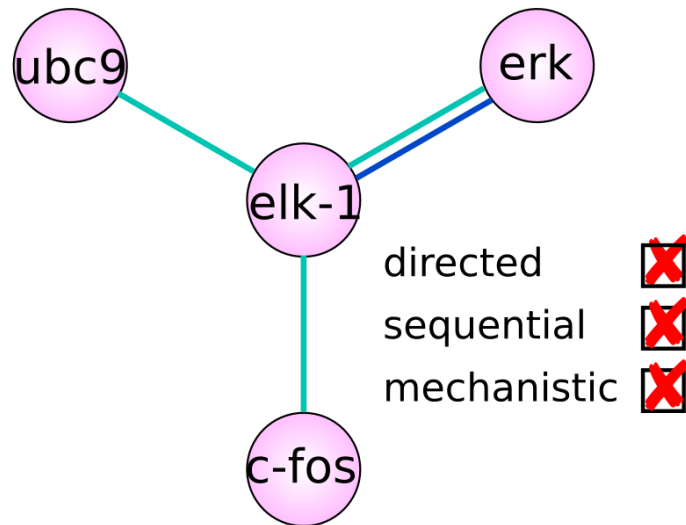


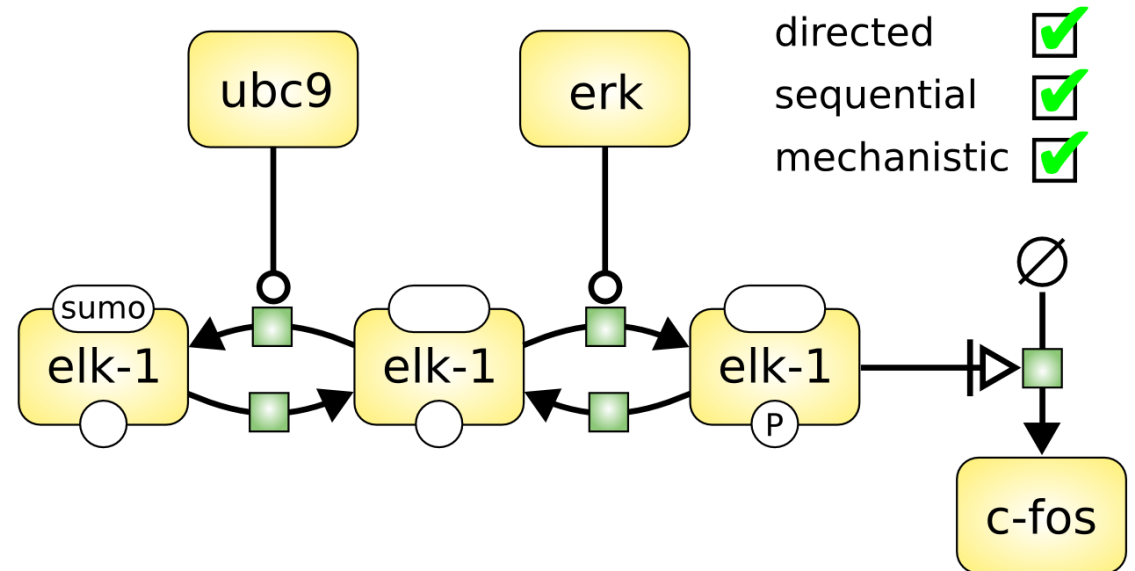
Modelling in Systems Biology, A few challenges

The four views of molecular systems biology

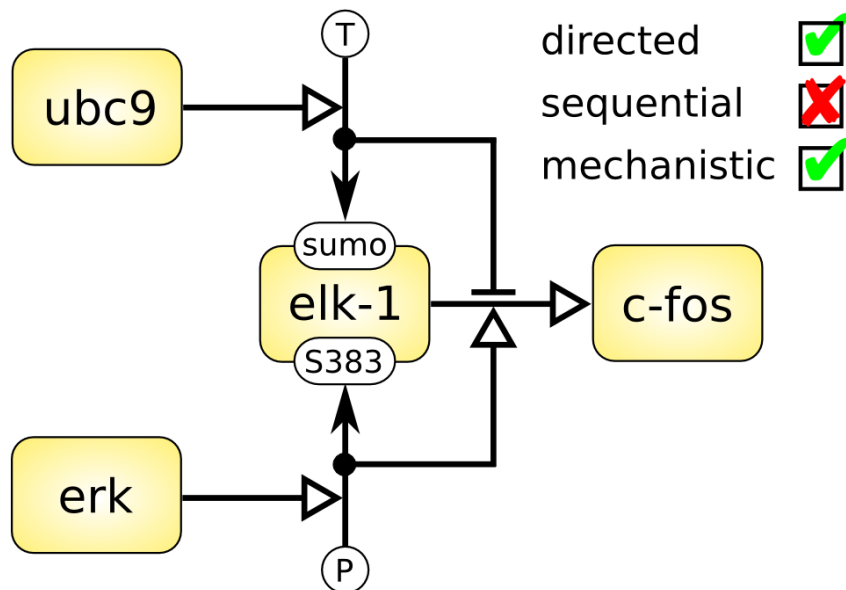
a interaction network



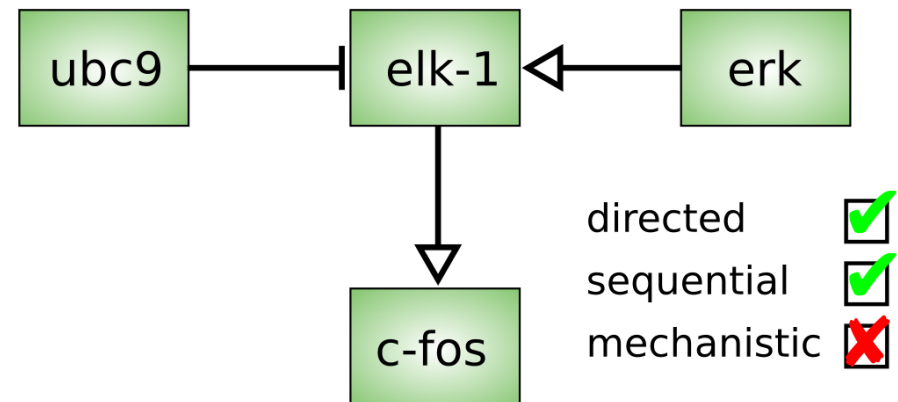
c process descriptions



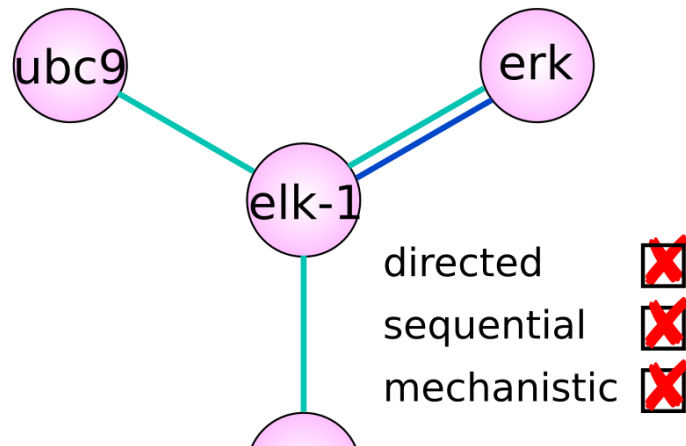
b entity relationships



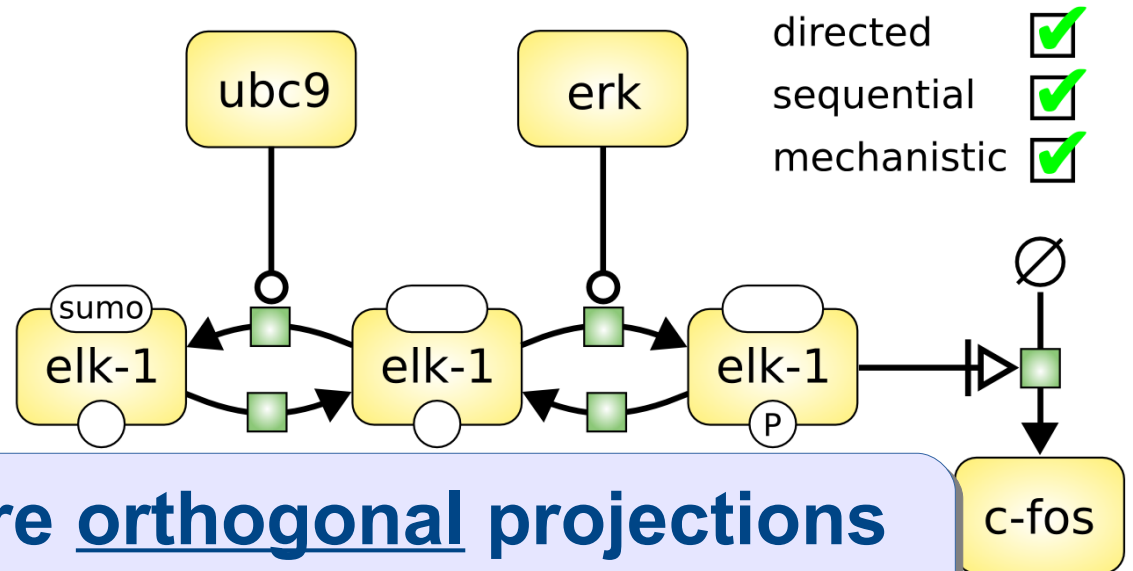
d activity flows



a interaction network

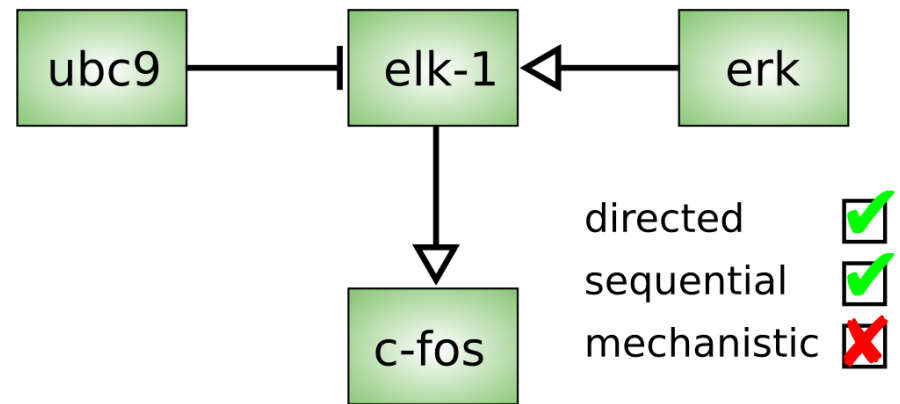
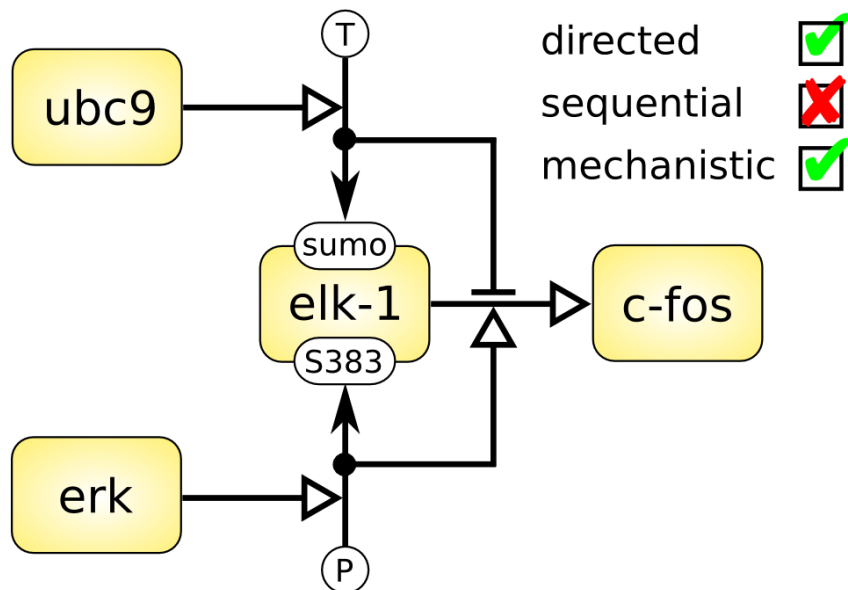


c process descriptions



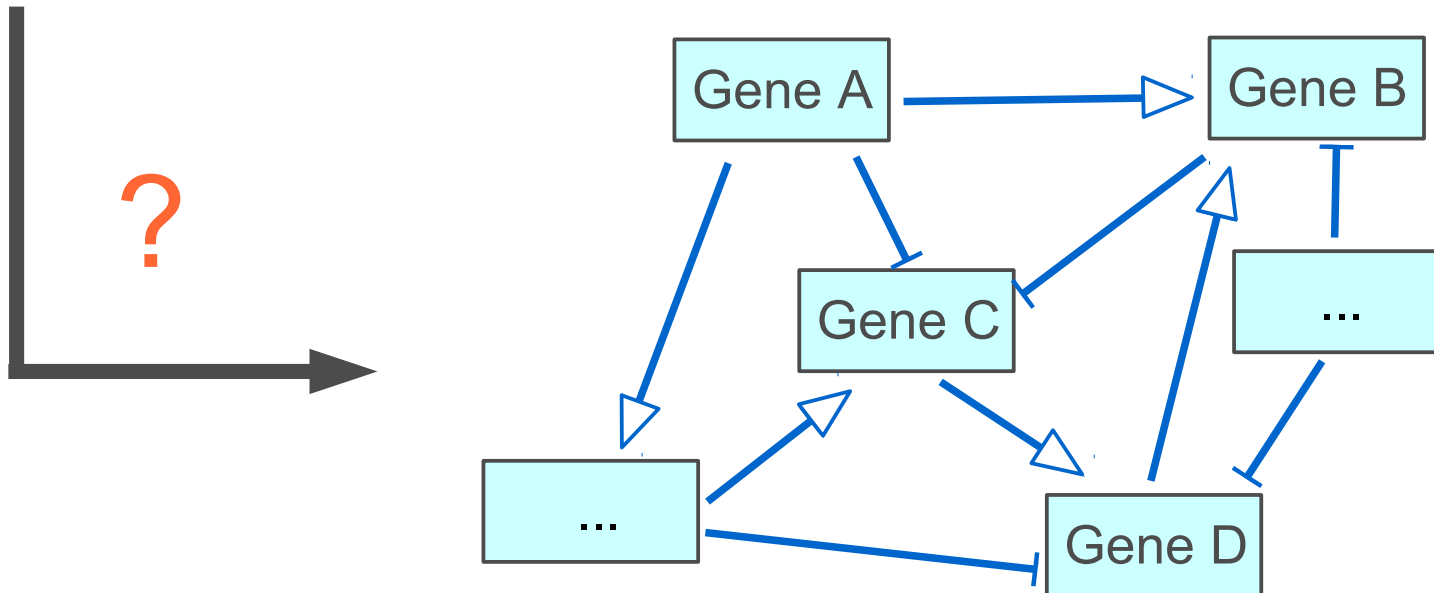
The four views are orthogonal projections of the underlying biological phenomena

b ent

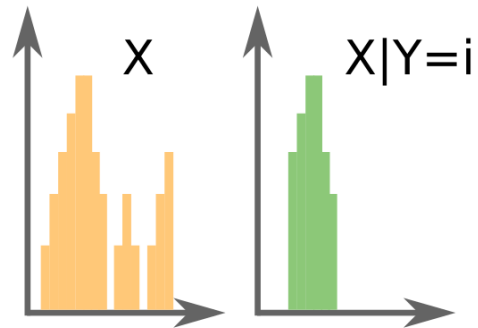


Reverse engineering is hard ...

