

BIMM 143
Structural Bioinformatics II

Lecture 12

Barry Grant
UC San Diego

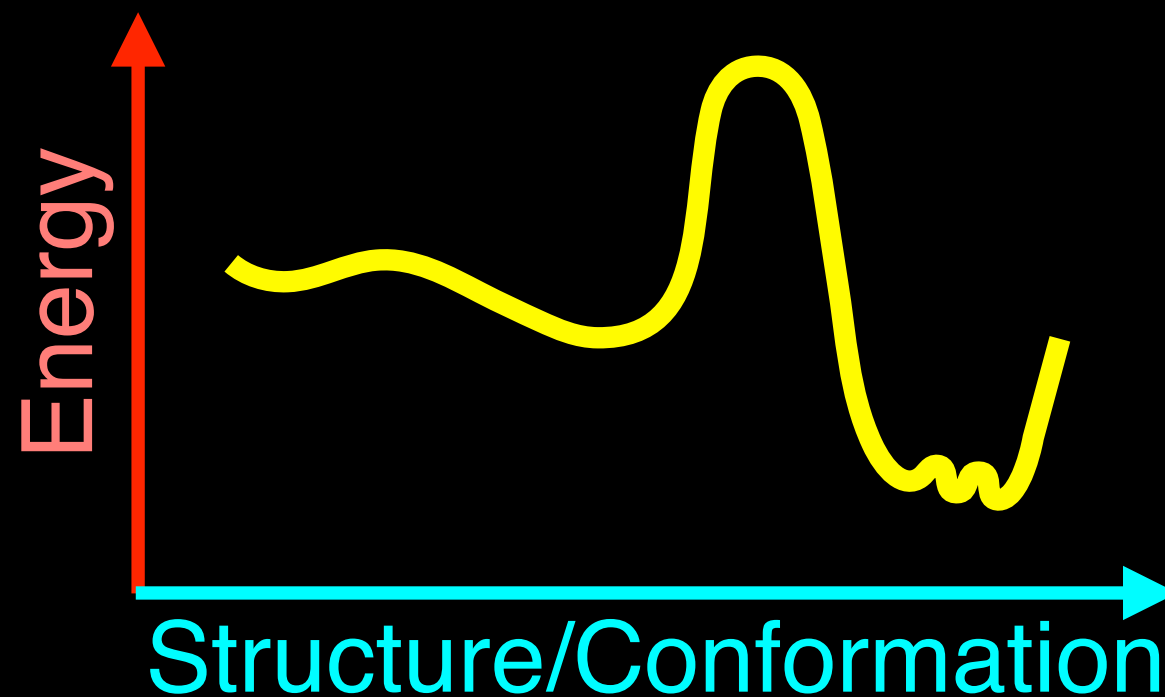
<http://thegrantlab.org/bimm143>

Next Up:

- **Overview of structural bioinformatics**
 - Motivations, goals and challenges
- **Fundamentals of protein structure**
 - Structure composition, form and forces
- **Representing, interpreting & modeling protein structure**
 - Visualizing and interpreting protein structures
 - Analyzing protein structures
 - Modeling energy as a function of structure
 - Drug discovery & Predicting functional dynamics

Key concept:

Potential functions describe a system's energy as a function of its structure



Two main approaches:

(1). Physics-Based

(2). Knowledge-Based

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(2). Knowledge-Based

For **physics** based potentials
energy terms come from physical theory

$$V(R) = E_{\text{bonded}} + E_{\text{non.bonded}}$$

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Sum of **bonded** and **non-bonded**
atom-type and position based terms

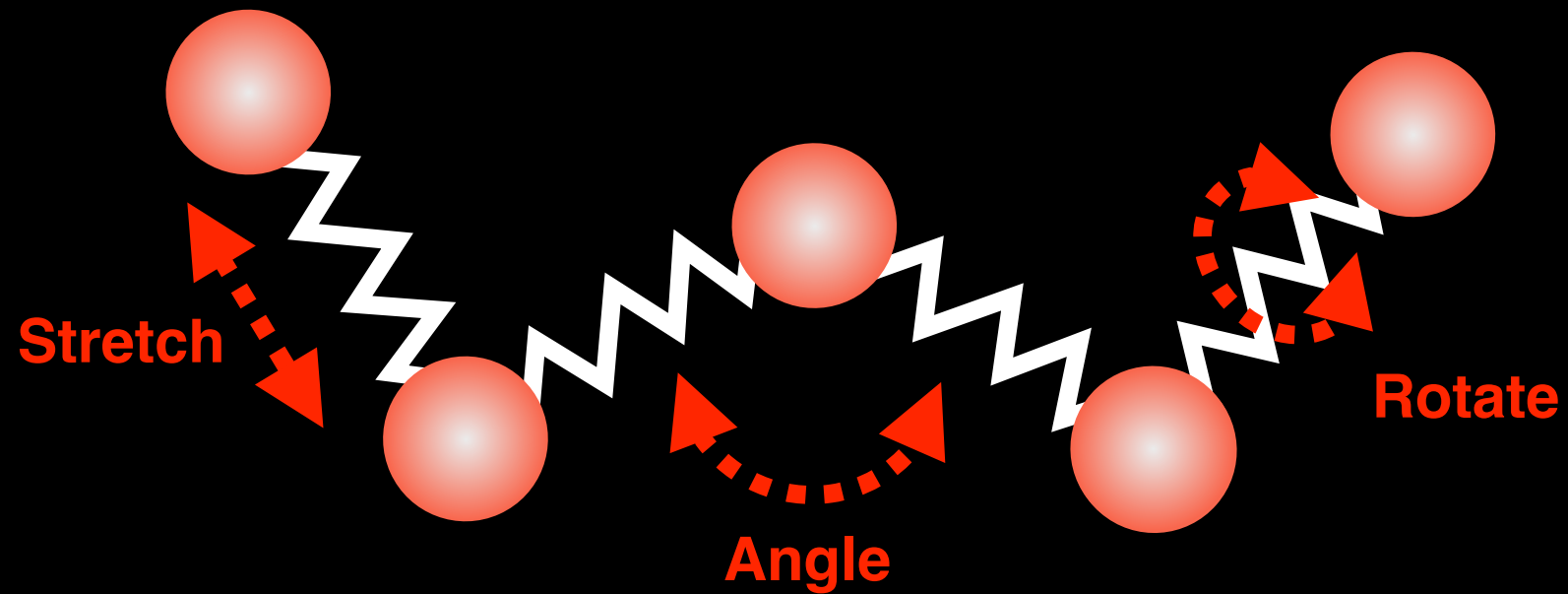
$$V(R) = E_{\text{bonded}} + E_{\text{non.bonded}}$$

E_{bonded} is itself a sum of three terms:

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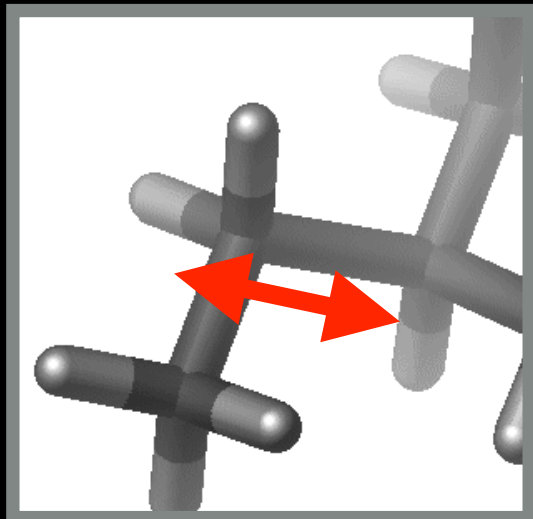
$$E_{\text{bond.stretch}} + E_{\text{bond.angle}} + E_{\text{bond.rotate}}$$



$$V(R) = E_{\text{bonded}} + E_{\text{non.bonded}}$$

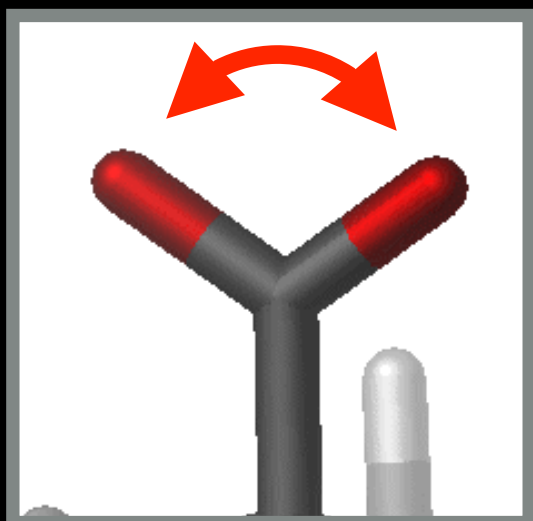
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$$E_{\text{bond.stretch}} + E_{\text{bond.angle}} + E_{\text{bond.rotate}}$$



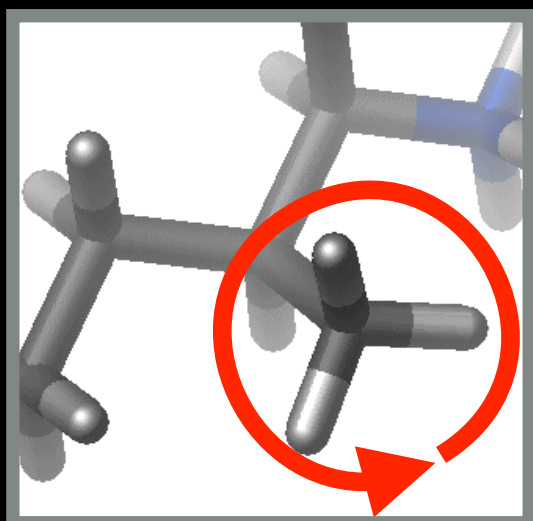
Bond Stretch

$E_{bond.stretch}$



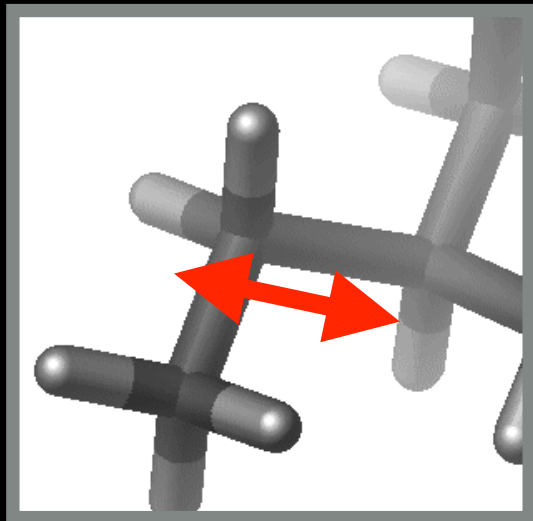
Bond Angle

$E_{bond.angle}$



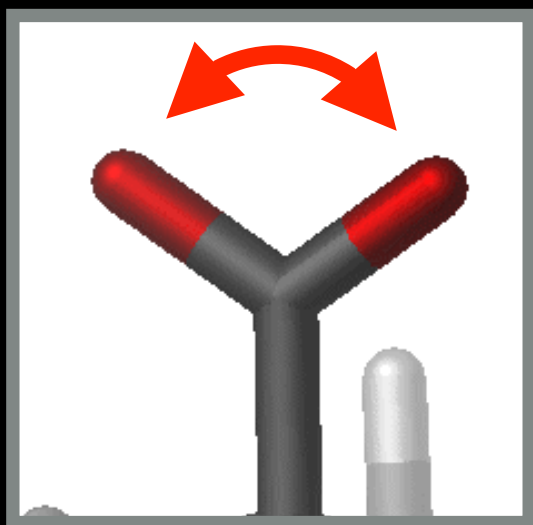
Bond Rotate

$E_{bond.rotate}$



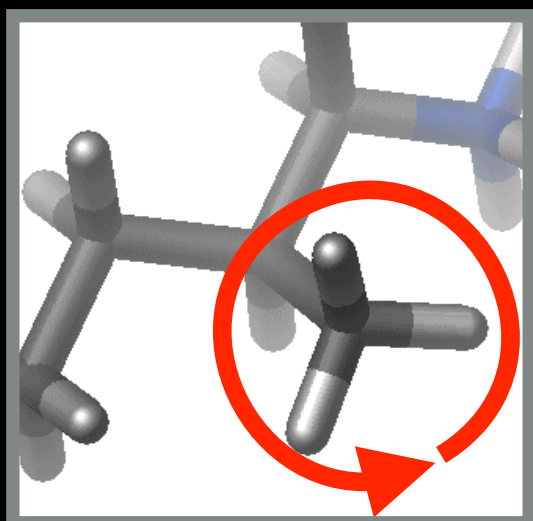
Bond Stretch

$$\sum_{\text{bonds}} K_i^{bs} (b_i - b_o)$$



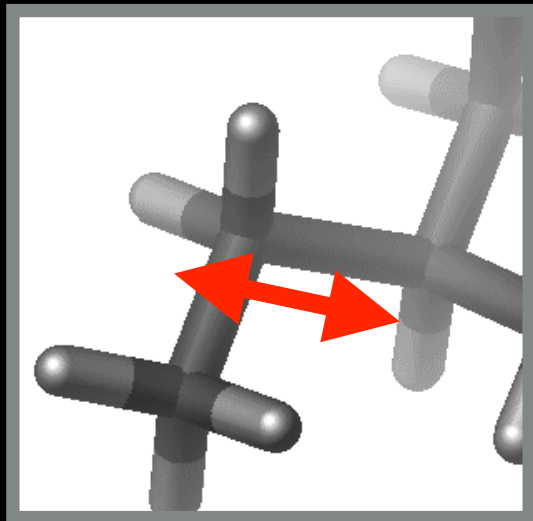
Bond Angle

$$\sum_{\text{angles}} K_i^{ba} (\theta_i - \theta_o)$$



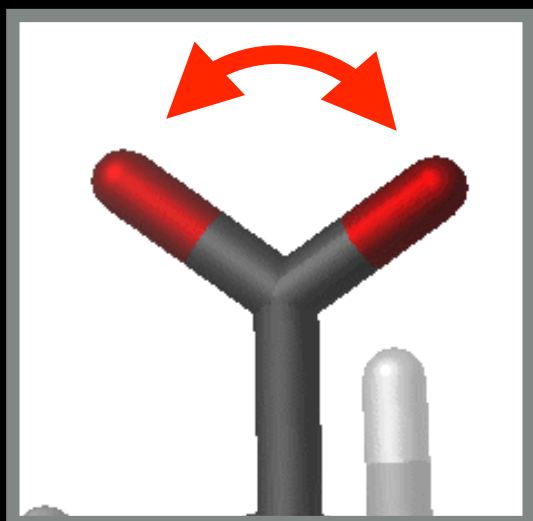
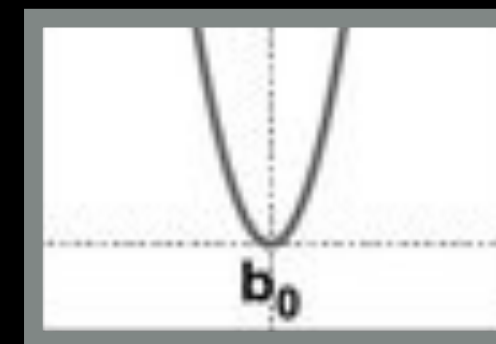
Bond Rotate

$$\sum_{\text{dihedrals}} K_i^{br} [1 - \cos(n_i \phi_i - \phi_o)]$$



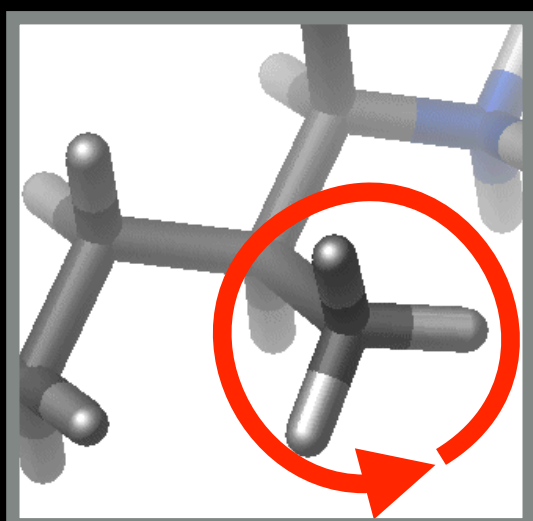
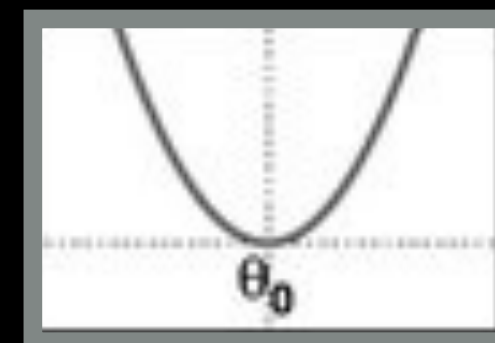
Bond Stretch

