Package ‘quincunx’

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'class-releases.R' 'class-sample_sets.R' 'class-scores.R'
'class-trait_categories.R' 'class-trait_categories.R' 'clear_cache.R'
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'is_pubmed_id.R' 'messages.R' 'n_pages.R' 'nr_to_na.R'
'offsets.R' 'open_in_dbsnp.R' 'open_in_pgs_catalog.R'
'open_in_pubmed.R' 'parse-ancestry_categories.R'
'parse-cohorts.R' 'parse-performance_metrics.R'
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'parse-scores.R' 'parse-trait_categories.R' 'parse-trait_categories.R'
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'unwrap_demographics.R' 'unwrap_efotrait.R' 'unwrap_interval.R'
'unwrap_publication.R' 'unwrap_sample.R' 'utils-pipe.R'
'warnings.R' 'write_xlsx.R'

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Author Ramiro Magno [aut, cre] (<https://orcid.org/0000-0001-5226-3441>),
Isabel Duarte [aut] (<https://orcid.org/0000-0003-0060-2936>),
Ana-Teresa Maia [aut] (<https://orcid.org/0000-0002-0454-9207>),
CINTESIS [cph, fnd]

Maintainer Ramiro Magno <ramiro.magno@gmail.com>

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**Description**

A dataset containing the ancestry categories defined in NHGRI-EBI GWAS Catalog framework (Table 1, doi: 10.1186/s1305901813962). Ancestry categories are assigned to samples with distinct and well-defined patterns of genetic variation. You will find these categories in the variable `ancestry_category` of the following objects: `scores`, `performance_metrics` and `sample_sets`. Ancestry categories (`ancestry_category`) are further clustered into ancestry classes (`ancestry_class`).

**Usage**

`ancestry_categories`

**Format**

A data frame with 19 ancestry categories (rows) and 6 columns:

- **ancestry_category**  Ancestry category.
- **ancestry_class**  To reduce the complexity associated with the many ancestry categories, some have been merged into higher-level groupings (ancestry_class). These groupings represent the current breadth of data in the PGS Catalog and are likely to change as more data is added.
- **ancestry_class_symbol**  3-letter code for the ancestry_class e.g. "EUR" or "MAE".
- **ancestry_class_colour**  Hexadecimal colour code associated with ancestry groupings (ancestry_class). This can be useful when visually communicating about ancestries.
- **definition**  Description of the ancestry category.
- **examples**  Examples of detailed descriptions of sample ancestries included in the category.
Source

Table 1 of Moralles et al. (2018): doi: 10.1186/s1305901813962
PGS Catalog Ancestry Documentation: http://www.pgscatalog.org/docs/ancestry/

Examples

ancestry_categories

---

**bind**

*Bind PGS Catalog objects*

Description

bind() binds together PGS Catalog objects of the same class. Note that bind() preserves duplicates whereas union() does not.

Usage

bind(x, ...)

Arguments

- x: An object of either class scores, publications, traits, performance_metrics, sample_sets, cohorts or trait_categories.
- ...: Objects of the same class as x.

Value

An object of the same class as x.

Examples

```r
# Get some 'scores' objects:
my_scores_1 <- get_scores(c('PGS000012', 'PGS000013'))
my_scores_2 <- get_scores(c('PGS000013', 'PGS000014'))

# NB: with 'bind()', PGS000013 is repeated (as opposed to 'union()')
bind(my_scores_1, my_scores_2)@scores
```
**clear_cache**

Clear quincunx cache of memoised functions

**Description**

quincunx uses memoised functions for the REST API calls. Use this function to reset the cache.

**Usage**

clear_cache()

**Value**

Returns a logical value, indicating whether the resetting of the cache was successful (TRUE) or not FALSE.

**Examples**

clear_cache()

---

**cohorts-class**

An S4 class to represent a set of cohorts

**Description**

The cohorts object consists of two tables (slots) that combined form a relational database of a subset of cohorts. Each cohort is an observation (row) in the cohorts table (first table).

**Slots**

cohorts A table of cohorts. Each cohort (row) is identified by its cohort_symbol. Columns:

- **cohort_symbol** Cohort symbol. Example: "CECILE".
- **cohort_name** Cohort full name. Example: "CECILE Breast Cancer Study".

pgs_ids A table of cohorts and their associated polygenic scores identifiers. Columns:

- **cohort_symbol** Cohort symbol. Example: "CECILE".
- **pgs_id** Polygenic Score (PGS) identifier.
- **stage** Sample stage: either "gwas/dev" or "eval".
get_ancestry_categories

Get ancestry categories and classes

Description

Retrieves ancestry categories and classes. This function simply returns the object `ancestry_categories`.

Usage

generate_ancestry_categories()

Value

A tibble with ancestry categories, classes and associated information. See `ancestry_categories` for details about each column.

Examples

generate_ancestry_categories()

get_cohorts

Get PGS Catalog Cohorts

Description

Retrieves cohorts via the PGS Catalog REST API. Please note that all `cohort_symbol` is vectorised, thus allowing for batch mode search.

Usage

generate_cohorts(
    cohort_symbol = NULL,
    verbose = FALSE,
    warnings = TRUE,
    progress_bar = TRUE
)

Arguments

cohort_symbol A cohort symbol or NULL if all cohorts are intended.
verbose A logical indicating whether the function should be verbose about the different queries or not.
warnings A logical indicating whether to print warnings, if any.
progress_bar Whether to show a progress bar indicating download progress from the REST API server.
get_performance_metrics

Value

A cohorts object.

Examples

# Get information about specific cohorts by their symbols (acronyms)
get_cohorts(cohort_symbol = c('23andMe', 'IPOBCS'))

# Get info on all cohorts (may take a few minutes to download)
## Not run:
get_cohorts()
## End(Not run)

get_performance_metrics

Get PGS Catalog Performance Metrics

Description

Retrieves performance metrics via the PGS Catalog REST API. The REST API is queried multiple times with the criteria passed as arguments (see below). By default all performance metrics that match the criteria supplied in the arguments are retrieved: this corresponds to the default option set_operation set to 'union'. If you rather have only the associations that match simultaneously all criteria provided, then set set_operation to 'intersection'.

Usage

get_performance_metrics(
  ppm_id = NULL,
  pgs_id = NULL,
  set_operation = "union",
  interactive = TRUE,
  verbose = FALSE,
  warnings = TRUE,
  progress_bar = TRUE
)

Arguments

ppm_id A character vector of PGS Catalog performance metrics accession identifiers.
pgs_id A character vector of PGS Catalog score accession identifiers.
set_operation Either 'union' or 'intersection'. This tells how performance metrics retrieved by different criteria should be combined: 'union' binds together all results removing duplicates and 'intersection' only keeps same performance metrics found with different criteria.
**get_publications**

- **interactive**: A logical. If all performance metrics are requested, whether to ask interactively if we really want to proceed.
- **verbose**: A logical indicating whether the function should be verbose about the different queries or not.
- **warnings**: A logical indicating whether to print warnings, if any.
- **progress_bar**: Whether to show a progress bar as the queries are performed.

**Details**

Please note that all search criteria are vectorised, thus allowing for batch mode search.

**Value**

A `performance_metrics` object.

