#### Phylogenetic reconstruction: criteria

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Need for an algorithm to find/construct the tree

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- Need for a criterion/score
  - Maximum Parsimony
  - Minimum Evolution or least squares (distance methods)
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  - Posterior Probability P(M|D)
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- Posterior Probability P(M|D)
  - Need for an algorithm to find/construct the tree
    - e.g.: try several topologies, (choose some branch lengths,) score the topologies, choose the one that has the best score

### Plan: Criteria for evaluating phylogenies

- Criteria for evaluating phylogenetic trees:
  - Parsimony
  - Distance methods
  - Maximum Likelihood
    - Using a Monte Carlo simulation
    - Using Felsenstein's pruning algorithm
  - Posterior probability (Bayesian approach)
- Conventions:
  - We're dealing with aligned sequence data
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#### Parsimony

 "The principle that the most acceptable explanation of an occurrence, phenomenon, or event is the simplest, involving the fewest entities, assumptions, or changes. In phylogenetics, for example, the preferred tree showing evolutionary relationships between species, molecules, or other entities is the one that requires the least amount of evolutionary change, that is, maximum parsimony."

Criteria: Maximum Parsimony

# Maximum parsimony

- Has been advocated strongly by some against model-based approaches: many controversies (see "The Troubled Growth of Statistical Phylogenetics", Felsenstein 2001)
- Edwards and Cavalli-Sforza (1963): the preferred evolutionary tree involves "the minimum net amount of evolution" = *Maximum parsimony tree*
- → For sequence data: find the phylogeny that involves the minimum number of substitutions
  - → We need a way to count the minimum number of substitutions on a phylogeny = compute the parsimony score of a phylogenetic tree

### Computing the parsimony score

Sp1 ATGCGCT...

Sp1 Sp3

Sp2 AGTCGCA...

Sp3 AGGTGCA...

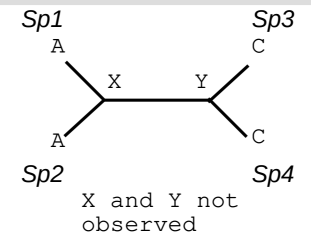
Sp4 ATGCCCT...

Sp4 Sp4

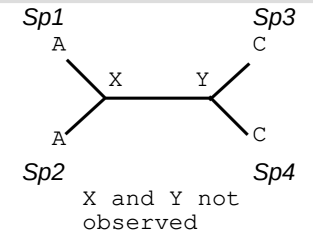
Parsimony score of a tree given an alignment: sum of the parsimony scores for each site

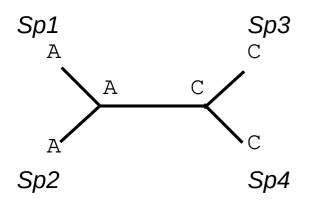
→ We assume that all sites are independent

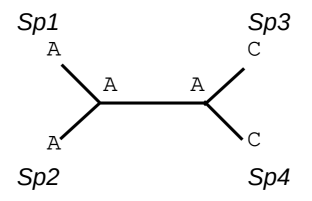
Species tree S Site i

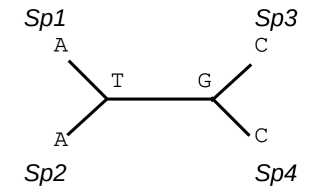


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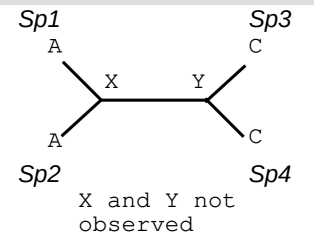


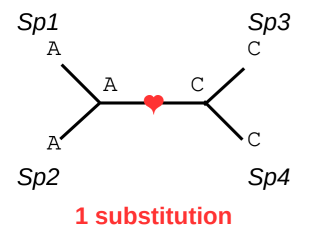


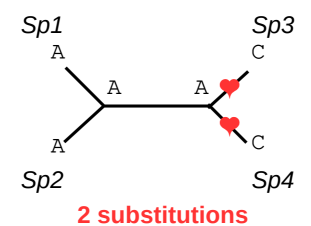


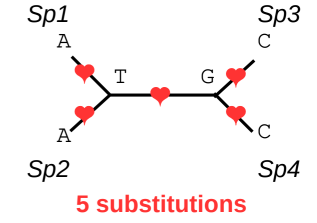


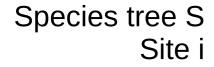
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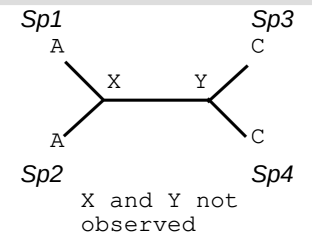


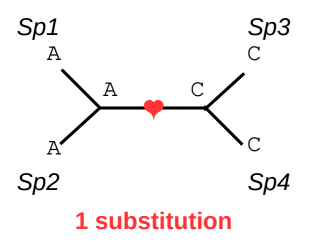


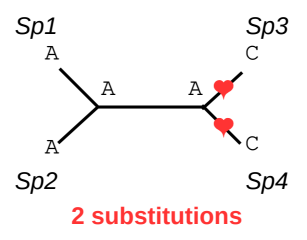


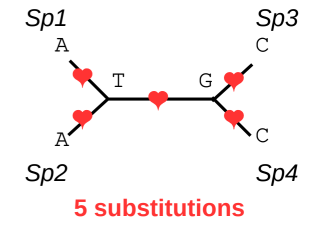












Parsimony score of species tree S for site i: 1

# Computing the parsimony score of a tree at one site

- Naive brute force approach: test all possible assignments on internal nodes
  - N internal nodes, 4 possibilities {A,C,G,T} per node
  - → 4<sup>N</sup> possibilities to try

- Fitch's algorithm (1971):
  - Arbitrarily root the tree
  - Compute, from the tips up, two elements per node:
    - P: The score of the underlying subtree
    - X: The set of states possible at that node, given the score P
  - Complexity: O(4N)

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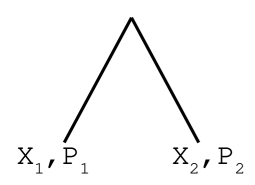
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For 2 internal nodes: 4\*2 =8

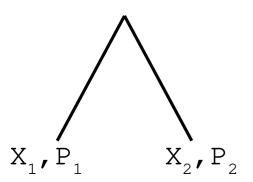
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Climbing up the tree: computing P and X for a node given its children

 $\underline{\mathbf{1}}^{\operatorname{st}}$  case:  $X_1 \cap X_2$  not empty

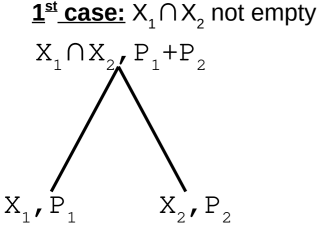


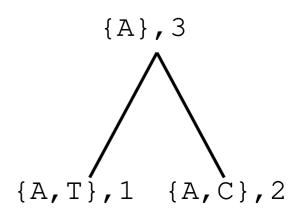
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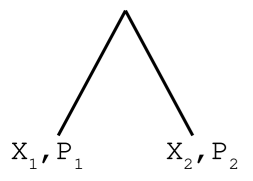
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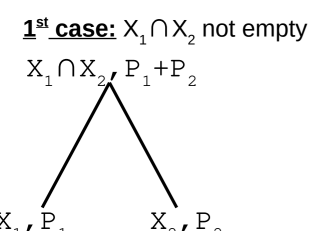


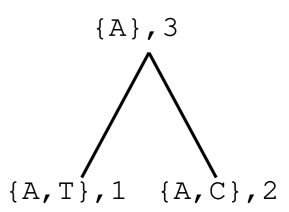
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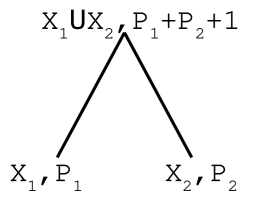
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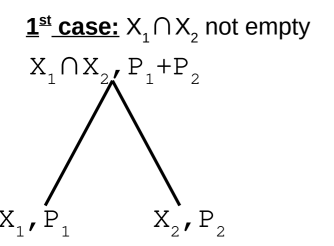


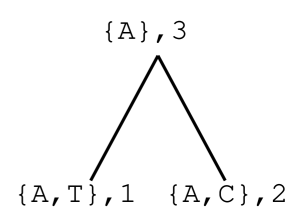
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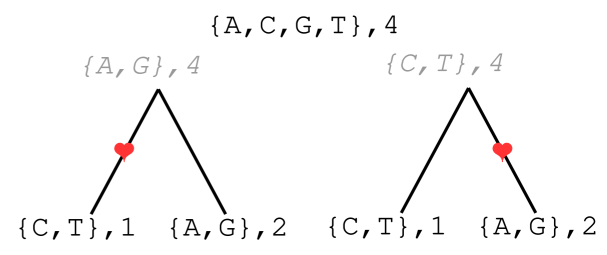
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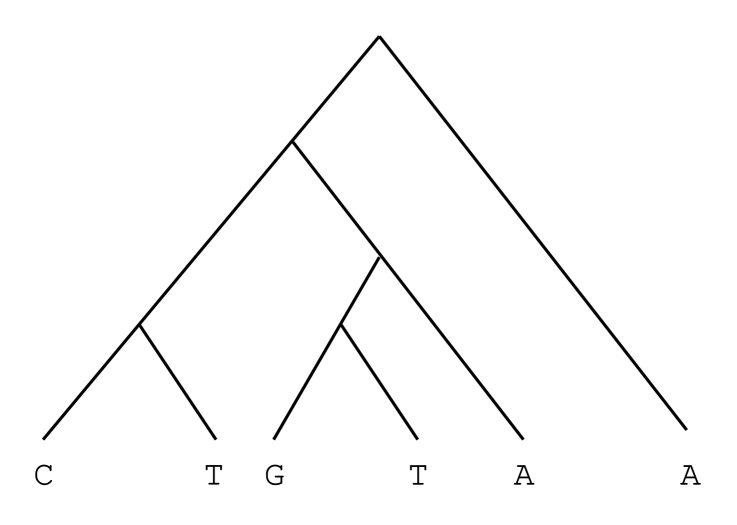
Climbing up the tree: computing P and X for a node given its children

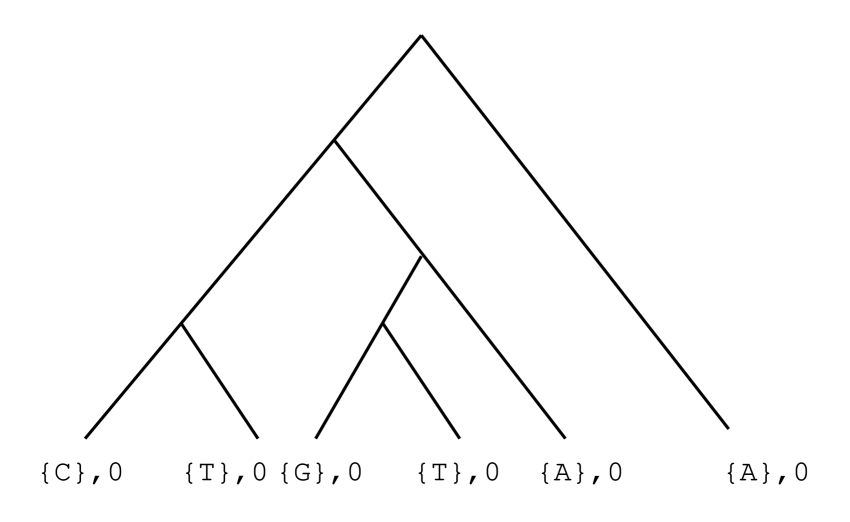


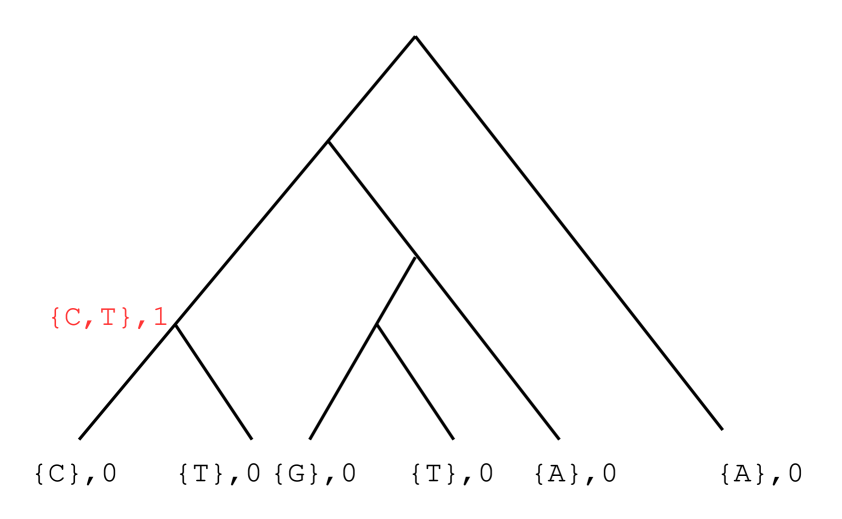


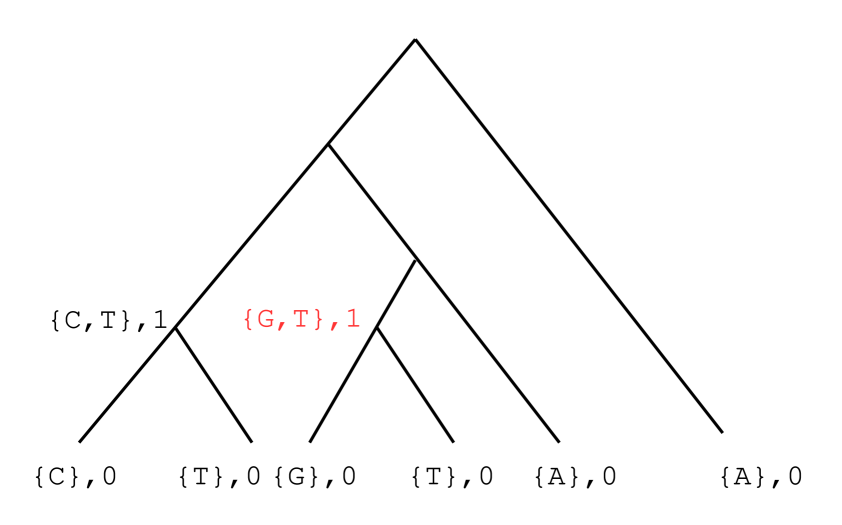
 $\begin{array}{ccc}
\mathbf{2}^{\operatorname{nd}} & \mathbf{case:} & \mathbf{X}_{1} \cap \mathbf{X}_{2} & \text{empty} \\
\mathbf{X}_{1} \mathbf{U} \mathbf{X}_{2} \mathbf{P}_{1} + \mathbf{P}_{2} + \mathbf{1} \\
\mathbf{X}_{1} \mathbf{P}_{1} \mathbf{X}_{2} \mathbf{P}_{2}
\end{array}$ 

