# Package 'DepLogo'

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<b>Description</b> Plots dependency logos from a set of aligned input sequences.
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Alphabet built alphabet

# Description

builts an object of class Alphabet from the given set of symbols and colors

# Usage

```
Alphabet(chars, cols)
```

# Arguments

chars set of symbols

cols set of colors; one for each symbol

# Value

the Alphabet object

# Author(s)

Martin Nettling

```
DNA <- Alphabet(c("A", "C", "G", "T"), c("green4", "blue", "orange", "red"))</pre>
```

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alphabet.dna

DNA alphabet

# Description

DNA alphabet

# Usage

alphabet.dna

# **Format**

An object of class list of length 2.

alphabet.dna.gap

DNA alphabet with gaps

# Description

DNA alphabet with gaps

# Usage

alphabet.dna.gap

# **Format**

An object of class list of length 2.

alphabet.protein

Amino acid alphabet

# Description

Amino acid alphabet

# Usage

alphabet.protein

## **Format**

An object of class list of length 2.

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alphabet.protein.gap Amino acid alphabet with gaps

# Description

Amino acid alphabet with gaps

# Usage

```
alphabet.protein.gap
```

# **Format**

An object of class list of length 2.

alphabet.rna

RNA alphabet

# Description

RNA alphabet

# Usage

alphabet.rna

# **Format**

An object of class list of length 2.

alphabet.rna.gap

RNA alphabet with gaps

# Description

RNA alphabet with gaps

# Usage

alphabet.rna.gap

## **Format**

An object of class list of length 2.

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colorchart

Plots a colorchart representation of a set of sequences

## **Description**

This function is a low-level plotting function (using image with add=TRUE, internally).

# Usage

```
colorchart(part, yoff, ic.scale = TRUE)
```

# **Arguments**

part the set of sequences as DLData object

yoff the offset in y-direction within the current plot

ic.scale ignored for colorcharts

## Value

the vertical (y) offset after this plot

## Author(s)

Jan Grau <grau@informatik.uni-halle.de>

```
# read data and create DLData object
seqs <- read.table(system.file("extdata", "cjun.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[,1], weights = log1p(seqs[, 2]) )

# create high-level plot
plot(NULL, xlim = c(1, ncol(data$data) - 1), ylim = c(0, nrow(data$data)),
    ylab = nrow(data$data), axes = FALSE)
# and add colorchart and axis
colorchart(data, yoff = nrow(data$data))
axis(1)</pre>
```

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Rectangles of averaged colors

# **Description**

Plots a representation of a set of sequences by rectangles of (scaled) averaged color values of the symbols at each position

## Usage

```
deprects(part, yoff, ic.scale = TRUE)
```

## **Arguments**

part the set of sequences as DLData object

yoff the offset in y-direction within the current plot

ic.scale if TRUE, alpha values of colors will be assigned based on "information content"

of the distribution at each position

#### **Details**

This function is a low-level plotting function (using rect, internally).

# Value

the vertical (y) offset after this plot

## Author(s)

Jan Grau <grau@informatik.uni-halle.de>

```
# read data and create DLData object
seqs <- read.table(system.file("extdata", "cjun.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[, 1],weights = log1p(seqs[, 2]) )

# create high-level plot
plot(NULL, xlim = c(1, ncol(data$data) - 1), ylim = c(0, nrow(data$data)),
    ylab = nrow(data$data), axes = FALSE)
# and add deprects and axis
deprects(data, yoff = nrow(data$data))
axis(1)</pre>
```

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DLData	Create DLData object
--------	----------------------

# Description

Creates a new DLData object from a set of input sequences.

