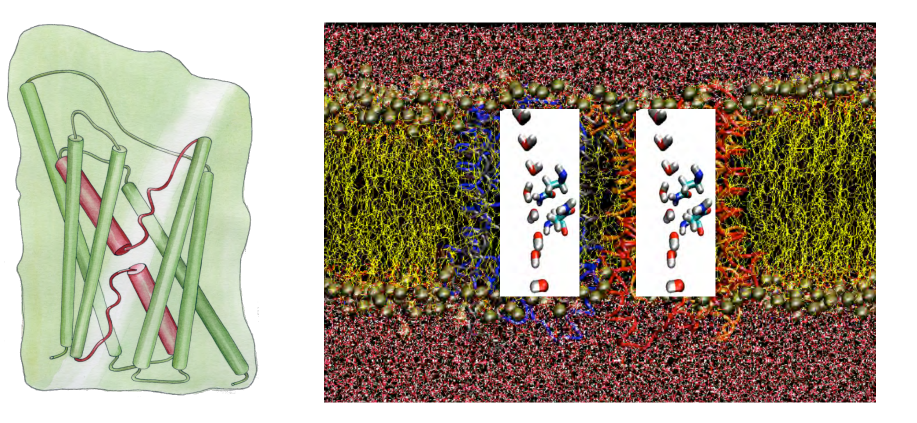
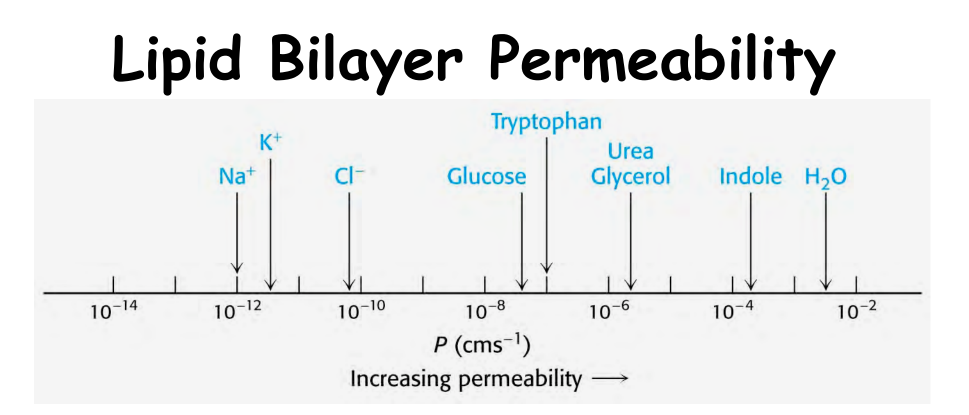
Analysis of Molecular Dynamics Simulations of Biomolecules

- 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





Water is an exception:

- •Small size
- ·Lack of charge
- Its high concentration

Water Transport Across Cell Membrane <u>Always passive; bidirectional; osmosis-driven</u>

Diffusion through lipid bilayers

slower, but enough for many purposes

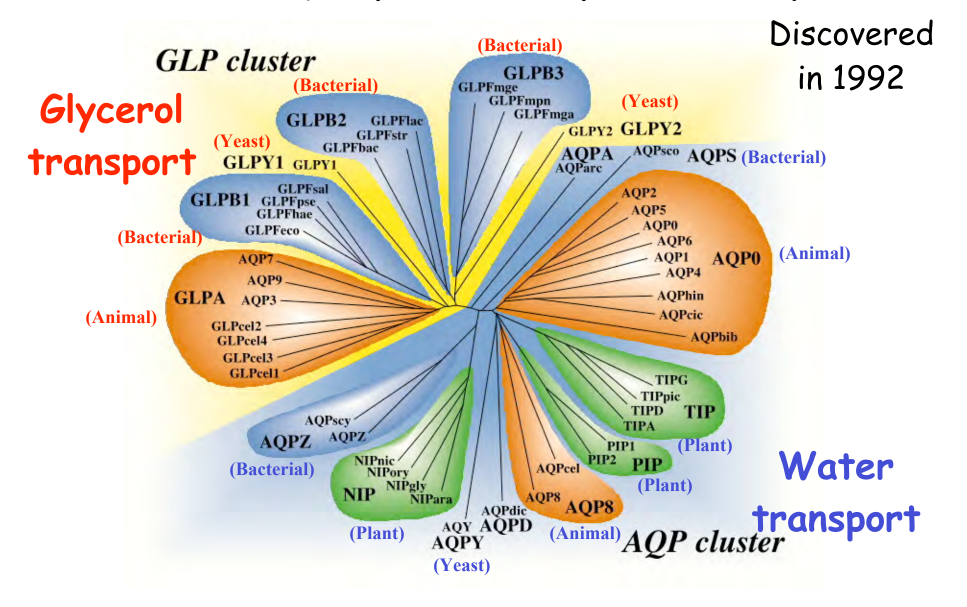
Channel-mediated

Large volumes of water needed to be transported (kidneys).

Fast adjustment of water concentration is necessary

(RBC, brain, lung).

The Aquaporin Superfamily

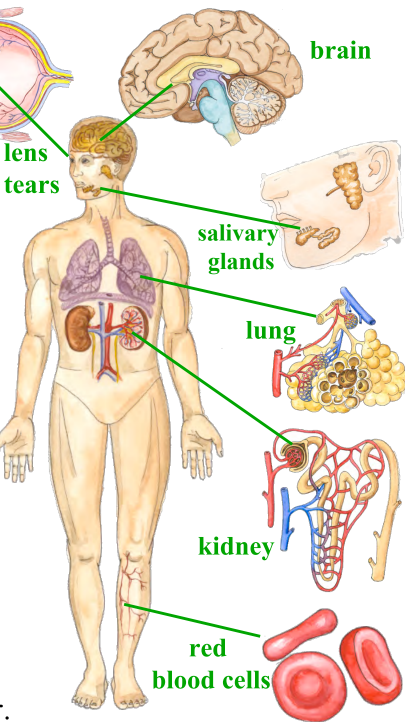


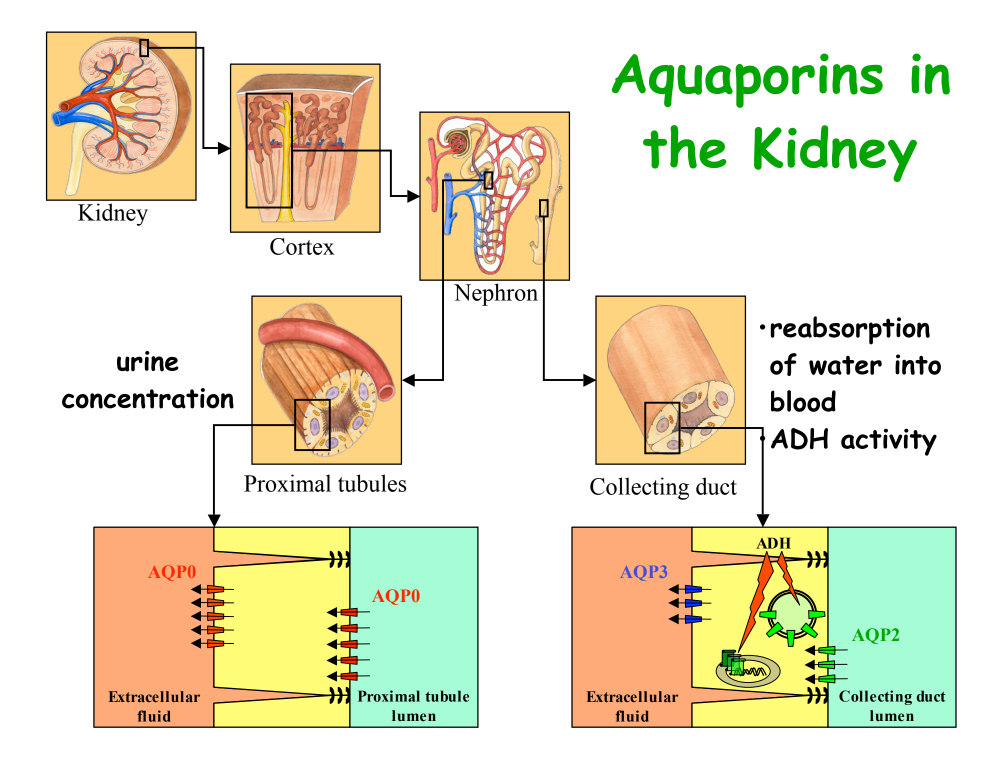
Heymann and Engel News Physiol. Sci. 14, 187 (1999)

