BIMM 143 Structural Bioinformatics

### Lecture 11

Barry Grant UC San Diego

http://thegrantlab.org/bimm143

http://www.ks.uiuc.edu/Development/Download/download.cgi

"Bioinformatics is the application of <u>computers</u> to the collection, archiving, organization, and analysis of <u>biological data</u>."

... A hybrid of biology and computer science

"Bioinformatics is the application of <u>computers</u> to the collection, archiving, organization, and analysis of <u>biological data</u>."

**Bioinformatics is computer aided biology!** 

"Bioinformatics is the application of <u>computers</u> to the collection, archiving, organization, and analysis of <u>biological data</u>."

Bioinformatics is computer aided biology! Goal: Data to Knowledge

### So what is structural bioinformatics?

# So what is structural bioinformatics?

... computer aided structural biology!

Aims to characterize and interpret biomolecules and their assembles at the molecular & atomic level

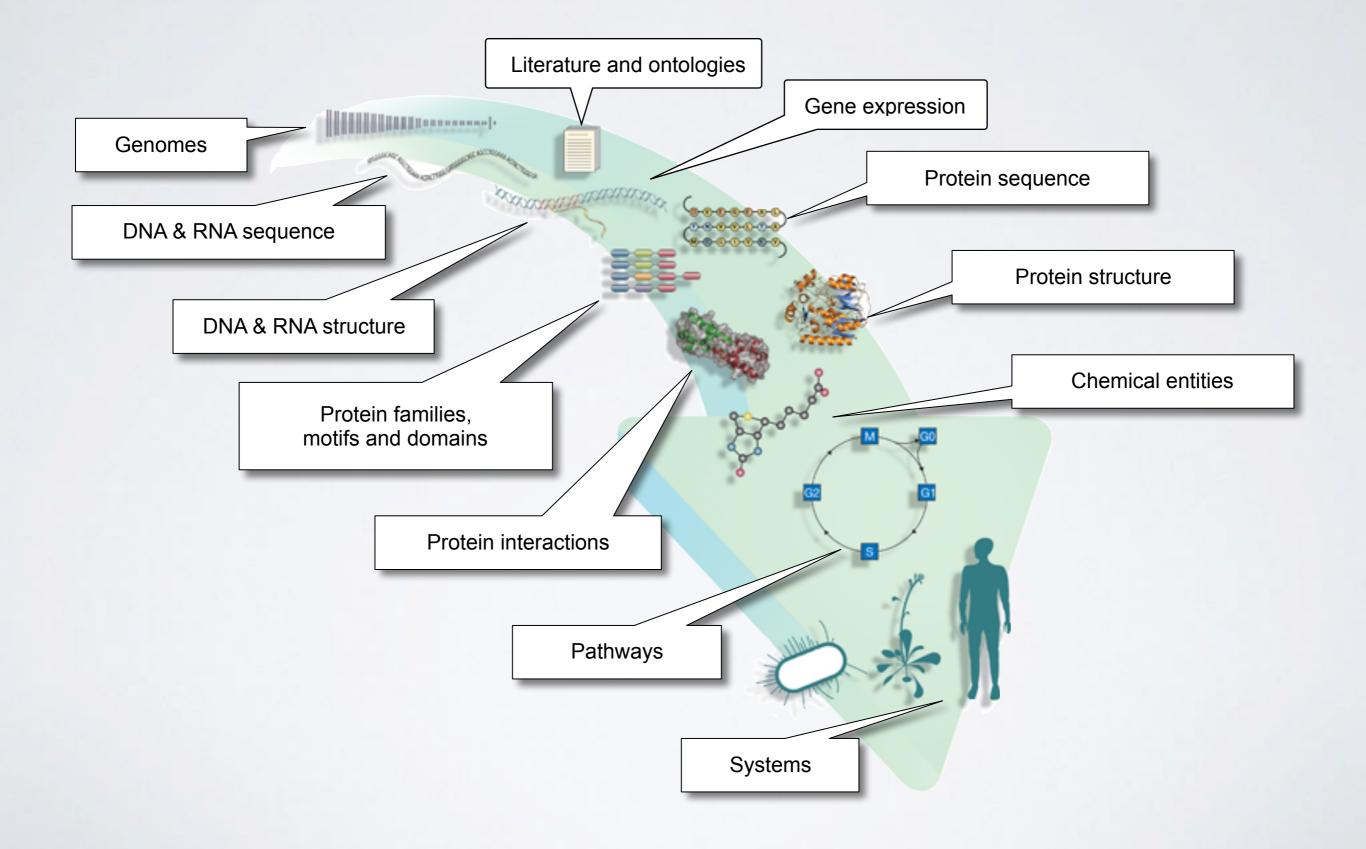
# Why should we care?

### Why should we care?

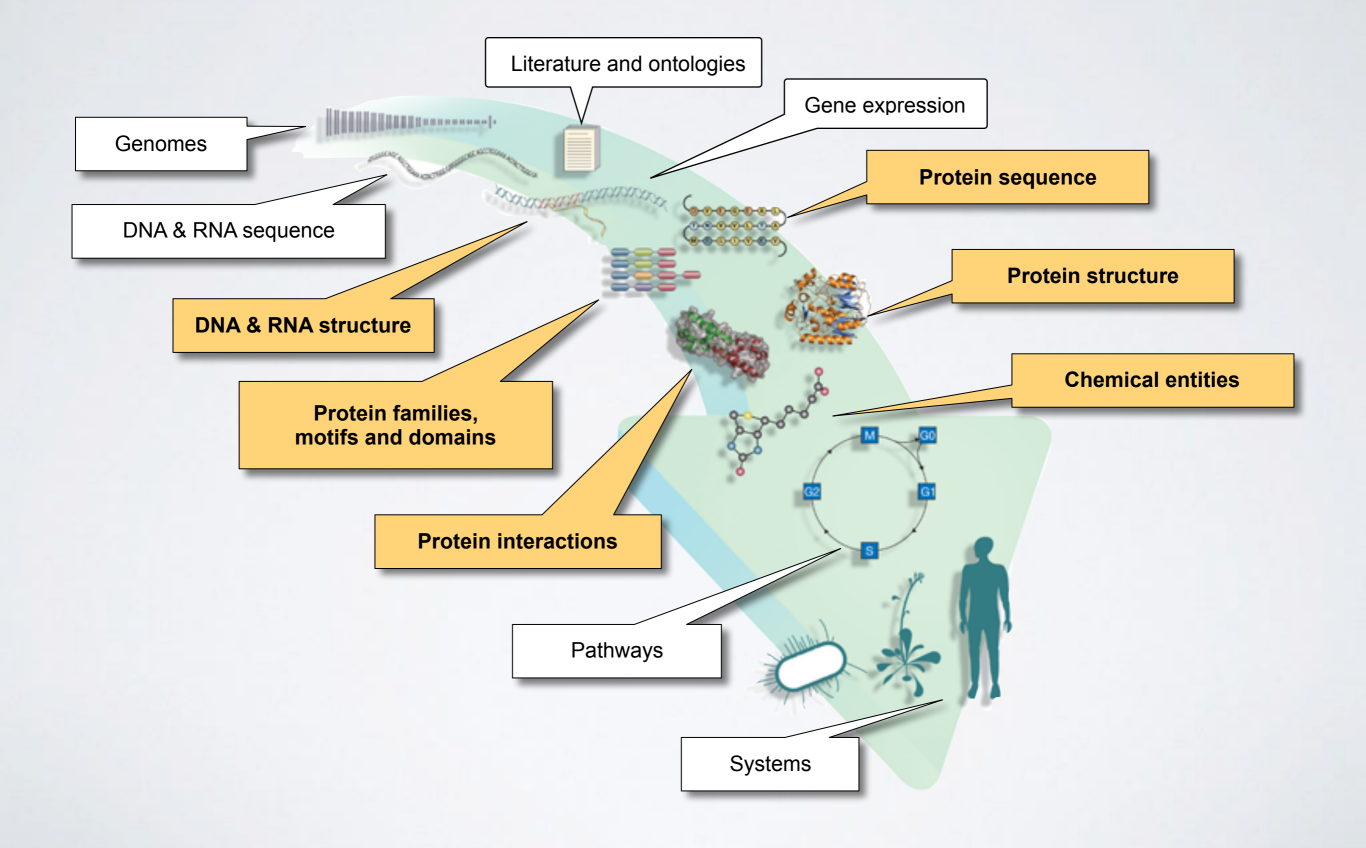
Because biomolecules are "nature's robots"

... and because it is only by coiling into specific 3D structures that they are able to perform their functions

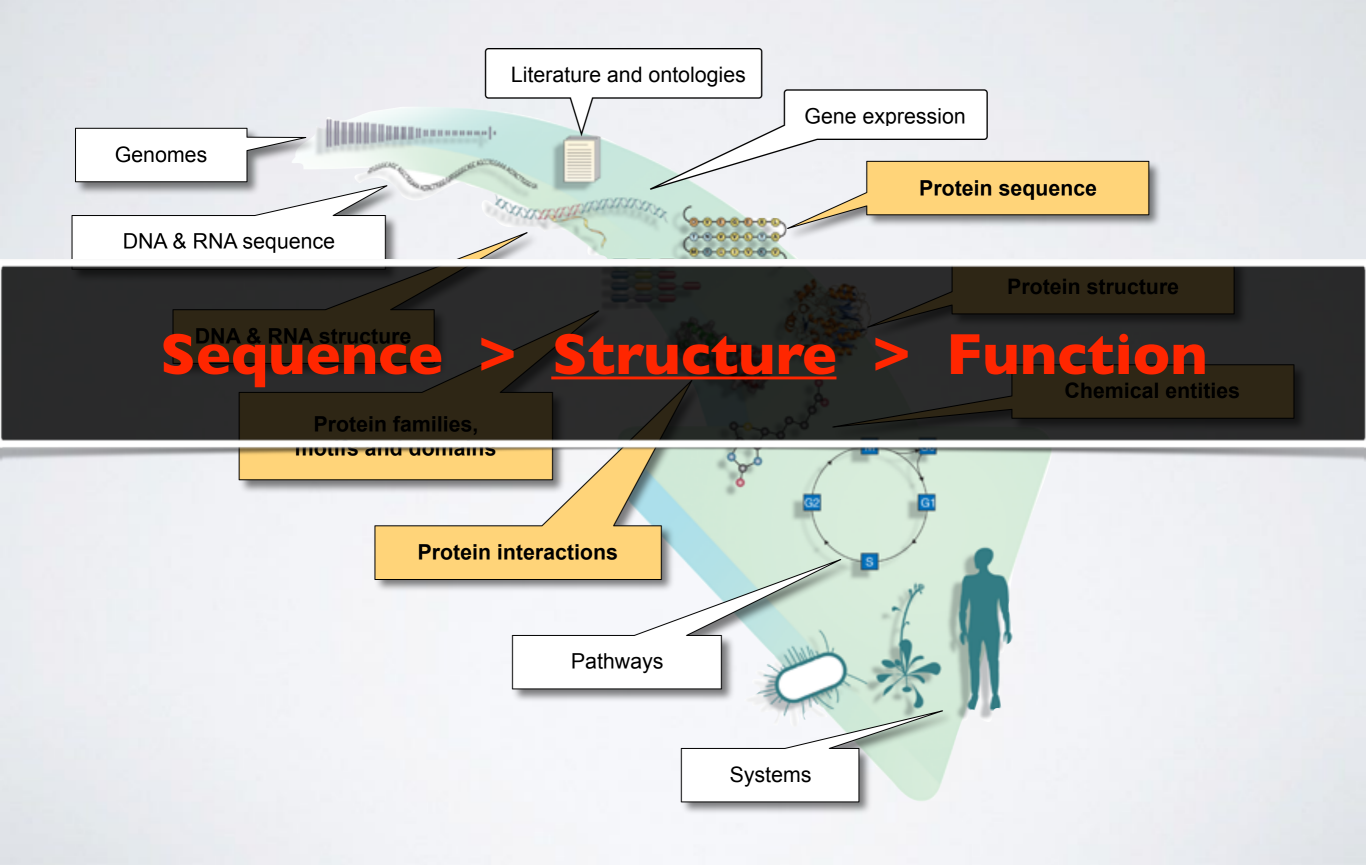
## BIOINFORMATICS DATA



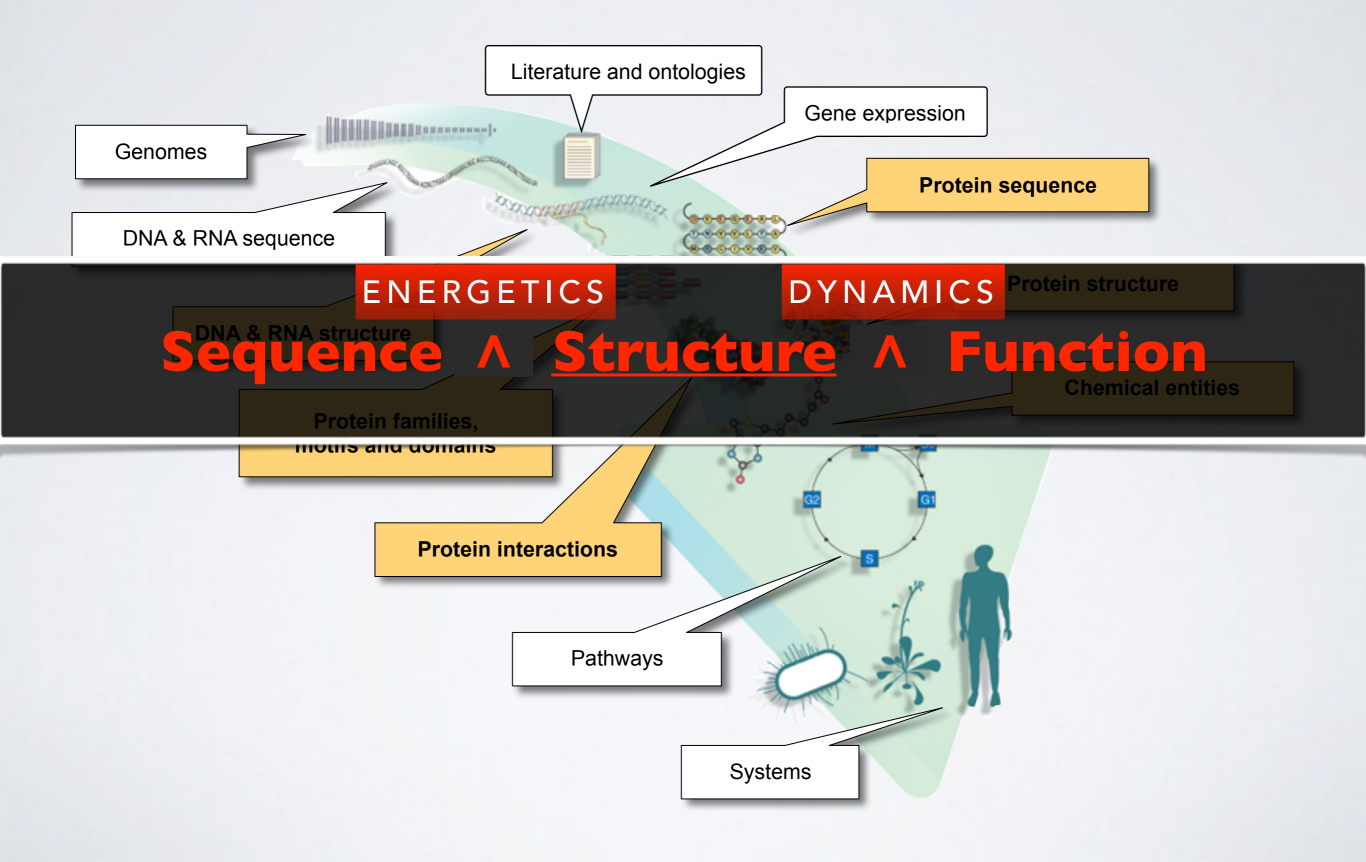
# STRUCTURAL DATA IS CENTRAL

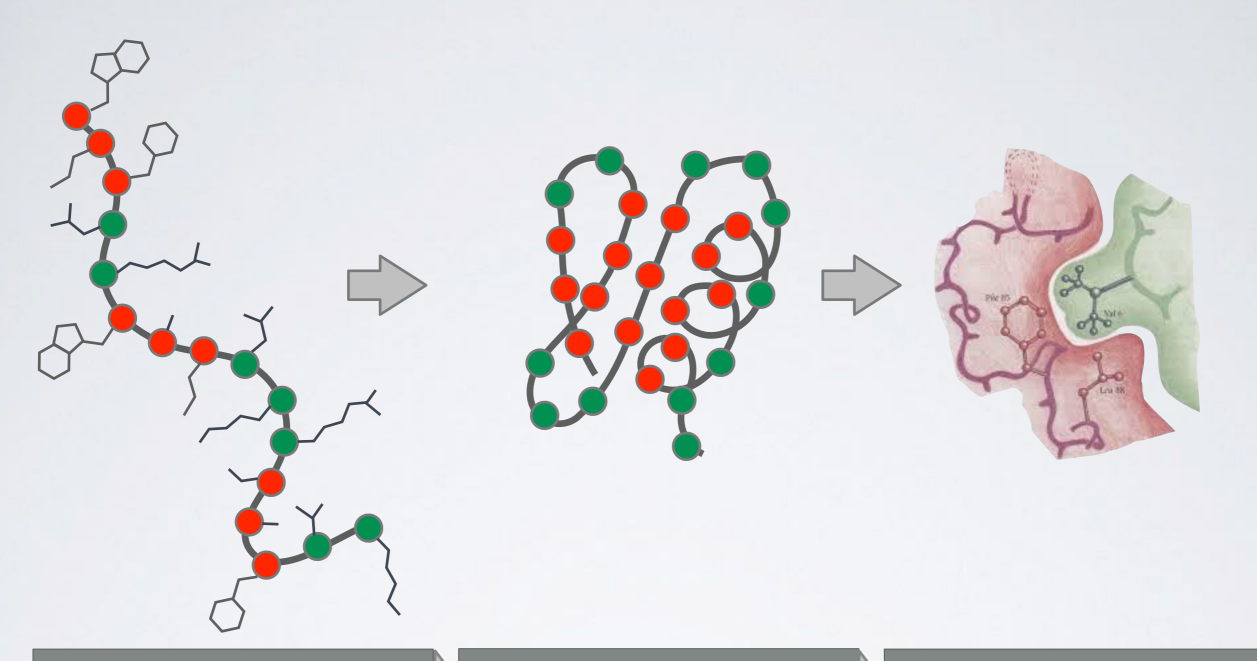


# STRUCTURAL DATA IS CENTRAL



# STRUCTURAL DATA IS CENTRAL





#### Sequence

- Unfolded chain of amino acid chain
- Highly mobile
- Inactive

### Structure

- Ordered in a precise 3D arrangment
- Stable but dynamic

### Function

- Active in specific ''conformations''
- Specific associations
   & precise reactions

# In daily life, we use machines with functional *structure* and *moving parts*





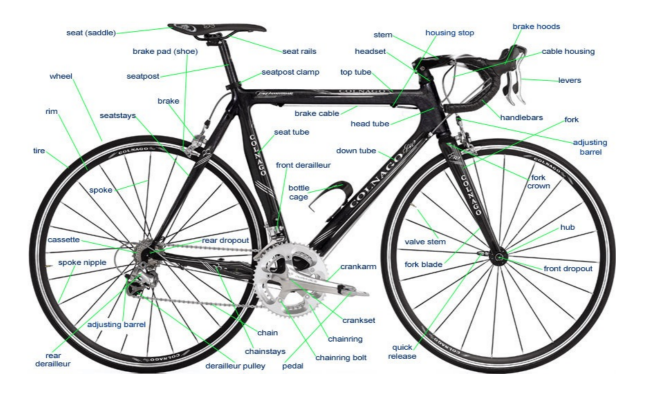
# Genomics is a great start ....

