Package ‘Mega2R’

December 21, 2023

Version 1.1.0

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Title Accessing and Processing a 'Mega2' Genetic Database

Description Uses as input genetic data that have been reformatted and
stored in a 'SQLite' database; this database is initially created
by the standalone 'mega2' C++ program (available freely from
<https://watson.hgen.pitt.edu/register/>). Loads and manipulates
data frames containing genotype, phenotype, and family
information from the input 'SQLite' database, and decompresses
needed subsets of the genotype data, on the fly, in a memory
efficient manner. We have also created several more functions
that illustrate how to use the data frames as well as perform
useful tasks: these permit one to run the 'pedgene' package to
carry out gene-based association tests on family data using
selected marker subsets, to run the 'SKAT' package to carry out
gene-based association tests using selected marker subsets, to
run the 'famSKATRC' package to carry out gene-based association
tests on families (optionally) and with rare or common variants
using selected marker subsets, to output the 'Mega2R' data as a
VCF file and related files (for phenotype and family data), and
to convert the data frames into CoreArray Genomic Data Structure
(GDS) format.

URL https://watson.hgen.pitt.edu/mega2/mega2r/

BugReports https://groups.google.com/forum/#!forum/mega2-users

Depends R (>= 3.5.0), SKAT, pedgene, gdsfmt

License GPL-2

LinkingTo Rcpp

biocViews Genetics

Imports AnnotationDbi, DBI, GenomeInfoDb, RSQLite, methods, famSKATRC,
kinship2

Suggests knitr, rmarkdown, formatR, TxDb.Hsapiens.UCSC.hg19.knownGene,
org.Hs.eg.db
R topics documented:

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applyFnToGenes

applyFnToGenes

Description

This function generates base pair ranges from its input arguments. Each range specifies a chromosome, a start base pair and end base pair. Typically, a range could be a gene transcript, though it could be a whole chromosome, or a run of base pairs on a chromosome. Once the ranges are generated, applyFnToRanges is called to find all the rows (i.e. markers) from the markers data frame that fall in each range. For these markers, a matrix of the genotypes is generated. Finally, the op function is called for each range with the arguments: markers, range, and ‘environment’.

Usage

```r
applyFnToGenes(op = function (markers, range, envir) {},
genes_arg = NULL,
ranges_arg = matrix(ncol = 3, nrow = 0),
chrs_arg = vector("integer", 0),
markers_arg = vector("character", 0),
type_arg = "TX",
fuzz_arg = 0,
envir = ENV)
```

Arguments

- **op**  
  Is a function of three arguments. It will be called repeatedly by applyFnToGenes in a try/catch context. The arguments are:
  - **markers**  
    Marker data for each marker selected. A marker is a data frame with the following 5 observations:
    - **locus_link**  
      is the ordinal ranking of this marker among all loci
    - **locus_link_fill**  
      is the position of corresponding marker genotype data in the unified_genotype_table
    - **MarkerName**  
      is the text name of the marker
    - **chromosome**  
      is the integer chromosome number
    - **position**  
      is the integer base pair position of marker
  - **range**  
    An indicator of which range argument these markers correspond to.
  - **envir**  
    An ‘environment’ holding Mega2R data frames and state data.
applyFnToGenes

genes_arg    a character vector of gene names. All the transcripts identified with the specified
gene in BioConductor Annotation,
TxDb.Hsapiens.UCSC.hg19.knownGene, are selected. This produces multiple "range" elements containing chromosome, start base pair, end base pair. (If
the gene name is ".", all the transcript will be selected.) Note: BioConductor
Annotation org.Hs.eg.db is used to convert from gene name to ENTREZ gene id.

ranges_arg   an integer matrix of three columns. The columns define a range: a chromosome
number, a start base pair value, and an end base pair value.

chrs_arg     an integer vector of chromosome numbers. All of the base pairs on each chro-
mosomes will be selected as a single range.

markers_arg  a data frame with the following 5 observations:
locus_link   is the ordinal ranking of this marker among all loci
locus_link_fill  is the position of corresponding marker genotype data in the
unified_genotype_table
MarkerName  is the text name of the marker
chromosome  is the integer chromosome number
position    is the integer base pair position of marker
type_arg    a character vector of length 1 that contains "TX" or does not. If it is "TX",
which is the default, the TX fields of BioConductor Annotation,
TxDb.Hsapiens.UCSC.hg19.knownGene are used to define the base pair ranges
and chromosome. Otherwise, the CDS fields are used.
fuzz_arg    is an integer vector of length one or two. The first argument is used to reduce the
start base pair selected from each transcript and the second to increase the end
base pair position. (If only one value is present, it is used for both adjustments.)
Note: The values can be positive or negative.

envir       an 'environment' that contains all the data frames created from the SQLite
database.

Value
None

Note
If you want subsequent calls to op to share information, data can be placed in a data frame that is
added to the 'environment'.

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)

show = function(m, r, e) {
  print(r)
  print(m)
  print(head(getgenotypes(m, envir = e)))}
applyFnToMarkers

apply function "show" to all transcripts on genes ELL2 and CARD15

# donttestcheck: time
applyFnToGenes(show, genes_arg = c("CEP104"))

# apply function "show" to all genotypes on chromosomes 11 for two base pairs ranges
applyFnToGenes(show, ranges_arg = matrix(c(1, 5000000, 10000000, 1, 10000000, 15000000), ncol = 3, nrow = 2, byrow = TRUE))

# apply function "show" to all genotypes for first marker in each chromosome
applyFnToGenes(show, markers_arg = ENV$markers[!duplicated(ENV$markers$chromosome), 3])

# apply function "show" to all genotypes on chromosomes 24 and 26
applyFnToGenes(show, chrs_arg = c(24, 26))

applyFnToMarkers

apply a function to the genotypes from a set of markers

Description

A matrix of the genotypes for all the specified markers is generated. Then, the call back function, op, is called with the markers, NULL (for the range), and the 'environment'.

Usage

applyFnToMarkers(op = function (markers, range, envir) {},
                  markers_arg,
                  envir = ENV)

Arguments

op Is a function of three arguments. It will be called once by applyFnToMarkers in a try/catch context. The arguments are:

markers Marker data for each marker in geno. A marker is a data frame with the following 5 observations:
    locus_link is the ordinal ranking of this marker among all loci
    locus_link_fill is the position of corresponding marker genotype data in the unified_genotype_table
    MarkerName is the text name of the marker
    chromosome is the integer chromosome number
    position is the integer base pair position of marker
range NULL: to indicate no explicit range was specified.
applyFnToRanges

**apply a function to all the genotypes for markers found in several specified ranges**

**Description**

First, for each range, determine the markers that fall between the start and end base pair of the range. Then, for each set of markers generate a matrix of the genotypes of those markers. Finally, the op function is called for each range with the arguments: markers, range, and 'environment'.

