

Package: epigraphdb (via r-universe)

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Title Interface Package for the 'EpiGraphDB' Platform

Version 0.2.3

Description The interface package to access data from the 'EpiGraphDB' <<https://epigraphdb.org>> platform. It provides easy access to the 'EpiGraphDB' platform with functions that query the corresponding REST endpoints on the API <<https://api.epigraphdb.org>> and return the response data in the 'tibble' data frame format.

URL <https://mrcieu.github.io/epigraphdb-r/>

BugReports <https://github.com/MRCIEU/epigraphdb-r/issues>

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GET /confounder

Usage

```
confounder(
  exposure_trait = NULL,
  outcome_trait = NULL,
  type = c("confounder", "intermediate", "reverse_intermediate", "collider"),
  pval_threshold = 1e-05,
  mode = c("table", "raw")
)
```

Arguments

- exposure_trait** A trait name, e.g. "Body mass index", leaving `exposure_trait` as `NULL` will return MR information related to a specific outcome. **NOTE:** `exposure_trait` and `outcome_trait` cannot be both `NULL`.
- outcome_trait** A trait name, e.g. "Coronary heart disease", leaving `outcome_trait` as `NULL` will return MR information related to a specific `exposure_trait`. **NOTE:** `exposure_trait` and `outcome_trait` cannot be both `NULL`.

type	One in ["confounder", "intermediate", "reverse_intermediate", "collider"] Refer to the confounder view in web application for details
pval_threshold	P-value threshold
mode	If mode = "table", returns a data frame (a tibble as per tidyverse convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by httr .

Value

Data from GET /confounder

Examples

```
## Not run:  
confounder(exposure_trait = "Body mass index", outcome_trait = "Coronary heart disease")  
  
## End(Not run)
```

cypher

Send a query in Cypher to EpiGraphDB

Description

NOTE: this function is intended for advanced uses. Regular users are recommended to use standard query functions

Usage

```
cypher(query, mode = c("table", "raw"))
```

Arguments

query	A Cypher query.
mode	If mode = "table", returns a data frame (a tibble as per tidyverse convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by httr .

Examples

```
## Not run:  
cypher("MATCH (n:Gwas) RETURN n LIMIT 2")  
  
## End(Not run)
```

`drugs_risk_factors` *Drugs for risk factors*

Description

`GET /drugs/risk-factors`

Usage

```
drugs_risk_factors(trait, pval_threshold = 1e-08, mode = c("table", "raw"))
```

Arguments

- `trait` A trait name
- `pval_threshold` P-value threshold
- `mode` If `mode = "table"`, returns a data frame (a `tibble` as per `tidyverse` convention). If `mode = "raw"`, returns a raw response from EpiGraphDB API with minimal parsing done by `httr`.

Value

Data from `GET /drugs/risk-factors`

Examples

```
## Not run:
drugs_risk_factors(trait = "Body mass index")

## End(Not run)
```

`genetic_cor` *Genetic correlations between traits*

Description

`GET /genetic-cor`

Usage

```
genetic_cor(trait, cor_coef_threshold = 0.8, mode = c("table", "raw"))
```

Arguments

trait name of the trait, e.g. "body mass index"
cor_coef_threshold correlation coefficient threshold
mode If mode = "table", returns a data frame (a **tibble** as per **tidyverse** convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by **httr**.

Value

Data from GET /genetic_cor

Examples

```

## Not run:
genetic_cor(trait = "Body mass index") %>%
  dplyr::glimpse()

## End(Not run)

# Use a different threshold
## Not run:
genetic_cor(trait = "Body mass index", cor_coef_threshold = 0.4) %>%
  dplyr::glimpse()

## End(Not run)

```

Description

[GET /literature/gwas](#)

Usage

```
literature_gwas(trait, semmed_predicate = NULL, mode = c("table", "raw"))
```

Arguments

trait A trait name
semmed_predicate Either NULL which returns entries from all predicates, or a SemMed predicate e.g. "DIAGNOSES" or "ASSOCIATED_WITH"
mode If mode = "table", returns a data frame (a **tibble** as per **tidyverse** convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by **httr**.

