

### Identification of dynamical models of genetic networks

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#### Outline

- The problem of genetic network identification
- A traditional approach: Boolean networks
- Identification of Ordinary Differential Equation (ODE) models
  - The general problem
  - Linearization methods (steady-state, time series)
  - Boolean-like methods (time series)
- Identification of stochastic models
  - An overall view
  - Focus: The Finite State Projection (FSP) Method
- Conclusions

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# The problem of genetic network identification



#### Gene networks



#### A GENE REGULATORY NETWORK



(Wikipedia)

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#### Example: Gene network of E.coli



(Martinez-Antonio et al, J Mol Biol, 2008)

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# About the statement of the identification problem

- Goal: Estimate a mathematical model of a network of genes from experimental observations of the system
  - Why ? What model do we want ? What do we want from that model ?
- The model should the describe structure and behavior of the network
  - Structure: genes and their interconnection
  - Behavior: inhibition vs. activation, dynamics
- Several different problems depending on the context
  - What data ?
  - What prior knowledge ?
  - What use of the model ?

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#### Scale

- Different levels of detail:
  - genes, but also mRNA, transcription factors, protein complexes...
  - expression: binding, DNA unfolding, transcription, translation, ...



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 $g_{_{4}}$ 

Coding Strand

Strand

Template

(Wikipedia)

 $G_{a}$ 

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### Information

- Modelling framework depends on available data...
  - Type, quality, quantity
  - System excitation, experimental conditions



• ... and on the use of the model



#### GFP fusions (courtesy of Z.Lygerou)



- Analysis (learning how cells function) & Prediction (response of an organism to perturbations/stimuli)
- Control (industrial exploitation, targeted chemicals for medical therapies...)
- Engineering of new functions ("Synthetic biology")

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### Modelling tradeoffs

- Qualitative vs. quantitative
- Mechanistic vs. phenomenological
- Fitting accuracy vs. predictive power (overfitting!)

- Complexity vs. identifiability
- Static vs. dynamic
- Black-box vs. grey-box vs. white-box



## The identification circle

- Model hypothesis:
  - Choice of modelling framework
  - Formulate model hypotheses
- Experiment design:
  - Choose experimental setup
  - Design most informative experiments
- Estimation:
  - Execute experiment and get data
  - Find model(s) that explains data
- Validation:
  - Inspect results
  - Evaluate predictive capability

Today's focus: formal statement of gene network inference problems and solution with selected methods







#### A traditional approach: Boolean networks



#### **Boolean models**

- Formalism to model regulatory effects (mutual activation, inhibition) from qualitative gene expression data
- N Boolean variables representing n genes

$$(X_1,X_2\ldots,X_n)\in\{0,1\}^n$$

 $X_i = 0$  gene not expressed

 $X_i = 1$  gene expressed

Boolean regulation function

 $X_i$  expressed iff  $b_i(X) = 1$ 

• Dynamic Boolean networks (discrete time):

 $X_i(t+1) = b_i(X(t))$  i = 1, ..., n t = 0, 1, 2, ...

• Network structure captured by gene interaction graph







#### Motivation: Microarray data

• Gene expression profiling by DNA microarrays:

- Isogenic cell populations placed in microscopic wells containing probes for specific mRNA molecules (genes)
- Combined with the use of fluorescence reporters, binding of mRNA-specific probes leads to fluorescence of the cells in the corresponding well
- Thousands of genes for wild-type and mutated cells can be observed in parallel, at low temporal resolution (one microarray prepared per measurement time)



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- Goal of identification: reconstruct logical interactions among genes
  - Network "structure" (graph edges) and "dynamics" (regulation rules)
  - From dynamical ON/OFF time series (system observed in transient) or from steady state ON/OFF data (system equilibrium for different perturbations)
  - Established methods exist (e.g. REVEAL)

Two words on inference of

**Boolean models** 

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Perturbation

#### **REVerse Engineering ALgorithm**

#### (Liang et al, 1998)

#### Based on information-theoretic concepts

 $X_1, \ldots, X_n$  random variables  $H(X_i)$  entropy ("variability") of  $X_i$  $M(X_i, X_i)$  mutual information of  $X_i$  and  $X_i$ generalizations to sets of variables

 $\frac{M(X_i,X_j)}{H(X_i)} \in (0,1)$  $0 = X_i$  is independent of  $X_i$  $1 = X_i$  is fully determined by  $X_i$ 

- Functions of probability distribution of X •
- Estimated from the observed trajectories of X
- Used to determine the effective inputs of a Boolean update map, e.g. • If  $\frac{M(X_1(t+1), [X_2(t), X_3(t)])}{H(X_1(t+1))} = 1$  then  $X_1(t+1) = b_1(X_2(t), X_3(t))$
- Specific form of update map determined from the observed transitions
- May cope with noise (measurement error)
- Worst case: evaluation of all possible combinations of inputs
  - Bound complexity with maximum allowable number of inputs





#### Simulation example

P (incorrect solution)

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•Vast literature on Boolean model analysis (Kauffman, ...)

- •Unsatisfactory description of quantitative phenomena, may lead to poor results
- •Starting point for quantitative dynamical modelling



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#### Identification of ODE models



### The model family

- Formalism to model average gene expression dynamics based on ensemble gene expression data from a population of cells
- Vector of concentrations:  $x = (x_1, \ldots, x_n) \in \mathbb{R}_{\geq 0}^n$
- ODE model:  $\dot{x}_i = f_i(x, u, \theta) \Gamma_i(x, u, \theta)$ 
  - $f_i \ge 0$  synthesis rate functions
  - $\Gamma_i \geq 0$  degradation rate functions
  - $heta \in \Theta$  unknown parameters (and structure)
    - u(t) perturbation input
- Common situation: unregulated degradation,  $\Gamma_i(x_i) = \gamma_i x_i$
- Depending on the identification approach, specific (parametric) form for rate functions

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### Model family: examples

• Linear model plus saturation (Jaeger et al, Nature 2004):



#### The data

• Measurement model

$$y_i(t) = h_i(x_i(t), e),$$
   

$$\begin{cases} h_i & \text{output function} \\ e & (random) \text{ measurement noise} \end{cases}$$

(not always used in full detail)

• Data set

$$\mathcal{D} = \{y^m(t_k): k = 1, ..., K, m = 1, ..., M\}$$

- K measurement times
- *M* time series (possibly different inputs)
- Sometimes, corresponding synthesis rates f also known (observed or inferred)

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### The problem

- Identification: find "the best" model of the data in a family of alternatives
- Typical formulation: optimization of a (problem-dependent) cost function

minimize  $C(\theta|\mathcal{D})$  with respect to  $\theta \in \Theta$ 

- Cost function describes the ability of a model to explain the data
  - Minimization of the data fitting error, then validation
  - Penalization of overly complicated models to avoid overfitting
- In general, cost function is non-convex
  - Non-uniqueness of the solution
  - Optimization heuristics are needed



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#### Synthetic gene regulatory network in Yeast (Cantone et al., Cell 2009)

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## Linearization methods: Steady state

- Working assumption:
  - all concentrations converge to an equilibrium
  - small, fixed perturbations modify the system equilibrium
  - perturbations are known, equilibria can be measured



- What perturbations ?
  - Changes in concentration of chemicals in the medium
  - Gene knockout/overexpression
- Idea: Infer local dynamics around unperturbed equilibrium from several known perturbations of the system

