# **CSAR**

# April 20, 2011

ChipseqScore Calculate read-enrichment scores for each nucleotide position

# **Description**

Calculate read-enrichment scores for each nucleotide position

# Usage

ChIPseqScore(control, sample, backg = 1, file = NA, norm =  $300 \times 10^6$ , test = '

# **Arguments**

control	data.frame structure obtained by mappedReads2Nhits
sample	data.frame structure obtained by mappedReads2Nhits
backg	Due low coverage in the control, there could be regions with no hits. Any region with a hit value lower than backg in the control will be set to the value of backg
file	Name of the file where you wan to save the results (if desired)
norm	Integer value. Number of hits will be reported by number of hits per norm nucleotides
test	Use a score based on the poisson distribution ("Poisson") or in the ratio ("Ratio")
times	To be memory efficient, CSAR will only upload to the RAM memory fragments of length times. A bigger value means more RAM memory needed but whole process will be faster
digits	Number of decimal digits used to report the score values

# **Details**

Different sequencing efforts yield different number of sequenced reads, for this reason the "number of hits" at each nucleotide position is normalized by the total number of nucleotides sequenced. Subsequently, the number of hits for the sample is normalize to have the same mean and variance than the control, for each chromosome independently or for the whole set of chromosomes (depending of the value of normEachChrInd). Due low coverage, there could be regions with no hits. Any region with a hit value lower than backg in the control will be set to the value of backg For each nucleotide position, a read-enrichment score will be calculated with the Poisson test, or with the ratio.

2 CSAR-package

#### Value

A list to be used for other functions of the CSAR package

chr Chromosme names
chrL Chromosme length (bp)

filenames Name of the files where the score values are storaged

digits Score values storaged on the files need to be divided by 10^digits

#### Author(s)

```
Jose M Muino, < jose.muino@wur.nl>
```

#### References

Muino et al. (submitted). Plant ChIP-seq Analyzer: An R package for the statistical detection of protein-bound genomic regions.

Kaufmann et al.(2009). Target genes of the MADS transcription factor SEPALLATA3: integration of developmental and hormonal pathways in the Arabidopsis flower. PLoS Biology; 7(4):e1000090.

#### See Also

CSAR-package

## **Examples**

```
##For this example we will use the a subset of the SEP3 ChIP-seq data (Kaufmann, 2009)
data("CSAR-dataset");
##We calculate the number of hits for each nucleotide posotion for the control and sample
```

##We calculate the number of hits for each nucleotide posotion for the control and sample
nhitsS<-mappedReads2Nhits(sampleSEP3\_test,file="sampleSEP3\_test",chr=c("CHR1v01212004"),c
nhitsC<-mappedReads2Nhits(controlSEP3\_test,file="controlSEP3\_test",chr=c("CHR1v01212004")</pre>

##We calculate a score for each nucleotide position
test<-ChIPseqScore(control=nhitsC,sample=nhitsS)</pre>

CSAR-package

Statistical tools for the analysis of ChIP-seq data

# Description

Statistical tools for ChIP-seq data analysis.

The package is oriented to plant organisms, and compatible with standard file formats in the plant research field.

CSAR-package 3

#### **Details**

Package: CSAR
Type: Package
Version: 1.0

Date: 2009-11-09 License: Artistic-2.0

LazyLoad: yes

#### Author(s)

Jose M Muino

Maintainer: Jose M Muino <jose.muino@wur.nl>

#### References

Muino et al. (submitted). Plant ChIP-seq Analyzer: An R package for the statistical detection of protein-bound genomic regions.

Kaufmann et al.(2009). Target genes of the MADS transcription factor SEPALLATA3: integration of developmental and hormonal pathways in the Arabidopsis flower. PLoS Biology; 7(4):e1000090.

