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Molecular Evolution and Phylogenetics ...a very very short course

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1. Introduction

1.1. Three basic questions

- Why use phylogenies?
 - Like astronomy, biology is an **historical** science!
 - The knowledge of the past is important to solve many questions related to biological patterns and processes.
- Can we know the past?
 - We can postulate alternative evolutionary scenarios (hypothesis)
 - Obtain the proper dataset and get statistical confidence
- What means to know "...the phylogeny"?
 - The ancestral-descendant relationships (tree topology)
 - The distances between them (tree branch lengths)

Phylogenies are working hypotheses!!!



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2. Tree Terminology

2.1. Topology, branches, nodes & root

• Nodes & branches. Trees contain internal and external nodes and branches. In molecular phylogenetics, external nodes are sequences representing genes, populations or species!. Sometimes, internal nodes contain the ancestral information of the clustered species. A branch defines the relationship between sequences in terms of descent and ancestry.





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- **Root** is the common ancestor of all the sequences.
- **Topology** represents the branching pattern. Branches **can rotate** on internal nodes. Instead of the singular aspect, the following trees represent a single phylogeny.



The topology is the same!!



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- **Taxa**. (*plural of taxon or operaqtional taxonomic unit (OTU)*) Any group of organisms, populations or sequences considered to be sufficiently distinct from other of such groups to be treated as a separate unit.
- **Polytomies**. Sometimes trees does not show fully bifurcated (binary) topologies. In that cases, the tree is considered **not resolved**. Only the relationships of species 1-3, 4 and 5 are known.



Polytomies can be solved by using more sequences, more characters or both!!!



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2.2. Rooted & Unrooted trees

Trees can be **rooted** or **unrooted** depending on the explicit definition or not of **outgroup** sequence or taxa.

• **Outgroup** is any group of sequences used in the analysis that is not included in the sequences under study (**ingroup**).



- Unrooted trees show the topological relationships among sequences althoug it is impossible to deduce wether nodes (n_i) represent a primitive or derived evolutionary condition.
- **Rooted trees** show the evolutionary basal and derived evolutionary relationships among sequences.

Rooting by outgroup is frequent in molecular phylogenetics!!



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2.3. Cladograms & Phylograms

Trees showing branching order exclusively (**cladogenesis**) are principally the interest of systematists¹ to make inferences on taxonomy². Those interesting in the evolutionary processes emphasize on branch lengths information (**anagenesis**).



- **Dendrogram** is a branching diagram in the form of a tree used to depict degrees of relationship or resemblance.
- **Cladogram** is a branching diagram depicting the hierarchical arrangement of taxa defined by cladistic methods (the distribution of shared derived characters -synapomorphies-).



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¹The study of biological diversity.

²The theory and practice of describing, naming and classifying organisms

- **Phylogram** is a phylogenetic tree that indicates the relationships between the taxa and also conveys a sense of time or rate of evolution. The temporal aspect of a phylogram is missing from a cladogram or a generalized dendogram.
- **Distance scale** represents the number of differences between sequences (e.g. 0.1 means 10 % differences between two sequences)



Rooted and unrooted phylograms or cladograms are frequently used in molecular systematics!



2.4. Consensus trees

It is frequent to obtain alternative phylogenetic hypothesis from a single data set. In such a case, it is usefull to summarize common or average relationships among the original set of trees. A number of different types of consensus trees have been proposed;

• The **majority rule consensus** tree uses a simple majority of relationships among the fundamental trees.



A consensus tree is a summary of how well the original trees agrees.

A helpfull manual covering these and other concepts of the section can be obtained in [20, 12].





3. Homology

The Origin of Species. Charles Darwin. Chapter 14

What can be more curious than that the hand of a man, formed for grasping, that of a mole for digging, the leg of the horse, the paddle of the porpoise, and the wing of the bat, should all be constructed on the same pattern, and should include similar bones, in the same relative positions?

Why should similar bones have been created to form the wing and the leg of a bat, used as they are for such totally different purposes, namely flying and walking?

How inexplicable are the cases of serial homologies on the ordinary view of creation!



Since Darwin homology was the result of descent with modification from a common ancestor.