$$\sum_{\text{bonds}} K_i^{bs} (b_i - b_o)$$



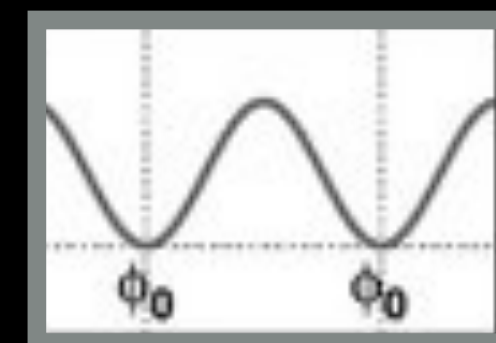
Bond Angle

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Bond Rotate

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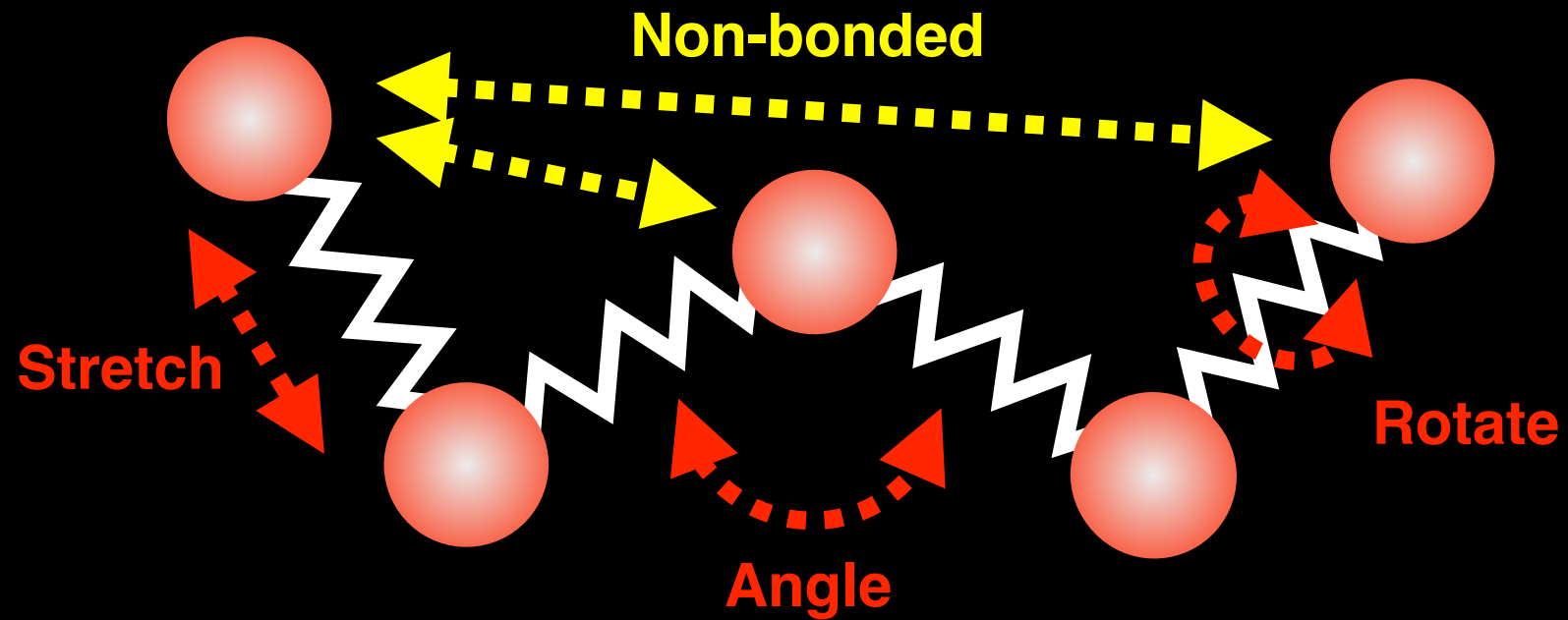
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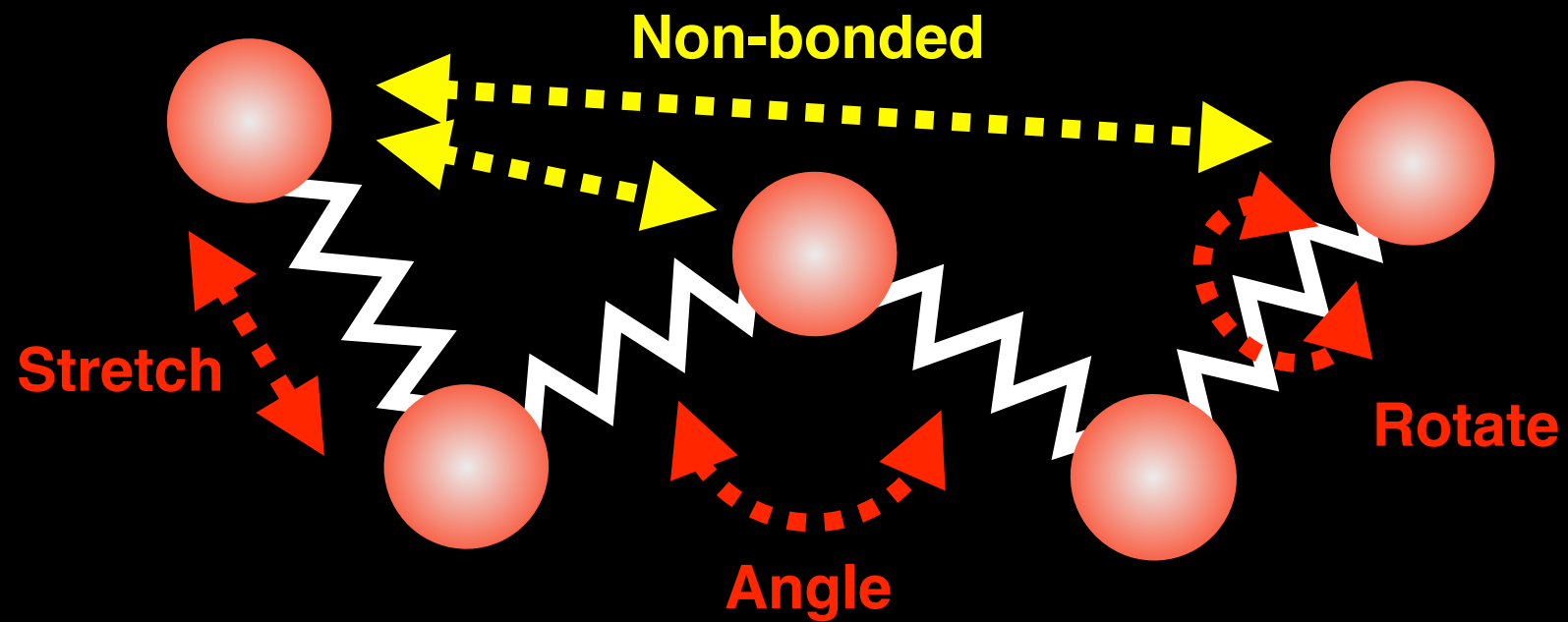
$$E_{\text{van.der.Waals}} + E_{\text{electrostatic}}$$



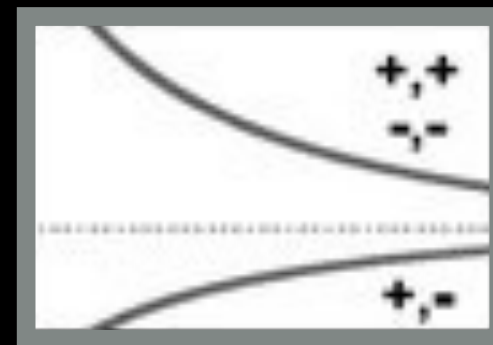
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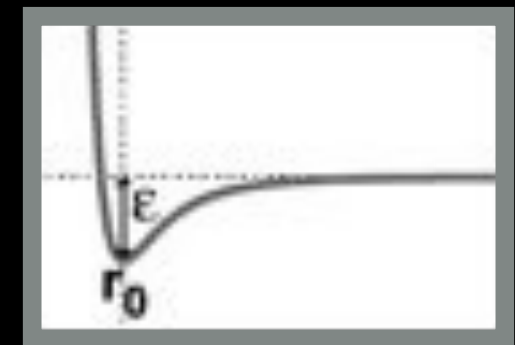
$$E_{\text{van.der.Waals}} + E_{\text{electrostatic}}$$



$$E_{electrostatic} = \sum_{pairs.i.j} \frac{q_i q_j}{\epsilon r_{ij}}$$



$$E_{van.der.Waals} = \sum_{pairs.i.j} \left[\epsilon_{ij} \left(\frac{r_{o.ij}}{r_{ij}} \right)^{12} - 2\epsilon_{ij} \left(\frac{r_{o.ij}}{r_{ij}} \right)^6 \right]$$



Total potential energy

The potential energy can be given as a sum of terms for: Bond stretching, Bond angles, Bond rotations, van der Waals and Electrostatic interactions between atom pairs

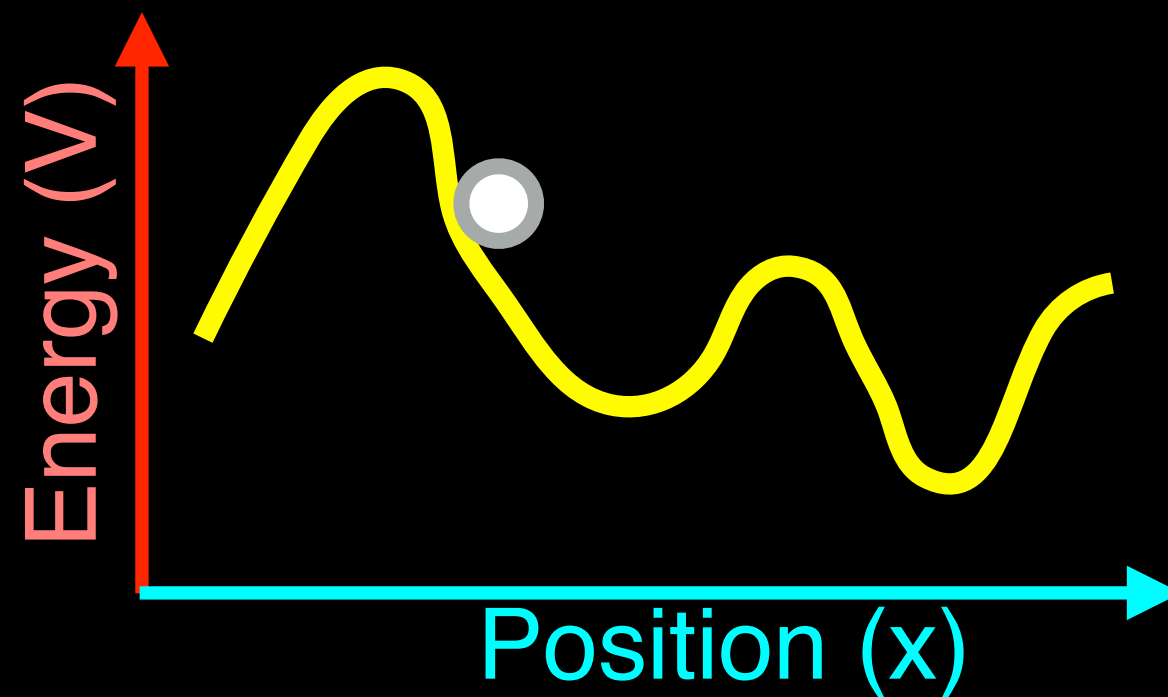
$$V(R) = E_{\text{bond.stretch}} + E_{\text{bond.angle}} + E_{\text{bond.rotate}} + E_{\text{van.der.Waals}} + E_{\text{electrostatic}}$$

E_{bonded}

$E_{\text{non.bonded}}$

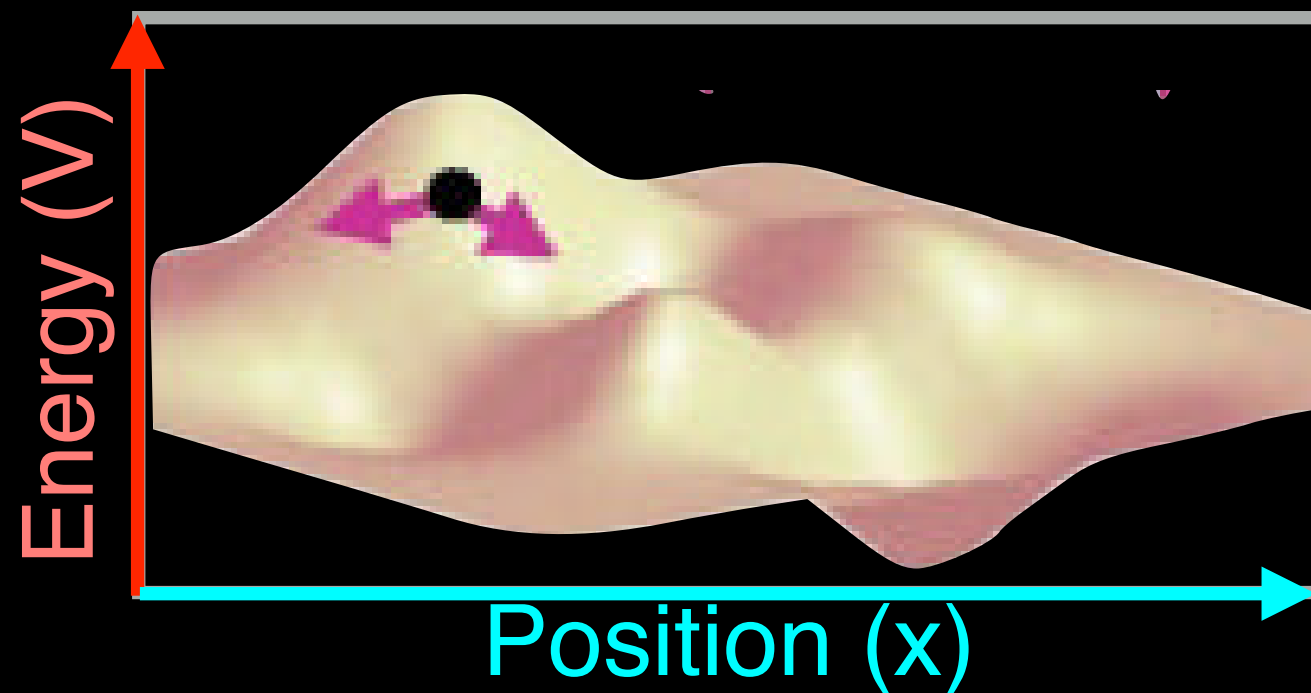
Potential energy surface

Now we can calculate the **potential energy surface** that fully describes the energy of a molecular system as a function of its geometry



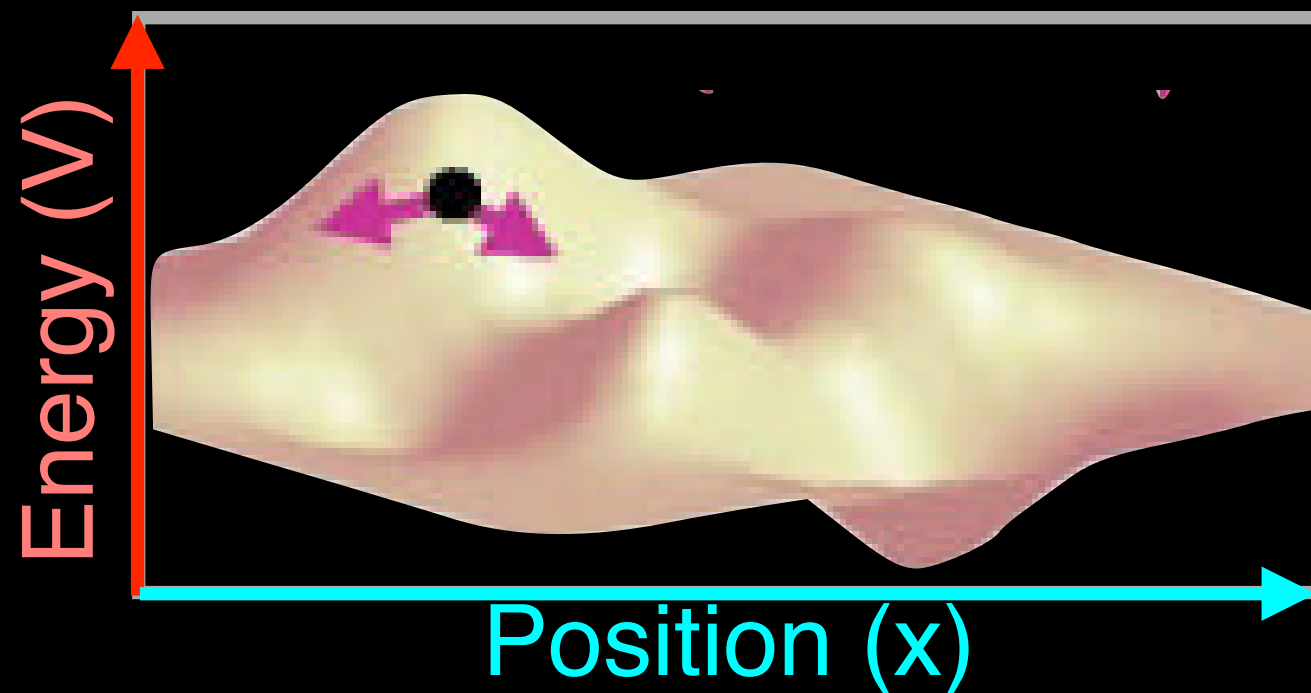
Potential energy surface

Now we can calculate the **potential energy surface** that fully describes the energy of a molecular system as a function of its geometry



Key concept:

Now we can calculate the **potential energy surface** that fully describes the energy of a molecular system as a function of its geometry



