# Package 'rtfbsdb'

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# Description

The motif library from CisBP web site. Link: http://cisbp.ccbr.utoronto.ca/

# **Objects from the Class**

Objects can be created by calls of the form CisBP.extdata, CisBP.zipload, CisBP.download.

#### **Slots**

```
\mbox{species:}\ \mbox{String indicating the species name defined in the CisBP dataset.}
```

zip.file: String indicating the filename of temporary data file.

zip.url: String indicating the download source.

zip.date: String indicating the download date.

file.tfinfo: String indicating the TF filename, defulat is TF\_Information.txt.

#### **Extends**

Class "tfbs.db", directly.

# Methods

**tfbs.createFromCisBP** Build a tfbs object by querying the meta file of CisBP dataset and subsetting the results.

CisBP.group Get the statistical summary by grouping the field in the CisBP dataset.

CisBP.getTFinformation Get the TF Information stored in the CisBP dataset.

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#### References

Weirauch, M. T., Yang, A., Albu, M., Cote, A. G., Montenegro-Montero, A., Drewe, P., ... & Hughes, T. R. (2014). Determination and inference of eukaryotic transcription factor sequence specificity. Cell, 158(6), 1431-1443.

#### See Also

```
CisBP.getTFinformation, CisBP.group, tfbs.createFromCisBP
```

#### **Examples**

```
showClass("CisBP.db")
```

CisBP.download

Download CisBP dataset.

## **Description**

Download TF data file from CisBP dataset and store it to temporary folder

## Usage

```
CisBP.download(species = "Homo_sapiens",
    url = "http://cisbp.ccbr.utoronto.ca/bulk_archive.php")
```

## **Arguments**

species String, indicating the species name in the CisBP dataset url String, the URL of bulk dowbnloads from CisBP dataset,

default is http://cisbp.ccbr.utoronto.ca/bulk\_archive.php

#### **Details**

The dowload function has been confirmed in the web site of cisbp.ccbr.utoronto.ca o June, 2015.

#### Value

A CisBP object (class name: "CisBP.db") is returned with four items:

species String indicating the species name

zip.file String indicating the filename of temporary data file.

zip.url String indicating the download source

file.tfinfo String indicating the TF filename, default is TF\_Information.txt.

#### References

Weirauch, M. T., Yang, A., Albu, M., Cote, A. G., Montenegro-Montero, A., Drewe, P., ... & Hughes, T. R. (2014). Determination and inference of eukaryotic transcription factor sequence specificity. Cell, 158(6), 1431-1443.

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#### See Also

See Also as CisBP.zipload, CisBP.extdata.

## **Examples**

```
#download zebra fish dataset
db1 <- CisBP.download("Danio_rerio");
#download Felis_catus dataset
db2 <- CisBP.download("Felis_catus");</pre>
```

CisBP.extdata

Load internal CisBP dataset.

## **Description**

Build a CisBP object from the internal zip file stored in this package

#### Usage

```
CisBP.extdata(species)
```

## **Arguments**

species String, only valid for human and mouse species, i.e. Homo\_sapiens, Mus\_musculus,

or Drosophila\_melanogaster

#### **Details**

The CisBP data for Homo\_sapiens and Mus\_musculus are delivered by this package. When you use the newest dataset, you should download it from the website by CisBP.download.

#### Value

A CisBP object (class name: "CisBP.db") is returned with four items:

species String indicating the species name defined in the CisBP dataset.

zip.file String indicating the filename of temporary data file.

zip.url String indicating the download source

file.tfinfo String indicating the TF filename, default is TF\_Information.txt.

#### See Also

See Also as CisBP.zipload, CisBP.download.

#### **Examples**

```
#reading human data from extension data file in the package
db.human <- CisBP.extdata("Homo_sapiens")

#reading Drosophila_melanogaster from extension data file in the package
db.dm3 <- CisBP.extdata("dm3")</pre>
```

CisBP.getTFinformation

Get TF information with PWM status

## **Description**

Get TF information with PWM status

#### Usage

```
CisBP.getTFinformation(cisbp.db, tf.information.type = NA)
```

#### **Arguments**

```
cisbp.db A CisBP object ("CisBP.db") including the TF_Information.txt.

tf.information.type

Number, indicating which TF meta file will be used. Available values are 1 for TF_Information.txt, 2 for TF_Information_all_motifs.txt and 3 for F_Information_all_motifs_plus.txt.
```

# **Details**

Three TF information files in CisBP dataset.

- 1: TF\_Information.txt : (direct motifs) or (no direct but inferred motifs with 90%)
- 2: TF\_Information\_all\_motifs.txt: (direct motifs) and (inferred motifs above the threshold)
- 3: F\_Information\_all\_motifs\_plus.txt: All motifs

The following parts are copied from RAEDME.txt in zipped CisBP data file.

TF\_Information.txt, TF\_Information\_all\_motifs.txt, TF\_Information\_all\_motifs\_plus.txt - These files contain information on the TFs.

'TF\_Information.txt' contains, for each TF, all directly determined motifs (see below). If a TF does not have a directly determined motif, this file will also include its best inferred motif. 'Best' is defined as the motif(s) obtained from the most similar TF (based on the

'TF\_Information\_all\_motifs.txt' is a superset of 'TF\_Information.txt'. It also includes any motif that can be inferred for a given TF, given the TF family-specific threshold. For example, if a TF has a directly determined motif, and two TFs with motifs with 90 TF\_Information\_all\_motifs.txt will include all three motifs. Likewise, if a TF does not have a direct motif, but has two TFs with 90

'TF\_Information\_all\_motifs\_plus.txt' is a superset of the other two files. It contains all motifs for a given TF, which includes all direct motifs, and all inferred motifs above the threshold.

#### Value

A data frame returned with the status indicating PWM data is existing or not

TF\_ID Internal CisBP ID for the TF. Each gene has a unique TF ID

Family\_ID Internal CisBP ID for the TF family. A family is the unique set of DNA binding

domains (DBDs) present in the protein.

TSource\_ID Internal CisBP ID for the source of the TF (i.e. where its genome sequence was

obtained).

Motif\_ID Internal CisBP ID for the associated motif.

MSource\_ID Internal CisBP ID for the source of the motif (i.e. which database or study it

came from)

DBID External ID of the RBP (e.g., Ensembl ID)

TF\_Name Name of the TF
TF\_Species Species of the TF

TF\_Status Motif status of the TF. 'D' stands for directly determined motif. 'I' indicates that

the motif is inferred from another TF, based on DBD similarity (see Weirauch

et al. 2013 for details). 'N' means no motif is available.

Family\_Name Name of the TF's family

DBDs The unique set of DBDs (Pfam names) present in the TF

DBD\_Count Number of unique DBDs in the TF

Cutoff Cutoff used to infer motifs for the TF family

DBID Motif ID from the associated database or study

Motif\_Type Experimental assay used to determine the motif

MSource\_Identifier

ID for the source of the motif (i.e., its project name)

MSource\_Type Internal CisBP ID for the motif category
MSource\_Author First author for the source of the motif
MSource\_Year Year of publication of the motif source

PMID Pubmed ID of the motif source

MSource\_Version

Version of the source (i.e. database build)

TFSource\_Name Source of the TF (i.e. where did the genome build come from?)

TFSource\_Year Year the genome data was downloaded
TFSource\_Month Month the genome data was downloaded
TFSource\_Day Day the genome data was downloaded

motif\_existing Status indicating PWM data is existing or not

#### See Also

See Also as CisBP.group, CisBP.extdata, CisBP.zipload, CisBP.download

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#### **Examples**

```
# Load the internal CisBP dataset
db_human <- CisBP.extdata("Homo_sapiens");

df.tfinfo <- CisBP.getTFinformation( db_human, tf.information.type = 2)
show(head(df.tfinfo));</pre>
```

CisBP.group

Summarize the motif number.

## Description

Get the statistical summary by grouping the field in the CisBP dataset.

## Usage

#### **Arguments**

cisbp.db A CisBP object ("CisBP.db") including the TF\_Information.txt.

group.by String, indicating which field will be used to group values. Available values are tf\_name, tf\_species, tf\_status, family\_name, motif\_type and msource\_id.

tf.information.type

Number, indicating which TF meta file will be used. Available values are 1 for

Number, indicating which TF meta file will be used. Available values are 1 for TF\_Information.txt, 2 for TF\_Information\_all\_motifs.txt and 3 for F\_Information\_all\_motifs\_plus.txt.

# **Details**

Three TF information files in CisBP dataset.

```
1: TF_Information.txt : (direct motifs) or (no direct but inferred motifs with 90%)
```

2: TF\_Information\_all\_motifs.txt: (direct motifs) and (inferred motifs above the threshold)

3: F\_Information\_all\_motifs\_plus.txt: All motifs

# Value

A data frame returned includes two columns

group\_by Values of grouping field number Counts of group value

# See Also

```
See Also as tfbs.createFromCisBP
```

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#### **Examples**

```
# Load the internal CisBP dataset
db_human <- CisBP.extdata("Homo_sapiens");

# Group the motif count by the column of family_name in TF_Information.txt
gr1 <- CisBP.group(db_human, group.by="family_name", tf.information.type=1 );

# Group the motif count by the column of tf_status in TF_Information.txt
gr2 <- CisBP.group(db_human, group.by="tf_status", tf.information.type=1 );

# Group the motif count by the column of tf_status
# in TF_Information_all_motifs.txt
gr3 <- CisBP.group(db_human, group.by="tf_status", tf.information.type=2);

# Group the motif count by the column of tf_status
# in F_Information_all_motifs_plus.txt
gr4 <- CisBP.group(db_human, group.by="tf_status", tf.information.type=3);</pre>
```

CisBP.zipload

Load the zipped CisBP file.

