

# Package ‘DiffLogo’

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**Type** Package

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```

**Depends** R (>= 3.4), stats, cba

**Imports** grDevices, graphics, utils, tools

**Suggests** knitr, testthat, seqLogo, MotifDb

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**Description** DiffLogo is an easy-to-use tool to visualize motif differences.

**License** GPL (>= 2)

**URL** <https://github.com/mgledi/DiffLogo/>

**BugReports** <https://github.com/mgledi/DiffLogo/issues>

**biocViews** Software, SequenceMatching, MultipleComparison,  
MotifAnnotation, Visualization, Alignment

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'seqLogo.R' 'stackHeights.R' 'utilities.R'  
'diffSeqLogoSupport.R' ' pwmAlignment.R'

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alignPwmSets	<i>Multiple PWMs alignment</i>
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**Description**

Align two sets of pwms

**Usage**

```
alignPwmSets(left_pwms_set, left_alignment, right_pwms_set,  
             right_alignment, try_reverse_complement)
```

**Arguments**

left\_pwms\_set list of pwms(matrixes)  
left\_alignment alignment of left\_pwms\_set.  
right\_pwms\_set list of pwms;  
right\_alignment alignment of right\_pwms\_set.  
try\_reverse\_complement if true(default), also try reverse complement.

**Value**

list - alignment of concatenation of left\_pwms\_set and right\_pwms\_set

**Author(s)**

Lando Andrey

---

Alphabet	<i>built alphabet</i>
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---

**Description**

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

**Usage**

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

**Arguments**

chars set of symbols  
cols set of colors; one for each symbol  
supportReverseComplement boolean whether the alphabet supports reverse complementation (like DNA/RNA) or not (like ASN)

**Value**

the Alphabet object

**Author(s)**

Martin Nettling

**Examples**

```
DNA = Alphabet(c("A", "C", "G", "T"), c("green4", "blue", "orange", "red"), TRUE)
```

ASN

*ASN alphabet*

**Description**

the amino acid alphabet (20 symbols), i.e. A, C, D, E, F, G, H, I, K, L, M, N, P, Q, R, S, T, V, W, Y

**Usage**

ASN

**Format**

An object of class Alphabet of length 4.

**Author(s)**

Martin Nettling

**Examples**

ASN

baseDistributionPwm

*Generates a PWM*

**Description**

Generates a PWM consisting of only the uniform distribution or the given base\_distribution (if defined).

**Usage**

```
baseDistributionPwm(pwm_length, alphabet_length,
base_distribution = NULL)
```

**Arguments**

pwm_length	the number of positions
alphabet_length	the alphabet size
base_distribution	optional base distribution for each PWM position

**Value**

a PWM

---

calculatePvalue	<i>p-value that two PWM-positions are from the same distribution</i>
-----------------	--

---

**Description**

Calculates the p-value for the null-hypothesis that two given probability vectors p1, p2 calculated from n1/n2 observations arise from the same distribution

**Usage**

```
calculatePvalue(p1, p2, n1, n2, stackHeight = shannonDivergence,
               numberOfPermutations = 100, plotGammaDistributionFit = FALSE)
```

**Arguments**

p1	first probability vector with one probability for each symbol of the alphabet
p2	second probability vector with one probability for each symbol of the alphabet
n1	number of observations for the calculation of p1
n2	number of observations for the calculation of p2
stackHeight	function for the calculation of a divergence measure for two probability vectors
numberOfPermutations	the number of permutations to perform for the calculation of stackHeights
plotGammaDistributionFit	if TRUE the fit of a gamma distribution to the sampled stackHeights is plotted

**Value**

a numeric p-value

**Author(s)**

Hendrik Treutler

## Examples

```
p1 <- c(0.2, 0.3, 0.1, 0.4)
p2 <- c(0.2, 0.1, 0.3, 0.4)
n1 <- 100
n2 <- 200
numberOfPermutations = 100
plotGammaDistributionFit = TRUE

pValue <- calculatePvalue(p1 = p1, p2 = p2, n1 = n1, n2 = n2, stackHeight = shannonDivergence, numberOfPermutati
```

---

`createDiffLogoObject` *DiffLogo object*

---

## Description

Creates a DiffLogo object

## Usage

```
createDiffLogoObject(pwm1, pwm2, stackHeight = shannonDivergence,
baseDistribution = normalizedDifferenceOfProbabilities,
alphabet = DNA, align_pwms = FALSE,
unaligned_penalty = divergencePenaltyForUnaligned,
try_reverse_complement = TRUE, base_distribution = NULL,
length_normalization = FALSE, unaligned_from_left = 0,
unaligned_from_right = 0)
```

