dispRity manual

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dispRity is a package for calculating disparity in R. It allows to summarise ordinated matrices (e.g. MDS, PCA, PCO, PCoA) into single values.

Contents

1	Before starting 1			
	1.1	Glossary	1	
	1.2	Installation		
	1.3	Data		
	1.4	A quick go through	4	
2	Pacl	kage specificities	5	
	2.1	The dispRity objects	5	
	2.2	Modular functions	7	
3				
	3.1	cust.series	8	
	3.2	time.series	9	
	3.3	boot.matrix		
	3.4	dispRity		
	3.5		14	
	3.6	·	15	
	3.7		19	
	3.8		19	
4	Developments 20			
		•	20	
			20	

1 Before starting

1.1 Glossary

Because this package is aimed to be multidisciplinary, many names or terms used in this tutorial might be non-familiar to certain fields. Here is a list of what are the exact meaning of these term:

• Ordinated space: it designates the mathematical multidimensional object studied here. In morphometrics, this one is often referred as being the morphospace. However it can also be the the cladisto-space for cladistic data or the eco-space in ecology, etc. In practice, this term designates an ordinated matrix where the columns represent the dimensions of the ordinated space (often – but not necessarily – > 3!) and the rows represent the elements within this space.

- Elements: it designates the rows of the ordinated space, elements can be either taxa, field sites, countries, etc...
- **Dimensions**: it designates the columns of the ordinated space. The dimensions can also be referred to as axis.
- Series: it designates sub-samples of the ordinated space. Basically a series contain the same number of dimensions as the morphospace but might contain a smaller number of elements. For example, if our ordinated space is composed of birds and mammals (i.e. the elements) and 50 dimensions, we can create two series of just mammals or birds as elements (but the same 50 dimensions) to look at the difference in disparity between both groups.

1.2 Installation

You can install this package easily if you use the latest version of R and devtools.

```
if(!require(devtools)) install.packages("devtools")
install_github("TGuillerme/dispRity", ref = "release")
library(dispRity)
```

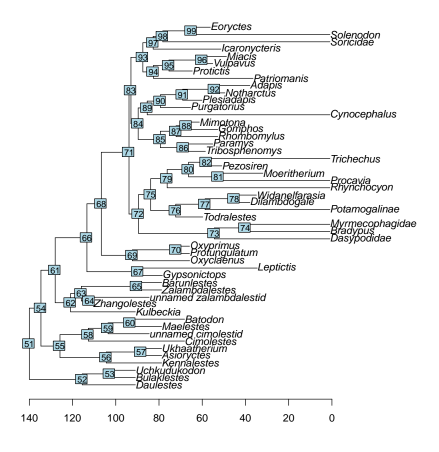
Note that we use the release branch here which is version 0.1.2. For the piping-hot (but potentially full of bugs) version, you can change the argument ref = "release" to ref = "master". This package depends mainly on the ape package and the timeSliceTree::paleotree function.

1.3 Data

In this tutorial we are going to use a subset of the ordinated cladistic data from Beck and Lee (2014) that contains 50 taxa (elements) ordinated using their cladistic distance (i.e. the distance between their discrete morphological characters). Note that this data is more oriented towards palaebiology analysis but that it can apply to other disciplines. Please refer to the GitHub page: github.com/TGuillerme/dispRity for other vignettes covering some specific example.

```
## Loading the package
library(dispRity)
## Loading required package: paleotree
## Setting the random seed for repeatability
set.seed(123)
## Loading the ordinated matrix containing 50 taxa
data(BeckLee_mat50)
dim(BeckLee_mat50)
## [1] 50 48
head(BeckLee_mat50[,1:5])
##
                    [,1]
                                 [,2]
                                            [,3]
                                                     [,4]
## Cimolestes
             -0.4087147 0.0139690317 0.26268300 0.2297096 0.1310953
## Maelestes
## Batodon -0.6923194 0.3308625215 -0.10175223 -0.1899656 0.1003108
## Bulaklestes -0.6802291 -0.0134872777 0.11018009 -0.4103588 0.4326298
## Daulestes -0.7386111 0.0009001369 0.12006449 -0.4978191 0.4741342
## Uchkudukodon -0.5105254 -0.2420633915 0.44170317 -0.1172972 0.3602273
```

```
## Loading another ordinated matrix containing 50 tips + 49 nodes
data(BeckLee_mat99)
dim(BeckLee_mat99)
## [1] 99 97
head(BeckLee_mat99[,1:5], 2)
##
                    [,1]
                                          [,3]
                                                     [, 4]
                               [,2]
## Cimolestes -0.6082437 -0.0323683 0.08458885 -0.4338448 -0.30536875
## Maelestes -0.5730206 -0.2840361 0.01308847 -0.1258848 0.06123611
tail(BeckLee_mat99[,1:5], 2)
##
              [,1]
                        [,2]
                                   [,3]
                                              [,4]
                                                         [.5]
## n48 -0.05529018 0.4799330 0.04118477 0.04944912 -0.3558830
## n49 -0.13067785 0.4478168 0.11956268 0.13800340 -0.3222785
## Loading a list of first and last occurrence data
data(BeckLee_ages)
head(BeckLee_ages)
              FAD LAD
## Adapis 37.2 36.8
## Asioryctes 83.6 72.1
## Leptictis 33.9 33.3
## Miacis
            49.0 46.7
## Mimotona 61.6 59.2
## Notharctus 50.2 47.0
## Loading the phylogeny
data(BeckLee_tree)
plot(BeckLee_tree) ; nodelabels(cex=0.8) ; axisPhylo(root=140)
```

