The Tropical Disease Initiative

An open source approach to drug discovery



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



Tuesday, July 12, 2011

TD asterystory



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

2009

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

14 Mar 2005	14 Mar 2005						
I think TDI is a uni it	que and very interesting project. I would like so much to make something for						
So, where are we	going? What's happening? What can we do?						
	I still trust in open source drug discovery. :-))						
Luca Brivio	ram merested in beginning rese						
9 Mar 2005 I'm a programmer, not a the list active :)	disease for underserved populatic Regards, however, confused. If someone will tell me where to begin on, I'd be greatful.						
GNU started with RMS Linux started with Line You need someone gr	u Adam Huber u						
sending patches							
I know this is chicken-eq papers or the website.	gg, but someone needs to point this out, since I haven't seen this brought up in	n the					
critical mass for things t	nd you might consider merging into the bios.net effort mentioned already. Together, you just might reach the ritical mass for things to take off. Consider this like when people jumped off the HURD project to come ogether and make linux work.						
Daniel Amelang							
	Stephen Mark Maurer						

	14 Mar 2005		
	I think TDI is a uniq it	ue and very interesting project. I v	16 Feb 2005
	So, where are we	going? What's happening? Wha	Hi,
		ource drug discovery. :-))	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!
9 Mar 2005 I'm a programmer, not a		disease for underserved populati however, confused. If someone will tell me where te	Regards, Jacob Lester
GNU started with RMS. Linux started with Linu You need someone gre		Adam Huber	
l kno Dape	ers or the website.		s out, since I haven't seen this brought up in the
And you might consider merging into the bios.net effort me critical mass for things to take off. Consider this like when together and make linux work.			
Dani	el Amelang	Stonho	n Mark Mauror
		Stephie	n Mark Maurer

	I am interested in beginning research focused on tropical and infectious					
	it	16 Feb 2005				
	I still trust in open s Luca Brivio r 2005	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.	bottlenecks are? ootential avenues to explore, n!			
the li GNU Linu You	st active :) started with RMS. x started with Linu need someone gre	begin on, I'd be greatful. Thank you kindly, Adam Huber				
l kno pape And critic	w this is chicken-egers or the website. you might consider i	nerging into the bios.net effort mentioned already. Together, you just might reach the take off. Consider this like when people jumped off the HURD project to come	stic that the rest			
Dani	el Amelang	Stonbon Mark Mauror				
		Stephen Mark Maurer				

	14 Mar 2005							
	I think TDI is a unic it	que and very interesti	ng project. I v	16 Feb 2005				
	So, where are we I still trust in open s Luca Brivio	Hello, My name is Adam H I am interested in be	luber and I ar	m a medical student arch focused on trop	at UNSW in Sydney Au pical and infectious seemingly matches TDI)	oote Istralia. <mark>n!</mark>	Itlenecks are? ential avenues to explore,	
l'm a	r 2005 programmer, not a st active :)	bioinformatician, but	I stumbled ac	cross your site and t	hought I'd say somethin	ig to keep		
Linu <mark>You</mark>	x started with Linu	. He gave us progra is. He released an o eat in the field to rele	perating sys	stem for us to play		ple start		
	w this is chicken-eg ers or the website.	g, but someone need	ls to point this	s out, since I haven'	t seen this brought up in	n the		
critic	, ,	merging into the bios take off. Consider th work.		tic that the rest				
Dani	el Amelang		Stonho	n Mark Mai	Iror			
			Olehile	n Mark Mau				

14 Mar 2005					
I think TDI is a unique and very interest it	ing project. I v 16 Feb 2005				
I am interested in b	Hi. Huber and I am a medical student at UNSW in Sydney Australia. eginning research focused on tropical and infectious erved populations (A mission that seemingly matches TDI). I am,				
I'm a programmer, not a bioinformatician, but the list active :) GNU started with RMS. He gave us progra Linux started with Linus. He released an o					
I know this is chicken-egg, but someone need papers or the website.	19 Jan 2005				
And you might consider merging into the bios critical mass for things to take off. Consider the together and make linux work.	If we do the science well, I'm optimistic that the rest of TDI will fall into place.				
Daniel Amelang	Stephen Mark Maurer				

Open Source without a Kernel?