## Arguments

<code>pwm1</code>	representation of the first position weight matrix (PWM) of type <code>pwm</code> , <code>data.frame</code> , or <code>matrix</code>
<code>pwm2</code>	representation of the second position weight matrix (PWM) of type <code>pwm</code> , <code>data.frame</code> , or <code>matrix</code>
<code>stackHeight</code>	function for the height of a stack at position i
<code>baseDistribution</code>	function for the heights of the individual bases
<code>alphabet</code>	of type <code>Alphabet</code>
<code>align_pwms</code>	if True, will align and extend pwms.
<code>unaligned_penalty</code>	is a function for localPwmAlignment.
<code>try_reverse_complement</code>	if True, alignment will try reverse complement pwms
<code>base_distribution</code>	is a vector of length <code>nrow(pwm)</code> that is added to unaligned columns of pwms for comparing. If <code>NULL</code> , uniform distribution is used
<code>length_normalization</code>	If true, will minimize the average divergence between PWMs. Otherwise will minimize the sum of divergences between positions. In both cases unaligned positions are compared to <code>base_distribution</code> and are counted when computing the alignment length.

```

unaligned_from_left
    the number of unaligned positions on the left
unaligned_from_right
    the number of unaligned positions on the right

```

**Value**

DiffLogo object

**Author(s)**

Martin Nettling

**Examples**

```

motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

pwm1 = motifs[[motif_names[[1]]]]
pwm2 = motifs[[motif_names[[2]]]]

diffLogoObj = createDiffLogoObject(pwm1 = pwm1, pwm2 = pwm2)
diffLogo(diffLogoObj)

```

**differenceOfICs**

*normalized information content differences*

**Description**

information content differences normalized by the sum of absolute information content differences for the given pair of probability vectors

**Usage**

`differenceOfICs(p1, p2)`

**Arguments**

<code>p1</code>	probability vector representing the first symbol distribution
<code>p2</code>	probability vector representing the second symbol distribution

**Value**

a vector with one result for each symbol

**Author(s)**

Martin Nettling

## Examples

```

motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder,"/",name,".pwm",sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

pwm1 = motifs[[motif_names[[1]]]]
pwm2 = motifs[[motif_names[[2]]]]

diffLogoFromPwm(pwm1 = pwm1, pwm2 = pwm2, baseDistribution = differenceOfICs)

```

**diffLogo**

*Draw DiffLogo*

## Description

Draws the difference of two sequence logos.

## Usage

```
diffLogo(diffLogoObj, ymin = 0, ymax = 0, sparse = FALSE,
         diffLogoConfiguration = list())
```

## Arguments

diffLogoObj	a DiffLogoObject created by the function createDiffLogoObject
ymin	minimum value on the y-axis
ymax	maximum value on the y-axis
sparse	if TRUE margins are reduced and tickmarks are removed from the logo
diffLogoConfiguration	list of configuration parameters (see function diffLogoTableConfiguration(...))

## Value

none (draws difference logo)

## Author(s)

Martin Nettling

## Examples

```

motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

pwm1 = motifs[[motif_names[[1]]]]
pwm2 = motifs[[motif_names[[2]]]]

diffLogoObj = createDiffLogoObject(pwm1 = pwm1, pwm2 = pwm2)
diffLogo(diffLogoObj)

```

`diffLogoFromPwm`      *Draw DiffLogo from PWM*

## Description

Draws the difference of two sequence logos.

## Usage

```
diffLogoFromPwm(pwm1, pwm2, ymin = 0, ymax = 0,
  stackHeight = shannonDivergence,
  baseDistribution = normalizedDifferenceOfProbabilities,
  sparse = FALSE, alphabet = DNA, align_pwms = FALSE,
  unaligned_penalty = divergencePenaltyForUnaligned,
  try_reverse_complement = TRUE, base_distribution = NULL,
  length_normalization = FALSE)
```

## Arguments

<code>pwm1</code>	representation of the first position weight matrix (PWM) of type <code>pwm</code> , <code>data.frame</code> , or <code>matrix</code>
<code>pwm2</code>	representation of the second position weight matrix (PWM) of type <code>pwm</code> , <code>data.frame</code> , or <code>matrix</code>
<code>ymin</code>	minimum value on the y-axis
<code>ymax</code>	maximum value on the y-axis
<code>stackHeight</code>	function for the height of a stack at position i
<code>baseDistribution</code>	function for the heights of the individual bases
<code>sparse</code>	if TRUE margins are reduced and tickmarks are removed from the logo
<code>alphabet</code>	of type <code>Alphabet</code>
<code>align_pwms</code>	if true, <code>DiffLogo</code> will align pwms before plotting
<code>unaligned_penalty</code>	is a function for <code>localPwmAlignment</code> .

```

try_reverse_complement
    if True, alignment will try reverse complement pwms
base_distribution
    is a vector of length nrow(pwm) that is added to unaligned columns of pwms for
    comparing. If NULL, uniform distribution is used
length_normalization
    If true, will minimize the average divergence between PWMs. Otherwise will
    minimize the sum of divergences between positions. In both cases unaligned
    positions are compared to base_distribution and are counted when computing
    the alignment length.

