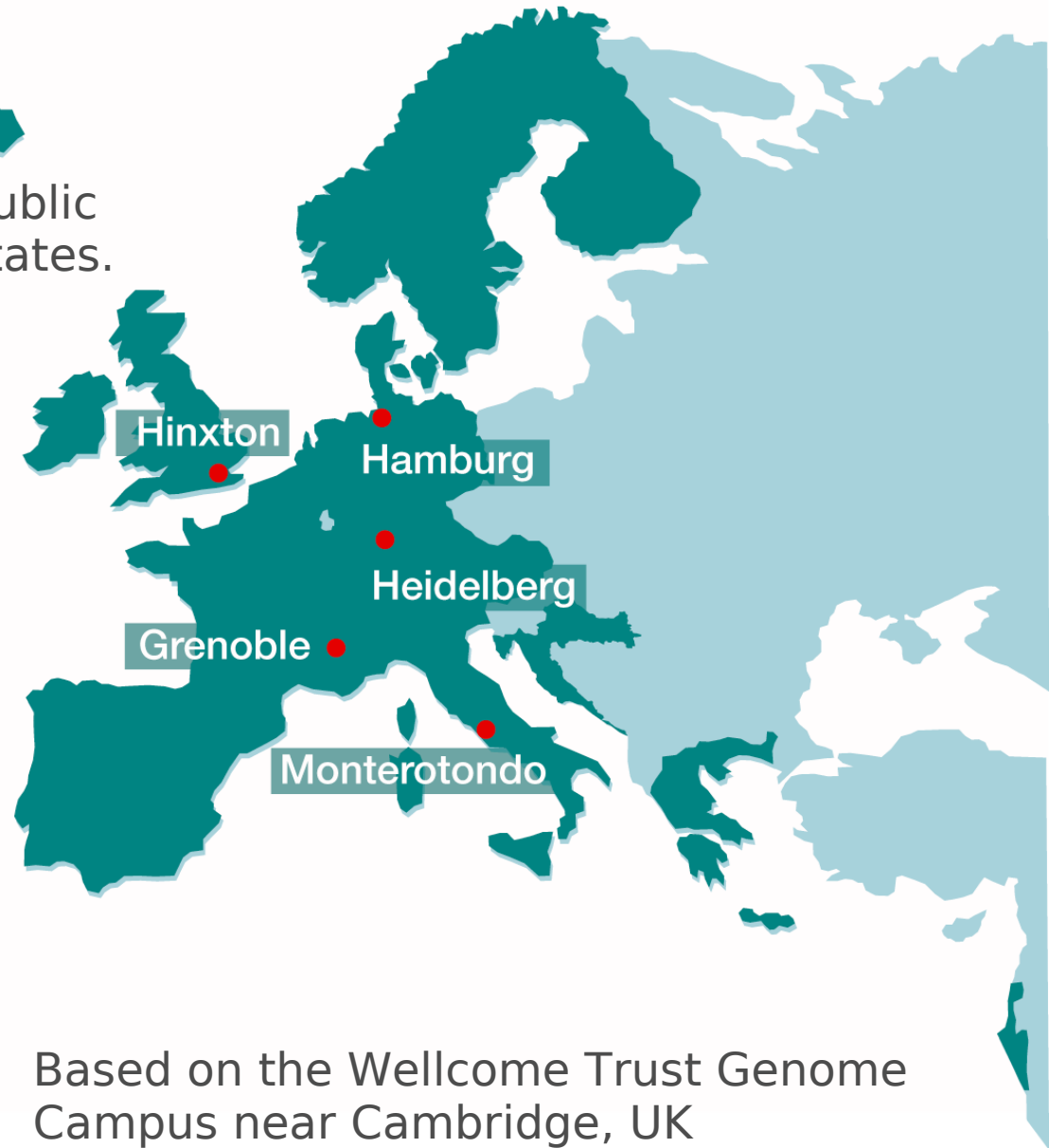


Modélisation du comportement des protéines allostériques impliquées dans la plasticité synaptique

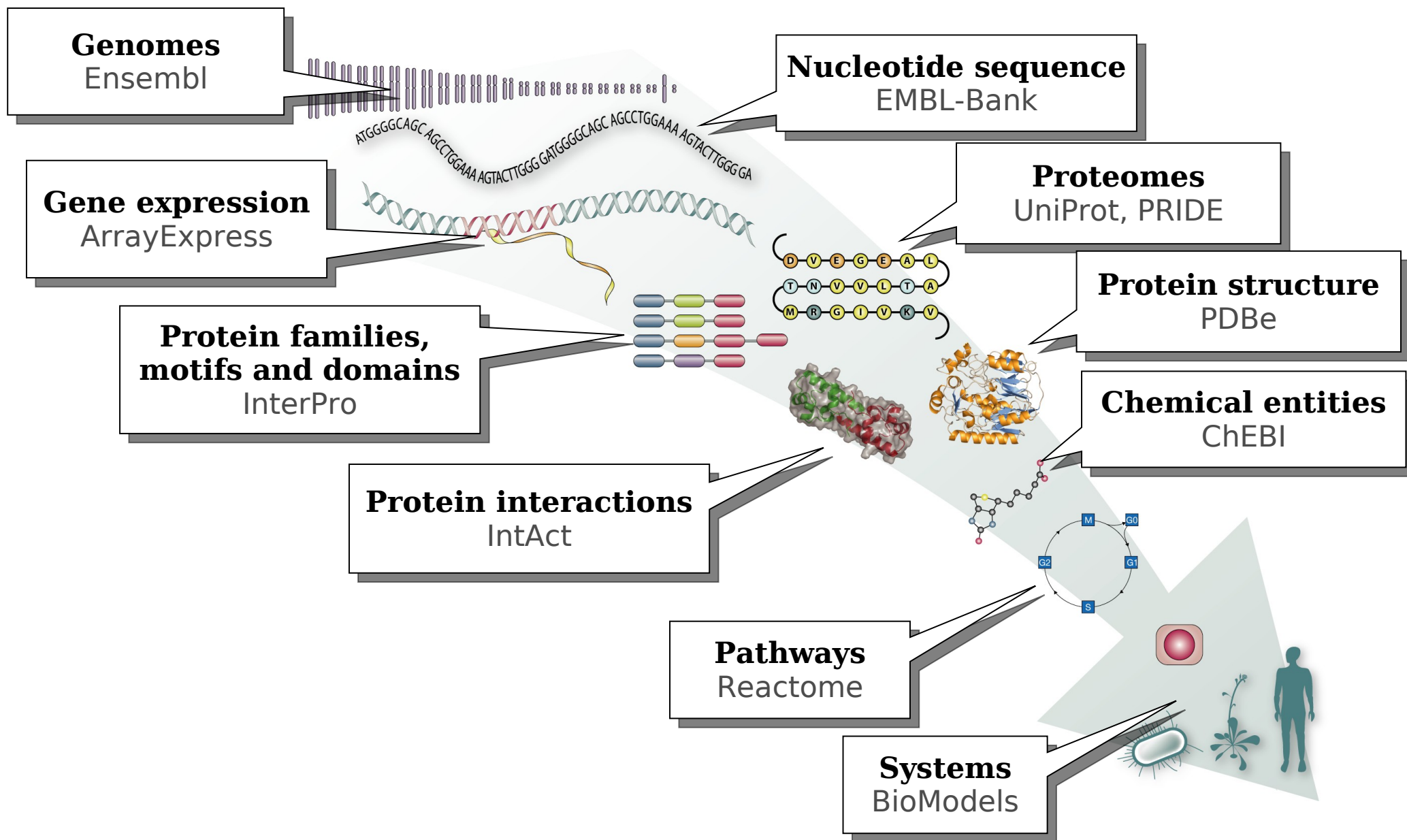
Nicolas Le Novère, EMBL-EBI

EBI is part of EMBL

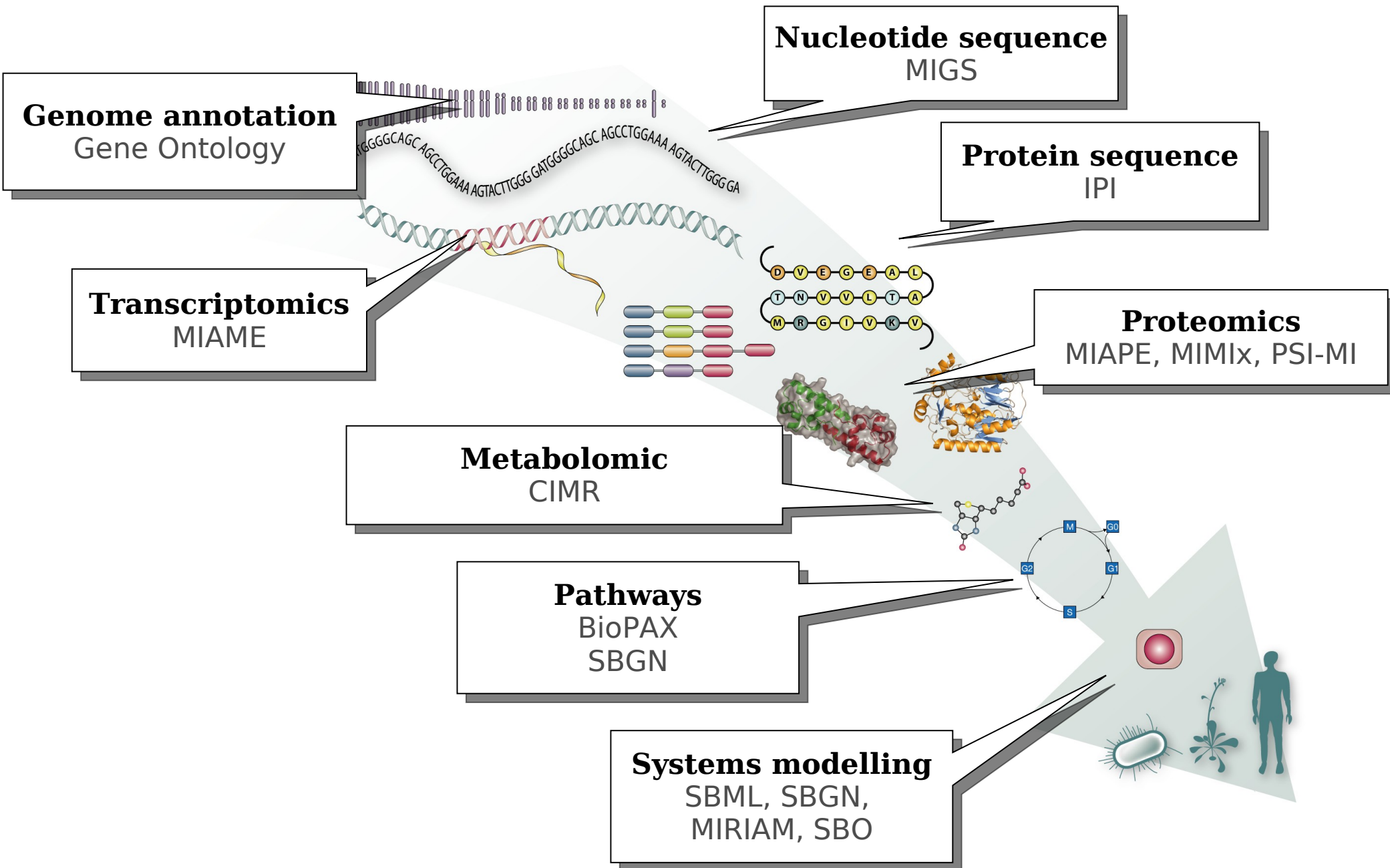
- Non-profit organization
- Part of the European Molecular Biology Laboratory (EMBL), a basic research institute funded by public research monies from 19 member states.



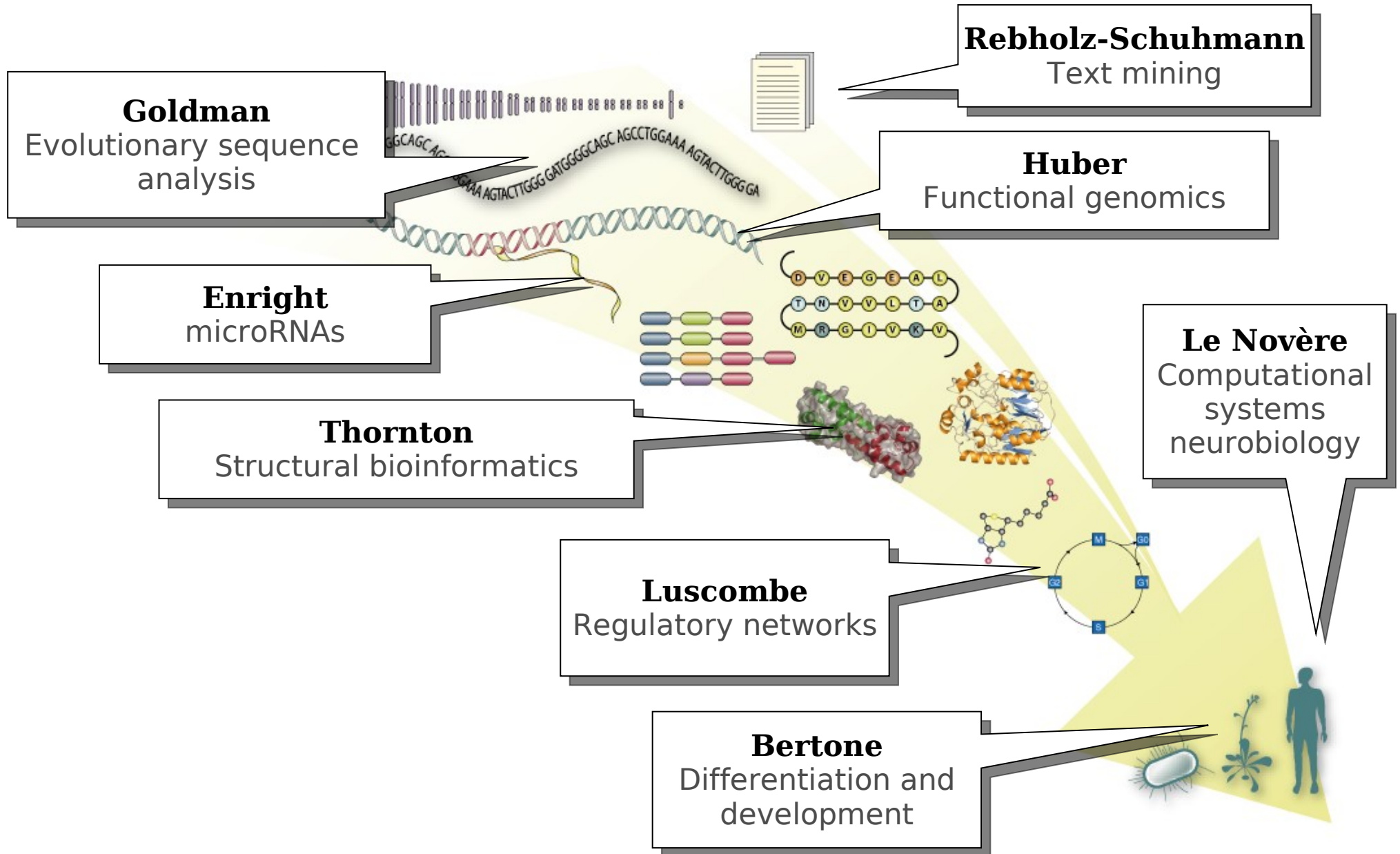
EBI Data resources: molecules to systems

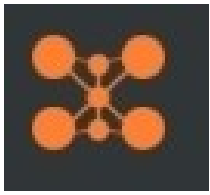


Standards development

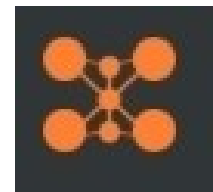


Research groups



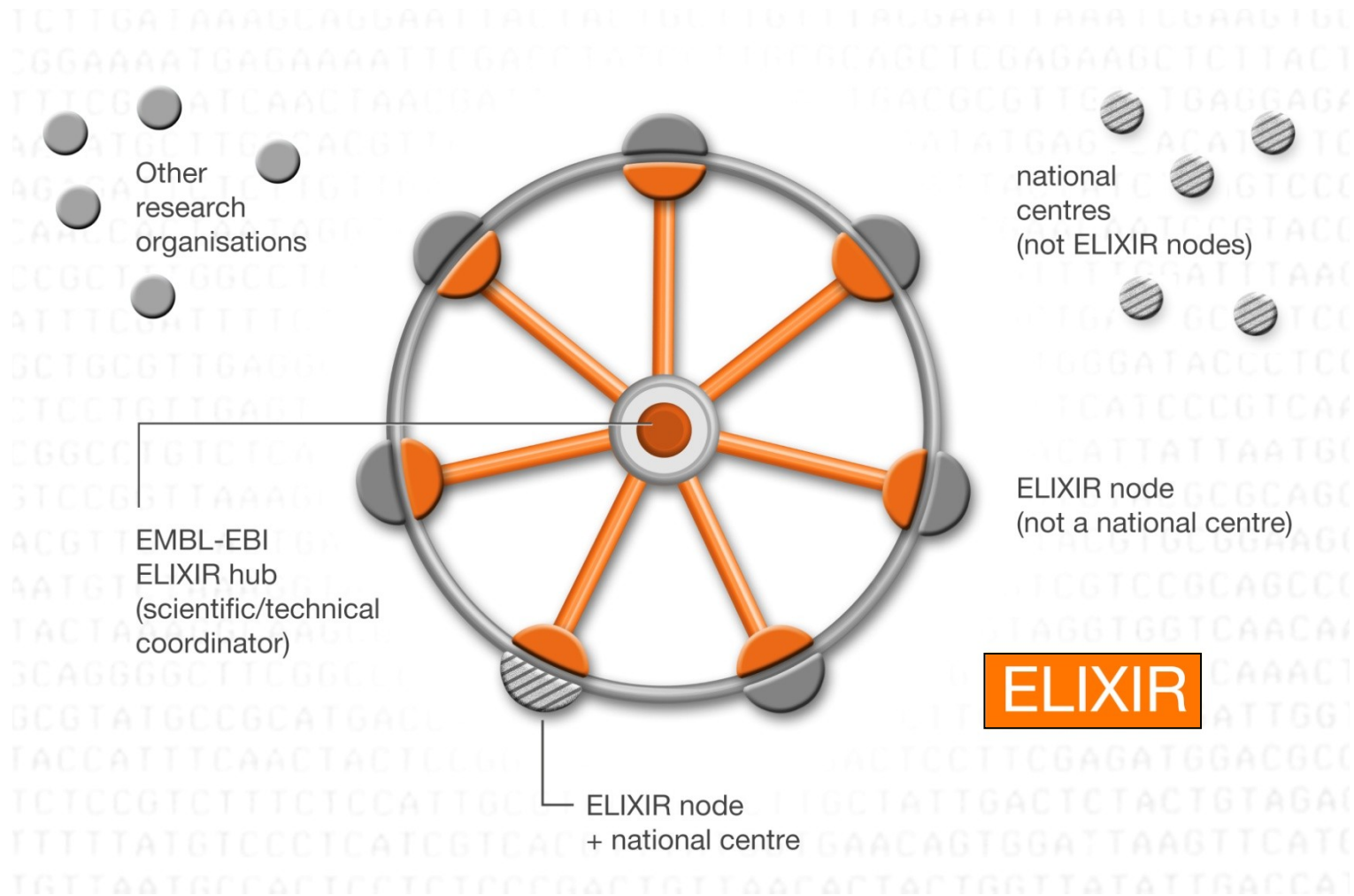
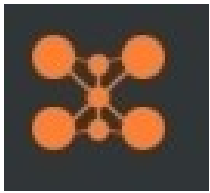


- An EU Framework 7 Preparatory Phase Project
- Coordinated by Prof Janet Thornton, Director EMBL-EBI
- To construct a plan for the operation of a **sustainable** infrastructure for biological information in Europe
- €4.5 million grant awarded May 2007, three year term
- 32 member consortium engaging many of Europe's main bioinformatics funding agencies and research institutes
- Deliverables are memoranda of understanding to fund the implementation phase which could cost €500 million
- Interested parties should register as stake-holders via the ELIXIR Website: www.elixir-europe.org



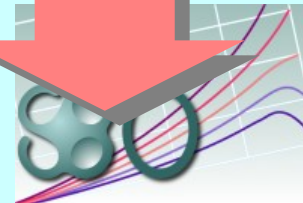
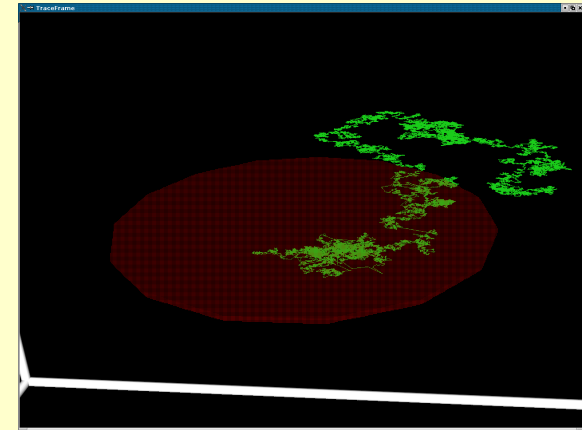
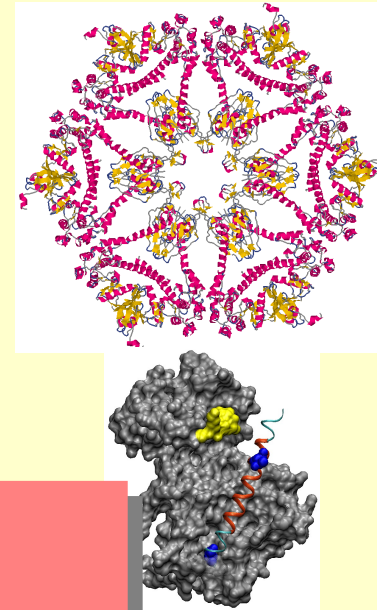
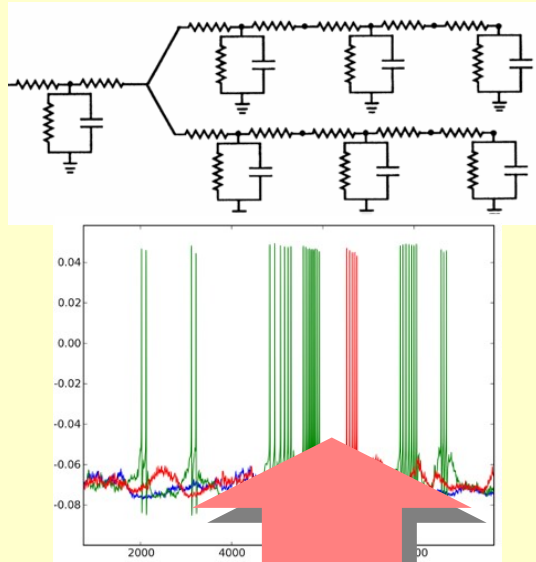
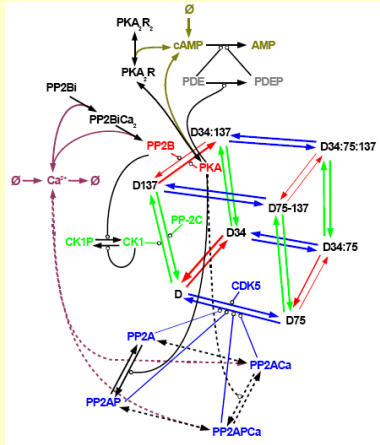
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including
Bordeaux 2