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Homology Molecular Evolution **Evolutionary Models** Homology Distance Methods Maximum Parsimony Species sharing a common ancestral states Searching Trees Tree Confidence PC Lab Sp1 Sp2 Sp3 OUG Sp4 Phylogenetic Links Credits Homologous Title Page character states Sp1-Sp2 Sp3-Sp4 Page 11 of 60 Inferred tree Go Back Full Screen Close Quit

3.1. Homoplasy

- Similarity among species could represent true homology (just by sharing the same ancestral state) or, **homoplastic** events like **convergence**, **par**allelism or reversals;
- **Homology** is a *posteriori* tree construction definition.

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• Convergences are ...

Convergent evolution



Homoplasy can provide misleading evidence of phylogenetic relationships!! (if mistakenly interpreted as homology).



• Parallels are ...

Parallel evolution



Homoplasy can provide misleading evidence of phylogenetic relationships!! (if mistakenly interpreted as homology).



• **Reversions** are ...



Homoplasy can provide misleading evidence of phylogenetic relationships!! (if mistakenly interpreted as homology).



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3.2. Similarity

- For molecular sequence data, **homology** means that two sequences or even two characters within sequences are descended from a common ancestor.
- This term is frequently mis-used as a synonym of **similarity**.
- as in two sequences were 70% homologous.
- This is totally incorrect!
- Sequences show a certain amount of similarity.
- From this similarity value, we can probably infer that the sequences are homologous or not.
- Homology is like pregnancy. You are either pregnant or not.
- Two sequences are either homologous or they are not.



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3.3. Sequence homology

In molecular studies it is important to distinguish among kinds of **homology**[6];

- **Ortholog**: Homologous genes that have diverged from each other after speciation events (e.g., human β and chimp β -globin).
- Paralog: Homologous genes that have diverged from each other after gene duplication events (e.g., β- and γ-globin)





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- **Xenolog**: Homologous genes that have diverged from each other after lateral gene transfer events (e.g., antibiotic resistance genes in bacteria).
- **Homolog**: Genes that are descended from a common ancestor (e.g., all globins).







• **Positional homology**: Common ancestry of specific amino acid or nucleotide positions in different genes.

11 50462 Homo.sapie VGLLGRTGSGKSTLLSAFLRLLNTEG-EIQI: Mus.muscul VGLLGRTGSGKSTLLSAFLRMLNIKG-DIET: Fugu.rubri MGLLGRTGSGKSTLLSALLRLASTDG-EISI: Ciona.inte VGIVGRTGAGKSSLISTLFRLNEYSKGSVMI: Droso.mela VGIVGRTGAGKSSLIGALFRLANEG-EIFI: Anoph.gamb VGIVGRTGAGKSSLIGALFRLAQVEG-EIRL: Caeno.eleg VGIVGRTGAGKSSLIJALFRILEADGGSIEI: Sacch.cere IGIVGRTGAGKSSTITALFRFLEPETGHIKI: Arabi.thal IGIVGRTGSGKTTLISALFRLVEPVGGKIVV. Oryza.sati IGVVGRTGSGKSTLVQALFRLVEPVEGHIV. Plasm.falc IGIVGKSGAGKSTMILSILGLIGTTRGRITT:



4. Molecular Evolution

4.1. Molecular clock

The **molecular clock hypothesis** postulates that for any given macromolecule (a protein or DNA sequence), the rate of evolution *-measured as the mean number* of amino acids or nucleotide sequence change per site per year- is approximately constant over time in all the evolutionary lineages [21].



Fig. 8.3. Rates of amino acid substitution in the fibrinopeptides, hemoglobin, and cytochrome c. Comparisons for which no adequate time coordinate is available are indicated by numbered crosses. Point 1 represents a date of 1200 \pm 75 MV (million years) for the separation of plants and animals, based on a linear extrapolation of the cytochrome c curve. Points 2–10 refer to events in the evolution of the globin family. The δ/β separation is at point 3, γ/β is at 4, and α/β is at 500 MV (carp/lamprey). From Dickerson (1971).



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This hypothesis has estimulated much interest in the use of macromolecules in evolutionay studies for two reasons:

- Sequences can be used as molecular markers to **date** evolutionary events.
- The degree of rate change among sequences and lineages can provide insights on **mechanisms** of molecular evolution. For example, a large increase in the rate of evolution in a protein in a particular lineage may indicate adaptive evolution.

Substitution rate estimation

It is based on the number of a substitution (distance) and divergence time (fossil calibration),



There is no universal clock

It is known that **clock variation** exists for:

- different molecules, depending on their functional constraints,
- different regions in the same molecule,

Rates of amino acid substitution at the surface and heme pocket regions of the hemogle	obin
α - and β -chains (Kimura and Ohta, 1973b).	