Value

Data from GET /literature/gwas

Examples

```
## Not run:
literature_gwas(trait = "Body mass index")

## End(Not run)
```

mappings_gene_to_protein

Return protein uniprot_id from associated genes

Description

POST /mappings/gene-to-protein

Usage

```
mappings_gene_to_protein(
  gene_name_list = NULL,
  gene_id_list = NULL,
  by_gene_id = FALSE,
  mode = c("table", "raw")
)
```

Arguments

gene_name_list	List of HGNC symbols of the genes (default)
gene_id_list	List of Ensembl gene IDs (when by_gene_id == TRUE)
by_gene_id	Search for gene ids (Ensembl gene IDs) instead of gene names (HGNC symbols)
mode	If mode = "table", returns a data frame (a tibble as per tidyverse convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by httr .

Value

Data from POST /mappings/gene-to-protein

Examples

```
# By HGNC symbols
## Not run:
mappings_gene_to_protein(gene_name_list = c("GCH1", "MYOF"))

## End(Not run)

# By Ensembl IDs
## Not run:
mappings_gene_to_protein(gene_id_list = c("ENSG00000162594", "ENSG00000113302"), by_gene_id = TRUE)

## End(Not run)
```

meta_nodes_list *List meta nodes (e.g. Gwas, Gene, etc.)*

Description

GET /meta/nodes/list

Usage

```
meta_nodes_list(mode = c("raw"))
```

Arguments

mode If mode = "table", returns a data frame (a **tibble** as per **tidyverse** convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by **httr**.

Value

Data from GET /meta/nodes/list

Examples

```
## Not run:
meta_nodes_list()

## End(Not run)
```

meta_nodes_list_node *List nodes under a meta node*

Description

`GET /meta/nodes/{meta_node}/list`

Usage

```
meta_nodes_list_node(
  meta_node,
  full_data = TRUE,
  limit = 10,
  offset = 0,
  mode = c("table", "raw")
)
```

Arguments

<code>meta_node</code>	Name of a meta node (e.g. Gwas). Use <code>meta_nodes_list</code> to get the full list of meta nodes.
<code>full_data</code>	When <code>False</code> , only return the <code>id</code> and <code>name</code> fields (their specific names differ in specific nodes) for a node. This is useful if you want your queries to return results faster with smaller amount of data requested.
<code>limit</code>	Max number of items to retrieve.
<code>offset</code>	Number of items to skip. Use <code>limit</code> and <code>offset</code> in combination to do pagination.
<code>mode</code>	If <code>mode = "table"</code> , returns a data frame (a <code>tibble</code> as per <code>tidyverse</code> convention). If <code>mode = "raw"</code> , returns a raw response from EpiGraphDB API with minimal parsing done by <code>httr</code> .

Value

Data from `GET /meta/nodes/{meta_node}/list`

Examples

```
# List the first 5 Gwas nodes, with only id and name fields
## Not run:
meta_nodes_list_node(meta_node = "Gwas", full_data = FALSE, limit = 5)

## End(Not run)

# List the 6th - 10th Disease nodes, with full properties
## Not run:
meta_nodes_list_node(meta_node = "Disease", full_data = TRUE, limit = 5, offset = 0)

## End(Not run)
```

meta_nodes_search_node

Search a node by its id field, or its name field

Description

GET /meta/nodes/{meta_node}/search

Usage

```
meta_nodes_search_node(  
  meta_node,  
  id = NULL,  
  name = NULL,  
  limit = 10,  
  full_data = TRUE,  
  mode = c("table", "raw")  
)
```

