

Package ‘MungeSumstats’

June 23, 2022

Type Package

Title Standardise summary statistics from GWAS

Version 1.4.5

Description The *MungeSumstats* package is designed to facilitate the standardisation of GWAS summary statistics. It reformats inputted summary statistics to include SNP, CHR, BP and can look up these values if any are missing. It also performs dozens of QC and filtering steps to ensure high data quality and minimise inter-study differences.

URL <https://github.com/neurogenomics/MungeSumstats>

BugReports <https://github.com/neurogenomics/MungeSumstats/issues>

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Depends R(>= 4.1)

Imports magrittr, data.table, utils, R.utils, dplyr, stats,
GenomicRanges, IRanges, GenomeInfoDb, BSgenome, Biostrings,
VariantAnnotation, googleAuthR, httr, jsonlite, methods,
parallel, rtracklayer, RCurl

biocViews SNP, WholeGenome, Genetics, ComparativeGenomics,
GenomeWideAssociation, GenomicVariation, Preprocessing

RoxygenNote 7.2.0

Encoding UTF-8

Roxygen list(markdown = TRUE)

Suggests SNPlocs.Hsapiens.dbSNP144.GRCh37,
SNPlocs.Hsapiens.dbSNP144.GRCh38,
BSgenome.Hsapiens.1000genomes.hs37d5,
BSgenome.Hsapiens.NCBI.GRCh38, BiocGenerics, S4Vectors,
rmarkdown, markdown, knitr, testthat (>= 3.0.0), UpSetR,
BiocStyle, covr, seqminer, Rsamtools, MatrixGenerics, badger

Config/testthat/edition 3

VignetteBuilder knitr

git_url <https://git.bioconductor.org/packages/MungeSumstats>

git_branch RELEASE_3_15

git_last_commit Oda13c2

git_last_commit_date 2022-06-08

Date/Publication 2022-06-23

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| | |
|-------------------|--|
| check_ldsc_format | <i>Ensures that parameters are compatible with LDSC format</i> |
|-------------------|--|

Description

Format summary statistics for direct input to Linkage Disequilibrium Score (LDSC) regression without the need to use their `munge_sumstats.py` script first.

Usage

```
check_ldsc_format(
    sumstats_dt,
    ldsc_format,
    convert_n_int,
    allele_flip_check,
    compute_z,
    compute_n
)
```

Arguments

| | |
|--------------------------------|--|
| <code>sumstats_dt</code> | data table obj of the summary statistics file for the GWAS. |
| <code>ldsc_format</code> | Binary Ensure that output format meets all requirements to be fed directly into LDSC without the need for additional munging. Default is FALSE |
| <code>convert_n_int</code> | Binary, if N (the number of samples) is not an integer, should this be rounded? Default is TRUE. |
| <code>allele_flip_check</code> | Binary Should the allele columns be checked against reference genome to infer if flipping is necessary. Default is TRUE. |
| <code>compute_z</code> | Whether to compute Z-score column from P. Default is FALSE. Note that imputing the Z-score for every SNP will not correct be perfectly correct and may result in a loss of power. This should only be done as a last resort. |
| <code>compute_n</code> | Whether to impute N. Default of 0 won't impute, any other integer will be imputed as the N (sample size) for every SNP in the dataset. Note that imputing the sample size for every SNP is not correct and should only be done as a last resort. N can also be inputted with "ldsc", "sum", "giant" or "metal" by passing one of these for this field or a vector of multiple. Sum and an integer value creates an N column in the output whereas giant, metal or ldsc create an Neff or effective sample size. If multiples are passed, the formula used to derive it will be indicated. |

Details

[LDSC documentation.](#)

Value

Formatted summary statistics

Source

[LDSC GitHub](#)

| | |
|---------------|---|
| compute_nsize | <i>Check for N column if not present and user wants, impute N based on user's sample size. NOTE this will be the same value for each SNP which is not necessarily correct and may cause issues down the line. N can also be inputted with "ldsc", "sum", "giant" or "metal" by passing one or multiple of these.</i> |
|---------------|---|

Description

Check for N column if not present and user wants, impute N based on user's sample size. **NOTE** this will be the same value for each SNP which is not necessarily correct and may cause issues down the line. N can also be inputted with "ldsc", "sum", "giant" or "metal" by passing one or multiple of these.

Usage

```
compute_nsize(
  sumstats_dt,
  imputation_ind = FALSE,
  compute_n = c("ldsc", "giant", "metal", "sum"),
  standardise_headers = FALSE,
  force_new = FALSE,
  return_list = TRUE
)
```

Arguments

| | |
|----------------|--|
| sumstats_dt | data table obj of the summary statistics file for the GWAS. |
| imputation_ind | Binary Should a column be added for each imputation step to show what SNPs have imputed values for differing fields. This includes a field denoting SNP allele flipping (flipped). Note these columns will be in the formatted summary statistics returned. Default is FALSE. |
| compute_n | How to compute per-SNP sample size (new column "N"). <ul style="list-style-type: none"> • 0: N will not be computed. • >0: If any number >0 is provided, that value will be set as N for every row. Note: Computing N this way is incorrect and should be avoided if at all possible. • "sum": N will be computed as: cases (N_CAS) + controls (N_CON), so long as both columns are present. |

- "ldsc": N will be computed as effective sample size: $N_{\text{eff}} = (N_{\text{CAS}} + N_{\text{CON}}) * (N_{\text{CAS}} / (N_{\text{CAS}} + N_{\text{CON}})) / \text{mean}((N_{\text{CAS}} / (N_{\text{CAS}} + N_{\text{CON}})) * (N_{\text{CAS}} + N_{\text{CON}}))$
- "giant": N will be computed as effective sample size: $N_{\text{eff}} = 2 / (1/N_{\text{CAS}} + 1/N_{\text{CON}})$.
- "metal": N will be computed as effective sample size: $N_{\text{eff}} = 4 / (1/N_{\text{CAS}} + 1/N_{\text{CON}})$.

standardise_headers

Standardise headers first.

force_new

If "Neff" (or "N") already exists in sumstats_dt, replace it with the recomputed version.

return_list

Return the sumstats_dt within a named list (default: TRUE).

