

# Package ‘motifRG’

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**Title** A package for discriminative motif discovery, designed for high throughput sequencing dataset

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**Description** Tools for discriminative motif discovery using regression methods

**Imports** Biostrings,IRanges,seqLogo,parallel,methods,grid,graphics,XVector

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**License** Artistic-2.0

**LazyLoad** yes

**biocViews** Transcription,MotifDiscovery

**Depends** R (>= 2.15), Biostrings (>= 2.26), IRanges, seqLogo, parallel, methods, grid, graphics, BSgenome, XVector, BSgenome.Hsapiens.UCSC.hg19

**NeedsCompilation** no

## R topics documented:

ctcf.motifs . . . . .	2
findMotif . . . . .	2
findMotifFasta . . . . .	4
findMotifFgBg . . . . .	4
getSequence . . . . .	5
Motif-class . . . . .	5
motifLatexTable . . . . .	6
plotMotif . . . . .	7
refinePWMMotif . . . . .	8
refinePWMMotifExtend . . . . .	9
summaryMotif . . . . .	10
YY1.control . . . . .	11
YY1.peak . . . . .	11

**Index**

**12**

ctcf.motifs

*CTCF motifs predicted by motifRG***Description**

The output produced by [findMotif](#).

**Details**

A list with following elements: motifs:a list motif descriptions of class [Motif-class](#). category:input binary specification of foreground/background. mask.motifs:if `mask=T`, then `mask.motifs` contain the description of motif is based on motif matches after the input sequences being masked by previous motifs. In this case, "motifs" contained the unmasked motif descriptions.

**References**

Unpublished

findMotif

*De-novo discovery of discriminative motifs***Description**

The function searches motifs that discriminate the given foreground and background sequences.

**Usage**

```
findMotif(all.seq, category, weights = rep(1, length(all.seq)),
  start.width=6,min.cutoff=5, min.ratio=1.3,
  min.frac=0.01, both.strand=TRUE, flank=2, max.motif=5,
  mask=TRUE,other.data=NULL, start.nmer=NULL,
  enriched.only=F,n.bootstrap = 5, bootstrap.pvalue=0.1,is.parallel =
  TRUE,mc.cores = 4,min.info=10,max.width=15,discretize=TRUE)
```

**Arguments**

<code>all.seq</code>	DNAStringSet; foreground and background sequences.
<code>category</code>	numeric vector; specify which sequences are foreground (with value 1), and background (value 0).
<code>weights</code>	numeric vector: the weights for all sequences. Default: 1
<code>start.width</code>	logical; the width for enumerating seed patterns
<code>min.cutoff</code>	numeric; the score cutoff required for seed selection. All scores are negative, the lower the better.

min.ratio	numeric; the minimum fold change of motif occurrences in foreground vs background.
min.frac	numeric; the minimum fraction of fg/bg sequences containing the candidate motifs
both.strand	logical; if true, search both strands
flank	integer; the length for step-wise pattern extension at both ends on candidate motifs
max.motif	integer; the maximum number of output motifs
mask	logical; if true, mask previous motifs when searching for the next motif
other.data	if not NULL, a matrix with additional terms for the regression model for bias adjustment
start.nmer	if not NULL, a matrix with counts for user specified seed pattern in each sequence
enriched.only	logical; if true, only predict enriched motif
n.bootstrap	integer; the number of bootstrapping tests to estimate score variance
bootstrap.pvalue	numeric: the bootstrap t.test pvalues to determine the significance of improvement
is.parallel	logical; if true, runs in parallel mode, and requires "parallel" library
mc.cores	integer; the number of CPUs for parallel run
min.info	minimal information content for the motif to prevent it from being too degenerate
max.width	maximum width of the motif for extension
discretize	logical default TRUE

### Value

return a list with following elements:

motifs	a list motif descriptions of class <b>Motif-class</b>
.	.
category	input binary specification of foreground/background
mask.motifs	if mask=T, then mask.motifs contain the description of motif is based on motif matches after the input sequences being masked by previous motifs. In this case, "motifs" contained the unmasked motif descriptions.

