

Package ‘OmnipathR’

March 29, 2022

Type Package

Title OmniPath web service client and more

Version 3.2.8

Description A client for the OmniPath web service
(<https://www.omnipathdb.org>) and many other resources.
It also includes functions to transform and pretty print
some of the downloaded data, functions to access a number
of other resources such as BioPlex, ConsensusPathDB, EVEX,
Gene Ontology, Guide to Pharmacology (IUPHAR/BPS), Harmonizome,
HTREdb, Human Phenotype Ontology, InWeb InBioMap, KEGG Pathway,
Pathway Commons, Ramiowski et al. 2015, RegNetwork, ReMap, TF
census, TRRUST and Vinayagam et al. 2011. Furthermore, OmnipathR
features a close integration with the NicheNet method for
ligand activity prediction from transcriptomics data, and its
R implementation `nichenetr` (available only on github).

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URL <https://saezlab.github.io/OmnipathR/>

BugReports <https://github.com/saezlab/OmnipathR/issues>

biocViews GraphAndNetwork, Network, Pathways, Software,
ThirdPartyClient, DataImport, DataRepresentation,
GeneSignaling, GeneRegulation, SystemsBiology, Transcriptomics,
SingleCell, Annotation, KEGG

Encoding UTF-8

VignetteBuilder knitr

Depends R(>= 4.0)

Imports checkmate, crayon, curl, digest, dplyr, httr, igraph,
jsonlite, later, logger, magrittr, progress, purrr, rappdirs,
readr(>= 2.0.0), readxl, rlang, stats, stringr, tibble, tidyverse,
tidyselect, tools, utils, xml2, yaml

Suggests BiocStyle, dnet, ggplot2, ggraph, gprofiler2, knitr, mlrMBO,
parallelMap, ParamHelpers, Rgraphviz, rmarkdown, smoof,
supraHex, testthat

RoxigenNote 7.1.2**git_url** <https://git.bioconductor.org/packages/OmnipathR>**git_branch** RELEASE_3_14**git_last_commit** f3fbd35**git_last_commit_date** 2022-02-23**Date/Publication** 2022-03-29**Author** Alberto Valdeolivas [aut] (<<https://orcid.org/0000-0001-5482-9023>>),Denes Turei [cre, aut] (<<https://orcid.org/0000-0002-7249-9379>>),Attila Gabor [aut] (<<https://orcid.org/0000-0002-0776-1182>>)**Maintainer** Denes Turei <turei.denes@gmail.com>**R topics documented:**

| | |
|--|----|
| .omnipath_options_defaults | 6 |
| all_uniprots | 6 |
| ancestors | 7 |
| annotated_network | 8 |
| annotation_categories | 9 |
| bioplex1 | 10 |
| bioplex2 | 10 |
| bioplex3 | 11 |
| bioplex_all | 12 |
| bioplex_hct116_1 | 13 |
| bma_motif_es | 14 |
| bma_motif_vs | 15 |
| consensuspathdb_download | 15 |
| consensuspathdb_raw_table | 16 |
| descendants | 17 |
| enzsub_graph | 18 |
| evex_download | 19 |
| filter_by_resource | 20 |
| filter_intercell_network | 20 |
| find_all_paths | 23 |
| get_annotation_resources | 24 |
| get_complex_genes | 25 |
| get_complex_resources | 26 |
| get_db | 26 |
| get_enzsub_resources | 27 |
| get_interaction_resources | 28 |
| get_intercell_categories | 29 |
| get_intercell_generic_categories | 29 |
| get_intercell_resources | 30 |
| get_ontology_db | 31 |
| get_resources | 32 |
| get_signed_ptms | 32 |