#### Description

Build a CisBP object from the zipped CisBP file.

## Usage

```
CisBP.zipload(zip.file, species = "Homo_sapiens")
```

#### **Arguments**

zip.file String, indicating the zipped file data

species String, indicating the species name in the CisBP database

# **Details**

The zip data canbe downloaded from the web site, please check CisBP.download.

## Value

A CisBP object (class name: "CisBP.db") is returned with four items:

species String indicating the species name

zip. file String indicating the filename of temporary data file.

zip.url String indicating the download source

file.tfinfo String indicating the TF filename, default is TF\_Information.txt.

## See Also

See Also as CisBP.extdata, CisBP.download.

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#### **Examples**

```
# Download the dataset
db1 <- CisBP.download("Arabidopsis_thaliana");

# Loading the zip file, the db2 and db3 have same TF data.
# Here is an example to show how to use CisBP.zipload.
# We dont nee to download it by CisBP.download and then load it
# by CisBP.zipload
db2 <- CisBP.zipload(db1@zip.file, species="Arabidopsis thaliana");</pre>
```

print.tfbs.enrichment Print the brief enrichment results

# **Description**

Print the brief enrichment results.

## Usage

```
## S3 method for class 'tfbs.enrichment'
print(x, ..., pv.threshold=0.05, pv.adj=NA )
```

#### **Arguments**

## **Details**

This command shows the calling parameters and significant motifs from the result object. The significant motifs are selected by the corrected p-value cutoff(0.05) and at most 20 significant motifs are listed. The adjust method of p-value is defined in the calling function.

#### Value

No return values.

## See Also

See also as tfbs.enrichmentTest.

# **Examples**

```
#See example in tfbs.enrichmentTest
```

print.tfbs.finding

Print scanning result of TF sites.

# Description

Print scanning result of TF sites.

# Usage

```
## S3 method for class 'tfbs.finding' print(x, ...)
```

# Arguments

x The result obtained by tfbs.scanTFsite.

... Additional arguments affecting the print produced.

#### **Details**

This function shows a brief information including calling parameters and enriched motifs.

#### Value

No return values.

#### See Also

```
See Also as tfbs.scanTFsite
```

# **Examples**

```
#See example in tfbs.scanTFsite
```

```
summary.tfbs.enrichment
```

Summarize the enrichment result

## **Description**

Return the significant motifs based on the adjust p-values using multiple comparisons.

# Usage

```
## S3 method for class 'tfbs.enrichment'
summary(object, pv.threshold = 0.05, pv.adj = NA, ...)
```

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## **Arguments**

object The result obtained by tfbs.enrichmentTest.

pv.threshold The p-value threshold for significant motifs.

pv.adj P-values adjust method for p.adjust function. The available values are "holm",

"hochberg", "hommel", "bonferroni", "BH", "BY", "fdr" or "none".

... Additional arguments affecting the summary produced.

#### **Details**

A data frame with 6 colums is returned.

#### Value

The results is a data frame including 6 columns,

 $\begin{array}{ll} \text{motif.id} & \text{Motif ID} \\ \text{tf.name} & \text{TF Name} \end{array}$ 

Npos Read count in positive loci. expected Read count in negative loci.

fe.ratio The ratio of read counts between positive loci and negative loci.

starch Cpmporessed Bed filename

pvalue p-value

pv.adj adjusted p-value by multiple comparson method.

## See Also

See also as tfbs.enrichmentTest.

```
summary.tfbs.finding Summarize scanning results.
```

# **Description**

Return a data frame with summarized TF sites for every motif if the calling parameter is "matches".

## Usage

```
## S3 method for class 'tfbs.finding'
summary(object, ...)
```

## Arguments

object The result obtained by tfbs.scanTFsite.

... Additional arguments affecting the summary produced.

#### **Details**

 $\hbox{summary in class of tfbs.} \hbox{finding is returned}.$ 

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#### Value

This function will return a data frame with summarized TF sites for every motif if the calling parameter is "matches", otherwise, NULL will be returned.

#### See Also

See Also as tfbs.scanTFsite

tfbs

Create a tfbs object from the supplied PWM files.

# Description

Create a tfbs object from the supplied PWM files.

## Usage

```
tfbs(filenames,
    names,
    species="Homo_sapiens",
    tf_info = NULL,
    tf_missing = NULL, ...)
```

## **Arguments**

filenames	Vector of PWM files
names	Vector of unique gene symbols.
species	String indicating species name
tf_info	Data frame including meta information copied from CisBP data file for all existing motifs., Default: NULL
tf_missing	Data frame including meta information copied from CisBP data file for missing motifs., Default: NULL
	Parameters, such as pseudocount, force_even, and the parameters used in read.table function.

# **Details**

Load the PWM files to build a "tfbs" object.

## Value

A tfbs object (class: "tfbs") including all PWM matrics. The all attributes are as follows:

TFID Vector of non-unique ID for TF.
species String indicating the species name
ntfs Number of motifs in matrix.
pwm A list including PWM matics.
filename Vector of PWM filename.

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mgisymbols	Unique gene symbols for TF.		
tf_info	Data frame, including extra information for all existing PWMs, it maybe different with motif dataset, default:NULL.		
tf_missing	Data frame, including extra information for missing PWMs, it maybe different with motif dataset, default:NULL.		
distancematrix	$Distance\ matrix\ between\ motifs\ returned\ by\ \verb"tfbs.clusterMotifs",\ default: NULL.$		
expressionlevel			
	Data frame indicatig the result of expression level returned by ${\tt tfbs.getExpression}$ , default: NULL.		
cluster	Matrix with 2 columns returned by tfbs.clusterMotifs, 1st column is the index of motifs and 2nd column is the group number of clustering, default:NULL.		

The tfbs object can be created by the function of tfbs, tfbs.dirs, tfbs.createFromCisBP.

#### See Also

```
tfbs, tfbs.dirs, tfbs.create From CisBP\\
```

# **Examples**

# Description

Tfbs object is a collection of motif PWM data. Some functions are provided based on the PWM and GENCODE data, such as clustering, search and compare.

# **Objects from the Class**

Objects can be created by calls of the function of tfbs.createFromCisBP, tfbs.dirs and tfbs.

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#### **Slots**

species String indicating the species name

ntfs Number of motifs in matrix.

pwm A list including PWM matics.

filename Vector of PWM filename.

mgisymbols Unique gene symbols for TF.

**tf\_info** Data frame, including extra information for all existing PWMs, it maybe different with motif dataset, default:NULL.

**tf\_missing** Data frame, including extra information for missing PWMs, it maybe different with motif dataset, default:NULL.

**distancematrix** Distance matrix between motifs returned by tfbs.clusterMotifs, default:NULL.

**expressionlevel** Data frame indicatig the result of expression level returned by tfbs.selectExpressedMotifs or tfbs.getExpression, default:NULL.

**cluster** Matrix with 3 columns returned by tfbs.clusterMotifs, 1st column is the index of motifs, 2nd column is the group number of clustering, 3rd column is selected flag by the function tfbs.selectByGeneExp or tfbs.selectByRandom. default:NULL.

#### Methods

tfbs.importMotifs Import the licensed motifs or other missing motifs for tfbs object

tfbs.getExpression Estimate gene expression of target TF.

**tfbs.selectExpressedMotifs** Select the expressed motifs in GRO-seq, PRO-seq or RNA-seq experimental data.

tfbs.clusterMotifs Cluster the specified motifs and drawing the heatmap.

tfbs.scanTFsite Find TF sites from genome data within the BED ranges.

tfbs.enrichmentTest Comparative TFBS search with the BED ranges

tfbs.selectByGeneExp Select the motifs with minimum p-value from each group of clustering.

tfbs.selectByRandom Select the motifs randomly from each group of clustering.

tfbs.drawLogosForClusters Draw the motif logos by one group per page.

tfbs.drawLogo Draw the logo for a single TF motif.

#### See Also

The class definition of tfbs.

## **Examples**

```
showClass("tfbs")
```

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tfbs.clusterMotifs Clustering the specified motifs and drawing the heatmap.

## **Description**

Clustering the specified motifs and drawing the heatmap.

