

NEXT UP:

- Overview of structural bioinformatics
 - Major motivations, goals and challenges
- Fundamentals of protein structure
 - Composition, form, forces and dynamics
- Representing and interpreting protein structure
 - Modeling energy as a function of structure
- Example application areas
 - drug discovery & Predicting functional dynamics

THETRADITIONAL EMPIRICAL PATH TO DRUG DISCOVERY

Compound library

(commercial, in-house, synthetic, natural)

High throughput screening (HTS)

Hit confirmation

Lead compounds

(e.g., µM K_d)

Lead optimization

(Medicinal chemistry)

Animal and clinical

← Potent drug candidates evaluation (nM K_d)

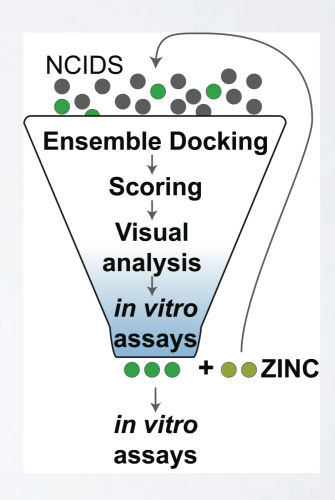
COMPUTER-AIDED LIGAND DESIGN

Aims to reduce number of compounds synthesized and assayed

Lower costs

Reduce chemical waste

Facilitate faster progress



Two main approaches:

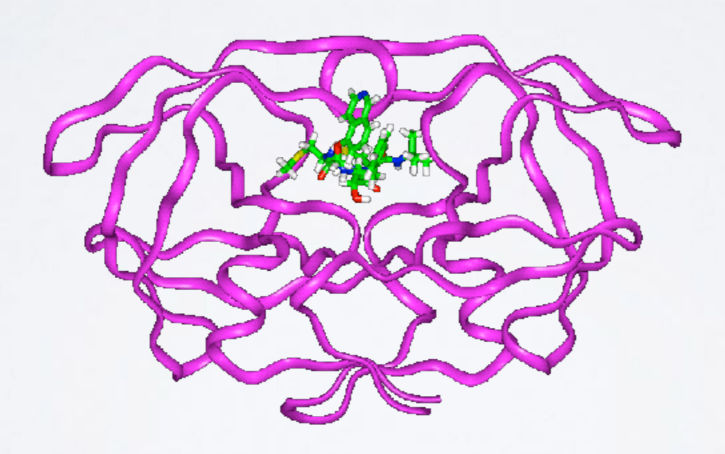
- (1). Receptor/Target-Based
- (2). Ligand/Drug-Based

Two main approaches:

- (1). Receptor/Target-Based
- (2). Ligand/Drug-Based

SCENARIO I: RECEPTOR-BASED DRUG DISCOVERY

Structure of Targeted Protein Known: Structure-Based Drug Discovery

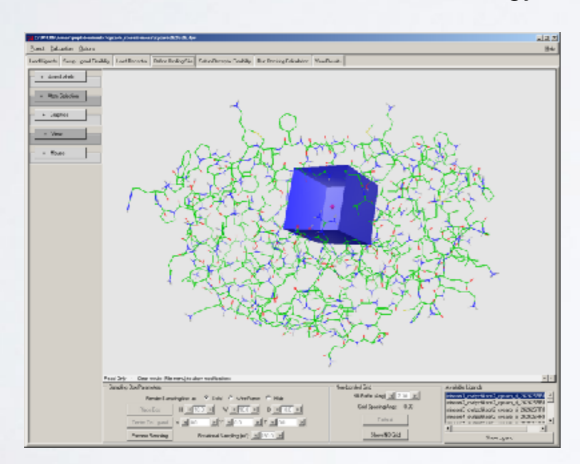


HIV Protease/KNI-272 complex

PROTEIN-LIGAND DOCKING

Structure-Based Ligand Design

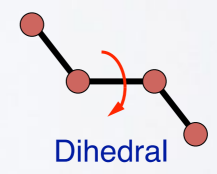
Docking software
Search for structure of lowest energy



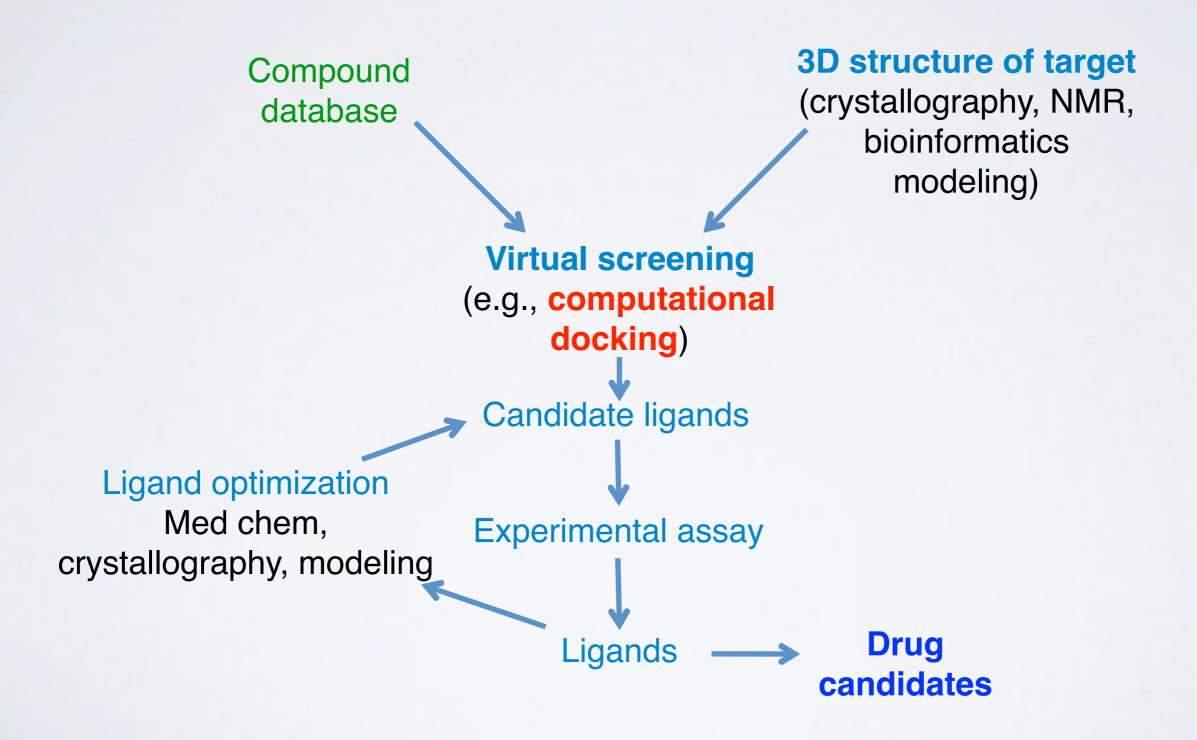
Potential function
Energy as function of structure







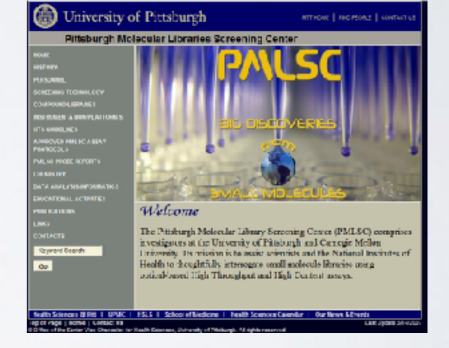
STRUCTURE-BASED VIRTUAL SCREENING



COMPOUND LIBRARIES





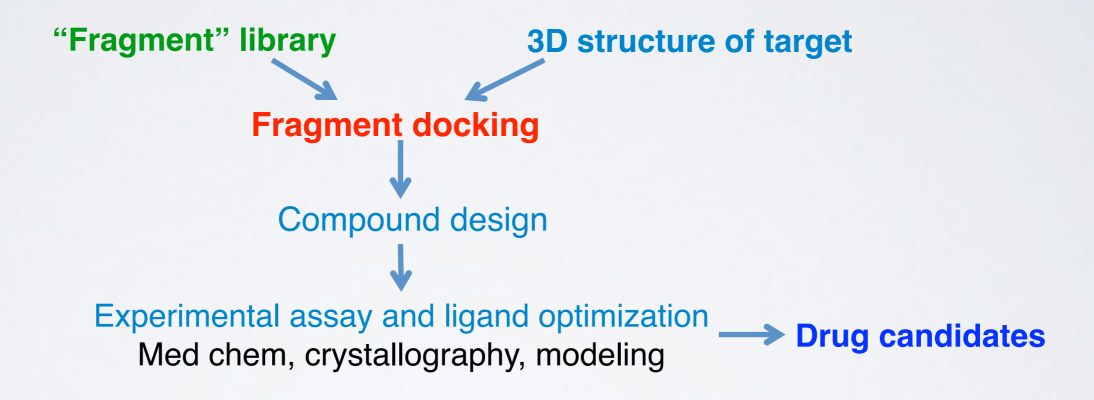


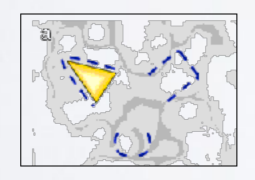
Commercial (in-house pharma)

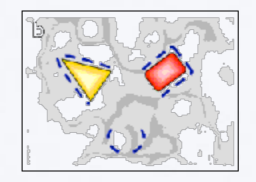
Government (NIH)

