Building a Pharmacological Lexicon: Small Molecule Discovery in Academia

John S. Lazo, Linda S. Brady and Ray Dingledine

Department of Pharmacology, Pittsburgh Molecular Library Screening Center, University of Pittsburgh Drug Discovery Institute, University of Pittsburgh, Pittsburgh, PA 15261 (JSL);

Project Team Leader, Molecular Screening Centers Network, Division of Neuroscience and Basic Behavioral Science, National Institute of Mental Health, 6001 Executive Blvd, Room 7204, Bethesda MD 20892-9641 (LSB);

Department of Pharmacology, Emory Chemical-Biology Discovery Center, Emory University, Atlanta, GA 30322 (RD)
Running title: Small Molecule Library Screening

Corresponding authors: John S. Lazo; Department of Pharmacology, Pittsburgh Molecular Library Screening Center, University of Pittsburgh, Biomedical Science Tower 3, Suite 10040, Pittsburgh, PA 15261-0001; Fax: 412-648-2229; Email: lazo@pitt.edu or Raymond Dingledine, Department of Pharmacology, Emory Chemical-Biology Center, 1510 Clifton Rd, Emory University, Atlanta, GA 30322; Email: rdingledine@pharm.emory.edu

22 text pages
2 tables
6 figures
14 references
86 words in abstract
4217 words in body of text (introduction, results, discussion combined)

Abbreviations: DMSO, dimethyl sulfoxide; HCS, high content screening; HTS, high throughput screening; MLSCN, Molecular Library Screening Centers Network; OED, Oxford English Dictionary; TR-FRET, time-resolved fluorescence resonance energy transfer; GPCR, G protein–coupled receptor; MLSMR, Molecular Libraries Small Molecule Repository; SAR, structure-activity relationship; CMLD, Chemical Methodologies and Library Development.
ABSTRACT

Small molecules are powerful pharmacological tools to dissect biological events. There is now considerable interest in expanding efforts to identify and use small molecules targeting proteins encoded in the genomes of humans and pathogenic organisms. Integration of the principles of molecular pharmacology with contemporary high throughput and high content screening technologies is essential for the success of these discovery activities. We present some of the challenges and opportunities provided by the Molecular Library Screening Centers Network (MLSCN), which is a National Institutes of Health Roadmap Initiative.
“Four hundred years ago there was not such convenience available on any English bookshelf.”

— Simon Winchester, *Professor and the Madman.*

Can you image writing an article or a book without a dictionary or a thesaurus? How would you find the definition of exon or intron; prion or ion? That is what William Shakespeare faced when he wrote “*In a Midsummer Night’s Dream*”. The first English dictionaries appeared about the time of his death and were arranged by subject not alphabetically. The English language was spoken and written but it was not defined. Perhaps pharmacologists a few decades from now will have similar thoughts when they consider the status of the Pharmacology and Chemical Biology of today. We take for granted that small molecules have biological effects but we lack a readily accessible annotated database of the pharmacological effects of chemicals. One attempt to remedy this is an effort by the National Institutes of Health’s Roadmap Initiative. The purpose of this article is to explore the potential and describe recent activities of the Roadmap’s Molecular Library Screening Center Network (MLSCN) (www.mli.nih.gov/mlscn/).

The power of readily available lexicons is evident even to a nonscientist. Consider the venerable Oxford English Dictionary (OED). Some would argue that the current hegemony of the English language rests largely on the development of the OED, which was a radical experiment sponsored by the Philological Society of London. On April 26, 1878, the Philological Society invited Professor James Murray to edit what would become the OED. The uniqueness of the project was effectively captured in the book “Professor and the Madman” written by Simon Winchester (Winchester 1999) and it was partially the inspiration for this article. Professor Murray needed to convince English-speaking people to scan the literature for the first use of words. The thought of
asking tens of thousands of individuals to voluntarily comb through their book collections and local libraries and, without any compensation, donate their findings to an Oxford Professor might seem futile or even mad. Nonetheless, starting with A and proceeding over several decades they defined 414,825 words and illustrated them with 1,827,306 textual examples, finally fixing the magnitude of the English language. The madman in this story, by the way, was the prime OED contributor, Dr. William Chester Minor, an American Civil War captain and Yale trained physician, who was also an inmate at an asylum for the criminally insane. The OED model of public contribution to build language lexicons lives on in the current Wikipedia experiment (http://en.wikipedia.org).

