Package 'motifbreakR'

November 17, 2025

Title A Package For Predicting The Disruptiveness Of Single Nucleotide Polymorphisms On Transcription Factor Binding Sites

Version 2.24.0

Description We introduce motifbreakR, which allows the biologist to judge in the first place whether the sequence surrounding the polymorphism is a good match, and in the second place how much information is gained or lost in one allele of the polymorphism relative to another. MotifbreakR is both flexible and extensible over previous offerings; giving a choice of algorithms for interrogation of genomes with motifs from public sources that users can choose from; these are 1) a weighted-sum probability matrix, 2) log-probabilities, and 3) weighted by relative entropy. MotifbreakR can predict effects for novel or previously described variants in public databases, making it suitable for tasks beyond the scope of its original design.

Lastly, it can be used to interrogate any genome curated within Bioconductor (currently there are 32 species, a total of 109 versions).

Depends R (>= 4.4.0), grid, MotifDb

Imports methods, grDevices, stringr, parallel, BiocGenerics, S4Vectors (>= 0.9.25), IRanges, GenomeInfoDb, GenomicRanges, Biostrings, BSgenome, rtracklayer, VariantAnnotation, BiocParallel, motifStack, Gviz, matrixStats, TFMPvalue, SummarizedExperiment, pwalign, DT, bsicons, BiocFileCache, biomaRt, bslib, shiny, vroom

Suggests BSgenome. Hsapiens. UCSC. hg19,

SNPlocs. Hsapiens.dbSNP155.GRCh37, knitr, rmarkdown,

BSgenome.Drerio.UCSC.danRer7, BiocStyle,

BSgenome. Hsapiens. 1000 genomes. hs37d5,

BSgenome. Hsapiens. UCSC. hg19. masked,

BSgenome. Hsapiens. NCBI. GRCh38,

BSgenome. Hsapiens. UCSC. hg38. masked, BSgenome. Hsapiens. UCSC. hg38

VignetteBuilder knitr

Encoding UTF-8

License GPL-2

BugReports https://github.com/Simon-Coetzee/motifbreakR/issues

biocViews ChIPSeq, Visualization, MotifAnnotation, Transcription

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calculatePvalue

Calculate the significance of the matches for the reference and alternate alleles for the for their PWM

Description

Calculate the significance of the matches for the reference and alternate alleles for the for their PWM

Usage

```
calculatePvalue(
  results,
  background = c(A = 0.25, C = 0.25, G = 0.25, T = 0.25),
  granularity = NULL,
  BPPARAM = BiocParallel::SerialParam()
)
```

encodemotif 3

Arguments

results The output of motifbreakR that was run with filterp=TRUE

background Numeric Vector; the background probabilities of the nucleotides

granularity Numeric Vector; the granularity to which to round the PWM, larger values compromise full accuracy for speed of calculation. A value of NULL does no rounding.

BPPARAM a BiocParallel object see register and see getClass("BiocParallelParam")

for additional parameter classes. Try BiocParallel::registered() to see what's available and for example BiocParallel::bpparam("SerialParam") would allow sorial avaluation.

would allow serial evaluation.

Details

This function is intended to be used on a selection of results produced by motifbreakR, and this can be (although not always) a very memory and time intensive process if the algorithm doesn't converge rapidly.

Value

a GRanges object. The same Granges object that was input as results, but with Refpvalue and Altpvalue columns in the output modified from NA to the p-value calculated by TFMsc2pv. Additionally a pvalueEffect column that indicates "strong" when the lower p-value (between ref and alt) is an order of magnitude or more different from the higher p-value, otherwise weak.

Source

H\'el\'ene Touzet and Jean-St\'ephane Varr\'e (2007) Efficient and accurate P-value computation for Position Weight Matrices. Algorithms for Molecular Biology, **2: 15**.

See Also

See TFMsc2pv from the TFMPvalue package for information about how the p-values are calculated.

Examples

```
data(example.results)
rs1006140 <- example.results[example.results$SNP_id %in% "rs1006140"]
# low granularity for speed; 1e-6 or 1e-7 recommended for accuracy
rs1006140 <- calculatePvalue(rs1006140, BPPARAM=BiocParallel::SerialParam(), granularity = 1e-4)</pre>
```

encodemotif MotifDb object containing motif information from the known and discovered motifs for the ENCODE TF ChIP-seq datasets.