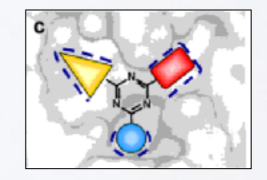
Academia

FRAGMENTAL STRUCTURE-BASED SCREENING



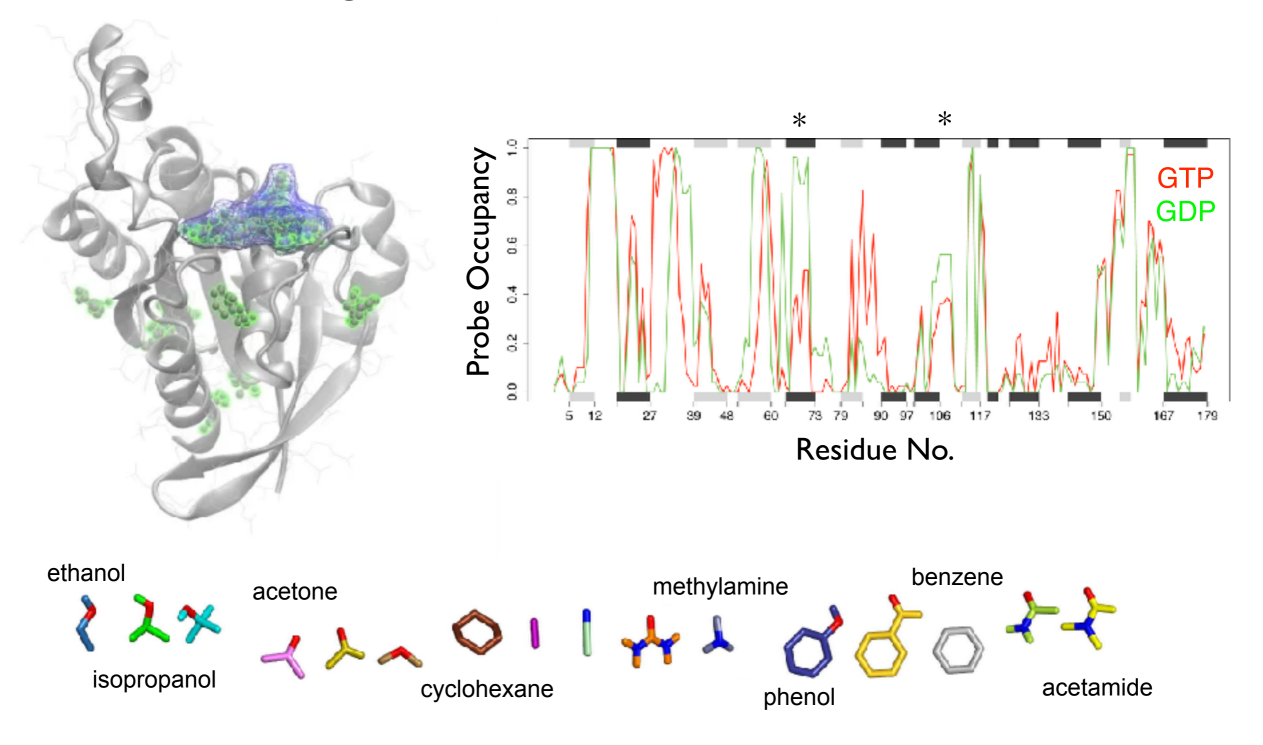






Multiple non active-site pockets identified

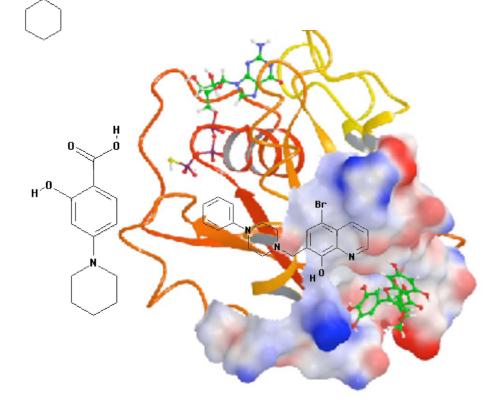
Small organic probe fragment affinities map multiple potential binding sites across the structural ensemble.



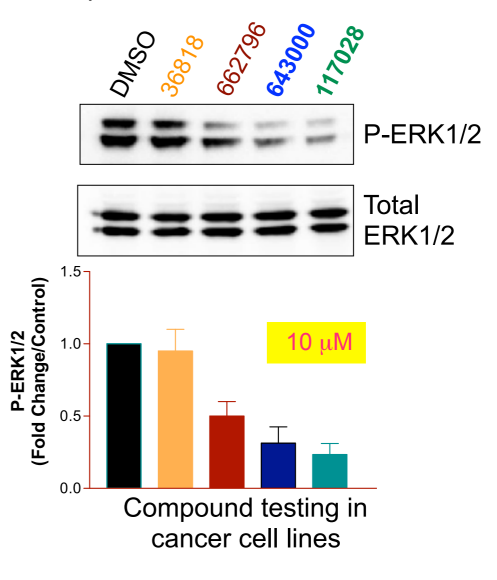
Ensemble docking & candidate inhibitor testing

Top hits from ensemble docking against distal pockets were tested for inhibitory effects on basal ERK activity in glioblastoma cell lines.

Ensemble computational docking

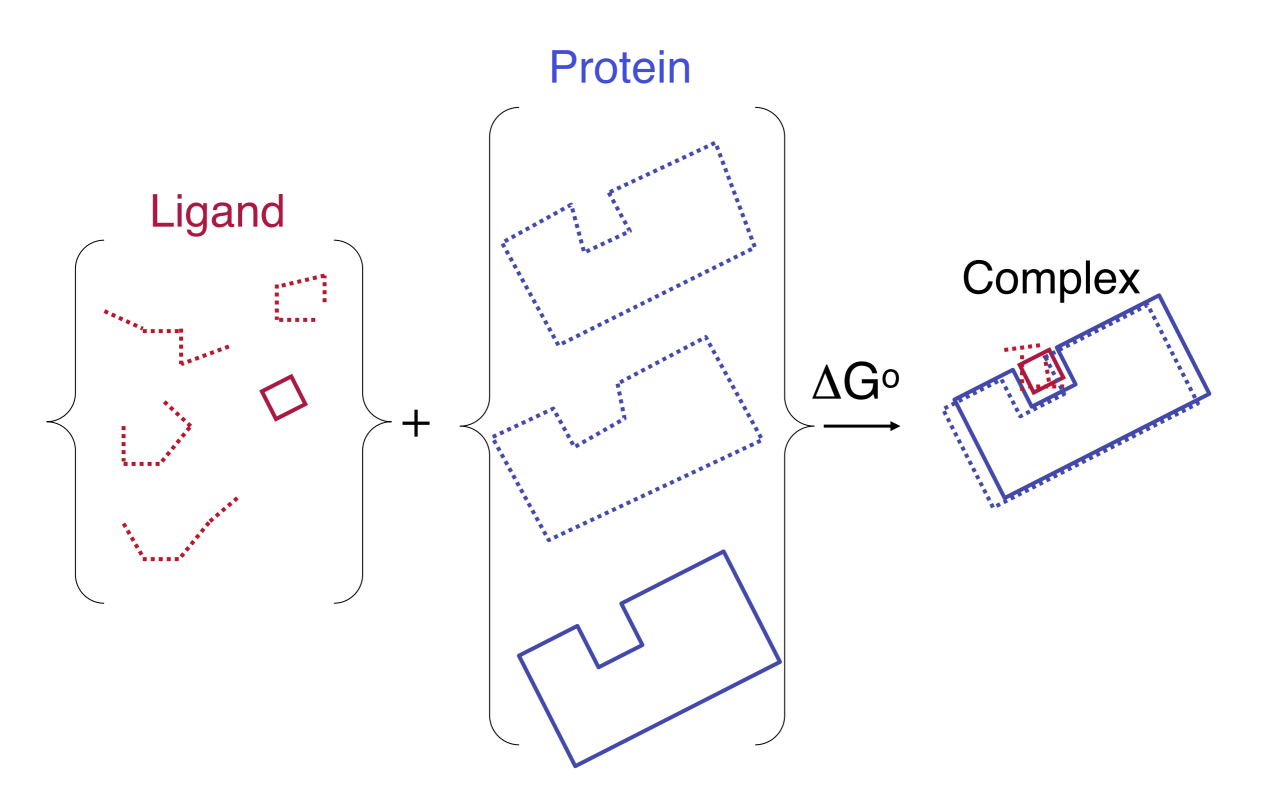


Compound effect on U251 cell line



PLoS One (2011, 2012)

Proteins and Ligand are Flexible



COMMON SIMPLIFICATIONS USED IN PHYSICS-BASED DOCKING

Quantum effects approximated classically

Protein often held rigid

Configurational entropy neglected

Influence of water treated crudely

Two main approaches:

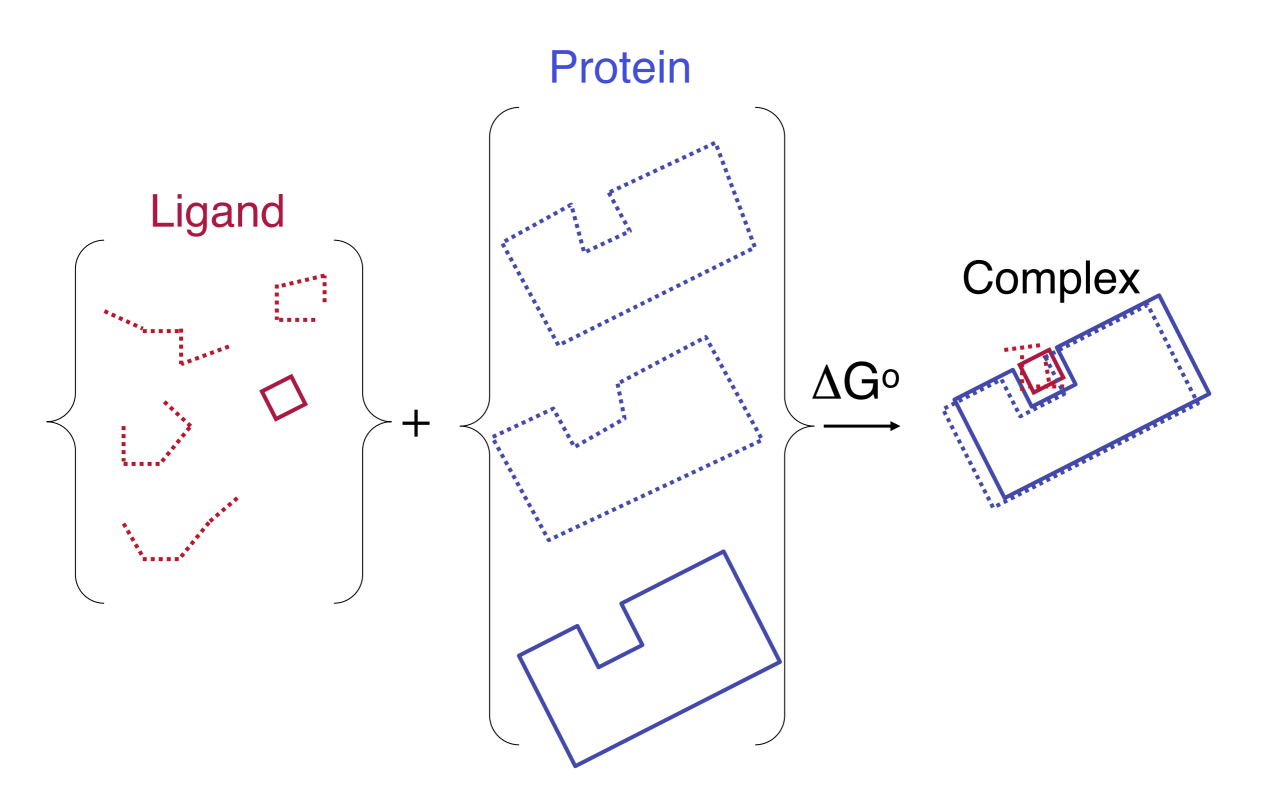
- (1). Receptor/Target-Based
- (2). Ligand/Drug-Based

