



Network Reconstruction & Network Analysis

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Content

- Network reconstruction
 - Boolean models
 - Correlation-Based Approaches: REVEAL / ARACNE
 - Example
- Quantitative network analysis

Networks

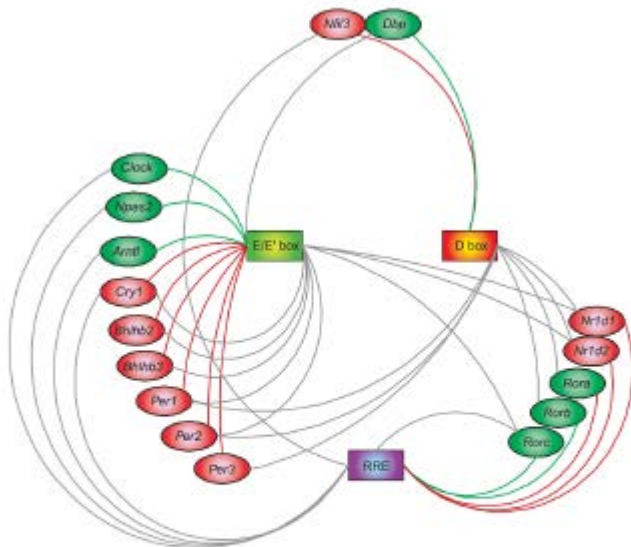
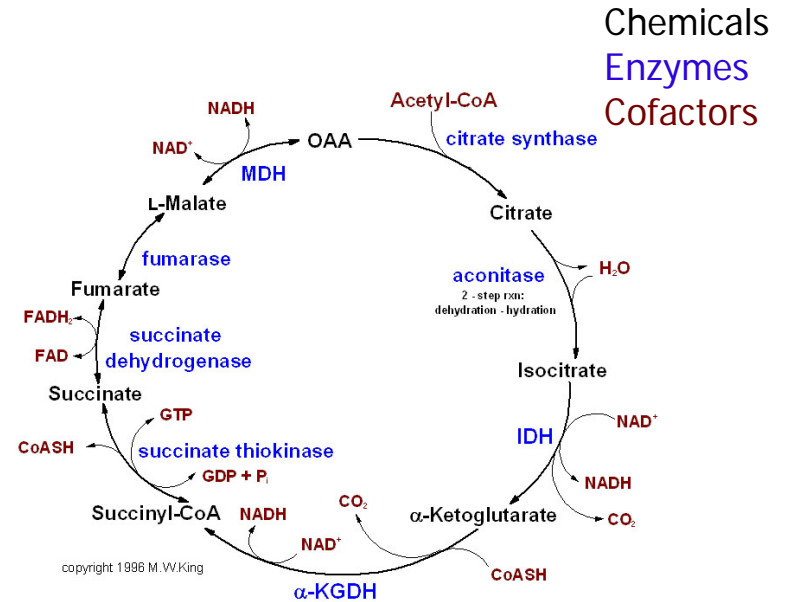


Abbildung 2: Zentrale Gene der zirkadianen Uhr und deren wechselseitiger Einfluss. [UHC⁺05] (Kästen: Cis-Elemente/Grüne Ovale: Positiv regulierende Gene/Rote Ovale: Negativ regulierende Gene/Regulationsrichtung 1: Von Gen über farbige Kante zu Cis-Element/Regulationsrichtung 2: Von Cis-Element über graue Kante zu Gen)



Chemicals
Enzymes
Cofactors

How do we know?
What does the network tell us?

Approaches to Network Reconstruction

- By many, many small-scale experiments
- By mathematical modeling from [high-throughput data sets](#)
- By evolutionary inference from model organisms
- By curation from the literature (see first bullet)

Reconstruction from Indirect High-Throughput Data

- Network reconstruction, re-engineering, inference, ...
- Idea: Derive network from indirect observations
 - **Network**: Links and their effect (strength, activation, ...)
 - We usually assume the players (genes, metabolites, ...) to be given
 - **Observation**: High-throughput measurements
 - Here: Transcriptome, microarrays, RNA-Seq
 - **Indirect**: We try to infer **mechanistic causality by correlation**
- Dynamic networks
 - Nodes get states (active / passive)
 - Current states **determine future states** of nodes
 - Leads to dynamic behavior
- Warning: All current methods are **highly reductionist**