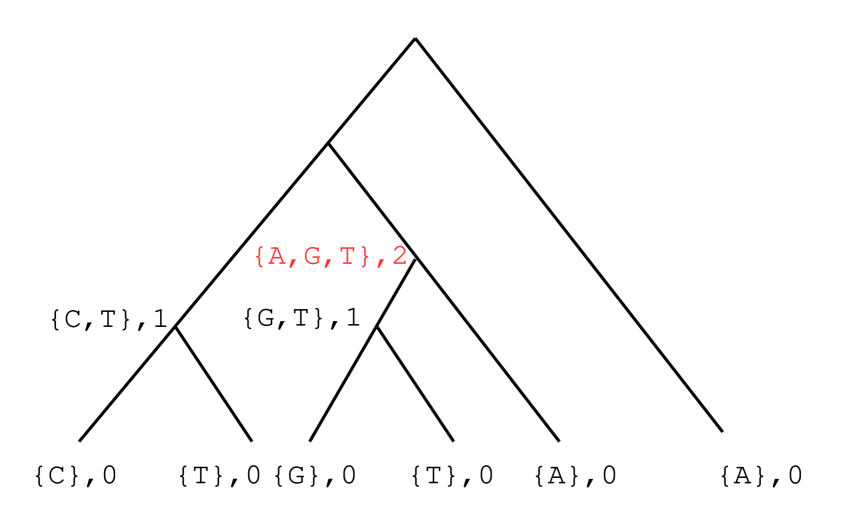


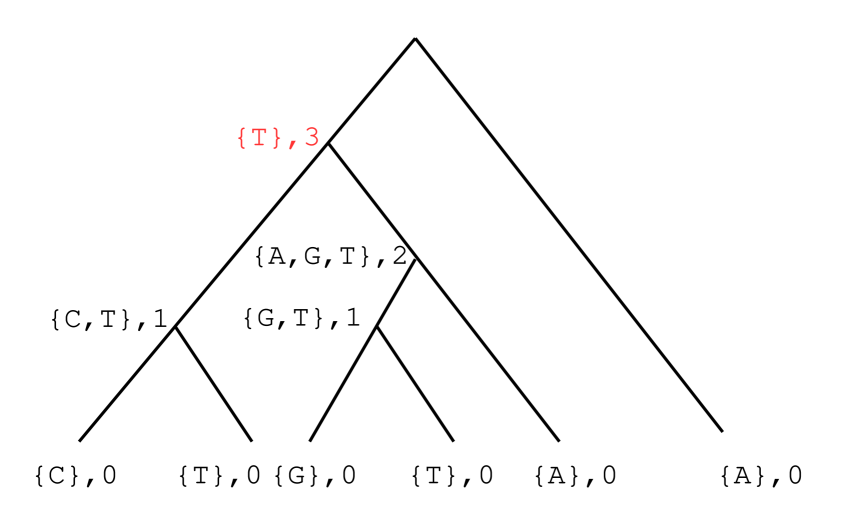


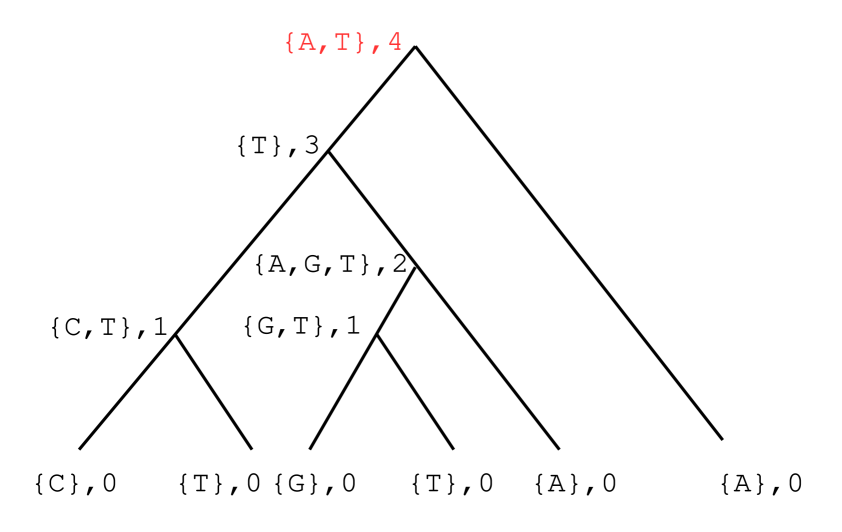


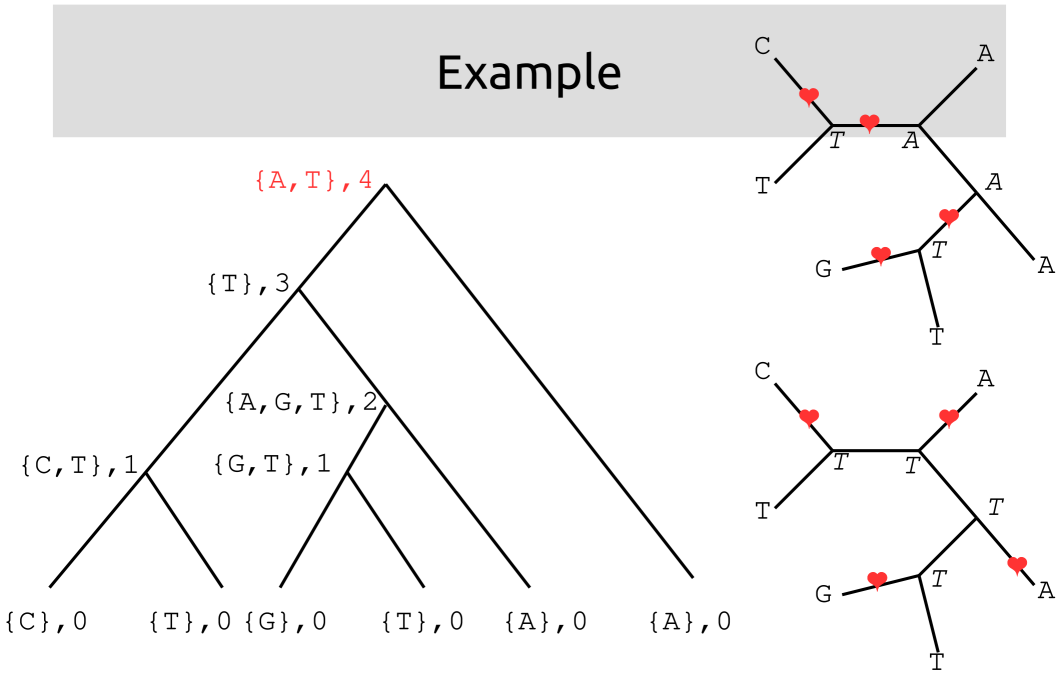












Several possible scenarios

#### Conclusion on maximum parsimony

- Involves looking for the tree with the least amount of change
- Computing the parsimony score of a tree can be done with Fitch's algorithm
- Searching through topology space for the most parsimonious tree can be done with algorithms similar to those used for Maximum Likelihood reconstruction (see Alexis's talk)

### Plan: Criteria for evaluating phylogenies

- Criteria for evaluating phylogenetic trees:
  - Parsimony
  - Distance methods
  - Maximum Likelihood
    - Using a Monte Carlo simulation
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#### Distance methods