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Description

From the abstract: "Recent advances in technology have led to a dramatic increase in the number of available transcription factor ChIP-seq and ChIP-chip data sets. Understanding the motif content of these data sets is an important step in understanding the underlying mechanisms of regulation. Here we provide a systematic motif analysis for 427 human ChIP-seq data sets using motifs curated from the literature and also discovered de novo using five established motif discovery tools. We use a systematic pipeline for calculating motif enrichment in each data set, providing a principled way for choosing between motif variants found in the literature and for flagging potentially problematic data sets. Our analysis confirms the known specificity of 41 of the 56 analyzed factor groups and reveals motifs of potential cofactors. We also use cell type-specific binding to find factors active in specific conditions. The resource we provide is accessible both for browsing a small number of factors and for performing large-scale systematic analyses. We provide motif matrices, instances and enrichments in each of the ENCODE data sets. The motifs discovered here have been used in parallel studies to validate the specificity of antibodies, understand cooperativity between data sets and measure the variation of motif binding across individuals and species."

Usage

encodemotif

Format

```
MotifDb object of length 2064; to access metadata use mcols(encodemotif)
providerName Name provided by ENCODE
providerId Same as providerName
dataSource "ENCODE-motif"
geneSymbol Gene symbol for the transcription factor
geneId Entrez gene id for the transcription factor
geneIdType "ENTREZ"
proteinId UNIPROT id for the transcription factor
proteinIdType "UNIPROT"
organism "Hsapiens"
sequenceCount NA not available
bindingSequence Consensus sequence for the motif
bindingDomain NA incomplete
tfFamily NA incomplete
experimentType occurs in two forms:
     For motifs that were discovered in this study, the format is cellType_source-LabMetadata: MotifFinder#Locat:
     for example H1-hESC_encode-Myers_seq_hsa_v041610.2_r1:MEME#2#Intergenic.
     For motifs that were "known" the format tends to be TF_source_sourceId for example
     AP1_jaspar_MA0099.2.
pubmedID "24335146" see Source for more details
```

Details

Load with data(encodemotif)

example.results 5

Value

```
MotifList-class object
```

Source

Pouya Kheradpour and Manolis Kellis (2013 December 13) Systematic discovery and characterization of regulatory motifs in ENCODE TF binding experiments. Nucleic Acids Research, doi:10.1093/nar/gkt1249

See Also

```
http://compbio.mit.edu/encode-motifs/
```

Examples

```
data(encodemotif)
encodemotif
```

example.results

Example Results from motifbreakR

Description

This contains example results from motifbreaker for use in examples from the help docs

Usage

```
example.results
```

Format

GRanges output from motifbreakR

Value

GRanges object. See motifbreakR for information on it's structure.

Examples

```
data(example.results)
example.results
```

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expo	rtM	IBD	ea

Export motifbreakR variants to bed file

Description

Export motifbreakR variants to bed file

Usage

```
exportMBbed(results, file, name = NULL, color = "effect_size")
```

Arguments

results The output of motifbreakR

file Character; the file name of the destination file

name Character; name for the BED track, defaults to "motifbreakR results"

color Character; one of ref_sig (Refpvalue), alt_sig (Altpvalue), best_sig (lowest

between Refpvalue and Altpvalue), (each of which require pre-computation of p-values with calculatePvalue), or ref_score (pctRef), alt_score (pctAlt), best_score (highest between pctRef and pctAlt), or the default value of ef-

fect_size (alleleDiff).

Value

exportMBbed produces an output BED file, with diverging color scale for effect_size (blue representing stronger binding in REF, red representing stronger binding in ALT), or a sequential color scale otherwise (low values as purple, high values as yellow). The score column is either the effect_size (alleleDiff column), the -log10(p-value) (capped at 10), corresponding to Refpvalue, Altpvalue, or the best match of the two, or the score pctRef, pctAlt, or the highest match of the two. The name column is formatted SNP_id:REF/ALT:providerId. Additionally a color key is returned indicating the range of values for each color output.

See Also

See exportMBtable for the function that exports the full motifbreakR results as a tab or comma separated table file.

Examples

```
data(example.results)
example.results
exportMBbed(example.results, file = "mb_test_output.bed", color = "effect_size")
```

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exportMBtable Export motifbreakR results to csv or tsv	exportMBtable	Export motifbreakR results to csv or tsv	
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Description

Export motifbreakR results to csv or tsv

Usage

```
exportMBtable(results, file, format = "tsv")
```

Arguments

results The output of motifbreakR

file Character; the file name of the destination file

format Character; one of tsv (tab separated values) or csv (comma separated values)

Value

exportMBtable produces an output file containing the output of the motifbreakR function.