Region	a-chain	β -chain
Surface	1.4 (18)	2.7 (23)
Heme pocket	0.17 (19)	0.24 (21)

Note: The rate represents 'per amino acid site per year'. The values in the table should be multiplied by 10⁻⁹. The figures in brackets are the number of amino acid sites involved.





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• different base position (synonimous-nonsynonimous),







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- different genomes in the same cell,
- different regions of genomes,
- different taxonomic groups for the same gene (lineage effects)



Fig. 7.14 Numbers of synonymous and non-synonymous substitutions for 49 genes from three mammalian orders: primates, rodents and artiodactyls, the phylogenetic relationships of which approximate a 'star phylogeny'. Note that, in both cases, rodents have accumulated more substitutions than primates or artiodactyls. Adapted from Ohta (1995).



5. Evolutionary Models

5.1. Multiple Hits

- The mutational change of DNA sequences varies with region. Even considering protein coding sequence alone, the patterns of nucleotide substitution at the first, second or third codon position are not the same.
- When two DNA sequences are derived from a common ancestral sequence, the descendant sequences gradually diverge by nucleotide substitution.
- A simple measure of sequence divergence is the proportion $p = N_d/N_t$ of nucleotide sites at which the two sequences are different.





• When p is large, it gives an underestimate of the number of of substitutions, because it does not take into account **multiple substitutions**.



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- Sequences may saturate due to multiple changes (hits) at the same position after lineage splitting.
- In the worst case, data may become random and all the **phylogenetic information** about relationships can be lost!!!





5.2. Models of nucleotide substitution

a

• In order to estimate **the number of nucleotide substitutions ocurred** it is necessary to use a mathematical model of nucleotide substitution. The model would consider the nucleotide frequencies and the instantaneous rate's change among them.

	Designation	Rate params	Base frequencies	Number of free params
	JC	a=b=c=d=e=f	$\pi_A=\pi_C=\pi_G=\pi_T$	1
	K80, K2P	a=c=d=f, b=e	$\pi_A = \pi_C = \pi_G = \pi_T$	2
	TrNef	a=c=d=f, b, e	$\pi_A = \pi_C = \pi_G = \pi_T$	3
	K81, K3ST	a=f, b=e, c=d	$\pi_A=\pi_C=\pi_G=\pi_T$	3
	TVMef	a, c, d, f, b=e	$\pi_A=\pi_C=\pi_G=\pi_T$	5
	TiMef	a=f, c=d, b, e	$\pi_A=\pi_C=\pi_G=\pi_T$	4
m 1.1	SYM	a, b, c, d, e, f	$\pi_A=\pi_C=\pi_G=\pi_T$	6
Transitions	F81	a=b=c=d=e	$\pi_A, \pi_C, \pi_G, \pi_T$	4
	HKY	a=c=d=f, b=e	$\pi_A, \pi_C, \pi_G, \pi_T$	5
T	TrN	a=c=d=f, b, e	$\pi_A, \pi_C, \pi_G, \pi_T$	6
Transversions	K81uf	a=f, b=e, c=d	$\pi_A, \pi_G, \pi_G, \pi_T$	6
	TVM	a, c, d, f, b=e	$\pi_A, \pi_C, \pi_G, \pi_T$	8
Transitions	TiM	a=f, c=d, b, e	$\pi_A, \pi_C, \pi_G, \pi_T$	7
Tanonono	GTR, REV	a, b, c, d, e, f	$\pi_A, \pi_G, \pi_G, \pi_T$	9



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• Interrrelationships among models for estimating the number of nucleotide substitutions among a pair of DNA sequences







• For constructing phylogenetic trees from distance measures, sophisticated distances are not neccesary more efficient.

Table 3.3 Observed numbers of the 10 pairs of nucleotides between the DNA sequences for the human and Rhesus monkey mitochondrial cytochrome b genes.

	Transition			Transversion				Identical Pair				T -+-
Codon Position	TC	AG	TA	TG	CA	CG	ΤT	CC	AA	GG	n_d	10ta (n)
First	21	22	5	1	5	4	68	93	100	56	58	375
Second	20	3	6	1	0	2	140	87	71	45	32	375
Third	60	16	6	5	49	2	11	122	102	2	138	375
All	101	41	17	7	54	8	219	302	273	103	228	112

Note: The numbers at the first, second, and third codon positions are shown separately.