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### Linearized dynamics

• True dynamics without perturbation

$$\dot{x} = \phi(x, u), \qquad u(t) \equiv 0 \text{ implies } x(t) \rightarrow x^*$$

• Linearization about equilibrium

$$\frac{d}{dt}(x-x^*)=\phi(x,u)=D_x\phi(x^*)(x-x^*)+D_u\phi(x^*)u+\mathsf{h.o.t.}$$

• Perturbed equilibria

$$u(t) \equiv \overline{u} \text{ implies } x(t) \rightarrow x^* + \overline{x}, \text{ where }$$

$$0 = D_x \phi(x^*)(x^* + ar{x} - x^*) + D_u \phi(x^*)ar{u} + h.o.t \simeq Aar{x} + Bar{u}$$





### Identification of linearized model

• Perform repeated perturbation experiments until equilibrium

$$u(t) \equiv \overline{u}_m \text{ implies } x(t) \rightarrow \overline{x}_m, \quad m = 1, \dots, M$$

• Collect observed results in data matrices

$$U = [\bar{u}_1, \dots, \bar{u}_M], \quad Y = [\bar{y}_1, \dots, \bar{y}_M], \text{ where } \bar{y}_m = \bar{x}_m + e_m$$

• Solve the least-squares problem

minimize ||AY + BU|| with respect to A

Solution well defined if B known and M large enough





#### Discussion

- A is network regulation matrix, B is (known?) perturbation effect A<sub>i,j</sub> > 0 gene j induces expression of gene i (x<sub>j</sub> ↑ ⇒ x<sub>i</sub> ↑)
  A<sub>i,j</sub> < 0 gene j inhibits expression of gene i (x<sub>j</sub> ↑ ⇒ x<sub>i</sub> ↓)
  A<sub>i,j</sub> = 0 gene j is not affected by gene i (x<sub>j</sub> indep. of x<sub>i</sub>)
- Explicit solution (Frobenius norm):

$$\widehat{A} = BUY^T (YY^T)^{-1}$$

warning: no zero elements (Overfitting !)

• Penalization of complexity: several strategies, e.g. "the Lasso":

$$\begin{array}{ll} \min & ||AY + BU|| & \min & \sum_{i,j} |A_{i,j}| \\ \text{s.t.} & \sum_{j} \mathbf{1}(A_{i,j} \neq 0) \le n_{max} \ \forall i & \text{s.t.} & ||AY + BU|| \le \epsilon \end{array}$$

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• Assumes linear dynamics (system evolving near equilibrium)

$$\frac{d}{dt}(x-x^*) = A(x-x^*) + Bu$$

- Allows for time-dependent (small) perturbation experiments
- Attempts to solve the equation

$$\dot{Y} = AY + BU$$

with the following time-course data (from a single experiment)

- $Y = [y(t_1), \dots, y(t_K)], U = [u(t_1), \dots, u(t_K)], \quad y(t_k) = x(t_k) x^* + e_k$
- In practice derivatives not known, resort to discretized dynamics

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### Identification from time-series

• Discretized linear dynamics (uniform measurement sampling)

$$x(t_{k+1}) = A^d x(t_k) + B^d u(t_k)$$

Solution of the approximate equality

$$Y^{+} = \begin{bmatrix} A^{d} & B^{d} \end{bmatrix} \begin{bmatrix} Y^{-} \\ U \end{bmatrix}, \qquad \qquad Y^{+} = \\ Y^{-} = \\ U = \end{bmatrix}$$

$$Y^+ = [y(t_2), \dots y(t_K)],$$
  
 $Y^- = [y(t_1), \dots y(t_{K-1})],$   
 $U = [u(t_1), \dots, u(t_{K-1})]$ 

- Also identifies perturbation matrix
- Regularized solution via Principal Component Analysis (PCA)
- Conversion to continuous-time network parameters

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## Identification of IRMA via TSNI: Results







ODE NETWORK INFERENCE (NIR & TSNI)



#### Figure 5. Reverse Engineering of the IRMA Gene Network from Steady-State and Time Series Experimental Data Using the ODE-Based Approach

The true network shows the regulatory interactions among genes in IRMA. Dashed lines represent protein-protein interactions. Directed edges with an arrow end represent activation, whereas a dash end represents inhibition.

(A and B) Inferred network using the TSNI reverse-engineering algorithm and the switch-on and switch-off time series experiments. Solid gray lines represent inferred interactions that are not present in the real network, or that have the wrong direction (FP, false positive). PPV [Positive Predictive Value = TP/(TP + FP)] and Se [Sensitivity = TP/(TP + FN)] values show the performance of the algorithm for an unsigned directed graph. TP, true positive; FN, false negative. The random PPV for the unsigned directed graph is equal to 0.40.

(C and D) Inferred network using the NIR reverse-engineering algorithm and the steady-state experimental data from network gene overexpression in cells grown in galactose or glucose medium, respectively.

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#### **Experimental validation: Example**

Perturbed Gene
 Perturbation Data

Empty Vector Data



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#### Discussion

- Returns a map of interactions and interaction strengths around nominal conditions
- In practice, linear assumption can be limiting:
  - Many interesting behaviors (e.g. switching) are inherently nonlinear
  - To observe these, experiments "excite" nonlinear system dynamics
- Still, generalizations of the linear model (e.g. piecewise affine models, as we will see) can be interesting

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#### **Boolean-like methods**

- Quantitative nonlinear modelling preserving the network "logics"
- Recall Boolean update map:

$$X_i^+ = b_i(X), \quad$$
 where  $b_i = igvee_I igwedge_j X_{I,j}', \quad X_{I,j}' \in \{X_j, 
eg X_j\}$ 

• Algebraic equivalent (Plahte et al, 1998): apply the transformation  $s^+(x_j) = rac{x_j^d}{x_j^d + \eta^d}$ 

$$egin{aligned} X_j &
ightarrow \sigma^+(x_j) \ &
expr(X) &
ightarrow 1 - expr(x) \ expr(X) \wedge expr'(X) &
ightarrow expr(x) \cdot expr'(x) \end{aligned}$$

Boolean-like model: define ODE

$$\dot{x}_i = \kappa_{0,i} + \kappa_{1,i} b_i(x) - \gamma_i x_i$$

 $b_i(x)$  algebraic equivalent of  $b_i(X)$ 



 $s^{-}(x_i) = 1 - s^{+}(x_i)$ 

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 $s^+$ 

S

### Example (Boolean model)



#### Gene Expressed when

- G2 not expressed 1
- G1 expressed or G4 not expressed 2
- G4 expressed and G1 not expressed  $b_3(X) = X_4 \land \neg X_1$ 3
- 4 G2 expressed

#### **Boolean model**

$$b_1(X) = \neg X_2$$

$$b_2(X) = X_1 \vee \neg X_4$$

$$b_4(X)=X_2$$

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### Example (Boolean-like ODE)



#### Gene More active when

- 1 G2 low
- 2 G1 high or G4 low
- 3 G4 high and G1 low
- 4 G2 high

#### ODE model $b_1(x) = s^-(x_2)$ $b_2(x) = 1 - s^-(x_1) \cdot s^+(x_4)$ $b_3(x) = s^+(x_4) \cdot s^-(x_1)$ $b_4(x) = s^+(x_2)$





## Plausibility ?

- Experimental evidence that often (Gjuvsland et al, 2007)
  - Transcription factors combine into Boolean-like input functions
  - Sigmoidal functions relate transcription factor concentrations and transcription rates
  - Post-transcriptional, transport, (and reaction) processes at equilibrium can be described by sigmoidal functions
- Still a phenomenological framework, but ...
  - Easy to interpret biologically
  - Quite accurate and flexible

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# Tractability ?

