Package 'PWMEnrich'

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Description A toolkit of high-level functions for DNA motif scanning and enrichment analysis built upon Biostrings. The main functionality is PWM enrichment analysis of already known PWMs (e.g. from databases such as MotifDb), but the package also implements high-level functions for PWM scanning and visualisation. The package does not perform ``de novo" motif discovery, but is instead focused on using motifs that are either experimentally derived or computationally constructed by other tools.

License LGPL (>= 2)

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PWMEnrich-package

PWMEnrich: PWM enrichment analysis

Description

A toolkit of high-level functions for DNA motif scanning and enrichment analysis built upon Biostrings. The main functionality is PWM enrichment analysis of already known PWMs (e.g. from databases such as MotifDb), but the package also implements high-level functions for PWM scanning and visualisation. The package does not perform "de novo" motif discovery, but is instead focused on using motifs that are either experimentally derived or computationally constructed by other tools.

.inputParamMotifs

Normalizes the motifs input argument for multiple functions

Description

Normalizes the motifs input argument for multiple functions

Usage

.inputParamMotifs(motifs)

Arguments

motifs

a list of motifs either as frequency matrices (PFM) or as PWM objects. If PFMs are specified they are converted to PWMs using uniform background.

.inputParamSequences 5

.inputParamSequences Normalize the sequences input argument

Description

Normalize the sequences input argument

Usage

```
.inputParamSequences(sequences)
```

Arguments

sequences a set of sequences to be scanned, a list of DNAString or other scannable objects

.inputPFMfromMatrixOrPWM

Check the frequency matrix input parameter for motifSimilarity

Description

Check the frequency matrix input parameter for motifSimilarity

Usage

```
.inputPFMfromMatrixOrPWM(m)
```

Arguments

m either a PWM object or a matrix

Value

corresponding PFM

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.normalize.bg.seq

check consistency of bg.seq input parameter

Description

check consistency of bg.seq input parameter

Usage

```
.normalize.bg.seq(bg.seq)
```

Arguments

bg.seq

a set of background sequences, either a list of DNAString object or DNAStringSet object

.normargPfm

Input parameter normalization for PWMUnscaled

Description

This function is from Biostrings package. A Position Frequency Matrix (PFM) is also represented as an ordinary matrix. Unlike a PWM, it must be of type integer (it will typically be the result of consensusMatrix()).

Usage

```
.normargPfm(x)
```

Arguments

Х

a frequency matrix

.normargPriorParams

Input parameter normalization function for PWMUnscaled

Description

This function is from Biostrings package

Usage

```
.normargPriorParams(prior.params)
```

Arguments

```
prior.params Typical 'prior.params' vector: c(A=0.25, C=0.25, G=0.25, T=0.25)
```

affinitySequenceSet 7

 $affinity {\tt Sequence Set}$

Calculate total affinity over a set of sequences

Description

Calculate total affinity over a set of sequences

Usage

```
affinitySequenceSet(scores, seq.len, pwm.len)
```

Arguments

scores affinity scores for individual sequences

seq.len lengths of sequences pwm.len lengths of PWMs

 $as. data. frame, \verb|MotifEnrichmentReport-method| \\ Convert\ a\ MotifEnrichmentReport\ into\ a\ data. frame\ object$

Description

Convert a MotifEnrichmentReport into a data.frame object

Usage

```
## S4 method for signature 'MotifEnrichmentReport'
as.data.frame(x, row.names = NULL, optional = FALSE, ...)
```

Arguments

```
x the MotifEnrichmentReport object
row.names unused
optional unused
```

unused

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cloverPvalue1seq

Calculate the Clover P-value as described in the Clover paper

Description

This function only take one background sequence as input, it also just calculates the P-value so it is more efficient.

Usage

```
cloverPvalue1seq(
  scores,
  seq.len,
  pwm.len,
  bg.fwd,
  bg.rev,
  B = 1000,
  verbose = TRUE,
  clover = NULL
)
```

Arguments

the affinity scores for individual sequences
lengths of sequences
lengths of PWMs
the raw score of forward strand
the raw scores of reverse strand
the number of random replicates
if to give verbose progress reports
the clover scores if already calculated

Value

P-value

cloverScore 9

cloverScore	Calculate the Clover score using the recursive formula from Frith et al

Description

Calculate the Clover score using the recursive formula from Frith et al

Usage

```
cloverScore(scores, lr3 = FALSE, verbose = FALSE)
```

Arguments

scores a matrix of average odds scores, where columns are motifs, and rows sequences

1r3 if to return a matrix of LR3 scores, where columns correpond to motifs, and

rows to subset sizes

verbose if to produce verbose output of progress

Value

the LR4 score, which is the mean of LR3 scores over subset sizes

colMedians	Calculate medians of columns	

Description

Calculate medians of columns

Usage

colMedians(x)

Arguments

x a matrix

colSds

Calculate standard deviations of columns

Description

Calculate standard deviations of columns

Usage

colSds(x)

Arguments

Χ

a matrix

concatenateSequences Concatenata DNA sequences into a single character object

Description

Concatenata DNA sequences into a single character object

Usage

concatenateSequences(sequences)

Arguments

sequences

either a list of DNAString objects, or a DNAStringSet

Value

a single character string

cutoffZscore 11

Description

The Z-score is calculated separately for each sequence

Usage

```
cutoffZscore(scores, seq.len, pwm.len, bg.P)
```

Arguments

scores	the hit counts for the sequences
seq.len	the length distribution of sequences
pwm.len	the length distribution of the PWMs

bg.P background probabilities of observing a motif hit at nucleotide resolution (scaled

to sequence length, not 2 * length)

Value

Z-score

cutoffZscoreSequenceSet

Z-score calculation for cutoff hits for group of sequences

Description

The Z-score is calculated as if the sequence came for one very long sequence

Usage

```
cutoffZscoreSequenceSet(scores, seq.len, pwm.len, bg.P)
```

Arguments

scores	the hit counts for the sequences
seq.len	the length distribution of sequences
pwm.len	the length distribution of the PWMs

bg.P background probabilities of observing a motif hit at nucleotide resolution

Value

Z-score

12 empiricalPvalue

divideRows

Divide each row of a matrix with a vector

Description

Divide each row of a matrix with a vector

Usage

```
divideRows(m, v)
```

Arguments

m matrix to be divided

v the vector to use for division

DNAStringSetToList

Convert DNAStringSet to list of DNAString objects

Description

as.list doesn't seem to always work for DNAStringSets, so implementing this ourselves.

Usage

```
DNAStringSetToList(x)
```

Arguments

Χ

an object of class DNAStringSet

empiricalPvalue

Calculate the empirical P-value by affinity of cutoff.

Description

This is the new backend function for empirical P-values for either affinity or cutoff. The function only works on single sequences.

Usage

```
empiricalPvalue(
   scores,
   seq.len,
   pwm.len,
   bg.fwd,
   bg.rev,
   cutoff = NULL,
   B = 10000,
   verbose = FALSE,
   exact.length = FALSE)
```

Arguments

scores	the scores obtained for the sequence
seq.len	the length of the sequence, if a single value will take a single sequence of given length. If a vector of values, will take sequences of given lengths and joint them together
pwm.len	the lengths of PWMs
bg.fwd	raw odds scores for the forward strand of background
bg.rev	raw odds scores for the reverse strand of background
cutoff	if not NULL, will use hit count above this cutoff. The cutoff should be specified in log2.
В	the number of random replicates
verbose	if to give verbose progress reports
exact.length	if to take into consideration that the actual sequence lengths differ for different PWMs. For very long sequences (i.e. seq.len » pwm.len) this make very little difference, however the run time with exact.length is much longer.

 ${\tt empiricalPvalueSequenceSet}$

Empirical P-value for a set of sequences

Description

Calculate empirical P-value for a set of sequences, using either affinity or cutoff. When cutoff is used, the score is a number of motif hits above a certain log-odds cutoff.

Usage

```
empiricalPvalueSequenceSet(
   scores,
   seq.len,
   pwm.len,
   bg.fwd,
   bg.rev,
   cutoff = NULL,
   B = 10000,
   verbose = FALSE
)
```

Arguments

scores	a matrix of scores, rows for sequences, columns for PWMs
seq.len	the lengths of sequences
pwm.len	the lengths of PWMs
bg.fwd	raw odds scores for the forward strand of background
bg.rev	raw odds scores for the reverse strand of background
cutoff	if not NULL, will use hit count above this cutoff. The cutoff should be specified in log2.
В	the number of random replicates
verbose	if to give verbose progress reports

getBackgroundFrequencies

Get the four nucleotides background frequencies

Description

Estimate the background frequencies of A,C,G,T on a set of promoters from an organism

Usage

```
getBackgroundFrequencies(organism = "dm3", pseudo.count = 1, quick = FALSE)
```

Arguments

organism either a name of the organisms for which the background should be compiled

(supported names are "dm3", "mm9" and "hg19"), a BSgenome object, DNAStringSet,

or list of DNAString objects

pseudo.count the number to which the frequencies sum up to, by default 1

quick if to preform fitting on a reduced set of 100 promoters. This will not give as

good results but is much quicker than fitting to all the promoters (~10k). Usage

of this parameter is recommended only for testing and rough estimates.

