

Model Based Target Identification from Gene Expression with Gaussian Processes

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Outline

Motivation

Probabilistic Model for TF Activity

Cascade Differential Equations

Discussion and Future Work

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Probabilistic Model for TF Activity

Cascade Differential Equations

Discussion and Future Work

Can a Biologist Fix a Radio? Lazebnik (2002)

The Case for Systems Biology

"It is difficult to find a black cat in a dark room, especially if there is no cat."

- ▶ Biological systems are immensely complicated.
- ▶ Lazebnik argues the need for models that are quantitative.
 - ▶ Such models should be predictive of biological behaviour.
 - ▶ Such models need to be combined with biological data.
- ▶ Systems biology:
 - ▶ Build mechanistic models (based on biochemical knowledge) of the system.
 - ▶ Identify modules, submodules, and parameterize the models.

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Coregulation of Gene Expression

The Case for Computational Biology

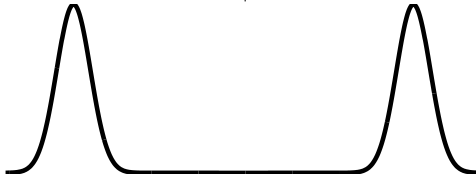
- ▶ Gene Expression to Transcriptional Regulation.
- ▶ A “data exploration” problem (computational biology/bioinformatics):
 - ▶ Use gene expression data to speculate on coregulated genes.
 - ▶ Traditionally use clustering of gene expression profiles.
- ▶ Contrast with (computational) systems biology approach:
 - ▶ Detailed mechanistic model of the system is created.
 - ▶ Fit parameters of the model to data.
 - ▶ Problematic for large data (genome wide).
 - ▶ Need to deal with unobserved biochemical species (TFs).

General Approach

Broadly Speaking: Two approaches to modeling

data modeling

mechanistic modeling



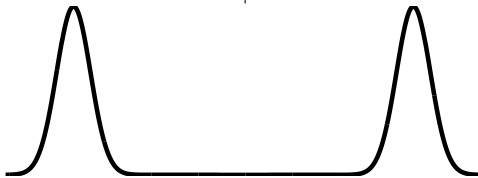
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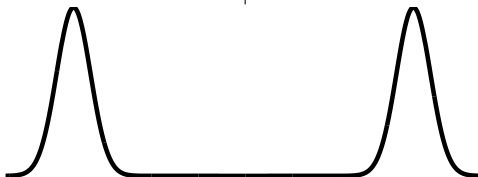
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impose physical laws



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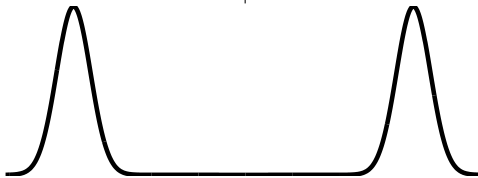
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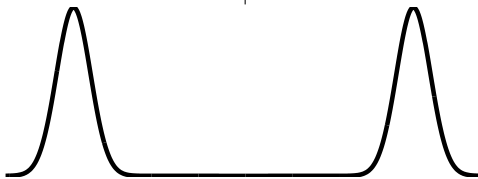
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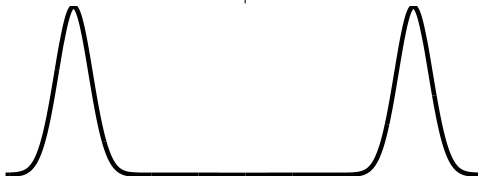
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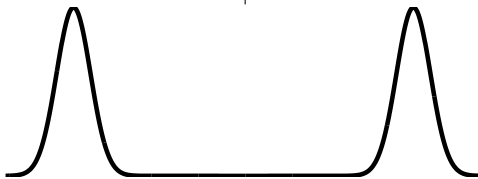
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differential equations



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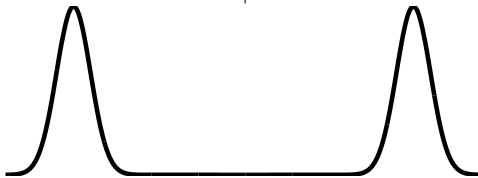
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adaptive models
PCA, clustering

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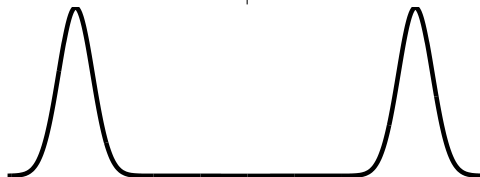
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SDE, ODE models



A Hybrid Approach

Introduce aspects of systems biology to computational models

- ▶ We advocate an approach *between* systems and computational biology.
- ▶ Introduce aspects of systems biology to the computational approach.
 - ▶ There is a computational penalty, but it may be worth paying.
 - ▶ Ideally there should be a smooth transition from pure computational (PCA, clustering, SVM classification) to systems (non-linear (stochastic) differential equations).
 - ▶ This work is one part of that transition.

Radiation Damage in the Cell

- ▶ Radiation can damage molecules including DNA.
- ▶ Most DNA damage is quickly repaired—single strand breaks, backbone break.
- ▶ Double strand breaks are more serious—a complete disconnect along the chromosome.
- ▶ Cell cycle stages:
 - ▶ G_1 : Cell is not dividing.
 - ▶ G_2 : Cell is preparing for mitosis, chromosomes have divided.
 - ▶ S: Cell is undergoing mitosis (DNA synthesis).
- ▶ Main problem is in G_1 . In G_2 there are two copies of the chromosome. In G_1 only one copy.

p53 “Guardian of the Cell”

- ▶ Responsible for Repairing DNA damage
- ▶ Activates DNA Repair proteins
- ▶ Pauses the Cell Cycle (prevents replication of damage DNA)
- ▶ Initiates *apoptosis* (cell death) in the case where damage can't be repaired.
- ▶ Large scale feedback loop with NF- κ B.

p53 DNA Damage Repair

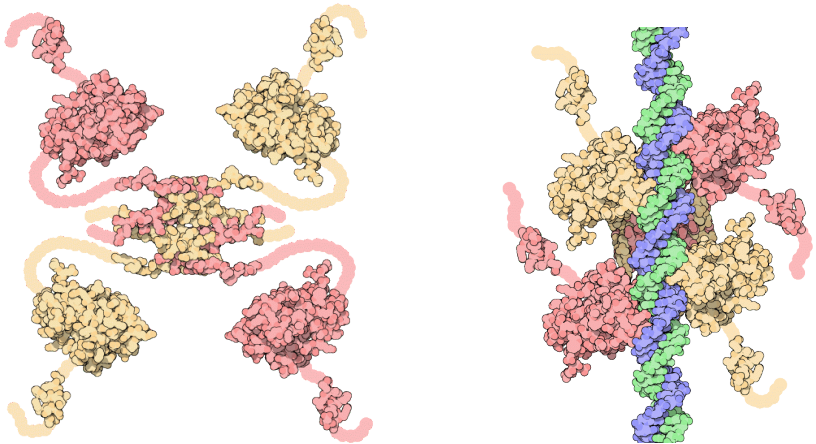


Figure: p53. *Left* unbound, *Right* bound to DNA. Images by David S. Goodsell from <http://www.rcsb.org/> (see the "Molecule of the Month" feature).

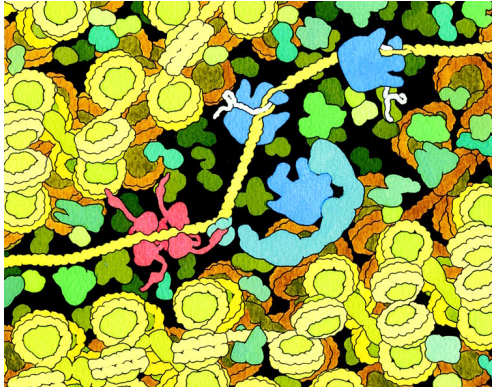


Figure: Repair of DNA damage by p53. Image from Goodsell (1999).

Some p53 Targets

DDB2 DNA Damage Specific DNA Binding Protein 2. (also governed by C/ EBP-beta, E2F1, E2F3,...).

p21 Cycline-dependent kinase inhibitor 1A (CDKN1A). A regulator of cell cycle progression. (also goverened by SREBP-1a, Sp1, Sp3,...).

hPA26/SESN1 sestrin 1 Cell Cycle arrest.

BIK BCL2-interacting killer. Induces cell death (apoptosis)

TNFRSF10b tumor necrosis factor receptor superfamily, member 10b. A transducer of apoptosis signals.

Modelling Assumption

- Assume p53 affects targets as a single input module network motif (SIM).

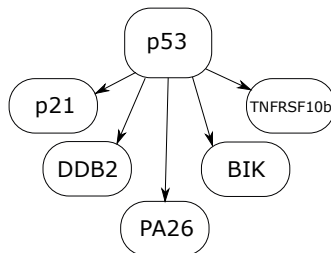


Figure: p53 SIM network motif as modelled by Barenco et al. 2006.

Standard Approach

Clustering of Gene Expression Profiles

- ▶ Assume that coregulated genes will cluster in the same groups.
- ▶ Perform clustering, and look for clusters containing target genes.
- ▶ These are candidates, look for confirmation in the literature etc.

Mathematical Model

- Differential equation model of system.

$$\frac{dx_j(t)}{dt} = B_j + S_j f(t) - D_j x_j(t)$$

rate of mRNA transcription, baseline transcription rate,
transcription factor activity, mRNA decay

- We have observations of $x_j(t)$ from gene expression. .

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$$\frac{dx_j(t)}{dt} = B_j + S_j f(t) - D_j x_j(t)$$
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- Reorder differential equation.

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- Jointly estimate $f(t)$ at observations of time points along with $\{B_j, D_j, S_j\}_{j=1}^g$.

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- Fit parameters by maximum likelihood or MCMC sampling.

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- Clustering model is equivalent to assuming D_j , B_j , and S_j are v. large.

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- Reorder differential equation and ignore gradient term.

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- We have observations of $x_j(t)$ from gene expression.
- Reorder differential equation and ignore gradient term.
- This suggests genes are scaled and offset versions of the TF.
- By normalizing data and clustering we hope to find those TFs.

Response of p53

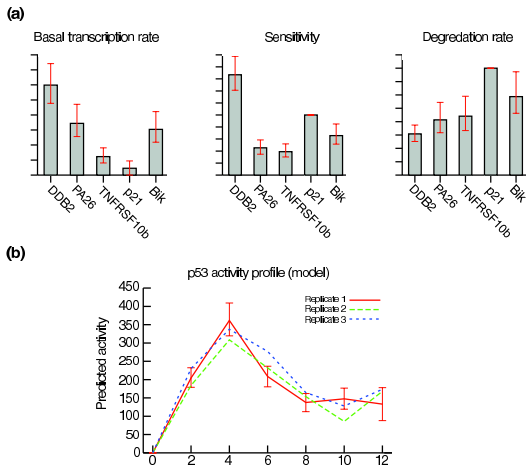


Figure: Results from Barenco et al. (2006). Top is parameter estimates. Bottom is inferred profile.

Response to p53 ...

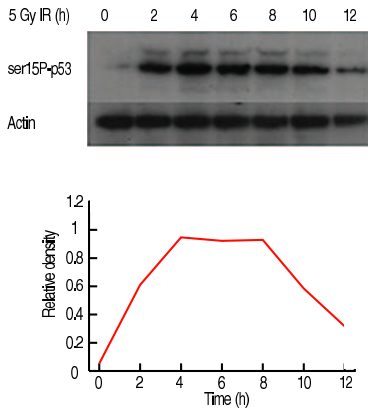


Figure: Results from Barenco et al. (2006). Activity profile of p53 was measured by Western blot to determine the levels of ser-15 phosphorylated p53 (ser15P-p53).

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Probabilistic Model for TF Activity

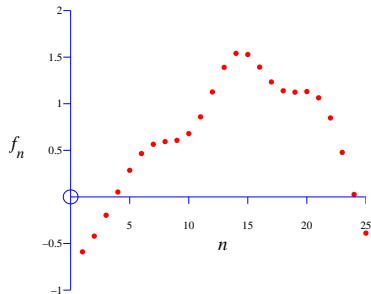
Cascade Differential Equations

Discussion and Future Work

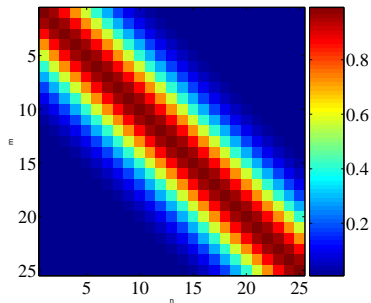
Probabilistic Model for $f(t)$

- ▶ We introduce a probabilistic model for $f(t)$.
- ▶ It is known as a Gaussian process, but we can think of it as a multivariate Gaussian (also known as a multivariate normal) distribution.
- ▶ The distribution has a mean vector, \mathbf{m} and a covariance matrix, \mathbf{K} .
- ▶ We will consider the mean to be zero: $\mathbf{m} = 0$.
- ▶ The covariance matrix will be structured to give correlation between samples.
- ▶ We will sample 25 points from the Gaussian distribution.
- ▶ Samples are governed by a 25×25 correlation matrix.

Gaussian Distribution Sample



(a) A 25 dimensional correlated random variable (values plotted against index)



(b) colormap showing correlations between dimensions

Figure: A sample from a 25 dimensional Gaussian distribution.

The covariance matrix

- ▶ Covariance matrix shows correlation between points f_m and f_n if n is near to m .
- ▶ Less correlation if n is distant from m .
- ▶ Our ordering of points means that the *function appears smooth*.
- ▶ In practice covariance matrix is computed as a function of time—index is equivalent to time.
- ▶ Different covariance functions give different characteristics.
- ▶ Because the models are *probabilistic* we can sample different characteristics.

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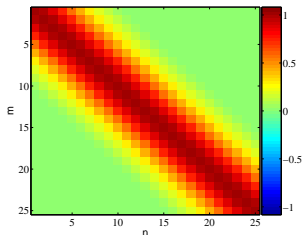
Covariance Functions

Where did this covariance matrix come from?

RBF Kernel Function

$$k(t, t') = \alpha \exp \left(-\frac{\|t - t'\|^2}{2\ell^2} \right)$$

- Covariance matrix is built using the *inputs* to the function t .
- For the example above it was based on Euclidean distance.
- The covariance function is also known as a kernel.



