# The Generalized Neighbor Joining method

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# Challenge

We would like to assemble the fungi tree of life.

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1500+ fungal species



http://ocid.nacse.org/research/aftol/about.php

# Many problems to be solved....

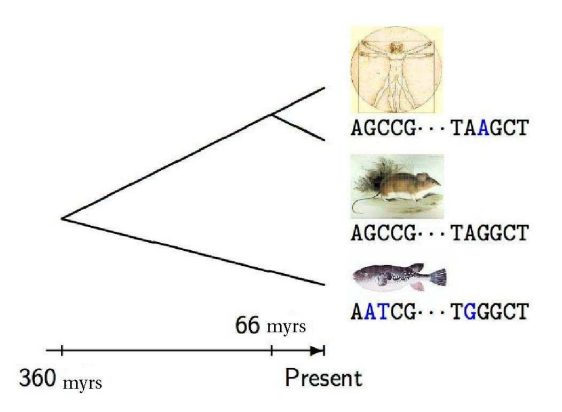


http://tolweb.org/tree?group=fungi

Zygomycota is not monophyletic. The position of some lineages such as that of Glomales and of Engodonales-Mortierellales is unclear, but they may lie outside Zygomycota as independent lineages basal to the Ascomycota-Basidiomycota lineage (Bruns et al., 1993).

# Phylogeny

Phylogenetic trees describe the evolutionary relations among groups of organisms.



### **Constructing trees from sequence data**

"Ten years ago most biologists would have agreed that all organisms evolved from a single ancestral cell that lived 3.5 billion or more years ago. More recent results, however, indicate that this family tree of life is far more complicated than was believed and may not have had a single root at all." (W. Ford Doolittle, (June 2000) *Scientific American*).

Since the proliferation of Darwinian evolutionary biology, many scientists have sought a coherent explanation from the evolution of life and have tried to reconstruct phylogenetic trees. Methods to reconstruct a phylogenetic tree from DNA sequences include:

- The maximum likelihood estimation (MLE) methods: They describe evolution in terms of a discrete-state continuous-time Markov process. The substitution rate matrix can be estimated using the expectation maximization (EM) algorithm. (for eg. Dempster, Laird, and Rubin (1977), Felsenstein (1981)).
- Distance based methods: It computes pair-wise distances, which can be obtained easily, and combinatorially reconstructs a tree. The most popular method is the neighbor-joining (NJ) method. (for eg. Saito and Nei (1987), Studier and Keppler (1988)).

#### However

**The MLE methods**: An exhaustive search for the ML phylogenetic tree is computationally prohibitive for large data sets.

**The NJ method**: The NJ phylogenetic tree for large data sets loses so much sequence information.

Goal:

- Want an algorithm for phylogenetic tree reconstruction by combining the MLE method and the NJ method.
- Want to apply methods to very large datasets.

**Note**: An algebraic view of these discrete stat problems might help solve this problem.

# The generalized neighbor-joining mathod

**The GNJ method**: in 2005, Levy, Y., and Pachter introduced the **generalized neighbor-joining (GNJ) method**, which reconstructs a phylogenetic tree based on comparisons of subtrees rather than pairwise distances

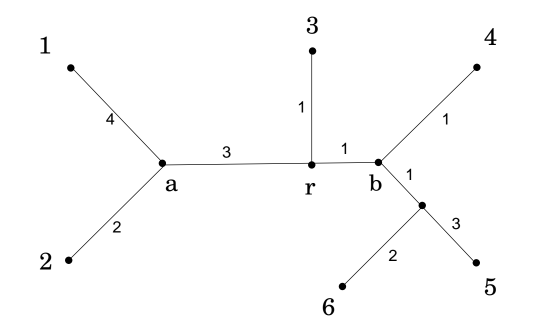
- The GNJ method is a method combined with the MLE method and the NJ method.
- The GNJ method uses more sequence information: the resulting tree should be more accurate than the NJ method.
- The computational time: polynomial in terms of the number of DNA sequences.

# The GNJ method

#### MJOIN is available at http://bio.math.berkeley.edu/mjoin/.

#### **Distance Matrix**

A distance matrix for a tree T is a matrix D whose entry  $D_{ij}$  stands for the mutation distance between i and j.



