

# Force Fields for Classical Molecular Dynamics simulations of Biomolecules

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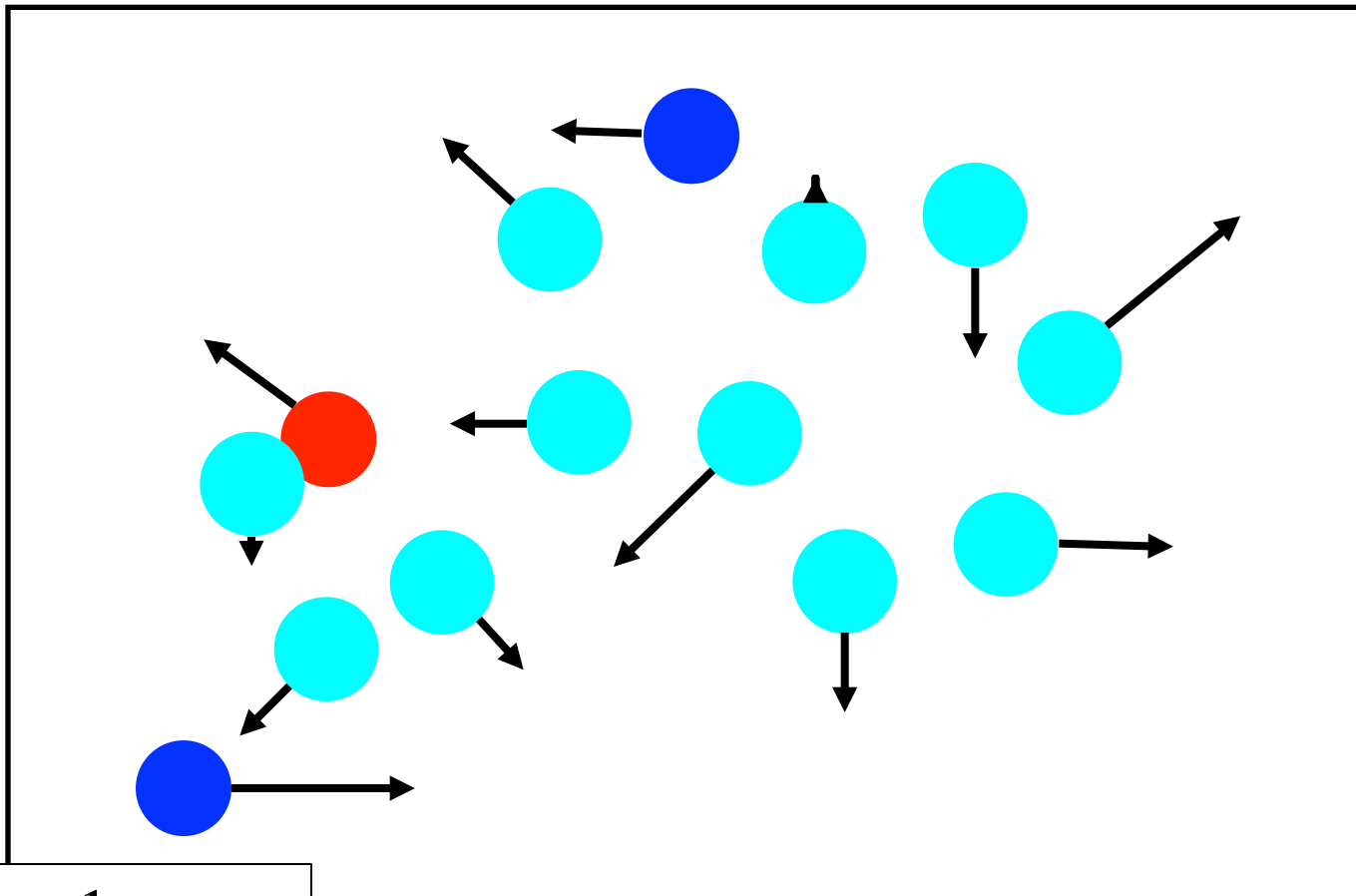
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# Classical Force Field Parameters

- Topology and structure files
- Parameter files
- Where do all the numbers needed by an MD code come from?
- Where to find these numbers and how to change them if needed.
- How to make topology files for ligands, cofactors, special amino acids, ...
- How to develop / put together missing parameters.

# Classical Molecular Dynamics

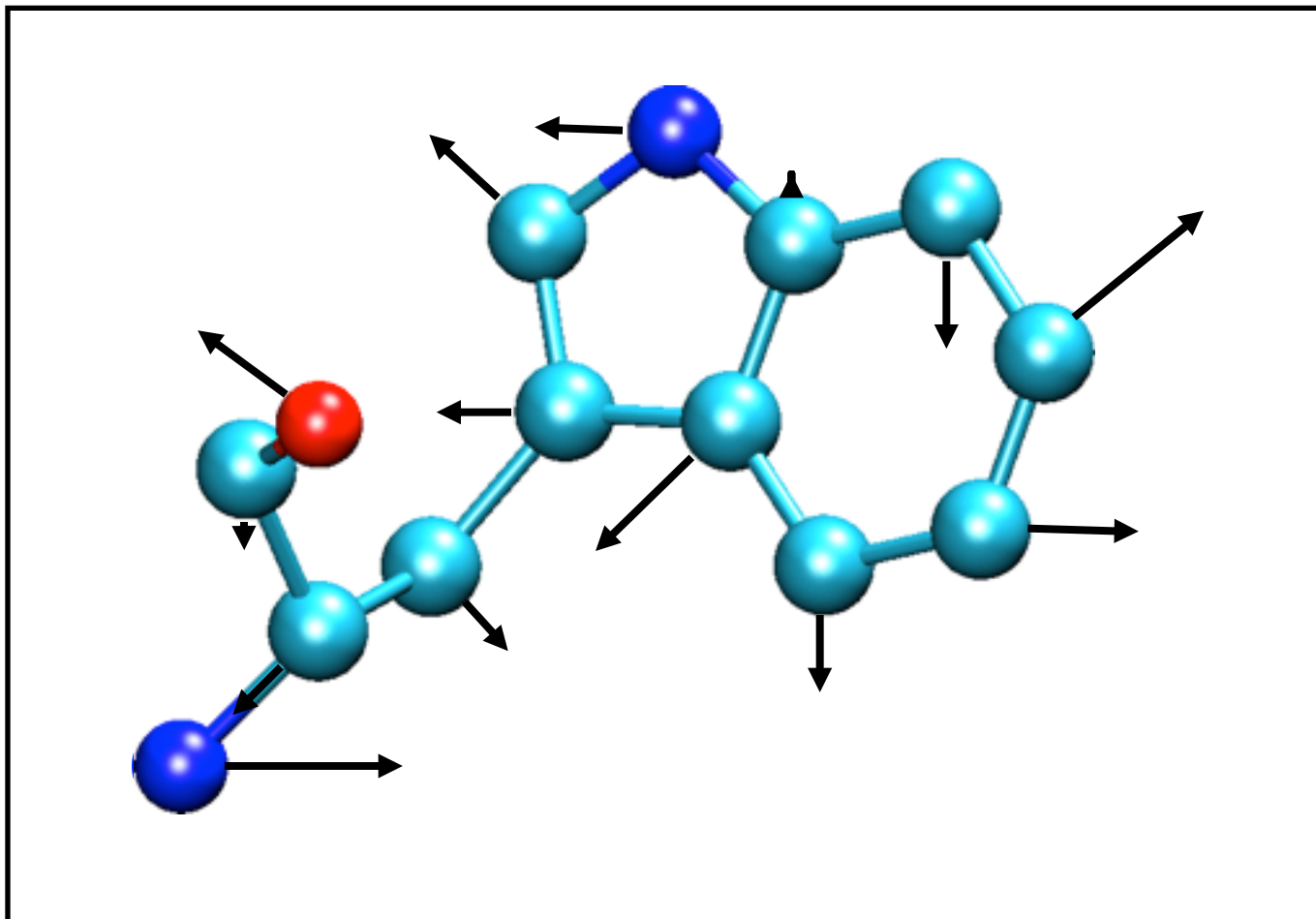


$$U(r) = \frac{1}{4\pi\epsilon_0} \frac{q_i q_j}{r_{ij}}$$

Coulomb interaction

$$U(r) = \epsilon_{ij} \left[ \left( \frac{R_{min,ij}}{r_{ij}} \right)^{12} - \left( \frac{R_{min,ij}}{r_{ij}} \right)^6 \right]$$