## Usage

```
tfbs.clusterMotifs(tfbs,
    method = c("agnes", "apcluster"),
    pdf.heatmap = NA,
    group.k = NA,
    apcluster.q = 0.99,
    ncores = 1,
    plot.style = c("rtfbsdb", "apcluster"),
    BG = log(c(0.25, 0.25, 0.25, 0.25)),
    ...)
```

## **Arguments**

tfbs	A tfbs object ("tfbs") returned by tfbs.createFromCisBP, tfbs.dirs or other functions.	
method	String, availabe values are "agnes" and "apcluster".	
pdf.heatmap	String, a PDF filename for heatmap.	
group.k	Integer, if the method of agnes is used to do clustering, the parameter of $k$ is optional to use as preset group number.	
apcluster.q	Numeric value between 0 and 1, if the method of appluster is used to do clustering, the parameter of q is optional to use as preset group number.	
ncores	Number, the number of cores to use simultaneously.	
plot.style	String indicating the heatmap is plotted by the apcluster package or not if the method apcluster is used.	
BG	The log value of probabilities for nucleotide A, C, G and T as Backgroud computing.	
	The parameters used in function apcluster.	

#### **Details**

 $This \ result \ of \ clustering \ will \ be \ used \ in \ the \ tfbs. drawLogosForClusters, \ tfbs. selectByGeneExp, \ tfbs. enrichmentTest.$ 

tfbs@cluster will be updated by the clustering matrix which 1st column is the index of motifs and 2nd column is the group number of clustering.

## Value

A matrix with 2 columns is returned, 1st column is the index of motifs and 2nd column is the group number of clustering.

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#### See Also

See Also as tfbs.selectByGeneExp and tfbs.selectByRandom

#### **Examples**

## **Description**

Build a tfbs object by querying the meta file of CisBP dataset and subsetting the results.

#### Usage

```
tfbs.createFromCisBP(cisbp.db,
    motif_id = NULL,
    tf_name = NULL,
    tf_status = NULL,
    family_name = NULL,
    motif_type = NULL,
    msource_id = NULL,
    tf.information.type = 1)
```

# **Arguments**

cisbp.db	A CisBP object("CisBP.db"), including the file of TF_Information.txt.
motif_id	String, indicating the Motif_ID field will be used to select motifs.
tf_name	String, indicating the TF_name field will be used to select motifs.
tf_status	String, indicating the TF_Status field will be used to select motifs.
family_name	String, indicating the Family_Name field will be used to select motifs.

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```
motif_type String, indicating the Motif_Type field will be used to select motifs.

msource_id String, indicating the MSource_Identifier field will be used to select motifs.

tf.information.type

Number,indicating which TF meta file will be used. Available values are 1
for TF_Information.txt, 2 for TF_Information_all_motifs.txt and 3 for
```

TF\_Information\_all\_motifs\_plus.txt.

#### **Details**

The function includes three steps to build a tfbs object:

1) Searching the TF information and PWM files in the CisBP dataset according to the criteria specified by the parameters of *tf\_name*, *tf\_status*, *family\_name*, *motif\_type* and *msource\_id*.

#### Value

A tfbs object is returned with PWM matrices, see Also as "tfbs"

#### See Also

See Also as tfbs

## **Examples**

tfbs.createFromMotifDb

Create TF object by querying the MotifDb dataset.

#### **Description**

Create TF object from the MotifDb object which is a annotated collection of Protein-DNA binding sequenece motifs. The subset of MotifDb can be obtained by the methods in the MotifDb package or by the criteria in this function.

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# Usage

```
tfbs.createFromMotifDb(motifDB= NULL,
      organism = "Hsapiens",
      geneSymbol = NULL,
      tfFamily = NULL,
      providerName = NULL,
      providerId = NULL,
      dataSource = NULL,
      geneId = NULL,
      geneIdType = NULL,
      proteinId = NULL,
      proteinIdType = NULL,
      sequenceCount = NULL,
      bindingSequence = NULL,
      bindingDomain = NULL,
      experimentType = NULL,
      pubmedID = NULL,
      pseudocount = -7)
```

check full list.

# Arguments

motifDB

	MOCITOD	Would be object of subset of Mould b.
	organism	String, species, use command print (table (values (MotifDb)\$organism)) to check full list.
	geneSymbol	String, gene symbol is used as Motif_ID in "tfbs", use command print (table (values (MotifDb)\$geneSymbol)) to check full list.
	tfFamily	String, TF family is used as TF_Name in "tfbs", use command print (table (values (MotifDb)\$tfFamily)) to check full list.
	providerName	String, use command print (table (values (MotifDb)\$providerName)) to check full list.
	providerId	String, use command print (table (values (MotifDb)\$providerId)) to check full list.
	dataSource	String, use command print (table (values (MotifDb)\$dataSource)) to check full list.
	geneId	String, use command print (table (values (MotifDb)\$geneId)) to check full list.
	geneIdType	String, use command print (table (values (MotifDb)\$geneIdType)) to check full list.
	proteinId	String, use command print (table (values (MotifDb)\$proteinId)) to check full list.
	proteinIdType	String, use command print (table (values (MotifDb)\$proteinIdType)) to check full list.
	sequenceCount	String, use command print (table (values (MotifDb)\$sequenceCount)) to check full list.
bindingSequence		
		String, use command print (table (values (MotifDb)\$bindingSequence)) to

MotifDb object or subset of MotifDb.

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bindingDomain String, use command print (table (values (MotifDb)\$bindingDomain)) to check full list.

experimentType String, use command print (table (values (MotifDb)\$experimentType)) to check full list.

pubmedID String, use command print (table (values (MotifDb)\$pubmedID)) to check full list.

pseudocount Number, log value for zero value in PWM matrix, default is -7.

#### **Details**

Two methods to make a subset obeject.

- 1.Using the methods provided by the MotifDB package, please check the maunal of MotifDb package.
- 2. Searching the meta information and PWM matrices in the MotifDb object according to the criteria specified by the parameters of *organism*, *geneSymbol*, *tfFamily*, *providerName*, etc.

#### Value

A tfbs object is returned with PWM matrices, see Also as "tfbs"

## Author(s)

MotifDB is authored by Paul Shannon.

#### References

The MotifDB is in Bioconductor. http://bioconductor.org/packages/release/bioc/html/MotifDb.html

## See Also

```
See Also as "tfbs"
```

# Examples

```
library(rtfbsdb);

if( require(MotifDb) )
{
    # Load the subset of MotifDb generated by the method in 'MotifDb' package
    mdb.sox4 <- query (MotifDb,'sox4');
    tfs0 <- tfbs.createFromMotifDb(mdb.sox4, organism=NULL);

# Load the subset of MotifDb generated by the method in 'MotifDb' package
    mdb.human <- query(MotifDb, 'hsapiens');
    tfs1 <- tfbs.createFromMotifDb(mdb.human, organism=NULL);
}

# Load all motifs.
tfs2 <- tfbs.createFromMotifDb(organism=NULL);
show(MotifDb);</pre>
```

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```
head(unique(tfs2@tf_info$organism), n=30);
head(unique(tfs2@tf_info$tfFamily), n=30);
head(unique(tfs2@tf_info$dataSource), n=30);
head(unique(tfs2@tf_info$experimentType), n=30);
head(unique(tfs2@tf_info$geneIdType), n=30);
head(unique(tfs2@tf_info$bindingDomain), n=30);
# Load all motifs of mouse species.
tfs4 <- tfbs.createFromMotifDb(organism="Mmusculus");</pre>
# Default: Load all motifs of human species.
tfs5 <- tfbs.createFromMotifDb();</pre>
# Load all motifs of Drosophila.melanogaster species.
tfs.Drosophila.melanogaster <- tfbs.createFromMotifDb(organism="Dmelanogaster");</pre>
# TFBS scanning
tfs.ap2 <- tfbs.createFromMotifDb(organism = "Hsapiens", tfFamily="AP2");</pre>
gen.bed <- data.frame(chr="chr19",</pre>
      start=round(runif(10,1000000, 2000000)),
      stop=0,
      name=""
      score=0,
      strand=".");
gen.bed$stop <- gen.bed$start + 3000;</pre>
file.twoBit <- system.file("extdata","hg19.chr19.2bit", package="rtfbsdb")</pre>
t1 <- tfbs.scanTFsite( tfs.ap2,</pre>
      file.twoBit,
      gen.bed,
      file.prefix="test.db",
      ncores = 1);
```

tfbs.db-class

Class "tfbs.db"

# Description

Abstract class for motif dataset. The CisBP class is a son class of tfbs.db.

# Objects from the Class

Now code or function can be used to create this class.

## **Slots**

species: Species name.

#### Methods

No methods defined with class "tfbs.db" in the signature.

tfbs.dirs 21

#### See Also

```
"CisBP.db" inherits this class.
```

#### **Examples**

```
showClass("tfbs.db")
```

tfbs.dirs

Create a tfbs object from the folders.

## **Description**

Create a tfbs object from all the PWM files found in the supplied folders.

## Usage

```
tfbs.dirs(...,
    species = "Homo_sapiens",
    args.read.motif = NULL,
    pattern = glob2rx("*.pwm"),
    recursive = FALSE)
```

## **Arguments**

... Multiple strings, one or more folders can be used in this function.

species String, including the species name.

args.read.motif

List, including *pseudocount*, *force\_even* or other parameters used in read. table

function.

pattern String, a character vector specifying regular expression and wlidcards.

recursive Logical, indicating the loading recursively descends into subfolders or not, de-

fault: FALSE.

