



**Mathematics**  
**describes only a tiny part of life,**

But

**Mathematics\* Creates**

our

**Standard of Living**

*\*e.g., Electricity, Computers, Fluid Dynamics, Optics, Structural Mechanics, ....*



**How can we use mathematics to describe biological systems?**

*I believe some biology is*  
**Physics ‘as usual’**  
**‘Guess and Check’**

*But you have to know which biology!*

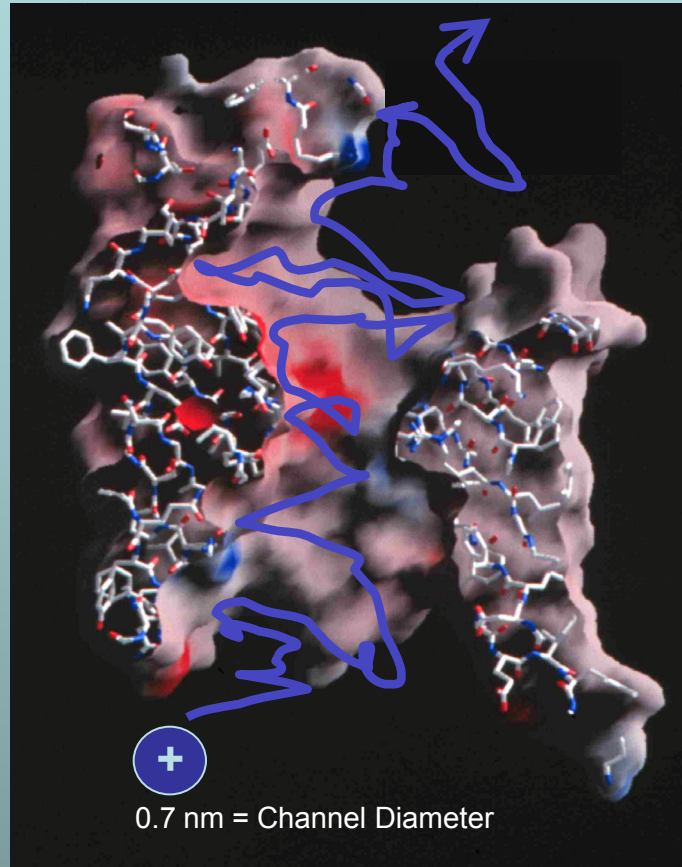
# Ion Channels are the Valves of Cells

Ion Channels are the Main Controllers of Biological Function

## Selectivity

Different Ions  
carry  
Different Signals

Chemical Bonds are lines  
Surface is Electrical Potential  
Red is negative (acid)  
Blue is positive (basic)



0.7 nm = Channel Diameter

~30 Å

Figure of ompF porin by Raimund Dutzler

## Ions in Water

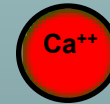
are the

Liquid of Life

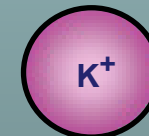
*Hard Spheres*



Na<sup>+</sup>



Ca<sup>++</sup>



K<sup>+</sup>



3 Å

# Ion Channels are Biological Devices

Natural nano-valves\* for atomic control of biological function

Ion channels coordinate contraction of cardiac muscle, allowing the heart to function as a pump

Ion channels coordinate contraction in skeletal muscle

Ion channels control all electrical activity in cells

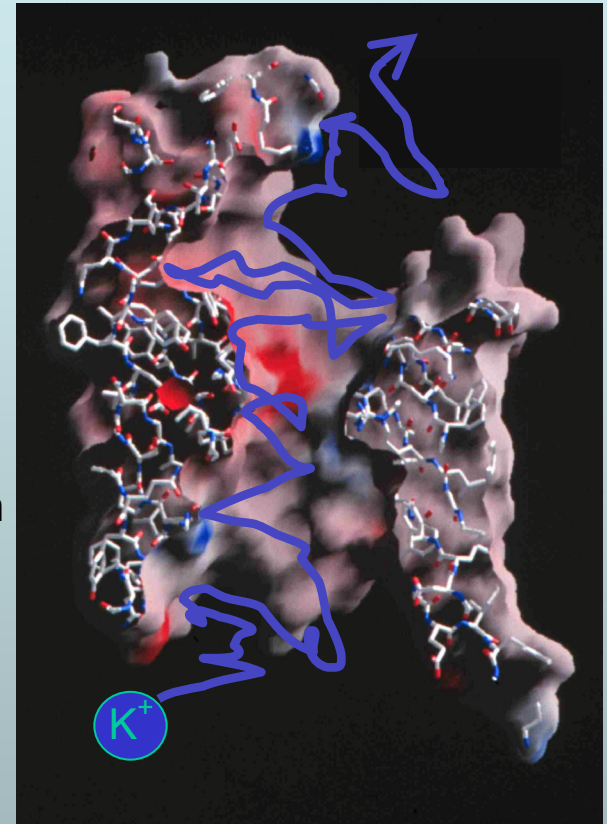
Ion channels produce signals of the nervous system

Ion channels are involved in secretion and absorption in all cells: kidney, intestine, liver, adrenal glands, etc.

Ion channels are involved in thousands of diseases and many drugs act on channels

Ion channels are proteins whose genes (blueprints) can be manipulated by molecular genetics

Ion channels have structures shown by x-ray crystallography in favorable cases



← ~30 Å →

\*nearly pico-valves: diameter is 400 – 900 picometers

# Thousands of Molecular Biologists Study Channels every day,

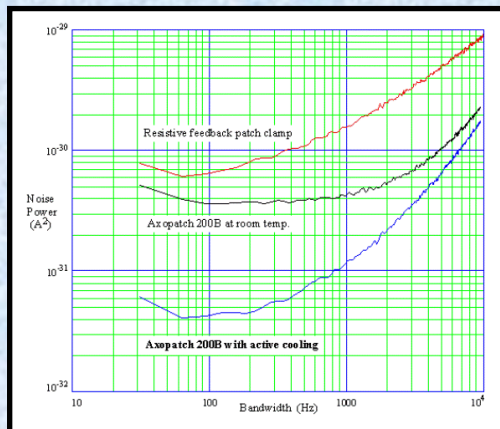
## One protein molecule at a time

This number is not an exaggeration.

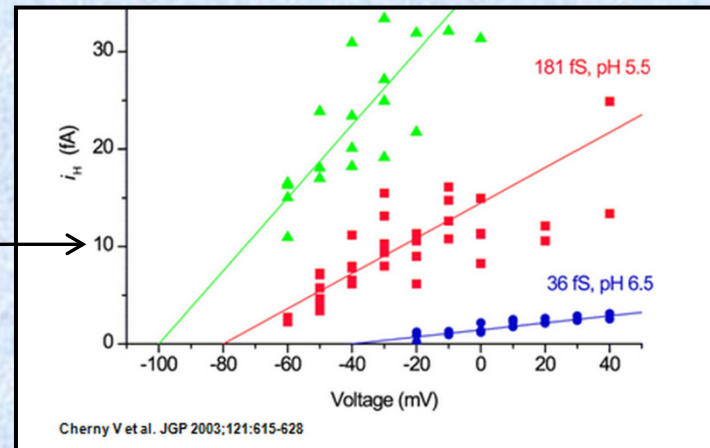
We have sold >10,000 AxoPatch amplifiers



AxoPatch 200B



**Femto-amps**  
( $10^{-15}$  A)



## Ion Channel Monthly

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Popular publications for March ([view most recent](#))

1. [Molecular basis of infrared detection by snakes](#). *Nature*
2. [The Angelman Syndrome Protein Ube3A Regulates Synapse Development by Ubiquitinating Arc](#). *Cell*
3. [AMPA receptors--another twist?](#) *Science*
4. [Molecular Basis of Calcium Signaling in Lymphocytes: STIM and ORAI](#). *Annu Rev Immunol*
5. [Neurological Channelopathies](#). *Annu Rev Neurosci*
6. [New antiarrhythmic drugs for treatment of atrial fibrillation](#). *Lancet*
7. [A Glial Signal Consisting of Gliomedin and NrCAM Clusters Axonal Na\(+\) Channels during the Formation of Nodes of Ranvier](#). *Neuron*
8. [Small Molecule Activators of TRPM1.3](#). *Chem Biol*
9. [Truncated \(beta\)-amyloid peptide channels provide an alternative mechanism for Alzheimer's Disease and Down syndrome](#). *Proc Natl Acad Sci U S A*
10. [Modelling the molecular mechanisms of synaptic plasticity using systems biology approaches](#). *Nat Rev Neurosci*

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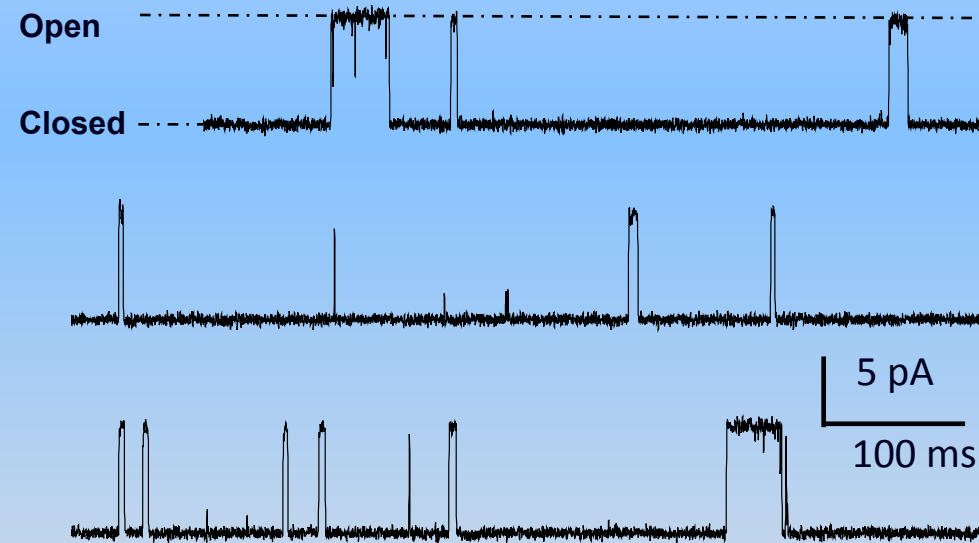
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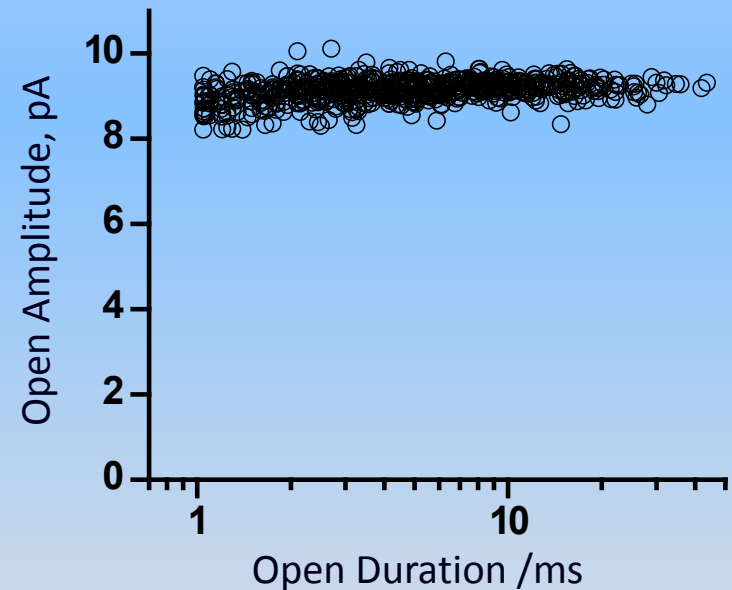
Why not forward  
this to your colleagues?  
They'll thank you  
for it!

# Channel Structure Does Not Change once the channel is open

Current vs. time



Amplitude vs. Duration



Lowpass Filter = 1 kHz    Sample Rate = 20 kHz

Typical Raw Single Channel Records

Ca<sup>2+</sup> Release Channel of Inositol Trisphosphate Receptor: slide and data from Josefina Ramos-Franco. Thanks!

*Channels are only Holes*  
*Why can't we understand and build them?*

***Where to start?***

**Why not compute all the atoms?**

# Multiscale Issues

*more later*

Computational Scale	Biological Scale	Ratio
<u>Time</u> $10^{-15}$ sec	$10^{-4}$ sec	$10^{11}$
<u>Space</u> $10^{-11}$ m	$10^{-5}$ m	$10^6$
<u>Spatial Resolution</u>	<i>Three Dimensional</i> $(10^6)^3$	$10^{18}$
<u>Solute Concentration</u>		$10^{11}$

Biological Scales Occur Together  
so must be

Computed Together

This may be impossible in simulations

Physicists and Engineers rarely try



## Multiscale Issues

**It may not be possible to deal accurately**

with

**Ratios of Scales**

of

**$10^{11}$   $10^6$   $10^{18}$   $10^{12}$**

*All at Once*

***Physicists and Engineers rarely try!***

**Computational Biology is  
NOT doing 'Physics as Usual'**

# *Why can't we understand and build channels?*

**Uncalibrated Simulations  
will not make devices that  
actually work**

*Unpopular view because*

**Calibration is Hard Work**

*particularly for Non-Ideal systems*

*with*

***Correlations, Finite Size effects, and Flows***

***Where do we start?***

**Physics 'As Usual'**

**'Guess and Check'**

***Stochastic 'Derivation'***

**Later**

*will include biological adaptation  
of Correlations and Crowded Charge*

# We start with Langevin equations of charged particles



*Opportunity  
and Need*

Simplest stochastic trajectories  
are  
Brownian Motion of Charged Particles

*Einstein, Smoluchowski, and Langevin ignored charge  
and therefore  
do not describe Brownian motion of ions in solutions*

We use

## Theory of Stochastic Processes

*to go*

## from Trajectories to Probabilities

*Once we learn to count Trajectories of Brownian Motion of Charge,  
we can count trajectories of Molecular Dynamics*

# Langevin Equations

Positive cation,  
e.g.,  $p = \text{Na}^+$

$$\ddot{x}_k^p - \frac{f_k^p(\tilde{x}; q_k)}{m} = -\gamma \dot{x}_k^p + \sqrt{\frac{2\gamma kT}{m}} \dot{w}_k^p$$

Negative anion,  
e.g.,  $n = \text{Cl}^-$

$$\ddot{x}_k^n - \frac{f_k^n(\tilde{x}; q_k)}{m} = -\gamma \dot{x}_k^n + \sqrt{\frac{2\gamma kT}{m}} \dot{w}_k^n$$

Newton's Law

Friction & Noise

## Electric Force

from all charges including  
Permanent charge of **Protein**,  
Dielectric Boundary charges,  
Boundary condition charge

# Electric Force from Poisson Equation

Excess  
'Chemical'  
Force

**Electric Force**  
from all charges including  
**P**ermanent charge of **P**rotein,  
Dielectric Boundary charges,  
Boundary condition charge

$$f_k^P(\vec{\mathbf{x}}) = f_{xs} + q_k(\vec{\mathbf{x}}) \operatorname{div} \left( \epsilon(\vec{\mathbf{x}}) \vec{\mathbf{E}} \right) = \frac{e}{\epsilon_0} \mathbf{P}(\mathbf{x}) + \frac{e}{\epsilon_0} \sum_i z_i \rho_i(\mathbf{x})$$

Implicit Solvent  
'Primitive' Model  
or  
Primitive Solvent Model

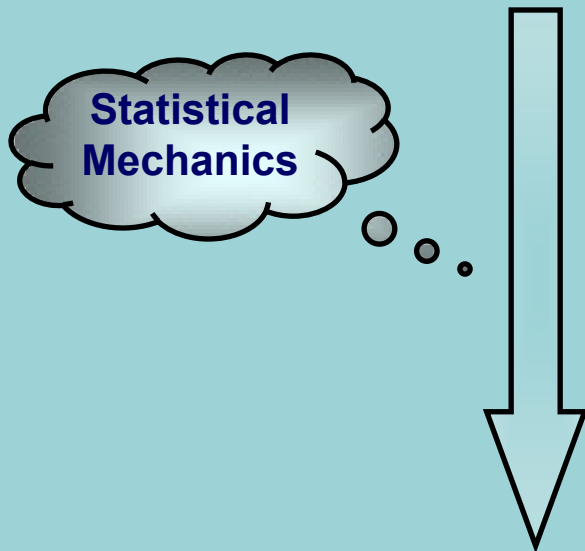
# Equilibrium

## Thermodynamics

Configurations

Boltzmann Distribution

$$\lim N, V \rightarrow \infty$$



***Thermodynamics***

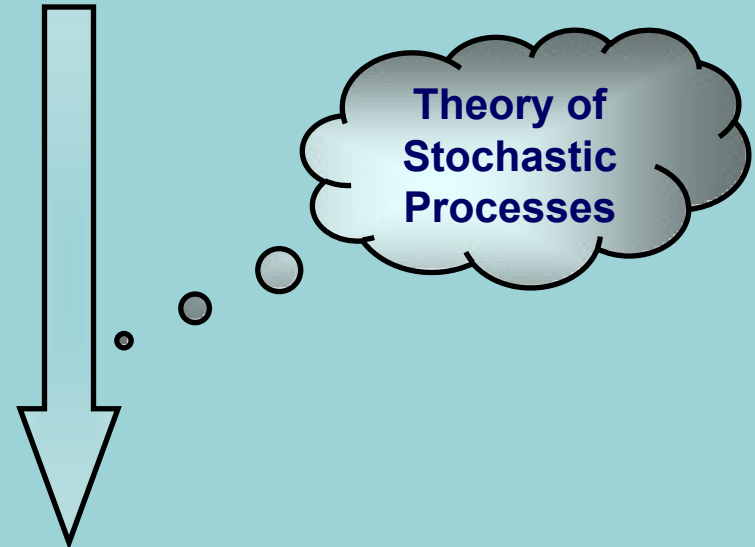
# Nonequilibrium

Schuss, Nadler, Singer & Eisenberg

Trajectories

Fokker Planck Equation

Finite OPEN System



***Device Equation***

# ***From Trajectories to Probabilities***

## **Main Result of Theory of Stochastic Processes**

*Joint* probability density of position and velocity

$$p(\tilde{x}, \tilde{v}) = \mathbf{Pr} \left\{ \left\{ x_j, v_j \right\}_{j=1}^{2N} \right\}; \quad N = \text{Number of Particles}$$

satisfies a Fokker Planck equation

$$0 = \sum_j \mathbf{L}_j^p p(\tilde{x}, \tilde{v}) + \sum_j \mathbf{L}_j^n p(\tilde{x}, \tilde{v})$$

with Fokker Planck Operator

$$\mathbf{L}_j^c p = -v_j^c \cdot \nabla_{x_j^c} p + \nabla v_j^c \cdot \left( \gamma v_j^c - \frac{f_j^c}{m_j^c} \right) p + \nabla \cdot \nabla_{v_j^c} \frac{\gamma kT}{m_j^c} p$$

*Coordinates are positions and velocities of  $N$  particles in  $12N$  dimensional phase space*



# Conditional PNP

Electric Force  $\nabla \bar{\phi}$  depends on Conditional Density of Charge

$$\nabla_y \cdot \left[ \frac{\epsilon_0 \epsilon(y)}{e} \nabla_y \bar{\phi}(y|x) \right] = P(y)$$

Permittivity, Dielectric Coefficient, Charge on Electron

Channel Protein

$$+ \rho_+(y|x) - \rho_-(y|x)$$

Closures or Approximations Needed

Nernst-Planck gives UNconditional Density of Charge

$$\nabla_y \cdot \left[ \frac{1}{m\gamma(x)} \rho_+(x) \left[ e \nabla_y \bar{\phi}(y|x) \Big|_{y=x} - (\text{Other Forces}) \right] \right] = 0$$

Mass

Friction

# Poisson-Nernst-Planck (PNP)

## Poisson's Equation

$$-\epsilon_0 \nabla \cdot \left( \epsilon(\mathbf{x}) \nabla \phi(\mathbf{x}) \right) = eP(\mathbf{x}) + e \sum_i z_i \rho_i(\mathbf{x})$$

Dielectric Coefficient  $\epsilon(\mathbf{x})$   
 Permittivity  $\epsilon_0$   
 Channel Protein  $P(\mathbf{x})$   
 Proton charge Valence  $z_i$   
 Number Densities  $\rho_i(\mathbf{x})$

## Drift-diffusion & Continuity Equation

$$\nabla \cdot \mathbf{J}(\mathbf{x}) = 0; \quad -\mathbf{J}_i(\mathbf{x}) = D_i(\mathbf{x}) \rho_i(\mathbf{x}) \frac{1}{kT} \nabla \mu_i(\mathbf{x})$$

Flux  $\mathbf{J}_i(\mathbf{x})$   
 Diffusion Coefficient  $D_i(\mathbf{x})$   
 Thermal Energy  $kT$   
 Chemical Potential  $\mu_i(x)$

$$\mu_i(\mathbf{x}) = z_i e \phi(\mathbf{x}) + kT \ln \left( \frac{\rho_i(\mathbf{x})}{\rho^*} \right) + \mu_i^{\text{ex}}(\mathbf{x})$$

Valence Proton charge  $z_i$   
 Thermal Energy  $kT$   
 Chemical Correlations  $\mu_i^{\text{ex}}(\mathbf{x})$

# Semiconductor Equations: One Dimensional PNP

## Poisson's Equation

$$-\frac{\epsilon_0}{A(x)} \frac{d}{dx} \left( \epsilon(x) A(x) \frac{d\phi}{dx} \right) = eP(x) + e \sum_i z_i \rho_i(x)$$

Dielectric Coefficient  $\epsilon_0$   
 Cross sectional Area  $A(x)$   
 Permanent Charge of Protein  $P(x)$   
 Valence Proton charge  $z_i$   
 Number Densities  $\rho_i(x)$

## Drift-diffusion & Continuity Equation

$$\frac{dJ_i}{dx} = 0 \quad -J_i = D_i(x) A(x) \rho_i(x) \frac{d\mu_i}{dx}$$

Flux  $J_i$   
 Diffusion Coefficient  $D_i(x)$   
 Number Densities  $\rho_i(x)$   
 Chemical Potential  $\mu_i$

## Chemical Potential $\mu_i(x)$

$$\mu_i(x) = z_i e \phi(x) + kT \ln \left( \frac{\rho_i(x)}{\rho^*} \right) + \underbrace{\mu_i^{\text{ex}}(x)}_{\text{Special Chemistry}}$$

valence proton charge  $z_i$   
 Thermal Energy  $kT$   
 Special Chemistry  $\mu_i^{\text{ex}}(x)$

Counting at low resolution gives  
'Semiconductor Equations'

# Poisson-Nernst-Planck (PNP)

*Ions are Points in PNP  
contains only the  
Correlations of Means*

Gouy-Chapman, (nonlinear) Poisson-Boltzmann,  
Debye-Hückel,

are siblings with similar resolution

but at equilibrium, without current or flux of any species

**Devices do not exist at equilibrium**

How do we check the theory?