	Gene A	Gene B	Gene C	Gene D	...
Phenotype X	✓	✗	✓	✗	
Phenotype Y	✓	✗	✗	✓	
Phenotype Z	✗	✓	✓	✗	
...					

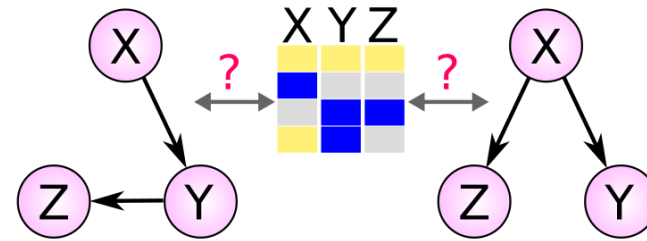


b Information theoretic



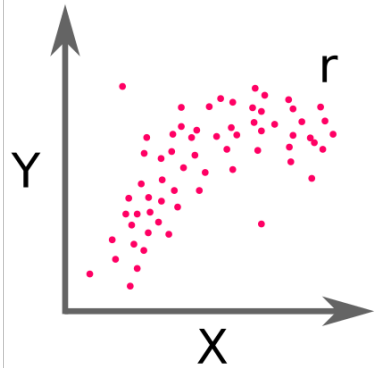
$$I(X, Y) = H(X) - H(X|Y)$$

c Bayesian inference



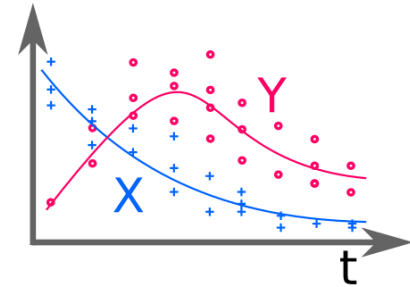
$$P(X|Y) = \frac{P(Y|X)P(X)}{P(Y)}$$

a Correlation

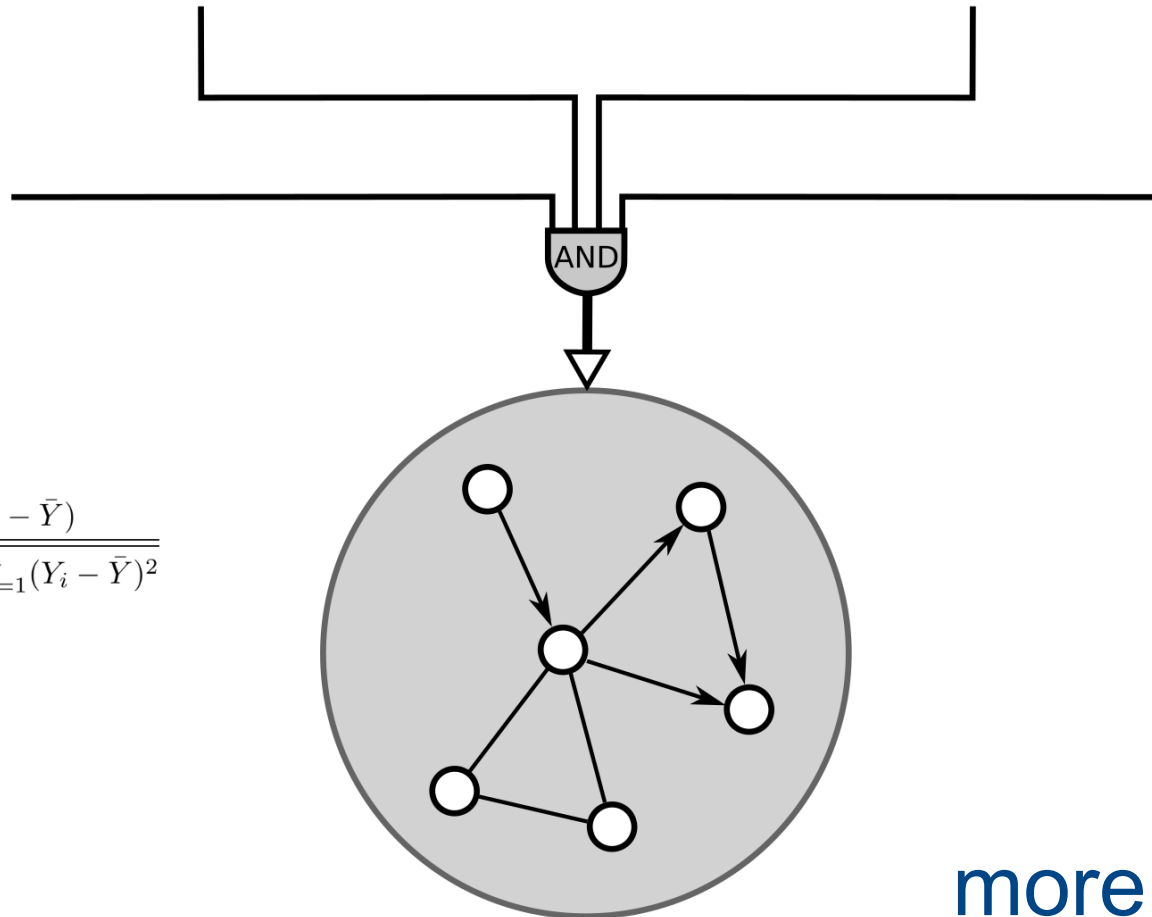


$$r = \frac{\sum_{i=1}^n (X_i - \bar{X})(Y_i - \bar{Y})}{\sqrt{\sum_{i=1}^n (X_i - \bar{X})^2} \sqrt{\sum_{i=1}^n (Y_i - \bar{Y})^2}}$$

c Differential equation



$$\frac{dx_i}{dt} = \sum_{j=1}^n a_{i,j} x_j$$



more on Monday

Bridging Bioinformatics and Systems Biology

Big historical divide: Origin of scientists (math/bio Vs eng/physics), journals, conferences, scientific societies etc.

Frontiers tend to blur: Network inference from omics data, parametrisation of models, optimisation (e.g. whole-genome metabolic models), precision medicine (perturbations and drugs explained in mechanistic context)



SysMod attempts to bring back systems modelling to ISCB (society) and ISMB (conference)

Improve the conversation between communities of bioinformatics/genomics and modelling

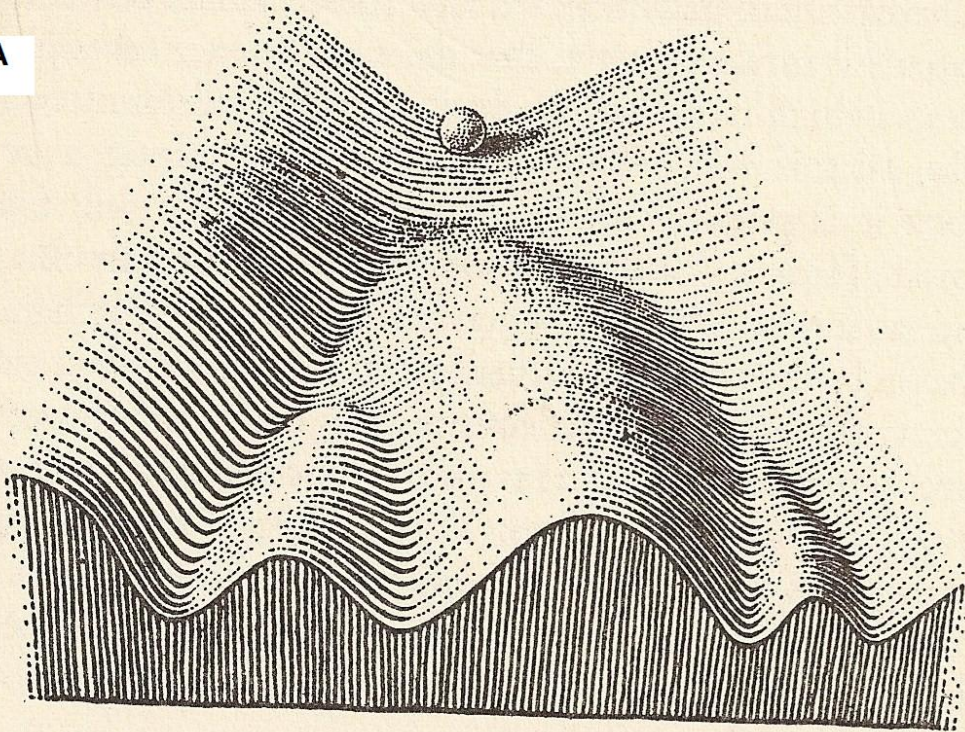
Annual meeting (Prague, 23 July 2017)

Mailing list (sysmod@googlegroups.com)

? (your idea)