• General Boolean-like model:

$$\dot{x}_i = \kappa_i^1 + \kappa_i^2 b_i(x) - \gamma_i x_i$$
, where  $b_i = \sum_I \prod_j s^{\pm}(x_j | \theta_{I,j})$ 

- Structure identification: based on data, decide
  - The number of summands
  - The sigmoids in each product
  - The signs of the sigmoids
- ... combinatorial explosion and identifiability issues !!
- Parameter identification: parameters of each sigmoid, rates
- Intractable problem. But, good starting point
  - Structured expression
  - Reduction to specific families of Boolean-like functions
  - Approximation

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#### **Piecewise Affine models**

• Idea: abstract nonlinearities sigmoids by hard thresholds (switches)



- Dynamical models with Boolean-type events
- Coarse approximation, but when applicable, powerful analysis (de Jong et al. 2004) & identification (Porreca et al, 2009) tools!

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#### Example: double-inhibition network

Courtesy of G.Ferrari-Trecate (apologies for notational changes...)



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#### PWA models: key features

• thresholds split  $\Omega$  into hyperrectangular domains  $\Delta^1, \Delta^2, \ldots$ :



switching thresholds and rate parameters define the interactions





• interaction: activation/inhibition based on changes in  $\kappa_i$ 





#### PWA models: key features cont'd

decoupling  $\Rightarrow$  local 1<sup>st</sup> order dynamics for each concentration: if no switches occur over  $[t_{k_0}, t_{k_1}]$  there exist  $\kappa \ge 0, \gamma > 0$ such that





Data can be split in *segments*  $S_j$ generated by rate parameters  $(\kappa^j, \gamma)$ 



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#### Data segmentation and classification

 $y_i(t_k) = x_i(t_k) + e_k,$  Given one time series  $i=1,\ldots,n$ Variable sampling time  $e_k \sim \mathcal{N}(0, \sigma^2)$  $k = 1, \ldots, K$ Extends to multiple time series Use statistical procedures to Find segments with exponential behavior in each concentration profile (fit parameters and check that fitting residuals are compatible with noise) Partition data into sets with the same experiential model  $x_2$  $\mathcal{F}_1$  $\mathcal{F}_{A}$  $\mathcal{F}_3$  $\mathcal{F}_2$ 

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#### Threshold reconstruction

Find *minimal* sets of thresholds that separate data clusters (multicuts)

- Find all thresholds that separate two clusters
- Define and exploit partial order relations among multicuts to find the minimal ones
- Combinatorial number of multicuts: exploit branch-and-bound optimization



#### **Optimal models**

- Search of minimal multicuts: complexity reduction
- Identifiability issues:
  - Cannot discriminate certain models on the basis of the data (pool of equivalent models providing alternative biological hypotheses)
  - Cannot fix thresholds, only bounds can be established



Three minimal multicuts = three possible GRNs

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#### Example: carbon starvation in E.coli



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#### Model and simulation

#### (Ropers et al., Biosystems, 2006)



### Identification from simulated data

Cut #	Variable	Threshold value	Interaction	Correct? (Y/N)
1	CRP	0.61	activator of the synthesis of Fis	Ν
2	CRP	0.64	activator of the synthesis of Fis	Ν
3	CRP	0.71	inhibitor of the synthesis of Stable RNAs	N
4	CRP	0.74	inhibitor of the synthesis of Fis	Ν
5	Fis	0.09	inhibitor of the synthesis of CRP	Y
6	Fis	0.23	activator of the synthesis of Fis	Ν
7	Fis	0.49	activator of the synthesis of Stable RNAs	Y
8	Fis	0.75	inhibite of the synthesis of GyrAB	
9	GyrAB	0.48	activation of the sonther is c Fig	N
10	GyrAB	0.50	activator of the synthesis of Fis	Y
11	GyrAB	0.55	inhibitor of the synthesis of Stable RNAs	N
12	GyrAB	0.67	activator of the synthesis of CRP	N
13	Stable RNAs	0.04	inhibitor of the synthesis of Fis, activator of the synthesis of Stable RNAs	N
14	Stable RNAs	0.18	inhibitor of the synthesis of CRP	Ν
15	Stable RNAs	0.53	inhibitor of the synthesis of Fis	Ν
16	Stable RNAs	0.55	activator of the synthesis of GyrAB	N
17	Stable RNAs	0.64	inhibitor of the synthesis of Fis, activator of the synthesis of Stable RNAs	N
18	Signal	0.50	inhibitor of the synthesis of Fis	Y

#### Minimal multicuts found

	Multicut composed of cuts #:	Correct? (Y/N)
	1, 5, 7, 8, 10	N, Y, Y, Y, Y
•	1, 7, 8, 10, 12	N, Y, Y, Y, N
	5, 7, 8, 10, 18	Y, Y, Y, Y, Y
	7, 8, 10, 12, 18	Y, Y, Y, N, Y
	7, 8, 10, 14, 18	Y, Y, Y, N, Y

the best minimal multicut captures all interactions that are identifiable from the data

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#### Discussion

- •Under the working hypotheses, effective method for structural and parametric identification of network dynamics
- •Neat framework for identifiability and system analysis
- Hardly compatible with soft nonlinearities
  - Requires generalization
  - Some concepts (invalidation of structural hypotheses) can be transported to certain smooth nonlinear models





#### Models with unate structure

- Framework for systematic selection of plausible quantitative models
- Unate functions: Boolean rules monotone in each input variable
  - Transcription factors with unambiguous role on every given gene
  - Most known rules (only experimentally observable rules? ↔ identifiability)
- Boolean-like ODE model: preserves monotonicity properties

• Model:  
$$b_i(x) = \prod_{l=1}^{n_i} \tau_l, \quad \tau_l = 1 - \prod_{j \in J_l} (1 - s^{\pm}(x_j)) \quad \text{where} \quad s^{\pm}(x_j) = \begin{cases} s^+(x_j), & \text{or} \\ s^-(x_j), \end{cases}$$

• Sign pattern:

$$p = (p_1, \dots p_n), \qquad p_j = \begin{cases} 1, & \text{if } s^{\pm}(x_j) = s^+(x_j), \\ -1 & \text{if } s^{\pm}(x_j) = s^-(x_j), \quad j = 1, \dots, n \\ 0 & \text{if } j \notin J_l \ \forall l \end{cases}$$

Example, p = (-1, 1):  $s^{-}(x_1)s^{+}(x_2)$ ,  $1 - s^{+}(x_1)s^{-}(x_2)$ ,  $s^{-}(x_1)(1 - s^{+}(x_1)s^{-}(x_2))$ , ...

b(x) is nondecreasing (resp. nonincreasing) in  $x_j$  if  $p_j = 1$  (resp.  $p_j = -1$ ) ... and so is any synthesis rate  $g_i(x) = \kappa_{0,i} + \kappa_{1,i}b_i(x)$ , provided  $\kappa_{0,i}, \kappa_{1,i} \ge 0$ 



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# Sign patterns: definitions and properties