### **Distance Matrix**

	1	2	3	4	5	6
1	0	6	8	9	12	11
2	6	0	6	7	10	9
3	8	6	0	3	6	5
4	9	7	3	0	5	4
5	12	10	6	5	0	5
6	11	9	5	4	5	0

Table 1: Distance matrix D for the example.

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# Definitions

**Def.** A distance matrix D is a **metric** iff D satisfies:

- Symmetric:  $D_{ij} = D_{ji}$  and  $D_{ii} = 0$ .
- Triangle Inequality:  $D_{ik} + D_{jk} \ge D_{ij}$ .

**Def.** D is an **additive metric** iff there exists a tree T s.t.

- Every edge has a positive weight and every leaf is labeled by a distinct species in the given set.
- For every pair of i, j,  $D_{ij}$  = the sum of the edge weights along the path from i to j.

Also we call such T an **additive tree**.

## **Neighbor Joining method**

**Def.** We call a pair of two distinct leaves  $\{i, j\}$  a **cherry** if there is exactly one intermediate node on the unique path between i and j.

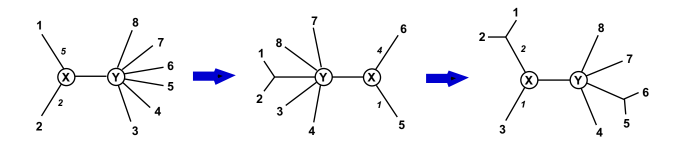
Thm. [Saitou-Nei, 1987 and Studier-Keppler, 1988]

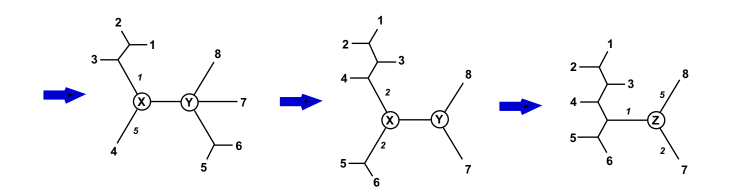
Let  $A \in \mathbb{R}^{n \times n}$  such that  $A_{ij} = D(ij) - (r_i + r_j)/(n-2)$ , where  $r_i := \sum_{k=1}^{n} D(ik)$ .  $\{i^*, j^*\}$  is a cherry in T if  $A_{i^*j^*}$  is a minimum for all i and j.

#### **Neighbor Joining Method**:

**Input.** A tree matric D. **Output.** An additive tree T. **Idea.** Initialize a star-like tree. Then find a cherry  $\{i, j\}$  and compute branch length from the interior node x to i and from x to j. Repeat this process recursively until we find all cherries.

# **Neighbor Joining Method**





# The GNJ method

- Extended the Neighbor Joining method with the total branch length of m-leaf subtrees.
- Increasing  $2 \le m \le n-2$ , since there are more data, a reconstructed tree from GNJ method gets closer to the true tree than the Saito-Nei NJ method.
- The time complexity of GNJ method is  $O(n^m)$ .

**Note**: If m = 2, then GNJ method is the Neighbor Joining method with pairwise distances.

#### Notation and definitions

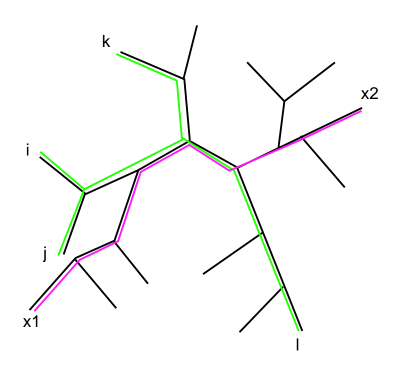
**Notation.** Let [n] denote the set  $\{1, 2, ..., n\}$  and  $\binom{[n]}{m}$  denote the set of all *m*-element subsets of [n].

**Def.** A *m*-dissimilarity map is a function  $D : {\binom{[n]}{m}} \to \mathbb{R}_{\geq 0}$ .

In the context of phylogenetic trees, the map  $D(i_1, i_2, ..., i_m)$  measures the weight of a subtree that spans the leaves  $i_1, i_2, ..., i_m$ .

Denote  $D(i_1 i_2 ... i_m) := D(i_1, i_2, ..., i_m).$ 

# Weights of Subtrees in ${\cal T}$



D(ijkl) is the total branch length of the subtree in green. Also  $D(x_1x_2)$  is the total branch length of the subtree in pink and it is also a pairwise distance between  $x_1$  and  $x_2$ .