Hand-on time!

https://bioboot.github.io/bimm143_W18/lectures/#12

You can use the classroom computers or your own laptops. If you are using your laptops then you will need to install VMD and MGLTools

Proteins and Ligand are Flexible



HTTP://129.177.232.111:3848/PCA-APP/

HTTPS://DCMB-GRANT-SHINY.UMMS.MED.UMICH.EDU/PCA-APP/

HTTP://BIO3D.UCSD.EDU/PCA-APP/

Two main approaches:

- (1). Receptor/Target-Based
- (2). Ligand/Drug-Based

Scenario 2

Structure of Targeted Protein Unknown: Ligand-Based Drug Discovery

e.g. MAP Kinase Inhibitors

Using knowledge of existing inhibitors to discover more

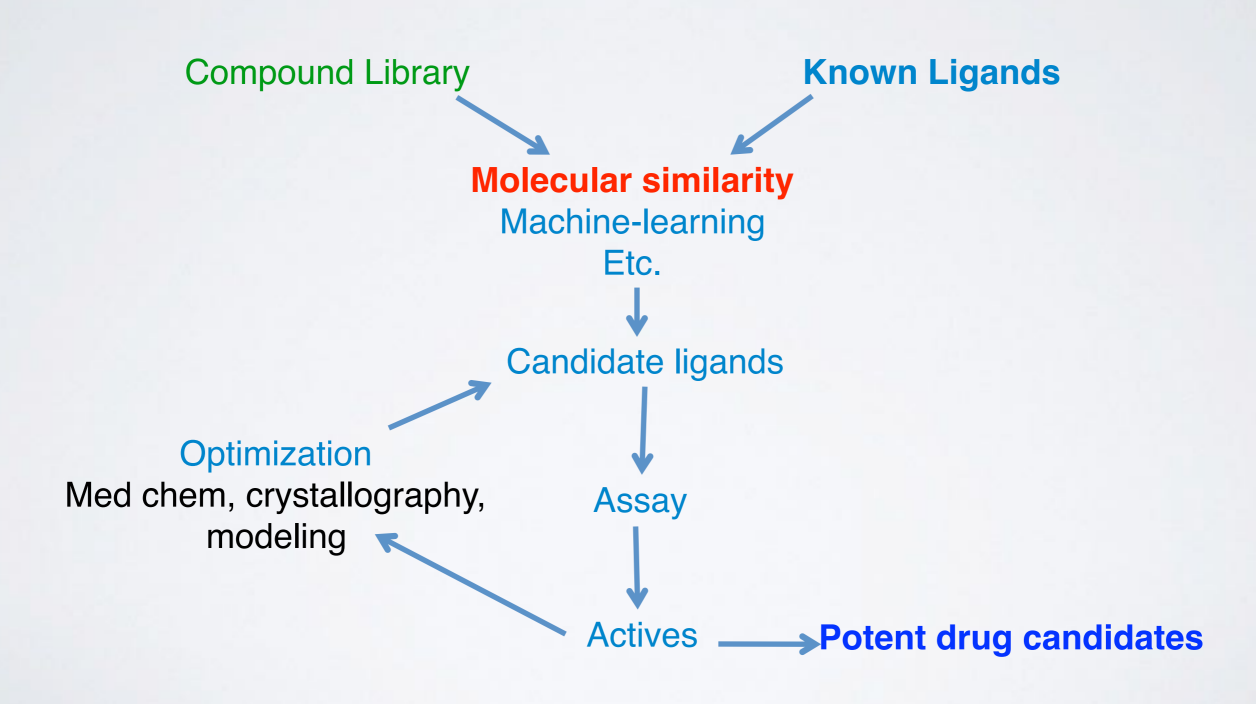
Why Look for Another Ligand if You Already Have Some?

Experimental screening generated some ligands, but they don't bind tightly enough

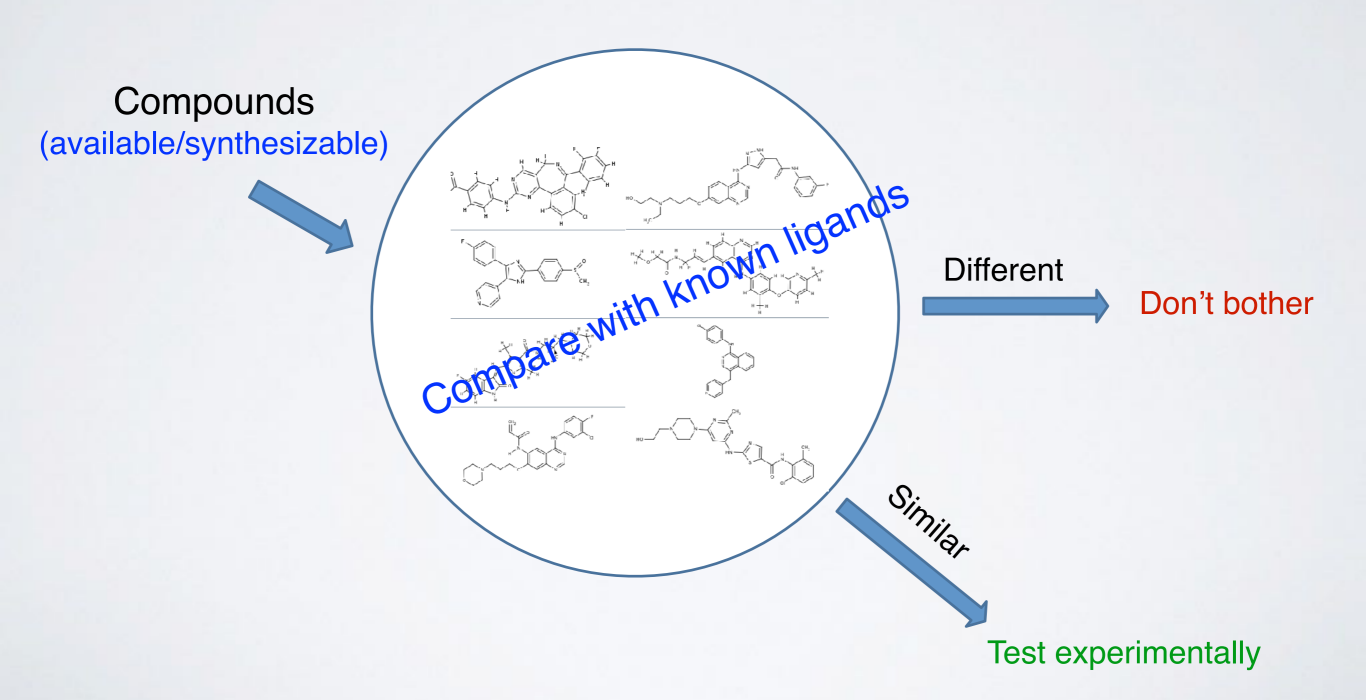
A company wants to work around another company's chemical patents

An high-affinity ligand is toxic, is not well-absorbed, difficult to synthesize etc.

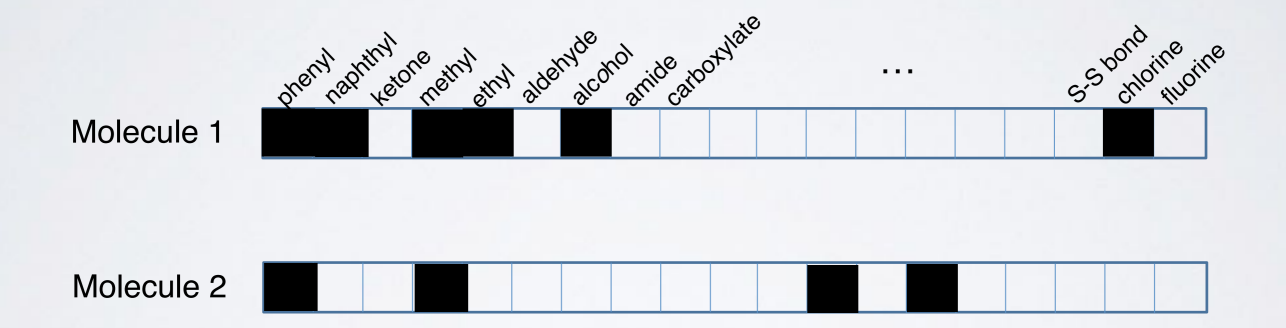
LIGAND-BASED VIRTUAL SCREENING



CHEMICAL SIMILARITY LIGAND-BASED DRUG-DISCOVERY



CHEMICAL FINGERPRINTS BINARY STRUCTURE KEYS



CHEMICAL SIMILARITY FROM FINGERPRINTS

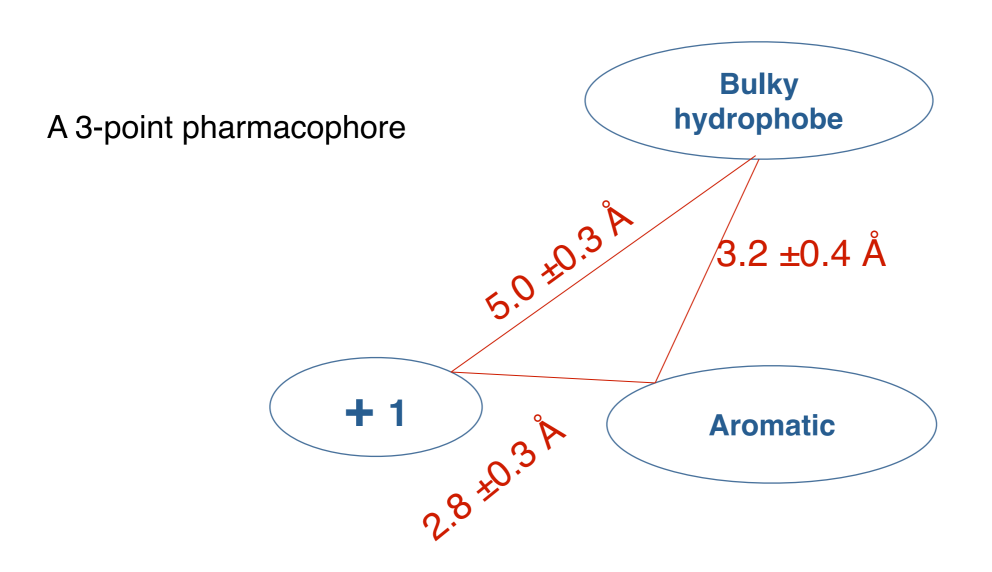


Tanimoto Similarity (or Jaccard Index), T

$$T \equiv \frac{N_I}{N_U} = 0.25$$



Pharmacophore Models Φάρμακο (drug) + Φορά (carry)

