Identification of cell signaling pathways based on biochemical reaction kinetics repositories

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Introduction

Cell signaling allows cells to respond to signals that come from its environment changing its behaviour accordingly.

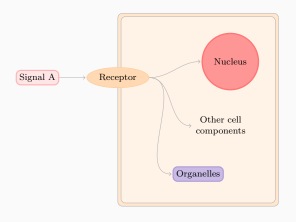
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Understanding the functioning of cell signaling is important in many biological areas.



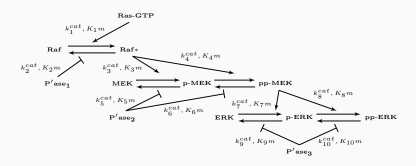
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We call the path of a signal a cell signaling pathway.

Cell Signaling Pathways

A cell signaling network can be characterized by a sequence of chemical reactions



Mathematical Models of Signaling Networks

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Using biochemical kinetics, we can model the concentration change of chemical species over time of a pathway.

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As the input, a description of a biological experiment and a set of experimental measurements are given. A possible output to the problem is composed by:

- a model composed by a set of chemical reactions that are relevant for the biological experiment;
- information about the reaction rate constants of the model.

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Hence, it is desirable to construct a method that can systematically modify these models and choose the one that better represents the experiment.

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On her work, the problem of identification of cell signaling pathways is treated as a feature selection problem.

Feature Selection for Identification of Signaling Pathways

The methodology proposed by Wu defines the set of features as a set of chemical reactions that can be added to a starting model.

Results of Wu's Methodology

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The cost function is defined as the minimum distance between experimental and simulated data. To find this distance a Simulated Annealing procedure is used.

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What we Propose on this Project

We proposed to create a methodology that uses a feature selection approach for identification of signaling pathways, tackling the difficulty of penalizing complex models.

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We used Bayesian approaches of model selection that allow us to create estimates of p(M|D) or p(D|M).

• Study state of the art Bayesian algorithms for signaling network model selection.

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- Implementation and comparison of cost functions for model selection.

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- Formulate systematic modifications to a model as the search space of a feature selection model.

Fundamental Concepts

Kinetics Modeling of Chemical Reactions

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- first order interaction kinetics;
- second order interaction kinetics;
- Michaelis-Menten enzymatic kinetics.

Kinetic Modeling of First Order Iteration

A first order reaction:

$$A \xrightarrow{k_1} B$$

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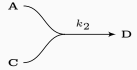
$$A \xrightarrow{k_1} B$$

has rate of:

$$k_1[A]$$
.

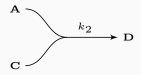
Kinetic Modeling of Second Order Iteration

A second order reaction:



Kinetic Modeling of Second Order Iteration

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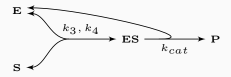


has rate of:

$$k_2[A][C]$$
.

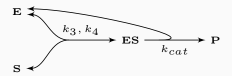
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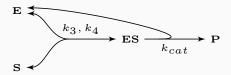
An enzymatic reaction:



Can be divided in two first order reactions plus a second order reaction.

Kinetic Modeling of Enzymatic Reactions

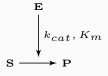
An enzymatic reaction:



Can be divided in two first order reactions plus a second order reaction. However, with the appropriate assumptions, it is possible to use a Michaelis-Menten simplification of this reaction.

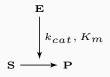
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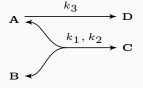
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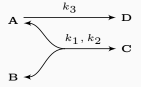
$$k_{cat} \frac{[E][S]}{K_M + [S]}.$$

Suppose we want to model the kinetics of A on these reactions:



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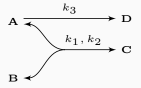
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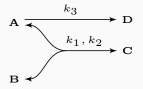
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- A \longrightarrow D, with rate $k_3[A]$.

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Then, the differential equation that models the concentration change of A is:

$$\frac{d[A]}{dt} = -k_1[A][B] + k_2[C] - k_3[D].$$

Bayesian Methods for Biochemical Model Selection

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For both methods, we resort to Metropolis-Hastings algorithm to generate samples of distributions.

