Phylogenetic inference and likelihood 2

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(With thanks to Mark Holder and Paul Lewis for slides)



Combining probabilities

 Multiply probabilities if the component events must happen simultaneously (i.e. where you would naturally use the word AND when describing the problem)

Using 2 dice, what is the probability of



$$(1/6) \times (1/6) = 1/36$$

AND rule in phylogenetics

probability of A at tip given ancestral state A probability of C at tip given ancestral state A we have observed these states pretend (for now) we also know p_1 these ancestral states $Pr(A, A, C, A, A) = p_0 \ p_1 \ p_2 \ p_3 \ p_4$ probability of A at end of edge given A at root probability of A at tip given One use of the AND rule in phylogenetics A at root is to combine probabilities associated with individual branches to produce the overall probability probability of the data for one site. of A at root

Combining probabilities

 Add probabilities if the component events are mutually exclusive (i.e. where you would naturally use the word OR in describing the problem)

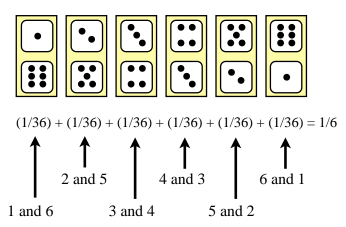
Using one die, what is the probability of



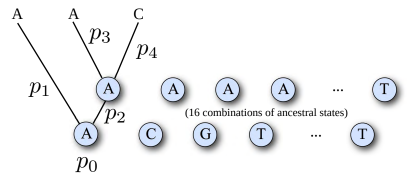
$$(1/6) + (1/6) = 1/3$$

Combining AND and OR

What is the probability that the sum of two dice is 7?



Using both AND and OR in phylogenetics



AND rule used to compute probability of the observed data for *each combination* of ancestral states.

OR rule used to combine different combinations of ancestral states.

Independence

This is always true...

$$Pr(A \text{ and } B) = Pr(A) Pr(B|A)$$
joint probability

conditional probability

If we can say this...

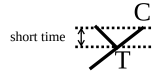
$$Pr(B|A) = Pr(B)$$

...then events A and B are **independent** and we can express the joint probability as the product of Pr(A) and Pr(B)

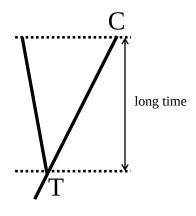
$$Pr(A \text{ and } B) = Pr(A) Pr(B)$$

Non-independence in molecular evolution

The state present in the descendant is **not independent** of the state in the ancestor



less probable



more probable

Conditional Independence

Assume both A and B depend on C:

$$Pr(A|C) \neq Pr(A)$$
 $Pr(B|C) \neq Pr(B)$

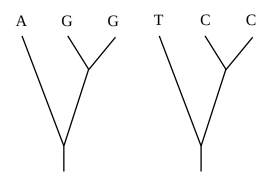
If we can say this...

$$Pr(B|A,\mathbb{C}) = Pr(B|\mathbb{C})$$

...then events A and B are **conditionally independent** and we can express the joint (conditional) probability as the product of Pr(A|C) and Pr(B|C)

$$Pr(A \text{ and } B|\mathbb{C}) = Pr(A|\mathbb{C}) Pr(B|\mathbb{C})$$

Conditional independence in molecular evolution



Paul O. Lewis (2017 Woods Hole Workshop in Molecular Evolution)

The site data patterns AGG and TCC are assumed by most models to be conditionally independent.

The patterns both depend on the underlying tree (including edge lengths) and the substitution model.

Pr(AGG and TCC|tree, model) = Pr(AGG|tree, model) Pr(TCC|tree, model)

Likelihood calculated from a single sequence

 $\Pr(A) = \pi_A$

First 32 nucleotides of the ψη-globin gene of gorilla:

 $Pr(C) = \pi_C$ $Pr(G) = \pi_G$

 $Pr(T) = \pi_T$

GAAGTCCTTGAGAAATAAACTGCACACACTGG

$$L = \pi_G \pi_A \pi_A \pi_G \pi_T \pi_C \pi_C \pi_T \pi_T \pi_G \pi_A \pi_G \pi_A \pi_A \pi_A \pi_T \pi_A \pi_A \pi_A \pi_C \pi_T \pi_G \pi_C \pi_A \pi_C \pi_A \pi_C \pi_A \pi_C \pi_T \pi_G \pi_G$$

$$= \pi_A^{12} \pi_C^{7} \pi_G^{7} \pi_T^{6}$$
Note that we are assuming independence among sites here

$$\log L = 12\log(\pi_A) + 7\log(\pi_C) + 7\log(\pi_G) + 6\log(\pi_T)$$

We can already see by eye-balling this that a model allowing **unequal** base frequencies will **fit better** than a model that assumes **equal** base frequencies because there are about twice as many As as there are Cs, Gs and Ts.