```

**Value**

none (draws difference logo)

**Author(s)**

Martin Nettling

**Examples**

```

motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

pwm1 = motifs[[motif_names[[1]]]]
pwm2 = motifs[[motif_names[[2]]]]

diffLogoFromPwm(pwm1 = pwm1, pwm2 = pwm2)

```

**diffLogoTable** *Draw DiffLogo-table*

**Description**

Draws a table of DiffLogos.

**Usage**

```
diffLogoTable(PWMs, sampleSizes = NULL, alphabet = DNA,
  configuration = list(), ...)
```

**Arguments**

PWMs	a list/vector of position weight matrices (PWMs) each of type pwm, data.frame, or matrix
sampleSizes	the number of sequences behind each PWM
alphabet	the alphabet of the given PWMs
configuration	list of (probably part of) of configuration options. See diffLogoTableConfiguration.
...	set of parameters passed to the function 'axis' for plotting

**Value**

none (draws table of difference logos)

**Author(s)**

Martin Nettling

**Examples**

```
motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

diffLogoTable(motifs)
```

**diffLogoTableConfiguration**

*Configuration object for diffLogoTable*

**Description**

Default configuration list for diffLogoTable

**Usage**

```
diffLogoTableConfiguration(alphabet, stackHeight = shannonDivergence,
  baseDistribution = normalizedDifferenceOfProbabilities,
  uniformYaxis = TRUE, sparse = TRUE, showSequenceLogosTop = TRUE,
  enableClustering = TRUE, treeHeight = 0.5, margin = 0.02,
  ratio = 1, align_pwms = FALSE, multiple_align_pwms = TRUE,
  unaligned_penalty = divergencePenaltyForUnaligned,
  try_reverse_complement = TRUE, length_normalization = FALSE,
  number_of_permutations = 100)
```

**Arguments**

alphabet        used alphabet of type Alphabet  
 stackHeight      function for the height of a stack at position i  
 baseDistribution  
                   function for the heights of the individual bases  
 uniformYaxis     if TRUE each DiffLogo is plotted with the same scaling of the y-axis  
 sparse            if TRUE margins are reduced and tickmarks are removed from the logo  
 showSequenceLogosTop  
                   if TRUE the classical sequence logos are drawn above each column of the table  
 enableClustering  
                   if TRUE the motifs are reordered, so that similar motifs have a small vertical  
                   and horizontal distance in the table  
 treeHeight        the height of the plotted cluster tree above the columns of the table; set equal to  
                   zero to omit the cluster tree  
 margin            the space reseverved for labels  
 ratio             the ratio of the plot; this is needed to determine the margin sizes correctly  
 align\_pwms        if True, will align and extend pwms in each cell of diffLogoTable independently.  
 multiple\_align\_pwms  
                   if True, will align and extend pwms in the diffLogoTable jointly.  
 unaligned\_penalty  
                   is a function for localPwmAlignment.  
 try\_reverse\_complement  
                   if True, alignment will try reverse complement pwms  
 length\_normalization  
                   if True, divergence between pwms is divided by length of pwms.  
 numberOfPermutations  
                   number of permutations for the permutation test for the calculation of p-values

**Value**

list of parameters

**Author(s)**

Lando Andrey

**Examples**

```
diffLogoTableConfiguration(DNA)
```

---

DNA	<i>DNA alphabet</i>
-----	---------------------

---

**Description**

the DNA alphabet, i.e. A, C, G, T

**Usage**

DNA

**Format**

An object of class Alphabet of length 4.

**Author(s)**

Martin Nettling

**Examples**

DNA

---

---

drawDiffLogoTable	<i>Draws a table of DiffLogos</i>
-------------------	-----------------------------------

---

**Description**

Draws a table of DiffLogos.

**Usage**

drawDiffLogoTable(diffLogoTableObj, ...)

**Arguments**

diffLogoTableObj

the diffLogoTable-Object created by function prepareDiffLogoTable(...)