Boolean Network Models

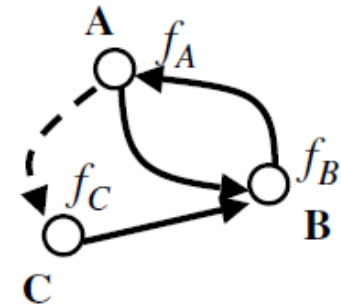
- Definition

A *Boolean Network* is a digraph $G=(V,E)$ where

- Every node has an associated Boolean state (on/off)
- Every node is labeled with a *Boolean function over the states of all incoming nodes*

- Usage

- Vertices = genes
- Edge (v,w) models an effect of v on w
- The state of a node v is determined by its Boolean function over all “incoming” states
- Simplistic: No cofactors, no cellular context, no binding affinity, no time, no kinetics, ...



$$f_A(B) = B$$

$$f_B(A, C) = A \text{ and } C$$

$$f_C(A) = \text{not } A$$

Boolean Network

Network Dynamics

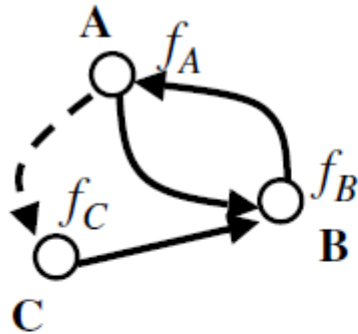
- Definition

A *Dynamic Boolean Network (DBN)* is a Boolean network where every node v is assigned a *sequence of states* V_0, V_1, V_2, \dots such that the state of v_t is defined over the Boolean function of v applied to the states w_{t-1} of all incoming nodes w

- Remarks

- Models the state of every gene (on / off) over time
- States at time point t (only) **depend on states at time point $t-1$**
 - No buffering, synchronized time, ...
- **Deterministic**: Given all states at any time point t and the Boolean functions, any state at any later time point can be uniquely determined

Example

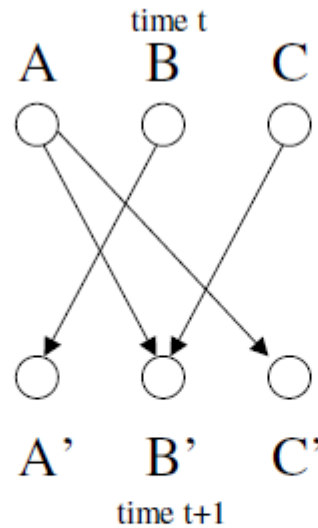


$$f_A(B) = B$$

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Boolean Network



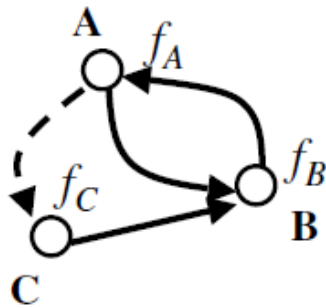
Wiring Diagram

State	INPUT			OUTPUT		
	A	B	C	A'	B'	C'
1	0	0	0	0	0	1
2	0	0	1	0	0	1
3	0	1	0	1	0	1
4	0	1	1	1	0	1
5	1	0	0	0	0	0
6	1	0	1	0	1	0
7	1	1	0	1	0	0
8	1	1	1	1	1	0

Transition table

Source: Filkov, „Modeling Gene Regulation“, 2003

Example



$$f_A(B) = B$$

$$f_B(A, C) = A \text{ and } C$$

$$f_C(A) = \text{not } A$$

Boolean Network

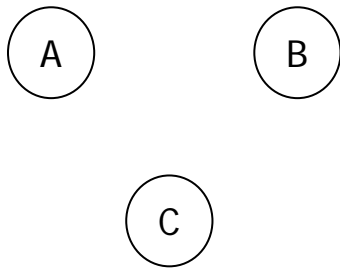
genes time	A	B	C
0	1	1	0
1	1	0	0
2	0	0	0
3	0	0	1
4	0	0	1
5

Network Analysis

- Many things can be analyzed using DBN
- For instance, an **attractor** is a (set of) states towards which the network state converges
 - Point attractor: State which cannot be left any more
 - Cyclic attractor: A series of states which will repeat forever
 - Probability of attractors depend largely on size of network and complexity of Boolean functions
- Skipped – we want to **reconstruct networks**

Network Reconstruction

- Assume we know all genes, but **not their relationships**
- Assume we observe the states of n genes over m time points (a matrix S; the **observations**)
- Can we **re-engineer the Boolean function** of every gene given a sequence of states?