## **Details**

Two parameters in the list of args.read.motif can be used: pseudocount: log value for zero value in PWM matrix, default is -7. force\_even: whether the PWM matrix with odd size needs to be even.

# Value

A tfbs object collecting all the PWM files in the specified folders. For the details of tfbs object, please see tfbs

#### See Also

The structure of tfbs object is described in "tfbs"

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# **Examples**

tfbs.drawLogo

Draw single motif logo.

#### **Description**

Draw the motif logos in two models, 1 logo within a page or 1 group within one page.

## Usage

## **Arguments**

tfbs	A tfbs object("tfbs")	
file.pdf	String, the file name of PDF report.	
index	Vector of number, indicating the motif index.	
tf_id	Vector of string, indicating the TF_ID string, TF_ID is one motif attribute in TF_Information.txt. (Default=NULL).	
motif_id	Vector of string, indicating the Motif_ID string, Motif_ID is one motif attribute in TF_Information.txt. (Default=NULL).	
tf_name	Vector of string, indicating the TF_Name string, TF_Name is one motif attribute in TF_Information.txt. (Default=NULL).	
family_name	Vector of string, indicating Family_Name string, Family_Name is one motif attribute in TF_Information.txt. (Default=NULL).	
tf_status	String, indicating the TF_status value, TF_status is one motif attribute in TF_Information.txt. (Default=NULL).	
groupby	String, indicating the group field is applied to print the motif, each group is printed in one page, the available values are NA, "Family_Name", "TF_Name", "TF_Status" or "Motif_Type". (Default=NA).	

# Details

Multiple selection is provided for outputting logos. The selected motifs by each criteria will be combined into one set.

Draw the motif logos in two models:

(1) 1 logo within a page (2) 1 group within one page. The motif logos are splitted if motif count is greater than 10.

#### Value

No return values.

#### See Also

```
See Also as "tfbs"
```

#### **Examples**

```
db <- CisBP.extdata("Homo_sapiens");</pre>
tfs <- tfbs.createFromCisBP(db);</pre>
\label{eq:motif_id} \text{motif\_id} \quad <- \text{ c( "M5604\_1.01", "M5441\_1.01", "M5162\_1.01");}
           <- c( "T093250_1.01", "T093251_1.01", "T093252_1.01", "T093253_1.01");</pre>
family_name<- c( "p53", "Homeodomain", "Paired box", "Pipsqueak");</pre>
#Draw 10 motif logos from first one.
tfbs.drawLogo(tfs, file.pdf="test-drawLogo1.pdf", index=c(1:10) );
#Draw logos for specified Motif_ID, or TF_ID, or TF_Name, or Family_Name
tfbs.drawLogo(tfs, file.pdf="test-drawLogo2.pdf",
      motif_id = motif_id,
      tf_id = tf_id,
      tf_name = "AP-2"
      family_name = family_name,
      groupby = "TF_Status");
#Draw logos for specified TF_Status
tfbs.drawLogo(tfs, file.pdf="test-drawLogo3.pdf", tf_status="D",
      groupby="TF_Status");
#unlink("test-drawLogo1.pdf");
#unlink("test-drawLogo2.pdf");
#unlink("test-drawLogo3.pdf");
```

tfbs.drawLogosForClusters

Draw the motif logos by clustering.

# **Description**

Draw the motif logos by one cluster per page.

#### Usage

```
tfbs.drawLogosForClusters(tfbs, file.pdf, nrow.per.page=6 )
```

#### **Arguments**

```
tfbs A tfbs object("tfbs").

file.pdf String indicating a PDF eilname.

nrow.per.page Number indicating row count in each page.
```

#### **Details**

It is different with tfbs.drawLogo which is capable of printing out motif logos in group. This group is calculated by the tfbs.clusterMotifs, not is classfied by any group filed.

#### Value

No return value.

#### See Also

```
See Also as tfbs.clusterMotifs
```

## **Examples**

tfbs.enrichmentTest

Comparative TS sites between positive and negative TRE loci

# Description

Comparative TS sites between positive and negative TRE loci for all motifs.

# Usage

```
tfbs.enrichmentTest(tfbs,
     file.twoBit,
     positive.bed,
     negative.bed=NA,
      file.prefix=NA,
      use.cluster=FALSE,
     ncores=1,
     gc.correction=TRUE,
     gc.correction.pdf=NA,
     gc.min.sample = 500,
     gc.robust.rep=NA,
      threshold = 6,
      threshold.type = c("score", "fdr"),
      gc.groups=1,
     background.order=2,
     background.length=100000,
     pv.adj = p.adjust.methods)
```

# Arguments

tfbs	A tfbs object, see also "tfbs"		
file.twoBit	String, the file name of genome data( e.g. hg19.2bit, mm10.2bit)		
positive.bed	Data frame, bed-formatted TRE loci.		
negative.bed	Data frame, bed-formatted background loci. If not specified, the genomic loci adjacent to positive one are randomly extracted as the negative bed.		
file.prefix	String, the prefix for outputted BED file, no bed files output if NA		
use.cluster	Clustering matrix with 2 columns, 1st column is the index of motifs and 2nd column is the group number of clustering. It can be obtained from tfbs.clusterMotifs. If no clustering matrix, all motifs are used to do the comparson. see <i>details</i>		
ncores	Number, comupting nodes in parallel environment.(default=1)		
gc.correction	Logical value, if the difference between positive and negative TREs is significant, the resampling will be applied to the correction for the negative TREs. (default=TRUE)		
gc.correction.pdf			
	String, indicating the pdf file name if the GC correction is checked. (default=NA)		
gc.min.sample	Number, indicating minimum sample size when resamping the negative genomic loci for the GC correct. (default=500)		
gc.robust.rep	Number, indicating whether resampling background set multiple times is applied to get the median of binding sites. (default=NA)		
threshold	Numeric value, if 'score' is specified in threshold.type, only binding sites with scores above this threshold are returned, if 'fdr' is specified in threshold.type, only binding sites with FDR (False Discovery Rate) less than this value can be selected. Default value is 6 for 'score' and 0.1 for 'fdr'.		
threshold.type	String value, two options are available. only sites with scores above this threshold are returned, not be used if NA. (default = 'score')		
gc.groups	Numeric value,indicating number of quantiles to group sequences into in rtfbs package. (default = 1)		

background.order

Number, order of Markov model to build background.(default=2).

background.length

Number, length of the sequence to simulate background.(default=100000).

pv.adj

String, P-values correct method for p.adjust function. The available values are "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr" or "none".

(default="bonferroni").

#### Details

(1)The difference of GC contents between positive.bed and negative.bed is checked before the comparson. The p-value of Wilcoxon-Mann-Whitney test shows this difference and helps the user to determine whether the GC correction is necessary. If the difference is very significant, please set gc.correction to do GC content correction by resampling the TREs from negative bed data based on the frequency of TREs in negative bed data. Use the parameter of gc.correction.pdf to output vioplot figurs in a pdf file if you want to check the visualized difference.

(2)The clustering matix indicates which motifs in the 1st column are slected to do comparson and which clustering group in the 2nd columns are applied to adjust p-values for multiple comparsons. The function applys the p-values adjust for each clustering group. If no clustering information, all motifs in the tfbs object will be selected and adjusted as one group, which is the most conservative method.

(3)This function heavily relys on some Unix commands to operate bed data and gencode data. Please make sure the following commands work normally in R terminal.

starch, sort-bed, bedtools, two Bit To Fa.

The function Sys. which and system help you to locate these commands and test its availability.

## Value

A object with the class name of "tfbs.enrichment" will be resturned in this comparson function. It includes one list of parameters parm and one data frame of results result.

result is a data frame with the following columns:

motif.id Motif ID. tf.name TF name.

Npos TF site count found in positive ranges. expected TF site count found in negative ranges.

fe.ratio Ratio of fold enrichment.

pvalue p-value calculated by fisher test.

pv. adj p-value corrected by the multiple correction.

starch Binary filename of detected TF sites.

The result can be outputted to a report by the function tfbs.reportEnrichment.

