

# kdetrees Manual

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## 1 Introduction

KDETrees is a tool for finding discordant phylogenetic trees. It takes as input a collection of trees (as a `ape::multiPhylo` object), and produces a *score* for each tree. High scores mean the tree is relatively similar to other trees in the sample, while low scores indicate that the tree in question may be discordant with the others. Low scoring trees are identified as putative *outliers*, and their contribution to the score calculation is removed. The result object contains scores for each tree, and a list of the putative outlier trees. If you use the program in your research, please cite our paper:

G. Weyenberg, P. Huggins, C. Schardl, D.K. Howe, and R. Yoshida.  
kdetrees: Nonparametric estimation of phylogenetic tree distributions. *Bioinformatics*, 2014.

## 2 Using kdetrees

The simplest method of using the software is the `kdetrees.complete` function. This is a convenience function which will do all the steps of the analysis at once: importing trees from file, performing the analysis, and producing output files. Basic use is to simply pass it the filename of a Newick file containing the trees to be analyzed. It will write several result files to the R working directory (`getwd`).

The call below assumes there is a file containing newick formatted trees named `trees.tre` in the current working directory. It will write out 4 files: `outliers.tre` a newick file containing only the trees identified as outliers; `results.csv` a csv files with the density estimates; `plot.png` and `hist.png` are diagnostic images. The names of these output files can be customized with further arguments to `kdetrees.complete`.

```
> kdetrees.complete("trees.tre")
```

The `kdetrees.complete` function also accepts any of the parameters accepted by the `kdetrees` function, as described in Sections 2.2 and 3.

## 2.1 Importing Trees

Trees may be imported using any of the methods provided by the `ape` package. (See `?read.tree` and `?read.nexus` for examples.) In the following examples, many functions are a part of the `ape` package, and it is recommended that you import it. For example, to load the example `apicomplexa` dataset, I placed the Newick tree strings into a file named `apicomplexa.tre` file and ran the commands:

```
> library(ape)
> apicomplexa <- read.tree("apicomplexa.tre")
```

*NB: The `apicomplexa` dataset is already included in the `kdetrees` package as example data, you do not need to import it. This is purely an example showing how to import your own trees.*

## 2.2 Running `kdetrees`

The simplest way to run `kdetrees` is to call the function of the same name, with the list of trees as the first argument.

```
> result <- kdetrees(apicomplexa)
> result
```

```
Call: kdetrees(trees = apicomplexa)
```

```
Density estimates:
```

	Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
	0.9963	65.8700	82.6400	77.3000	94.4900	114.5000

```
Cutoff: 22.92261
```

```
Outliers detected:
```

```
[1] 488.tre 497.tre 515.tre 546.tre 547.tre 641.tre 660.tre
[8] 662.tre 728.tre 747.tre 773.tre 780.tre
```

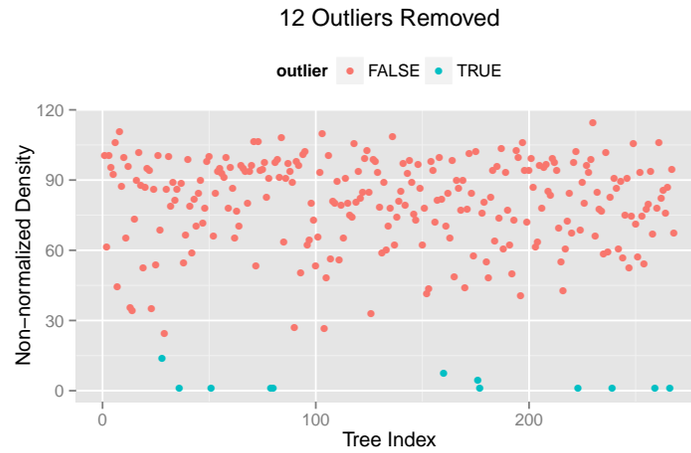
There are 3 main settings which control the method used in the analysis: the outlier detection tuning parameter (`k`), the distance computation method (`distance`), and whether or not to include branch length information in the distance calculation (`topo.only`). The default options are `distance="geodesic"`, `topo.only=FALSE`, and `k=1.5`.