Model ranking using LRT or AIC

Likelihood Ratio Tests (LRT) and the Akaike Information Criterion (AIC) provide two ways to evaluate whether an **unconstrained** model fits the data significantly better than a **constrained** version of the same model.

Find maximum logL under the unconstrained model:

$$\begin{split} \log L_{\text{unconstrained}} &= 12 \log(\pi_A) + 7 \log(\pi_C) + 7 \log(\pi_G) + 6 \log(\pi_T) \\ &= 12 \log(0.375) + 7 \log(0.219) + 7 \log(0.219) + 6 \log(0.187) \\ &= -43.1 \end{split}$$

This model has 3 estimated parameters

Find maximum logL under the constrained model:

$$\begin{split} \log L_{\text{constrained}} &= 12 \log(\pi_A) + 7 \log(\pi_C) + 7 \log(\pi_G) + 6 \log(\pi_T) \\ &= 12 \log(0.25) + 7 \log(0.25) + 7 \log(0.25) + 6 \log(0.25) \\ &= -44.4 \end{split}$$

This model has 0 estimated parameters

Likelihood Ratio Test (LRT)

Calculate the likelihood ratio test statistic:

$$\begin{split} R &= -2 \left[\log(L_{\text{constrained}}) - \log(L_{\text{unconstrained}}) \right] \\ &= -2 \left[-44.4 - (-43.1) \right] \\ &= 2.6 \end{split}$$
 (Note that the log-likelihoods used in the test statistic have been *maximized* under each model separately)

"unconstrained" does fit better than "constrained" (-43.1 > -44.4), but not significantly better (P = 0.457, chi-squared with 3 d.f.*)

*The number of degrees of freedom equals the difference between the two models in the number of estimated parameters. In this case, unconstrained has 3 parameters and constrained has 0, so d.f. = 3 - 0 = 3

Comparing models in phylogenetics can be challenging, as topologies are not nested within one another.

We will discuss appropriate statistical approaches later in the course.

Likelihood of the simplest tree

sequence 1 _____ sequence 2

To keep things simple, assume that the sequences are only 2 nucleotides long:

GA

froot (arbitrary)

$$L = L_1 L_2$$
 $= \left[\left(\frac{1}{4} \right) \left(\frac{1}{4} + \frac{3}{4} e^{-4\alpha t} \right) \right] \left[\left(\frac{1}{4} \right) \left(\frac{1}{4} - \frac{1}{4} e^{-4\alpha t} \right) \right]$

Pr(G)

Pr(G|G, αt)

Pr(A)

Pr(G|A, αt)

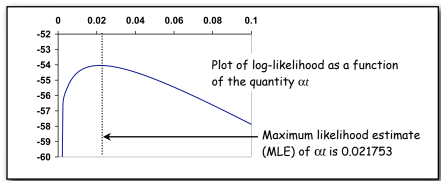
Note that we are NOT assuming independence here

Maximum likelihood estimation

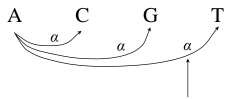
First 32 nucleotides of the $\psi\eta$ -globin gene of gorilla and orangutan:

gorilla GAAGTCCTTGAGAAATAAACTGCACACACTGG orangutan GGACTCCTTGAGAAATAAACTGCACACACTGG

$$L = \left[\left(\frac{1}{4} \right) \left(\frac{1}{4} + \frac{3}{4} e^{-4\alpha t} \right) \right]^{30} \left[\left(\frac{1}{4} \right) \left(\frac{1}{4} - \frac{1}{4} e^{-4\alpha t} \right) \right]^2$$



