

# KDETrees Simulations

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## 1 Introduction

Here we present the code for the simulations found in the KDETrees article. These simulations compare the ability of KDETrees to find trees which were generated by a non-contained coalescent process, in a dataset consisting mostly of trees generated by a contained coalescent process. We test two scenarios: all contained coalescent trees are contained in a single species tree; and the contained coalescent trees are sampled from a mixed distribution, where the trees are contained in one of 5 different species trees. We also compare the performance of our method with that of the previously published Phylo-MCOA method.

The coalescent trees are generated using the methods found in the Dendropy python module. The script which generates the trees, as well as the species trees can be found in the `sim` directory of the KDETrees package.

## 2 Comparison of KDETrees and Phylo-MCOA

In this simulation we generate a test dataset which consists of one “outlier” tree, which is generated by an unconstrained coalescent process, and 100 “non-outlier” trees which are generated by constrained coalescent processes. The outlier trees used are those in the `species1.nex` file.

```
> out.trees <- read.nexus("sim/species1k.nex")
```

The non-outlier trees are created by the `tresim.py` script. This example call to the script will generate 100 coalescent trees with effective population size 800 for each species tree found in `species.nex`.

```
> system2("sim/treesim.py", c("-s", "sim/species.nex", "-n", 800, "-N", 100))
```

Thus, each simulated dataset consists of a single tree from `species1k.nex`, along with the output of a call to `tresim.py`.

The simulation routine does the following for each outlier tree: (1) generate the non-outlier trees, (2) call `kdetrees`, in both dissimilarity and topological modes, (3) call `pMCOA` in both modes, (4) determines whether or not the outlier tree is identified.

```
> sim <- function(neff, out.trees, sp.file="species.nex", ncoal=100, ...){
+   found.kde <- integer(2)
+   found.pmcoa <- integer(2)
+   for (tree in .uncompressTipLabel(out.trees)){
+     trees <- read.tree(text=system2("sim/treesim.py",
+                                     c("-n", neff, "-s", sp.file, "-N", ncoal), stdout=TRUE))
+     res.topo <- kdetrees(c(c(tree), trees), ..., use.blen=FALSE)
+     res.diss <- kdetrees(c(c(tree), trees), ..., use.blen=TRUE)
+     found.kde <- found.kde + c((1 %in% res.topo$outliers), (1 %in% res.diss$outliers))
+
+     res.topo <- detect.complete.outliers(pMCOA(c(c(tree), trees), distance="nodal"))
+   }
+ }
```

```

+   res.diss <- detect.complete.outliers(pMCOA(c(c(tree),trees),distance="patristic"))
+   found.pmcoa <- found.pmcoa + c(res.topo$TFgn[1], res.diss$TFgn[1])
+ }
+ res <- c(found.kde,found.pmcoa)/length(out.trees)
+ names(res) <- c("kde.topo","kde.diss","pmcoa.topo","pmcoa.diss")
+ res
+ }

```

The following wrapper runs the simulation for various effective population sizes using `mclapply`.

```

> sim2 <- function(out.trees,neff,nout=5,sp.file,ncoal=100){
+   result <- mclapply(neff,sim,out.trees,sp.file,ncoal,nout,mc.cores=4L)
+   out <- stack(as.data.frame(t(simplify2array(result))))
+   out$neff <- rep(neff,4)
+   out
+ }

```

Instead of the published pMCOA routines, we use a slightly modified version which removes most of the progress reporting present in the published script.

```

> source("sim/my-pmcoa.R")

```

We run the simulation at 10 equally log-spaced values of the effective population size.

```

> neff <- round(exp(seq(log(500),log(3000),len=10)))

```

This runs 100 simulation replicates with the contained coalescent trees all generated using the single species tree shown in Figure 1.

```

> result <- sim2(out.trees[1:100],neff,sp.file="sim/species.nex",ncoal=100)

```

This run uses 5 species trees (see Figure 2) to generate the contained coalescent trees.

```

> mix.result <- sim2(out.trees[1:100],neff,sp.file="species5.nex",ncoal=20)

```

Format the results as a data frame suitable for use with `ggplot2`. Results are shown in Figure 3.

```

> sim1.res <- rbind(cbind(result,model="single"),cbind(mix.result,model="mixed"))
> sim1.res$Distance <- sim1.res$Method <- sim1.res$ind
> levels(sim1.res$Method) <- list(kdetrees=c("kde.outliers","kde.topo"),
+                               pMCOA=c("pmcoa.outliers","pmcoa.topo"))
> levels(sim1.res$Distance) <- list(topographical=c("pmcoa.topo","kde.topo"),
+                                   dissimilarity=c("pmcoa.outliers","kde.outliers"))

```

### 3 Distribution of Tree Scores

Also of interest to us is the distribution of tree scores, depending on if the tree in question is an outlier tree or one of the contained coalescent trees. If these score distributions have significant overlap, then it will be difficult for the method to distinguish between the two categories of trees. On the other hand, if the distributions have little to no overlap, then identification of outliers will be easier.

