# \*BEAST in BEAST 2.4.x

# Estimating Species Tree from Multilocus Data

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# 1 Introduction

In this tutorial we describe a full Bayesian framework for species tree estimation. The statistical methodology described in this tutorial is known by the acronym \*BEAST (pronounced "star beast") (Heled and Drummond 2010).

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 v2.4.x, which has support for multiple partitions. It is available for download from http://beast2.org (Bouckaert et al. 2014).
- **Tracer** this program is used to explore the output of BEAST (and other Bayesian MCMC programs). It summarizes graphically and quantitively the distributions of continuous parameters and provides diagnostic information for the particular MCMC chain. At the time of writing, the current version is v1.6. It is available for download from
- http://tree.bio.ed.ac.uk/software/tracer.
- **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/figtree.

## 2 \*BEAST analysis

This tutorial will guide you through the analysis of three loci sampled from 26 individuals representing nine species of pocket gophers. This is a subset of previously published data (Belfiore et al. 2008). The objective of this tutorial is to estimate the species tree that is most probable given the multi-individual multi-locus sequence data. The species tree has nine taxa, whereas each gene tree has 26 taxa. \*BEAST 2 will co-estimate three gene trees embedded in a shared species tree (see Heled and Drummond 2010 for details).

The first step in the analysis 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 (Bayesian Evolutionary Analysis Utility). This is a user-friendly program for setting the evolutionary model and options for the MCMC analysis. The second step will be to actually run BEAST using the XML input file that contains the data, model and MCMC chain settings. The final step will be to explore the output of BEAST in order to diagnose problems and to summarize the results.

## 2.1 BEAUti

Start up BEAUti to set up the configuration file for the analysis.

#### Set up BEAUti for \*BEAST 2.1.1

BEAUti uses templates to populate the BEAST XML configuration file with the necessary elements. This template also allows BEAUti to set up the graphical user interface accordingly, having the right tabs and settings.

\*BEAST uses a non-standard template to generate the analysis XML, which means that to use BEAUti for \*BEAST, the first thing to do is to change the template. Choose the File/Template/StarBeast item (Fig. 1). Keep in mind that when changing a template, BEAUti deletes all previously imported data and starts with a clean template. So, if you already loaded some data, a warning message will pop up indicating that this data will be lost if you switch templates.

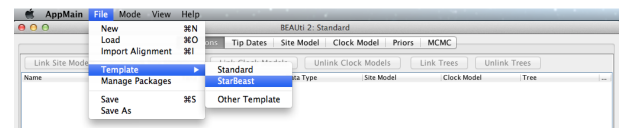


Figure 1: Select a new template in BEAUti.

#### Loading the NEXUS file 2.1.2

\*BEAST is a multi-individual, multi-locus method. The data for each locus is stored as one alignment in its own NEXUS file. Taxa names in each alignment have to be unique, but duplicates across alignments are allowed.

To load a NEXUS format alignment, simply select the **Import Alignment** option from the **File** menu. Select three files called 26.nex, 29.nex, 47.nex by holding shift key. You can find the files in the examples/nexus directory in the directory where BEAST was installed. Alternatively, click the links above to download the data. After the data is opened in your web browser each time, right click with the mouse and separately save them as 26.nex, 29.nex, 47.nex.

Each file contains an alignment of sequences from an independent locus. The 26.nex looks like this (content has been truncated):

```
#NEXUS
[TB026oLong]
BEGIN DATA;
   DIMENSIONS NTAX =26 NCHAR=614;
   FORMAT DATATYPE = DNA GAP = - MISSING = ?;
   MATRIX
   Orthogeomys_heterodus
                                 ATTCTAGGCAAAAAGAGCAATGC ...
   Thomomys bottae awahnee a
                                 ??????????????????ATGCTG ...
   Thomomys bottae awahnee b
                                 ??????????????????ATGCTG ...
   Thomomys_bottae_xerophilus
                                 ??????????????????ATGCTG ...
                                 ??????????????AGCAATGCT ...
   Thomomys_bottae_cactophilus
    . . . . . .
```

```
END;
```

Once loaded, the three partitions are displayed in the main panel. You can double click any alignment (partition) to show its detail. Figure 2 shows the way BEAUti looks after loading the appropriate alignments.

