Lecture 2.2

Bayesian Phylogenetics I

The Bayesian framework

Bayesian phylogenetic analysis

- Bayesian phylogenetic analysis was developed in the mid 1990s
- Now one of the most widely used methods
- Bayes's theorem (1763)
- Reverend Thomas Bayes



Image probably not of Thomas Bayes

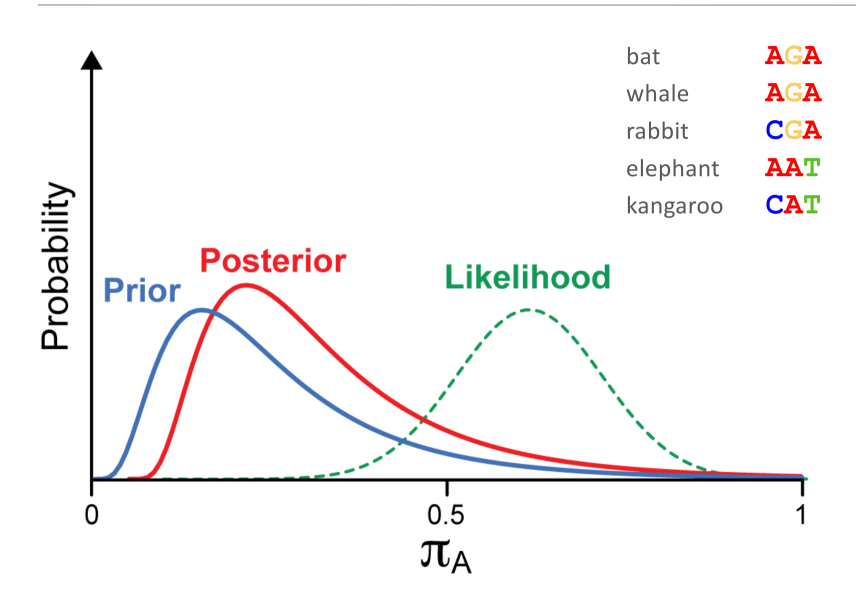
Contrast with frequentist statistics (likelihood)

Bayesian phylogenetic analysis

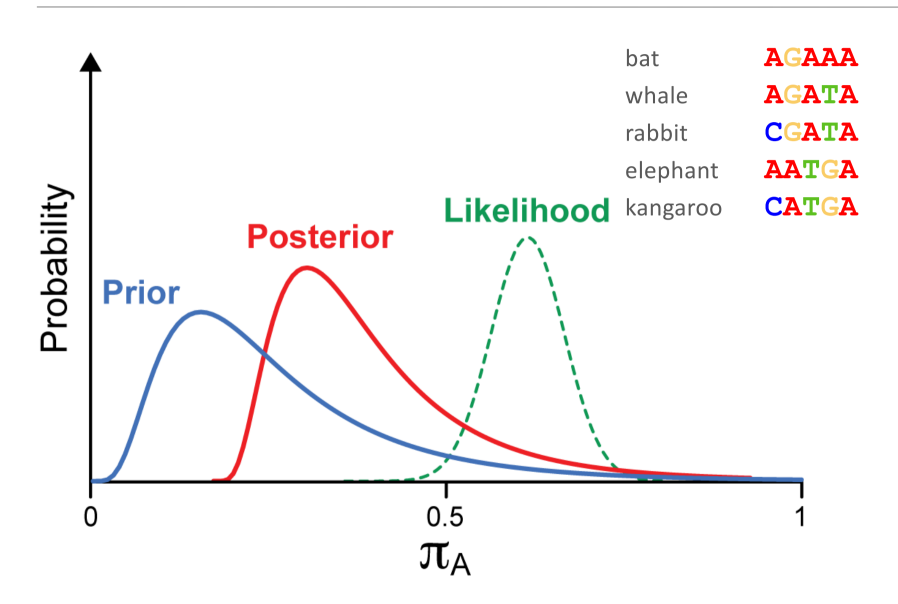
$$Pr(\theta \mid D) \propto Pr(\theta) Pr(D \mid \theta)$$

- Parameters have distributions
- Before the data are observed, each parameter has a prior probability distribution
 - Reflect our prior expectations (and uncertainty) about values of parameters (without knowledge of the data)
- Likelihood of the data is computed
- Prior probability distribution is combined (updated) with the likelihood to yield the posterior probability distribution