#### Track Bike - DL175

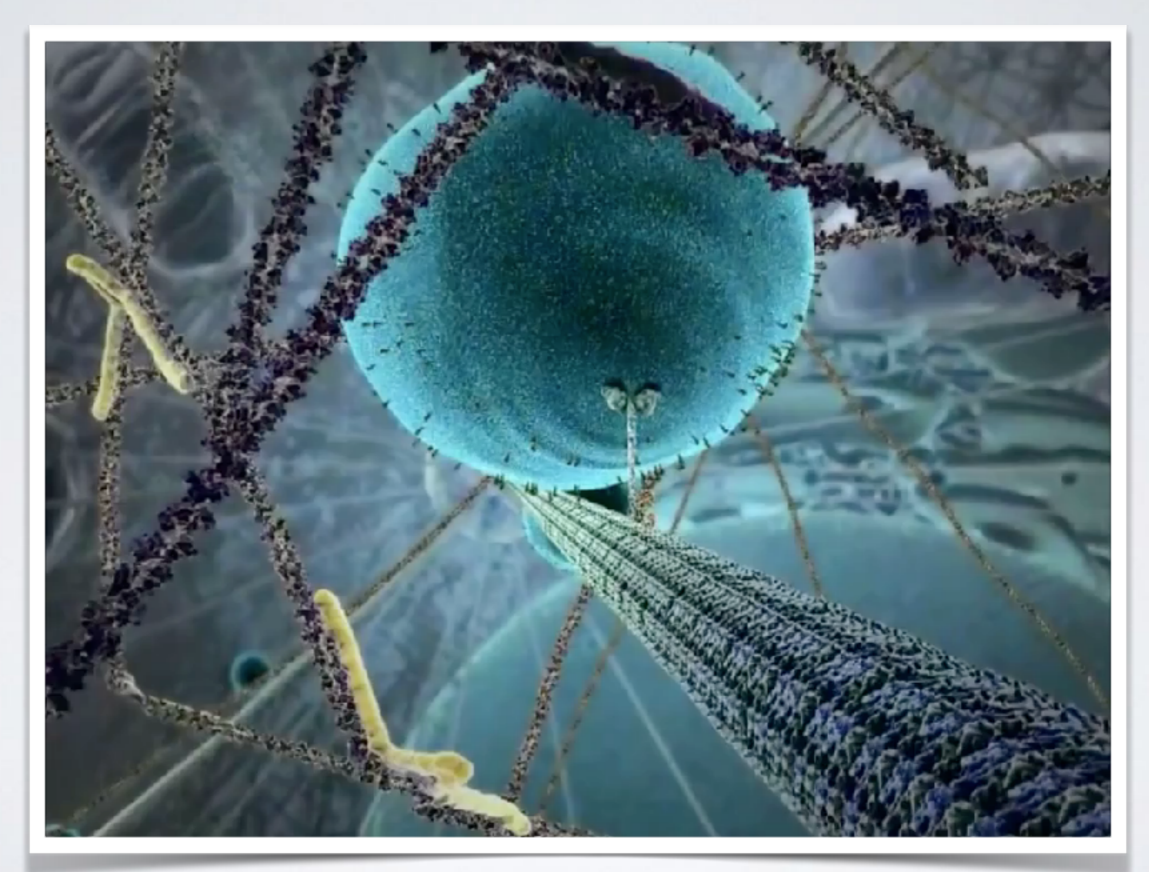
REF. NO.	IBM NO.	DESCRIPTION
1	156011	Track Frame 21", 22", 23", 24", Team Red
2	157040	Fork for 21" Frame
2	157039	Fork for 22" Frame
2 2 2 3 4	157038	Fork for 23" Frame
2	157037	Fork for 24" Frame
з	191202	Handlebar TTT Competition Track Alloy 15/16"
		Handlebar Stem, TTT, Specify extension
5	191278	Expander Bolt
5 6 7	191272	Clamp Bolt
	145841	Headset Complete 1 x 24 BSC
8	145842	Ball Bearings
9	190420	175 Raleigh Pistard Seta Tubular Prestavalve 27"
10	190233	Rim, 27" AVA Competition (36H) Alloy Prestavalve
11	145973	Hub, Large Flange Campagnolo Pista Track Alloy (pairs)
12	190014	Spokes, 11 5/8"
13	145837	Sleeve
14	145636	Ball Bearings
15	145170	Bottom Bracket Axle
16	145838	Cone for Sleeve
17	146473	L.H. Adjustable Cup
18	145833	Lockring
19	145239	Straps for Toe Clips
20	145834	Fixing Bolt
21	145835	Fixing Washer
22	145822	Dustcap
23	145823	R.H. and L.H. Crankset with Chainwheel
24	146472	Fixed Cup
25	145235	Toe Clips, Christophe, Chrome (Medium)
26	145684	Pedals, Extra Light, Pairs
27	123021	Chain
28	145980	Seat Post
29		Seat Post Bolt and Nut
30	167002	Saddle, Brooks
31	145933	Track Sprocket, Specify 12, 13, 14, 15, or 16 T.

 But a parts list is not enough to understand how a bicycle works

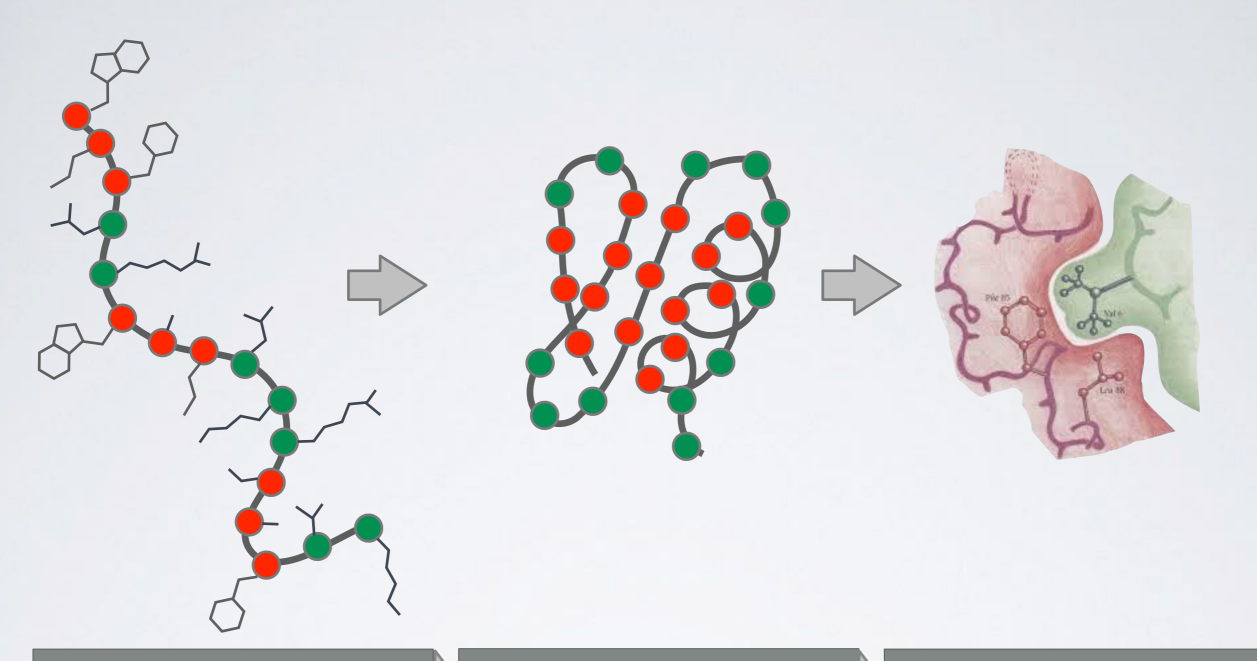
# ... but not the end



- We want the full spatiotemporal picture, and an ability to control it
- Broad applications, including drug design, medical diagnostics, chemical manufacturing, and energy



Extracted from The Inner Life of a Cell by Cellular Visions and Harvard [YouTube link: <u>https://www.youtube.com/watch?v=y-uuk4Pr2i8</u>]



#### Sequence

- Unfolded chain of amino acid chain
- Highly mobile
- Inactive

### Structure

- Ordered in a precise 3D arrangment
- Stable but dynamic

### Function

- Active in specific ''conformations''
- Specific associations
   & precise reactions

### **KEY CONCEPT: ENERGY LANDSCAPE**

5

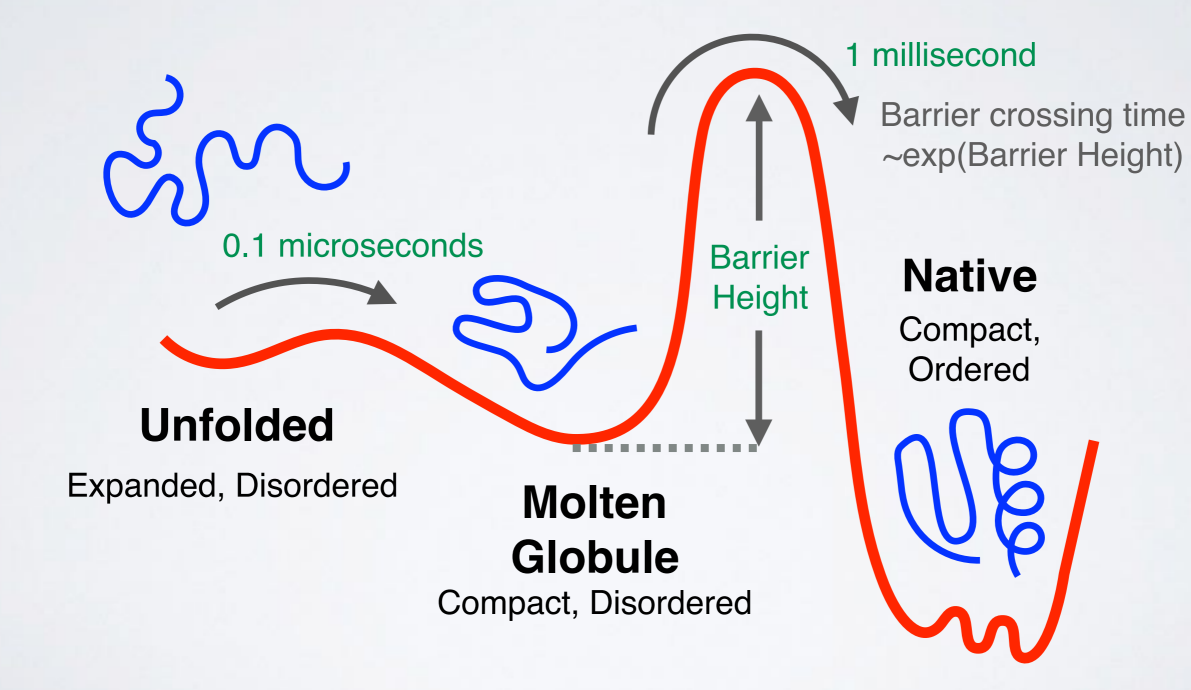


Expanded, Disordered

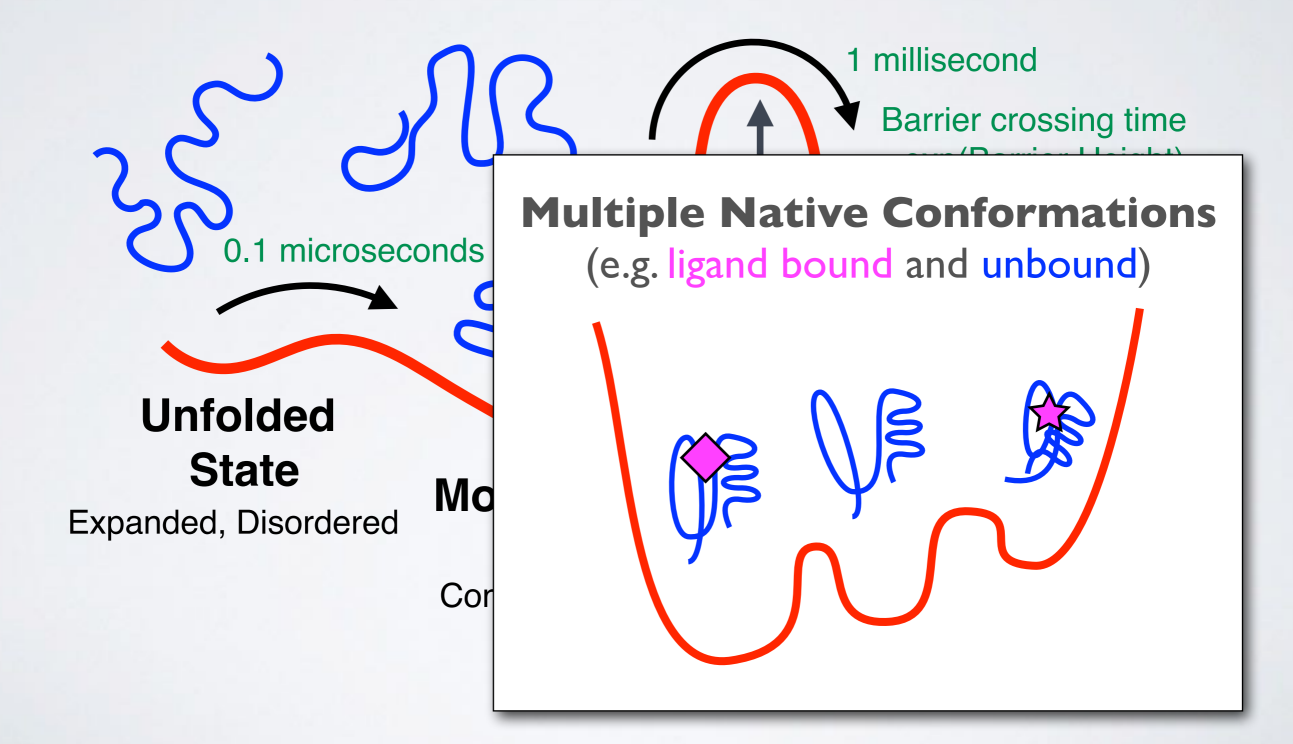
#### Native

Compact, Ordered

### **KEY CONCEPT: ENERGY LANDSCAPE**



### KEY CONCEPT: ENERGY LANDSCAPE



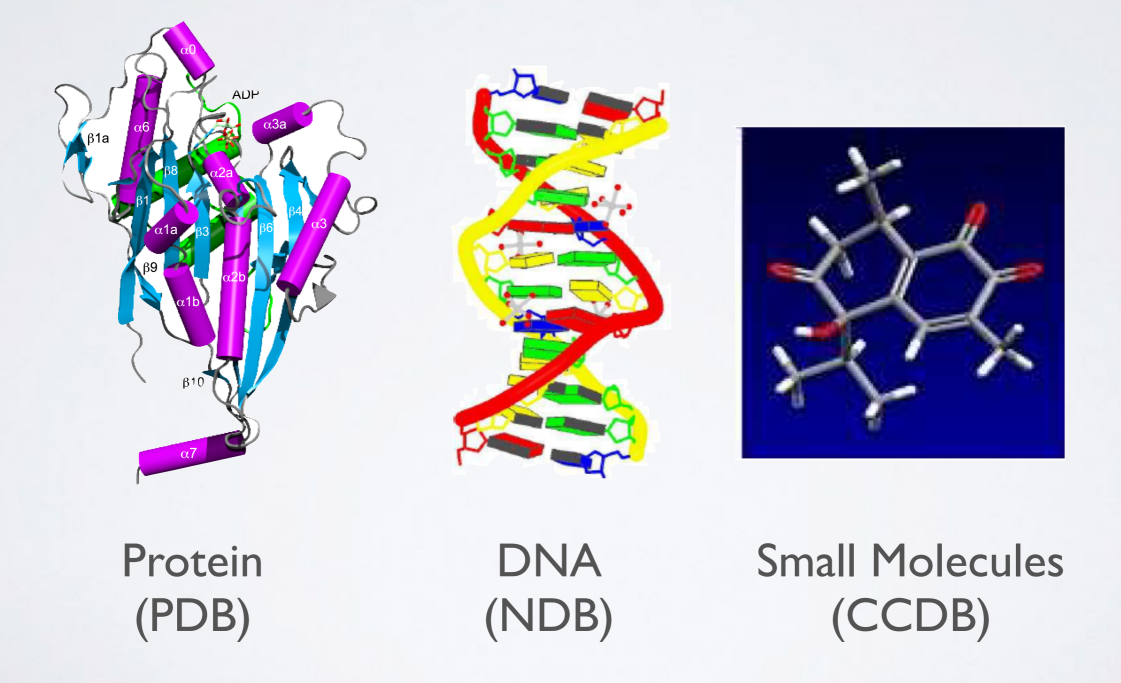
# **OUTLINE:**

- 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
  - Predicting functional dynamics & drug discovery

# **OUTLINE:**

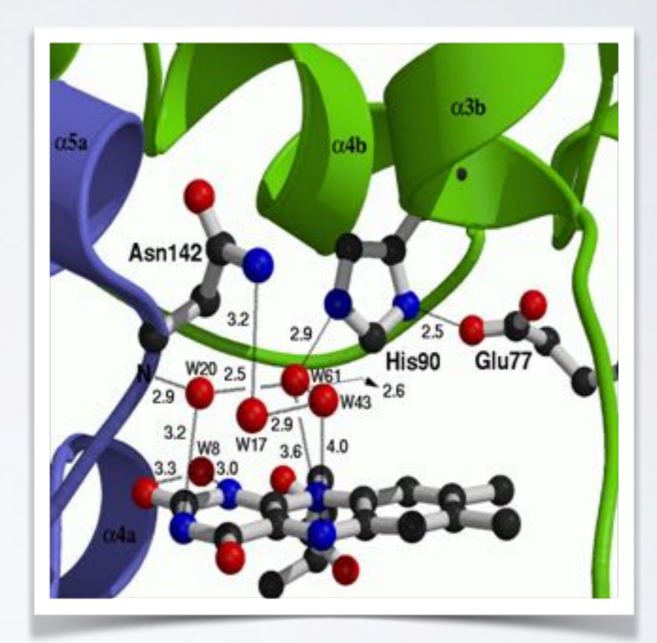
- Overview of structural bioinformatics
  - Major motivations, goals and challenges
- Fundamentals of protein structure
  - Composition, form, forces and dynamics
- Representing and interpreting protein structure
  - Modeling energy as a function of structure
- Example application areas
  - Predicting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

### TRADITIONAL FOCUS **PROTEIN**, **DNA** AND **SMALL MOLECULE** DATA SETS WITH **MOLECULAR STRUCTURE**



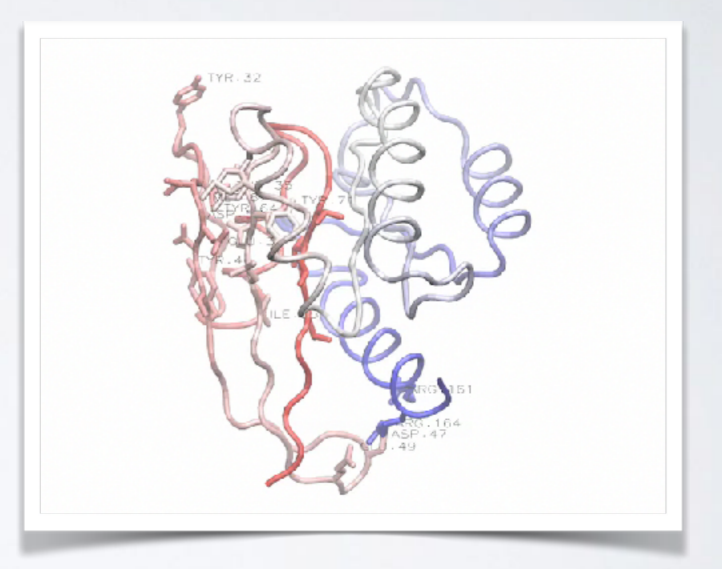
### Motivation 1: Detailed understanding of molecular interactions

Provides an invaluable structural context for conservation and mechanistic analysis leading to functional insight.



### Motivation 1: Detailed understanding of molecular interactions

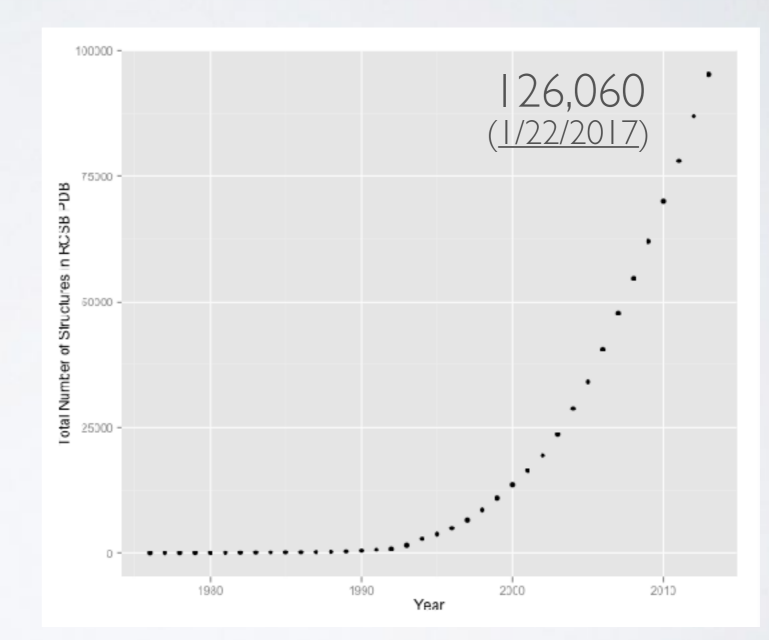
Computational modeling can provide detailed insight into functional interactions, their regulation and potential consequences of perturbation.