<http://sysmod.info>

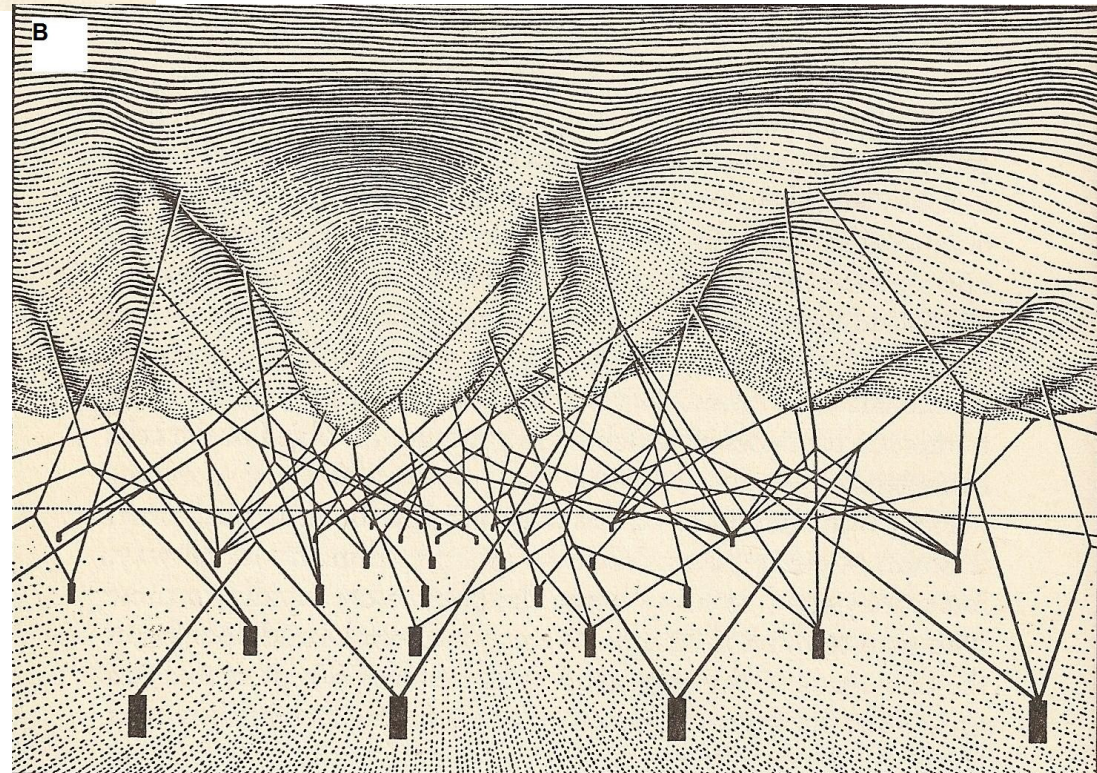
A



Emergent properties and the gene-system-phenotype puzzle

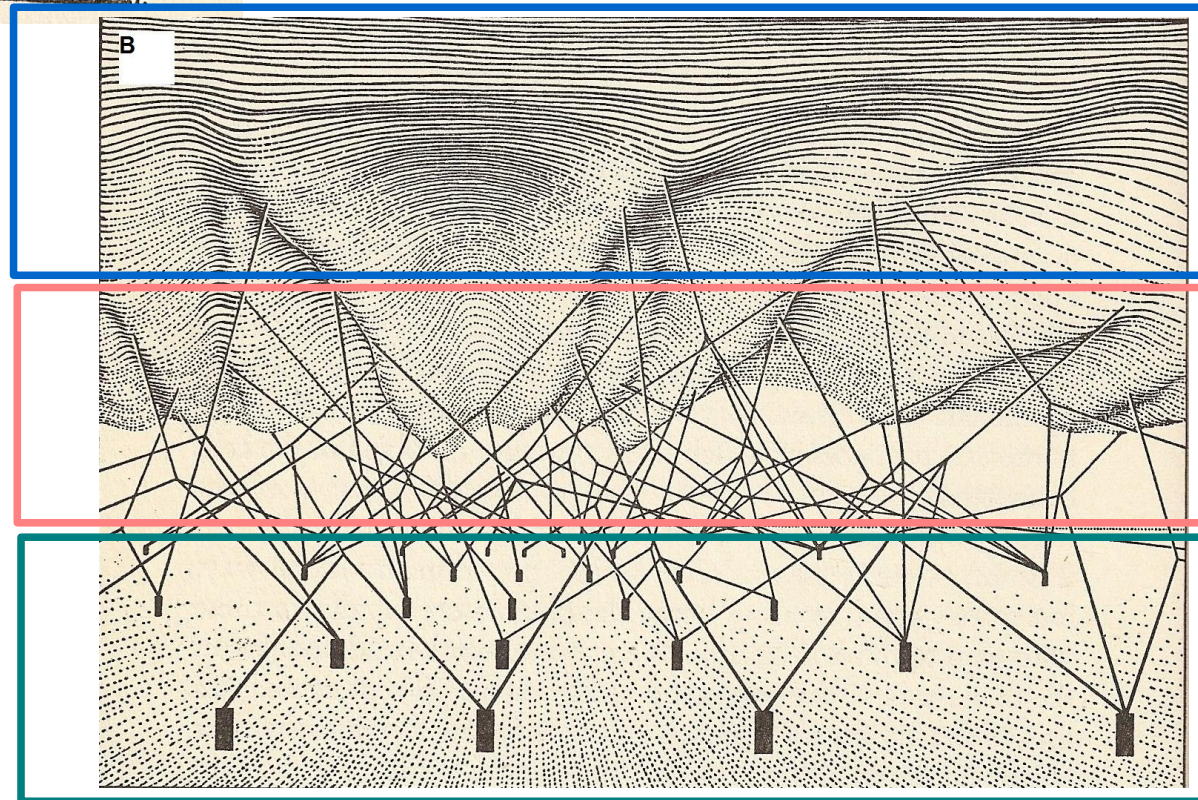
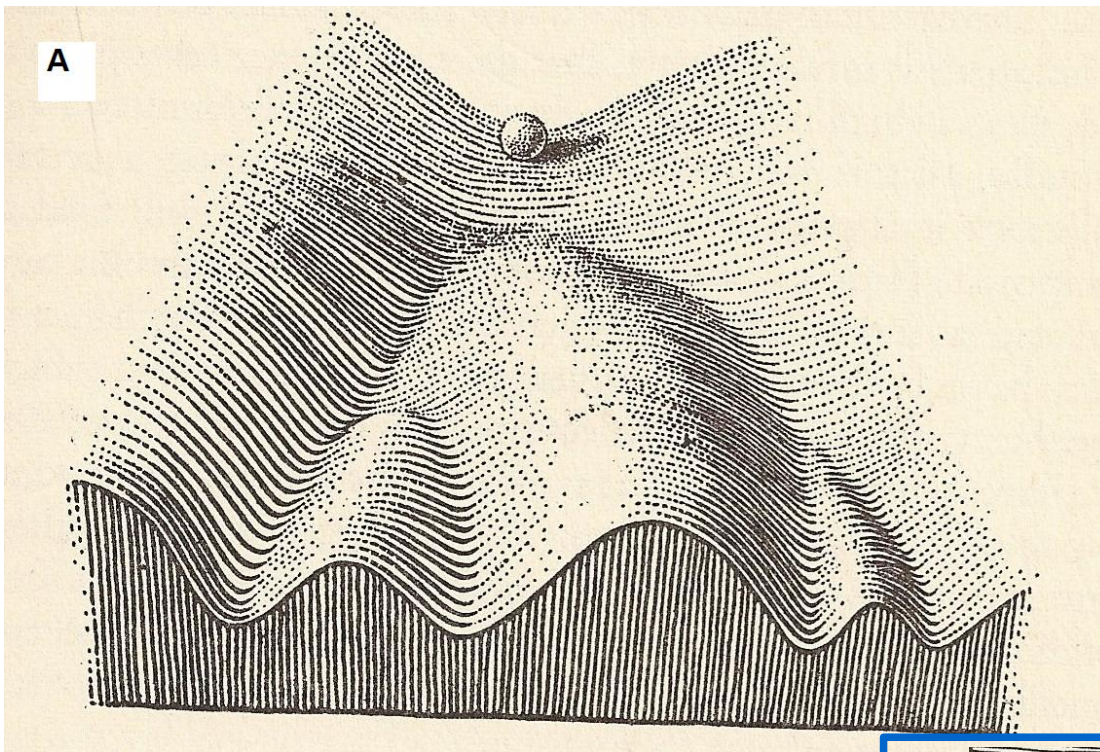
Waddington C.H., Kacser H (1957)
The Strategy of the Genes:
A Discussion of Some Aspects of
Theoretical Biology.
George Allen & Unwin

B



Emergent properties and the gene-system-phenotype puzzle

Waddington C.H., Kacser H (1957)
The Strategy of the Genes:
A Discussion of Some Aspects of
Theoretical Biology.
George Allen & Unwin



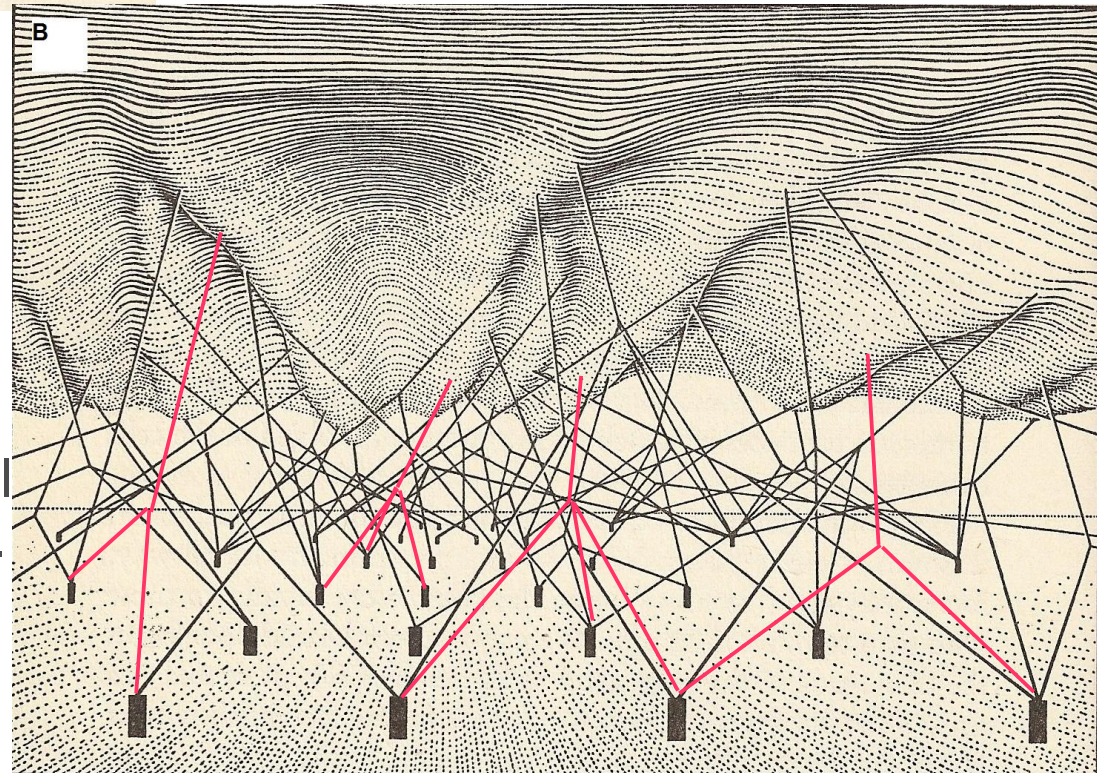
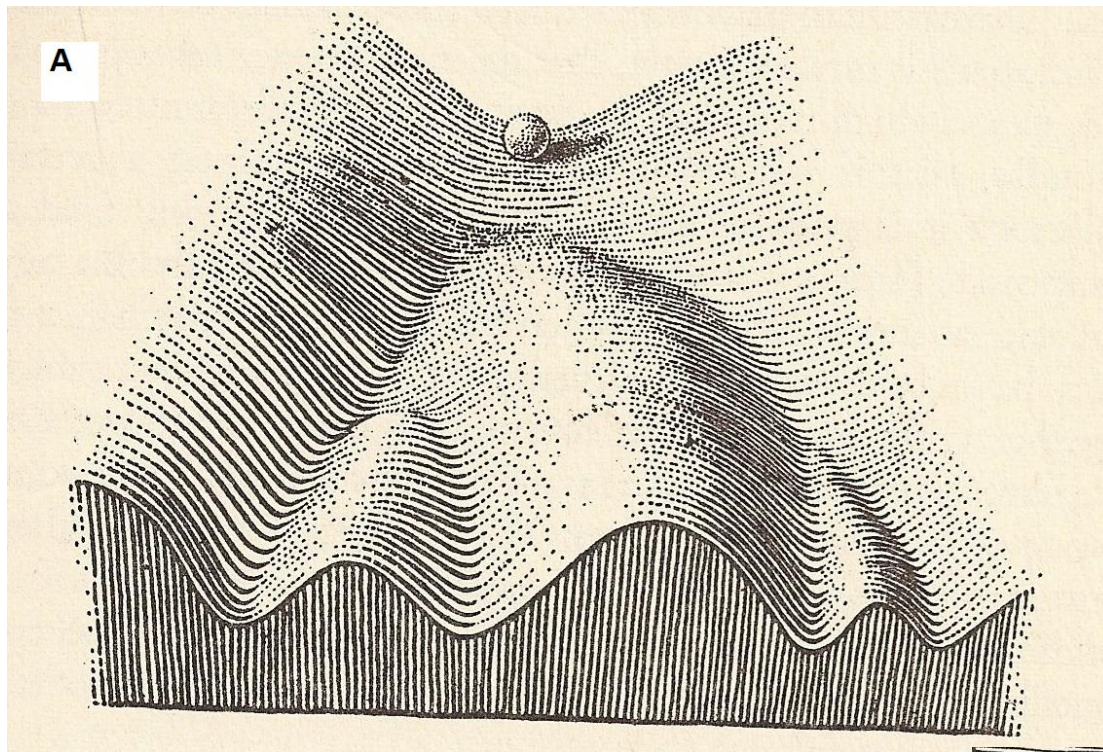
physiology ← phenotype

systems biology ← system

genetics ← genotype

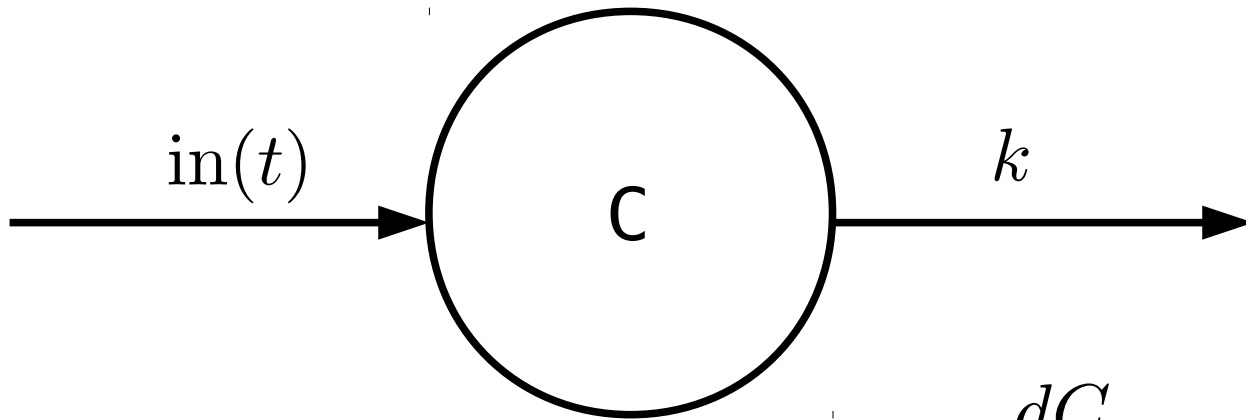
Emergent properties and the gene-system-phenotype puzzle

Waddington C.H., Kacser H (1957)
The Strategy of the Genes:
A Discussion of Some Aspects of
Theoretical Biology.
George Allen & Unwin

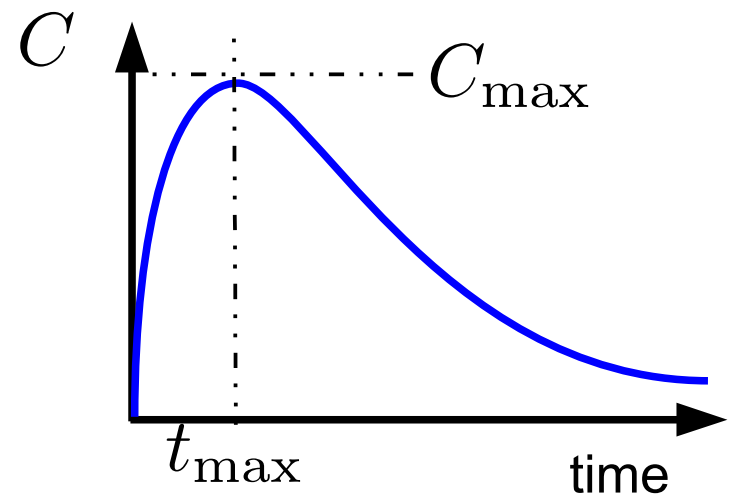
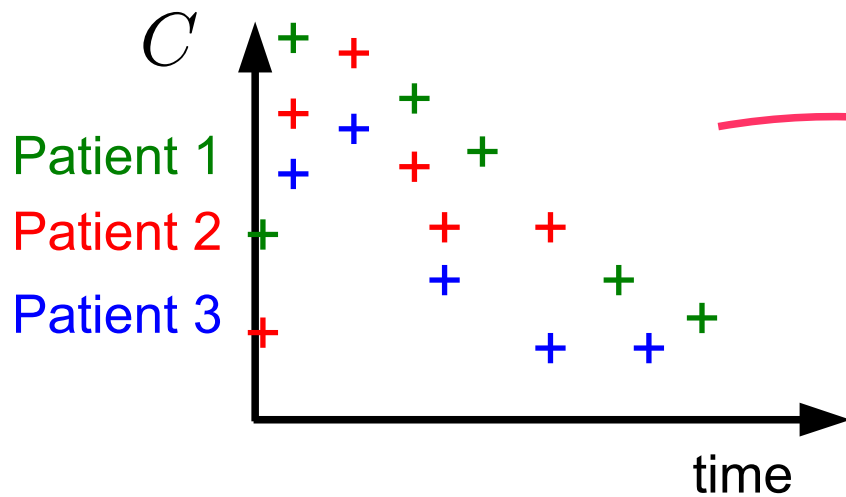


Many networks can theoretically generate the same phenotype, and this happens, in a synchronous (sister cells with same phenotype but different transcript/prote/metabol/omes) and diachronous manner (*omes of a cell changes over time but same phenotype).