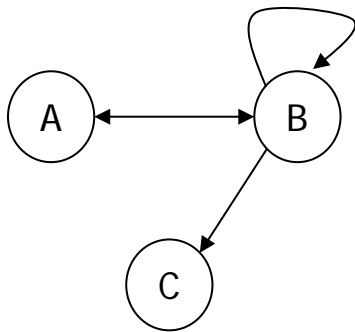


S

genes time	A	B	C
0	1	1	0
1	0	0	1
2	1	0	1
3	1	1	0
4	0	0	1
5

Network Reconstruction

- Assume we know all genes, but not their relationships
- Assume we observe the states of n genes over m time points (a matrix S ; the observations)
- Can we re-engineer the Boolean function of every gene given a sequence of states?



$f(A) = \text{not } B$

$f(B) = A \text{ and not } B$

$f(C) = B$

genes time	A	B	C
0	1	1	0
1	0	0	1
2	1	0	0
3	1	1	0
4	0	0	1
5

Formal Problem

- Definition

Let S_t , $0 \leq t \leq m$, be the vector of all observed states of all genes V at time point t . A DBN $G=(V,E)$ with functions f_1, \dots, f_n , $n=|V|$, is called

- *consistent with S_t iff $S_t = [f_1(S_{t-1}), f_2(S_{t-1}), \dots, f_n(S_{t-1})]$*
- *consistent with S iff it is consistent for all S_t , $1 \leq t \leq m$*

- The Boolean network reconstruction problem

*Given an observation S over a set V , find a **DBN $G=(V,E)$** that is consistent with S .*

- Remark

- Reconstruction means finding the functions f_1, \dots, f_n
- This also determines **network topology** (nodes appearing in a f_i)

Solutions

- Clearly, there are many observations S for **which no consistent G** exists
 - Recall that DBN are deterministic
 - Imagine S_t, S_{t+1} and S_u, S_{u+1} with $S_t = S_u$ but $S_{t+1} \neq S_{u+1}$
- Also, there are many observation S for which **more than one consistent G** exists
- Every time point narrows the options for G – the longer S , the less (or no) consistent G 's exist

Optimal Networks

- Definition
 - For a DBN G , let $\text{size}(G)$ be the total number of variables (edges) appearing in the Boolean functions of G
 - A DBN G is minimal for observation S , if G is consistent with S and there is no G' which is also consistent with S and $\text{size}(G') < \text{size}(G)$
- Remark
 - Parsimony assumption: Small models are better
 - Thus, the smallest network is the best – functions are as simple as possible, nothing is inferred that is not enforced by the data
 - Not necessarily unique

Naive Algorithm

```
N = V;
for k=1..n                                # length of functions
  for every n in N                          # all unexplained nodes
    test all functions f of size k for n on S;
    if f is consistent for n on S
      N := N \ n;                            # n is explained
      Add f to network;
    end if;
  end for;
end for;
```

- Exhaustive algorithm for finding minimal networks
- **Very complex** (AND, OR, NOT, no paranthesis)
 - k=1: $2n$ functions
 - k=2: $2 \cdot 2n \cdot 2n = O(n^2)$ functions
 - ...
 - General: $O(2^{2k-1} \cdot n^k)$ functions

Pros and Cons

- Application (transcriptome data)
 - Perform **time-series gene expression** experiments
 - **Brutally discretize** each measurement: Genes are on or off
 - Reconstruct DBN
- Pros: Simple
- Cons
 - Binary values are not capturing reality
 - Synchronized, clocked time is nonsense
 - No quantification (It needs $2 \cdot A$ and one B to regulate C)
 - Only small networks are computable
 - ...

Content

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Towards Reality

- There are **less complex & more robust** algorithms
- REVEAL replaces Boolean functions by mutual information; correlations rather than deterministic switching
 - Liang, S., S. Fuhrman and R. Somogyi (1998). Reveal, a general reverse engineering algorithm for inference of genetic network architectures. Pacific Symposium on Biocomputing., Hawaii, US.
- ARACNE is even simpler: Only removal of some (presumably indirect) correlations
 - Margolin, A. A., I. Nemenman, K. Basso, C. Wiggins, G. Stolovitzky, R. D. Favera and A. Califano (2006). "ARACNE: an algorithm for the reconstruction of gene regulatory networks in a mammalian cellular context." BMC Bioinformatics 7((Suppl 1), S7).

