#### **BINF 731**

# **Protein Structure Analysis**

Iosif Vaisman

2015

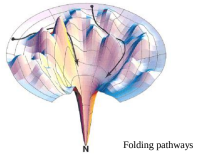
# Protein Modeling Methods

- Ab initio methods:
  - solution of a protein folding problem search in conformational space
- Energy-based methods: energy minimization molecular simulation

fold recogniion

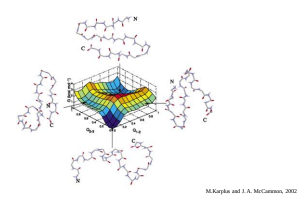
Knowledge-based methods:
 homology modeling

## **HP Lattice Models**

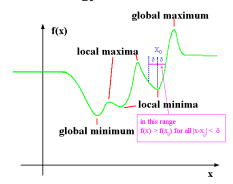


Chan & Dill, 1998

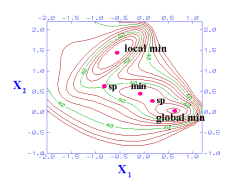
## Free energy surface in protein simulation



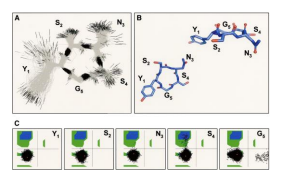
# **Energy Minimazation**



# **Energy Minimazation**

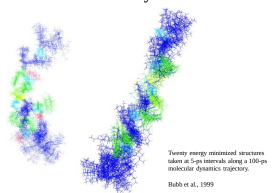


# Molecular dynamics



YSNSG cyclopeptide as observed along the 20 ns molecular dynamics trajectory (Thevenard et al., 2006)

# Molecular dynamics



# Molecular Dynamics

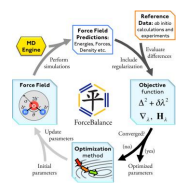
- Model system
- Initial conditions
- Boundary conditions
- Integration algorithm
- Constraints
- Ensemble
- Results

### Potential Energy Function and Force Field

$$egin{aligned} V(ec{R}) &= \sum_{ ext{bonds}} K_{ ext{d}} (d-d_0)^2 + \sum_{ ext{Urey-Bradley}} K_{ ext{UB}} (S-S_0)^2 + \ \sum_{ ext{angle}} K_{ heta} ( heta- heta_0)^2 + \sum_{ ext{dihedrals}} K_{\chi} (1+\cos(n\chi-\delta)) + \ \sum_{ ext{impropers}} K_{\phi} (\phi-\phi_0)^2 + \ \sum_{ ext{nonbond}} \left\{ \in_{ij} \left[ \left( rac{R_{ij}^{ ext{min}}}{r_{ij}} 
ight)^{12} - \left( rac{R_{ij}^{ ext{min}}}{r_{ij}} 
ight)^6 \right] + rac{q_i q_j}{ec{ec{e}_i r_{ij}}} 
ight\} \end{aligned}$$

M.Karplus and J. A. McCammon, 2002

## Force Field Development and Parametrization



L.-P. Wang et al., 2014

# Molecular Dynamics

$$F_{i}=m_{i} a_{i}$$

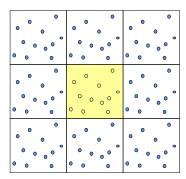
$$a_{i} = dv_{i} / dt$$

$$v_{i} = dr_{i} / dt$$

$$- dE / dr_{i} = F_{i}$$

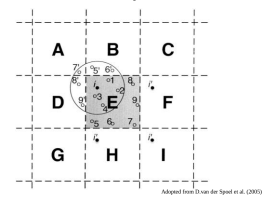
$$- dE / dr_{i} = m_{i} d^{2}r_{i} / dt^{2}$$