**Examples**

```r
# Get performance metrics catalogued with identifier 'PPM000001'
get_performance_metrics(ppm_id = 'PPM000001')

# Get performance metrics associated with polygenic score id 'PGS000001'
get_performance_metrics(pgs_id = 'PGS000001')

# To retrieve all catalogued performed metrics in PGS Catalog you simply
# leave the parameters `ppm_id` and `pgs_id` as `NULL`.
# Not run:
get_performance_metrics()
# End(Not run)
```

---

**get_publications**  
*Get PGS Catalog Publications*

**Description**

Retrieves PGS publications via the PGS Catalog REST API. The REST API is queried multiple times with the criteria passed as arguments (see below). By default all publications that match the criteria supplied in the arguments are retrieved: this corresponds to the default option `set_operation` set to 'union'. If you rather have only the associations that match simultaneously all criteria provided, then set `set_operation` to 'intersection'.
Usage

get_publications(
  pgp_id = NULL,
  pgs_id = NULL,
  pubmed_id = NULL,
  author = NULL,
  set_operation = "union",
  interactive = TRUE,
  verbose = FALSE,
  warnings = TRUE,
  progress_bar = TRUE
)

Arguments

pgp_id A character vector of PGS Catalog publication accession identifiers.
pgs_id A character vector of PGS Catalog score accession identifiers.
pubmed_id An integer vector of PubMed identifiers.
author A character vector of author names, any author in the list of authors in a publication, e.g. 'Mavaddat'.
set_operation Either 'union' or 'intersection'. This tells how publications retrieved by different criteria should be combined: 'union' binds together all results removing duplicates and 'intersection' only keeps same publications found with different criteria.
interactive A logical. If all publications are requested, whether to ask interactively if we really want to proceed.
verbose A logical indicating whether the function should be verbose about the different queries or not.
warnings A logical indicating whether to print warnings, if any.
progress_bar Whether to show a progress bar as the queries are performed.

Details

Please note that all search criteria are vectorised, thus allowing for batch mode search. For more details see the help vignette: vignette("getting-pgs-publications",package = "quincunx").

Value

A publications object.

Examples

# Get PGS publications by their identifier
get_publications(pgp_id = c('PGP000001', 'PGP000002'))

# By polygenic score identifier
get_publications(pgs_id = 'PGS000003')
# By PubMed identifier
get_publications(pubmed_id = '30554720')

# By author's last name
get_publications(author = 'Natarajan')

get_releases
Get PGS Catalog Releases

Description
This function retrieves PGS Catalog release information. Note that the columns pgs_id, ppm_id
and pgp_id contain in each element a vector. These columns can be unnested using unnest_longer
(see Examples).

Usage
get_releases(
  date = "latest",
  verbose = FALSE,
  warnings = TRUE,
  progress_bar = TRUE
)

Arguments
date One or more dates formatted as "YYYY-MM-DD" for respective releases, "latest"
for the latest release, or "all" for all releases.
verbose Whether to print information about the underlying requests to the REST API
server.
warnings Whether to print warnings about the underlying requests to the REST API server.
progress_bar Whether to show a progress bar indicating download progress from the REST
API server.

Value
A data frame where each row is a release. Columns are:

  date  Release date.
  n_pgs Number of released Polygenic Score (PGS) identifiers (pgs_id).
  n_ppm Number of released Performance Metric (PPM) identifiers (ppm_id).
  n_pgp Number of released PGS Catalog Publication (PGP) identifiers (pgp_id).
  pgs_id Released Polygenic Score (PGS) identifiers.
  ppm_id Released Performance Metric (PPM) identifiers.
  pgp_id Released PGS Catalog Publication (PGP) identifiers.
  notes News about the release.
**get_sample_sets**

**Examples**

```r
# Get the latest release
get_releases()
get_releases(date = 'latest')

# Get all releases
get_releases(date = 'all')

# Get a specific release by date
get_releases(date = '2020-08-19')
```

---

**Description**

Retrieves sample sets via the PGS Catalog REST API. The REST API is queried multiple times with the criteria passed as arguments (see below). By default all sample sets that match the criteria supplied in the arguments are retrieved: this corresponds to the default option `set_operation` set to 'union'. If you rather have only the associations that match simultaneously all criteria provided, then set `set_operation` to 'intersection'.

**Usage**

```r
get_sample_sets(
  pss_id = NULL,
  pgs_id = NULL,
  set_operation = "union",
  interactive = TRUE,
  verbose = FALSE,
  warnings = TRUE,
  progress_bar = TRUE
)
```

**Arguments**

- `pss_id` A character vector of PGS Catalog sample sets accession identifiers.
- `pgs_id` A character vector of PGS Catalog score accession identifiers.
- `set_operation` Either 'union' or 'intersection'. This tells how performance metrics retrieved by different criteria should be combined: 'union' binds together all results removing duplicates and 'intersection' only keeps same sample sets found with different criteria.
- `interactive` A logical. If all sample sets are requested, whether to ask interactively if we really want to proceed.
get_scores

verbose A logical indicating whether the function should be verbose about the different queries or not.

warnings A logical indicating whether to print warnings, if any.

progress_bar Whether to show a progress bar indicating download progress from the REST API server.

Details

Please note that all search criteria are vectorised, thus allowing for batch mode search.

Value

A sample_sets object.

Examples

# Search by PGS identifier
get_sample_sets(pgs_id = 'PGS000013')

# Search by the PSS identifier
get_sample_sets(pss_id = 'PSS000068')

get_scores

Get PGS Catalog Scores

Description

Retrieves polygenic scores via the PGS Catalog REST API. The REST API is queried multiple times with the criteria passed as arguments (see below). By default all scores that match the criteria supplied in the arguments are retrieved: this corresponds to the default option set_operation set to 'union'. If you rather have only the associations that match simultaneously all criteria provided, then set set_operation to 'intersection'.

Usage

get_scores(
  pgs_id = NULL,
  efo_id = NULL,
  pubmed_id = NULL,
  set_operation = "union",
  interactive = TRUE,
  verbose = FALSE,
  warnings = TRUE,
  progress_bar = TRUE
)

get_traits

Get PGS Catalog Traits

Description

Retrieves traits via the PGS Catalog REST API. The REST API is queried multiple times with the criteria passed as arguments (see below). By default all traits that match the criteria supplied in the arguments are retrieved: this corresponds to the default option set_operation set to 'union'. If you rather have only the traits that match simultaneously all criteria provided, then set set_operation to 'intersection'.

Arguments

pgs_id A character vector of PGS Catalog score accession identifiers.

efo_id A character vector of EFO identifiers.

pubmed_id An integer vector of PubMed identifiers.

set_operation Either 'union' or 'intersection'. This tells how scores retrieved by different criteria should be combined: 'union' binds together all results removing duplicates and 'intersection' only keeps same scores found with different criteria.

interactive A logical. If all scores are requested, whether to ask interactively if we really want to proceed.

verbose A logical indicating whether the function should be verbose about the different queries or not.

warnings A logical indicating whether to print warnings, if any.

progress_bar Whether to show a progress bar as the queries are performed.

Details

Please note that all search criteria are vectorised, thus allowing for batch mode search.

Value

A scores object.

