# Divergence Dating Tutorial with BEAST 2.2.x

Alexei Drummond, Andrew Rambaut, Remco Bouckaert, and Walter Xie January 29, 2015

#### 1 Introduction

This tutorial introduces the BEAST software for Bayesian evolutionary analysis through a simple tutorial. The tutorial involves co-estimation of a gene phylogeny and associated divergence times in the presence of calibration information from fossil evidence.

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 v2.2.x, which has support for multiple partitions. It is available for download from http://www.beast2.org/.
- 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 v1.6. It is available for download from

http://tree.bio.ed.ac.uk/software/.

• **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 v1.4.2. It is available for download from http://tree.bio.ed.ac.uk/software/.

This tutorial will guide you through the analysis of an alignment of sequences sampled from twelve primate species (see Figure 1). The goal is to

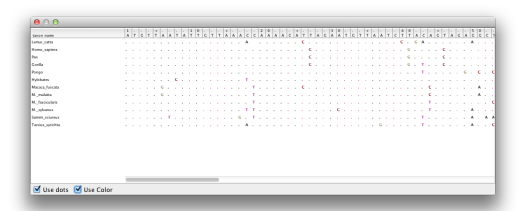


Figure 1: Part of the alignment for primates.

estimate the phylogeny, the rate of evolution on each lineage and the ages of the uncalibrated ancestral divergences.

The first step will be to convert a NEXUS file with a DATA or CHARAC-TERS block into a BEAST XML input file. This is done using the program BEAUti (which 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 generated by BEAUTi, which contains the data, model and analysis settings. The final step is to explore the output of BEAST in order to diagnose problems and to summarize the results.

## 2 BEAUti

The program BEAUti is a user-friendly program for setting the model parameters for BEAST. Run BEAUti by double clicking on its icon. Once running, BEAUti will look similar irrespective of which computer system it is running on. For this tutorial, the Mac OS X version is used in the figures but the Linux and Windows versions will have the same layout and functionality.

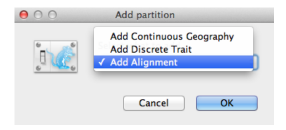


Figure 2: Add Partition window (Only appear if related packages are installed).

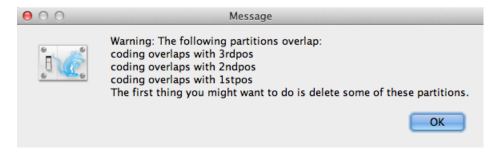


Figure 3: Warning message window (Only appear if there is any coding overlaps in the partitions).

## 2.1 Loading the NEXUS file

To load a NEXUS format alignment, simply select the Import Alignment... option from the File menu, or drag the file into the middle of Partitions panel.

The example file called primate-mtDNA.nex is available from the examples/nexus/directory for Mac and Linux and examples/nexus/for Windows inside the directory where BEAST was installed. This file contains an alignment of sequences of 12 species of primates.

An Add Partition window (Figure 2) would pop up if the related package is installed. If you are using "pure" BEAST 2, you can go to the next paragraph. Otherwise, select Add Alignment and click **OK** to continue.

If there is any coding overlaps in the partitions, the warning message window (Figure 3) will appear. Read and click **OK** to continue.

Once loaded, five character partitions are displayed in the main panel (Figure 4). The alignment is divided into a protein coding part and a non-coding part, and the coding part is divided in codon positions 1, 2 and 3.

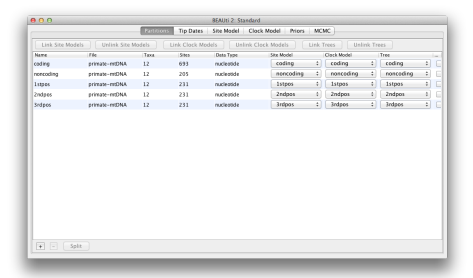


Figure 4: A screenshot of the data tab in BEAUti. This and all following screenshots were taken on an Apple computer running Mac OS X and will look slightly different on other operating systems.

You must remove the 'coding' partition before continuing to the next step as it refers to the same nucleotides as partitions '1stpos', '2ndpos' and '3rdpos'. To remove the 'coding' partition select the row and click the '-' button at the bottom of the table. You can view the alignment by double clicking the partition.