We also have two powerful examples of the consequences of freely available lexicons in biology: PubMed and the Human Genome Project. It is difficult to remember the days before Web-based searches of hundreds of genomes with publicly available free software programs. Nevertheless, the initiation of the Human Genome Project was surrounded with controversy (Palca 1989; Roberts 1989; Roberts 1990) that is reminiscent of the contemporary public debate over funding precipitated by the new NIH Roadmap Initiative (Bravo 2006; Marks 2006; Weissmann 2006). Public debate about the allocation of NIH funds would seem to be an inevitable and productive process associated with public funding mechanisms and has been addressed elsewhere (Lazo 2006). In this review, we explore how small molecule screening and the MLSCN could affect the future of Pharmacology.

Should I use chemistry or genetics to probe protein function? The field of pharmacology, now more than 100 years old, has always been at the interface between chemistry and biology. The earliest pharmacologists relied on crude organ preparations or animal models to probe how recently isolated or extracted compounds like cocaine, morphine or atropine functioned. The advent of more
reductionist approaches revealed molecular, even atomic, interactions of small molecules with protein targets. The Human Genome Project resulted in the discovery of thousands of new genes encoding tens of thousands of proteins. Now many are attempting to integrate the knowledge obtained from molecular biological and genetic studies in the context of the whole organism. How can we systematically and relatively quickly develop an understanding of the biological functions of all these new proteins? Sophisticated gene knockout technologies can produce highly specific deletion of individual proteins, although homeostatic compensation must always be borne in mind. Moreover, knockout technologies are currently restricted to just a few species and are often irreversible, leading to a hypomorph lacking half or all of the gene product. Like silencing RNA strategies, gene knockout usually eliminates the entire protein, which reduces the value of these approaches for studying splice variants or the functions of multiple independent domains within an individual protein.

As an alternative to the genetics approach, small molecules can provide powerful tools to dissect biological events. They allow one to reversibly affect protein function in a graded rather than all-or-none fashion, modify the subcellular location of a macromolecule, or disrupt a specific protein-protein interaction. The motto of small molecule users is “any species, any place (in the body), any time (during development)”. Moreover, small molecules can be designed that inhibit or potentiate protein function, and can be used to probe the function of individual subunits in a multimeric protein complex, or even different domains of the same protein subunit. Finally, a selective, potent small molecule targeted to a disease-related protein can serve as a lead for therapeutic development.

However, achieving sufficient target selectivity is often very difficult, making the small molecule approach complementary to genetics rather than a substitute.
The recognition that small molecules can be powerful reagents has fueled enthusiasm for using high throughput screening (HTS) and high content screening (HCS) methods to identify biologically friendly small molecules that would be readily available to scientists. Evidence for the power of such endeavors are found throughout the contemporary pharmacological literature (Oltersdorf et al. 2005; Huang et al. 2006; Sanna et al. 2006).

**Differentiating Academic and Industry Goals.** The phrase “compound screening” sometimes evokes a disapproving remark in the halls of academia even though other screening activities such as screening a DNA library for expression differences or novel genes are well accepted and occasionally even admired. This might reflect a belief that compound screening is something that should be relegated to an industrial environment, being an applied task that is not hypothesis driven. That small molecule screening is reagent building cannot be denied but there are many chemical biology screens that provide valuable probes and yield unique biological insights. Not to seek and use these reagents seems foolish. Indeed, innovative and high impact advances in therapeutics will likely come from aggressive efforts to provide a bridge that allows translation of advances in basic science to novel therapeutics and marketable products. This bridge is the identification of novel small molecules and their mechanisms of action that specifically perturb the function of disease-related proteins studied in academic laboratories. Three developments in the past five years have made the small molecule-based chemical genetics approach a realistic goal in academic institutions: the development of commercially available small molecule libraries, a reduction in cost of screening instrumentation, and the beginning of a flow of talent from industry back to academia. Currently there are over 30 academic small molecule screening centers, including the MLSCN
centers (see http://web.memberclicks.com/mc/directory/viewSaveSearch.do?searchId=48085&hidOrgID=sbs; Gordon, 2007).