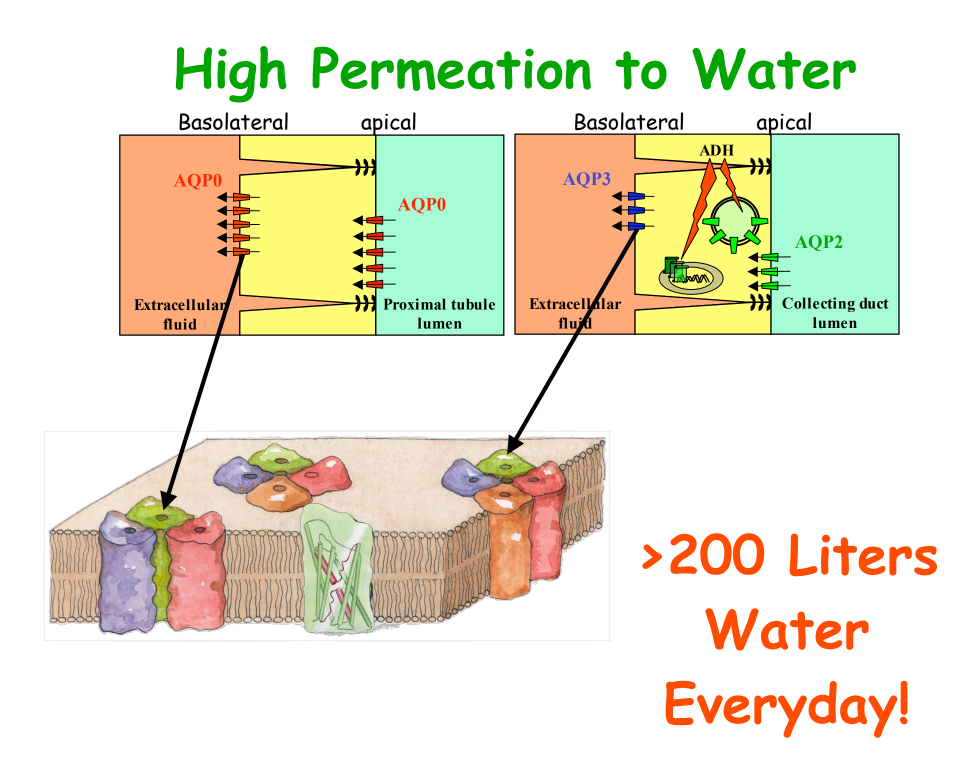
Aquaporins in Human Body

Aquaporin-o	Eye: lens fiber cells	Fluid balance of the
Aquaporin-1	Red blood cells Kidney: proximal tubules Eye: ciliary epithelium Brain: choriod plexus	Osmotic protection Concentration of urine Aqueous humor Production of CSF Alveolar hydration
	Lung: alveolar	, i i i i i i i i i i i i i i i i i i i
Aquaporin-2	epithelial cells. Kidney: collecting ducts	ADH hormone activity
Aquaporin-3	Kidney: collecting ducts Trachea: epithelial cells	Reabsorption of water Secretion of water
Aquaporin-4	Kidney: collecting ducts Brain: ependymal cells Brain: hypothalamus Lung: bronchial	Reabsorption of water CSF fluid balance Osmosensing function?
Aquaporin-5	epithelium Salivary glands Lacrimal glands	Bronchial fluid Production of saliva secretion Production of tears
Aquaporin-6	Kidney	Very low water permeability!
Aquaporin-7	Testis and sperm	
Aquaporin-8	Testis, pancreas, liver	
Aquaporin-9	Leukocytes	
Aquaporin-10		

Additional members are suspected to exist.



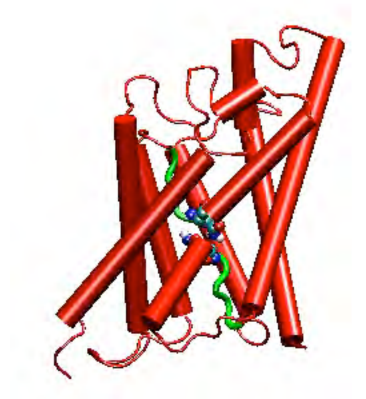


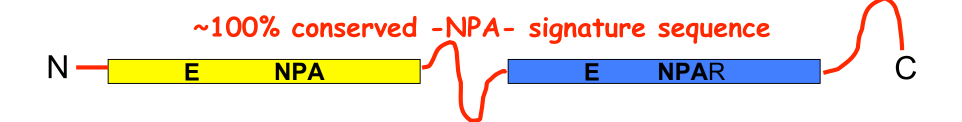


Tetrameric pore Monomeric pores Perhaps ions??? Water, glycerol, ... Aquaporins of known structure: GlpF - E. coli glycerol channel (aquaglycerolporin) AQP1 - Mammalian aquaporin-1 (pure water channel) AgpZ and AQPO (2004)

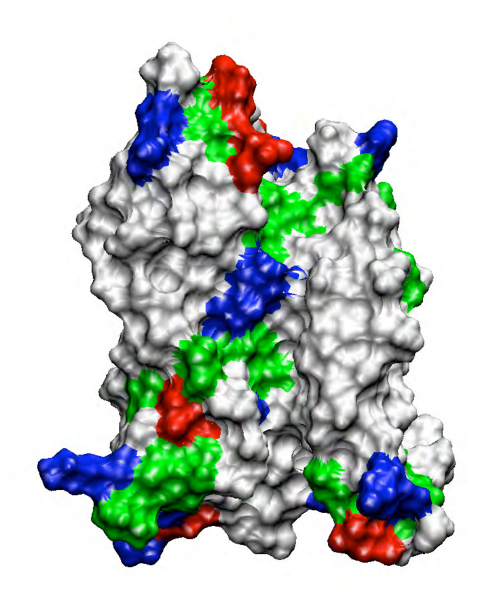
Functionally Important Features

- Tetrameric architecture
- Amphipatic channel interior
- Water and glycerol transport
- Protons, and other ions are excluded
- Conserved asparagine-prolinealanine residues; NPA motif
- Characteristic half-membrane spanning structure





A Semi-hydrophobic channel

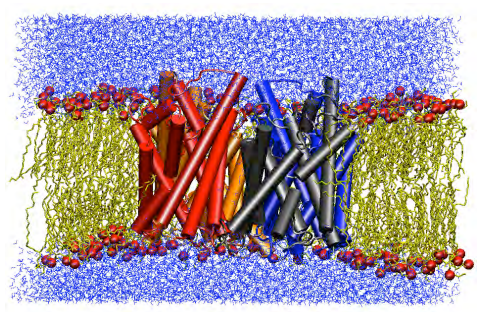


Molecular Dynamics Simulations

Protein: Lipids (POPE) Water: Total:

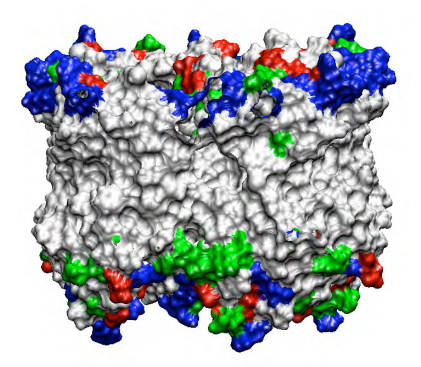
- ~ 15,000 atoms
- Lipids (POPE): ~ 40,000 atoms
 - ~ 51,000 atoms

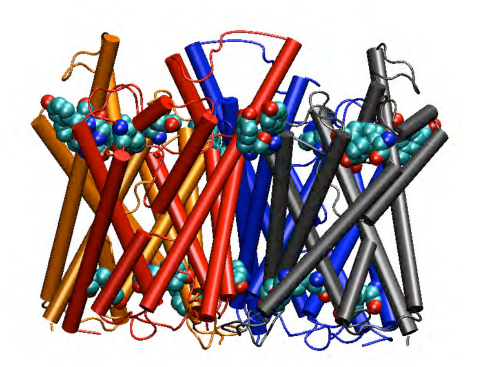
~ 106,000 atoms



NAMD, CHARMM27, PME NpT ensemble at 310 K Ins equilibration, 4ns production 10 days /ns - 32-proc Linux cluster 3.5 days/ns - 128 O2000 CPUs 0.35 days/ns - 512 LeMieux CPUs

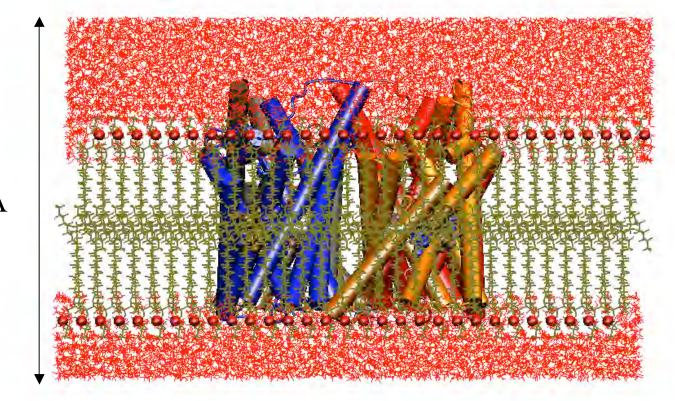
Protein Embedding in Membrane





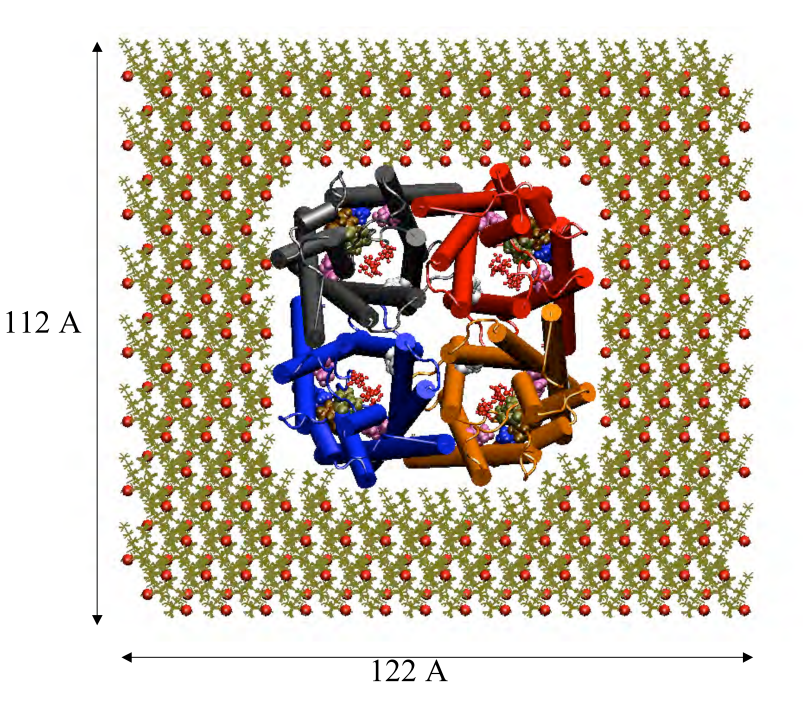
Hydrophobic surface of the protein Ring of Tyr and Trp

