Documentation of the R package epiphylo

Xavier Didelot

June 16, 2014

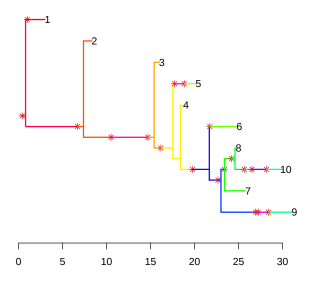
1 Simulation

A pathogen has an effective within-host population size of $N_e = 100$ and a generation time g = 1 day, so that $N_e g = 100/365$ years. The basic reproduction number is R = 1. To following command simulates an outbreak of this pathogen:

> simu <- simulateOutbreak(R=1,neg=100/365,pi=0.5)</pre>

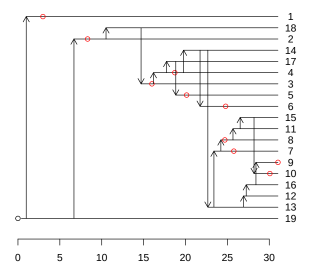
This simulation contains both the transmission tree between infected hosts and the within-host phlogenetic tree of each host. This can be visualised as a colored phlogenetic tree, where each host is represented by a unique color:

> plotBothTree(simu)



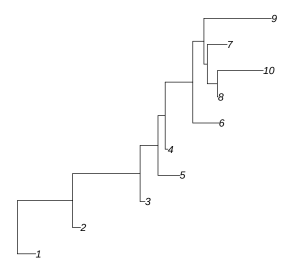
The transmission tree can be extracted and plotted separately from the phylogeny:

- > ttree<-ttreeFromFullTree(simu)
- > plotTTree(ttree)



The phylogenetic tree can be extracted and converted into a phylo object from the ape package:

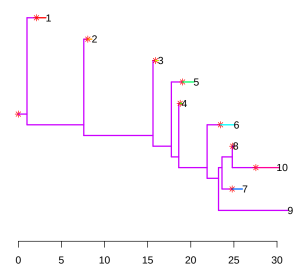
- > library(ape)
- > ptree<-ptreeFromFullTree(simu)
- > p<-phyloFromPtree(ptree)
- > plot(p)



2 Inference of transmission tree given a phylogeny

A phylo object can be turned into a phylogenetic tree and complemented with a wild guess at the transmission tree in order to provide the starting point of the MCMC procedure:

- > ptree<-ptreeFromPhylo(p)</pre>
- > full<-makeFullTreeFromPTree(ptree)</pre>
- > plotBothTree(full)



The MCMC procedure to infer the transmission tree given the phylogenetic tree can be run as follows:

> record<-inferTTree(ptree,mcmcIterations=100)</pre>

This returns a record of all MCMC iterations. Let's see what the transmission tree looked like at the end of the MCMC:

- > lastIteration<-record[[length(record)]]</pre>
- > plotBothTree(lastIteration\$tree)