...

optional parameters for functon axis

**Value**

none (draws difference logo)

## Examples

```

motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

diffLogoTableObj = prepareDiffLogoTable(motifs)
drawDiffLogoTable(diffLogoTableObj)

```

### enrichDiffLogoObjectWithPvalues

*Enriches a difflogo object with p-values*

## Description

Enriches a difflogo object with p-values which quantifies the probability that two PWM-positions are from the same distribution

## Usage

```
enrichDiffLogoObjectWithPvalues(diffLogoObj, n1, n2,
  stackHeight = shannonDivergence, number_of_permutations = 100)
```

## Arguments

diffLogoObj	matrix of difflogo objects
n1	the number of sequences behind the first pwm behind the given difflogo object
n2	the number of sequences behind the second pwm behind the given difflogo object
stackHeight	function for the calculation of a divergence measure for two probability vectors
number_of_permutations	the number of permutations to perform for the calculation of stackHeights

## Value

enriched difflogo object

## Examples

```

motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

```

```

pwm1 = motifs[[motif_names[[1]]]]
pwm2 = motifs[[motif_names[[2]]]]
n1 <- 100
n2 <- 100
diffLogoObj = createDiffLogoObject(pwm1 = pwm1, pwm2 = pwm2)
diffLogoObj = enrichDiffLogoObjectWithPvalues(diffLogoObj, n1, n2)

```

**enrichDiffLogoTableWithPvalues***Enriches a matrix of difflogo objects with p-values***Description**

Enriches a matrix of difflogo objects with p-values which quantifies the probability that two PWM-positions are from the same distribution

**Usage**

```
enrichDiffLogoTableWithPvalues(diffLogoObjMatrix, sampleSizes,
stackHeight = shannonDivergence, number_of_permutations = 100)
```

**Arguments**

diffLogoObjMatrix	matrix of difflogo objects
sampleSizes	number of sequences behind the pwms behind the given difflogo objects
stackHeight	function for the calculation of a divergence measure for two probability vectors
number_of_permutations	the number of permutations to perform for the calculation of stackHeights

**Value**

matrix of difflogo objects enriched with p-values

**Author(s)**

Martin Nettling

**Examples**

```

motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}
sampleSizes <- c(100, 150, 200, 250)
names(sampleSizes) <- motif_names

diffLogoTableObj = prepareDiffLogoTable(motifs);
diffLogoTableObj$diffLogoObjMatrix = enrichDiffLogoTableWithPvalues(diffLogoTableObj$diffLogoObjMatrix, samp
```

**extendPwmsFromAlignmentVector**  
*Extend pwms with respect to alignment*

## Description

Extends pwms by adding base\_distribution to both sides, so that they keep aligned, but have equal length.

## Usage

```
extendPwmsFromAlignmentVector(pwms, alignment_vector,
                               base_distribution = NULL)
```

## Arguments

<b>pwms</b>	is a list of matrixes
<b>alignment_vector</b>	is a list of shifts (\$shift) and orientations (\$direction)
<b>base_distribution</b>	is a vector of length nrow(pwm) that is added to unaligned columns of pwms for comparing. If NULL, uniform distribution is used

## Value

extended pwms

## Author(s)

Lando Andrey

## Examples

```
file1 = system.file("extdata/homer/Max.motif", package = "DiffLogo")
file2 = system.file("extdata/homer/c-Myc.motif", package = "DiffLogo")
pwm1 = getPwmFromFile(file1)
pwm2 = getPwmFromFile(file2)

pwms <- list(pwm1, pwm2)
multiple_pwms_alignment = multipleLocalPwmsAlignment(pwms)
aligned_pwms = extendPwmsFromAlignmentVector(pwms, multiple_pwms_alignment$alignment$vector)
```

---

FULL_ALPHABET	<i>Complete character alphabet</i>
---------------	------------------------------------

---

**Description**

the alphabet of all 26 characters

**Usage**

`FULL_ALPHABET`

**Format**

An object of class `Alphabet` of length 4.

**Author(s)**

Hendrik Treutler

**Examples**

`FULL_ALPHABET`

---

---

<code>getAlphabetFromCharacters</code>	<i>returns the alphabet which fits to the given characters</i>
--	--

---

**Description**

returns the alphabet which fits to the given characters

**Usage**

`getAlphabetFromCharacters(characters)`

**Arguments**

`characters` a character vector of characters

**Value**

an alphabet of type `Alphabet`

**Examples**

`alphabet = getAlphabetFromSequences(c("A", "A", "C", "C", "G", "G", "T", "T"))`

`getAlphabetFromSequences`

*returns the alphabet which fits to the given sequences*

### Description

returns the alphabet which fits to the given sequences

### Usage

```
getAlphabetFromSequences(sequences)
```

### Arguments

sequences	a character vector of sequences
-----------	---------------------------------

### Value

an alphabet of type Alphabet

### Examples

```
alphabet = getAlphabetFromSequences("AACC GGTT")
```

`getPwmFromAlignment`     *Create PWM from alignment*

### Description

Creates a matrix-representation of a PWM from a set of sequences

### Usage

```
getPwmFromAlignment(alignment, alphabet = NULL, pseudoCount = 0)
```

### Arguments

alignment	a vector or list of sequences each with equal length
alphabet	of type Alphabet
pseudoCount	the number of pseudo-observations for each character in the alphabet