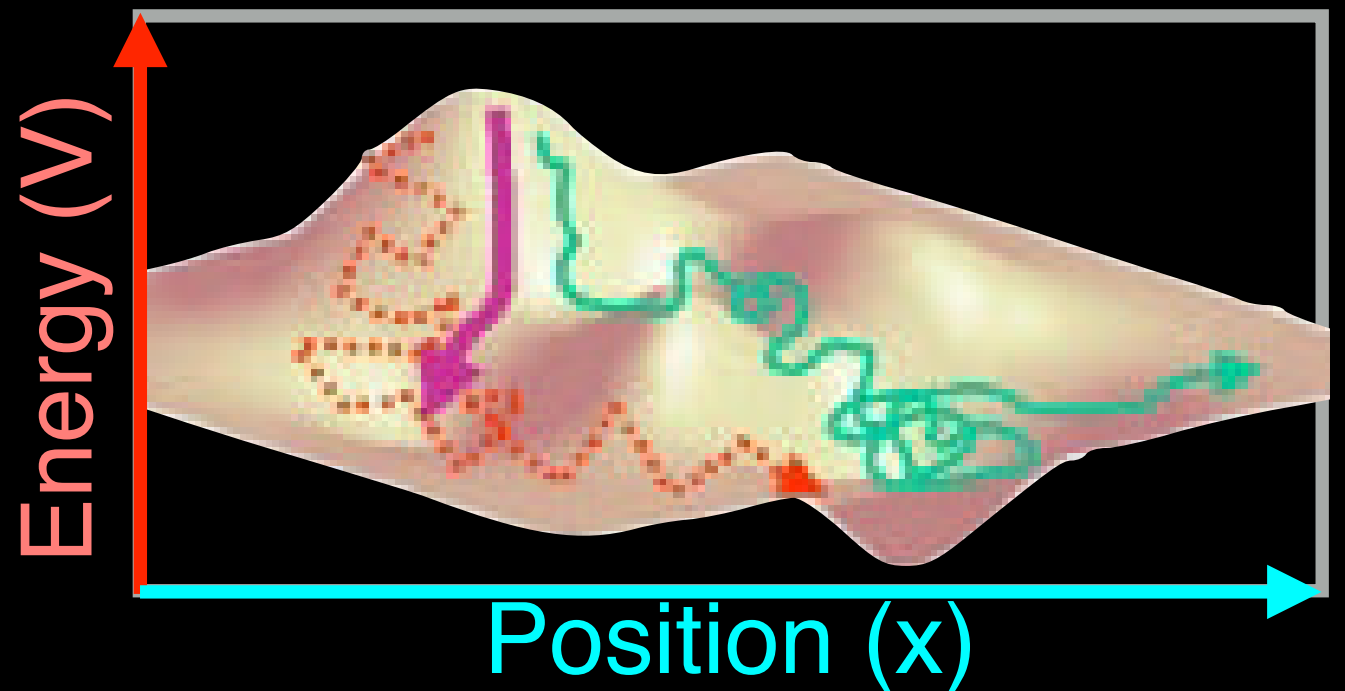
- The **forces** are the gradients of the energy

$$F(x) = -dV/dx$$

Moving Over The Energy Surface

- **Energy Minimization** drops into local minimum
- **Molecular Dynamics** uses thermal energy to move smoothly over surface
- **Monte Carlo Moves** are random. Accept with probability:

$$\exp(-\Delta V/dx)$$



PHYSICS-ORIENTED APPROACHES

Weaknesses

Fully physical detail becomes computationally intractable

Approximations are unavoidable

(Quantum effects approximated classically, water may be treated crudely)

Parameterization still required

Strengths

Interpretable, provides guides to design

Broadly applicable, in principle at least

Clear pathways to improving accuracy

Status

Useful, widely adopted but far from perfect

Multiple groups working on fewer, better approxs

Force fields, quantum

entropy, water effects

Moore's law: hardware improving

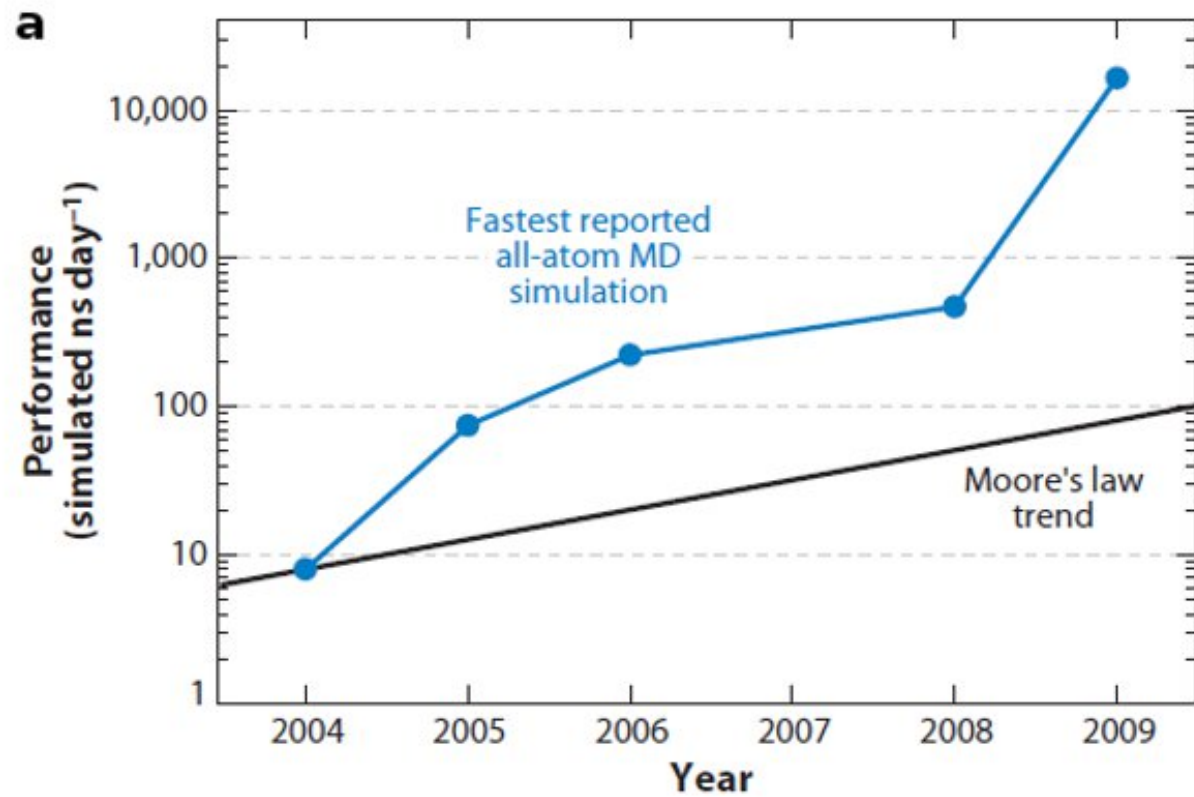
HOW COMPUTERS HAVE CHANGED

DATE	COST	SPEED	MEMORY	SIZE
1967	\$40M	0.1 MHz	1 MB	HALL
2013	\$4,000	1 GHz	10 GB	LAPTOP
CHANGE	10,000	10,000	10,000	10,000

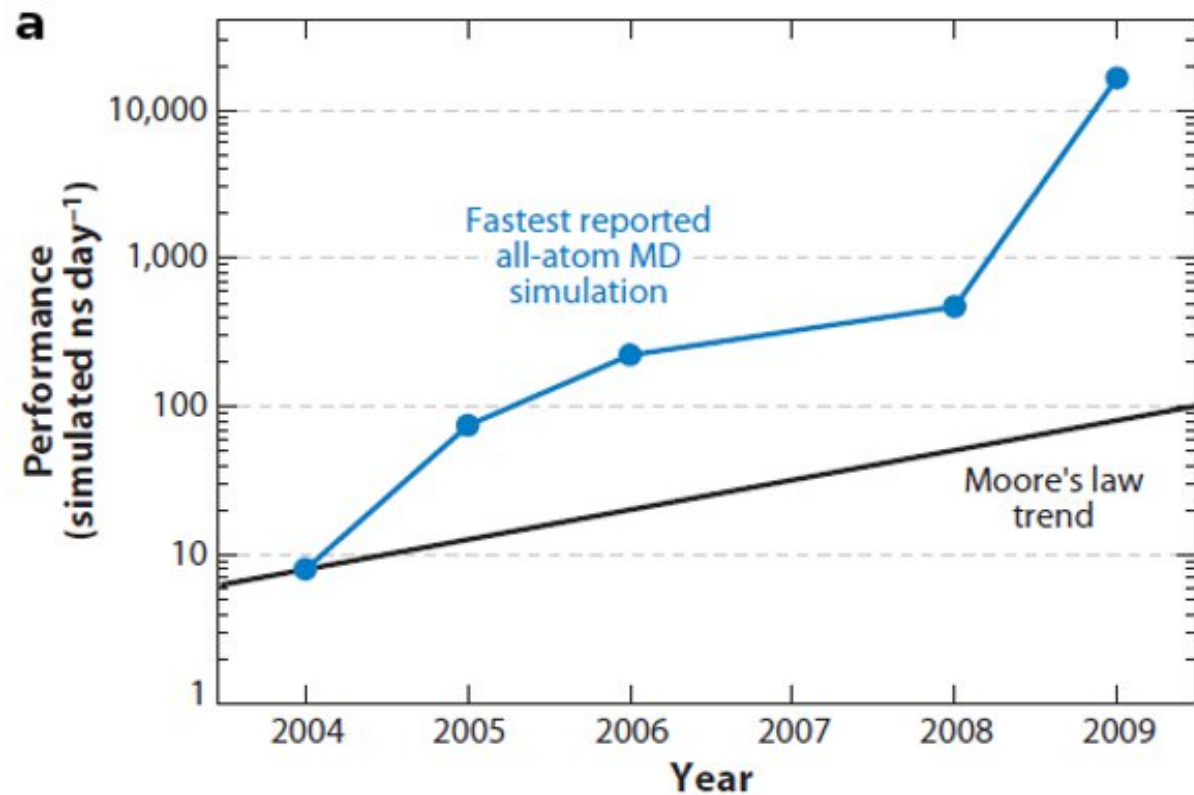
If cars were like computers then a new Volvo would cost \$3, would have a top speed of 1,000,000 km/hr, would carry 50,000 adults and would park in a shoebox



SIDE-NOTE: GPUS AND ANTON SUPERCOMPUTER



SIDE-NOTE: GPUS AND ANTON SUPERCOMPUTER



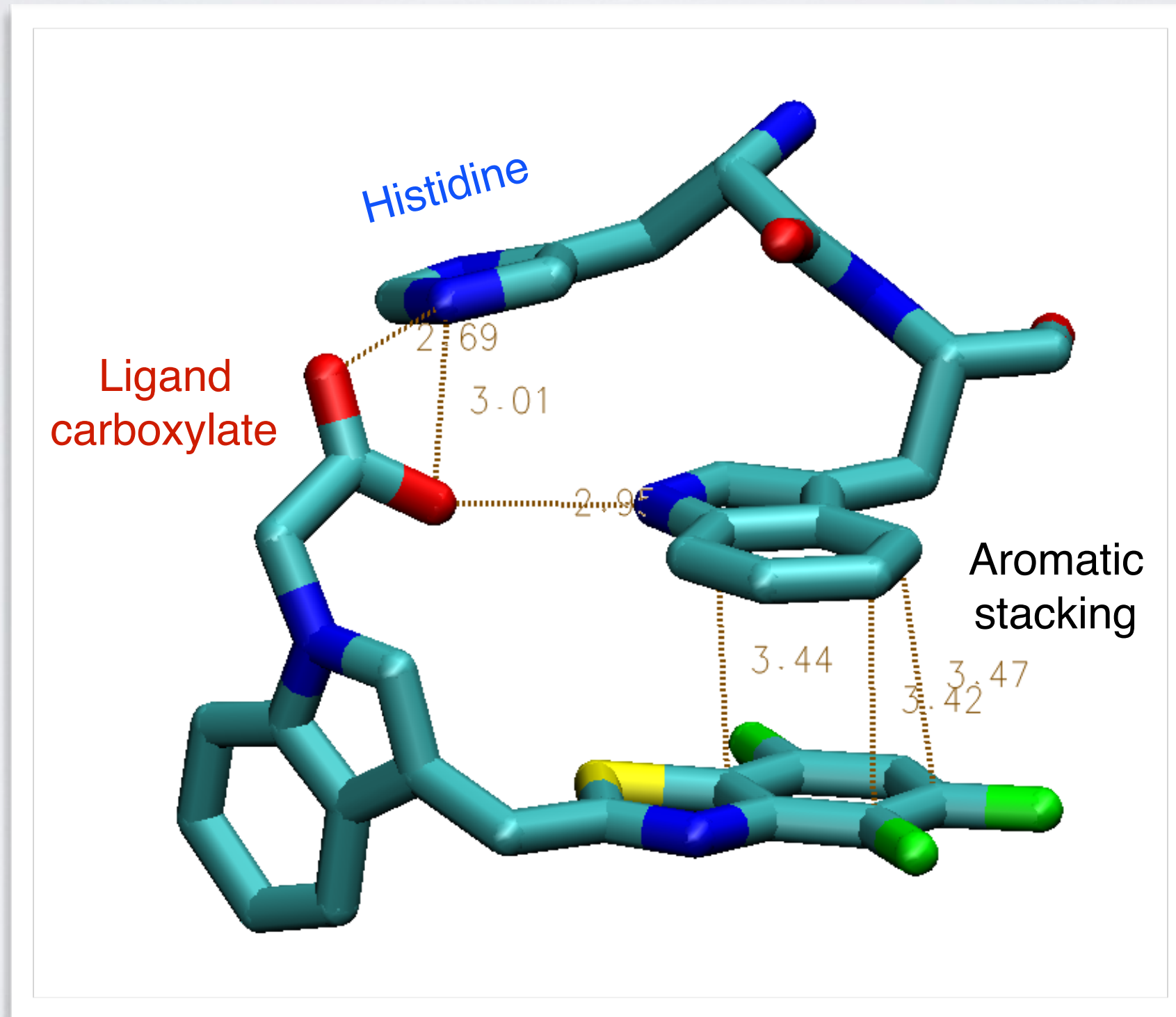
POTENTIAL FUNCTIONS DESCRIBE A SYSTEMS
ENERGY AS A FUNCTION OF ITS **STRUCTURE**

Two main approaches:

(1). Physics-Based

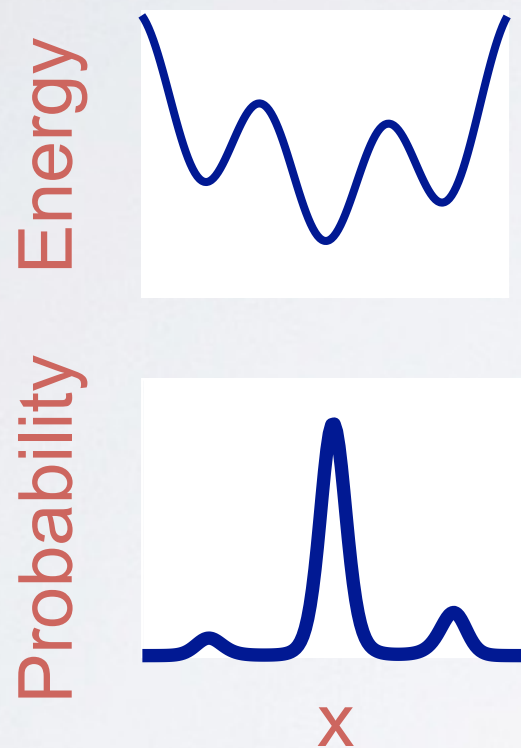
(2). Knowledge-Based

KNOWLEDGE-BASED DOCKING POTENTIALS



ENERGY DETERMINES **PROBABILITY** (STABILITY)

Basic idea: Use probability as a proxy for energy



Boltzmann:

$$p(r) \propto e^{-E(r)/RT}$$

Inverse Boltzmann:

$$E(r) = -RT \ln [p(r)]$$

Example: ligand **carboxylate O** to protein **histidine N**

Find all protein-ligand structures in the PDB with a ligand carboxylate **O**

1. For each structure, histogram the distances from **O** to every histidine **N**
2. Sum the histograms over all structures to obtain $p(r_{\text{O-N}})$
3. Compute $E(r_{\text{O-N}})$ from $p(r_{\text{O-N}})$

KNOWLEDGE-BASED POTENTIALS

Weaknesses

Accuracy limited by availability of data

Strengths

Relatively easy to implement

Computationally fast

Status

Useful, far from perfect

May be at point of diminishing returns

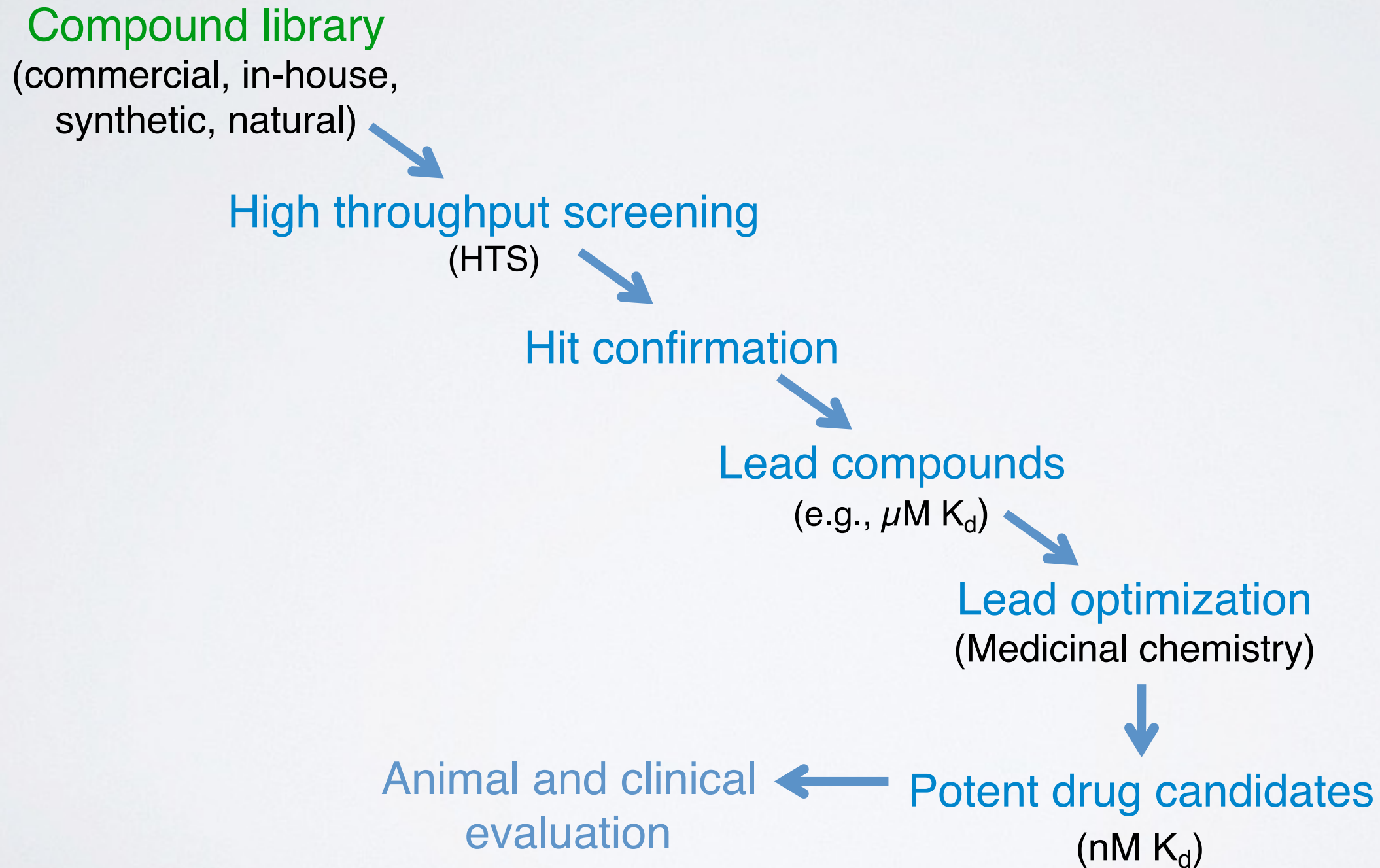
(not always clear how to make improvements)