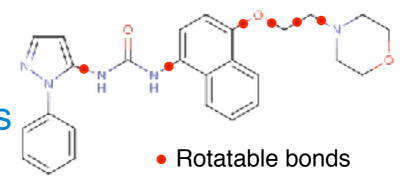


Molecular Descriptors

More abstract than chemical fingerprints

Physical descriptors

molecular weight charge dipole moment number of H-bond donors/acceptors number of rotatable bonds hydrophobicity (log P and clogP)



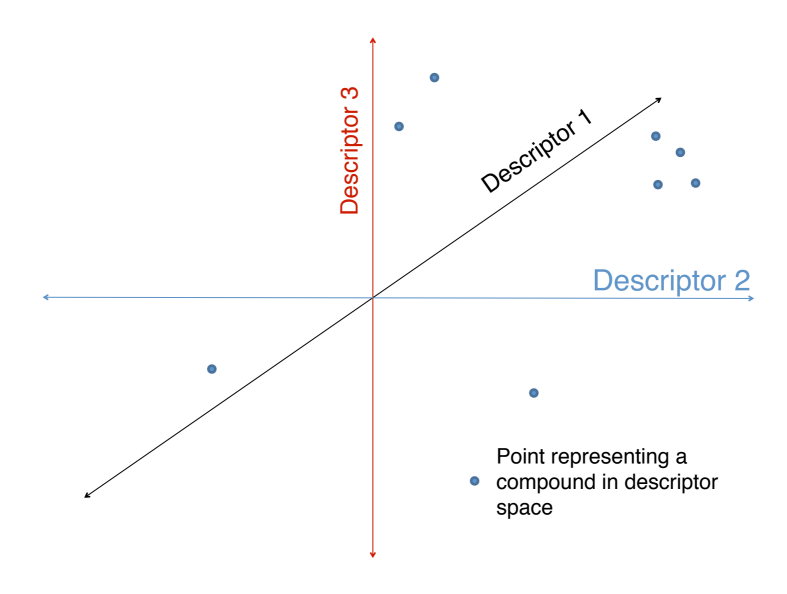
Topological

branching index measures of linearity vs interconnectedness

Etc. etc.

A High-Dimensional "Chemical Space"

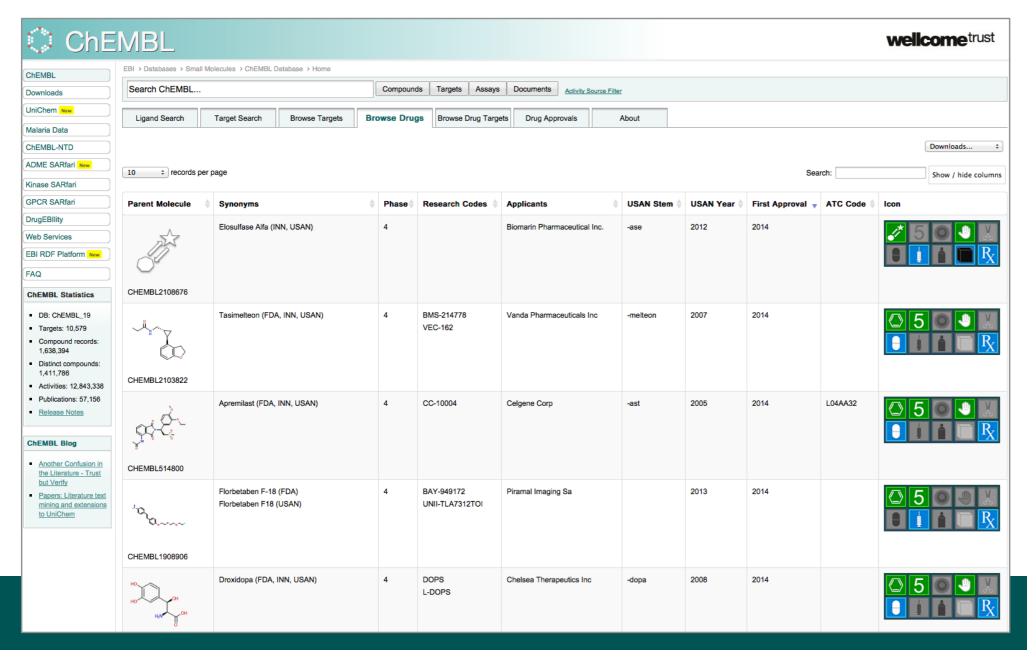
Each compound is at a point in an n-dimensional space Compounds with similar properties are near each other



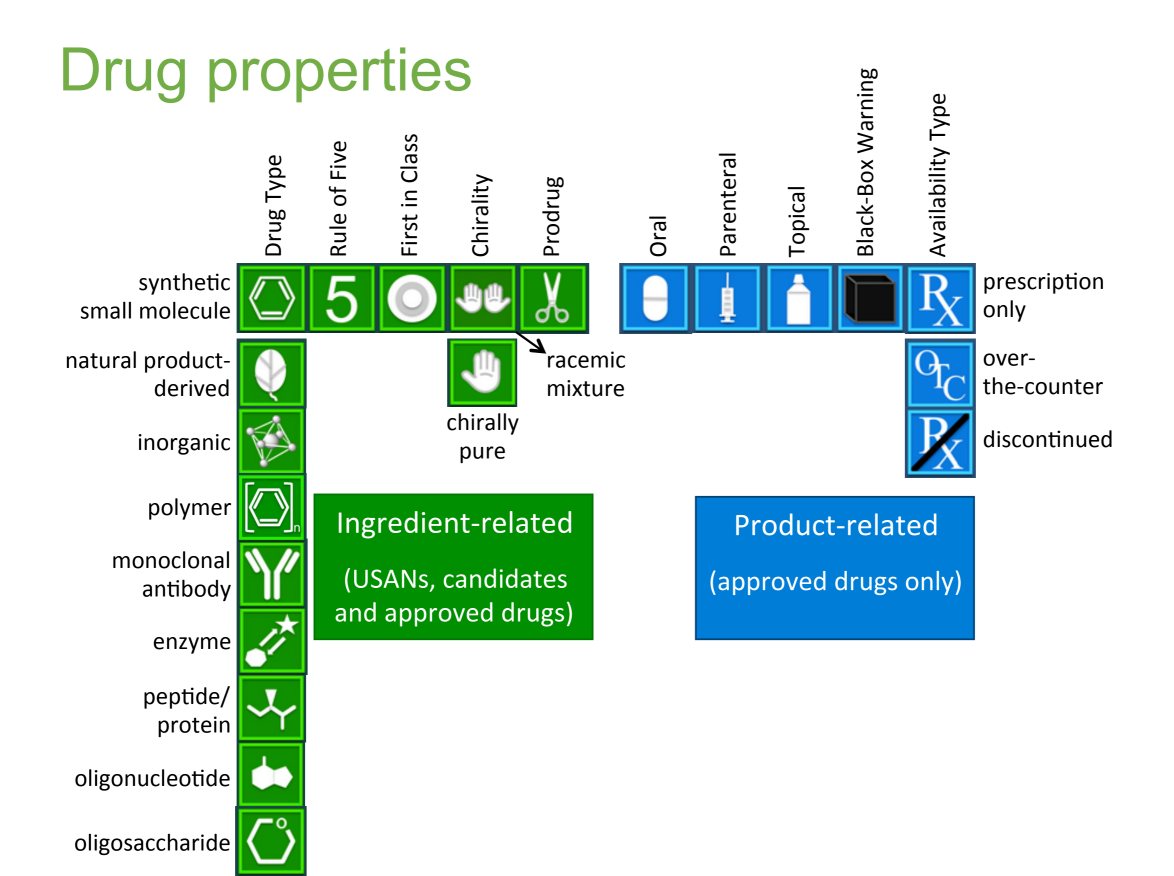
Apply multivariate statistics and machine learning for descriptorselection. (e.g. partial least squares, PCA, support vector machines, random forest, deep learning etc.)

Approved drugs and clinical candidates

- Catalogue approved drugs and clinical candidates from FDA Orange Book, and USAN applications
- Small molecules and biotherapeutics







LIPINSKI'S RULE OF FIVE

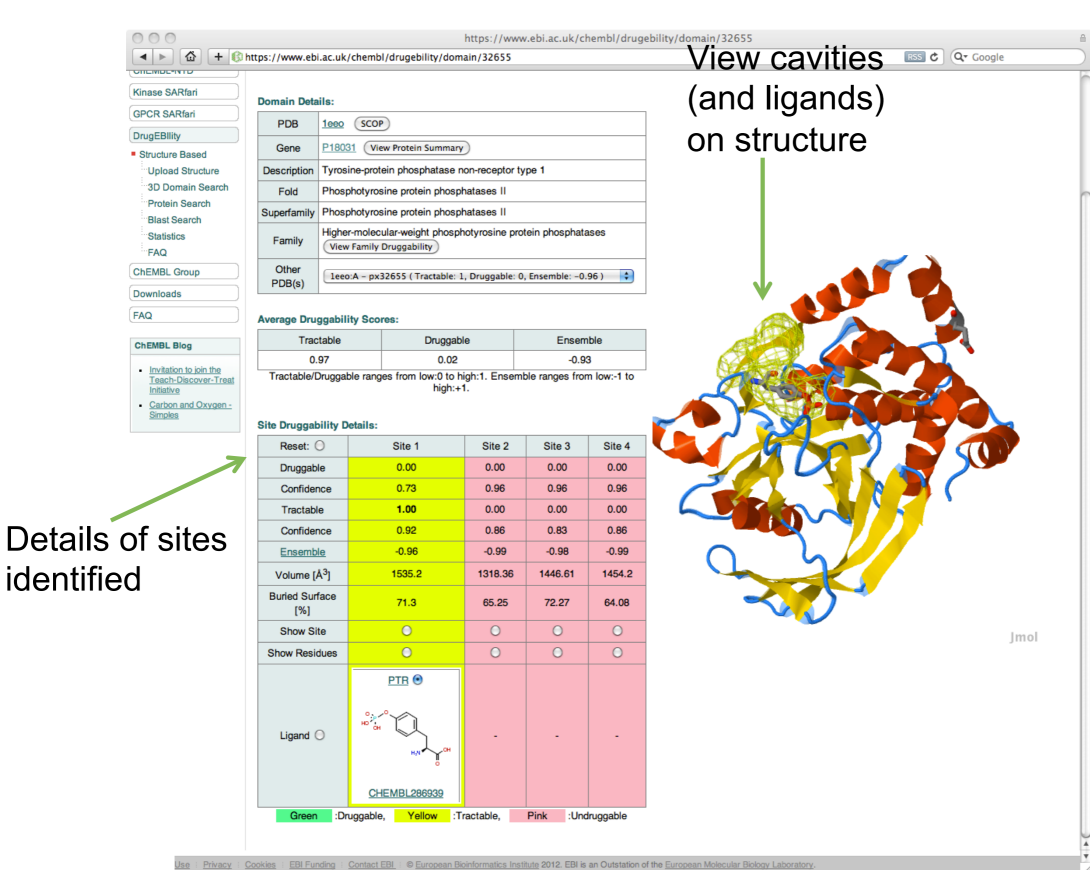
Lipinski's rule of five states that, in general, an orally active drug has no more than one violation of the following criteria:

- Not more than 5 hydrogen bond donors (nitrogen or oxygen atoms with one or more hydrogen atoms)
- Not more than 10 hydrogen bond acceptors (nitrogen or oxygen atoms)
- A molecular mass less than 500 daltons
- An octanol-water partition coefficient log P not greater than 5

Rules for drug discovery success

- Set of approved drugs or medicinal chemistry compounds and their targets can be used to derive rules for drug discovery success (or failure):
 - What features make a successful drug target?
 - What features make a protein druggable by small molecules?
 - What features of a compound contribute to good oral bioavailability?
 - What chemical groups may be associated with toxicity?