- Distance-based approaches:
  - least squares methods,
  - Minimum evolution method,
  - Neighbor Joining.

```
Sp1 ATGCGCT...
Sp2 AGTCGCA...
Sp3 AGGTGCA...
Sp4 ATGCCCT...
```

```
Sp1 ATGCGCT...

Sp2 AGTCGCA...

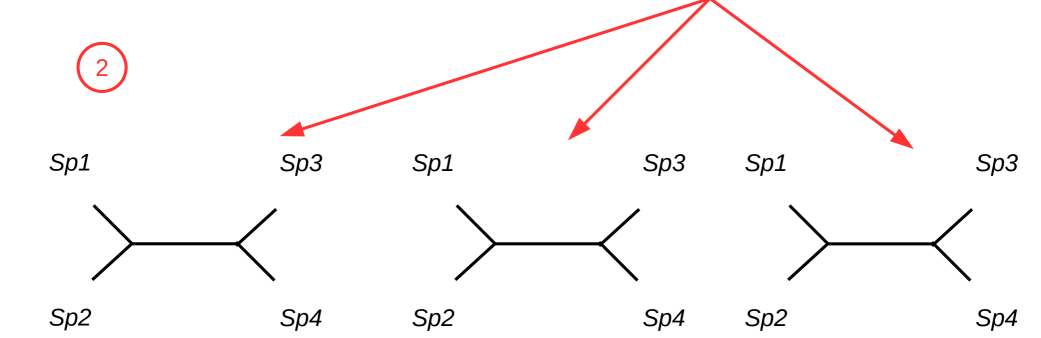
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	Sp1	Sp2	Sp3	Sp4
Sp1	0	0.1	0.2	0.15
Sp2	0.1	0	0.3	0.01
Sp3	0.2	0.3	0	0.6
Sp4	0.15	0.01	0.6	0

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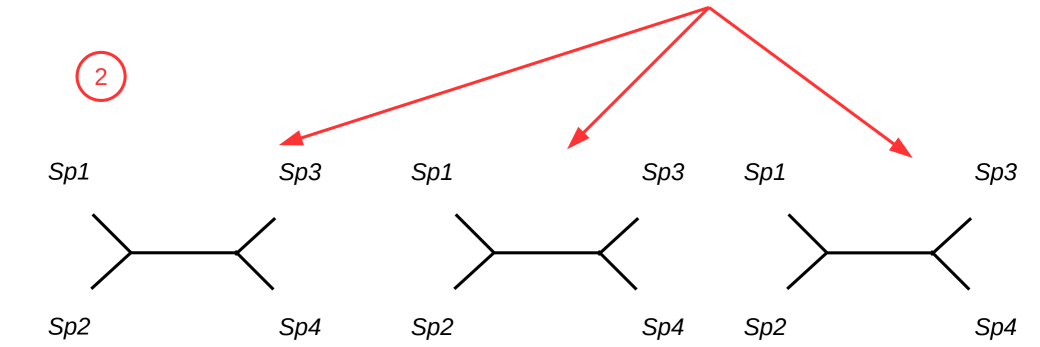
#### Uses a distance matrix:

Sp1 ATGCGCT...

Sp2 AGTCGCA...

How to compute the distance matrix?

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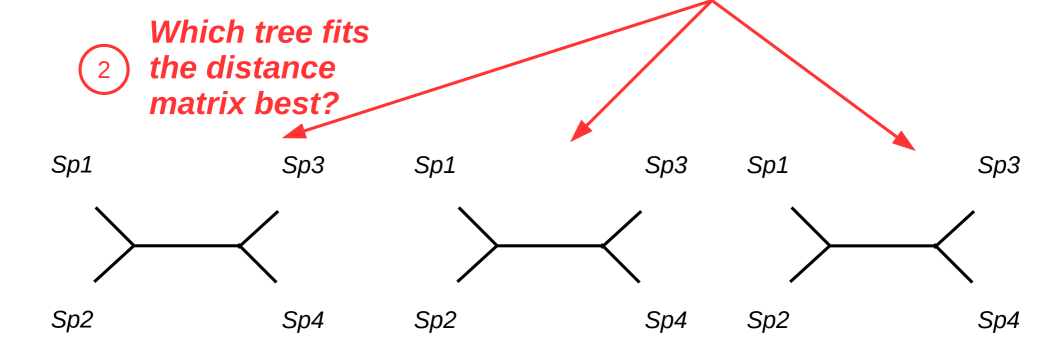


### Minimum Evolution or least squares: distance methods

Sp1	ATGCGCT	(1)
Sp2	AGTCGCA	How to compute
Cn 2	7 CCTCC7	How to compute

SPS	AGGIGCA	the distance
Sp4	ATGCCCT	matrix?

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Simply count differences (observed divergence)

Criteria: Minimum Evolution or least squares

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Simply count differences (observed divergence)

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 ATGCGCT  $Sp2$  AGTCGCA  $\longrightarrow$  d(Sp1-Sp2) = 3/7~0.43

- Use a model of sequence evolution
  - $\rightarrow$  cf. talk on models
  - -Advantages:
    - Hidden substitutions are taken into account
    - Parameters of the model of substitution can be estimated in the Maximum Likelihood framework

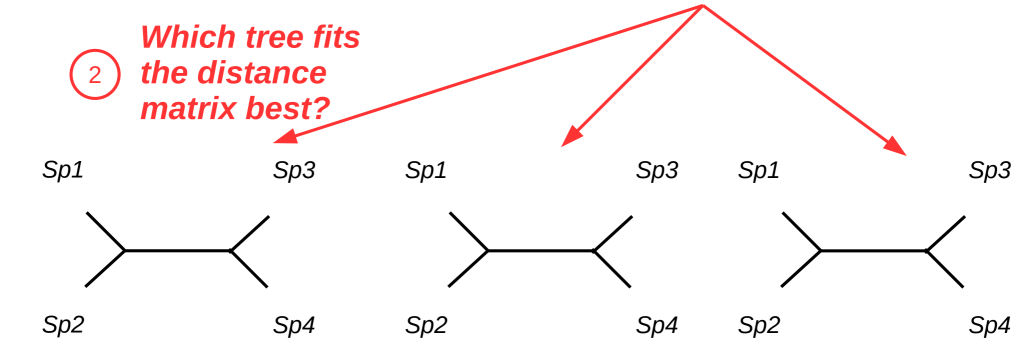
## Minimum Evolution or least squares: distance methods

### Uses a distance matrix:

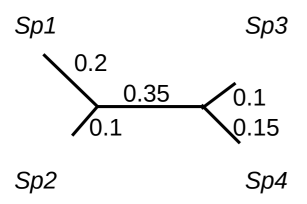
(1)	AIGCGCI	Spi
How to compute	AGTCGCA	Sp2
the distance	AGGTGCA	Sp3

Sp4 ATGCCCT... matrix?

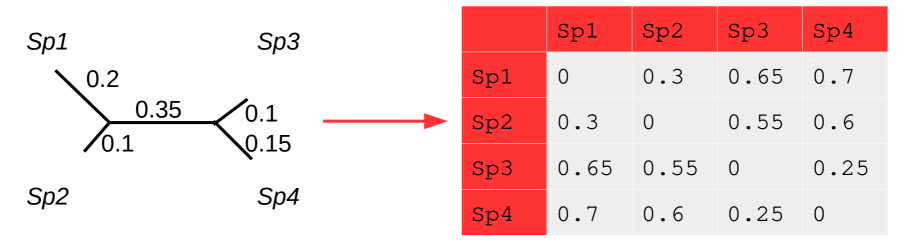
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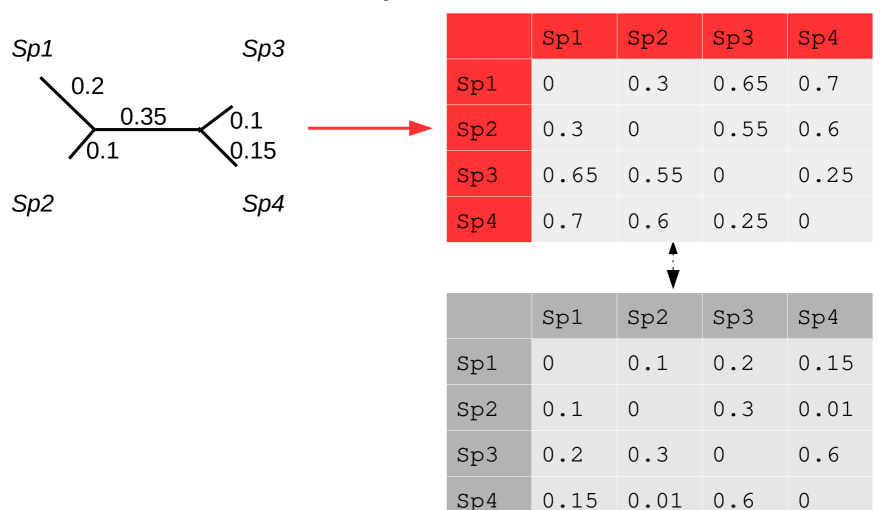
A tree implies distances between tips: compare those patristic distances to sequence-based distances



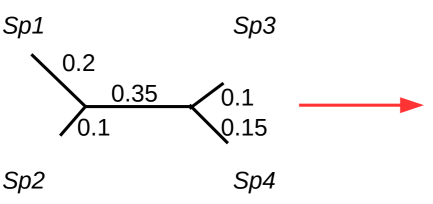
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Sp3	0.65	0.55	0	0.25
Sp4	0.7	0.6	0.25	0
		<b>A</b>		

score	=	$(0-0)^2 + (0.3-0.1)^2 + (0.65-$
0.2)2+		

With ULS: Unweighted Least Squares (other criteria have been proposed)

▼					
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Criteria: Minimum Evolution or least squares

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Using the ULS criterion, we can compute the fit between a sequence-based distance matrix and any tree (topology + branch lengths), thanks to the patristic matrix trick.

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But how can we pick branch lengths on the topology?

ULS provides a mathematical way to find the optimal branch lengths on a given topology! This involves some simple matrix algebra (solving a set of linear equations).

## Searching for the best tree using Unweighted Least Squares

- We now know how to compute the ULS score of a tree topology. It involves:
  - Matrix algebra to find the best branch lengths
  - Computing the score<sub>ULS</sub> for that tree
- Given a set of tree topologies, we can compute the "best" tree topology according to the ULS criterion: it is the one with the lowest score
- How to obtain a set of tree topologies to score is tackled later in the course (see Alexis's talk)

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## Minimum evolution criterion

- Motivation similar to parsimony
- **Hypothesis:** the true tree should be the shortest tree
- → Idea:
  - Given a matrix of pairwise distances and a set of tree topologies to evaluate
  - Match pairwise distances onto each tree topology
  - Sum the branch lengths on each tree
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- → Idea:
  - Given a matrix of pairwise distances and a set of tree topologies to evaluate
  - Match pairwise distances onto each tree topology: Use least-squares fitting!
  - Sum the branch lengths on each tree
  - Your best estimate is the tree with the smallest sum of branch lengths

**NTCCCCT** 

## Minimum Evolution or least squares: distance methods

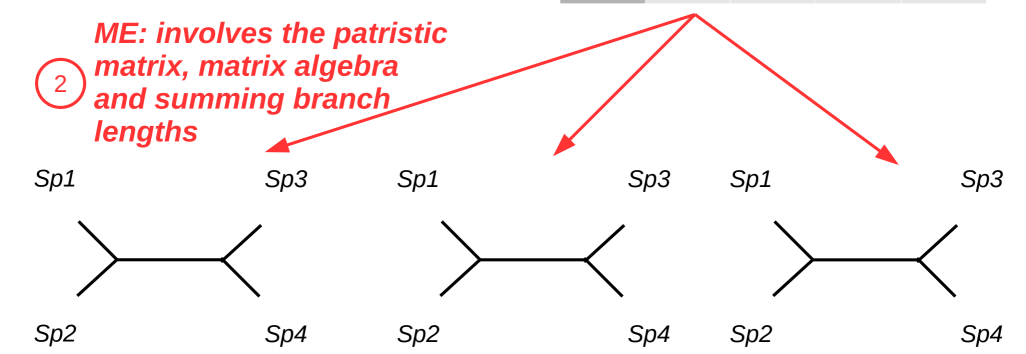
### Uses a distance matrix:

SPI	AIGCGCI	
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## Minimum evolution criterion

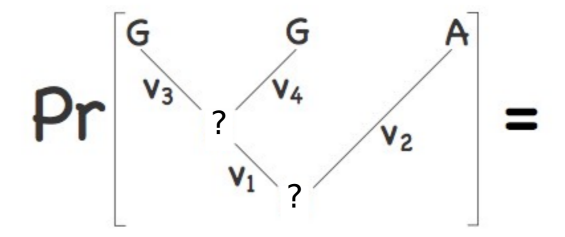
- To obtain a Minimum Evolution tree, at some point we have to use Least Squares estimation to assign branch lengths to a tree topology
  - → hybrid approach where two different criteria are mixed up
- However, Minimum evolution works pretty well in practice
- Neighbor-Joining (Saitou and Nei, 1987) is a famous heuristic algorithm for finding the Minimum Evolution tree (not seen in our course, but has been very widely used); see Gascuel and Steel, 2006 for a clear explanation

## Summary on distance methods

- Distance methods are the fastest phylogenetic methods available, notably thanks to Neighbor Joining and other algorithms to build a tree given a distance matrix (e.g. BioNJ, Weighbor, FastME...)
- Can be based on models of sequence evolution to compute pairwise distances
- Better than Maximum Parsimony when sequences are divergent, but less accurate than Maximum Likelihood or Bayesian Inference
- The main reason is that distance methods do not use the entire data matrix together, but look at it pair of sequences by pair of sequences

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$$\pi_i \times p_{ij}(v_1) \times p_{iA}(v_2) \times p_{jG}(v_3) \times p_{jG}(v_4)$$

 $\pi_i$  Stationary frequencies

 $p_{ij}(v)$  Transition probabilities

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The likelihood is a quantity that is proportional to the probability of observing/realizing the data under a fully specified model/hypothesis.

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 $Likelihood(Parameter) \propto Probability(Data|Parameter)$ 

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 $Likelihood(Parameter) \propto Probability(\mathbf{X}|Parameter)$ 

#### What is likelihood?

The likelihood is a quantity that is proportional to the probability of observing/realizing the data, X, under a fully specified model/hypothesis, $\Theta$ .

 $Likelihood(\Theta) \propto Probability(\mathbf{X}|\Theta)$ 

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$$L(\tau, \nu, \Phi) \propto f(\mathbf{X} \mid \tau, \nu, \Phi)$$