Embedding GlpF in Membrane



77 A

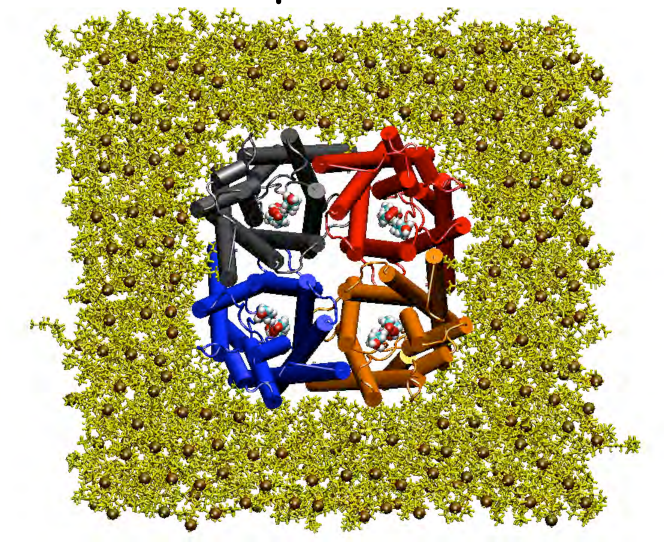
GlpF in VMD



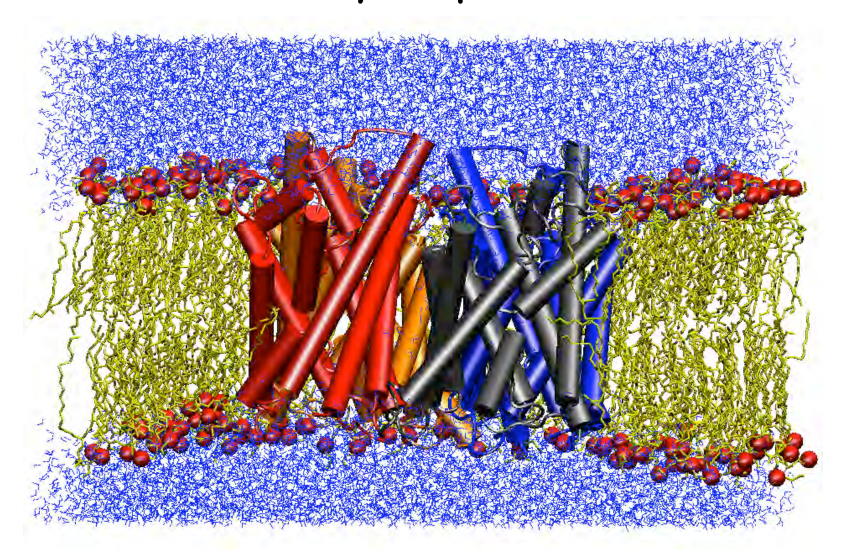
A Recipe for Membrane Protein Simulations

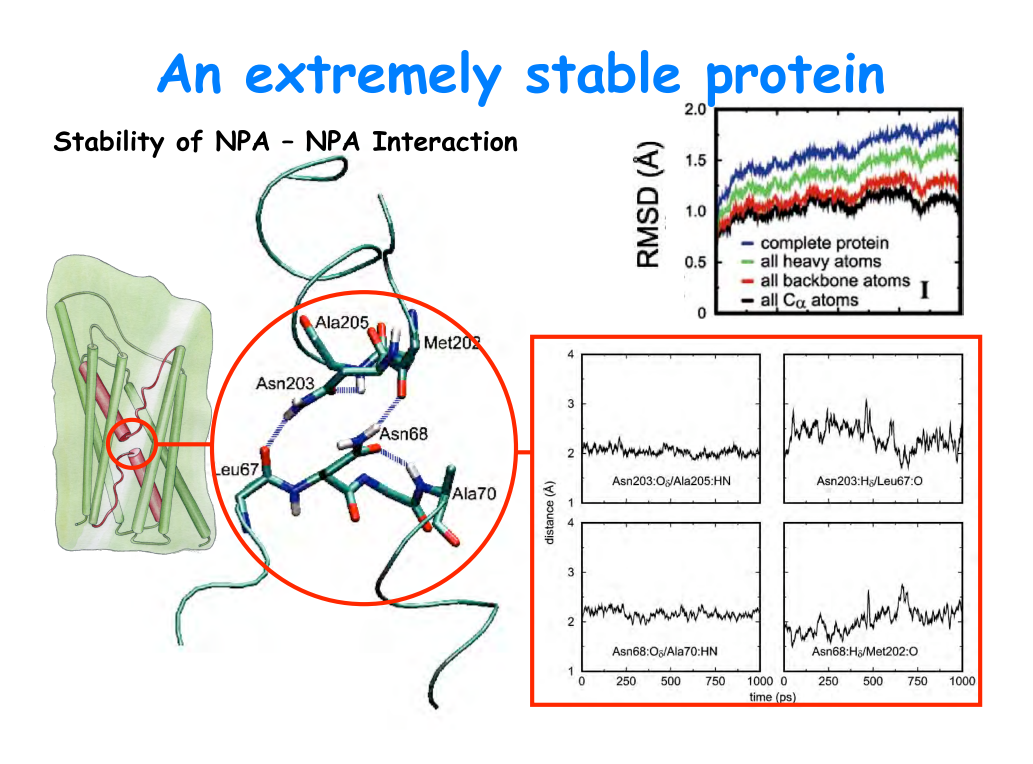
- Insert your protein into a hydrated lipid bilayer.
- Fix the protein; minimize the rest and run a short "constantpressure" MD to bring lipids closer to the protein and fill the gap between the protein and lipids.
- Watch water molecules; if necessary apply constraints to prevent them from penetrating 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

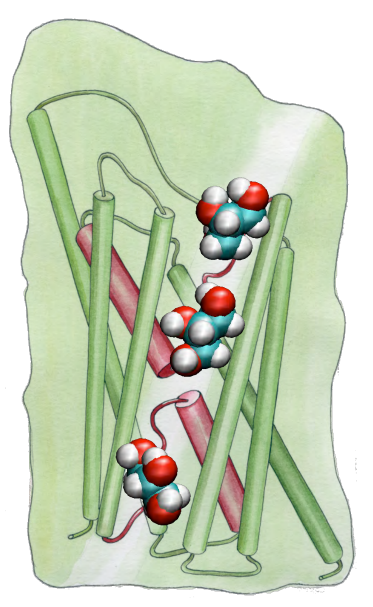


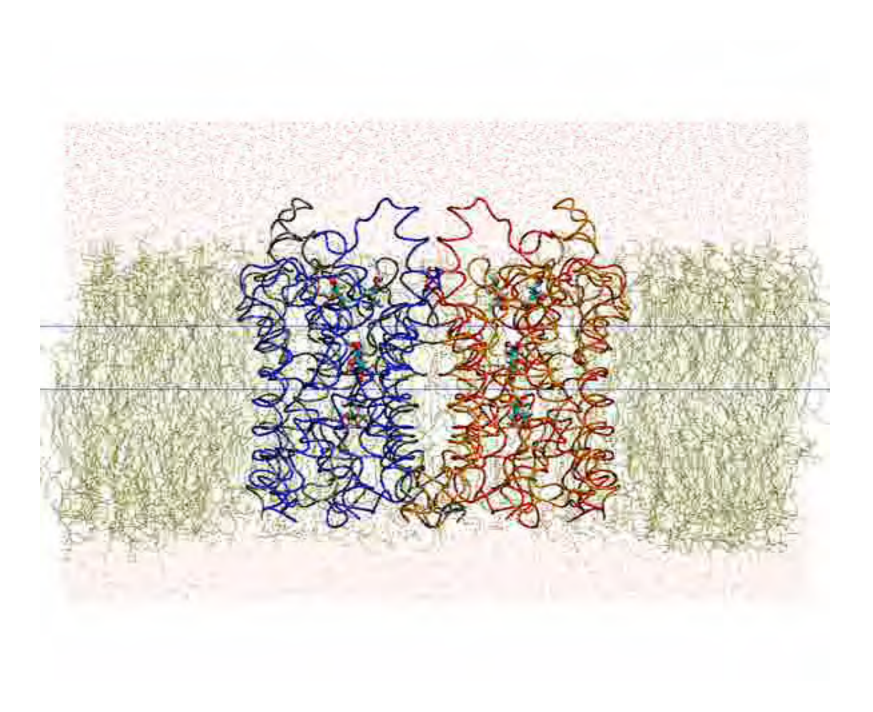
Adjustment of Membrane Thickness to the Protein Hydrophobic Surface