Druggability prediction



Examples

nature

Vol 460 16 July 2009 doi:10.1038/nature08160

ARTICLES

The genome of the blood fluke *Schistosoma* mansoni

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Schistosoma mansoni is responsible for the neglected tropical disease schistosomiasis that affects 210 million people in 76 countries. Here we present analysis of the 363 megabase nuclear genome of the blood fluke. It encodes at least 11,809 genes, with an unusual intron size distribution, and new families of micro-exon genes that undergo frequent alternative splicing. As the first sequenced flatworm, and a representative of the Lophotrochozoa, it offers insights into early events in the evolution of the animals, including the development of a body pattern with bilateral symmetry, and the development of tissues into organs. Our analysis has been informed by the need to find new drug targets. The deficits in lipid metabolism that make schistosomes dependent on the host are revealed, and the identification of membrane receptors, ion channels and more than 300 proteases provide new insights into the biology of the life cycle and new targets. Bioinformatics approaches have identified metabolic chokepoints, and a chemogenomic screen has pinpointed schistosome proteins for which existing drugs may be active. The information generated provides an invaluable resource for the research community to develop much needed new control tools for the treatment and eradication of this important and neglected disease.

Schistosomiasis is a neglected tropical disease that ranks with malaria and tuberculosis as a major source of morbidity affecting approximately 210 million people in 76 countries, despite strenuous control efforts¹. It is caused by blood flukes of the genus *Schistosoma* (phylum Platyhelminthes), which exhibit dioecy and have complex life cycles comprising several morphologically distinct phenotypes in definitive human and intermediate snail hosts. *Schistosoma mansoni*, one of the three major human species, occurs across much of sub-Saharan Africa, parts of the Middle East, Brazil, Venezuela and some West Indian islands. The mature flukes dwell in the human portal vasculature, depositing eggs in the intestinal wall that either

pass to the gut lumen and are voided in the faeces, or travel to the liver where they trigger immune-mediated granuloma formation and peri-portal fibrosis². Approximately 280,000 deaths per annum are attributable to schistosomiasis in sub-Saharan Africa alone³. However, the disease is better known for its chronicity and debilitating morbidity⁴. A single drug, praziquantel, is almost exclusively used to treat the infection but this does not prevent reinfection, and with the large-scale control programmes in place, there is concern about the development of drug resistance. Indeed, resistance can be selected for in the laboratory and there are reports of increased drug tolerance in the field⁵.

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NATURE CHEMISTRY | ARTICLE





Quantifying the chemical beauty of drugs

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Affiliations | Contributions | Corresponding author

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Received 01 September 2011 | Accepted 02 December 2011 | Published online 24 January 2012



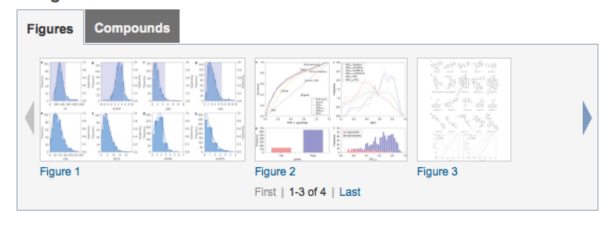
Abstract

Abstract · References · Author information · Supplementary Information

Drug-likeness is a key consideration when selecting compounds during the early stages of drug discovery. However, evaluation of drug-likeness in absolute terms does not reflect adequately the whole spectrum of compound quality. More worryingly, widely used rules may inadvertently foster undesirable molecular property inflation as they permit the encroachment of rule-compliant compounds towards their boundaries. We propose a measure of drug-likeness based on the concept of desirability called the quantitative estimate of drug-likeness (QED). The empirical rationale of QED reflects the underlying distribution of molecular properties. QED is intuitive, transparent, straightforward to implement in many practical settings and allows compounds to be ranked by their relative merit. We extended the utility of QED by applying it to the problem of molecular target druggability assessment by prioritizing a large set of published bioactive compounds. The measure may also capture the abstract notion of aesthetics in medicinal chemistry.

Subject terms: Pharmacology • Theoretical chemistry

At a glance



Target prediction models

- Active compounds from ChEMBL can be used to train target prediction models
- Variety of methods used
 - Multi-Category Naïve Bayesian Classifier (e.g., ChEMBL)
 - Chemical similarity between ligand sets (e.g., SEA)
 - 3D similarity between ligands (e.g., SwissTargetPrediction)
 - Protein and ligand descriptors (e.g., Proteochemometric models)
- Open source tools available for many methods
 - E.g., Scikit-learn with RDKit

Examples at: https://github.com/chembl/mychembl/blob/master/ipython_notebooks

Examples





Mycobacterial Dihydrofolate Reductase Inhibitors Identified Using Chemogenomic Methods and *In Vitro* Validation

Grace Mugumbate¹, Katherine A. Abrahams², Jonathan A. G. Cox², George Papadatos¹, Gerard van Westen¹, Joël Lelièvre³, Szymon T. Calus², Nicholas J. Loman², Lluis Ballell³, David Barros³, John P. Overington¹*, Gurdyal S. Besra²*

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OPEN ACCESS

Citation: Mugumbate G, Abrahams KA, Cox JAG, Papadatos G, van Westen G, Lellèvre J, et al. (2015) Mycobacterial Dihydrofolate Reductase Inhibitors Identified Using Chemogenomic Methods and In Vitro Validation. PLoS ONE 10(3): e0121492. doi:10.1371/ journal.pone.0121492

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Data Availability Statement: All relevant data are within the paper and its Supporting Information files.

Funding: GM and GvW are grateful to EMBL and Marie Curie Actions for funding this work. GSB acknowledges support in the form of a Personal Research Chair from Mr. James Bardrick, a Royal Society Wolfston Research Merit Award, The Wellcome Trust (681569/Z/06/Z), and the Medical Research Council (MR/K012118/1). The research also received funding from the European Union's 7th framework programme (FP7-2007–2013) under grant agreement ORCHID no. 261378. The funders had no

Abstract

The lack of success in target-based screening approaches to the discovery of antibacterial agents has led to reemergence of phenotypic screening as a successful approach of identifying bioactive, antibacterial compounds. A challenge though with this route is then to identify the molecular target(s) and mechanism of action of the hits. This target identification, or deorphanization step, is often essential in further optimization and validation studies. Direct experimental identification of the molecular target of a screening hit is often complex, precisely because the properties and specificity of the hit are not yet optimized against that target, and so many false positives are often obtained. An alternative is to use computational, predictive, approaches to hypothesize a mechanism of action, which can then be validated in a more directed and efficient manner. Specifically here we present experimental validation of an in silico prediction from a large-scale screen performed against Mycobacterium tuberculosis (Mtb), the causative agent of tuberculosis. The two potent anti-tubercular compounds studied in this case, belonging to the tetrahydro-1,3,5-triazin-2-amine (THT) family, were predicted and confirmed to be an inhibitor of dihydrofolate reductase (DHFR), a known essential Mtb gene, and already clinically validated as a drug target. Given the large number of similar screening data sets shared amongst the community, this in vitro validation of these target predictions gives weight to computational approaches to establish the mechanism of action (MoA) of novel screening hit.

Introduction

The human pathogen, Mycobacterium tuberculosis (Mtb) is the causative agent of tuberculosis (TB), an infectious disease that is widespread, infecting around one third of the world's population [1]. The discovery of streptomycin in 1943, and the subsequent discovery and

ARTICLE

doi:10.1038/nature11159

Large-scale prediction and testing of drug activity on side-effect targets

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Discovering the unintended 'off-targets' that predict adverse drug reactions is daunting by empirical methods alone. Drugs can act on several protein targets, some of which can be unrelated by conventional molecular metrics, and hundreds of proteins have been implicated in side effects. Here we use a computational strategy to predict the activity of 656 marketed drugs on 73 unintended 'side-effect' targets. Approximately half of the predictions were confirmed, either from proprietary databases unknown to the method or by new experimental assays. Affinities for these new off-targets ranged from 1 nM to 30 μ M. To explore relevance, we developed an association metric to prioritize those new off-targets that explained side effects better than any known target of a given drug, creating a drug-target-adverse drug reaction network. Among these new associations was the prediction that the abdominal pain side effect of the synthetic oestrogen chlorotrianisene was mediated through its newly discovered inhibition of the enzyme cyclooxygenase-1. The clinical relevance of this inhibition was borne out in whole human blood platelet aggregation assays. This approach may have wide application to de-risking toxicological liabilities in drug discovery.

Adverse drug reactions (ADRs) can limit the use of otherwise effective drugs. Next to lack of efficacy, they are the leading cause for attrition in clinical trials of new drugs1-3 and are more prominent still in the failure of molecules to advance from pre-clinical research into human trials4. Some ADRs are caused by modulation of the primary target of a drug5, others result from non-specific interactions of reactive metabolites⁶. In many cases, however, ADRs are caused by unintended activity at off-targets. Notorious examples of off-target toxicity include that of the appetite suppressant fenfluramine-phentermine (fenphen), which was withdrawn from the market after numerous patient deaths. These owed to the activation of the 5-hydroxytryptamine-2B (5-HT_{2B}) receptor by one of its metabolites, norfenfluramine, leading to proliferative valvular heart disease7. Similarly, well-known drugs, such as the antihistamine terfenadine, have been withdrawn because they caused arrhythmias and death, which have been attributed to their off-target inhibition of the human ether-à-go-go-related gene potassium channel (hERG, also known as KCNH2)^{8,9}. Prediction of unknown off-target drug interactions might prevent such disastrous drug toxicities, which are often detected only after fatalities in the clinic, and might allow safer molecules to be prioritized for pre-clinical development. Methods to systematically predict off-targets, and associate these with side effects, have thus attracted intense interest 10-16, frequently in the form of either chemical genomics 17,18 or informatics 19-26 approaches.