The mission and architecture of academic and industrial compound screening can be readily differentiated. A significant fraction of the research and development budget for compound screening in large pharmaceutical companies is focused on identifying potential “blockbuster” drugs. Thus, 90% of the current commercial research and development resources are spent on only 10% of the current worldwide human disease burden (Munos 2006). Academic investigators are often less encumbered by profit motive limitations when they select targets for screening. Moreover, investigators in the non-profit sector often are just seeking potent biological probes and the resulting compounds frequently lack the requisite pharmacokinetic and metabolic profiles for a good drug. Thus, academics are free to interrogate a chemical library that may not be viewed as “drug-like” but still has pharmacologically unique components. Funding for academic scientists often has longer time lines, measured in half decades, than that for industrial scientists, who must place a premium on flexibility and rapid assay development and analysis as corporate goals change. Finally, the product of academic screening is by its nature open access, such as in PubChem (see below) or ChemBank, whereas the result of an industrial screening exercise generally remains proprietary.

The differences in core missions between the public and private sector means there is opportunity for mutual benefits, or collaborations. Indeed there are some interesting public-private partnerships that have already emerged (Munos 2006). One of these is the Medicines for Malaria Venture,
which was established in 1999 to discover and develop new and affordable antimalarial drugs. This group has brought 40 public and private institutions together in a network comprising 300 scientists. Another example is the GAVI Alliance (www.gavialliance.org), which brings together private foundations, national governments, UNICEF, WHO, The World Bank, the vaccine industry and public health institutions in a $3 billion effort to increase children's access to vaccines in poor countries. The Initiative on Public-Private Partnership for Health (www.ippph.org) lists 92 different public-private partnerships focused on neglected diseases. These recent developments, most within the last five years, could have a major impact on the research focus of some academic centers. Furthermore, academic centers that support small molecule screening should be in an excellent position to train the next generation of drug discovery scientists.

**NIH Roadmap and Contemporary Pharmacology.** The NIH has launched an ambitious program to optimize their entire research portfolio, called the Roadmap Initiative (http://nihroadmap.nih.gov). One of the three main areas of the initiative is the New Pathways to Discovery component, which is expected to empower the research community with small molecule compounds for tools to perturb genes and pathways, as imaging probes in basic or clinical applications, or as starting points for the development of new therapeutics for human disease. To achieve this goal, the NIH has established ten new MLSCN Centers. The eventual goal of each MLSCN Center is to conduct 15 assays with 300,000 compounds each year, and to deposit these 4.5 million assay results on PubChem for public use.

**How do I get involved in the MLSCN?**
Overview of target-to-probe process. Academic investigators typically bring a zen-like intensity to their attempts to understand the biological functions of their favorite proteins. When it becomes important to develop a small molecule tool for more detailed exploration, the MLSCN stands ready to help. R03 and R21 NIH grant mechanisms are available to prepare your assay for HTS, often in collaboration with one of the existing MLSCN Centers (see below). In your own laboratory (or in collaboration with one of the MLSCN Centers) it is first necessary to convert your benchtop assay from microfuge tube or microscope slide to 96 or 384 well plate format, by carrying out the experiments explained in the next section. From this point, the process is summarized in Figure 1. The next step is to submit an R03 or X01 grant application to NIH (www.mli.nih.gov/funding/assay_solic_fund_ops.php). Once the project is accepted and assigned to a Center, further assay optimization and, if necessary, miniaturization, is carried out at the Center, then a high throughput primary screen of up to 100,000 compounds is performed generally at a single compound concentration (typically 10-30 \( \mu \text{M} \)), although one center now routinely engages in concentration profiling as a primary screening approach (Inglese et al., 2006). Hits are confirmed in one or more secondary screens, concentration-response relations are obtained to inform structure-activity analysis, and a decision is made whether to invest medicinal chemistry resources of the network to the project. Assuming a positive answer, a wide range of cheminformatics tools is brought to bear, both to identify commercially available analogs of promising hits and to assist medicinal chemists in the design and synthesis of novel analogs for bioassay. One or more iterations of synthesis and bioassay are performed until the desired goal of the project is achieved. All screening data are deposited in a public database, PubChem (see below). Thus the dual deliverables of the MLSCN are novel, useful chemical probes for the biomedical investigator and a public database relating chemical structures to biological effects.
From bench to robot. Adaptation of a successful bench assay to high throughput mode requires optimizing the robustness and stability of the assay readouts, taking into consideration cost per well as well as labor costs. From a practical point of view, it is well to keep in mind that if reagent costs are even as low as $0.20 per well, a 100,000 compound primary screen (without duplicates) will cost $20,000 plus labor. The MLSCN is gearing up for 300,000 compound screens, and for this reason it is highly worthwhile to spend time up-front optimizing the assay. A detailed description of assay optimization for high throughput screening has been compiled by investigators at Eli Lilly and the Chemical Genomics Center at NIH, available at http://www.ncgc.nih.gov/guidance/manual_toc.html. Very briefly, the process typically begins by optimizing the concentration of protein and substrate in each well for a biochemical assay, or the number of cells per well in the case of cell-based assays. The parameters to optimize are cost of reagents, the signal to noise or signal to background ratio, and the Z' factor. The Z' factor measures the quality of the assay itself without intervention of test compounds. This measure of assay robustness is calculated from equation 1.

