

Package ‘MungeSumstats’

August 11, 2025

Type Package

Title Standardise summary statistics from GWAS

Version 1.16.0

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>,
<https://al-murphy.github.io/MungeSumstats/>

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

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Depends R(>= 4.1)

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GenomeInfoDb, IRanges, ieugwasr(>= 1.0.1), BSgenome,
Biostrings, stringr, VariantAnnotation, methods, parallel,
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BSgenome.Hsapiens.1000genomes.hs37d5,
BSgenome.Hsapiens.NCBI.GRCh38, BiocGenerics, S4Vectors,
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Author Alan Murphy [aut, cre] (ORCID: <<https://orcid.org/0000-0002-2487-8753>>),

Brian Schilder [aut, ctb] (ORCID:

<<https://orcid.org/0000-0001-5949-2191>>),

Nathan Skene [aut] (ORCID: <<https://orcid.org/0000-0002-6807-3180>>)

Maintainer Alan Murphy <alanmurph94@hotmail.com>

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axel	<i>axel downloader</i>
------	------------------------

Description

R wrapper for axel, which enables multi-threaded download of a single large file.

Usage

```
axel(  
  input_url,  
  output_path,  
  background = FALSE,  
  nThread = 1,  
  force_overwrite = FALSE,  
  quiet = TRUE,  
  alternate = TRUE,  
  check_certificates = FALSE  
)
```

Arguments

input_url	input_url.
output_path	output_path.
background	Run in background
nThread	Number of threads to parallelize over.
force_overwrite	Overwrite existing file.
quiet	Run quietly.
alternate	alternate,
check_certificates	check_certificates

Value

Path where the file has been downloaded

See Also

<https://github.com/axel-download-accelerator/axel/>

Other downloaders: [downloader\(\)](#)

check_allele_flip	<i>Ensure A1 & A2 are correctly named, if GWAS SNP constructed as Alternative/Reference or Risk/Nonrisk alleles these SNPs will need to be converted to Reference/Alternative or Nonrisk/Risk. Here non-risk is defined as what's on the reference genome (this may not always be the case).</i>
-------------------	--

Description

Ensure A1 & A2 are correctly named, if GWAS SNP constructed as Alternative/Reference or Risk/Nonrisk alleles these SNPs will need to be converted to Reference/Alternative or Nonrisk/Risk. Here non-risk is defined as what's on the reference genome (this may not always be the case).

Usage

```
check_allele_flip(
  sumstats_dt,
  path,
  ref_genome,
  rsids,
  allele_flip_check,
  allele_flip_drop,
  allele_flip_z,
  allele_flip_frq,
  bi_allelic_filter,
  flip_frq_as_biallelic,
  imputation_ind,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files,
  standardise_headers = FALSE,
  mapping_file,
  dbSNP
)
```

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.

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-bi-allelic SNPs be removed. Default is TRUE.
flip_frq_as_biallelic	Binary Should non-bi-allelic SNPs frequency values be flipped as 1-p despite there being other alternative alleles? Default is FALSE but if set to TRUE, this allows non-bi-allelic SNPs to be kept despite needing flipping.
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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations
standardise_headers	Run standardise_sumstats_column_headers_crossplatform first.
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.
dbSNP	version of dbSNP to be used for imputation (144 or 155).

Value

A list containing two data tables:

- sumstats_dt: the modified summary statistics data.table object.
- rsids: snpsById, filtered to SNPs of interest if loaded already. Or else NULL.
- log_files: log file list

check_allele_merge	<i>Ensure that A1:A2 or A1/A2 or A1>A2 or A2>A1 aren't merged into 1 column</i>
--------------------	---

Description

Ensure that A1:A2 or A1/A2 or A1>A2 or A2>A1 aren't merged into 1 column

Usage

```
check_allele_merge(sumstats_dt, path)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS
path	Filepath for the summary statistics file to be formatted

Value

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

check_bi_allelic	<i>Remove non-biallelic SNPs</i>
------------------	----------------------------------

Description

Remove non-biallelic SNPs

Usage

```
check_bi_allelic(
  sumstats_dt,
  path,
  ref_genome,
  bi_allelic_filter,
  rsids,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files,
  dbSNP
)
```

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.
bi_allelic_filter	Binary Should non-bi-allelic SNPs be removed. Default is TRUE.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations
dbSNP	version of dbSNP to be used for imputation (144 or 155).

Value

A list containing two data tables:

- sumstats_dt: the modified summary statistics data table object
- rsids: snpsById, filtered to SNPs of interest if loaded already. Or else NULL.
- log_files: log file list

check_bp_range	<i>Ensure that the Base-pair column values are all within the range for the chromosome</i>
----------------	--

Description

Ensure that the Base-pair column values are all within the range for the chromosome

Usage

```
check_bp_range(
  sumstats_dt,
  path,
  ref_genome,
  log_folder_ind,
  imputation_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files
)
```

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.
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 .tsv.gz. Default is FALSE.
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.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations

Value

list containing sumstats_dt, the modified summary statistics data table object and the log file list

check_chr	<i>Standardize the CHR column</i>
-----------	-----------------------------------

Description

Maps chromosome names to the default Ensembl/NCBI naming style and removes SNPs with non-standard CHR entries. Optionally, also removes SNPs on user-specified chromosomes.

Usage

```
check_chr(
  sumstats_dt,
  log_files,
  check_save_out,
  rmv_chr,
  nThread,
  tabix_index,
  log_folder_ind
)
```

Arguments

sumstats_dt	data.table with summary statistics
log_files	list of locations for all log files
check_save_out	list of parameters for saved files
rmv_chr	Chromosomes to exclude from the formatted summary statistics file. Use NULL if no filtering is necessary. Default is c("X", "Y", "MT") which removes all non-autosomal SNPs.
nThread	Number of threads to use for parallel processes.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
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 .tsv.gz. Default is FALSE.

Value

list containing the updated summary statistics data.table and the updated log file locations list

check_col_order	<i>Ensure that the first three columns are SNP, CHR, BP in that order and then A1, A2 if present</i>
-----------------	--

Description

Ensure that the first three columns are SNP, CHR, BP in that order and then A1, A2 if present

Usage

```
check_col_order(sumstats_dt, path)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS
path	Filepath for the summary statistics file to be formatted

Value

list containing sumstats_dt, the modified summary statistics data table object

check_drop_indels	<i>Drop Indels from summary statistics</i>
-------------------	--

Description

Drop Indels from summary statistics

Usage

```
check_drop_indels(
  sumstats_dt,
  drop_indels,
  path,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files
)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS
drop_indels	Binary, should any indels found in the sumstats be dropped? These can not be checked against a reference dataset and will have the same RS ID and position as SNPs which can affect downstream analysis. Default is False.
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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.

Value

list containing sumstats_dt, the modified summary statistics data table object

Source

```
sumstats_dt <- MungeSumstats::formatted_example() sumstats <- check_drop_indels(sumstats_dt
= sumstats_dt, drop_indels = TRUE)
```

check_dup_bp

*Ensure all rows have unique positions, drop those that don't***Description**

Ensure all rows have unique positions, drop those that don't

Usage

```
check_dup_bp(
  sumstats_dt,
  bi_allelic_filter,
  check_dups,
  indels,
  path,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files
)
```

Arguments

bi_allelic_filter	Binary Should non-bi-allelic SNPs be removed. Default is TRUE.
check_dups	whether to check for duplicates - if formatting QTL datasets this should be set to FALSE otherwise keep as TRUE. Default 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.
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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations

Value

list containing sumstats_dt, the modified summary statistics data table object and log files list

check_dup_col	<i>Ensure that no columns are duplicated</i>
---------------	--

Description

Ensure that no columns are duplicated

Usage

```
check_dup_col(sumstats_dt, path)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS
path	Filepath for the summary statistics file to be formatted

Value

list containing sumstats_dt, the modified summary statistics data table object

check_dup_row	<i>Ensure all rows are unique based on SNP,CHR,BP,A1,A2, drop those that aren't</i>
---------------	---

Description

Ensure all rows are unique based on SNP,CHR,BP,A1,A2, drop those that aren't

Usage

```
check_dup_row(
  sumstats_dt,
  check_dups,
  path,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files
)
```

Arguments

check_dups	whether to check for duplicates - if formatting QTL datasets this should be set to FALSE otherwise keep as TRUE. Default is TRUE.
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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations

Value

list containing sumstats_dt, the modified summary statistics data table object and log files list

check_dup_snp	<i>Ensure all rows have unique SNP IDs, drop those that don't</i>
---------------	---

Description

Ensure all rows have unique SNP IDs, drop those that don't

Usage

```
check_dup_snp(
  sumstats_dt,
  indels,
  path,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files,
  bi_allelic_filter,
  check_dups
)
```

Arguments

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.
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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations
bi_allelic_filter	Binary Should non-bi-allelic SNPs be removed. Default is TRUE.
check_dups	whether to check for duplicates - if formatting QTL datasets this should be set to FALSE otherwise keep as TRUE. Default is TRUE.

Value

list containing sumstats_dt, the modified summary statistics data table object and log files list

check_effect_columns_nonzero

Ensure that the standard error (se) is positive for all SNPs

Description

Ensure that the standard error (se) is positive for all SNPs

Usage

```
check_effect_columns_nonzero(
  sumstats_dt,
  path,
  effect_columns_nonzero,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files
)
```

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.
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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations

Value

list containing sumstats_dt, the modified summary statistics data table object and the log file list

check_empty_cols	<i>Check for empty columns</i>
------------------	--------------------------------

Description

Empty columns contain only ".", NA, or 0

Usage

```
check_empty_cols(sumstats_dt, sampled_rows = NULL, verbose = TRUE)
```

Arguments

sampled_rows	First N rows to sample. Set NULL to use full sumstats_file. when determining whether cols are empty.
verbose	Print messages.

