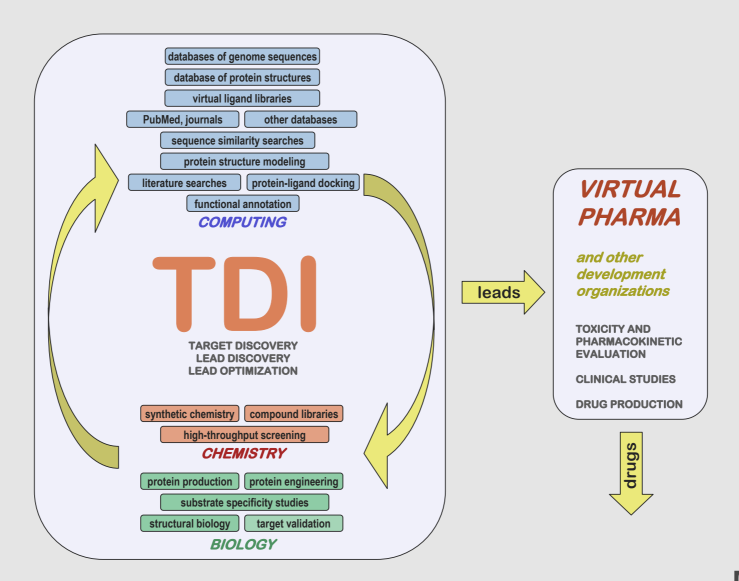
## A "kernel" for the Tropical Disease Initiative

An open source approach to drug discovery



### Marc A. Marti-Renom

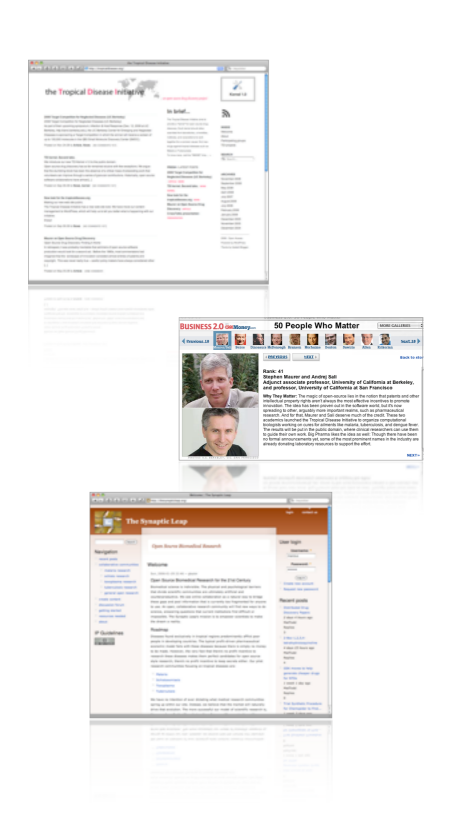
http://sgu.bioinfo.cipf.es

Structural Genomics Unit Bioinformatics Department

Prince Felipe Resarch Center (CIPF), Valencia, Spain



# TU a sterystory



### 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

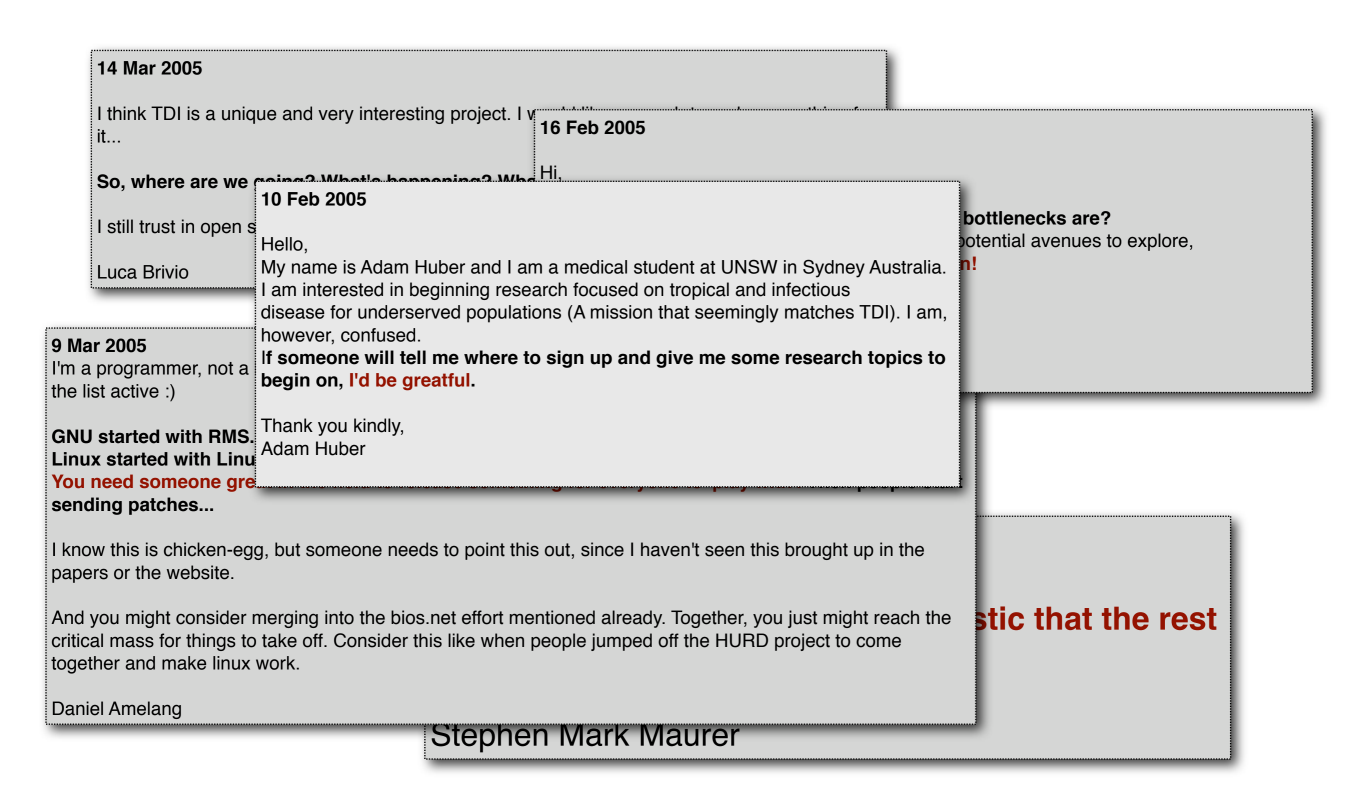
- .Maurer and Sali 41th in "50 Who Matter"
- .TSL web site http://TheSynapticLeap.org