| | |
|--|----|
| giant_component | 33 |
| go_annot_download | 34 |
| go_annot_slim | 35 |
| go_ontology_download | 36 |
| guide2pharma_download | 37 |
| harmonizome_download | 38 |
| hpo_download | 39 |
| htridb_download | 39 |
| import_all_interactions | 40 |
| import_dorothea_interactions | 42 |
| import_intercell_network | 43 |
| import_kinaseextra_interactions | 46 |
| import_ligrecextra_interactions | 47 |
| import_lncrna_mrna_interactions | 49 |
| import_mirnatarget_interactions | 50 |
| import_omnipath_annotations | 51 |
| import_omnipath_complexes | 52 |
| import_omnipath_enzsub | 53 |
| import_omnipath_interactions | 54 |
| import_omnipath_intercell | 56 |
| import_pathwayextra_interactions | 58 |
| import_post_translational_interactions | 59 |
| import_tf_mirna_interactions | 60 |
| import_tf_target_interactions | 62 |
| import_transcriptional_interactions | 63 |
| inbiomap_download | 64 |
| inbiomap_raw | 65 |
| interaction_graph | 66 |
| intercell_categories | 67 |
| intercell_consensus_filter | 67 |
| is_ontology_id | 68 |
| kegg_info | 69 |
| kegg_open | 70 |
| kegg_pathways_download | 71 |
| kegg_pathway_annotations | 72 |
| kegg_pathway_download | 73 |
| kegg_pathway_list | 74 |
| kegg_picture | 75 |
| kegg_process | 76 |
| load_db | 77 |
| nichenet_build_model | 78 |
| nichenet_expression_data | 78 |
| nichenet_gr_network | 79 |
| nichenet_gr_network_evex | 80 |
| nichenet_gr_network_harmonizome | 81 |
| nichenet_gr_network_htridb | 82 |
| nichenet_gr_network_omnipath | 83 |
| nichenet_gr_network_pathwaycommons | 84 |

| | |
|---|-----|
| nichenet_gr_network_regnetwork | 84 |
| nichenet_gr_network_remap | 85 |
| nichenet_gr_network_trrust | 86 |
| nichenet_ligand_activities | 87 |
| nichenet_ligand_target_links | 88 |
| nichenet_ligand_target_matrix | 90 |
| nichenet_lr_network | 91 |
| nichenet_lr_network_guide2pharma | 92 |
| nichenet_lr_network_omnipath | 93 |
| nichenet_lr_network_ramilowski | 94 |
| nichenet_main | 95 |
| nichenet_networks | 97 |
| nichenet_optimization | 99 |
| nichenet_remove_orphan_ligands | 100 |
| nichenet_results_dir | 101 |
| nichenet_signaling_network | 101 |
| nichenet_signaling_network_cpdb | 103 |
| nichenet_signaling_network_evex | 104 |
| nichenet_signaling_network_harmonizome | 104 |
| nichenet_signaling_network_inbiomap | 105 |
| nichenet_signaling_network_omnipath | 106 |
| nichenet_signaling_network_pathwaycommons | 107 |
| nichenet_signaling_network_vinayagam | 107 |
| nichenet_test | 108 |
| nichenet_workarounds | 109 |
| obo_parser | 109 |
| OmnipathR | 111 |
| omnipath_cache_autoclean | 112 |
| omnipath_cache_clean | 112 |
| omnipath_cache_clean_db | 113 |
| omnipath_cache_download_ready | 114 |
| omnipath_cache_filter_versions | 115 |
| omnipath_cache_get | 116 |
| omnipath_cache_key | 117 |
| omnipath_cache_latest_or_new | 117 |
| omnipath_cache_latest_version | 119 |
| omnipath_cache_load | 119 |
| omnipath_cache_move_in | 120 |
| omnipath_cache_remove | 121 |
| omnipath_cache_save | 123 |
| omnipath_cache_search | 124 |
| omnipath_cache_set_ext | 125 |
| omnipath_cache_update_status | 126 |
| omnipath_cache_wipe | 127 |
| omnipath_get_config_path | 127 |
| omnipath_load_config | 128 |
| omnipath_log | 129 |
| omnipath_logfile | 129 |