## Link/Unlink partition models

Since the sequences are linked (i.e. they are all from the mitochondrial genome which is not believed to undergo recombination in birds and mammals) they share the same ancestry, so the partitions should share the same time-tree in the model. For the sake of simplicity, we will also assume the partitions share the same evolutionary rate for each branch, and hence the same "clock model". We will restrict our modelling of rate heterogeneity to among-site heterogeneity within each partition, and also allow the partitions to have different mean rates of evolution.

So, at this point we will need to link the clock model and tree. In the

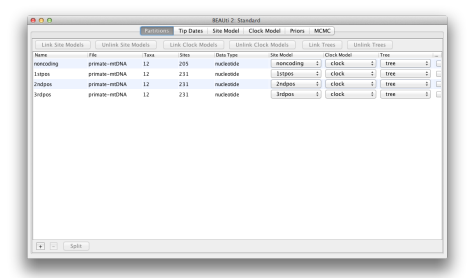


Figure 5: A screenshot of the Partitions tab in BEAUti after linking and renaming the clock model and tree.

Partitions panel, select all four partitions in the table (or none, by default all partitions are affected) and click the Link Trees button and then the Link Clock Models button (see Figure 5). Then click on the first drop-down menu in the Clock Model column and rename the shared clock model to 'clock'. Likewise rename the shared tree to 'tree'. This will make following options and generated log files more easy to read.

## 2.2 Setting the substitution model

The next step is to set up the substitution model. Then, select the **Site Models** tab at the top of the main window (we skip the **Tip Dates** tab since all taxa are from contemporary samples). This will reveal the evolutionary model settings for BEAST. The options available depend on whether the data are nucleotides, or amino acids, binary data, or general data. The settings that will appear after loading the primate nucleotide alignment will be the default values for nucleotide data so we need to make some changes.

Most of the models should be familiar to you. First, set the **Gamma** Category Count to 4 and then tick the 'estimate' box for the **Shape** pa-

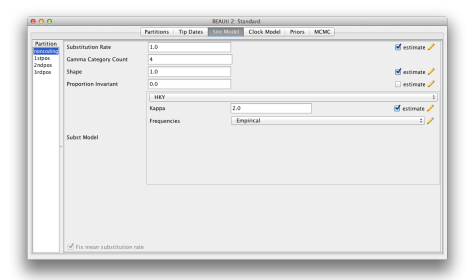


Figure 6: A screenshot of the site model tab in BEAUti.

rameter. This will allow rate variation between sites in each partition to be modelled. Note that 4 to 6 categories works sufficiently well for most data sets, while having more categories takes more time to compute for little added benefit. We leave the Proportion Invariant entry set to zero.

Then select **HKY** from the **Subst Model** drop-down menu. Ideally, a substitution model should be selected that fit the data best for each partition, but here for the sake of simplicity we use HKY for all partitions. Further, select **Empirical** from the **Frequencies** drop-down menu. This will fix the frequencies to the proportions observed in the data (for each partition individually, once we unlink the site models). This approach means that we can get a good fit to the data without explicitly estimating these parameters. We do it here simply to make the log files a bit shorter and more readable in later parts of the exercise.

Finally check the 'estimate' box for the **Substitution rate** parameter and select the **Fix mean mutation rate** check box. This will allow the individual partitions to have their relative rates estimated for unlinked the site models (Figure 6).

At last, hold 'shift' key to select all site models on the left side, and click **OK** to clone the setting from **noncoding** into **1stpos**, **2ndpos** and **3rdpos** 

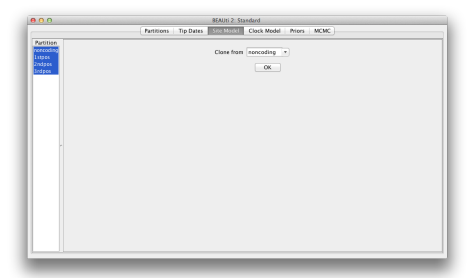


Figure 7: clone configuration from one site model to others.

(Figure 7). Go through each site model, as you can see, their configurations are same now.

#### 2.3 Setting the clock model

The next step is to select the **Clock Models** tab at the top of the main window. This is where we select the molecular clock model. For this exercise we are going to leave the selection at the *default* value of a **strict molecular clock**, because this data is very clock-like, and does not need rate variation among branches to be included in the model.

