

Analyse statistique de réseaux biologiques Sophie S. Schbath

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Analyse statistique de réseaux biologiques

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Séminaire Proba-Stat, Orsay, 23 juin 2011

Part 1 Introduction

The network revolution

Nature of the data :

- n individuals (n large),
- but also n² couples.

Many scientific fields :

sociology, physics, "internet", biology.

Biological networks :

protein-protein interaction networks, regulatory networks, metabolic networks.

Gene regulatory networks

- nodes = genes
- edges : regulations (directed)

Protein-protein interaction networks

- nodes = proteins
- edges : physical interaction

Metabolic networks

- nodes = chemical compounds
- edges : chemical reactions or enzyme (directed, hyper-edges)

- nodes = enzymes
- edges : consecutiveness in the metabolic network



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Main characteristics :

- several thousands of nodes (n)
- sparsity (nb of edges = O(n))
- heterogeneous connexions
- nodes may be coloured (biological function, class of reaction, cellular localization etc.)

The network revolution (fol.)

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 - but also n^2 couples.
- Many scientific fields : sociology, physics, "internet", biology.

Biological networks :

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Statistical aspects :

- network inference,

- statistical properties of given networks (degrees, diameter, clustering coefficient, modules, motifs etc.),

- random graph models.

Looking for local structures

- Breaking-down complex networks into functional modules or basic building blocks : [Shen-Orr et al. (02)]
 → network motifs : topological motifs and/or coloured motifs.
- Focus on exceptional motifs = motifs appearing more frequently than expected.

[*Milo et al. (02), Shen-Orr et al. (02), Zhang et al. (05), Lacroix et al. (06), Lee et al. (07), Taylor et al. (07)*]

Network motifs (1/2)

- Topological motif : connected pattern of interconnection
 - \rightarrow an occurrence in the network is an isomorphic subgraph



Ex : particular regulatory units like feed-forward loop or bi-fan motifs.

Interpretation of over-represented topological motifs : they are thought to reflect functional units which combine to regulate the cellular behavior as a whole.

Network motifs (2/2)

Coloured motif : multiset of node colours, e.g. {•, •, •}

 \rightarrow an occurrence in the network is a connected subgraph with the appropriate node colours



Interpretation of over-represented coloured motifs : they are thought to reflect groups of cooperative enzymes (reaction networks). Introduction

Coloured motifs : example



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Statistics for biological networks

Orsay, 23 juin 2011 13 / 56

How to assess the exceptionality of a motif?

Step 1 **To count** the observed number of occurrences $N_{obs}(\mathbf{m})$ of a given motif \mathbf{m} (out of my scope)

Its significance is assessed with the *p*-value $\mathbb{P}\{N(\mathbf{m}) \ge N_{obs}(\mathbf{m})\}\$ (the probability to get as much occurrences at random)

Step 2 To choose an appropriate random graph modelStep 3 To get the distribution of the count *N*(m) under this model

State of the art (1/2)

Analytical approaches :

- The most popular random graph model is the Erdös-Rényi model (nodes are connected independently with proba p)
- Some theoretical works exist on Poisson and Gaussian approximations of topological motif count distribution (see [Janson et al. (00)] for an overview)

BUT

- only for particular motifs ("balanced" property),
- the Erdös-Rényi model is not a good model for biological networks (e.g. it does not fit the degrees).

State of the art (2/2)

Simulated approaches :

- Random networks are generated by edge swapping, (degrees are preserved)
- Empirical distributions for motif counts are obtained leading either to *p*-values or to *z*-scores

BUT

- huge number of simulations required to estimate tiny p-values,
- z-scores are compared to N(0, 1) which is not always appropriate,
- edge swapping does not define a probabilistic random graph model.

SSB contributions (1/2)

- To propose probabilistic random graph models
 - adapted for biological networks,
 - allowing probabilistic calculations,
 - with efficient estimation procedures.

