

Package ‘sesameData’

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Type Package

Title Supporting Data for SeSAmE Package

Description Provides supporting annotation and test data for SeSAmE package. This includes chip tango addresses, mapping information, performance annotation, and trained predictor for Infinium array data. This package provides user access to essential annotation data for working with many generations of the Infinium DNA methylation array. Current we support human array (HM27, HM450, EPIC), mouse array (MM285) and the Horvath-MethylChip40 (Mammal40) array.

Version 1.25.0

License Artistic-2.0

Depends R (>= 4.2.0), ExperimentHub, AnnotationHub

Imports utils, readr, stringr, GenomicRanges, S4Vectors, IRanges, GenomeInfoDb

Suggests BiocGenerics, sesame, testthat, knitr, rmarkdown

biocViews ExperimentData, MicroarrayData, Genome, ExperimentHub, MethylationArrayData

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BugReports <https://github.com/zwdzwd/sesame/issue>

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build_GENCODE_gtf	<i>build GENCODE gtf</i>
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Description

build GENCODE gtf

Usage

build_GENCODE_gtf(x)

Arguments

x	GENCODE ftp url
---	-----------------

Value

GRangesList

df_master

*Master data frame for all object to cache***Description**

This is an internal object which will be updated on every new release library(ExperimentHub) `eh <- query(ExperimentHub(localHub=FALSE), c("sesameData", "v1.13.1")) data.frame(name=eh$title, eh=names(eh))`

Details

Cache location is default to `/Users/zhouw3/Library/Caches/org.R-project.R/R/ExperimentHub/`

Value

master sheet of sesameData objects

extend

*Extend a GRanges***Description**

source: <https://support.bioconductor.org/p/78652/>

Usage

```
extend(gr, upstream = 0, downstream = 0)
```

Arguments

gr	a GenomicRanges::GRanges
upstream	distance to expand upstream
downstream	distance to expand downstream

Value

a GenomicRanges::GRanges

```
inferPlatformFromProbeIDs
      infer platform from Probe_IDs
```

Description

infer platform from Probe_IDs

Usage

```
inferPlatformFromProbeIDs(Probe_IDs, silent = FALSE)
```

Arguments

Probe_IDs	probe IDs
silent	suppress message

Value

a platform code

Examples

```
sesameDataCache("probeIDSignature")
inferPlatformFromProbeIDs(c("cg14620903", "cg22464003"))
```

```
sesameDataCache      Cache SeSAmE data
```

Description

Cache SeSAmE data

Usage

```
sesameDataCache(data_titles = NULL)
```

Arguments

data_titles	data to cache, if not given will cache all
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Value

TRUE

Examples

```
sesameDataCache("genomeInfo.hg38")
```

sesameDataCacheAll	<i>Cache all SeSAmE data</i>
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Description

Cache all SeSAmE data

Usage

```
sesameDataCacheAll(data_titles = NULL)
```

Arguments

data_titles data to cache, if not given will cache all

Value

TRUE

Examples

```
sesameDataCache("genomeInfo.hg38")
```

sesameDataGet	<i>Get SeSAmE data</i>
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Description

Get SeSAmE data

Usage

```
sesameDataGet(title, verbose = FALSE)
```

Arguments

title title of the data
verbose whether to output ExperimentHub message

Value

data object

Examples

```
sesameDataCache("EPIC.1.SigDF")  
EPIC.1.SigDF <- sesameDataGet('EPIC.1.SigDF')
```

sesameDataGet_checkEnv

Check whether the title exists in cacheEnv

Description

Check whether the title exists in cacheEnv

Usage

```
sesameDataGet_checkEnv(title)
```

Arguments

title the title to check

Value

the data associated with the title or NULL if title doesn't exist

sesameDataGet_resetEnv

Empty cache environment to free memory

Description

When this function is called sesameDataGet will retrieve all data from disk again instead of using the in-memory cache, i.e., sesameData:::cacheEnv.

Usage

```
sesameDataGet_resetEnv()
```

Details

Note this is different from sesameDataClearHub which empties the ExperimentHub on disk.