8

TDI flowchart



Drug Discovery pipeline



Non-Profit organizations

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

11

Exploratory	Disco	overy	Preclinical	lopment	
	Lead identification	Lead optimization	Transition Pl	hase I Phase	II Phase III
PSAC antagonist	Dihydrofolate reductase	Novel macrolides	lsoquine (improved aminoquinoline)	OZ + PQP RBx11160/ OZ277 + piperaquine	Chlorproguanil- dapsone (Lapdap) -artesunate (CDA)
<i>Pf</i> enoyl-ACP reductase (Fab i)	New dicationic molecules	4(1H)- pyridones Backups		AQ-13 new aminoquinoline	Paediatric coartem
Cyclofarnesyl sequiterpenes	Pf protein farnesyl- transferase (Pf-PFT)	Falcipain (cysteine protease)		Pyronarid artesunat	
	Next generation antimalarials	Entantio- selective 8-amino- quinolines	EuArtekin (dil	piperaquine)	
		Novel imidazolidine -diones			
	MMV active s	upport ended	MMV/GSK portfo	olio 🔲 New pro	jects to be added
	MMV active s	upport ended	MMV/GSK portfe	olio 🛛 🔲 New pro	jects to be added
		Novel imidazolidine -diones			



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 of Tropical Diseases



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.

14

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.

14

"Unprofitable" Diseases and Global DALY (in 1000's)

Malaria*	46,486	Trichuriasis	I,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	Poliomyelitise	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.

Comparative docking



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
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	١,409	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

http://tropicaldisease.org Creative Commons (no viral!)

L. major Histone deacetylase 2 + Vorinostat

Template 1t64A a human HDAC8 protein.



PDB	60	Template	000	Model	G	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.



PDB	ŝ	Template	000	Model	¢	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
	9	Z		DB00495 Zidovud	ine				ĥ	
		1		Small Molecule; Ap	oproved				HN H	СН3
				Drug categories:						
				Anti-HIV Agents						
~				Antimetabolites						
		100		Nucleoside and Nu	eoside and Nucleotide Reverse Transcriptase					
		TR		Inhibitors						
07				Drug indication:						
		1 Y		For the treatm	ent of huma	n immunovi	rus (HIV)) infection	s.	

P. falciparum thymidylate kinase + zidovudine

NMR Water-LOGSY and STD experiments



22

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

TDI's kernel

http://tropicaldisease.org/kernel



TDI reading... http://tropicaldisease.org

The Economist (2004) http://www.economist.com/node/2724420



Maurer et al. PLoS Medicine (2004) 1:e56

Neglected Diseases

Finding Cures for Tropical Diseases: Is Open Source an Answer? Stephen M. Maurer*, Arti Rai, Andrej Sali

nly about 1% of newly developed drugs are for University of the second secon dengue fever, and leishmainsis [1]. While patent incentives and commercial pharmaceutical houses have made Western health care the envy of the world, the commercial model only works if companies can sell enough patented products to cover their research and development (R&D) products to cover their research and development (R&D) costs. The model fails in the developing world, where few patients can afford to pay patented prices for drugs. It is easy (and correct) to say that Western governments could solve this problem by paying existing institutions to focus on curres for tropical diseases. But sadly, there is not enough political will for this to happen. In any case, grant

and patent incentives were never designed with tropical incentives were never designed with tropical diseases in mind. Two main kinds of proposals have been suggested for tackling the problem. The first is to ask sponsors governments and charities-to subsidize developing governments and charites—to subsidize developing-country purchases at a guaranteed price [23,4]. In the second approach, charities create nonprofit venture-capital firms ("Virtual Pharmas"), which look for promising drug candidates and then push drug development through contracts with corporate partners. In this article, we discuss Contracts with copy counts particular, in this and cut, we calcular the problems with these two approaches and suggest a third, "open source," approach to drug development, called the Tropical Diseases Initiative (TDI). We envise TDI as a decentralized, Web-based, community-wide effort where scientists from laboratories, universities, institutes, and corporations can work together for a common cause (see www.tropicaldisease.org).

Why Open Source?