See Also

See exportMBbed for the function that exports the motifbreakR results as a BED file, colored by selected score.

Examples

```
data(example.results)
example.results
exportMBtable(example.results, file = "mb_test_output.tsv", format = "tsv")
```

factorbook MotifDb object containing motif information from around the genomic regions bound by 119 human transcription factors in Factorbook.

Description

From the abstract: "Chromatin immunoprecipitation coupled with high-throughput sequencing (ChIP-seq) has become the dominant technique for mapping transcription factor (TF) binding regions genome-wide. We performed an integrative analysis centered around 457 ChIP-seq data sets on 119 human TFs generated by the ENCODE Consortium. We identified highly enriched sequence motifs in most data sets, revealing new motifs and validating known ones. The motif sites (TF binding sites) are highly conserved evolutionarily and show distinct footprints upon DNase I digestion. We frequently detected secondary motifs in addition to the canonical motifs of the TFs, indicating tethered binding and cobinding between multiple TFs. We observed significant position and orientation preferences between many cobinding TFs. Genes specifically expressed in a cell line are often associated with a greater occurrence of nearby TF binding in that cell line. We observed

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cell-line-specific secondary motifs that mediate the binding of the histone deacetylase HDAC2 and the enhancer-binding protein EP300. TF binding sites are located in GC-rich, nucleosome-depleted, and DNase I sensitive regions, flanked by well-positioned nucleosomes, and many of these features show cell type specificity. The GC-richness may be beneficial for regulating TF binding because, when unoccupied by a TF, these regions are occupied by nucleosomes in vivo. We present the results of our analysis in a TF-centric web repository Factorbook (http://factorbook.org) and will continually update this repository as more ENCODE data are generated."

Usage

factorbook

Format

```
MotifDb object of length 79; to access metadata use mcols(factorbook)
```

```
providerName Name listed in meme output of 'Supp TableS2.pdf' for the citation indicated below providerId Same as providerName
dataSource "FactorBook"
geneSymbol NA these motifs don't have a direct 1 to 1 relationship with a transcription factor geneId NA
geneIdType NA
proteinId NA
proteinIdType NA
organism "Hsapiens"
sequenceCount NA
bindingSequence Consensus sequence for the motif
bindingDomain NA
tfFamily NA
experimentType NA
pubmedID "22955990" see Source for more details
```

Details

Load with data(factorbook)

Value

MotifList-class object

Source

J Wang, J Zhuang, S Iyer, XY Lin, et al. (2012) Sequence features and chromatin structure around the genomic regions bound by 119 human transcription factors. Genome Research, **22** (**9**), 1798-1812, doi:10.1101/gr.139105.112

See Also

http://factorbook.org

Examples

data(factorbook)
factorbook

findSupportingRemapPeaks

Find Corresponding TF Binding From The ReMap2022 Project

Description

Find Corresponding TF Binding From The ReMap2022 Project

Usage

findSupportingRemapPeaks(results, genome, TFClass = FALSE)

Arguments

results The output of motifbreakR

genome Character; one of: hg38 or hg19 for Homo sapiens, mm10 or mm39 for Mus

musculus, dm6 for Drosophila melanogaster, TAIR10_TF or TAIR10_HISTONE

for Arabidopsis thaliana

TFClass Logical; The user may optionally query an expanded motif/transcription fac-

tor relationship encompassing the entire potential transcription factor family as

implemented by ${\tt MotifDb}$ based on TFClass.

Details

TFClass argument works for objects loaded in from the MotifDb package. hg19 and mm39 are data from liftOver.

The ReMap catalogues (2022, 2020, 2018, 2015) are under CC BY-NC 4.0 international license, as described in ReMap.

The CC BY-NC 4.0 license correspond to the following terms: Attribution — You must give appropriate credit, provide a link to the license, and indicate if changes were made. You may do so in any reasonable manner, but not in any way that suggests the licensor endorses you or your use. Non-Commercial — You may not use the material for commercial purposes. No additional restrictions — You may not apply legal terms or technological measures that legally restrict others from doing anything the license permits.

