## Package ‘motifTestR’

**May 30, 2024**

**Title**  Perform key tests for binding motifs in sequence data

**Version**  1.0.2

**Description**  Taking a set of sequence motifs as PWMs, test a set of sequences for over-representation of these motifs, as well as any positional features within the set of motifs. Enrichment analysis can be undertaken using multiple statistical approaches. The package also contains core functions to prepare data for analysis, and to visualise results.

**License**  GPL-3

**Encoding**  UTF-8

**URL**  [https://github.com/smped/motifTestR](https://github.com/smped/motifTestR)

**BugReports**  [https://github.com/smped/motifTestR/issues](https://github.com/smped/motifTestR/issues)

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motifTestR-package

motifTestR: Perform Key Analyses on Transcription Factor Binding Motifs

Description

The package motifTestR has been designed for two primary analyses of TFBMs, testing for positional bias and overall enrichment.

Details

The package motifTestR provides two primary functions for testing TFBMs within a set of sequences

• `testMotifPos()` for detecting positional bias within a set of test sequences

• `testMotifEnrich()` for testing overall enrichment of a TFBM within a set of test sequences

The main functions rely on lower-level functions such as:

• `countPwmMatches()` simply counts the number of matches within an XStringSet

• `getPwmMatches()` returns the position of matches within an XStringSet

• `makeRMRanges()` which produces a set of random, matching ranges based on key characteristics of the set of test sequences/ranges

A simple utility function is provided to enable visualisation of results

• `plotMatchPos()` enables visualisation of the matches within a set of sequences using multiple strategies
ar_er_peaks

Author(s)
Stevie Pederson

See Also
Useful links:
• https://github.com/smped/motifTestR
• Report bugs at https://github.com/smped/motifTestR/issues

ar_er_peaks  A set of peaks with AR and ER detected

Description
A set of ChIP-Seq peaks where AR and ER were both detected

Usage
data("ar_er_peaks")

Format
An object of class GRanges of length 229.

Details
The subset of peaks found on chr1, and which contained signal from AR and ER, along with H3K27ac signal were taken from GSE123767. Peaks were resized to a uniform width of 400bp after downloading

Generation of these ranges is documented in system.file("scripts/ar_er_peaks.R", package = "motifTestR")

Source

Examples
data("ar_er_peaks")
ar_er_peaks
ar_er_seq  Sequences from peaks with AR and ER detected

Description
The genomic sequences obtained from the ar_er_peaks

Usage
data("ar_er_seq")

Format
An object of class DNAStringSet of length 229.

Details
These sequences represent the sequences obtained from BSgenome.Hsapiens.UCSC.hg19 for the peaks supplied as ar_er_peaks
Generation of these sequences is documented in system.file("scripts/ar_er_peaks.R", package = "motifTestR")

Examples
data("ar_er_seq")
ar_er_seq

countPwmMatches  Count the matches to a PWM within an XStringSet

Description
Count the matches to a PWM within an XStringSet

Usage
countPwmMatches(
pwm,
stringset,
rc = TRUE,
min_score = "80%",
mc.cores = 1,
...
)
Arguments

- **pwm**: A Position Weight Matrix
- **stringset**: An XStringSet
- **rc**: logical(1) Also find matches using the reverse complement of pwm
- **min_score**: The minimum score to return a match
- **mc.cores**: Passed to `mclapply` when analysing a list of PWMs

Details

Will simply count the matches within an XStringSet and return an integer. All matches are included.

Value

An integer vector

Examples

```r
## Load the example PWM
data("ex_pwm")
esr1 <- ex_pwm$ESR1

## Load the example Peaks
data("ar_er_seq")
countPwmMatches(esr1, ar_er_seq)

## Count all PWMs
countPwmMatches(ex_pwm, ar_er_seq)
```

---

### ex_pwm  

**Example Position Weight Matrices**

Description

Example Position Weight Matrices

Usage

```r
data("ex_pwm")
```

Format

An object of class `list` of length 5.
Details

This object contains 5 PWMs taken from HOOCOMOCOv11-coreA for examples and testing.