- Given data pairs:  $(x^1, g^1), \ldots, (x^m, g^m)$ , with  $g^k = g(x^k|p)$
- Definition: p is *inconsistent* if the property

$$p_j(x_j^k - x_j^l) \ge 0, \ j = 1, \ldots, n \implies g(x^k|p) - g(x^l|p) \ge 0$$

is falsified for some k,l

• Definition: subpattern and superpattern

11-11 1 1 -1 -1 1 -1 -1 1 1 -1 -1 -1 4 Superpatterns 11-10 1 -1 -1 0 3 Pattern 10 - 102 00-10 1000 1 Subpatterns 0000 Ω

- Subpatterns of inconsistent patterns are also inconsistent
- Superpatterns of consistent patterns are also consistent
- Minimal consistent and maximal inconsistent patterns exist

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Complexity

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Example. g(x|p),  $x = (x_1, x_2)$ , unknown  $p = (p_1, p_2)$ . Given  $(x, g_i)$ ,  $(x', g'_i)$  with  $x_1 > x'_1$ ,  $x_2 < x'_2$ ,  $g_i > g'_i$ . Can exclude:  $p = (-1, 1) = (\text{sign}(x'_1 - x_1), \text{sign}(x'_2 - x_2))$ . Can also exclude: p = (0, 1), p = (-1, 0), p = (0, 0). Note: Parameter values play no role here!

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# Identification via sign patterns: rationale

Given protein concentrations & synthesis rates: (recall  $\dot{x}_i = g_i(x) - \gamma_i(x)$ )

- Step 1: Exploit monotonicity properties to invalidate sign patterns
  - Extract invalid sign patterns from data
  - Infer the set of minimal consistent sign patterns

Next, given candidate unate model structures S(p) for every p:

- Step 2: Search best fitting model structure with valid sign pattern
  - Enumerate valid sign patterns p of increasing level of complexity
  - For every valid p at the current level of complexity, fit (the parameters of) every model in S(p) to the data
  - Return all models that pass a statistical test on the fitting residuals. If none, go to next complexity level.

In practice, S(p) shall be a subset of unate models with pattern p

- Exploitation of a priori knowledge
- Computational limitations

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# Algorithm 1: original version (full data)

- Protein concentrations & synthesis rates
- Time-course noisy data, known variance:

$$\tilde{x}_{i}^{k} = x_{i}^{k} + e_{i}^{k} \qquad \tilde{g}_{i}^{k} = g_{i}^{k} + \epsilon_{i}^{k}$$
$$x_{i}^{k} = x_{i}(t_{k}) \qquad g_{i}^{k} = g(x(t_{k}))$$

with  $k = 1, \ldots, m$  and zero-mean Gaussian noise

$$v_e(x_i^k) = \operatorname{var}(e_i^k) \quad v_\epsilon(g_i^k) = \operatorname{var}(\epsilon_i^k)$$

Computation of  $\overline{P}$ : set  $\overline{P} = \emptyset$ . For all indices  $k, l \in \{1, ..., m\}$ :

If g<sup>k</sup>−g<sup>l</sup> <0, define the sign pattern p
=(p
<sub>1</sub>,...,p
<sub>n</sub>) by setting p
<sub>j</sub> = sign(x<sub>i</sub><sup>k</sup>−x<sub>i</sub><sup>l</sup>), with j=1,...,n, and include p
 in P

Computation of  $P^*$ : define  $\overline{\ell} = \max\{C(\overline{p}): \overline{p} \in \overline{P}\}$ . Initialize  $P^* = \emptyset$ . For increasing values of complexity  $\ell = 0, ..., \min\{n, \overline{\ell} + 1\}$ :

- (II) Generate all patterns p of complexity  $\ell$ . For each such p,
- (III) Check if p is consistent by verifying that there is no  $\bar{p} \in \bar{P}$  such that  $p \sqsubseteq \bar{p}$ . If this is the case,
- (IV) Check if p is minimal consistent by verifying that there is no  $p^* \in P^*$  such that  $p^* \sqsubseteq p$ . If this is the case, include p in  $P^*$ .

Algorithm 1 Two-step identification.

- Step 1. (Selection of consistent model structures)
  - I. Set  $\bar{P} = \emptyset$ . For all indices  $k, l \in \{1, ..., m\}$ , if  $\tilde{g}_i^k \tilde{g}_i^l < -N\sigma_{g_i}^{k,l}$ then define  $\bar{p} = (\bar{p}_1, ..., \bar{p}_n)$  by

$$\bar{p}_{j} = \begin{cases} -1, & \text{if } \tilde{x}_{j}^{k} - \tilde{x}_{j}^{l} \le -N\sigma_{x_{j}}^{k,l}, \\ 1, & \text{if } \tilde{x}_{j}^{k} - \tilde{x}_{j}^{l} \ge N\sigma_{x_{j}}^{k,l}, \\ 0, & \text{otherwise}, \end{cases} \quad j = 1, \dots, n,$$

and include  $\bar{p}$  in  $\bar{P}$ .

II–IV. Execute the computation of  $P^*$  from the resulting  $\overline{P}$ , as described in Section 2.2.

Step 2. (Identification of best consistent models) Set  $\mathscr{P} = \emptyset$ . Define  $\ell^* = \min\{C(p^*): p^* \in P^*\}$ . For  $\ell = \ell^*$  to *n*:

- V. Generate patterns p such that  $C(p) = \ell$  and  $p^* \sqsubseteq p$  for some  $p^* \in P^*$ . For each such p, execute VI.
- VI. For all  $s \in S(p)$ , fit the model  $g_i(\cdot)$  with sign pattern p and structure s by solving the nonlinear regression problem

$$\delta = \min_{\theta} \sum_{k=1}^{m} w_k \left( \tilde{g}_i^k - g_i(\tilde{x}^k) \right)^2. \tag{8}$$

If  $\delta < \tau(\alpha)$ , include the fitted model in  $\mathscr{P}$ . VII. If  $\mathscr{P} \neq \emptyset$  return  $\mathscr{P}$  and exit.

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#### Comments

- Separate identification of regulation function of each gene
- Practicality requires use of a sub-hierarchy of unate structures
- Hierarchical search of model structures of increasing complexity
  - Stops when a good model is found (statistical test on the model residuals)
  - Favors simple over complicated models
  - Returns pool of biological alternatives
- What is a statistically good model?
  - Under the null hypothesis that the estimated model is correct, the fitting residual is distributed as  $\chi^2(m)$
  - Use this property to define confidence levels (threshold on the fitting residuals) on the model estimate
- Limitations: Nonconvex parameter fitting, Data requirements





# Case study: unate models with canalizing structure



- Goal: use a priori knowledge to reduce the family of network structures
- Intuition: many Boolean expression rules are unlikely/uncommon
- Evidence: (Szallasi et al 1998, Kauffman et al 2004, ...) out of 139 gene activation rules analyzed in (Harris et al., 2002), 99% are "Canalizing Functions", 95% are HD rarchi ally Canalizing Functions", 90% are "Ho U H1"
  - CFs: at least one (canalizing) value of at least one (canalizing) variable determines the value of the function
  - HCFs: when the canalizing variable takes its non-canalizing value, a second variable is canalizing, etc.

