Package `genio`  

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Title Genetics Input/Output Functions

Version 1.0.25

Description Implements readers and writers for file formats associated with genetics data. Reading and writing Plink BED/BIM/FAM and GCTA binary GRM formats is fully supported, including a lightning-fast BED reader and writer implementations. Other functions are ‘readr’ wrappers that are more constrained, user-friendly, and efficient for these particular applications; handles Plink and Eigenstrat tables (FAM, BIM, IND, and SNP files). There are also make functions for FAM and BIM tables with default values to go with simulated genotype data.

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Encoding UTF-8

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Imports readr (>= 2.0.0), tibble, dplyr, Rcpp

LinkingTo Rcpp

Suggests testthat, knitr, rmarkdown, BEDMatrix, snpStats, lobstr

VignetteBuilder knitr

URL https://github.com/OchoaLab/genio

BugReports https://github.com/OchoaLab/genio/issues

NeedsCompilation yes

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count_lines

Count the number of lines of a file

Description

This function returns the number of lines in a file. It is intended to result in fast retrieval of numbers of individuals (from FAM or equivalent files) or loci (BIM or equivalent files) when the input files are extremely large and no other information is required from these files. This code uses C++ to quickly counts lines (like linux’s wc -l but this one is cross-platform).

Usage

count_lines(file, ext = NA, verbose = TRUE)
**delete_files_grm**

Arguments

- **file**: The input file path to read (a string).
- **ext**: An optional extension. If NA (default), file is expected to exist as-is. Otherwise, if file doesn’t exist and the extension was missing, then this extension is added.
- **verbose**: If TRUE (default), writes a message reporting the file whose lines are being counted (after adding extensions if it was needed).

Details

Note: this function does not work correctly with compressed files (they are not uncompressed prior to counting newlines).

Value

The number of lines in the file.

Examples

```r
# count number of individuals from an existing plink *.fam file
file <- system.file("extdata", "sample.fam", package = "genio", mustWork = TRUE)
n_ind <- count_lines(file)

# count number of loci from an existing plink *.bim file
file <- system.file("extdata", "sample.bim", package = "genio", mustWork = TRUE)
m_loci <- count_lines(file)
```

---

**delete_files_grm**

**Delete all GCTA binary GRM files**

Description

This function deletes each of the GCTA binary GRM files (grm.bin, grm.N.bin, and grm.id extensions) given the shared base file path, warning if any of the files did not exist or if any were not successfully deleted.

Usage

```r
delete_files_grm(file)
```

Arguments

- **file**: The shared file path (excluding extensions: grm.bin, grm.N.bin, or grm.id).
delete_files_phen

Delete PHEN files

Description

This function deletes a PHEN files given the base file path (without extension), warning if the file did not exist or if it was not successfully deleted.

Usage

deleter_files_phen(file)

Arguments

file The base file path (excluding phen extension).

Value

Nothing

Examples

# if you want to delete "data.phen", run like this:
# delete_files_phen("data")

# The following example is more awkward
# because (only for these examples) the package must create *temporary* files to actually delete

define dummy GRM files
file <- tempfile('delete-me-test') # no extension
# add each extension and create empty files
file.create( paste0(file, ".grm.bin") )
file.create( paste0(file, ".grm.N.bin") )
file.create( paste0(file, ".grm.id") )

# delete the GRM files we just created
delete_files_grm(file)
### delete_files_plink

`delete_files_plink`  
Delete all Plink binary files

---

**Description**

This function deletes each of the Plink binary files (`bed`, `bim`, `fam` extensions) given the shared base file path, warning if any of the files did not exist or if any were not successfully deleted.

**Usage**

```r
delete_files_plink(file)
```

**Arguments**

- `file`  
The shared file path (excluding extensions: `bed`, `bim`, `fam`).

**Value**

Nothing

**Examples**

```r
# if you want to delete "data.bed", "data.bim" and "data.fam", run like this:
# delete_files_plink("data")

# The following example is more awkward
# because (only for these examples) the package must create *temporary* files to actually delete

# create dummy BED/BIM/FAM files
file <- tempfile('delete-me-test') # no extension
# add each extension and create empty files
file.create( paste0(file, '.bed') )
file.create( paste0(file, '.bim') )
file.create( paste0(file, '.fam') )

# delete the BED/BIM/FAM files we just created
delete_files_plink(file)
```
Description

This package fully supports reading and writing Plink BED/BIM/FAM and GCTA GRM files, as illustrated below. These functions make it easy to create dummy annotation tables to go with simulated genotype data too. Lastly, there is functionality to read and write Eigenstrat tables.

Author(s)

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See Also

Useful links:

- https://github.com/OchoaLab/genio
- Report bugs at https://github.com/OchoaLab/genio/issues

Examples

```r
# read existing BED/BIM/FAM files

file <- system.file( "extdata", 'sample.bed', package = "genio", mustWork = TRUE )
# read genotypes and annotation tables
plink_data <- read_plink( file )
# genotypes
X <- plink_data$X
# locus annotations
bim <- plink_data$bim
# individual annotations
fam <- plink_data$fam

# the same works without .bed extension
file <- sub( '\\.[^\s]+\.', '', file ) # remove extension
plink_data <- read_plink( file )

# write data into new BED/BIM/FAM files
file_out <- tempfile( 'delete-me-example' )
write_plink( file_out, X, bim, fam )

# delete example files when done
delete_files_plink( file_out )

# read sample GRM files
file <- system.file( "extdata", 'sample.grm.bin', package = "genio", mustWork = TRUE )
```
file <- sub( '\.grm\bin\$', '', file ) # remove extension from this path on purpose
obj <- read_grm( file )
# the kinship matrix
kinship <- obj$kinship
# the pair sample sizes matrix
M <- obj$M
# the fam and ID tibble
fam <- obj$fam

# write data into new GRM files
write_grm( file_out, kinship, M = M, fam = fam )

# delete example files when done
delete_files_grm( file_out )

# other functions not shown here allow reading and writing individual files,
# creating dummy tables to go with simulated genotypes,
# requiring the existence of these files,
# and reading and writing of Eigenstrat tables too.

---

**geno_to_char**

Convert a genotype matrix from numeric to character codes

**Description**

Given the genotype matrix `X` and `bim` table (as they are parsed by `read_plink()`), this outputs a matrix of the same dimensions as `X` but with the numeric codes (all values in 0, 1, 2) translated to human-readable character codes (such as 'A/A', 'A/G', 'G/G', depending on which are the two alleles at the locus as given in the `bim` table, see return value).

**Usage**

`geno_to_char(X, bim)`

**Arguments**

- `X`  
  The genotype matrix. It must have values only in 0, 1, 2, and NA.

- `bim`  
  The variant table. It is required to have the same number of rows as `X`, and to have at least two named columns `ref` and `alt` (alleles 1 and 2 in a plink BIM table). These alleles can be arbitrary strings (i.e. not just SNPs but also indels, any single or multicharacter code, or even blank strings) except the forward slash character ('/') is not allowed anywhere in these strings (function stops if a slash is present), since in the output it is the delimiter string. `ref` and `alt` alleles must be different at each locus.
ind_to_fam

Value

The genotype matrix reencoded as strings. At one locus, if the two alleles (ref and alt) are 'A' and 'B', then the genotypes in the input are encoded as characters as: 0 -> 'B/B', 1 -> 'B/A', and 2 -> 'A/A'. Thus, the numeric encoding counts the reference allele dosage. NA values in input X remain NA in the output. If the input genotype matrix had row and column names, these are inherited by the output matrix.

