

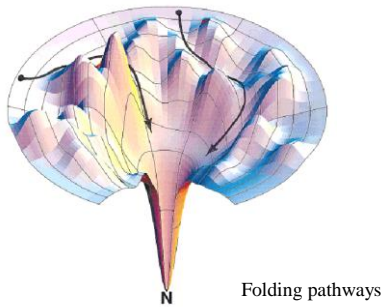
# Protein Structure Analysis

Iosif Vaisman

2015

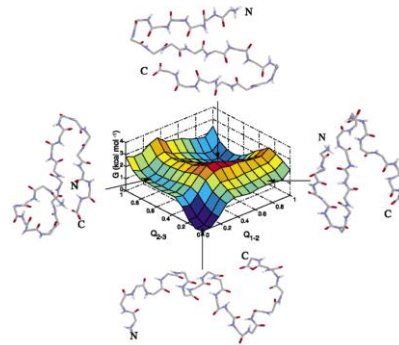
- **Ab initio methods:**  
solution of a protein folding problem  
search in conformational space
- **Energy-based methods:**  
energy minimization  
molecular simulation
- **Knowledge-based methods:**  
homology modeling  
fold recognition

## HP Lattice Models



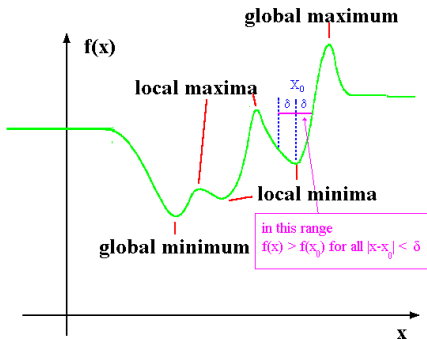
Chan & Dil, 1998

## Free energy surface in protein simulation

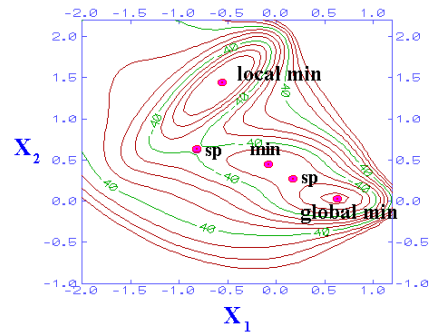


M.Karplus and J. A. McCammon, 2002

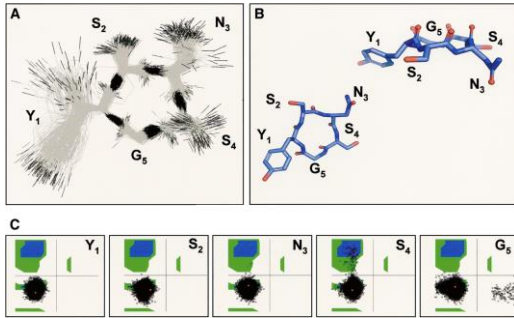
## Energy Minimization



## Energy Minimization

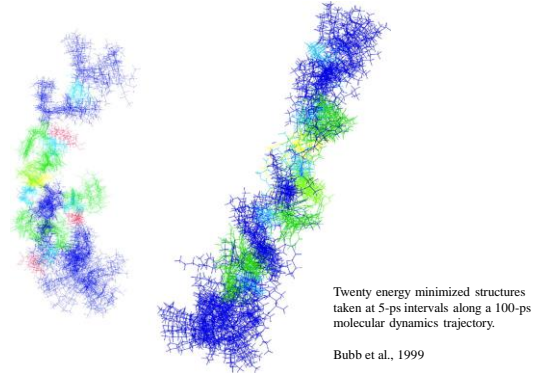


## Molecular dynamics



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

## Molecular dynamics



## Molecular Dynamics

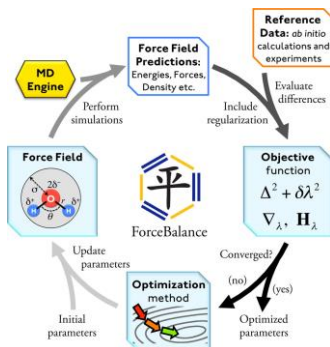
- Model system
- Initial conditions
- Boundary conditions
- Integration algorithm
- Constraints
- Ensemble
- Results