number of substitutions = rate \times time

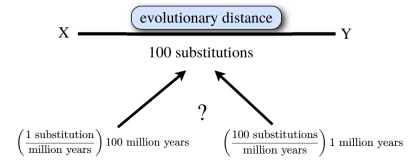


This is the rate at which an existing A changes to a T

Overall substitution rate is 3α , so the expected number of substitutions (ν) is

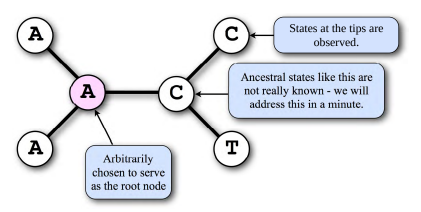
$$v = 3\alpha t$$

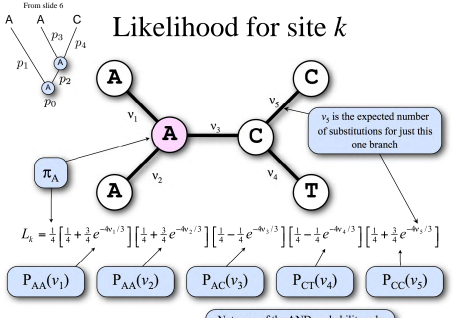
Rate and time are confounded



Likelihood of an unrooted tree

(data shown for only one site)



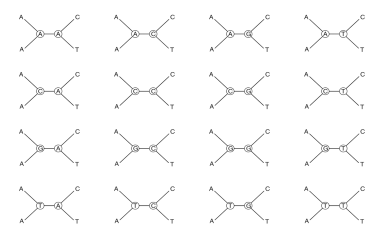


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Note use of the AND probability rule

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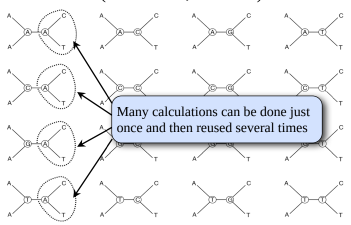
Brute force approach would be to calculate L_k for all 16 combinations of ancestral states and sum them



Note use of the OR probability rule

Pruning algorithm

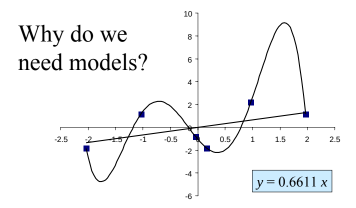
(same result, less time)



Felsenstein, J. 1981. Evolutionary trees from DNA sequences: a maximum likelihood approach. *Journal of Molecular Evolution* 17:368-376

Substitution Models

$y = -1.5972 x^5 + 23.167 x^4 - 126.18 x^3 + 319.17 x^2 - 369.22 x + 155.67$



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Models

- Models help us intelligently interpolate between our observations for purposes of making predictions
- Adding parameters to a model generally increases its fit to the data
- Underparameterized models lead to poor fit to observed data points
- Overparameterized models lead to poor prediction of future observations
- Criteria for choosing models include likelihood ratio tests, AIC, BIC, Bayes Factors, etc.
 - all provide a way to choose a model that is neither underparameterized nor overparameterized

Jukes-Cantor (JC69) model

- The four bases (A, C, G, T) are expected to be **equally** frequent in sequences ($\pi_A = \pi_C = \pi_G = \pi_T = 0.25$)
- Assumes **same rate** for all types of substitution $(r_{A\leftrightarrow C} = r_{A\leftrightarrow G} = r_{A\leftrightarrow T} = r_{C\leftrightarrow G} = r_{C\leftrightarrow T} = r_{G\leftrightarrow T} = \alpha)$
- Usually described as a **1-parameter** model (the parameter being the edge length)
 - Remember, however, that each edge in a tree can have its own length, so there are really as many parameters in the model as there are edges in the tree!
- Assumes substitution is a **Markov** process...

Jukes, T. H., and C. R. Cantor. 1969. Evolution of protein molecules. Pages 21-132 in H. N. Munro (ed.), Mammalian Protein Metabolism. Academic Press, New York.

What is a Markov process?

