

Latent Force Models with Gaussian Processes

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Outline

Motivation and Review

Covariance Functions

Motion Capture Example

ODE Model of Transcriptional Regulation

Cascade Differential Equations

Discussion and Future Work

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Styles of Machine Learning

- ▶ Urs Hölzle keynote talk at NIPS 2005.
 - ▶ In the future NIPS ML algorithms will have to run on massive data sets.
 - ▶ Let the data do the work!
- ▶ Alternative paradigm:
 - ▶ Very scarce data: computational biology, human motion.
 - ▶ How to generalize from scarce data?

General Approach

- Two approaches to modeling.

data modeling

mechanistic modeling



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data modeling

classification (SVM)

mechanistic modeling

differential equations



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dimensionality reduction (PCA)

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climate, weather models



General Approach

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dimensionality reduction (PCA)

clustering

mechanistic modeling

differential equations

climate, weather models

finite element analysis



Dimensionality Reduction

- ▶ Linear relationship between the data, $\mathbf{X} \in \mathbb{R}^{N \times d}$, and a reduced dimensional representation, $\mathbf{F} \in \mathbb{R}^{N \times q}$, where $q \ll d$.

$$\mathbf{X} = \mathbf{F}\mathbf{W} + \epsilon,$$

$$\epsilon \sim \mathcal{N}(\mathbf{0}, \Sigma)$$

- ▶ Integrate out $\mathbf{F}$, optimize with respect to $\mathbf{W}$.
- ▶ For Gaussian prior, $\mathbf{F} \sim \mathcal{N}(\mathbf{0}, \mathbf{I})$
 - ▶ and $\Sigma = \sigma^2 \mathbf{I}$ we have probabilistic PCA (Tipping and Bishop, 1999).
 - ▶ and Σ constrained to be diagonal, we have factor analysis.

Dimensionality Reduction: Temporal Data

- ▶ Deal with temporal data with a temporal latent prior.
- ▶ Independent Gauss-Markov priors over each $f_i(t)$ leads to : Rauch-Tung-Striebel (RTS) smoother (Kalman filter).
- ▶ More generally consider a Gaussian process (GP) prior,

$$p(\mathbf{F}|\mathbf{t}) = \prod_{i=1}^q \mathcal{N}(\mathbf{f}_{:,i} | \mathbf{0}, \mathbf{K}_{f_{:,i}, f_{:,i}}).$$

- ▶ Given the covariance functions for $\{f_i(t)\}$ we have an implied covariance function across all $\{x_i(t)\}$ —(ML: semi-parametric latent factor model (Teh et al., 2005), Geostatistics: linear model of coregionalization).
- ▶ Rauch-Tung-Striebel smoother has been preferred
 - ▶ linear computational complexity in N .
 - ▶ Advances in sparse approximations have made the general GP framework practical. (Titsias, 2009; Snelson and Ghahramani, 2006; Quiñero Candela and Rasmussen, 2005).

Zero mean Gaussian distribution

- ▶ A multi-variate Gaussian distribution is defined by a mean and a covariance matrix.

$$N(\mathbf{f}|\mu, \mathbf{K}) = \frac{1}{(2\pi)^{\frac{N}{2}} |\mathbf{K}|^{\frac{1}{2}}} \exp\left(-\frac{(\mathbf{f} - \mu)^T \mathbf{K}^{-1} (\mathbf{f} - \mu)}{2}\right).$$

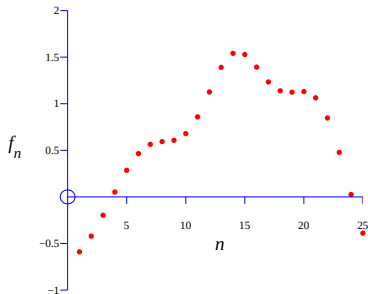
- ▶ We will consider the special case where the mean is zero,

$$N(\mathbf{f}|\mathbf{0}, \mathbf{K}) = \frac{1}{(2\pi)^{\frac{N}{2}} |\mathbf{K}|^{\frac{1}{2}}} \exp\left(-\frac{\mathbf{f}^T \mathbf{K}^{-1} \mathbf{f}}{2}\right).$$

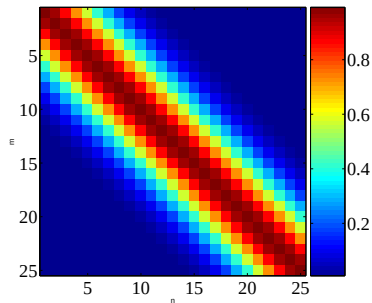
Multi-variate Gaussians

- ▶ We will consider a Gaussian with a particular structure of covariance matrix.
- ▶ Generate a single sample from this 25 dimensional Gaussian distribution, $\mathbf{f} = [f_1, f_2 \dots f_{25}]$.
- ▶ We will plot these points against their index.

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*.
- ▶ Combine covariance function with training data by conditioning on one point given the other.
- ▶ Let's consider the marginal density for variables indexed by 1 and 2.

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Prediction of f_2 from f_1

demGpCov2D([1 2])

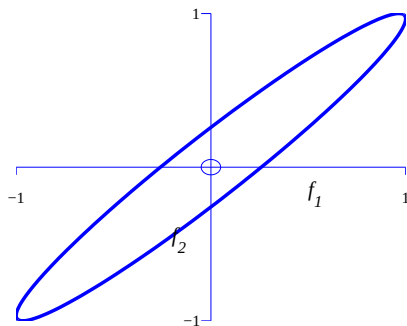


Figure: Covariance for $\begin{bmatrix} f_1 \\ f_2 \end{bmatrix}$ is $\mathbf{K}_{12} = \begin{bmatrix} 1 & 0.966 \\ 0.966 & 1 \end{bmatrix}$.

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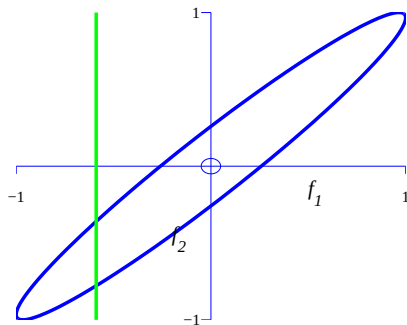


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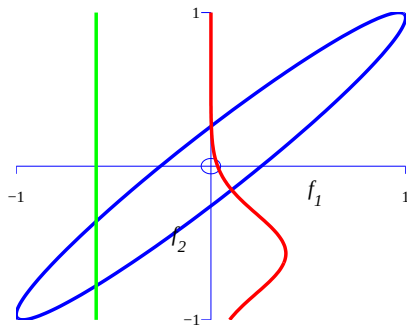


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Prediction of f_5 from f_1

demGpCov2D([1 5])

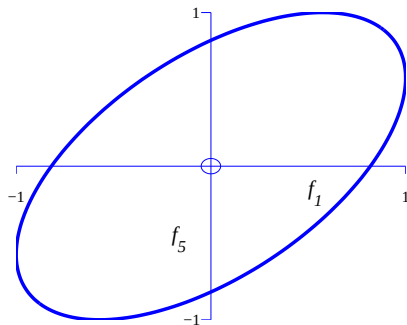


Figure: Covariance for $\begin{bmatrix} f_1 \\ f_5 \end{bmatrix}$ is $\mathbf{K}_{15} = \begin{bmatrix} 1 & 0.574 \\ 0.574 & 1 \end{bmatrix}$.

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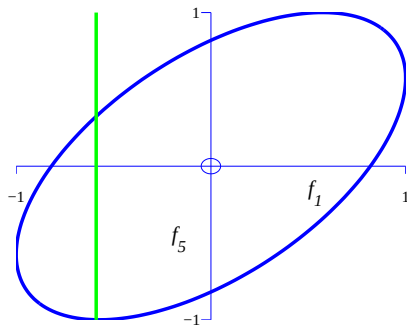


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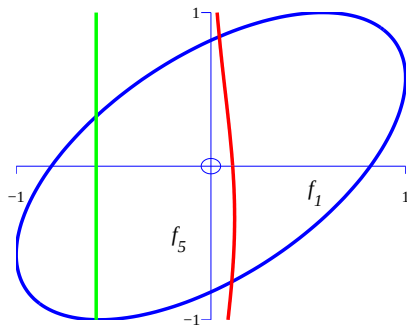


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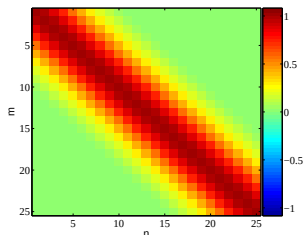
Covariance Functions

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

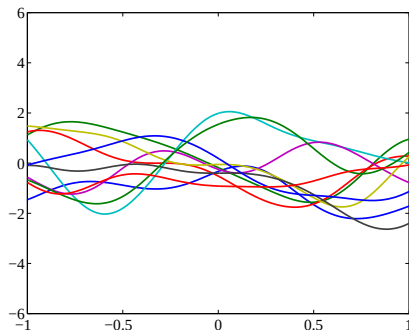


Figure: RBF kernel with $\ell = 10^{-\frac{1}{2}}$, $\alpha = 1$

Covariance Samples

demCovFuncSample

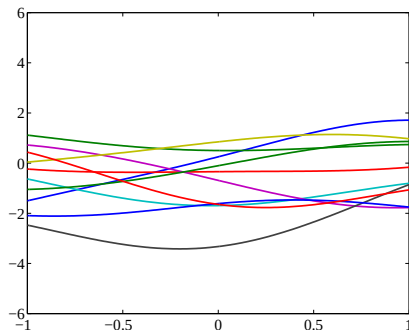


Figure: RBF kernel with $\ell = 1$, $\alpha = 1$

Covariance Samples

demCovFuncSample

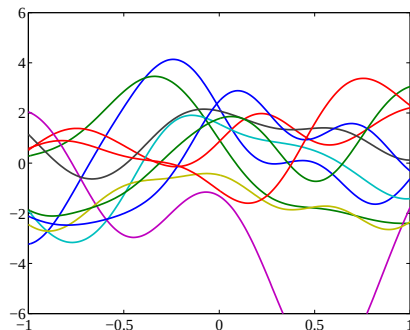


Figure: RBF kernel with $\ell = 0.3$, $\alpha = 4$

Covariance Samples

demCovFuncSample

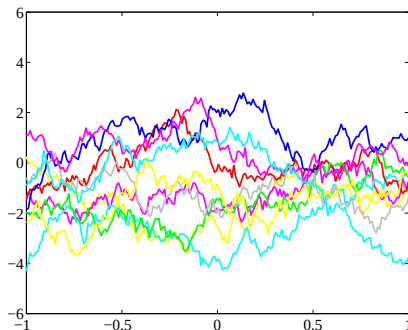


Figure: Ornstein-Uhlenbeck (stationary Gauss-Markov) covariance function $\ell = 1$, $\alpha = 4$

Back to Latent Force Models!

- ▶ These models rely on the latent variables to provide the dynamic information.
- ▶ We now introduce a further dynamical system with a *mechanistic* inspiration.
- ▶ Physical Interpretation:
 - ▶ the latent functions, $f_i(t)$ are q forces.
 - ▶ We observe the displacement of d springs to the forces.,
 - ▶ Interpret system as the force balance equation, $\mathbf{X}\mathbf{D} = \mathbf{F}\mathbf{S} + \epsilon$.
 - ▶ Forces act, e.g. through levers — a matrix of sensitivities, $\mathbf{S} \in \mathbb{R}^{q \times d}$.
 - ▶ Diagonal matrix of spring constants, $\mathbf{D} \in \mathbb{R}^{d \times d}$.
 - ▶ Original System: $\mathbf{W} = \mathbf{S}\mathbf{D}^{-1}$.

- Add a damper and give the system mass.

$$\mathbf{F}\mathbf{S} = \ddot{\mathbf{X}}\mathbf{M} + \dot{\mathbf{X}}\mathbf{C} + \mathbf{X}\mathbf{D} + \epsilon.$$

- Now have a second order mechanical system.
- It will exhibit inertia and resonance.
- There are many systems that can also be represented by differential equations.
 - When being forced by latent function(s), $\{f_i(t)\}_{i=1}^q$, we call this a *latent force model*.