#### **Examples**

##For this example we will use the a subset of the SEP3 ChIP-seq data (Kaufmann, 2009) data("CSAR-dataset");

##We calculate the number of hits for each nucleotide posotion for the control and sample
nhitsS<-mappedReads2Nhits(sampleSEP3\_test, file="sampleSEP3\_test", chr=c("CHR1v01212004"), c
nhitsC<-mappedReads2Nhits(controlSEP3\_test, file="controlSEP3\_test", chr=c("CHR1v01212004")</pre>

##We calculate a score for each nucleotide position
test<-ChIPseqScore(control=nhitsC,sample=nhitsS)</pre>

##We calculate the candidate read-enriched regions
win<-sigWin(test)</pre>

##We generate a wig file of the results to visualize tehm in a genome browser score2wig(test,file="test.wig")

##We calculate relative positions of read-enriched regions regarding gene position
d<-distance2Genes(win=win,gff=TAIR8\_genes\_test)</pre>

##We calculate table of genes with read-enriched regions, and their location genes<-genesWithPeaks(d)

##We calculate two sets of read-enrichment scores through permutation
permutatedWinScores(nn=1,sample=sampleSEP3\_test,control=controlSEP3\_test,fileOutput="test
permutatedWinScores(nn=2,sample=sampleSEP3\_test,control=controlSEP3\_test,fileOutput="test

###Next function will get all permutated score values generated by permutatedWinScores fu ##This represent the score distribution under the null hypotesis and therefore it can be 4 distance2Genes

```
nulldist<-getPermutatedWinScores(file="test",nn=1:2)</pre>
```

##From this distribution, several cut-off values can be calculated to control the error of ##Several functions in R can be used for this purpose.

##In this package we had implemented a simple method for the control of the error based of
getThreshold(winscores=win\$score,permutatedScores=nulldist,FDR=.01)

distance2Genes

Calculate relative positions of read-enriched regions regarding gene position

# Description

Calculate relative positions of read-enrichment regions regarding gene position

#### Usage

```
distance2Genes(win, qff, t = -\log(0.05), d1 = -3000, d2 = 1000)
```

# **Arguments**

win	Data.frame structure obtained with the function sigWin
gff	Data.frame structure obtained after loading a desired gff file
t	Integer. Only distances of read-enriched regions with a score bigger than $\ensuremath{\text{t}}$ will be considered
d1	Negative integer. Minimum relative position regarding the start of the gene to be considered
d2	Positive integer. Maximum relative position regarding the end of the gene to be considered

### Value

data.frame structure where each row represents one relative position, and each column being:

peakName read-enriched region name
p1 relative position regarding the start of the gene
p2 relative position regarding the end of the gene
gene name of the gene
le length (bp) of the gene

#### Author(s)

Jose M Muino, < jose.muino@wur.nl>

#### References

Muino et al. (submitted). Plant ChIP-seq Analyzer: An R package for the statistical detection of protein-bound genomic regions.

Kaufmann et al.(2009). Target genes of the MADS transcription factor SEPALLATA3: integration of developmental and hormonal pathways in the Arabidopsis flower. PLoS Biology; 7(4):e1000090.

genesWithPeaks 5

#### See Also

genesWithPeaks, CSAR-package

# **Examples**

```
##For this example we will use the a subset of the SEP3 ChIP-seq data (Kaufmann, 2009)
data("CSAR-dataset");
##We calculate the number of hits for each nucleotide posotion for the control and sample
nhitsS<-mappedReads2Nhits(sampleSEP3_test,file="sampleSEP3_test",chr=c("CHR1v01212004"),c
nhitsC<-mappedReads2Nhits(controlSEP3_test,file="controlSEP3_test",chr=c("CHR1v01212004")

##We calculate a score for each nucleotide position
test<-ChIPseqScore(control=nhitsC,sample=nhitsS)

##We calculate the candidate read-enriched regions
win<-sigWin(test)

##We calculate relative positions of read-enriched regions regarding gene position
d<-distance2Genes(win=win,gff=TAIR8_genes_test)</pre>
```

genesWithPeaks

Provide table of genes with read-enriched regions, and their location

# Description

Provide table of genes with read-enriched regions, and their location

# Usage

```
genesWithPeaks(distances)
```

# **Arguments**

distances data

data.frame structure obtained by distances2Genes

#### **Details**

This function report for each gene, the maximum peak score in different regions near of the gene. The input of the function is the distances between genes and peaks calculated by distance2Genes