See Also

read_plink(), read_bed(), read_bim().

Examples

# a numeric/dosage genotype matrix with two loci (rows)
# and three individuals (columns)
X <- rbind( 0:2, c(0, NA, 2) )
# corresponding variant table (minimal case with just two required columns)
library(tibble)
bim <- tibble( ref = c('A', 'G'), alt = c('C', 'GT') )

# genotype matrix translated as characters
X_char <- geno_to_char( X, bim )
X_char

ind_to_fam

Convert an Eigenstrat IND tibble into a Plink FAM tibble

Description

This function takes an existing IND tibble and creates a FAM tibble with the same information and dummy values for missing data. In particular, the output FAM tibble will contain these columns with these contents (IND only contain id, sex, and label, so there is no loss of information):

- fam: IND label
- id: IND id
- pat: 0 (missing paternal ID)
- mat: 0 (missing maternal ID)
- sex: IND sex converted to Plink integer codes via sex_to_int()
- peno: 0 (missing phenotype)

Usage

ind_to_fam(ind)

Arguments

ind The input Eigenstrat IND tibble to convert.
make_bim

Value
A Plink FAM tibble.

See Also

sex_to_int()

Eigenstrat IND format reference: https://github.com/DReichLab/EIG/tree/master/CONVERTF

Plink FAM format reference: https://www.cog-genomics.org/plink/1.9/formats#fam

Examples

# create a sample IND tibble
library(tibble)
ind <- tibble(
  id = 1:3,
  sex = c('U', 'M', 'F'),
  label = c(1, 1, 2)
)
# convert to FAM
fam <- ind_to_fam(ind)
# inspect:
fam

make_bim
Create a Plink BIM tibble

Description
This function simplifies the creation of Plink BIM-formatted tibbles, which autocompletes missing information if a partial tibble is provided, or generates a completely made up tibble if the number of individuals is provided. The default values are most useful for simulated genotypes, where IDs can be made up but must be unique, and there are no chromosomes, positions, or particular reference or alternative alleles.

Usage

make_bim(tib, n = NA)

Arguments

tib          The input tibble (optional). Missing columns will be autocompleted with reasonable values that are accepted by Plink and other external software. If missing, all will be autocompleted, but n is required.

n            The desired number of loci (rows). Required if tib is missing; otherwise it is ignored.
Details

Autocompleted column values:

- chr: 1 (all data is on a single chromosome)
- id: 1:n
- posg: 0 (missing)
- pos: 1:n
- ref: 1
- alt: 2

Note that n is either given directly or obtained from the input tibble.

Value

The input tibble with autocompleted columns and columns in default order, or the made up tibble if only the number of individuals was provided. The output begins with the standard columns in standard order: chr, id, posg, pos, ref, alt. Additional columns in the input tibble are preserved but placed after the standard columns.

See Also

Plink BIM format reference: [https://www.cog-genomics.org/plink/1.9/formats#bim](https://www.cog-genomics.org/plink/1.9/formats#bim)

Examples

```r
# create a synthetic tibble for 10 loci
# (most common use case)
bim <- make_bim(n = 10)

# manually create a partial tibble with only chromosomes defined
library(tibble)
bim <- tibble(chr = 0:2)
# autocomplete the rest of the columns
bim <- make_bim(bim)
```

Description

This function simplifies the creation of Plink FAM-formatted tibbles, which autocompletes missing information if a partial tibble is provided, or generates a completely made up tibble if the number of individuals is provided. The default values are most useful for simulated genotypes, where IDs can be made up but must be unique, and there are no parents, families, gender, or phenotype.
Usage

make_fam(tib, n = NA)

Arguments

tib  The input tibble (optional). Missing columns will be autocompleted with reasonable values that are accepted by Plink and other external software. If missing, all will be autocompleted, but n is required.
n  The desired number of individuals (rows). Required if tib is missing; otherwise it is ignored.

Details

Autocompleted column values:

- fam: 1:n
- id: 1:n
- pat: 0 (missing)
- mat: 0 (missing)
- sex: 0 (missing)
- pheno: 0 (missing)

Note that n is either given directly or obtained from the input tibble.

Value

The input tibble with autocompleted columns and columns in default order, or the made up tibble if only the number of individuals was provided. The output begins with the standard columns in standard order: fam, id, pat, mat, sex, pheno. Additional columns in the input tibble are preserved but placed after the standard columns.

See Also

Plink FAM format reference: https://www.cog-genomics.org/plink/1.9/formats#fam

Examples

# create a synthetic tibble for 10 individuals
# (most common use case)
fam <- make_fam(n = 10)

# manually create a partial tibble with only phenotypes defined
library(tibble)
fam <- tibble(pheno = 0:2)
# autocomplete the rest of the columns
fam <- make_fam(fam)
read_bed  

Read a genotype matrix in Plink BED format

Description

This function reads genotypes encoded in a Plink-formatted BED (binary) file, returning them in a standard R matrix containing genotypes encoded numerically as dosages (values in \( c(0, 1, 2, NA) \)). Each genotype per locus (\( m \) loci) and individual (\( n \) total) counts the number of reference alleles, or NA for missing data. No *.fam or *.bim files are read by this basic function. Since BED does not encode the data dimensions internally, these values must be provided by the user.

Usage

```r
read_bed(
  file,
  names_loci = NULL,
  names_ind = NULL,
  m_loci = NA,
  n_ind = NA,
  ext = "bed",
  verbose = TRUE
)
```

Arguments

- **file**: Input file path. *.bed extension may be omitted (will be added automatically if file doesn’t exist but file.bed does). See ext option below.
- **names_loci**: Vector of loci names, to become the row names of the genotype matrix. If provided, its length sets \( m \) loci below. If NULL, the returned genotype matrix will not have row names, and \( m \) loci must be provided.
- **names_ind**: Vector of individual names, to become the column names of the genotype matrix. If provided, its length sets \( n \) ind below. If NULL, the returned genotype matrix will not have column names, and \( n \) ind must be provided.
- **m_loci**: Number of loci in the input genotype table. Required if names_loci = NULL, as its value is not deducible from the BED file itself. Ignored if names_loci is provided.
- **n_ind**: Number of individuals in the input genotype table. Required if names_ind = NULL, as its value is not deducible from the BED file itself. Ignored if names_ind is provided.
- **ext**: The desired file extension (default "bed"). Ignored if file points to an existing file. Set to NA to force file to exist as-is.
- **verbose**: If TRUE (default) function reports the path of the file being read (after autocompleting the extension).
Details

The code enforces several checks to validate data given the requested dimensions. Errors are thrown if file terminates too early or does not terminate after genotype matrix is filled. In addition, as each locus is encoded in an integer number of bytes, and each byte contains up to four individuals, bytes with fewer than four are padded. To agree with other software (plink2, BEDMatrix), byte padding values are ignored (may take on any value without causing errors).

This function only supports locus-major BED files, which are the standard for modern data. Format is validated via the BED file’s magic numbers (first three bytes of file). Older BED files can be converted via Plink.