Value

list("sumstats_dt"=sumstats_dt)

Examples

```
sumstats_dt <- MungeSumstats::formatted_example()
sumstats_dt2 <- MungeSumstats::compute_nsize(sumstats_dt=sumstats_dt,
                                             compute_n=10000)
```

download_vcf

Download VCF file and its index file from Open GWAS

Description

Ideally, we would use [gwasvcf](#) instead but it hasn't been made available on CRAN or Bioconductor yet, so we can't include it as a dep.

Usage

```
download_vcf(
  vcf_url,
  vcf_dir = tempdir(),
  vcf_download = TRUE,
  download_method = "download.file",
  force_new = FALSE,
  quiet = TRUE,
  nThread = 1
)
```

Arguments

| | |
|-----------------|---|
| vcf_url | Remote URL to VCF file. |
| vcf_dir | Where to download the original VCF from Open GWAS. <i>WARNING</i> : This is set to <code>tempdir()</code> by default. This means the raw (pre-formatted) VCFs be deleted upon ending the R session. Change this to keep the raw VCF file on disk (e.g. <code>vcf_dir="/raw_vcf"</code>). |
| vcf_download | Download the original VCF from Open GWAS. |
| download_method | "axel" (multi-threaded) or "download.file" (single-threaded) . |
| force_new | Overwrite a previously downloaded VCF with the same path name. |
| quiet | Run quietly. |
| nThread | Number of threads to parallelize over. |

Value

List containing the paths to the downloaded VCF and its index file.

Examples

```
#only run the examples if user has internet access:
if(try(is.character(getURL("www.google.com")))==TRUE){
vcf_url <- "https://gwas.mrcieu.ac.uk/files/ieu-a-298/ieu-a-298.vcf.gz"
out_paths <- download_vcf(vcf_url = vcf_url)
}
```

find_sumstats

Search Open GWAS for datasets matching criteria

Description

For each argument, searches for any datasets matching a case-insensitive substring search in the respective metadata column. Users can supply a single character string or a list/vector of character strings.

Usage

```
find_sumstats(
  ids = NULL,
  traits = NULL,
  years = NULL,
  consortia = NULL,
  authors = NULL,
  populations = NULL,
  categories = NULL,
  subcategories = NULL,
```

```

    builds = NULL,
    pmids = NULL,
    min_sample_size = NULL,
    min_ncase = NULL,
    min_ncontrol = NULL,
    min_nsnp = NULL,
    include_NAs = FALSE,
    access_token = check_access_token()
  )

```

Arguments

| | |
|-----------------|--|
| ids | List of Open GWAS study IDs (e.g. c("prot-a-664", "ieu-b-4760")). |
| traits | List of traits (e.g. c("parkinson", "Alzheimer")). |
| years | List of years (e.g. seq(2015,2021) or c(2010, 2012, 2021)). |
| consortia | List of consortia (e.g. c("MRC-IEU", "Neale Lab")). |
| authors | List of authors (e.g. c("Elsworth", "Kunkle", "Neale")). |
| populations | List of populations (e.g. c("European", "Asian")). |
| categories | List of categories (e.g. c("Binary", "Continuous", "Disease", "Risk factor")). |
| subcategories | List of categories (e.g. c("neurological", "Immune", "cardio")). |
| builds | List of genome builds (e.g. c("hg19", "grch37")). |
| pmids | List of PubMed ID (exact matches only) (e.g. c(29875488, 30305740, 28240269)). |
| min_sample_size | Minimum total number of study participants (e.g. 5000). |
| min_ncase | Minimum number of case participants (e.g. 1000). |
| min_ncontrol | Minimum number of control participants (e.g. 1000). |
| min_nsnp | Minimum number of SNPs (e.g. 200000). |
| include_NAs | Include datasets with missing metadata for size criteria (i.e. min_sample_size, min_ncase, or min_ncontrol). |
| access_token | Google OAuth2 access token. Used to authenticate level of access to data |

Details

By default, returns metadata for all studies currently in Open GWAS database.

Value

(Filtered) GWAS metadata table.

Examples

```

# Only run the examples if user has internet access:
if(try(is.character(getURL("www.google.com")))==TRUE){
### By ID
metagwas <- find_sumstats(ids = c(

```

```
    "ieu-b-4760",
    "prot-a-1725",
    "prot-a-664"
  ))
  ### By ID and sample size
  metagwas <- find_sumstats(
    ids = c("ieu-b-4760", "prot-a-1725", "prot-a-664"),
    min_sample_size = 5000
  )
  ### By criteria
  metagwas <- find_sumstats(
    traits = c("alzheimer", "parkinson"),
    years = seq(2015, 2021)
  )
}
```

| | |
|-------------------|--------------------------|
| formatted_example | <i>Formatted example</i> |
|-------------------|--------------------------|

Description

Returns an example of summary stats that have had their column names already standardised with [standardise_header](#).

Usage

```
formatted_example(
  path = system.file("extdata", "eduAttain0kbay.txt", package = "MungeSumstats"),
  formatted = TRUE,
  sorted = TRUE
)
```

Arguments

| | |
|-----------|--|
| path | Path to raw example file. Default to built-in dataset. |
| formatted | Whether the column names should be formatted (default:TRUE). |
| sorted | Whether the rows should be sorted by genomic coordinates (default:TRUE). |

Value

sumstats_dt

Examples

```
sumstats_dt <- MungeSumstats::formatted_example()
```

| | |
|-----------------|--|
| format_sumstats | <i>Check that summary statistics from GWAS are in a homogeneous format</i> |
|-----------------|--|