Grant et al. PLoS. Comp. Biol. (2010)

### Motivation 2: Lots of structural data is becoming available

Structural Genomics has contributed to driving down the cost and time required for structural determination



Data from: <u>http://www.rcsb.org/pdb/statistics/</u>

### Motivation 2: Lots of structural data is becoming available

Structural Genomics has contributed to driving down the cost and time required for structural determination

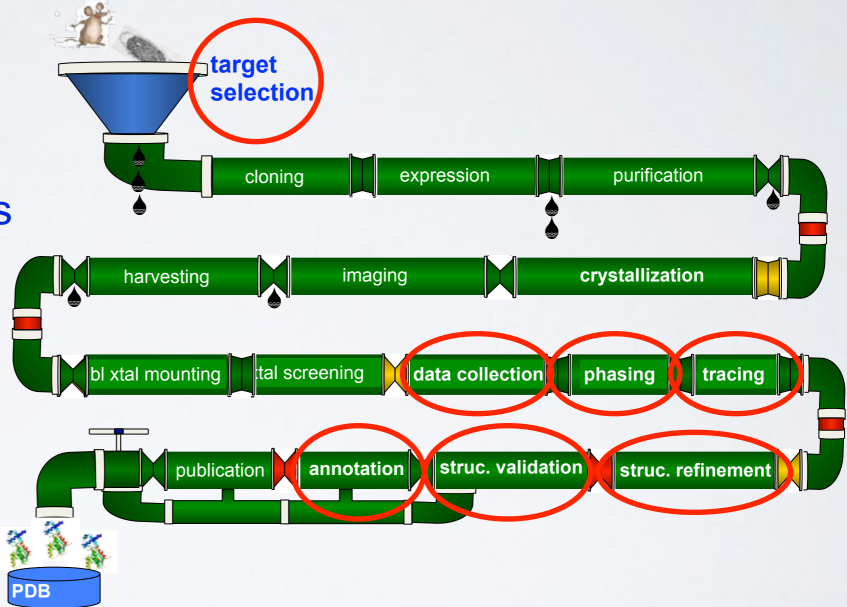
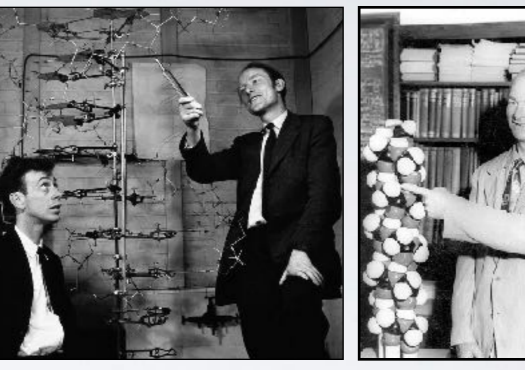


Image Credit: "Structure determination assembly line" Adam Godzik

### **Motivation 3:**

Theoretical and computational predictions have been, and continue to be, enormously valuable and influential!





### SUMMARY OF KEY MOTIVATIONS

#### Sequence > Structure > Function

 Structure determines function, so understanding structure helps our understanding of function

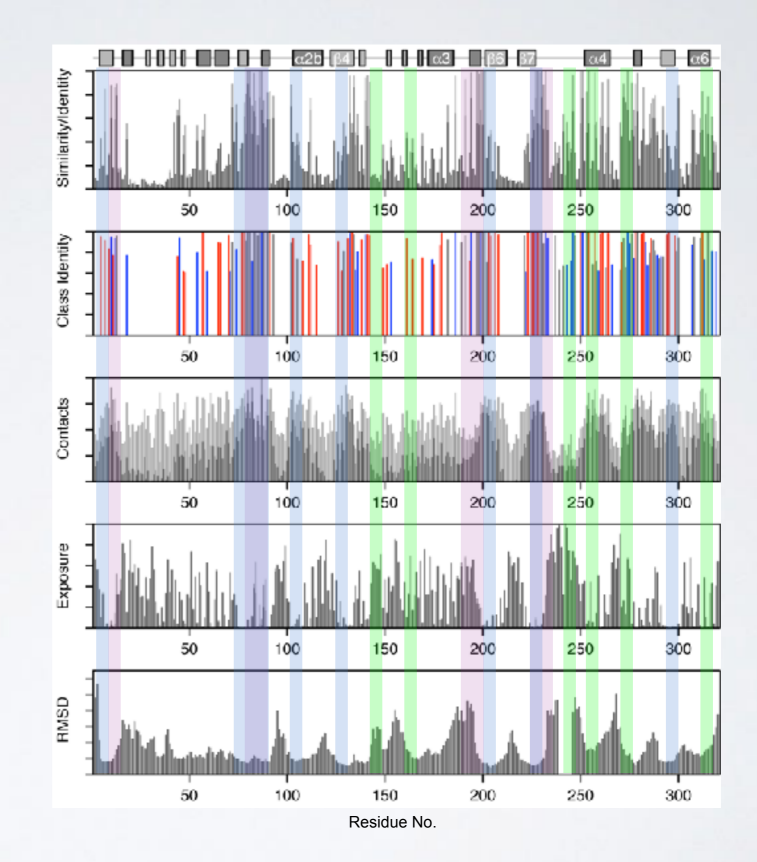
#### Structure is more conserved than sequence

Structure allows identification of more distant evolutionary relationships

#### Structure is encoded in sequence

 Understanding the determinants of structure allows design and manipulation of proteins for industrial and medical advantage

- Analysis
- Visualization
- Comparison
- Prediction
- Design



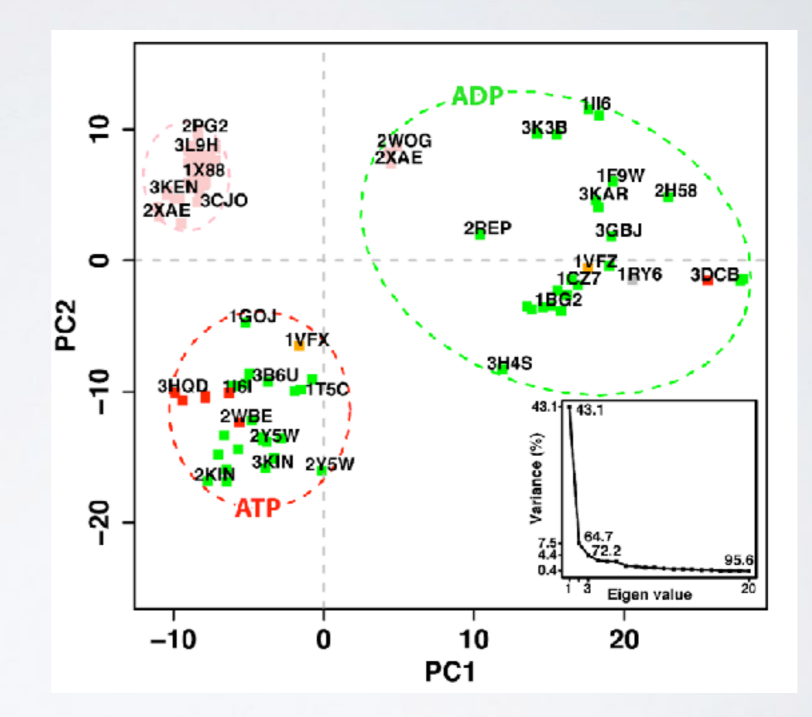
Grant et al. JMB. (2007)

- Analysis
- Visualization
- Comparison
- Prediction
- Design



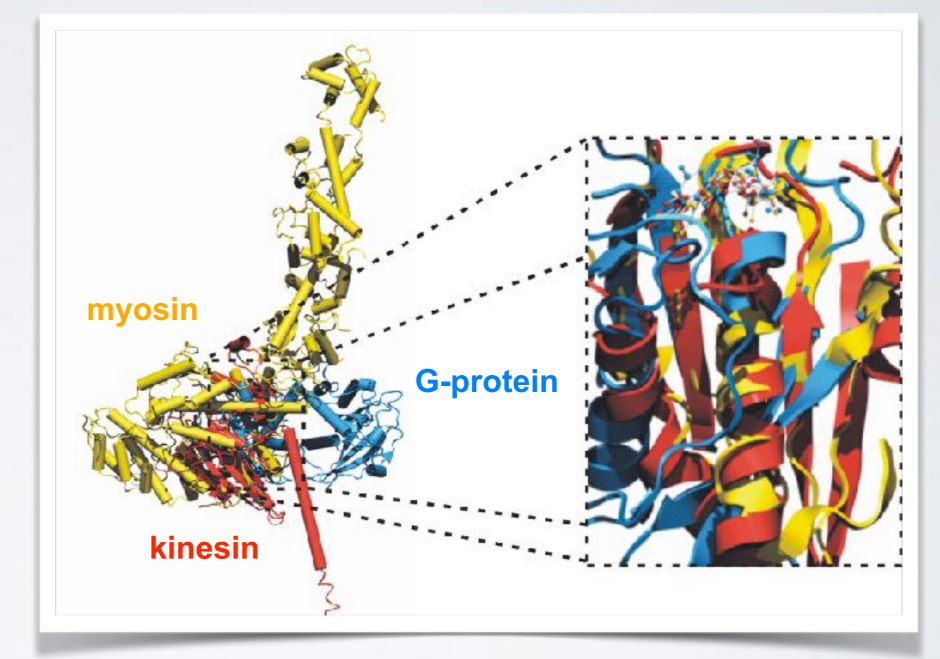
Scarabelli and Grant. PLoS. Comp. Biol. (2013)

- Analysis
- Visualization
- Comparison
- Prediction
- Design



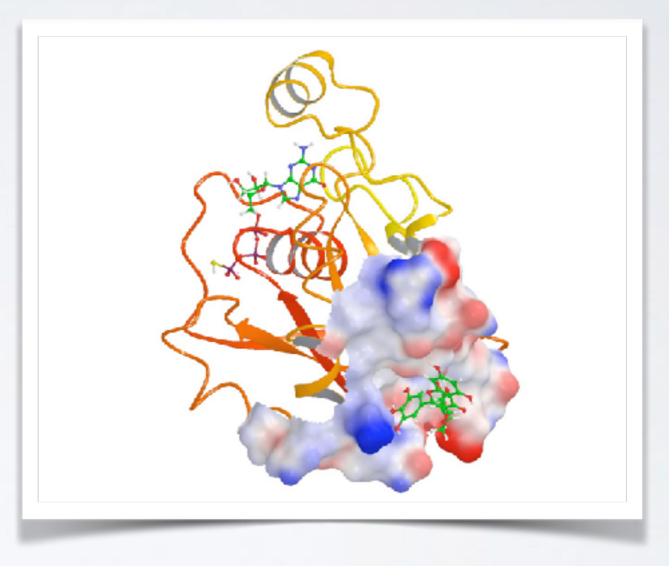
Scarabelli and Grant. PLoS. Comp. Biol. (2013)

- Analysis
- Visualization
- Comparison
- Prediction
- Design



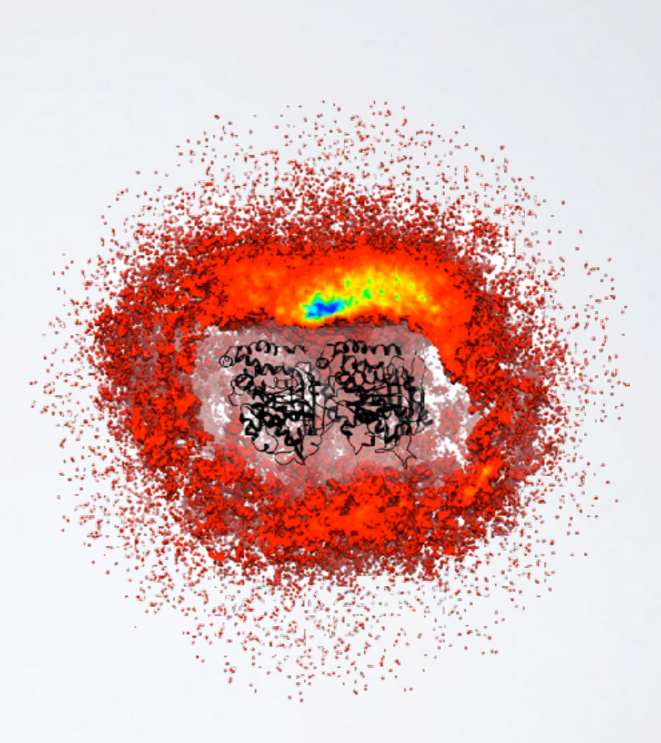
#### Grant et al. unpublished

- Analysis
- Visualization
- Comparison
- Prediction
- Design



Grant et al. PLoS One (2011, 2012)

- Analysis
- Visualization
- Comparison
- Prediction
- Design



Grant et al. PLoS Biology (2011)

# MAJOR RESEARCH AREAS AND CHALLENGES

Include but are not limited to:

- Protein classification
- Structure prediction from sequence
- Binding site detection
- Binding prediction and drug design
- Modeling molecular motions
- Predicting physical properties (stability, binding affinities)
- Design of structure and function
- etc...

With applications to Biology, Medicine, Agriculture and Industry

### 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
  - Predicting functional dynamics & drug discovery

### HIERARCHICAL STRUCTURE OF PROTEINS

#### Primary > Secondary > Tertiary > Quaternary

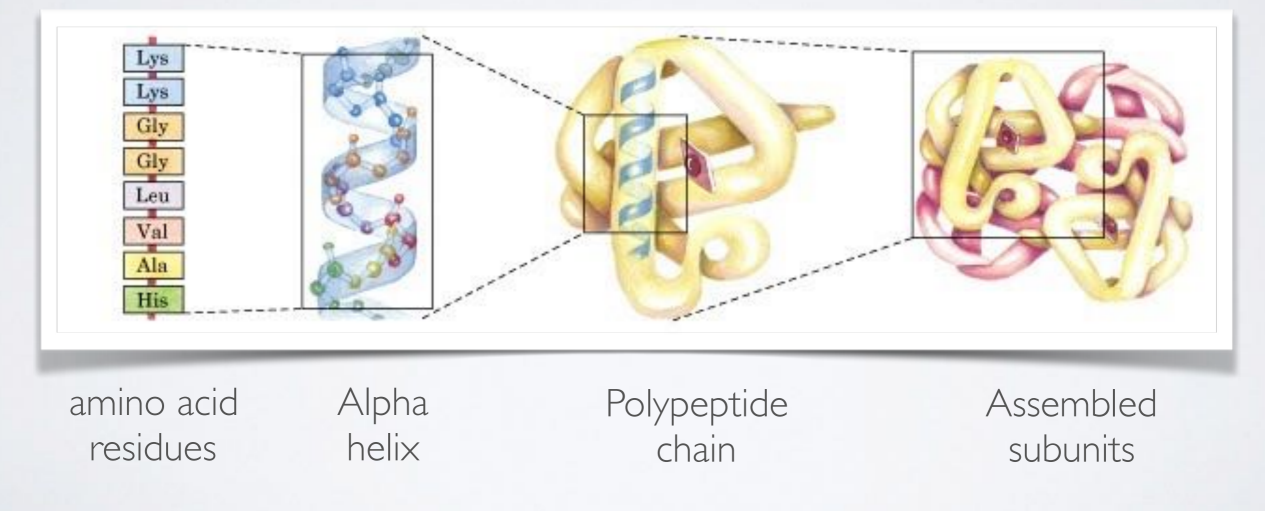
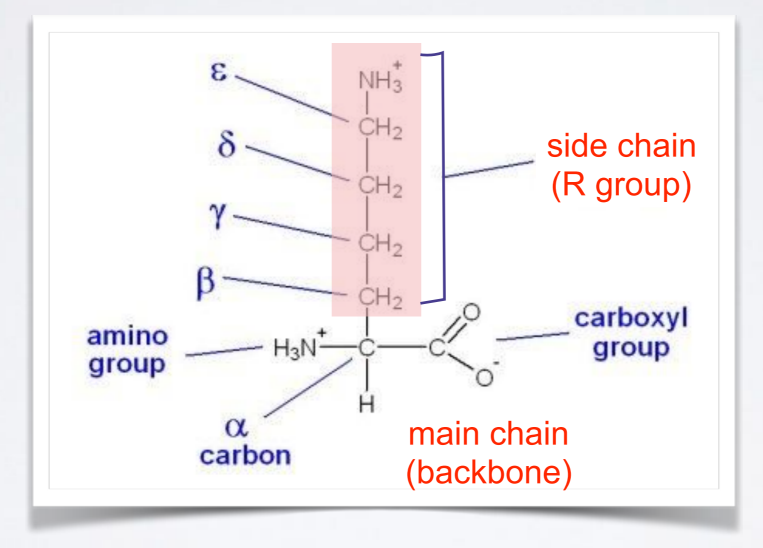
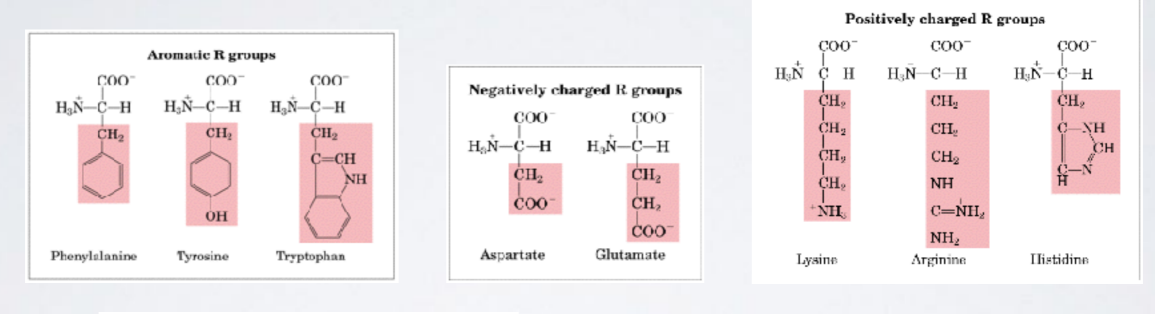


