

PAML lab slides:

Review some keynotes

Part 1: PAML Introduction

Part 2: Real data exercises

Review some Keynotes

1. LRT is accurate and powerful, even in small data sets
2. Parameter estimation is more difficult, and can be very sensitive to assumptions
3. Bayes site prediction is very difficult and requires much more data than LRT and parameter estimation
 1. Site prediction is NOT reliable from a few, similar sequences
 2. Site prediction can be sensitive to assumed ω dist.
4. There is an optimal window of sequence divergence
5. Adding lineages is the most efficient way to improve power and accuracy
6. We recommend **robustness analyses**:
 1. Judicious use multiple models (M0-M3, M1a-M2a, M7-M8)
 2. Evaluate sensitivity to model parameters (incl. tree!)
 3. Attempt to identify "consensus sets of sites"
7. Check for and discard results from local optima

PAML (Phylogenetic Analysis by Maximum Likelihood)

A program package by Ziheng Yang
(Demonstration by Joseph Bielawski)

What does PAML do?

Features include:

- estimating synonymous and nonsynonymous rates
- testing hypotheses concerning d_N/d_S rate ratios
- various amino acid-based likelihood analysis
- ancestral sequence reconstruction (DNA, codon, or AAs)
- various clock models
- simulating nucleotide, codon, or AA sequence data sets
- and more

Downloading PAML

PAML download files are at:

<ftp://abacus.gene.ucl.ac.uk/pub/paml/>

For windows, Macs, and Unix/Linux

Programs in the package

baseml	for bases
baseml-g	continuous-gamma for bases
codeml	aaml (for amino acids) & codonml (for codons)
evolver	simulation, tree distances
yn00	d_N and d_S by YN00
chi2	chi square table
pamp	parsimony (Yang and Kumar 1996)
mcmctree	Bayes MCMC tree (Yang & Rannala 1997). Slow

Running PAML programs

1. Sequence data file
2. Tree file
3. Control file (*.ctl)

Running PAML programs: the sequence file

```
4 20
sequence_1 TCATT CTATC TATCG TGATG
sequence_2 TCATT CTATC TATCG TGATG
sequence_3 TCATT CTATC TATCG TGATG
sequence_4 TCATT CTATC TATCG TGATG
```



```
4 20
sequence_1TCATTCTATCTATCGTGATG
sequence_2TCATTCTATCTATCGTGATG
sequence_3TCATTCTATCTATCGTGATG
sequence_4TCATTCTATCTATCGTGATG
```

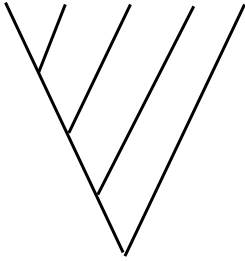


Format = plain text in "PHYLIP" format

Running PAML programs: the tree file

Format = parenthetical notation

```
((((1, 2), 3), 4), 5)
```

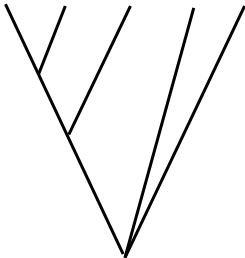


This is a rooted tree

Running PAML programs: the tree file

Format = parenthetical notation

```
(( (1, 2), 3), 4, 5)
```



This is an **unrooted** tree

Running PAML programs: the tree file

Format = parenthetical notation

Examples:

```
(( (1,2), 3), 4, 5);
```

```
(( ((1,2), 3), 4), 5);
```

```
(( (1:0.1, 2:0.2):0.8, 3:0.3):0.7, 4:0.4, 5:0.5);
```

```
(( (Human:0.1, Chimpanzee:0.2):0.8, Gorilla:0.3):0.7,  
Orangutan:0.4, Gibbon:0.5);
```