Examples

# By 'pgs_id'
gscores(pgs_id = 'PGS000088')

# By 'efo_id'
gscores(efo_id = 'EFO_0007992')

# By 'pubmed_id'
gscores(pubmed_id = '25748612')
Usage

get_traits(
  efo_id = NULL,
  trait_term = NULL,
  exact_term = TRUE,
  include_children = FALSE,
  set_operation = "union",
  interactive = TRUE,
  verbose = FALSE,
  warnings = TRUE,
  progress_bar = TRUE
)

Arguments

efo_id A character vector of EFO identifiers.

trait_term A character vector of terms to be matched against trait identifiers (efo_id), trait descriptions, synonyms thereof, externally mapped terms, or even trait categories.

exact_term A logical value, indicating whether to match the trait_term exactly (TRUE) or not (FALSE).

include_children A logical value, indicating whether to include child traits or not.

set_operation Either 'union' or 'intersection'. This tells how performance metrics retrieved by different criteria should be combined: 'union' binds together all results removing duplicates and 'intersection' only keeps same sample sets found with different criteria.

interactive A logical. If all traits are requested, whether to ask interactively if we really want to proceed.

verbose A logical indicating whether the function should be verbose about the different queries or not.

warnings A logical indicating whether to print warnings, if any.

progress_bar Whether to show a progress bar indicating download progress from the REST API server.

Details

Please note that all search criteria are vectorised, thus allowing for batch mode search.

Value

A traits object.
get_trait_categories

Examples

# Get a trait by its EFO identifier
get_traits(efo_id = 'EFO_0004631')

# Get a trait by matching a term in EFO identifier ('efo_id'), label, # description synonyms, categories, or external mapped terms
get_traits(trait_term = 'stroke', exact_term = FALSE)

# Get a trait matching its name ('trait') exactly (default)
get_traits(trait_term = 'stroke', exact_term = TRUE)

# Get traits, excluding its children traits (default)
get_traits(trait_term = 'breast cancer')

# Get traits, including its children traits (check column 'is_child' for # child traits)
get_traits(trait_term = 'breast cancer', include_children = TRUE)

get_trait_categories Get PGS Catalog Trait Categories

Description

Retrieves all trait categories via the PGS Catalog REST API.

Usage

get_trait_categories(verbose = FALSE, warnings = TRUE, progress_bar = TRUE)

Arguments

verbose A logical indicating whether the function should be verbose about the different queries or not.
warnings A logical indicating whether to print warnings, if any.
progress_bar Whether to show a progress bar indicating download progress from the REST API server.

Value

A trait_categories object.

Examples

get_trait_categories(progress_bar = FALSE)
Number of PGS Catalog entities

Description

This function returns the number of entities in a PGS Catalog object. To avoid ambiguity with `dplyr::n()` use `quincunx::n()`.

Usage

```r
n(x, unique = FALSE)
```

## S4 method for signature 'scores'
```
n(x, unique = FALSE)
```

## S4 method for signature 'publications'
```
n(x, unique = FALSE)
```

## S4 method for signature 'traits'
```
n(x, unique = FALSE)
```

## S4 method for signature 'performance_metrics'
```
n(x, unique = FALSE)
```

## S4 method for signature 'sample_sets'
```
n(x, unique = FALSE)
```

## S4 method for signature 'cohorts'
```
n(x, unique = FALSE)
```

## S4 method for signature 'trait_categories'
```
n(x, unique = FALSE)
```

## S4 method for signature 'releases'
```
n(x, unique = FALSE)
```

Arguments

- `x` A `scores`, `publications`, `traits`, `performance_metrics`, `sample_sets`, `cohorts`, `trait_categories` or `releases` object.
- `unique` Whether to count only unique entries (TRUE) or not (FALSE).

Value

An integer scalar.
open_in_dbsnp

Examples

# Return the number of polygenic scores in a scores object:
my_scores <- get_scores(pgs_id = c('PGS000007', 'PGS000007', 'PGS000042'))
n(my_scores)

# If you want to count unique scores only, then use the 'unique' parameter:

# Total number of curated publications in the PGS Catalog:
all_pub <- get_publications(interactive = FALSE, progress_bar = FALSE)
n(all_pub)

# Total number of curated traits in the PGS Catalog:
all_traits <- get_traits(interactive = FALSE, progress_bar = FALSE)
n(all_traits)

open_in_dbsnp

Browse dbSNP from SNP identifiers.

Description

This function launches the web browser at dbSNP and opens a tab for each SNP identifier.

Usage

open_in_dbsnp(variant_id)

Arguments

variant_id A variant identifier, a character vector.

Value

Returns TRUE if successful. Note however that this function is run for its side effect.

Examples

open_in_dbsnp('rs56261590')
open_in_pgs_catalog  

Browse PGS Catalog entities from the PGS Catalog Web Graphical User Interface

Description

This function launches the web browser and opens a tab for each identifier on the PGS Catalog web graphical user interface: https://www.pgscatalog.org/.

Usage

open_in_pgs_catalog(
  identifier = NULL,
  pgs_catalog_entity = c("pgs", "pgp", "pss", "efo")
)

Arguments

identifier A vector of identifiers. The identifiers can be: PGS, PGP, PSS or EFO identifiers.
pgs_catalog_entity Either 'pgs' (default), 'pgp', 'pss', 'efo'. This argument indicates the type of the identifiers passed in identifier.

Value

Returns TRUE if successful, or FALSE otherwise. But note that this function is run for its side effect.

Examples

# Open in PGS scores Catalog Web Graphical User Interface
open_in_pgs_catalog(c("PGS000001", "PGS000002"))

# Open PGS Catalog Publications
open_in_pgs_catalog(c("PGP000001", "PGP000002"),
  pgs_catalog_entity = 'pgp')

# Open Sample Sets (PSS)
open_in_pgs_catalog(c("PSS000001", "PSS000002"),
  pgs_catalog_entity = 'pss')

# Open EFO traits (EFO)
open_in_pgs_catalog(c("EFO_0001645", "MONDO_0007254"),
  pgs_catalog_entity = 'efo')
open_in_pubmed

Browse PubMed from PubMed identifiers.

Description

This function launches the web browser and opens a tab for each PubMed citation.

Usage

```r
open_in_pubmed(pubmed_id)
```

Arguments

- `pubmed_id` A PubMed identifier, either a character or an integer vector.

Value

Returns TRUE if successful. Note however that this function is run for its side effect.

Examples

```r
open_in_pubmed(c("26301688", "30595370"))
```

performance_metrics-class

An S4 class to represent a set of PGS Catalog Performance Metrics

Description

The performance_metrics object consists of nine tables (slots) that combined form a relational database of a subset of performance metrics. Each performance metric is an observation (row) in the scores table (first table).

Slots

- `performance_metrics` A table of PGS Performance Metrics (PPM). Each PPM (row) is uniquely identified by the ppm_id column. Columns:
  - `ppm_id` A PGS Performance Metrics identifier. Example: "PPM000001".
  - `pgs_id` Polygenic Score (PGS) identifier.
  - `reported_trait` The author-reported trait that the PGS has been developed to predict. Example: "Breast Cancer".
  - `covariates` Comma-separated list of covariates used in the prediction model to evaluate the PGS.
**comments** Any other information relevant to the understanding of the performance metrics.