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Author(s)

Robert Stojnic, Diego Diez

Examples

```
## Not run:
   getBackgroundFrequencies("dm3")
## End(Not run)
```

getPromoters

Get the promoter sequences either for a named organism such as "dm3" or a BSgenome object

Description

Get the promoter sequences either for a named organism such as "dm3" or a BSgenome object

Usage

```
getPromoters(organismOrGenome)
```

Arguments

```
organism0rGenome either organism name, e.g. "dm3", or BSgenome object
```

Value

a list of: promoters - DNAStringSet of (unique) promoters; organism - name of species; version - genome version

gevPerSequence

Apply GEV background normalization per every sequence

Description

Apply GEV background normalization per every sequence

Usage

```
gevPerSequence(scores, seq.len, pwm.len, bg.loc, bg.scale, bg.shape)
```

Arguments

scores	affinity scores for the PWMs, can contain scores for more than one sequence (as rows), P-values are extracted separately
seq.len	the length distribution of the sequences
pwm.len	the lengths of PWMs
bg.loc	list of linear regression for location parameter
bg.scale	list of linear regression for scale parameter
bg.shape	list of linear regression for shape parameter

groupReport,MotifEnrichmentResults-method

Generate a motif enrichment report for the whole group of sequences together

Description

Generate a motif enrichment report for the whole group of sequences together

Usage

```
## S4 method for signature 'MotifEnrichmentResults'
groupReport(obj, top = 0.05, bg = TRUE, by.top.motifs = FALSE, ...)
```

Arguments

obj	a MotifEnrichmentResults object
top	what proportion of top motifs should be examined in each individual sequence (by default 0.05 , i.e. 5%)
bg	if to use background corrected P-values to do the ranking (if available)
by.top.motifs	if to rank by the proportion of sequences where the motif is within 'top' percentage of motifs
	unused

Value

a MotifEnrichmentReport object containing a table with the following columns:

- 'rank' The rank of the PWM's enrichment in the whole group of sequences together
- 'target' The name of the PWM's target gene, transcript or protein complex.
- 'id' The unique identifier of the PWM (if set during PWM creation).
- 'raw.score' The raw score before P-value calculation
- 'p.value' The P-value of motif enrichment (if available)
- 'top.motif.prop' The proportion (between 0 and 1) of sequences where the motif is within top proportion of enrichment motifs.

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Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
    ###
    # load the pre-compiled lognormal background
    data(PWMLogn.dm3.MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")

# scan two sequences for motif enrichment
sequences = list(DNAString("GAAGTATCAAGTGACCAGTAAGTCCCAGATGA"),
    DNAString("AGGTAGATAGAACAGTAGGCAATGAAGCCGATG"))

res = motifEnrichment(sequences, PWMLogn.dm3.MotifDb.Dmel)

# produce a report for all sequences taken together
r.default = groupReport(res)

# produce a report where the last column takes top 1% motifs
r = groupReport(res, top=0.01)

# view the results
r

# plot the top 10 most enriched motifs
plot(r[1:10])
}
```

keepFinite

Replace all infinite values by 0

Description

Replace all infinite values by 0

Usage

```
keepFinite(x)
```

Arguments

Х

a vector of values

logNormPval	Calculate the P-value from lognormal distribution with background of equal length
-------------	---

Description

Calculate the P-value from lognormal distribution with background of equal length

Usage

```
logNormPval(scores, seq.len, pwm.len, bg.mean, bg.sd, bg.len, log = FALSE)
```

Arguments

scores	affinity scores for the PWMs, can contain scores for more than one sequence (as rows), P-values are extracted separately
seq.len	the length distribution of the sequences
pwm.len	the leggths of PWMs
bg.mean	the mean values from the background for PWMs
bg.sd	the sd values from the background
bg.len	the length distribution of the background (we currently support only constant length)
log	if to produce log p-values

 ${\tt logNormPvalSequenceSet}$

Lognormal P-value for a set of sequences

Description

Lognormal P-value for a set of sequences

Usage

```
logNormPvalSequenceSet(scores, seq.len, pwm.len, bg.mean, bg.sd, bg.len)
```

Arguments

scores	a matrix of per-sequence affinity scores
seq.len	lengths of sequences
pwm.len	lengths of pwms
bg.mean	mean background at length of bg.len
bg.sd	standard deviation of background at length of bg.len
bg.len	the length for which mean and sd are calculated

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Value

P-value

makeBackground

Make a background for a set of position frequency matrices

Description

This is a convenience front-end function to compile new backgrounds for a set of PFMs. Currently only supports D. melanogaster, but in the future should support other common organisms as well.

Usage

```
makeBackground(
  motifs,
  organism = "dm3",
  type = "logn",
  quick = FALSE,
  bg.seq = NULL,
  ...
)
```

Arguments

motifs

a list of position frequency matrices (4xL matrices)

organism

either a name of the organisms for which the background should be compiled (currently supported names are "dm3", "mm9" and "hg19"), or a BSgenome object (see BSgenome package).

type

the type of background to be compiled. Possible types are:

- "logn" estimate a lognormal background
- "cutoff" estimate a Z-score background with fixed log-odds cutoff (in log2)
- "pval" estimate a Z-score background with a fixed P-value cutoff. Note that this may require a lot of memory since the P-value of motif hits is first estimated from the empirical distribution.
- "empirical" create an empirical P-value background. Note that this may require a lot of memory (up to 10GB in default "slow" mode (quick=FALSE) for 126 JASPAR motifs and 1000 D. melanogaster promoters).
- "GEV" estimate a generalized extreme value (GEV) distribution background by fitting linear regression to distribution parameters in log space

quick

if to preform fitting on a reduced set of 100 promoters. This will not give as good results but is much quicker than fitting to all the promoters (~10k). Usage of this parameter is recommended only for testing and rough estimates.

bg.seq

a set of background sequences to use. This parameter overrides the "organism" and "quick" parameters.

. . .

other named parameters that backend function makePWM***Background functions take.

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Author(s)

Robert Stojnic, Diego Diez

Examples

```
# load in the two example de-novo motifs
motifs = readMotifs(system.file(package = "PWMEnrich", dir = "extdata", file = "example.transfac"),
 remove.acc = TRUE)
## Not run:
 # construct lognormal background
 bg.logn = makeBackground(motifs, organism="dm3", type="logn")
 # alternatively, any BSgenome object can also be used
 if(requireNamespace("BSgenome.Dmelanogaster.UCSC.dm3"))
   bg.logn = makeBackground(motifs, organism=Dmelanogaster, type="logn")
 # construct a Z-score of hits with P-value background
 bg.pval = makeBackground(motifs, organism="dm3", type="pval", p.value=1e-3)
 # now we can use them to scan for enrichment in sequences (in this case there is a consensus
 # Tin binding site).
 motifEnrichment(DNAString("TGCATCAAGTGTGTAGTG"), bg.logn)
 motifEnrichment(DNAString("TGCATCAAGTGTGTAGTG"), bg.pval)
## End(Not run)
```

makePriors

Make priors from background sequences

Description

These priors serve both as background nucleotide frequencies and pseudo-counts for PWMs.

Usage

```
makePriors(bg.seq, bg.pseudo.count)
```

Arguments

```
bg.seq a set of background sequences
bg.pseudo.count
the total pseudocount shared between nucleotides
```

Examples

```
# some example sequences
sequences = list(DNAString("AAAGAGAGTGACCGATGAC"), DNAString("ACGATGAGGATGAC"))
# make priors with pseudo-count of 1 shared between them
makePriors(sequences, 1)
```

makePWMCutoffBackground

Make a cutoff background

Description

Make a background based on number of motifs hits above a certain threshold.

Usage

```
makePWMCutoffBackground(
  bg.seq,
  motifs,
  cutoff = log2(exp(4)),
  bg.pseudo.count = 1,
  bg.source = "",
  verbose = TRUE
)
```

Arguments

bg.seq	a set of background sequences, either a list of DNAString object or DNAS-
	tringSet object

motifs a set of motifs, either a list of frequency matrices, or a list of PWM objects. If

frequency matrices are given, the background distribution is fitted from bg.seq.

cutoff the cutoff at which the background should be made, i.e. at which a motif hit is

called significant

bg.pseudo.count

the pseudo count which is shared between nucleotides when frequency matrices

are given

bg.source a free-form textual description of how the background was generated

verbose if to produce verbose output

Examples

```
## Not run:
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
   data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")

# make background for MotifDb motifs using 2Kb promoters of all D. melanogaster transcripts
```

```
# using a cutoff of 5
if(requireNamespace("BSgenome.Dmelanogaster.UCSC.dm3"))
makePWMCutoffBackground(Dmelanogaster$upstream2000, MotifDb.Dmel.PFM, cutoff=log2(exp(5)))
}
## End(Not run)

makePWMEmpiricalBackground

Make an empirical P-value background
```

Description

Make a background appropriate for empirical P-value calculation. The provided set of background sequences is contcatenated into a single long sequence which is then scanned with the motifs and raw scores are saved. This object can be very large.

Usage

```
makePWMEmpiricalBackground(
  bg.seq,
  motifs,
  bg.pseudo.count = 1,
  bg.source = "",
  verbose = TRUE,
   ...
)
```

Arguments

bg.seq	a set of background sequences, either a list of DNAString object or DNAStringSet object	
motifs	a set of motifs, either a list of frequency matrices, or a list of PWM objects. If frequency matrices are given, the background distribution is fitted from bg.seq.	
bg.pseudo.count		
	the pseudo count which is shared between nucleotides when frequency matrices are given	
bg.source	a free-form textual description of how the background was generated	
verbose	if to produce verbose output	
	currently unused (this is for convenience for makeBackground function)	

Details

For reliable P-value calculation the size of the background set needs to be at least seq.len / min.P.value. For instance, to get P-values at a resolution of 0.001 for a single sequence of 500bp, we would need a background of at least 500/0.001 = 50kb. This ensures that we can make 1000 independent 500bp samples from this background to properly estimate the P-value. For a group of sequences, we would take seq.len to be the total length of all sequences in a group.

Examples