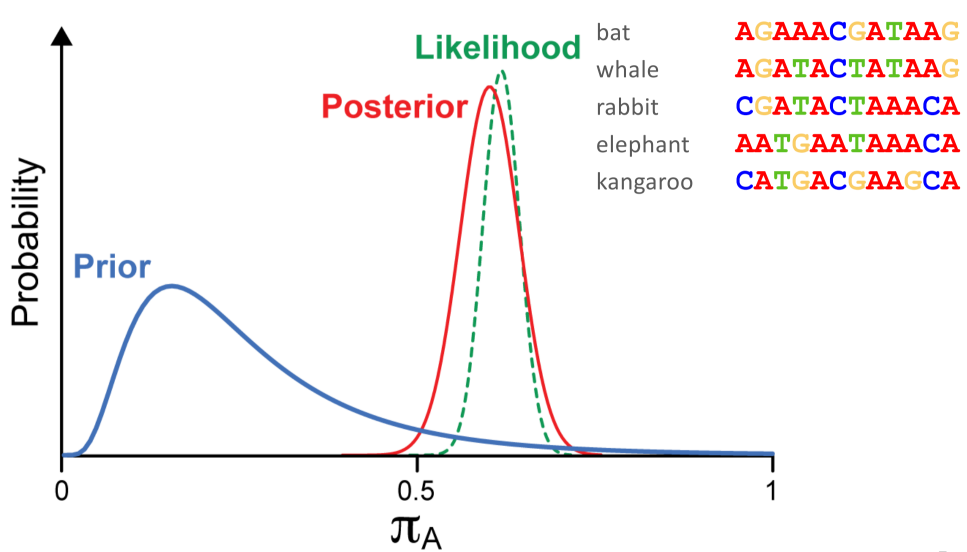
Simple example



Simple example



Simple example



Bayesian inference

Prior

Specified by user, independent of data

Likelihood

Calculated from data

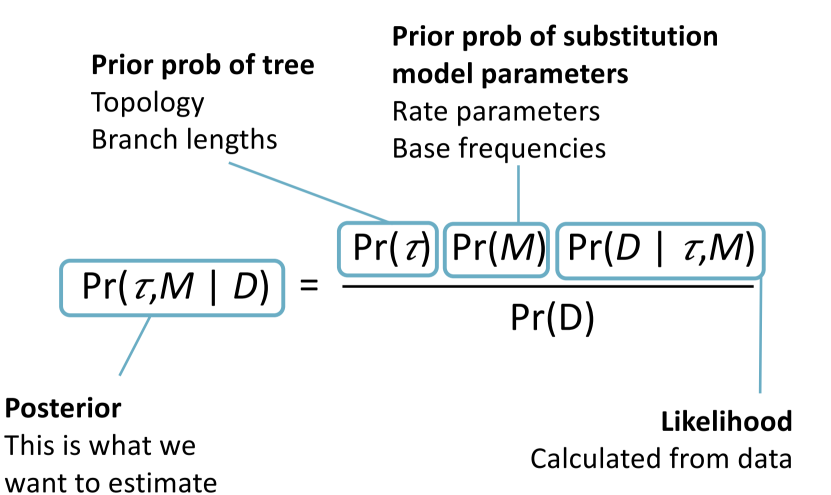
$$Pr(\theta \mid D) = \frac{Pr(\theta) Pr(D \mid \theta)}{Pr(D)}$$

Posterior

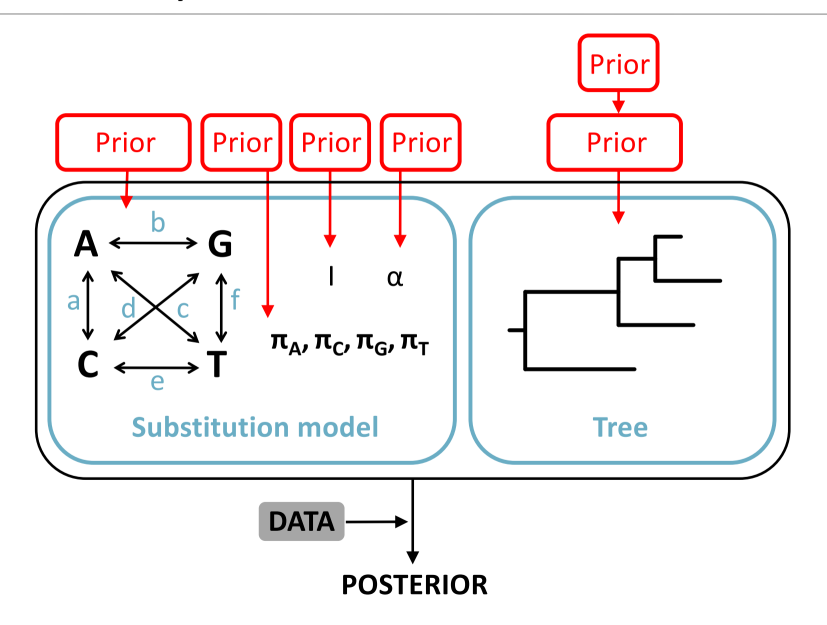
This is what we want to estimate

normalising constant marginal likelihood of the data model likelihood

Bayesian inference



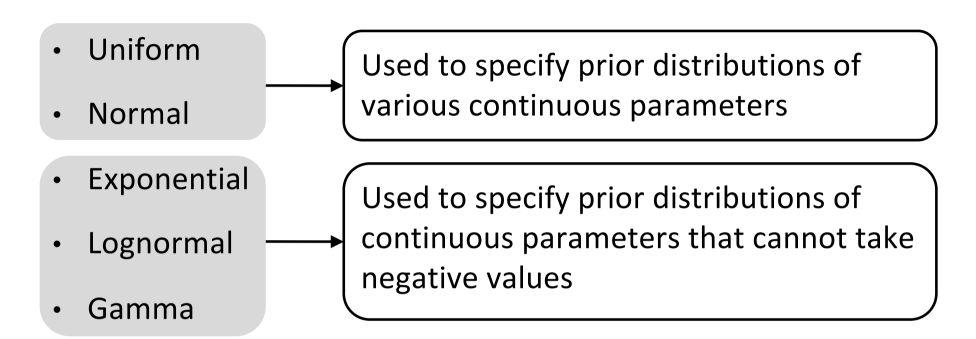
Bayesian hierarchical model





Priors

- Priors are chosen in the form of probability distributions
- Reflect our prior expectations (and uncertainty) about values of parameters (without knowledge of the data)
 - Past observations
 - Personal beliefs
 - Use of a biological model