Running PAML programs: the “*.ctl” file

codeml.ctl

```

seqfile = seqfile.txt * sequence data filename
treefile = tree.txt * tree structure file name
outfile = results.txt * main result file name

noisy = 9 * 0,1,2,3,9: how much rubbish on the screen
verbose = 1 * 1:detailed output
runmode = 0 * 0:user defined tree

seqtype = 1 * 1:codons
CodonFreq = 2 * 0:equal, 1:F1X4, 2:F3X4, 3:F61

model = 0 * 0:one omega ratio for all branches

NSSites = 0 * 0:one omega ratio (M0 in Tables 2 and 4)
* 1:neutral (M1 in Tables 2 and 4)
* 2:selection (M2 in Tables 2 and 4)
* 3:discrete (M3 in Tables 2 and 4)
* 7:beta (M7 in Tables 2 and 4)
* 8:beta&w; (M8 in Tables 2 and 4)

icode = 0 * 0:universal code

fix_kappa = 0 * 1:kappa fixed, 0:kappa to be estimated
kappa = 2 * initial or fixed kappa

fix_omega = 0 * 1:omega fixed, 0:omega to be estimated
omega = 5 * initial omega

*set ncatG for models M3, M7, and M8!!!
*ncatG = 3 * # of site categories for M3 in Table 4
*ncatG = 10 * # of site categories for M7 and M8 in Table 4

```

Exercises:

Maximum Likelihood Methods for Detecting Adaptive Protein Evolution

Joseph P. Bielawski and Ziheng Yang

in

Statistical methods in Molecular Evolution (R. Nielsen, ed.), Springer Verlag
Series in Statistics in Health and Medicine. New York, New York.

Exercises:

	Method/model	program	dataset
1	Pair-wise ML method	codeml	<i>Drosophila GstD1</i>
2	Pair-wise ML method	codeml	<i>Drosophila GstD1</i>
3	M0 and “branch models”	codeml	<i>Ldh</i> gene family
4	M0 and “site models”	codeml	HIV-2 <i>nef</i> genes

Exercise 1: ML estimation of the d_N/d_S (ω) ratio “by hand” for *GstD1*

Dataset: *GstD1* genes of *Drosophila melanogaster* and *D. simulans* (600 codons).

Objective: Use codeml to evaluate the likelihood the *GstD1* sequences for a variety of fixed ω values.

- 1- Plot log-likelihood scores against the values of ω and determine the maximum likelihood estimate of ω .
- 2- Check your finding by running codeml's hill-climbing algorithm.

Exercise 1

Part 2: Real data exercises

```

seqfile = seqfile.txt * sequence data filename
outfile = results.txt * main result file name

noisy = 9 * 0,1,2,3,9: how much rubbish on the screen
verbose = 1 * 1:detailed output
runmode = -2 * -2:pairwise

seqtype = 1 * 1:codons
CodonFreq = 3 * 0:equal, 1:FIX4, 2:F3X4, 3:F61
model = 0 *
NSsites = 0 *
icode = 0 * 0:universal code

fix_kappa = 0 * 1:kappa fixed, 0:kappa to be estimated
kappa = 2 * initial or fixed kappa

fix_omega = 1 * 1:omega fixed, 0:omega to be estimated
omega = 0.001 * 1st fixed omega value [CHANGE THIS]

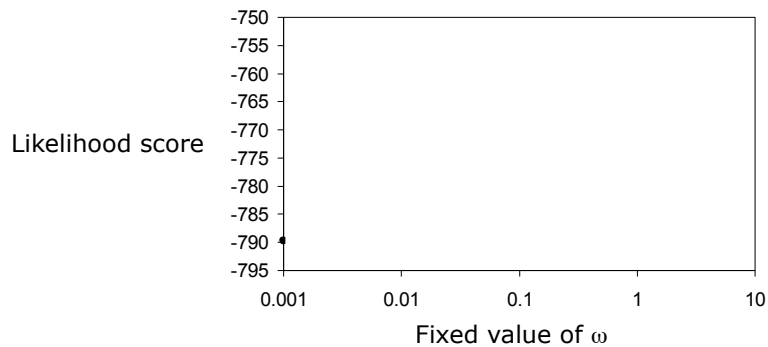
*alternate fixed omega values
*omega = 0.005 * 2nd fixed value
*omega = 0.01 * 3rd fixed value
*omega = 0.05 * 4th fixed value
*omega = 0.10 * 5th fixed value
*omega = 0.20 * 6th fixed value
*omega = 0.40 * 7th fixed value
*omega = 0.80 * 8th fixed value
*omega = 1.60 * 9th fixed value
*omega = 2.00 * 10th fixed value

```

Exercise 1

Part 2: Real data exercises

Plot results: likelihood score vs. omega (log scale)



Exercise 1

If you forget what to do, there is a “step-by-step” guide on the course web site.

