Getting started with Quartet

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This document should contain all you need to get started measuring tree distances with 'Quartet.' If you get stuck, please let me know so I can improve this documentation.

Loading trees

Instructions for loading phylogenetic trees into R can be found in a separate vignette. For these examples, we'll enter two simple trees by hand:

```
tree1 <- ape::read.tree(text = '(A, ((B, (C, (D, E))), ((F, G), (H, I))));')
tree2 <- ape::read.tree(text = '(A, ((B, (C, (D, (H, I)))), ((F, G), E)));')</pre>
```

Calculating distances

We can calculate distances between pairs of trees using the 'Quartet' package.

First we'll install the package. We can either install the stable version from the CRAN repository:

```
install.packages('Quartet')
```

or the development version, from GitHub – which will contain the latest features but may not be as extensively tested:

```
devtools::install_github('ms609/Quartet')
```

Then we'll load the package into R's working environment:

```
library('Quartet')
```

Now the package's functions are available within R. Let's proceed to calculate some tree distances.

Pairs of trees

Calculating the distance between two trees is a two stage process. For a quartet distance, we first have to calculate the status of each quartet:

```
statuses <- QuartetStatus(tree1, tree2)
```

Then we convert these counts into a distance metric (or similarity measure) that suits our needs – perhaps the Quartet Divergence:

```
QuartetDivergence(statuses, similarity = FALSE)
```

[1] 0.6031746

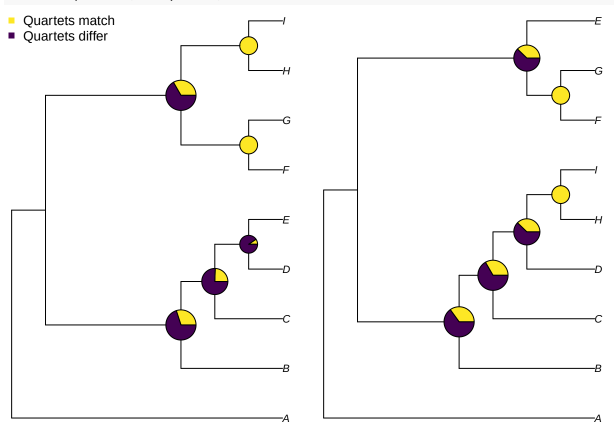
We can calculate all similarity metrics at once using:

SimilarityMetrics(statuses, similarity = TRUE)

```
##
        DoNotConflict ExplicitlyAgree StrictJointAssertions
## [1,]
            0.3968254
                            0.3968254
                                                   0.3968254
##
        SemiStrictJointAssertions SymmetricDifference MarczewskiSteinhaus
## [1,]
                        0.3968254
                                             0.3968254
                                                                  0.2475248
        SteelPenny QuartetDivergence SimilarityToReference
##
## [1,] 0.3968254
                           0.3968254
                                                  0.3968254
```

It can be instructive to visualize how each split in the tree is contributing to the quartet similarity:

VisualizeQuartets(tree1, tree2)



Rather than using quartets, we might want to use partitions as the basis of our comparison:

SimilarityMetrics(SplitStatus(tree1, tree2))

```
##
        DoNotConflict ExplicitlyAgree StrictJointAssertions
## [1,]
             0.3333333
                               0.3333333
                                                        0.3333333
##
        {\tt SemiStrictJointAssertions} \ \ {\tt SymmetricDifference} \ \ {\tt MarczewskiSteinhaus}
## [1,]
                           0.3333333
                                                 0.3333333
                                                                              0.2
##
         SteelPenny QuartetDivergence SimilarityToReference
## [1,] 0.3333333
                              0.3333333
                                                      0.3333333
```

Multiple comparisons

If you have more than two trees to compare, you can send a list of trees (class: list or multiPhylo) to the distance comparison function.

You can calculate the similarity between one tree and a forest of other trees:

```
library('TreeTools', quietly = TRUE, warn.conflicts = FALSE)
oneTree <- CollapseNode(as.phylo(0, 11), 14)</pre>
twoTrees <- structure(list(bal = BalancedTree(11), pec = PectinateTree(11)),</pre>
                       class = 'multiPhylo')
status <- SharedQuartetStatus(twoTrees, cf = oneTree)</pre>
QuartetDivergence(status)
##
                    pec
         bal
## 0.4939394 0.6272727
Or between one tree and (itself and) all other trees in the forest:
forest <- as.phylo(0:5, 11)</pre>
names(forest) <- letters[1:6]</pre>
status <- SharedQuartetStatus(forest)</pre>
QuartetDivergence(status)
                                 С
## 1.0000000 0.9757576 0.9757576 0.9333333 0.9121212 0.9333333
Or between each pair of trees in a forest:
status <- ManyToManyQuartetAgreement(forest)</pre>
QuartetDivergence(status, similarity = FALSE)
##
## a 0.00000000 0.02424242 0.02424242 0.06666667 0.08787879 0.06666667
## b 0.02424242 0.00000000 0.02424242 0.08787879 0.06666667 0.06666667
## c 0.02424242 0.02424242 0.00000000 0.08484848 0.08484848 0.04242424
## d 0.06666667 0.08787879 0.08484848 0.00000000 0.04242424 0.04242424
## e 0.08787879 0.06666667 0.08484848 0.04242424 0.00000000 0.04242424
## f 0.06666667 0.06666667 0.04242424 0.04242424 0.04242424 0.00000000
Or between one list of trees and a second:
status <- TwoListQuartetAgreement(forest[1:4], forest[5:6])</pre>
QuartetDivergence(status, similarity = FALSE)
## a 0.08787879 0.06666667
## b 0.06666667 0.06666667
## c 0.08484848 0.04242424
## d 0.04242424 0.04242424
```

Other calculations

To calculate how many quartets are unique to a certain tree (akin to the partitionwise equivalent ape::prop.clades), use:

```
interestingTree <- as.phylo(42, 7)
referenceTrees <- list(BalancedTree(7), PectinateTree(7))
status <- CompareQuartetsMulti(interestingTree, referenceTrees)</pre>
```

status['x_only'] = 23 quartets are resolved in a certain way in interestingTree, but not resolved that way in any referenceTrees.

What next?

You may wish to:

- Read more about Quartet distances
- $\bullet\,$ Review alternative distance measures and corresponding functions
- Interpret or contextualize tree distance metrics