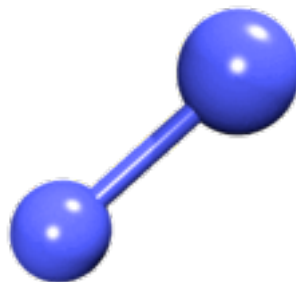
# Classical Molecular Dynamics



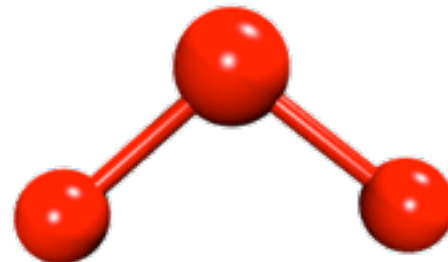
Bond definitions, atom types, atom names, parameters, ....

# Energy Terms Described in

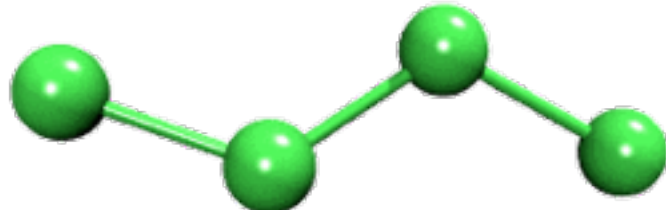
Bond



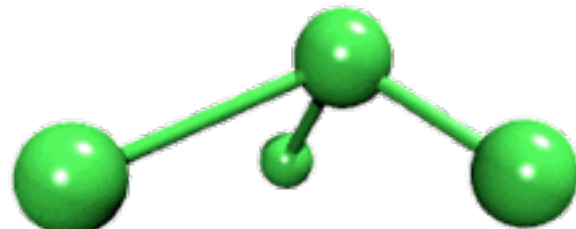
Angle



Dihedral



Improper



# The Potential Energy Function

$$U(\vec{R}) = \underbrace{\sum_{bonds} k_i^{bond} (r_i - r_0)^2}_{U_{bond}} + \underbrace{\sum_{angles} k_i^{angle} (\theta_i - \theta_0)^2}_{U_{angle}} + \underbrace{\sum_{dihedrals} k_i^{dih} [1 + \cos(n_i \phi_i + \delta_i)]}_{U_{dihedral}} + \underbrace{\sum_i \sum_{j \neq i} 4\epsilon_{ij} \left[ \left( \frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left( \frac{\sigma_{ij}}{r_{ij}} \right)^6 \right] + \sum_i \sum_{j \neq i} \frac{q_i q_j}{\epsilon r_{ij}}}_{U_{nonbond}}$$

$U_{bond}$  = oscillations about the equilibrium bond length

$U_{angle}$  = oscillations of 3 atoms about an equilibrium bond angle

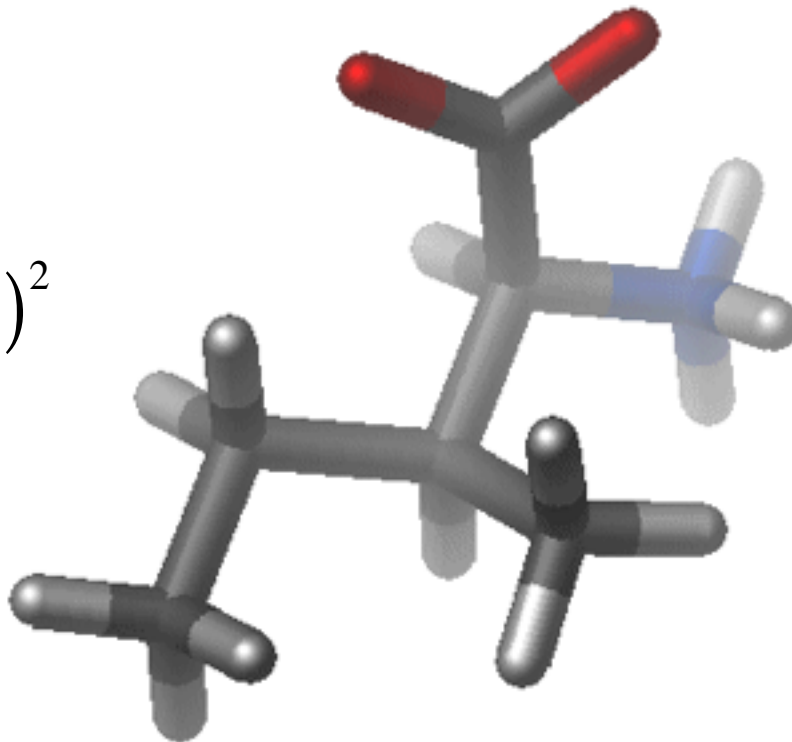
$U_{dihedral}$  = torsional rotation of 4 atoms about a central bond

$U_{nonbond}$  = non-bonded energy terms (electrostatics and Lenard-Jones)