Arguments

meta_node	Name of a meta node (e.g. Gwas). Use <code>meta_nodes_list</code> to get the full list of meta nodes.
id	The id field of a node (e.g. "ieu-a-2" for a Gwas). Use EpiGraphDB web UI to get a sense of what those ids are for entities.
name	The name field of a node (e.g. "body mass index" for a Gwas). Use EpiGraphDB web UI to get a sense of what those names are for entities.
limit	Max number of items to retrieve.
full_data	When False, only return the id and name fields (their specific names differ in specific nodes) for a node. This is useful if you want your queries to return results faster with smaller amount of data requested.
mode	If <code>mode = "table"</code> , returns a data frame (a <code>tibble</code> as per <code>tidyverse</code> convention). If <code>mode = "raw"</code> , returns a raw response from EpiGraphDB API with minimal parsing done by <code>httr</code> .

Value

Data from GET /meta/nodes/{meta_node}/search

Examples

```
# Search Gwas nodes  
## Not run:  
meta_nodes_search_node(meta_node = "Gwas", name = "body mass index")  
  
## End(Not run)
```

<code>meta_rels_list</code>	<i>List meta rels (e.g. MR, etc.)</i>
-----------------------------	---------------------------------------

Description

`GET /meta/rels/list`

Usage

```
meta_rels_list(mode = c("raw"))
```

Arguments

<code>mode</code>	If <code>mode = "table"</code> , returns a data frame (a <code>tibble</code> as per <code>tidyverse</code> convention). If <code>mode = "raw"</code> , returns a raw response from EpiGraphDB API with minimal parsing done by <code>httr</code> .
-------------------	--

Value

Data from `GET /meta/rels/list`

Examples

```
## Not run:  
meta_rels_list()  
  
## End(Not run)
```

<code>meta_rels_list_rel</code>	<i>List relationships under a meta relationship</i>
---------------------------------	---

Description

`GET /meta/rels/{meta_rel}/list`

Usage

```
meta_rels_list_rel(meta_rel, limit = 10, offset = 0, mode = c("table", "raw"))
```

Arguments

<code>meta_rel</code>	Name of a meta relationship (e.g. MR). Use <code>meta_rels_list</code> to get the full list of meta relationships.
<code>limit</code>	Max number of items to retrieve.
<code>offset</code>	Number of items to skip. Use <code>limit</code> and <code>offset</code> in combination to do pagination.
<code>mode</code>	If <code>mode = "table"</code> , returns a data frame (a <code>tibble</code> as per <code>tidyverse</code> convention). If <code>mode = "raw"</code> , returns a raw response from EpiGraphDB API with minimal parsing done by <code>httr</code> .

Value

Data from GET /meta/rels/{meta_rel}/list

Examples

```
# List the first 5 MR relationships
## Not run:
meta_rels_list_rel(meta_rel = "MR_EVE_MR", limit = 5)

## End(Not run)
```

mr

Return information related to Mendelian Randomisation

Description

GET /mr

Usage

```
mr(
  exposure_trait = NULL,
  outcome_trait = NULL,
  pval_threshold = 1e-05,
  mode = c("table", "raw")
)
```

Arguments

<code>exposure_trait</code>	A trait name, e.g. "Body mass index", leaving <code>exposure_trait</code> as NULL will return MR information related to a specific outcome. NOTE: <code>exposure_trait</code> and <code>outcome_trait</code> cannot be both NULL.
<code>outcome_trait</code>	A trait name, e.g. "Coronary heart disease", leaving <code>outcome_trait</code> as NULL will return MR information related to a specific <code>exposure_trait</code> . NOTE: <code>exposure_trait</code> and <code>outcome_trait</code> cannot be both NULL.

pval_threshold P-value threshold
 mode If mode = "table", returns a data frame (a **tibble** as per **tidyverse** convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by **httr**.