#### See Also

```
print.tfbs.enrichment, summary.tfbs.enrichment, tfbs.reportEnrichment.
```

## **Examples**

```
library(rtfbsdb);
file.twoBit <- system.file("extdata","hg19.chr19.2bit", package="rtfbsdb")</pre>
db <- CisBP.extdata("Homo_sapiens");</pre>
tfs <- tfbs.createFromCisBP(db, family_name="AP-2");</pre>
#make two dummy BED data frame for positive loci and negative loci
pos.bed <- data.frame(chr="chr19",</pre>
      start=round(runif(1000,1000000, 2000000)),
      stop=0,
      name="",
      score=0,
      strand=".");
pos.bed$stop <- pos.bed$start + round(runif(1000, 20, 30));</pre>
neg.bed <- data.frame(chr="chr19",</pre>
      start=round(runif(8000, 800000, 1800000)),
      stop=0,
      name=""
      score=0,
      strand=".");
neg.bed$stop <- neg.bed$start + round(runif(8000, 20, 30));</pre>
t1 <- tfbs.enrichmentTest( tfs,</pre>
      file.twoBit,
      pos.bed,
      neg.bed,
      gc.correction=TRUE,
      gc.min.sample = 1000,
      ncores = 1); #ncores=3
#Show a brief result
t1;
#Show the comparson results of all motifs
show(t1$result);
summary(t1);
#Output the result to one pdf report.
tfbs.reportEnrichment(tfs, t1, file.pdf="test-tfbs-enrich-all.pdf", sig.only=FALSE);
file.ELF1 <- system.file("extdata", "Chipseq-k562-chr19-ELF1.bed", package="rtfbsdb")</pre>
pos.bed<- read.table(file.ELF1)</pre>
tfs <- tfbs.createFromCisBP(db, family_name="Ets");</pre>
t2 <- tfbs.enrichmentTest( tfs,</pre>
      file.twoBit,
      pos.bed,
      neg.bed,
      gc.correction=TRUE,
```

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```
gc.robust.rep=5,
     ncores = 1); #ncores=3
show(t2)
#Output the result to one pdf report.
tfbs.reportEnrichment(tfs, t2, file.pdf="test-tfbs-enrich-both.pdf",
      sig.only=TRUE, enrichment.type="both");
#plot QQ-like plot for the pvalues of all motifs
tfbs.plotEnrichment(tfs, t2, file.pdf="plot-tfbs-enrich-both.pdf",
     enrichment.type="both",
     plot.title="QQ plot",
      top.motif.labels=3, bottom.motif.labels=3, plot.type="polar", color.scheme=1);
t3 <- tfbs.enrichmentTest( tfs,
     file.twoBit,
     pos.bed.
     gc.correction=TRUE,
     gc.robust.rep=5,
     ncores = 1); #ncores=3
show(t3)
#Output the result to one pdf report.
tfbs.reportEnrichment(tfs, t3, file.pdf="test-elf1-enrich-depleted.pdf",
      sig.only=TRUE, enrichment.type="depleted");
#plot QQ-like plot for the pvalues of all motifs
tfbs.plotEnrichment(tfs, t3, file.pdf="plot-elf1-enrich-enriched.pdf",
     enrichment.type="enriched",
      plot.title="QQ plot",
      top.motif.labels=3, plot.type="nonpolar", color.scheme=2);
```

tfbs.getExpression

Estimate gene expression of target TF.

## **Description**

Gets expression level of target TF. USE extra\_info\$DBID to find gene information encoded by GENCODE

#### Usage

```
tfbs.getExpression(tfbs,
    file.twoBit,
    file.gencode.gtf,
    file.bigwig.plus=NA,
    file.bigwig.minus=NA,
    file.bam=NA,
```

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```
seq.datatype = c("GRO-seq", "PRO-seq", "RNA-seq"),
use.strand=FALSE,
ncores =1 )
```

## **Arguments**

tfbs A tfbs object("tfbs").

file.twoBit String, indicating the binary data of sequence. (e.g. hg19.2bit, mm10.2bit)

file.gencode.gtf

Gencode RDATA file encoded by ths package.

file.bigwig.plus

String, indicating bigwig file for strand plus(+) if seq.datatype is GRO-seq or

PRO-seq.

file.bigwig.minus

String, indicating bigwig file for strand minus(-) if seq.datatype is GRO-seq

or PRO-seq.

file.bam String, indicating BAM file for rna reads if seq. datatype is RNA-seq.

seq.datatype String, indicating which kind of seq data is applied to this function, three values

are available: GRO-seq, PRO-seq and RNA-seq. (Default=GRO-seq)

use.strand Logical, indicating whether same strandedness is required when getting the

number of mapped reads from BAM files for RNA-seq.

ncores Number, comupting nodes in parallel environment.

#### **Details**

1) For each motif, the occurance ranges can be queried by the gene ID in the GENCODE database( for human, gencode.v19.annotation.gtf, for mouse: gencode.vM3.annotation.gtf). After the searching, one range obtianed from the merge of the multiple ranges will be used to detect the reads count in the specified bigwig files(including plus and minus). The probability of each motif can be calcuated by the reads count and lambda.

The lambda is determined by the following formulation:

```
r.lambda = 0.04 * sum(reads_in_all_chromosomes)/10751533/1000.
```

2) This function heavily relys on some Unix commands to operate bed data and gencode data. Please make sure the following commands work normally in R terminal.

```
awk, zcat, sort-bed, twoBitInfo, bedtools.
```

The function Sys. which and system help you to locate these commands and test its availability.

# Value

A tbfs object with new expression data frame including the follwing columns:

Motif\_ID Motif\_ID from CisBP dataset or other data source.

DBID DBID from CisBP dataset or other data source.

txID Transcript ID from GTF file transcript which is mapped by DBID.

chr String, chromosome name of transcript which is mapped by DBID.

txStart Integer, start postion of transcript. txEnd Integer, end postion of transcript.

txLength or exonLen

Integer, the length of transcript for GRO-seq data and PRO-seq data or sum

length of exons on the transcript for RNA-seq data.

strand String, + or -, indicating the strand direction.

reads The reads number queried by BigWig function from the bigwig files( plus and

minus)

lambda The lambda parameter in poison distribution.

reads.RPKM The RPKM value of reads column.

lambda.RPKM The RPKM value of lambda column.

p.pois The probability calculated based on Poisson distribution.

#### See Also

```
See Also as "tfbs"
```

# **Examples**

tfbs.importMotifs

Import motifs to tfbs object

#### **Description**

Import licensed motifs or the source other than Cis-BP to tfbs object

## Usage

# Arguments

tfbs	$A \ tfbs \ object \ ("tfbs") \ returned \ by \ tfbs. create From CisBP, \ tfbs, \ tfbs. dirs.$	
format	String value indicating predefined format or string vector containing some tags to define a customize format. Predefined formats include <b>pwm.matrix</b> , <b>jaspar</b> , <b>meme</b> , <b>mscan</b> , <b>transfac</b> and <b>HOCOMOCO</b> .	
filenames	Vector of file names.	
motif_ids	Vector of motif ID only for the format <b>pwm.matrix</b> . Theses motif IDs are assigned to the motif data loaded from 'pwm.matrix' files. If these motifs are missing or licensed motifs, these IDs have to be in accordance with the TF information which can be exported from 'TF_information.txt' by the function CisBP.getTFinformation.	
PPM.format	Boolean value indicating whether the matrix data is format by PPM(position probability matrix) or PFM(position frequency matrix), otherwise, the original data is PWM format. Default: FALSE.	
skip.lines	Number indicating specified non-empty lines are skipped from the head of data file. This parameter is not used for the format 'pwm.matrix'. Default: 0.	
pseudocount	Number value indicating to replace -Inf in PWM. Default: -7.	
force_even	Boolean value indicating whether to append a row to make the row number even. Default: FALSE.	
•••	Optional paramaters used in the function read.table for the 'pwm.matrix' format.	

## **Details**

Two goals to import the motifs from the sources other than Cis-BP.

- 1) Fill the missing motifs mainly licensed in the Cis-BP database.
- 2) Make use of the different source.

Five predefined formats are available, 'pwm.matrix', 'jaspar', 'meme', 'mscan', 'transfac' and 'HOCOMOCO'.

## 'pwm.matrix'

Single text file containing 5 colums, Position, A, C, G and T as shown below.

```
Pos A C G T
1 0.2 0.0 0.4 0.4
2 0.0 0.4 0.4 0.2
3 0.2 0.2 0.2 0.4
4 0.6 0.0 0.4 0.0
5 0.0 0.4 0.0 0.6
6 0.0 0.4 0.4 0.2
7 0.0 0.0 1.0 0.0
8 0.2 0.0 0.0 0.8
```

```
9 0.0 0.6 0.0 0.4
10 0.6 0.4 0.0 0.0
11 0.0 1.0 0.0 0.0
```

#### 'jaspar'

The package use a predifined template to load the 'jaspar' format as show below.

```
> Mycn
A [ 0 29 0 2 0 0 ]
C [31 0 30 1 3 0 ]
G [ 0 0 0 28 0 31]
T [ 0 2 1 0 28 0 ]
```

#### 'meme'

The 'meme' format reference: <a href="http://meme-suite.org/">http://meme-suite.org/</a>

The package use a predifined template to load the 'meme' format as show below.

```
MOTIF crp alternative name
```

```
letter-probability matrix: alength= 4 w= 19 nsites= 17 E= 4.1e-009
0.000000 0.176471 0.000000 0.823529
0.000000 0.058824 0.647059 0.294118
0.000000 0.058824 0.000000 0.941176
0.176471 0.000000 0.764706 0.058824
0.823529 0.058824 0.000000 0.117647
0.294118   0.176471   0.176471   0.352941
0.294118 0.352941 0.235294 0.117647
0.117647 0.235294 0.352941 0.294118
0.529412 0.000000 0.176471 0.294118
0.058824 0.235294 0.588235 0.117647
0.176471 0.235294 0.294118 0.294118
0.000000 0.058824 0.117647 0.823529
0.058824 0.882353 0.000000 0.058824
0.764706 0.000000 0.176471 0.058824
0.058824 0.882353 0.000000 0.058824
0.823529 0.058824 0.058824 0.058824
0.176471 0.411765 0.058824 0.352941
0.411765 0.000000 0.000000 0.588235
0.352941 0.058824 0.000000 0.588235
```