[*Daudin, Picard, Robin (08)*]. A mixture model for random graphs. *Statis. Comput.* [*Birmelé (07)*]. A scale-free graph model based on bipartite graphs. *Disc. Appl. Math.* [*Mariadassou, Robin, Vacher (10)*]. Uncovering structure in valued graphs : a variational approach. *Ann. Appl. Statist.*

[Latouche, Birmele, Ambroise (10)] Overlapping Stochastic Block Models with Application to the French Political Blogosphere. Annals of Applied Statistics [Daudin, Pierre, Vacher (10).] Model for Heterogeneous Random Networks Using Continuous Latent Variables and an Application to a Tree–Fungus Network. Biometrics [Latouche, Birmele, Ambroise (11)] Variational Bayesian Inference and Complexity Control for Stochastic Block Models. Statistical Modelling [Gazal, Daudin, Robin (11)]. Accuracy of variational estimates for random graph mixture models. J. Comput. Comput. Simul.

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SSB contributions (2/2)

• To provide general analytical results on motif count distribution :

- mean and variance of the count in a wide class of random graph models,
- relevant distribution to approximate the count distribution.

[*Matias, Schbath, Birmelé, Daudin and Robin (06)*] Network motifs : mean and variance for the count, *REVSTAT.* 4 31–51. [*Picard, Daudin, Schbath and Robin (08)*] Assessing the exceptionality of network motifs, *J. Comput. Biol.* [*Schbath, Lacroix and Sagot (09)*] Assessing the exceptionality of coloured motifs in networks, *EURASIP*

Part 2

Mixture model for random graphs (Stochastic Block model)

Random graphs

- A random graph G is defined by :
 - a set \mathcal{V} of fixed vertices with $|\mathcal{V}| = n$,
 - a set of random edges $\mathbf{X} = \{X_{ij}, (i, j) \in \mathcal{V}^2\}$ such that

$$X_{ij} = \begin{cases} 1 & \text{if } i \text{ and } j \text{ are connected,} \\ 0 & \text{otherwise} \end{cases}$$

- and a distribution on X_{ij} .
- Examples :
 - the Erdös-Rényi model,
 - the Stochastic Block Model (=mixture of ER models),
 - the Expected Degree Distribution model.

Erdös-Rényi model

- Edges X_{ij}'s are independent ...
- ... and identically distributed according to $\mathcal{B}(p)$

$$\mathbb{P}(X_{ij}=1)=p$$

Degrees are Poisson distributed

$$\mathcal{K}_i := \sum_{j \neq i} X_{ij} \sim \mathcal{B}(n-1,p) pprox \mathcal{P}((n-1)p)$$

 Bad fit of Erdös-Rényi model on biological networks due to heterogeneous connection probabilities along the network.

Stochastic Block Model (or "Mixnet")

- Vertices are spread into Q groups.
- Conditionally to the group of vertices, edges are independent and

$$X_{ij} \mid \{i \in q, j \in \ell\} \sim \mathcal{B}(\pi_{q,\ell})$$

 $\pi_{q,\ell}$ is the connection probability between groups q and ℓ . • Degrees are distributed according to a Poisson mixture

$$\mathcal{K}_i \sim \sum_q lpha_q \mathcal{B}(n-1, \overline{\pi}_q) ext{ with } \overline{\pi}_q = \sum_\ell lpha_\ell \pi_{q,\ell}$$

Introduced by [Nowicki and Snijers (2001)]

Mixnet fit

- *E. coli* reaction network : 605 vertices, 1782 edges. (data curated by V. Lacroix and M.-F. Sagot).
- Degrees : Poisson mixture versus empirical distribution



Mixnet flexibility



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Mixnet : estimation procedure

[Daudin, Picard and Robin (Stat. Comput. 08)]

Classical maximum likelihood procedures fail

- log-likelihood L(X) not calculable because of hidden groups (Z, Z_i is the group of node i).
- EM algorithm, classical to fit mixture models, cannot be used because P(Z | X) is not computable (all vertices are potentially connected, no local dependence).