Value

Data from GET /mr

Examples

```
# Returns a data frame
## Not run:
mr(exposure_trait = "Body mass index", outcome_trait = "Coronary heart disease")

## End(Not run)

# Returns raw response
## Not run:
mr(
  exposure_trait = "Body mass index", outcome_trait = "Coronary heart disease",
  mode = "raw"
) %>% str()

## End(Not run)

# Use a different threshold
## Not run:
mr(exposure_trait = "Body mass index", pval_threshold = 1e-8)

## End(Not run)
```

obs_cor

Observational correlations between traits

Description

GET /obs-cor

Usage

```
obs_cor(trait, cor_coef_threshold = 0.8, mode = c("table", "raw"))
```

Arguments

trait name of the trait, e.g. "body mass index"
 cor_coef_threshold correlation coefficient threshold

mode If mode = "table", returns a data frame (a **tibble** as per **tidyverse** convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by **httr**.

Value

Data from GET /obs-cor

Examples

```
## Not run:
obs_cor(trait = "Body mass index (BMI)") %>%
  dplyr::glimpse()

## End(Not run)

# Use a different threshold
## Not run:
obs_cor(trait = "Body mass index (BMI)", cor_coef_threshold = 0.8) %>%
  dplyr::glimpse()

## End(Not run)
```

Description

GET /ontology/gwas-efo

Usage

```
ontology_gwas_efo(
  trait = NULL,
  efo_term = NULL,
  fuzzy = TRUE,
  mode = c("table", "raw")
)
```

Arguments

trait	trait name, e.g. "body mass"
efo_term	EFO term, e.g. "systolic blood pressure"
fuzzy	whether query with exact matching (FALSE) or fuzzy matching (default, TRUE)
mode	If mode = "table", returns a data frame (a tibble as per tidyverse convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by httr .

Value

Data from GET /ontology/gwas-efo

Examples

```
## Not run:
ontology_gwas_efo(trait = "blood", fuzzy = FALSE)

## End(Not run)

## Not run:
ontology_gwas_efo(efo_term = "blood pressure", fuzzy = FALSE)

## End(Not run)
```

pathway

Pathway evidence

Description

[GET /pathway](#)

Usage

```
pathway(trait, pval_threshold = 1e-05, mode = c("table", "raw"))
```

Arguments

trait	A trait name
pval_threshold	P-value threshold
mode	If mode = "table", returns a data frame (a tibble as per tidyverse convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by httr .

Value

Data from GET /pathway

Examples

```
## Not run:
pathway(trait = "Body mass index")

## End(Not run)
```

pqt1*Return information related to the pQTL analysis*

Description`GET /pqt1/`**Usage**

```
pqt1(  
  query,  
  rtype = c("mrres", "simple", "sglmr", "inst", "sense"),  
  pvalue = 0.05,  
  searchflag = c("traits", "proteins"),  
  mode = c("table", "raw")  
)
```

Arguments

query	(Required) A protein coding gene name or a trait name, e.g. "ADAM19" or "Inflammatory bowel disease" which cannot be NULL.
rtype	(Optional) A type of data to be extracted, which can be one of these options: <ol style="list-style-type: none">1. simple: Basic summary2. mrres: MR results (DEFAULT)3. sglmr: Single SNP MR results4. inst: SNP information5. sense: Sensitivity analysis NOTE: mrres is a DEFAULT option.
pvalue	(Optional) A pvalue threshold for MR results with the DEFAULT set to 0.05. NOTE: this threshold applies to any rtype chosen.
searchflag	(Required) A flag to indicate whether you are searching for proteins or traits which cannot be NULL. If query is a protein name, then this flag should be "proteins"; if query is a trait, this flag should be "traits". NOTE: if the wrong flag is chosen for query, there will be no result returned.
mode	If mode = "table", returns a data frame (a tibble as per tidyverse convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by httr .