## Potential Energy Function and Force Field

$$V(\vec{R}) = \sum_{\text{bonds}} K_d (d - d_0)^2 + \sum_{\text{Urey-Bradley}} K_{UB} (S - S_0)^2 + \sum_{\text{angle}} K_\theta (\theta - \theta_0)^2 + \sum_{\text{dihedrals}} K_\chi (1 + \cos(n\chi - \delta)) + \sum_{\text{impropers}} K_\phi (\phi - \phi_0)^2 + \sum_{\text{nonbond}} \left\{ \epsilon_{ij} \left[ \left( \frac{R_{ij}^{\text{min}}}{r_{ij}} \right)^{12} - \left( \frac{R_{ij}^{\text{min}}}{r_{ij}} \right)^6 \right] + \frac{q_i q_j}{\epsilon_i r_{ij}} \right\}$$

M.Karplus and J. A. McCammon, 2002

## Force Field Development and Parametrization



L.-P. Wang et al., 2014

## Molecular Dynamics

$$F_i = m_i a_i$$

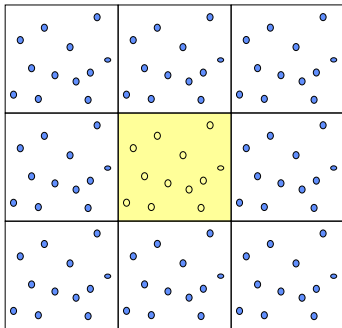
$$a_i = dv_i / dt$$

$$v_i = dx_i / dt$$

$$-dE / dx_i = F_i$$

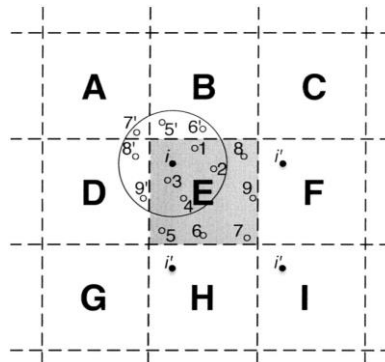
$$-dE / dx_i = m_i d^2x_i / dt^2$$

## Periodic Boundary Conditions



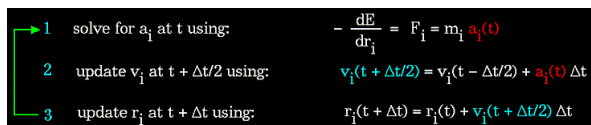
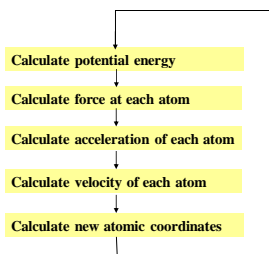
J. Jensen, [animation](#)

## Periodic Boundary Conditions



Adopted from D.van der Spoel et al. (2005)

## MD cycle and integration algorithm



## Characteristic Time Scales for Protein Motions

event	spatial extent (nm)	amplitude (nm)	time (s)	appropriate simulations
bond-length vibration	0.2-0.5	0.001-0.01	10 <sup>-14</sup> -10 <sup>-13</sup>	QM methods
elastic vibration of globular domain	1.0-2.0	0.005-0.05	10 <sup>-12</sup> -10 <sup>-11</sup>	conventional MD
rotation of solvent-exposed side chains	0.5-1.0	0.5-1.0	10 <sup>-11</sup> -10 <sup>-10</sup>	conventional MD
torsional libration of buried groups	0.5-1.0	0.05	10 <sup>-11</sup> -10 <sup>-9</sup>	conventional MD
hinge bending (relative motion of globular domains)	1.0-2.0	0.1-0.5	10 <sup>-11</sup> -10 <sup>-7</sup>	Langevin dynamics, enhanced sampling MD methods?
rotation of buried side chains	0.5	0.5	10 <sup>-4</sup> -1	enhanced sampling MD methods?
allosteric transitions	0.5-4.0	0.1-0.5	10 <sup>-5</sup> -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 <sup>-8</sup> -10 <sup>-5</sup>	Brownian dynamics?
rigid-body (helix) motions	1.0-5.0	1.0-5.0	10 <sup>-8</sup> -10 <sup>-6</sup>	enhanced sampling MD methods?
helix-coil transitions	>1.0	>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 ( $\mu$ V $T$ ):