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Gaussian Process priors and Latent Force Models

Driven Harmonic Oscillator

- ▶ For Gaussian process we can compute the covariance matrices for the output displacements.
- ▶ For one displace the model is

$$m_k \ddot{x}_k(t) + c_k \dot{x}_k(t) + d_k x_k(t) = b_k + \sum_{i=0}^M s_{ik} f_i(t), \quad (1)$$

where, m_k is the k th diagonal element from $\mathbf{M}$ and similarly for c_k and d_k . s_{ik} is the i , k th element of $\mathbf{S}$.

- ▶ Model the latent forces as q independent, GPs with RBF covariances

$$k_{f_i f_l}(t, t') = \exp \left(-\frac{(t - t')^2}{\sigma_i^2} \right) \delta_{il}.$$

Covariance for ODE Model

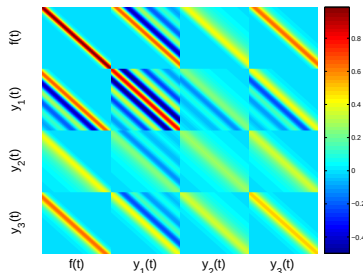
- RBF Kernel function for $f(t)$

$$x_j(t) = \frac{1}{m_j \omega_j} \sum_{i=1}^q S_{ji} \exp(-\alpha_j t) \int_0^t f_i(u) \exp(\alpha_j u) \sin(\omega_j(t-u)) du$$

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

Damping ratios:

ζ_1	ζ_2	ζ_3
0.125	2	1



Covariance for ODE Model

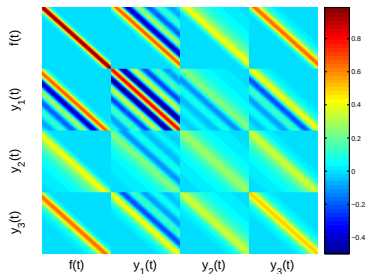
► Analogy

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

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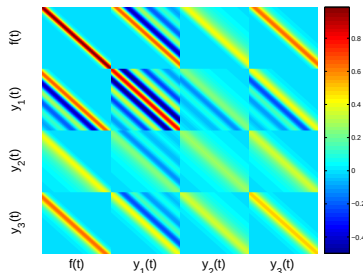
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Joint Sampling of $x(t)$ and $f(t)$

► lfmSample

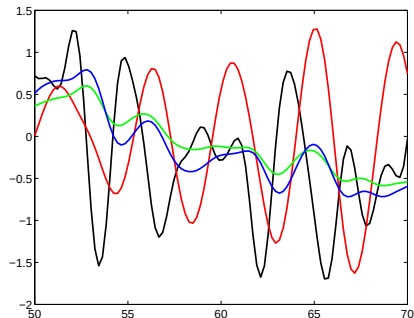


Figure: Joint samples from the ODE covariance, *black*: $f(t)$, *red*: $x_1(t)$ (underdamped), *green*: $x_2(t)$ (overdamped), and *blue*: $x_3(t)$ (critically damped).

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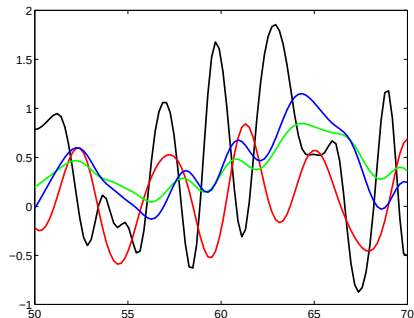


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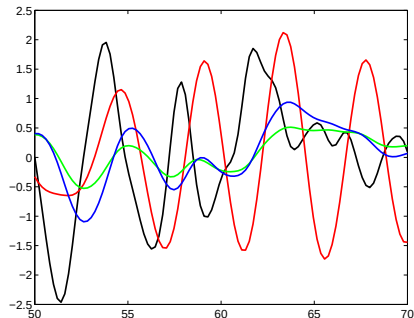


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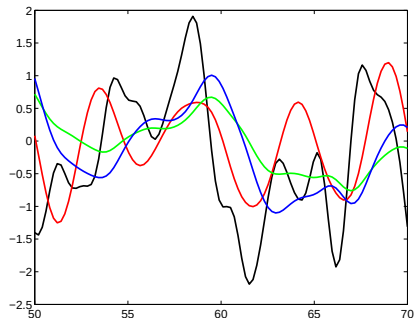


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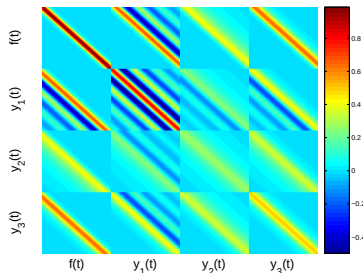
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Mauricio Alvarez and David Luengo (Álvarez et al., 2009)

- ▶ Motion capture data: used for animating human motion.
- ▶ Multivariate time series of angles representing joint positions.
- ▶ Objective: generalize from training data to realistic motions.
- ▶ Use 2nd Order Latent Force Model with mass/spring/damper (resistor inductor capacitor) at each joint.

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Prediction of Test Motion

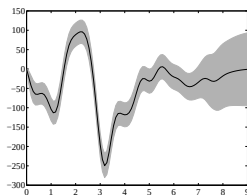
- ▶ Model left arm only.
- ▶ 3 balancing motions (18, 19, 20) from subject 49.
- ▶ 18 and 19 are similar, 20 contains more dramatic movements.
- ▶ Train on 18 and 19 and testing on 20
- ▶ Data was down-sampled by 32 (from 120 fps).
- ▶ Reconstruct motion of left arm for 20 given other movements.
- ▶ Compare with GP that predicts left arm angles given other body angles.

Mocap Results

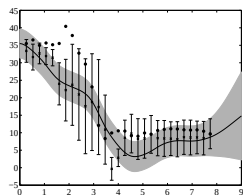
Table: Root mean squared (RMS) angle error for prediction of the left arm's configuration in the motion capture data. Prediction with the latent force model outperforms the prediction with regression for all apart from the radius's angle.

Angle	Latent Force Error	Regression Error
Radius	4.11	4.02
Wrist	6.55	6.65
Hand X rotation	1.82	3.21
Hand Z rotation	2.76	6.14
Thumb X rotation	1.77	3.10
Thumb Z rotation	2.73	6.09

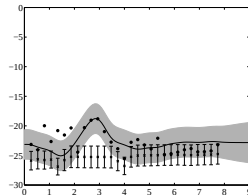
Mocap Results II



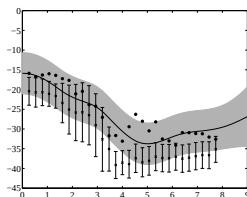
(a) Inferred Latent Force



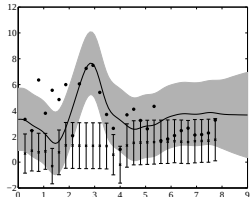
(b) Wrist



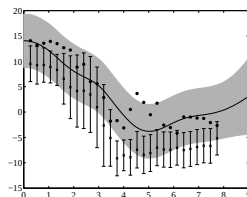
(c) Hand X Rotation



(d) Hand Z Rotation



(e) Thumb X Rotation



(f) Thumb Z Rotation

Figure: Predictions from LFM (solid line, grey error bars) and direct regression (crosses with stick error bars).

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Example: Transcriptional Regulation

- ▶ First Order Differential Equation

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

- ▶ Can be used as a model of gene transcription: Barenco et al., 2006; Gao et al., 2008.
- ▶ $x_j(t)$ – concentration of gene j 's mRNA
- ▶ $f(t)$ – concentration of active transcription factor
- ▶ Model parameters: baseline B_j , sensitivity S_j and decay D_j
- ▶ Application: identifying co-regulated genes (targets)
- ▶ Problem: how do we fit the model when $f(t)$ is not observed?

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- Problem: how do we fit the model when $f(t)$ is not observed?

Example: Transcriptional Regulation

- First Order Differential Equation

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

- Can be used as a model of gene transcription: Barenco et al., 2006; Gao et al., 2008.
- $x_j(t)$ – concentration of gene j 's mRNA
- $f(t)$ – concentration of active transcription factor
- Model parameters: baseline B_j , sensitivity S_j and decay D_j
- Application: identifying co-regulated genes (targets)
- Problem: how do we fit the model when $f(t)$ is not observed?

Covariance for Transcription Model

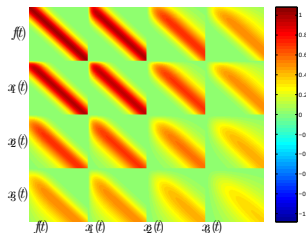
RBF covariance function for $f(t)$

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

- 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`

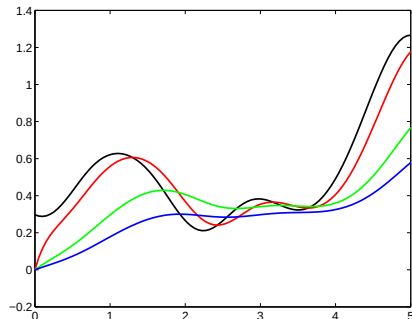


Figure: Joint samples from the ODE covariance, *black*: $f(t)$, *red*: $x_1(t)$ (high decay/sensitivity), *green*: $x_2(t)$ (medium decay/sensitivity) and *blue*: $x_3(t)$ (low decay/sensitivity).

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

► `simSample`

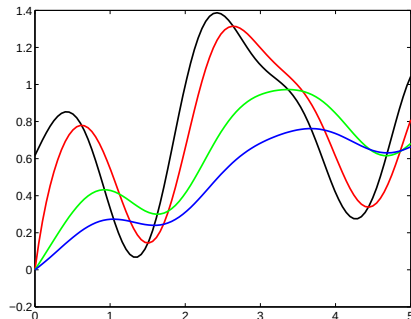


Figure: Joint samples from the ODE covariance, *black*: $f(t)$, *red*: $x_1(t)$ (high decay/sensitivity), *green*: $x_2(t)$ (medium decay/sensitivity) and *blue*: $x_3(t)$ (low decay/sensitivity).

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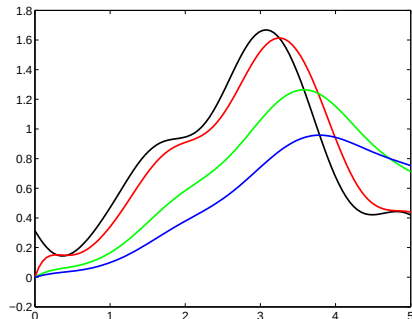


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Joint Sampling of $f(t)$ and $x(t)$

► `simSample`

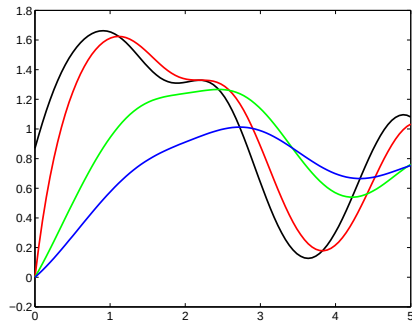
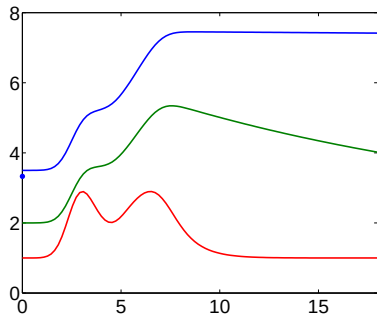


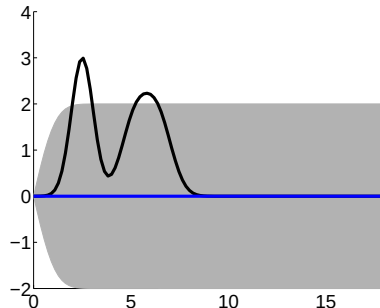
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Artificial Example: Inferring $f(t)$

Inferring TF activity from artificially sampled genes.



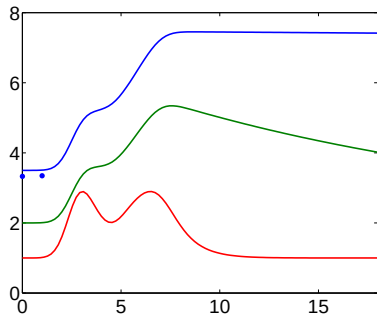
True “gene profiles” and noisy observations.



