
grasp Documentation

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A Simple GRASP (grasp.nhlbi.nih.gov) API based on SQLAlchemy and Pandas.

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For an introduction see the [github readme](#)

For table information see the **'[wiki <https://github.com/MikeDacre/grasp/wiki>](https://github.com/MikeDacre/grasp/wiki)'** _

SUBMODULES

1.1 grasp.tables

GRASP table descriptions in SQLAlchemy ORM.

These tables do not exist in the GRASP data, which is a single flat file. By separating the data into these tables querying is much more efficient.

This submodule should only be used for querying.

```
class grasp.tables.SNP (**kwargs)
    Bases: sqlalchemy.ext.declarative.api.Base
```

An SQLAlchemy Table for GRASP SNPs.

Study and phenotype information are pushed to other tables to minimize table size and make querying easier.

Table Name: 'snps'

int
The integer of this class is the ID number of the row

str
The string of this class is the 'chr:pos' of the SNP

hvg_ids
A list of HGVS IDs for this SNP

ConservPredTFBS

CreationDate

EgtlMethMetabStudy

HUPfield

HumanEnhancer

InGene

InLincRNA

InMiRNA

InMiRNABS

LSSNP

LastCurationDate

NHLBIkey

NearestGene

ORegAnno

PolyPhen2

RNAedit

SIFT

UniProt

chrom

columns = `OrderedDict([('id', ('BigInteger', 'ID')), ('snpid', ('String', 'SNPid')), ('chrom', ('String', 'chr')), ('pos', ('Int`

A description of all columns in this table.

dbSNPClinStatus

dbSNPMAF

dbSNPfxn

dbSNPinfo

dbSNPvalidation

get_variant_info (*fields='dbsnp', pandas=True*)

Use the myvariant API to get info about this SNP.

Note that this service can be very slow. It will be faster to query multiple SNPs.

Parameters

- **fields** – Choose fields to display from: [‘docs.myvariant.info/en/latest/doc/data.html#available-fields’](https://docs.myvariant.info/en/latest/doc/data.html#available-fields) Good choices are ‘dbsnp’, ‘clinvar’, or ‘gwassnps’ Can also use ‘grasp’ to get a different version of this info.
- **pandas** – Return a dataframe instead of dictionary.

Returns A dictionary or a dataframe.

hvg_ids

The HVGS ID from myvariant.

id

paper_loc

phenotype_cats

phenotype_desc

population

population_id

pos

pval

snp_loc

Return a simple string containing the SNP location.

snpid

study

study_id

study_snpid

```
class grasp.tables.Phenotype (**kwargs)
```

Bases: sqlalchemy.ext.declarative.api.Base

An SQLAlchemy table to store the primary phenotype.

Table Name: 'phenos'

Columns: phenotype: The string phenotype from the GRASP DB, unique. alias: A short representation of the phenotype, not unique. studies: A link to the studies table.

int

The ID number.

str

The name of the phenotype.

alias

id

phenotype

studies

```
class grasp.tables.PhenoCats (**kwargs)
```

Bases: sqlalchemy.ext.declarative.api.Base

An SQLAlchemy table to store the lists of phenotype categories.

Table Name: 'pheno_cats'

Columns: category: The category from the grasp database, unique. alias: An easy to use alias of the category, not unique. snps: A link to all SNPs in this category. studies: A link to all studies in this category.

int

The PhenoCat ID

str

The category name

alias

category

id

snps

studies

```
class grasp.tables.Platform(platform)
```

Bases: sqlalchemy.ext.declarative.api.Base

An SQLAlchemy table to store the platform information.

Table Name: 'platforms'

Columns: platform: The name of the platform from GRASP. studies: A link to all studies using this platform.

int

The ID number of this platform

str

The name of the platform

id

platform

studies

class `grasp.tables.Population(population)`

Bases: `sqlalchemy.ext.declarative.api.Base`

An SQLAlchemy table to store the platform information.

Table Name: 'populations'

Columns: `population`: The name of the population. `studies`: A link to all studies in this population. `snps`: A link to all SNPs in this populations.

int

Population ID number

str

The name of the population

id

population

1.2 grasp.db

Functions for managing the GRASP database.

`get_session()` is used everywhere in the module to create a connection to the database. `initialize_database()` is used to build the database from the GRASP file. It takes about an hour 90 minutes to run and will overwrite any existing database.

`grasp.db.get_session(echo=False)`

Return a session and engine, uses config file.

Parameters `echo` – Echo all SQL to the console.

Returns

A SQLAlchemy session and engine object corresponding to the grasp database for use in querying.

Return type session, engine

`grasp.db.initialize_database(study_file, grasp_file, commit_every=250000, progress=False)`

Create the database quickly.

Study_file Tab delimited GRASP study file, available here:
github.com/MikeDacre/grasp/blob/master/grasp_studies.txt

Grasp_file Tab delimited GRASP file.

Commit_every How many rows to go through before committing to disk.

Progress Display a progress bar (db length hard coded).

1.3 grasp.config

Manage a persistent configuration for the database.

`grasp.config.config = <configparser.ConfigParser object>`

A globally accessible ConfigParser object, initialized with CONFIG_FILE.

`grasp.config.CONFIG_FILE = '/Users/dacre/.grasp'`

The PATH to the config file.

`grasp.config.init_config(db_type='', db_file='', db_host='', db_user='', db_pass='')`

Create an initial config file.

Parameters

- **db_type** – 'sqlite/mysql/postgresql'
- **db_file** – PATH to sqlite database file
- **db_host** – Hostname for mysql or postgresql server
- **db_user** – Username for mysql or postgresql server
- **db_pass** – Password for mysql or postgresql server (not secure)

Returns NoneType

Return type None

`grasp.config.init_config_interactive()`

Interact with the user to create a new config.