Computer Aided Drug Discovery

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 - Drug discovery & Predicting functional dynamics

THE TRADITIONAL EMPIRICAL PATH TO DRUG DISCOVERY



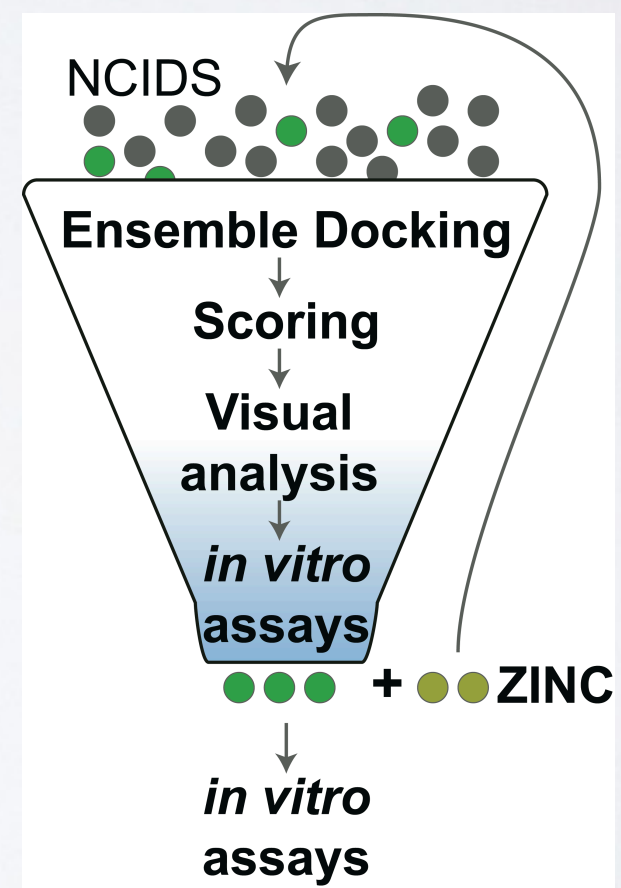
COMPUTER-AIDED LIGAND DESIGN

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

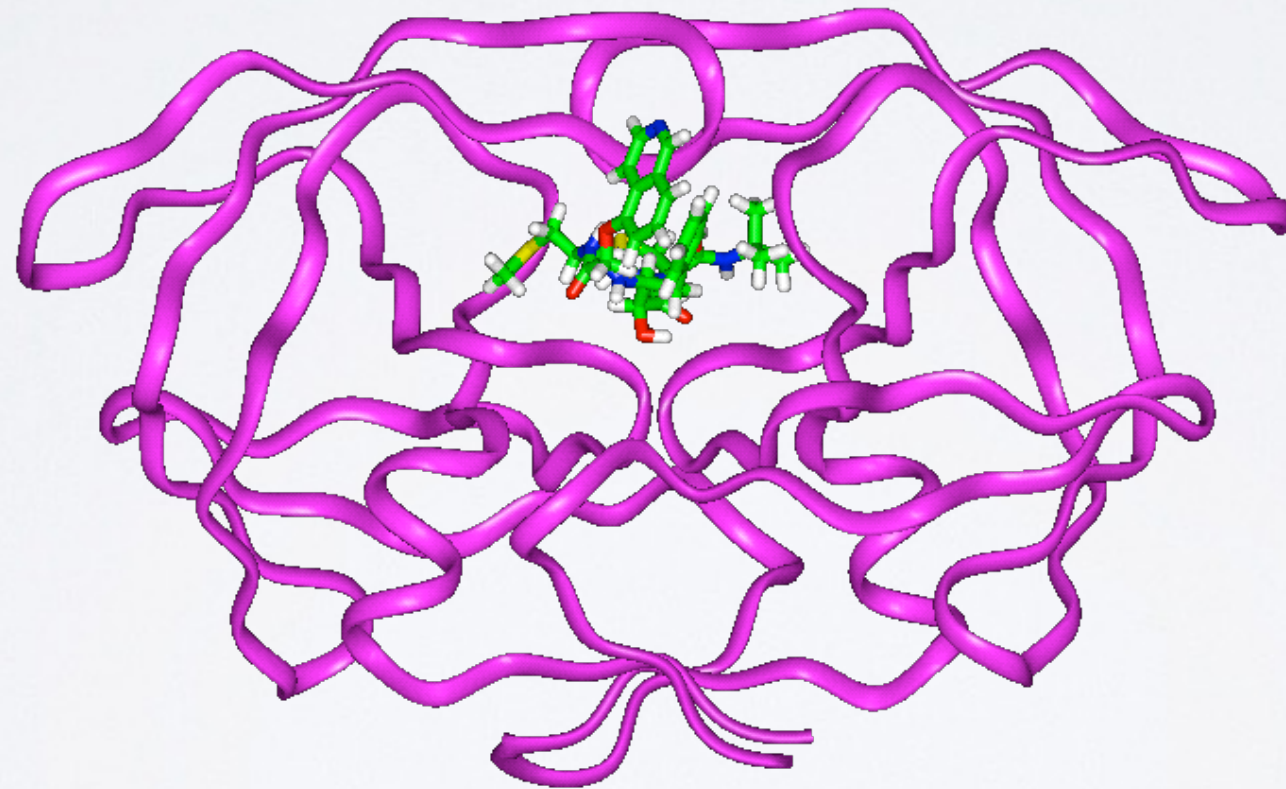
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SCENARIO I: RECEPTOR-BASED DRUG DISCOVERY

Structure of Targeted Protein Known: **Structure-Based Drug Discovery**



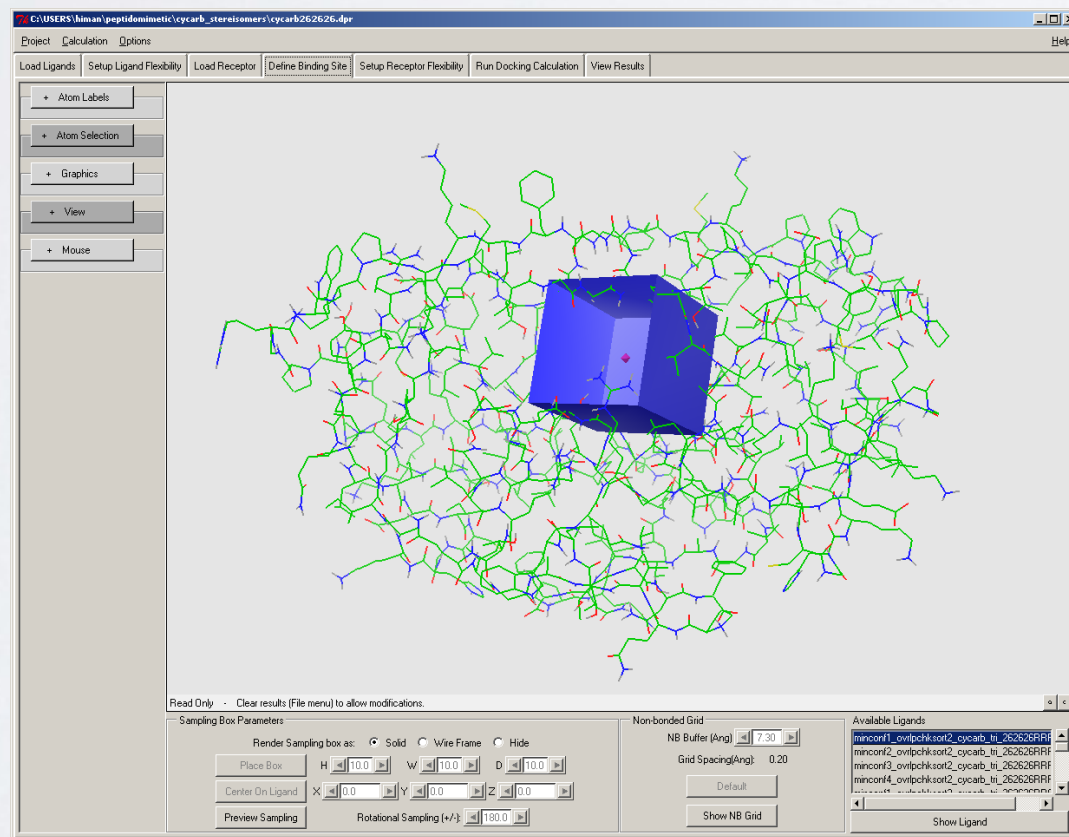
HIV Protease/KNI-272 complex

PROTEIN-LIGAND DOCKING

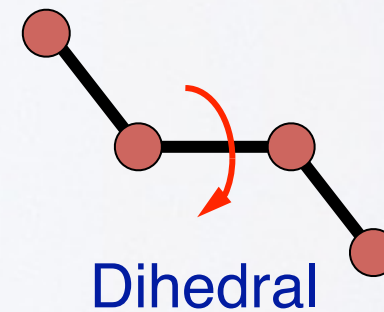
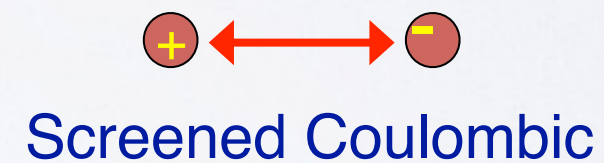
Structure-Based Ligand Design

Docking software

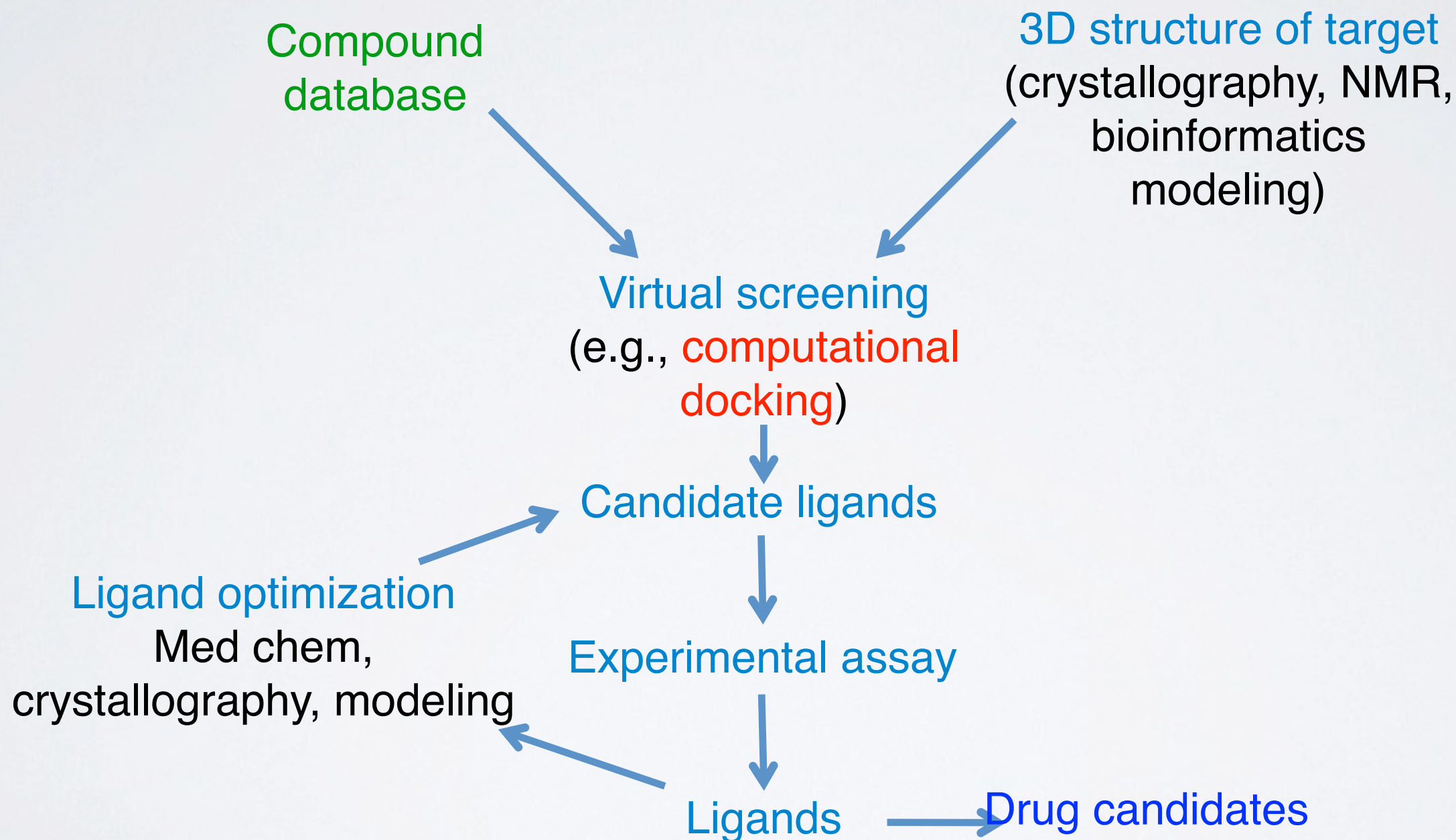
Search for structure of lowest energy



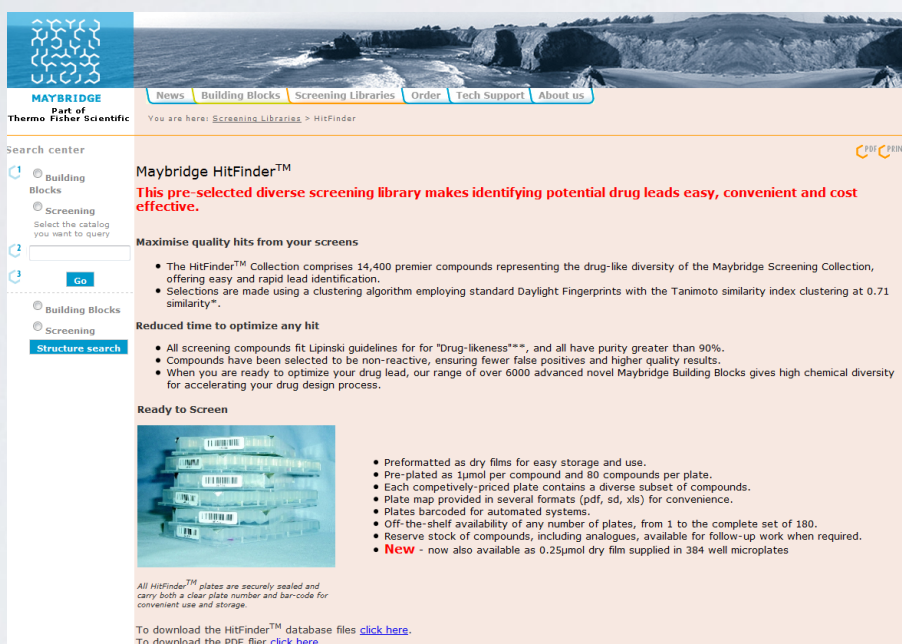
Potential function
Energy as function of structure



STRUCTURE-BASED VIRTUAL SCREENING



COMPOUND LIBRARIES



The screenshot shows the Maybridge HitFinder website, which is part of Thermo Fisher Scientific. The page features a navigation menu with links for News, Building Blocks, Screening Libraries, Order, Tech Support, and About us. The main content area is titled "Maybridge HitFinder™" and describes a pre-selected diverse screening library. It highlights the library's size (14,400 compounds) and its focus on drug-like diversity. A "Ready to Screen" section lists key features such as preformatted dry films, 1µmol per compound, and 80 compounds per plate. The page also includes a search center with options for Building Blocks, Screening, and Structure search.

Commercial
(in-house pharma)



The screenshot displays the NIH Molecular Libraries Small Molecule Repository website, operated by BioFocus, a Galapagos Company. The page is titled "A Roadmap Initiative" and features a "Welcome" message. It describes the repository's role in collecting samples for high-throughput biological screening and distributing them to the NIH Molecular Libraries Probe Production Centers Network. A "Registered Users Login" section is visible on the left. The page also includes a "BioFocus, a Galapagos company" logo and a copyright notice for 2007 Galapagos NV.