| | |
|---|-----|
| omnipath_msg | 130 |
| omnipath_reset_config | 131 |
| omnipath_save_config | 131 |
| omnipath_set_cachedir | 132 |
| omnipath_set_console_loglevel | 133 |
| omnipath_set_logfile_loglevel | 134 |
| omnipath_set_loglevel | 134 |
| omnipath_show_db | 135 |
| omnipath_unlock_cache_db | 136 |
| ontology_ensure_id | 136 |
| ontology_ensure_name | 137 |
| ontology_name_id | 137 |
| pathwaycommons_download | 138 |
| pivot_annotations | 139 |
| preppi_download | 140 |
| preppi_filter | 141 |
| print_bma_motif_es | 142 |
| print_bma_motif_vs | 143 |
| print_interactions | 144 |
| print_path_es | 145 |
| print_path_vs | 146 |
| ramilowski_download | 147 |
| regnetwork_directions | 147 |
| regnetwork_download | 148 |
| relations_list_to_table | 149 |
| relations_table_to_graph | 150 |
| relations_table_to_list | 151 |
| remap_dorothea_download | 152 |
| remap_filtered | 153 |
| remap_tf_target_download | 154 |
| resources_colname | 155 |
| simplify_intercell_network | 155 |
| swap_relations | 156 |
| tfcensus_download | 157 |
| translate_ids | 158 |
| trrust_download | 160 |
| uniprot_full_id_mapping_table | 161 |
| uniprot_id_mapping_table | 162 |
| unique_intercell_network | 163 |
| vinayagam_download | 164 |
| walk_ontology_tree | 165 |
| zenodo_download | 166 |

.omnipath_options_defaults*Default values for the package options***Description**

These options describe the default settings for OmnipathR so you do not need to pass these parameters at each function call. Currently the only option useful for the public web service at omnipathdb.org is “omnipath.license”. If you are a for-profit user set it to “commercial” to make sure all the data you download from OmniPath is legally allowed for commercial use. Otherwise just leave it as it is: “academic”. If you don’t use omnipathdb.org but within your organization you deployed your own pypath server and want to share data with a limited availability to outside users, you may want to use a password. For this you can use the “omnipath.password” option. Also if you want the R package to work from another pypath server instead of omnipathdb.org, you can change the option “omnipath.url”.

Usage

```
.omnipath_options_defaults
```

Format

An object of class `list` of length 13.

all_uniprot*A table with all UniProt IDs***Description**

Retrieves a table from UniProt with all proteins for a certain organism.

Usage

```
all_uniprot(fields = "id", reviewed = TRUE, organism = 9606)
```

Arguments

| | |
|-----------------------|---|
| <code>fields</code> | Character vector of fields as defined by UniProt. For possible values please refer to https://www.uniprot.org/help/uniprotkb%5Fcolumn%5Fnames |
| <code>reviewed</code> | Retrieve only reviewed (‘TRUE’), only unreviewed (‘FALSE’) or both (‘NULL’). |
| <code>organism</code> | Integer, NCBI Taxonomy ID of the organism (by default 9606 for human). |

Value

Data frame (tibble) with the requested UniProt entries and fields.

Examples

```
human_swissprot_ac <- all_uniprot(fields = 'entry name')
human_swissprot_ac
# # A tibble: 20,396 x 1
#   `Entry name`<chr>
#   1 OR4K3_HUMAN
#   2 O52A1_HUMAN
#   3 O2AG1_HUMAN
#   4 O10S1_HUMAN
#   5 O11G2_HUMAN
# # . with 20,386 more rows
```

ancestors

All ancestors in the ontology tree

Description

Starting from the selected nodes, recursively walks the ontology tree until it reaches the root. Collects all visited nodes, which are the ancestors (parents) of the starting nodes.

Usage

```
ancestors(
  terms,
  db_key = "go_basic",
  ids = TRUE,
  relations = c("is_a", "part_of", "occurs_in", "regulates", "positively_regulates",
    "negatively_regulates")
)
```

Arguments

| | |
|------------------------|---|
| <code>terms</code> | Character vector of ontology term IDs or names. A mixture of IDs and names can be provided. |
| <code>db_key</code> | Character: key to identify the ontology database. For the available keys see omnipath_show_db . |
| <code>ids</code> | Logical: whether to return IDs or term names. |
| <code>relations</code> | Character vector of ontology relation types. Only these relations will be used. |

Details

Note: this function relies on the database manager, the first call might take long because of the database load process. Subsequent calls within a short period should be faster. See [get_ontology_db](#).

Value

Character vector of ontology IDs. If the input terms are all root nodes, NULL is returned. The starting nodes won't be included in the result unless some of them are ancestors of other starting nodes.

