

# Accurate parameter optimization leads to predictive dynamical models for systems biology

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Signaling Biology

Modeling and data fitting framework

Optimization for parameter estimation

Results on the angiontensin signaling pathway

#### Signaling Biology

Modeling and data fitting framework

Optimization for parameter estimation

Results on the angiontensin signaling pathway

- ► A Ligand binds a receptor in the cell surface and leads to a signal.
- The bound receptor-ligand complex leads to a cascade of reactions (enzymatic catalysis, phosphorylation,...) up to some effector molecule that leads to a cellular response.



- ► A Ligand binds a receptor and signals.
- The bound receptor-ligand complex leads to a cascade of reactions.
- G Protein Coupled Receptor (GPRC) : Family of receptor, widely targeted by drugs.
- Two main pathways : G protein pathway and β-arrestin (signal vs internalization)



- ► A Ligand binds a receptor and signals.
- The bound receptor-ligand complex leads to a cascade of reactions.
- ▶ G Protein Coupled Receptor (GPRC)
- Two pathways : G protein and β-arrestin.
- More complex issues : β-arrestin induced pathway leads to a different signal on the same effector.



- ► A Ligand binds a receptor and signals.
- The bound receptor-ligand complex leads to a cascade of reactions.
- ▶ G Protein Coupled Receptor (GPRC)
- Complex interactions between G protein and β-arrestin pathways.

#### Drug discovery

 Signaling through one pathway and not another one : Bias (synthetic hormone, mutant receptor, small molecules...)



- A Ligand binds a receptor and signals.
- The bound receptor-ligand complex leads to a cascade of reactions.
- ► G Protein Coupled Receptor (GPRC)
- Complex interactions between G protein and β-arrestin pathways.

#### Computational Modeling

- Help deciphering the intricate effect of each pathway.
- Quantify the precise effect of a specific couple Ligand-Receptor.



#### GPCR signaling through ERK phosphorylation

The extracellular signal-regulated kinase ERK is activated both by the G protein and the  $\beta$ -arrestin pathway but (*Ahn et al. J Biol Chem (2004)*) :

- The spatial distribution are distinct.
- The kinetics are distinct.



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Transient and sustained ERK activation have been shown to regulate cell fates such as **growth** and **differentiation**.(*Sasagawa et al. Nat Cell Biol* (2005))



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- The spatial distribution are distinct.
- ► The kinetics are distinct.

 $\beta$ -arrestin 2 dependent ERK pathway can be activated **independently** of G proteins with a mutant receptor (*Wei et al. PNAS (2004)*).



#### Case study : Angiotensin receptor

- Angiotensin II type 1A receptor (AT1AR) transfected in cultured human embryonic kidney (HEK 293 cells).
- ERK phosphorylation data : Phosphorylated ERK in immunoblots, quantified by densitometry (*Kim et al. PNAS* 2005)
- DAG accumulation and PKC activity data, measured in real time by FRET sensors.
- Four perturbed conditions in addition to control :
  - $\beta$ -arrestin 2 siRNA
  - G protein-coupled receptor kinases (GRK2/3 and GRK5/6) siRNA
  - PKC inhibitor.

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 Starting point : graph of interaction of molecules (based on biological knowledge, literature)



#### Graph of interaction of molecules

 Law of mass-action : Ordinary Differential Equations (ODE) produce timedependent trajectories, that depend on parameters (kinetic rate, initial condition)

$$\frac{d[B]}{dt} = k_0[A] - k_1[B].$$
$$\frac{d[B]}{dt} = k_0[A][C] - k_1[B].$$



- Graph of interaction of molecules.
- ► Law of mass-action : ODE.
- Quality of the model based on the introduction of a cost function (based on statistical error model, or heuristic arguments).



- Graph of interaction of molecules.
- ► Law of mass-action : ODE.
- Cost function.
- Optimization of the cost function (Frequentist / Bayesian approach). Numerical search.