To test for clock-likeness, you can (i) run the analysis with a relaxed clock model and check how much variation among rates are implied by the data (see coefficient of variation for more on this), or (ii) perform a model comparison between a strict and relaxed clock using path sampling, or (iii) use a random local clock model [2] which explicitly considers whether each branch in the tree needs its own branch rate.

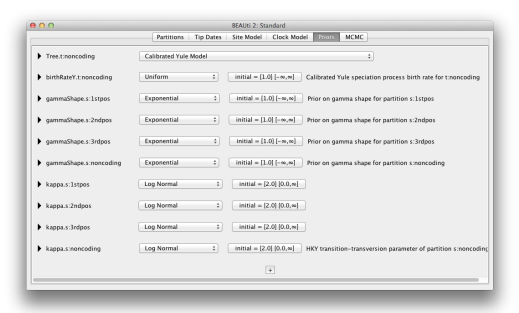


Figure 8: A screenshot of the Priors tab in BEAUti.

#### 2.4 Priors

The **Priors** tab allows priors to be specified for each parameter in the model. The model selections made in the site model and clock model tabs, result in the inclusion of various parameters in the model, and these are shown in the priors tab (see Figure 8).

Here we also specify that we wish to use the Calibrated Yule model [4] as the tree prior. The Yule model is a simple model of speciation that is generally more appropriate when considering sequences from different species. Select the Calibrated Yule Model from the **Tree prior** dropdown menu.

#### 2.4.1 Defining the calibration node

We now need to specify a prior distribution on the calibrated node, based on our fossil knowledge. This is known as calibrating our tree. To define an extra prior, press the small + button below list of priors. If it is not visible in your view, please scroll down the panel to the bottom to find + button.

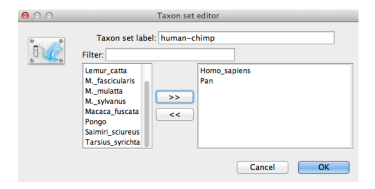


Figure 9: Taxon set editor in BEAUti.

You will see a dialog that allows you to define a subset of the taxa in the phylogenetic tree. Once you have created a taxa set you will be able to add calibration information for its most recent common ancestor (MRCA) later on.

Name the taxa set by filling in the taxon set label entry. Call it human-chimp, since it will contain the taxa for *Homo sapiens* and *Pan*. In the list below you will see the available taxa. Select each of the two taxa in turn and press the >> arrow button. (Figure 9). Click OK and the newly defined taxa set will be added in to the prior list. As this is a calibrated node to be used in conjunction with the Calibrated Yule prior, monophyly must be enforced, so select the checkbox marked Monophyletic. This will constrain the tree topology so that the human-chimp grouping is kept monophyletic during the course of the MCMC analysis.

To encode the calibration information we need to specify a distribution for the MRCA of human-chimp. Select the **Log-normal** distribution from the drop down menu to the right of the newly added human-chimp.prior. Click on the black triangle and a graph of the probability density function will appear, along with parameters for the log normal distribution. We are going to set M=1.78 and S=0.085 which will specify a distribution centred at about 6 million years with a standard deviation of about 0.5 million years. This will give a central 95% probability range covering 5-7 Mya. This roughly corresponds to the current consensus estimate of the date of the most recent common ancestor of humans and chimpanzees (Figure 10).

We should convince ourselves that the priors shown in the priors panel really reflect the prior information we have about the parameters of the

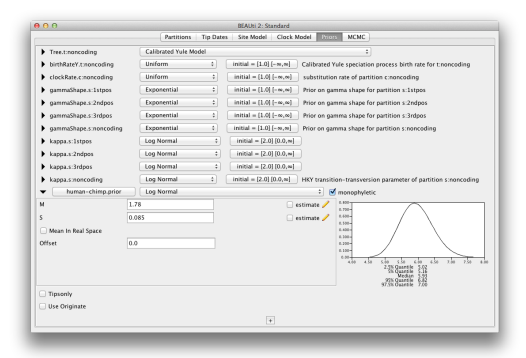


Figure 10: A screenshot of the calibration prior options in the Priors panel in BEAUti.