Government (NIH)



The screenshot shows the Pittsburgh Molecular Libraries Screening Center (PMLSC) website, part of the University of Pittsburgh. The page features a navigation menu with links for HOME, HISTORY, PERSONNEL, SCREENING TECHNOLOGY, COMPOUND LIBRARIES, INSTRUMENTATION/PLATFORMS, HTS GUIDELINES, APPROVED PMLSC ASSAY PROTOCOLS, PMLSC PROBE REPORTS, CHEMISTRY, DATA ANALYSIS/INFORMATICS, EDUCATIONAL ACTIVITIES, PUBLICATIONS, LINKS, and CONTACTS. The main content area is titled "PMLSC" and includes a "Welcome" message. It describes the center's mission to assist scientists and the National Institutes of Health in interrogating small molecule libraries using optical-based High Throughput and High Content assays. The page also includes a "Keyword Search" field and a "Go" button.

Academia

COMMON SIMPLIFICATIONS USED IN PHYSICS-BASED DOCKING

Quantum effects approximated classically

Protein often held rigid

Configurational entropy neglected

Influence of water treated crudely

Do it Yourself!

Hand-on time!

https://bioboot.github.io/bimm143_S19/lectures/#13

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

Two main approaches:

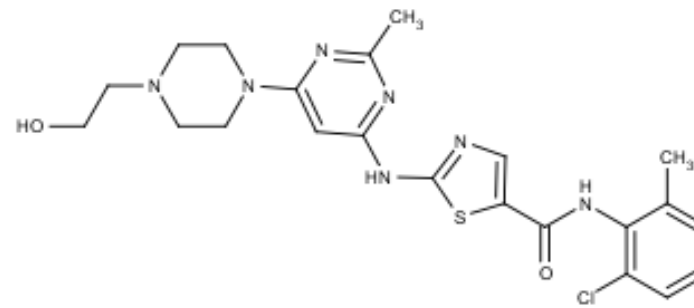
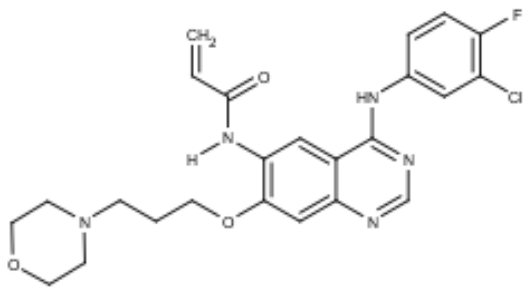
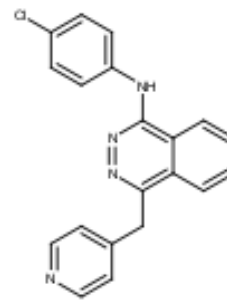
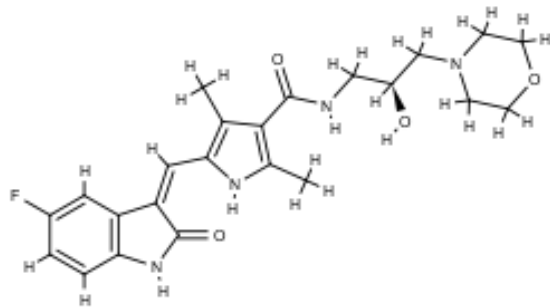
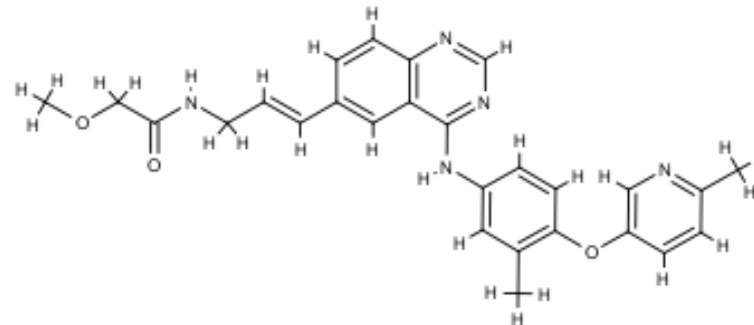
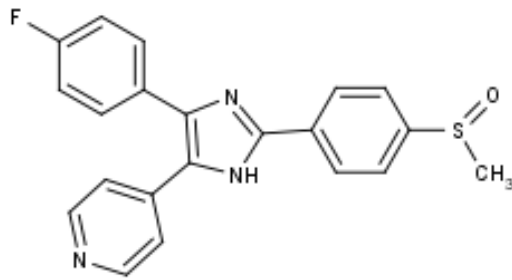
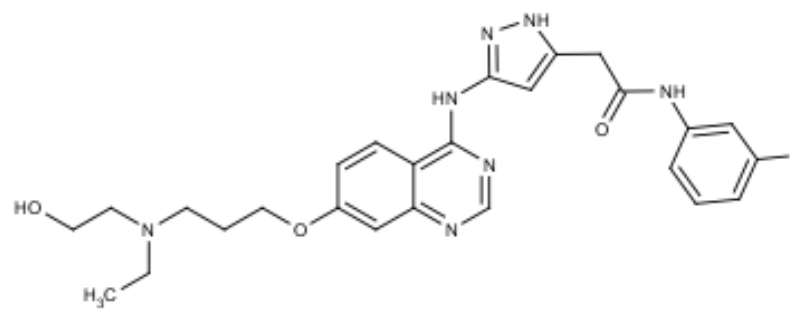
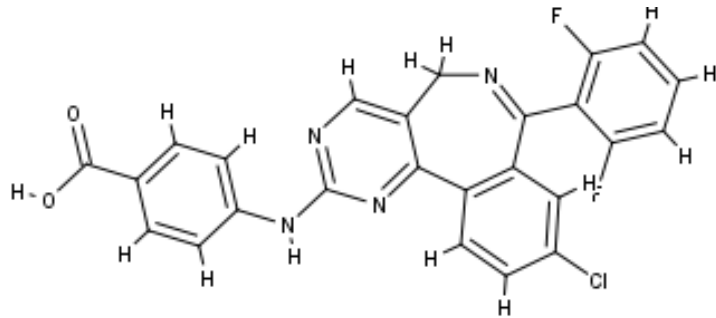
(1). Receptor/Target-Based

(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

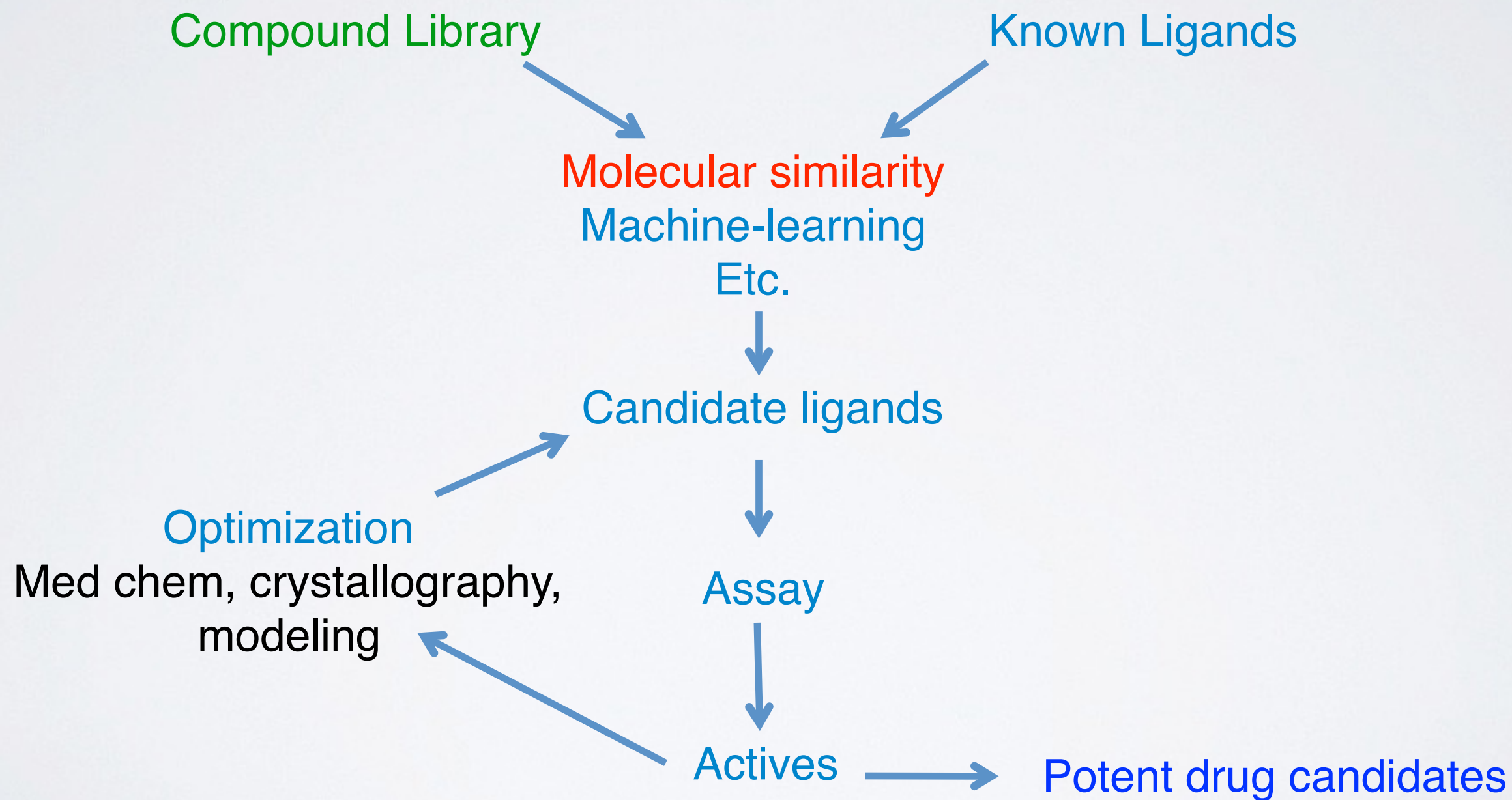
Why Look for Another Ligand if You Already Have Some?

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.

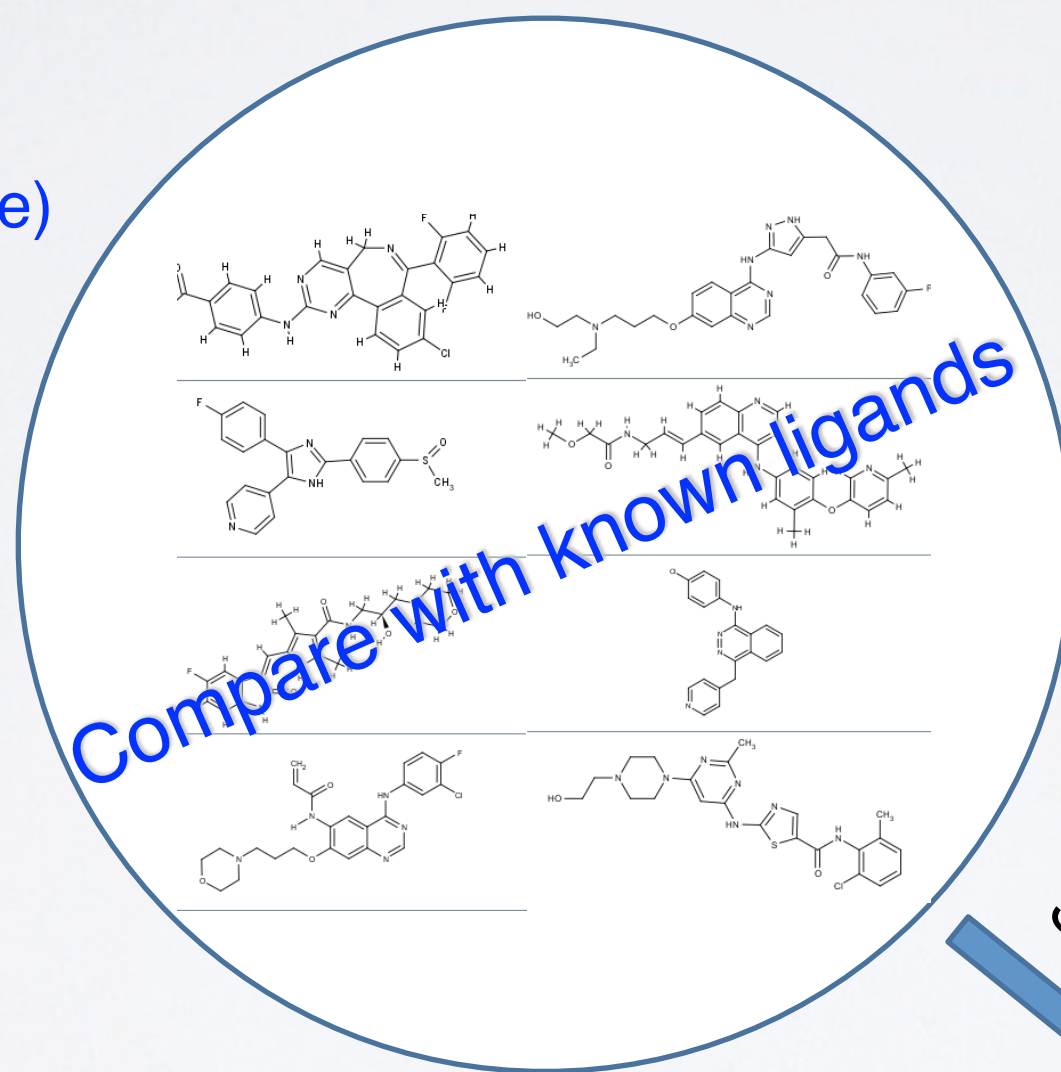
LIGAND-BASED VIRTUAL SCREENING



CHEMICAL SIMILARITY

LIGAND-BASED DRUG-DISCOVERY

Compounds
(available/synthesizable)



Different

Don't bother

Similar

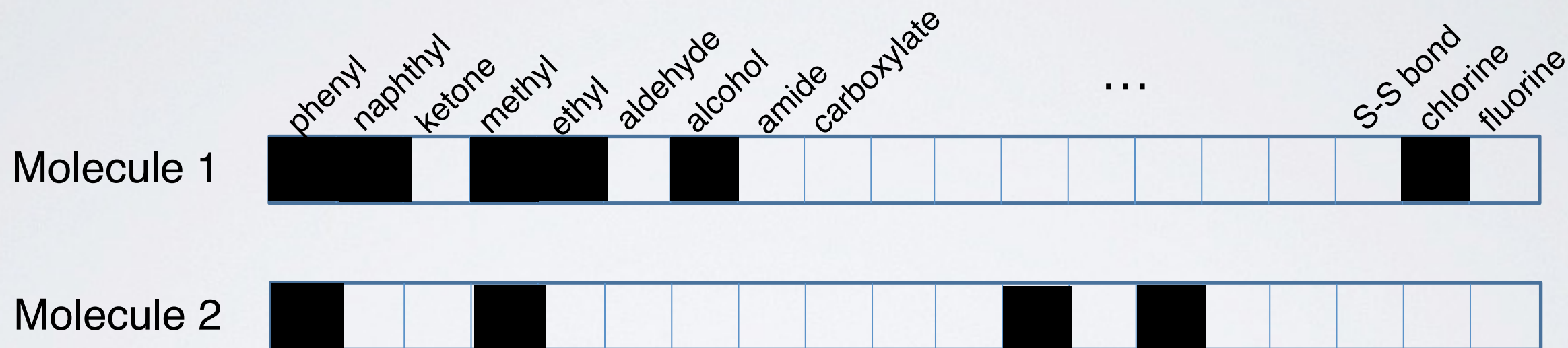
Test experimentally

CHEMICAL FINGERPRINTS

BINARY STRUCTURE KEYS



CHEMICAL SIMILARITY FROM FINGERPRINTS



Tanimoto Similarity
(or Jaccard Index), T

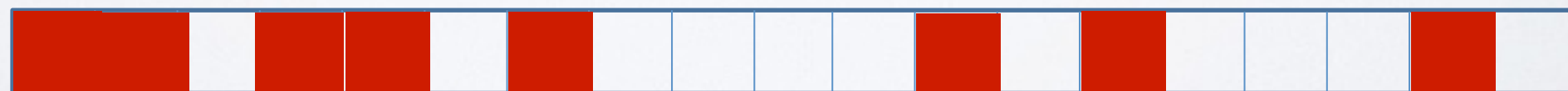
$$T \equiv \frac{N_I}{N_U} = 0.25$$

Intersection



$N_I=2$

Union

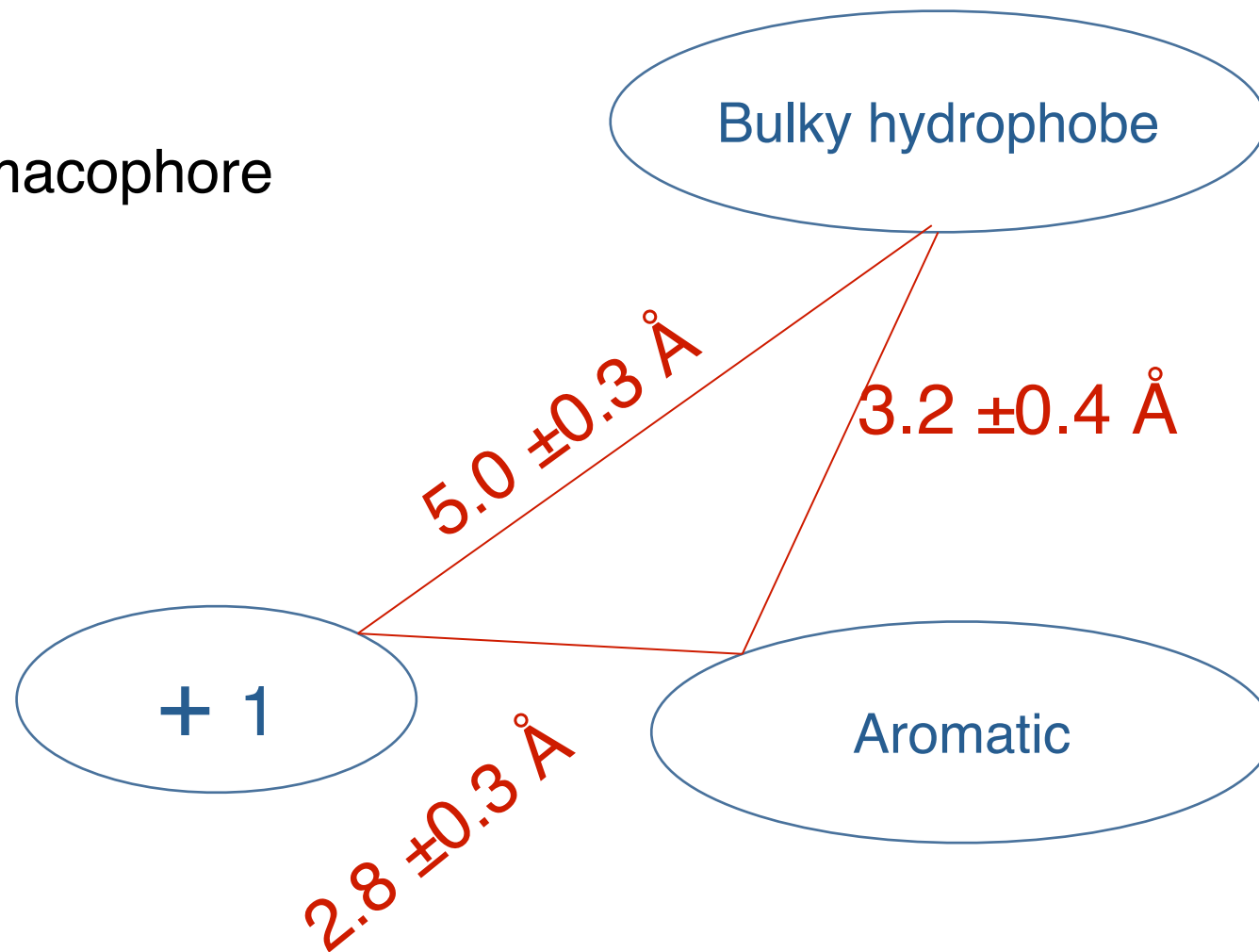


$N_U=8$

Pharmacophore Models

Φάρμακο (drug) + Φορά (carry)