The idea behind asking sponsors to subsidize developing The idea behind asking sponsors to subsidize developing country purchases at a guaranteed price is that this will prop up drug prices and restore incentives for developing new drugs [2,3,4]. In other words, it is a way of fixing the patent problem. However, subsidies have an important weakness: it is almost impossible to correctly determine how large the subsidy should be. In principle, the most cost-effective solution is to set a subsidy that just covers expected R&D costs. But how large is that? R&D costs are very poorly known, with published estimates ranging from \$100 to \$500 million per drug [5]. If the subsidy is set too low, companies cannot cover their R&D costs and postime with here. over their R&D costs and nothing will happen. Set the cover their K&D costs and nothing will happen. Set the subsidy too high, and the sponsor's cost skyrocket. To date, no sponsor has tried to implement these proposals. In the "Virtual Pharma" approach, governments and philanthropies fund teams to search out and subsidize the most promising private and academic research. Examples

include the Institute for One World Health (www.iowh.org), a not-for-profit pharmaceutical company funded mainly through private sources and the Gates Foundation, and the brough private sources and the Gates Foundation, and the Drugs for Neglected Diseases Initiative (www.dndi.org), a public sector not-for-profit organization designed to mobilize resources for R&D of new drugs for neglected diseases. Virtual Pharmas have clearly started to bear fruit, and are responsible for most candidate treatments for tropical are responsible to most cantinear treatments of tropical diseases currently under development. For example, the Drugs for Neglected Diseases Initiative has a portfolio of nine projects spread out across the drug development pipeline for the treatment of leishmaniasis, sleeping sickness, Chagas

Box 1. Possible Licenses for TDI Discoveries

Open access, freely available online

disease, and malaria [6]. But Virtual Pharmas face three important problems. The first is similar to the problem faced by subsidy proposals: guessing private-sector R&D costs. One needs to understand what a product costs in order to negotiate the best possible price-and guessing wrong is likely to be expensive. Second, Virtual Pharma's development pipelines will run dry without more upstream

Citation: Maurer SM, Rai A, Sali A (2004) Finding cures for tropical diseases: Is ope Copyright: © 2004 Maurer et al. This is an open-access article distribute terms of the Creative Commons Attribution License, which permit use, distribution, and reproduction in any medium, provided the origi properly cited. Creative Commons Attribution License, which permits u , and reproduction in any medium, provided the original

hen M. Maurer is in the Goldman School of Public Policy, University of Califor eley, California, United States of America. Arti Rai is in the School of Law, Duk versity, Durham, North Carolina, United States of America. Andrej Sali is in the partments of Biopharmaceutical Sciences and Pharmaceutical Chemistry and stitute for Ouan

Competing Interests: The authors declare that they hav To whom correspondence should be addressed. E-mail: mau DOI: 10.1371/journal.nmed.0010056