Examples

```
ancestors('GO:0005035', ids = FALSE)
# [1] "molecular_function"
# [2] "transmembrane signaling receptor activity"
# [3] "signaling receptor activity"
# [4] "molecular transducer activity"
```

annotated_network *Network interactions with annotations*

Description

Annotations are often useful in a network context, e.g. one might want to label the interacting partners by their pathway membership. This function takes a network data frame and joins an annotation data frame from both the left and the right side, so both the source and target molecular entities will be labeled by their annotations. If one entity has many annotations these will yield many rows, hence the interacting pairs won't be unique across the data frame any more. Also if one entity has really many annotations the resulting data frame might be huge, we recommend to be careful with that. Finally, if you want to do the same but with intercell annotations, there is the [import_intercell_network](#) function.

Usage

```
annotated_network(network = NULL, annot = NULL, ...)
```

Arguments

| | |
|---------|---|
| network | Behaviour depends on type: if list, will be passed as arguments to import_omnipath_interactions to obtain a network data frame; if a data frame or tibble, it will be used as a network data frame; if a character vector, will be assumed to be a set of resource names and interactions will be queried from these resources. |
| annot | Either the name of an annotation resource (for a list of available resources call get_annotation_resources), or an annotation data frame. If the data frame contains more than one resources, only the first one will be used. |
| ... | Column names selected from the annotation data frame (passed to <code>dplyr::select</code> , if empty all columns will be selected.) |

Value

A data frame of interactions with annotations for both interacting entities.

Examples

```
signalink_with_pathways <-  
  annotated_network('SignaLink3', 'SignaLink_pathway')
```

annotation_categories *Annotation categories and resources*

Description

A full list of annotation resources, keys and values.

Usage

```
annotation_categories()
```

Value

A data frame with resource names, annotation key labels and for each key all possible values.

Examples

```
annot_cat <- annotation_categories()  
annot_cat  
# # A tibble: 46,307 x 3  
#   source      label    value  
#   <chr>       <chr>    <chr>  
# 1 connectomeDB2020 role    ligand  
# 2 connectomeDB2020 role    receptor  
# 3 connectomeDB2020 location ECM  
# 4 connectomeDB2020 location plasma membrane  
# 5 connectomeDB2020 location secreted  
# 6 KEGG-PC     pathway  Alanine, aspartate and glutamate metabolism  
# 7 KEGG-PC     pathway  Amino sugar and nucleotide sugar metabolism  
# 8 KEGG-PC     pathway  Aminoacyl-tRNA biosynthesis  
# 9 KEGG-PC     pathway  Arachidonic acid metabolism  
# 10 KEGG-PC    pathway  Arginine and proline metabolism
```

bioplex1*Downloads the BioPlex version 1.0 interaction dataset***Description**

This dataset contains ~24,000 interactions detected in HEK293T cells using 2,594 baits. More details at <https://bioplex.hms.harvard.edu/interactions.php>.

Usage

```
bioplex1()
```

Value

Data frame (tibble) with interactions.

See Also

- [bioplex2](#)
- [bioplex3](#)
- [bioplex_hct116_1](#)
- [bioplex_all](#)

Examples

```
bioplex_interactions <- bioplex1()
nrow(bioplex_interactions)
# [1] 23744
colnames(bioplex_interactions)
# [1] "GeneA"          "GeneB"          "UniprotA"        "UniprotB"
# [5] "SymbolA"        "SymbolB"        "p_wrong"        "p_no_interaction"
# [9] "p_interaction"
```

bioplex2*Downloads the BioPlex version 2.0 interaction dataset***Description**

This dataset contains ~56,000 interactions detected in HEK293T cells using 5,891 baits. More details at <https://bioplex.hms.harvard.edu/interactions.php>

Usage

```
bioplex2()
```

Value

Data frame (tibble) with interactions.

See Also

- [bioplex1](#)
- [bioplex3](#)
- [bioplex_hct116_1](#)
- [bioplex_all](#)

Examples

```
bioplex_interactions <- bioplex2()
nrow(bioplex_interactions)
# [1] 56553
colnames(bioplex_interactions)
# [1] "GeneA"          "GeneB"          "UniprotA"        "UniprotB"
# [5] "SymbolA"        "SymbolB"        "p_wrong"        "p_no_interaction"
# [9] "p_interaction"
```

bioplex3

Downloads the BioPlex version 3.0 interaction dataset

Description

This dataset contains ~120,000 interactions detected in HEK293T cells using 10,128 baits. More details at <https://bioplex.hms.harvard.edu/interactions.php>.