00				BEAUti 2: StarBe	ast			
	Partit	ions Taxon sets	Site Mode	I Clock Model	Multi Species Coale	scent Priors MCI	МС	
Link Site M	odels Unlink	Site Models	Link Clock N	lodels Unlink	Clock Models	Link Trees Unl	ink Trees	
Name	File	Taxa	Sites	Data Type	Site Model	Clock Model	Tree	
26	26	26	614	nucleotide	26	\$ 26	\$ 26	\$
29	29	26	601	nucleotide	29	\$ 29	\$ 29	\$
47	47	26	819	nucleotide	47	\$ 47	\$ 47	\$

Figure 2: Data partition panel after loading alignments.

For multi-locus analyses, BEAST can link or unlink substitutions models across the loci by clicking buttons on the top of **Partitions** panel. The default of \*BEAST is unlinking all models: substitution model, clock model, and tree models. For this tutorial we shall leave all models unlinked.

Note that you should only unlink the tree model across data partitions that are actually genetically unlinked. For example, in most organisms all the mitochondrial genes are effectively linked due to a lack of recombination and they should be set up to use the same tree model in a \*BEAST analysis. It could also be that for this analysis a linked model would perform better.

#### 2.1.3 Guess or import trait(s) from a mapping file

Each taxon in a \*BEAST analysis is associated with a species. Typically, the species name is already embedded inside the taxon. The species name should be easy to extract; place it either at the beginning or the end, separated by a "special" character which does not appear in names. For example, aria\_334259, coast\_343436 (using an underscore) or 10x017b.wrussia, 2x305b.eastis (using a dot).

We need to tell BEAUti somehow which lineages in the alignments go with taxa in the species tree. Select the **Taxon Set** panel, and a list of taxa from the alignments is shown together with a default guess by BEAUti. In this case, the guess is not very good, so we want to change this. You can either change each of the entries in the table manually, have a mapping stored in a file or have BEAUti guess the taxon names.

If the mapping is to be read from a file one needs to specify a proper trait file. The first row is always specified as the keyword **traits** in the first column followed by the keyword **species** in the second column, separated by tab. The rest of the rows map each individual taxon name to a species name; the taxon name in the first column and species name in the second column separated by tab. For example:

traits species
taxon1 speciesA
taxon2 speciesA
taxon3 speciesB
...

Alternatively, pressing the **Guess** button in BEAUti, a dialog is shown where you can choose from several ways to try to detect the species from the name of the taxon. In our case, splitting the name on the underscore character (\_) and selecting the second group will give us the mapping that we need (Fig. 3).

	Deutisiana Taura esta Cita Mad	al - Clash Madel - Nubi Georgias Castararent - Prises - MCN
	Partitions Taxon sets Site Mod	el Clock Model Multi Species Coalescent Priors MCM
lter:		
	Taxon	Species/Population
	Orthogeomys_heterodus	0
	Thomomys_bottae_albatus	Т
	Thomomys_bottae_alpinus	Т
	Thomomys_bottae_awahnee_a	T
	Thomomys_bottae_awahnee_b	Т
	Thomomys_bottae_bottae	T
	Thomomy lotter caston hiller	Guess taxon sets
	Thomorry	
	Thomomy: Thomomy: use everything	after first 💠
	Thomomy: Thomomy:	
	Thomomy	
	Thomomy	
	Thomomy 💽 split on character	and take group(s): 2 ‡
	Thomomy	
	Thomomy	
	Thomomy	
		A( +)( -) 1( +)(
	Thomomy Ouse regular expression	on ^(.+)[\.](.*)\$
	Thomomy	
	Thomomy	
	Thomomy	
	Thomomy: O read from file	File Browse ?
	Thomomy	
	Thomomy:	
		OK Cancel
	Fill down	Guess

Figure 3: Selecting taxon sets in BEAUti using the guess dialog from the Taxon set panel.

### 2.1.4 Setting up the substitution model

The next thing to do is to click on the **Site Model** tab at the top of the main window. This will reveal the evolutionary model settings for BEAST. Exactly which options appear depends on whether the data are nucleotides, amino acids, binary data, or general data. The settings that appear after loading the dataset are always the default values.

Thus, we have to pick a substitution model that will work better on real data than the default JC69. For this analysis, we select each partition of the data on the left side of the panel and set up the same substitution model for all three partitions: 'HKY' for **Substitution Model** and 'Empirical' for the **Frequencies** (Fig. 4).

#### 2.1.5 Setting up the clock model

Then, click on the **Clock Model** tab at the top of the main window. In this analysis, we use the 'Strict Clock' molecular clock model. Your model options should now look as displayed in Figure 5.

The **estimate** check box is unchecked for the first clock model and checked for the rest clock models, because we wish to estimate the substitution rate of each subsequent locus relative to the first locus whose rate is fixed to 1.0.

