# Package 'haploR'

October 9, 2023

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Type Package						
Title Query 'HaploReg', 'RegulomeDB'						
Version 4.0.7						
<b>Date</b> 2023-10-08						
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Description A set of utilities for querying  'HaploReg' <a href="https://pubs.broadinstitute.org/mammals/haploreg/haploreg.php">https://pubs.broadinstitute.org/mammals/haploreg/haploreg.php</a> , 'RegulomeDB' <a href="https://www.regulomedb.org/regulome-search/">https://www.regulomedb.org/regulome-search/</a> > 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						
<b>Depends</b> R (>= $3.4.0$ )						
Encoding UTF-8						
License GPL-3						
NeedsCompilation no						
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Repository CRAN						
<b>Date/Publication</b> 2023-10-09 13:50:02 UTC						
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as.num

Converts vector of strings to numeric vector

## 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

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

## 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"]])
```

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#### **Arguments**

snp A SNP of interest.

url A url to HaploReg. Default: <a href="https://pubs.broadinstitute.org/mammals/haploreg/detail\_v4.1.php?query=a">https://pubs.broadinstitute.org/mammals/haploreg/detail\_v4.1.php?query=a</a>

#### Value

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

## **Examples**

```
tables <- getExtendedView(snp="rs10048158")
tables</pre>
```

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: <a href="https://pubs.broadinstitute.org/mammals/haploreg/haploreg.php">https://pubs.broadinstitute.org/mammals/haploreg/haploreg.php</a>

#### Value

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

#### **Examples**

```
studies <- getStudyList()
studies</pre>
```

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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,
  1dThresh = 0.8,
  1dPop = "EUR",
  epi = "vanilla",
  cons = "siphy",
  genetypes = "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 getHaploRegStudyList(). 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.
genetypes	Show position relative to. Possible values: gencode for Gencode genes; refseq for RefSeq genes; both for both. Default: gencode.
url	HaploReg url address. Default: <a href="https://pubs.broadinstitute.org/mammals/haploreg/haploreg.php">https://pubs.broadinstitute.org/mammals/haploreg/haploreg.php</a>
timeout	A timeout parameter for curl. Default: 100

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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: A11.
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)</pre>
```

queryRegulome

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

## 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 httr::GET. Default: 100

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#### Value

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

#### **Examples**

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

regulomeSearch

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

#### **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

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## Value

```
a list with the following items: - guery_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)</pre>
```

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