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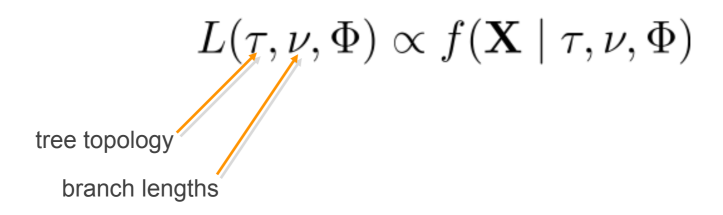
$$L(\Theta) \propto f(\mathbf{X}|\Theta)$$

$$L(\tau,\nu,\Phi) \propto f(\mathbf{X} \mid \tau,\nu,\Phi)$$
 tree topology

#### What is likelihood?

The likelihood is a quantity that is proportional to the probability of observing/realizing the data, X, under a fully specified model/hypothesis, $\Theta$ .

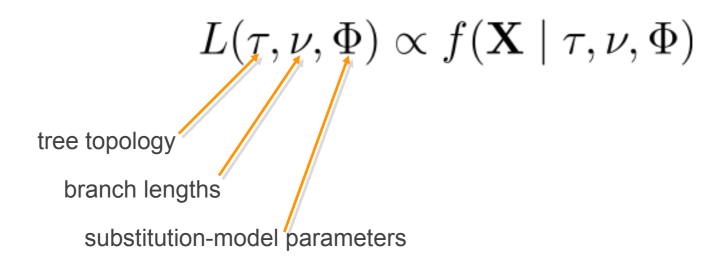
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For phylogenetic problems, the likelihood is proportional to the probability of observing the sequence alignment, **X**, under a fully specified phylogenetic model.

$$L(\tau, \nu, \Phi) \propto f(\mathbf{X} \mid \tau, \nu, \Phi)$$

A model that has a high likelihood score on a data set is a model that has a high probability of simulating this data set.

substitution-model parameters

The likelihood is a score that measures the fit of the model to the data, providing a basis for comparing different hypotheses/parameter values on the same data

OK, so what does likelihood *really* mean?

Consider a simple simulation experiment to develop our intuition

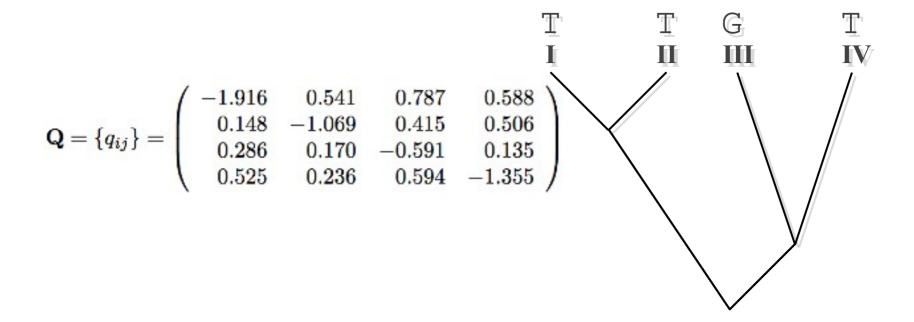
#### OK, so what does likelihood *really* mean?

Consider a simple simulation experiment to develop our intuition Imagine we have a fully specified phylogenetic model for four species:

$$\mathbf{Q} = \{q_{ij}\} = \begin{pmatrix} -1.916 & 0.541 & 0.787 & 0.588 \\ 0.148 & -1.069 & 0.415 & 0.506 \\ 0.286 & 0.170 & -0.591 & 0.135 \\ 0.525 & 0.236 & 0.594 & -1.355 \end{pmatrix}$$

#### OK, so what does likelihood *really* mean?

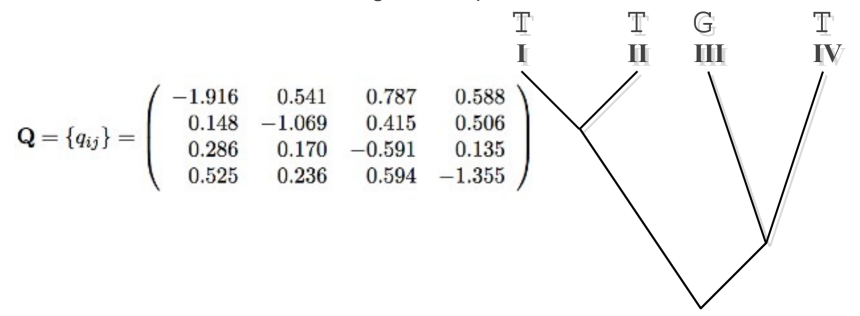
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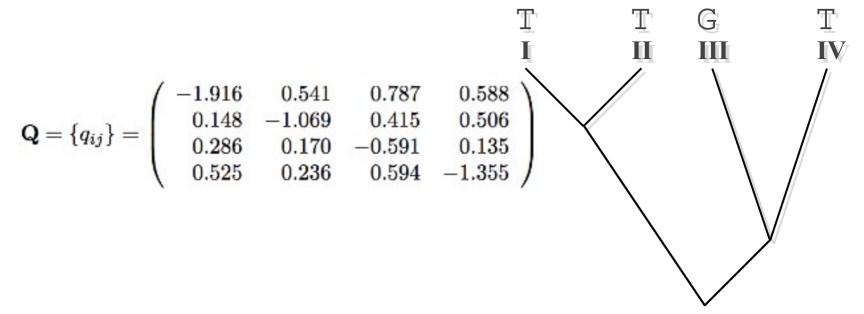
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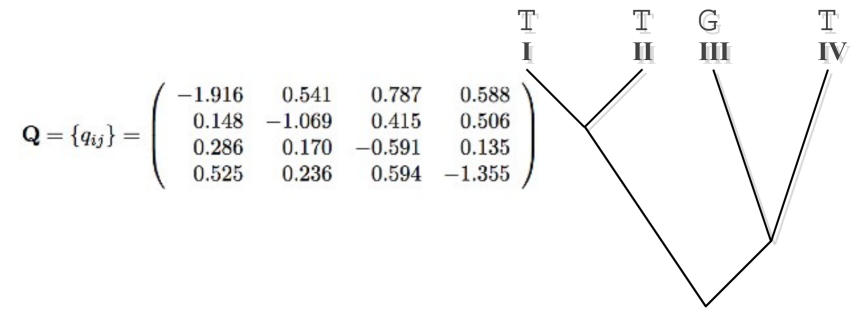
The stationary frequencies have been estimated from the rate matrix:

$$\mathbf{P}(100.0) = \begin{pmatrix} 0.138 & 0.188 & 0.495 & 0.179 \\ 0.138 & 0.188 & 0.495 & 0.179 \\ 0.138 & 0.188 & 0.495 & 0.179 \\ 0.138 & 0.188 & 0.495 & 0.179 \end{pmatrix}$$

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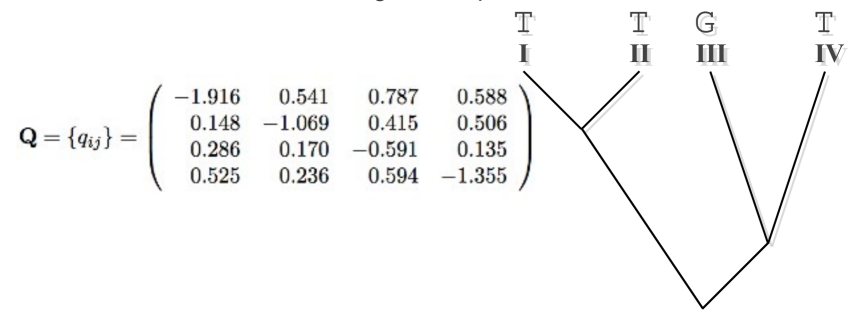
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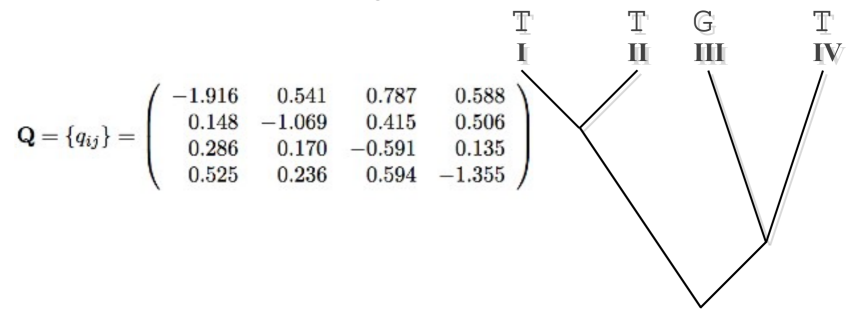
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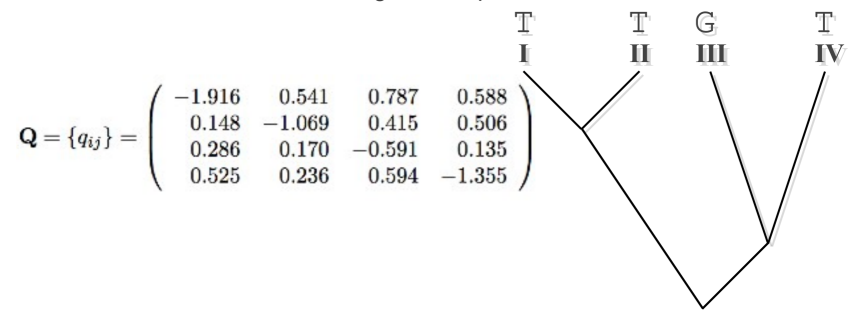
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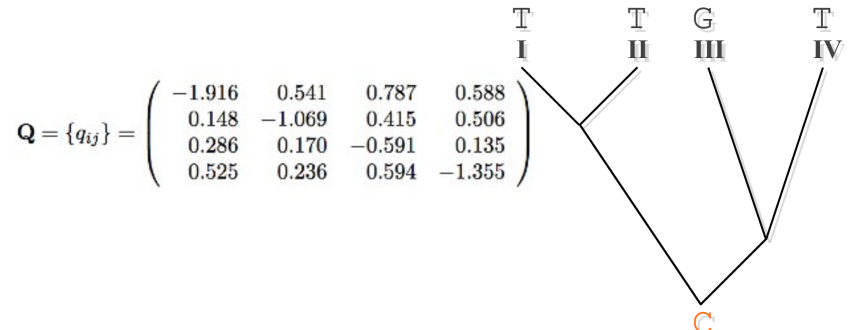
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Pick a starting state at the root using  $x \sim dnUniform(0,1)$  x=0.246

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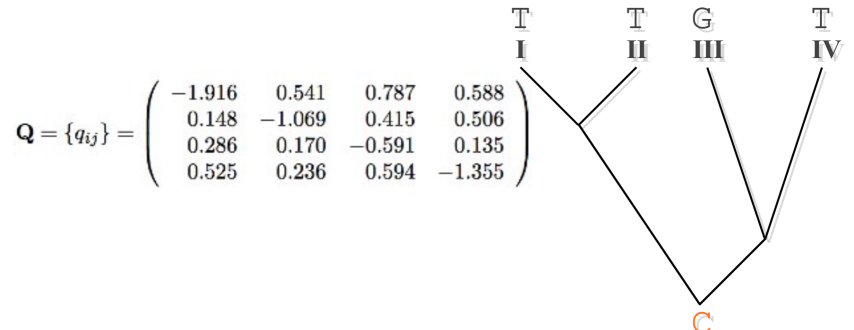
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We randomly selected state C

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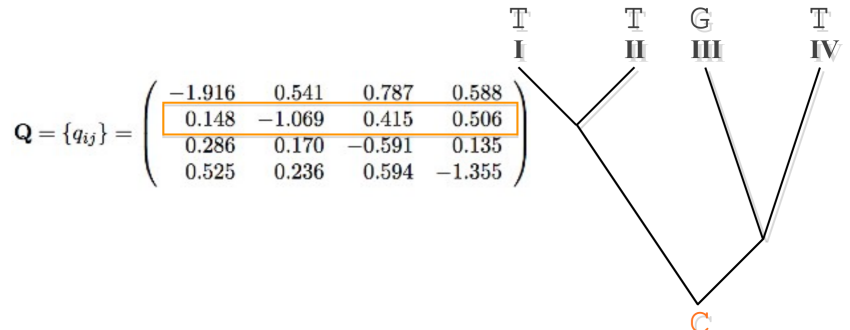
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Simulate histories along each branch in a pre-order traversal

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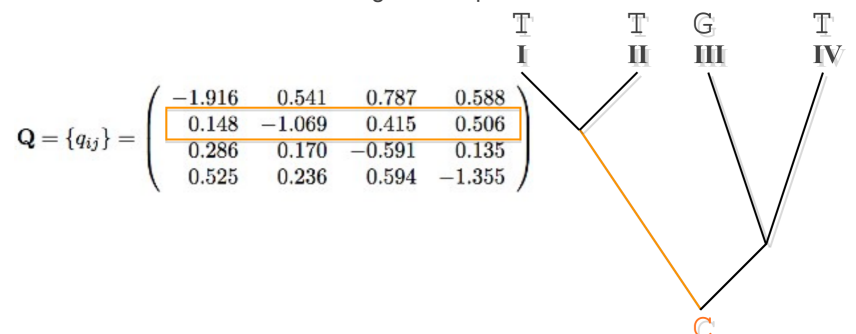
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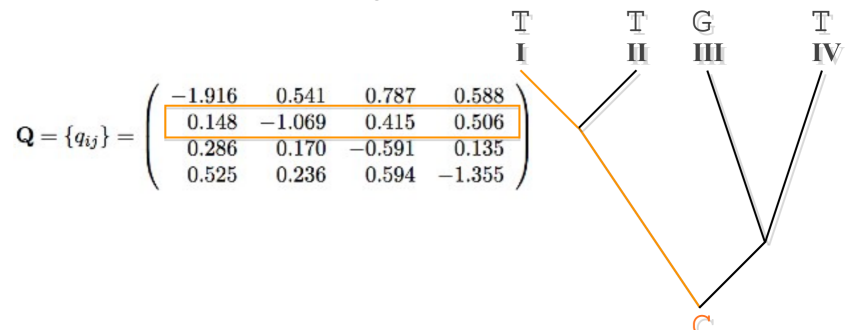
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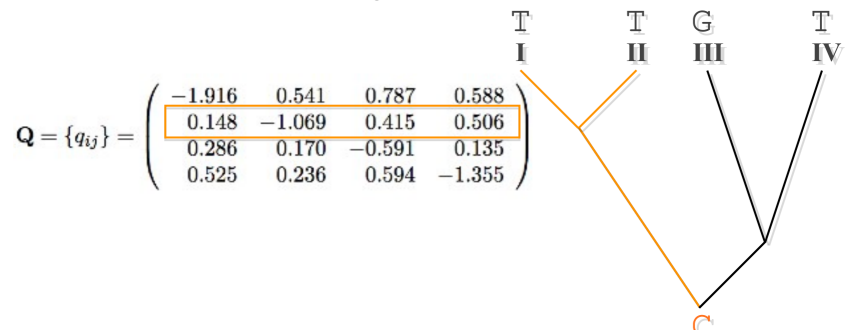
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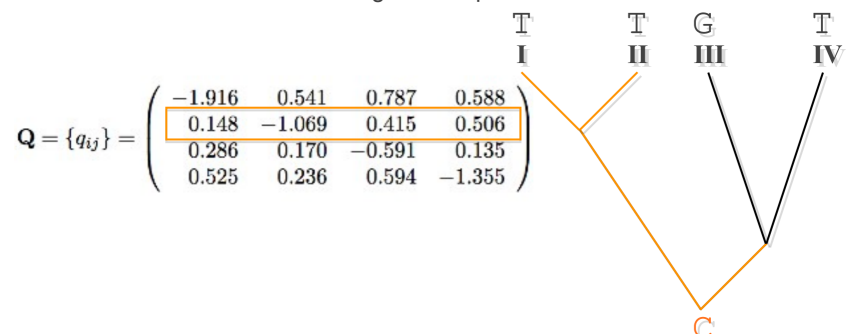
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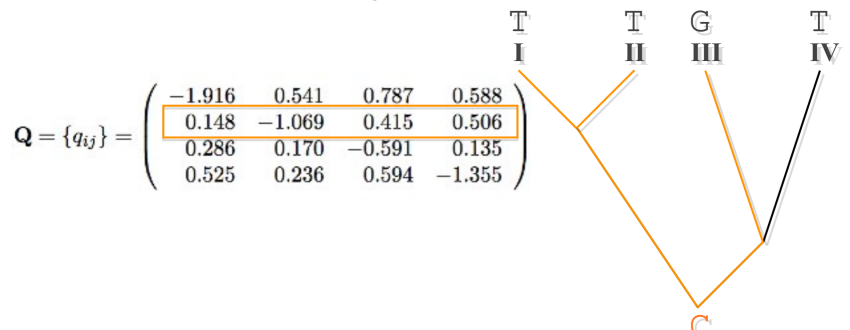
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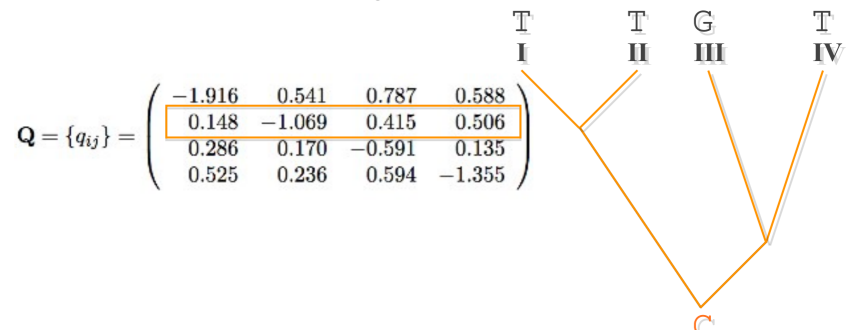
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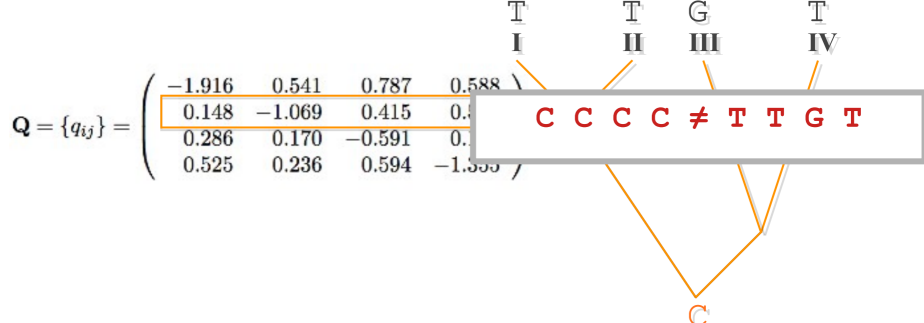
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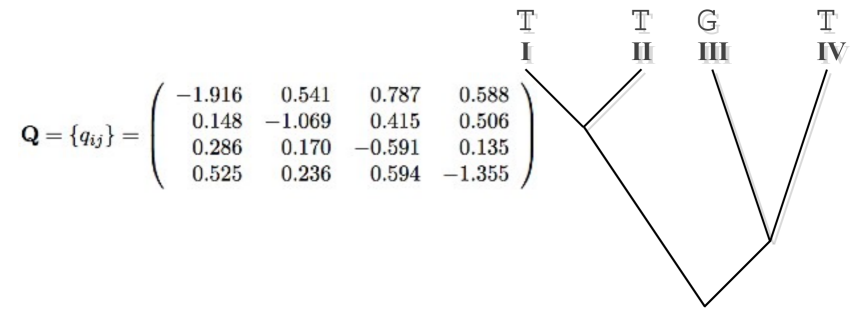
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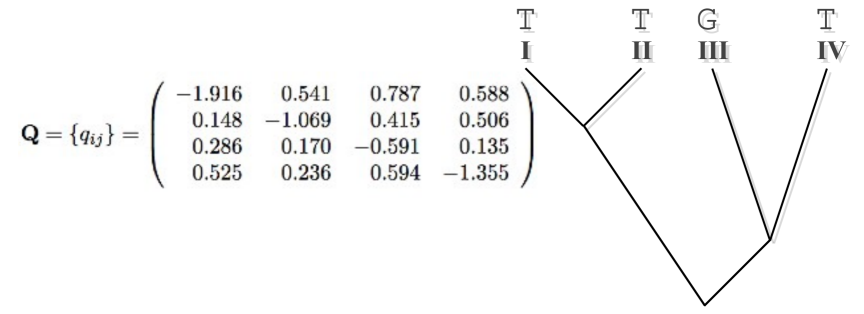


We repeat the simulation 100,000,000 times and record the frequency of outcomes that match the observed tip states.

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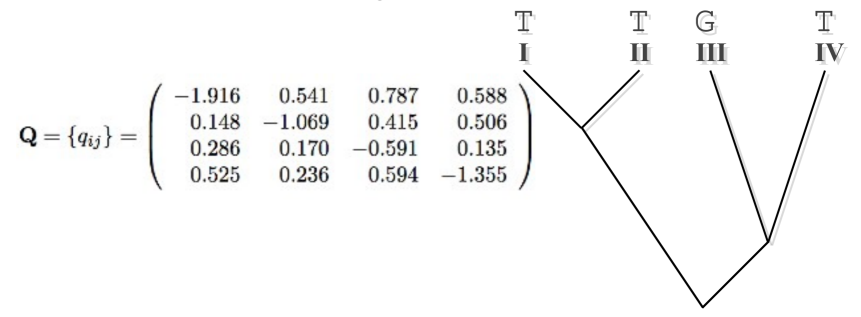
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The observed site pattern, TTGT, is one of  $4^4 = 256$  possible site patterns.

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What is the likelihood of observing the site pattern under this model?

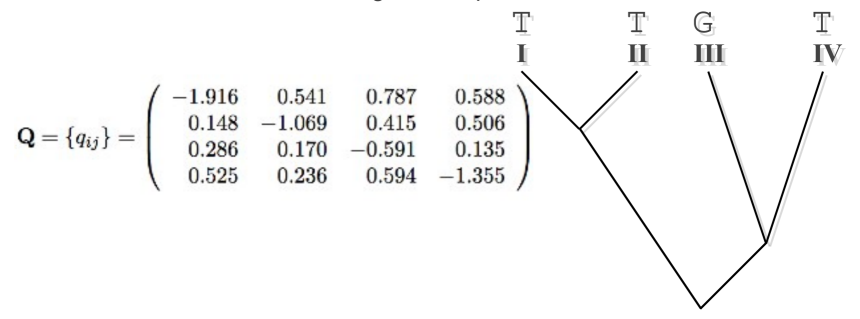


Of the 100,000,000 simulations, 850,358 realize the observed data (TTGT); the estimated probability of the observed data is therefore 0.00850358.

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Of the 100,000,000 simulations, 850,358 realize the observed data (TTGT); the estimated probability of the observed data is therefore 0.00850358.

The probability includes all of the possible histories that can give rise to the data.

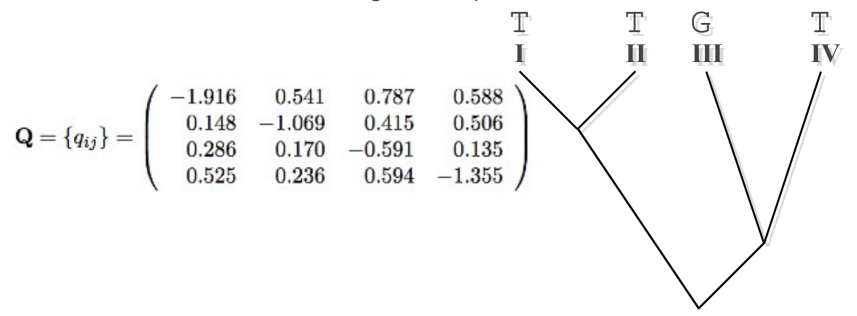
Number of Changes								
Pattern	0	1	2	3	4	5	6	7
TTGT	0.000000	0.881408	0.075358	0.037817	0.004725	0.000622	0.000062	0.000007

Number of Changes

#### OK, so what does likelihood *really* mean?

The four species have the following observed site pattern: TTGT

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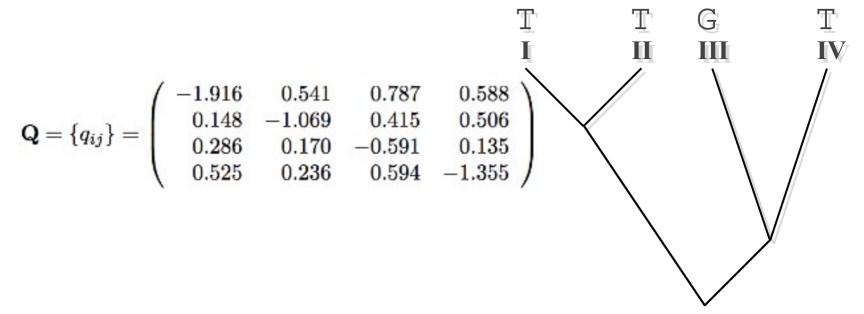


Using Monte Carlo simulation to estimate site likelihood is pedagogical, but too inefficient for the analysis of real data.

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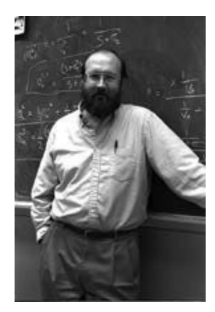
e.g., there are >1,000,000 possible site patterns for a tree with 10 species.

# Plan: Criteria for evaluating phylogenies

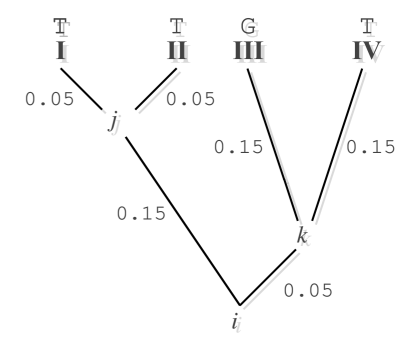
- Criteria for evaluating phylogenetic trees:
  - Parsimony
  - Distance methods
  - Maximum Likelihood
    - Using a Monte Carlo simulation
    - Using Felsenstein's pruning algorithm
  - Posterior probability (Bayesian approach)
- Conventions:
  - We're dealing with aligned sequence data
  - gaps are not taken into account

### The Felsenstein Pruning Algorithm

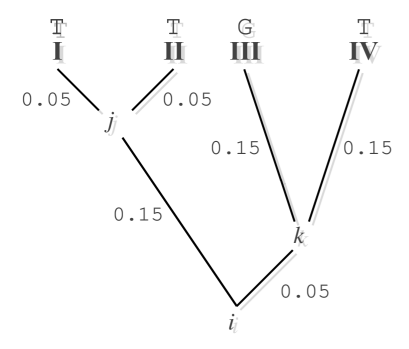
Make the math mirror the tree to avoid redundant calculations.

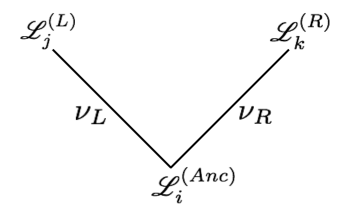


Joe Felsenstein (c.1981)

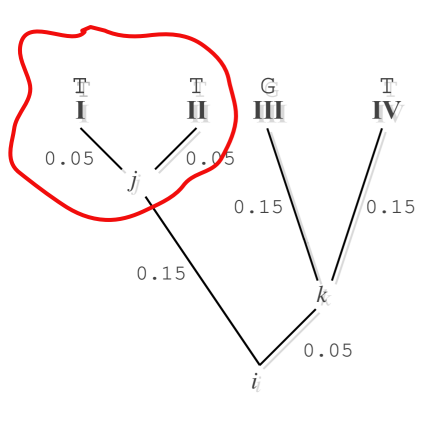


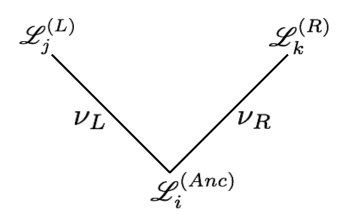
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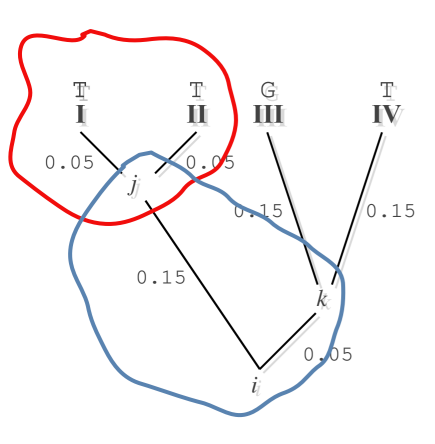


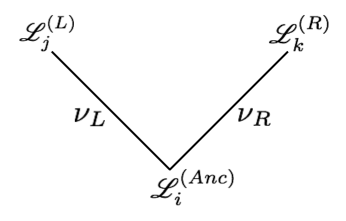
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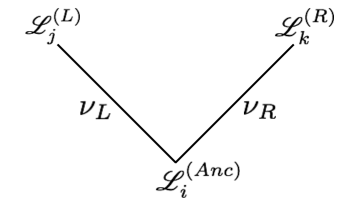