# Usage

```
DLData(
    sequences,
    weights = NULL,
    symbols = alphabet.dna$alphabet,
    colors = alphabet.dna$colors,
    delim = "",
    sortByWeights = !is.null(weights),
    axis.labels = NULL
)
```

#### **Arguments**

sequences	the input sequences, may be provided as i) character vector or ii) a data. frame with sequences organized in rows and one symbol per column
weights	weights associated with the sequences, numeric vector of the same length as sequences has sequences
symbols	the symbols (alphabet) over which the sequences are defined
colors	colors for each of the symbols, not necessarily unique
delim	delimiter between the symbols in the input sequences, ignored if sequences as a $\mbox{data.frame}$
sortByWeights	if TRUE, sequences will be ordered by their weight in decreasing order
axis.labels	the labels of the individual sequence positions; if NULL, indexes from 1 to to total number of positions will be used

#### **Details**

Sequences may either be provided as a character vector or as a data. frame. All symbols occurring in these sequences need to be defined and assigned to colors, which are used for plotting later. Colors do not need to be unique, but symbols with identical colors may become indistinguishable in subsequent plots (which might even be desired, for instance, when visualizing protein properties instead of amino acids). Sequences may have an associated weight, which is used to order sequences, e.g., for creating chunks/blocks of sequences in subsequent plots (see chunks parameter of plotDeplogo).

## Value

the DLData object

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#### Author(s)

Jan Grau <grau@informatik.uni-halle.de>

#### See Also

plotDeplogo

## **Examples**

filter.by.conservation

Filters columns (sequence positions) by conservation

# Description

Filters columns based on the relative information content of each column which is the standard information content normalized to the interval [0,1], where 0 corresponds to uniform distribution and 1 to perfect conservation of one nucleotide or amino acid, respectively.

#### Usage

```
filter.by.conservation(relative.ic)
```

# **Arguments**

relative.ic the maximum relative information content allowed to retain a position

#### Value

function that, given a DLData object, returns TRUE for every column that does not exceed the specified relative information content

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## Author(s)

Jan Grau <grau@informatik.uni-halle.de>

# **Examples**

```
fun <- filter.by.conservation(relative.ic = 0.9)</pre>
```

filter.by.dependencies

Filters columns (sequence positions) by dependency

# **Description**

Filters columns based on the average or maximum mutual information of a column to all other columns. Mutual information is normalized to to interval [0,1], where 0 corresponds to independence and 1 to perfect dependence.

# Usage

```
filter.by.dependencies(mi.threshold, use.max = FALSE)
```

# **Arguments**

mi.threshold the minimum average or maximum mutual information required

use.max if TRUE, the maximum and otherwise the average mutual information will be

considered

# Value

function that, given a DLData object, returns TRUE for every column that does exceed the specified average mutual information

# Author(s)

Jan Grau <grau@informatik.uni-halle.de>

```
fun <- filter.by.dependencies(mi.threshold = 0.3)</pre>
```

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filter.by.gaps

Filters columns (sequence positions) by gaps

## Description

Filters columns (sequence positions) by gaps

## Usage

```
filter.by.gaps(percent.gap)
```

# Arguments

percent.gap the maximum fraction of gaps allowed to retain a column

# Value

function that, given a DLData object, returns TRUE for every column that does not exceed the specified number of gaps

## Author(s)

Jan Grau <grau@informatik.uni-halle.de>

# **Examples**

```
fun <- filter.by.gaps(percent.gap = 0.1)</pre>
```

filterColumns

Filters data columns by some filter function

# **Description**

Filters the columns of the input data, i.e., positions of input sequences, by a filter function that, given a DLData object, returns a list containing i) as element \$selected a vector with entries TRUE for every column that should be retained in the filtered data and ii) as element \$range the range of values obtained for the filtering criterion.

# Usage

```
filterColumns(data, filter.fun)
```

# Arguments

data the data as DLData object

filter.fun the filter function

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## Value

a DLData object containing the filtered columns and the indexes of the remaining in its axis.labels field

# Author(s)

Jan Grau <grau@informatik.uni-halle.de>

#### See Also

```
filter.by.gaps
filter.by.dependencies
filter.by.conservation
```

## **Examples**

getDeps

Compute dependencies between positions

# Description

Computes the dependencies (as measures by mutual information) between all positions (columns) of discrete data. Specifically, it returns for each pair of positions (i,j) the mutual information  $I(X_i,X_j)$  multiplied by the number N of sequences (rows), which may also be used for testing the statistical significance of mutual information values, as for large N,  $2*N*I(X_i,X_j)$  is approximately chi squared.