#### 'mscan'

The 'mscan' format reference: http://www.cisreg.ca/cgi-bin/mscan/MSCAN The package use a predifined template to load the 'mscan' format as show below.

```
>mef2
10 0 0 0 22 0 6 2 3 4 22 10
0 2 12 0 0 0 0 0 0 0 0 0 0
9 20 2 0 0 0 0 0 0 0 0 10
3 0 8 22 0 22 16 20 19 18 0 2
>myf
```

```
0 16
                     0
                        6
                            0
      2 15
                 0
                   15
                        0
                           0
                              10
                                   0
1
   7
             0
                 9
                                   0 16
     10
          1
                    1
                        0 16
                               6
          0
                     0 10
                           0
                               0 10
```

#### 'transfac'

The transfac format reference: http://www.cisreg.ca/cgi-bin/mscan/MSCAN
The package use a predifined template to load the 'transfac' format as show below.

```
AC
    MA0001.1
ХΧ
    AGL3
ΙD
XX
DE
    MA0001.1 AGL3; from JASPAR
P0
                  C
                          G
                                 Τ
           Α
                                 2
1
           0
                 94
                          1
                 75
2
           3
                          0
                                19
3
          79
                  4
                          3
                                11
4
          40
                  3
                          4
                                50
5
                                29
          66
                  1
                          1
6
                  2
                          0
          48
                                47
7
          65
                  5
                          5
                                22
8
          11
                  2
                          3
                                81
9
                  3
          65
                        28
                                 1
10
           0
                  3
                        88
                                 6
ХΧ
CC
    program: jaspar
ХΧ
//
```

The package implemented a simple parser to load motifs from the different sources. This parser basically reads the data file word by word according to the format rules defined in advance for different source.

The format rules are comprised of some fixed vocabularies and tags defined by the parser or the user. So there are two kinds of tags to describe a motif format, predefined tags and user-defined tags. All tags start with the dollar(\$) symbol and meet the requirement of program identifier, such as \$Express\_Pvalue,\$D1.

The predefined tags include the following names, which define the motif information and control the parser's cursor.

```
Start-stop characters for a line or a motif
[1]
      '>' '[' ']'
                     control
 [2]
      $SKIP n
                     control
                               Ignore or skip the rest part or next n lines.
[3]
      $REPEAT
                               Repeat use the current format until it can't be matched.
                     control
[4]
      $EOM
                               End of Motif.
                     control
                               Line of Motif.
[5]
      $LOM
                     control
[6]
      $Motif_ID
                     variable
                               Motif_ID required in the 'tfbs' object.
[7]
      $TF_Name
                     variable
                               TF_Name required in the 'tfbs' object.
[8]
                               Multiple A nucleobase frequencies in one row
      $A+
                     variable
[9]
      $C+
                     variable
                               Multiple C nucleobase frequencies in one row
[10]
      $G+
                     variable
                               Multiple G nucleobase frequencies in one row
                               Multiple T nucleobase frequencies in one row
[11]
      $T+
                     variable
```

[12]	\$A	variable	Single <i>A</i> nucleobase frequence in one column
[13]	\$C	variable	Single <i>C</i> nucleobase frequence in one column
[14]	\$G	variable	Single <i>G</i> nucleobase frequence in one column
[15]	\$T	variable	Single <i>T</i> nucleobase frequence in one column

The user-defined tags reprsent the variables which values can be collected to the 'tf\_info' slot in the tfbs object ("tfbs"). e.g. \$Description, \$anyword.

The following section shows 5 predefined format:

# 'jaspar'

```
>$Motif_ID $TF_Name
A [ $A+ ]
C [ $C+ ]
G [ $G+ ]
T [ $T+ ]
```

## 'meme'

```
MOTIF $Motif_ID $TF_Name $SKIP
letter-probability matrix: $SKIP
$A $C $G $T $REPEAT
URL $SKIP
```

## 'mscan'

```
>$Motif_ID $TF_Name $SKIP
$A+
$C+
$G+
$T+
```

## 'transfac'

```
AC $Motif_ID

XX $SKIP

ID $TF_Name

XX $SKIP

DE $Description

PO A C G T

$LOM $A $C $G $T $REPEAT

XX $SKIP

CC $SKIP $REPEAT

// $EOM
```

## 'HOCOMOCO'

```
>$Motif_ID,
$A $C $G $T $REPEAT
```

#### Value

A new tfbs object ("tfbs") merged with licensed motifs.

#### References

```
1.http://rsat01.biologie.ens.fr/rsa-tools/help.convert-matrix.html
2.http://www.cisreg.ca/cgi-bin/mscan/MSCAN
3.http://meme-suite.org/
4.http://hocomoco.autosome.ru/downloads
```

#### See Also

tfbs.createFromCisBP

#### **Examples**

```
require(rtfbsdb);
db <- CisBP.extdata("Homo_sapiens");</pre>
tfs <- tfbs.createFromCisBP(db, family_name="AP-2");</pre>
tfs;
#It is test code
#motif_ids <- c( "M2938_1.02", "M2940_1.02", "M2940_2.02", "M4056_1.02" );</pre>
#path <- system.file("extdata", package="rtfbsdb");</pre>
#file_pwms <- paste(path, c(</pre>
       "fake_M2938_1.02.pwm",
#
       "fake_M2940_1.02.pwm",
#
       "M3591_1.01.pwm",
       "M3590_1.01.pwm",
#
       ), sep="/");
#tfs <- tfbs.importMotifs(tfs, 'pwm.matrix', file_pwms, motif_ids, header=T );</pre>
#show(tfs);
## import 2 motifs to fill the licensed motifs in Cis-BP and 1 new motif fromother source
motif\_ids <- c( \ "M2938\_1.02", \ "M3591\_1.01", \ "M3590\_1.01" \ );
path <- system.file("extdata", package="rtfbsdb");</pre>
file_pwms <- paste(path, c(</pre>
      "fake_M2938_1.02.pwm",
      "M3591_1.01.pwm",
      "M3590_1.01.pwm"), sep="/");
tfs <- tfbs.importMotifs(tfs, 'pwm.matrix', file_pwms, motif_ids, header=TRUE );</pre>
show(tfs);
## Data is copied from http://rsat01.biologie.ens.fr/rsa-tools/help.convert-matrix.html
data.transfac <- system.file("extdata", "pwm.example.transfac.txt", package="rtfbsdb");</pre>
tfs.transfac <- tfbs.importMotifs(tfs, "transfac", data.transfac, skip.lines=0);</pre>
```

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```
show(tfs.transfac);
## Data from
## http://jaspar.genereg.net/html/DOWNLOAD/JASPAR_CORE/pfm/nonredundant/pfm_all.txt
data.jaspar <- system.file("extdata", "pwm.example.jaspar.2015.txt", package="rtfbsdb");</pre>
tfs.jaspar <- tfbs.importMotifs(tfs, "jaspar", data.jaspar, skip.lines=0);</pre>
show(tfs.jaspar);
## Data from
## http://jaspar.genereg.net/html/DOWNLOAD/ARCHIVE/JASPAR2010/JASPAR_CORE/pfms/pfms_all.txt
data.mscan <- system.file("extdata", "pwm.example.mscan.jaspar2010.txt", package="rtfbsdb");</pre>
tbs <- tfbs();
tfs.mscan <- tfbs.importMotifs(tbs, "mscan", data.mscan, skip.lines=0);</pre>
show(tfs.mscan);
## Data from http://meme-suite.org/doc/download.html?man_type=web
data.meme <- system.file("extdata", "pwm.example.meme.Homo_sapiens.txt", package="rtfbsdb");</pre>
tbs <- tfbs();
tfs.meme <- tfbs.importMotifs(tbs, "meme", data.meme, skip.lines=5);</pre>
show(tfs.meme);
## Data from http://www.nature.com/nature/journal/v527/n7578/full/nature15518.html
format.style <- c("Base $Motif_ID $TF_Name $Experiment</pre>
$Ligand_sequbatch $Seed $Multinomial $Cycle $Is_matrix $Comment
$Genomic_pvalue $Enrichment_pvalue $Category $SKIP",
"A $A+",
"C $C+",
"G $G+",
"T $T+" );
data.file <- system.file("extdata", "pwm.example.nature15518.s1.txt", package="rtfbsdb");</pre>
tbs <- tfbs();
tfs.nature15518 <- tfbs.importMotifs(tbs, format.style, data.file, skip.lines=19)
show(tfs.nature15518);
## HOCOMOCO_10
## Data from http://hocomoco.autosome.ru/downloads
## The matrix is PWM format.
library(rtfbsdb)
format.style <- c(">$Motif_ID",
"$A $C $G $T $REPEAT");
data.file <- system.file("extdata", "HOCOMOCO_10_pwms_HUMAN_mono.txt", package="rtfbsdb");</pre>
tbs <- tfbs();</pre>
tfs.HOCOMOCO.human <- tfbs.importMotifs(tbs, format.style, data.file, skip.lines=0, PPM.format=FALSE)
show(tfs.HOCOMOCO.human);
```

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# **Description**

Output enrichment results to a PDF report which includes motif names, counts of TF site, p-value, enrichment ratio and motif logos.

# Usage