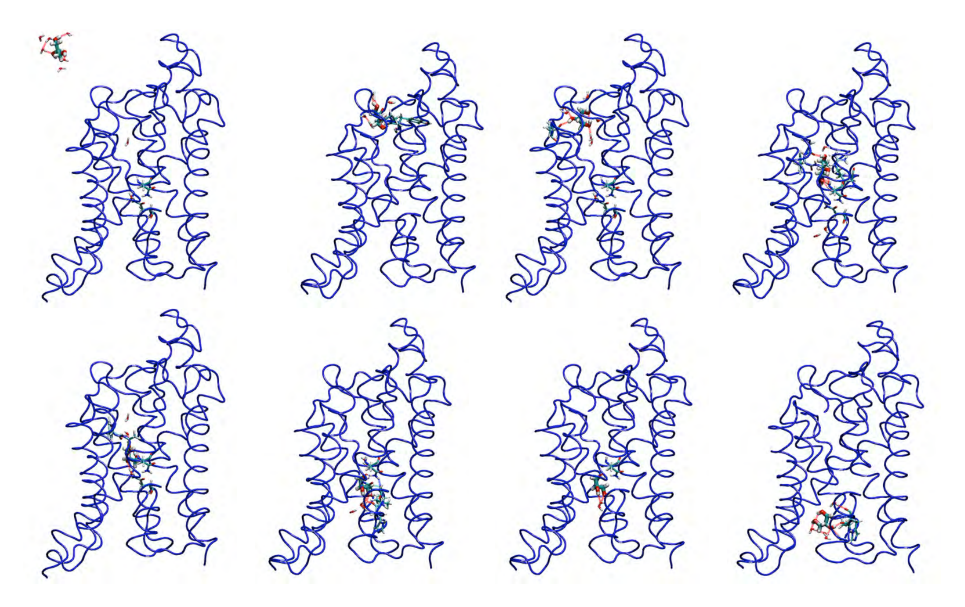


Glycerol-Saturated GlpF





Description of full conduction pathway



Complete description of the conduction pathway



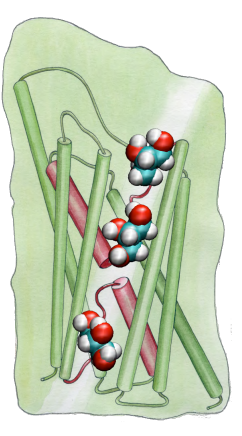
Details of Protein-Substrate Interaction Are Always Important