**Thm.** [Levy, Y., Pachter, 2005] Let  $D_m$  be an *m*-dissimilarity map on *n* leaves of a tree *T*,  $D_m : {[n] \choose m} \to \mathbb{R}_{\geq 0}$  corresponding *m*-subtree weights, and define

$$\mathbf{S}(\mathbf{ij}) := \sum_{\mathbf{X} \in \binom{[\mathbf{n}] \setminus \{\mathbf{i}, \mathbf{j}\}}{\mathbf{m} - 2}} \mathbf{D}_{\mathbf{m}}(\mathbf{ijX}).$$

Then S(ij) is a tree metric.

Furthermore, if T' is based on this tree metric S(ij) then T' and T have the same tree topology and there is an invertible linear map between their edge weights.

**Note.** This means that if we reconstruct T', then we can reconstruct T.

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#### **Neighbor Joining with Subtree Weights**

**Input**: *n* DNA sequences and an integer  $2 \le m \le n-2$ .

**Output**: A phylogenetic tree T with n leaves.

- 1. Compute all *m*-subtree weights via the ML method.
- 2. Compute S(ij) for each pair of leaves i and j.
- 3. Apply Neighbor Joining method with a tree metric S(ij) and obtain additive tree T'.
- 4. Using a one-to-one linear transformation, obtain a weight of each internal edge of T and a weight of each leaf edge of T.

## Complexity

**Lemma.** [Levy, Pachter, Y.] If  $m \ge 3$ , the time complexity of this algorithm is  $O(n^m)$ , where n is the number of leaves of T and if m = 2, then the time complexity of this algorithm is  $O(n^3)$ .

**Sketch of Proof**: If  $m \ge 3$ , the computation of S(ij) is  $O(n^m)$  (both steps are trivially parallelizable). The subsequent neighbor-joining is  $O(n^3)$  and edge weight reconstruction is  $O(n^2)$ . If m = 2, then the subsequent neighbor-joining is  $O(n^3)$  which is greater than computing S(ij). So, the time complexity is  $O(n^3)$ .

**Note**: The running time complexity of the algorithm is  $O(n^3)$  for both m = 2 and m = 3.

# **Cherry Picking Theorem**

**Thm.** [Levy, Pachter, Y.] Let T be a tree with n leaves and no nodes of degree 2 and let m be an integer satisfying  $2 \le m \le n-2$ . Let  $D: {[n] \choose m} \to \mathbb{R}_{\ge 0}$  be the m-dissimilarity map corresponding to the weights of the subtrees of size m in T. If  $Q_D(a^*b^*)$  is a minimal element of the matrix

$$Q_D(ab) = \left(\frac{n-2}{m-1}\right) \sum_{X \in \binom{[n-i-j]}{m-2}} D(ijX) - \sum_{X \in \binom{[n-i]}{m-1}} D(iX) - \sum_{X \in \binom{[n-j]}{m-1}} D(jX)$$

then  $\{a^*, b^*\}$  is a cherry in the tree T.

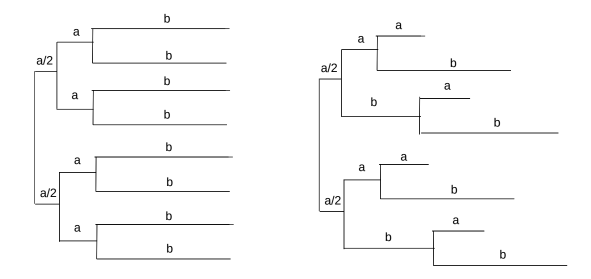
**Note.** The theorem by Saitou-Nei and Studier-Keppler is a corollary from Cherry Picking Theorem.

# Simulation Results

With the Juke Cantor model.

## Consider two tree models...

Modeled from Strimmer and von Haeseler.



T1

**T**2

We generate 500 replications with the Jukes-Cantor model via a software evolver from PAML package.

The number represents a percentage which we got the same tree topology.