Image from: http://www.ncbi.nlm.nih.gov/books/NBK21581/

### RECAP: AMINO ACID NOMENCLATURE



### AMINO ACIDS CAN BE GROUPED BY THE Physiochemical properties



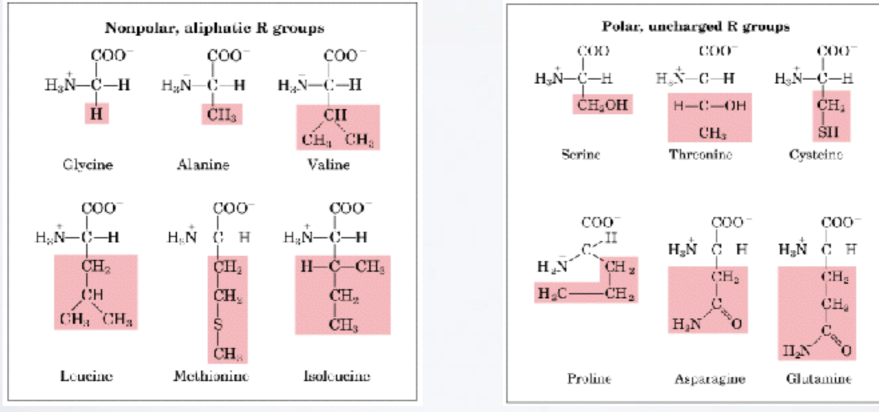


Image from: http://www.ncbi.nlm.nih.gov/books/NBK21581/

### AMINO ACIDS POLYMERIZE THROUGH PEPTIDE BOND FORMATION

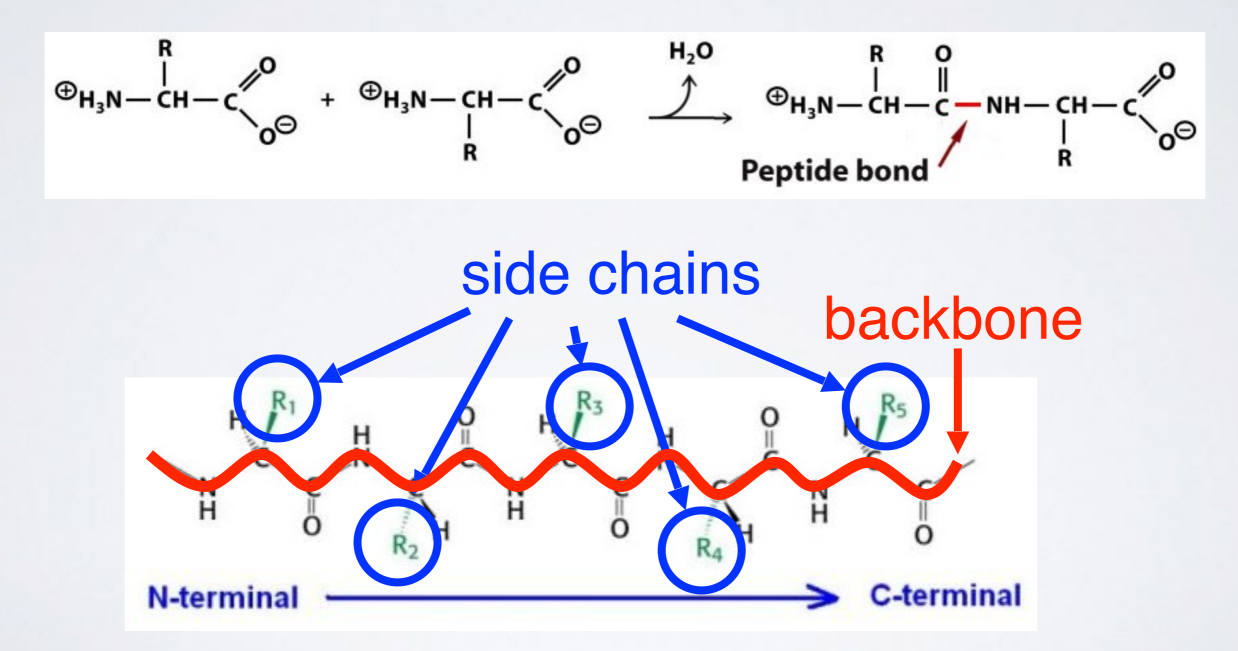


Image from: http://www.ncbi.nlm.nih.gov/books/NBK21581/

### PEPTIDES CAN ADOPT DIFFERENT CONFORMATIONS BY VARYING THEIR PHI & PSI BACKBONE TORSIONS

Ψ

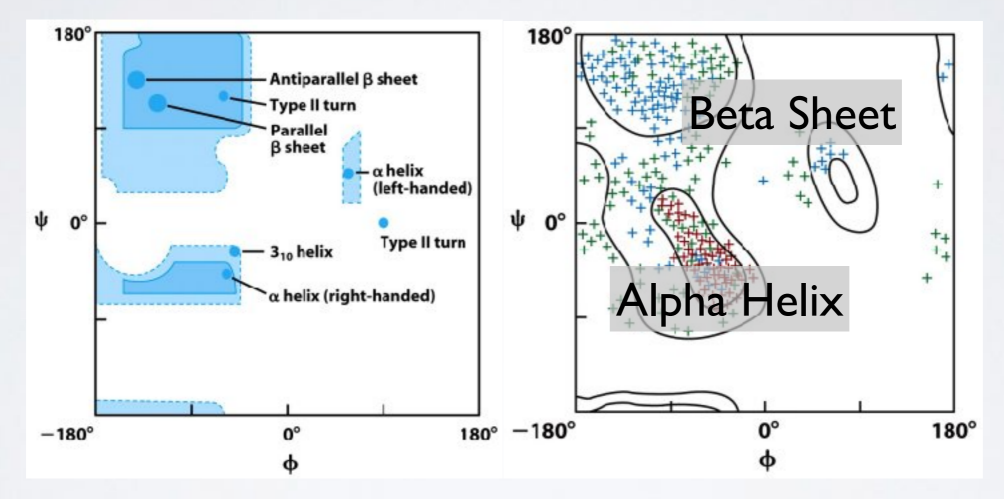
П

**N-terminal** 

Bond angles and lengths are largely invariant C-terminal

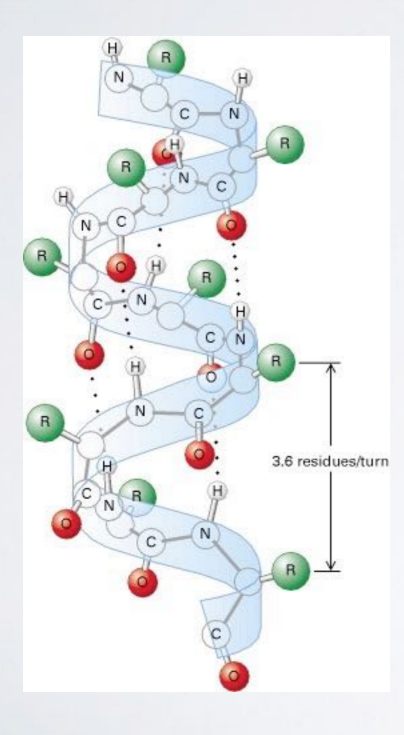
Peptide bond is planer (Ca, C, O, N, H, Ca all lie in the same plane)

# PHI vs PSI PLOTS ARE KNOWN AS RAMACHANDRAN DIAGRAMS



- Steric hindrance dictates torsion angle preference
- Ramachandran plot show preferred regions of φ and ψ dihedral angles which correspond to major forms of secondary structure

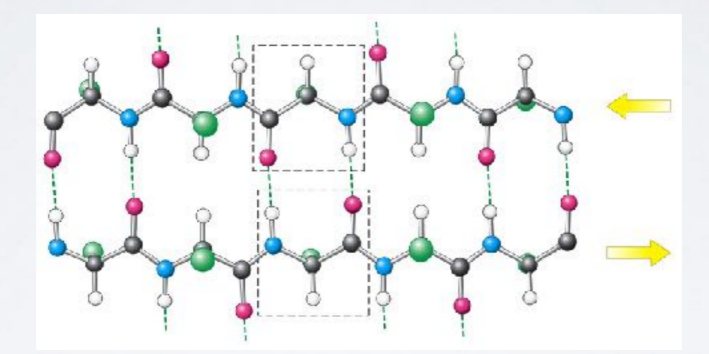
### MAJOR SECONDARY STRUCTURE TYPES ALPHA HELIX & BETA SHEET



#### a-helix

- Most common from has <u>3.6 residues per</u> <u>turn</u> (number of residues in one full rotation)
- Hydrogen bonds (dashed lines) between residue <u>i and i+4</u> stabilize the structure
- The side chains (in green) protrude
   outward
- 3<sub>10</sub>-helix and π-helix forms are less common

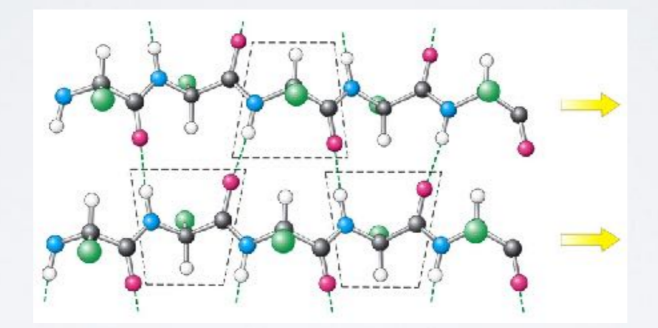
### MAJOR SECONDARY STRUCTURE TYPES ALPHA HELIX & **BETA SHEET**



#### In **antiparallel** $\beta$ -sheets

- Adjacent β-strands run in <u>opposite</u> directions
- Hydrogen bonds (dashed lines) between NH and CO stabilize the structure
- The side chains (in green) are above and below the sheet
   Image from: <u>http://www.ncbi.nlm.nih.gov/books/NBK21581/</u>

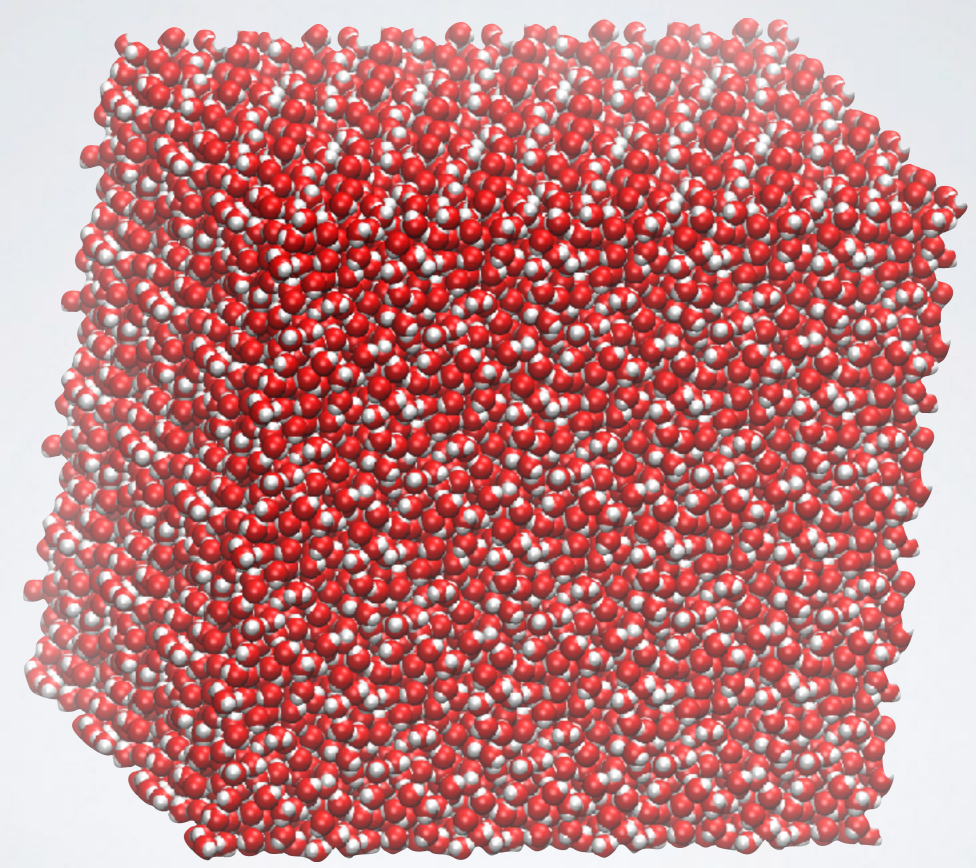
### MAJOR SECONDARY STRUCTURE TYPES ALPHA HELIX & **BETA SHEET**



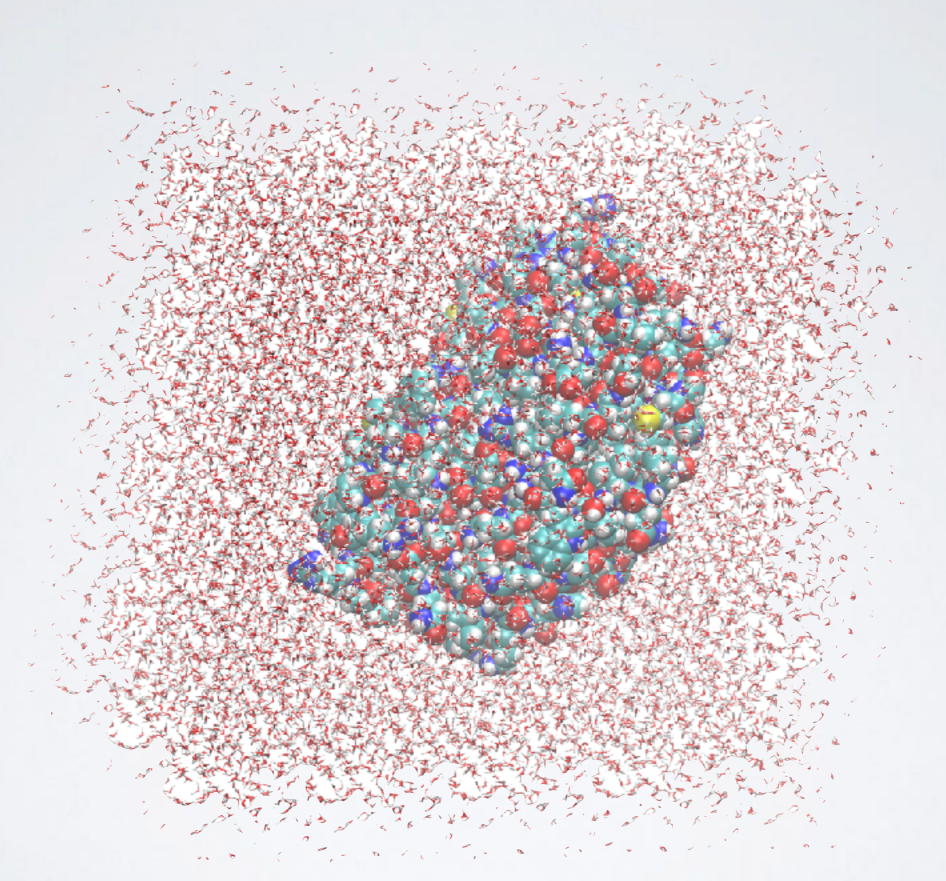
#### In **parallel** $\beta$ -sheets

- Adjacent β-strands run in <u>same</u> direction
- Hydrogen bonds (dashed lines) between NH and CO stabilize the structure
- The side chains (in green) are above and below the sheet
   Image from: <u>http://www.ncbi.nlm.nih.gov/books/NBK21581/</u>

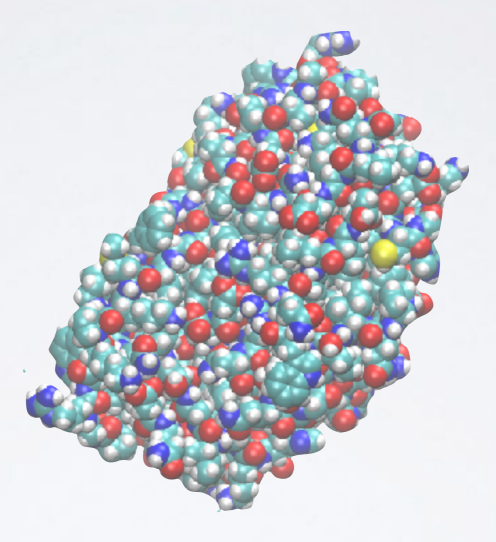
### What Does a Protein Look like?



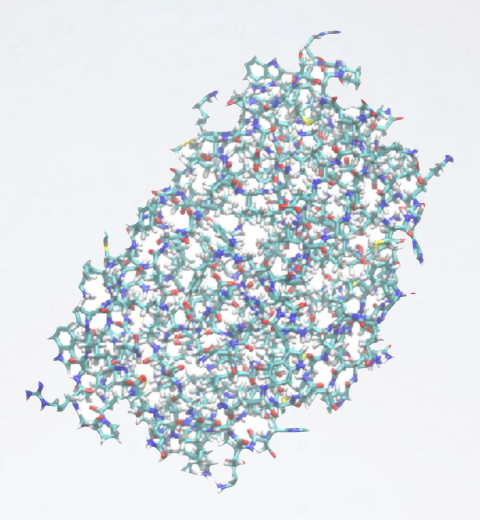
• Proteins are stable (and hidden) in water



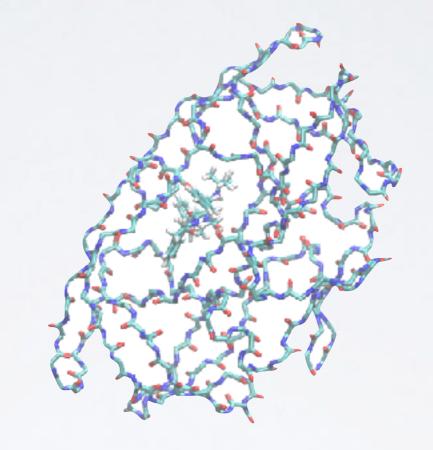
Proteins closely interact with water



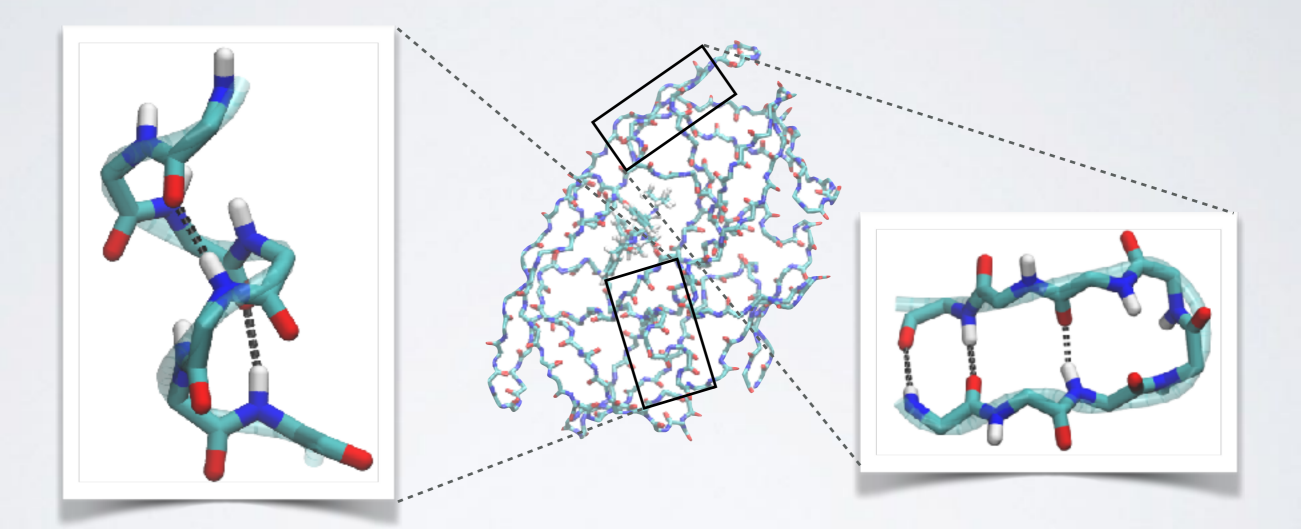
Proteins are close packed solid but flexible objects (globular)



 Due to their large size and complexity it is often hard to see whats important in the structure



 Backbone or main-chain representation can help trace chain topology

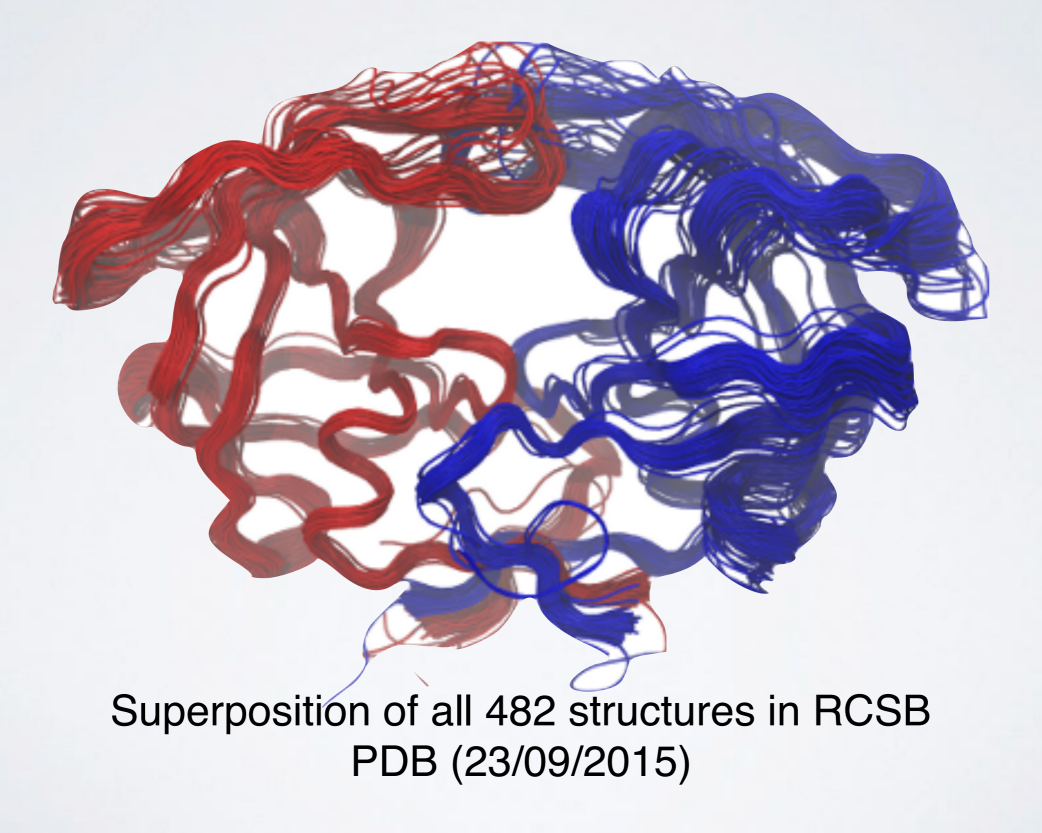


 Backbone or main-chain representation can help trace chain topology & reveal secondary structure