- 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

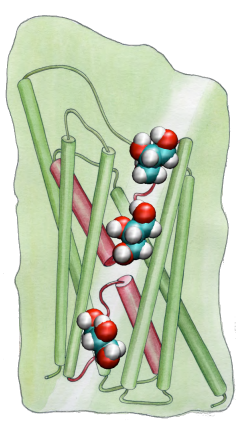
```
{set frame 0}{frame < 100}{incr frame}{
animate goto $frame
set donor [atomselect top
"name 0 N and within 2 of
(resname GCL and name HO)"]
lappend [$donor get index] list1
set acceptor [atomselect top
"resname GCL and name 0 and
within 2 of (protein and name HN HO)"]
lappend [$acceptor get index]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

. . .



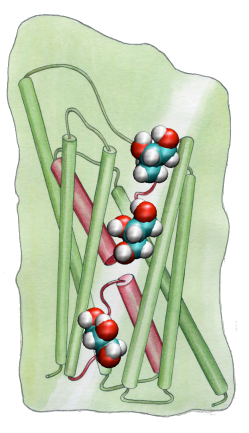
Channel Hydrogen Bonding Sites

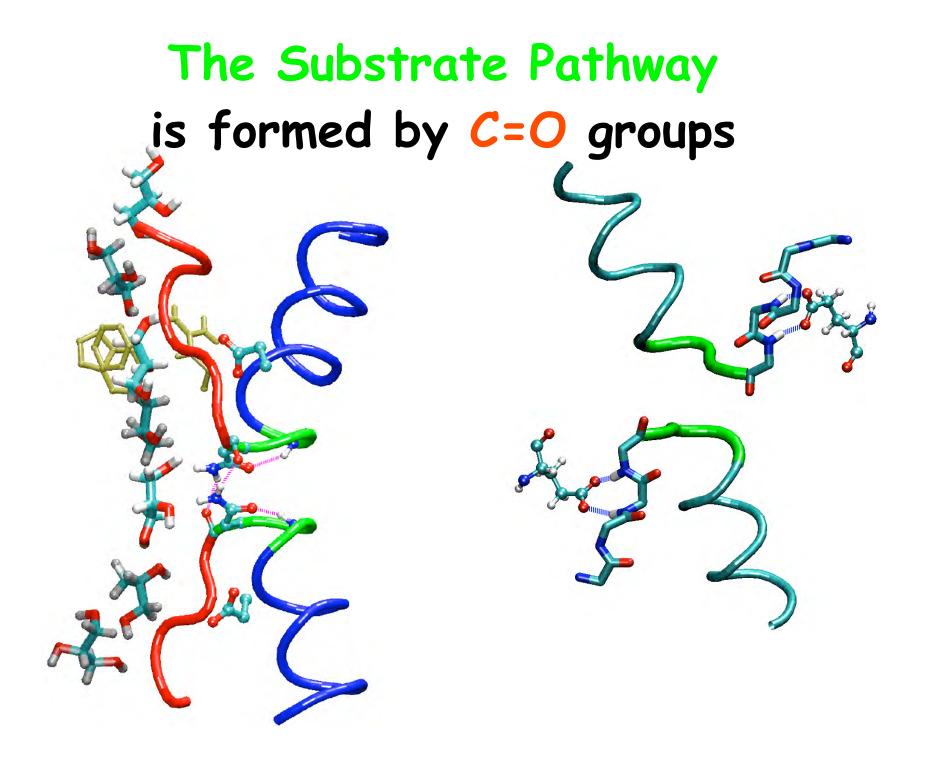
GLN	41	OE1 NE2	LEU	197	0
TRP	48	O NE1	THR	198	0
GLY	64	0	GLY	199	0
ALA	65	0	PHE	200	0
HIS	66	O ND1	ALA	201	0
LEU	67	0	ASN	203	ND2
ASN	68	ND2			
ASP	130	OD1	LYS	33	HZ1 HZ3
GLY	133	0	GLN	41	HE21
SER	136	0	TRP	48	HE1
TYR	138	0	HIS	66	HD1
PRO	139	ΟΝ	<u>ASN</u>	68	HD22
ASN	140	OD1 ND2	TYR	138	HN
HIS	142	ND1	ASN	140	HN HD21 HD22
THR	167	OG1	HIS	142	HD1
GLY	195	0	GLY	199	HN
PRO	196	0	<u>ASN</u>	203	HN HD21HD22
			<u>ARG</u>	206	HE HH21HH22



Channel Hydrogen Bonding Sites

GLN	41	OE1 NE2	LEU	197	0
TRP	48	O NE1	THR	198	0
GLY	64	0	GLY	199	0
ALA	65	0	PHE	200	0
HIS	66	O ND1	ALA	201	0
LEU	67	0	ASN	203	ND2
ASN	68	ND2			