\[
Z' = 1 - 3\left(\frac{SD_s + SD_b}{\mu_s - \mu_b}\right)
\]  

Where the s subscript refers to the maximum assay signal (e.g., in the presence of a screening concentration of agonist), b the minimum signal (e.g., in the absence of agonist), SD the standard deviation and \(\mu\) the mean signal in each condition. The signal to background ratio (S:B) is defined as \(\mu_s/\mu_b\), and the signal to noise ratio (S:N) is defined by equation 2:

\[
(S:N) = \frac{(\mu_s - \mu_b)}{\sqrt{(SD_s^2 + SD_b^2)}}
\]

A simple example is shown in figure 2A for a cell-based cytotoxicity assay using A549 lung tumor...
cells in culture. The assay is based on the ability of healthy, viable cells to reduce resazurin to resorufin, which is excited at 560 nm and emits fluorescence at 590 nm. Injured or dead cells lack active reducing enzymes. Various numbers of cells (50 to 150,000) were seeded into 96 well plates and treated with vehicle or 1 µM doxorubicin for 24 hr. Resazurin was then added to each well and the reaction allowed to proceed in the culture incubator for 15 min to 24 hr before reading the plate in an Analyst HT multimode plate reader. Best results were obtained after 3.5 to 4 hr incubation. A plot of Z’ and S:B against cell density after 3.5 hr incubation (Fig. 2A) indicates that 10,000 to 20,000 cells per well yield an optimum Z’ (=0.8) and signal-to-background (S:B) ratio of 12, although these robustness measures are not much reduced if only 4,000 cells per well are used. As a rule of thumb, conditions resulting in Z’>0.5 and S:B > 5 produce an assay robust enough for most high throughput screens.

The next step in assay optimization is to determine whether plate-to-plate and week-to-week variability is adequate. Figure 2B and C show results from four 96-well plates repeated on three separate weeks, each week representing a different cell splitting. The assay shows acceptable stability over the three week period, without evidence of “edge effects” (not shown). It is important to evaluate the effect of DMSO on the assay because DMSO is the vehicle for the compound libraries and so is often present at concentrations of 0.5-2% in the assay. In this case, Z’ was reduced to ~0.65 in the presence of 1% DMSO (not shown). Finally, it is important to show that a positive control, here doxorubicin, affects the assay readout in a stable manner over a multi-day test period. Some assays, for example of novel targets, may not have positive controls. Screening a small library such as the Library of Pharmacologically Active Compounds (LOPAC) often results in identification of a compound that can be used as a positive control for the purpose of assessing
assay quality, even if its mechanism of action is not the desired one. Figure 2D shows that the IC\textsubscript{50}
of doxorubicin varied less than 2-fold over this three week period. This cell viability assay was
viewed as ready for high throughput screening of small molecules applied in a final concentration of
25 µM (1% DMSO).