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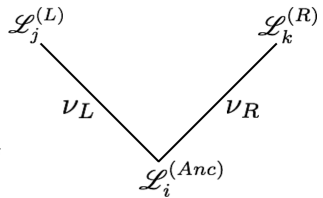
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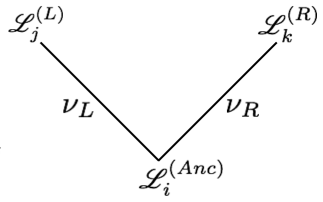
The conditional likelihoods  $\mathcal{L}_i^{(Anc)}$  are the probabilities of the observations above some point in the tree (a local ancestral node, Anc), conditional on state i at that node.



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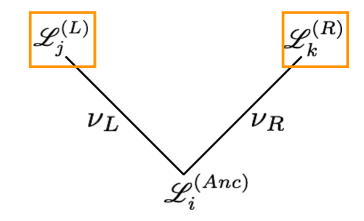
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$$\mathcal{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathcal{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathcal{L}_{k}^{(R)}\right)$$

where  $i \in (A,C,G,T)$ 

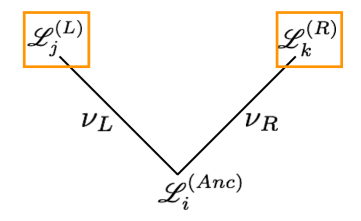
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These conditional likelihoods are the likelihoods of the observations above the end of each branch...

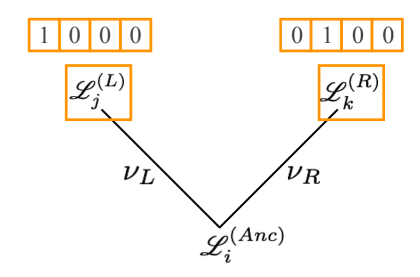
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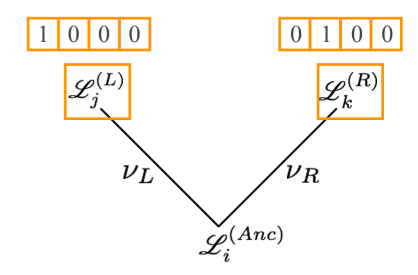


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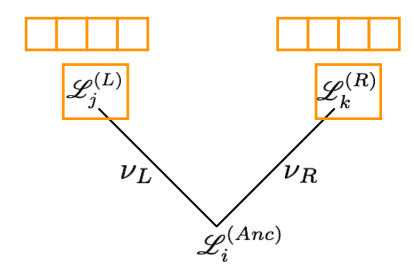


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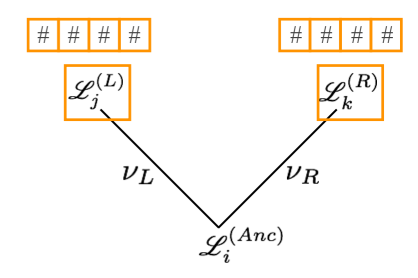
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#### The Felsenstein Pruning Algorithm



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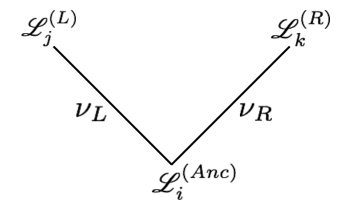
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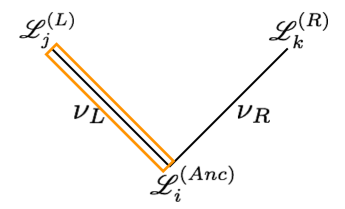
So, *how* do we calculate the conditional likelihoods?



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#### The Felsenstein Pruning Algorithm

So, how do we calculate the conditional likelihoods?

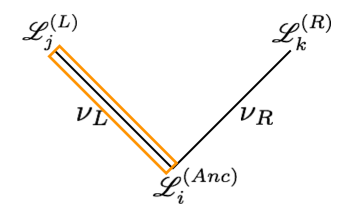


$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_{L}) \mathscr{L}_{j}^{(L)}\right)$$

Let's first focus on the left descendant branch.

#### The Felsenstein Pruning Algorithm

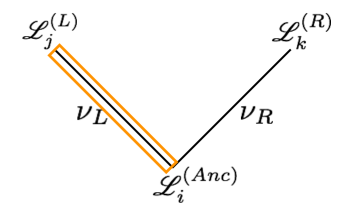
So, how do we calculate the conditional likelihoods?



$$\mathcal{L}_i^{(Anc)} = \left( \sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathcal{L}_j^{(L)} \right)$$
 for each possible end state sum... ...the product of the transition probabilities of changes from  $i$  to  $j$  over branch  $L$ ... ...and the conditional likelihood of each end state,  $j$ .

## The Felsenstein Pruning Algorithm

So, how do we calculate the conditional likelihoods?

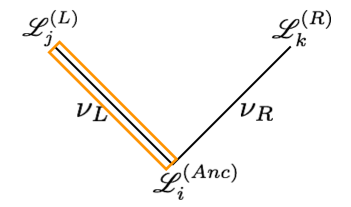


$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right)$$

Recall that the transition probabilities,  $\mathbf{P}_{ij}(v_L)$ , of histories ending in state j that were initiated from state i and run over branch L are either approximated (by Monte Carlo simulation) or solved by exponentiating the product of the instantaneous-rate matrix,  $\mathbf{Q}$ , and branch length,  $v_L$ 

#### The Felsenstein Pruning Algorithm

So, *how* do we calculate the conditional likelihoods?



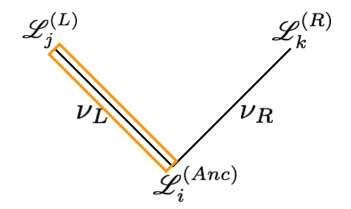
$$\mathcal{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathcal{L}_{j}^{(L)}\right) \mathbf{Q} = q_{ij} = \begin{pmatrix} -\mu a\pi_C & \mu b\pi_G & \mu c\pi_T \\ \mu a\pi_A & -\mu d\pi_G & \mu e\pi_T \\ \mu b\pi_A & \mu d\pi_C & -\mu f\pi_T \\ \mu c\pi_A & \mu e\pi_C & \mu f\pi_G & - \end{pmatrix}$$

$$\mathbf{P}_{ij}(\nu) = e^{\mathbf{Q}\nu}$$

Recall that the transition probabilities,  $\mathbf{P}_{j}(v_{L})$ , of histories ending in state j that were initiated from state i and run over branch L are either approximated (by Monte Carlo simulation) or solved by exponentiating the product of the instantaneous-rate matrix,  $\mathbf{Q}$ , and branch length,  $v_{L}$ 

#### The Felsenstein Pruning Algorithm

So, how do we calculate the conditional likelihoods?

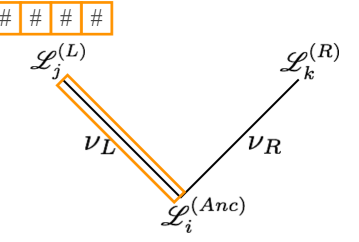


$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right)$$

The probability that the process is in state j at the end of the branch, times...

## The Felsenstein Pruning Algorithm

So, *how* do we calculate the conditional likelihoods?



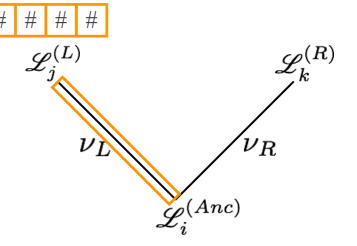
$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (\mathrm{A,C,G,T})} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right)$$

The probability that the process is in state j at the end of the branch, times...

...the probability of everything 'above' the end of the branch, conditional on state *j*.

## The Felsenstein Pruning Algorithm

So, *how* do we calculate the conditional likelihoods?

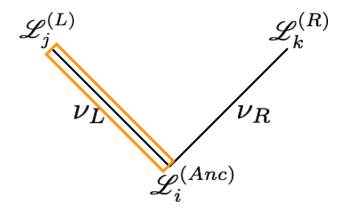


$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in ( ext{A,C,G,T})} p_{ij}(
u_L) \mathscr{L}_{j}^{(L)} \right)$$

We iteratively compute this product for each of the four j end states, A,C,G,T...

#### The Felsenstein Pruning Algorithm

So, how do we calculate the conditional likelihoods?

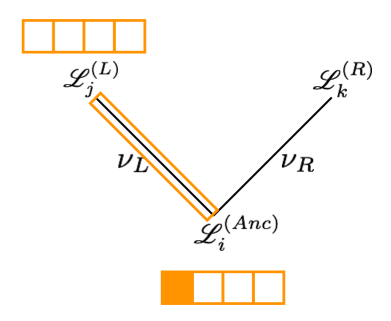


$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A, C, G, T)} p_{ij}(\nu_{L}) \mathscr{L}_{j}^{(L)}\right)$$

We iteratively compute this product for each of the four j end states, A,C,G,T... and then we sum these four terms because they are mutually exclusive.

## The Felsenstein Pruning Algorithm

First, set the start state, i, to i = A

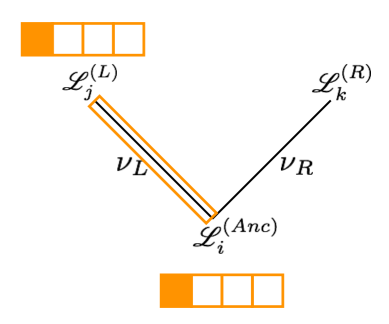


$$\mathcal{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_{L}) \mathcal{L}_{j}^{(L)}\right)$$
 $\mathcal{L}_{A}^{(Anc)} = \mathcal{L}_{A}^{(L)}$ 

The transition probabilities  $p_{AA}(v_L)$  over branch  $v_L$  are calculated by exponentiating the product of the instantaneous rate matrix,  $\mathbf{Q}$ , and branch length  $v_L$ .

## The Felsenstein Pruning Algorithm

First, set the start state, i, to i = A and set the end state, j, to j = A

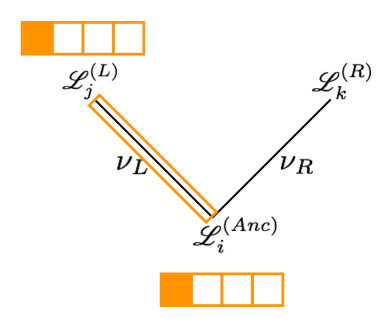


$$egin{aligned} \mathscr{L}_{i}^{(Anc)} &= \left(\sum_{j \in ( ext{A,C,G,T})} p_{ij}(
u_{L}) \mathscr{L}_{j}^{(L)} 
ight) \ \mathscr{L}_{ ext{A}}^{(Anc)} &= \mathscr{L}_{ ext{A}}^{(L)} \end{aligned}$$

The transition probabilities  $p_{AA}(v_L)$  over branch  $v_L$  are calculated by exponentiating the product of the instantaneous rate matrix,  $\mathbf{Q}$ , and branch length  $v_L$ .

#### The Felsenstein Pruning Algorithm

First, set the start state, i, to i = A and set the end state, j, to j = A next set the end state, j, to j = C...

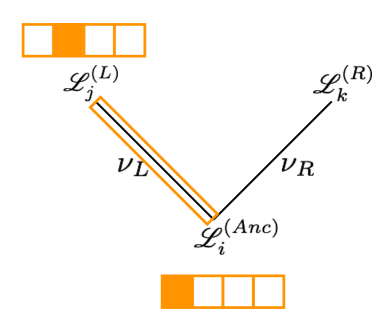