### Value

PWM as matrix

### Author(s)

Hendrik Treutler

**Examples**

```
motif_folder= "extdata/alignments"
motif_name = "calamodulin_1"
fileName = paste(motif_folder,"/",motif_name,".txt",sep="")
file = system.file(fileName, package = "DiffLogo")
motif = getPwmFromAlignment(readLines(file), ASN, 1)
seqLogo(pwm = motif, alphabet=ASN)
```

**getPwmFromAlignmentFile***generates a pwm from an alignment file***Description**

generates a pwm from an alignment file

**Usage**

```
getPwmFromAlignmentFile(filename, alphabet = NULL)
```

**Arguments**

filename	the alignment file
alphabet	the desired alphabet of type Alphabet

**Value**

a pwm

**Examples**

```
fileName = "extdata/alignments/calamodulin_1.txt"
file = system.file(fileName, package = "DiffLogo")
pwm = getPwmFromAlignmentFile(file)
```

**getPwmFromFastaFile**    *generates a pwm from a FASTA file***Description**

generates a pwm from a FASTA file

**Usage**

```
getPwmFromFastaFile(filename, alphabet = NULL)
```

**Arguments**

filename	the FASTA file
alphabet	the desired alphabet of type Alphabet

**Value**

a pwm

**Examples**

```
fileName = "extdata/alignments/F-box_bacteria.seq.fa"
file = system.file(fileName, package = "DiffLogo")
pwm = getPwmFromFastaFile(file)
```

**getPwmFromFile**      *generates a pwm from a file of different formats*

**Description**

Generates a pwm from a file of different formats. Supported formats are FASTA files (.fa, .fasta), alignment files (.txt, .text, .al, .alignment), PWM files (.pwm), JASPAR / Position Frequency Matrix files (.pfm), and homer files (.motif).

**Usage**

```
getPwmFromFile(filename)
```

**Arguments**

filename      the file

**Value**

a pwm

**Examples**

```
fileName = "extdata/pwm/H1-hESC.pwm"
file = system.file(fileName, package = "DiffLogo")
pwm = getPwmFromFile(file)
```

**getPwmFromHomerFile**      *generates a pwm from a homer file*

**Description**

generates a pwm from a homer file

**Usage**

```
getPwmFromHomerFile(filename)
```

**Arguments**

filename      the homer file

**Value**

a pwm

**Examples**

```
fileName = "extdata/homer/CTCF_Zf_CD4.motif"
file = system.file(fileName, package = "DiffLogo")
pwm = getPwmFromHomerFile(file)
```

---

getPwmFromPfmOrJasparFile

*generates a pwm from a jaspar file*

---

**Description**

generates a pwm from a jaspar file

**Usage**

```
getPwmFromPfmOrJasparFile(filename)
```

**Arguments**

filename        the jaspar file

**Value**

a pwm

**Examples**

```
fileName = "extdata/pfm/ctcf_jaspar.pfm"
file = system.file(fileName, package = "DiffLogo")
pwm = getPwmFromPfmOrJasparFile(file)
```

---

getPwmFromPwmFile

*generates a pwm from a pwm file*

---

**Description**

generates a pwm from a pwm file

**Usage**

```
getPwmFromPwmFile(filename)
```

**Arguments**

filename        the pwm file

**Value**

```
a pwm
```

**Examples**

```
fileName = "extdata/pwm/H1-hESC.pwm"
file = system.file(fileName, package = "DiffLogo")
pwm = getPwmFromPwmFile(file)
```

`getSequencesFromAlignmentFile`

*extracts the sequences from an alignment file*

**Description**

extracts the sequences from an alignment file

**Usage**

```
getSequencesFromAlignmentFile(filename)
```

**Arguments**

filename	the alignment file
----------	--------------------

**Value**

a vector of sequences

**Examples**

```
fileName = "extdata/alignments/calmodulin_1.txt"
file = system.file(fileName, package = "DiffLogo")
sequences = getSequencesFromAlignmentFile(file)
```

`getSequencesFromFastaFile`

*extracts the sequences from a FASTA file*

**Description**

extracts the sequences from a FASTA file

**Usage**

```
getSequencesFromFastaFile(filename)
```

**Arguments**

filename	the FASTA file
----------	----------------

**Value**

a vector of sequences

**Examples**

```
fileName = "extdata/alignments/F-box_bacteria.seq.fa"
file = system.file(fileName, package = "DiffLogo")
sequences = getSequencesFromFastaFile(file)
```

---

informationContent     *information content*

---

**Description**

the information content for the given probability vector

**Usage**

```
informationContent(p)
```

**Arguments**

p                    probability vector representing the symbol distribution

**Value**

an object consisting of height a ylab

**Author(s)**

Martin Nettling

**Examples**

```
motif_folder= "extdata/pwm"
motif_name = "HepG2"
fileName = paste(motif_folder,"/",motif_name,".pwm",sep="")
file = system.file(fileName, package = "DiffLogo")
motif = getPwmFromPwmFile(file)
seqLogo(pwm = motif, stackHeight = informationContent)
```

**localPwmAlignment**      *Align pwms*

## Description

Finds best local alignment for two PWMs.