- Graph of interaction of molecules.
- ► Law of mass-action : ODE.
- Cost function.
- Optimization.
- Validation data, prediction and experimental design...

Signaling Biology

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The major difficulties are due to

- ► Large dimension of parameter space (10 100) and state space (> 10).
- Few molecule concentrations measured, and not in absolute numbers.
- Large ODE's may be numerically costly to simulate if they are stiff.
- Parameters can be non-identifiable (non-convexity, presence of many local minima)

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A variety of methods can be employed : local/global methods and deterministic/stochastic methods, hybrid method.

- Gradient descent methods with many random initial start (D2D, Raue A., et al. Bioinformatics (2015)).
- Hybrid local and global method, based on heuristics (HYPE, T. Bourquard & A. Poupon)

#### H.Y.P.E



GENETIC ALGORITHM:

· A random set of n parameter sets is chosen, called parents.

A random set of parameters is optimized using Genetic algorithm. The resulting parameter set is optimized using CMA-ES. This operation is repeated until a sufficient number of parameter sets giving small errors is obtained.

#### Critical assessment of methods

How to judge different method? How to asses the quality of a fit?

- Toy models with *in silico* simulated data / Benchmark models.
- Absolute value of the error function.
- **Speed** of the algorithm.
- Convergence curve (number of best error function value over number of runs/function evaluation.
- Robustness of the minima.

#### Critical assessment of methods

How to judge different method? How to asses the quality of a fit?

Why/How to deal with Non-Identifiability (NI)?

- It slows down the numerical search and leads to unreliable results.
- Theoretical NI : reduction / algebraic relations.
- Numerical NI : distinguish between structural and practical NI. Calculate sensitivity and one-dimensional profile likelihood.

# Toy models









# Toy models : HYPE gives comparable results to high quality optimization methods



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#### Full model (Heitzler el al. MSB 2012) : 3 pathways

A model that fit the data : role of **G** protein-coupled receptor kinases (GRK) in the balance of signals.



#### Full model (Heitzler el al. MSB 2012) : Control + 4 pert.



#### The model fits the data...



#### In black : control experiments. In Red : perturbated experiments.

#### The model correctly predicts some validation data...



#### Remark

Good correlation between error and prediction.

# ...but the convergence is poor, and the identifiability a serious issue !



#### Critical assessments :

- Convergence curve. Identifiability of parameters.
- Parsimonious use of parameters. Model selection.

From 50 parameters...



... to 22 parameters !



The reduced model still fit (reasonably) well...



- ▶ The reduced model still fit well...
- and the convergence is better...



- ▶ The reduced model still fit well...
- ▶ and the convergence is better...
- but the identifiability is still poor !



- The reduced model still fit well...
- and the convergence is better...
- but the identifiability is still poor! Let's reduced further?





















#### Results

With model selection criteria :

- a model with 19 parameters
- good convergence properties (10% of runs reached the optima)
- most parameters are identifiable













4.0 4.0 22.6 Magael 40 8 81 813 km\_(0)

#### What do we learn (so far)?

The three pathways are a necessary condition to reproduce the data.



#### What do we learn (so far)?

- Three pathways are necessary.
- An internalization pathway, independent of the β-arrestin signaling pathway, is mandatory.



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- Three pathways are necessary.
- An independent internalization pathway is mandatory.
- A minimal model with three reversible pathway with 10 parameter is able to fit the phospho ERK data and its parameter are all identifiable.



## What do we learn (so far)?

- Three pathways are necessary.
- An independent internalization pathway is mandatory.
- A minimal model can fit the phospho ERK data and is identifiable.
- The best model able to fit
  all data present non-identifiability
  - ⇒ Experimental design.



#### Conclusion

- A full model able to fit the data (*Heitzler el al. MSB 2012*).
- Accurate parameter estimation leads to accurate prediction.
- Further improvements with model reduction/selection.
- Parameter identifiability with a good fit can be achieved.
- We have shed light on the importance of three pathways in GPCR signaling, and its regulations.

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#### Thanks for your attention !

#### validation data