• Indeed, by using sophisticated models distances show higher variance values.

Table 3.4 Estimates (\hat{d}) of the number of nucleotide substitutions per site between the human and Rhesus monkey mitochondrial cytochrome b genes for the first, second, and third codon positions $(\hat{d} \times 100)$.

Position in Codon	p	Jukes-Cantor	Kimura	Tajima-Nei	Tamura-Nei
First	15.5 ± 1.9	17.3 ± 2.4	17.8 ± 2.5	18.0 ± 2.6	17.9 ± 2.5
Second	8.5 ± 1.4	9.1 ± 1.6	9.2 ± 1.7	9.2 ± 1.7	9.3 ± 1.7
Third	36.8 ± 2.5	50.6 ± 4.9	52.3 ± 5.4	66.5 ± 9.4	87.9 ± 39.0

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• Of course, corrected distances are greather than the observed.



FIGURE 3.1. Estimates of the number of nucleotide substitutions obtained by different distance measures when actual nucleotide substitution follows the Tamura-Nei model. The nucleotide frequencies assumed are $g_A = 0.3$, $g_T = 0.4$, $g_C = 0.2$, and $g_G = 0.1$; and the two transition/transversion rate ratios assumed are $\alpha_1/\beta = 4$ and $\alpha_2/\beta = 8$.



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Distance correction methods share several assumptions:

- All nucleotide sites change independently.
- The substitution rate is constant over time and in different lineages
- The base composition is at equilibrium (all sequences have the same base frequencies)
- The conditional probabilities of nucleotide substitutions are the same for all sites and do not change over time.

While these assumptions make the methods tractable, they are in many cases unrealistic.



6. Distance Methods

Distance matrix methods is a major family of phylogenetic methods trying to fit a tree to a matrix of pairwise distance [1, 5]. Distance are generally corrected distances.

- The best way of thinking about distance matrix methods is to consider distances as estimates of the branch length separating that pair of species.
- Branch lengths are not simply a function of time, they reflect expected amounts of evolution in different branches of the tree.
- Two branches may reflect the same elapsed time (sister taxa), but they can have different expected amounts of evolution.
- The product $r_i * t_i$ is the branch length
- The main distance-based tree-building methods are cluster analysis, least square and minimum evolution.
- They rely on different assumptions, and their success or failure in retrieving the correct phylogenetic tree depends on how well any particular data set meet such assumptions.



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6.1. **Cluster Analysis**

Cluster analysis derived from clustering algorithms popularized by Sokal and Sneath [16]

6.1.1. UPGMA

One of the most popular distance approach is the **unweighted pair-group** method with arithmetic mean (UPGMA), which is also the simplest method for tree reconstruction [10].

- 1. Given a matrix of pairwise distances, find the clusters (taxa) i and jsuch that d_{ij} is the minimum value in the table.
- 2. Define the depth of the branching between i and j (l_{ij}) to be $d_{ij}/2$
- 3. If i and j are the last 2 clusters, the tree is complete. Otherwise, create a new cluster called u.
- 4. Define the distance from u to each other cluster $(k, \text{ with } k \neq i \text{ or } j)$ to be an average of the distances d_{ki} and d_{kj}
- 5. Go back to step 1 with one less cluster; clusters i and j are eliminated, and cluster u is added.

The variants of UPGMA are in the step 4. Weighted PGMA(WPGM:: $d_{ku} =$ $d_{ki}+d_{kj}/2$). Complete linkage $(d_{ku} = max(d_{ki}, d_{kj}))$. Single linkage $(d_{ku} = max(d_{ki}, d_{kj}))$. $min(d_{ki}, d_{kj}).$



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	Bsu	Bst	Lvi	Amo	Mlu						- Alt
Bsu Bst	_	0.1715	0.2147 0.2991	0.3091 0.3399 0.2795	0.2326 0.2058 0.3943						Introduction
Amo			_		0.3943						Tree Terminology
Mlu											Homology
	Bsu-Bst	Lvi	Amo	Mlu							Molecular Evoluti
Bsu-Bst Lvi Amo		0.2569	0.3245 0.2795	0.2192 0.3943 0.4289							Evolutionary Mod Distance Methods
Mlu											Maximum Parsim
	Bsı	ı-Bst-Mlu	Lvi	Amo						Bet	Searching Trees
Bsu-Bst-l Lvi	Mlu	10 <u></u>	0.3027	0.3593 0.2795						Mlu	Tree Confidence
Amo										I vi	PC Lab
	Bsı	ı-Bst-Mlu	Lvi-Amo							Amo	Phylogenetic Link
Bsu-Bst-l Lvi-Amo	Mlu	. <u></u> .	0.3310			0.20	0.15	0.10	0.05	J	Credits
						0.20	0.15	0.10	0.05	0.00	