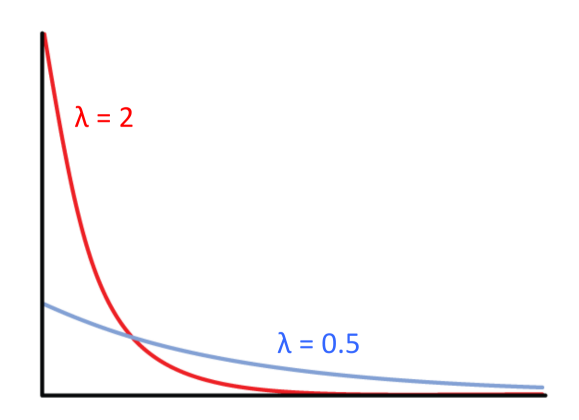


- Beta
- Dirichlet

- Uniform
- Normal
- Exponential
- Lognormal
- Gamma
- Beta
- Dirichlet

Parameters

• λ = rate of decay



- Uniform
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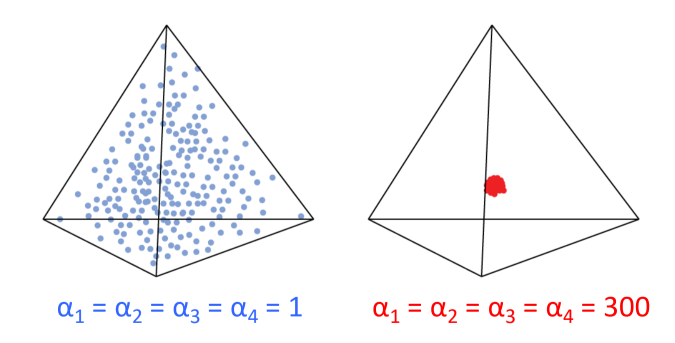
Used to specify the prior distributions of simplex parameters

- Base frequencies
- Relative rates in GTR model

- Uniform
- Normal
- Exponential
- Lognormal
- Gamma
- Beta
- Dirichlet

Parameters

• α_1 , α_2 , ... = shape parameters



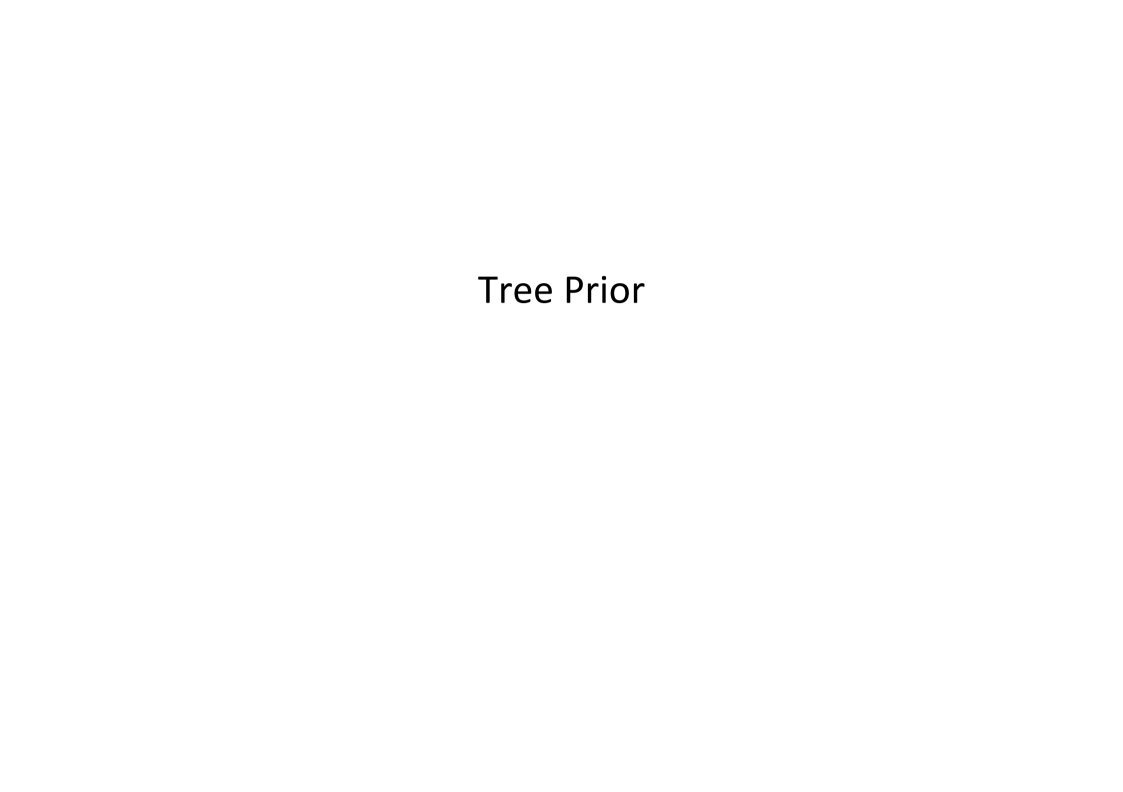
Discrete distributions

- Bernoulli distribution
- Binomial
- Multinomial
- Poisson

Default priors

	BEAST2	MrBayes
Rate matrix parameters	Gamma(0.05,10)	Dirichlet(1,1,1,1,1,1)
Base frequencies	Uniform(0,1)	Dirichlet(1,1,1,1)
Shape parameter (α)	Exponential(1)	Exponential(2)
Proportion invariable	Uniform(0,1)	Uniform(0,1)

Can specify uninformative priors where appropriate



Tree prior

- 1. Use a **flat prior** (*MrBayes*)
 - All trees have equal probability
 - Also need a prior for branch lengths or node times

- 2. Use a **biological model** (*BEAST* and *MrBayes*)
 - Among species: speciation model
 - Within species: coalescent model

