Package: BinMat (via r-universe)

August 30, 2024

```
Type Package
Title Processes Binary Data Obtained from Fragment Analysis (Such as
     AFLPs, ISSRs, and RFLPs)
Version 0.1.5
Description A molecular genetics tool that processes binary data from
     fragment analysis. It consolidates replicate sample pairs,
     outputs summary statistics, and produces hierarchical
     clustering trees and nMDS plots. This package was developed
     from the publication available here:
     <https://www.sciencedirect.com/science/article/pii/S1049964420306538>.
     The GUI version of this package is available on the R Shiny
     online server at:
     <https://clarkevansteenderen.shinyapps.io/BINMAT/> or it is
     accessible via GitHub by typing: shiny::runGitHub(``BinMat",
     ``clarkevansteenderen") into the console in R. Two real-world
     datasets accompany the package: an AFLP dataset of Bunias
     orientalis samples from Tewes et. al. (2017)
     <https:
     //besjournals.onlinelibrary.wiley.com/doi/full/10.1111/1365-2745.12869>,
     and an ISSR dataset of Nymphaea specimens from Reid et. al.
     (2021)
     <https://www.sciencedirect.com/science/article/pii/S0304377021000218>
     . The authors of these publications are thanked for allowing
     the use of their data.
License GPL-3
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RoxygenNote 7.1.2
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     3.1.4)
Suggests knitr, rmarkdown
```

13

VignetteBuilder knitr

NeedsCompilation no

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Contents

BinMatInput_ordinat	ion							 								
BinMatInput_reps .								 								
bunias_orientalis .								 								
check.data								 								
consolidate								 								
errors								 								
group.names								 								
nmds								 								
nymphaea								 								
peak.remove																
peaks.consolidated								 								
peaks.original								 								. 1
scree								 								. 1
shepard								 								. 1
upgma								 								. 1

BinMatInput_ordination

Example input data containing a consolidated binary matrix with groups

Description

Example input data containing a consolidated binary matrix with groups

Usage

Index

data(BinMatInput_ordination)

BinMatInput_reps 3

Format

A dataframe with columns for loci, and rows of replicate pairs. Grouping information is in the second column.

Examples

```
data(BinMatInput_ordination)
mat = BinMatInput_ordination
group.names(mat)
scree(mat)
shepard(mat)
clrs = c("red", "green", "black")
nmds(mat, colours = clrs, labs = TRUE)
```

BinMatInput_reps

Example input data containing a binary matrix comprising replicate pairs

Description

Example input data containing a binary matrix comprising replicate pairs

Usage

```
data(BinMatInput_reps)
```

Format

A dataframe with columns for loci, and rows of replicate pairs.

```
data(BinMatInput_reps)
mat = BinMatInput_reps
check.data(mat)
cons = consolidate(mat)
pks = peaks.consolidated(cons)
err = errors(cons)
rem = peak.remove(cons, 4)
clust = upgma(cons)
```

4 check.data

bunias_orientalis

Example input file of Bunias orientalis AFLP data, taken from Tewes et. al. (2017). This dataset has already been consolidated, and can be used as input for the generation of an nMDS plot. The paper can be found here: https://besjournals.onlinelibrary.wiley.com/doi/full/10.1111/1365-2745.12869

Description

Example input file of Bunias orientalis AFLP data, taken from Tewes et. al. (2017). This dataset has already been consolidated, and can be used as input for the generation of an nMDS plot. The paper can be found here: https://besjournals.onlinelibrary.wiley.com/doi/full/10.1111/1365-2745.12869

Usage

```
data(bunias_orientalis)
```

Format

A dataframe with columns for loci, and rows of replicate pairs. Grouping information is in the second column.

check.data

Checks binary matrix for unwanted characters.

Description

Checks for unwanted values (other than 1, 0, and ?).

Usage

check.data(x)

Arguments

Χ

A CSV file containing replicate pairs of binary data.

Value

Index positions where unwanted values occur (row, column).

consolidate 5

Examples

```
data(BinMatInput_reps)
mat = BinMatInput_reps
check.data(mat)
```

consolidate

Consolidates replicate pairs in a binary matrix.

Description

Reads in a binary matrix comprising replicate pairs and consolidates each pair into a consensus read. For each replicate pair at each locus, $1 \& 1 \rightarrow 1$ (shared presence), $0 \& 0 \rightarrow 0$ (shared absence), $0 \& 1 \rightarrow 2$ (ambiguity).

Usage

```
consolidate(x)
```

Arguments

Χ

A CSV file containing replicate pairs of binary data. See the example input file "BinMatInput_reps".

Value

Consolidated binary matrix.