Value

The m-by-n genotype matrix.

See Also

read_plink() for reading a set of BED/BIM/FAM files.

geno_to_char() for translating numerical genotypes into more human-readable character encodings.

Plink BED format reference: https://www.cog-genomics.org/plink/1.9/formats#bed

Examples

# first obtain data dimensions from BIM and FAM files
# all file paths
file_bed <- system.file("extdata", 'sample.bed', package = "genio", mustWork = TRUE)
file_bim <- system.file("extdata", 'sample.bim', package = "genio", mustWork = TRUE)
file_fam <- system.file("extdata", 'sample.fam', package = "genio", mustWork = TRUE)
# read annotation tables
bim <- read_bim(file_bim)
fam <- read_fam(file_fam)

# read an existing Plink *.bim file
# pass locus and individual IDs as vectors, setting data dimensions too
X <- read_bed(file_bed, bim$id, fam$id)
X

# can specify without extension
file_bed <- sub("\.[bed$]", "", file_bed) # remove extension from this path on purpose
file_bed # verify .bed is missing
X <- read_bed(file_bed, bim$id, fam$id) # loads too!
X
Description

This function reads a standard Plink *.bim file into a tibble with named columns. It uses `readr::read_table()` to do it efficiently.

Usage

```r
read_bim(file, verbose = TRUE)
```

Arguments

- `file` Input file (whatever is accepted by `readr::read_table()`). If file as given does not exist and is missing the expected *.bim extension, the function adds the .bim extension and uses that path if that file exists. Additionally, the .gz extension is added automatically if the file (after *.bim extension is added as needed) is still not found and did not already contain the .gz extension and adding it points to an existing file.
- `verbose` If TRUE (default) function reports the path of the file being loaded (after auto-completing the extensions).

Value

A tibble with columns: chr, id, posg, pos, ref, alt

See Also

- `read_plink()` for reading a set of BED/BIM/FAM files.
- Plink BIM format reference: [https://www.cog-genomics.org/plink/1.9/formats#bim](https://www.cog-genomics.org/plink/1.9/formats#bim)

Examples

```r
# to read "data.bim", run like this:
# bim <- read_bim("data")
# this also works
# bim <- read_bim("data.bim")

# The following example is more awkward
# because package sample data has to be specified in this weird way:

# read an existing Plink *.bim file
file <- system.file("extdata", 'sample.bim', package = "genio", mustWork = TRUE)
bim <- read_bim(file)
bim

# can specify without extension
```
read_eigenvec

```r
file <- sub('\\.bim$', '', file) # remove extension from this path on purpose
file # verify .bim is missing
bim <- read_bim(file) # loads too!
```

---

### Description

This function reads a Plink eigenvec file, parsing columns strictly. First two must be ‘fam’ and ‘id’, which are strings, and all remaining columns (eigenvectors) must be numeric.

### Usage

```r
read_eigenvec(file, ext = "eigenvec", comment = ",", verbose = TRUE)
```

### Arguments

- **file**: The input file path, potentially excluding extension.
- **ext**: File extension (default "eigenvec") can be changed if desired. Set to NA to force file to exist as-is.
- **comment**: A string used to identify comments. Any text after the comment characters will be silently ignored. Passed to `readr::read_table()`. "#" (default) works for Plink 2 eigenvec files, which have a header lines that starts with this character (the header is therefore ignored).
- **verbose**: If TRUE (default) function reports the path of the file being written (after auto-completing the extension).

### Value

A list with two elements:

- **eigenvec**: A numeric R matrix containing the parsed eigenvectors
- **fam**: A tibble with two columns, fam and id, which are the first two columns of the parsed file.

### See Also

- `write_eigenvec()` for writing an eigenvec file.
- Plink 1 eigenvec format reference: [https://www.cog-genomics.org/plink/1.9/formats#eigenvec](https://www.cog-genomics.org/plink/1.9/formats#eigenvec)
- Plink 2 eigenvec format reference: [https://www.cog-genomics.org/plink/2.0/formats#eigenvec](https://www.cog-genomics.org/plink/2.0/formats#eigenvec)
- GCTA eigenvec format reference: [https://cnsgenomics.com/software/gcta/#PCA](https://cnsgenomics.com/software/gcta/#PCA)
Examples

# to read "data.eigenvec", run like this:
# data <- read_eigenvec("data")
# this also works
# data <- read_eigenvec("data.eigenvec")

# either way you get a list with these two items:
# numeric eigenvector matrix
# data$eigenvec
# fam/id tibble
# data$fam

# The following example is more awkward
# because package sample data has to be specified in this weird way:

# read an existing *.eigenvec file created by GCTA
file <- system.file("extdata", 'sample-gcta.eigenvec', package = "genio", mustWork = TRUE)
data <- read_eigenvec(file)
# numeric eigenvector matrix
data$eigenvec
# fam/id tibble
data$fam

# can specify without extension
file <- sub(\".eigenvec\", '', file) # remove extension from this path on purpose
file # verify .eigenvec is missing
data <- read_eigenvec(file) # load it anyway!
data$eigenvec

# read an existing *.eigenvec file created by Plink 2
file <- system.file("extdata", 'sample-plink2.eigenvec', package = "genio", mustWork = TRUE)
data <- read_eigenvec(file)
# numeric eigenvector matrix
data$eigenvec
# fam/id tibble
data$fam


---

read_fam

Read Plink *.fam files

Description

This function reads a standard Plink *.fam file into a tibble with named columns. It uses readr::read_table() to do it efficiently.

Usage

read_fam(file, verbose = TRUE)
Arguments

file  Input file (whatever is accepted by `readr::read_table()`). If file as given does not exist and is missing the expected *.fam extension, the function adds the .fam extension and uses that path if that file exists. Additionally, the .gz extension is added automatically if the file (after *.fam extension is added as needed) is still not found and did not already contain the .gz extension and adding it points to an existing file.

verbose  If TRUE (default) function reports the path of the file being loaded (after autocompleting the extensions).

Value

A tibble with columns: fam, id, pat, mat, sex, pheno.

See Also

`read_plink()` for reading a set of BED/BIM/FAM files.

Plink FAM format reference: [https://www.cog-genomics.org/plink/1.9/formats#fam](https://www.cog-genomics.org/plink/1.9/formats#fam)

Examples

```r
# to read "data.fam", run like this:
# fam <- read_fam("data")
# this also works
# fam <- read_fam("data.fam")

# The following example is more awkward
# because package sample data has to be specified in this weird way:

# read an existing Plink *.fam file
file <- system.file("extdata", 'sample.fam', package = "genio", mustWork = TRUE)
fam <- read_fam(file)
fam

# can specify without extension
file <- sub("\..fam\", ",", file) # remove extension from this path on purpose
file # verify .fam is missing
fam <- read_fam(file) # load it anyway!
fam
```

Description

This function reads a GCTA Genetic Relatedness Matrix (GRM, i.e. kinship) set of files in their binary format, returning the kinship matrix and, if available, the corresponding matrix of pair sample sizes (non-trivial under missingness) and individuals table.
read_grm

Usage

read_grm(name, n_ind = NA, verbose = TRUE)

Arguments

name
The base name of the input files. Files with that base and extensions .grm.bin, .grm.N.bin, and .grm.id are read if they exist. Only .grm.bin is absolutely required; .grm.id can be substituted by the number of individuals (see below); .grm.N.bin is entirely optional.

n_ind
The number of individuals, required if the file with the extension .grm.id is missing. If the file with the .grm.id extension is present, then this n_ind is ignored.

verbose
If TRUE (default), function reports the path of the files being loaded.