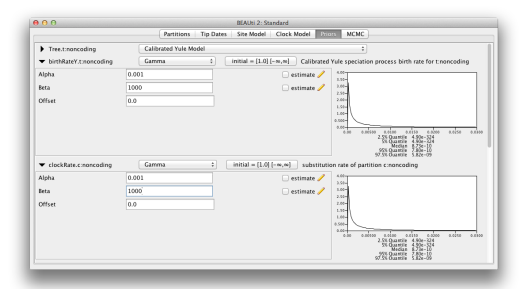


Figure 11: Gamma prior.

model. Finally we will also specify some diffuse "uninformative" but proper priors on the overall molecular clock rate (clockRate) and the speciation rate (birthRateY) of the Yule tree prior. For each of them, select Gamma from the drop-down menu and using the arrow button, expand the view to reveal the parameters of the Gamma prior. For both the clock rate and the Yule birth rate set the Alpha (shape) parameter to 0.001 and the Beta (scale) parameter to 1000 (Figure 11).

By default each of the gamma shape parameters has an exponential prior distribution with a mean of 1. This implies (see Figure 3.7) we expect some rate variation. By default the kappa parameters for the HKY model have a log normal(1,1.25) prior distribution, which broadly agrees with empirical evidence [6] on the range of realistic values for transition/transversion bias. These default priors are kept since they are suitable for this particular analysis.

#### 2.5 Setting the MCMC options

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

Firstly we have the **Chain Length**. 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 set the **chain length** to 6,000,000 as this will run reasonably quickly on most modern computers (a few minutes).

The **Store Every** field determines how often the state is stored to file. Storing the state periodically is useful for situations where the computing environment is not very reliable and a BEAST run can be interrupted. Having a stored copy of the recent state allows you to resume the chain instead of restarting from the beginning, so you do not need to get through burn-in again. The **Pre Burnin** field specifies the number of samples that are not logged at the very beginning of the analysis. We leave the **Store Every** and **Pre Burnin** fields set to their default values. Below these are the details of the log files. Each one can be expanded by clicking the black triangle.

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 accuracy of the analysis. Sample too infrequently and the log file will not record sufficient 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 / 10,000.

For this exercise we will set the trace  $\log$  and the tree  $\log$  frequency to 1,000 and the screen  $\log$  to 10,000. Also specify Primates.log as the file name of the trace  $\log$  file and Primates.trees as the file name of the tree  $\log$  file. Make sure that the file name filed of the screen  $\log$  is left empty, or the screen  $\log$  will not be written to the screen.

• If you are using the Windows operating system 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.

### 2.6 Generating the BEAST XML file

We are now ready to create the BEAST XML file. To do this, select the **Save** option from the **File** menu. 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.

## 3 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 v2.2.0, 2002-2014
Bayesian Evolutionary Analysis Sampling Trees
Designed and developed by
Remco Bouckaert, Alexei J. Drummond, Andrew Rambaut and Marc A. Suchard

Department of Computer Science University of Auckland remco@cs.auckland.ac.nz 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://beast2.org/

Source code distributed under the GNU Lesser General Public License:  ${\tt http://github.com/CompEvol/beast2}$ 

BEAST developers:
Alex Alekseyenko, Trevor Bedford, Erik Bloomquist, Joseph Heled,
Sebastian Hoehna, Denise Kuehnert, Philippe Lemey, Wai Lok Sibon Li,
Gerton Lunter, Sidney Markowitz, Vladimir Minin, Michael Defoin Platel,
Oliver Pybus, Chieh-Hsi Wu, Walter Xie

Thanks to: Roald Forsberg, Beth Shapiro and Korbinian Strimmer

Random number seed: 777

12 taxa 898 sites 413 patterns

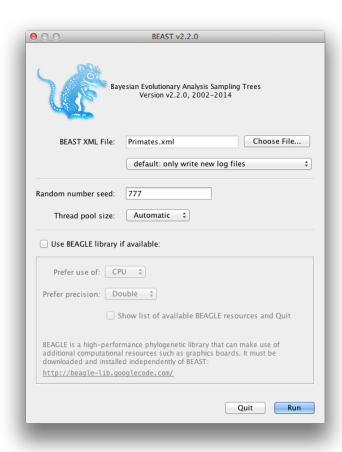


Figure 12: A screenshot of BEAST.

```
TreeLikelihood uses beast.evolution.likelihood.BeerLikelihoodCore4
TreeLikelihood uses beast.evolution.likelihood.BeerLikelihoodCore4
TreeLikelihood uses beast.evolution.likelihood.BeerLikelihoodCore4
TreeLikelihood uses beast.evolution.likelihood.BeerLikelihoodCore4
```