# Interactions between bonded atoms

$$V_{angle} = K_\theta (\theta - \theta_o)^2$$

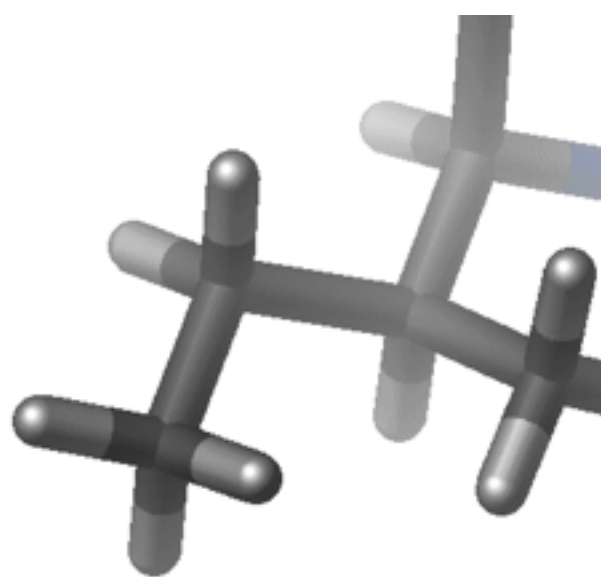
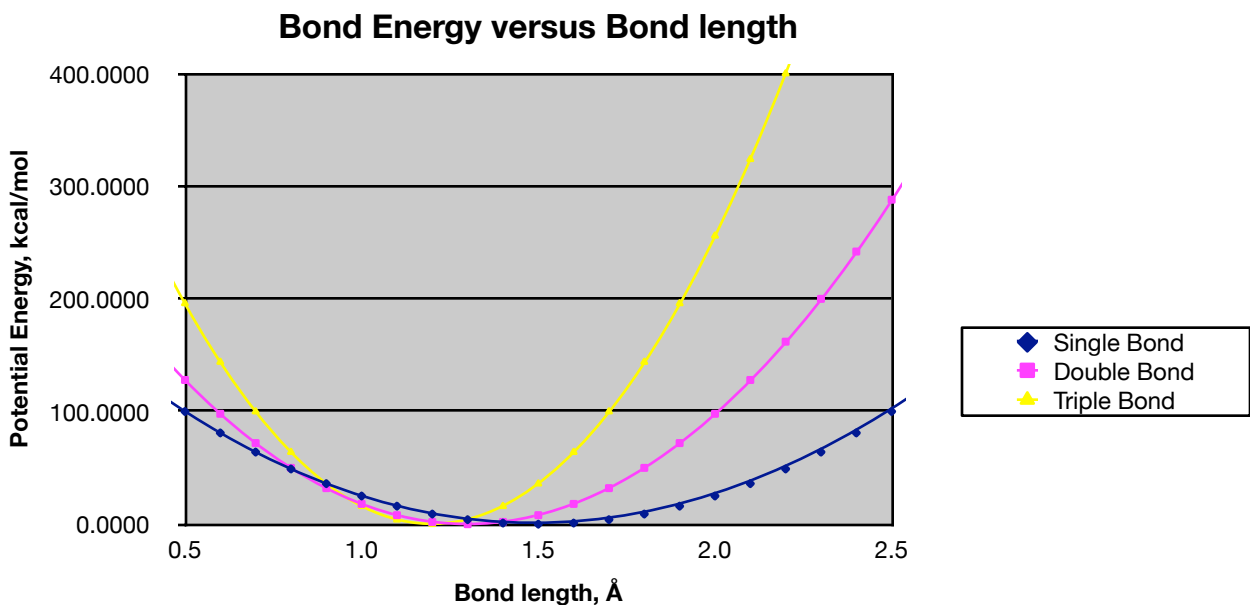
$$V_{bond} = K_b (b - b_o)^2$$



$$V_{dihedral} = K_\phi (1 + \cos(n\phi - \delta))$$

$$V_{bond} = K_b(b - b_o)^2$$

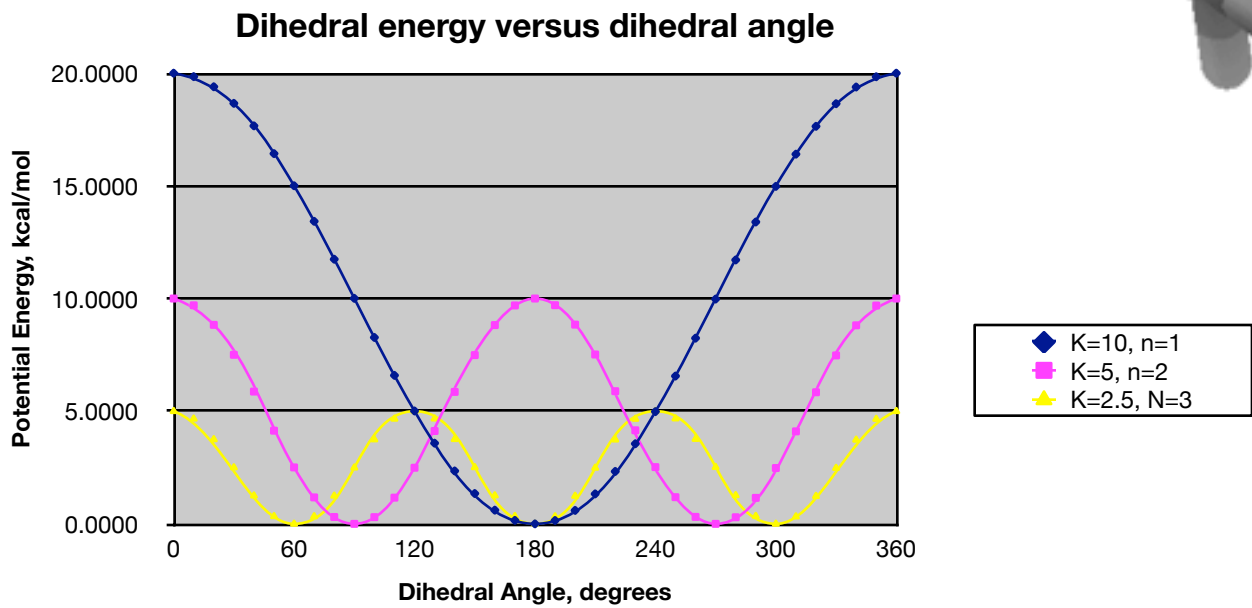
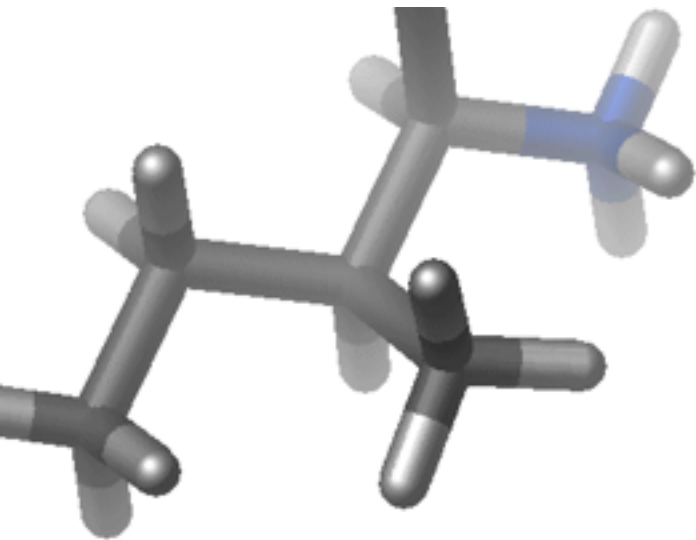
Chemical type	$K_{bond}$	$b_o$
C-C	100 kcal/mole/ $\text{\AA}^2$	1.5 $\text{\AA}$
C=C	200 kcal/mole/ $\text{\AA}^2$	1.3 $\text{\AA}$
C≡C	400 kcal/mole/ $\text{\AA}^2$	1.2 $\text{\AA}$



*Bond angles* and *improper* terms have similar quadratic forms, but with softer spring constants. The force constants can be obtained from vibrational analysis of the molecule (experimentally or theoretically).