The smallest distance in the first table is 0.1715 substitutions per sequence position separating *Bacillus subtilis* and *B. stearothermophilus*. The distance between Bsu-Bst to Lvi (*Lactobacillus viridescens*) is (0.2147+0.2991)/2=0.2569. In the second table, joins Bsu-Bst to Mlu(*Micrococcus luteus*) at the depth 0.1096(=0.2192/2). The distances Bsu-Bst-Mlu to Lvi is (2*0.2569+0.3943)/3=0.3027. Notice that this value is identical to (Bsu:Lvi+Bst:Lvi+Mlu:Lvi)/3. Each taxon in the original data table contributes equally to the averages, this is why the method called **unweighted**

UPGMA method supposes a cloclike behaviour of all the lineages, giving a rooted and ultrametric tree.

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6.1.2. NJ (Neighboor Joining)

A variety of methods related to cluster analysis have been proposed that will correctly reconstruct additive trees, whether the data are ultrametric or not. NJ removes the assumption that the data are ultrametric.

- 1. For each terminal node *i* calculate its net divergence (r_i) from all the other taxa using $\mapsto r_i = \sum_{k=1}^{N} d_{ik}^{-3}$.
- 2. Create a rate-corrected distance matrix (**M**) in which the elements are defined by $\mapsto M_{ij} = d_{ij} (r_i + r_j)/(N-2)^4$.
- 3. Define a new node u whose three branches join nodes i, j and the rest of tree. Define the lengths of the tree branches from u to i and $j \mapsto v_{iu} = d_{ij}/2 + ((r_i r_j)/[2(N-2)]; v_{ju} = d_{ij} v_{iu}$
- 4. Define the distance from u to each other terminal node (for all $k \neq i$ or $j \mapsto d_{ku} = (d_{ik} + d_{jk} d_{ij})/2$
- 5. Remove distances to nodes i and j from the matrix, decrease N by 1
- 6. If more than 2 nodes remain, go back to step 1. Otherwise, the tree is fully defined except for the length of the branch joining the two remaining nodes $(i \text{ and } j) \mapsto v_{ij} = d_{ij}$

³N is the number of terminal nodes

⁴Only the values i and j for which M_{ij} is minimum need to be recorded, saving the entire matrix is unnecessary



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The main virtue of neighbor-joining is its efficiency. It can be used on very large data sets for which other phylogenetic analysis are computationally prohibitive.

	Bsu	Bst	Lvi	Amo	Mlu	R	R/3
Bsu	100 <u>1-1</u> 000	0.1715	0.2147	0.3091	0.2326	0.9279	0.309
Bst	-0.4766	100 A	0.2991	0.3399	0.2058	1.0163	0.338
Lvi	-0.4905	-0.4356		0.2795	0.3943	1,1876	0.395
Amo	-0.4527	-0.4514	-0.5689		0.4289	1.3574	0.452
Mlu	-0.4972	-0.5535	-0.4221	-0.4441	_	1.2616	0.420

Lvi to node 1 distance = 0.2795/2 + (0.3959 - 0.4525)/2 = 0.1114Amo to node 1 distance = 0.2795 - 0.1114 = 0.1681

	Bsu	Bst	Mlu	Node 1	R	R/2
Bsu		0.1715	0.2326	0.1222	0.5263	0.2631
Bst	-0.3701		0.2058	0.1798	0.5571	0.2785
Mlu	-0.3856	-0.4278		0.2719	0.7103	0.3551
Node 1	-0.4278	-0.3856	-0.3701		0.5739	0.2869

Bsu to node 2 distance = 0.1222/2 + (0.2631 – 0.2869)/2 = 0.0492 node 1 to node 2 distance = 0.1222 – 0.0492 = 0.0730

	Bst	Mlu	Node 2	R	R/1
Bst		0.2058	0.1146	0.3204	0.3204
Mlu	-0.5116		0.1912	0.3970	0.3970
Node 2	-0.5116	-0.5116	_	0.3058	0.3058

Bst to node 3 distance = 0.1146/2 + (0.3204 - 0.3058)/2 = 0.0646node 2 to node 3 distance = 0.1146 - 0.0646 = 0.0500

	Mlu	Node 3
Mlu		0.1412
Node 3		

Mlu to node 3 distance = 0.1412



Unlike the UPGMA, NJ does not assume that all lineages evolve at the same rate and produces an unrooted tree.



