Statistical and computational methods for bioinformatics and social network analysis

or how did I learn to stop worrying and love the bomb

George G Vega Yon

University of Southern California, Department of Preventive Medicine

October 11, 2019





Statistical and computational methods for bioinformatics and social network analysis

- ▶ We live in a non-*IID* world.
- ► Some times, looking the whole helps understanding the parts.
- ▶ We have the computational tools to do such.

Contents



Paper 1: Exponential Random Graph Models for Small Networks

Paper 2: On the prediction of gene functions using phylogenetic trees

Exponential Random Graph Models for Small Networks

Joint with: Andrew Slaughter and Kayla de la Haye

What are Exponential Random Graph Models

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Exponential Family Random Graph Models, aka ERGMs are:

What are Exponential Random Graph Models

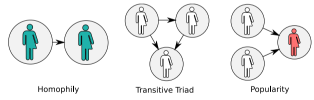
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Exponential Family Random Graph Models, aka ERGMs are:

► Statistical models of (social) networks

Exponential Family Random Graph Models, aka ERGMs are:

- ► Statistical models of (social) networks
- ▶ In simple terms: statistical inference on what network patterns/structures/motifs govern social networks



A vector of model parameters

A vector of sufficient statistics

$$\Pr\left(\mathbf{Y} = \mathbf{y} \mid \theta, \mathbf{X}\right) = \frac{\exp\left\{\theta^{\mathbf{t}} s\left(\mathbf{y}, \mathbf{X}\right)\right\}}{\sum_{\mathbf{y}' \in \mathcal{Y}} \exp\left\{\theta^{\mathbf{t}} s\left(\mathbf{y}', \mathbf{X}\right)\right\}}, \quad \forall \mathbf{y} \in \mathcal{Y}$$
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The normalizing constant

All possible networks



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$$\mathsf{All\ possible\ networks}$$

$$\mathsf{Constant}$$

The normalizing constant has $2^{n(n-1)}$ terms!



ERGMs: State of the Art

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ERGMs: State of the Art



Medium-large (dozens to a couple of thousand vertices) networks

- ► Maximum Pseudo Likelihood (MPLE)

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- ► Markov Chain Monte Carlo (MCMC) based approaches like MC-MLE or Robbins-Monro Stochastic Approximation.

 details
- ► Maximum Pseudo Likelihood (MPLE)

large-huge networks (up to the millions of vertices)

- ► Semi-parametric bootstrap
- Conditional joint estimation (like snowball sampling, a.k.a. divide and conquer)
- ► Equilibrium Expectation Algorithm (millions of vertices)

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What about small networks?

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We see small networks everywhere

► Families and friends

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- ► Families and friends
- ► Small teams

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- ► Families and friends
- ► Small teams
- ► Egocentric networks



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- ▶ It is not MLE...



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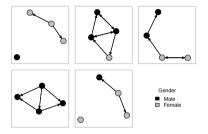


Figure 1 Random sample of 5 networks simulated using the ergmito package

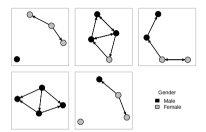


Figure 1 Random sample of 5 networks simulated using the ergmito package

| | Bernoulli | Full model |
|-----------------------|-----------|------------|
| Edge-count | -0.69* | -1.70** |
| | (0.27) | (0.54) |
| Homophily (on Gender) | | 1.59^{*} |
| | | (0.64) |
| AIC | 78.38 | 73.34 |
| BIC | 80.48 | 77.53 |
| Log Likelihood | -38.19 | -34.67 |
| Num. networks | 5 | 5 |

Standard errors in parenthesis. *** p < 0.001, ** p < 0.01, * p < 0.05

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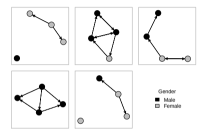


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| We performed a large simulation study prove | comparing MC-MLE (ergm) with MLE (ergmito). |

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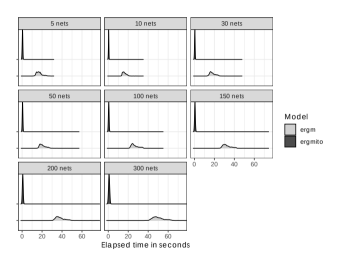
Standard errors in parenthesis. ***p < 0.001, **p < 0.01, *p < 0.05

| | | P(Type I error) | | |
|-------------|----------------|-----------------|---------------|-----------|
| Sample size | N. Simulations | MC-MLE (ergm) | MLE (ergmito) | χ^2 |
| 5 | 2,189 | 0.084 | 0.057 | 11.71 *** |
| 10 | 2,330 | 0.070 | 0.045 | 12.46 *** |
| 15 | 2,395 | 0.084 | 0.066 | 5.55 * |
| 20 | 2,430 | 0.074 | 0.060 | 3.58 |
| 30 | 2,460 | 0.057 | 0.052 | 0.67 |
| 50 | 2,495 | 0.046 | 0.044 | 0.17 |
| 100 | 2,499 | 0.048 | 0.048 | 0.00 |

Table 2 Empirical Type I error rates. The χ^2 statistic is from a 2-sample test for equality of proportions, and the significance levels are given by *** p < 0.001, ** p < 0.01, and * p < 0.05. The lack of fitted samples in some levels is due to failure of the estimation method.