# Value

data.frame structure with each coloumn being:

name of the gene

max3kb1kb maximum score value for the region 3Kb upstream to 1Kb dowstream u3000 maximum score value for the region 3Kb upstream to 2Kb upstream

u2000	maximum score value for the region 2Kb upstream to 1Kb upstream
u1000	maximum score value for the region 1Kb upstream to 0Kb upstream
d0	maximum score value for the region 0Kb upstream to 0Kb dowstream
d1000	maximum score value for the region 0Kb dowstream to 1Kb dowstream

#### Author(s)

Jose M Muino, < jose.muino@wur.nl>

#### References

Muino et al. (submitted). Plant ChIP-seq Analyzer: An R package for the statistical detection of protein-bound genomic regions.

Kaufmann et al.(2009). Target genes of the MADS transcription factor SEPALLATA3: integration of developmental and hormonal pathways in the Arabidopsis flower. PLoS Biology; 7(4):e1000090.

#### See Also

distance2Genes,CSAR-package

# **Examples**

```
##For this example we will use the a subset of the SEP3 ChIP-seq data (Kaufmann, 2009)
data("CSAR-dataset");
##We calculate the number of hits for each nucleotide posotion for the control and sample
nhitsS<-mappedReads2Nhits(sampleSEP3_test,file="sampleSEP3_test",chr=c("CHR1v01212004"),c
nhitsC<-mappedReads2Nhits(controlSEP3_test,file="controlSEP3_test",chr=c("CHR1v01212004")

##We calculate a score for each nucleotide position
test<-ChIPseqScore(control=nhitsC,sample=nhitsS)

##We calculate the candidate read-enriched regions
win<-sigWin(test)

##We calculate relative positions of read-enriched regions regarding gene position
d<-distance2Genes(win=win,gff=TAIR8_genes_test)

##We calculate table of genes with read-enriched regions, and their location
genes<-genesWithPeaks(d)</pre>
```

```
getPermutatedWinScores
```

Obtain the read-enrichment score distribution under the null hypothesis

# Description

Obtain the read-enrichment score distribution under the null hypothesis

*getPermutatedWinScores* 

## Usage

```
getPermutatedWinScores(file, nn)
```

# **Arguments**

file Name of the file generated by permutatedWinScores

nn ID for the multiple permutation process

#### Value

Numeric vector of score values under permutation

#### Author(s)

```
Jose M Muino, < jose.muino@wur.nl>
```

#### References

Muino et al. (submitted). Plant ChIP-seq Analyzer: An R package for the statistical detection of protein-bound genomic regions.

Kaufmann et al.(2009). Target genes of the MADS transcription factor SEPALLATA3: integration of developmental and hormonal pathways in the Arabidopsis flower. PLoS Biology; 7(4):e1000090.

#### See Also

CSAR-package, permutatedWinScores

# **Examples**

```
##For this example we will use the a subset of the SEP3 ChIP-seq data (Kaufmann, 2009)
data("CSAR-dataset");
##We calculate the number of hits for each nucleotide posotion for the control and sample
nhitsS<-mappedReads2Nhits(sampleSEP3_test,file="sampleSEP3_test",chr=c("CHR1v01212004"),c
nhitsC<-mappedReads2Nhits(controlSEP3_test,file="controlSEP3_test",chr=c("CHR1v01212004")</pre>
```

##We calculate two sets of read-enrichment scores through permutation
permutatedWinScores(nn=1,sample=sampleSEP3\_test,control=controlSEP3\_test,fileOutput="test
permutatedWinScores(nn=2,sample=sampleSEP3\_test,control=controlSEP3\_test,fileOutput="test

###Next function will get all permutated score values generated by permutatedWinScores fu
##This represent the score distribution under the null hypotesis and therefore it can be
nulldist<-getPermutatedWinScores(file="test",nn=1:2)</pre>