Value

A list with named elements:

- kinship: The symmetric $n$-times-$n$ kinship matrix (GRM). Has IDs as row and column names if the file with extension .grm.id was available.
- M: The symmetric $n$-times-$n$ matrix of pair sample sizes (number of non-missing loci pairs), if the file with extension .grm.N.bin was available. Has IDs as row and column names if the file with extension .grm.id was available.
- fam: A tibble with two columns: fam and id, same as in Plink FAM files. Returned if the file with extension .grm.id was available.

See Also

Greatly adapted from sample code from GCTA: https://cnsgenomics.com/software/gcta/
#MakingaGRM

Examples

# to read "data.grm.bin" and etc, run like this:
# obj <- read_grm("data")
# obj$kinship # the kinship matrix
# obj$M # the pair sample sizes matrix
# obj$fam # the fam and ID tibble

# The following example is more awkward
# because package sample data has to be specified in this weird way:

# read an existing set of GRM files
file <- system.file("extdata", 'sample.grm.bin', package = "genio", mustWork = TRUE)
file <- sub("\.grm\.bin$", '', file) # remove extension from this path on purpose
obj <- read_grm(file)
stopifnot( !is.null(obj$kinship ) ) # the kinship matrix
stopifnot( !is.null(obj$M ) ) # the pair sample sizes matrix
stopifnot( !is.null(obj$fam ) ) # the fam and ID tibble
**read_ind**

*Read Eigenstrat *.ind files*

**Description**

This function reads a standard Eigenstrat *.ind file into a tibble. It uses `readr::read_table()` to do it efficiently.

**Usage**

```r
read_ind(file, verbose = TRUE)
```

**Arguments**

- **file**: Input file (whatever is accepted by `readr::read_table()`). If file as given does not exist and is missing the expected *.ind extension, the function adds the .ind extension and uses that path if that file exists. Additionally, the .gz extension is added automatically if the file (after *.ind extension is added as needed) is still not found and did not already contain the .gz extension and adding it points to an existing file.

- **verbose**: If TRUE (default), function reports the path of the file being loaded (after auto-completing the extensions).

**Value**

A tibble with columns: id, sex, label.

**See Also**

Eigenstrat IND format reference: [https://github.com/DReichLab/EIG/tree/master/CONVERTF](https://github.com/DReichLab/EIG/tree/master/CONVERTF)

**Examples**

```r
# to read "data.ind", run like this:
# ind <- read_ind("data")
# this also works
# ind <- read_ind("data.ind")

# The following example is more awkward
# because package sample data has to be specified in this weird way:

# read an existing Eigenstrat *.ind file
file <- system.file("extdata", 'sample.ind', package = "genio", mustWork = TRUE)
ind <- read_ind(file)
ind

# can specify without extension
file <- sub("\ind$", '', file) # remove extension from this path on purpose
file # verify .ind is missing
```
read_matrix

Read a numerical matrix file into an R matrix

Description

Reads a matrix file under strict assumptions that it is entirely numeric and there are no row or column names present in this file. It uses `readr::read_table()` to do it efficiently. Intended for outputs such as those of admixture inference approaches.

Usage

```r
read_matrix(file, ext = "txt", verbose = TRUE)
```

Arguments

- **file**: Input file (whatever is accepted by `readr::read_table()`). If file as given does not exist and is missing the expected extension (see `ext` below), the function adds the extension and uses that path if that file exists. Additionally, the `.gz` extension is added automatically if the file (after the extension is added as needed) is still not found and did not already contain the `.gz` extension and adding it points to an existing file.

- **ext**: The desired file extension. Ignored if `file` points to an existing file. Set to `NA` to force `file` to exist as-is.

- **verbose**: If `TRUE` (default) function reports the path of the file being loaded (after autocompleting the extensions).

Value

A numeric matrix without row or column names.

See Also

`write_matrix()`, the inverse function.

Examples

```r
# to read "data.txt", run like this:
# mat <- read_matrix("data")
# this also works
# mat <- read_matrix("data.txt")

# The following example is more awkward
# because package sample data has to be specified in this weird way:
```
# read an existing matrix *.txt file
file <- system.file("extdata", 'sample-Q3.txt', package = "genio", mustWork = TRUE)
mat <- read_matrix(file)
mat

# can specify without extension
file <- sub('\.txt$', '', file) # remove extension from this path on purpose
file # verify .txt is missing
mat <- read_matrix(file) # load it anyway!
mat

---

## read_phen

**Read *.phen files**

### Description

This function reads a standard *.phen file into a tibble. It uses `readr::read_table()` to do it efficiently. GCTA and EMMAX use this format.

### Usage

```r
read_phen(file, verbose = TRUE)
```

### Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>file</code></td>
<td>Input file (whatever is accepted by <code>readr::read_table()</code>). If file as given does not exist and is missing the expected *.phen extension, the function adds the .phen extension and uses that path if that file exists. Additionally, the .gz extension is added automatically if the file (after *.phen extension is added as needed) is still not found and did not already contain the .gz extension and adding it points to an existing file.</td>
</tr>
<tr>
<td><code>verbose</code></td>
<td>If TRUE (default), function reports the path of the file being loaded (after autocompleting the extensions).</td>
</tr>
</tbody>
</table>

### Value

A tibble with columns: fam, id, pheno.

### See Also

GCTA PHEN format reference: https://cnsgenomics.com/software/gcta/#GREMLanalysis
Examples

# to read "data.phen", run like this:
# phen <- read_phen("data")
# this also works
# phen <- read_phen("data.phen")

# The following example is more awkward
# because package sample data has to be specified in this weird way:

# read an existing plink *.phen file
file <- system.file("extdata", 'sample.phen', package = "genio", mustWork = TRUE)
phen <- read_phen(file)
phen

# can specify without extension
file <- sub('\\.phen$', '', file) # remove extension from this path on purpose
file # verify .phen is missing
phen <- read_phen(file) # load it anyway!
phen

read_plink

Read genotype and sample data in a Plink BED/BIM/FAM file set.

Description

This function reads a genotype matrix ($X$, encoded as reference allele dosages) and its associated locus ($bim$) and individual ($fam$) data tables in the three Plink files in BED, BIM, and FAM formats, respectively. All inputs must exist or an error is thrown. This function is a wrapper around the more basic functions `read_bed()`, `read_bim()`, `read_fam()`, which simplifies data parsing and additionally better guarantees data integrity. Below suppose there are $m$ loci and $n$ individuals.

Usage

```r
read_plink(file, verbose = TRUE)
```

Arguments

- **file**: Input file path, without extensions (each of .bed, .bim, .fam extensions will be added automatically as needed). Alternatively, input file path may have .bed extension (but not .bim, .fam, or other extensions).
- **verbose**: If TRUE (default), function reports the paths of the files being read (after auto-completing the extensions).