Covariance Samples

demCovFuncSample

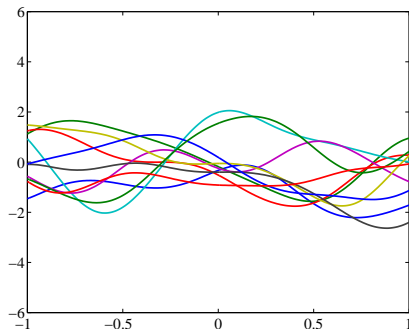


Figure: RBF kernel with $\ell = 10^{-\frac{1}{2}}$, $\alpha = 1$

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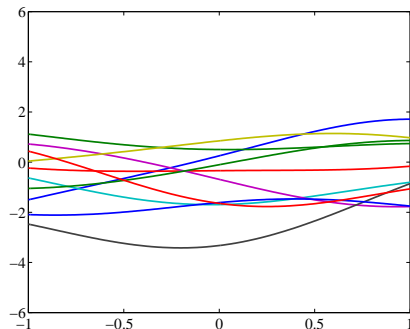


Figure: RBF kernel with $\ell = 1$, $\alpha = 1$

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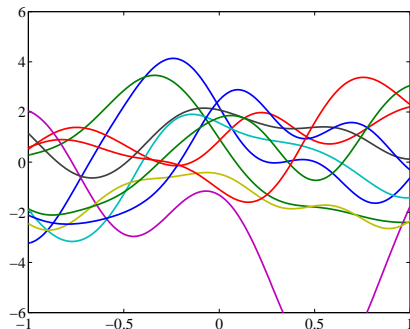


Figure: RBF kernel with $\ell = 0.3$, $\alpha = 4$

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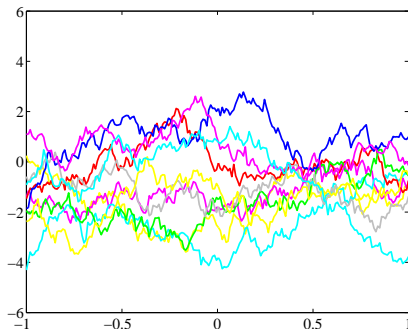


Figure: Ornstein-Uhlenbeck (stationary Gauss-Markov) covariance function $\ell = 1$, $\alpha = 4$

Example: Transcriptional Regulation

- First Order Differential Equation

$$\frac{dx_j(t)}{dt} = B_j + S_j f(t) - D_j x_j(t)$$

- It turns out that our Gaussian process assumption for $f(t)$, implies $x(t)$ is also a Gaussian process.
- The new Gaussian process is over $f(t)$ and all its targets: $x_1(t), x_2(t), \dots$ etc.
- Our new covariance matrix gives correlations between all these functions.
- This gives us a *probabilistic* model for transcriptional regulation.

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Covariance for Transcription Model

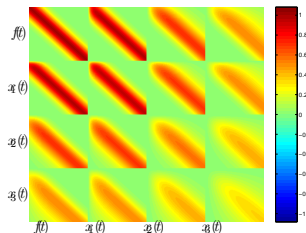
RBF covariance function for $f(t)$

$$x_i(t) = \frac{B_i}{D_i} + S_i \exp(-D_i t) \int_0^t f(u) \exp(D_i u) du.$$

- Joint distribution for $x_1(t)$, $x_2(t)$, $x_3(t)$, and $f(t)$.

- Here:

D_1	S_1	D_2	S_2	D_3	S_3
5	5	1	1	0.5	0.5



Covariance for Transcription Model

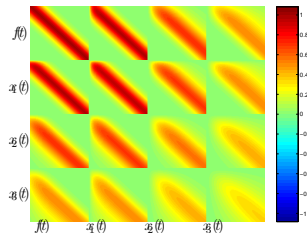
RBF covariance function for $f(t)$

$$x = b/d + \sum_i \mathbf{e}_i^\top \mathbf{f} \quad \mathbf{f} \sim \mathcal{N}(\mathbf{0}, \Sigma_i) \rightarrow x \sim \mathcal{N}\left(b/d, \sum_i \mathbf{e}_i^\top \Sigma_i \mathbf{e}_i\right)$$

- Joint distribution for $x_1(t)$, $x_2(t)$, $x_3(t)$, and $f(t)$.

- Here:

D_1	S_1	D_2	S_2	D_3	S_3
5	5	1	1	0.5	0.5



Covariance for Transcription Model

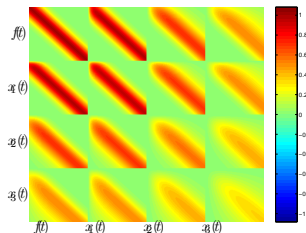
RBF covariance function for $f(t)$

$$x_i(t) = \frac{B_i}{D_i} + S_i \exp(-D_i t) \int_0^t f(u) \exp(D_i u) du.$$

- Joint distribution for $x_1(t)$, $x_2(t)$, $x_3(t)$, and $f(t)$.

- Here:

D_1	S_1	D_2	S_2	D_3	S_3
5	5	1	1	0.5	0.5



Joint Sampling of $f(t)$ and $x(t)$

► `simSample`

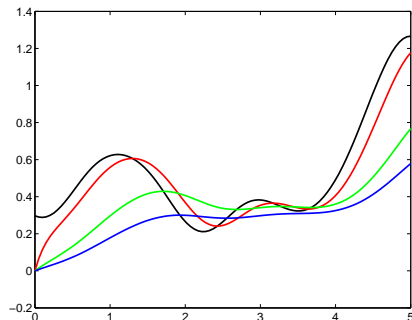


Figure: Joint samples from the ODE covariance, *black*: $f(t)$, *red*: $x_1(t)$ (high decay/sensitivity), *green*: $x_2(t)$ (medium decay/sensitivity) and *blue*: $x_3(t)$ (low decay/sensitivity).

Joint Sampling of $f(t)$ and $x(t)$

► `simSample`

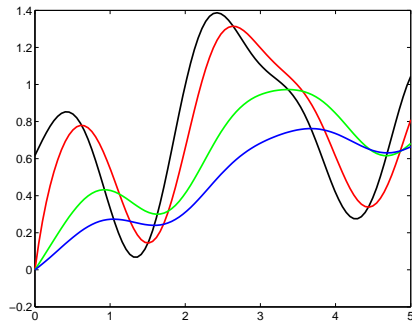


Figure: Joint samples from the ODE covariance, *black*: $f(t)$, *red*: $x_1(t)$ (high decay/sensitivity), *green*: $x_2(t)$ (medium decay/sensitivity) and *blue*: $x_3(t)$ (low decay/sensitivity).

Joint Sampling of $f(t)$ and $x(t)$

► `simSample`

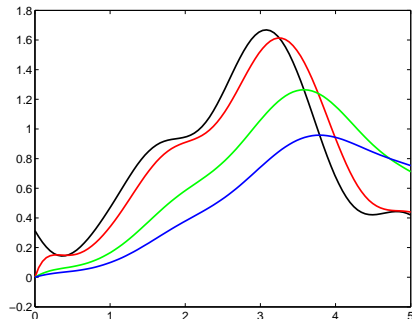


Figure: Joint samples from the ODE covariance, *black*: $f(t)$, *red*: $x_1(t)$ (high decay/sensitivity), *green*: $x_2(t)$ (medium decay/sensitivity) and *blue*: $x_3(t)$ (low decay/sensitivity).

Joint Sampling of $f(t)$ and $x(t)$

► `simSample`

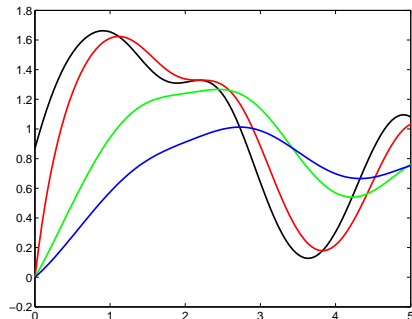
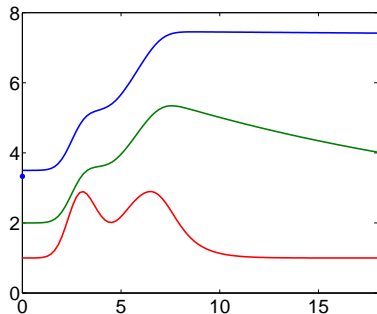


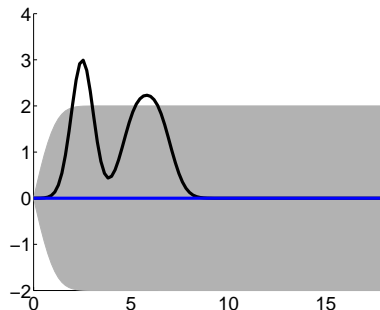
Figure: Joint samples from the ODE covariance, *black*: $f(t)$, *red*: $x_1(t)$ (high decay/sensitivity), *green*: $x_2(t)$ (medium decay/sensitivity) and *blue*: $x_3(t)$ (low decay/sensitivity).

Artificial Example: Inferring $f(t)$

Inferring TF activity from artificially sampled genes.



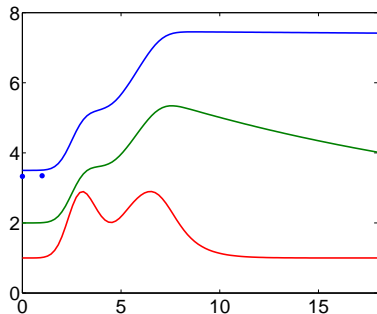
True “gene profiles” and noisy observations.



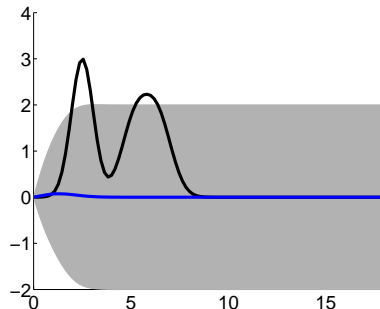
Inferred transcription factor activity.

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Inferring TF activity from artificially sampled genes.



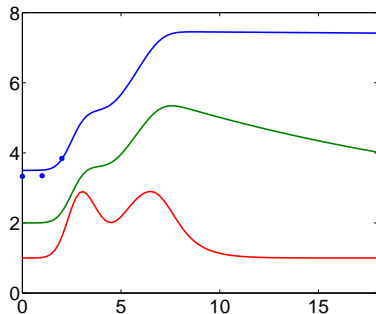
True “gene profiles” and noisy observations.



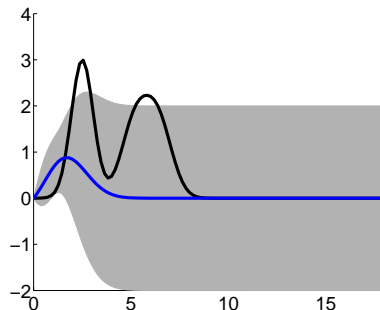
Inferred transcription factor activity.

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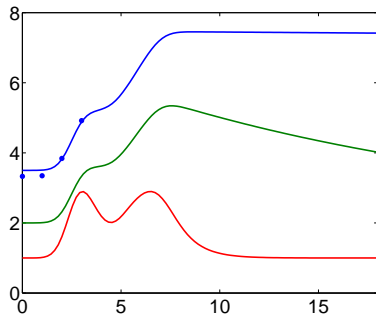
True “gene profiles” and noisy observations.



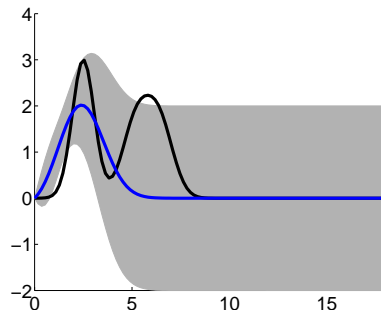
Inferred transcription factor activity.

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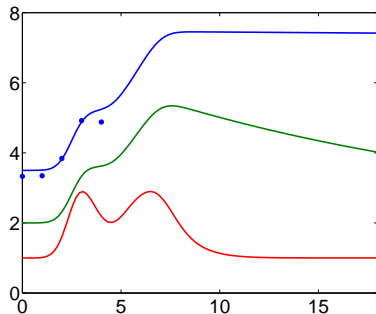
True “gene profiles” and noisy observations.



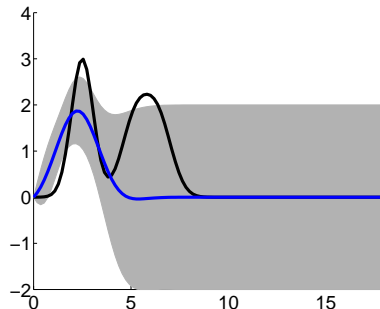
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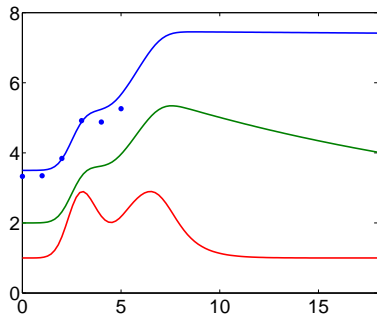
True “gene profiles” and noisy observations.



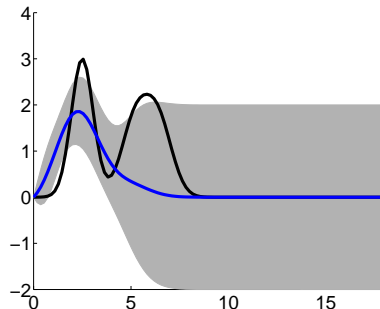
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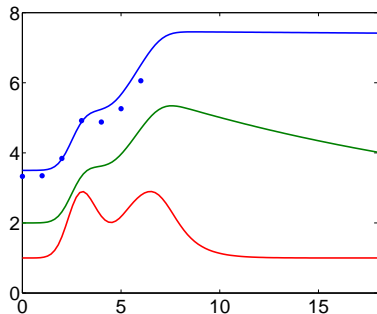
True “gene profiles” and noisy observations.



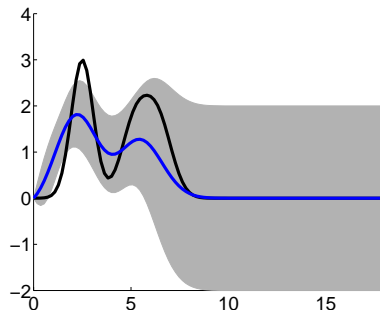
Inferred transcription factor activity.

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Inferring TF activity from artificially sampled genes.



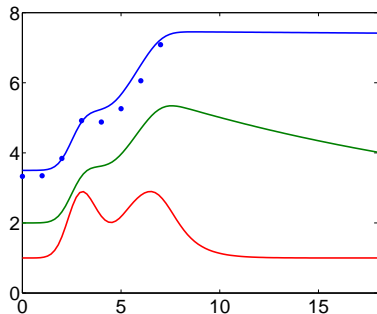
True “gene profiles” and noisy observations.



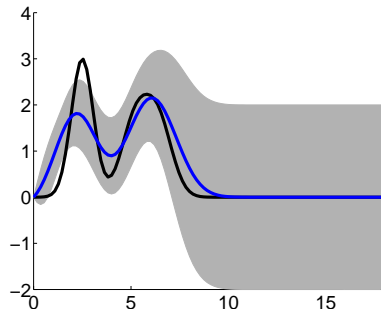
Inferred transcription factor activity.

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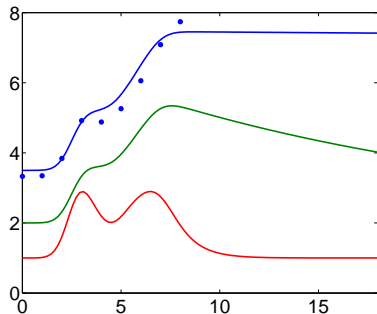
True “gene profiles” and noisy observations.



