |
| NMR           | K\textsubscript{d}-dependent | Homogeneous     | Fragment-based lead discovery                           | Smaller space to sample, lead-likeness                                                    | Target size limits, soluble proteins only, lower throughput, labeled protein may be required |
|               |                | Heterogeneous: immobilized protein | Fragment-based lead discovery                           | Smaller space to sample, enables membrane proteins to be used, lead-likeness              | Target size limits, labeled protein required                                          |
| X-ray         |                | Fragment-based lead discovery | Fragment-based lead discovery                           | Smaller space to sample, lead-likeness                                                    | Long validation for new proteins, difficulties associated with crystallization      |
| PCR           | K\textsubscript{d}-dependent | Heterogeneous: immobilized protein | Ultra-HTS of vast DNA-templated libraries              | Improved sampling of drug-like chemistry space (higher throughput and ability to analyze mixtures) | Limited chemistry scope, immobilization of target                                      |
| SPR           | K\textsubscript{off}-dependent | Heterogeneous: immobilized compounds | Fragment-based lead discovery and traditional libraries | Applicable for both fragment-like and drug-like compounds                                 | Immobilization of compounds                                                           |

\(^a\) K\textsubscript{off}, off-rate (dissociation rate constant); K\textsubscript{d}, dissociation constant.

The automated ligand identification system (ALIS) screening platform (NeoGenesis; http://www.neogenesis.com/) was one of the earliest affinity selection technologies capable of screening very large numbers of compounds. The key step to ALIS is the separation of the desired protein–ligand complex from non-binding components and dissociation of the complex by reverse phase chromatography with mass detection. As part of a lead discovery effort against the anti-infective target *Escherichia coli* dihydrofolate reductase, a total of 3 750 000 compounds (in mixtures of 2000), were screened to identify compound 1 (Figure 1) [4]. The advantages of improved sampling of chemistry space were demonstrated by the discovery of a single compound (2) from a multimillion member library for β-secretase (BACE1), an enzyme that is crucial in Alzheimer’s disease [5\textsuperscript{*}]. The compound was selective, possessed a unique binding mode and could be optimized to form a potent inhibitor (3) of the enzyme [6]. The generality of the ALIS platform has been demonstrated against a diverse set of targets [7].
A related technique, termed Speedscreen (Novartis; http://www.novartis.com/), has also been disclosed. In practice, this approach includes the same sequence of events as ALIS, but each step is carried out completely decoupled from the last. Unlike ALIS, Speedscreen takes advantage of the parallel capabilities of using a 96-well format [8]. A typical screening campaign is comprised of approximately 600 000 compounds in mixtures of 400 tested against soluble and globular proteins [9].

Researchers at Abbott (http://www.abbott.com/) have used ultrafiltration to separate binders for the oncology target Bcl-XL from non-binders, in mixtures of 2400 compounds, as part of a study comparing results from a fluorescence polarization assay and an affinity selection/mass spectrometry (ASMS) platform. From the 263 382 compounds that were screened in duplicate, 29 binders with affinities below 100 μM were identified by ASMS [10]. The authors suggested that it is valuable to apply different screening methods for lowering the false-negative rate. Similar conclusions can be made from the screening statistics for far upstream element binding protein (FBP) [11].

Capillary electrophoresis (CE) has also been shown to be a reliable tool for studying molecular interactions [12,13]. CE has been used to interrogate crude natural product broths for potential leads [14]. A binding event can be discerned by monitoring the migration times of the protein–ligand complex versus that of the protein alone as a reference point. Upon complex formation conformational changes of the protein expose different residues to the surface, an event that can change the mobility of the complex in relation to its charged environment.

Frontal affinity chromatography coupled to a mass spectrometer (FAC-MS) is based on the immobilization of a target onto a solid support [15,16]. Small molecules are infused continuously over the immobilized target and are monitored by a mass spectrometer. Depending on their affinity strength, the extracted ion currents for the binders are observed at later elution times when compared to a non-interacting control compound. Furthermore, the shape of the individual extracted ion currents can provide crude approximations of relative affinities under competitive conditions. In a study to discover molecules that bind to the surface spike protein of severe acute respiratory syndrome coronavirus (SARS-CoV), two small molecules have been reported. Compounds 4 and 5 were identified from the extracts of more than 121 Chinese herbs using the FAC-MS approach [17].

By using the resolving power of Fourier transform ion cyclotron resonance mass spectrometry, researchers have
been able to study non-covalent interactions of RNA–
ligand complexes without the need for a separation step
before detection [18,19]. The platform, termed multi-
target affinity/specificity screening, allows for the multi-
plexing of RNA constructs and small molecules; however,
the generality of this technique for other targets remains
to be demonstrated.

As part of the quest to screen increasing numbers of small
molecules, DNA-tagged libraries have been devised and
affinity screened against immobilized targets. Binders are
then amplified by PCR to reveal the chemical identity.
This technology is rather new and its limitations (syn-
thetic, false-positive or false-negative rates) are not yet
clear (more details can be found at http://www.nuevolu-
tion.com or http://www.praecis.com).

