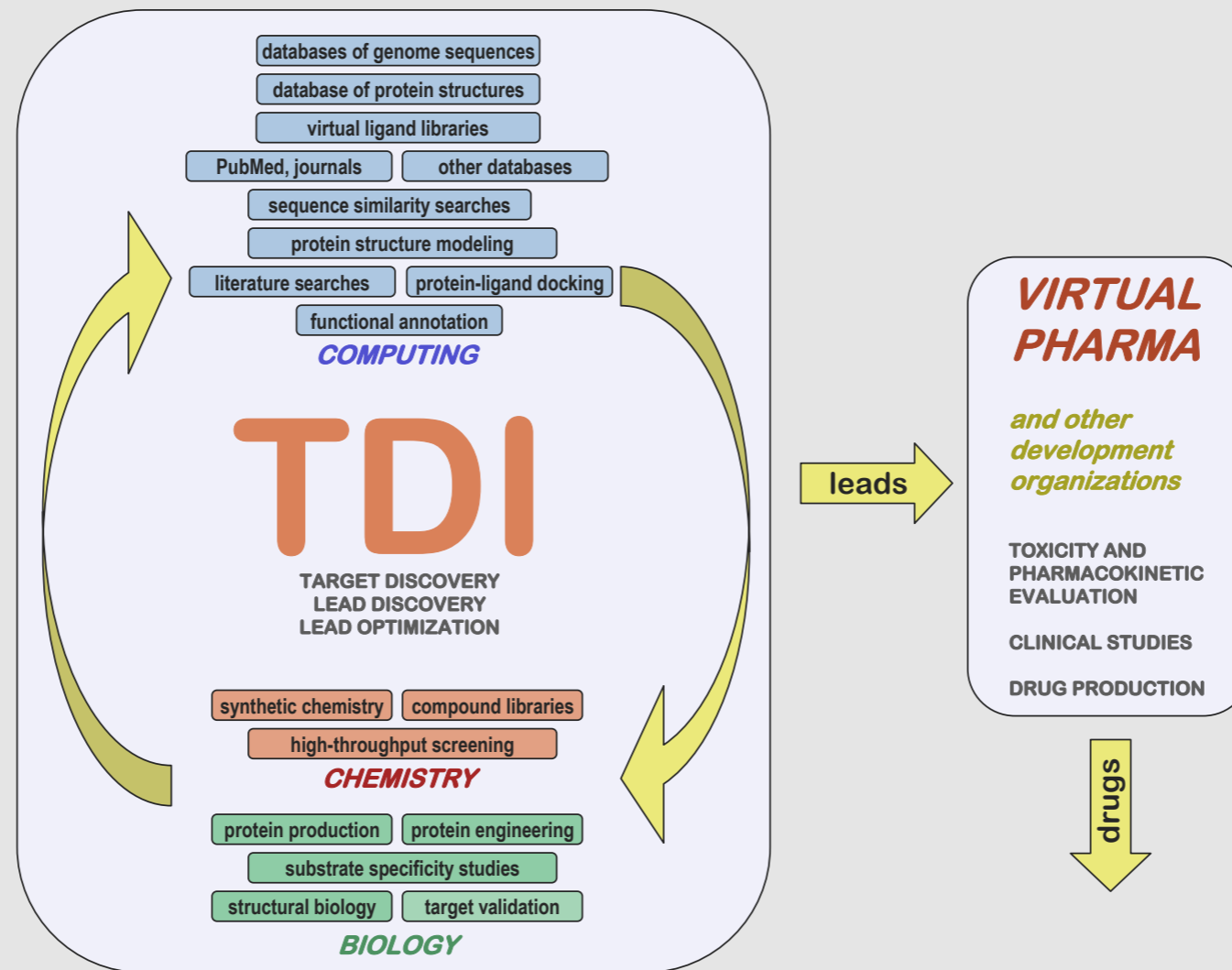


A “kernel” for the Tropical Disease Initiative

An open source approach to drug discovery



Marc A. Marti-Renom

<http://sgu.bioinfo.cipf.es>

Structural Genomics Unit
Bioinformatics Department
Prince Felipe Research Center (CIPF), Valencia, Spain



TDI *a story*



2004

.Steve Maurer (Berkeley) and Arti Rai (Duke)
.PLoS Medicine, Dec. 2004. Vol 1(3):e56

2005

.TDI web site <http://TropicalDisease.org>
.Ginger Taylor and The Synaptic Leap



2006

.Maurer and Sali 41th in “50 Who Matter”
.TSL web site <http://TheSynapticLeap.org>



2008

.TDI kernel <http://TropicalDisease.org/kernel>

Initial feed-back...

14 Mar 2005

I think TDI is a unique and very interesting project. I would like so much to make something for it...

So, where are we going? What's happening? What can we do?

I still trust in open source drug discovery. :-))

Luca Brivio

any, the bottlenecks are?
areas and potential avenues to explore,
action Plan!

9 Mar 2005

I'm a programmer, not a
the list active :)

**GNU started with RMS.
Linux started with Linu
You need someone gre
sending patches...**

I am interested in beginning rese
disease for underserved populatio
however, confused.

**If someone will tell me where to
begin on, I'd be grateful.**

Regards,
Jacob Lester

Thank you kindly,
Adam Huber

I know this is chicken-egg, but someone needs to point this out, since I haven't seen this brought up in the papers or the website.

And you might consider merging into the bios.net effort mentioned already. Together, you just might reach the critical mass for things to take off. Consider this like when people jumped off the HURD project to come together and make linux work.

Daniel Amelang

stic that the rest

Stephen Mark Maurer

Initial feed-back...

14 Mar 2005

I think TDI is a unique and very interesting project. I v
it...

So, where are we going? What's happening? What

I still trust in open source drug discovery. :-))

Luca Brivio

16 Feb 2005

Hi,

It would be interesting to know what, if any, the bottlenecks are?

The Wiki site contains many interesting ideas and potential avenues to explore, but from what I can see it is **lacking an Action Plan!**

Regards,
Jacob Lester

9 Mar 2005

I'm a programmer, not a
the list active :)

**GNU started with RMS.
Linux started with Linu
You need someone gre
sending patches...**

I am interested in beginning rese
disease for underserved populatio
however, confused.

**If someone will tell me where to
begin on, I'd be grateful.**

Thank you kindly,
Adam Huber

I know this is chicken-egg, but someone needs to point this out, since I haven't seen this brought up in the papers or the website.

And you might consider merging into the bios.net effort mentioned already. Together, you just might reach the critical mass for things to take off. Consider this like when people jumped off the HURD project to come together and make linux work.

Daniel Amelang

stic that the rest

Stephen Mark Maurer

Initial feed-back...

14 Mar 2005

I think TDI is a unique and very interesting project. I v
it...

So, where are we going? What's happening? What

I still trust in open s

Luca Brivio

16 Feb 2005

Hi,

10 Feb 2005

Hello,

My name is Adam Huber and I am a medical student at UNSW in Sydney Australia. I am interested in beginning research focused on tropical and infectious disease for underserved populations (A mission that seemingly matches TDI). I am, however, confused.

If someone will tell me where to sign up and give me some research topics to begin on, I'd be grateful.

Thank you kindly,
Adam Huber

9 Mar 2005

I'm a programmer, not a
the list active :)

**GNU started with RMS.
Linux started with Linu
You need someone gre
sending patches...**

I know this is chicken-egg, but someone needs to point this out, since I haven't seen this brought up in the papers or the website.

And you might consider merging into the bios.net effort mentioned already. Together, you just might reach the critical mass for things to take off. Consider this like when people jumped off the HURD project to come together and make linux work.

Daniel Amelang

Stephen Mark Maurer

bottlenecks are?
potential avenues to explore,
n!

stic that the rest

Initial feed-back...

14 Mar 2005

I think TDI is a unique and very interesting project. I v
it...

So, where are we going? What's happening? Who

I still trust in open s

Luca Brivio

16 Feb 2005

Hi,

10 Feb 2005

Hello,

My name is Adam Huber and I am a medical student at UNSW in Sydney Australia.
I am interested in beginning research focused on tropical and infectious
disease for underserved populations (A mission that seemingly matches TDI). I am,

bottlenecks are?

potential avenues to explore,

n!

9 Mar 2005

I'm a programmer, not a bioinformatician, but I stumbled across your site and thought I'd say something to keep the list active :)

GNU started with RMS. He gave us programming/administration tools to play with.

Linux started with Linus. He released an operating system for us to play with.

You need someone great in the field to release something for everyone to 'play with'. Then people start sending patches...

I know this is chicken-egg, but someone needs to point this out, since I haven't seen this brought up in the papers or the website.

And you might consider merging into the bios.net effort mentioned already. Together, you just might reach the critical mass for things to take off. Consider this like when people jumped off the HURD project to come together and make linux work.

Daniel Amelang

stic that the rest

Stephen Mark Maurer

Initial feed-back...

14 Mar 2005

I think TDI is a unique and very interesting project. I v
it...

So, where are we going? What's happening? What

I still trust in open s

Luca Brivio

16 Feb 2005

Hi,

10 Feb 2005

Hello,

My name is Adam Huber and I am a medical student at UNSW in Sydney Australia.
I am interested in beginning research focused on tropical and infectious
disease for underserved populations (A mission that seemingly matches TDI). I am,

bottlenecks are?

potential avenues to explore,

n!

9 Mar 2005

I'm a programmer, not a bioinformatician, but I stumbled across your site and thought I'd say something to keep the list active :)

GNU started with RMS. He gave us programming/administration tools to play with.

Linux started with Linus. He released an operating system for us to play with.

You need someone great in the field to release something for everyone to 'play with'. Then people start sending patches...

I know this is chicken-egg, but someone needs
papers or the website.

And you might consider merging into the bios
critical mass for things to take off. Consider th
together and make linux work.

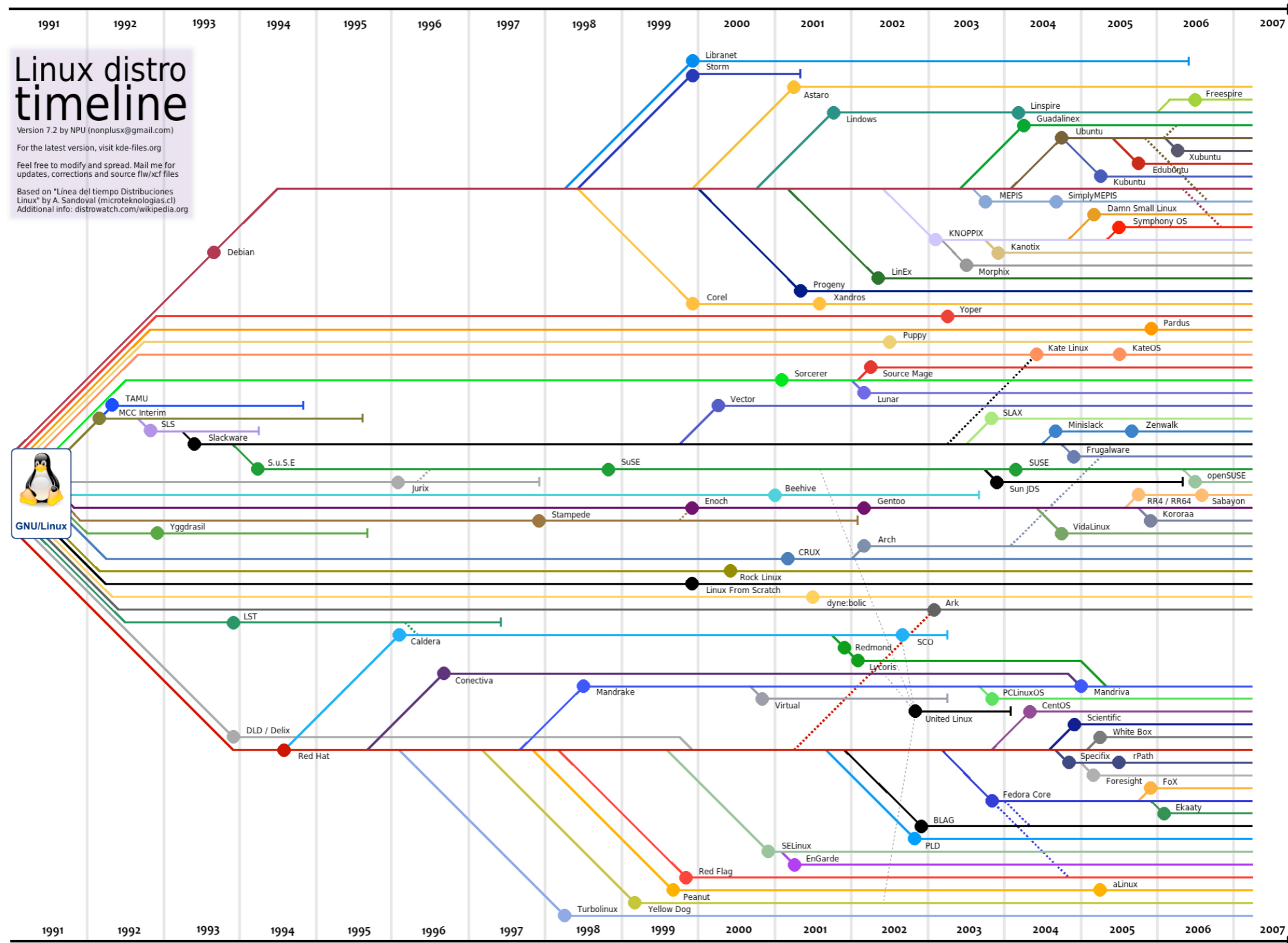
Daniel Amelang

19 Jan 2005

**If we do the science well, I'm optimistic that the rest
of TDI will fall into place.**

Stephen Mark Maurer

Open Source without a Kernel?