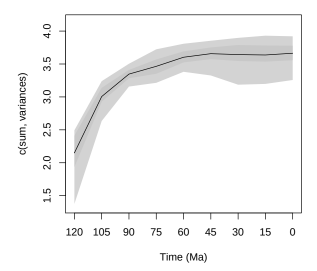


1.4 A quick go through

Here is a really crude and quick analysis to go through the package, showing some of its features. Note that all these features will be discussed in more details below.

```
## Splitting the data
sliced_data <- time.series(BeckLee_mat99, BeckLee_tree, method = "continuous",</pre>
    model = "acctran", time = rev(seq(from=0, to=130, by=15)),
    FADLAD = BeckLee_ages)
## Some tips have no FAD/LAD and are assumed to be single points in time.
## Bootstrapping the data
bootstrapped_data <- boot.matrix(sliced_data, 100)</pre>
## Calculating disparity
sum_of_variances <- dispRity(bootstrapped_data, metric = c(sum, variances))</pre>
## Summarising the results
summary(sum_of_variances)
                                       25%
                                             75% 97.5%
     series n observed mean 2.5%
## 1
                  2.699 2.150 1.376 1.937 2.409 2.493
## 2 105 11 3.275 3.006 2.634 2.928 3.137 3.239
```

```
## 3
         90 18
                  3.534 3.347 3.157 3.284 3.410 3.504
## 4
         75 19
                  3.660 3.465 3.214 3.352 3.568 3.723
                  3.805 3.603 3.382 3.525 3.691 3.807
## 5
         60 20
## 6
         45 14
                  3.956 3.657 3.326 3.575 3.751 3.853
## 7
                  4.055 3.643 3.186 3.546 3.788 3.897
         30 10
## 8
         15 10
                  4.055 3.637 3.198 3.537 3.780 3.930
          0 10
## 9
                  4.055 3.663 3.257 3.557 3.777 3.921
plot(sum_of_variances)
## Testing the effect of time on disparity
summary(test.dispRity(sum_of_variances, test = aov, comparisons = "all"))
##
                Df Sum Sq Mean Sq F value Pr(>F)
                 8 199.71 24.964
                                    755.5 <2e-16 ***
## series
                   29.44
## Residuals
                            0.033
               891
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```



2 Package specificities

2.1 The dispRity objects

Disparity analysis can involve a lot of shuffling around with many matrices (especially when bootstrapping the data) which can be a bit impractical to visualise and quickly jam your R console. For example, we can have a look at the structure of the object created in the quick example:

```
str(sum_of_variances)
## That's a more than 4000 lines of output!
```

Therefore this package proposes a new class of object called dispRity objects. These objects allow to easily use a S3 method functions such as summary.dispRity (just called as summary; see section 3.5) or plot.dispRity (just called as plot; see section 3.6). But also, this allows to use the S3 method for printing dispRity objects via print.dispRity that allows to summarise the content of the objects similar to the phylo class objects (see print.phylo::ape).

```
## What is the class of the sum_of_variances object?
class(sum_of_variances)
## [1] "dispRity"
## What is in the object?
names(sum_of_variances)
## [1] "data"
                   "disparity" "elements" "series"
                                                        "call"
## Summarising it using the S3 method print.dispRity:
sum_of_variances
## Disparity measurements across 9 series for 99 elements
## Series:
## 120, 105, 90, 75, 60, 45 ...
## Disparity calculated as: c(sum, variances) for 97 dimensions.
## Data was split using continuous method.
## Data was bootstrapped 100 times, using the full bootstrap method.
## This displays some basic information about the object content
```

Note however, that it is always possible to recall the full object using the argument all=TRUE:

```
## Displaying the full object
print(sum_of_variances, all = TRUE)
```

Finally, some utility functions such as get.dispRity or extract.dispRity allows to access to some specific content of the object:

```
## Extracting some specific series from the disparity object
series_1_and_4 <- get.dispRity(sum_of_variances, what = c(1,4))</pre>
series_1_and_4
## Disparity measurements across 2 series for 24 elements
## Series:
## 120, 75.
## Disparity calculated as: c(sum, variances) for 97 dimensions.
## Data was split using continuous method.
## Data was bootstrapped 100 times, using the full bootstrap method.
## The observed disparity
extract.dispRity(sum_of_variances)
        120
                 105
                           90
                                    75
                                             60
                                                      45
                                                                30
                                                                         15
## 2.698895 3.274613 3.533596 3.660471 3.804800 3.955998 4.054575 4.054575 4.054575
## The list of bootstrapped scores of disparity
str(extract.dispRity(sum_of_variances, observed = FALSE))
## List of 9
## $ 120: num [1:100] 2.41 1.9 2.46 2.24 1.49 ...
## $ 105: num [1:100] 2.86 2.83 3.14 2.78 3.07 ...
## $ 90 : num [1:100] 3.48 3.51 3.38 3.42 3.31 ...
## $ 75 : num [1:100] 3.22 3.45 3.48 3.7 3.44 ...
## $ 60 : num [1:100] 3.59 3.76 3.46 3.43 3.5 ...
```

```
## $ 45 : num [1:100] 3.67 3.47 3.53 3.62 3.55 ...

## $ 30 : num [1:100] 3.32 3.79 3.7 3.87 3.56 ...

## $ 15 : num [1:100] 3.81 3.5 3.86 3.32 3.34 ...

## $ 0 : num [1:100] 3.9 3.67 3.59 3.65 3.63 ...
```

2.2 Modular functions

This package aims to be a modular package where users can personalise some aspects of the package fairly easily. In this version 0.1.2 only one function is fully modular (dispRity; see section 3.4) but more will be hopefully available in the near future (see section 4).

The modular idea is that users can use implemented tools to help facilitating their own personalised function. For example, the dispRity function intake a metric argument designating how disparity should be calculated. This argument can take some already implemented functions such as mean but also some completely new ones such as the sum divided by length:

```
## Disparity measured as the mean value of the ordinated matrix
summary(dispRity(sliced_data, metric = mean))
    series n observed
##
## 1
      120 5 -0.005
       105 11
## 2
                -0.010
## 3
       90 18 -0.010
## 4
        75 19
               -0.001
## 5
        60 20
                 0.005
## 6
        45 14
               0.013
## 7
       30 10
               0.017
## 8
        15 10
               0.017
## 9
        0 10
                 0.017
## A function for measuring the sum divided by the length (the mean!)
sum.divide.by.length <- function(X) sum(X)/length(X)</pre>
## Disparity measured as the mean divided by the length
summary(dispRity(sliced_data, metric = sum.divide.by.length))
##
    series n observed
## 1
     120 5 -0.005
## 2
     105 11 -0.010
## 3
        90 18
                -0.010
## 4
        75 19
                -0.001
## 5
        60 20
               0.005
## 6
        45 14
                 0.013
## 7
        30 10
                 0.017
## 8
       15 10
                 0.017
## 9
        0 10
                 0.017
## Giving the exact same results!
```

For more information concerning this modularity, please refer to the GitHub page: github.com/TGuillerme/dispRity for the vignette about the metric implementation in dispRity.

3 Functions

One of the first steps in disparity analysis is to split the ordinated space into sub-samples of this space (hereafter *series*). These series correspond to regions of the ordinated space that we want to compare to

each other.

3.1 cust.series

The cust.series function allows to create these series according to factors determined by the user. In this example, we can split the matrix based on the phylogeny and separate to different groups of taxa: the crown mammals (all the living mammals and their direct ancestor) and the stem mammals (all the mammals that have no direct offspring).

```
## We want to separate the species around the node 71 (see phylogeny above).
## All the descendant of these node are crown and all the ancestors are stem
crown <- extract.clade(BeckLee_tree, node = 71)$tip.label</pre>
stem <- drop.tip(BeckLee_tree, tip = crown)$tip.label</pre>
## We then have to feed this information in a data frame with one column
factors <- as.data.frame(</pre>
    matrix(data = c(rep("crown", length(crown)), rep("stem", length(stem))),
   ncol = 1, dimnames = list(c(crown, stem), "Group")))
## We then can use the customised series function to create the two series
## i.e. the two regions of the ordinated space
crown_stem <- cust.series(BeckLee_mat50, factors)</pre>
## This created a dispRity object containing two series: crown and stem
class(crown_stem)
## [1] "dispRity"
crown_stem
## 2 custom series for 50 elements
## Series:
## Group.crown, Group.stem.
## This object contains three elements
names(crown_stem)
## [1] "data"
                 "elements" "series"
## With "data" being the list sub-matrices
str(crown_stem$data)
## List of 2
## $ Group.crown: num [1:30, 1:48] 0.3079 0.6531 0.5089 -0.1652 -0.0419 ...
   ..- attr(*, "dimnames")=List of 2
   ....$ : chr [1:30] "Dasypodidae" "Bradypus" "Myrmecophagidae" "Todralestes" ...
##
    ....$ : NULL
##
## $ Group.stem : num [1:20, 1:48] -0.739 -0.68 -0.511 -0.477 -0.473 ...
    ..- attr(*, "dimnames")=List of 2
     ....$ : chr [1:20] "Daulestes" "Bulaklestes" "Uchkudukodon" "Kennalestes" ...
##
     ....$ : NULL
## "elements" being the list of taxa in the original ordinated matrix
str(crown_stem$elements)
## chr [1:50] "Cimolestes" "Maelestes" "Batodon" "Bulaklestes" "Daulestes" ...
```