```
## Not run:
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
   data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")
  # make empirical background by saving raw scores for each bp in the sequence. This can be
  # very large in memory!
  if(requireNamespace("BSgenome.Dmelanogaster.UCSC.dm3"))
     makePWMEmpiricalBackground(Dmelanogaster$upstream2000[1:100], MotifDb.Dmel.PFM)
}
## End(Not run)
```

makePWMGEVBackground Make a GEV background distribution

Description

Construct a lognormal background distribution for a set of sequences. Sequences concatenated are binned in 'bg.len' chunks and lognormal distribution fitted to them.

Usage

```
makePWMGEVBackground(
 bg.seq,
 motifs,
 bg.pseudo.count = 1,
 bg.len = seq(200, 2000, 200),
 bg.source = "",
 verbose = TRUE,
  fit.log = TRUE
)
```

Arguments

a set of background sequences, either a list of DNAString object or DNASbg.seq

tringSet object

motifs a set of motifs, either a list of frequency matrices, or a list of PWM objects. If

frequency matrices are given, the background distribution is fitted from bg.seq.

bg.pseudo.count

the pseudo count which is shared between nucleotides when frequency matrices

are given

bg.len the length range of background chunks

bg.source a free-form textual description of how the background was generated

verbose if to produce verbose output fit.log if to fit log odds (instead of odds)

Examples

```
## Not run:
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
   data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")

# make background for MotifDb motifs using 2kb promoters of all D. melanogaster transcripts
   if(requireNamespace("BSgenome.Dmelanogaster.UCSC.dm3"))
        makePWMGEVBackground(Dmelanogaster$upstream2000, MotifDb.Dmel.PFM)
}

## End(Not run)
```

makePWMLognBackground Make a lognormal background distribution

Description

Construct a lognormal background distribution for a set of sequences. Sequences concatenated are binned in 'bg.len' chunks and lognormal distribution fitted to them.

Usage

```
makePWMLognBackground(
  bg.seq,
  motifs,
  bg.pseudo.count = 1,
  bg.len = 250,
  bg.len.sizes = 2^(0:4),
  bg.source = "",
  verbose = TRUE,
  algorithm = "default"
)
```

Arguments

bg.seq a set of background sequences, either a list of DNAString object or DNAS-

tringSet object

motifs a set of motifs, either a list of frequency matrices, or a list of PWM objects. If

frequency matrices are given, the background distribution is fitted from bg.seq.

bg.pseudo.count

the pseudo count which is shared between nucleotides when frequency matrices

are given

bg.len background sequences will be split into tiles of this length (default: 250bp)

bg.len.sizes background tiles will be joined into bigger tiles containing this much smaller

tiles. The default is 2^(0:4), which with bg. len translates into 250bp, 500bp, 1000bp, 1500bp, 2000bp, 4000bp. Note this is only used in the "human" algo-

rithm.

bg. source a free-form textual description of how the background was generated

verbose if to produce verbose output

algorithm type of algorithm to use, valid values are: "default" and "human".

Examples

```
## Not run:
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
   data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")

# make background for MotifDb motifs using 2kb promoters of all D. melanogaster transcripts
   if(requireNamespace("BSgenome.Dmelanogaster.UCSC.dm3"))
      makePWMLognBackground(Dmelanogaster$upstream2000, MotifDb.Dmel.PFM)
}

## End(Not run)
```

makePWMPvalCutoffBackground

Construct a cutoff background from empirical background

Description

This function takes already calculated empirical background distribution and chooses cutoff for each motif based on P-value cutoff for individual sites.

Usage

```
makePWMPvalCutoffBackground(bg.p, p.value = 0.001, bg.source = "")
```

Arguments

bg.p an object of class PWMEmpiricalBackground

p.value the P-value used to find cuttoffs for each of the motifs

bg.source textual description of background source

Value

an object of type PWMCutoffBackground

Examples

```
## Not run:
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
   data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")

# make empirical background - here we use only 100 sequences for illustrative purposes
   if(requireNamespace("BSgenome.Dmelanogaster.UCSC.dm3"))
```

```
bg.p = makePWMEmpiricalBackground(Dmelanogaster$upstream2000[1:100], MotifDb.Dmel.PFM)

# use the empirical background to pick a threshold and make cutoff background
makePWMPvalCutoffBackground(bg.p, 0.001)
}

## End(Not run)
```

 ${\tt makePWMPvalCutoffBackgroundFromSeq}$

Construct a P-value cutoff background from a set of sequences

Description

This function creates a P-value cutoff background for motif enrichment.

Usage

```
makePWMPvalCutoffBackgroundFromSeq(
  bg.seq,
  motifs,
  p.value = 0.001,
  bg.pseudo.count = 1,
  bg.source = "",
  verbose = TRUE
)
```

Arguments

bg.seq a set of background sequences, either a list of DNAString object or DNAS-

tringSet object

motifs a set of motifs, either a list of frequency matrices, or a list of PWM objects. If

frequency matrices are given, the background distribution is fitted from bg.seq.

p.value the P-value used to find cuttoffs for each of the motifs

bg.pseudo.count

the pseudo count which is shared between nucleotides when frequency matrices

are given

bg. source textual description of background source

verbose if to print verbose output

Value

an object of type PWMCutoffBackground

makeStartEndPos 27

Examples

```
## Not run:
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
   data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")

   # use the empirical background to pick a threshold and make cutoff background
   makePWMPvalCutoffBackground(Dmelanogaster$upstream2000, 0.001)
}

## End(Not run)
```

makeStartEndPos

Divide total.len into fragments of length len by providing start, end positions

Description

Divide total.len into fragments of length len by providing start,end positions

Usage

```
makeStartEndPos(total.len, len)
```

Arguments

total.len total available length to be subdivided

len size of the individual chunk

Value

a data.frame containing paired up start,end positions

```
matrixShuffleZscorePerSequence
```

Obtain z-score for motif column shuffling

Description

All PWMs are shuffled at the same time. This function would be too slow to produce empirical P-values, thus we return a z-score from a small number of shuffles.

Usage

```
matrixShuffleZscorePerSequence(scores, sequences, pwms, cutoff = NULL, B = 30)
```

28 maxAligned

Arguments

scores a set of already calculated scores

sequences either one sequence or a list/set of sequences (objects of type DNAString or

DNAStringSet)

pwms a list of PWMs

cutoff if NULL, will use affinity, otherwise will use number of hits over this log2 odds

cutoff

B number of replicates, i.e. PWM column shuffles

Details

The z-scores are calculated for each sequence individually.

maxAligned Ret	urned the aligned motif parts
----------------	-------------------------------

Description

This function takes the offset of first motif relative to second and chops off the end of both motifs that are not aligned. It returns a list containing only the columns that align.

Usage

```
maxAligned(m1, m2, offset)
```

Arguments

m1 frequency matrix of first motif m2 frequency matrix of second motif

offset a number of nucleotides by which the first motif is offsetted compared to the

second

Value

a list of column-trimmed motifs m1, m2

motifDiffEnrichment 29

motifDiffEnrichment

Differential motif enrichment

Description

Test for differential enrichment between two groups of sequences

Usage

```
motifDiffEnrichment(
    sequences1,
    sequences2,
    pwms,
    score = "autodetect",
    bg = "autodetect",
    cutoff = log2(exp(4)),
    verbose = TRUE,
    res1 = NULL,
    res2 = NULL
)
```

Arguments

sequences1

First set of sequences. Can be either a single sequence (an object of class DNAS-tring), or a list of DNAString objects, or a DNAStringSet object.

sequences2

Second set of sequences. Can be either a single sequence (an object of class DNAString), or a list of DNAString objects, or a DNAStringSet object.

pwms

this parameter can take multiple values depending on the scoring scheme and background correction used. When the method parameter is set to "autodetect", the following default algorithms are going to be used:

- if pwms is a list containing either frequency matrices or a list of PWM objects then the "affinity" algorithm is selected. If frequency matrices are given, they are converted to PWMs using uniform background. For best performance, convert frequency matrices to PWMs before calling this function using realistic genomic background.
- Otherwise, appropriate scoring scheme and background correction are selected based on the class of the object (see below).

score

this parameter determines which scoring scheme to use. Following scheme as available:

- "autodetect" default value. Scoring method is determined based on the type of pwms parameter.
- "affinity" use threshold-free affinity scores without a background. The
 pwms parameter can either be a list of frequency matrices, PWM objects, or a
 PWMLognBackground object.

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"cutoff" - use number of motif hits above a score cutoff as a measure of enrichment. No background correction is performed. The pwms parameter can either be a list of frequency matrices, PWM objects, or a PWMCutoffBackground object.

bg

this parameter determines which background correction to use, if any.

- "autodetect" default value. Background correction is determined based on the type of the pwms parameter.
- "logn" use a lognormal distribution background pre-computed for a set of PWMs. This requires pwms to be of class PWMLognBackground.
- "z" use a z-score for the number of significant motif hits compared to background number of hits. This requires pwms to be of class PWMCutoffBackground.
- "none" no background correction

cutoff the score cutoff for a significant motif hit if scoring scheme "cutoff" is selected.

verbose if to produce verbose output

res1 the output of motifEnrichment if already calculated for sequences1

res2 the output of motifEnrichment if already calculated for sequences2

Details

This function calls motifEnrichment on two groups of sequences and calculates the difference statistics when possible.

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
    # load the background file for drosophila and lognormal correction
    data(PWMLogn.dm3.MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")

# get the differential enrichment
    diff = motifDiffEnrichment(DNAString("TGCATCAAGTGTGTAGTGTGAGATTAGT"),
        DNAString("TGAACGAGTAGGACGATGAGAGATTGATG"), PWMLogn.dm3.MotifDb.Dmel, verbose=FALSE)

# motifs differentially enriched in the first sequence (with lognormal background correction)
    head(sort(diff$group.bg, decreasing=TRUE))

# motifs differentially enriched in the second sequence (with lognormal background correction)
    head(sort(diff$group.bg))
}
motifEcdf

**Calculate the empirical distribution score distribution for a set of mo-
```

Description

Calculate the empirical distribution score distribution for a set of motifs

tifs

Usage

```
motifEcdf(
  motifs,
  organism = NULL,
  bg.seq = NULL,
  quick = FALSE,
  pseudo.count = 1
)
```

Arguments

motifs	a set of motifs, either a list of frequency matrices, or a list of PWM objects. If frequency matrices are given, the background distribution is fitted from bg.seq.
organism	either a name of the organisms for which the background should be compiled (supported names are "dm3", "mm9" and "hg19"), or a BSgenome object (see BSgenome package).
bg.seq	a set of background sequence (either this or organism needs to be specified!). Can be a DNAString or DNAStringSet object.
quick	if to do the fitting only on a small subset of the data (only in combination with organism). Useful only for code testing!
pseudo.count	the pseudo count which is shared between nucleotides when frequency matrices are given

Value

a list of ecdf objects (see help page for ecdf for usage).

motifEnrichment Motif enrichment

Description

Calculate motif enrichment using one of available scoring algorithms and background corrections.