Generation of this motif list is documented in `system.file("scripts/ex_pwm.R", package = "motifTestR")`

Examples

```r
data("ex_pwm")
ex_pwm$ESR1
```

---

**getPwmMatches**  
*Find all PWM matches within an XStringSet*

Description

Find all PWM matches within a set of sequences

Usage

```r
getPwmMatches(
  pwm,  
  stringset,  
  rc = TRUE,  
  min_score = "80%",  
  best_only = FALSE,  
  break_ties = c("all", "random", "first", "last", "central"),  
  mc.cores = 1,  
  ...  
)
```

Arguments

- **pwm**: A Position Weight Matrix, list of PWMs or universalmotif list
- **stringset**: An XStringSet
- **rc**: logical(1) Also find matches using the reverse complement of pwm
- **min_score**: The minimum score to return a match
- **best_only**: logical(1) Only return the best match
- **break_ties**: Method for breaking ties when only returning the best match Ignored when all matches are returned (the default)
- **mc.cores**: Passed to `mclapply` if passing multiple PWMs
- **...**: Passed to `matchPWM`
Details

Taking a set of sequences as an XStringSet, find all matches above the supplied score (i.e. threshold) for a single Position Weight Matrix (PWM), generally representing a transcription factor binding motif. By default, matches are performed using the PWM as provided and the reverse complement, however this can easily be disable by setting `rc = FALSE`.

The function relies heavily on `matchPWM` and `Views` for speed.

When choosing to return the best match (`best_only = TRUE`), only the match with the highest score is returned for each sequence. Should there be tied scores, the best match can be chosen as either the first, last, most central, all tied matches, or choosing one at random (the default).

Value

A DataFrame with columns: `seq`, `score`, `direction`, `start`, `end`, `fromCentre`, `seq_width`, and `match`

The first three columns describe the sequence with matches, the score of the match and whether the match was found using the forward or reverse PWM. The columns `start`, `end`, and `width` describe the where the match was found in the sequence, whilst `from_centre` defines the distance between the centre of the match and the centre of the sequence being queried. The final column contains the matching fragment of the sequence as an XStringSet.

When passing a list of PWMs, a list of the above DataFrames will be returned.

Examples

```r
## Load the example PWM
data("ex_pwm")
esr1 <- ex_pwm$ESR1

## Load the example Peaks
data("ar_er_seq")

## Return all matches
getPwmMatches(esr1, ar_er_seq)

## Just the best match
getPwmMatches(esr1, ar_er_seq, best_only = TRUE)

## Apply multiple PWMs as a list
getPwmMatches(ex_pwm, ar_er_seq, best_only = TRUE)
```

## Regions from hg19 with high N content

A GRanges object with regions annotated as telomeres or centromeres
makeRMRanges

Usage

data("hg19_mask")

Format

An object of class GRanges of length 345.

Details

The regions defined as centromeres or telomeres in hg19, taken from AnnotationHub objects "AH107360" and "AH107361". These were combined with regions containing Ns from the UCSC 2bit file, and regions with Ns in the BSgenome.Hsapiens.UCSC.hg19 were retained.

Generation of these ranges is documented in system.file("scripts/hg19_mask.R", package = "motifTestR")

Source

The package AnnotationHub and https://hgdownload.cse.ucsc.edu/goldenpath/hg19/bigZips/hg19.fa.masked.gz

Examples

data("hg19_mask")
hg19_mask

makeRMRanges Form a set of random, matching ranges for bootstrapping or permuting

Description

Form a set of ranges from y which (near) exactly match those in x for use as a background set requiring matching

Usage

makeRMRanges(x, y, ...)

## S4 method for signature 'GRanges,GRanges'
makeRMRanges(
x, y,
exclude = GRanges(),
n_iter = 1,
n_total = NULL,
replace = TRUE,
...,
makeRMRanges

```
makeRMRanges(
  x,
  y,
  exclude = GRanges(),
  n_iter = 1,
  n_total = NULL,
  replace = TRUE,
  mc.cores = 1,
  ...,
  force_ol = TRUE,
  unlist = TRUE
)
```

Arguments

- **x**: GRanges/GRangesList with ranges to be matched
- **y**: GRanges/GRangesList with ranges to select random matching ranges from
- **...**: Not used
- **exclude**: GRanges of ranges to omit from testing
- **n_iter**: The number of times to repeat the random selection process
- **n_total**: Setting this value will over-ride anything set using n_iter. Can be vector of any length, corresponding to the length of x, when x is a GRangesList
- **replace**: logical(1) Sample with our without replacement when creating the set of random ranges.
- **force_ol**: logical(1) Enforce an overlap between every site in x and y
- **mc.cores**: Passed to mclapply
- **unlist**: logical(1) Return as a sorted GRanges object, or leave as a GRangesList

Details

This function uses the width distribution of the 'test' ranges (i.e. x) to randomly sample a set of ranges with matching width from the ranges provided in y. The width distribution will clearly be exact when a set of fixed-width ranges is passed to x, whilst random sampling may yield some variability when matching ranges of variable width.