Drug discovery modelling: pharmacometrics

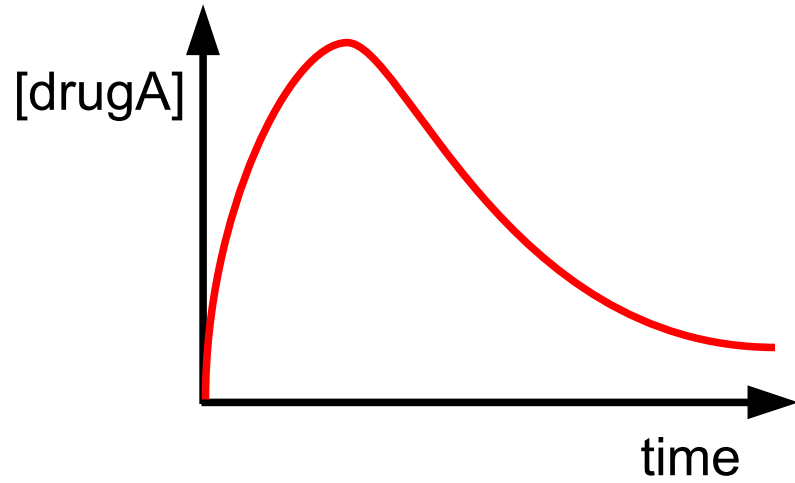


$$\frac{dC}{dt} = \text{in}(t) - k \times C$$

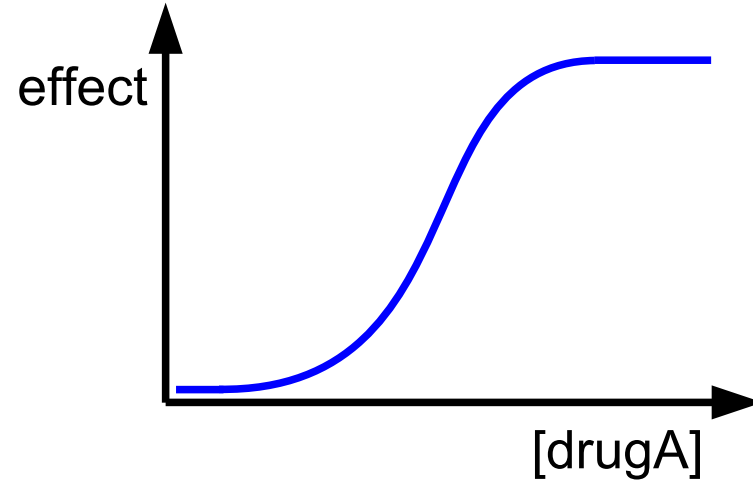


Drug discovery modelling: pharmacometrics

PK

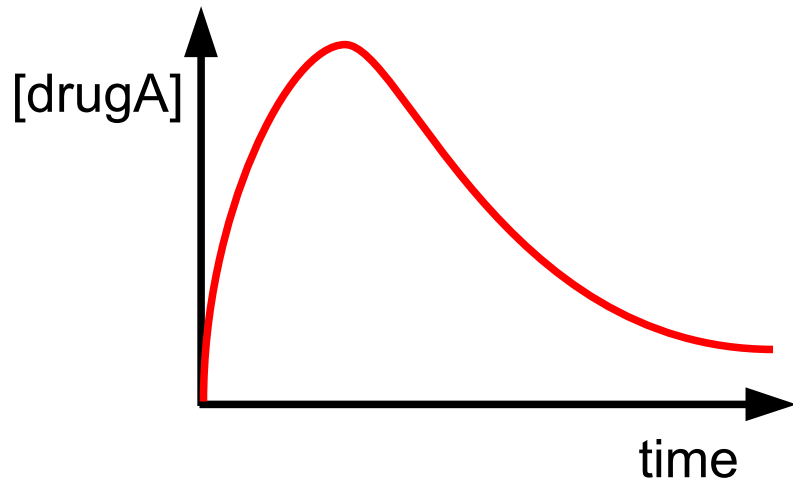


PD

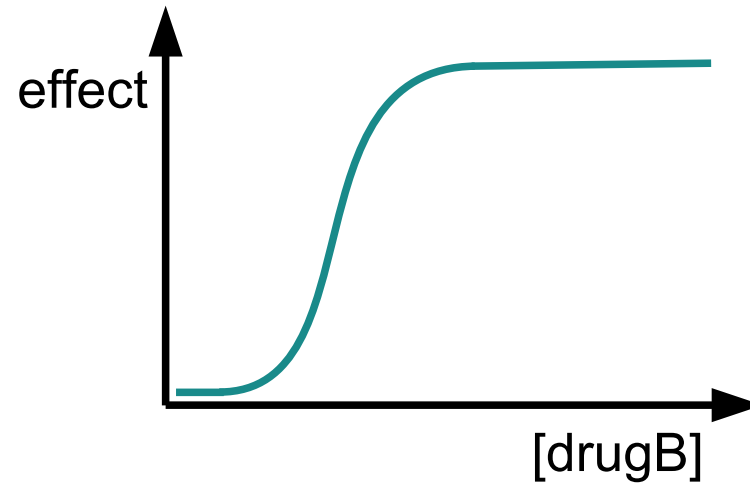
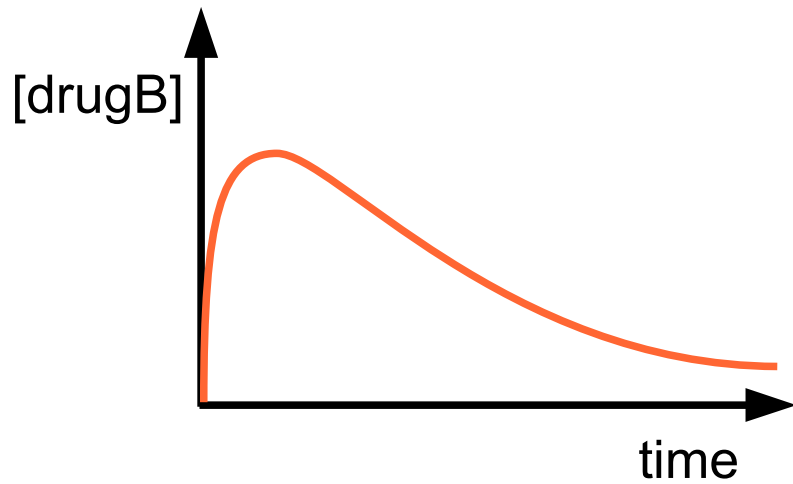
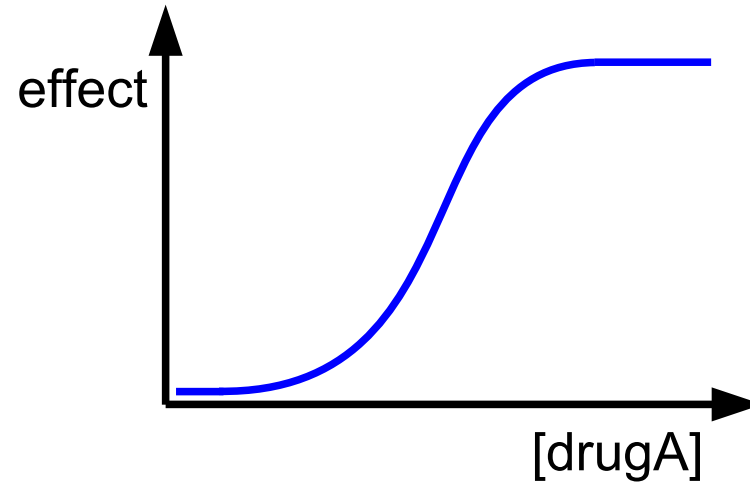


Drug discovery modelling: pharmacometrics

PK

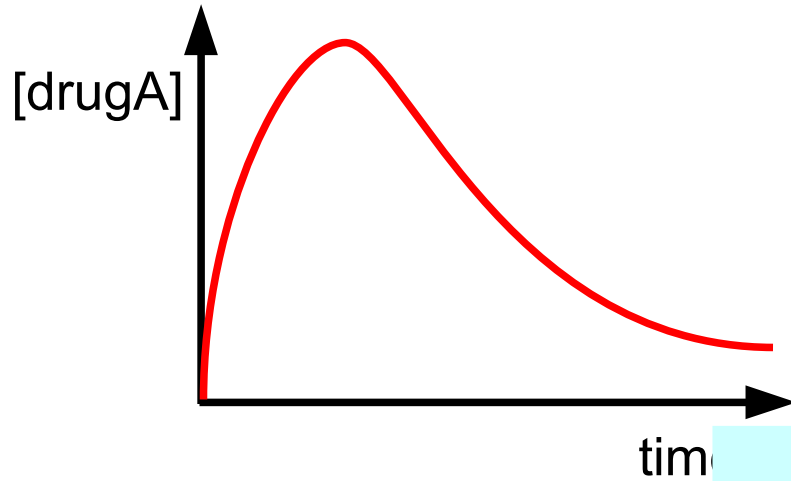


PD

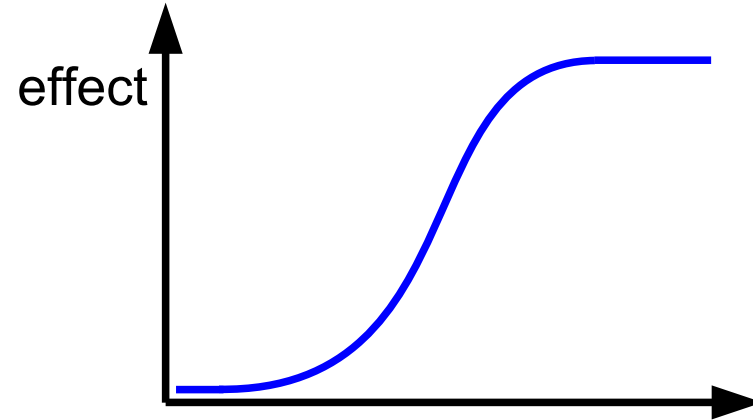


Drug discovery modelling: pharmacometrics

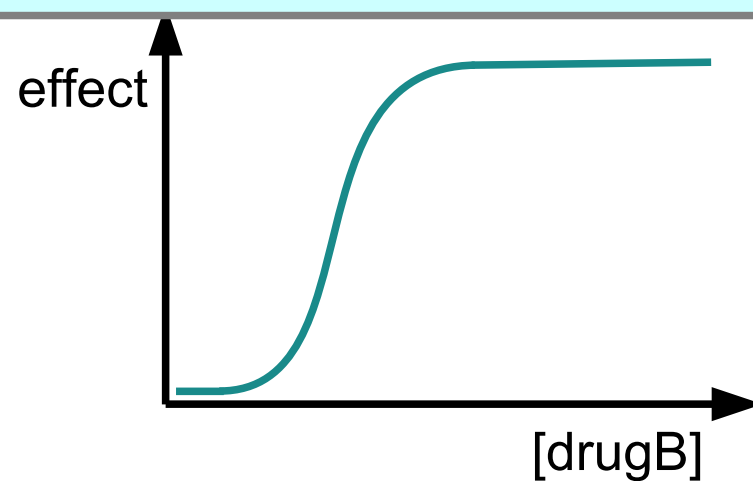
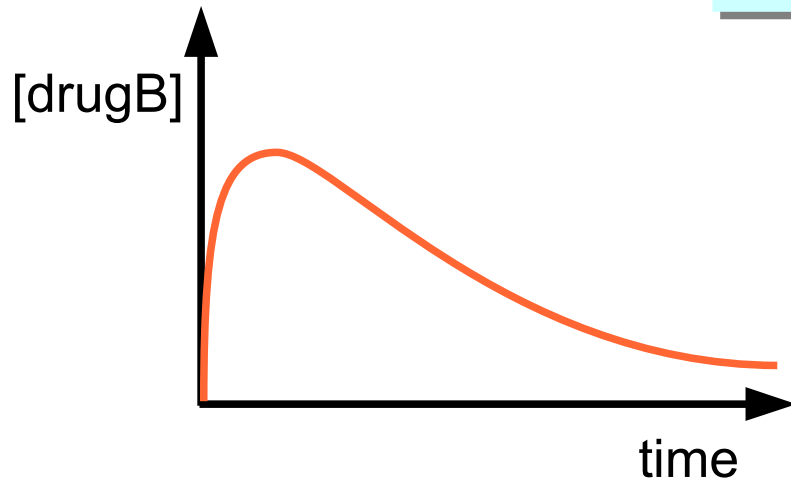
PK