Paper 1 Simulation Studies: Elapsed time

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▶ more results

Paper 1: Exponential Random Graph Models for Small Networks



Key takeaways

- ▶ New extension of ERGMs using exact statistics for small networks (families, teams, etc.)
- ▶ Performance: Same (un)bias, Lower Type I error rates, (way) faster.
- ▶ Opens the door the new methods, e.g. Mixed effects, LRT, etc.

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Next steps

- ► Revisit measurement of goodness-of-fit.
- ► Explore extending this method for (very) large networks.

On the prediction of gene functions using phylogenetic trees

Joint with: Paul D Thomas, Paul Marjoram, Huaiyu Mi, Duncan Thomas, and John Morrison

Genes

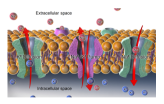


How we organize the information about genes (according to the Gene Ontology Project)

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Molecular function

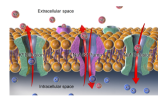
Active transport GO:0005215



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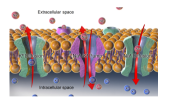
Cellular component

Mitochondria GO:0004016



How we organize the information about genes (according to the Gene Ontology Project)

Molecular function Active transport GO:0005215



Cellular component Mitochondria GO:0004016



Biological process Heart contraction GO:0060047







(filling)

The Gene Ontology Project

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- \blacktriangleright Of all annotations, about \sim 500,000 are on human genes.
- ▶ Knowledge about gene functions can accelerate bio-medical research.

Example of GO term

| Accession | GO:0060047 |
|------------------|---|
| Name | heart contraction |
| Ontology | biological_process |
| Synonyms | heart beating, cardiac contraction, hemolymph circulation |
| Alternate | IDs None |
| Definition | The multicellular organismal process in which the heart decreases in volume |
| | in a characteristic way to propel blood through the body. Source: GOC:dph |

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You know what is interesting about this function?

These four species have a gene with that function...



Felis catus pthr10037



Anolis carolinensis pthr11521



Oryzias latipes pthr11521



Equus caballus pthr24356

These four species have a gene with that function... and two of these are part of the same evolutionary tree!



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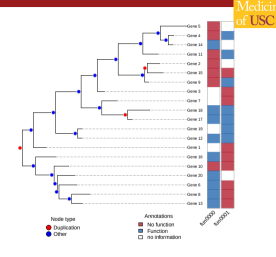


Figure 2 Random annotated phylogenetic tree.

We can use

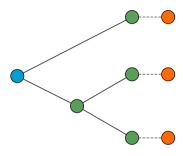
the evolutionary tree

to infer presence/absence of

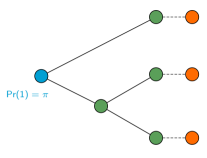
gene functions (annotations)!

An evolutionary model of gene functions





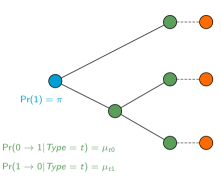
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An evolutionary model of gene functions

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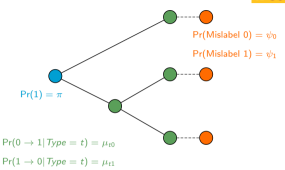
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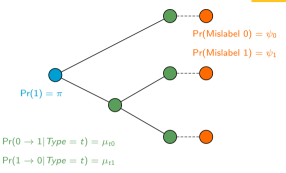
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We implemented the model using Felsenstein's' pruning algorithm (linear complexity) in the R package aphylo • more.

| | (1) | (2) |
|--------------------|---------|------|
| Mislab. prob. | | |
| ψ_0 | 0.23 | 0.25 |
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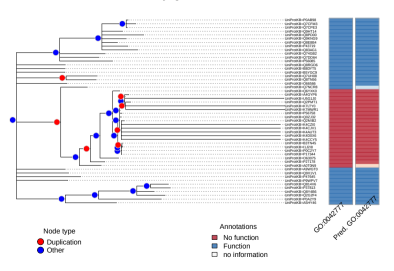
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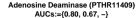
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- ▶ Took about 5 minutes each.

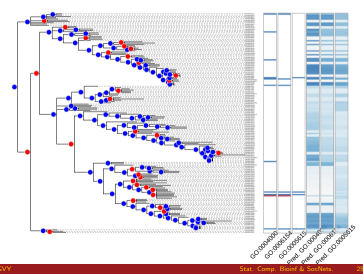
Annotated Phylogenetic Tree



Prediction with real data: Out-of-sample prediction

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Key takeaways

- ► Yet another model for predicting gene functions using phylogenetics.
- ▶ Big difference, this is computationally scalable. SIFTER (our benchmark) would take about 66 years (yes, years) to estimate a model for 100 families of size 300, we take about 5 minutes.
- ► Meaningful biological results.
- ▶ Preliminary accuracy results comparable to state-of-the-art phylo-based models.

