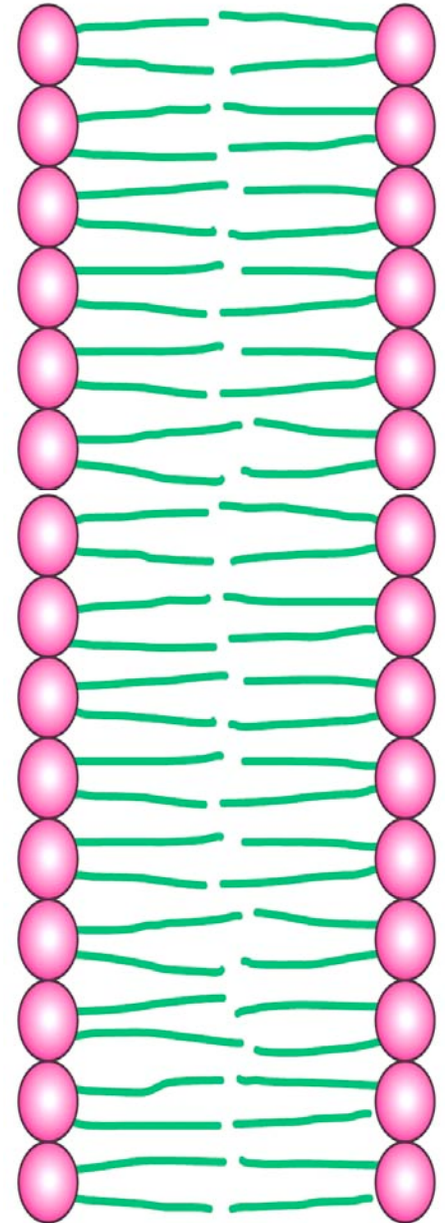
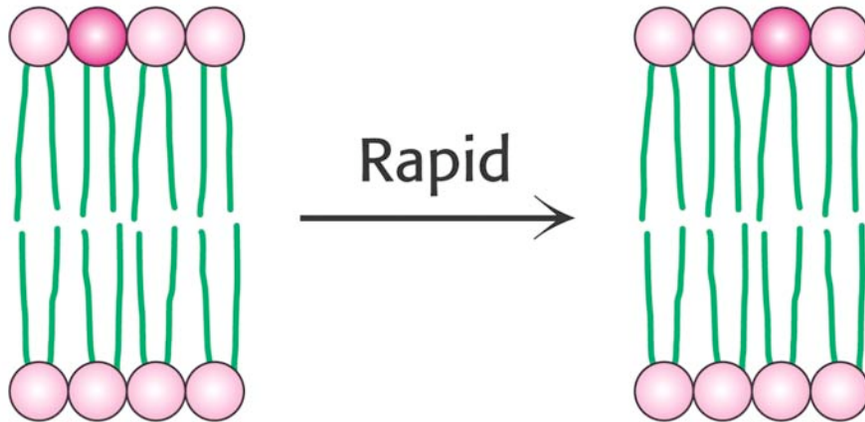


Lipid Bilayers Are Excellent For Cell Membranes

- Hydrophobic interaction is the driving force
- Self-assembly in water
- Tendency to close on themselves
- Self-sealing (a hole is unfavorable)
- Extensive: up to millimeters



Lipid Diffusion in Membrane

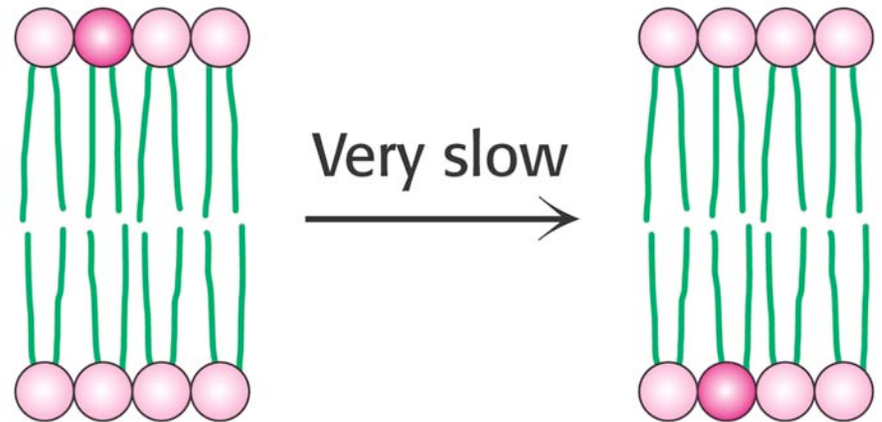


Lateral diffusion

$$D = 1 \mu\text{m}^2 \cdot \text{s}^{-1}$$

$$50 \text{ \AA} \text{ in } \sim 2.5 \times 10^{-5} \text{ s}$$

$$D_{\text{lip}} = 10^{-8} \text{ cm}^2 \cdot \text{s}^{-1}$$
$$D_{\text{wat}} = 2.5 \times 10^{-5} \text{ cm}^2 \cdot \text{s}^{-1}$$



Transverse diffusion
(flip-flop)

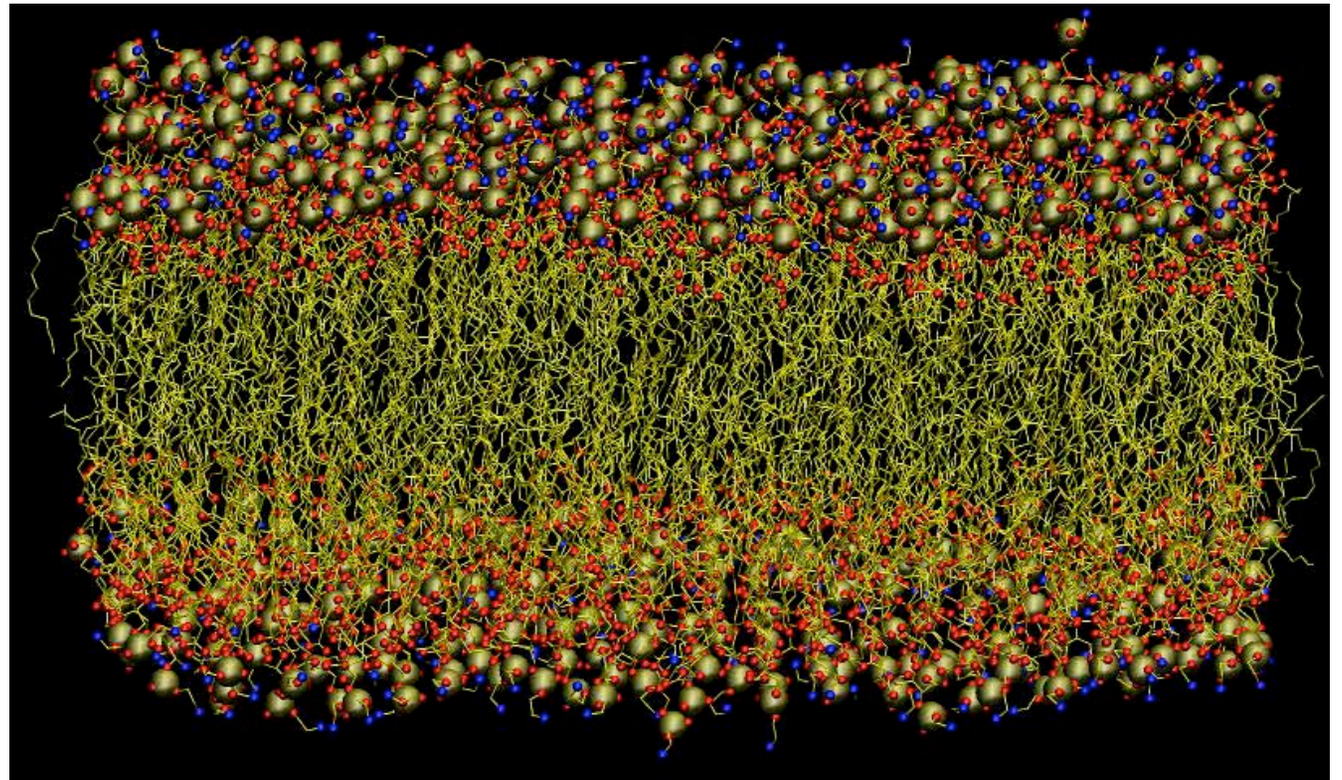
Once in several hours!
(10^4 s)

**~9 orders of magnitude
difference**

Technical difficulties in Simulations of Biological Membranes

- Time scale
- Heterogeneity of biological membranes ☹️

60 x 60 Å
Pure POPE
5 ns
~100,000
atoms



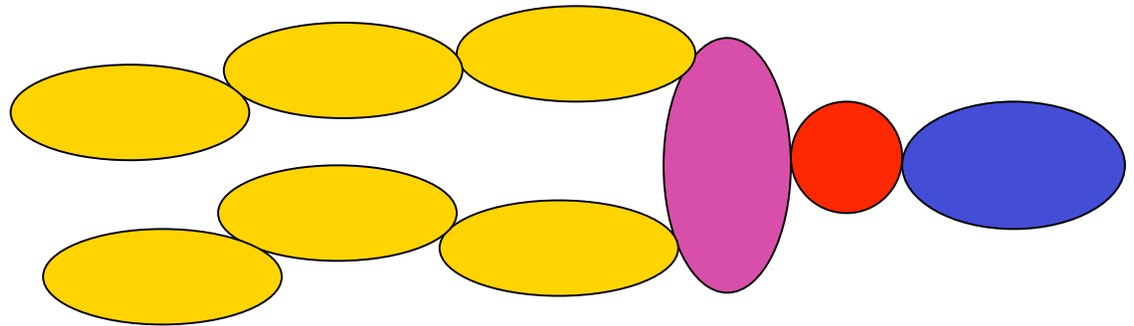
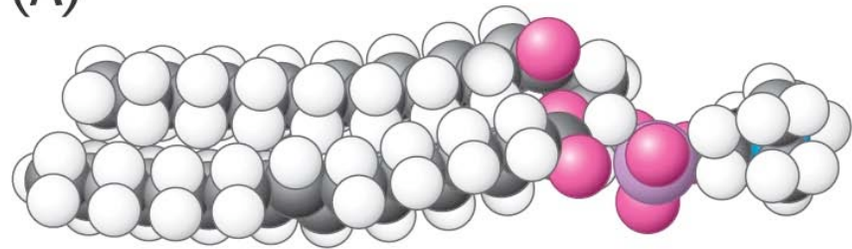
Coarse grain modeling of lipids

150 particles

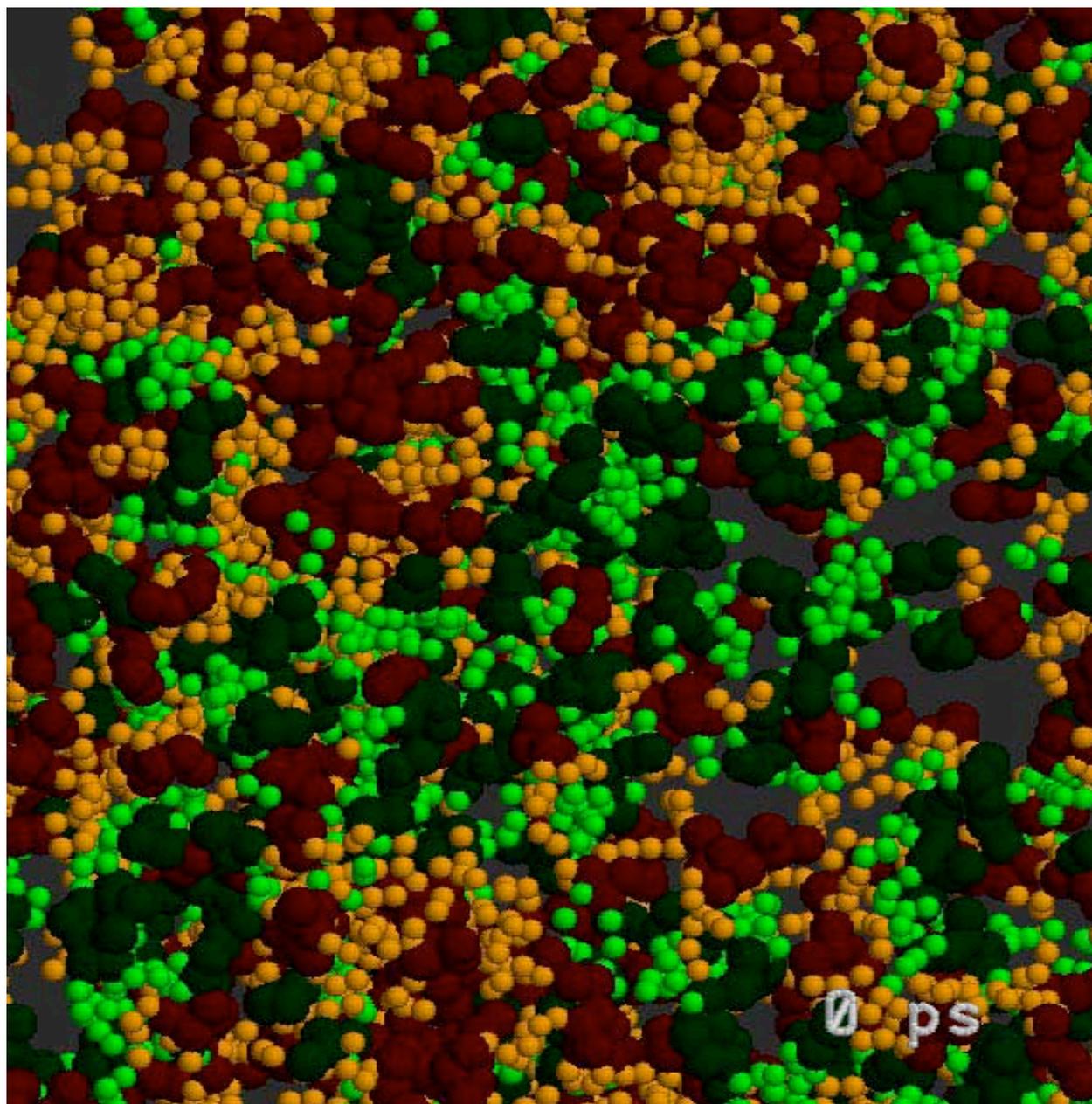


9 particles!