A 3-point pharmacophore



Molecular Descriptors

More abstract than chemical fingerprints

Physical descriptors

molecular weight

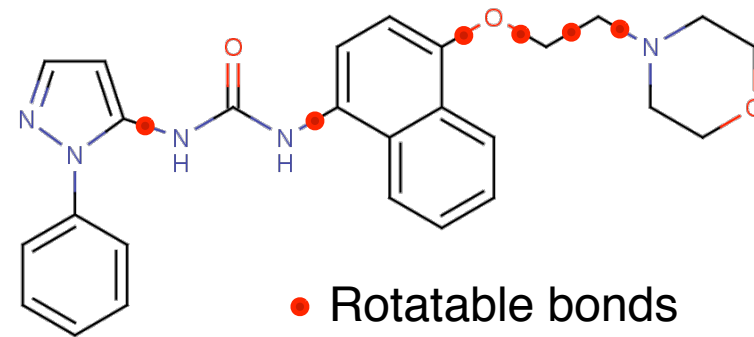
charge

dipole moment

number of H-bond donors/acceptors

number of rotatable bonds

hydrophobicity (log P and clogP)



Topological

branching index

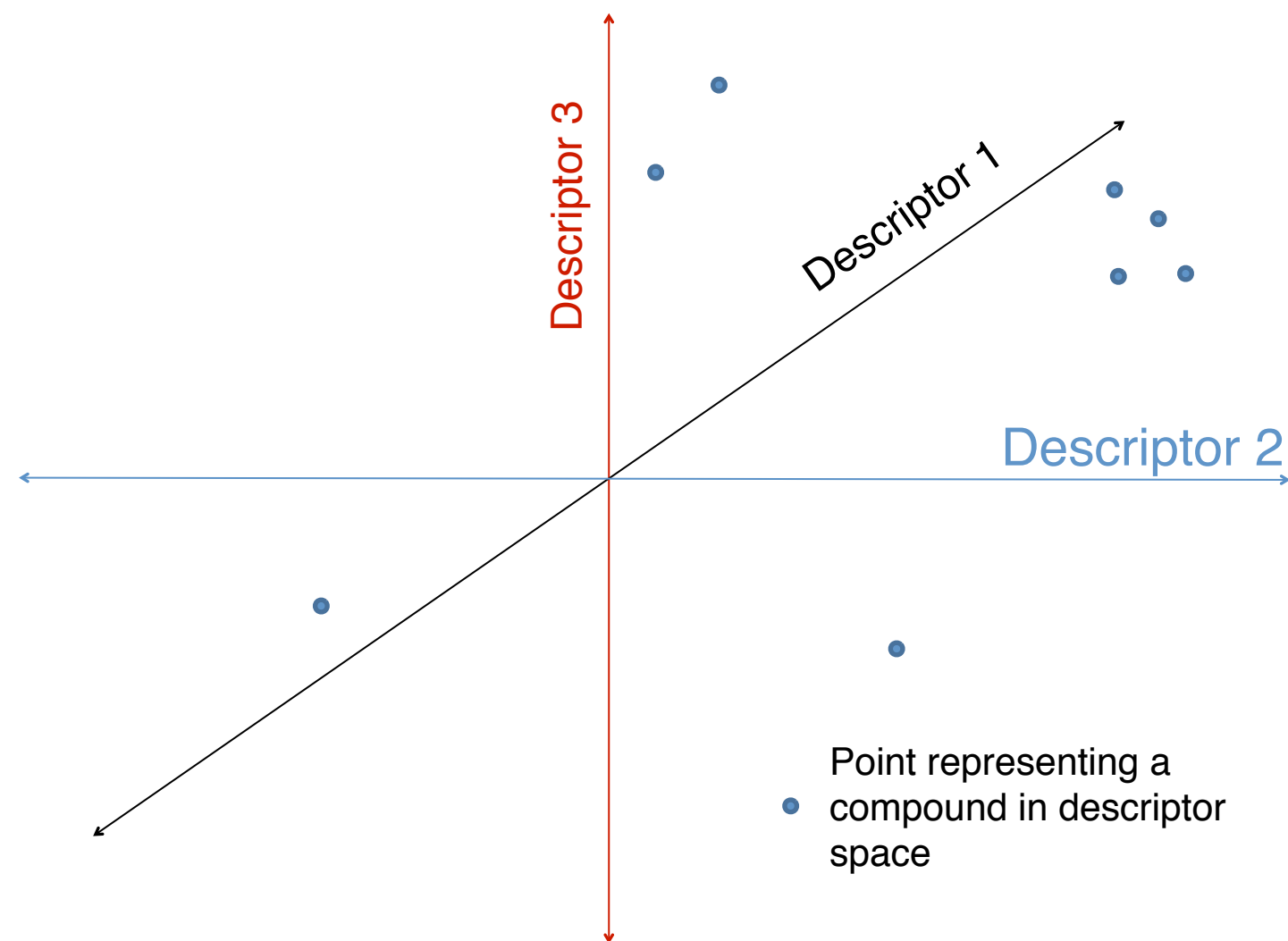
measures of linearity vs interconnectedness

Etc. etc.

A High-Dimensional “Chemical Space”

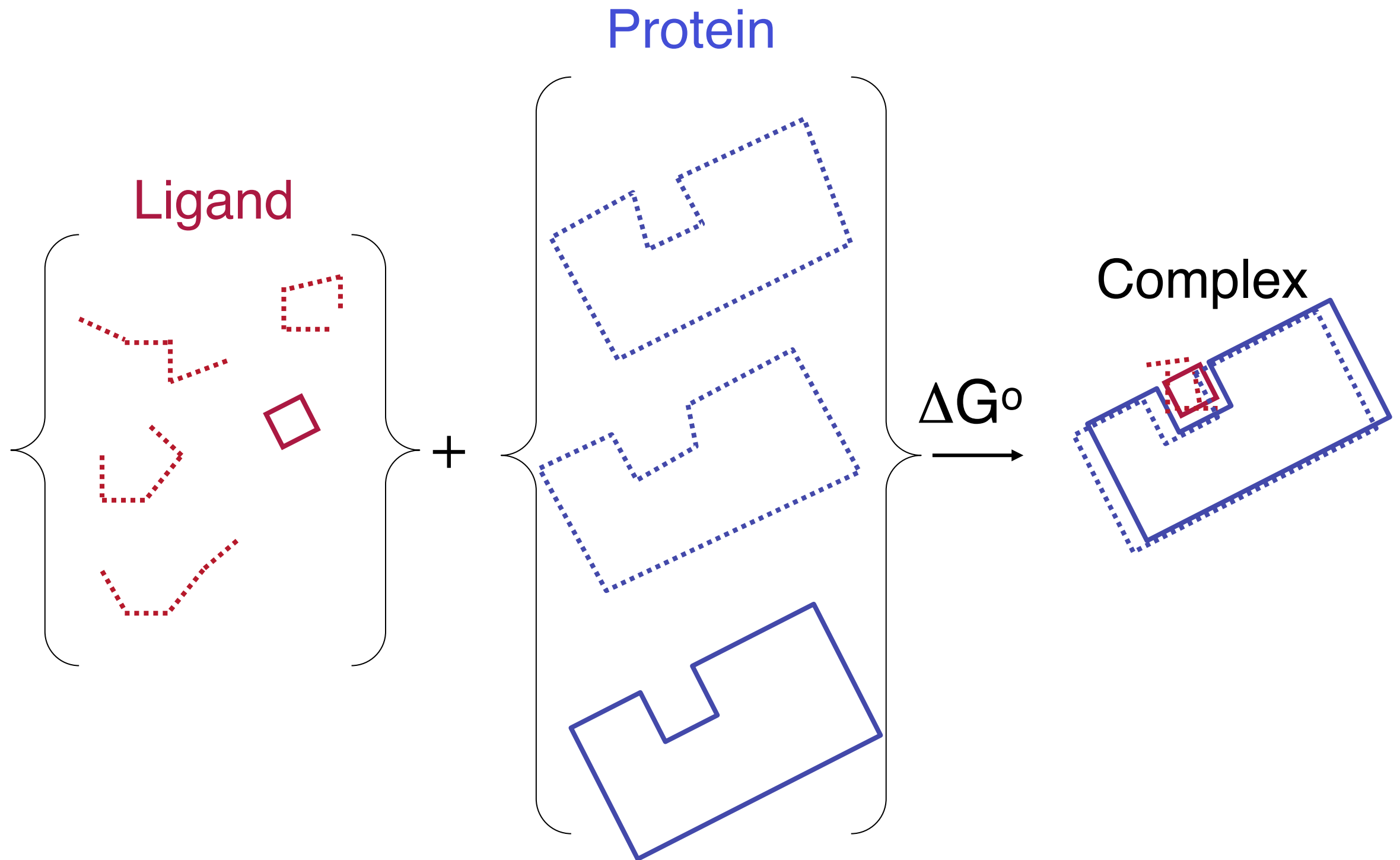
Each compound is a point in an n-dimensional space

Compounds with similar properties are near each other



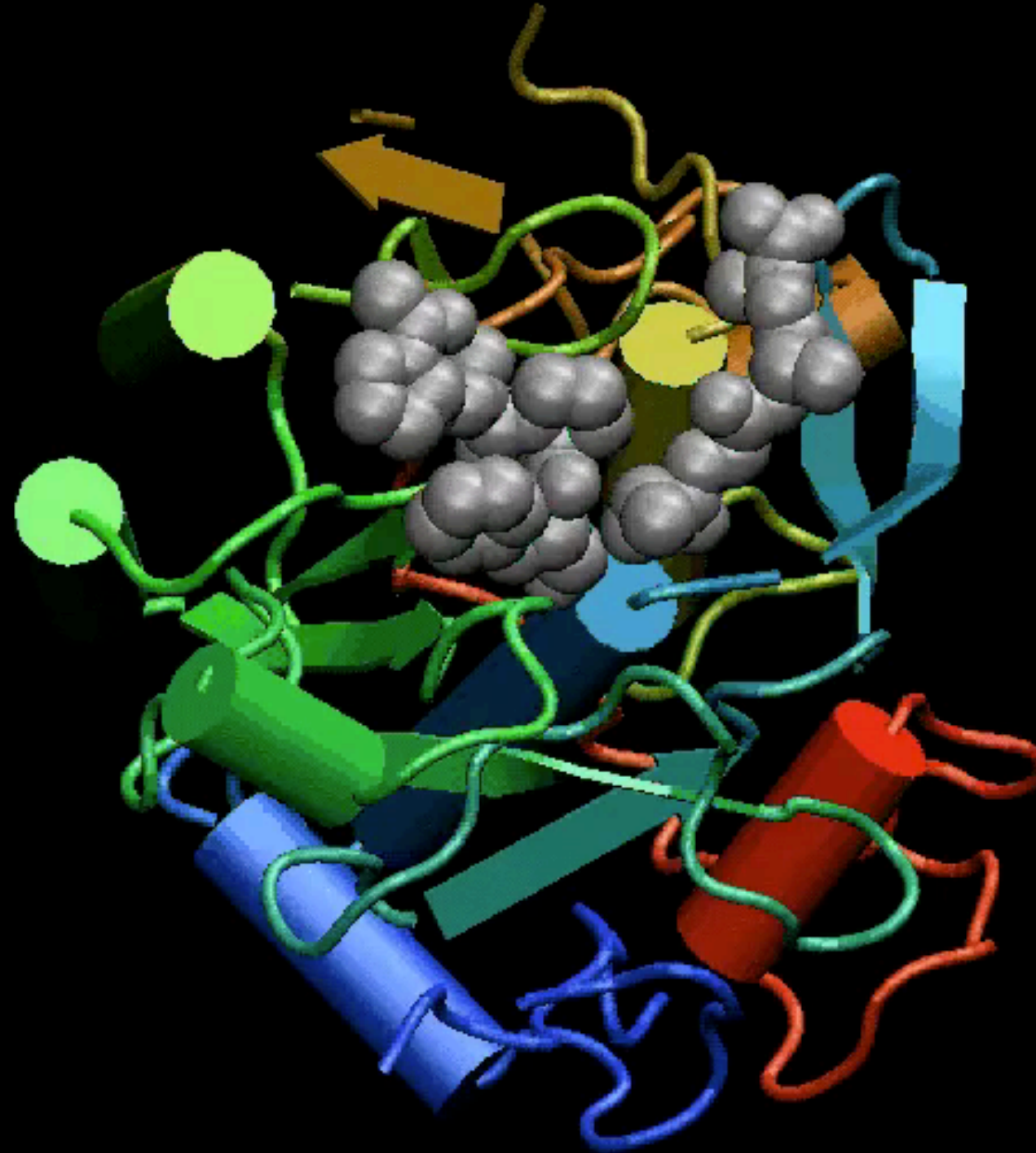
Apply multivariate statistics and machine learning for descriptor-selection. (e.g. partial least squares, PCA, support vector machines, random forest, deep learning etc.)

Proteins and Ligand are Flexible



NMA (Normal Mode Analysis) is a bioinformatics method to predict the intrinsic dynamics of biomolecules

Do it Yourself!



https://bioboot.github.io/bimm143_S19/lectures/#12

NMA in Bio3D

- Normal Mode Analysis (NMA) is a bioinformatics method that can predict the major motions of biomolecules.

```
pdb <- read.pdb("1hel")  
modes <- nma( pdb )  
m7 <- mktrj(modes, mode=7, file="mode_7.pdb")
```

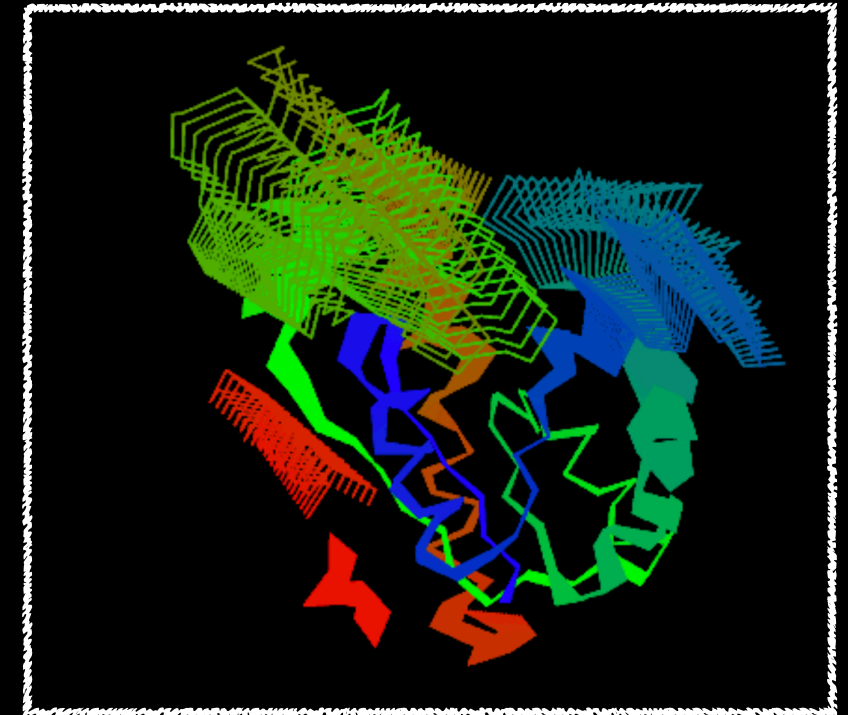
Then you can open the resulting `mode_7.pdb` file in VMD
> Use "TUBE" representation and hit the play button...

```
view(m7, col=vec2color(rmsf(m7)))
```

```
...
```

Bio3D view()

- If you want the 3D viewer in your R markdown you can install the development version of **bio3d.view**
- In your R console:



```
> install.packages("devtools")  
> devtools::install_bitbucket("Grantlab/bio3d-view")  
> install.packages("rgl")
```

- To use in your R session:

```
> library("bio3d.view")  
> pdb <- read.pdb("5p21")  
> view(pdb)  
> view(pdb, "overview", col="sse")  
> view(m7)
```

SideNote: **view()**

- If you want the interactive 3D viewer in Rmd rendered to **output: html_output** document:

```
```{r}
library(bio3d.view)
library(rgl)
```
```

```
```{r}
modes <- nma(read.pdb("1hel"))
m7 <- mktrj(modes, mode=7, file="mode_7.pdb")

view(m7, col=vec2color(rmsf(m7)))
rglwidget(width=500, height=500)
```
```

Do it Yourself!