**Usage**

```r
applyFnToRanges(op = function (markers, range, envir) {},
                 ranges_arg = NULL,
                 indices_arg = NULL,
                 fuzz_arg = 0,
                 envir = ENV)
```

**Arguments**

- `markers_arg`: a data frame with the following 5 observations:
  - `locus_link`: is the ordinal ranking of this marker among all loci
  - `locus_link_fill`: is the position of corresponding marker genotype data in the `unified_genotype_table`
  - `MarkerName`: is the text name of the marker
  - `chromosome`: is the integer chromosome number
  - `position`: is the integer base pair position of marker

- `envir`: an 'environment' that contains all the data frames created from the SQLite database.

**Value**

None

**Examples**

```r
db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)
show = function(m, r, e) {
  print(r)
  print(m)
  print(head(getgenotypes(m, envir = e)))
}

# apply function "show" to all genotypes > 5,000,000 bp
applyFnToMarkers(show, ENV$markers[ENV$markers$position > 5000000,])
```
applyFnToRanges

Arguments

- **op**
  - Is a function of three arguments. It will be called repeatedly by `applyFnToRanges` in a try/catch context. The arguments are:
    - **markers** Marker data for each marker in `geno`. A marker is a data frame with the following 5 observations:
      - **locus_link** is the ordinal ranking of this marker among all loci
      - **locus_link_fill** is the position of corresponding marker genotype data in the `unified_genotype_table`
      - **MarkerName** is the text name of the marker
      - **chromosome** is the integer chromosome number
      - **position** is the integer base pair position of marker
    - **range** An indicator of which range argument of `applyFnToRanges` these markers correspond to.
    - **envir** An 'environment' holding Mega2R data frames and state data.

- **ranges_arg**
  - is a data frame that contains at least 4 observations: a name, a chromosome, a start base pair position and an end base pair position.

- **indices_arg**
  - is a vector of 3 integers that specify the location of chromosome, start base pair column and end base pair column of the `ranges_arg` data frame. An optional fourth integer indicates the column containing the name of the ranges.

- **fuzz_arg**
  - is an integer vector of length one or two. The first argument is used to reduce the start base pair selected from each range and the second to increase the end base pair position. (If only one value is present, it is used for both changes.) Note: The values can be positive or negative.

- **envir**
  - an 'environment' that contains all the data frames created from the SQLite database.

Value

None

Note

If the `ranges_arg` and `indices_arg` are NULL or missing, then the default ranges that have been set by `setRanges` are used. If `setRanges` has not been called, a default set of the ranges is used.

Examples

```r
db = system.file("exdata", "seqsim.db", package="Mega2R")
ENV = read.Mega2DB(db)

show = function(m, r, e) {
  print(r)
  print(m)
  print(head(getgenotypesraw(m, envir = e)))
}

# apply function "show" to all genotypes on chromosomes 1 for two base pair
```

# ranges
applyFnToRanges(show,
    ranges_arg =
    matrix(c(1, 2244000, 2245000,
        1, 3762500, 3765000),
        ncol = 3, nrow = 2, byrow = TRUE),
    indices_arg = 1:3)

# apply function "show" to all genotypes on chromosomes 1 for two base pair
# ranges
applyFnToRanges(show,
    ranges_arg =
    matrix(c(1, 2240000, 2245000, "range1",
        1, 3760000, 3765000, "range2"),
        ncol = 4, nrow = 2, byrow = TRUE),
    indices_arg = 1:4)

clean_mega2rtutorial_data

remove tutorial data

Description
This function removes the Mega2R tutorial (inst/exdata) data that was copied to the specified directory.

Usage
clean_mega2rtutorial_data(dir = file.path(tempdir(), "Mega2Rtutorial"))

Arguments

   dir               The directory to remove the tutorial data to. By default, this is tempdir()/Mega2Rtutorial

Value
None

Examples
clean_mega2rtutorial_data()
computeDosage

computeDosage function

Description

Convert the genotypesraw() allele patterns of 0x10001, 0x10002 (or 0x20001), 0x20002, 0 to the numbers 0, 1, 2, 9 for each marker. (Reverse, the order iff allele "1" has the minor allele frequency.)

Usage

computeDosage(markers_arg, range_arg, envir)

Arguments

- markers_arg a data.frame with the following 5 observations:
  - locus_link is the ordinal ranking of this marker among all loci
  - locus_link_fill is the position of corresponding genotype data in the unified_genotype_table
  - MarkerName is the text name of the marker
  - chromosome is the integer chromosome number
  - position is the integer base pair position of marker

- range_arg one row of a ranges_arg. The latter is a data frame of at least three integer columns. The columns indicate a range: a chromosome number, a start base pair value, and an end base pair value.

- envir 'environment' containing SQLite database and other globals especially the phenotype_table, phe.

Value

a matrix of samples X markers for all the markers that have nonzero changes.

See Also

DOfamSKATRC

Examples

```r
db = system.file("exdata", "seqsim.db", package="Mega2R")
ENV = init_famSKATRC(db, verbose = TRUE)
dimDosage = function(m, r, e) {print(dim(computeDosage(m, r, e)))}
applyFnToRanges(dimDosage, ENV$refRanges[50:60, ], ENV$refIndices, envir=ENV)
# This will use return dosage matrices for the markers in the ranges 50 - 60,
# but is basically ignores the results.
```
dbmega2_import  

*read Mega2 SQLite database into R*

---

**Description**

Read the fields of SQLite database tables that are required for Mega2R into data frames. These data frames are stored in an ‘environment’ which is returned. This function also adds some state data, extra data frames, and computed data frames to the ‘environment’.

**Usage**

```r
dbmega2_import(dbname,  
    bpPosMap = NULL,  
    verbose = FALSE)
```

**Arguments**

- `dbname` file path to SQLite database.
- `bpPosMap` index that specifies which map in the map_table should be used for marker chromosome/position. If it is `NULL`, the internal variable `base_pair_position_index` is used instead. `showMapNames()` shows the association between map name and map number.
- `verbose` print out statistics on the name/size of each table read and show column headers. Also, save the verbose value for use by other Mega2R functions.

**Value**

`envir` an environment that contains all the data frames made from the SQLite database.