Usage

```
motifEnrichment(
  sequences,
  pwms,
  score = "autodetect",
  bg = "autodetect",
  cutoff = NULL,
  verbose = TRUE,
  motif.shuffles = 30,
  B = 1000,
  group.only = FALSE
)
```

Arguments

sequences

the sequences to be scanned for enrichment. Can be either a single sequence (an object of class DNAString), or a list of DNAString objects, or a DNAStringSet object.

pwms

this parameter can take multiple values depending on the scoring scheme and background correction used. When the method parameter is set to "autodetect", the following default algorithms are going to be used:

- if pwms is a list containing either frequency matrices or a list of PWM objects then the "affinity" algorithm is selected. If frequency matrices are given, they are converted to PWMs using uniform background. For best performance, convert frequency matrices to PWMs before calling this function using realistic genomic background.
- Otherwise, appropriate scoring scheme and background correction are selected based on the class of the object (see below).

score

this parameter determines which scoring scheme to use. Following scheme as available:

- "autodetect" default value. Scoring method is determined based on the type of pwms parameter.
- "affinity" use threshold-free affinity score. The pwms parameter can either be a list of frequency matrices, PWM objects, or a PWMLognBackground object.
- "cutoff" use number of motif hits above a score cutoff. The pwms parameter can either be a list of frequency matrices, PWM objects, or a PWMCutoffBackground object.
- "clover" use the Clover algorithm (Frith et al, 2004). The Clover score of a single sequence is identical to the affinity score, while for a group of sequences is an average of products of affinities over all sequence subsets.

bg

this parameter determines how the raw score is compared to the background distribution.

- "autodetect" default value. Background correction is determined based on the type of the pwms parameter.
- "logn" use a lognormal distribution background pre-computed for a set of PWMs. This requires pwms to be of class PWMLognBackground.
- "z" use a z-score for the number of significant motif hits compared to background number of hits. This requires pwms to be of class PWMCutoffBackground.
- "pval" use empirical P-value based on a set of background sequences.
 This requires pwms to be of class PWMEmpiricalBackground. Note that PWMEmpiricalBackground objects tend to be very large so that the empirical P-value can be calculated in reasonable time.
- "ms" shuffle columns of motif matrices and use that as basis for P-value calculation. Note that since the sequences need to rescanned with all of the new shuffled motifs this can be very slow. Also, this also works only no *individual* sequences, not groups.
- "none" no background correction

cutoff

the score cutoff for a significant motif hit if scoring scheme "cutoff" is selected.

verbose if to print verbose output

motif.shuffles number of times to shuffle motifs if using "ms" background correction

B number of replicates when calculating empirical P-value

group.only if to return statistics only for the group of sequences, not individual sequences.

In the case of empirical background the P-values for individual sequences are not calculated (thus saving time), for other backgrounds they are calculated but

not returned.

Details

This function provides and interface to all algorithms available in PWMEnrich to find motif enrichment in a single or a group of sequences with/without background correction.

Since for all algorithms the first step involves calculating raw scores without background correction, the output always contains the scores without background correction together with (optional) background-corrected scores.

Unless otherwise specified the scores are returned both separately for each sequence (without/with background) and for the whole group of sequences (without/with background).

To use a background correction you need to supply a set of PWMs with precompiled background distribution parameters (see function makeBackground). When such an object is supplied as the pwm parameter, the scoring scheme and background correction are automatically determined.

There are additional packages with already pre-computed background (e.g. see package PWMEnrich.Dmelanogaster.backgr Please refer to (Stojnic & Adryan, 2012) for more details on the algorithms.

Value

a MotifEnrichmentResults object containing a subset following elements:

- "score" scoring scheme used
- · "bg" background correction used
- "params" any additional parameters
- "sequences" the set of sequences used
- "pwms" the set of pwms used
- "sequence.nobg" per-sequence scores without any background correction. For "affinity" and "clover" a matrix of mean affinity scores; for "cutoff" number of significant hits above a cutoff
- "sequence.bg" per-sequence scores after background correction. For "logn" and "pval" the P-value (smaller is better); for "z" and "ms" background corrections the z-scores (bigger is better).
- "group.nobg" aggregate scores for the whole group of sequences without background correction. For "affinity" and "clover" the mean affinity over all sequences in the set; for "cutoff" the total number of hits in all sequences.
- "group.bg" aggregate scores for the whole group of sequences with background correction. For "logn" and "pval", the P-value for the whole group (smaller is better); for "z" and "ms" the z-score for the whole set (bigger is better).

• "sequence.norm" - (only for "logn") the length-normalized scores for each of the sequences. Currently only implemented for "logn", where it returns the values normalized from LogN(0,1)distribution

• "group.norm" - (only for "logn") similar to sequence.norm, but for the whole group of sequences

References

- R. Stojnic & B. Adryan: Identification of functional DNA motifs using a binding affinity lognormal background distribution, submitted.
- MC Frith et al: Detection of functional DNA motifs via statistical over-representation, Nucleid Acid Research (2004).

Examples

}

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
  # load the pre-compiled lognormal background
  data(PWMLogn.dm3.MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")
  # scan two sequences for motif enrichment
  sequences = list(DNAString("GAAGTATCAAGTGACCAGTAGATTGAAGTAGACCAGTC"),
    DNAString("AGGTAGATAGAACAGTAGGCAATGGGGGAAATTGAGAGTC"))
  res = motifEnrichment(sequences, PWMLogn.dm3.MotifDb.Dmel)
  # most enriched in both sequences (lognormal background P-value)
  head(motifRankingForGroup(res))
  # most enriched in both sequences (raw affinity, no background)
  head(motifRankingForGroup(res, bg=FALSE))
  # most enriched in the first sequence (lognormal background P-value)
  head(motifRankingForSequence(res, 1))
  # most enriched in the first sequence (raw affinity, no background)
  head(motifRankingForSequence(res, 1, bg=FALSE))
  ###
 # Load the pre-compiled background for hit-based motif counts with cutoff of P-value = 0.001
 data(PWMPvalueCutoff1e3.dm3.MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")
  res.count = motifEnrichment(sequences, PWMPvalueCutoff1e3.dm3.MotifDb.Dmel)
  # Enrichment in the whole group, z-score for the number of motif hits
  head(motifRankingForGroup(res))
  # First sequence, sorted by number of motif hits with P-value < 0.001
  head(motifRankingForSequence(res, 1, bg=FALSE))
```

MotifEnrichmentReport-class

A report class with formatted results of motif enrichment

Description

The columns stored in this object will depend on the type of the report (either for group of sequences, or individual sequences).

Slots

d: a DataFrame object that contains the main tabular report data pwms: a list of PWM objects corresponding to rows of d

MotifEnrichmentResults-class

A wrapper class for results of motifEnrichment() that should make it easier to access the results.

Description

Note that this is only a wrapper around a list which is the return value in PWMEnrich 1.3 and as such it provides the same interface as a list (for backward compatibility), with some additional methods.

Slots

res: a list of old results with elements such as: sequence.bg, sequence.nobg, group.bg, group.nobg

motifIC

Information content for a PWM or PFM

Description

Information content for a PWM or PFM

Usage

```
motifIC(
  motif,
  prior.params = c(A = 0.25, C = 0.25, G = 0.25, T = 0.25),
  bycol = FALSE
)
```

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Arguments

motif a matrix of frequencies, or a PWM object

prior.params the prior parameters to use when a matrix is given (ignored if motif is already a

PWM)

bycol if to return values separately for each column

Value

information content in bits (i.e. log2)

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
   data(MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")
   data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")

# the nucleotide distribution is taken from the PWM (in this case genomic background)
   motifIC(MotifDb.Dmel[["ttk"]])
# information content with default uniform background because the input is a matrix,
# not PWM object
motifIC(MotifDb.Dmel.PFM[["ttk"]])
}
```

motifPrAUC

Calculate PR-AUC for motifs ranked according to some scoring scheme

Description

Note that this function asssumes that smaller values are better!