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6.2. Pros & Cons of Distance Methods

• Pros:

- They are very fast,
- There are a lot of models to correct for multiple,
- LRT may be used to search for the best model.
- Cons:
 - Information about evolution of particular characters is lost



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7. Maximum Parsimony

Most biologists are familiar with the usual notion of **parsimony** in science, which essentially maintains that simpler hypotheses are prefereable to more complicated ones and that *ad hoc* hypotheses should be avoided whenever possible. The principle of *maximum parsimony* (MP) searches for a tree that requires **the smallest number of evolutionary changes** to explain differences observed among OTUs.

In general, parsimony methods operate by selecting trees that minimize the total tree length: the number of evolutionary steps (transformation of one character state to another) require to explain a given set of data.

In mathematical terms: from the set of possible trees, find all trees τ such that $L_{(\tau)}$ is **minimal**

$$L_{(\tau)} = \sum_{k=1}^{B} \sum_{j=1}^{N} w_j.diff(x_{k'j}, x_{k''j})$$

Where $L_{(\tau)}$ is the length of the tree, *B* is the number of branches, *N* is the number of characters, k' and k'' are the two nodes incident to each branch k, $x_{k'j}$ and $x_{k''j}$ represent either element of the input data matrix or optimal character-state assignments made to internal nodes, and diff(y, z) is a function specifying the cost of a transformation from state y to state z along any branch. The coefficient w_j assigns a weight to each character. Note also that diff(y, z) needs not to be equal diff(z, y).⁵

⁵For methods that yield unrooted trees diff(y, z) = diff(z, y).

Determining the length of the tree is computed by algorithmic methods [4, 15]. However, we will show how to calculate the length of a particular tree topology $((W,Y),(X,Z))^6$ for a specific site of a sequence, using Fitch (A) and transversion parsimony (B)⁷:

 Seq. WACAGGAT...
 (A)
 (B)

 Seq. XACACGCT...
 $\begin{bmatrix} 0 & 1 & 1 & 1 \\ 1 & 0 & 1 & 1 \\ 1 & 1 & 0 & 1 \\ 1 & 1 & 0 & 1 \\ 1 & 1 & 1 & 0 \end{bmatrix}$ $tv4 = \begin{bmatrix} 0 & 4 & 1 & 4 \\ 4 & 0 & 4 & 1 \\ 1 & 4 & 0 & 4 \\ 4 & 1 & 4 & 0 \end{bmatrix}$

- With equal costs, the minimum is 2 steps, achieved by 3 ways (internal nodes "A-C", "C-C", "G-C"),
- The alternative trees ((W,X),(Y,Z)) and ((W,Z),(Y,X)) also have 2 steps,
- Therefore, the character is said to be **parsimony-uninformative**,⁸
- With 4:1 ts:tv weighting scheme, the minimum length is 5 steps, achived by two reconstructions (internal nodes "A-C" and "G-C"),
- By evaluating the alternative topologies finds a minimum of 8 steps,



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⁶Newick format

⁷Matrix character states: A,C,G,T

 $^{^{8}\}mathrm{A}$ site is informative, only it favors one tree over the others



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• Therefore, under unequal costs, the character **becomes informative**. The use of unequal costs may provide more information for phylogenetic reconstruction,