**Examples**

```r
db = system.file("exdata", "seqsim.db", package="Mega2R")
ENV = dbmega2_import(db, verbose = TRUE)
ENV = dbmega2_import(db)
```

---

**DOfamSKATRC**  

*DofamSKATRC call back function*

---

**Description**

Convert the genotypesraw() allele patterns of 0x10001, 0x10002 (or 0x20001), 0x20002, 0 to the numbers 0, 1, 2, 9 for each marker. (Reverse, the order iff allele "1" has the minor allele frequency.) Ignore markers that have no variants. Finally, invoke `famSKAT_RC` with the converted genotype matrix. Save information about the range and the p.value calculated by `famSKAT_RC` in `envir$famSKATRC_results`. If you want to change the argument values to this function they should be changed instead when calling the Mega2FamSKATRC function.
Usage

DOfamSKATRC(
  markers_arg,
  range_arg,
  envir,
  pheno = 3,
  id = NULL,
  covariates = NULL,
  sqrtweights_c = NULL,
  sqrtweights_r = NULL,
  binomialimpute = TRUE,
  acc = 1e-06,
  maf = 0.05,
  phi = c(0, 0.2, 0.5, 0.9)
)

Arguments

markers_arg a data.frame with the following 5 observations:
  locus_link is the ordinal ranking of this marker among all loci
  locus_link_fill is the position of corresponding genotype data in the unified_genotype_table
  MarkerName is the text name of the marker
  chromosome is the integer chromosome number
  position is the integer base pair position of marker

range_arg one row of a ranges_arg. The latter is a data frame of at least three integer columns. The columns indicate a range: a chromosome number, a start base pair value, and an end base pair value.

envir 'environment' containing SQLite database and other globals especially the phenotype_table, phe.

pheno is an index into the phenotypes_table to select the phenotype. Missing phenotypes are represented by NA.

id a vector of individuals to be included in the test, a subset of the family members. If NULL is given, all members will be used.

covariates a matrix of covariates for the phenotype.

sqrtweights_c weight function for common variants, if NULL use weight set in init_famSKAT

sqrtweights_r weight function for rare variants, if NULL use weight set in init_famSKAT.

binomialimpute if TRUE, impute missing genotypes using a binomial distribution.

acc accuracy used in Davies approximation.

maf threshold used to separate rare from common variants.

phi a vector of ratios ratios; each indicates the contribution of rare variants.

Value

None
Note

This function accumulates output in the data frame, `envir$famSKATRC_results`. It will print out the lines as they are generated if `envir$verbose` is TRUE. It does not write the data frame to a file. You must save the data frame. You also must initialize the data frame when necessary.

See Also

`init_famSKATRC`, `Mega2famSKATRC`

Examples

```r
db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = init_famSKATRC(db, verbose = TRUE)
ENV$famSKATRC_results = ENV$famSKATRC_results[0, ]
Mega2famSKATRC(gs=1:1, envir=ENV, pheno=3)
# this sets one of the many arguments for DOfamSKATRC
# but basically prepares the ENV for the direct use of DOfamSKATRC (below).

# donttestcheck: try this below instead if there is time
Mega2famSKATRC(genes=c("CEP104"), envir=ENV, pheno=3 )

# DOfamSKATRC is called within Mega2famSKATRC. init_famSKATRC and Mega2famSKATRC need to be
# called to set up the environment for famSKAT_RC to run. BUT, you should ignore DOfamSKATRC
# and use Mega2famSKATRC instead.
# applyFnToRanges(DOfamSKATRC, ENV$refRanges[50:60, ], ENV$refIndices, envir=ENV)
# this will use all the default argument values for DOfamSKATRC
```

DOpedgene

**pedgene call back function**

Description

First, ignore call backs that have less than two markers. Second, convert the genotypes raw() patterns of 0x10001, 0x10002 (or 0x20001), 0x20002, 0 from the genotype matrix to the numbers 0, 1, 2, 0 for each marker. (Reverse, the order if allele "1" has the minor allele frequency.) Next, prepend the pedigree and person columns of the family data to this modified genotype matrix. Finally, invoke pedgene with the family data and genotype matrix for several different weights. Save the kernel and burden, value and p-value for each measurement in `envir$pedgene_results`.

Usage

```r
DOpedgene(markers_arg, range_arg, envir = ENV)
```
Arguments

markers_arg   a data.frame with the following 5 observations:
   locus_link is the ordinal ranking of this marker among all loci
   locus_link_fill is the position of corresponding genotype data in the unified_genotype_table
   MarkerName is the text name of the marker
   chromosome is the integer chromosome number
   position is the integer base pair position of marker

range_arg one row of a ranges_arg. The latter is a data frame of at least three integer columns. The columns indicate a range: a chromosome number, a start base pair value, and an end base pair value.

envir 'environment' containing SQLite database and other globals

Value

None

Note

This function appends output to the data frame, envir$pedgene_results. It will print out the lines as they are generated if envir$verbose is TRUE. The data frame envir$pedgene_results is initialized by init_pedgene, and is appended to each time DOpedgene is run.

See Also

init_pedgene

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = init_pedgene(db)
ENVverbose = TRUE
applyFnToRanges(DOpedgene, ENV$refRanges[50:60,], ENV$refIndices)

# donttestcheck: try this below if there is time
applyFnToGenes(DOpedgene, genes_arg = c("CEP104"))
Description

Convert the genotypesraw() allele patterns of 0x10001, 0x10002 (or 0x20001), 0x20002, 0 to the numbers 0, 1, 2, 9 for each marker. (Reverse the order iff allele "1" has the minor allele frequency.) Ignore markers that have no variants (unless allMarkers is TRUE). Finally, invoke SKAT with the converted genotype matrix, Null model saved in envir$obj, and any additionally supplied arguments. Save information about the range and the p.value calculated by SKAT in envir$SKAT_results.

Usage

DOSKAT(markers_arg, range_arg, envir, ...)

Arguments

- **markers_arg** a data.frame with the following 5 observations:
  - `locus_link` is the ordinal ranking of this marker among all loci
  - `locus_link_fill` is the position of corresponding genotype data in the `unified_genotype_table`
  - `MarkerName` is the text name of the marker
  - `chromosome` is the integer chromosome number
  - `position` is the integer base pair position of marker

- **range_arg** one row of a ranges_arg. The latter is a data frame of at least three integer columns. The columns indicate a range: a chromosome number, a start base pair value, and an end base pair value.

- **envir** 'environment' containing SQLite database and other globals

... extra arguments for SKAT

Value

None

Note

This function accumulates output in the data frame, envir$SKAT_results. It will print out the lines as they are generated if envir$verbose is TRUE. It does not write the data frame to a file. You must save the data frame. You also must initialize the data frame when necessary.