Value

gc() output

Examples

```
sesameDataGet_resetEnv()
```

sesameDataHas	<i>Whether sesameData has</i>
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Description

Whether sesameData has

Usage

```
sesameDataHas(data_titles)
```

Arguments

data_titles data titles to check

Value

a boolean vector the same length as data_titles

Examples

```
sesameDataHas(c("EPIC.address", "EPIC.address.Nonexist"))
```

sesameDataList	<i>List all SeSAmE data</i>
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Description

List all SeSAmE data

Usage

```
sesameDataList(filter = NULL, full = FALSE)
```

Arguments

filter keyword to filter title, optional
full whether to display all columns

Value

all titles from SeSAmE Data

Examples

```
sesameDataList("KYCG")
```

 sesameData_annoProbes *Annotate Probes by Probe ID*

Description

Columns in the manifests will be added to the annotation. Please note that if unfound, the annotation will be NA. The probe will always be kept in the output.

Usage

```
sesameData_annoProbes(
  Probe_IDs,
  regs = NULL,
  collapse = TRUE,
  chooseOne = FALSE,
  column = NULL,
  sep = ", ",
  return_ov_probes = FALSE,
  return_ov_features = FALSE,
  out_name = NULL,
  platform = NULL,
  genome = NULL,
  silent = FALSE
)
```

Arguments

Probe_IDs	a character vector of probe IDs
regs	a GenomicRanges::GRanges object against which probes will be annotated, default to genes if not given
collapse	whether to collapse multiple regs into one
chooseOne	choose an arbitrary annotation if multiple exist default to FALSE. which concatenates all with ", "
column	which column in regs to annotate, if not given return all overlapping probes
sep	the delimiter for collapsing
return_ov_probes	if TRUE will return overlapping probes in a GRanges object.
return_ov_features	if TRUE will return overlapping features in a GRanges object.
out_name	column header of the annotation, use column if not given
platform	EPIC, MM285 etc. will infer from Probe_IDs if not given
genome	hg38, mm10, ... will infer if not given. For additional mapping, download the GRanges object from http://zwdzwd.github.io/InfiniumAnnotation and provide the following argument ..., genome = sesameAnno_buildManifestGRanges("downloaded_file"),... to this function.
silent	suppress messages

Value

a GRanges with annotated column. If a probe has no overlap with regs, it will be included in the results with NA. But if a probe is not included in the manifest (due to mappability), it won't be included in the results.

Examples

```
library(GenomicRanges)
sesameDataCache(c(
  "genomeInfo.mm10", "MM285.address",
  "genomeInfo.hg38", "Mamma140.address"))

regs = sesameData_getTxnGRanges("mm10")
Probe_IDs = names(sesameData_getManifestGRanges("MM285"))
anno = sesameData_annoProbes(Probe_IDs, promoters(regs), column="gene_name")

## get all genes associated with a probe set
genes = sesameData_getTxnGRanges("hg38", merge2gene = TRUE)
anno = sesameData_annoProbes(
  c("cg14620903", "cg22464003"), genes, return_ov_features=TRUE)
```

sesameData_check_genome

Find genome assembly version(s) supported for a platform

Description

Find genome assembly version(s) supported for a platform

Usage

```
sesameData_check_genome(genome, platform)
```

Arguments

genome	mm10, hg38, ..., or NULL
platform	HM27, HM450, EPIC, EPICv2, MSA, ...

Value

genome as string

Examples

```
sesameData_check_genome(NULL, "Mamma140")
```

```
sesameData_check_platform
```

Check platform code

Description

Note: custome platforms lead to error here.

Usage

```
sesameData_check_platform(platform = NULL, probes = NULL, silent = TRUE)
```

Arguments

platform	input platform
probes	probes by which the platform may be guessed
silent	suppress message

Value

platform code

Examples

```
sesameData_check_platform("HM450")
```

```
sesameData_getGenomeInfo
```

Get genome info files

Description

Get genome info files

Usage

```
sesameData_getGenomeInfo(genome)
```

Arguments

genome	hg38, mm10, or GRanges with a metadata(genome)[["genome"]]
--------	--

Value

a list of genome info files

Examples

```
sesameDataCache("genomeInfo.hg38")
res <- sesameData_getGenomeInfo("hg38")
```

```
sesameData_getManifestGRanges
  get Infinium manifest GRanges
```

Description

Note that some unaligned probes are not included. For full manifest, please visit <http://zwdzwd.github.io/InfiniumAnnotation>

Usage

```
sesameData_getManifestGRanges(platform, genome = NULL)
```

Arguments

platform	Mammal40, MM285, EPIC, and HM450
genome	hg38, mm10, ... will infer if not given. For additional mapping, download the GRanges object from http://zwdzwd.github.io/InfiniumAnnotation and provide the following argument ..., genome = sesameAnno_buildManifestGRanges("downloaded_file"),... to this function.