. PLoS Medicine | www.plosmedicine.org

Neglected Diseases section focuses attention either on a specific disea

ribes a novel strategy for approaching neglected health issues in general

December 2004 | Volume 1 | Issue 3 | e56

TDI's kernel

http://tropicaldisease.org/kernel

L. Orti et al., Nat Biotechnol 27, 320 (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 discovery for neglected diseases; scoond, the determination of a large number of organisms that cause tropical diseases; scoond, the determination of a large number of compound libraries, including already-approved drugs, and durith, the creation of compound libraries, including already-approved drugs and torth, the availability of them. Specifically, the pipeline diseases; nodeline indication of a large number of compound libraries, including already-approved drugs and corts, the availability of them. Specifically, the pipeline diseases; no deline indication of a large number of compound libraries, including already-approved drugs and torth, the availability of them. Specifically, the pipeline diseases; no deline indication of a large number of compound libraries, including already-approved drugs and corts, the availability of them. Specifically, the pipeline diseases; no deline indication of a large number of compound libraries, including already-approved drugs and corts, the availability of proteins from the pathogen genomes with the Tropical Disease Initiative You are browsing version 1.3 (2008/05/07) of the TDI Kenel. Endrig sile predictor to pproved thops (weathing interpret): 1918 o. Template at Notet o Upped East Suptor Saldor Santor 2010 Generation 2010A minima 2012/2010 Services A228 Ottops 201005 or compound instarces, including already-paproved trugs; and fourth, the valiability of improved bisinformatics analysis, including imethods for comparative proteins structure modeling, binding site identification, virtue gleand screening and drug design. Therefore, we are now in a position to increase the oddle of identifying neglected trupical diseases. Here we encourage a collaboration among scientists to engelected trupical diseases. Here we encourage a collaboration among scientists to engelected trupical diseases. Here we encourage a collaboration among scientists to engelected trupical diseases. Here we encourage a collaboration among scientists to engelected trupical diseases. Here we encourage a collaboration among scientists to engelected trupical diseases. Here we encourage a collaboration among scientists to engelected trupical diseases. Here we encourage a collaboration among scientists to engelected trupical diseases. Here we encourage a collaboration among scientists to engelected trupical diseases. Here we for open source code evelopment, we suggest that the TDI kernel may help overcome a major stumbiling blocki, in this case, for open source drug discovery: the absence of a critical mass of precisiting work hat volunteers can build on incrementally. 5.000 C Figure 1 TDI kernel snapshot of the web page for the Plasmodium fabciparum thymiolytate kinase target (http:// torpicatidisease ong/kenel/GHASL.D. Our computational pipeline predicted that thymiolytate kinase from *P. fabciparum* binds ATM (13-xido-3-deodythymildine-5-monophosphate), a supra-structure of the zidovatine drug approved for the treatment of HV infection. The binding of this ligned to a site on the kinase was experimentally validated by one-dimensional Water-LOSSYP and estivation treafer difference¹⁰ MMB energiments and saturation transfer difference¹⁰ NMR experiments. that volunteers can build on incrementally. predictions based on this limited open source context, where results are made This kernel complements several other experimental testing. Thus, we encourage other investigators to donate their expertise available with limited or no restrictions A freely downloadable version of the TDI ⁶⁰ including collaborative web portals (e.g., http://www.thesynapticleap.org/), public-We hope the testing will occur within the kernel is available in accordance with the Science Commons protocol for implementing open access data (http://sciencecommons. org/project/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 diseases² By minimizing restrictions on the data, including viral terms that would be inherited by all derivative works, we hope to attract as many eybealls as we possibly can to use and improve the kernel. open access data (http://science Table 1 TDI kernel genomes Transcripts^b Modeled targets^c Similar^d ycobacterium leprae 1,605 xoplasma gondi 7,793 we possibly can to use and improve the kernel 3,070 Although many of the drugs in the kernel are 19,607 1,386 proprietary under diverse types of rights, we believe that the existence of public domain 11,714 pairs of targets and compounds will reduce the royalties that patent owners can charge and sponsors must pay. This should decre the large sums of money governments an overnments and 320 VOLUME 27 NUMBER 4 APRIL 2009 NATURE BIOTECHNOLOGY

