Relaxed Phylogenetics and Dating with Confidence

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Introduction

This practical will introduce the BEAST software for Bayesian evolutionary analysis, with a focus on estimating phylogenies and divergence times when you have calibration information from fossil evidence or other prior knowledge.

You will need the following software at your disposal:

- **BEAST** this package contains the BEAST program, BEAUti, TreeAnnotator and other utility programs. This tutorial is written for BEAST v1.6.x, which has support for multiple partitions. It is available for download from http://beast.bio.ed.ac.uk/.
- Tracer this program is used to explore the output of BEAST (and other Bayesian MCMC programs). It graphically and quantitively summarizes the distributions of continuous parameters and provides diagnostic information. At the time of writing, the current version is v1.5. It is available for download from http://beast.bio.ed.ac.uk/.
- FigTree this is an application for displaying and printing molecular phylogenies, in particular those obtained using BEAST. At the time of writing, the current version is v1.3.1. It is available for download from http://tree.bio.ed.ac.uk/.

Rates and dates

This tutorial will guide you through the analysis of an alignment of sequences sampled from twelve primate species. The goal is to estimate the phylogeny as well as the rate of evolution on each lineage based on dates of divergence of their host species.

The first step will be to convert a NEXUS file with a DATA or CHARACTERS block into a BEAST XML input file. This is done using the program BEAUti (this stands for Bayesian Evolutionary Analysis Utility). This is a user-friendly program for setting the evolutionary model and options for the MCMC analysis. The second step is to actually run BEAST using the input file that contains the data, model and settings.

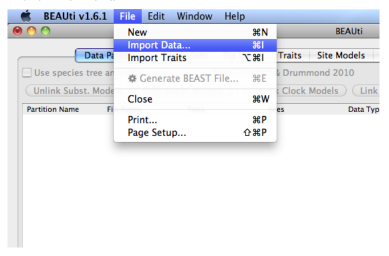
The final step is to explore the output of BEAST in order to diagnose problems and to summarize the results.

BEAUti

The program BEAUti is a user-friendly program for setting the model parameters for BEAST. Run BEAUti by double clicking on its icon.

Loading the NEXUS file

To load a NEXUS format alignment, simply select the Import Alignment... option from the File menu:



Select the file called primates.nex. This file contains an alignment of sequences of 12 species of primates. It looks like this (the lines have been truncated):

#NEXUS

begin data;

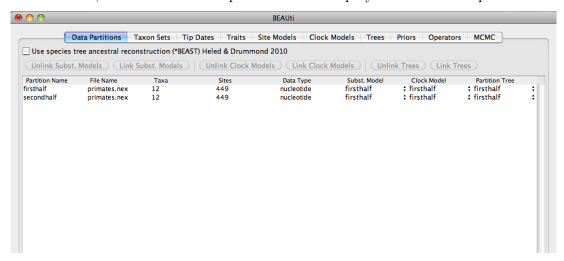
dimensions ntax=12 nchar=898;

format datatype=dna interleave=no gap=-;

matrix

AAGTTTCATTGGAGCCACCACTCTTATAATTGCCCATGGCCTCACC Tarsius_syrichta Lemur_catta AAGCTTCATAGGAGCAACCATTCTAATAATCGCACATGGCCTTACA Homo_sapiens AAGCTTCACCGGCGCAGTCATTCTCATAATCGCCCACGGGCTTACA Pan AAGCTTCACCGGCGCAATTATCCTCATAATCGCCCACGGACTTACA Gorilla AAGCTTCACCGGCGCAGTTGTTCTTATAATTGCCCACGGACTTACA Pongo AAGCTTCACCGGCGCAACCACCCTCATGATTGCCCATGGACTCACA Hylobates AAGCTTTACAGGTGCAACCGTCCTCATAATCGCCCACGGACTAACC Macaca_fuscata AAGCTTTTCCGGCGCAACCATCCTTATGATCGCTCACGGACTCACC M_mulatta AAGCTTTTCTGGCGCAACCATCCTCATGATTGCTCACGGACTCACC M_fascicularis AAGCTTCTCCGGCGCAACCACCCTTATAATCGCCCACGGGCTCACC