There is also a “HelpFile” to help you to get what you need from the output of codeml

Exercise 2: Investigating the sensitivity of the d_N/d_S ratio to assumptions

Dataset: *GstD1* genes of *Drosophila melanogaster* and *D. simulans* (600 codons).

Objective:

- 1- Test effect of transition / transversion ratio (κ)
- 2- Test effect of codon frequencies (π_i 's)
- 3- Determine which assumptions yield the largest and smallest values of S , and what is the effect on ω

"CodonFreq=" is used to specify the equilibrium codon frequencies

Fequal: - 1/61 each for the standard genetic code

- CodonFreq = 0
- number of parameters in the model = 0

F3x4: - calculated from the average nucleotide frequencies at the three codon positions

- CodonFreq = 2
- number of parameters in the model = 9

F61 - also called "ftable"; empirical estimate of each codon frequency

- CodonFreq = 3
- number of parameters in the model = 61

Example: A → C

AAA → CAA

AAA → ACA

AAA → AAC

	Target codon (nucleotide)			NP
	CAA	ACA	AAC	
No bias	1/61	1/61	1/61	0
F1×4	$\pi_C(\pi_A)^2$	$\pi_C(\pi_A)^2$	$\pi_C(\pi_A)^2$	3
MG	π_C^1	π_C^2	π_C^3	9
F3×4 (GY)	$\pi_C^1\pi_A^2\pi_A^3$	$\pi_A^1\pi_C^2\pi_A^3$	$\pi_A^1\pi_A^2\pi_C^3$	9
F61 (GY)	π_{CAA}	π_{ACA}	π_{AAC}	61

Exercise 2

Part 2: Real data exercises

```

seqfile = seqfile.txt * sequence data filename
outfile = results.txt * main result file name

noisy = 9 * 0,1,2,3,9: how much rubbish on the screen
verbose = 1 * 1:detailed output
runmode = -2 * -2:pairwise

seqtype = 1 * 1:codons
CodonFreq = 0 * 0:equal, 1:F1X4, 2:F3X4, 3:F61 [CHANGE THIS]
model = 0 *
NSsites = 0 *
icode = 0 * 0:universal code

fix_kappa = 1 * 1:kappa fixed, 0:kappa to be estimated [CHANGE THIS]
kappa = 1 * fixed or initial value

fix_omega = 0 * 1:omega fixed, 0:omega to be estimated
omega = 0.5 * initial omega value

* Codon bias = none (equal); Ts/Tv bias = none (fixed at 1)
* Codon bias = none (equal); Ts/Tv bias = Yes (estimate by ML)

* Codon bias = yes (F3x4); Ts/Tv bias = none (fixed at 1)
* Codon bias = yes (F3x4); Ts/Tv bias = Yes (estimate by ML)

* Codon bias = yes (F61); Ts/Tv bias = none (fixed at 1)
* Codon bias = yes (F61); Ts/Tv bias = Yes (estimate by ML)

```

Exercise 2

Part 2: Real data exercises

Further details for about the assumptions tested in ACTIVITY 2

Assumption set 1: (Codon bias = none; Ts/Tv bias = none)
CodonFreq=0; kappa=1; fix_kappa=1

Assumption set 2: (Codon bias = none; Ts/Tv bias = Yes)
CodonFreq=0; kappa=1; fix_kappa=0

Assumption set 3: (Codon bias = yes [F3x4]; Ts/Tv bias = none)
CodonFreq=2; kappa=1; fix_kappa=1