Examples

```
data(BinMatInput_reps)
mat = BinMatInput_reps
cons = consolidate(mat)
```

errors

Calculates Jaccard and Euclidean error rates.

Description

Calculates the Jaccard and Euclidean error rates for the dataset. Jaccard's error does not take shared absences of bands as being biologically meaningful. JE = (f10 + f01)/(f10 + f01 + f11) and EE = (f10 + f01)/(f10 + f01 + f11 + f00). At each locus, f01 and f10 indicates a case where a 0 was present in one replicate, and a 1 in the other. f11 indicates the shared presence of a band in both replicates, and f00 indicates a shared absence. For example, if a replicate pair comprises Rep1 = 00101 and Rep2 = 01100, JE = (1+1)/(1+1+1) = 2/3 = 0.67, EE = (1+1)/(1+1+1+2) = 2/5 = 0.4.

6 group.names

Usage

```
errors(x)
```

Arguments

Χ

Consolidated binary matrix.

Value

JE (Jaccard Error), EE (Euclidean Error), and standard deviations.

Examples

```
data(BinMatInput_reps)
mat = BinMatInput_reps
cons = consolidate(mat)
errors(cons)
```

group.names

Outputs group names specified in the input file for the creation of an nMDS plot.

Description

Returns group names in the uploaded consolidated binary data. This will help in knowing which colours are assigned to which group name.

Usage

```
group.names(x)
```

Arguments

х

Consolidated binary matrix with grouping information in column 2.

Value

Scree plot.

```
mat = BinMatInput_ordination
group.names(mat)
```

nmds 7

nmds

Creates a non-metric multidimensional scaling plot (nMDS).

Description

Creates an nMDS plot from a consolidated binary matrix with grouping information. Colours and shapes of plotted points need to be specified. For example, if there are two groups, then: clrs = c("red", "blue"), sh = c(16, 16). This assigns red to the first group name, and blue to the second. Both will have a pch shape of 16 (round dot). These two vectors are then passed to the function nmds() as: colours = clrs, shapes = sh.

Usage

```
nmds(
    x,
    dist_meth = "binary",
    k_val = 2,
    pt_size = 1,
    colours = c("dodgerblue", "black", "red", "green3", "orange", "darkblue", "gold2",
    "darkgreen", "darkred", "grey", "darkgrey", "magenta", "darkorchid", "purple",
    "brown", "coral3", "turquoise", "deeppink", "lawngreen", "deepskyblue", "tomato",
    "yellow", "yellowgreen", "royalblue", "olivedrab", "midnightblue", "indianred1",
    "darkturquoise"),
    labs = FALSE,
    legend_pos = "right",
    include_ellipse = FALSE,
    ellipse_type = "norm",
    dimension1 = 1,
    dimension2 = 2
)
```

Arguments

X	Consolidated binary matrix with grouping information in the second column.
dist_meth	Distance method. Set to "binary" by default. Other options are "euclidean", "maximum", "manhattan", "canberra", or "minkowski".
k_val	Number of dimensions for the nMDS plot. Set to 2 by default.
pt_size	Point size for symbols on the plot. Set to 1 by default.
colours	Vector containing colours to be assigned to groups. This can be changed to the options available in the RColorBrewer palette set (e.g. "Set1"). See http://applied-r.com/rcolorbrewer-palettes/ for more palette options. Alternatively, the colours can be set manually using, for example, c("red", "green", "blue"), thereby setting a colour for each group in your dataset. There are 28 default colours that will be set automatically to your groups.
labs	Indicate whether labels should appear on the graph or not (TRUE or FALSE). Default = FALSE.

8 nymphaea

legend_pos	Indicate the position of the legend. Default = "right", but other options are "left", "bottom", "top", or "none"
include_ellipse	
	Indicate whether ellipses should be included around groups. Default = FALSE.
ellipse_type	Select the type of ellipses to include around groups. Options are "convex", "confidence", "t", "norm", and "euclid". See the ggpubr::ggscatter() function documentation for more details.
dimension1	Indicate the first dimension to plot $(1, 2, \text{ or } 3)$ for the x axis. If $k = 2$, the first two dimensions will automatically be plotted. If $k = 3$, select between the three.
dimension2	Indicate the second dimension to plot (1, 2, or 3) for the y axis

Value

nMDS plot.