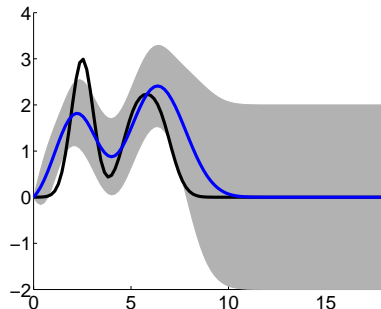
Inferred transcription factor activity.

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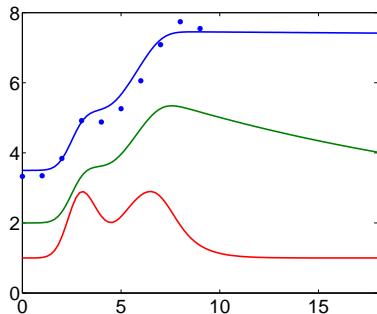
True “gene profiles” and noisy observations.



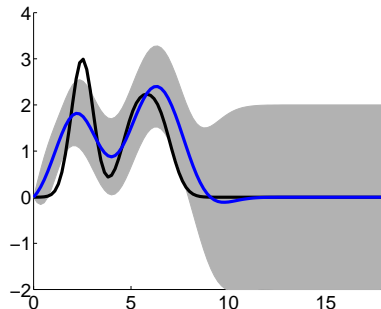
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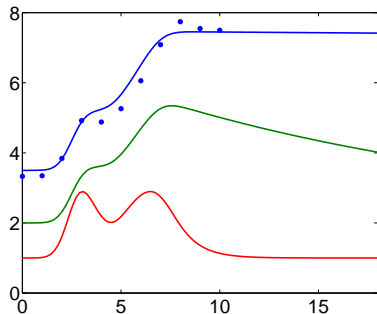
True “gene profiles” and noisy observations.



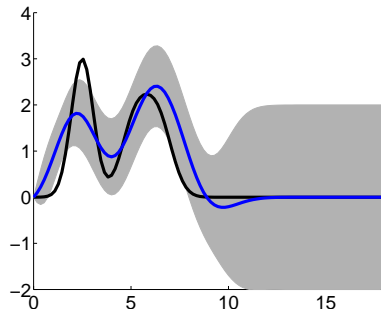
Inferred transcription factor activity.

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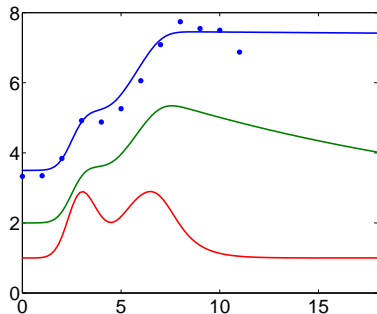
True “gene profiles” and noisy observations.



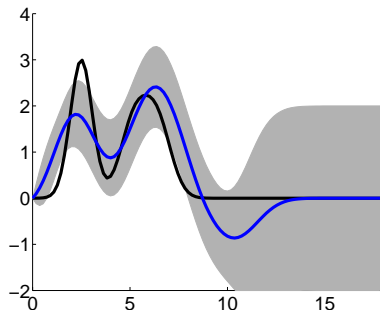
Inferred transcription factor activity.

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Inferring TF activity from artificially sampled genes.



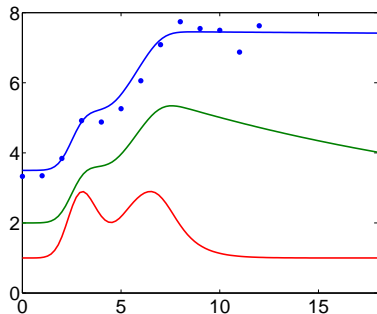
True “gene profiles” and noisy observations.



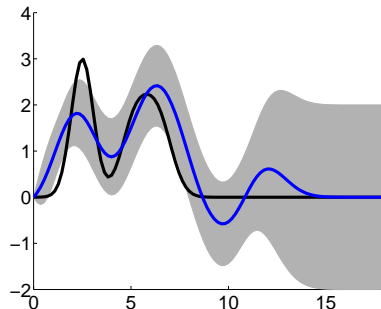
Inferred transcription factor activity.

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Inferring TF activity from artificially sampled genes.



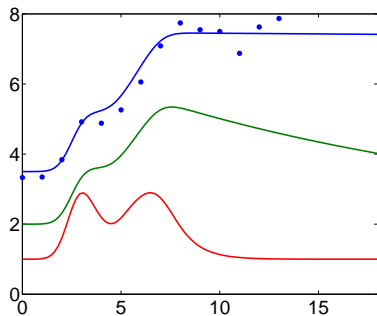
True “gene profiles” and noisy observations.



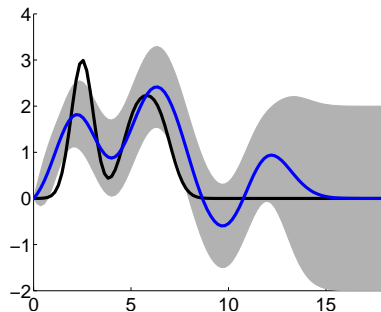
Inferred transcription factor activity.

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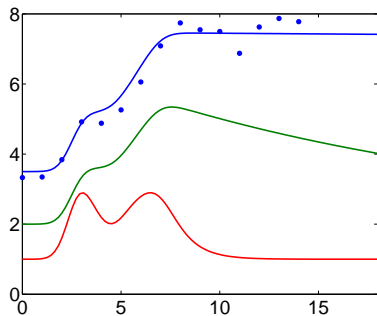
True “gene profiles” and noisy observations.



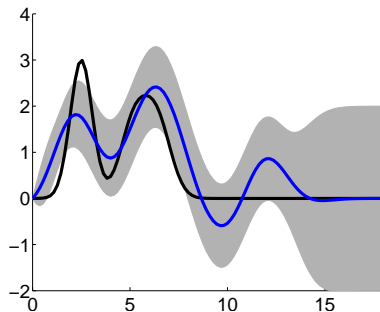
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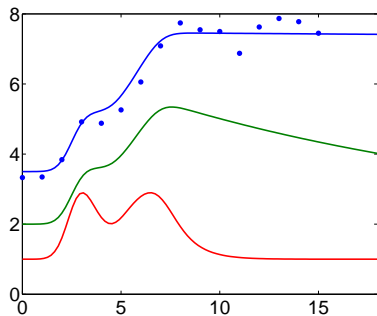
True “gene profiles” and noisy observations.



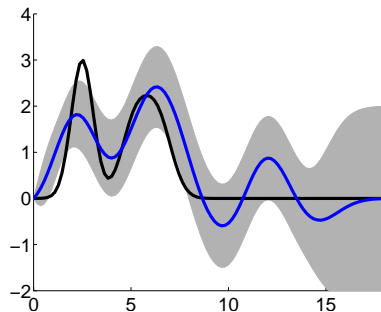
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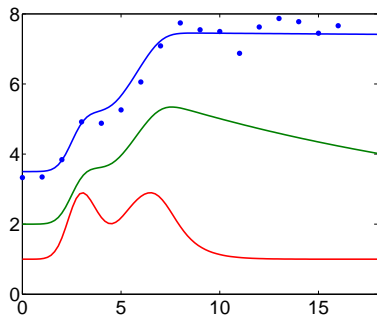
True “gene profiles” and noisy observations.



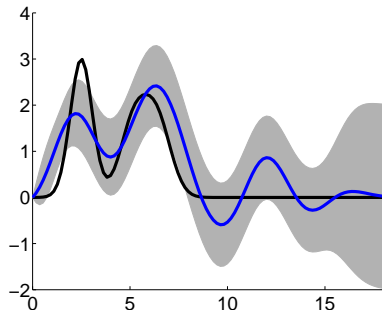
Inferred transcription factor activity.

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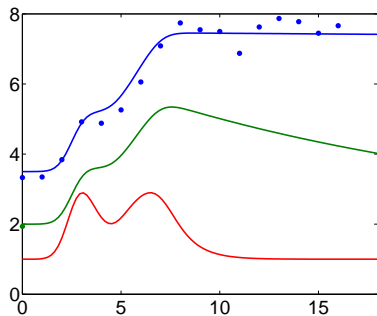
True “gene profiles” and noisy observations.



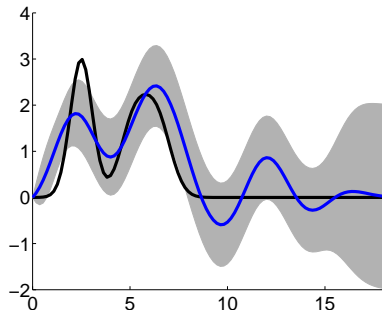
Inferred transcription factor activity.

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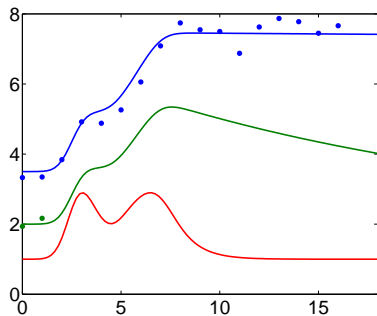
True “gene profiles” and noisy observations.



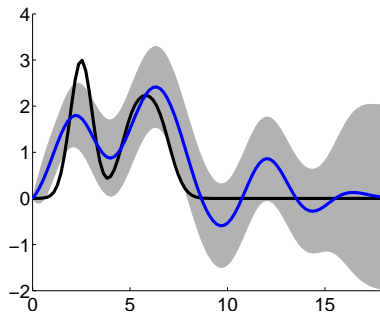
Inferred transcription factor activity.

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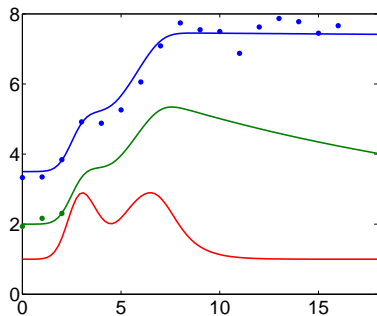
True “gene profiles” and noisy observations.



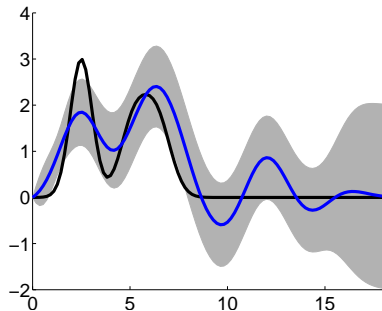
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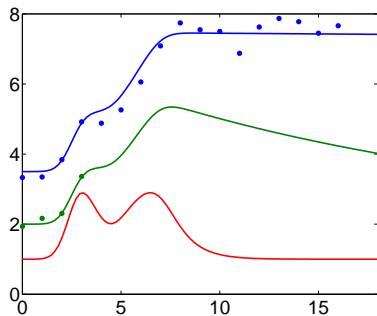
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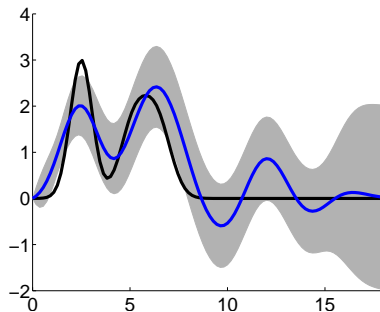
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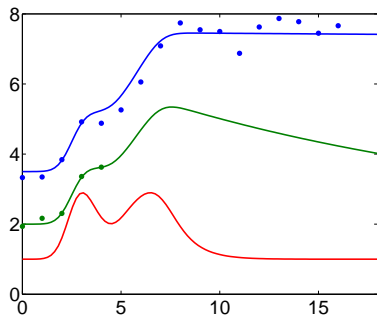
True “gene profiles” and noisy observations.



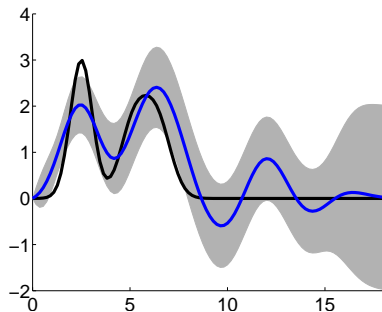
Inferred transcription factor activity.

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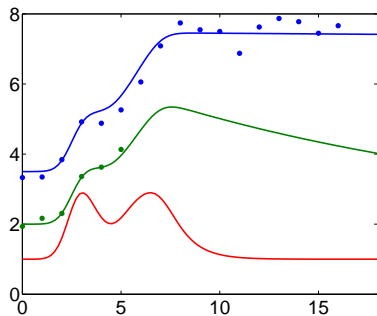
True “gene profiles” and noisy observations.



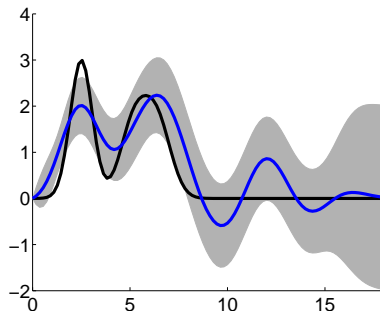
Inferred transcription factor activity.

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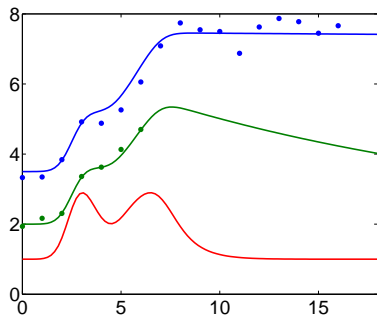
True “gene profiles” and noisy observations.



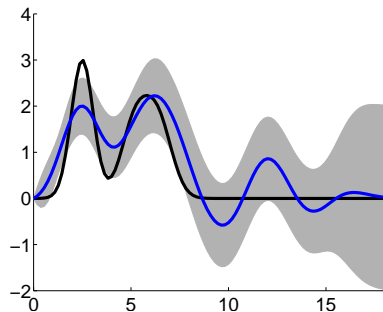
Inferred transcription factor activity.

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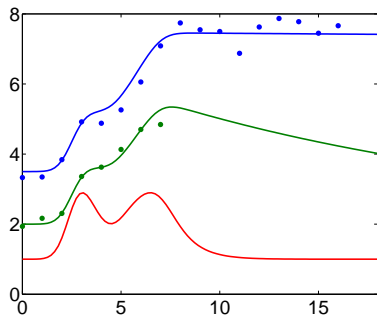
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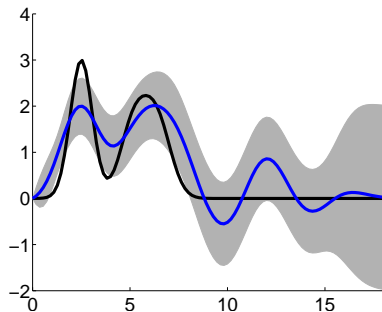
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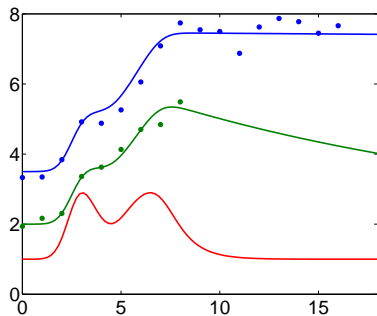
True “gene profiles” and noisy observations.