Bouckaert RR, Heled J, Kuehnert D, Vaughan TG, Wu C-H, Xie D, Suchard MA, Rambaut A, Drummond AJ (2014) BEAST 2: A software platform for Bayesian evolutionary analysis. PLoS Computational Biology 10(4): e1003537

Heled J, Drummond AJ (2012) Calibrated Tree Priors for Relaxed Phylogenetics and Divergence Time Estimation. Systematic Biology 61(1):138-149.

Hasegawa M, Kishino H, Yano T (1985) Dating the human-ape splitting by a molecular clock of mitochondrial DNA. Journal of Molecular Evolution 22:160-174.

Writing file /Prim	nates log							
Writing file /Prin								
Sample		SS(posterior)	likelihood		prior			
0	-7924.3599	N	-7688.4922	-23	35.8676			
10000	-5529.0700	2.0	-5459.1993	-6	39.8706			
20000	-5516.8159	3.0	-5442.3372	-7	74.4786			
30000	-5516.4959	4.0	-5439.0839	-7	77.4119			
40000	-5521.1160	5.0	-5445.6047	-75.5113				
50000	-5520.7350	6.0	-5444.6198	-76.1151				
60000	-5512.9427	7.0	-5439.2561	-73.6866 2m39s/Msamples				
70000	-5513.8357	8.0	-5437.9432	-7	75.8924 2m3	39s/Msample	es	
5990000	-5516.6832	474.6	-5442.5945					
6000000	-5512.3802	472.2	-5440.8928	-7	71.4874 2m4	10s/Msample	es	
Operator				Tuning	#accept	#reject		prob.acc
ScaleOperator(treeScaler.t:tree)				0.703	39935	174155	214090	0.187
ScaleOperator(treeRootScaler.t:tree)				0.644	37329	177166	214495	0.174
Uniform(UniformOperator.t:tree)					479419	1668915	2148334	0.223
SubtreeSlide(SubtreeSlide.t:tree)				9.922	272787	801404	1074191	0.254
Exchange(narrow.t:tree)					744	1074261	1075005	0.001
Exchange(wide.t:tree)					9	214594	214603	0.000
WilsonBalding(WilsonBalding.t:tree)				0.050	4	214548 5375	214552 7114	0.000
ScaleOperator(KappaScaler.s:noncoding)				0.352	1739			0.244
DeltaExchangeOperator(FixMeanMutationRatesOperator)				0.425 0.375	17277 1729	126203 5428	143480 7157	0.120 0.242
ScaleOperator(gammaShapeScaler.s:noncoding)				0.375	58005	5428 156128	214133	0.242
ScaleOperator(CalibratedYuleBirthRateScaler.t:tree)				0.706	50080	164952	214133	0.271
ScaleOperator(StrictClockRateScaler.c:clock) UpDownOperator(strictClockUpDownOperator.c:clock)				0.706	50809	163882	214691	0.233
ScaleOperator(KappaScaler.s:1stpos)				0.369	1816	5388	7204	0.252
ScaleOperator(gammaShapeScaler.s:1stpos)				0.44	1927	5129	7056	0.232
ScaleOperator(KappaScaler.s:2ndpos)				0.332	1964	5301	7265	0.273
ScaleOperator(gammaShapeScaler.s:2ndpos)				0.332	2033	5177	7203	0.270
ScaleOperator(KappaScaler.s:3rdpos)				0.505	1424	5860	7210	0.282
ScaleOperator(gammaShapeScaler.s:3rdpos)				0.267	1569	5536	7105	0.133
bearcoperator (gammabhapebearer.s.orapos)				0.201	1000	3330	1100	0.221

Total calculation time: 964.067 seconds

Note that there is some useful information at the start concerning the alignments and which tree likelihoods are used. Also, all citations relevant for the analysis are mentioned at the start of the run, which can easily be copied to manuscripts reporting about the analysis. Then follows reporting of the chain, which gives some real time feedback on progress of the chain.