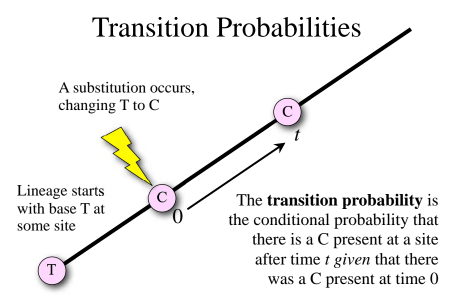
A substitution occurs, changing T to C

Lineage starts with base T at

To predict which base will be present after some time *t* we need know only which base was present at time 0 (C in this case).

If it is irrelevant that there was a T present at this site before time 0, then this is a Markov model.

some site



Jukes-Cantor transition probabilities

Here is the probability that a site starting in state T will end up in state G after time t when the individual substitution rates are all α :

$$P_{TG}(t) = \frac{1}{4} \left(1 - e^{-4\alpha t} \right) = \Pr(G|T, \alpha t)$$

The JC69 model has only one unknown quantity: αt

(The symbol *e* represents the base of the natural logarithms: its value is 2.718281828459045...)

Where does a transition probability formula such as this come from?

"ACHNyons" vs. substitutions

ACHN =
"Anything
Can Happen
Now"

When an *achnyon* occurs, any base can appear in a sequence.

Note: achnyon is *my term* for this make-believe event. You will not see this term in the literature.

T

If the base that A / \
appears is different C G
from the base that
was already there, then a
substitution event has occurred.

The rate (α) at which any *particular* substitution occurs will be 1/4 the achnyon rate (μ). That is, $\alpha = \mu/4$

(or $\mu = 4\alpha$)

The Poisson distribution

Probability distribution on the number of events when:

- 1. events are assumed to be independent,
- 2. the *rate* of events some constant, μ , and
- 3. the process continues for some duration of time, t.

The expectation of the number of events is $\nu = \mu t$.

Note that ν can be any non-negative number, but the Poisson is a discrete distribution – it gives the probabilities of the number of events (and this number will always be a non-negative integer).

Poisson distibution can be used to explain statistical regularities of rare events

P (k events in interval) = $\frac{e^{-\nu}\nu^k}{k!}$

- \triangleright ν is the average number of events per interval (rate times time)
- ightharpoonup e is the number 2.71828... (Euler's number) the base of the natural logarithms
- ▶ k takes values 0, 1, 2, ...
- $k! = k * (k-1) * (k-2) * \dots * 2 * 1$ is the factorial of k.

from wikipedia

Deriving a transition probability

Calculate the probability that a site currently T will change to G over time t when the rate of this particular substitution is α :

$$Pr(zero \ achnyons) = e^{-\mu t}$$
 (Poisson probability of zero events)

$$Pr(at least 1 achnyon) = 1 - e^{-\mu t}$$

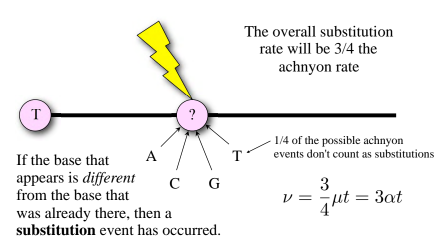
Pr(last achnyon results in base G) = $\frac{1}{4}$

Pr(end in G | start in T) =
$$\frac{1}{4} (1 - e^{-\mu t})$$

Remember that the rate (α) of any particular substitution is one fourth the achnyon rate (μ) :

$$P_{GT}(t) = \frac{1}{4} \left(1 - e^{-4\alpha t} \right)$$

Expected number of substitutions



Transition Probabilities: Remarks

$$P_{TA}(t) = \frac{1}{4}(1 - e^{-4\alpha t})$$

$$P_{TC}(t) = \frac{1}{4}(1 - e^{-4\alpha t})$$

$$P_{TG}(t) = \frac{1}{4}(1 - e^{-4\alpha t})$$

$$P_{TT}(t) = \frac{1}{4}(1 - e^{-4\alpha t})$$

These should add to 1.0 because T *must* change to something!

$$1 - e^{-4\alpha t}$$

Doh! Something must be wrong here...