Variational approach (iterative procedure)

- maximization of L(X) KL(P(Z | X), Q_R(Z)) where Q_R is the best approximation of P(Z | X) within a class of 'nice' distributions.
 ⇒ estimator of P(Z_i = q|X).
- analytical expressions for $\widehat{\alpha}_{q}$ and $\widehat{\pi}_{q,\ell}$

Choice of Q: heuristic penalized likelihood criterion inspired from BIC (ICL)

Part 3

Assessing the significance of topological motif frequencies

Topological motifs

Let **m** be a motif of size k (connected graph with k vertices, $k \ll n$).

- m is defined by its adjacency matrix (also denoted by m) :
 m_{uv} = 1 iff nodes u ↔ v (m_{uv} = 0 otherwise).
- Let R(m) be the set of non redundant permutations of m (so-called "versions").
- Ex : 3 versions of the V motif at a **fixed** position (*i*, *j*, *k*).



Occurrences of a motif

- Let α = (i₁,..., i_k) ∈ I_k be a possible position of **m** in *G*.
 G_α denotes the subgraph (V_{i1},..., V_{ik}).
- Non strict occurrences :

m occurs at position $\alpha \Leftrightarrow \mathbf{m} \subseteq \mathbf{G}_{\alpha}$

• Random indicator of occurrence : $Y_{\alpha}(\mathbf{m})$

$$Y_{\alpha}(\mathbf{m}) = \mathbf{1}\{\mathbf{m} \text{ occurs at position } \alpha\} = \prod_{1 \le u, v \le k} X_{i_u i_v}^{m_{uv}}.$$

• The total count *N*(**m**) of motif **m** is then :

$$N(\mathbf{m}) = \sum_{lpha \in I_k} \sum_{\mathbf{m}' \in \mathcal{R}(\mathbf{m})} Y_{lpha}(\mathbf{m}')$$

• Warning : $N(\mathbf{m})
eq$ number of induced subgraphs (" $\mathbf{m} = \mathbf{G}_{lpha}$ ").

Expected count and variance

Under assumptions (H1) and (H2) on the random graph model :

(H1) Stationary assumption : D(X_{i₁,j₁},...,X_{iℓ,jℓ}) = D(X_{i¹₁,j¹},...,X_{iℓjℓ})
(H2) Independence of disjoint occurrences
we have [*Picard, Daudin, Koskas, Schbath, Robin (08)*]

$$\mathbb{E}N(\mathbf{m}) = \binom{n}{k} |\mathcal{R}(\mathbf{m})| \mu(\mathbf{m}).$$

where $\mu(\mathbf{m}) := \mathbb{E} Y_{\alpha}(\mathbf{m}) = \mathbb{P}(\mathbf{m} \text{ occurs at } \alpha)$ and

$$\operatorname{Var} \mathcal{N}(\mathbf{m}) = \sum_{s=0}^{k} \mathcal{C}(n, k, s) \sum_{\substack{\mathbf{m}' \Omega \mathbf{m}'' \\ s}} \mu(\mathbf{m}' \Omega \mathbf{m}'') - [\mathbb{E} \mathcal{N}(\mathbf{m})]^{2}.$$

where $\mathbf{m}' \Omega \mathbf{m}''$ is a **super-motif** composed of the union of two overlapping occurrences of \mathbf{m}' and \mathbf{m}'' sharing *s* common vertices.