Value

empty_cols

check_four_step_col	<i>Ensure that CHR:BP:A2:A1 aren't merged into 1 column</i>
---------------------	---

Description

Ensure that CHR:BP:A2:A1 aren't merged into 1 column

Usage

```
check_four_step_col(sumstats_dt, path)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS
path	Filepath for the summary statistics file to be formatted

Value

list containing sumstats_dt, the modified summary statistics data table object

check_frq	<i>Ensure all SNPs have frq score above threshold</i>
-----------	---

Description

Ensure all SNPs have frq score above threshold

Usage

```
check_frq(
  sumstats_dt,
  path,
  FRQ_filter,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files
)
```

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.
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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations

Value

list containing sumstats_dt, the modified summary statistics data table object and the log file list

check_frq_maf	<i>Check that FRQ column refers to minor/effect allele frequency not major</i>
---------------	--

Description

Check that FRQ column refers to minor/effect allele frequency not major

Usage

```
check_frq_maf(sumstats_dt, frq_is_maf)
```

Arguments

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.
------------	---

Value

sumstats_dt, the modified summary statistics data table object

check_info_score	<i>Ensure all SNPs have info score above threshold</i>
------------------	--

Description

Ensure all SNPs have info score above threshold

Usage

```
check_info_score(  
    sumstats_dt,  
    INFO_filter,  
    log_folder_ind,  
    check_save_out,  
    tabix_index,  
    nThread,  
    log_files  
)
```

Arguments

- INFO_filter numeric The minimum value permissible of the imputation information score (if present in sumstats file). Default 0.9.
- 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 .tsv.gz. Default is FALSE.
- tabix_index Index the formatted summary statistics with **tabix** for fast querying.
- nThread Number of threads to use for parallel processes.
- log_files list of log file locations.

Value

list containing sumstats_dt, the modified summary statistics data table object and the log file list

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,
  save_format,
  convert_n_int,
  allele_flip_check,
  compute_z,
  compute_n
)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS.
save_format	Output format of sumstats. Options are NULL - standardised output format from MungeSumstats, LDSC - output format compatible with LDSC and openGWAS - output compatible with openGWAS VCFs. Default is NULL. NOTE - If LDSC format is used, the naming convention of A1 as the reference (genome build) allele and A2 as the effect allele will be reversed to match LDSC (A1 will now be the effect allele). See more info on this here . Note that any effect columns (e.g. Z) will be in relation to A1 now instead of A2.
convert_n_int	Binary, if N (the number of samples) is not an integer, should this be rounded? Default is TRUE.
allele_flip_check	Binary Should the allele columns be checked against reference genome to infer if flipping is necessary. Default is TRUE.
compute_z	Whether to compute Z-score column. Default is FALSE. This can be computed from Beta and SE with (Beta/SE) or P ($Z := \text{sign}(\text{BETA}) * \sqrt{\text{stats::qchisq}(P, 1, \text{lower} = \text{FALSE})}$). Note that imputing the Z-score from P for every SNP will not be perfectly correct and may result in a loss of power. This should only be done as a last resort. Use 'BETA' to impute by BETA/SE and 'P' to impute by SNP p-value.
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.

Details

[LDSC documentation.](#)

Value

Formatted summary statistics

Source

[LDSC GitHub](#)

check_miss_data	<i>Remove SNPs with missing data</i>
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Description

Remove SNPs with missing data

Usage

```
check_miss_data(  
  sumstats_dt,  
  path,  
  log_folder_ind,  
  check_save_out,  
  tabix_index,  
  nThread,  
  log_files,  
  drop_na_cols  
)
```

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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations
drop_na_cols	A character vector of column names to be checked for missing values. Rows with missing values in any of these columns (if present in the dataset) will be dropped. If NULL, all columns will be checked for missing values. Default columns are SNP, chromosome, position, allele 1, allele2, effect columns (frequency, beta, Z-score, standard error, log odds, signed sumstats, odds ratio), p value and N columns.

Value

list containing sumstats_dt, the modified summary statistics data table object and a log file list.

check_multi_gwas	<i>Ensure that only one model in GWAS sumstats or only one trait tested</i>
------------------	---

Description

Ensure that only one model in GWAS sumstats or only one trait tested

Usage

```
check_multi_gwas(
  sumstats_dt,
  path,
  analysis_trait,
  ignore_multi_trait,
  mapping_file
)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS
path	Filepath for the summary statistics file to be formatted
analysis_trait	If multiple traits were studied, name of the trait for analysis from the GWAS. Default is NULL
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

list containing sumstats_dt, the modified summary statistics data table object

check_multi_rs_snp	<i>Ensure that SNP ids don't have multiple rs ids on one line</i>
--------------------	---

Description

Ensure that SNP ids don't have multiple rs ids on one line

Usage

```
check_multi_rs_snp(
  sumstats_dt,
  path,
  remove_multi_rs_snp,
  imputation_ind,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files
)
```

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. |
| 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. |
| 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. |
| 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 .tsv.gz. Default is FALSE. |
| tabix_index | Index the formatted summary statistics with tabix for fast querying. |
| nThread | Number of threads to use for parallel processes. |
| log_files | list of log file locations |

Value

list containing sumstats_dt, the modified summary statistics data table object and the log file list.

check_no_allele	<i>Ensure that A1 & A2 are present, if not can find it with SNP and other allele</i>
-----------------	--

Description

More care needs to be taken if one of A1/A2 is present, before imputing the other allele flipping needs to be checked

Usage

```
check_no_allele(
  sumstats_dt,
  path,
  ref_genome,
  rsids,
  imputation_ind,
  allele_flip_check,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files,
  bi_allelic_filter,
  dbSNP
)
```

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.
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.
allele_flip_check	Binary Should the allele columns be checked against reference genome to infer if flipping is necessary. Default is TRUE.

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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations
bi_allelic_filter	Binary Should non-bi-allelic SNPs be removed. Default is TRUE.
dbSNP	version of dbSNP to be used for imputation (144 or 155).

Value

A list containing two data tables:

- sumstats_dt: the modified summary statistics data table object
- rsids: snpsById, filtered to SNPs of interest if loaded already. Or else NULL.
- allele_flip_check: does the dataset require allele flip check
- log_files: log file list
- bi_allelic_filter: should multi-allelic SNPs be filtered out

check_no_chr_bp	<i>Ensure that CHR and BP are missing if SNP is present, can find them</i>
-----------------	--

Description

Ensure that CHR and BP are missing if SNP is present, can find them

Usage

```
check_no_chr_bp(
  sumstats_dt,
  path,
  ref_genome,
  rsids,
  imputation_ind,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files,
  dbSNP
)
```

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.
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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations
dbSNP	version of dbSNP to be used for imputation (144 or 155).

Value

A list containing two data tables:

- sumstats_dt : the modified summary statistics data table object
- rsids : snpsById, filtered to SNPs of interest if loaded already. Or else NULL
- log_files : log file list

check_no_rs_snp	<i>Ensure that SNP appears to be valid RSIDs (starts with rs)</i>
-----------------	---

Description

Ensure that SNP appears to be valid RSIDs (starts with rs)

Usage

```

check_no_rs_snp(
  sumstats_dt,
  path,
  ref_genome,
  snp_ids_are_rs_ids,
  indels,
  imputation_ind,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files,
  dbSNP
)

```

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.
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.
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.
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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations
dbSNP	version of dbSNP to be used for imputation (144 or 155).

Value

list containing sumstats_dt, the modified summary statistics data table object and the log file list.

check_no_snp

Ensure that SNP is present if not can find it with CHR and BP

Description

Ensure that SNP is present if not can find it with CHR and BP

Usage

```
check_no_snp(
  sumstats_dt,
  path,
  ref_genome,
  indels,
  imputation_ind,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files,
  dbSNP,
  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.
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.
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.
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.

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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations
dbSNP	version of dbSNP to be used for imputation (144 or 155).
verbose	should messages be printed. Default it TRUE.

Value

list containing sumstats_dt, the modified summary statistics data table object and the log files list

check_numeric	<i>Check numeric columns</i>
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Description

Checks for any columns that should be numeric, and ensures that they are indeed numeric.

Usage

```
check_numeric(sumstats_dt, cols = c("P", "SE", "FRQ", "MAF", "BETA"))
```

Arguments

sumstats_dt	Summary stats with column names already standardised by format_sumstats .
cols	Names of columns that should be numeric. If any of these columns are not actually present in sumstats_dt, they will be skipped.

Value

sumstats_dt

check_n_int	<i>Ensure that the N column is all integers</i>
-------------	---

Description

Ensure that the N column is all integers

Usage

```
check_n_int(sumstats_dt, path, convert_n_int, imputation_ind)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS
path	Filepath for the summary statistics file to be formatted
convert_n_int	Binary, if N (the number of samples) is not an integer, should this be rounded? Default is TRUE.
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.

Value

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

check_n_num	<i>Ensure all SNPs have N less than X std dev below mean</i>
-------------	--

Description

In case some SNPs were genotyped by a specialized genotyping array and have substantially more samples than others. These will be removed.

Usage

```
check_n_num(
  sumstats_dt,
  path,
  N_std,
  N_dropNA = FALSE,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files
)
```

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.
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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations

Value

list containing sumstats_dt, the modified summary statistics data table object and the log file list

check_on_ref_genome	<i>Ensure all SNPs are on the reference genome</i>
---------------------	--

Description

Ensure all SNPs are on the reference genome

Usage

```
check_on_ref_genome(
  sumstats_dt,
  path,
  ref_genome,
  on_ref_genome,
  indels = indels,
  rsids,
  imputation_ind,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files,
  dbSNP
)
```

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.
on_ref_genome	Binary Should a check take place that all SNPs are on the reference genome by SNP ID. Default 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.
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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations
dbSNP	version of dbSNP to be used for imputation (144 or 155).

Value

A list containing two data tables:

- `sumstats_dt` : the modified summary statistics data table object
- `rsids` : `snpsById`, filtered to SNPs of interest if loaded already. Or else NULL
- `log_files` : log file list

check_pos_se	<i>Ensure that the standard error (se) is positive for all SNPs Also impute se if missing</i>
--------------	---

Description

Ensure that the standard error (se) is positive for all SNPs Also impute se if missing

Usage

```
check_pos_se(
  sumstats_dt,
  path,
  pos_se,
  log_folder_ind,
  imputation_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files,
  impute_se
)
```

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.
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.
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 .tsv.gz. Default is FALSE.
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.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations
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.

Value

list containing sumstats_dt, the modified summary statistics data table object and the log file list

check_range_p_val	<i>Ensure that the p values are not >1 and if so set to 1</i>
-------------------	--

Description

Ensure that the p values are not >1 and if so set to 1

Usage

```
check_range_p_val(sumstats_dt, convert_large_p, convert_neg_p, imputation_ind)
```

Arguments

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

Value

list containing sumstats_dt, the modified summary statistics data table object

Source

```
sumstats_dt <- MungeSumstats::formatted_example() sumstats_dt$P[1:3] <- 5 sumstats_dt$P[6:10]
<- -5 sumstats <- check_range_p_val(sumstats_dt = sumstats_dt, convert_large_p = TRUE,
convert_neg_p = TRUE, imputation_ind = TRUE)
```

check_row_snp	<i>Ensure all rows have SNPs beginning with rs or SNP, drop those that don't</i>
---------------	--

Description

Ensure all rows have SNPs beginning with rs or SNP, drop those that don't

Usage

```
check_row_snp(
  sumstats_dt,
  path,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files
)
```

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.
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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations

Value

list containing sumstats_dt, the modified summary statistics data table object and log file list

check_save_path	<i>Check if save path and log folder is appropriate</i>
-----------------	---

Description

Check if save path and log folder is appropriate

Usage

```
check_save_path(
  save_path,
  log_folder,
  log_folder_ind,
  tabix_index,
  write_vcf = FALSE,
  verbose = TRUE
)
```

Arguments

save_path	File path to save formatted data. Defaults to <code>tempfile(fileext=".tsv.gz")</code> .
log_folder	Filepath to the directory for the log files and the log of MungeSumstats messages to be stored. Default is a temporary directory. Note the name of the log files (log messages and log outputs) are now the same as the name of the file specified in the save path parameter with the extension <code>'_log_msg.txt'</code> and <code>'_log_output.txt'</code> respectively.
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.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
write_vcf	Whether to write as VCF (TRUE) or tabular file (FALSE).
verbose	Print messages.