00			BEAUti 2: StarBeast	
	Partitions	Taxon sets Site Model	Clock Model   Multi Species Coalescent   Priors   MCMC	
Partition 26	Substitution Rate	1.0		🗌 estimate 🥖
29 47	Gamma Category Count	0		
	Proportion Invariant	0.0		🗌 estimate 🥖
		НКҮ		\$
		Карра	2.0	🗹 estimate 🥖
		Frequencies	Empirical	÷ 🥖
	Subst Model			

Figure 4: Setting up substitution and site models for the gopher alignments.

0 0	BEAUti 2: StarBeast	
	Partitions Taxon sets Site Model Clock Model Multi Species Coalescent Priors MCMC	
Partition 26 29 47 Strict Clock Clock.rate	1.0	÷
000	BEAUti 2: StarBeast	
	Partitions Taxon sets Site Model Clock Model Multi Species Coalescent Priors MCMC	
Partition 26 29 47 Clock.rate	1.0	♦ ♦ ♦ ♦
000	BEAUti 2: StarBeast	
	Partitions Taxon sets Site Model Clock Model Multi Species Coalescent Priors MCMC	
Partition 26 29 47 Clock.rate	1.0	🔹 estimate 🥒

Figure 5: Setting up clock models for the gopher alignments.

### 2.1.6 Multi Species Coalescent

The **Multi Species Coalescent** panel allows settings of the multi-species coalescent model to be specified. Users can only configure the species tree prior, not the gene tree priors, which are automatically specified by the multispecies coalescent. Currently, three population size models for species tree (under the entry **Pop Function**) are available: 'Continuous-linear and constant root', 'Continuous-linear', and 'Constant'. In this analysis, since we don't have much information for the population change at the root, we use 'linear and constant root' model. In \*BEAST, the population sizes are assigned gamma priors, and users cannot change the type to a different distribution. But the scale parameter of the gamma prior (called **Population Mean**) can be assigned a hyperprior (tick 'estimate', Fig. 6).

The **Ploidy** item determines the type of sequence (mitochondrial, nuclear, X, Y). This matters since different modes of inheritance give rise to different effective population sizes, for example in the case of mitochondrial DNA we would only deal with the half of the available population, since this type of DNA is only passed on by the mother. In the case of this analysis the data is autosomal nuclear, which is the default option (Fig. 6).

00	BEAUti 2: StarBeast	
	Partitions         Taxon sets         Site Model         Clock Model         Multi Species Coalescent         Priors         MCMC	
▼ Species Tree Popu	lation Size	
Pop Function	linear_with_constant_root \$	
Gamma Parameter	1.0	🗹 estimate 🥖
▼ Gene Tree 26		
Ploidy	autosomal_nuclear 🛊	
▼ Gene Tree 29		
Ploidy	autosomal_nuclear 🛟	
▼ Gene Tree 47		
Ploidy	autosomal_nuclear ‡	

Figure 6: Setting up the multi species coalescent model parameters.

#### 2.1.7 Priors

The **Priors** panel allows priors for each parameter in the model to be specified.

Several models are available as species tree priors, however, only the Yule or the birth-death model are suitable for \*BEAST analyses. Leave the tree prior setting at the default 'Yule Model'. This prior specifies the distribution of the speciation times.

The default priors that BEAST sets for other parameters would allow the analysis to work. However, some of these are inappropriate for this analysis. Therefore change the priors as follows:

- birthRate.t:Species: set the prior to 'Log Normal' with M of 4.0 and S of 1.25;
- **popMean** (scale parameter of the gamma priors for population sizes): set the prior to 'Log Normal' with M of -5.0 and S of 1.25;
- clockRate.c:29: set the prior to 'Exponential' with mean 1.0;
- clockRate.c:45: set the prior to 'Exponential' with mean 1.0.

The resulting configuration is shown in Figure 7.

#### 2.1.8 Operators

The **Operators** panel (normally hidden) is used to configure technical settings that affect the efficiency of the MCMC program. In MCMC the operators define which values will be proposed for the next step in the chain given the current state. You can take a look at them if you go to **View/Show Operators panel**, but do not edit the settings.