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

PLOS REGLECTED 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^a}, Stephen M. Maurer⁴, Arti K. Rai⁵, Ginger Taylor⁶, Matthew H. Todd⁷, Antonio Pineda-Lucena², Andrej Sali³*, Marc A. Marti-Renom¹ utural Genonics Unit, Bioriformatics and Genonics Department, Centro de Investigación Principe Felipe, Valencia, Spain, 2 Structural Biology Laboratory, Medi nistry: Oppartment, Centro de Investigación Principe Felipe, Valencia, Spain, 3 Department of Bioengineering and Therapeutic Science, Department macutical Chemistry, and California Institute for Quantitative Bioscience, University of California, Sintrafoco, San Francisco, California, United States of Am Gould School of Law University of Southern California, University of Law California, University of Southern California, University of Southern California, University of Law California 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 leuise. We argue that the stumbing 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 gene 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 teceived 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 use, distribution, and reproduction in any medium, provided the original author and source are credited. use, distribution, and reproduction in any medium, provided the original author and source are credited. Funding: MMA-8 achoneldegs the support from 5 speak Ministerio & Editoxiconi y Cencia grant (BD00207/6470). AS acknowledges the support from the Sandler Family. Supporting Toundation and the National Institutes of Health (BD1 GMS4762, US4 GM07464, PD1 AU33707, and PD1 GM1790). AP4-acknowledges the support from 5 aparts Ministerio de Educación y Cencia grant (AFA200-61864). RC acknowledges the support from the Famon y Cajal Program of the Spanish Ministerio de Educación y Ciencia. We are alor garatelí for computer hardware gifts to AS from Ron Conway, Mile Homer, Intel, BM, Hewlett-Rackard, and Netkop. The Industria Ada or ong garatelí for computer hardware gifts to AS from Ron Conway, Mile Homer, Intel, BM, Hewlett-Rackard, and Netkop. The Industria Ada or ong garatelí for computer hardware gifts to AS from Ron Conway, Mile Homer, Intel, BM, Hewlett-Rackard, and Netkop. The Industria Ada or ong estimation of the manuscipt. Competing Interests: The authors have declared that no competing interests exist. * E-mail: sali@salilab.org (AS); mmarti@cipf.es (MAM-R) Current address: DuPont Knowledge Center, Hyderabad, India Introduction There is a lack of high-quality protein drug targets and drug target 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 [8], identifying binding sites for small molecule [9,10], suggesting drug drag builts of therefore, we are now in a position to leverage this information by identifying potential protein targets for drug discovery. All, and optimizing these leads [13–15]. The absence of an experimentally determined structure, comparative modeling can provide useful models for sequences that are detectably related to havow protein structures [3,4]. Approximately half of known protein structures [3,6]. This coverage www.plosntds.org 1 April 2009 | Volume 3 | Issue 4 | e418

OpenPool/Lab GSK

http://ntdpool.org



Gamo et al. Nature (2010) vol. 465 (7296) pp. 305-10

ARTICLES

nature

Thousands of chemical starting points for antimalarial lead identification

Francisco-Javier Gamo¹, Laura M. Sanz¹, Jaume Vidal¹, Cristina de Cozar¹, Emilio Alvarez¹, Jose-Luis Lavandera¹, Dana E. Vanderwall², Darren V. S. Green³, Vinod Kumar⁴, Samiul Hasan⁴, James R. Brown⁴, Catherine E. Peishoff⁵, Lon R. Cardon⁶ & Jose F. Garcia-Bustos¹

Malaria is a devastating infection caused by protozoa of the genus Plasmodium. Drug resistance is widespread, no new chemical class of antimalarials has been introduced into clinical practice since 1996 and there is a recent rise of parasite strains with reduced sensitivity to the newest drugs. We screened nearly 2 million compounds in GlaxoSmithKline's chemical library for inhibitors of P. falciparum, of which 13,533 were confirmed to inhibit parasite growth by at least 80% at 2 µM concentration. More than 8,000 also showed potent activity against the multidrug resistant strain Dd2. Most (82%) compounds originate from internal company projects and are new to the malaria community. Analyses using historic assay data suggest several novel mechanisms of antimalarial action, such as inhibition of protein kinases and host-nathogen interaction related targets. Chemical structures and associated data are hereby made public to encourage additional drug lead identification efforts and further research into this disease

With approximately 243 million cases and 863,000 attributed deaths Tres Cantos antimalarial compound set (TCAMS) reported globally in 2009 (ref. 1), malaria is one of the most severe infectious diseases, primarily affecting the world's most disadvantaged populations. Of the four typically recognized *Plasmodium* species causing disease in humans, *Plasmodium falciparum* causes most mortality, mainly in children below the age of 5, and Plasmodium vivax most morbidity, additionally representing a reservoir of latent infection that hampers current control and future elimination efforts2. No new class of antimalarials has been introduced into clinical practice since 1996 (ref. 3), owing to the intrinsic difficulties in discovering and developing new antimicrobials, as well as a relative lack of public and private resource commitment towards antimalarial research. Today, the last class of widely efficacious drugs, the artemisinins, is being compromised by the rise of *P. falciparum* strains with reduced clinical response to artemisinin-containing drug combinations4-6. The genomics revolution has not yet led to new antimalarial medicines and target-based lead discovery has produced disappoint-ing results, generally for lack of whole-cell activity as documented for antibacterials². To secure that property in all chemical starting points for new antimalarial leads, we have tested the approximately 2 million-compound library used for high throughput screening at GlaxoSmithKline (GSK) for inhibitors of *P. falciparum*'s intraerythrocytic cycle, the Plasmodium species causing the highest mortality and the parasite growth phase responsible for disease symptoms as well as being amenable to *in vitro* culture. Here we describe 13,533 compounds confirmed to inhibit parasite growth by more than 80% at $2\,\mu$ M concentration. Only 15% displayed some cytotoxicity in that they inhibited proliferation of the HepG2 human hepatoma cell line by more than 50% at 10 μ M. All of these proven plasmodial inhibitors, of which 82% were previously proprietary and thus unknown to the general research community, are hereby made public to accelerate the pace of drug development for malaria.