Usage

```
motifPrAUC(seq.res)
```

Arguments

seq.res

a matrix where each column represents a PWM and each row a result for a

different sequence.

```
\verb|motifRankingForGroup,MotifEnrichmentResults-method|\\
```

Get a ranking of motifs by their enrichment in the whole set of sequences

Description

Get a ranking of motifs by their enrichment in the whole set of sequences

Usage

```
## S4 method for signature 'MotifEnrichmentResults'
motifRankingForGroup(
  obj,
  bg = TRUE,
  id = FALSE,
  order = FALSE,
  rank = FALSE,
  unique = FALSE,
  ...
)
```

Arguments

obj	a MotifEnrichmentResults object
bg	if to use background corrected P-values to do the ranking (if available)
id	if to show PWM IDs instead of target TF names
order	if to output the ordering of PWMs instead of actual P-values or raw values
rank	if the output should be rank of a PWM instead of actual P-values or raw values
unique	if TRUE, only the best rank is taken for each TF (only when id = FALSE, order = FALSE)
	currently unused

Value

a vector of P-values or raw enrichments sorted such that the first motif is most enriched

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
    ###
    # load the pre-compiled lognormal background
    data(PWMLogn.dm3.MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")

# scan two sequences for motif enrichment
    sequences = list(DNAString("GAAGTATCAAGTGACCAGTAAGTCCCAGATGA"),
```

```
DNAString("AGGTAGATAGAACAGTAGGCAATGAAGCCGATG"))
res = motifEnrichment(sequences, PWMLogn.dm3.MotifDb.Dmel)

# most enriched in both sequences (sorted by lognormal background P-value)
head(motifRankingForGroup(res))

# Return a non-redundant set of TFs
head(motifRankingForGroup(res, unique=TRUE))

# sorted by raw affinity instead of P-value
head(motifRankingForGroup(res, bg=FALSE))

# show IDs instead of target TF names
head(motifRankingForGroup(res, id=TRUE))

# output the rank instead of P-value
head(motifRankingForGroup(res, rank=TRUE))
}
```

motifRankingForSequence,MotifEnrichmentResults-method

Get a ranking of motifs by their enrichment in one specific sequence

Description

Get a ranking of motifs by their enrichment in one specific sequence

Usage

```
## S4 method for signature 'MotifEnrichmentResults'
motifRankingForSequence(
  obj,
  seq.id,
  bg = TRUE,
  id = FALSE,
  order = FALSE,
  rank = FALSE,
  unique = FALSE,
  ...
)
```

Arguments

obj	a MotifEnrichmentResults object
seq.id	either the sequence number or sequence name
bg	if to use background corrected P-values to do the ranking (if available)
id	if to show PWM IDs instead of target TF names

motifRecoveryAUC 39

order	if to output the ordering of PWMs instead of actual P-values or raw values
rank	if the output should be rank of a PWM instead of actual P-values or raw values
unique	if TRUE, only the best rank is taken for each TF (only when id = FALSE, order = FALSE)
	currently unused

Value

a vector of P-values or raw enrichments sorted such that the first motif is most enriched

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
  # load the pre-compiled lognormal background
  data(PWMLogn.dm3.MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")
  # scan two sequences for motif enrichment
   sequences = list(DNAString("GAAGTATCAAGTGACCAGTAAGTCCCAGATGA"),
    DNAString("AGGTAGATAGAACAGTAGGCAATGAAGCCGATG"))
   res = motifEnrichment(sequences, PWMLogn.dm3.MotifDb.Dmel)
  # most enriched in the second sequences (sorted by lognormal background P-value)
  head(motifRankingForSequence(res, 2))
  # return unique TFs enriched in sequence 2
  head(motifRankingForSequence(res, 2, unique=TRUE))
  # sorted by raw affinity instead of P-value
  head(motifRankingForSequence(res, 2, bg=FALSE))
  # show IDs instead of target TF names
  head(motifRankingForSequence(res, 2, id=TRUE))
  # output the rank instead of P-value
  head(motifRankingForSequence(res, 2, rank=TRUE))
}
```

motifRecoveryAUC

Calculate Recovery-AUC for motifs ranked according to some scoring scheme

Description

Note that this function asssumes that smaller values are better!

Usage

```
motifRecoveryAUC(seq.res)
```

40 motifScores

Arguments

seq.res a matrix where each column represents a PWM and each row a result for a

different sequence.

motifScores

Motif affinity or number of hits over a threshold

Description

Scan a number of sequences either to find overall affinity, or a number of hits over a score threshold.

Usage

```
motifScores(
  sequences,
  motifs,
  raw.scores = FALSE,
  verbose = TRUE,
  cutoff = NULL
)
```

Arguments

sequences a set of sequences to be scanned, a list of DNAString or other scannable objects

motifs a list of motifs either as frequency matrices (PFM) or as PWM objects. If PFMs

are specified they are converted to PWMs using uniform background.

raw. scores if to return raw scores (odds) for each position in the sequence. Note that scores

for forward and reverse strand are concatenated into a single long vector of

scores (twice the length of the sequence)

verbose if to print verbose output

cutoff if not NULL, will count number of matches with score above value specified

(instead of returning the average affinity). Can either be one value, or a vector

of values for each of the motifs.

Value

if raw.scores=FALSE, returns a matrix of mean scores (after cutoff if any), where columns are motifs. The returned values are either mean odd scores (not log-odd), or number of hits above a threshold; otherwise if raw.scores=TRUE, returns a list of raw score values (before cutoff)

Examples

 $\begin{tabular}{ll} motifScoresBigMemory & This is a memory intensive version of motifScore() which is about 2 \\ & times faster \end{tabular}$

Description

The parameters and functionality are the same as motifScores. Please refer to documentation of this function for detailed explanation of functionality.

Usage

```
motifScoresBigMemory(
   sequences,
   motifs,
   raw.scores = FALSE,
   verbose = TRUE,
   cutoff = NULL,
   seq.all = NULL
)
```

Arguments

sequences set of input sequences
motifs set of input PWMs or PFMs
raw.scores if to return scores for each base-pair
verbose if to produce verbose output

42 motifSimilarity

cutoff	the cutoff for calling binding sites (in base 2 log).
seq.all	already concatenated sequences if already available (used to internally speed up things)

Details

This function is not meant to be called directly, but is indirectly called by motifScores() once a global parameters useBigMemory is set.

See Also

motifScores

motifSimilarity	Calculates similarity between two PFMs.	
-----------------	---	--

Description

This function calculates the normalized motif correlation as a measure of motif frequency matrix similarity.

Usage

```
motifSimilarity(m1, m2, trim = 0.4, self.sim = FALSE)
```

Arguments

m1	matrix with four rows representing the frequency matrix of first motif
m2	matrix with four rows representing the frequency matrix of second motif
trim	bases with information content smaller than this value will be trimmed off both motif ends
self.sim	if to calculate self similarity (i.e. without including offset=0 in alignment)

Details

This score is essentially a normalized version of the sum of column correlations as proposed by Pietrokovski (1996). The sum is normalized by the average motif length of m1 and m2, i.e. (ncol(m1)+ncol(m2))/2. Thus, for two idential motifs this score is going to be 1. For unrelated motifs the score is going to be typically around 0.

Motifs need to aligned for this score to be calculated. The current implementation tries all possible ungapped alignment with a minimal of two basepair matching, and the maximal score over all alignments is returned.

Motif 1 is aligned both to Motif 2 and its reverse complement. Thus, the motif similarities are the same if the reverse complement of any of the two motifs is given.

References

Pietrokovski S. Searching databases of conserved sequence regions by aligning protein multiplealignments. Nucleic Acids Res 1996;24:3836-3845.

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
   data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")

# calculate the similarity of tin and vnd motifs (which are almost identical)
   motifSimilarity(MotifDb.Dmel.PFM[["tin"]], MotifDb.Dmel.PFM[["vnd"]])

# similarity of two unrelated motifs
   motifSimilarity(MotifDb.Dmel.PFM[["tin"]], MotifDb.Dmel.PFM[["ttk"]])
}
```

names,MotifEnrichmentReport

Names of variables

Description

Columns stored in the motif enrichment report

Usage

```
## S4 method for signature 'MotifEnrichmentReport'
names(x)

## S4 method for signature 'MotifEnrichmentReport'
x$name

## S4 method for signature 'MotifEnrichmentReport'
x[i, j, ..., drop = TRUE]
```

Arguments

```
x the MotifEnrichmentReport object
name the variable name
i the row selector
j unused
... unused
drop unused (always FALSE)
```

Value

the names of the variables

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```
{\it names}\,, {\it MotifEnrichmentResults}\\ {\it Names}\,\,of\,variables
```

Description

Name of different pieces of information associated with MotifEnrichmentResults

Usage

```
## S4 method for signature 'MotifEnrichmentResults'
names(x)
## S4 method for signature 'MotifEnrichmentResults'
x$name
```

Arguments

x the MotifEnrichmentResults object name the variable name

Value

the names of the variables

names, PWM

Names of variables

Description

Name of different pieces of information associated with PWM Returns the motif length, i.e. the number of columns in the PWM.

Usage

```
## S4 method for signature 'PWM'
names(x)

## S4 method for signature 'PWM'
x$name

## S4 method for signature 'PWM'
length(x)
```

Arguments

x the PWM objectname the variable name

Value

the names of the variables

names, PWMCutoffBackground

Names of variables

Description

Name of different pieces of information associated with PWMCutoffBackground

Usage

```
## $4 method for signature 'PWMCutoffBackground'
names(x)
## $4 method for signature 'PWMCutoffBackground'
x$name
```

Arguments

x the PWMCutoffBackground object

name the variable name

Value

the names of the variables

 ${\tt names,PWMEmpiricalBackground}$

Names of variables

Description

Name of different pieces of information associated with PWMEmpiricalBackground

Usage

```
## S4 method for signature 'PWMEmpiricalBackground'
names(x)
## S4 method for signature 'PWMEmpiricalBackground'
x$name
```

Arguments

x the PWMEmpiricalBackground object

name the variable name

Value

the names of the variables

names, PWMGEVBackground

Names of variables

Description

Name of different pieces of information associated with PWMGEVBackground

Usage

```
## $4 method for signature 'PWMGEVBackground'
names(x)
## $4 method for signature 'PWMGEVBackground'
x$name
```

Arguments

x the PWMGEVBackground object

name the variable name

Value

the names of the variables

names, PWMLognBackground

Names of variables

Description

Name of different pieces of information associated with PWMLognBackground

Usage

```
## S4 method for signature 'PWMLognBackground'
names(x)
## S4 method for signature 'PWMLognBackground'
x$name
```

Arguments

x the PWMLognBackground object name the variable name

Value

the names of the variables

PFMtoPWM

Convert frequencies into motifs using PWMUnscaled

Description

Note that this function is deprecated and replaced by toPWM().