Is it possible? ...

- 1. In silico drug discovery**
2. Chemistry
3. Stem cell lines

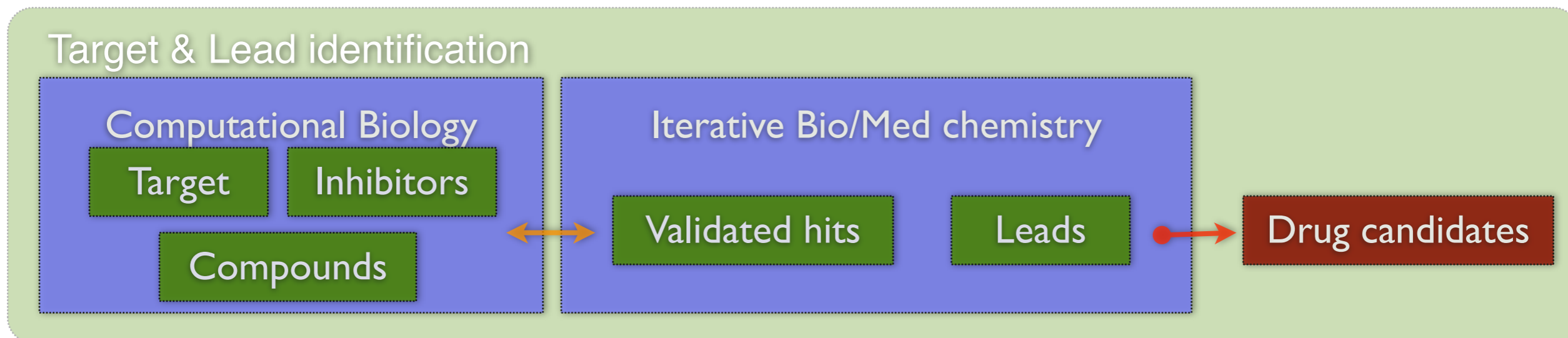
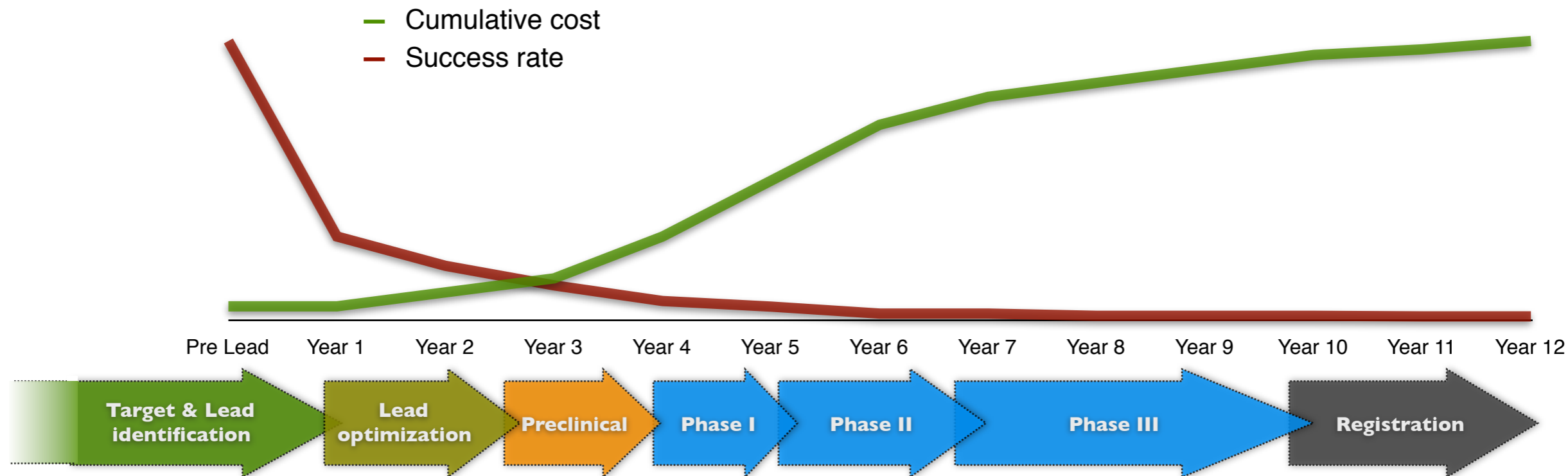
NEXT STEPS

4. Phase I to III trials
5. Phase IV trials

AMBITIOUS GOALS

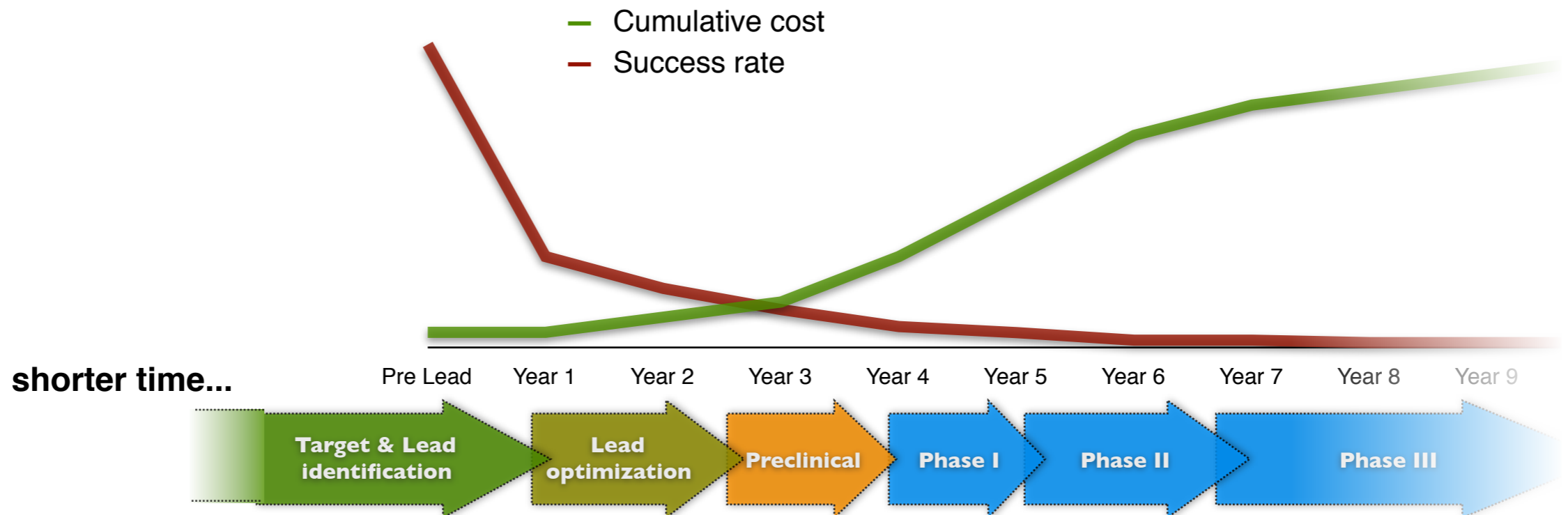
Maurer, Stephen M., "Open Source Drug Discovery: Finding a Niche (or Maybe Several) (April 2007)

Drug Discovery pipeline



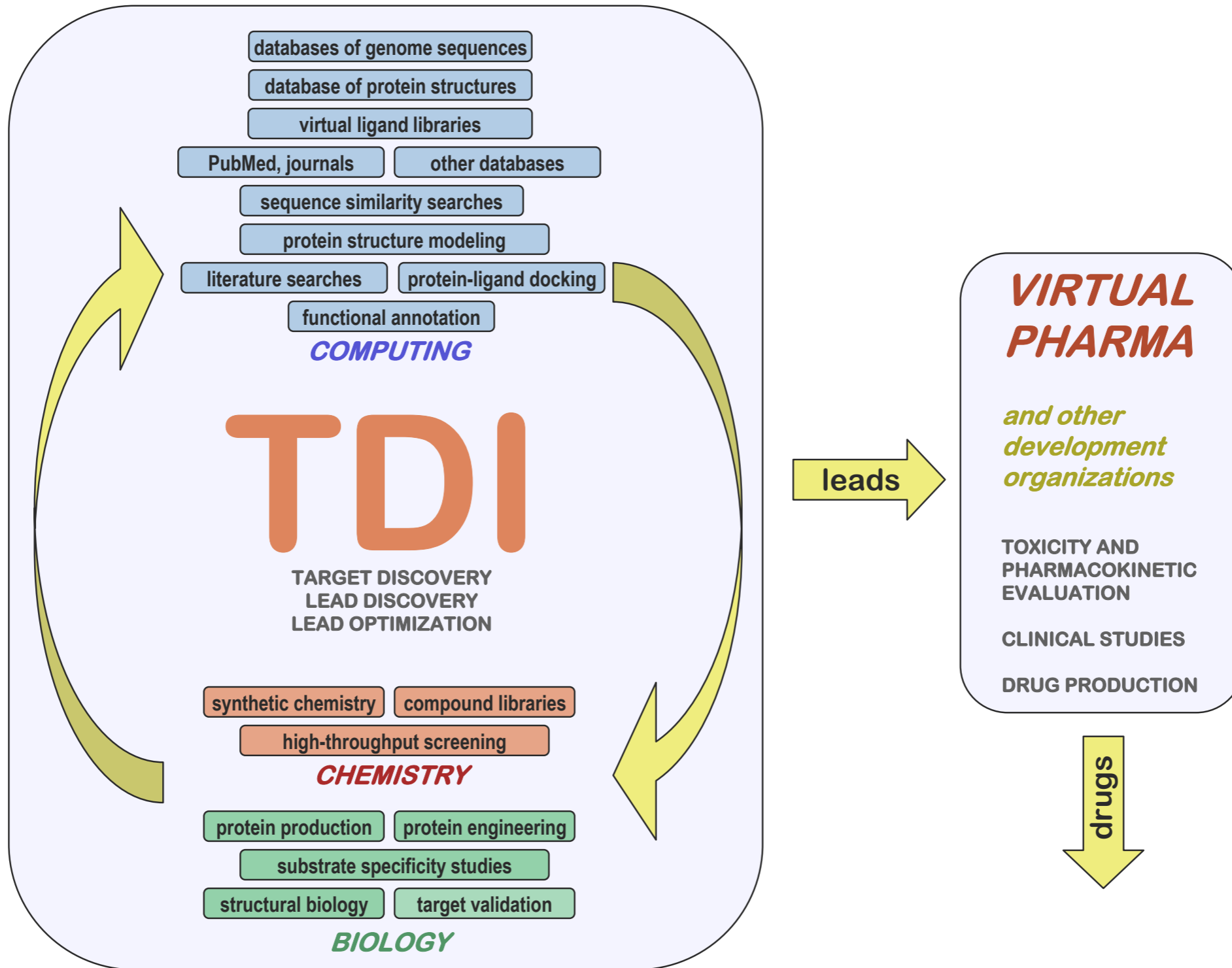
Adapted from: - Nwaka & Ridley. (2003) *Nature Reviews. Drug Discovery*. 2:919
- Austin, Brady, Insel & collins. (2004) *Science*. 306:1138