Hand-on time!

https://bioboot.github.io/bimm143_S19/lectures/#13

Focus on **section 3** & **4** exploring **NMA** and **PCA apps**

Reference Slides

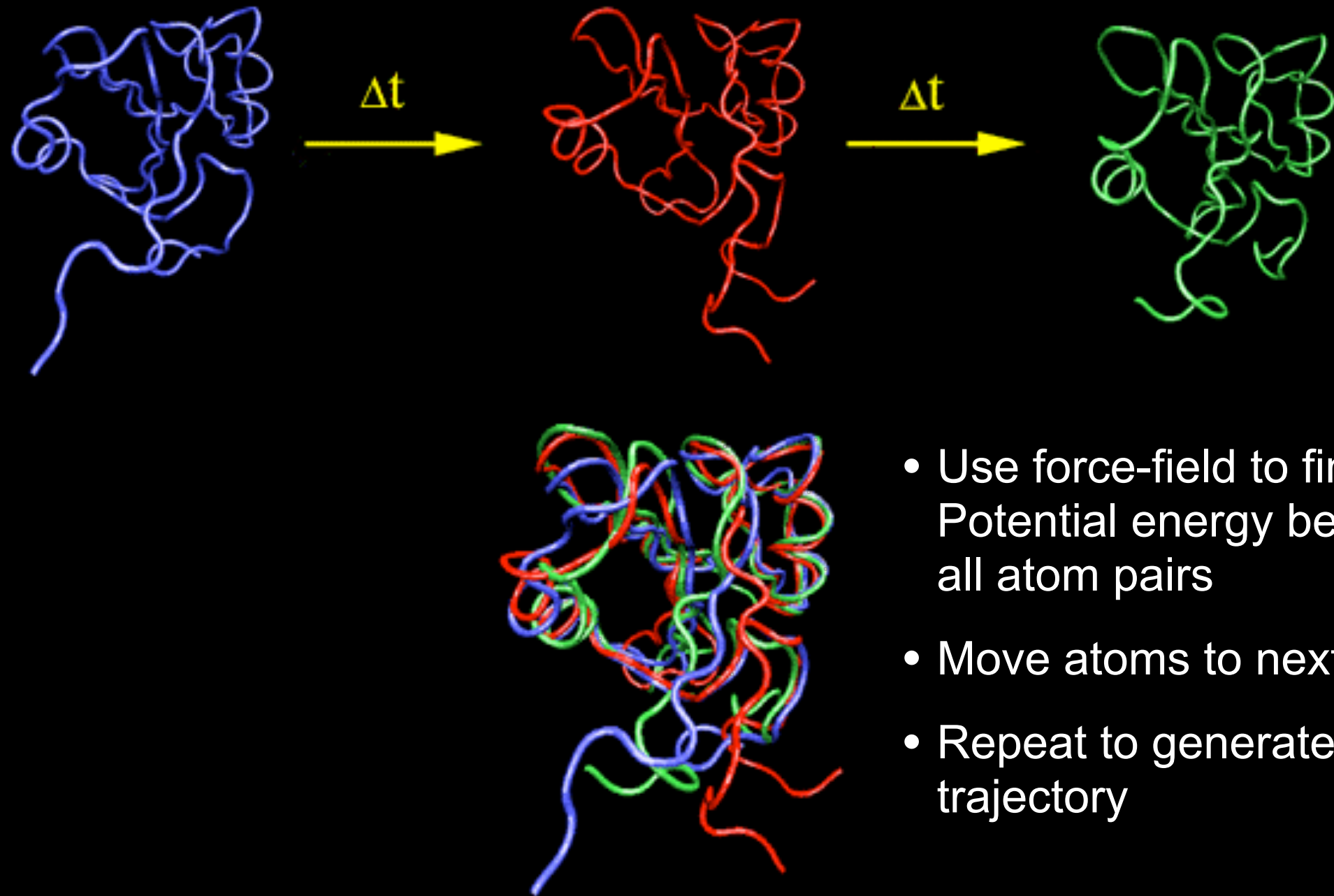
Molecular Dynamics (MD) and Normal Mode Analysis
(NMA) Background and Cautionary Notes

[[Muddy Point Assessment](#)]

PREDICTING FUNCTIONAL DYNAMICS

- Proteins are intrinsically flexible 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 mapping of structure to function
 - Molecular dynamics (MD) and normal mode analysis (NMA) are two major methods for predicting and characterizing molecular motions and their properties

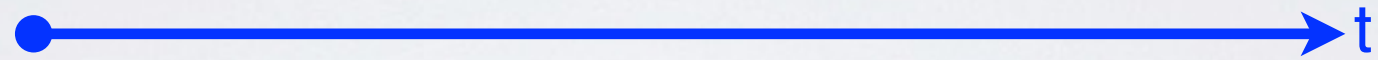
MOLECULAR DYNAMICS SIMULATION



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

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

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



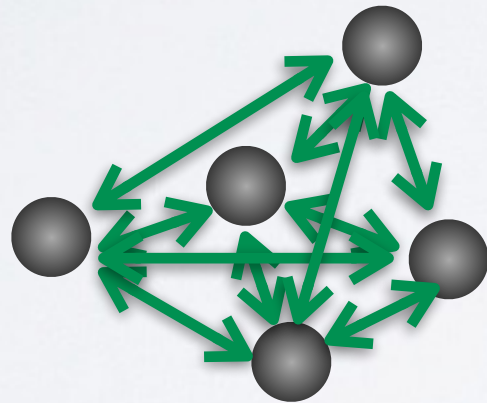
- ▶ Divide **time** into discrete (~ 1 fs) **time steps** (Δt)
(for integrating equations of motion, see below)



- ▶ Divide **time** into discrete (~ 1 fs) **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

$$m_i \frac{d^2 \vec{R}_i}{dt^2} = -\vec{\nabla}_i E(\vec{R})$$

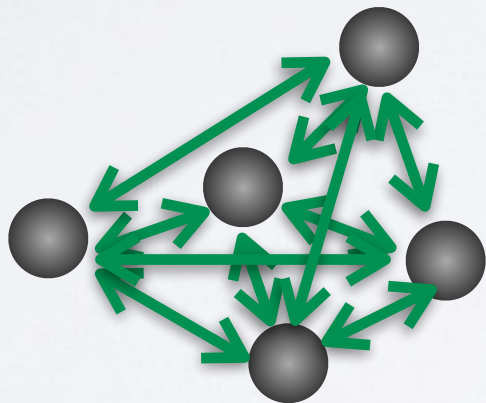
Empirical force field

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

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



- ▶ At each time step calculate pair-wise atomic **forces** ($\mathbf{F}(t)$)
(by evaluating force-field gradient)



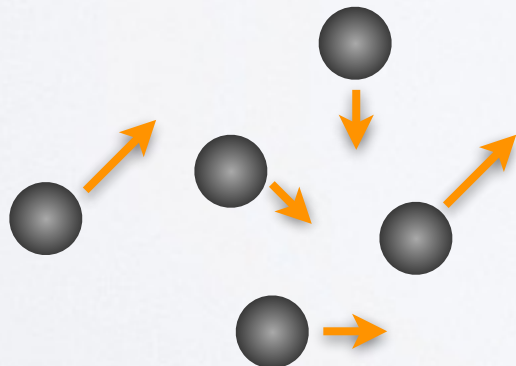
Nucleic motion described classically

$$m_i \frac{d^2 \vec{R}_i}{dt^2} = -\vec{\nabla}_i E(\vec{R})$$

Empirical force field

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

- ▶ Use the forces to calculate velocities and move atoms to new positions
(by integrating numerically via the “leapfrog” scheme)



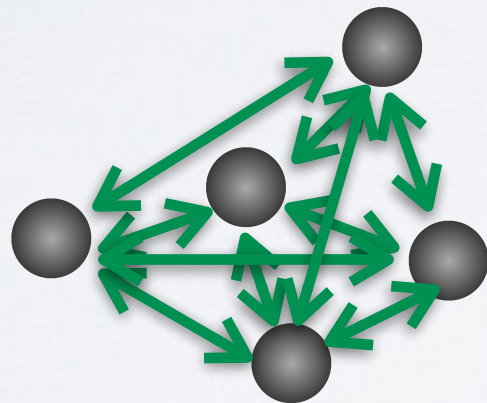
$$\begin{aligned} \mathbf{v}\left(t + \frac{\Delta t}{2}\right) &= \mathbf{v}\left(t - \frac{\Delta t}{2}\right) + \frac{\mathbf{F}(t)}{m} \Delta t \\ \mathbf{r}(t + \Delta t) &= \mathbf{r}(t) + \mathbf{v}\left(t + \frac{\Delta t}{2}\right) \Delta t \end{aligned}$$

BASIC ANATOMY OF A MD SIMULATION

- ▶ Divide **time** into discrete (~ 1 fs) **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

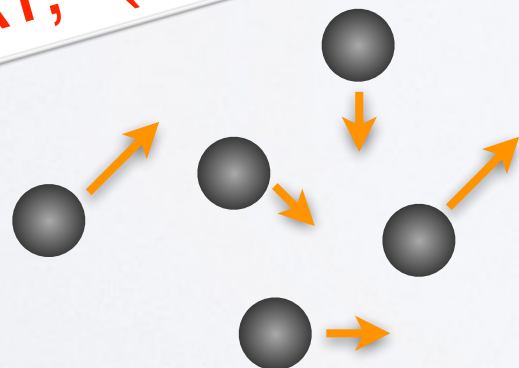
$$m_i \frac{d^2 \vec{R}_i}{dt^2} = -\vec{\nabla}_i E(\vec{R})$$

Empirical force field

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

- ▶ Use the forces to calculate velocities and move atoms to new positions
(the integration is done numerically via the “leapfrog” scheme)

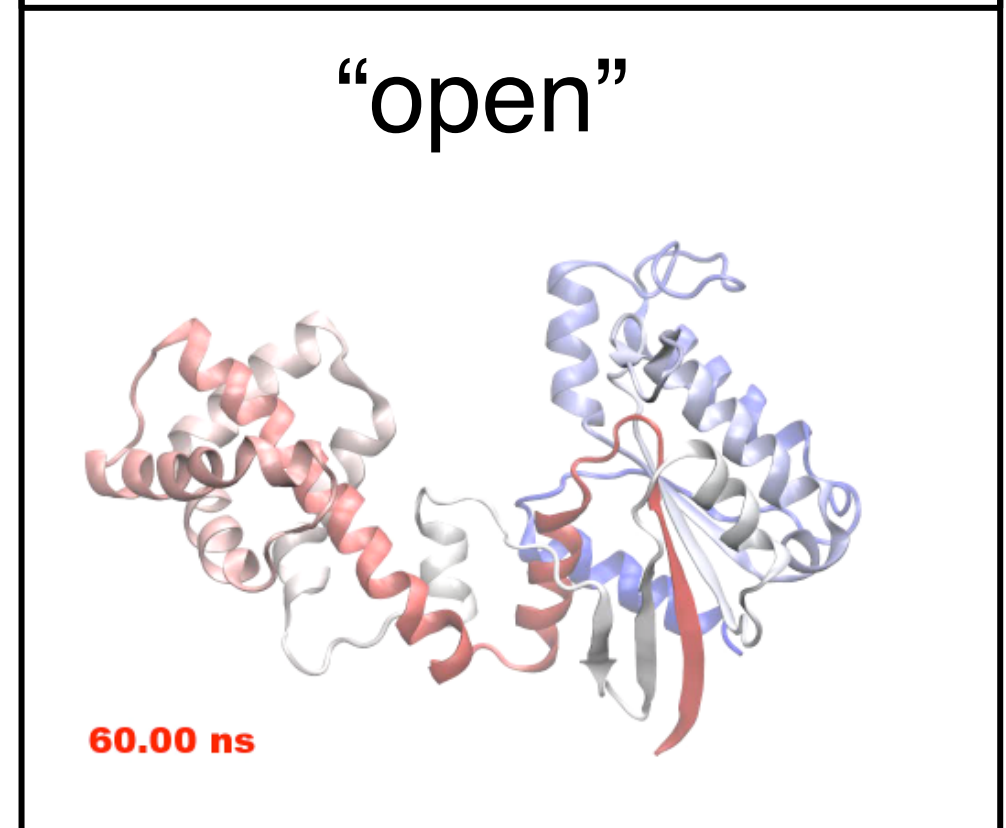
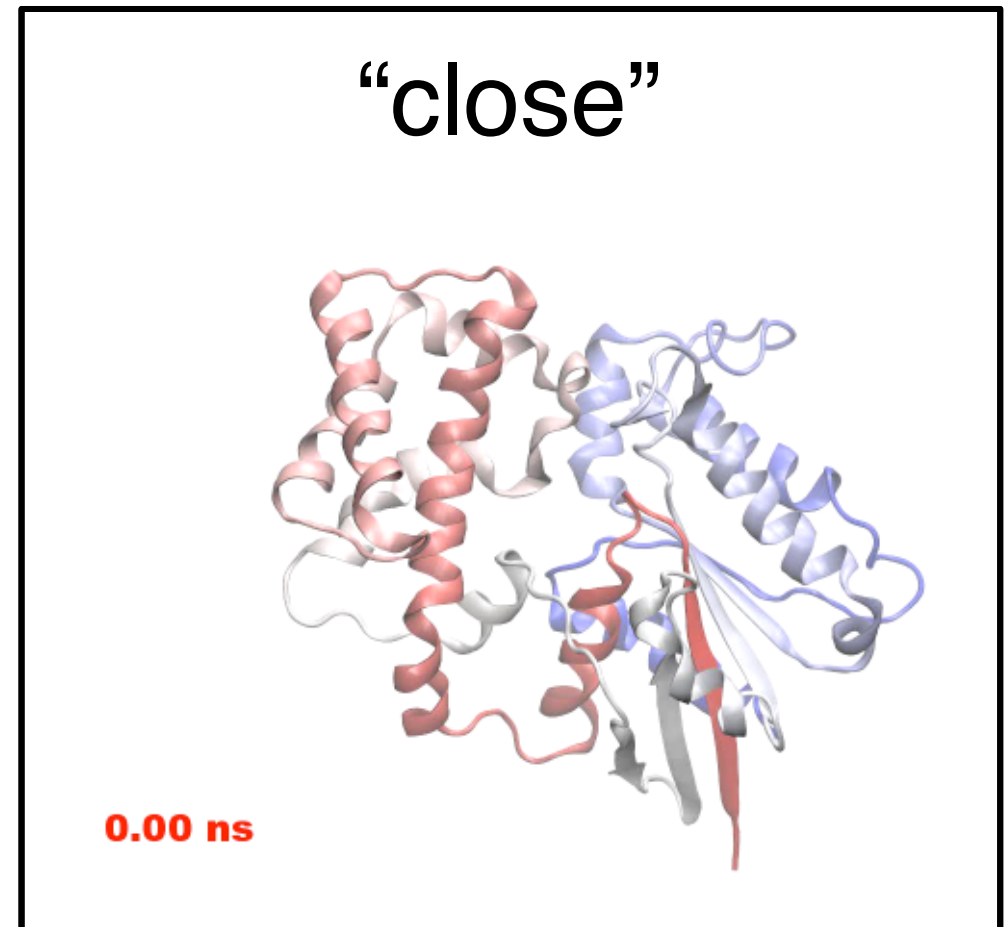
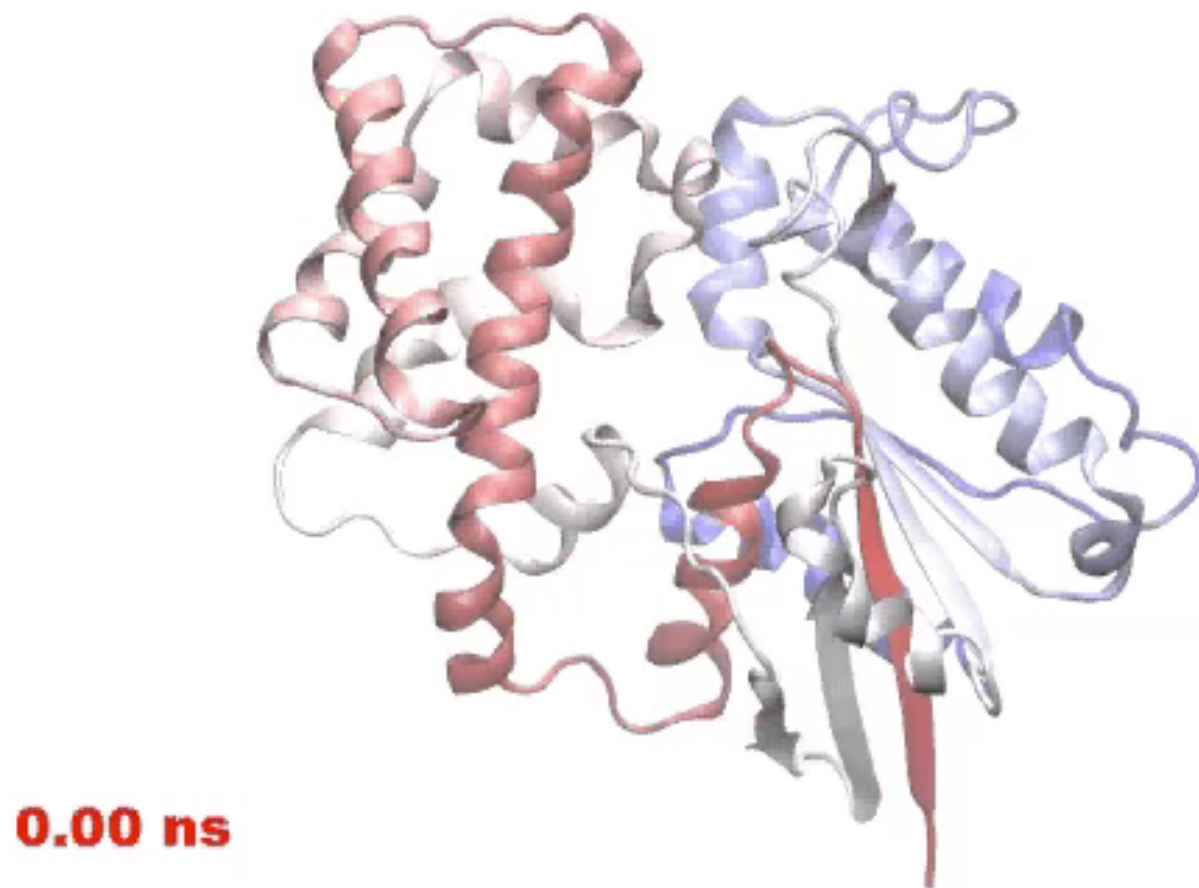
REPEAT, (iterate many, many times... 1ms = 10¹² time steps)



$$\begin{aligned} \mathbf{v}\left(t + \frac{\Delta t}{2}\right) &= \mathbf{v}\left(t - \frac{\Delta t}{2}\right) + \frac{\mathbf{F}(t)}{m} \Delta t \\ \mathbf{r}(t + \Delta t) &= \mathbf{r}(t) + \mathbf{v}\left(t + \frac{\Delta t}{2}\right) \Delta t \end{aligned}$$