# Compare with Biological Function!

Our task is to  
**Discover & Understand, Control & Improve**  
Biological Function



Inverse  
Problem

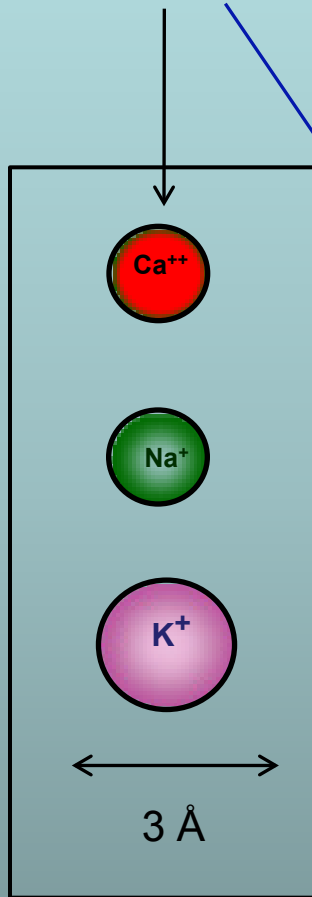
That means **Selectivity**

Ions are not Ideal

*Potassium  $K^+$   $\neq$   $Na^+$  Sodium*

# Channels are Selective

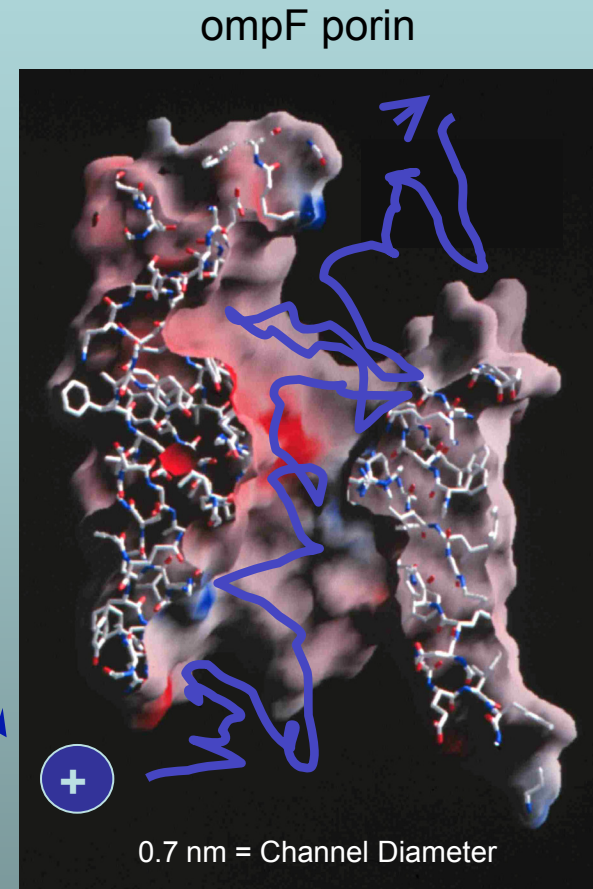
Different Ions Carry Different Signals through Different Channels



## Diameter matters

Diameter is the Only Difference between  $\text{K}^+$  and  $\text{Na}^+$

*In ideal solutions  $\text{K}^+ = \text{Na}^+$*



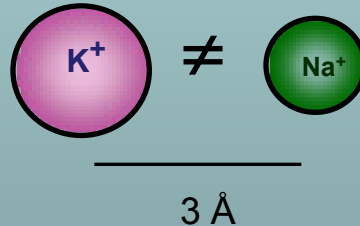
$\sim 30 \text{ \AA}$

Flow time scale is 0.1 msec to 1 min

Figure of *ompF* porin by Raimund Dutzler

# Channels are Selective because Ions are NOT Ideal

*Potassium  $K^+$   $\neq$   $Na^+$  Sodium*



**Ideal Electrolytes are Identical**  
*if they have the same charge*

*Modelers and Mathematicians, Bioengineers: this is reverse engineering*

*How does the*  
**Channel control Selectivity?**

# **Inverse Problems**

*Many answers are possible*

**Central Issue**

**Which answer is right?**

**Core Math Problem has actually been solved  
using methods for the**

**Inverse Problem of a Blast Furnace**

Burger, Eisenberg and Engl (2007) SIAM J Applied Math 67: 960-989



How does the  
Channel control Selectivity?

**Inverse Problems: many answers possible**

Central Issue

Which answer is right?

Key is

**ALWAYS**

Large Amount of Data

from

Many Different Conditions

**Goal:**

# **Understand Selectivity**

well enough to

## **Fit Large Amounts of Data**

from many solutions and concentrations

and to

## **Make a Calcium Channel**

**Atomic Scale**

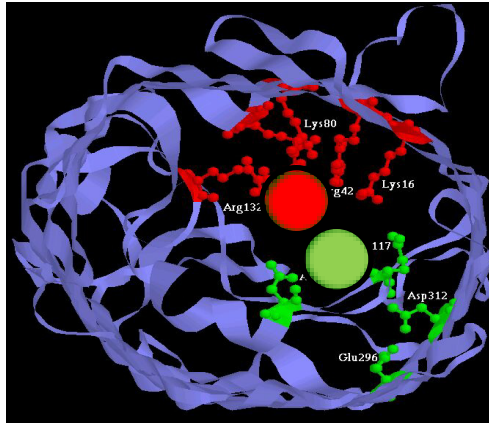


**Macro Scale**

Experiments have built

**Atomic Scale**

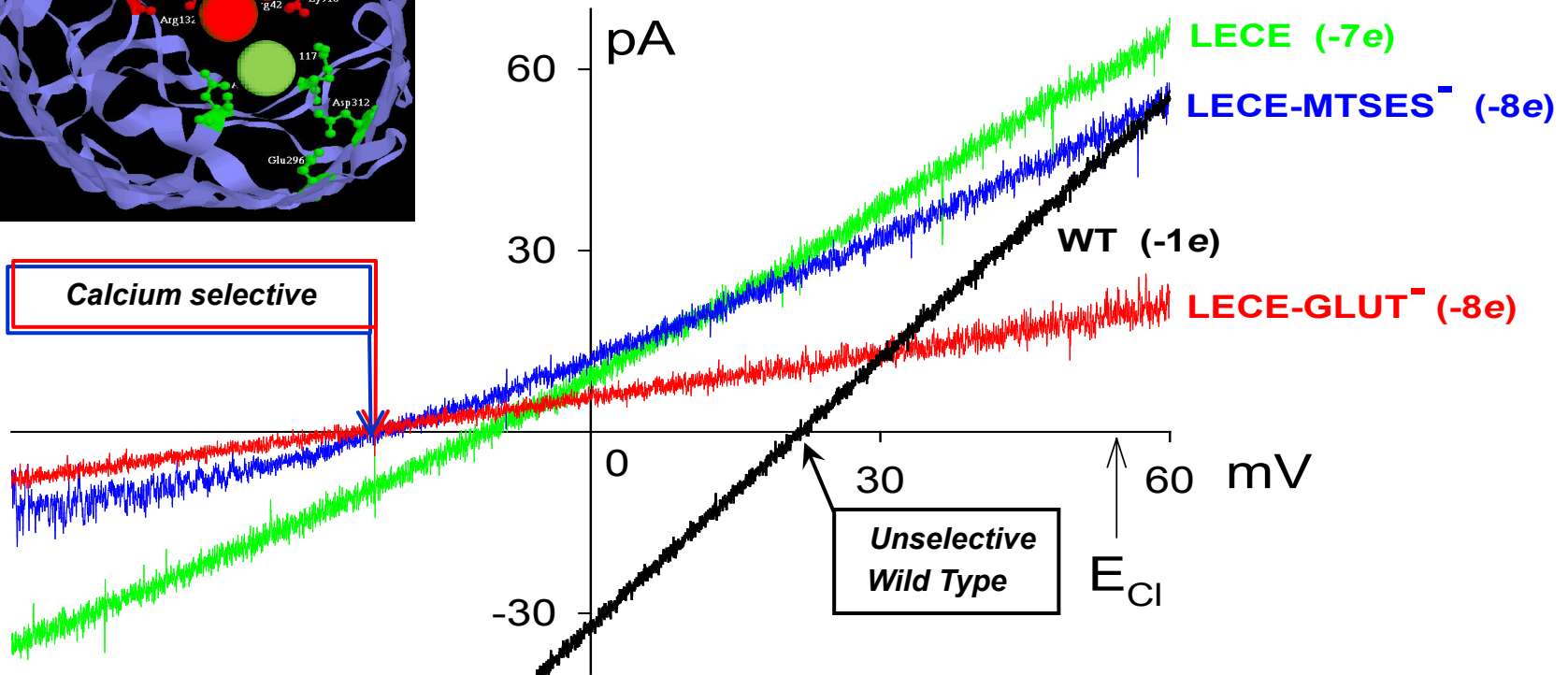
# Two Synthetic Calcium Channels



**Calcium selective**

Designed by Theory

Glutathione derivatives



As density of permanent charge increases, channel becomes calcium selective

$E_{\text{rev}} \rightarrow E_{\text{Ca}}$  in 0.1M || 1.0 M  $\text{CaCl}_2$

**Macro Scale**

built by Henk Miedema, Wim Meijberg of BioMade Corp., Groningen, Netherlands

27

Miedema et al, *Biophys J* 87: 3137–3147 (2004)

*Comparison with Experiments shows  
Potassium  $K^+$   $\neq$  Sodium  $Na^+$*

**Must include Biological  
Adaptation!**

*Working Hypothesis*

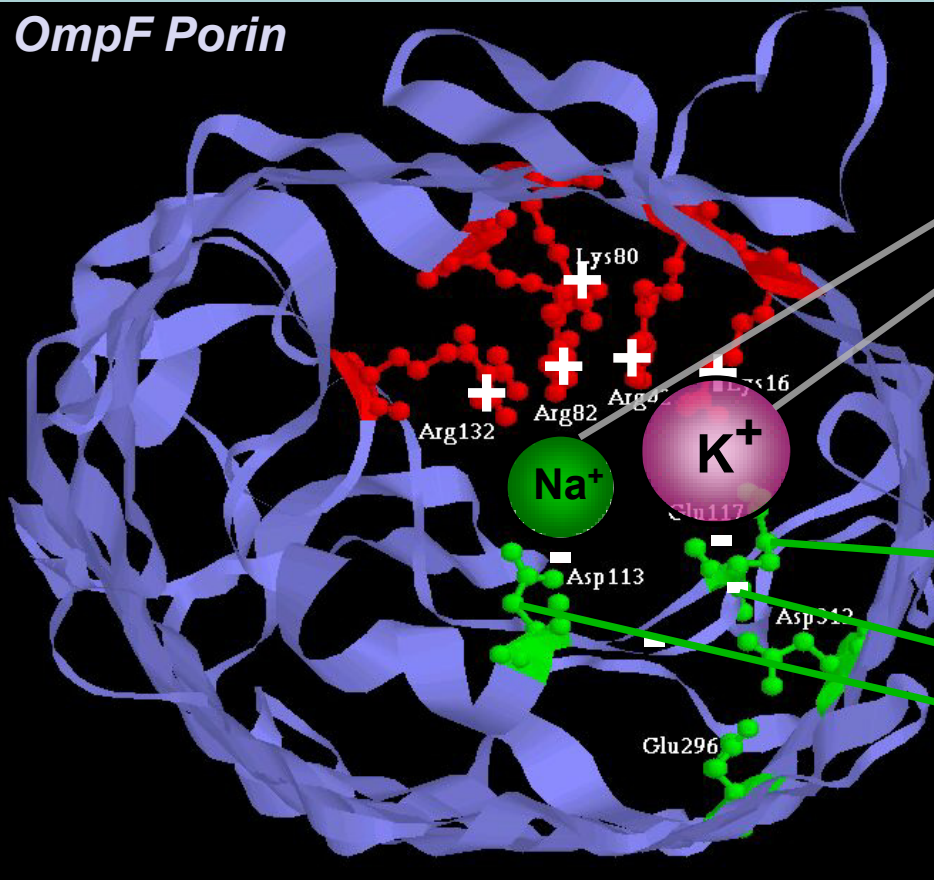
Biological Adaptation is

**Crowded Ions *and* Side Chains**

# Active Sites of Proteins are Very Charged

7 charges ~ 20 M net charge =  $1.2 \times 10^{22} \text{ cm}^{-3}$

liquid **Water** is **55 M**  
solid **NaCl** is **37 M**

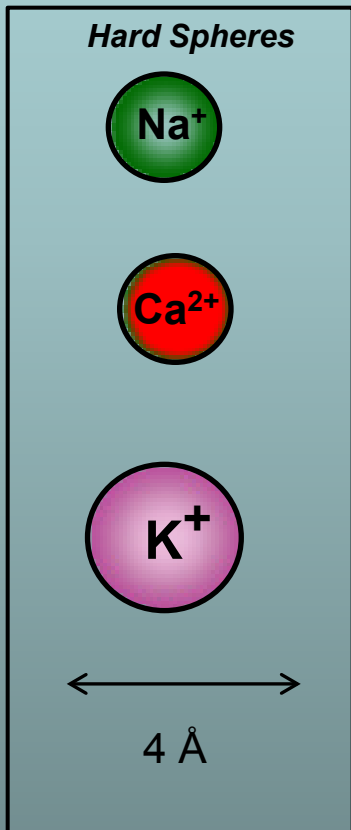


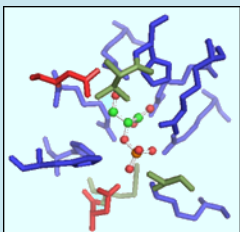
Ions are Crowded

Induced Fit of Side Chains

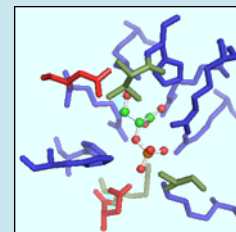
Selectivity Filters and Gates of Ion Channels are Active Sites

Figure adapted from Tilman Schirmer





# Charge Density 22 M



		#AA	MS_A^3	CD_MS+	CD_MS-	CD_MSt
<b>EC1:Oxidoreductases</b>	Average	47.2	1,664.74	<b>7.58</b>	<b>2.82</b>	<b>10.41</b>
	Median	45.0	1,445.26	<b>6.12</b>	<b>2.49</b>	<b>8.70</b>
<b>EC2:Transferases</b>	Average	33.8	990.42	<b>13.20</b>	<b>6.63</b>	<b>19.83</b>
	Median	32.0	842.43	<b>8.18</b>	<b>6.71</b>	<b>14.91</b>
<b>EC3:Hydrolases</b>	Average	24.3	682.88	<b>13.14</b>	<b>13.48</b>	<b>26.62</b>
	Median	20.0	404.48	<b>11.59</b>	<b>12.78</b>	<b>23.64</b>
<b>EC4:Lyases</b>	Average	38.2	1,301.89	<b>13.16</b>	<b>6.60</b>	<b>19.76</b>
	Median	28.0	822.73	<b>10.81</b>	<b>4.88</b>	<b>16.56</b>
<b>EC5:Isomerases</b>	Average	31.6	1,027.15	<b>24.03</b>	<b>11.30</b>	<b>35.33</b>
	Median	34.0	989.98	<b>9.05</b>	<b>7.76</b>	<b>16.82</b>
<b>EC6:Ligases</b>	Average	44.4	1,310.03	<b>9.25</b>	<b>9.93</b>	<b>19.18</b>
	Median	49.0	1,637.98	<b>8.32</b>	<b>7.95</b>	<b>17.89</b>
<b>Total</b> <i>n= 150</i>	Average	36.6	1,162.85	<b>13.39</b>	<b>8.46</b>	<b>21.86</b>
	Median	33.0	916.21	<b>8.69</b>	<b>7.23</b>	<b>16.69</b>



**EC#:** Enzyme Commission Number based on chemical reaction catalyzed  
**#AA:** Number of residues in the functional pocket  
**MS\_A^3:** Molecular Surface Area of the Functional Pocket (Units Angstrom^3)  
**CD\_MS+:** Charge Density (**positive**)  
**CD\_MS-:** Charge Density (**negative**)  
**CD\_MSt:** Total Charge density

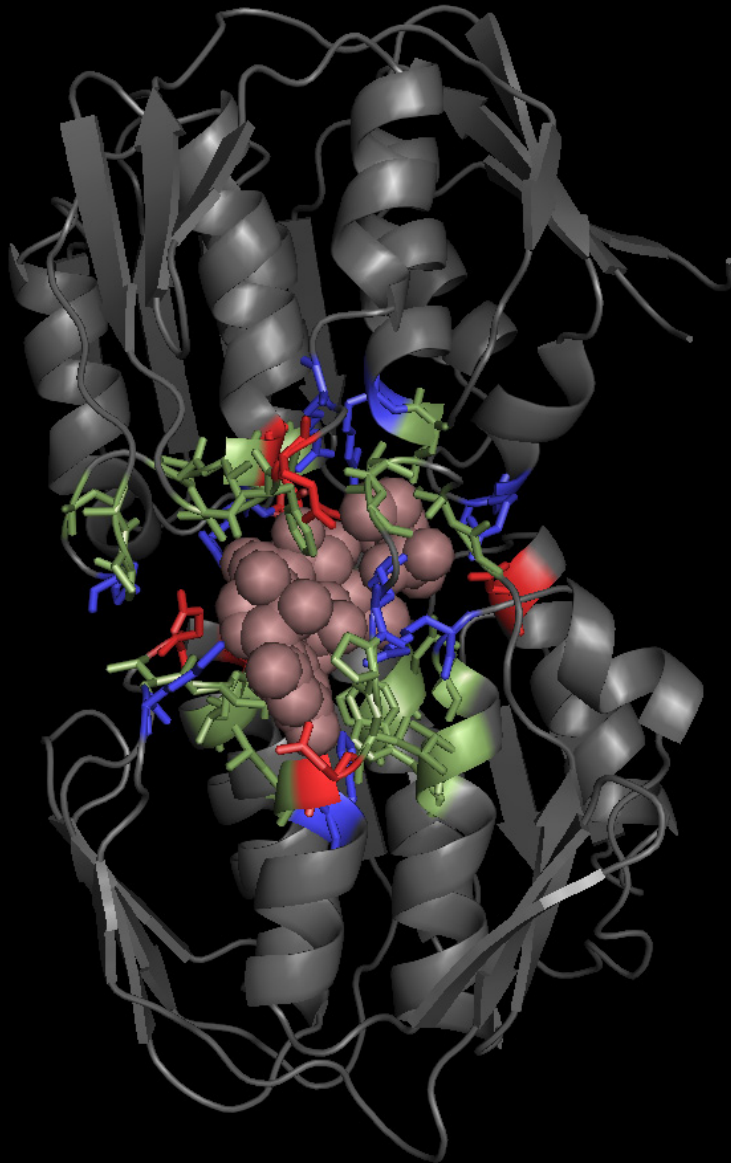
Jimenez-Morales,  
 Liang,  
 Eisenberg

# EC2: TRANSFERASES

Average Charged Density: 19.8 Molar

Example:  
UDP-N-ACETYLGLUCOSAMINE  
ENOLPYRUVYL TRANSFERASE  
(PDB:1UAE)

Functional Pocket Molecular Surface Volume:  
1462.40 Å<sup>3</sup>  
Density Charge: 19.3 Molar (11.3 M+ . 8 M-)



Green: Functional pocket residues

Blue: Basic = Positive charged = R+K+H

Red: Acid = Negative charged = E + D

Brown URIDINE-DIPHOSPHATE-N-  
ACETYLGLUCOSAMINE

*Jimenez-Morales, Liang, Eisenberg*

*Working Hypothesis*

Biological Adaptation is

**Crowded Ions *and* Side Chains**

**Everything interacts**



*Working Hypothesis*

# **Interactions in Channels**

come mostly from

***Finite Size Effects***

**Chemically Specific Properties**

come from

**Diameter and Charge**

learned from Doug Henderson, J.-P. Hansen, Stuart Rice, among others...  
Thanks!

## Bulk Solutions:

Interactions come mostly from  
Finite Size Effects

# Chemically Specific Properties

of ions (e.g. activity = free energy per mole)  
are known to come from interactions of their

## Diameter and Charge

and dielectric 'constant' of ionic solution

Atomic Detail 

'Primitive Implicit Solvent Model'

learned from Doug Henderson, J.-P. Hansen, Stuart Rice, among others...

Thanks!