Drug Discovery pipeline



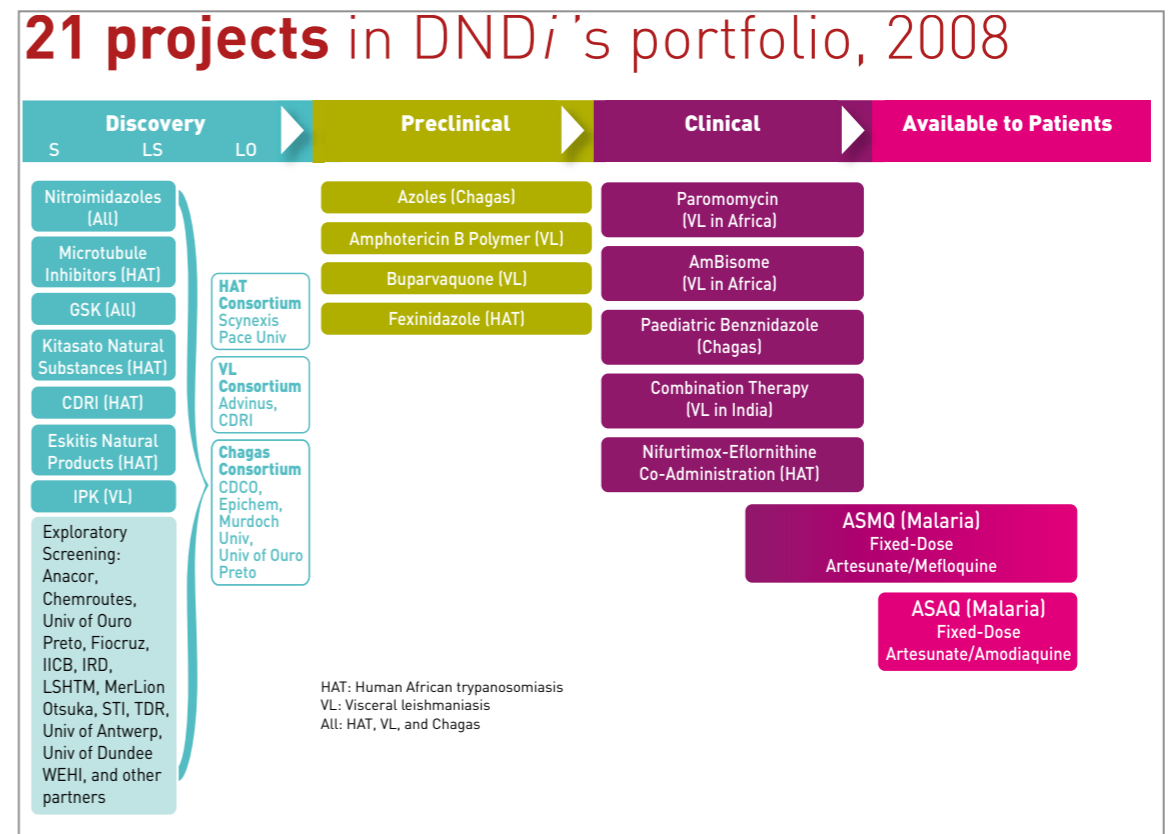
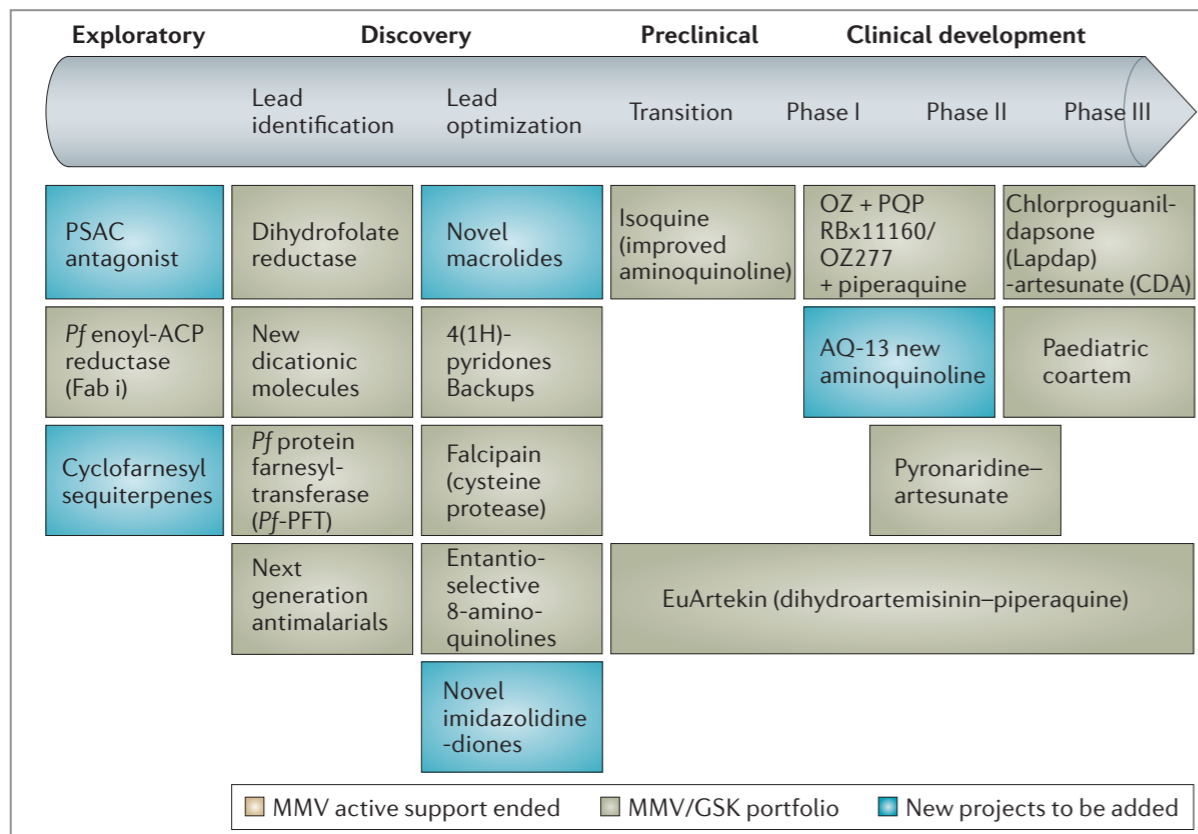
- + Completeness of genome projects (eg, Malaria)
- + New and more complete biological databases
- + New software and computers (cheaper and faster)
- + Internet == more people == less cost

TDI flowchart



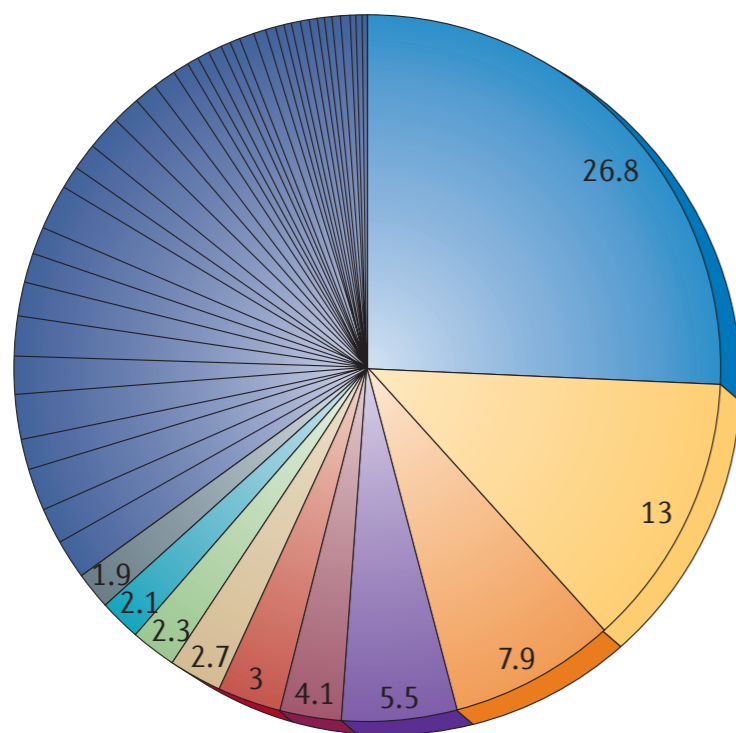
Non-Profit organizations

Open-Source + Out-Source = low cost business model



Munos (2006) Nature Reviews. Drug Discovery.

Number of (new) targets?



- Rhodopsin-like GPCRs
- Nuclear receptors
- Ligand-gated ion channels
- Voltage-gated ion channels
- Penicillin-binding protein
- Myeloperoxidase-like
- Sodium: neurotransmitter symporter family
- Type II DNA topoisomerase
- Fibronectin type III
- Cytochrome P450

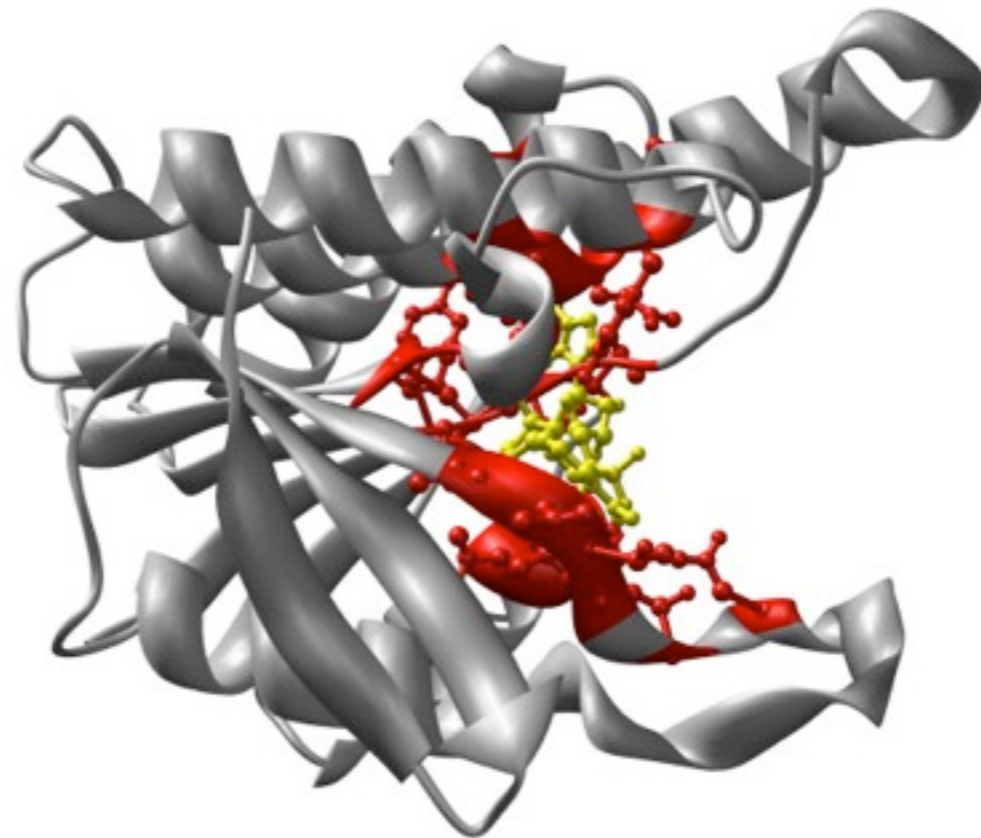
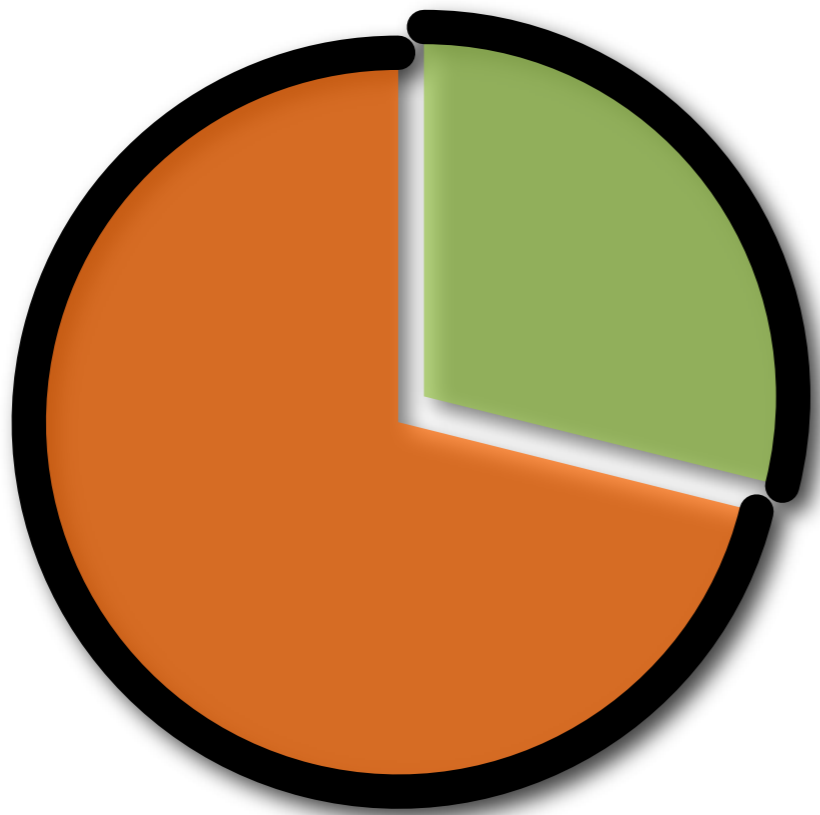
“... of 361 new molecular entities approved by the FDA between 1989 and 2000, 76% targeted a precedented drugged domain and only 6% targeted a previously undrugged domain ...”