8 getThreshold

getThreshold	Calculate the threshold value corresponding to control FDR at a desired level
	strea tevet

#### **Description**

Calculate the threshold value corresponding to control FDR at a desired level

# Usage

```
getThreshold(winscores, permutatedScores, FDR)
```

#### **Arguments**

winscores Numeric vector with score values obtained from the sigWin function permutatedScores

Numeric vector with the permutated read-enrichment score values

FDR Numeric value with the desired FDR control

#### **Details**

This is a very simple function to obtain the threshold value of our test statistic controlling FDR at a desired level. Other functions implemented in R (eg: multtest) could be more sophisticated. Basically, for each possible threshold value, the proportion of error type I is calculated assuming that the permutated score distribution is a optimal estimation of the score distribution under the null hypothesis. This is, the proportion of permutated scores exceding the considered threshold value is used as an estimation of the error type I of our statistic. FDR is obtained as the ratio of the proportion of error type I by the proportion of significant tests.

## Value

A table with the columns being:

threshold The threshold value

p-value The p-value obtained from the permutated score ditribution

FDR The FDR control obtained using threshold

# Author(s)

```
Jose M Muino, <jose.muino@wur.nl>
```

#### References

Muino et al. (submitted). Plant ChIP-seq Analyzer: An R package for the statistical detection of protein-bound genomic regions.

Kaufmann et al.(2009). Target genes of the MADS transcription factor SEPALLATA3: integration of developmental and hormonal pathways in the Arabidopsis flower. PLoS Biology; 7(4):e1000090.

#### See Also

CSAR-package,getPermutatedWinScores, sigWin

IoadMappedReads 9

#### **Examples**

```
##For this example we will use the a subset of the SEP3 ChIP-seq data (Kaufmann, 2009)
data("CSAR-dataset");
\#\#We calculate the number of hits for each nucleotide posotion for the control and sample
nhitsC<-mappedReads2Nhits(controlSEP3_test,file="controlSEP3_test",chr=c("CHR1v01212004")
##We calculate a score for each nucleotide position
test<-ChIPseqScore (control=nhitsC, sample=nhitsS)
##We calculate the candidate read-enriched regions
win<-sigWin(test)
##We calculate two sets of read-enrichment scores through permutation
permutatedWinScores(nn=1,sample=sampleSEP3_test,control=controlSEP3_test,fileOutput="test
permutatedWinScores(nn=2,sample=sampleSEP3_test,control=controlSEP3_test,fileOutput="test
###Next function will get all permutated score values generated by permutatedWinScores fu
##This represent the score distribution under the null hypotesis and therefore it can be
nulldist<-getPermutatedWinScores(file="test",nn=1:2)</pre>
##From this distribution, several cut-off values can be calculated to control the error of
```

##In this package we had implemented a simple method for the control of the error based of

loadMappedReads

Load mapped reads

##Several functions in R can be used for this purpose.

getThreshold(winscores=win\$score,permutatedScores=nulldist,FDR=.01)

#### Description

This function load the output file of a read mapping software (eg:SOAP)

## Usage

```
loadMappedReads(file, format = "SOAP", header = FALSE)
```

## **Arguments**

file File name to load

format Format of the file. "SOAP" for the output of the soap software and "MAQ" for

the maq software. Other user formats can be provided as a character vector for the file column names. Columns named: "Nhits", "lengthRead", "strand",

"chr", and "pos" are needed.

header Logical value indicating if the first line of the file should be skipped (TRUE) or

not (FALSE)

#### Value

data.frame structure that can be used by mappedReads2Nhits

10 mappedReads2Nhits

#### Author(s)

```
Jose M Muino, < jose.muino@wur.nl>
```

#### References

Muino et al. (submitted). Plant ChIP-seq Analyzer: An R package for the statistical detection of protein-bound genomic regions.

Kaufmann et al.(2009). Target genes of the MADS transcription factor SEPALLATA3: integration of developmental and hormonal pathways in the Arabidopsis flower. PLoS Biology; 7(4):e1000090.