**publications** A table of publications. Each publication (row) is uniquely identified by the column `ppg_id`. Columns:

- `ppm_id` A PGS Performance Metrics identifier. Example: "PPM000001".
- `pgp_id` PGS Publication identifier. Example: "PGP000001".
- `pubmed_id` PubMed identifier. Example: "25855707".
- `publication_date` Publication date. Example: "2020-09-28". Note that the class of `publication_date` is `Date`.
- `publication` Abbreviated name of the journal. Example: "Am J Hum Genet".
- `author_fullname` First author of the publication. Example: "Mavaddat N".
- `doi` Digital Object Identifier (DOI). This variable is also curated to allow unpublished work (e.g. preprints) to be added to the catalog. Example: "10.1093/jnci/djv036".

**sample_sets** A table of sample sets. Each sample set (row) is uniquely identified by the column `pss_id`. Columns:

- `ppm_id` A PGS Performance Metrics identifier. Example: "PPM000001".
- `pss_id` A PGS Sample Set identifier. Example: "PSS000042".

**samples** A table of samples. Each sample (row) is uniquely identified by the combination of values from the columns: `ppm_id`, `pss_id`, and `sample_id`. Columns:

- `ppm_id` A PGS Performance Metrics identifier. Example: "PPM000001".
- `pss_id` A PGS Sample Set identifier. Example: "PSS000042".
- `sample_id` Sample identifier. This is a surrogate key to identify each sample.
- `stage` Sample stage: should be always Evaluation ("eval").
- `sample_size` Number of individuals included in the sample.
- `sample_cases` Number of cases.
- `sample_controls` Number of controls.
- `sample_percent_male` Percentage of male participants.
- `phenotype_description` Detailed phenotype description.
- `ancestry_category` Author reported ancestry is mapped to the best matching ancestry category from the NHGRI-EBI GWAS Catalog framework (see `ancestry_categories`) for possible values.
- `ancestry` A more detailed description of sample ancestry that usually matches the most specific description described by the authors (e.g. French, Chinese).
- `country` Author reported countries of recruitment (if available).
- `ancestry_additional_description` Any additional description not captured in the other columns (e.g. founder or genetically isolated populations, or further description of admixed samples).
- `study_id` Associated GWAS Catalog study accession identifier, e.g., "GCST002735".
- `pubmed_id` PubMed identifier.
- `cohorts_additional_description` Any additional description about the samples (e.g. sub-cohort information).

**demographics** A table of sample demographics’ variables. Each demographics’ variable (row) is uniquely identified by the combination of values from the columns: `ppm_id`, `pss_id`, `sample_id`, and `variable`. Columns:
ppm_id A PGS Performance Metrics identifier. Example: "PPM000001".

pss_id A PGS Sample Set identifier. Example: "PSS000042".

sample_id Sample identifier. This is a surrogate identifier to identify each sample.

variable Demographics variable. Following columns report about the indicated variable.

estimate_type Type of statistical estimate for variable.

estimate The variable’s statistical value.

unit Unit of the variable.

variability_type Measure of statistical dispersion for variable, e.g. standard error (se) or standard deviation (sd).

variability The value of the measure of dispersion.

interval_type Type of statistical interval for variable: range, iqr (interquartile), ci (confidence interval).

interval_lower Interval lower bound.

interval_upper Interval upper bound.

cohorts A table of cohorts. Each cohort (row) is uniquely identified by the combination of values from the columns: ppm_id, sample_id and cohort_symbol. Columns:

ppm_id A PGS Performance Metrics identifier. Example: "PPM000001".

sample_id Sample identifier. This is a surrogate key to identify each sample.

cohort_symbol Cohort symbol.

cohort_name Cohort full name.

gs_effect_sizes A table of effect sizes per standard deviation change in PGS. Examples include regression coefficients (betas) for continuous traits, odds ratios (OR) and/or hazard ratios (HR) for dichotomous traits depending on the availability of time-to-event data. Each effect size is uniquely identified by the combination of values from the columns: ppm_id and effect_size_id. Columns:

ppm_id A PGS Performance Metrics identifier. Example: "PPM000001".

effect_size_id Effect size identifier. This is a surrogate identifier to identify each effect size.

estimate_type_long Long notation of the effect size (e.g. Odds Ratio).

estimate_type Short notation of the effect size (e.g. OR).

estimate The estimate’s value.

unit Unit of the estimate.

variability_type Measure of statistical dispersion for variable, e.g. standard error (se) or standard deviation (sd).

variability The value of the measure of dispersion.

interval_type Type of statistical interval for variable: range, iqr (interquartile), ci (confidence interval).

interval_lower Interval lower bound.

interval_upper Interval upper bound.

gs_classification_metrics A table of classification metrics. Examples include the Area under the Receiver Operating Characteristic (AUROC) or Harrell’s C-index (Concordance statistic). Columns:

ppm_id A PGS Performance Metrics identifier. Example: "PPM000001".
classification_metrics_id Classification metric identifier. This is a surrogate identifier to identify each classification metric.

estimate_type_long Long notation of the classification metric (e.g. Concordance Statistic).

estimate_type Short notation classification metric (e.g. C-index).

estimate The estimate’s value.

unit Unit of the estimate.

variability_type Measure of statistical dispersion for variable, e.g. standard error (se) or standard deviation (sd).

variability The value of the measure of dispersion.

interval_type Type of statistical interval for variable: range, iqr (interquartile), ci (confidence interval).

interval_lower Interval lower bound.

interval_upper Interval upper bound.

pgs_other_metrics A table of other metrics that are neither effect sizes nor classification metrics. Examples include: R² (proportion of the variance explained), or reclassification metrics. Columns:

ppm_id A PGS Performance Metrics identifier. Example: "PPM000001".

other_metrics_id Other metric identifier. This is a surrogate identifier to identify each metric.

estimate_type_long Long notation of the metric. Example: "Proportion of the variance explained".

estimate_type Short notation metric. Example: "R²".

estimate The estimate’s value.

unit Unit of the estimate.

variability_type Measure of statistical dispersion for variable, e.g. standard error (se) or standard deviation (sd).

variability The value of the measure of dispersion.

interval_type Type of statistical interval for variable: range, iqr (interquartile), ci (confidence interval).

interval_lower Interval lower bound.

interval_upper Interval upper bound.

---

pgp_to_pgs  

Map PGP identifiers to PGS identifiers

Description

Map PGP identifiers to PGS identifiers.
Usage

pgp_to_pgs(
    pgp_id = NULL,
    verbose = FALSE,
    warnings = TRUE,
    progress_bar = TRUE
)

Arguments

pgp_id  A character vector of PGS Catalog Publication identifiers, e.g., "PGP000001". If NULL then returns results for all PGP identifiers in the Catalog.
verbose  A logical indicating whether the function should be verbose about the different queries or not.
warnings  A logical indicating whether to print warnings, if any.
progress_bar  Whether to show a progress bar as the queries are performed.

Value

A data frame of two columns: pgp_id and pgs_id.

Examples

## Not run:
pgp_to_pgs('PGP000001')
pgp_to_pgs(c('PGP000017', 'PGP000042'))
## End(Not run)

---

pgp_to_ppm  Map PGP identifiers to PPM identifiers

Description

Map PGP identifiers to PPM identifiers.

Usage

pgp_to_ppm(
    pgp_id = NULL,
    verbose = FALSE,
    warnings = TRUE,
    progress_bar = TRUE
)
### Arguments

- **pgp_id**
  - A character vector of PGS Catalog Publication identifiers, e.g., "PGP000001". If NULL then returns results for all PGP identifiers in the Catalog.