Value

the results GenomicRanges object output by motifbreakR containing with the additional columns: matchingBindingEvent

The name of the transcription factor that binds over the motif, or NA if none matchingCellType

A list corresponding in length to the number of transcription factors in matchingBindingEvent indicating the biotype/celltype that the transcription factor binding was found in.

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See Also

associateTranscriptionFactors for information about TFClass. https://remap.univ-amu.fr/ for details about ReMap2022.

Examples

hocomoco

MotifDb object containing motif information from Homo Sapiens Comprehensive Model Collection (HOCOMOCO) of transcription factor (TF) binding models

Description

From the abstract: "We present the Homo sapiens comprehensive model collection (HOCOMOCO, http://autosome.ru/HOCOMOCO/, http://cbrc.kaust.edu.sa/hocomoco/) containing carefully hand-curated TFBS models constructed by integration of binding sequences obtained by both lowand high-throughput methods. To construct position weight matrices to represent these TFBS models, we used ChIPMunk software in four computational modes, including newly developed periodic positional prior mode associated with DNA helix pitch. We selected only one TFBS model per TF, unless there was a clear experimental evidence for two rather distinct TFBS models. We assigned a quality rating to each model. HOCOMOCO contains 426 systematically curated TFBS models for 401 human TFs, where 172 models are based on more than one data source."

Usage

hocomoco

Format

```
MotifDb object of length 426; to access metadata use mcols(hocomoco)

providerName Name provided by HOCOMOCO

providerId ID provided by HOCOMOCO including experiment type
dataSource "HOCOMOCO"
geneSymbol Gene symbol for the transcription factor
geneId Entrez gene id for the transcription factor
geneIdType "ENTREZ"

proteinId UNIPROT id for the transcription factor
proteinIdType "UNIPROT"

organism "Hsapiens"

sequenceCount Number of sequences evaluated for producing the PWM
bindingSequence Consensus sequence for the motif
```

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bindingDomain NA incomplete

tfFamily NA incomplete

experimentType from http://autosome.ru/HOCOMOCO/Details.php#200 quoted here:

"TFBS model identification modes

To construct TFBS models ChIPMunk was run four times: two times (f1) and (f2) with uniform model positional prior and two times (si) and (do) with informative model positional prior.

The min-to-max (f1) model length estimation mode was used with the min length of 7 bp and increasing it by 1 bp until the default max length of 25 bp was reached following the optimal length selection procedure as in Kulakovskiy and Makeev, Biophysics, 2009. For max-to-min (f2) model length estimation mode we started from 25 bp and searched for the best alignment decreasing the length by 1 bp until the minimal length of 7 bp. We also used the single (si) and double box (do) model positional priors in order to simulate DNA helix turn. For a single box, the positional weights are to be distributed as $\cos 2(pi \text{ n} / \text{T})$, where T=10.5 is the DNA helix pitch, n is the coordinate within the alignment, and the center of the alignment of the length L is at n=0. During the internal cycle of PWM optimization the PWM column scores are multiplied by prior values so the columns closer to the center of the alignment (n=0) receive no score penalty while the columns around (n = 5,6,-5,-6) contribute much less to the score of the PWM under optimization. The single box model prior was used along with the min-to-max length estimation mode (si). We also used the double box model prior with a shape prior equal to $\sin 2(pin / T)$, which was used to search for possibly longer double box models in the max-to-min length estimation mode (do).

Model quality assignment

The resulting models were rated (from A to F) according to their quality. Model quality rates from A-to-D were assigned to proteins known to be TFs, including those listed in Schaefer et al., Nucleic Acids Research, 2011 with addition of a number of proteins having relevant models and sufficient evidence to be TFs. The ratings were assigned by human curation according to the following criteria:

Relevant distribution of position-specific information content over alignment columns, which means a model LOGO representation displaying well formed core positions with a high information content surrounded by flanking letters with lower information content; the information content at flanking positions decreasing with the distance from the model core.

"Stability", which means that in more than one of the ChIPMunk modes we obtained models with a similar length, consensus, and comparable number of aligned binding sites, along with a similar shape of model LOGO representation. "Similarity" of the model to the binding sequence consensus for this TF given in the UniProt or other databases, which means similarity of the shape of the model LOGO and TFBS lengths to those of other TFs from the same TF family. "A total number of binding sites" was also considered as a quality measure, as a large set of binding regions (mostly but not limited to ChIP-Seq and parallel SELEX) implies that there are many observations of each letter in any position of the alignment, particularly many observations of non-consensus letters in core positions. In positions with low information content, where there is no strong consensus, all variants have many observations, and thus the observed letter frequencies are less dependent on statistical fluctuations.