See Also

init_SKAT, Mega2SKAT

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = init_SKAT(db, verbose = TRUE, allMarkers = FALSE)
Mega2SKAT(ENV$phe[, 3] - 1 ~ 1, "D", gs=1:1)

# donttestcheck: try this below instead if there is time
Mega2SKAT(ENV$phe[, 3] - 1 ~ 1, "D", kernel = "linear.weighted", ...
weights.beta = c(0.5, 0.5), genes=c("CEP104")

# DOSKAT is called internally to Mega2SKAT. init_SKAT and Mega2SKAT need to be
# called to set up the environment for DOSKAT to run. You should ignore DOSKAT
# and use Mega2SKAT instead
#
applyFnToRanges(DOSKAT, ENV$refRanges[50:60, ], ENV$refIndices)

dump_mega2rtutorial_data

**dump tutorial data**

**Description**

This function retrieves data stored in the Mega2rtutorial (inst/exdata). It dumps them in the specified directory.

**Usage**

dump_mega2rtutorial_data(dir = file.path(tempdir(), "Mega2Rtutorial"))

**Arguments**

- **dir**: The directory to store the tutorial data to. By default, this is tempdir()/Mega2Rtutorial

**Value**

None

**Examples**

dump_mega2rtutorial_data()

getgenotypes

**fetch genotype character matrix for specified markers**

**Description**

This function calls a C++ function that does all the heavy lifting. It passes the arguments necessary for the C++ function: some from the caller’s arguments and some from data frames that are in the "global" environment, **envir**. From the markers_arg argument, it fetches the locus_index and the index in the **unified_genotype_table**. It also passes the allele nucleotide separator argument. From the "global" environment, **envir**, it gets a bit vector of compressed genotype information, the alleles for each marker, and some bookkeeping related data. Note: This function also contains a dispatch/switch on the type of compression in the genotype vector. A different C++ function is called when there is compression versus when there is no compression.
Usage

getgenotypes(markers_arg, sepstr = "", envir = ENV)

Arguments

markers_arg  a data.frame with the following 5 observations:
  locus_link  is the ordinal ranking of this marker among all loci
  locus_link_fill  is the position of corresponding genotype data in the \textit{unified_genotype_table}
  MarkerName  is the text name of the marker
  chromosome  is the integer chromosome number
  position  is the integer base pair position of marker

sepstr  separator string inserted between the alleles (default is none). When present, this is typically a space, a tab or "/".

envir  an environment that contains all the data frames created from the SQLite database.

Details

The \textit{unified_genotype_table} contains one raw vector for each person. In the vector there are two bits for each genotype. This function creates an output matrix by fixing the marker and collecting genotype information for each person and then repeating for all the needed markers. (Currently, this appears slightly faster than a scan which fixes the person and iterates over markers.)

Value

a matrix of genotypes represented as two allele pairs. The matrix has one column for each marker in \textit{markers_arg} argument. There is one row for each person in the family \textit{(fam)} table.

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)

getgenotypes(ENV$markers)

Description

This function calls a C++ function that does all the heavy lifting. It passes the arguments necessary for the C++ function: some from the caller’s arguments and some from data frames that are in the "global" environment, \textit{envir}. From its markers_arg argument, it gets the locus_index and the index in the \textit{unified_genotype_table}. From the "global" environment, \textit{envir}, it gets a bit vector of compressed genotype information, and some bookkeeping related data. Note: This function also contains a dispatch/switch on the type of compression in the genotype vector. A different C++ function is called when there is compression versus when there is no compression.
getgenotypesdos

Usage

getgenotypesdos(markers_arg, envir = ENV)

Arguments

markers_arg a data.frame with the following 5 observations:
  locus_link is the ordinal ranking of this marker among all loci
  locus_link_fill is the position of corresponding genotype data in the unified_genotype_table
  MarkerName is the text name of the marker
  chromosome is the integer chromosome number
  position is the integer base pair position of marker

envir an environment that contains all the data frames created from the SQLite database.

Details

The unified_genotype_table contains one raw vector for each person. In the vector, there are two bits for each genotype. This function creates an output matrix by fixing the marker and collecting genotype information for each person and then repeating for all the specified markers.

Value

a list of 3 values, named "ncol", "zero", "geno".

geno is a matrix of dosages as integers. The value 0 is given to the Major allele value, 1 is given to the heterozygote value, and 2 is given to the Minor allele. In the matrix, there is usually one column for each marker in the markers_arg argument. But if there would be only the one allele 0 or 2 in the column, the column is ignored not present. There is one row for each person in the family (fam) table.

ncol Is the count of the actual number of columns in the geno matrix.

zero Is a vector with one entry per marker. The value will be 0 if the marker is not in the geno matrix. Otherwise the value is the column number in the geno matrix where the marker data appears.

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)

getgenotypesdos(ENV$markers[ENV$markers$chromosome == 1,])
getgenotypesgenabel

process the genotype matrix for specified markers and return the corresponding GenABEL genotype matrix

Description

This function calls a C++ function that does all the heavy lifting. It passes the arguments necessary for the C++ function: some from the caller’s arguments and some from data frames that are in the "global" environment, envir. From its markers_arg argument, it gets the locus_index and the index in the unified_genotype_table. From the "global" environment, envir, it gets a bit vector of compressed genotype information, allele information, and some bookkeeping related data. Note: This function also contains a dispatch/switch on the type of compression in the genotype vector. A different C++ function is called when there is compression versus when there is no compression.

Usage

getgenotypesgenabel(markers_arg, envir = ENV)

Arguments

- **markers_arg** a data.frame with the following 5 observations:
  - **locus_link** is the ordinal ranking of this marker among all loci
  - **locus_link_fill** is the position of corresponding genotype data in the unified_genotype_table
  - **MarkerName** is the text name of the marker
  - **chromosome** is the integer chromosome number
  - **position** is the integer base pair position of marker

- **envir** an environment that contains all the data frames created from the SQLite database.

Details

This function reads the genotype data in Mega2 compressed format and converts it to the GenABEL compressed format. The unified_genotype_table contains one raw vector for each person. In the vector, there are two bits for each genotype; each byte has the data for 4 markers. In GenABEL, there is one raw vector per marker, and each byte has the data for 4 persons. The C++ function does the conversion as well as adjusts the bits’ contents. For example, in GenABEL the genotype represented by bits == 0, is what Mega2 represents with 2. Doing the conversion in C++ is 10 - 20 times faster than converting the Mega2 data to PLINK .tped files and then having GenABEL read in and process/convert those files.

Value

the GenABEL gwaa.data-class object component that contains the genotype data.

Note

This function is called from Mega2ENVGenABEL; it is not intended to be called by the programmer.
getgenotypesraw

Examples

db = system.file("exdata", "seqsim.db", package="Mega2R")
ENV = read.Mega2DB(db)

aa = getgenotypesgenabel(ENV$markers[ENV$markers$chromosome == 1,])

aa

getgenotypesraw fetch genotype integer matrix for specified markers

Description

This function calls a C++ function that does all the heavy lifting. It passes the arguments necessary
for the C++ function: some from the caller’s arguments and some from data frames that are in
the "global" environment, envir. From its markers_arg argument, it gets the locus_index and the
index in the unified_genotype_table. From the "global" environment, envir, it gets a bit vector of
compressed genotype information, and some bookkeeping related data. Note: This function also
contains a dispatch/switch on the type of compression in the genotype vector. A different C++
function is called when there is compression versus when there is no compression.