Uses readline autocompletion to make setup easier.

`grasp.config.write_config()`

Write the current config to CONFIG_FILE.

1.4 grasp.query

A mix of functions to make querying the database and analyzing the results faster.

`grasp.query.get_studies(primary_phenotype=None, pheno_cats=None, pheno_cats_alias=None, primary_pop=None, has_disc_pop=None, has_rep_pop=None, only_disc_pop=None, only_rep_pop=None, query=False, count=False, pandas=False)`

Return a list of studies filtered by phenotype and population.

There are two ways to query both phenotype and population.

Phenotype: GRASP provides a 'primary phenotype' for each study, which are fairly poorly curated. They also provide a list of phenotype categories, which are well curated. The problem with the categories is that there are multiple per study and some are to general to be useful. If using categories be sure to post filter the study list.

Note: I have made a list of aliases for the phenotype categories to make them easier to type. Use `pheno_cats_alias` for that.

Population: Each study has a primary population (list available with 'get_populations') but some studies also have other populations in the cohort. GRASP indexes all population counts, so those can be used to query also. To query these use `has_` or `only_` (exclusive) parameters, you can query either discovery populations or replication populations. Note that you cannot provide both `has_` and `only_` parameters for the same population type.

For doing population specific analyses most of the time you will want the `excl_disc_pop` query.

Argument Description: Phenotype Arguments are ‘primary_phenotype’, ‘pheno_cats’, and ‘pheno_cats_alias’.

Only provide one of pheno_cats or pheno_cats_alias

Population Arguments are *primary_pop*, *has_disc_pop*, *has_rep_pop*, *only_disc_pop*, *only_rep_pop*.

primary_pop is a simple argument, the others use bitwise flags for lookup.

The easiest way to use the following parameters is with the `_ref.PopFlag` object. It uses py-flags. For example:

```
pops = _ref.PopFlag.eur | _ref.PopFlag.afr
```

In addition you can provide a list of strings corresponding to PopFlag attributes.

Note: the *only_* parameters work as ANDs, not ORs. So *only_disc_pop*=‘eurlafr’ will return those studies that have BOTH european and african discovery populations, but no other discovery populations. On the other hand, *has_* works as an OR, and will return any study with any of the specified populations.

Parameters

- **primary_phenotype** – Phenotype of interest, string or list of strings.
- **pheno_cats** – Phenotype category of interest.
- **pheno_cats_alias** – Phenotype category of interest.
- **primary_pop** – Query the primary population, string or list of strings.
- **has_disc_pop** – Return all studies with these discovery populations
- **has_rep_pop** – Return all studies with these replication populations
- **only_disc_pop** – Return all studies with ONLY these discovery populations
- **only_rep_pop** – Return all studies with ONLY these replication populations
- **query** – Return the query instead of the list of study objects.
- **count** – Return a count of the number of studies.
- **pandas** – Return a dataframe of study information instead of the list.

Returns A list of study objects, a query, or a dataframe.

`grasp.query.get_snps(studies, pandas=True)`

Return a list of SNPs in a single population in a single phenotype.

Studies A list of studies.

Pandas Return a dataframe instead of a list of SNP objects.

Returns Either a DataFrame or list of SNP objects.

`grasp.query.get_variant_info(snp_list, fields='db SNP', pandas=True)`

Use the myvariant API to get info about this SNP.

Note that this service can be very slow.

Snp_list A list of SNP objects or ‘chr:loc’

Fields Choose fields to display from: docs.myvariant.info/en/latest/doc/data.html#available-fields
Good choices are ‘db SNP’, ‘clinvar’, or ‘gwas SNPs’ Can also use ‘grasp’ to get a different version of this info.

Pandas Return a dataframe instead of dictionary.

`grasp.query.get_pop_flags(pop_flags)`
Merge a list, string, int, or PopFlag series.

`grasp.query.get_phenotypes(list_only=False, dictionary=False, table=False)`
Return all phenotypes from the phenotype table.

List_only Return a simple text list instead of a list of Phenotype objects.

Dictionary Return a dictionary of phenotype=>ID

Table Return a pretty table for printing.

`grasp.query.get_phenotype_categories(list_only=False, dictionary=False, table=False)`
Return all phenotype categories from the PhenoCats table.

List_only Return a simple text list instead of a list of Phenotype objects.

Dictionary Return a dictionary of phenotype=>ID

Table Return a pretty table for printing.

`grasp.query.get_populations(list_only=False, dictionary=False, table=False)`
Return all phenotypes from the phenotype table.

List_only Return a simple text list instead of a list of Phenotype objects.

Dictionary Return a dictionary of population=>ID

Table Return a pretty table for printing.

`grasp.query.get_population_flags(list_only=False, dictionary=False, table=False)`
Return all population flags available in the PopFlags class.

List_only Return a simple text list instead of a list of Phenotype objects.

Dictionary Return a dictionary of population=>ID

Table Return a pretty table for printing.

`grasp.query.get_study_columns(list_only=False, dictionary=False, table=False)`
Return all columns in the Study table.

List_only Return a simple text list instead of a list of Phenotype objects.

Dictionary Return a dictionary of population=>ID

Table Return a pretty table for printing.

`grasp.query.get_snp_columns(list_only=False, dictionary=False, table=False)`
Return all columns in the SNP table.

List_only Return a simple text list instead of a list of Phenotype objects.

Dictionary Return a dictionary of population=>ID

Table Return a pretty table for printing.

REFERENCE

ref.py holds some simple lookups and the *PopFlags* classes that don't really go anywhere else. Holds reference objects for use elsewhere in the module.

class `grasp.ref.PopFlag`

A simplified bitwise flag system for tracking populations.

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