# Dihedral Potential

$$V_{dihedral} = K_\phi (1 + \cos(n\phi - \delta))$$



$$\delta = 0^\circ$$

# Nonbonded Parameters

$$\sum_{\text{non-bonded}} \frac{q_i q_j}{4\pi D r_{ij}} + \epsilon_{ij} \left[ \left( \frac{R_{min,ij}}{r_{ij}} \right)^{12} - \left( \frac{R_{min,ij}}{r_{ij}} \right)^6 \right]$$

$q_i$ : partial atomic charge

$D$ : dielectric constant

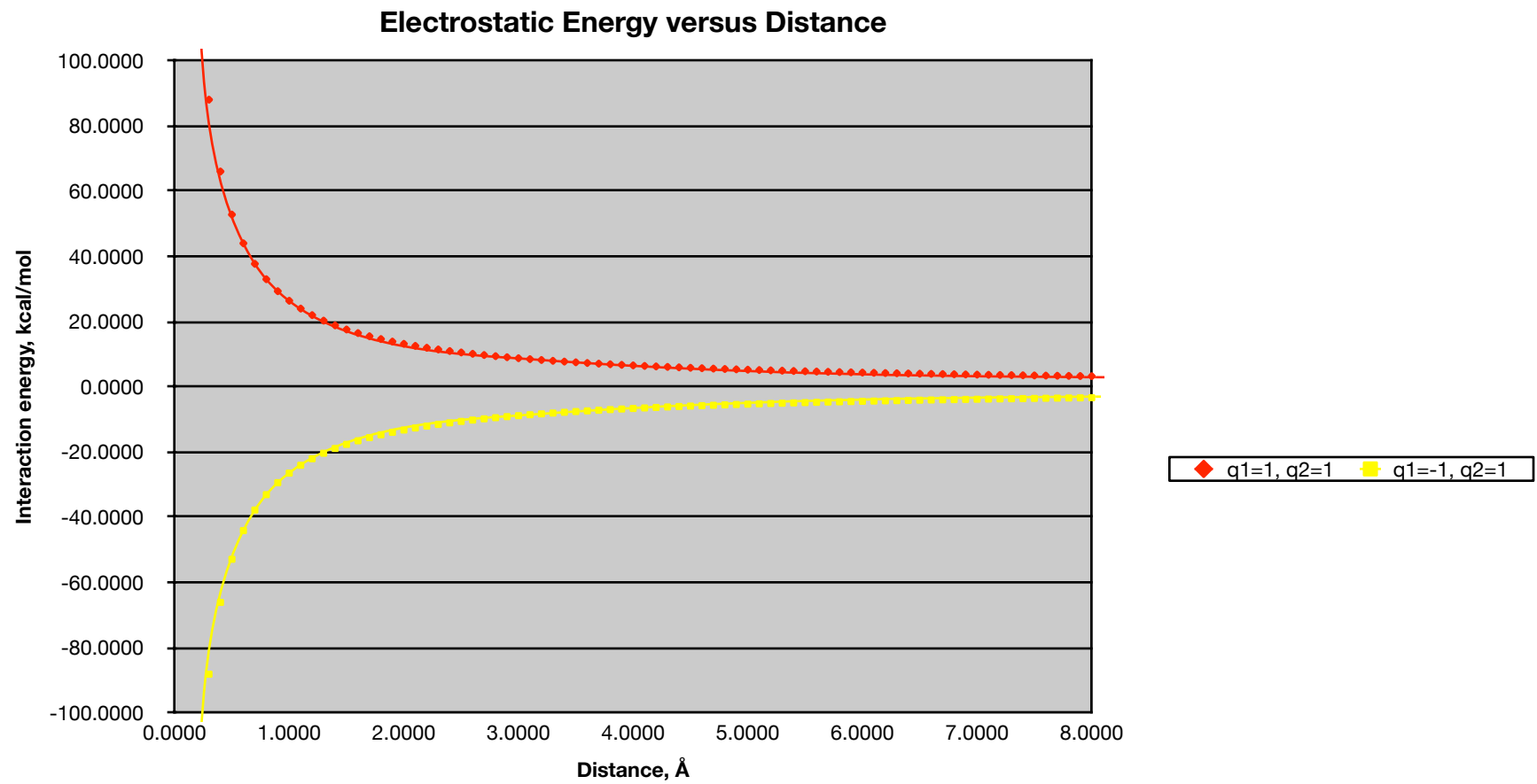
$\epsilon$ : Lennard-Jones (LJ, vdW) well-depth

$R_{min}$ : LJ radius ( $R_{min}/2$  in CHARMM)

Combining rules (CHARMM, Amber)

$$R_{min,i,j} = R_{min,i} + R_{min,j}$$

$$\epsilon_{i,j} = \text{SQRT}(\epsilon_i * \epsilon_j)$$



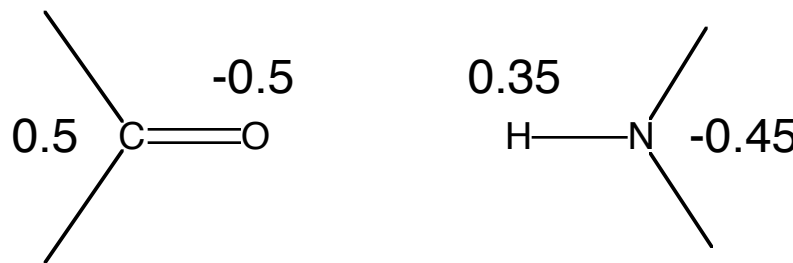
Note that the effect is long range.

# Charge Fitting Strategy

CHARMM- Mulliken\*

AMBER(ESP/RESP)

Partial atomic charges



\*Modifications based on interactions with TIP3 water

# CHARMM Potential Function

$$U(\vec{R}) = \underbrace{\sum_{bonds} k_i^{bond} (r_i - r_0)^2}_{U_{bond}} + \underbrace{\sum_{angles} k_i^{angle} (\theta_i - \theta_0)^2}_{U_{angle}} + \underbrace{\sum_{dihedrals} k_i^{dihed} [1 + \cos(n_i \phi_i + \delta_i)]}_{U_{dihedral}} + \underbrace{\sum_i \sum_{j \neq i} 4\epsilon_{ij} \left[ \left( \frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left( \frac{\sigma_{ij}}{r_{ij}} \right)^6 \right]}_{U_{nonbond}} + \sum_i \sum_{j \neq i} \frac{q_i q_j}{\epsilon r_{ij}}$$

The diagram illustrates the components of the CHARMM potential function. The potential energy  $U(\vec{R})$  is the sum of four terms: bond stretching, angle bending, dihedral rotation, and non-bonded interactions. Each term is labeled with its name in blue and has a corresponding red arrow pointing to it from a box labeled "parameters". The bond stretching term is labeled  $U_{bond}$ , the angle bending term is labeled  $U_{angle}$ , the dihedral rotation term is labeled  $U_{dihedral}$ , and the non-bonded interaction term is labeled  $U_{nonbond}$ . The boxes for "geometry" and "parameters" are highlighted in red, while the boxes for "PDB file", "Topology PSF file", and "Parameter file" are white.

# File Format/Structure

- The structure of a pdb file
- The structure of a psf file
- The topology file
- The parameter file
- Connection to potential energy terms

# Structure of a PDB file

index	resname		chain	resid	x	y	z	segname			
	name										
ATOM	22	N	ALA	B	3	-4.073	-7.587	-2.708	1.00	0.00	BH
ATOM	23	HN	ALA	B	3	-3.813	-6.675	-3.125	1.00	0.00	BH
ATOM	24	CA	ALA	B	3	-4.615	-7.557	-1.309	1.00	0.00	BH
ATOM	25	HA	ALA	B	3	-4.323	-8.453	-0.704	1.00	0.00	BH
ATOM	26	CB	ALA	B	3	-4.137	-6.277	-0.676	1.00	0.00	BH
ATOM	27	HB1	ALA	B	3	-3.128	-5.950	-0.907	1.00	0.00	BH
ATOM	28	HB2	ALA	B	3	-4.724	-5.439	-1.015	1.00	0.00	BH
ATOM	29	HB3	ALA	B	3	-4.360	-6.338	0.393	1.00	0.00	BH
ATOM	30	C	ALA	B	3	-6.187	-7.538	-1.357	1.00	0.00	BH
ATOM	31	O	ALA	B	3	-6.854	-6.553	-1.264	1.00	0.00	BH
ATOM	32	N	ALA	B	4	-6.697	-8.715	-1.643	1.00	0.00	BH
ATOM	33	HN	ALA	B	4	-6.023	-9.463	-1.751	1.00	0.00	BH
ATOM	34	CA	ALA	B	4	-8.105	-9.096	-1.934	1.00	0.00	BH
ATOM	35	HA	ALA	B	4	-8.287	-8.878	-3.003	1.00	0.00	BH
ATOM	36	CB	ALA	B	4	-8.214	-10.604	-1.704	1.00	0.00	BH
ATOM	37	HB1	ALA	B	4	-7.493	-11.205	-2.379	1.00	0.00	BH
ATOM	38	HB2	ALA	B	4	-8.016	-10.861	-0.665	1.00	0.00	BH
ATOM	39	HB3	ALA	B	4	-9.245	-10.914	-1.986	1.00	0.00	BH
ATOM	40	C	ALA	B	4	-9.226	-8.438	-1.091	1.00	0.00	BH
ATOM	41	O	ALA	B	4	-10.207	-7.958	-1.667	1.00	0.00	BH