Value

A named list with items in this order: $X$ (genotype matrix, see description in return value of `read_bed()`), $bim$ (tibble, see `read_bim()`), $fam$ (tibble, see `read_fam()`). $X$ has row and column names corresponding to the id values of the $bim$ and $fam$ tibbles.
**read_snp**

See Also

- `read_bed()`, `read_bim()`, and `read_fam()` for individual parsers of each input table, including a description of each object returned.
- `geno_to_char()` for translating numerical genotypes into more human-readable character encodings.

Plink BED/BIM/FAM format reference: [https://www.cog-genomics.org/plink/1.9/formats](https://www.cog-genomics.org/plink/1.9/formats)

Examples

```r
# to read "data.bed" etc, run like this:
# obj <- read_plink("data")
# this also works
# obj <- read_plink("data.bed")
#
# you get a list with these three items:
# genotypes
# obj$X
# locus annotations
# obj$bim
# individual annotations
# obj$fam

# The following example is more awkward
# because package sample data has to be specified in this weird way:

# first get path to BED file
file <- system.file("extdata", "sample.bed", package = "genio", mustWork = TRUE)

# read genotypes and annotation tables
plink_data <- read_plink(file)
# genotypes
plink_data$X
# locus annotations
plink_data$bim
# individual annotations
plink_data$fam

# the same works without .bed extension
file <- sub("\.bed", ",", file) # remove extension
# it works!
plink_data <- read_plink(file)
```

---

**read_snp**

*Read Eigenstrat *.snp files*

**Description**

This function reads a standard Eigenstrat *.snp file into a tibble. It uses `readr::read_table()` to do it efficiently.
read_snp

Usage

read_snp(file, verbose = TRUE)

Arguments

file Input file (whatever is accepted by readr::read_table()). If file as given does not exist and is missing the expected *.snp extension, the function adds the .snp extension and uses that path if that file exists. Additionally, the .gz extension is added automatically if the file (after *.snp extension is added as needed) is still not found and did not already contain the .gz extension and adding it points to an existing file.

verbose If TRUE (default), function reports the path of the file being loaded (after auto-completing the extensions).

Value

A tibble with columns: id, chr, posg, pos, ref, alt

See Also

Eigenstrat SNP format reference: https://github.com/DReichLab/EIG/tree/master/CONVERTF

Examples

# to read "data.snp", run like this:
# snp <- read_snp("data")
# this also works
# snp <- read_snp("data.snp")

# The following example is more awkward
# because package sample data has to be specified in this weird way:

# read an existing Eigenstrat *.snp file
file <- system.file("extdata", 'sample.snp', package = "genio", mustWork = TRUE)
snp <- read_snp(file)

# can specify without extension
file <- sub('.snp$', '', file) # remove extension from this path on purpose
file # verify .snp is missing
snp <- read_snp(file) # load it anyway!
snp
require_files_grm

Description

This function checks that each of the GCTA binary GRM files (grm.bin, grm.N.bin, and grm.id extensions) are present, given the shared base file path, stopping with an informative message if any of the files is missing. This function aids troubleshooting, as various downstream external software report missing files differently and sometimes using confusing or obscure messages.

Usage

require_files_grm(file)

Arguments

file The shared file path (excluding extensions: grm.bin, grm.N.bin, or grm.id).

Value

Nothing

Examples

# to require all of "data.grm.bin", "data.grm.N.bin", and "data.grm.id", run like this:
# (stops if any of the three files is missing)
# require_files_grm("data")

# The following example is more awkward
# because package sample data has to be specified in this weird way:

# check that the samples we want exist
# start with bed file
file <- system.file("extdata", 'sample.grm.bin', package = "genio", mustWork = TRUE)
# remove extension
file <- sub(\\.grm\\.bin\$, '', file)
# since all sample.grm.(bin,N.bin,id) files exist, this will not stop with error messages:
require_files_grm(file)
require_files_phen  

**Description**

This function checks that the PHEN file is present, given the base file path, stopping with an informative message if the file is missing. This function aids troubleshooting, as various downstream external software report missing files differently and sometimes using confusing or obscure messages.

**Usage**

```
require_files_phen(file)
```

**Arguments**

- `file`  
The base file path (excluding phen extensions).

**Value**

Nothing

**Examples**

```r
# to require "data.phen", run like this:
# (stops if file is missing)
# require_files_phen("data")

# The following example is more awkward
# because package sample data has to be specified in this weird way:

# check that the samples we want exist
# get path to an existing phen file
file <- system.file("extdata", 'sample.phen', package = "genio", mustWork = TRUE)
# remove extension
file <- sub('\\.phen$', '', file)
# since sample.phen file exist, this will not stop with error messages:
require_files_phen(file)
```
require_files_plink

Description

This function checks that each of the Plink binary files (BED/BIM/FAM extensions) are present, given the shared base file path, stopping with an informative message if any of the files is missing. This function aids troubleshooting, as various downstream external software report missing files differently and sometimes using confusing or obscure messages.

Usage

require_files_plink(file)

Arguments

file The shared file path (excluding extensions bed, bim, fam).

Value

Nothing

Examples

# to require all of "data.bed", "data.bim", and "data.fam", run like this:
# (stops if any of the three files is missing)
# require_files_plink("data")

# The following example is more awkward
# because package sample data has to be specified in this weird way:

# check that the samples we want exist
# start with bed file
file <- system.file("extdata", 'sample.bed', package = "genio", mustWork = TRUE)
# remove extension
file <- sub('\..bed$', '', file)
# since all sample.(bed,bim,fam) files exist, this will not stop with error messages:
require_files_plink(file)
sex_to_char

Convert integer sex codes to character codes

Description

This function accepts the integer sex codes accepted by Plink and turns them into the character codes accepted by Eigenstrat. Only upper-case characters are returned. Cases outside the table below are mapped to `U` (unknown) with a warning. The correspondence is:

- 0: U (unknown)
- 1: M (male)
- 2: F (female)

Usage

sex_to_char(sex)

Arguments

sex 
Integer vector of sex codes

Value

The converted character vector of sex codes

See Also

sex_to_int()

Eigenstrat IND format reference: https://github.com/DReichLab/EIG/tree/master/CONVERTF

Plink FAM format reference: https://www.cog-genomics.org/plink/1.9/formats#fam

Examples

```r
# verify the mapping above
sex_int <- 0:2
sex_char <- c('U', 'M', 'F') # expected values
stopifnot(
  all(
    sex_to_char( sex_int ) == sex_char
  )
)
```
sex_to_int

Convert character sex codes to integer codes

Description
This function accepts the character sex codes accepted by Eigenstrat and turns them into the integer
codes accepted by Plink. Matching is case insensitive. Cases outside the table below are mapped to
0 (unknown) with a warning. The correspondence is:

• U: 0 (unknown)
• M: 1 (male)
• F: 2 (female)

Usage
sex_to_int(sex)

Arguments
sex Character vector of sex codes

Value
The converted numeric vector of sex codes

See Also
sex_to_char()

Eigenstrat IND format reference: https://github.com/DReichLab/EIG/tree/master/CONVERTF
Plink FAM format reference: https://www.cog-genomics.org/plink/1.9/formats#fam

Examples
# verify the mapping above
sex_char <- c('U', 'm', 'f') # mixed case works!
sex_int <- 0:2 # expected values
stopifnot(
  all(
    sex_to_int(sex_char) == sex_int
  )
)
tidy_kinship  
Create a tidy version of a kinship matrix

Description

A square symmetric kinship matrix is transformed into a tibble, with a row per unique element in the kinship matrix, and three columns: ID of row, ID of column, and the kinship value.