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Statistics for biological networks

Variance

By definition $\mathbb{V}arN(\mathbf{m}) = \mathbb{E}N^2(\mathbf{m}) - [\mathbb{E}N(\mathbf{m})]^2$. We then calculate

$$\mathbb{E}N^{2}(\mathbf{m}) = \mathbb{E}\left(\sum_{\alpha,\beta\in I_{k}}\sum_{\mathbf{m}',\mathbf{m}''\in\mathcal{R}(\mathbf{m})}Y_{\alpha}(\mathbf{m}')Y_{\beta}(\mathbf{m}'')\right),$$

$$= \mathbb{E}\left(\sum_{s=0}^{k}\sum_{|\alpha\cap\beta|=s}\sum_{\mathbf{m}',\mathbf{m}''\in\mathcal{R}(\mathbf{m})}Y_{\alpha\cup\beta}(\mathbf{m}'_{s}\mathbf{m}'')\right)$$

$$= \sum_{s=0}^{k}C(n,k,s)\sum_{\mathbf{m}'\Omega\mathbf{m}''}\mu(\mathbf{m}'\Omega\mathbf{m}''),$$

where $\mathbf{m}' \Omega \mathbf{m}''$ is a **super-motif** composed of the union of two overlapping occurrences of \mathbf{m}' and \mathbf{m}'' sharing *s* common vertices.

Candidate random graph models

• Erdös-Rényi model (ER) : Edges X_{ij} 's are i.i.d. $\sim \mathcal{B}(p)$

 $\mu(\mathbf{m}) = \boldsymbol{p}^{\boldsymbol{e}(\mathbf{m})}$

Candidate random graph models

Erdös-Rényi model (ER) : Edges X_{ij}'s are i.i.d. ∼ B(p)

 $\mu(\mathbf{m}) = p^{e(\mathbf{m})}$

• Mixture of ER model (Mixnet/SBM) with Q groups, proportions $\alpha_1, \ldots, \alpha_Q$ and connection probabilities $\pi_{q,\ell}$

$$\mu(\mathbf{m}) = \sum_{c_1=1}^{Q} \dots \sum_{c_k=1}^{Q} \alpha_{c_1} \dots \alpha_{c_k} \prod_{1 \le u < v \le k} \pi_{c_u, c_v}^{m_{uv}}.$$

Motif count distribution

- Exact distribution unknown.
- Several approximations exist in the literature under specific conditions (motif and model) :
 - Poisson distribution [Bollobas (81), Barbour (82), Karónski and Ruciński (83)]
 - Gaussian distribution [Barbour et al. (87)]
 - Compound Poisson distribution [Stark (01)]

Compound Poisson distribution

• Distribution of
$$\sum_{i=1}^{Z} T_i$$
 when $Z \sim \mathcal{P}(\lambda)$ and T_i 's iid.

- Particularly adapted for the count of clumping events : Z is the number of clumps and T_i is the size of the *i*-th clump.
- All network motifs are overlapping : they occur in clumps.
- We proposed to use a Geometric-Poisson(λ, a) distribution, i.e. when T_i ≈ G(1 − a)
 - analogy with sequence motifs [S. (95)],
 - (λ, a) can be calculated according to EN(m) and VarN(m):

$$m{a} = rac{\mathbb{E} N(m{m}) - \mathbb{V} N(m{m})}{\mathbb{E} N(m{m}) + \mathbb{V} N(m{m})} \quad ext{ and } \quad \lambda = (1-m{a}) \mathbb{E} N(m{m}).$$

Simulation study

Model = mixnet with 2 groups, n = 200, etc. For expectedly frequent motifs :



Gaussian (-), Poisson (-) and Geometric-Poisson (-)

Simulation study (fol.)

For expectedly rare motifs :



Gaussian (-), Poisson (-) and Geometric-Poisson (-)

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Topological motifs

Application to the *H. pylori* PPI network

- PPI network : 706 proteins and 1420 interactions (edges).
- Mixnet was fitted to the network \rightarrow 4 groups of connectivity.