**Ions in Water are the Liquid of Life**

**They are not ideal solutions**

**Everything  
Interacts  
with  
Everything**

***For Modelers and Mathematicians***

**Tremendous Opportunity for Applied Mathematics**

**Chun Liu's Energetic Variational Principle**

***EnVarA***

# Variational Principles Deal with Interactions Consistently and Automatically

*Chun Liu,*  
*with Yunkyong Hyon, and Bob Eisenberg*

## EnVarA

$$\overbrace{\frac{\delta E}{\delta \vec{x}}}^{\text{Conservative 'Force'}} - \overbrace{\frac{1}{2} \frac{\delta \Delta}{\delta \vec{u}}}^{\text{Dissipative 'Force'}} = 0$$

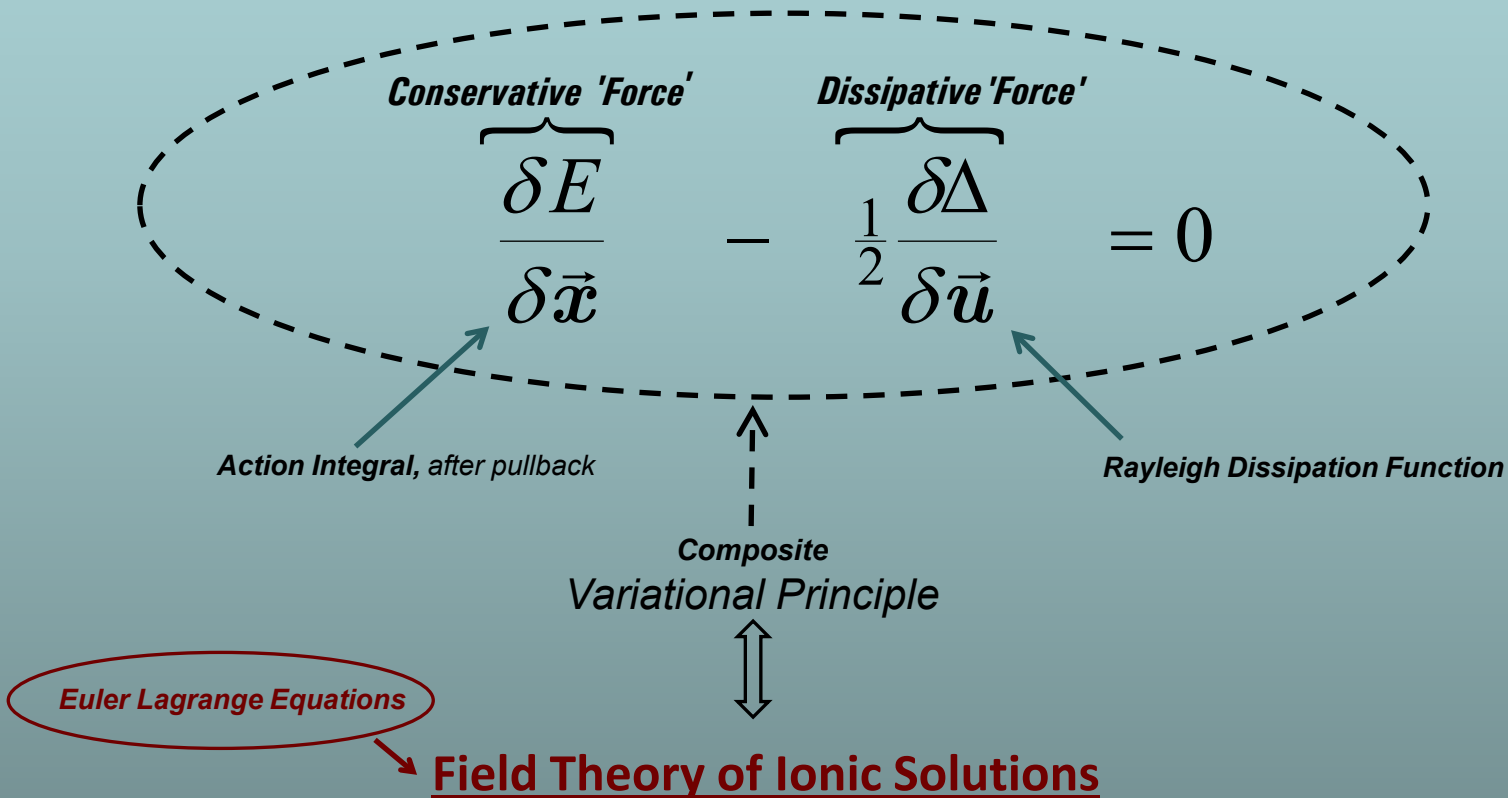
**New Component** (or Scale)  
implies  
**New Field Equations (Euler Lagrange)**  
by  
**Algebra Alone**  
*No new Assumptions*

# Energetic Variational Approach

## *EnVarA*

Chun Liu, Rolf Ryham, Yunkyong Hyon, and Bob Eisenberg

*Mathematicians and Modelers: two different 'partial' variations written in one framework, using a 'pullback' of the action integral*



that allows boundary conditions and flow and deals with Interactions of Components self-consistently

# Variational Analysis of Ionic Solution

## EnVarA

Generalization of Chemical Free Energy

$$E(\text{Primitive Phase}; t) = \int \left[ \underbrace{\frac{1}{2} \rho |\vec{u}_{IP}|^2}_{\text{Hydrodynamic Kinetic Energy}} + \underbrace{w(\rho)}_{\text{Hydrodynamic Potential Energy Equation of State}} \right] d\vec{x}$$

**Macroscopic** (hydrodynamic)

$$+ \lambda \left[ \underbrace{\frac{1}{2} \epsilon |\nabla \phi|^2}_{\text{Electrostatic}} + \underbrace{k_B T (c_n \log c_n + c_p \log c_p)}_{\text{Entropy}} + \underbrace{E(\text{Solid Spheres})}_{\text{Finite Size Effect}} \right] d\vec{x}$$

Dielectric Coefficient from Poisson Eq. →  $\epsilon$   
 Number Densities →  $c_n, c_p$   
 Lennard Jones →  $E(\text{Solid Spheres})$   
 Lagrange Multiplier →  $\lambda$

**Microscopic** (atomic)

# Dissipation Principle for Ions

Dissipative

$$\begin{aligned}
 & \frac{d}{dt} \int \left\{ k_B T \sum_{i=n,p} c_i \log c_i + \frac{1}{2} \left( \rho_0 + \sum_{i=n,p} z_i e c_i \right) \phi + \sum_{i,j=n,p} \frac{c_i}{2} \int \tilde{\Psi}_{i,j} c_j d\vec{y} \right\} d\vec{x} \\
 & = - \int \left\{ \sum_{i=n,p} \frac{D_i c_i}{k_B T} \left| k_B T \frac{\nabla c_i}{c_i} + z_i e \nabla \phi - \sum_{j=n,p} \nabla \int \tilde{\Psi}_{i,j} c_j d\vec{y} \right|^2 \right\} d\vec{x} \\
 & \hspace{15em} \text{Conservative}
 \end{aligned}$$

Annotations for the Dissipative part:
 

- $\frac{d}{dt}$ : time
- $k_B T$ : Thermal Energy
- $\sum_{i=n,p} c_i \log c_i$ : Number Density
- $\rho_0$ : Permanent Charge of protein
- $\sum_{i=n,p} z_i e c_i$ : valence proton charge
- $\int \tilde{\Psi}_{i,j} c_j d\vec{y}$ : Hard Sphere Terms

$c_i$  number density;  $k_B T$  thermal energy;  $D_i$  diffusion coefficient;  $n$  negative;  $p$  positive;  $z_i$  valence

# Field Equations with Lennard Jones Spheres

Non-equilibrium variational field theory *EnVarA*

## Nernst Planck Diffusion Equation

for number density  $c_n$  of negative  $n$  ions; positive ions are analogous

Diffusion Coefficient

$$\frac{\partial c_n}{\partial t} = \nabla \cdot \left[ D_n \left\{ \nabla c_n + \frac{c_n}{k_B T} \left( z_n e \nabla \phi - \int \frac{12 \epsilon_{n,n} (a_n + a_n)^{12} (\vec{x} - \vec{y})}{|\vec{x} - \vec{y}|^{14}} c_n(\vec{y}) d\vec{y} \right. \right. \right. \\ \left. \left. \left. - \int \frac{6 \epsilon_{n,p} (a_n + a_p)^{12} (\vec{x} - \vec{y})}{|\vec{x} - \vec{y}|^{14}} c_p(\vec{y}) d\vec{y} \right) \right\} \right],$$

Thermal Energy

Coupling Parameters

Ion Radii

Number Densities

## Poisson Equation

Dielectric Coefficient

$$\nabla \cdot (\epsilon \nabla \phi) = - \left( \rho_0 + \sum_{i=1}^N z_i e c_i \right) \quad i = n \text{ or } p$$

valence proton charge

Permanent Charge of Protein



# Energetic Variational Approach

*EnVarA across biological scales: molecules, cells, tissues*

developed by Chun Liu

with

(1) Hyon, Eisenberg

Ions in

Channels

(2) Bezanilla, Hyon, Eisenberg

Conformation Change of

Voltage Sensor

(3) Ryham, Eisenberg, Cohen

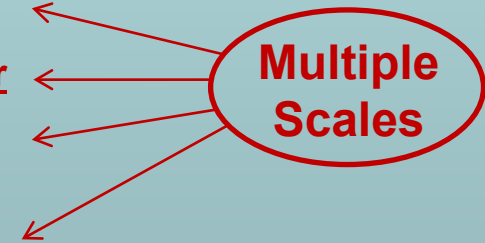
Virus fusion to

Cells

(4) Mori, Eisenberg

Water flow in

Tissues



creates a new

## Multiscale Field Theory of Interacting Components

that allows boundary conditions and flow  
and deals with

Ions in solutions self-consistently

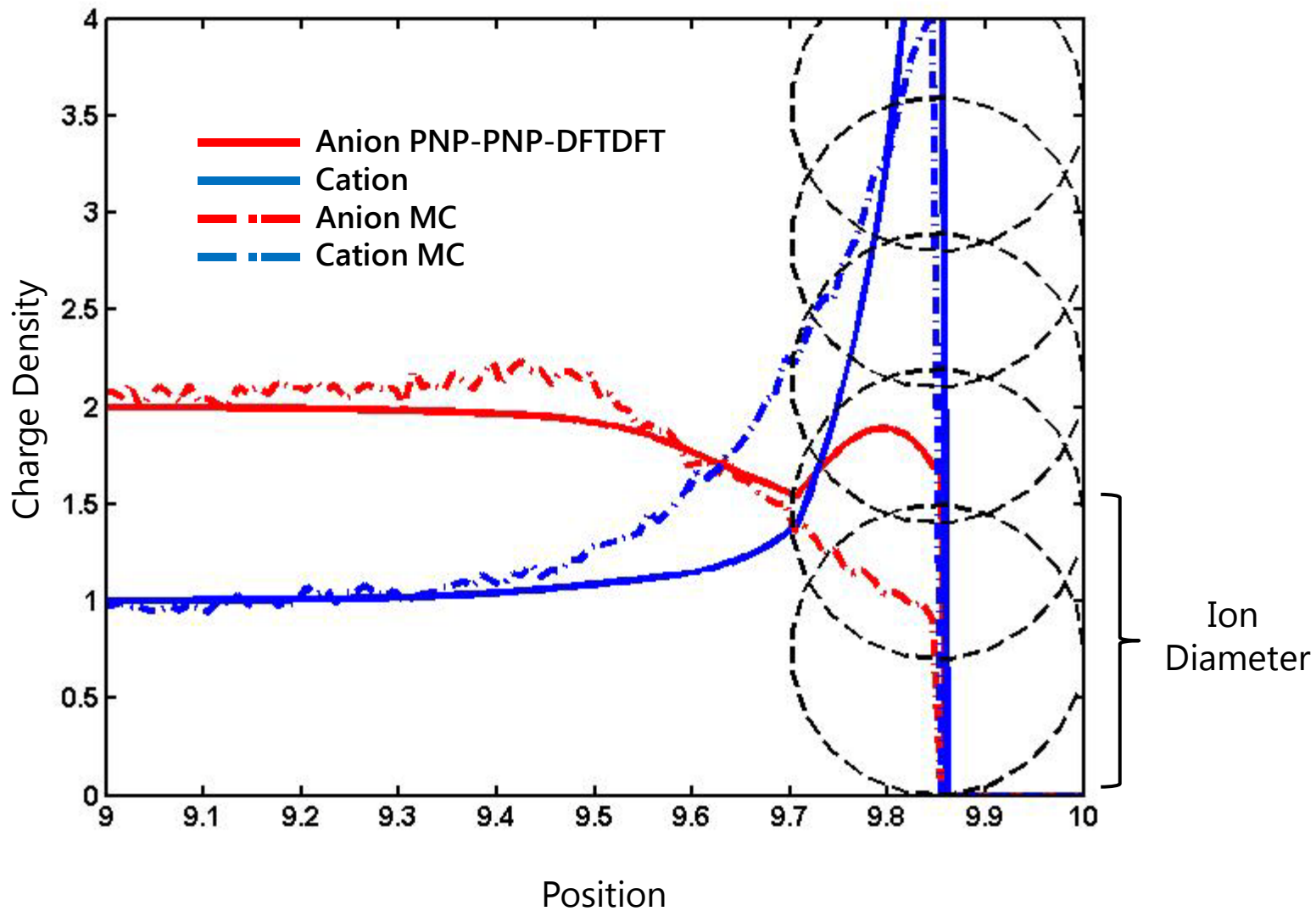
# **Energetic Variational Approach**

developed by Chun Liu

## **Preliminary Results** and **Provocations**

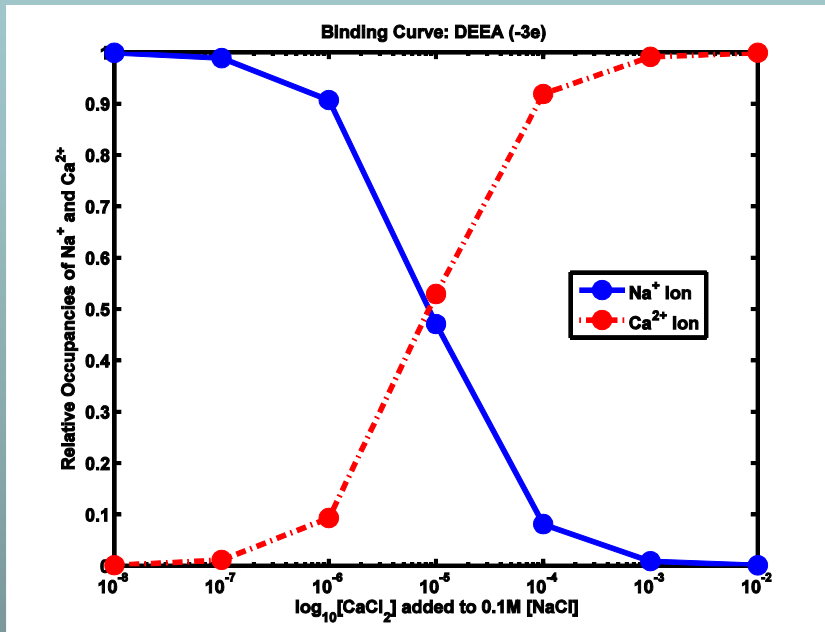
# Layering: Classical Interaction Effect

Comparison between PNP-DFT and MC

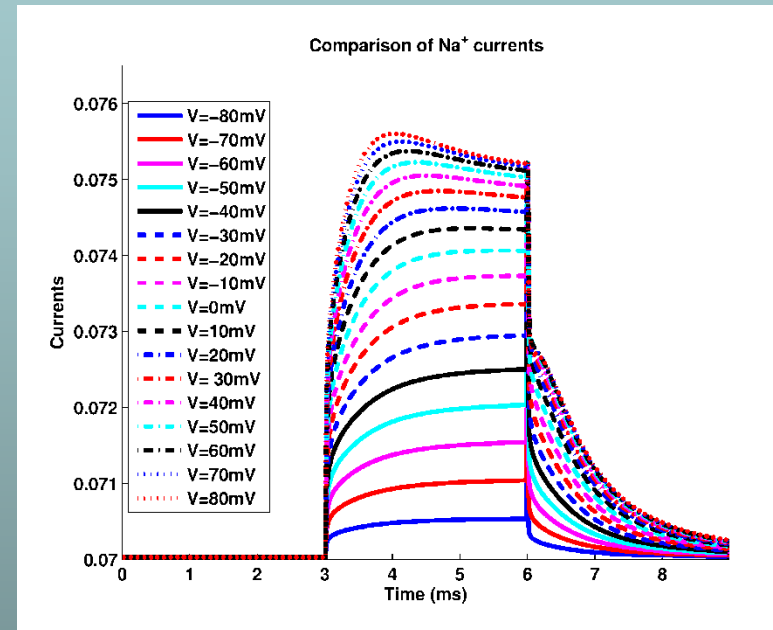


# Nonequilibrium Computations with Variational Field Theory *EnVarA*

Binding Curves



Current Voltage **Time** Curves



# Energetic Variational Approach *EnVarA*

**New mechanisms\***  
(e.g., active transport)  
**can be added**

**\*if they define an energy and its variation**

**Energy defined by simulations or theories or experiments is OK**

*Full micro/macro treatment is needed for an Atomic Model, with closure, as in liquid crystals*

**Back to the Calcium Channel**

**Then, the Sodium Channel**

# Selectivity Filter

Crowded with Charge

L type Ca Channel

Selectivity Filter

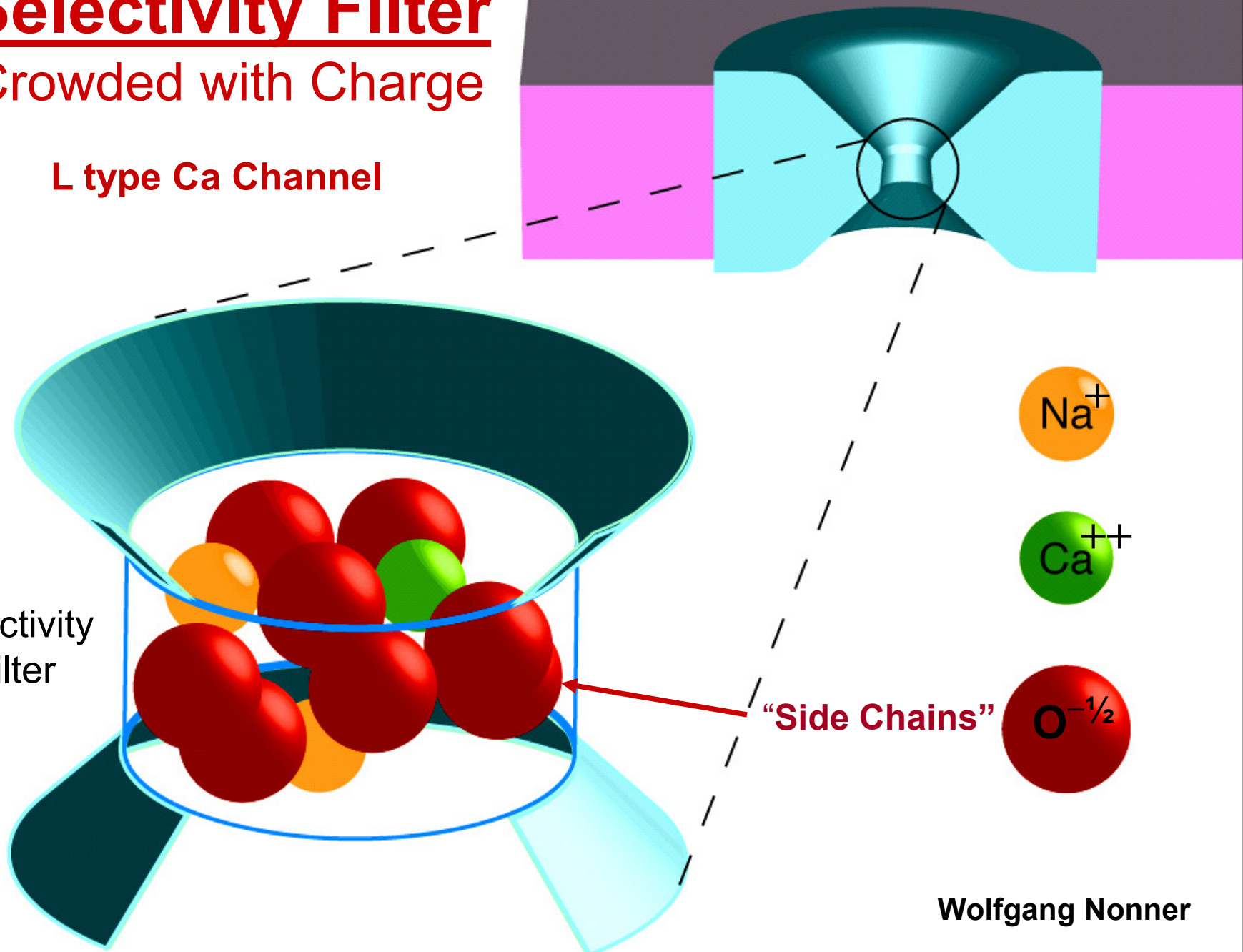
“Side Chains”

Na<sup>+</sup>

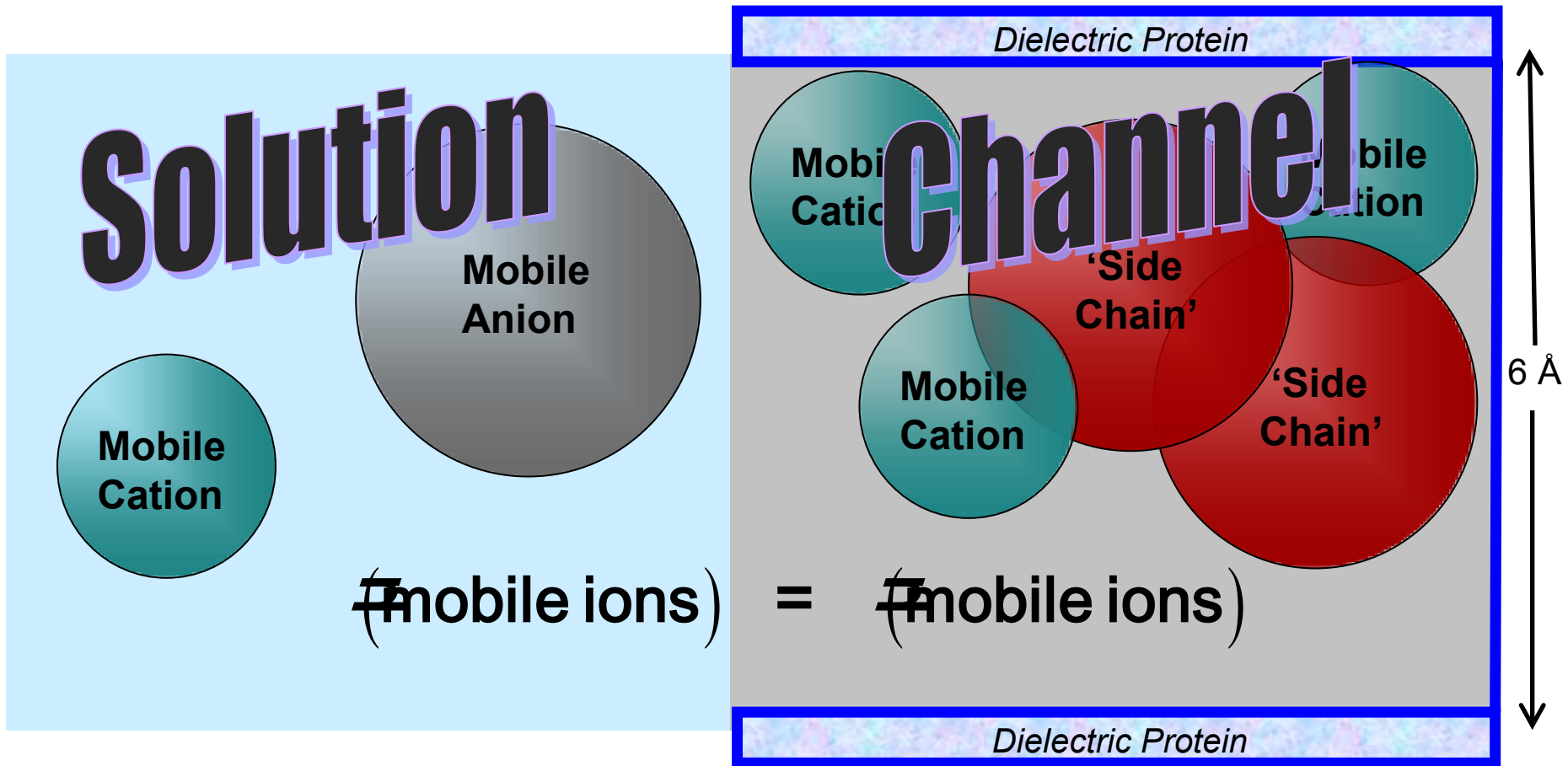
Ca<sup>++</sup>

O<sup>-1/2</sup>

Wolfgang Nonner



# ***Ion 'Binding' in Crowded Channel***



## ***Classical Donnan Equilibrium of Ion Exchanger***

*large mechanical forces*

Side chains move within channel to their equilibrium position of minimal free energy.

We compute the Tertiary Structure as the structure of minimal free energy.



