Package ‘rsnps’

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**Title**  Get ‘SNP’ (‘Single-Nucleotide’ ‘Polymorphism’) Data on the Web

**Description**  A programmatic interface to various ‘SNP’ ‘datasets’
on the web: ‘OpenSNP’ (<https://opensnp.org>), and ‘NBCIs’ ‘dbSNP’ database
are included for searching for ‘NCBI’. For ‘OpenSNP’, functions are included
for getting ‘SNPs’, and data for ‘genotypes’, ‘phenotypes’, annotations,
and bulk downloads of data by user.

**Version**  0.4.0

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**URL** https://docs.ropensci.org/rsnps/,
        https://github.com/ropensci/rsnps/

**BugReports** https://github.com/ropensci/rsnps/issues/

**VignetteBuilder** knitr

**Encoding** UTF-8

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rsnps-package  Get SNP (Single-Nucleotide Polymorphism) Data on the Web

Description

This package gives you access to data from OpenSNP (https://opensnp.org) via their API (https://opensnp.org/faq#api) and NCBI’s dbSNP SNP database (https://www.ncbi.nlm.nih.gov/snp).

NCBI Authentication

This applies the function `ncbi_snp_query()`:

You can optionally use an API key, if you do it will allow higher rate limits (more requests per time period)

If you don’t have an NCBI API key, get one at https://www.ncbi.nlm.nih.gov/account/

Create your key from your account. After generating your key set an environment variable as `ENTREZ_KEY` in `.Renviron.`

`ENTREZ_KEY='youractualkeynotthisstring'`

You can optionally pass in your API key to the key parameter in NCBI functions in this package. However, it’s much better from a security perspective to set an environment variable.
allgensnp

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allgensnp

Get openSNP genotype data for all users at a particular snp.

Description

Get openSNP genotype data for all users at a particular snp.

Usage

allgensnp(snp = NA, ...)

Arguments

snp (character) A SNP name

... Curl options passed on to crul::HttpClient

Value

data.frame of genotypes for all users at a certain SNP

See Also

Other opensnp-fxns: allphenotypes(), annotations(), download_users(), fetch_genotypes(), genotypes(), phenotypes_byid(), phenotypes(), users()

Examples

## Not run:
x <- allgensnp(snp = 'rs7412')
head(x)

## End(Not run)
allphenotypes

Get all openSNP phenotypes, their variations, and how many users have data available for a given phenotype.

Description

Either return data.frame with all results, or output a list, then call the characteristic by id (parameter = "id") or name (parameter = "characteristic").

Usage

allphenotypes(df = FALSE, ...)

Arguments

df

Return a data.frame of all data. The column known_variations can take multiple values, so the other columns id, characteristic, and number_of_users are replicated in the data.frame. Default: FALSE

...

Curl options passed on to crul::HttpClient

Value

data.frame of results, or list if df=FALSE

See Also

Other opensnp-fxns: allgensnp(), annotations(), download_users(), fetch_genotypes(), genotypes(), phenotypes_byid(), phenotypes(), users()

Examples

## Not run:
# Get all data
allphenotypes(df = TRUE)

# Output a list, then call the characterisitc of interest by 'id' or
# 'characteristic'
datalist <- allphenotypes()
names(datalist) # get list of all characteristics you can call
datalist[["ADHD"]]["ADHD"] # get data.frame for 'ADHD'
datalist[[c("mouth size","SAT Writing")]["ADHD"] # get data.frame for 'ADHD'

## End(Not run)
Annotations

Get all openSNP phenotypes, their variations, and how many users have data available for a given phenotype.

Description
Either return data.frame with all results, or output a list, then call the characteristic by id (parameter = "id") or name (parameter = "characteristic").