tv4: 1+0+0+4+4=9





equal: 1+0+1+1+1=4 equal: 1+0+1+1+1=3 tv4: 1+0+1+4+4=10 tv4: 1+0+4+1+1=7



equal: 1+1+1+1+1=5

tv4: 4+4+4+4=20



equal: 1+1+0+0+0=2

tv4: 4+4+0+0+0=8

equal: 0+1+1+0+0=2

tv4: 0+1+4+0+0=5





equal: 1+1+1+1+1=5

equal: 1+1+1+1+1=5 tv4: 4+4+4+4=20 tv4: 4+4+1+1+1=11







equal: 0+1+0+1+1=3

tv4: 0+1+0+4+4=9



equal: 0+1+1+1+1=4tv4: 0+1+1+4+4=10





equal: 1+1+1+1+1=5 tv4: 4+4+4+4=20 tv4: 4+4+1+0+0=9

equal: 1+1+1+0+0=3

equal: 1+1+1+1+1=5 tv4: 4+4+4+4=20 tv4 4+4+0+1+1=10



equal: 0+1+1+1+1=4

tv4: 0+1+4+1+1=7



7.1. Pros & Cons of MP

- Pros:
 - Does not depend on an explicit model of evolution,
 - At least gives both, a tree and the associated hypotheses of character evolution,
 - If homoplasy is rare, gives reliable results,

• Cons:

- May give misleading results if homplasy is common (*Long branch attraction effect*)
- Underestimate branch lengths
- Parsimony is often justified by phylosophical, instead statistical grounds.



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8. Searching Trees

8.1. How many trees are there?

The obvious method for searching the most parsimonious tree is to consider all possible trees, one after another, and evaluate them. We will see that this procedure becomes impossible for more than a few number of taxa (\sim 11). Felsenstein [2] deduced that:

$$B(T) = \prod_{i=3}^{T} (2i-5)$$

An unrooted, fully resolved tree has:

- T terminal nodes, T-2 internal nodes,
- 2T-3 branches; T-3 interior and T peripheral,
- B(T) alternative topologies,
- Adding a **root**, adds one more **internal node** and one more **internal branch**,
- Since the root can be placed along any 2T 3 branches, the number of possible rooted trees becomes,

$$B(T) = (2T - 3) \prod_{i=3}^{T} (2i - 5)$$



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OTUs	Rooted trees	Unrooted trees
2	1	1
3	3	1
4	15	3
5	105	15
6	954	105
7	10,395	954
8	$135,\!135$	10,395
9	2,027,025	$135,\!135$
10	$34,\!459,\!425$	$2,\!027,\!025$
11	$> 654 \mathrm{x} 10^{6}$	$> 34 \mathrm{x} 10^{6}$
15	$> 213 \text{x} 10^{12}$	$> 7 \mathrm{x} 10^{12}$
20	$> 8 \times 10^{21}$	$> 2 \times 10^{20}$
50	$> 6 \times 10^{81}$	$> 2 \mathrm{x} 10^{76}$

The observable universe has about 8.8×10^{77} atoms

There is not memory neither time to evaluate all the trees!!

For 11 or fewer taxa, a brute-force **exhaustive search** is feasible!! For more than 11 taxa an **heuristic search** is the best solution!!



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8.2. Exhaustive search methods

- Every possible tree is examined; the shortest tree will always be found,
- Taxon addition sequence is important only in that the algorithm needs to remember where it is,
- Search will also generate **a list** of the lenths of all possible trees, which can be plotted as an histogram,





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8.3. Heuristic search methods

When a data set is **too large to permit the use of exact methods**, optimal trees must be sought via heuristic approaches that **sacrifice the guarantee of optimality in favor of reduced computing time**

Two kind of algorithms can be used:

- 1. Greedy Algorithms
- 2. Branch Swapping Algorithms



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8.3.1. Greedy Algorithms



Strategies of this sort are often called *the greedy algorithm* because they seize the first improvement that they see. Two major algorithms exist:

- Stepwise Addition,
- Star Decomposition⁹

Both algoritms are prone to entrapment in local optima



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⁹The most common star decomposition method is the NJ algorithm

Stepwise Addition

- Use addition sequence similar to that for an exhaustive search, but at each addition, determines the shortest tree, and add the next taxon to that tree.
- Addition sequence will affect the tree topology that is found!



A greedy stepwise-addition search applied to the example in Figure 7.2. The best four-taxon tree is determined by evaluating the lengths of the three trees obtained by joining Jacon D to Iree 1 containing only the first three taxa. Taxa E and F are then connected to the fire and seven possible locations, respectively, on Irees 4 and 9, with only the shortest trees found during each tap being used for the next step. In this cample, the 233-dep tree obtained is not a global optimum (see Figure 7.2.), Circled numbers indicate the order in which phylogenetic trees an ocaluated in the stepwise-addition search.



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Star Decomposition

- Start with all taxa in an unresolved (star) tree,
- Form pairs of taxa, and determine length of tree with paired taxa.