HCF

H0 U H1)) Unate

#### We focus on Ho U H1

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CF

#### The class H<sub>0</sub> U H<sub>1</sub>

• Class H<sub>0</sub>: 
$$b_i(X) = X'_{j_1} \land X'_{j_2} \land \cdots \land X'_{j_\ell}$$
  $X'_{l,j} \in \{X_j, \neg X_j\}$ 

• Class H<sub>1</sub>: 
$$b_i(X) = X'_{j_1} \wedge X'_{j_2} \wedge \cdots \wedge X'_{j_{\ell-2}} \wedge (X'_{j_{\ell-1}} \vee X'_{j_\ell})$$

• Boolean-like ODE model with H<sub>0</sub> U H<sub>1</sub>-structure:

$$egin{aligned} \dot{x}_i &= \kappa_i^1 + \kappa_i^2 b_i(x) - \gamma_i x_i \ b_i(x) &= egin{cases} s^{\pm}(x_{j_1}) \cdot s^{\pm}(x_{j_2}) \cdots s^{\pm}(x_{j_\ell}) \ s^{\pm}(x_{j_1}) \cdot s^{\pm}(x_{j_2}) \cdots s^{\pm}(x_{j_{\ell-2}}) ig(1 - s^{\mp}(x_{j_{\ell-1}}) \cdot s^{\mp}(x_{j_\ell})ig) \end{aligned}$$

Structure:  $\ell$ ,  $(j_1, j_2, ..., j_{\ell})$ ,  $H_0$  vs.  $H_1$ 

Parameters:  $\kappa_i^1$ ,  $\kappa_i^2$ , sigmoids' parameters (threshold, cooperativity)



### Identification of Ho U H1 models

• Given concentration and synthesis rate measurements

$$y_i(t_k) = x_i(t_k)(1 + e_{i,k}) \qquad z_i(t_k) = f_i(x(t_k))(1 + \epsilon_{i,k}) \qquad i = 1, \dots, n$$
  
$$e_{i,k} \sim \mathcal{N}(0, \sigma_e^2) \qquad \epsilon_{i,k} \sim \mathcal{N}(0, \sigma_\epsilon^2) \qquad k = 1, \dots, K$$

• For known degradation rate, can compute synthesis rates from x:

$$f_i(x) = \kappa_i^1 + \kappa_i^2 b_i(x) = \dot{x}_i + \gamma_i x_i$$
 (Ronen et al 2002, Brown et al 2008,...)

#### • Estimate

- Structure:  $\ell, (j_1, j_2, ..., j_\ell), H_0$  vs.  $H_1$
- Parameters:  $\kappa_i^1$ ,  $\kappa_i^2$ ,  $\theta_j$  (possibly depending on *i*)

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#### **Performance results**

We attempted identification of this system with 90 equally spaced data points over a time interval such that the product concentrations of the core genes complete three full oscillations. Measurements  $\tilde{x}_i^k$  and  $\tilde{g}_i^k$  were artificially corrupted by Gaussian noise samples according to the observation model (7), with  $v_e(x_i^k) =$  $(\sigma_e x_i^k)^2$  and  $v_{\epsilon}(g_i^k) = (\sigma_{\epsilon} g_i^k)^2$ , for the different noise levels  $\sigma_e =$  $\sigma_{\epsilon} = 0.01, 0.03, 0.05, 0.07$ . This corresponds to noise roughly within 3%, 10%, 15% and 20% of the actual values of  $x_i^k$  and  $g_i^k$ . The performance of Algorithm 1 (with N=6 and  $\alpha=0.95$ ) for the various noise levels and all genes is conveyed by the scores on the performance indices R, S, A and D (Table 1). These ware computed as described in Section 2.3.4 on the basis of dentificat on runs with the same system evolution, but with different random outcomes of the noise. Each run (MATLAB V.7 R.14) took on an average roughly 5 min on a Windows XP workstation with Pentium 3.20 GHz processor and 2.00 GB RAM. Computational time ranged from  $\sim 2$  s for the identification of  $g_3$  to  $\sim 4$  min for the identification of g<sub>6</sub>. Step 1 always performs very reliably, i.e. index R is constantly

	Index	Range
Stop 1	R eliability	[0,1]
Step I	S electivity	[0,1]
Stop 2	A ccuracy	[0,1]
Step 2	D ispersion	≥1

#### Description

Probability that the true p is deemed consistent Percentage of sign patterns eliminated from the search in Step 2 Probability that the true structure is In the pool of identified models Average number of models in the pool



		$\sigma_{\ell}, \sigma_{\epsilon}$	0.01	0.03	0.05	0.07
Gene 1	Step 1	R S	1 0.92	1 0.92	1 0.92	1 0.91
	Step 2	A D	0.90 1	0.92 1	0.91 1	0.89 1
Gene 2	Step 1	R S	1 0.92	1 0.92	1 0.92	1 0.91
	Step 2	A D	0.93 1	0.92 1	0.89 1	0.89 1
Gene 3	Step 1	R S	1 0.92	1 0.92	1 0.92	1 0.92
	Step 2	A D	0.93 1	0.93 1	0.93 1	0.92 1
	Step 1	R S	1 0.94	1	1 0.87	1
Gene 4	Step 2	A D	0.94 1	0.94 1	0.93 1.02	0.89 1.44
Gene 5	Sep 1	R	1	1	1	1
	Step 2	A D	0.95	0.94 1	0.91 1.79	0.83
	Step 1	R	1	1	1	1
Gene 6	Step 2	A D	0.89 1	0.92 1.02	0.85 2.76	0.42 2.74

#### Simulated identification on E.coli model Signal 6-gene carbon starvation CRP activation → GyrAB response network → Fis Supercoiling Model in exponential growth phase crp → TopA Stable RNAs topA All but third equation rrn have Ho U H1-structure ilators and regulatory interactions takbal reg (all have unate structure) place during the transmon from stationary to exponential growth phase in E. Coli. $\dot{x}_1 = \kappa_1^1 + \kappa_1^2 - \gamma_1 x_1$ (Ropers et al, Biosystems 2006) $\dot{x}_2 = \kappa_2^1 + \kappa_2^3 \sigma^-(x_3) - \gamma_2 x_2$ $\dot{x}_3 = \kappa_3^1 \sigma^-(x_3) + \kappa_3^2 \sigma^+(x_4) \sigma^-(x_5) \sigma^-(x_3) - \gamma_3 x_3$ $\dot{x}_4 = \kappa_4 (1 - \sigma^+(x_4) \sigma^-(x_5)) \sigma^-(x_3) - \gamma_4 x_4$ $x_1, x_2, x_3, x_4, x_5, x_6 =$ $\dot{x}_5 = \kappa_5 \ \sigma^+(x_4) \ \sigma^-(x_5) \ \sigma^+(x_3) - \gamma_5 \ x_5$ Cya, CRP, Fis, GyrAB, TopA Stable RNAs $\dot{x}_6 = \kappa_6^1 \sigma^+(x_3) + \kappa_6^2 - \gamma_6 x_6$

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#### Identification scenario



- Simulated data collected every 10 min
- Measurements over 1200 min
- Various noise levels
- Performance from 100 simulated runs
   Realistic parameters and initial cond.
  - Dynamics excited in the experiment:
  - $g_1 = \kappa_{0,1}, \qquad g_4 \simeq \kappa_{1,4} \sigma^-(x_4) \sigma^-(x_3),$  $g_2 = \kappa_{0,2} + \kappa_{1,2} \sigma^-(x_3), \qquad g_5 \simeq \kappa_{1,5} \sigma^+(x_4) \sigma^+(x_3),$
  - $g_3 \simeq \kappa_{0,3} + \kappa_{1,3}\sigma^+(x_4)\sigma^-(x_3), \ g_6 = \kappa_{0,6} + \kappa_{1,6}\sigma^+(x_3).$
- All excited dynamics have H<sub>0</sub> U H<sub>1</sub>structure