Value

Corrected save_path, the file type, the separator, corrected log_folder, the log file extension.

check_signed_col	<i>Ensure that there is at least one signed column in summary statistics file Impute beta if user requests</i>
------------------	--

Description

Ensure that there is at least one signed column in summary statistics file Impute beta if user requests

Usage

```
check_signed_col(
  sumstats_dt,
  impute_beta,
  log_folder_ind,
  rsids,
  imputation_ind,
  check_save_out,
  tabix_index,
  log_files,
  nThread
)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS
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 (for Z & SE approach) 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.
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 .tsv.gz. Default is FALSE.
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.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
log_files	list of log file locations
nThread	Number of threads to use for parallel processes.

Value

null

check_small_p_val	<i>Ensure that the non-negative p-values are not 5e-324 or lower, if so set to 0</i>
-------------------	--

Description

Ensure that the non-negative p-values are not 5e-324 or lower, if so set to 0

Usage

```
check_small_p_val(sumstats_dt, convert_small_p, imputation_ind)
```

Arguments

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

Value

list containing sumstats_dt, the modified summary statistics data table object

Source

```
sumstats_dt <- MungeSumstats::formatted_example() sumstats_dt$P[1:3] <- 5e-324 sumstats_dt$P[6:10]
<- "5e-324" sumstats <- check_small_p_val(sumstats_dt = sumstats_dt, convert_small_p
= TRUE, imputation_ind = TRUE)
```

check_strand_ambiguous

Remove SNPs with strand-ambiguous alleles

Description

Remove SNPs with strand-ambiguous alleles

Usage

```
check_strand_ambiguous(
  sumstats_dt,
  path,
  ref_genome,
  strand_ambig_filter,
  log_folder_ind,
  check_save_out,
  tabix_index,
  nThread,
  log_files
)
```

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.
strand_ambig_filter	Binary Should SNPs with strand-ambiguous alleles be removed. 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 .tsv.gz. Default is FALSE.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
nThread	Number of threads to use for parallel processes.
log_files	list of log file locations

Value

list containing sumstats_dt, the modified summary statistics data table object and the log file list

check_tabular	<i>Ensure valid tabular format</i>
---------------	------------------------------------

Description

Ensure valid tabular format

Usage

```
check_tabular(header)
```

Arguments

header	The summary statistics file for the GWAS
--------	--

Value

Whether the file is tabular

check_two_step_col	<i>Ensure that CHR:BP aren't merged into 1 column</i>
--------------------	---

Description

Ensure that CHR:BP aren't merged into 1 column

Usage

```
check_two_step_col(sumstats_dt, path)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS
path	Filepath for the summary statistics file to be formatted

Value

list containing sumstats_dt, the modified summary statistics data table object

check_vcf	<i>Check if the inputted file is in VCF format</i>
-----------	--

Description

Check if the inputted file is in VCF format

Usage

```
check_vcf(header)
```

Arguments

header	Header of the GWAS summary statistics file.
--------	---

Value

Whether the file is vcf or not

check_vital_col	<i>Ensure that all necessary columns are in the summary statistics file</i>
-----------------	---

Description

Ensure that all necessary columns are in the summary statistics file

Usage

```
check_vital_col(sumstats_dt)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS
-------------	--

Value

null

check_zscore	<i>Check for Z-score column</i>
--------------	---------------------------------

Description

The following ensures that a Z-score column is present. The Z-score formula we used here is a R implementation of the formula used in **LDSC's** `munge_sumstats.py`:

Usage

```
check_zscore(
    sumstats_dt,
    imputation_ind,
    compute_z = "BETA",
    force_new_z = FALSE,
    standardise_headers = FALSE,
    mapping_file
)
```

Arguments

<code>sumstats_dt</code>	data table obj of the summary statistics file for the GWAS.
<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). Note these columns will be in the formatted summary statistics returned. Default is FALSE.
<code>compute_z</code>	Whether to compute Z-score column. Default is FALSE. This can be computed from Beta and SE with (Beta/SE) or P ($Z := \text{sign}(\text{BETA}) * \sqrt{\text{stats::qchisq}(P, 1, \text{lower} = \text{FALSE})}$). Note that imputing the Z-score from P for every SNP will not be perfectly correct and may result in a loss of power. This should only be done as a last resort. Use 'BETA' to impute by BETA/SE and 'P' to impute by SNP p-value.
<code>force_new_z</code>	When a "Z" column already exists, it will be used by default. To override and compute a new Z-score column from P set <code>force_new_z=TRUE</code> .
<code>standardise_headers</code>	Run <code>standardise_sumstats_column_headers_crossplatform</code> first.
<code>mapping_file</code>	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 <code>data(sumstatsColHeaders)</code> for default mapping and necessary format.

Details

```
np.sqrt(chi2.isf(P, 1))
```

The R implementation is adapted from the `GenomicSEM::munge` function, after optimizing for speed using `data.table`:

```
sumstats_dt[,Z:=sign(BETA)*sqrt(stats::qchisq(P,1,lower=FALSE))]
```

NOTE: `compute_z` is set to `TRUE` by default to ensure standardisation of the "Z" column (which can be computed differently in different datasets).

Value

```
list("sumstats_dt"=sumstats_dt)
```

column_dictionary	<i>Map column names to positions.</i>
-------------------	---------------------------------------

Description

Useful in situations where you need to specify columns by index instead of name (e.g. awk queries).

Usage

```
column_dictionary(file_path)
```

Arguments

file_path	Path to full summary stats file (or any really file you want to make a column dictionary for).
-----------	--

Value

Named list of column positions.

Source

Borrowed function from [echotabix](#).

```
eduAttainOkbayPth <- system.file("extdata", "eduAttainOkbay.txt", package = "MungeSumstats")
) tmp <- tempfile(fileext = ".tsv") file.copy(eduAttainOkbayPth, tmp) cdict <- MungeSumstats::column_dictionary(tmp)
```

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"). • 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: $N_{\text{eff}} = (N_{\text{CAS}} + N_{\text{CON}}) * (N_{\text{CAS}} / (N_{\text{CAS}} + \text{mean}((N_{\text{CAS}} / (N_{\text{CAS}} + N_{\text{CON}})) * (N_{\text{CAS}} + N_{\text{CON}})) = \max(N_{\text{CAS}} + N_{\text{CON}})))$. • "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)
```

compute_sample_size	<i>Compute (effective) sample size</i>
---------------------	--

Description

Computes sample sum (as new column "N") or effective sample size (ESS) (as new column "Neff"). Computing ESS is important as it takes into account the proportion of cases to controls (i.e. class imbalance) so as not to overestimate your statistical power.

Usage

```
compute_sample_size(
  sumstats_dt,
  method = c("ldsc", "giant", "metal", "sum"),
  force_new = FALSE,
  append_method_name = FALSE
)
```

Arguments

sumstats_dt	Summary statistics data.table.
method	Method for computing (effective) sample size. • "ldsc" : $Neff = (N_{CAS} + N_{CON}) * (N_{CAS} / (N_{CAS} + N_{CON})) / mean((N_{CAS} / (N_{CAS} + N_{CON}))[(N_{CAS} + N_{CON}) == max(N_{CAS} + N_{CON})])$ bulik/ldsc GitHub Issue bulik/ldsc GitHub code • "giant" : $Neff = 2 / (1/N_{CAS} + 1/N_{CON})$ Winkler et al. 2014, Nature

	• "metal" : $Neff = 4 / (1/N_{CAS} + 1/N_{CON})$ Willer et al. 2010, Bioinformatics • "sum" : $N = N_{CAS} + N_{CON}$ Simple summation of cases and controls that does not account for class imbalance. • "\<integer\>" : $N = \text{\<integer\>}$ If method is a positive integer, it will be used as N for every row.
force_new	If "Neff" (or "N") already exists in sumstats_dt, replace it with the recomputed version.
append_method_name	should Neff column have an indicator to explain the method that makes it., Default is FALSE unless multiple methods are passed

Details

There are many different formulas for calculating ESS, but LDSC is probably the best method available here, as it doesn't assume that the proportion of controls:cases is 2:1 (as in GIANT) or 4:1 (as in METAL).

Value

A data.table with a new column "Neff" or "N"

compute_sample_size_n *Add user supplied sample size*

Description

Add user supplied sample size

Usage

```
compute_sample_size_n(sumstats_dt, method, force_new = FALSE)
```

Arguments

sumstats_dt	Summary statistics data.table.
method	Method for computing (effective) sample size. • "ldsc" : $Neff = (N_{CAS} + N_{CON}) * (N_{CAS} / (N_{CAS} + N_{CON})) / mean((N_{CAS} / (N_{CAS} + N_{CON}))[(N_{CAS} + N_{CON}) == max(N_{CAS} + N_{CON})])$ bulik/ldsc GitHub Issue bulik/ldsc GitHub code

- "giant" :
 $N_{eff} = 2/(1/N_{CAS} + 1/N_{CON})$
[Winkler et al. 2014, Nature](#)
- "metal" :
 $N_{eff} = 4/(1/N_{CAS} + 1/N_{CON})$
[Willer et al. 2010, Bioinformatics](#)
- "sum" :
 $N = N_{CAS} + N_{CON}$
 Simple summation of cases and controls that does not account for class imbalance.
- "<integer>" :
 $N = \text{<integer>}$
 If method is a positive integer, it will be used as N for every row.

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

Value

No return

compute_sample_size_neff
Compute Neff/N

Description

Compute Neff/N

Usage

```
compute_sample_size_neff(
  sumstats_dt,
  method,
  force_new = FALSE,
  append_method_name = FALSE
)
```

Arguments

sumstats_dt Summary statistics data.table.

method Method for computing (effective) sample size.

- "ldsc" :
 $N_{eff} = (N_{CAS} + N_{CON}) * (N_{CAS} / (N_{CAS} + N_{CON})) / \text{mean}((N_{CAS} / (N_{CAS} + N_{CON}))[(N_{CAS} + N_{CON}) == \max(N_{CAS} + N_{CON})])$
[bulik/ldsc GitHub Issue](#) [bulik/ldsc GitHub code](#)

- "giant" :
 $N_{eff} = 2/(1/N_{CAS} + 1/N_{CON})$
Winkler et al. 2014, Nature
 - "metal" :
 $N_{eff} = 4/(1/N_{CAS} + 1/N_{CON})$
Willer et al. 2010, Bioinformatics
 - "sum" :
 $N = N_{CAS} + N_{CON}$
Simple summation of cases and controls that does not account for class imbalance.
 - "<integer>" :
 $N = \text{integer}$
If method is a positive integer, it will be used as N for every row.
- force_new If "Neff" (or "N") already exists in sumstats_dt, replace it with the recomputed version.
- append_method_name should Neff column have an indicator to explain the method that makes it., Default is FALSE unless multiple methods are passed

Value

No return

convert_sumstats	<i>Convert summary statistics to desired object type</i>
------------------	--

Description

Convert summary statistics to desired object type

Usage

```
convert_sumstats(  
  sumstats_dt,  
  return_format = c("data.table", "vranges", "granges")  
)
```

Arguments

return_format Object type to convert to; "data.table", "GenomicRanges" or "VRanges"(default is "data.table").

Value

Summary statistics in the converted format

DF_to_dt	<i>DataFrame to data.table</i>
----------	--------------------------------

Description

Efficiently convert [DataFrame](#) to [data.table](#).

Usage

```
DF_to_dt(DF)
```

Arguments

DF [DataFrame](#) object.

Value

VCF data in [data.table](#) format.

Source

[Solution from Bioc forum](#)

downloader	<i>downloader wrapper</i>
------------	---------------------------

Description

R wrapper for [axel](#) (multi-threaded) and [download.file](#) (single-threaded) download functions.

Usage

```
downloader(  
  input_url,  
  output_path,  
  download_method = "axel",  
  background = FALSE,  
  force_overwrite = FALSE,  
  quiet = TRUE,  
  show_progress = TRUE,  
  continue = TRUE,  
  nThread = 1,  
  alternate = TRUE,  
  check_certificates = TRUE,  
  timeout = 10 * 60  
)
```

Arguments

input_url	input_url.
output_path	output_path.
download_method	"axel" (multi-threaded) or "download.file" (single-threaded) .
background	Run in background
force_overwrite	Overwrite existing file.
quiet	Run quietly.
show_progress	show_progress.
continue	continue.
nThread	Number of threads to parallelize over.
alternate	alternate,
check_certificates	check_certificates
timeout	How many seconds before giving up on download. Passed to download.file. Default: 10*60 (10min).