When both x and y are GRanges objects, they are implicitly assumed to both represent similar ranges, such as those overlapping a promoter or enhancer. When passing two GRangesList objects, both objects are expected to contain ranges annotated as belonging to key features, such that the list elements in y must encompass all elements in x. For example if x contains two elements named 'promoter' and 'intron', y should also contain elements named 'promoter' and 'intron' and these will be sampled as matching ranges for the same element in x. If elements of x and y are not named, they are assumed to be in matching order.
The default behaviour is to assume that randomly-generated ranges are for iteration, and as such, ranges are randomly formed in multiples of the number of 'test' ranges provided in \( x \). The column \textit{iteration} will be added to the returned ranges. Placing any number into the \textit{n_total} argument will instead select a total number of ranges as specified here. In this case, no \textit{iteration} column will be included in the returned ranges.

Sampling is assumed to be with replacement as this is most suitable for bootstrapping and related procedures, although this can be disabled by setting \textit{replace} = \texttt{FALSE}.

**Value**

A GRanges or GRangesList object

**Examples**

```r
## Load the example peaks
data("ar_er_peaks")
sq <- seqinfo(ar_er_peaks)
## Now sample size-matched ranges for two iterations from chr1
makeRMRanges(ar_er_peaks, GRanges(sq)[1], n_iter = 2)
## Or simply sample 100 ranges if not planning any iterative analyses
makeRMRanges(ar_er_peaks, GRanges(sq)[1], n_total = 100)
```

---

**plotMatchPos**  
*Plot Motif Match Positions*

**Description**

Plot the distribution of motif matches across sequences

**Usage**

```r
plotMatchPos(
    matches,  
    binwidth = 10,  
    abs = FALSE,  
    use_totals = FALSE,  
    type = c("density", "cdf", "heatmap"),  
    geom = c("smooth", "line", "point", "col"),  
    cluster = FALSE,  
    w = 0.1,  
    heat_fill = NULL,  
    ...  
)
```
plotMatchPos

Arguments

matches Output from `getPwmMatches`
binwidth Width of bins to use when plotting
abs logical(1) Plot absolute distances from centre
use_totals logical(1). If TRUE, plots will use total counts. The default (FALSE) plots probabilities.
type Plot match density, the CDF or a binned heatmap
geom Type of geom to be used for line plots. Ignored for heatmaps
cluster logical(1) Cluster motifs when drawing a heatmap. If TRUE a dendrogram will be added to the LHS of the plot
w Relative width of the dendrogram on (0, 1)
heat_fill scale_fill_continuous object for heatmaps. If not provided, `scale_fill_viridis_c()` will be added to the heatmap.
... Passed to individual geom* functions

Details

Multiple options are provided for showing the distribution of PWM matches within a set of sequences, using either the smoothed probability density, the probability CDF or as a heatmap. Distances can be shown as symmetrical around the centre or using absolute distances from the central position within the sequences.

Heatmaps are only enabled for comparisons across multiple PWMs, with optional clustering enabled. If adding a dendrogram for clustering, the returned plot object will be a patchwork object.

Value

A ggplot2 object

Examples

```r
## Load the example PWM
data("ex_pwm")
esr1 <- ex_pwm$ESR1

## Load the example sequences from the peaks
data("ar_er_seq")

## Just the best match
bm <- getPwmMatches(esr1, ar_er_seq, best_only = TRUE)
plotMatchPos(bm, se = FALSE)

## Matches can also be shown by distance from centre
plotMatchPos(bm, abs = TRUE)

## Cumulative Probability plots are also implemented
plotMatchPos(bm, type = "cdf", geom = "line", colour = "red") + geom_abline(intercept = 0.5, slope = 1/ 400)
```
testMotifEnrich

Test motif enrichment using a background set of sequences

Description

Test for motif enrichment within a set of sequences using a background set to derive a NULL hypothesis

Usage

```r
testMotifEnrich(
  pwm,  
  stringset,  
  bg,  
  var = "iteration",  
  model = c("quasipoisson", "hypergeometric", "poisson", "iteration"),  
  sort_by = c("p", "none"),  
  mc.cores = 1,
  ...)
```

Arguments

- `pwm`: A Position Weight Matrix or list of PWMs
- `stringset`: An XStringSet with equal sequence widths
- `bg`: An XStringSet with the same sequence widths as the test XStringset
- `var`: A column in the mcols element of bg, usually denoting an iteration number
- `model`: The model used for analysis
- `sort_by`: Column to sort results by
- `mc.cores`: Passed to mclapply
- `...`: Passed to getPwmMatches or countPwmMatches

Details

This function offers four alternative models for assessing the enrichment of a motif within a set of sequences, in comparison to a background set of sequences. Selection of the BG sequences plays an important role and, in conjunction with the question being addressed, determines the most appropriate model to use for testing, as described below.

