## Command Reference for MrBayes ver. 3.2

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Sumt -- Summarizes trees from MCMC analysis
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Unlink -- Unlinks parameters across character partitions

Version -- Shows program version

Commands that should be in a NEXUS file (data block, trees block or taxa block) include:

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Begin -- Denotes beginning of block in file
Dimensions -- Defines size of character matrix
End -- Denotes end of a block in file
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Endblock -- Alternative way of denoting end of a block
Format -- Defines character format in data block
Matrix -- Defines matrix of characters in data block

Taxlabels -- Defines taxon labels

Translate -- Defines alternative names for taxa

Tree -- Defines a tree

Note that this program supports the use of the shortest unambiguous spelling of the above commands (e.g., "exe" instead of "execute").

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About

This command provides some general information about the program.

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Acknowledgments

This command shows the authors' acknowledgments.

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Calibrate

This command dates a terminal or interior node in the tree. The format is

```
calibrate <node_name> = <age_prior>
```

where <node\_name> is the name of a defined interior constraint node or the name of a terminal node (tip) and <age\_prior> is a prior probability distribution on the age of the node. The latter can either be a fixed date, a uniform probability distribution between a minimum and a maximum age, or an offset exponential distribution with a minimum age and a rate parameter. The format is:

```
calibrate <node_name> = fixed(<age>)
calibrate <node_name> = uniform(<min_age>,<max_age>)
calibrate <node_name> = offsetexponential(<min_age>,<rate>)
```

For instance, assume that we had three fossil terminals in our analyses named 'FossilA', 'FossilB', and 'FossilC'. Assume further that we want to fix the

age of FossilA to 100.0 million years, we think that FossilB is somewhere between 100.0 and 200.0 million years old, and that FossilC is at least 300.0 million years old, possibly older but relatively unlikely to be more than 400.0 million years old. Then we might use the commands:

calibrate FossilA = fixed(100) FossilB = uniform(100,200)
calibrate FossilC = offsetexponential(300,0.01)

Note that it is possible to give more than one calibration for each 'calibrate' statement. The offset exponential distribution has its expectation at the minimum age plus the inverse of the rate parameter. That is, the offsetexponential(300,0.01) has the minimum 300, the expectation 400 = = 300 + 1/0.01 and the mode < 400, so that most of the probability distribution is on values smaller than 400.

To actually use the calibrations to obtain dated trees, you also need to set a clock model using relevant 'brlenspr' and 'nodeagepr' options of the 'prset' command. You may also want to examine the 'clockvarpr' and 'clockratepr' options. Furthermore, you need to activate the relevant constraint(s) using 'topologypr', if you use any dated interior nodes in the tree.

You may wish to remove a calibration from an interior or terminal node, which has previously been calibrated. You can do that using

calibrate <node\_name> = unconstrained

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Charset

This command defines a character set. The format for the charset command is

charset <name> = <character numbers>

For example, "charset first\_pos =  $1-720\3$ " defines a character set called "first\_pos" that includes every third site from 1 to 720. The character set name cannot have any spaces in it. The slash (\) is a nifty way of telling the program to assign every third (or second, or fifth, or whatever) character to the character set. This option is best used not from the command line, but rather as a line in the mrbayes block of a file. Note that you can use "." to stand in for the last character (e.g., charset  $1-.\3$ ).

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#### Charstat

This command shows the status of all the characters. The correct usage is

charstat

After typing "charstat", the character number, whether it is excluded or included, and the partition identity are shown. The output is paused every 100 characters. This pause can be turned off by setting autoclose to "yes" (set autoclose=yes).

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#### Citations

This command shows a thorough list of citations you may consider using when publishing the results of a MrBayes analysis.

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#### Comparetree

This command compares the trees in two files, called "filename1" and "filename2". It will output a bivariate plot of the split frequencies as well as plots of the tree distance as a function of the generation. The plots can be used to get a quick indication of whether two runs have converged onto the same set of trees. The "Comparetree" command will also produce a ".pairs" file and a ".dists" file (these file endings are added to the end of the "Outputname"). The ".pairs" file contains the paired split frequencies from the two tree samples; the ".dists" file contains the tree distance values.

Note that the "Sumt" command provides a different set of convergence diagnostics tools that you may also want to explore. Unlike "Comparetree", "Sumt" can compare more than two tree samples and will calculate consensus trees and split frequencies from the pooled samples.

-- If this option is set to 'Yes', then a proportion of the

-- Name of the file to which 'comparetree' results will be

### Options:

Relburnin

		samples will be discarded as burnin when calculating summary
		statistics. The proportion to be discarded is set with
		Burninfrac (see below). When the Relburnin option is set to
		'No', then a specific number of samples is discarded instead.
		This number is set by Burnin (see below). Note that the
		burnin setting is shared with the 'mcmc', 'sumt', 'sump' and 'plot' commands.
<b>D</b> 111	rnin	Determines the number of samples (not generations) that will
Dui		be discarded when summary statistics are calculated. The
		value of this option is only relevant when Relburnin is set
		to 'No'.
D	rninFrac	
Bur	rninfrac	Determines the fraction of samples that will be discarded
		when summary statistics are calculated. The value of this
		option is only relevant when Relburnin is set to 'Yes'.
		Example: A value for this option of 0.25 means that 25% of
		the samples will be discarded.
Mir	npartfreq	The minimum probability of partitions to include in summary
		statistics.
Fi]	lename1	The name of the first tree file to compare.
Fi?	lename2	The name of the second tree file to compare.

## Current settings:

Parameter	Options	Current Setting
Relburnin	Yes/No	Yes
Burnin	<number></number>	0
Burninfrac	<number></number>	0.25
Minpartfreq	<number></number>	0.00
Filename1	<name></name>	temp.t
Filename2	<name></name>	temp.t
Outputname	<name></name>	temp.comp

printed.

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#### Constraint

This command defines a tree constraint. The format for the constraint command is

constraint <name> [hard|negative|partial] = <taxon list> [:<taxon list>]

There are three types of constraint implemented in MrBayes. The type of the constraint is specified by using one of the three keywords 'hard', 'negative', or 'partial' right after the name of the constraint. If no type is specified, then the constraint is assumed to be 'hard'.

In a rooted tree, a 'hard' constraint forces the taxa in the list to form a monophyletic group. In an unrooted tree, the taxon split that separates the taxa in the list from other taxa is forced to be present. The interpretation of this depends on whether the tree is rooted on a taxon outside the list or a taxon in the list. If the outgroup is excluded, the taxa in the list are assumed to form a monophyletic group, but if the outgroup is included, it is the taxa that are not in the list that are forced together.

A 'negative' constraint bans all the trees that have the listed taxa in the same subtree. In other words, it is the opposite of a hard constraint.

A 'partial' or backbone constraint is defined in terms of two sets of taxa separated by a colon character. The constraint forces all taxa in the first list to form a monophyletic group that does not include any taxon in the second list. Taxa that are not included in either list can be placed in any position on the tree, either inside or outside the constrained group. In an unrooted tree, the two taxon lists can be switched with each other with no effect. For a rooted tree, it is the taxa in the first list that have to be monophyletic, that is, these taxa must share a common ancestor not shared with any taxon in the second list. The taxa in the second list may or may not fall in a monophyletic group depending on the rooting of the tree.

A list of taxa can be specified using a taxset, taxon names, taxon numbers, or any combination of the above, sepatated by spaces. The constraint is treated as an absolute requirement of trees, that is, trees that are not compatible with the constraint have zero prior (and hence zero posterior) probabilty.

If you are interested in inferring ancestral states for a particular node, you need to 'hard' constrain that node first using the 'constraint' command. The same applies if you wish to calibrate an interior node in a dated analysis. For more information on how to infer ancestral states, see the help for the 'report' command. For more on dating, see the 'calibrate' command.

It is important to note that simply defining a constraint using this command is not sufficient for the program to actually implement the constraint in an analysis. You must also enforce the constraints using 'prset topologypr = constraints (<list of constraints>)'. For more information on this, see the help on the 'prset' command.

## Examples:

constraint myclade = Homo Pan Gorilla

Defines a hard constraint forcing Homo, Pan, and Gorilla to form a monophyletic group or a split that does not include any other taxa.

constraint forbiddenclade negative = Homo Pan Gorilla

Defines a negative constraint that associates all trees where Homon, Pan, and Gorilla form a monophyletic group with zero posterior probability. In other words, such trees will not be sampled during MCMC.

constraint backbone partial = Homo Gorilla : Mus

Defines a partial constraint that keeps Mus outside of the clade defined by the most recent common ancestor of Homo and Gorilla. Other taxa are allowed to sit anywhere in the tree.

To define a more complex constraint tree, simply combine constraints into a list when issuing the 'prset topologypr' command.

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Ctype

This command sets the character ordering for standard-type data. The correct usage is:

ctype <ordering>:<characters>

The available options for the <ordering> specifier are:

unordered -- Movement directly from one state to another is

allowed in an instant of time.

ordered -- Movement is only allowed between adjacent characters.

For example, perhaps only between 0 <-> 1 and 1 <-> 2

for a three state character ordered as 0 - 1 - 2.

irreversible -- Rates of change for losses are 0.

The characters to which the ordering is applied is specified in manner that is identical to commands such as "include" or "exclude". For example,

ctype ordered: 10 23 45

defines charactes 10, 23, and 45 to be of type ordered. Similarly,

ctype irreversible: 54 - 67 71-92

defines characters 54 to 67 and characters 71 to 92 to be of type irreversible. You can use the "." to denote the last character, and "all" to denote all of the characters. Finally, you can use the specifier "\" to apply the ordering to every n-th character or you can use predefined charsets to specify the character.

Only one ordering can be used on any specific application of ctype. If you want to apply different orderings to different characters, then you need to use ctype multiple times. For example,

ctype ordered: 1-50

ctype irreversible: 51-100

sets characters 1 to 50 to be ordered and characters 51 to 100 to be irreversible.

The ctype command is only sensible with morphological (here called "standard") characters. The program ignores attempts to apply char-

acter	orderings	to	other	types	of	characters,	such	as	DNA	characters.	
											· – – –
Databi	reaks										

This command is used to specify breaks in your input data matrix. Your data may be a mixture of genes or a mixture of different types of data. Some of the models implemented by MrBayes account for nonindependence at adjacent characters. The autocorrelated gamma model, for example, allows rates at adjacent sites to be correlated. However, there is no way for such a model to tell whether two sites, adjacent in the matrix, are actually separated by many kilobases or megabases in the genome. The databreaks command allows you to specify such breaks. The correct usage is:

databreaks <br/> <break 1> <break 2> <break 3> ...

For example, say you have a data matrix of 3204 characters that include nucleotide data from three genes. The first gene covers characters 1 to 970, the second gene covers characters 971 to 2567, and the third gene covers characters 2568 to 3204. Also, let's assume that the genes are not directly adjacent to one another in the genome, as might be likely if you have mitochondrial sequences. In this case, you can specify breaks between the genes using:

databreaks 970 2567;

The first break, between genes one and two, is after character 970 and the second break, between genes two and three, is after character 2567.

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Delete

This command deletes taxa from the analysis. The correct usage is:

delete <name and/or number and/or taxset> ...

A list of the taxon names or taxon numbers (labelled 1 to ntax in the order in the matrix) or taxset(s) can be used. For example, the following:

delete 1 2 Homo\_sapiens

deletes taxa 1, 2, and the taxon labelled Homo\_sapiens from the analysis. You can also use "all" to delete all of the taxa. For example,

delete all

deletes all of the taxa from the analysis. Of course, a phylogenetic analysis that does not include any taxa is fairly uninteresting.

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Disclaimer

This command shows the disclaimer for the program. In short, the disclaimer states that the authors are not responsible for any silly things you may do to your computer or any unforseen but possibly nasty things the computer program may inadvertently do to you.