Description

Check that summary statistics from GWAS are in a homogeneous format

Usage

```
format_sumstats(  
  path,  
  ref_genome = NULL,  
  convert_ref_genome = NULL,  
  convert_small_p = TRUE,  
  convert_large_p = TRUE,  
  convert_neg_p = TRUE,  
  compute_z = FALSE,  
  force_new_z = FALSE,  
  compute_n = 0L,  
  convert_n_int = TRUE,  
  impute_beta = FALSE,  
  impute_se = FALSE,  
  analysis_trait = NULL,  
  INFO_filter = 0.9,  
  FRQ_filter = 0,  
  pos_se = TRUE,  
  effect_columns_nonzero = FALSE,  
  N_std = 5,  
  N_dropNA = TRUE,  
  rmv_chr = c("X", "Y", "MT"),  
  rmv_chrPrefix = TRUE,  
  on_ref_genome = TRUE,  
  strand_ambig_filter = FALSE,  
  allele_flip_check = TRUE,  
  allele_flip_drop = TRUE,  
  allele_flip_z = TRUE,  
  allele_flip_frq = TRUE,  
  bi_allelic_filter = TRUE,  
  snp_ids_are_rs_ids = TRUE,  
  remove_multi_rs_snp = FALSE,  
  frq_is_maf = TRUE,  
  indels = TRUE,  
  sort_coordinates = TRUE,  
  nThread = 1,  
  save_path = tempfile(fileext = ".tsv.gz"),  
  write_vcf = FALSE,  
)
```

```

tabix_index = FALSE,
return_data = FALSE,
return_format = "data.table",
ldsc_format = FALSE,
log_folder_ind = FALSE,
log_mungesumstats_msgs = FALSE,
log_folder = tempdir(),
imputation_ind = FALSE,
force_new = FALSE,
mapping_file = sumstatsColHeaders
)

```

Arguments

| | |
|--------------------|--|
| path | Filepath for the summary statistics file to be formatted. A dataframe or datatable of the summary statistics file can also be passed directly to MungeSumstats using the path parameter. |
| ref_genome | name of the reference genome used for the GWAS ("GRCh37" or "GRCh38"). Argument is case-insensitive. Default is NULL which infers the reference genome from the data. |
| convert_ref_genome | name of the reference genome to convert to ("GRCh37" or "GRCh38"). This will only occur if the current genome build does not match. Default is not to convert the genome build (NULL). |
| convert_small_p | Binary, should non-negative p-values $\leq 5e-324$ be converted to 0? Small p-values pass the R limit and can cause errors with LDSC/MAGMA and should be converted. Default is TRUE. |
| convert_large_p | Binary, should p-values >1 be converted to 1? P-values >1 should not be possible and can cause errors with LDSC/MAGMA and should be converted. Default is TRUE. |
| convert_neg_p | Binary, should p-values <0 be converted to 0? Negative p-values should not be possible and can cause errors with LDSC/MAGMA and should be converted. Default is TRUE. |
| compute_z | Whether to compute Z-score column from P. Default is FALSE. Note that imputing the Z-score for every SNP will not correct be perfectly correct and may result in a loss of power. This should only be done as a last resort. |
| force_new_z | When a "Z" column already exists, it will be used by default. To override and compute a new Z-score column from P set force_new_z=TRUE. |
| compute_n | Whether to impute N. Default of 0 won't impute, any other integer will be imputed as the N (sample size) for every SNP in the dataset. Note that imputing the sample size for every SNP is not correct and should only be done as a last resort. N can also be inputted with "ldsc", "sum", "giant" or "metal" by passing one of these for this field or a vector of multiple. Sum and an integer value creates an N column in the output whereas giant, metal or ldsc create an Neff or effective sample size. If multiples are passed, the formula used to derive it will be indicated. |

| | |
|------------------------|---|
| convert_n_int | Binary, if N (the number of samples) is not an integer, should this be rounded? Default is TRUE. |
| impute_beta | Binary, whether BETA should be imputed using other effect data if it isn't present in the sumstats. Note that this imputation is an approximation so could have an effect on downstream analysis. Use with caution. The different methods MungeSumstats will try and impute beta (in this order or priority) are: 1. log(OR) 2. Z x SE Default value is FALSE. |
| impute_se | Binary, whether the standard error should be imputed using other effect data if it isn't present in the sumstats. Note that this imputation is an approximation so could have an effect on downstream analysis. Use with caution. The different methods MungeSumstats will try and impute se (in this order or priority) are: 1. BETA / Z 2. abs(BETA/ qnorm(P/2)) Default is FALSE. |
| analysis_trait | If multiple traits were studied, name of the trait for analysis from the GWAS. Default is NULL. |
| INFO_filter | numeric The minimum value permissible of the imputation information score (if present in sumstats file). Default 0.9. |
| FRQ_filter | numeric The minimum value permissible of the frequency(FRQ) of the SNP (i.e. Allele Frequency (AF)) (if present in sumstats file). By default no filtering is done, i.e. value of 0. |
| pos_se | Binary Should the standard Error (SE) column be checked to ensure it is greater than 0? Those that are, are removed (if present in sumstats file). Default TRUE. |
| effect_columns_nonzero | Binary should the effect columns in the data BETA,OR (odds ratio),LOG_ODDS,SIGNED_SUMSTAT be checked to ensure no SNP=0. Those that do are removed(if present in sumstats file). Default FALSE. |
| N_std | numeric The number of standard deviations above the mean a SNP's N is needed to be removed. Default is 5. |
| N_dropNA | Drop rows where N is missing.Default is TRUE. |
| rmv_chr | vector or character The chromosomes on which the SNPs should be removed. Use NULL if no filtering necessary. Default is X, Y and mitochondrial. |
| rmv_chrPrefix | Remove "chr" or "CHR" from chromosome names. Default is TRUE. |
| on_ref_genome | Binary Should a check take place that all SNPs are on the reference genome by SNP ID. Default is TRUE. |
| strand_ambig_filter | Binary Should SNPs with strand-ambiguous alleles be removed. Default is FALSE. |
| allele_flip_check | Binary Should the allele columns be checked against reference genome to infer if flipping is necessary. Default is TRUE. |
| allele_flip_drop | Binary Should the SNPs for which neither their A1 or A2 base pair values match a reference genome be dropped. Default is TRUE. |
| allele_flip_z | Binary should the Z-score be flipped along with effect and FRQ columns like Beta? It is assumed to be calculated off the effect size not the P-value and so will be flipped i.e. default TRUE. |