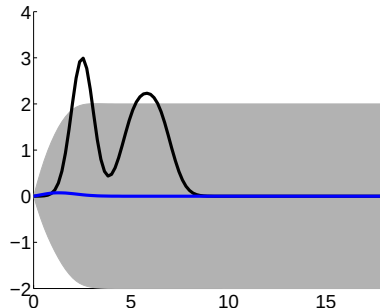
Inferred transcription factor activity.

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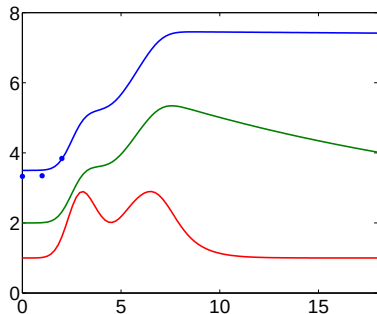
True “gene profiles” and noisy observations.



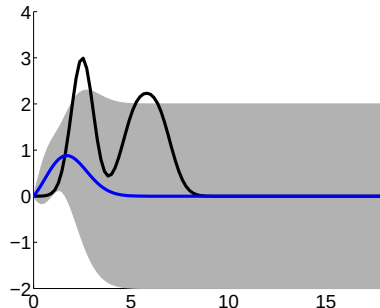
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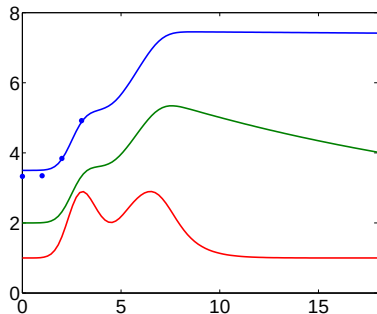
True “gene profiles” and noisy observations.



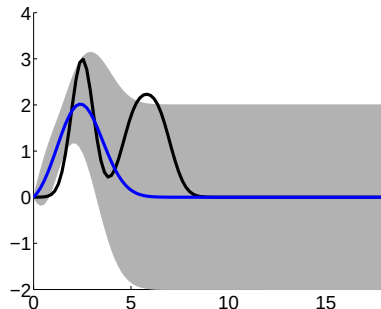
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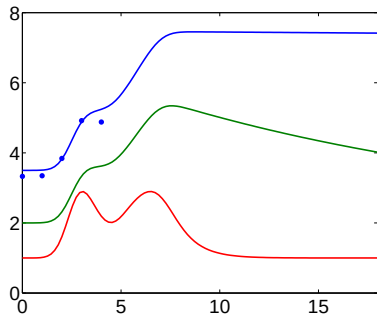
True “gene profiles” and noisy observations.



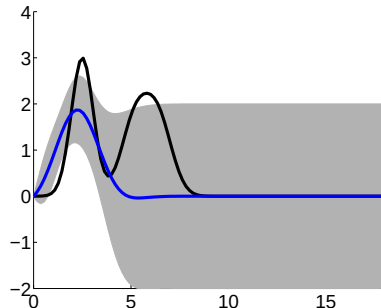
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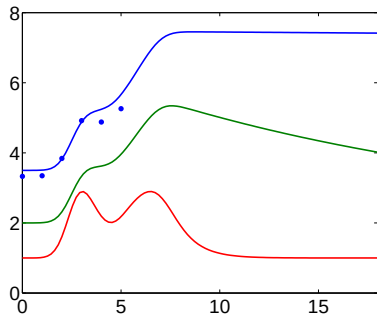
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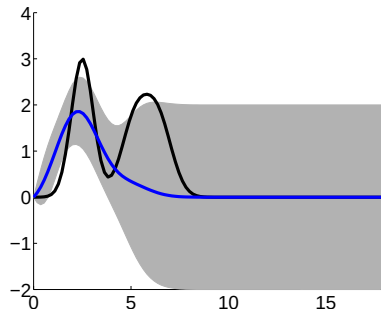
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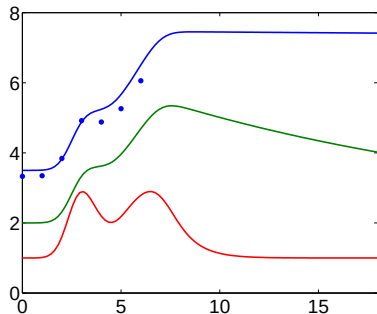
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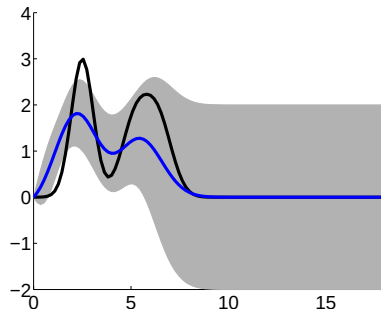
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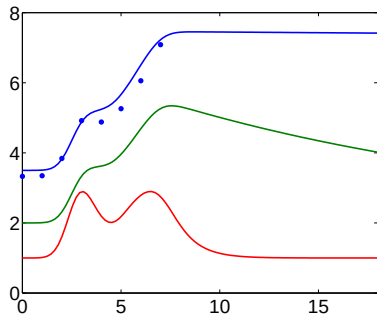
True “gene profiles” and noisy observations.



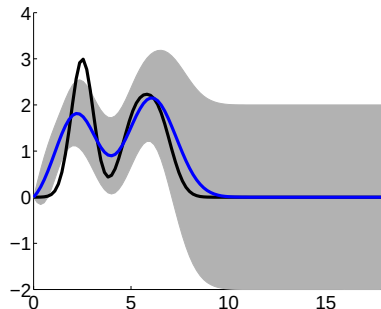
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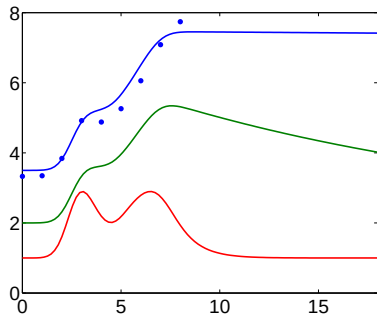
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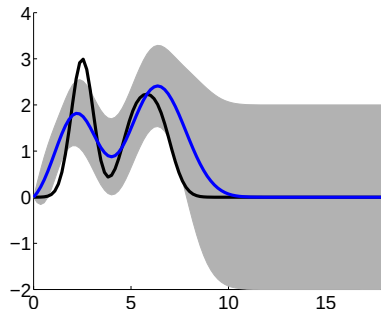
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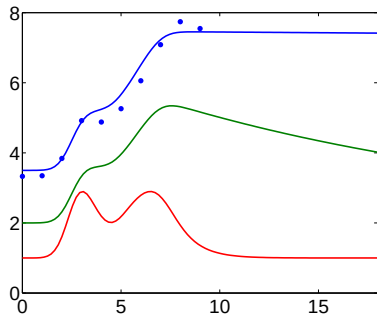
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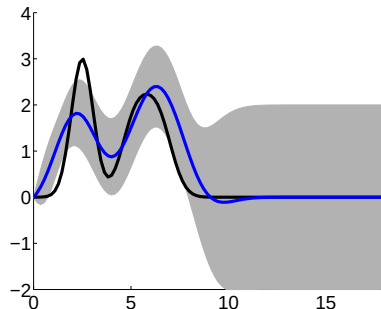
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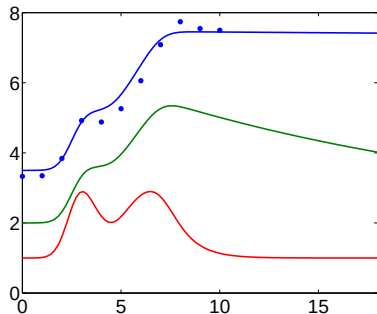
True “gene profiles” and noisy observations.



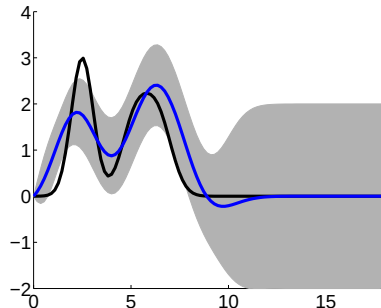
Inferred transcription factor activity.

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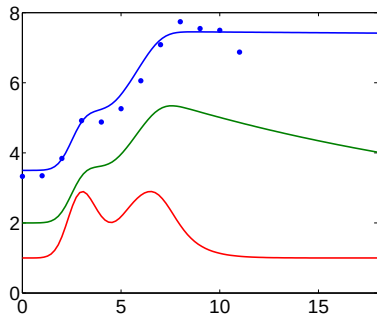
True “gene profiles” and noisy observations.



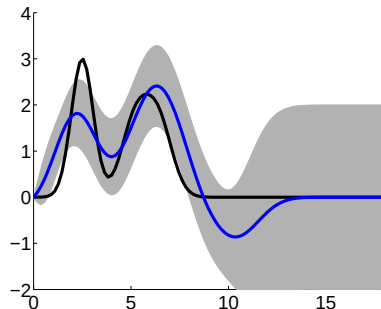
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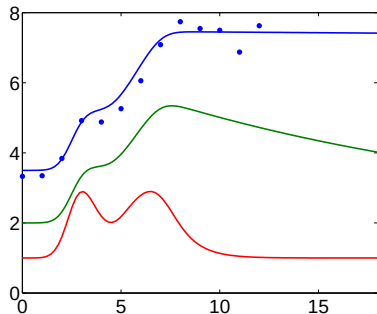
True “gene profiles” and noisy observations.