## Usage

```
getDeps(data, ...)
## S3 method for class 'DLData'
getDeps(data, ...)
## S3 method for class 'data.frame'
getDeps(data, alphabet, ...)
```

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# **Arguments**

data the data for computing mutual information. Either a DLData object or a data.frame;

In the latter case, the symbols of the alphabet must be provided as a second pa-

rameter

... the symbols of the alphabet as character vector, only if data is a data.frame

alphabet only required when called on a data.frame

# Value

a matrix of the mutual information values, where the diagonal is fixed to zero

# Author(s)

Jan Grau <grau@informatik.uni-halle.de>

## **Examples**

```
data <- DLData(c("ACGT", "ATTA"))
deps <- getDeps(data)</pre>
```

getPWM

Position weight matrix from DLData object

# **Description**

Determines the position weight matrix from a DLData object as relative frequency of symbols in each column of the data slot.

#### Usage

```
getPWM(part)
## S3 method for class 'DLData'
getPWM(part)
```

## **Arguments**

part the DLData object

## Value

the position weight matrix, where columns correspond to positions (columns of the DLData\$data slot) and rows to symbols

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## Author(s)

Jan Grau <grau@informatik.uni-halle.de>

## **Examples**

```
data <- DLData(c("ACGT", "ATTA"))
getPWM(data)</pre>
```

logo

Sequence logo

## **Description**

Plots a representation of a set of sequences as a sequence logo

# Usage

```
logo(part, yoff, ic.scale = TRUE)
```

# Arguments

part the set of sequences as DLData object

yoff the offset in y-direction within the current plot

ic.scale if TRUE, symbols are scaled by "information content" of the distribution at each

position

## **Details**

This function is a low-level plotting function (using polygon, internally).

## Value

the vertical (y) offset after this plot

# Author(s)

Jan Grau <grau@informatik.uni-halle.de>

```
# read data and create DLData object
seqs <- read.table(system.file("extdata", "cjun.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[, 1], weights = log1p(seqs[,2]) )

# create high-level plot
plot(NULL, xlim = c(1, ncol(data$data) - 1), ylim = c(0, nrow(data$data)),
    ylab = nrow(data$data), axes = FALSE)</pre>
```

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```
# and add sequence logo and axis
logo(data, yoff = nrow(data$data))
axis(1)
```

partition

Paritions data by most inter-dependent positions

## **Description**

Partitions data by the nucleotides at the most inter-dependent positions as measures by pairwise mutual information. Partitioning is performed recursively on the resulting subsets until i) the number of sequences in a partition is less then minElements, ii) the average pairwise dependency between the current position and numBestForSorting other positions with the largest mutual information value drops below threshold, or iii) maxNum recursive splits have already been performed. If splitting results in smaller partitions than minElements, these are added to the smallest partition with more than minElements sequences.

# Usage

```
partition(
  data,
  minElements = 10,
  threshold = 0.1,
  numBestForSorting = 3,
 maxNum = 6,
  sortByWeights = NULL,
  partition.by = NULL
)
## S3 method for class 'DLData'
partition(
  data,
 minElements = 10,
  threshold = 0.1,
  numBestForSorting = 3,
 maxNum = 6,
  sortByWeights = NULL,
  partition.by = NULL
)
```

#### **Arguments**

data the data as DLData object
minElements the minimum number of elements to perform a further split
threshold the threshold on the average mutual information value

numBestForSorting

the number of dependencies to other positions considered

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maxNum the maximum number of recursive splits

sortByWeights if TRUE, partitions are ordered by their average weight value, if false by fre-

quency of symbols at the partitioning position otherwise. If NULL, the \$sortByWeights

value of the DLData object is used

partition.by specify fixed positions to partition by

#### Value

the partitions as list of DLData objects

#### Author(s)

Jan Grau <grau@informatik.uni-halle.de>

## **Examples**

```
# create DLData object
seqs <- read.table(system.file("extdata", "cjun.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[, 1], weights = log1p(seqs[,2]) )

# partition data using default parameters
partitions <- partition(data)

# partition data using a threshold of 0.3 on the mutual
# information value to the most dependent position,
# sorting the resulting partitions by weight
partitions2 <- partition(data = data, threshold = 0.3, numBestForSorting = 1, sortByWeights = TRUE)</pre>
```

plotBlocks

Plots blocks of data

# Description

Plots the blocks of data in data by successive, vertically arranged sub-plots of the function provided as block. fun. If data is a single DLData object, one block is plotted. Further arguments are provided to block. fun.

# Usage

