

Package ‘haploR’

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Type Package

Title Query 'HaploReg', 'RegulomeDB'

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Description A set of utilities for querying

'HaploReg' <<https://pubs.broadinstitute.org/mammals/haploreg/haploreg.php>>,

'RegulomeDB' <<https://www.regulomedb.org/regulome-search/>>

web-based tools. The package connects to

'HaploReg', 'RegulomeDB' searches and downloads results, without opening web pages, directly from R environment.

Results are stored in a data frame that can be directly used in various kinds of downstream analyses.

RoxygenNote 7.1.1

Suggests knitr, rmarkdown, openxlsx

VignetteBuilder knitr

Imports httr, XML, tibble, RUnit, plyr, DT, RCurl, RJSONIO, methods

Depends R (>= 3.4.0)

Encoding UTF-8

License GPL-3

NeedsCompilation no

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as.num	<i>Converts vector of strings to numeric vector</i>
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Description

Converts vector of strings to numeric vector

Usage

```
as.num(x, na.strings = "NA")
```

Arguments

x	Input vector of strings.
na.strings	A string which represents NA. Default: "NA"

Value

A numeric vector

Examples

```
library(haploR)
as.num(c("1", "2", "X"), na.strings="X")
```

getExtendedView	<i>This function queries HaploReg web-based tool in order to Extended view for SNP of interest</i>
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Description

This function queries HaploReg web-based tool in order to Extended view for SNP of interest

Usage

```
getExtendedView(snp, url = Haploreg.settings[["extended.view.url"]])
```

Arguments

snp A SNP of interest.
url A url to HaploReg. Default: <https://pubs.broadinstitute.org/mammals/haploreg/detail_v4.1.php?query=d

Value

A list of tables t1, t2, ..., etc depending on information contained in HaploReg database.

Examples

```
tables <- getExtendedView(snp="rs10048158")  
tables
```

getStudyList *This function queries HaploReg web-based tool in order to see a list of GWAS.*

Description

This function queries HaploReg web-based tool in order to see a list of GWAS.

Usage

```
getStudyList(url = Haploreg.settings[["study.url"]])
```

Arguments

url A url to HaploReg. Default: <https://pubs.broadinstitute.org/mammals/haploreg/haploreg.php>

Value

A list of studies. Each study is itself a list of two: name, id.

Examples

```
studies <- getStudyList()  
studies
```

 queryHaploreg

This function queries HaploReg web-based tool and returns results.

Description

This function queries HaploReg web-based tool and returns results.

Usage

```
queryHaploreg(
  query = NULL,
  file = NULL,
  study = NULL,
  ldThresh = 0.8,
  ldPop = "EUR",
  epi = "vanilla",
  cons = "siphy",
  genotypes = "gencode",
  url = Haploreg.settings[["base.url"]],
  timeout = 100,
  encoding = "UTF-8",
  querySNP = FALSE,
  fields = NULL,
  verbose = FALSE
)
```

Arguments

query	Query (a vector of rsIDs).
file	A text file (one refSNP ID per line).
study	A particular study. See function <code>getHaploRegStudyList(...)</code> . Default: NULL.
ldThresh	LD threshold, r2 (select NA to only show query variants). Default: 0.8.
ldPop	1000G Phase 1 population for LD calculation. Can be: "AFR", "AMR", "ASN". Default: "EUR".
epi	Source for epigenomes. Possible values: vanilla for ChromHMM (Core 15-state model); imputed for ChromHMM (25-state model using 12 imputed marks); methyl for H3K4me1/H3K4me3 peaks; acetyl for H3K27ac/H3K9ac peaks. Default: vanilla.
cons	Mammalian conservation algorithm. Possible values: gerp for GERP, siphy for SiPhy-omega, both for both. Default: siphy.
genotypes	Show position relative to. Possible values: gencode for Gencode genes; refseq for RefSeq genes; both for both. Default: gencode.
url	HaploReg url address. Default: < https://pubs.broadinstitute.org/mammals/haploreg/haploreg.php >
timeout	A timeout parameter for curl. Default: 100

encoding	sets the encoding for correct retrieval web-page content. Default: UTF-8
querySNP	A flag indicating to return query SNPs only. Default: FALSE
fields	A set of fields to extract. Refer to the package vignette for available fields. Default: All.
verbose	Verbosing output. Default: FALSE.

Value

A data frame (table) with results similar to HaploReg uses.

Examples

```
library(haploR)
data <- queryHaploreg(c("rs10048158", "rs4791078"))
head(data)
```

queryRegulome	<i>This function queries RegulomeDB web-based tool and returns results in a data frame.</i>
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Description

This function queries RegulomeDB web-based tool and returns results in a data frame.

Usage

```
queryRegulome(
  query = NULL,
  genomeAssembly = "GRCh37",
  limit = 1000,
  timeout = 100
)
```

Arguments

query	Query (a vector of rsIDs or exact one query region in rsid or like "chr1:39492461-39492462").
genomeAssembly	Genome assembly built: can be GRCh37 (default) or GRCh38.
limit	It controls how many variants will be queried and returned for a large region. It can be a number (1000 by default) or "all". Please note that large number or "all" may get yourself hurt because you could get timeout or may even crash the server.
timeout	A timeout parameter for <code>httr::GET</code> . Default: 100

Value

a data frame (table) OR a list with the following items: - query_coordinates - features - regulome_score - variants - nearby_snps, - assembly

Examples

```
data <- queryRegulome(c("rs4791078", "rs10048158"))
head(data)
```

regulomeSearch	<i>This function queries RegulomeDB web-based tool and returns results in a data frame.</i>
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Description

This function queries RegulomeDB web-based tool and returns results in a data frame.

This function queries RegulomeDB web-based tool and returns results in a data frame.

Usage

```
regulomeSearch(
  query = NULL,
  genomeAssembly = NULL,
  limit = 1000,
  timeout = 100
)

regulomeSummary(
  query = NULL,
  limit = 1000,
  genomeAssembly = NULL,
  timeout = 100
)
```

Arguments

query	Query (a vector of rsIDs).
genomeAssembly	Genome assembly built: can be GRCh37 or GRCh38
limit	It controls how many variants will be queried and returned for a large region. It can be a number (1000 by default) or "all". Please note that large number or "all" may get yourself hurt because you could get timeout or may even crash the server.
timeout	A timeout parameter for httr::GET. Default: 100

Value

a list with the following items: - `query_coordinates` - `features` - `regulome_score` - `variants` - `nearby_snps`,
- `assembly`

a data frame (table)

Examples

```
data <- regulomeSearch("rs4791078")  
head(data)  
data <- regulomeSummary(c("rs4791078", "rs10048158"))  
head(data)
```

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