#### 2.1.9 Setting the MCMC options

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

First, we need to set the **Chain Length**. This is the number of steps the MCMC will make in the chain before finishing. The appropriate length of the chain depends on the size of the dataset, the complexity of the model and the accuracy of the answer required. The default value of 10,000,000 is entirely arbitrary and should be adjusted according to the size of your dataset. For this dataset let's set the chain length to

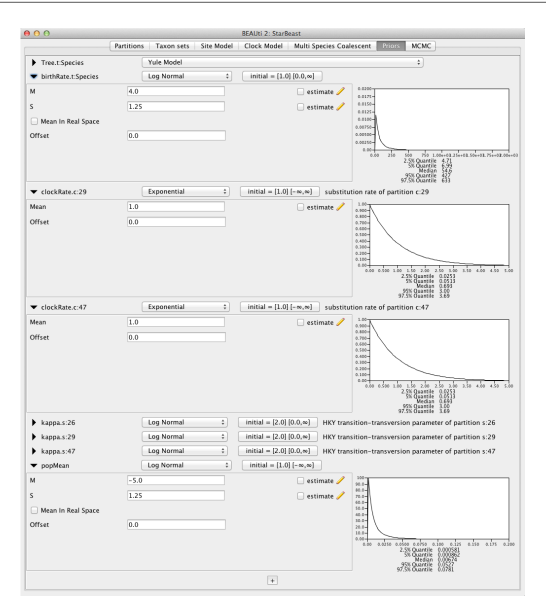


Figure 7: Setting up parameter priors.

5,000,000 as this will run reasonably quickly on most modern computers (a few minutes).

The next options specify how often should the parameter values in the Markov chain be displayed on the screen and recorded in the log file. The screen output is simply for monitoring the program's progress and can thus be set to any value (although if set too small, i.e. if displaying on screen too often, 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 the log frequency should be set to no less than 5,000,000 / 10,000 = 500.

For this exercise we will set the the tracelog to 5,000 and the screenlog to 10,000. Set the file name of

the tracelog to gopher.log, that of the speciesTreeLogger to gopher.species.trees, and that of the treelog.t:26, treelog.t:29, treelog.t:47 to gopher.gene\$(tree).trees.

00				BEAUti 2: StarB	east			
(	Partitions	Taxon sets	Site Model	Clock Model	Multi Species Coalescent	Priors	мсмс	
Chain Length	50000	00						
Store Every	5000							
Pre Burnin	0							
Num Initialization Attempt	s 10							
▼ tracelog								
File Name	gopher	.log						
Log Every	5000							
Mode	autoo	letect			\$			
Sort	smar	t			* *			
Sanitise Headers								
posterior likelihood prior speciescoalescent TreeHeight.Species								
<ul> <li>speciesTreeLogger</li> </ul>								
File Name	gopher	.species.trees						
Log Every	5000							
Mode	tree				\$			
Sort	none				\$			
Sanitise Headers								
SpeciesTreeLoggerX								J
screenlog								
Sample From Prior								

The resulting configuration is shown in Figure 8.

Figure 8: Setting up the MCMC parameters.

If you are using Windows then we suggest you add the suffix .txt to both of these files (i.e. to specify output file names to gopher.log.txt and gopher.species.trees.txt) so that Windows recognizes these as text files.

### 2.1.10 Generating the BEAST XML file

We are now ready to create the BEAST XML file. To do this, either select the **File/Save** or **File/Save As** option from the **File** menu. Save the file with an appropriate name (we usually end the filename with .xml, i.e. gopher.xml). We are now ready to run the file through BEAST.

### 2.2 BEAST

Now run BEAST. Provide your newly created XML file as input by clicking **Choose File** ..., and then click **Run** (Fig. 9).

BEAST will then run until the specified chain length is reached, and until it has finished reporting infor-

mation on the screen. The actual result files are saved to the disk in the same location as your input file. The output to the screen will look something like this:

BEAST v2.4.2, 2002-2016 Bayesian Evolutionary Analysis Sampling Trees Designed and developed by Remco Bouckaert, Alexei J. Drummond, Andrew Rambaut & Marc A. Suchard

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> Downloads, Help & Resources: http://beast2.org/

Source code distributed under the GNU Lesser General Public License: 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\includeimage, Beth Shapiro and Korbinian Strimmer