Transition Probabilities: Remarks

$$P_{TA}(t) = \frac{1}{4}(1 - e^{-4\alpha t})$$

$$P_{TC}(t) = \frac{1}{4}(1 - e^{-4\alpha t})$$

$$P_{TG}(t) = \frac{1}{4}(1 - e^{-4\alpha t})$$

$$P_{TT}(t) = \frac{1}{4}(1 - e^{-4\alpha t}) + e^{-4\alpha t}$$

Forgot to account for the possibility of *no* acnyons over time *t*

Equilibrium frequencies

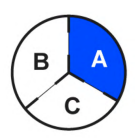
- The JC69 model assumes that the frequencies of the four bases (A, C, G, T) are equal
- The equilibrium relative frequency of each base is thus 0.25
- Why are they called *equilibrium* frequencies?

Equilibrium Frequencies

Imagine a bottle of perfume has been spilled in room A.

The doors to the other rooms are closed, so the perfume has, thus far, not been able to spread.

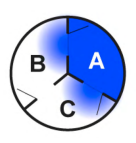
What would happen if we opened all the doors?



Equilibrium Frequencies

If the doors are suddenly opened, the perfume would begin diffusing from the area of highest concentration to lowest.

Molecules of perfume go both ways through open doors, but more pass one way than another, leading to a net flow from room A to rooms B and C.

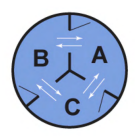


In the instant that the doors are opened, A is losing perfume molecules at *twice the rate* each of the other rooms is gaining molecules. As diffusion progresses, however, the rate of loss from A drops, approaching an equilibrium.

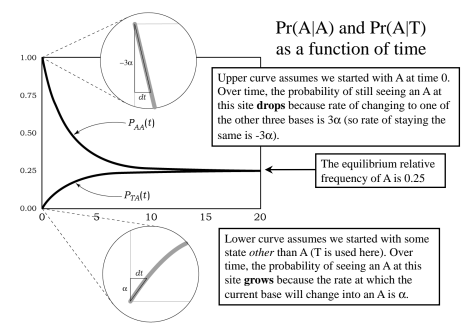
Equilibrium Frequencies

Eventually, all 3 rooms have essentially the same concentration of perfume.

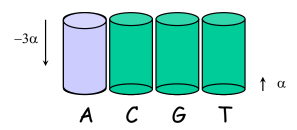
Molecules still move through open doors, but now the rates are the same in all directions.



Back to sequence evolution: assume a sequence began with only A nucleotides (a poly-A sequence). Over time, substitution would begin converting some of these As to Cs, Gs, and Ts, just as the perfume diffused into adjacent rooms.



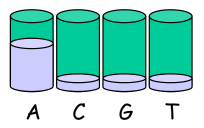
Water analogy (time 0)



- •Start with container A completely full and others empty
- Imagine that all containers are connected by tubes that allow same rate of flow between any two
- Initially, A will be losing water at 3 times the rate that C (or G or T) gains water

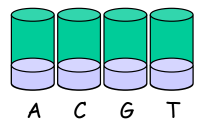
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Water analogy (after some time)



A's level is not dropping as fast now because it is now also *receiving* water from C, G and T

Water analogy (after a very long time)



Eventually, all containers are one fourth full and there is zero *net* volume change – **stationarity** (equilibrium) has been achieved

(Thanks to Kent Holsinger for this analogy)

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JC69 rate matrix

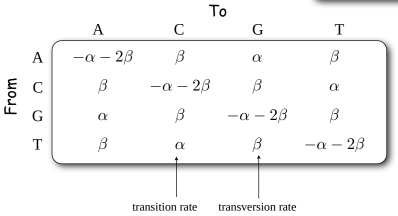
1 parameter: α

		То			
		A	C	G	T
From	A	$\sqrt{-3\alpha}$	α	α	α
	C	α	-3α	lpha	α
	G	$ \alpha $	lpha	-3α	α
	T	α	α	α	-3α

Jukes, T. H., and C. R. Cantor. 1969. Evolution of protein molecules. Pages 21-132 in H. N. Munro (ed.), Mammalian Protein Metabolism. Academic Press, New York.

K80 (or K2P) rate matrix

2 parameters: $\alpha \beta$



Kimura, M. 1980. A simple method for estimating evolutionary rate of base substitutions through comparative studies of nucleotide sequences. Journal of Molecular Evolution 16:111-120.