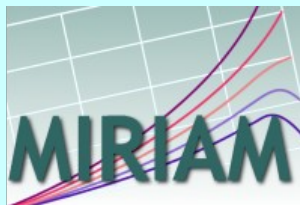


compneur group themes and projects

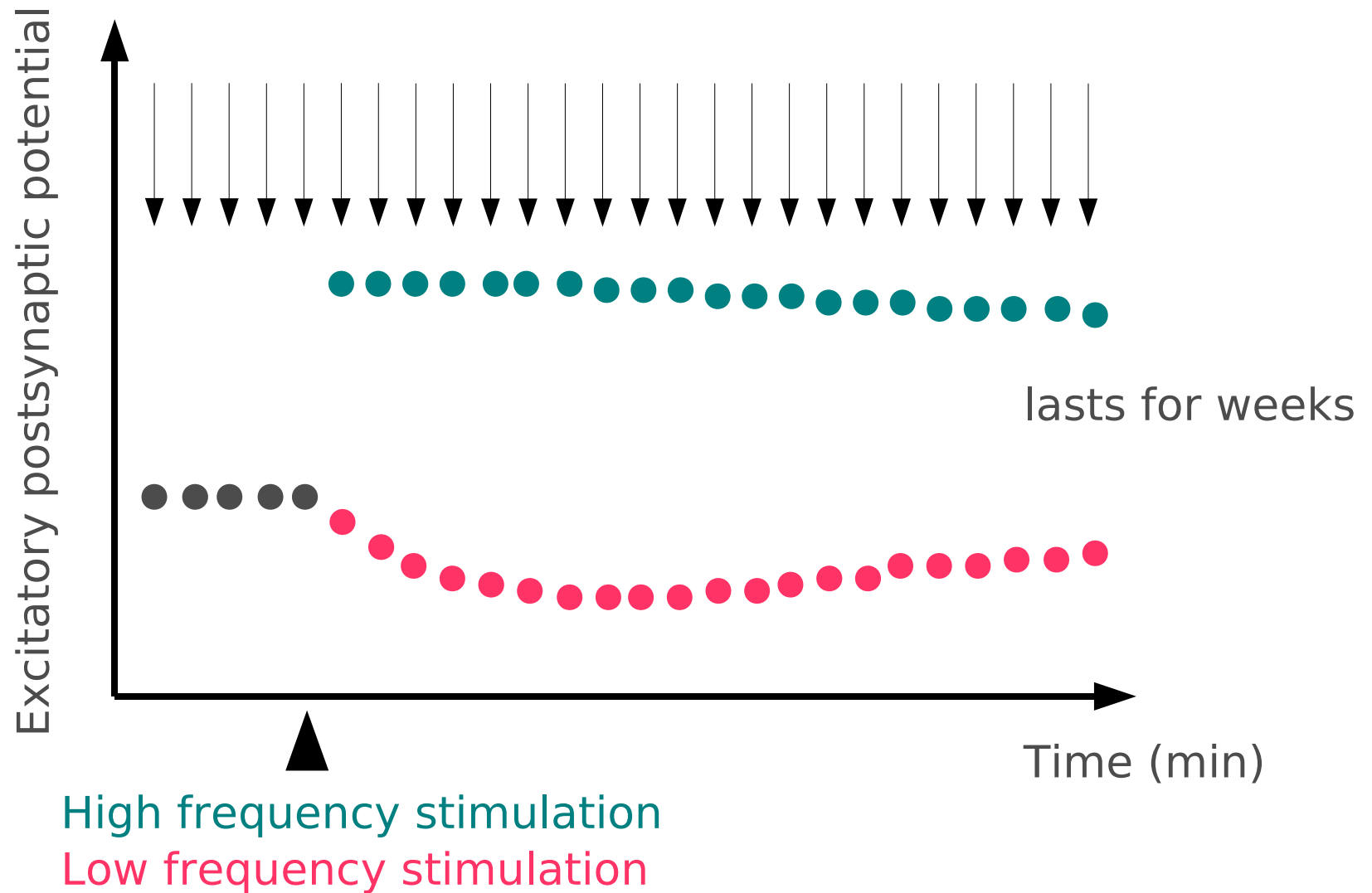
Computational Neurobiology



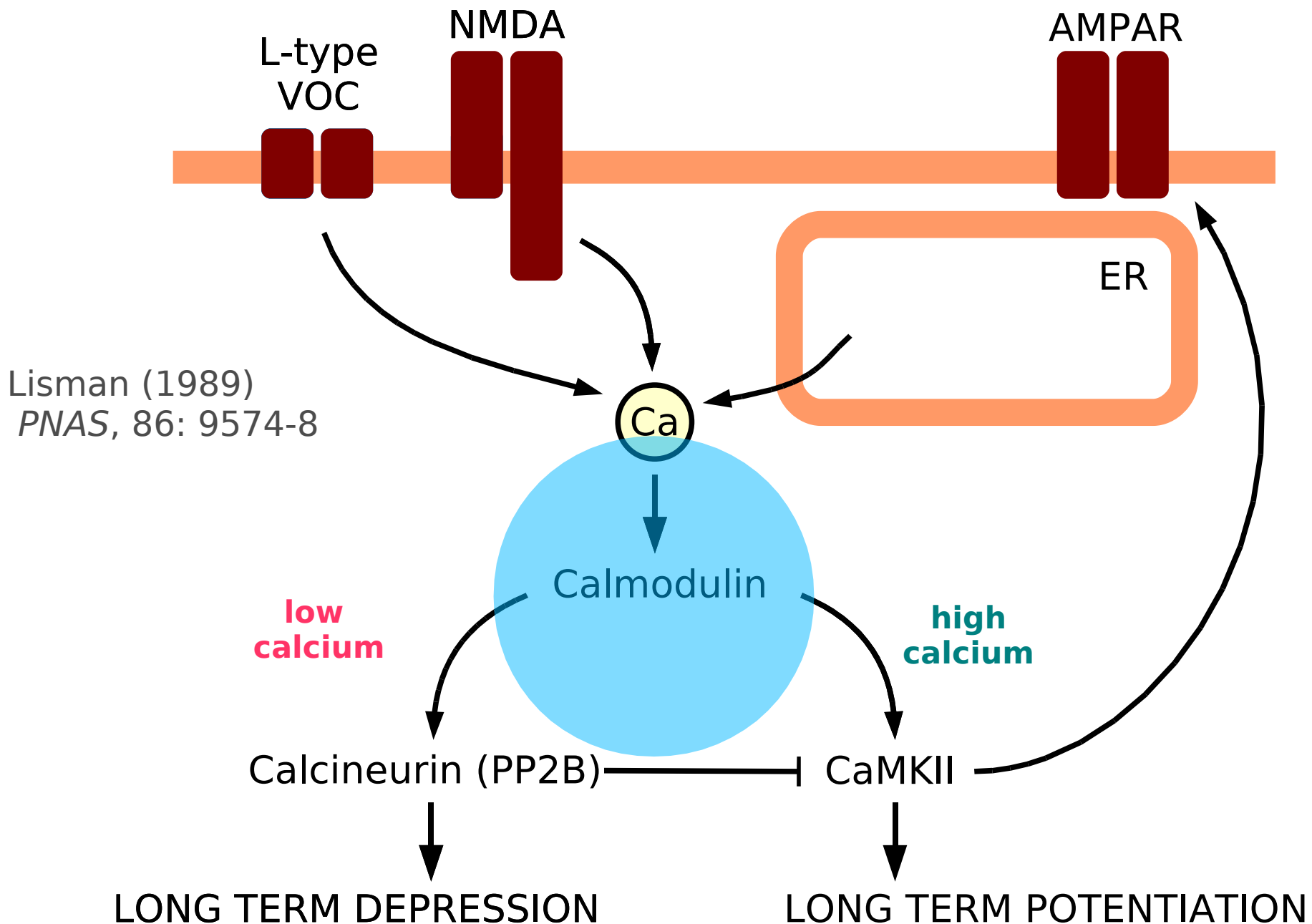
Computational Systems Biology



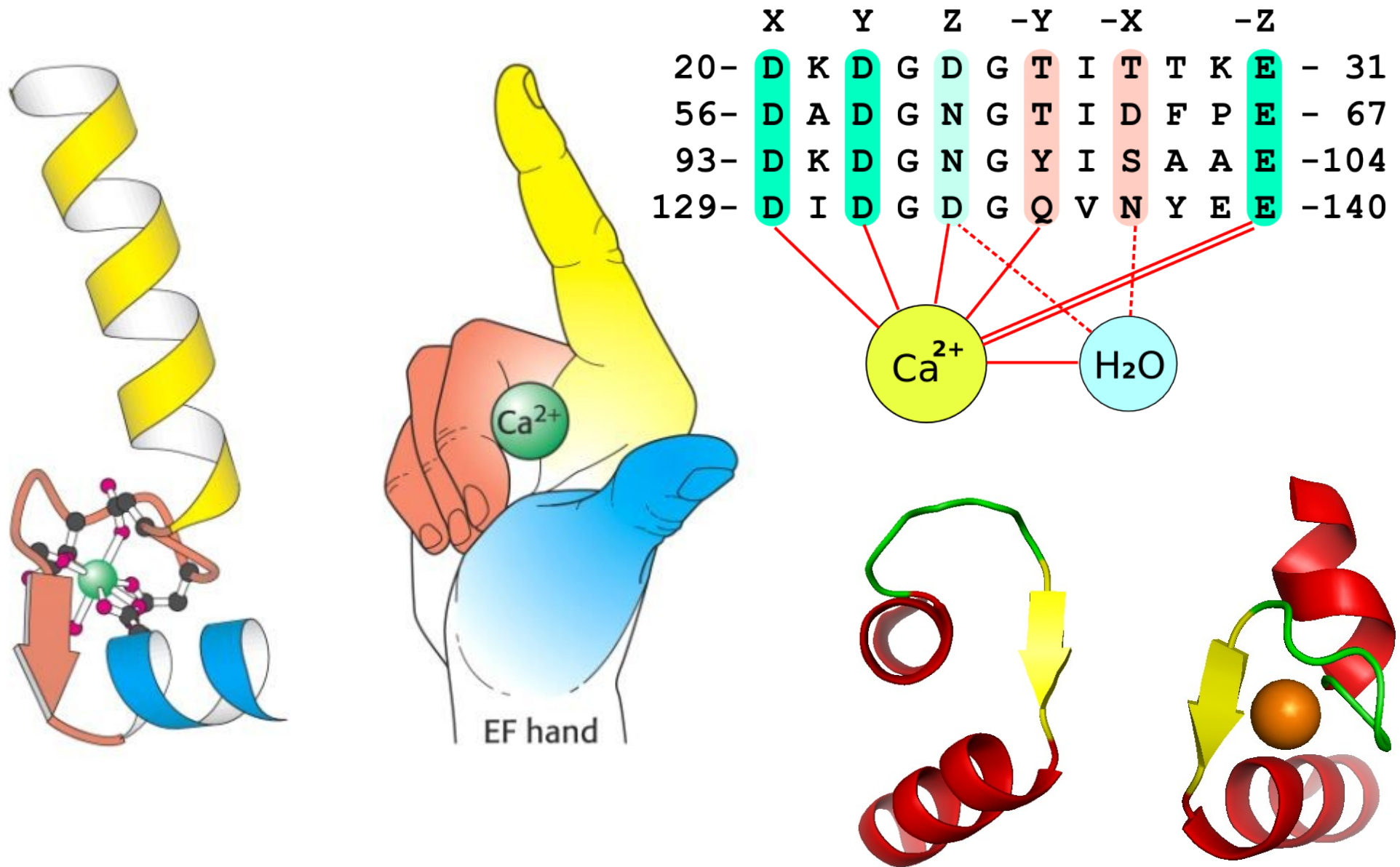
Bidirectional synaptic plasticity



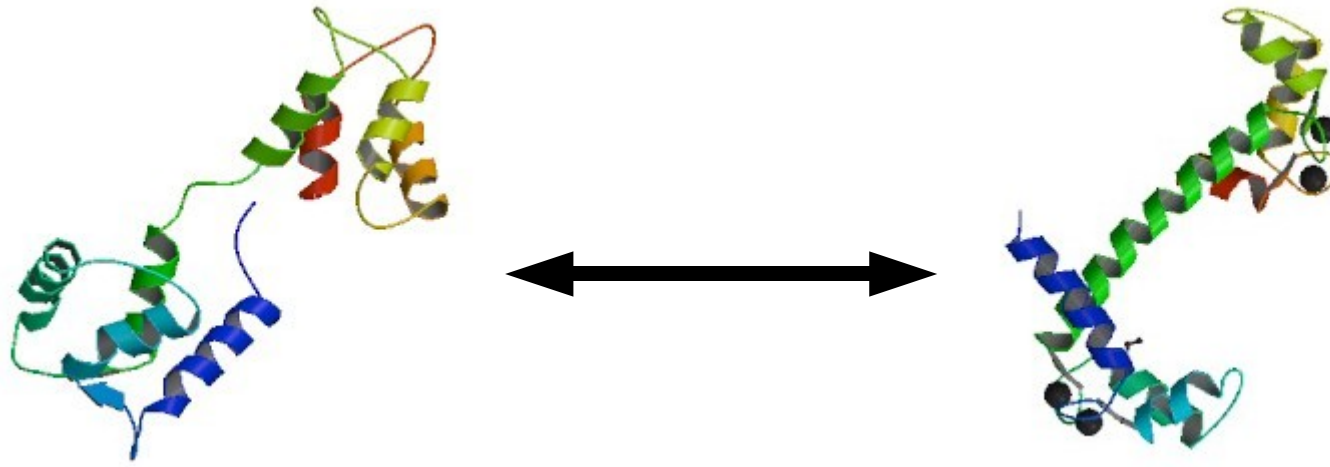
Calmodulin, the memory switch



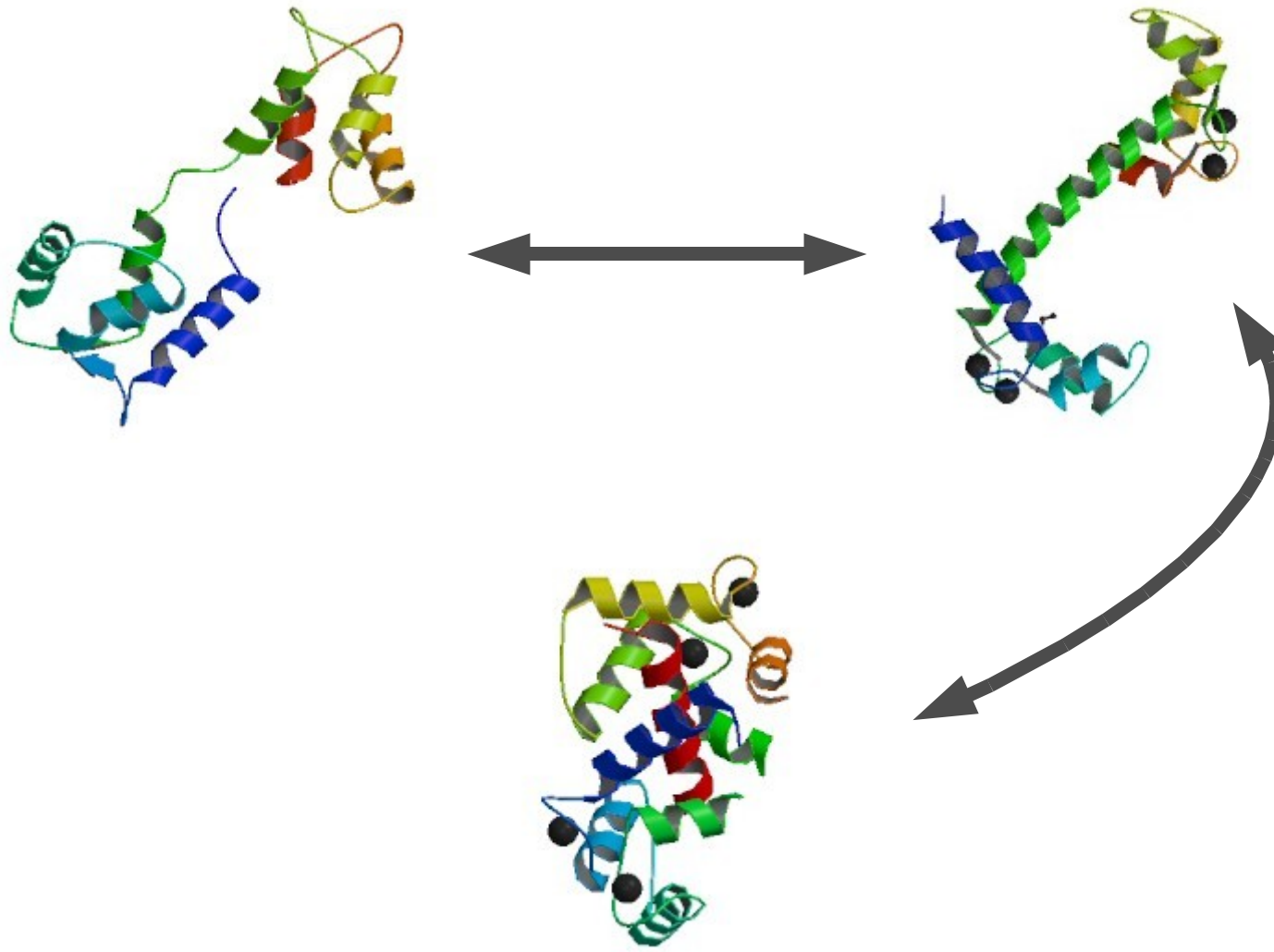
Structure of Calmodulin



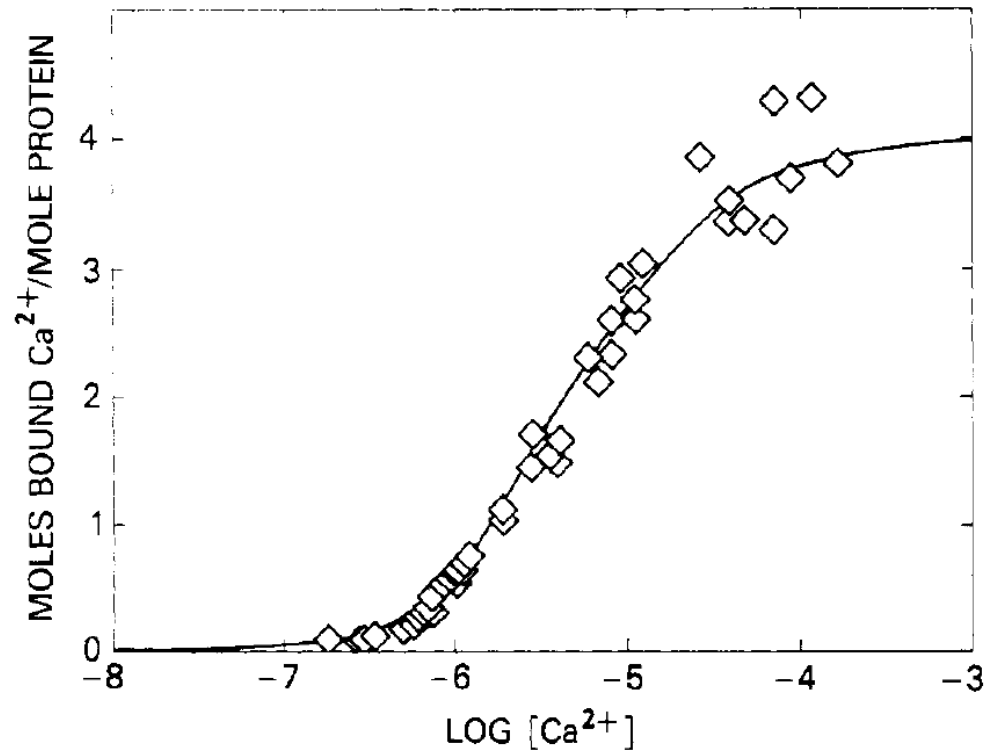
State transitions of calmodulin



State transitions of calmodulin

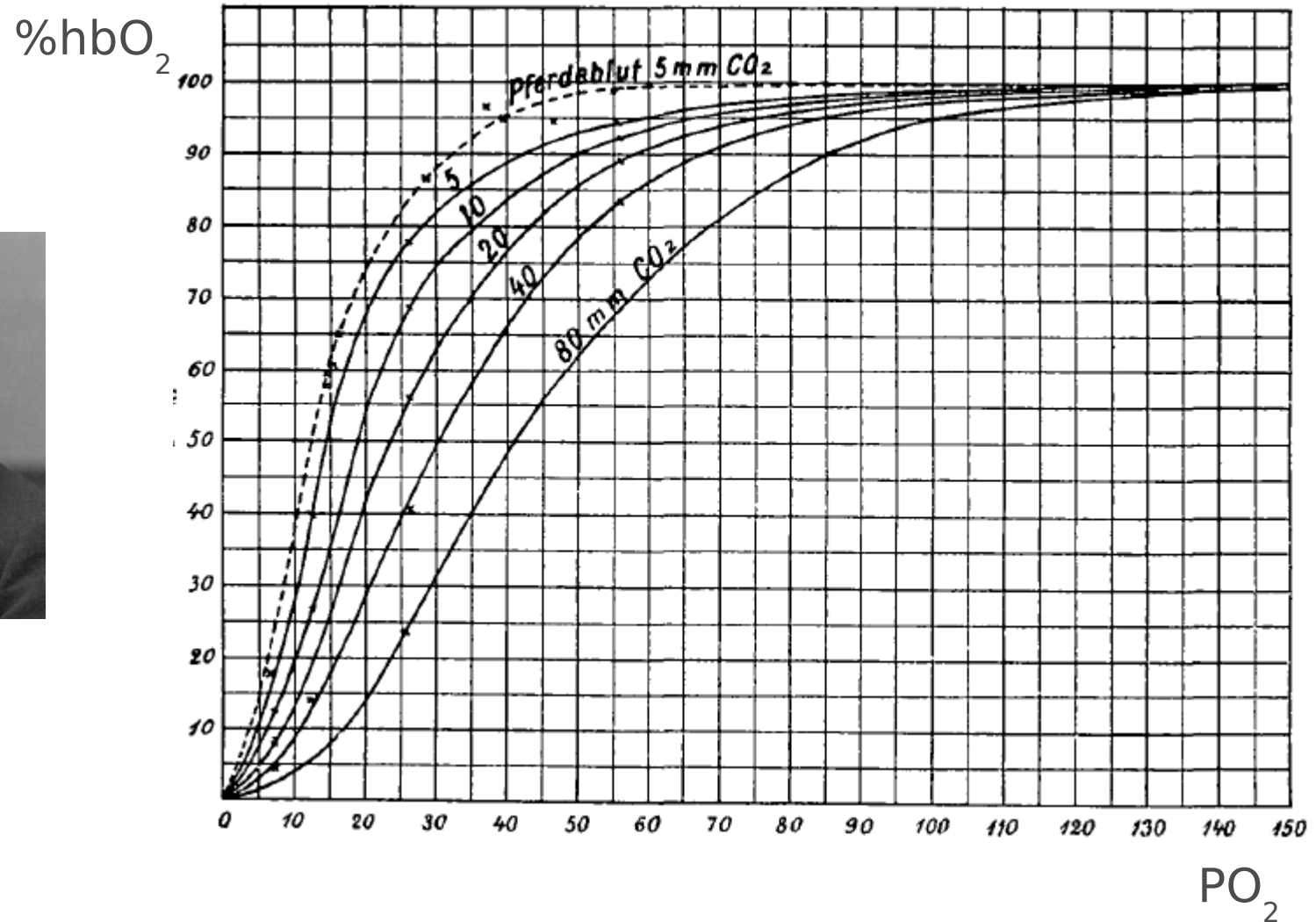


Calmodulin is ultra-sensitive



Crouch and Klee (1980)
Biochemistry, 19: 3692-3698

Origins of cooperativity: Bohr



Bohr C (1903) Theoretische behandlung der quantitativen verhältnisse bei der sauerstoff aufnahme des hämoglobins *Zentralbl Physiol* 17: 682

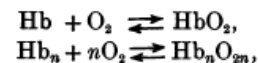
The possible effects of the aggregation of the molecules of hæmoglobin on its dissociation curves. By A. V. HILL.

In a previous communication Barcroft and I gave evidence which seemed to us to prove conclusively that dialysed hæmoglobin consists simply of molecules containing each one atom of iron. The molecular weight is therefore $Hb = 16,660$. These experiments have not been published yet, but I shall assume the results.

Other observers (Reid, Roaf, Hüfner and Gansser) working on different solutions have obtained divergent results. The method used by all of them was the direct estimation of the osmotic pressure, by means of a membrane permeable to salts, but not to hæmoglobin. The method involves a relatively large error, because the quantity measured is small. It is doubtful however whether this can explain the discordant results.

Our work led me to believe that the divergence between the results of different observers was due to an aggregation of the hæmoglobin molecules by the salts present in the solution, a consequent lowering of the number of molecules, and an increase in the average molecular weight as observed by the osmotic pressure method. To test this hypothesis I have applied it to several of the dissociation curves obtained by Barcroft and Camis with hæmoglobin in solutions of various salts, and with hæmoglobin prepared by Bohr's method.

The equation for the reaction would be



where Hb_n represents the aggregate of n molecules of Hb . I have supposed that in every solution there are many different sized aggregates, corresponding to many values of n .

If there were in the solution only Hb and Hb_2 the dissociation curve would be

$$y = \lambda \frac{K'x^2}{1 + K'x^2} + (100 - \lambda) \frac{Kx}{1 + Kx} \dots\dots\dots (A),$$

where $\lambda\%$ is as Hb_2 , $(100 - \lambda)\%$ as Hb , K' is the equilibrium constant of the reaction $Hb_2 + 2O_2 \rightleftharpoons Hb_2O_4$ and K that of $Hb + O_2 \rightleftharpoons HbO_2$: K has the value '125 (Barcroft and Roberts).

Origins of cooperativity: Hill

Hill (1910) *J Physiol* 40: iv-vii.

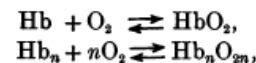
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$$y = 100 \frac{Kx^n}{1 + Kx^n} \dots\dots\dots (B)$$

would satisfy the observations.

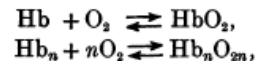
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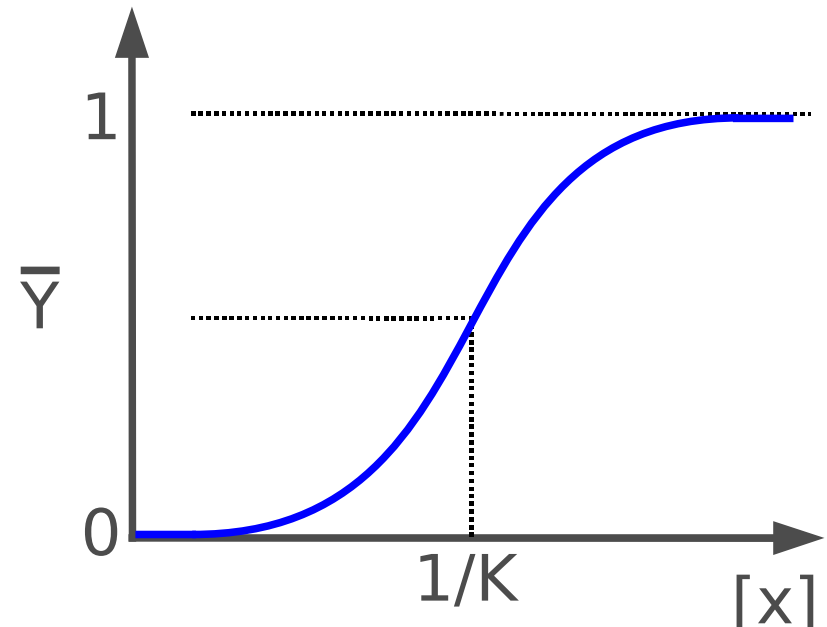
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$$\bar{Y} = \frac{K^n [X]^n}{1 + K^n [X]^n}$$

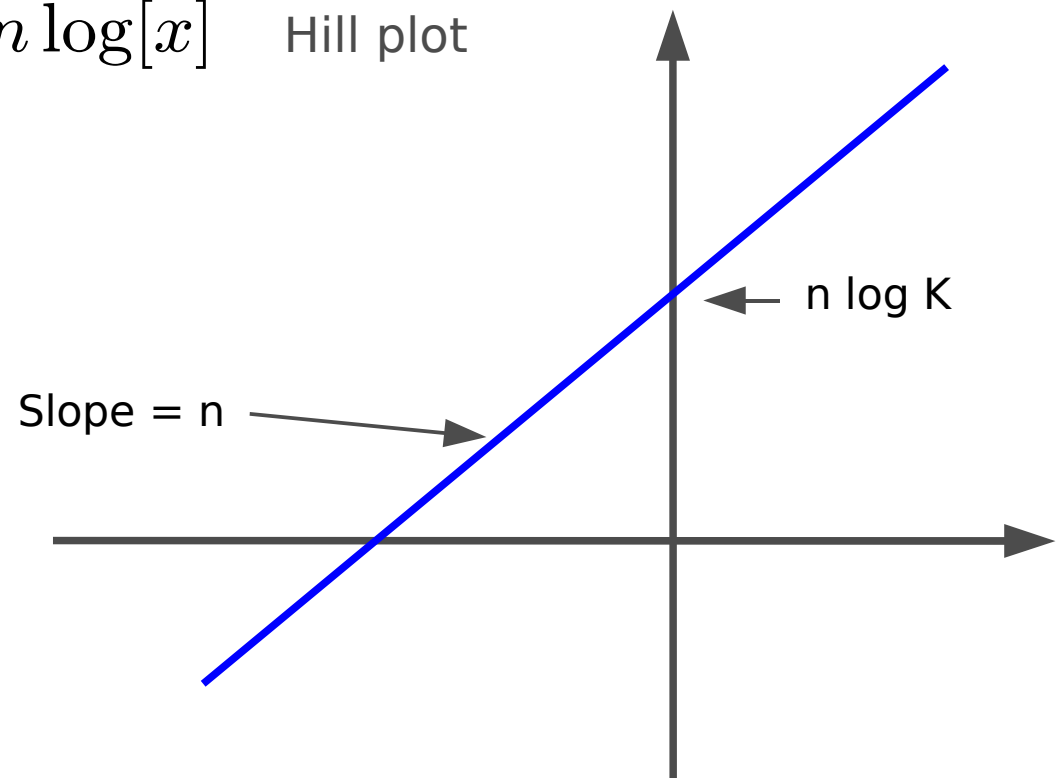
Hill equation

$$\log \frac{\bar{Y}}{1 - \bar{Y}} = n \log K + n \log [x]$$

Hill plot

Effect increases in function of
the signal to the power of n:
n>1, ultra-sensitive
n<1, infra-sensitive

BUT cooperativity of ligand,
not of binding sites: unique affinity



Origins of cooperativity: Adair-Klotz

THE HEMOGLOBIN SYSTEM.

VI. THE OXYGEN DISSOCIATION CURVE OF HEMOGLOBIN.*

By G. S. ADAIR.

With the Collaboration of A. V. BOCK and H. FIELD, JR.

(From the Medical Laboratories of the Massachusetts General Hospital,
Boston.)

(Received for publication, January 7, 1925.)