```
## and finally, "series" containing information on the series type (custom)
## and names (crown and stem)
crown_stem$series
## [1] "custom" "Group.crown" "Group.stem"
```

3.2 time.series

Another way to split the ordinated space (maybe more relevant to palaeobiologists) is to do it according to time. The time.series function allows to create series that contain all the elements present at specific points or during specific periods in time. This functions needs as input an ordinated space and a matching phylogenetic tree. Two types of time series can be performed by using the method option:

- discrete time series (or time-binning) using method = "discrete";
- 2. continuous time series (or time-slicing) using method = "continuous".

For the time-slicing method details see Guillerme and Cooper (in prep.). For both methods, the function intakes the time argument which can be a vector of numeric values for:

- 1. defining the boundaries of the time bins (when method = "discrete");
- 2. defining the time slices (when using method = "continuous").

Otherwise, the time argument can be set as a single numeric value for automatically generating a given number of equidistant time-bins/slices. Additionally, it is also possible to input a data frame containing the First/Last Occurrence Data (FAD/LAD) for taxa that span over a longer time than the tips/nodes age. Here is an example for method = "discrete":

```
## Generating three time bins containing the data present every 40 Ma
time_bins <- time.series(data = BeckLee_mat50, tree = BeckLee_tree,
   method = "discrete", time = c(120, 80, 40, 0))
## No FAD/LAD table has been provided.
## Every tips are assumed to be single points in time.
## Note that the function provides a warning saying that tips where single
## points in time (no FAD/LAD information). We can fix that by adding the
## age data for the taxa that have some longer occurrence spans.
time_bins <- time.series(data = BeckLee_mat50, tree = BeckLee_tree,</pre>
   method = "discrete", time = c(120, 80, 40, 0), FADLAD = BeckLee_ages)
## Some tips have no FAD/LAD and are assumed to be single points in time.
## To entirely avoid the warning we could collect the occurrence span data
## for all the taxa but that's not necessary. The function automatically
## assumes no occurrence span time (i.e. single points in time) for all taxa
## by default.
time_bins
## 3 discrete series for 50 elements
## Series:
## 120-80, 80-40, 40-0.
```

This generated indeed a list of 3 sub matrices. Note that we can also generate equivalent results by just telling the function that we want three time-bins (series) as follow:

We now have three time bins of 44.50368 million years each.

When using this method, the oldest boundary of the first bin (or the first slice, see below) is automatically generated as being the root age + 1% of the tree length as long as at least three elements are present at that point in time. The algorithm adds an extra 1% tree length until reaching the required minimum of three elements. It is also possible to include nodes in each bin by using inc.nodes = TRUE and providing a matrix that contains the ordinated distance between tips AND nodes.

For the time-slicing method (method = "continuous"), the idea is really similar. This option intakes a matrix that contains the ordinated distance between taxa *AND* nodes and an assumed evolutionary model via the model argument:

- 1. "acctran" where the data chosen on each time slice is always the one of the offspring
- 2. "deltran" where the data chosen on each time slice is always the one of the descendant
- 3. "punctuated" where the data chosen on each time slice is randomly chosen between the offspring or the descendant
- 4. "gradual" where the data chosen on each time slice is either the offspring or the descendant depending on branch length

```
## Generating four time slices every 40 million years assuming a gradual
## evolution model
time_slices <- time.series(data = BeckLee_mat99, tree = BeckLee_tree,</pre>
   method = "continuous", model = "gradual", time = c(120, 80, 40, 0),
   FADLAD = BeckLee_ages)
## Some tips have no FAD/LAD and are assumed to be single points in time.
time_slices
## 4 continuous series for 99 elements
## Series:
## 120, 80, 40, 0.
## Note that in the same way as for the discrete method, we can also
## automatically generate the slices
time.series(data = BeckLee_mat99, tree = BeckLee_tree, method = "continuous",
   model = "gradual", time = 4)
## No FAD/LAD table has been provided.
## Every tips are assumed to be single points in time.
## 4 continuous series for 99 elements
## Series:
## 133.51104, 89.00736, 44.50368, 0.
```

3.3 boot.matrix

Once we obtain our different series, we might want to bootstrap and rarefy it (i.e. pseudo-replicating the data). The bootstrap will allow us to make each subsample more robust to outliers and the rarefaction will allow us to compare slices with the same number of elements to get rid of eventual sampling problems. The boot.matrix allows to bootstraps and rarefy ordinated matrices in a fast and easy way. The default options will bootstrap the matrix 1000 times without rarefaction. The number of bootstrap pseudo-replicates can be defined using the bootstraps option (see below).