# Solved with Metropolis Monte Carlo

**MMC Simulates Location of Ions**  
both the mean and the variance

**Produces Equilibrium Distribution**  
**of location**  
**of Ions and 'Side Chains'**

MMC yields Boltzmann Distribution with correct Energy, Entropy and Free Energy

## Other methods

give nearly identical results:

Equilibrium Multiscale

**MSA** (mean spherical approximation)

**SPM** (primitive solvent model)

**DFT** (density functional theory of fluids),

Non-equilibrium Multiscale

**DFT-PNP** (Poisson Nernst Planck)

**EnVarA...** (Energy Variational Approach)

*etc*

# Metropolis Monte Carlo

## Simulates Location of Ions

both the mean and the variance

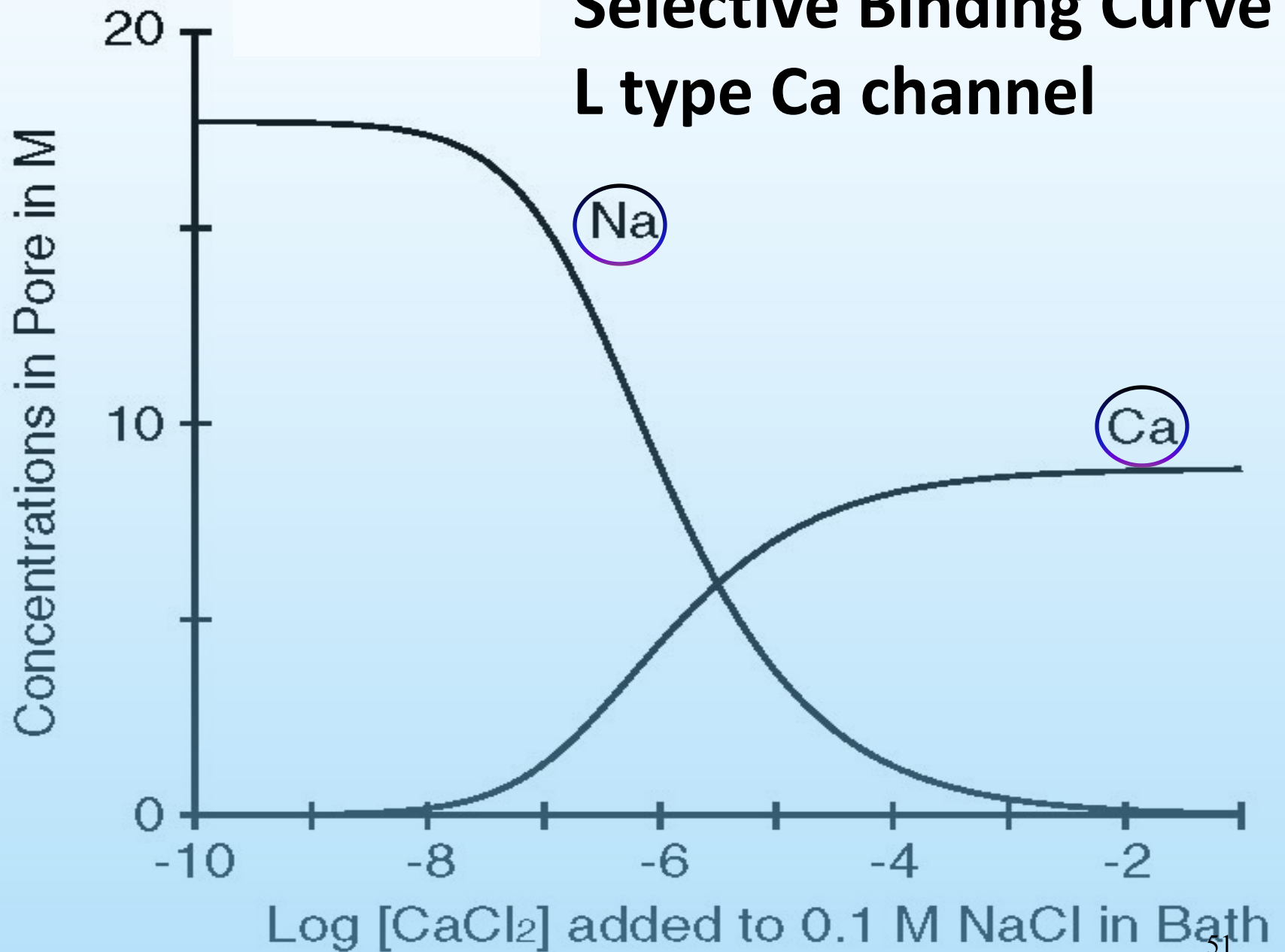
### Details:

- 1) Start with Configuration  $A$ , with computed energy  $E_A$
- 2) Move an ion to location  $B$ , with computed energy  $E_B$
- 3) If spheres overlap,  $E_B \rightarrow \infty$  and configuration is rejected
- 4) If spheres do not overlap,  $E_B \rightarrow 0$  and configuration is accepted
- 5) If  $E_B < E_A$  : accept new configuration.
- 6) If  $E_B > E_A$  : accept new configuration with probability  $\exp[-(E_A - E_B)/k_B T]$

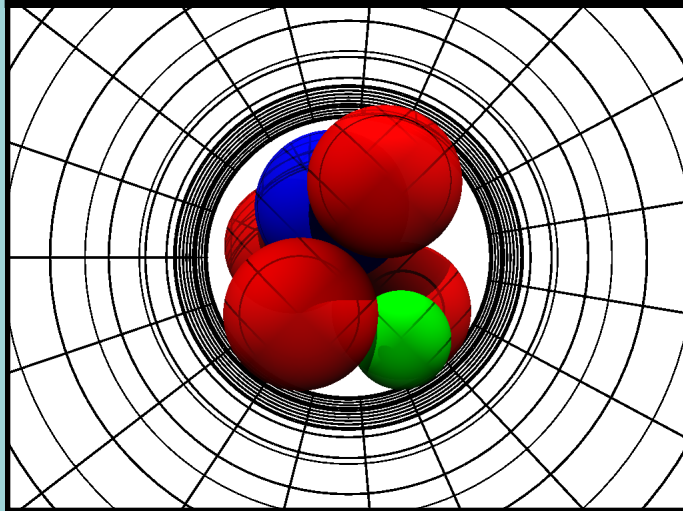
### Key idea

MMC chooses configurations with a Boltzmann probability and weights them evenly instead of choosing them from uniform distribution and then weighting them with  $\exp(-E/k_B T)$

# Selective Binding Curve L type Ca channel

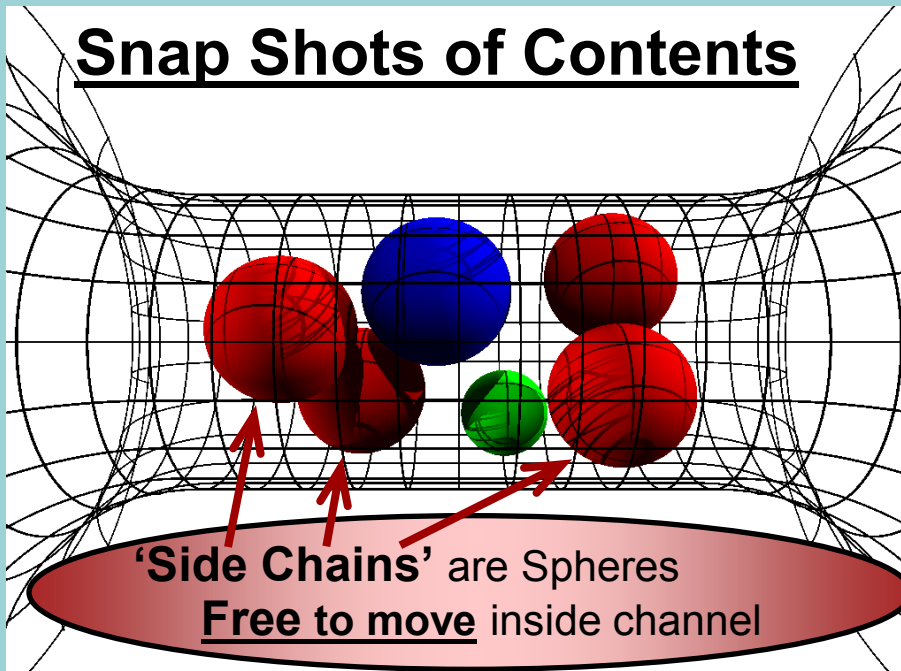


## Radial Crowding is Severe



6Å

## Snap Shots of Contents



# Crowded Ions

## Ion Diameters

'Pauling' Diameters

Ca<sup>++</sup>

1.98 Å

Na<sup>+</sup>

2.00 Å

K<sup>+</sup>

2.66 Å

## 'Side Chain' Diameter

Lysine K

3.00 Å

D or E

2.80 Å

Channel Diameter 6 Å

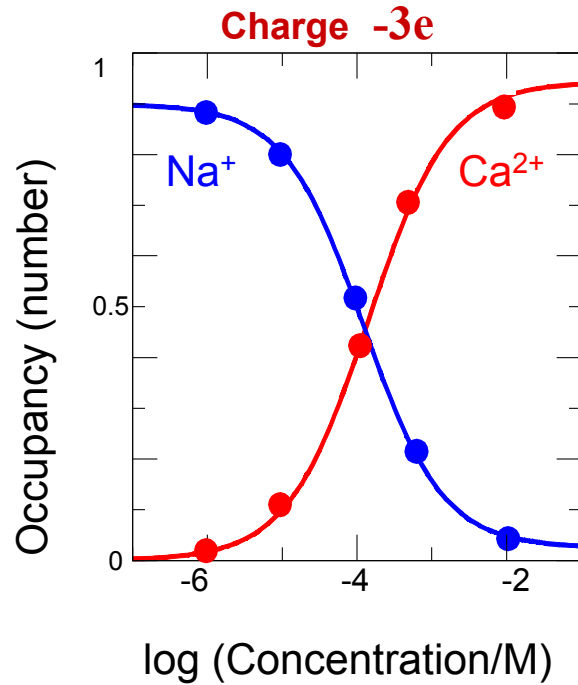
*Parameters are Fixed in all calculations  
in all solutions for all mutants*

Experiments and Calculations done at pH 8

52

# Ca Channel

E  
E  
E  
A



EEEE has full biological selectivity  
in similar simulations

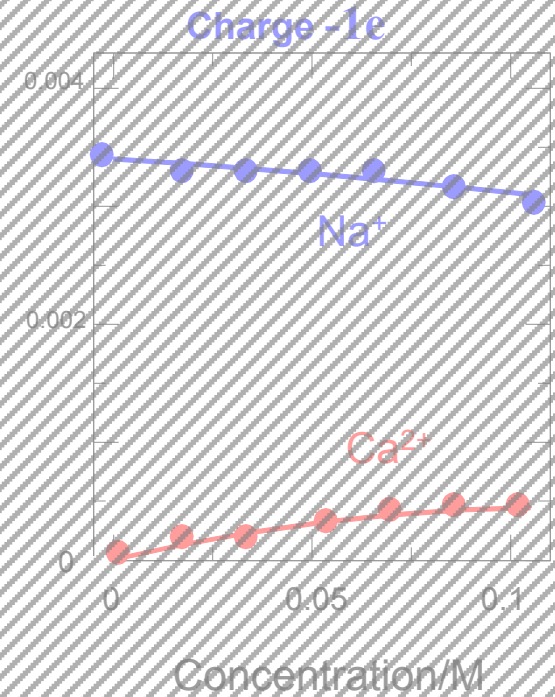
Mutation



Same Parameters

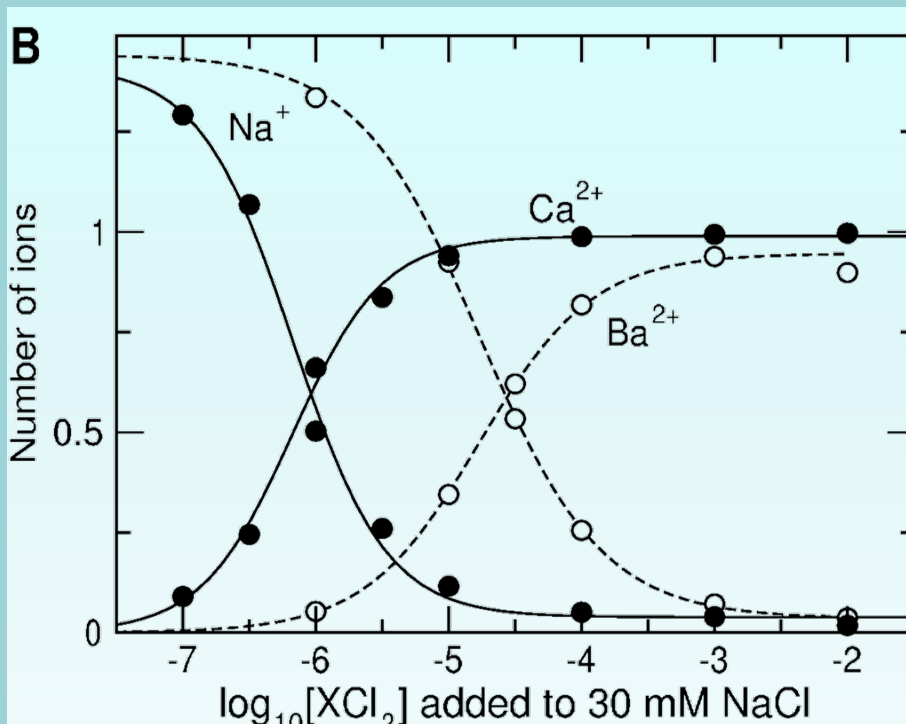
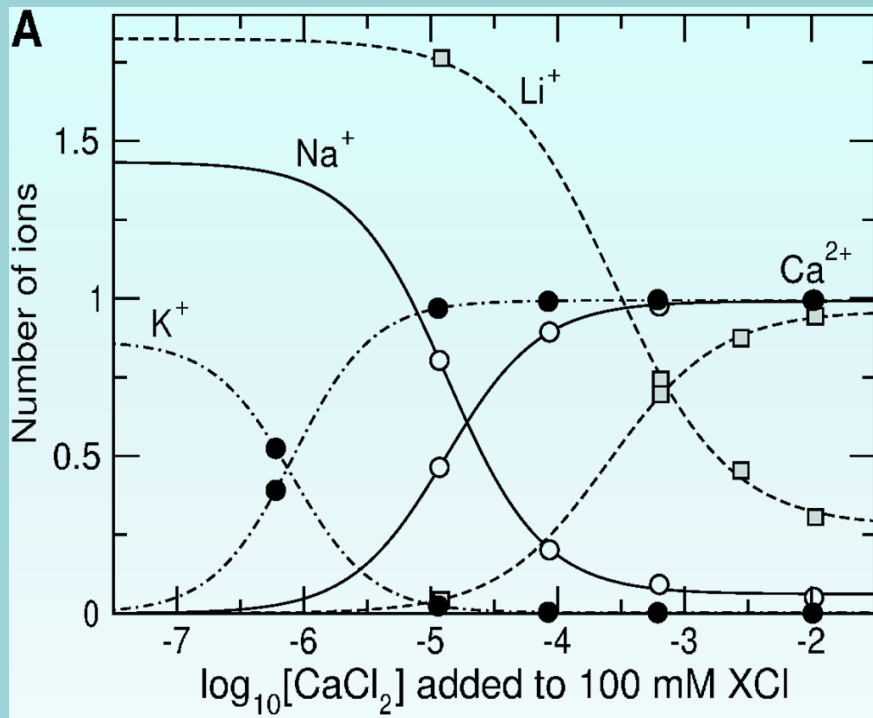
# Na Channel

D  
E  
K  
A



Boda, et al

# Na, K, Li, Ca, Ba Binding in Calcium Channel

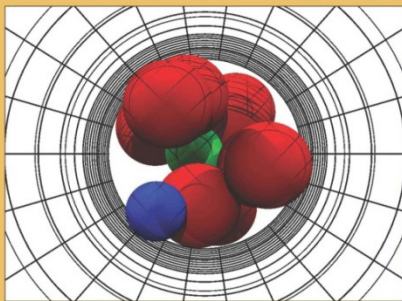
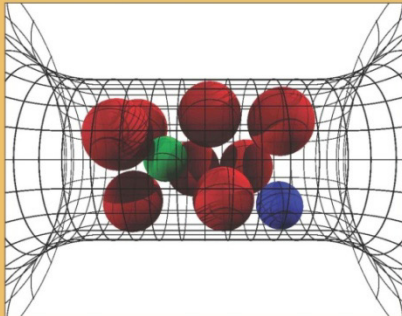


# Calcium Channel

*has been examined in ~35 papers, e.g.,*

# JGP

The Journal of General Physiology  
Vol 133 • No 5 • May 2009



[www.jgp.org](http://www.jgp.org)

- Nonner, W., D. P. Chen, and B. Eisenberg. 1998. Anomalous Mole Fraction Effect, Electrostatics, and Binding in Ionic Channels. *Biophysical Journal* 74:2327-2334.
- Nonner, W., L. Catacuzzeno, and B. Eisenberg. 2000. Binding and Selectivity in L-type Ca Channels: a Mean Spherical Approximation. *Biophysical Journal* 79:1976-1992.
- Nonner, W., D. Gillespie, D. Henderson, and B. Eisenberg. 2001. Ion accumulation in a biological calcium channel: effects of solvent and confining pressure. *J Physical Chemistry B* 105:6427-6436.
- Boda, D., W. Nonner, D. Henderson, B. Eisenberg, and D. Gillespie. 2008. Volume exclusion in calcium selective channels. *Biophys. J.:biophysj*.107.122796.
- Boda, D., M. Valisko, B. Eisenberg, W. Nonner, D. Henderson, and D. Gillespie. 2006. Effect of Protein Dielectric Coefficient on the Ionic Selectivity of a Calcium Channel. *Journal of Chemical Physics* 125:034901.
- Boda, D., T. Varga, D. Henderson, D. Busath, W. Nonner, D. Gillespie, and B. Eisenberg. 2004. Monte Carlo simulation study of a system with a dielectric boundary: application to calcium channel selectivity. *Molecular Simulation* 30:89-96.
- Boda, D., M. Valisko, B. Eisenberg, W. Nonner, D. Henderson, and D. Gillespie. 2007. The combined effect of pore radius and protein dielectric coefficient on the selectivity of a calcium channel. *Physical Review Letters* 98:168102.

Most of the papers are available at

[ftp://ftp.rush.edu/users/molebio/Bob\\_Eisenberg/Reprints](ftp://ftp.rush.edu/users/molebio/Bob_Eisenberg/Reprints)

<http://www.phys.rush.edu/RSEisenberg/physioeis.html>

# Selectivity

comes from

**Electrostatic Interaction**

and

**Steric Competition for Space**



Repulsion

Location and Strength of Binding Sites  
Depend on Ionic Concentration and  
Temperature, etc

***Rate Constants are Variables***



*for*  
***Biologists:***  
***a Word Picture***

**How does Calcium Selectivity Work?**  
**qualitatively**

# How does it work qualitatively?

*(for biologists)*

**2 Ca<sup>++</sup>**

are

**LESS CROWDED**

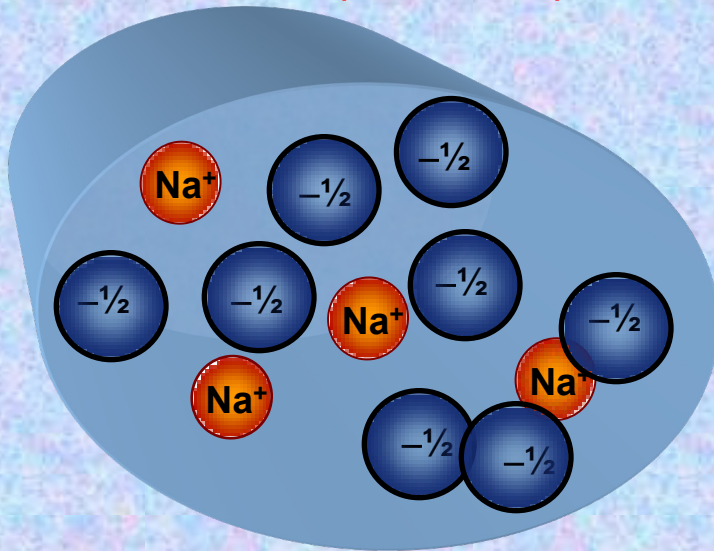
**than 4 Na<sup>+</sup>**

# Selectivity from Crowded Charges

2  $\text{Ca}^{++}$  are less crowded than 4  $\text{Na}^+$

## Ca Channel Filled with $\text{Na}^+$

(not to scale)



### Channel Protein

Glutamate Oxygens = 4e

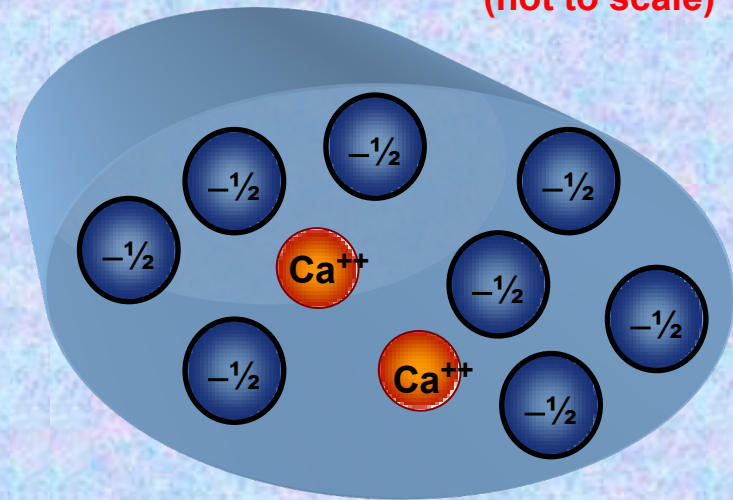
8 of  $-\frac{1}{2}$  charge each

Volume  $0.38 \text{ nm}^3$

Dielectric Constant 64

## Ca Channel Filled with $\text{Ca}^{++}$

(not to scale)



### Outside the Filter

Bulk Solution

$\text{NaCl}$  and  $\text{CaCl}_2$

**Ionic Selectivity in Protein Channels**  
**Crowded Charge Mechanism**

**4 Negative Charges**  
**of glutamates of protein**

**DEMAND**

**4 Positive Charges**  
**nearby**

either 4 Na<sup>+</sup> or 2 Ca<sup>++</sup>

# Ionic Selectivity in Protein Channels

## Crowded Charge Mechanism

*Simplest Version: MSA*

2  $\text{Ca}^{++}$  are **LESS CROWDED** than 4  $\text{Na}^+$ ,

$\text{Ca}^{++}$  **SHIELDS BETTER** than  $\text{Na}^+$ , so

**Protein Prefers  $\text{Ca}^{++}$**

*because*

**$\text{Ca}^{++}$  is less crowded**

# What does the protein do?

Channel and Contents  
form a

**Self-Organized Structure**

with Side Chains at position of  
Minimum Free Energy

Protein Fits the Substrate

**“Induced Fit Model of Selectivity”**

# What does the protein do?

*(for biologists)*

Certain **MEASURES** of structure are  
Powerful **DETERMINANTS** of Function

e.g., Volume, Dielectric Coefficient, etc.