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Exclude
This command excludes characters from the analysis. The correct usage is
  exclude <number> <number> <number>
or
  exclude <number> - <number>
or
  exclude <charset>
or some combination thereof. Moreover, you can use the specifier "\" to
exclude every nth character. For example, the following
  exclude 1-100\3
would exclude every third character. As a specific example,
  exclude 2 3 10-14 22
excludes sites 2, 3, 10, 11, 12, 13, 14, and 22 from the analysis. Also,
  exclude all
excludes all of the characters from the analysis. Excluding all characters
does not leave you much information for inferring phylogeny.
Execute
This command executes a file called <file name>. The correct usage is:
  execute <file name>
For example,
  execute replicase.nex
would execute the file named "replicase.nex". This file must be in the
same directory as the executable.
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______
Help
This command provides useful information on the use of this program. The
correct usage is
  help
which gives a list of all available commands with a brief description of
each or
  help <command>
which gives detailed information on the use of <command>.
```

#### Include

This command includes characters that were previously excluded from the analysis. The correct usage is

include <number> <number> <number>

or

include <number> - <number>

or

include <charset>

or some combination thereof. Moreover, you can use the specifier "\" to include every nth character. For example, the following  $\[ \frac{1}{2} + \frac{1$ 

include  $1-100\3$ 

would include every third character. As a specific example,

include 2 3 10-14 22

includes sites 2, 3, 10, 11, 12, 13, 14, and 22 from the analysis. Also,

include all

includes all of the characters in the analysis. Including all of the characters (even if many of them are bad) is a very total-evidence-like thing to do. Doing this will make a certain group of people very happy. On the other hand, simply using this program would make those same people unhappy.

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Link

This command links model parameters across partitions of the data. The correct usage is:

link <parameter name> = (<all> or <partition list>)

The list of parameters that can be linked includes:

Tratio -- Transition/transversion rate ratio
Revmat -- Substitution rates of GTR model
Omega -- Nonsynonymous/synonymous rate ratio

Statefreq -- Character state frequencies

Shape -- Gamma shape parameter

Pinvar -- Proportion of invariable sites

Correlation -- Correlation parameter of autodiscrete gamma

Switchrates -- Switching rates for covarion model

Brlens -- Branch lengths of tree

Topology -- Topology of tree

Speciationrates -- Speciation rates for birth-death process Extinctionrates -- Extinction rates for birth-death process

Theta -- Parameter for coalescence process
Growthrate -- Growth rate of coalescence process

Aamodel -- Aminoacid rate matrix

Cpprate -- Rate of Compound Poisson Process (CPP)

Cppmultdev -- Standard dev. of CPP rate multipliers (log scale)

Cppevents -- CPP events

TK02var -- Variance increase in TK02 relaxed clock model
TK02branchrates -- Branch rates of TK02 relaxed clock model
Igrvar -- Variance increase in IGR relaxed clock model
Igrbrlens -- Branch lengths of IGR relaxed clock model

For example,

link shape=(all)

links the gamma shape parameter across all partitions of the data. You can use "showmodel" to see the current linking status of the characters. For more information on this command, see the help menu for link's converse, unlink ("help unlink");

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Loq

This command allows output to the screen to also be output to a file. The useage is:

log start/stop filename=<name> append/replace

The options are:

Start/Stop -- Starts or stops logging of output to file.

Append/Replace -- Either append to or replace existing file.

Filename -- Name of log file (currently, the name of the log

file is "log.out").

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Lset

This command sets the parameters of the likelihood model. The likelihood function is the probability of observing the data conditional on the phylogenetic model. In order to calculate the likelihood, you must assume a model of character change. This command lets you tailor the biological assumptions made in the phylogenetic model. The correct usage is

lset <parameter>=<option> ... <parameter>=<option>

For example, "lset nst=6 rates=gamma" would set the model to a general model of DNA substition (the GTR) with gamma-distributed rate variation across sites.

Options:

-- This option allows you to apply the lset commands to specific partitions. This command should be the first in the list of commands specified in lset. Moreover, it only makes sense to be using this command if the data have been partitioned. A default partition is set on execution of a matrix. If the data are homogeneous (i.e., all of the same data type), then this partition will not subdivide the characters. Up to 30 other partitions can be defined, and you can switch among them using "set partition=<partition name>". Now, you may want to specify different models to different partitions of the data. Applyto allows you to do this. For example, say you have partitioned the data by codon position, and you want to apply a nst=2 model to the first two partitions and nst=6 to the last. This could be implemented in two uses of lset:

lset applyto=(1,2) nst=2

lset applyto=(3) nst=6

The first applies the parameters after "applyto" to the first and second partitions. The second lset applies nst=6 to the third partition. You can also use applyto=(all), which attempts to apply the parameter settings to all of the data partitions. Importantly, if the option is not consistent with the data in the partition, the program will not apply the lset option to that partition.

Nucmodel

-- This specifies the general form of the nucleotide substitution model. The options are "4by4" [the standard model of DNA substitution in which there are only four states (A,C,G,T/U)], "doublet" (a model appropriate for modelling the stem regions of ribosomal genes where the state space is the 16 doublets of nucleotides), "codon" (the substitution model is expanded around triplets of nucleotides -- a codon), and "Protein" (triplets of nucleotides are translated to amino acids, which form the basis of the substitution model).

Nst

-- Sets the number of substitution types: "1" constrains all of the rates to be the same (e.g., a JC69 or F81 model); "2" allows transitions and transversions to have potentially different rates (e.g., a K80 or HKY85 model); "6" allows all rates to be different, subject to the constraint of time-reversibility (e.g., a GTR model). Finally, 'nst' can be set to 'mixed', which results in the Markov chain sampling over the space of all possible reversible substitution models, including the GTR model and all models that can be derived from it model by grouping the six rates in various combinations. This includes all the named models above and a large number of others, with or without name.

Code

-- Enforces the use of a particular genetic code. The default is the universal code. Other options include "vertmt" for vertebrate mitocondrial DNA, "mycoplasma", "yeast", "ciliates", and "metmt" (for metazoan mitochondrial DNA except vertebrates).

Ploidy

-- Specifies the ploidy of the organism. Options are "Haploid", "Diploid" or "Zlinked". This option is used when a coalescence prior is used on trees.

Rates

- -- Sets the model for among-site rate variation. In general, the rate at a site is considered to be an unknown random variable. The valid options are:
  - \* equal -- No rate variation across sites.
  - \* gamma -- Gamma-distributed rates across sites. The rate at a site is drawn from a gamma distribution. The gamma distribution has a single parameter that describes how much rates vary.
  - -- Autocorrelated rates across sites. The marginal rate distribution is gamma, but adjacent sites have correlated rates.
  - \* propinv -- A proportion of the sites are invariable.
  - \* invgamma -- A proportion of the sites are invariable while the rate for the remaining sites are drawn from a gamma distribution.

Note that MrBayes versions 2.0 and earlier supported options that allowed site specific rates (e.g., ssgamma). In versions 3.0 and later, site specific rates are allowed, but set using the 'prset ratepr' command for each partition.

Ngammacat -- Sets the number of rate categories for the gamma distribution. The gamma distribution is continuous. However, it is virtually impossible to calculate likelihoods under the continuous gamma distribution. Hence, an approximation to the continuous gamma is used; the gamma distribution is broken into ncat categories of equal weight (1/ncat). The mean rate for each category represents the rate for the entire category. This option allows you to specify how many rate categories to use when approximating the gamma. The approximation is better as ncat is increased. In practice, "ncat=4" does a reasonable job of approximating the continuous gamma.

Nbetacat

- -- Sets the number of rate categories for the beta distribution. A symmetric beta distribution is used to model the stationary frequencies when morphological data are used. This option specifies how well the beta distribution will be approximated.
- Omegavar -- Allows the nonsynonymous/synonymous rate ratio (omega) to vary across codons. Ny98 assumes that there are three classes, with potentially different omega values (omega1, omega2, omega3): omega2 = 1; 0 < omega1 <1; and omega3 > 1. Like the Ny98 model, the M3 model has three omega classes. However, their values are less constrained, with omega1 < omega2 < omega3. The default (omegavar = equal) has no variation on omega across sites.
- Covarion -- This forces the use of a covarion-like model of substitution for nucleotide or amino acid data. The valid options are "yes" and "no". The covarion model allows the rate at a site to change over its evolutionary history. Specifically, the site is either on or off. When it is off, no substitutions are possible. When the process is on, substitutions occur according to a specified substitution model (specified using the other lset options).
- Coding -- This specifies how characters were sampled. If all site patterns had the possibility of being sampled, then "all" should be specified (the default). Otherwise "variable" (only variable characters had the possibility of being sampled), "noabsence" (characters for which all taxa were coded as absent were not sampled), and "nopresence" (characters for which all taxa were coded as present were not sampled. "All" works for all data types. However, the others only work for morphological (all/variable) or restriction site (all/variable/noabsence/nopresence) data.
- Parsmodel -- This forces calculation under the so-called parsimony model described by Tuffley and Steel (1998). The options are "yes" or "no". Note that the biological assumptions of this model are anything but parsimonious. In fact, this model assumes many more parameters than the next most complicated model implemented in this program. If you really believe that the parsimony model makes the biological assumptions described by Tuffley and Steel, then the parsimony method is miss-named.

## Default model settings:

Parameter	Options	Current Setting
Nucmodel Nst	4by4/Doublet/Codon/Protein 1/2/6/Mixed	4by4 1
Code	Universal/Vertmt/Mycoplasma/	
	Yeast/Ciliates/Metmt	Universal
Ploidy	Haploid/Diploid/Zlinked	Diploid
Rates	Equal/Gamma/Propinv/Invgamma/Adgamma	Equal
Ngammacat	<number></number>	4
Nbetacat	<number></number>	5
Omegavar	Equal/Ny98/M3	Equal

Covarion No/Yes No

Coding All/Variable/Noabsencesites/

Nopresencesites All Parsmodel No/Yes No

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#### Manual

This command allows you to generate a text file containing help information on all the available commands. This text file can be used as an up-to-date command reference. You can set the name of the text file using the "filename" option; the default is "commref\_mb<version>.txt".

Parameter Options Current Setting
-----Filename <name> commref\_mb3.2.0.txt

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#### Mcmc

This command starts the Markov chain Monte Carlo (MCMC) analysis. The posterior probability of phylogenetic trees (and other parameters of the substitution model) cannot be determined analytically. Instead, MCMC is used to approximate the posterior probabilities of trees by drawing (dependent) samples from the posterior distribution. This program can implement a variant of MCMC called "Metropolis-coupled Markov chain Monte Carlo", or MCMCMC for short. Basically, "Nchains" are run, with Nchains - 1 of them heated. The chains are labelled 1, 2, ..., Nchains. The heat that is applied to the i-th chain is B = 1 / (1 + temp X i). B is the power to which the posterior probability is raised. When B = 0, all trees have equal probability and the chain freely visits trees. B = 1 is the "cold" chain (or the distribution of interest). MCMCMC can mix better than ordinary MCMC; after all of the chains have gone through one cycle, two chains are chosen at random and an attempt is made to swap the states (with the probability of a swap being determined by the Metropolis et al. equation). This allows the chain to potentially jump a valley in a single bound. The correct usage is

mcmc <parameter> = <value> ... <parameter> = <value>

For example,

mcmc ngen=100000 nchains=4 temp=0.5

performs a MCMCMC analysis with four chains with the temperature set to  $0.5.\ \mathrm{The}$  chains would be run for  $100,000\ \mathrm{cycles}.$ 

## Options:

Ngen -- This option sets the number of cycles for the MCMC algorithm. This should be a big number as you want the chain to first reach stationarity, and then remain there for

enough time to take lots of samples.

Nruns -- How many independent analyses are started simultaneously.

Nchains -- How many chains are run for each analysis for the MCMCMC variant. The default is 4: 1 cold chain and 3 heated chains.

If Nchains is set to 1, MrBayes will use regular MCMC sampling, without heating.

Temp -- The temperature parameter for heating the chains. The higher

the temperature, the more likely the heated chains are to move between isolated peaks in the posterior distribution. However, excessive heating may lead to very low acceptance rates for swaps between different chains. Before changing the default setting, however, note that the acceptance rates of swaps tend to fluctuate during the burn-in phase of the run.

Reweight

-- Here, you specify three numbers, that respectively represent the percentage of characters to decrease in weight, the percentage of characters to increase in weight, and the increment. An increase/decrease in weight is acheived by replicating/removing a character in the matrix. This is only done to non-cold chains. The format for this parameter is "reweight=(<number>,<number>)" or "reweight=(<number>, <number>,<number>)".

Swapfreq

-- This specifies how often swaps of states between chains are attempted. You must be running at least two chains for this option to be relevant. The default is Swapfreq=1, resulting in Nswaps (see below) swaps being tried each generation of the run. If Swapfreg is set to 10, then Nswaps swaps will be tried every tenth generation of the run.

Nswaps

-- The number of swaps tried for each swapping generation of the chain (see also Swapfreg).

Samplefreq

-- This specifies how often the Markov chain is sampled. You can sample the chain every cycle, but this results in very large output files. Thinning the chain is a way of making these files smaller and making the samples more independent.

Printfreq

-- This specifies how often information about the chain is printed to the screen.

Printall

-- If set to NO, only cold chains in a MCMC analysis are printed to screen. If set to YES, both cold and heated chains will be output. This setting only affects the printing to screen, it does not change the way values are written to file.