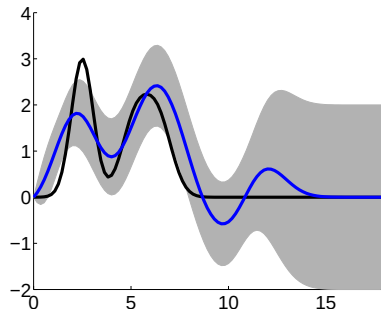
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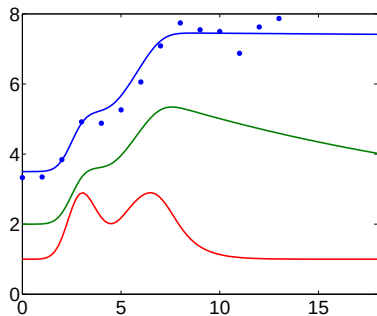
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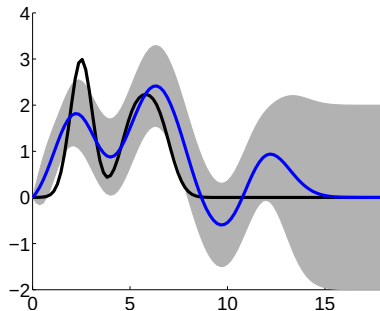
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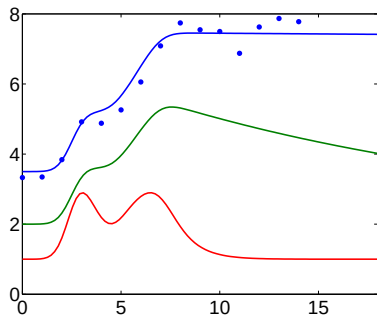
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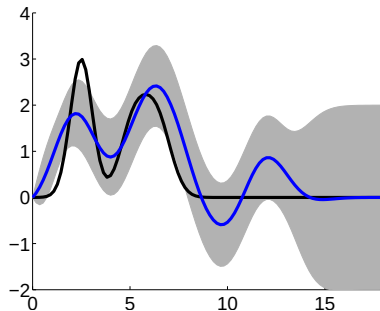
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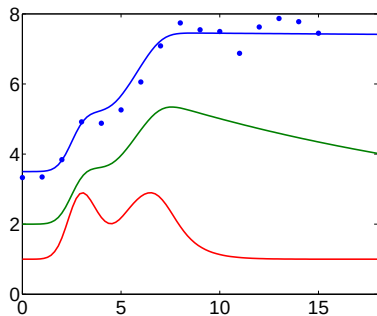
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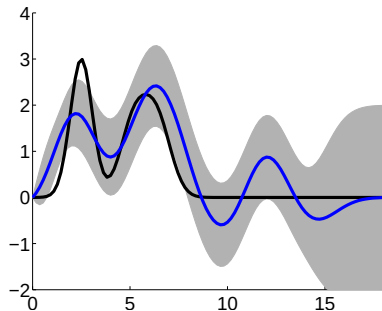
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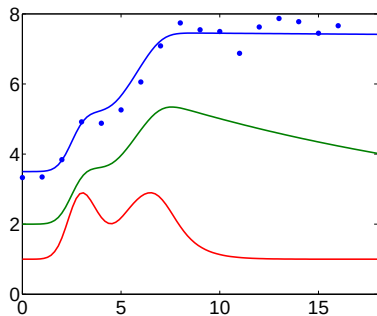
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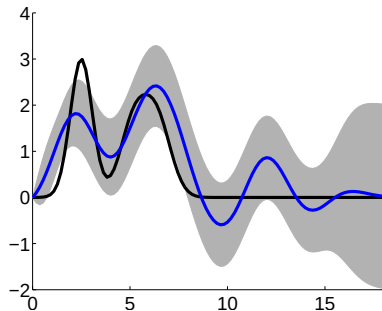
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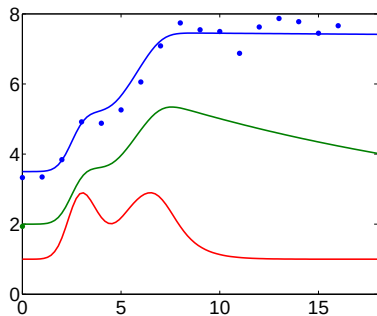
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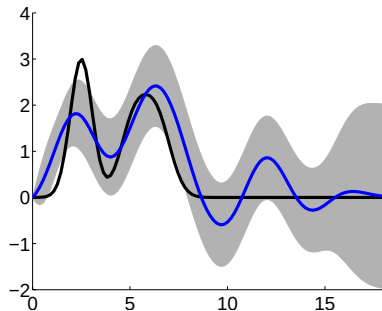
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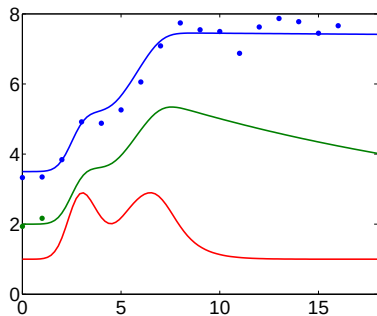
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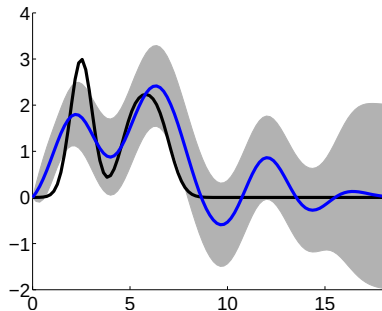
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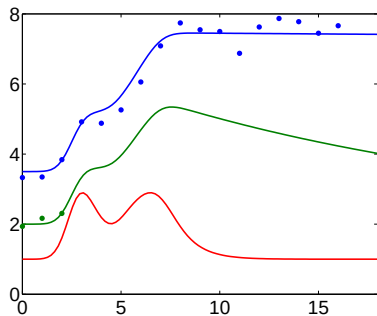
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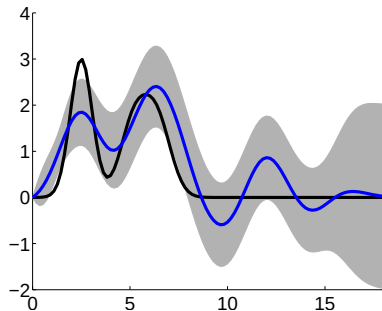
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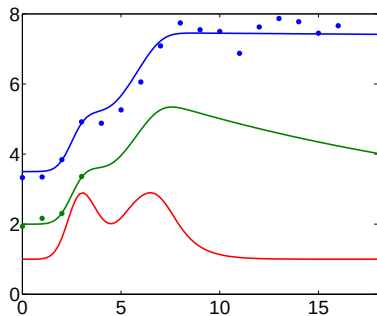
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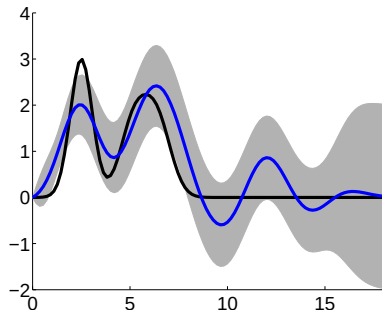
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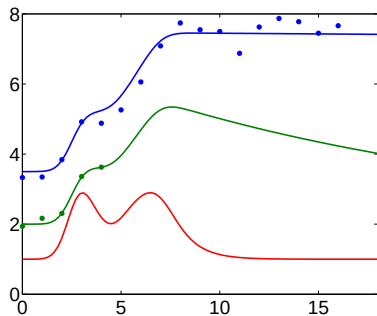
True “gene profiles” and noisy observations.



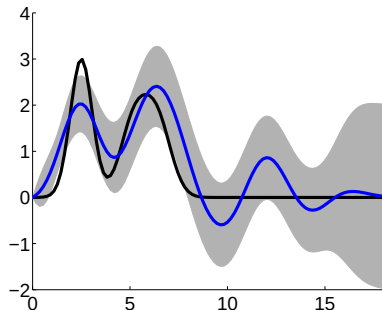
Inferred transcription factor activity.

Artificial Example: Inferring $f(t)$

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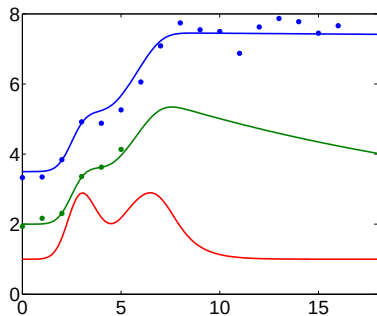
True “gene profiles” and noisy observations.



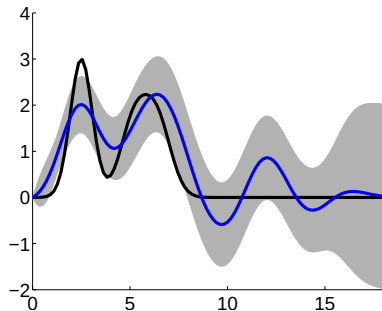
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Inferring TF activity from artificially sampled genes.



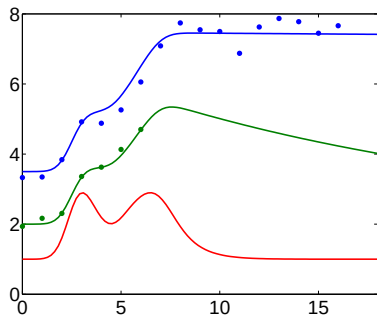
True “gene profiles” and noisy observations.



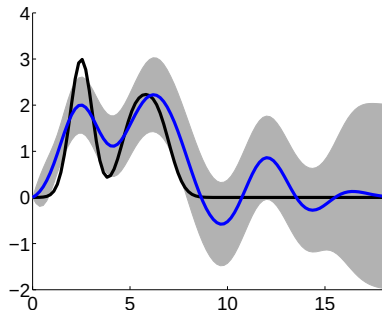
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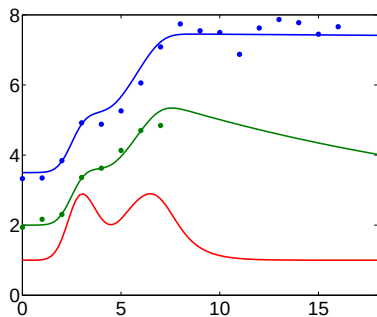
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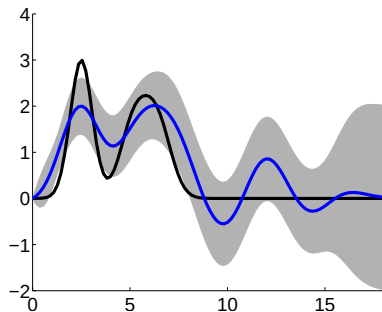
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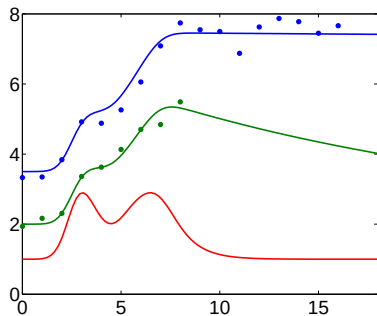
True “gene profiles” and noisy observations.



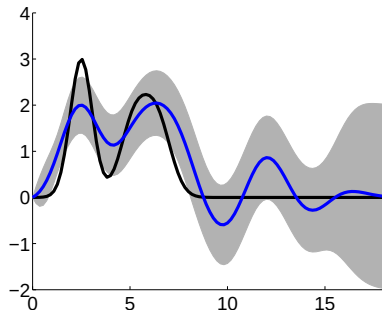
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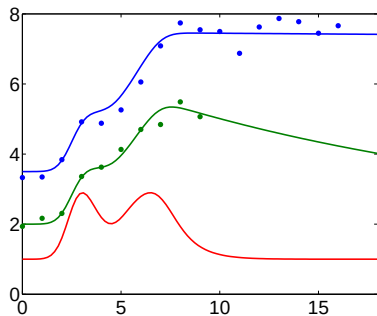
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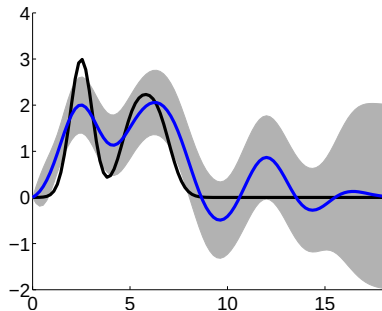
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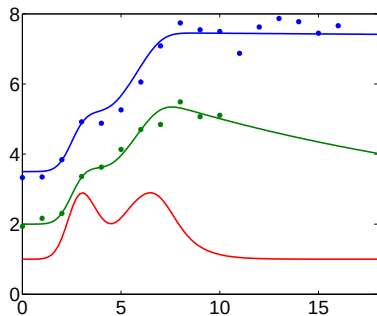
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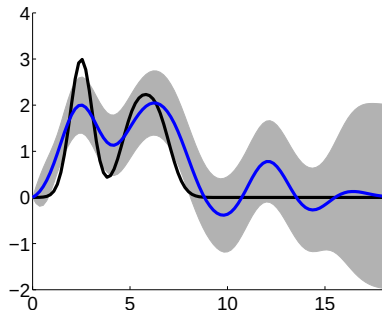
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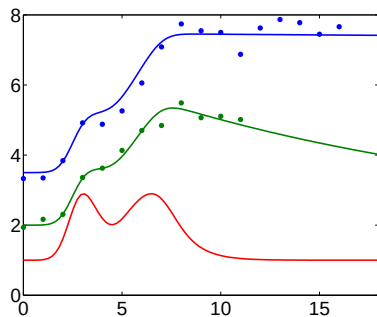
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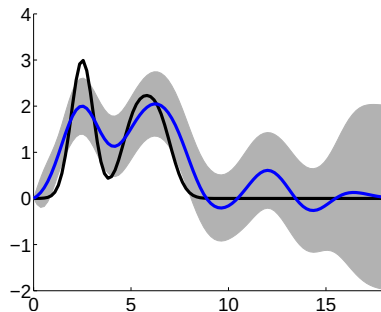
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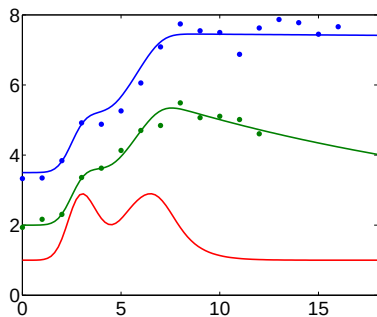
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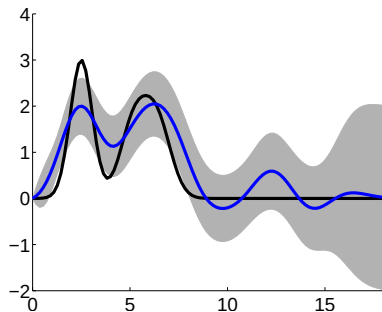
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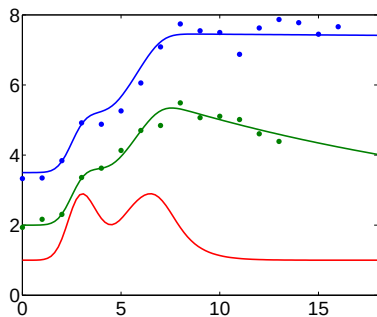
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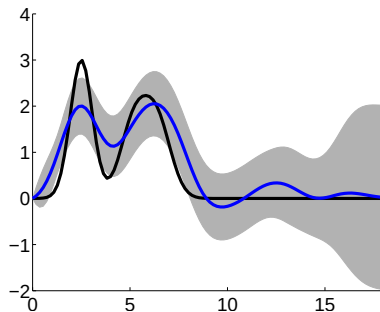
Inferred transcription factor activity.