Affinity detection of fragments for lead
discovery

As the cost, efficiency, scalability and lead delivery-rate
issues with HTS became apparent over the past decade,
new terms such as lead-likeness, fragment-likeness and
fragment-based drug discovery have surfaced (see Glossary).
Significant resources and research have been dedi-
cated to identifying techniques that can measure the
binding of very small (molecular weight \(\sim 120–300\) Da)
molecules (fragments) to macromolecules. The inherent
difficulties of using biochemical tools to detect weak
ligands, often at millimolar concentrations, necessitated
the search for affinity-type readouts. Over the years,
NMR (Abbott, Vertex; http://www.vpharm.com/) and
X-ray crystallography (Abbott, Astex http://www.astex-
therapeutics.com/ and Plexxikon http://www.plexxikon.-
com/) have emerged as the methods of choice to conduct
fragment-based lead discovery. An intriguing alternative
for the covalent trapping of fragments in a \(K_d\)-dependent
manner has also been developed by Sunesis (thiol-reactive
probes; http://www.sunesis.com/).

The fragment-based approach is ideal for finding new
‘warheads’, small functional groups or ensembles of func-
tional groups that bind to crucial residues conserved
within protein families. These advantages have been
demonstrated with protein tyrosine phosphatase 1B
(PTP1B), a validated but historically challenging target
implicated in type II diabetes. A protein-based NMR
fragment screen (which detects changes in the NMR
spectra of a protein on ligand binding) has been used to
identify several phosphate mimics: oxalylarylamino-
benzoic acids (6; Figure 2) [20], which proved not to be
cell permeable, and isoxazolcarboxylic acids (7) [21*].
The latter was linked with a noncompetitive fragment
identified by another NMR screen, which resulted in the
development of a monocharged PTP1B inhibitor (8) with
cellular activity. In a search for aminoglycoside mimetics
against the \(E.\ coi\) A-site RNA, an NMR-based screen of a
10 000 member library was used to identify several dif-
ferent chemical series, two of which (2-aminoquinolines
and aminopyridines) could rapidly be advanced to more
potent derivatives in the low micromolar range [22].
Structure activity relationships determined by NMR
(SAR by NMR) have been instrumental in the develop-
ment of Abbott’s matrix metalloproteinase program, iden-
tifying the key biphenyl group [23]. Fragment linking
with hydroxamic acids followed by medicinal chemistry
optimization ultimately resulted in the development
candidate ABT-518 (9) for treating cancer. Site-directed
fragment screening and fragment evolution (see Glossary)
was also used to discover and optimize leads for the anti-
apoptotic Bcl-2 family, ultimately leading to ABT-737
(10). Compound 10 was well tolerated and was shown to
cause tumor regression in multiple cancer cell lines either
as a single agent or in combination with chemotherapeut-
tics [24]. Protein-based NMR fragment screens have also
recently been extended to insoluble proteins [25].

Unlike protein-based NMR, the SHAPEs technique
relies on the detection of changes in NMR spectra of
the small molecules themselves. Advantages over the
protein-based method are that no limits are imposed
on the size of the protein, there is no requirement for
labeled protein, and lower reagent consumption. On the
downside, no structural information on the protein’s
binding site or on the relative binding mode of different
fragments can be discerned. The SHAPEs method has
been applied pre-HTS, owing to minimal development
time, or post-HTS for failed targets, as exemplified by
Jnk3 MAP kinase and adipocyte lipid-binding protein
[26]. Close integration with X-ray crystallography and
molecular modeling was reported to be essential in opti-
mization of the initial fragment hits.

Over the past decade, data analysis, crystal soaking tech-
nology and throughput have matured to the level where
X-ray crystallography can be applied to the parallel detec-
tion of weakly binding fragments. Although drawbacks
such as false-negatives [27], long development times for
novel unknown proteins, and protein consumption still
remain, structure-guided fragment evolution makes this
technology an attractive option. Fragments can simply be
screened in small mixtures (<10 members) or can be pre-
filtered by biochemical assays to limit the set of scaffolds
for crystallization. A proof-of-principle for the latter
approach has recently been disclosed for phosphodies-
terases. In that study, 1-phenyl-4-pyrazolecarboxylic
esters (11) were developed using a 20 000 member scaf-
fold library and structure-based optimization [28]. A more
comprehensive study of five targets (p38, CDK2, throm-
bin, PTP1B and RNase A) demonstrated that weak
fragments can indeed be identified using high-through-
put crystallography [29*]. Chemical series evolution from
fragments (12) to leads (13) was illustrated by the p38
program [30]. Although these studies clearly show the
potential of the underlying technology in lead generation,
the strong reliance on structure-based design for optimization makes the discovery of truly novel molecular frameworks unlikely.

Using thiol-reactive probes, researchers at Sunesis were able to screen for new chemical motifs that were active against PTP1B with the aid of ‘breakaway tethering’ [31*]. A cysteine residue was introduced outside of the enzyme active site and covalently modified with a flexible linker holding a thiol end group. The protein was then interrogated with 15,000 fragments and the screen unveiled the 1,2-pyrazinedicarboxylic acid monoamide moiety (14), which was an equipotent and competitive inhibitor of phosphotyrosine in biochemical assays. A different study in which electrophiles were screened against PTP1B found that 15 selectively reacts with a cysteine 8 Å away from the catalytic site [32*]. This covalent modification reduces the enzymatic activity of
PTP1B via an allosteric perturbation of the neighboring phosphotyrosine-binding site.