```
plotBlocks(
  data,
  show.number = TRUE,
  block.fun = deprects,
  ic.scale = TRUE,
  add = FALSE,
  ...
)
```

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```
## S3 method for class 'DLData'
plotBlocks(
  data,
  show.number = TRUE,
 block.fun = deprects,
  ic.scale = TRUE,
  add = FALSE,
)
## S3 method for class 'list'
plotBlocks(
  data,
  show.number = TRUE,
  block.fun = deprects,
  ic.scale = TRUE,
  add = FALSE,
)
```

## **Arguments**

data the data, a single DLData object or a list of DLData objects

show.number if true, the number of sequences (in total) in data is displayed on the left side

of the plot

block.fun the function called for each of the blocks

ic.scale if TRUE, output of block. fun may be scaled by "information content"

add if TRUE, the plot is added to an existing plot

... if add=FALSE forwarded to the internal call to plot

# Author(s)

Jan Grau <grau@informatik.uni-halle.de>

# See Also

deprects logo colorchart

```
# read data and create DLData object
seqs <- read.table(system.file("extdata", "cjun.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[, 1], weights = log1p(seqs[, 2]) )</pre>
```

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```
# plot all data
plotBlocks(data)

# partition data
partitions <- partition(data, threshold = 0.3)
# and plot partitions
plotBlocks(partitions)

# or plot partitions as sequence logos
plotBlocks(partitions, block.fun = logo)</pre>
```

plotDeparcs

Plots a graph representation of dependency values

## **Description**

Plots a representation of dependency values as arcs between the sequence positions. Internally, dependency values are computed using getDeps on the data object.

# Usage

```
plotDeparcs(
  data,
  axis.at.bottom = TRUE,
  add.legend = TRUE,
  show.pvals = FALSE,
  axis.labels = NULL,
  threshold = 0.1
)
```

## **Arguments**

the DLData object containing the data

axis.at.bottom if TRUE, the x-axis is shown at the bottom (side=1) of the plot, and at the top (side=3) otherwise

add.legend if TRUE a legend of the color scale is added to the plot

show.pvals if TRUE, -log10 p-values (computed by pchisq) are shown instead of mutual information values

axis.labels the labels of the x-axis

threshold threshold in mutual information values, edges below this value are not shown; ignored in show.pvals=TRUE

# Author(s)

Jan Grau <grau@informatik.uni-halle.de>

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## **Examples**

plotDeplogo

Plot a dependency logo

# **Description**

Plots a dependency logo

## Usage

```
plotDeplogo(
  data,
  dep.fun = plotDeparcs,
  block.fun = deprects,
  summary.fun = logo,
  weight.fun = NULL,
  chunks = NULL,
  chunk.height = 800,
  summary.height = 100,
  minPercent = 0.03,
  threshold = 0.1,
  numBestForSorting = 3,
  maxNum = 6,
  sortByWeights = NULL,
  dep.fun.legend = TRUE,
  show.dependency.pvals = FALSE,
  axis.labels = NULL,
  weight.ratio = 5,
  partition.by = NULL,
)
## S3 method for class 'DLData'
plotDeplogo(
  data,
  dep.fun = plotDeparcs,
```

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```
block.fun = deprects,
  summary.fun = logo,
 weight.fun = NULL,
  chunks = NULL,
  chunk.height = 800,
  summary.height = 100,
 minPercent = 0.03,
  threshold = 0.1,
  numBestForSorting = 3,
 maxNum = 6,
  sortByWeights = NULL,
  dep.fun.legend = TRUE,
  show.dependency.pvals = FALSE,
  axis.labels = NULL,
 weight.ratio = 5,
  partition.by = NULL,
)
```

# **Arguments**

data the data, currently implemented for DLData objects
dep.fun the function for plotting the representation of dependency values (as computed

by getDeps)

block.fun the function for plotting a representation of the individual partitions of the data

generated in dependency logos.