Induced Fit Model of Selectivity

Atomic Structure is not pre-formed

Atomic Structure is an important output of the simulation

# What does the protein do?

Protein maintains

Mechanical Forces\*

Volume of Pore

Dielectric Coefficient/Boundary

Permanent Charge

*\* Driving force for conformation changes ??*



Binding Sites\* are **outputs**  
of our Calculations

## **Induced Fit Model of Selectivity**

**Our model has no preformed  
structural binding sites**

but

**Selectivity is very Specific**

\*Selectivity is in the Depletion Zone,  
NOT IN THE BINDING SITE  
of the DEKA Na Channel

# Challenge

from leading biophysicists

**Walter Stühmer and Stefan Heinemann**

Max Planck Institutes, Göttingen, Leipzig

**Can a physical theory explain the mutation  
Calcium Channel into Sodium Channel?**



# DEKA Sodium Channel

has very different properties from Ca channel,

e.g., 'binding' curve,  
Na<sup>+</sup> vs Ca<sup>++</sup> selectivity  
Na<sup>+</sup> vs K<sup>+</sup> selectivity

# Sodium Channel

specifically, the

## DEKA Sodium Channel 6 Å

Aspartate

Glutamate

Lysine

Alanine

D

E

K

A

Acid

Acid

Basic

Aliphatic

Negative

Negative

Positive

Neutral

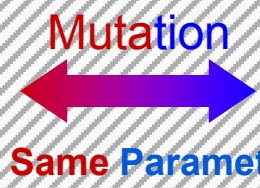
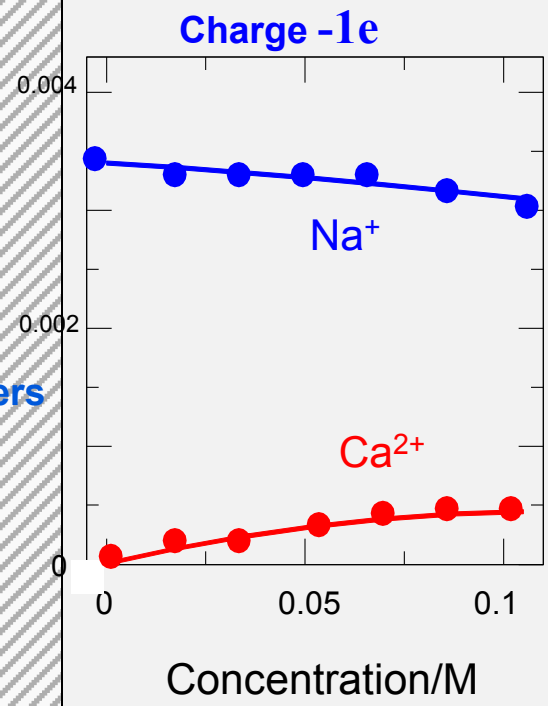
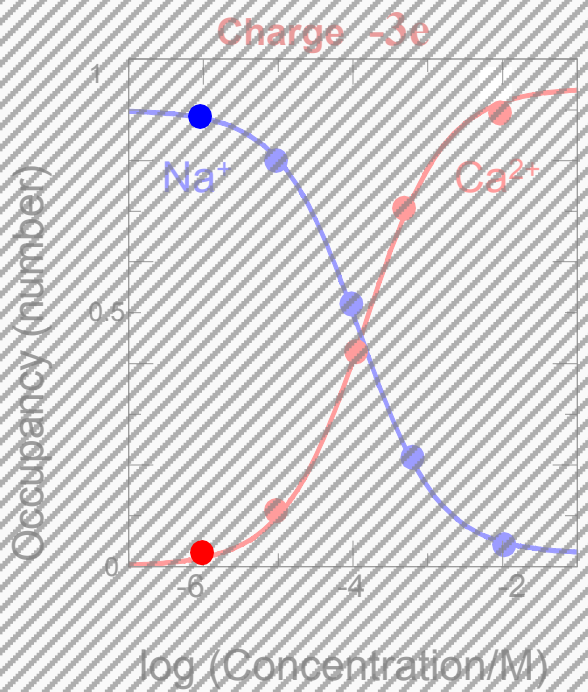
QUALITATIVELY DIFFERENT Properties from the Calcium Channel

# Ca Channel

# Na Channel



Same Parameters



Same Parameters

EEEE has full biological selectivity in similar simulations

**Nothing was changed**  
from the  
*EEEE Ca channel*  
except the amino acids

Calculated DEKA Na Channel  
Selects

$\text{Ca}^{2+}$  vs.  $\text{Na}^{+}$  and also  $\text{K}^{+}$  vs.  $\text{Na}^{+}$

# Selectivity

comes from

**Electrostatic Interaction**

and

**Steric Competition for Space**



Repulsion

Location and Strength of Binding Sites  
Depend on Ionic Concentration and  
Temperature, etc

***Rate Constants are Variables***

# Miracle

**We can actually compute the  
Structures that determine Selectivity**



# New Miracle???

**Can *EnVarA* actually compute the  
Function of these systems?**

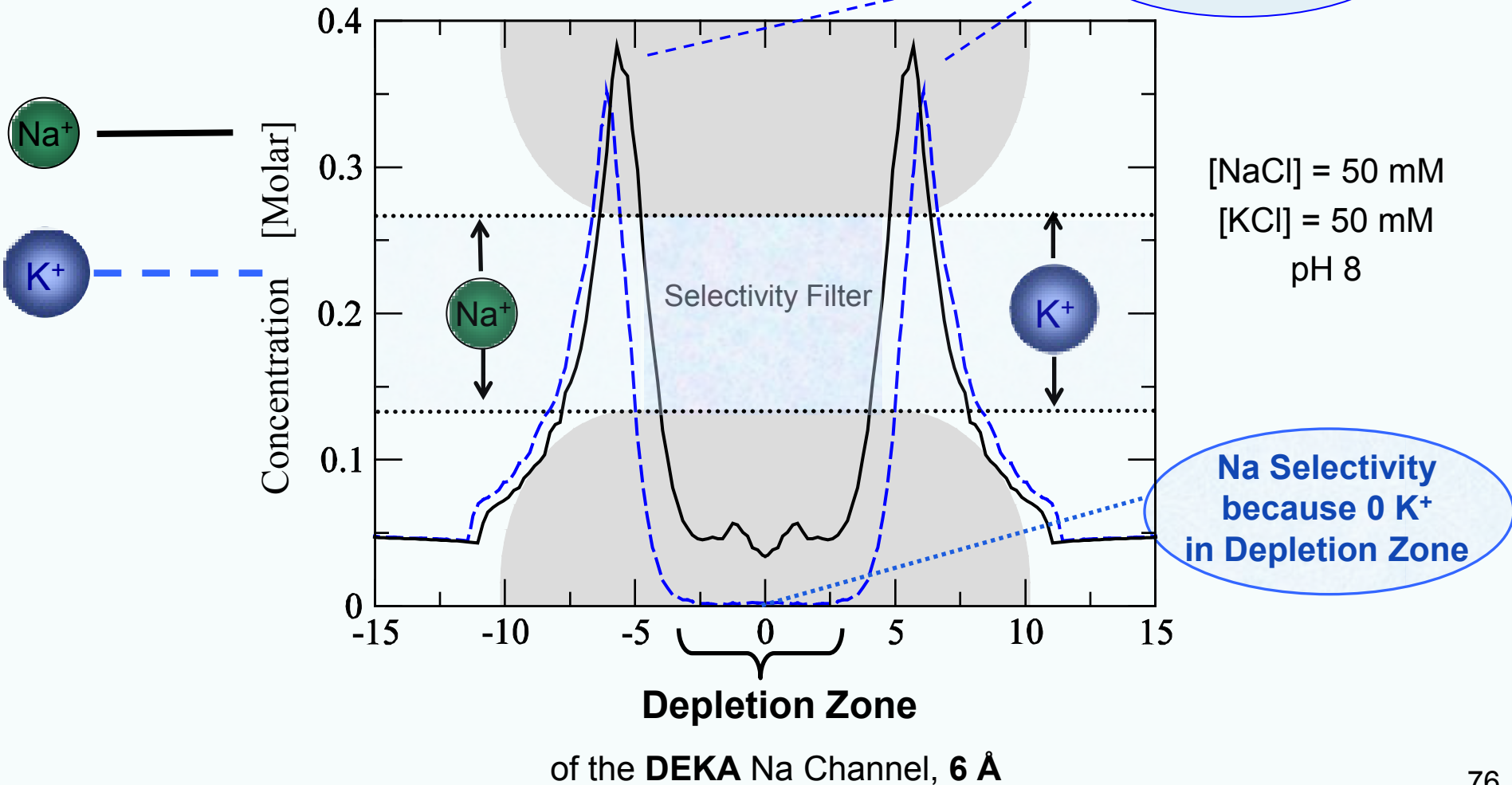


# How?

How does the DEKA Na Channel Select  $\text{Na}^+$  vs.  $\text{K}^+$  ?

# Size Selectivity is in the Depletion Zone

## Na<sup>+</sup> vs. K<sup>+</sup> Occupancy

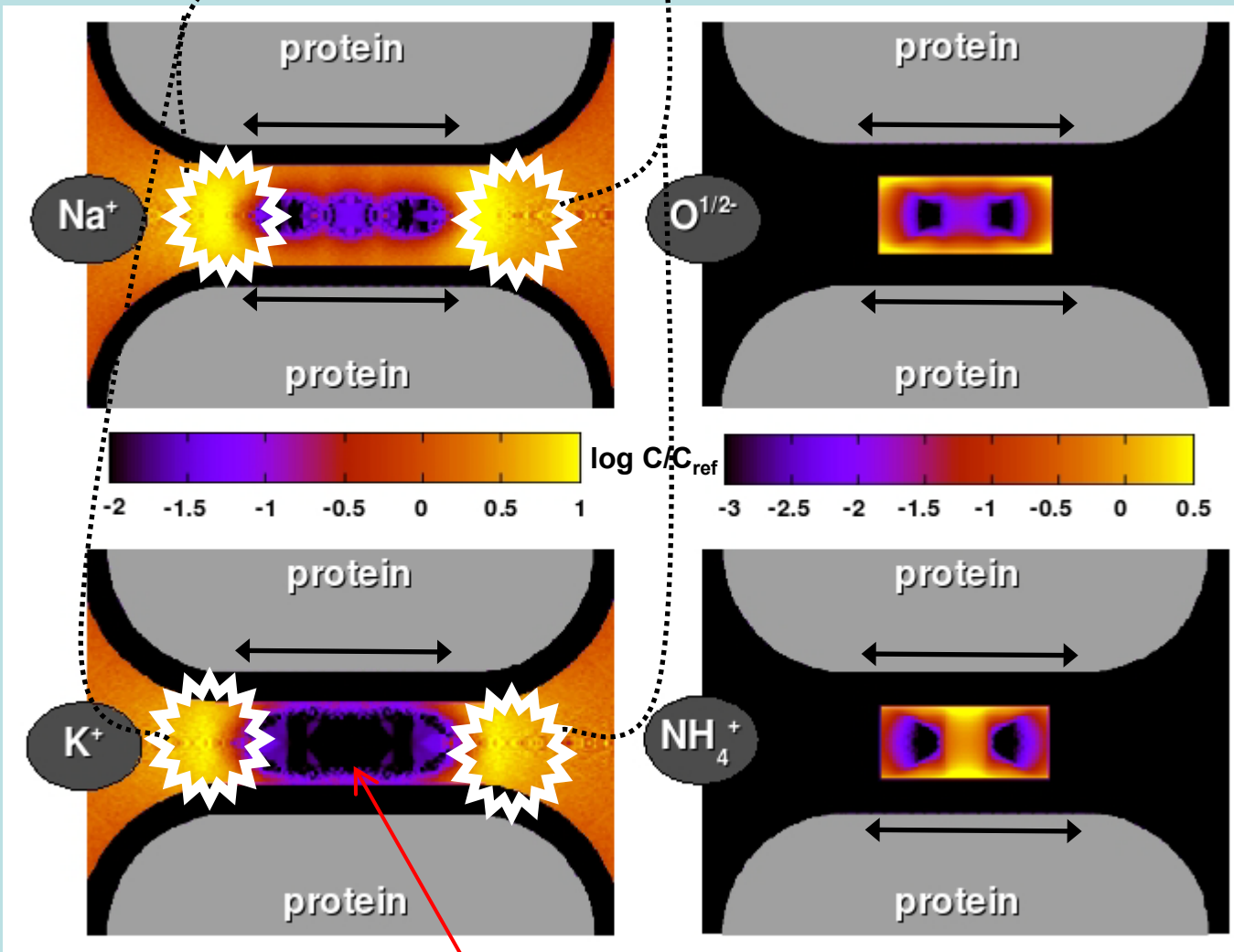


# Size Selectivity

# Binding Sites

\*Binding Sites are outputs of our INDUCED FIT Model of Selectivity, *not structural inputs*

[NaCl] = [KCl] = 50 mM

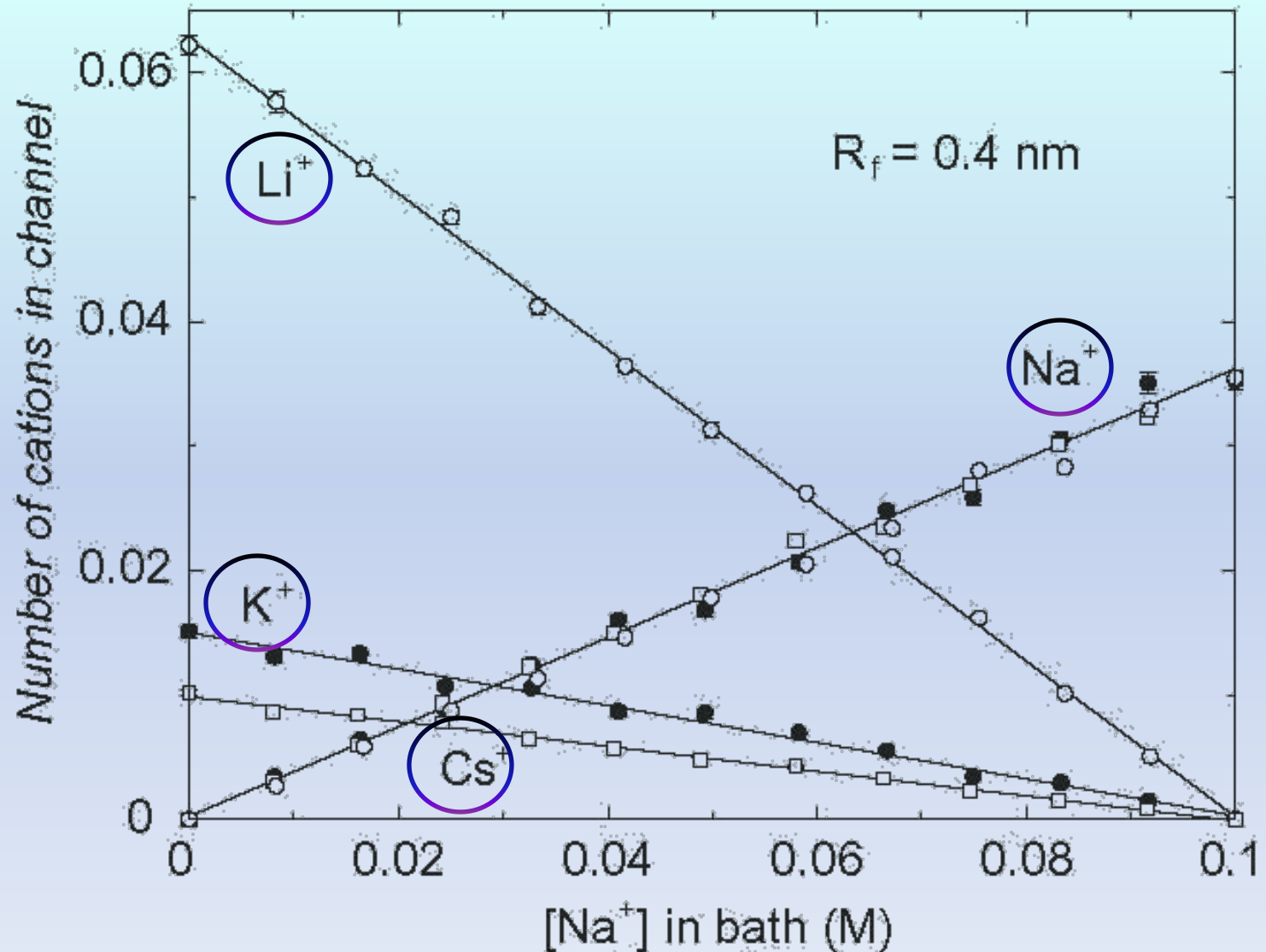


Na vs K Size Selectivity is in **Depletion Zone**

BLACK = Depletion=0

Ion Diameter	
Ca <sup>++</sup>	1.98 Å
Na <sup>+</sup>	2.00 Å
K <sup>+</sup>	2.66 Å
'Side Chain' Diameter	
NH <sub>4</sub> <sup>+</sup>	3.00 Å
Lys or K	pH 8
O <sup>1/2-</sup> D or E	2.80 Å
	pH 8
Na Channel DEKA 6 Å	

# Na, K, Li, Cs Binding in Sodium channel



# What do the Variables do?

What happens  
if we  
**Vary Diameter**  
and  
**Vary Dielectric Coefficient?**

Inverse Problem  
**We discover**  
**Orthogonal Control Variables\***  
in simulations of the Na channel,  
but not the Ca channel.

*\*These emerge as outputs. They are not inputs.*



# Control Variables

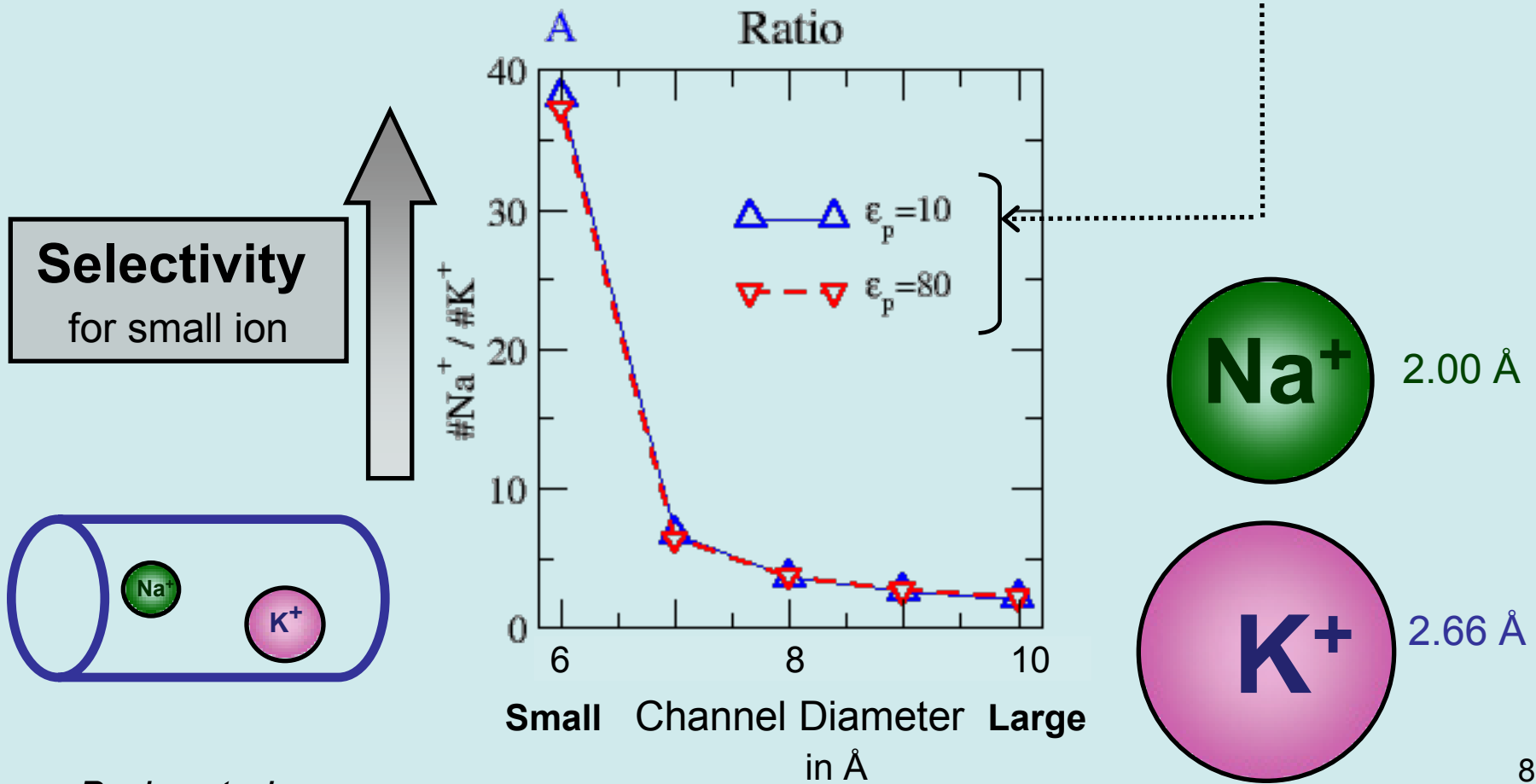
Selectivity  $Na^+$  vs  $K^+$

## Selectivity Depends on Structure

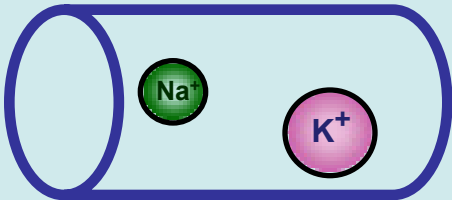
Depends **STEEPLY** on channel diameter

Depends only on channel diameter

# Na<sup>+</sup> vs K<sup>+</sup> (size) **Selectivity** (*ratio*) Depends on Channel Size, *not* Protein Dielectric Coefficient\*



**Selectivity**  
for small ion



Boda, et al

\*in DEKA Na Channel

# Control Variables

Conductance of DEKA Na<sup>+</sup> channel

## Conductance Depends Steeply on Dielectric

Contents of Channel depend only on dielectric

*but*

Selectivity does not depend on Dielectric

Selectivity depends **only** on Structure

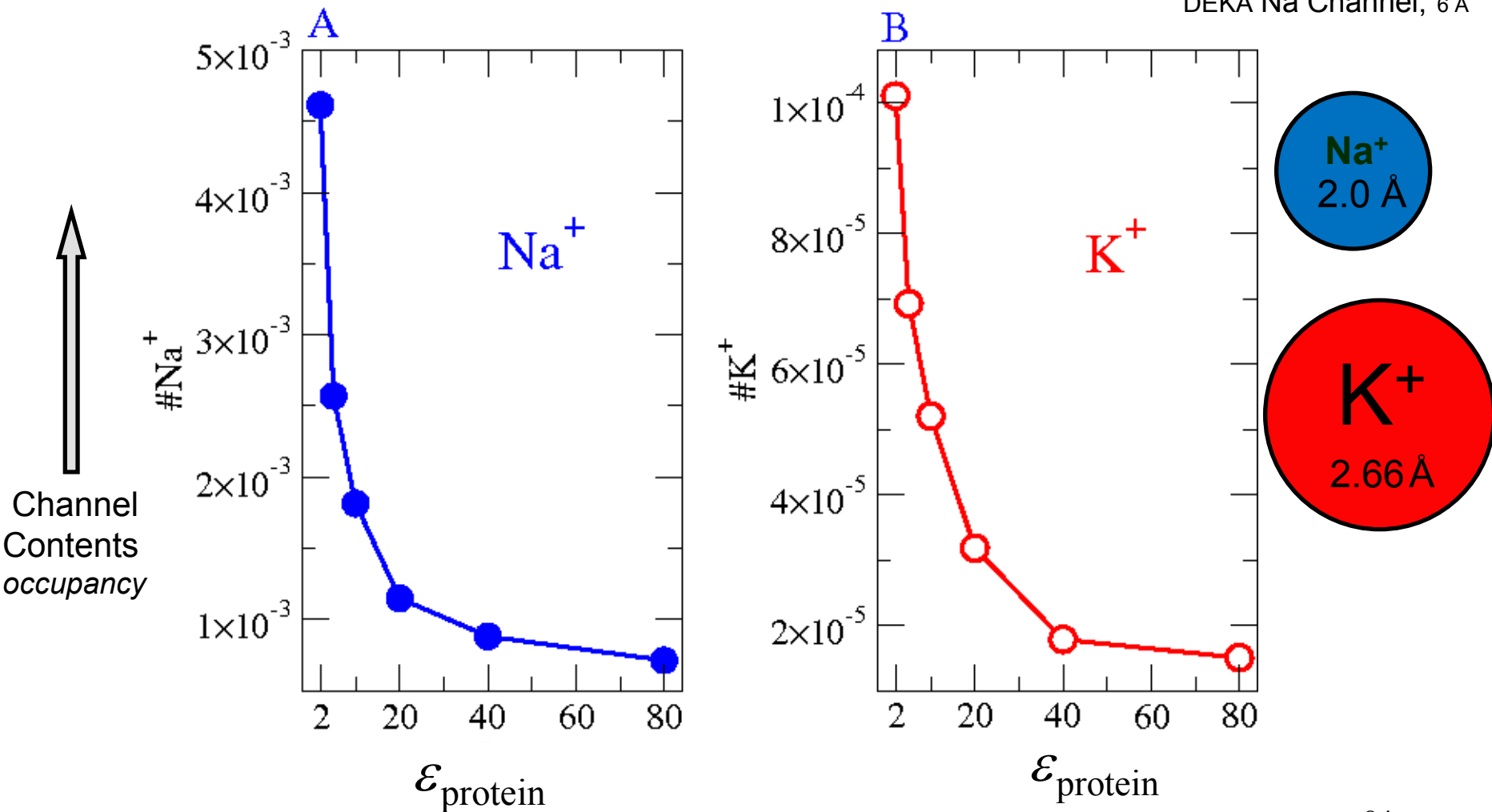
Control Variable

# Channel Contents (occupancy)

depends on

## Protein Polarization (dielectric)

DEKA Na Channel, 6 Å



**Static  
Structure**

**Channel Diameter**

and

**Dielectric Coefficient**

emerge as

**Orthogonal Control Variables\***

in simulations of the Na channel,  
but not the Ca channel.