# See Also

CSAR-package

# **Examples**

```
##We load the mapped reads:
#sample<-loadMappedReads(file=file,format="SOAP",w=300,header=F)
##where file is the name and path of the output file of the mapping process.</pre>
```

mappedReads2Nhits Calculate number of overlapped extended reads per nucleotide position

# Description

Calculate number of overlapped extended reads per nucleotide position

# Usage

```
mappedReads2Nhits(input, file , chr = c("chr1", "chr2", "chr3", "chr4", "chr5"),
```

### **Arguments**

input	data loaded with loadMappedReads or an AlignedRead object from the Short-Read package
file	Name of the file where the results will be saved. If NA the results will not be saved in a file.
chr	Character vector containing the chromosome names as identified on input.
chrL	Numeric vector containing the length (bp) of the chromosomes. It should be in the same order than ${\tt chr}$
W	Integer corresponding to the desired length of the extended reads. An advised value will be the average fragment length of the DNA submitted to sequence (usually 300 bp).

considerStrand

Character value.

"Minimum"=>Default value. Report the minimum number of hits at each nucleotide position for both strands.

mappedReads2Nhits 11

"Foward"=> Report the number of hits at each nucleotide position for the "foward" strands (the one denoted as "+" in q).

"Reverse"=>Report the number of hits at each nucleotide position for the "reverse" strands (the one denoted as "-" in q).

"Sum"=>Report the sum of number of hits at each nucleotide position for both strands.

uniquelyMapped

Logic value, If TRUE, only consider uniquely mapped reads.

uniquePosition

Logic value. If TRUE, only consider reads mapped in different positions.

#### Value

A list to be used for other functions of the CSAR package

chr	Chromosme names
chrL	Chromosme length (bp)
chrL_0	Number of nucleotide positions with at least one extended read
chrL_0	Number of nucleotide positions with at least one extended read
filenames	Name of the files where the Nhits values are storaged
c1	Sum of all the Nhits values for each chromosome
c2	Sum of all the Nhits square values for each chromosome

# Author(s)

```
Jose M Muino, < jose.muino@wur.nl>
```

## References

Muino et al. (submitted). Plant ChIP-seq Analyzer: An R package for the statistical detection of protein-bound genomic regions.

Kaufmann et al.(2009). Target genes of the MADS transcription factor SEPALLATA3: integration of developmental and hormonal pathways in the Arabidopsis flower. PLoS Biology; 7(4):e1000090.

## See Also

CSAR-package

# **Examples**

```
#For this example we will use the a subset of the SEP3 ChIP-seq data (Kaufmann, 2009)
data("CSAR-dataset");
```

 12 permutatedWinScores

permutatedWinScores

Calculate scores for permutated read-enriched regions

#### **Description**

Calculate scores for permutated read-enriched regions

#### **Usage**

```
permutatedWinScores(nn = 1, control, sample, fileOutput, chr = c("chr1", "chr2",
```

#### **Arguments**

ID to identify each permutation nn control data.frame structure obtained by loading the mapped reads with the function LoadMappedReads() data.frame structure obtained by loading the mapped reads with the function sample LoadMappedReads() fileOutput Name of the file were the results will be written Character vector containing the chromosome names as identified on q. chr Numeric vector containing the length (bp) of the chromosomes. It should be in chr<sub>L</sub> the same order than chr Integer corresponding to the desired length of the extended reads.

considerStrand

Character value.

"Minimum"=>Default value. Report the minimum number of hits at each nucleotide position for both strands.

"Foward"=> Report the number of hits at each nucleotide position for the "foward" strands (the one denoted as "+" in q).

"Reverse"=>Report the number of hits at each nucleotide position for the "reverse" strands (the one denoted as "-" in q).

"Sum"=>Report the sum of number of hits at each nucleotide position for both strands.

uniquelyMapped

Logic value, If TRUE, only consider unquely mapped reads.

uniquePosition

Logic value. If TRUE, only consider reads mapped in different positions.