To do this, we generate 500 coalescent trees contained in the tree in Figure 1, and run `kdetrees` on these trees alone to find the distribution of the non-outlier contained coalescent trees. To find the distribution of outlier tree scores, one non-contained tree from `species1k.nex` is appended to the contained trees, and the analysis re-run. This process is repeated for many outlier trees, and their scores are recorded. Results of the simulation are summarized in Figure 4.

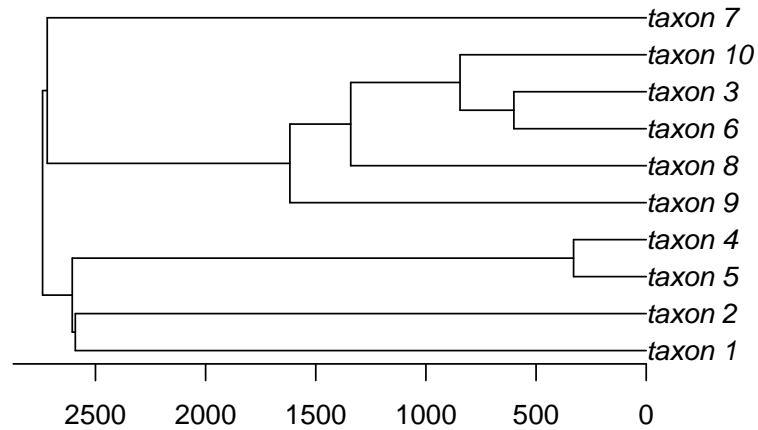


Figure 1: Species tree used for the single contained coalescent distribution simulation.

```
> simtrees <- read.tree(text=system2("./treesim.py",
+                                     c("-n",1500,"-s","species.nex","-N",500),stdout=TRUE))
> sim2.res <- kdetrees(simtrees,use.blen=TRUE)
> dsim <- function(tree) kdetrees(c(c(tree),simtrees),use.blen=TRUE)$density[1]
> sim2.resa <- simplify2array(mclapply(out.trees[1:100],dsim,mc.cores=4L))
> sim2 <- rbind(cbind(data.frame(value=unname(sim2.res$density)),type="coalescent"),
+               cbind(data.frame(value=unname(sim2.resa)),type="outlier"))
```