ASP	130	OD1	LYS	33	HZ1 HZ3
GLY	133	0	GLN	41	HE21
SER	136	0	TRP	48	HE1
TYR	138	0	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	0	GLY	199	HN
PRO	196	0	<u>ASN</u>	203	HN HD21HD22
			<u>ARG</u>	206	HE HH21HH22





The Substrate Pathway is formed by C=O groups

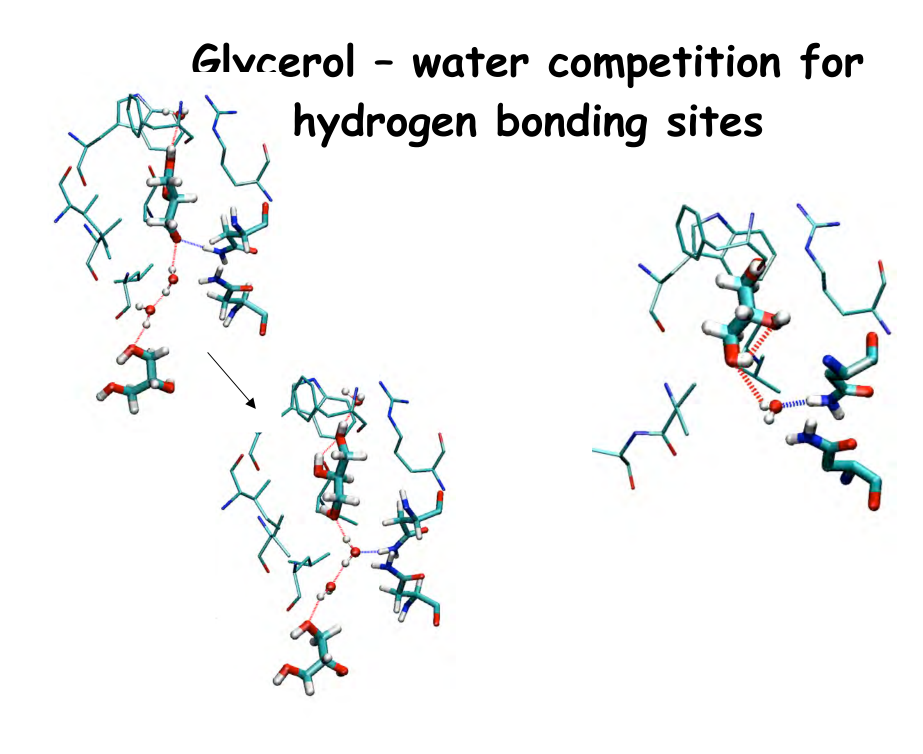
NPAR

Non-helical motifs are stabilized by two glutamate residues.

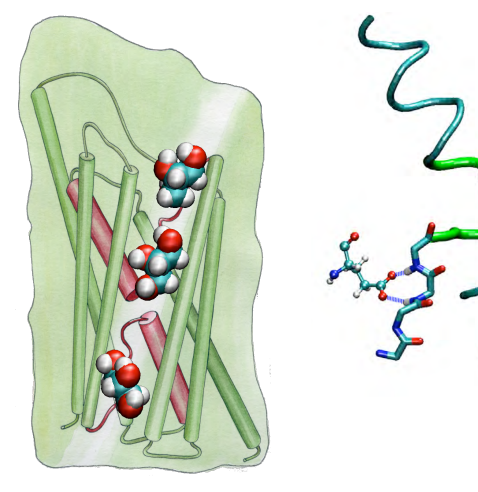
NPA

Ν

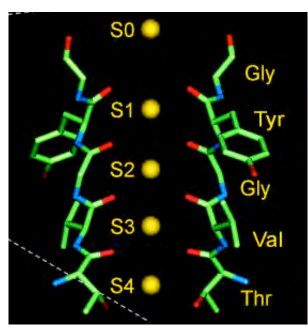
Conservation of Glutamate Residue in Human Aquaporins



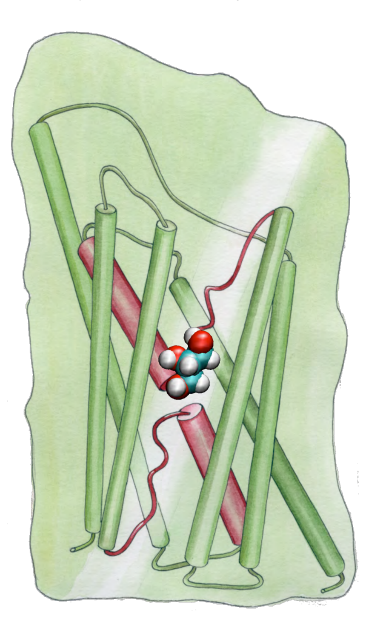
Revealing the Functional Role of Reentrant Loops

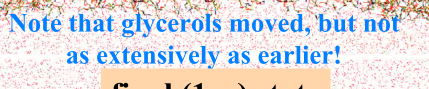


Potassium channel



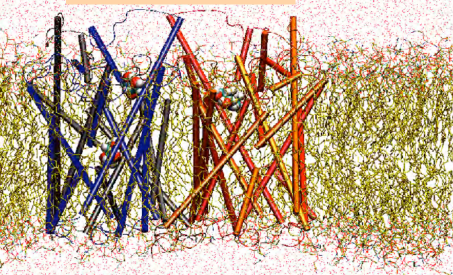
Single Glycerol per channel

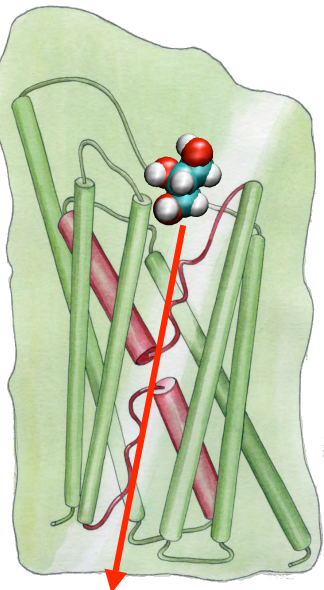




initial state

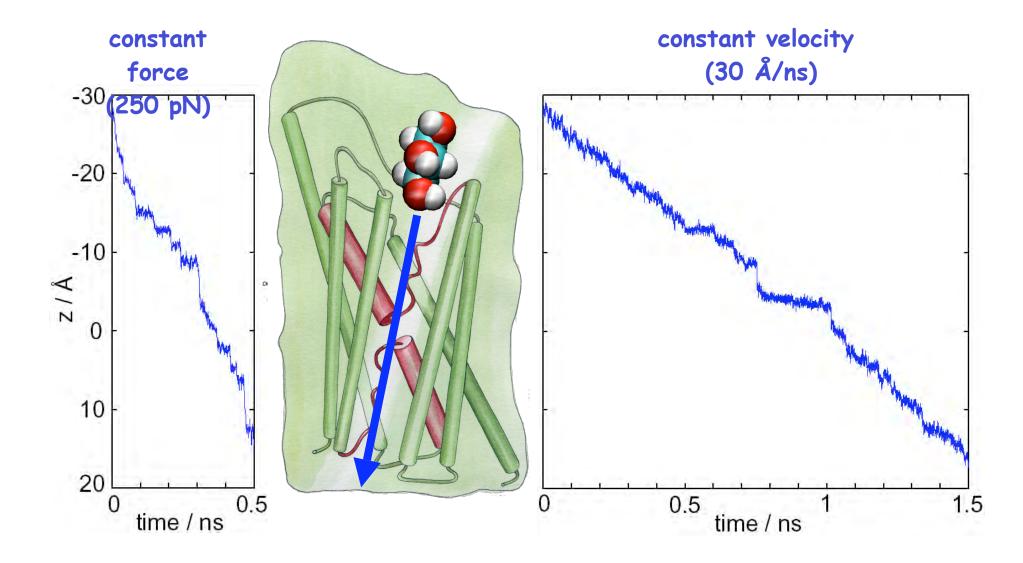
final (1ns) state



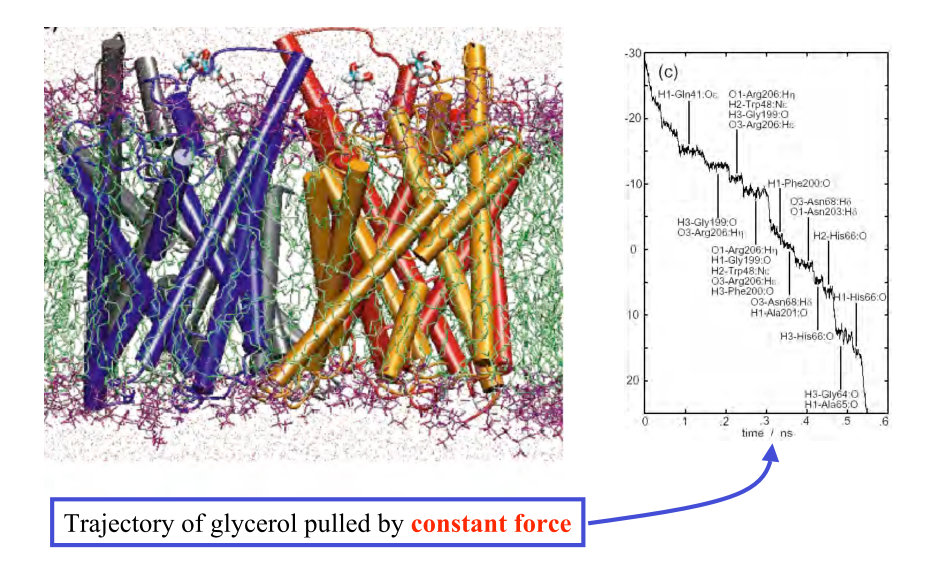


We need to enforce an entire conduction event.

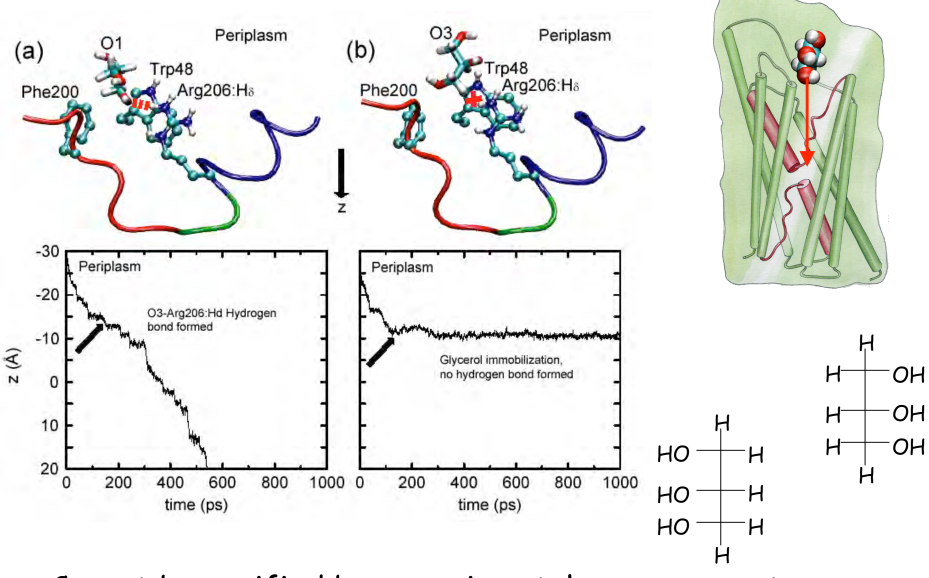
Steered Molecular Dynamics



SMD Simulation of Glycerol Passage

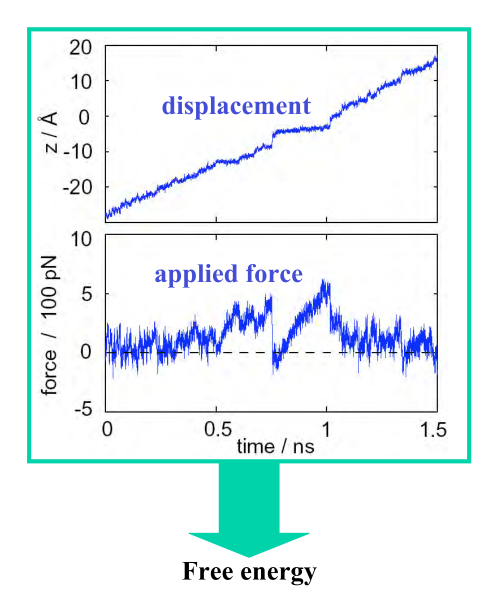


Evidence for Stereoselectivity of Glycerol



Cannot be verified by experimental measurements