Priors on rooted trees

Speciation model

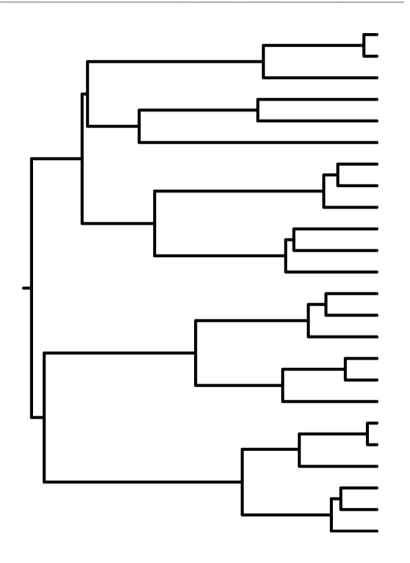
 Tree shape described by a stochastic branching process

Yule process

- The root lineage splits into two
- Lineages split at a constant rate
- Simulates speciation process

Birth-death process

Allow lineages to go extinct



Speciation model

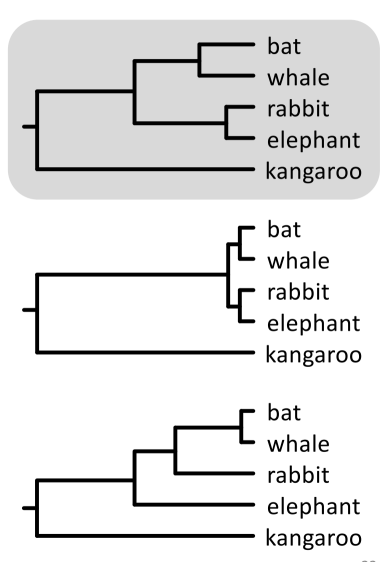
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Birth-death process

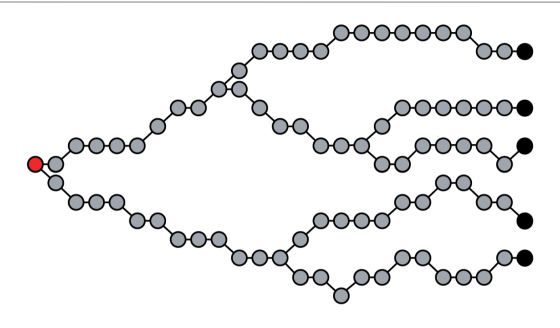
Allow lineages to go extinct

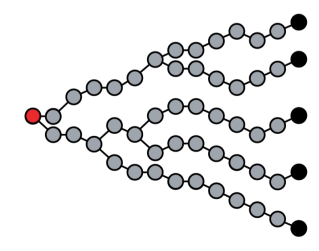


Coalescent model

Constant size

Exponential growth





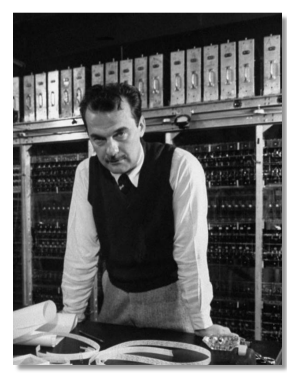
Choosing a tree prior

- Test whether inferences are robust to the choice of tree prior
- Mixed data sets: multiple sequences from each species
 - Birth-death prior generally works well
- Compare tree priors using Bayesian model selection