Assumption set 4: (Codon bias = yes [F3x4]; Ts/Tv bias = Yes)
CodonFreq=2; kappa=1; fix_kappa=0

Assumption set 5: (Codon bias = yes [F61]; Ts/Tv bias = none)
CodonFreq=3; kappa=1; fix_kappa=1

Assumption set 6: (Codon bias = yes [F61]; Ts/Tv bias = Yes)
CodonFreq=3; kappa=1; fix_kappa=0

Complete this table (If you forget what to do, there is a “step-by-step” guide on the course web-site.)

Table E2: Estimation of d_S and d_N between *Drosophila melanogaster* and *D. simulans* *GstD1* genes

Assumptions	κ	S	N	d_S	d_N	ω	ℓ
Fequal + $\kappa = 1$	1.0	?	?	?	?	?	?
Fequal + $\kappa = \text{estimated}$?	?	?	?	?	?	?
F3x4 + $\kappa = 1$	1.0	?	?	?	?	?	?
F3x4 + $\kappa = \text{estimated}$?	?	?	?	?	?	?
F61 + $\kappa = 1$	1.0	?	?	?	?	?	?
F61 + $\kappa = \text{estimated}$?	?	?	?	?	?	?

κ = transition/transversion rate ratio

S = number of synonymous sites

N = number of nonsynonymous sites

$\omega = d_N/d_S$

ℓ = log likelihood score

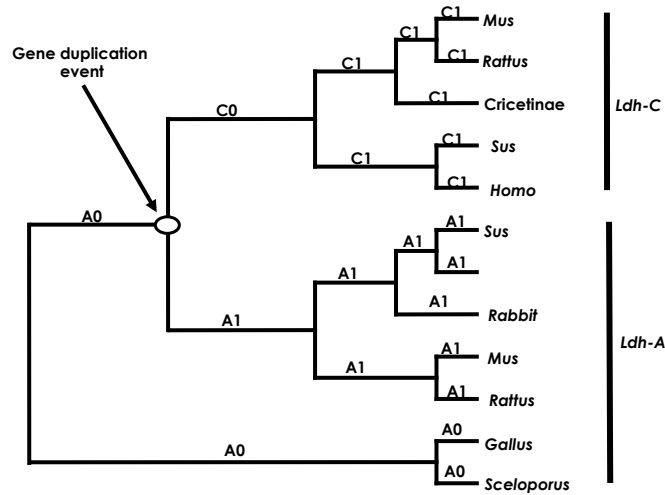
Exercise 3: Test hypotheses about molecular evolution of *Ldh*

Dataset: The *Ldh* gene family is an important model system for molecular evolution of isozyme multigene families. The rate of evolution is known to have increased in *Ldh-C* following the gene duplication event

Objective: Use LRTs to evaluate the following hypotheses:

- 1- The mutation rate of *Ldh-C* has increased relative to *Ldh-A*,
- 2- A burst of positive selection for functional divergence occurred following the duplication event that gave rise to *Ldh-C*
- 3- There was a long term shift in selective constraints following the duplication event that gave rise to *Ldh-C*

Exercise 3



- $H_0: \omega_{A0} = \omega_{A1} = \omega_{C1} = \omega_{C0}$
 $H_1: \omega_{A0} = \omega_{A1} = \omega_{C1} \neq \omega_{C0}$
 $H_2: \omega_{A0} = \omega_{A1} \neq \omega_{C1} = \omega_{C0}$
 $H_3: \omega_{A0} \neq \omega_{A1} \neq \omega_{C1} = \omega_{C0}$