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

## Temperature in molecular dynamics

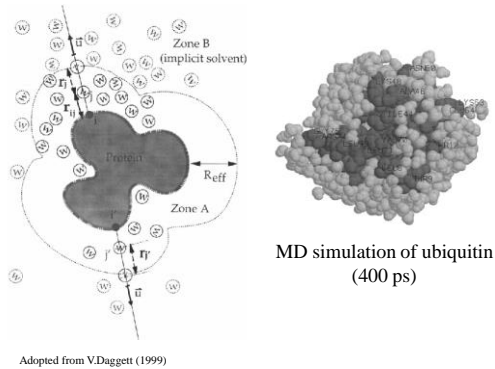
$$U_{kin} = \sum \frac{1}{2} m_i v_i^2 = \frac{3}{2} NkT$$

$N$  – number of atoms

$k$  – Boltzmann constant

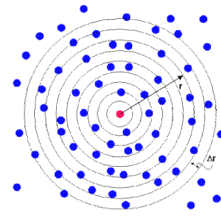
$T$  – absolute temperature

## MD of proteins: Solvent model

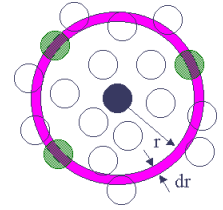


MD simulation of ubiquitin (400 ps)

## MD of proteins: radial distribution functions

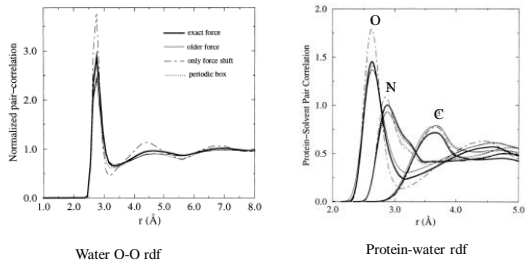


$$N_{\text{ideal}}(r) = \frac{N}{V} \times V_{\text{shell}}(r) = \rho \times 4\pi r^2 dr$$

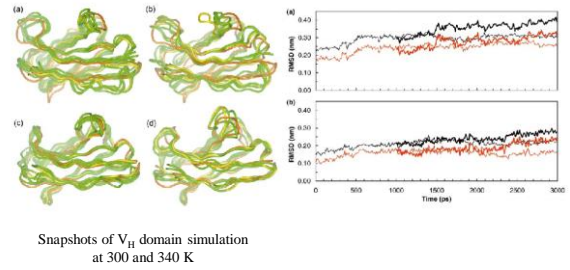


$$g(r) = \frac{N(r)}{N_i(r)} = \frac{N(r)}{\rho \times 4\pi r^2 dr}$$

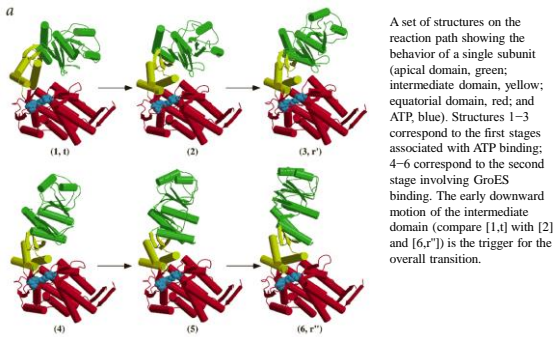
## MD of proteins: radial distribution functions