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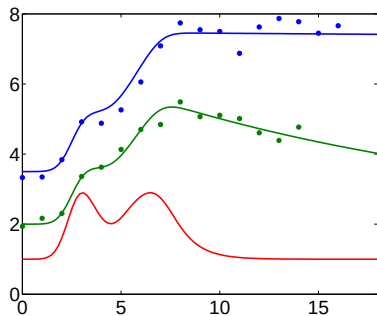
True “gene profiles” and noisy observations.



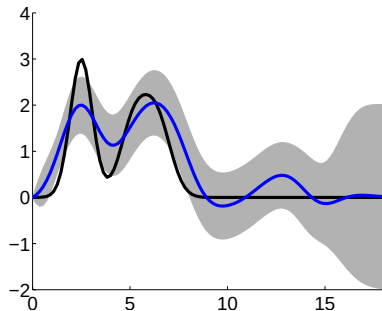
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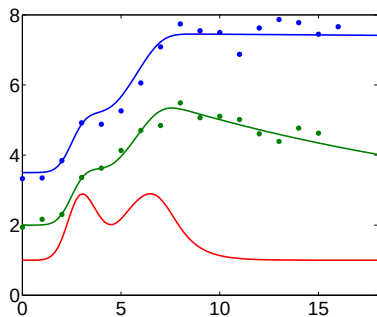
True “gene profiles” and noisy observations.



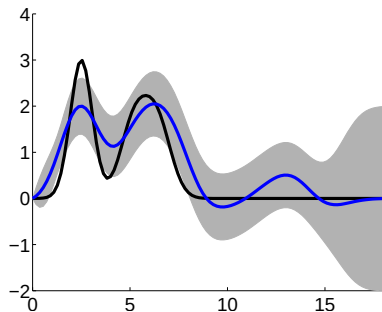
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Inferring TF activity from artificially sampled genes.



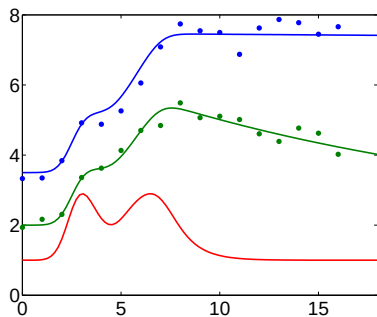
True “gene profiles” and noisy observations.



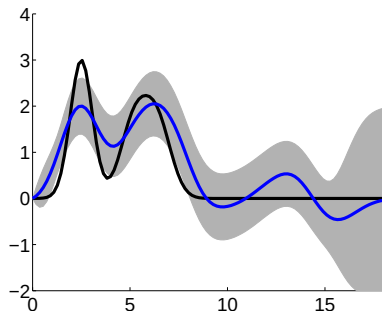
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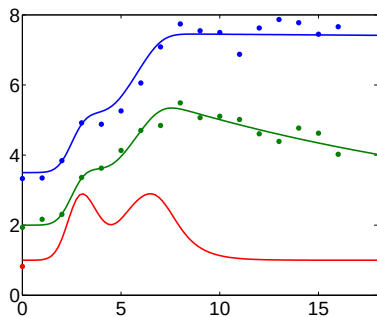
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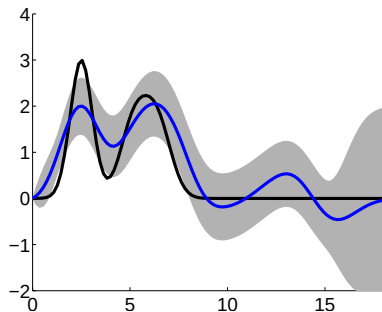
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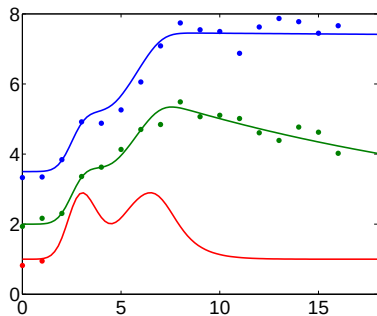
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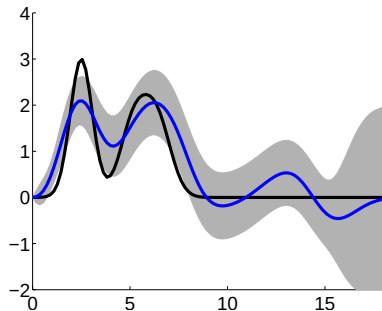
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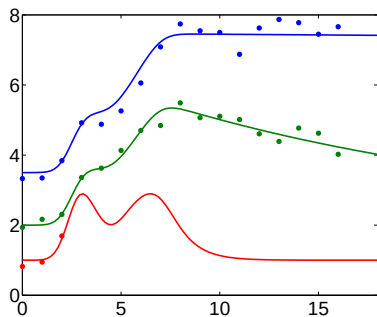
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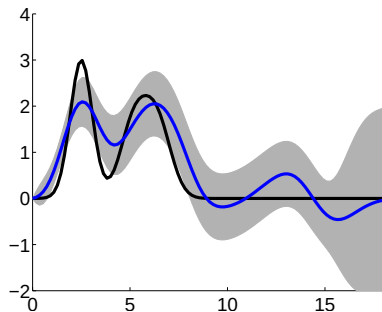
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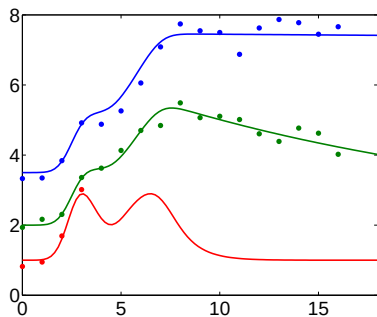
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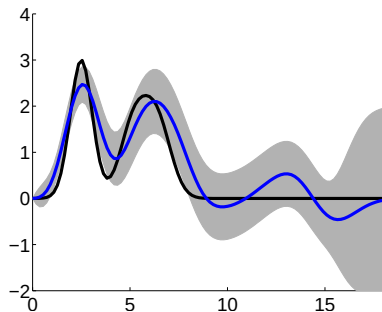
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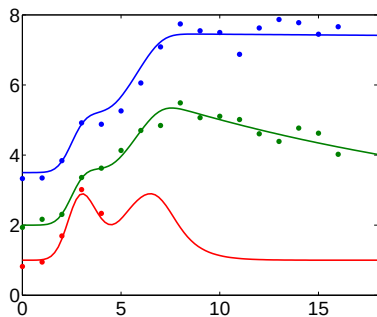
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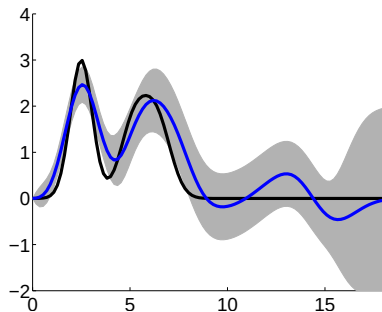
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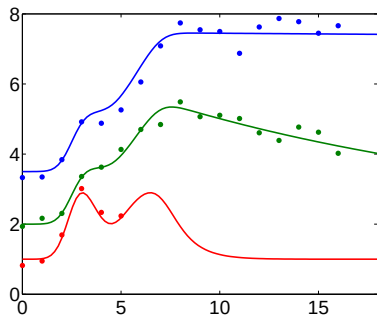
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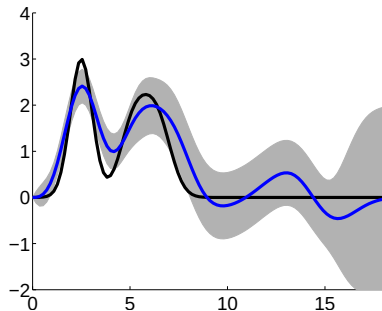
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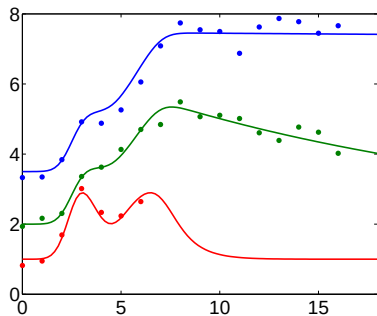
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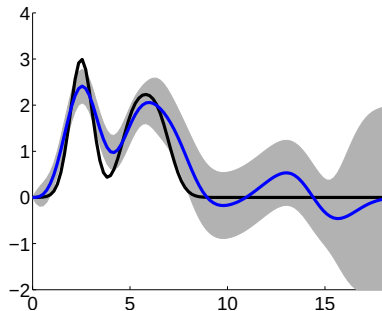
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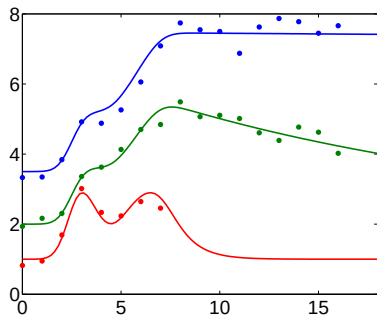
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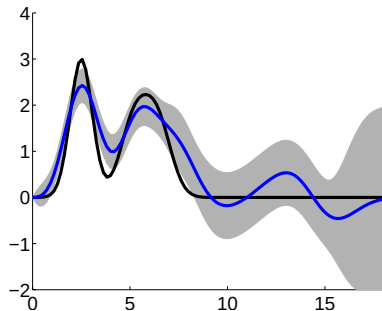
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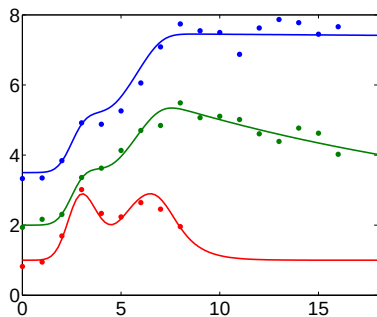
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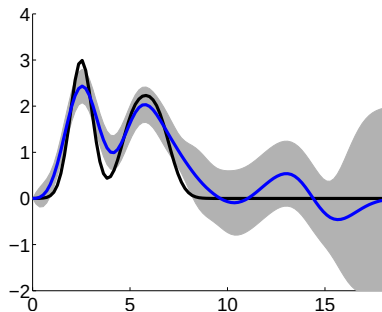
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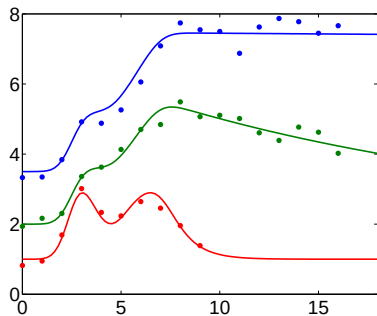
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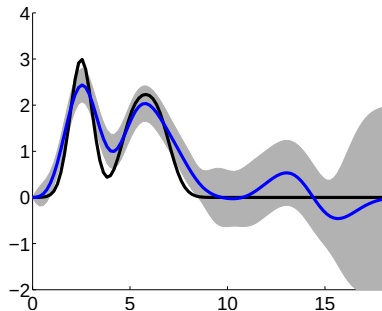
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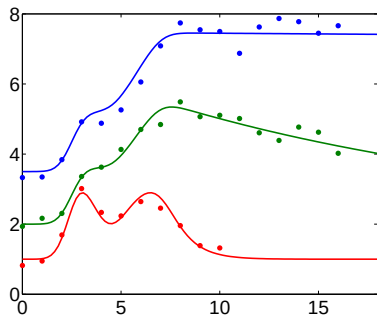
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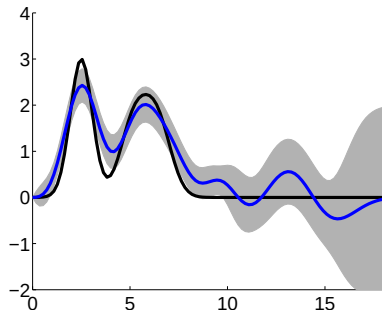
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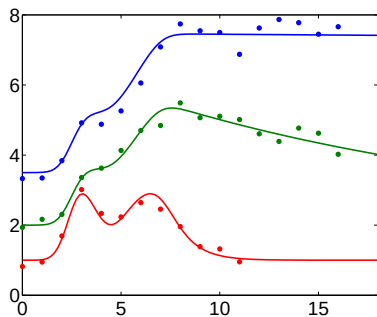
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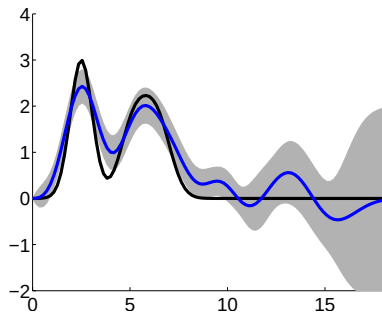
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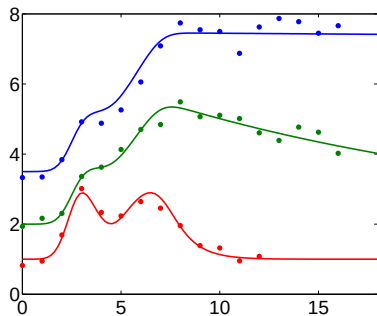
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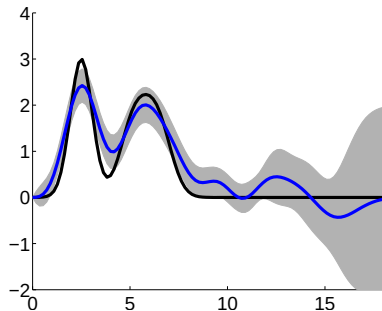
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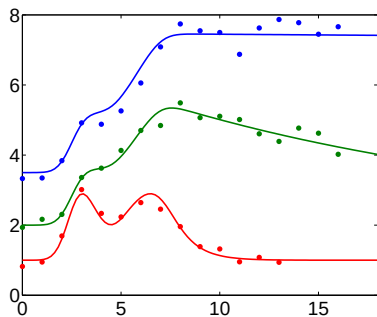
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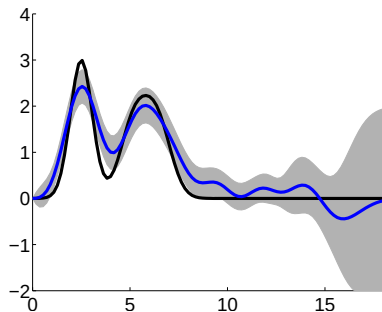
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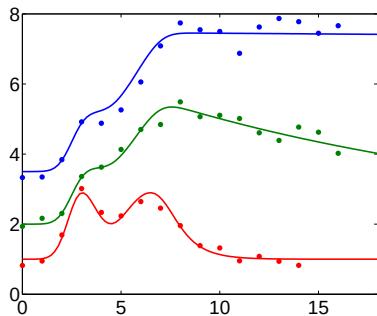
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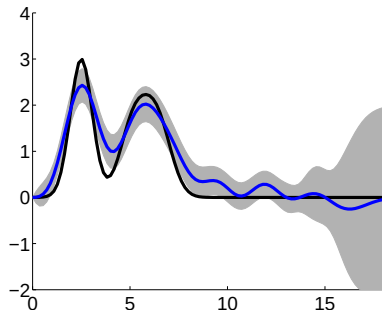
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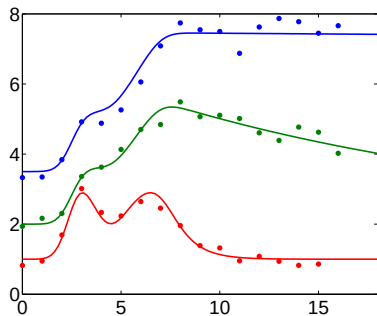
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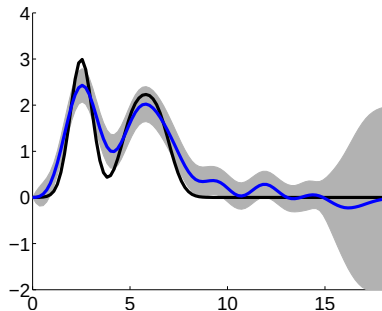
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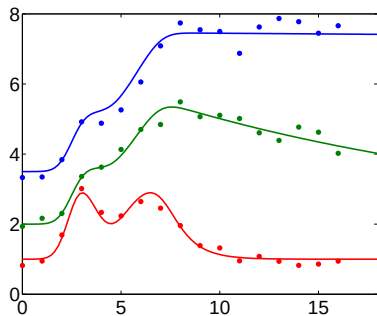
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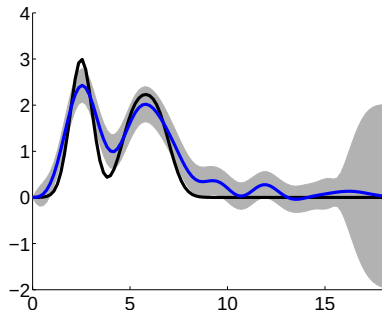
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True “gene profiles” and noisy observations.