K80 rate matrix

(looks different, but actually the same)

2 parameters: κ

All I've done is re-parameterize the rate matrix, letting κ equal the *transition/transversion rate ratio*

$$\longrightarrow \kappa = \frac{\alpha}{\beta}$$

Note: the K80 model is identical to the JC69 model if $\kappa = 1$ ($\alpha = \beta$)

Transition/transversion ratio (tratio) versus

Transition/transversion *rate* ratio (kappa)



Cobbler analogy:

- 4 cobblers in a factory make loafers
- 8 cobblers in the factory make work boots
- all cobblers produce the same number of shoes per unit time, regardless of shoe type
- what is the loafer/boot *rate ratio* and how does that compare to the loafer/boot *ratio*?

The loafer/boot *rate ratio* is 1.0 because each cobbler cranks out shoes at the same rate.

The loafer/boot *ratio*, however, is 0.5 because there are twice as many cobblers making boots as there are cobblers making loafers.

There are 8 possible transversion-type substitutions and only 4 possible transition-type substitutions: the transition/transversion ratio is thus 0.5 when the transition/transversion rate ratio is 1.

F81 rate matrix

4 parameters: μ π_A π_C π_G

	A	C	G	T
A	$-\mu(1-\pi_A)$	$\pi_C \mu$	$\pi_G \mu$	$\pi_T \mu$
C	$\pi_A \mu$	$-\mu(1-\pi_C)$	$\pi_G \mu$	$\pi_T \mu$
G	$\pi_A \mu$	$\pi_C \mu$	$-\mu(1-\pi_G)$	$\pi_T \mu$
T	$\pi_A \mu$	$\pi_C \mu$	$\pi_G \mu$	$-\mu(1-\pi_T)$

Note: the F81 model is identical to the JC69 model if all base frequencies are equal

HKY85 rate matrix

	A	C	G	T
Α	_	$\pi_C eta$	$\pi_Geta\kappa$	$\pi_T eta$
C	$\pi_A \beta$	=	$\pi_G eta$	$\pi_Teta\kappa$
G	$\pi_Aeta\kappa$	$\pi_C eta$	=	$\pi_T eta$
T	$\pi_A \beta$	$\pi_C eta \kappa$	$\pi_G eta$	_ /

5 parameters:

 κ β $\pi_{\rm A}$ $\pi_{\rm C}$ $\pi_{\rm G}$

A dash means equal to negative sum of other elements on the same row

Note: the HKY85 model is identical to the F81 model if $\kappa = 1$. If, in addition, all base frequencies are equal, it is identical to JC69.

Hasegawa, M., H. Kishino, and T. Yano. 1985. Dating of the human-ape splitting by a molecular clock of mitochondrial DNA. Journal of Molecular Evolution 21:160-174

F84 vs. HKY85

F84 model:

 μ rate of process generating *all types of substitutions* $k\mu$ rate of process generating *only transitions* Becomes F81 model if k = 0

HKY85 model:

 β rate of process generating *only transversions* $\kappa\beta$ rate of process generating *only transitions* Becomes F81 model if $\kappa = 1$

F84 first used in Felsenstein's PHYLIP package in 1984, first published by: Kishino, H., and M. Hasegawa. 1989. Evaluation of the maximum likelihood estimate of the evolutionary tree topologies from DNA sequence data, and the branching order in hominoidea. Journal of Molecular Evolution 29: 170-179.

GTR rate matrix

9 parameters:

 π_{A} π_{C} π_{G} a b c d e

Identical to the F81 model if a = b = c = d = e = f = 1. If, in addition, all the base frequencies are equal, GTR is identical to JC69. If $a = c = d = f = \beta$ and $b = e = \kappa \beta$, GTR becomes the HKY85 model.

Lanave, C., G. Preparata, C. Saccone, and G. Serio. 1984. A new method for calculating evolutionary substitution rates. Journal of Molecular Evolution 20:86-93.