- **verbose**
  - A logical indicating whether the function should be verbose about the different queries or not.

- **warnings**
  - A logical indicating whether to print warnings, if any.

- **progress_bar**
  - Whether to show a progress bar as the queries are performed.

### Value

A data frame of two columns: pgp_id and ppm_id.

### Examples

```r
## Not run:
pgp_to_ppm('PGP000001')
pgp_to_ppm(c('PGP000017', 'PGP000042'))
## End(Not run)
```

---

**pgp_to_pss**  
*Map PGP identifiers to PSS identifiers*

### Description

Map PGP identifiers to PSS identifiers.

### Usage

```r
pgp_to_pss(
  pgp_id = NULL,
  verbose = FALSE,
  warnings = TRUE,
  progress_bar = TRUE
)
```

### Arguments

- **pgp_id**
  - A character vector of PGS Catalog Publication identifiers, e.g., "PGP000001". If NULL then returns results for all PGP identifiers in the Catalog.

- **verbose**
  - A logical indicating whether the function should be verbose about the different queries or not.

- **warnings**
  - A logical indicating whether to print warnings, if any.

- **progress_bar**
  - Whether to show a progress bar as the queries are performed.
pgs_to_pgp

Value

A data frame of two columns: pgp_id and pss_id.

Examples

```r
## Not run:
pgp_to_pss('PGP000001')
pgp_to_pss(c('PGP000017', 'PGP000042'))
## End(Not run)
```

pgs_to_pgp

Map PGS identifiers to PGP identifiers

Description

Map PGS identifiers to PGP identifiers.

Usage

```r
pgs_to_pgp(
  pgs_id = NULL,
  verbose = FALSE,
  warnings = TRUE,
  progress_bar = TRUE
)
```

Arguments

- `pgs_id`: A character vector of PGS identifiers, e.g., "PGS000001". If NULL then returns results for all PGS identifiers in the Catalog.
- `verbose`: A logical indicating whether the function should be verbose about the different queries or not.
- `warnings`: A logical indicating whether to print warnings, if any.
- `progress_bar`: Whether to show a progress bar as the queries are performed.

Value

A data frame of two columns: pgs_id and pgp_id.

Examples

```r
## Not run:
pgs_to_pgp('PGS000001')
pgs_to_pgp(c('PGS000017', 'PGS000042'))
## End(Not run)
```
**pgs_to_ppm**  
*Map PGS identifiers to PPM identifiers*

**Description**
Map PGS identifiers to PPM identifiers.

**Usage**

```
pgs_to_ppm(pgs_id, verbose = FALSE, warnings = TRUE, progress_bar = TRUE)
```

**Arguments**

- `pgs_id` A character vector of PGS identifiers, e.g., "PGS000001".
- `verbose` A logical indicating whether the function should be verbose about the different queries or not.
- `warnings` A logical indicating whether to print warnings, if any.
- `progress_bar` Whether to show a progress bar as the queries are performed.

**Value**
A data frame of two columns: `pgs_id` and `ppm_id`.

**Examples**

```r
## Not run:
pgs_to_ppm("PGS000001")
pgs_to_ppm(c("PGS000017", "PGS000042"))
## End(Not run)
```

---

**pgs_to_pss**  
*Map PGS identifiers to PSS identifiers*

**Description**
Map PGS identifiers to PSS identifiers.

**Usage**

```
pgs_to_pss(pgs_id = NULL, verbose = FALSE, warnings = TRUE, progress_bar = TRUE)
```

**Examples**

```r
## Not run:
pgs_to_pss()
```
pgs_to_study

Arguments

- **pgs_id**: A character vector of PGS identifiers, e.g., "PGS000001". If NULL then returns results for all PGS identifiers in the Catalog.
- **verbose**: A logical indicating whether the function should be verbose about the different queries or not.
- **warnings**: A logical indicating whether to print warnings, if any.
- **progress_bar**: Whether to show a progress bar as the queries are performed.

Value

A data frame of two columns: pgs_id and pss_id.

Examples

```r
## Not run:
pgs_to_pss('PGS000001')
pgs_to_pss(c('PGS000017', 'PGS000042'))
## End(Not run)
```

pgs_to_study  Map PGS identifiers to GWAS study identifiers

Description

Map PGS identifiers to GWAS study identifiers. Retrieves GWAS study identifiers associated with samples used in the discovery stage of queried PGS identifiers.

Usage

```r
pgs_to_study(
  pgs_id = NULL,
  verbose = FALSE,
  warnings = TRUE,
  progress_bar = TRUE
)
```

Arguments

- **pgs_id**: A character vector of PGS Catalog score accession identifiers, e.g., "PGS000001". If NULL then returns results for all PGS identifiers in the Catalog.
- **verbose**: A logical indicating whether the function should be verbose about the different queries or not.
- **warnings**: A logical indicating whether to print warnings, if any.
- **progress_bar**: Whether to show a progress bar as the queries are performed.
ppm_to_pgp

Value

A data frame of two columns: pgs_id and study_id.

Examples

```r
## Not run:
pgs_to_study('PGS000001')
# Unmappable pgs ids will be missing, e.g., PGS000023
pgs_to_study(c('PGS000013', 'PGS000023'))
## End(Not run)
```

ppm_to_pgp  

Map PPM identifiers to PGP identifiers

Description

Map PPM identifiers to PGP identifiers.

Usage

```r
ppm_to_pgp(ppm_id, verbose = FALSE, warnings = TRUE, progress_bar = TRUE)
```

Arguments

- `ppm_id`: A character vector of PPM identifiers, e.g., "PPM000001".
- `verbose`: A logical indicating whether the function should be verbose about the different queries or not.
- `warnings`: A logical indicating whether to print warnings, if any.
- `progress_bar`: Whether to show a progress bar as the queries are performed.

Value

A data frame of two columns: ppm_id and pgp_id.

Examples

```r
## Not run:
ppm_to_pgp('PPM000001')
ppm_to_pgp(c('PPM000017', 'PPM000042'))
## End(Not run)
```
### ppm_to_pgs

**Map PPM identifiers to PGS identifiers**

#### Description

Map PPM identifiers to PGS identifiers.

#### Usage

```r
ppm_to_pgs(ppm_id, verbose = FALSE, warnings = TRUE, progress_bar = TRUE)
```

#### Arguments

- `ppm_id` A character vector of PPM identifiers, e.g., "PPPM000001".
- `verbose` A logical indicating whether the function should be verbose about the different queries or not.
- `warnings` A logical indicating whether to print warnings, if any.
- `progress_bar` Whether to show a progress bar as the queries are performed.

#### Value

A data frame of two columns: `ppm_id` and `pgs_id`.

#### Examples

```r
## Not run:
ppm_to_pgs("PPM000001")
ppm_to_pgs(c("PPM000017", "PPM000042"))

## End(Not run)
```

### ppm_to_pss

**Map PPM identifiers to PSS identifiers**

#### Description

Map PPM identifiers to PSS identifiers.

#### Usage

```r
ppm_to_pss(ppm_id, verbose = FALSE, warnings = TRUE, progress_bar = TRUE)
```

#### Value

A data frame of two columns: `ppm_id` and `pgs_id`.
Arguments

- **ppm_id**: A character vector of PPM identifiers, e.g., "PPM000001".
- **verbose**: A logical indicating whether the function should be verbose about the different queries or not.
- **warnings**: A logical indicating whether to print warnings, if any.
- **progress_bar**: Whether to show a progress bar as the queries are performed.