Class of drug target	Species	Number of molecular targets
Targets of approved drugs	Pathogen and human	324
Human genome targets of approved drugs	Human	266
Targets of approved small-molecule drugs	Pathogen and human	248
Targets of approved small-molecule drugs	Human	207
Targets of approved oral small-molecule drugs	Pathogen and human	227
Targets of approved oral small-molecule drugs	Human	186
Targets of approved therapeutic antibodies	Human	15
Targets of approved biologicals	Pathogen and human	76

Where are new the targets for tropical diseases?

Overington et al. How many drug targets are there?. Nature reviews Drug discovery (2006) vol. 5 (12) pp. 993-6

Predicting binding sites in protein structure models.



UCSF

Duke
UNIVERSITY

PRINCIPE FELIPE
CENTRO DE INVESTIGACION
CERAMIO DE INGENIERIA

Need is High in the Tail

- DALY Burden Per Disease in Developed Countries
- DALY Burden Per Disease in Developing Countries



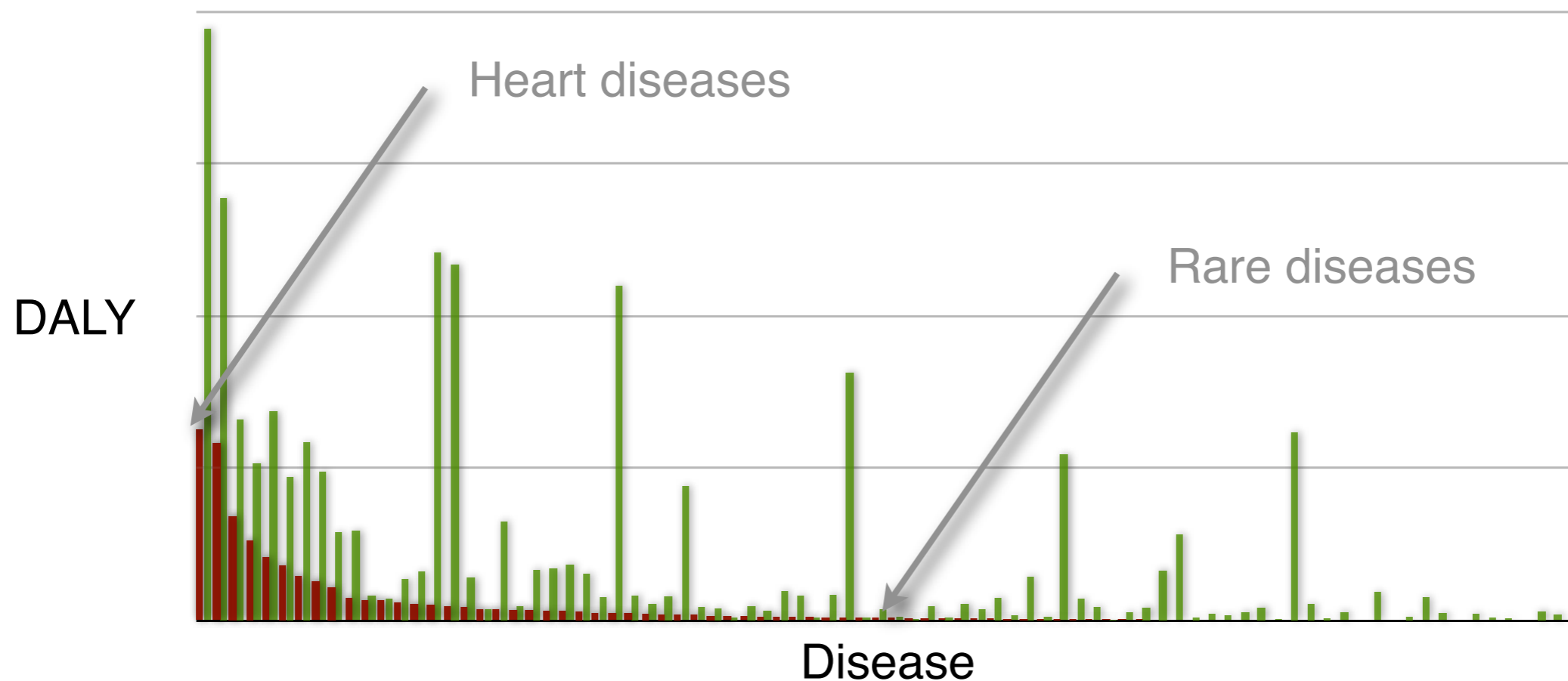
Disease data taken from WHO, *World Health Report 2004*
DALY - Disability adjusted life years

DALY is not a perfect measure of market size, but is certainly a good measure for importance.

DALYs for a disease are the sum of the years of life lost due to premature mortality (YLL) in the population and the years lost due to disability (YLD) for incident cases of the health condition. The DALY is a health gap measure that extends the concept of potential years of life lost due to premature death (PYLL) to include equivalent years of 'healthy' life lost in states of less than full health, broadly termed disability. One DALY represents the loss of one year of equivalent full health.

Need is High in the Tail

- DALY Burden Per Disease in Developed Countries
- DALY Burden Per Disease in Developing Countries



Disease data taken from WHO, *World Health Report 2004*
DALY - Disability adjusted life years

DALY is not a perfect measure of market size, but is certainly a good measure for importance.

DALYs for a disease are the sum of the years of life lost due to premature mortality (YLL) in the population and the years lost due to disability (YLD) for incident cases of the health condition. The DALY is a health gap measure that extends the concept of potential years of life lost due to premature death (PYLL) to include equivalent years of 'healthy' life lost in states of less than full health, broadly termed disability. One DALY represents the loss of one year of equivalent full health.

“Unprofitable” Diseases and Global DALY (in 1000’s)

Malaria*	46,486	Trichuriasis	1,006
Tetanus	7,074	Japanese encephalitis	709
Lymphatic filariasis*	5,777	Chagas Disease*	667
Syphilis	4,200	Dengue*	616
Trachoma	2,329	Onchocerciasis*	484
Leishmaniasis*	2,090	Leprosy*	199
Ascariasis	1,817	Diphtheria	185
Schistosomiasis*	1,702	Poliomyelitis	151
Trypanosomiasis*	1,525	Hookworm disease	59

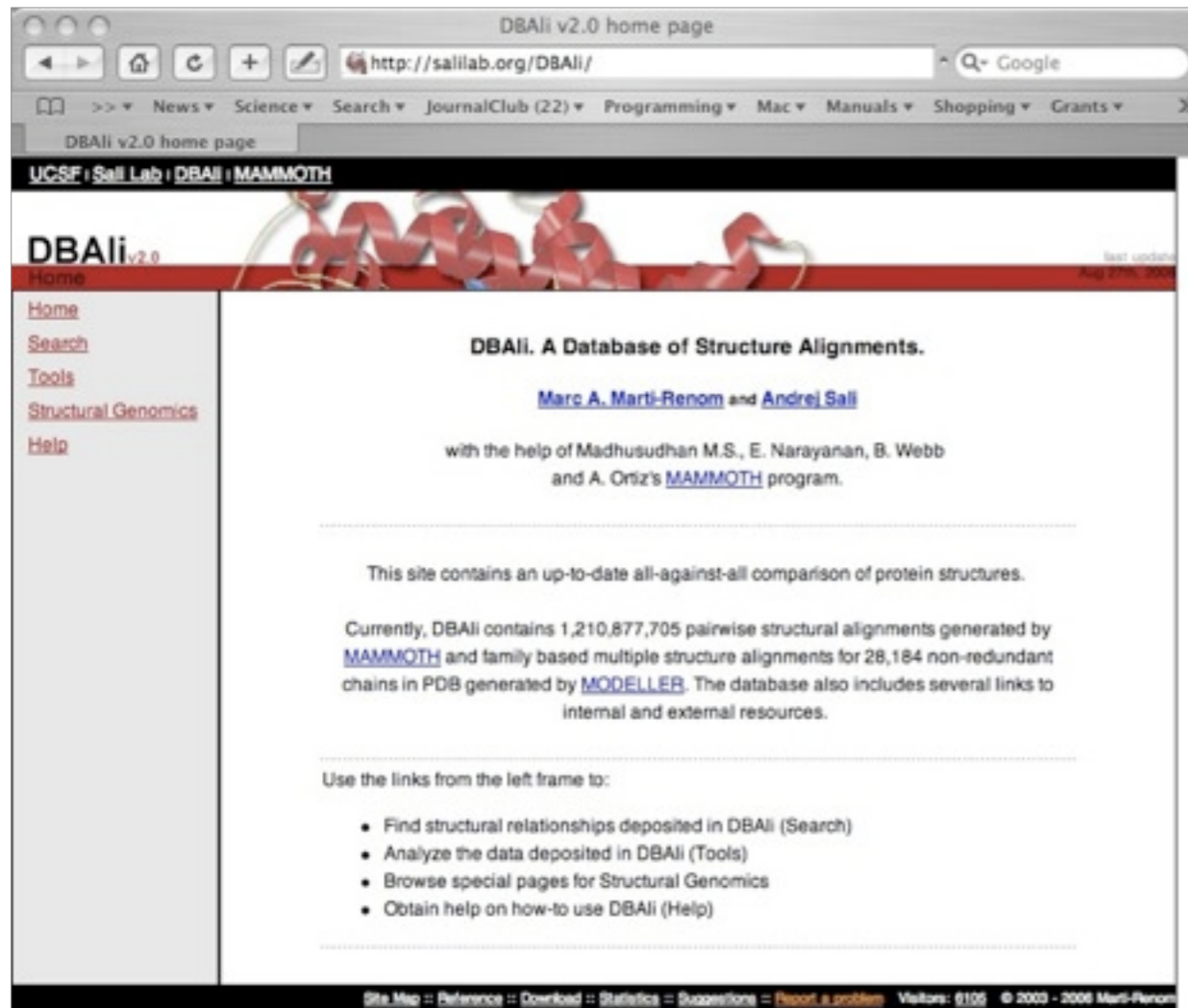
Disease data taken from WHO, *World Health Report 2004*

DALY - Disability adjusted life year in 1000’s.

* Officially listed in the WHO Tropical Disease Research [disease portfolio](#).

DBAli_{v2.0} database

<http://www.dbali.org>



- ✓ Fully-automatic
- ✓ Data is kept up-to-date with PDB releases
- ✓ Tools for “on the fly” classification of families.
- ✓ Easy to navigate
- ✓ Provides tools for structure analysis

Does not provide a stable classification similar to that of CATH or SCOP

Uses MAMMOTH for similarity detection

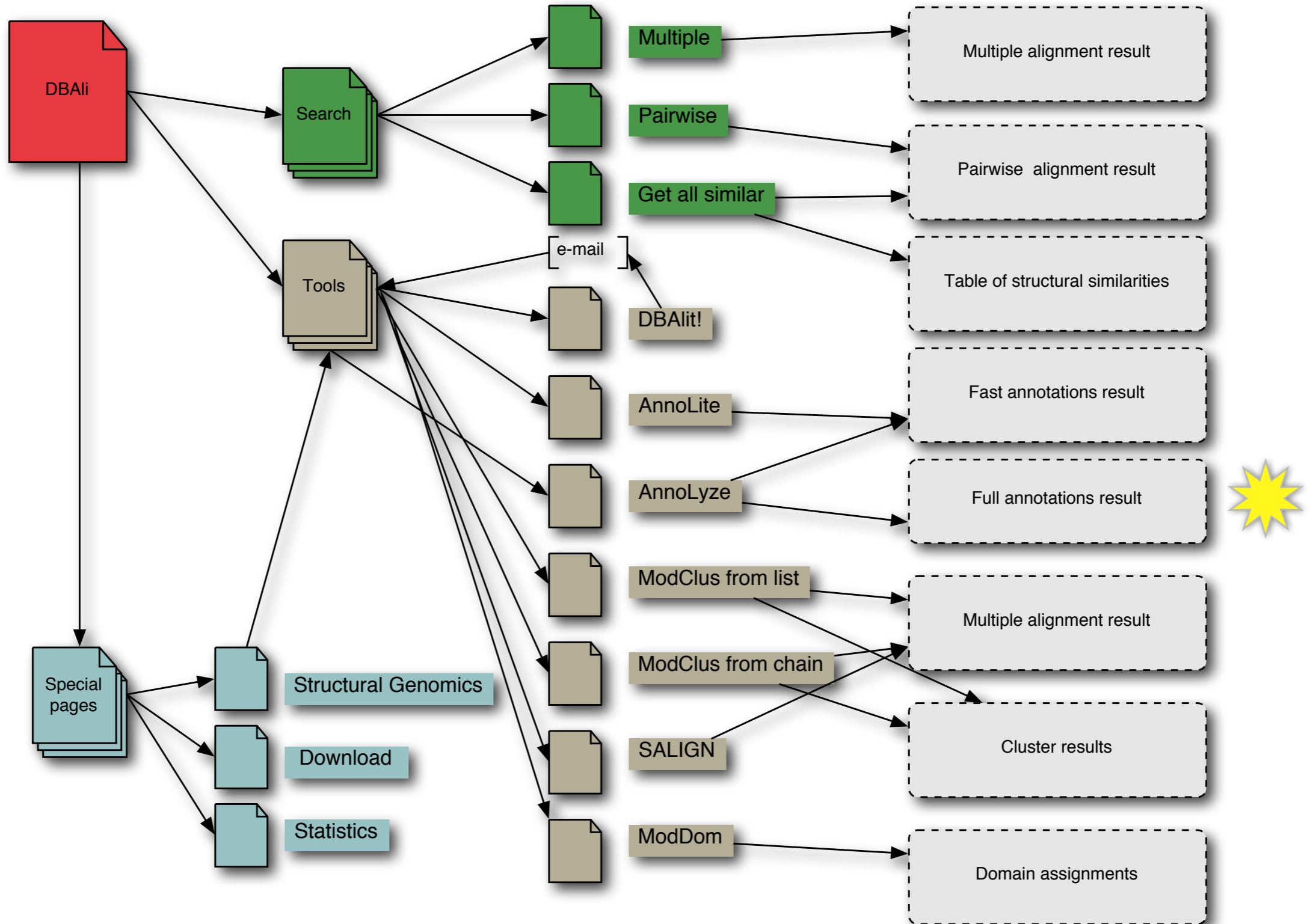
- ✓ VERY FAST!!!
- ✓ Good scoring system with significance