```
tfbs.plotEnrichment(tfbs, r.comp,
    file.pdf,
    enrichment.type = c ("both", "enriched", "depleted"),
    plot.type=c("nonpolar", "polar"),
    options=list() )
```

# **Arguments**

```
tfbs A tfbs object, see also "tfbs"

r.comp A result object from the function of tfbs.enrichmentTest

file.pdf String, the file name of PDF report.

enrichment.type

String, three values are available for significant motifs to be drawn out.(default="both").

plot.type String, indicating whether the depleted motifs are shown at the negative side.(default="nonpolar")

options List, containing the pre-defined parameters to control the plot, see details.
```

#### Details

The function draws a QQ-like plot which all motifs are ordered by its p-values along X axis and -log10(pvalue) are drawn along Y axis. If plot type is 'ploar', depleted motifs are seperated from enriched motifs and drawn at the negative side of Y axis. The motif color is determined by the enrichment ratio.

The parameter options has the following options:

```
options=list(
title = "",
width = 7,
height = 7,
y.max = NULL,
xlab = "Order",
ylab = "-log10(p-value)",
top.motif.labels = 5,
bottom.motif.labels = 5,
color.scheme = 2,
abline = NULL,
zoom.tick = 1,
zoom.label = 1,
zoom.motif.logo = 1,
zoom.legend.label=1,
zoom.motif.label = 1 );
```

- title: String, the plot title.
- width: Number, page width, default is 7 inches.

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- height: Number, page height, default is 7 inches.
- y.max: Number, the maximum value for Y axis. The points larger than this value will be drawn at the border.
- xlab: String, the label for X axis.
- ylab: String, the label for Y axis.
- top.motif.labels: String or number, the labels for top motifs.
- bottom.motif.labels: String or number, the labels for top motifs in the depleted side.
- color.scheme: Number, 1 or 2 are available, color scheme for the points based on enrichment ratios. (default=2).
- abline: Number, indicating to draw horizontal lines for significant p-values.
- zoom. tick: Number, the ratio to zoom in or out the ticks on the X and Y axis. (default=1).
- zoom. label: Number, the ratio to zoom in or out the labels on the X and Y axis. (default=1).
- zoom.motif.logo: Number, the ratio to zoom in or out the motif logos located at the top or bottom. (default=1)
- zoom.motif.label: Number, the ratio to zoom in or out the motif labels located at the top or bottom. (default=1)
- zoom. legend. label: Number, the ratio to zoom in or out the legend label. (default=1)

#### Value

No return values.

#### See Also

```
tfbs.enrichmentTest, summary.tfbs.enrichment.
```

# **Examples**

```
# see examples in tfbs.enrichmentTest
```

tfbs.reportEnrichment Output report for enrichment results.

### **Description**

Output enrichment results to a PDF report which includes motif names, counts of TF site, p-value, enrichment ratio and motif logos.

# Usage

```
tfbs.reportEnrichment(tfbs, r.comp,
    file.pdf = NA,
    report.size = "letter",
    report.title = "",
    enrichment.type = c ("both", "enriched", "depleted"),
    sig.only = TRUE,
    pv.threshold = 0.05,
    pv.adj = NA,
    sorted = c ("pvalue", "enrich.ratio"))
```

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## **Arguments**

tfbs A tfbs object, see also "tfbs"

r.comp A result object from the function of tfbs.enrichmentTest

file.pdf String, the file name of PDF report.
report.size String, the page size ( default="letter")

report.title String, the report title.

enrichment.type

String, three values are available for significant motifs to be printed out.(default="both").

sig.only String, indicating whether only significant motifs are outputted or not.(default=TRUE).

pv.threshold Numeric value, indicating whether the different threshold of p-value is applied

to select the significant motifs.

pv.adj String, indicating whether the different correction metod of p-value is applied to

select the significant motifs.

sorted String, indicating which field is used to sort the results and print in the report.

(default="pvalue")

#### **Details**

The table with 7 columns is outputted into a PDF report within letter size.

Two color bars are used to display p-values and enrichment ratios. Motif logos are shown visually in each row.

#### Value

No return values.

#### See Also

```
tfbs.enrichmentTest, summary.tfbs.enrichment.
```

# **Examples**

```
# see examples in tfbs.enrichmentTest
```

tfbs.reportFinding Make report for scanning results.

### **Description**

Output a PDF report includes motif names, counts of TF site and motif logos.

# Usage

# **Arguments**

tfbs A tfbs object, see also "tfbs"

r.scan A result object from the function of tfbs.scanTFsite

file.pdf String, the file name of PDF report.

report.size String, the page size ( default="letter")

report.title String, the report title.

### **Details**

The table with 4 columns is outputted into a PDF report within letter size. Motif logos are shown visually in each row.

#### Value

No return values.

#### See Also

```
tfbs.scanTFsite, print.tfbs.finding
```

### **Examples**

```
#See example in tfbs.scanTFsite
```

tfbs.scanTFsite

Find TF sites from genome data within the BED loci

# **Description**

Find TF sites from genome data within the BED loci. Please notice that this package does not provided genome data such as hg19.2bit, mm10.2bit.

# Usage

```
tfbs.scanTFsite(tfbs,
    file.twoBit,
    gen.bed,
    return.type=c("matches", "maxscore", "posteriors", "maxposterior", "writedb"),
    file.prefix=NA,
    usemotifs = NA,
    ncores = 1,
    threshold = 6,
    threshold.type = c("score", "fdr"),
    gc.groups = NA,
    background.order = 2,
    background.length = 1000000)
```

#### **Arguments**

tfbs A tfbs object ("tfbs") returned by tfbs.createFromCisBP, tfbs, tfbs.dirs. String, the file name of genome data(e.g. hg19.2bit or mm10.2bit) file.twoBit gen.bed Data frame, bed-formatted loci information with 6 columns String, five available values explained in th details(default = "matches") return.type file.prefix String, the prefix for outputted file, only used when the return.type is writedb usemotifs Vector indicating indexes of motif to be used in scanning. Number, computing nodes in parallel environment (default = 1). ncores Numeric value, if 'score' is specified in threshold.type, only binding sites threshold with scores above this threshold are returned, if 'fdr' is specified in threshold. type, only binding sites with FDR (False Discovery Rate) less than this value can be selected. Default value is 6 for 'score' and 0.1 for 'fdr'. threshold. type String value, two options are available. only sites with scores above this threshold are returned, not be used if NA. (default = 'score') Numeric value, indicating number of quantiles to group sequences into in rtfbs gc.groups package (default = 1). background.order Numeric value, indicating the order of Markov model to build in rtfbs package (default = 2).

background.length

Numeric value, indicating length of the sequence to simulate in rtfbs package (default = 100000)

## **Details**

- (1) Five options are availabl for the function of tfbs.scanTFsite as follows.
  - matches: returns all matching TF sites for all motifs.
  - writedb: writes a bed file with matches sites. Assumes that sort-bed and starch tools are available in \$PATH
  - maxscore: returns the max score(posterior difference between motif model and background) in each bed-formatted loci.
  - posteriors: returns the posteriors at each position in bed-formatted loci.
  - $\bullet$  maxposterior: returns the max(posterior) in each bed-formatted loci.
- (2) In order to make the binary file with the parameter of writedb, make sure that starchcat and sort-bed command (in BEDOPS) can be accessed from R environment. If not, please put the folder in \$PATH.
- (3)This function heavily relys on some Unix commands to operate bed data and gencode data. Please make sure the following commands work normally in R terminal.

starch, starchcat, sort-bed, twoBitInfo, twoBitToFa.

The function Sys. which and system help you to locate these commands and test its availability.

#### Value

A list object will be returned with the class name of tfbs.finding. The object wraps four sub-list as follows:

- 1) parm:Calling parameters(fdr, threshold), gc.groups...).
- 2) bed: Calling bed-formatted loci(gen.bed).
- 3) summary: A data frame including summrized information about matched TF sites for all motifs.
- 4) result: Scanning results which data type is depend on the parameter of return. type.

The option of *matches* returns a list including the result of every motif, which result is BED style data frame with the following columns.

chrom Chromosome chromStart Start position

chromEnd Chromosome end position

name

score The score is given by the log likelihood ratio against the Marklov model(backgound).

strand Strand

peakStart Start position of the matched BED region
peakEnd End position of the matched BED region

The option of writedb will return a binary BED filename in which store all bed ranges.

The option of *posteriors* will return a list for each motif returned by score.ms function. Scores represent the motif 'match score', or the product of the probability of observing each base under the motif or background models. Scores are returned under the motif model for all positions in the sequence, on both forward and reverse strands, and under the background model.

The option of *maxposterior* will return a probability matrix which the row indicates the target loci and the column indicates the motif.

The option of *maxscore* will return a score (poserior difference) matrix which the row indicates the target loci and the column indicates the motif.

## See Also