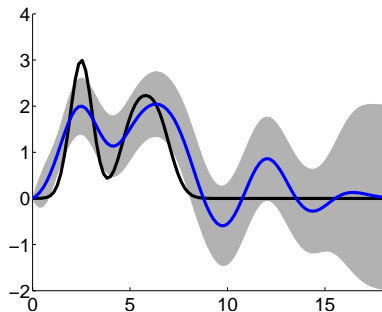
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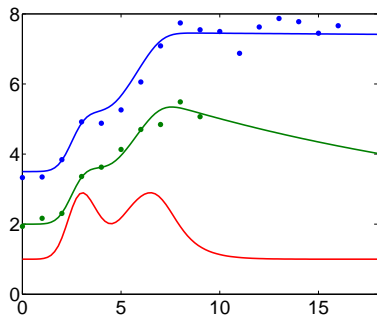
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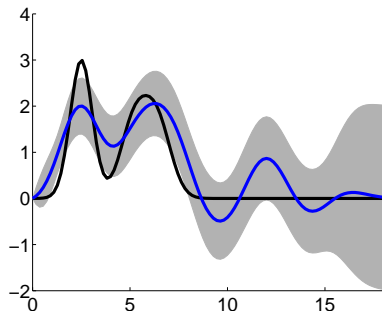
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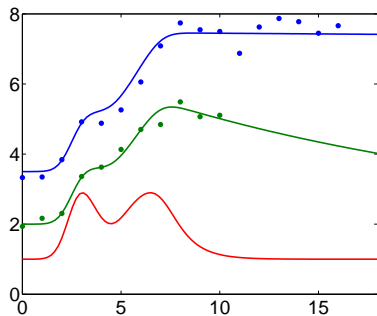
True “gene profiles” and noisy observations.



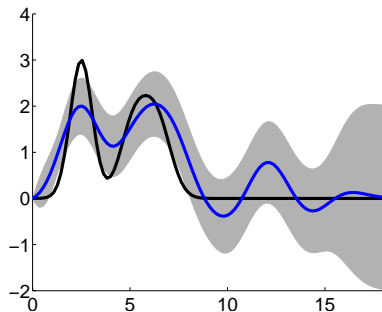
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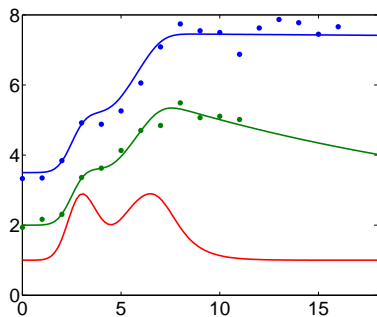
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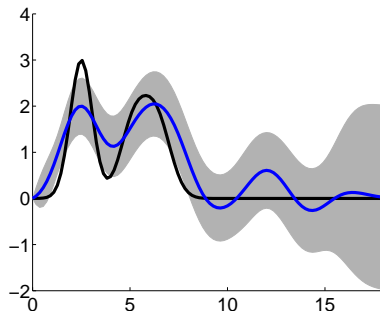
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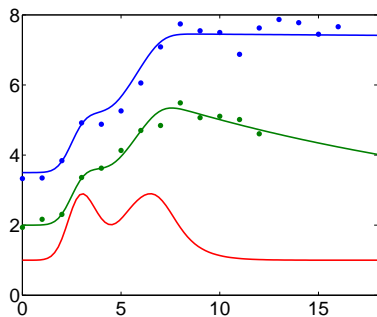
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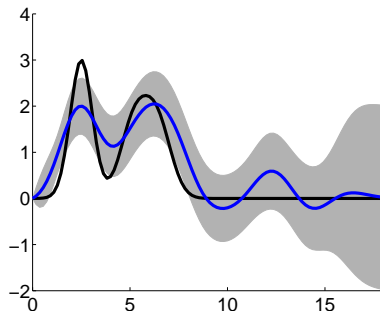
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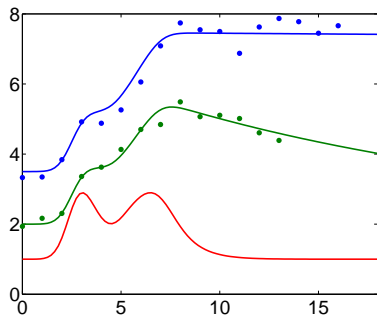
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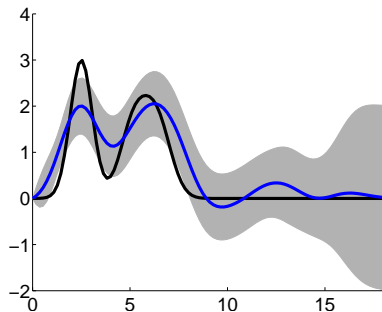
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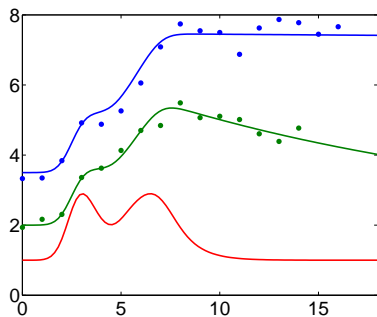
True “gene profiles” and noisy observations.



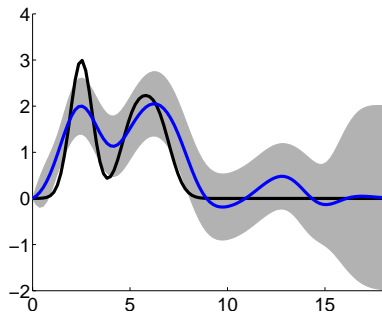
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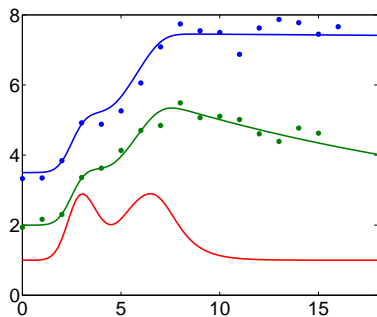
True “gene profiles” and noisy observations.



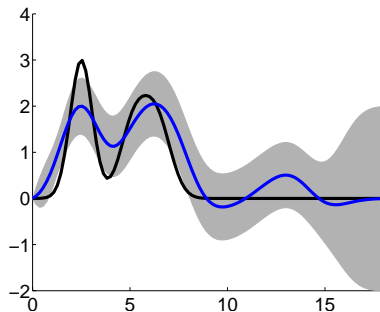
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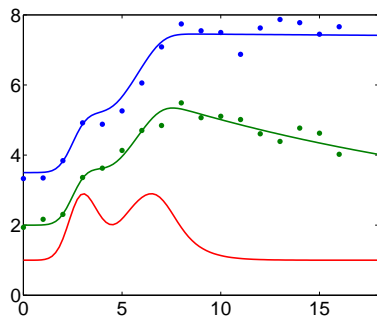
True “gene profiles” and noisy observations.



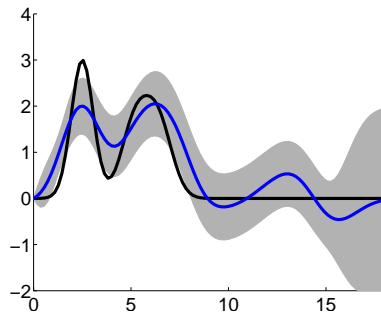
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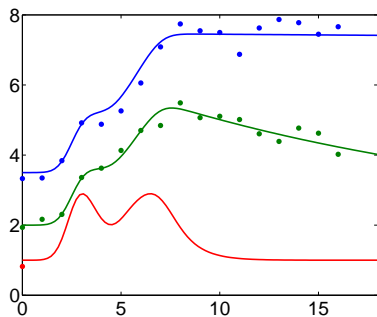
True “gene profiles” and noisy observations.



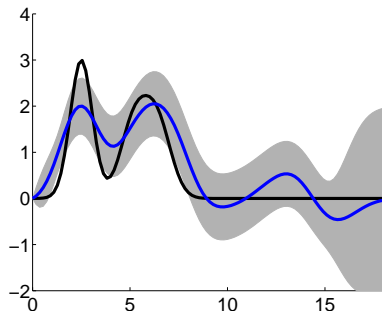
Inferred transcription factor activity.

Artificial Example: Inferring $f(t)$

Inferring TF activity from artificially sampled genes.



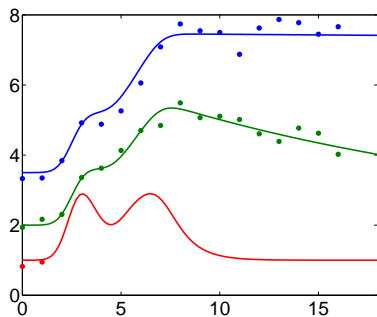
True “gene profiles” and noisy observations.



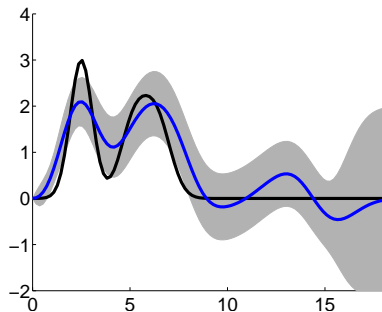
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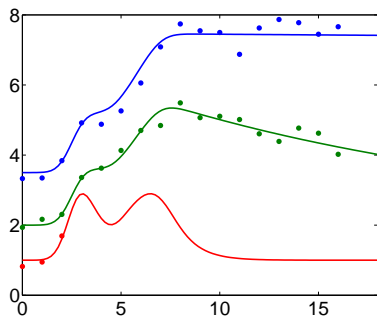
True “gene profiles” and noisy observations.



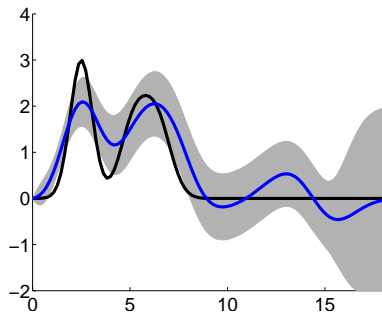
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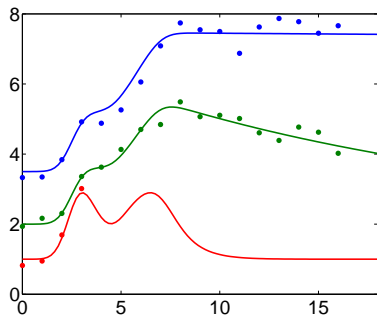
True “gene profiles” and noisy observations.



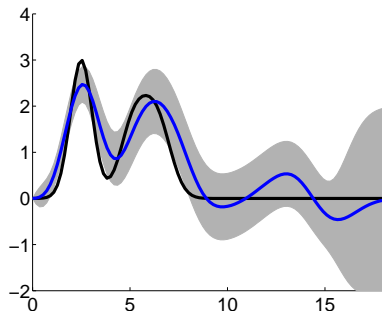
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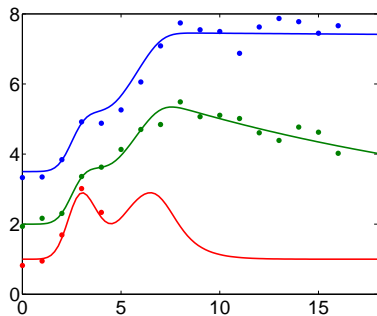
True “gene profiles” and noisy observations.



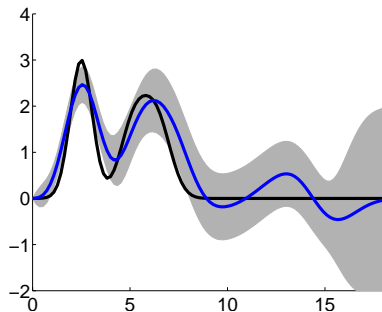
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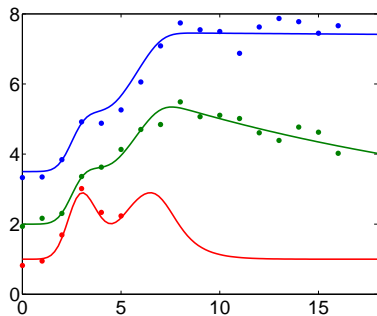
True “gene profiles” and noisy observations.



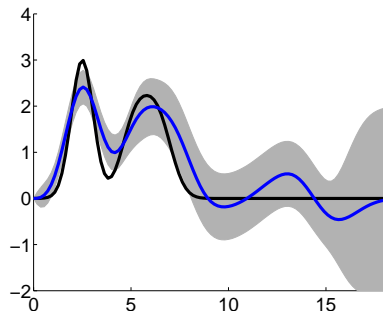
Inferred transcription factor activity.

Artificial Example: Inferring $f(t)$

Inferring TF activity from artificially sampled genes.



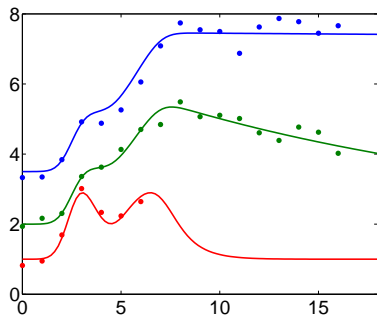
True “gene profiles” and noisy observations.



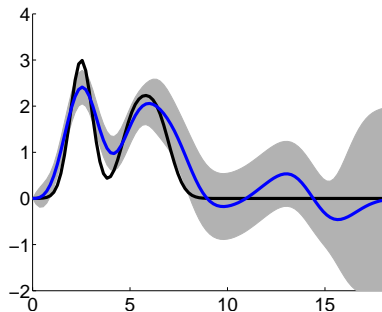
Inferred transcription factor activity.

Artificial Example: Inferring $f(t)$

Inferring TF activity from artificially sampled genes.



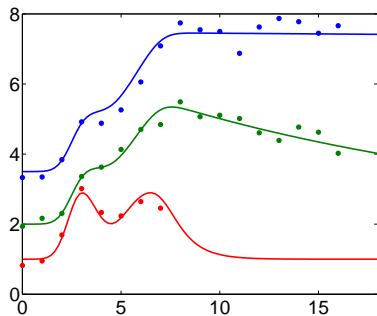
True “gene profiles” and noisy observations.