Foundations

- Definition

*Let X, Y be two discrete random variables. The **mutual information** $MI(X, Y)$ is defined as*

$$MI(X, Y) = \sum_{x \in X} \sum_{y \in Y} p(x, y) * \log \left(\frac{p(x, y)}{p(x) * p(y)} \right)$$

- Remark

- Measure the variable's mutual dependency
- Dependency: Deviation of $p(X, Y)$ from $p(X) * p(Y)$
- How much does the state of X determines the state of Y ?
- Many similar measures (information gain, conditional entropy, cross entropy, ...)

Example

$$MI(X, Y) = \sum_{x \in X} \sum_{y \in Y} p(x, y) * \log \left(\frac{p(x, y)}{p(x) * p(y)} \right)$$

p(x,y)	y=0 p(y=0)=0.6	y=1 p(y=1)=0.4
x=0; p(x=0)=0.2	0,12	0,08
x=1; p(x=1)=0.8	0,48	0,32

MI(X,Y)=0

p(x,y)	y=0 p(y=0)=0.6	y=1 p(y=1)=0.4
x=0; p(x=0)=0.2	0,18	0,03
x=1; p(x=1)=0.8	0,05	0,74

MI(X,Y)=0,53

Two more Facts

- With a little math, we find

$$MI(X,Y) = H(X) - H(X|Y) = H(Y) - H(Y|X)$$

- $H(X)$: Entropy of X
 - $H(X|Y)$: **Conditional entropy** of X given Y
-
- It follows that the **maximal value of $MI(X,Y) = H(X)$ ($H(Y)$)**
 - $H(X|Y) = 0$, which means that **X (Y) completely determines Y (X)**
-
- MI can be extended to sets of three, four, ... variables
 - Like Boolean functions over three, four, ... variables
 - Multivariate mutual information

REVEAL

```
N = V;
for k=1..n                # number of nodes/variables
  for every X in N        # all unexplained nodes
    find subset T=(Y1,...Yk) with MI(X,Y1,...Yk) = H(X);
    if T exists
      N := N \ X;         # n is explained
    end for;
  end for;
end for;
```

- Again, we have observations of n genes at m time points
 - Or m different conditions, treatments, ...
- Again, we discretize expression values to 0 or 1
 - More bins are possible
- $MI(X,Y)$ means looking at pairs $(x_1,y_0), (x_2,y_1), \dots$