(A)



Also, increasing the time step by orders of magnitude.



by: J. Siewert-Jan Marrink and Alan E. Mark, University of Groningen, The Netherlands

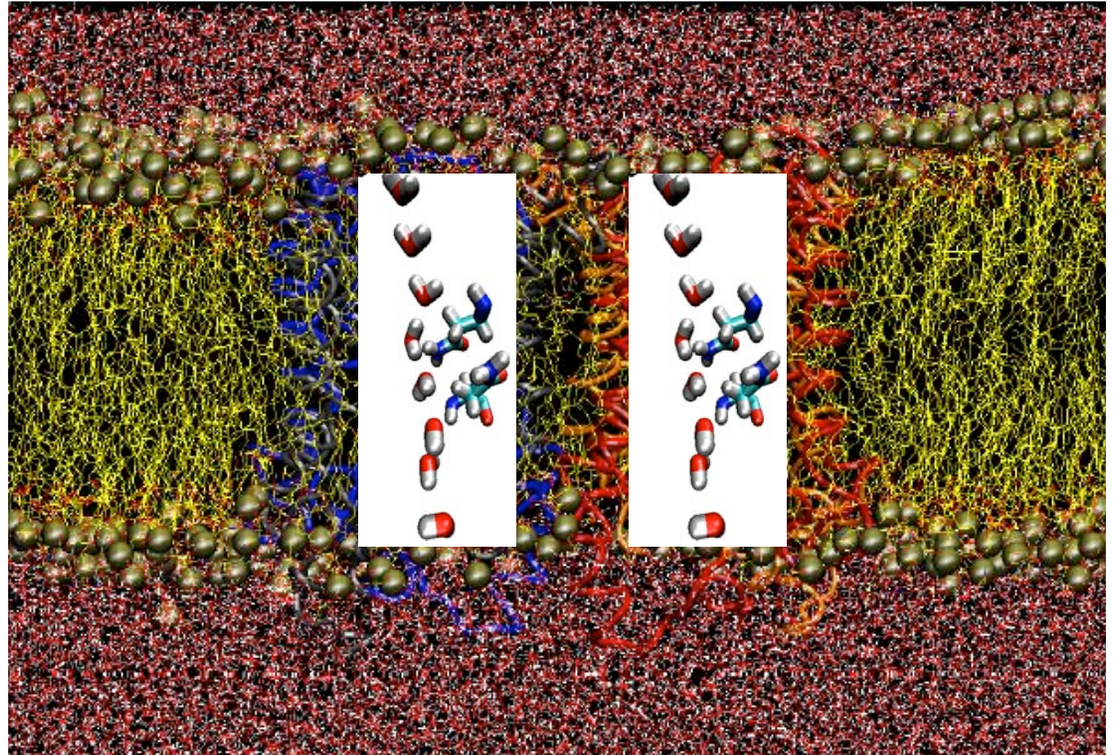
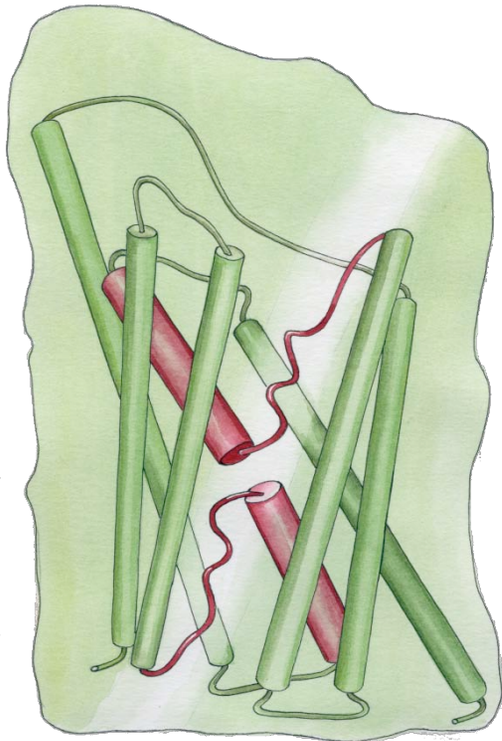
Analysis of Molecular Dynamics Simulations of Biomolecules

- A very complicated arrangement of hundreds of groups interacting with each other
- Where to start to look at?
- What to analyze?
- How much can we learn from simulations?

It is very important to get acquainted with your system

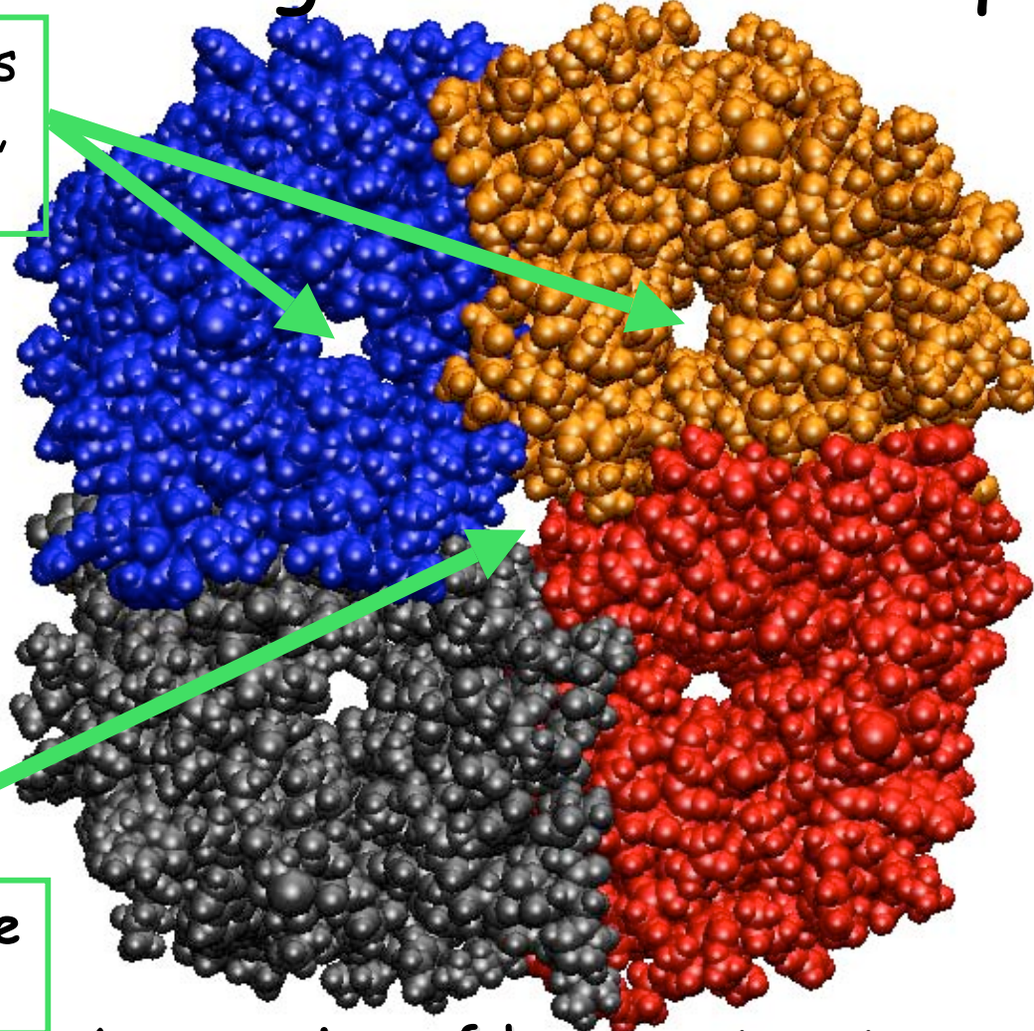
Aquaporins

Membrane water channels



Structural Organization of Aquaporins

Monomeric pores
Water, glycerol,
urea, H₂S, ...



Tetrameric pore
????

Aquaporins of known structure:

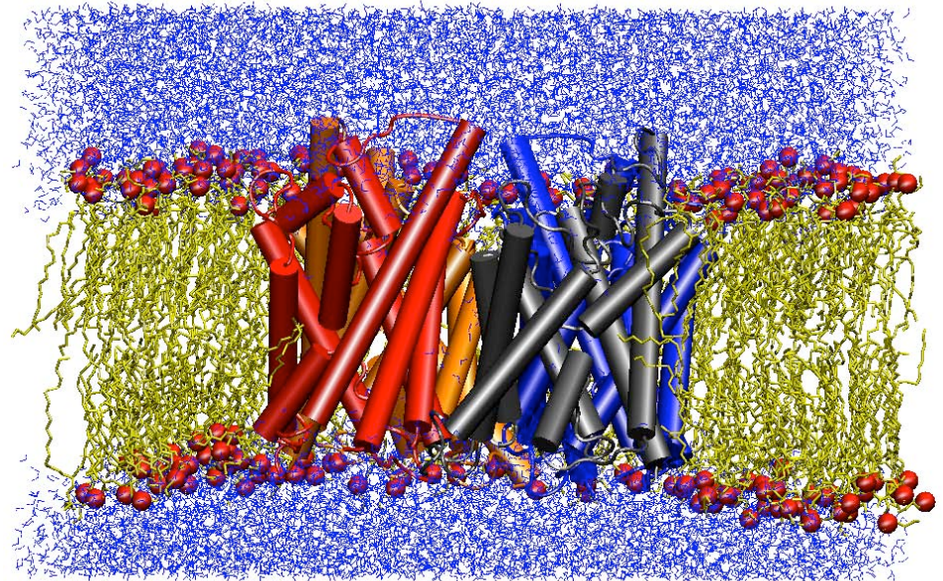
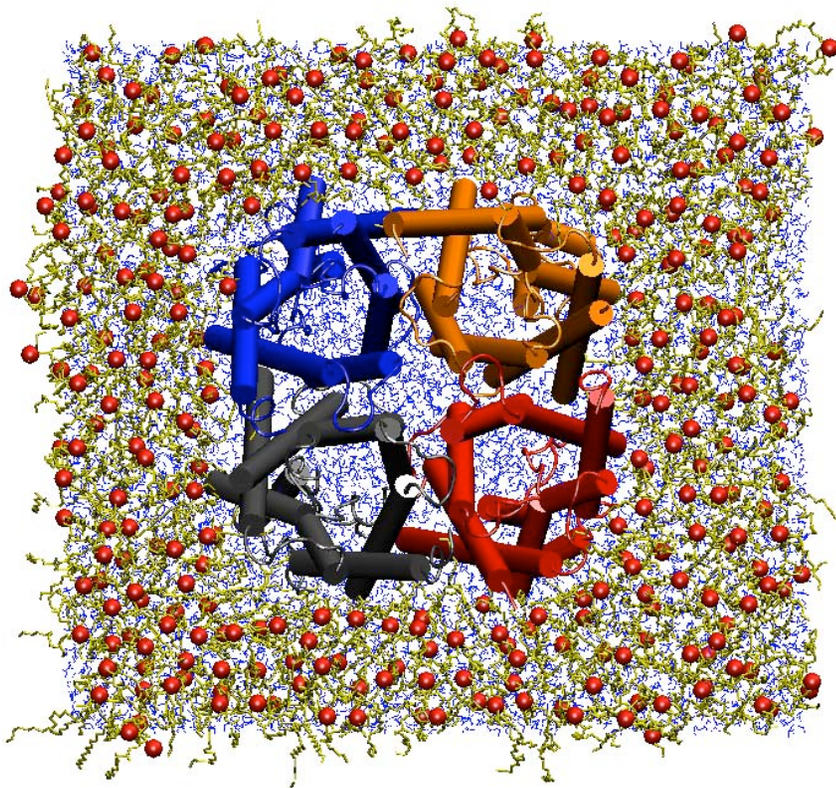
GlpF - E. coli glycerol channel

AQP1 - Mammalian aquaporin-1

AqpZ and AQP0 (2004, 2005), AqpM (2005), soPIP2 (2006)

Molecular Dynamics Simulations

Protein: ~ 15,000 atoms
Lipids (POPE): ~ 40,000 atoms
Water: ~ 51,000 atoms
Total: ~ 106,000 atoms