Once loaded, the two character partitions are displayed in the main panel:

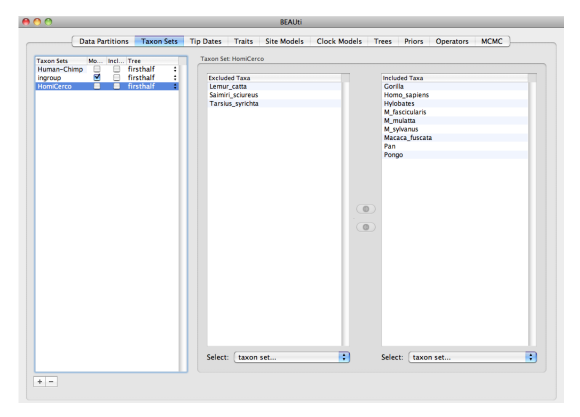


Defining the calibration nodes

Select the **Taxon Sets** tab at the top of the main window. You will see the panel that allows you to create sets of taxa. Once you have created a taxa set you will be able to add calibration information for its most recent common ancestor (MRCA) later on. Press the small "plus" button at the bottom left of the panel. This will create a new taxon set.

Rename it by double-clicking on the entry that appears (it will initially be called untitled1). Call it ingroup (it will contain all taxa except the lemur, which will form the outgroup). In the next table along you will see the available taxa. Select all taxa and press the green arrow button. Move the Lemur back into the excluded taxa set. Since we know that lemur is the outgroup, we will set select the checkbox in the Monophyletic? column. This will ensure that the ingroup is kept monophyletic during the course of the MCMC analysis.

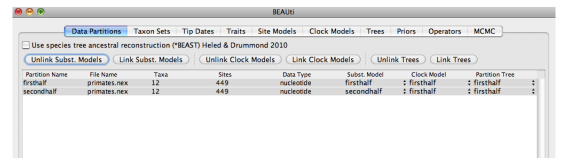
Now repeat the whole procedure creating a set called Human-Chimp that contains only Homo_sapiens and Pan taxa. The screen should look like this:



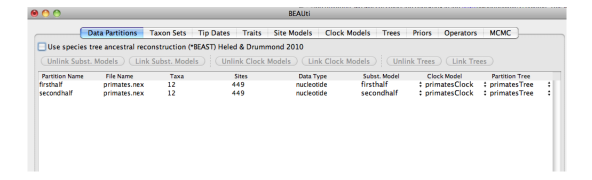
Finally, create a taxon group that contains everything under the hominoid/cercopithecoid split (i.e. everything except Lemur, Saimiri and Tarsius). Call this taxon set something like HomiCerco.

Unlink partition models

At this point we will need to unlink the substitution model so that each parameter is estimated separately for the two partitions. To do this return to **Data Partitions** panel, select both partitions in the table and click the **Unlink Subst Models** button.



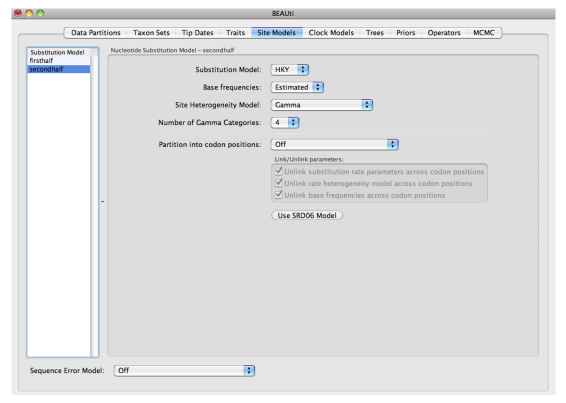
And you can also change the partition model name in its corresponding panel (e.g. **Clock Models** panel), and make the final partitions as illustrated below:



Setting the substitution model

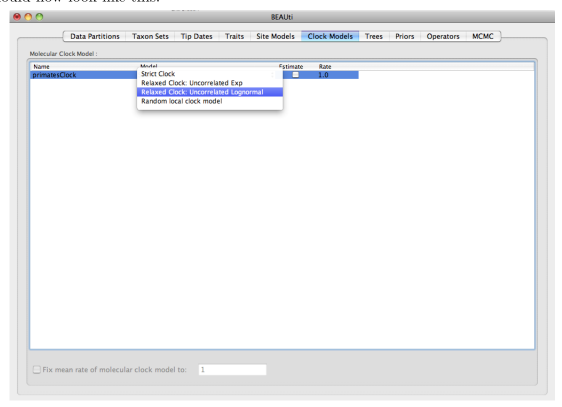
The next thing to do is to click on the **Site Models** tab at the top of the main window. This will reveal the evolutionary model settings for BEAST. Exactly which options appear depend on whether the data are nucleotides, or amino acids, or binary data, or general data. The settings that will appear after loading the Primates data set will be the default values so we need to make some changes.

Most of the models should be familiar to you. For this analysis, we will configure two substitution models by clicking **each of them in turn** on the left side to make the following change: select **Gamma** under the **Site Heterogeneity Model** menu which will allow rate variation across sites in the associated partition. Remember to do this for both partitions.



Setting the clock model

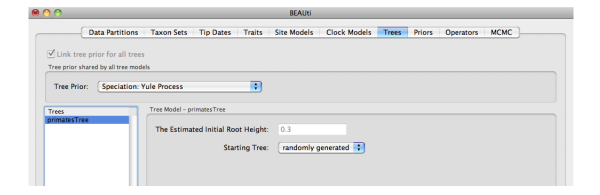
Second, we will do is to click on the **Clock Models** tab at the top of the main window, and to change the molecular clock model to **Relaxed Clock: Uncorrelated Log-normal** so as to account for lineage-specific rate heterogeneity. Your model options should now look like this:



The **Estimate** check box is required to be checked, because we wish to estimate the clock rate (and in doing so the divergence times). But this will be automatically checked, in this case, when we put a proper prior on **tmcra** statistics appeared in **Priors** panel.

Trees

The **Trees** tab allows priors to be specified for each parameter in the model. The first thing to do is to specify that we wish to use the **Yule** model as the tree prior. This is a simple model of speciation that is generally more appropriate when considering sequences from different species. Select this from the **Tree prior** dropdown menu.



Priors

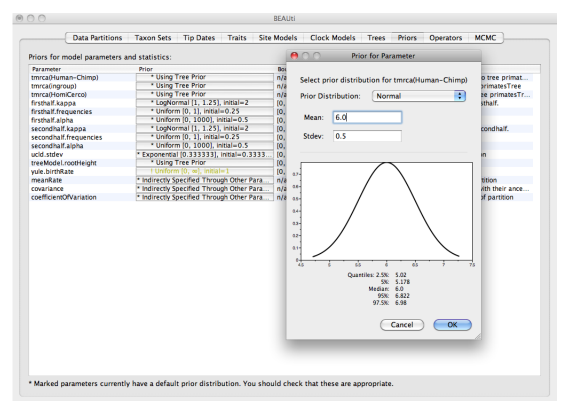
The **Priors** tab allows priors to be specified for each parameter in the model. The first thing to do is to specify that we wish to use the Yule model as the tree prior. This is a simple model of speciation that is generally more appropriate when considering sequences from different species. Select this from the **Tree prior** dropdown menu.

We now need to specify a prior distribution for some of the divergence times, based on our prior fossil knowledge. This is known as calibrating our tree. We will actually use two calibrations in this analysis. Click on the button in the table next to tmrca(human-chimp), A dialog box will appear allowing you to specify a prior for the MRCA of species.

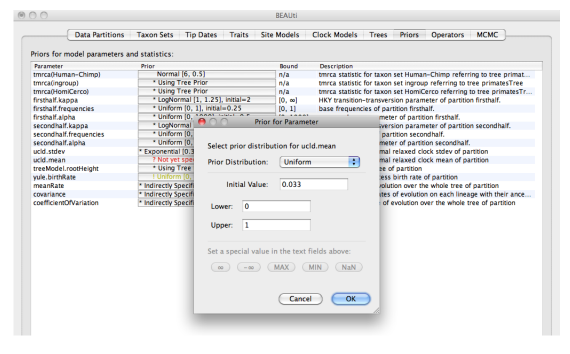
Select the **Normal** distribution. We are going to assume a normal distribution centered at 6 million years with a standard deviation of 0.5 million years. This will give a central 95% range of about 5-7 My. This corresponds to the consensus estimate of the date of the most recent common ancestor of humans and chimps.