Quality A was assigned to high confidence models complying with all four criteria listed in the section above. Quality B was assigned to models built from large sequence sets that failed no more than one out of the three remaining criteria. Quality C was assigned to models built from small sequence sets but (with a number of specifically marked exceptions) complying with the three remaining criteria. Quality D models missed part of the known consensus sequence or had no clearly significant core positions in the TFBS model. Quality E (error) was assigned to models for proteins not convincingly shown to be TFs or to models exhibiting an irrelevant

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LOGO shape or a wrong consensus sequence. Quality F (failure) was assigned to TFs for which there was no reliable model identified."

pubmedID "23175603" see Source for more details

Details

Load with data(hocomoco)

Value

```
MotifList-class object
```

Source

Kulakovskiy, I.V., Medvedeva, Y.A., Schaefer, U., Kasianov, A.S., Vorontsov, I.E., Bajic, V.B. and Makeev, V.J. (2013) HOCOMOCO: a comprehensive collection of human transcription factor binding sites models. Nucleic Acids Research, 41, D195–D202.

See Also

```
http://autosome.ru/HOCOMOCO/http://cbrc.kaust.edu.sa/hocomoco/
```

Examples

data(hocomoco)
hocomoco

homer

MotifDb object containing motif information from motif databases included in HOMER.

Description

From the website: "Homer includes several motif databases that are used to help annotate results and conduct searches for known motifs. HOMER contains a custom motif database based on independent analysis of mostly ChIP-Seq data sets which is heavily utilized in the software." See http://homer.salk.edu/homer/motif/motifDatabase.html for more information on how these files were generated, and Homer's sources.

Usage

homer

Format

```
MotifDb object of length 247; to access metadata use mcols(homer)

providerName Name provided HOMER

providerId Factor Name provided by HOMER

dataSource "HOMER"

geneSymbol Symbol provided by HOMER
```

```
geneIdType "ENTREZ"

proteinId UNIPROT id for the transcription factor

proteinIdType "UNIPROT"

organism "Hsapiens"

sequenceCount NA

bindingSequence Consensus sequence for the motif

bindingDomain DBD provided by HOMER

tfFamily NA

experimentType The Celltype, IP, Assay, and GEO id if applicable for the motif

pubmedID "20513432" see Source for more details
```

Details

Load with data(homer)

Value

MotifList-class object

Source

Heinz S, Benner C, Spann N, Bertolino E et al. (2010 May 28) Simple Combinations of Lineage-Determining Transcription Factors Prime cis-Regulatory Elements Required for Macrophage and B Cell Identities. Mol Cell, **38(4):576-589**. PMID: 20513432

See Also

http://homer.salk.edu/homer/index.html http://homer.salk.edu/homer/motif/motifDatabase. html http://homer.salk.edu/homer/motif/HomerMotifDB/homerResults.html

Examples

data(homer)
homer

motifbreakR

Predict The Disruptiveness Of Single Nucleotide Polymorphisms On Transcription Factor Binding Sites.

Description

Predict The Disruptiveness Of Single Nucleotide Polymorphisms On Transcription Factor Binding Sites.

Usage

```
motifbreakR(
   snpList,
   pwmList,
   threshold = 0.85,
   filterp = FALSE,
   method = "default",
   show.neutral = FALSE,
   verbose = FALSE,
   bkg = c(A = 0.25, C = 0.25, G = 0.25, T = 0.25),
   BPPARAM = bpparam()
)
```

Arguments

snpList The output of snps.from.rsid or snps.from.file

pwmList An object of class MotifList containing the motifs that you wish to interrogate

threshold Numeric; the maximum p-value for a match to be called or a minimum score

threshold

filterp Logical; filter by p-value instead of by pct score.

method Character; one of default, log, ic, or notrans; see details.

show.neutral Logical; include neutral changes in the output

verbose Logical; if running serially, show verbose messages

bkg Numeric Vector; the background probabilities of the nucleotides used with method=log

method=ic

BPPARAM a BiocParallel object see register and see getClass("BiocParallelParam")

for additional parameter classes. Try BiocParallel::registered() to see what's availible and for example BiocParallel::bpparam("SerialParam")

would allow serial evaluation.