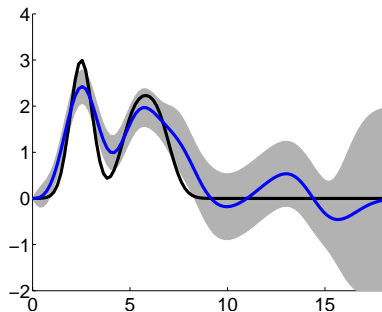
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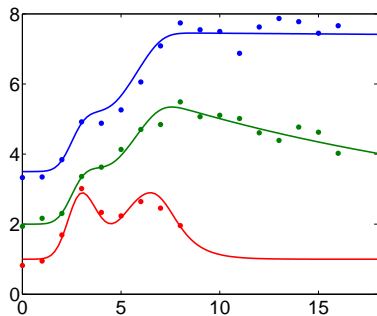
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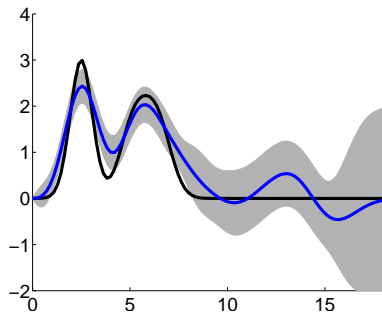
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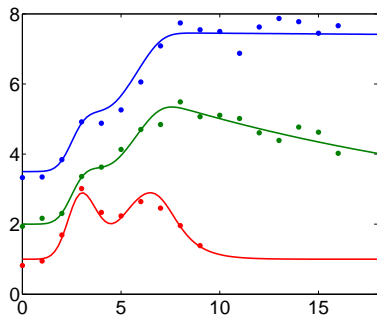
True “gene profiles” and noisy observations.



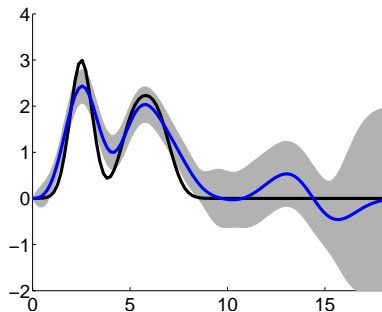
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Artificial Example: Inferring $f(t)$

Inferring TF activity from artificially sampled genes.



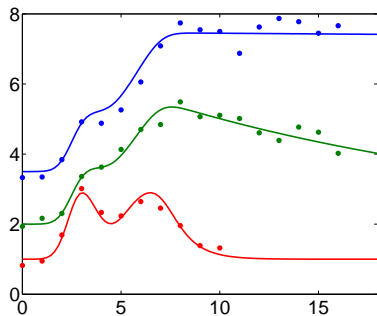
True “gene profiles” and noisy observations.



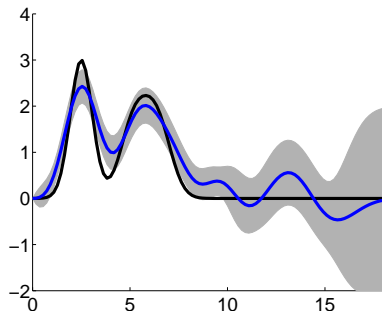
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Inferring TF activity from artificially sampled genes.



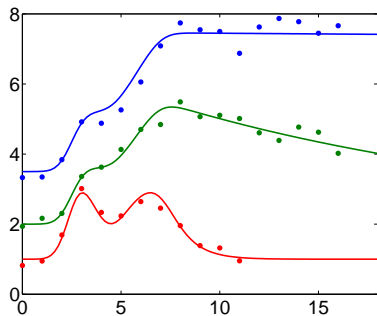
True “gene profiles” and noisy observations.



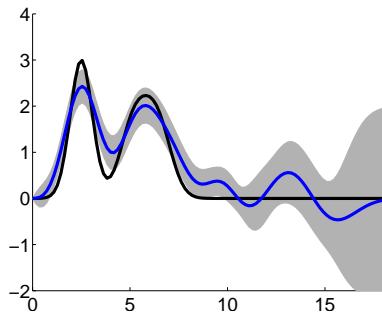
Inferred transcription factor activity.

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Inferring TF activity from artificially sampled genes.



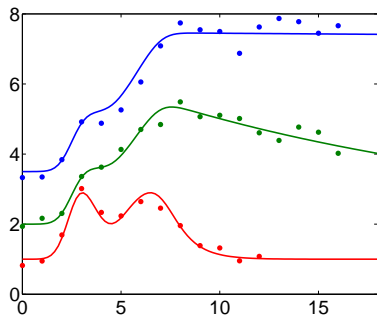
True “gene profiles” and noisy observations.



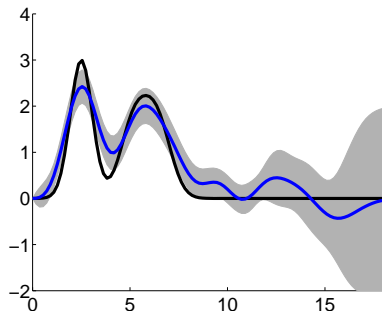
Inferred transcription factor activity.

Artificial Example: Inferring $f(t)$

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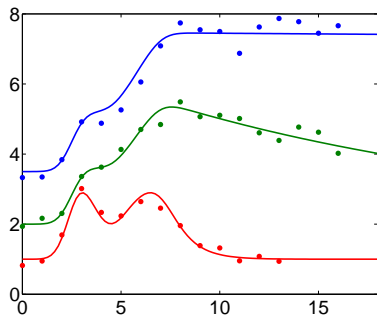
True “gene profiles” and noisy observations.



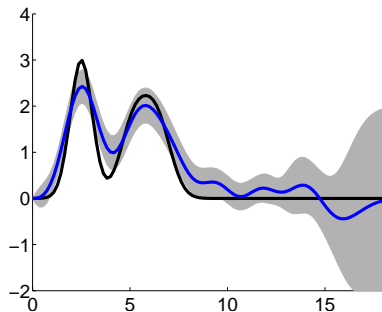
Inferred transcription factor activity.

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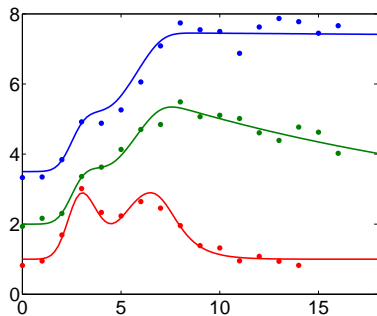
True “gene profiles” and noisy observations.



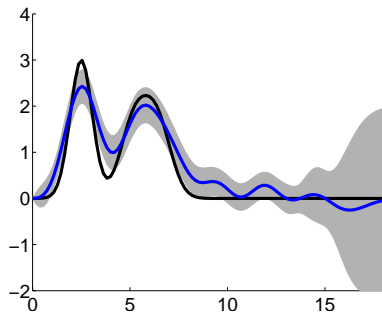
Inferred transcription factor activity.

Artificial Example: Inferring $f(t)$

Inferring TF activity from artificially sampled genes.



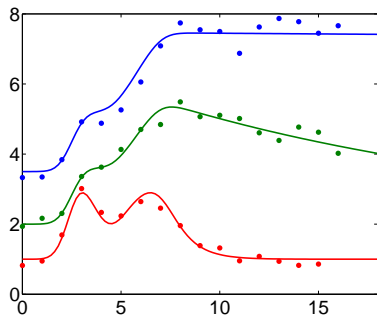
True “gene profiles” and noisy observations.



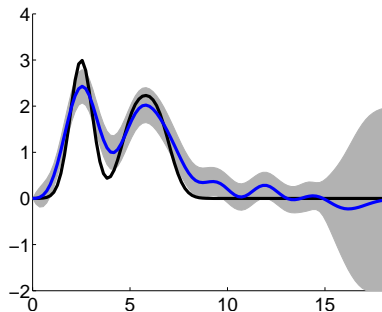
Inferred transcription factor activity.

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Inferring TF activity from artificially sampled genes.



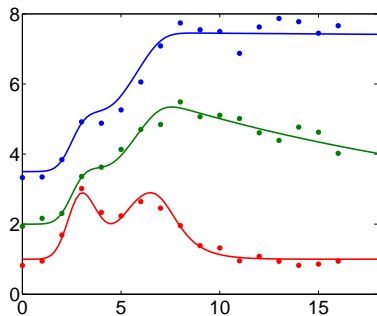
True “gene profiles” and noisy observations.



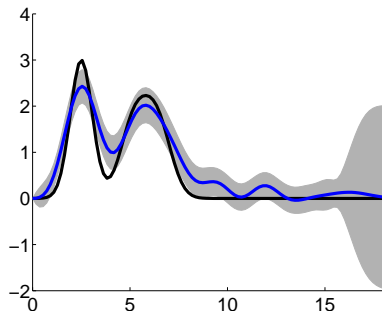
Inferred transcription factor activity.

Artificial Example: Inferring $f(t)$

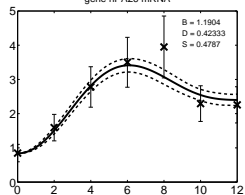
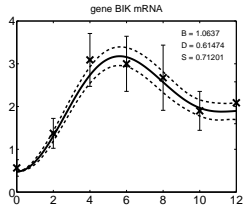
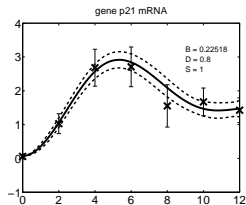
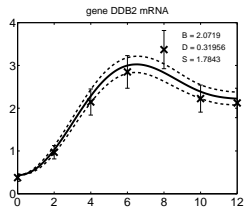
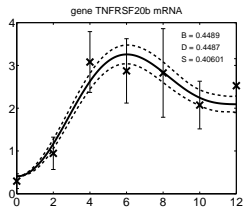
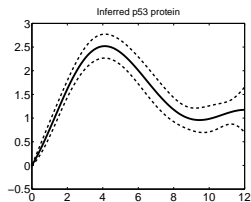
Inferring TF activity from artificially sampled genes.



True “gene profiles” and noisy observations.



Inferred transcription factor activity.

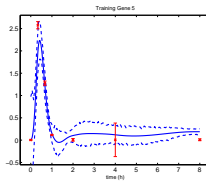
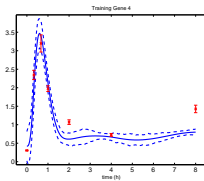
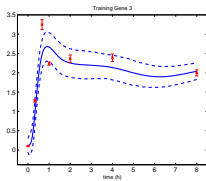
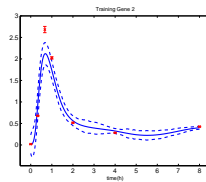
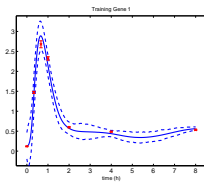
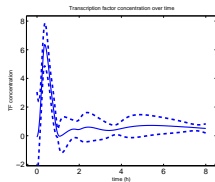


Ranking with ERK Signalling

- ▶ Target Ranking for Elk-1.
- ▶ Elk-1 is phosphorylated by ERK from the EGF signalling pathway.
- ▶ Predict concentration of Elk-1 from known targets.
- ▶ Rank other targets of Elk-1.

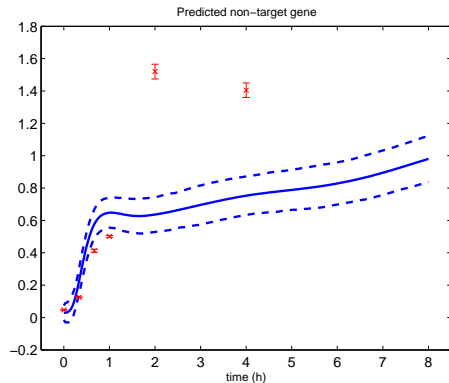
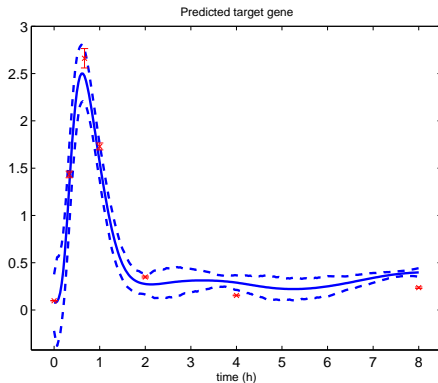
Elk-1 (MLP covariance)

Jennifer Withers



Elk-1 target selection

Fitted model used to rank potential targets of Elk-1



Outline

Motivation

Probabilistic Model for TF Activity

Cascade Differential Equations

Discussion and Future Work

Antti Honkela

- ▶ Transcription factor protein also has governing mRNA.
- ▶ This mRNA can be measured.
- ▶ In signalling systems this measurement can be misleading because it is activated (phosphorylated) transcription factor that counts.
- ▶ In development phosphorylation plays less of a role.

Collaboration with Furlong Lab in EMBL Heidelberg.

- ▶ Mesoderm development in *Drosophila melanogaster* (fruit fly).
- ▶ Mesoderm forms in triploblastic animals (along with ectoderm and endoderm). Mesoderm develops into muscles, and circulatory system.
- ▶ The transcription factor Twist initiates *Drosophila* mesoderm development, resulting in the formation of heart, somatic muscle, and other cell types.
- ▶ Wildtype microarray experiments publicly available.
- ▶ Can we use the cascade model to predict viable targets of Twist?

We take the production rate of active transcription factor to be given by

$$\begin{aligned}\frac{df(t)}{dt} &= \sigma y(t) - \delta f(t) \\ \frac{dx_j(t)}{dt} &= B_j + S_j f(t) - D_j x_j(t)\end{aligned}$$

The solution for $f(t)$, setting transient terms to zero, is

$$f(t) = \sigma \exp(-\delta t) \int_0^t y(u) \exp(\delta u) du .$$

Covariance for Translation/Transcription Model

RBF covariance function for $y(t)$

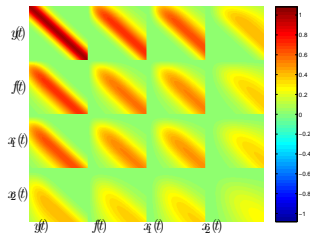
$$f(t) = \sigma \exp(-\delta t) \int_0^t y(u) \exp(\delta u) du$$

$$x_i(t) = \frac{B_i}{D_i} + S_i \exp(-D_i t) \int_0^t f(u) \exp(D_i u) du.$$

- Joint distribution for $x_1(t)$, $x_2(t)$, $f(t)$ and $y(t)$.