MD Prediction of Functional Motions

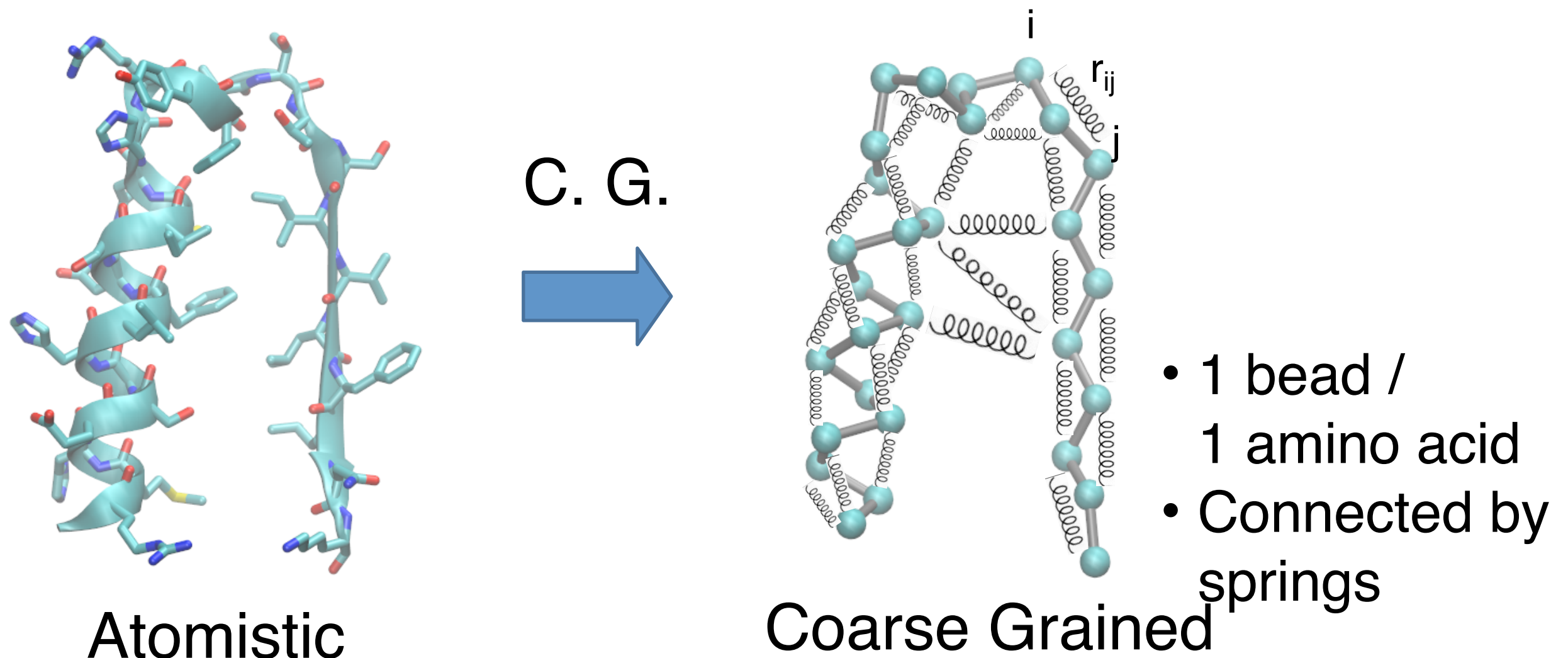
Accelerated MD simulation of
nucleotide-free transducin alpha subunit



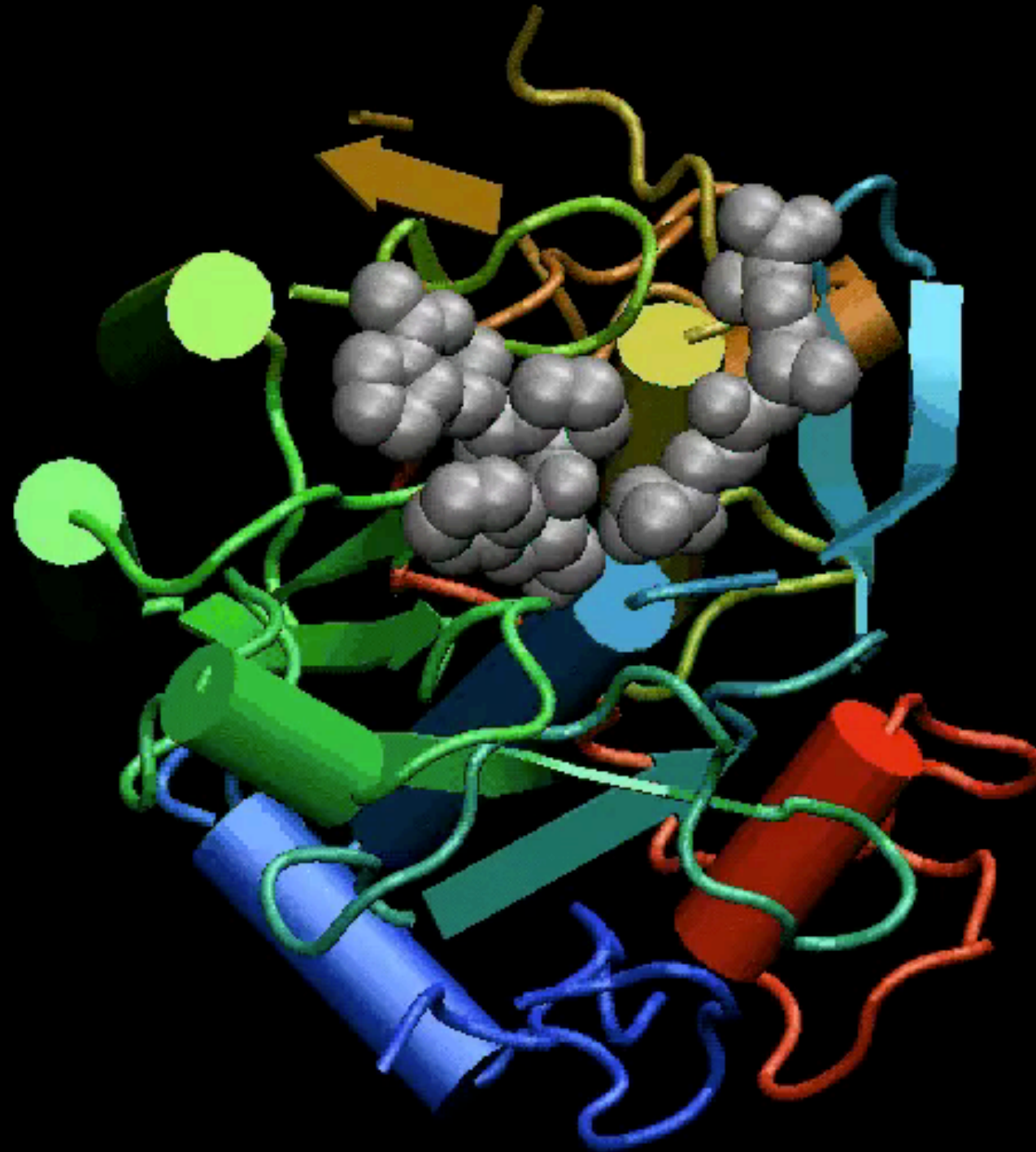
Yao and Grant, Biophys J. (2013)

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 it Yourself!

Hand-on time!

https://bioboot.github.io/bimm143_S19/lectures/#13

Focus on **section 3** & **4** exploring **NMA** and **PCA apps**

ACHIEVEMENTS

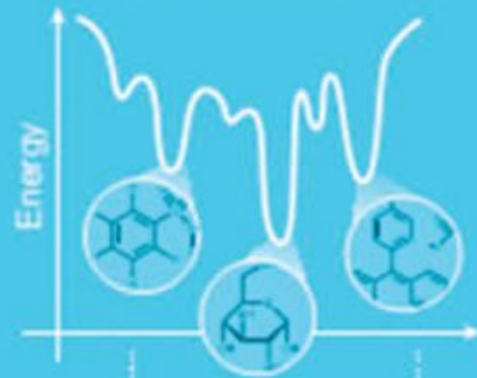
Computational power



Data coverage and community resources



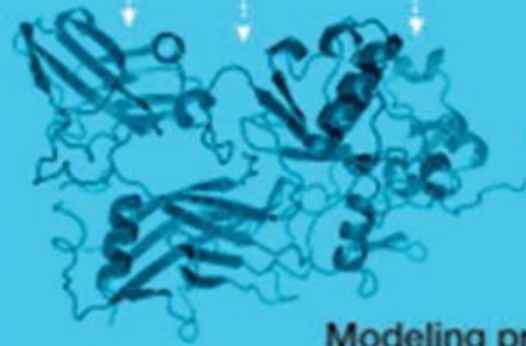
Chemical systems biology and small-molecule docking simulations



Objective method assessment



Correlated mutations



Modeling protein structure

CHALLENGES

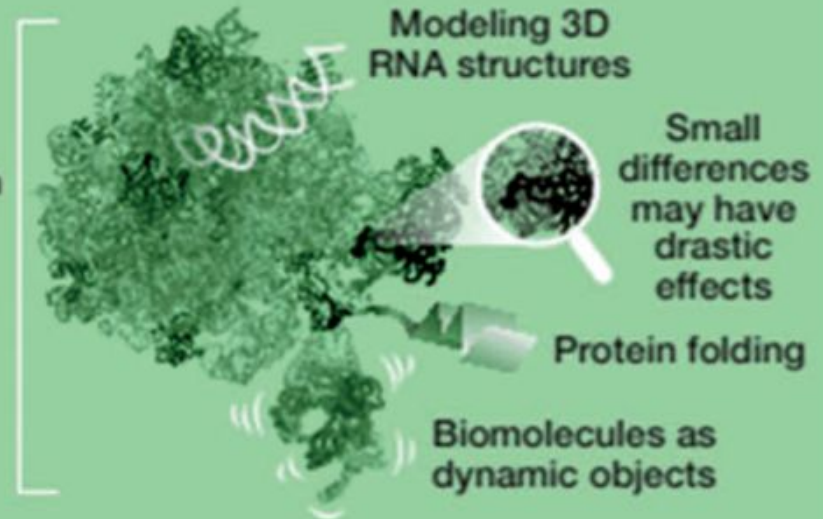
Accessibility and integration of data and methods



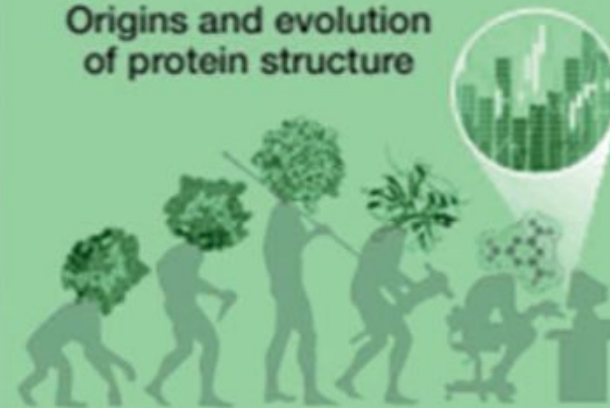
Protein engineering and synthetic biology



Modeling multi-domain proteins and large assemblies



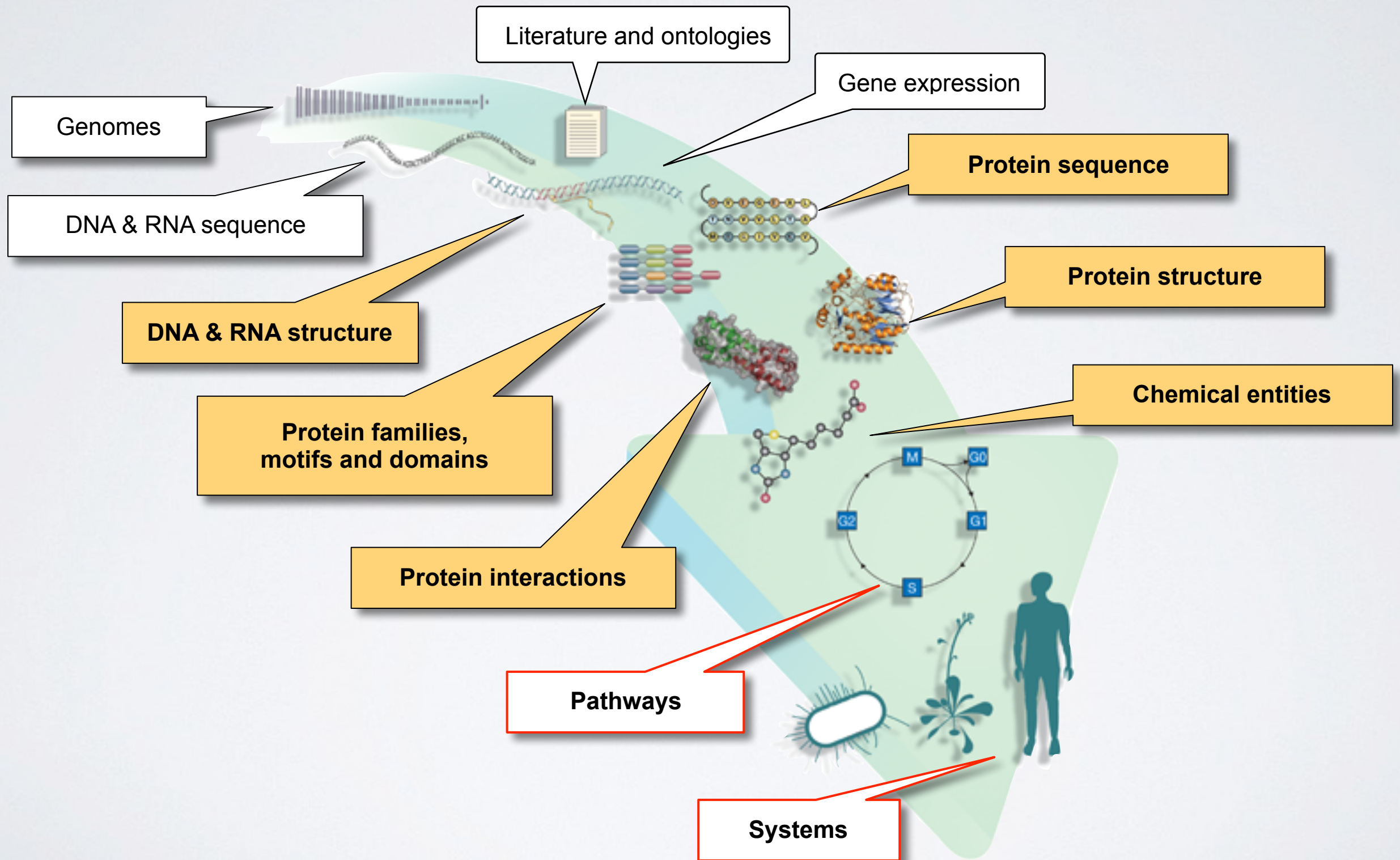
Origins and evolution of protein structure



Integration with systems biology



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
- Explored how to visualize protein structure with VMD and use R to perform more advanced structural bioinformatics analysis!
- 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

[[Muddy Point Assessment](#)]

CAUTIONARY NOTES

- A model is never perfect

A model that is not quantitatively accurate in every respect does not preclude one from establishing results relevant to our understanding of biomolecules as long as the biophysics of the model are properly understood and explored.

- Calibration of parameters is an ongoing imperfect process

Questions and hypotheses should always be designed such that they do not depend crucially on the precise numbers used for the various parameters.

- A computational model is rarely universally right or wrong

A model may be accurate in some regards, inaccurate in others. These subtleties can only be uncovered by comparing to all available experimental data.