PGS Catalog access with quincunx

auincunx

demographics

• pss id

· sample id

estimate type

variability_type

interval type

· interval lower

· interval_upper

variable

estimate

variability

• unit

Introduction

The **PGS Catalog** is a service provided by the EMBL-EBI and NHGRI that offers a manually curated and freely available database of published polygenic scores (PGS): https://www.pgscatalog.org/.

The PGS Catalog data provided by the **RESTful API** is organised around five core entities:

- **PGS** Polygenic Scores
- **PGP** PGS Publications
- **PSS** PGS Sample Sets
- **PPM** PGS Performance Metrics
- FFO traits

Get PGS Catalog Entities

quincunx facilitates the access to the Catalog via the RESTful API, allowing you to programmatically retrieve data directly into R. Each of the five entities is mapped to an S4 object of a class of the same name.

PGS CATALOG RETRIEVAL FUNCTIONS **S4 CLASSES** scores get_scores() get_publications() publications sample sets get sample sets() performance metrics get_performance_metrics() get traits() traits

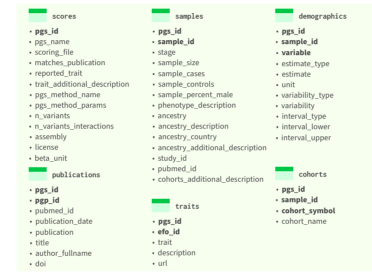
Query criteria for retrieval functions, e.g., PGS can be gueried by either pgs_id, efo_id or pubmed_id. These correspond to the criteria exposed by the PGS Catalog REST API: https://www.pgscatalog.org/rest/.

Search by	Example	PGS PGP PSS PPM EFO
pgs_id	"PGS000001"	
pgp_id	"PGP000001"	
pss_id	"PSS000001"	
ppm_id	"PPM000001"	
efo_id	"EFO_0000249"	
pubmed_id	"25855707"	
author	"Mavaddat"	
trait_term	"Alzheimer"	

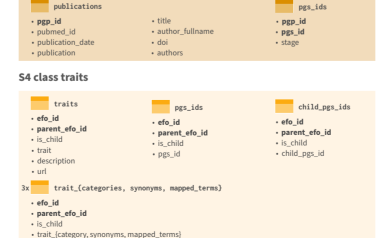
PGS Catalog Entities in R

PGS Catalog entities are represented as S4 classes in R. Each class represents a relational database of tidy data tables. All objects start with a table with the same name as the class. Combination of variables indicated in bold renders each row unique in each table.

S4 class scores



S4 class publications



S4 class sample sets

sample sets • pss id • pss id · pgs name · sample id · scoring_file · matches publication reported trait trait additional description

- · pgs method name
- · pgs_method_params
- · n variants
- · n variants interactions
- assembly
- beta unit
- · sample id
- · cohort_symbol
- cohort_name

samples

- - ancestry

- cohorts
- pss id

- · sample size
- sample cases
- · sample controls
- · sample_percent_male
- · phenotype_description
- · ancestry_description
- · ancestry country
- · ancestry additional description study_id
- pubmed id
- · cohorts_additional_description

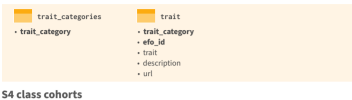
S4 class performance metrics

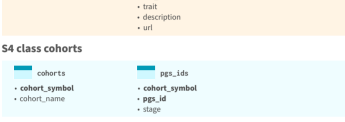


Other S4 Entities

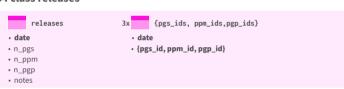
Besides the five PGS Catalog entities, there are three other objects that can be retrieved from the REST API: trait_categories, cohorts and releases.

S4 class trait_categories

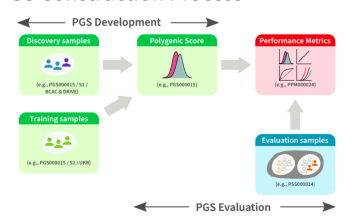




S4 class releases



PGS Construction Process



Samples and Polygenic Scores (PGS) are annotated according to their utilisation context in the PGS construction process, i.e. the stage variable in quincunx:

- "discovery"
- "training"
- "development" ("discovery" or "training")
- "evaluation"

Cohorts, Samples and Sample Sets

Cohorts

A cohort is a group of individuals with a shared characteristic. Cohorts are identified in quincunx by the cohort symbol variable.