NAMD, CHARMM27, PME

NpT ensemble at 310 K

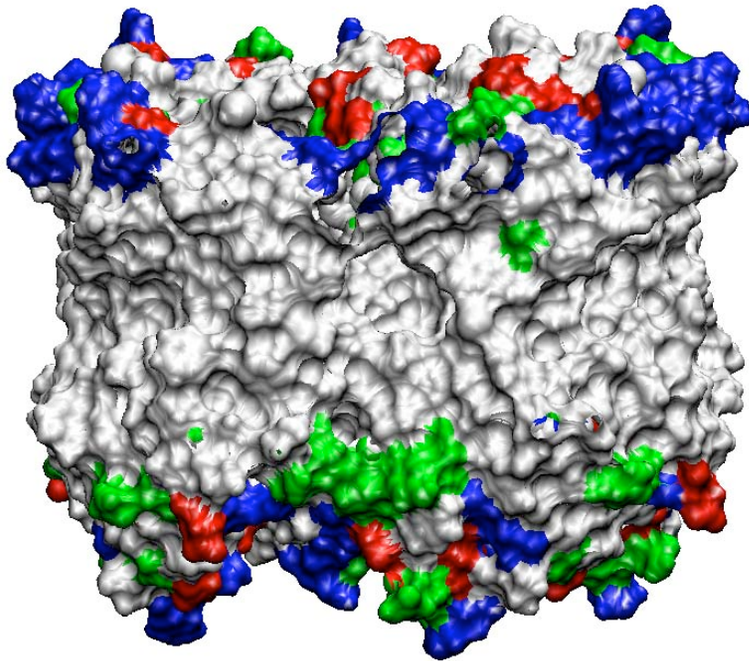
1ns equilibration, 4ns production

10 days /ns - 32-proc Linux cluster

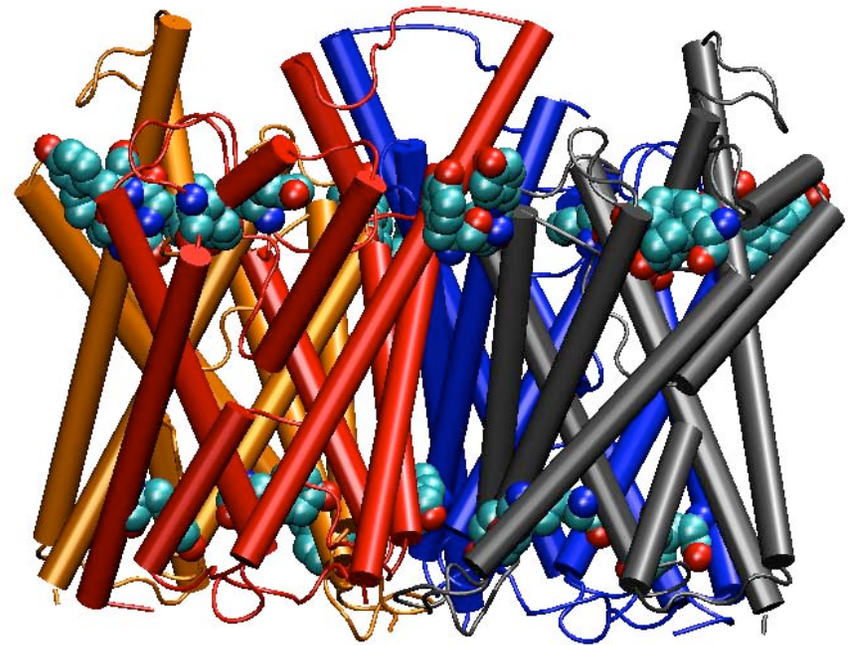
3.5 days/ns - 128 O2000 CPUs

0.35 days/ns - 512 LeMieux CPUs

Protein Embedding in Membrane



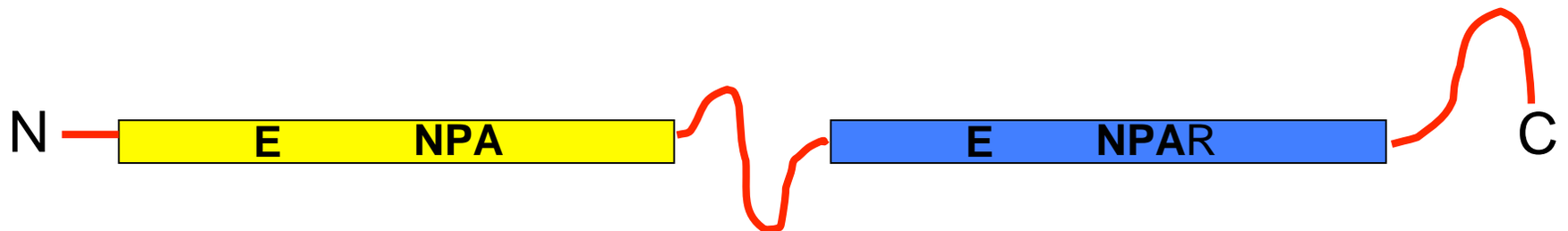
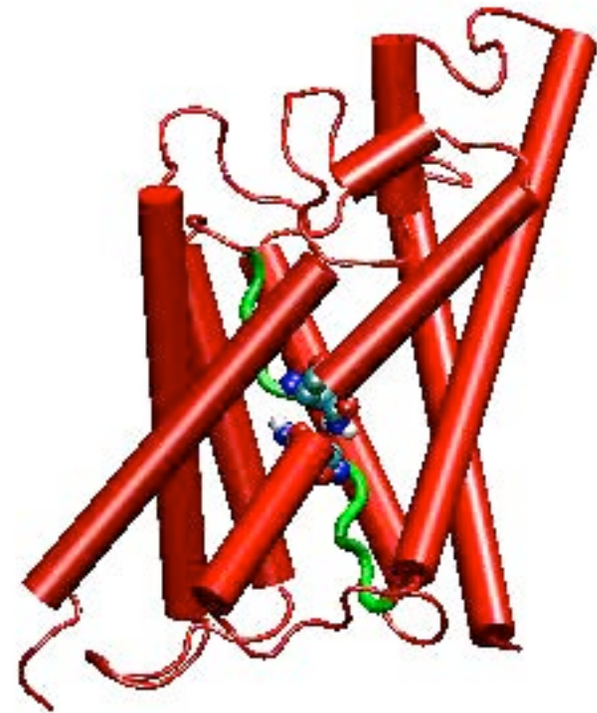
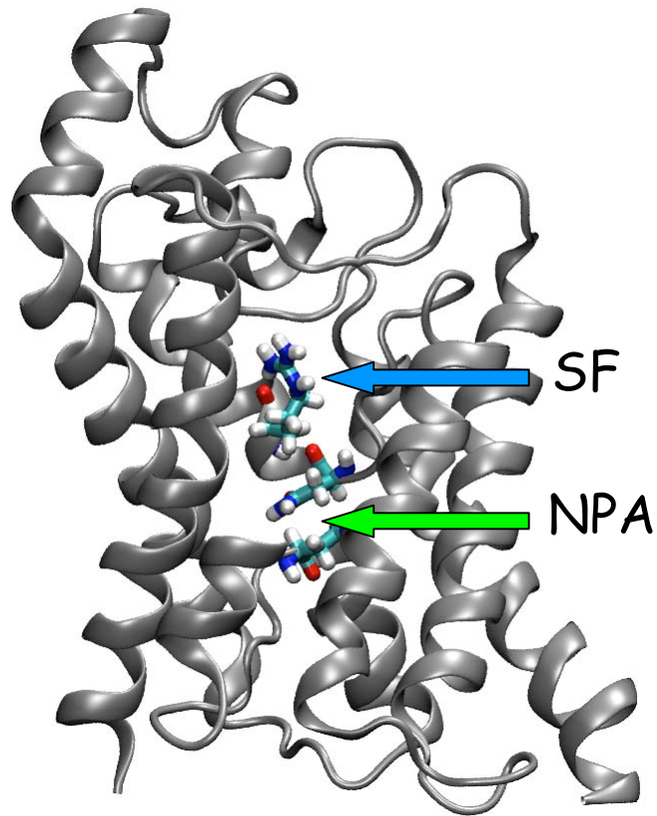
Hydrophobic surface
of the protein

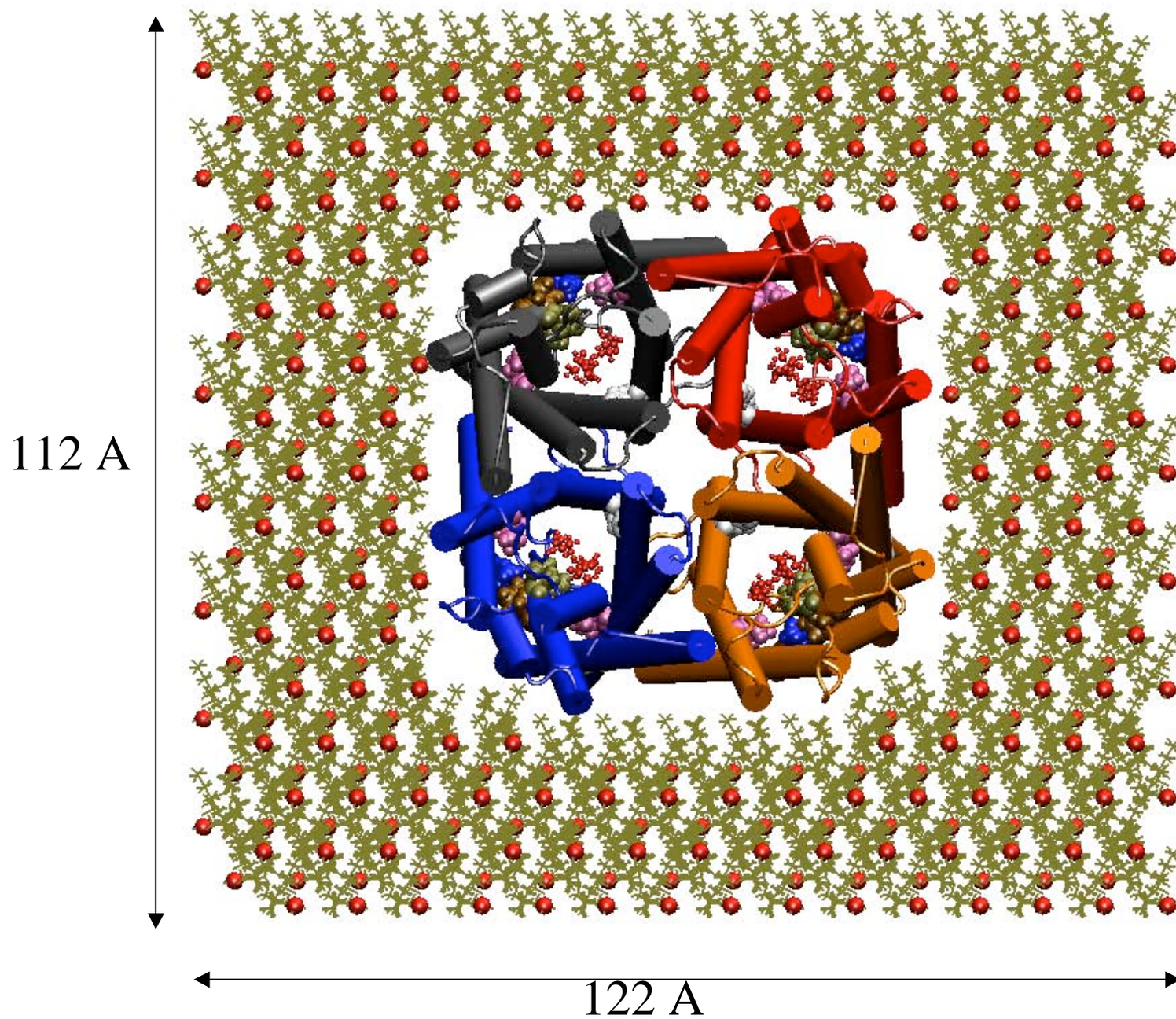


Ring of
Tyr and Trp

GlpF in VMD

Structurally Conserved Features

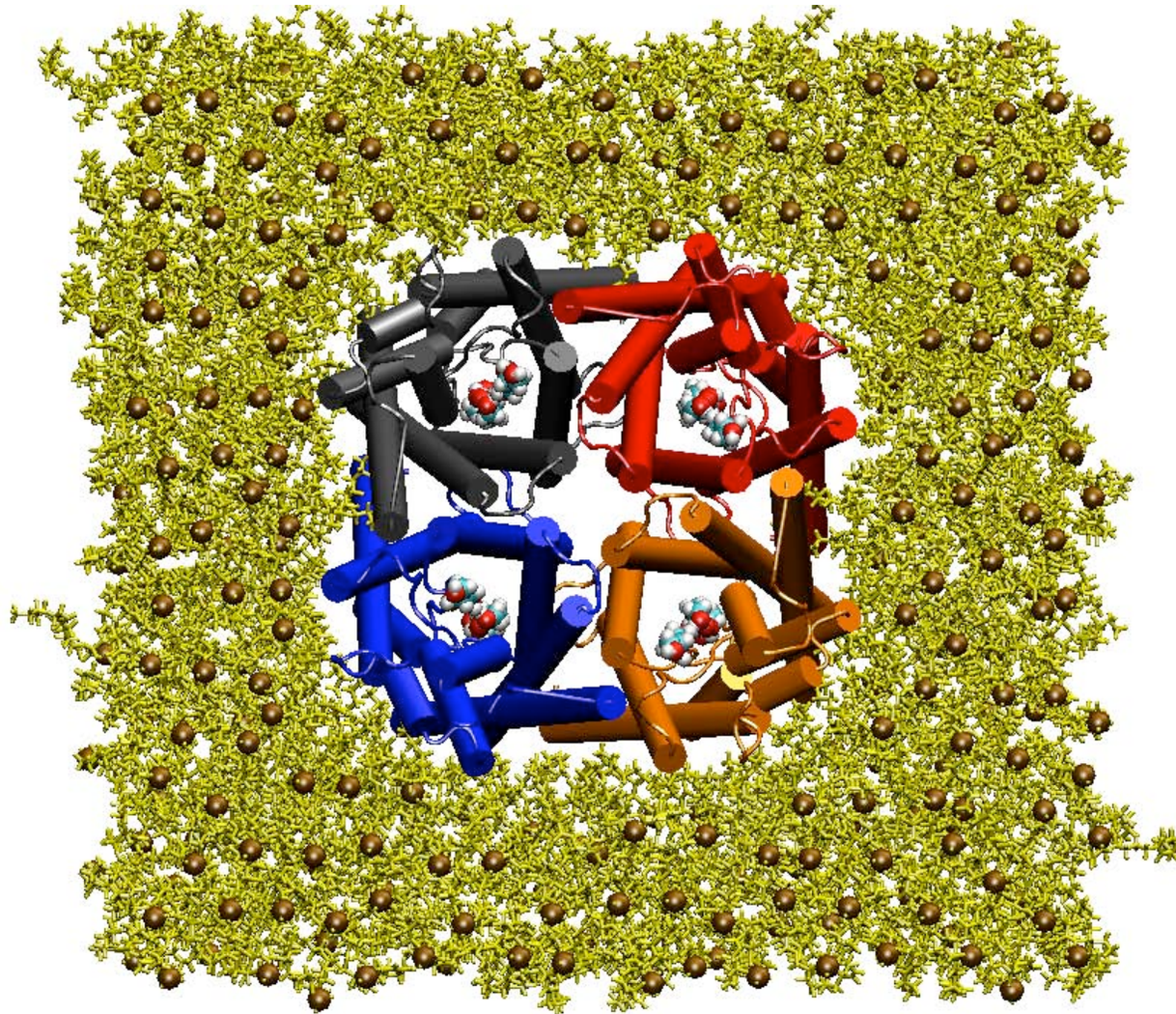




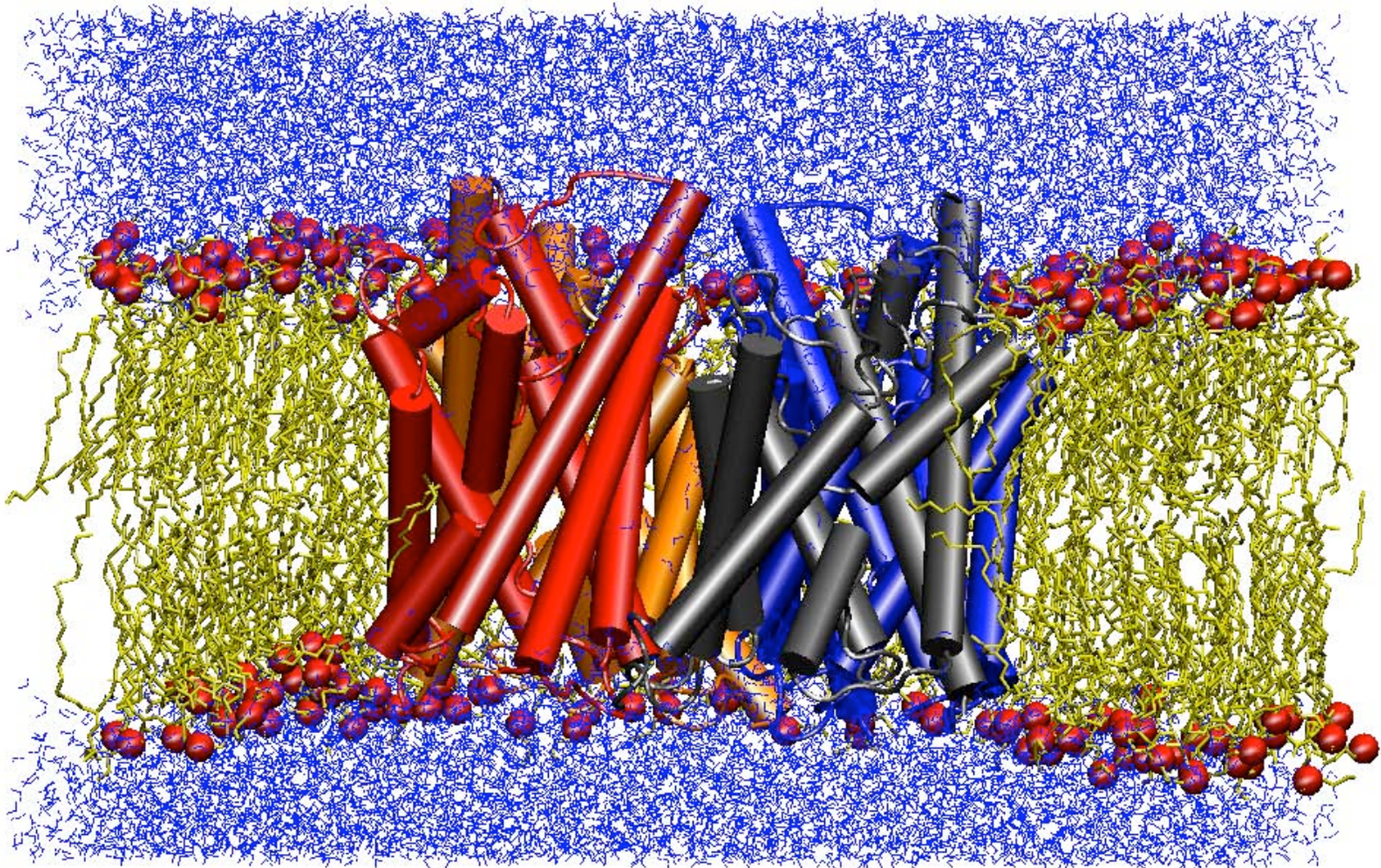
A Recipe for Membrane Protein Simulations

- Insert your protein into a hydrated lipid bilayer.
- Fix the protein; minimize the rest and run a short “constant-pressure” MD to bring lipids closer to the protein and fill the gap between the protein and lipids.
- Watch water molecules; if necessary apply constraints to prevent them from penetrating into the open gaps between lipids and the protein.
- Monitor the volume of your simulation box until it is almost constant. Do not run the system for too long during this phase.
- Now release the protein, minimize the whole system, and start an NpT simulation of the whole system.
- If desired, you may switch to an NVT simulation, when the system reaches a stable volume.