This work gives the oxygen dissociation curves of solutions previously investigated in regard to their acid-binding and base-

Adair (1925) *J Biol Chem* 63: 529

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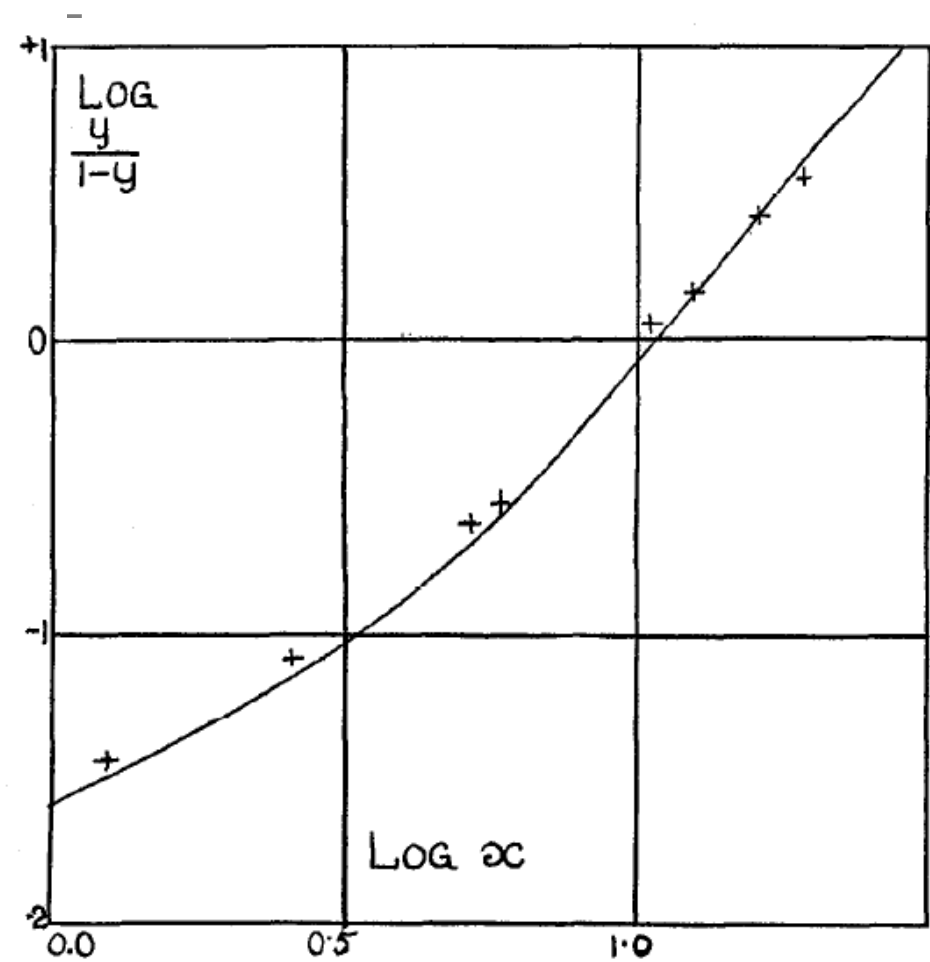


FIG. 2. Test of formula (6). Curve drawn from 6 experimental points from Table IV.

Origins of cooperativity: Adair-Klotz

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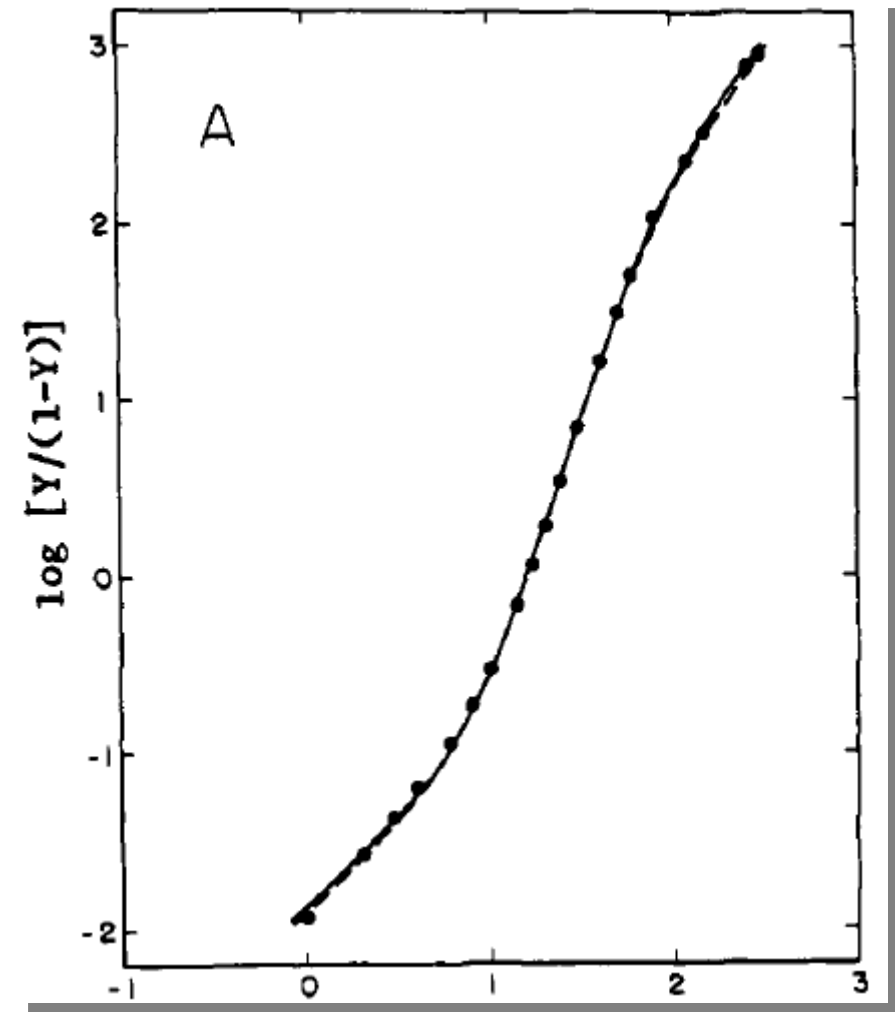
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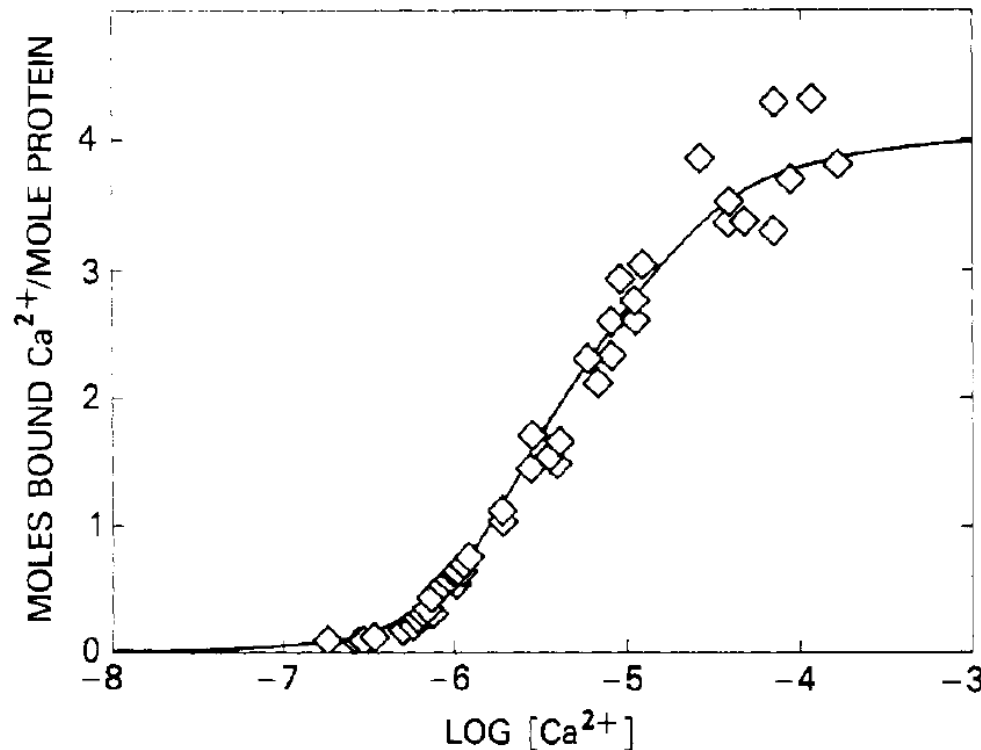


Imai (1973) *Biochemistry* 12: 798-808

Adair-Klotz model applied to Calmodulin

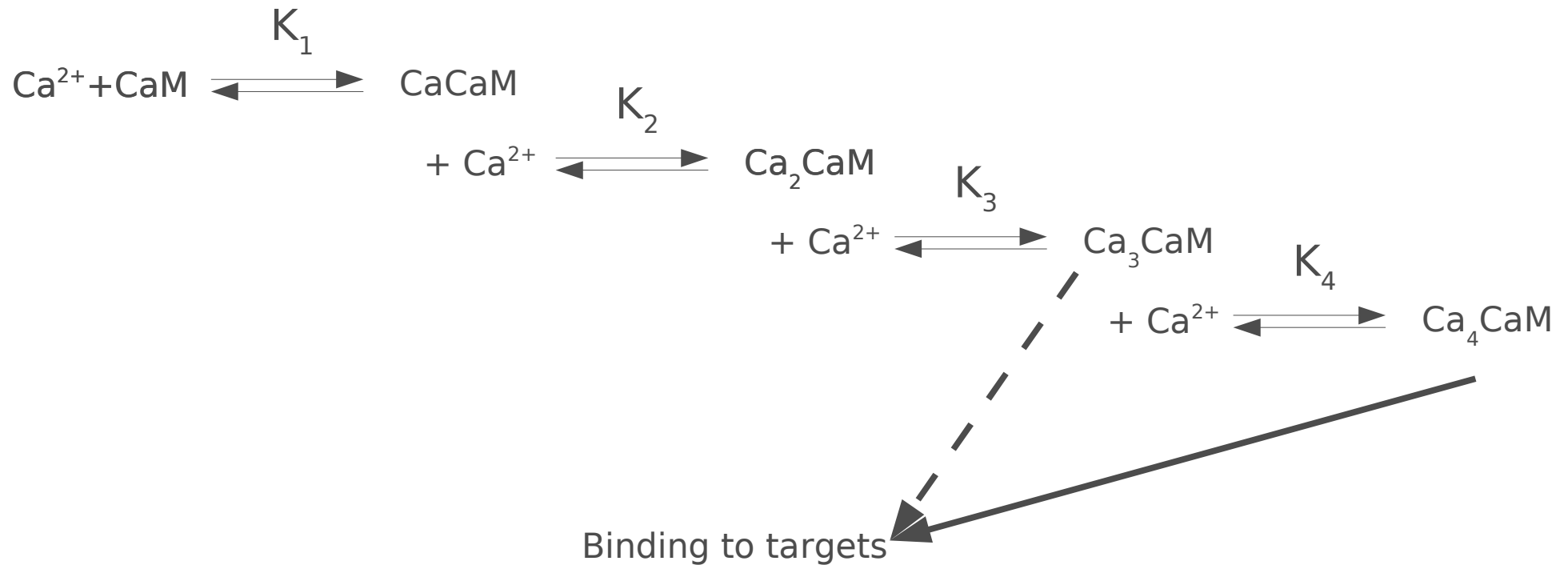
Klotz (1946) The Application of the Law of Mass Action to Binding by Proteins. Interactions with Calcium. *Arch Biochem*, 9:109–117.

$$\bar{Y} = \frac{1}{n} \frac{K_1[Ca] + 2K_1K_2[Ca]^2 + 3K_1K_2K_3[Ca]^3 + 4K_1K_2K_3K_4[Ca]^4}{1 + K_1[Ca] + K_1K_2[Ca]^2 + K_1K_2K_3[Ca]^3 + K_1K_2K_3K_4[Ca]^4}$$

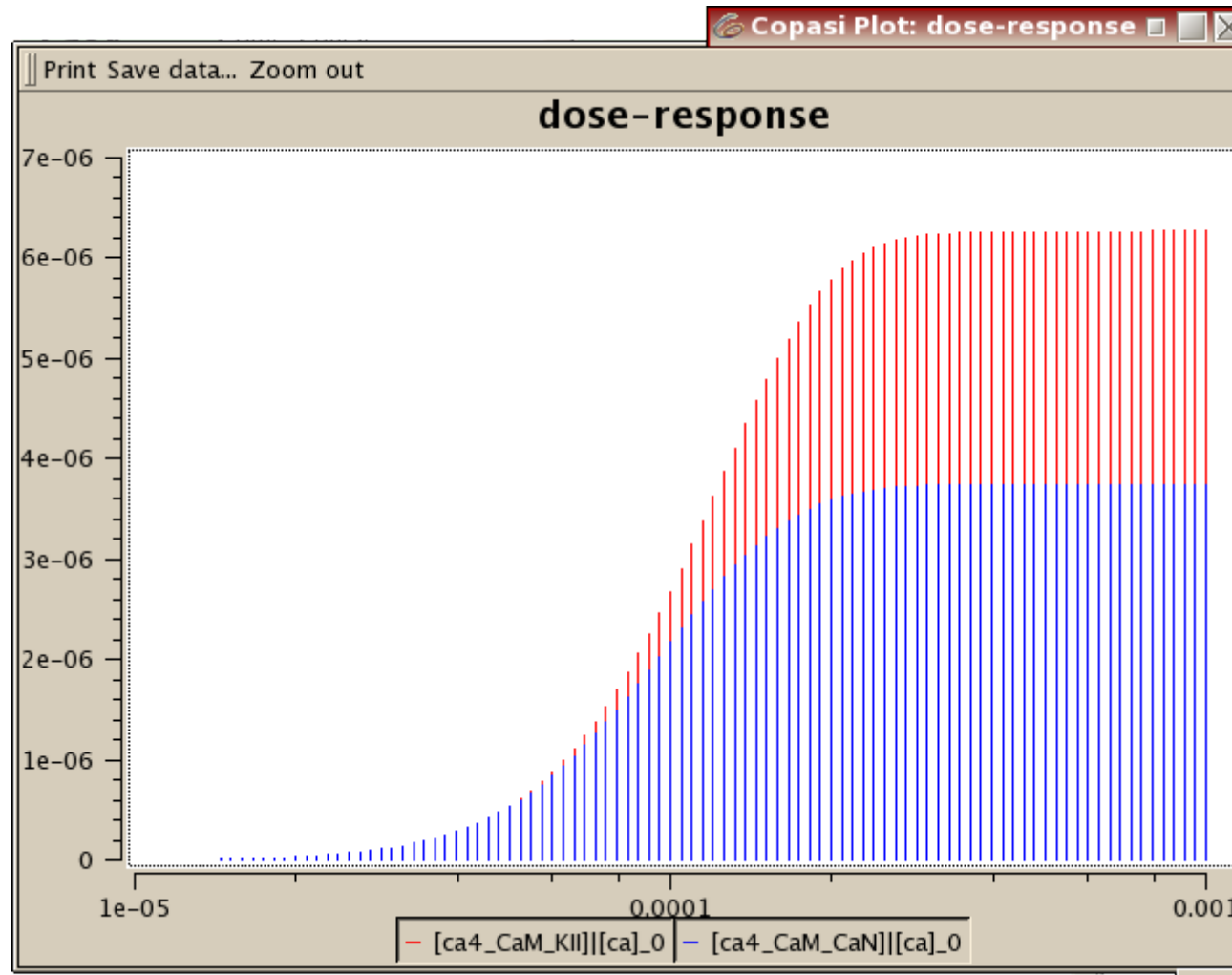


Crouch and Klee (1980)
Biochemistry, 19: 3692-3698

Corresponding induced-fit model



That does not work ...



- Calmodulin bound to three calcium activates calcineurin
 - Kincaid and Vaughan (1986). *PNAS*, 83: 1193-1197
- Calmodulin bound to two calcium can bind CaMKII
 - Shifman et al (2006). *PNAS*, 103: 13968-13973
- Calmodulin affinity for calcium increases once bound to CaMKII
 - Shifman et al (2006) [but many previous reports on other targets: e.g. Burger et al (1983). *JBC*, 258: 14733-14739 ; Olwin et (1984). *JBC* 259: 10949-10955]
- Calcium activates both LTP and LTD through calmodulin
 - Lisman (1989) *PNAS*, 86: 9574-9578
 - High $[Ca^{2+}]$ (high freq) $\cong$ CaMKII ; Low $[Ca^{2+}]$ (low freq) $\cong$ Calcineurin

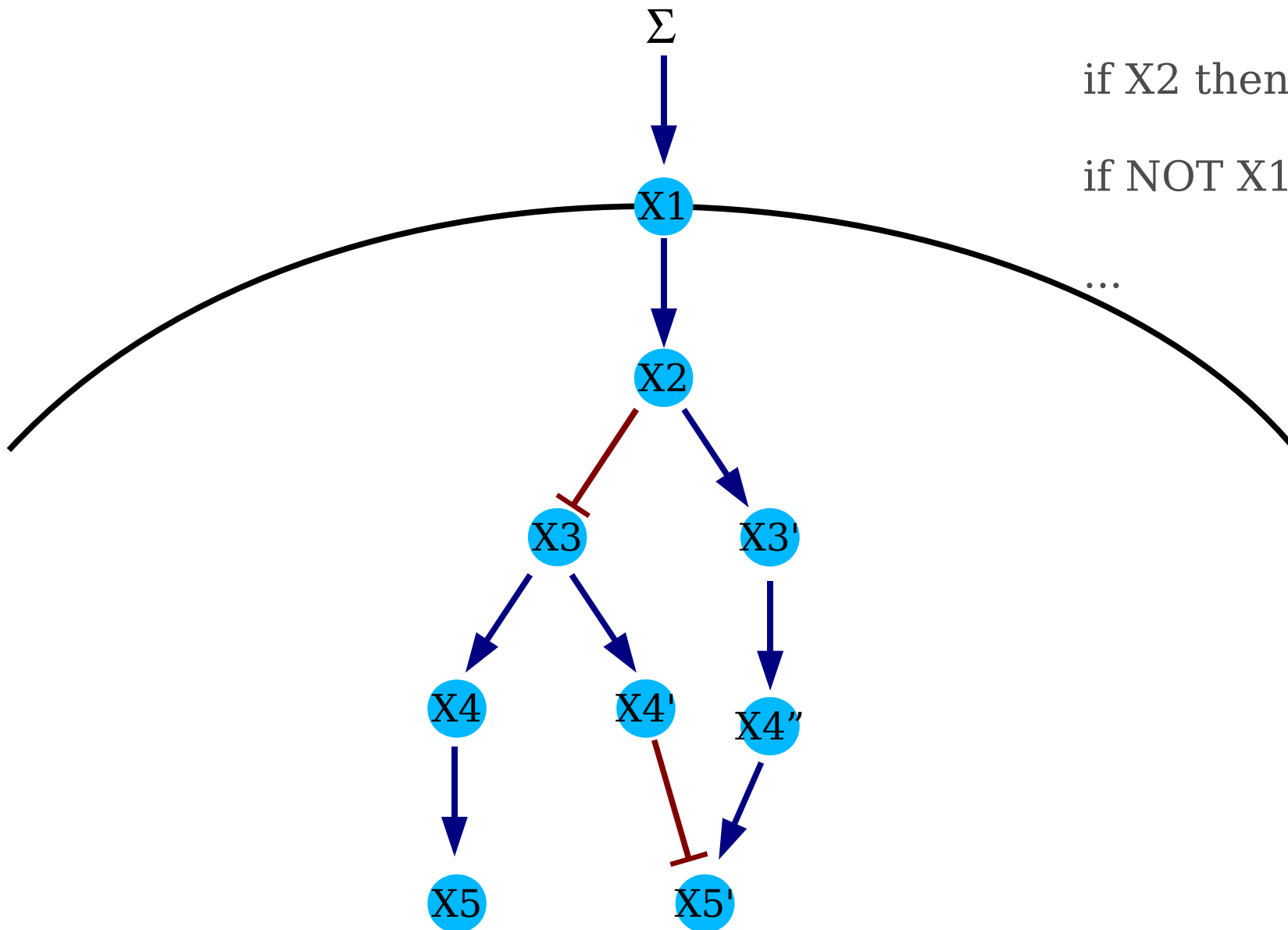
Mistake: signals, activity flow and induction

if X1 then X2

if X2 then not X3

if NOT X1 then NOT X2

...

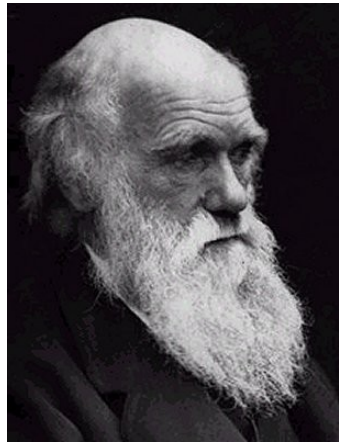


- Induction = BAD
(Lamarck's first law, instructive theory of antibody formation, directed axonal growth, induced-fit ...)

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- Physically meaningless. Calcium has no inertia. Calcium cannot “trigger” a conformational change

Induction Vs Selection

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(Lamarck's first law, instructive theory of antibody formation, directed axonal growth, induced-fit ...)
- Physically meaningless. Calcium has no inertia. Calcium cannot “trigger” a conformational change
- Selection = GOOD
(natural selection, clonal selection, synapse stabilisation, conformation stabilisation ...)

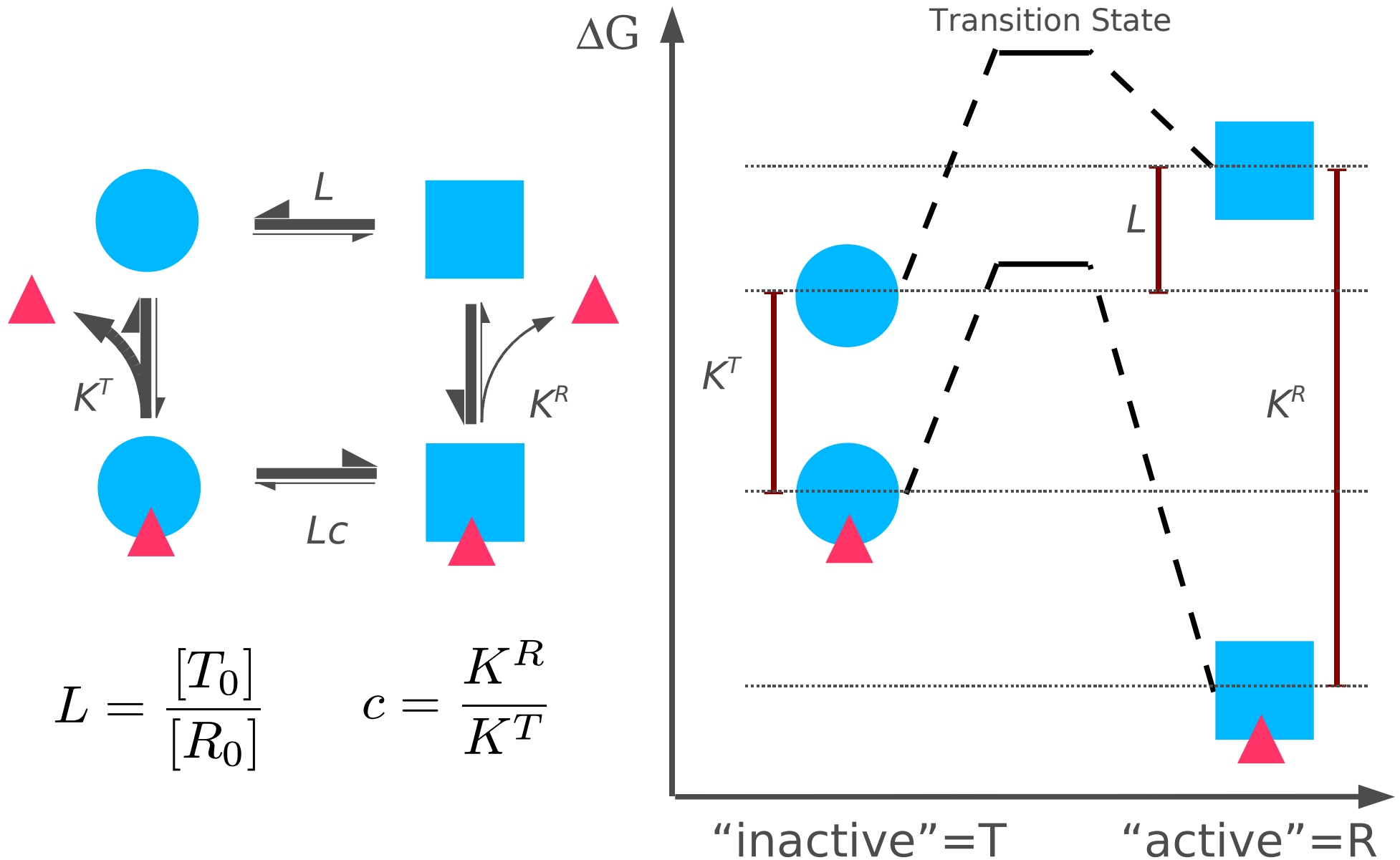


Allostery and state selection

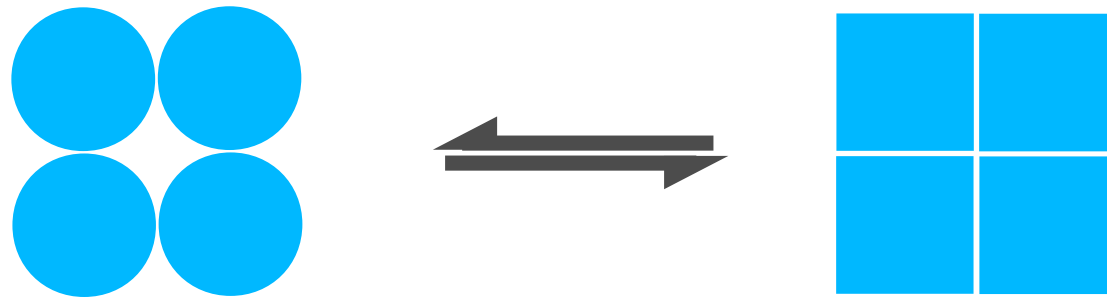
- ➔ Monod, Wyman, Changeux (1965). On the nature of allosteric transitions: a plausible model.
J Mol Biol, 12: 88-118



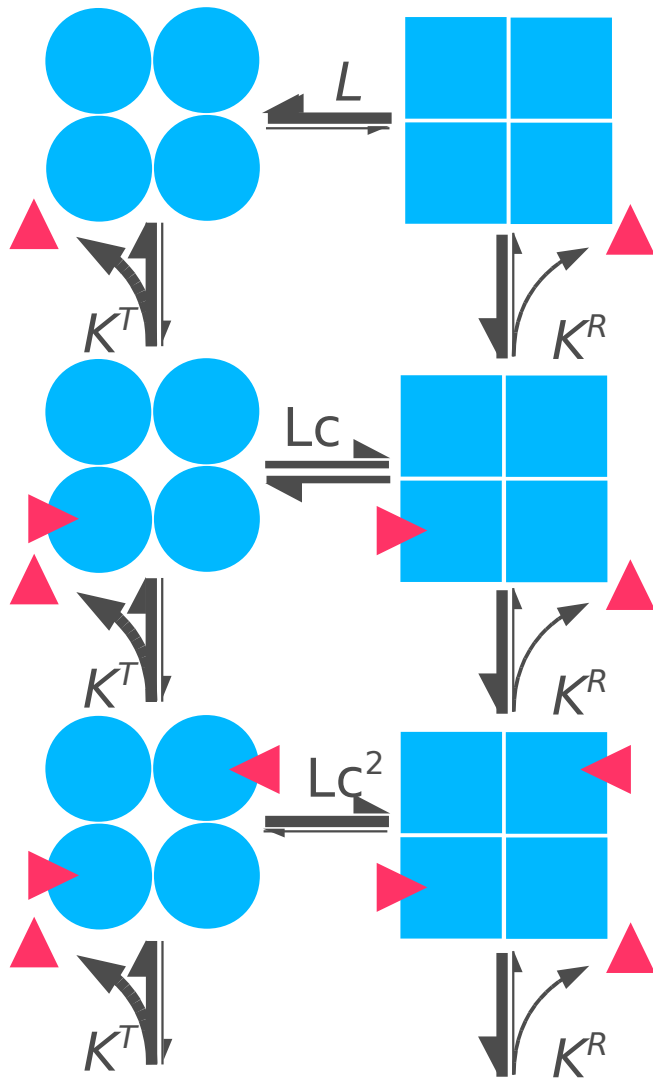
Modulation of thermal equilibria $\neq$ induced-fit