Ortiz AR, (2002) *Protein Sci.* 11 pp2606

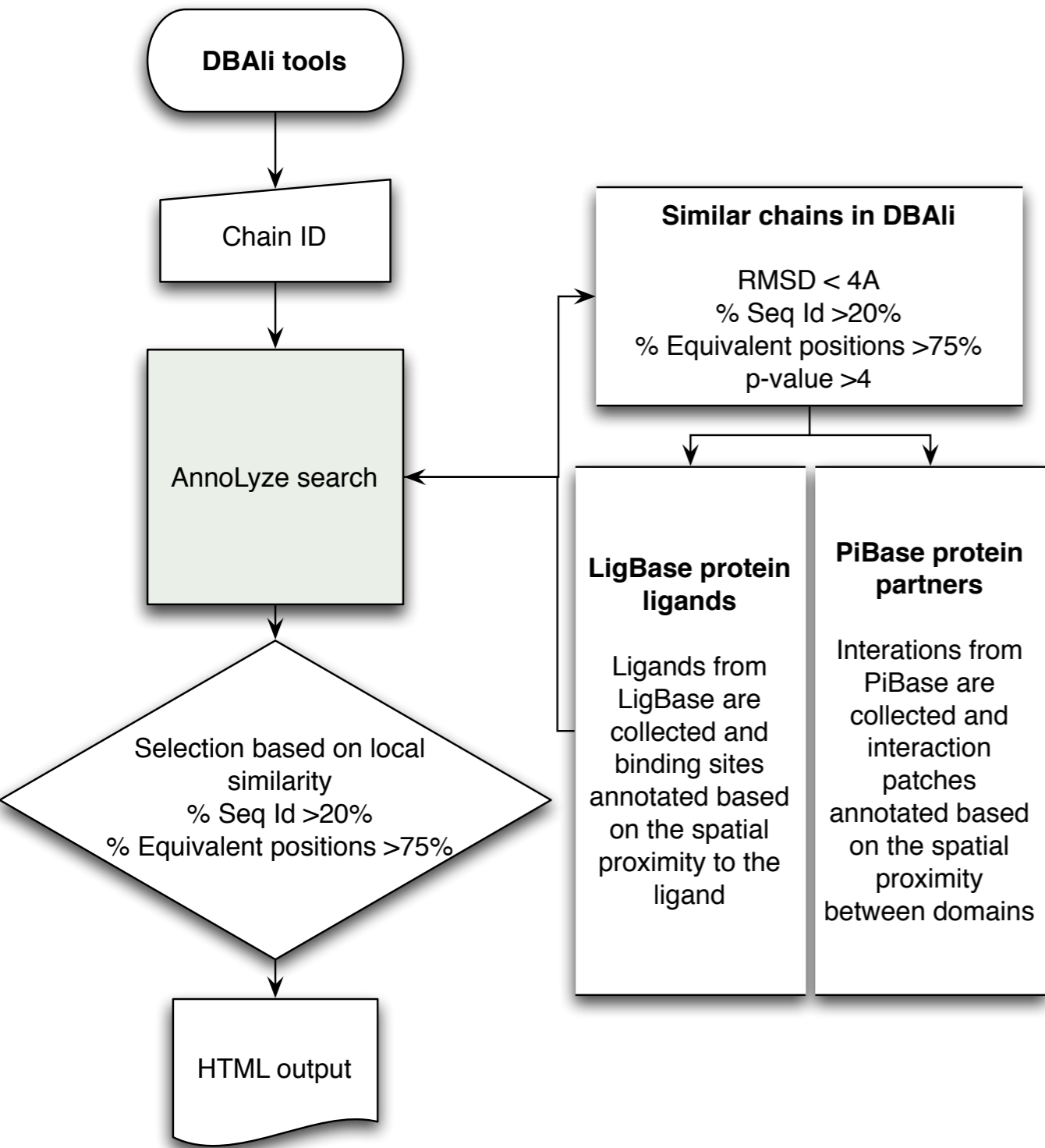
Marti-Renom et al. 2001. *Bioinformatics.* 17, 746

DBAli_{v2.0} database

<http://www.dbali.org>



Method



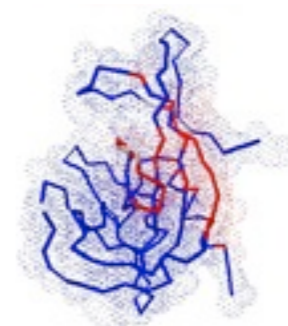
Inherited ligands: 4

Ligand	Av. binding site seq. id.	Av. residue conservation	Residues in predicted binding site (size proportional to the local conservation)
MO2	59.03	0.185	48 49 52 62 63 66 67 113 116
CRY	20.00	0.111	23 29 31 37 44 48 49 83 85 94 96 103 121
BOG	20.00	0.111	19 20 21 48 49 51 96 98 136
ACY	15.87	0.163	23 29 31 37 44 45 81 83 85 94 96 98 103 121 135



Inherited partners: 1

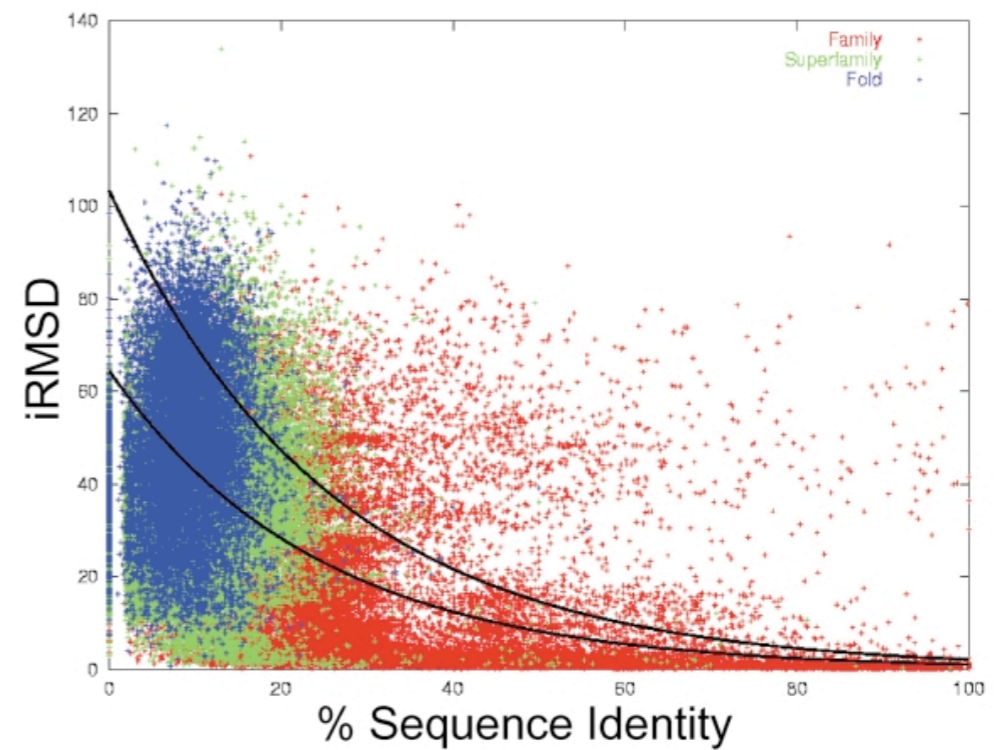
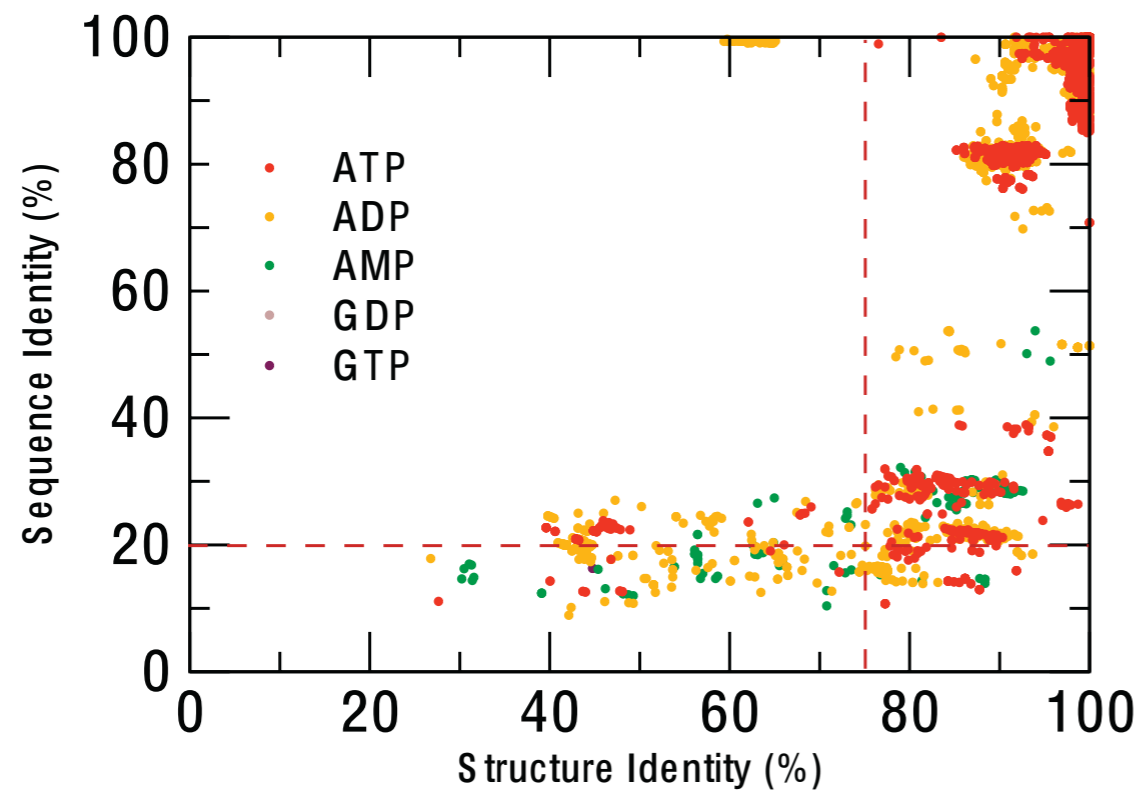
Partner	Av. binding site seq. id.	Av. residue conservation	Residues in predicted binding site (size proportional to the local conservation)
d.113.1.1	23.68	0.948	19 20 50 51 52 53 54 55 56 57 58 77 78 79 80 81 82 83 84 85 93 95 97 99 134 135 138 142 145



Scoring function

Ligands

Partners



Aloy *et al.* (2003) J.Mol.Biol. 332(5):989-98.