Usage

```
bioplex3()
```

Value

Data frame (tibble) with interactions.

See Also

- [bioplex1](#)
- [bioplex2](#)
- [bioplex_hct116_1](#)
- [bioplex_all](#)

Examples

```
bioplex_interactions <- bioplex3()
nrow(bioplex_interactions)
# [1] 118162
colnames(bioplex_interactions)
# [1] "GeneA"          "GeneB"          "UniprotA"        "UniprotB"
# [5] "SymbolA"        "SymbolB"        "p_wrong"        "p_no_interaction"
# [9] "p_interaction"
```

bioplex_all

Downloads all BioPlex interaction datasets

Description

BioPlex provides four interaction datasets: version 1.0, 2.0, 3.0 and HCT116 version 1.0. This function downloads all of them, merges them to one data frame, removes the duplicates (based on unique pairs of UniProt IDs) and separates the isoform numbers from the UniProt IDs. More details at <https://bioplex.hms.harvard.edu/interactions.php>.

Usage

```
bioplex_all(unique = TRUE)
```

Arguments

| | |
|--------|--|
| unique | Logical. Collapse the duplicate interactions into single rows or keep them as they are. In case of merging duplicate records the maximum p value will be chosen for each record. |
|--------|--|

Value

Data frame (tibble) with interactions.

See Also

- [bioplex1](#)
- [bioplex2](#)
- [bioplex3](#)
- [bioplex_hct116_1](#)

Examples

```
bioplex_interactions <- bioplex_all()
bioplex_interactions
# # A tibble: 195,538 x 11
#   UniprotA IsoformA UniprotB IsoformB GeneA GeneB SymbolA SymbolB
#   <chr>      <int> <chr>      <int> <dbl> <dbl> <chr>    <chr>
# 1 A0AV02        2 Q5K4L6       NA 84561 11000 SLC12A8 SLC27A3
# 2 A0AV02        2 Q8N5V2       NA 84561 25791 SLC12A8 NGEF
# 3 A0AV02        2 Q9H6S3       NA 84561 64787 SLC12A8 EPS8L2
# 4 A0AV96        2 000425       2 54502 10643 RBM47  IGF2BP3
# 5 A0AV96        2 000443       NA 54502 5286  RBM47  PIK3C2A
# 6 A0AV96        2 043426       NA 54502 8867  RBM47  SYNJ1
# 7 A0AV96        2 075127       NA 54502 26024 RBM47  PTCD1
# 8 A0AV96        2 095208       2 54502 22905 RBM47  EPN2
# 9 A0AV96        2 095900       NA 54502 26995 RBM47  TRUB2
# 10 A0AV96       2 P07910       2 54502 3183  RBM47  HNRNPC
# # . with 195,528 more rows, and 3 more variables: p_wrong <dbl>,
# #   p_no_interaction <dbl>, p_interaction <dbl>
```

bioplex_hct116_1

Downloads the BioPlex HCT116 version 1.0 interaction dataset

Description

This dataset contains ~71,000 interactions detected in HCT116 cells using 5,522 baits. More details at <https://bioplex.hms.harvard.edu/interactions.php>.

Usage

```
bioplex_hct116_1()
```

Value

Data frame (tibble) with interactions.

See Also

- [bioplex1](#)
- [bioplex2](#)
- [bioplex3](#)
- [bioplex_all](#)

Examples

```
bioplex_interactions <- bioplex_hct116_1()
nrow(bioplex_interactions)
# [1] 70966
colnames(bioplex_interactions)
# [1] "GeneA"           "GeneB"           "UniprotA"        "UniprotB"
# [5] "SymbolA"         "SymbolB"         "p_wrong"         "p_no_interaction"
# [9] "p_interaction"
```

bma_motif_es

BMA motifs from a sequence of edges

Description

These motifs can be added to a BMA canvas.

Usage

```
bma_motif_es(edge_seq, G, granularity = 2)
```

Arguments

- edge_seq An igraph edge sequence.
- G An igraph graph object.
- granularity Numeric: granularity value.