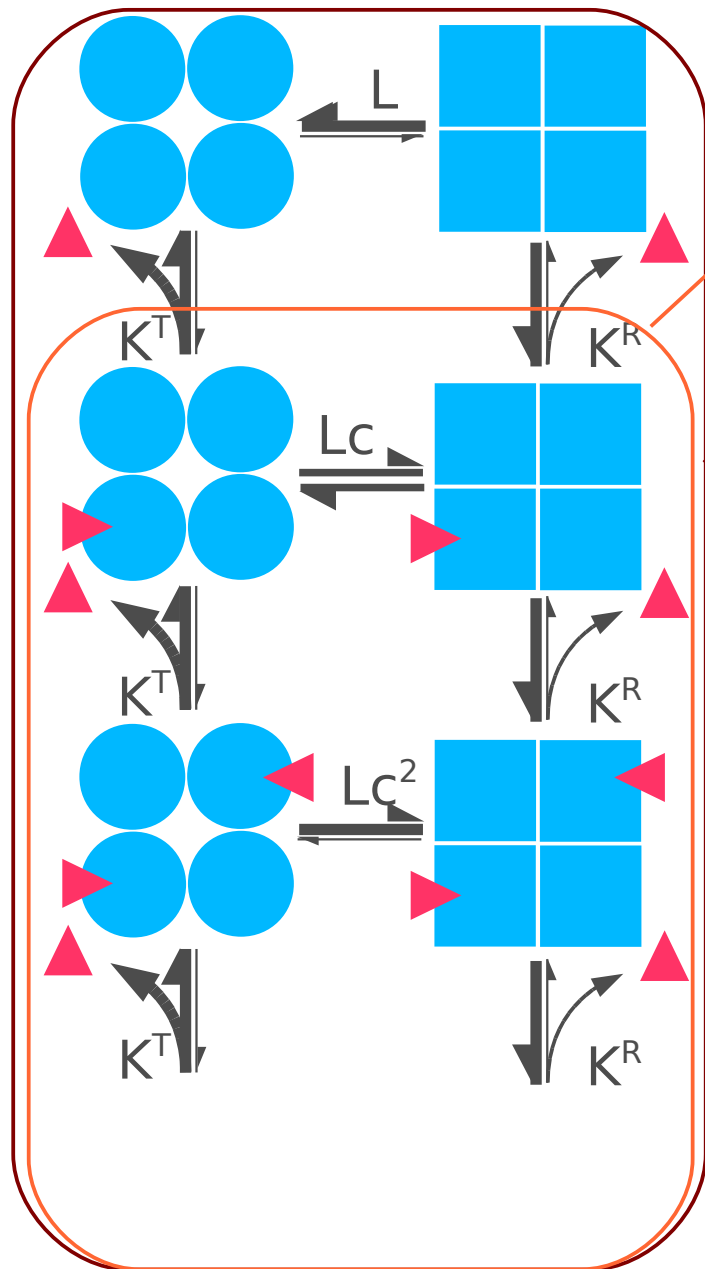
Concerted transitions $\neq$ sequential model



Monod-Wyman-Changeux model



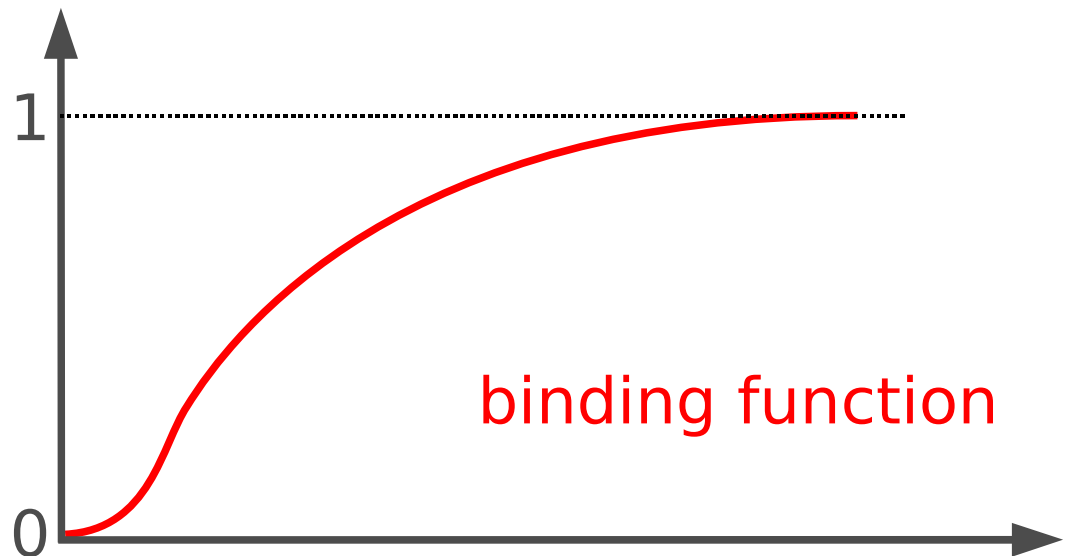
Monod-Wyman-Changeux model



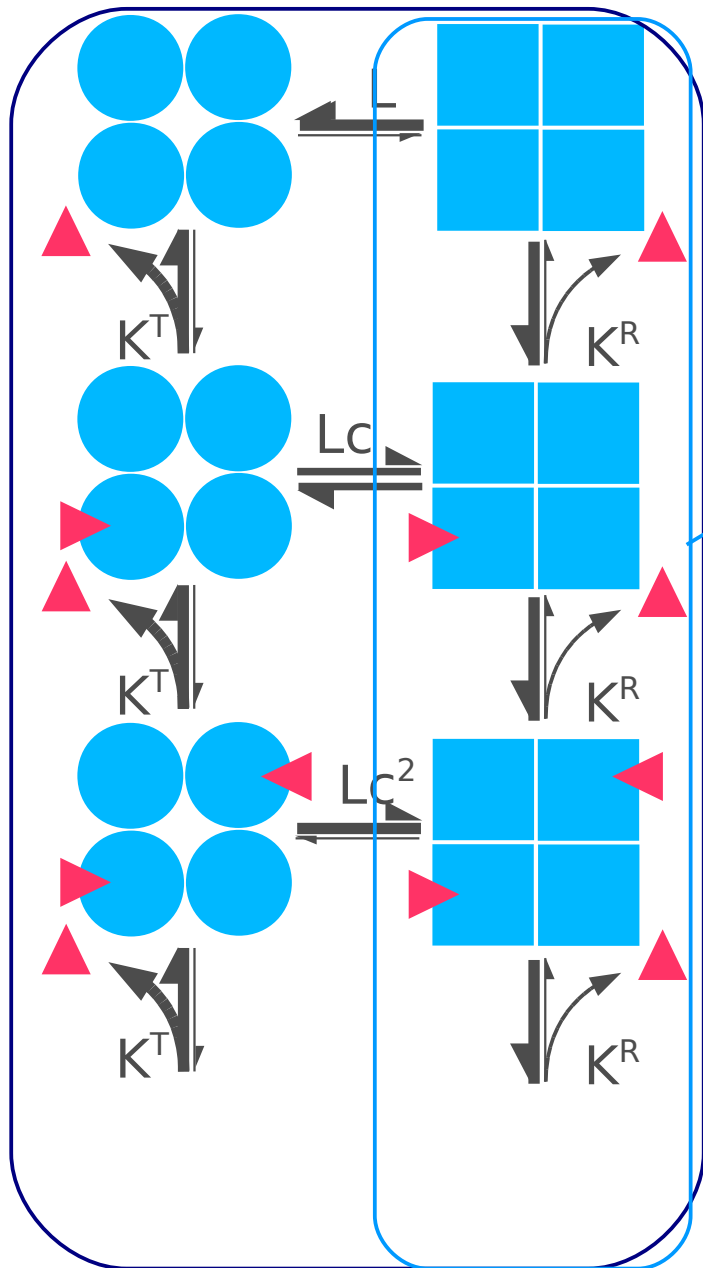
this
that

$$\alpha = \frac{[x]}{K^R}$$

$$\bar{Y} = \frac{\alpha(1 + \alpha)^{n-1} + Lc\alpha(1 + c\alpha)^{n-1}}{(1 + \alpha)^n + L(1 + c\alpha)^n}$$



Monod-Wyman-Changeux model

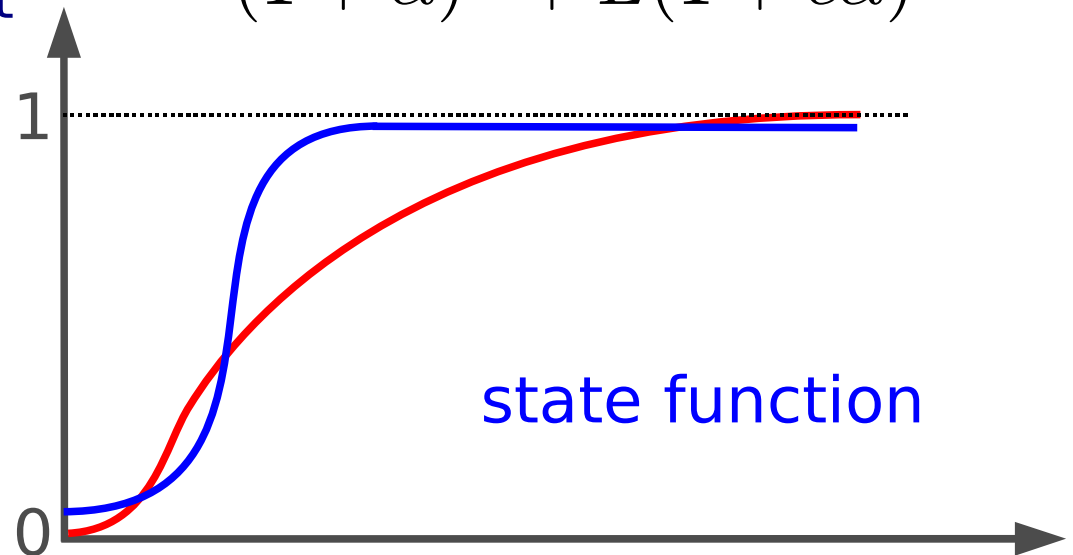


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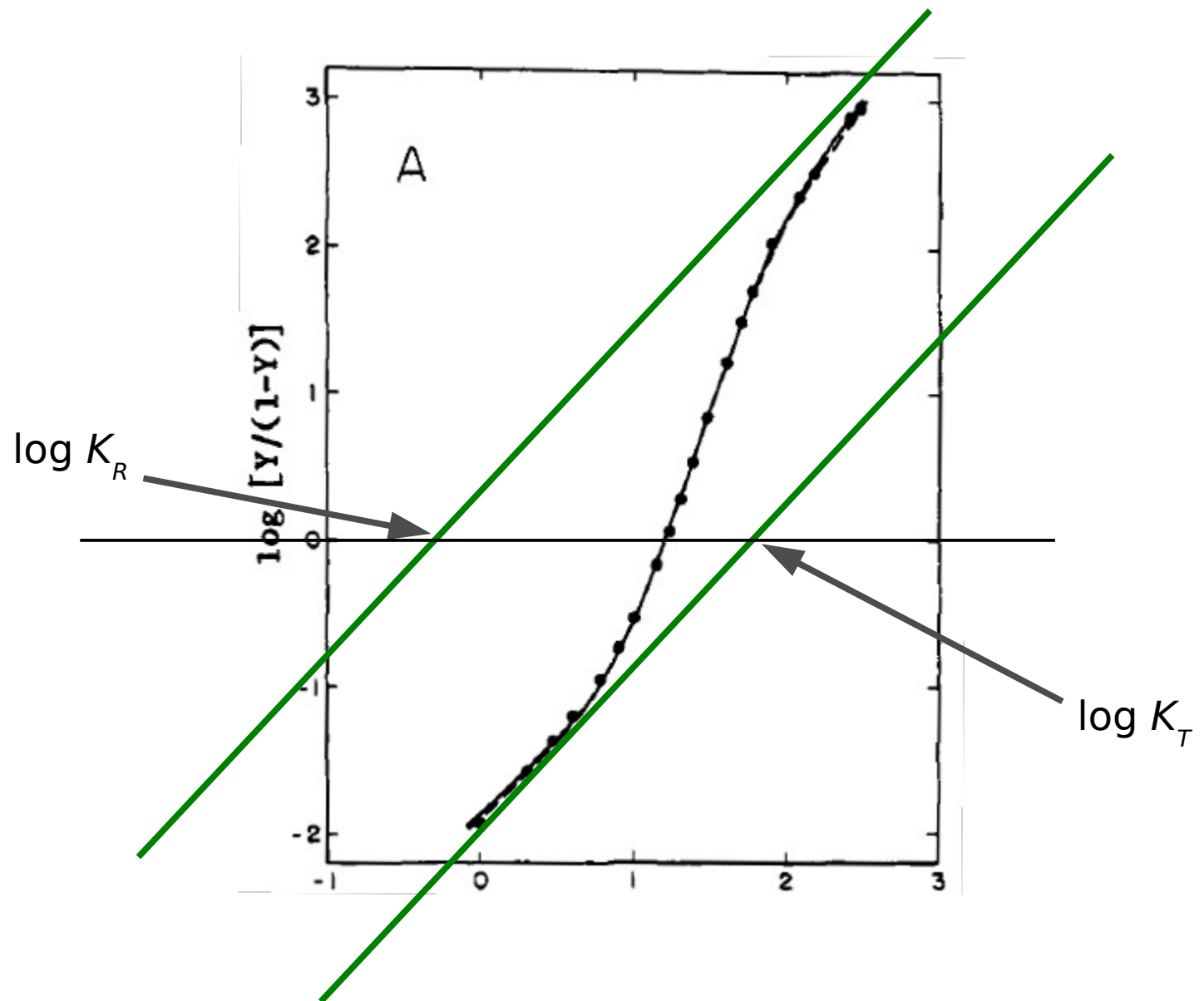
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this
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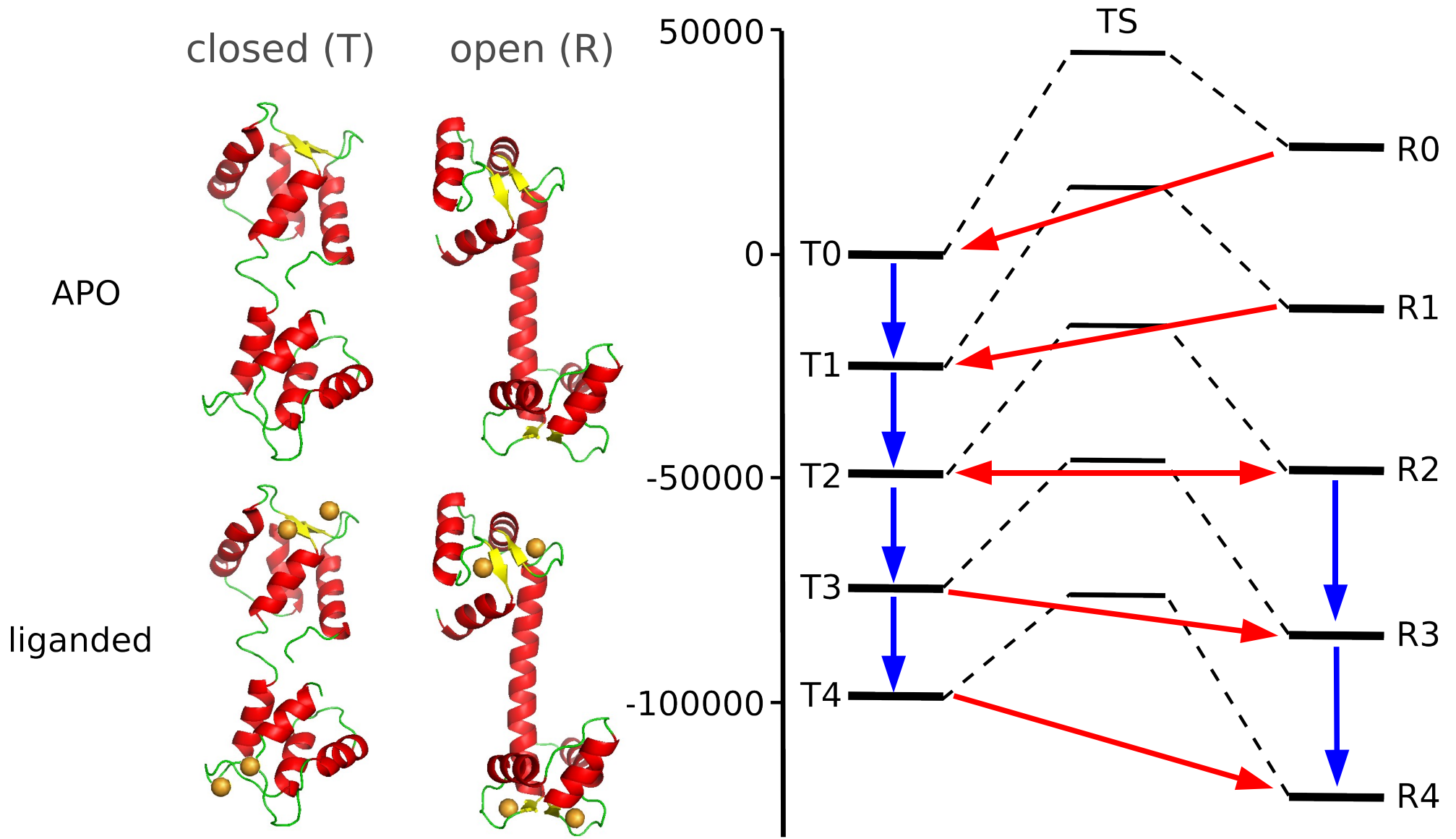
$$\bar{R} = \frac{(1 + \alpha)^n}{(1 + \alpha)^n + L(1 + c\alpha)^n}$$