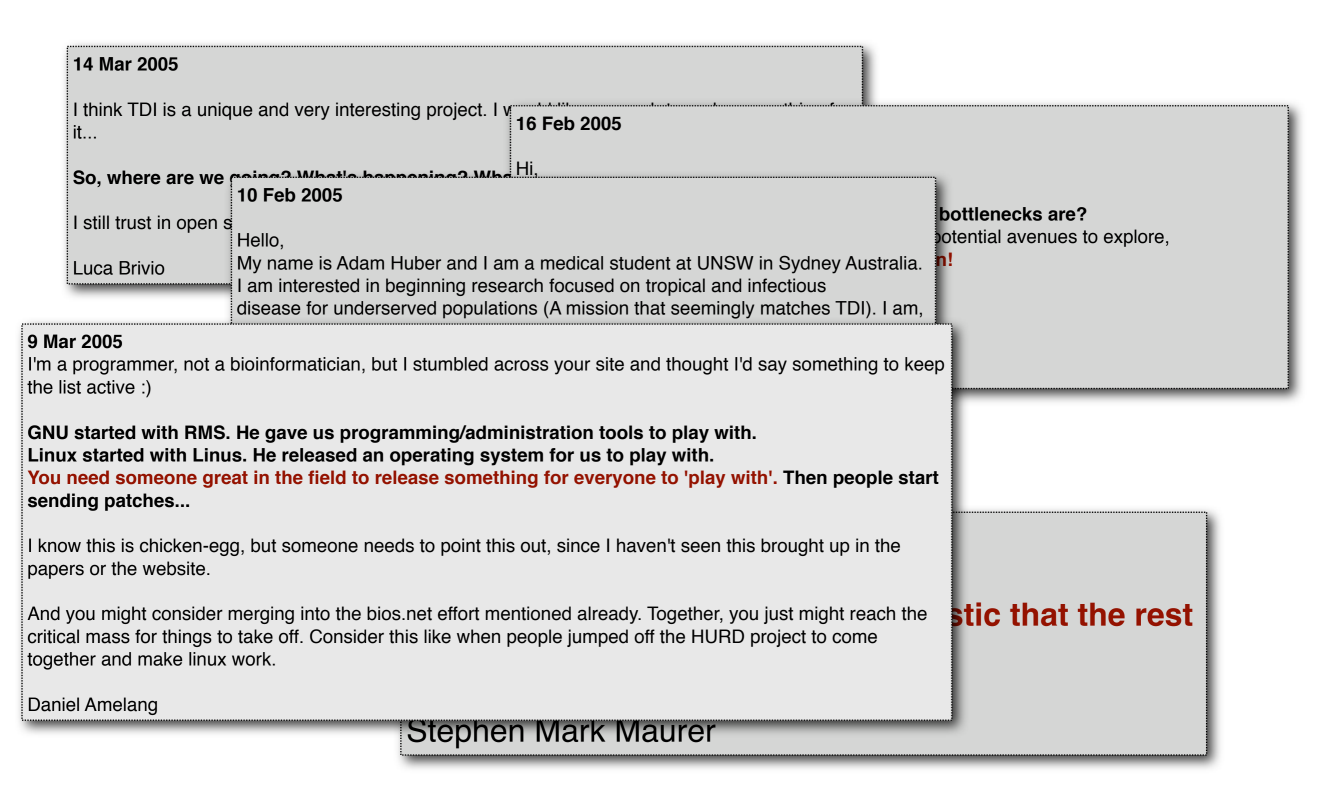
### 2008

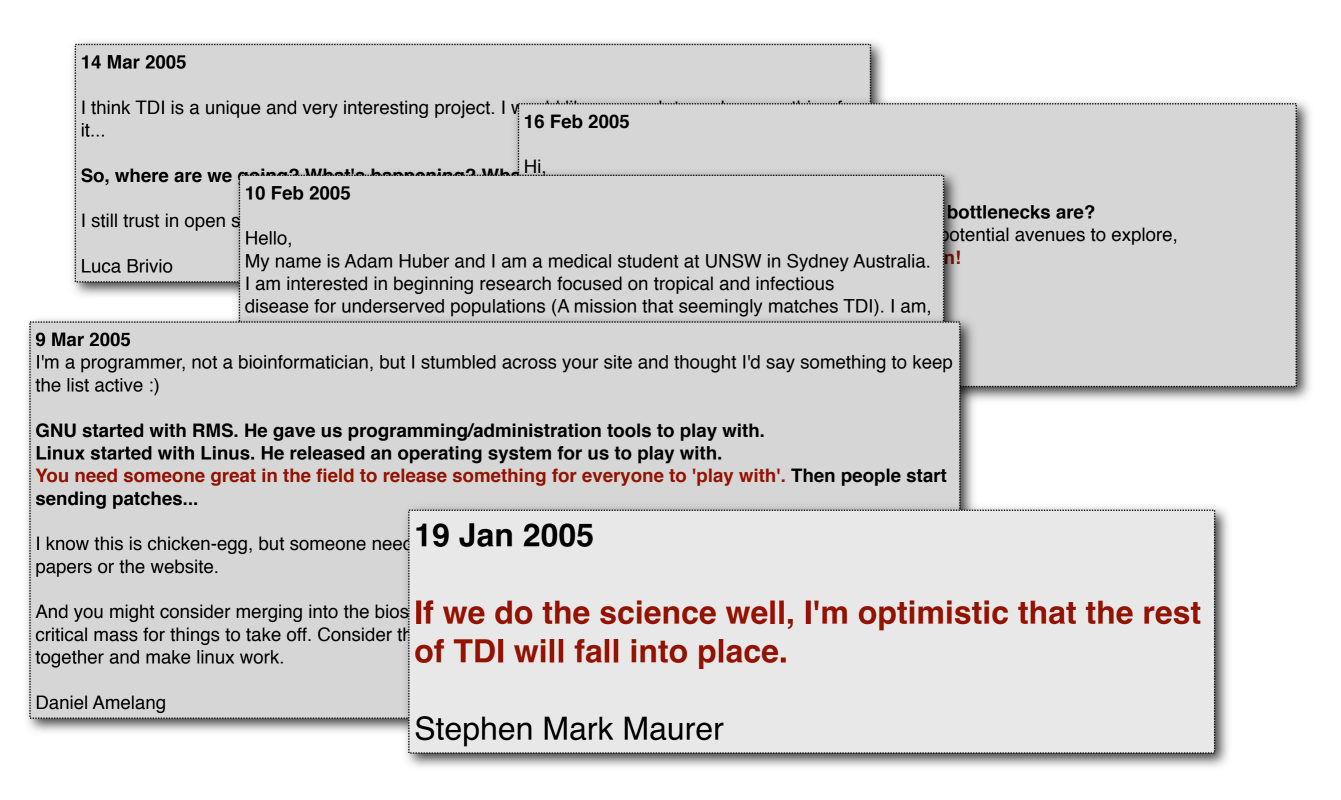
.TDI kernel http://TropicalDisease.org/kernel