Use this as a "reference" model

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#### Results on *E.coli*

Note that the expression of gene 1 obeys trivial dynamics. Correspondingly, a constant model for  $q_1$  is returned by the preprocessing Step 0 in roughly 95% of the runs. This is summarized by the accuracy index A. In the remaining runs the algorithm rules out the constant model, i.e. the true pattern is not in the patterns deemed consistent and a model with correct structure cannot be found in Step 2. For the remaining genes, the values of reliability R and selectivity S witness that Step 1 is still very effective and robust to noise. Step 2 includes the correct model structure in a small pool of identified models in all cases with a moderate performance decay at increased noise levels. For gene 4 only, this duc 1pt when the noise level raises above 5% ( $\sigma_e = \sigma_{\epsilon}$ ) (0), possibly due to a limited excitation of the expression dynamics. Finally, for gene 5, the limited accuracy of Step 2 (A = 0.14) at the lowest noise level is due to convergence to local minima in the solution of the nonconvex optimization (8). With low noise, the local minima are more pronounced and the solver currently used cannot escape them. This limitation could be ameliorated by a randomized optimization strategy ([28]). To conclude we mention that, whenever the identifiable model structure was estimated correctly, the corresponding parameter estimates were generally accurate (best accuracy being obtained with lowest noise, results not shown).

		$\sigma_e, \sigma_e$	0.01	0.03	0.05	0.07
	Step 1	R S	_	_	_	_
Gene 1	Step 2	A D	0.95	0.95 -	0.96	0.95
	Step 1	R	1	1	1	1
Gene 2	Step 2	A D	0.98	0.97	0.95	0.94
Con	Step 1	R S	1 0.81	$\begin{array}{c}1\\0.58\end{array}$	$\begin{array}{c}1\\0.54\end{array}$	$\begin{array}{c} 1 \\ 0.50 \end{array}$
Genes	Step 2	A D	0.95 1	0.93 1.39	$0.87 \\ 2.47$	$0.58 \\ 2.84$
Canad	Step 1	$R \\ S$	$\begin{array}{c}1\\0.60\end{array}$	$\begin{array}{c}1\\0.50\end{array}$	$\begin{array}{c}1\\0.44\end{array}$	$\begin{array}{c} 1 \\ 0.37 \end{array}$
Gene 4	Stop 9	A	0.93	0.16	0	0
	Step 2	D	1.24	4.31	_	-
Cana 5	Step 2	D R S	1.24 1 0.73	4.31 1 0.66	1 0.61	1 0.54
Gene 5	Step 2 Step 1 Step 2	D R S A D	1.24 1 0.73 0.14 1	4.31 1 0.66 0.84 1	- 1 0.61 0.88 1	- 1 0.54 0.79 1
Gene 5	Step 2 Step 1 Step 2 Step 1	D R S A D R S	1.24 1 0.73 0.14 1 0.75	4.31 1 0.66 0.84 1 1 0.67	1 0.61 0.88 1 0.64	

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## Algorithm 2: extension to partial data

- Assuming only protein concentrations are available:
  - 1. Reconstruct missing information (synthesis rates, variances)
  - 2. Apply Algorithm 1 (unchanged)
- Option 1: Deconvolution

 $\dot{x}_i(t) = -\gamma_i x_i(t) + g_i(t), \quad g_i(t) = \kappa_{0,i} + \kappa_{1,i} b_i(x(t))$  is a forcing input

- Well established (Bayesian) methods for regularized estimates
- Severe over- and under-smoothing observed in practice
- Option 2 (our choice): Pata fitting + Boots rapping

Choose basis functions for  $x_i(\cdot)$ , e.g. cubic splines

Compute estimate  $\hat{x}_i$  by fitting data  $\tilde{x}_i^k$ , and  $\hat{x}_i = \dot{\hat{x}}_i$  by explicit differentiation

Reconstruct the synthesis rates  $\tilde{g}_i^k = \dot{\hat{x}}_i(t_k) + \gamma_i \tilde{x}_i^k$ 

Utilize the fitting errors  $\tilde{x}_i^k - \hat{x}_i(t_k)$  to reproduce the noise statistics

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#### **Residual resampling**

- Randomized procedure to infer statistics of any functional of the regression curve
- Applicable to any type of regression curve (But sensitive to this choice!)



Algorithm 2 Bootstrap spline-based resampling.

1: compute the spline 
$$\hat{x}_i(t)$$
 from  $\{\tilde{x}_i^k\}$  using weights  $\{w^k\}$   
2: let  $R = \{w^k(\tilde{x}_i^k - \hat{x}_i(t_k)), k = 1, \dots, m\}$ 

3: for 
$$r = 1$$
 to  $N_r$  do

- extract with replacement m residuals  $\{\varepsilon^k\}$  from R 4:
- 5:
- let  $\tilde{x}_i^{k(r)} = \hat{x}_i(t_k) + \varepsilon^k / w^k, k = 1, \dots, m$ compute the spline  $\hat{x}_i^{(r)}(t)$  from  $\{\tilde{x}_i^{k(r)}\}$  using weights  $\{w^k\}$ 6:

7: let 
$$\hat{g}_i^{\kappa(r)} = \hat{x}_i^{(r)}(t_k) + \gamma_i \hat{x}_i^{(r)}(t_k), k = 1, \dots, m$$

9: let 
$$\hat{g}_i^k = \frac{1}{N_r} \sum_r \hat{g}_i^{k(r)}, \ \hat{v}_\epsilon(g_i^k) = \frac{1}{N_r - 1} \sum_r (\hat{g}_i^k - \hat{g}_i^{k(r)})^2$$
 and  $\hat{v}_\epsilon(x_i^k) = \frac{1}{m-1} \sum_{\varepsilon \in R} (\varepsilon/w^k)^2$ 







**Experiment on IRMA** 

Synthetic gene network in Yeast (Cantone et al., Cell 2009)





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#### Mathematical model

Letting  $[CBF1] = x_1$ ;  $[GAL4] = x_2$ ;  $[SWI5] = x_3$ ;  $[GAL80] = x_4$ ;  $[ASH1] = x_5$ , (Cantone *et al.*, Cell 2009) the evolution of the mRNAs concentrations were modelled as follows:

$$\frac{dx_1}{dt} = \alpha_1 + v_1 \left( \frac{x_3^{h_1}(t-\tau)}{(k_1^{h_1} + x_3^{h_1}(t-\tau)) \cdot \left(1 + \frac{x_5^{h_2}}{k_2^{h_2}}\right)} \right) - d_1 x_1, \tag{1}$$

$$\frac{dx_2}{dt} = \alpha_2 + v_2 \left( \frac{x_1^{h_3}}{k_3^{h_3} + x_1^{h_3}} \right) - (d_2 - \Delta(\beta_1))x_2, \tag{2}$$

$$\frac{dx_3}{dt} = \alpha_3 + \widehat{v}_3 \left( \frac{x_2^{h_4}}{\widehat{k_4}^{h_4} + x_2^{h_4}(1 + \frac{x_4^4}{\widehat{\gamma}^4})} \right) - d_3 x_3, \tag{3}$$

$$\frac{dx_4}{dt} = \alpha_4 + v_4 \left( \frac{x_3^{h_5}}{k_5^{h_5} + x_3^{h_5}} \right) - (d_4 - \Delta(\beta_2)) x_4,$$

$$\frac{dx_5}{dt} = \alpha_5 + v_5 \left( \frac{x_3^{h_6}}{k_6^{h_6} + x_3^{h_6}} \right) - d_5 x_5,$$
(5)