| | |
|------------------------|---|
| allele_flip_frq | Binary should the frequency (FRQ) column be flipped along with effect and z-score columns like Beta? Default TRUE. |
| bi_allelic_filter | Binary Should non-biallelic SNPs be removed. Default is TRUE. |
| snp_ids_are_rs_ids | Binary Should the supplied SNP ID's be assumed to be RSIDs. If not, imputation using the SNP ID for other columns like base-pair position or chromosome will not be possible. If set to FALSE, the SNP RS ID will be imputed from the reference genome if possible. Default is TRUE. |
| remove_multi_rs_snp | Binary Sometimes summary statistics can have multiple RSIDs on one row (i.e. related to one SNP), for example "rs5772025_rs397784053". This can cause an error so by default, the first RS ID will be kept and the rest removed e.g. "rs5772025". If you want to just remove these SNPs entirely, set it to TRUE. Default is FALSE. |
| frq_is_maf | Conventionally the FRQ column is intended to show the minor/effect allele frequency (MAF) but sometimes the major allele frequency can be inferred as the FRQ column. This logical variable indicates that the FRQ column should be renamed to MAJOR_ALLELE_FRQ if the frequency values appear to relate to the major allele i.e. >0.5. By default this mapping won't occur i.e. is TRUE. |
| indels | Binary does your Sumstats file contain Indels? These don't exist in our reference file so they will be excluded from checks if this value is TRUE. Default is TRUE. |
| sort_coordinates | Whether to sort by coordinates of resulting sumstats |
| nThread | Number of threads to use for parallel processes. |
| save_path | File path to save formatted data. Defaults to <code>tempfile(fileext=".tsv.gz")</code> . |
| write_vcf | Whether to write as VCF (TRUE) or tabular file (FALSE). |
| tabix_index | Index the formatted summary statistics with <code>tabix</code> for fast querying. |
| return_data | Return data <code>.table</code> , <code>GRanges</code> or <code>VRanges</code> directly to user. Otherwise, return the path to the save data. Default is FALSE. |
| return_format | If <code>return_data</code> is TRUE. Object type to be returned (" <code>data.table</code> ", " <code>vranges</code> ", " <code>granges</code> "). |
| ldsc_format | Binary Ensure that output format meets all requirements to be fed directly into LDSC without the need for additional munging. Default is FALSE |
| log_folder_ind | Binary Should log files be stored containing all filtered out SNPs (separate file per filter). The data is outputted in the same format specified for the resulting sumstats file. The only exception to this rule is if output is vcf, then log file saved as <code>.tsv.gz</code> . Default is FALSE. |
| log_mungesumstats_msgs | Binary Should a log be stored containing all messages and errors printed by MungeSumstats in a run. Default is FALSE |
| log_folder | Filepath to the directory for the log files and the log of MungeSumstats messages to be stored. Default is a temporary directory. |

| | |
|----------------|--|
| imputation_ind | Binary Should a column be added for each imputation step to show what SNPs have imputed values for differing fields. This includes a field denoting SNP allele flipping (flipped). On the flipped value, this denoted whether the alleles were switched based on MungeSumstats initial choice of A1, A2 from the input column headers and thus may not align with what the creator intended. Note these columns will be in the formatted summary statistics returned. Default is FALSE. |
| force_new | If a formatted file of the same names as save_path exists, formatting will be skipped and this file will be imported instead (default). Set force_new=TRUE to override this. |
| mapping_file | MungeSumstats has a pre-defined column-name mapping file which should cover the most common column headers and their interpretations. However, if a column header that is in your file is missing of the mapping we give is incorrect you can supply your own mapping file. Must be a 2 column dataframe with column names "Uncorrected" and "Corrected". See data(sumstatsColHeaders) for default mapping and necessary format. |

Value

The address for the modified sumstats file or the actual data dependent on user choice. Also, if log files wanted by the user, the return in both above instances are a list.

Examples

```
# Pass path to Educational Attainment Okbay sumstat file to a temp directory

eduAttainOkbayPth <- system.file("extdata", "eduAttainOkbay.txt",
  package = "MungeSumstats"
)

## Call uses reference genome as default with more than 2GB of memory,
## which is more than what 32-bit Windows can handle so remove certain checks

is_32bit_windows <-
  .Platform$OS.type == "windows" && .Platform$r_arch == "i386"
if (!is_32bit_windows) {
  reformatted <- format_sumstats(
    path = eduAttainOkbayPth,
    ref_genome = "GRCh37"
  )
} else {
  reformatted <- format_sumstats(
    path = eduAttainOkbayPth,
    ref_genome = "GRCh37",
    on_ref_genome = FALSE,
    strand_ambig_filter = FALSE,
    bi_allelic_filter = FALSE,
    allele_flip_check = FALSE
  )
}
# returned location has the updated summary statistics file
```

get_genome_builds *Infer genome builds*

Description

Infers the genome build of summary statistics files (GRCh37 or GRCh38) from the data. Uses SNP (RSID) & CHR & BP to get genome build.

Usage

```
get_genome_builds(
  sumstats_list,
  header_only = TRUE,
  sampled_snps = 10000,
  names_from_paths = FALSE,
  nThread = 1
)
```

Arguments

| | |
|------------------|---|
| sumstats_list | A named list of paths to summary statistics, or a named list of data.table objects. |
| header_only | Instead of reading in the entire sumstats file, only read in the first N rows where N=sampled_snps. This should help speed up cases where you have to read in sumstats from disk each time. |
| sampled_snps | Downsample the number of SNPs used when inferring genome build to save time. |
| names_from_paths | Infer the name of each item in sumstats_list from its respective file path. Only works if sumstats_list is a list of paths. |
| nThread | Number of threads to use for parallel processes. |

Details

Iterative version of get_genome_build.