Exercise 3

```

seqfile = seqfile.txt * sequence data filename
treefile = tree.txt * tree structure file name [CHANGE THIS]
outfile = results.txt * main result file name

noisy = 9 * 0,1,2,3,9: how much rubbish on the screen
verbose = 1 * 1:detailed output
runmode = 0 * 0:user defined tree

seqtype = 1 * 1:codons
CodonFreq = 2 * 0:equal, 1:F1X4, 2:F3X4, 3:F61

model = 0 * 0:one omega ratio for all branches
* 1:separate omega for each branch
* 2:user specified dN/dS ratios for branches

NSsites = 0 *

icode = 0 * 0:universal code

fix_kappa = 0 * 1:kappa fixed, 0:kappa to be estimated
kappa = 2 * initial or fixed kappa

fix_omega = 0 * 1:omega fixed, 0:omega to be estimated
omega = 0.2 * initial omega

*H0 in Table 3:
*model = 0
*(X02152Hom,U07178Sus,(M22585rab,((NM017025Rat,U13687Mus),
*((AF070995C,(X04752Mus,U07177Rat)),(U95378Sus,U13680Hom)),(X53828OG1,
* U28410OG2)))));

*H1 in Table 3:
*model = 2
*(X02152Hom,U07178Sus,(M22585rab,((NM017025Rat,U13687Mus),((AF070995C,
*(X04752Mus,U07177Rat)),(U95378Sus,U13680Hom))#1,(X53828OG1,U28410OG2))
* ));

*H2 in Table 3:
*model = 2
*(X02152Hom,U07178Sus,(M22585rab,((NM017025Rat,U13687Mus),((AF070995C
* #1,(X04752Mus #1,U07177Rat #1)#1,(U95378Sus #1,U13680Hom #1)
* #1)#1,(X53828OG1,U28410OG2)))));

*H3 in Table 3:
*model = 2
*(X02152Hom,U07178Sus,(M22585rab,((NM017025Rat,U13687Mus),((AF070995C
* #1,(X04752Mus #1,U07177Rat #1)#1,(U95378Sus #1,U13680Hom #1)
* #1)#1,(X53828OG1 #2,U28410OG2 #2)#2)))));
    
```

Complete this table (If you forget what to do, there is a "step-by-step" guide on the course web-site.)

Table E3: Parameter estimates under models of variable ω ratios among lineages and LRTs of their fit to the *Ldh-A* and *Ldh-C* gene family.

Models	ω_{A0}	ω_{A1}	ω_{C1}	ω_{C0}	ℓ	LRT
H ₀ : $\omega_{A0} = \omega_{A1} = \omega_{C1} = \omega_{C0}$?	$= \omega_{A,0}$	$= \omega_{A,0}$	$= \omega_{A,0}$?	?
H ₁ : $\omega_{A0} = \omega_{A1} = \omega_{C1} \neq \omega_{C0}$?	$= \omega_{A,0}$	$= \omega_{A,0}$?	?	?
H ₂ : $\omega_{A0} = \omega_{A1} \neq \omega_{C1} = \omega_{C0}$?	$= \omega_{A,0}$?	$= \omega_{C,1}$?	?
H ₃ : $\omega_{A0} \neq \omega_{A1} \neq \omega_{C1} = \omega_{C0}$?	?	?	$= \omega_{C,1}$?	?

The topology and branch specific ω ratios are presented in Figure 5.

H₀ v H₁: df = 1

H₀ v H₂: df = 1

H₂ v H₃: df = 1

$\chi_{df=1, \alpha=0.05}^2 = 3.841$

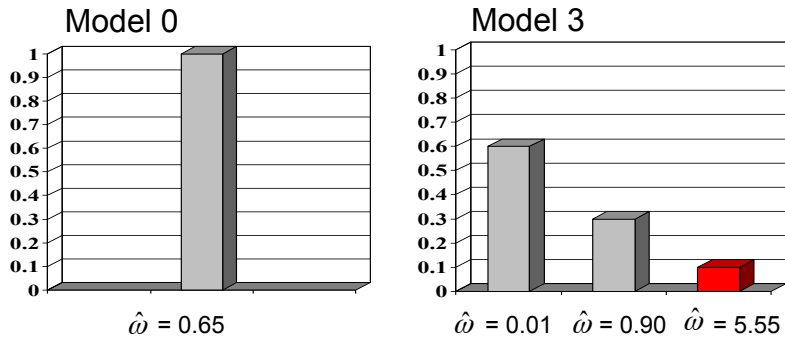
Exercise 4: Testing for adaptive evolution in the *nef* gene of human HIV-2 (Start tonight, but finish as homework)