Value

Character: BMA motifs as a single string.

Examples

```
interactions <- import_omnipath_interactions(resources = 'ARN')
graph <- interaction_graph(interactions)
motifs <- bma_motif_es(igraph::E(graph)[1], graph)
```

| | |
|--------------|--|
| bma_motif_vs | <i>Prints a BMA motif to the screen from a sequence of nodes, which can be copy/pasted into the BMA canvas</i> |
|--------------|--|

Description

Intended to parallel print_path_vs

Usage

```
bma_motif_vs(node_seq, G)
```

Arguments

- | | |
|----------|--------------------------|
| node_seq | An igraph node sequence. |
| G | An igraph graph object. |

Value

Character: BMA motifs as a single string.

Examples

```
interactions <- import_omnipath_interactions(resources = 'ARN')
graph <- interaction_graph(interactions)
bma_string <- bma_motif_vs(
  igraph::all_shortest_paths(
    graph,
    from = 'ULK1',
    to = 'ATG13'
  )$res,
  graph
)
```

| | |
|--------------------------|--|
| consensuspathdb_download | <i>Retrieves the ConsensusPathDB network</i> |
|--------------------------|--|

Description

Compiles a table of binary interactions from ConsensusPathDB (<http://cpdb.molgen.mpg.de/>) and translates the UniProtKB ACs to Gene Symbols.

Usage

```
consensuspathdb_download(complex_max_size = 4, min_score = 0.9)
```

Arguments

| | |
|------------------|--|
| complex_max_size | Numeric: do not expand complexes with a higher number of elements than this. ConsensusPathDB does not contain conventional interactions but lists of participants, which might be members of complexes. Some records include dozens of participants and expanding them to binary interactions result thousands, sometimes hundreds of thousands of interactions from one single record. At the end, this process consumes >10GB of memory and results rather unusable data, hence it is recommended to limit the complex sizes at some low number. |
| min_score | Numeric: each record in ConsensusPathDB comes with a confidence score, expressing the amount of evidences. The default value, a minimum score of 0.9 retains approx. the top 30 percent of the interactions. |

Value

Data frame (tibble) with interactions.

Examples

```
## Not run:
cpdb_data <- consensuspathdb_download(
  complex_max_size = 1,
  min_score = .99
)
nrow(cpdb_data)
# [1] 252302
colnames(cpdb_data)
# [1] "databases"    "references"   "uniprot_a"    "confidence"   "record_id"
# [6] "uniprot_b"    "in_complex"  "genesymbol_a" "genesymbol_b"
cpdb_data
# # A tibble: 252,302 x 9
#   databases references uniprot_a confidence record_id uniprot_b in_com
#   <chr>      <chr>      <chr>        <dbl>     <int> <chr>    <lgl>
# 1 Reactome  NA        SUMF2_HU.     1         1 SUMF1_HU. TRUE
# 2 Reactome  NA        SUMF1_HU.     1         1 SUMF2_HU. TRUE
# 3 DIP,Reac. 22210847,. STIM1_HU.   0.998     2 TRPC1_HU. TRUE
# 4 DIP,Reac. 22210847,. TRPC1_HU.   0.998     2 STIM1_HU. TRUE
# # . with 252,292 more rows, and 2 more variables: genesymbol_a <chr>,
# #   genesymbol_b <chr>
## End(Not run)
```

Description

Downloads interaction data from ConsensusPathDB

Usage

```
consensuspathdb_raw_table()
```

Value

Data frame (tibble) with interactions.

Examples

```
cpdb_raw <- consensuspathdb_raw_table()
```

descendants

All descendants in the ontology tree

Description

Starting from the selected nodes, recursively walks the ontology tree until it reaches the leaf nodes. Collects all visited nodes, which are the descendants (children) of the starting nodes.

Usage

```
descendants(
  terms,
  db_key = "go_basic",
  ids = TRUE,
  relations = c("is_a", "part_of", "occurs_in", "regulates", "positively_regulates",
    "negatively_regulates")
)
```

Arguments

| | |
|-----------|---|
| terms | Character vector of ontology term IDs or names. A mixture of IDs and names can be provided. |
| db_key | Character: key to identify the ontology database. For the available keys see omnipath_show_db . |
| ids | Logical: whether to return IDs or term names. |
| relations | Character vector of ontology relation types. Only these relations will be used. |

Details

Note: this function relies on the database manager, the first call might take long because of the database load process. Subsequent calls within a short period should be faster. See [get_ontology_db](#).