Details

motifbreakR works with position probability matrices (PPM). PPM are derived as the fractional occurrence of nucleotides A,C,G, and T at each position of a position frequency matrix (PFM). PFM are simply the tally of each nucleotide at each position across a set of aligned sequences. With a PPM, one can generate probabilities based on the genome, or more practically, create any number of position specific scoring matrices (PSSM) based on the principle that the PPM contains information about the likelihood of observing a particular nucleotide at a particular position of a true transcription factor binding site. What follows is a discussion of the three different algorithms that may be employed in calls to the **motifbreakR** function via the method argument.

Suppose we have a frequency matrix M of width n (i.e. a PPM as described above). Furthermore, we have a sequence s also of length n, such that $s_i \in \{A, T, C, G\}, i = 1, \dots n$. Each column of M contains the frequencies of each letter in each position.

Commonly in the literature sequences are scored as the sum of log probabilities:

Equation 1

$$F(s, M) = \sum_{i=1}^{n} \log(\frac{M_{s_i, i}}{b_{s_i}})$$

where b_{s_i} is the background frequency of letter s_i in the genome of interest. This method can be specified by the user as method='log'.

As an alternative to this method, we introduced a scoring method to directly weight the score by the importance of the position within the match sequence. This method of weighting is accessed by specifying method='ic' (information content). A general representation of this scoring method is given by:

Equation 2

$$F(s, M) = p_s \cdot \omega_M$$

where p_s is the scoring vector derived from sequence s and matrix M, and w_M is a weight vector derived from M. First, we compute the scoring vector of position scores p

Equation 3

$$p_s = (M_{s_i,i}) \quad \text{where} \quad \frac{i = 1, \dots n}{s_i \in \{A,C,G,T\}}$$

and second, for each M a constant vector of weights $\omega_M = (\omega_1, \omega_2, \dots, \omega_n)$.

There are two methods for producing ω_M . The first, which we call weighted sum, is the difference in the probabilities for the two letters of the polymorphism (or variant), *i.e.* Δp_{s_i} , or the difference of the maximum and minimum values for each column of M:

Equation 4.1

$$\omega_i = \max\{M_i\} - \min\{M_i\}$$
 where $i = 1, \dots n$

The second variation of this theme is to weight by relative entropy. Thus the relative entropy weight for each column i of the matrix is given by:

Equation 4.2

$$\omega_i = \sum_{j \in \{A,C,G,T\}} M_{j,i} \log_2(\frac{M_{j,i}}{b_i})$$
 where $i = 1, \dots n$

where b_i is again the background frequency of the letter i.

Thus, there are 3 possible algorithms to apply via the method argument. The first is the standard summation of log probabilities (method='log'). The second and third are the weighted sum and information content methods (method='default' and method='ic') specified by equations 4.1 and 4.2, respectively. **motifbreakR** assumes a uniform background nucleotide distribution (b) in equations 1 and 4.2 unless otherwise specified by the user. Since we are primarily interested in the difference between alleles, background frequency is not a major factor, although it can change the results. Additionally, inclusion of background frequency introduces potential bias when collections of motifs are employed, since motifs are themselves unbalanced with respect to nucleotide composition. With these cautions in mind, users may override the uniform distribution if so desired. For all three methods, **motifbreakR** scores and reports the reference and alternate alleles of the sequence ($F(s_{\text{REF}}, M)$ and $F(s_{\text{ALT}}, M)$), and provides the matrix scores $p_{s_{\text{REF}}}$ and $p_{s_{\text{ALT}}}$ of the SNP (or variant). The scores are scaled as a fraction of scoring range 0-1 of the motif matrix, M. If either of $F(s_{\text{REF}}, M)$ and $F(s_{\text{ALT}}, M)$ is greater than a user-specified threshold (default value of 0.85) the SNP is reported. By default **motifbreakR** does not display neutral effects, ($\Delta p_i < 0.4$) but this behaviour can be overridden.

Additionally, now, with the use of TFMPvalue-package, we may filter by p-value of the match. This is unfortunately a two step process. First, by invoking filterp=TRUE and setting a threshold

at a desired p-value e.g 1e-4, we perform a rough filter on the results by rounding all values in the PWM to two decimal place, and calculating a scoring threshold based upon that. The second step is to use the function calculatePvalue() on a selection of results which will change the Refpvalue and Altpvalue columns in the output from NA to the p-value calculated by TFMsc2pv. This can be (although not always) a very memory and time intensive process if the algorithm doesn't converge rapidly.