**Dynamic  
Structure**

*\*These emerge as outputs. They are not inputs.*

Supplementary Material

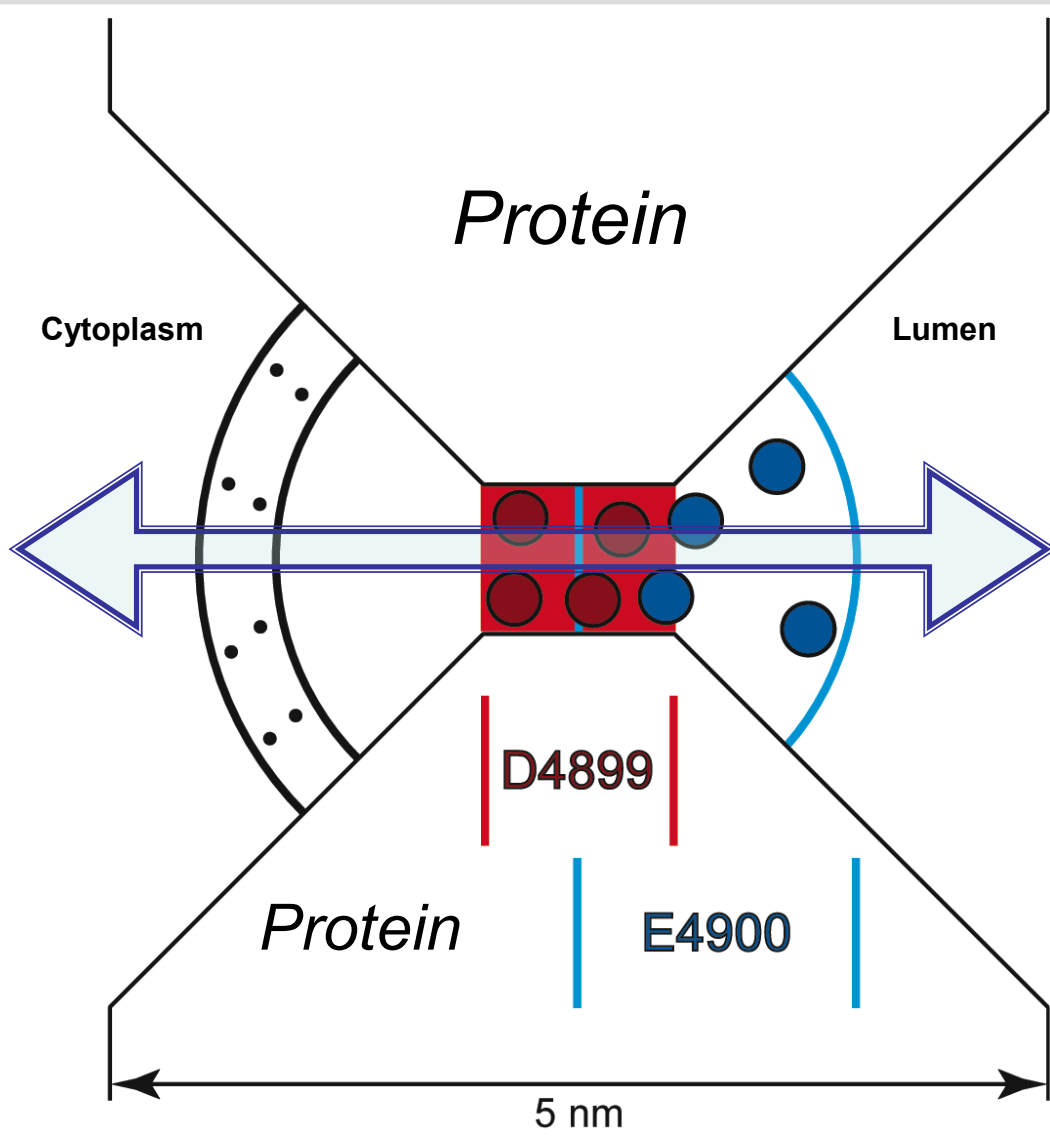
# RyR Channel: Current Voltage Curves

*Best Evidence is from the*  
***RyR Receptor***

**Gillespie, Meissner, Le Xu, et al,**  
*not Bob Eisenberg*

- **More than 120 combinations of solutions & mutants**
- **7 mutants with significant effects fit successfully**

# The Geometry



## Selectivity Filter

- is 10 Å long and 8 Å in diameter
- confines four **D4899** negative amino acids.

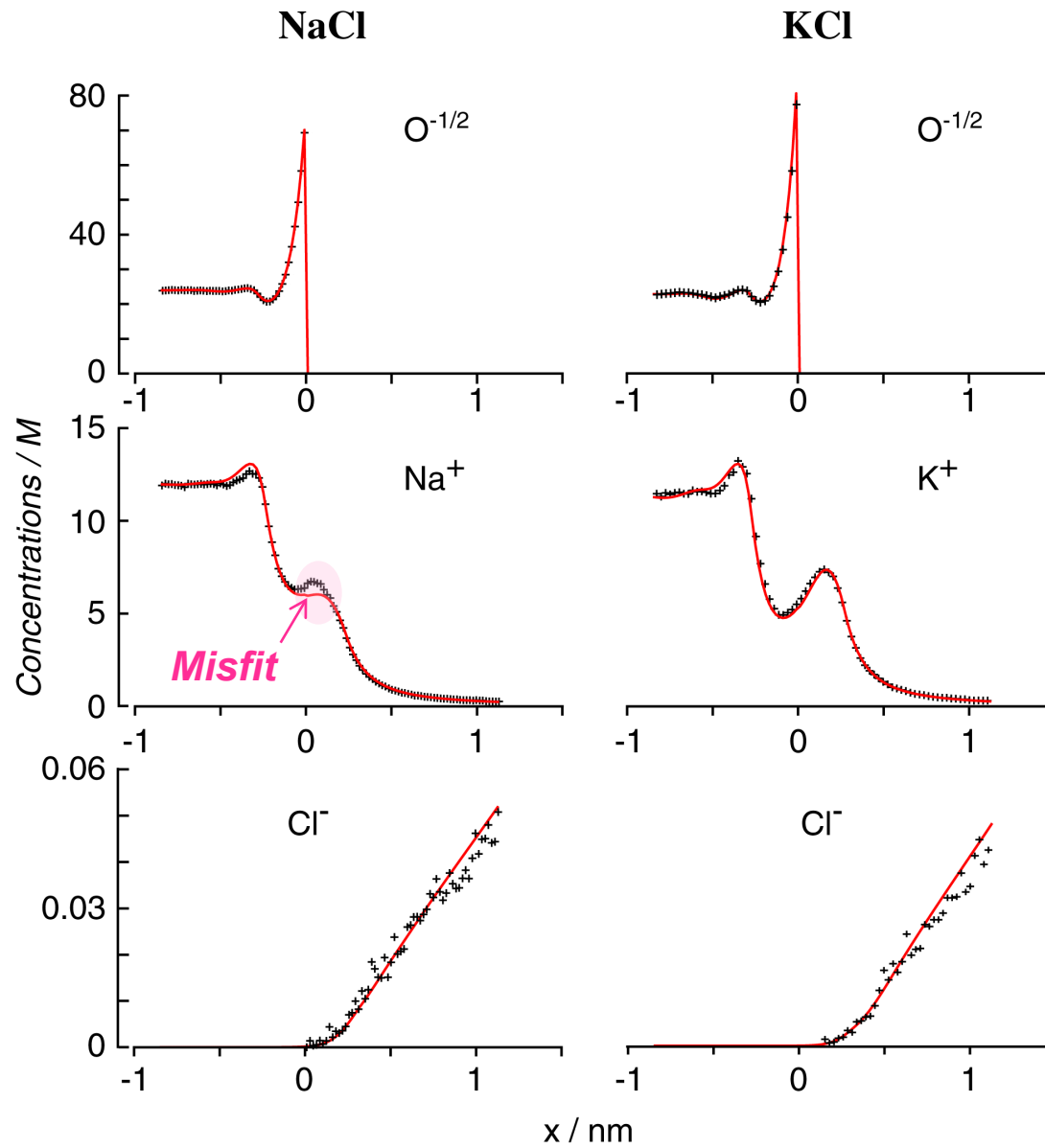
Four **E4900** positive amino acids are on luminal side, overlapping D4899.

**Cytosolic distributed charge**

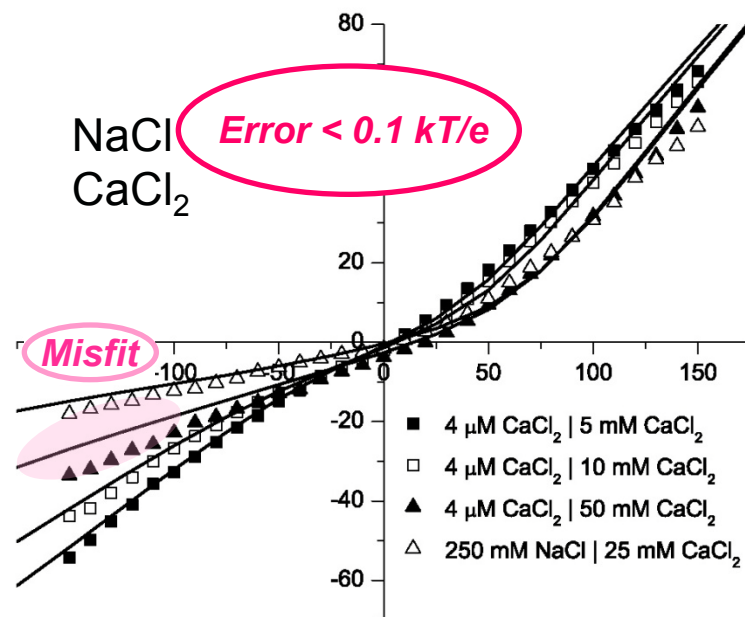
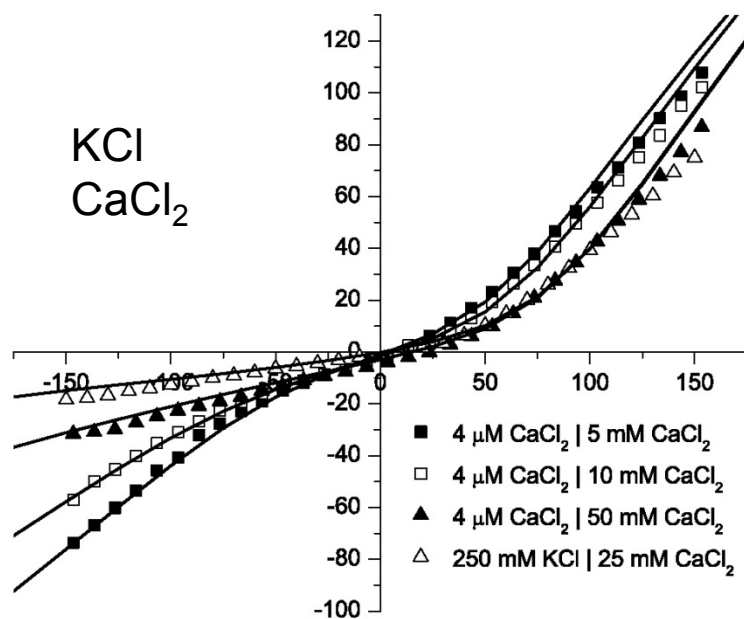


# DFT/PNP vs Monte Carlo Simulations

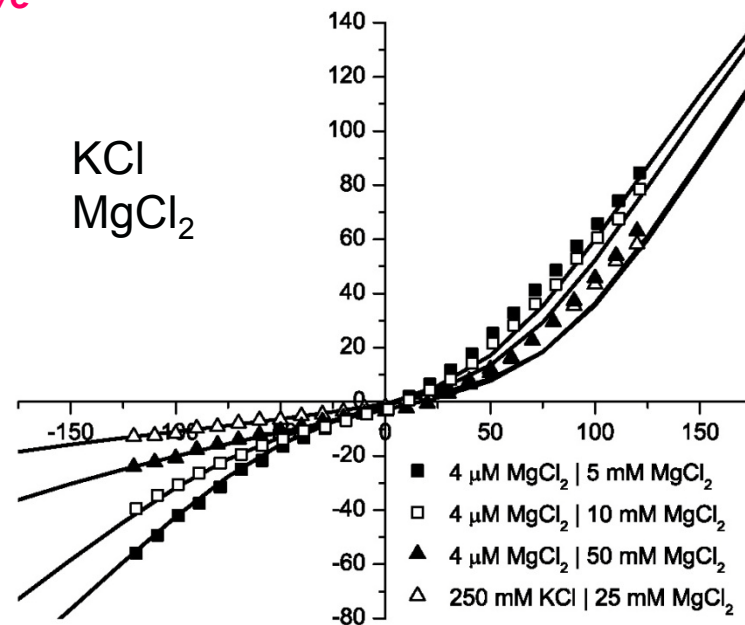
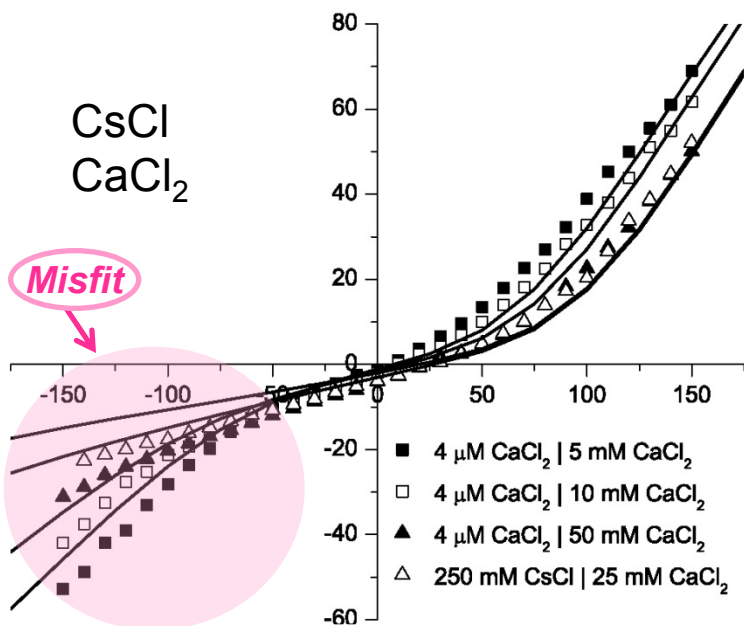
## Concentration Profiles

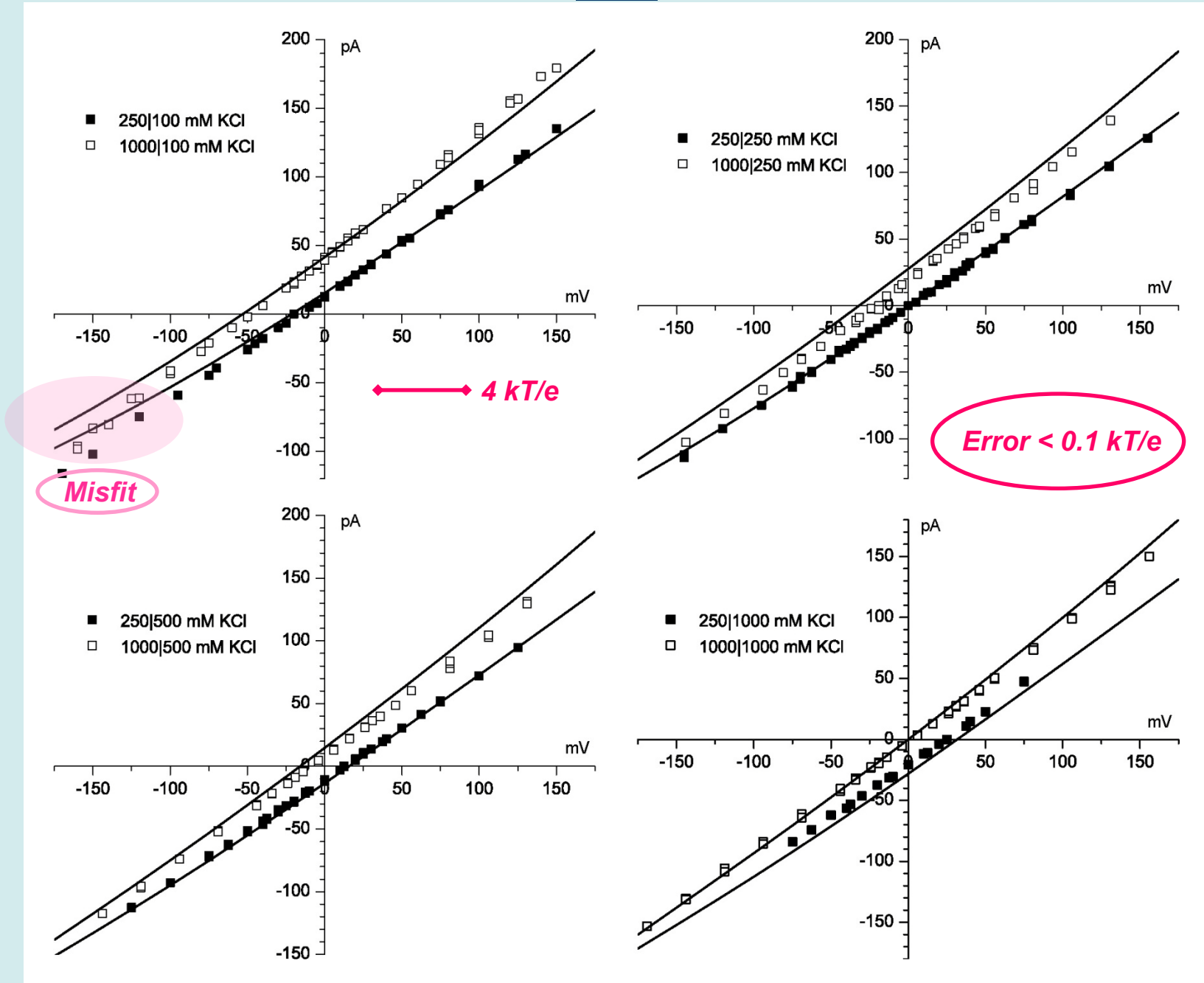


# Divalents



↔ 2 kT/e

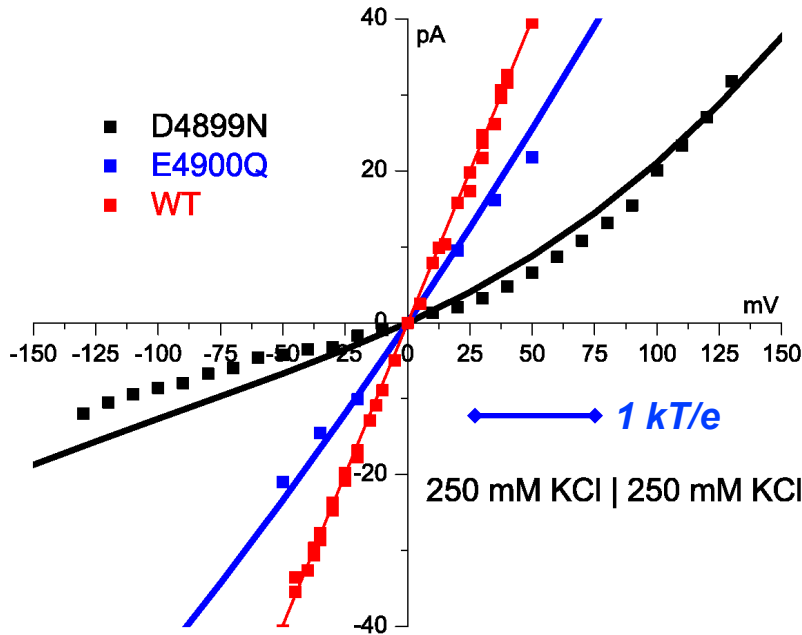




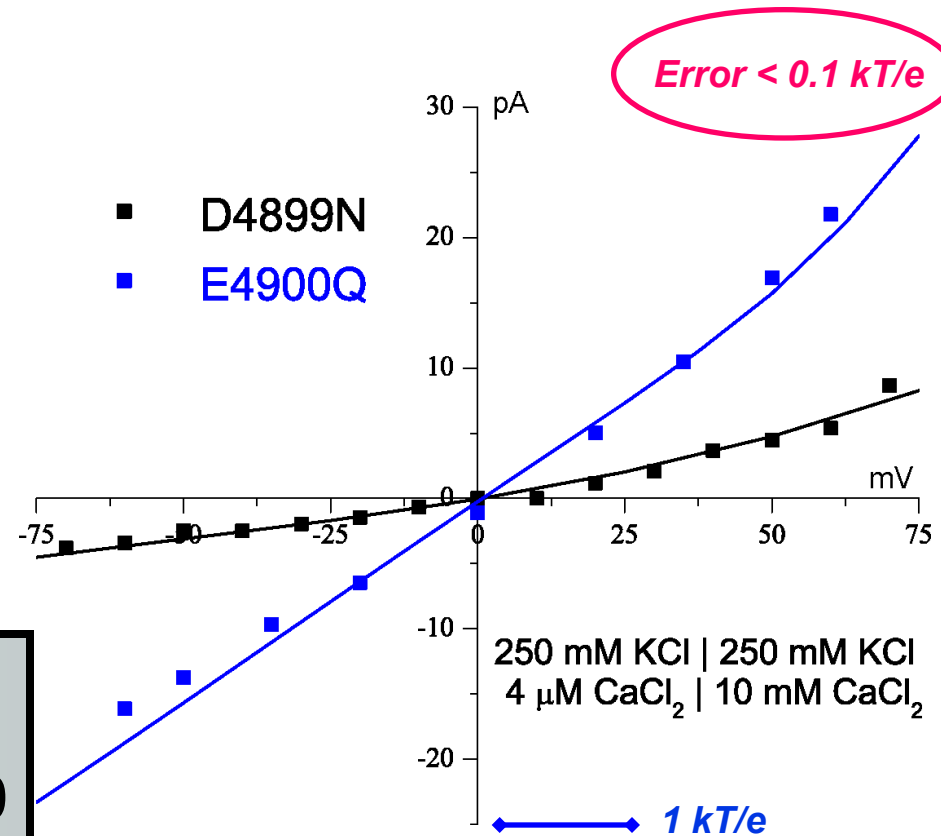
# Theory fits Mutation with Zero Charge

*No parameters adjusted*

## Theory Fits Mutant in K



## Theory Fits Mutant in K + Ca



**Protein charge density**  
wild type\* **13 M**  $\Rightarrow$  **0 M** in D4899

Water is 55 M

*\*some wild type curves not shown, 'off the graph'*

Gillespie *et al*

*J Phys Chem* 109 15598 (2005)