Inferred transcription factor activity.

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

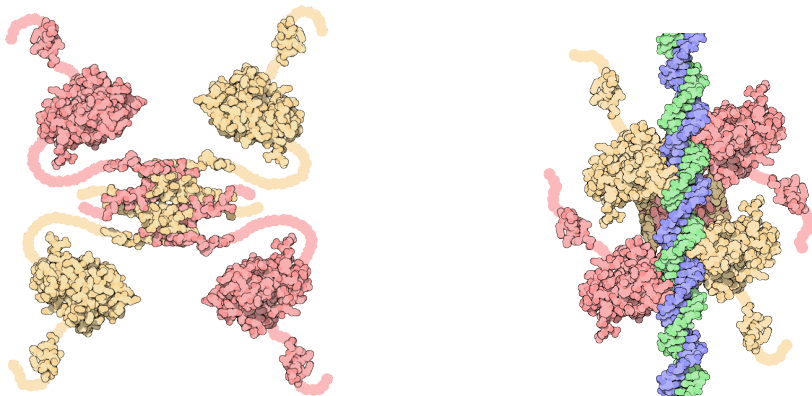


Figure: p53. *Left* unbound, *Right* bound to DNA. Images by David S. Goodsell from <http://www.rcsb.org/> (see the “Molecule of the Month” feature).

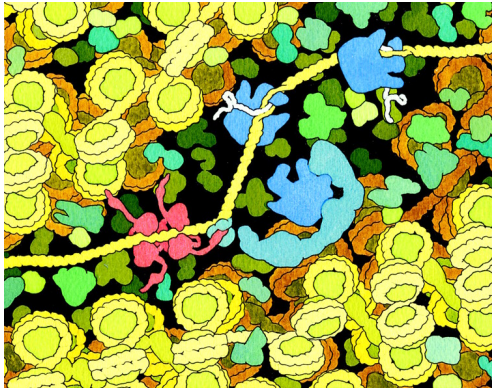


Figure: Repair of DNA damage by p53. Image from Goodsell (1999).

Response of p53 to Ionizing Radiation

- ▶ Experiment by Barenco et al. 2006.
- ▶ Human leukemia cell line (MOLT4) containing functional p53 and harvested protein and RNA at regular intervals after irradiation.
- ▶ The time course was performed in triplicate, and mRNA concentrations measured using Affymetrix U133A microarrays.

Modelling Assumption

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

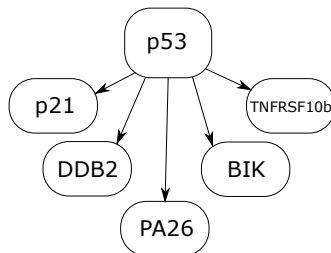


Figure: p53 SIM network motif as modelled by Barenco et al. 2006.

Ordinary Differential Equation Model

- First Order Differential Equation

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

- Proposed by Barenco et al. (2006).
- $x_j(t)$ – concentration of gene j 's mRNA
- $f(t)$ – concentration of active transcription factor
- Model parameters: baseline B_j , sensitivity S_j and decay D_j
- Application: identifying co-regulated genes (targets)
- Problem: how do we fit the model when $f(t)$ is not observed?

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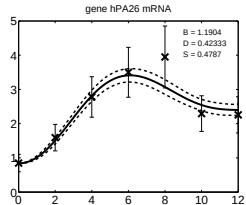
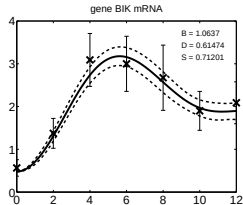
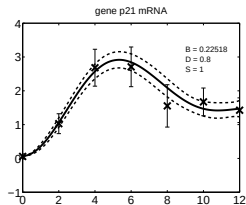
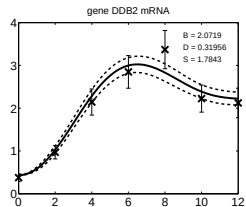
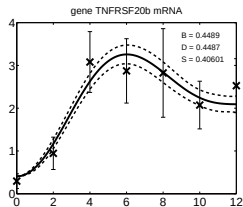
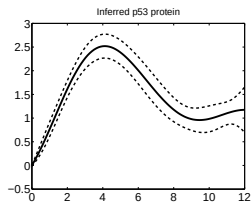
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Outline

Motivation and Review

Covariance Functions

Motion Capture Example

ODE Model of Transcriptional Regulation

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 $y(t)$

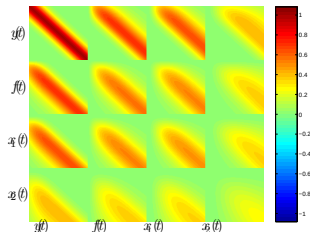
$$f(t) = \sigma \exp(-\delta t) \int_0^t y(u) \exp(\delta u) du$$

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

- 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`

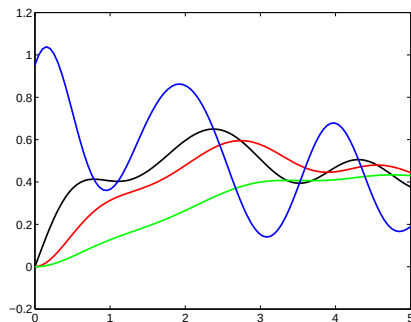


Figure: Joint samples from the ODE covariance, *blue*: $y(t)$ (mRNA of TF), *black*: $f(t)$ (TF concentration), *red*: $x_1(t)$ (high decay target) and *green*: $x_2(t)$ (low decay target)

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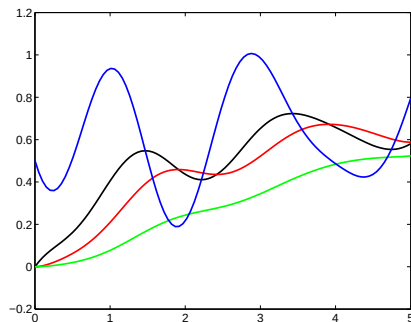


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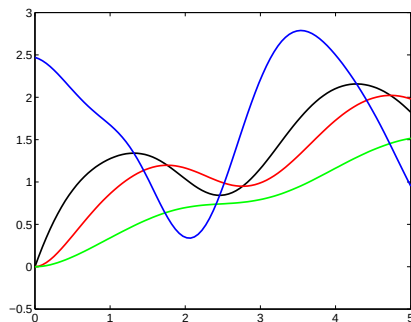


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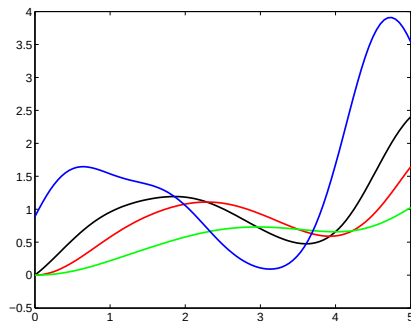


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Twist Results

- ▶ 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.