With Metropolis-Hastings, we can generate a sample of a distribution $p(\lambda)$ doing the following:

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- 4. Set $\lambda_t = \lambda^*$ with probability min(1, r) and $\lambda_t = \lambda_{t-1}$ otherwise;
- 5. Increase *t* by one and repeat from Step 2 if not reached iteration limit.

Model Selection

Marginal Likelihood Estimation

Likelihood of Data Given Model and Parameters

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$$p(\mathbf{D}|M, \boldsymbol{\theta}) = p_{\mathcal{N}_{(\vec{0}, \Sigma)}}(\phi(M, \boldsymbol{\theta}) - \boldsymbol{D}).$$

Where $\phi(M, \theta)$ is the simulated observation.

Marginal Likelihood of Data

We can marginalize the likelihood to obtain:

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Calculating this integral is hard, therefore we resort to estimating another integral.

Power-posterior distributions

We define a power-posterior distribution as:

$$p_{\beta}(\boldsymbol{\theta}) = \frac{p(\boldsymbol{D}|\boldsymbol{\theta}, M)^{\beta} p(\boldsymbol{\theta}|M)}{\int_{\boldsymbol{\Theta}} p(\boldsymbol{D}|\boldsymbol{\theta}, M)^{\beta} p(\boldsymbol{\theta}|M) d\boldsymbol{\theta}},$$

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Note that:

$$p_{\beta=0}(\theta)=p(\theta|M),$$

and that

$$p_{\beta=1}(\theta) = \frac{p(\boldsymbol{D}, \theta|M)}{\int_{\Theta} p(\boldsymbol{D}, \theta|M) d\theta} = \frac{p(\theta|\boldsymbol{D}, M)p(\boldsymbol{D}|M)}{p(\boldsymbol{D}|M)} = p(\theta|\boldsymbol{D}, M).$$

The Thermodynamic Integral

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$$\ln p(\mathbf{D}|M) = \int_0^1 \mathbb{E}_{p_{\beta}(\boldsymbol{\theta})}[\ln p(\mathbf{D}|\boldsymbol{\theta}, M)]d\beta.$$

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$$\sum_{t=0}^{T-1} (\beta_{t+1} - \beta_t) \frac{\mathbb{E}_{p_{\beta_{t+1}}(\boldsymbol{\theta})}[\log p(\boldsymbol{D}|M,\boldsymbol{\theta})] + \mathbb{E}_{p_{\beta_t}(\boldsymbol{\theta})}[\log p(\boldsymbol{D}|M,\boldsymbol{\theta})]}{2}$$

To produce the estimates of

$$\mathbb{E}_{p_{\beta_t}(\boldsymbol{\theta})}[\log p(\boldsymbol{D}|M,\boldsymbol{\theta})]$$
 for $t \in \{0,\ldots,T\}$

we need to produce samples of the power-posteriors $p_{eta_t}(heta)$.

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On the second sampling step, called posterior shaped burn-in, we use the covariance of the current sample times some constant as the covariance of the jump distribution.

On the third step, we perform the Populational Monte Carlo Markov Chain sampling. This algorithm allows us to mix samples from different power posteriors.

Ranking with Approximate Bayesian Computation

Approximate Bayesian Computation (ABC) is a method that allows one to obtain samples of a distribution close to $p(\theta, M|D)$.

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- 4. Repeat until some iteration limit.

The result of the algorithm is a sample of the distribution

$$p(\theta, M|d(\phi(M, \theta), \mathbf{D}) < \epsilon).$$

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We can use the accepted parameters of tolerance ϵ and model M to estimate

$$p(M|d(\phi(M, \boldsymbol{\theta}), \boldsymbol{D}) < \epsilon).$$

Development of SigNetMS

The SigNetMS Software

To choose a cost function, we needed to compare the ABC-SMC and Marginal Likelihood approaches for model selection.

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- ABC-SysBio is a software that implements ABC-SMC
- BioBayes is a software that implements the estimation of the marginal likelihood.

However, the usage of BioBayes in our context was cumbersome.

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The SigNetMS software

SigNetMS is a Python package and command line software that estimates the marginal likelihood of data given a model, p(D|M).