Value

Data from GET /pqt1/

Examples

```
# Returns a data frame of MR results, while searching for proteins
## Not run:
pqtL(query = "ADAM19", searchflag = "proteins")

## End(Not run)

# Returns a data frame with SNP information, while searching for traits
## Not run:
pqtL(
  query = "Inflammatory bowel disease",
  rtype = "inst",
  searchflag = "traits"
)

## End(Not run)

# Change a pvalue threshold (the default is 0.05)
## Not run:
pqtL(
  query = "Inflammatory bowel disease",
  rtype = "inst",
  pvalue = 1.0,
  searchflag = "traits"
)

## End(Not run)

# Returns raw response if mode="raw"
## Not run:
pqtL(
  query = "ADAM19", searchflag = "proteins",
  mode = "raw"
) %>% str()

## End(Not run)
```

pqtL_list

Return a list of all proteins/exposures or traits/outcomes available in the database

Description

GET /pqtL/list/

Usage

```
pqtL_list(flag = c("exposures", "outcomes"), mode = c("table", "raw"))
```

Arguments

<code>flag</code>	(Optional) A flag which indicates whether the list of exposures (if "exposures") or outcomes (if "outcomes") should be returned. The DEFAULT is "exposures".
<code>mode</code>	If <code>mode = "table"</code> , returns a data frame (a <code>tibble</code> as per <code>tidyverse</code> convention). If <code>mode = "raw"</code> , returns a raw response from EpiGraphDB API with minimal parsing done by <code>httr</code> .

Value

Data from GET /pqtl/list/

Examples

```
# Returns a list of available proteins (exposures)
## Not run:
pqtl_list()

## End(Not run)

# Returns a list of available traits (outcomes)
## Not run:
pqtl_list(flag = "outcomes")

## End(Not run)
```

pqtl_pleio

Return information related to the pleiotropy of SNPs

Description

GET /pqtl/pleio/

Usage

```
pqtl_pleio(
  rsid = NULL,
  prflag = c("proteins", "count"),
  mode = c("table", "raw")
)
```

Arguments

<code>rsid</code>	(Required) A SNP identified by rsID which cannot be NULL.
<code>prflag</code>	(Optional) A flag which determines whether the number (if "count") or names (if "proteins") of the associated proteins should be returned. The DEFAULT value is "proteins".
<code>mode</code>	(Optional) If <code>mode = "table"</code> , returns a data frame (a <code>tibble</code> as per <code>tidyverse</code> convention). If <code>mode = "raw"</code> , returns a raw response from EpiGraphDB API with minimal parsing done by <code>httr</code> .

Value

Data from GET /pqtl/pleio/

Examples

```
# Returns a data frame of associated proteins
## Not run:
pqtl_pleio(rsid = "rs1260326")

## End(Not run)

# Returns a number of associated proteins
## Not run:
pqtl_pleio(rsid = "rs1260326", prflag = "count")

## End(Not run)
```

protein_in_pathway

For the list of proteins, returns their associated pathway data

Description

POST /protein/in-pathway

Usage

```
protein_in_pathway(uniprot_id_list, mode = c("table", "raw"))
```

Arguments

uniprot_id_list	A list of protein UniProt IDs
mode	If mode = "table", returns a data frame (a tibble as per tidyverse convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by httr .

Value

Data from POST /protein/in-pathway

Examples

```
## Not run:
protein_in_pathway(uniprot_id_list = c("014933", "060674", "P32455"))

## End(Not run)
```

query_epigraphdb*Send data request to an EpiGraphDB API endpoint*

Description

This is a general purpose function to send data request which can be used when there has not been an R equivalent package function to an API endpoint. Underneath this is a wrapper around `httr` functions with better handling of returned status.

Usage

```
query_epigraphdb(  
  route,  
  params = NULL,  
  mode = c("raw", "table"),  
  method = c("GET", "POST"),  
  retry_times = 3,  
  retry_pause_min = 1  
)
```

Arguments

<code>route</code>	An EpiGraphDB API endpoint route, e.g. <code>"/mr"</code> or <code>"/confounder"</code> . Consult the EpiGraphDB API documentation .
<code>params</code>	A list of parameters associated with the query endpoint.
<code>mode</code>	<code>c("raw", "table")</code> , if <code>"table"</code> then the query handler will try to convert the returned data to a tibble dataframe. NOTE: The default mode is <code>"raw"</code> which will NOT convert the returned response to a dataframe. This is different to functions that query topic endpoints which default to return a dataframe. Explicitly specify <code>mode = "table"</code> when needed.
<code>method</code>	Type of HTTP (GET, POST, PUT, etc.) method. NOTE: When sending a POST request where a specific parameter is specified as a list on the API, and if the equivalent in R is a vector of length 1, you should wrap this parameter in <code>I()</code> , e.g. <code>I(c("APOE"))</code> to avoid auto unboxing. For details, please refer to httr::POST
<code>retry_times</code>	Number of times the function will retry the request to the API.
<code>retry_pause_min</code>	Minimum number of seconds to wait for the next retry.