Usage

tidy_kinship(kinship, sort = TRUE)

Arguments

kinship  
The n-by-n symmetric kinship matrix

sort  
If TRUE (default), rows are sorted ascending by kinship value. Otherwise, order is moving along the upper triangle row-by-row

Value

A tibble with \( n \times (n + 1) / 2 \) rows (the upper triangle, including the diagonal), and 3 columns with names: id1, id2, kinship.

Examples

```r
# create a symmetric matrix
kinship <- matrix(
  c(
    0.5, 0.1, 0.0,
    0.1, 0.5, 0.2,
    0.0, 0.2, 0.6
  ),
  nrow = 3
)
# add names (best for tidy version)
colnames(kinship) <- paste0('pop', 1:3)ownames(kinship) <- paste0('pop', 1:3)
# this returns tidy version
kinship_tidy <- tidy_kinship( kinship )
# test colnames
stopifnot( colnames( kinship_tidy ) == c('id1', 'id2', 'kinship') )
# test row number
stopifnot( nrow( kinship_tidy ) == 6 )
# inspect it
kinship_tidy
```
write_bed

write_bed

Write a genotype matrix into Plink BED format

Description

This function accepts a standard R matrix containing genotypes (values in \(\{0, 1, 2, \text{NA}\}\)) and writes it into a Plink-formatted BED (binary) file. Each genotype per locus (m loci) and individual (n total) counts the number of alternative alleles or NA for missing data. No *.fam or *.bim files are created by this basic function.

Usage

write_bed(file, X, verbose = TRUE, append = FALSE)

Arguments

- **file**: Output file path. .bed extension may be omitted (will be added automatically if it is missing).
- **X**: The m-by-n genotype matrix. Row and column names, if present, are ignored.
- **verbose**: If TRUE (default), function reports the path of the file being written (after auto-completing the extension).
- **append**: If TRUE, appends variants onto the file. (Default is FALSE).

Details

Genotypes with values outside of \([0, 2]\) cause an error, in which case the partial output is deleted. However, beware that decimals get truncated internally, so values that truncate to 0, 1, or 2 will not raise errors. The BED format does not accept fractional dosages, so such data will not be written as expected.

Value

Nothing

See Also

write_plink() for writing a set of BED/BIM/FAM files.

Plink BED format reference: https://www.cog-genomics.org/plink/1.9/formats#bed

Examples

```R
# to write an existing matrix `X` into file "data.bed", run like this:
# write_bed("data", X)
# this also works
# write_bed("data.bed", X)

# The following example is more detailed but also more awkward
```
write_bim

Write Plink *.bim files

Description

This function writes a tibble with the right columns into a standard Plink *.bim file. It uses \texttt{readr::write_tsv()} to do it efficiently.

Usage

\begin{verbatim}
write_bim(file, tib, verbose = TRUE, append = FALSE)
\end{verbatim}

Arguments

- \texttt{file}  
  Output file (whatever is accepted by \texttt{readr::write_tsv()}). If file is missing the expected *.bim extension, the function adds it.

- \texttt{tib}  
  The tibble or data.frame to write. It must contain these columns: chr, id, posg, pos, ref, alt. Throws an error if any of these columns are missing. Additional columns are ignored. Columns are automatically reordered in output as expected in format.

- \texttt{verbose}  
  If TRUE (default), function reports the path of the file being written (after auto-completing the extension).

- \texttt{append}  
  If TRUE, appends rows onto the file. (Default is FALSE).

Value

The output tib invisibly (what \texttt{readr::write_tsv()} returns).

See Also

\texttt{write_plink()} for writing a set of BED/BIM/FAM files.
Plink BIM format reference: \url{https://www.cog-genomics.org/plink/1.9/formats#bim}
write_eigenvec

Examples

# to write an existing table 'bim' into file "data.bim", run like this:
# write_bim("data", bim)
# this also works
# write_bim("data.bim", bim)

# The following example is more detailed but also more awkward
# because (only for these examples) the package must create the file in a *temporary* location

# create a dummy tibble with the right columns
library(tibble)
tib <- tibble(
    chr = 1:3,
    id = 1:3,
    posg = 0,
    pos = 1:3,
    ref = 'A',
    alt = 'B'
)
# a dummy file
file_out <- tempfile("delete-me-example", fileext = ".bim") # will also work without extension
# write the table out in *.bim format (no header, columns in right order)
write_bim(file_out, tib)
# delete output when done
file.remove(file_out)

write_eigenvec

Write eigenvectors table into a Plink-format file

Description

This function writes eigenvectors in Plink 1 (same as GCTA) format (table with no header, with first two columns being fam and id), which is a subset of Plink 2 format (which optionally allows column names and does not require fam column). Main expected case is eigenvec passed as a numeric matrix and fam provided to complete first two missing columns. However, input eigenvec may also be a data.frame already containing the fam and id columns, and other reasonable intermediate cases are also handled. If both eigenvec and fam are provided and contain overlapping columns, those in eigenvec get overwritten with a warning.

Usage

write_eigenvec(file, eigenvec, fam = NULL, ext = "eigenvec", verbose = TRUE)

Arguments

file The output file name (possibly without extension)
write_eigenvec

eigenvec A matrix or tibble containing the eigenvectors to include in the file. Column names other than fam and id can be anything and are all treated as eigenvectors (not written to file).

fam An optional fam table, which is used to add the fam and id columns to eigenvec (which overwrite columns of the same name in eigenvec if present, after a warning is produced). Individuals in fam and eigenvec are assumed to be the same and in the same order.

ext Output file extension. Since the general "covariates" file format in GCTA and Plink are the same as this, this function may be used to write more general covariates files if desired, in which case users may wish to change this extension for clarity.

verbose If TRUE (default), function reports the path of the file being written (after auto-completing the extension).

Value

Invisibly, the final eigenvec data.frame or tibble written to file, starting with columns fam and id (merged from the fam input, if it was passed) followed by the rest of columns in the input eigenvec.

See Also

read_eigenvec() for reading an eigenvec file.

Plink 1 eigenvec format reference: [https://www.cog-genomics.org/plink/1.9/formats#eigenvec](https://www.cog-genomics.org/plink/1.9/formats#eigenvec)
Plink 2 eigenvec format reference: [https://www.cog-genomics.org/plink/2.0/formats#eigenvec](https://www.cog-genomics.org/plink/2.0/formats#eigenvec)
GCTA eigenvec format reference: [https://cnsgenomics.com/software/gcta/#PCA](https://cnsgenomics.com/software/gcta/#PCA)

Examples

# to write an existing matrix 'eigenvec' and optional 'fam' tibble into file "data.eigenvec",
# run like this:
# write_eigenvec("data", eigenvec, fam = fam)
# this also works
# write_eigenvec("data.eigenvec", eigenvec, fam = fam)

# The following example is more detailed but also more awkward
# because (only for these examples) the package must create the file in a *temporary* location

# create dummy eigenvectors matrix, in this case from a small identity matrix
# number of individuals
n <- 10
eigenvec <- eigen(diag(n))$vectors
# subset columns to use top 3 eigenvectors only
eigenvec <- eigenvec[, 1:3]
# dummy fam data
library(tibble)
fam <- tibble(fam = 1:n, id = 1:n)

# write this data to .eigenvec file
# output path without extension
write_fam

```rile <- tempfile('delete-me-example')
eigenvec_final <- write_eigenvec( file, eigenvec, fam = fam )
# inspect the tibble that was written to file (returned invisibly)
eigenvec_final

# remove temporary file (add extension before deletion)
file.remove( paste0( file, '.eigenvec' ) )
```

---

**write_fam**

*Write Plink *.fam files*

**Description**

This function writes a tibble with the right columns into a standard Plink *.fam* file. It uses `readr::write_tsv()` to do it efficiently.

