Using BISSE with diversitree

Rich G. FitzJohn

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1 Introduction

This demo is in a very early stage of development. It does not aim to be a complete reference to the package, but should give some useful hints. The code all assumes that the diversitree package is loaded.

> library(diversitree)

2 Simulating phylogenies

diversitree includes code for simulating phylogenies, under both a character-dependent BISSE-type model and a character-independent model (as in Nee et al., 1994). The tree simulation code is likely to change a reasonable amount soon. It will be useful to have a simulated tree to demonstrate the code; the following generates a birth-death tree following the BISSE model with 203 species, using $\lambda_0 = 0.1$, $\lambda_1 = 0.2$, $\mu_0 = \mu_1 = 0.03$, and $q_{01} = q_{10} = 0.01$, starting in state 0 and running for 60 time units.

```
> pars <- c(0.1, 0.2, 0.03, 0.03, 0.01, 0.01)
> set.seed(2)
> phy <- tree.bisse(pars, max.t = 60, x0 = 0)
```

3 Running BISSE

The way diversitree runs BISSE, you first construct a likelihood function with make.bisse, then use this in a maximum likelihood or MCMC approach to do your inference. The make.bisse takes as its first two arguments a tree and set of character states:

```
> lik <- make.bisse(phy, phy$tip.state)
```

The object "lik" is the likelihood function; given a vector of parameters (in the order λ_0 , λ_1 , μ_0 , μ_1 , q_{01} , q_{10}) it computes the likelihood of the tree and character data:

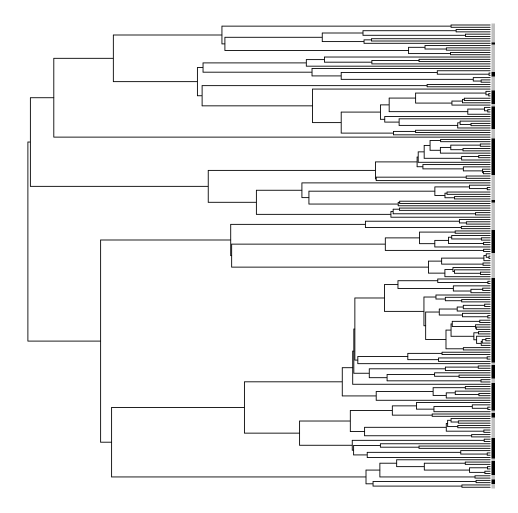


Figure 1: A random BISSE tree. Gray indicates state 0 (low speciation) and black indicates state 1 (high speciation).

```
> lik(pars)
[1] -660.0173
```

To do an maximum likelihood analysis, we need to find the set of parameters that maximises the likelihood. The find.mle function provides an interface to R's built-in optim that simplifies this process. To perform an optimisation, we need a starting point. The starting.point function does a very poor heuristic search for a sensible starting point, based on the character-independent birth-death fit.

```
> p <- starting.point(phy)</pre>
> p
                                                       q01
   lambda0
               lambda1
                               muO
                                           mu1
                                                                    q10
0.16721949 0.16721949 0.08017315 0.08017315 0.01740927 0.01740927
Run the optimisation (this may take up to a minute, depending on the speed of your computer).
> fit <- find.mle(lik, p)</pre>
> fit
$par
    lambda0
                 lambda1
                                  mu0
                                                mu1
0.099400026 0.196255619 0.023972305 0.032610865 0.010040429 0.009863226
$1nLik
[1] -659.9226
$counts
function gradient
      54
                54
$convergence
[1] 0
$message
[1] "CONVERGENCE: REL_REDUCTION_OF_F <= FACTR*EPSMCH"
attr(,"class")
[1] "mle.bisse"
```

The object returned by find.mle will change in the future, but the key bits are the element par with the ML parameters and lnLik with the best log-likelihood value. It would be preferable to start this search from multiple starting points, and I will add something to automate that to some degree soon.

3.1 Constrained models

To constrain a model, use the function constrain. This can be used to set parameters to a constant, to be the same as other parameters, or even to arbitrary expressions of free parameters. This is done through a formula interface that is still undergoing some change. As an example, consider the model where $\lambda_1 = \lambda_0$:

```
> lik.equal.l <- constrain(lik, lambda1 ~ lambda0)</pre>
```

The second argument to constrain describes the constraint. Additional arguments could impose different constraints; for example, setting μ_0 to zero.

```
> lik.equal.1.2 <- constrain(lik, lambda1 ~ lambda0, mu0 ~ 0)
```

This can be optimised as before, starting this time at the full model's MLE (find.mle here automatically does the filtering of the paramters, but you can alternatively specify a correct-length parameter vector).

This model has tweaked the extinction rates to get a suitable diversification rate, and settled on a λ that is intermediate between the true rates. The likelihood has dropped 3.5 units; a likelihood ratio test gives this a p value of 0.008:

In a second example, consider preventing $0 \rightarrow 1$ transitions:

```
> lik.no.01 <- constrain(lik, q01 ~ 0)
> fit.no.01 <- find.mle(lik.no.01, fit$par)
> fit.no.01[1:2]
```

```
$par
   lambda0
              lambda1
                              mu0
                                          mu1
                                                      q10
0.10568736 0.23544593 0.05834708 0.14417320 0.02559457
$lnLik
[1] -673.0052
This model is strongly rejected (p < 0.001)
> anova(fit, no.01 = fit.no.01)
           lnLik
                      AIC
                            ChiSq Pr(>|Chi|)
      Df
       6 -659.92 1331.85
full
no.01 5 -673.01 1356.01
                            26.17 3.134e-07
```