Value

ref_genome the genome build of the data

Examples

```
# Pass path to Educational Attainment Okbay sumstat file to a temp directory
eduAttainOkbayPth <- system.file("extdata", "eduAttainOkbay.txt",
  package = "MungeSumstats"
)
```

```
sumstats_list <- list(ss1 = eduAttainOkbayPth, ss2 = eduAttainOkbayPth)

## Call uses reference genome as default with more than 2GB of memory,
## which is more than what 32-bit Windows can handle so remove certain checks
is_32bit_windows <-
  .Platform$OS.type == "windows" && .Platform$r_arch == "i386"
if (!is_32bit_windows) {

  #multiple sumstats can be passed at once to get all their genome builds:
  #ref_genomes <- get_genome_builds(sumstats_list = sumstats_list)
  #just passing first here for speed
  sumstats_list_quick <- list(ss1 = eduAttainOkbayPth)
  ref_genomes <- get_genome_builds(sumstats_list = sumstats_list_quick)
}
```

hg19ToHg38

UCSC Chain file hg19 to hg38

Description

UCSC Chain file hg19 to hg38, .chain.gz file, downloaded from <https://hgdownload.cse.ucsc.edu/goldenpath/hg19/liftOver/> on 09/10/21

Format

gunzipped chain file

Details

UCSC Chain file hg19 to hg38, .chain.gz file, downloaded on 09/10/21 To be used as a back up if the download from UCSC fails.

hg19ToHg38.over.chain.gz

NA

Source

The chain file was downloaded from <https://hgdownload.cse.ucsc.edu/goldenpath/hg19/liftOver/>
`utils::download.file('ftp://hgdownload.cse.ucsc.edu/goldenPath/hg19/liftOver/hg19ToHg38.over.chain.`

hg38ToHg19

UCSC Chain file hg38 to hg19

Description

UCSC Chain file hg38 to hg19, .chain.gz file, downloaded from <https://hgdownload.cse.ucsc.edu/goldenpath/hg19/liftOver/> on 09/10/21

Format

gunzipped chain file

Details

UCSC Chain file hg38 to hg19, .chain.gz file, downloaded on 09/10/21 To be used as a back up if the download from UCSC fails.

hg38ToHg19.over.chain.gz

NA

Source

The chain file was downloaded from <https://hgdownload.cse.ucsc.edu/goldenpath/hg38/liftOver/> `utils::download.file('ftp://hgdownload.cse.ucsc.edu/goldenPath/hg38/liftOver/hg38ToHg19.over.chain.gz')`

ieu-a-298

Local ieu-a-298 file from IEU Open GWAS

Description

Local ieu-a-298 file from IEU Open GWAS, downloaded on 09/10/21.

Format

gunzipped tsv file

Details

Local ieu-a-298 file from IEU Open GWAS, downloaded on 09/10/21. This is done in case the download in the package vignette fails.

ieu-a-298.tsv.gz

NA

Source

The file was downloaded with: `MungeSumstats::import_sumstats(ids = "ieu-a-298", ref_genome = "GRCH37")`

| | |
|-----------------|---|
| import_sumstats | <i>Import full genome-wide GWAS summary statistics from Open GWAS</i> |
|-----------------|---|

Description

Requires internet access to run.

Usage

```
import_sumstats(
  ids,
  vcf_dir = tempdir(),
  vcf_download = TRUE,
  save_dir = tempdir(),
  write_vcf = FALSE,
  download_method = "download.file",
  quiet = TRUE,
  force_new = FALSE,
  force_new_vcf = FALSE,
  nThread = 1,
  parallel_across_ids = FALSE,
  ...
)
```

Arguments

| | |
|------------------------------|--|
| <code>ids</code> | List of Open GWAS study IDs (e.g. <code>c("prot-a-664", "ieu-b-4760")</code>). |
| <code>vcf_dir</code> | Where to download the original VCF from Open GWAS. <i>WARNING:</i> This is set to <code>tempdir()</code> by default. This means the raw (pre-formatted) VCFs be deleted upon ending the R session. Change this to keep the raw VCF file on disk (e.g. <code>vcf_dir="/raw_vcf"</code>). |
| <code>vcf_download</code> | Download the original VCF from Open GWAS. |
| <code>save_dir</code> | Directory to save formatted summary statistics in. |
| <code>write_vcf</code> | Whether to write as VCF (TRUE) or tabular file (FALSE). |
| <code>download_method</code> | "axel" (multi-threaded) or "download.file" (single-threaded) . |
| <code>quiet</code> | Run quietly. |
| <code>force_new</code> | If a formatted file of the same names as <code>save_path</code> exists, formatting will be skipped and this file will be imported instead (default). Set <code>force_new=TRUE</code> to override this. |

force_new_vcf Overwrite a previously downloaded VCF with the same path name.
 nThread Number of threads to use for parallel processes.
 parallel_across_ids If parallel_across_ids=TRUE and nThread>1, then each ID in ids will be processed in parallel.
 ... Additional arguments passed to [format_sumstats](#).

Value

Either a named list of data objects or paths, depending on the arguments passed to `format_sumstats`.

Examples

```
#only run the examples if user has internet access:
if(try(is.character(getURL("www.google.com")))==TRUE){
### Search by criteria
metagwas <- find_sumstats(
  traits = c("parkinson", "alzheimer"),
  min_sample_size = 5000
)
### Only use a subset for testing purposes
ids <- (dplyr::arrange(metagwas, nsnp))$id

### Default usage
## You can supply \code{import_sumstats()}
## with a list of as many OpenGWAS IDs as you want,
## but we'll just give one to save time.

## Call uses reference genome as default with more than 2GB of memory,
## which is more than what 32-bit Windows can handle so remove certain checks
## commented out down to runtime
# datasets <- import_sumstats(ids = ids[1])
}
```

index_tabular

Tabix-index file: table

Description

Convert summary stats file to tabix format.

Usage

```
index_tabular(
  path,
  chrom_col = "CHR",
  start_col = "BP",
  end_col = start_col,
  verbose = TRUE
)
```

Arguments

path Path to GWAS summary statistics file.
 chrom_col column for chromosome
 start_col column for start position
 end_col column for end position (is the same as start for snps)
 verbose Print messages.

Value

Path to tabix-indexed tabular file

Source

Borrowed function from [echotabix](#).

See Also

Other tabix: [index_vcf\(\)](#)

Examples

```

sumstats_dt <- MungeSumstats::formatted_example()
sumstats_dt <- MungeSumstats::sort_coords(sumstats_dt = sumstats_dt)
path <- tempfile(fileext = ".tsv")
MungeSumstats::write_sumstats(sumstats_dt = sumstats_dt, save_path = path)

indexed_file <- MungeSumstats::index_tabular(path = path)

```

is_tabix

Is tabix

Description

Is a file bgz-zip compressed and tabix-indexed.

Usage

```
is_tabix(path)
```

Arguments

path Path to file.

Value

logical: whether the file is tabix-indexed or not.

| | |
|----------|------------------------------|
| liftover | <i>Genome build liftover</i> |
|----------|------------------------------|

Description

Transfer genomic coordinates from one genome build to another.