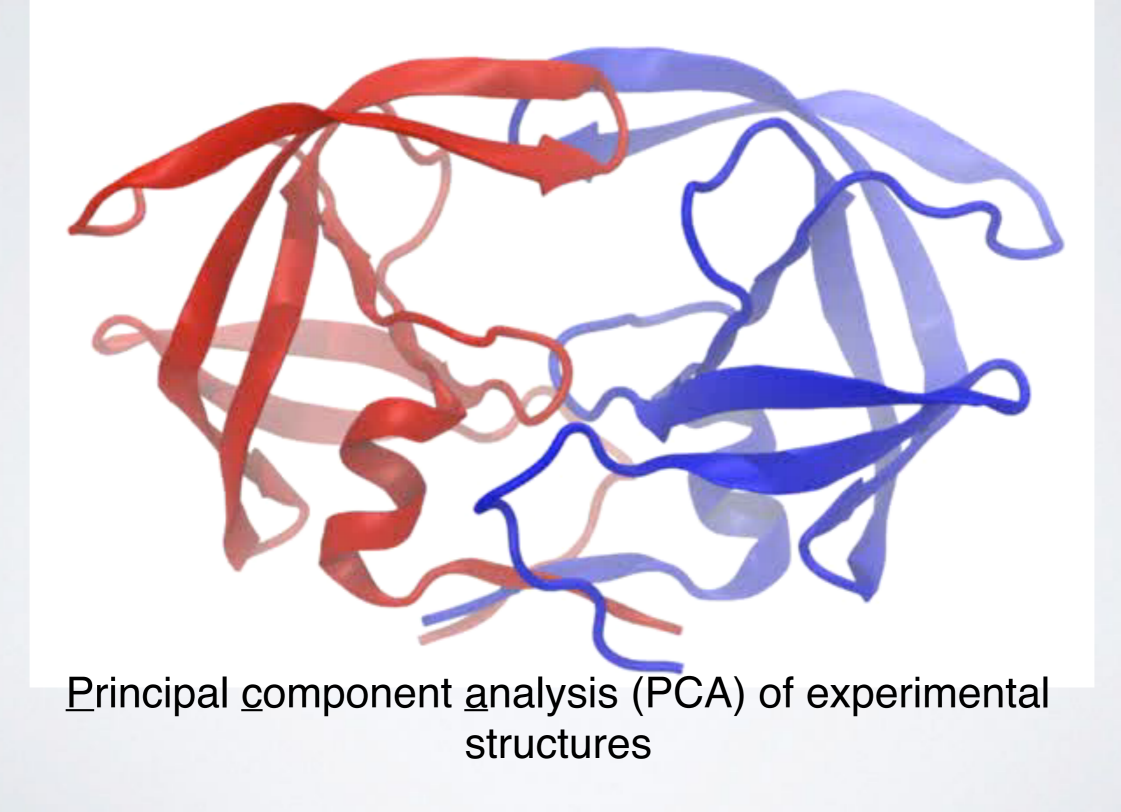


- Simplified secondary structure representations are commonly used to communicate structural details
- Now we can clearly see 2°, 3° and 4° structure
- Coiled chain of connected secondary structures

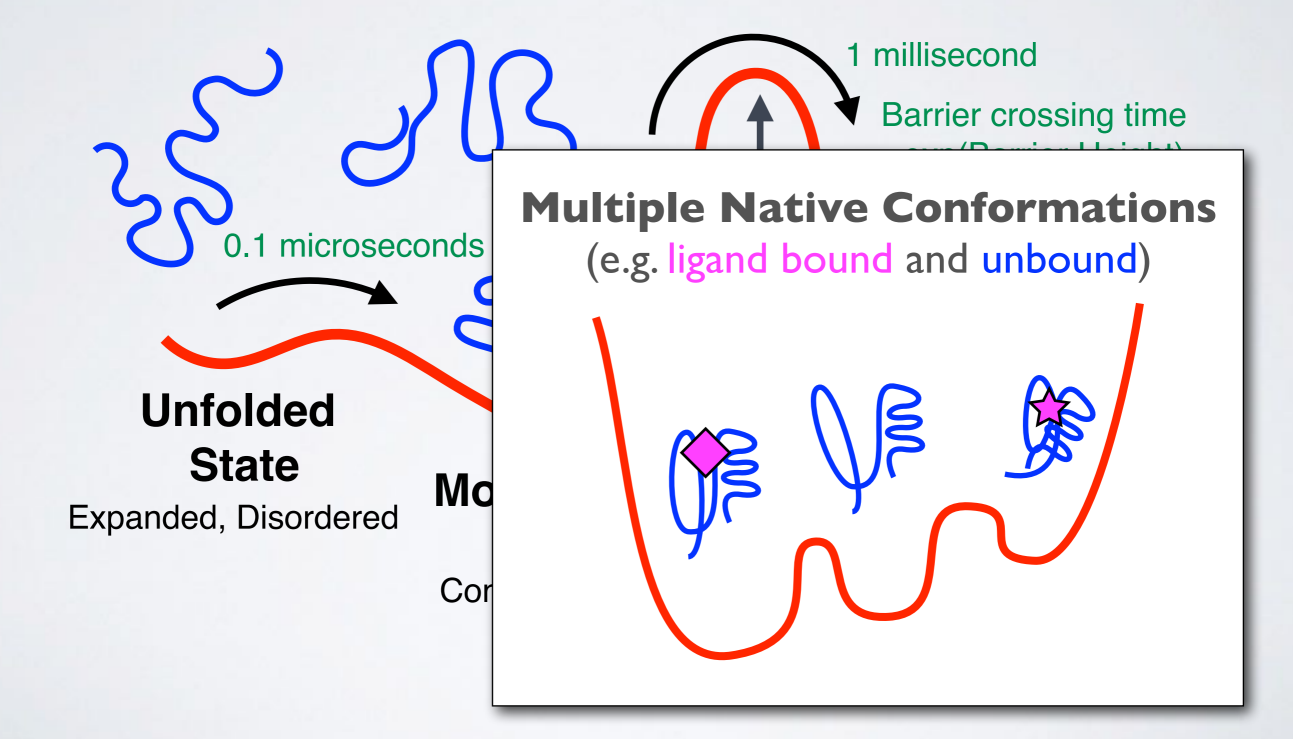
### DISPLACEMENTS REFLECT INTRINSIC FLEXIBILITY



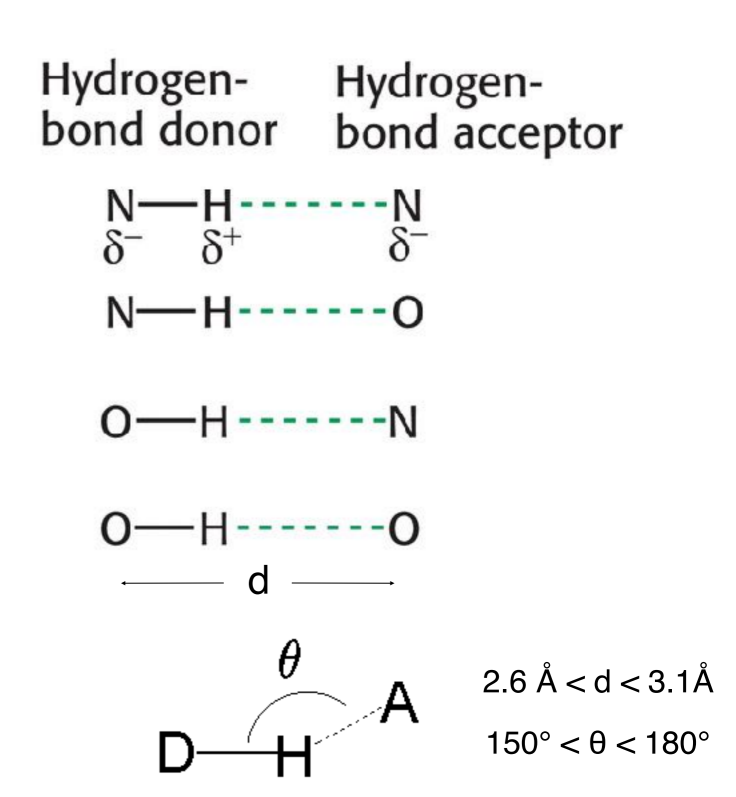
### DISPLACEMENTS REFLECT INTRINSIC FLEXIBILITY

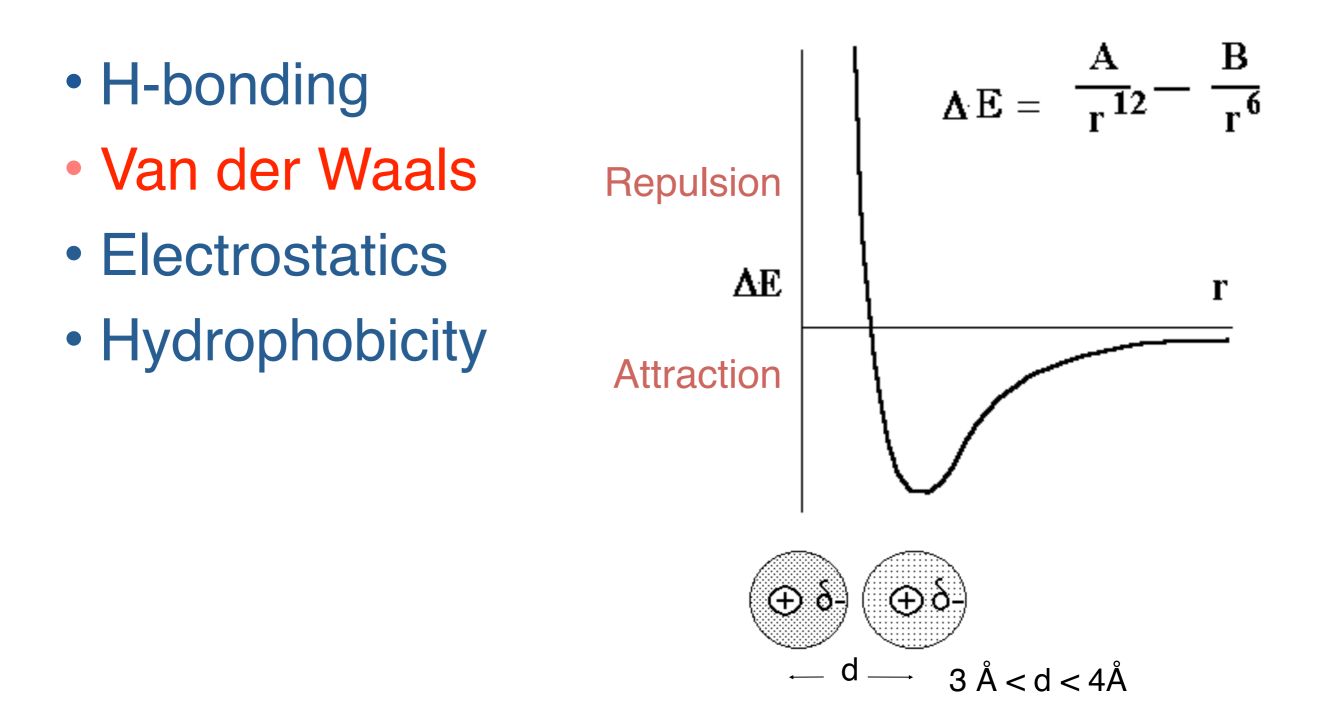


### KEY CONCEPT: ENERGY LANDSCAPE

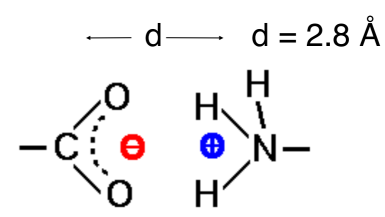


- H-bonding
- Van der Waals
- Electrostatics
- Hydrophobicity





- H-bonding
- Van der Waals
- Electrostatics
- Hydrophobicity

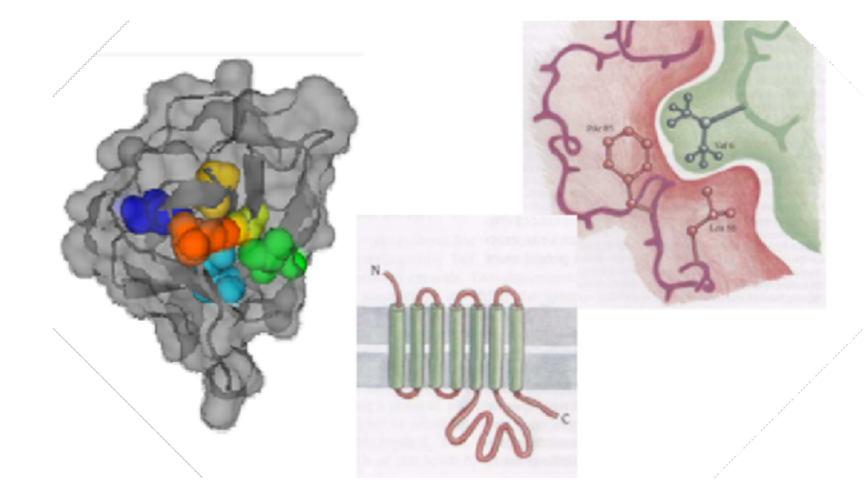


#### carboxyl group and amino group

(some time called IONIC BONDs or SALT BRIDGEs)

$\begin{array}{c} \mathbf{Co} \\ \mathbf{q}_1 & \mathbf{q}_2 \\ \mathbf{O} & \mathbf{r} & \mathbf{O} \\ \mathbf{Co} & \mathbf{r} \end{array} \mathbf{E} \end{array}$	ulomb's law = Kq1q2 Dr	E = Energy k = constant D = Dielectric constant (vacuum = 1; $H_2O = 80$ ) $q_1 \& q_2 =$ electronic charges (Coulombs) r = distance (Å)
--	------------------------------	--

- H-bonding
- Van der Waals
- Electrostatics
- Hydrophobicity



The force that causes hydrophobic molecules or nonpolar portions of molecules to aggregate together rather than to dissolve in water is called <u>Hydrophobicity</u> (*Greek, "water fearing"*). This is not a separate bonding force; rather, it is the result of the energy required to insert a nonpolar molecule into water.

# Hand-on time! https://tinyurl.com/bimm143-L11

DO IN LOUIS OFF

Focus on section 1 to 3 and user your sticky notes for problems and questions please!

### NEXT UP:

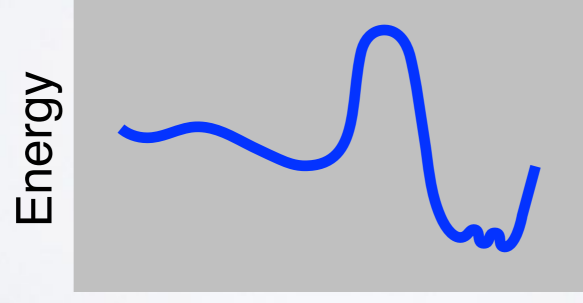
- Overview of structural bioinformatics
  - Major motivations, goals and challenges
- Fundamentals of protein structure
  - Composition, form, forces and dynamics
- Representing and interpreting protein structure
  - Modeling energy as a function of structure
- Example application areas
  - Predicting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

### **KEY CONCEPT:** POTENTIAL FUNCTIONS DESCRIBE A SYSTEMS **ENERGY** AS A FUNCTION OF ITS **STRUCTURE**

Two main approaches: (1). **Physics-Based** (2). **Knowledge-Based** 

### KEY CONCEPT: POTENTIAL FUNCTIONS DESCRIBE A SYSTEMS ENERGY AS A FUNCTION OF ITS STRUCTURE

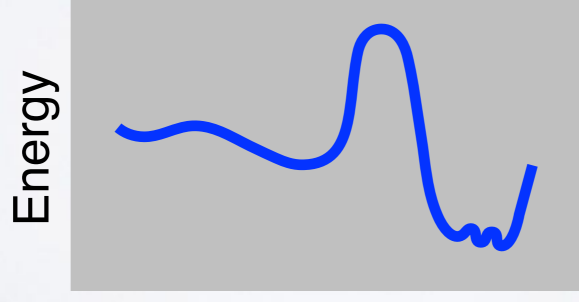
Two main approaches: (1). **Physics-Based** (2). **Knowledge-Based** 



Structure/Conformation

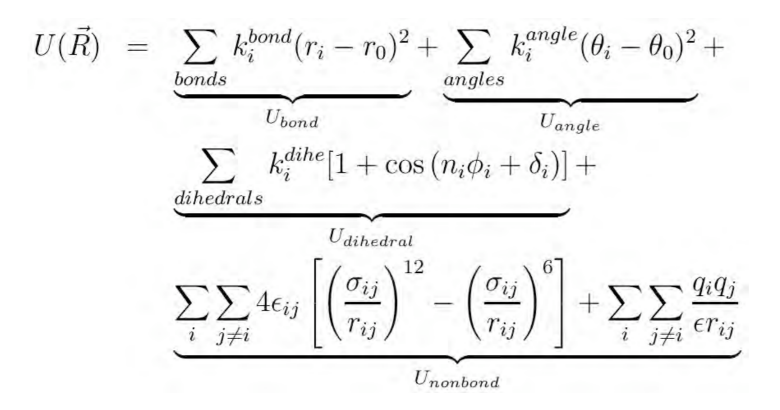
### KEY CONCEPT: POTENTIAL FUNCTIONS DESCRIBE A SYSTEMS ENERGY AS A FUNCTION OF ITS STRUCTURE

Two main approaches: (1). Physics-Based (2). Knowledge-Based



Structure/Conformation

### PHYSICS-BASED POTENTIALS ENERGY TERMS FROM PHYSICAL THEORY

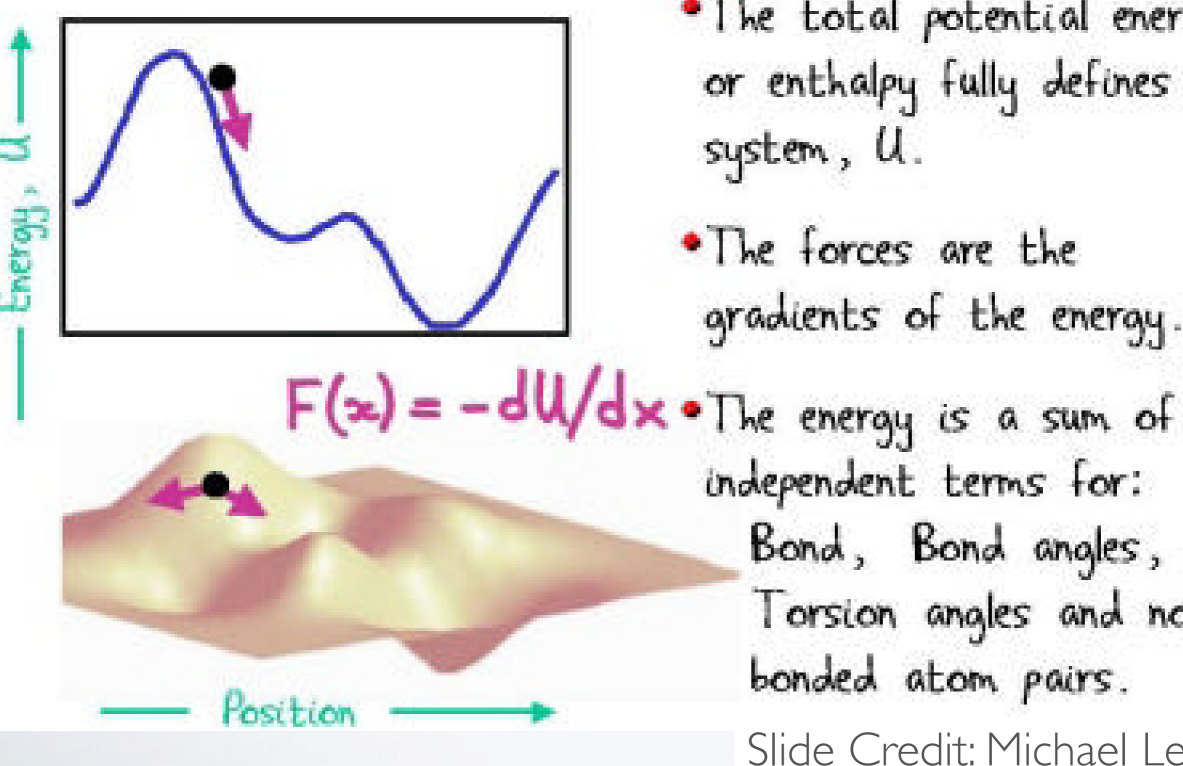


+..

 $U_{bond}$  = oscillations about the equilibrium bond length  $U_{angle}$  = oscillations of 3 atoms about an equilibrium bond angle  $U_{dihedral}$  = torsional rotation of 4 atoms about a central bond  $U_{nonbond}$  = non-bonded energy terms (electrostatics and Lenard-Jones)

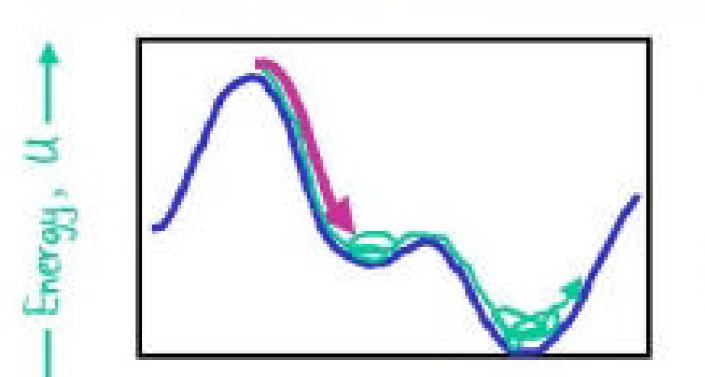
CHARMM P.E. function, see: http://www.charmm.org/

## TOTAL POTENTIAL ENERGY



- The total potential energy or enthalpy fully defines the system, U.
- The forces are the gradients of the energy.
- independent terms for:
  - Bond, Bond angles, Torsion angles and nonbonded atom pairs.
  - Slide Credit: Michael Levitt

# MOVING OVER THE ENERGY SURFACE



Position

- 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 (-∆U/kT).