Lipid-Protein Packing During the Initial NpT Simulation

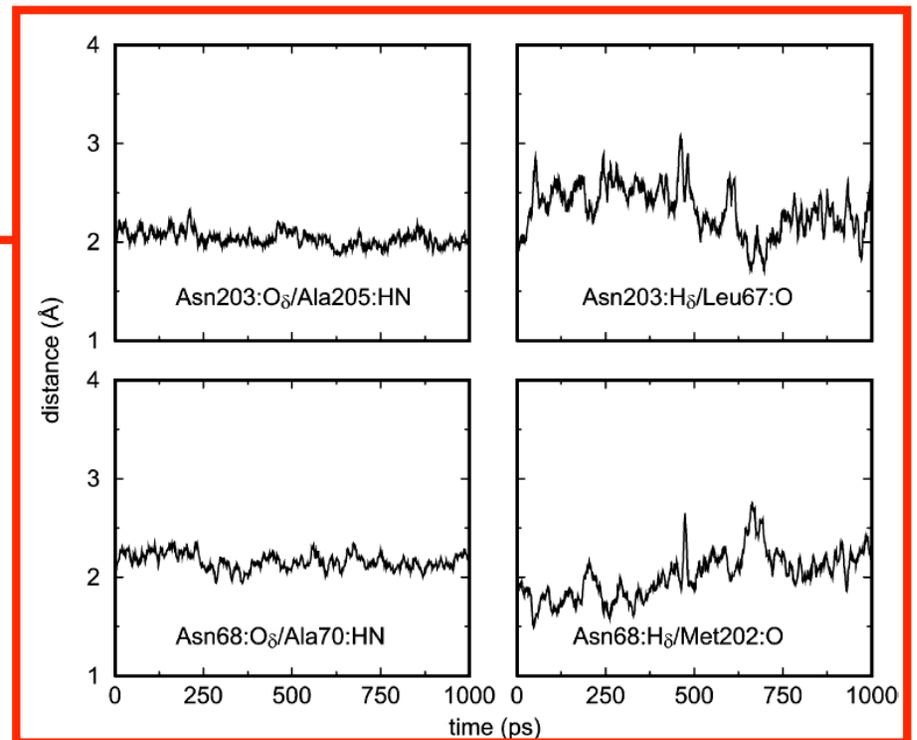
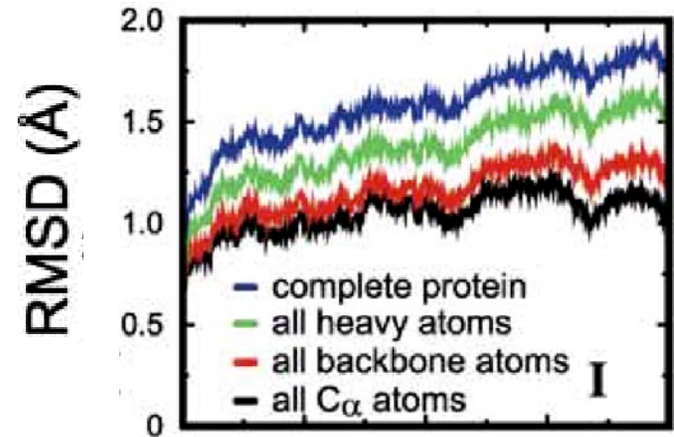
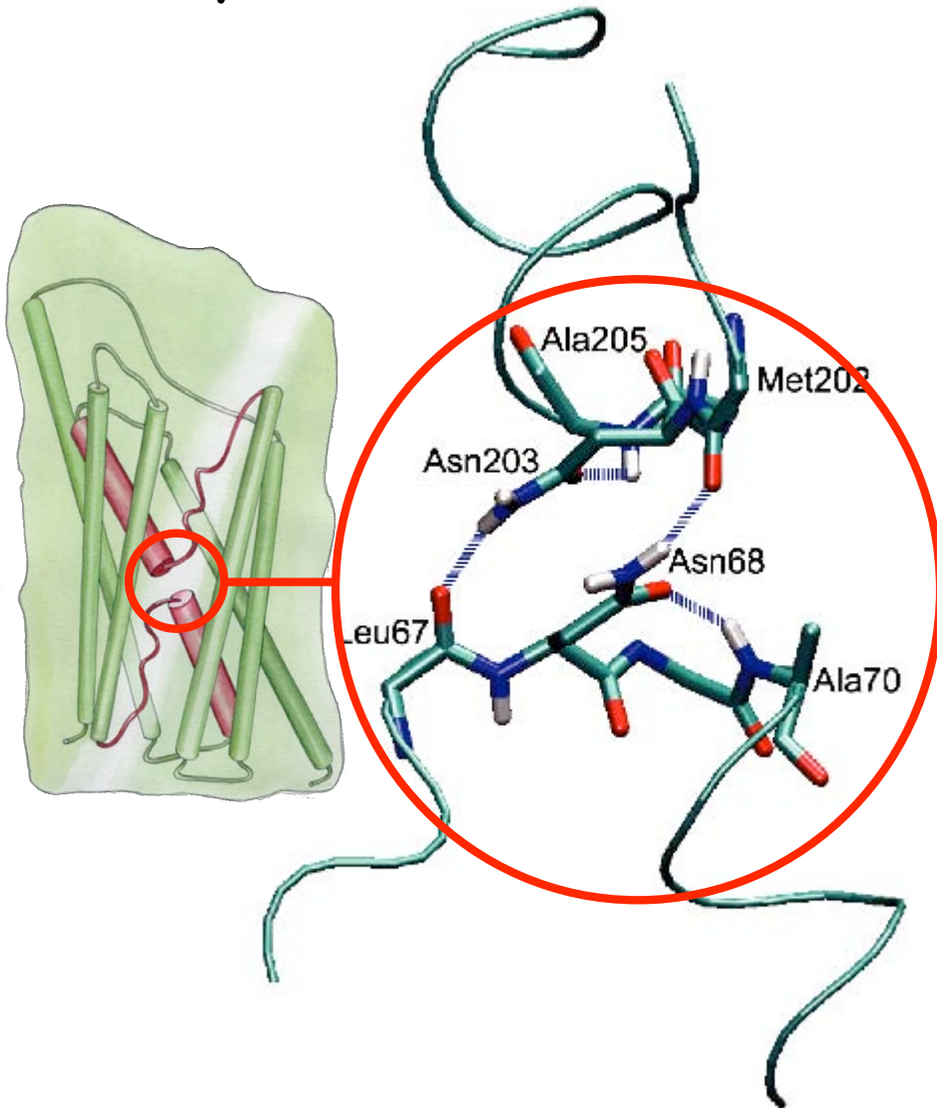