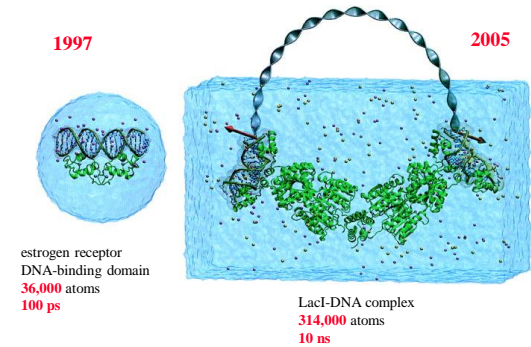
## MD of proteins: mobile regions



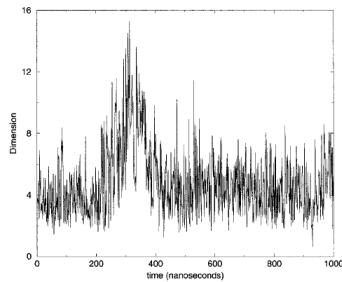
## MD of proteins: Conformational change



## MD of proteins: scale of simulation



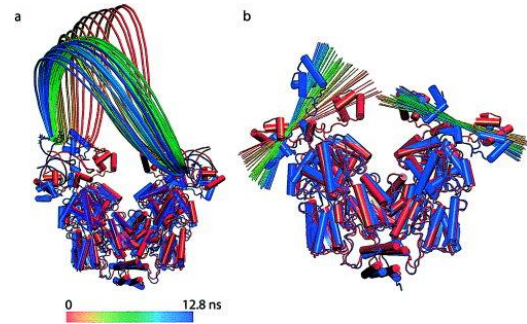
## MD of proteins: long runs



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

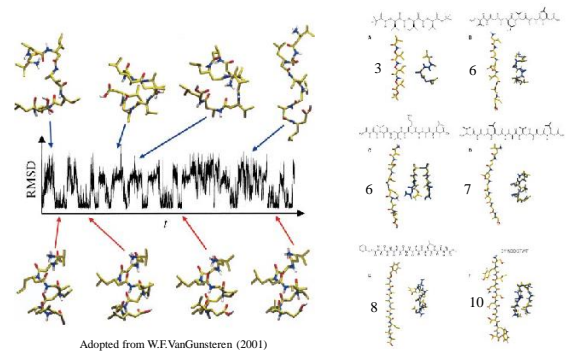
Simulation setup			Performance, ps/day					
System	FF	virtH	Water	Coulomb	LJ	ia32	x86-64	ppc
Vil	G	no	TIP3P	cutoff 0.8	cutoff 0.8	9744	9574	14,385
Vil	G	yes	TIP3P	cutoff 0.8	cutoff 0.8	16,900	16,895	23,681
Vil	G	yes	TIP3P	RF 1.0	cutoff 1.0	10,308	9719	12,934

1999

System (PDB ID)	Number of atoms	Approximate performance (ps/machine-day)
*DHFR (5DFR)	23,558	17.4
aSFP (1SFP)	48,423	11.7
FtsZ (1FSZ)	98,236	5.7
T7Lig (1A01)	116,650	5.5
bILAP (1BPM)	132,362	4.8

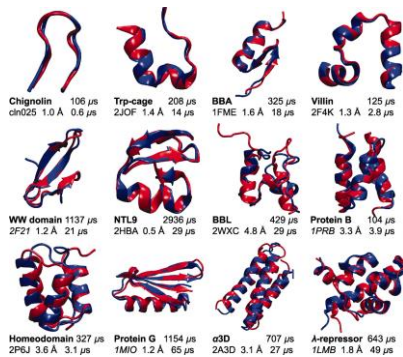
2014

## MD: Reversible folding of peptides



Adopted from W.E.VanGunsteren (2001)

## MD: Reversible folding of small proteins



K. Lindorff-Larsen et al., 2011

## MD: Reversible folding of small proteins