Slide Credit: Michael Levitt

### 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

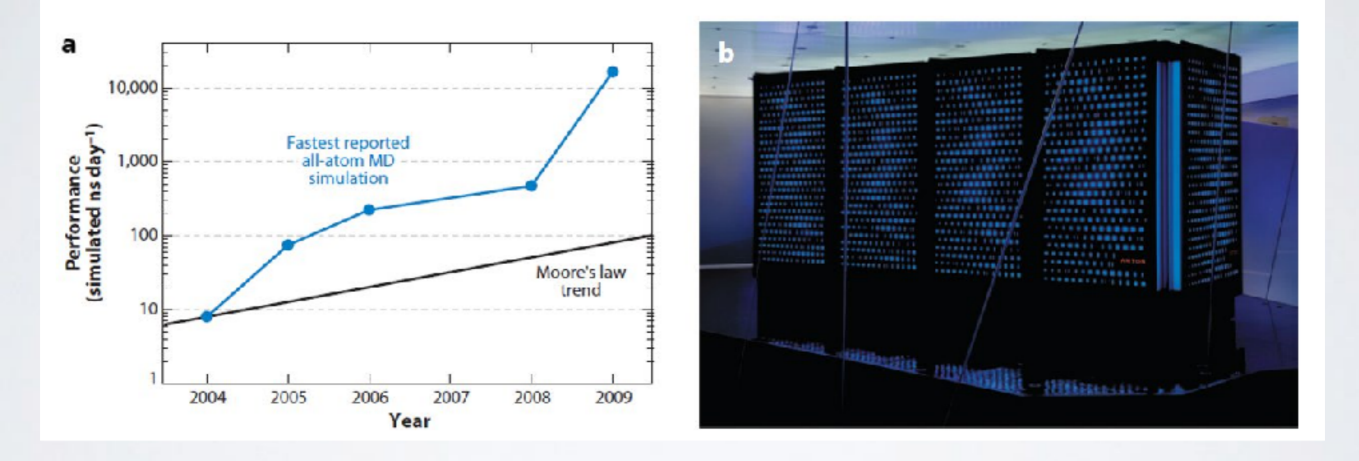
#### Status

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

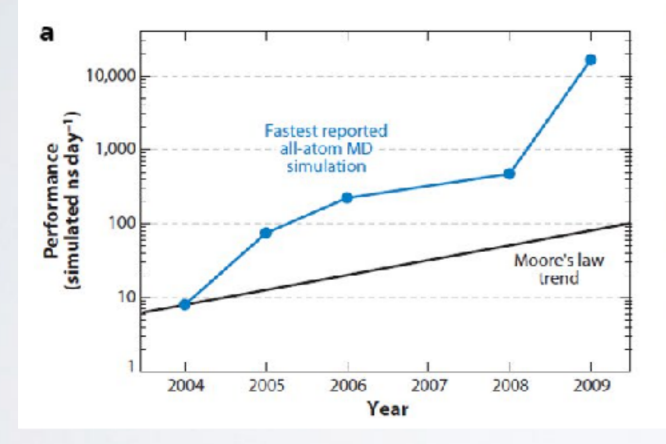
1967	HUOM	0.1 MH	1 MB	HALL
1013	14,000	1643	10 68	LAPTOP
CHANNEE		10,000	10,000	10,000

It cars were the computers then a new Volic would cost \$3, would have a top speed of 1,000,000 Km/hr, would carry 50,000 adults and would park in a shocker.

# SUPERCOMPUTER



# SUPERCOMPUTER

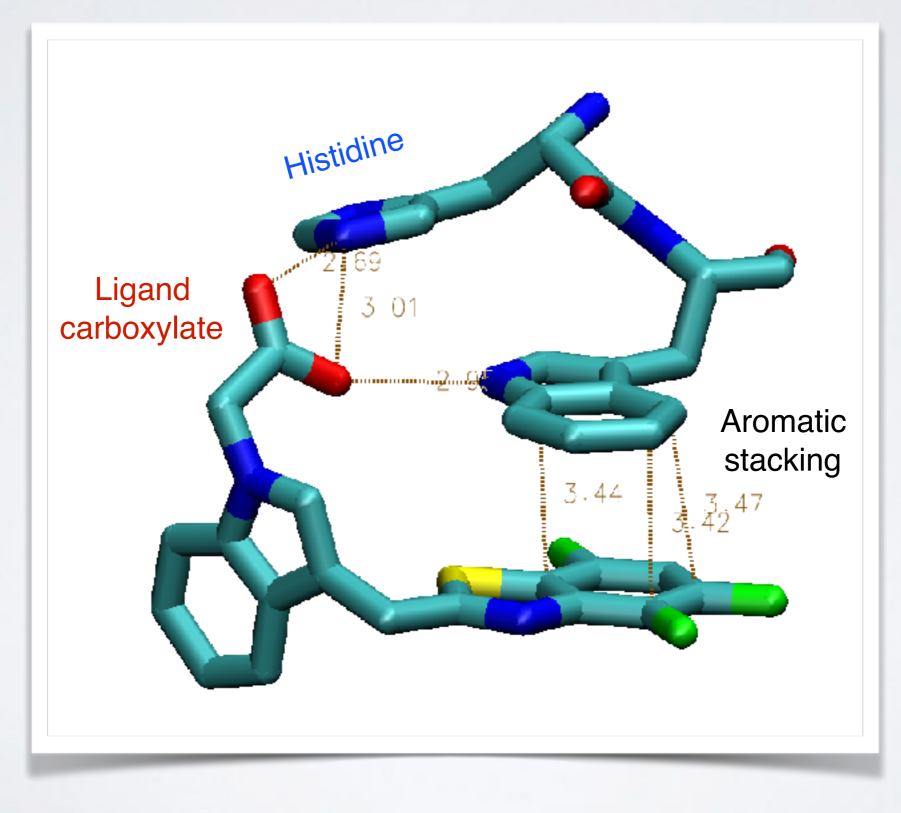




### **KEY CONCEPT:** POTENTIAL FUNCTIONS DESCRIBE A SYSTEMS **ENERGY** AS A FUNCTION OF ITS **STRUCTURE**

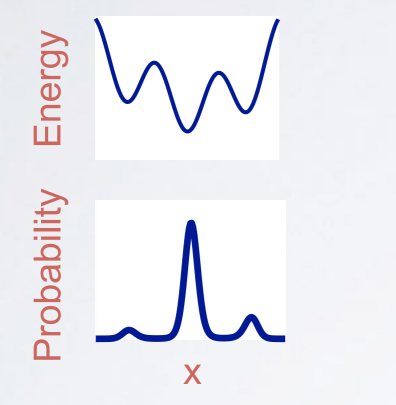
Two main approaches: (1). Physics-Based (2). Knowledge-Based

### 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:  $E(r) = -RT \ln \left[ p(r) \right]$ 

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

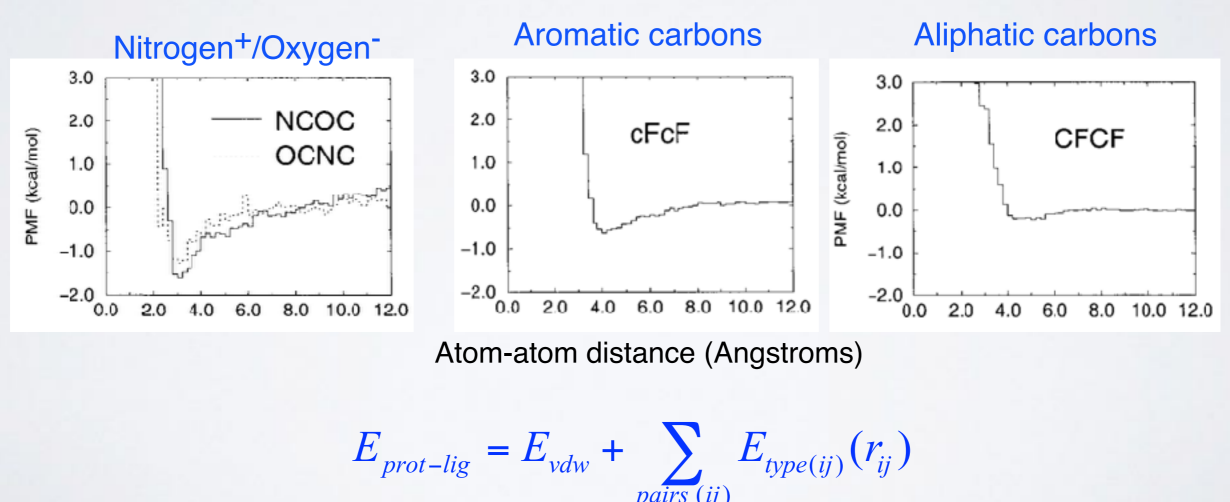
Find all protein-ligand structures in the PDB with a ligand carboxylate O

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

# KNOWLEDGE-BASED DOCKING POTENTIALS

"PMF", Muegge & Martin, J. Med. Chem. (1999) 42:791

A few types of atom pairs, out of several hundred total



### 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)

# Hand-on time! https://tinyurl.com/bimm143-L11

Do in Louis Solar

Focus on section 4 & 5

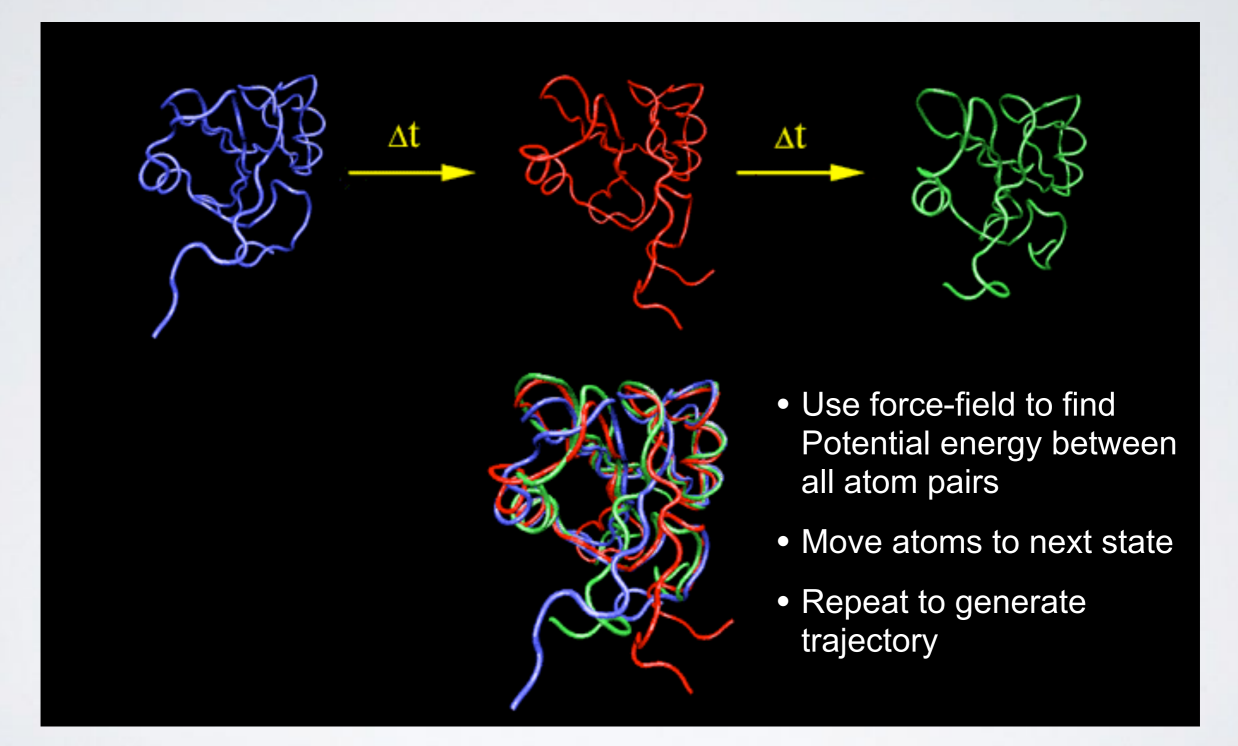
# NEXT UP:

- 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** 
    - Predicting functional dynamics & drug discovery

### 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**



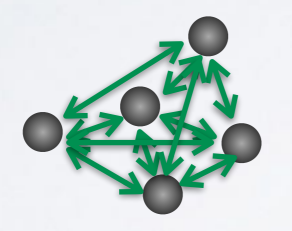
McCammon, Gelin & Karplus, Nature (1977) [ See: <u>https://www.youtube.com/watch?v=ui1ZysMFcKk</u> ] Divide time into discrete (~1fs) time steps (Δt) (for integrating equations of motion, see below)

										$\rightarrow$	ŧ
									1		L

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  

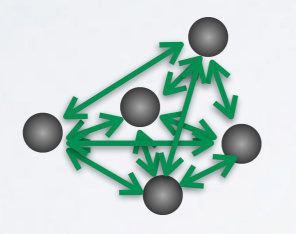
$$m_i \frac{d^2}{dt^2} \vec{R}_i = -\vec{\nabla}_i E(\vec{R})$$
Empirical force field  

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

Divide time into discrete (~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  

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

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

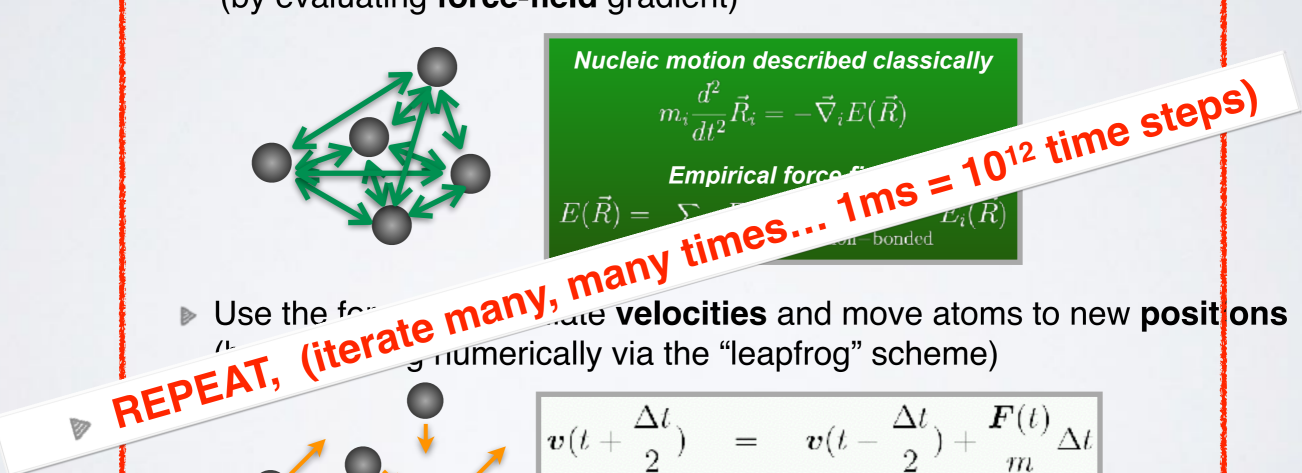
#### **BASIC ANATOMY OF A MD SIMULATION**

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

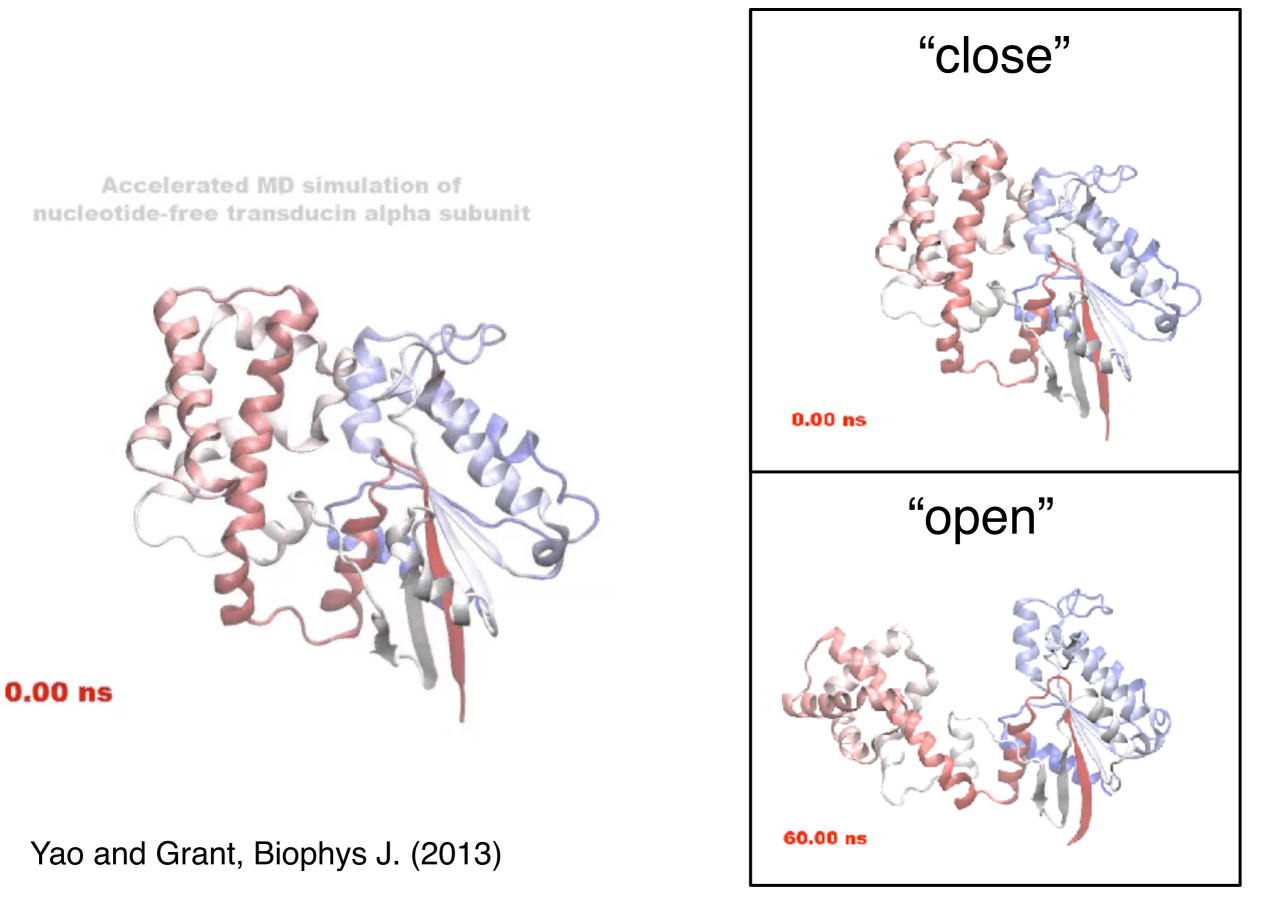
#### 

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

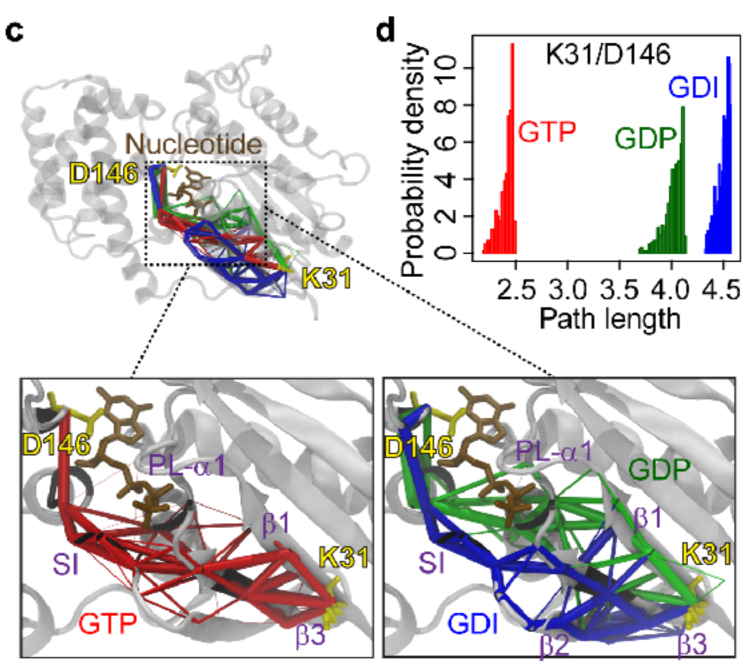
At each time step calculate pair-wise atomic forces (F(t)) (by evaluating force-field gradient)