Value

GRanges

Examples

```
sesameDataCache("Mammal40.address")
res <- sesameData_getManifestGRanges("Mammal40")
```

```
sesameData_getProbesByGene
  Get Probes by Genes or Gene Promoters
```

Description

Get probes mapped to a gene. All transcripts for the gene are considered. The function takes a gene name as appears in UCSC RefGene database. The platform and reference genome build can be changed with 'platform' and 'genome' options. The function returns a vector of probes that falls into the given gene.

Usage

```
sesameData_getProbesByGene(  
  gene_name = NULL,  
  platform = NULL,  
  promoter = FALSE,  
  upstream = 1500,  
  downstream = 1500,  
  genome = NULL  
)
```

Arguments

gene_name	gene name, if NULL return all genes
platform	EPIC or HM450
promoter	if TRUE, use TSS instead of the whole gene
upstream	number of bases to expand upstream of target gene
downstream	number of bases to expand downstream of target gene
genome	hg38 or hg19

Value

GRanges containing probes that fall into the given gene

Examples

```
## download needed data  
sesameDataCache(c("Mammal40.address", "genomeInfo.hg38"))  
  
## get all probes overlapping with DNMT3A  
probes <- sesameData_getProbesByGene(  
  'DNMT3A', "Mammal40", upstream=500, downstream=500)  
  
## get the promoter-associated probes  
probes <- sesameData_getProbesByGene('DNMT3A', "Mammal40", promoter = TRUE)
```

sesameData_getProbesByRegion

Get probes by genomic region

Description

The function takes a genomic coordinate and output the a vector of probes on the specified platform that falls in the given genomic region.

Usage

```
sesameData_getProbesByRegion(
  regs,
  chrm = NULL,
  beg = 1,
  end = -1,
  platform = NULL,
  chrm_to_exclude = NULL,
  genome = NULL
)
```

Arguments

regs	GRanges
chrm	chromosome, when given regs are ignored
beg	begin, 1 if omitted
end	end, chromosome end if omitted
platform	EPICv2, EPIC, HM450, ...
chrm_to_exclude	chromosome to exclude.
genome	hg38, mm10, ... will infer if not given. For additional mapping, download the GRanges object from http://zwdzwd.github.io/InfiniumAnnotation and provide the following argument ..., genome = sesameAnno_buildManifestGRanges("downloaded_file"),... to this function.

Value

GRanges of selected probes

Examples

```
## download needed data
sesameDataCache(c("Mammal40.address", "genomeInfo.hg38"))

## get probes in a region
library(GenomicRanges)
probes = sesameData_getProbesByRegion(
  GRanges('chr5', IRanges(135313937, 135419936)), platform = 'Mammal40')

## get all probes on chromosome 5
probes = sesameData_getProbesByRegion(chrm = "chr5", platform = "Mammal40")

## get all probes on chromosome X
probes = sesameData_getProbesByRegion(chrm = 'chrX', platform = "Mammal40")

## get all probes on both chromosome X and Y
probes = sesameData_getProbesByRegion(
  chrm = c('chrX', 'chrY'), platform = "Mammal40")
```

```
## get all autosomal probes
probes = sesameData_getProbesByRegion(
  chrm_to_exclude = c("chrX", "chrY"), platform = "Mammal40")
```

sesameData_getTxnGRanges

convert GRangesList to transcript GRanges

Description

convert GRangesList to transcript GRanges

Usage

```
sesameData_getTxnGRanges(genome = NULL, gr1 = NULL, merge2gene = FALSE)
```

Arguments

genome	hg38, mm10, ...
gr1	GRangesList object
merge2gene	merge transcript to genes

Value

a GRanges object

Examples

```
## all mm10 transcripts
txns <- sesameData_getTxnGRanges("mm10")

## verified protein-coding transcripts
txns[(txns$transcript_type == "protein_coding" & txns$level <= 2)]

## merged to genes
sesameData_getTxnGRanges("mm10", merge2gene = TRUE)
```

```
sesameData_txnToGeneGRanges
      convert transcript GRanges to gene GRanges
```

Description

convert transcript GRanges to gene GRanges

Usage

```
sesameData_txnToGeneGRanges(txns)
```

Arguments

txns GRanges object

Value

a GRanges object

Examples

```
txns <- sesameData_getTxnGRanges("mm10")
genes <- sesameData_txnToGeneGRanges(txns)
```

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