Adjustment of Membrane Thickness to the Protein Hydrophobic Surface

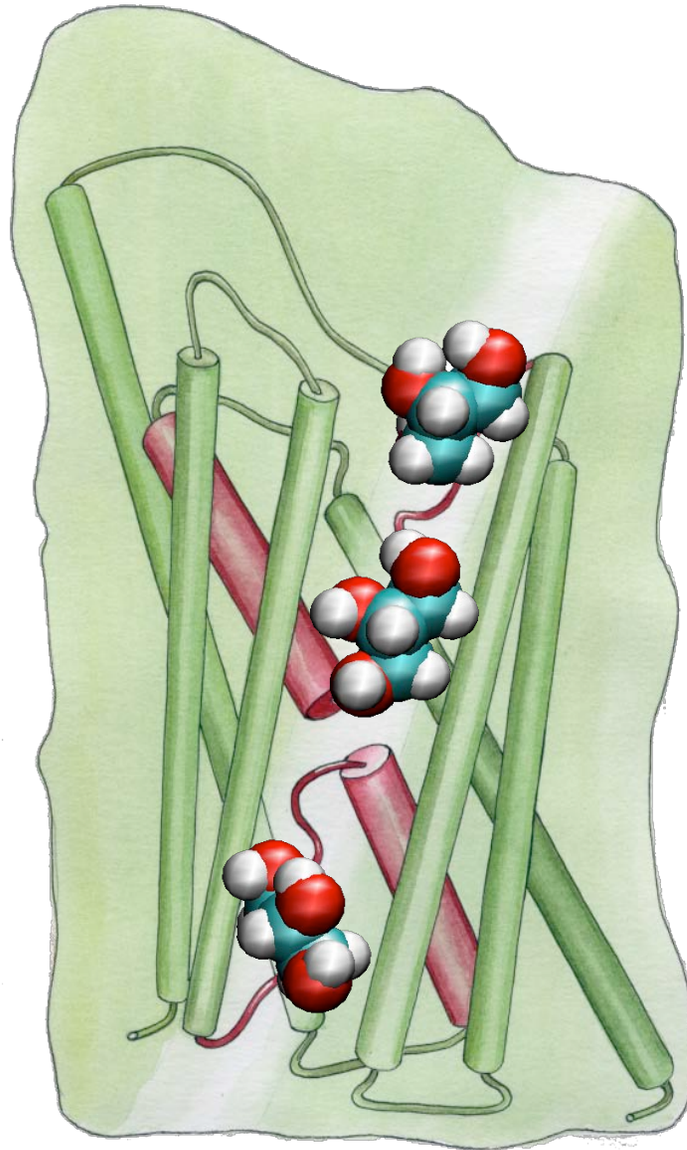


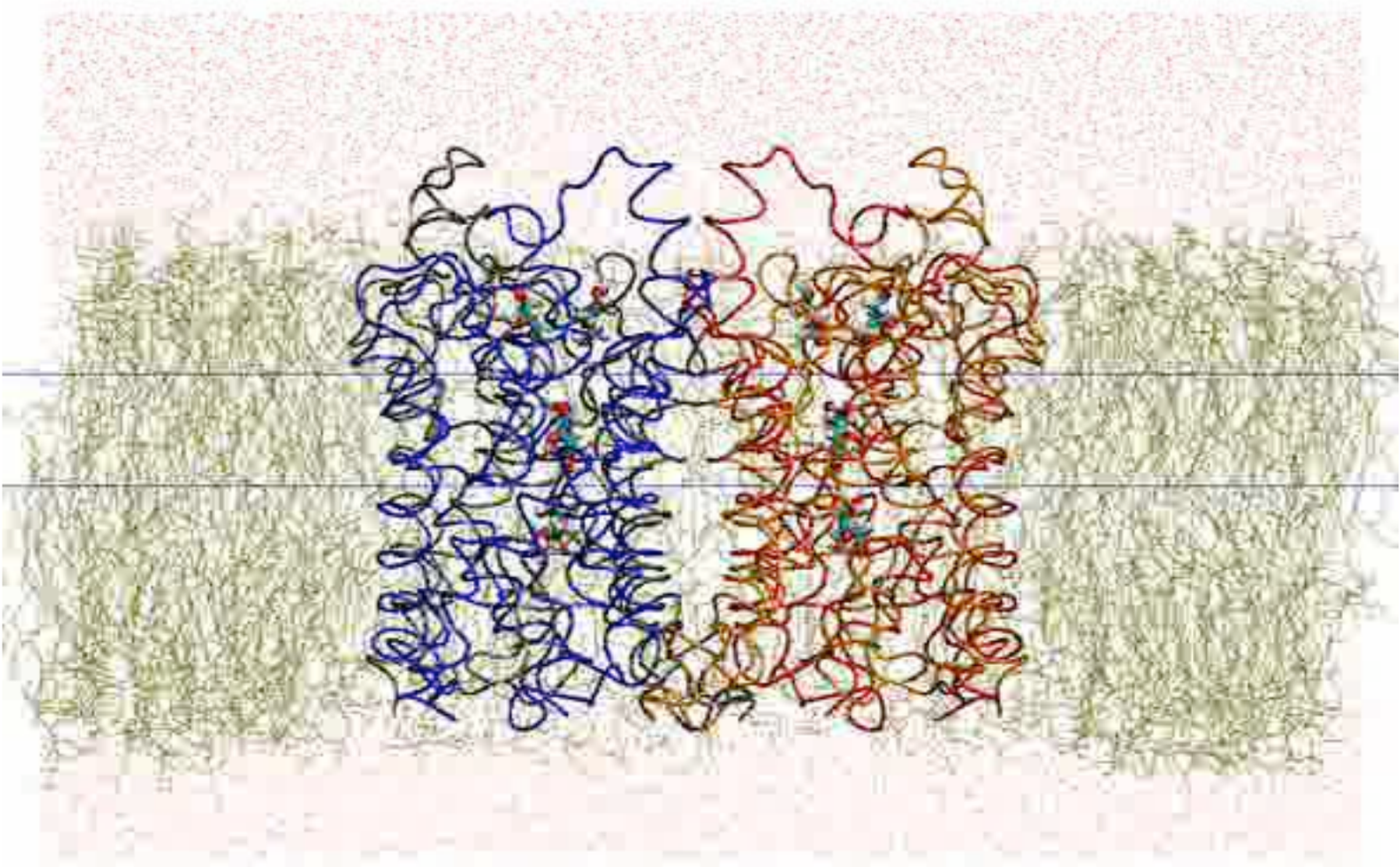
An extremely stable protein

Stability of NPA - NPA Interaction

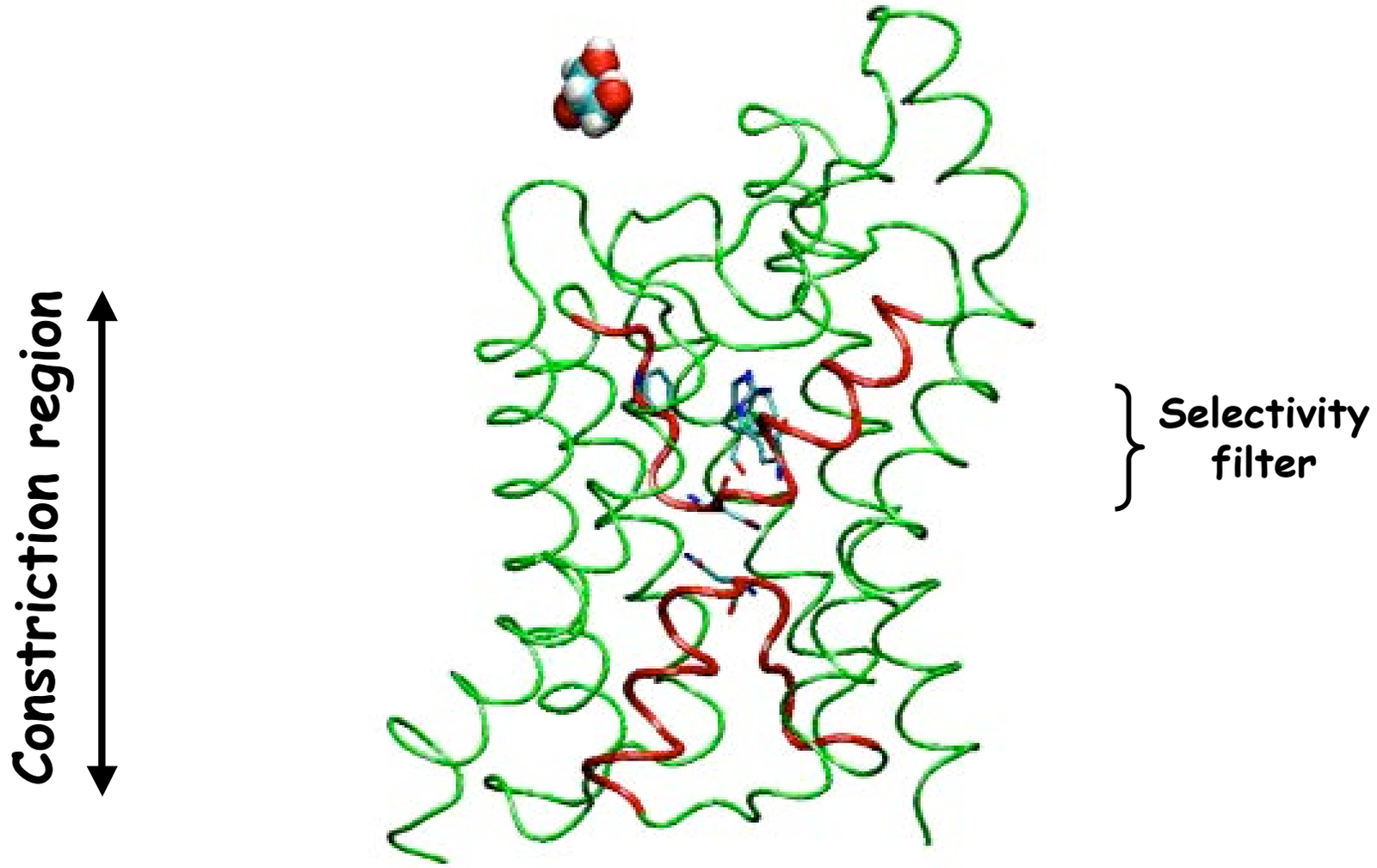


Glycerol-Saturated GlpF





Complete description of the conduction pathway



Details of Protein-Substrate Interaction Are Always Important

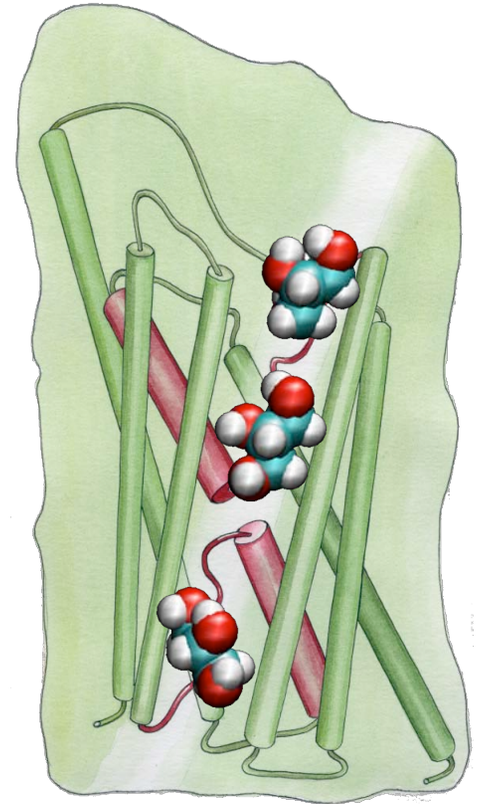
- Identify those groups of the protein that are directly involved in the main function of the protein.
- Look at the interaction of these primary residues with other groups in the protein.
- Look at buried charged residues inside the protein; they must have an important role.
- Backbone hydrogen bonds are mainly responsible for stabilization of secondary structure elements in the protein; side chain hydrogen bonds could be functionally important.

Channel Hydrogen Bonding Sites

...

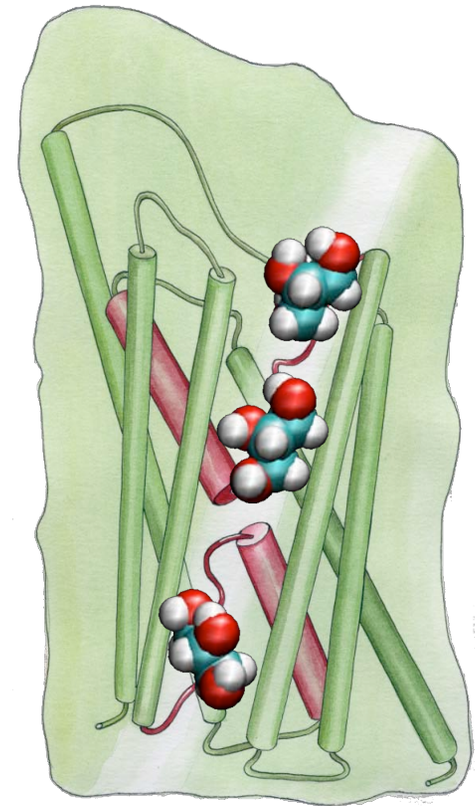
```
{set frame 0}{frame < 100}{incr frame}{  
  animate goto $frame  
  set donor [atomselect top  
    "name O N and within 2 of  
    (resname GCL and name HO)"]  
  lappend [$donor get index] list1  
  set acceptor [atomselect top  
    "resname GCL and name O and  
    within 2 of (protein and name HN HO)"]  
  lappend [$acceptor get index] list2  
}
```

...



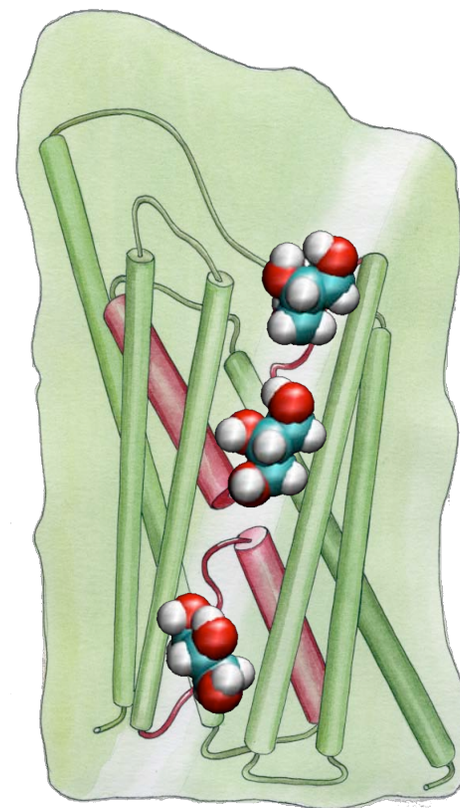
Channel Hydrogen Bonding Sites

GLN	41	OE1 NE2	LEU	197	O
TRP	48	O NE1	THR	198	O
GLY	64	O	GLY	199	O
ALA	65	O	PHE	200	O
HIS	66	O ND1	ALA	201	O
LEU	67	O	ASN	203	ND2
ASN	68	ND2			
ASP	130	OD1	LYS	33	HZ1 HZ3
GLY	133	O	GLN	41	HE21
SER	136	O	TRP	48	HE1
TYR	138	O	HIS	66	HD1
PRO	139	O N	<u>ASN</u>	68	HD22
ASN	140	OD1 ND2	<u>TYR</u>	138	HN
HIS	142	ND1	ASN	140	HN HD21 HD22
THR	167	OG1	HIS	142	HD1
GLY	195	O	GLY	199	HN
PRO	196	O	<u>ASN</u>	203	HN HD21HD22
			<u>ARG</u>	206	HE HH21HH22

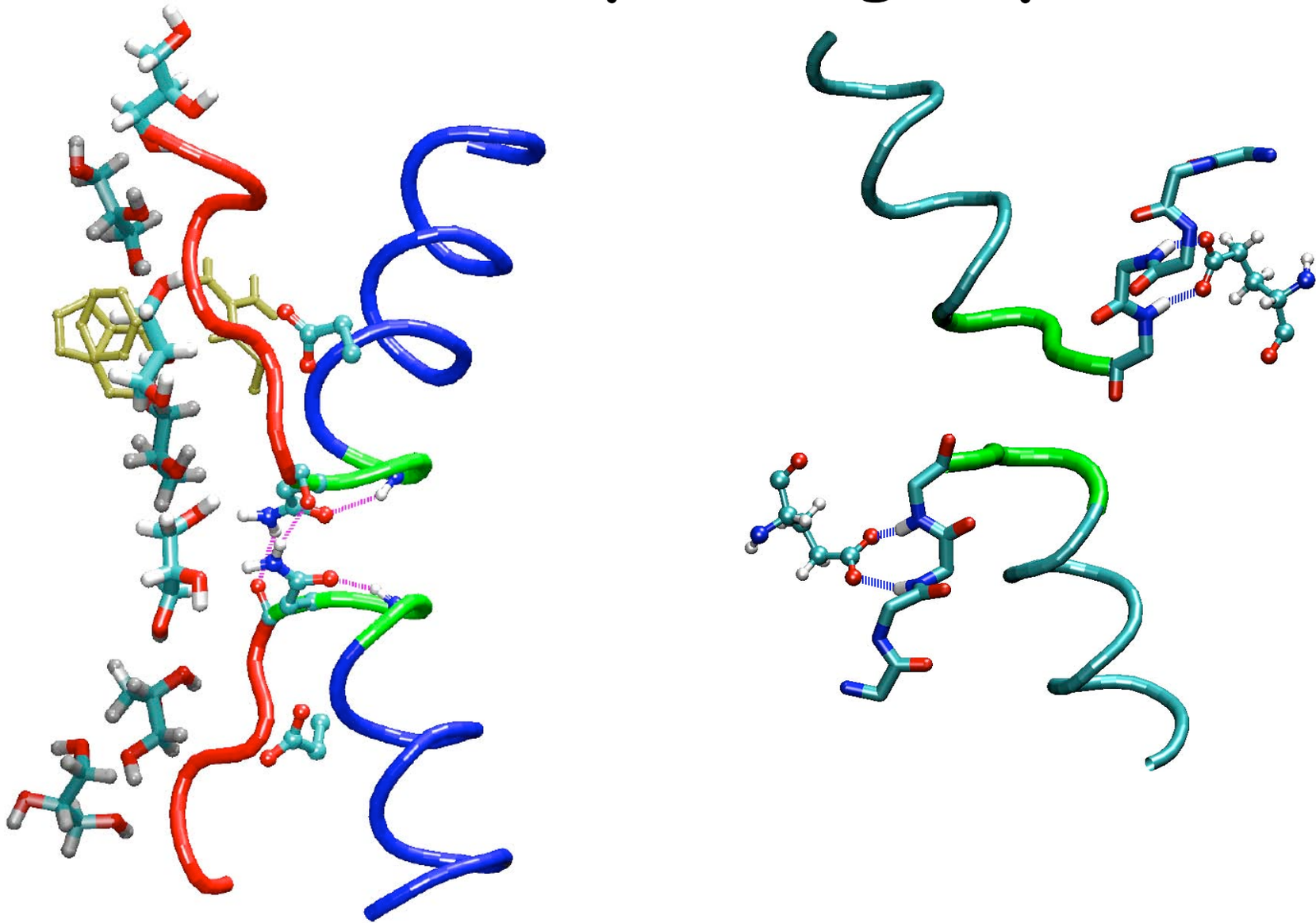


Channel Hydrogen Bonding Sites

GLN	41	OE1 NE2	LEU	197	O
TRP	48	O NE1	THR	198	O
GLY	64	O	GLY	199	O
ALA	65	O	PHE	200	O
HIS	66	O ND1	ALA	201	O
LEU	67	O	ASN	203	ND2
ASN	68	ND2			
ASP	130	OD1	LYS	33	HZ1 HZ3
GLY	133	O	GLN	41	HE21
SER	136	O	TRP	48	HE1
TYR	138	O	HIS	66	HD1
PRO	139	O N	<u>ASN</u>	68	HD22
ASN	140	OD1 ND2	TYR	138	HN
HIS	142	ND1	ASN	140	HN HD21 HD22
THR	167	OG1	HIS	142	HD1
GLY	195	O	GLY	199	HN
PRO	196	O	<u>ASN</u>	203	HN HD21HD22
			<u>ARG</u>	206	HE HH21HH22

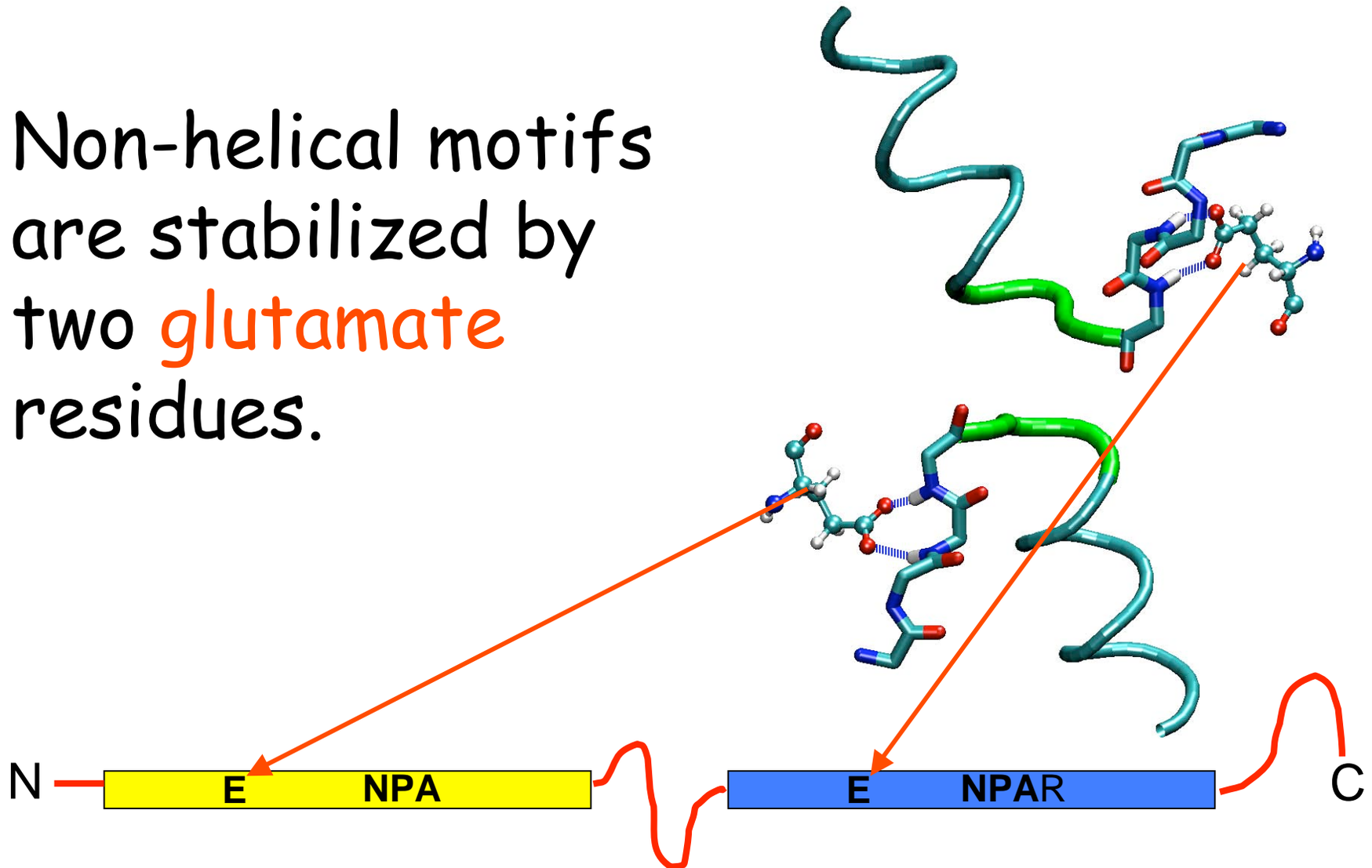


The Substrate Pathway
is formed by $C=O$ groups

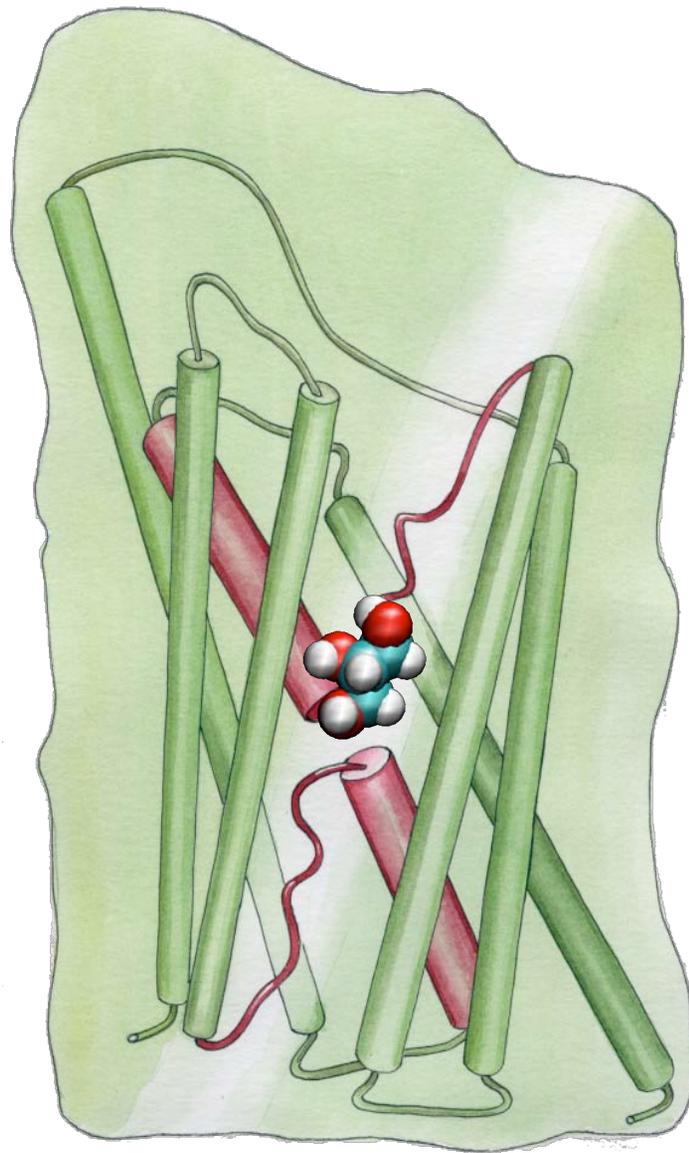


The Substrate Pathway is formed by $C=O$ groups

Non-helical motifs
are stabilized by
two **glutamate**
residues.