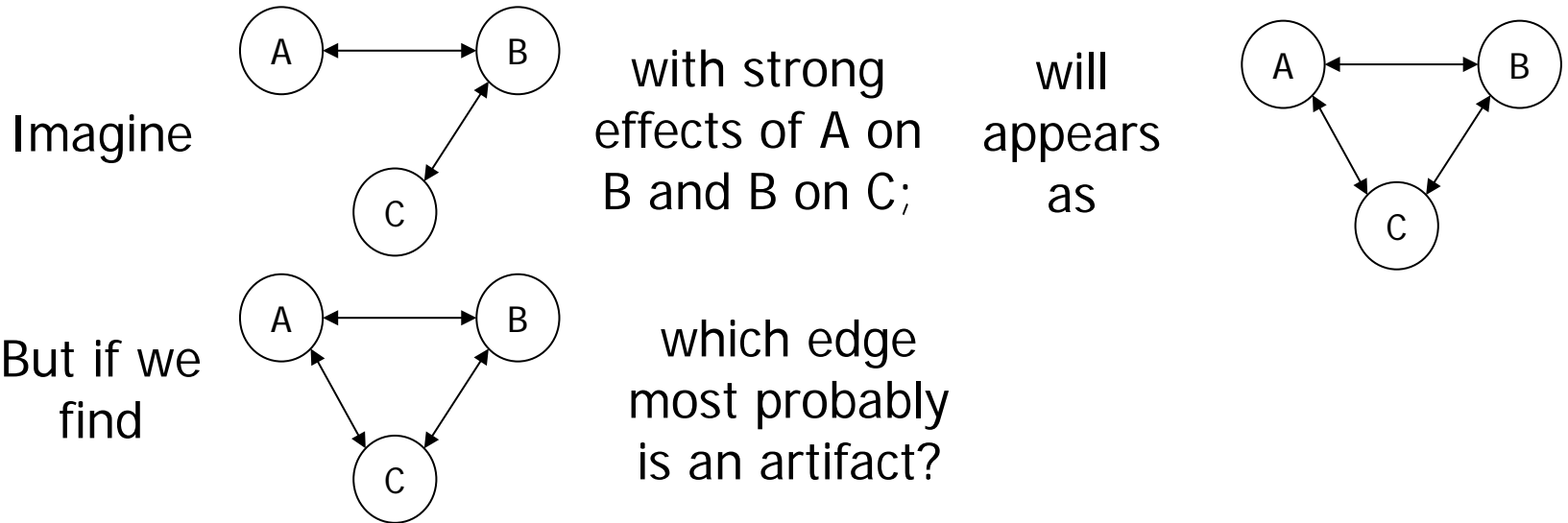
REVEAL in Practice

- In the formulation given, REVEAL would be as strict as Boolean functions
 - Dependencies must be perfect
- In the presence of noise, one must be satisfied with **almost maximal MI**
 - I.e., $|MI(X,Y)-H(X)| < \epsilon$
- Can be extended to **more than two possible states**
 - Less strict discretization, more realistic model
- Most other restrictions of DBN remain

ARACNE

- **Fast** variation of REVEAL which (a) considers each pair in isolation and (b) gives up model minimality
- **Idea**
 - Compute mutual information between all pairs of genes
 - This gives a **complete network**
 - Remove **edges where $|MI(X,Y)-H(X)| > \epsilon$**
 - ϵ can be estimated from the distribution of MI – created at random?
 - Remove certain **indirect effects** (“data processing inequalities”)
- Under certain assumptions, ARACNE provably converges to the true network
 - Given unlimited input, no loops
 - “True”: Under all networks obeying our simplifying assumptions

Data Processing Inequalities



- Assumption: If $MI(X,Z) \leq \min(MI(X,Y),MI(Y,Z))$, then the correlation between **X-Z is an indirect effect** and removed
- Procedural: In **every triangle**, remove the smallest edge
 - But in which order should triangles be visited?

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Reconstructing the Mammalian Clock

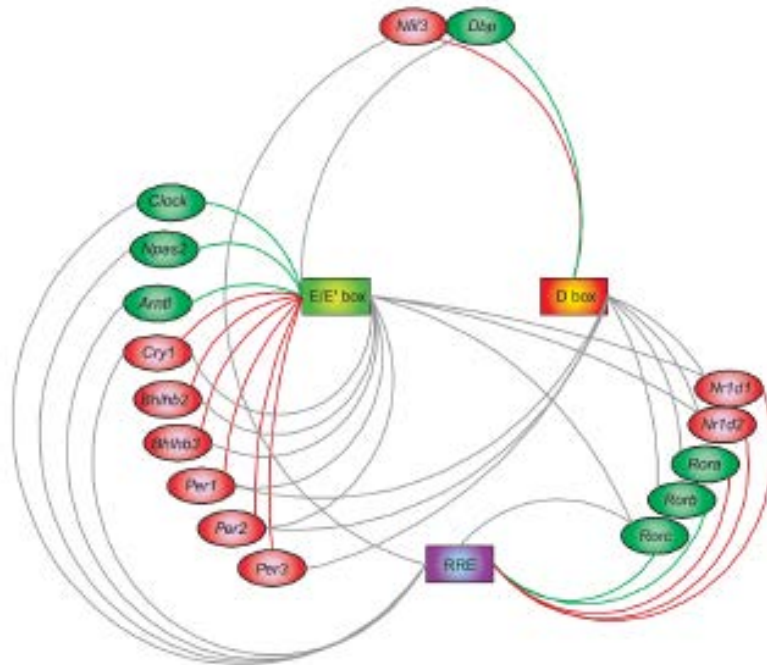


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- DA Sven Lund, 2015
- Data
 - ~630 rather unspecific arrays from GEO
 - Compared to two time-resolved clock-specific experiments
- Reconstruction quality of three algorithms
 - Aracne, Bayes Networks, Time-Delay Aracne