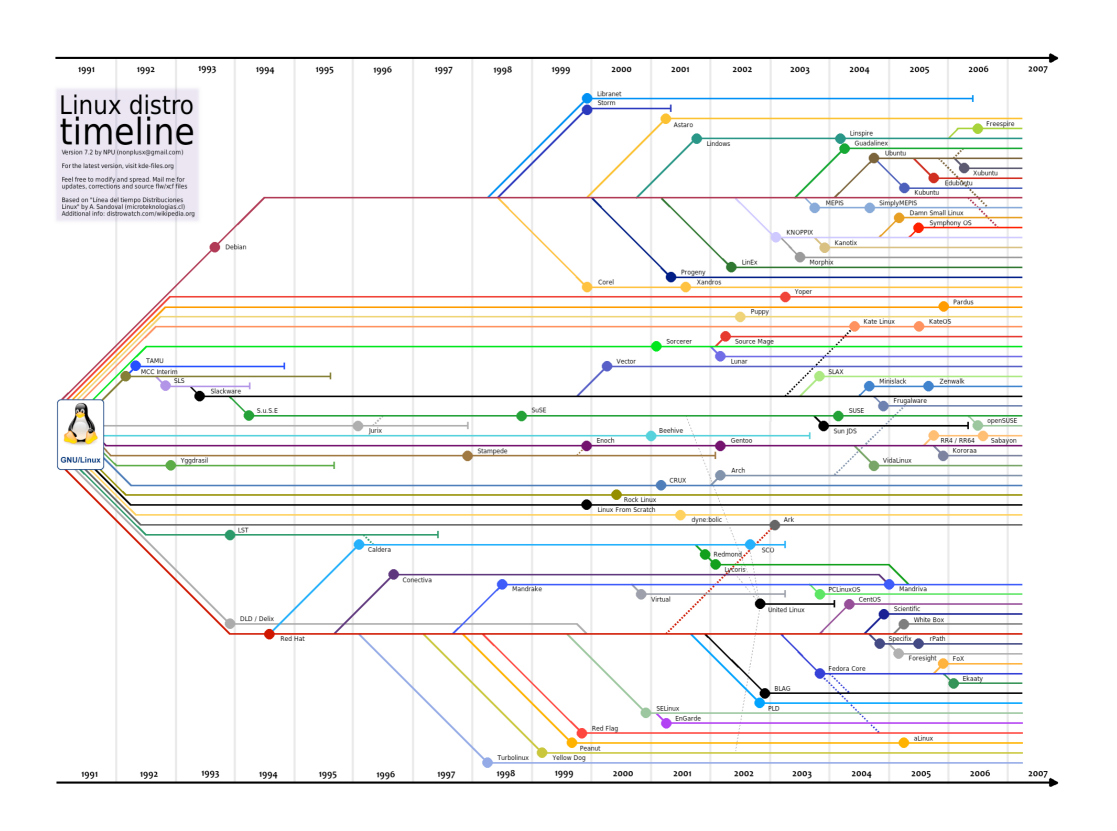








# **Open Source without a Kernel?**



## Is it possible? ...

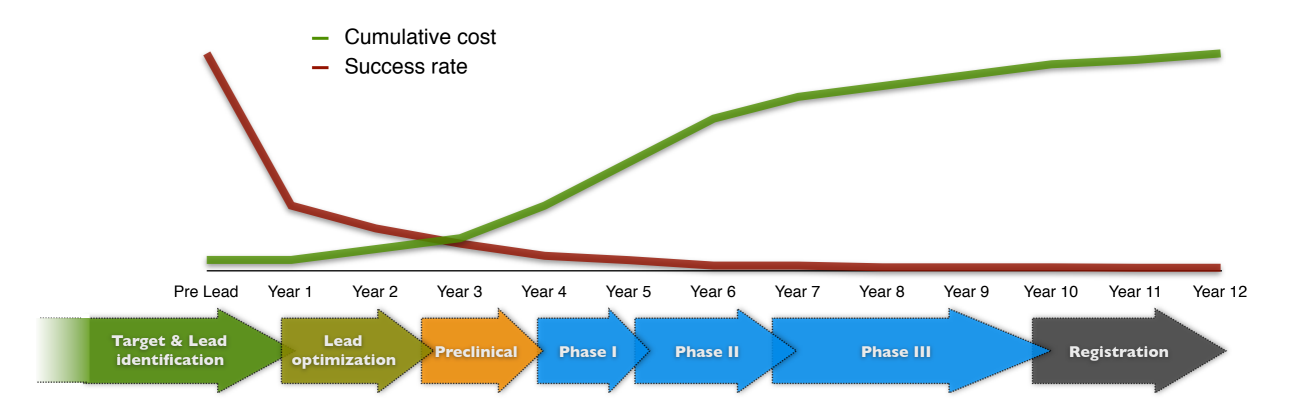
- 1. In silico drug discovery
- 2. Chemistry
- 3. Stem cell lines