The input expected by SigNetMS

The input to SigNetMS includes:

- An SBML file model;
- An XML file with prior distributions of parameters;
- An XML file with experimental data;

The output produced by SigNetMS

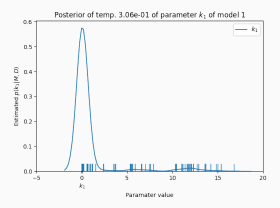
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 $Fast\ integration\ and\ parameter\ sampling$

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The first implementation of SigNetMS did not cope with larger instances of model selection. We tackled this problem in two ways:

- change the representation of the system of ordinary differential equations;
- implement parallelization.

Changing the representation of the system of ordinary differential equations

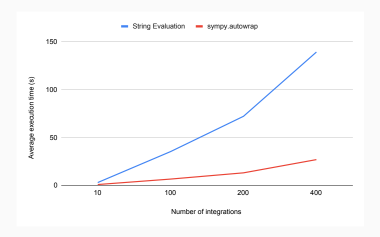
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Changing the representation of the system of ordinary differential equations

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We used SymPy to provide automatically generated code that allowed us to create a C function to represent the system of ODEs.

Comparing the representation of the system of ordinary differential equations



Parallelizing the sampling of multiple power posteriors

The first two phases of the sampling procedure occurs independently between different power posteriors.

Parallelizing the sampling of multiple power posteriors

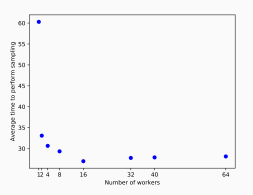
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Experiments and Results

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- Comparison between ABC-SysBio and SigNetMS
- Solving model selection as a feature selection instance

 ${\tt Comparison \ between \ ABC-SySBio \ and \ SigNetMS}$

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ABC-SysBio and SigNetMS use different Bayesian approaches for model selection. The first creates an estimate of p(M|D), and the second creates an estimate of p(D|M) (the marginal likelihood).

To compare both software we ran an experiment based on the following procedure:

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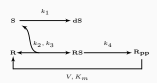
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- Rank the four models.

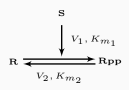
The instance

This instance is originally from Vyshemirsky and Girolami (2007), in which they present results of Annealing Melting Integration.

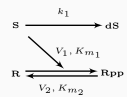
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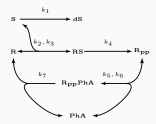
The "correct" model



The incorrect model

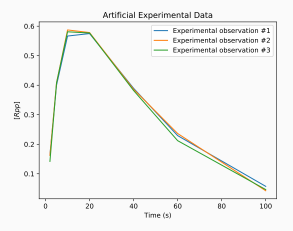


The simplification model



The generalization model

The instance



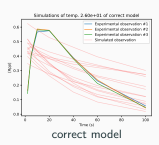
The ABC-SysBio software returned the following ranking of models:

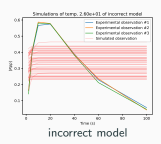
1. incorrect model

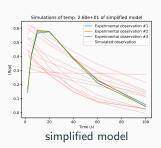
- 1. incorrect model
- 2. simplification model

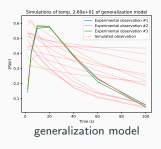
- 1. incorrect model
- 2. simplification model
- 3. generalization model

- 1. incorrect model
- 2. simplification model
- 3. generalization model
- 4. correct model









The ranking returned by SigNetMS on the first experiment is:

1. correct model

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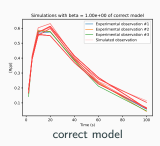
- 1. correct model
- 2. simplification model

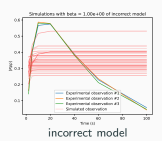
The ranking returned by SigNetMS on the first experiment is:

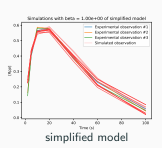
- 1. correct model
- 2. simplification model
- 3. generalization model

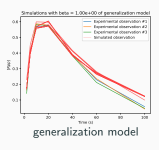
The ranking returned by SigNetMS on the first experiment is:

- 1. correct model
- 2. simplification model
- 3. generalization model
- 4. incorrect model









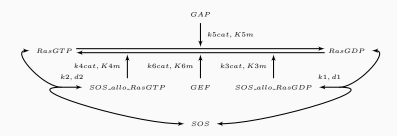
Simulations generated by the correct model $% \left(1\right) =\left(1\right) \left(1\right) \left($

 ${\tt Model \ selection \ as \ a \ feature \ selection \ problem}$

After defining that SigNetMS is our software choice for a cost function, we are able to experiment the approach of solving a model selection instance as a feature selection problem.