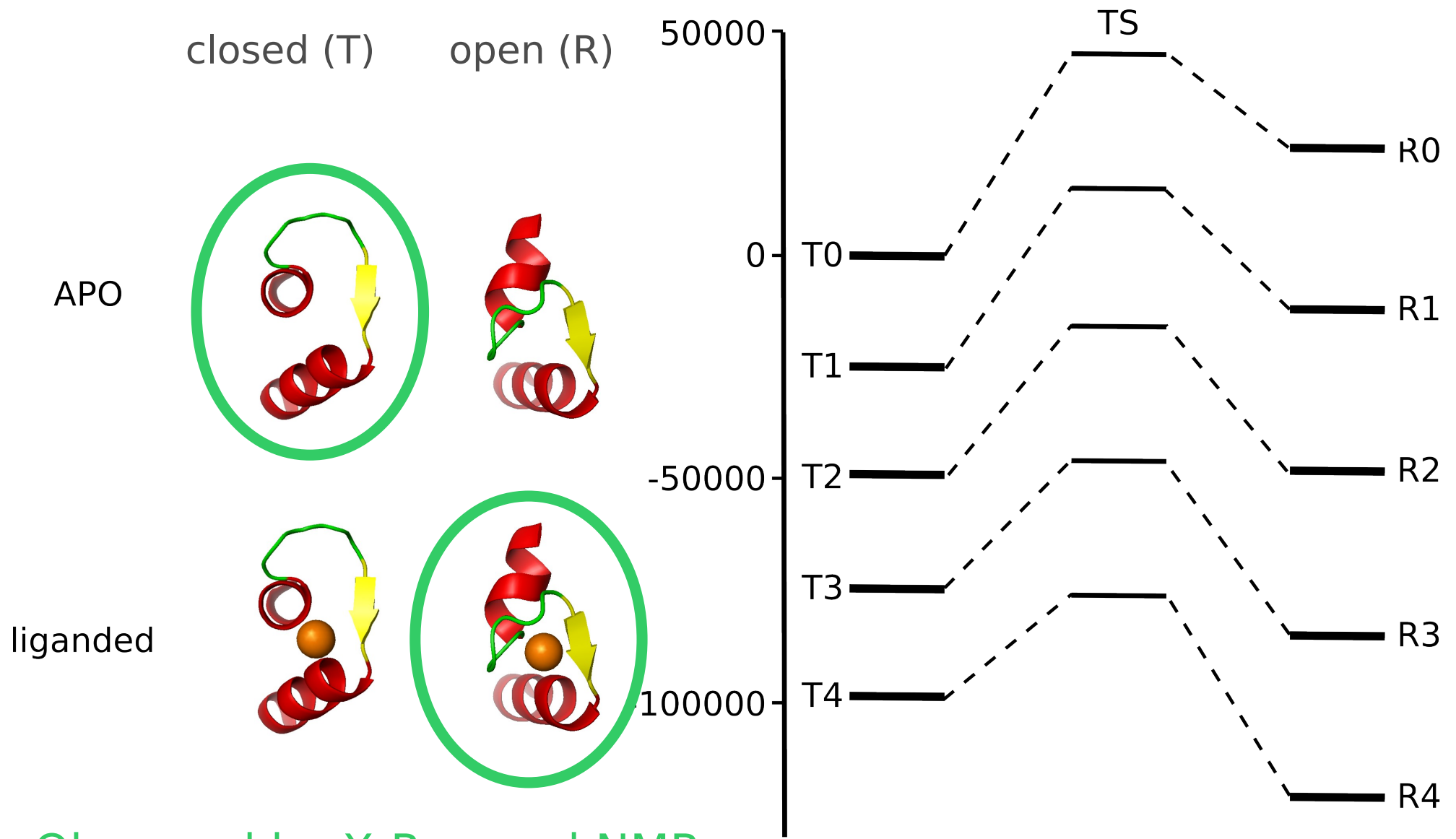
“Hill” Plot for MWC model



Concerted transition

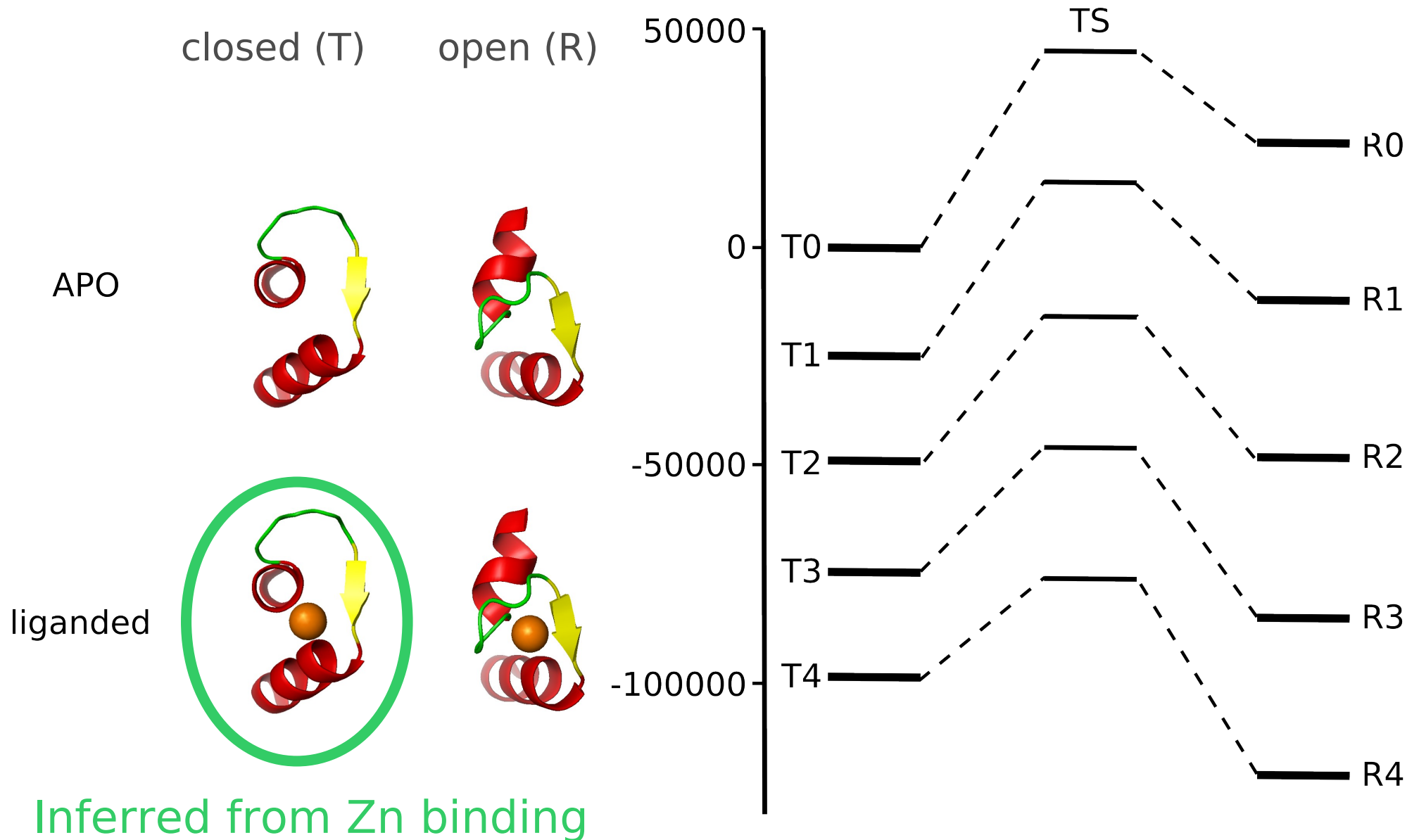


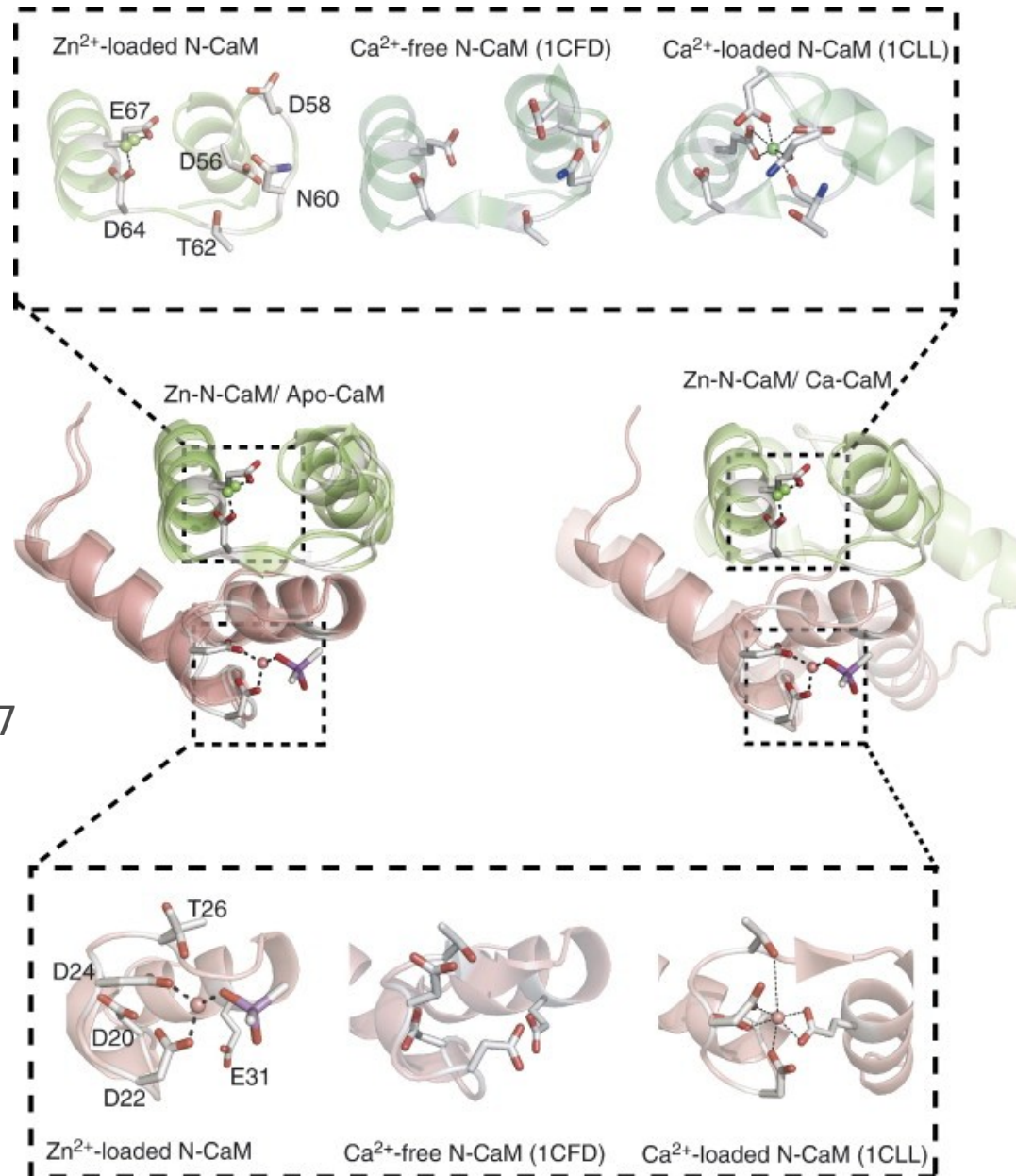
Observation Vs. Prediction



Observed by X-Ray and NMR

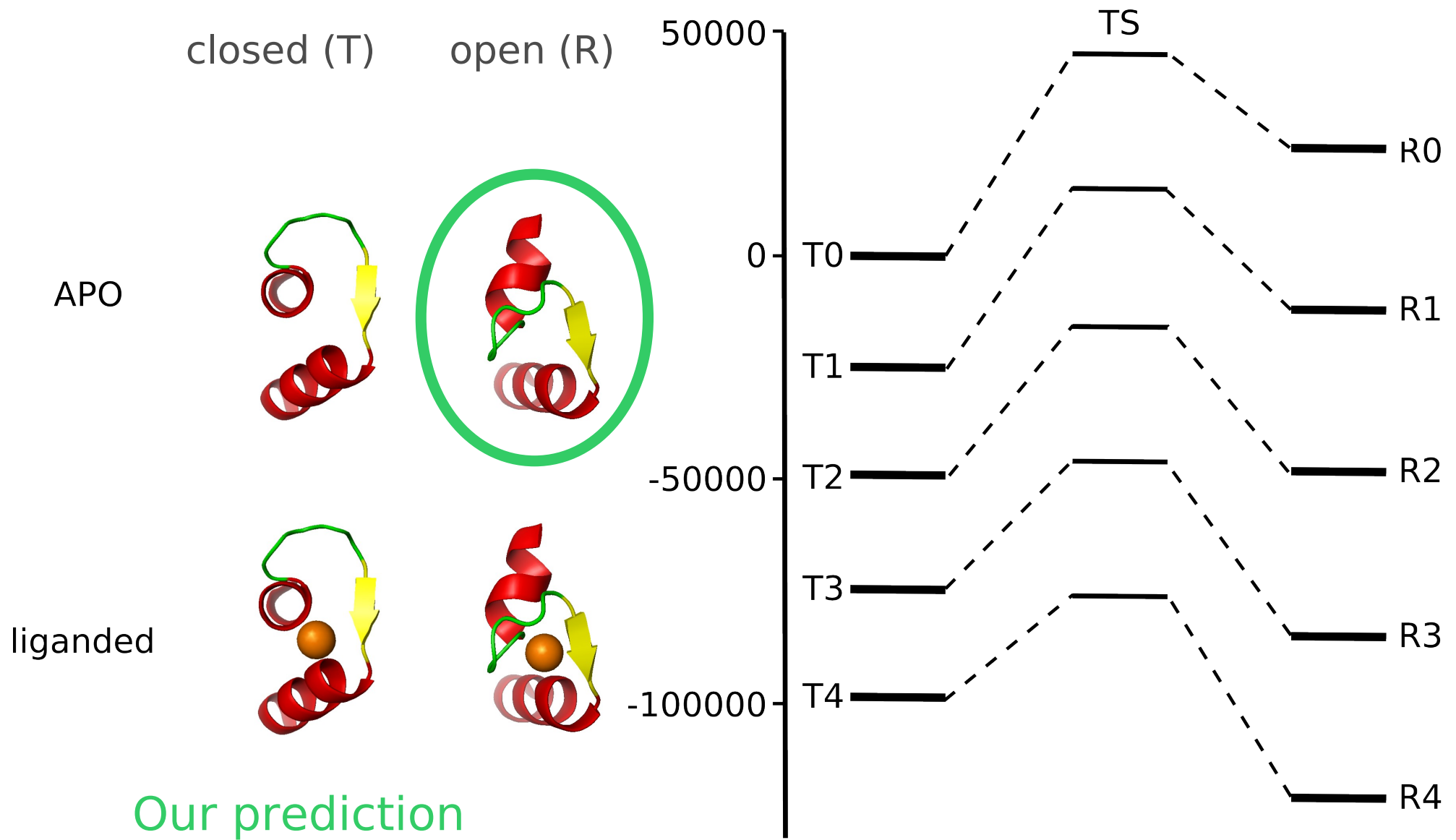
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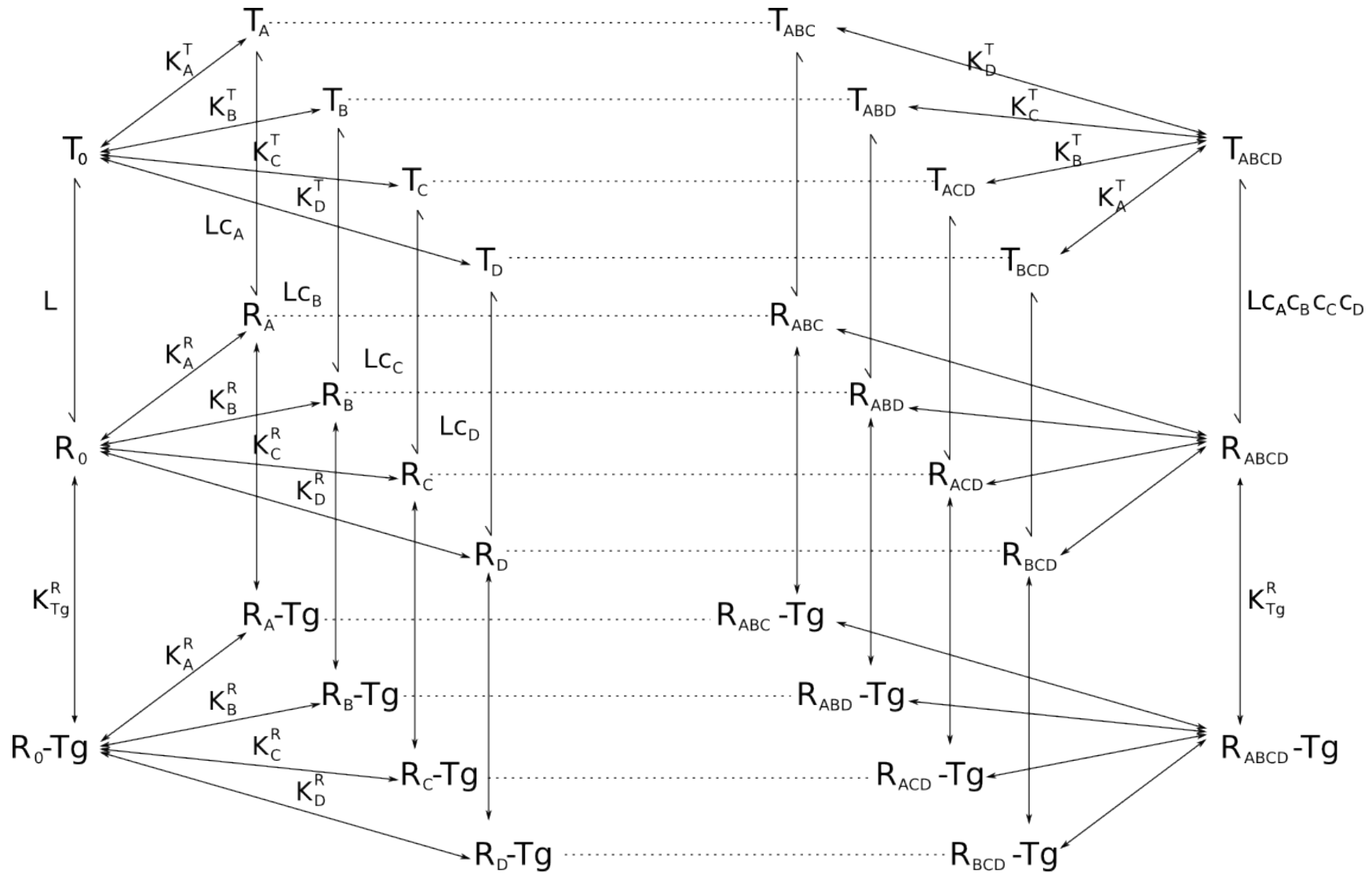


Warren *et al.* (2007).
J Mol Biol 374: 517-527

Observation Vs. Prediction



Full mechanistic thermodynamic model



320 reactions

Extended MWC model necessary for Calmodulin

Based on Rubin and Changeux
(1966) *J Mol Biol*, 21: 265-274

$$\bar{Y} = \frac{\alpha(1 + \alpha)^{n-1} + L \left(\frac{1 + d\beta}{1 + \beta} \frac{1 + e\gamma}{1 + \gamma} \right)^n c\alpha(1 + c\alpha)^{n-1}}{(1 + \alpha)^n + L \left(\frac{1 + d\beta}{1 + \beta} \frac{1 + e\gamma}{1 + \gamma} \right)^n (1 + c\alpha)^n}$$

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Based on Rubin and Changeux
(1966) *J Mol Biol*, 21: 265-274

$$\bar{Y} = \frac{\alpha(1 + \alpha)^{n-1} + L \left(\frac{1 + d\beta}{1 + \beta} \frac{1 + e\gamma}{1 + \gamma} \right)^n c\alpha(1 + c\alpha)^{n-1}}{(1 + \alpha)^n + L \left(\frac{1 + d\beta}{1 + \beta} \frac{1 + e\gamma}{1 + \gamma} \right)^n (1 + c\alpha)^n}$$

$$\bar{Y} = \frac{\frac{1}{n} \left(\sum_i \left(\alpha_i \prod_{j \neq i} (1 + \alpha_j) \right) + L \prod_k \left(\frac{1 + e_k \gamma_k}{1 + \gamma_k} \right) \sum_i \left(c_i \alpha_i \prod_{j \neq i} (1 + c_j \alpha_j) \right) \right)}{\prod_i (1 + \alpha_i) + L \prod_k \left(\frac{1 + e_k \gamma_k}{1 + \gamma_k} \right) \prod_i (1 + c_i \alpha_i)}$$

Any number of different
sites per protomer.
Several protomers can
be carried by one subunit

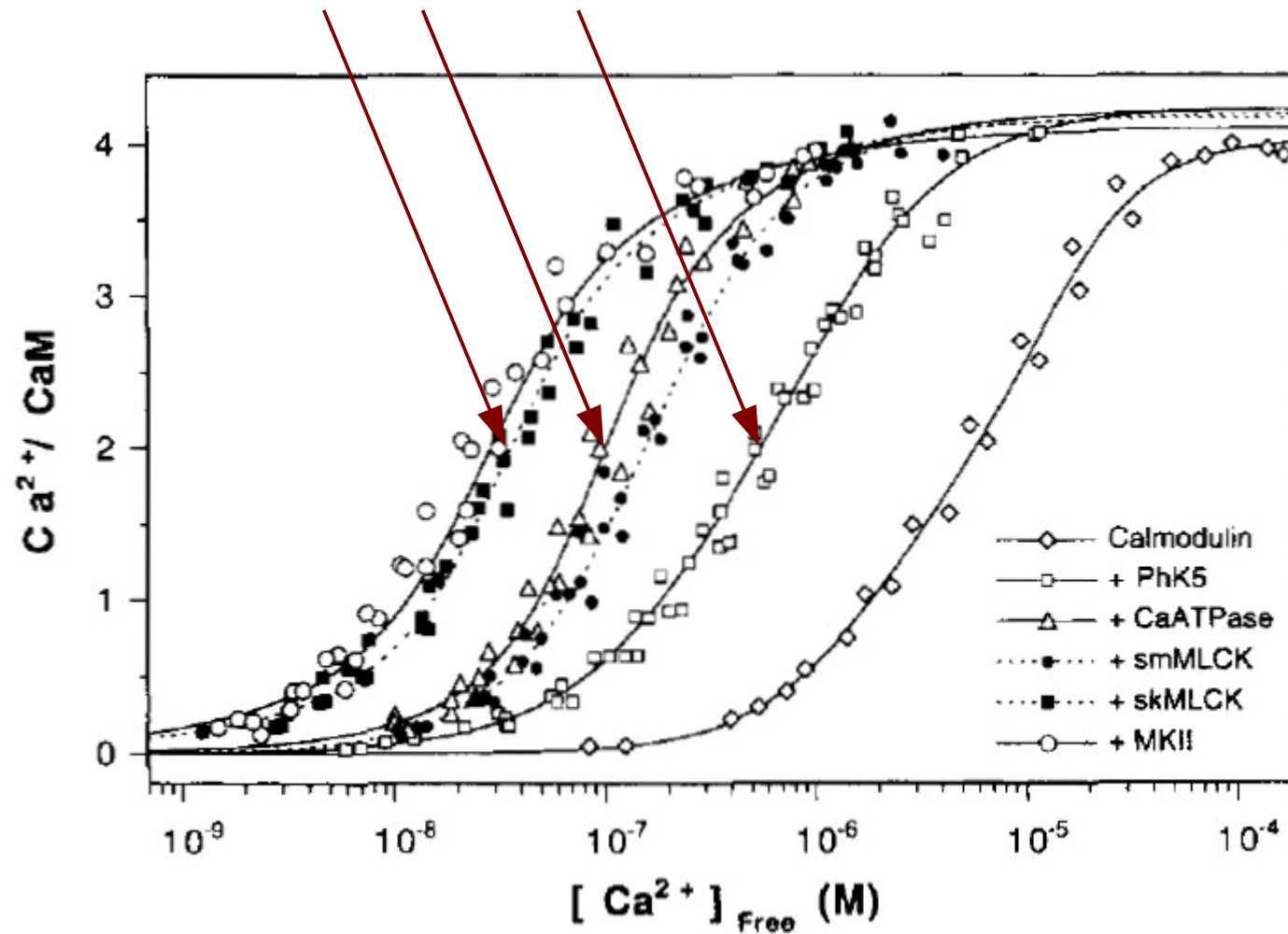
- $\alpha_i = [\text{ligand}] / K_{i,\text{lig}}^R$
- $\gamma_k = [\text{modulator}] / K_{k,\text{mod}}^R$
- $c_i = K_{i,\text{lig}}^R / K_{i,\text{lig}}^T$
- $e_k = K_{k,\text{mod}}^R / K_{k,\text{mod}}^T$

Simplification of the model for finding L and c

- Hypothesis for the whole model: free energy of conformational transition is evenly distributed: c is unique
- Additional simplification to determine L : affinities are identical

$$\bar{Y} = \frac{\alpha(1 + \alpha)^3 + L\left(\frac{1 + \gamma e}{1 + \gamma}\right)c\alpha(1 + c\alpha)^3}{(1 + \alpha)^4 + L\left(\frac{1 + \gamma e}{1 + \gamma}\right)(1 + c\alpha)^4}$$

Targets as allosteric effectors



Peersen et al. (1997) *Prot Sci*, 6: 794-807

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- Model constraints for the determination of c and L
 - Ca binding in presence of target: none, skMLCK, PhK5, CaATPase (Peersen et al (1997) Prot Sci 6: 794-807). Concentration at 50% saturation.
 - 100 000 parameter sets plus least-square
 - 13 identical minima. e for skMLCK is 10^{-15} , which can be taken as skMLCK binding only to R state.

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$L=20670$
 $c=3.96.10^{-3}$

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Relaxation of the model for finding *Ki*

- Determination of individual affinities:

$$\bar{Y} = 0.25 \frac{\sum_i \left(\alpha_i \prod_j (1 + \alpha_j) \right) + L \sum_i \left(c \alpha_i \prod_j (1 + c \alpha_j) \right)}{\prod_i (1 + \alpha_i) + L \prod_i (1 + c \alpha_i)}$$

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- Model constraints for calcium dissociation constants
 - Complete CaM (Bayley et al (1996) *Prot Sci* 5: 1215-1228)
 - N and C term Mutants (Shifman et al (2006) *PNAS*, 103: 13968-13973)
 - R-only - skMLCK (Peersen et al (1997) *Prot Sci* 6: 794-807)
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 - Systematic logarithmic sampling of the affinity space (coarse-grained, 50 values per affinity, then refined 66 values per affinity) = 25 millions parameter sets

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$$K_A^R = 8.32 \cdot 10^{-6}$$

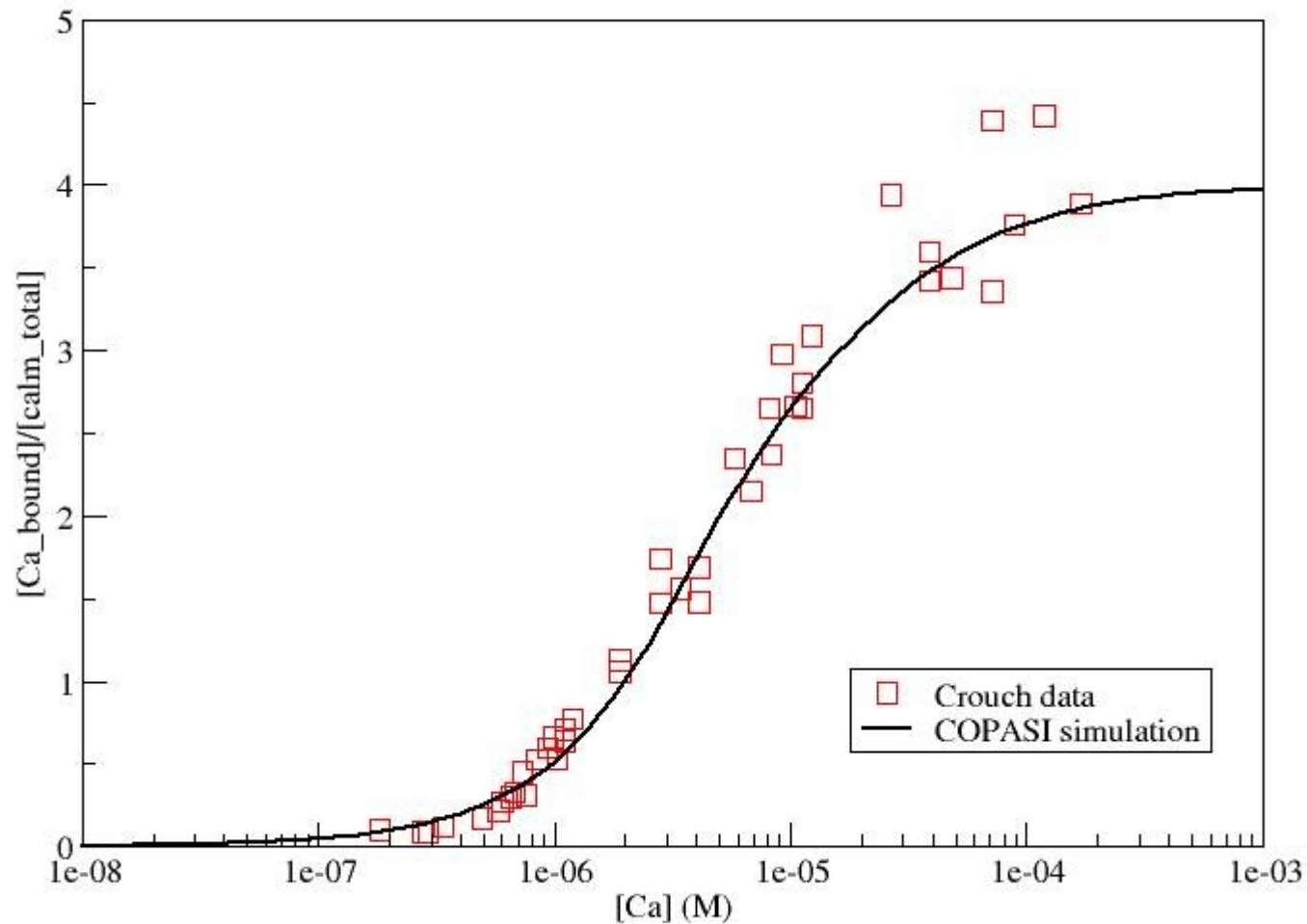
$$K_B^R = 1.66 \cdot 10^{-8}$$

$$K_C^R = 1.74 \cdot 10^{-5}$$

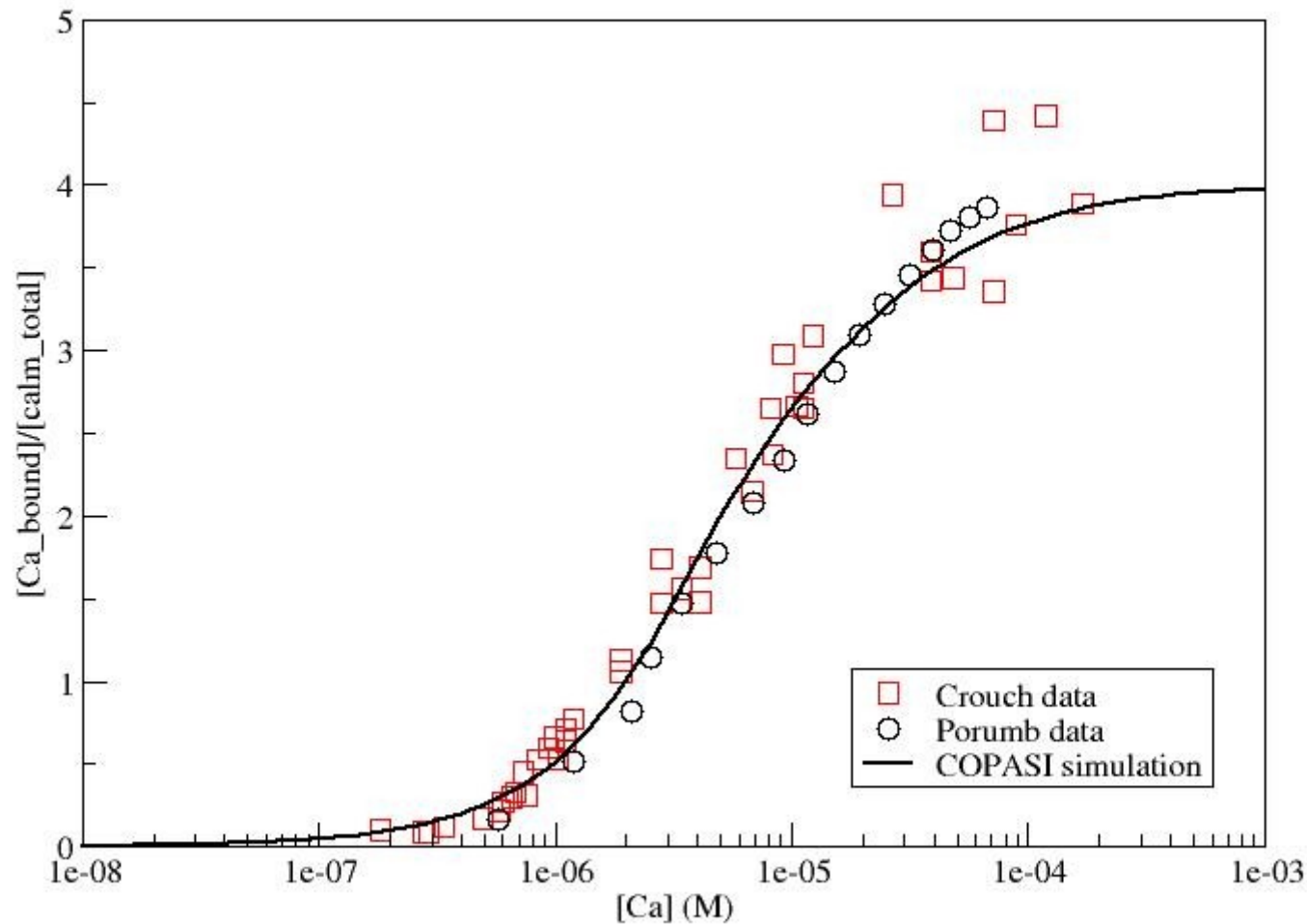
$$K_D^R = 1.45 \cdot 10^{-8}$$

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 - Complete CaM (Bayley et al (1996) *Prot Sci* 5: 1215-1228)
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Comparison with experiments