Results

Kennzahl	Verfahren	TP	TN	FP	FN	Recall	Precision
\bar{x}	Pearson	53.75	20.00	41.00	21.25	0.72	0.57
s	Pearson	4.979	8.718	8.718	4.979	0.068	0.070
\bar{x}	Bayes	36.00	33.50	27.50	39.00	0.48	0.57
s	Bayes	12.739	10.282	10.282	12.739	0.170	0.020
\bar{x}	ARACNE	18.88	48.00	13.00	56.13	0.25	0.59
s	ARACNE	5.515	-----	-----	-----	-----	-----

Kennzahl	Datenquelle	TP	TN	FP	FN	Recall	Precision
\bar{x}	GEO	45.00	26.00	35.00	30.00	0.60	0.57
s	GEO	17.550	16.480	16.480	17.550	0.235	0.034
\bar{x}	Korenčić	35.67	36.22	24.78	39.33	0.48	0.60
s	Korenčić	16.462	12.940	12.940	16.462	0.219	0.037
\bar{x}	Hogenesch	30.89	36.67	24.33	44.11	0.41	0.55
s	Hogenesch	15.648	12.708	12.708	15.648	0.208	0.094

- Filtering of ARACNE **reduces recall a lot**, while precision increases only marginally
- Data set **size outweighs specificity** – reconstruction about as good using many untargeted arrays or using fewer targeted arrays

Content

- Network reconstruction
- Quantitative network analysis
 - A model of transcriptional regulation
 - Metabolic network models
 - Kinetic modeling

Networks as Equations

- REVEAL / ARACNE infer relationships based on correlation
- Alternative: Describe states as **sets of (linear) equations**
 - **No discretization**
 - Extensibility: Incorporate different types of experiments (binding, epigenetic status, mRNA translation, ...)
 - Still many limitations: Synchronized time, no kinetics, ...
- We look at one simple approach in between reconstruction and analysis
 - We assume the network topology to be given
 - We infer the (probable) **strengths of different effects**
 - Schacht et al. (2014). "Estimating the activity of transcription factors by the effect on their target genes." Bioinformatics

Approach

- Assume a network $G=(V,E)$, where V consists of a set of **transcription factors T and a set of genes G**
 - Transcription factors regulate genes, but not vice versa
 - We also ignore that a TF may regulate TFs (other or even itself)
 - Each gene g is regulated by all its incoming TFs (and no others)
- Measurements: m observations for n nodes (genes / TFs)
- We model the expression values of all genes as **linear combinations** of the values of its regulating TFs

$$g_{i,j} = \beta_0 + \sum_{t=1}^{|T|} \beta_t * \delta_{t,j} * e_{t,i}$$

Model

Sought

$$g_{i,j} = \beta_0 + \sum_{t=1}^{|T|} \beta_t * \delta_{t,j} * e_{t,i}$$

Given

- $g_{i,j}$: Expression of gene i in observation S_j
- β_0 : Fixed additive offset (sample differences)
- β_t : Global activity parameter for transcription factor t
- $\delta_{t,j}$: Observation specific (S_j) effect on t
- $e_{t,i}$: Effect strength of TF t on gene i
 - Set to 0 if there is no edge between g and t

Optimization

- Typically, these (large) systems cannot be solved exactly
- Instead, minimize the error

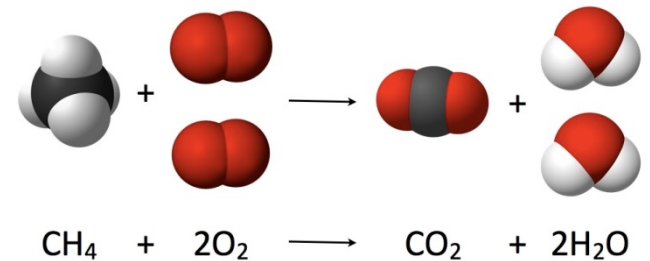
$$\left| g_{i,j} - \left(\beta_0 + \sum_{t=1}^{|T|} \beta_t * \delta_{t,j} * e_{t,i} \right) \right| = \min$$

- ... under a set of constraints
- Several solvers available

Many Other Models

- Stoichiometric networks

- Model the turnover of molecules
 - Especially metabolism
- Needs to consider enzymatic effects
- What will a **network produce** given a certain input?
- Is a network in **flux balance**?



- Kinetic networks

- Takes into account reaction rates: How many in what time
 - No linear relationship
- Leads to **systems of differential equations**
- Can predict system **behavior in time** under realistic assumptions