Results for Twi using the Cascade model

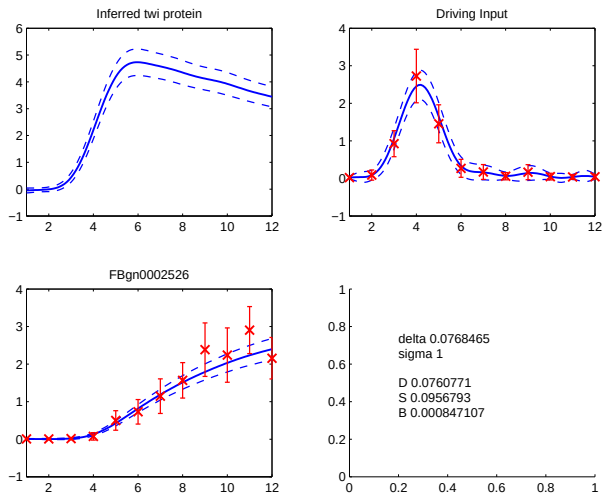


Figure: Model for flybase gene identity FBgn0002526.

Results for Twi using the Cascade model

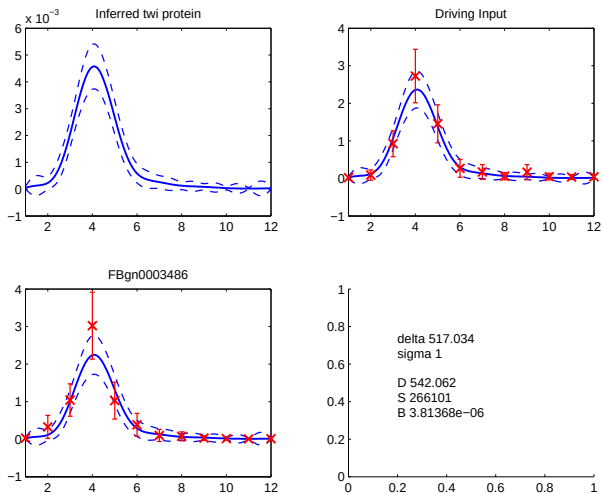


Figure: Model for flybase gene identity FBgn0003486.

Results for Twi using the Cascade model

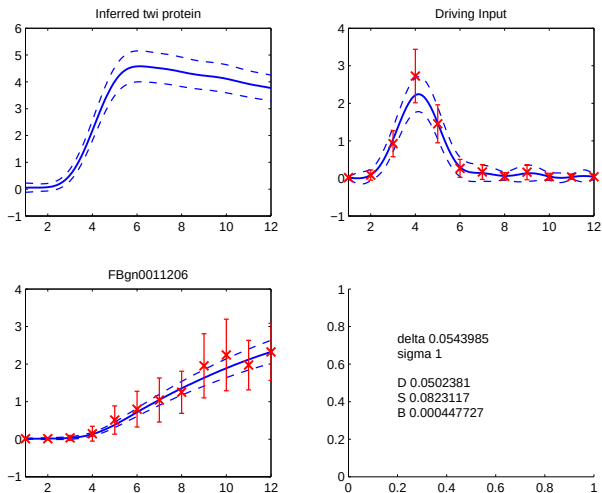


Figure: Model for flybase gene identity FBgn0011206.

Results for Twi using the Cascade model

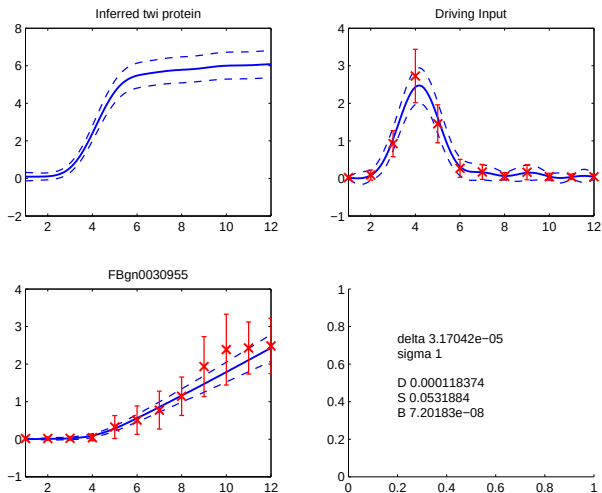


Figure: Model for flybase gene identity FBgn00309055.

Results for Twi using the Cascade model

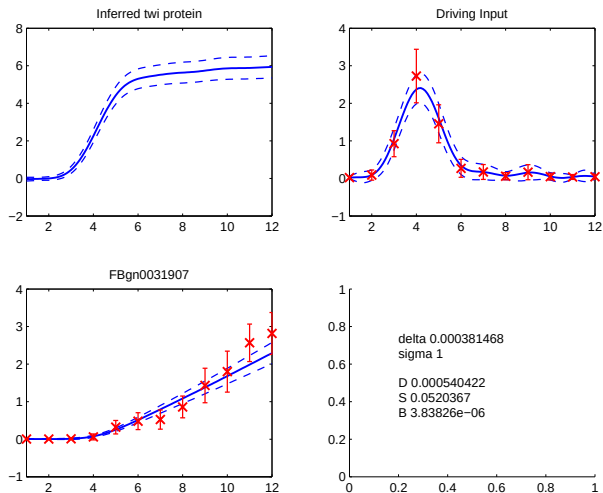


Figure: Model for flybase gene identity FBgn0031907.

Results for Twi using the Cascade model

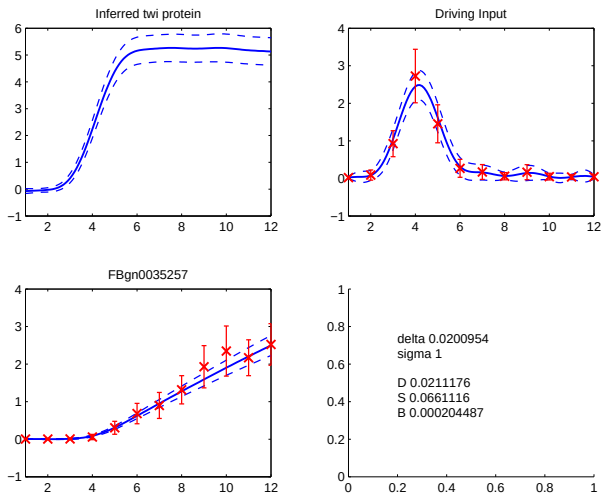


Figure: Model for flybase gene identity FBgn0035257.

Results for Twi using the Cascade model

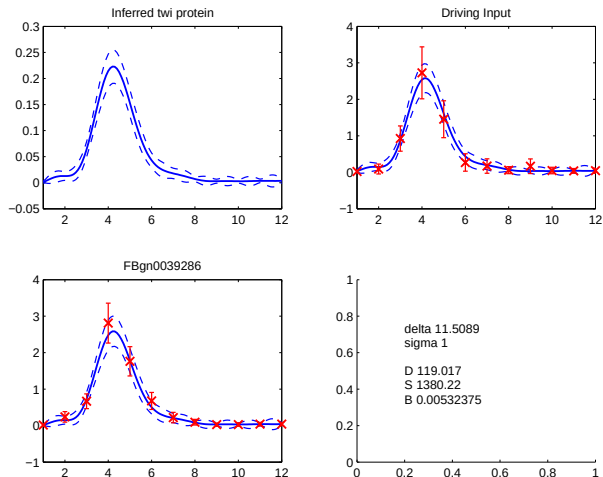


Figure: Model for flybase gene identity FBgn0039286.

Results of Ranking

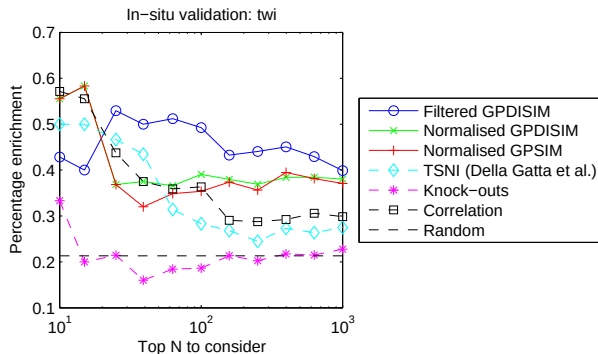


Figure: Percentage enrichment for top N targets for relevant terms in *Drosophila* in situs.

Results of Ranking

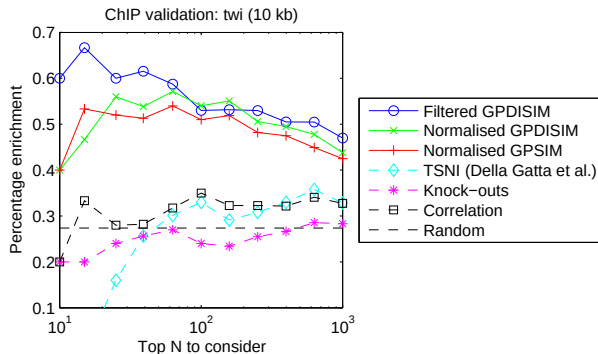


Figure: Percentage enrichment for top N targets for ChIP-chip confirmed targets.

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

Motivation and Review

Covariance Functions

Motion Capture Example

ODE Model of Transcriptional Regulation

Cascade Differential Equations

Discussion and Future Work

Discussion and Future Work

- ▶ Integration of probabilistic inference with mechanistic models.
- ▶ Ongoing/other work:
 - ▶ Non linear response and non linear differential equations.
 - ▶ Scaling up to larger systems
 - ▶ Robotics applications
 - ▶ Applications to other types of system, e.g. spatial systems.
 - ▶ Stochastic differential equations

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PDE Example

Efficient Approximations

Non-linear Response

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- ▶ Can extend the concept to latent functions in PDEs.
- ▶ Jura data: concentrations of heavy metal pollutants from the Swiss Jura.
- ▶ Consider a latent function that represents how the pollutants were originally laid down (initial condition).
- ▶ Assume pollutants diffuse at different rates resulting in the concentrations observed in the data set.

$$\frac{\partial x_q(\mathbf{x}, t)}{\partial t} = \sum_{j=1}^d \kappa_q \frac{\partial^2 x_q(\mathbf{x}, t)}{\partial x_j^2},$$

- ▶ Latent function $f_r(\mathbf{x})$ represents the concentration of pollutants at time zero (i.e. the system's initial condition).

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- The solution to the system (Polyanin, 2002) is then given by

$$x_q(\mathbf{x}, t) = \sum_{r=1}^R S_{rq} \int_{\mathbb{R}^d} f_r(\mathbf{x}') G_q(\mathbf{x}, \mathbf{x}', t) d\mathbf{x}'$$

where $G_q(\mathbf{x}, \mathbf{x}', t)$ is the Green's function given as

$$G_q(\mathbf{x}, \mathbf{x}', t) = \frac{1}{2^d \pi^{d/2} T_q^{d/2}} \exp \left[- \sum_{j=1}^d \frac{(x_j - x'_j)^2}{4 T_q} \right],$$

with $T_q = \kappa_q t$.