Whereas the informatics methods have never been tested systematically on a large scale, in principle they can be deployed against thousands of targets. Here we present a large-scale, prospective evaluation of safety target prediction using one such method, the similarity ensemble approach (SEA)^{25–27}. SEA calculates whether a molecule will bind to a target based on the chemical features it shares with those of known ligands, using a statistical model to control for random similarity. Because SEA relies only on chemical similarity, it can be applied systematically and, for those targets that have known ligands,

comprehensively. For 656 drugs approved for human use (Supplementary Table 1), targets were predicted from among 73 proteins (Supplementary Table 2 and Methods) with established association of ADRs^{22,28}, for which assays were available at Novartis. Encouragingly, many of the predictions were confirmed, often at pharmacologically relevant concentrations. This motivated us to develop a guilt-by-association metric that linked the new targets to the ADRs of those drugs for which they are the primary or well-known off-targets, creating a drug-target-ADR network. The applicability and the limitations of this approach will be considered.

Testing the predictions

The 656 drugs were computationally screened for their likelihood to bind to 73 targets (Supplementary Table 2) using SEA25-27. The targets belong to the Novartis in vitro safety panels based on their association with ADRs 22,28. Here we insisted that they also be described in the ChEMBL database29, enabling correspondence with SEA predictions (Supplementary Table 2). ChEMBL annotates more than 285,000 ligands modulating more than 1,500 different human targets with affinities better than 30 µM. SEA calculated the similarity of each drug versus each set of ligands for the 73 targets, comparing the overall set similarity to a model of such expected at random. For instance, the sodium channel blocker aprindine loosely resembled the set of histamine H1 ligands; although no single H1 ligand was strongly similar to the drug (Table 1), the overall similarity of the set was much greater than expected at random, leading to a highly significant SEA expectation value (E value) of 5×10^{-26} between aprinidine and H₁ receptor ligands. Only 1,644 of the more than 47,000 possible drugtarget pairs had significant E values. Of these, 403 were already known in ChEMBL and so were trivially confirmed; we do not consider these further. Of the remaining 1,241 predictions, 348 (28%) were unknown to ChEMBL, but could be found in proprietary ligand-target databases that were unavailable to SEA (see Methods). The remaining

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^{*}These authors contributed equally to this work.

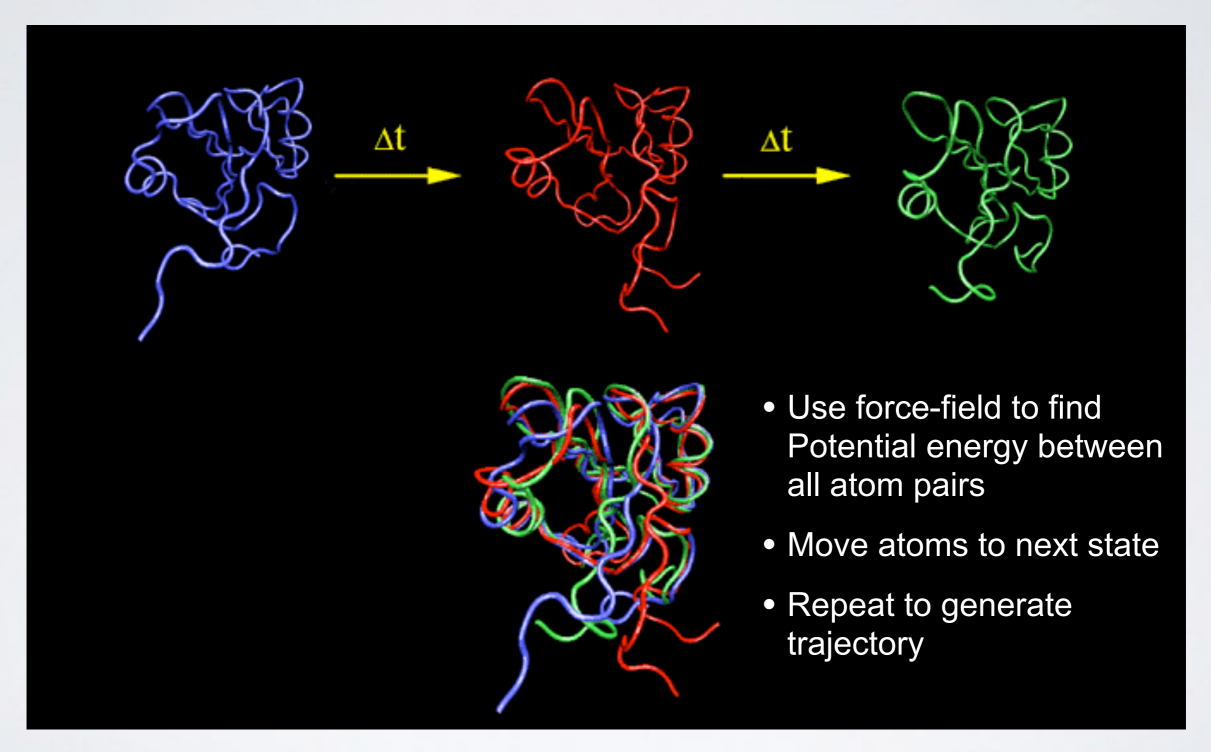
NEXT UP:

- Overview of structural bioinformatics
 - Major motivations, goals and challenges
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 - Modeling energy as a function of structure
- Example application areas
 - Drug discovery & predicting <u>functional dynamics</u>

PREDICTING FUNCTIONAL DYNAMICS

- Proteins are <u>intrinsically flexible</u> molecules with internal motions that are often intimately coupled to their biochemical function
 - E.g. ligand and substrate binding, conformational activation, allosteric regulation, etc.
- Thus knowledge of dynamics can provide a deeper understanding of the <u>mapping of structure to</u> <u>function</u>
 - Molecular dynamics (MD) and normal mode analysis
 (NMA) are two major methods for predicting and characterizing molecular motions and their properties

MOLECULAR DYNAMICS SIMULATION



McCammon, Gelin & Karplus, *Nature* (1977)

[See: https://www.youtube.com/watch?v=ui1ZysMFcKk]

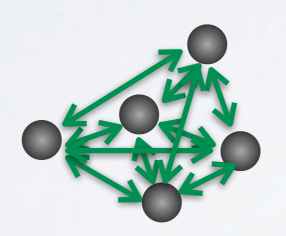
Divide time into discrete (~1fs) time steps (Δt) (for integrating equations of motion, see below)



Divide time into discrete (~1fs) time steps (Δt) (for integrating equations of motion, see below)



At each time step calculate pair-wise atomic forces (*F(t)*) (by evaluating force-field gradient)



Nucleic motion described classically

$$m_i rac{d^2}{dt^2} ec{R}_i = - ec{
abla}_i E(ec{R})$$

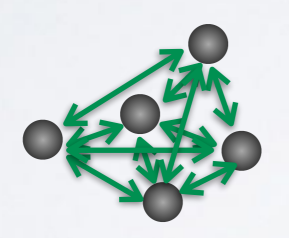
Empirical force field

$$E(\vec{R}) = \sum_{\text{bonded}} E_i(\vec{R}) + \sum_{\text{non-bonded}} E_i(\vec{R})$$

Divide time into discrete (~1fs) time steps (Δt) (for integrating equations of motion, see below)



At each time step calculate pair-wise atomic forces (F(t)) (by evaluating force-field gradient)



Nucleic motion described classically
$$m_i rac{d^2}{dt^2} ec{R}_i = - ec{
abla}_i E(ec{R})$$

Empirical force field $E(ec{R}) = \sum_{ ext{bonded}} E_i(ec{R}) + \sum_{ ext{non-bonded}} E_i(ec{R})$

Use the forces to calculate velocities and move atoms to new positions (by integrating numerically via the "leapfrog" scheme)

$$\mathbf{v}(t + \frac{\Delta t}{2}) = \mathbf{v}(t - \frac{\Delta t}{2}) + \frac{\mathbf{F}(t)}{m} \Delta t$$

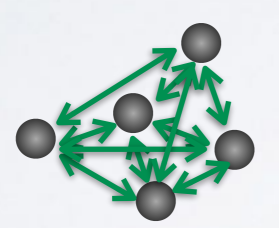
$$\mathbf{r}(t + \Delta t) = \mathbf{r}(t) + \mathbf{v}(t + \frac{\Delta t}{2}) \Delta t$$

BASIC ANATOMY OF A MD SIMULA

Divide time into discrete (~1fs) time steps (Δt) (for integrating equations of motion, see below)

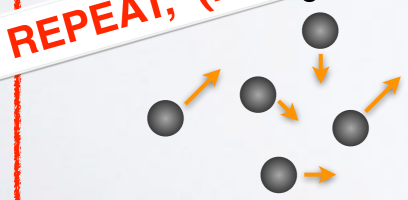


 \triangleright At each time step calculate pair-wise atomic forces (F(t)) (by evaluating **force-field** gradient)



$$m_i rac{d^2}{dt^2} ec{R}_i = - ec{
abla}_i E(ec{R})$$

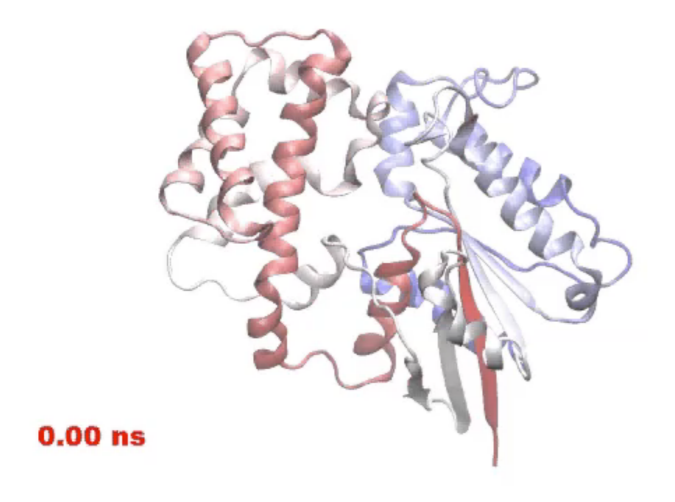
Use the formany times... $E(\vec{R}) = \sum_{i=1}^{m} \sum_{j=1}^{m} \sum_{k=1}^{m} \sum_{k=1}^{m} \sum_{j=1}^{m} \sum_{j=1}^{m} \sum_{j=1}^{m} \sum_{k=1}^{m} \sum_{j=1}^{m} \sum_{j=$