Value

Local path to downloaded file.

Source

Suggestion to avoid 'proc\$get_built_file() : Build process failed'

See Also

Other downloaders: [axel\(\)](#)

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 = FALSE,
  timeout = 10 * 60,
  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 tempdir() 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. vcf_dir="/raw_vcf").
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.
timeout	How many seconds before giving up on download. Passed to download.file. Default: 10*60 (10min).
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)
}
```

drop_duplicate_cols	<i>Drop duplicate columns</i>
---------------------	-------------------------------

Description

Drop columns with identical names (if any exist) within a data.table.

Usage

```
drop_duplicate_cols(dt)
```

Arguments

dt	data.table
----	------------

Value

Null output

drop_duplicate_rows	<i>Drop duplicate rows</i>
---------------------	----------------------------

Description

Drop rows with duplicate values across all columns.

Usage

```
drop_duplicate_rows(dt, verbose = TRUE)
```

Arguments

dt	data.table
verbose	Print messages.

Value

Filtered dt.

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
)
```

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).

Details

To authenticate, you need to generate a token from the OpenGWAS website. The token behaves like a password, and it will be used to authorise the requests you make to the OpenGWAS API. Here are the steps to generate the token and then have ieugwasr automatically use it for your queries:

1. Login to <https://api.opengwas.io/profile/>
2. Generate a new token
3. Add OPENGWAS_JWT=<token> to your .Renviron file, thi can be edited in R by running `usethis::edit_r_environ()`
4. Restart your R session
5. To check that your token is being recognised, run `ieugwasr::get_opengwas_jwt()`. If it returns a long random string then you are authenticated.
6. To check that your token is working, run `ieugwasr::user()`. It will make a a request to the API for your user information using your token. It should return a list with your user information. If it returns an error, then your token is not working.
7. Make sure you have submitted use

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
# and if access token has been added
if(try(is.character(getURL("www.google.com")))==TRUE && ieugwasr::get_opengwas_jwt()!=""){
### 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", "eduAttainOkbay.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,
  chain_source = "ensembl",
  local_chain = 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,
  es_is_beta = TRUE,
  impute_se = FALSE,
  analysis_trait = NULL,
  ignore_multi_trait = FALSE,
  INFO_filter = 0.9,
  FRQ_filter = 0,
  pos_se = TRUE,
  effect_columns_nonzero = FALSE,
  N_std = 5,
  N_dropNA = TRUE,
  chr_style = "Ensembl",
  rmv_chr = c("X", "Y", "MT"),
  on_ref_genome = TRUE,
  infer_eff_direction = TRUE,
  eff_on_minor_alleles = FALSE,
  strand_ambig_filter = FALSE,
  allele_flip_check = TRUE,
  allele_flip_drop = TRUE,
  allele_flip_z = TRUE,
  allele_flip_frq = TRUE,
  bi_allelic_filter = TRUE,
  flip_frq_as_biallelic = FALSE,
  snp_ids_are_rs_ids = TRUE,
  remove_multi_rs_snp = FALSE,
  frq_is_maf = TRUE,
  indels = TRUE,
  drop_indels = FALSE,
  drop_na_cols = c("SNP", "CHR", "BP", "A1", "A2", "FRQ", "BETA", "Z", "OR", "LOG_ODDS",
    "SIGNED_SUMSTAT", "SE", "P", "N"),
  dbSNP = 155,
  check_dups = 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,
save_format = NULL,
log_folder_ind = FALSE,
log_mungesumstats_msgs = FALSE,
log_folder = tempdir(),
imputation_ind = FALSE,
force_new = FALSE,
mapping_file = sumstatsColHeaders,
rmv_chrPrefix = NULL
)

```

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).
chain_source	source of the chain file to use in liftover, if converting genome build ("ucsc" or "ensembl"). Note that the UCSC chain files require a license for commercial use. The Ensembl chain is used by default ("ensembl").
local_chain	Path to local chain file to use instead of downloading. Default of NULL i.e. no local file to use. NOTE if passing a local chain file make sure to specify the path to convert from and to the correct build like GRCh37 to GRCh38. We can not sense check this for local files. The chain file can be submitted as a gz file (as downloaded from source) or unzipped.
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. Default is FALSE. This can be computed from Beta and SE with (Beta/SE) or P ($Z := \text{sign}(\text{BETA}) * \sqrt{\text{stats::qchisq}(P, 1, \text{lower} = \text{FALSE})}$). Note that imputing the Z-score from P for every SNP will not be perfectly correct and may result in a loss of power. This should only be done as a last resort. Use 'BETA' to impute by BETA/SE and 'P' to impute by SNP p-value.
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 (for Z & SE approach) 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.
es_is_beta	Binary, whether to map ES to BETA. We take BETA to be any BETA-like value (including Effect Size). If this is not the case for your sumstats, change this to FALSE. Default is TRUE.
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.
ignore_multi_trait	If you have multiple traits (p-values) in the study but you want to ignore these and instead use a standard named p-value, set to TRUE. By default is FALSE which will check for multi-traits.
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.
chr_style	Chromosome naming style to use in the formatted summary statistics file ("NCBI", "UCSC", "dbSNP", or "Ensembl"). The NCBI and Ensembl styles both code chromosomes as 1-22, X, Y, MT; the UCSC style is chr1-chr22, chrX, chrY, chrM; and the dbSNP style is ch1-ch22, chX, chY, chMT. Default is Ensembl.
rmv_chr	Chromosomes to exclude from the formatted summary statistics file. Use NULL if no filtering is necessary. Default is c("X", "Y", "MT") which removes all non-autosomal SNPs.
on_ref_genome	Binary Should a check take place that all SNPs are on the reference genome by SNP ID. Default is TRUE.
infer_eff_direction	Binary Should a check take place to ensure the alleles match the effect direction? Default is TRUE.
eff_on_minor_alleles	Binary Should MungeSumstats assume that the effects are majoritively measured on the minor alleles? Default is FALSE as this is an assumption that won't be appropriate in all cases. However, the benefit is that if we know the majority of SNPs have their effects based on the minor alleles, we can catch cases where the allele columns have been mislabelled.
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-bi-allelic SNPs be removed. Default is TRUE.
flip_frq_as_biallelic	Binary Should non-bi-allelic SNPs frequency values be flipped as 1-p despite there being other alternative alleles? Default is FALSE but if set to TRUE, this allows non-bi-allelic SNPs to be kept despite needing flipping.

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.
drop_indels	Binary, should any indels found in the sumstats be dropped? These can not be checked against a reference dataset and will have the same RS ID and position as SNPs which can affect downstream analysis. Default is False.
drop_na_cols	A character vector of column names to be checked for missing values. Rows with missing values in any of these columns (if present in the dataset) will be dropped. If NULL, all columns will be checked for missing values. Default columns are SNP, chromosome, position, allele 1, allele2, effect columns (frequency, beta, Z-score, standard error, log odds, signed sumstats, odds ratio), p value and N columns.
dbSNP	version of dbSNP to be used for imputation (144 or 155).
check_dups	whether to check for duplicates - if formatting QTL datasets this should be set to FALSE otherwise keep as 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 tempfile(fileext=".tsv.gz").
write_vcf	Whether to write as VCF (TRUE) or tabular file (FALSE).
tabix_index	Index the formatted summary statistics with tabix for fast querying.
return_data	Return data.table, GRanges or VRanges directly to user. Otherwise, return the path to the save data. Default is FALSE.
return_format	If return_data is TRUE. Object type to be returned ("data.table", "vranges", "granges").
ldsc_format	DEPRECATED, do not use. Use save_format="LDSC" instead.
save_format	Output format of sumstats. Options are NULL - standardised output format from MungeSumstats, LDSC - output format compatible with LDSC and openGWAS - output compatible with openGWAS VCFs. Default is NULL. NOTE - If LDSC format is used, the naming convention of A1 as the reference (genome build)

	allele and A2 as the effect allele will be reversed to match LDSC (A1 will now be the effect allele). See more info on this here . Note that any effect columns (e.g. Z) will be in relation to A1 now instead of A2.
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 .tsv.gz. 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. Note the name of the log files (log messages and log outputs) are now the same as the name of the file specified in the save_path parameter with the extension '_log_msg.txt' and '_log_output.txt' respectively.
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.
rmv_chrPrefix	Is now deprecated, do not use. Use chr_style instead - chr_style = 'Ensembl' will give the same result as rmv_chrPrefix=TRUE used to give.

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
## Using dbSNP = 144 for speed as it's smaller but you should use 155 unless
## you know what you are doing and need 144

is_32bit_windows <-
  .Platform$OS.type == "windows" && .Platform$r_arch == "i386"
if (!is_32bit_windows) {
  reformatted <- format_sumstats(
    path = eduAttainOkbayPth,
    ref_genome = "GRCh37",
    dbSNP = 144
  )
} 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,
    dbSNP=144
  )
}
# returned location has the updated summary statistics file

```

get_chain_file	<i>Download chain file for liftover</i>
----------------	---

Description

Download chain file for liftover

Usage

```

get_chain_file(
  from = c("hg38", "hg19"),
  to = c("hg19", "hg38"),
  chain_source = c("ucsc", "ensembl"),
  save_dir = tempdir(),
  verbose = TRUE
)

```

Arguments

from	genome build converted from ("hg38", "hg19")
to	genome build converted to ("hg19", "hg38")
chain_source	chain file source used ("ucsc" as default, or "ensembl")
save_dir	where is the chain file saved? Default is a temp directory
verbose	extra messages printed? Default is TRUE

Value

loaded chain file for liftover

Source

- UCSC chain files
- Ensembl chain files

get_eff_frq_allele_combns
<i>Get combinations of uncorrected allele and effect (and frq) columns</i>

Description

Get combinations of uncorrected allele and effect (and frq) columns

Usage

```
get_eff_frq_allele_combns(  
  mapping_file = sumstatsColHeaders,  
  eff_frq_cols = c("BETA", "OR", "LOG_ODDS", "SIGNED_SUMSTAT", "Z", "FRQ")  
)
```

Arguments

- | | |
|--------------|--|
| 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. |
| eff_frq_cols | Corrected effect or frequency column names found in a sumstats. Default of BETA, OR, LOG_ODDS, SIGNED_SUMSTAT, Z and FRQ. |

Value

datatable containing uncorrected and corrected combinations

get_genome_build	<i>Infers the genome build of the summary statistics file (GRCh37 or GRCh38) from the data. Uses SNP (RSID) & CHR & BP to get genome build.</i>
------------------	---

Description

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

Usage

```
get_genome_build(
  sumstats,
  nThread = 1,
  sampled_snps = 10000,
  standardise_headers = TRUE,
  mapping_file = sumstatsColHeaders,
  dbSNP = 155,
  header_only = FALSE,
  allele_match_ref = FALSE,
  ref_genome = NULL,
  chr_filt = NULL
)
```

Arguments

sumstats	data table/data frame obj of the summary statistics file for the GWAS ,or file path to summary statistics file.
nThread	Number of threads to use for parallel processes.
sampled_snps	Downsample the number of SNPs used when inferring genome build to save time.
standardise_headers	Run standardise_sumstats_column_headers_crossplatform.
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.
dbSNP	version of dbSNP to be used (144 or 155). Default is 155.
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.

allele_match_ref	Instead of returning the genome_build this will return the proportion of matches to each genome build for each allele (A1,A2).
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.
chr_filt	Internal for testing - filter reference genomes and sumstats to specific chromosomes for testing. Pass a list of chroms in format: c("1","2"). Default is NULL i.e. no filtering

Value

ref_genome the genome build of the data

get_genome_builds	<i>Infer genome builds</i>
-------------------	----------------------------

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,
  dbSNP = 155,
  nThread = 1,
  chr_filt = NULL
)
```

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.

dbSNP	version of dbSNP to be used (144 or 155). Default is 155.
nThread	Number of threads to use for parallel processes.
chr_filt	Internal for testing - filter reference genomes and sumstats to specific chromosomes for testing. Pass a list of chroms in format: c("1","2"). Default is NULL i.e. no filtering

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,
    dbSNP=144)
}
```

get_unique_name_log_file

Simple function to ensure the new entry name to a list doesn't have the same name as another entry

Description

Simple function to ensure the new entry name to a list doesn't have the same name as another entry

Usage

```
get_unique_name_log_file(name, log_files)
```

name	proposed name for the entry
log_files	list of log file locations

a unique name (character)

get_vcf_sample_ids	Get VCF sample ID(s)
--------------------	----------------------

Get VCF sample ID(s)

```
get_vcf_sample_ids(path)
```

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.

sample_id

granges_to_dt *GenomicRanges* to *data.table*

Convert a GRanges into a data.table.

```
granges_to_dt(gr)
```

gr A **GRanges** object.