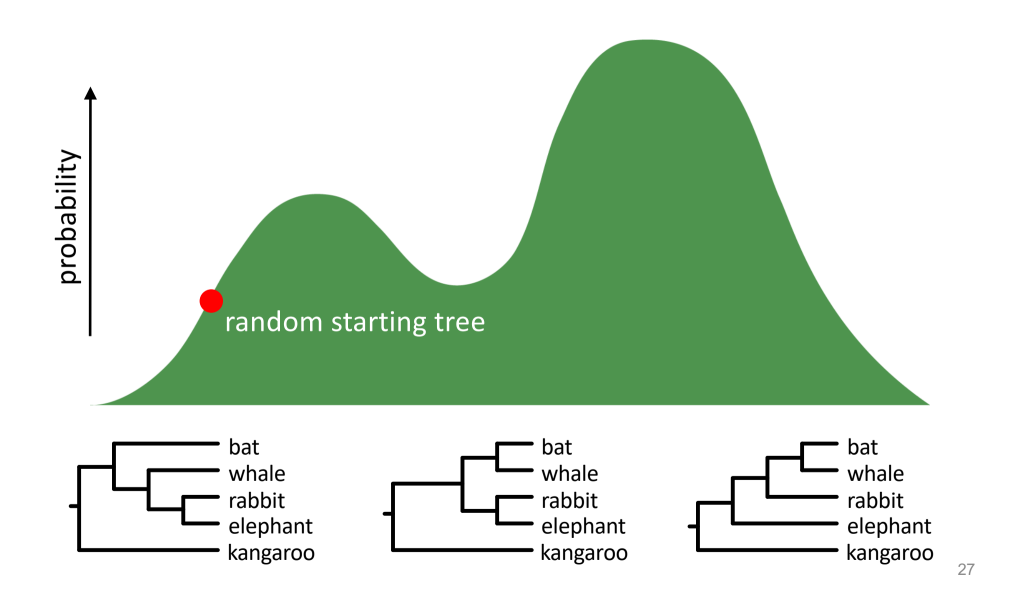


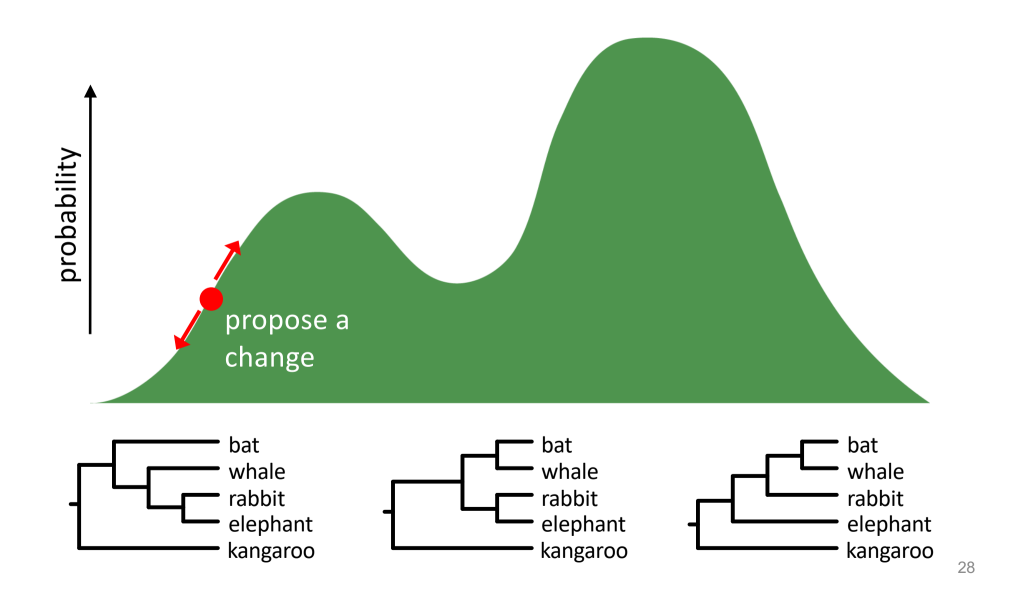
Estimating the posterior

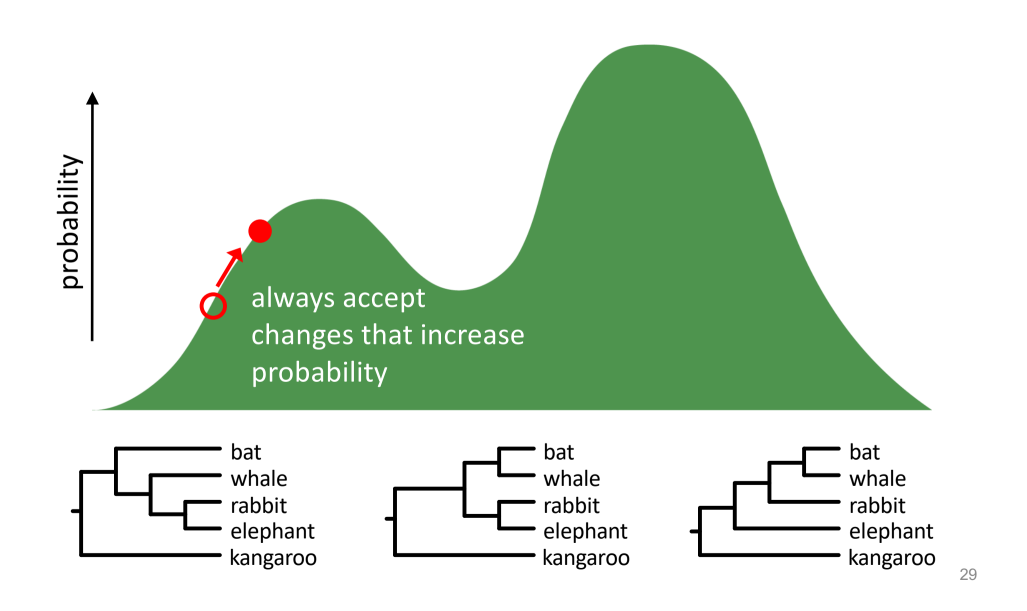
- Impossible to obtain the posterior directly
- Instead, the posterior can be estimated using Markov chain Monte Carlo simulation
- This is usually done using the Metropolis-Hastings algorithm