Another example is an assay being developed to screen for potentiators or novel agonists of the EP2
prostanoid receptor, which activates adenylyl cyclase. An HEK293 cell line that expresses human
EP2 receptors was used to optimize a time-resolved fluorescence resonance energy transfer
(TRFRET)-based immunoassay for cyclic AMP formation in response to the selective EP2 agonist,
butaprost. The method depends upon competition by cell-derived cAMP for binding of labeled
cAMP to a cAMP antibody. The FRET donor, an anti-cAMP antibody conjugated to europium
cryptate, is excited at 337 nm and emits at 620 nm. The half-life of europium emission is several
hundred µsec or longer, so in practice a time delay of 50-100 us is imposed between excitation and
emission readings to allow the intrinsic fluorescence of library compounds (typical half-life of
hundreds of nanosec) to subside. The FRET acceptor is a cAMP molecule conjugated to a
fluorophore that is excited at 620 nm and emits at 665 nm. The TR-FRET signal is then the ratio
(x10\textsuperscript{4}) of emission readings at 665 and 620 nm. A ratiometric measure reduces well-to-well
variability due to the presence of colored compounds, phenol red in culture medium, etc. As
expected the FRET signal decreases as cAMP concentration rises (Fig 3D). In the experiment
HEK293 or C6 glioma cells seeded into a 384 well plate were treated for 30 min with vehicle (0.3%
DMSO), forskolin (a strong activator of adenylyl cyclase), or butaprost, all in the presence of 200
µM IBMX to block phosphodiesterases. Fig 3A,B show that forskolin strongly reduces the FRET
signal (i.e., elevates cAMP) in both HEK and C6 cell lines, but butaprost is an effective activator of
cAMP production only in the HEK293 cell line. A cell density of 3,000 cells per well provided the optimal signal to background ratio (Fig 3A). Experiments with butaprost or forskolin as agonist indicate adequate signal stability across a 384 well plate (Fig 3C), with $Z'=0.62$ and S:B=9. The next steps for assay development would involve evaluation of plate-to-plate and week-to-week stability.

What does the MLSCN offer pharmacologists?

**Diverse screening platforms.** The MLSCN offers the research community an opportunity to leverage biology and chemistry resources to identify small molecule probes for innovative or challenging targets with the potential for insight into biological pathways that impact public health. The MLSCN initiative is a consortium of ten centers that, in its initial pilot phase, is developing the capacity to screen a large set of shared compounds (> 100,000) maintained in a central repository, in a highly diverse set of assays solicited from the scientific community. Simultaneously, the centers are developing informatics and chemistry capacities to optimize "hits" identified in the initial screening to produce chemical probes that can be used for *in vitro* studies to interrogate the targets or phenotypes studied in the assays. All of the HTS screening data from the MLSCN assays, including functional and pharmacologic selectivity data, are deposited into PubChem (http://pubchem.ncbi.nlm.nih.gov/), an open access database that provides biological and chemical information to researchers for use in studying biology and disease.

**Access to MLSCN resources.** Launched in June of 2005, the ten MLSCN centers comprise a network of shared management, experience and expertise that provides greater capability and productivity for biological discovery than would be found in any single academic center.
Collectively, the centers network has an array of cutting-edge HTS technology and detection platforms that include ultra HTS capabilities afforded by Kalypsys technology, quantitative HTS approaches (Inglese et al., 2006), high through microscopy for phenotypic screening, NMR-based approaches, and flow cytometry that allows for multiplexing of related assay targets (Table 1). The comprehensive technology and detection platforms give the network the ability to implement a diversity of target-based (e.g., G-protein coupled receptors, ion channels, transporters, kinases, enzymes), cell-based, phenotypic high content screens, and model organism-based assays that address areas of pharmacologic research in need of a chemical probe to dissect signaling pathways and drive biological discovery. More detailed information about the capabilities and expertise of each of the centers can be found at http://www.mli.nih.gov/mlscn/descriptions.php.

The network of centers has gained experience implementing a diverse array of HTS assays and has developed the capacity to conduct nearly 150 assay campaigns/year against the library of 150,000 compounds, generating over 10 million data points in PubChem to date. Collectively, the centers can conduct most assays including biochemical and cell based, phenotypic, multiplexed, infectious agent-based, and a limited number of whole organism-based assays. The NIH is soliciting HTS assay projects directed towards novel proteins, cellular phenotypes, biological functions, and disease mechanisms from the research community (http://grants.nih.gov/grants/guide/notice-files/NOT-RM-07-003.html). This program provides scientists access to MLSCN resources at no cost to the investigator. HTS assay projects are selected for implementation on the basis of peer review. Projects are prioritized based on the significance of the biological target, need for a chemical probe, and HTS readiness, and the assays are then assigned to each MLSCN center based on its expertise and technology. Collaboration among the assay provider, MLSCN biologists, and chemists is
critical for development of a critical path work plan for progressing from assay to probe, which
describes assay automation, secondary and counter-screening assays, and identification of ‘hits’
that are amenable to SAR, computational modeling, and synthetic chemistry to optimize small
molecule probes (Figure 4). Through February 2007, 92 assays have been selected for HTS
campaigns by the MLSCN. The assays span a variety of target classes (Figure 5A) and disease
relevance (Fig 5B).