Motif	N _{obs}	$\mathbb{E}_{mixnet} N$	$\sigma_{mixnet}(N)$	$\mathbb{P}(\mathcal{GP} \leq \mathit{N_{obs}})$	$\mathbb{P}(\mathcal{GP} \geq \mathit{N_{obs}})$
\mathbf{V}	14113	13602	2659		4.06 10 ⁻¹
∇	75	66.9	20.4		3.31 10 ⁻¹
	98697	94578	27039		4.1210 ⁻¹
	112490	93741	27257		2.34 10 ⁻¹
	1058	516.6	208.7		1.33 10 ⁻²
Z	3535	2897	1120		2.63 10 ⁻¹
	79	34.8	20.0		3.11 10 ⁻²
\square	0	0.17	0.45	8.5 10 ⁻¹	

Part 3

Assessing the significance of coloured motif frequencies

Coloured motifs

Graph : n nodes coloured with colours in C

Coloured motif m of size *k* is a multiset of *k* colours $\{m_1, \ldots, m_k\} \in C^k$.

Multiplicity of colour c in m: $s_m(c) = s(c)$.

Indicator of occurrence at position α : $Y_{\alpha}(\mathbf{m})$

Number of occurrences : $N(\mathbf{m}) = \sum_{\alpha \in I_k} Y_{\alpha}(\mathbf{m}).$



Model for coloured graph

- Topology : Erdös-Rényi model with probability p
- Colours : Let *f* be a probability measure on *C*; Nodes are coloured independently in color *c* ∈ *C* with probability *f*(*c*).

This model allows to derive analytical formulas for mean and (co)variance of motif counts [*Schbath, Lacroix, Sagot (09)*]

The motif count distribution is then approximated by a Geometric-Poisson distribution.

 \longrightarrow approximate *p*-value $\mathbb{P}(N(\mathbf{m}) \ge N^{\text{obs}}(\mathbf{m}))$.

Coloured motifs

Coloured motifs : Expected count

$$\mathbb{E}N(\mathbf{m}) = \sum_{\alpha \in I_k} \mathbb{E}Y_{\alpha}(\mathbf{m}) = \binom{n}{k} \mathbb{P}(\mathbf{m} \text{ occurs at } \alpha)$$
$$= \binom{n}{k} g(k, p) \times \underbrace{\frac{k!}{\prod_{c \in \mathcal{C}} s(c)!} \prod_{i=1}^k f(m_i)}_{:=\gamma(\mathbf{m})}$$

where g(k, p) is the probability for an ER(p) graph of size k to be connected [*Gilbert, 59*]:

$$g(k,p) = 1 - \sum_{i=1}^{k-1} {\binom{k-1}{i-1}} g(i,p)(1-p)^{i(k-i)}.$$

(g(1, p) = 1).

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Coloured motifs : Variance of the count (1/2)

Let us just compute $\mathbb{E}N^2(\mathbf{m})$.

$$\mathbb{E}N^{2}(\mathbf{m}) = \sum_{\alpha \in I_{k}} \sum_{\beta \in I_{k}} \mathbb{E}[Y_{\alpha}(\mathbf{m})Y_{\beta}(\mathbf{m})].$$
$$= \sum_{\ell=0}^{k} \sum_{|\alpha \cap \beta| = \ell} \underbrace{\mathbb{P}(\mathbf{m} \text{ occurs at } \alpha \text{ and } \beta)}_{=K(\alpha,\beta) \times \mathsf{Q}_{\mathbf{m}}(\alpha,\beta)}.$$

where

$$\begin{array}{lll} \mathcal{K}(\alpha,\beta) &=& \mathbb{P}(\mathcal{G}(\alpha) \text{ and } \mathcal{G}(\beta) \text{ are connected}) \\ \mathcal{Q}_{\mathbf{m}}(\alpha,\beta) &=& \mathbb{P}(\mathcal{C}(\alpha)=\mathcal{C}(\beta)=\{m_1,\ldots,m_k\}). \end{array}$$