```
boot.matrix(data = BeckLee_mat50)

## Bootstrapped ordinated matrix with 50 elements
## 1

## Data was bootstrapped 1000 times, using the full bootstrap method.

## As we can see, the output is also a dispRity object that is summarised
## automatically, and gives information on the data as well as the number of
## bootstraps and the bootstraps methods.
```

Additionally, this function allows to control the bootstrap algorithm through the boot.type argument. Currently two algorithms are implemented:

- 1. "full" where the bootstrapping is entirely stochastic (all the data is bootstrapped)
- 2. "single" where only one random elements is replaced by one other random elements each pseudo-replication

This function also allows to rarefy the data using the rarefaction argument. The default argument is FALSE but it can be set to TRUE to fully rarefy the data (i.e. remove n elements for the number of pseudoreplicates, where n varies from the maximum number of elements present in the dataset to a minimum of 3 elements). It can also be set to a fix numeric value (or a set of numeric values). Finally, one last argument, rm.last.axis allows to remove a certain amount of dimensions (or axis) for the ordinated space. This can be logical argument where FALSE (default) will not remove any dimension and TRUE will remove the last dimensions that bear up to 5% of the total ordinated space's variance.

```
## Bootstrapping with the single bootstrap method
boot.matrix(BeckLee_mat50, boot.type = "single")

## Bootstrapped ordinated matrix with 50 elements
## 1

## Data was bootstrapped 1000 times, using the single bootstrap method.

## Bootstrapping with the full rarefaction
boot.matrix(BeckLee_mat50, bootstraps = 20, rarefaction = TRUE)

## Bootstrapped ordinated matrix with 50 elements
## 1

## Data was bootstrapped 20 times, using the full bootstrap method.

## Data was fully rarefied (down to 3 elements).

## Or with a set number of rarefaction levels
boot.matrix(BeckLee_mat50, bootstraps = 20, rarefaction = c(6:8,3))

## Bootstrapped ordinated matrix with 50 elements
## 1

## Data was bootstrapped 20 times, using the full bootstrap method.

## Data was bootstrapped 20 times, using the full bootstrap method.

## Data was rarefied with a maximum of 6, 7, 8 and 3 elements
```

```
## And removing the last axis (default)
boot.matrix(BeckLee_mat50, rm.last.axis = TRUE)

## Bootstrapped ordinated matrix with 50 elements
## 1
## Data was bootstrapped 1000 times, using the full bootstrap method.
## The 6 last axis were removed from the original ordinated data.

## Or with a fix value (50%)
boot.matrix(BeckLee_mat50, rm.last.axis = 0.5)

## Bootstrapped ordinated matrix with 50 elements
## 1
## Data was bootstrapped 1000 times, using the full bootstrap method.
## The 35 last axis were removed from the original ordinated data.
```

Of course, one could be interested in directly supplying the sub-sampled matrices generated above directly to this function. In fact, it can also deal with a list of matrices or with a dispRity object output from the cust.series or time.series functions.

```
## Bootstrap and full rarefaction on the crown/stem series
crown_stemBS <- boot.matrix(crown_stem, bootstraps = 100, rarefaction = TRUE)</pre>
## Bootstrap on the time binning/slicing series
time_binsBS <- boot.matrix(time_bins, bootstraps = 100)</pre>
time_slicesBS <- boot.matrix(time_slices, bootstraps = 100)</pre>
## Note that all these objects are of class dispRity
crown_stemBS
## Bootstrapped ordinated matrix with 50 elements
## Series:
## Group.crown, Group.stem.
## Data was split using custom method.
## Data was bootstrapped 100 times, using the full bootstrap method.
## Data was fully rarefied (down to 3 elements).
time_binsBS
## Bootstrapped ordinated matrix with 50 elements
## Series:
## 120-80, 80-40, 40-0.
## Data was split using discrete method.
## Data was bootstrapped 100 times, using the full bootstrap method.
time_slicesBS
## Bootstrapped ordinated matrix with 99 elements
## Series:
## 120, 80, 40, 0.
## Data was split using continuous method.
## Data was bootstrapped 100 times, using the full bootstrap method.
```

3.4 dispRity

This function is a modular function that allow to simply (and quickly!) calculate disparity from a matrix.

Because disparity can be measured in many ways, this function is a tool to measure disparity *as defined by the user*. In fact, the dispRity function intakes two main arguments: the data and the disparity metric. The disparity metric is a function or a set of functions that summarises the ordinated matrix to a single value that represents, in our example, the diversity of morphologies.

The dispRity algorithm decomposes the metrics functions into three levels that correspond to the dimensions of the output of each metric function. For more details on this algorithm please refer to the metric vignette.

In practice, the dispRity function intakes one or more functions as a definition of disparity. Several of these functions will be already implemented in other packages (such as stats::median, base::sum, etc.); some others are implemented in this package (listed in ?dispRity.metric) and finally some others will be defined by the users. The make.metric function is designed to help users create and test their own disparity metric functions. In practice, the use of these metrics in the dispRity function, is pretty easy:

```
dispRity(BeckLee_mat50, metric = mean)
## Disparity measurements across 1 series for 50 elements
## Disparity calculated as: mean for 48 dimensions.
## This defines disparity as the mean value of the morphospace (using
## mean::base).
## It is also possible to combine multiple functions:
dispRity(BeckLee_mat50, metric = c(sum, variances))
## Disparity measurements across 1 series for 50 elements
## Disparity calculated as: c(sum, variances) for 48 dimensions.
## Defining disparity as the sum (sum::base) of the variances
## (variances::dispRity) of each dimension of the morphospace.
dispRity(BeckLee_mat50, metric = c(prod, centroids))
## Disparity measurements across 1 series for 50 elements
## Disparity calculated as: c(prod, centroids) for 48 dimensions.
## For the product (prod::base) of the distances between each elements and the
## centroid of the morphospace (centroids::dispRity).
## Or user defined ones:
total.range <- function(X) abs(range(X)[1]-range(X)[2])</pre>
dispRity(BeckLee_mat50, metric = c(total.range, centroids))
## Disparity measurements across 1 series for 50 elements
## Disparity calculated as: c(total.range, centroids) for 48 dimensions.
## For the range (total.range, user defined) of the distances between each
## elements and the centroid of the morphospace (centroids::dispRity).
## Or more complex ones:
vars1 <- dispRity(BeckLee_mat50, metric = c(sd, variances, var))</pre>
## For the standard deviation (sd::stats) of the variance (variances::dispRity)
## of each column of the variance/covariance matrix (var::stats).
```

```
## Note that the order of the function is not important since the levels of
## each function are automatically detected by the dispRity algorithm.
vars2 <- dispRity(BeckLee_mat50, metric = c(variances, var, sd))
all(summary(vars) == summary(vars2))
## Error in summary(vars): object 'vars' not found</pre>
```

Note that these functions do not directly output the disparity values but only the summary of the dispRity objects. To display the results, see the section 3.5.

In these examples we used only simple ordinated matrix but of course, it might be more interesting to directly use the dispRity objects we generated in the steps above. Thus we can calculate the bootstrapped and rarefied disparity in each sub-sample of the morphospace. In this example we are going to define disparity as being the sum of the variances in each dimension)=.

```
## Disparity in crown and stem mammals:
disp_crown_stemBS <- dispRity(crown_stemBS, metric = c(sum, variances))

## Disparity through time:
disp_time_binsBS <- dispRity(time_binsBS, metric = c(median, centroids))
disp_time_slicesBS <- dispRity(time_slicesBS, metric = c(median, centroids))</pre>
```

Note that the computational time was longer for the rarefied data (crown_stemBS) since it had to calculate the disparity on all the rarefied matrix, each bootstrapped 100 times. A faster parallel version will be implemented in future releases (see section ??).

3.5 summary

As shown in these examples above however, even though the disparity has been calculated in many way, the function doesn't directly outputs the results, rather just a summary of the analysis.

This function intakes a dispRity object plus various options namely the quantile values for the confidence intervals levels; the cent.tend for the central tendency to use for summarising the results and two *visual* options which are whether to recall the dispRity options and how much digits are wanted in the results.

```
## Let's have a look at the disparity-through-time using in the time-binned data
summary(disp_time_binsBS)
##
    series n observed mean 2.5%
                                     25%
                                           75% 97.5%
                 1.203 1.103 0.895 1.057 1.178 1.236
## 1 120-80 8
## 2 80-40 27
                 1.344 1.322 1.275 1.306 1.339 1.366
## 3
      40-0 16
               1.353 1.323 1.265 1.304 1.346 1.368
## The 50 and 95 quantiles are calculated and the central tendency is the mean by default.
## Note that the function also displays the observed disparity (non-bootstrapped).
## We can change that easily by specifying different values in the options
summary(disp_time_binsBS, quantile = 88, cent.tend = sd, rounding = 1)
    series n observed sd 6% 94%
## 1 120-80 8
                  1.2 0.1 1.0 1.2
## 2 80-40 27
                   1.3 0.0 1.3 1.4
## 3
     40-0 16
                  1.4 0.0 1.3 1.4
## Also, if we happen to have forgotten what was in the disp_time_binsBS object (as in
## which options where used to modify the original matrix) we can use the recall option:
summary(disp_time_binsBS, recall = TRUE)
```