Following the same procedure set a calibration of 24 +/-0.5 million (stdev) for the hominoid-cercopithecoid split.

Although we created a taxon set for the ingroup (tmrca(ingroup) in the prior table), we are not going to put an informative prior on this. We can then estimate this divergence time based on the other calibrations.



And the clock model parameters will appear when the clock rate is estimated. The priors table should now look like this:

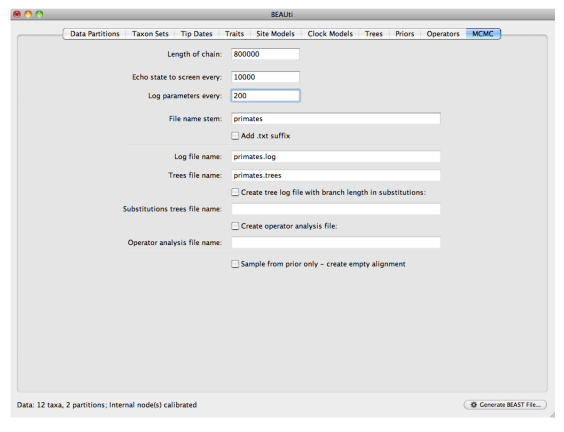


Setting the MCMC options

Ignore the **Operators** tab as this just contains technical settings effecting the efficiency of the MCMC program (see Notes for details).

The next tab, **MCMC**, provides more general settings to control the length of the MCMC and the file names.

Firstly we have the **Length of chain**. This is the number of steps the MCMC will make in the chain before finishing. How long this should be depends on the size of the data set, the complexity of the model and the quality of answer required. The default value of 10,000,000 is entirely arbitrary and should be adjusted according to the size of your data set. For this data set let's initially set the chain length to 800,000 as this will run reasonably quickly on most modern computers (a few minutes).



The next options specify how often the parameter values in the Markov chain should be displayed on the screen and recorded in the log file. The screen output is simply for monitoring the programs progress so can be set to any value (although if set too small, the sheer quantity of information being displayed on the screen will actually slow the program down). For the log file, the value should be set relative to the total length of the chain. Sampling too often will result in very large files with little extra benefit in terms of the precision of the analysis. Sample too infrequently and the log file will not contain much information about the distributions of the parameters. You probably want to aim to store no more than 10,000 samples so this should be set to no less than

chain length / 10000.

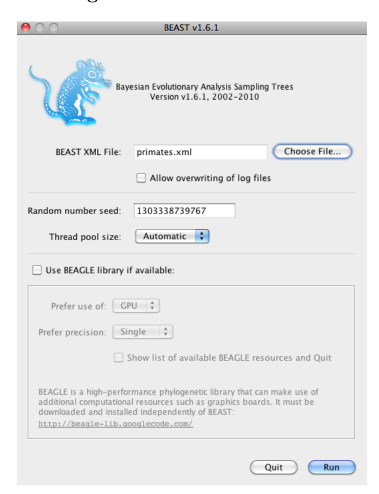
For this exercise we will set the screen log to 10,000 and the file log to 200. The final two options give the file names of the log files for the sampled parameters and the trees. These will be set to a default based on the name of the imported NEXUS file.

• If you are using windows then we suggest you add the suffix .txt to both of these (so, Primates.log.txt and Primates.trees.txt) so that Windows recognizes these as text files.

Generating the BEAST XML file

We are now ready to create the BEAST XML file. To do this, either select the **Generate BEAST File...** option from the **File** menu or click the similarly labelled button at the bottom of the window. Check the default priors, and save the file with an appropriate name (we usually end the filename with .xml, i.e., Primates.xml). We are now ready to run the file through BEAST.