- Here:

δ	D_1	S_1	D_2	S_2
1	5	5	0.5	0.5



Joint Sampling of $y(t)$, $f(t)$, and $x(t)$

► `disimSample`

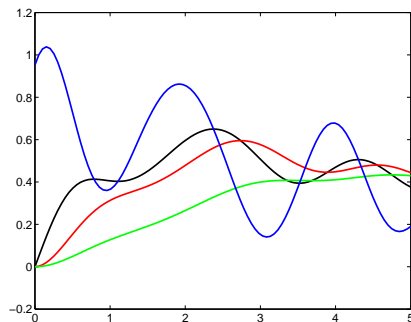


Figure: Joint samples from the ODE covariance, *blue*: $y(t)$ (mRNA of TF), *black*: $f(t)$ (TF concentration), *red*: $x_1(t)$ (high decay target) and *green*: $x_2(t)$ (low decay target)

Joint Sampling of $y(t)$, $f(t)$, and $x(t)$

► `disimSample`

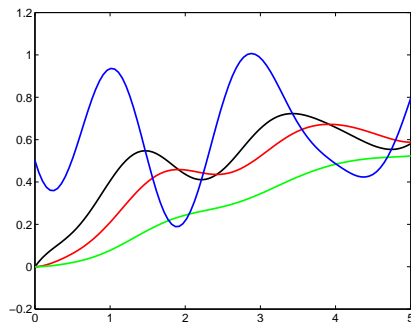


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Joint Sampling of $y(t)$, $f(t)$, and $x(t)$

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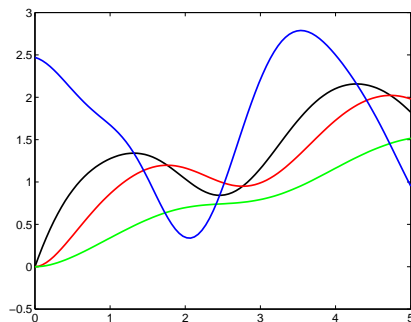


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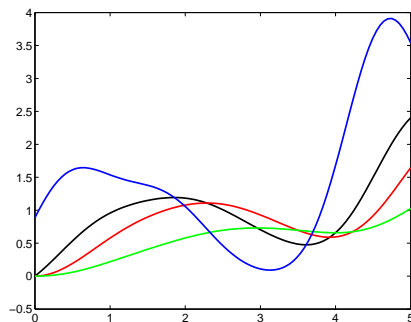


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Twist Results

- ▶ Use mRNA of Twist as driving input.
- ▶ For each gene build a cascade model that forces Twist to be the only TF.
- ▶ Compare fit of this model to a baseline (e.g. similar model but sensitivity zero).
- ▶ Rank according to the likelihood above the baseline.
- ▶ Compare with correlation, knockouts and time series network identification (TSNI) (Della Gatta et al., 2008).

Results for Twi using the Cascade model

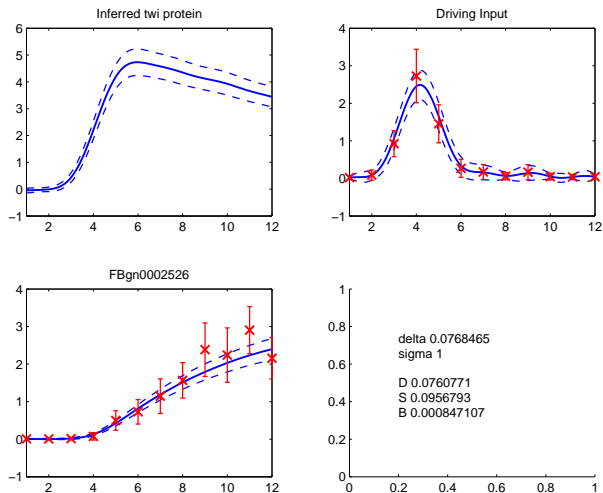


Figure: Model for flybase gene identity FBgn0002526.

Results for Twi using the Cascade model

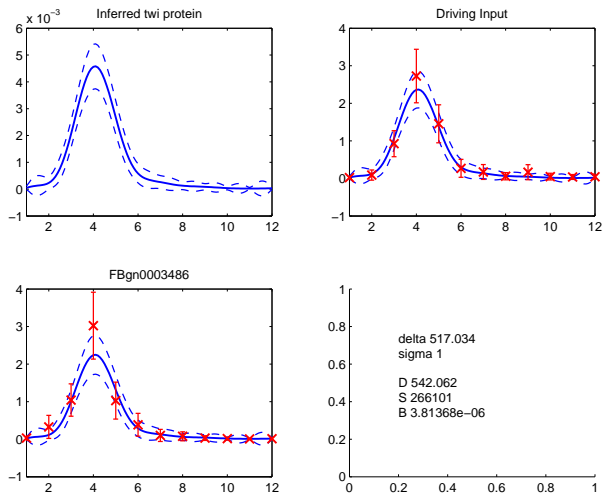


Figure: Model for flybase gene identity FBgn0003486.

Results for Twi using the Cascade model

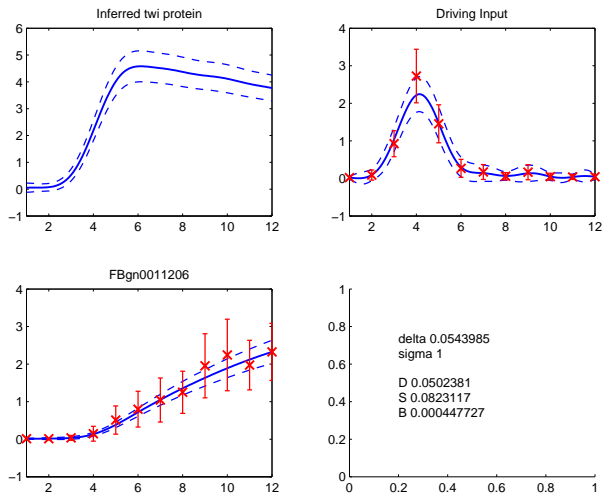


Figure: Model for flybase gene identity FBgn0011206.

Results for Twi using the Cascade model

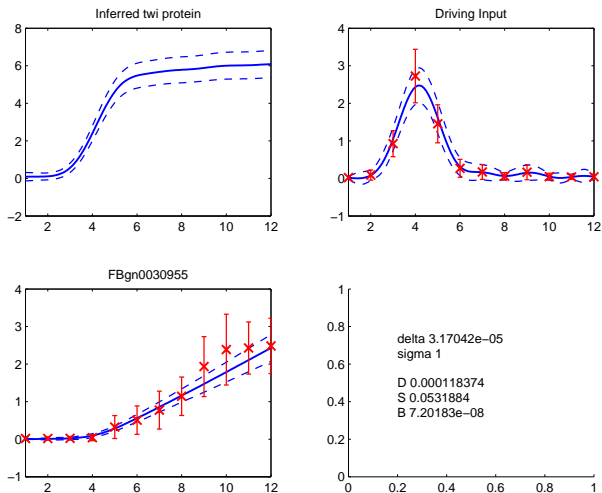


Figure: Model for flybase gene identity FBgn00309055.

Results for Twi using the Cascade model

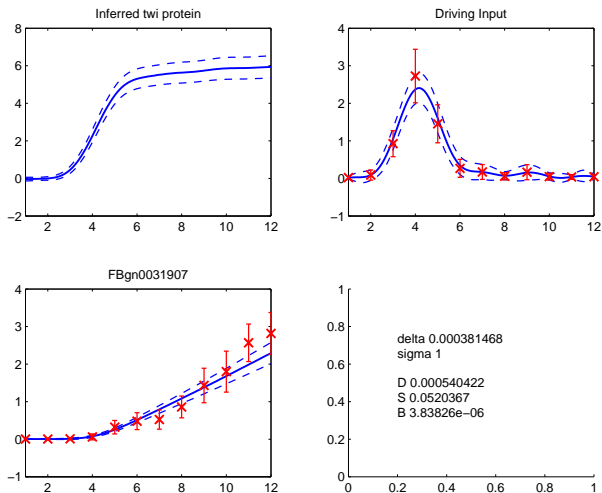


Figure: Model for flybase gene identity FBgn0031907.

Results for Twi using the Cascade model

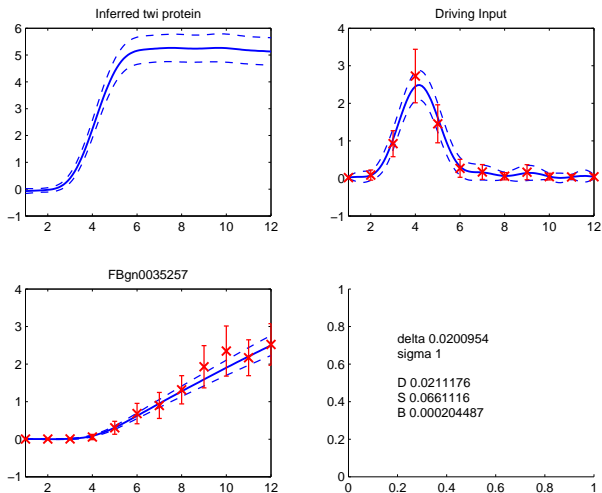


Figure: Model for flybase gene identity FBgn0035257.

Results for Twi using the Cascade model

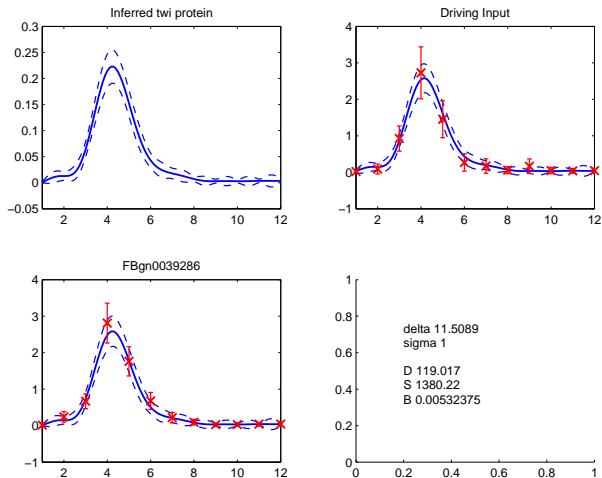


Figure: Model for flybase gene identity FBgn0039286.

Results of Ranking

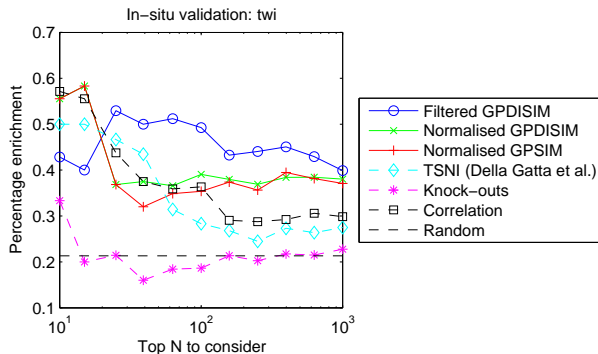


Figure: Percentage enrichment for top N targets for relevant terms in *Drosophila* in situs.

Results of Ranking

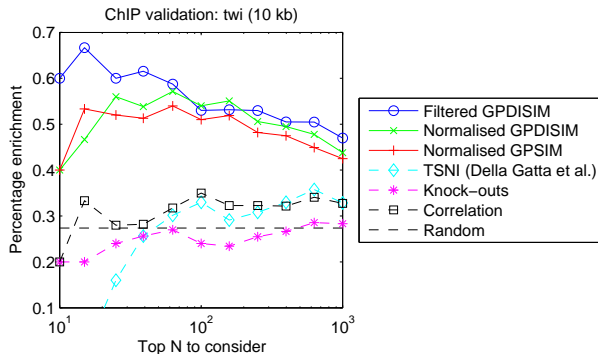


Figure: Percentage enrichment for top N targets for ChIP-chip confirmed targets.

Summary

- ▶ Cascade models allow genomewide analysis of potential targets given only expression data.
- ▶ Once a set of potential candidate targets have been identified, they can be modelled in a more complex manner.
- ▶ We don't have ground truth, but evidence indicates that the approach *can* perform as well as knockouts.

Outline

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Cascade Differential Equations

Discussion and Future Work

Discussion and Future Work

- ▶ Integration of probabilistic inference with mechanistic models.
- ▶ Applications in modeling gene expression.
- ▶ Cascade model introduces model of translation.
- ▶ Ongoing/other work:
 - ▶ Non linear response and non linear differential equations.
 - ▶ Scaling up to larger systems.
 - ▶ Stochastic differential equations.

Acknowledgements

- ▶ Investigators: Neil Lawrence and Magnus Rattray
- ▶ Researchers: Pei Gao, Antti Honkela, Guido Sanguinetti, and Jennifer Withers
- ▶ Martino Barenco and Mike Hubank at the Institute of Child Health in UCL (p53 pathway).
- ▶ Charles Girardot and Eileen Furlong of EMBL in Heidelberg (mesoderm development in *D. Melanogaster*).

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Nonlinear Response

Nonlinear Response Models

Consider the following modification to the model,

$$\frac{dx_j(t)}{dt} = B_j + S_j g(f(t)) - D_j x_j(t),$$

where $g(\cdot)$ is a non-linear function. The differential equation can still be solved,

$$x_j(t) = \frac{B_j}{D_j} + S_j \int_0^t e^{-D_j(t-u)} g_j(f(u)) du$$

MAP-Laplace Approximation

Based on Laplace's method,

$$p(\mathbf{f} \mid \mathbf{x}) = N(\hat{\mathbf{f}}, \mathbf{A}^{-1}) \propto \exp\left(-\frac{1}{2}(\mathbf{f} - \hat{\mathbf{f}})^T \mathbf{A}(\mathbf{f} - \hat{\mathbf{f}})\right)$$

where $\hat{\mathbf{f}} = \operatorname{argmax}_{\mathbf{f}} p(\mathbf{f} \mid \mathbf{x})$ and $\mathbf{A} = -\nabla \nabla \log p(\mathbf{f} \mid \mathbf{y})|_{\mathbf{f}=\hat{\mathbf{f}}}$ is the Hessian of the negative posterior at that point. To obtain $\hat{\mathbf{f}}$ and \mathbf{A} ,

we define the following function $\psi(\mathbf{f})$ as:

$$\log p(\mathbf{f} \mid \mathbf{x}) \propto \psi(\mathbf{f}) = \log p(\mathbf{x} \mid \mathbf{f}) + \log p(\mathbf{f})$$

MAP-Laplace Approximation

Assigning a GP prior distribution to $f(t)$, it then follows that

$$\log p(\mathbf{f}) = -\frac{1}{2}\mathbf{f}^T \mathbf{K}^{-1}\mathbf{f} - \frac{1}{2}\log |\mathbf{K}| - \frac{n}{2}\log 2\pi$$

where \mathbf{K} is the covariance matrix of $f(t)$. Hence,

$$\begin{aligned}\nabla\psi(\mathbf{f}) &= \nabla \log p(\mathbf{x}|\mathbf{f}) - \mathbf{K}^{-1}\mathbf{f} \\ \nabla\nabla\psi(\mathbf{f}) &= \nabla\nabla \log p(\mathbf{x}|\mathbf{f}) - \mathbf{K}^{-1} = -\mathbf{W} - \mathbf{K}^{-1}\end{aligned}$$

Estimation of $\psi(\mathbf{f})$

Newton's method is applied to find the maximum of $\psi(\mathbf{f})$ as

$$\begin{aligned}\mathbf{f}^{new} &= \mathbf{f} - (\nabla \nabla \psi(\mathbf{f}))^{-1} \nabla \psi(\mathbf{f}) \\ &= (\mathbf{W} + \mathbf{K}^{-1})^{-1} (\mathbf{W}\mathbf{f} - \nabla \log p(\mathbf{x}|\mathbf{f}))\end{aligned}$$