$$\mathcal{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_{L}) \mathcal{L}_{j}^{(L)}\right)$$
 $\mathcal{L}_{A}^{(Anc)} = \mathcal{L}_{C}^{(Anc)}$ 

Then we do the same thing for end state j = C...

## The Felsenstein Pruning Algorithm

First, set the start state, i, to i = A and set the end state, j, to j = A next set the end state, j, to j = C then set the end state, j, to j = G...

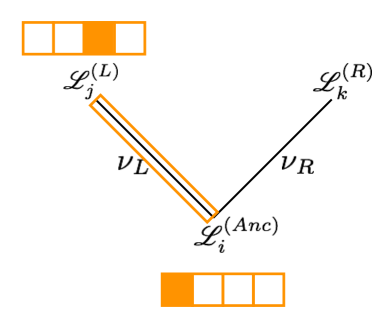


$$egin{aligned} \mathscr{L}_{i}^{(Anc)} &= \left(\sum_{j \in ( ext{A,C,G,T})} p_{ij}(
u_L) \mathscr{L}_{j}^{(L)} 
ight) \ \mathscr{L}_{ ext{A}}^{(Anc)} &= \mathscr{L}_{ ext{G}}^{(Anc)} \end{aligned}$$

Then we do the same thing for end state j = G...

## The Felsenstein Pruning Algorithm

First, set the start state, i, to i = A and set the end state, j, to j = A next set the end state, j, to j = C then set the end state, j, to j = G then set the end state, j, to j = G

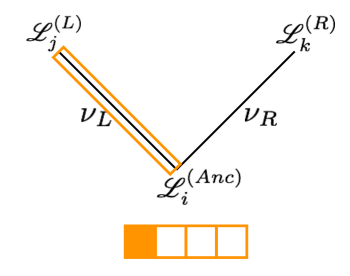


$$egin{aligned} \mathscr{L}_{i}^{(Anc)} &= \left(\sum_{j \in ( ext{A,C,G,T})} p_{ij}(
u_L) \mathscr{L}_{j}^{(L)} 
ight) \ \mathscr{L}_{ ext{A}}^{(Anc)} &= \mathscr{L}_{ ext{T}}^{(Anc)} \end{aligned}$$

Finally, we do the same thing for end state j = T

## The Felsenstein Pruning Algorithm

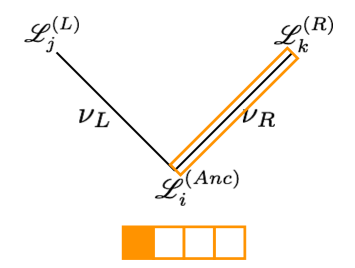
Next, we sum the fractional likelihoods for each of the four end states  $j = \{A, C, G, T\}...$ 



$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right)$$

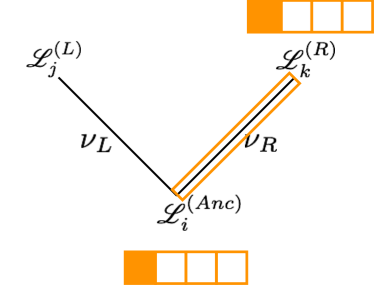
Then we sum the fractional likelihoods for each of the four end states  $j = \{A, C, G, T\}...$ 

## The Felsenstein Pruning Algorithm



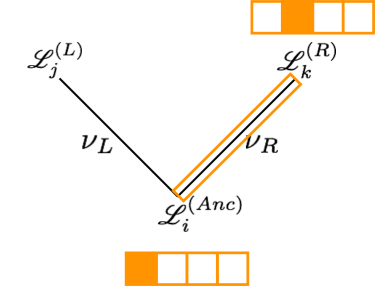
$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm



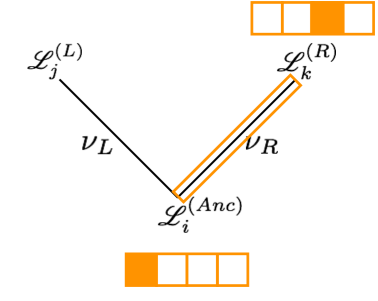
$$\mathcal{L}_i^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathcal{L}_j^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathcal{L}_k^{(R)}\right)$$

## The Felsenstein Pruning Algorithm



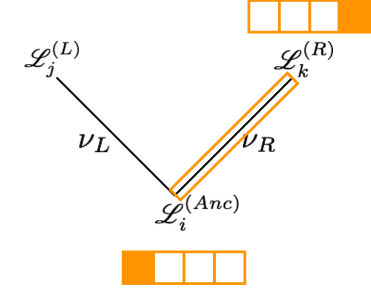
$$\mathcal{L}_i^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathcal{L}_j^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathcal{L}_k^{(R)}\right)$$

## The Felsenstein Pruning Algorithm



$$\mathcal{L}_i^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathcal{L}_j^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathcal{L}_k^{(R)}\right)$$

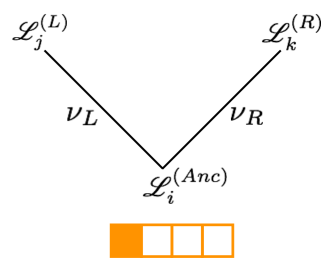
## The Felsenstein Pruning Algorithm



$$\mathcal{L}_i^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathcal{L}_j^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathcal{L}_k^{(R)}\right)$$

#### The Felsenstein Pruning Algorithm

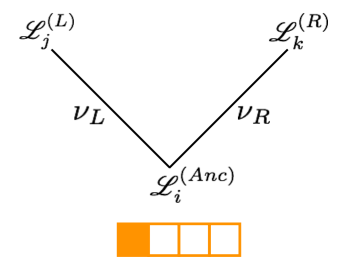
Finally, we calculate  $\mathcal{L}_A^{(Anc)}$  as the product of the conditional likelihoods of the two descendant branches, which makes explicit the assumption that substitutions along these two lineages are independent.



$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

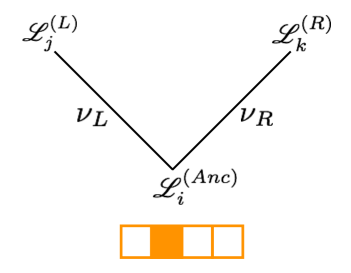
#### The Felsenstein Pruning Algorithm

Now we have computed  $\mathscr{L}_{A}^{(Anc)}$ ...



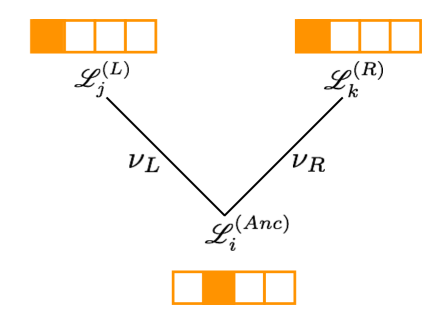
$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A, C, G, T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A, C, G, T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

#### The Felsenstein Pruning Algorithm



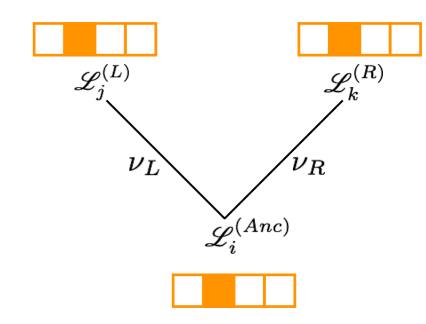
$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm



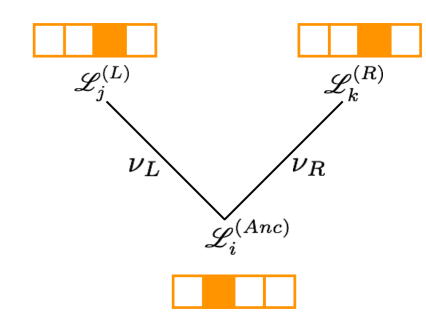
$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm



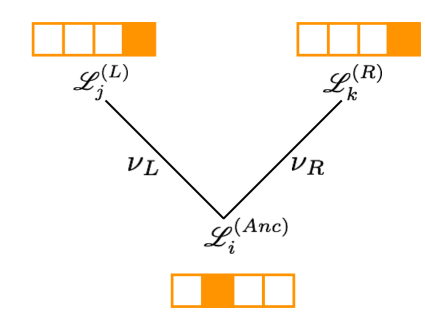
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## The Felsenstein Pruning Algorithm



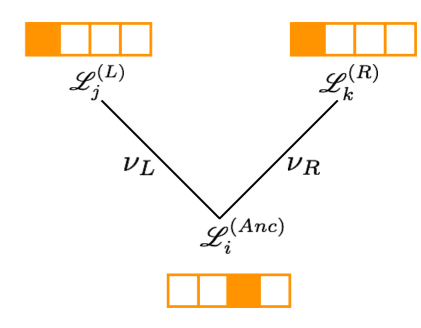
$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm



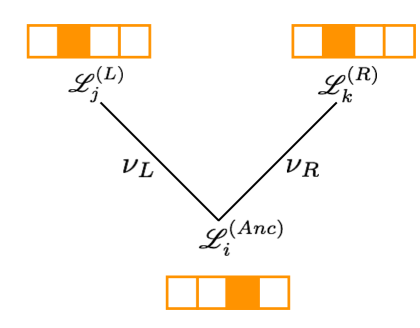
$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm



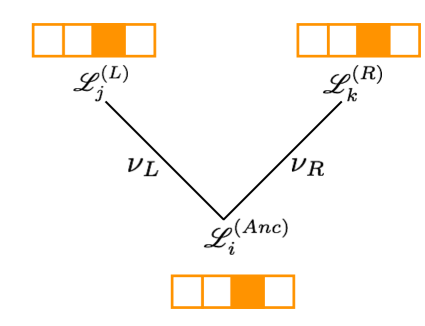
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## The Felsenstein Pruning Algorithm



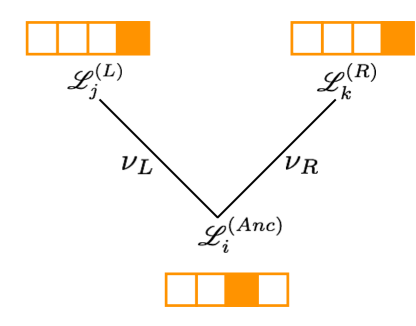
$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm



$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

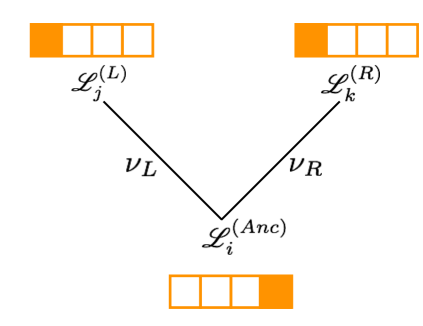
## The Felsenstein Pruning Algorithm



$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm

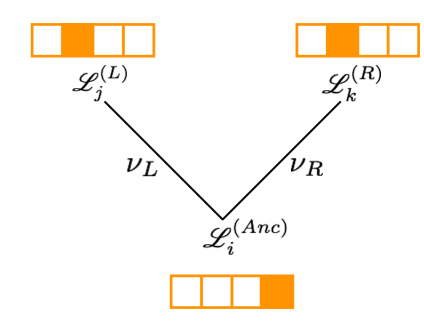
So, now we need to repeat the entire process for i = C to compute  $\mathcal{L}_{C}^{(Anc)}$  for i = G to compute  $\mathcal{L}_{G}^{(Anc)}$  and finally for i = T to compute  $\mathcal{L}_{T}^{(Anc)}$ ...



$$\mathcal{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathcal{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathcal{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm

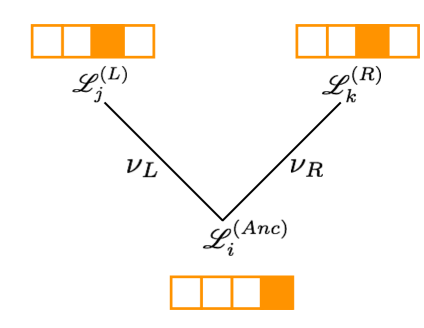
So, now we need to repeat the entire process for i = C to compute  $\mathcal{L}_{C}^{(Anc)}$  for i = G to compute  $\mathcal{L}_{G}^{(Anc)}$  and finally for i = T to compute  $\mathcal{L}_{T}^{(Anc)}$ ...



$$\mathcal{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathcal{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathcal{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm

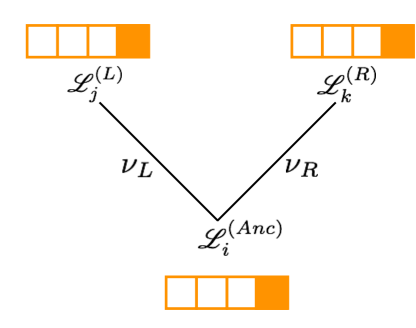
So, now we need to repeat the entire process for i = C to compute  $\mathscr{L}_{C}^{(Anc)}$  for i = G to compute  $\mathscr{L}_{G}^{(Anc)}$  and finally for i = T to compute  $\mathscr{L}_{T}^{(Anc)}$ ...



$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm

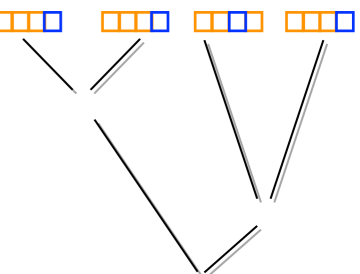
So, now we need to repeat the entire process for i = C to compute  $\mathscr{L}_{C}^{(Anc)}$  for i = G to compute  $\mathscr{L}_{G}^{(Anc)}$  and finally for i = T to compute  $\mathscr{L}_{T}^{(Anc)}$ ...



$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm

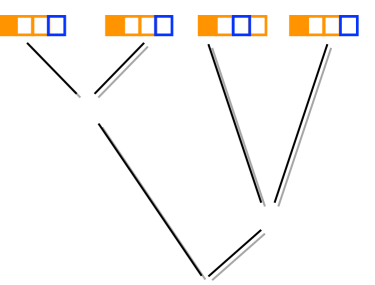
Then we need to repeat the entire process for each of the more inclusive nodes toward the root, where the conditional likelihoods of the tips are first recorded...



$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A, C, G, T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A, C, G, T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

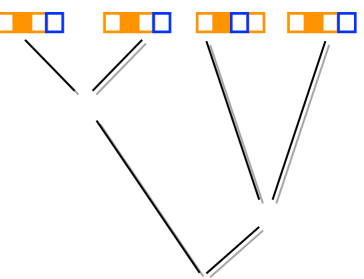
#### The Felsenstein Pruning Algorithm

Then we need to repeat the entire process for each of the more inclusive nodes toward the root, where the conditional likelihoods of the tips are first recorded...