• We attempt identification in the class of models with  $H_0 \cup H_1$ -structure

- Different but similar analytical form
- Test for flexibility of the approach
- Known delays can be accounted for

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#### **Results: full data**

- Comparison with TSNI (Cantone et al., Cell 2009)
- True protein concentrations (very few data points)
- Rates simulated from the model ("what-if" performance test)
- Evaluation of network reconstruction performance, but not of parameter fit
- PPV=TD/TD+FD and Se=TD/TD+FU (T=True, D=Detected, U=Undetected edges)



Fig. 1. (a) True network of interactions in IRMA. Results obtained by (b) the TSNI algorithm (Cantone *et al.*, 2009) and by (c) Algorithm 1. Grey arcs (respectively, grey-end markers) denote incorrect direction (respectively, sign) of the inferred interactions. Values of PPV and Se for the signed directed graph, when different from the unsigned case, appear in square brackets. The three values of Se in (c) refer to increasing noise levels, while dashed and dotted arcs denote interactions inferred only for  $\sigma_{\epsilon} < 0.3$  and  $\sigma_{\epsilon} < 0.1$ , respectively.

#### Porreca et al, Bioinformatics 2010



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### **Results: partial data**



Fig. 1: (a) True network of interactions in IRMA. Results obtained by (b) the TSNI algorithm [27] and by (c) Algorithms 1 and 2. Gray edges denote incorrect direction of the inferred interactions.

- Additional assumptions (no self-regulation)
- Loss of accuracy
  - Parameter estimates (when applicable, not shown)
  - Sign of interaction (possibly due to low data quality)

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- Direction of regulation (bad!)
- Still better than TSNI...

#### To be compared with...



, INRIA

#### Identification of stochastic models: A quick view



### Introduction: stochastic gene expression

- At the cell level, protein synthesis depends on random events
  - Binding/unbinding of activators/repressors and RNApol to DNA, ...
  - Environmental conditions (temperature, availability of free RNAP,...)
- Classical stochastic gene expression model:
  - Describes the formation and degradation of single molecules
  - Time resolution, no spatial resolution (homogeneous reaction volume)



 $P[x_1 \text{ increases by } 1 \text{ in } \delta t] = \lambda_1 \delta t + o(\delta t)$  $P[x_1 \text{ decreases by } 1 \text{ in } \delta t] = \gamma_1 x_1 \delta t + o(\delta t)$   $\mathbf{x}_1 =$ number of mRNA molecules

 $\mathbf{x}_2 =$ number of protein molecules

 $\lambda_1, \lambda_2 = \text{prob.}$  of molecule formation per unit time  $\gamma_1, \gamma_2 = \text{prob.}$  of molecule degradation per unit time

 $P[x_2 \text{ increases by } 1 \text{ in } \delta t] = \lambda_2 x_1 \delta t + o(\delta t)$  $P[x_2 \text{ decreases by } 1 \text{ in } \delta t] = \gamma_2 x_2 \delta t + o(\delta t)$ 

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### **Regulation and noise**

Example: regulated gene expression and protein degradation



- This modelling framework describes the random nature of the events *internal* to the gene expression mechanism (*intrinsic noise*)
- Random fluctuations of the event rates, due to changes *external* to the gene expression mechanism, are not modelled (*extrinsic noise*)
   [Many contributors: Paulsson, Elowitz, Alon, Arkin, ...]



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#### Example: bistable switch

• Network that admits two distinct stable equilibria



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### Simulated probability of the state



(Convergence to different equilibria with different probabilities)





## **The Chemical Master Equation**

• In general, for any system described via a discrete-valued continuous-time Markov chain, the following holds:

$$\dot{p}(\mathbf{x};t) = \sum_{\mu=1}^{M} p(\mathbf{x} - s_{\mu};t) a_{\mu}(\mathbf{x} - s_{\mu}) - p(\mathbf{x};t) \cdot \sum_{\mu=1}^{M} a_{\mu}(\mathbf{x})$$

 $\mathbf{x}$  = random vector of the number of molecules of every species, one per entry  $\mu$  = reaction index (from 1 to M possible reactions)

$$s_{\mu}$$
 = state change associated to the  $\mu$ -th reaction

- $a_{\mu}$  = propensity (prob. per unit time) of  $\mu$ -th reaction (state dependent)
- Infinite-dimensional linear equation in the probabilities p
- In general, no closed-form but only approximate solutions
  - Stochastic simulation (also known as the Gillespie algorithm)
  - Analytical approximations (Langevin eq., Finite State Projection)

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# Identification of stochastic gene expression models from population snapshot data

• Consider a network with *given structure* and *unknown parameters* 

$$\dot{p}_{\theta,u}(\mathbf{x};t) = \sum_{\mu=1}^{M} p_{\theta,u}(\mathbf{x}-s_{\mu};t) a_{\mu;\theta}(\mathbf{x}-s_{\mu};u(t)) - p_{\theta,u}(\mathbf{x};t) \cdot \sum_{\mu=1}^{M} a_{\mu;\theta}(\mathbf{x};u(t))$$

• Assume that, for a given function h, the following data is available:

$$y^{m}(t_{k}) = h(p_{u^{m}}(\cdot, t_{k}), e_{k,m}), \qquad k = 1, \dots, K, \quad m = 1, \dots, M$$

• (Parameter) identification is typically formulated as the optimization

$$\hat{\theta} = \arg\min_{\theta \in \Theta} \sum_{k,m} D_{k,m} \Big( y^m(t_k), h\big( p_{\theta,u^m}(\cdot; t_k), \epsilon_{k,m} \big) \Big)$$

for suitable distance(s) / fitting cost(s)

• Several hypotheses on model structure can be tested based on fitting result (Khammash & van Oudenaarten)

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#### Population snapshot data: microfluidics

Fig. 1. Quantitative imaging of a YFP-fusion library. (A) Each library strain has a YFP translationally fused to the C terminus of a protein in its native chromosomal position. (B) A poly(dimethylsiloxane) (PDMS) microfluidic chip is used for imaging 96 library strains. E. coli cells of each strain are injected into separate lanes and immobilized on a polylysinecoated coverslip for automated fluorescence imaging with single-molecule sensitivity. (C to E) Representative fluorescence images overlaid on phase-contrast images of three library strains, with respective single-cellprotein level histograms that are fit to gamma distributions with parameters a and b. Protein levels are determined by deconvolution (18). The protein copy number per average cell volume, or the concentration, was determined as described in the main text and the SOM (18). (C) The cytoplasmic protein Adk uniformly distributed intracellularly. (D) The membrane protein AtpD distributed on the cell periphery. (E) The predicted DNAbinding protein YjiE with clear intercellular localization. Single YjiE-YFPs can be visualized because they are localized. Note that, unlike (C) and



(D), the gamma distribution asymmetrically peaks near zero if a is close to or less than unity.