***Vaccination  
against  
Traditional Models***

**Traditional Biochemistry  
and  
Traditional Molecular Dynamics  
Assume  
Ideal Solutions**

Ions in Water and Life are NOT ideal

**Life Occurs in ~130 mM salt solutions**

Ions in Water are the Liquid of Life

**No gas phase models of  
traditional channel biochemistry**

***Liquids are not Gases***

***Rate Constants are Variables***

**No discussions of individual trajectories of  
Structural Biologists**

***Counting and Statistics are essential***



# **Computation Starts From Crystal Structure** *when available* *but*

**Crystal Structures cannot determine Selectivity**  
*because*

- 1) Crystal Structures are measured in only one unphysiological solution**
- 2) Crystal Structures are not accurate enough**
- 3) Crystal Structures do not give entropy**

# Selectivity

Depends Sensitive on Self-organized Structure and their Flexibility

Induced Structure is Different in Different Solutions

so

**Structure must be Computed!**

**Rate constants are variables that change dramatically with conditions**



# Supplementary Material

Ions in Water are the Liquid of Life. They are not ideal solutions

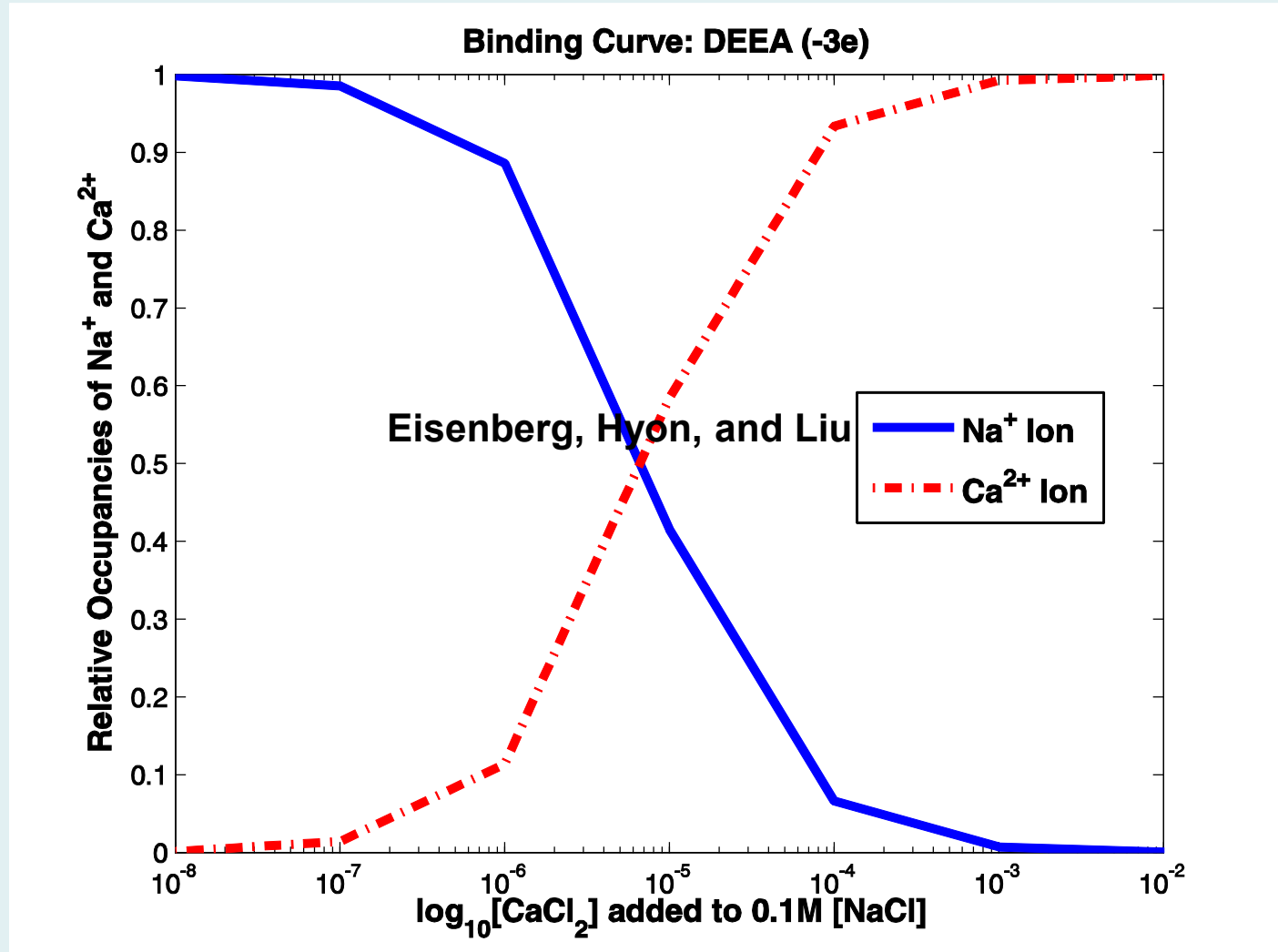
**Chemically Specific Properties**  
of Ionic Solutions come from  
**Interactions**

*Molecular Dynamics Force Fields are Calibrated  
assuming no interactions with concentrations*

**Force Fields must be REcalibrated  
in each Biological Solution**

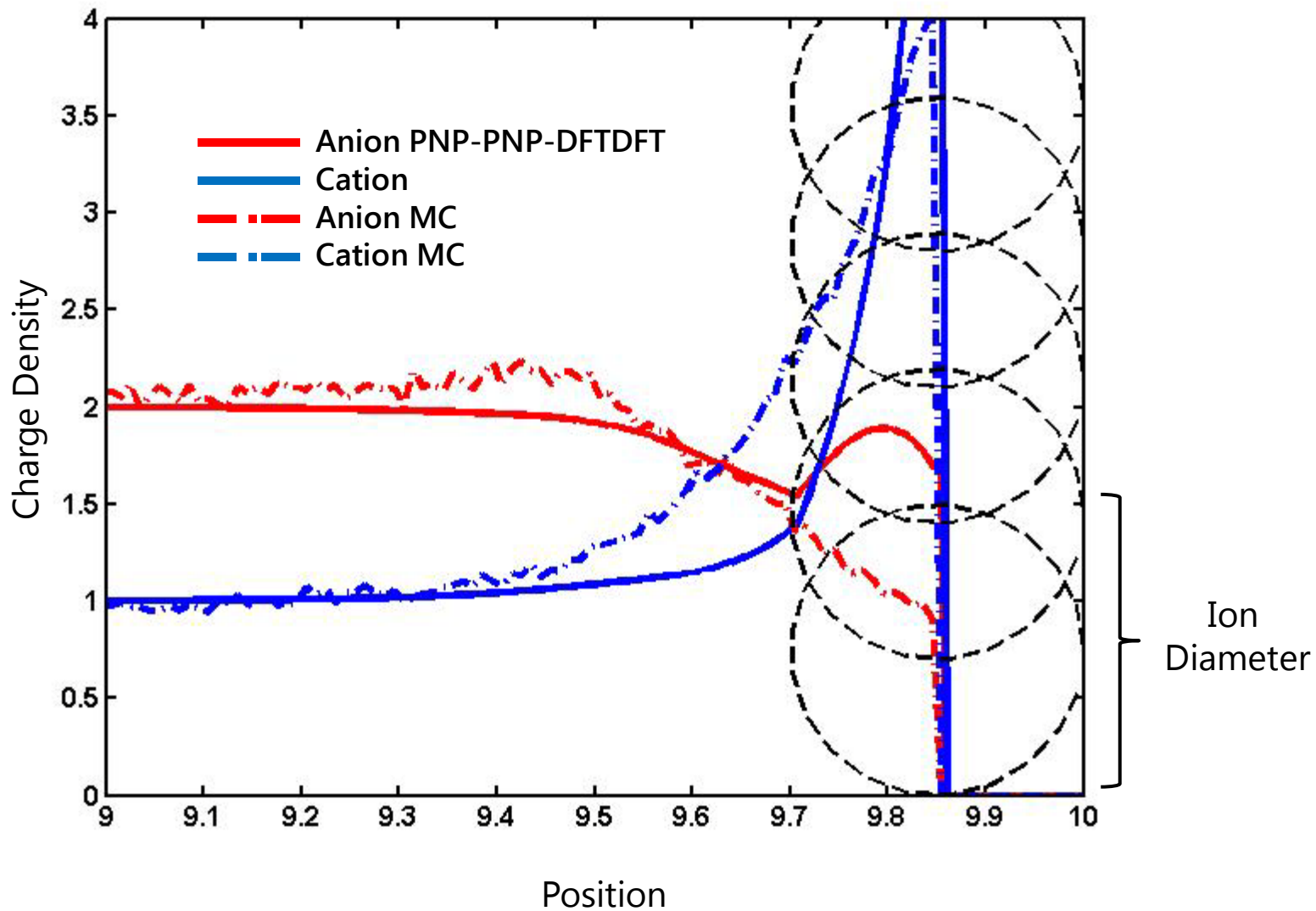
# Ca<sup>2+</sup> and Na<sup>+</sup> Binding Curves

## DEEA Calcium Channel



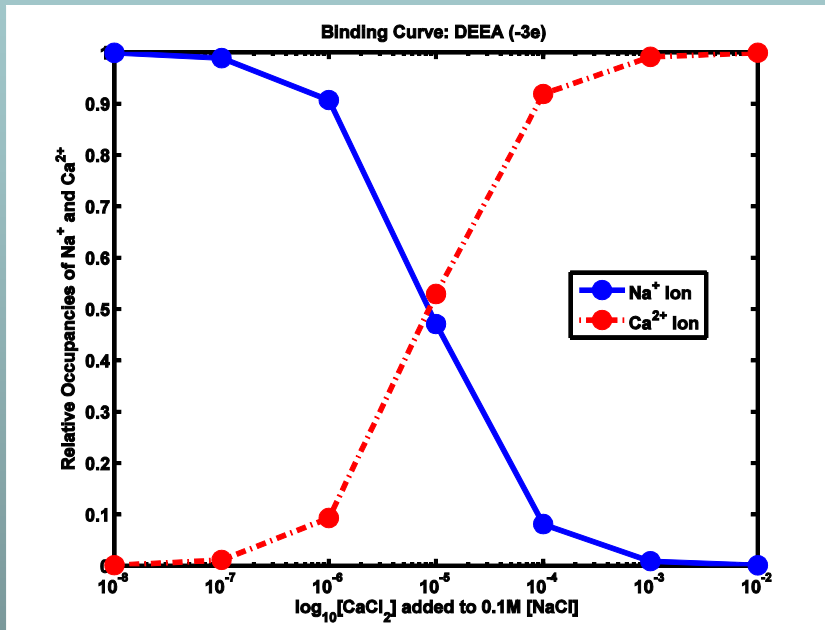
# Layering: Classical Interaction Effect

Comparison between PNP-DFT and MC

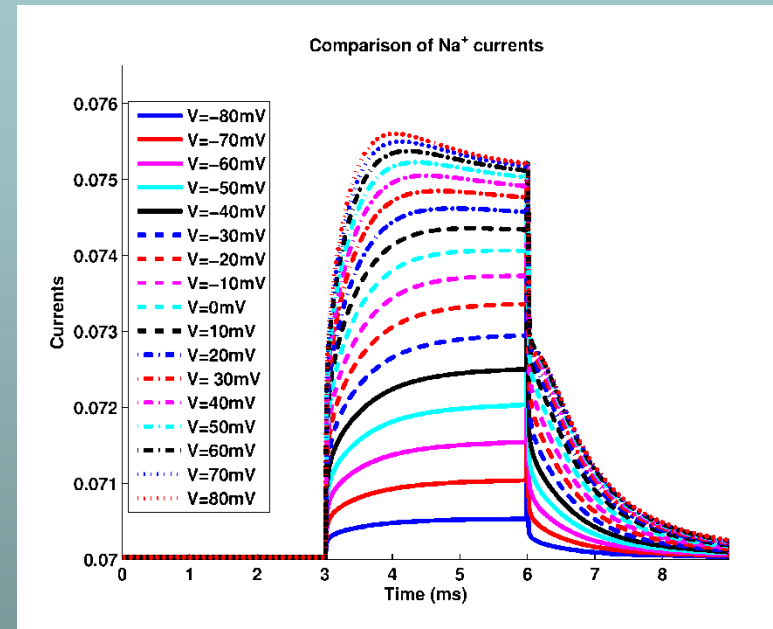


# Nonequilibrium Computations with Variational Field Theory *EnVarA*

Binding Curves



Current Voltage Time Curves







# Inactivation is Important

**Many diseases produced by changes in details of inactivation.**

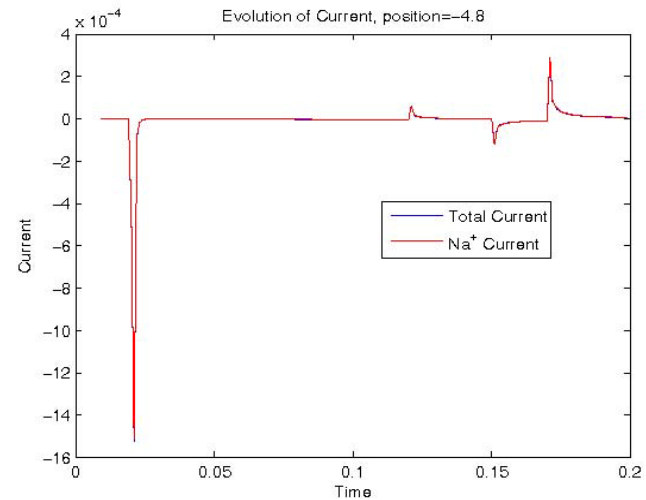
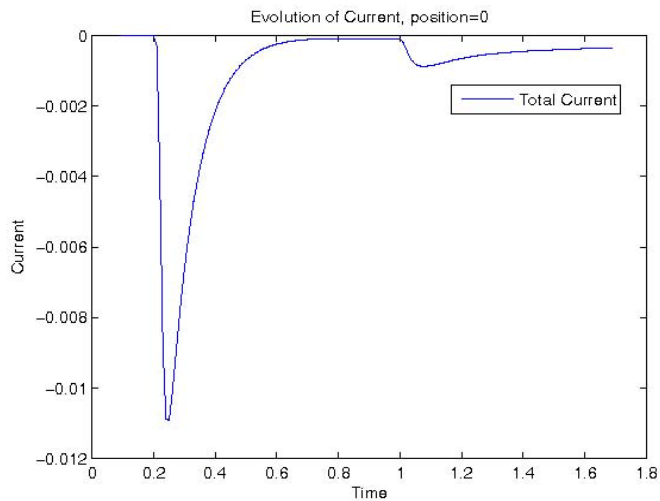
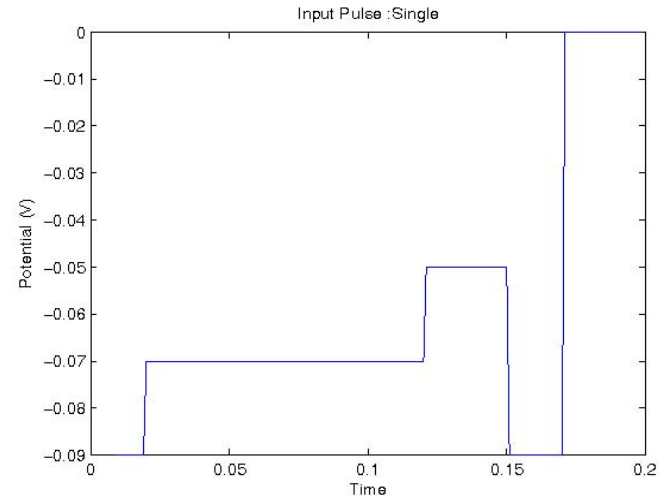
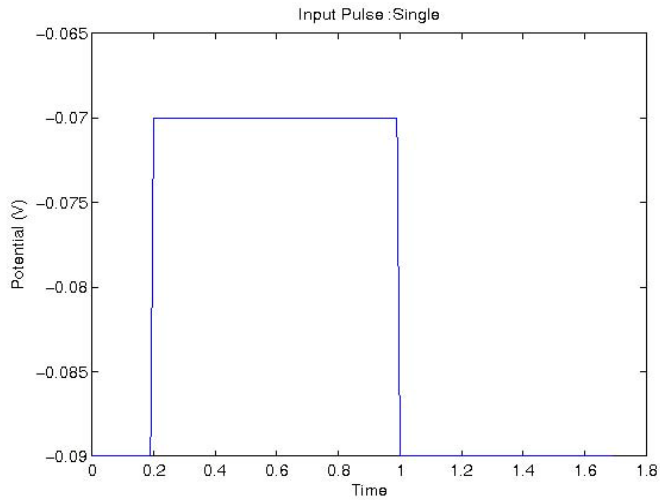
**Energetics of Brain determined by details of inactivation\***