Value

A data frame of two columns: ppm_id and pss_id.

Examples

```r
## Not run:
ppm_to_pss('PPM000001')
ppm_to_pss(c('PPM000017', 'PPM000042'))

## End(Not run)
```

Map PSS identifiers to PGP identifiers

Description

Map PSS identifiers to PGP identifiers. This is a slow function because it starts by downloading first all Performance Metrics, as this is the linkage between PSS and PGP.

Usage

```r
pss_to_pgp(pss_id, verbose = FALSE, warnings = TRUE, progress_bar = TRUE)
```

Arguments

- **pss_id**: A character vector of PSS identifiers, e.g., "PSS000001".
- **verbose**: A logical indicating whether the function should be verbose about the different queries or not.
- **warnings**: A logical indicating whether to print warnings, if any.
- **progress_bar**: Whether to show a progress bar as the queries are performed.

Value

A data frame of two columns: pss_id and pgp_id.
### pss_to_pgs

**Map PSS identifiers to PGS identifiers**

**Description**

Map PSS identifiers to PGS identifiers. This is a slow function because it starts by downloading first all Performance Metrics, as this is the linkage between PSS and PGS.

**Usage**

```r
pss_to_pgs(pss_id, verbose = FALSE, warnings = TRUE, progress_bar = TRUE)
```

**Arguments**

- **pss_id**: A character vector of PSS identifiers, e.g., "PSS000001".
- **verbose**: A logical indicating whether the function should be verbose about the different queries or not.
- **warnings**: A logical indicating whether to print warnings, if any.
- **progress_bar**: Whether to show a progress bar as the queries are performed.

**Value**

A data frame of two columns: pss_id and pgs_id.

**Examples**

```r
## Not run:
pss_to_pgp('PSS000001')
pss_to_pgp(c('PSS000017', 'PSS000042'))
## End(Not run)
```
pss_to_ppm  
Map PSS identifiers to PPM identifiers

Description
Map PSS identifiers to PPM identifiers. This is a slow function because it starts by downloading first all Performance Metrics.

Usage
pss_to_ppm(pss_id, verbose = FALSE, warnings = TRUE, progress_bar = TRUE)

Arguments
- **pss_id**: A character vector of PSS identifiers, e.g., "PSS000001".
- **verbose**: A logical indicating whether the function should be verbose about the different queries or not.
- **warnings**: A logical indicating whether to print warnings, if any.
- **progress_bar**: Whether to show a progress bar as the queries are performed.

Value
A data frame of two columns: pss_id and ppm_id.

Examples
```r
# Not run:
pss_to_ppm('PSS000001')
pss_to_ppm(c('PSS000017', 'PSS000042'))
# End(Not run)
```

publications-class  
An S4 class to represent a set of PGS Catalog Publications

Description
The publications object consists of two tables (slots), each a table that combined form a relational database of a subset of PGS Catalog Publications. Each publication is an observation (row) in the publications table (first table).
Slots

publications  A table of publications. Each publication (row) is uniquely identified by the pgp_id column. Columns:

  pgp_id  PGS Publication identifier. Example: "PGP000001".
  pubmed_id  PubMed identifier. Example: "25855707".
  publication_date  Publication date. Example: "2020-09-28". Note that the class of publication_date is Date.
  publication  Abbreviated name of the journal. Example: "Am J Hum Genet".
  title  Publication title.
  author_fullname  First author of the publication. Example: "Mavaddat N".
  doi  Digital Object Identifier (DOI). This variable is also curated to allow unpublished work (e.g. preprints) to be added to the catalog. Example: "10.1093/jnci/djv036".
  authors  Concatenated list of all the publication authors.

gps_ids  A table of publication and associated PGS identifiers. Columns:

  pgp_id  PGS Publication identifier. Example: "PGP000001".
  pgs_id  Polygenic Score (PGS) identifier.
  stage  PGS stage: either "gwas/dev" or "eval".

Description

This function imports a PGS scoring file. For more information about the scoring file schema check vignette("pgs-scoring-file",package = "quincunx").

Usage

read_scoring_file(source, protocol = "http", metadata_only = FALSE)

Arguments

source  PGS scoring file. This can be specified in three forms: (i) a PGS identifier, e.g. "PGS000001", (ii) a path to a local file, e.g. "~/PGS000001.txt" or "~/PGS000001.txt.gz" or (iii) a direct URL to the PGS Catalog FTP server, e.g. "http://ftp.ebi.ac.uk/pub/databases/spot/pgs/scores/PGS000001/ScoringFiles/PGS000001.txt.gz".

protocol  Network protocol for communication with the PGS Catalog FTP server: either "http" or "ftp".

metadata_only  Whether to read only the comment block (header) from the scoring file.
Value

The returned value is a named list. The names are copied from the arguments passed in source. Each element of the list contains another list of two elements: "metadata" and "data". The "metadata" element contains data parsed from the header of the PGS scoring file. The "data" element contains a data frame with as many rows as variants that constitute the PGS score. The columns can vary. There are mandatory and optional columns. The mandatory columns are those that identify the variant, effect allele (effect_allele), and its respective weight (effect_weight) in the score. The columns that identify the variant can either be the rsID or the combination of chr_name and chr_position. The "data" element will be NULL if argument metadata_only is TRUE. For more information about the scoring file schema check vignette("pgs-scoring-file",package = "quincunx").

Examples

```r
## Not run:
# Read a PGS scoring file by PGS ID
# (internally, it translates the PGS ID
to the corresponding FTP URL)
try(read_scoring_file("PGS000655"))

# Equivalent to `read_scoring_file("PGS000655")`
url <- paste0(
  "http://ftp.ebi.ac.uk/",
  "pub/databases/spot/pgs/scores/",
  "PGS000655/ScoringFiles/",
  "PGS000655.txt.gz"
)
read_scoring_file(url)

# Reading from a local file
try(read_scoring_file("~/PGS000655.txt.gz"))

## End(Not run)
```

---

**releases-class**

An S4 class to represent a set of PGS Catalog Releases

Description

The releases object consists of four tables (slots) that combined form a relational database of a subset of PGS Catalog releases. Each release is an observation (row) in the releases table (first table).

Slots

- `releases` A table of PGS Catalog releases. Each release (row) is uniquely identified by the release date (date). Columns:
sample_sets-class

Date

Release date.

n_pgs

Number of newly released Polygenic Scores.

n_ppm

Number of newly released PGS Performance Metrics.

n_pgp

Number of newly released PGS Publications.

pgs_ids

A table of released Polygenic Scores (PGS) identifiers. Columns:

date

Release date.

pgs_id

Polygenic Score (PGS) identifier. Example: "PGS000001".

ppm_ids

A table of the released PGS Performance Metrics identifiers. Columns:

date

Release date.

ppm_id

A PGS Performance Metrics identifier. Example: "PPM000001".

pgp_ids

A table of the released PGS Publication identifiers. Columns:

date

Release date.

pgp_id

PGS Publication identifier. Example: "PGP000001".

Description

The sample_sets object consists of four tables (slots) that combined form a relational database of a subset of PGS Catalog sample sets. Each sample set is an observation (row) in the sample_sets table (first table).