**Usage**

```r
write_fam(file, tib, verbose = TRUE)
```

**Arguments**

- **file**: Output file (whatever is accepted by `readr::write_tsv()`). If file is missing the expected *.fam* extension, the function adds it.
- **tib**: The tibble or data.frame to write. It must contain these columns: `fam`, `id`, `pat`, `mat`, `sex`, `pheno`. Throws an error if any of these columns are missing. Additional columns are ignored. Columns are automatically reordered in output as expected in format.
- **verbose**: If `TRUE` (default), function reports the path of the file being written (after auto-completing the extension).

**Value**

The output tibble invisibly (what `readr::write_tsv()` returns).

**See Also**

- `write_plink()` for writing a set of BED/BIM/FAM files.
- Plink FAM format reference: [https://www.cog-genomics.org/plink/1.9/formats#fam](https://www.cog-genomics.org/plink/1.9/formats#fam)
Examples

```r
# to write an existing table `fam` into file "data.fam", run like this:
# write_fam("data", fam)
# this also works
# write_fam("data.fam", fam)

# The following example is more detailed but also more awkward
# because (only for these examples) the package must create the file in a *temporary* location

# create a dummy tibble with the right columns
library(tibble)
tib <- tibble(
  fam = 1:3,
  id = 1:3,
  pat = 0,
  mat = 0,
  sex = 1,
  pheno = 1
)

# a dummy file
file_out <- tempfile("delete-me-example", fileext = ".fam") # will also work without extension
# write the table out in *.fam format (no header, columns in right order)
write_fam(file_out, tib)
# delete output when done
file.remove(file_out)
```

---

**write_grm**

Write GCTA GRM binary files

**Description**

This function writes a GCTA Genetic Relatedness Matrix (GRM, i.e. kinship) set of files in their binary format, given a kinship matrix and, if available, the corresponding matrix of pair sample sizes (non-trivial under missingness) and individuals table.

**Usage**

```r
write_grm(name, kinship, M = NULL, fam = NULL, verbose = TRUE)
```

**Arguments**

- `name`: The base name of the output files. Files with that base and extensions `.grm.bin`, `.grm.N.bin`, and `.grm.id` may be created depending on the data provided.
- `kinship`: The symmetric n-times-n kinship matrix to write into file with extension `.grm.bin`.
- `M`: The optional symmetric n-times-n matrix of pair sample sizes to write into file with extension `.grm.N.bin`. 
write_grm

fam

The optional data.frame or tibble with individual annotations (columns with names `fam` and `id`, subset of columns of Plink FAM) to write into file with extension `.grm.id`. If `fam` is NULL but `kinship` has non-NULL column or row names, these are used as the second (`id`) value in the output table (the first (`fam`) column is set to the missing value in this case).

verbose

If TRUE (default), function reports the path of the files being written.

See Also

`read_grm()`

Examples

# to write existing data `kinship`, `M`, and `fam` into files "data.grm.bin" etc, run like this:
# write_grm("data", kinship, M = M, fam = fam )

# The following example is more detailed but also more awkward
# because (only for these examples) the package must create the file in a *temporary* location

# create dummy data to write
# kinship for 3 individuals
kinship <- matrix(
  c(0.6, 0.2, 0.0,
     0.2, 0.5, 0.1,
     0.0, 0.1, 0.5
  ),
  nrow = 3
)

# pair sample sizes matrix
M <- matrix( c(10, 9, 8,
               9, 9, 7,
               8, 7, 8
  ),
  nrow = 3
)

# individual annotations table
library(tibble)
fam <- tibble(  fam = 1:3,
                id = 1:3
  )

# dummy files to write and delete
name <- tempfile("delete-me-example") # no extension
# write the data now!
write_grm( name, kinship, M = M, fam = fam )
# delete outputs when done
delete_files_grm( name )
write_ind

Write Eigenstrat *.ind files

Description

This function writes a tibble with the right columns into a standard Eigenstrat *.ind file. It uses `readr::write_tsv()` to do it efficiently.

Usage

\[
\text{write_ind(file, tib, verbose = TRUE)}
\]

Arguments

- **file**: Output file (whatever is accepted by `readr::write_tsv()`). If file is missing the expected *.ind extension, the function adds it.
- **tib**: The tibble or data.frame to write. It must contain these columns: id, sex, label. Throws an error if any of these columns are missing. Additional columns are ignored. Columns are automatically reordered in output as expected in format.
- **verbose**: If `TRUE` (default), function reports the path of the file being written (after auto-completing the extension).

Value

The output tib invisibly (what `readr::write_tsv()` returns).

See Also

Eigenstrat IND format reference: [https://github.com/DReichLab/EIG/tree/master/converter](https://github.com/DReichLab/EIG/tree/master/converter)

Examples

\[
\begin{align*}
\text{# to write an existing table `ind` into file "data.ind", run like this:} \\
\text{# write_ind("data", ind)} \\
\text{# this also works} \\
\text{# write_ind("data.ind", ind)} \\
\text{# The following example is more detailed but also more awkward} \\
\text{# because (only for these examples) the package must create the file in a *temporary* location} \\
\text{# create a dummy tibble with the right columns} \\
\text{library(tibble)} \\
\text{tib <- tibble(} \\
\text{  \quad \text{\hspace{1cm} id = 1:3,}} \\
\text{  \quad \text{\hspace{1cm} sex = 1,}} \\
\text{  \quad \text{\hspace{1cm} label = 1}} \\
\text{)} \\
\text{# a dummy file}
\end{align*}
\]
write_matrix <- tempfile('delete-me-example', fileext = '.ind') # will also work without extension
# write the table out in *.ind format (no header, columns in right order)
write_ind(file_out, tib)
# delete output when done
file.remove(file_out)

write_matrix

Write a matrix to a file without row or column names

Description

The inverse function of read_matrix(), this writes what is intended to be a numeric matrix to a tab-delimited file without row or column names present. It uses readr::write_tsv() to do it efficiently. Intended for outputs such as those of admixture inference approaches.

Usage

write_matrix(file, x, ext = "txt", verbose = TRUE, append = FALSE)

Arguments

file Output file (whatever is accepted by readr::write_tsv()). If file is missing the expected extension (see below), the function adds it.

x The matrix to write. Unlike read_matrix(), this is not in fact required to be a matrix or be strictly numeric; anything that coerces to tibble or data.frame is acceptable.

ext The desired file extension. If NA, no extension is added. Works if file already contains desired extension.

verbose If TRUE (default), function reports the path of the file being written (after autocompleting the extension).

append If TRUE, appends rows onto the file. (Default is FALSE).