Examples

```
mat = BinMatInput_ordination
group.names(mat)
clrs = c("red", "green", "black")
nmds(mat, colours = clrs, labs = TRUE, include_ellipse = TRUE)
```

nymphaea	Example input file of Nymphaea ISSR data, taken from
	Reid et. al. (2021). This dataset has already been
	consolidated, and can be used as input for the genera-
	tion of an nMDS plot. The paper can be found here:
	<https: article="" pii="" s0304377021000218="" science="" www.sciencedirect.com=""></https:>

Description

Example input file of Nymphaea ISSR data, taken from Reid et. al. (2021). This dataset has already been consolidated, and can be used as input for the generation of an nMDS plot. The paper can be found here: https://www.sciencedirect.com/science/article/pii/S0304377021000218

Usage

data(nymphaea)

Format

A dataframe with columns for loci, and rows of replicate pairs. Grouping information is in the second column

peak.remove 9

peak.remove	Removes samples with peaks equal to or less than a specified threshold value.
peak.remove	

Description

Removes samples with a peak number less than a specified value.

Usage

```
peak.remove(x, thresh)
```

Arguments

x Binary matrix - consolidated or original.thresh Peak threshold value for removal.

Value

Filtered dataset, and either the row name/s or row number/s of samples that were removed.

Examples

```
mat = BinMatInput_ordination
new = peak.remove(mat, 4)
```

peaks.consolidated

Calculates peak numbers for a consolidated data set (total, maximum, and minimum).

Description

Returns total, maximum, and minimum number of peaks in the binary matrix.

Usage

```
peaks.consolidated(x)
```

Arguments

x Binary matrix comprising replicate pairs.

Value

Peak information.

10 scree

Examples

```
data(BinMatInput_reps)
mat = BinMatInput_reps
cons = consolidate(mat)
peaks.consolidated(cons)
```

peaks.original

Calculates peak numbers for the data set with all replicates (total, maximum, and minimum).

Description

Returns total, maximum, and minimum number of peaks in the binary matrix.

Usage

```
peaks.original(x)
```

Arguments

Х

Binary matrix comprising replicate pairs.

Value

Peak information.

Examples

```
data(BinMatInput_reps)
mat = BinMatInput_reps
peaks.original(mat)
```

scree

Draws a scree plot.

Description

Creates a scree plot for the nMDS. This indicates the optimum number of dimensions to use to minimise the stress value. The stress value is indicated by a red dotted line at 0.15. Values equal to or below this are considered acceptable.

Usage

```
scree(x, dimensions = 4, dist_meth = "binary")
```

shepard 11

Arguments

x Consolidated binary matrix with grouping information in column 2.

dimensions Number of dimensions to plot. Set to 4 by default.

dist_meth Distance method. Set to "binary" by default. Other options are "euclidean",

"maximum", "manhattan", "canberra", or "minkowski".

Value

Scree plot.

Examples

```
mat = BinMatInput_ordination
scree(mat)
```

shepard

Creates a shepard plot.

Description

Creates a Shepard plot for the nMDS. This indicates the 'goodness of fit' of the original distance matrix vs the ordination representation. A high R-squared value is favourable.

Usage

```
shepard(x, k_val = 2, dist_meth = "binary")
```

Arguments

x Consolidated binary matrix.

k_val Number of dimensions. Set to 2 by default.

dist_meth Distance method. Set to "binary" by default. Other options are "euclidean",

"maximum", "manhattan", "canberra", or "minkowski".

Value

Shepard plot.

```
mat = BinMatInput_ordination
shepard(mat)
```

12 upgma

upgma

Draws a hierarchical clustering tree (UPGMA).

Description

Creates a UPGMA hierarchical clustering tree, with a specified number of bootstrap repetitions.

Usage

```
upgma(
    x,
    bts = 10,
    size = 0.55,
    lab_size = 0.55,
    method = "binary",
    hclust = "average",
    fromFile = FALSE
)
```

Arguments

x	Consolidated binarx matrix.
bts	Bootstrap replications. Set to 10 by default.
size	Size of plot. Set to 0.55 by default.
lab_size	Size of label text. Set to 0.55 by default.
method	Distance method. Set to 'binary' (=Jaccard distance) by default.
hclust	Clustering method. Set to 'average' (=UPGMA) by default
fromFile	Indicates whether the binary data used by the function has been consolidated by BinMat, or whether it comes from the user's own file. Set to FALSE by default (in the assumption that the data has been consolidated by BinMat, and that that object is being passed to the function).

Value

UPGMA tree

```
data(BinMatInput_reps)
mat = BinMatInput_reps
cons = consolidate(mat)
clust = upgma(cons)
```

Index

```
* datasets
    BinMatInput_ordination, 2
    BinMatInput_reps, 3
    bunias_orientalis, 4
    nymphaea, 8
BinMatInput_ordination, 2
BinMatInput_reps, 3
bunias_orientalis, 4
check.data,4
consolidate, 5
errors, 5
{\tt group.names}, {\color{red} 6}
nmds, 7
nymphaea, 8
peak.remove, 9
peaks.consolidated, 9
peaks.original, 10
scree, 10
shepard, 11
upgma, 12
```