Value

A data.table object.

Source

Code adapted from GenomicDistributions.

hg19ToHg38	<i>UCSC Chain file hg19 to hg38</i>
------------	-------------------------------------

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.gz')`

hg38ToHg19	<i>UCSC Chain file hg38 to hg19</i>
------------	-------------------------------------

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/ucsc::download.file\('ftp://hgdownload.cse.ucsc.edu/goldenPath/hg38/liftOver/hg38ToHg19.over.chain.gz'\)](https://hgdownload.cse.ucsc.edu/goldenpath/hg38/liftOver/ucsc::download.file('ftp://hgdownload.cse.ucsc.edu/goldenPath/hg38/liftOver/hg38ToHg19.over.chain.gz'))

ieu-a-298	<i>Local ieu-a-298 file from IEU Open GWAS</i>
-----------	--

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, downlaoded 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

ids	List of Open GWAS study IDs (e.g. c("prot-a-664", "ieu-b-4760")).
vcf_dir	Where to download the original VCF from Open GWAS. <i>WARNING:</i> This is set to tempdir() 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. vcf_dir="/raw_vcf").
vcf_download	Download the original VCF from Open GWAS.
save_dir	Directory to save formatted summary statistics in.
write_vcf	Whether to write as VCF (TRUE) or tabular file (FALSE).
download_method	"axel" (multi-threaded) or "download.file" (single-threaded) .
quiet	Run quietly.
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.
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.

...

Arguments passed on to `format_sumstats`

`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).

`chain_source` source of the chain file to use in liftover, if converting genome build ("ucsc" or "ensembl"). Note that the UCSC chain files require a license for commercial use. The Ensembl chain is used by default ("ensembl").

`local_chain` Path to local chain file to use instead of downloading. Default of NULL i.e. no local file to use. NOTE if passing a local chain file make sure to specify the path to convert from and to the correct build like GRCh37 to GRCh38. We can not sense check this for local files. The chain file can be submitted as a gz file (as downloaded from source) or unzipped.

`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. Default is FALSE. This can be computed from Beta and SE with (Beta/SE) or $P (Z := \text{sign}(\text{BETA}) * \sqrt{\text{stats::qchisq}(P, 1, \text{lower} = \text{FALSE})})$. **Note** that imputing the Z-score from P for every SNP will not be perfectly correct and may result in a loss of power. This should only be done as a last resort. Use 'BETA' to impute by BETA/SE and 'P' to impute by SNP p-value.

`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 (for Z & SE approach) 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.
- es_is_beta** Binary, whether to map ES to BETA. We take BETA to be any BETA-like value (including Effect Size). If this is not the case for your sumstats, change this to FALSE. Default is TRUE.
- 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.
- ignore_multi_trait** If you have multiple traits (p-values) in the study but you want to ignore these and instead use a standard named p-value, set to TRUE. By default is FALSE which will check for multi-traits.
- 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.
- chr_style** Chromosome naming style to use in the formatted summary statistics file ("NCBI", "UCSC", "dbSNP", or "Ensembl"). The NCBI and Ensembl styles both code chromosomes as 1-22, X, Y, MT; the UCSC style is chr1-chr22, chrX, chrY, chrM; and the dbSNP style is ch1-ch22, chX, chY, chMT. Default is Ensembl.
- rmv_chrPrefix** Is now deprecated, do not use. Use chr_style instead - chr_style = 'Ensembl' will give the same result as rmv_chrPrefix=TRUE used to give.
- rmv_chr** Chromosomes to exclude from the formatted summary statistics file. Use NULL if no filtering is necessary. Default is c("X", "Y", "MT") which removes all non-autosomal SNPs.
- on_ref_genome** Binary Should a check take place that all SNPs are on the reference genome by SNP ID. Default is TRUE.

- `infer_eff_direction` Binary Should a check take place to ensure the alleles match the effect direction? Default is TRUE.
- `eff_on_minor_alleles` Binary Should MungeSumstats assume that the effects are majoritively measured on the minor alleles? Default is FALSE as this is an assumption that won't be appropriate in all cases. However, the benefit is that if we know the majority of SNPs have their effects based on the minor alleles, we can catch cases where the allele columns have been mislabelled.
- `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-bi-allelic SNPs be removed. Default is TRUE.
- `flip_frq_as_biallelic` Binary Should non-bi-allelic SNPs frequency values be flipped as 1-p despite there being other alternative alleles? Default is FALSE but if set to TRUE, this allows non-bi-allelic SNPs to be kept despite needing flipping.
- `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.
- `drop_indels` Binary, should any indels found in the sumstats be dropped? These can not be checked against a reference dataset and will have the same

- RS ID and position as SNPs which can affect downstream analysis. Default is False.
- `drop_na_cols` A character vector of column names to be checked for missing values. Rows with missing values in any of these columns (if present in the dataset) will be dropped. If NULL, all columns will be checked for missing values. Default columns are SNP, chromosome, position, allele 1, allele2, effect columns (frequency, beta, Z-score, standard error, log odds, signed sumstats, odds ratio), p value and N columns.
- `dbSNP` version of dbSNP to be used for imputation (144 or 155).
- `check_dups` whether to check for duplicates - if formatting QTL datasets this should be set to FALSE otherwise keep as TRUE. Default is TRUE.
- `sort_coordinates` Whether to sort by coordinates of resulting sumstats
- `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.
- `return_data` Return data.table, GRanges or VRanges directly to user. Otherwise, return the path to the save data. Default is FALSE.
- `return_format` If `return_data` is TRUE. Object type to be returned ("data.table", "vranges", "granges").
- `ldsc_format` DEPRECATED, do not use. Use `save_format="LDSC"` instead.
- `save_format` Output format of sumstats. Options are NULL - standardised output format from MungeSumstats, LDSC - output format compatible with LDSC and openGWAS - output compatible with openGWAS VCFs. Default is NULL. **NOTE** - If LDSC format is used, the naming convention of A1 as the reference (genome build) allele and A2 as the effect allele will be reversed to match LDSC (A1 will now be the effect allele). See more info on this [here](#). Note that any effect columns (e.g. Z) will be in relation to A1 now instead of A2.
- `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 .tsv.gz. 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. Note the name of the log files (log messages and log outputs) are now the same as the name of the file specified in the save path parameter with the extension '_log_msg.txt' and '_log_output.txt' respectively.
- `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.
- `mapping_file` MungeSumstats has a pre-defined column-name mapping file which should cover the most common column headers and their interpreta-

tions. 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

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,
  overwrite = TRUE,
  remove_tmp = TRUE,
  verbose = TRUE
)
```

Arguments

path	Path to GWAS summary statistics file.
chrom_col	Name of the chromosome column in sumstats_dt (e.g. "CHR").
start_col	Name of the starting genomic position column in sumstats_dt (e.g. "POS", "start").
end_col	Name of the ending genomic position column in sumstats_dt (e.g. "POS", "end"). Can be the same as start_col when sumstats_dt only contains SNPs that span 1 base pair (bp) each.
overwrite	A logical(1) indicating whether dest should be over-written, if it already exists.
remove_tmp	Remove the temporary uncompressed version of the file (.tsv).
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()
path <- tempfile(fileext = ".tsv")
MungeSumstats::write_sumstats(sumstats_dt = sumstats_dt, save_path = path)
indexed_file <- MungeSumstats::index_tabular(path = path)
```

index_vcf	<i>Tabix-index file: VCF</i>
-----------	------------------------------

Description

Convert summary stats file to tabix format

Usage

```
index_vcf(path, verbose = TRUE)
```

Arguments

path	Path to VCF.
verbose	Print messages.

Value

Path to tabix-indexed tabular file

Source

Borrowed function from [echotabix](#).

See Also

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

Examples

```
eduAttainOkbayPth <- system.file("extdata", "eduAttainOkbay.txt",
                                package = "MungeSumstats")
sumstats_dt <- data.table::fread(eduAttainOkbayPth, nThread = 1)
sumstats_dt <-
MungeSumstats::standardise_sumstats_column_headers_crossplatform(
  sumstats_dt = sumstats_dt)$sumstats_dt
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)
```

infer_effect_column *Infer if effect relates to A1 or A2 if ambiguously named*

Description

Three checks are made to infer which allele the effect/frequency information relates to if they are ambiguous (named A0, A1 and A2 or equivalent):

1. Check if ambiguous naming conventions are used (i.e. allele 0, 1 and 2 or equivalent). If not exit, otherwise continue to next checks. This can be checked by using the mapping file and splitting A1/A2 mappings by those that contain 0, 1 or 2 (ambiguous) or doesn't contain 0, 1 or 2 e.g. effect, tested (unambiguous so fine for MSS to handle as is).
2. Look for effect column/frequency column where the A0/A1/A2 explicitly mentioned, if found then we know the direction and should update A0/A1/A2 naming so A2 is the effect column. We can look for such columns by getting every combination of A0/A1/A2 naming and effect/frq naming.
3. If not found in 2, a final check should be against the reference genome, whichever of A0, A1 and A2 has more of a match with the reference genome should be taken as **not** the effect allele. There is an assumption in this but is still better than guessing the ambiguous allele naming.

Usage

```
infer_effect_column(
  sumstats_dt,
  dbSNP = 155,
  sampled_snps = 10000,
  mapping_file = sumstatsColHeaders,
  nThread = nThread,
  ref_genome = NULL,
  on_ref_genome = TRUE,
  infer_eff_direction = TRUE,
  eff_on_minor_alleles = FALSE,
  return_list = TRUE
)
```

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS.
dbSNP	version of dbSNP to be used for imputation (144 or 155).
sampled_snps	Downsample the number of SNPs used when inferring genome build to save time.
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.
nThread	Number of threads to use for parallel processes.
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.
on_ref_genome	Binary Should a check take place that all SNPs are on the reference genome by SNP ID. Default is TRUE.
infer_eff_direction	Binary Should a check take place to ensure the alleles match the effect direction? Default is TRUE.
eff_on_minor_alleles	Binary Should MungeSumstats assume that the effects are majoritively measured on the minor alleles? Default is FALSE as this is an assumption that won't be appropriate in all cases. However, the benefit is that if we know the majority of SNPs have their effects based on the minor alleles, we can catch cases where the allele columns have been mislabelled.
return_list	Return the sumstats_dt within a named list (default: TRUE).