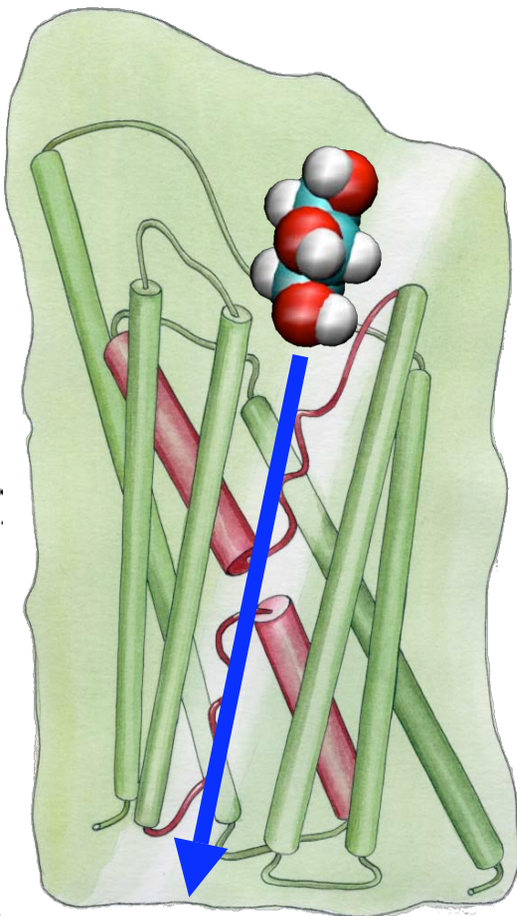
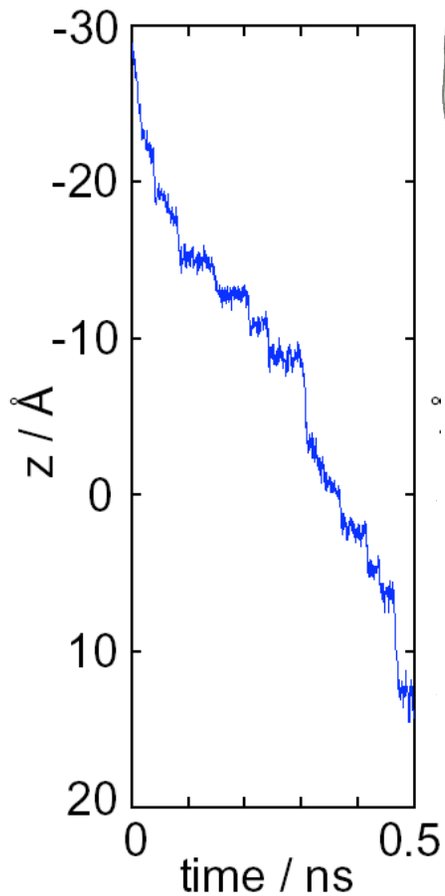


Single Glycerol per channel

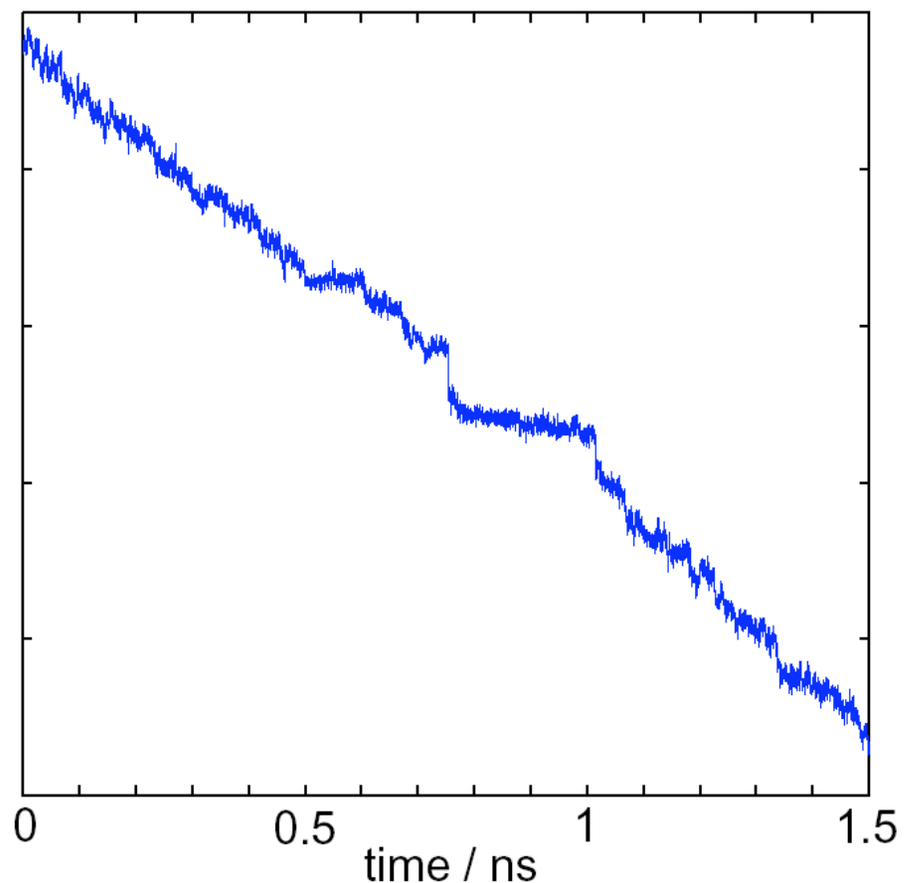


Steered Molecular Dynamics

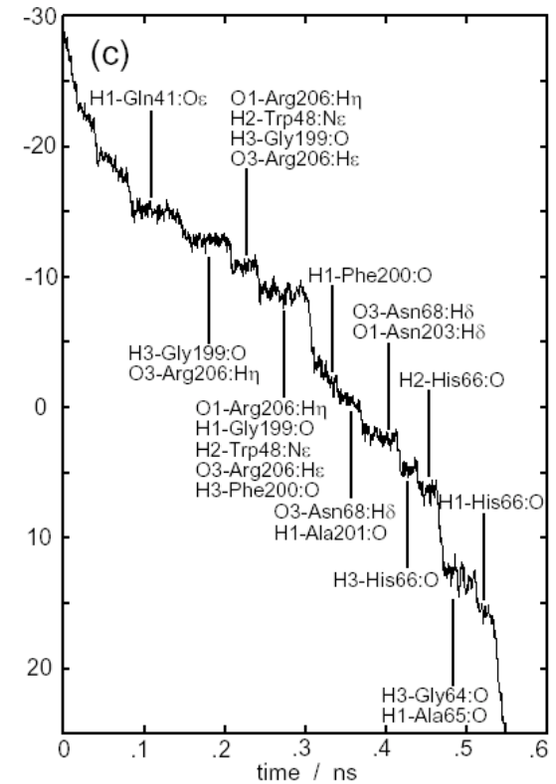
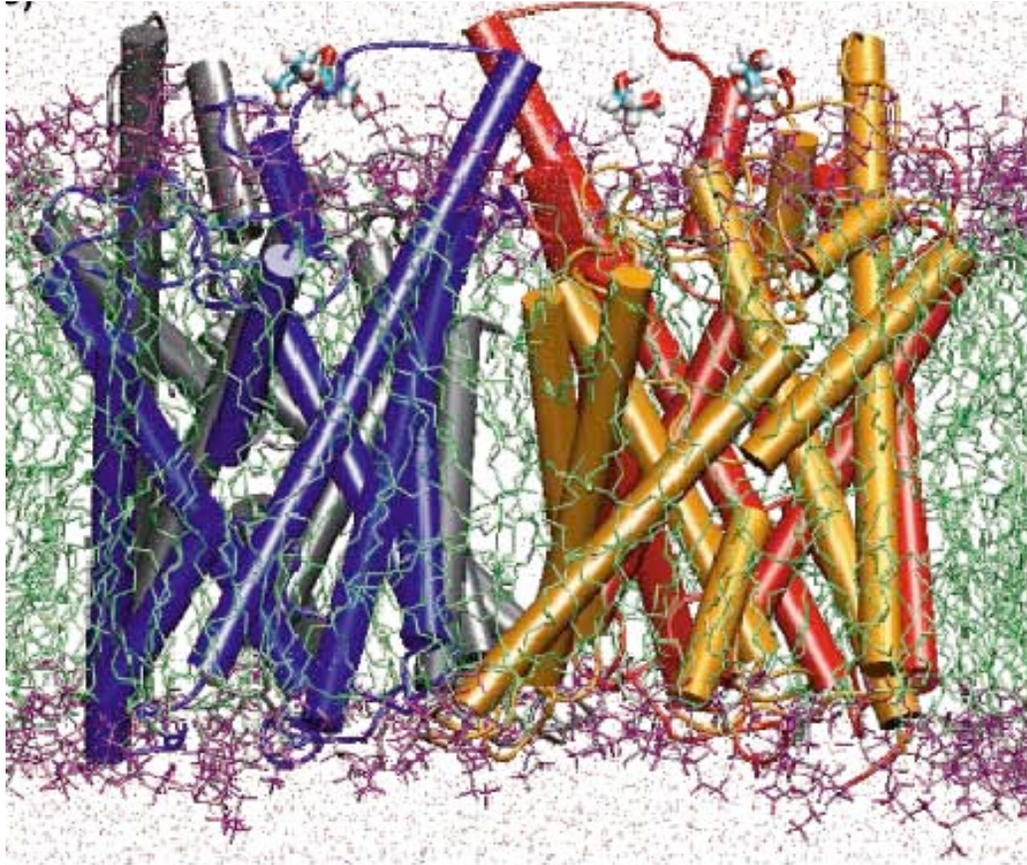
constant force
(250 pN)



constant velocity
(30 $\text{\AA}/\text{ns}$)

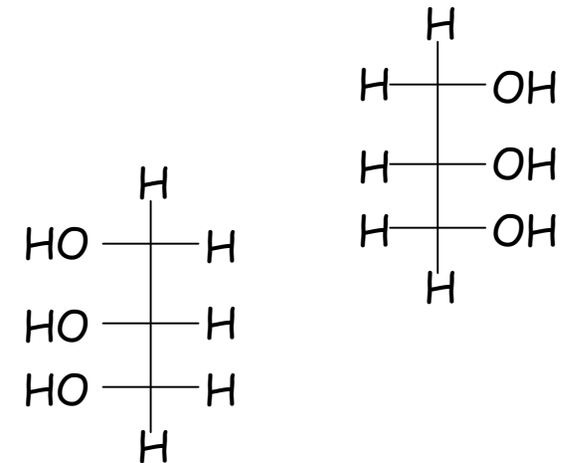
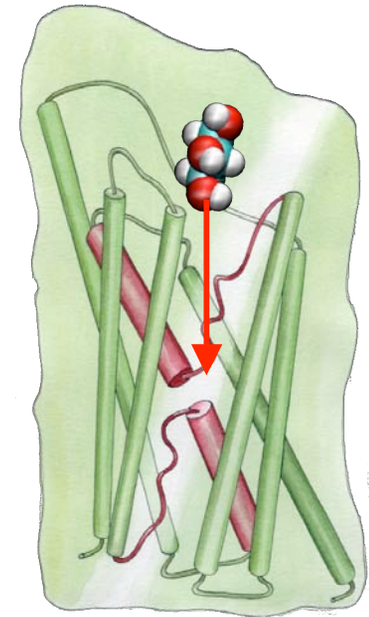
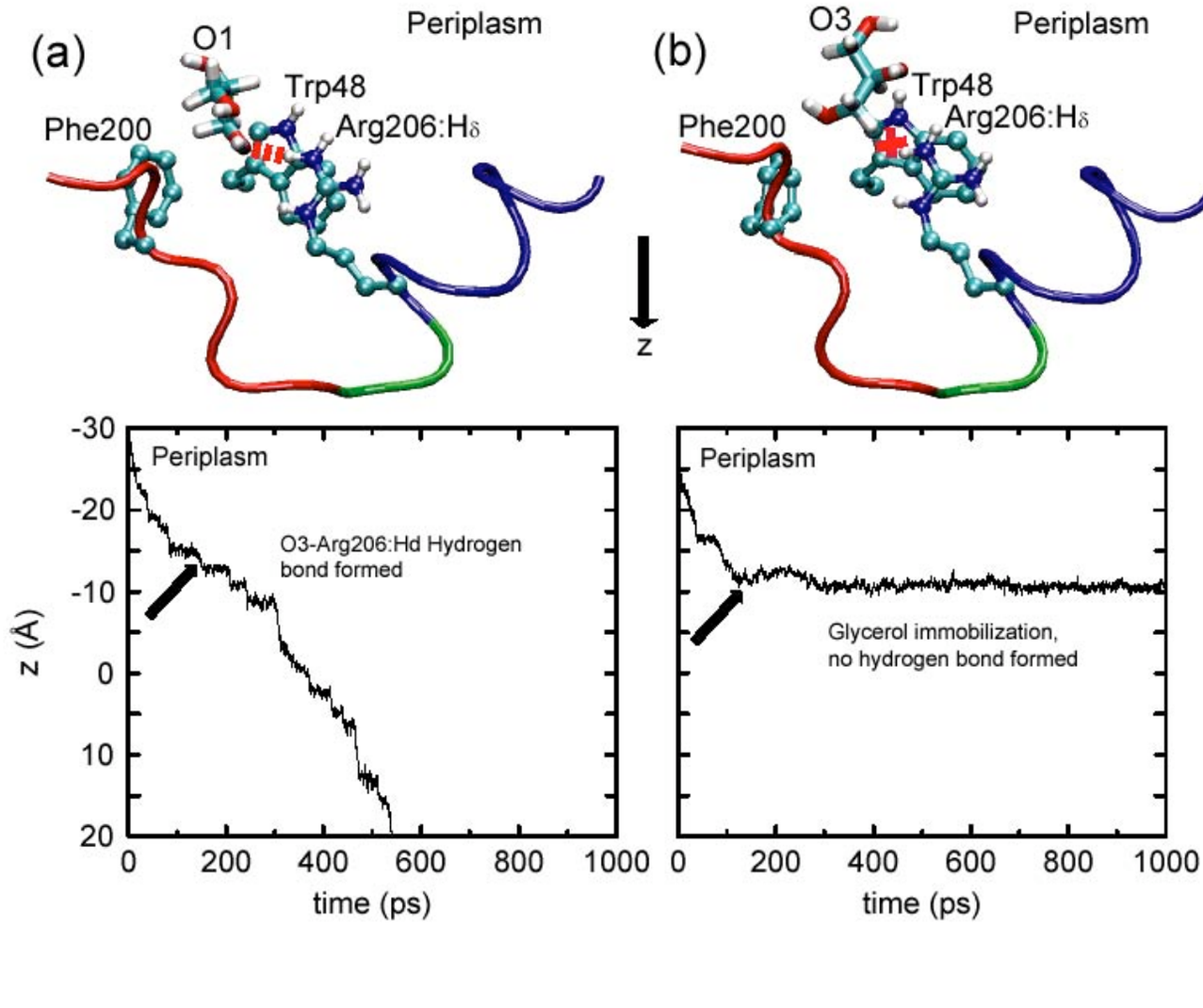