### Examples

```
MD.peak.seq <- readDNAStringSet(system.file("extdata", "MD.peak.fa", package="motifRG"))
MD.control.seq <- readDNAStringSet(system.file("extdata", "MD.control.fa", package="motifRG"))
category <- c(rep(1, length(MD.peak.seq)), rep(0, length(MD.control.seq)))
MD.motifs <- findMotif(append(MD.peak.seq, MD.control.seq), category, max.motif=3, enriched=TRUE)

### Get summary of motifs
```

```
summaryMotif(MD.motifs$motifs, MD.motifs$category)

### plot the dinucleotide representation of the first motif
plotMotif(MD.motifs$motifs[[1]]@match$pattern)

### Create table of motifs in Latex
motifLatexTable(MD.motifs, main="MD motifs")

### Create table of motifs in Html
motifHtmlTable(MD.motifs)
```

**findMotifFasta***Wrapper function for findMotif using fasta input***Description**

Perform motif search on two input fasta files. This is a wrapper function for `findMotif`

**Usage**

```
findMotifFasta(fg.file, bg.file, ...)
```

**Arguments**

<code>fg.file</code>	character; foreground fasta filename
<code>bg.file</code>	character; background fasta filename
<code>...</code>	Other parameters passed to <code>findMotif</code>

**findMotifFgBg***Wrapper function for findMotif using two sequence datasets***Description**

Perform motif search on two sequence datasets. This is a wrapper function for `findMotif`.

**Usage**

```
findMotifFgBg(fg.seq, bg.seq, ...)
```

**Arguments**

<code>fg.seq</code>	DNAStringSet; foreground sequences
<code>bg.seq</code>	DNAStringSet; background sequences
<code>...</code>	Other parameters passed to <code>findMotif</code>

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getSequence	<i>Fetch genomic sequences from GRanges Object</i>
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## Description

The function fetch genomic sequences given the coordinates and strand information

## Usage

```
getSequence(gr, genome)
```

## Arguments

gr	GRanges object; coordinates of sequences.
genome	BSgenome object

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Motif-class	<i>Motif objects</i>
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## Description

A Motif object contains general motif characteristics and details of motif match

## Details

A motif object has the following slots: score:absolute z-value based on the logistic regression model for the motif. sign:the sign of the motif:plus for enriched motif in the foreground sequences, and negative for depleted motif count:a numeric vector holding the number of matches in each sequence match:a data.frame with the following columns: match.strand:the strand on which the match is found; pattern:the motif match pattern; seq.id: on which sequence the match is found; pos:the position relative to sequence start of the match. pattern:the motif pattern consensus:the motif consensus pattern determined by the majority votes at each position using the following rule: the most dominate single nucleotide if its frequency is greater than 0.6, or the two most dominate nucleotide if combined frequency is greater than 0.8, or the three most dominate nucleotide if combined frequency is greater than 0.95

## See Also

[findMotif](#) [summaryMotif](#) [plotMotif](#) [motifLatexTable](#)

**motifLatexTable**      *create of table of motifs*

## Description

create a latex table to be embedded in a latex document

## Usage

```
motifLatexTable(motifs, main="", prefix="motif", dir=".", height=1,
width=3,enriched.only=F, plot.pwm= TRUE,
summary.cols=c(1,7,8,9),use.mask=TRUE)
motifHtmlTable(motifs, dir="html", prefix="motif", enriched.only=F,
plot.pwm= TRUE, summary.cols=c(1,7,8,9),use.mask=TRUE)
```

## Arguments

<b>motifs</b>	result of findMotif
<b>main</b>	The title of table
<b>prefix</b>	The prefix for the filenames of motif logos
<b>dir</b>	The directory for storing motif logo files
<b>height, width</b>	size of the sequence logo
<b>enriched.only</b>	If true, list only enriched motifs
<b>plot.pwm</b>	If true, plot PWM logo instead of di-nucleotide logo
<b>summary.cols</b>	The selected columns of summary table created by summaryMotif included in the table
<b>use.mask</b>	If true, use masked motif match summary statistics

## Value

**motifLatexTable** outputs a latex table to the stdout console. **motifHtmlTable** outputs a html file named as <prefix>.html in "dir" directory.

## See Also

[findMotif](#)

## Examples

```
data(ctcf.motifs)
### Create table of motifs in Latex
motifLatexTable(ctcf.motifs, main="CTCF motifs", dir="motif")

### Create table of motifs in Html
motifHtmlTable(ctcf.motifs, dir="Html")
```

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<code>plotMotif</code>	<i>plot motif sequence matches</i>
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## Description

plot aligned sequences, revealing the independent position specificity and dependency among adjacent positions.