>>> It is an ascii, fixed-format file <<

“No connectivity information”

# Looking at File Structures

- PDB file
- Topology file
- PSF file
- Parameter file

# Parameter Optimization Strategies

**Check if it has been parameterized by somebody else**

Literature

Google

**Minimal optimization**

By analogy (direct transfer of known parameters)

Quick, starting point

**Maximal optimization**

Time-consuming

Requires appropriate experimental and target data

**Choice based on goal of the calculations**

Minimal

database screening

NMR/X-ray structure determination

Maximal

free energy calculations, mechanistic studies,  
subtle environmental effects

# Getting Started

- Identify previously parameterized compounds
- Access topology information – assign atom types, connectivity, and charges – **annotate changes**

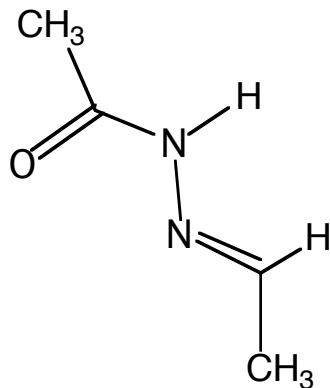
## CHARMM topology (parameter files)

top\_all22\_model.inp (par\_all22\_prot.inp)  
top\_all22\_prot.inp (par\_all22\_prot.inp)  
top\_all22\_sugar.inp (par\_all22\_sugar.inp)  
top\_all27\_lipid.rtf (par\_all27\_lipid.prm)  
top\_all27\_na.rtf (par\_all27\_na.prm)  
top\_all27\_na\_lipid.rtf (par\_all27\_na\_lipid.prm)  
top\_all27\_prot\_lipid.rtf (par\_all27\_prot\_lipid.prm)  
top\_all27\_prot\_na.rtf (par\_all27\_prot\_na.prm)  
toph19.inp (param19.inp)

NA and lipid force fields have new LJ parameters for the alkanes, representing increased optimization of the protein alkane parameters. Tests have shown that these are compatible (e.g. in protein-nucleic acid simulations). For new systems it is suggested that the new LJ parameters be used. Note that only the LJ parameters were changed; the internal parameters are identical.

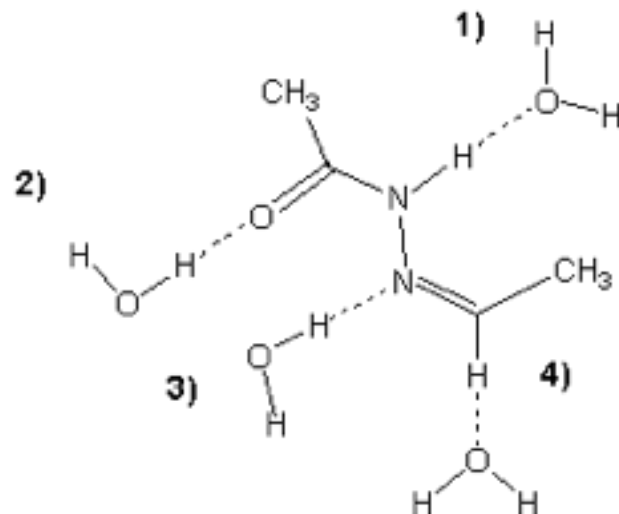
# Partial Charge Assignment

- Most important aspect for ligands
- Different force fields might take different philosophies
  - AMBER: RESP charges at the HF/6-31G level
    - Overestimation of dipole moments
    - Easier to set up
  - CHARMM: Interaction based optimization
    - TIP3P water representing the environment
    - Could be very difficult to set up
- Conformation dependence of partial charges
- Lack of polarization
- Try to be consistent within the force field
- pKa calculations for titratable residues



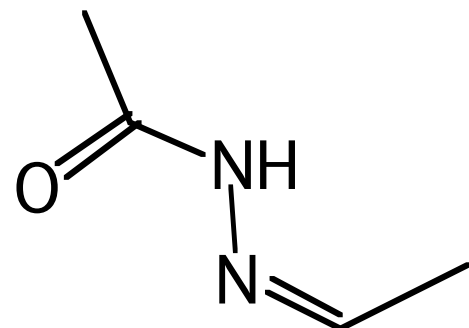
Starting charges??  
 Mulliken population analysis  
 Analogy comparison

Final charges (methyl, vary  $q_C$  to maintain integer charge,  $q_H = 0.09$ )  
 interactions with water (HF/6-31G\*, monohydrates!)

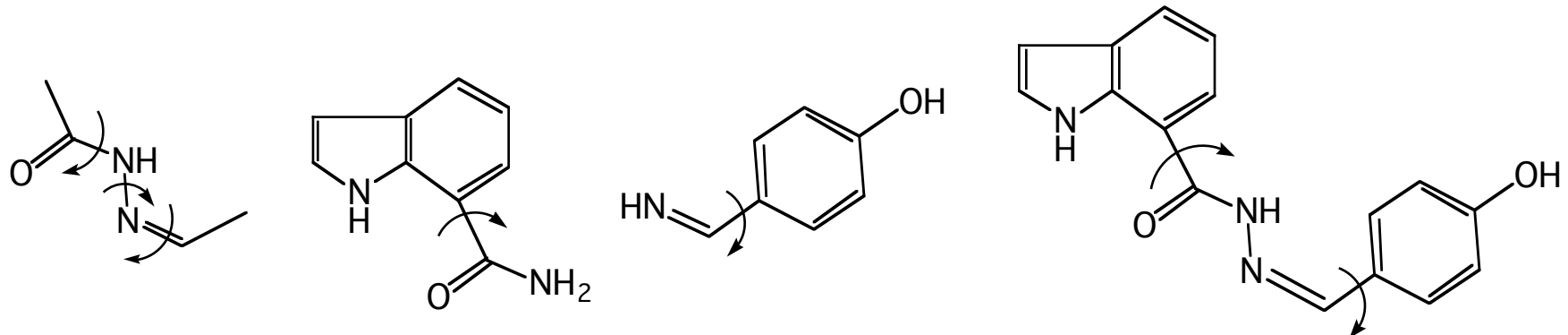
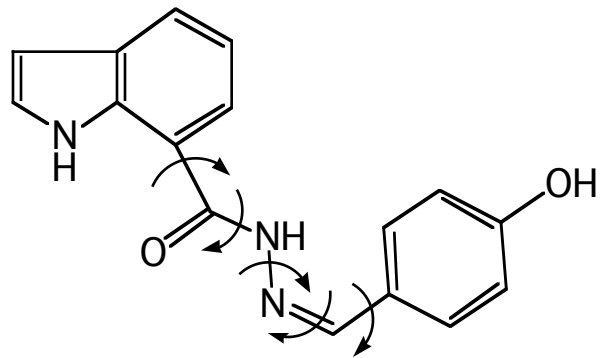