Details

Also, if eff_on_minor_alleles=TRUE, check 3 will be used in all cases. However, This assumes that the effects are majoritively measured on the minor alleles and should be used with caution as this

is an assumption that won't be appropriate in all cases. However, the benefit is that if we know the majority of SNPs have their effects based on the minor alleles, we can catch cases where the allele columns have been mislabelled. IF `eff_on_minor_alleles=TRUE`, checks 1 and 2 will be skipped.

Value

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

Examples

```
sumstats <- MungeSumstats::formatted_example()
#for speed, don't run on_ref_genome part of check (on_ref_genome = FALSE)
sumstats_dt2<-infer_effect_column(sumstats_dt=sumstats,on_ref_genome = FALSE)
```

<code>is_tabix</code>	<i>Is tabix</i>
-----------------------	-----------------

Description

Is a file bgz-compressed and tabix-indexed.

Usage

```
is_tabix(path)
```

Arguments

`path` Path to file.

Value

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

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

Description

Transfer genomic coordinates from one genome build to another.

Usage

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

Arguments

sumstats_dt	data table obj of the summary statistics file for the GWAS.
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).
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.
chain_source	chain file source used ("ucsc" as default, or "ensembl")
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.
chrom_col	Name of the chromosome column in sumstats_dt (e.g. "CHR").
start_col	Name of the starting genomic position column in sumstats_dt (e.g. "POS", "start").
end_col	Name of the ending genomic position column in sumstats_dt (e.g. "POS", "end"). Can be the same as start_col when sumstats_dt only contains SNPs that span 1 base pair (bp) each.
as_granges	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").
local_chain	Path to local chain file to use instead of downloading. Default of NULL i.e. no local file to use. NOTE if passing a local chain file make sure to specify the path to convert from and to the correct build like GRCh37 to GRCh38. We can not

sense check this for local files. The chain file can be submitted as a gz file (as downloaded from source) or unzipped.

verbose Print messages.

Value

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

Source

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

Examples

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

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

<code>list_sumstats</code>	<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

- | | |
|----------------------------|--|
| <code>save_dir</code> | Top-level directory to recursively search for summary statistics files within. |
| <code>pattern</code> | Regex pattern to search for files with. |
| <code>ids_from_file</code> | Try to extract dataset IDs from file names. If FALSE, will infer IDs from the directory names instead. |
| <code>verbose</code> | 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,
  dbSNP = c(144, 155),
  msg = NULL,
  chr_filt = 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)
dbSNP	version of dbSNP to be used (144 or 155)
msg	Optional name of the column missing from the dataset in question. Default is NULL
chr_filt	Internal for testing - filter reference genomes and sumstats to specific chromosomes for testing. Pass a list of chroms in format: c("1","2"). Default is NULL i.e. no filtering.

Value

data table of snpsById, filtered to SNPs of interest.

Source

```
sumstats_dt <- formatted_example()
rsids <- MungeSumstats::load_ref_genome_data(snps = sumstats_dt$SNP, ref_genome = "GRCh37", dbSNP=144)
```

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

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, dbSNP = c(144, 155), msg = NULL)
```

Arguments

ref_genome	name of the reference genome used for the GWAS (GRCh37 or GRCh38)
dbSNP	version of dbSNP to be used (144 or 155)
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",dbSNP=144)
```

logs_example	<i>Example logs file</i>
--------------	--------------------------

Description

Example logs file produced by [format_sumstats](#).

Usage

```
logs_example(read = FALSE)
```

Arguments

read	Whether to read the logs file into memory.
------	--

Value

Path to logs file.

Source

```
eduAttainOkbayPth <- system.file("extdata", "eduAttainOkbay.txt", package = "MungeSumstats")
sumstats_dt <- data.table::fread(eduAttainOkbayPth) ##### Introduce values that need
to be fixed ##### sumstats_dt$Pval[10:15] <- 5 sumstats_dt$Pval[20:22] <- -5 sumstats_dt$Pval[23:25]
<- "5e-324" ss_path <- tempfile() data.table::fwrite(sumstats_dt, ss_path) log_folder
<- tempdir() reformatted <- MungeSumstats::format_sumstats( path = ss_path, ref_genome
= "GRCh37", log_folder = log_folder, log_mungesumstats_msgs = TRUE, log_folder_ind =
TRUE,) file.copy(reformatted$log_files$MungeSumstats_log_msg, "inst/extdata",overwrite
= TRUE)
```

make_allele_upper	Ensure A1 and A2 are upper case
-------------------	---------------------------------

Description

Ensure A1 and A2 are upper case

Usage

```
make_allele_upper(sumstats_dt, log_files)
```

Arguments

log_files list of log file locations

Value

list containing sumstats_dt, the modified summary statistics data table object and the log file list

messenger	Print messages
-----------	----------------

Description

Print messages with option to silence.

Usage

```
messenger(..., v = TRUE)
```

Arguments

- ... Message input.
- v Whether to print messages.

Value

Null output.

message_parallel	<i>Send messages to console even from within parallel processes</i>
------------------	---

Description

Send messages to console even from within parallel processes

Usage

message_parallel(...)

Value

A message

parse_dropped_chrom	<i>Parse number of SNPs dropped due to being on chrom X, Y or MT</i>
---------------------	--

Description

Support function for [parse_logs](#).

Usage

parse_dropped_chrom(1)

Arguments

- 1 Lines of text from log file.

Value

Numeric

`parse_dropped_duplicates`*Parse number of SNPs dropped due to being duplicates*

Description

Support function for [parse_logs](#).

Usage

```
parse_dropped_duplicates(1)
```

Arguments

1 Lines of text from log file.

Value

Numeric

`parse_dropped_INFO`*Parse number of SNPs dropped due to being below the INFO threshold*

Description

Support function for [parse_logs](#).

Usage

```
parse_dropped_INFO(1)
```

Arguments

1 Lines of text from log file.

Value

Numeric

parse_dropped_nonA1A2	<i>Parse number of SNPs dropped due to not matching the ref genome A1 or A2</i>
-----------------------	---

Description

Support function for [parse_logs](#).

Usage

parse_dropped_nonA1A2(1)

Arguments

1 Lines of text from log file.

Value

Numeric

parse_dropped_nonBiallelic	<i>Parse number of SNPs dropped due to not being bi-allelic</i>
----------------------------	---

Description

Support function for [parse_logs](#).

Usage

parse_dropped_nonBiallelic(1)

Arguments

1 Lines of text from log file.

Value

Numeric

parse_dropped_nonRef	<i>Parse number of SNPs dropped due to being in the ref genome</i>
----------------------	--

Description

Support function for [parse_logs](#).

Usage

```
parse_dropped_nonRef(1)
```

Arguments

1	Lines of text from log file.
---	------------------------------

Value

Numeric

parse_flipped	<i>Parse number of SNPs flipped to align with the ref genome</i>
---------------	--

Description

Support function for [parse_logs](#).

Usage

```
parse_flipped(1)
```

Arguments

1	Lines of text from log file.
---	------------------------------

Value

Numeric

parse_genome_build	<i>Genome build inferred from the summary statistics</i>
--------------------	--

Description

Support function for [parse_logs](#).

Usage

parse_genome_build(1)

Arguments

1 Lines of text from log file.

Value

Character

parse_idStandard	<i>Standardised IEU MRC OpenGWAS ID</i>
------------------	---

Description

Support function for [parse_logs](#).

Usage

parse_idStandard(1)

Arguments

1 Lines of text from log file.

Value

Character

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

<code>save_dir</code>	Top-level directory to recursively search for log files within.
<code>pattern</code>	Regex pattern to search for files with.
<code>verbose</code>	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)
```

parse_pval_large	<i>Parse number of SNPs with p-values >1</i>
------------------	---

Description

Support function for [parse_logs](#).

Usage

```
parse_pval_large(1)
```

Arguments

1	Lines of text from log file.
---	------------------------------

Value

Numeric

parse_pval_neg	<i>Parse number of SNPs with p-values <0</i>
----------------	---

Description

Support function for [parse_logs](#).

Usage

```
parse_pval_neg(1)
```

Arguments

1	Lines of text from log file.
---	------------------------------

Value

Numeric

parse_pval_small	<i>Parse number of SNPs with non-negative p-values $\leq 5e-324$</i>
------------------	---

Description

Support function for [parse_logs](#).

Usage

```
parse_pval_small(1)
```

Arguments

1	Lines of text from log file.
---	------------------------------

Value

Numeric

parse_report	<i>Parse "Summary statistics report" metrics</i>
--------------	--

Description

Support function for [parse_logs](#).

Usage

```
parse_report(1, entry = 1, line = 1)
```

Arguments

1 Lines of text from log file.

Value

Numeric

parse_snps_freq_05	<i>Parse number/percent of SNPs with <i>FREQ</i> values >0.5</i>
--------------------	---

Description

Support function for [parse_logs](#).

Usage

```
parse_snps_freq_05(1, percent = FALSE)
```

Arguments

1 Lines of text from log file.

Value

Numeric

`parse_snps_not_formatted`*Parse number of SNPs not correctly formatted*

Description

Support function for [parse_logs](#).

Usage

```
parse_snps_not_formatted(1)
```

Arguments

1 Lines of text from log file.

Value

Numeric

`parse_time`*Parse the total time taken the munge the file*

Description

Support function for [parse_logs](#).

Usage

```
parse_time(1)
```

Arguments

1 Lines of text from log file.

Value

Character

preview_sumstats	<i>Preview formatted sum stats saved to disk</i>
------------------	--

Description

Prints the first n lines of the sum stats.

Usage

```
preview_sumstats(save_path, nrows = 5L)
```

Arguments

save_path File path to save formatted data. Defaults to tempfile(fileext=".tsv.gz").

Value

No return

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/ALScvf.vcf.gz")

raw_eduAttainOkbay	GWAS Educational Attainment Okbay 2016 - Subset
--------------------	---

Description

GWAS Summary Statistics on Educational Attainment by Okbay et al 2016: PMID: 27898078
PMCID: 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 subsetted 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 arebi/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(tm
#DT read removes the .0's #remove those not on ref genome and withbi/tri allelic rmv <-
c("rs192818565","rs79925071","rs1606974","rs1871109", "rs73074378","rs7955289") eduAttainOkbay
<- eduAttainOkbay[!MarkerName data.table::fwrite(eduAttainOkbay,file=tmp,sep="\t")
eduAttainOkbay <- readLines(tmp) writeLines(eduAttainOkbay,"inst/extdata/eduAttainOkbay.txt")

read_header	Read in file header
-------------	---------------------

Description

Read in file header

Usage

read_header(path, n = 2L, skip_vcf_metadata = FALSE, nThread = 1)

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.
n	integer. The (maximal) number of lines to read. Negative values indicate that one should read up to the end of input on the connection.
skip_vcf_metadata	logical, should VCF metadata be ignored
nThread	Number of threads to use for parallel processes.

Value

First n lines of the VCF header

Examples

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

read_log_pval	<i>Read -log10 p-value column</i>
---------------	-----------------------------------

Description

Parse p-value column in VCF file.of other general -log10 p-values

Usage