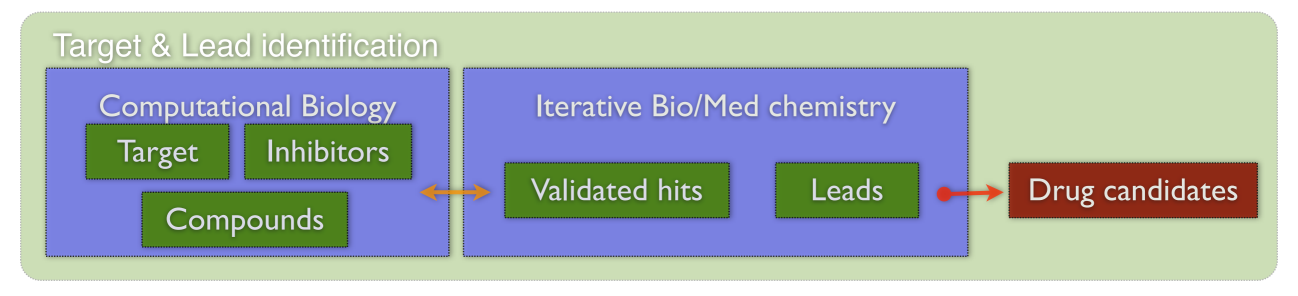
**NEXT STEPS** 

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

**AMBITIOUS GOALS** 

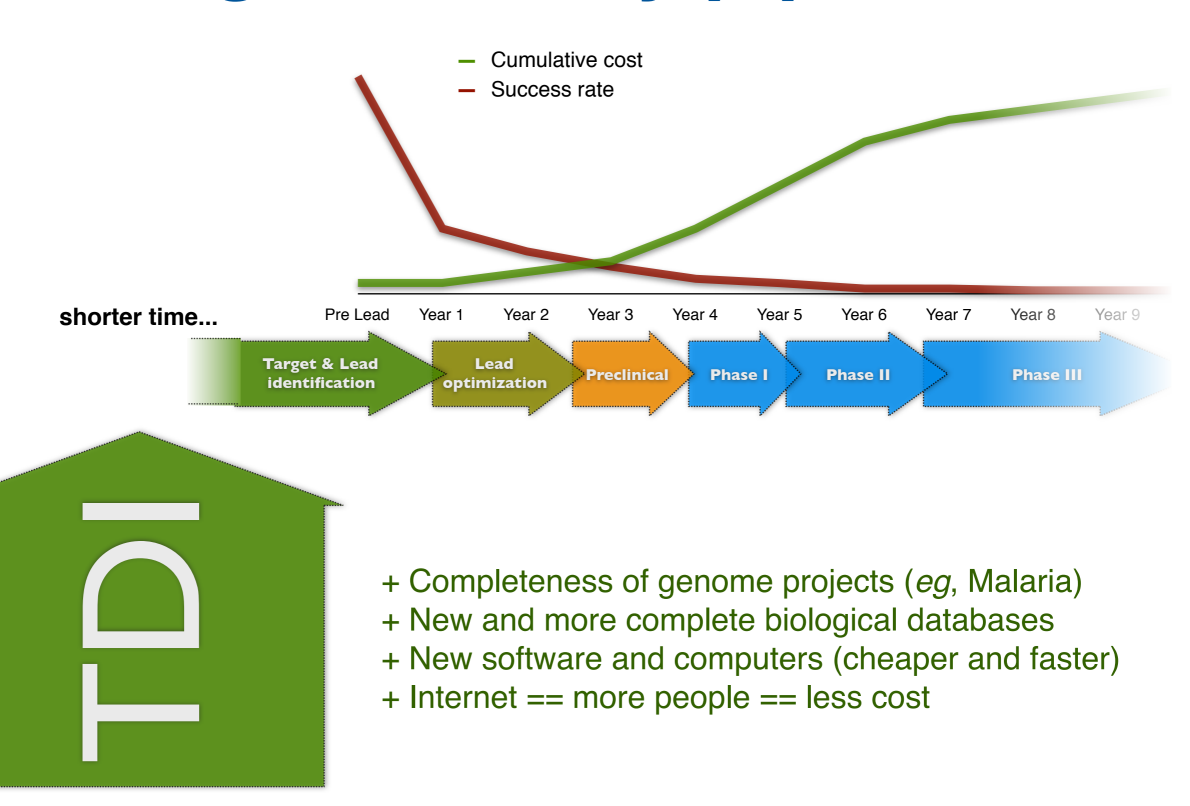
# Drug Discovery pipeline



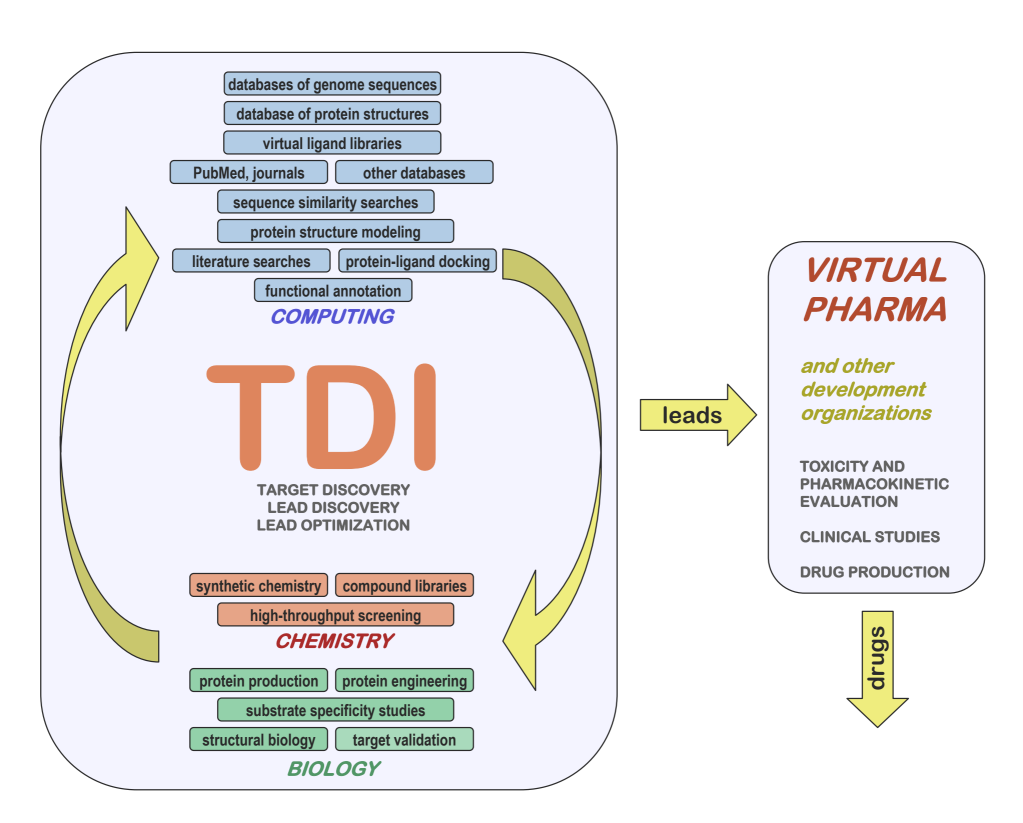


Adapted from: - Nwaka & Ridley. (2003) *Nature Reviews. Drug Discovery.* **2**:919 - Austin, Brady, Insel & collins. (2004) *Science.* **306**:1138

# Drug Discovery pipeline

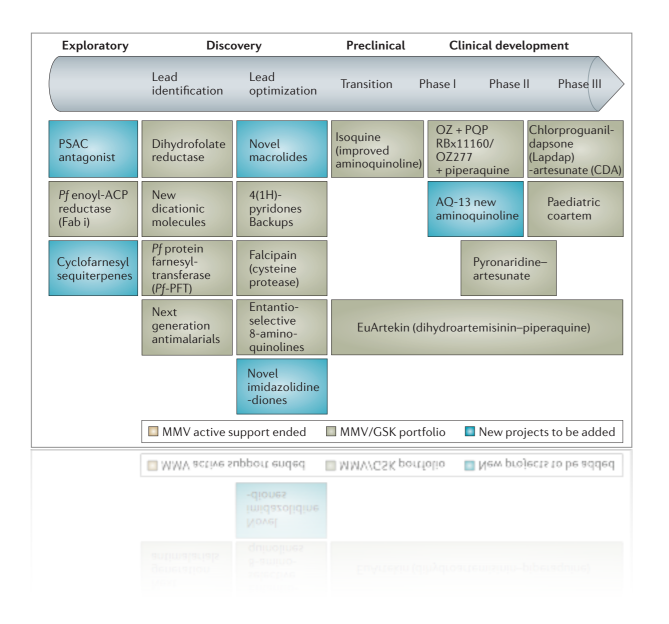


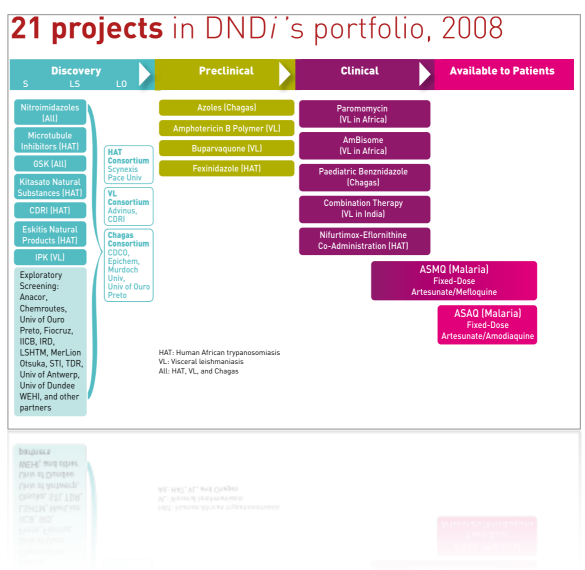
# **TDI flowchart**



# Non-Profit organizations

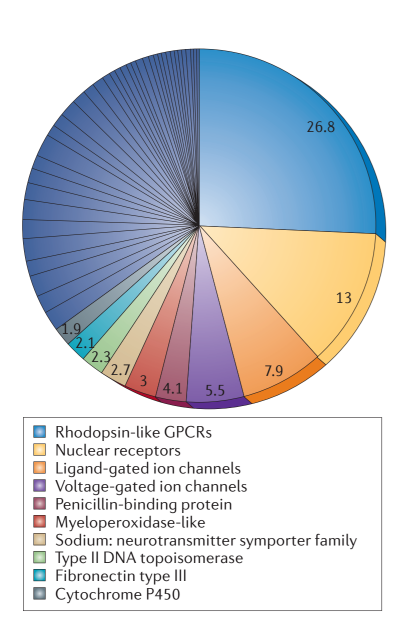
Open-Source + Out-Source = low cost business model





Munos (2006) Nature Reviews. Drug Discovery.