SMD Simulation of Glycerol Passage



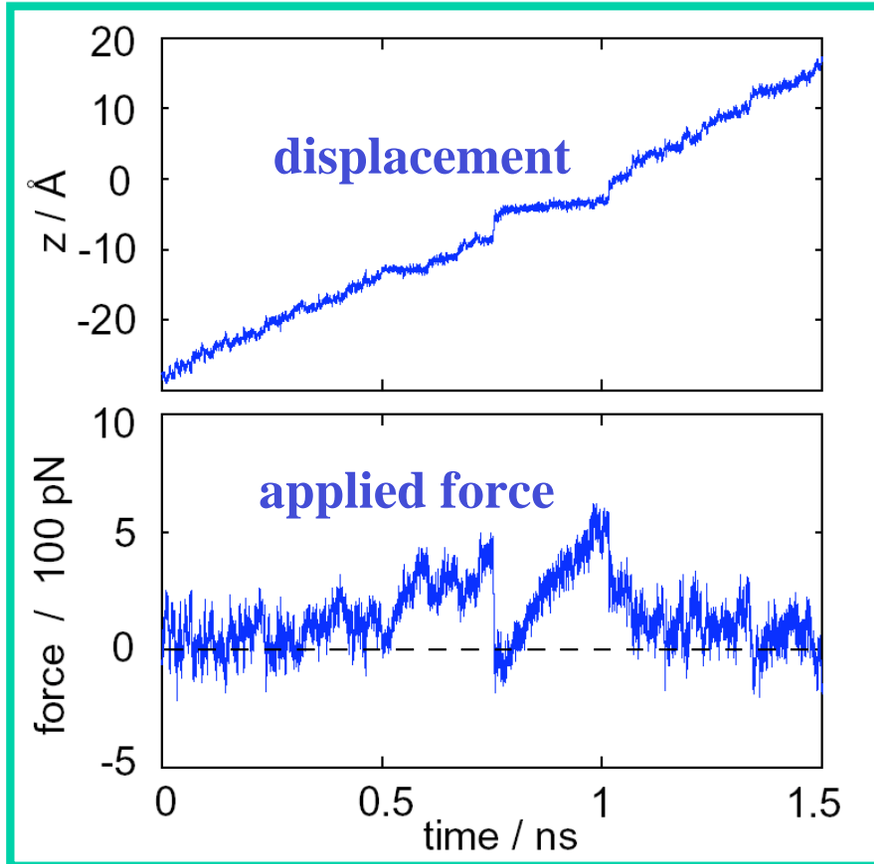
Trajectory of glycerol pulled by **constant force**