In addition, $\mathbf{A} = -\nabla \nabla \psi(\hat{\mathbf{f}}) = \mathbf{W} + \mathbf{K}^{-1}$ where \mathbf{W} is the negative Hessian matrix. Hence, the Laplace approximation to the posterior is a Gaussian with mean $\hat{\mathbf{f}}$ and covariance matrix \mathbf{A}^{-1} as

$$p(\mathbf{f} | \mathbf{x}) \simeq N(\hat{\mathbf{f}}, \mathbf{A}^{-1}) = N(\hat{\mathbf{f}}, (\mathbf{W} + \mathbf{K}^{-1})^{-1})$$

Model Parameter Estimation

The marginal likelihood is useful for estimating the model parameters θ and covariance parameters ϕ

$$p(\mathbf{x}|\theta, \phi) = \int p(\mathbf{x}|\mathbf{f}, \theta) p(\mathbf{f}|\phi) d\mathbf{f} = \int \exp(\psi(\mathbf{f})) d\mathbf{f}$$

Using Taylor expansion of $\psi(\mathbf{f})$,

$$\log p(\mathbf{x}|\theta, \phi) = \log p(\mathbf{x}|\hat{\mathbf{f}}, \theta, \phi) - \frac{1}{2}\mathbf{f}^T \mathbf{K}^{-1} \mathbf{f} - \frac{1}{2} \log |\mathbf{I} + \mathbf{K} \mathbf{W}|$$

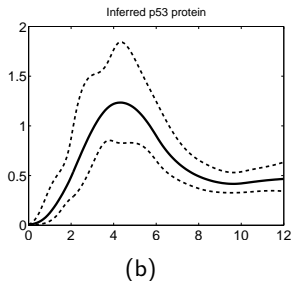
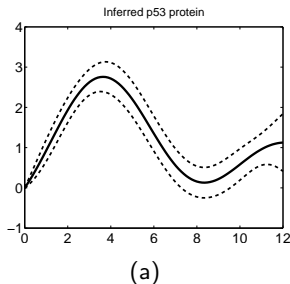
The parameters $\eta = \{\theta, \phi\}$ can be then estimated by using

$$\frac{\partial \log p(\mathbf{x}|\eta)}{\partial \eta} = \frac{\partial \log p(\mathbf{x}|\eta)}{\partial \eta} \Big|_{\text{explicit}} + \frac{\partial \log p(\mathbf{x}|\eta)}{\partial \hat{\mathbf{f}}} \frac{\partial \hat{\mathbf{f}}}{\partial \eta}$$

- The Michaelis-Menten activation model uses the following non-linearity

$$g_j(f(t)) = \frac{e^{f(t)}}{\gamma_j + e^{f(t)}},$$

where we are using a GP $f(t)$ to model the log of the TF activity.



Valdiation of Laplace Approximation

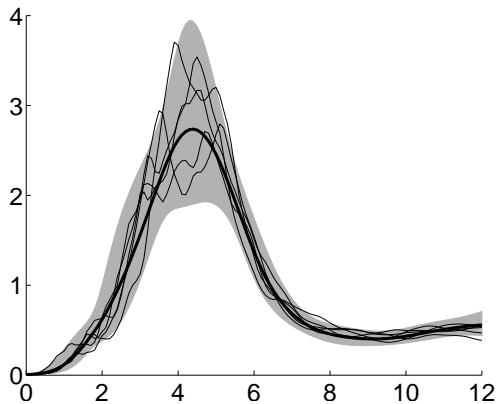


Figure: Laplace approximation error bars along with samples from the true posterior distribution.

SOS Response

- ▶ DNA damage may occur as a result of activity of antibiotics.
- ▶ LexA is bound to the genome preventing transcription of the SOS genes.
- ▶ RecA protein is stimulated by single stranded DNA, inactivates the LexA repressor.
- ▶ This allows several of the LexA targets to transcribe.
- ▶ The SOS pathway may be essential in antibiotic resistance Cirz et al. (2005).
- ▶ Aim is to target these proteins to produce drugs to increase efficacy of antibiotics Lee et al. (2005).

LexA Experimental Description

- ▶ Data from Courcelle et al. (2001)
- ▶ UV irradiation of *E. coli*. in both wild-type cells and *lexA1* mutants, which are unable to induce genes under LexA control.
- ▶ Response measured with two color hybridization to cDNA arrays.

Their Model

Given measurements of gene expression at N time points $(t_0, t_1, \dots, t_{N-1})$, the temporal profile of a gene i , $x_i(t)$, that solves the ODE in Eq. 1 can be approximated by

$$x_i(t) = x_i^0 e^{-\delta_i t} + \frac{B_i}{D_i} + S_i e^{-\delta_i t} \frac{1}{D_i} \sum_{j=0}^{N-2} (e^{D_i t_{j+1}} - e^{D_i t_j}) \frac{1}{\gamma_i + \bar{f}_j}$$

where $\bar{f}_j = \frac{(f(t_j) + f(t_{j+1}))}{2}$ on each subinterval (t_j, t_{j+1}) , $j = 0, \dots, N-2$. This is under the simplifying assumption that $f(t)$ is a piece-wise constant function on each subinterval (t_j, t_{j+1}) .

Khanin et al. (2006) Results Reminder

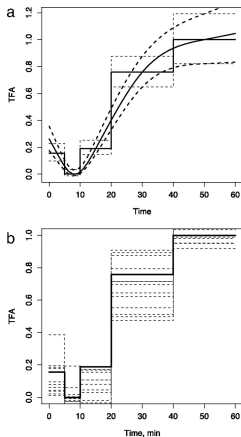


Figure: Fig. 2 from Khanin et al. (2006): Reconstructed activity level of master repressor LexA, following a UV dose of 40 J/m².

Their Results

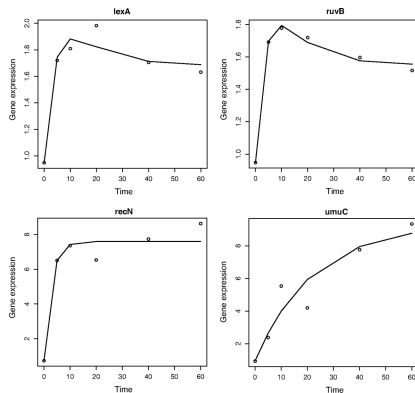


Figure: Fig. 3 from Khanin et al. (2006): Reconstructed profiles for four genes in the LexA SIM.

- We can use the same model of repression,

$$g_j(f(t)) = \frac{1}{\gamma_j + e^{f(t)}}$$

In the case of repression we have to include the transient term,

$$x_j(t) = \alpha_j e^{-D_j t} + \frac{B_j}{D_j} + S_j \int_0^t e^{-D_j(t-u)} g_j(f(u)) du$$

Results for the repressor LexA

Pei Gao

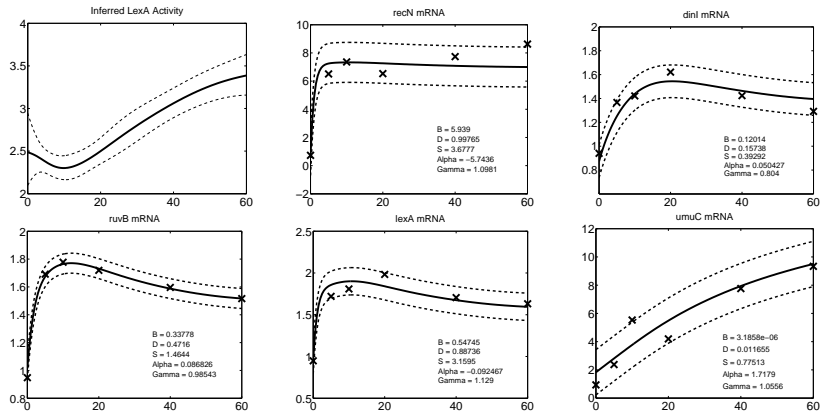


Figure: Our results using an MLP kernel. To appear at ECCB08 Gao et al. (2008).

Michalis Titsias

- ▶ Sample in Gaussian processes

$$p(\mathbf{f}|\mathbf{x}) \propto p(\mathbf{x}|\mathbf{f}) p(\mathbf{f})$$

- ▶ Likelihood relates GP to data through

$$x_j(t) = \alpha_j e^{-D_j t} + \frac{B_j}{D_j} + S_j \int_0^t e^{-D_j(t-u)} g_j(f(u)) du$$

- ▶ We use *control points* for fast sampling.

MCMC for Non Linear Response

The Metropolis-Hastings algorithm

- ▶ Initialize $\mathbf{f}^{(0)}$
- ▶ Form a Markov chain. Use a proposal distribution $Q(\mathbf{f}^{(t+1)}|\mathbf{f}^{(t)})$ and accept with the M-H step

$$\min \left(1, \frac{p(\mathbf{x}|\mathbf{f}^{(t+1)})p(\mathbf{f}^{(t+1)})}{p(\mathbf{x}|\mathbf{f}^{(t)})p(\mathbf{f}^{(t)})} \frac{Q(\mathbf{f}^{(t)}|\mathbf{f}^{(t+1)})}{Q(\mathbf{f}^{(t+1)}|\mathbf{f}^{(t)})} \right)$$

- ▶ \mathbf{f} can be very *high dimensional* (hundreds of points)
- ▶ How do we choose the proposal $Q(\mathbf{f}^{(t+1)}|\mathbf{f}^{(t)})$?
 - ▶ Can we use the GP prior $p(\mathbf{f})$ as the proposal?

Sampling using control points

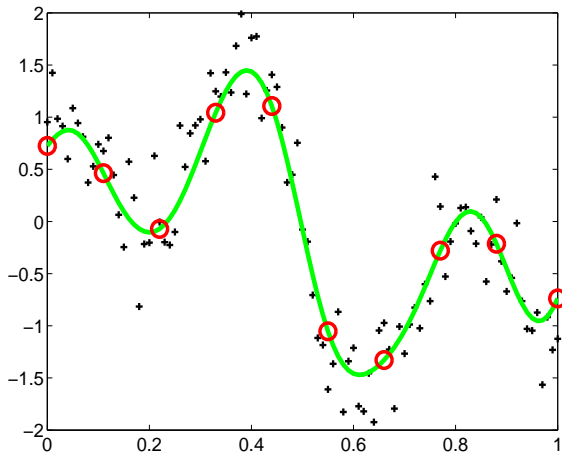
- ▶ Separate the points in \mathbf{f} into two groups:
 - ▶ few control points \mathbf{f}_c
 - ▶ and the large majority of the remaining points $\mathbf{f}_\rho = \mathbf{f} \setminus \mathbf{f}_c$
- ▶ Sample the control points \mathbf{f}_c using a proposal $q\left(\mathbf{f}_c^{(t+1)}|\mathbf{f}_c^{(t)}\right)$
- ▶ Sample the remaining points \mathbf{f}_ρ using the conditional GP prior $p\left(\mathbf{f}_\rho^{(t+1)}|\mathbf{f}_c^{(t+1)}\right)$
- ▶ The whole proposal is

$$Q\left(\mathbf{f}^{(t+1)}|\mathbf{f}^{(t)}\right) = p\left(\mathbf{f}_\rho^{(t+1)}|\mathbf{f}_c^{(t+1)}\right) q\left(\mathbf{f}_c^{(t+1)}|\mathbf{f}_c^{(t)}\right)$$

- ▶ Its like sampling from the prior $p(\mathbf{f})$ but imposing random walk behaviour through the control points

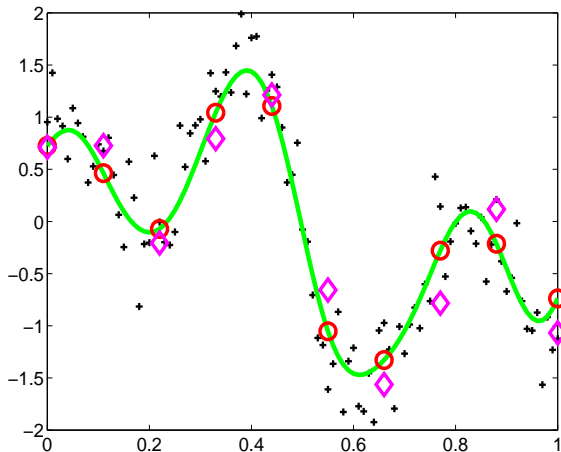
Sampling using control points: Regression-Examples

Sample 121 points using 10 control points



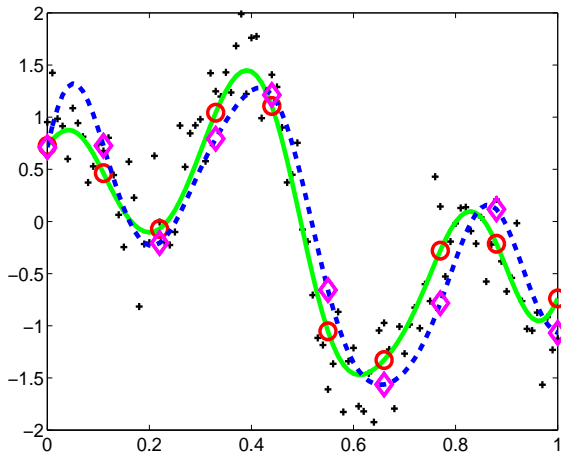
Sampling using control points: Regression-Examples

Sample 121 points using 10 control points



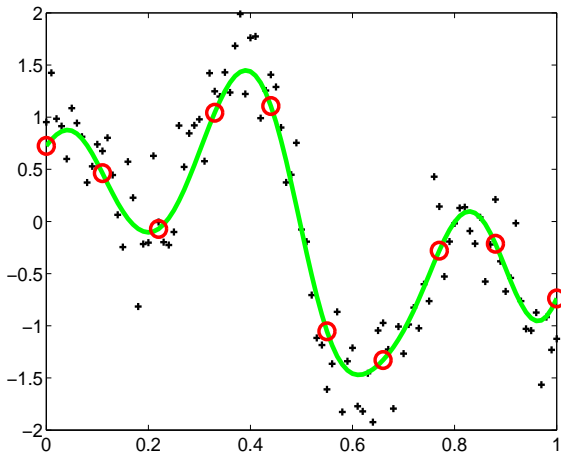
Sampling using control points: Regression-Examples

Sample 121 points using 10 control points



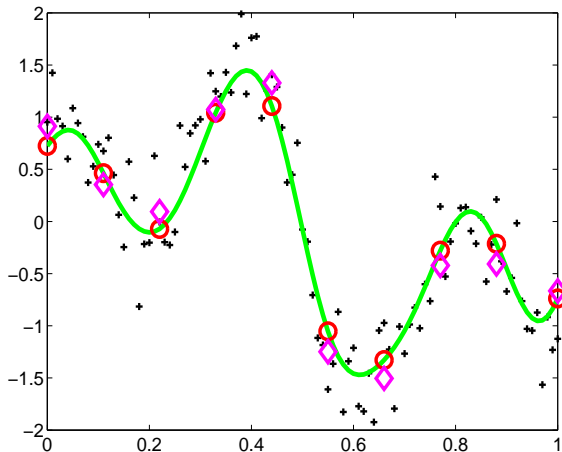
Sampling using control points: Regression-Examples