Slots

sample_sets

A table of sample sets. Each sample set (row) is uniquely identified by the column pss_id. Columns:

pss_id

A PGS Sample Set identifier. Example: "PSS000042".

samples

A table of samples. Each sample (row) is uniquely identified by the combination of values from the columns: pss_id and sample_id. Columns:

pss_id

A PGS Sample Set identifier. Example: "PSS000042".

sample_id

Sample identifier. This is a surrogate key to identify each sample.

stage

Sample stage: should be always Evaluation ("eval").

sample_size

Number of individuals included in the sample.

sample_cases

Number of cases.

sample_controls

Number of controls.

sample_percent_male

Percentage of male participants.

phenotype_description

Detailed phenotype description.

ancestry_category

Author reported ancestry is mapped to the best matching ancestry category from the NHGRI-EBI GWAS Catalog framework (see ancestry_categories) for possible values.
**ancrecy**  A more detailed description of sample ancestry that usually matches the most specific description described by the authors (e.g. French, Chinese).

**country**  Author reported countries of recruitment (if available).

**ancestry_additional_description**  Any additional description not captured in the other columns (e.g. founder or genetically isolated populations, or further description of admixed samples).

**study_id**  Associated GWAS Catalog study accession identifier, e.g., "GCST002735".

**pubmed_id**  PubMed identifier.

**cohorts_additional_description**  Any additional description about the samples (e.g. subcohort information).

**demographics**  A table of sample demographics’ variables. Each demographics’ variable (row) is uniquely identified by the combination of values from the columns: pss_id, sample_id, and variable. Columns:

- **pss_id**  A PGS Sample Set identifier. Example: "PSS000042".
- **sample_id**  Sample identifier. This is a surrogate identifier to identify each sample.
- **variable**  Demographics variable. Following columns report about the indicated variable.
- **estimate_type**  Type of statistical estimate for variable.
- **estimate**  The variable’s statistical value.
- **unit**  Unit of the variable.
- **variability_type**  Measure of statistical dispersion for variable, e.g. standard error (se) or standard deviation (sd).
- **variability**  The value of the measure of dispersion.
- **interval_type**  Type of statistical interval for variable: range, iqr (interquartile), ci (confidence interval).
- **interval_lower**  Interval lower bound.
- **interval_upper**  Interval upper bound.

**cohorts**  A table of cohorts. Each cohort (row) is uniquely identified by the combination of values from the columns: pss_id, sample_id and cohort_symbol. Columns:

- **pss_id**  A PGS Sample Set identifier. Example: "PSS000042".
- **sample_id**  Sample identifier. This is a surrogate key to identify each sample.
- **cohort_symbol**  Cohort symbol.
- **cohort_name**  Cohort full name.

---

**scores-class**  An S4 class to represent a set of PGS Catalog Polygenic Scores

**Description**

The scores object consists of six tables (slots) that combined form a relational database of a subset of PGS Catalog polygenic scores. Each score is an observation (row) in the scores table (the first table).
**Scores**

A table of polygenic scores. Each polygenic score (row) is uniquely identified by the `pgs_id` column. Columns:

- **pgs_id**: Polygenic Score (PGS) identifier. Example: "PGS000001".
- **pgs_name**: This may be the name that the authors describe the PGS with in the source publication, or a name that a curator of the PGS Catalog has assigned to identify the score during the curation process (before a PGS identifier has been given). Example: PRS77_BC.
- **matches_publication**: Indicate if the PGS data matches the published polygenic score (TRUE). If not (FALSE), the authors have provided an alternative polygenic for the Catalog and some other data, such as performance metrics, may differ from the publication.
- **reported_trait**: The author-reported trait that the PGS has been developed to predict. Example: "Breast Cancer".
- **trait_additional_description**: Any additional description not captured in the other columns. Example: "Femoral neck BMD (g/cm²)".
- **pgs_method_name**: The name or description of the method or computational algorithm used to develop the PGS.
- **pgs_method_params**: A description of the relevant inputs and parameters relevant to the PGS development method/process.
- **n_variants**: Number of variants used to calculate the PGS.
- **n_variants_interactions**: Number of higher-order variant interactions included in the PGS.
- **assembly**: The version of the genome assembly that the variants present in the PGS are associated with. Example: GRCh37.
- **license**: The PGS Catalog distributes its data according to EBI’s standard Terms of Use. Some PGS have specific terms, licenses, or restrictions (e.g. non-commercial use) that we highlight in this field, if known.

**Publications**

A table of publications. Each publication (row) is uniquely identified by the `pgp_id` column. Columns:

- **pgs_id**: Polygenic Score (PGS) identifier.
- **pgp_id**: PGS Publication identifier. Example: "PGP000001".
- **pubmed_id**: PubMed identifier. Example: "25855707".
- **publication_date**: Publication date. Example: "2020-09-28". Note that the class of `publication_date` is `Date`.
- **publication**: Abbreviated name of the journal. Example: "Am J Hum Genet".
- **title**: Publication title.
- **author_fullname**: First author of the publication. Example: 'Mavaddat N'.
- **doi**: Digital Object Identifier (DOI). This variable is also curated to allow unpublished work (e.g. preprints) to be added to the catalog. Example: "10.1093/jnci/djv036".

**Samples**

A table of samples. Each sample (row) is uniquely identified by the combination of values from the columns: `pgs_id` and `sample_id`. Columns:

- **pgs_id**: Polygenic score identifier. An identifier that starts with 'PGS' and is followed by six digits, e.g. 'PGS000001'.
- **sample_id**: Sample identifier. This is a surrogate key to identify each sample.
stage  Sample stage: either "discovery" or "training".
sample_size  Number of individuals included in the sample.
sample_cases  Number of cases.
sample_controls  Number of controls.
sample_percent_male  Percentage of male participants.
phenotype_description  Detailed phenotype description.
ancestry_category  Author reported ancestry is mapped to the best matching ancestry category from the NHGRI-EBI GWAS Catalog framework (see ancestry_categories) for possible values.
ancestry  A more detailed description of sample ancestry that usually matches the most specific description described by the authors (e.g. French, Chinese).
country  Author reported countries of recruitment (if available).
ancestry_additional_description  Any additional description not captured in the other columns (e.g. founder or genetically isolated populations, or further description of admixed samples).
study_id  Associated GWAS Catalog study accession identifier, e.g., "GCST002735".
pubmed_id  PubMed identifier.
cohorts_additional_description  Any additional description about the samples (e.g. sub-cohort information).
demographics  A table of sample demographics’ variables. Each demographics’ variable (row) is uniquely identified by the combination of values from the columns: pgs_id, sample_id and variable. Columns:
    pgs_id  Polygenic Score (PGS) identifier.
sample_id  Sample identifier. This is a surrogate identifier to identify each sample.
variable  Demographics variable. Following columns report about the indicated variable.
estimate_type  Type of statistical estimate for variable.
estimate  The variable’s statistical value.
unit  Unit of the variable.
avariability_type  Measure of statistical dispersion for variable, e.g. standard error (se) or standard deviation (sd).
avariability  The value of the measure of dispersion.
interval_type  Type of statistical interval for variable: range, iqr (interquartile), ci (confidence interval).
interval_lower  Interval lower bound.
interval_upper  Interval upper bound.
cohorts  A table of cohorts. Each cohort (row) is uniquely identified by the combination of values from the columns: pgs_id, sample_id and cohort_symbol. Columns:
    pgs_id  Polygenic Score (PGS) identifier.
sample_id  Sample identifier. This is a surrogate key to identify each sample.
cohort_symbol  Cohort symbol.
cohort_name  Cohort full name.
traits  A table of EFO traits. Each trait (row) is uniquely identified by the combination of the columns pgs_id and efo_id. Columns:
**setop**

- `pgs_id` Polygenic Score (PGS) identifier.
- `efo_id` An EFO identifier.
- `trait` Trait name.
- `description` Detailed description of the trait from EFO.
- `url` External link to the EFO entry.