# Number of (new) targets?



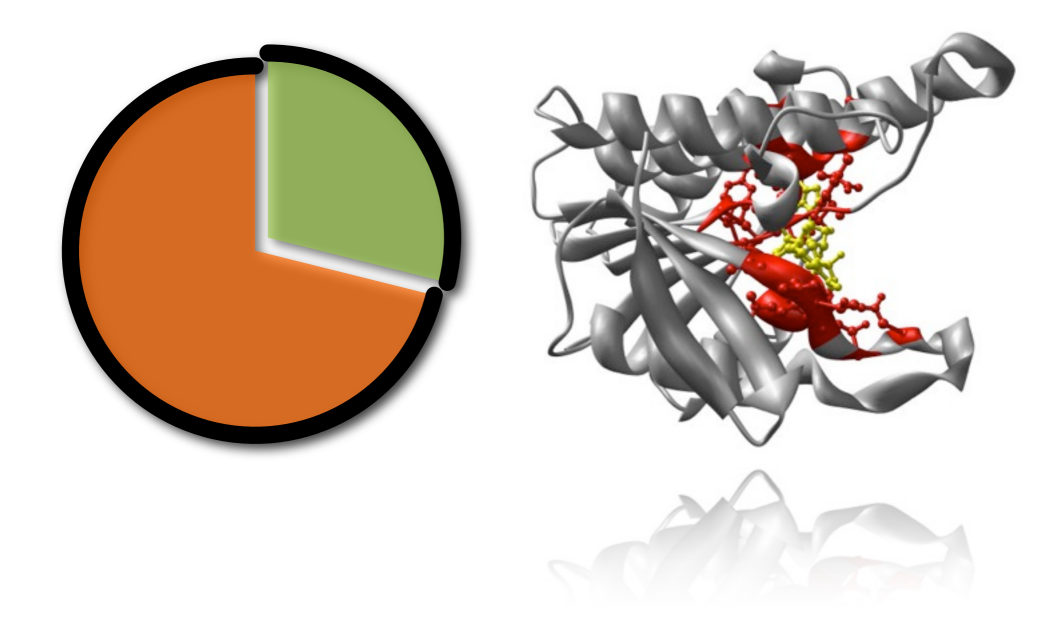
"... of 361 new molecular entities approved by the FDA between 1989 and 2000, 76% targeted a precedented drugged domain and only 6% targeted a previously undrugged domain ..."

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.











# 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

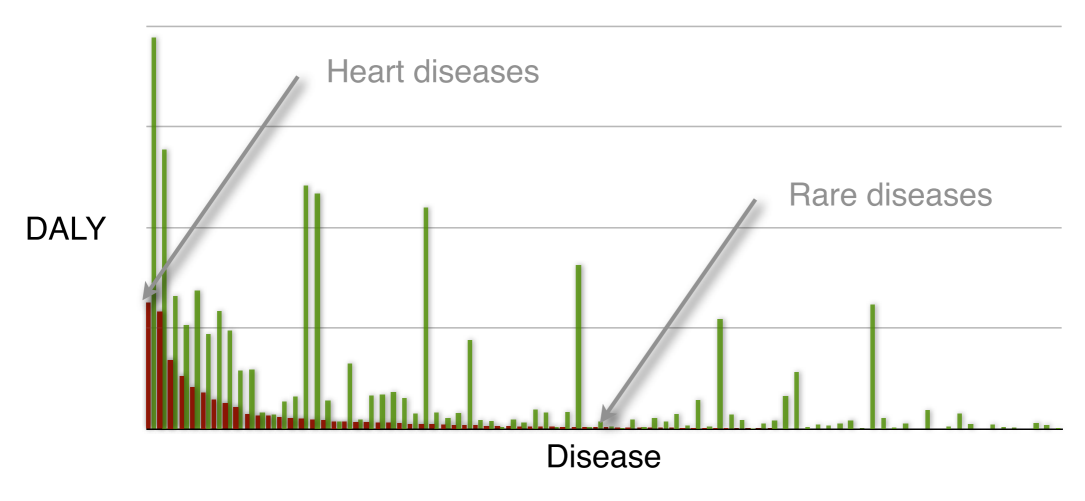
DALY - Disability adjusted life years

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, <u>World Health Report 2004</u>
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
Tetanus	7,074
Lymphatic filariasis*	5,777
Syphilis	4,200
Trachoma	2,329
Leishmaniasis*	2,090
Ascariasis	1,817
Schistosomiasis*	1,702
Trypanosomiasis*	1,525

Trichuriasis	1,006
Japanese encephalitis	709
Chagas Disease*	667
Dengue*	616
Onchocerciasis*	484
Leprosy*	199
Diphtheria	185
Poliomyelitise	151
Hookworm disease	59

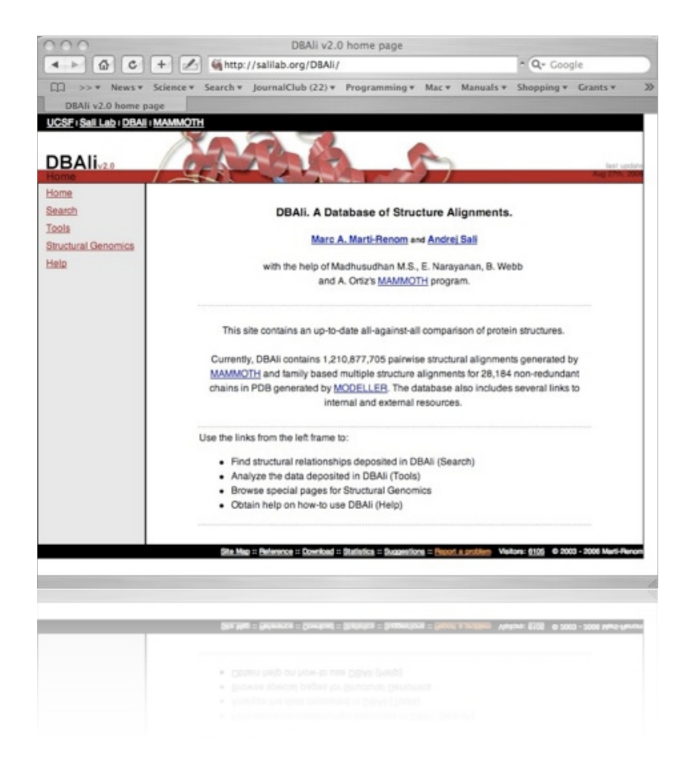
Disease data taken from WHO, World Health Report 2004

DALY - Disability adjusted life year in 1000's.

<sup>\*</sup> Officially listed in the WHO Tropical Disease Research disease portfolio.