Ι	a/b	m=2	m=3	4	fastDNAml
500	0.01/0.07	68.2	76.8	80.4	74.8
	0.02/0.19	54.2	61.2	73.6	55.6
	0.03/0.42	10.4	12.6	23.8	12.6
1000	0.01/0.07	94.2	96	97.4	96.6
	0.02/0.19	87.6	88.6	96.2	88
	0.03/0.42	33.4	35	52.4	33.6

Table 2: Success Rates for the model  $T_1$ .

	a/b	m=2	3	m=4	fastDNAml
500	0.01/0.07	84.4	86	85.6	88.4
	0.02/0.19	68.2	72	73.2	88.4
	0.03/0.42	18.2	29.2	36.2	87.4
1000	0.01/0.07	95.6	97.8	97.4	99.4
	0.02/0.19	88.4	89.6	93.4	99.8
	0.03/0.42	40	48.2	57.6	96.6

Table 3: Success Rates for the model  $T_2$ .

# A unifying framework: Algebraic Statistics

# What is Algebraic Statistics?

**Algebraic Statistics** is to apply computational commutative algebraic techniques to statistical problems.

The algebraic view of discrete statistical models has been applied in many statistical problems, including:

- conditional inference [Diaconis and Sturmfels 1998]
- disclosure limitation [Sullivant 2005]
- the maximum likelihood estimation [Hosten et al 2004]
- parametric inference [Pachter and Sturmfels 2004]
- phylogenetic invariants [Allman and Rhodes 2003, Eriksson 2005, etc].

#### **Algebraic statistical models**

An algebraic statistical model arises as the image of a polynomial map

$$\mathbf{f} : \mathbb{R}^d \to \mathbb{R}^m, \ \theta = (\theta_1, \dots, \theta_d) \mapsto (p_1(\theta), p_2(\theta), \dots, p_m(\theta)).$$

The unknowns  $\theta_1, \ldots, \theta_d$  represent the model parameters.

In the view of algebraic geometry, statistical models are **algebraic varieties**, sets of points where all given polynomials vanish at the same time.

**Note**: The phylogenetic models are also algebraic varieties.

Note: The MLE problem is a polynomial optimization problem over the image of  ${\bf f}.$ 

#### **Jukes-Cantor Model**

Consider the Jukes-Cantor (JC) model.

The JC model has substitution rate matrix:

$$Q = \begin{pmatrix} -3\alpha & \alpha & \alpha & \alpha \\ \alpha & -3\alpha & \alpha & \alpha \\ \alpha & \alpha & -3\alpha & \alpha \\ \alpha & \alpha & \alpha & -3\alpha \end{pmatrix}$$

where  $\alpha \geq 0$  is a parameter. The corresponding substitution matrix equals

$$\theta(t) = \frac{1}{4} \begin{pmatrix} 1+3e^{-4\alpha t} & 1-e^{-4\alpha t} & 1-e^{-4\alpha t} & 1-e^{-4\alpha t} \\ 1-e^{-4\alpha t} & 1+3e^{-4\alpha t} & 1-e^{-4\alpha t} & 1-e^{-4\alpha t} \\ 1-e^{-4\alpha t} & 1-e^{-4\alpha t} & 1+3e^{-4\alpha t} & 1-e^{-4\alpha t} \\ 1-e^{-4\alpha t} & 1-e^{-4\alpha t} & 1-e^{-4\alpha t} & 1-e^{-4\alpha t} \end{pmatrix}$$

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However, they are not polynomials.... But we can do the following: Introduce the new two parameters

$$\pi_\mathbf{i}\,=\,\frac{\mathbf{1}}{\mathbf{4}}(\mathbf{1}-\mathbf{e}^{-\mathbf{4}\alpha_\mathbf{i}\mathbf{t}_\mathbf{i}})\quad\text{and}\quad\mu_\mathbf{i}\,=\,\frac{\mathbf{1}}{\mathbf{4}}(\mathbf{1}+\mathbf{3}\mathbf{e}^{-\mathbf{4}\alpha_\mathbf{i}\mathbf{t}_\mathbf{i}}).$$

These parameters satisfy the linear constraint

 $\mu_{\mathbf{i}} + \mathbf{3}\pi_{\mathbf{i}} = \mathbf{1},$ 

and the branch length  $t_i$  of the *i*th edge can be recovered as follows:

$$\mathbf{3}lpha_{\mathbf{i}}\mathbf{t}_{\mathbf{i}} = -\frac{\mathbf{1}}{\mathbf{4}}\cdot\log\det\left(\mathbf{ heta}^{\mathbf{i}}
ight) = -\frac{\mathbf{3}}{\mathbf{4}}\cdot\log(\mathbf{1}-\mathbf{4}\pi_{\mathbf{i}}).$$