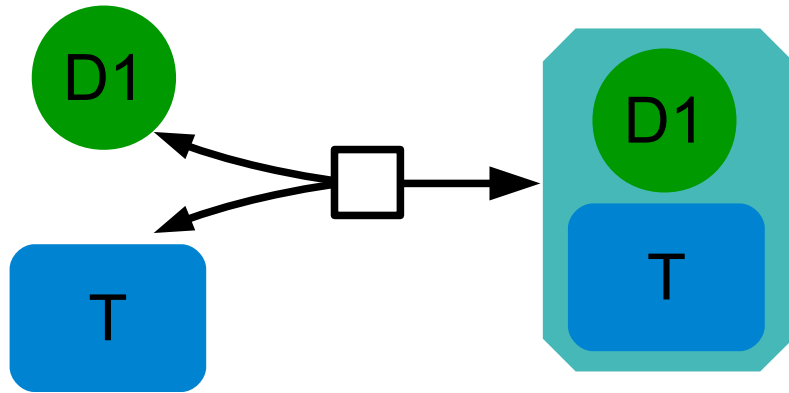
PD



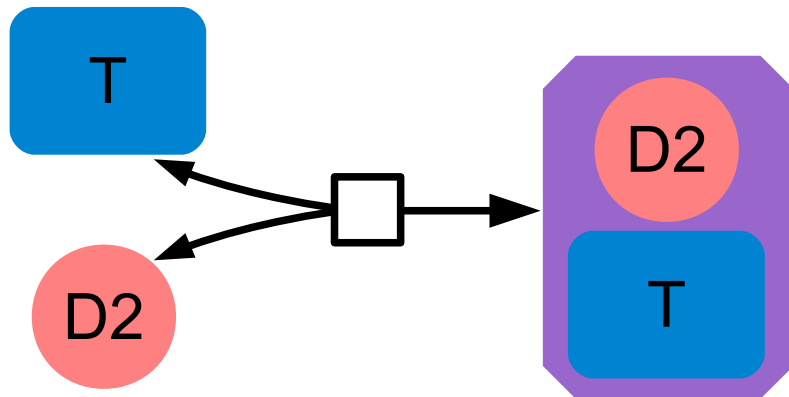
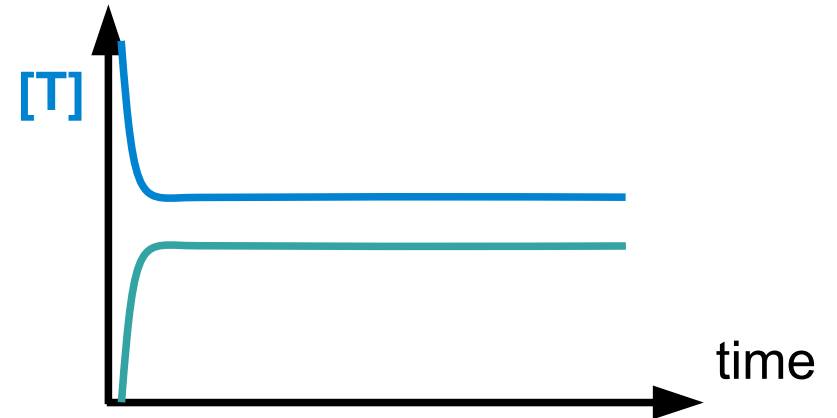
Effect of A+B?



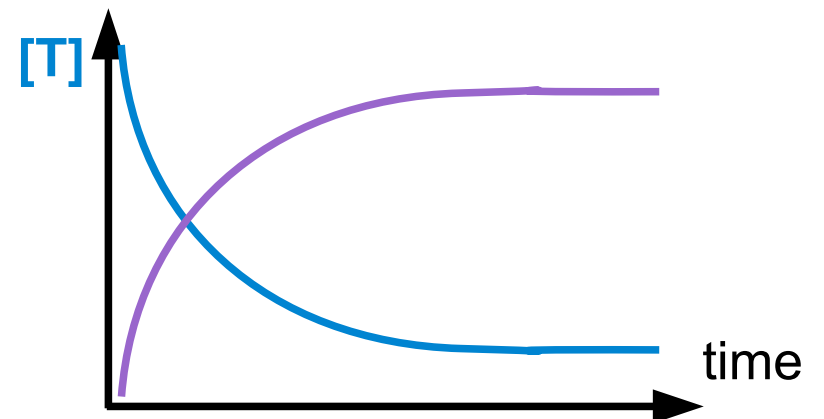
Systems modelling



$$\frac{d[T]}{dt} = -k1 \times [T] \times [D1]$$

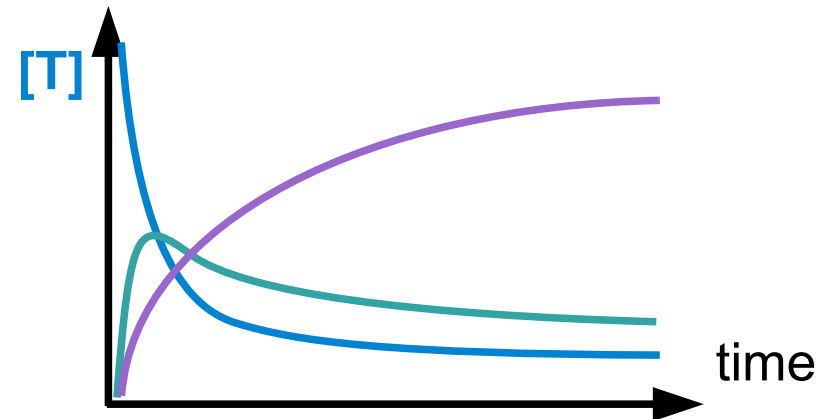
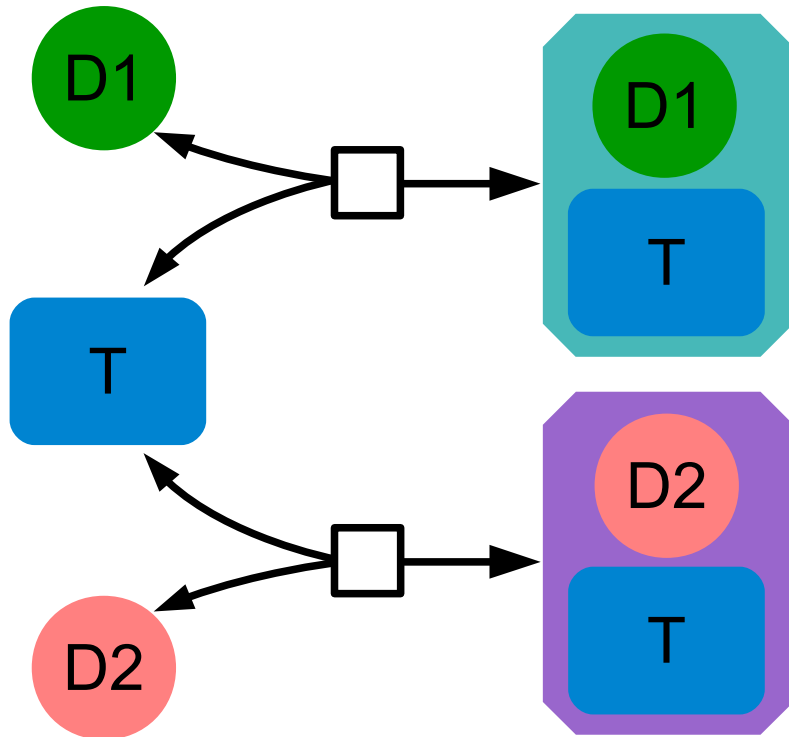


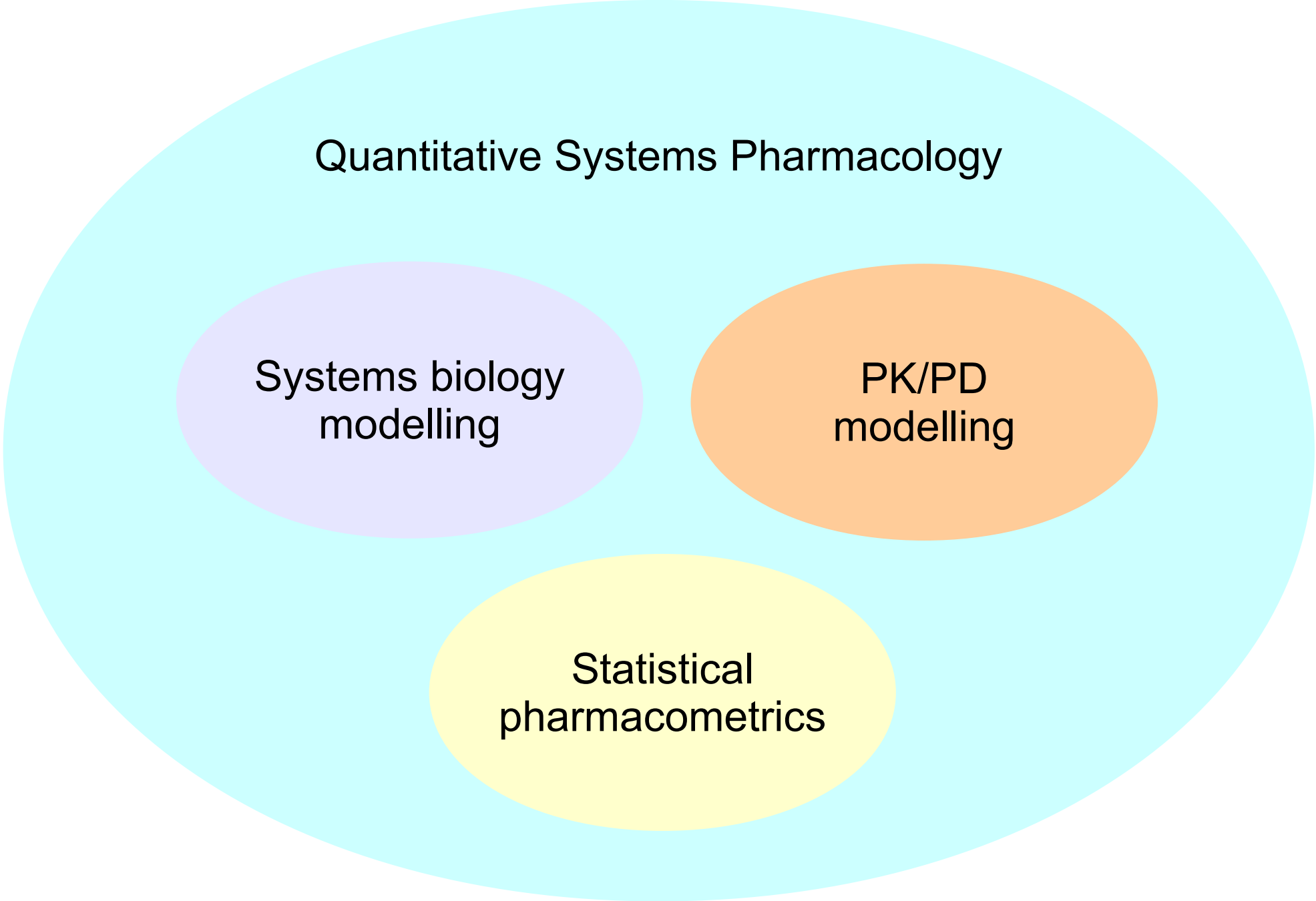
$$\frac{d[T]}{dt} = -k2 \times [T] \times [D2]$$