summary. fun the function for plotting a representation of the summary plot for (one chunk of)

the data

weight. fun the function for plotting a representation of the weights values of the sequences

within one partition

chunks the size of chunks the data is split into. The sum of the chunk sizes must not

be greater than the number of data points in data; The default value of NULL

corresponds to one chunk containing all data points

chunk.height the (relative) height of the parts of the plot representing each of the chunks, one

height for each chunk

summary.height the (relative) height of the block summaries in the plot

minPercent the minimum percentage of the (sub) data set that may constitute its own parti-

tion in the dependency logo

threshold the threshold on the dependency value for further splits

numBestForSorting

the number of dependencies between position i and all other positions when

computing the dependency value of position i

maxNum the maximum number of splits allowed

sortByWeights are partitions sorted by their average weight (descending)

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```
dep.fun.legend if TRUE, a legend of the color scale used for plotting the dependency values in dep.fun is added to the plot

show.dependency.pvals

is TRUE, p-values are used for plotting dependency values in dep.fun instead of mutual information values

axis.labels labels for the x-axis, vector of the same length as the individual sequences

weight.ratio the factor by which the plotting width for the main plot is larger than for weight.fun partition.by specify fixed positions to partition by

forwarded to the high-level plot that contains the blocks plotted by block.fun
```

#### **Details**

The function dep. fun provided for plotting the representation of dependencies is currently implemented in plotDeparcs and plotDepmatrix. Custom implementations must have the same signature as these functions and create a single plot without using layout (or similar).

The functions block fun and summary fun provided for plotting the representation of individual partitions of the data generated in dependency logos are currently implemented in deprects, colorchart, and logo. Custom implementations must have the same signature as these functions and create a single plot without using layout (or similar).

The function weight.fun for plotting a representation of the weights values of the sequences within one partition is currently implemented in subLines and subBoxes. Custom implementations must have the same signature as these functions and create a single plot without using layout (or similar).

#### Value

a list of DLData objects with the partitions created for the dependency logo

#### Author(s)

Jan Grau <grau@informatik.uni-halle.de>

```
# read data and create DLData object
seqs <- read.table(system.file("extdata", "cjun.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[, 1],weights = log1p(seqs[, 2]) )

# plot default dependency logo
plotDeplogo(data)

# refine threshold for clearer picture
plotDeplogo(data, threshold = 0.3)

# customize different parts of the plot
plotDeplogo(data, threshold = 0.3, dep.fun = plotDepmatrix, block.fun = colorchart)

# add plots of the weights</pre>
```

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```
plotDeplogo(data, weight.fun = subBoxes)
```

plotDepmatrix

Plots a matrix representation of dependency values

## **Description**

Plots a representation of dependency values as a triangular matrix rotated by 45 degrees. Internally, dependency values are computed using getDeps on the data object.

## Usage

```
plotDepmatrix(
  data,
  axis.at.bottom = TRUE,
  add.legend = TRUE,
  show.pvals = FALSE,
  axis.labels = NULL,
  threshold = 0.1
)
```

## **Arguments**

```
data the DLData object containing the data

axis.at.bottom if TRUE, the x-axis is shown at the bottom (side=1) of the plot, and at the top (side=3) otherwise

add.legend if TRUE a legend of the color scale is added to the plot

show.pvals if TRUE, -log10 p-values (computed by pchisq) are shown instead of mutual information values

axis.labels the labels of the x-axis

threshold ignored
```

## Author(s)

Jan Grau <grau@informatik.uni-halle.de>

```
# create DLData object
seqs <- read.table(system.file("extdata", "cjun.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[, 1], weights = log1p(seqs[, 2]) )
# plot using default parameters
plotDepmatrix(data)
# plot with axis at top, without a legend (color scale), and using p-values
plotDepmatrix(data, axis.at.bottom = FALSE, add.legend = FALSE, show.pvals = TRUE)</pre>
```

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replaceColors

Replaces colors in DLData object

# Description

Replaces colors in DLData object

# Usage

```
replaceColors(data, colors)
## S3 method for class 'DLData'
replaceColors(data, colors)
```

# **Arguments**

data the data

colors the new colors

# Value

the modified DLData object

# Author(s)

Jan Grau <grau@informatik.uni-halle.de>

#### See Also

replaceColors

```
# read data and create DLData object
seqs <- read.table(system.file("extdata", "cjun.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[, 1], weights = log1p(seqs[, 2]))
replaceColors(data, c("red", "green", "blue", "yellow"))</pre>
```

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revcom

Reverse complement

# Description

Determine the reverse complementary DLData object. Only works for DNA or RNA. Data may include gap symbols.