Evidence for **Stereoselectivity** of Glycerol



Cannot be verified by experimental measurements

Free Energy Calculation in SMD



Free energy

SMD simulation
a **non-equilibrium** process

$$\Delta G \leq \langle W \rangle$$

**One needs to discount
irreversible work**

$$e^{-\Delta G / k_B T} = \left\langle e^{-W / k_B T} \right\rangle$$

Jarzynski, *PRL* 1997

Hummer, *PNAS*, *JCP* 2001

Liphardt, et al., *Science* 2002

Constructing the Potential of Mean Force

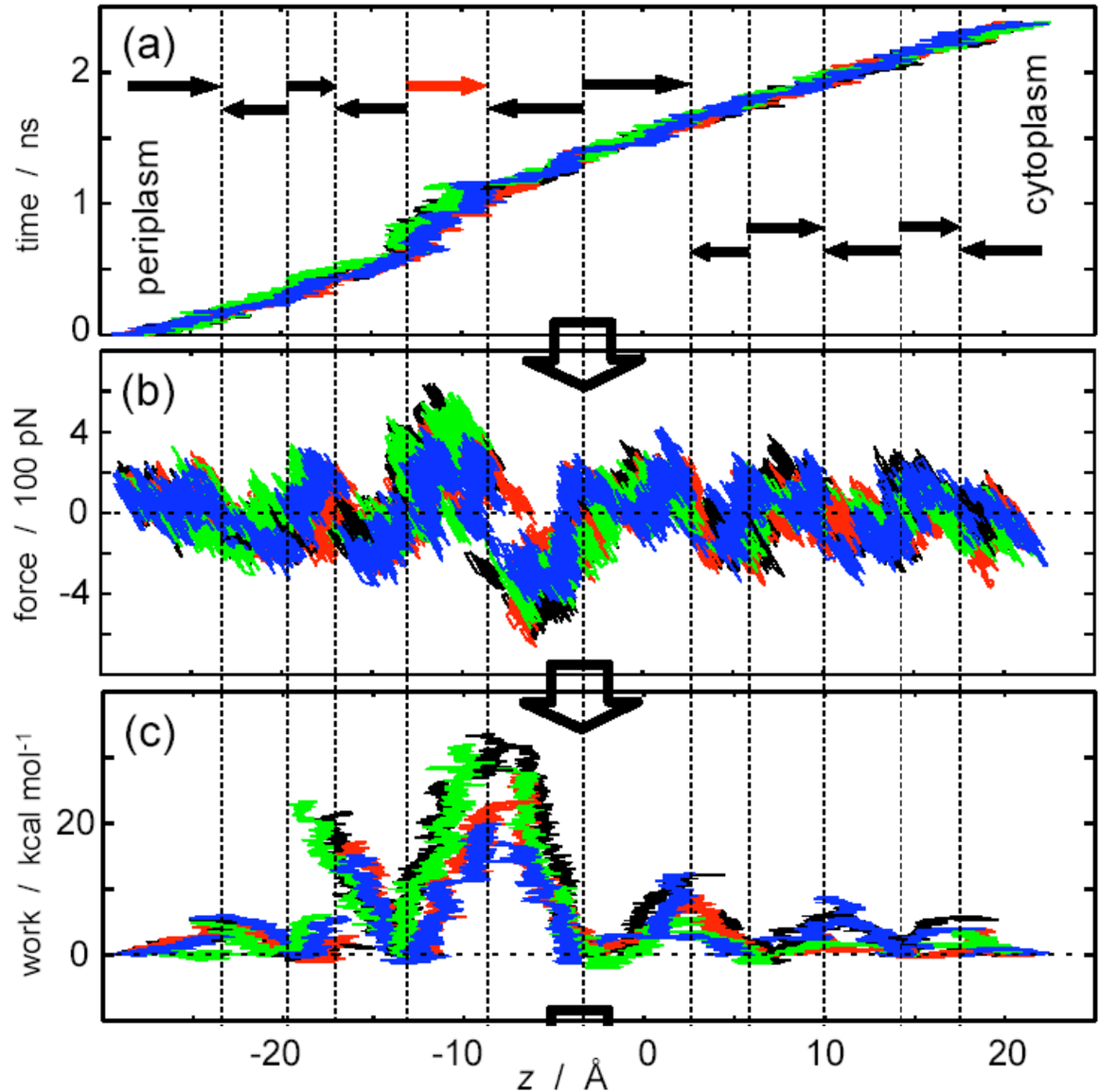
4 trajectories

$v = 0.03, 0.015 \text{ \AA/ps}$

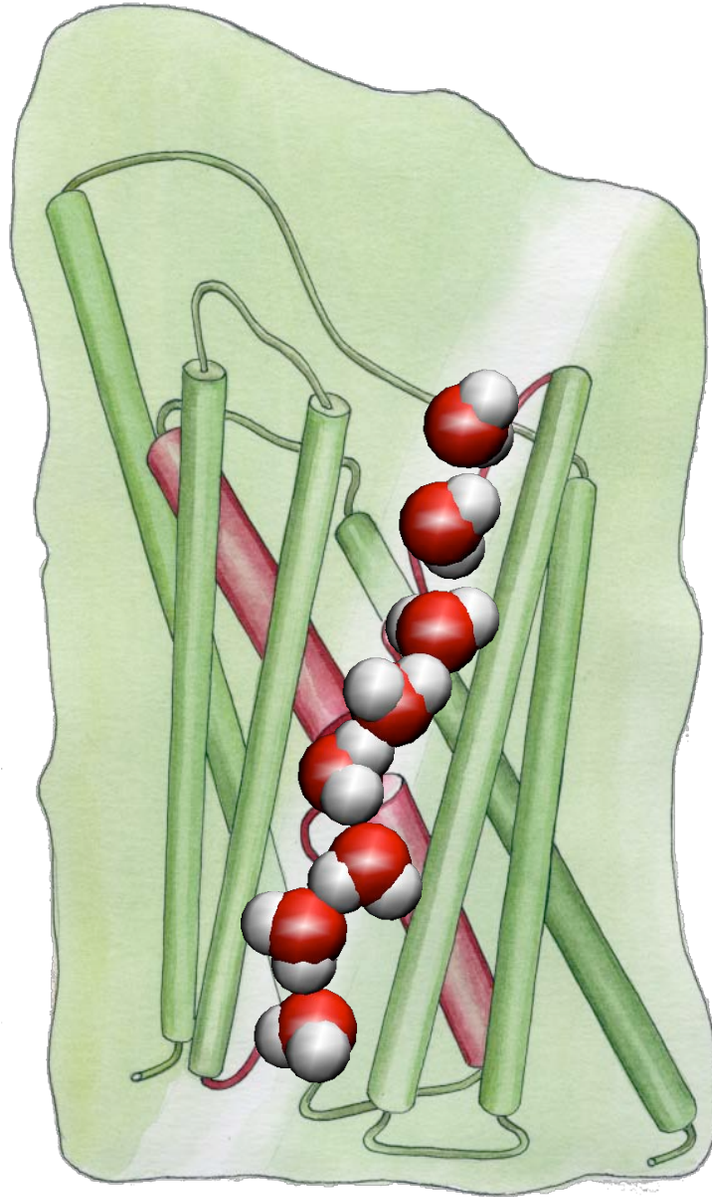
$k = 150 \text{ pN/\AA}$

$$f(t) = -k[z(t) - z_0 - vt]$$

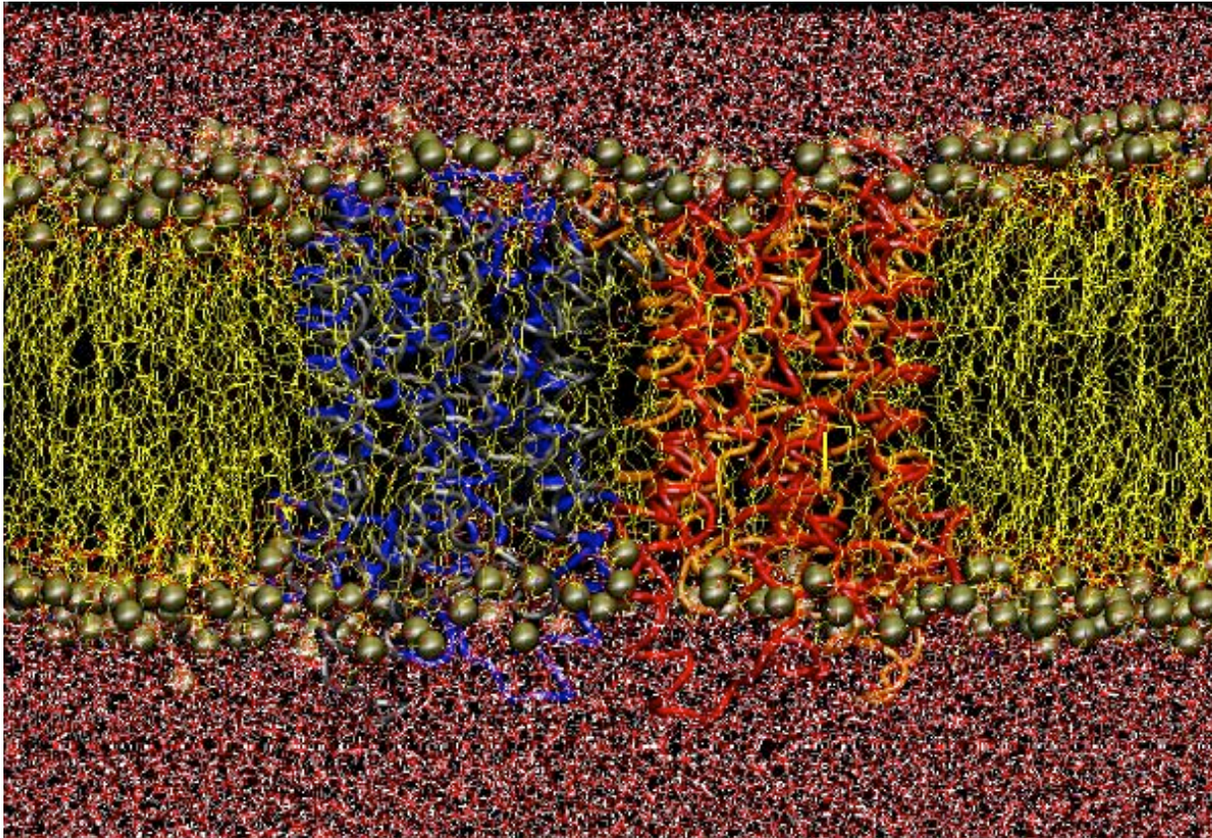
$$W(t) = \int_0^t dt' v f(t')$$



Glycerol-Free GlpF

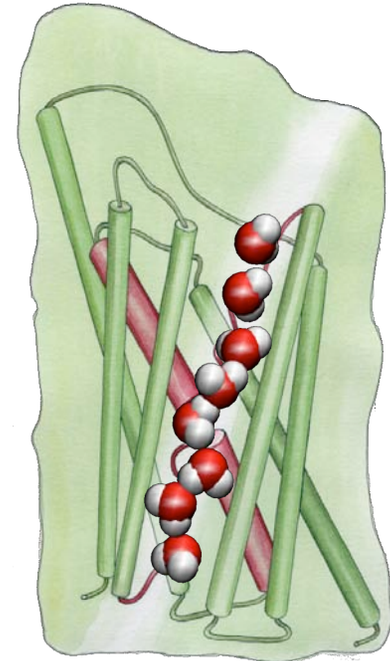


Water permeation

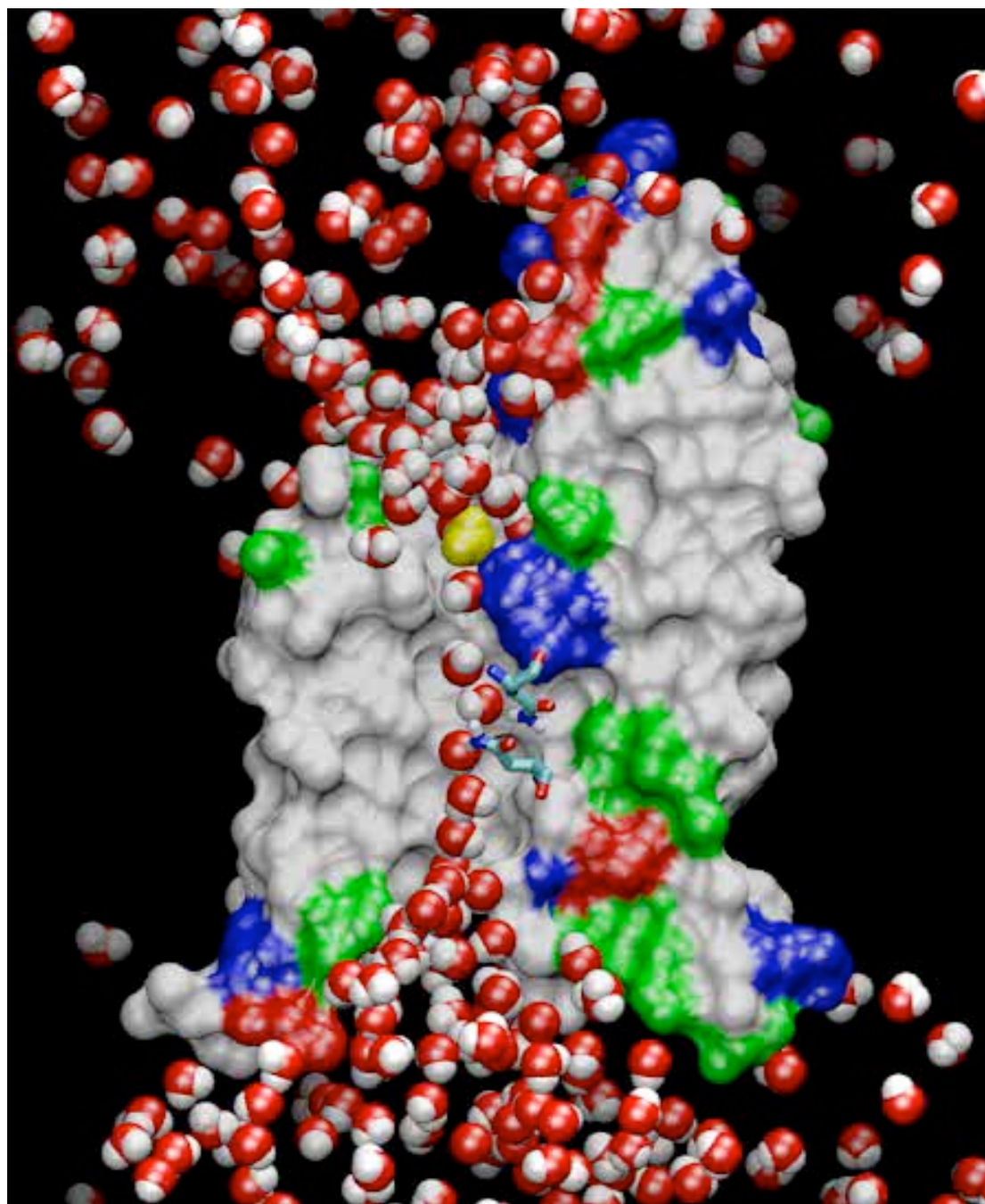


18 water conducted
In 4 monomers in 4 ns
1.125 water/monomer/ns
Exp. = $\sim 1-2$ /ns

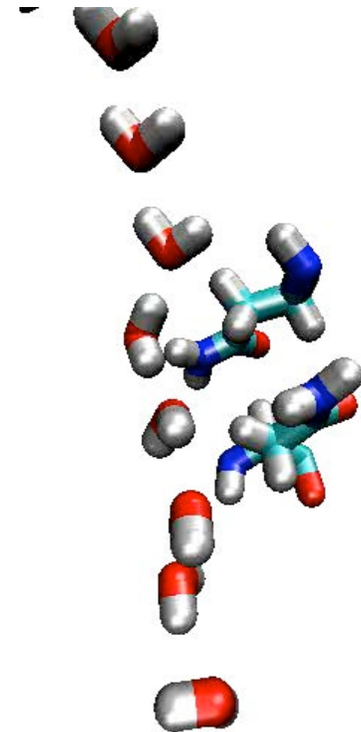
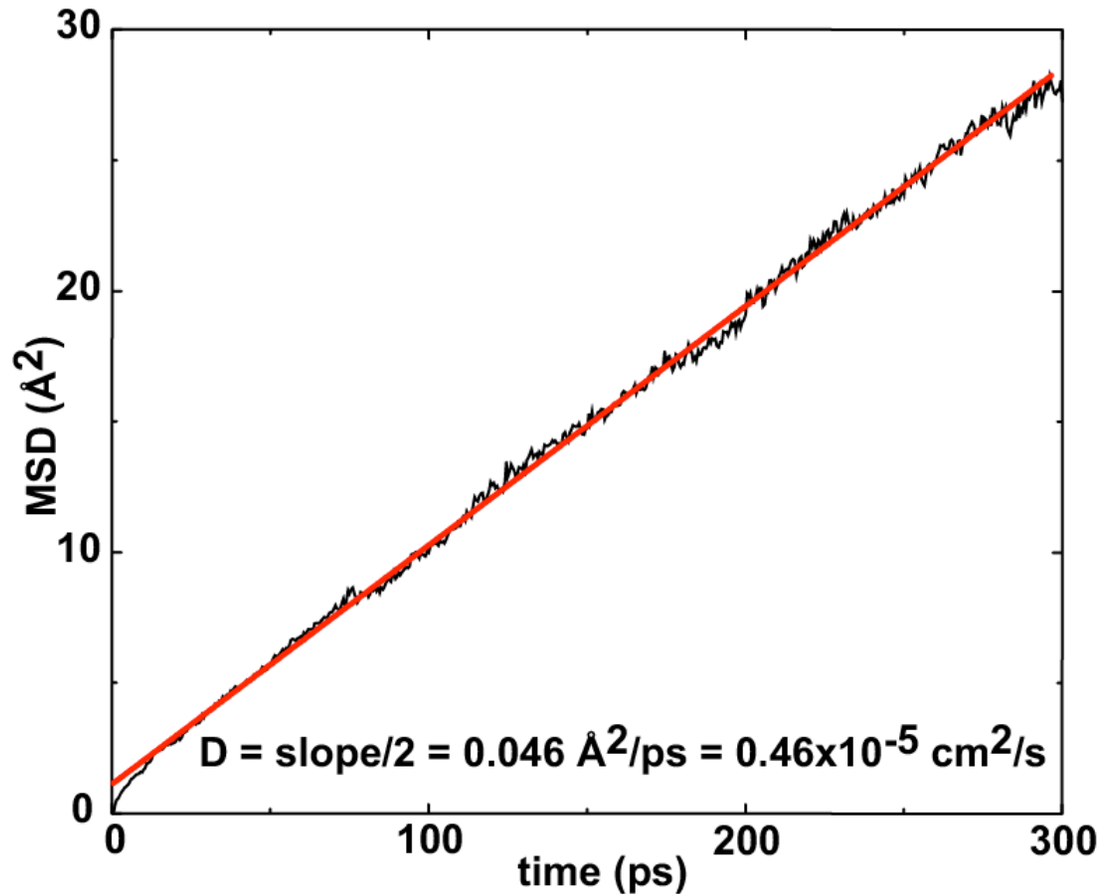
5 nanosecond
Simulation



7-8 water
molecules in each
channel



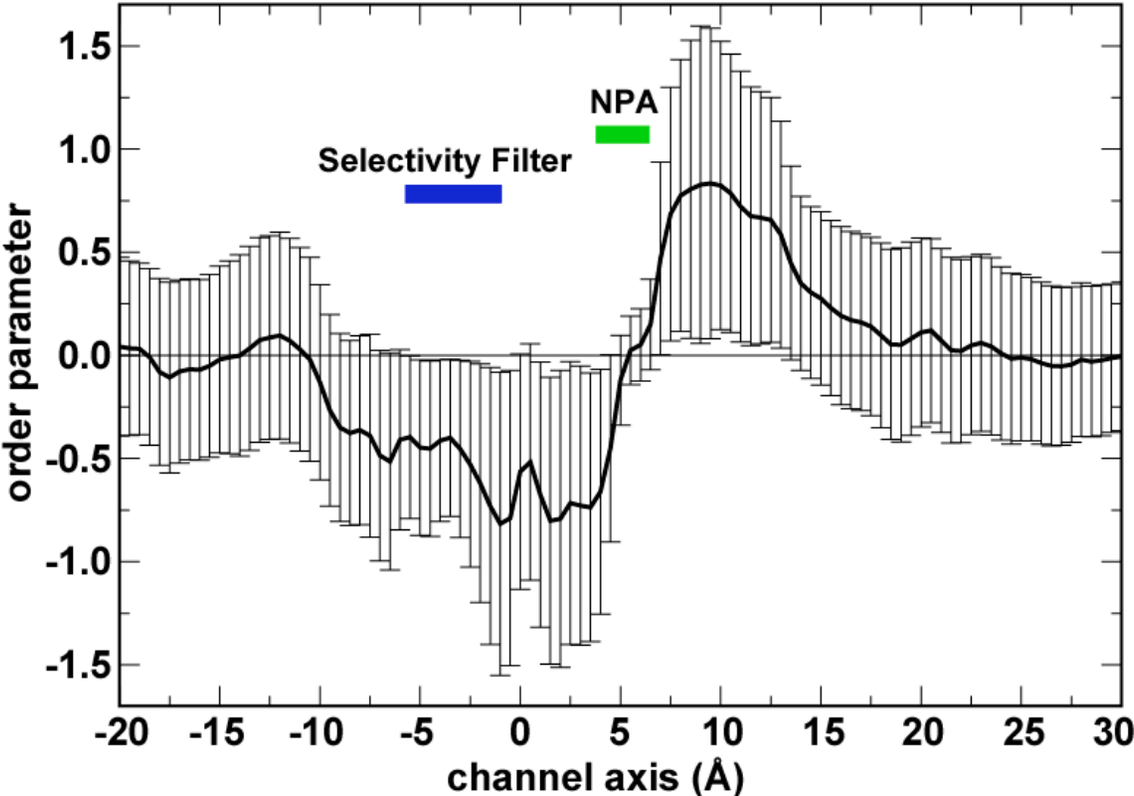
Diffusion of Water in the channel



One dimensional diffusion: $2Dt = \langle (z_t - z_0)^2 \rangle$

Experimental value for AQP1: $0.4-0.8 \text{ e-}5$

channel region (20 Å)

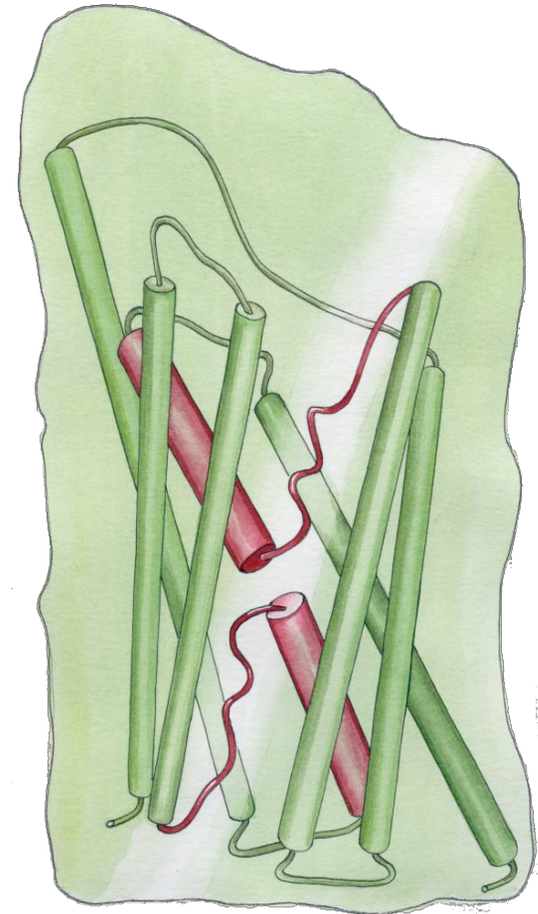
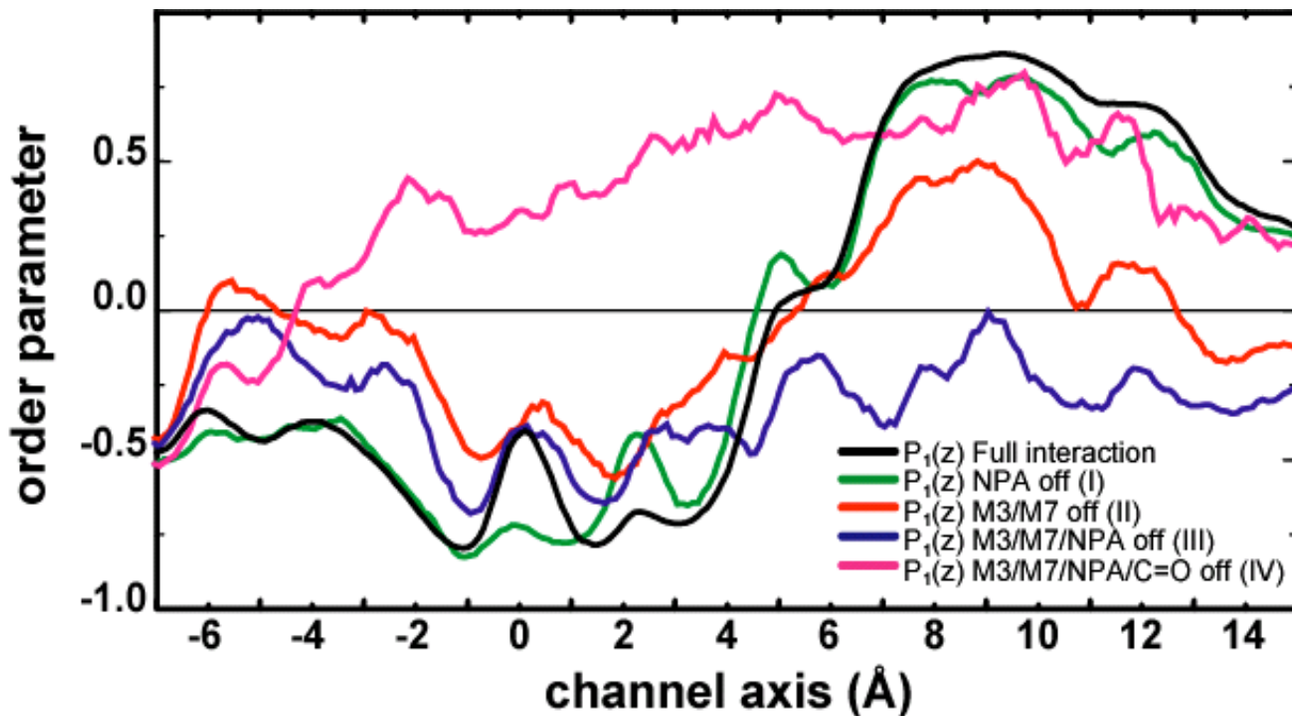


R E M E M B E R:

One of the most useful advantages of simulations over experiments is that you can modify the system as you wish: You can do modifications that are not even possible at all in reality!

This is a powerful technique to test hypotheses developed during your simulations. **Use it!**

Electrostatic Stabilization of Water Bipolar Arrangement



Proton transfer through water