Usage
annotations(
  snp = NA,
  output = c("all", "plos", "mendeley", "snpedia", "metadata"),
  ...
)

Arguments

  snp      SNP name.
  output   Name the source or sources you want annotations from (options are: 'plos', 'mendeley', 'snpedia', 'metadata'). 'metadata' gives the metadata for the response.
  ...     Curl options passed on to curl::HttpClient

Value
data.frame of results

See Also
Other opensnp-fxns: allgensnp(), allphenotypes(), download_users(), fetch_genotypes(), genotypes(), phenotypes_byid(), phenotypes(), users()

Examples
## Not run:
# Get all data
# get just the metadata
annotations(snp = 'rs7903146', output = 'metadata')

# just from plos
annotations(snp = 'rs7903146', output = 'plos')

# just from snpedia
annotations(snp = 'rs7903146', output = 'snpedia')

# get all annotations
download_users

Description

Download openSNP user files.

Usage

download_users(name = NULL, id = NULL, dir = "~/", ...)  

Arguments

name User name
id User id
dir Directory to save file to
... Curl options passed on to curl::HttpClient

Value

File downloaded to directory you specify (or default), nothing returned in R.

See Also

Other opensnp-fxns: allgensnp(), allphenotypes(), annotations(), fetch_genotypes(), genotypes(), phenotypes_byid(), phenotypes(), users()

Examples

## Not run:
# Download a single user file, by id
download_users(id = 14)

# Download a single user file, by user name
download_users(name = 'kevinmcc')

# Download many user files
lapply(c(14,22), function(x) download_users(id=x))
read_users(id=14, nrows=5)

## End(Not run)
fetch_genotypes

Description

Download openSNP genotype data for a user

Usage

fetch_genotypes(url, rows = 100, filepath = NULL, quiet = TRUE, ...)

Arguments

url (character) URL for the download. See example below of function use.
rows (integer) Number of rows to read in. Useful for getting a glimpse of the data. Negative and other invalid values are ignored, giving back all data. Default: 100
filepath (character) If none is given the file is saved to a temporary file, which will be lost after your session is closed. Save to a file if you want to access it later.
quiet (logical) Should download progress be suppressed. Default: TRUE
...
Further args passed on to download.file()

Details

Beware, not setting the rows parameter means that you download the entire file, which can be large (e.g., 15MB), and so take a while to download depending on your connection speed. Therefore, rows is set to 10 by default to sort of protect the user.

Internally, we use download.file() to download each file, then read.table() to read the file to a data.frame.

Value

data.frame for a single user, with four columns:

- rsid (character)
- chromosome (integer)
- position (integer)
- genotype (character)

See Also

Other opensnp-fxns: allgensnp(), allphenotypes(), annotations(), download_users(), genotypes(), phenotypes_byid(), phenotypes(), users()
Examples

```r
## Not run:
# get a data.frame of the users data
data <- users(df = TRUE)
head( data[[1]] ) # users with links to genome data
mydata <- fetch_genotypes(url = data[[1]][1,"genotypes.download_url"],
                          file="~/myfile.txt")

# see some data right away
mydata

# Or read in data later separately
read.table("~/myfile.txt", nrows=10)

## End(Not run)
```

---

**genotypes**

*Get openSNP genotype data for one or multiple users.*

**Description**

Get openSNP genotype data for one or multiple users.

**Usage**

```r
genotypes(snp = NA, userid = NA, df = FALSE, ...)
```

**Arguments**

- `snp` SNP name.
- `userid` ID of openSNP user.
- `df` Return data.frame (TRUE) or not (FALSE). Default: FALSE
- `...` Curl options passed on to `cru::HttpClient`

**Value**

List (or data.frame) of genotypes for specified user(s) at a certain SNP.

**See Also**

Other opensnp-fxns: `allgensnp()`, `allphenotypes()`, `annotations()`, `download_users()`, `fetch_genotypes()`, `phenotypes_byid()`, `phenotypes()`, `users()`
get_frequency

Examples

```r
## Not run:
genotypes(snp='rs9939609', userid=1)
genotypes('rs9939609', userid='1,6,8', df=TRUE)
genotypes('rs9939609', userid='1-2', df=FALSE)
## End(Not run)
```

get_frequency  Internal function to get the frequency of the variants from different studies.

Description

Internal function to get the frequency of the variants from different studies.