4980000	-3827.2093	279.2	-4293.2486	2	2.1146 24s/M	samples		
4990000	-3842,9689	276.3	-4295.6213		7.6877 24s/M			
5000000	-3798.1962	278.2	-4280.7865		2.0904 24s/M			
						1		
Operator			Tuning	#accept	#reject	Pr(m)	Pr(acc m)	
NodeReheight(Rehe	ight.t:Species)		-	205317	486572	0.1386	0.2967	
ScaleOperator(pop	SizeBottomScaler	.t:Species)	0.1902	10224	26338	0.0074	0.2796	
ScaleOperator(pop	MeanScale.t:Spec:	ies)	0.4890	5795	16242	0.0044	0.2630	
UpDownOperator(up	down.all.Species	)	0.5029	38885	108717	0.0295	0.2634	
ScaleOperator(Yul	eBirthRateScaler	.t:Species)	0.2416	7155	15125	0.0044	0.3211	
ScaleOperator(tre	eScaler.t:26)	-	0.7580	8510	42962	0.0102	0.1653	
ScaleOperator(tre	eRootScaler.t:26	)	0.4424	11487	39392	0.0102	0.2258	
Uniform(UniformOp	erator.t:26)		-	287517	225590	0.1024	0.5603	
SubtreeSlide(Subt	reeSlide.t:26)		0.5560	418	256419	0.0512	0.0016	Try decreasing size to about 0.278
Exchange(narrow.t	:26)		-	115438	140923	0.0512	0.4503	
Exchange(wide.t:20	6)		-	1363	49988	0.0102	0.0265	
WilsonBalding(Wil:	sonBalding.t:26)		-	1882	49890	0.0102	0.0364	
ScaleOperator(Str	ictClockRateScale	er.c:29)	0.4744	13355	37595	0.0102	0.2621	
ScaleOperator(tre	eScaler.t:29)		0.7567	8137	42785	0.0102	0.1598	
ScaleOperator(tree	eRootScaler.t:29	)	0.4101	12191	38887	0.0102	0.2387	
Uniform(UniformOp	erator.t:29)		-	296262	216280	0.1024	0.5780	
SubtreeSlide(Subt	reeSlide.t:29)		0.5724	339	255443	0.0512	0.0013	Try decreasing size to about 0.286
Exchange(narrow.t	:29)		-	117912	138109	0.0512	0.4606	
Exchange(wide.t:2	9)		-	2229	49303	0.0102	0.0433	
WilsonBalding(Wil:	sonBalding.t:29)		-	2459	49001	0.0102	0.0478	
UpDownOperator(up	down.29)		0.7793	10314	40694	0.0102	0.2022	
ScaleOperator(Str	ictClockRateScal	er.c:47)	0.5490	12974	38097	0.0102	0.2540	
ScaleOperator(tree	eScaler.t:47)		0.7363	8746	42523	0.0102	0.1706	
ScaleOperator(tree	eRootScaler.t:47	)	0.4742	7774	43670	0.0102	0.1511	
Uniform(UniformOp	erator.t:47)		-	276942	234361	0.1024	0.5416	
SubtreeSlide(Subt	reeSlide.t:47)		0.5044	307	255119	0.0512	0.0012	Try decreasing size to about 0.252
Exchange(narrow.t	:47)		-	91846	164747	0.0512	0.3579	
Exchange(wide.t:4	7)		-	731	50281	0.0102	0.0143	
WilsonBalding(Wils	sonBalding.t:47)		-	937	50377	0.0102	0.0183	
UpDownOperator(up	down.47)		0.7397	10853	40537	0.0102	0.2112	
UpDownOperator(st	rictClockUpDownOj	perator.c:47)	0.7278	9987	41373	0.0102	0.1945	
UpDownOperator(st	rictClockUpDownOj	perator.c:29)	0.7761	10040	40724	0.0102	0.1978	
ScaleOperator(Kap	paScaler.s:26)		0.3401	469	1197	0.0003	0.2815	
ScaleOperator(Kap			0.2955	436	1231	0.0003	0.2615	
ScaleOperator(Kap	paScaler.s:47)		0.3199	379	1298	0.0003	0.2260	
ScaleOperator(pop	SizeTopScaler.t:	Species)	0.1910	23665	54936	0.0157	0.3011	

Tuning: The value of the operator's tuning parameter, or '-' if the operator can't be optimized.

#accept: The total number of times a proposal by this operator has been accepted. #reject: The total number of times a proposal by this operator has been rejected.

Pr(m): The probability this operator is chosen in a step of the MCMC (i.e. the normalized weight).

 $\Pr(acc|m)$ : The acceptance probability (#accept as a fraction of the total proposals for this operator).