It should also be noted that the larger the BG set of sequences, the larger the computational burden, and results can take far longer to return. For many millions of background sequences, this may run beyond an hour.

Descriptions of Models and Use Cases:
Hypergeometric Tests:

Hypergeometric tests are best suited to the use case where the test set of sequences represents a subset of a larger set, with a specific feature or behaviour, whilst the BG set may be the remainder of the set without that feature. For example, the test set may represent ChIP-Seq binding sites where signal changes in response to treatment, whilst the BG set represents the sites where no changed signal was observed. Testing is one-sided, for enrichment of motifs within the test set.

Due to these relatively smaller sized datasets, setting model = "hypergeometric", will generally return results quickly.

Poisson Tests:

This approach requires a set of background sequences which should be much larger than the test set of sequences. The parameters for a Poisson model are estimated in a per-sequence manner on the set of BG sequences, and the observed rate of motif-matches within the test set is then tested using poisson.test.

This approach assumes that all matches follow a Poisson distribution, which is often true, but data can also be overdispersed. Given that this model can also return results relatively quickly, is it primarily suitable for data exploration, but not for final results.

Quasi-Poisson Test:

The quasipoisson model allows for over-dispersion and will return more conservative results than using the standard Poisson model. Under the method currently implemented here, BG sequences should be divided into blocks (i.e. iterations), identical in size to the test set of sequences. Model parameters are estimated per iteration across the BG set of sequences, with the rate of matches in the test being compared against these.

It is expected that the BG set will matched for the features of interest and chosen using makeRMRanges with a large number of iterations, e.g. n_iter = 1000. Due to this parameterisation, quasipoisson approaches can be computationally time-consuming, as this is effectively an iterative approach.

Iteration:

Setting the model as "iteration" performs a non-parametric analysis, with the exception of returning Z-scores under the Central Limit Theorem. Mean and SD of matches is found for each iteration, and used to return Z scores, with p-values returned from both a Z-test and from comparing observed values directly to sampled values obtained from the BG sequences. Sampled values are calculated directly and as such, are limited in precision.

As for the QuasiPoisson model, a very large number of iterations is expected to be used, to ensure the CLT holds, again making this a computationally demanding test. Each iteration/block is expected to be identically-sized to the test set, and matched for any features as appropriate using makeRMRanges().

Value

A data.frame with columns: sequences, matches, expected, enrichment, and p, with additional columns Z, est_bg_rate (Poisson), odds_ratio (Hypergeometric) or Z, sd_bg, n_iter and iter_p (Iterations). The numbers of sequences and matches refer to the test set of sequences, whilst expected is the expected number of matches under the Poisson or iterative null distribution. The ratio of matches to expected is given as enrichment, along with the Z score and p-value. Whilst the Z score is only representative under the Poisson model, it is used to directly estimate p-values under the iterative approach. Under this latter approach, the sd of the matches found in the background
sequences is also given, along with the number of iterations and the p-values from permutations testing the one-sided hypothesis hypothesis for enrichment.

It may also be worth noting that if producing background sequences using makeRMRanges with replace = TRUE and force.ol = TRUE, the iterative model corresponds to a bootstrap, given that the test sequences will overlap the background sequences and background ranges are able to be sampled with replacement.

See Also
makeRMRanges(), getPwmMatches(), countPwmMatches()

Examples

```r
## Load the example peaks & the sequences
data("ar_er_peaks")
data("ar_er_seq")
sq <- seqinfo(ar_er_peaks)
## Now sample size-matched ranges 10 times larger. In real-world analyses,
## this set should be sampled as at least 1000x larger, ensuring features
## are matched to your requirements. This example masks regions with known N
## content, including centromeres & telomeres
data("hg19_mask")
set.seed(305)
bg_ranges <- makeRMRanges(
  ar_er_peaks, GRanges(sq)[1], exclude = hg19_mask, n_iter = 10
)

## Convert ranges to DNAStringSets
library(BSgenome.Hsapiens.UCSC.hg19)
genome <- BSgenome.Hsapiens.UCSC.hg19
bg_seq <- getSeq(genome, bg_ranges)

## Test for enrichment of the ESR1 motif
data("ex_pwm")
esr1 <- ex_pwm$ESR1
testMotifEnrich(esr1, ar_er_seq, bg_seq, model = "poisson")

## Test all motifs
testMotifEnrich(ex_pwm, ar_er_seq, bg_seq, model = "poisson")
```