Coloured motifs

Coloured motifs : variance of the count (2/2)

color term :

$$Q_{\mathbf{m}}(\alpha,\beta) = \mathbb{P}(C(\alpha) = C(\beta) = \{m_1,\ldots,m_k\}).$$
$$= \sum_{\mathbf{m}^* \subset \mathbf{m}} \frac{\gamma(\mathbf{m}^*)[\gamma(\mathbf{m}^-)]^2}{s(\mathbf{m}^*)}$$

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connectedness term :

$$\begin{split} \mathcal{K}(\alpha,\beta) &= & \mathbb{P}(\mathcal{G}(\alpha) \text{ and } \mathcal{G}(\beta) \text{ are connected}) \\ &= & \left\{ \begin{array}{ll} g(k,p), & \text{ if } \ell = k \\ g^2(k,p), & \text{ if } \ell = 0 \text{ or } 1. \end{array} \right. \end{split}$$

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(

$$egin{array}{rcl} \mathcal{K}(lpha,eta) &=& \mathbb{P}(\mathcal{G}(lpha) ext{ and } \mathcal{G}(eta) ext{ are connected}) \ &=& \left\{egin{array}{ll} g(k,p), & ext{if } \ell=k \ g^2(k,p), & ext{if } \ell=0 ext{ or } 1. \ ext{ad-hoc polynoms otherwise} \ 4p^3-3p^4 & ext{if } \ell=3 ext{ and } k=3 \ ext{ etc.} \end{array}
ight.$$

Geometric-Poisson approximation (1/2)

Both parameters of the GP distribution can be derived from the first 2 moments of the count.



Gaussian (red curve), Geometric-Poisson (green curve)

Geometric-Poisson approximation (2/2)

Both parameters of the PA distribution can be derived from the first 2 moments of the count.



Gaussian (red curve), Geometric-Poisson (green curve)

Part 4

Comparison of reaction networks (ongoing work with S. Robin and L. Benaroya)

Aim

Let G_1 and G_2 be two coloured graphs of size n_1 and n_2 (typically reaction networks from 2 different species).

Each graph is caracterized by the count vector of *M* given motifs of size $k : \mathbf{N}_g = (N_{g1}, N_{g2}, \dots, N_{gM})$, for g = 1, 2.

Questions :

- Do they share common exceptional motifs?
- Have both graphs similar *k*-motif compositions?
- Whose motifs are the most discriminant?
- Idea : to define a motif-based distance taking care of
 - the deviations from the models,
 - the dependence between motif counts.

Motif-based distance

Normalization by the size of the graphs : Since $\mathbb{E}N(\mathbf{m}) = \binom{n}{k} \mathbb{P}(\mathbf{m} \text{ occurs at } \alpha)$, we define :

$$\widetilde{N}_{gm} = {\binom{n_g}{k}}^{-1} N_{gm}, \quad g = 1, 2$$

Box-Cox Transformation to make the counts "more" Gaussian :

$$N^*_{gm}=2(\sqrt{\widetilde{N}_{gm}}-1), \quad g=1,2$$

Euclidian distance on z-scores :

$$d^2(\boldsymbol{\mathsf{N}}_1^*,\boldsymbol{\mathsf{N}}_2^*) = ||(\boldsymbol{\Sigma}_1^*)^{-1/2}(\boldsymbol{\mathsf{N}}_1^*-\mathbb{E}\boldsymbol{\mathsf{N}}_1^*)-(\boldsymbol{\Sigma}_2^*)^{-1/2}(\boldsymbol{\mathsf{N}}_2^*-\mathbb{E}\boldsymbol{\mathsf{N}}_2^*)||_2^2$$

where $\mathbb{E}\mathbf{N}_{g}^{*}$ and the covariance matrix $\Sigma_{g}^{*} = (\operatorname{Cov}(N_{g}^{*}(\mathbf{m}_{i}), N_{g}^{*}(\mathbf{m}_{j})))_{i,j}$ can be calculated from $\mathbb{E}\mathbf{N}_{g}$ and the covariance matrix $(\operatorname{Cov}(N_{g}(\mathbf{m}_{i}), N_{g}(\mathbf{m}_{j})))_{i,j}$ (previous part).