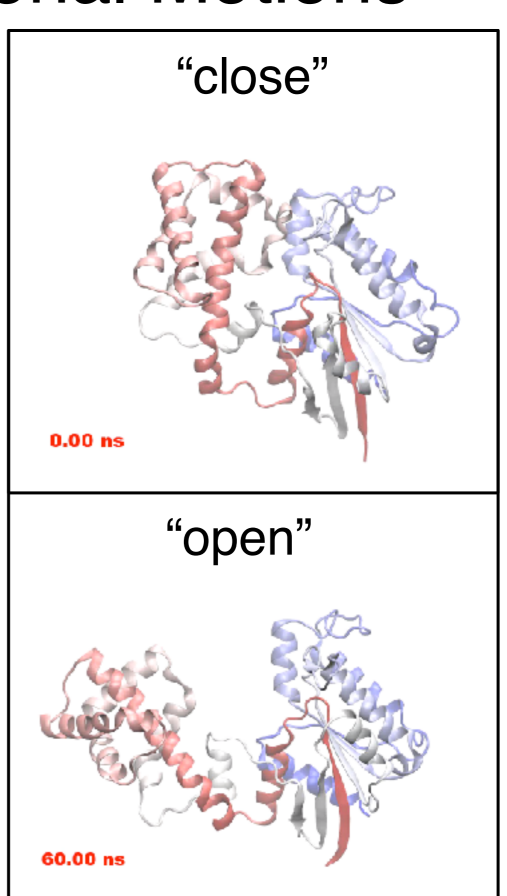
$$egin{array}{cccc} oldsymbol{v}(t+rac{\Delta t}{2}) & = & oldsymbol{v}(t-rac{\Delta t}{2})+rac{oldsymbol{F}(t)}{m}\Delta t \ oldsymbol{r}(t+\Delta t) & = & oldsymbol{r}(t)+oldsymbol{v}(t+rac{\Delta t}{2})\Delta t \end{array}$$

MD Prediction of Functional Motions

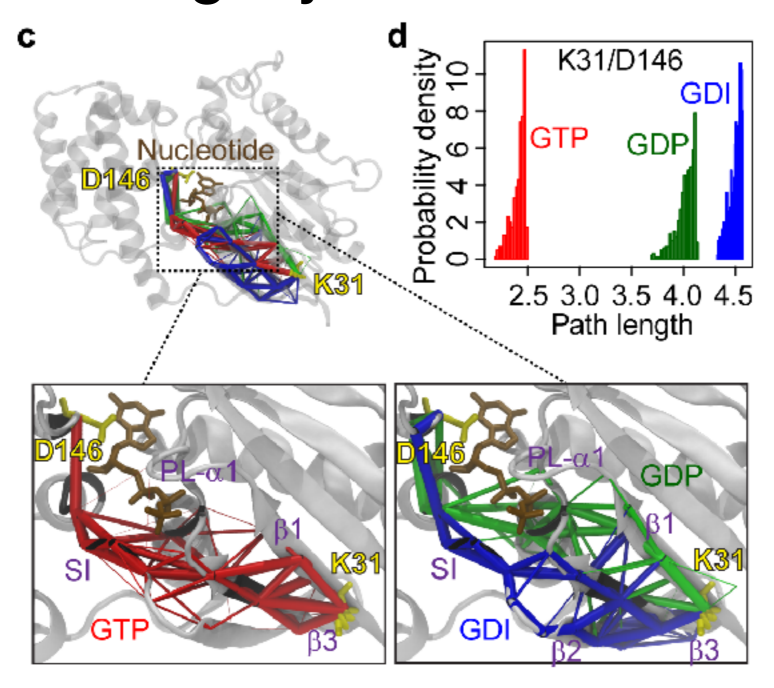
Accelerated MD simulation of nucleotide-free transducin alpha subunit



Yao and Grant, Biophys J. (2013)

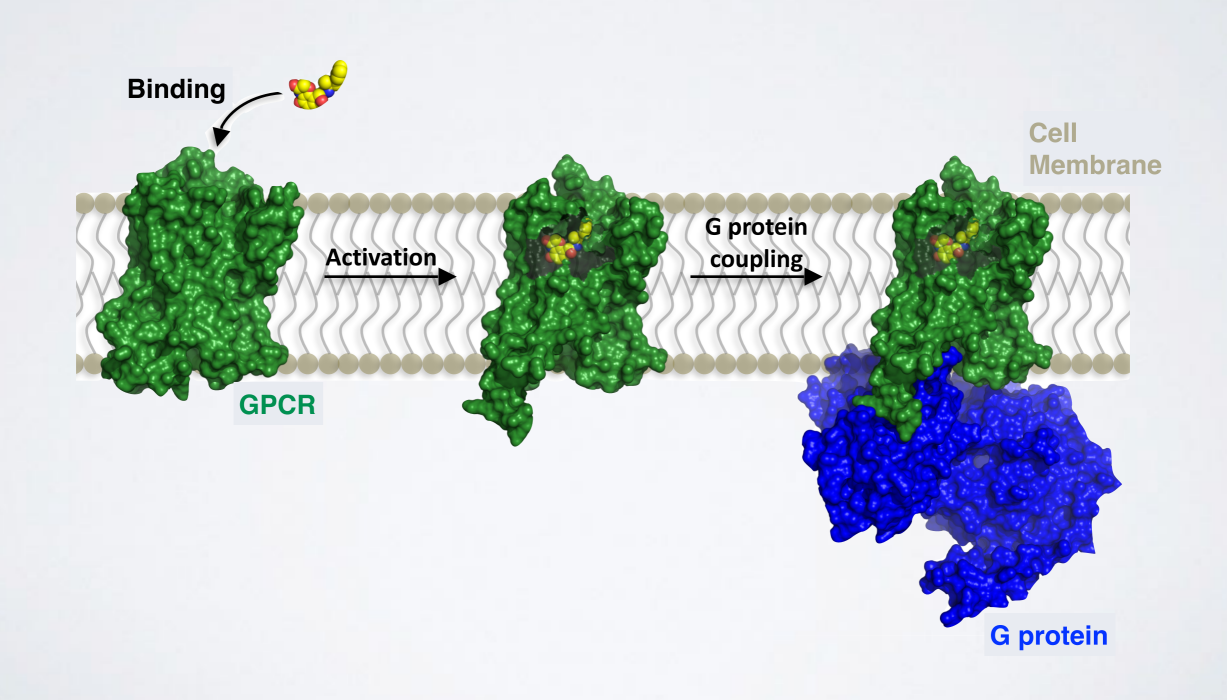


Simulations Identify Key Residues Mediating Dynamic Activation

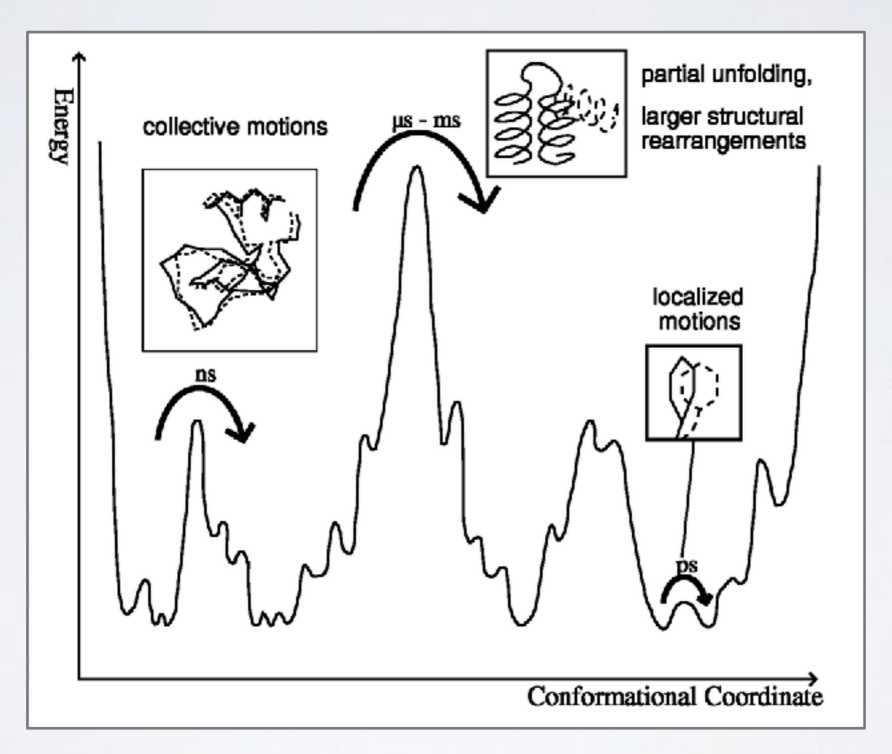


Yao ... Grant, <u>Journal of Biological Chemistry</u> (2016)

EXAMPLE APPLICATION OF MOLECULAR SIMULATIONS TO GPCRS



PROTEINS JUMP BETWEEN MANY, HIERARCHICALLY ORDERED "CONFORMATIONAL SUBSTATES"



H. Frauenfelder et al., *Science* **229** (1985) 337

MOLECULAR DYNAMICS IS VERY EXPENSIVE

Example: F₁-ATPase in water (183,674 atoms) for 1 nanosecond:

=> 10⁶ integration steps

=> 8.4 * 10¹¹ floating point operations/step [n(n-1)/2 interactions]

Total: 8.4 * 10¹⁷ flop

(on a 100 Gflop/s cpu: ca 25 years!)

... but performance has been improved by use of:

multiple time stepping ca. 2.5 years

fast multipole methods ca. 1 year

parallel computers ca. 5 days

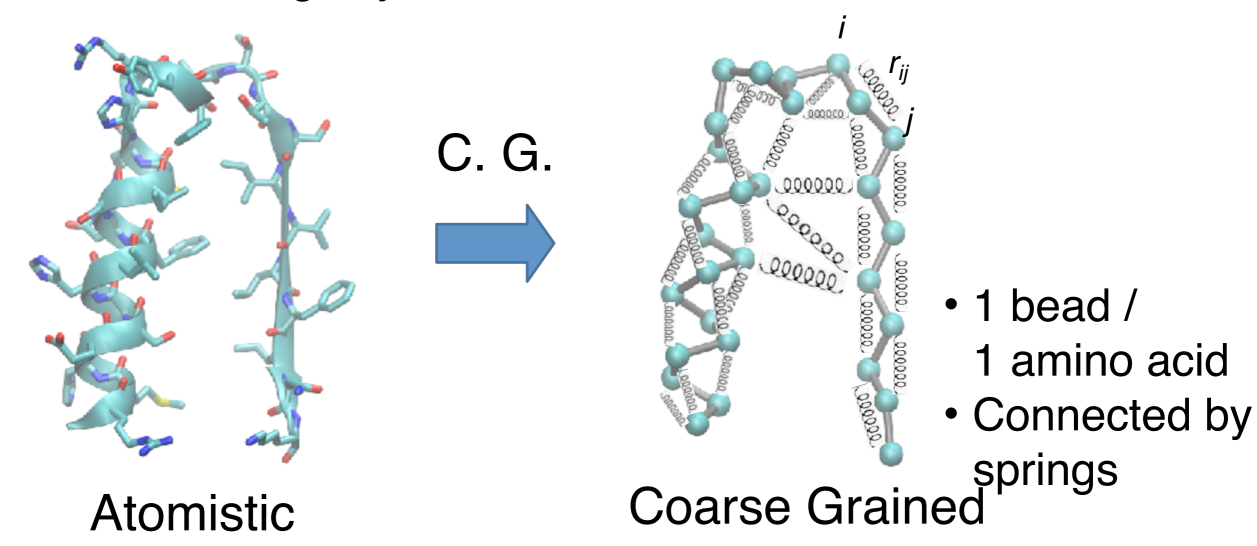
modern GPUs ca. 1 day

(Anton supercomputer ca. minutes)

