# Package 'MatrixRider' 

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Type Package<br>Title Obtain total affinity and occupancies for binding site matrices on a given sequence<br>Version 1.37.0<br>Date 2015-11-02<br>Author Elena Grassi<br>Maintainer Elena Grassi [grassi.e@gmail.com](mailto:grassi.e@gmail.com)<br>Description Calculates a single number for a whole sequence that reflects the propensity of a DNA binding protein to interact with it. The DNA binding protein has to be described with a PFM matrix, for example gotten from Jaspar.<br>biocViews GeneRegulation, Genetics, MotifAnnotation<br>License GPL-3<br>Depends R (>= 3.1.2)<br>Imports methods, TFBSTools, IRanges, XVector, Biostrings<br>Suggests RUnit, BiocGenerics, BiocStyle, JASPAR2014<br>LinkingTo IRanges, XVector, Biostrings, S4Vectors<br>NeedsCompilation yes<br>git_url https://git.bioconductor.org/packages/MatrixRider<br>git_branch devel<br>git_last_commit e362023<br>git_last_commit_date 2024-04-30<br>Repository Bioconductor 3.20<br>Date/Publication 2024-05-24

## Contents

MatrixRider-package ..... 2
getSeqOccupancy ..... 2
Index ..... 4

MatrixRider-package Calculate total affinity and occupancies for binding site matrices on a given sequence

## Description

Calculates a single number for a whole sequence that reflects the propensity of a DNA binding protein to interact with it. The DNA binding protein has to be described with a PFM matrix, for example gotten from Jaspar.

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## Examples

library(JASPAR2014)
library(TFBSTools)
library(Biostrings)
pfm <- getMatrixByID(JASPAR2014,"MA0004.1")
\#\# The following sequence has a single perfect match
\#\# thus it gives the same results with all cutoff values.
sequence <- DNAString("CACGTG")
getSeqOccupancy (sequence, pfm, 0.1)
getSeqOccupancy (sequence, pfm, 1)
getSeqOccupancy Computes the total affinity or the occupancy at a given cutoff

## Description

The affinity/occupancy is calculated on the given DNAString for the given PFMatrix (or all the ones in the PFMatrix list given).

## Usage

getSeqOccupancy (sequence, pfm, cutoff)

## Arguments

sequence
pfm A PFMatrix or a PFMatrixList object with the matrixes whose affinity will be calculated. The background (bg,XMatrix-method) of the given pfm is used to perform affinity calculations.
cutoff numeric(1); between 0 and 1 (included): 0 corresponds to total affinity (i.e. summing all the affinities) while 1 to summing only values corresponding to the perfect match for a given PFMatrix. See vignette for details on how scores are calculated. If MatrixRider is installed, open the vignette with vignette("MatrixRider").

## Value

numeric; the resulting total affinity calculated on the given fasta. If a PFMatrixList has been passed then a named numeric vector with the affinities for all the PFMs. The vignette has all the details on the calculations (such as PFM to PWM conversion and pseudocounts).

## Examples

> library(JASPAR2014)
library(TFBSTools)
library(Biostrings)
pfm <- getMatrixByID(JASPAR2014,"MA0004.1")
\#\# The following sequence has a single perfect match
\#\# thus it gives the same results with all cutoff values.
sequence <- DNAString("CACGTG")
getSeqOccupancy (sequence, pfm, 0.1)
getSeqOccupancy (sequence, pfm, 1)
pfm2 <- getMatrixByID(JASPAR2014,"MA0005.1")
pfms <- PFMatrixList(pfm, pfm2)
names(pfms) <- c(name(pfm), name(pfm2))
\#\# This calculates total affinity for both the PFMatrixes.
getSeqOccupancy (sequence, pfms, 0)

## Index

```
* MatrixRider
    MatrixRider-package, 2
* getSeqOccupancy
    getSeqOccupancy, 2
bg, XMatrix-method, 2
DNAString,2
getSeqOccupancy,2
getSeqOccupancy,DNAString,PFMatrix, numeric-method
    (getSeqOccupancy), 2
getSeqOccupancy,DNAString, PFMatrixList, numeric-method
    (getSeqOccupancy), 2
getSeqOccupancy-methods
    (getSeqOccupancy), 2
MatrixRider (MatrixRider-package), 2
MatrixRider-package, 2
PFMatrix, 2
PFMatrixList, 2, 3
```