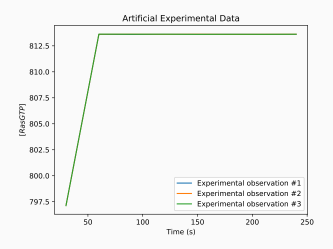
We proposed a Ras switch pathway to experiment on.

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The concentration of activated Ras was measured at the time steps of 30, 60, 90, 120, 150, 180, 210, and 240 seconds.

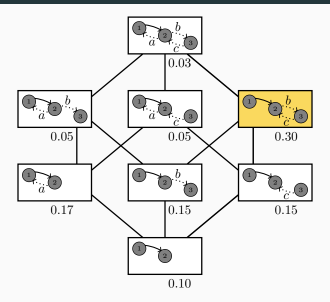
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The feature selection problem consists in finding the best subset of a set of features, S, given a cost function c.

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If we define the set of feature as a set of reactions, we can create a feature selection instance that represents a model selection instance.

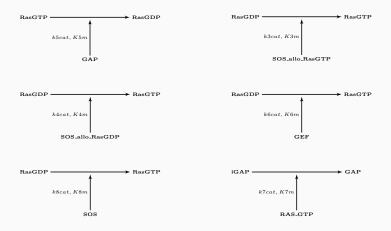


In the instance we prepared, the base model has zero reactions,

In the instance we prepared, the base model has zero reactions, and the set of features ${\cal S}$ is composed by 10 reactions,

In the instance we prepared, the base model has zero reactions, and the set of features S is composed by 10 reactions, 8 of them present on the correct model.





The search space $\mathcal{P}(S)$, has 2^{10} . Therefore, a heuristic is necessary to traverse the space.

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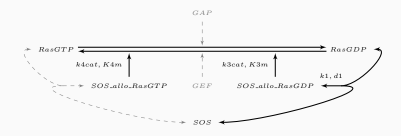
In the SFS procedure, we start from the bottom of the search space.

In the SFS procedure, we start from the bottom of the search space. And for every iteration, we select the best adjacent model that has one more reaction.

Characteristic Vector	Score	Cost function time (seconds)
000000000	330721.05	851.3
0010000000	245681.93	1083.4
0010010000	211.62	4257.4
0011010000	-1.32	5007.71
0011011000	-4.27	4458.7
0111011000	-7.90	5035.7

The found model is contained in the correct model:

The found model is contained in the correct model:



Simulations generated by the found model

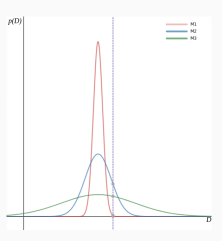
Simulations generated by the correct model $% \left\{ 1\right\} =\left\{ 1\right\} \left\{ 1\right\} =\left\{ 1\right\} \left\{ 1\right\} \left\{$

Results of the search

In this experiment, we experienced a known feature of marginal likelihood approaches:

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In this experiment, we experienced a known feature of marginal likelihood approaches: intermediate complexity models are preferred.



Conclusions

The main contributions of this work are:

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- the implementation of the SigNetMS software;
- the comparison between SigNetMS and ABC-SysBio;
- the experimentation of feature selection on model selection using a marginal likelihood approach to define the cost function.

We also suggest a few topics for future related work:

efficiency improvements on SigNetMS;

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- treatment of numerical instabilities on numerical integrations of SigNetMS;

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- treatment of numerical instabilities on numerical integrations of SigNetMS;
- solving the model selection problem as a U-Curve problem;
- experimentation on heterogeneous conditions of experimental measurements;
- application of the methodology on real instances.

Thank you!