Running BEAST



Now run BEAST and when it asks for an input file, provide your newly created XML file as input. BEAST will then run until it has finished reporting information to the screen. The actual results files are save to the disk in the same location as your input file. The output to the screen will look something like this:

```
BEAST v1.6.1, 2002-2010
       Bayesian Evolutionary Analysis Sampling Trees
                 Designed and developed by
   Alexei J. Drummond, Andrew Rambaut and Marc A. Suchard
               Department of Computer Science
                   University of Auckland
                  alexei@cs.auckland.ac.nz
             Institute of Evolutionary Biology
                  University of Edinburgh
                     a.rambaut@ed.ac.uk
              David Geffen School of Medicine
           University of California, Los Angeles
                     msuchard@ucla.edu
                Downloads, Help & Resources:
                  http://beast.bio.ed.ac.uk
Source code distributed under the GNU Lesser General Public License:
             http://code.google.com/p/beast-mcmc
                     BEAST developers:
Alex Alekseyenko, Erik Bloomquist, Joseph Heled, Sebastian Hoehna,
Philippe Lemey, Wai Lok Sibon Li, Gerton Lunter, Sidney Markowitz,
Vladimir Minin, Michael Defoin Platel, Oliver Pybus, Chieh-Hsi Wu, Walter Xie
     Roald Forsberg, Beth Shapiro and Korbinian Strimmer
Random number seed: 1303338739767
Parsing XML file: primates.xml
 File encoding: MacRoman
Read alignment: alignment
  Sequences = 12
      Sites = 898
   Datatype = nucleotide
Site patterns 'firsthalf.patterns' created from positions 1-449 of alignment 'alignment'
  pattern count = 227
Site patterns 'secondhalf.patterns' created from positions 450-898 of alignment 'alignment'
  pattern count = 231
Using Yule prior on tree
Creating the tree model, 'treeModel'
  initial tree topology = ((((((Gorilla, Tarsius_syrichta), Pan), M_sylvanus), (M_mulatta, Macaca_fuscata)), ((((Homo_sapiens, Saim:
  tree height = 311.73866128365125
Using discretized relaxed clock model.
  over sampling = 1
  parametric model = logNormalDistributionModel
```

Creating state frequencies model: Initial frequencies = {0.25, 0.25, 0.25, 0.25}

Creating state frequencies model: Initial frequencies = {0.25, 0.25, 0.25, 0.25}

rate categories = 22

Creating site model.