Printmax

-- The maximum number of chains to print to screen.

Mcmcdiagn

-- Determines whether acceptance ratios of moves and swaps will be printed to file. The file will be named similarly to the '.p' and '.t' files, but will have the ending '.mcmc'. If more than one independent analysis is run simultaneously (see Nruns below), convergence diagnostics for tree topology will also be printed to this file. The convergence diagnostic used is the average standard deviation in partition frequency values across independent analyses. The Burnin setting (see below) determines how many samples will be discarded as burnin before calculating the partition frequencies. The Minpartfreq setting (see below) determines the minimum partition frequency required for a partition to be included in the calculation. As the independent analyses approach stationarity (converge), the value of the diagnostic is expected to approach zero.

Diagnfreq

-- The number of generations between the calculation of MCMC diagnostics (see Mcmcdiagn above).

Diagnstat

-- The statistic to use for run-time convergence diagnostics. Choices are 'Avgstddev' for average standard deviation of split frequencies and 'Maxstddev' for maximum standard deviation of split frequencies.

Savetrees

-- If you are using a relative burnin for run-time convergence diagnostics, tree samples need to be deleted from split frequency counters as the cut-off point for the burnin moves during the run. If 'Savetrees' is set to 'No', tree samples to be discarded are read back in from file. If 'Savetrees' is set to 'Yes', the tree samples to be removed will be stored in the internal memory instead. This can use up a lot of memory in large analyses.

Minpartfreq -- The minimum frequency required for a partition to be included in the calculation of the topology convergence diagnostic. The partition is included if the minimum frequency is reached in at least one of the independent tree samples that are compared. -- If this option is set to YES, acceptance ratios for moves are Allchains recorded for all chains, cold or heated. By default, only the acceptance ratios for the cold chain are recorded. -- If this option is set to YES, topological convergence diag-Allcomps nostics are calculated over all pairwise comparisons of runs. If it is set to NO, only the overall value is reported. -- If this option is set to YES, then a proportion of the sampled Relburnin values will be discarded as burnin when calculating the convergence diagnostic. The proportion to be discarded is set with Burninfrac (see below). When the Relburnin option is set to NO, then a specific number of samples will be discarded instead. This number is set by Burnin (see below). Burnin -- Determines the number of samples (not generations) that will be discarded when convergence diagnostics are calculated. The value of this option is only relevant when Relburnin is set to NO. BurninFrac -- Determines the fraction of samples that will be discarded when convergence diagnostics are calculated. The value of this option is only relevant when Relburnin is set to YES. Example: A value for this option of 0.25 means that 25% of the samples will be discarded. Stoprule -- If this option is set to NO, then the chain is run the number of generations determined by Ngen. If it is set to YES, and topological convergence diagnostics are calculated (Mcmcdiagn is set to YES), then the chain will be stopped before the predetermined number of generations if the convergence diagnostic falls below the stop value. -- The critical value for the topological convergence diagnostic. Stopval Only used when Stoprule and Mcmcdiagn are set to yes, and more than one analysis is run simultaneously (Nruns > 1). -- If this parameter is set to 'Yes', all the current parameter Checkpoint values of all chains will be printed to a check-pointing file every 'Checkfreq' generation of the analysis. The file will be named <Filename>.ckp and allows you to restart the analysis from the last check point. This can be handy if you are running a long analysis and want to extend it, or if there is a risk that a long analysis will be inadvertently interupted by hardware failure or other factors that are out of your control. Checkfreq -- The number of generations between check-pointing. See the 'Checkpoint' parameter above for more information. Filename -- The name of the files that will be generated. Two files are generated: "<Filename>.t" and "<Filename>.p". The .t file contains the trees whereas the .p file contains the sampled values of the parameters. -- The starting values for the model parameters are set to Startparams arbitrary or random values when the parameters are created. These starting values can be altered using the 'Startvals' command. The 'Startparams=reset' option allows you to reset the starting values to the default at the start of the analysis, overriding any previous user-defined starting values. Under the default option, 'current', the chains will use the current starting values. -- The starting tree(s) for the chain can either be randomly Starttree

selected or user-defined. It might be a good idea to start from randomly chosen trees; convergence seems

likely if independently run chains, each of which started from different random trees, converge to the same answer. If you want the chain to start from user-defined trees instead, you first need to read in your tree(s) from a Nexus file with a 'trees' block, and then you need to set the starting tree(s) using the 'Startvals' command. Finally, you need to make sure that 'Starttree' is set to 'current'. If you do not set the starting tree(s), the chains will start with random trees. Setting 'Starttree' to 'random' causes new starting trees to be drawn randomly at the start of the run, overwriting any previous user-defined starting trees.

Nperts

-- This is the number of random perturbations to apply to the user starting tree. This allows you to have something between completely random and user-defined trees start the chain.

Savebrlens

-- This specifies whether branch length information is saved on the trees.

Data

-- When Data is set to NO, the chain is run without data. This should be used only for examining induced priors. DO NOT SET 'DATA' TO 'NO' UNLESS YOU KNOW WHAT YOU ARE DOING!

Ordertaxa

-- Determines whether taxa should be ordered before trees are printed to file. If set to 'Yes', terminals in the sampled trees will be reordered to match the order of the taxa in the data matrix as closely as possible. By default, trees will be printed without reordering of taxa.

**Append** 

-- Set this to 'Yes' to append the results of the current run to a previous run. MrBayes will first read in the results of the previous run (number of generations and sampled splits) and will then continue that run where you left it off. Make sure that the output file names used in the previous run are the same as those in the current run.

Autotune

-- Set this to 'Yes' to autotune the proposals that change substitution model parameters. When set to 'No', the tuning parameters are fixed to their starting values. Note that the autotuning occurs independently for each chain. The target acceptance rate for each move can be changed using the 'Propset' command.

Tunefreq

-- When a proposal has been tried 'Tunefreq' times, its tuning parameter is adjusted to reach the target acceptance rate if 'Autotune' is set to 'Yes'.

Parameter	Options	Current Setting	
Ngen	<number></number>	1000000	
Nruns	<number></number>	2	
Nchains	<number></number>	4	
Temp	<number></number>	0.100000	
Reweight	<number>,<number></number></number>	0.00 v 0.00 ^	
Swapfreq	<number></number>	1	
Nswaps	<number></number>	1	
Samplefreq	<number></number>	500	
Printfreq	<number></number>	500	
Printall	Yes/No	Yes	
Printmax	<number></number>	8	
Mcmcdiagn	Yes/No	Yes	
Diagnfreq	<number></number>	5000	
Diagnstat	Avgstddev/Maxstddev	Avgstddev	
Minpartfreq	<number></number>	0.10	
Allchains	Yes/No	No	
Allcomps	Yes/No	No	
Relburnin	Yes/No	Yes	

Burnin	<number></number>	0
Burninfrac	<number></number>	0.25
Stoprule	Yes/No	No
Stopval	<number></number>	0.05
Savetrees	Yes/No	No
Checkpoint	Yes/No	Yes
Checkfreq	<number></number>	100000
Filename	<name></name>	temp.out. <p t=""></p>
Startparams	Current/Reset	Current
Starttree	Current/Random/	Current
	Parsimony	
Nperts	<number></number>	0
Data	Yes/No	Yes
Ordertaxa	Yes/No	No
Append	Yes/No	No
Autotune	Yes/No	Yes
Tunefreq	<number></number>	100

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### Mcmcp

This command sets the parameters of the Markov chain Monte Carlo (MCMC) analysis without actually starting the chain. This command is identical in all respects to Mcmc, except that the analysis will not start after this command is issued. For more details on the options, check the help menu for Mcmc.

Parameter	Options	Current Setting	
Ngen	<number></number>	1000000	
Nruns	<number></number>	2	
Nchains	<number></number>	4	
Temp	<number></number>	0.100000	
Reweight	<number>,<number></number></number>	0.00 v 0.00 ^	
Swapfreq	<number></number>	1	
Nswaps	<number></number>	1	
Samplefreq	<number></number>	500	
Printfreq	<number></number>	500	
Printall	Yes/No	Yes	
Printmax	<number></number>	8	
Mcmcdiagn	Yes/No	Yes	
Diagnfreq	<number></number>	5000	
Diagnstat	Avgstddev/Maxstddev	Avgstddev	
Minpartfreq	<number></number>	0.10	
Allchains	Yes/No	No	
Allcomps	Yes/No	No	
Relburnin	Yes/No	Yes	
Burnin	<number></number>	0	
Burninfrac	<number></number>	0.25	
Stoprule	Yes/No	No	
Stopval	<number></number>	0.05	
Savetrees	Yes/No	No	
Checkpoint	Yes/No	Yes	
Checkfreq	<number></number>	100000	
Filename	<name></name>	temp.out. <p t=""></p>	
Startparams	Current/Reset	Current	
Starttree	Current/Random/	Current	
	Parsimony		
Nperts	<number></number>	0	
Data	Yes/No	Yes	

Ordertaxa Yes/No No
Append Yes/No No
Autotune Yes/No Yes
Tunefreq <number> 100

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Outgroup

This command assigns a taxon to the outgroup. The correct usage is:

outgroup <number>/<taxon name>

For example, "outgroup 3" assigns the third taxon in the matrix to be the outgroup. Similarly, "outgroup Homo\_sapiens" assings the taxon "Homo\_sapiens" to be the outgroup (assuming that there is a taxon named "Homo\_sapiens" in the matrix). Only a single taxon can be assigned to be the outgroup.

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Pairs

This command is used to specify pairs of nucleotides. For example, your data may be RNA sequences with a known secondary structure of stems and loops. Substitutions in nucleotides involved in a Watson-Crick pairing in stems are not strictly independent; a change in one changes the probability of a change in the partner. A solution to this problem is to expand the model around the pair of nucleotides in the stem. This command allows you to do this. The correct usage is:

pairs <NUC1>:<NUC2>, <NUC1>:<NUC2>,..., <NUC1>:<NUC2>;

For example,

pairs 30:56, 31:55, 32:54, 33:53, 34:52, 35:51, 36:50;

specifies pairings between nucleotides 30 and 56, 31 and 55, etc. Only nucleotide data (DNA or RNA) may be paired using this command. Note that in order for the program to actually implement a "doublet" model involving a 16 X 16 rate matrix, you must specify that the structure of the model is 16 X 16 using "lset nucmodel=doublet".

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Partition

This command allows you to specify a character partition. The format for this command is

partition <name> = <num parts>:<chars in first>, ...,<chars in last>

For example, "partition by\_codon = 3:1st\_pos,2nd\_pos,3rd\_pos" specifies a partition called "by\_codon" which consists of three parts (first, second, and third codon positions). Here, we are assuming that the sites in each partition were defined using the charset command. You can specify a partition without using charset as follows:

partition by\_codon = 3:1 4 6 9 12,2 5 7 10 13,3 6 8 11 14

However, we recommend that you use the charsets to define a set of characters and then use these predefined sets when defining the partition. Also, it makes more sense to define a partition as a line in the mrbayes

block than to issue the command from the command line (then again, you may be a masochist, and want to do extra work).

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Plot

This command plots specified parameters in the .p file or one of the .p files created during an MCMC analysis. An x-y graph of the parameter over the course of the chain is created. The command can be useful for visually diagnosing convergence for many of the parameters of the phylogenetic model. The parameter to be plotted is specified by the "parameter" option. Several parameters can be plotted at once by using the "match" option, which has a default value of "perfect". For example, if you were to set "parameter = pi" and "match = consistentwith", then all of the state frequency parameters would be plotted. You can also set "match=all", in which case all of the parameters are plotted.

Note that the "Sump" command provides a different set of convergence diagnostics tools that you may also want to explore. Unlike "Plot", "Sump" can compare two or more parameter samples and will calculate convergence diagnostics as well as parameter summaries for the pooled sample.

#### Options:

Relburnin	If this option is set to 'Yes', then a proportion of the samples will be discarded as burnin when creating the plot. The proportion to be discarded is set with Burninfrac (see Burninfrac below). When the Relburnin option is set to 'No', then a specific number of samples is discarded instead. This number is set by Burnin (see below). Note that the burnin setting is shared across the 'comparetree', 'sump' and 'sumt' commands.
Burnin	Determines the number of samples (not generations) that will be discarded when summary statistics are calculated. The value of this option is only relevant when Relburnin is set to 'No'.
Burninfrac	Determines the fraction of samples that will be discarded when creating a plot. The value of this parameter is only relevant when Relburnin is set to 'Yes'. Example: A value of

this option of 0.25 means that 25% of the samples will be discarded. Filename  $\,$  -- The name of the file to plot.

Parameter -- Specification of parameters to be plotted. See above for

details.

 $\hbox{\tt Match} \qquad \quad \hbox{\tt -- Specifies how to match parameter names to the Parameter}$ 

specification. See above for details.

### Current settings:

Parameter	Options	Current Setting
Relburnin Burnin Burninfrac Filename Parameter Match	Yes/No <number> <number> <name> <name> Perfect/Consistentwith/All</name></name></number></number>	Yes 0 0.25 temp.p lnL Perfect

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#### Prset

This command sets the priors for the phylogenetic model. Remember that in a Bayesian analysis, you must specify a prior probability distribution for the parameters of the likelihood model. The prior distribution represents your prior beliefs about the parameter before observation of the data. This command allows you to tailor your prior assumptions to a large extent.

### Options:

Applyto

-- This option allows you to apply the prset commands to specific partitions. This command should be the first in the list of commands specified in prset. Moreover, it only makes sense to be using this command if the data have been partitioned. A default partition is set on execution of a matrix. If the data are homogeneous (i.e., all of the same data type), then this partition will not subdivide the characters. Up to 30 other partitions can be defined, and you can switch among them using "set partition=<partition name>". Now, you may want to specify different priors to different partitions of the data. Applyto allows you to do this. For example, say you have partitioned the data by codon position, and you want to fix the statefregs to equal for the first two partitions but apply a flat Dirichlet prior to the statefreqs of the last. This could be implemented in two uses of prset:

```
prset applyto=(1,2) statefreqs=fixed(equal)
prset applyto=(3) statefreqs=dirichlet(1,1,1,1)
```

The first applies the parameters after "applyto" to the first and second partitions. The second prset applies a flat Dirichlet to the third partition. You can also use applyto=(all), which attempts to apply the parameter settings to all of the data partitions. Importantly, if the option is not consistent with the data in the partition, the program will not apply the prset option to that partition.

Tratiopr

-- This parameter sets the prior for the transition/transversion rate ratio (tratio). The options are:

```
prset tratiopr = beta(<number>, <number>)
prset tratiopr = fixed(<number>)
```

The program assumes that the transition and transversion rates are independent gamma-distributed random variables with the same scale parameter when beta is selected. If you want a diffuse prior that puts equal emphasis on transition/transversion rate ratios above 1.0 and below 1.0, then use a flat Beta, beta(1,1), which is the default. If you wish to concentrate this distribution more in the equal-rates region, then use a prior of the type beta(x,x), where the magnitude of x determines how much the prior is concentrated in the equal rates region. For instance, a beta(20,20) puts more probability on rate ratios close to 1.0 than a beta(1,1). If you think it is likely that the transition/transversion rate ratio is 2.0, you can use a prior of the type beta(2x,x), where x determines how strongly the prior is concentrated on

tratio values near 2.0. For instance, a beta(2,1) is much more diffuse than a beta(80,40) but both have the expected tratio 2.0 in the absence of data. The parameters of the Beta can be interpreted as counts: if you have observed x transitions and y transversions, then a beta(x+1,y+1) is a good representation of this information. The fixed option allows you to fix the tratio to a particular value.

Revmatpr

-- This parameter sets the prior for the substitution rates of the GTR model for nucleotide data. The options are:

```
prset revmatpr = dirichlet(<number>,<number>,...,<number>)
prset revmatpr = fixed(<number>,<number>,...,<number>)
```

The program assumes that the six substitution rates are independent gamma-distributed random variables with the same scale parameter when dirichlet is selected. The six numbers in brackets each corresponds to a particular substitution type. Together, they determine the shape of the prior. The six rates are in the order A<->C, A<->G, A<->T, C<->G, C<->T, and G<->T. If you want an uninformative prior you can use dirichlet(1,1,1,1,1,1), also referred to as a 'flat' Dirichlet. This is the default setting. If you wish a prior where the C<->T rate is 5 times and the A<->G rate 2 times higher, on average, than the transversion rates, which are all the same, then you should use a prior of the form dirichlet(x,2x,x,x,5x,x), where x determines how much the prior is focused on these particular rates. For more info, see tratiopr. The fixed option allows you to fix the substitution rates to particular values.

Revratepr

-- This parameter sets the prior for each substitution rate of the GTR model subspace when 'nst' is set to 'mixed' (see the 'lset' command). The only option is

```
prset revratepr = symdir(<number>)
```

which will associate each independent rate in the rate matrix with a modified symmetric Dirichlet prior, where a singleton rate has the specified alpha parameter, while a rate that applies to n pairwise substitution types has an alpha that is n times the specified number. The higher the specified number, the more focused the prior will be on equal rates. The default value is 1, which gives an effect similar to a flat Dirichlet.

Aamodelpr

-- This parameter sets the rate matrix for amino acid data. You can either fix the model by specifying aamodelpr= fixed(<model name>), where <model name> is 'poisson' (a glorified Jukes-Cantor model), 'jones', 'dayhoff', 'mtrev', 'mtmam', 'wag', 'rtrev', 'cprev', 'vt', 'blosum', 'equalin' (a glorified Felsenstein 1981 model), or 'gtr'. You can also average over the first ten models by specifying aamodelpr= mixed. If you do so, the Markov chain will sample each model according to its probability. The sampled model is reported as an index: poisson(0), jones(1), dayhoff(2), mtrev(3), mtmam(4), wag(5), rtrev(6), cprev(7), vt(8), or blosum(9). The 'Sump' command summarizes the MCMC samples and calculates the posterior probability estimate for each of these models.

Aarevmatpr

-- This parameter sets the prior for the substitution rates of the GTR model for amino acid data. The options are:

prset revmatpr = dirichlet(<number>,<number>,...,<number>)
prset revmatpr = fixed(<number>,<number>,...,<number>)

The options are the same as those for 'Revmatpr' except that they are defined over the 190 rates of the time-reversible GTR model for amino acids instead of over the 6 rates of the GTR model for nucleotides. The rates are in the order A<->R, A<->N, etc to Y<->V. In other words, amino acids are listed in alphabetic order based on their full name. The first amino acid (Alanine) is then combined in turn with all amino acids following it in the list, starting with amino acid 2 (Arginine) and finishing with amino acid 20 (Valine). The second amino acid (Arginine) is then combined in turn with all amino acids following it, starting with amino acid 3 (Asparagine) and finishing with amino acid 20 (Valine), and so on.

Omegapr

-- This parameter specifies the prior on the nonsynonymous/ synonymous rate ratio. The options are:

```
prset omegapr = uniform(<number>,<number>)
prset omegapr = exponential(<number>)
prset omegapr = fixed(<number>)
```

This parameter is only in effect if the nucleotide substitution model is set to codon using the 1set command (lset nucmodel=codon). Moreover, it only applies to the case when there is no variation in omega across sites (i.e., "lset omegavar=equal").

Ny98omega1pr

-- This parameter specifies the prior on the nonsynonymous/ synonymous rate ratio for sites under purifying selection. The options are:

```
prset Ny98omega1pr = beta(<number>,<number>)
prset Ny98omegalpr = fixed(<number>)
```

This parameter is only in effect if the nucleotide substitution model is set to codon using the 1set command (lset nucmodel=codon). Moreover, it only applies to the case where omega varies across sites using the model of Nielsen and Yang (1998) (i.e., "lset omegavar=ny98"). If fixing the parameter, you must specify a number between 0 and 1.

Ny98omega3pr -- This parameter specifies the prior on the nonsynonymous/ synonymous rate ratio for positively selected sites. The options are:

```
prset Ny98omega3pr = uniform(<number>,<number>)
prset Ny98omega3pr = exponential(<number>)
prset Ny98omega3pr = fixed(<number>)
```

This parameter is only in effect if the nucleotide substitution model is set to codon using the lset command (lset nucmodel=codon). Moreover, it only applies to the case where omega varies across sites according to the NY98 model. Note that if the NY98 model is specified that this parameter must be greater than 1, so you should not specify a uniform(0,10) prior, for example.

M3omegapr

-- This parameter specifies the prior on the nonsynonymous/ synonymous rate ratios for all three classes of sites for the M3 model. The options are:

```
prset M3omegapr = exponential
prset M3omegapr = fixed(<number>,<number>,<number>)
```

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This parameter is only in effect if the nucleotide substitution model is set to codon using the lset command (lset nucmodel=codon). Moreover, it only applies to the case where omega varies across sites using the M3 model of Yang et al. (2000) (i.e., "lset omegavar=M3"). Under the exponential prior, the four rates (dN1, dN2, dN3, and dS) are all considered to be independent draws from the same exponential distribution (the parameter of the exponential does not matter, and so you don't need to specify it). The rates dN1, dN2, and dN3 are taken to be the order statistics with dN1 < dN2 < dN3. These three rates are all scaled to the same synonymous rate, dS. The other option is to simply fix the three rate ratios to some values.

Codoncatfreqs -- This parameter specifies the prior on frequencies of sites under purifying, neutral, and positive selection. The options are:

```
prset codoncatfregs = dirichlet(<num>,<num>,<num>)
prset codoncatfreqs = fixed(<number>,<number>,<number>)
```

This parameter is only in effect if the nucleotide substitution model is set to codon using the 1set command (lset nucmodel=codon). Moreover, it only applies to the case where omega varies across sites using the models of Nielsen and Yang (1998) (i.e., "lset omegavar=ny98") or Yang et al. (2000) (i.e., "lset omegavar=M3") Note that the sum of the three frequencies must be 1.

Statefreqpr

-- This parameter specifies the prior on the state frequencies. The options are:

```
prset statefreqpr = dirichlet(<number>)
prset statefreqpr = dirichlet(<number>,...,<number>)
prset statefreqpr = fixed(equal)
prset statefreqpr = fixed(empirical)
prset statefreqpr = fixed(<number>,...,<number>)
```

For the dirichlet, you can specify either a single number or as many numbers as there are states. If you specify a single number, then the prior has all states equally probable with a variance related to the single parameter passed in.

Shapepr

-- This parameter specifies the prior for the gamma shape parameter for among-site rate variation. The options are:

```
prset shapepr = uniform(<number>,<number>)
prset shapepr = exponential(<number>)
prset shapepr = fixed(<number>)
```

-- This parameter specifies the prior for the proportion of invariable sites. The options are:

```
prset pinvarpr = uniform(<number>,<number>)
prset pinvarpr = fixed(<number>)
```

Note that the valid range for the parameter is between 0 and 1. Hence, "prset pinvarpr=uniform(0,0.8)" is valid while "prset pinvarpr=uniform(0,10)" is not. The default setting is "prset pinvarpr=uniform(0,1)".

Ratecorrpr

-- This parameter specifies the prior for the autocorrelation parameter of the autocorrelated gamma distribution for among-site rate variation. The options are:

```
prset ratecorrpr = uniform(<number>,<number>)
prset ratecorrpr = fixed(<number>)
```

Note that the valid range for the parameter is between -1 and 1. Hence, "prset ratecorrpr=uniform(-1,1)" is valid while "prset ratecorrpr=uniform(-11,10)" is not. The default setting is "prset ratecorrpr=uniform(-1,1)".

Covswitchpr

-- This option sets the prior for the covarion switching rates. The options are:

```
prset covswitchpr = uniform(<number>,<number>)
prset covswitchpr = exponential(<number>)
prset covswitchpr = fixed(<number>,<number>)
```

The covarion model has two rates: a rate from on to off and a rate from off to on. The rates are assumed to have independent priors that individually are either uniformly or exponentially distributed. The other option is to fix the switching rates, in which case you must specify both rates. (The first number is off->on and the second is on->off).

Symdirihyperpr - This option sets the prior for the stationary frequencies of the states for morphological (standard) data. There can be as many as 10 states for standard data. However, the labelling of the states is somewhat arbitrary. For example, the state "1" for different characters does not have the same meaning. This is not true for DNA characters, for example, where a "G" has the same meaning across characters. The fact that the labelling of morphological characters is arbitrary makes it difficult to allow unequal characterstate frequencies. MrBayes gets around this problem by assuming that the states have a dirichlet prior, with all states having equal frequency. The variation in the dirichlet can be controlled by this parameter--symdirihyperpr. Symdirihyperpr specifies the distribution on the variance parameter of the dirichlet. The valid options are:

```
prset Symdirihyperpr = uniform(<number>,<number>)
prset Symdirihyperpr = exponential(<number>)
prset Symdirihyperpr = fixed(<number>)
prset Symdirihyperpr = fixed(infinity)
```

Topologypr

If "fixed(infinity)" is chosen, the dirichlet prior is fixed such that all character states have equal frequency. -- This parameter specifies the prior probabilities of phylogenies. The options are:

```
prset topologypr = uniform
prset topologypr = speciestree
prset topologypr = constraints(<list>)
prset topologypr = fixed(<treename>)
```

If the prior is selected to be "uniform", the default, then all possible trees are considered a priori equally probable. The 'speciestree' option is used when the topology is constrained to fold inside a species tree together with other (gene) trees. The constraints option allows you to specify complicated prior probabilities on trees (constraints are discussed more fully in "help constraint"). Note that you must specify a list of constraints that you wish to be

obeyed. The list can be either the constraints' name or number. Finally, you can fix the topology to that of a user tree defined in a trees block. Branch lengths will still be sampled as usual on the fixed topology.

Brlenspr

-- This parameter specifies the prior probability distribution on branch lengths. The options are:

```
prset brlenspr = unconstrained:uniform(<num>,<num>)
prset brlenspr = unconstrained:exponential(<number>)
prset brlenspr = clock:uniform
prset brlenspr = clock:birthdeath
prset brlenspr = clock:coalescence
prset brlenspr = clock:speciestreecoalescence
prset brlenspr = fixed(<treename>)
```

Trees with unconstrained branch lengths are unrooted whereas clock-constrained trees are rooted. The option after the colon specifies the details of the probability density of branch lengths. If you choose a birth-death or coalescence prior, you may want to modify the details of the parameters of those processes (speciation rate, extinction rate and sample probability for the birth-death prior; population size and clock rate parameter for the coalescence prior). When gene trees are constrained to fold inside species trees, the appropriate branch length prior is 'clock:speciestreecoalescence'. Under this model, it is possible to control whether the population size is constant or varies across the species tree using the 'popvarpr' setting. Branch lengths can also be fixed but only if the topology is fixed.

Treeagepr

-- This parameter specifies the prior probability distribution on the tree age when a uniform prior is used on the branch lengths of a clock tree.

The options are:

```
prset treeagepr = Gamma(<num>,<num>)
prset treeagepr = Exponential(<number>)
prset treeagepr = Fixed(<number>)
```

(And, yes, we know the exponential is a special case of the gamma distribution.) The tree age is simply the age of the most recent common ancestor of the tree. If the clock rate is fixed to 1.0, which is the default, the tree age is equivalent to the expected number of substitutions from the root to the tip of the tree. The tree age prior ensures that the joint prior probability distribution for the uniform prior model of branch lengths on a clock tree is proper. The default setting is 'Exponential(1)'. If the root node in the tree is calibrated, the root calibration replaces the tree age prior.

Speciationpr -- This parameter sets the prior on the net speciation rate, that is, lambda - mu in the birth-death model. Options are:

```
prset speciationpr = uniform(<number>,<number>)
prset speciationpr = exponential(<number>)
prset speciationpr = fixed(<number>)
```

This parameter is only relevant if the birth-death process is selected as the prior on branch lengths.

Extinctionpr -- This parameter sets the prior on the relative extinction

rate, that is, mu / lambda in the birth-death model; values of this parameter are in the range (0,1). Prior options are:

prset extinctionpr = beta(<number>,<number>)
prset extinctionpr = fixed(<number>)

This parameter is only relevant if the birth-death process is selected as the prior on branch lengths.

SampleStrat

-- This parameter sets the strategy under which species where sampled in the analysis. This is used with the birth-death prior on trees (see Höhna et al, 2011).

Sampleprob

-- This parameter sets the fraction of species that are sampled in the analysis. This is used with the birth-death prior on trees (see Yang and Rannala, 1997).

Popsizepr

-- This parameter sets the prior on the population size component of the coalescent parameter. The options are:

prset popsizepr = uniform(<number>,<number>)
prset popsizepr = exponential(<number>)
prset popsizepr = fixed(<number>)

This parameter is only relevant if the coalescence process is selected as the prior on branch lengths. Note that the setting of 'ploidy' in 'lset' is important for how this parameter is interpreted.

Popvarpr

-- In a gene tree - species tree model, this parameter determines whether the population size is the same for the entire species tree ('popvarpr = equal', the default), or varies across branches of the species tree ('popvarpr=variable').

Nodeagepr

-- This parameter specifies the assumptions concerning the age of the terminal and interior nodes in the tree. The default model ('nodeagepr = unconstrained') assumes that all terminal nodes are of the same age while the age of interior nodes is unconstrained. The alternative ('nodeagepr = calibrated') option derives a prior probability distribution on terminal and interior node ages from the calibration settings (see the 'calibrate' command). The 'nodeagepr' parameter is only relevant for clock trees.

Clockratepr

-- This parameter specifies the prior assumptions concerning the base substitution rate of the tree, measured in expected number of substitutions per site per time unit. The default setting is 'Fixed(1.0)', which effectively means that the time unit is the number of expected substitutions per site. If you apply age constraints to the tree, the default setting changes automatically to 'Exponential(<x>)', where '<x>' is set such that the expectation of the exponential is ten times the age of the maximum age constraint. This will give you a very vague prior, which may or may not be adequate for your particular problem.

If you do not have any age calibrations in the tree, you can still calibrate the tree using 'Clockratepr'. For instance, if you know that your sequence data evolve at a rate of 0.20 substitutions per million years, you might calibrate the tree by fixing the substitution rate to 0.20 using

```
prset clockratepr = fixed(0.20)
```

after which the tree will be calibrated using millions of years as the unit.

You can also assign a prior probability distribution to the substitution rate, accommodating the uncertainty about its value. You can choose between normal, lognormal, exponential and gamma distributions for this purpose. For instance, if you would like to associate the substitution rate with a normal distribution truncated at 0, so that only positive values are allowed, and with mean 0.20 and standard deviation of 0.02, you would use

prset clockratepr = normal(0.20,0.02)

The lognormal distribution is parameterized in terms of the mean and standard deviation on the log scale (natural logs). For instance,

prset clockratepr = lognormal(-1.61,0.10)

specifies a lognormal distribution with a mean of log values of -1.61 and a standard deviation of log values of 0.10. In such a case, the mean value of the lognormal distribution is equal to  $e^{(-1.61 + 0.10^2/2)} = 0.20$ .

Note that the 'Clockratepr' parameter has no effect on non-clock trees.

Clockvarpr

-- This parameter allows you to specify the type of clock you are assuming. The default is 'strict', which corresponds to the standard clock model where the evolutionary rate is constant throughout the tree. You can also use 'cpp', which invokes a relaxed clock model where the rate evolves according to a Compound Poisson Process (CPP) model (see Huelsenbeck et al., 2000) or 'tk02', which invokes the Brownian Motion (TK02) model described by Thorne and Kishino (2002). Finally, you can use a model where each branch has an independent rate drawn from a scaled gamma distribution, such that there is a specified variance in the effective height of the tree in the prior, the Independent Gamma Rate (IGR) model (LePage et al., 2007). Each of the relaxed clock models has additional parameters with priors. For the CPP model, it is 'cppratepr' and 'cppmultdevpr'; for the TKO2 model, it is 'tk02varpr'; for the IGR model, it is 'igrvarpr'. The 'clockvarpr' parameter is only relevant for clock trees.

For backward compatibility, 'bm' is allowed as a synonym of 'tk02', and 'ibr' as a synonym of 'igr'.

Cppratepr

-- This parameter allows you to specify a prior probability distribution on the rate of the Poisson process generating changes in the evolutionary rate in the CPP relaxed clock model. You can either fix the rate or associate it with an exponential prior using

```
prset cppratepr = fixed(<number>)
prset cppratepr = exponential(<number>)
```

For instance, if you fix the rate to 2, then on a branch with the length equual to one expresed in terms of average expected number of substitution per site, you expect to see, on average, two rate-modifying events. If you put an exponential (0.1) on the rate, you will be estimating the rate against a prior probability distribution where the expected rate is  $10 \ (= 1/0.1)$ .

Cppmultdevpr -- This parameter allows you to specify the standard deviation of the log-normal distribution from which the rate multipliers of the CPP relaxed clock model are drawn. The standard deviation is given on the log scale. The default value of 1.0 thus corresponds to rate multipliers varying from 0.37 (1/e) to 2.7 (e) when they are +/- one standard deviation from the expected mean. The expected mean of the logarithm of the mulpliers is fixed to 0, ensuring that the expected mean rate is 1.0. You can change the default value by using

prset cppmultdevpr = fixed(<number>)

TK02varpr

where <number> is the standard deviation on the log scale. -- This parameter allows you to specify the prior probability distribution for the variance of the rate multiplier in the Thorne-Kishino ('Brownian motion') relaxed clock model. Specifically, the parameter specifies the rate at which the variance increases with respect to the base rate of the clock. If you have a branch of a length corresponding to 0.4 expected changes per site according to the base rate of the clock, and the tk02var parameter has a value of 2.0, then the rate multiplier at the end of the branch will be drawn from a lognormal distribution with a variance of 0.4\*2.0 (on the linear, not the logarithm scale). The mean is the same as the rate multiplier at the start of the branch (again on the linear scale).

You can set the parameter to a fixed value, or specify that it is drawn from an exponential or uniform distribution:

```
prset tk02varpr = fixed(<number>)
prset tk02varpr = exponential(<number>)
prset tk02varpr = uniform(<number>,<number>)
```

For backward compatibility, 'bmvarpr' is allowed as a synonym of 'tko2varpr'.

Igrvarpr

-- This parameter allows you to specify a prior on the variance of the gamma distribution from which the branch lengths are drawn in the independent branch rate (IGR) relaxed clock model. Specifically, the parameter specifies the rate at which the variance increases with respect to the base rate of the clock. If you have a branch of a length corresponding to 0.4 expected changes per site according to the base rate of the clock, and the igrvar parameter has a value of 2.0, then the effective branch length will be drawn from a distribution with a variance of 0.4\*2.0.

You can set the parameter to a fixed value, or specify that it is drawn from an exponential or uniform distribution:

```
prset igrvarpr = fixed(<number>)
prset igrvarpr = exponential(<number>)
prset igrvarpr = uniform(<number>,<number>)
```

For backward compatibility, 'ibrvarpr' is allowed as a synonym of 'igrvarpr'.

Ratepr

-- This parameter allows you to specify the site specific rates model or any other model that allows different partitions to evolve at different rates. First, you must have defined a partition of the characters. For example, you may define a partition that divides the characters by codon position, if

you have DNA data. You can also divide your data using a partition that separates different genes from each other. The next step is to make the desired partition the active one using the set command. For example, if your partition is called "by\_codon", then you make that the active partition using "set partition=by\_codon". Now that you have defined and activated a partition, you can specify the rate multipliers for the various partitions. The options are:

```
prset ratepr = fixed
prset ratepr = variable
prset ratepr = dirichlet(<number>,<number>,...,<number>)
```

If you specify "fixed", then the rate multiplier for that partition is set to 1 (i.e., the rate is fixed to the average rate across partitions). On the other hand, if you specify "variable", then the rate is allowed to vary across partitions subject to the constraint that the average rate of substitution across the partitions is 1. You must specify a variable rate prior for at least two partitions, otherwise the option is not activated when calculating likelihoods. The variable option automatically associates the partition rates with a dirichlet(1,...,1)prior. The dirichlet option is an alternative way of setting a partition rate to be variable, and also gives accurate control of the shape of the prior. The parameters of the Dirichlet are listed in the order of the partitions that the ratepr is applied to. For instance, "prset applyto=(1,3,4) ratepr = dirichlet(10,40,15)" would set the Dirichlet parameter 10 to partition 1, 40 to partition 3, and 15 to partition 4. The Dirichlet distribution is applied to the weighted rates; that is, it weights the partition rates according to the number of included characters in each partition.

## Default model settings:

Parameter	Options	Current Setting
Tratiopr	Beta/Fixed	Beta(1.0,1.0)
Revmatpr	Dirichlet/Fixed	Dirichlet(1.0,1.0,)
Aamodelpr	Fixed/Mixed	Fixed(Poisson)
Aarevmatpr	Dirichlet/Fixed	Dirichlet(1.0,1.0,)
Omegapr	Dirichlet/Fixed	<pre>Dirichlet(1.0,1.0)</pre>
Ny98omega1pr	Beta/Fixed	Beta(1.0,1.0)
Ny98omega3pr	Uniform/Exponential/Fixed	<pre>Exponential(1.0)</pre>
M3omegapr	Exponential/Fixed	Exponential
Codoncatfreqs	Dirichlet/Fixed	Dirichlet(1.0,1.0,1.0)
Statefreqpr	Dirichlet/Fixed	Dirichlet(1.0,1.0,1.0,1.0)
Shapepr	Uniform/Exponential/Fixed	Uniform(0.0,200.0)
Ratecorrpr	Uniform/Fixed	Uniform(-1.0,1.0)
Pinvarpr	Uniform/Fixed	Uniform(0.0,1.0)
Covswitchpr	Uniform/Exponential/Fixed	Uniform(0.0,100.0)
Symdirihyperpr	Uniform/Exponential/Fixed	Fixed(Infinity)
Topologypr	Uniform/Constraints/Fixed	Uniform
Brlenspr	Unconstrained/Clock/Fixed	<pre>Unconstrained:Exp(10.0)</pre>
Treeagepr	Exponential/Gamma/Fixed	<pre>Exponential(1.0)</pre>
Speciationpr	Uniform/Exponential/Fixed	<pre>Exponential(1.0)</pre>
Extinctionpr	Beta/Fixed	Beta(1.0,1.0)
SampleStrat	Random/Diversity/Cluster	Random
Sampleprob	<number></number>	1.00
Popsizepr	Lognormal/Gamma/Uniform/	Lognormal(4.6,2.3)

Normal/Fixed

Popvarpr Equal/Variable Equal

Nodeagepr Unconstrained/Calibrated Unconstrained

Clockratepr Fixed/Normal/Lognormal/

Exponential/Gamma Fixed(1.00)

Clockvarpr Strict/Cpp/TK02/Igr Strict

Cppratepr Fixed/Exponential Exponential(0.10)
Cppmultdevpr Fixed Fixed(0.40)

TK02varpr Fixed/Exponential/Uniform Exponential(10.00)
Igrvarpr Fixed/Exponential/Uniform Exponential(10.00)

Ratepr Fixed/Variable=Dirichlet Fixed

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### Propset

This command allows the user to change the details of the MCMC samplers (moves) that update the state of the chain. The useage is:

propset <move name>\$<tuning-parameter>=<value>

Assume we have a topology parameter called 'Tau{all}', which is sampled by the move 'ExtTBR(Tau{all})' (note that the parameter name is included in the move name). This move has three tuning parameters: (1) 'prob', the relative proposal probability (a weight defining its probability relative to other moves); (2) 'p\_ext', the extension probability; and (3) 'lambda', the tuning parameter of the branch length multiplier. A list of the tuning parameters is available by using 'Showmoves' (see below). To change the relative proposal probability to 20 and the extension probability to 0.7, use:

propset etbr(tau{all})\$prob=20 etbr(tau{all})\$p\_ext=0.7

This change would apply to all chains in all runs. It is also possible to set the tuning parameters of individual runs and chains using the format:

propset <move\_name>\$<tuning-parameter>(<run>,<chain>)=<value>

where <run> and <chain> are the index numbers of the run and chain for which you want to change the value. If you leave out the index of the run, the change will apply to all runs; if you leave out the index of the chain, the change will similarly apply to all chains. To switch off the exttbr(tau{all}) move in chain 2 of all runs, use:

propset etbr(tau{all})\$prob(,2)=0

It is important to note that all moves are not available until the model has been completely defined. Any change to the model will cause all proposal tuning parameters to return to their default values. To see a list of all the moves that are currently switched on for the model, use 'showmoves'. You can also see other available moves by using 'showmoves allavailable=yes'. A list of the moves for each parameter in the model is available by using the command 'Showparams'. If you change proposal probabilities, make sure that all parameters that are not fixed in your model have at least one move switched on.

One word of warning: You should be extremely careful when modifying any of the chain parameters using 'propset'. It is quite possible to completely wreck any hope of achieving convergence by inappropriately setting the tuning parameters. In general, you want to set move tuning parameters such that the acceptance rate of the move is intermediate (we suggest targeting the range 10% to 70% acceptance, if possible). If the acceptance rate is

outside of this range, the MCMC chain will probably not sample that parameter very efficiently. The acceptance rates for all moves in the cold chain(s) are summarized at the end of each run in the screen output. The acceptance rates (potentially for all chains, cold and heated) are also printed to the .mcmc file if Mcmc convergence diagnostics are turned on (using 'Mcmc' or 'Mcmcp').

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Ouit

This command quits the program. The correct usage is:

quit

It is a very easy command to use properly.

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## Report

This command allows you to control how the posterior distribution is reported. For rate parameters, it allows you to choose among several popular parameterizations. The report command also allows you to request printing of some model aspects that are usually not reported. For instance, if a node is constrained in the analysis, MrBayes can print the probabilities of the ancestral states at that node. Similarly, if there is rate variation in the model, MrBayes can print the inferred site rates, and if there is omega variation, MrBayes can print the inferred omega (positive selection) values for each codon. In a complex model with several partitions, each partition is controlled separately using the same 'Applyto' mechanism as in the 'Lset' and 'Prset' commands.

### Options:

Applyto

-- This option allows you to apply the report commands to specific partitions. This command should be the first in the list of commands specified in 'report'.

For example,

report applyto=(1,2) tratio=ratio

report applyto=(3) tratio=dirichlet

would result in the transition and transversion rates of the first and second partitions in the model being reported as a ratio and the transition and transversion rates of the third partition being reported as proportions of the rate sum (the Dirichlet parameterization).

Tratio

-- This specifies the report format for the transition and transversion rates of a nucleotide substituion model with nst=2. If 'ratio' is selected, the rates will be reported as a ratio (transition rate/transversion rate). If 'dirichlet' is selected, the transition and transversion rates will instead be reported as proportions of the rate sum. For example, if the transition rate is three times the transversion rate and 'ratio' is selected, this will reported as a single value, '3.0'. If 'dirichlet' is selected instead, the same rates will be reported using two values, '0.75 0.25'. The sum of the Dirichlet values is always 1. Although the Dirichlet format may be unfamiliar to some users, it is more convenient for specifying priors than the ratio format.

Revmat

-- This specifies the report format for the substitution rates of a GTR substitution model for nucleotide or amino acid data. If

'ratio' is selected, the rates will be reported scaled to the G-T rate (for nucleotides) or the Y-V rate (for amino acids). If 'dirichlet' is specified instead, the rates are reported as proportions of the rate sum. For instance, assume that the C-T rate is twice the A-G rate and four times the transversion rates, which are equal. If the report format is set to 'ratio', this would be reported as '1.0 2.0 1.0 1.0 4.0 1.0' since the rates are reported in the order rAC, rAG, rAT, rCG, rCT, rGT and scaled relative to the last rate, the G-T rate. If 'dirichlet' is selected instead, the same rates would have been reported as '0.1 0.2  $0.1\ 0.1\ 0.4\ 0.1'$  since the rates are now scaled so that they sum to 1.0. The Dirichlet format is the parameterization used for formulating priors on the rates.

Ratemult

-- This specifies the report format used for the rate multiplier of different model partitions. Three formats are available. If 'scaled' is selected, then rates are scaled such that the mean rate per site across partitions is 1.0. If 'ratio' is chosen, the rates are scaled relative to the rate of the first partition. Finally, if 'dirichlet' is chosen, the rates are given as proportions of the rate sum. The latter is the format used when formulating priors on the rate multiplier.

Tree

-- This specifies the report format used for the tree(s). Two options are available. 'Topology' results in only the topology being printed to file, whereas 'brlens' causes branch lengths to to be printed as well.

- Ancstates -- If this option is set to 'yes', MrBayes will print the probability of the ancestral states at all constrained nodes. Typically, you are interested in the ancestral states of only a few characters and only at one node in the tree. To perform such an analysis, first define and enforce a topology constraint using 'constraint' and 'prset topologypr = constraints (...)'. Then put the character(s) of interest in a separate partition and set MrBayes to report the ancestral states for that partition. For instance, if the characters of interest are in partition 2, use 'report applyto=(2) ancstates=yes' to force MrBayes to print the probability of the ancestral states of those characters at the constrained node to the '.p' file.
- Siterates -- If this option is set to 'yes' and the relevant model has rate variation across sites, then the site rates, weighted over rate categories, will be reported to the '.p' file.
- Possel -- If this option is set to 'yes' and the relevant model has omega variation across sites, the probability that each model site (codon in this case) is positively selected will be written to file.
- Siteomega -- If this option is set to 'yes' and the relevant model has omega variation across sites, the weighted omega value (over omega categories) for each model site will be reported to file.

# Default report settings:

Parameter	Options	Current Setting
Tratio Revmat Ratemult Tree Ancstates Siterates Possel Siteomega	Ratio/Dirichlet Ratio/Dirichlet Scaled/Ratio/Dirichlet Brlens/Topology Yes/No Yes/No Yes/No Yes/No Yes/No	Ratio Dirichlet Scaled Brlens No No No

#### Restore

This command restores taxa to the analysis. The correct usage is:

restore <name and/or number and/or taxset> ...

A list of the taxon names or taxon numbers (labelled 1 to ntax in the order in the matrix) or taxset(s) can be used. For example, the following:

restore 1 2 Homo\_sapiens

restores taxa 1, 2, and the taxon labelled Homo\_sapiens to the analysis. You can also use "all" to restore all of the taxa. For example,

restore all

restores all of the taxa to the analysis.

-----

This command is used to set some general features of the model or program behavior. The correct usage is

set <parameter>=<value> ... <parameter>=<value>

#### Available options:

Seed

-- Sets the seed number for the random number generator. The random number seed is initialized haphazardly at the beginning of each MrBayes session. This option allows you to set the seed to some specific value, thereby allowing you to exactly repeat an analysis. If the analysis uses swapping between cold and heated chains, you must also set the swap seed (see below) to exactly repeat the analysis.

Swapseed

-- Sets the seed used for generating the swapping sequence when Metropolis-coupled heated chains are used. This seed is initialized haphazardly at the beginning of each MrBayes session. This option allows you to set the seed to some specific value, thereby allowing you to exactly repeat a swap sequence. See also the 'Seed' option.

Dir

-- The working directory. Specifies the absolute or relative path to the working directory. If left empty, the working directory is the current directory.

Partition

-- Set this option to a valid partition id, either the number or name of a defined partition, to enforce a specific partitioning of the data. When a data matrix is read in, a partition called "Default" is automatically created. It divides the data into one part for each data type. If you only have one data type, DNA for instance, the default partition will not divide up the data at all. The default partition is always the first partition, so 'set partition=1' is the same as 'set partition=default'.

Speciespartition -- Set this option to a valid speciespartition id, either the number or name of a defined speciespartition, to enforce a specific partitioning of taxa to species. When a data matrix is read in, a speciespartition called "Default" is automatically created. It assigns one taxon for each species. The

default speciespartition is always the first speciespartition, so 'set speciespartition=1' is the same as

'set speciespartition=default'.

-- If autoclose is set to 'yes', then the program will not prompt Autoclose you during the course of executing a file. This is particularly useful when you run MrBayes in batch mode.

-- If nowarnings is set to yes, then the program will not prompt Nowarnings you when overwriting or appending an ouput file that is already present. If 'nowarnings=no' (the default setting), then the program propts the user before overwriting output files.

-- When nowarnings is set to yes, then MrBayes will by default Autoreplace overwrite output files that already exists. This may cause irrecoverable loss of previous results if you have not removed or renamed the files from previous runs. To override this behavior, set autooverwrite to no, in which case new output will be appended to existing files instead.

Quitonerror -- If quitonerror is set to yes, then the program will quit when an error is encountered, after printing an error message. If quitonerror is set to no (the default setting), then the program will wait for additional commands from the command line after the error message is printed.

Scientific -- Set this option to 'Yes' to write sampled values to file in scientific format and to 'No' to write them in fixed format. Fixed format is easier for humans to read but you risk losing precision for small numbers. For instance, sampled values that are less than 1E-6 will print to file as '0.000000' if fixed format is used and 'precision' is set to 6.

-- Precision allows you to set the number of decimals to be prin-Precision ted when sampled values are written to file. Precision must be in the range 3 to 15.

### Current settings:

Parameter	Options	Current Setting
Seed	<pre><number></number></pre>	1321443066
Swapseed	<number></number>	1321443066
Dir	<name></name>	11 11
Partition	<name></name>	11 11
Speciespartition	<name></name>	11 11
Autoclose	Yes/No	No
Nowarnings	Yes/No	No
Autoreplace	Yes/No	Yes
Quitonerror	Yes/No	No
Sientific	Yes/No	Yes
Precision	<number></number>	15

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#### Showbeagle

This command shows available BEAGLE resources.

# Showmatrix

This command shows the character matrix currently in memory.

#### Showmcmctrees

This command shows the current trees used by the Markov chains. is "showmcmctrees".

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Showmodel

This command shows the current model settings. The correct usage is

showmodel

After typing "showmodel", the modelling assumptions are shown on a partition-by-partition basis.

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Showmoves

This command shows the MCMC samplers (moves) that are switched on for the parameters in the current model. The basic usage is

showmoves

If you want to see all available moves, use

showmoves allavailable=yes

If you want to change any of the tuning parameters for the moves, use the 'propset' command.

-----

Showparams

This command shows all of the parameters in the current model. The basic usage is

showparams

The parameters are listed together with their priors, the available moves, and the current value(s), which will be used as the starting values in the next mcmc analysis.

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Showusertrees

This command shows the currently defined user trees. The correct usage is "shownsertrees"

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Speciespartition

Defines a partition of tips into species. The format for the speciespartition command is

Speciespartition <name> = <species name>:<taxon list> ,...,<sp nm>:<tx lst>

The command enumerates comma separated list of pairs consisting of 'species name' and 'taxon list'. The 'taxon list' is a standard taxon list, as used by the 'Taxset' command. This means that you can use either the index or the name of a sequence ('taxon'). Ranges are specified using a dash, and a period can be used as a synonym of the last sequence in the matrix.

For exammple: speciespartition species = SpeciesA: 1, SpeciesB: 2-. Here, we name two species. SpeciesA is represented by a single sequence while SpeciesB is represented by all remaining sequences in the matrix. Each sequence is specified by its row index in the data matrix.

As with ordinary partitioning you may define multiple species partitioning scheme. You have to use command 'set speciespartition' to enable use of one of them.

Currently defined Speciespartitions:

Number	Speciespartition	name	Number	of	species
Q q					

This command is used to start stepping-stone sampling, which is an efficient and accurate method for estimating the marginal likelihood of the currently specified model. It is considerably more accurate than the harmonic mean of the likelihoods from a standard MCMC run on the model (calculated by the 'Sump' command) but it requires a separate MCMC-like run. To be more specific, stepping-stone sampling uses importance sampling to estimate each ratio in a series of discrete steps bridging the posterior and prior distributions. The importance distributions that are used are called power posterior distributions, and are defined as prior\*(likelihood^beta). By varying beta from 1 to 0, we get a series of distributions that connect the posterior (beta = 1) to the prior (beta = 0).

The power posterior distributions are sampled using MCMC. First, we start a standard MCMC chain on the posterior distribution, and let it run until we have reached the criterion specified by the 'Burninss' option. After this, we step through the power posterior distributions until we reach the prior distribution. In each of the 'Nsteps' steps, we sample from a new power posterior distribution with a distinct beta value. The beta values correspond to 'Nsteps' evenly spaced quantiles in a Beta distribution with the parameters 'Alpha' and 1.0. For the first sampling step, the beta value is equal to the last quantile, i.e., it is close to 1.0. For each successive step, the beta value takes on the value of the next quantile, in decreasing order, until it reaches the value of 0.0.

The 'Ss' procedure uses the same machinery as the standard 'Mcmc' algorithm, and shares most of its parameters with the 'Mcmc' and 'Mcmcp' commands. All 'Mcmc' parameters, except those related to burnin, have the same meaning and usage in the 'Ss' command as they have in the 'Mcmc' command. However, the 'Mcmc' burnin parameters are ignored. Instead, the 'Ss' command uses its own burnin parameter, 'Burninss' (see below for details). The 'Ss' command also has its own parameters for specifying the number of steps and the shape of the Beta distribution from which the beta values are computed (see below).

Note that the 'Ngen' parameter of 'Mcmc' is used to set the maximum number of generations processed, including both the burnin and the following steps in the stepping-stone sampling phase. For instance, assume that 'Burninss' is set to '-1', 'Nsteps' to '49', 'Ngen' to '10000000' and 'Samplefreq' to '1000'. We will then get 1,000 samples in total (1,000,000 / 1,000). These will fall into 50 bins, one of which represents the burnin and is discarded. Each step in the algorithm will thus be represented by 20 samples.

More information on 'Mcmc' parameters is available in the help for the 'Mcmc'

and 'Mcmcp' commands. Only the exclusive 'Ss' parameters are listed below. These can only be set up using the 'Ss' command, while the parameters shared with 'Mcmc' and 'Mcmcp' can also be set up using those commands.

The correct usage is

ss <parameter>=<value> ... <parameter>=<value>

Note that a command:

ss <setting parameters shared with mcmc> <setting exclusive ss parameters>

would be equivalent to executing two commands:

mcmcp <setting parameters shared with mcmc>;
ss <setting exclusive ss parameters>;

For more information on the stepping-stone algorithm, see:

Xie, W., P. O. Lewis, Y. Fan, L. Kuo, and M.-H. Chen. 2011. Improving marginal likelihood estimation for Bayesian phylogenetic model selection. Systematic Biology 60:150-160.

### Available paremeters:

(NB: Only exclusive ss parameters listed here. For additional parameters, see help on 'mcmc' or 'mcmcp'.

Alpha

-- The beta values used in the stepping-stone sampling procedure correspond to evenly spaced quantiles from a Beta('Alpha',1.0) distribution. The parameter 'Alpha' determines the skewness of the beta values. If 'Alpha' is set to '1.0', the beta values would be spaced uniformly on the interval (0.0,1.0). However, better results are obtained if the beta values are skewed. Empirically, it was observed that 'Alpha' values in the range of 0.3 to 0.5 produce the most accurate results.

Burninss

-- Fixed number of samples discarded before sampling of the first step starts. 'Burninss' can be specified using either a positive or a negative number. If the number is positive, it is interpreted as the number of samples to discard as burnin. If the number is negative, its absolute value is interpreted as the length of the burnin in terms of the length of each of the following steps in the stepping-stone algorithm. For instance, a value of '-1' means that the length of the burnin is the same as the length of each of the subsequent steps.

Nsteps

Startvals

-- Number of steps in the stepping-stone algorithm. Typically, a number above 30 is sufficient for accurate results.

### Current settings:

Parameter	Options	Current Setting
Alpha BurninSS Nsteps	<number> <number> <number></number></number></number>	0.40 -1 50

Use this command to change the current values for parameters in your model. These values will be used as the starting values in the next mcmc analysis.

The basic format is:

```
startvals <param>=(<value_1>,<value_2>,...,<value_n>)
```

for all substitution model parameters. The format is slightly different for parameters that are written to a tree file:

startvals <param>=<tree\_name>

This version of the command will look for a tree with the specified name among the trees read in previously when parsing a tree block. The information stored in that tree will be used to set the starting value of the parameter. The parameters that are set using this mechanism include topology and branch length parameters, as well as relaxed clock branch rates, cpp events and cpp branch rate multipliers.

The above versions of the command will set the value for all runs and chains. You can also set the value for an individual run and chain by using the format

```
startvals <param>(<run>, <chain>)=(<value 1>,...)
```

where <run> is the index of the run and <chain> the index of the chain. If the run index is omitted, the values will be changed for all runs. Similarly, if the chain index is omitted, all chains will be set to the specified value. For example, if we wanted to set the values of the stationary frequency parameter  $pi\{1\}$  to (0.1,0.1,0.4,0.4) for all chains in run 1, and to (0.3,0.3,0.2,0.2) for chain 3 of run 2, we would use

startvals pi{1}(1,)=(0.1,0.1,0.4,0.4) pi{1}(2,3)=(0.3,0.3,0.2,0.2)

During an MCMC analysis, MrBayes prints the sampled parameter values to one or more tab-delimited text files, one for each independent run in your analysis. The command 'Sump' summarizes the information in this parameter file or these parameter files. By default, the root of the parameter file name(s) is assumed to be the name of the last matrix-containing nexus file. MrBayes also remembers the number of independent runs in the last analysis that you set up, regardless of whether you actually ran it. For instance, if there were two independent runs, which is the initial setting when you read in a new matrix, MrBayes will assume that there are two parameter files with the endings '.run1.p' and '.run2.p'. You can change the root of the file names and the number of runs using the 'Filename' and 'Nruns' settings.

When you invoke the 'Sump' command, three items are output: (1) a generation plot of the likelihood values; (2) estimates of the marginal likelihood of the model; and (3) a table with the mean, variance, and 95 percent credible interval for the sampled parameters. All three items are output to screen. The table of marginal likelihoods is also printed to a file with the ending '.lstat' and the parameter table to a file with the ending '.pstat'. For some model parameters, there may also be a '.mstat' file.

When running 'Sump' you typically want to discard a specified number or fraction of samples from the beginning of the chain as the burn in. This is done using the same mechanism used by the 'mcmc' command. That is, if you run an mcmc analysis with a relative burn in of 25 % of samples for convergence diagnostics, then the same burn in will be used for a subsequent sump command, unless a different burn in is specified. That is, issuing

### sump

immediately after 'mcmc', will result in using the same burn in settings as for the 'mcmc' command. All burnin settings are reset to default values every time a new matrix is read in, namely relative burnin ('relburnin=yes') with 25 % of samples discarded ('burninfrac = 0.25').

## Options:

Relburnin	If this option is set to 'Yes', then a proportion of the samples will be discarded as burnin when calculating summary statistics. The proportion to be discarded is set with 'Burninfrac' (see below). When the 'Relburnin' option is set to 'No', then a specific number of samples is discarded instead. This number is set by 'Burnin' (see below). Note that the burnin setting is shared across the 'sumt', 'sump', and 'mcmc' commands.
Burnin	Determines the number of samples (not generations) that will be discarded when summary statistics are calculated. The value of this option is only applicable when 'Relburnin' is set to 'No'.
Burninfrac	Determines the fraction of samples that will be discarded when summary statistics are calculated. The setting only takes effect if 'Relburnin' is set to 'Yes'.
Nruns	Determines how many '.p' files from independent analyses that will be summarized. If Nruns > 1 then the names of the files are derived from 'Filename' by adding '.run1.p', '.run2.p', etc. If Nruns=1, then the single file name is obtained by adding '.p' to 'Filename'.
Filename	The name of the file to be summarized. This is the base of the file name to which endings are added according to the current setting of the 'Nruns' parameter. If 'Nruns' is 1, then only '.p' is added to the file name. Otherwise, the endings will be '.run1.p', '.run2.p', etc.
Outputname	Base name of the file(s) to which 'Sump' results will be printed.
Hpd	Determines whether credibility intervals will be given as the region of Highest Posterior Density ('Yes') or as the interval containing the median 95 % of sampled values ('No').
Minprob	Determines the minimum probability of submodels to be included in summary statistics. Only applicable to models that explore submodel spaces, like 'nst=mixed' and 'aamodelpr=mixed'.

# Current settings:

Parameter	Options	Current Setting
Relburnin Burnin Burninfrac Nruns Filename Outputname Hpd Minprob	Yes/No <number> <number> <number> <name> <name> <name> <number> <name> <number></number></name></number></name></name></name></number></number></number>	Yes 0 0.25 2 temp.p<.run <i>.p&gt; &lt;.pstat etc&gt; 5391905 0.050</i>
Sumt		

This command is used to produce summary statistics for trees sampled during

a Bayesian MCMC analysis. You can either summarize trees from one individual analysis, or trees coming from several independent analyses. In either case, all the sampled trees are read in and the proportion of the time any single taxon bipartition (split) is found is counted. The proportion of the time that the bipartition is found is an approximation of the posterior probability of the bipartition. (Remember that a taxon bipartition is defined by removing a branch on the tree, dividing the tree into those taxa to the left and right of the removed branch. This set is called a taxon bipartition.) The branch length of the bipartition is also recorded, if branch lengths have been saved to file. The result is a list of the taxon bipartitions found, the frequency with which they were found, the posterior probability of the bipartition and, the mean and variance of the branch lengths or node depths, and various other statistics.

The key to the partitions is output to a file with the suffix '.parts'. The summary statistics pertaining to bipartition probabilities are output to a file with the suffix '.tstat', and the statistics pertaining to branch or node parameters are output to a file with the suffix '.vstat'.

A consensus tree is also printed to a file with the suffix '.con.tre' and printed to the screen as a cladogram, and as a phylogram if branch lengths have been saved. The consensus tree is either a 50 percent majority rule tree or a majority rule tree showing all compatible partitions. If branch lengths have been recorded during the run, the '.con.tre' file will contain a consensus tree with branch lengths and interior nodes labelled with support values. By default, the consensus tree will also contain other summary information in a format understood by the program 'FigTree'. To use a simpler format understood by other tree-drawing programs, such as 'TreeView', set 'Conformat' to 'Simple'.

MrBayes alo produces a file with the ending ".trprobs" that contains a list of all the trees that were found during the MCMC analysis, sorted by their probabilities. This list of trees can be used to construct a credible set of trees. For example, if you want to construct a 95 percent credible set of trees, you include all of those trees whose cumulative probability is less than or equal to 0.95. You have the option of displaying the trees to the screen using the "Showtreeprobs" option. The default is to not display the trees to the screen; the number of different trees sampled by the chain can be quite large. If you are analyzing a large set of taxa, you may actually want to skip the calculation of tree probabilities entirely by setting 'Calctreeprobs' to 'No'.

When calculating summary statistics you probably want to skip those trees that were sampled in the initial part of the run, the so-called burn-in period. The number of skipped samples is controlled by the 'Relburnin', 'Burnin', and 'Burninfrac' settings, just as for the 'Mcmc' command. Since version 3.2.0, the burn-in settings are shared across the 'Sumt', 'Sump' and 'Mcmc' commands. That is, changing the burn-in setting for one command will change the settings for subsequent calls to any of the other commands.

If you are summarizing the trees sampled in several independent analyses, such as those resulting from setting the 'Nruns' option of the 'Mcmc' command to a value larger than 1, MrBayes will also calculate convergence diagnostics for the sampled topologies and branch lengths. These values can help you determine whether it is likely that your chains have converged.

The 'Sumt' command expands the 'Filename' according to the current values of the 'Nruns' and 'Ntrees' options. For instance, if both 'Nruns' and 'Ntrees' are set to 1, 'Sumt' will try to open a file named '<Filename>.t'. If 'Nruns' is set to 2 and 'Ntrees' to 1, then 'Sumt' will open two files, the first named '<Filename>.run1.t' and the second '<Filename>.run2.t', etc. By default,

the 'Filename' option is set such that 'Sumt' automatically summarizes all the results from your immediately preceding 'Mcmc' command. You can also use the 'Sumt' command to summarize tree samples in older analyses. If you want to do that, remember to first read in a matrix so that MrBayes knows what taxon names to expect in the trees. Then set the 'Nruns', 'Ntrees' and 'Filename' options appropriately if they differ from the MrBayes defaults.

## Options:

Relburnin	 If this option is set to YES, then a proportion of the samples will be discarded as burnin when calculating summary statistics. The proportion to be discarded is set with Burninfrac (see below). When the Relburnin option is set to NO, then a specific number of samples is discarded instead. This number is set by Burnin (see below). Note that the burnin setting is shared across the 'sumt', 'sump', and 'mcmc' commands.
Burnin	 Determines the number of samples (not generations) that will be discarded when summary statistics are calculated. The value of this option is only relevant when Relburnin is set to NO.
BurninFrac	 Determines the fraction of samples that will be discarded when summary statistics are calculated. The value of this option is only relevant when Relburnin is set to YES. Example: A value for this option of 0.25 means that 25% of the samples will be discarded.
Nruns	 Determines how many '.t' files from independent analyses that will be summarized. If Nruns > 1 then the names of the files are derived from 'Filename' by adding '.run1.t', '.run2.t', etc. If Nruns=1 and Ntrees=1 (see below), then only '.t' is added to 'Filename'.
Ntrees	 Determines how many trees there are in the sampled model. If 'Ntrees' > 1 then the names of the files are derived from 'Filename' by adding '.tree1.t', '.tree2.t', etc. If there are both multiple trees and multiple runs, the filenames will be ' <filename>.tree1.run1.t', '<filename>.tree1.run2.t', etc.</filename></filename>
Filename	 The name of the file(s) to be summarized. This is the base of the file name, to which endings are added according to the current settings of the 'Nruns' and 'Ntrees' options.
Minpartfreq	 The minimum probability of partitions to include in summary statistics.
Contype	 Type of consensus tree. 'Halfcompat' results in a 50% majority rule tree, 'Allcompat' adds all compatible groups to such a tree.
Conformat	 Format of consensus tree. The 'Figtree' setting results in a consensus tree formatted for the program FigTree, with rich summary statistics. The 'Simple' setting results in a simple consensus tree written in a format read by a variety of programs.
Outputname	 Base name of the file(s) to which 'sumt' results will be printed. The default is the same as 'Filename'.
	Determines whether tree probabilities should be calculated. Determines whether tree probabilities should be displayed on
Hpd	 Determines whether credibility intervals will be given as the

## Current settings:

Parameter Options Current Setting

region of Highest Posterior Density ('Yes') or as the interval containing the median 95 % of sampled values ('No').

Relburnin Yes/No Burnin <number> Yes Burninfrac <number> 0.25 Nruns <number> Ntrees <number>

Filename <name> temp.t<.run<i>.t>

Minpartfreq <number>
Contype Halfcompat/Allcompat 0.10 Halfcompat Figtree

Outputname <name> temp.t.stat<.parts etc>

Calctreeprobs Yes/No Yes Showtreeprobs Yes/No No Hpd Yes/No Yes

#### Taxastat

This command shows the status of all the taxa. The correct usage is

#### taxastat

After typing "taxastat", the taxon number, name, and whether it is excluded or included are shown.

#### Taxset

This command defines a taxon set. The format for the taxset command

taxset <name> = <taxon names or numbers>

For example, "taxset apes = Homo Pan Gorilla Orang gibbon" defines a taxon set called "apes" that includes five taxa (namely, apes). You can assign up to 30 taxon sets. This option is best used not from the command line but rather as a line in the mrbayes block of a file.

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### Unlink

This command unlinks model parameters across partitions of the data. The correct usage is:

unlink <parameter name> = (<all> or <partition list>)

A little background is necessary to understand this command. Upon execution of a file, a default partition is set up. This partition referenced either by its name ("default") or number (0). If your data are all of one type, then this default partition does not actually divide up your characters. However, if your datatype is mixed, then the default partition contains as many divisions as there are datatypes in your character matrix. Of course, you can also define other partitions, and switch among them using the set command ("set partition=<name/number>"). Importantly, you can also assign model parameters to individual partitions or to groups of them using the "applyto" option in 1set and prset. When the program attempts to perform an analysis, the model is set for individual partitions. If the same parameter applies to differpartitions and if that parameter has the same prior, then the program

will link the parameters: that is, it will use a single value for the parameter. The program's default, then, is to strive for parsimony. However, there are lots of cases where you may want unlink a parameter across partitions. For example, you may want a different transition/ transversion rate ratio to apply to different partitions. This command allows you to unlink the parameters, or to make them different across partitions. The converse of this command is "link", which links together parameters that were previously told to be different. The list of parameters that can be unlinked includes:

```
-- Transition/transversion rate ratio
  Tratio
               -- Substitution rates of GTR model
  Revmat
  Omega
               -- Nonsynonymous/synonymous rate ratio
  Statefreq
               -- Character state frequencies
               -- Gamma shape parameter
  Pinvar
               -- Proportion of invariable sites
  Correlation
               -- Correlation parameter of autodiscrete gamma
  Switchrates
               -- Switching rates for covarion model
  Brlens
               -- Branch lengths of tree
  Topology
               -- Topology of tree
  Speciationrates -- Speciation rates for birth-death process
  Ratemultiplier -- Rate multiplier for partitions
  Extinctionrates -- Extinction rates for birth-death process
         -- Parameter for coalescence process
  Growthrate
               -- Growth rate of coalescence process
  Aamodel
               -- Aminoacid rate matrix
  Cpprate
               -- Rate of Compound Poisson Process (CPP)
  Cppmultdev
               -- Standard dev. of CPP rate multipliers (log scale)
  Cppevents
               -- CPP events
  TK02var
               -- Variance increase in TK02 relaxed clock model
  TK02branchrates -- Branch rates of TK02 relaxed clock model
          -- Variance increase in IGR relaxed clock model
  Igrbranchlens -- Branch lengths of IGR relaxed clock model
For example,
  unlink shape=(all)
unlinks the gamma shape parameter across all partitions of the data.
You can use "showmodel" to see the current linking status of the
______
This command shows the release version of the program.
______
*******************
  3. 'Data' or 'tree' block commands (in #NEXUS file)
*******************
Begin
```

This command is used to format data or commands in the program. The correct usage is

begin <data or mrbayes>;

The two valid uses of the "begin" command, then, are

```
begin data;
begin mrbayes;
```

The "data" specifier is used to specify the beginning of a data block; your character data should follow. For example, the following is an example of a data block for four taxa and ten DNA sites:

```
begin data;
  dimensions ntax=4 nchar=10;
  format datatype=dna;
  matrix
  taxon_1 AACGATTCGT
  taxon_2 AAGGATTCCA
  taxon_3 AACGACTCCT
  taxon_4 AAGGATTCCT;
end;
```

The other commands -- dimensions, format, and matrix -- are discussed in the appropriate help menu. The only thing to note here is that the block begins with a "begin data" command. The "mrbayes" command is used to enter commands specific to the MrBayes program into the file. This allows you to automatically process commands on execution of the program. The following is a simple mrbayes block:

```
begin mrbayes;
  charset first = 1-10\3;
  charset second = 2-10\3;
  charset third = 3-10\3;
end;
```

This mrbayes block sets off the three "charset" commands, used to predefine some blocks of characters. The mrbayes block can be very useful. For example, in this case, it would save you the time of typing the character sets each time you executed the file. Also, note that every "begin <data or mrbayes>" command ends with an "end". Finally, you can have so-called foreign blocks in the file. An example of a foreign block would be "begin paup". The program will simply skip this block. This is useful because it means that you can use the same file for MrBayes, PAUP\* or MacClade (although it isn't clear why you would want to use those other programs).

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### Dimensions

This command is used in a data block to define the number of taxa and characters. The correct usage is

dimensions ntax=<number> nchar=<number>

The dimensions must be the first command in a data block. The following provides an example of the proper use of this command:

```
begin data;
  dimensions ntax=4 nchar=10;
  format datatype=dna;
  matrix
  taxon_1 AACGATTCGT
```

```
taxon_2 AAGGATTCCA
     taxon_3 AACGACTCCT
     taxon_4 AAGGATTCCT
  end;
Here, the dimensions command tells MrBayes to expect a matrix with four
taxa and 10 characters.
______
______
End
This command is used to terminate a data or mrbayes block. The correct
usage is
  end;
For more information on this, check the help for the "begin" command.
______
______
Endblock
This is an older, deprecated version of "End", see that command.
Format
This command is used in a data block to define the format of the char-
acter matrix. The correct usage is
  format datatype=<name> ... <parameter>=<option>
The format command must be the second command in a data block. The following
provides an example of the proper use of this command:
  begin data;
     dimensions ntax=4 nchar=10;
     format datatype=dna gap=-;
    matrix
    taxon_1 AACGATTCGT
     taxon_2 AAGGAT--CA
     taxon_3 AACGACTCCT
     taxon_4 AAGGATTCCT
  end;
Here, the format command tells MrBayes to expect a matrix with DNA char-
acters and with gaps coded as "-".
The following are valid options for format:
        -- This parameter MUST BE INCLUDED in the format command. More-
Datatype
           over, it must be the first parameter in the line. The
           datatype command specifies what type of characters are
            in the matrix. The following are valid options:
              Datatype = Dna: DNA states (A,C,G,T,R,Y,M,K,S,W,H,B,
                       V,D,N)
              Datatype = Rna: DNA states (A,C,G,U,R,Y,M,K,S,W,H,B,
                       V,D,N)
              Datatype = Protein: Amino acid states (A,R,N,D,C,Q,E,
```

G,H,I,L,K,M,F,P,S,T,W,Y,V)

```
Datatype = Mixed(<type>:<range>,...,<type>:<range>): A
                            mixture of the above datatypes. For example,
                             "datatype=mixed(dna:1-100,protein:101-200)"
                            would specify a mixture of DNA and amino acid
                            characters with the DNA characters occupying
                            the first 100 sites and the amino acid char-
                            acters occupying the last 100 sites.
Interleave -- This parameter specifies whether the data matrix is in
              interleave format. The valid options are "Yes" or "No",
              with "No" as the default. An interleaved matrix looks like
                 format datatype=dna gap=- interleave=yes;
                 matrix
                 taxon 1 AACGATTCGT
                 taxon_2 AAGGAT--CA
taxon_3 AACGACTCCT
                 taxon 4 AAGGATTCCT
                 taxon_1 CCTGGTAC taxon_2 CCTGGTAC
                 taxon_3
                          ---GGTAG
                 taxon 4 ---GGTAG
           -- This parameter specifies the format for gaps. Note that
Gap
              gap character can only be a single character and that it
              cannot correspond to a standard state (e.g., A,C,G,T,R,Y,
              M,K,S,W,H,B,V,D,N for nucleotide data).
           -- This parameter specifies the format for missing data. Note
Missing
              that the missing character can only be a single character and
              cannot correspond to a standard state (e.g., A,C,G,T,R,Y,
              M,K,S,W,H,B,V,D,N for nucleotide data). This is often an
              unnecessary parameter to set because many data types, such
              as nucleotide or amino acid, already have a missing char-
              acter specified. However, for morphological or restriction
              site data, "missing=?" is often used to specify ambiguity
              or unobserved data.
Matchchar -- This parameter specifies the matching character for the
              matrix. For example,
                 format datatype=dna gap=- matchchar=.;
                 matrix
                 taxon 1 AACGATTCGT
                 taxon 2 ..G...--CA
                 taxon_3 .....C...C.
                 taxon_4 ..G....C.
              is equivalent to
                 format datatype=dna gap=-;
                 matrix
                 taxon 1 AACGATTCGT
                 taxon_2 AAGGAT--CA
                 taxon_3 AACGACTCCT
```

Datatype = Restriction: Restriction site (0,1) states

Datatype = Standard: Morphological (0,1) states
Datatype = Continuous: Real number valued states

```
taxon_4 AAGGATTCCT
;
```

The only non-standard NEXUS format option is the use of the "mixed", "restriction", "standard" and "continuous" datatypes. Hence, if you use any of these datatype specifiers, a program like PAUP\* or MacClade will report an error (as they should because MrBayes is not strictly NEXUS compliant).

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Matrix

This command specifies the actual data for the phylogenetic analysis. The character matrix should follow the dimensions and format commands in a data block. The matrix can have all of the characters for a taxon on a single line:

```
begin data;
      dimensions ntax=4 nchar=10;
      format datatype=dna gap=-;
      matrix
      taxon 1 AACGATTCGT
      taxon_2 AAGGAT--CA
taxon_3 AACGACTCCT
taxon_4 AAGGATTCCT
   end;
or be in "interleaved" format:
   begin data;
      dimensions ntax=4 nchar=20;
      format datatype=dna gap=- interleave=yes;
      matrix
      taxon_1 AACGATTCGT
      taxon_2 AAGGAT--CA
      taxon_3 AACGACTCCT
      taxon_4 AAGGATTCCT
      taxon_1 TTTTCGAAGC
      taxon_2 TTTTCGGAGC
      taxon_3 TTTTTGATGC
      taxon_4 TTTTCGGAGC
   end;
```

Note that the taxon names must not have spaces. If you really want to indicate a space in a taxon name (perhaps between a genus and species name), then you might use an underline ("\_"). There should be at least a single space after the taxon name, separating the name from the actual data on that line. There can be spaces between the characters.

If you have mixed data, then you specify all of the data in the same matrix. Here is an example that includes two different data types:

```
begin data;
  dimensions ntax=4 nchar=20;
  format datatype=mixed(dna:1-10,standard:21-30) interleave=yes;
  matrix
  taxon_1 AACGATTCGT
```

```
taxon_2 AAGGAT--CA
taxon_3 AACGACTCCT
taxon_4 AAGGATTCCT

taxon_1 0001111111
taxon_2 0111110000
taxon_3 1110000000
taxon_4 1000001111
;
end;
```

The matrix command is terminated by a semicolon.

Finally, just a note on data presentation. It is much easier for others to (1) understand your data and (2) repeat your analyses if you make your data clean, comment it liberally (using the square brackets), and embed the commands you used in a publication in the mrbayes block. Remember that the data took a long time for you to collect. You might as well spend a little time making the data file look nice and clear to any that may later request the data for further analysis.

-----

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#### Taxlabels

This command defines taxon labels. It could be used within taxa block.

-----

#### Translate

This command is used by MrBayes to specify the mapping between taxon names and taxon numbers in a Nexus tree file. For instance,

```
translate
```

- 1 Homo,
- 2 Pan,
- 3 Gorilla,
- 4 Hylobates;

establishes that the taxon labeled 1 in the trees that follow is Homo, the taxon labeled 2 is Pan, etc.

\_\_\_\_\_\_

### Tree

This command is used by MrBayes to write trees to a nexus tree file. Trees are written in the Newick format. For instance,

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

describes an unrooted tree with taxa 1 and 2 being more closely related to each other than to taxa 3 and 4. If branch lengths are saved to file, they are given after a colon sign immediately following the terminal taxon or the interior node they refer to. An example of an unrooted tree with branch lengths is:

```
tree ((1:0.064573,2:0.029042):0.041239,3:0.203988,4:0.187654);
```

Trees that are rooted (clock trees) are written with a basal dichotomy instead of a basal trichotomy. If the tree described above had been rooted

on the branch leading to taxon 4, it would have been represented as: tree (((1,2),3),4);