Figure 25 Heuristic tree selection using star decomposition method. At each step, the optimality criterion is evaluated for each possible joining of a pair of lineages leading away from the central node. The best tree found during each step becomes the starting point for the next step.



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8.3.2. Branch Swapping Algorithms

It may be possible to improve the *greedy* solutions by performing sets of predefined rearrangements, or branch swappings. Examples of branch swapping algorithms are:

- NNI Nearest Neighbor Interchange,
- SPR Subtree Pruning and Regrafting,
- TBR Tree Bisection and Reconnection.



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9. Tree Confidence

9.1. Non-parametric bootstrapping

- For many simple distributions there are simple equations for calculating confidence intervals around an estimate (e.g., std error of the mean)
- Trees, however are rather complicated structures, and it is extremely difficult to develop equations for confidence intervals around a phylogeny.
- One way to measure the confidence on a phylogenetic tree is by means of the **bootstrap** non-parametric method of resampling the same sample many times.





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- Each sample from the original sample is a **pseudoreplicate**. By generation many hundred or thousand pseudoreplicates, a *majority consensus rule tree* can be obtained.
- High bootstrap values > 90% is indicative of strong **phylogenetic signal**.
- Bootstrap can be viewed as a way of exploring the robustness of phylogenetic inferences to perturbations
- **Jackkniffe** is another non-parametric resampling method that differentiates from bootstrap in the way of sampling. Some proportion of the characters are randomly selected and deleted (withouth replacement).
- Another technique used exclusively for parsimony is by means of **Decay** index or **Bremmer support**. This is the length difference between the shortest tree including the group and the shortest tree excluding the group (The extra-steps required to overturn a group.¹⁰
- $\bullet~\mathbf{DI}\ \&\ \mathbf{BPs}$ generally correlates!!



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 $^{^{10}}$ See [19] for a practical example using PAUP*[17]

10. PC Lab

10.1. Download Programs

- PHYLIP 3.6 http://evolution.genetics.washington.edu/phylip.html
- MEGA 3.0 http://www.megasoftware.net
- TREE-PUZZLE http://www.tree-puzzle.de/
- MODELTEST http://darwin.uvigo.es/
- MrBayes http://morphbank.ebc.uu.se/mrbayes/download.php
- TreeView http://taxonomy.zoology.gla.ac.uk/rod/treeview.html



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11. Phylogenetic Links

- Software:
 - The Felsenstein node http://evolution.genetics.washington.edu/phylip/software. html
 - The R. Page Lab. http://taxonomy.zoology.gla.ac.uk/software/software.html
- Courses:
 - Molecular Systematics and Evolution of Microorganisms. http://www.dbbm. fiocruz.br/james/index.html
 - Workshop on Molecular Evolution http://workshop.molecularevolution.org/
 - P. Lewis MCB/EEB Course http://www.eeb.uconn.edu/Courses/EEB372/
- Tools:
 - Clustalw at EBI http://www.ebi.ac.uk/clustalw/
 - Phylip Web http://cbrmain.cbr.nrc.ca:8080/cbr/jsp/ServicePage_e.jsp?id=38
 - Phylip Dochttp://www.hgmp.mrc.ac.uk/Registered/Help/phylip/phylip.html



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12. Credits

This presentation is based on:¹¹

- Major Book or Chapters References:
 - Swofford, D. L. et al. 1996. Phylogenetic inference [18].
 - Harvey, P. H. et al. 1996. New Uses for New Phylogenies [7].
 - Li, W. S. 1997 . Molecular Evolution [9].
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 - Salemi, M. & Vandamme, A. (ed.) 2003. The phylogenetic handbook [14].
 - $-\,$ Felsenstein, J. 2004. Inferring phylogenies [3] .
- On Line Phylogenetic Resources:
 - http://www.dbbm.fiocruz.br/james/index.html.Molecular Systematics and Evolution of Microorganisms. The Natural History Museum, London and Instituto Oswaldo Cruz, FIOCRUZ.
 - Peter Foster's "The Idiot's Guide to the Zen of Likelihood in a Nutshell in Seven Days for Dummies" at http://www.bioinf.org/molsys/data/idiots.pdf
- Slides Production:
 - Latex and pdfscreen package.



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 $^{^{11}\}mathrm{HJD}$ take responsibility for innacuracies of this presentation.

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