Rate Heterogeneity

Green Plant rbcL

First 88 amino acids (translation is for Zea mays)

MSPQT- Chara	-ETKASVGFKAGV (green alga; land plant lineage)	KDYKLTYYTPEYETKDTDILAAFRVTP AAAGATTACAGATTACTATACTCCTGAGTATAAACTAAAGATACTGACATTTTAGCTGCATTTCGTGTAACTCCA
Chlorella	(green alga)	CC.TTCCC.ACTC.TAGCAGT
Volvox	(green alga)	TC.TACACGT.GTAC
Conocephalum	(liverwort)	TCTGT
Bazzania	(moss)	TCTGAG.GCGATGAA
Anthoceros	(hornwort)	TCC.TCTCG.GCGTGAG.C.T.AA.GT
Osmunda	(fern)	TCGCAA.GC
Lycopodium	(club "moss")	.GG
Ginkgo	(gymnosperm; Ginkgo biloba)	
Picea	(gymnosperm; spruce)	TAC.GCGTGACAT
Iris	(flowering plant)	
Asplenium	(fern; spleenwort)	TCC.GTCCCACGCCTCGATCGA.GC
Nicotiana	(flowering plant; tobacco)	GAGT
T. A. G. G	M. C. T. G. G. G. A. C. T. G. G. G. T. A. G. G. G. T. G. G. A. C. T. G. G. A. C. G. G. A. C. G.	ACGCGCTCCT
All f	four bases are observed at some sites	while at other sites, only one base is observed

Site-specific rates

Each defined subset (e.g. gene, codon position) has its own relative rate

Subset 1

Subset 2

 r_1 applies to subset 1 (e.g. sites 1 - 1000)

 r_2 applies to subset 2 (e.g. sites 1001-2000)

Relative rates have mean 1:

More generally:

$$\frac{r_1 + r_2}{2} = 1$$

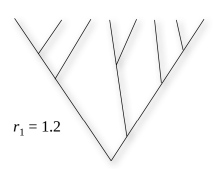
$$r_1 p(r_1) + r_2 p(r_2) = 1$$

62.

Site-specific rates

$$L = \Pr(D_1|r_1) \cdots \Pr(D_{1000}|r_1) \quad \Pr(D_{1001}|r_2) \cdots \Pr(D_{2000}|r_2)$$

Gene 1



Gene 2



 $r_2 = 0.8$

Site-specific rates

JC69 transition probabilities that would be used for every site if rate *homogeneity* were assumed:

$$P_{ii}(t) = \frac{1}{4} + \frac{3}{4}e^{-4\alpha t}$$

$$P_{ij}(t) = \frac{1}{4} - \frac{1}{4}e^{-4\alpha t}$$

Site specific rates

JC69 transition probabilities that would be used for sites in **gene 1**:

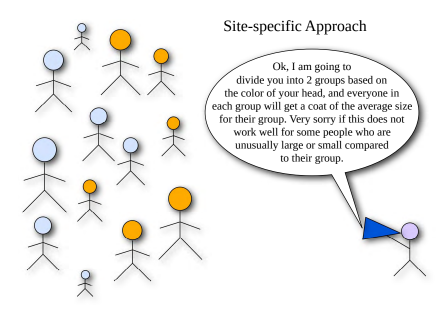
$$P_{ii}(t) = \frac{1}{4} + \frac{3}{4}e^{-4r_1\alpha t}$$

$$P_{ij}(t) = \frac{1}{4} - \frac{1}{4}e^{-4r_1\alpha t}$$

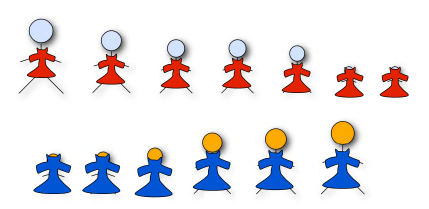
JC69 transition probabilities that would be used for sites in gene 2:

$$P_{ii}(t) = \frac{1}{4} + \frac{3}{4}e^{-4r_2\alpha t}$$

$$P_{ij}(t) = \frac{1}{4} - \frac{1}{4}e^{-4r_2\alpha t}$$



Site-specific Approach

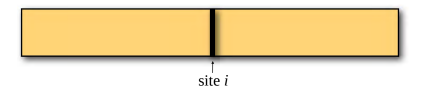


Good: costs less: need to buy just one coat for every person

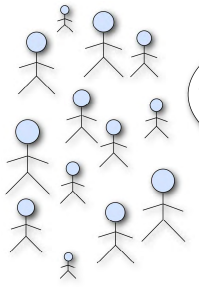
<u>Bad:</u> every person in a group has to wear the same size coat, so the fit will be poor for some people if they are much bigger or smaller than the average size for the group in which they have been placed