Creating HKY substitution model. Initial kappa = 2.0

Creating HKY substitution model. Initial kappa = 2.0

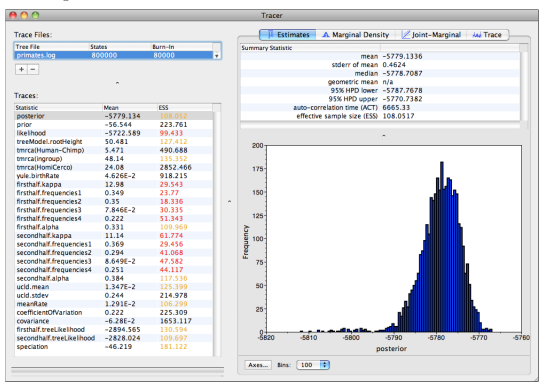
4 category discrete gamma with initial shape = 0.5

```
Creating site model.
  4 category discrete gamma with initial shape = 0.5
TreeLikelihood(treeModel) using native nucleotide likelihood core
  Ignoring ambiguities in tree likelihood.
  With 227 unique site patterns.
Branch rate model used: discretizedBranchRates
TreeLikelihood(treeModel) using native nucleotide likelihood core
  Ignoring ambiguities in tree likelihood.
  With 231 unique site patterns.
Branch rate model used: discretizedBranchRates
Creating swap operator for parameter branchRates.categories (weight=10.0)
Likelihood is using -1 threads.
Creating the MCMC chain:
  chainLength=800000
  autoOptimize=true
  autoOptimize delayed for 8000 steps
# BEAST v1.6.1, Build r3651
# Generated Thu Apr 21 16:27:15 NZST 2011 [seed=1303338739767]
state Posterior
                  Prior
                               Likelihood rootHeight ucld.mean
0 -115044.0238 -105244.7369 -9799.2869 311.739
                                                     3.3E-2
10000 -6019.0942 -48.1460
                             -5970.9482 31.2013
                                                      1.13161E-2
20000 -5932.1777
                  -53.7968
                               -5878.3809
                                            58.3426
                                                         7.80428E-3 0.11 hours/million states
30000 -5864.1836
                  -55.6386
                               -5808.5450 54.7400
                                                        9.7303E-3
                                                                      0.09 hours/million states
40000 -5845.2874
                  -58.9120
                               -5786.3753
                                            82.1531
                                                         6.86446E-3
                                                                      0.08 hours/million states
                                                         7.84354E-3 0.08 hours/million states
50000 -5798.2689
                  -60.2217
                               -5738.0472
                                            87.4058
                  -57.1298
                                                        1.17545E-2 0.07 hours/million states
60000 -5779.7566
                               -5722.6268 49.1015
70000 -5787.4056
                  -60.5876
                                -5726.8180
                                            81.5675
                                                         1.03161E-2 0.07 hours/million states
80000 -5785.8794
                  -58.9768
                                                         9.80333E-3 0.07 hours/million states
                               -5726.9027
                                            70.4040
90000 -5778.6354
                  -59.2618
                                -5719.3737
                                             77.9236
                                                         1.0054E-2
                                                                       0.07 hours/million states
100000 -5780.4106
                  -60.2610
                                -5720.1496
                                            78.7802
                                                           8.80928E-3 0.07 hours/million states
. . . . . .
790000 -5772.8359
                    -59.0540
                                 -5713.7819
                                             77.1089
                                                           9.1281E-3
                                                                        0.06 hours/million states
800000 -5774.2917
                    -57.2419
                                 -5717.0498
                                              61.2605
                                                           9.77329E-3
                                                                       0.06 hours/million states
Operator analysis
                                                 Tuning
                                                                     Time
                                                                              Time/Op Pr(accept) Performance suggestion
Operator
                                                          Count
scale(firsthalf.kappa)
                                                  0.536
                                                          701
                                                                     232
                                                                              0.33
                                                                                       0.2596
                                                                                                   good
firsthalf.frequencies
                                                 0.059
                                                          677
                                                                     206
                                                                              0.3
                                                                                       0.2851
                                                                                                   good
scale(firsthalf.alpha)
                                                 0.598
                                                         719
                                                                     220
                                                                              0.31
                                                                                       0.3255
                                                                                                   good
                                                                                                   good
scale(secondhalf.kappa)
                                                 0.555
                                                         685
                                                                     222
                                                                              0.32
                                                                                       0.3109
                                                 0.058
                                                         661
                                                                                       0.3238
secondhalf.frequencies
                                                                     216
                                                                              0.33
                                                                                                   good
scale(secondhalf.alpha)
                                                 0.598
                                                          680
                                                                     210
                                                                              0.31
                                                                                       0.3015
                                                                                                   good
scale(ucld.mean)
                                                 0.697
                                                         21192
                                                                     7410
                                                                              0.35
                                                                                       0.2786
                                                                                                   good
scale(ucld.stdev)
                                                 0.274
                                                         21350
                                                                     7342
                                                                              0.34
                                                                                       0.3526
                                                                                                   good
                                                                                                   good
subtreeSlide(treeModel)
                                                 4.405
                                                         105795
                                                                     18763
                                                                              0.18
                                                                                       0.337
Narrow Exchange(treeModel)
                                                          106685
                                                                     19150
                                                                              0.18
                                                                                       0.0004
                                                                                                   very low
Wide Exchange(treeModel)
                                                          21301
                                                                     2071
                                                                                       0.0
                                                                                                   very low
                                                                              0.1
                                                                                                   very low
wilsonBalding(treeModel)
                                                                              0.16
                                                          21229
                                                                     3431
                                                                                       0.0
scale(treeModel.rootHeight)
                                                 0.853
                                                         21224
                                                                     1334
                                                                              0.06
                                                                                       0.218
                                                                                                   good
uniform(nodeHeights(treeModel))
                                                          213114
                                                                     43645
                                                                              0.2
                                                                                       0.222
                                                                                                   good
scale(yule.birthRate)
                                                 0.268
                                                          21594
                                                                     697
                                                                              0.03
                                                                                       0.2761
                                                                                                   good
up:ucld.mean down:nodeHeights(treeModel)
                                                 0.596
                                                         21448
                                                                    7551
                                                                              0.35
                                                                                       0.2417
                                                                                                   slightly high
Try setting scaleFactor to about 0.586
                                                          70928
                                                                     16433
                                                                                       0.6553
swapOperator(branchRates.categories)
                                                                              0.23
                                                                                                   high No suggestions
randomWalkInteger(branchRates.categories)
                                                          70818
                                                                     13509
                                                                              0.19
                                                                                       0.9432
                                                                                                   very high
Try increasing windowSize to about 2.0
uniformInteger(branchRates.categories)
                                                          71199
                                                                     14008
                                                                              0.2
                                                                                       0.7518
                                                                                                   high
```