At the end, an operator analysis is printed, which lists all operators used in the analysis together with how often the operator was tried, accepted, and rejected (see columns #total, #accept and #reject respectively). The acceptance rate is the proportion of times an operator is accepted when it is selected for doing a proposal. In general, an acceptance rate that is high,

say over 0.5 indicates the proposals are conservative and do not explore the parameter space efficiently. On the other hand a low acceptance rate indicates that proposals are too aggressive and almost always result in a state that is rejected because of its low posterior. Both too high and too low acceptance rates result in low ESS values. An acceptance rate of 0.234 is the target (based on very limited evidence provided by [3]) for many (but not all) operators implemented in BEAST.

Some operators have a tuning parameter, for example the scale factor of a scale parameter. If the final acceptance rate is not near the target, BEAST will suggest a new value for the tuning parameter, which is printed in the operator analysis. In this case, all acceptance rates are good for the operators that have tuning parameters. Operators without tuning parameters include the wide exchange and Wilson-Balding operators for this analysis. Both these operators attempt to change the topology of the tree with large steps, but since the data supports a single topology overwhelmingly, these radical proposals are almost always rejected.

## 4 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** (Figure 13).

Remember that MCMC is a stochastic algorithm so the actual numbers will not be exactly the same as those depicted in the figure.

On the left hand side is a list of the different quantities that BEAST has logged to file. There are traces for the posterior (this is the natural logarithm of the product of the tree likelihood and the prior density), 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 the clockRate parameter in the lefthand list to look at the average rate of evolution (averaged over the whole tree and all sites). Tracer will plot a (marginal posterior) histogram for the selected statistic and also give you summary statistics such as the mean and median. The 95% HPD stands for highest posterior density interval and represents the most compact interval

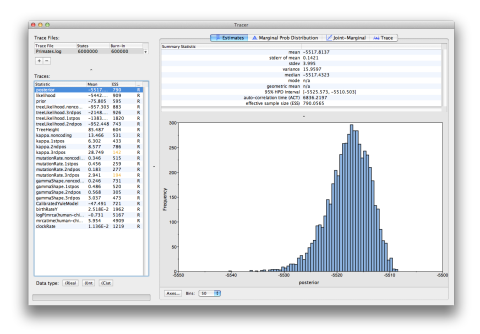


Figure 13: A screenshot of Tracer v1.6.

on the selected parameter that contains 95% of the posterior probability. It can be loosely thought of as a Bayesian analog to a confidence interval. The TreeHeight parameter gives the marginal posterior distribution of the age of the root of the entire tree.

Select the TreeHeight parameter and then Ctrl-click mrcatime (human-chimp) (Command-click on Mac OS X). This will show a display of the age of the root and the calibration MRCA we specified earlier in BEAUti. You can verify that the divergence that we used to calibrate the tree (mrcatime(human-chimp)) has a posterior distribution that matches the prior distribution we specified (Figure 14).

## 5 Marginal posterior estimates

To show the relative rates for the four partitions, select the mutationRate parameter for each of the four partitions, and select the marginal density tab in Tracer. Figure 15 shows the marginal densities for the relative substitution rates. The plot shows that codon positions 1 and 2 have substantially

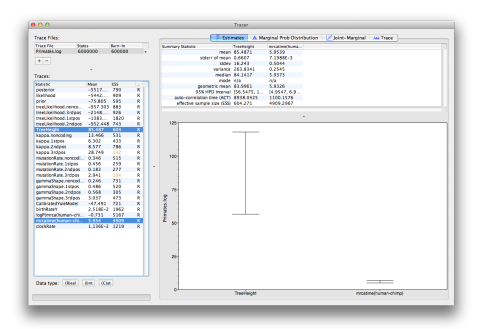


Figure 14: A screen shot of the 95% HPD intervals of the root height and the user-specified (human-chimp) MRCA in Tracer.

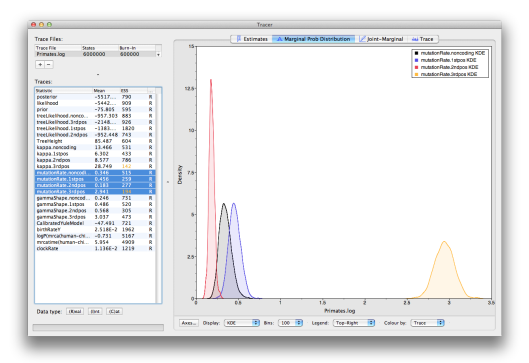


Figure 15: A screenshot of the marginal posterior densities of the relative substitution rates of the four partitions (relative to the site-weighted mean rate).

different rates (0.456 versus 0.183) and both are far slower than codon position 3 with a relative rate of 2.941. The noncoding partition has a rate intermediate between codon positions 1 and 2 (0.346). Taken together this result suggests strong purifying selection in both the coding and noncoding regions of the alignment.