**Energetics determined by time overlap of Na and K currents**

**\*Alle, Roth, and Geiger. Science (2009) 325:1405-8.**

# Sodium Conductance and Inactivation

## Variational Computation in Fixed Structure



# Energetic Variational Analysis

*EnVarA*

Chun Liu, Yunkyong Hyon and Bob Eisenberg

**New Interpretations**

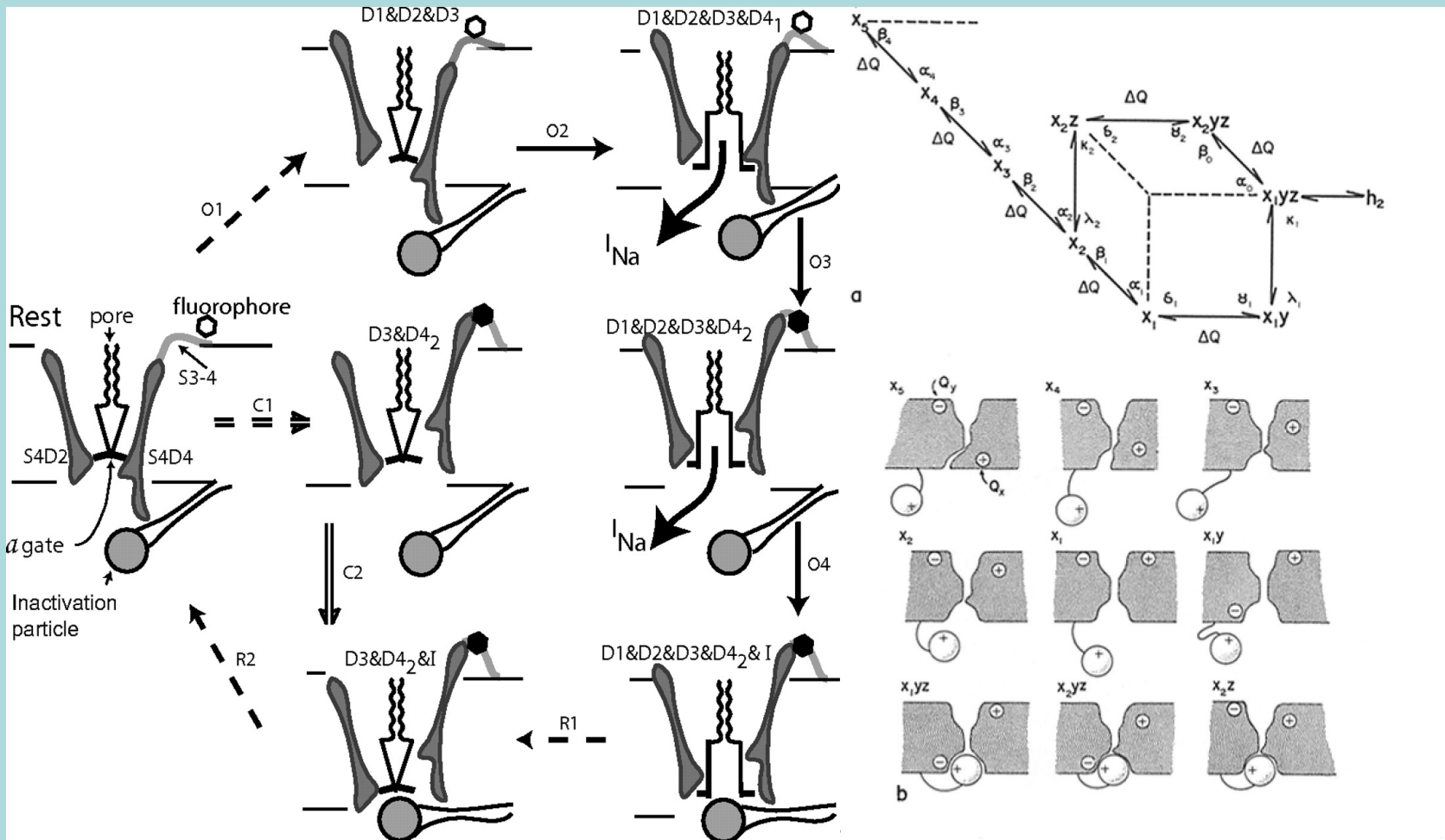
likely to be

**Controversial**

but

**Quantitative and Testable**

# Channel Activation and Inactivation 'Ball and Chain' Model



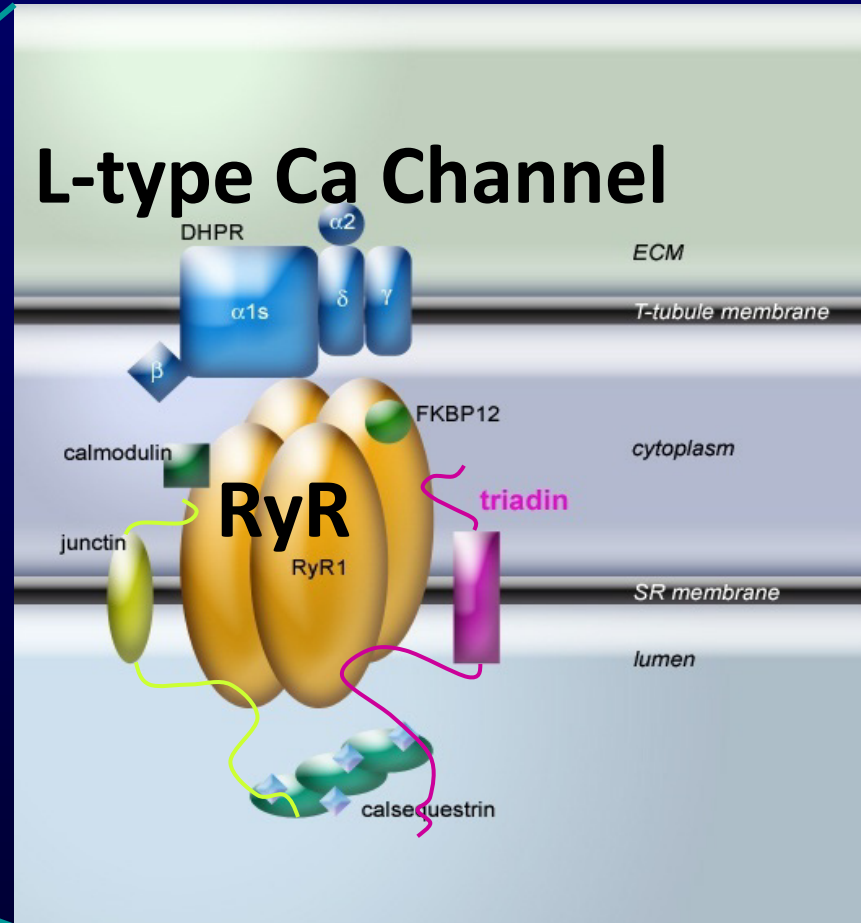
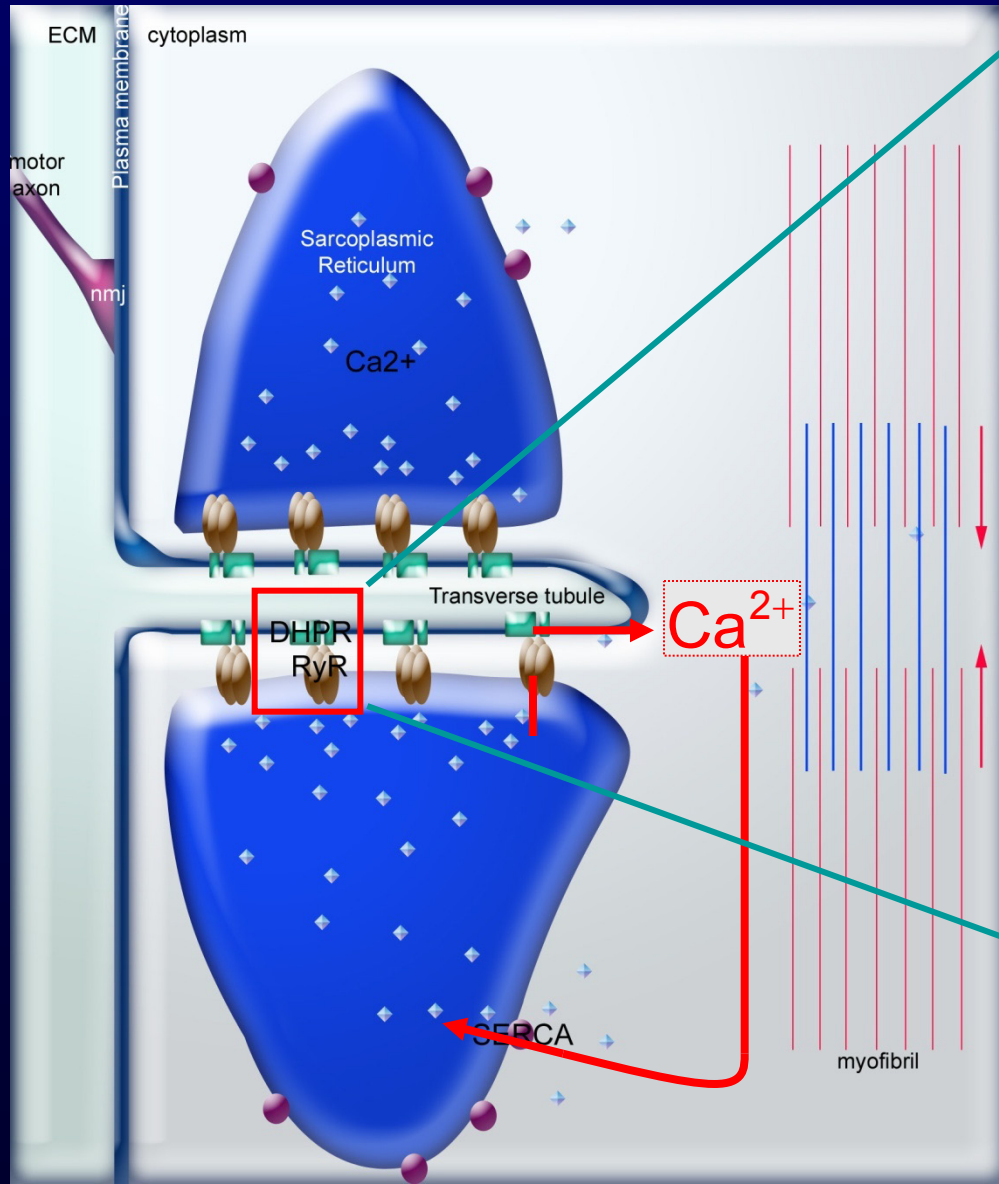
Armstrong PNAS 2006 103:17991

Armstrong & Bezanilla J Gen Physiol 1977 70:567

Existing Models are Structural and Mechanical  
with no quantitative results

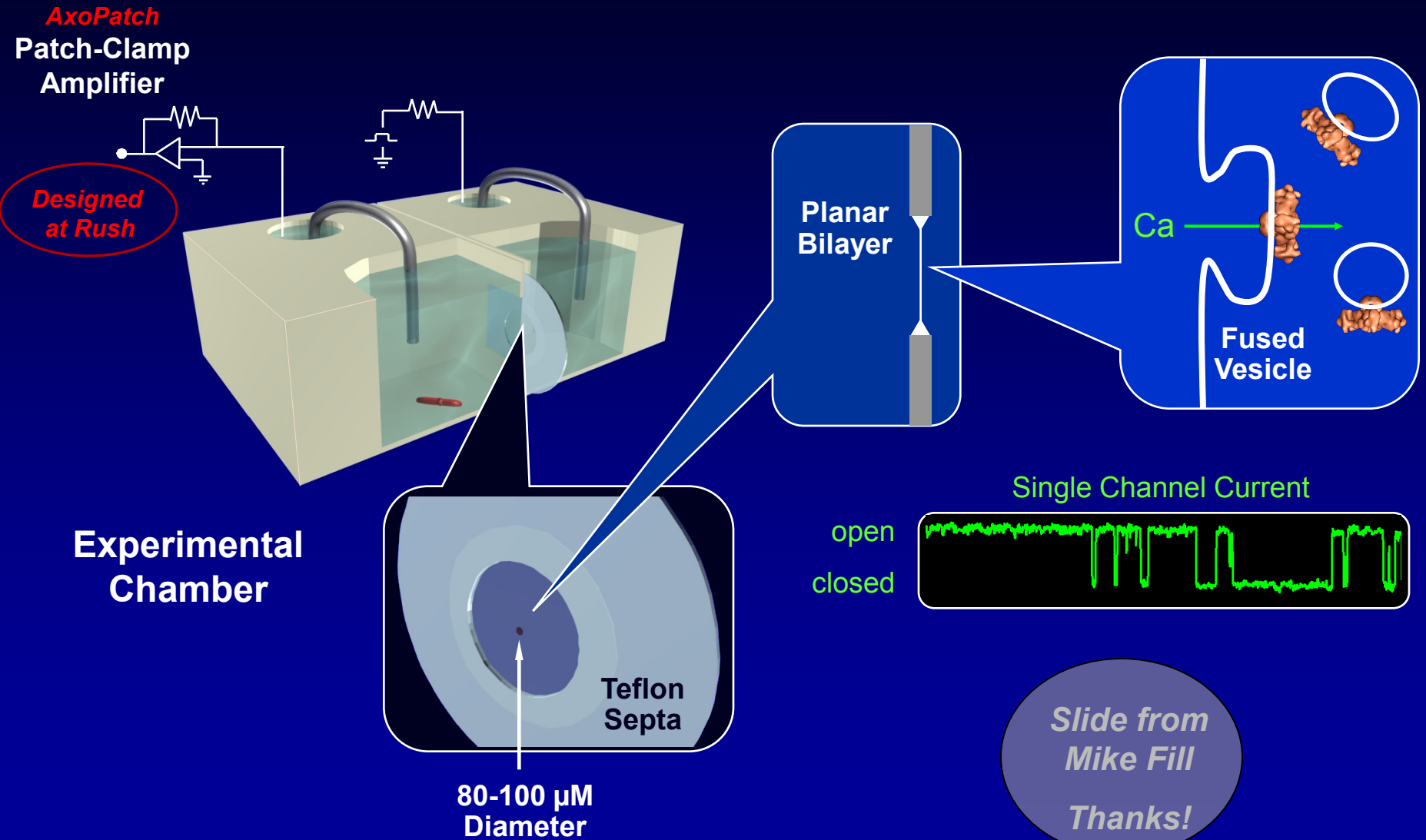
Channels are parts of Machines,  
e.g., Excitation-Contraction Coupling

L type Ca Channel    RyR ryanodine receptor



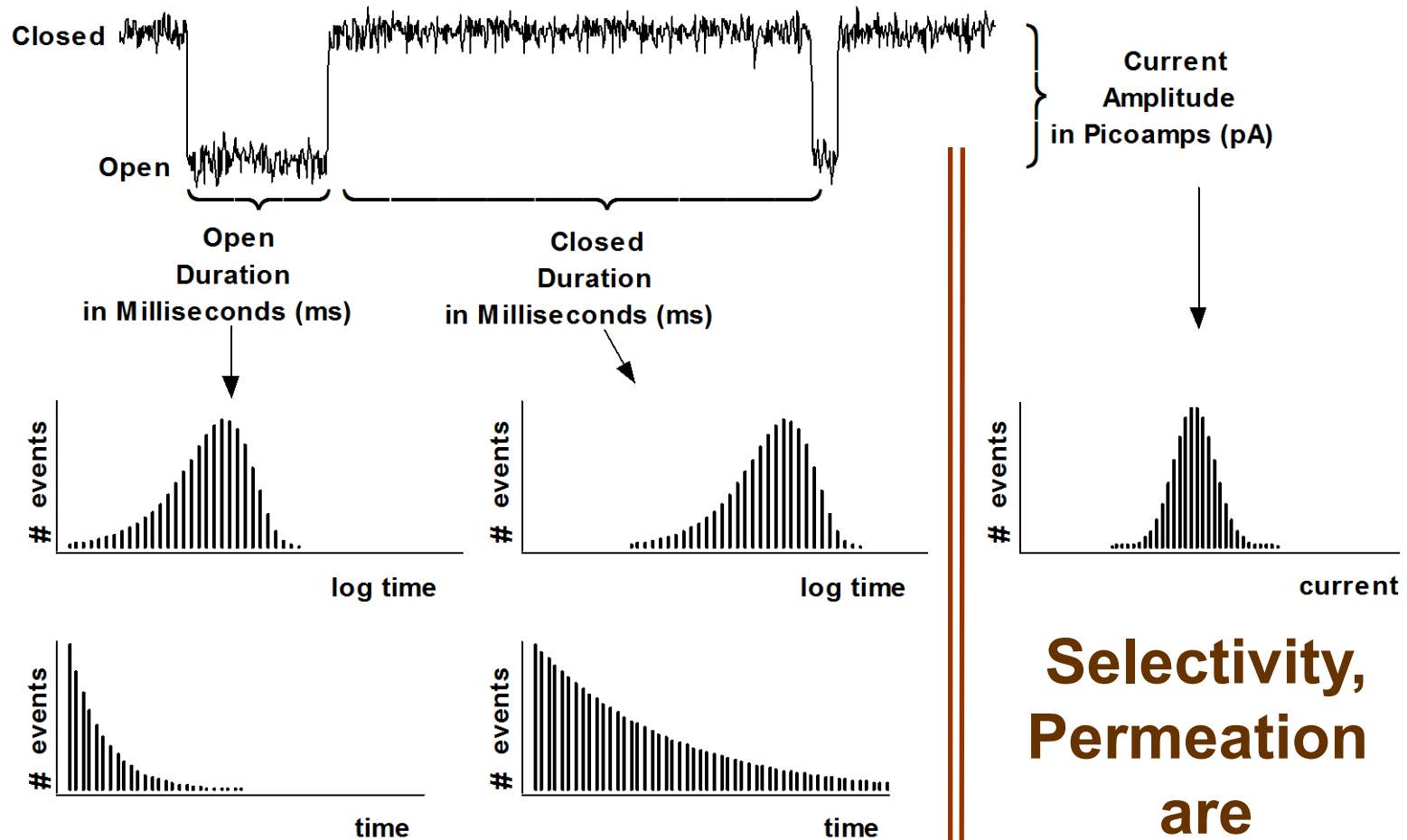
Thanks for the figure to  
László Csernoch, Debrecen, Hungary  
Isabelle Marty, Grenoble, France

# Function of SINGLE isolated RyR Channels in Artificial Planar Lipid Bilayers



*Slide from  
Mike Fill  
Thanks!*

# Gating and Permeation



**Gating is Time Behavior**

**Selectivity,  
Permeation  
are  
Amplitude**



*For Modelers and Mathematicians: This is reverse engineering!*

## Central Problem

**How does the channel control Selectivity?**

## Inverse Problem for Selectivity

Badly posed, many answers are possible,  
simultaneously over and under determined  
with noise and systematic error

**Core Math Problem has actually been solved**  
using methods for the  
**Inverse Problem of a Blast Furnace**

*Burger, Eisenberg and Engl (2007) SIAM J Applied Math 67: 960-989*

# *Channels are Selective*

**Different Types of Channels**

**use**

**Different Types of Ions**

**for**

**Different Information**

**New Interpretations**

likely to be

**Controversial**

but

**Quantitative and Testable**

# Time Dependence

is

## Important

**Many diseases produced by inactivation**

**Energetics of Brain**

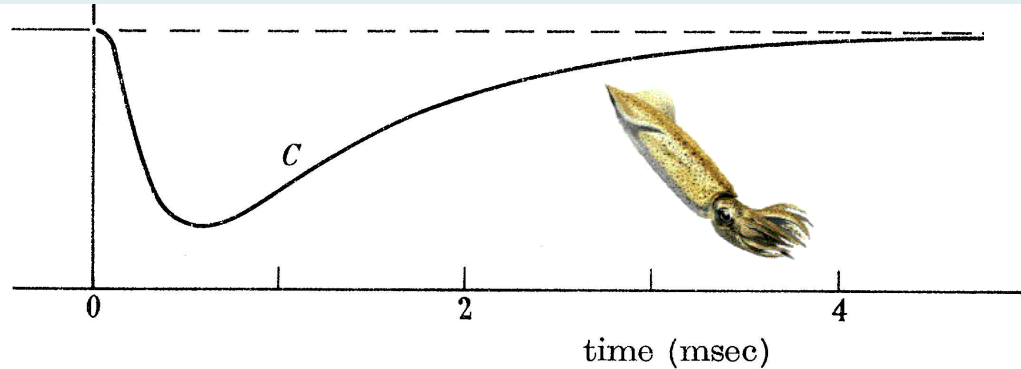
**determined by inactivation\***

**\*Energetics determined by time overlap of Na and K currents**

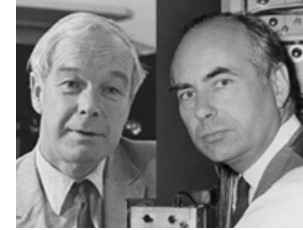
*Alle, Roth, and Geiger. Science (2009) 325:1405-8.*

# Time Dependent Sodium Conductance

## Inactivation in Squid Axon (nerve fiber)



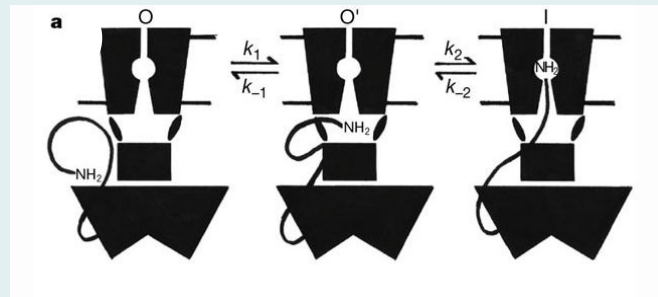
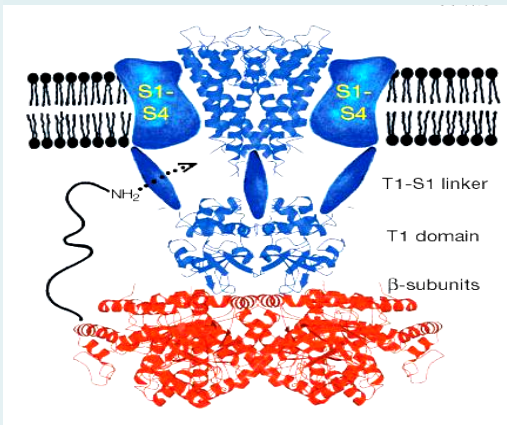
Hodgkin Huxley



J. Physiol (1952)  
116:497

FIGURE 9. Separation of current into components carried by Na and K, from Hodgkin & Huxley (1952a, figure 5). A depolarization of 56 mV was applied at  $t = 0$ ; the temperature was  $8.5^{\circ}\text{C}$ . Outward current is shown upwards.

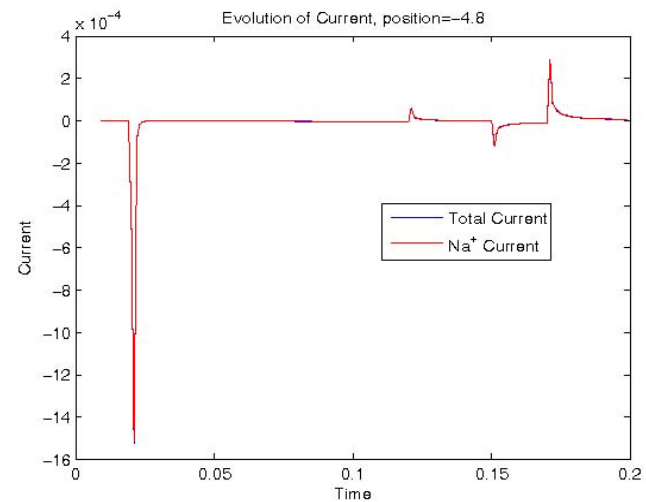
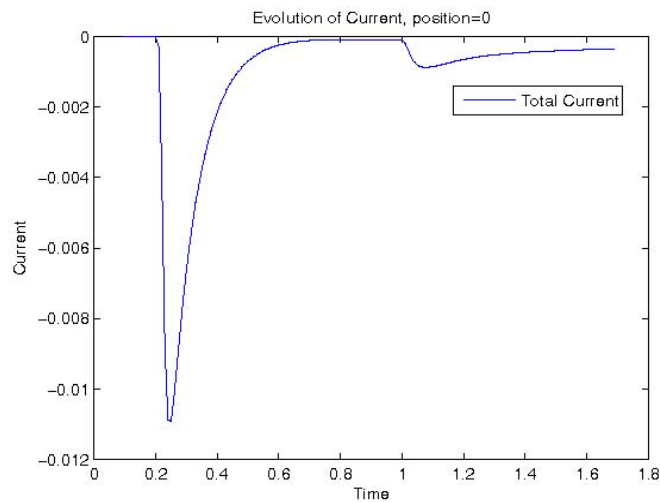
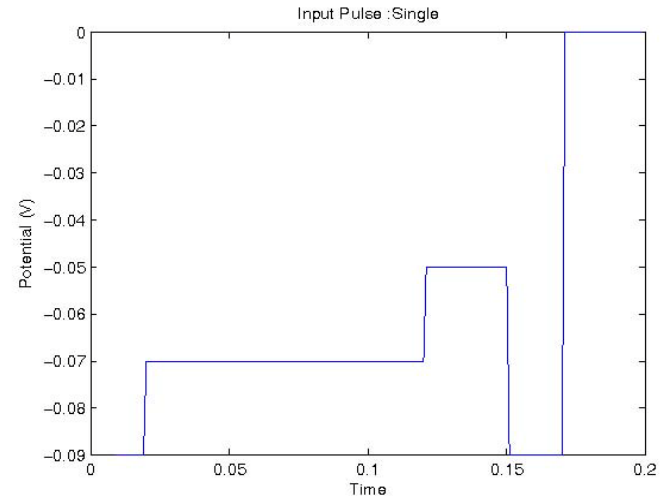
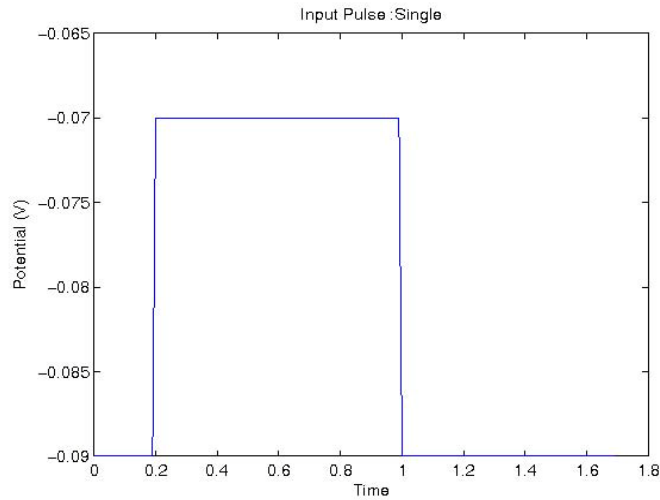
### Conventional Explanation: Elaborate Structural Change



$$\begin{aligned}
 C_1 &\xrightleftharpoons[\beta]{4\alpha} C_2 \xrightleftharpoons[2\beta x_1]{3\alpha x_1} C_3 \xrightleftharpoons[3\beta x_1^2]{2\alpha x_1^2} C_4 \xrightleftharpoons[4\beta x_1^3]{\alpha x_1^3} C_5 \\
 &\delta \parallel \gamma \\
 C_6 &\xrightleftharpoons[2\delta x_1 y_1]{3\alpha x_1 y_1} C_7 \xrightleftharpoons[2\beta x_1 y_1]{2\alpha x_1 y_1} C_8 \xrightleftharpoons[3\beta x_1 y_1^2]{\alpha x_1 y_1^2} C_9 \\
 &2\delta x_1 y_1 \parallel \gamma x_1 y_1 \\
 C_{10} &\xrightleftharpoons[2\beta x_1 y_1^2]{2\alpha x_1 y_1^2} C_{11} \xrightleftharpoons[2\beta x_1 y_1^3]{\alpha x_1 y_1^3} C_{12} \\
 &3\delta x_1 y_1^2 \parallel \gamma x_1 y_1^2 \\
 C_{13} &\xrightleftharpoons[4\delta x_1 y_1^3]{\alpha x_1 y_1^3} C_{14} \\
 &\parallel \gamma x_1 y_1^3 \\
 &O \\
 &\parallel \\
 &C_1
 \end{aligned}$$

# Sodium Conductance and Inactivation in Fixed Structure

## Variational Computation



# Multiscale Issues are the key if we want to actually build channels that work

Computational Scale	Biological Scale	Ratio
<u>Time</u> $10^{-15}$ sec	$10^{-4}$ sec <i>Action Potential</i>	$10^{11}$
<u>Space</u> $10^{-11}$ m	$10^{-5}$ m <i>Side Chains of Proteins</i>	$10^6$
<u>Spatial Resolution</u>	<i>Three Dimensional</i> $(10^6)^3$	$10^{18}$
<u>Solute Concentration</u>	$10^{-11}$ to 20 Molar	$10^{12}$