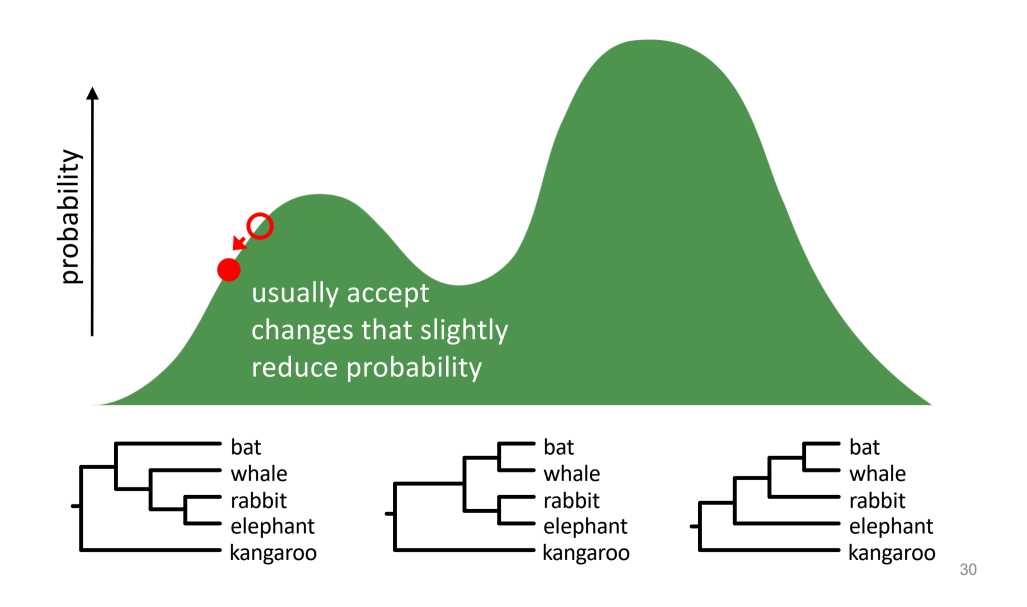


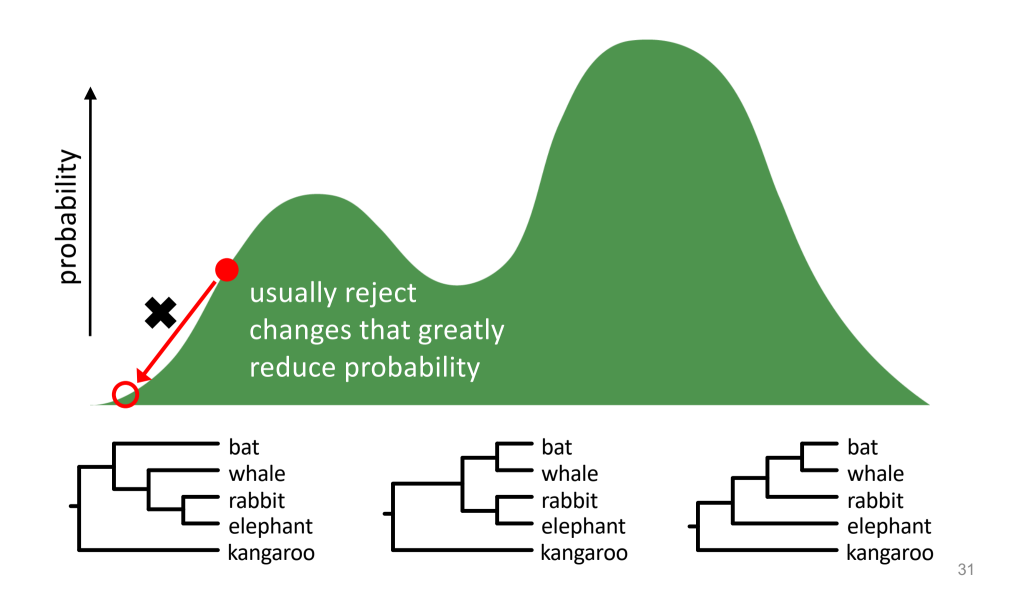
Nicholas Metropolis Los Alamos, 1953

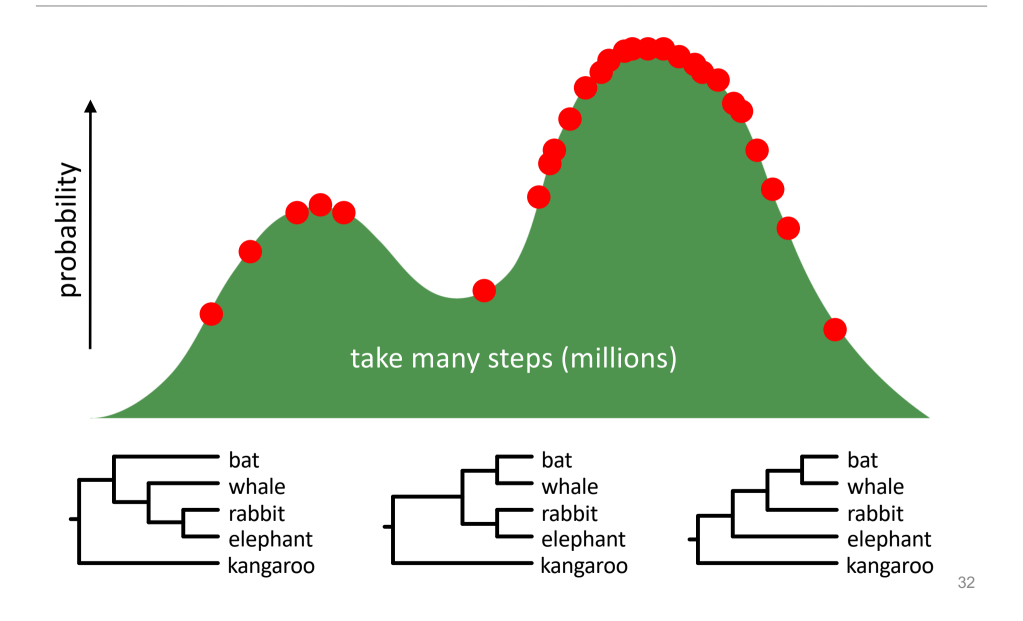




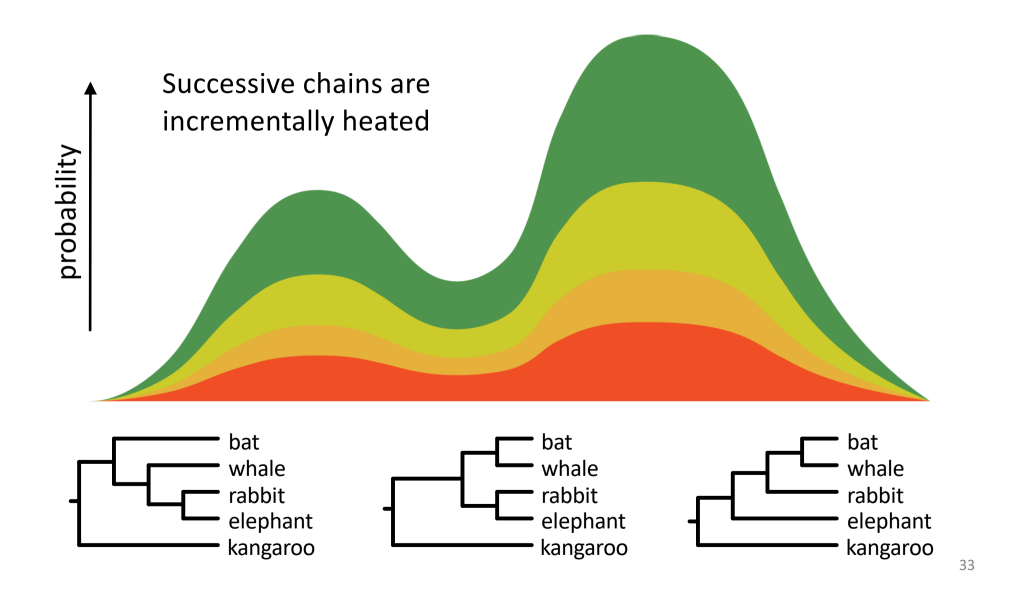




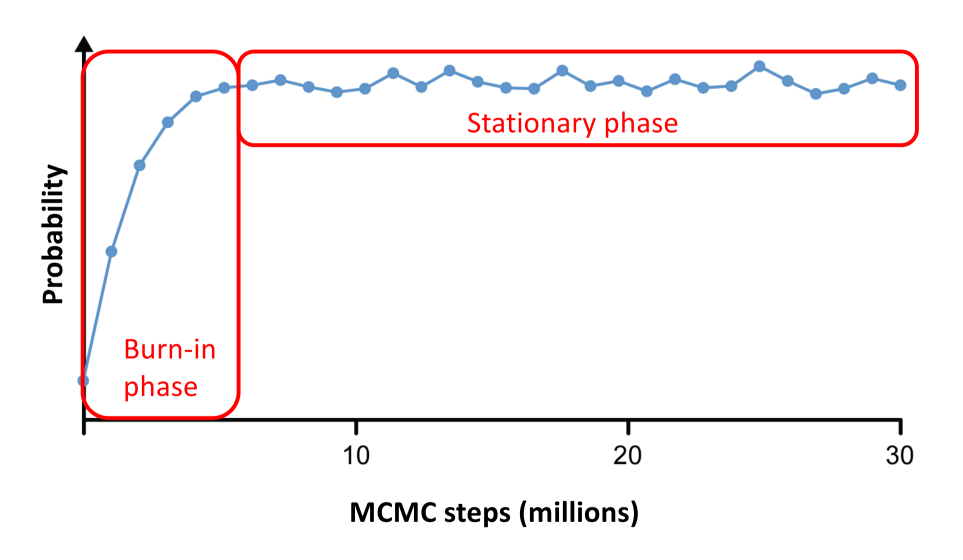


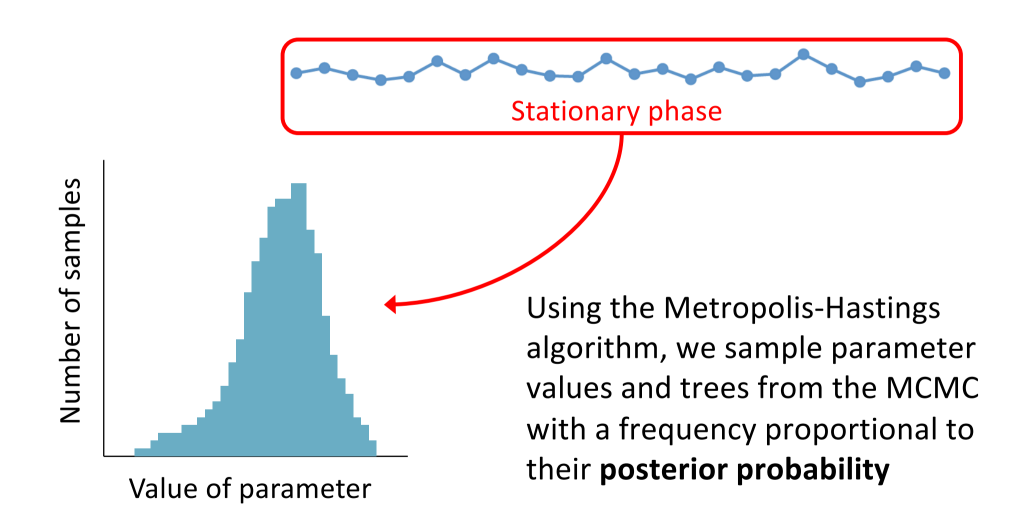


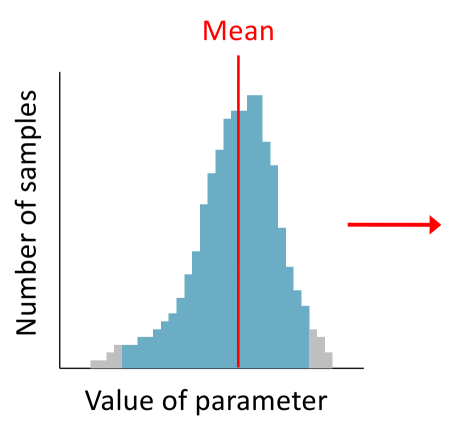
Metropolis-coupled MCMC



- Output from a Bayesian phylogenetic analysis:
 - A list of the **parameter values** visited by the Markov chain (.p file in *MrBayes*, .log file in *BEAST*)
 - A list of the **trees** visited by the Markov chain (.t file in *MrBayes*, .trees file in *BEAST*)