Systems modelling

$$\frac{d[T]}{dt} = -k1 \times [T] \times [D1] - k2 \times [T] \times [D2]$$





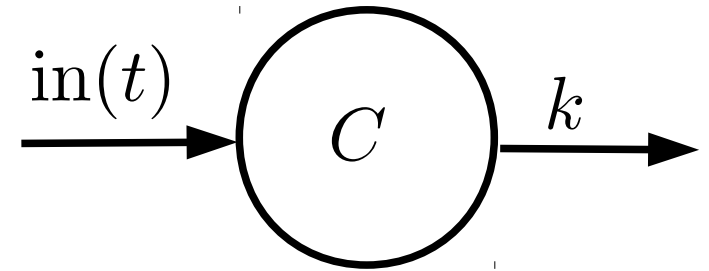
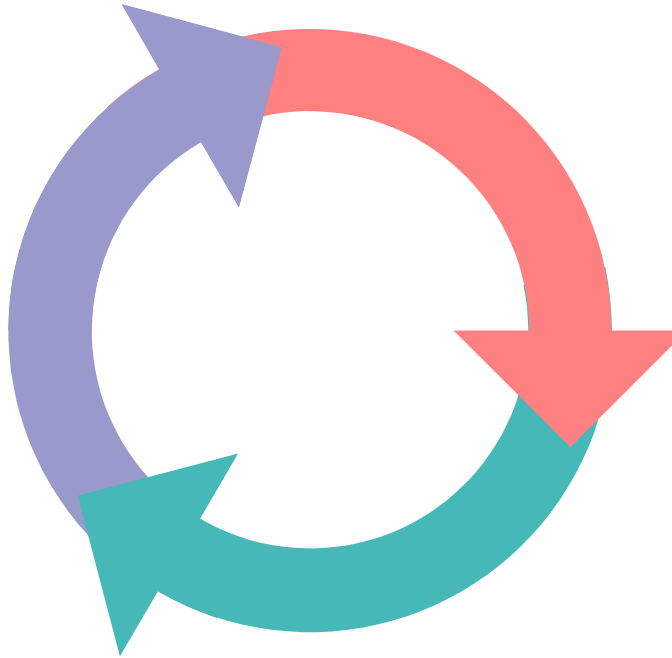
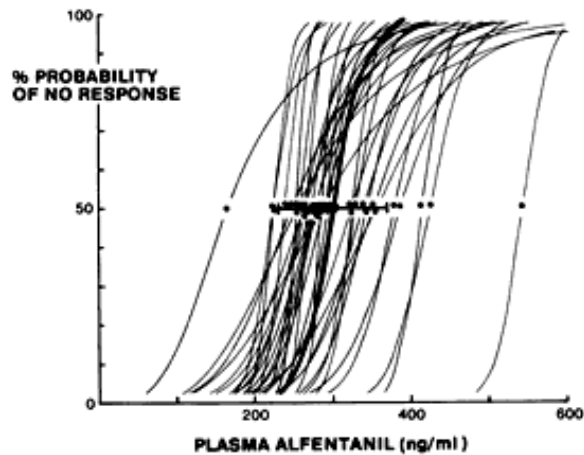
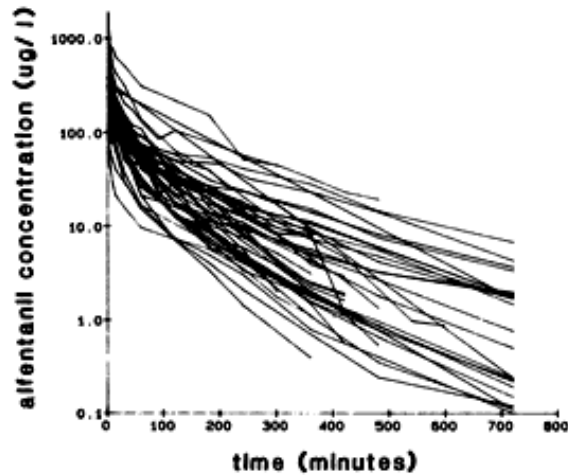
Quantitative Systems Pharmacology

Systems biology
modelling

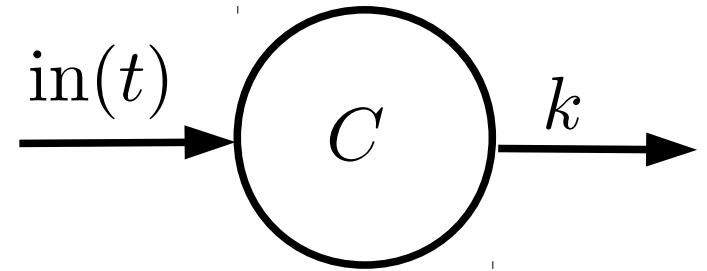
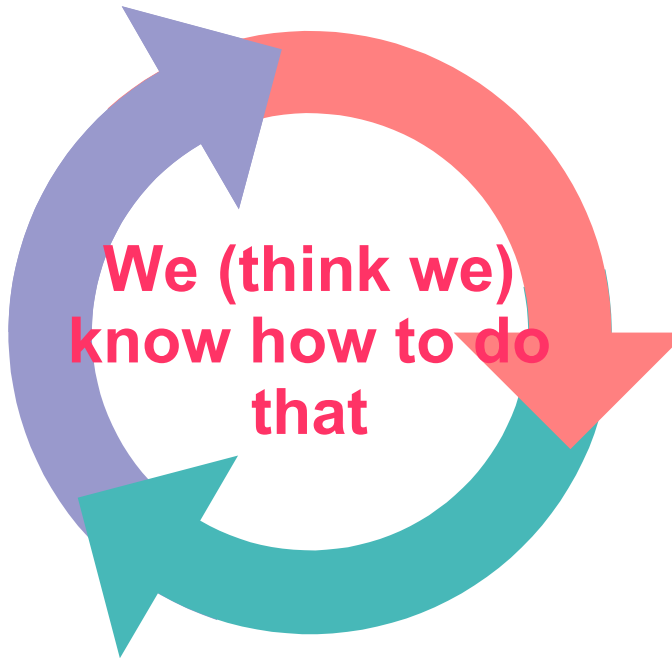
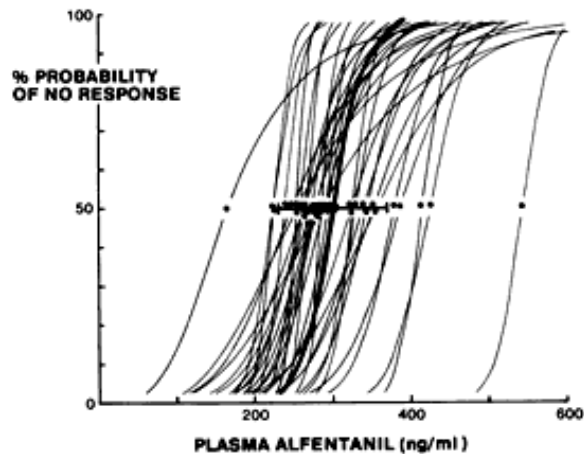
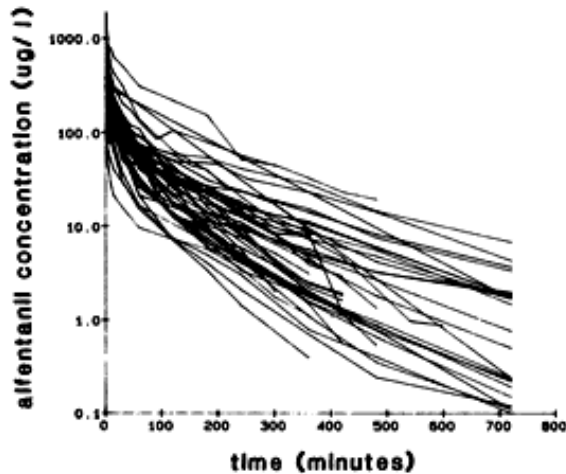
PK/PD
modelling

Statistical
pharmacometrics

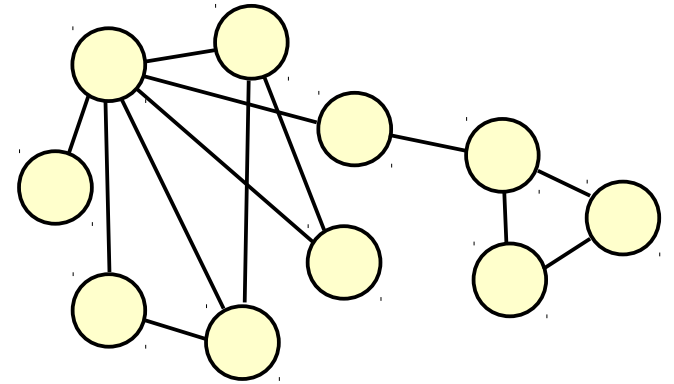
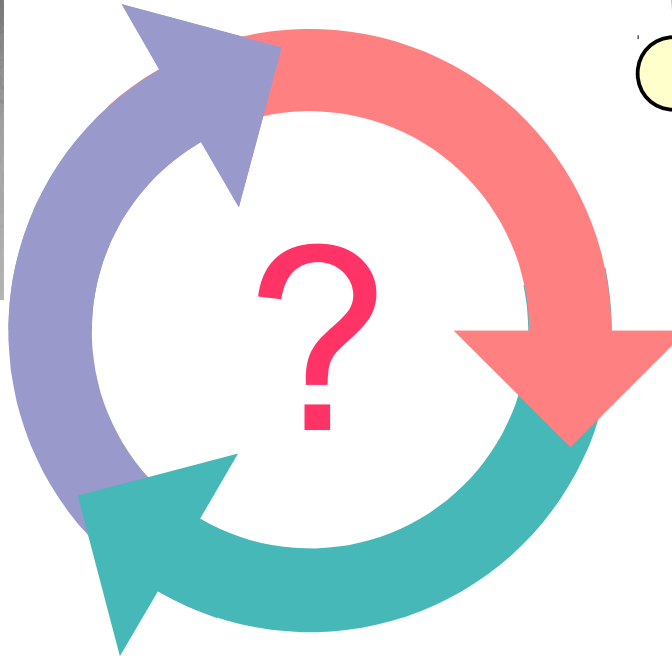
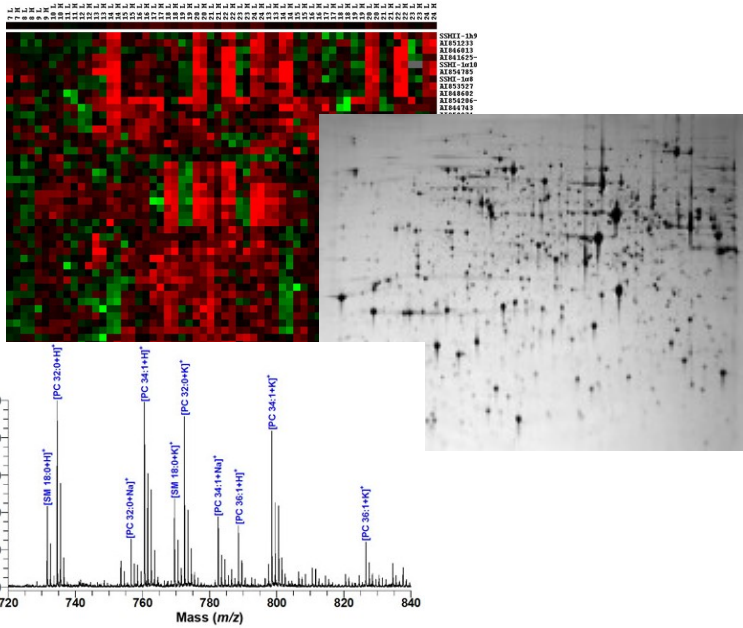
Drug discovery and pharmacometrics models



Drug discovery and pharmacometrics models



Drug discovery and omics



Quantitative Systems Pharmacology

The diagram consists of a large light blue rounded rectangle containing three smaller ovals: a purple one for 'Systems biology modelling', an orange one for 'PK/PD modelling', and a yellow one for 'Statistical pharmacometrics'. To the right of this rectangle is a red question mark. Below the rectangle, outside the light blue area, is a green oval labeled 'Pharmacogenomics'.