## Usage

```
plotMotif(match, logodds=F, entropy=F, bg.ld=NULL, alphabet=c("A", "C", "G", "T"), has.box=T, ...)
```

## Arguments

<code>match</code>	motif match to be plotted. character or <a href="#">DNAStringSet</a> object.
<code>logodds</code>	logical; if true, plot the enrichment/depletion of a adjacent pair relative to the independent model.
<code>entropy</code>	logical; if true, areas outside the core region of the motif are dimmed
<code>bg.ld</code>	Experimental features: background dinucleotide logodds against independent model. if <code>logodds=T</code> , then the background logodds will be substracted
<code>alphabet</code>	the alphabets used in the sequence. Do not change its value
<code>has.box</code>	logical; if true, plot the boundaries of the motif
<code>...</code>	other arguments passed to the lower level plot function

## Details

X-axis refers to the positions of the motifs.

Y-axis correspond to the alphabets.

Letter sizes define the frequencies of the nucleotides at a given position.

Edges between the letters specify the dinucleotide relationship. The depth of the color correspond to the dinucleotide frequency. If `logodds=T`, thinner edges will be plotted between dependent pairs. The edge is colored red if the pair is depleted (relative to the expected frequency if the pair is independent), and green if the pair is enriched. The gradient of color red/green correspond to the level of dependency.

## Examples

```
data(ctcf.motifs)
### plot the dinucleotide representation of the first motif
plotMotif(ctcf.motifs$motifs[[1]]@match$pattern)
plotMotif(ctcf.motifs$motifs[[1]]@match$pattern, logodds=TRUE)
plotMotif(ctcf.motifs$motifs[[1]]@match$pattern, logodds=TRUE, entropy=TRUE)
```

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<b>refinePWMMotif</b>	<i>create a PWM (Position Weight Model) model given a initial set of motif matches and input sequences</i>
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## Description

Create a PWM model given a initial set of motif matches and input sequences

## Usage

```
refinePWMMotif(motifs=NULL, seqs, pwm.ld= NULL, max.iter=50,
tol=10^-4, mod="oops", null=rep(0.25, 4),pseudo=1, weights=rep(1,
length(seqs)), motif.weights=NULL)
```

## Arguments

<b>motifs</b>	The initial set of motif matches. character vector or DNAStringSet object
<b>seqs</b>	Input sequences. character vector or DNAStringSet object
<b>pwm.ld</b>	The initial PWM matrixes in logodds transformation. Either "motifs" or "pwm.ld" is not NULL
<b>max.iter</b>	Maximum number of iterations for refinement
<b>tol</b>	Convergence criteria. The percentage of total PWM scores improvement required for convergence.
<b>mod</b>	Motif occurrence model. If mod=="oops", assume one motif match per sequence. If mod=="zoops", assume zero or one motif match per sequence.
<b>null</b>	A numeric vector specifying the background model
<b>pseudo</b>	Pseudo counts for PWM construction
<b>weights</b>	a numeric vector specifying the weights for all sequences. Default: 1 for all sequences
<b>motif.weights</b>	a numeric vector specifying the weights for initial sets of motifs. Default: NULL

## Value

Return a list with two elements:

<b>model</b>	a list with two elements. "prob": PWM model, sum of columns add to 1. "logodd": PWM model in logodds form, log2 of original matrix subtract the background model
.	.
<b>match</b>	a data.frame specifying the motif matches in each sequence. Columns are: "match": the sequence of the match, "score": PWM score, "strand", the strand of the match in the input sequence, "pos": start position of the motif match. If multiple matches are allowed, then "seq.id" specifies the index of the input sequence for the motif match.
<b>score</b>	Total PWM score of the motif matches

**See Also**

[findMotif](#) [refinePWMMotifExtend](#)

**Examples**

```
ctcf.seq <- readDNAStringSet(system.file("extdata","ctcf.fa", package="motifRG"))
data(ctcf.motifs)
### refine PWM model based on motif matches
pwm.match <- refinePWMMotif(ctcf.motifs$motifs[[1]]@match$pattern, ctcf.seq)
### plot traditional motif logo
library("seqLogo")
seqLogo(pwm.match$model$prob)
### plot dinucleotide motif logo
plotMotif(pwm.match$match$pattern)
### automatically extend PWM model
pwm.match.extend <-refinePWMMotifExtend(ctcf.motifs$motifs[[1]]@match$pattern, ctcf.seq)
### plot the new motif matches
plotMotif(pwm.match.extend$match$pattern)
```