Usage

```
liftover(
  sumstats_dt,
  convert_ref_genome,
  ref_genome,
  imputation_ind = TRUE,
  chrom_col = "CHR",
  start_col = "BP",
  end_col = start_col,
  as_granges = FALSE,
  style = "NCBI",
  verbose = TRUE
)
```

Arguments

| | |
|---------------------------------|--|
| <code>sumstats_dt</code> | data table obj of the summary statistics file for the GWAS. |
| <code>convert_ref_genome</code> | name of the reference genome to convert to ("GRCh37" or "GRCh38"). This will only occur if the current genome build does not match. Default is not to convert the genome build (NULL). |
| <code>ref_genome</code> | name of the reference genome used for the GWAS ("GRCh37" or "GRCh38"). Argument is case-insensitive. Default is NULL which infers the reference genome from the data. |
| <code>imputation_ind</code> | Binary Should a column be added for each imputation step to show what SNPs have imputed values for differing fields. This includes a field denoting SNP allele flipping (flipped). On the flipped value, this denoted whether the alleles were switched based on MungeSumstats initial choice of A1, A2 from the input column headers and thus may not align with what the creator intended. Note these columns will be in the formatted summary statistics returned. Default is FALSE. |
| <code>chrom_col</code> | Name of the chromosome column in <code>sumstats_dt</code> (e.g. "CHR"). |
| <code>start_col</code> | Name of the starting genomic position column in <code>sumstats_dt</code> (e.g. "POS", "start"). |
| <code>end_col</code> | Name of the ending genomic position column in <code>sumstats_dt</code> (e.g. "POS", "end"). Can be the same as <code>start_col</code> when <code>sumstats_dt</code> only contains SNPs that span 1 base pair (bp) each. |
| <code>as_granges</code> | Return results as GRanges instead of a data.table (default: FALSE). |

| | |
|---------|--|
| style | Style to return GRanges object in (e.g. "NCBI" = 4; "UCSC" = "chr4"); (default: "NCBI"). |
| verbose | Print messages. |

Value

Lifted summary stats in `data.table` or [GRanges](#) format.

Source

[liftOver](#)
[UCSC chain files](#)

Examples

```
sumstats_dt <- MungeSumstats::formatted_example()

sumstats_dt_hg38 <- liftOver(sumstats_dt=sumstats_dt,
                             ref_genome = "hg19",
                             convert_ref_genome="hg38")
```

| | |
|---------------|---------------------------------------|
| list_sumstats | <i>List munged summary statistics</i> |
|---------------|---------------------------------------|

Description

Searches for and lists local GWAS summary statistics files munged by [format_sumstats](#) or [import_sumstats](#).

Usage

```
list_sumstats(
  save_dir = getwd(),
  pattern = "*.tsv.gz$",
  ids_from_file = TRUE,
  verbose = TRUE
)
```

Arguments

| | |
|---------------|--|
| save_dir | Top-level directory to recursively search for summary statistics files within. |
| pattern | Regex pattern to search for files with. |
| ids_from_file | Try to extract dataset IDs from file names. If FALSE, will infer IDs from the directory names instead. |
| verbose | Print messages. |

Value

Named vector of summary stats paths.

Examples

```
save_dir <- system.file("extdata", package = "MungeSumstats")
munged_files <- MungeSumstats::list_sumstats(save_dir = save_dir)
```

load_ref_genome_data *Load the reference genome data for SNPs of interest*

Description

Load the reference genome data for SNPs of interest

Usage

```
load_ref_genome_data(snps, ref_genome, msg = NULL)
```

Arguments

| | |
|------------|---|
| snps | Character vector SNPs by rs_id from sumstats file of interest. |
| ref_genome | Name of the reference genome used for the GWAS (GRCh37 or GRCh38) |
| msg | Optional name of the column missing from the dataset in question. Default is NULL |

Value

data table of snpsById, filtered to SNPs of interest.

load_snp_loc_data *Loads the SNP locations and alleles for Homo sapiens extracted from NCBI dbSNP Build 144. Reference genome version is dependent on user input.*

Description

Loads the SNP locations and alleles for Homo sapiens extracted from NCBI dbSNP Build 144. Reference genome version is dependent on user input.

Usage

```
load_snp_loc_data(ref_genome, msg = NULL)
```

Arguments

ref_genome name of the reference genome used for the GWAS (GRCh37 or GRCh38)
msg Optional name of the column missing from the dataset in question

Value

SNP_LOC_DATA SNP positions and alleles for Homo sapiens extracted from NCBI dbSNP Build 144

Examples

```
SNP_LOC_DATA <- load_snp_loc_data("GRCH37")
```

| | |
|------------|----------------------------------|
| parse_logs | <i>Parse data from log files</i> |
|------------|----------------------------------|

Description

Parses data from the log files generated by [format_sumstats](#) or [import_sumstats](#) when the argument `log_mungesumstats_msgs` is set to TRUE.

Usage

```
parse_logs(  
  save_dir = getwd(),  
  pattern = "MungeSumstats_log_msg.txt$",  
  verbose = TRUE  
)
```

Arguments

save_dir Top-level directory to recursively search for log files within.
pattern Regex pattern to search for files with.
verbose Print messages.

Value

[data.table](#) of parsed log data.

Examples

```
save_dir <- system.file("extdata", package = "MungeSumstats")  
log_data <- MungeSumstats::parse_logs(save_dir = save_dir)
```

| | |
|------------|--|
| raw_ALSvcf | <i>GWAS Amyotrophic lateral sclerosis ieu open GWAS project - Subset</i> |
|------------|--|

Description

VCF (VCFv4.2) of the GWAS Amyotrophic lateral sclerosis ieu open GWAS project Dataset: ebi-a-GCST005647. A subset of 99 SNPs

Format

vcf document with 528 items relating to 99 SNPs

Details

A VCF file (VCFv4.2) of the GWAS Amyotrophic lateral sclerosis ieu open GWAS project has been subsetting here to act as an example summary statistic file in VCF format which has some issues in the formatting. MungeSumstats can correct these issues and produced a standardised summary statistics format.