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Sequential distance

Onsider all single motif sets $(\dim(\mathbf{N}_1^*)=\dim(\mathbf{N}_2^*)=1)$, and take

$$\widehat{\boldsymbol{\mathsf{m}}}^1 = \text{argmax}_{\boldsymbol{\mathsf{m}}_1,\dots,\boldsymbol{\mathsf{m}}_M} \boldsymbol{\mathit{d}}^2(\boldsymbol{\mathsf{N}}_1^*,\boldsymbol{\mathsf{N}}_2^*)$$

② Consider all motif pairs $(\widehat{\mathbf{m}}^1, \mathbf{m}_j)$ with $\mathbf{m}_j \neq \widehat{\mathbf{m}}^1$, and take

$$\widehat{\boldsymbol{m}}^2 = \text{argmax}_{\boldsymbol{m}_{j} \neq \widehat{\boldsymbol{m}}^1} \boldsymbol{d}^2(\boldsymbol{N}_1^*, \boldsymbol{N}_2^*)$$



Exemple (1/2)

Reaction networks with threshold 3 on the EC numbers :

	Escherichia coli	Buchnera aphidicola
number of nodes	886	248
number of edges	4630	473
number of colors	107	62
motifs of size 3	6402	597

Exemple (2/2)

Rank		Motif		Cumulative distance
1	{ 2.7.1	2.7.4	6.3.4}	300.90
2	${1.1.1}$	1.3.1	1.14.14}	577.15
3	{1.1.1	1.1.1	1.6.1}	835.41
4	{1.1.1	1.6.1	2.5.1}	1029.83
5	{1.1.1	2.3.1	2.3.1}	1177.35
6	{2.3.1	2.3.1	2.5.1}	1324.18
7	{2.7.1	2.7.4	2.7.9}	1467.98
8	{2.7.1	2.7.2	2.7.4}	1606.38
9	{ 2.7.4	2.7.4	6.3.1}	1747.93
10	{ 2.7.4	2.7.4	2.7.10}	1876.69
11	{1.1.1	1.2.1	1.6.1}	2003.00
12	{ 2.7.4	2.7.4	3.6.4}	2127.12
13	{2.3.1	3.1.2	6.2.1}	2250.14
14	{2.7.1	2.7.4	6.3.3}	2372.24
15	{1.1.1	1.6.1	3.5.1}	2489.55

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Another approach

- To model the vector $\mathbf{N} = (N_1, N_2, \dots, N_M)$
- Need for a "multidimensional (compound) Poisson distribution with given covariance matrix"
- Our choice = the multivariate Poisson-log normal distribution from [*Aitchison and Ho, 89*] :

$$egin{array}{rcl} \mathsf{N}_m &\sim & \mathcal{P}(oldsymbol{e}^{\lambda_m}) \ \mathsf{\Lambda} &\sim & \mathcal{N}(oldsymbol{\mu},oldsymbol{\Sigma}) \end{array}$$

- (μ, Σ) can be explicitly derived from the expectation and covariance matrix of N.
- Distance = euclidian distance between Λ_1 and Λ_2
- Λ is estimated by $\mathbb{E}(\Lambda \mid \mathbf{N})$

Limitations :

- Σ may be not positive
- No analytical expression for $\mathbb{E}(\Lambda \mid \mathbf{N})$ (Monte Carlo)

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R package (Mixer) and C++ program (Mixnet) on www.ssbgroup.fr An R package nemo soon available for network motif analysis.

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Sophie Schbath (INRA -MIG)

Statistics for biological networks

Comparison of reaction networks

Multivariate Poisson-log normal distribution



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Orsay, 23 juin 2011 54 / 56

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