# Comparison of analogy and optimized charges

Name	Type	Analogy	Optimized
C1	CT3	-0.27	-0.27
H11	HA3	0.09	0.09
H12	HA3	0.09	0.09
H13	HA3	0.09	0.09
C2	C	0.51	0.58
O2	O	-0.51	-0.50
N3	NH1	-0.47	-0.32
H3	H	0.31	0.33
N4	NR1	0.16	-0.31
C5	CEL1	-0.15	-0.25
H51	HEL1	0.15	0.29
C6	CT3	-0.27	-0.09
H61	HA	0.09	0.09
H62	HA	0.09	0.09
H63	HA	0.09	0.09

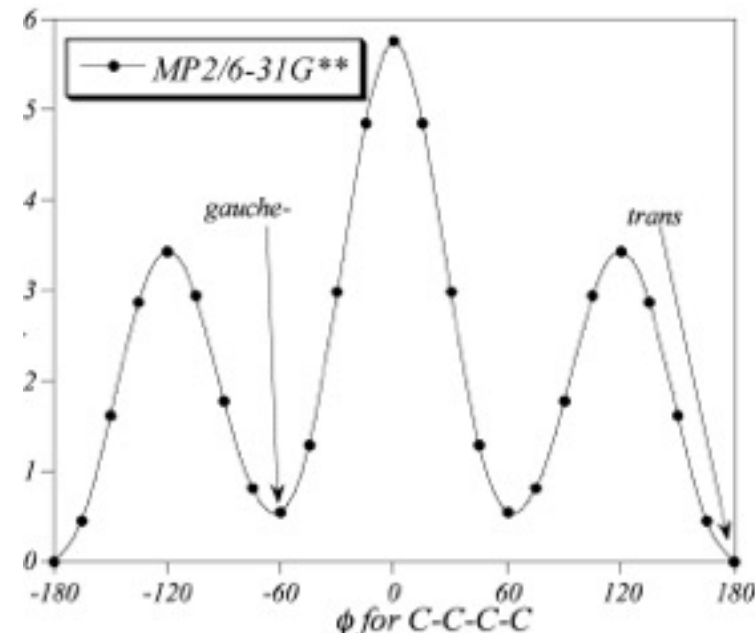
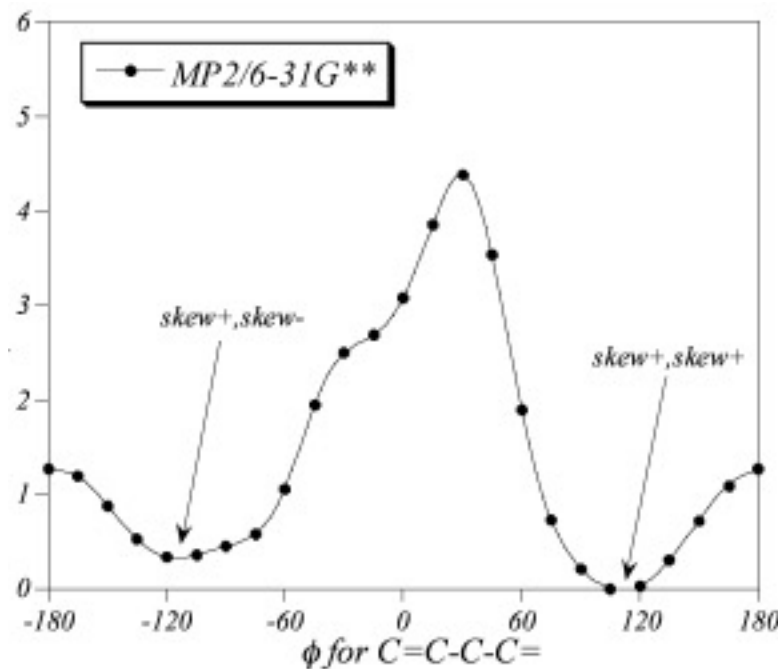
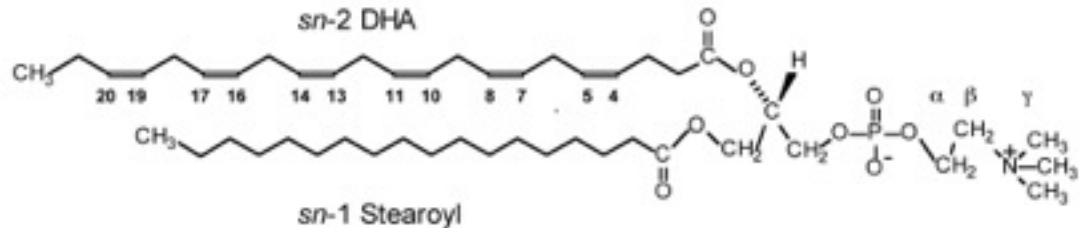
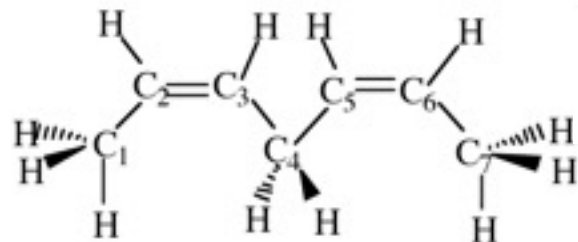


Dihedral optimization based on QM potential energy surfaces (HF/6-31G\* or MP2/6-31G\*).



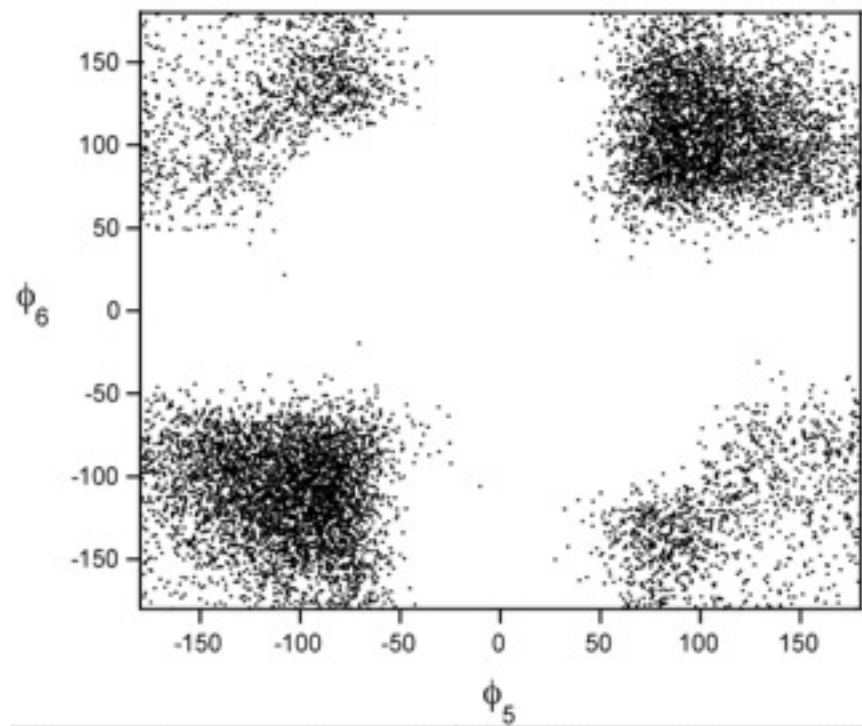
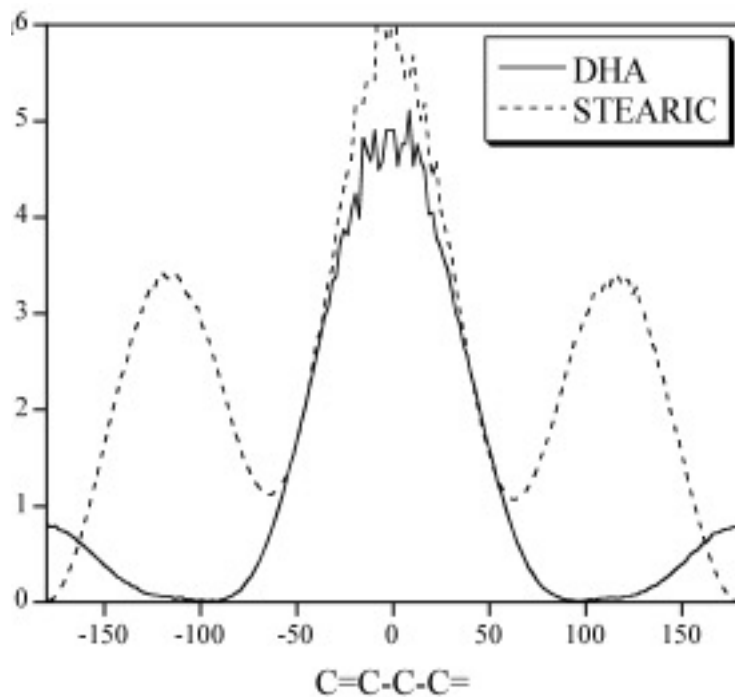
# Parameterization of unsaturated lipids