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Next steps

▶ Adapt the model to incorporate joint estimation of functions using pseudo-likelihood.

$$P(a,b,c) \approx P(a,b)P(b,c)P(a,c)$$

▶ Make the model hierarchical when pooling trees: different mutation rates.

Statistical and computational methods for bioinformatics and social network analysis

or how did I learn to stop worrying and love the bomb

George G Vega Yon

University of Southern California, Department of Preventive Medicine

October 11, 2019



Thanks!

References I



Dodd, D. M. B. (1989). Reproductive isolation as a consequence of adaptive divergence in drosophila pseudoobscura. *Evolution*, 43(6), 1308–1311. Retrieved from http://www.jstor.org/stable/2409365

Here are some by-products of my research here at USC

- ► The slurmR R package
- ► The pruner C++ library
- ► The fmcmc R package

Sufficient statistics have various forms

| Representation | Description |
|---|--|
| $\bigcirc \longleftrightarrow \bigcirc$ | Mutual Ties (Reciprocity) $\sum_{i\neq j} y_{ij} y_{ji}$ |
| | Transitive Triad (Balance) $\sum_{i \neq j \neq k} y_{ij} y_{jk} y_{ik}$ |
| → | Homophily $\sum_{i \neq j} y_{ij} 1(x_i = x_j)$ |
| | Covariate Effect for Incoming Ties $\sum_{i\neq j} y_{ij} x_j$ |
| → | Four Cycle $\sum_{i \neq j \neq k \neq l} y_{ij} y_{jk} y_{kl} y_{li}$ |

One of the most popular methods for estimating ERGMs is the MC-MLE approach (citations here) This consists on the following steps

- 1. Start from a sensible guess on what should be the population parameters (usually done using pseudo-MLE estimation)
- 2. While the algorithm doesn't converge, do:
 - 2.1 Simulate a stream of networks with the current state of the parameter, θ_t
 - 2.2 Using the law of large numbers, approximate the ratio of likelihoods based on the parameter θ_t , this is the objective function
 - 2.3 Update the parameter by a Newton-Raphson step
 - 2.4 Next iteration



- ► Implements estimation of ERGMs using exact statistics for small networks
- ▶ Meta-programming allows specifying likelihood (and gradient) functions for joint models
- ► Includes tools for simulating, and post-estimation checks
- ► Getting ready for CRAN!





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- ► Each group could have from 5 to 300 small networks

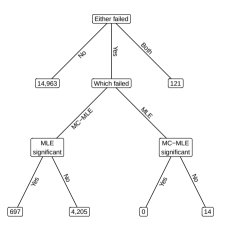




We performed a simulation study with the following features:

- ▶ Draw 20,000 samples of groups of small networks
- ► Each group had prescribed: (model parameters, number of networks, sizes of the networks)
- ► Each group could have from 5 to 300 small networks
- ▶ We estimated the models using MC-MLE and MLE.

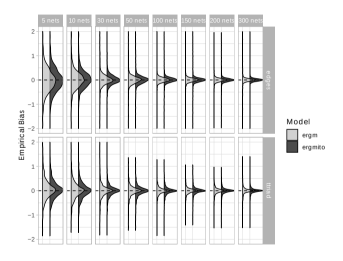






Paper 1 Simulation Studies: Empirical Bias

Keck School of Medicine of USC





An evolutionary model of gene functions (algorithmic view)

Keck School of Medicine of USC

```
Data: A phylogenetic tree, \{\pi, \mu, \psi\} (Model probabilities)
Result: An annotated tree
for n \in PostOrder(N) do
   Nodes gain/loss function depending on their parent;
   switch class of n do
       case root node do
           Gain function with probability \pi:
       case interior node do
           if Parent has the function then Keep it with prob. (1 - \mu_1):
           else Gain it with prob. \mu_0:
   end
   Finally, we allow for mislabeling:
   if n is leaf then
       if has the function then Mislabel with prob. \psi_1:
       else Mislabel with prob. \psi_0:
end
```



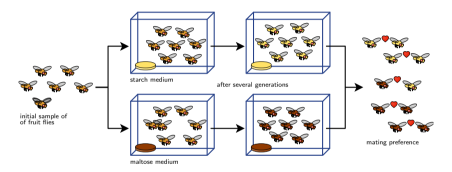


Figure 3 11989DoddDodd (): After one year of isolation, flies showed a significant level or assortativity in mating (wikimedia)



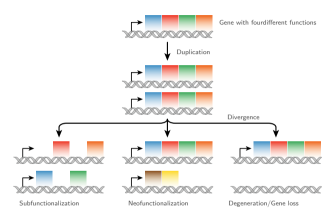


Figure 4 A key part of molecular innovation, gene duplication provides opportunity for new functions to emerge (wikimedia)



GGVY

- ▶ Pruning algorithm implemented in C++ using the pruner template library (implemented in this project).
- ▶ The estimation is done using either Maximum Likelihood, Maximum A Posteriory, or MCMC.
- ► The MCMC estimation is done via the fmcmc R package using adaptive MCMC (also implemented as part of this project)

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