^{3.028016666666667} minutes

Analyzing the results

Run the program called **Tracer** to analyze the output of BEAST. When the main window has opened, choose **Import Trace File...** from the **File** menu and select the file that BEAST has created called **Primates.log**. You should now see a window like the following:



Remember that MCMC is a stochastic algorithm so the actual numbers will not be exactly the same.

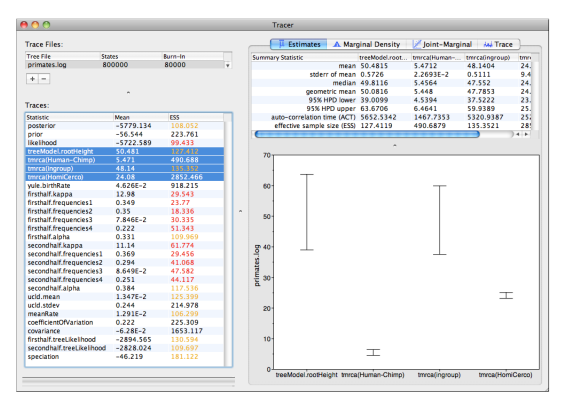
On the left hand side is a list of the different quantities that BEAST has logged. There are traces for the posterior (this is the log of the product of the tree likelihood and the prior probabilities), and the continuous parameters. Selecting a trace on the left brings up analyses for this trace on the right hand side depending on tab that is selected. When first opened, the 'posterior' trace is selected and various statistics of this trace are shown under the Estimates tab. In the top right of the window is a table of calculated statistics for the selected trace.

Select meanRate to look at the rate of evolution averaged over the whole tree. Tracer will plot a (marginal posterior) distribution for the selected parameter and also give you statistics such as the mean and median. The 95% HPD stands for highest posterior density interval and represents the most compact interval on the selected parameter that contains 95% of the posterior probability. It can be thought of as a Bayesian analog to a confidence interval.

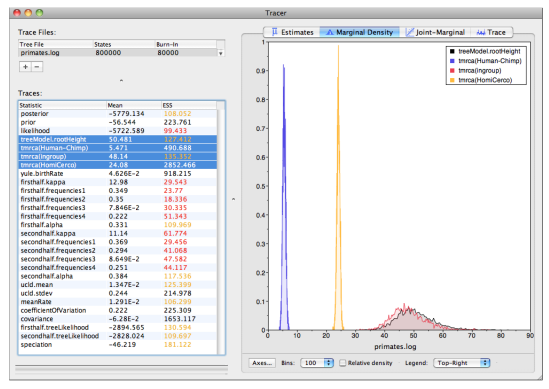
Questions

	ne rate of molecular					,
What	sources of error does	s this estima	te include?			
	pefficientOfVaria varies from lineage t		-			
D 4	ha mata of accolation	difference beta	m ti alla, ama an	ant difference	+ lim == = i ==	the too
Does t	he rate of evolution	differ substa	ntially amor	ngst differen	t lineages in	the tre
Does t	he rate of evolution	differ substa	ntially amor	ngst differen	t lineages in	the tre
Does t	he rate of evolution	differ substa	ntially amor	ngst differen	t lineages in	the tre
Selecti	ng the treeModel.	rootHeight	parameter			

Select the treeModel.rootHeight parameter and the next three (hold shift whilst selecting). This will show a display of the age of the root and the three MRCAs we specified in BEAUti. The parameter that we used to calibrate the tree (tmrca(human-chimp)) will have posterior distributions very similar to the prior distributions that we specified.