Dataset: 44 *nef* alleles from a study population of 37 HIV-2 infected people living in Lisbon, Portugal. The *nef* gene in HIV-2 has received less attention than HIV-1, presumably because HIV-2 is associated with reduced virulence and pathogenicity relative to HIV-1

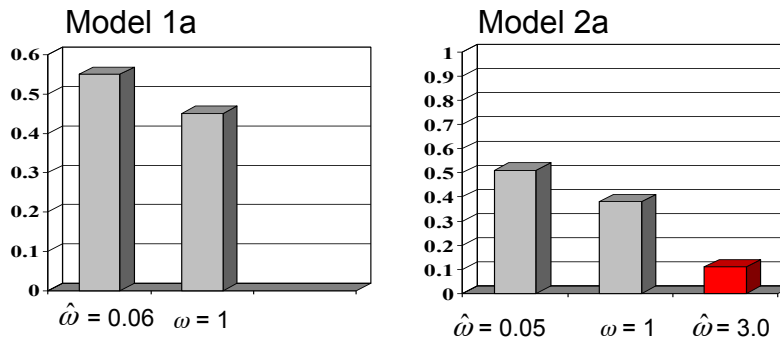
Objectives: 1- Learn to use LRTs to test for sites evolving under positive selection in the *nef* gene.

2- If you find significant evidence for positive selection, then identify the involved sites by using empirical Bayes methods.

H_0 : uniform selective pressure among sites (M0)
 H_1 : variable selective pressure among sites (M3)
 Compare $2\Delta l = 2(l_1 - l_0)$ with a χ^2 distribution



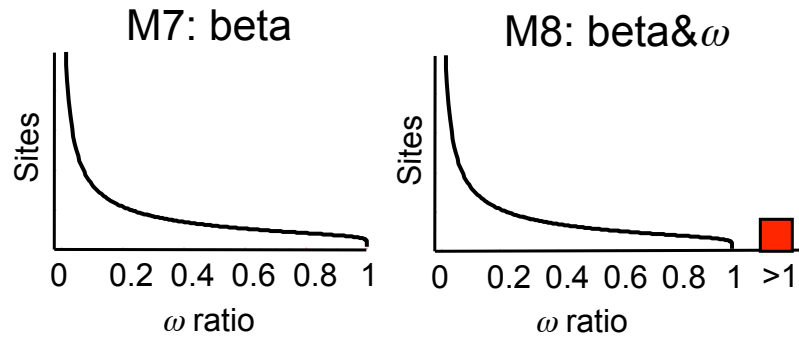
H_0 : variable selective pressure but NO positive selection (M1)
 H_1 : variable selective pressure with positive selection (M2)
 Compare $2\Delta l = 2(l_1 - l_0)$ with a χ^2 distribution



H_0 : Beta distributed variable selective pressure (M7)

H_1 : Beta plus positive selection (M8)