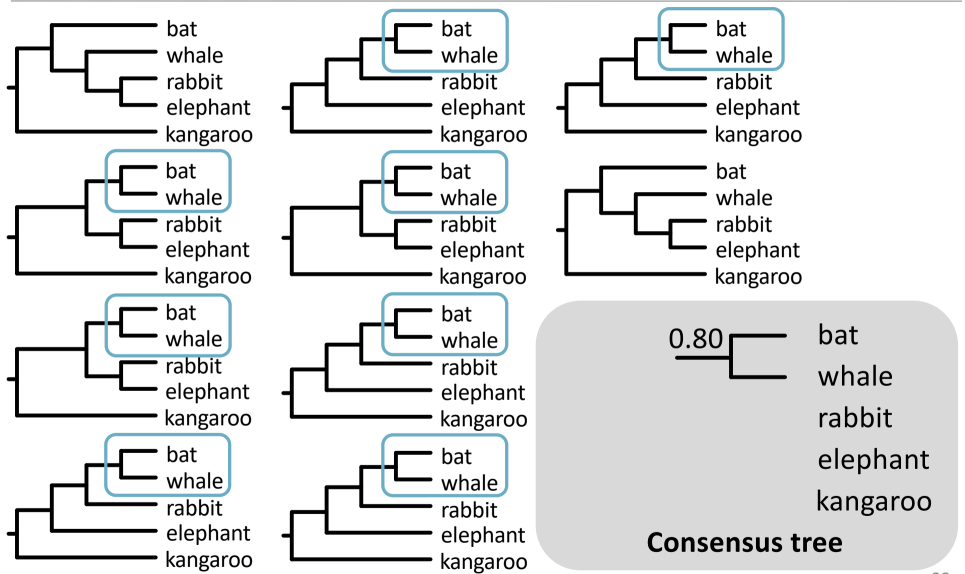


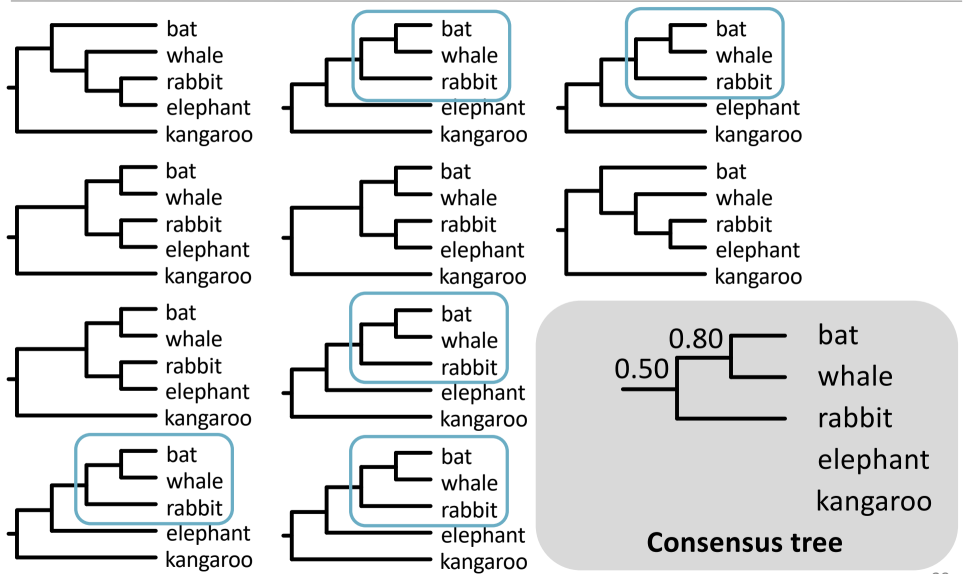
Take the mean of the sampled values

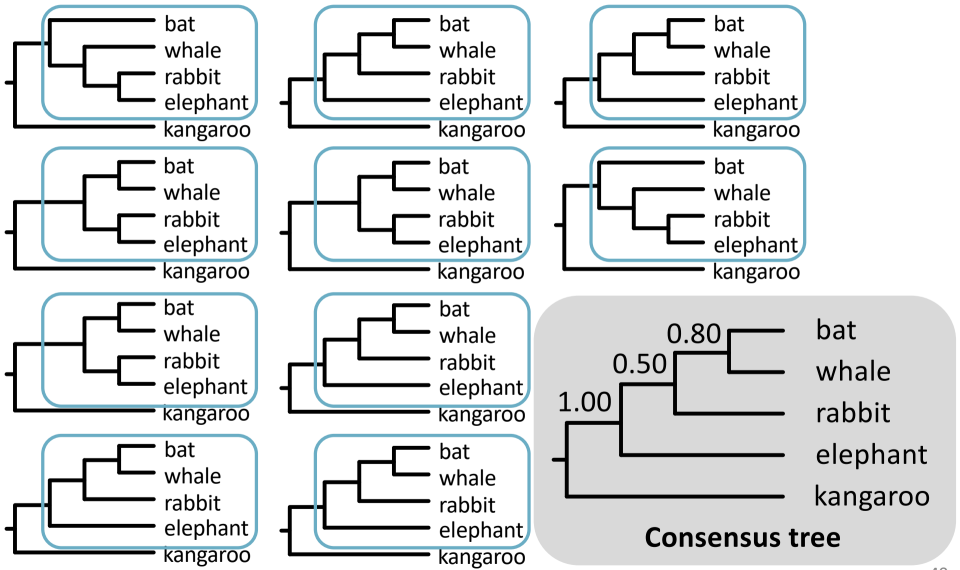
Mean posterior estimate

Take the 'central' 95% of the sampled values

95% credibility interval







- Majority-rule consensus tree (MrBayes)
 Shows all nodes with posterior probability >0.50
- Maximum a posteriori (MAP) tree
 Sampled tree with highest posterior probability
- Maximum clade credibility (MCC) tree (BEAST/TreeAnnotator)
 Sampled tree with highest sum or product of posterior node probabilities

Useful references