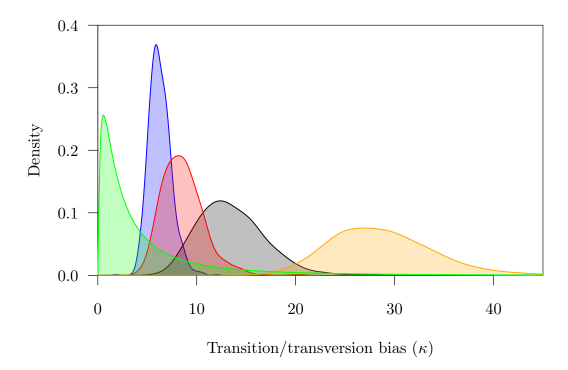


Figure 16: The marginal prior and posterior densities for the shape  $(\alpha)$  parameters. The prior is in gray. The posterior density estimate for each partition is also shown: noncoding (orange) and first (red), second (green) and third (blue) codon positions.

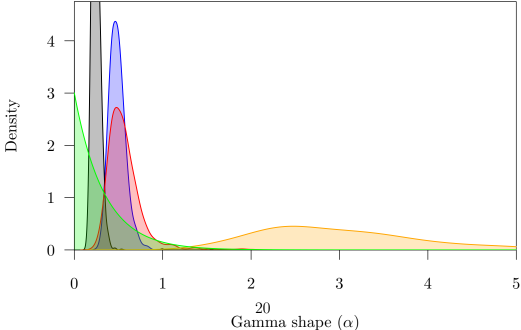


Figure 17: The marginal prior and posterior densities for the transition/transversion bias  $(\kappa)$  parameters. The prior is in gray. The posterior density estimate for each partition is also shown: noncoding (orange) and

## Questions

What is the estimated rate of molecular evolution for this gene tree (include the 95% HPD interval)?

What sources of error does this estimate include? How old is the root of the tree (give the mean and the 95% HPD range)?

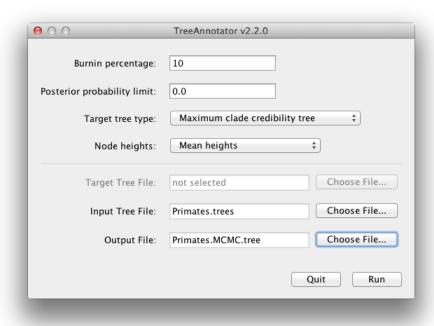


Figure 18: A screenshot of TreeAnnotator.

# 6 Obtaining an estimate of the phylogenetic tree

BEAST also produces a posterior sample of phylogenetic time-trees along with its sample of parameter estimates. These need to be summarized using the program **TreeAnnotator**. This will take the set of trees and find the best supported one. It will then annotate this representative summary tree with the mean ages of all the nodes and the corresponding 95% HPD ranges. It will also calculate the posterior clade probability for each node. Run the TreeAnnotator program and set it up as depicted in Figure 18.

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 6,000,000 steps sampling every 1,000 steps. Thus the trees file will contain 6,000 trees and so to specify a 10% burnin in the top text field.

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.

The **Target tree type** specifies the tree topology that will be annotated. 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.

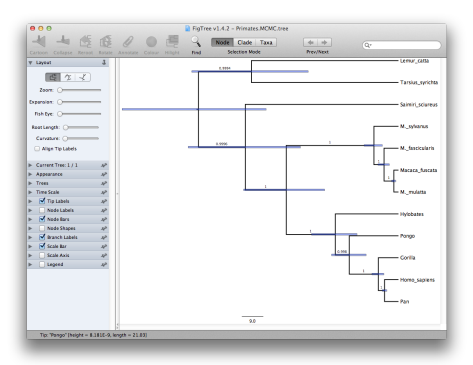
For node heights, the default is Common Ancestor Heights, which calculates the height of a node as the mean of the MRCA time of all pairs of nodes in the clade. For trees with large uncertainty in the topology and thus many clades with low support, some other methods can result in trees with negative branch lengths. In this analysis, the support for all clades in the summary tree is very high, so this is no issue here. 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 and select a file for the output (here we called it Primates.MCC.tree). Now press Run and wait for the program to finish.