Benchmark

	Number of chains
Initial set*	78,167
LigBase**	30,126
Non-redundant set***	4,948 (8,846 ligands)

**all PDB chains larger than 30 aminoacids in length (8th of August, 2006)*

***annotated with at least one ligand in the LigBase database*

****not two chains can be structurally aligned within 3Å, superimposing more than 75% of their Ca atoms, result in a sequence alignment with more than 30% identity, and have a length difference inferior to 50aa*

Sensitivity .vs. Precision

	Optimal cut-off	Sensitivity (%) Recall or TPR	Precision (%)
Ligands	30%	71.9	13.7

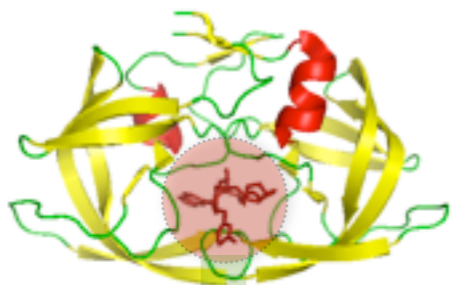
$$\text{Sensitivity} = \frac{TP}{TP + FN} \quad \text{Precision} = \frac{TP}{TP + FP}$$

~90-95% of residues correctly predicted

Comparative docking

Expansion

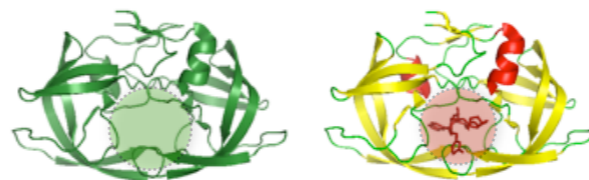
co-crystallized protein/ligand



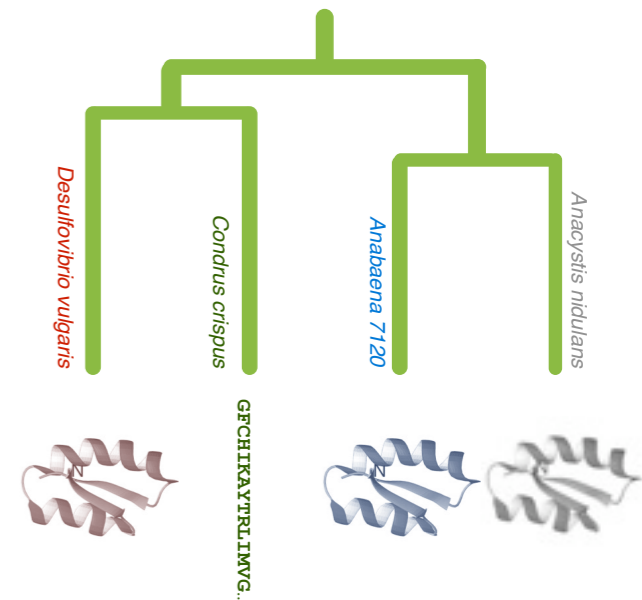
crystallized protein

2. Inheritance

model



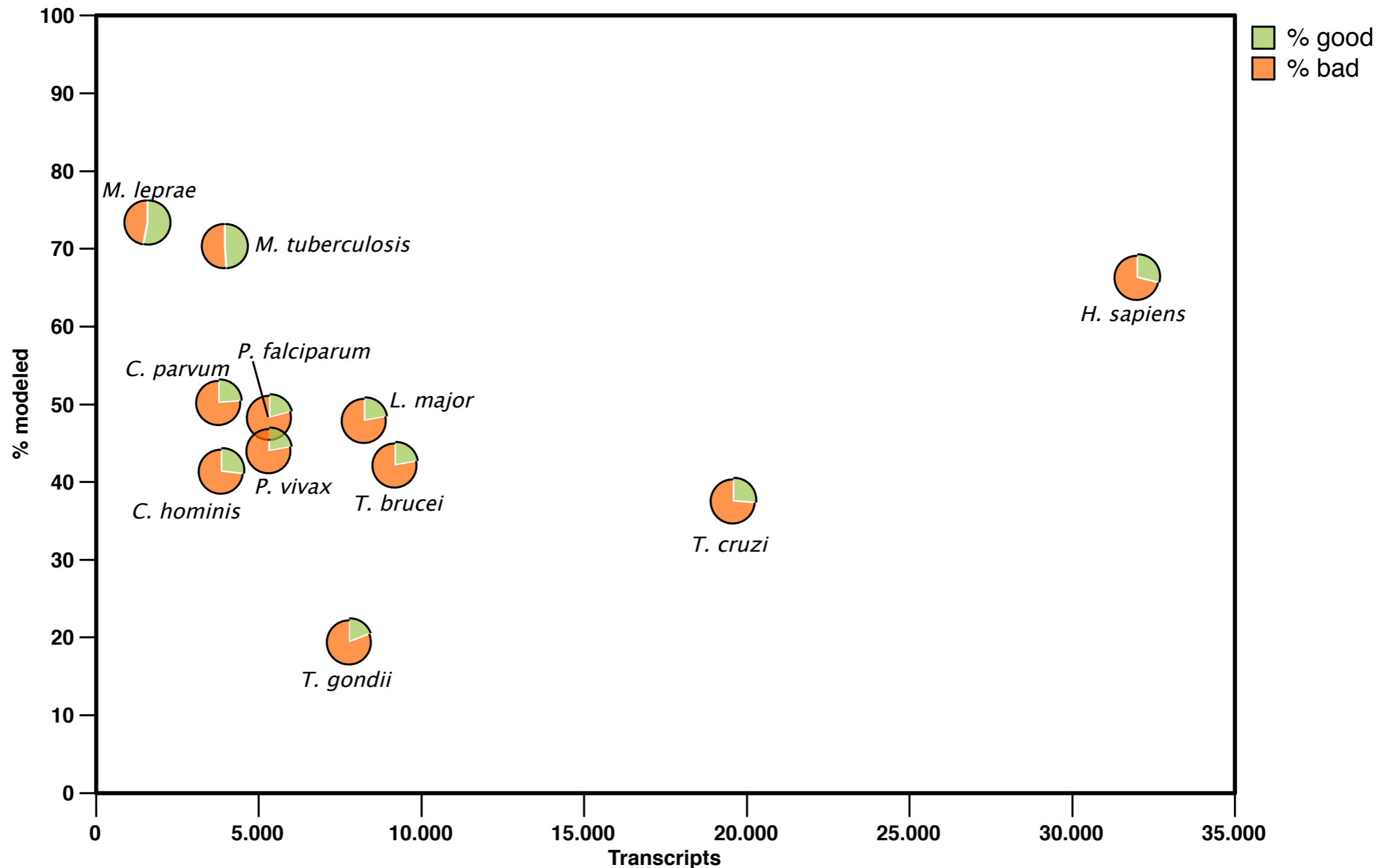
template



1. Modeling

Modeling Genomes

data from models generated by ModPipe (Eswar, Pieper & Sali)



A good model has MPQS of 1.0 or higher

Summary table

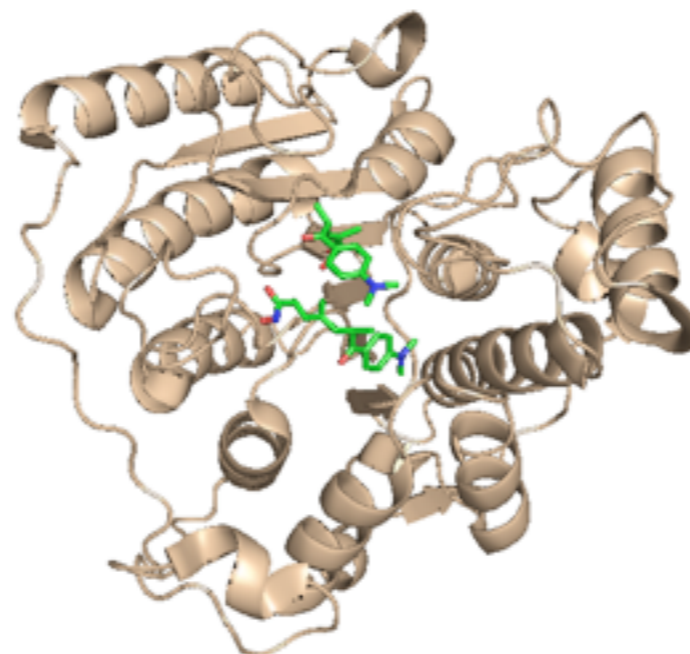
models with inherited ligands

29,271 targets with good models, 297 inherited a ligand/substance similar to a known drug in DrugBank

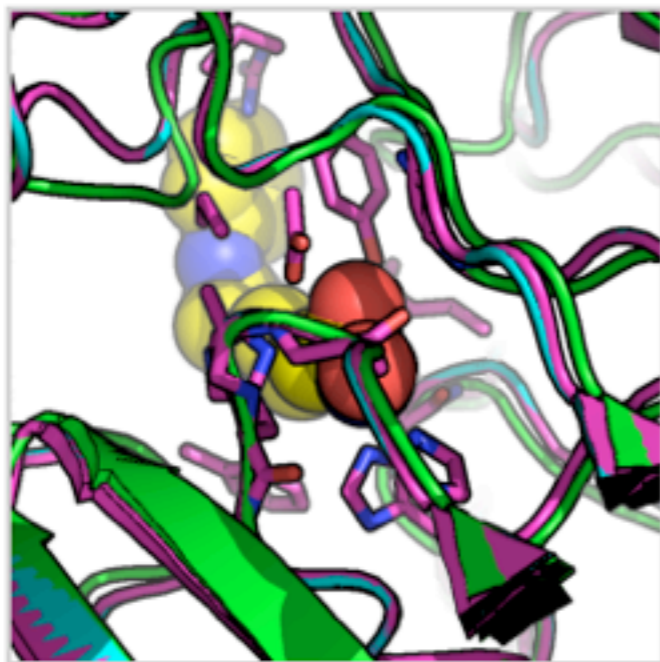
	Transcripts	Modeled targets	Selected models	Inherited ligands	Similar to a drug	Drugs
<i>C. hominis</i>	3,886	1,614	666	197	20	13
<i>C. parvum</i>	3,806	1,918	742	232	24	13
<i>L. major</i>	8,274	3,975	1,409	478	43	20
<i>M. leprae</i>	1,605	1,178	893	310	25	6
<i>M. tuberculosis</i>	3,991	2,808	1,608	365	30	10
<i>P. falciparum</i>	5,363	2,599	818	284	28	13
<i>P. vivax</i>	5,342	2,359	822	268	24	13
<i>T. brucei</i>	7,793	1,530	300	138	13	6
<i>T. cruzi</i>	19,607	7,390	3,070	769	51	28
<i>T. gondii</i>	9,210	3,900	1,386	458	39	21
TOTAL	68,877	29,271	11,714	3,499	297	143

L. major Histone deacetylase 2 + Vorinostat

Template 1t64A a human HDAC8 protein.



PDB	IO	Template	IO	Model	IO	Ligand	Exact	SupStr	SubStr	Similar
1c3sA	83.33/80.00	1t64A	36.00/1.47	LmjF21.0680.1.pdb	90.91/100.00	SHH	DB02546	DB02546	DB02546	DB02546



[DB02546](#) Vorinostat

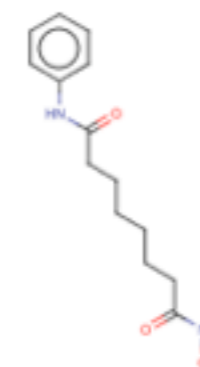
Small Molecule; Approved; Investigational

Drug categories:

Anti-Inflammatory Agents, Non-Steroidal
Anticarcinogenic Agents
Antineoplastic Agents
Enzyme Inhibitors

Drug indication:

For the treatment of cutaneous manifestations in patients with cutaneous T-cell lymphoma who have progressive, persistent or recurrent disease on or following two systemic therapies.



L. major Histone deacetylase 2 + Vorinostat

Literature

Proc. Natl. Acad. Sci. USA
Vol. 93, pp. 13143–13147, November 1996
Medical Sciences

Apicidin: A novel antiprotozoal agent that inhibits parasite histone deacetylase

(cyclic tetrapeptide / Apicomplexa / antiparasitic / malaria / coccidiosis)

SANDRA J. DARKIN-RATTRAY*[†], ANNE M. GURNETT*, ROBERT W. MYERS*, PAULA M. DULSKI*,
TAMI M. CRUMLEY*, JOHN J. ALLOCCO*, CHRISTINE CANNOVA*, PETER T. MEINKE[‡], STEVEN L. COLLETTI[‡],
MARIA A. BEDNAREK[‡], SHEO B. SINGH[§], MICHAEL A. GOETZ[§], ANNE W. DOMBROWSKI[§],
JON D. POLISHOOK[§], AND DENNIS M. SCHMATZ*

Departments of *Parasite Biochemistry and Cell Biology, [‡]Medicinal Chemistry, and [§]Natural Products Drug Discovery, Merck Research Laboratories, P.O. Box 2000, Rahway, NJ 07065

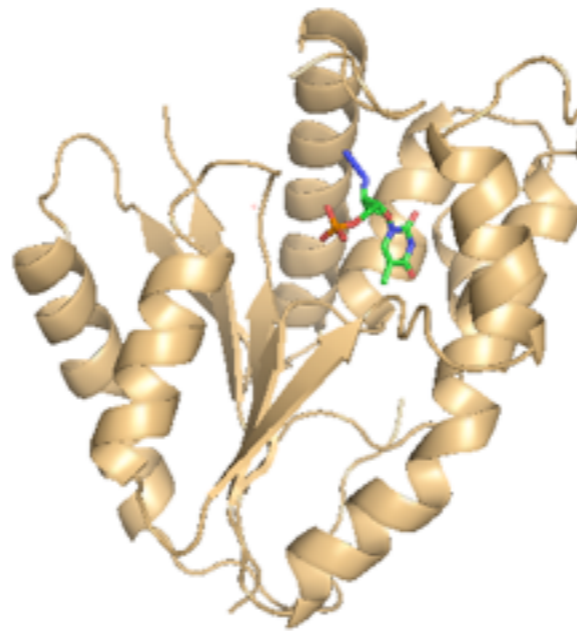
ANTIMICROBIAL AGENTS AND CHEMOTHERAPY, Apr. 2004, p. 1435–1436
0066-4804/04/\$08.00+0 DOI: 10.1128/AAC.48.4.1435–1436.2004
Copyright © 2004, American Society for Microbiology. All Rights Reserved.