## Usage

```
localPwmAlignment(pwm_left, pwm_right, divergence = shannonDivergence,
  unaligned_penalty = divergencePenaltyForUnaligned,
  try_reverse_complement = TRUE, base_distribution = NULL,
  length_normalization = FALSE)
```

## Arguments

pwm_left	first PWM, a matrix of type matrix
pwm_right	first PWM, a matrix of type matrix
divergence	is a measure of difference between two pwm columns. Smaller is more similar. If you want to use non-uniform background distribution, provide your own function.
unaligned_penalty	distance for unaligned columns at edges of matrixes. See divergencePenaltyForUnaligned as an example for providing your own function
try_reverse_complement	If false the alignment will not be performed on reverse complements. If true, the input pwms should have column order of ACTG/ACGU.
base_distribution	is a vector of length nrow(pwm) that is added to unaligned columns of pwms for comparing. If NULL, uniform distribution is used
length_normalization	If true, will minimize the average divergence between PWMs. Otherwise will minimize the sum of divergences between positions. In both cases unaligned positions are compared to base_distribution and are counted when computing the alignment length.

## Value

list of length two containing the alignment and the divergence

## Author(s)

Lando Andrey

---

**lossOfAbsICDifferences**

*the change of information content*

---

## Description

the change of information content for the given probability vectors

## Usage

```
lossOfAbsICDifferences(p1, p2)
```

## Arguments

- |    |  |
|----|--|
| p1 | probability vector representing the first symbol distribution  |
| p2 | probability vector representing the second symbol distribution |

## Value

an object consisting of height and ylab

## Author(s)

Martin Nettling

## Examples

```
motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

pwm1 = motifs[[motif_names[[1]]]]
pwm2 = motifs[[motif_names[[2]]]]

diffLogoFromPwm(pwm1 = pwm1, pwm2 = pwm2, stackHeight = lossOfAbsICDifferences)
```

---

**multipleLocalPwmsAlignment**

*Multiple PWMs alignment*

---

## Description

Creates a multiple alignment of pwms

**Usage**

```
multipleLocalPwmsAlignment(pwms, divergence = shannonDivergence,
  unaligned_penalty = divergencePenaltyForUnaligned,
  try_reverse_complement = TRUE, base_distribution = NULL,
  length_normalization = FALSE)
```

**Arguments**

<code>pwms</code>	list of pwms
<code>divergence</code>	Divergence measure.
<code>unaligned_penalty</code>	is a function for localPwmAlignment.
<code>try_reverse_complement</code>	if True, alignment will try reverse complement pwms
<code>base_distribution</code>	is a vector of length nrow(pwm) that is added to unaligned columns of pwms for comparing. If NULL, uniform distribution is used
<code>length_normalization</code>	If true, will minimize the average divergence between PWMs. Otherwise will minimize the sum of divergences between positions. In both cases unaligned positions are compared to base_distribution and are counted when computing the alignment length.

**Value**

list

**Author(s)**

Lando Andrey

**Examples**

```
file1 = system.file("extdata/homer/Max.motif", package = "DiffLogo")
file2 = system.file("extdata/homer/c-Myc.motif", package = "DiffLogo")
pwm1 = getPwmFromFile(file1)
pwm2 = getPwmFromFile(file2)

multiple_pwms_alignment = multipleLocalPwmsAlignment(list(pwm1, pwm2))
```

`normalizedDifferenceOfProbabilities`  
*normalized probability differences*

**Description**

probability differences normalized by the sum of absolute probability differences for the given pair of probability vectors

**Usage**

```
normalizedDifferenceOfProbabilities(p1, p2)
```

**Arguments**

p1	probability vector representing the first symbol distribution
p2	probability vector representing the second symbol distribution

**Value**

a vector with one result for each symbol

**Author(s)**

Martin Nettling

**Examples**

```
motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

pwm1 = motifs[[motif_names[[1]]]]
pwm2 = motifs[[motif_names[[2]]]]

diffLogoFromPwm(pwm1 = pwm1, pwm2 = pwm2, baseDistribution = normalizedDifferenceOfProbabilities)
```

normalizePWM

*normalizes the given pwm*

**Description**

normalizes the given pwm to column-sums of 1.0

**Usage**

```
normalizePWM(pwm)
```

**Arguments**

pwm	a pwm
-----	-------

**Value**

a normalized pwm

## Examples

```
pwm = matrix(1:40, nrow = 4, dimnames = list(c("A", "C", "G", "T"), 1:10))
pwm = normalizePWM(pwm)
```

**prepareDiffLogoTable** *Prepare a table of difflogo objects*

## Description

Prepares a DiffLogoTable and generates an object that contains the hierarchical clustering and a matrix of prepared difference logos.