Value

a GRanges object containing:

REF the reference allele for the variant
the alternate allele for the variant
snpPos the coordinates of the variant

motifPos The position of the motif relative the the variant

geneSymbol the geneSymbol corresponding to the TF of the TF binding motif

dataSource the source of the TF binding motif

providerName, providerId

the name and id provided by the source

seqMatch the sequence on the $5' \rightarrow 3'$ direction of the "+" strand that corresponds to DNA

at the position that the TF binding motif was found.

pctRef The score as determined by the scoring method, when the sequence contains the

reference variant allele, normalized to a scale from 0 - 1. If filterp = FALSE,

this is the value that is thresholded.

pctAlt The score as determined by the scoring method, when the sequence contains the

alternate variant allele, normalized to a scale from 0 - 1. If filterp = FALSE,

this is the value that is thresholded.

scoreRef The score as determined by the scoring method, when the sequence contains the

reference variant allele

scoreAlt The score as determined by the scoring method, when the sequence contains the

alternate variant allele

Refpvalue p-value for the match for the pctRef score, initially set to NA. see calculatePvalue

for more information

Altpvalue p-value for the match for the pctAlt score, initially set to NA. see calculatePvalue

for more information

altPos the position, relative to the reference allele, of the alternate allele

alleleDiff The difference between the score on the reference allele and the score on the

alternate allele

alleleEffectSize

The ratio of the alleleDiff and the maximal score of a sequence under the

PWM

effect one of weak, strong, or neutral indicating the strength of the effect.

each SNP in this object may be plotted with plotMB

See Also

See snps.from.rsid and snps.from.file for information about how to generate the input to this function and plotMB for information on how to visualize it's output

motifbreakR_motif

Examples

motifbreakR_motif

MotifDb object containing motif information from the motif databases of HOCOMOCO, Homer, FactorBook and ENCODE

Description

This object contains all the MotifList-class objects that were generated for this package. See the individual help sections for hocomoco, homer, factorbook, and encodemotif, for how the data is formatted.

Usage

```
motifbreakR_motif
```

Format

MotifDb object of length 2816; to access metadata use mcols(motifbreakR_motif)

Details

Load with data(motifbreakR_motif)

Value

```
MotifList-class object
```

Source

Kulakovskiy,I.V., Medvedeva,Y.A., Schaefer,U., Kasianov,A.S., Vorontsov,I.E., Bajic,V.B. and Makeev,V.J. (2013) HOCOMOCO: a comprehensive collection of human transcription factor binding sites models. Nucleic Acids Research, **41**, D195–D202.

Heinz S, Benner C, Spann N, Bertolino E et al. (2010 May 28) Simple Combinations of Lineage-Determining Transcription Factors Prime cis-Regulatory Elements Required for Macrophage and B Cell Identities. Mol Cell, **38(4):576-589**. PMID: 20513432

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J Wang, J Zhuang, S Iyer, XY Lin, et al. (2012) Sequence features and chromatin structure around the genomic regions bound by 119 human transcription factors. Genome Research, **22** (**9**), 1798-1812, doi:10.1101/gr.139105.112

Pouya Kheradpour and Manolis Kellis (2013 December 13) Systematic discovery and characterization of regulatory motifs in ENCODE TF binding experiments. Nucleic Acids Research, doi:10.1093/nar/gkt1249

See Also

hocomoco, homer, factorbook, and encodemotif

Examples

```
data(motifbreakR_motif)
motifbreakR_motif
```

plotMB

Plot a genomic region surrounding a genomic variant, and potentially disrupted motifs

Description

Plot a genomic region surrounding a genomic variant, and potentially disrupted motifs

Usage

```
plotMB(
   results,
   rsid,
   reverseMotif = TRUE,
   effect = c("strong", "weak"),
   altAllele = NULL
)
```

Arguments

results The output of motifbreakR

rsid Character; the identifier of the variant to be visualized

reverseMotif Logical; if the motif is on the "-" strand show the the motifs as reversed FALSE

or reverse complement TRUE

effect Character; show motifs that are strongly effected c("strong"), weakly effected

c("weak"), or both c("strong", "weak")

altAllele Character; The default value of NULL uses the first (or only) alternative allele for

the SNP to be plotted.

Details

plotMB produces output showing the location of the SNP on the chromosome, the surrounding sequence of the + strand, the footprint of any motif that is disrupted by the SNP or SNV, and the DNA sequence motif(s). The altAllele argument is included for variants like rs1006140 where multiple alternate alleles exist, the reference allele is A, and the alternate can be G,T, or C. plotMB only plots one alternate allele at a time.