- For latent function given by a GP with the RBF covariance function this is tractable.

$$k_{x_p x_q}(\mathbf{x}, \mathbf{x}', t) = \sum_{r=1}^R \frac{S_{rp} S_{rq} |\mathbf{L}_r|^{1/2}}{|\mathbf{L}_{rp} + \mathbf{L}_{rq} + \mathbf{L}_r|^{1/2}} \\ \times \exp \left[-\frac{1}{2} (\mathbf{x} - \mathbf{x}')^\top (\mathbf{L}_{rp} + \mathbf{L}_{rq} + \mathbf{L}_r)^{-1} (\mathbf{x} - \mathbf{x}') \right],$$

where $\mathbf{L}_{rp}$, $\mathbf{L}_{rq}$ and $\mathbf{L}_r$ are diagonal isotropic matrices with entries $2\kappa_p t$, $2\kappa_q t$ and $1/\ell_r^2$ respectively. The covariance function between the output and latent functions is given by

$$k_{x_q f_r}(\mathbf{x}, \mathbf{x}', t) = \frac{S_{rq} |\mathbf{L}_r|^{1/2}}{|\mathbf{L}_{rq} + \mathbf{L}_r|^{1/2}} \\ \times \exp \left[-\frac{1}{2} (\mathbf{x} - \mathbf{x}')^\top (\mathbf{L}_{rq} + \mathbf{L}_r)^{-1} (\mathbf{x} - \mathbf{x}') \right].$$

Mauricio Alvarez

- ▶ Replicate experiments in (Goovaerts, 1997, pp. 248,249):
 - ▶ *Primary variable* (Cd, Cu, Pb, Co) predicted in conjunction with *secondary variables* (Ni and Zn for Cd; Pb, Ni, and Zn for Cu; Cu, Ni, and Zn for Pb; Ni and Zn for Co).¹
- ▶ Condition on the secondary variables to improve prediction for primary variables.
- ▶ Compare results for the diffusion kernel with independent GPs and “ordinary co-kriging” (Goovaerts, 1997, pp. 248,249).

¹Data available at <http://www.ai-geostats.org/>.

Table: Mean absolute error and standard deviation for ten repetitions of the experiment for the Jura dataset. IGP stands for independent GPs, GPK stands for GP diffusion kernel, OCK for ordinary co-kriging. For the Gaussian process with diffusion kernel, we learn the diffusion coefficients and the length-scale of the covariance of the latent function.

Metals	IGPs	GPK	OCK
Cd	0.5823 ± 0.0133	0.4505 ± 0.0126	0.5
Cu	15.9357 ± 0.0907	7.1677 ± 0.2266	7.8
Pb	22.9141 ± 0.6076	10.1097 ± 0.2842	10.7
Co	2.0735 ± 0.1070	1.7546 ± 0.0895	1.5

Outline

PDE Example

Efficient Approximations

Non-linear Response

Mauricio Alvarez

- Solutions to these differential equations is normally as a convolution.

$$x_i(t) = \int f(u) k_i(u - t) du + h_i(t)$$

$$x_i(t) = \int_0^t f(u) g_i(u) du + h_i(t)$$

- Convolution Processes (Higdon, 2002; Boyle and Frea, 2005).
- Convolutions lead to $N \times d$ size covariance matrices $O(N^3 d^3)$ complexity, $O(N^2 d^2)$ storage.
- Model is conditionally independent over $\{x_i(t)\}_{i=1}^d$ given $f(t)$.

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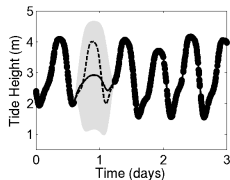
- ▶ Can assume conditional independence given $\{f(t_i)\}_{i=1}^k$.
(Álvarez and Lawrence, 2009)
 - ▶ Result is very similar to PITC approximation (Quiñonero Candela and Rasmussen, 2005).
 - ▶ Reduces to $O(N^3 dk^2)$ complexity, $O(N^2 dk)$ storage.
 - ▶ Can also do a FITC style approximation (Snelson and Ghahramani, 2006).
 - ▶ Reduces to $O(Ndk^2)$ complexity, $O(Ndk)$ storage.

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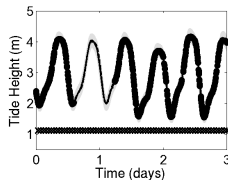
- ▶ Network of tide height sensors in the solent — tide heights are correlated.
- ▶ Data kindly provided by Alex Rogers (see Osborne et al., 2008).
- ▶ $d = 3$ and $N = 1000$ of the 4320 for the training set.
- ▶ Simulate sensor failure by knocking out one sensor for a given time.
- ▶ For the other two sensors we used all 1000 training observations.
- ▶ Take $k = 100$.

Tide Height Results

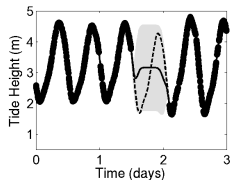
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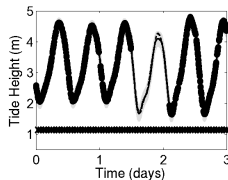
(a) Bramblemet Independent



(b) Bramblemet PITC



(c) Cambermet Independent

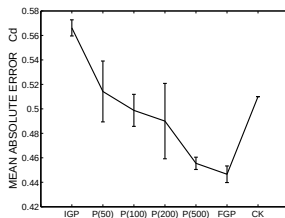


(d) Cambermet PITC

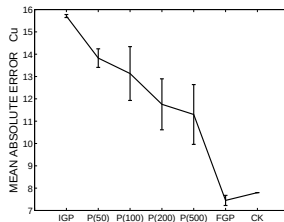
Mauricio Alvarez

- ▶ Jura dataset — concentrations of several heavy metals.
- ▶ Prediction 259 data, validation 100 data points.
- ▶ Predict *primary variables* (cadmium and copper) at prediction locations in conjunction with some *secondary variables* (nickel and zinc for cadmium; lead, nickel and zinc for copper) (Goovaerts, 1997, p. 248,249).

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(a) Cadmium



(b) Copper

Figure: Mean absolute error. IGP stands for independent GP, $P(M)$ stands for PITC with M inducing values, FGP stands for full GP and CK stands for ordinary co-kriging.

Outline

PDE Example

Efficient Approximations

Non-linear Response

Models of non-linear regulation

- Non-linear Activation: Michaelis-Menten Kinetics

$$\frac{dx_i(t)}{dt} = B_i + \frac{S_i f(t)}{\gamma_i + f(t)} - D_i x_i(t)$$

used by Rogers and Girolami (2006)

- Non-linear Repression

$$\frac{dx_i(t)}{dt} = B_i + \frac{S_i}{\gamma_i + f(t)} - D_i x_i(t)$$

used by Khanin et al., 2006, PNAS 103

Models of non-linear regulation

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- Non-linear Repression

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MAP Laplace Approximation

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,

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

Use Laplace's method (Laplace, 1774),

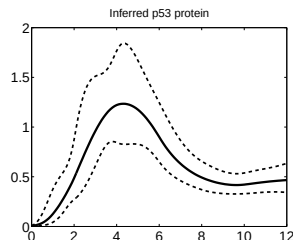
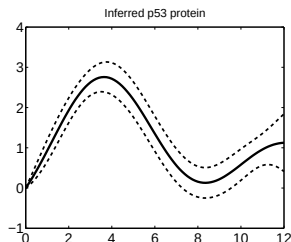
$$p(\mathbf{f} | \mathbf{x}) = N(\hat{\mathbf{f}}, \mathbf{A}^{-1}) \propto \exp\left(-\frac{1}{2} (\mathbf{f} - \hat{\mathbf{f}})^T \mathbf{A} (\mathbf{f} - \hat{\mathbf{f}})\right)$$

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

- 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.



(a)

Validation of Laplace Approximation

Michalis Titsias

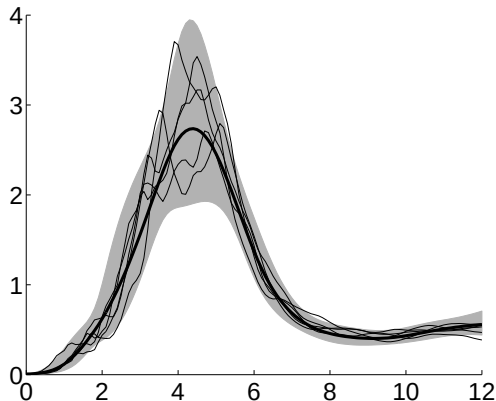


Figure: Laplace approximation error bars along with samples from the true posterior distribution.

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. (Titsias et al., 2009)

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 $\mathbf{f}_\rho$ using the conditional GP prior $p\left(\mathbf{f}_\rho^{(t+1)}|\mathbf{f}_c^{(t+1)}\right)$
- ▶ The whole proposal is

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

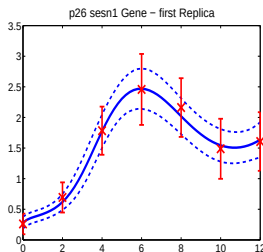
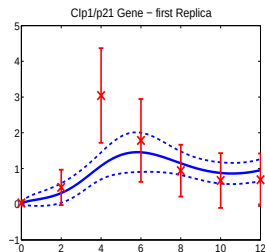
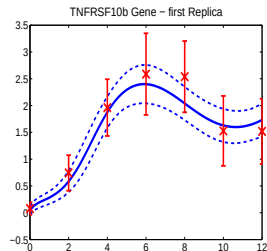
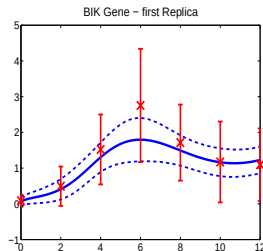
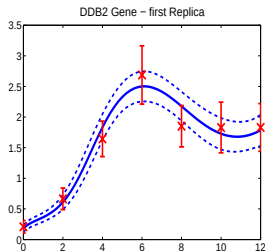
- ▶ Its like sampling from the prior $p(\mathbf{f})$ but imposing random walk behaviour through the control points.

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

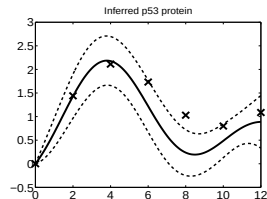
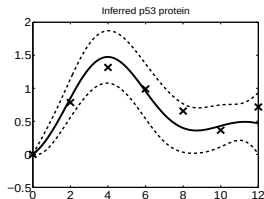
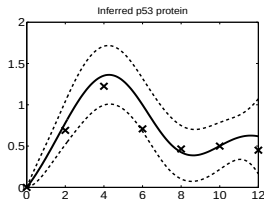
$$\frac{dx_j(t)}{dt} = B_j + S_j \frac{\exp(f(t))}{\exp(f(t)) + \gamma_j} - D_j x_j(t)$$

- ▶ 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%

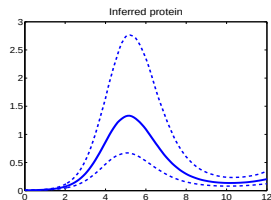
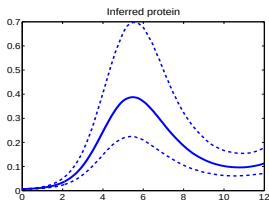
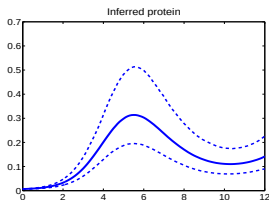
Data used by Barenco et al. (2006): Predicted gene expressions for the 1st replica



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

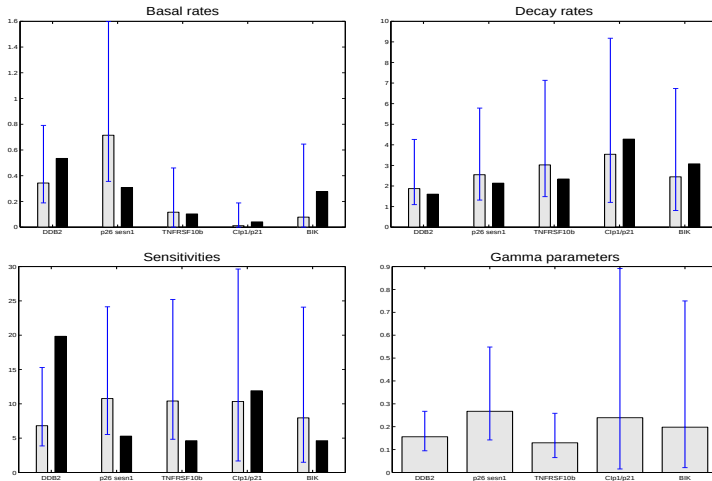


Linear model (Barenco et al. predictions are shown as crosses)



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