**stages_tally** A table of sample sizes and number of samples sets at each stage.

- `pgs_id` Polygenic Score (PGS) identifier.
- `stage` Sample stage: either "gwas", "dev" or "eval".
- `sample_size` Sample size.
- `n_sample_sets` Number of sample sets (only meaningful for the evaluation stage "eval").

**ancestry_frequencies** This table describes the ancestry composition at each stage.

- `pgs_id` Polygenic Score (PGS) identifier.
- `stage` Sample stage: either "gwas", "dev" or "eval".
- `ancestry_class_symbol` Ancestry class symbol.
- `frequency` Ancestry fraction (percentage).

**multi_ancestry_composition** A table of a breakdown of the ancestries included in multi-ancestries.

- `pgs_id` Polygenic Score (PGS) identifier.
- `stage` Sample stage: either "gwas", "dev" or "eval".
- `multi_ancestry_class_symbol` Multi-ancestry class symbol.
- `ancestry_class_symbol` Ancestry class symbol.

---

**setup**

*Set operations on PGS Catalog objects*

**Description**

Performs set union, intersection, and (asymmetric!) difference on two objects of either class `scores`, `publications`, `traits`, `performance_metrics`, `sample_sets`, `cohorts` or `trait_categories`. Note that `union()` removes duplicated entities, whereas `bind()` does not.

**Usage**

```r
union(x, y, ...)
intersect(x, y, ...)
setdiff(x, y, ...)
setequal(x, y, ...)
```

**Arguments**

- `x, y` Objects of either class `scores`, `publications`, `traits`, `performance_metrics`, `sample_sets`, `cohorts` or `trait_categories`.
- `...` other arguments passed on to methods.
Value

In the case of union(), intersect(), or setdiff(): an object of the same class as x and y. In the case of setequal(), a logical scalar.

Examples

```r
# Get some 'scores' objects:
my_scores_1 <- get_scores(c('PGS000012', 'PGS000013'))
my_scores_2 <- get_scores(c('PGS000013', 'PGS000014'))

# union()
# NB: with 'union()', PGS000013 is not repeated.
union(my_scores_1, my_scores_2)@scores

# intersect()

intersect(my_scores_1, my_scores_2)@scores

# setdiff()

setdiff(my_scores_1, my_scores_2)@scores

# setequal()

setequal(my_scores_1, my_scores_2)
setequal(my_scores_1, my_scores_1)
setequal(my_scores_2, my_scores_2)
```

<table>
<thead>
<tr>
<th>stages</th>
<th>Study stages</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

A dataset containing the various study stages assigned to samples in the PGS Catalog.

Usage

stages
study_to_pgs

Format

A data frame with 5 stages (rows) and 4 columns:

- **stage**: Study stage.
- **symbol**: One-letter symbol for the stage, or a comma separated combination thereof.
- **name**: Stage name.
- **definition**: Stage description.

Source

https://www.pgscatalog.org/docs/ancestry

Examples

stages

<table>
<thead>
<tr>
<th>study_to_pgs</th>
<th>Map GWAS studies identifiers to PGS identifiers</th>
</tr>
</thead>
</table>

Description

Map GWAS studies identifiers to PGS identifiers.

Usage

study_to_pgs(study_id, verbose = FALSE, warnings = TRUE, progress_bar = TRUE)

Arguments

- **study_id**: A character vector of GWAS Catalog study accession identifiers, e.g., "GCST001937".
- **verbose**: A logical indicating whether the function should be verbose about the different queries or not.
- **warnings**: A logical indicating whether to print warnings, if any.
- **progress_bar**: Whether to show a progress bar as the queries are performed.

Value

A data frame of two columns: study_id and pgs_id.

Examples

```r
## Not run:
study_to_pgs("GCST001937")
study_to_pgs(c("GCST000998", "GCST000338"))
## End(Not run)
```
traits-class

An S4 class to represent a set of PGS Catalog Traits

Description

The traits object consists of six slots, each a table (tibble), that combined form a relational database of a subset of PGS Catalog traits. Each trait is an observation (row) in the traits table — main table. All tables have the column efo_id as primary key.

Slots

traits A table of traits. Columns:
- efo_id An EFO identifier.
- parent_efo_id An EFO identifier of the parent trait.
- is_child Is this trait obtained because it is a child of other trait?
- trait Trait name.
- description Detailed description of the trait from EFO.
- url External link to the EFO entry.

pgs_ids A table of associated polygenic score identifiers. Columns:
- efo_id An EFO identifier.
- parent_efo_id An EFO identifier of the parent trait.
- is_child Is this trait obtained because it is a child of other trait?
- pgs_id Polygenic Score (PGS) identifier.

child_pgs_ids A table of polygenic score identifiers associated with the child traits. Columns:
- efo_id An EFO identifier.
- parent_efo_id An EFO identifier of the parent trait.
- is_child Is this trait obtained because it is a child of other trait?
- child_pgs_id Polygenic Score (PGS) identifiers associated with child traits.

trait_categories A table of associated trait categories. Columns:
- efo_id An EFO identifier.
- parent_efo_id An EFO identifier of the parent trait.
- is_child Is this trait obtained because it is a child of other trait?
- trait_category Trait category name.

trait_synonyms A table of associated trait synonyms. Columns:
- efo_id An EFO identifier.
- parent_efo_id An EFO identifier of the parent trait.
- is_child Is this trait obtained because it is a child of other trait?
- trait_synonyms Trait synonyms.

trait_mapped_terms A table of associated external references, identifiers or other terms. Columns:
- efo_id An EFO identifier.
- parent_efo_id An EFO identifier of the parent trait.
- is_child Is this trait obtained because it is a child of other trait?
- trait_mapped_terms Trait mapped terms.
trait_categories-class

An S4 class to represent a set of PGS Catalog Trait Categories

Description

The trait_categories object consists of two tables (slots) that combined form a relational database of a subset of PGS Catalog trait categories. Each score is an observation (row) in the trait_categories table (first table).

Slots

trait_categories A table of trait categories. Columns:
- trait_category Trait category name.

traits A table of associated traits. Columns:
- trait_category Trait category name.
- efo_id An EFO identifier.
- trait Trait name.
- description Detailed description of the trait from EFO.
- url External link to the EFO entry.

write_xlsx Export a PGS Catalog object to xlsx

Description

This function exports a PGS Catalog object to Microsoft Excel xlsx file. Each table (slot) is saved in its own sheet.

Usage

write_xlsx(x, file = stop("\"file\" must be specified"))

Arguments

x A scores, publications, traits, performance_metrics, sample_sets, cohorts, trait_categories or releases object.

file A file name to write to.

Value

No return value, called for its side effect.
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