Sample 121 points using 10 control points



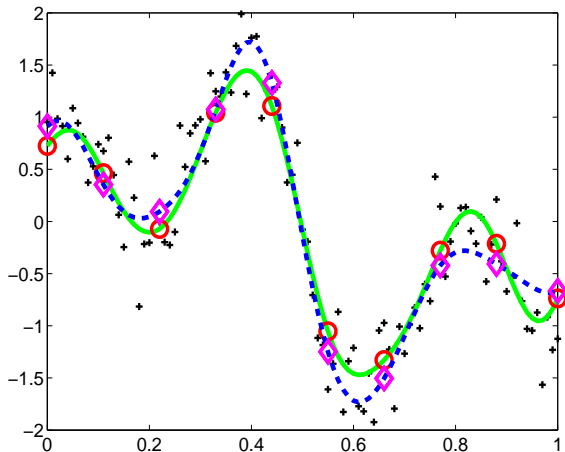
Sampling using control points: Regression-Examples

Sample 121 points using 10 control points



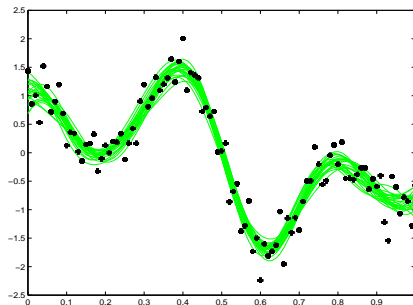
Sampling using control points: Regression-Examples

Sample 121 points using 10 control points



Sampling using control points

Few samples drawn during MCMC



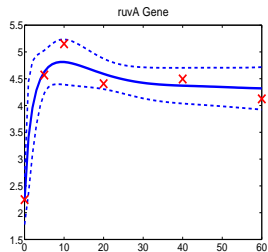
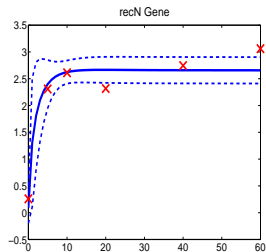
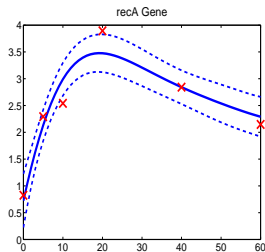
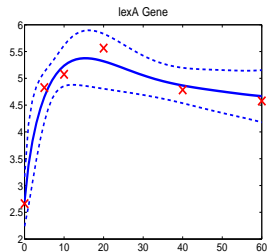
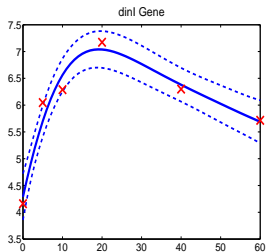
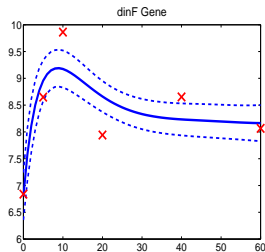
Results on SOS System

- ▶ Again consider the Michaelis-Menten kinetic equation

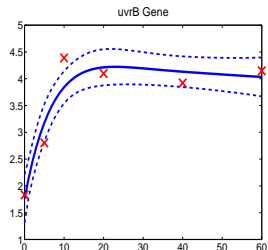
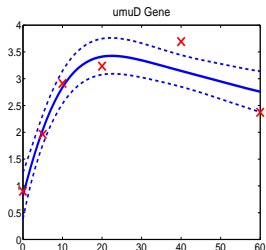
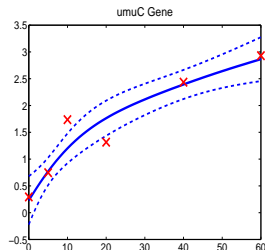
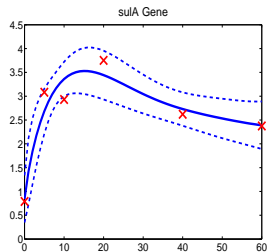
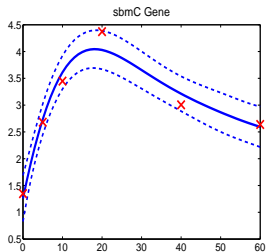
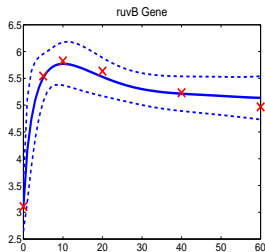
$$\frac{dx_j(t)}{dt} = B_j + S_j \frac{1}{\exp(f(t)) + \gamma_j} - D_j x_j(t)$$

- ▶ We have 14 genes (5 kinetic parameters each)
- ▶ Gene expressions are available for $T = 6$ time slots
- ▶ TF (\mathbf{f}) is discretized using 121 points
- ▶ MCMC details:
 - ▶ 6 control points are used (placed in a equally spaced grid)
 - ▶ Running time was 5 hours for 2 million sampling iterations plus burn in
 - ▶ Acceptance rate for \mathbf{f} after burn in was between 15% – 25%

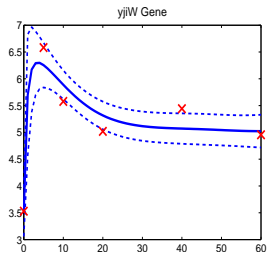
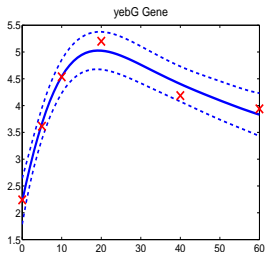
Results in E.coli data: Predicted gene expressions



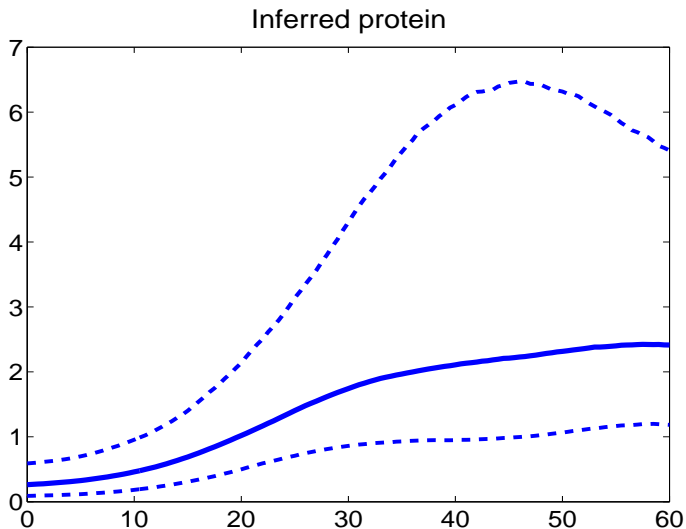
Results in E.coli data: Predicted gene expressions



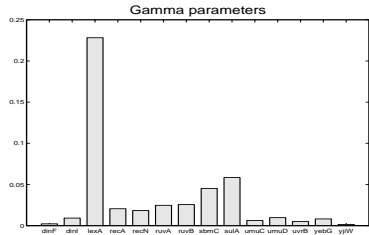
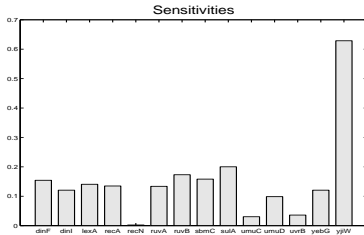
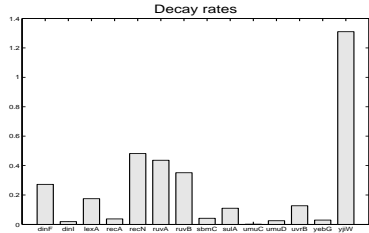
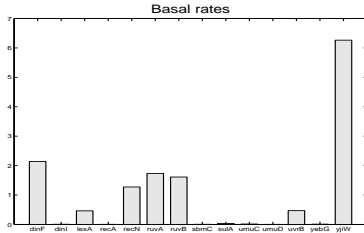
Results in E.coli data: Predicted gene expressions



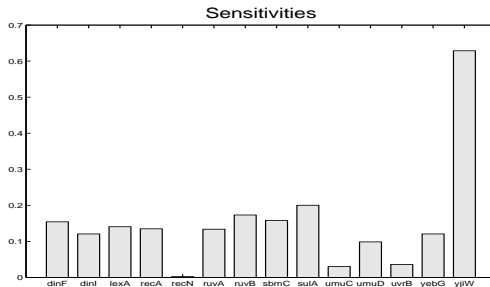
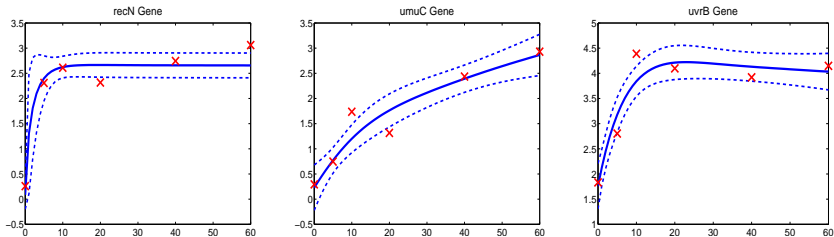
Results in E.coli data: Protein concentration



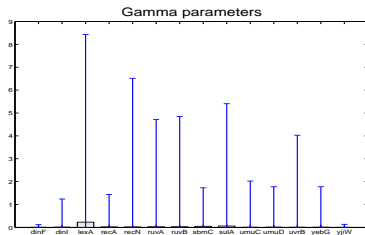
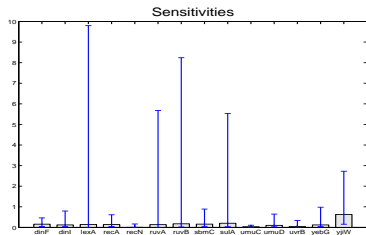
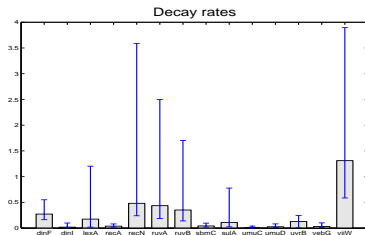
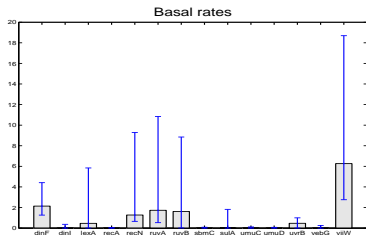
Results in E.coli data: Kinetic parameters



Results in E.coli data: Genes with low sensitivity value



Results in E.coli data: Confidence intervals for the kinetic parameters

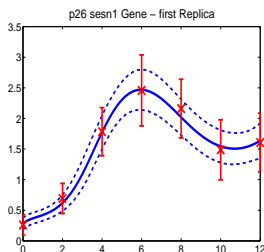
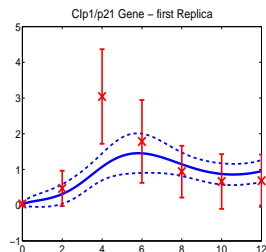
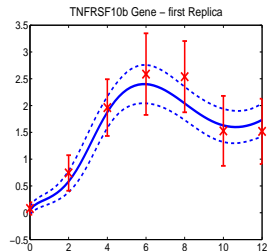
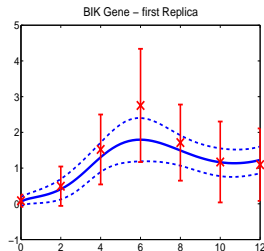
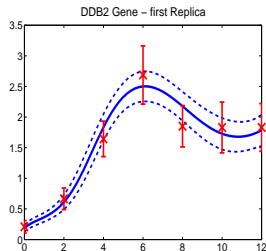


- ▶ One transcription factor (p53) that acts as an activator. We consider the Michaelis-Menten kinetic equation

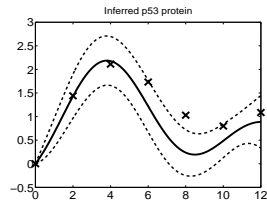
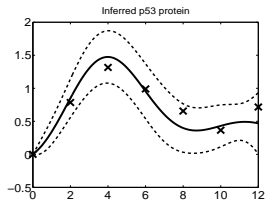
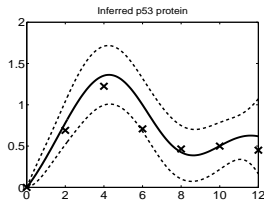
$$\frac{dx_j(t)}{dt} = B_j + S_j \frac{\exp(f(t))}{\exp(f(t)) + \gamma_j} - D_j x_j(t)$$

- ▶ We have 5 genes
- ▶ Gene expressions are available for $T = 7$ times and there are 3 replicas of the time series data
- ▶ TF (\mathbf{f}) is discretized using 121 points
- ▶ MCMC details:
 - ▶ 7 control points are used (placed in a equally spaced grid)
 - ▶ Running time 4/5 hours for 2 million sampling iterations plus burn in
 - ▶ Acceptance rate for \mathbf{f} after burn in was between 15% – 25%

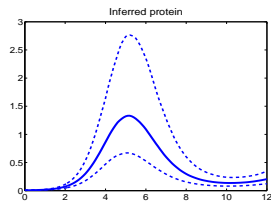
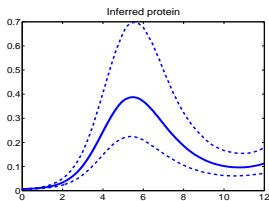
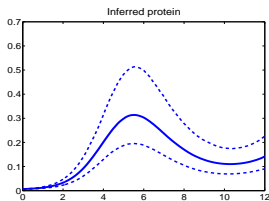
Data used by Barenco et al. (2006): Predicted gene expressions for the 1st replica



Data used by Barenco et al. (2006): Protein concentrations

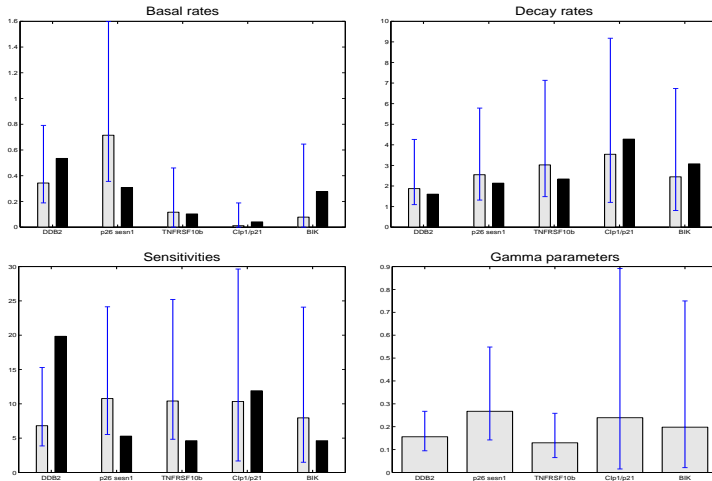


Linear model (Barenco et al. predictions are shown as crosses)



Nonlinear (Michaelis-Menten kinetic equation)

p53 Data Kinetic parameters



Our results (grey) compared with Barenco et al. (2006) (black).
Note that Barenco et al. use a linear model