```
## Disparity calculated as: c(median, centroids) for 48 dimensions.
## Data was split using discrete method.
## Data was bootstrapped 100 times, using the full bootstrap method.
  series n observed mean 2.5% 25% 75% 97.5%
## 1 120-80 8 1.203 1.103 0.895 1.057 1.178 1.236
## 2 80-40 27
                 1.344 1.322 1.275 1.306 1.339 1.366
      40-0 16
                 1.353 1.323 1.265 1.304 1.346 1.368
## Note that the recall just prints text. The output of summary.dispRity is always a
## data.frame
summary_table <- summary(disp_time_binsBS, recall = TRUE)</pre>
## Disparity calculated as: c(median, centroids) for 48 dimensions.
## Data was split using discrete method.
## Data was bootstrapped 100 times, using the full bootstrap method.
class(summary_table)
## [1] "data.frame"
## Finally we can see the results of the rarefaction analysis for the crown/stem data
head(summary(disp_crown_stemBS))
##
          series n observed mean 2.5%
                                          25%
                                               75% 97.5%
## 1 Group.crown 3
                       NA 1.894 1.245 1.863 2.058 2.190
## 2 Group.crown 4
                        NA 1.932 1.591 1.874 2.034 2.145
## 3 Group.crown 5
                        NA 1.919 1.634 1.816 2.029 2.087
## 4 Group.crown 6
                       NA 1.928 1.616 1.862 2.014 2.101
## 5 Group.crown 7
                        NA 1.937 1.753 1.865 1.998 2.057
## 6 Group.crown 8
                        NA 1.927 1.769 1.865 1.991 2.043
## This outputs a longer table with all the variations of the number of crown/stem taxa
## from the maximum (30 and 20) to the minimum (3). Note that the observed disparity
## is NA for rarefied data.
```

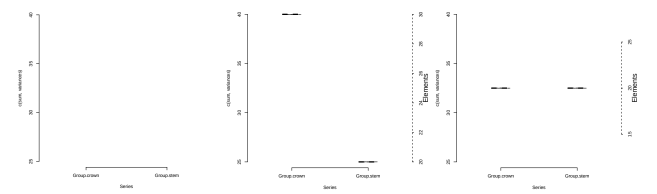
3.6 plot

This final function allows to visualise the disparity results in a often nicer fashion than just a table (even though the exact same data is displayed). The plot.dispRity option intakes the same options as summary.dispRity along side with various graphical options described in the function manual. Here, let's just have a look a few of these options.

Let's start with plotting the difference in disparity between the crown and the stem mammals. For this we are going to simply specify the type of plots using the type option that allows to choose between the "continuous" or "discrete" method. In our case, we are interested in looking at the results in a discrete way. Note that the plot type can be left missing. The function will then use "discrete" as default or "continuous" if the input data came from the time.series function with method = "continuous".

The grey squares represent the confidence intervals (50 in dark grey and 90 in light grey) and the dot represents the mean. However, it might be interesting to see how many taxa are present in each group. This can be done via the elements option. A third interesting option is to plot the rarefied data if a rarefaction was applied on the data with the boot.matrix function. For displaying the rarefied data we can use the rarefaction option. This option can be either FALSE (default) for always plotting the maximum number of taxa per series or TRUE to always plot the minimum. Additionally, it can be a specific number to plot a specific number of taxa. Let say we want both groups to have 20 taxa maximum. We can see now that the confidence intervals overlap more than previously

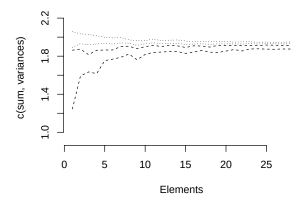
```
## Graphical options
quartz(width = 15, height = 5); par(mfrow = (c(1,3)), bty = "n")
## Plotting the score for both groups
plot(disp_crown_stemBS, ylim = c(25,40))
## Same one with the number of taxa for each data set
plot(disp_crown_stemBS, elements = TRUE, ylim = c(25,40))
## Same one with rarefied data (note how that affects the elements data as well!)
plot(disp_crown_stemBS, elements = TRUE, rarefaction = 20, ylim = c(25,40))
```



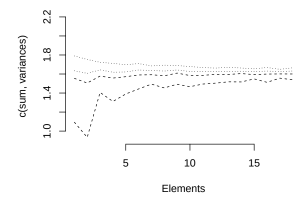
To understand how the number of taxa affects each series, we can use the rarefaction = "plot" function to plot the rarefaction curves.

```
## Graphical options
par(bty = "n")
## The rarefaction curves
plot(disp_crown_stemBS, elements = TRUE, rarefaction = "plot")
## Note that the function automatically draws the curves for each series and splits
## the plot screen accordingly.
```

Group.crown

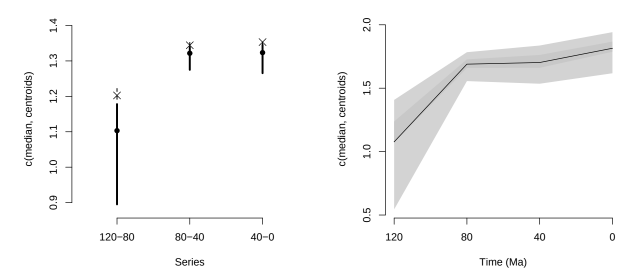


Group.stem



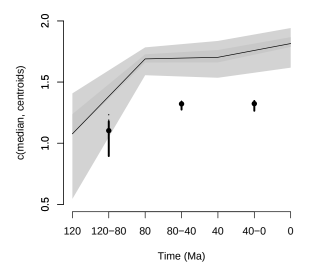
The different dashed lines represent the different confidence intervals around each rarefaction curve. Additionally, for the type = "discrete" option, we might also decide to plot many boxplots (which can be a bit messy). The discrete_type option allows to switch between "box" or "line" for showing the results in one or the other format. We can also plot the observed data when existing (i.e. not with some rarefaction levels) by using observed = TRUE. Let's look at that with the time binned data (even though there is only three time bins). Finally, we can use the type = "continuous" option that comes in handy for plotting continuous data like in the time sliced analysis (obviously...).