Samples

A sample is a group of participants associated with none, one or more catalogued cohorts. The selection from a cohort can be either a subset or its totality. Samples are not identified in PGS Catalog with a global unique identifier, but quincunx assigns a surrogate identifier (sample_id) to allow relations between tables.

Possible compositions of samples:









Admixture of cohorts. (e.g., PGS000011 / S2 /

A cohort subset Another but disjoint subset (e.g., PSS000020 / S1 / MHI / N=862) (e.g., PSS000020 / S2 / MHI / N=2,333)

any cohort

Sample Sets

A sample set is a group of samples used in a polygenic score evaluation. Each sample set is identified in the PGS Catalog by a unique sample set identifier (PSS ID).

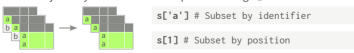


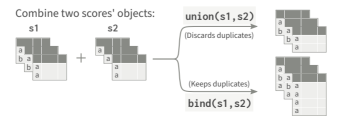
Manipulate Cases of S4 Entities

Get a scores object s consisting of two polygenic scores (PGS):

```
s <- get_scores(pgs_id = c('a', 'b'))
```

Subset object **s** by either identifier or position using `[`:







Polygenic scoring file

PGS scoring files are provided by the PGS Catalog to allow computation of polygenic scores by users. These files are hosted at the PGS Catalog FTP server: http://ftp.ebi.ac.uk/pub/databases/spot/pgs/ scores/. They are labelled by their respective PGS Score ID (e.g. PGS000001.txt.gz). For more details please visit: https://rmagno.eu/ quincunx/articles/pgs-scoring-file.html.

File Format

Each scoring file contains variant identification, effect alleles and respective score weights. The file is formatted as a gzipped tab-delimited text file, with a header containing brief metada about the score. You can read PGS scoring files into R with read_scoring_file().

PGS000117.txt.gz 1 ### PGS CATALOG SCORING FILE - see www.pgscatalog.org/downloads/#dl_ftp for... 2 ## POLYGENIC SCORE (PGS) INFORMATION # PGS ID = PGS000117 4 # Reported Trait = Cardiovascular Disease 5 # Original Genome Build = GRCh37 6 # Number of Variants = 267863 7 ## SOURCE INFORMATION 8 # PGP TD = PGP000054 9 # Citation = Elliott J et al. JAMA (2020). doi:10.1001/jama.2019.22241 chr_name chr_position effect_allele reference_allele effect_weight 10 rsTD 11 rs11240779 1 808631 0 00077622 12 rs1921 949608 -0.00583829 13 rs2710890 958905 -0.00182583 14 rs4970349 -0.001855691

Columns

The following table lists all possible columns in a PGS scoring file. A few columns are required (R), and most are optional (O); either the rsID alone or the combination of chr_name and chr_position are required, with the other being optional.

Column (Requirement)	Description	Example
rsID (R/O)	dbSNP Accession ID	"rs554219"
chr_name (R/O)	Chromosome name	"11"
chr_position (R/O)	Chromosome position	69516874
effect_allele (R)	Effect allele	"G"
reference_allele (O)	Reference allele	"C"
effect_weight (R)	Variant weight	0.117
locus_name (O)	Locus name	"CCND1"
weight_type (O)	Type of weight	"log(OR)", "beta_cox"
allelefrequency_effect (O)	Effect allele frequency	0.410
is_interaction (O)	Variant interaction?	TRUE or FALSE
is_recessive (O)	Recessive inheritance model?	TRUE or FALSE
is_haplotype (O)	Is effect allele a haplotype?	TRUE or FALSE
is_diplotype (O)	Is effect allele a diplotype?	TRUE or FALSE
imputation_method (O)	Imputation method	TODO
variant_description (O)	Variant description	TODO
inclusion_criteria (O)	Score inclusion criteria	TODO
OR (O)	Odds Ratio	1.12
HR (O)	Hazard Ratio	1.08