Competitive affinity screens for cytokine hormone interleukin-2 (IL-2) using fragments in the presence of a previously known inhibitor revealed a series of compounds that show cooperative binding with a previously known molecule [33]. Fragment decomposition of the same known inhibitor and additional competitive fragment screens were invaluable for the development of nanomolar inhibitors with cellular activity [34]. Fragment libraries with exchangeable thiols unveiled an allosteric binding site for caspases [35] as well as agonist and antagonist fragments for the G-protein-coupled receptor C5a [36]. These results elegantly demonstrate one of the key advantages of affinity-related techniques: the ability to study or screen different enzyme forms and to find compounds with affinity for allosteric or adjacent binding sites.

Fragment-based lead discovery has been shown to be successful over the years, delivering ‘efficient’ leads (relatively high free-energy of binding per heavy atom [37**]). As the approach is relatively new, it remains to be seen whether the smaller and more hydrophilic lead entry points for medicinal chemistry actually yield smaller drug candidates that fail less often in the clinic [38].

Surface plasmon resonance in lead generation

The focus of the affinity techniques discussed so far has been either to increase the throughput for traditional libraries or to enable the detection of weakly binding fragments with low throughput. Both help sampling of discovery space but represent opposite extremes. The immobilization of small organic molecules on microarrays enables the use of SPR to detect weak interactions. The method is practically an extension of the established BIacore™ technology to fragments, and may hold promise for HTS of lead-like small molecules. A model study using factor VIIa and a focused set of immobilized compounds confirmed binding of the positive control (\(\beta\)-aminoaztreonamidine, \(\sim 70 \mu M\)) and identified several new fragment ligands [39]. Follow-up biochemical activities or the tendency to form crystals, however, had no correlation with the observed SPR signal. This developing approach may mature into a semi-quantitative screening technique or a qualitative filter.

Secondary assay applications

High-throughput affinity methods other than NMR or X-ray crystallography have found their way into many applications. These include mechanistic and competition studies, the determination of binding kinetics, and compound profiling or selectivity screens.

The use of homogenous affinity selection has recently been shown to distinguish direct or allosteric competitive behavior. It could also establish compound ranking in mixture format through titration experiments of unknown compounds with a ligand [40]. The same method can also be used for ranking new derivatives in optimization studies. SPR analysis of binding kinetics with BIacore™ has been the standard for the determination of association and dissociation rate constants and \(K_d\)s [41,42]. The major drawback — the requirement for immobilizing one of the binding partners, typically the protein — is compensated for by the lack of need for radiolabeled materials.

Selectivity is a particularly crucial factor in the development of drug candidates for ATP-dependent enzymes. There has been a significant effort to express and develop assays for a growing number of kinases and their mutants for counter-screening studies, owing to a few early reports on finding off-target liabilities (potential adverse effects) in distant sections of the kinase dendogram. Immobilized inhibitors can be used to map ATP-binding site interactions with many targets. Conceivably, each inhibitor might have to be linked via multiple orientations to avoid false-negatives arising from linker interference. Nevertheless, several surprisingly tight off-target interactions were identified for the ary1-imidazole and pyrido[2,3-d]pyrimidine kinase inhibitor classes [43–45]. A semihomogenous variation of this technique has also been developed, which requires the immobilization of only a small number of compounds to study all ATP-competitive small molecules with any enzymes [46**]. In this approach, highly promiscuous (inhibitory constant (IC\(_{50}\) < 1 \(\mu M\)) ‘bait’ molecules are immobilized on the solid support via a biotinylated linker. The T7 phage tagged protein is introduced along with the ‘study compound’ free in solution. If the free compound does not bind to the protein, elution of the macromolecule is slower because it can extensively interact with the bait compounds on the solid support. The amount of protein bound to the support is quantified by either phage plaque assays or quantitative PCR for high sensitivity and can be plotted as a function of the test compound concentration. An entire enzyme panel can be run in a single experiment and, unlike biochemical assays, the result is not dependent on ATP concentration. Of course, when interesting novel interactions of known compounds with new targets are found, these profiling techniques can also serve as lead discovery tools to initiate chemistry efforts to derive new therapeutics.

Conclusions

Today’s high-throughput screens, although frequently effective, might not be sufficient for the pharmaceutical industry’s future lead discovery needs. Affinity methods aim at easing this problem in various ways and new advances in these technologies will surely follow. Better success rates will not only improve the lead discovery process, but will also enhance our understanding of small-molecule binding to proteins. To this end, a protein
‘druggability’ index has been proposed based on NMR fragment screening data [47]. The predictive classification of proteins via small-molecule interactions is a highly intriguing concept, but further theoretical and experimental studies on the required sampling rate of the different discovery libraries (drug-like or fragments) might be necessary before this major milestone can be reached.

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