Usage

```
PFMtoPWM(
  motifs,
  id = names(motifs),
  name = names(motifs),
  seq.count = NULL,
  ...
)
```

Arguments

```
motifs a list of motifs represented as matrices of frequencies (PFM)

id the set of IDs for the motifs (defaults to names of the 'motifs' list)

name the set of names for the motifs (defaults to names of the 'motifs' list)

seq.count if frequencies in the motifs are normalized to 1, provides a vector of sequence counts (e.g. for MotifDb motifs)

... other parameters to PWMUnscaled
```

Examples

```
## Not run:
if (requireNamespace("PWMEnrich.Dmelanogaster.background")) {
   data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")

# convert to PWM with uniform background
   PFMtoPWM(MotifDb.Dmel.PFM)

# get background for drosophila (quick mode on a reduced dataset)
   prior = getBackgroundFrequencies("dm3", quick=TRUE)

# convert with genomic background
   PFMtoPWM(MotifDb.Dmel.PFM, prior.params=prior)
}

## End(Not run)
```

Description

Plots a graphical version of the motif enrichment report. Note that all values are plotted, if you want to plot only a subset of a report, first select this subset (see examples).

Usage

```
## S4 method for signature 'MotifEnrichmentReport,missing'
plot(
    x,
    y,
    fontsize = 14,
    id.fontsize = fontsize,
    header.fontsize = fontsize,
    widths = NULL,
    ...
)
```

Arguments

x a MotifEnrichmentReport object

y unused

fontsize font size to use in the plot id.fontsize font size to use for the motif IDs

header.fontsize

font size of the header

widths the relative widths of columns

... unused if(requireNamespace("PWMEnrich.Dmelanogaster.background")) ####

load the pre-compiled lognormal background data(PWMLogn.dm3.MotifDb.Dmel,

package = "PWMEnrich.Dmelanogaster.background")

scan two sequences for motif enrichment sequences = list(DNAString("GAAGTATCAAGTGACCAGTA

DNAString("AGGTAGATAGAACAGTAGGCAATGAAGCCGATG"))
res = motifEnrichment(sequences, PWMLogn.dm3.MotifDb.Dmel)
produce a report for all sequences taken together r = groupReport(res)

plot the top 10 most enriched motifs plot(r[1:10])

plot, PWM, missing-method

Plotting for the PWM class

Description

This function produces a sequence logo (via package seqLogo).

Usage

```
## S4 method for signature 'PWM,missing'
plot(x, y, ...)
```

Arguments

x the PWM object

y unused

... other parameters to pass to seqLogo's plot function

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
  data(MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")

# plot the tinman motif from MotifDb
  plot(MotifDb.Dmel[["tin"]])
}
```

50 plotMotifScores

plotMotifScores

Plot the raw motifs scores as returned by motifScores()

Description

This function visualises the motif scores for one or more sequences. Sequences are drawn as lines, and scores are plotted as triangles at both sides of the line (corresponding to the two strands). The width of the base of the triangle corresponds to motif width and the height to the motif log(score) that is positive and greater than the cutoff parameter (if specified). All scores have the same y-axis, so the heights of bars are comparable between sequences and motifs.

Usage

```
plotMotifScores(
  scores,
  sel.motifs = NULL,
  seq.names = NULL,
  cols = NULL,
  cutoff = NULL,
  log.fun = log2,
  main = "",
  legend.space = 0.3,
 max.score = NULL,
  trans = 0.5,
  text.cex = 0.9,
  legend.cex = 0.9,
  motif.names = NULL,
  seq.len.spacing = 8,
  shape = "rectangle"
)
```

Arguments

scores

the list of motifs scores. Each element of the list is a matrix of scores for one sequences. The columns in the matrix correspond to different motifs. Each column contains the odds (not log-odds!) scores over both strands. For example, for a sequence of length 5, scores for a 3 bp motifs could be: c(0.1, 1, 4, NA, NA, 1, 0.3, 2, NA, NA). The first 3 numbers are odds scores starting at first three bases, and the second lot of 3 numbers is the scores starting at the same positions but with the reverse complement of the motif. The last two values are NA on both strands because we do not support partial motif hits.

sel.motifs

a vector of motif names. Use this parameter to show the motif hits to only a subset of motifs for which the scores are available.

seq.names

a vector of sequence names to show in the graph. If none specified, the sequences will be named Sequence 1, Sequence 2, ...

plotMotifScores 51

cols a vector of colours to use to colour code motif hits. If none are specified, the

current palette will be used.

cutoff either a single value, or a vector of values. The values are PWM cutoffs after

log.fun (see below). Only motif scores above these cutoffs will be shown. If a single values is specified, it will be used for all PWMs, otherwise the vector

needs to specify one cutoff per PWM.

log. fun the logarithm function to use to calculate log-odds. By default log2 is used for

consistency with Biostrings.

main the main title

legend. space the proportion of horizontal space to reserve for the legend. The default is 30%.

max.score the maximal log-odds score used to scale all other scores. By default this values

is automatically determined, but it can also be set manually to make multiple

plots comparable.

trans the level of transparency. By default 50% transparency to be able to see over-

lapping binding sites

text.cex the scaling factor for sequence names

legend.cex the scaling factor for the legend

motif.names optional vector of motif names to show instead of those present as column names

in scores

seq.len.spacing

the spacing (in bp units) between the end of the sequence line and the text show-

ing the length in bp

shape the shape to use to draw motif occurances, valid values are "rectangle" (default),

"line" and "triangle"

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
    ###
    # Load Drosophila PWMs
    data(MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")

# two sequences of interest
    sequences = list(DNAString("GAAGTATCAAGTGACCAGGTGAAGTCCCAGATGA"),
        DNAString("AGGTAGATAGAACAGTAGGCAATGAAGCCGATG"))

# select the tinman and snail motifs
    pwms = MotifDb.Dmel[c("tin", "sna")]

# get the raw score that will be plotted
    scores = motifScores(sequences, pwms, raw.scores=TRUE)

# plot the scores in both sequences, green for tin and blue for sna
    plotMotifScores(scores, cols=c("green", "blue"))
}
```

52 plotMultipleMotifs

plotMultipleMotifs Plot mulitple motifs in a single plot

Description

Individual motif logos are plotted on a rows x cols grid. This function is a convenience interface for the seqLogoGrid function that deals with viewpoint placement in a matrix-like grid layout.

Usage

```
plotMultipleMotifs(
  pwms,
  titles = names(pwms),
  rows = ceiling(sqrt(length(pwms))),
  cols = ceiling(sqrt(length(pwms))),
  xmargin.scale = 0.4,
  ymargin.scale = 0.4,
  ...
)
```

Arguments

pwms	a list of PWM objects or frequency matrices
titles	a characater vector of titles for each of the plots
rows	number of rows in the grid
cols	number or cols in the grid
xmargin.scale	the scaling parameter for the X-axis margin. Useful when plotting more than one logo on a page $$
ymargin.scale	the scaling parameter for the Y-axis margin. Useful when plotting more than one logo on a page
• • •	other parameters passed to seqLogoGrid()

Details

By default will try to make a square grid plot that would fit all the motifs and use list names as captions.

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plotPFM

Plot a PFM (not PWM) using seqLogo

Description

```
Plot a PFM (not PWM) using seqLogo
```

Usage

```
plotPFM(pfm, ...)
```

Arguments

pfm a matrix where rows are the four nucleotides
... additional parameters for plot()

plotTopMotifsGroup,MotifEnrichmentResults-method

Plot the top N enrichment motifs in a group of sequences

Description

Plot the top N enrichment motifs in a group of sequences

Usage

```
## S4 method for signature 'MotifEnrichmentResults'
plotTopMotifsGroup(obj, n, bg = TRUE, id = FALSE, ...)
```

Arguments

obj	a MotifEnrichmentResults object
n	the number of top ranked motifs to plot
bg	if to use background corrected P-values to do the ranking (if available)
id	if to show PWM IDs instead of target TF names
	other parameters passed to plotMultipleMotifs()

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
    ###
    # load the pre-compiled lognormal background
    data(PWMLogn.dm3.MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")

# scan two sequences for motif enrichment
    sequences = list(DNAString("GAAGTATCAAGTGACCAGTAAGTCCCAGATGA"),
        DNAString("AGGTAGATAGAACAGTAGGCAATGAAGCCGATG"))

res = motifEnrichment(sequences, PWMLogn.dm3.MotifDb.Dmel)

# plot the top 4 motifs in a 2x2 grid
    plotTopMotifsGroup(res, 4)

# plot top 3 motifs in a single row
    plotTopMotifsGroup(res, 3, row=1, cols=3)
}
```

 ${\it plot} {\it TopMotifs Sequence}, {\it MotifEnrichment Results-method} \\ {\it Plot the top N enrichment motifs in a single sequence}$

Description

Plot the top N enrichment motifs in a single sequence

Usage