```
read_log_pval(
  sumstats_dt,
  mapping_file = sumstatsColHeaders,
  return_list = TRUE
)
```

Arguments

sumstats_dt	Summary stats data.table.
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.
return_list	Binary, whether to return the dt in a list or not - list is standard for the format_sumstats() function.

Value

Null output.

read_sumstats	<i>Determine summary statistics file type and read them into memory</i>
---------------	---

Description

Determine summary statistics file type and read them into memory

Usage

```
read_sumstats(
  path,
  nrows = Inf,
  standardise_headers = FALSE,
  samples = 1,
  sampled_rows = 10000L,
  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: • 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).
sampled_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.
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,
  as_datatable = TRUE,
  save_path = NULL,
  tabix_index = FALSE,
  samples = 1,
  which = NULL,
  use_params = TRUE,
  sampled_rows = 10000L,
  download = TRUE,
  vcf_dir = tempdir(),
  download_method = "download.file",
  force_new = FALSE,
  mt_thresh = 100000L,
  nThread = 1,
  verbose = TRUE
)
```

Arguments

- | | |
|--------------|--|
| path | Path to local or remote VCF file. |
| as_datatable | Return the data as a data.table (default: TRUE) or a VCF (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	Genomic ranges to be added if supplied. Default is NULL.
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.
download	Download the VCF (and its index file) to a temp folder before reading it into R. This is important to keep TRUE when nThread>1 to avoid making too many queries to remote file.
vcf_dir	Where to download the original VCF from Open GWAS. <i>WARNING:</i> This is set to tempdir() 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. vcf_dir=". /raw_vcf").
download_method	"axel" (multi-threaded) or "download.file" (single-threaded) .
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.
mt_thresh	When the number of rows (variants) in the VCF is < mt_thresh, only use single-threading for reading in the VCF. This is because the overhead of parallelisation outweighs the speed benefits when VCFs are small.
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.
vcf <- VariantAnnotation::readVcf(file = path) N <- 1e5 vcf_sub <- vcf[1:N,] res <- microbenchmark::microb
"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](#)

[Discussion on VariantAnnotation GitHub](#)

Examples

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

#### Remote file ####
## Small GWAS (0.2Mb)
# path <- "https://gwas.mrcieu.ac.uk/files/ieu-a-298/ieu-a-298.vcf.gz"
# sumstats_dt2 <- read_vcf(path = path)

## Large GWAS (250Mb)
# path <- "https://gwas.mrcieu.ac.uk/files/ubm-a-2929/ubm-a-2929.vcf.gz"
# sumstats_dt3 <- read_vcf(path = path, nThread=11)

### Very large GWAS (500Mb)
# path <- "https://gwas.mrcieu.ac.uk/files/ieu-a-1124/ieu-a-1124.vcf.gz"
# sumstats_dt4 <- read_vcf(path = path, nThread=11)
```

read_vcf_genome	<i>Read VCF genome</i>
-----------------	------------------------

Description

Get the genome build of a remote or local VCF file.

Usage

```
read_vcf_genome(
  header = NULL,
  validate = FALSE,
  default_genome = "HG19/GRCh37",
  verbose = TRUE
)
```

Arguments

header	Header extracted by scanVcfHeader .
validate	Validate genome name using mapGenomeBuilds .
default_genome	When no genome can be extracted, default to this genome build.
verbose	Print messages.

Value

genome

<code>read_vcf_info</code>	<i>Read VCF: INFO column</i>
----------------------------	------------------------------

Description

Parse INFO column in VCF file.

Usage

`read_vcf_info(sumstats_dt)`

Arguments

`sumstats_dt` Summary stats data.table.

Value

Null output.

<code>read_vcf_markername</code>	<i>Read VCF: MarkerName column</i>
----------------------------------	------------------------------------

Description

Parse MarkerName/SNP column in VCF file.

Usage

`read_vcf_markername(sumstats_dt)`

Arguments

`sumstats_dt` Summary stats data.table.

Value

Null output.

read_vcf_parallel	<i>Read VCF: parallel</i>
-------------------	---------------------------

Description

Read a VCF file across 1 or more threads in parallel. If tilewidth is not specified, the size of each chunk will be determined by total genome size divided by ntile. By default, ntile is equal to the number of threads, nThread. For further discussion on how this function was optimised, see [here](#) and [here](#).

Usage

```
read_vcf_parallel(
  path,
  samples = 1,
  which = NULL,
  use_params = TRUE,
  as_datatable = TRUE,
  sampled_rows = 10000L,
  include_xy = FALSE,
  download = TRUE,
  vcf_dir = tempdir(),
  download_method = "download.file",
  force_new = FALSE,
  tilewidth = NULL,
  mt_thresh = 100000L,
  nThread = 1,
  ntile = nThread,
  verbose = TRUE
)
```

Arguments

path	Path to local or remote VCF file.
samples	Which samples to use: • 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	Genomic ranges to be added if supplied. Default is NULL.
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.
as_datatable	Return the data as a data.table (default: TRUE) or a VCF (FALSE).

sampled_rows	First N rows to sample. Set NULL to use full sumstats_file. when determining whether cols are empty.
download	Download the VCF (and its index file) to a temp folder before reading it into R. This is important to keep TRUE when nThread>1 to avoid making too many queries to remote file.
vcf_dir	Where to download the original VCF from Open GWAS. <i>WARNING:</i> This is set to tempdir() 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. vcf_dir="/raw_vcf").
download_method	"axel" (multi-threaded) or "download.file" (single-threaded) .
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.
tilewidth	The desired tile width. The effective tile width might be slightly different but is guaranteed to never be more than the desired width.
mt_thresh	When the number of rows (variants) in the VCF is < mt_thresh, only use single-threading for reading in the VCF. This is because the overhead of parallelisation outweighs the speed benefits when VCFs are small.
nThread	Number of threads to use for parallel processes.
ntile	The number of tiles to generate.
verbose	Print messages.

Value

VCF file.

Source

```
path <- "https://gwas.mrcieu.ac.uk/files/ieu-a-298/ieu-a-298.vcf.gz" ##### Single-threaded
#### vcf <- MungeSumstats:::read_vcf_parallel(path = path) ##### Parallel ##### vcf2 <-
MungeSumstats:::read_vcf_parallel(path = path, nThread=11)
```

register_cores	Register cores
----------------	----------------

Description

Register a multi-threaded instances using **BiocParallel**.

Usage

```
register_cores(workers = 1, progressbar = TRUE)
```

Arguments

- `workers` `integer(1)` Number of workers. Defaults to the maximum of 1 or the number of cores determined by `detectCores` minus 2 unless environment variables `R_PARALLELLY_AVAILABLECORES_FALLBACK` or `BIOCPARALLEL_WORKER_NUMBER` are set otherwise. For a SOCK cluster, `workers` can be a `character()` vector of host names.
- `progressbar` `logical(1)` Enable progress bar (based on `plyr::progress_text`).

Value

Null output.

<code>remove_empty_cols</code>	<i>Remove empty columns</i>
--------------------------------	-----------------------------

Description

Remove columns that are empty or contain all the same values in a `data.table`.

Usage

```
remove_empty_cols(sumstats_dt, sampled_rows = NULL, verbose = TRUE)
```

Arguments

- `sampled_rows` First N rows to sample. Set `NULL` to use full `sumstats_file`. when determining whether cols are empty.
- `verbose` Print messages.

Value

Null output.

<code>report_summary</code>	<i>Report info on current state of the summary statistics</i>
-----------------------------	---

Description

Prints report.

Usage

```
report_summary(sumstats_dt, orig_dims = NULL)
```

Arguments

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

Value

No return

select_vcf_fields	<i>Select VCF fields</i>
-------------------	--------------------------

Description

Select non-empty columns from each VCF field type.

Usage

```
select_vcf_fields(  
  path,  
  sampled_rows = 10000L,  
  which = NULL,  
  samples = NULL,  
  nThread = 1,  
  verbose = TRUE  
)
```

Arguments

path Path to local or remote VCF file.

sampled_rows First N rows to sample. Set NULL to use full sumstats_file. when determining whether cols are empty.

which Genomic ranges to be added if supplied. Default is NULL.

samples Which samples to use:

- 1 : Only the first sample will be used (*DEFAULT*).
- NULL : All samples will be used.
- c("<sample_id1>","<sample_id2>",...) : Only user-selected samples will be used (case-insensitive).

nThread Number of threads to use for parallel processes.

verbose Print messages.

Value

ScanVcfParam object.

sort_coords	<i>Sort sum stats</i>
-------------	-----------------------

Description

Sort summary statistics table by genomic coordinates.

Usage

```
sort_coords(  
  sumstats_dt,  
  sort_coordinates = TRUE,  
  sort_method = c("data.table", "GenomicRanges")  
)
```

Arguments

- | | |
|-------------|---|
| sumstats_dt | data.table obj of the summary statistics file for the GWAS. |
| sort_method | Method to sort coordinates by: <ul style="list-style-type: none">• "data.table" (default) Uses setorderv, which is much faster than "GenomicRanges" but less robust to variations in some sum stats files.• "GenomicRanges" Uses sort.GenomicRanges, which is more robust to variations in sum stats files but much slower than the "data.table" method. |
| sort_coords | Whether to sort by coordinates. |

Value

Sorted sumstats_dt

sort_coords_datatable	<i>Sort sum stats: data.table</i>
-----------------------	-----------------------------------

Description

Sort summary statistics table by genomic coordinates using a fast data.table-native strategy

Usage

```
sort_coords_datatable(  
  sumstats_dt,  
  chr_col = "CHR",  
  start_col = "BP",  
  end_col = start_col  
)
```

Arguments

- sumstats_dt [data.table](#) obj of the summary statistics file for the GWAS.
- chr_col Chromosome column name.
- start_col Genomic end position column name.

Value

Sorted sumstats_dt

sort_coord_genomicranges
<i>Sort sum stats: GenomicRanges</i>

Description

Sort summary statistics table by genomic coordinates using a slower (but in some cases more robust) GenomicRanges strategy

Usage

```
sort_coord_genomicranges(sumstats_dt)
```

Arguments

- sumstats_dt [data.table](#) obj of the summary statistics file for the GWAS.

Value

Sorted sumstats_dt

standardise_header	<i>Standardise the column headers in the Summary Statistics files</i>
--------------------	---

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,  
  convert_A0 = 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 from 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).
- convert_A0 Whether to convert A* (representing A0) to A1/A2. This should be done unless checking if A0 was present in the input as if you do it you can't infer this. Default is TRUE
- 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", "eduAttainOkbay.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 new A1 and A2 naming

```
a1_name <- c("NON", "RISK", "ALLELE") a2_name <- c("RISK", "ALLELE") all_delims <- c("_", ".", "", "
", "-") all_uncorr_a1 <- vector(mode="list", length = length(all_delims)) all_corr_a1
<- vector(mode="list", length = length(all_delims)) all_uncorr_a2 <- vector(mode="list", length
= length(all_delims)) all_corr_a2 <- vector(mode="list", length = length(all_delims))
for(i in seq_along(all_delims)){ delim <- all_delims[i] a1 <- unlist(paste(a1_name, collapse=delim))
a2 <- unlist(paste(a2_name, collapse=delim)) all_uncorr_a1[[i]] <- a1 all_uncorr_a2[[i]]
<- a2 all_corr_a1[[i]] <- "A1" all_corr_a2[[i]] <- "A2" } se_cols <- data.frame("Uncorrected"=c(unlist(a1
"Corrected"=c(unlist(all_corr_a1), unlist(all_corr_a2))) # Or another example ....
# shown is an example of adding columns for Standard Error (SE) se_cols <- data.frame("Uncorrected"=c("SE"
"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$Corrected
sumstatsColHeaders <- sumstatsColHeaders[order(sumstatsColHeaders$Corrected, sumstatsColHeaders$order
= TRUE), ] rownames(sumstatsColHeaders)<-1:nrow(sumstatsColHeaders) sumstatsColHeaders$ordering
<- NULL #manually move FREQUENCY to above MAR - github issue 95 frequency <- sumstatsColHeaders[sumstatsCo
maf <- sumstatsColHeaders[sumstatsColHeaders$Uncorrected=="MAF", ] if(as.integer(rownames(frequency))>
sumstatsColHeaders[as.integer(rownames(frequency)), ] <- maf sumstatsColHeaders[as.integer(rownames(ma
<- 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
```

supported_suffixes *List supported file formats*

Description

List supported file formats

Usage

```
supported_suffixes(
  tabular = TRUE,
  tabular_compressed = TRUE,
  vcf = TRUE,
```

```
vcf_compressed = TRUE
)
```

Arguments

- tabular Include tabular formats.
- tabular_compressed Include compressed tabular formats.
- vcf Include Variant Call Format.
- vcf_compressed Include compressed Variant Call Format.