Value

Data from an EpiGraphDB API endpoint.

Examples

```

# GET /mr
# equivalent to `mr(exposure_trait = "Body mass index", outcome_trait = "Coronary heart disease")`^
## Not run:
query_epigraphdb(
  route = "/mr",
  params = list(
    exposure_trait = "Body mass index",
    outcome_trait = "Coronary heart disease"
  ),
  mode = "table"
)

## End(Not run)

# GET /meta/nodes/Gwas/list
## Not run:
query_epigraphdb(
  route = "/meta/nodes/Gwas/list",
  params = list(
    limit = 5,
    offset = 0
  )
) %>% str(1)

## End(Not run)

# POST /protein/ppi
## Not run:
query_epigraphdb(
  route = "/protein/ppi",
  params = list(
    uniprot_id_list = c("P30793", "Q9NZM1", "095236")
  ),
  method = "POST"
)

## End(Not run)

# error handling
## Not run:
tryCatch(
  query_epigraphdb(
    route = "/mr",
    params = list(
      exposure_trait = NULL,
      outcome_trait = NULL
    ),
    retry_times = 0
  ),
  error = function(e) {
    message(e)
}

```

```

    }
)

## End(Not run)

```

xqtl_multi.snp_mr *Multi SNP QTL MR evidence*

Description

GET /xqtl/multi-snp-mr

Usage

```

xqtl_multi.snp_mr(
  exposure_gene = NULL,
  outcome_trait = NULL,
  mr_method = c("IVW", "Egger"),
  qtl_type = c("eQTL", "pQTL"),
  pval_threshold = 1e-05,
  mode = c("table", "raw")
)

```

Arguments

exposure_gene	Name of the exposure gene
outcome_trait	Name of the outcome trait
mr_method	"IVW" or "Egger"
qtl_type	"eQTL" or "pQTL"
pval_threshold	P-value threshold
mode	If mode = "table", returns a data frame (a tibble as per tidyverse convention). If mode = "raw", returns a raw response from EpiGraphDB API with minimal parsing done by httr .

Value

Data from GET /xqtl/multi-snp-mr

Examples

```

## Not run:
xqtl_multi.snp_mr(outcome_trait = "Coronary heart disease")

## End(Not run)

```

`xqtl_single.snp_mr` *Single SNP QTL MR evidence*

Description

`GET /xqtl/single-snp-mr`

Usage

```
xqtl_single.snp_mr(
  exposure_gene = NULL,
  outcome_trait = NULL,
  snp = NULL,
  qtl_type = c("eQTL", "pQTL"),
  pval_threshold = 1e-05,
  mode = c("table", "raw")
)
```

Arguments

<code>exposure_gene</code>	Name of the exposure gene
<code>outcome_trait</code>	Name of the outcome trait
<code>snp</code>	SNP rsid
<code>qtl_type</code>	"eQTL" or "pQTL"
<code>pval_threshold</code>	P-value threshold
<code>mode</code>	If <code>mode = "table"</code> , returns a data frame (a <code>tibble</code> as per <code>tidyverse</code> convention). If <code>mode = "raw"</code> , returns a raw response from EpiGraphDB API with minimal parsing done by <code>httr</code> .

Value

Data from GEET `/xqtl/single-snp-mr`

Examples

```
## Not run:
xqtl_single.snp_mr(outcome_trait = "Coronary heart disease")

## End(Not run)
```

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