**Access to a high-quality, compound repository.** The MLSMR, established in September 2004,
currently houses a collection of nearly 115,000 chemically diverse molecules with both proven
and unknown biological activities. Biofocus DPI/Gallapagos collects, maintains, and distributes the
The repository currently contains diverse compounds, targeted libraries (e.g., GPCRs, ion channels,
nuclear receptors, kinases, proteases, etc.), natural products, a specialty set of compounds with
known biological activity, including approved drugs, failed clinical candidates, veterinary
medications, toxins, metabolites, etc., and novel chemical structures from the Molecular Libraries
(Table 2). The Chemical Methodologies and Library Development (CMLD) centers, which are
supported by the National Institute of General Medical Sciences
([http://www.nigms.nih.gov/Initiatives/CMLD/Centers/](http://www.nigms.nih.gov/Initiatives/CMLD/Centers/)), have also begun to contribute novel
structures to the compound repository.

Over the next 3-5 years, the repository will grow to 500,000 compounds. There will be commercial
acquisitions based on rational selection strategies and efforts to obtain novel compounds from non-
commercial sources. The PSL initiative will continue to expand the library by generating small
molecules to explore areas of “chemical diversity space” through target-oriented synthesis of complex molecules, and natural products isolation and derivatization (http://grants.nih.gov/grants/guide/rfa-files/RFA-RM-06-003.html). In addition, the MLSMR is seeking unique chemical contributions from all sources meeting the criteria of at least 90% purity, sufficient water solubility for use in HTS, reasonable stability at room temperature, and preferably in solid form. Compounds obtained by high-throughput synthesis, medicinal or synthetic organic chemistry, and purified discrete natural products from microorganisms, plants or marine organisms are of interest (http://grants.nih.gov/grants/guide/notice-files/NOT-RM-07-005.html). Over time, the use of the shared library by the MLSCN should provide extensive biological annotation, generating a unique and rich dataset available in the public domain.

**Access to biological and chemical datasets.** PubChem is a public sector cheminformatics database of small organic molecule modulators developed by the National Center for Biotechnology Information and launched in September of 2004 (http://pubchem.ncbi.nlm.nih.gov/). It is an on-line resource providing comprehensive information on the biological activities of small molecules and search, retrieval, and data analysis tools such as structure search, structure-activity analysis and structure clustering to optimize the utility of chemical structure and bioactivity information, as well as integration with other NIH biomedical information sources such as PubMed and Genome, Protein, and Structure databases. To date, PubChem contains over 15.5 million substance records from more than 50 depositors (Figure 6A), over 6 million unique compound structures with links to bioassay descriptions, relevant literature, references, and assay data points (Fig 6D), and nearly 400 bioassay data sets, with more than 128 of these contributed by the MLSCN to date (Fig 6C). Examples of MLSCN bioassay summary data including target/assay description, assay protocol, and definition of compound bioactivity can be found at (http://pubchem.ncbi.nlm.nih.gov/assay/assay.cgi?aid=439;
Examples of probe reports generated by the MLSCN centers can be found at (http://molscreen.florida.scripps.edu/probes.html; http://ncgc.nih.gov/db/?aid=103). As the deposition of chemical structures and assays continue to rise over time, so does the number of users (Figure 6B), making this a valuable resource to the public and private sector.

Conclusions and summary.

The MLSCN, along with PubChem and the MLSMR, offers a new dimension in research opportunities for pharmacologists, chemists and biologists in the academic and non-profit sector. The sharing of small molecules, biological assays, and screening data with the larger scientific community represents a new public sector paradigm that promises to facilitate the understanding of basic biological mechanisms and shorten the timeline for drug development, with resulting benefits to public health, especially for rare and neglected disorders.