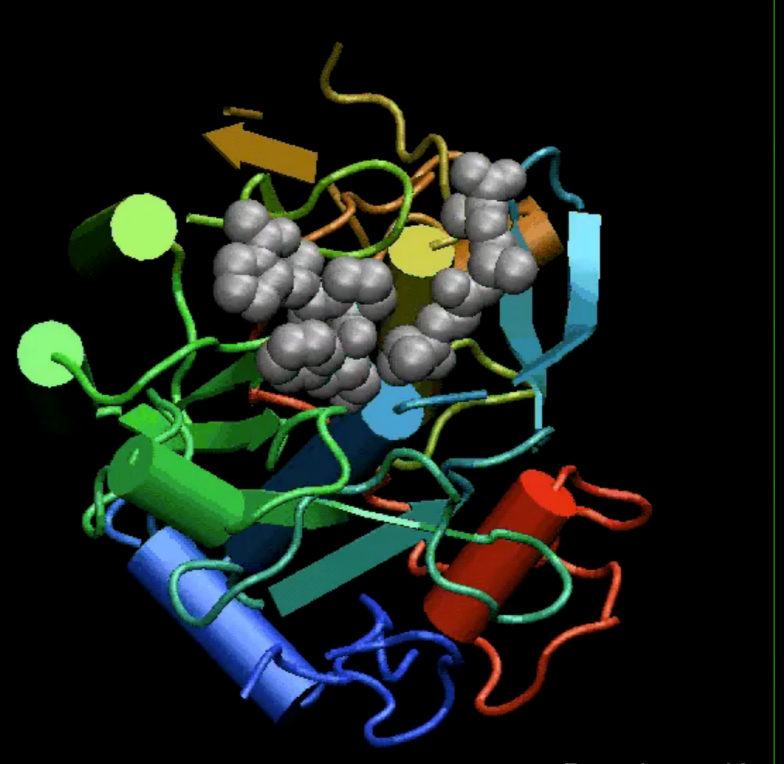
COARSE GRAINING: NORMAL MODE ANALYSIS

(NMA)

- MD is still time-consuming for large systems
- Elastic network model NMA (ENM-NMA) is an example of a lower resolution approach that finishes in seconds even for large systems.



NMA models the protein as a network of elastic strings



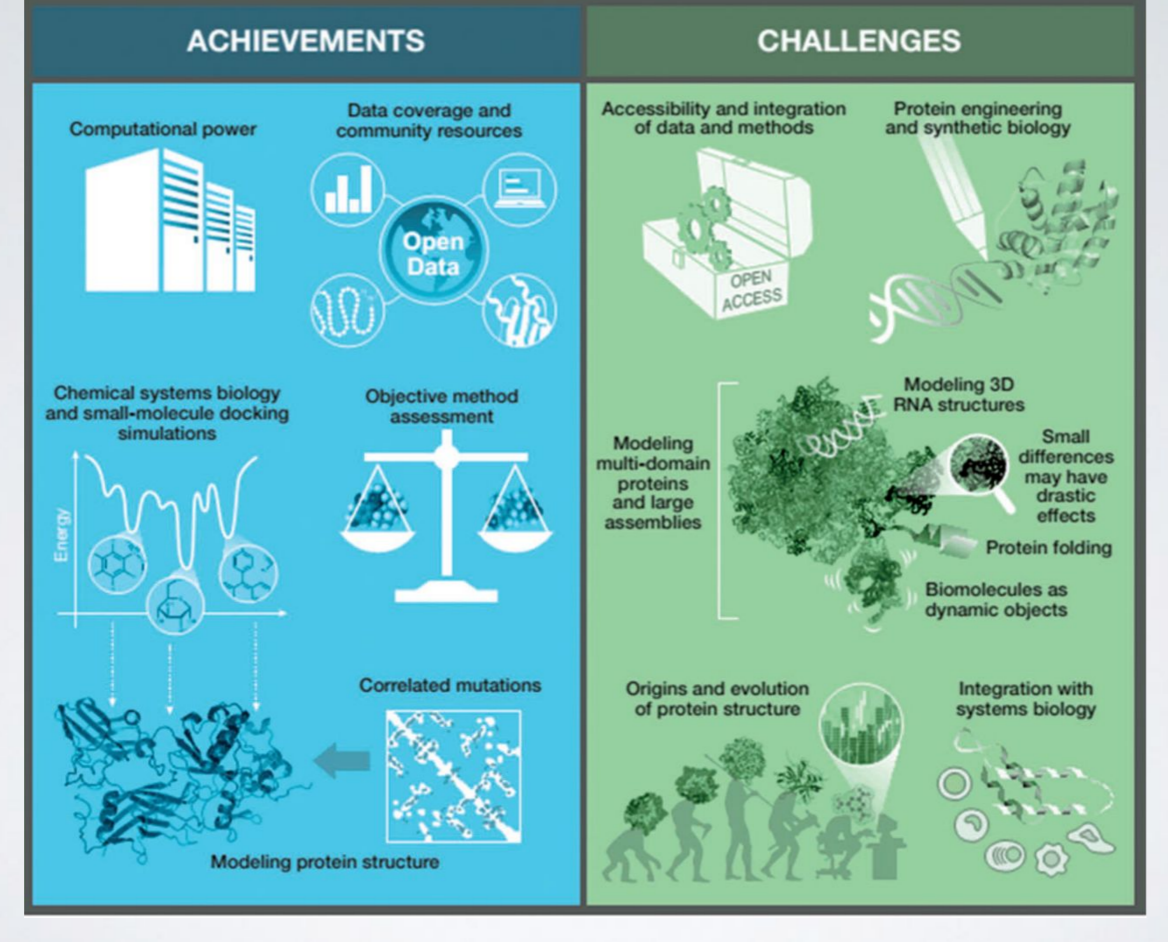
Proteinase K

DO 11 TOURS OF THE SERVICE OF THE SE

Hand-on time!

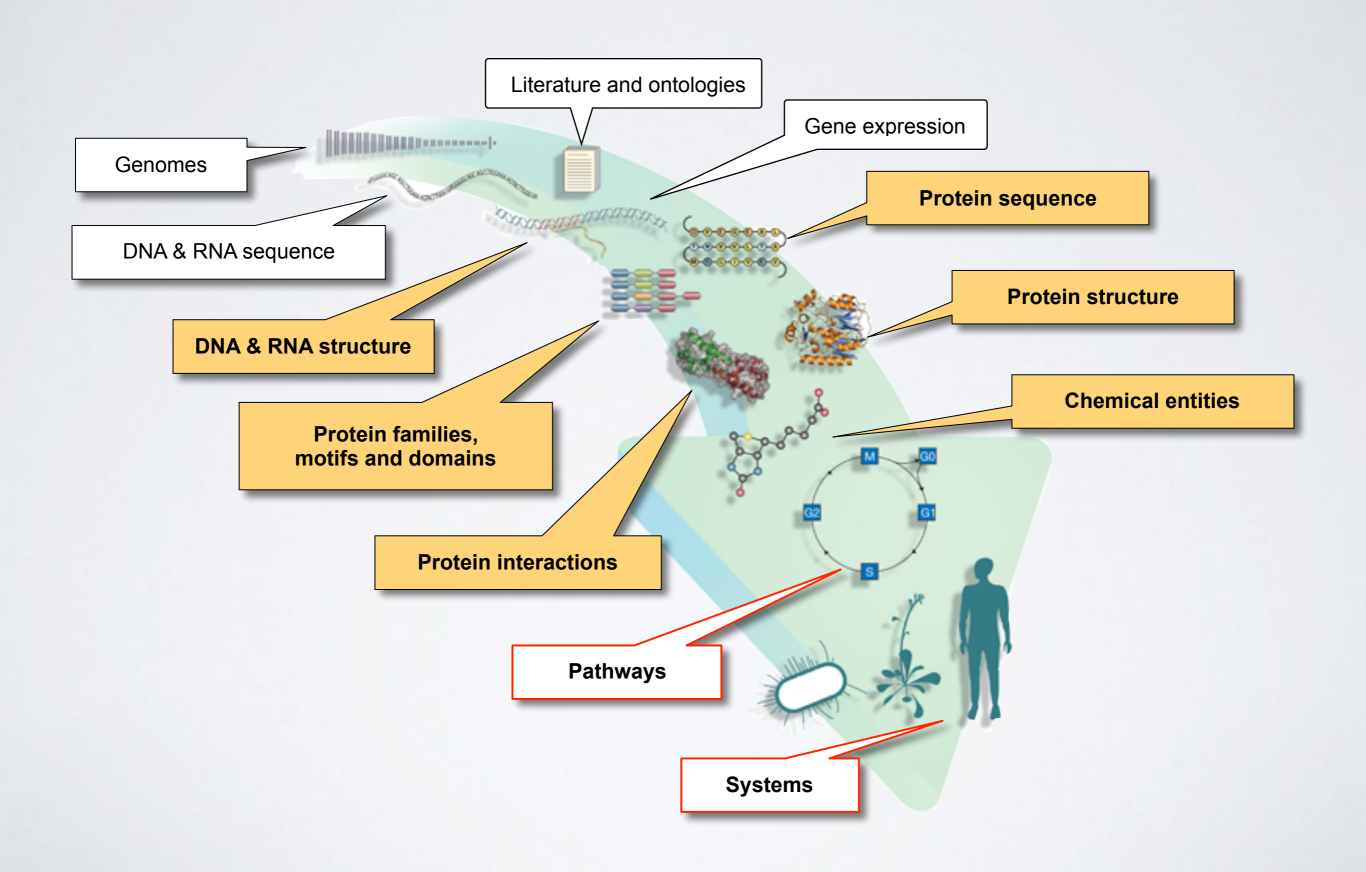
https://bioboot.github.io/bimm143_W18/lectures/#12

Focus on section 3 & 4 exploring PCA and NMA apps



Ilan Samish et al. Bioinformatics 2015;31:146-150

INFORMING SYSTEMS BIOLOGY?



SUMMARY

- Structural bioinformatics is computer aided structural biology
- Described major motivations, goals and challenges of structural bioinformatics
- Reviewed the fundamentals of protein structure
- Introduced both physics and knowledge based modeling approaches for describing the structure, energetics and dynamics of proteins computationally
- Introduced both structure and ligand based bioinformatics approaches for drug discovery and design