# **Periodic Boundary Conditions**

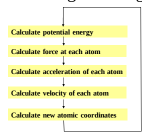


J. Jensen, animation

# **Periodic Boundary Conditions**



## MD cycle and integration algorithm



### Characteristic Time Scales for Protein Motions

| event  | spatial<br>extent (nm) | amplitude<br>(nm) | time (s)                          | appropriate simulations                          |
|--|------------------------|-------------------|-----------------------------------|--|
| bond-length vibration                                  | 0.2-0.5                | 0.001-0.01        | 10-14-10-13                       | QM methods                                       |
| elastic vibration of globular domain                   | 1.0-2.0                | 0.005-0.05        | 10-12-10-11                       | conventional MD                                  |
| rotation of solvent-exposed side chains                | 0.5-1.0                | 0.5-1.0           | 10-11-10-10                       | conventional MD                                  |
| torsional libration of buried groups                   | 0.5-1.0                | 0.05              | 10-11-10-9                        | conventional MD                                  |
| hinge bending (relative<br>motion of globular domains) | 1.0-2.0                | 0.1-0.5           | 10-11-10-7                        | Langevin dynamics, enhanced sampling MD methods? |
| rotation of buried side chains                         | 0.5                    | 0.5               | 10-4-1                            | enhanced sampling MD methods?                    |
| allosteric transitions                                 | 0.5-4.0                | 0.1-0.5           | 10-5-1                            | enhanced sampling MD methods?                    |
| local denaturation                                     | 0.5-1.0                | 0.5-1.0           | 10 <sup>-5</sup> -10 <sup>1</sup> | enhanced sampling MD methods?                    |
| loop motions   | 1.0-5.0                | 1.0-5.0           | 10-9-10-5                         | Brownian dynamics?                               |
| rigid-body (helix) motions                             |                        | 1.0-5.0           | 10-9-10-6                         | enhanced sampling MD methods?                    |
| helix-coil transitions                                 |                        | >5.0              | 10 <sup>-7</sup> -10 <sup>4</sup> | enhanced sampling MD methods?                    |
| protein association                                    | ≫1.0                   |                   |                                   | Brownian dynamics                                |

S. A. Adcock and J. A. McCammon, 2006

# MD Ensemble

#### $Microcanonical\ ensemble\ (NVE):$

The thermodynamic state characterized by a fixed number of atoms, N, a fixed volume, V, and a fixed energy, E. This corresponds to an isolated system.

#### Canonical Ensemble (NVT):

This is a collection of all systems whose thermodynamic state is characterized by a fixed number of atoms, N, a fixed volume, V, and a fixed temperature, T.

#### **Isobaric-Isothermal Ensemble (NPT):**

This ensemble is characterized by a fixed number of atoms, N, a fixed pressure, P, and a fixed temperature, T.

#### Grand canonical Ensemble ( $\mu VT$ ):

The thermodynamic state for this ensemble is characterized by a fixed chemical potential, m, a fixed volume, V, and a fixed temperature, T.

# Temperature in molecular dynamics

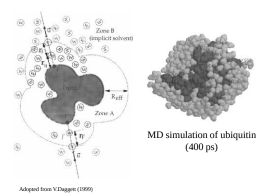
$$U_{kin} = \sum_{i=1}^{n} \frac{1}{2} m_i v_i^2 = \frac{3}{2} NkT$$