Free Energy Calculation in SMD



SMD simulation a non-equilibrium process

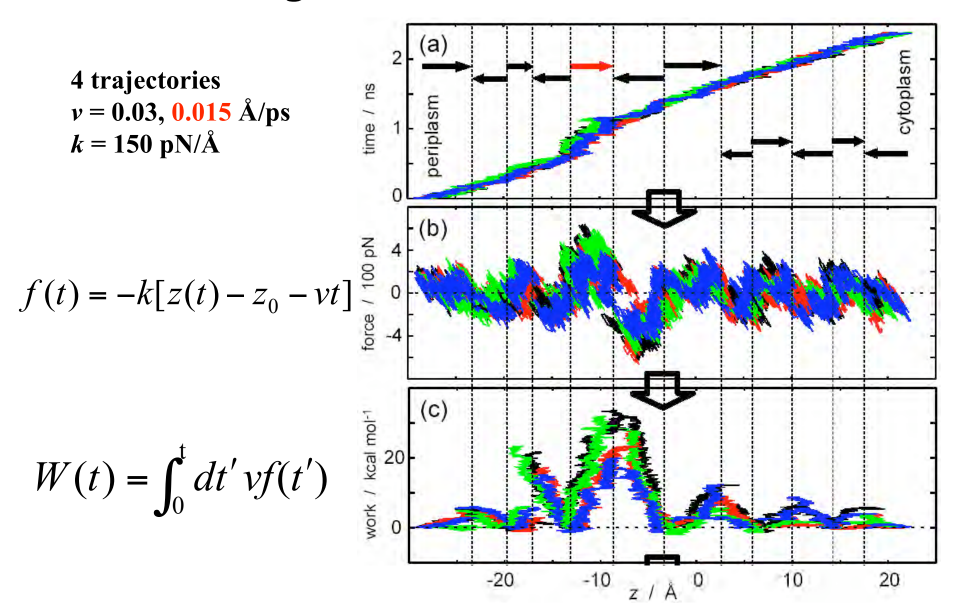
$$\Delta G \leq \left< W \right>$$

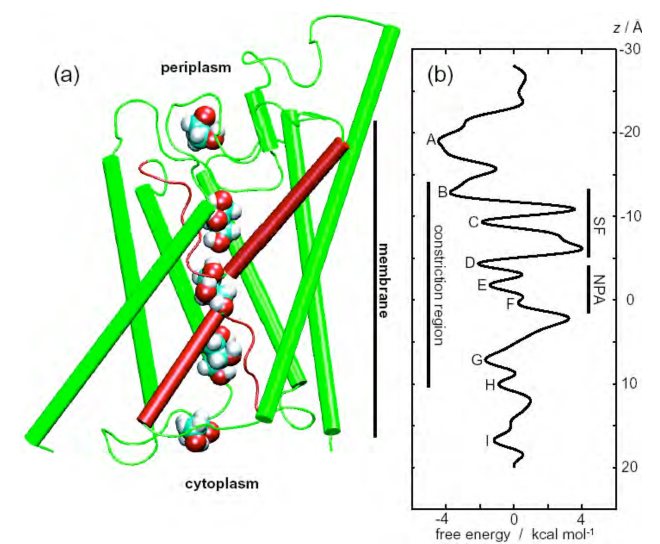
One needs to discount irreversible work

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

Jarzynski, *PRL* 1997 Hummer, *PNAS, JCP* 2001 Liphardt, et al., *Science* 2002

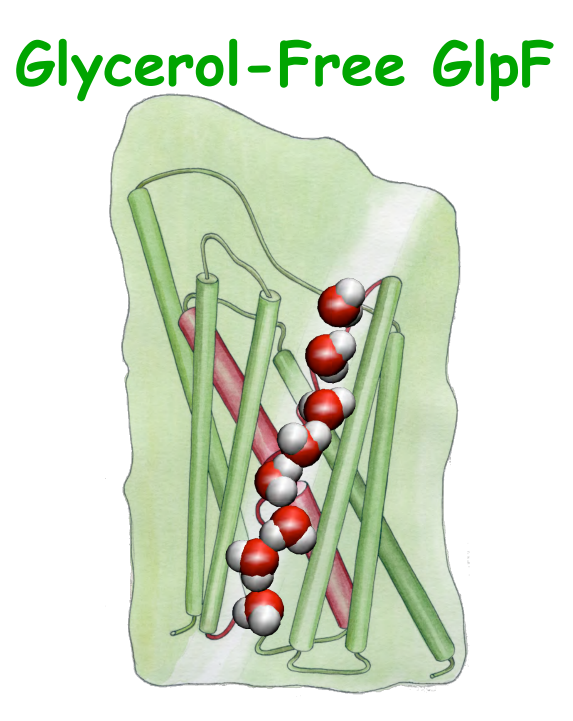
Constructing the Potential of Mean Force



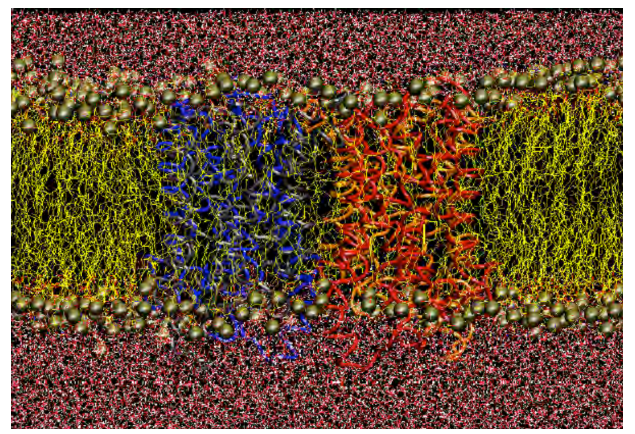


Features of the Potential of Mean Force

- Captures major features of the channel
- The largest barrier ~ 7.3 kcal/mol; exp.: 9.6±1.5 kcal/mol

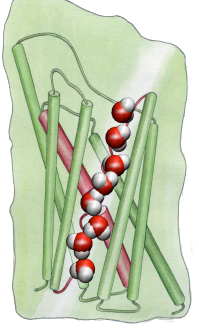


Water permeation

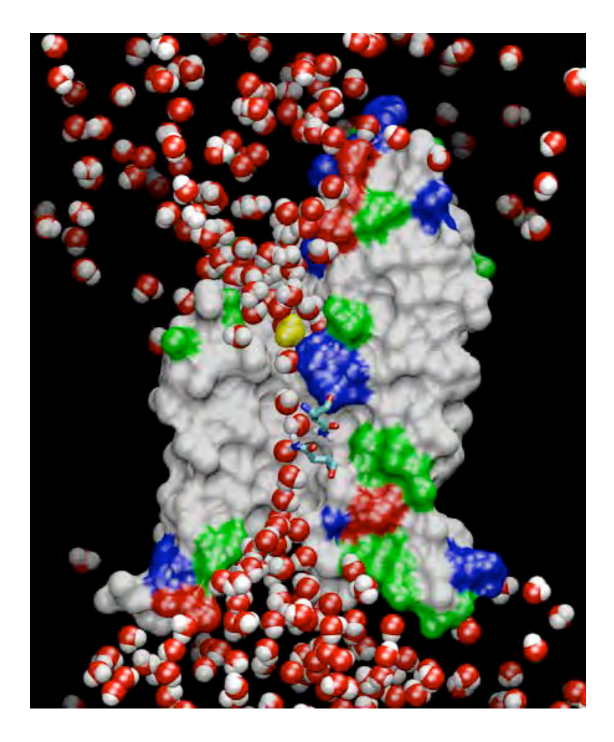


18 water conducted In 4 monomers in 4 ns 1.125 water/monomer/ns Exp. = ~ 1-2 /ns

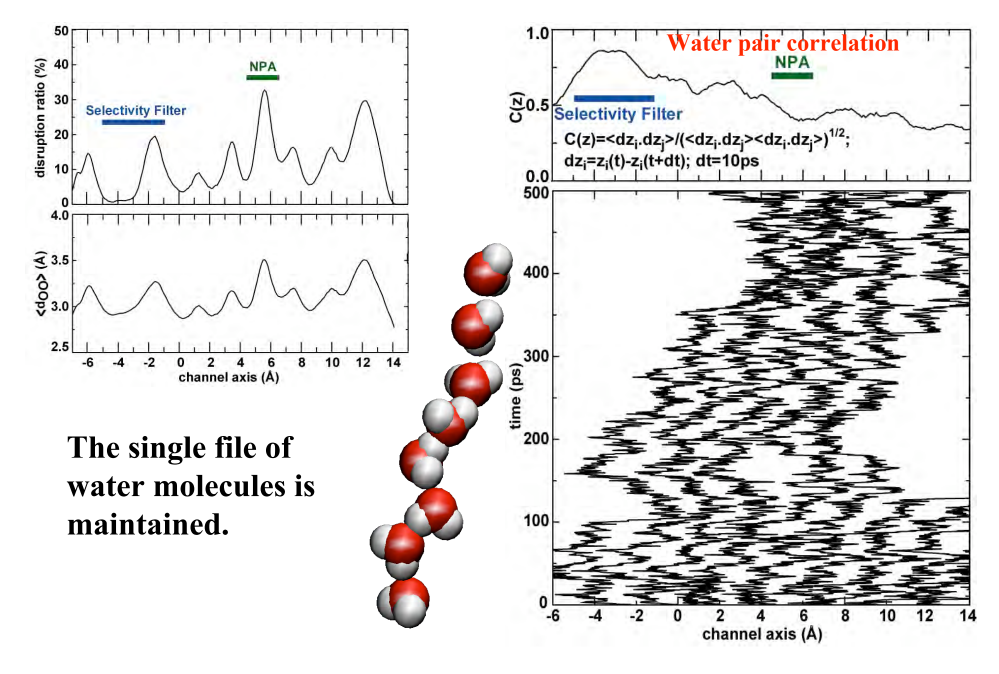
5 nanosecond Simulation



7-8 water molecules in each channel

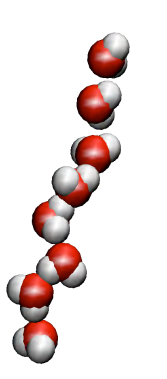


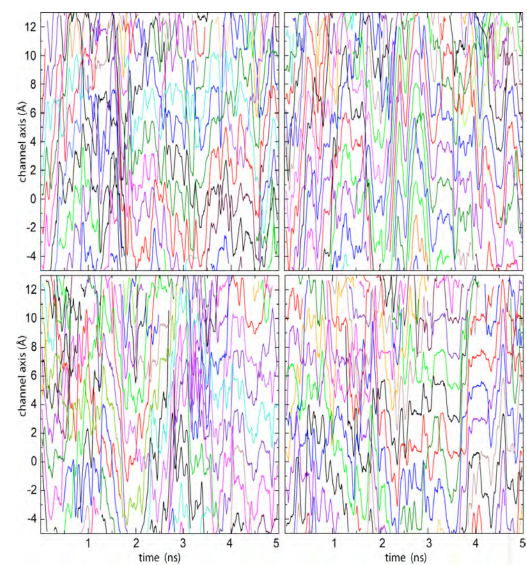
Correlated Motion of Water in the Channel



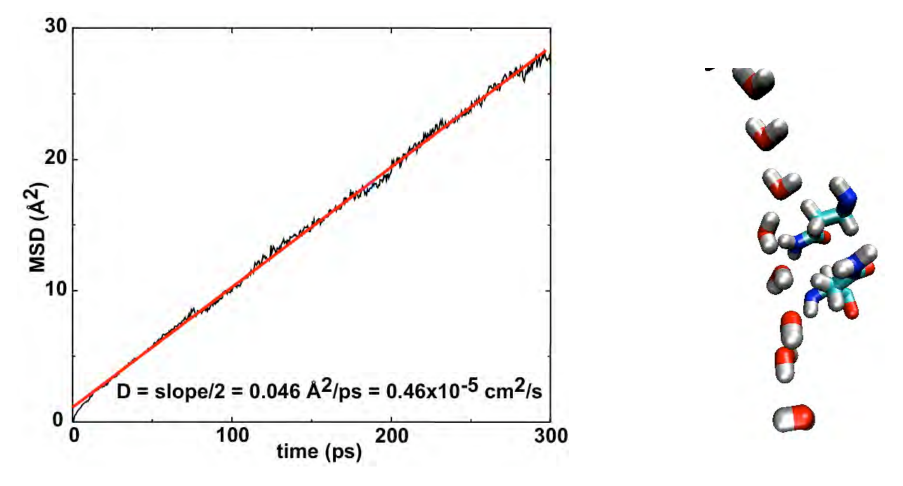
Correlated Motion of Water in the Channel

The single file of water molecules is maintained.





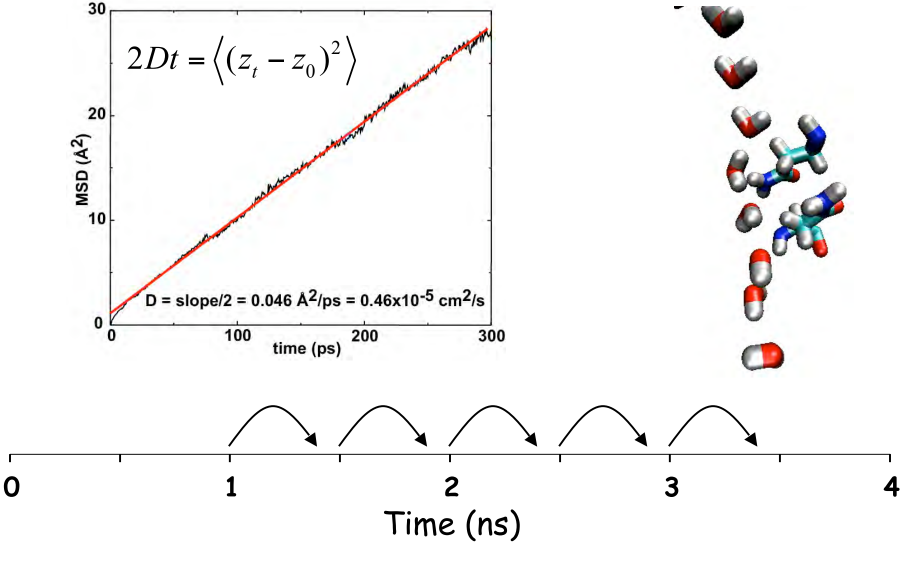