Mixture Models

All relative rates applied to every site



$$L_i = \Pr(D_i|r_1) \Pr(r_1) + \Pr(D_i|r_2) \Pr(r_2)$$
Common examples { Invariable sites (I) model Discrete Gamma (G) model}

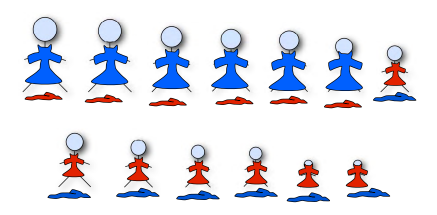


Mixture Model Approach

Ok, I am going to give
each of you 2 coats: use the one
that fits you best and throw away the
other one. This costs twice as much for me,
but on average leads to better fit for you. I
have determined the two sizes of coats
based on the distribution of your
sizes.



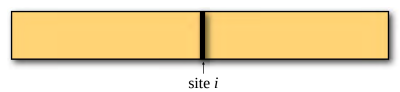
Mixture Model Approach



<u>Good:</u> every person experiences better fit because they can choose the size coat that fits best <u>Bad:</u> costs more because two coats much be provided for each person

Invariable Sites Model

A fraction p_{invar} of sites are assumed to be invariable (i.e. rate = 0.0)



$$L_i = \Pr(D_i|r_1)p_{\text{invar}} + \Pr(D_i|r_2)(1 - p_{\text{invar}})$$

$$r_1 = 0.0$$
 $r_2 = \frac{1}{1 - n}$

Allows for the possibility that any given site could be variable or invariable

Reeves, J. H. 1992. Heterogeneity in the substitution process of amino acid sites of proteins coded for by mitochondrial DNA. Journal of Molecular Evolution 35:17-31.

Invariable sites model

If site *i* is a *constant* site, both terms will contribute to the site likelihood:

$$A \rightarrow A$$

$$L_i = \Pr(D_i|0.0)p_{\text{invar}} + \Pr(D_i|r_2)(1 - p_{\text{invar}})$$

If site *i* is a *variable* site, there is no way to explain the data with a zero rate, so the first term is zero:

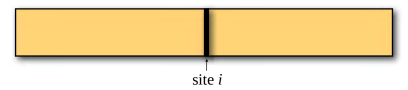
$$A \longrightarrow A$$

$$L_i = \Pr(D_i|\theta.\theta) \widehat{p_{\text{invar}}} + \Pr(D_i|r_2)(1 - p_{\text{invar}})$$

Paul O. Lewis (2017 Woods Hole Workshop in Molecular Evolution)

Discrete Gamma Model

No relative rate is exactly 0.0, and all are equally probable



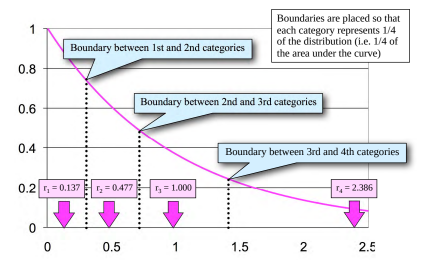
$$L = (\frac{1}{4}) \Pr(D_i|r_1) + (\frac{1}{4}) \Pr(D_i|r_2) + (\frac{1}{4}) \Pr(D_i|r_3) + (\frac{1}{4}) \Pr(D_i|r_4)$$

Relative rates are constrained to a discrete gamma distribution Number of rate categories can vary (4 used here)

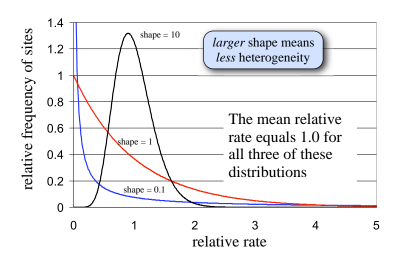
Yang, Z. 1993. Maximum-likelihood estimation of phylogeny from DNA sequences when substitution rates differ over sites. Molecular Biology and Evolution 10:1396-1401.

Yang, Z. 1994. Maximum likelihood phylogenetic estimation from DNA sequences with variable rates over sites: approximate methods. Journal of Molecular Evolution 39:306-314.

Relative rates in 4-category case



Gamma distributions



 $Lab\ on\ ML\ tree\ searching.$