# **MD** Prediction of Functional Motions

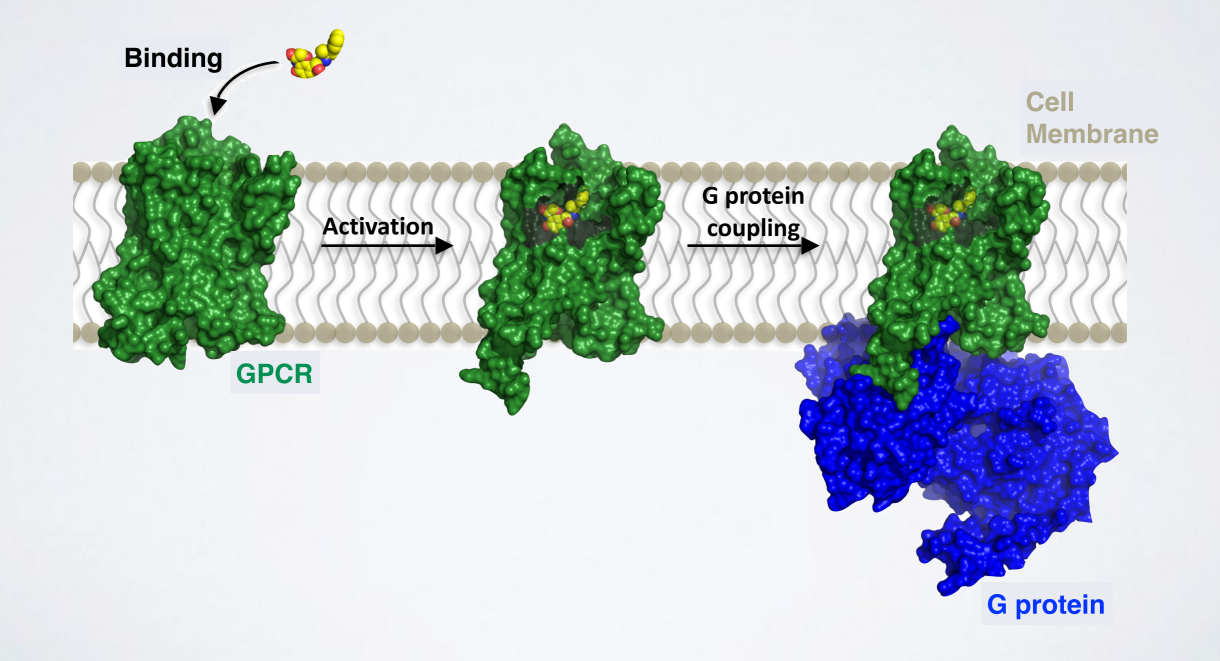


# Simulations Identify Key Residues Mediating Dynamic Activation

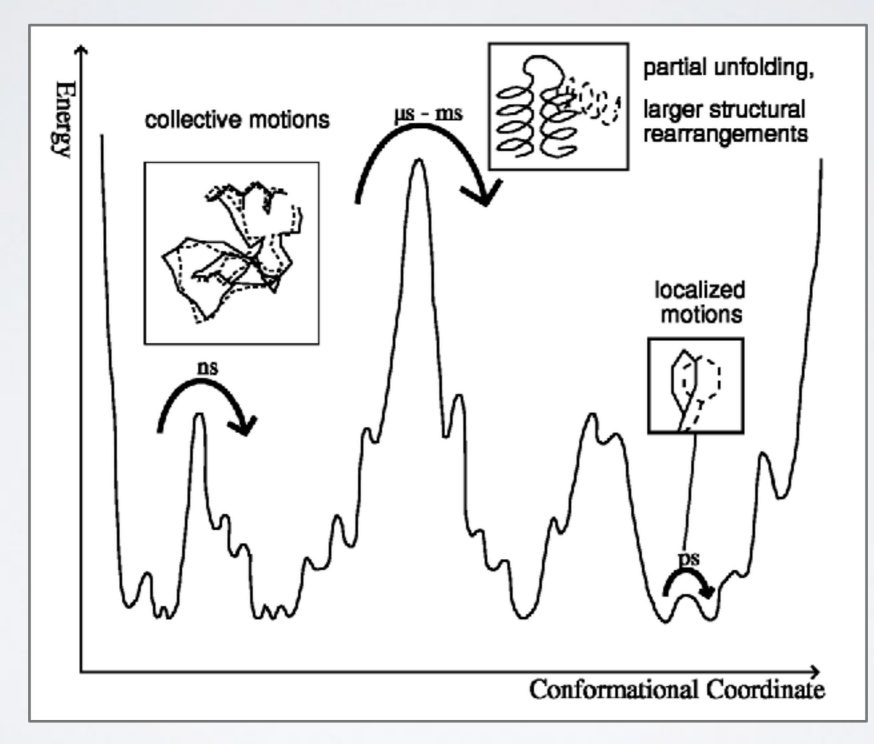


Yao ... Grant, <u>Journal of Biological Chemistry</u> (2016)

# EXAMPLE APPLICATION OF MOLECULAR SIMULATIONS TO GPCRS



#### PROTEINS JUMP BETWEEN MANY, HIERARCHICALLY ORDERED "CONFORMATIONAL SUBSTATES"



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

Improve this slide

#### **MOLECULAR DYNAMICS IS VERY**

**Example**: F<sub>1</sub>-ATPase in water (183,674 atoms) for 1 nanosecond:

=> 10<sup>6</sup> integration steps

=> 8.4 \* 10<sup>11</sup> floating point operations/step [n(n-1)/2 interactions]

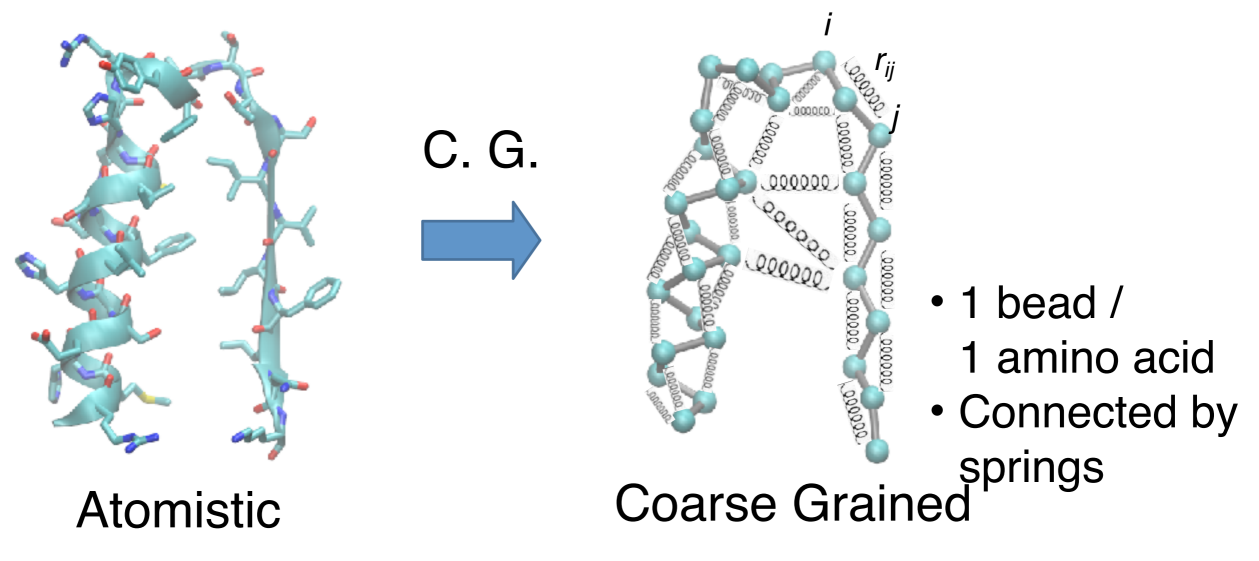
> Total: 8.4 \* 10<sup>17</sup> flop (on a 100 Gflop/s cpu: **ca 25 years!**)

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

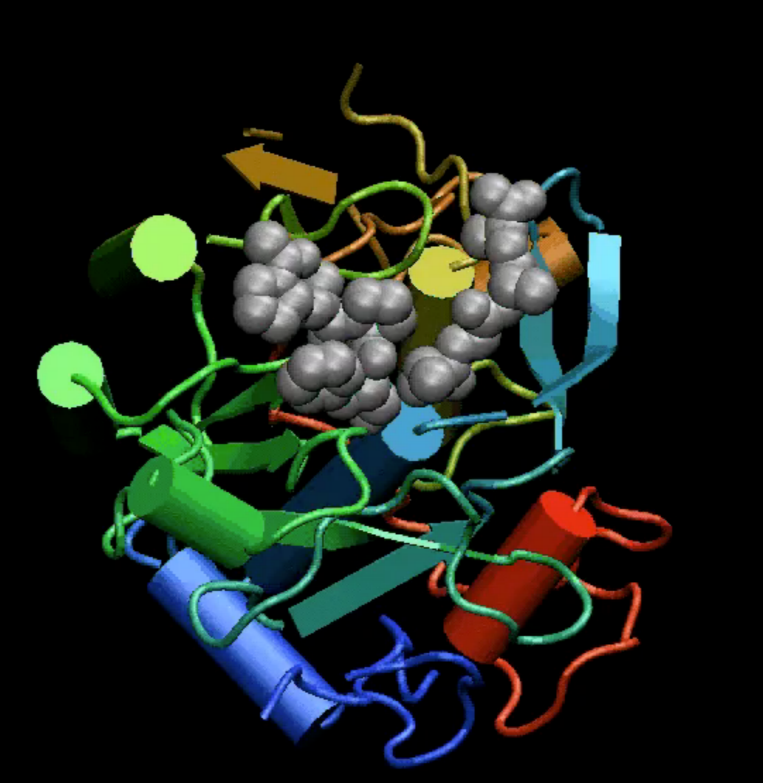
multiple time stepping	ca. 2.5 years
fast multipole methods	ca. 1 year
parallel computers	ca. 5 days
modern GPUs	ca. 1 day
(Anton supercomputer	ca. minutes)

#### 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

# Hand-on time! https://tinyurl.com/bimm143-L11

Do in Louis Solar

Focus on section 6 to 7

# 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

**Optional**: Stop here for Today!

# 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** 

Predicting functional dynamics & <u>drug discovery</u>

#### **CAUTIONARY NOTES**

 "Everything should be made as simple as it can be but not simpler"

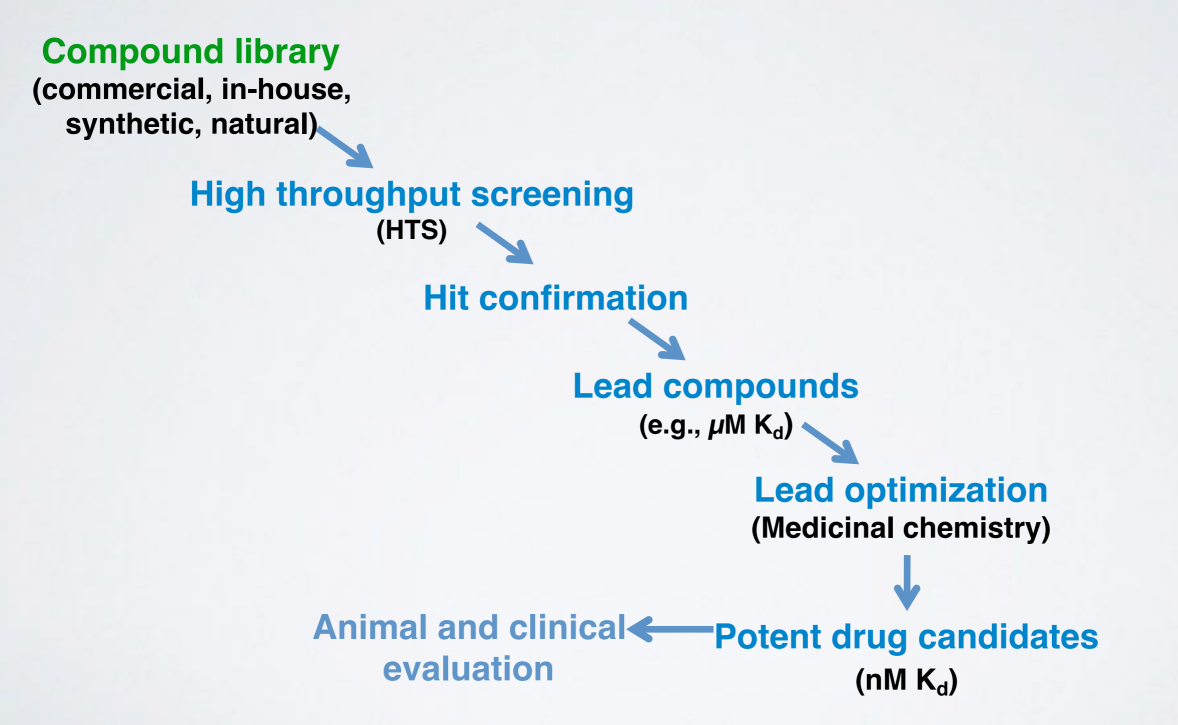
A model is **never perfect**. A model that is not quantitatively accurate in every respect does not preclude one from establishing results relevant to our understanding of biomolecules as long as the biophysics of the model are properly understood and explored.

 Calibration of the parameters is an ongoing and 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.

# THE TRADITIONAL EMPIRICAL PATH TO DRUG DISCOVERY



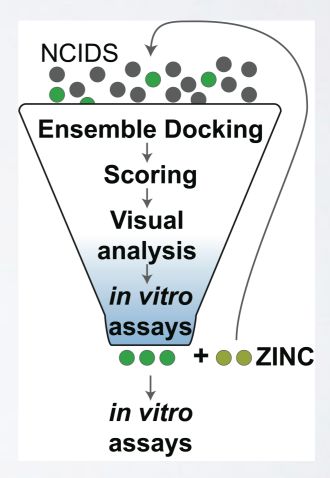
## COMPUTER-AIDED LIGAND DESIGN

Aims to reduce number of compounds synthesized and assayed

Lower costs

Reduce chemical waste

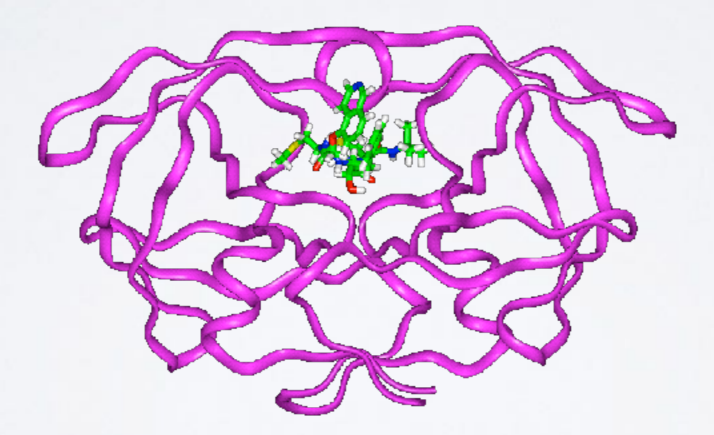
Facilitate faster progress



Two main approaches: (1). Receptor/Target-Based (2). Ligand/Drug-Based 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

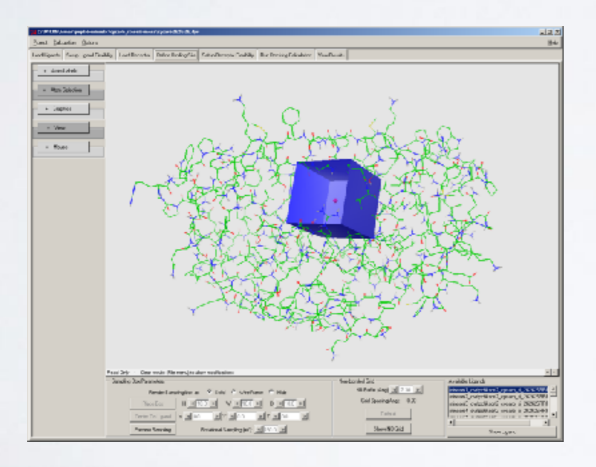


HIV Protease/KNI-272 complex

### PROTEIN-LIGAND DOCKING

#### Structure-Based Ligand Design

Docking software Search for structure of lowest energy

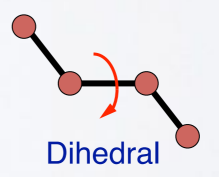


Potential function Energy as function of structure

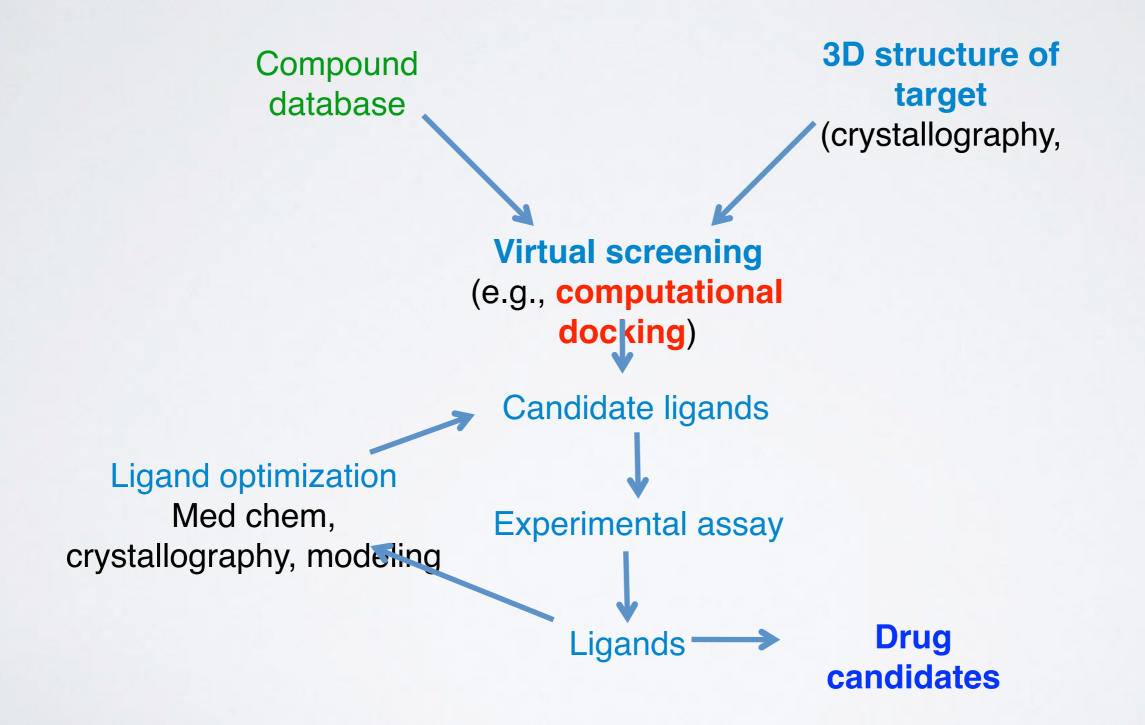




**Screened Coulombic** 



### STRUCTURE-BASED VIRTUAL SCREENING



# COMPOUND LIBRARIES



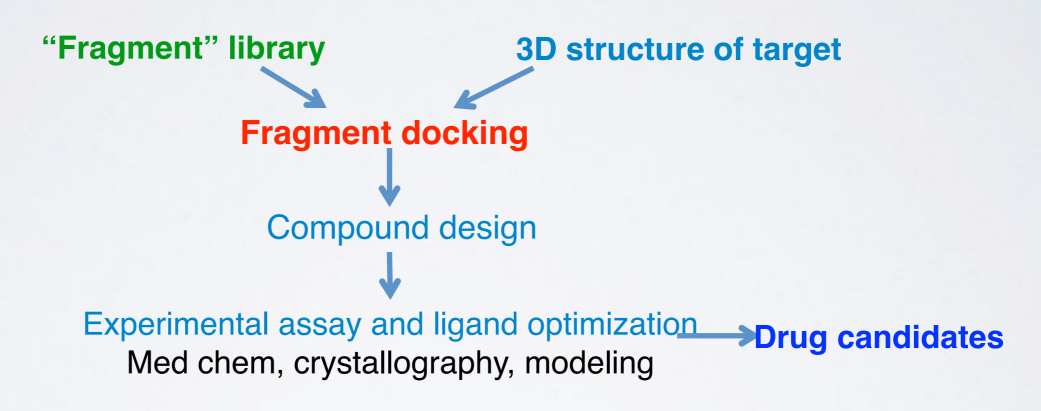
#### **Commercial** (in-house pharma)

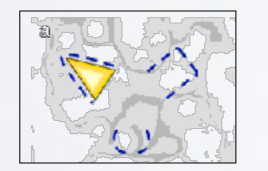
#### Government (NIH)