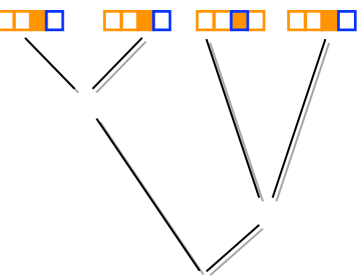
$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm



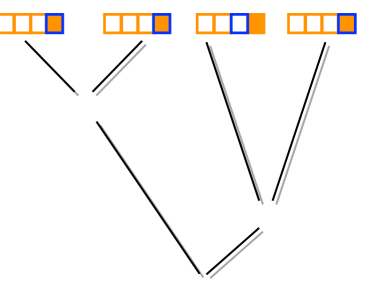
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## The Felsenstein Pruning Algorithm



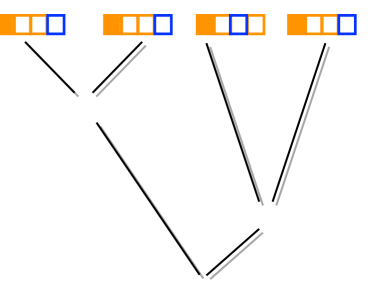
$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

#### The Felsenstein Pruning Algorithm



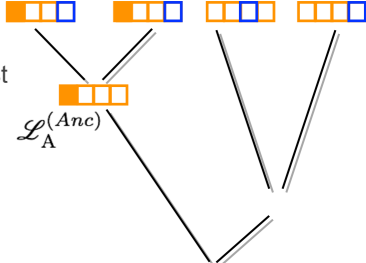
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#### The Felsenstein Pruning Algorithm



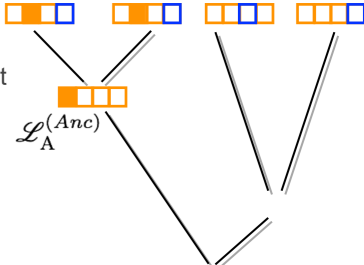
$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A,C,G,T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A,C,G,T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

## The Felsenstein Pruning Algorithm



$$\mathscr{L}_{i}^{(Anc)} = \left(\sum_{j \in (A, C, G, T)} p_{ij}(\nu_L) \mathscr{L}_{j}^{(L)}\right) \times \left(\sum_{k \in (A, C, G, T)} p_{ik}(\nu_R) \mathscr{L}_{k}^{(R)}\right)$$

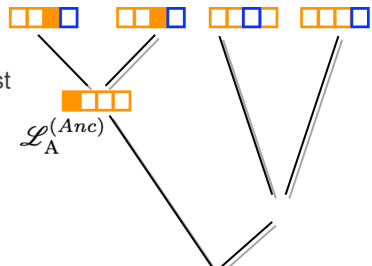
#### The Felsenstein Pruning Algorithm



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#### The Felsenstein Pruning Algorithm

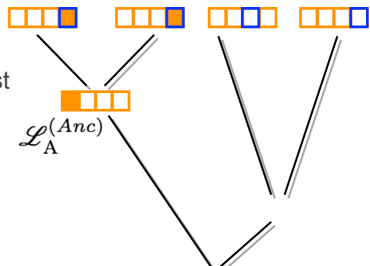
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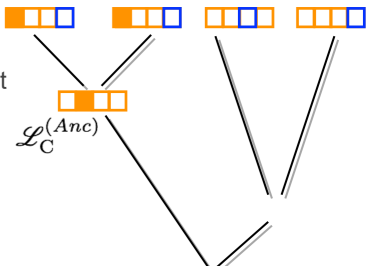
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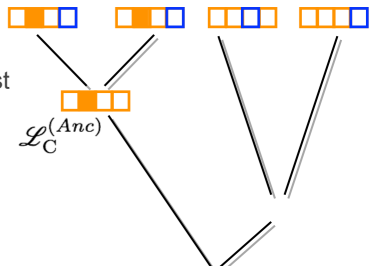
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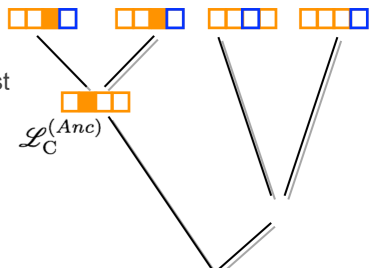
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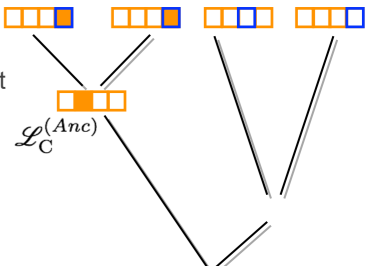
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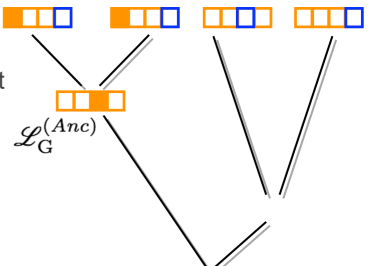
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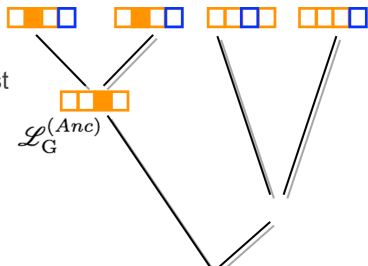
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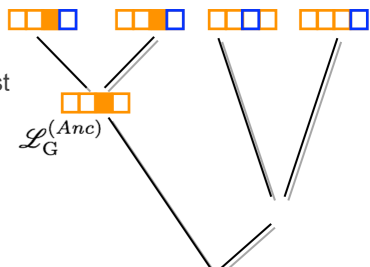
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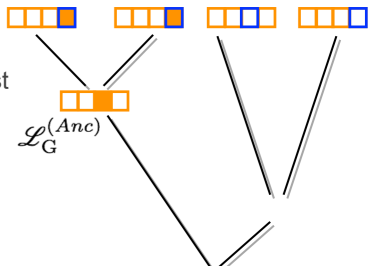
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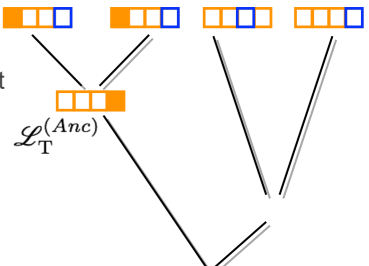
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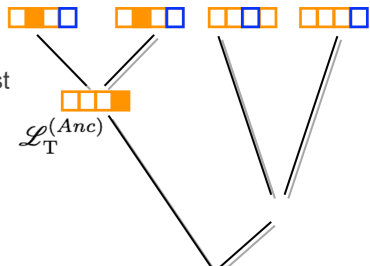
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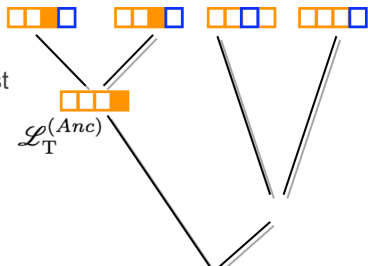
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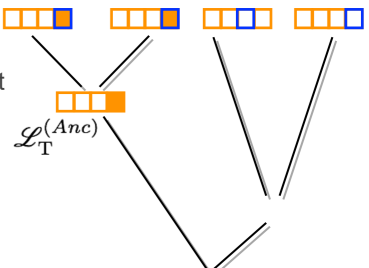
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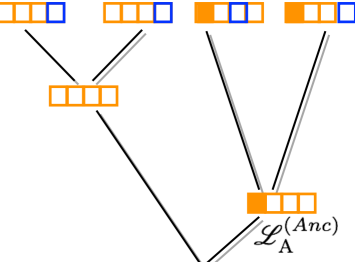
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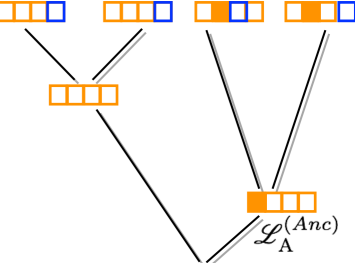
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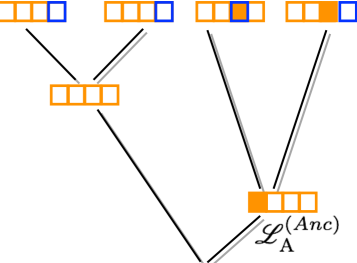
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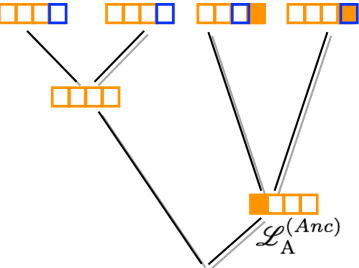
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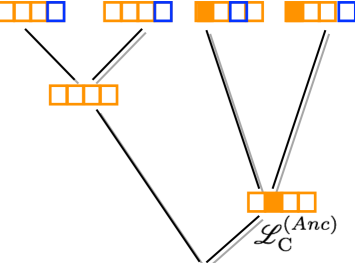
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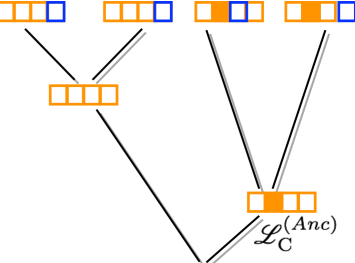
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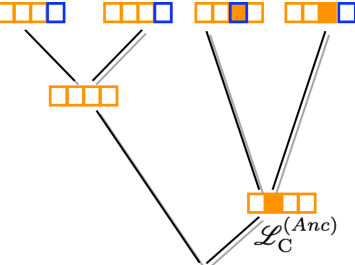
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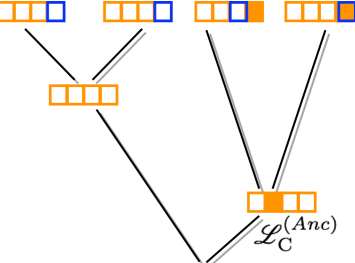
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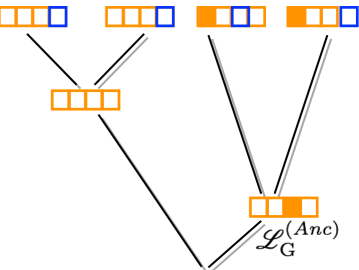
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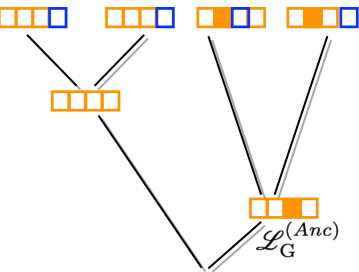
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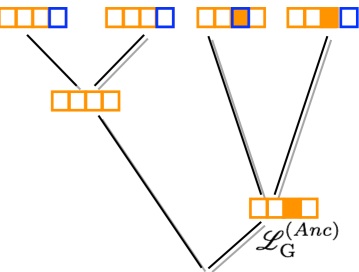
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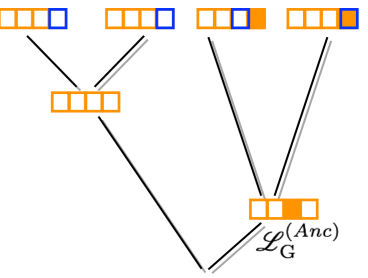
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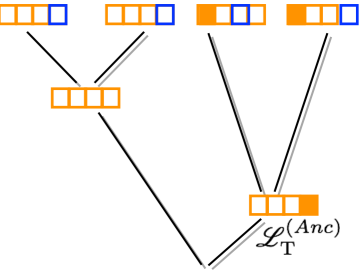
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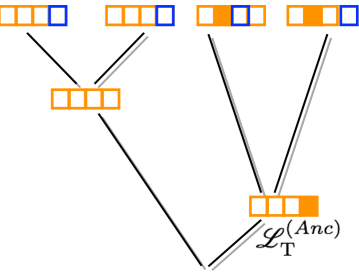
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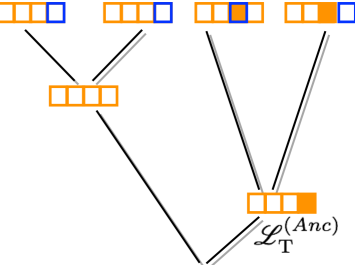
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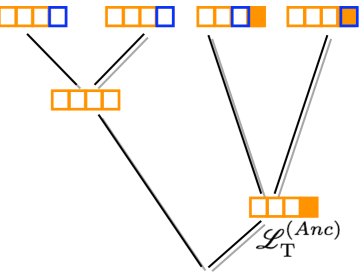
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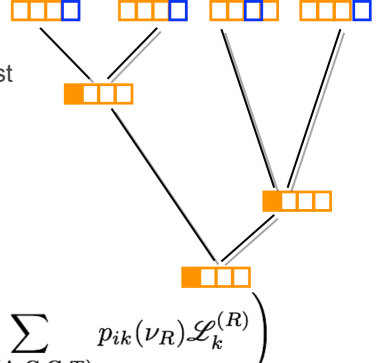
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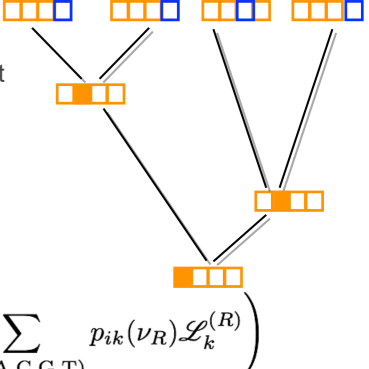
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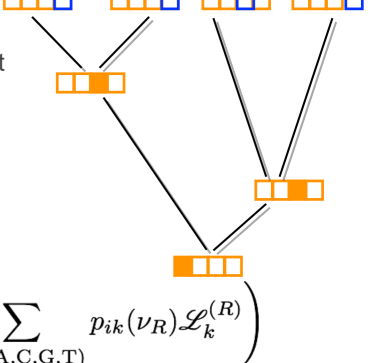
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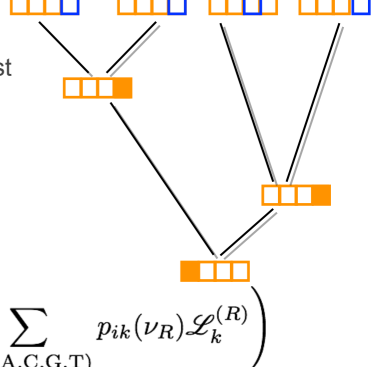
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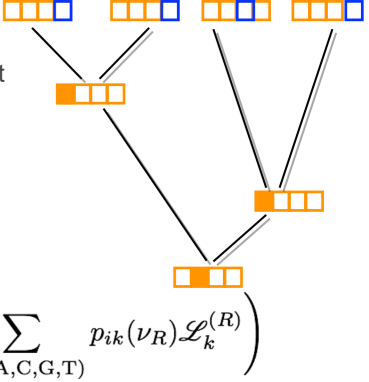
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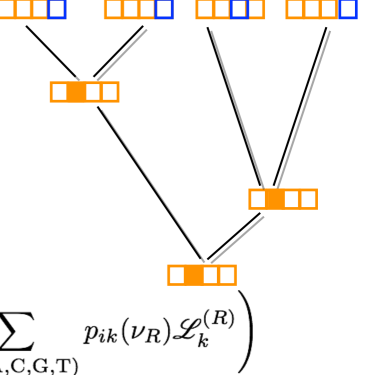
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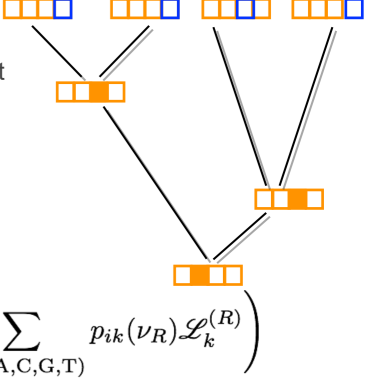
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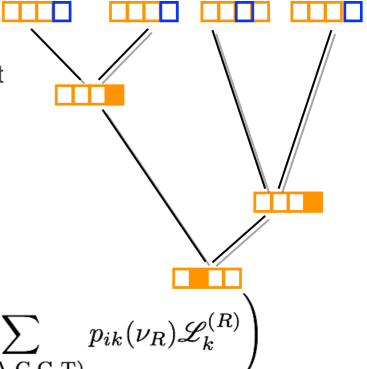
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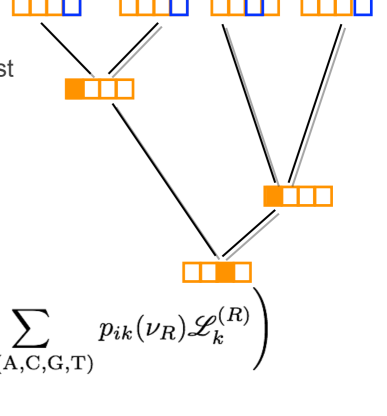
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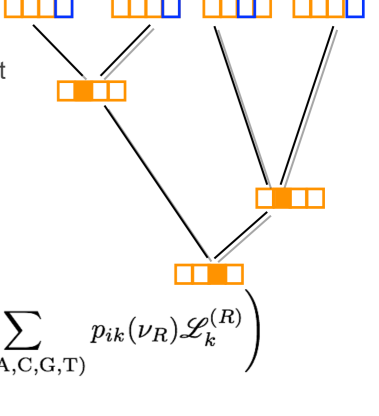
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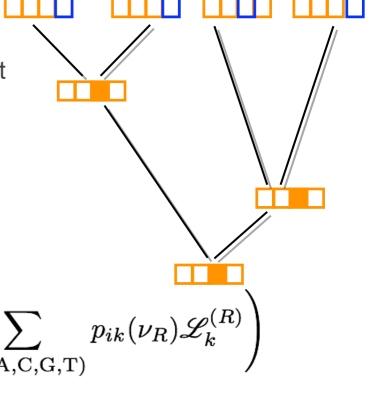
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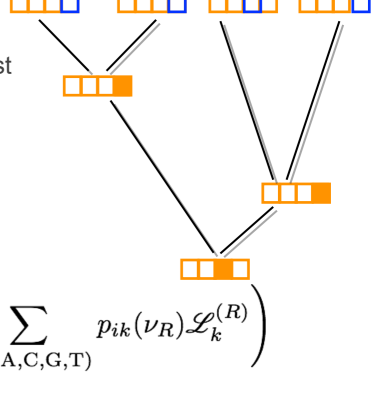
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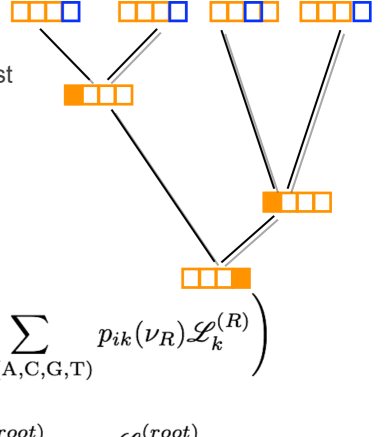
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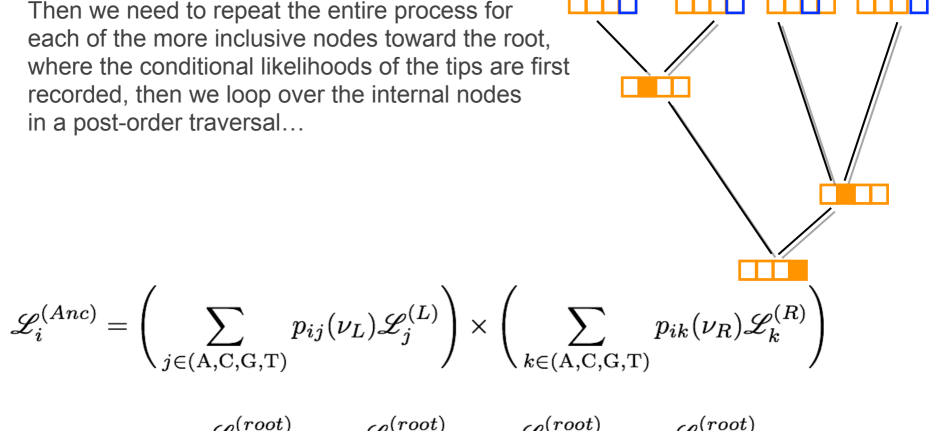
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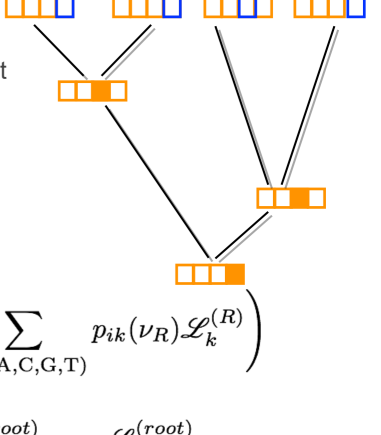
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  $\mathscr{L}_{\mathrm{C}}^{(root)}$   $\mathscr{L}_{\mathrm{G}}^{(root)}$   $\mathscr{L}_{\mathrm{T}}^{(root)}$ 

#### The Felsenstein Pruning Algorithm



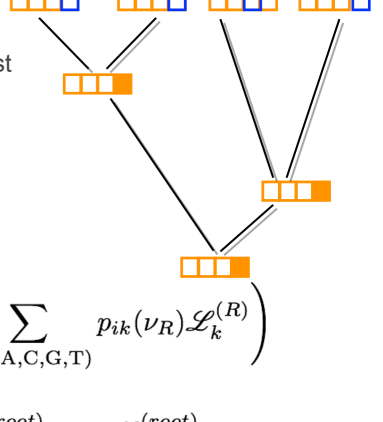
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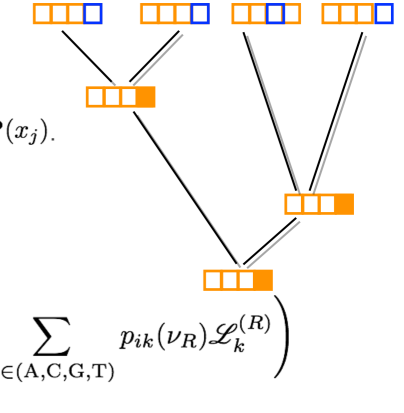
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$$\pi_{
m A} \mathcal{L}_{
m A}^{(root)}$$
  $\mathcal{L}_{
m C}^{(root)}$   $\mathcal{L}_{
m G}^{(root)}$   $\mathcal{L}_{
m T}^{(root)}$ 

The probability of everything in the tree above the root, *given* that the process is in state A,...

...times the probability that the process was in state A at the root.

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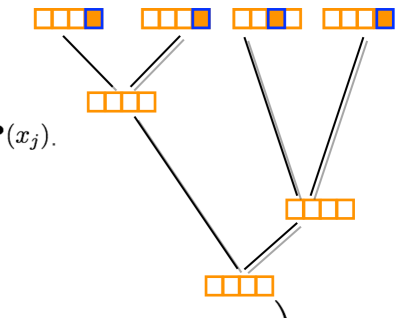
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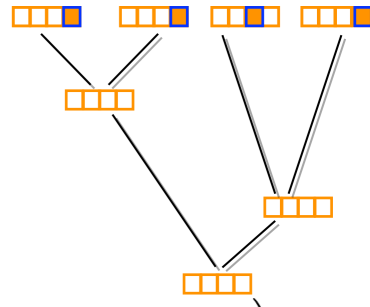
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...and then sum the four terms because these root states are mutually exclusive.