Vol. 48, No. 4

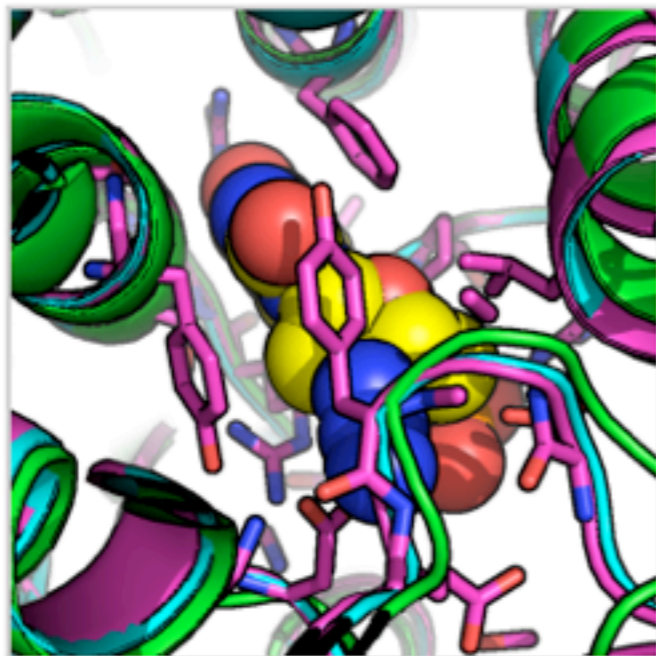
Antimalarial and Antileishmanial Activities of Aroyl-Pyrrolyl-Hydroxyamides, a New Class of Histone Deacetylase Inhibitors

P. falciparum thymidylate kinase + zidovudine

Template *3tmkA* a yeast thymidylate kinase.



PDB	IO	Template	IO	Model	IO	Ligand	Exact	SupStr	SubStr	Similar
2tmkB	100.00/100.00	3tmkA	41.00/1.49	PFL2465c.2.pdb	82.61/100.00	ATM		DB00495		DB00495



[DB00495](#) Zidovudine

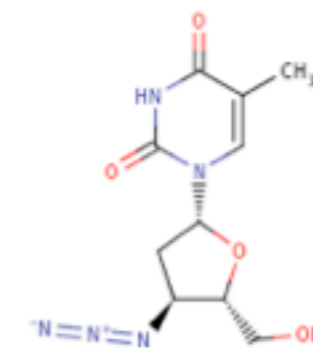
Small Molecule; Approved

Drug categories:

Anti-HIV Agents
Antimetabolites
Nucleoside and Nucleotide Reverse Transcriptase Inhibitors

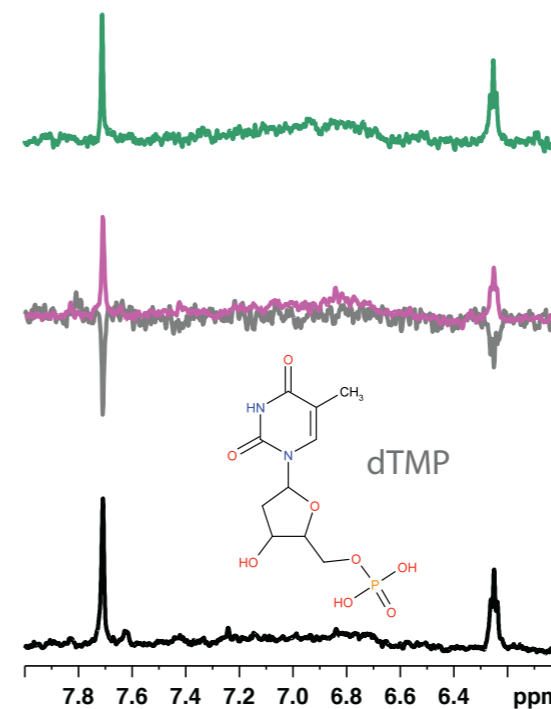
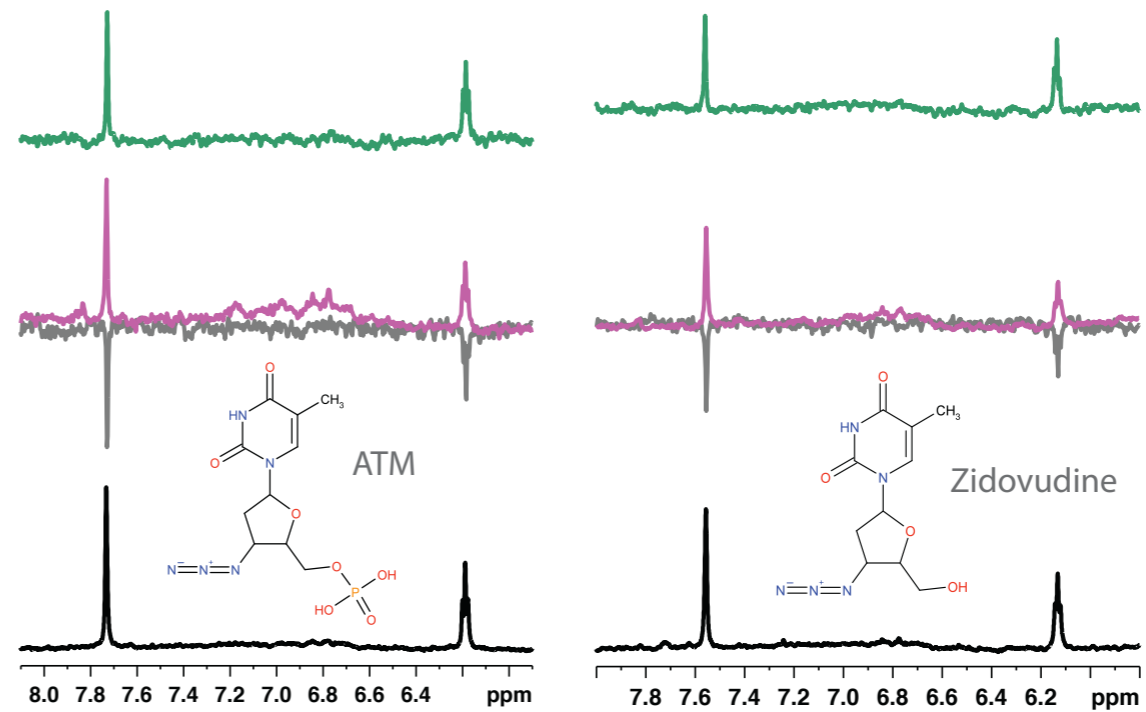
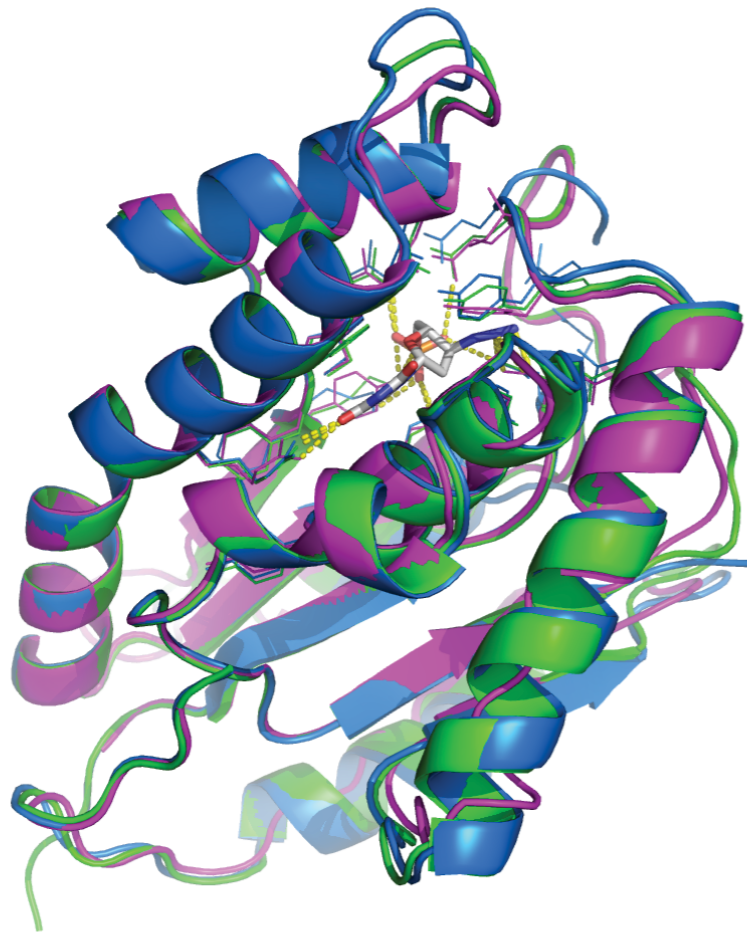
Drug indication:

For the treatment of human immunovirus (HIV) infections.



P. falciparum thymidylate kinase + zidovudine

NMR Water-LOGSY and STD experiments



Leticia Ortí, Rodrigo J. Carbajo, and Antonio Pineda-Lucena

TDI's kernel

<http://tropicaldisease.org/kernel>

TDI Kernel database » Q9GU59

http://tropicaldisease.org/kernel/q9gu59/

the **Tropical Disease Initiative** *an open source drug discovery project*

You are browsing version 1.0 (2008/05/01) of the TDI Kernel.

Posted on 05.07.08 to Target. Grab the feed. No comments yet. Add your thoughts or trackback from your own site. Edit this entry.

Putative histone deacetylase. predicted to bind 1 ligands [SHH]

UniPort id: **Q9GU59** [*C. parvum*]

Target keywords: ; Anticarcinogenic Agents; Antineoplastic Agents; Transcription; Chromatin regulator; Anti-inflammatory Agents, Non-Steroidal; Enzyme Inhibitors; Q9GU59; Transcription regulation; Nucleus

Do you consider this target suitable for drug discovery: ★★★★★ (No Ratings Yet)

Binding site prediction to approved drugs (need help reading this page?):

PDB	IC ₅₀	Template	IC ₅₀	Model	Score	Ligand	Exact	SupStr	SubStr	Similar
1c3sA	83.33/80.00	1t64A	37.00/1.47	cgd6_1380.1.pdb	90.91/100.00	SHH	DB02546	DB02546	DB02546	DB02546

DB02546 Vorinostat

Small Molecule; Approved; Investigational

Drug categories:

- Anti-inflammatory Agents, Non-Steroidal
- Anticarcinogenic Agents
- Antineoplastic Agents
- Enzyme Inhibitors

Drug indication:

For the treatment of cutaneous manifestations in patients with cutaneous T-cell lymphoma who have progressive, persistent or recurrent disease on or following two systemic therapies.

Show ligand **SHH**

OCTANEDIOICACIDHYDROXYAMIDEPHENYLAMIDE expanded from **1t64A** to template **1t64A** used for building a 3D model of **cgd6_1380.1.pdb**. Download the coordinates data/Q9GU59/Q9GU59_SHH.952.pdb

SEARCH KERNEL

Q Search...

Advanced Search

Browse the kernel

Download Q9GU59

Login / Register

Batch downloads

Help

Methods

CTDL BLOG IT!

Highest rated target:

- A7UD81 (5 out of 5)

2008 - Open Access.

Powered by WordPress.

Theme by Upstart Blogger.