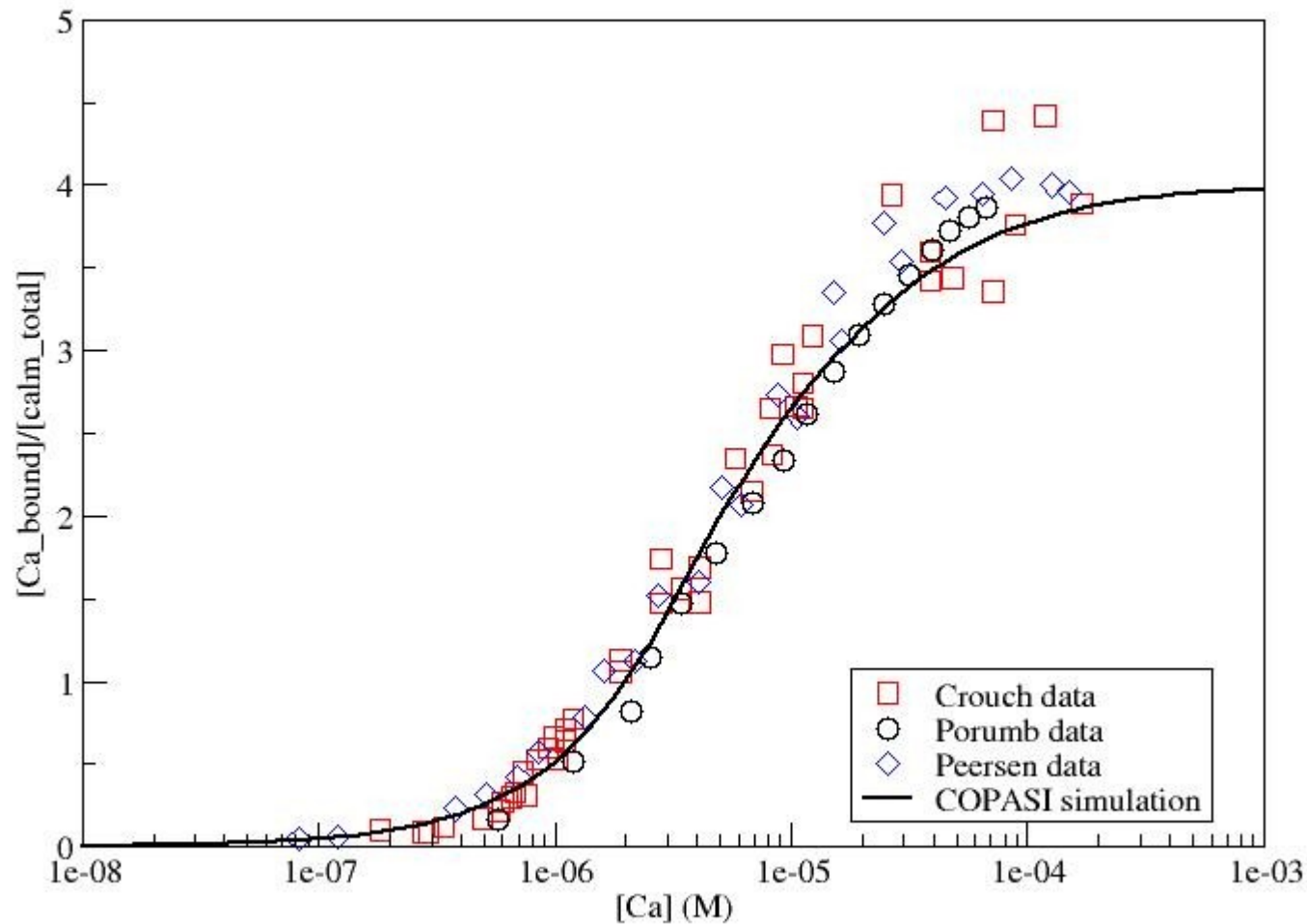


Comparison with experiments

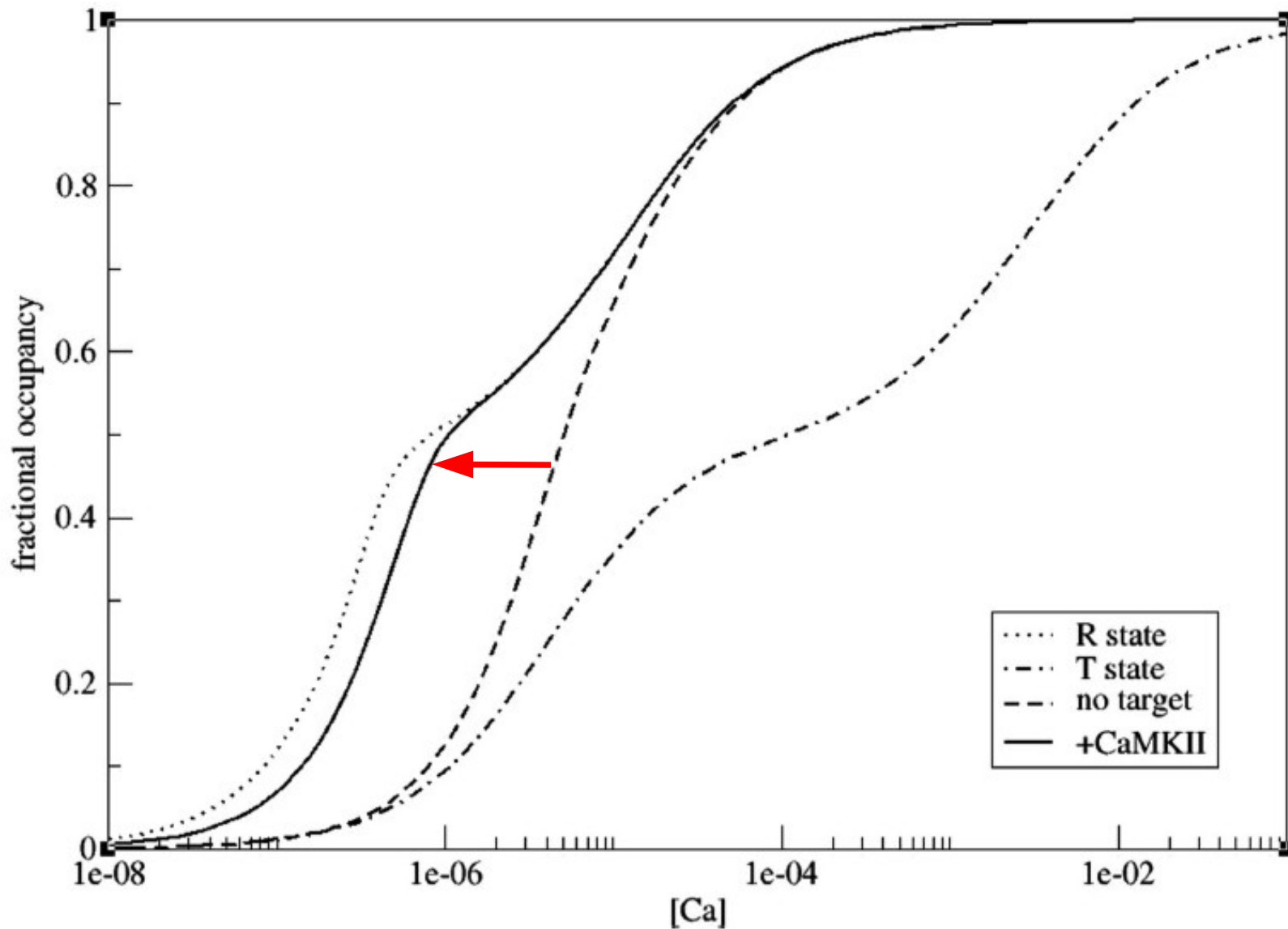


Porumb et al (1994) Anal Biochem 220: 227-237

Comparison with experiments



Binding to target increases the affinity for Ca^{2+}



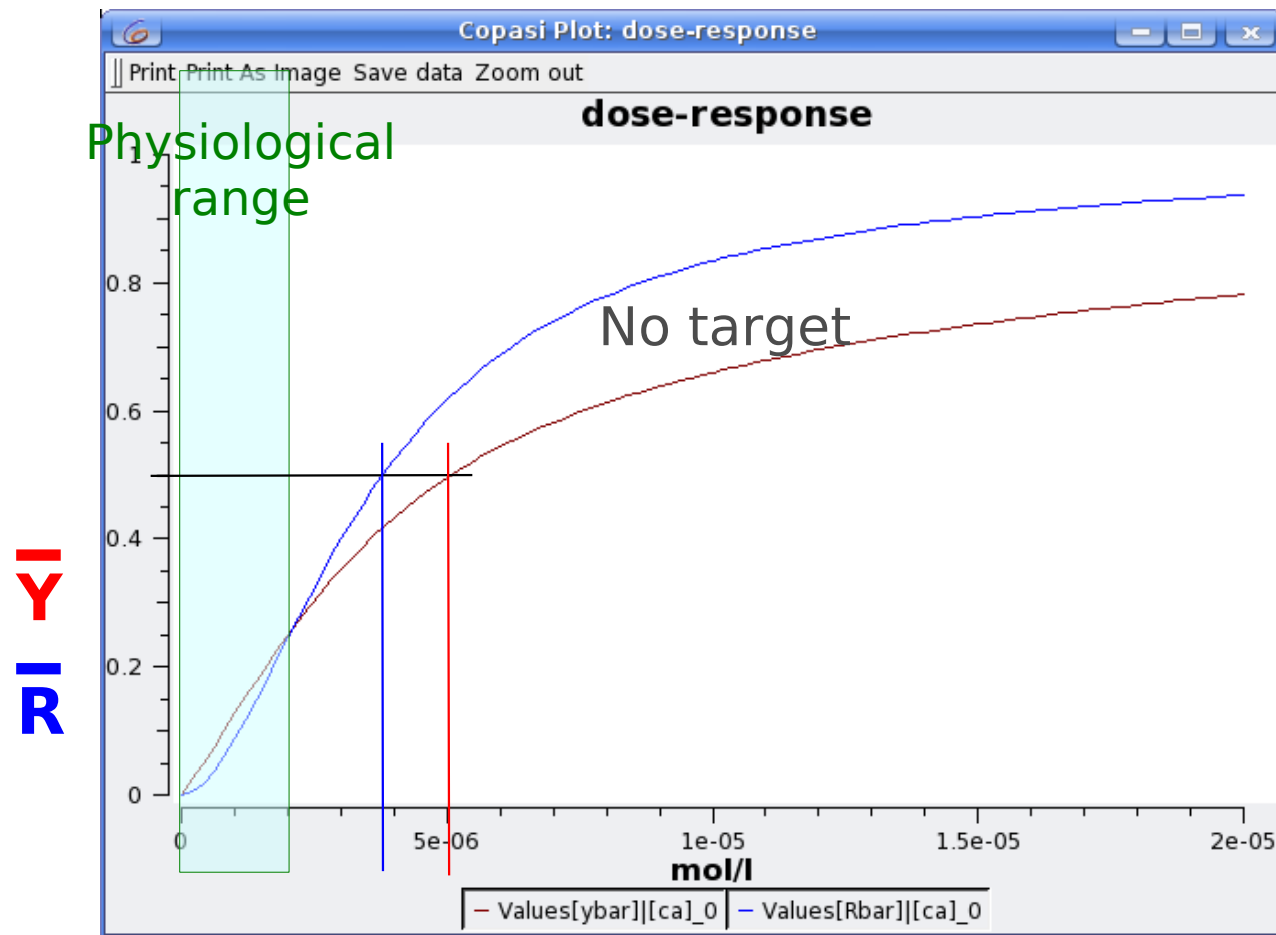
Activity of unsaturated calmodulin

- Fractional activity depends on the number of calcium ions bound. E.g.:

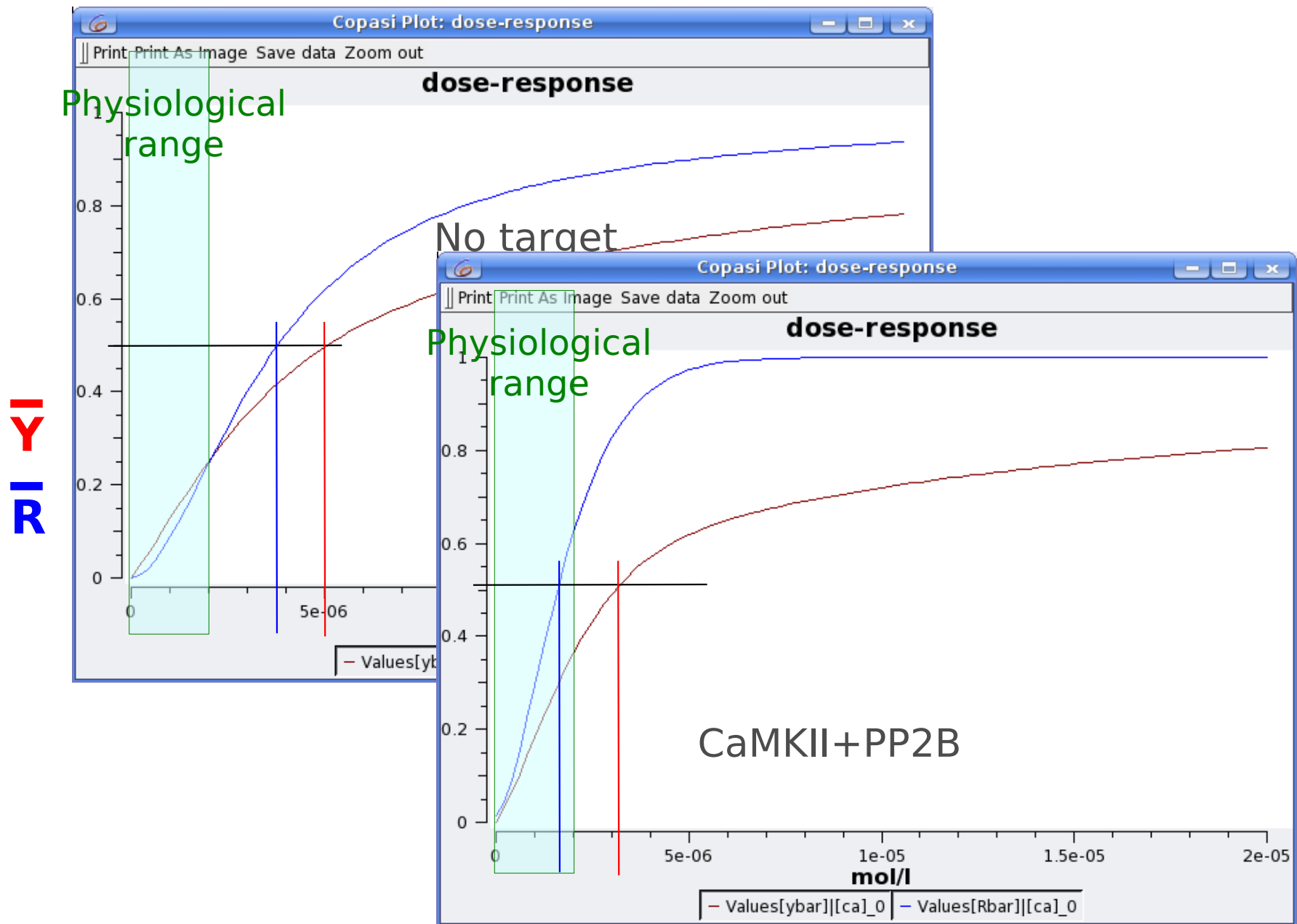
$$\frac{R_2}{T_2} = \frac{1}{L \cdot c^2}$$

- $R_0/T_0 = 1/20000$ (1/L)
- $R_1/T_1 = 1/170$
- $R_2/T_2 = 0.69$ → half-saturation = equi-probability
- $R_3/T_3 = 80$
- $R_4/T_4 = 10000$

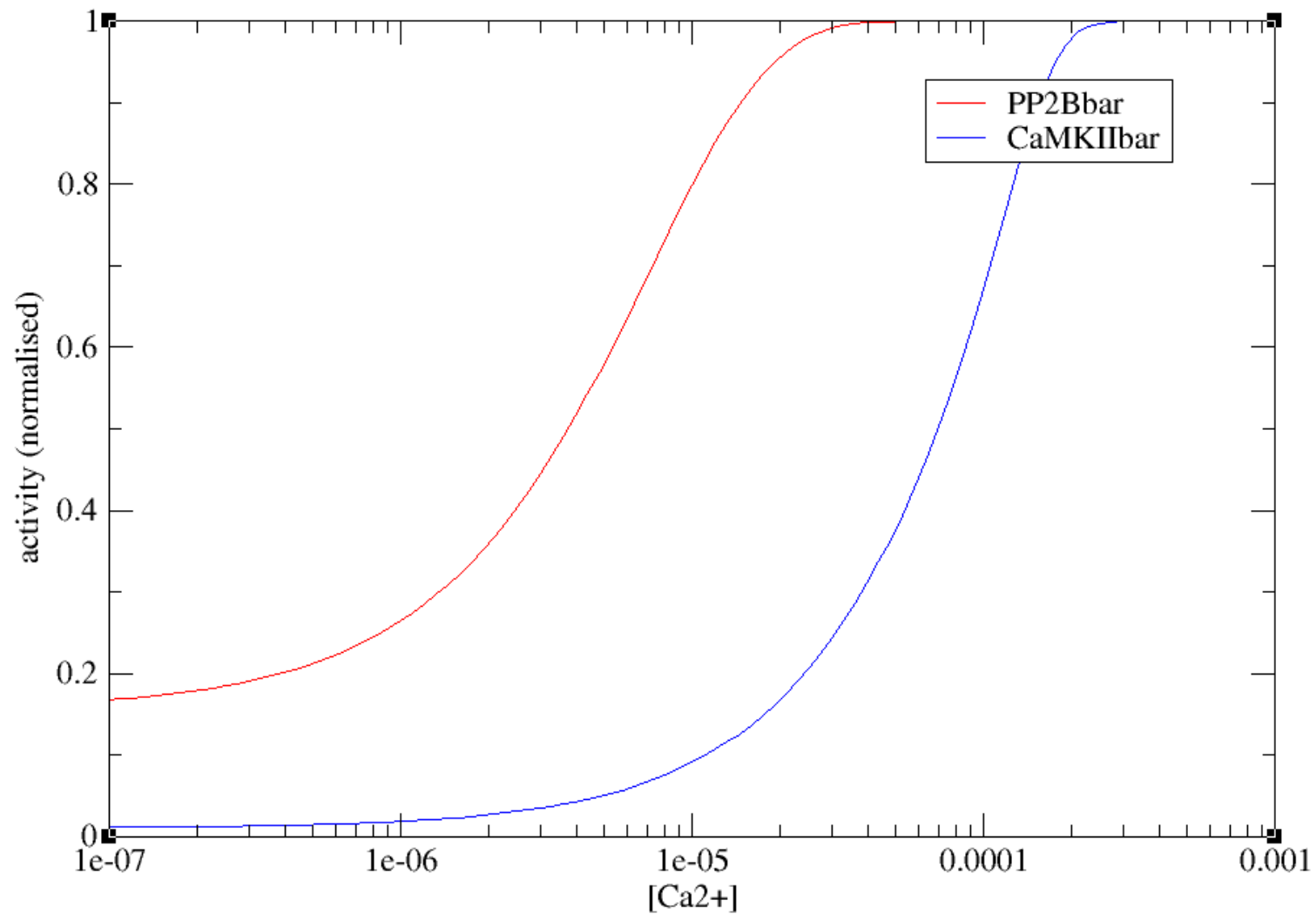
But ... we're out of the physiological range?



This is Systems Biology!