#### [Taniguchi et al., Science 329, 533 (2010)]





## Example: Identification of *lac* operon in *E.coli* $_{9}^{\prime}$



**Figure 3** Experimental identification of a simple construct (**A**) in which IPTG induces the production of GFP. (**B**) Experimentally measured histograms of *gfp* expression on two different days (solid blue and green lines—in arbitrary units) and the best determined parameter fit (red dashed lines). Here, each column corresponds to a different measurement time (0, 3, 4, or 5 h) after induction, and each row corresponds to a different level of extracellular IPTG induction (5, 10, or  $20 \,\mu$ M). In the parameter fits, different weights were applied to each experimental condition, shown as the values {*q*} in the histograms. (**C**) Predicted (red) and then measured (blue and green) fluorescence at 40 and  $100 \,\mu$ M.

#### •Parametric Markov chain model:

$$[IPTG]_{IN} = [IPTG]_{OUT} \cdot (1 - \exp(-nt)),$$

$$R_1 : \phi \xrightarrow{w_1} LacI, \quad R_2 : LacI \xrightarrow{w_2} \phi, \qquad w_1 = k_L, \qquad w_2 = \delta_L \cdot [LacI], \quad \delta_L = \delta_L^{(0)} + \delta_L^{(1)} [IPTG]_{IN}.$$

$$R_3 : \phi \xrightarrow{w_3} GFP, \quad R_4 : GFP \xrightarrow{w_4} \phi \qquad w_3([LacI]) = \frac{k_G}{1 + \alpha [LacI]^{\eta}} \qquad w_4 = \delta_G \cdot [GFP],$$

•Parameter identification by matching fluorescence histograms

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Fitting ...

8  $\mathbf{\Lambda}^* := \operatorname{argmin}_{\mathbf{\Lambda}} \left\{ \sum_{i} q_i \cdot \left\| f_{\text{Meas}}^{(i)} - f_{\text{Tot}}^{(i)} \right\|_1 \right\}$ 0



Total florescence: GFP + background (arbitrary units)

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#### ... and validation



• But how are probability distributions computed from the model for changing inputs and parameter values ?

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## Solving the CME: Finite State Projection

[ The material on Finite State Projection in these slides is borrowed from

M.Khammash,"The Chemical Master Equation in Gene Networks: Complexity and Approaches"

available at:

http://www.cds.caltech.edu/~murray/wiki/images/d/d9/Khammash\_master-15aug06.pdf ]

• The state of the system evolves on a lattice

- Each (discrete) state value has a probability that evolves over time
- Some (discrete) state values are traversed with larger probability over time
- Figure shows a simulated example for a system with two species



population of  $S_1$ 

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#### • Idea: How about focusing on the most likely states only ?



The projected system can be solved exactly!

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#### • Starting from the (infinite) matrix representation of the CME:

The states of the chemical system can be enumerated:

 $\mathbf{X} := [\mathbf{x}_1 \quad \mathbf{x}_2 \quad \mathbf{x}_3 \quad \dots \quad ]^T$ 

The probability density state velocity  $\mathbf{F}(\mathbf{X} \cdot : R \rightarrow t)$  defined by:  $\mathbf{P}(\mathbf{X}; t) := [p(\mathbf{x}_1; t) \quad p(\mathbf{x}_2; t) \quad p(\mathbf{x}_3, t) \quad \dots \quad ]^T$ 

The evolution of the probability density state vector is governed by

$$\dot{\mathbf{P}}(\mathbf{X};t) = \mathbf{A} \cdot \mathbf{P}(\mathbf{X};t)$$

... one has the following result, where only the states indexed by the indexing vector J are retained (next slide):

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Let  $J = [m_1 \dots m_N]$  be an indexing vector. We define  $\mathbf{A}_J$  to be the principle submatrix of  $\mathbf{A}$  defined by J.

**Theorem [Projection Error Bound]:** Consider any Markov process in which the probability distribution evolves according to the ODE:  $\dot{\mathbf{P}}(\mathbf{X};t) = \mathbf{A} \cdot \mathbf{P}(\mathbf{X};t).$ If for an indexing vector J:  $\mathbf{1}^T \exp(\mathbf{A}_J t) \mathbf{P}(\mathbf{X}_J; 0) \ge 1 - \epsilon$ , then

$$\begin{bmatrix} \mathbf{P}(\mathbf{X}_{J};t) \\ \mathbf{P}(\mathbf{X}_{J'};t) \end{bmatrix} - \begin{bmatrix} \exp(\mathbf{A}_{J}t) \ \mathbf{P}(\mathbf{X}_{J};0) \\ 0 \end{bmatrix} \Big\|_{1} \le \epsilon$$

Munsky B. and Khammash M., Journal of Chemical Physics, 2006

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#### The FSP algorithm

- Step 0. Define the propensity functions and stoichiometry for all reactions.
  - Choose the initial probability density function P(X, 0).
  - Choose the final time of interest, t.
  - Choose the total amount of acceptable error  $\epsilon$ .
  - Choose an initial finite set of states:  $V_0$  **DED**



• Step 2. If  $\Gamma_{J_i} \ge 1 - \epsilon$ : stop.

 $\exp(\mathbf{A}_{J_i}t) \mathbf{P}(\mathbf{X}_{J_i}; 0)$  approximates  $\mathbf{P}(\mathbf{X}_{J_i}; t)$  to within  $\epsilon$ .

• Step 3. Add more states to get  $X_{J_{i+1}}$ . Increment i. Go to step 1.



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## Identification from snapshot data: Other methods

Moment matching: [e.g. work by J.Hespana]

Instead of probabilities, consider vector of all moments z and a truncation z\*

 $z(t) = \begin{bmatrix} Ex(t) & Ex(t)^T x(t) & \dots \end{bmatrix}^T, \qquad z^*(t) = \begin{bmatrix} Ex(t) & Ex(t)^T x(t) \end{bmatrix}^T$ evolving according to the equations depending on the model parameters  $z(t) = Bz(t), \qquad z^*(t) \simeq B^* z^*(t)$ 

and fit the equation for z\* to the corresp. empirical statistics from many cells

- At stochastic steady state: [Taniguchi et al., Science 329, 533 (2010)]
  - System evolves until stochastic equilibrium where p does not change
  - Use asymptotic approximation with a Gamma distribution

$$p(x;t) 
ightarrow d(x)$$
 for  $t 
ightarrow +\infty$ 

to fit (combinations of the) model parameters

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### Discussion

• Evidence for fundamental role of intrinsic and extrinsic noise

(e.g. Elowitz et al, Science, 2002)

- Vast literature on linear stochastic system identification useful but not sufficient
- Identification of stochastic models of genetic networks still in its infancy, first results on problem analysis and solution methods

(Munsky, Khammash et al 2009)

- More to exploit from data
  - Single-cell tracks reveal time correlation that cannot be observed in population snapshot data
  - Methods exploiting time correlation being developed and applied

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#### Conclusions

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- Masses of data wait for being processed. Automated processing unavoidable
- Modern experimental techniques enable inference of quantitative dynamic models at population and (sometimes) single cell level, even more to come
- Numerous applications in medicine, (bio)chemical industry etc.
- A lot of work in progress for model identification methods
- Intriguing mathematical problems
- Nonstandard identification problems: a lot to use, a lot to invent

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## ... Thank you!

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