Systems biology
modelling

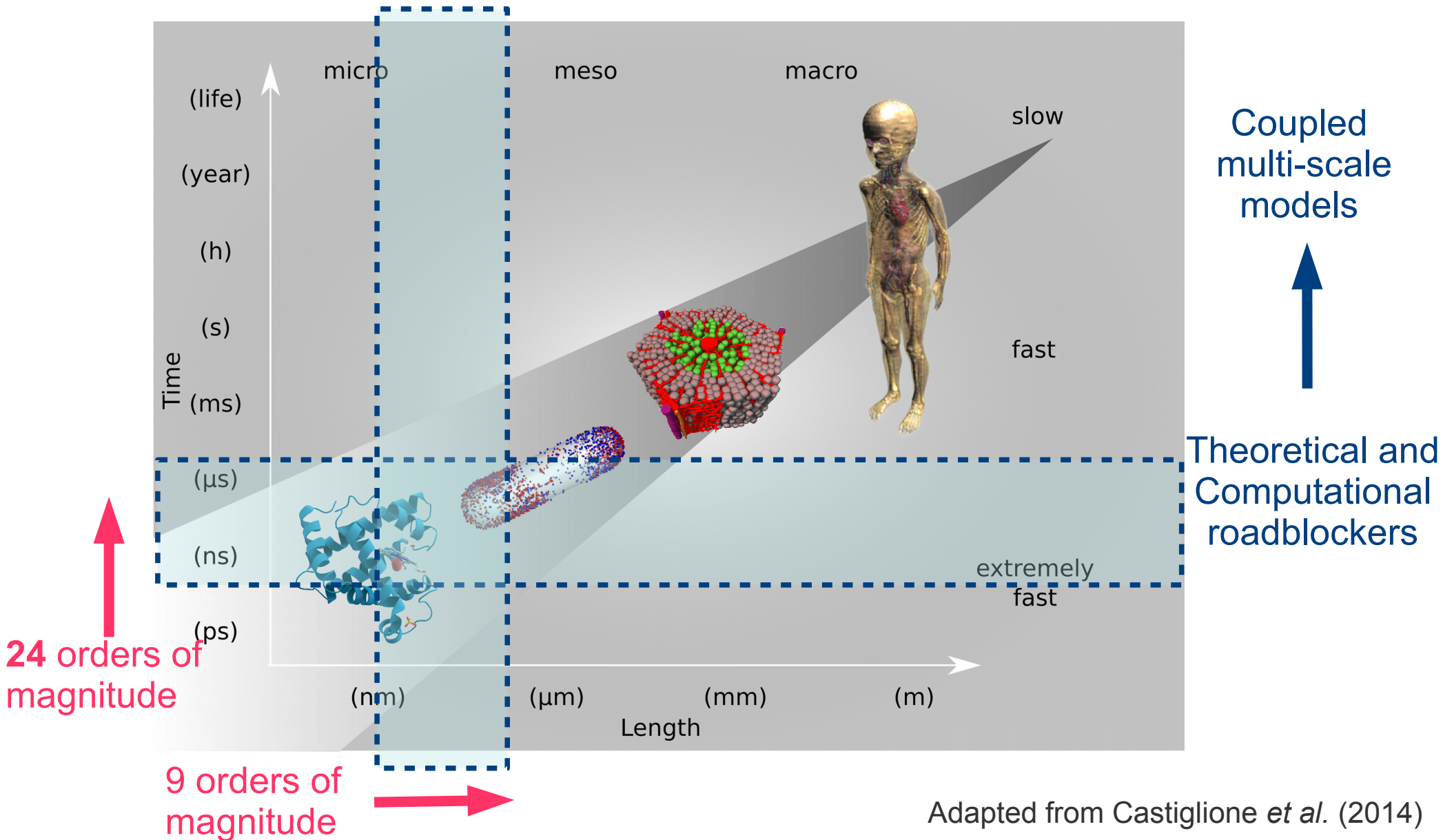
PK/PD
modelling

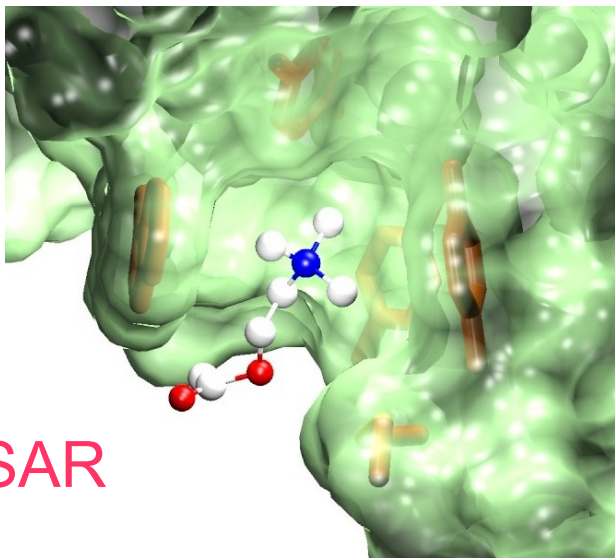
Statistical
pharmacometrics



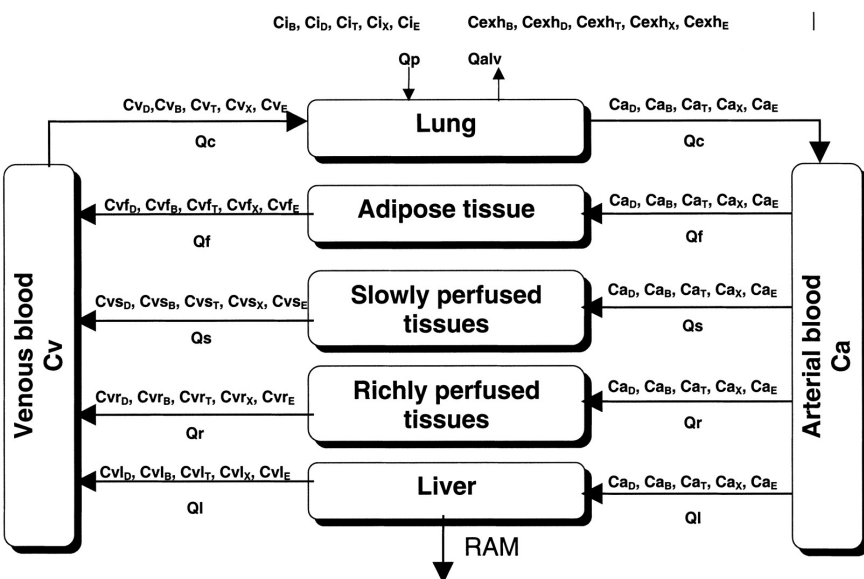
Pharmacogenomics

The problem of scales





QSAR



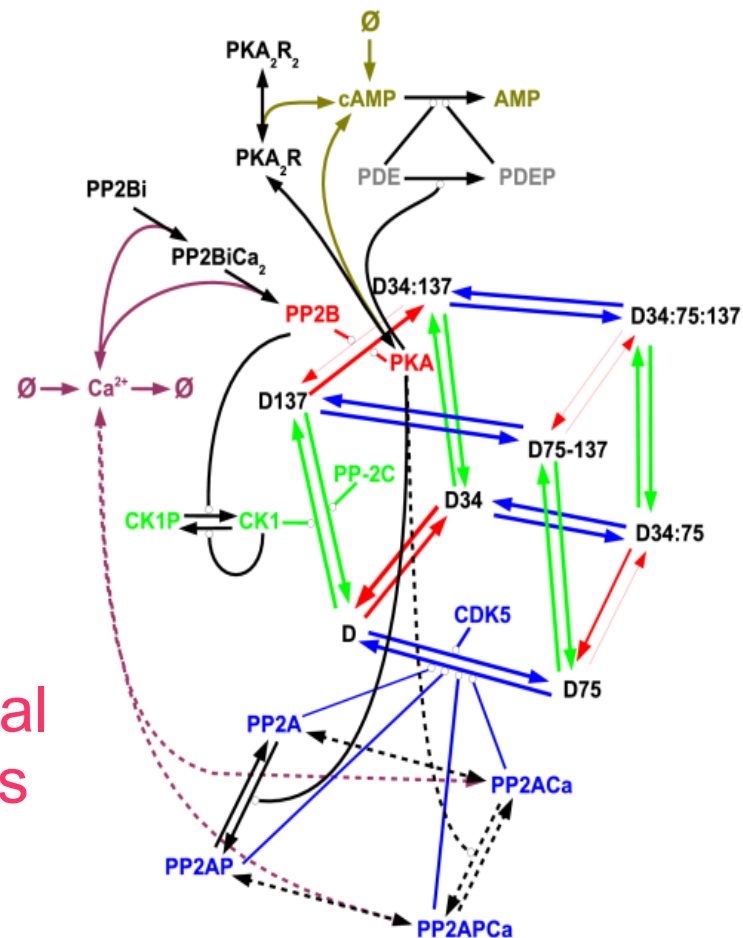
$$\text{RAM}_D = \frac{V_{\max_D} C_{vD}}{K_{mD} \left(1 + \frac{C_{vD}}{K_{iD}} + \frac{C_{vT}}{K_{iD}} + \frac{C_{vX}}{K_{iD}} + \frac{C_{vE}}{K_{iD}} \right) + C_{vD}}$$

$$\text{RAM}_B = \frac{V_{\max_B} C_{vB}}{K_{mB} \left(1 + \frac{C_{vD}}{K_{iB}} + \frac{C_{vT}}{K_{iB}} + \frac{C_{vX}}{K_{iB}} + \frac{C_{vE}}{K_{iB}} \right) + C_{vB}}$$

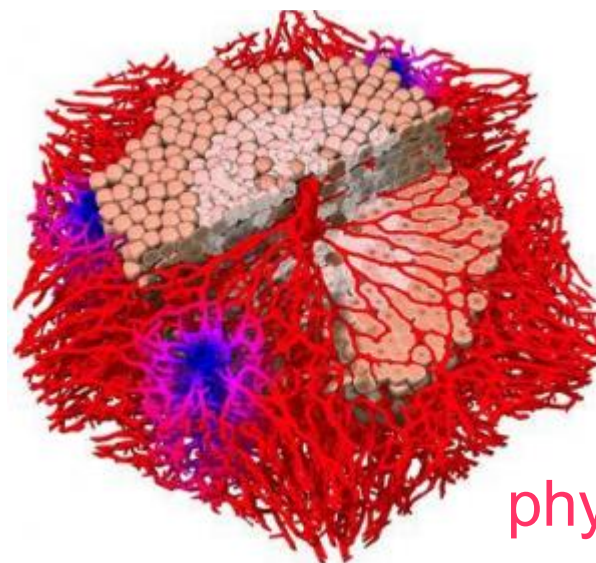
$$\text{RAM}_T = \frac{V_{\max_T} C_{vT}}{K_{mT} \left(1 + \frac{C_{vD}}{K_{iT}} + \frac{C_{vB}}{K_{iT}} + \frac{C_{vX}}{K_{iT}} + \frac{C_{vE}}{K_{iT}} \right) + C_{vT}}$$

$$\text{RAM}_E = \frac{V_{\max_E} C_{vE}}{K_{mE} \left(1 + \frac{C_{vD}}{K_{iE}} + \frac{C_{vB}}{K_{iE}} + \frac{C_{vT}}{K_{iE}} + \frac{C_{vX}}{K_{iE}} \right) + C_{vE}}$$

pharmacometrics

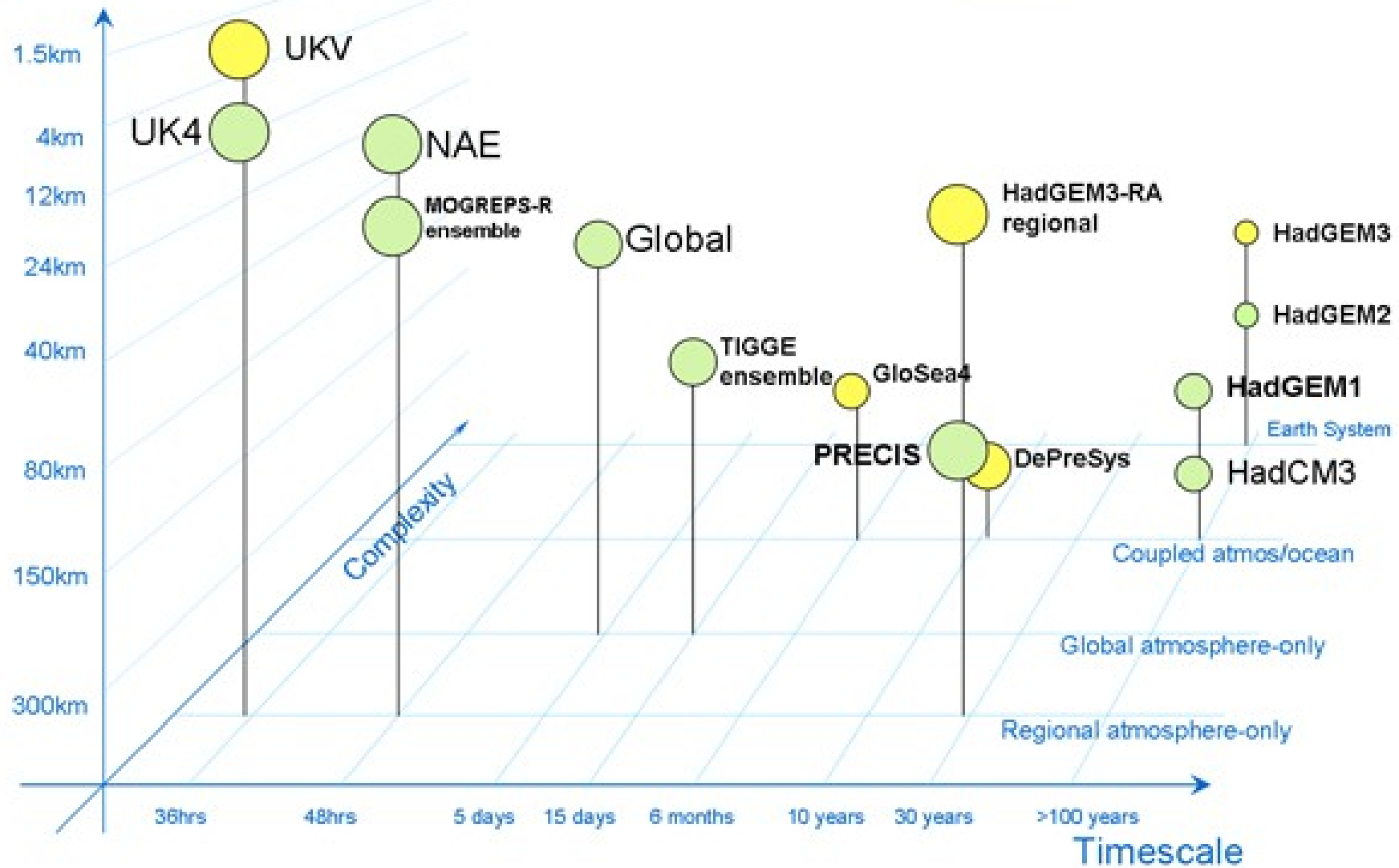


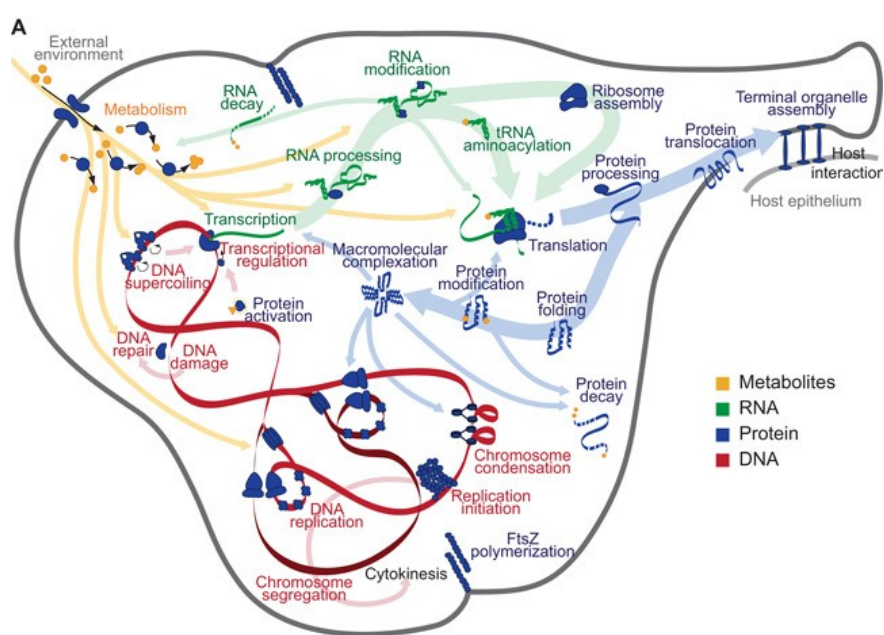
chemical kinetics



physiology

Atmospheric
grid length





Theory

2012 Jul 20

A Whole-Cell Computational Model Predicts Phenotype from Genotype

Jonathan R. Karr,^{1,4} Jayodita C. Sanghvi,^{2,4} Derek N. Macklin,² Miriam V. Gutschow,² Jared M. Jacobs,² Benjamin Bolival, Jr.,² Nacyra Assad-Garcia,³ John I. Glass,³ and Markus W. Covert^{2,4}