## Usage

```
prepareDiffLogoTable(PWMs, alphabet = DNA, configuration = list())
```

## Arguments

PWMs	a list/vector of position weight matrices (PWMs) each of type pwm, data.frame, or matrix
alphabet	the alphabet of the given PWMs
configuration	list of (probably part of) of configuration options. See diffLogoTableConfiguration.

## Value

matrix of difference logos

## Author(s)

Martin Nettling

## Examples

```
motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}
sampleSizes <- c(100, 150, 200, 250)

diffLogoTableObj = prepareDiffLogoTable(motifs);
```

probabilities	<i>probabilities</i>
---------------	----------------------

### Description

the given probabilities

### Usage

```
probabilities(p)
```

### Arguments

p	probability vector representing the symbol distribution
---	---

### Value

the given vector

### Author(s)

Martin Nettling

### Examples

```
motif_folder= "extdata/pwm"
motif_name = "HepG2"
fileName = paste(motif_folder,"/",motif_name,". pwm",sep="")
file = system.file(fileName, package = "DiffLogo")
motif = getPwmFromPwmFile(file)
seqLogo(pwm = motif, baseDistribution = probabilities)
```

pwmDivergence	<i>PWM divergence</i>
---------------	-----------------------

### Description

Counts PWM divergence as sum of divergencies of their columns.

### Usage

```
pwmDivergence(pwm_left, pwm_right, divergence = shannonDivergence)
```

### Arguments

pwm_left	is a PWM representation in type of matrix
pwm_right	is a PWM representation in type of matrix. The result is symmetric on pwm_left and pwm_right
divergence	is a Divergence function on columns.

**Value**

float - sum of divergences

**pwmsDistanceMatrix**      *Multiple PWMs alignment*

**Description**

Creates a distance matrix for pwms

**Usage**

```
pwmsDistanceMatrix(pwms, diagonal_value = 0,
                   bottom_default_value = NULL, divergence = shannonDivergence,
                   unaligned_penalty = divergencePenaltyForUnaligned,
                   try_reverse_complement = TRUE, base_distribution = NULL,
                   length_normalization = FALSE)
```

**Arguments**

pwms	list of pwms
diagonal_value	value to put on diagonal.
bottom_default_value	value to put on bottom triangle. Set to NULL to get symmetric distance matrix.
divergence	divergence measure.
unaligned_penalty	is a function for localPwmAlignment.
try_reverse_complement	if True, alignment will try reverse complement pwms
base_distribution	is a vector of length nrow(pwm) that is added to unaligned columns of pwms for comparing. If NULL, uniform distribution is used
length_normalization	is a vector of length nrow(pwm) that is added to unaligned columns of pwms for comparing. If NULL, uniform distribution is used

**Value**

list

**Author(s)**

Lando Andrey

---

```
reverseAlignmentVector  
Reverse for alignment vector
```

---

**Description**

Returns alignment vector as if all pwm were reverted.

**Usage**

```
reverseAlignmentVector(alignment_vector, pwms)
```

**Arguments**

alignment_vector	list of list which \$shift and \$orientation
pwms	list of matrixes.

**Value**

list - reversed alignment vector

**Author(s)**

Lando Andrey

---

RNA	<i>RNA alphabet</i>
-----	---------------------

---

**Description**

the RNA alphabet, i.e. A, C, G, U

**Usage**

```
RNA
```

**Format**

An object of class Alphabet of length 4.

**Author(s)**

Martin Nettling

**Examples**

```
RNA
```

**seqLogo***Draw sequence logo***Description**

Draws the classic sequence logo.