Compare $2\Delta l = 2(l_1 - l_0)$ with a χ^2 distribution



```

seqfile = seqfile.txt          * sequence data filename

* treefile = treefile_M0.txt    * SET THIS for tree file with ML branch lengths under M0
* treefile = treefile_M1.txt    * SET THIS for tree file with ML branch lengths under M1
* treefile = treefile_M2.txt    * SET THIS for tree file with ML branch lengths under M2
* treefile = treefile_M3.txt    * SET THIS for tree file with ML branch lengths under M3
* treefile = treefile_M7.txt    * SET THIS for tree file with ML branch lengths under M7
* treefile = treefile_M8.txt    * SET THIS for tree file with ML branch lengths under M8

outfile = results.txt          * main result file name
noisy = 9                       * lots of rubbish on the screen
verbose = 1                     * detailed output
runmode = 0                     * user defined tree
seqtype = 1                    * codons
CodonFreq = 2                  * F3X4 for codon frequencies
model = 0                      * one omega ratio for all branches

* NSsites = 0                  * SET THIS for M0
* NSsites = 1                  * SET THIS for M1
* NSsites = 2                  * SET THIS for M2
* NSsites = 3                  * SET THIS for M3
* NSsites = 7                  * SET THIS for M7
* NSsites = 8                  * SET THIS for M8

icode = 0                      * universal code
fix_kappa = 1                  * kappa fixed
* kappa = 4.43491              * SET THIS to fix kappa at MLE under M0
* kappa = 4.39117              * SET THIS to fix kappa at MLE under M1
* kappa = 5.08964              * SET THIS to fix kappa at MLE under M2
* kappa = 4.89033              * SET THIS to fix kappa at MLE under M3
* kappa = 4.22750              * SET THIS to fix kappa at MLE under M7
* kappa = 4.87827              * SET THIS to fix kappa at MLE under M8

fix_omega = 0                  * omega to be estimated
omega = 5                      * initial omega

* ncatG = 3                    * SET THIS for 3 site categories under M3
* ncatG = 10                   * SET THIS for 10 of site categories under M7 and M8

fix_branch = 2                 * fixed branch lengths from tree file

```

Complete this table (If you forget what to do, there is a "step-by-step" guide on the course web-site.)

Table E4: Parameter estimates and likelihood scores under models of variable ω ratios among sites for HIV-2 *nef* genes.

Nested model pairs	d_N/d_S^b	Parameter estimates ^c	PSS ^d	ℓ
M0: one-ratio (1) ^a	?	$\omega = ?$	N.A.	?
M3: discrete (5)	?	$p_0 = ?, p_1 = ?, (p_2 = ?)$ $\omega_0 = ?, \omega_1 = ?, \omega_2 = ?$? (?)	?
M1: neutral (1)	?	$p_0 = ?, (p_1 = ?)$ $\omega_0 = ?, (\omega_1 = 1)$	N.A.	?
M2: selection (3)	?	$p_0 = ?, p_1 = ?, (p_2 = ?)$ $\omega_0 = ?, (\omega_1 = 1), \omega_2 = ?$? (?)	?
M7: beta (2)	?	$p = ?, q = ?$	N.A.	?
M8: beta& ω (4)	?	$p_0 = ? (p_1 = ?)$ $p = ?, q = ?, \omega = ?$? (?)	?

^a The number after the model code, in parentheses, is the number of free parameters in the ω distribution.

^b This d_N/d_S ratio is an average over all sites in the HIV-2 *nef* gene alignment.

^c Parameters in parentheses are not free parameters.

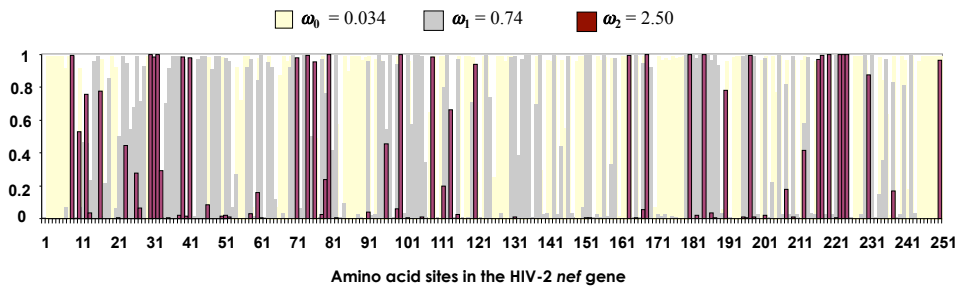
^d PSS is the number of positive selection sites (NEB). The first number is the PSS with posterior probabilities > 50%. The second number (in parentheses) is the PSS with posterior probabilities > 95%.

NOTE: Codeml now implements models M1a and M2a !

Exercise 4

Part 2: Real data exercises

Reproduce this plot



Major weaknesses:

- Poor tree search
- Poor user interface

Major strength:

- Sophisticated likelihood models