Usage

getgenotypesraw(markers_arg, envir = ENV)

Arguments

markers_arg a data.frame with the following 5 observations:
  locus_link is the ordinal ranking of this marker among all loci
  locus_link_fill is the position of corresponding genotype data in the unified_genotype_table
  MarkerName is the text name of the marker
  chromosome is the integer chromosome number
  position is the integer base pair position of marker

envir an environment that contains all the data frames created from the SQLite database.

Details

The unified_genotype_table contains one raw vector for each person. In the vector, there are two
bits for each genotype. This function creates an output matrix by fixing the marker and collecting
genotype information for each person and then repeating for all the needed markers.

Value

a matrix of genotypes represented as integers. Each 32 bit integer represents contains two allele
values: the high 16 bits contains the index of allele1 and the low 16 bits contains the index of
allele2. In the matrix, there is one column for each marker in the markers_arg argument. There is
one row for each person in the family (fam) table.
Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)

# two ints in upper/lower half integer representing allele # for all persons in chromosome 1
getgenotypesraw(ENV$markers[ENV$markers$chromosome == 1,])

init_famSKATRC

Description

This populates the R data frames with the specified Mega2 SQLite database. It initializes the fam(ily) table and makes sure the person entries are unique. Finally, it generates a kinship matrix from the family data. It also stores a weighting for the common and rare variant that may be used later if NULL is specified as a weight in Mega2famSKATRC. The common weighting is the function dbeta(maf, 1, 25). The rare weighting is the function dbeta(maf, 0.5, 0.5).

Usage

init_famSKATRC(db = NULL, verbose = FALSE, ALPHA = FALSE, ...)
### init_pedgene

**See Also**

- Mega2famSKATRC

**Examples**

```r
db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = init_famSKATRC(db, verbose = FALSE)
ls(ENV)
```

**Description**

This populates the R data frames from the specified Mega2 SQLite database.

**Usage**

```r
init_pedgene(db = NULL, verbose = FALSE, traitname = "default", ...)
```

**Arguments**

- `db`: specifies the path of a Mega2 SQLite database containing study data.
- `verbose`: TRUE indicates that diagnostic printouts should be enabled. This value is saved in the returned environment.
- `traitname`: Name of the affection status trait to use to set the case/control status; default value = "default".
- `...`: fed to `dbmega2_import()`: should be bpPosMap= to select from the maps of base pairs, if the default is not desired.

**Value**

"environment" containing data frames from an SQLite database and some computed values.

**Note**

`init_pedgene` calculates schaidPed and pedPer that are used later in the Dopedgene calculation. In addition, it initializes a matrix to aid in translating a genotype allele matrix to a genotype count matrix.

It also initializes the dataframe `envir$pedgene_results` to zero rows.

**See Also**

- DOpedgene
- Mega2pedgene
- mkfam
init_SKAT

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = init_pedgene(db, traitname = "default")
ls(ENV)

Description

This populates the R data frames from the specified Mega2 SQLite database. It then prunes the samples to only include members that have a definite case or control status. Undefined samples are ignored.

Usage

init_SKAT(db = NULL, verbose = FALSE, allMarkers = FALSE, ...)

Arguments

db specifies the path of a Mega2 SQLite database containing study data.
verbose TRUE indicates that diagnostic printouts should be enabled. This value is saved in the returned environment.
allMarkers TRUE means use all markers in a given transcript even if there is no variation. FALSE means ignore markers that show no variation; this is the default.
...

Value

"environment" containing data frames from an SQLite database and some computed values.

Note

init_SKAT creates a data frame, envir$phe, of phenotype observations. In addition, it initializes a matrix to aid in translating a genotype allele matrix to a genotype count matrix.

It also initializes the data frame envir$SKAT_results to zero rows.

See Also

Mega2SKAT
**Mega2ENVGenABEL**

**Examples**

```r
db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = init.SKAT(db, verbose = FALSE, allMarkers = FALSE)
ls(ENV)
```

---

**Mega2ENVGenABEL**

*generate gwaa.data-class object*

---

**Description**

create a gwaa.data-class object from the data frames in a Mega2 environment. This function is a front end that eventually calls a C++ Rcpp function that reads the genotype data in Mega2 compressed format and converts it to the GenABEL compressed format. The results of Mega2ENVGenABEL are/should be the same as Mega2GenABEL, but the calculation is much faster, typically a factor of 10 to 20.

**Usage**

```r
Mega2ENVGenABEL(
  markers = NULL,
  force = TRUE,
  makemap = FALSE,
  sort = TRUE,
  envir = ENV
)
```

**Arguments**

- **markers**
  - data frame of markers to be processed
- **force**
  - pass value to gwaa conversion function
- **makemap**
  - pass value to gwaa conversion function
- **sort**
  - pass value to gwaa conversion function
- **envir**
  - ‘environment’ containing SQLite database and other globals

**Value**

gwaa.data-class object created from Mega2R database

**Examples**

```r
## Not run:
require("GenABEL")
db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)
gwaa = Mega2ENVGenABEL(markers=ENVmarkers[1:10,])
```
str(gwaa)
head(summary(gwaa))

## End(Not run)

### Mega2famSKATRC

**execute the CRAN famSKAT_RC function on a subset of the gene transcripts**

**Description**

If the `gene` argument is NULL, execute the `famSKAT_RC` function on the first `gs` gene transcripts (default is `gs = 1:100`). Update the `envir$famSKATRC_results` data frame with the results. Otherwise, `gene` is a string vector of genes to process. The special value `"*"` stands for all the known genes.

**Usage**

```r
Mega2famSKATRC(gs = 1:100, genes = NULL, envir = ENV, ...)
```

**Arguments**

- `gs` a subrange of the default transcripts (`refRanges`) over which to calculate the `DOfamSKATRC` function.
- `genes` a list of genes over which to calculate the `DOfamSKATRC` function. The value, `"*"`, means use all the transcripts in the selected Bioconductor database. If genes is NULL, the `gs` range of the internal `refRanges` will be used.
- `envir` 'environment' containing SQLite database and other globals.
- `...` extra arguments that are acceptable to `famSKAT_RC`. These are listed with the `DOfamSKATRC` function.

**Value**

The data frame with the results is stored in the environment and named `famSKATRC_results`, viz. `envir$famSKATRC_results`

**Note**

A helper function `SKAT3arg` is defined for the 3 argument callback function which in turn calls `DOfamSKATRC` with the appropriate arguments (including those specific to the Mega2famSKATRC function).