Integer value. Number of hits will be reported by number of hits per norm norm

nucleotides

Any region with a hit value lower than backg in the control will be set to backg

the value of backg

Numeric value. Read-enriched regions are calculated as genomic regions with t

score values bigger than t

Integer value. The maximum gap allowed between regions. Regions that are

less than g bps away will be merged.

permutatedWinScores 13

times To be m	emory efficient, CSAR v	will only upload to the RA	M memory fragments
---------------	-------------------------	----------------------------	--------------------

of length times. A bigger value means more RAM memory needed but whole

process will be faster

digits Number of decimal digits used to report the score values

test Use a score based on the poisson distribution ("Poisson") or in the ratio ("Ratio")

#### **Details**

The parameter values should be the same than the one used in sigWin, ChIPseqScore, and mappedReads2Nhits. The label "control" and "sample" is asigned to each read to identify from which group they came. Labels are randomly permutated, and read-enriched regions for this new permuated dataset are calculated.

## Value

The file filePutput is created with the next columns:

chr Chromosome name

start Start of the read-enriched region end End of the read-enriched region

posPeak Position of the maximum score value on the read-enriched region

score Maximum score value on the read-enriched region

length Read-enriched region length

## Author(s)

Jose M Muino, < jose.muino@wur.nl>

# References

Muino et al. (submitted). Plant ChIP-seq Analyzer: An R package for the statistcal detection of protein-bound genomic regions.

Kaufmann et al.(2009). Target genes of the MADS transcription factor SEPALLATA3: integration of developmental and hormonal pathways in the Arabidopsis flower. PLoS Biology; 7(4):e1000090.

#### See Also

CSAR-package,getPermutatedWinScores

#### **Examples**

```
##For this example we will use the a subset of the SEP3 ChIP-seq data (Kaufmann, 2009)
data("CSAR-dataset");
##We calculate the number of hits for each nucleotide posotion for the control and sample
nhitsS<-mappedReads2Nhits(sampleSEP3_test,file="sampleSEP3_test",chr=c("CHR1v01212004"),c</pre>
```

```
nhitsC<-mappedReads2Nhits(controlSEP3_test, file="controlSEP3_test", chr=c("CHR1v01212004")
##We calculate two sets of read-enrichment scores through permutation</pre>
```

```
permutatedWinScores(nn=1, sample=sampleSEP3_test, control=controlSEP3_test, fileOutput="test permutatedWinScores(nn=2, sample=sampleSEP3_test)]
```

14 score2wig

sampleSEP3_test Partial dataset of a ChIP-seq experiment
--

# Description

Partial dataset of a Solexa DNA library obtained from a ChIP-seq experiment in Arabidopsis

#### Source

Kaufmann et al. (2009) Target Genes of the MADS Transcription Factor SEPALLATA3: Integration of Developmental and Hormonal Pathways in the \$Arabidopsis\$ Flower. PLoS Biol 7:e1000090

# **Examples**

```
data(CSAR-dataset)
```

score2wig	Save the read-enrichment scores at each nucleotide position in a .wig
	file format

# **Description**

Save the read-enrichment scores at each nucleotide position in a .wig file format that can be visualize by a genome browser (eg: Integrated Genome Browser)

# Usage

```
score2wig(experiment, file, t = 3, times = 1e6,description="CSAR track", name="C
```

# Arguments

experiment	Output of the function ChIPseqScore
file	Name of the output .wig file
t	Only nucleotide positions with a read-enrichment score bigger than $\ensuremath{\mathtt{t}}$ will be reported
times	To be memory efficient, CSAR will only upload to the RAM memory fragments of length times. A bigger value means more RAM memory needed but whole process will be faster
description	Character. It adds a description to the wig file. The description will be shown by the genome browser used to visualize the wig file.
name	Character. It adds a wig to the wig file. The name will be shown by the genome browser used to visualize the wig file.

# Value

None. Results are printed in a file

sigWin 15

#### Author(s)

```
Jose M Muino, < jose.muino@wur.nl>
```

#### References

Muino et al. (submitted). Plant ChIP-seq Analyzer: An R package for the statistical detection of protein-bound genomic regions.