Value

The output x, coerced into data.frame, invisibly (what readr::write_tsv() returns).

See Also

read_matrix(), the inverse function.
Examples

# to write an existing matrix `x` into file "data.txt", run like this:
# write_matrix( "data", x )
# this also works
# write_matrix( "data.txt", x )

# The following example is more detailed but also more awkward
# because (only for these examples) the package must create the file in a *temporary* location

# create a dummy matrix with the right columns
x <- rbind( 1:3, (0:2)/10, -1:1 )
# a dummy file
file_out <- tempfile( 'delete-me-example', fileext = '.txt' ) # will also work without extension
# write the matrix without header
write_matrix( file_out, x )
# delete output when done
file.remove( file_out )

write_phen

Write *.phen files

Description

This function writes a tibble with the right columns into a standard *.phen file. It uses \texttt{readr::write_tsv()} to do it efficiently. GCTA and EMMAX use this format.

Usage

write_phen(file, tib, verbose = TRUE)

Arguments

- \texttt{file}  
  Output file (whatever is accepted by \texttt{readr::write_tsv()}). If file is missing the expected *.phen extension, the function adds it.

- \texttt{tib}  
  The tibble or data.frame to write. It must contain these columns: \texttt{fam, id, pheno}. Throws an error if any of these columns are missing. Additional columns are ignored. Columns are automatically reordered in output as expected in format.

- \texttt{verbose}  
  If \texttt{TRUE} (default), function reports the path of the file being written (after auto-completing the extension).

Value

The output \texttt{tib} invisibly (what \texttt{readr::write_tsv()} returns).

See Also

GCTA PHEN format reference: \url{https://cnsgenomics.com/software/gcta/#GREMLanalysis}
Examples

# to write an existing table `phen` into file "data.phen", run like this:
# write_phen("data", phen)
# this also works
# write_phen("data.phen", phen)

# The following example is more detailed but also more awkward
# because (only for these examples) the package must create the file in a *temporary* location

# create a dummy tibble with the right columns
library(tibble)
tib <- tibble(
  fam = 1:3,
  id = 1:3,
  pheno = 1
)
# a dummy file
file_out <- tempfile("delete-me-example", fileext = ".phen") # will also work without extension
# write the table out in *.phen format (no header, columns in right order)
write_phen(file_out, tib)
# delete output when done
file.remove(file_out)

write_plink

Write genotype and sample data into a Plink BED/BIM/FAM file set.

Description

This function writes a genotype matrix (X) and its associated locus (bim) and individual (fam) data tables into three Plink files in BED, BIM, and FAM formats, respectively. This function is a wrapper around the more basic functions write_bed(), write_bim(), write_fam(), but additionally tests that the data dimensions agree (or stops with an error). Also checks that the genotype row and column names agree with the bim and fam tables if they are all present. In addition, if bim = NULL or fam = NULL, these are auto-generated using make_bim() and make_fam(), which is useful behavior for simulated data. Lastly, the phenotype can be provided as a separate argument and incorporated automatically if fam = NULL (a common scenario for simulated genotypes and traits). Below suppose there are m loci and n individuals.

Usage

write_plink(
  file,
  X,
  bim = NULL,
  fam = NULL,
  pheno = NULL,
  verbose = TRUE,
append = FALSE}
)

Arguments

file Output file path, without extensions (each of .bed, .bim, .fam extensions will be added automatically as needed).

X The m-by-n genotype matrix.

bim The tibble or data.frame containing locus information. It must contain m rows and these columns: chr, id, posg, pos, ref, alt. If NULL (default), it will be quietly auto-generated.

fam The tibble or data.frame containing individual information. It must contain n rows and these columns: fam, id, pat, mat, sex, pheno. If NULL (default), it will be quietly auto-generated.

pheno The phenotype to write into the FAM file assuming fam = NULL. This must be a length-n vector. This will be ignored (with a warning) if fam is provided.

verbose If TRUE (default) function reports the paths of the files being written (after auto-completing the extensions).

append If TRUE, appends loci onto the BED and BIM files (default FALSE). In this mode, all individuals must be present in each write (only loci are appended); the FAM file is not overwritten if present, but is required at every write for internal validations. If the FAM file already exists, it is not checked to agree with the FAM table provided.

Value

Invisibly, a named list with items in this order: X (genotype matrix), bim (tibble), fam (tibble). This is most useful when either BIM or FAM tables were auto-generated.

See Also

write_bed(), write_bim(), write_fam(), make_bim(), make_fam().

Plink BED/BIM/FAM format reference: https://www.cog-genomics.org/plink/1.9/formats

Examples

# to write existing data `X`, `bim`, `fam` into files "data.bed", "data.bim", and "data.fam", # run like this:
# write_plink("data", X, bim = bim, fam = fam)

# The following example is more detailed but also more awkward # because (only for these examples) the package must create the file in a *temporary* location

# here is an example for a simulation

# create 10 random genotypes
X <- rbinom(10, 2, 0.5)
# replace 3 random genotypes with missing values

write_snp

```r
X[sample(10, 3)] <- NA
# turn into 5x2 matrix
X <- matrix(X, nrow = 5, ncol = 2)

# simulate a trait for two individuals
pheno <- rnorm(2)

# write this data to BED/BIM/FAM files
# output path without extension
file_out <- tempfile('delete-me-example')
# here all of the BIM and FAM columns except 'pheno' are autogenerated
write_plink(file_out, X, pheno = pheno)

# delete all three outputs when done
delete_files_plink(file_out)
```

---

**write_snp**  
*Write Eigenstrat *.snp files*

**Description**

This function writes a tibble with the right columns into a standard Eigenstrat *.snp file. It uses `readr::write_tsv()` to do it efficiently.

**Usage**

```r
write_snp(file, tib, verbose = TRUE)
```

**Arguments**

- `file`: Output file (whatever is accepted by `readr::write_tsv()`). If file is missing the expected *.snp extension, the function adds it.
- `tib`: The tibble or data.frame to write. It must contain these columns: id, chr, posg, pos, ref, alt. Throws an error if any of these columns are missing. Additional columns are ignored. Columns are automatically reordered in output as expected in format.
- `verbose`: If TRUE (default), function reports the path of the file being written (after auto-completing the extension).

**Value**

The output tib invisibly (what `readr::write_tsv()` returns).

**See Also**

Eigenstrat SNP format reference: https://github.com/DReichLab/EIG/tree/master/CONVERTF
Examples

# to write an existing table `snp` into file "data.snp", run like this:
# write_snp("data", snp)
# this also works
# write_snp("data.snp", snp)

# The following example is more detailed but also more awkward
# because (only for these examples) the package must create the file in a *temporary* location

# create a dummy tibble with the right columns
library(tibble)
tib <- tibble(
  id = 1:3,
  chr = 1:3,
  posg = 0,
  pos = 1:3,
  ref = 'A',
  alt = 'B'
)

# a dummy file
file_out <- tempfile('delete-me-example', fileext = '.snp')  # will also work without extension
# write the table out in *.snp format (no header, columns in right order)
write_snp(file_out, tib)
# delete output when done
file.remove(file_out)
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