Value

File formats

to_granges	To GRanges
------------	------------

Description

Convert a [data.table](#) to [GRanges](#).

Usage

```
to_granges(
  sumstats_dt,
  seqnames.field = "CHR",
  start.field = "BP",
  end.field = "BP",
  style = c("NCBI", "UCSC")
)
```

Arguments

- sumstats_dt data table obj of the summary statistics file for the GWAS.
- seqnames.field A character vector of recognized names for the column in df that contains the chromosome name (a.k.a. sequence name) associated with each genomic range. Only the first name in seqnames.field that is found in colnames(df) is used. If no one is found, then an error is raised.
- start.field A character vector of recognized names for the column in df that contains the start positions of the genomic ranges. Only the first name in start.field that is found in colnames(df) is used. If no one is found, then an error is raised.
- end.field A character vector of recognized names for the column in df that contains the end positions of the genomic ranges. Only the first name in start.field that is found in colnames(df) is used. If no one is found, then an error is raised.
- style GRanges style to convert to, "NCBI" or "UCSC".

Value

GRanges object

to_vranges	<i>Convert to VRanges</i>
------------	---------------------------

Description

Convert to VRanges

Usage

```
to_vranges(sumstats_dt)
```

Arguments

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

Value

VRanges object

unlist_dt	<i>Unlist a data.table</i>
-----------	----------------------------

Description

Identify columns that are lists and turn them into vectors.

Usage

```
unlist_dt(dt, verbose = TRUE)
```

Arguments

dt data.table
verbose Print messages.

Value

dt with list columns turned into vectors.

validate_parameters	<i>Ensure that the input parameters are logical</i>
---------------------	---

Description

Ensure that the input parameters are logical

Usage

```
validate_parameters(  
  path,  
  ref_genome,  
  convert_ref_genome,  
  convert_small_p,  
  es_is_beta,  
  compute_z,  
  compute_n,  
  convert_n_int,  
  analysis_trait,  
  INFO_filter,  
  FRQ_filter,  
  pos_se,  
  effect_columns_nonzero,  
  N_std,  
  N_dropNA,  
  chr_style,  
  rmv_chr,  
  on_ref_genome,  
  infer_eff_direction,  
  eff_on_minor_alleles,  
  strand_ambig_filter,  
  allele_flip_check,  
  allele_flip_drop,  
  allele_flip_z,  
  allele_flip_frq,  
  bi_allelic_filter,  
  flip_frq_as_biallelic,  
  snp_ids_are_rs_ids,  
  remove_multi_rs_snp,  
  frq_is_maf,  
  indels,  
  drop_indels,  
  check_dups,  
  dbSNP,  
  write_vcf,  
  return_format,  
  ldsc_format,
```

```

    save_format,
    imputation_ind,
    log_folder_ind,
    log_mungesumstats_msgs,
    mapping_file,
    tabix_index,
    chain_source,
    local_chain,
    drop_na_cols,
    rmv_chrPrefix
)

```

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.
es_is_beta	Binary, whether to map ES to BETA. We take BETA to be any BETA-like value (including Effect Size). If this is not the case for your sumstats, change this to FALSE. Default is TRUE.
compute_z	Whether to compute Z-score column. Default is FALSE. This can be computed from Beta and SE with (Beta/SE) or P ($Z := \text{sign}(\text{BETA}) * \sqrt{\text{stats::qchisq}(P, 1, \text{lower} = \text{FALSE})}$). Note that imputing the Z-score from P for every SNP will not be perfectly correct and may result in a loss of power. This should only be done as a last resort. Use 'BETA' to impute by BETA/SE and 'P' to impute by SNP p-value.
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.

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.
chr_style	Chromosome naming style to use in the formatted summary statistics file ("NCBI", "UCSC", "dbSNP", or "Ensembl"). The NCBI and Ensembl styles both code chromosomes as 1-22, X, Y, MT; the UCSC style is chr1-chr22, chrX, chrY, chrM; and the dbSNP style is ch1-ch22, chX, chY, chMT. Default is Ensembl.
rmv_chr	Chromosomes to exclude from the formatted summary statistics file. Use NULL if no filtering is necessary. Default is c("X", "Y", "MT") which removes all non-autosomal SNPs.
on_ref_genome	Binary Should a check take place that all SNPs are on the reference genome by SNP ID. Default is TRUE.
infer_eff_direction	Binary Should a check take place to ensure the alleles match the effect direction? Default is TRUE.
eff_on_minor_alleles	Binary Should MungeSumstats assume that the effects are majoritively measured on the minor alleles? Default is FALSE as this is an assumption that won't be appropriate in all cases. However, the benefit is that if we know the majority of SNPs have their effects based on the minor alleles, we can catch cases where the allele columns have been mislabelled.
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-bi-allelic SNPs be removed. Default is TRUE.
flip_frq_as_biallelic	Binary Should non-bi-allelic SNPs frequency values be flipped as 1-p despite there being other alternative alleles? Default is FALSE but if set to TRUE, this allows non-bi-allelic SNPs to be kept despite needing flipping.
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.
drop_indels	Binary, should any indels found in the sumstats be dropped? These can not be checked against a reference dataset and will have the same RS ID and position as SNPs which can affect downstream analysis. Default is False.
check_dups	whether to check for duplicates - if formatting QTL datasets this should be set to FALSE otherwise keep as TRUE. Default is TRUE.
dbSNP	version of dbSNP to be used for imputation (144 or 155).
write_vcf	Whether to write as VCF (TRUE) or tabular file (FALSE).
return_format	If return_data is TRUE. Object type to be returned ("data.table", "vranges", "granges").
ldsc_format	DEPRECATED, do not use. Use save_format="LDSC" instead.
save_format	Output format of sumstats. Options are NULL - standardised output format from MungeSumstats, LDSC - output format compatible with LDSC and openGWAS - output compatible with openGWAS VCFs. Default is NULL. NOTE - If LDSC format is used, the naming convention of A1 as the reference (genome build) allele and A2 as the effect allele will be reversed to match LDSC (A1 will now be the effect allele). See more info on this here . Note that any effect columns (e.g. Z) will be in relation to A1 now instead of A2.
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.
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 .tsv.gz. 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
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.
tabix_index	Index the formatted summary statistics with tabix for fast querying.
chain_source	source of the chain file to use in liftover, if converting genome build ("ucsc" or "ensembl"). Note that the UCSC chain files require a license for commercial use. The Ensembl chain is used by default ("ensembl").
local_chain	Path to local chain file to use instead of downloading. Default of NULL i.e. no local file to use. NOTE if passing a local chain file make sure to specify the path to convert from and to the correct build like GRCh37 to GRCh38. We can not sense check this for local files. The chain file can be submitted as a gz file (as downloaded from source) or unzipped.
drop_na_cols	A character vector of column names to be checked for missing values. Rows with missing values in any of these columns (if present in the dataset) will be dropped. If NULL, all columns will be checked for missing values. Default columns are SNP, chromosome, position, allele 1, allele2, effect columns (frequency, beta, Z-score, standard error, log odds, signed sumstats, odds ratio), p value and N columns.
rmv_chrPrefix	Is now deprecated, do. not use. Use chr_style instead - chr_style = 'Ensembl' will give the same result as rmv_chrPrefix=TRUE used to give.

Value

No return

vcf2df

*VCF to DF***Description**Function to convert a **VariantAnnotation** CollapsedVCF/ExpandedVCF object to a data.frame.

Usage

```
vcf2df(
  vcf,
  add_sample_names = TRUE,
  add_rowranges = TRUE,
  drop_empty_cols = TRUE,
  unique_cols = TRUE,
  unique_rows = TRUE,
  unlist_cols = TRUE,
  sampled_rows = NULL,
  verbose = TRUE
)
```

Arguments

<code>vcf</code>	Variant Call Format (VCF) file imported into R as a VariantAnnotation CollapsedVCF / ExpandedVCF object.
<code>add_sample_names</code>	Append sample names to column names (e.g. "EZ" -> "EZ_ubm-a-2929").
<code>add_rowranges</code>	Include rowRanges from VCF as well.
<code>drop_empty_cols</code>	Drop columns that are filled entirely with: NA, ".", or "".
<code>unique_cols</code>	Only keep uniquely named columns.
<code>unique_rows</code>	Only keep unique rows.
<code>unlist_cols</code>	If any columns are lists instead of vectors, unlist them. Required to be TRUE when <code>unique_rows=TRUE</code> .
<code>sampled_rows</code>	First N rows to sample. Set NULL to use full <code>sumstats_file</code> . when determining whether cols are empty.
<code>verbose</code>	Print messages.

Value

data.frame version of VCF

Source

[Original code source](#)

vcfR:

```
if(!require("pinfsc50")) install.packages("pinfsc50") vcf_file <- system.file("extdata", "pinf_sc50.vcf.gz",
package = "pinfsc50") vcf <- read.vcfR( vcf_file, verbose = FALSE ) vcf_df_list <- vcfR::vcfR2tidy(vcf,
single_frame=TRUE) vcf_df <- data.table::data.table(vcf_df_list$dat)
```

Examples

```
#### VariantAnnotation ####
# path <- "https://github.com/brentp/vcfanno/raw/master/example/exac.vcf.gz"
path <- system.file("extdata", "ALSvcf.vcf",
                    package = "MungeSumstats")

vcf <- VariantAnnotation::readVcf(file = path)
vcf_df <- MungeSumstats::vcf2df(vcf = vcf)
```

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

Description

Write sum stats file to disk

Usage

```
write_sumstats(
  sumstats_dt,
  save_path,
  ref_genome = NULL,
  sep = "\t",
  write_vcf = FALSE,
  save_format = NULL,
  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> .
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.
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 NULL or "" 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).

save_format	Output format of sumstats. Options are NULL - standardised output format from MungeSumstats, LDSC - output format compatible with LDSC and openGWAS - output compatible with openGWAS VCFs. Default is NULL. NOTE - If LDSC format is used, the naming convention of A1 as the reference (genome build) allele and A2 as the effect allele will be reversed to match LDSC (A1 will now be the effect allele). See more info on this here . Note that any effect columns (e.g. Z) will be in relation to A1 now instead of A2.
tabix_index	Index the formatted summary statistics with tabix 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 save_path. 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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