**Usage**

```
seqLogo(pwm, sparse = FALSE, drawLines = 0.5,
        stackHeight = informationContent, baseDistribution = probabilities,
        alphabet = DNA, main = NULL)
```

**Arguments**

pwm	representation of a position weight matrix (PWM) of type pwm, data.frame, or matrix
sparse	if TRUE margins are reduced and tickmarks are removed from the logo
drawLines	distance between background lines
stackHeight	function for the height of a stack at position i
baseDistribution	function for the heights of the individual bases
alphabet	of type Alphabet
main	the main title for the plot

**Value**

none (draws sequence logo)

**Author(s)**

Martin Nettling

**Examples**

```
motif_folder= "extdata/pwm"
motif_name = "HepG2"
fileName = paste(motif_folder,"/",motif_name,".pwm",sep="")
file = system.file(fileName, package = "DiffLogo")
motif = getPwmFromPwmFile(file)
seqLogo(pwm = motif)
```

shannonDivergence	<i>shannon divergence</i>
-------------------	---------------------------

## Description

the shannon divergence for the given pair of probability vectors

## Usage

```
shannonDivergence(p1, p2)
```

## Arguments

p1	probability vector representing the first symbol distribution
p2	probability vector representing the second symbol distribution

## Value

an object consisting of height and ylab

## Author(s)

Martin Nettling

## Examples

```
motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

pwm1 = motifs[[motif_names[[1]]]]
pwm2 = motifs[[motif_names[[2]]]]

diffLogoFromPwm(pwm1 = pwm1, pwm2 = pwm2, stackHeight = shannonDivergence)
```

sumOfAbsICDifferences	<i>sum of absolute information content differences</i>
-----------------------	--

## Description

the sum of absolute information content differences for the given pair of probability vectors

## Usage

```
sumOfAbsICDifferences(p1, p2)
```

**Arguments**

- p1 probability vector representing the first symbol distribution  
 p2 probability vector representing the second symbol distribution

**Value**

an object consisting of height and ylab

**Author(s)**

Martin Nettling

**Examples**

```
motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder, "/", name, ".pwm", sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

pwm1 = motifs[[motif_names[[1]]]]
pwm2 = motifs[[motif_names[[2]]]]

diffLogoFromPwm(pwm1 = pwm1, pwm2 = pwm2, stackHeight = sumOfAbsICDifferences)
```

**sumOfAbsProbabilityDifferences**  
*sum of absolute probability differences*

**Description**

the sum of absolute probability differences for the given pair of probability vectors

**Usage**

```
sumOfAbsProbabilityDifferences(p1, p2)
```

**Arguments**

- p1 probability vector representing the first symbol distribution  
 p2 probability vector representing the second symbol distribution

**Value**

an object consisting of height and ylab

**Author(s)**

Martin Nettling

## Examples

```

motif_folder= "extdata/pwm"
motif_names = c("HepG2", "MCF7", "HUVEC", "ProgFib")
motifs = list()
for (name in motif_names) {
  fileName = paste(motif_folder,"/",name,".pwm",sep="")
  file = system.file(fileName, package = "DiffLogo")
  motifs[[name]] = getPwmFromPwmFile(file)
}

pwm1 = motifs[[motif_names[[1]]]]
pwm2 = motifs[[motif_names[[2]]]]

diffLogoFromPwm(pwm1 = pwm1, pwm2 = pwm2, stackHeight = sumOfAbsProbabilityDifferences)

```

**sumProbabilities**      *sum of probabilities, i.e. 1.0*

## Description

the sum of probabilities for the given probability vector, i.e. 1.0

## Usage

```
sumProbabilities(p)
```

## Arguments

p	probability vector representing the symbol distribution
---	---

## Value

an object consisting of height and ylab

## Author(s)

Martin Nettling

## Examples

```

motif_folder= "extdata/pwm"
motif_name = "HepG2"
fileName = paste(motif_folder,"/",motif_name,".pwm",sep="")
file = system.file(fileName, package = "DiffLogo")
motif = getPwmFromPwmFile(file)
seqLogo(pwm = motif, stackHeight = sumProbabilities)

```

switchDirection	<i>Switches between 'forward' and 'reverse'</i>
-----------------	---

**Description**

Switches between 'forward' and 'reverse'

**Usage**

```
switchDirection(direction)
```

**Arguments**

direction	either 'forward' or 'reverse'
-----------	-------------------------------

**Value**

either 'reverse' or 'forward'

twoSetsAveragePwmDivergenceFromAlignmentVector	<i>Average divergence between two sets.</i>
--	---

**Description**

Computes average pwm divergence from alignment vector for two datasets. This equals to average divergence between all pairs where one pwm comes from left set, and other comes from right

**Usage**

```
twoSetsAveragePwmDivergenceFromAlignmentVector(left_pwms_list,
left_pwms_alignment, right_pwms_list, right_pwms_alignment,
divergence = shannonDivergence)
```

**Arguments**

left_pwms_list	is a list of matrixes
left_pwms_alignment	is a list of shifts (\$shift) and orientations (\$direction)
right_pwms_list	is a list of matrixes
right_pwms_alignment	is a list of shifts (\$shift) and orientations (\$direction)
divergence	divergence measure.

**Value**

float

**Author(s)**

Lando Andrey

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