**See Also**

`init_famSKATRC`, `DOfamSKATRC`
Mega2gdsfmt  25

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = init_famSKATRC(db, verbose = FALSE)
ENV$verbose = FALSE
ENV$famSKATRC_results = ENV$famSKATRC_results[0, ]
Mega2famSKATRC(gs=50:60, envir=ENV, pheno=3)

# donttestcheck: try this below if there is time
Mega2famSKATRC(genes=c("CEP104"), envir=ENV, pheno=3 )

ENV$famSKATRC_results

---

Mega2gdsfmt  transcode mega2 to gdsfmt/SNP_ARRAY

Description

Reads the data frames in "envir" and builds a GDSFMT COREARRAY file from them.

Usage

Mega2gdsfmt(
  filename = "test.gds",
  markers = NULL,
  snp.order = FALSE,
  SeqArray = FALSE,
  envir = ENV
)

Arguments

filename  gdsfmt file to create
markers   data frame of markers to be processed
snp.order TRUE indicates that the "genotype" data matrix has SNP as the first index which
            changes more quickly than subsequent indices. FALSE indicates that SAMPLE
            is the the first index.
SeqArray  TRUE uses SeqArray labels for the gdsfmt vector elements. FALSE it uses
            labels shown in SNPRelate
envir     'environment' containing SQLite database and other globals

Value

writes the "filename" file containing the CoreArray data. Then returns an internal pointer, class
.gds, to the file data.
See Also
gdsfmt

Examples

```r
db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)
gdsfmtfile = file.path(where_mega2rtutorial_data(), "test.gds")
append_genotype_a = TRUE
append_genotype_b = append_genotype_c = FALSE
gn = Mega2gdsfmt(gdsfmtfile, envir=ENV)
gn
```

Mega2GenABEL
generate gwaa.data-class object from a Mega2R database

Description

Call the Mega2R functions to: create a .ped file, a .tfam file and a .phe file. Then call the GenABEL functions to process these files: the .ped and the .tfam file are processed by convert.snp.tped to produce a .ped.raw file. The latter is combined with a .phe (phenotype) file by load.gwaa.data to create a gwaa.data-class object in memory. All these files are deleted when the exits.

Usage

Mega2GenABEL(markers = NULL, mapno = 0, envir = ENV)

Arguments

- **markers**: data frame of markers to be processed
- **mapno**: specify which map index to use for physical distances
- **envir**: ‘environment’ containing SQLite database and other globals

Value
gwaa.data-class object generated from the Mega2R database

Examples

```r
## Not run:
require("GenABEL")
db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)
seqsimgwa = Mega2GenABEL(markers=ENV$markers[1:10,])

str(seqsimgwa)
head(summary(seqsimgwa))
```
Mega2GenABELtst

## End(Not run)

### Mega2GenABELtst

**compare two gwaa.data-class objects**

**Description**

Verify by fields, all the fields in two gwaa.data-class objects. Show more detailed marker information iff the coding values are different. (When comparing two gwaa.data-class objects, one native and one created via **Mega2R** sometimes when an allele frequency is .5 for both alleles, the allele order 1/2 vs 2/1 can not be currently be determined.)

**Usage**

\[
\text{Mega2GenABELtst}(\text{mega\_} = \text{mega}, \text{gwaa\_} = \text{srtda}, \text{full} = \text{TRUE}, \text{envir} = \text{ENV})
\]

**Arguments**

- `mega\_` name of first gwaa.data-class object
- `gwaa\_` name of second gwaa.data-class object
- `full` if TRUE convert genotypes to text as.character(gwaa\_@gtdata) and as.character(mega\_@gtdata). Then standardize the order for heterozygous alleles and finally compare. This step is optional because it can be rather slow.
- `envir` 'environment' containing SQLite database and other globals

**Value**

None

**Examples**

```
## Not run:
db = system.file("exdata", "seqsimm.db", package="Mega2R")
require("GenABEL")
ENV = read.Mega2DB(db)

y = Mega2ENVGenABEL()
Mega2GenABELtst(y, y, full = FALSE)

## End(Not run)

## Not run:
# donttestcheck: if you have more time, try ...
x = Mega2GenABEL()
Mega2GenABELtst(x, y, full = FALSE)

## End(Not run)
```
Mega2pedgene

Execute the pedgene function on a transcript ranges

Description

Execute the pedgene function on the first gs default gene transcript ranges (gs = 1:100). Update the envir$pedgene_results data frame with the results.

Usage

Mega2pedgene(gs = 1:100, genes = NULL, envir = ENV)

Arguments

gs a subrange of the default transcript ranges over which to calculate the Dopedsge function.

genes a list of genes over which to calculate the DOpedsge function. The value, "*", means use all the transcripts in the selected Bioconductor database. If genes is NULL, the gs range of the internal refRanges will be used.

envir 'environment' containing SQLite database and other globals

Value

None the data frame with the results is stored in the environment and named pedgene_results, viz. envir$pedgene_results

See Also

init_pedgene

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = init_pedgene(db)
ENV$verbose = TRUE
Mega2pedgene(gs = 50:60)
Mega2R-TBLS  

**Mega2R SQLite3 tables**

**Description**

This character vector indicates the names of the Mega2 SQLite3 database tables to load. (Not all of the existing tables are loaded.)

**Usage**

TBLS

**Format**

An object of class character of length 15.

**Author(s)**

Robert V Baron

---

Mega2R-TBLSFilter  

**Mega2R SQLite3 table filter**

**Description**

This list contains named values. The name corresponds to an SQLite database table. The value is a character string of column names from the "named" table that should be fetched. A table is in this list, if not all the database table columns are needed. The columns for each table are separated by commas.

**Usage**

TBLSFilter

**Format**

An object of class list of length 7.

**Note**

For the data base tables not in this list, all columns are stored in the corresponding data frame.

**Author(s)**

Robert V Baron
### Mega2Version

**Description**

This string indicates the current release of Mega2R

**Usage**

Mega2Version

**Format**

An object of class character of length 1.

**Author(s)**

Robert V Baron

---

### Mega2SKAT

**execute the CRAN SKAT function on a subset of the gene transcripts**

**Description**

Execute the SKAT function on the first `gs` default gene transcripts (`gs = 1:100`). Update the `en-vir$SKAT_results` data frame with the results.

**Usage**

Mega2SKAT(f, ty, gs = 1:100, genes = NULL, skat = SKAT::SKAT, envir = ENV, ...)