```
## Graphical options
quartz(width = 10, height = 5); par(mfrow = (c(1,2)), bty = "n")
## The disparity-through-time data in a time-binned way (with lines rather than
## boxes) with the observed data as crosses
plot(disp_time_binsBS, discrete_type = "line", observed = TRUE)
## The disparity-through-time data in a time-sliced ways
plot(disp_time_slicesBS)
```



We can even plot both results (discrete and continuous) on the same graph:

```
## Graphical options
par(bty = "n")
## First, plotting the continuous data
plot(disp_time_slicesBS, ylim=c(0.5,2))
## Second, adding the discrete data
par(new = TRUE)
plot(disp_time_binsBS, discrete_type = "line", ylim=c(0.5,2), xlab="", ylab="")
```

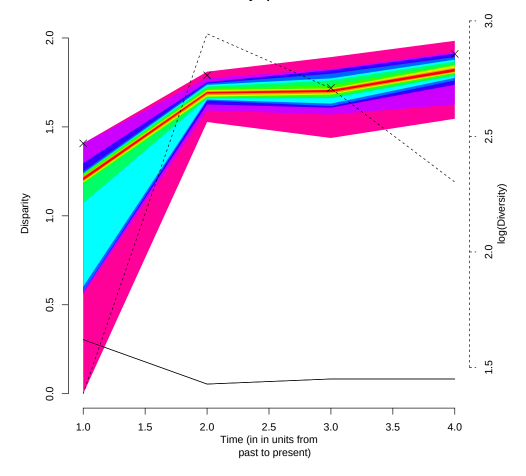


Note that the results vary in values but not in pattern. The change in values might be due to the fact that nodes data are not included in the discrete analysis.

I encourage you to play with the graphical options to have some prettier results. Note that most of the options from plot can be passed to plot.dispRity via

```
## Graphical options
par(bty = "n")
## A plot with many options!
plot(disp_time_slicesBS, quantile = c(seq(from=10, to=100, by=10)), cent.tend = sd,
    type = "continuous", elements = "log", col = c("black", rainbow(10)),
    ylab = c("Disparity", "log(Diversity)"), xlab = "Time (in in units from
    past to present)", time.series = FALSE, observed = TRUE, main = "Many options...")
```

Many options...



3.7 test.dispRity

3.8 utilities

tree.age

This function allows to calculate the age of each individual nodes and tips in a tree. It can either use the root age of the tree (if present as \$root.time) or else calculate the age using a user defined root age via the age argument. Also, it is possible to decide whether the time is calculated towards the past (e.g. million years ago) or towards the present (e.g. in time since the origin).

```
## This tree has a root age
BeckLee_tree$root.time
## [1] 139.074
```

```
## So we can get the age of each tips and nodes directly
head(tree.age(BeckLee_tree), 5)
##
              elements
      ages
           Daulestes
## 1 90.00
## 2 90.00 Bulaklestes
## 3 90.00 Uchkudukodon
## 4 77.85 Kennalestes
## 5 77.85
           Asioryctes
## But we can also decide to make the age relative (between 1 and 0)
head(tree.age(BeckLee_tree, age = 1), 5)
##
      ages
              elements
## 1 0.647
             Daulestes
## 2 0.647 Bulaklestes
## 3 0.647 Uchkudukodon
## 4 0.560 Kennalestes
## 5 0.560
           Asioryctes
## Or even relative, but from the root (i.e. how far are the nodes/tips
## from the root)
head(tree.age(BeckLee_tree, age = 1, order = "present"), 5)
                  elements
          ages
## 1 0.3528637
                 Daulestes
## 2 0.3528637 Bulaklestes
## 3 0.3528637 Uchkudukodon
## 4 0.4402271 Kennalestes
## 5 0.4402271 Asioryctes
```

4 Developments

As stated at the start of the demo, this version 0.1.2 is still in development and many parts are missing. Here are the new functionalities that will be implemented before the proper release version (v.1).

4.1 More user defined stuff

I intend also develop functions to help users to develop their own algorithms for the bootstrap method (via make.boot) or the evolutionary models (via make.model). Both functions will provide similar testing as the make.metric function.

4.2 Faster!

Finally, for long analysis, I intend to develop a parallel running version of the package. In fact, most of the internal functions are base on lapply functions that can be easily passed to snow::parLapply or similar parallel functions.

References

Beck, R. M. and M. S. Lee. 2014. Ancient dates or accelerated rates? Morphological clocks and the antiquity of placental mammals. Proceedings of the Royal Society B: Biological Sciences 281:1–10.

Guillerme, T. and N. Cooper. in prep. Mammalian morphological diversity does not increase in response to the cretaceous-paleogene mass extinction and the extinction of the (non-avian) dinosaurs. .