TDI's kernel

<http://tropicaldisease.org/kernel>

L. Orti *et al.*, *Nat Biotechnol* **27**, 320 (Apr, 2009).

L. Orti *et al.*, *PLoS Negl Trop Dis* **3**, e418 (2009).

CORRESPONDENCE

A kernel for the Tropical Disease Initiative

To the Editor:

Identifying proteins that are good drug targets and finding drug leads that bind to them is generally a challenging problem. It is particularly difficult for neglected tropical diseases, such as malaria and tuberculosis, where research resources are relatively scarce¹. Fortunately, several developments improve our ability to deal with drug discovery for neglected diseases: first, the sequencing of many complete genomes of organisms that cause tropical diseases; second, the determination of a large number of protein structures; third, the creation of compound libraries, including already-approved drugs and fourth, the availability of improved bioinformatics analysis, including methods for comparative protein structure modeling, binding site identification, virtual ligand screening and drug design. Therefore, we are now in a position to increase the odds of identifying high-quality drug targets and drug leads for neglected tropical diseases. Here we encourage a collaboration among scientists to engage in drug discovery for tropical diseases by providing a 'kernel' for the Tropical Disease Initiative (TDI, <http://www.tropicaldisease.org/>)². As the Linux kernel did for open source code development, we suggest that the TDI kernel may help overcome a major stumbling block, in this case, for open source drug discovery: the absence of a critical mass of preexisting work that volunteers can build on incrementally. This kernel complements several other initiatives on neglected tropical diseases³⁻⁵, including collaborative web portals (e.g., <http://www.thesyntacteam.org/>), public-

private partnerships (e.g., <http://www.mmv.org/>) and private foundations (e.g., <http://www.gatesfoundation.org/>); for an updated list of initiatives, see the TDI website above.

The TDI kernel was derived with our software pipeline^{6,7} for predicting structures of protein sequences by comparative modeling, localizing small-molecule binding sites on the surfaces of the models and predicting ligands that bind to them. Specifically, the pipeline linked 297 proteins from ten pathogen genomes with already approved drugs that were developed for treating other diseases (Table 1). Such links, if proven experimentally, may significantly increase the efficiency of target identification, target validation, lead discovery, lead optimization and clinical trials. Two of the kernel targets were tested for their binding to a known drug by NMR spectroscopy, validating one of our predictions (Fig. 1 and Supplementary Data online). It is difficult to assess the accuracy of our computational predictions based on this limited experimental testing. Thus, we encourage other investigators to donate their expertise and facilities to test additional predictions. We hope the testing will occur within the

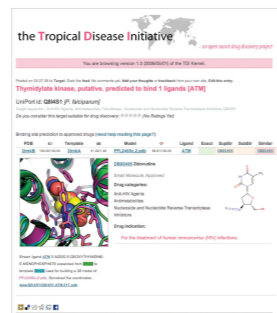


Figure 1 TDI kernel snapshot of the web page for the *Plasmodium falciparum* thymidylate kinase target (<http://tropicaldisease.org/kernel/q84411/>). Our computational pipeline predicted that thymidylate kinase from *P. falciparum* binds ATM (5'-azido-3'-deoxythymidine-5'-monophosphate), a supra-structure of the zidovudine drug approved for the treatment of HIV infection. The binding of this ligand to a site on the kinase was experimentally validated by one-dimensional Water-LOGSY⁸ and saturation transfer difference⁹ NMR experiments.

open source context, where results are made available with limited or no restrictions.

A freely downloadable version of the TDI kernel is available in accordance with the Science Commons protocol for implementing open access data (<http://sciencecommons.org/projects/publishing/open-access-data-protocol/>), which prescribes standard academic attribution and facilitates tracking of work but imposes no other restrictions. We do not seek intellectual property rights in the actual discoveries based on the TDI kernel, in the hope of reinvigorating drug discovery for neglected tropical diseases⁸. By minimizing restrictions on the data, including viral terms that would be inherited by all derivative works, we hope to attract as many eyeballs as we possibly can to use and improve the kernel. Although many of the drugs in the kernel are proprietary under diverse types of rights, we believe that the existence of public domain pairs of targets and compounds will reduce the royalties that patent owners can charge and sponsors must pay. This should decrease the large sums of money governments and

Table 1 TDI kernel genomes

Organism ^a	Transcripts ^b	Modeled targets ^c	Similar ^d	Exact ^e
<i>Cryptosporidium hominis</i>	3,886	666	20	13
<i>Cryptosporidium parvum</i>	3,806	742	24	13
<i>Leishmania major</i>	8,274	1,409	43	20
<i>Mycobacterium leprae</i>	1,605	893	25	6
<i>Mycobacterium tuberculosis</i>	3,991	1,608	30	10
<i>Plasmodium falciparum</i>	5,363	818	28	13
<i>Plasmodium vivax</i>	5,342	822	24	13
<i>Toxoplasma gondii</i>	7,793	300	13	6
<i>Trypanosoma cruzi</i>	19,607	3,070	51	28
<i>Trypanosoma brucei</i>	9,210	1,386	39	21
Total	68,877	11,714	297	143

^aOrganisms in bold are included in the World Health Organization (Geneva) Tropical Disease portfolio. ^bNumber of transcripts in each genome. ^cNumber of targets with at least one domain accurately modeled (that is, MDDPPE quality score of at least 1.0). ^dNumber of modeled targets with at least one predicted binding site for a molecule with a Tanimoto score¹⁰ of at least 0.9 to a drug in DrugBank¹¹. ^eNumber of modeled targets with at least one predicted binding site for a molecule in DrugBank.

320

VOLUME 27 NUMBER 4 APRIL 2009 NATURE BIOTECHNOLOGY

OPEN ACCESS Freely available online



A Kernel for Open Source Drug Discovery in Tropical Diseases

Leticia Orti^{1,2}, Rodrigo J. Carbajo², Ursula Pieper³, Narayanan Eswar^{3*}, Stephen M. Maurer⁴, Arti K. Rai⁵, Ginger Taylor⁶, Matthew H. Todd⁷, Antonio Pineda-Lucena², Andrej Sali^{3*}, Marc A. Marti-Renom^{1*}

1 Structural Genomics Unit, Bioinformatics and Genomics Department, Centro de Investigación Príncipe Felipe, Valencia, Spain, **2** Structural Biology Laboratory, Medical Chemistry Department, Centro de Investigación Príncipe Felipe, Valencia, Spain, **3** Department of Bioengineering and Therapeutic Sciences, Department of Pharmaceutical Chemistry, and California Institute for Quantitative Biosciences, University of California San Francisco, San Francisco, California, United States of America, **4** Gould School of Law, University of Southern California, Los Angeles, California, United States of America, **5** School of Law, Duke University, Durham, North Carolina, United States of America, **6** The Synaptic Leap, San Ramon, California, United States of America, **7** School of Chemistry, University of Sydney, Sydney, New South Wales, Australia

Abstract

Background: Conventional patent-based drug development incentives work badly for the developing world, where commercial markets are usually small to non-existent. For this reason, the past decade has seen extensive experimentation with alternative R&D institutions ranging from private-public partnerships to development prizes. Despite extensive discussion, however, one of the most promising avenues—open source drug discovery—has remained elusive. We argue that the stumbling block has been the absence of a critical mass of preexisting work that volunteers can improve through a series of granular contributions. Historically, open source software collaborations have almost never succeeded without such 'kernels'.

Methodology/Principal Findings: Here, we use a computational pipeline for: (i) comparative structure modeling of target proteins, (ii) predicting the localization of ligand binding sites on their surfaces, and (iii) assessing the similarity of the predicted ligands to known drugs. Our kernel currently contains 143 and 297 protein targets from ten pathogen genomes that are predicted to bind a known drug or a molecule similar to a known drug, respectively. The kernel provides a source of potential drug targets and drug candidates around which an online open source community can nucleate. Using NMR spectroscopy, we have experimentally tested our predictions for two of these targets, confirming one and invalidating the other.

Conclusions/Significance: The TDI kernel, which is being offered under the Creative Commons attribution share-alike license for free and unrestricted use, can be accessed on the World Wide Web at <http://www.tropicaldisease.org>. We hope that the kernel will facilitate collaborative efforts towards the discovery of new drugs against parasites that cause tropical diseases.

Citation: Orti L, Carbajo RJ, Pieper U, Eswar N, Maurer SM, et al. (2009) A Kernel for Open Source Drug Discovery in Tropical Diseases. *PLoS Negl Trop Dis* 3(4): e418. doi:10.1371/journal.pntd.0000418

Editor: Timothy G. Geary, McGill University, Canada

Received: December 29, 2008; **Accepted:** March 23, 2009; **Published:** April 21, 2009

Copyright: © 2009 Orti et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: MAM-R acknowledges the support from a Spanish Ministerio de Educación y Ciencia grant (B02007/66670). AS acknowledges the support from the Sandler Family Supporting Foundation and the National Institutes of Health (R01 GM47472, U54 GM074945, P01 AG035707, and P01 GM071790). AP-L acknowledges the support from a Spanish Ministerio de Ciencia e Innovación grant (SAF2008-01845). RJC acknowledges the support from the Ramon y Cajal Program of the Spanish Ministerio de Educación y Ciencia. We are also grateful for computer hardware gifts to AS from Ron Conway, Mike Homer, Intel, IBM, Hewlett-Packard, and NetApp. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: sali@lab.org (AS); mmarti@cpf.es (MAM-R)

† Current address: DuPont Knowledge Center, Hyderabad, India

Introduction

There is a lack of high-quality protein drug targets and drug leads for neglected diseases [1,2]. Fortunately, many genomes of organisms that cause tropical diseases have already been sequenced and published. Therefore, we are now in a position to leverage this information by identifying potential protein targets for drug discovery. Atomic-resolution structures can facilitate this task. In the absence of an experimentally determined structure, comparative modeling can provide useful models for sequences that are detectably related to known protein structures [3,4]. Approximately half of known protein sequences contain domains that can be currently predicted by comparative modeling [5,6]. This coverage

will increase as the number of experimentally determined structures grows and modeling software improves. A protein model can facilitate at least four important tasks in the early stages of drug discovery [7]: prioritizing protein targets for drug discovery [8], identifying binding sites for small molecules [9,10], suggesting drug leads [11,12], and optimizing these leads [13–15].

Here, we address the first three tasks by assembling our computer programs into a software pipeline that automatically and on large-scale predicts protein structures, their ligand binding sites, and known drugs that interact with them. As a proof of principle, we applied the pipeline to the genomes of ten organisms that cause tropical diseases ('target genomes'). We also experimentally tested two predicted drug-target interactions using Nuclear Magnetic

www.plosntds.org

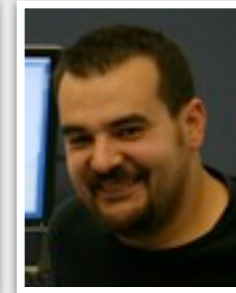
1

April 2009 | Volume 3 | Issue 4 | e418

Acknowledgments

<http://tropicaldisease.org>

<http://thesynapticleap.org>



COMPARATIVE MODELING

Andrej Sali

M. S. Madhusudhan

Narayanan Eswar

Min-Yi Shen

Ursula Pieper

Ben Webb

Maya Topf (Birbeck College)

MODEL ASSESSMENT

Francisco Melo (CU)

Alejandro Panjkovich (CU)

NMR

Antonio Pineda-Lucena

Leticia Ortí

Rodrigo J. Carbajo

MODEL ASSESSMENT

David Eramian

Min-Yi Shen

Damien Devos

MAMMOTH

Angel R. Ortiz

FUNCTIONAL ANNOTATION

Andrea Rossi (Rinat-Pfizer)

Fred Davis (Janelia Fram)

FUNCTIONAL ANNOTATION

Fatima Al-Shahrour

Joaquin Dopazo

FUNDING

Prince Felipe Research Center

Ministerio de Educación y Ciencia

STREP UE Grant

Marie Curie Reintegration Grant

BIOLOGY

Jeff Friedman (RU)

James Hudsped (RU)

Partho Ghosh (UCSD)

Alvaro Monteiro (Cornell U)

Stephen Krilis (St. George H)

Tropical Disease Initiative

Stephen Maurer (UC Berkeley)

Arti Rai (Duke U)

Andrej Sali (UCSF)

Ginger Taylor (TSL)

Matthew Todd (U Sydney)

CCPR Functional Proteomics

Patsy Babbitt (UCSF)

Fred Cohen (UCSF)

Ken Dill (UCSF)

Tom Ferrin (UCSF)

John Irwin (UCSF)

Matt Jacobson (UCSF)

Tack Kuntz (UCSF)

Andrej Sali (UCSF)

Brian Shoichet (UCSF)

Chris Voigt (UCSF)

EVA

Burkhard Rost (Columbia U)

Alfonso Valencia (CNB/UAM)

CAMP

Xavier Aviles (UAB)

Hans-Peter Nester (SANOFI)

Ernst Meinjohanns (ARPIDA)

Boris Turk (IJS)

Markus Gruetter (UE)

Matthias Wilmanns (EMBL)

Wolfram Bode (MPG)