#### The Felsenstein Pruning Algorithm

Upon reaching the root of the tree, the conditional likelihood of each state is 'weighted' by the prior probability (stationary frequency) of the corresponding state,  $\pi_i$   $\pi_i$  to give the unconditional probability of the data,  $x_i$   $x_i$ 



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→ We have computed the likelihood of site pattern TTGT using Felsenstein's pruning algorithm!

#### Calculating the likelihood of an alignment assumes independence of sites

We generally assume that the substitution process is independent across sites, which allows us to combine their probabilities as the product of the N site likelihoods.

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Although this is a convenient assumption, it is not very biologically realistic.

More complex models can relax it, cf Jeff Thorne's talk.

# Plan: Criteria for evaluating phylogenies

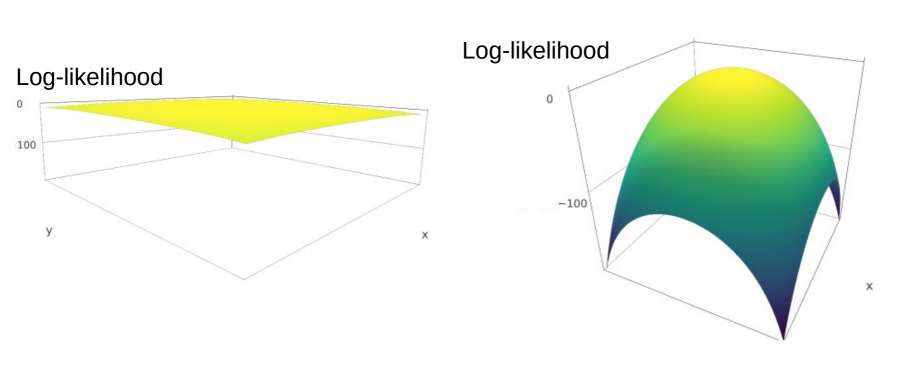
- Criteria for evaluating phylogenetic trees:
  - Parsimony
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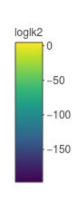
So far, we have presented all methods for phylogenetic reconstruction as attempting to find the best tree according to some criterion.

This optimization approach has some drawbacks.

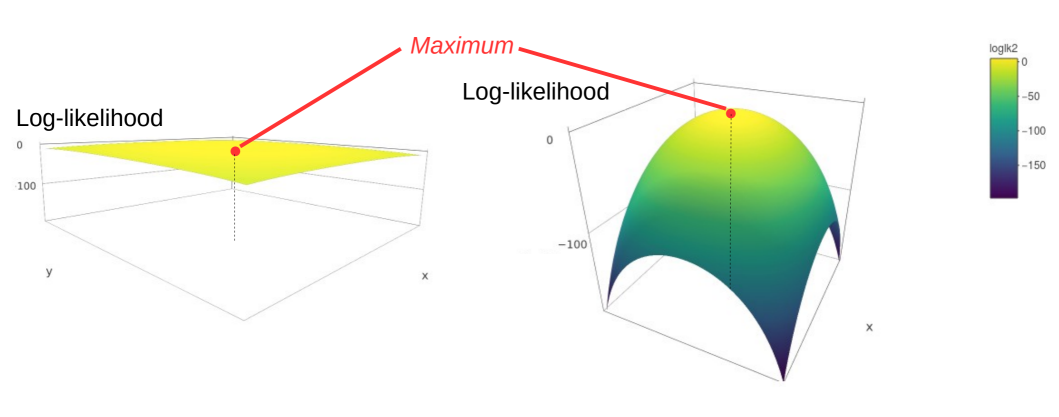
Let's see two examples.

**Ex. A**: likelihood surface for a simple model with two parameters:





**Ex. A**: likelihood surface for a simple model with two parameters:



**Ex. B**: Given an alignment, 9 trees have very similar likelihoods and are much more likely than all the other ones.

Taking the most likely tree only provides knowledge about 1 of those 9 almost equi-likely trees.

→ It would be better to take into account all 9 of them!

# Optimizing, integrating, sampling

- Optimizing: Looking for the topology and all other parameter values that are most likely
- Integrating: visiting all topologies and parameter values according to their probability

#### Pros:

- If we are interested only in the topology, we integrate over all "nuisance" parameters
- One gets confidence intervals for free
- Sampling: like integrating, but we do not attempt to be exhaustive in our exploration of parameter values

# Likelihood vs Bayesian approach for sampling

Likelihood :  $P(D|M,\theta)$ 

We want to sample parameter values  $\theta$  of model M.

How can we know that the 9 sets of parameter values  $\theta$  we have sampled are much more likely than all other  $\theta$ , without sampling everything?

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$$\sum_{\theta} P(\theta|D,M) = 1 \quad \rightarrow \text{True probability distribution.}$$
Posterior probability

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#### Simplifying the notation

When we write  $P(\theta|D,M)$ , we mean that we are computing the probability of parameter values  $\theta$  given model M and given data D.

Here *M* is used to represent the structure of the model.

#### For instance:

- The fact that all sites are independent
- The fact that they share a single value of the transition/transversion ratio
- The fact that we have a Birth-death prior on a chronogram
- Etc...

In some cases, we may want to integrate over different model structures M: e.g. a model with 1 value of the transition/transversion ratio, or 2, or 3...

In most cases, we use a single model M, so we will forget about it in most of the following:

$$P(\theta|D,M) \rightarrow P(\theta|D)$$

#### Bayes theorem:

$$P(\theta|D) = \frac{P(\theta \land D)}{P(D)} = \frac{P(D \land \theta)}{P(D)}$$

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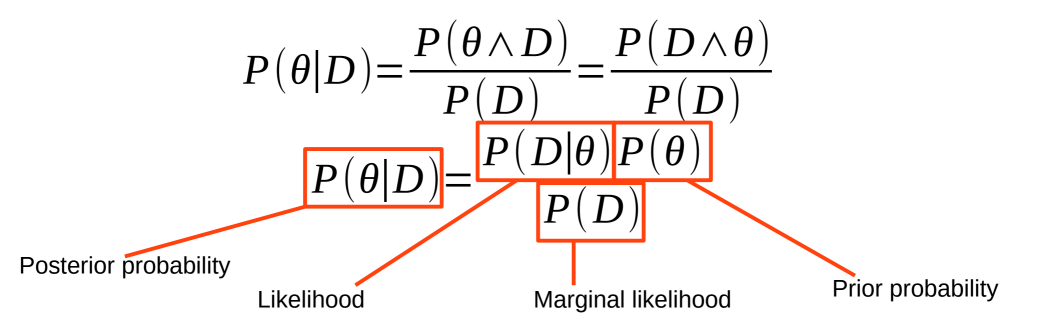
$$P(\theta|D) = \frac{P(D|\theta)P(\theta)}{P(D|\theta)d\theta}$$

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$$\begin{split} P(\theta|D) &= \frac{P(\theta \land D)}{P(D)} = \frac{P(D \land \theta)}{P(D)} \\ P(\theta|D) &= \frac{P(D|\theta)P(\theta)}{P(D)} \\ P(\theta|D) &= \frac{P(D|\theta)P(\theta)}{P(D|\theta)d\theta} \\ P(\theta|D) &= \frac{P(D|\theta)P(\theta)}{\int_{\theta} P(D|\theta)d\theta} \end{split} \quad \text{Hard to compute. In practice, we can ignore it.} \end{split}$$

In practice, we can often ignore it.

#### Bayes theorem:



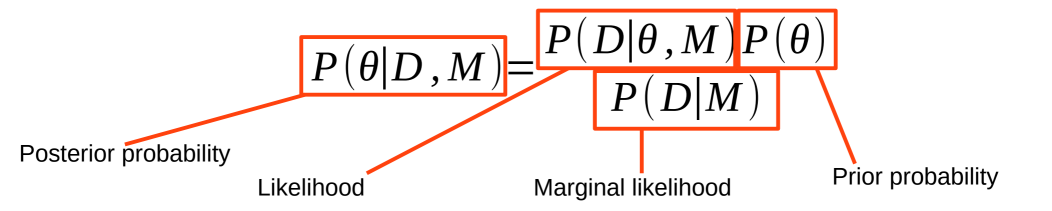
### The importance of prior probabilities

The Bayesian approach amounts to considering that all parameters of a model are random variables in a probabilistic world.

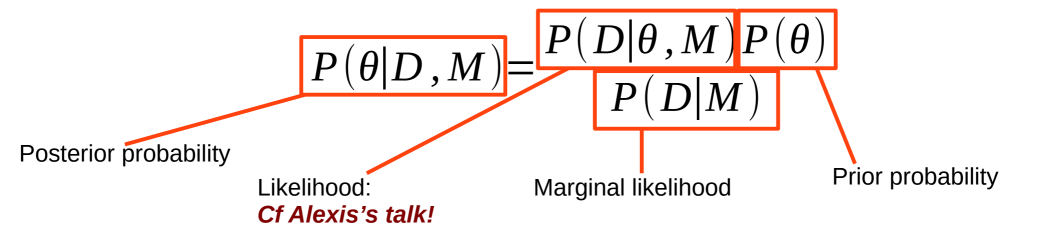
Therefore, one needs to assign probability distributions to those parameters: the **priors**.

$$P(\theta|D) = P(D|\theta) \frac{P(\theta)}{P(\theta)} / P(D)$$

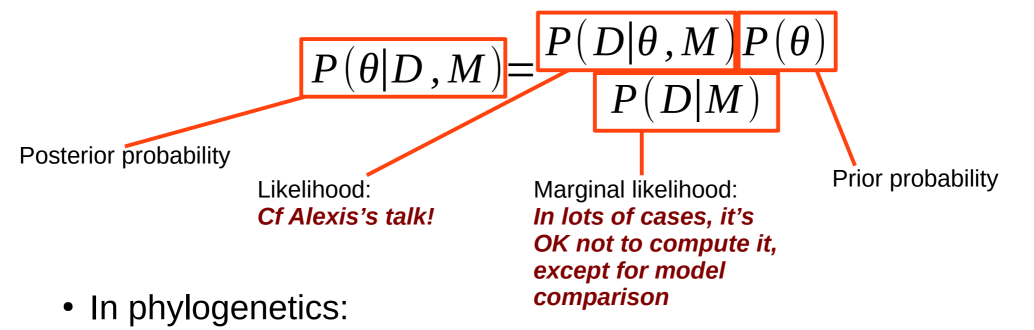
- → PROS: allows incorporating prior information coming from the analysis of other data: my conclusion relies on more than the tiny amount of data I have analyzed in a given experiment.
- → CONS: introduces prior information into the analysis: is my conclusion simply propagating my prior information?



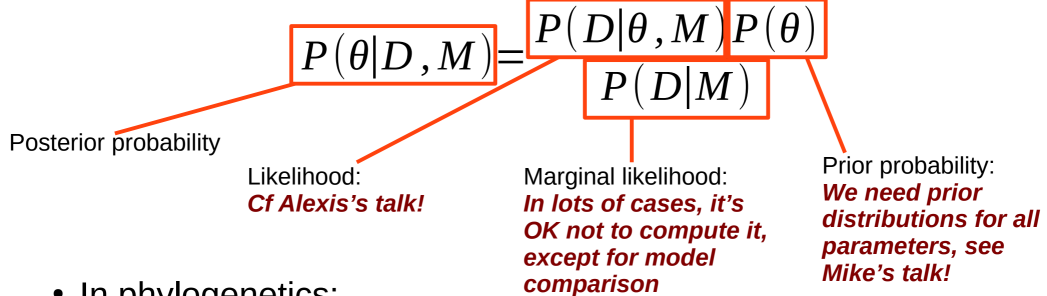
- In phylogenetics:
  - D: aligned sequence data
  - M: the model can be very complicated, but usually contains:
    - Topology
    - Branch lengths
    - Rate matrix
    - Etc...
  - $-\theta$ : the values of the parameters above



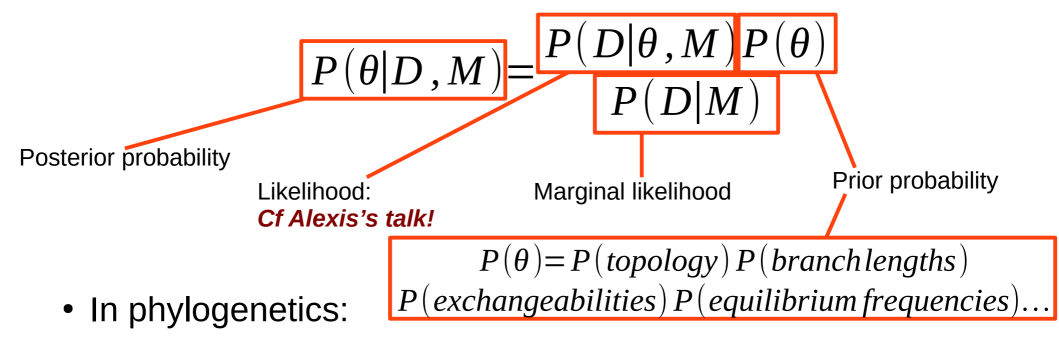
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## Smart sampling of $P(\theta|D)$

- Smart sampling = no need to sample a huge number of points!
- → Ideally : Sample trees and parameter values with a frequency equal to their probability.
- E.g.: we would sample one of our 9 trees 99% of the time! Several approaches:
  - Importance sampling
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Next time!

#### Conclusion

- One can perform inference according to the posterior probability of a probabilistic model
- That's what Bayesian inference is about
- It combines the likelihood with priors on parameter values
- In phylogenetics, the likelihood is typically computed thanks to Felsenstein's pruning algorithm (1981, cf Alexis)
- Then priors need to be defined for:
  - Topologies and branch lengths / chronograms
  - Rates
  - Other parameters... (cf Mike)
- When parameters are independent, to compute the prior of all parameter values, one only needs to compute the product over individual parameter priors

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Optimization algorithms (Alexis)

Bayesian inference and MCMC (Mike)