Total calculation time: 135.584 seconds

End likelihood: -3798.196207797654

	BEAST v2.4.2
Bay	yesian Evolutionary Analysis Sampling Trees Version v2.4.2, 2002–2016
BEAST XML File:	gopher.xml Choose File
	default: only write new log files
Random number seed:	1466686429679
Thread pool size:	Automatic ᅌ
Use BEAGLE library	if available:
Prefer use of: CP	PU ≎
Prefer precision: Do	ouble 🗘
2	Show list of available BEAGLE resources and Quit
	rmance phylogenetic library that can make use of
additional computationa	I resource's such as graphics boards. It must be ed independently of BEAST:

Figure 9: Launching BEAST.

## 3 Analyzing the results

### 3.1 Tracer

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 **gopher.log**. You should now see a window looking like the snapshot in Figure 10.

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 likelihood and the prior probabilities), and the continuous parameters. Selecting one trace from this list on the left-hand side brings up analyses for this trace on the right-hand side. When a file is first opened in Tracer, the 'posterior' trace is selected and various statistics of this trace are shown under the **Estimates** tab (top window on the right-hand side).

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 lower or upper 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.

Select the three 'TreeHeight' traces (hold **shift** key while selecting). This will display the age of the root for the three gene trees. If you switch the tab at the top of the window on the right-hand side to **Marginal Prob Distribution** then you will get a plot of the marginal posterior densities of each of these date estimates overlayed (Fig. 11).

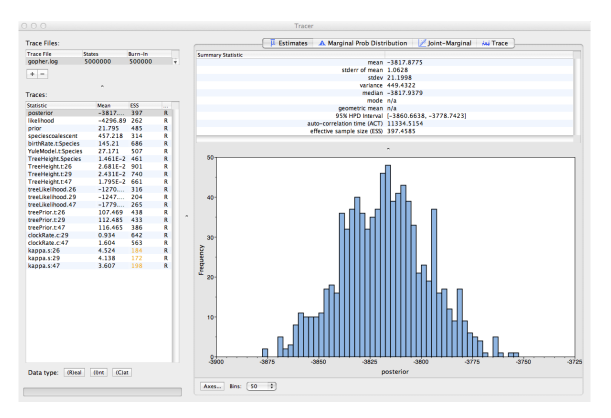


Figure 10: Tracer with the gopher data.

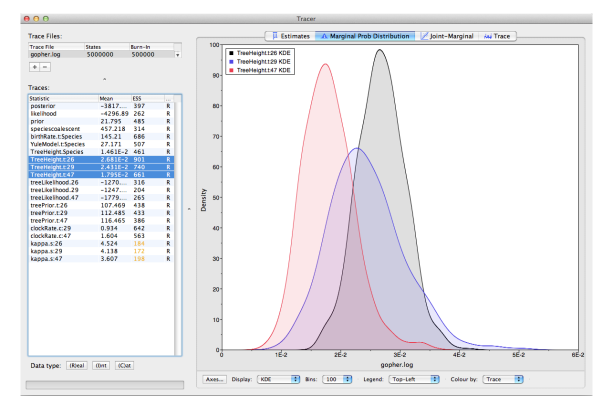


Figure 11: Tracer showing the root heights of the lineage trees.

## 3.2 TreeAnnotator

BEAST also produces a sample of plausible trees. These can be summarized using the program **TreeAnnotator**. This program will take the set of trees and identify a single tree that best represents the posterior distribution. It will then annotate this selected tree topology with the mean ages of all the nodes as well as the 95% HPD interval of divergence times for each clade in the selected tree. It will also calculate the posterior clade probability for each node. Run the TreeAnnotator program and set it up as shown in Figure 12.

000	TreeAnnotator v2.3.0	
Burnin percentage:	10	
Posterior probability limit:	0.5	
Target tree type:	Maximum clade credibility tree	÷ +
Node heights:	Median heights	•
Target Tree File:	not selected	Choose File
Input Tree File:	gopher.species.trees	Choose File
Output File:	gopher.species.tree	Choose File
	Qu	uit Run

Figure 12: Using TreeAnnotator to summarise the tree set.

The burnin is the number of trees to remove from the start of the sample. Unlike in Tracer where burnin is specified as the number of steps, in TreeAnnotator you need to specify the percentage of trees discarded. For this run, we set the burnin to 10 (i.e. we remove first 10% of the trees).

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. Setting this to zero will annotate all nodes. For this analysis, set it to 0.5. This means that only the nodes seen in the majority of trees will be annotated.

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 'Median heights' for **Node Heights**. This sets the heights (ages) of each node in the tree to the median height across the entire sample of trees for that clade.

For the input file, select the trees file that BEAST created (in this tutorial this was the gopher.species.trees) and select a file for the output (here we called it gopher.species.tre).

Now press **Run** and wait for the program to finish.