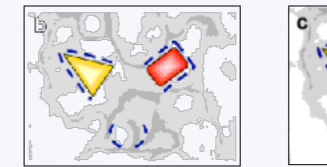
#### Academia

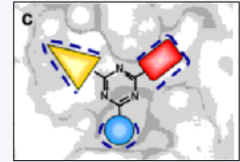
LINE USERSI MUS

# FRAGMENTAL STRUCTURE-BASED SCREENING





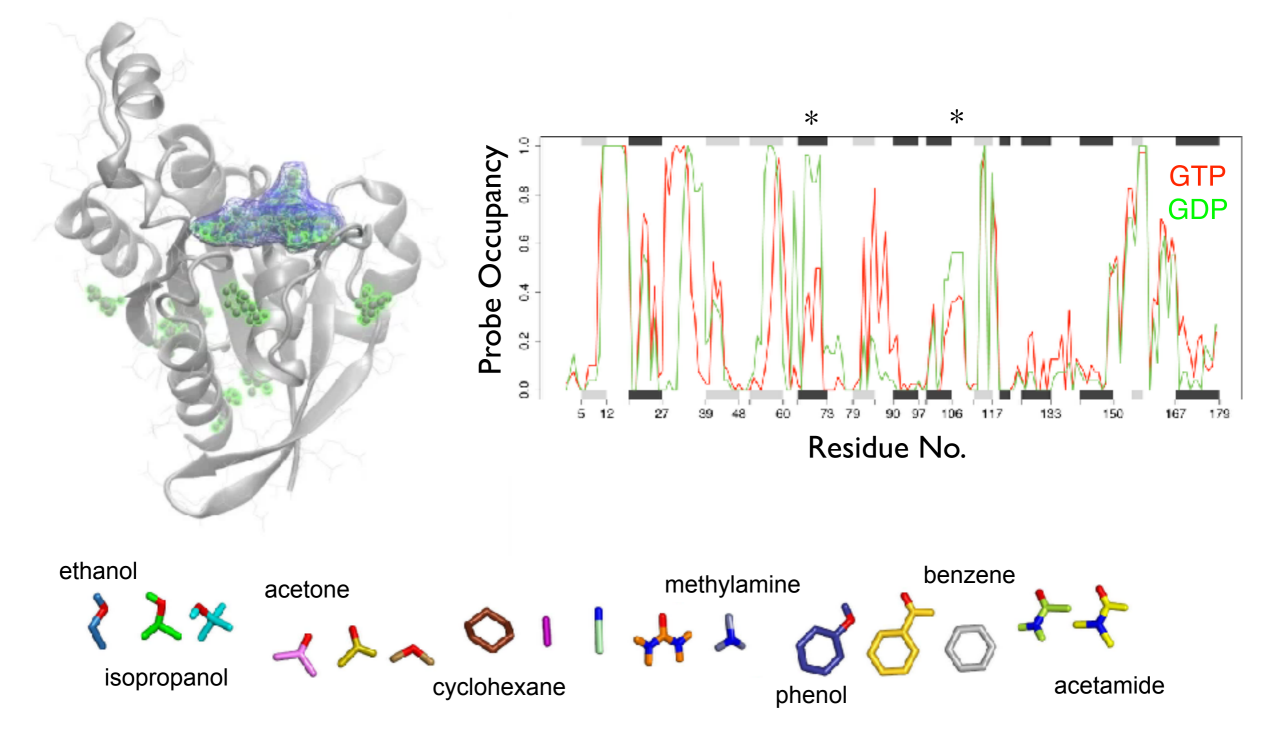




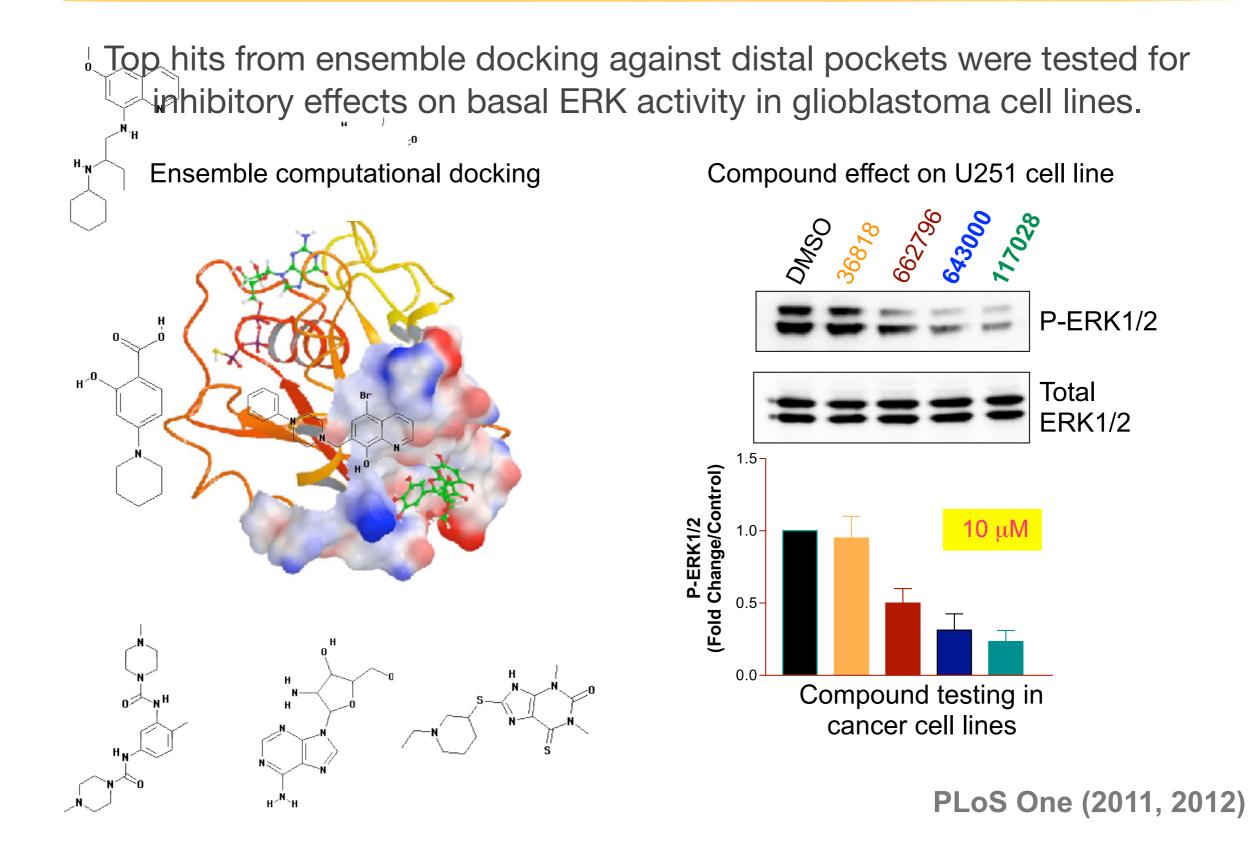
http://www.beilstein-institut.de/bozen2002/proceedings/Jhoti/jhoti.html

#### Multiple non active-site pockets identified

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

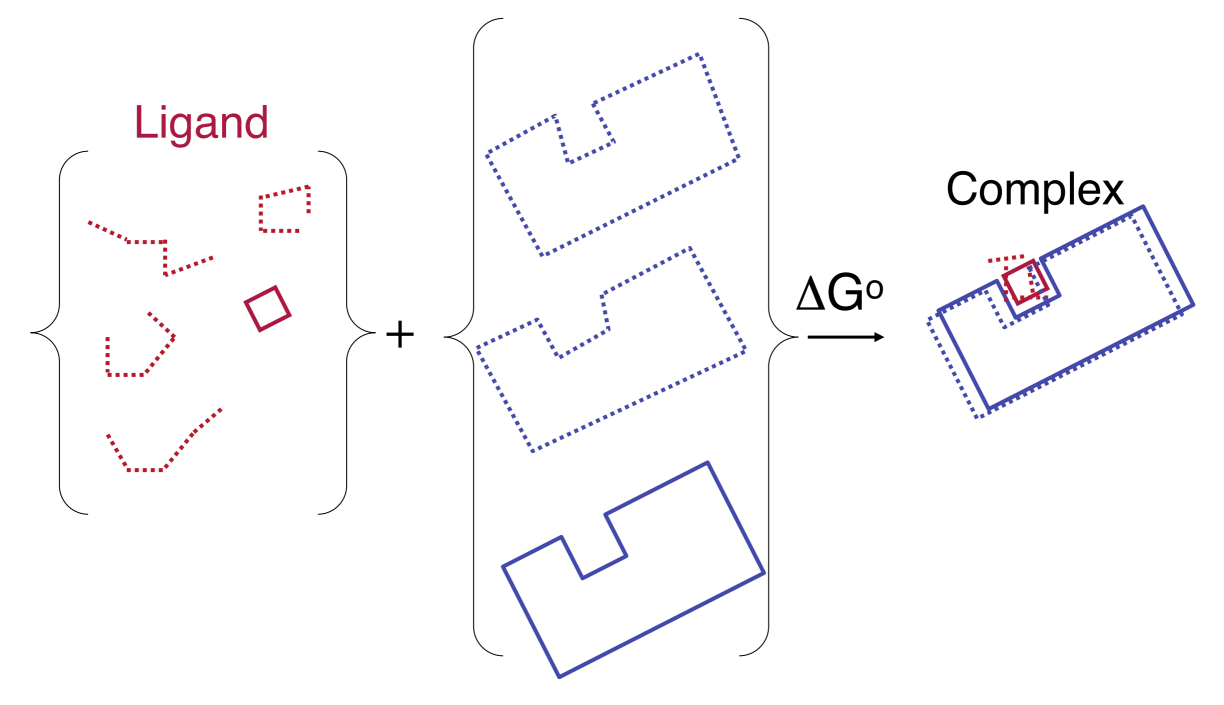


### Ensemble docking & candidate inhibitor testing



### Proteins and Ligand are Flexible

Protein



## 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

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

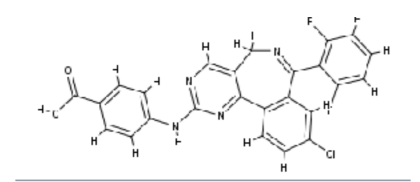
A company wants to work around another company's chemical patents

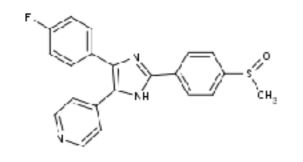
A high-affinity ligand is toxic, is not well-absorbed, etc.

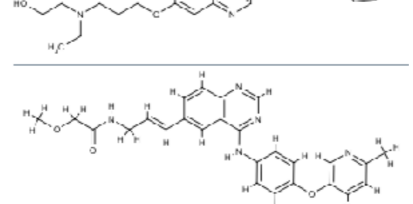
### **Scenario 2**

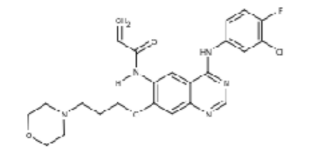
### Structure of Targeted Protein Unknown: Ligand-Based Drug Discovery

e.g. MAP Kinase Inhibitors



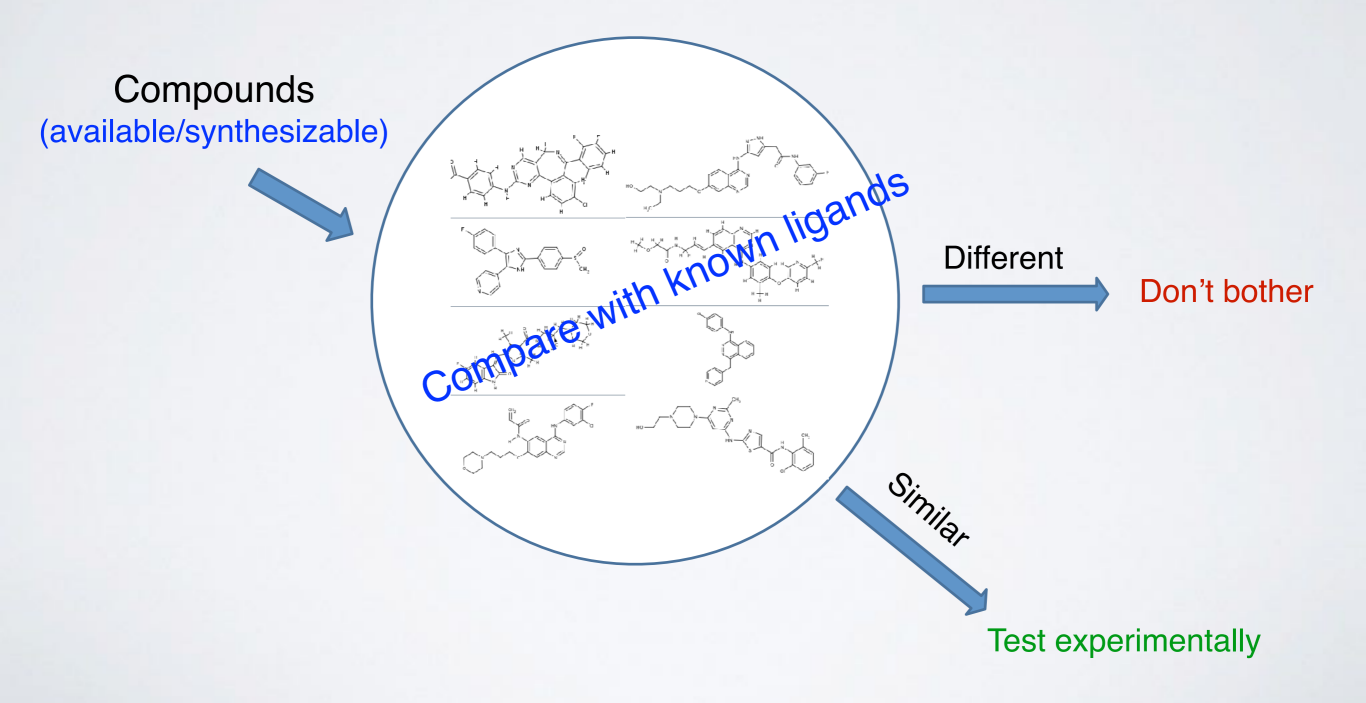






Using knowledge of existing inhibitors to discover more

## CHEMICAL SIMILARITY LIGAND-BASED DRUG-DISCOVERY



# CHEMICAL FINGERPRINTS BINARY STRUCTURE KEYS



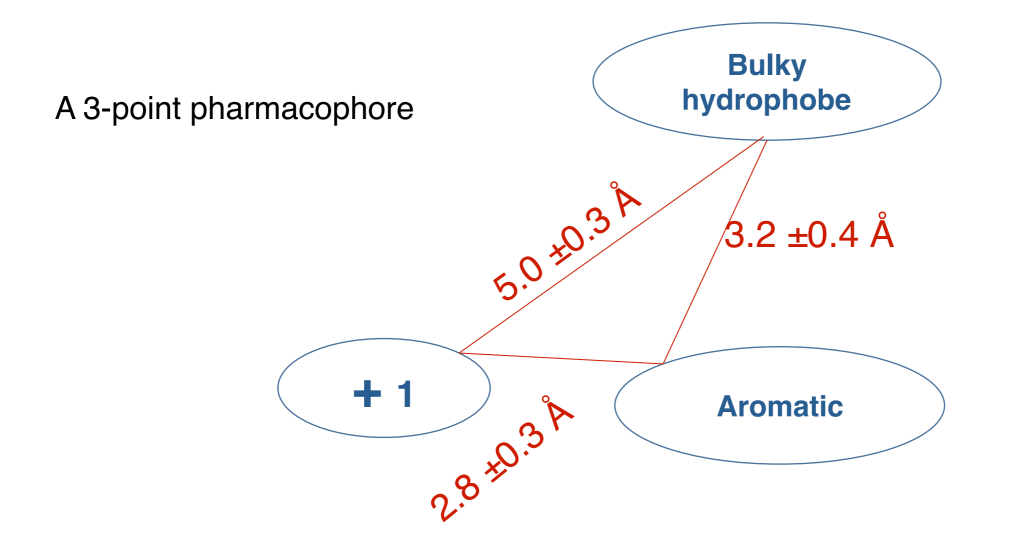
### CHEMICAL SIMILARITY FROM FINGERPRINTS



Tanimoto Similarity (or Jaccard Index), T  $T \equiv \frac{N_I}{N_U} = 0.25$ 



#### Pharmacophore Models Φάρμακο (drug) + Φορά (carry)

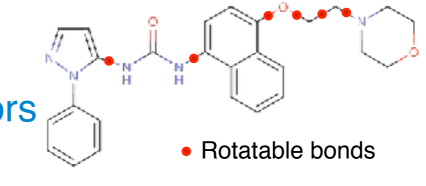


### **Molecular Descriptors**

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)

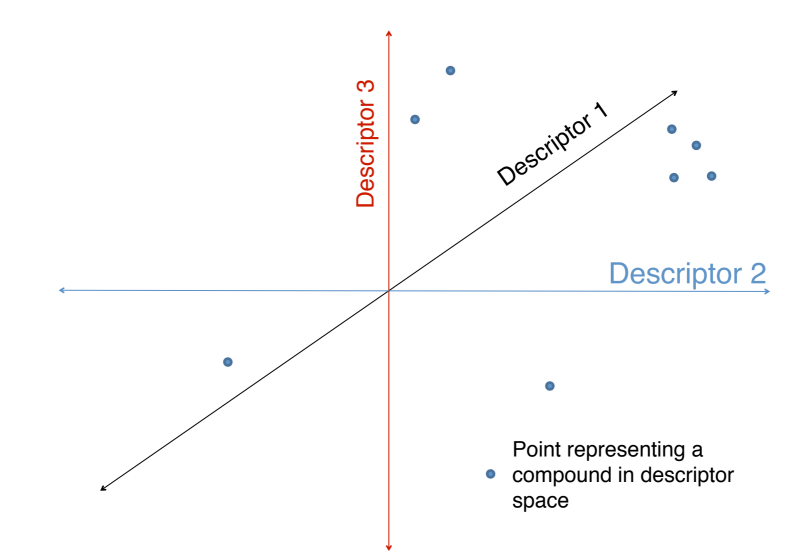


### Topological

branching index measures of linearity vs interconnectedness

#### Etc. etc.





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

### **CAUTIONARY NOTES**

 "Everything should be made as simple as it can be but not simpler"

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.

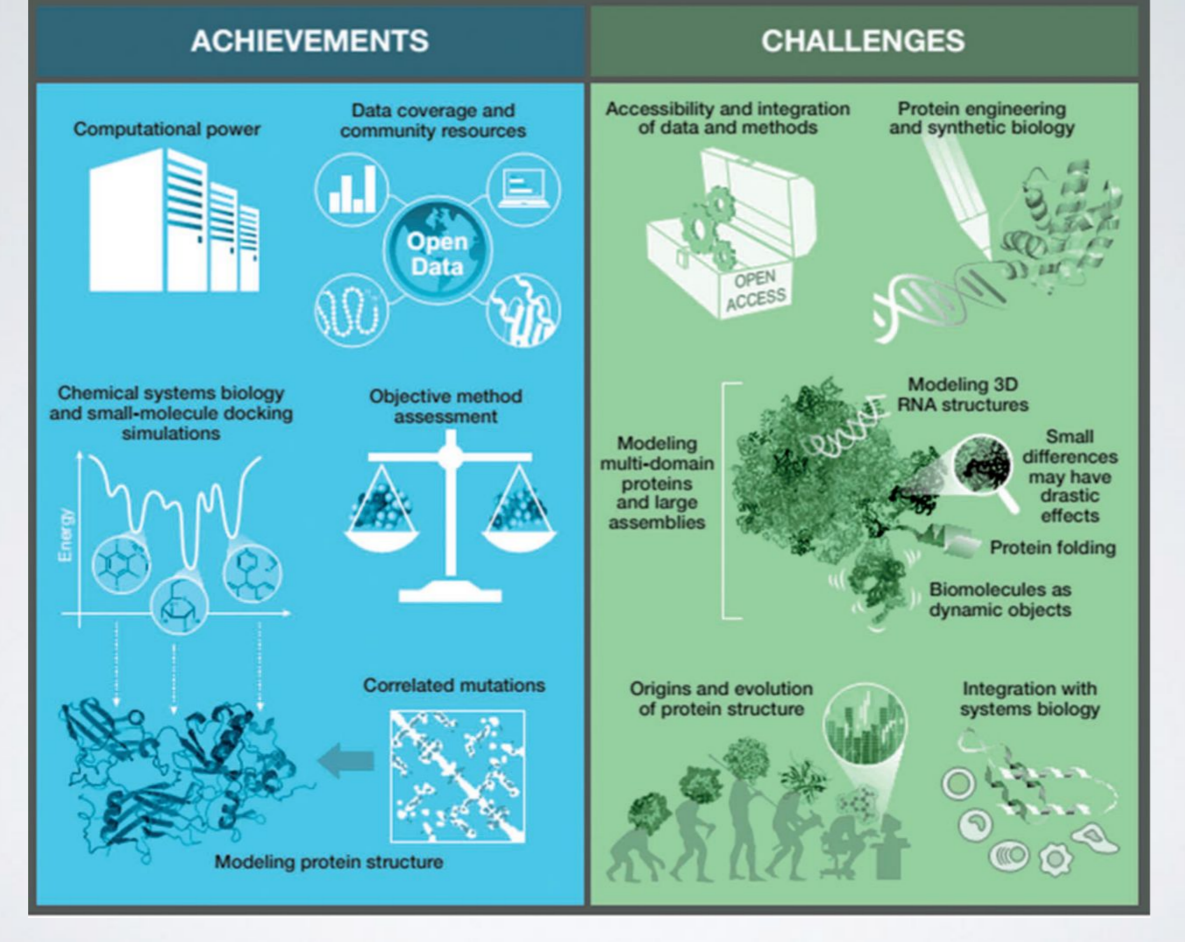
 Calibration of the parameters is an ongoing and imperfect process

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.

# 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



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

# INFORMING SYSTEMS BIOLOGY?