Kaufmann et al.(2009). Target genes of the MADS transcription factor SEPALLATA3: integration of developmental and hormonal pathways in the Arabidopsis flower. PLoS Biology; 7(4):e1000090.

#### See Also

CSAR-package

## **Examples**

```
##For this example we will use the a subset of the SEP3 ChIP-seq data (Kaufmann, 2009)
data("CSAR-dataset");
##We calculate the number of hits for each nucleotide position for the control and sample
nhitsS<-mappedReads2Nhits(sampleSEP3_test, file="sampleSEP3_test", chr=c("CHR1v01212004"), c
nhitsC<-mappedReads2Nhits(controlSEP3_test, file="controlSEP3_test", chr=c("CHR1v01212004"))
##Since we will not need the raw data anymore, we could delete it from the RAM memory
rm(sampleSEP3_test, controlSEP3_test); gc(verbose=FALSE)
##We calculate a score for each nucleotide position
test<-ChIPseqScore(control=nhitsC, sample=nhitsS)

##We generate a wig file of the results to visualize them in a genome browser
score2wig(test, file="test.wig")</pre>
```

sigWin

Calculate regions of read-enrichment

#### **Description**

Calculate regions of read-enrichment

# Usage

```
sigWin(experiment, t = -log(0.05), g = 100)
```

#### **Arguments**

experiment	Output of the function ChIPseqScore
t	Numeric value. Read-enriched regions are calculated as genomic regions with score values bigger than $\ensuremath{\text{t}}$
g	Integer value. The maximum gap allowed between regions. Regions that are less than g bps away will be merged.

16 sigWin

#### Value

A data.frame structure with the columns being:

chr Chromosome name

start Start of the read-enriched region end End of the read-enriched region

posPeak Position of the maximum score value on the read-enriched region

score Maximum score value on the read-enriched region

length Read-enriched region length

#### Author(s)

Jose M Muino, < jose.muino@wur.nl>

#### References

Muino et al. (submitted). Plant ChIP-seq Analyzer: An R package for the statistical detection of protein-bound genomic regions.

Kaufmann et al.(2009). Target genes of the MADS transcription factor SEPALLATA3: integration of developmental and hormonal pathways in the Arabidopsis flower. PLoS Biology; 7(4):e1000090.

# See Also

CSAR-package

# **Examples**

```
##For this example we will use the a subset of the SEP3 ChIP-seq data (Kaufmann, 2009)
data("CSAR-dataset");
```

##We calculate the number of hits for each nucleotide posotion for the control and sample
nhitsS<-mappedReads2Nhits(sampleSEP3\_test, file="sampleSEP3\_test", chr=c("CHR1v01212004"), c
nhitsC<-mappedReads2Nhits(controlSEP3\_test, file="controlSEP3\_test", chr=c("CHR1v01212004")</pre>

```
##We calculate a score for each nucleotide position
test<-ChIPseqScore(control=nhitsC,sample=nhitsS)</pre>
```

##We calculate the candidate read-enriched regions
win<-sigWin(test)</pre>

# **Index**

```
*Topic datasets
   sampleSEP3_test, 14
ChIPseqScore, 1
controlSEP3_test
       (sampleSEP3_test), 14
CSAR-package, 2
distance2Genes,4
genesWithPeaks, 5
getPermutatedWinScores, 6
{\tt getThreshold}, {\tt 8}
LoadBinCSAR (score2wig), 14
loadMappedReads, 9
mappedReads2Nhits, 10
mappedReads2Nhits_chr
       (mappedReads2Nhits), 10
permutatedWinScores, 12
pos2Nhits (mappedReads2Nhits), 10
pos2Nhits_old
       (mappedReads2Nhits), 10
sampleSEP3_test, 14
score2wig, 14
score_chr (ChIPseqScore), 1
sigWin, 15
sigWin_chr(sigWin), 15
TAIR8_genes_test
       (sampleSEP3_test), 14
```