---

**testMotifPos**

Test for a Uniform Distribution across a set of best matches

**Description**

Test for a Uniform Distribution across a set of best matches
Usage

testMotifPos(
  x,
  stringset,
  binwidth = 10,
  abs = FALSE,
  rc = TRUE,
  min_score = "80%",
  break_ties = "all",
  alt = c("greater", "less", "two.sided"),
  sort_by = c("p", "none"),
  mc.cores = 1,
  ...
)

Arguments

x A Position Weight Matrix, universalmotif object or list thereof. Alternatively can be a single DataFrame or list of DataFrames as returned by getPwmMatches with best_only = TRUE

stringset An XStringSet. Not required if matches are supplied as x

binwidth Width of bins across the range to group data into

abs Use absolute positions around zero to find symmetrical enrichment

rc logical(1) Also find matches using the reverse complement of pwm

min_score The minimum score to return a match

break_ties Choose how to resolve matches with tied scores

alt Alternative hypothesis for the binomial test

sort_by Column to sort results by

mc.cores Passed to mclapply

... Passed to matchPWM

Details

This function tests for an even positional spread of motif matches across a set of sequences, using the assumption (i.e. H~0~) that if there is no positional bias, matches will be evenly distributed across all positions within a set of sequences. Conversely, if there is positional bias, typically but not necessarily near the centre of a range, this function intends to detect this signal, as a rejection of the null hypothesis.

Input can be provided as the output from getPwmMatches setting best_only = TRUE if these matches have already been identified. If choosing to provide this object to the argument matches, nothing is required for the arguments pwm, stringset, rc, min_score or break_ties Otherwise, a Position Weight Matrix (PWM) and an XStringSet are required, along with the relevant arguments, with best matches identified within the function.

The set of best matches are then grouped into bins along the range, with the central bin containing zero, and tallied. Setting abs to TRUE will set all positions from the centre as absolute values,
returning counts purely as bins with distances from zero, marking this as an inclusive lower bound. Motif alignments are assigned into bins based on the central position of the match, as provided in the column `from_centre` when calling `getPwmMatches`.

The `binom.test` is performed on each bin using the alternative hypothesis, with the returned p-values across all bins combined using the Harmonic Mean p-value (HMP) (See `p.hmp`). All bins with raw p-values below the HMP are identified and the returned values for start, end, centre, width, matches in region, expected and enrichment are across this set of bins. The expectation is that where a positional bias is evident, this will be a narrow range containing a non-trivial proportion of the total matches.

It should also be noted that `binom.test()` can return p-values of zero, as beyond machine precision. In these instances, zero p-values are excluded from calculation of the HMP. This will give a very slight conservative bias, and assumes that for these extreme cases, neighbouring bins are highly likely to also return extremely low p-values and no significance will be lost.

**Value**

A data frame with columns `start`, `end`, `centre`, `width`, `total_matches`, `matches_in_region`, `expected`, `enrichment`, `prop_total`, `p` and `consensus_motif`. The total matches represent the total number of matches within the set of sequences, whilst the number observed in the final region are also given, along with the proportion of the total this represents. Enrichment is simply the ratio of observed to expected based on the expectation of the null hypothesis.

The consensus motif across all matches is returned as a Position Frequency Matrix (PFM) using `consensusMatrix`.

**Examples**

```r
## Load the example PWM
data("ex_pwm")
esr1 <- ex_pwm$ESR1

## Load the example sequences
data("ar_er_seq")

## Get the best match and use this data
matches <- getPwmMatches(esr1, ar_er_seq, best_only = TRUE)

## Test for enrichment in any position
testMotifPos(matches)

## Provide a list of PWMs, testing for distance from zero
testMotifPos(ex_pwm, ar_er_seq, abs = TRUE, binwidth = 10)
```

---

**Candidate Enhancer Regions from ZR-75-1 Cells**
**Description**

The chr1 subset of candidate enhancers for ZR-75-1 cells

**Usage**

```r
data("zr75_enh")
```

**Format**

An object of class GRanges of length 5237.

**Details**

These enhancers are the chr1 subset of enhancer regions for ZR-75-1 cells as identified by EnhancerAtlas 2.0

# Generation of these ranges is documented in `system.file("scripts/zr75_enh.R", package = "motifTestR")`

**Source**


**Examples**

```r
data("zr75_enh")
zr75_enh
```
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