N – number of atoms

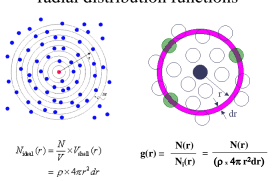
k – Boltzmann constant

T-absolute temperature

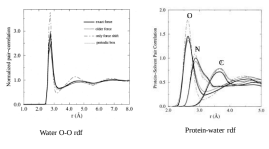
## MD of proteins: Solvent model



# MD of proteins: radial distribution functions

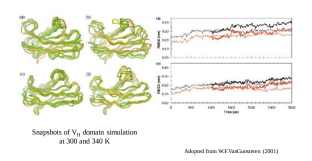


# MD of proteins: radial distribution functions

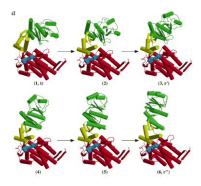


#### Adopted from V.Daggett (1999)

## MD of proteins: mobile regions

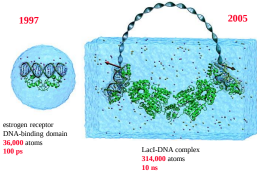


# MD of proteins: Conformational change



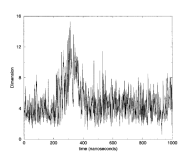
A set of structures on the reaction path showing the behavior of a single subunit (apical domain, green; intermediate domain, yellow; equatorial domain, red; and ATP, blue). Structures 1–3 correspond to the first stages associated with ATP binding; 4–6 correspond to the second stage involving GroES binding. The early downward motion of the intermediate domain (compare [1,1] with [2] and [6,7"]) is the trigger for the overall transition.

# MD of proteins: scale of simulation



Adopted from J.C.Phillips et al. (2005) M.Karplus and J. A. McCammon, 2002

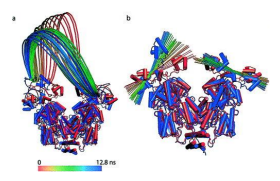
## MD of proteins: long runs



1 microsecond simulation of villin

Adopted from I.D.Kuntz and P.Kollman (2001)

# MD of proteins: long runs



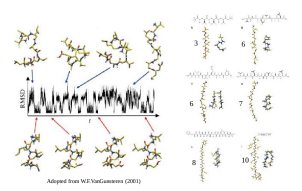
Adopted from J.C.Phillips et al. (2005)

# MD of proteins: performance

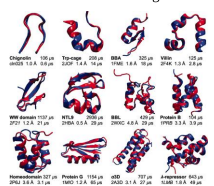
|      | Simulation | n setup |       |       |            | Performance, ps/day |        |        |        |
|------|------------|---------|-------|-------|------------|---------------------|--------|--------|--------|
| 66   | System     | FF      | virtH | Water | Coulomb    | LJ                  | ia32   | x86-64 | ppc    |
| 1999 | Vil        | G       | no    | TIP3P | cutoff 0.8 | cutoff 0.8          | 9744   | 9574   | 14,385 |
|      | Vil        | G       | yes   | TIP3P | cutoff 0.8 | cutoff 0.8          | 16,900 | 16,895 | 23,681 |
|      | Vil        | G       | yes   | TIP3P | RF 1.0     | cutoff 1.0          | 10,308 | 9719   | 12,934 |

|     | stem<br>DB ID) | Number<br>of atoms | Approximate performance (µs/machine-day) |
|-----|----------------|--------------------|--|
| *D  | HFR (5DFR)     | 23,558             | 17.4                                     |
| aSl | FP (1SFP)      | 48,423             | 11.7                                     |
| Fts | Z (1FSZ)       | 98,236             | 5.7                                      |
| T7  | Lig (1A01)     | 116,650            | 5.5                                      |
| bII | AP (1BPM)      | 132,362            | 4.8                                      |

# MD: Reversible folding of peptides

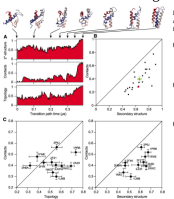


## MD: Reversible folding of small proteins



K. Lindorff-Larsen et al., 2011

# MD: Reversible folding of small proteins



Formation of topology, native contacts, and secondary structure during protein folding.

- (A) The three panels show the accumulation of native secondary structure, nonlocal native contacts, and native topology during a single folding event for α3D (B) The 24 transitions of α3D in a scatter plot are represented, with each of the black points corresponding to the time series integral for a single folding event event
  (C) Each point shows the average value
- Each point shows the average value over all folding and unfolding a vents observed for one protein. Each point is labeled with the PDB code of the relevant protein. Most proteins fall below the diagonal in these plots, showing that topology and secondary structure develop earlier than the full set of native contacts.