Usage

```r
get_frequency(Class, primary_info, study = "GnomAD")
```

Arguments

- **Class**  What kind of variant is the rsid. Accepted options are "snv", "snp" and "delins".
- **primary_info**  refsnp entry read in JSON format
- **study**  Study from which frequency information is obtained. Possibilities include: GnomAD (default), 1000Genomes, ALSPAC, Estonian, NorthernSweden, TWIN-SUK

get_gene_names  Internal function to get gene names.

Description

If multiple gene names are encountered they are collapsed with a "/".

Usage

```r
gene_names(primary_info)
```

Arguments

- **primary_info**  refsnp entry read in JSON format
ncbi_snp_query

cbi_snp_query

get_placements

Internal function to get the position, alleles, assembly, hgvs notation

Description

Internal function to get the position, alleles, assembly, hgvs notation

Usage

get_placements(primary_info)

Arguments

primary_info refsnp entry read in JSON format

ncbi_snp_query

Query NCBI’s refSNP for information on a set of SNPs via the API

Description

This function queries NCBI’s refSNP for information related to the latest dbSNP build and latest reference genome for information on the vector of snps submitted.

Usage

ncbi_snp_query(snps)

Arguments

snps (character) A vector of SNPs (rs numbers).

Details

This function currently pulling data for Assembly 38 - in particular note that if you think the BP position is wrong, that you may be hoping for the BP position for a different Assembly.

Note that you are limited in the to a max of one query per second and concurrent queries are not allowed. If users want to set curl options when querying for the SNPs they can do so by using http::set_config/http::with_config
Value

A dataframe with columns:

- `query`: The rs ID that was queried.
- `chromosome`: The chromosome that the marker lies on.
- `bp`: The chromosomal position, in base pairs, of the marker, as aligned with the current genome used by dbSNP. We add 1 to the base pair position in the BP column in the output data.frame to agree with what the dbSNP website has.
- `rsid`: Reference SNP cluster ID. If the rs ID queried has been merged, the up-to-date name of the ID is returned here, and a warning is issued.
- `gene`: If the rsid lies within a gene (either within the exon or introns of a gene), the name of that gene is returned here; otherwise, NA. Note that the gene may not be returned if the rsid lies too far upstream or downstream of the particular gene of interest.
- `alleles`: The alleles associated with the SNP if it is a SNV; otherwise, if it is an INDEL, microsatellite, or other kind of polymorphism the relevant information will be available here.
- `minor`: The allele for which the MAF is computed, given it is an SNV; otherwise, NA.
- `maf`: The minor allele frequency of the SNP, given it is an SNV. This is drawn from the current global reference population used by NCBI (GnomAD).
- `ancestral_allele`: allele as described in the current assembly
- `variation_allele`: difference to the current assembly
- `seqname`: Chromosome RefSeq reference.
- `hgvs`: full hgvs notation for variant
- `assembly`: which assembly was used for the annotations
- `ref_seq`: sequence in reference assembly

References


Examples

```r
## Not run:
## an example with both merged SNPs, non-SNV SNPs, regular SNPs,
## SNPs not found, microsatellite
SNPs <- c("rs332", "rs420358", "rs1837253", "rs1209415715", "rs111068718")
ncbi_snp_query(SNPs)
# ncbi_snp_query("123456") ##invalid: must prefix with 'rs'
ncbi_snp_query("rs420358")
ncbi_snp_query("rs332") # warning that its merged into another, try that
ncbi_snp_query("rs121909001")
ncbi_snp_query("rs1837253")
ncbi_snp_query("rs1209415715")
```
phenotypes

Get openSNP phenotype data for one or multiple users.

Description

Get openSNP phenotype data for one or multiple users.

Usage

phenotypes(userid = NA, df = FALSE, ...)

Arguments

userid ID of openSNP user.
df Return data.frame (TRUE) or not (FALSE). Default: FALSE
...

Curl options passed on to crul::HttpClient

Value

List of phenotypes for specified user(s).