**refinePWMMotifExtend** *create an extended PWM (Position Weight Model) model given a initial set of motif matches and input sequences*

**Description**

Create an extended PWM model given a initial set of motif matches and input sequences

**Usage**

```
refinePWMMotifExtend(motifs=NULL, seqs, pwm.ld=NULL, flank=3, extend.tol=10^-3, trim.rel.entropy=0)
```

**Arguments**

<b>motifs</b>	The initial set of motif matches. character vector or DNAStringSet object
<b>seqs</b>	Input sequences. character vector or DNAStringSet object
<b>pwm.ld</b>	The initial PWM matrixes in logodds transformation. Either "motifs" or "pwm.ld" is not NULL
<b>flank</b>	The number of bases for extension on both sides of the motif. The extension will be iterated if the there is sufficient signal in the flanking region.
<b>extend.tol</b>	Convergence criteria for extension.
<b>trim.rel.entropy</b>	cutoff to be used to trim the uninformative flanking of a PWM model based on relative entropy against a null distribution.
<b>null</b>	NULL background distribution
<b>max.width</b>	The maximum width of PWM
<b>...</b>	other arguments passed to function refinePWMMotif

## Details

Flanking regions with length equal to flank is still included in output for reference

## Value

Same type of object returned by [refinePWMMotif](#)

## See Also

[findMotif](#) [refinePWMMotif](#)

**summaryMotif**

*summarize a list of motifs*

## Description

Create a summary table of a list of motifs found by [findMotif](#)

## Usage

`summaryMotif(motifs, category)`

## Arguments

<code>motifs</code>	a list of motifs of class <a href="#">Motif-class</a>
<code>category</code>	a vector of 0 or 1, specifying which sequences are foreground and background. Input for <a href="#">findMotif</a>

## Value

A data.frame with following columns:

<code>scores</code>	scores for each Motif. All values are negative. The absolute scales of the scores reflect the discriminative power of the motif for separating the foreground and background. Statistically, they correspond to the Z-values of the predictor(counts of the motifs in this case) in the logistic regression model
<code>signs</code>	sign of the motifs. TRUE for enriched motifs, FALSE for depleted motifs
<code>fg.hits, bg.hits</code>	Total number of hits in the foreground, and background sequences. If the motif is scanned on both strands of the input sequences, the counts on both strands are added.
<code>fg.seq, bg.seq</code>	The number of sequences that contain at least one motif match in the foreground, and the background
.	
<code>ratio</code>	The enrichment/depleted ratio of motifs
<code>fg.frac, bg.frac</code>	The fraction of foreground/background sequences that contain at least one motif match

**See Also**

[findMotif](#) [motifLatexTable](#) [motifHtmlTable](#)

**Examples**

```
data(ctcf.motifs)
###plot the summary statics of motif matches after masking previous motif occurrences###
summaryMotif(ctcf.motifs$mask.motifs, ctcf.motifs$category)

###plot the summary statics of motif matches in the original sequences###
summaryMotif(ctcf.motifs$motifs, ctcf.motifs$category)
```

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YY1.control

*Background for YY1 ChIP-Seq peaks in HepG2*

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**Description**

Background regions randomly sampled in the flanking region of YY1 ChIP-Seq peaks in HepG2.

**References**

Unpublished

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YY1.peak

*YY1 ChIP-Seq peaks in HepG2*

---

**Description**

a subset of 5000 YY1 ChIP-Seq peaks in HepG2 from ENCODE

**References**

Unpublished

# Index

\*Topic **datasets**

  ctcf.motifs, 2

  YY1.control, 11

  YY1.peak, 11

class:Motif (Motif-class), 5

  ctcf.motifs, 2

  DNAStringSet, 7

  findMotif, 2, 2, 5, 6, 9–11

  findMotifFasta, 4

  findMotifFgBg, 4

  getSequence, 5

  Motif-class, 2, 3, 5, 10

  motifHtmlTable, 11

  motifHtmlTable (motifLatexTable), 6

  motifLatexTable, 5, 6, 11

  plotMotif, 5, 7

  refinePWMMotif, 8, 10

  refinePWMMotifExtend, 9, 9

  summaryMotif, 5, 10

YY1.control, 11

YY1.peak, 11