```
## S4 method for signature 'MotifEnrichmentResults'
plotTopMotifsSequence(obj, seq.id, n, bg = TRUE, id = FALSE, ...)
```

Arguments

obj	a MotifEnrichmentResults object
seq.id	either the sequence number or sequence name
n	the number of top ranked motifs to plot
bg	if to use background corrected P-values to do the ranking (if available)
id	if to show PWM IDs instead of target TF names
	other parameters passed to plotMultipleMotifs()

PWM-class 55

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
    ###
    # load the pre-compiled lognormal background
    data(PWMLogn.dm3.MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")

# scan two sequences for motif enrichment
    sequences = list(DNAString("GAAGTATCAAGTGACCAGTAAGTCCCAGATGA"),
        DNAString("AGGTAGATAGAACAGTAGGCAATGAAGCCGATG"))

res = motifEnrichment(sequences, PWMLogn.dm3.MotifDb.Dmel)

# plot the top 4 motifs in a 2x2 grid
    plotTopMotifsSequence(res, 1, 4)

# plot top 3 motifs in a single row
    plotTopMotifsSequence(res, 1, 3, row=1, cols=3)
}
```

PWM-class

A class that represents a Position Weight Matrix (PWM)

Description

A class that represents a Position Weight Matrix (PWM)

Slots

id: a systematic ID given to this PWM, could include the source, version, etc

name: the name of the transcription factor (TF) to which the PWM corresponds to

pfm: Position Frequency Matrix (PFM) from which the PWM is derived

prior.params: Defines prior frequencies of the four bases (A,C,G,T), a named vector. These will be added to individual values for the PFM and at the same time used as background probabilities

pwm: Final Position Weight Matrix (PWM) constructed using prior.params with logarithm base 2

PWMCutoffBackground-class

Hit count background distribution for a set of PWMs

Description

Hit count background distribution for a set of PWMs

Slots

bg. source: textual description of where the background distribution is derived from

bg.cutoff: the cutoff score used to find significant motif hits (in log2 odds), either a single value or a vector of values

bg.P: the density of significant motif hits per nucleotide in background

pwms: the pwms for which the background has been compiled

PWMEmpiricalBackground-class

Background for calculating empirical P-values

Description

This object contains raw scores for one very long sequence, thus it can be very large.

Slots

bg.source: textual description of where the background distribution is derived from

bg.fwd: affinity scores (odds) for the forward strand. PWMs as columns

bg.rev: affinity scores (odds) for the reverse strand. PWMs as columns

pwms: the pwms for which the background has been compiled

PWMGEVBackground-class

Generalized Extreme Values (GEV) background for P-values

Description

The three parameters of the GEV distribution are fitted by doing linear regression on log of sequence length.

Slots

bg.source: textual description of where the background distribution is derived from

bg.loc: linear regression model for estimating the location parameter based on log(L), list of lm objects of PWMs

 $\label{eq:bg.scale:linear regression model for estimating the scale parameter based on log(L), list of lm objects of PWMs$

bg.shape: linear regression model for estimating the shape parameter based on log(L), list of lm objects of PWMs

pwms: the pwms for which the background has been compiled

PWMLognBackground-class

Lognormal background distribution for a set of PWMs

Description

Lognormal background distribution for a set of PWMs

Slots

bg. source: textual description of where the background distribution is derived from

bg.len: the length to which the background is normalized to. This is a vector of values, can have a different value for each motif.

bg.mean: the mean value of the lognormal distribution at bg.len

bg.sd: the standard deviation of the lognormal distribution at bg.len

pwms: the pwms for which the background has been compiled

PWMUnscaled

Create a PWM from PFM

Description

The PWM function from Biostrings without unit scaling

Usage

```
PWMUnscaled(
    x,
    id = "",
    name = "",
    type = c("log2probratio", "prob"),
    prior.params = c(A = 0.25, C = 0.25, G = 0.25, T = 0.25),
    pseudo.count = prior.params,
    unit.scale = FALSE,
    seq.count = NULL
)
```

Arguments

x the integer count matrix representing the motif, rows as nucleotides
id a systematic ID given to this PWM, could include the source, version, etc
name the name of the transcription factor (TF) to which the PWM corresponds to

type	the type of PWM calculation, either as log2-odds, or posterior probability (frequency matrix)
prior.params	the pseudocounts for each of the nucleotides
pseudo.count	the pseudo-count values if different from priors
unit.scale	if to unit.scale the pwm (default is no unit scaling)
seq.count	if x is a normalised PFM (i.e. with probabilities instead of sequence counts),

if x is a normalised PFM (i.e. with probabilities instead of sequence counts),

then this sequence count will be used to convert x into a count matrix

Details

By default the Biostrings package scales the log-odds score so it is within 0 and 1. In this function we take a more traditional approach with no unit scaling and offer unit scaling as an additional parameter.

See ?PWM from Biostrings for more information on input arguments.

Value

a new PWM object representing the PWM

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
  data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")
   ttk = MotifDb.Dmel.PFM[["ttk"]]
   # make a PWM with uniform background
  PWMUnscaled(ttk, id="ttk-JASPAR", name="ttk")
  # custom background
  PWMUnscaled(ttk, id="ttk-JASPAR", name="ttk",
    prior.params=c("A"= 0.2, "C" = 0.3, "G" = 0.3, "T" = 0.2))
  # get background for drosophila (quick mode on a reduced dataset)
  prior = getBackgroundFrequencies("dm3", quick=TRUE)
  # convert using genomic background
  PWMUnscaled(ttk, id="ttk-JASPAR", name="ttk", prior.params=prior)
}
```

rankingProcessAndReturn

A helper function for motifRankingForGroup and motifRankingForSequence with the common code

readJASPAR 59

Description

A helper function for motifRankingForGroup and motifRankingForSequence with the common code

Usage

```
rankingProcessAndReturn(res, r, id, order, rank, unique, decreasing)
```

Arguments

res	the list of results from MotifEnrichmentResults object
r	the vector of raw results that needs to be processed

id if to return IDs instead of names order if to return the ordering of motifs rank if to return the rank of motifs

unique if to remove duplicates
decreasing specifies the sorting order

readJASPAR	Read motifs in JASPAR format	

Description

Read motifs in JASPAR format

Usage

```
readJASPAR(file, remove.ids = FALSE)
```

Arguments

file the filename

remove.ids if to strip JASPAR ID's from motif names, e.g. "MA0211.1 bap" would become

just "bap"

Value

a list of matrices representing motifs (with four nucleotides as rows)

60 readTRANSFAC

readMotifs

Read in motifs in JASPAR or TRANSFAC format

Description

The format is autodetected based on file format. If the autodetection fail then the file cannot be read.

Usage

```
readMotifs(file, remove.acc = FALSE)
```

Arguments

file the filename

remove.acc if to remove accession numbers. If TRUE, the AC entry in TRANSFAC files

is ignored, and the accession is stripped from JASPAR, e.g. motif with name "MA0211.1 bap" would become just "bap". If FALSE, botht he AC and ID are used to generate the TRANSFAC name and the original motif names are

preserved in JASPAR files.

Value

a list of 4xL matrices representing motifs (four nucleotides as rows)

Examples

```
# read in example TRANSFAC motifs without accession codes (just IDs)
readMotifs(system.file(package = "PWMEnrich", dir = "extdata", file = "example.transfac"),
    remove.acc = TRUE)

# read in the JASPAR insects motifs provided as example
readMotifs(system.file(package = "PWMEnrich", dir = "extdata", file = "jaspar-insecta.jaspar"),
    remove.acc = TRUE)
```

readTRANSFAC

Read in motifs in TRANSFAC format

Description

Read in motifs in TRANSFAC format

Usage

```
readTRANSFAC(file, remove.acc = TRUE)
```

Arguments

file the filename

remove.acc if to ignore transfac accession numbers

Value

a list of matrices representing motifs (with four nucleotides as rows)

registerCoresPWMEnrich

Register than PWMEnrich can use parallel CPU cores

Description

Certain functions (like motif scanning) can be parallelized in PWMEnrich. This function registers a number of parallel cores (via core package parallel) to be used in code that can be parallelized. After this function is called, all further PWMEnrich function calls will run in parallel if possible.

Usage

```
registerCoresPWMEnrich(numCores = NA)
```

Arguments

numCores number of cores to use (default to take all cores), or NULL if no parallel execu-

tion is to be used

Details

By default parallel execution is turned off. To turn it off after using it, call this function by passing NULL.

Examples

```
## Not run:
registerCoresPWMEnrich(4) # use 4 CPU cores in PWMEnrich
registerCoresPWMEnrich() # use maximal number of CPUs
registerCoresPWMEnrich(NULL) # do not use parallel execution
## End(Not run)
```

62 scanWithPWM

```
reverseComplement,PWM-method
```

Reverse complement for the PWM object

Description

Finds the reverse complement of the PWM

Usage

```
## S4 method for signature 'PWM'
reverseComplement(x, ...)
```

Arguments

```
x an object of type PWM
```

.. unused

Value

an object of type PWM that is reverse complement of x

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
   data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")

   reverseComplement(MotifDb.Dmel.PFM[["ttk"]]) # reverse complement of the ttk PWM
}
```

scanWithPWM

Scan the whole sequence on both strands

Description

The whole sequence is scanned with a PWM and scores returned beginning at each position. Partial motif matches are not done, thus the last #[length of motif]-1 scores are NA.

scanWithPWM 63

Usage

```
scanWithPWM(
  pwm,
  dna,
  pwm.rev = NULL,
  odds.score = FALSE,
  both.strands = FALSE,
  strand.fun = "mean"
)
```

Arguments

pwm	PWM object
dna	a DNAString or other sequence from Biostrings
pwm.rev	the reverse complement for a pwm (if it is already pre-computed)
odds.score	if to return raw scores in odds (not logodds) space
both.strands	if to return results on both strands
strand.fun	which function to use to summarise values over two strands (default is "mean")

Details

The function returns either an odds average (*not* log-odds average), maximal score on each strand, or scores on both strands.