- All C=C bonds are cis, what does rotation about neighboring single bonds look like?



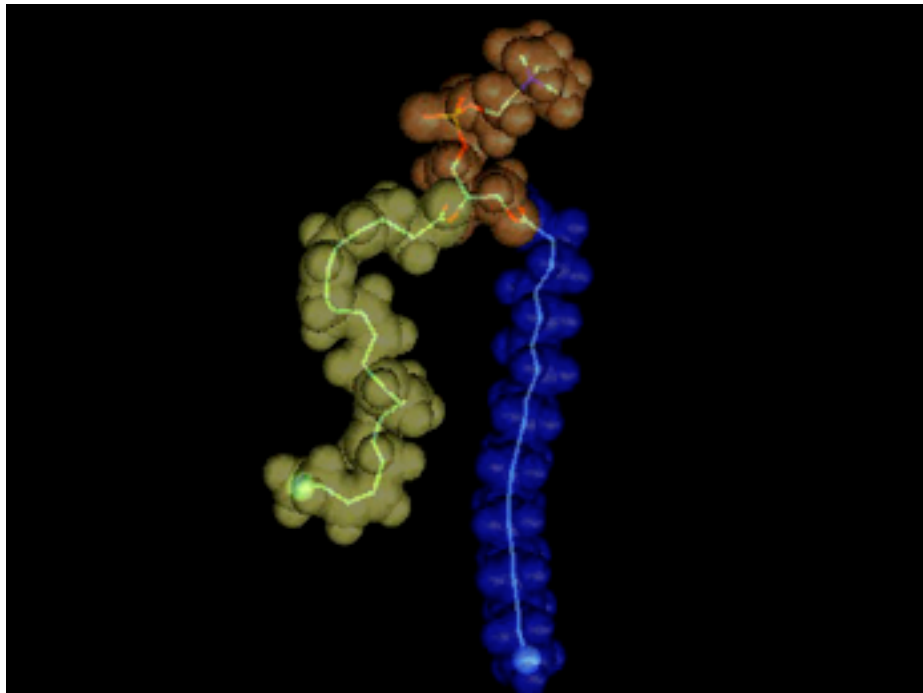
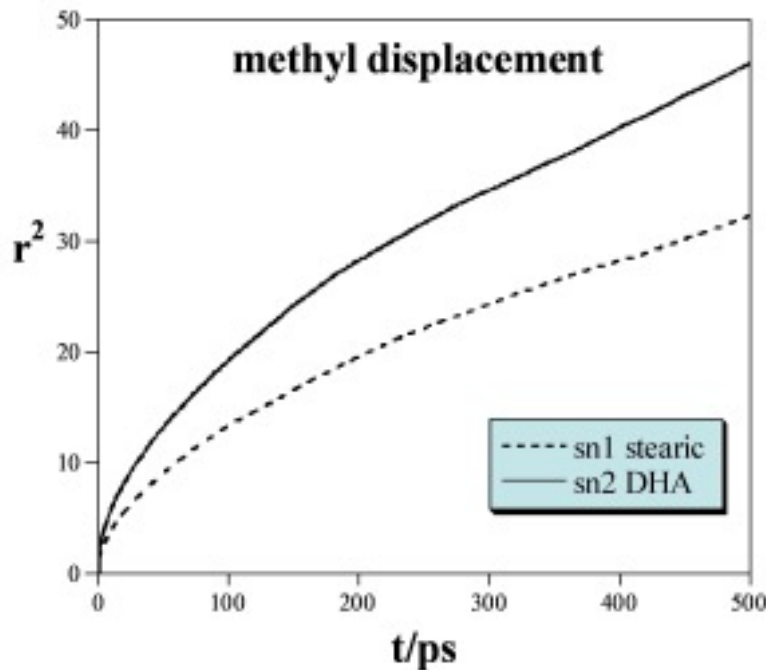
# DHA conformations from MD

- rotational barriers are extremely small
- many conformers are accessible w/ short lifetimes



# Dynamics of saturated vs. polyunsaturated lipid chains

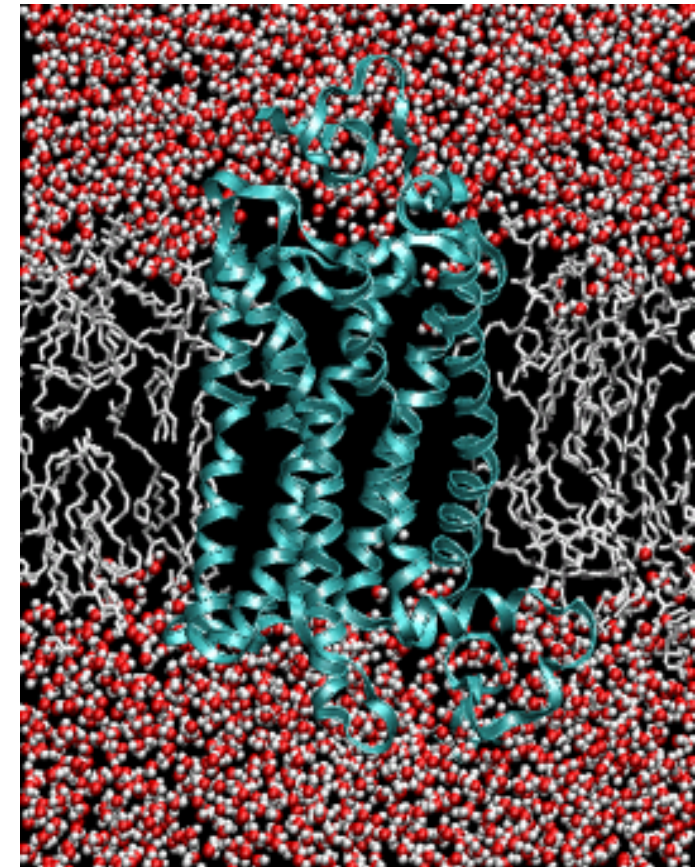
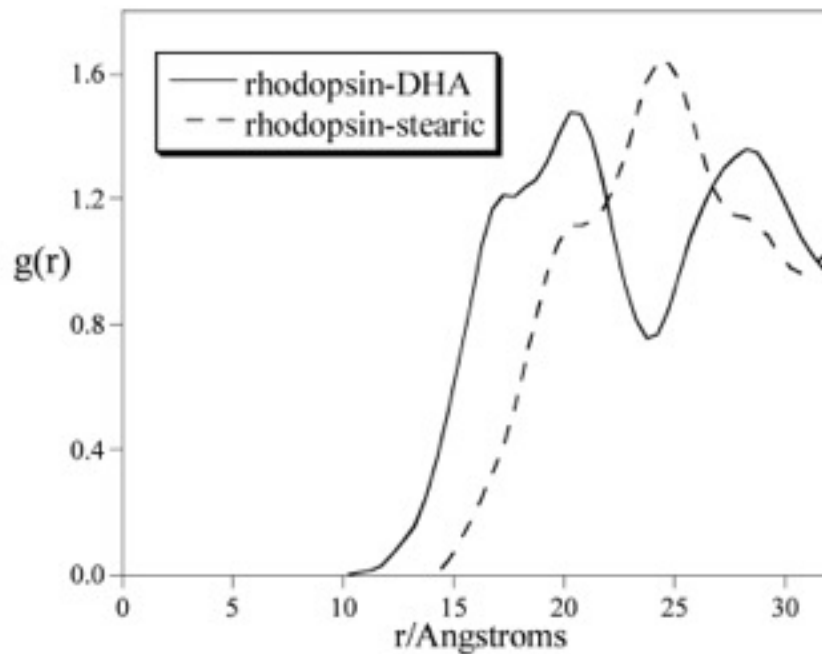
- $sn1$  stearic acid = blue
- $sn2$  DHA = yellow
- 500 ps of dynamics