### 3.3 Viewing the species tree

Finally, we can look at the tree in another program called **FigTree**. Run this program, and open the **gopher.species.tre** 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. Tick the **Node Bars** and try 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 'posterior'. You should end up with a tree similar to one displayed in Figure 13.

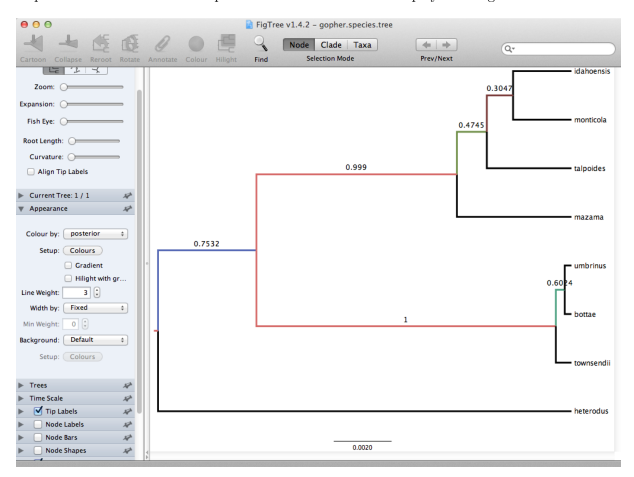


Figure 13: Figure representation of the gopher species tree.

Alternatively, you can load the species tree set gopher.species.trees into **DensiTree** and visualize it with following settings.

- Perhaps, the intensity of the individual trees is not high enough, so you might want to increase it by clicking the icon **Increase Intensity of Trees** in the button bar at the top. The text for each button shows when you hover over it.
- Set **Burn in** to 500 trees. If before you had a collapsed trees, after this setting the tree should not be collapsed anymore.
- Show a root-canal tree to better guide the eye over the set of trees.
- In **Clades** menu, select **Show clades** and display their mean and 95% HPD graphically, and posterior support using text. By default you may not be able to select the **Show clades** option. If that is the case, first click on the icon **Central** in the top right corner of the window.

You can see that there is information for too many clades being displayed. As most clades have very little support, they are not of interest and we would like to stop visualizing the information for those. Tick **Selected only**, then open the clade toolbar (menu **Window/View clade toolbar**), and select only highly supported clades (hold down the **shift** key to select several clades at a time). Also, select the clade consisting of heterodus, bottae, umbinus and townsendii, to show that heterodus is an outgroup of the tree, but there is some support ( $\sim 17\%$ ) that it is not.

• Drag the clade monticola and idahoensis up in the displayed tree so that the 95% HPD bar does not overlap with the one for mazama, monticola, idahoensis and talpoidis. Increase font size of the label for better readability (i.e. change the **Label** options).

You should now see an image similar to Figure 14.

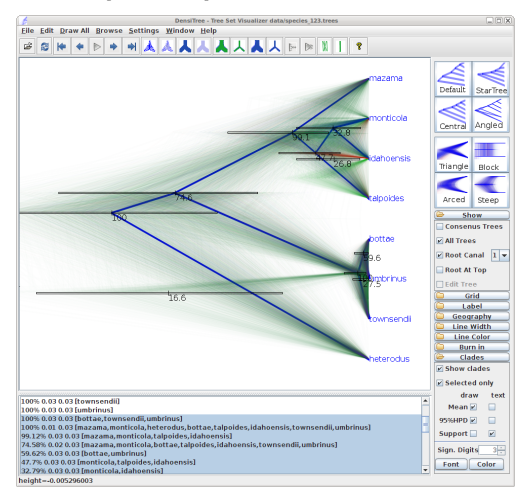


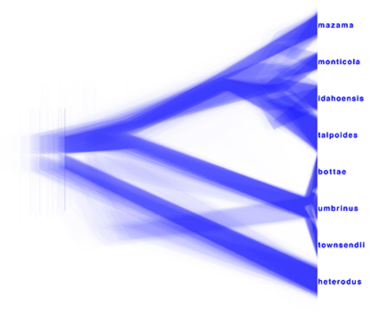
Figure 14: DensiTree representation of the species tree.

**DensiTree** can be used to show the population sizes of the each of the trees from gopher.species.trees as branch widths. Under Line Width, select 'BY\_METADATA\_PATTERN' option, for both upper and lower spinners and use

.\*dmv=.(^,\*).\*

for the upper (start of the branch) pattern and

#### .\*dmv=.^,\*,(^\}\*).\*



for the lower (end of the branch) pattern. This gives us visualisation as seen in Figure 15.