See Also

Other opensnp-fxns: allgensnp(), allphenotypes(), annotations(), download_users(), fetch_genotypes(), genotypes(), phenotypes_byid(), users()

Examples

## Not run:
phenotypes(userid =1)
phenotypes(userid = '1,6,8', df=TRUE)
phenotypes(userid = '1-8', df=TRUE)

# coerce to data.frame
library(plyr)
df <- ldply(phenotypes(userid = '1-8', df=TRUE))
head(df); tail(df)

# pass on curl options
phenotypes(1, verbose = TRUE)

## End(Not run)
### phenotypes_byid

Get all openSNP known variations and all users sharing that phenotype for one phenotype(-ID).

#### Description

Get all openSNP known variations and all users sharing that phenotype for one phenotype(-ID).

#### Usage

```r
phenotypes_byid(
  phenotypeid = NA,
  return_ = c("description", "knownvars", "users"),
  ...
)
```

#### Arguments

- **phenotypeid**
  ID of openSNP phenotype.
- **return_**
  Return data.frame (TRUE) or not (FALSE). Default: FALSE
- **...**
  Curl options passed on to crul::HttpClient

#### Value

List of description of phenotype, list of known variants, or data.frame of variants for each user with that phenotype.

#### See Also

Other opensnp-fxs: `allgensnp()`, `allphenotypes()`, `annotations()`, `download_users()`, `fetch_genotypes()`, `genotypes()`, `phenotypes()`, `users()`

#### Examples

```r
## Not run:
phenotypes_byid(phenotypeid=12, return_ = 'desc')
phenotypes_byid(phenotypeid=12, return_ = 'knownvars')
phenotypes_byid(phenotypeid=12, return_ = 'users')

# pass on curl options
phenotypes_byid(phenotypeid=12, return_ = 'desc', verbose = TRUE)

## End(Not run)
```
**read_users**  
*Read in openSNP user files from local storage.*

**Description**

Beware, these tables can be large. Check your RAM before executing. Or possibly read in a subset of the data. This function reads in the whole kitten kaboodle.

**Usage**

```r
read_users(name = NULL, id = NULL, path = NULL, ...)
```

**Arguments**

- `name` User name
- `id` User id
- `path` Path to file to read from.
- `...` Parameters passed on to `read.table()`

**Details**

If you specify a name or id, this function reads environment variables written in the function `download_users`, and then searches against those variables for the path to the file saved. Alternatively, you can supply the path.

**Value**

A data.frame.

**Examples**

```r
## Not run:
# dat <- read_users(name = "kevinmcc")
# head(dat)
# dat <- read_users(id = 285)

## End(Not run)
```
rsnps-defunct

---

**rsnps-defunct**

**Defunct functions in rsnps**

### Description

- **LDSearch()**: Function name changed to ld_search
- **ld_search()**: The Broad Institute took the service down, see https://www.broadinstitute.org/snap/snap
- **NCBI_snp_query()**: Function name changed to ncbi_snp_query
- **NCBI_snp_query2()**: Function name changed to ncbi_snp_query
- **ncbi_snp_summary()**: Function name changed to ncbi_snp_query
- **ncbi_snp_query2()**: Function name changed to ncbi_snp_query

---

**users**

**Get openSNP users.**

### Description

Get openSNP users.

### Usage

```r
users(df = FALSE, ...)
```

### Arguments

- **df**
  - Return data.frame (TRUE) or not (FALSE). Default: FALSE
- **...**
  - Curl options passed on to `crl::HttpClient`

### Value

List of openSNP users, their ID numbers, and XX if available.

### See Also

Other opensnp-fxns: `allgensnp()`, `allphenotypes()`, `annotations()`, `download_users()`, `fetch_genotypes()`, `genotypes()`, `phenotypes_byid()`, `phenotypes()`
Examples

## Not run:
# just the list
data <- users(df = FALSE)
data

# get a data.frame of the users data
data <- users(df = TRUE)
data[[1]] # users with links to genome data
data[[2]] # users without links to genome data

## End(Not run)
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