For example, this call uses topology-based dissimilarity map distance.

```
> kdetrees(apicomplexa, k=1.25, distance="diss", topo.only=TRUE)
```

One can plot or `hist` the result object to create diagnostic plots. The `plot` and `hist` methods use the `ggplot2` package, not base graphics, thus you can modify them as you see fit. See Figure 1 for example plots.

```
> plot(result)
```



```
> hist(result)
```

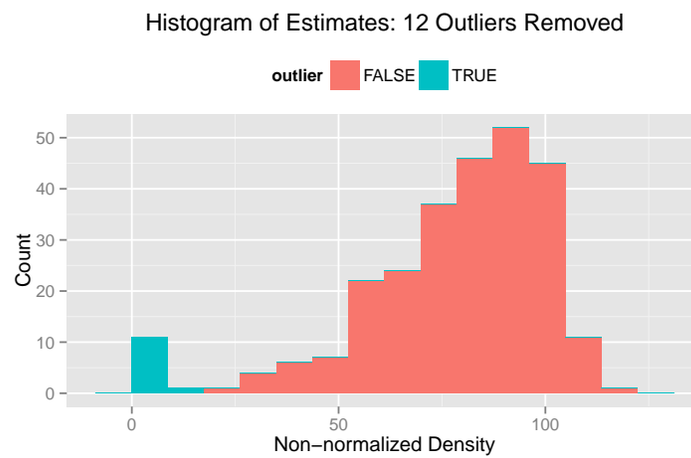


Figure 1: Diagnostic plots can be created with `plot` and `hist`.

## 2.3 Results

The result object is a list with three components, as well as several attributes that are used internally. The first element, `density`, has the computed score for each tree in the input list. This is the variable displayed in the diagnostic plots. The second element, `i`, contains the indices of the low scoring trees which were not included in the calculations. Finally, the `outliers` element contains the trees which were identified as outliers.

One might then wish to look at a plot of the putative outlier trees. Here I plot the lowest scoring tree in the apicomplexa dataset. It appears that something bad happened during the reconstruction of this tree, causing one branch to be much longer than the others.

```
> plot(result$outliers[[1]])
```

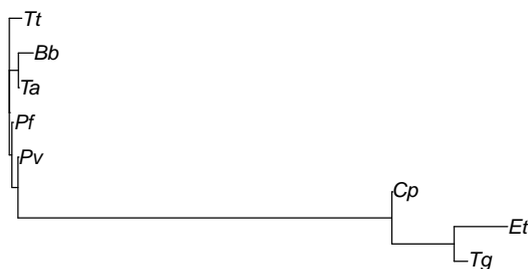


Figure 2: A plot of an outlying tree.

An `as.data.frame` method is provided for the result object, which can be used to export the results using the standard R methods. (Such as `write.csv`.) The list of outlier trees can be exported using `ape::write.tree`

```
> write.tree(result$outliers, file="outliers.tre")  
> result.df <- as.data.frame(result)  
> write.csv(result.df, file="scores.csv")
```

## 3 Advanced Options

### 3.1 Distance Methods

There are currently two methods implemented for the `distance` option in `kde-trees`: The "geodesic" method, based on work by Billera et al. [2001], as well

as that of Owen and Provan [2011]; and the "dissimilarity" map method, which utilizes pairwise tip-to-top distances.

Each of these distances can use either branch lengths supplied by the input trees, `topo.only=FALSE`, or the branch lengths can be ignored by setting `topo.only=TRUE`.

## 3.2 Bandwidth Selection

Currently, `kdtrees` uses an adaptive bandwidth method based on a nearest-neighbor calculation by default. It is possible to control the number of trees used to define the neighborhood, or disable the adaptive method entirely and provide a constant bandwidth, using the `bw` parameter.

If the `bw` parameter is supplied a list, the list is used as a set of parameters for a call to `bw.nn`. Currently, the only interesting setting that may be passed in this way is `prop`, which controls the proportion of the set of trees used to define the neighborhoods. For example, to change the neighborhood to include 50% of the sample we pass the following option.

```
> kdtrees(apicomplexa, bw=list(prop=0.5))
```

If `bw` is set to a single number, a constant bandwidth is used.

```
> kdtrees(apicomplexa, bw=6)
```

If `bw` is a vector, the values are used as bandwidths for the corresponding trees in the sample, possibly recycling the values if needed.

## 3.3 Command Line Interface

CLI use can be achieved by using the `Rscript` executable included with R. For example, this CLI command replicates the first example call in Section 2.

```
$ Rscript -e 'library(kdtrees); kdtrees.complete("trees.tre")'
```

The desired R commands can also be placed into a R script file (e.g. `myscript.R`), and run using

```
$ Rscript myscript.R
```

## References

- Louis J Billera, Susan P Holmes, and Karen Vogtmann. Geometry of the space of phylogenetic trees. *Advances in Applied Mathematics*, 27(4):733–767, 2001.
- Megan Owen and J Scott Provan. A fast algorithm for computing geodesic distances in tree space. *IEEE/ACM Transactions on Computational Biology and Bioinformatics (TCBB)*, 8(1):2–13, 2011.
- G. Weyenberg, P. Huggins, C. Schardl, D.K. Howe, and R. Yoshida. `kdtrees`: Nonparametric estimation of phylogenetic tree distributions. *Bioinformatics*, 2014.