Diffusion of Water in the channel



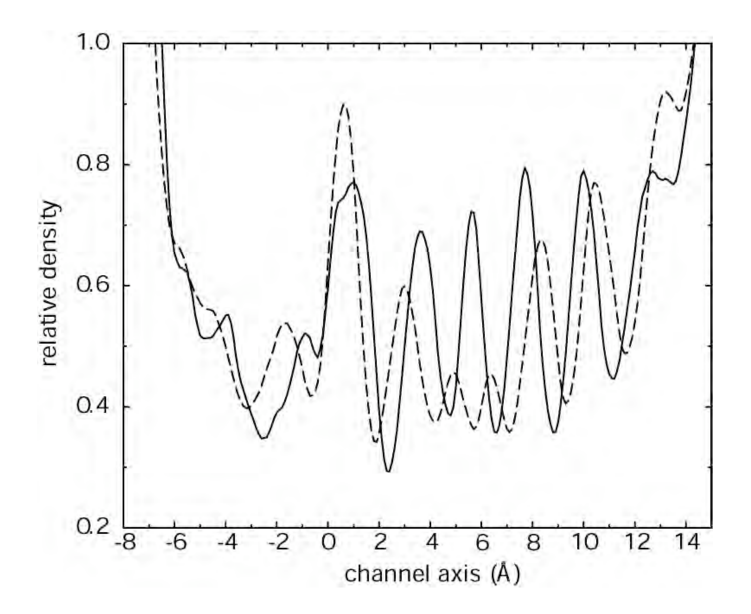
One dimensional diffusion: $2Dt = \left\langle (z_t - z_0)^2 \right\rangle$

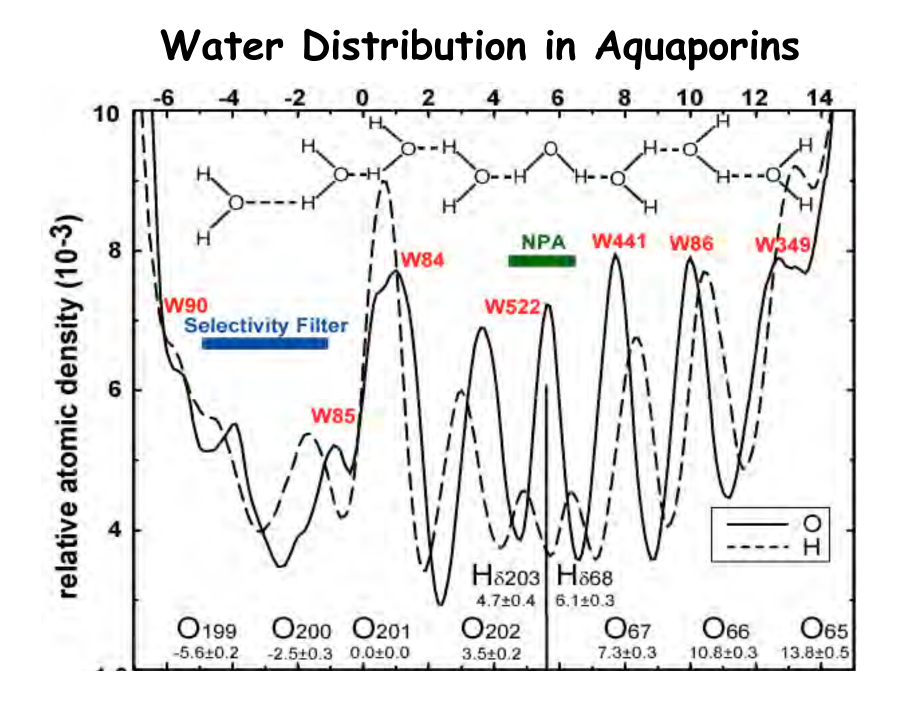
Experimental value for AQP1: 0.4-0.8 e-5

Diffusion of Water in the channel

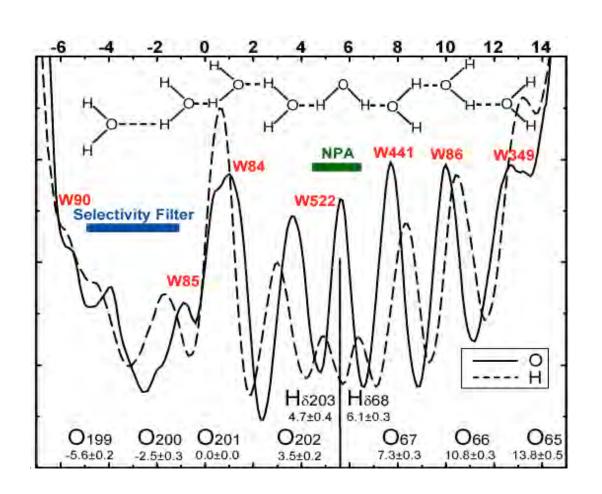


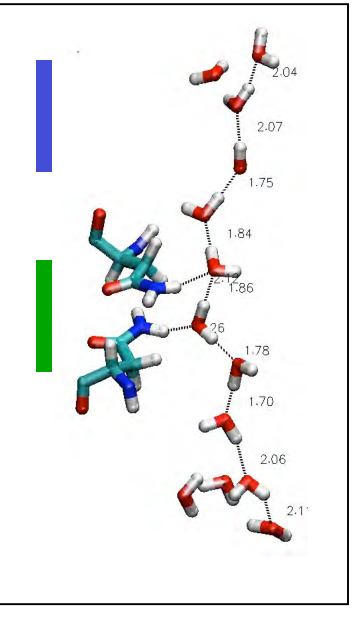
Improvement of statistics



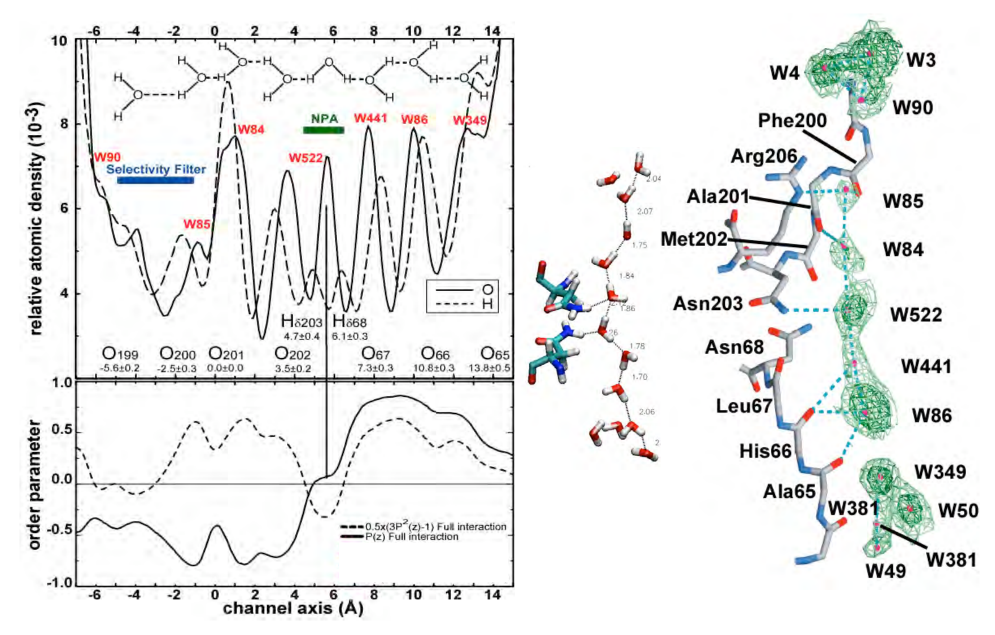


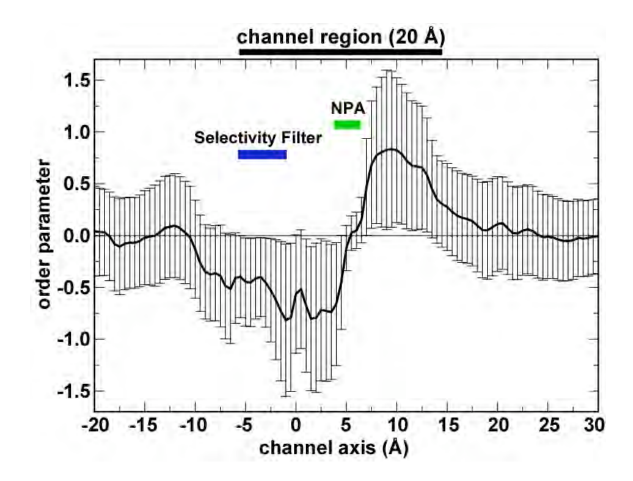
Water Bipolar Configuration in Aquaporins





Water Bipolar Configuration in Aquaporins



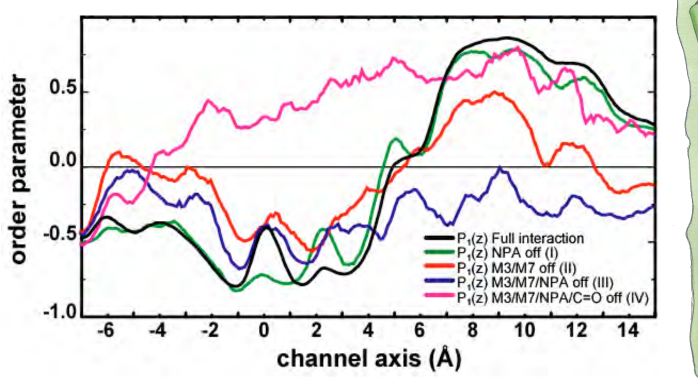


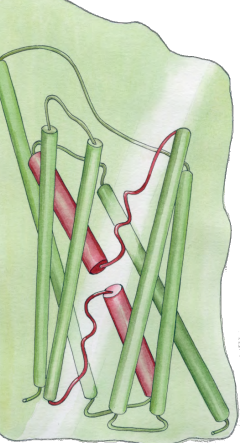
REMEMBER:

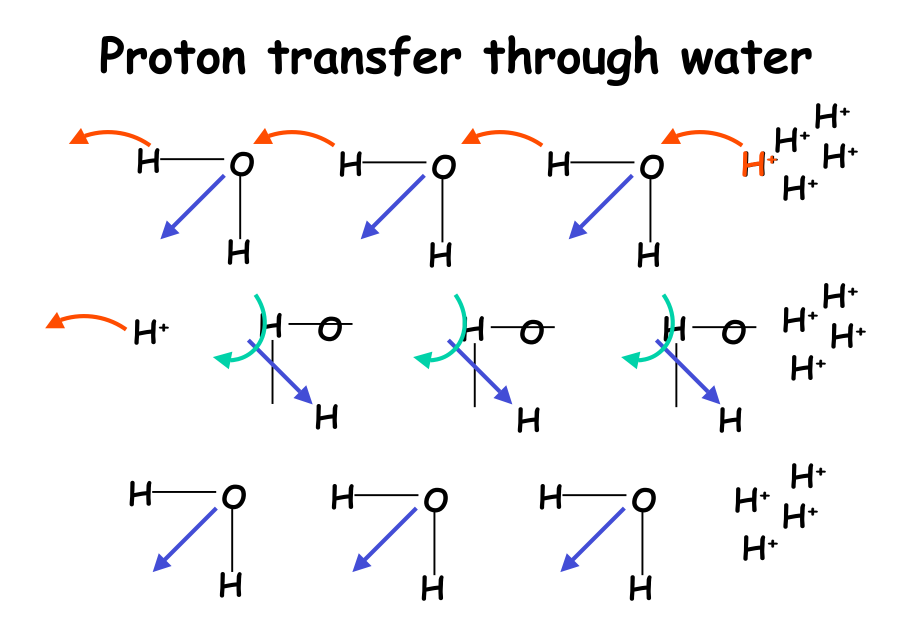
One of the most useful advantages of simulations over experiments is that you can modify the system as you wish: You can do modifications that are not even possible at all in reality!

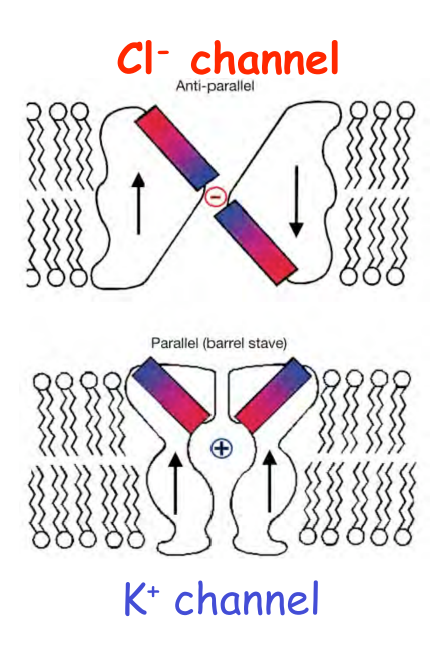
This is a powerful technique to test hypotheses developed during your simulations. Use it!

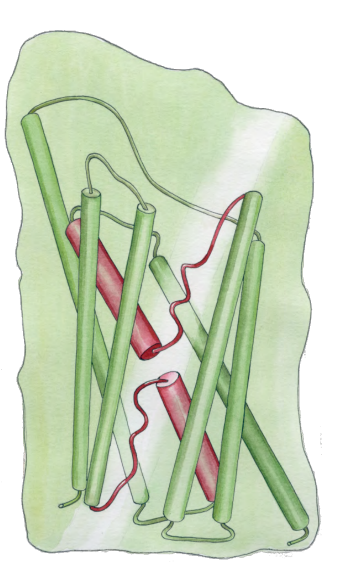
Electrostatic Stabilization of Water Bipolar Arrangement











Aquaporins

Proton Blocking by a Global Orientation Mechanism