Acknowledgements. We appreciate the assistance of Drs. Carson Loomis, Ingrid Li, Jamie Driscoll and Steve Bryant for providing the data for figures 4-6 and Table 2.
References


Footnotes

This work supported in part by NIH grants 5U54HG003918 (RD) and 5U54MH074411 (JSL).
Figure legends

**Figure 1.** Overview of the probe development process.

**Figure 2.** Optimization of an Alamar blue cytotoxicity assay for A549 cells treated with doxorubicin in 96 well plates. A. Z' factor and S:B as function of number of cells per well. B,C. Stability of assay over three weeks (4 plates per week). D. Doxorubicin concentration-inhibition curve is stable over three weeks. Yuhong Du and Haian Fu, unpublished.

**Figure 3.** Activation of EP2 receptors in HEK293 cell line measured by immuno-FRET based assay for cAMP. A,B, Raw TR-FRET signals show that HEK293 but not C6 glioma cells respond to butaprost, whereas forskolin activates cAMP formation strongly in both lines. C, Assay stability across 384 well plate (symbols as in A), with FRET signal converted to cAMP concentration by standard curve in D. R Dingledeine and J. Wetherington, unpublished.

**Figure 4.** HTS assay campaign: path from assay to probe.

**Figure 5.** Diversity of assay targets and disease relevance of HTS assays assigned to the MLSCN.

**Figure 6.** Growth of PubChem composition and usage
# Tables

## Table 1. Core capabilities of the MLSCN network

<table>
<thead>
<tr>
<th>Assay Expertise</th>
<th>Technologies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Enzymes/proteases</td>
<td>Ultra HTS</td>
</tr>
<tr>
<td>G-protein coupled receptors</td>
<td>Quantitative HTS</td>
</tr>
<tr>
<td>Kinases</td>
<td>High throughput microscopy</td>
</tr>
<tr>
<td>Ion channels/transporters</td>
<td>Flow cytometry</td>
</tr>
<tr>
<td>Cytotoxicity</td>
<td>NMR-based methods</td>
</tr>
<tr>
<td>Protein-protein interactions</td>
<td>Virtual screening</td>
</tr>
<tr>
<td>Protein misfolding/degradation</td>
<td></td>
</tr>
<tr>
<td>High-content screens</td>
<td></td>
</tr>
<tr>
<td>Yeast-based assays</td>
<td></td>
</tr>
<tr>
<td>Zebrafish</td>
<td></td>
</tr>
</tbody>
</table>
Table 2. Chemical composition of the Molecular Libraries repository

<table>
<thead>
<tr>
<th>Compound Class</th>
<th>number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diverse structures</td>
<td>109,829</td>
</tr>
<tr>
<td>Targeted libraries</td>
<td></td>
</tr>
<tr>
<td>GPCR</td>
<td>2,339</td>
</tr>
<tr>
<td>kinases</td>
<td>2,677</td>
</tr>
<tr>
<td>ion channels</td>
<td>1,676</td>
</tr>
<tr>
<td>Proteases</td>
<td>662</td>
</tr>
<tr>
<td>nuclease</td>
<td>110</td>
</tr>
<tr>
<td>Specialty sets</td>
<td>853</td>
</tr>
<tr>
<td>Non-commercial compounds</td>
<td></td>
</tr>
<tr>
<td>from PSLs, CMLDs, MLSCN hit followup</td>
<td>24,489</td>
</tr>
<tr>
<td>Natural products</td>
<td>&lt;1,000</td>
</tr>
<tr>
<td>Total</td>
<td>143,635</td>
</tr>
</tbody>
</table>
Research tool
PubChem

Hit optimization

ChemINFORMATICS

High throughput screening

Assay adaptation

MLSCN
you

Target identification

Fig 1
Fig 2
Fig 3
Assay Acceptance

Assay Implementation

Assay Validation

Primary Screen of Assay

Hit Confirmation

Hit Validation

Secondary Screen

Counter Screen

Purchased Analogs of Hits

Analogs by Synthetic Chemistry

Pharmacology of hits

Chemical Probe

Fig 4
Fig 5

A. Molecular target
- Enzyme
- Signaling Pathway
- Receptor or Protein
- Protein-Protein interaction
- Channel or Transporter
- Transcription factors
- Protein Folding
- Protein-DNA or RNA interaction
- Protein synthesis
- Other

B. Disease relevance
- Cancer
- Neurological Disorders
- Infectious diseases
- Immune disorders
- Metabolic disorders
- Cardiovascular disease
- Biology

Number of assays
Fig 6