```
print.tfbs.finding, summary.tfbs.finding, tfbs.reportFinding.
```

```
library(rtfbsdb);
file.twoBit <- system.file("extdata","hg19.chr19.2bit", package="rtfbsdb")
db <- CisBP.extdata("Homo_sapiens");
tfs <- tfbs.createFromCisBP(db, family_name="Ets");
gen.bed <- data.frame(chr="chr19",</pre>
```

```
start=round(runif(10,1000000, 2000000)),
      stop=0,
      name=""
      score=0,
      strand=".");
gen.bed$stop <- gen.bed$start + 3000;</pre>
t1 <- tfbs.scanTFsite( tfs,</pre>
     file.twoBit,
      gen.bed,
      file.prefix="test.db",
      ncores = 1);
#show a brief information about the result
#show the summary information in the result
show(t1$summary);
#show the matched TF sites for first motif
show(t1$result[[1]]);
#Output a PDF report for all motifs.
tfbs.reportFinding(tfs, t1, file.pdf="test-rtfbs-scan.pdf", report.title="ELF1");
file.ELF1 <- system.file("extdata","Chipseq-k562-chr19-ELF1.bed", package="rtfbsdb")</pre>
gen.bed<- read.table(file.ELF1)</pre>
t2 <- tfbs.scanTFsite( tfs,
      file.twoBit,
      gen.bed,
      file.prefix="test.db",
      return.type="writedb",
      ncores = 1);
t2
t3 <- tfbs.scanTFsite( tfs,
      file.twoBit,
      gen.bed,
      return.type="posteriors",
      ncores = 1);
t3
t4 <- tfbs.scanTFsite( tfs,
      file.twoBit,
      gen.bed,
      return.type="maxposterior",
      ncores = 1);
t4;
dim(t4$result);
t5 <- tfbs.scanTFsite( tfs,
      file.twoBit,
```

```
gen.bed,
  return.type="maxscore",
  ncores = 1);
t5;
```

tfbs.selectByGeneExp Motif selection by gene expression level.

# **Description**

Select the motifs with minimum p-value from each group of clustering.

# Usage

```
tfbs.selectByGeneExp(tfbs)
```

## Arguments

tfbs

A tfbs object ("tfbs") with the data frame of gene expression level.

### **Details**

The function of tfbs.getExpression should be successfully called and the results of gene expression should be returned before this function is called. The indexes of selected motifs will be used in the function of tfbs.enrichmentTest or tfbs.scanTFsite.

## Value

New tfbs object with the selected indices is returned(tfbs@cluster). The 3rd column of tfbs@cluster is added or updated as the select flag(1:selected, 0:unselected).

### See Also

See Also as tfbs.selectByRandom, tfbs.getExpression

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tfbs.selectByRandom

Random motif selection

# **Description**

Select the motifs randomly from each group of clustering.

# Usage

```
tfbs.selectByRandom(tfbs)
```

# **Arguments**

```
tfbs A tfbs object("tfbs").
```

#### **Details**

The indexes of selected motifs can be used in the function of tfbs.enrichmentTest or tfbs.scanTFsite.

## Value

New tfbs object with the selected indices is returned(tfbs@cluster). The 3rd column of tfbs@cluster is added or updated as the select flag(1:selected, 0:unselected).

#### See Also

See Also as tfbs.selectByGeneExp, tfbs.getExpression

```
db <- CisBP.extdata("Homo_sapiens");

tfs <- tfbs.createFromCisBP(db, family_name="AP-2");

tfs <- tfbs.clusterMotifs(tfs, pdf.heatmap="test-AP2-heatmap.pdf" );

tfs <- tfbs.selectByRandom(tfs );

show(tfs@cluster);</pre>
```

```
tfbs.selectExpressedMotifs
```

Select expressed Motifs for GRO-seq, PRO-seq and RNA-seq data

## **Description**

Select expressed Motifs for GRO-seq, PRO-seq and RNA-seq data

# Usage

```
tfbs.selectExpressedMotifs(tfbs,
    file.twoBit,
    file.gencode.gtf,
    file.bigwig.plus=NA,
    file.bigwig.minus=NA,
    file.bam=NA,
    seq.datatype= c("GRO-seq", "PRO-seq", "RNA-seq"),
    pvalue.threshold = 0.05,
    lowest.reads.RPKM = NA,
    include.DBID.missing=TRUE,
    use.strand = FALSE,
    ncores = 1 )
```

## **Arguments**

tfbs A tfbs object ("tfbs") returned by tfbs.createFromCisBP, tfbs, tfbs.dirs. file.bigwig.plus

String, indicating bigwig file for strand plus(+) if seq. datatype is GRO-seq or PRO-seq.

file.bigwig.minus

String, indicating bigwig file for strand minus(-) if seq.datatype is GRO-seq

or PRO-seq.

file.bam String, indicating BAM file for rna reads if seq. datatype is RNA-seq.

file.twoBit String, indicating the binary data of sequence. (e.g. hg19.2bit, mm10.2bit)

file.gencode.gtf

String, indicating Gencode GTF file downloaded from the Gencode web site.

seq.datatype String,indicating which kind of seq data is applied to this function, three values are available: GRO-seq, PRO-seq and RNA-seq. Default: GRO-seq

pvalue.threshold

Numeric, indicating p-value criteria for expressed motifs. Default: 0.05

lowest.reads.RPKM

Numeric, implying the motifs with lower reads than this threshold will be removed from expressed list. Default: NA

include.DBID.missing

Logical, indicating whether the TFs without association with GENCODE through the DBID are selected.

use.strand Logical, indicating whether same strandedness is required when getting the

number of mapped reads from BAM files for RNA-seq.

ncores Number, comuputing nodes in parallel environment for gencode data converting.

#### **Details**

1) If seq. datatype is GRO-seq or PRO-seq and the bigwig files are provided, the gene expression values are calculated through querying the TREs region from the GENCODE database( for human, gencode.v19.annotation.gtf, for mouse: gencode.vM3.annotation.gtf) and querying the reads count in the plus and minus bigWig files.

If seq. datatype is RNA-seq and the BAM file is provided, read counts for each TRE regions will be queried from the BAM file.

2) If the expressed TFs only is used in the tfbs object, the TFs with p-values corrected by Bonfrroni less than 0.05 will be selected.

The following part explains how to calculate the gene expression.

For each motif, the occurance ranges can be queried by the gene ID After the searching, one range obtained from the merge of the multiple ranges will be used to detect the reads count in the specified bigwig files(including plus and minus). The probability of each motif can be calcuated by the reads count and lambda.

The lambda is determined by the following formulation:

```
For GRO-seq and PRO-seq data:
r.lambda = 0.04 * sum(reads_in_all_chromosomes)/10751533/1000.
```

For RNA-seq data:

```
r.lambda = mode( reads_in_1000_bp_windows_cross_all_gene_deserts )/1000.
```

3) This function heavily relys on some Unix commands to operate bed data and gencode data. Please make sure the following commands work normally in R terminal.

```
awk, zcat, sort-bed, twoBitInfo, bedtools.
```

The function Sys. which and system help you to locate these commands and test its availability.

4) This function will be failed to get the reads count if the BAM file is not indexed. Please use the command samtools to make the index file for the BAM file

```
samtools index your_bam_file
```

#### Value

A new tfbs object ("tfbs") with the matrix of gene expression level.

```
library(rtfbsdb);
# Load the internal CisBP data set
db.human <- CisBP.extdata("Homo_sapiens");</pre>
# Create a tfbs object by querying the meta file of CisBP dataset.
tfs <- tfbs.createFromCisBP(db.human, motif_type="ChIP-seq",</pre>
      tf.information.type=1 );
file.bigwig.minus <- system.file("extdata",</pre>
      "GSM1480327_K562_PROseq_chr19_minus.bw", package="rtfbsdb")
file.bigwig.plus <- system.file("extdata",</pre>
      "GSM1480327_K562_PROseq_chr19_plus.bw", package="rtfbsdb")
hg19.twobit <- system.file("extdata","hg19.chr19.2bit", package="rtfbsdb")</pre>
gencode.gtf <- system.file("extdata",</pre>
      "gencode.v19.annotation.chr19.gtf.gz", package="rtfbsdb")
tfs1 <- tfbs.selectExpressedMotifs(tfs,</pre>
      hg19.twobit,
      gencode.gtf,
      file.bigwig.plus,
      file.bigwig.minus,
      seq.datatype = "PRO-seq",
      pvalue.threshold=0.001,
      include.DBID.missing=TRUE,
      ncore=1);
show(tfs1)
file.bam <- "/local/storage/projects/NHP/AllData/bams/H3_U.fastq.gz.sort.bam"</pre>
tfs2 <- tfbs.selectExpressedMotifs(tfs,</pre>
      hg19.twobit,
      gencode.gtf,
      file.bam = file.bam,
      seq.datatype = "RNA-seq",
      pvalue.threshold=0.01,
      include.DBID.missing=TRUE,
      ncore=1);
show(tfs2)
```

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