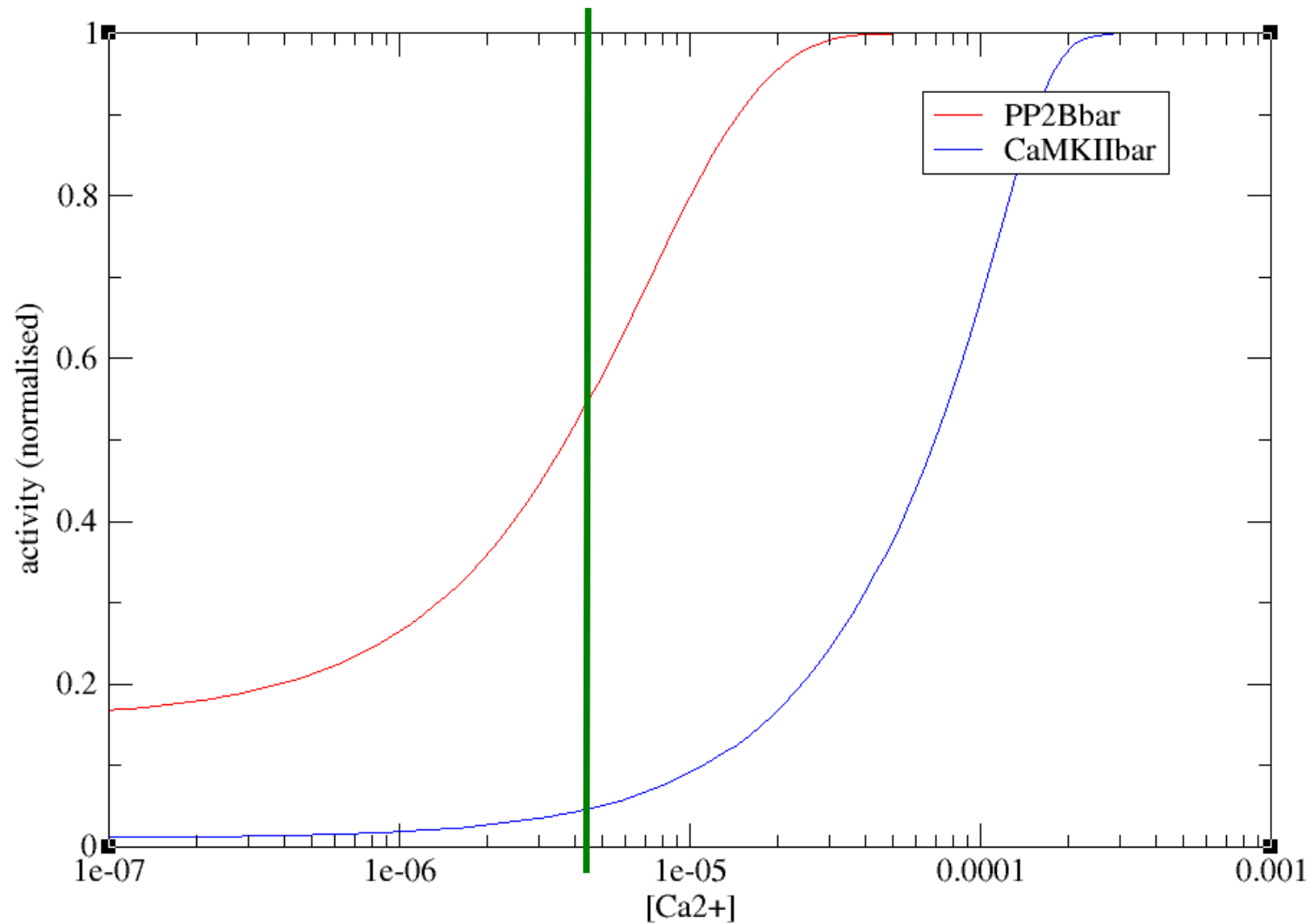


Bidirectional synaptic plasticity

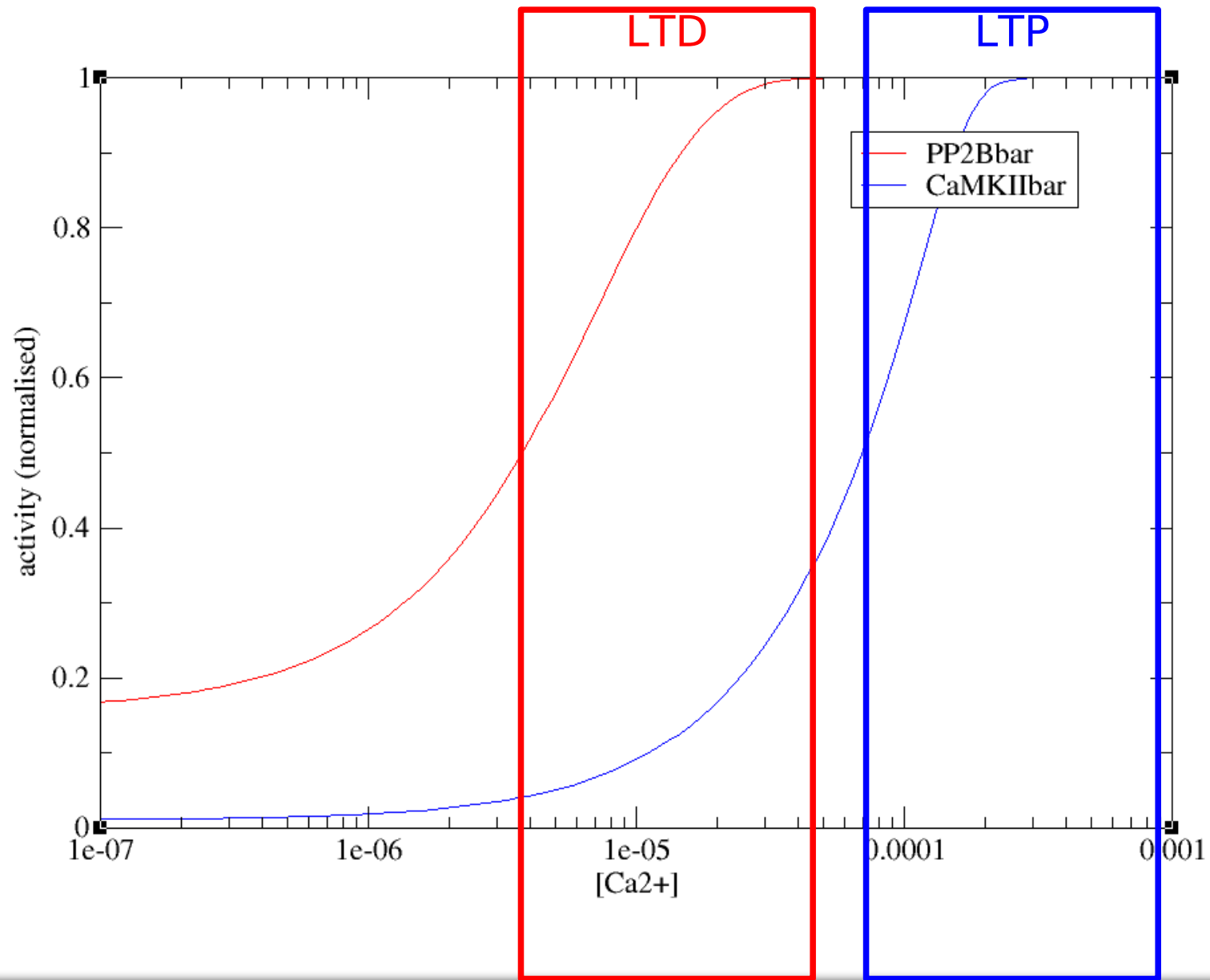


Bidirectional synaptic plasticity

half saturation of calmodulin

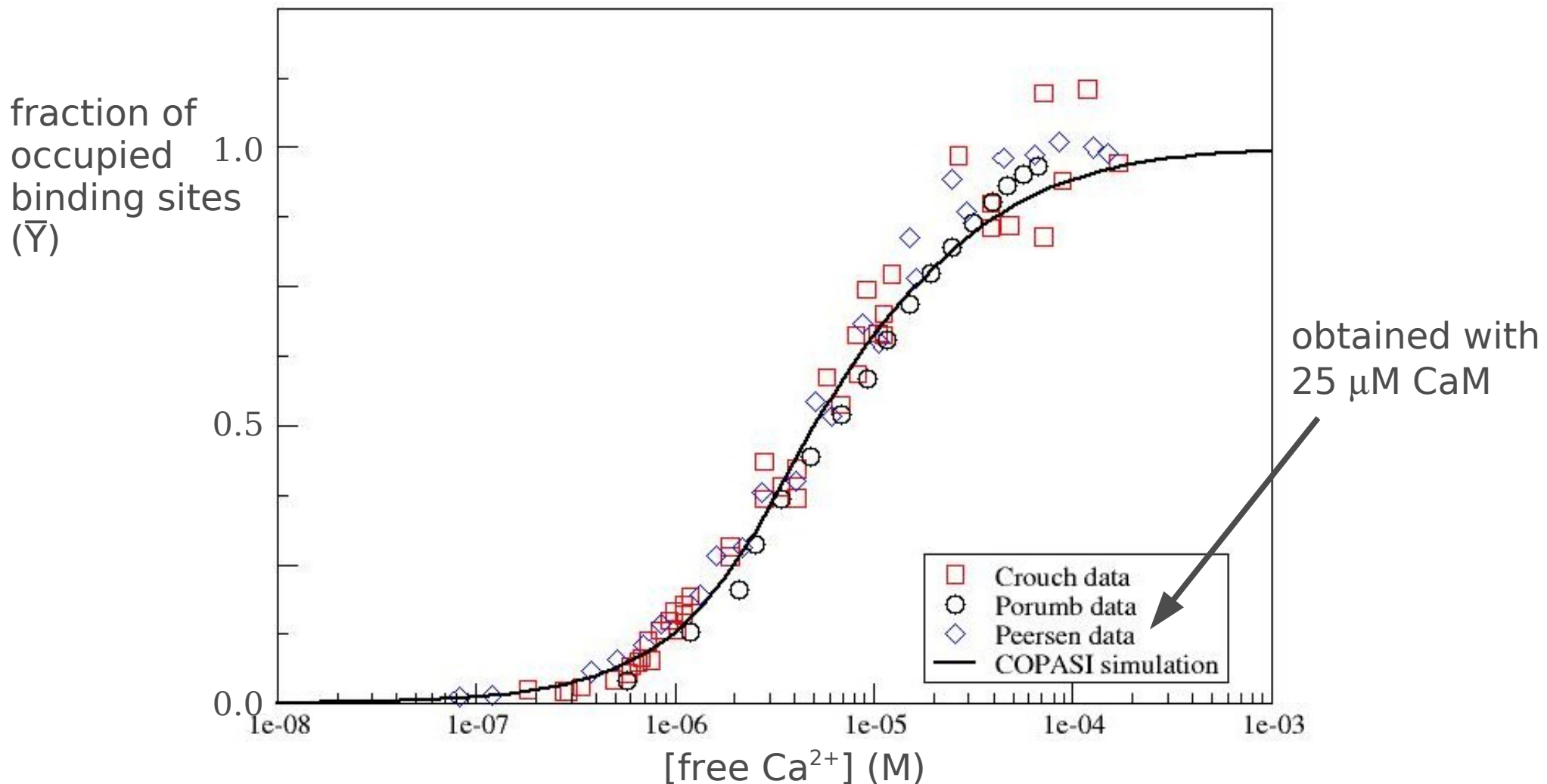


Bidirectional synaptic plasticity



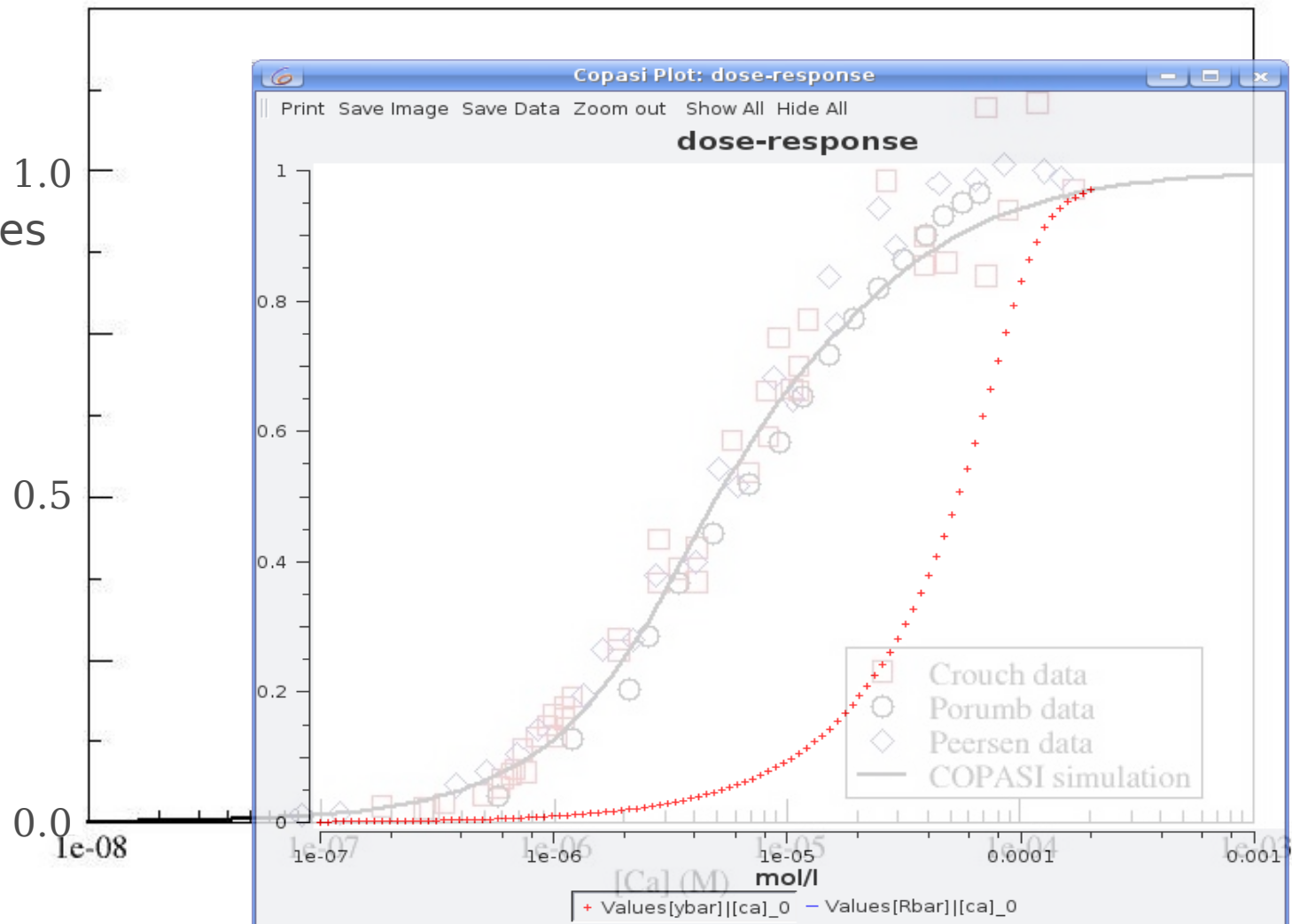
- We designed an allosteric model of Calmodulin, based on only two states for the EF hands, both binding calcium with different affinities, and a concerted transition for all 4 EF hands. We parametrised the model with experimental data-sets.
 - The model fits independent experimental datasets.
 - The affinity of CaM for calcium increases upon binding of the target.
 - CaM can be significantly in the open state even with less than 4 calcium bounds.
 - CaM can bind its targets even when with less than 4 calcium bounds.
- The model displays an activation of the sole PP2B at low concentration of calcium, while high concentrations activate both PP2N and CaMKII.

Allosteric model of Calmodulin function



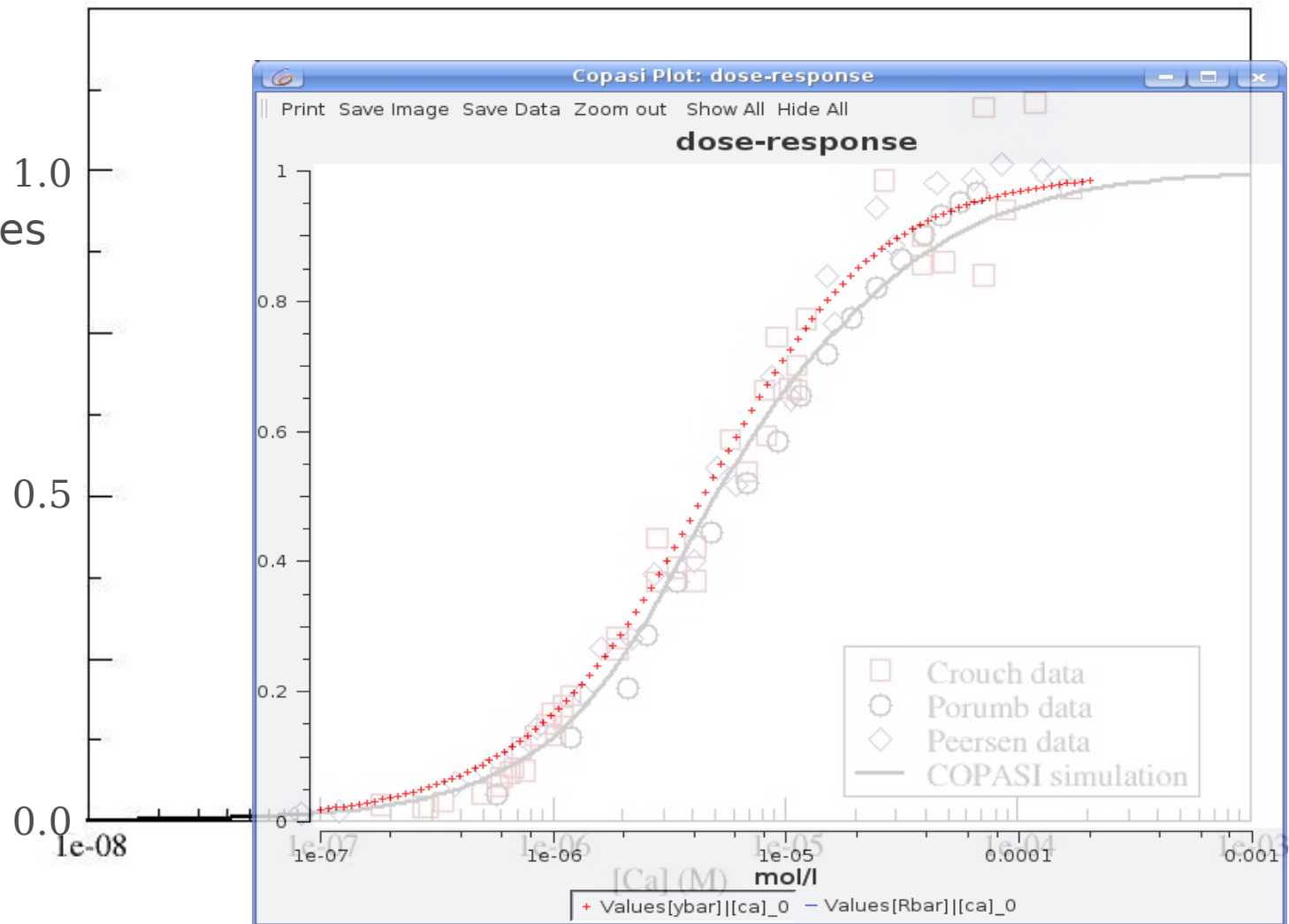
Calcium dose-response on 25 μM Calmodulin

fraction of
occupied
binding sites
($\bar{Y}$)

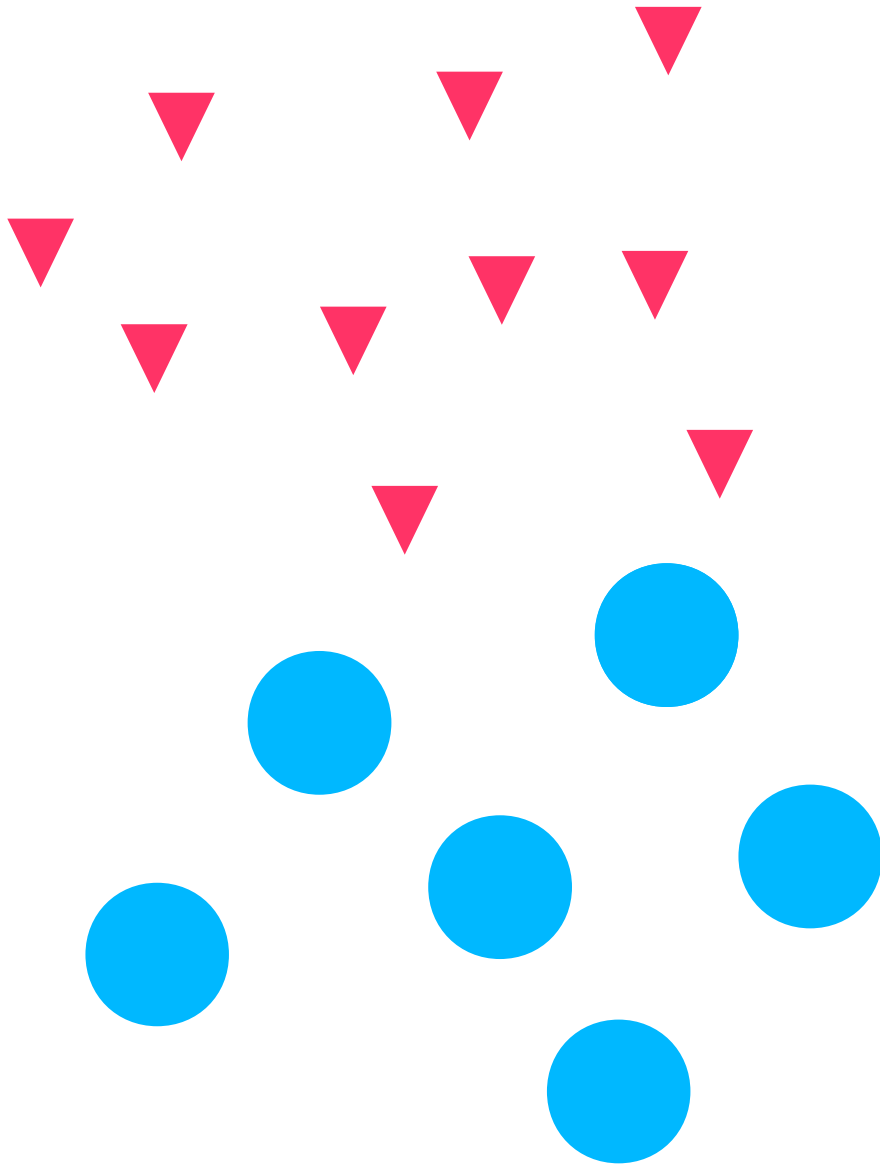


Calcium dose-response on 0.1 μM Calmodulin

fraction of
occupied
binding sites
($\bar{Y}$)

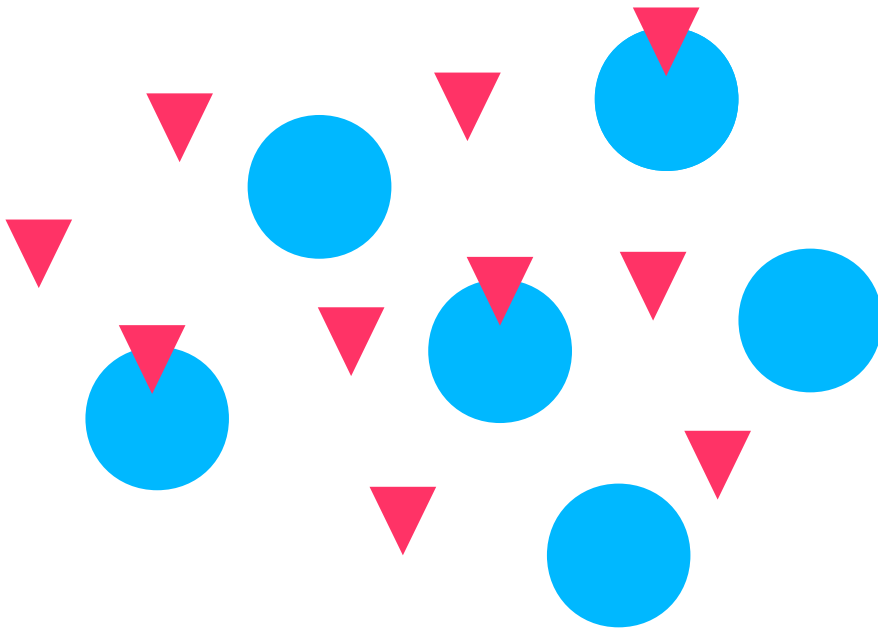
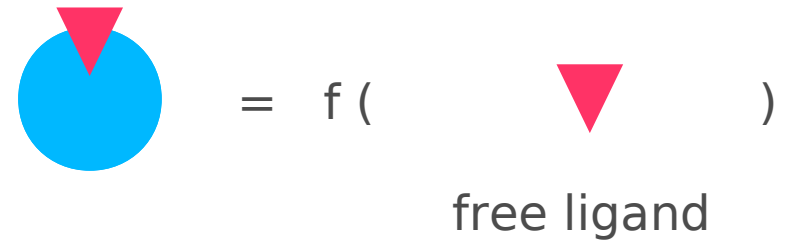


What is ligand depletion?



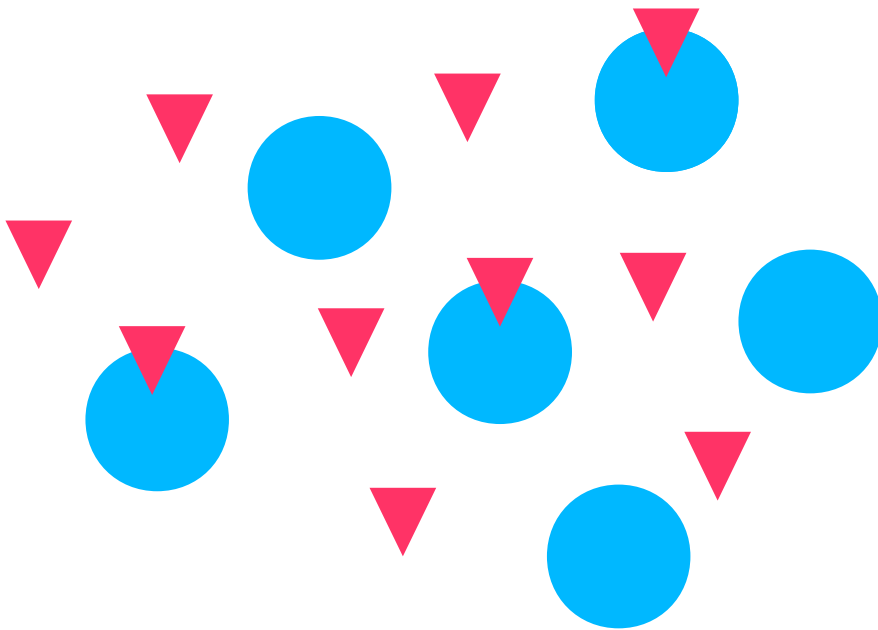
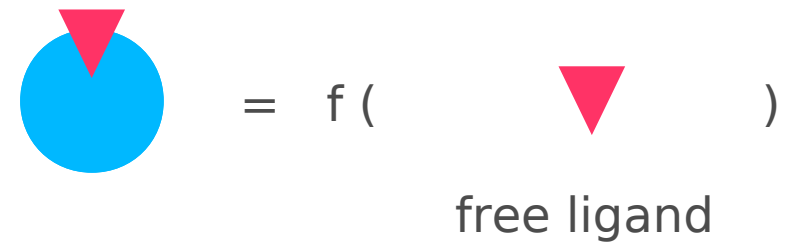
What is ligand depletion?

Chemistry (mass-action law)

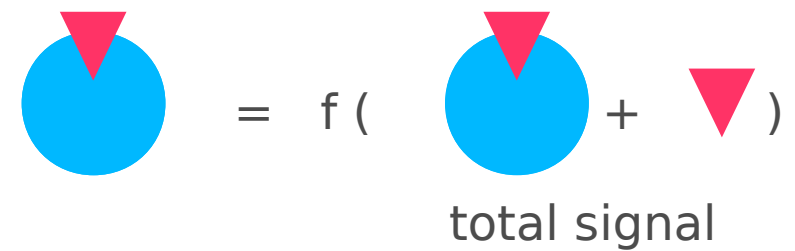


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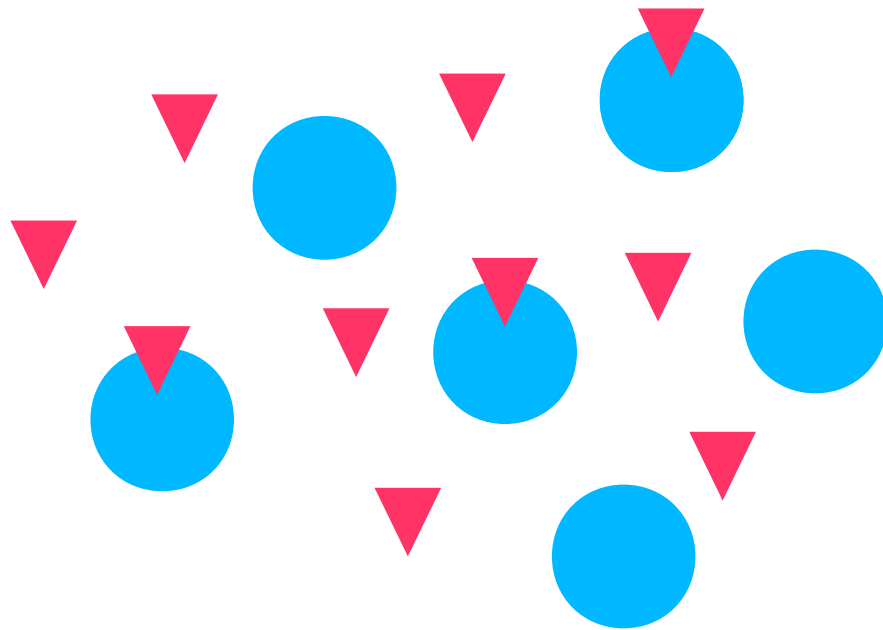
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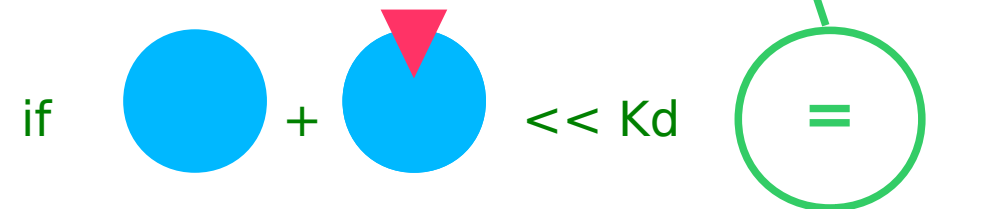
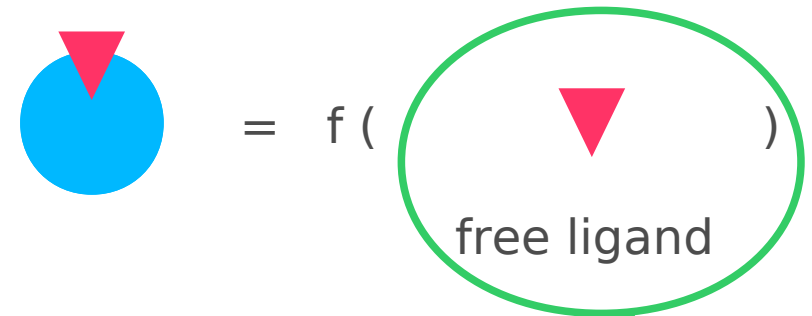
Cellular signalling



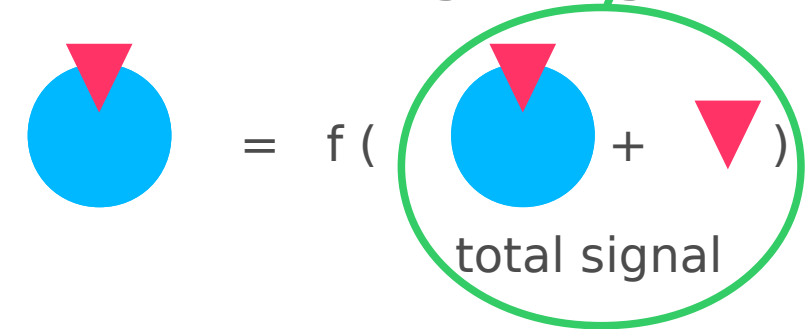
What is ligand depletion?



Chemistry (mass-action law)

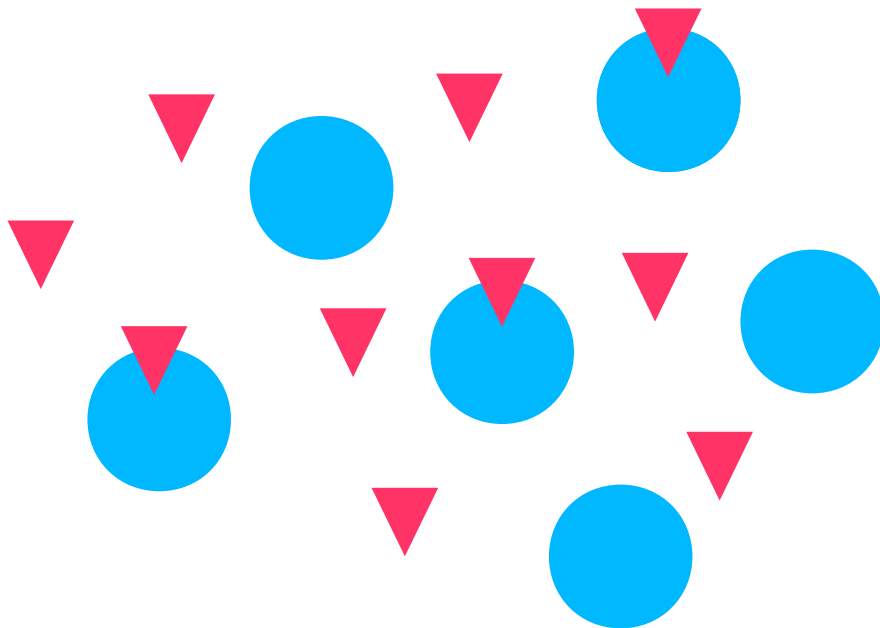


Cellular signalling

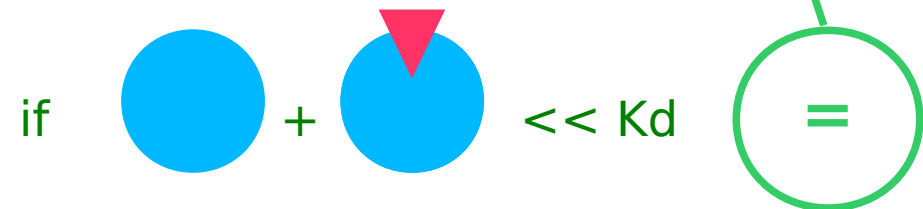
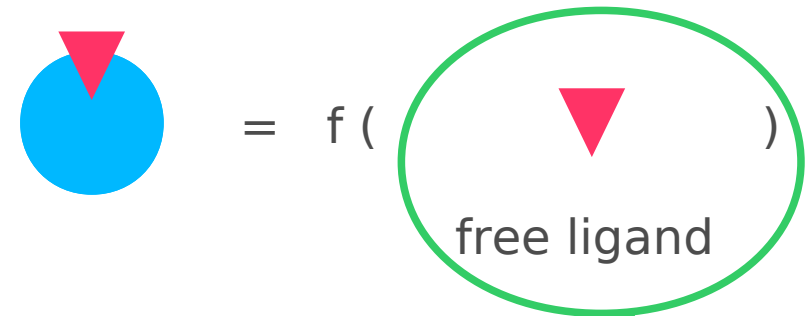


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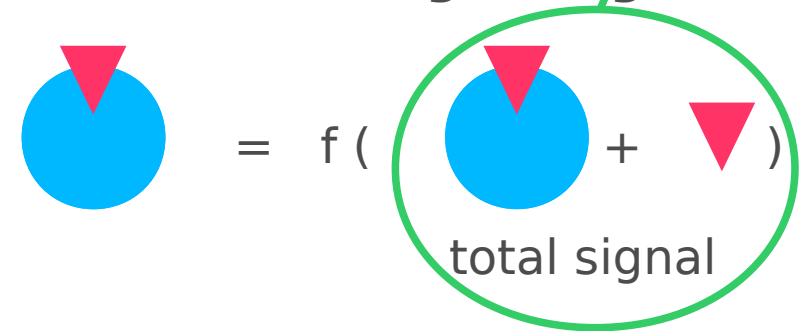
This is generally not the case in signalling:
Concentrations of sensors are in micromolar range, as are the dissociation constants.