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Value

plots a figure representing the results of motifbreakR at the location of a single SNP, returns invisible NULL.

See Also

See motifbreakR for the function that produces output to be visualized here, also snps.from.rsid and snps.from.file for information about how to generate the input to motifbreakR function.

Examples

```
data(example.results)
example.results
library(BSgenome.Hsapiens.UCSC.hg19)
plotMB(results = example.results, rsid = "rs1006140", effect = "strong", altAllele = "C")
```

shiny_motifbreakR

Run Shiny version of the motifbreakR package.

Description

Run Shiny version of the motifbreakR package.

Usage

```
shiny_motifbreakR()
```

Value

returns a shinyAppDir that launches the shiny app when printed.

Examples

```
library(motifbreakR)
app <- shiny_motifbreakR()
if (interactive()) {
   shiny::runApp(app)
}</pre>
```

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snps.from.file

Import SNPs from a BED file or VCF file for use in motifbreakR

Description

Import SNPs from a BED file or VCF file for use in motifbreakR

Usage

```
snps.from.file(
  file = NULL,
  dbSNP = NULL,
  search.genome = NULL,
  format = "bed",
  indels = FALSE,
  biomart.dataset = NULL,
  check.unnamed.for.rsid = FALSE
)

variants.from.file(
  file = NULL,
  dbSNP = NULL,
  search.genome = NULL,
  biomart.dataset = NULL,
  format = "bed"
)
```

Arguments

file Character; a character containing the path to a bed file or a vcf file see Details

for a description of the required format

dbSNP OPTIONAL; an object of class SNPlocs to lookup rsids; see availible.SNPs

in injectSNPs to check for available SNPlocs

search. genome an object of class BSgenome for the species you are interrogating; see available. genomes

for a list of species

format Character; one of bed or vcf

indels Logical; allow the import of indels.

biomart.dataset

object.

check.unnamed.for.rsid

Logical; check snps in the form chr:pos:ref:alt for corresponding rsid, lookup may be slow, requires either param dbSNP or biomart.dataset.

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Details

snps.from.file takes a character vector describing the file path to a bed file that contains the necissary information to generate the input for motifbreakR see httml#format1 for a complete description of the BED format. Our convention deviates in that there is a required format for the name field. name is defined as chromosome:start:REF:ALT or the rsid from dbSNP (if you've included the optional SNPlocs argument). For example if you were to include rs123 in it's alternate format it would be entered as chr7:24966446:C:A

Value

a GRanges object containing:

SNP_id The rsid of the snp with the "rs" portion stripped alleles_as_ambig

THE IUPAC ambiguity code between the reference and alternate allele for this

SNP

REF The reference allele for the SNP

ALT The alternate allele for the SNP

Functions

• variants.from.file(): Allows the use of indels by default

See Also

See motifbreakR for analysis; See snps.from.rsid for an alternate method for generating a list of variants.

Examples

snps.from.rsid

Import SNPs from rsid for use in motifbreakR

Description

Import SNPs from rsid for use in motifbreakR

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Usage

```
snps.from.rsid(
  rsid = NULL,
  dbSNP = NULL,
  search.genome = NULL,
  biomart.dataset = NULL)
```

Arguments

rsid Character; a character vector of rsid values from dbSNP

dbSNP an object of class SNPlocs to lookup rsids; see availible . SNPs in injectSNPs

to check for available SNPlocs

search.genome an object of class BSgenome for the species you are interrogating; see available.genomes

for a list of species.

biomart.dataset

a Mart object from useEnsembl specifying the snps biomart, which dataset i.e., hsapiens_snp, and which version i.e., 111 or GRCm39. This will override SNPlocs and must be compatible with search.genome selection, and will query from biomaRt, which may be considerably faster than a lookup from a SNPlocs object.

Details

snps.from.rsid take an rsid, or character vector of rsids and generates the required object to input into motifbreakR

Value

a GRanges object containing:

SNP_id The rsid of the snp with the "rs" portion stripped

alleles_as_ambig

THE IUPAC ambiguity code between the reference and alternate allele for this

SNP

REF The reference allele for the SNP
ALT The alternate allele for the SNP

See Also

See motifbreakR for analysis; See snps.from.file for an alternate method for generating a list of variants.

Examples

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