## 7 Visualizing the tree estimate

Finally, we can visualize 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. First of all, expend **Trees** option in the panel, and check **Order nodes** and choose *Ordering* by decreasing. 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. If you use a non strict clock model then under **Appearance** you can also tell FigTree to colour the branches by the rate. You should end up with something similar to Figure 19.

An alternative view of the tree can be made with DensiTree, which is part of Beast 2. The advantage of DensiTree is that it is able to visualize both uncertainty in node heights and uncertainty in topology. For this particular



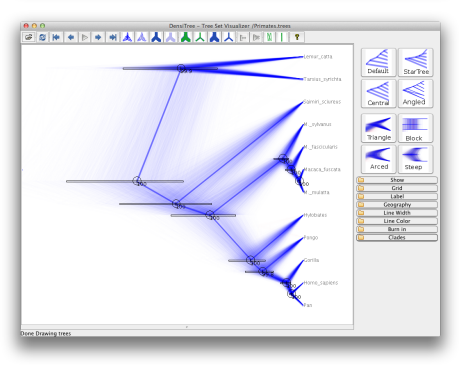


Figure 19: A screen shot of FigTree and Densi Tree.

dataset, the dominant topology is present in more than 99% of the samples. So, we conclude that this analysis results in a very high consensus on topology (Figure 19).

### Questions

- 1. Does the rate of evolution differ substantially amongst different lineages in the tree?
- 2. DensiTree has a clade bar (Menu Window/View clade toolbar) to show information on clades.
  - What is the support for the clade [Homo\_sapiens, Pan, Gorilla, Hylobates]?
- 3. You can browse through the topologies in Densi Tree using the Browse menu. The most popular topology has a support of over 99%.
  - What is the support for the second most popular topology?
- 4. Under the help menu, DensiTree shows some information.

  How many topologies are in the tree set?

## 8 Comparing your results to the prior

It is a good idea to rerun the analysis while sampling from the prior to make sure that interactions between priors are not affecting your prior information. The interaction between priors can be problematic especially when using calibrations since it means putting multiple priors on the tree.

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.

Divergence time estimation using "node dating" of the type described in this chapter has been applied to answer a variety of different questions in ecology and evolution. For example, node dating with fossils was used in determining the species diversity of cycads [5], analysing the rate of evolution in flowering plants [7], and investigating the origins of hot and cold desert cyanobacteria [1].

#### References

- [1] Justin Bahl, Maggie CY Lau, Gavin JD Smith, Dhanasekaran Vijaykrishna, S Craig Cary, Donnabella C Lacap, Charles K Lee, R Thane Papke, Kimberley A Warren-Rhodes, Fiona KY Wong, et al., Ancient origins determine global biogeography of hot and cold desert cyanobacteria, Nature communications 2 (2011), 163.
- [2] Alexei J Drummond and Marc A Suchard, Bayesian random local clocks, or one rate to rule them all, BMC biology 8 (2010), no. 1, 114.
- [3] A Gelman, G Roberts, and W Gilks, Efficient metropolis jumping hules, Bayesian statistics 5 (1996), 599–608.
- [4] Joseph Heled and Alexei J Drummond, Calibrated tree priors for relaxed phylogenetics and divergence time estimation, Syst Biol **61** (2012), no. 1, 138–49.
- [5] NS Nagalingum, CR Marshall, TB Quental, HS Rai, DP Little, and S Mathews, *Recent synchronous radiation of a living fossil*, Science **334** (2011), no. 6057, 796–799.

- [6] Michael S Rosenberg, Sankar Subramanian, and Sudhir Kumar, *Patterns of transitional mutation biases within and among mammalian genomes*, Molecular biology and evolution **20** (2003), no. 6, 988–993.
- [7] Stephen A Smith and Michael J Donoghue, Rates of molecular evolution are linked to life history in flowering plants, science **322** (2008), no. 5898, 86–89.