ALSvcf.vcf

NA

Source

The summary statistics VCF (VCFv4.2) file was downloaded from <https://gwas.mrcieu.ac.uk/datasets/ebi-a-GCST005647/> and formatted to a .rda with the following: #Get example VCF dataset, use GWAS Amyotrophic lateral sclerosis ALS_GWAS_VCF <- readLines("ebi-a-GCST005647.vcf.gz") #Subset to just the first 99 SNPs ALSvcf <- ALS_GWAS_VCF[1:528] writeLines(ALSvcf, "inst/extdata/ALSvcf.v")

| | |
|--------------------|--|
| raw_eduAttain0kbay | <i>GWAS Educational Attainment Okbay 2016 - Subset</i> |
|--------------------|--|

Description

GWAS Summary Statistics on Educational Attainment by Okbay et al 2016: PMID: 27898078 PMID: PMC5509058 DOI: 10.1038/ng1216-1587b. A subset of 93 SNPs

Format

txt document with 94 items

Details

GWAS Summary Statistics on Educational Attainment by Okbay et al 2016 has been subsetting here to act as an example summary statistic file which has some issues in the formatting. MungeSumstats can correct these issues.

eduAttainOkbay.txt

NA

Source

The summary statistics file was downloaded from <https://www.nature.com/articles/ng.3552> and formatted to a .rda with the following: #Get example dataset, use Educational-Attainment_Okbay_2016 link<-"Educational-Attainment_Okbay_2016/EduYears_Discovery_5000.txt" eduAttainOkbay<-readLines(link) #There is an issue where values end with .0, this 0 is removed in func #There are also SNPs not on ref genome or are bi/tri allelic #So need to remove these in this dataset as its used for testing tmp <- tempfile() writelines(eduAttainOkbay,con=tmp) eduAttainOkbay <- data.table::fread(tmp) #DT read removes the .0's #remove those not on ref genome and with bi/tri allelic rmv <- c("rs192818565","rs79925071","rs1606974","rs1871109", "rs73074378","rs7955289") eduAttainOkbay <- eduAttainOkbay[!MarkerName %in% c("rs192818565","rs79925071","rs1606974","rs1871109", "rs73074378","rs7955289")] #write to file data.table::fwrite(eduAttainOkbay,file=tmp,sep="\t") eduAttainOkbay <- readLines(tmp) writelines(eduAttainOkbay,"inst/extdata/eduAttainOkbay.txt")

read_sumstats

*Determine summary statistics file type and read them into memory***Description**

Determine summary statistics file type and read them into memory

Usage

```
read_sumstats(
  path,
  nrows = Inf,
  standardise_headers = FALSE,
  samples = 1,
  add_sample_names = FALSE,
  nThread = 1,
  mapping_file = sumstatsColHeaders
)
```

Arguments

| | |
|---------------------|--|
| path | Filepath for the summary statistics file to be formatted. A dataframe or datatable of the summary statistics file can also be passed directly to MungeSumstats using the path parameter. |
| nrows | integer. The (maximal) number of lines to read. If Inf, will read in all rows. |
| standardise_headers | Standardise headers first. |
| samples | Which samples to use: <ul style="list-style-type: none"> • 1 : Only the first sample will be used (<i>DEFAULT</i>). |

- NULL : All samples will be used.
- c("<sample_id1>", "<sample_id2>", ...) : Only user-selected samples will be used (case-insensitive).

`add_sample_names` Append sample names to column names (e.g. "EZ" → "EZ_ubm-a-2929").

`nThread` Number of threads to use for parallel processes.

`mapping_file` MungeSumstats has a pre-defined column-name mapping file which should cover the most common column headers and their interpretations. However, if a column header that is in your file is missing of the mapping we give is incorrect you can supply your own mapping file. Must be a 2 column dataframe with column names "Uncorrected" and "Corrected". See `data(sumstatsColHeaders)` for default mapping and necessary format.

Value

data.table of formatted summary statistics

Examples

```
path <- system.file("extdata", "eduAttain0kbay.txt",
  package = "MungeSumstats"
)
eduAttain0kbay <- read_sumstats(path = path)
```

read_vcf

Read in VCF file

Description

Read in a VCF file as a [VCF](#) or a [data.table](#). Can optionally save the VCF/data.table as well.

Usage

```
read_vcf(
  path,
  write_vcf = FALSE,
  save_path = NULL,
  tabix_index = FALSE,
  samples = 1,
  which = NULL,
  use_params = TRUE,
  sampled_rows = 1e+07,
  nThread = 1,
  verbose = TRUE
)
```

Arguments

| | |
|-------------|--|
| path | Filepath for the summary statistics file to be formatted. A dataframe or datatable of the summary statistics file can also be passed directly to MungeSumstats using the path parameter. |
| write_vcf | Whether to write as VCF (TRUE) or tabular file (FALSE). |
| save_path | File path to save formatted data. Defaults to tempfile(fileext=".tsv.gz"). |
| tabix_index | Index the formatted summary statistics with tabix for fast querying. |
| samples | Which samples to use: <ul style="list-style-type: none"> • 1 : Only the first sample will be used (<i>DEFAULT</i>). • NULL : All samples will be used. • c("<sample_id1>",<sample_id2>",<...>") : Only user-selected samples will be used (case-insensitive). |
| which | A GRanges describing the sequences and ranges to be queried. Variants whose POS lies in the interval(s) [start, end] are returned. If which is not specified all ranges are returned. |
| use_params | When TRUE (default), increases the speed of reading in the VCF by omitting columns that are empty based on the head of the VCF (NAs only). NOTE that that this requires the VCF to be sorted, bgzip-compressed, tabix-indexed, which read_vcf will attempt to do. |
| sampld_rows | First N rows to sample. Set NULL to use full sumstats_file. when determining whether cols are empty. |
| nThread | Number of threads to use for parallel processes. |
| verbose | Print messages. |

Value

The VCF file in data.table format.