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The parameters are simply the entries in the substitution matrix

$$\theta^{i} = \begin{pmatrix} \mu_{i} & \pi_{i} & \pi_{i} & \pi_{i} \\ \pi_{i} & \mu_{i} & \pi_{i} & \pi_{i} \\ \pi_{i} & \pi_{i} & \mu_{i} & \pi_{i} \\ \pi_{i} & \pi_{i} & \pi_{i} & \mu_{i} \end{pmatrix}$$

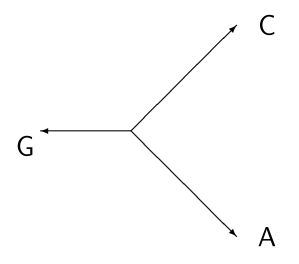
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The Jukes–Cantor model on the tree T with r edges and n leaves is the polynomial map

 $\mathbf{f} \, : \, \mathbb{R}^{\mathbf{r}} \, 
ightarrow \, \mathbb{R}^{4^{\mathbf{n}}}.$ 

#### Example

Suppose we have an unrooted tree T with leaves  $\{1, 2, 3\}$  with letters  $\Sigma = \{A, C, G, T\}$  at a single site. Want to estimate all parameters.



This model is a three-dimensional algebraic variety, given as the image of a trilinear map

$$\mathbf{f} : \mathbb{R}^3 \to \mathbb{R}^{64}.$$

#### Example cont

Let  $p_{123}$  be the probability of observing the same letter at all three leaves,  $p_{ij}$  the probability of observing the same letter at the leaves i, j and a different one at the third leaf, and  $p_{dis}$  the probability of seeing three distinct letters.

$p_{123}$	=	$\mu_1\mu_2\mu_3 + 3\pi_1\pi_2\pi_3,$
$p_{dis}$	=	$6\mu_1\pi_2\pi_3 + 6\pi_1\mu_2\pi_3 + 6\pi_1\pi_2\mu_3 + 6\pi_1\pi_2\pi_3,$
$p_{12}$	=	$3\mu_1\mu_2\pi_3 + 3\pi_1\pi_2\mu_3 + 6\pi_1\pi_2\pi_3,$
$p_{13}$	=	$3\mu_1\pi_2\mu_3 + 3\pi_1\mu_2\pi_3 + 6\pi_1\pi_2\pi_3,$
$p_{23}$	=	$3\pi_1\mu_2\mu_3 + 3\mu_1\pi_2\pi_3 + 6\pi_1\pi_2\pi_3.$

All 64 coordinates of f are given by these five trilinear polynomials, namely,

$$f_{AAA} = f_{CCC} = f_{GGG} = f_{TTT} = \frac{1}{4} \cdot p_{123},$$
  

$$f_{ACG} = f_{ACT} = \cdots = f_{GTC} = \frac{1}{24} \cdot p_{dis},$$
  

$$f_{AAC} = f_{AAT} = \cdots = f_{TTG} = \frac{1}{12} \cdot p_{12},$$
  

$$f_{ACA} = f_{ATA} = \cdots = f_{TGT} = \frac{1}{12} \cdot p_{13},$$
  

$$f_{CAA} = f_{TAA} = \cdots = f_{GTT} = \frac{1}{12} \cdot p_{23}.$$

This means that the Jukes–Cantor model is the image of the simplified map  $\mathbf{f}' : \mathbb{R}^3 \to \mathbb{R}^5, \ ((\mu_1, \pi_1), (\mu_2, \pi_2), (\mu_3, \pi_3)) \mapsto (\mathbf{p_{123}}, \mathbf{p_{dis}}, \mathbf{p_{12}}, \mathbf{p_{13}}, \mathbf{p_{23}}).$ 

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### Characterize the image of $f^\prime$