The function by default returns the score in log2 following the package Biostrings.

Value

a vector representing scores starting at each position, or a matrix with score in the two strands

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
   data(MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")

   ttk = MotifDb.Dmel[["ttk"]]

# odds average over the two strands expressed as log2-odds
   scanWithPWM(ttk, DNAString("CGTAGGATAAAGTAACT"))

# log2-odds scores on both strands
   scanWithPWM(ttk, DNAString("CGTAGGATAAAGTAACT"), both.strands=TRUE)
}
```

64 seqLogoGrid

seqLogoGrid	Draw a motif logo on an existing viewport

Description

This function comes from the seqLogo package. It has been modified to remove some unneccessary code as suggested by W Huber (https://stat.ethz.ch/pipermail/bioconductor/2010-September/035267.html).

Usage

```
seqLogoGrid(
  pwm,
  ic.scale = TRUE,
  xaxis = TRUE,
  yaxis = TRUE,
  xfontsize = 10,
  yfontsize = 10,
  xmargin.scale = 1,
  ymargin.scale = 1,
  title = "",
  titlefontsize = 15
```

Arguments

pwm	numeric The 4xW position weight matrix.
ic.scale	logical If TRUE, the height of each column is proportional to its information content. Otherwise, all columns have the same height.
xaxis	logical If TRUE, an X-axis will be plotted.
yaxis	logical If TRUE, a Y-axis will be plotted.
xfontsize	numeric Font size to be used for the X-axis.
yfontsize	numeric Font size to be used for the Y-axis.
xmargin.scale	the scaling parameter for the X-axis margin. Useful when plotting more than one logo on a page
ymargin.scale	the scaling parameter for the Y-axis margin. Useful when plotting more than one logo on a page
title	to be shown on the top
titlefontsize	the fontsize of the title

Details

Use this function for more advanced plotting where the viewports are directly set up and maintained (see package grid).

sequenceReport,MotifEnrichmentResults-method

Generate a motif enrichment report for a single sequence

Description

Generate a motif enrichment report for a single sequence

Usage

```
## S4 method for signature 'MotifEnrichmentResults'
sequenceReport(obj, seq.id, bg = TRUE, ...)
```

Arguments

```
obj a MotifEnrichmentResults object
seq.id the sequence index or name
bg if to use background corrected P-values to do the ranking (if available)
... unused
```

Value

a MotifEnrichmentReport object containing a table with the following columns:

- 'rank' The rank of the PWM's enrichment in the sequence
- 'target' The name of the PWM's target gene, transcript or protein complex.
- 'id' The unique identifier of the PWM (if set during PWM creation).
- 'raw.score' The raw score before P-value calculation
- 'p.value' The P-value of motif enrichment (if available)

Examples

```
if(requireNamespace("PWMEnrich.Dmelanogaster.background")){
    ###
    # load the pre-compiled lognormal background
    data(PWMLogn.dm3.MotifDb.Dmel, package = "PWMEnrich.Dmelanogaster.background")

# scan two sequences for motif enrichment
    sequences = list(DNAString("GAAGTATCAAGTGACCAGTAAGTCCCAGATGA"),
        DNAString("AGGTAGATAGAACAGTAGGCAATGAAGCCGATG"))

res = motifEnrichment(sequences, PWMLogn.dm3.MotifDb.Dmel)

# reports for the two sequences
    r1 = sequenceReport(res, 1)
    r2 = sequenceReport(res, 2)
```

```
# view the results
r1
r2

# plot the top 10 most enriched motifs in the first, and then second sequence
plot(r1[1:10])
plot(r2[1:10])
}
```

 $show, \verb|MotifEnrichmentReport-method| \\ show \textit{ method for MotifEnrichmentReport}$

Description

show method for MotifEnrichmentReport

Usage

```
## S4 method for signature 'MotifEnrichmentReport'
show(object)
```

Arguments

object

the MotifEnrichmentReport object

```
show, \verb|MotifEnrichmentResults-method| \\ show \textit{ method for MotifEnrichmentResults}
```

Description

show method for MotifEnrichmentResults

Usage

```
## S4 method for signature 'MotifEnrichmentResults'
show(object)
```

Arguments

object

the MotifEnrichmentResults object

show,PWM-method 67

show,PWM-method

show method for PWM

Description

show method for PWM

Usage

```
## S4 method for signature 'PWM'
show(object)
```

Arguments

object

the PWM object

```
show, \verb"PWMCutoffBackground-method" \\ show \textit{ method for PWMCutoffBackground}
```

Description

show method for PWMCutoffBackground

Usage

```
## S4 method for signature 'PWMCutoffBackground'
show(object)
```

Arguments

object

the PWMCutoffBackground object

 $show, \verb"PWMEmpiricalBackground-method" \\ show \textit{ method for PWMEmpiricalBackground}$

Description

show method for PWMEmpiricalBackground

Usage

```
## S4 method for signature 'PWMEmpiricalBackground'
show(object)
```

Arguments

object the PWMEmpiricalBackground object

 $show, {\tt PWMGEVBackground-method} \\ show\ method\ for\ PWMGEVBackground$

Description

show method for PWMGEVBackground

Usage

```
## S4 method for signature 'PWMGEVBackground'
show(object)
```

Arguments

object the PWMGEVBackground object

```
show, \verb"PWMLognBackground-method" \\ show \textit{ method for PWMLognBackground}
```

Description

show method for PWMLognBackground

Usage

```
## S4 method for signature 'PWMLognBackground'
show(object)
```

Arguments

object

the PWMLognBackground object

toPWM

Convert motifs into PWMs

Description

Convert motifs into PWMs

Usage

```
toPWM(
  motifs,
  ids = names(motifs),
  targets = names(motifs),
  seq.count = 50,
  prior = c(A = 0.25, C = 0.25, G = 0.25, T = 0.25),
  ...
)
```

Arguments

motifs	a list of motifs either as position probability matrices (PPM) or frequency matirces (PFMs)
ids	the set of IDs for the motifs (defaults to names of the 'motifs' list)
targets	the set of target TF names for the motifs (defaults to names of the 'motifs' list)
seq.count	provides a vector of sequence counts for probability matrices (PPMs). Default it 50.
prior	frequencies of the four letters in the genome. Default is uniform background.
	other parameters to PWMUnscaled

Examples

```
## Not run:
if (requireNamespace("PWMEnrich.Dmelanogaster.background")) {
   data(MotifDb.Dmel.PFM, package = "PWMEnrich.Dmelanogaster.background")

   toPWM(MotifDb.Dmel.PFM) # convert to PWM with uniform background

   # get background for drosophila (quick mode on a reduced dataset)
   prior = getBackgroundFrequencies("dm3", quick=TRUE)
   toPWM(MotifDb.Dmel.PFM, prior=prior) # convert with genomic background
}

## End(Not run)
```

tryAllMotifAlignments *Try all motif alignments and return max score*

Description

This function tries all offsets of motif1 compared to motif2 and returns the maximal (unnormalized) correlation score.

Usage

```
tryAllMotifAlignments(m1, m2, min.align = 2, exclude.zero = FALSE)
```

Arguments

m1 frequency matrix of motif 1 m2 frequency matrix of motif 2

min.align minimal number of basepairs that need to align

exclude.zero if to exclude offset=0, useful for calculating self-similarity

Details

The correlation score is essentially the sum of correlations of individual aligned columns as described in Pietrokovski (1996).

Value

single maximal score

References

Pietrokovski S. Searching databases of conserved sequence regions by aligning protein multiplealignments. Nucleic Acids Res 1996;24:3836-3845.

useBigMemoryPWMEnrich If to use a faster implementation of motif scanning that requires abount 5 to 10 times more memory

Description

If to use a faster implementation of motif scanning that requires abount 5 to 10 times more memory

Usage

```
useBigMemoryPWMEnrich(useBigMemory = FALSE)
```

Arguments

```
useBigMemory a boolean value denoting if to use big memory implementation
```

Examples

```
## Not run:
useBigMemoryPWMEnrich(TRUE) # switch to big memory implementation globally
useBigMemoryPWMEnrich(FALSE) # switch back to default implementation
## End(Not run)
```

```
[,PWMCutoffBackground-method
```

Get the background for a subset of PWMs

Description

Get the background for a subset of PWMs

Usage

```
## S4 method for signature 'PWMCutoffBackground' x[i, j, ..., drop = TRUE]
```

Arguments

```
x the PWMCutoffBackground object
i the indicies of PWMs
j unused
... unused
drop unused
```

 $\hbox{\verb|[,PWMEmpiricalBackground-method||}$

Get the background for a subset of PWMs

Description

Get the background for a subset of PWMs

Usage

```
## S4 method for signature 'PWMEmpiricalBackground' x[i, j, ..., drop = TRUE]
```

Arguments

drop

x the PWMEmpiricalBackground object
i the indicies of PWMs
j unused
... unused

[,PWMGEVBackground-method

Get the background for a subset of PWMs

Description

Get the background for a subset of PWMs

unused

Usage

```
## S4 method for signature 'PWMGEVBackground' x[i, j, ..., drop = TRUE]
```

Arguments

drop

the PWMGEVBackground object
the indicies of PWMs
unused
unused

unused

```
\hbox{\tt [,PWMLognBackground-method]}
```

Get the background for a subset of PWMs

Description

Get the background for a subset of PWMs

Usage

```
## S4 method for signature 'PWMLognBackground' x[i, j, ..., drop = TRUE]
```

Arguments

X	the PWMLognBackground object
i	the indicies of PWMs
j	unused
	unused
drop	unused

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