## DBAliv2.0 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

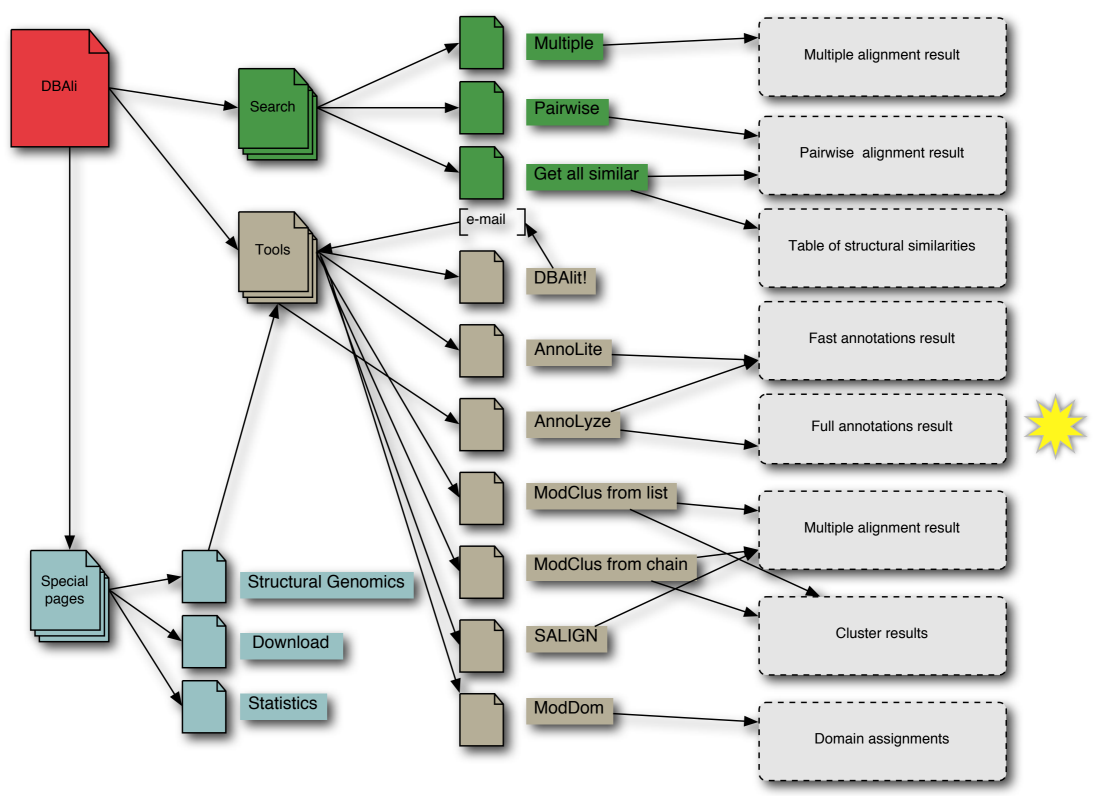
- ✓ VERY FAST!!!
- √ Good scoring system with significance

Ortiz AR, (2002) Protein Sci. 11 pp2606

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

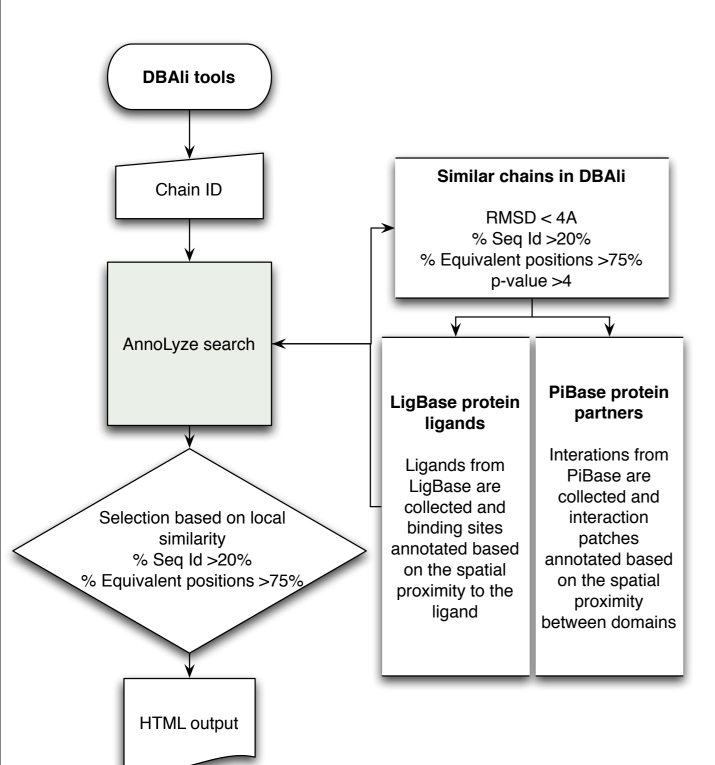
## DBAliv2.0 database

http://www.dbali.org



Marti-Renom et al. BMC Bioinformatics (2007) Volume 8. Suppl S4

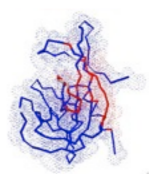
# Method



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
8OG	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

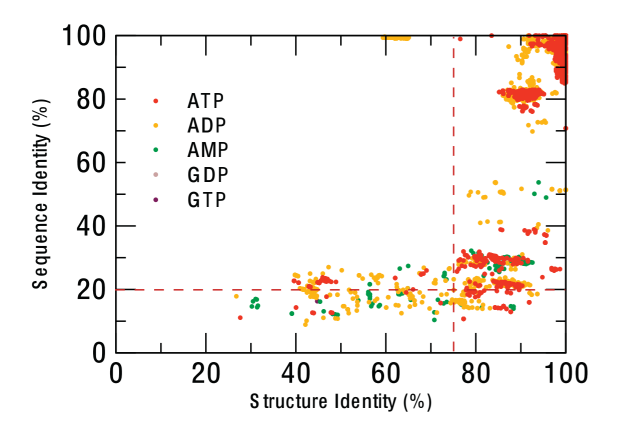


herited p	artners:1		
Partner	Av. binding site seq. id.	Av. residue conservation	Residues in predicted binding site (size proportional to the local conservation)
1.113.1.1	23.68	0.948	19 20 50 51 52 53 54 55 56 57 58 77 78 79 80 81 82 83 84 85 93 95 97 99 134 135 138 142
			145

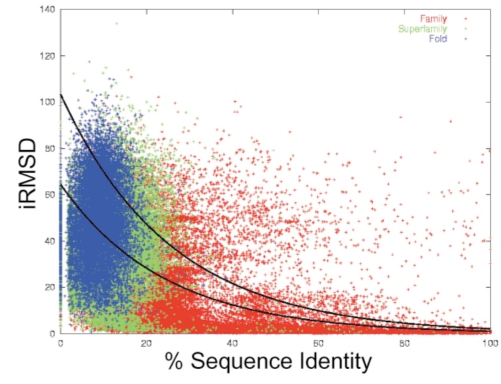


# Scoring function

## 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 3A, superimposing more than 75% of their Cα atoms, result in a sequence alignment with more than 30% identity, and have a length difference inferior to 50aa