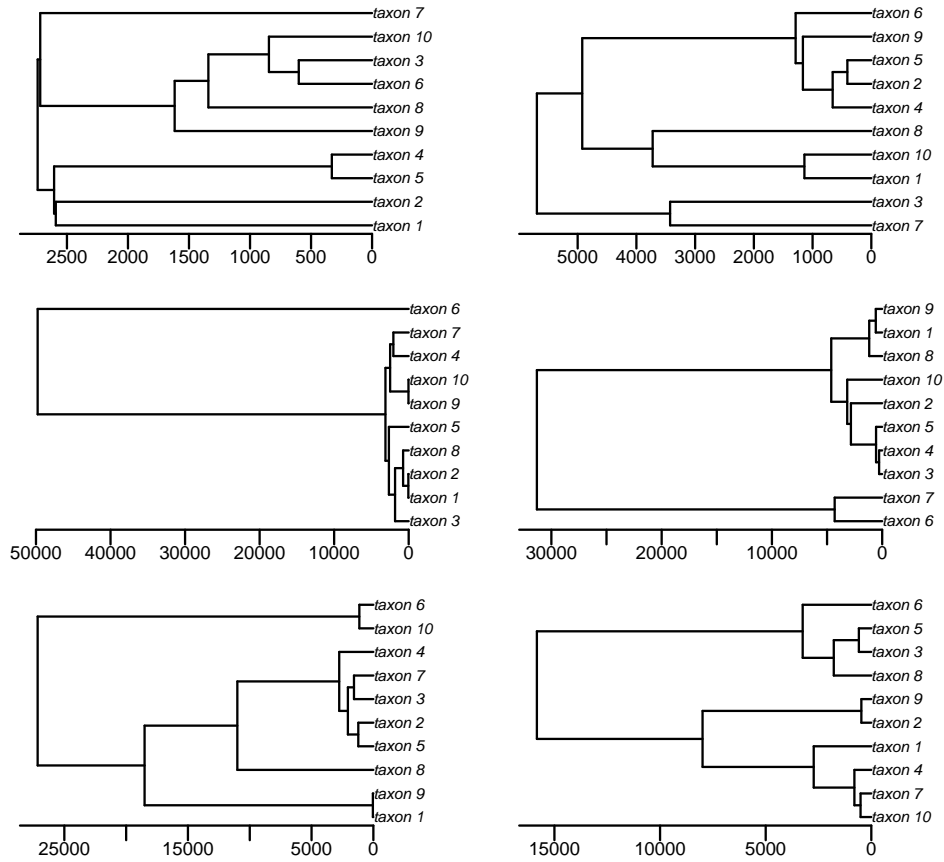


Figure 2: Species trees used for mixed coalescent simulation.

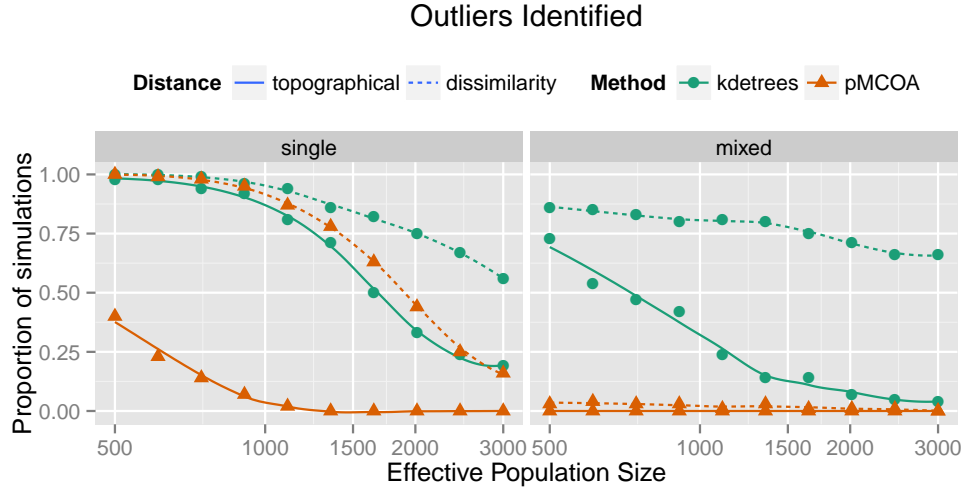


Figure 3: Results of simulation comparing KDETrees with Phylo-MCOA. We find that in this situation, the branch-length methods outperform the topology-only methods for both KDETrees and Phylo-MCOA. More striking however is the complete failure of Phylo-MCOA in the mixed distribution scenario, while KDETrees still performs quite well.

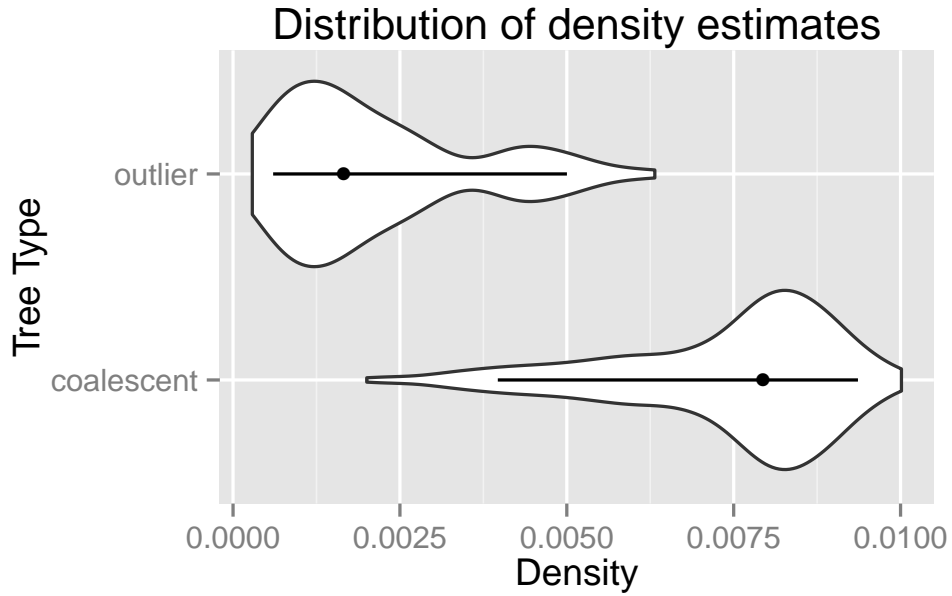


Figure 4: Comparison of the distribution of density scores for outlier trees and the contained coalescent trees.