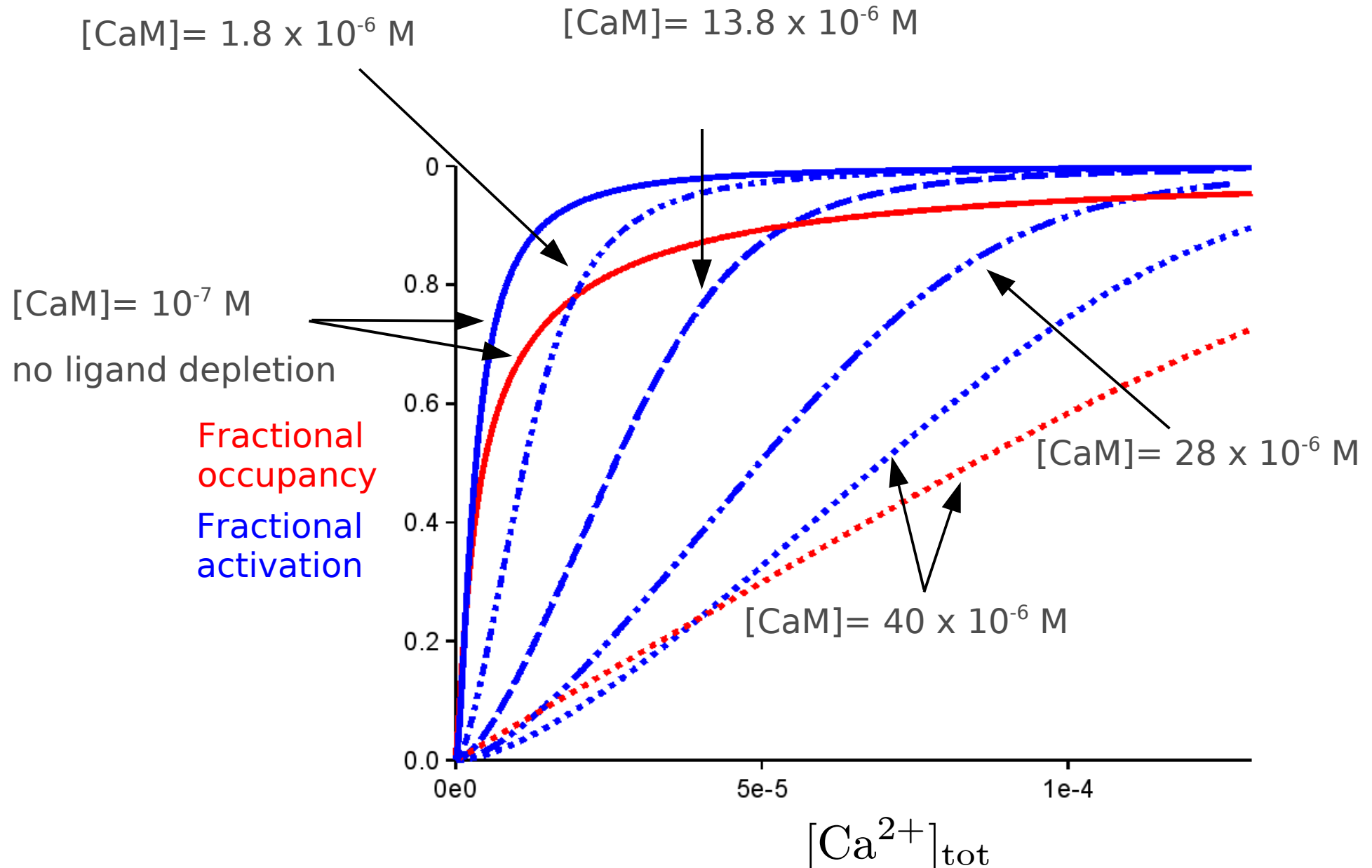
Chemistry (mass-action law)



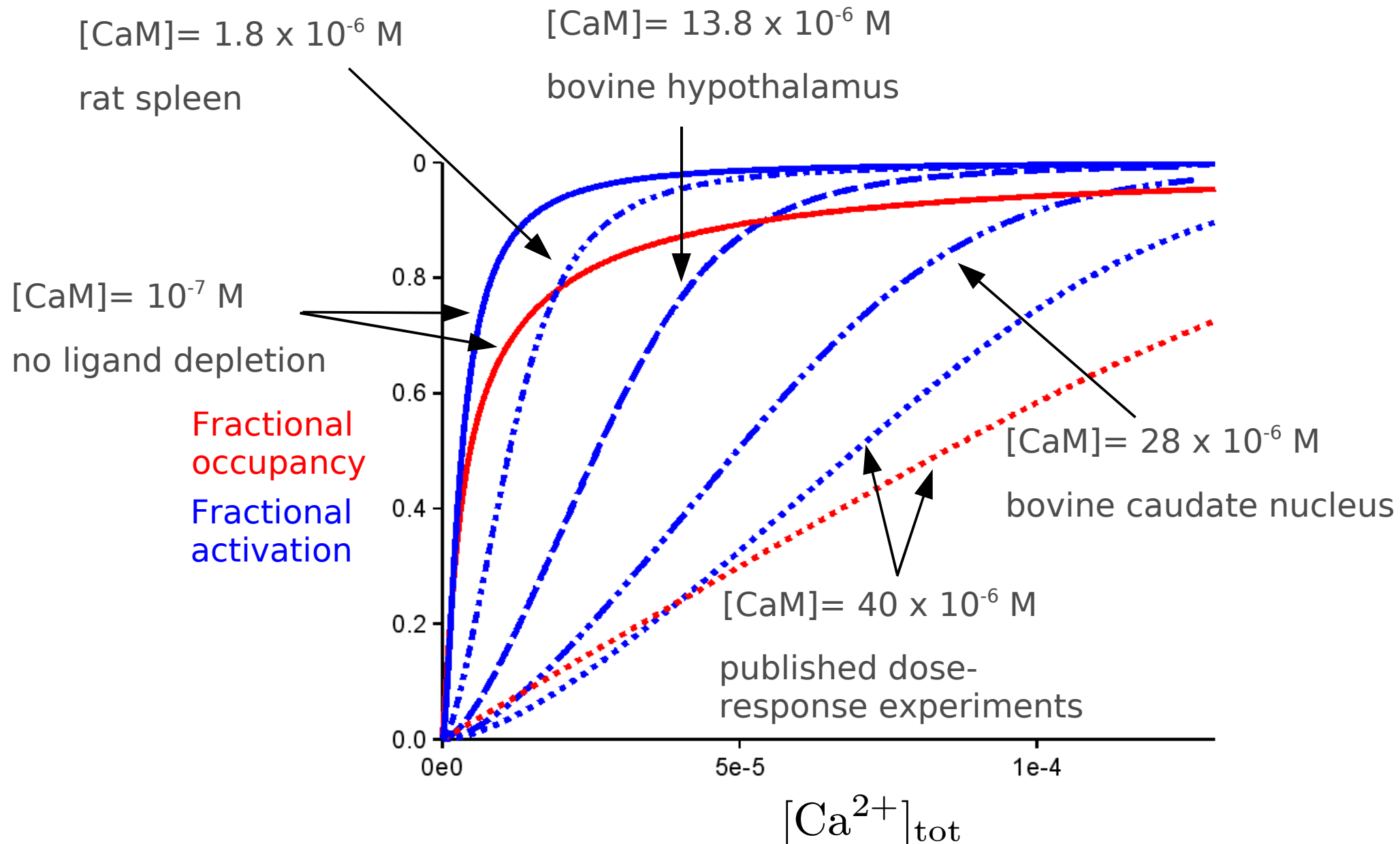
Cellular signalling



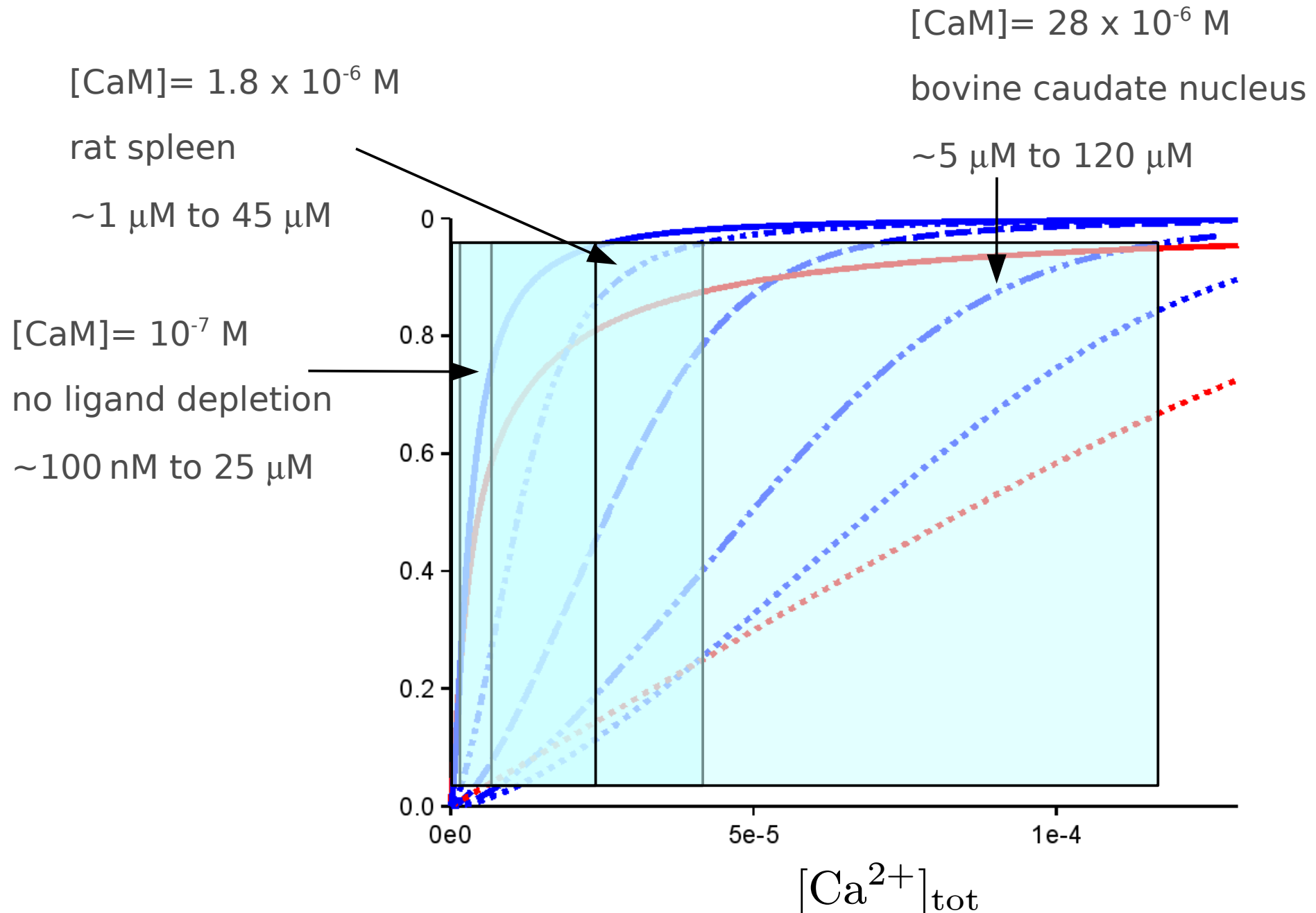
Dose-response depends on Calmodulin concentration



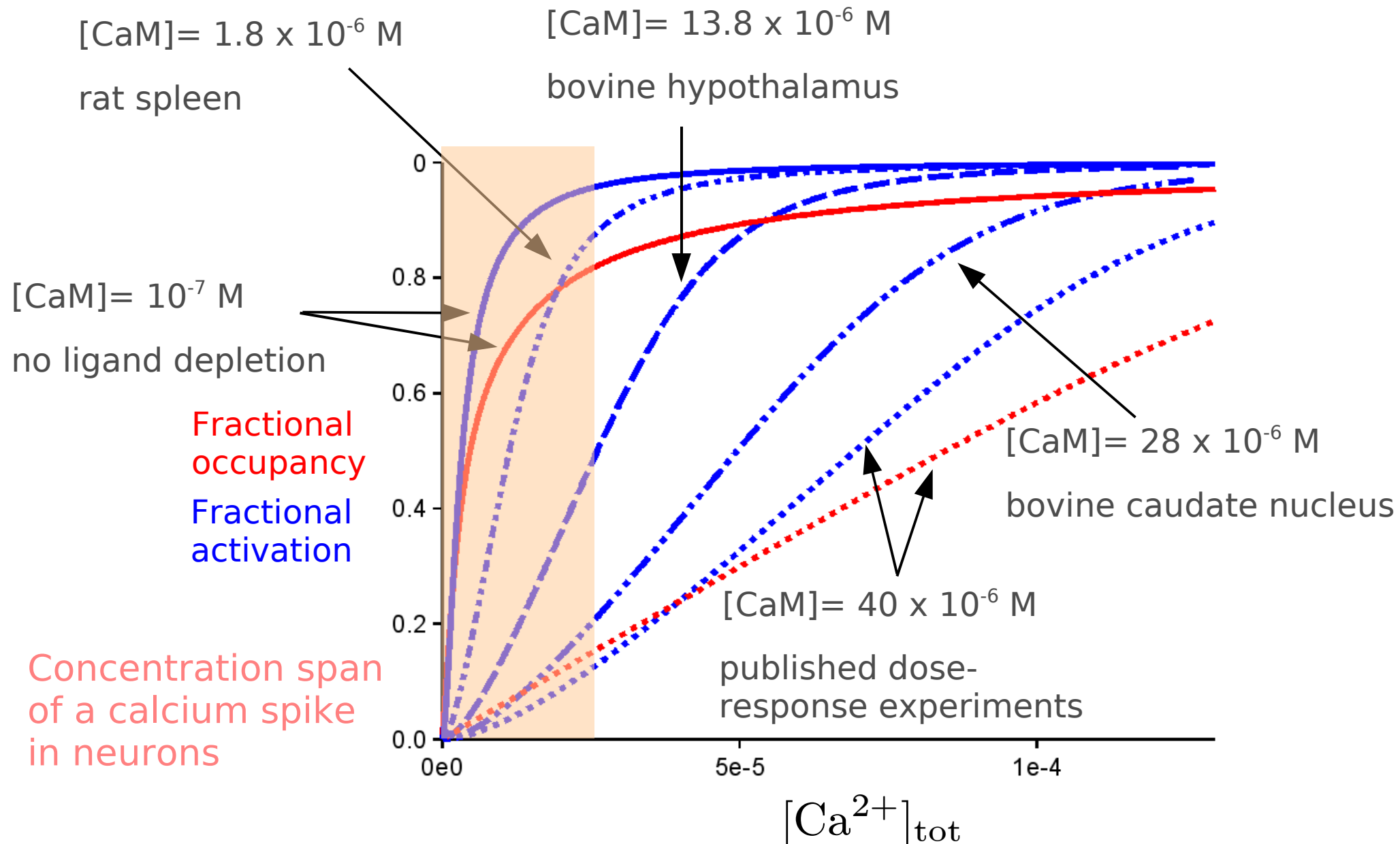
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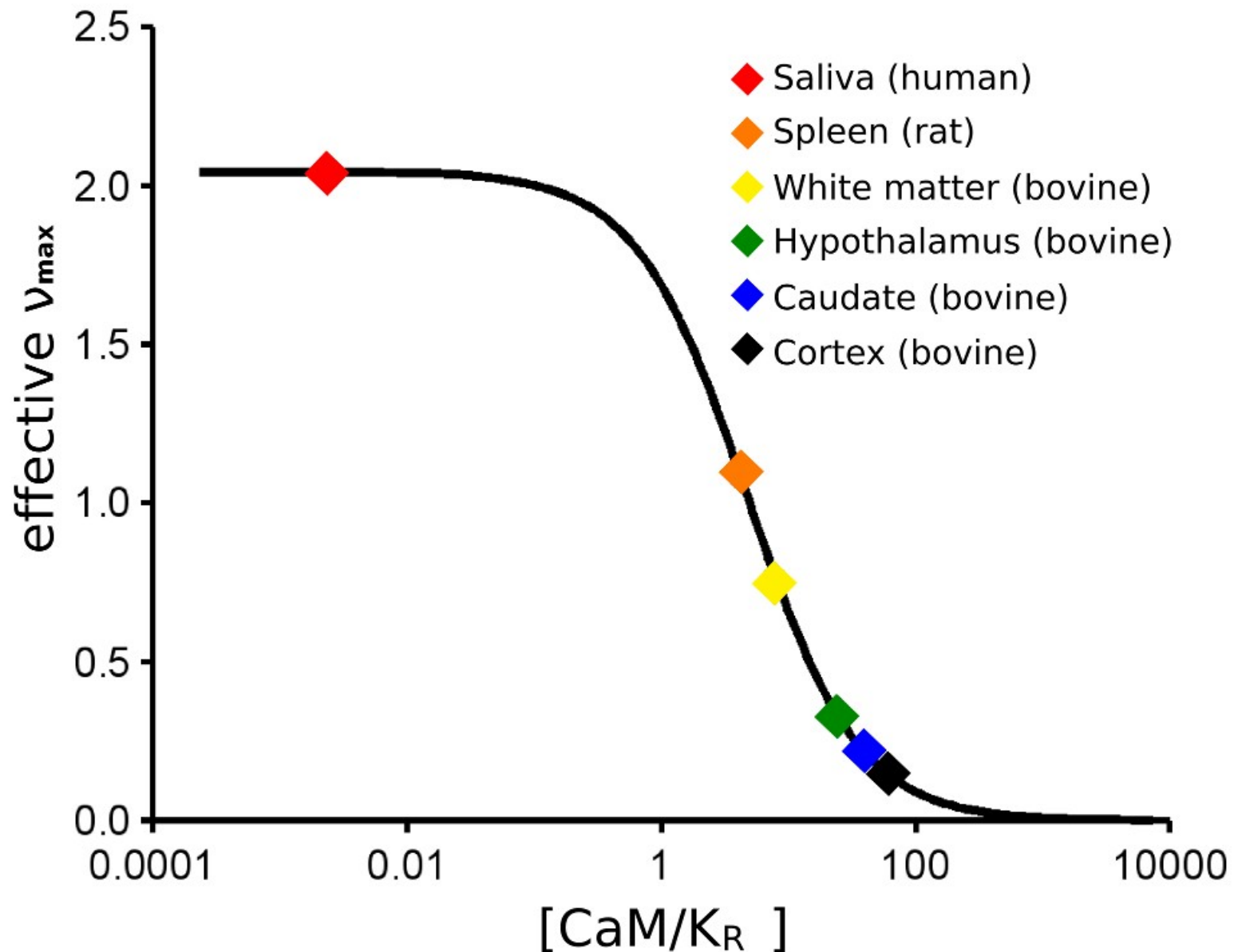
Ligand-depletion modifies sensitivity



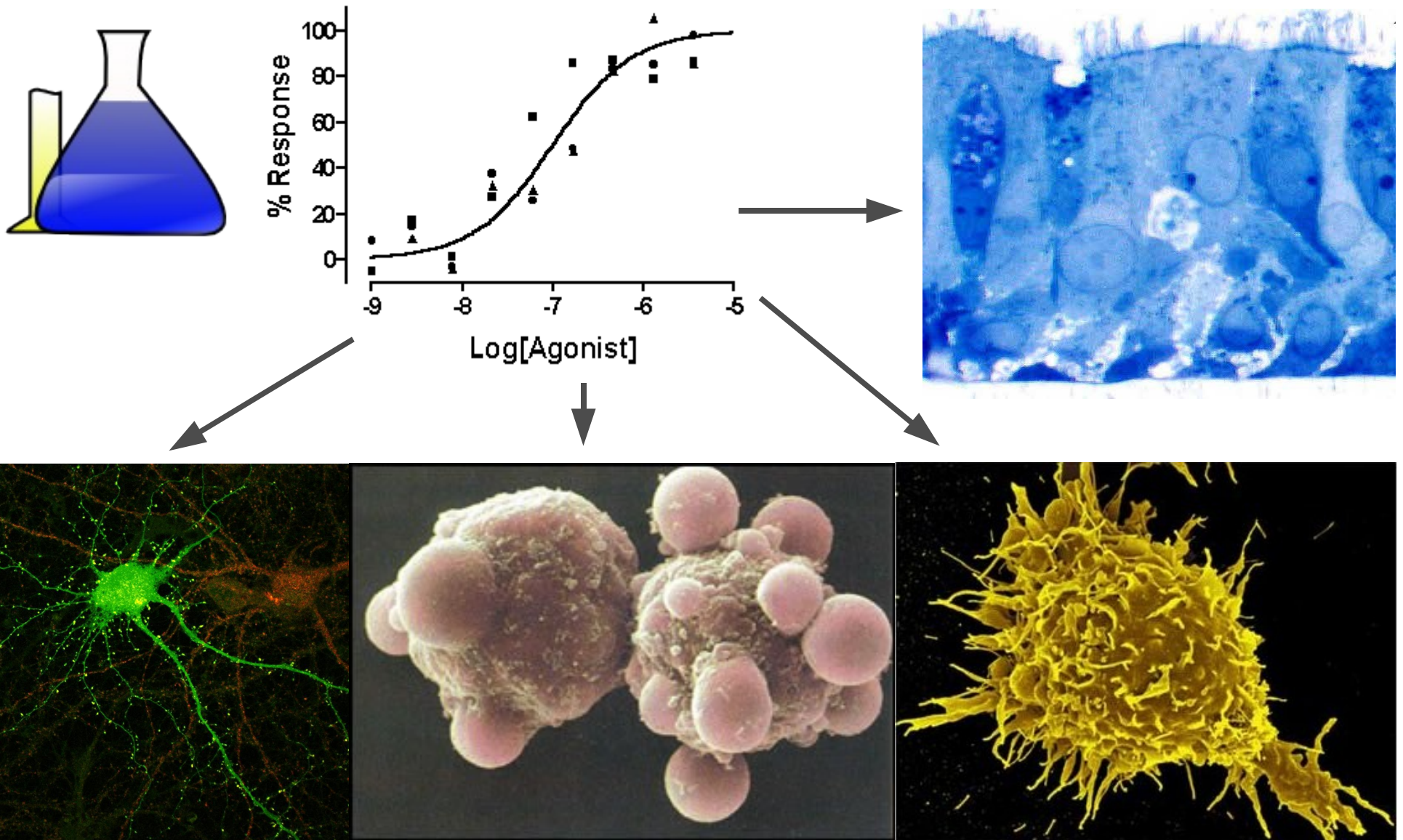
But we cannot build a large $[Ca^{2+}]$ in neurons ...



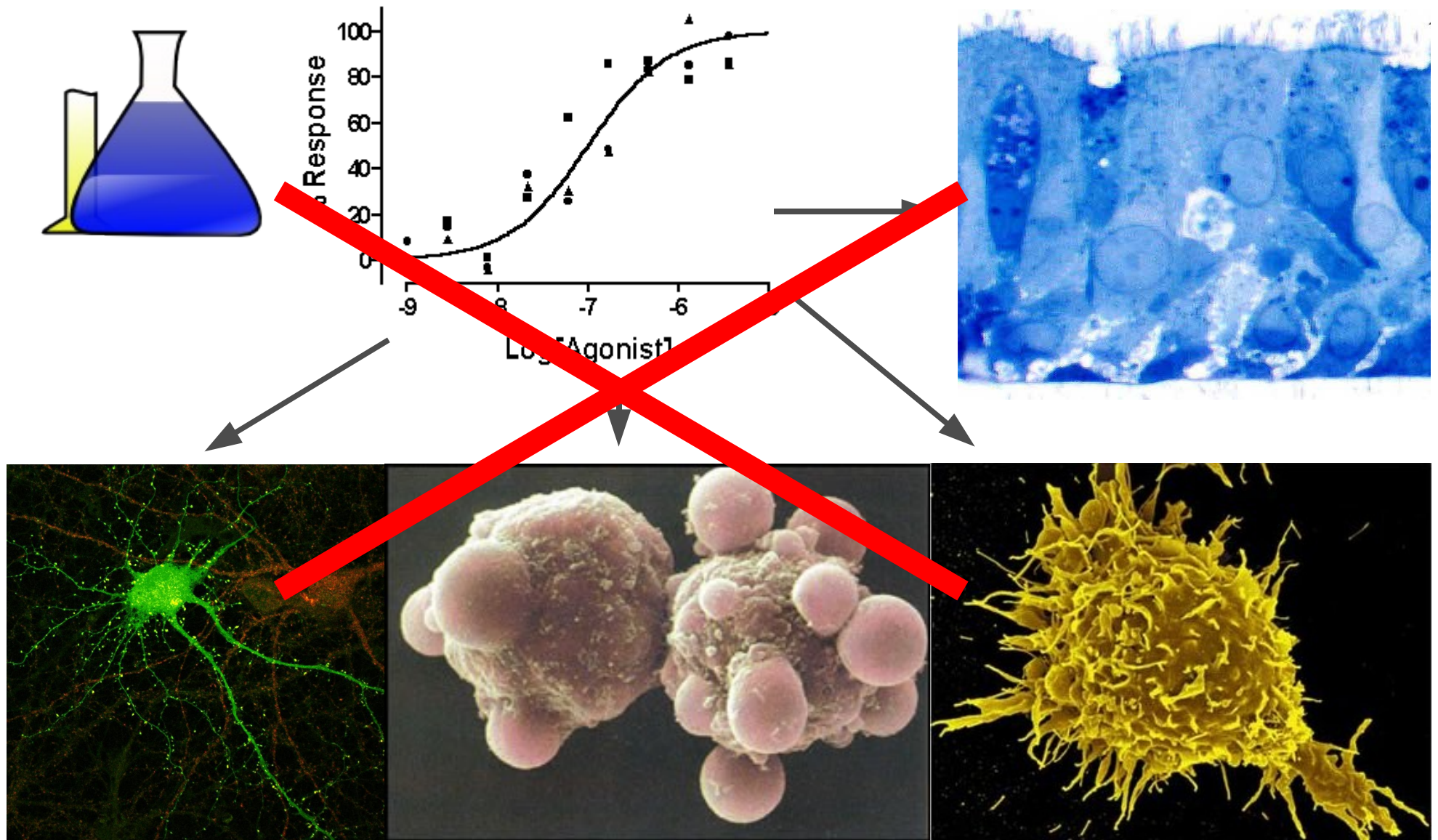
Ligand-depletion decreases effective cooperativity



How general is a dose-response?



A “dose-response” cannot be reused directly!



- Dose-responses are the basic characterisations of “systems”, but also at the core of pharmacological treatments. Here we show that:
 - A “dose-response” cannot be reused directly in models of signalling systems. Instead one needs to build “mechanistic” models and run parameter-fitting approaches.
 - Ligand depletion decreases the effective cooperativity of transducers *in situ*
 - Ligand depletion increases the dynamic range
- Modifying the concentration of the sensor may be a powerful way to quickly adapt to a new environment, and switch from a measurement mode to a detection mode.

Acknowledgements

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Melanie Stefan



Stuart Edelstsein

CompNeur group



Melanie Stefan, postdoc



Stuart Edelstein, visitor



Michele Mattioni, predoc



Lu Li, predoc



Benedetta Baldi, predoc



Christine Hoyer, predoc

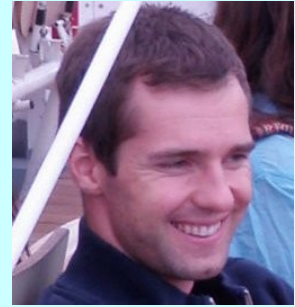


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Sarah Keating
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Chen Li, developer



Nick Juty, curator



Nicolas Rodriguez
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Marine Dumousseau
developer