**Arguments**

- **f**
  SKAT_Null_Model formula. If this is non NULL, `envir$obj` is initialized by calling `SKAT_Null_Model(f, out_type = ty)`. If you need to specify additional arguments to the Model viz. (data, Adjustment, n.Resampling, type.Resampling) or need to use a different model viz. SKAT_NULL_emmaX, SKAT_Null_Model_ChrX set the formula to NULL, then before Mega2SKAT is called, build the model you need and assign it to `ENV$obj`.

- **ty**
  type of phenotype C/D = Continuous/Binary 5 (internal type 1/2)

- **gs**
  a subrange of the default transcripts (refRanges) over which to calculate the DOSKAT function.

- **genes**
  a list of genes over which to calculate the DOSKAT function. The value, "*", means use all the transcripts in the selected Bioconductor database. If genes is NULL, the gs range of the internal refRanges will be used.
skat alternate SKAT function, viz. SKATBinary, SKAT_CommonRare. If it is also necessary is to pass additional arguments to the SKAT function, they may be added to the end of the Mega2SKAT function and will be passed. See examples

envir 'environment' containing SQLite database and other globals
... extra arguments for SKAT

Value
None the data frame with the results is stored in the environment and named SKAT_results, viz. envir$SKAT_results

Note
The SKAT_Null_Model is called if the formula, f, is not NULL. A helper function SKAT3arg is defined for the 3 argument callback function which in turn calls DOSKAT with the appropriate arguments (including those additional to the Mega2SKAT function).

See Also
init_SKAT

Examples

```r
#donttestcheck: try this below if there is time
Mega2SKAT(ENV$phe[, 3] - 1 ~ 1, "D", kernel = "linear.weighted",
weights.beta = c(0.5, 0.5), gs=50:60 )
```

```
# donttestcheck: try this below if there is time
Mega2SKAT(ENV$phe[, 3] - 1 ~ 1, "D", kernel = "linear.weighted",
weights.beta = c(0.5, 0.5), genes=c("CEP104") )
```

```
ENV$SKAT_results
```

---

**Mega2VCF**

*generate a VCF file set for a collection of markers*

**Description**

Generate a VCF file from the specified Mega2 SQLite database. The file is named "prefix".vcf If the markers argument is.null(), the entire envir$markers set is used, otherwise markers argument MUST be rows of the markers (envir$markers) data frame. In addition, several other files are generated to hold additional database information: "prefix".fam, "prefix".freq, "prefix".map, "prefix".phe, and "prefix".pen, which contain the pedigree, allele frequency, marker genetic and physical map position, member phenotype and phenotype penetrance data.
Usage

Mega2VCF(
  prefix,
  markers = NULL,
  mapno = 0,
  alleleOrder = "default",
  envir = ENV
)

Arguments

prefix  prefix of output files including the VCF file (see Description section above). This prefix can include a path.
markers markers selected to be in the VCF output file
mapno specify which map index to use for genetic distances. The function showMapNames() will print out the internal map numbers corresponding to all the maps in the Mega2 database.
alleleOrder how to order alleles in VCF file. 'default' is Mega2order, 'minor' is minor allele freq first, 'major' is major allele freq first, and 'name' is ascending ascii character order of allele name.
envir 'environment' containing SQLite database and other globals

Value

None

Note

This code in this package illustrates how to extract the various kinds of data in the Mega2 data frames to use for further processing. Some of the data internal representations are a bit quirky but the code "explains" it all.

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)
vcfdir = file.path(where_mega2rtutorial_data(), "vcfr")
if (!dir.exists(vcfdir)) dir.create(vcfdir)
vcffile = file.path(where_mega2rtutorial_data(), "vcfr", "vcf.01")
Mega2VCF(vcffile, ENV$markers[ENV$markers$chromosome == 1, ][1:10,], envir = ENV)
list.files(vcfdir)
assemble pedigree information into a data frame

Description

Generate a data frame with a row for each person. The observations are:

- **pedigree**: family pedigree name
- **person**: person name
- **father**: father of person
- **mother**: mother of person
- **sex**: sex of person
- **trait**: value of case/control phenotype for person

Usage

```r
mkfam(brkloop = FALSE, traitname = "default", envir = ENV)
```

Arguments

- **brkloop**: I haven’t needed to set this TRUE yet. Maybe never will. If loops are broken, a person will be replaced by a dopple ganger in the same family with a different father/mother. The number of persons per family will be different when there are broken loops. Also, the person_link numbers will be different for all the persons after the first loop is broken.
- **traitname**: Name of the trait to use as case/control value; by default, "default"
- **envir**: An ’environment’ that contains all the data frames created from the SQLite database.

Value

data frame that is described above

Note

The columns of this data frame come by selecting the values after merging the data frames: `pedigree_table`, `person_table`, and `trait_table`.

Also, the father and mother columns from `person_table` are translated from the row index in the `person_table` to the corresponding name.

This function stores the data frame in the ‘environment’ and also returns it. The function `setfam()` stores the data frame into the ‘environment’ and adjusts the `genotype_table` and the `phenotype_table`.

Examples

```r
db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)
```
fam = mkfam()
fam

---

**Description**

Create the markers data frame. It contains 5 observations:

- **locus_link**: locus offset of this marker
- **locus_link_fill**: locus offset plus an accumulating fudge factor that jumps with each new chromosome because the count of markers per chromosome is force to be a multiple of 4. (This value corresponds to the offset of the marker in the unified_genotype_table.)
- **MarkerName**: name of the marker
- **chromosome**: chromosome number of the marker
- **position**: base pair position of the marker (selected by bpPosMap[below])

**Usage**

```
mkMarkers(bpPosMap = 1, envir = ENV)
```

**Arguments**

- **bpPosMap**: An integer that indicates the map (index) to use to merge the chromosome/position fields from the map_table data frame to the marker_table data frame. See showMapNames() for the string name to index mapping.
- **envir**: an environment that contains all the data frames created from the SQLite database.

**Details**

Select a map (index) from the map_table to merge with the select marker_table data frame to make the marker data frame. See showMapNames() for the string name to index mapping.

**Value**

None

**Examples**

```
db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db, verbose = FALSE)

mkMarkers(1)
ENV$markers
```
**mkphenotype**

`mkphenotype` generate a phenotype data frame

**Description**

Convert data in phenotype_table to a data frame of columns that are phenotypes. The columns may be affection status or quantitative values.

**Usage**

```r
mkphenotype(envir)
```

**Arguments**

- `envir` "environment" containing SQLite database and other globals

**Value**

is a data frame with FID column, then IID column, and then an additional column for each phenotype.

**Examples**

```r
db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)
out = mkphenotype()
```

------

**read.Mega2DB**

load Mega2 database and initialize family data frame and markers data frame

**Description**

Call dbmega2_import() with the specified database and create an 'environment', with the SQLite table data loaded into data frames. Also run mkfam() to create the pedigree data frame `fam` and then store it with setfam(). setfam() modifies the unified_genotype_table (and phenotype_table) to match the family members that remain.