¹Graduate Program in Biophysics

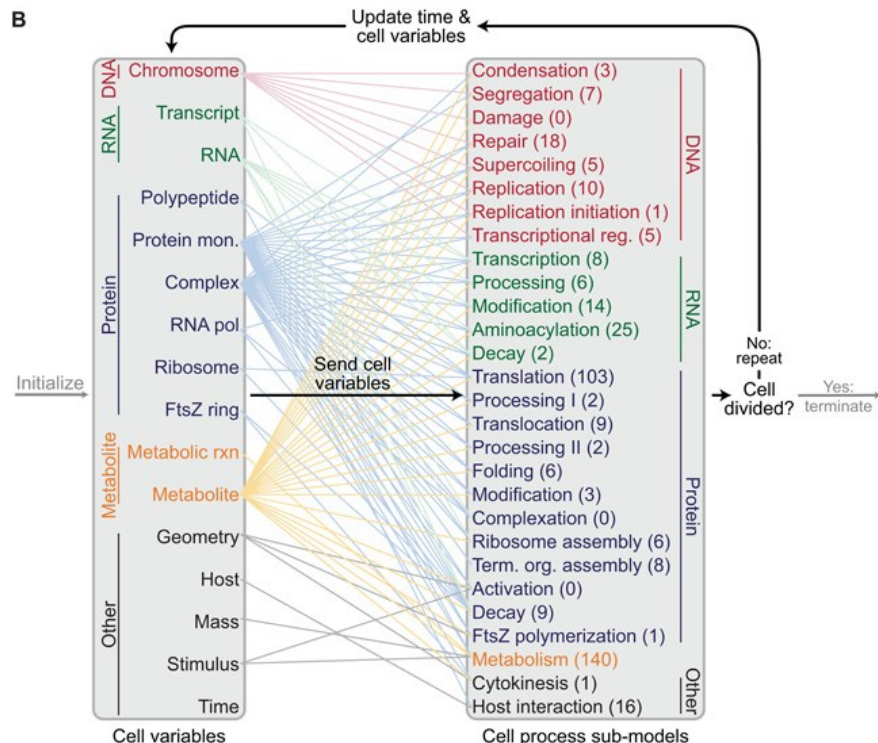
²Department of Bioengineering
Stanford University, Stanford, CA 94305, USA

³J. Craig Venter Institute, Rockville, MD 20850, USA

⁴These authors contributed equally to this work

*Correspondence: mcovert@stanford.edu

<http://dx.doi.org/10.1016/j.cell.2012.05.044>

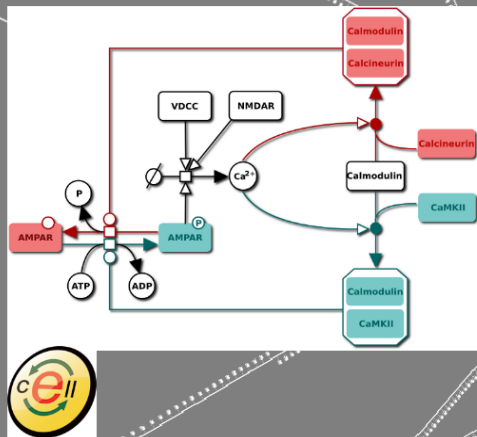


SUMMARY

Understanding how complex phenotypes arise from individual molecules and their interactions is a primary challenge in biology that computational approaches are poised to tackle. We report a whole-cell computational model of the life cycle of the human pathogen *Mycoplasma genitalium* that includes all of its molecular components and their interactions. An integrative approach to modeling that combines diverse mathematics enabled the simultaneous inclusion of fundamentally different cellular processes and experimental measurements. Our whole-cell model accounts for all annotated gene functions and was validated against a broad range of data. The model provides insights into many previously unobserved cellular behaviors, including *in vivo* rates of protein-DNA association

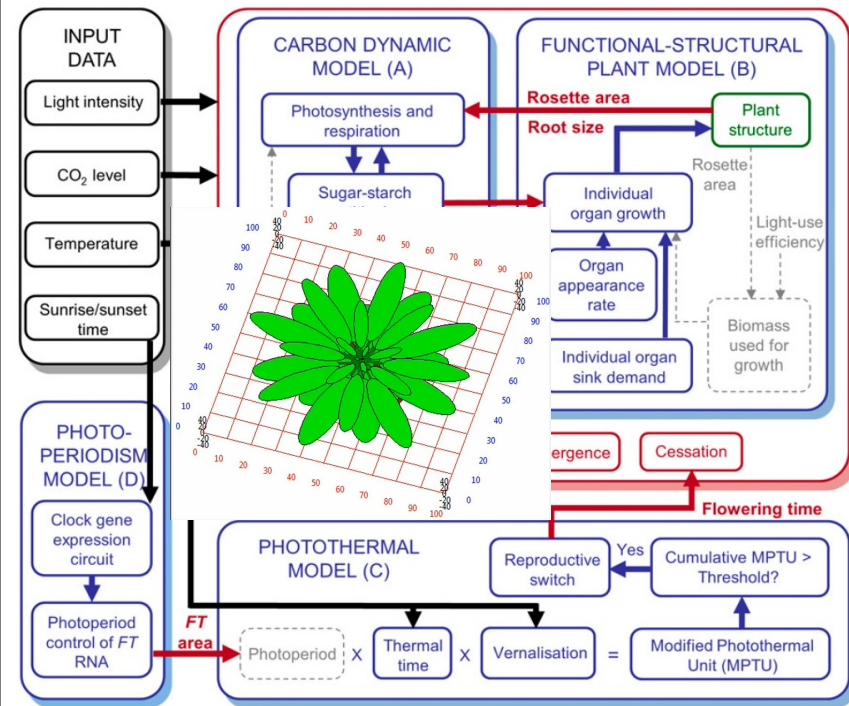
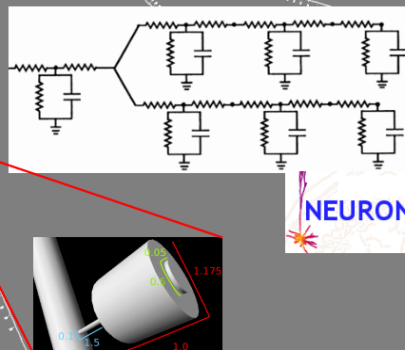
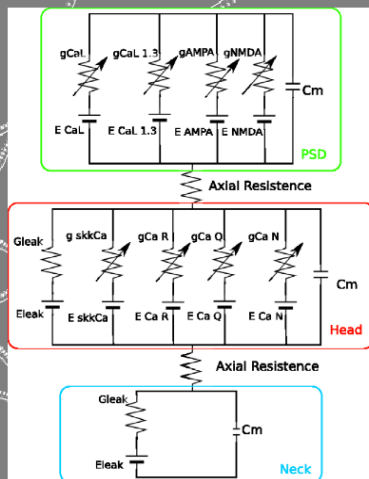
First, until recently, not enough has been known about the individual molecules and their interactions to completely model any one organism. The advent of genomics and other high-throughput measurement techniques has accelerated the characterization of some organisms to the extent that comprehensive modeling is now possible. For example, the mycoplasmas, a genus of bacteria with relatively small genomes that includes several pathogens, have recently been the subject of an exhaustive experimental effort by a European consortium to determine the transcriptome (Güell et al., 2009), proteome (Kühner et al., 2009), and metabolome (Yus et al., 2009) of these organisms.

The second limiting factor has been that no single computational method is sufficient to explain complex phenotypes in terms of molecular components and their interactions. The first approaches to modeling cellular physiology, based on ordinary differential equations (ODEs) (Atlas et al., 2008; Browning et al., 2004; Castellanos et al., 2004, 2007; Domach et al., 1984; Tomita et al., 1999), were limited by the difficulty in obtaining the necessary model parameters. Subsequently, alternative

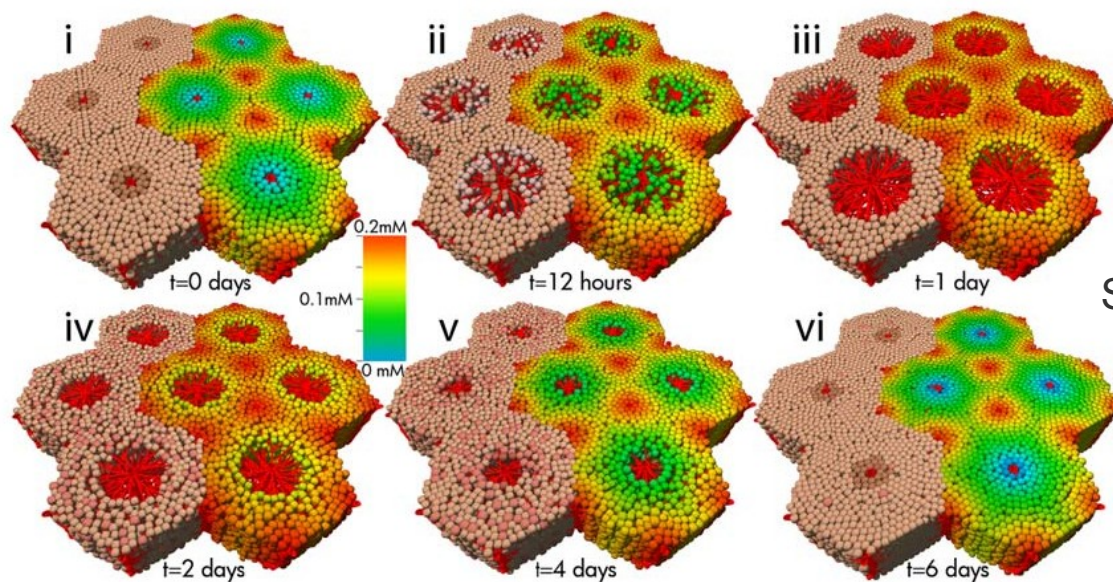


1504 spines
4761 compartments
16362 channels

Mattioni, Le Novère (2013)



Chew *et al.* (2014)



Schliess *et al.* (2014)

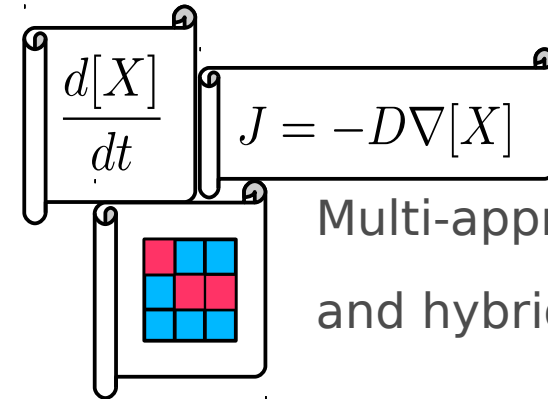
Models need to be developed differently

Modular



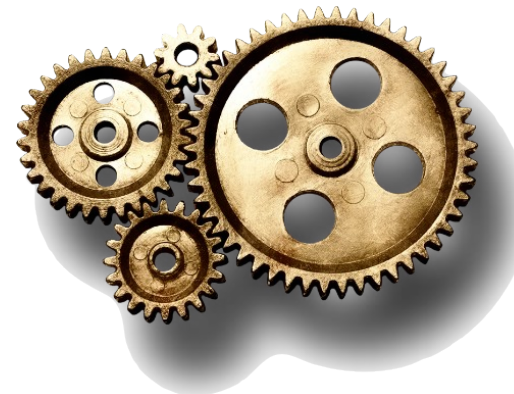
Collaborative

Built on
knowledge-bases



Multi-approach
and hybrid models

Versioned



Multi-software
Coupled
simulations

Models need to be developed differently

Modular

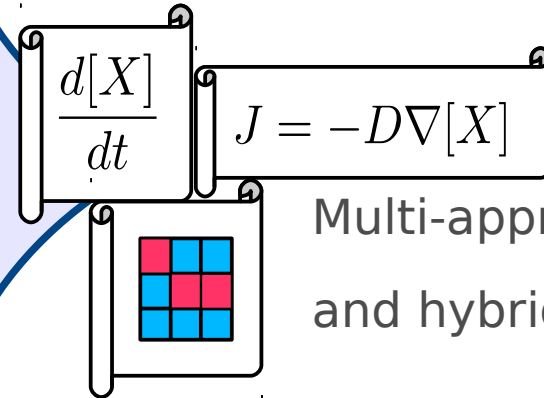


Collaborative

Built on
knowledge-bases

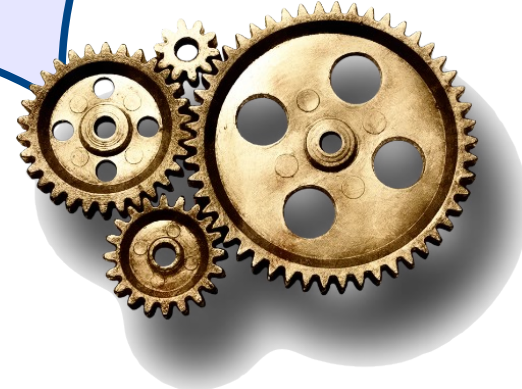


Standards!



Multi-approach
and hybrid models

Versioned



Multi-software
Coupled
simulations

Edda Klipp, Wolfram Liebermeister,
Christoph Wierling and Axel Kowald

Systems Biology

A Textbook

Second Edition



DYNAMIC MODELS



STEPHEN P. ELLNER AND
JOHN GUCKENHEIMER