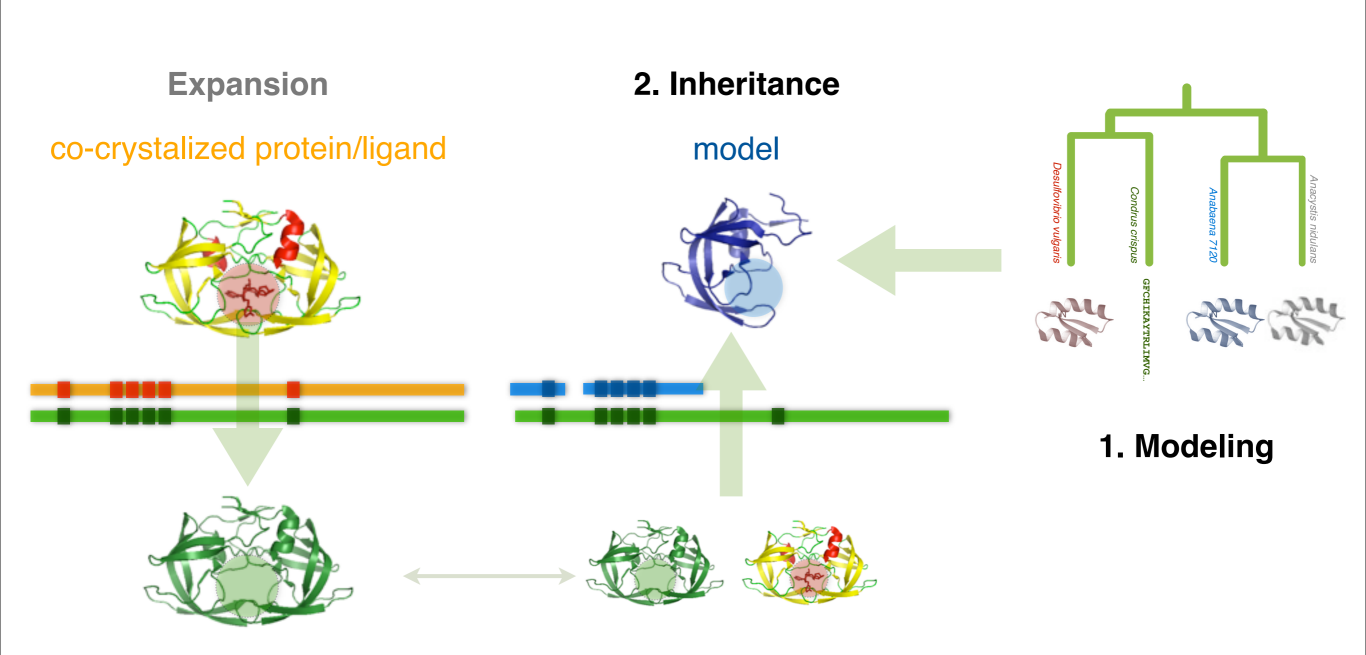
# Sensitivity .vs. Precision

	Optimal cut-off		Precision (%)	
Ligands	30%	71.9	13.7	

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

~90-95% of residues correctly predicted

# Comparative docking



24

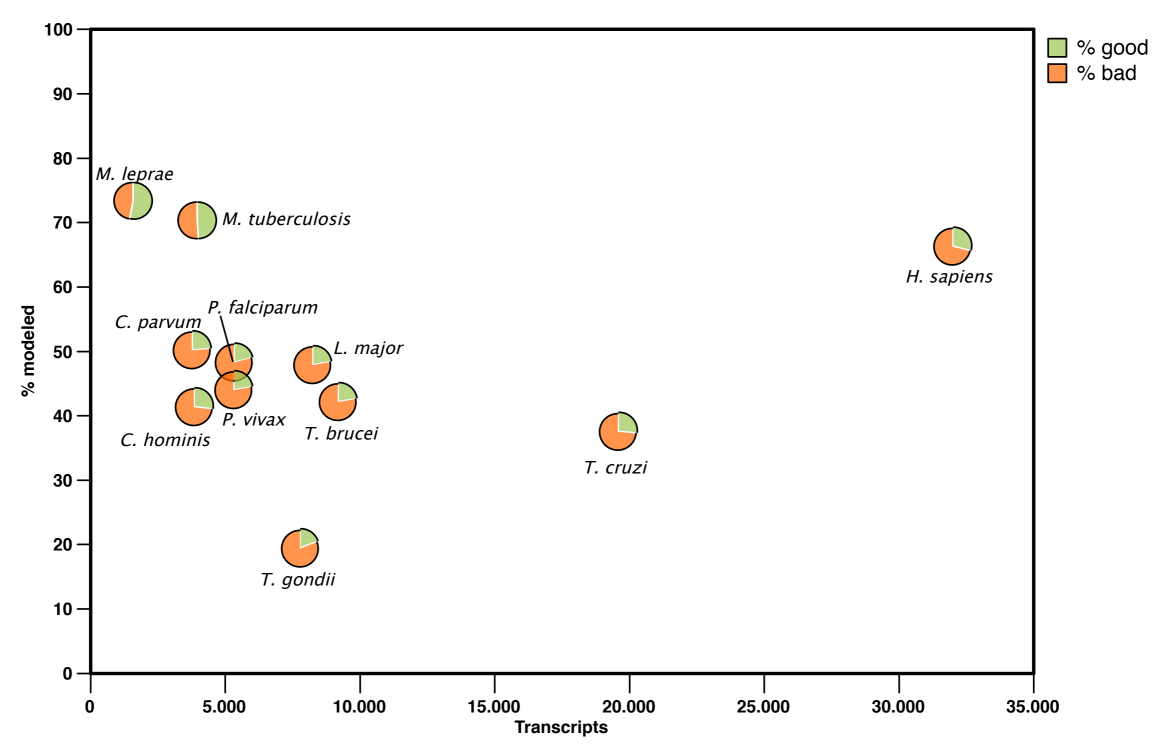
template

crystalized

protein

# Modeling Genomes

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



# Summary table

models with inherited ligands

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

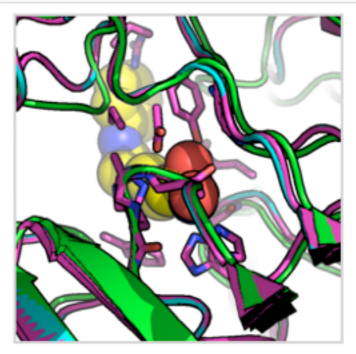
	Transcripts	Modeled targets	Selected models	Inherited ligands	Similar to a drug	Drugs
C. hominis	3,886	1,614	666	197	20	13
C. parvum	3,806	1,918	742	232	24	13
L. major	8,274	3,975	1,409	478	43	20
M. leprae	1,605	1,178	893	310	25	6
M. tuberculosis	3,991	2,808	1,608	365	30	10
P. falciparum	5,363	2,599	818	284	28	13
P. vivax	5,342	2,359	822	268	24	13
T. brucei	7,793	1,530	300	138	13	6
T. cruzi	19,607	7,390	3,070	769	51	28
T. gondii	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	EO	Template	666	Model	⇔	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

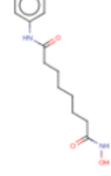


#### DB02546 Vorinostat

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

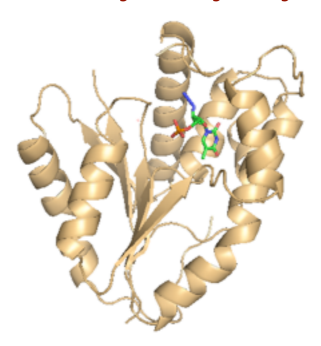
ANTIMICROBIAL AGENTS AND CHEMOTHERAPY, Apr. 2004, p. 1435–1436 0066-4804/04/\$08.00+0 DOI: 10.1128/AAC.48.4.1435–1436.2004 Copyright © 2004, American Society for Microbiology. All Rights Reserved.