Figure 15: DensiTree representation of the consensus species tree with population sizes indicated as line widths.

Similarly, DensiTree can be used to show the population sizes of the summary tree gopher.species.tre from TreeAnnotator. Load this file to DensiTree and explore the options in the spinners under Line Width. Notice, that dmt stands for "demographic times", dmv for "demographic values". These are the population parameters that are estimated for each node in the tree. With the continous-linear population size model, the first value dmv1 is the population size at the branch start and the second dmv2 is the population size at the branch end; where start and end are relative to going *backwards* in time, that is start is the more recent time, end the more ancient.

Choose 'dmv1' for the upper (start/bottom of a branch) and 'dmv2' for the lower spinner (end/top of a branch). The resulting tree should look like the one in the left picture in Figure 16. You can see that at the nodes, the population size estimates do not quite match. This is true for the areas of the tree with little posterior support for the clades (see Figure 14 to check which clades have little support). In the right picture in Figure 16 the top is matched up with the bottom of the branch above (more ancestral), using the 'Make fit to bottom' option for the lower spinner. This looks a bit prettier, but may not be quite accurate.

Alternatively to TreeAnnotator, a consensus tree can be generated by biopy (https://github.com/jheled/biopy). An example is shown in Figure 17 where 1-norm option has been applied to generate the left picture, and 2-norm to obtain the right picture.

### 3.4 Comparing your results to the prior

Using BEAUti, set up the same analysis but under the **MCMC** options, select the **Sample From Prior** option. Re-run the analysis with BEAST. This will allow you to visualize the full prior distribution in

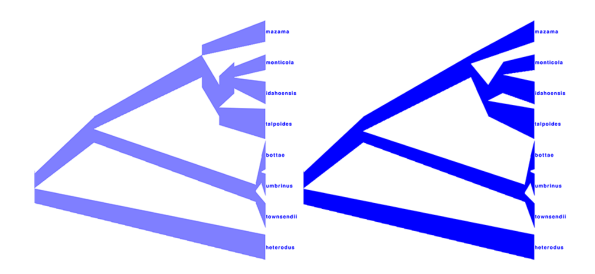


Figure 16: DensiTree representation of the consensus species tree from TreeAnnotator.

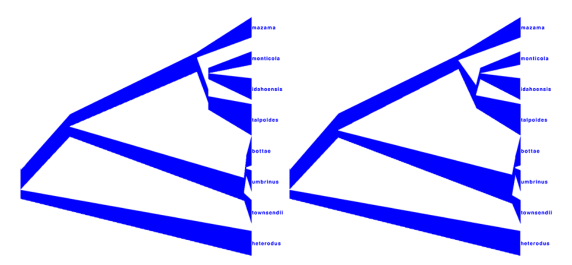


Figure 17: DensiTree representation of the consensus species tree from biopy.

the absence of your sequence data. Summarize the trees from the full prior distribution and compare the summary to the posterior summary tree.

You can also compare the log files from both analyses by loading the log file from the actual analysis and the log file from the prior sampling run into Tracer. Then select both of the files on the left side using the **shift** key. Now you can select different statistics from the traces and compare their marginal probability distributions. Note that Tracer will not allow you to compare any of the likelihoods as they are always NaN (not a number) in the case of prior sampling.

Figure 18 shows an example of comparing the traces of the species tree height. From the plot we can conclude that the sequences contain quite a bit of information about this parameter, as the posterior moved away from the prior and is much more focused on a specific value.

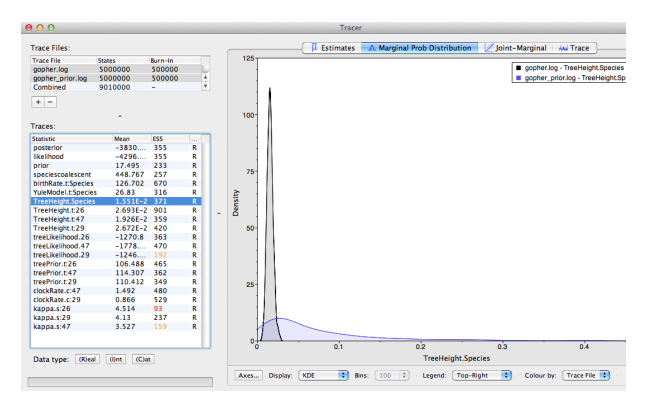


Figure 18: Tracer showing the comparison between the prior and posterior distribution of the species tree height.

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## **Relevant References**

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