**Usage**

```r
read.Mega2DB(db, ...)
```
**Arguments**

- **db**  
  specify SQLite database to load
- 
  additional arguments to pass to dbmega2_import

**Value**

an 'environment' that contains all the data frames created from the SQLite database.

**Note**

By default, mkfam will remove one of each person that was replicated to break loops in the pedigree, see mkfam for details. If you want to leave loops broken, the code is available, but you will have to write your own version of read.Mega2DB with a different invocation of mkfam().

**Examples**

```
db = system.file("exdata", "seqsim.db", package="Mega2R")
ENV = read.Mega2DB(db, verbose = TRUE)
```

---

**setAnnotations**

*set default name of transcription database and name of database mapping gene name to entrez gene id*

**Description**

This function takes two string parameters: one to specify entrez gene ids to transcripts, the other to map gene names to entrez gene id's.

**Usage**

```
setAnnotations(txdb, entrezGene, envir = ENV)
```

**Arguments**

- **txdb**  
  name of Bioconductor transcription database.
- **entrezGene**  
  name of Bioconductor mapping of gene name or gene alias to entrez gene id
- **envir**  
  an 'environment' that contains all the data frames created from the SQLite database.

**Value**

None

**Note**

Mega2R will take care to load the necessary databases, but you will have to install them from Bioconductor. This is explained at length in the package Vignette.
Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)

setAnnotations("TxDb.Hsapiens.UCSC.hg19.knownGene", "org.Hs.eg.db")

ENV$txdb
ENV$entrezGene

setfam replace the pedigree data frame

Description

You should first modify the fam data frame to filter the members you need to remove. (For example, you might want to delete members that have an unknown case/control status.) This function takes a new data frame of pedigree information and replaces the fam data frame in the 'environment' with it. Additionally, changing fam data frame will filter the genotypes data frame to only contain persons matching those in the fam data frame. setfam also filters for the phenotype data records.

Usage

setfam(fam, envir = ENV)

Arguments

fam data frame of family information filtered from fam data frame (generated by mkfam).
envir an 'environment' that contains all the data frames created from the SQLite database.

Value

None

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)

fam = mkfam()
# remove founders
fam = fam[ !( (fam[, 5] == fam[, 6]) & (fam[, 5] == 0)), ]
setfam(fam)

ENV$fam
Description

This function sets the default list of ranges used by `applyFnToRanges`. `applyFnToRanges` examines each range and the set of markers that fall within the range will be processed.

Usage

```r
setRanges(ranges, indices, envir = ENV)
```

Arguments

- `ranges`: a data frame that contains at least 4 observations: a name, a chromosome, a start base pair position and an end base pair position.
- `indices`: a vector of 3 or 4 integers that specify the chromosome column, start base pair column, end base pair column of range data frame and lastly the name column. If the vector only contains 3 integers, a name will be generated from the three range elements and it will be appended to the ranges and the last range column will be added to the indices.
- `envir`: an 'environment' that contains all the data frames created from the SQLite database.

Value

None

Examples

```r
db = system.file("exdata", "seqsim.db", package="Mega2R")
ENV = read.Mega2DB(db)

ranges = matrix(c(1, 2240000, 2245000, 1, 2245000, 2250000, 1, 3760000, 3761000, 1, 3761000, 3762000, 1, 3762000, 3763000, 1, 3763000, 3764000, 1, 3764000, 3765000, 1, 3765000, 3766000, 1, 3766000, 3767000, 1, 3767000, 3768000, 1, 3768000, 3769000, 1, 3769000, 3770000),
nocol = 3, nrow = 12, byrow = TRUE)

setRanges(ranges, 1:3)
```
showMapNames

ranges = matrix(c(1, 2240000, 2245000, 
                  1, 2245000, 2250000, 
                  1, 3760000, 3761000, 
                  1, 3761000, 3762000, 
                  1, 3762000, 3763000, 
                  1, 3763000, 3764000, 
                  1, 3764000, 3765000, 
                  1, 3765000, 3763760, 
                  1, 3763760, 3767000, 
                  1, 3767000, 3768000, 
                  1, 3768000, 3769000, 
                  1, 3769000, 3770000), 
                  ncol = 3, nrow = 12, byrow = TRUE)
ranges = data.frame(ranges)
ranges$name = LETTERS[1:12]
names(ranges) = c("chr", "start", "end", "name")
setRanges(ranges, 1:4)
ENV$refRanges

Description
Mega2R allows several different physical and genetic maps to be stored and used to select positions. This function shows the association between map number and map name.

Usage
showMapNames(envir = ENV)

Arguments

envir an environment that contains all the data frames created from the SQLite database.

Value
None

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)
showMapNames()
**showMega2ENV**

*Mega2R environment, viz. data frames and related info.*

**Description**

Mega2R uses an environment to store the data frames when it reads SQLite database tables. This function shows the data frames and their sizes; it also shows the count of samples and markers in the database. Note: It is not necessary to provide an argument, if the environment is named `ENV`.

**Usage**

```r
showMega2ENV(envir = ENV)
```

**Arguments**

- `envir` an environment that contains all the data frames created from the SQLite database.

**Value**

None

**Examples**

```r
db = system.file("exdata", "seqsim.db", package="Mega2R")
ENV = read.Mega2DB(db)
showMega2ENV()
```

---

**showPhenoNames**

*show the association between index no and phenotype*

**Description**

Mega2R stores several phenotypes, both affective and quantitative. This function displays the mapping between phenotype (name), index, and the phenotype type (affection or quantitative).

**Usage**

```r
showPhenoNames(envir = ENV)
```

**Arguments**

- `envir` an environment that contains all the data frames created from the SQLite database.
Value

None

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)
showPhenoNames()


uniqueFamMember regenerate fam data frame with unique values in member column

Description

Reads the fam data frame in "envir" and returns a new one with unique entries in the member column

Usage

uniqueFamMember(envir = ENV)

Arguments

envir 'environment' containing SQLite database and other globals

Value

a data frame with columns the same as the "fam" data frame but with the member column containing unique entries

See Also

mkfam

Examples

db = system.file("exdata", "seqsimm.db", package="Mega2R")
ENV = read.Mega2DB(db)
setfam(uniqueFamMember(envir = ENV))
where_mega2rtutorial_data

show directory of tutorial data

Description

This function shows the directory the Mega2Rtutorial (inst/exdata) was copied to.

Usage

where_mega2rtutorial_data(dir = file.path(tempdir(), "Mega2Rtutorial"))

Arguments

dir The directory to store the tutorial data to. By default, this is tempdir()/Mega2Rtutorial

Value

dir tutorial to hold vignette

Examples

directory = where_mega2rtutorial_data()
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