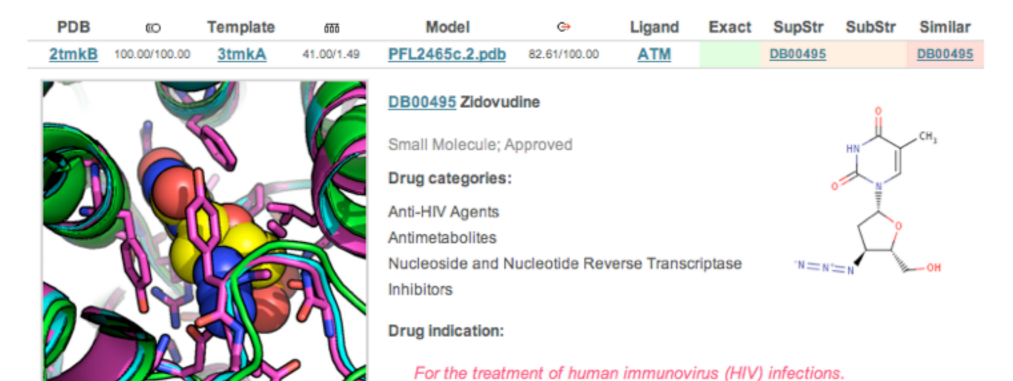
Vol. 48, No. 4

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

## P. falciparum tymidylate kinase + zidovudine

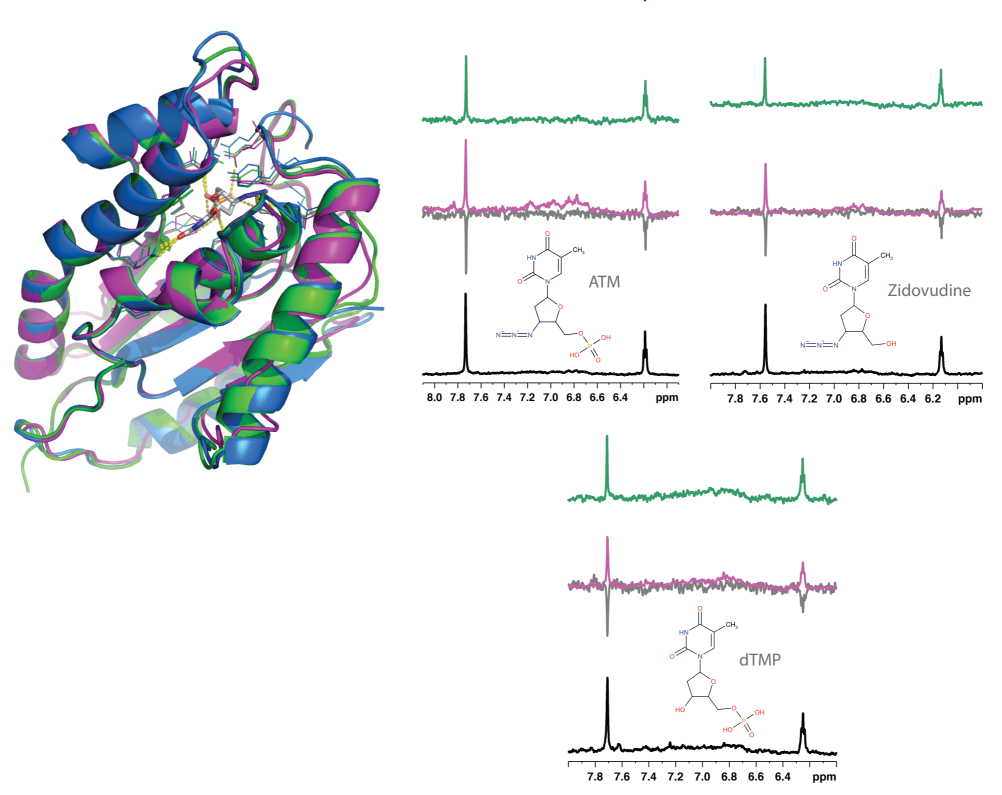
Template 3tmkA a yeast tymidylate kinase.





## P. falciparum thymidylate kinase + zidovudine

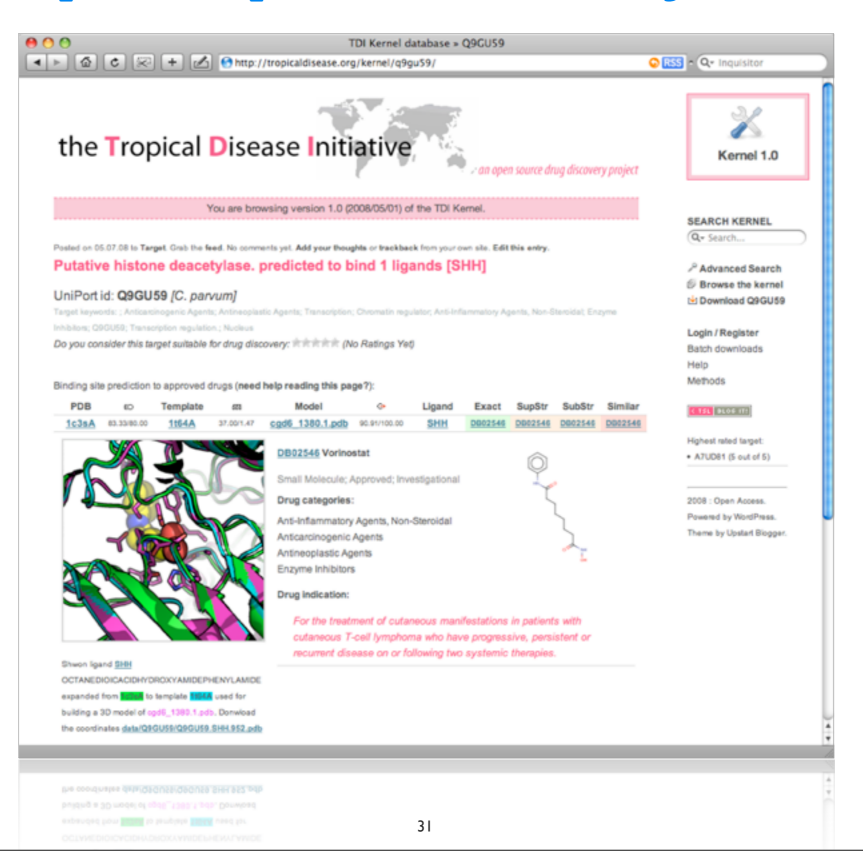
NMR Water-LOGSY and STD experiments



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

## TDI's kernel

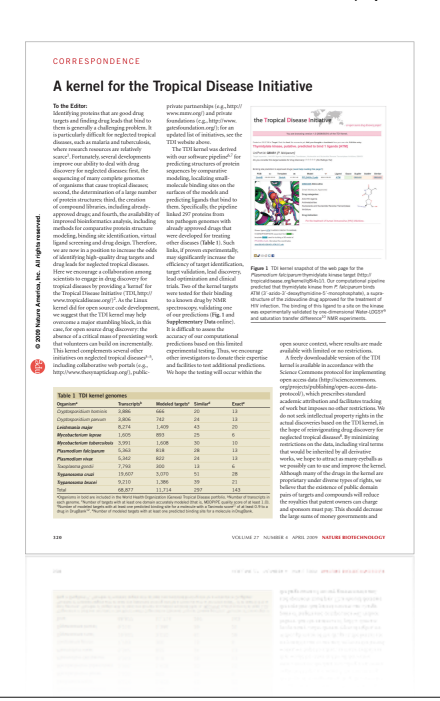
http://tropicaldisease.org/kernel



## TDI'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).



# Acknowledgments

http://tropicaldisease.org
http://thesynapticleap.org



**Andrej Sali** 

M. S. Madhusudhan

**Narayanan Eswar** 

Min-Yi Shen

**Ursula Pieper** 

Ben Webb

Maya Topf (Birbeck College)

### **MODEL ASSESSMENT**

David Eramian Min-Yi Shen Damien Devos

#### **FUNCTIONAL ANNOTATION**

Andrea Rossi (Rinat-Pfizer) Fred Davis (Janelia Fram)

#### **FUNDING**

Prince Felipe Research Center

Ministerio de Educación y Ciencia

STREP UE Grant

Marie Curie Reintegration Grant

### **MODEL ASSESSMENT**

Francisco Melo (CU) Alejandro Panjkovich (CU)

#### **NMR**

Antonio Pineda-Lucena Leticia Ortí Rodrigo J. Carbajo

MAMMOTH Angel R. Ortiz

FUNCTIONAL ANNOTATION
Fatima Al-Shahrour
Joaquin Dopazo

#### **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)

#### FVA

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)