Do the following linear change of coordinates:

$$\begin{aligned} q_{111} &= p_{123} + \frac{1}{3} p_{dis} - \frac{1}{3} p_{12} - \frac{1}{3} p_{13} - \frac{1}{3} p_{23} = (\mu_1 - \pi_1)(\mu_2 - \pi_2)(\mu_3 - \pi_3) \\ q_{110} &= p_{123} - \frac{1}{3} p_{dis} + p_{12} - \frac{1}{3} p_{13} - \frac{1}{3} p_{23} = (\mu_1 - \pi_1)(\mu_2 - \pi_2)(\mu_3 + 3\pi_3) \\ q_{101} &= p_{123} - \frac{1}{3} p_{dis} - \frac{1}{3} p_{12} + p_{13} - \frac{1}{3} p_{23} = (\mu_1 - \pi_1)(\mu_2 + 3\pi_2)(\mu_3 - \pi_3) \\ q_{011} &= p_{123} - \frac{1}{3} p_{dis} - \frac{1}{3} p_{12} - \frac{1}{3} p_{13} + p_{23} = (\mu_1 + 3\pi_1)(\mu_2 - \pi_2)(\mu_3 - \pi_3) \\ q_{000} &= p_{123} + p_{dis} + p_{12} + p_{13} + p_{23} = (\mu_1 + 3\pi_1)(\mu_2 + 3\pi_2)(\mu_3 - \pi_3). \end{aligned}$$

This model is the hypersurface in  $\Delta_4$  whose ideal equals

$$\mathbf{P}_{\mathbf{f}'} = \langle \mathbf{q}_{000} \mathbf{q}_{111}^2 - \mathbf{q}_{011} \mathbf{q}_{101} \mathbf{q}_{110} \rangle.$$

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### MLE with the JC model

Suppose  $(U_{123}, U_{dis}, U_{12}, U_{13}, U_{23})$  is the observed data.

Then the MLE with the JC model for  $\boldsymbol{T}$  is

 $\max \ p_{123}^{U_{123}} p_{dis}^{U_{dis}} p_{12}^{U_{12}} p_{13}^{U_{13}} p_{23}^{U_{23}} \ \text{ subject to } \theta \in \pmb{\Theta}.$ 

**Note**: The image of  $\mathbf{f}'$  is a hyper-surface over  $\mathbb{R}^5$ .

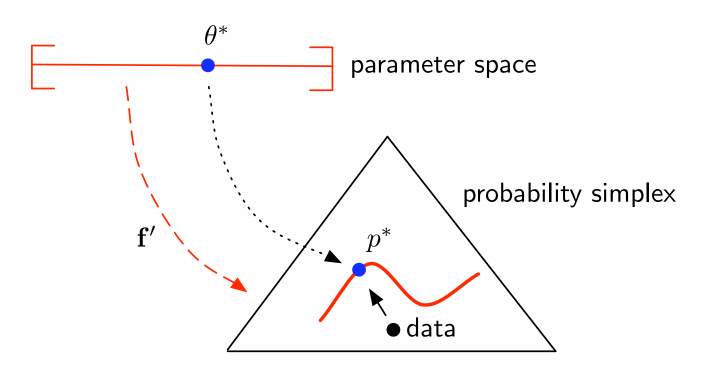


Figure 1:  $\theta^*$  is the global maxima and  $p^*$  is an image under  $\mathbf{f}'$ .

## **Commutative algebraic methods to phylogenetics.**

Using the algebraic techniques with the JC model with triplets, interval arithmetics, and the GNJ method, one can reconstruct a phylogenetic tree from DNA sequences (Sainudiin and Y. 2005).

One can find more tree invariants with the JC model, the Kimura 2parameter model (K80), and the Kimura 3-parameter model (K81) at http://www.math.tamu.edu/~lgp/small-trees/small-trees.html.

Using these invariants and the GNJ method one can reconstruct a phylogenetic tree from DNA sequences (Contois and Levy, 2005).

One can find more applications of algebra to computational biology at our new book **Algebraic Statistics for Computational Biology** edited by Pachter and Sturmfels, Cambridge University Press 2005.

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- A. Hobolth (Bioinformatics, NCSU) and R. Yoshida, "Maximum likelihood estimation of phylogenetic tree and substitution rates via generalized neighbor-joining and the EM algorithm", *Algebraic Biology* 2005, Computer Algebra in Biology, edited by H. Anai and K. Horimoto, vol. 1 (2005) p41 - 50, Universal Academy Press, INC.. (Also available at arXiv:q-bio.QM/0511034.)
- R. Sainudiin (Statistics, Oxford) and R. Yoshida, "Applications of Interval Methods to Phylogenetic trees" *Algebraic Statistics for Computational Biology* edited by Lior Pachter and Bernd Sturmfels, (2005) Cambridge University Press, p359 - 374.

# Thank you....