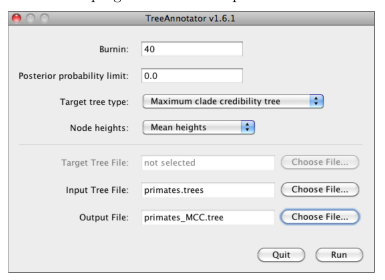


If you switch the tab at the top of the window to **Marginal Density** then you will get a plot of the marginal posterior densities of each of these date estimates overlayed:



Obtaining an estimate of the phylogenetic tree

BEAST also produces a sample of plausible trees along with its sample of parameter estimates. These need to be summarized using the program **TreeAnnotator** (see Notes for details). This will take the set of trees and find the best supported one. It will then annotate this summary tree with the mean ages of all the nodes and the HPD ranges. It will also calculate the posterior clade probability for each node. Run the TreeAnnotator program and set it up to look like this:



The burnin is the number of trees to remove from the start of the sample. Unlike **Tracer** which specifies the number of steps as a burnin, in **TreeAnnotator** you need to specify the actual number of trees. For this run, you specified a chain length of 800,000 steps sampling every 200 steps. Thus the trees file will contain 4000 trees and so to specify a 1% burnin use the value 40.

The **Posterior probability limit** option specifies a limit such that if a node is found at less than this frequency in the sample of trees (i.e., has a posterior probability less than this limit), it will not be annotated. The default of 0.5 means that only nodes seen in the majority of trees will be annotated. Set this to zero to annotate all nodes.

For **Target tree type** you can either choose a specific tree from a file or ask TreeAnnotator to find a tree in your sample. The default option, **Maximum clade credibility tree**, finds the tree with the highest product of the posterior probability of all its nodes.

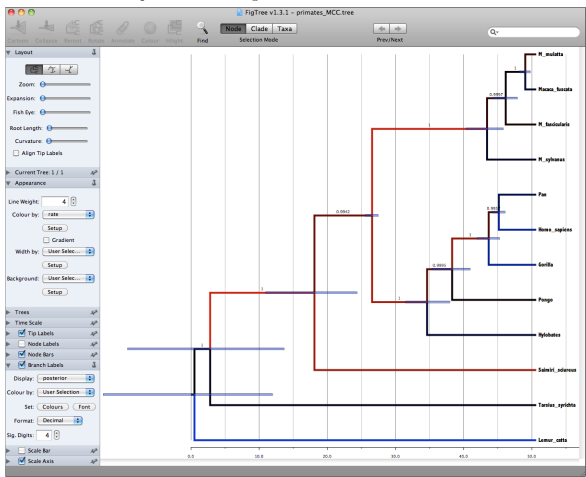
Choose **Mean heights** for node heights. This sets the heights (ages) of each node in the tree to the mean height across the entire sample of trees for that clade.

For the input file, select the trees file that BEAST created (by default this will be called Primates.trees) and select a file for the output (here we called it Primates_MCC.tree).

Now press Run and wait for the program to finish.

Viewing the Tree

Finally, we can look at the tree in another program called **FigTree**. Run this program, and open the Primates.MCC.tree file by using the Open command in the File menu. The tree should appear. You can now try selecting some of the options in the control panel on the left. Try selecting **Node Bars** to get node age error bars. Also turn on **Branch Labels** and select **posterior** to get it to display the posterior probability for each node. Under **Appearance** you can also tell FigTree to colour the branches by the rate. You should end up with something like this:



Which branch has the fastest rate of evolution and what is the estimated rate?

Which branch has the slowest rate of evolution and what is the estimated rate?)

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Comparing your results to the prior

Using BEAUti, set up the same analysis but under the MCMC options, select the **Sample from prior only** option. This will allow you to visualize the full prior distribution in the absence of your sequence data. Summarize the trees from the full prior distribution and compare the summary to the posterior summary tree.