Vol 465|20 May 2010|doi:10.1038/nature0910

The 1,986,056 compounds present in GSK's screening collection in January 2009 were tested for inhibition of *P. falciparum* 3D7 at $2\,\mu$ M under *in vitro* conditions described in Methods. 19,451 primary hits inhibiting parasite growth by more than 80% were obtained. Fresh samples of these primary hits were tested in two independent experi ments and compounds displaying 80% or higher inhibition of parasite growth in at least two of the three assay runs were considered confirmed hits. 13,533 compounds were identified using this protocol (confirmation rate > 70%). We did not detect any compounds in this set as non-specific inhibitors of the biochemical readout sys-tem by testing directly for inhibition of lactate dehydrogenase (LDH) in P. falciparum extracts (Methods). Evidence of cytotoxicity against human hepatoma HepG2 cells (a widely used in vitro marker for liver toxicity8), or interference with the luciferase reporter system used in the cytotoxicity assay (Methods), was observed in just 1,982 of the compounds when tested at 10 μ M. This relative lack of non-specific cell toxicity is probably due in part to the low (2 μ M) primary screen-ing concentration used⁹. Estimation of the concentrations producing 50% inhibition of *P. falciparum* growth (XC₅₀, see Methods) indicated that most compounds are sub-micromolar inhibitors. The full compound set (TCAMS) and data table (Supplementary Table 1 and available at http://www.ebi.ac.uk/chemblntd) contains 13,533 compound entries. We have detected 139 of these as variations in salt form or stereochemistry of 68 parent structures, which make good internal controls for the biological assay data. They appear as differ-ent compounds with the same structure. When the stereochemistry is resolved it shows in the SMILES structural code in Supplementary Table 1 and in the Chembl-NTD database (http://www.ebi.ac.uk/ chemblntd).

Representatives from all but one class of clinically used antimalarials have been recovered in the screen, providing additional validation

305

10.0

¹Tres Cantos Medicines Development Campus, GlaxoSmithKline, Severo Ochoa 2, 28760 Tres Cantos, Spain. ²Computational and Structural Chemistry, GlaxoSmithKline, Five Moore Drive, Research Triangle Park, North Carolina 27709-3398, USA. ²Computational and Structural Chemistry, GlaxoSmithKline, Breise Mood Road, Hertfordshire, Stevenage SG1 2NV, UK. ²Computational Biology, Quantitative Sciences, GlaxoSmithKline, Stowas Mithkline, Steven ²Computational and Structural Chemistry, GlaxoSmithKline, 1250 South Collegeville, Road, Collegeville, Road, Sub, Marking Stevenson, Ste land Road. King of Prussia. Pennsylvania 19406. USA

©2010 Macmillan Publishers Limited. All rights reserved

20010 We will be a straight provide the straight of the straig

Acknowledgments

http://tropicaldisease.org
http://sgu.bioinfo.cipf.es

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

MAMMOTH Angel R. Ortiz

FUNCTIONAL ANNOTATION Fatima Al-Shahrour Joaquin Dopazo Tropical Disease Initiative Stephen Maurer (UC Berkeley) Arti Rai (Duke U) Andrej Sali (UCSF) Ginger Taylor (TSL) Matthew Todd (U Sydney)

MODEL ASSESSMENT David Eramian Min-Yi Shen Damien Devos GeMoA Lluís Ballell (GSK) Olivier Neyrolles (IPBS) Matthias Wilmanns (EMBL) Ginger Taylor (TSL) Brigitte Cicquel (IP)