# Usage

```
revcom(data)
## S3 method for class 'DLData'
revcom(data)
```

# **Arguments**

data

the data

#### Value

the reverse complement

## Author(s)

Jan Grau <grau@informatik.uni-halle.de>

# **Examples**

```
data <- DLData(c("ACGT", "ATTA"))
revcom(data)</pre>
```

subBoxes

Plots weights as boxplots

# Description

Plots a representation of the weights of a list of DLData objects. Each entry of the list is shown as an independent boxplot.

# Usage

```
subBoxes(sub.parts, range, axis.above = TRUE, axis.below = TRUE)
```

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## **Arguments**

```
sub.parts a list of DLData objects

range the range of values shown in the plot (i.e., the xlim value of the call to plot)

axis.above if TRUE, an axis at the top of the plot (side=3) is shown

axis.below if TRUE, an axis at the bottom of the plot (side=1) is shown
```

## Author(s)

Jan Grau <grau@informatik.uni-halle.de>

# **Examples**

```
# read data and create DLData object
seqs <- read.table(system.file("extdata", "nrsf.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[, 1], weights = log1p(seqs[, 2]))
# create dependency logo with plotted weights
plotDeplogo(data, threshold = 0.03, weight.fun = subBoxes)</pre>
```

subLines

Plots weights as lines

## Description

Plots a representation of the weights of a list of DLData objects. Each entry of the list is shown as an independent line with the median value shown as a red vertical line. Plots of list entries are separated by horizontal grey lines.

## Usage

```
subLines(sub.parts, range, axis.above = TRUE, axis.below = TRUE)
```

# Arguments

```
sub.parts a list of DLData objects

range the range of values shown in the plot (i.e., the xlim value of the call to plot)

axis.above if TRUE, an axis at the top of the plot (side=3) is shown

axis.below if TRUE, an axis at the bottom of the plot (side=1) is shown
```

## Author(s)

Jan Grau <grau@informatik.uni-halle.de>

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## **Examples**

```
# read data and create DLData object
seqs <- read.table(system.file("extdata", "nrsf.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[, 1], weights = log1p(seqs[, 2]) )
# create dependency logo with plotted weights
plotDeplogo(data, threshold = 0.03, weight.fun = subLines)</pre>
```

suggestColors

Suggests colors for symbols

# **Description**

Suggests colors for the symbols in data based on the co-occurrence of symbols at common positions, weighted by the dependency values at those positions. The idea is to assign similar colors only to symbols that either mostly occur at different positions or that are present at positions with low inter-dependencies to other positions.

## Usage

```
suggestColors(data)
## S3 method for class 'DLData'
suggestColors(data)
```

# **Arguments**

data

the data

#### Value

the colors

## Author(s)

Jan Grau <grau@informatik.uni-halle.de>

## See Also

replaceColors

```
# read data and create DLData object
seqs <- read.table(system.file("extdata", "cjun.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[, 1] ,weights = log1p(seqs[, 2]) )
suggestColors(data)</pre>
```

26 summary.DLData

summary.DLData

Summarizing DLData objects

## Description

summary method for class "DLData". The summary includes the number of sequences, the consensus sequence and the number of sequences in object that match the consensus.

#### Usage

```
## S3 method for class 'DLData'
summary(object, delete.gaps = FALSE, ...)
```

# **Arguments**

```
object an object of class "DLData"
```

delete.gaps if gaps should be removed from the consensus ... further arguments passed to or from other methods

## Value

a list with elements members containing the number of sequences, consensus containing the consensus sequences, and equal.consensus containing the number of sequences in object that are identical to consensus

## Author(s)

Jens Keilwagen, Jan Grau <grau@informatik.uni-halle.de>

```
seqs <- read.table(system.file("extdata", "cjun.txt", package = "DepLogo"),
    stringsAsFactors = FALSE)
data <- DLData(sequences = seqs[, 1], weights = log1p(seqs[, 2]) )
summary(data)</pre>
```

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