

Package ‘MEDIPSData’

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Type Package

Title Example data for the MEDIPS package.

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Description Data for the MEDIPS package, consisting of chromosome 22 MeDIP and control/Input sample data from DNA methylation analysis of human embryonic stem cells and mapped to the human genome hg19 using bowtie.

License GPL (>= 2)

LazyLoad yes

biocViews ExperimentData, Genome, SequencingData

NeedsCompilation no

R topics documented:

CS	2
DE_Input	2
DE_MeDIP	3
hESCs_Input	3
hESCs_MeDIP	4
mart_gene	5
resultTable	5

Index

7

CS

*COUPLING SET***Description**

This is a CpG coupling set generated by the MEDIPS package based on the human chromosome 22 (hg19) and with a window size of 100bp.

Usage

```
data(CS)
```

Examples

```
## Not run:  
data(CS)  
library(MEDIPS)  
CS  
  
## End(Not run)
```

DE_Input

*control data set from definitive endoderm***Description**

This is a MEDIPS SET object created from Input-seq control data derived from definitive endoderm as presented by Chavez et al. 2010. The parameter settings are: BSgenome= "BSgenome.Hsapiens.UCSC.hg19" extend= 300 shift= 0 uniq= T ws= 100 chr.select= "chr22"

Usage

```
data(DE_Input)
```

References

Chavez, L., Jozefczuk, J., Grimm, C., Dietrich, J., Timmermann, B., Herwig, R., Adjaye, J. (2010): Computational analysis of genome-wide DNA methylation during the differentiation of human embryonic stem cells along the endodermal lineage, *Genome Research*. 20(10):1441-50

Examples

```
## Not run:  
data(DE_Input)  
library(MEDIPS)  
DE_Input  
  
## End(Not run)
```

DE_MeDIP	<i>Concatenated set of three MeDIP-seq data sets (replicates) from definitive endoderm</i>
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Description

This is a concatenated set of three MEDIPS SET objects created from MeDIP-seq data derived from definitive endoderm as presented by Chavez et al. 2010. The parameter settings are: BSgenome= "BSgenome.Hsapiens.UCSC.hg19" extend= 300 shift= 0 uniq= T ws= 100 chr.select= "chr22"

Usage

```
data(DE_MeDIP)
```

References

Chavez, L., Jozefczuk, J., Grimm, C., Dietrich, J., Timmermann, B., Herwig, R., Adjaye, J. (2010): Computational analysis of genome-wide DNA methylation during the differentiation of human embryonic stem cells along the endodermal lineage, *Genome Research.* 20(10):1441-50

Examples

```
## Not run:  
data(DE_MeDIP)  
library(MEDIPS)  
DE_MeDIP  
  
## End(Not run)
```

hESCs_Input	<i>control data set from human embryonic stem cells</i>
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Description

This is a MEDIPS SET object created from Input-seq control data derived from human embryonic stem cells as presented by Chavez et al. 2010. The parameter settings are: BSgenome= "BSgenome.Hsapiens.UCSC.hg19" extend= 300 shift= 0 uniq= T ws= 100 chr.select= "chr22"

Usage

```
data(hESCs_Input)
```

References

Chavez, L., Jozefczuk, J., Grimm, C., Dietrich, J., Timmermann, B., Herwig, R., Adjaye, J. (2010): Computational analysis of genome-wide DNA methylation during the differentiation of human embryonic stem cells along the endodermal lineage, *Genome Research.* 20(10):1441-50

Examples

```
## Not run:
data(hESCs_Input)
library(MEDIPS)
hESCs_Input

## End(Not run)
```

hESCs_MeDIP

Concatenated set of three MeDIP-seq data sets (replicates) from human embryonic stem cells

Description

This is a concatenated set of three MEDIPS SET objects created from MeDIP-seq data derived from human embryonic stem cells as presented by Chavez et al. 2010. The parameter settings are: BSgenome= "BSgenome.Hsapiens.UCSC.hg19" extend= 300 shift= 0 uniq= T ws= 100 chr.select= "chr22"

Usage

```
data(hESCs_MeDIP)
```

References

Chavez, L., Jozefczuk, J., Grimm, C., Dietrich, J., Timmermann, B., Herwig, R., Adjaye, J. (2010): Computational analysis of genome-wide DNA methylation during the differentiation of human embryonic stem cells along the endodermal lineage, *Genome Research.* 20(10):1441-50

Examples

```
## Not run:
data(hESCs_MeDIP)
library(MEDIPS)
hESCs_MeDIP

## End(Not run)
```

mart_gene

An annotation object generated by accessing biomaRt using the MEDIPS.getAnnotation function of the MEDIPS package.

Description

This is an annotation object generated by accessing biomaRt using the MEDIPS.getAnnotation function of the MEDIPS package: `mart_gene = MEDIPS.getAnnotation(mart="ensembl", dataset=c("hsapiens_gene_ensembl", annotation=c("GENE"), chr=22))` The annotation object contains genomic coordinates of human genes on chromosome 22.

Usage

```
data(mart_gene)
```

Examples

```
## Not run:  
data(mart_gene)  
data(resultTable)  
library(MEDIPS)  
resultTable = MEDIPS.setAnnotation(regions=resultTable, annotation=mart_gene)  
  
## End(Not run)
```

resultTable

A result table as returned by the MEDIPS.meth function of the MEDIPS package

Description

This is a result table as returned by the MEDIPS.meth function using the following command: `mr.edgeR = MEDIPS.meth(MSet1=hESCs, MSet2=DE, CSet=CS, ISet1=hESCs.Input, ISet2=DE.Input, p.adj="bonferroni", diff.method="edgeR", prob.method="poisson", CNV=F, MeDIP=T)` where hESCs, DE, and CS are data objects included in this data package.

Usage

```
data(resultTable)
```

Examples

```
## Not run:  
data(resultTable)  
library(MEDIPS)  
mr.edgeR.s = MEDIPS.selectSig(results=mr.edgeR, p.value=0.05, adj=T, ratio=NULL, bg.counts=NULL, CNV=F)  
mr.edgeR.s  
  
## End(Not run)
```

Index

*Topic **datasets**

CS, [2](#)
DE_Input, [2](#)
DE_MeDIP, [3](#)
hESCs_Input, [3](#)
hESCs_MeDIP, [4](#)
mart_gene, [5](#)
resultTable, [5](#)

CS, [2](#)

DE_Input, [2](#)
DE_MeDIP, [3](#)

hESCs_Input, [3](#)
hESCs_MeDIP, [4](#)

mart_gene, [5](#)

resultTable, [5](#)