Source

```
##### Benchmarking ##### library(VCFwrenchR) library(VariantAnnotation) path <- "https://gwas.mrcieu.ac.uk/files/ieu-a-298/ieu-a-298.vcf.gz"
vcf <- VariantAnnotation::readVcf(file = path) N <- 1e5 vcf_sub <- vcf[1:N,] res <- microbenchmark::microbenchmark("vcf2df"={dat1 <- MungeSumstats::vcf2df(vcf = vcf_sub)}, "VCFwrenchR"={dat2 <- as.data.frame(x = vcf_sub)}, "VRanges"={dat3 <- data.table::as.data.table(methods::as(vcf_sub, "VRanges"))}, times=1)
```

[Discussion on VariantAnnotation GitHub](#)

Examples

```
##### Local file #####
path <- system.file("extdata", "ALSvcf.vcf", package="MungeSumstats")
sumstats_dt <- read_vcf(path = path)

##### Remote file #####
path <- "https://gwas.mrcieu.ac.uk/files/ieu-a-298/ieu-a-298.vcf.gz"
sumstats_dt2 <- read_vcf(path = path)
```

standardise_header *Standardise the column headers in the Summary Statistics files*

Description

Use a reference data table of common column header names (stored in `sumstatsColHeaders` or user inputted mapping file) to convert them to a standard set, i.e. chromosome -> CHR. This function does not check that all the required column headers are present. The amended header is written directly back into the file

Usage

```
standardise_header(
  sumstats_dt,
  mapping_file = sumstatsColHeaders,
  uppercase_unmapped = TRUE,
  return_list = TRUE
)
```

Arguments

`sumstats_dt` data table obj of the summary statistics file for the GWAS.

`mapping_file` MungeSumstats has a pre-defined column-name mapping file which should cover the most common column headers and their interpretations. However, if a column header that is in your file is missing of the mapping we give is incorrect you can supply your own mapping file. Must be a 2 column dataframe with column names "Uncorrected" and "Corrected". See `data(sumstatsColHeaders)` for default mapping and necessary format.

`uppercase_unmapped` For columns that could not be identified in the `mapping_file`, return them in the same format they were input as (without forcing them to uppercase).

`return_list` Return the `sumstats_dt` within a named list (default: TRUE).

Value

list containing `sumstats_dt`, the modified summary statistics data table object

Examples

```
sumstats_dt <- data.table::fread(system.file("extdata", "eduAttain0kbay.txt",
                                             package = "MungeSumstats"))
sumstats_dt2 <- standardise_header(sumstats_dt=sumstats_dt)
```

| | |
|--------------------|--|
| sumstatsColHeaders | <i>Summary Statistics Column Headers</i> |
|--------------------|--|

Description

List of uncorrected column headers often found in GWAS Summary Statistics column headers. Note the effect allele will always be the A2 allele, this is the approach done for VCF(<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC>) This is enforced with the column header corrections here and also the check allele flipping test.

Usage

```
data("sumstatsColHeaders")
```

Format

dataframe with 2 columns

Source

The code to prepare the .Rda file from the marker file is: # Most the data in the below table comes from the LDSC github wiki data("sumstatsColHeaders") # Make additions to sumstatsColHeaders using github version of MungeSumstats-# shown is an example of adding columns for Standard Error (SE) #se_cols <- data.frame("Uncorrected"=c("SE", "se", "STANDARD.ERROR", # "STANDARD_ERROR", "STANDARD_ERROR", "Corrected"=rep("SE", 5)) #sumstatsColHeaders <- rbind(sumstatsColHeaders, se_cols) #Once additions are made, order & save the new mapping dataset #now sort ordering -important for logic that # uncorrected=corrected comes first sumstatsColHeaders\$ordering <- sumstatsColHeaders\$Uncorrected sumstatsColHeaders <- sumstatsColHeaders[order(sumstatsColHeaders\$Corrected, sumstatsColHeaders\$ordering = TRUE),] rownames(sumstatsColHeaders)<-1:nrow(sumstatsColHeaders) sumstatsColHeaders\$ordering <- NULL #manually move FRWQUENCY to above MAR - github issue 95 frequency <- sumstatsColHeaders[sumstatsColHeaders\$Uncorrected=="MAF",] if(as.integer(rownames(frequency))> sumstatsColHeaders[as.integer(rownames(frequency)),] <- maf sumstatsColHeaders[as.integer(rownames(maf))> sumstatsColHeaders[as.integer(rownames(maf))> maf sumstatsColHeaders[as.integer(rownames(maf))> maf <- frequency } usethis::use_data(sumstatsColHeaders, overwrite = TRUE, internal=TRUE) save(sumstatsColHeaders, file="data/sumstatsColHeaders.rda") # You will need to restart your r session for effects to take account

| | |
|----------------|-------------------------------------|
| write_sumstats | <i>Write sum stats file to disk</i> |
|----------------|-------------------------------------|

Description

Write sum stats file to disk

Usage

```
write_sumstats(
  sumstats_dt,
  save_path,
  sep = "\t",
  write_vcf = FALSE,
  tabix_index = FALSE,
  nThread = 1,
  return_path = FALSE,
  save_path_check = FALSE
)
```

Arguments

| | |
|-----------------|--|
| sumstats_dt | data table obj of the summary statistics file for the GWAS. |
| save_path | File path to save formatted data. Defaults to <code>tempfile(fileext=".tsv.gz")</code> . |
| sep | The separator between columns. Defaults to the character in the set <code>[\t ; :]</code> that separates the sample of rows into the most number of lines with the same number of fields. Use <code>NULL</code> or <code>""</code> to specify no separator; i.e. each line a single character column like <code>base::readLines</code> does. |
| write_vcf | Whether to write as VCF (TRUE) or tabular file (FALSE). |
| tabix_index | Index the formatted summary statistics with <code>tabix</code> for fast querying. |
| nThread | The number of threads to use. Experiment to see what works best for your data on your hardware. |
| return_path | Return <code>save_path</code> . This will have been modified in some cases (e.g. after compressing and tabix-indexing a previously un-compressed file). |
| save_path_check | Ensure path name is valid (given the other arguments) before writing (default: FALSE). |

Value

If `return_path=TRUE`, returns `save_path`. Else returns `NULL`.

Source

VariantAnnotation::writeVcf has some unexpected/silent file renaming behavior

Examples

```
path <- system.file("extdata", "eduAttain0kbay.txt",
  package = "MungeSumstats"
)
eduAttain0kbay <- read_sumstats(path = path)
write_sumstats(
  sumstats_dt = eduAttain0kbay,
  save_path = tempfile(fileext = ".tsv.gz")
)
```

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