3.2 Markov Chain Monte Carlo (MCMC)

The likelihood function can also be used to perform Markov Chain Monte Carlo (MCMC). I will use an exponential prior with a rate of 5.7 (having a mean twice the character independent diversification rate). Running this will take some time (allow up to 1 s per step on this tree). The output is a large data frame with the contents of the chain.

```
> r <- 1/(2 * (p[1] - p[3]))
> ans <- mcmc(lik, fit$par, nsteps = 100, w = rep(0.1, 6), lower = rep(0,
+ 6), upper = rep(Inf, 6), prior = r, fail.value = -Inf)</pre>
```

4 Skeleton trees

To sample from the simulated phylogeny above to make a skeleton tree, we can do this:

```
> set.seed(1)
> n.taxa <- length(phy$tip.label)
> keep <- sort(sample(n.taxa, 50))
> phy.s <- drop.tip(phy, setdiff(seq_len(n.taxa), keep))</pre>
```

A need likelihood function is needed for this tree. We also need to specify what the sampling fraction is (close to 20% for both).

Pass this in to make.bisse and construct a new likelihood function that accounts for the sampling:

```
> lik.s <- make.bisse(phy.s, phy.s$tip.state, sampling.f = sampling.f)
This can then be optimised, as before:
> fit.s <- find.mle(lik.s, p)</pre>
> fit.s[1:2]
$par
    lambda0
                 lambda1
                                  mu0
                                               mu1
                                                            q01
                                                                         q10
0.080669502 0.227324661 0.000000000 0.080338704 0.010907117 0.005236501
$lnLik
[1] -193.9340
The fits compare reasonably well from the full tree and sampled tree
> round(rbind(full = fit$par, sampled = fit.s$par), 3)
        lambda0 lambda1
                            mu0
                                  mu1
                                         q01
full
          0.099
                   0.196 0.024 0.033 0.010 0.010
          0.081
                   0.227 0.000 0.080 0.011 0.005
sampled
```

5 Terminally unresolved trees

For this section, I will show how the shorebird example in the manuscript can be performed. The file Thomas-tree.nex contains the shorebird supertree assembled by Thomas et al. (2004) in nexus format and the file Lislevand-states.csv contains a measure of the level of sexual dimorphism, computed from the data in Lislevand et al. (2007). The tree contains many polytomies, so I am converting it into a terminally unresolved tree, with the polytomies.to.clades function.

```
> tree <- read.nexus("data/Thomas-tree.nex")
> tree2 <- polytomies.to.clades(tree)</pre>
```

> names(states) <- tree\$tip.label</pre>

Terminally unresolved trees can be assembled manually from a phylogeny that contains only the exemplar tips using the make.clade.tree function.

```
> states <- read.csv("data/Lislevand-states.csv", as.is = TRUE)
> states <- structure(states$dimorph, names = states$species)

Every species needs tip data
> states <- states[tree$tip.label]</pre>
```

This is all the information make.bisse needs. Construct a likelihood function by categorising states with 15% dimorphism being the cut-off:

```
> lik.sb <- make.bisse(tree2, abs(states) > 0.15)
```

This starting point comes from ape's function bd.ext, but this cannot be automated, as it does not return an object.

```
> p \leftarrow c(0.147, 0.147, 0, 0, 0.02, 0.02)
> names(p) \leftarrow c("lambda0", "lambda1", "mu0", "mu1", "q01", "q10")
```

Find the ML point (this will take 1-2 minutes, and will produce warnings that can be ignored).

```
> fit.sb <- find.mle(lik.sb, p)
> fit.sb[1:2]
```

\$par

```
lambda0 lambda1 mu0 mu1 q01 q10 0.05303850 0.20341423 0.00000000 0.00000000 0.04235417 0.28197098
```

\$lnLik

```
[1] -633.7833
```

It looks from this that the speciation rate and the character transition rate in state 1 are higher than the corresponding rates in state 0 ($\lambda_1 > \lambda_0$, $q_{10} > q_{01}$). The significance of this can be tested with a likelihood ratio test:

```
> lik.sb.l <- constrain(lik.sb, lambda1 ~ lambda0)
> lik.sb.q <- constrain(lik.sb, q10 ~ q01)
> fit.sb.l <- find.mle(lik.sb.l, fit.sb$par)
> fit.sb.q <- find.mle(lik.sb.q, fit.sb$par)</pre>
```

These constrained models are significantly worse fits than the full model (λ : p = 0.0005, q: p = 0.01).

> anova(fit.sb, equal.1 = fit.sb.1, equal.q = fit.sb.q)

```
Df lnLik AIC ChiSq Pr(>|Chi|)
full 6 -633.78 1279.57
equal.1 5 -639.88 1289.76 12.19 0.0004801
equal.q 5 -636.88 1283.77 6.20 0.0127691
```

This analysis can also be done with MCMC (not shown)

```
> ans <- mcmc(lik.sb, fit.sb$par, nsteps = 10000, w = rep(0.1,
+ 6), lower = rep(0, 6), upper = rep(Inf, 6), prior = 1/p[1] *
+ 2, fail.value = -Inf)</pre>
```

References

- Lislevand T., Figuerola J., and Székely T. 2007. Avian body sizes in relation to fecundity, mating system, display behavior, and resource sharing. Ecology 88:1605.
- Nee S., May R.M., and Harvey P.H. 1994. The reconstructed evolutionary process. Philos. Trans. R. Soc. Lond. B Biol. Sci. 344:305–311.
- Thomas G.H., Wills M.A., and Székely T. 2004. A supertree approach to shorebird phylogeny. BMC Evolutionary Biology 4:28.