*Movie courtesy of Mauricio Carrillo Tripp*

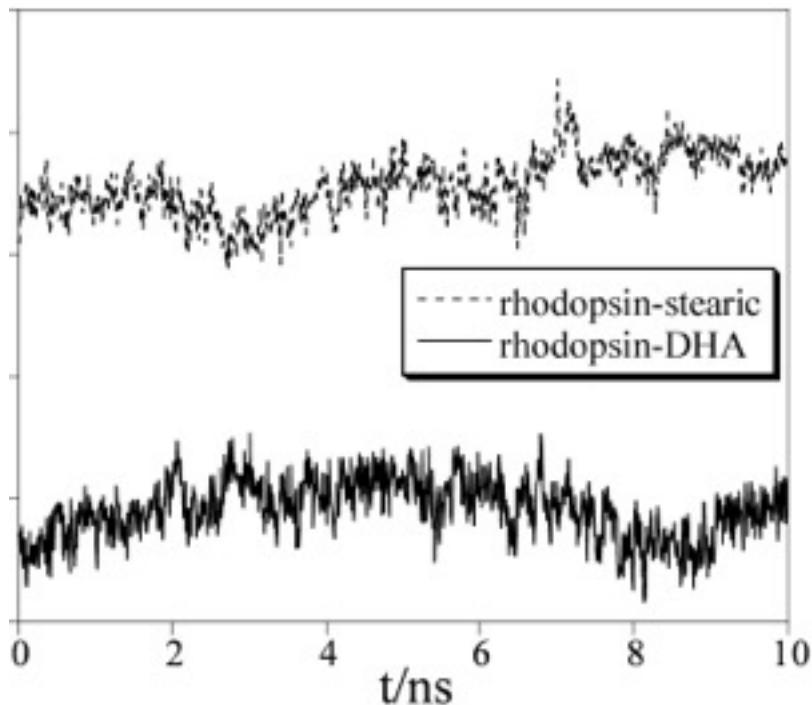
# Lipid-protein interactions

- Radial distribution around protein shows distinct layering of acyl chains



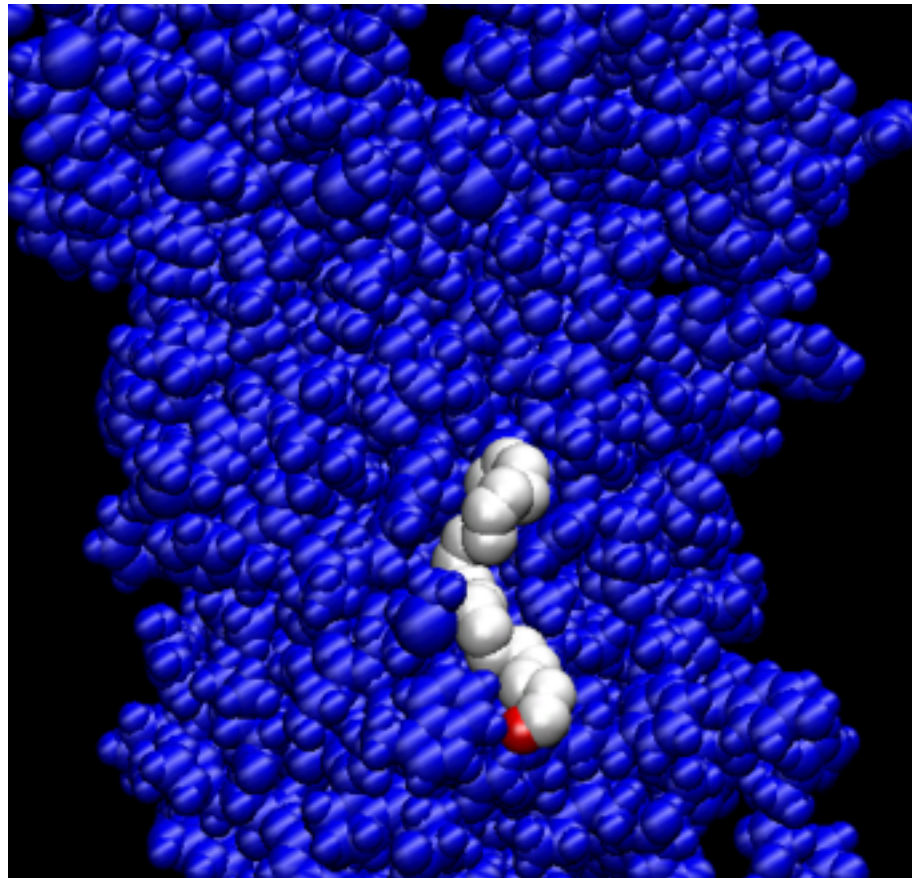
# Lipid-protein interactions

- Decomposition of non-bonded interaction shows rhodopsin is strongly attracted to unsaturated chain
- All hydrophobic residues are stabilized by DHA



<u>resname</u>	$U_{DHA}$	$U_{stearic}$	<u>ratio</u>
PHE	-44.9	-22.6	2.0
ILE	-30.0	-10.1	3.0
VAL	-24.0	-9.6	2.5
LEU	-23.1	-13.0	1.8
MET	-22.8	-9.7	2.4
TYR	-18.6	-10.4	1.8
ALA	-11.4	-3.0	3.8
TRP	-10.3	-2.4	4.2

# Origin of protein:DHA attraction



- Flexibility of the DHA chain allows solvation of the rough protein surface to occur with little intra-molecular energy cost

# Major Recent Developments

- New set of lipid force field parameters for CHARMM (CHARMM32<sup>+</sup>)
  - Pastor, B. Brooks, MacKerell
- Polarizable force field
  - Roux, MacKerell