Value

Character vector of ontology IDs. If the input terms are all leaves NULL is returned. The starting nodes won't be included in the result unless some of them are descendants of other starting nodes.

Examples

```
descendants('GO:0005035', ids = FALSE)
# [1] "tumor necrosis factor-activated receptor activity"
# [2] "TRAIL receptor activity"
# [3] "TNFSF11 receptor activity"
```

enzsub_graph

*Enzyme-substrate graph***Description**

Transforms the a data frame with enzyme-substrate relationships (obtained by [import_omnipath_enzsub](#)) to an igraph graph object.

Usage

```
enzsub_graph(enzsub)
```

Arguments

| | |
|--------|--|
| enzsub | Data frame created by import_omnipath_enzsub |
|--------|--|

Value

An igraph directed graph object.

See Also

- [import_omnipath_enzsub](#)
- [giant_component](#)
- [find_all_paths](#)

Examples

```
enzsub <- import_omnipath_enzsub(resources = c('PhosphoSite', 'SIGNOR'))
enzsub_g <- enzsub_graph(enzsub = enzsub)
```

Description

Downloads interactions from EVEX, a versatile text mining resource (<http://evexdb.org>). Translates the Entrez Gene IDs to Gene Symbols and combines the interactions and references into a single data frame.

Usage

```
evex_download(
  min_confidence = NULL,
  remove_negatives = TRUE,
  top_confidence = NULL
)
```

Arguments

| | |
|-------------------------------|--|
| <code>min_confidence</code> | Numeric: a threshold for confidence scores. EVEX confidence scores span roughly from -3 to 3. By providing a numeric value in this range the lower confidence interactions can be removed. If NULL no filtering performed. |
| <code>remove_negatives</code> | Logical: remove the records with the "negation" attribute set. |
| <code>top_confidence</code> | Confidence cutoff as quantile (a number between 0 and 1). If NULL no filtering performed. |

Value

Data frame (tibble) with interactions.

Examples

```
evex_interactions <- evex_download()
evex_interactions
# # A tibble: 368,297 x 13
#   general_event_id source_entrezge. target_entrezge. confidence negation
#   <dbl> <chr>           <chr>           <dbl>      <dbl>
# 1 98    8651          6774            -1.45      0
# 2 100   8431          6774            -1.45      0
# 3 205   6261          6263            0.370     0
# 4 435   1044          1045            -1.09     0
# . with 368,287 more rows, and 8 more variables: speculation <dbl>,
#   coarse_type <chr>, coarse_polarity <chr>, refined_type <chr>,
#   refined_polarity <chr>, source_genesymbol <chr>,
#   target_genesymbol <chr>, references <chr>
```

filter_by_resource*Filters OmniPath data by resources***Description**

Keeps only those records which are supported by any of the resources of interest.

Usage

```
filter_by_resource(data, resources = NULL)
```

Arguments

- | | |
|-----------|--|
| data | A data frame downloaded from the OmniPath web service (interactions, enzyme-substrate or complexes). |
| resources | Character vector with resource names to keep. |

Value

The data frame filtered.

Examples

```
interactions <- import_omnipath_interactions()
signor <- filter_by_resource(interactions, resources = 'SIGNOR')
```

filter_intercell_network*Quality filter an intercell network***Description**

The intercell database of OmniPath covers a very broad range of possible ways of cell to cell communication, and the pieces of information, such as localization, topology, function and interaction, are combined from many, often independent sources. This unavoidably result some weird and unexpected combinations which are false positives in the context of intercellular communication. [import_intercell_network](#) provides a shortcut (`high_confidence`) to do basic quality filtering. For custom filtering or experimentation with the parameters we offer this function.

Usage

```
filter_intercell_network(
  network,
  transmitter_topology = c("secreted", "plasma_membrane_transmembrane",
    "plasma_membrane_peripheral"),
  receiver_topology = "plasma_membrane_transmembrane",
  min_curation_effort = 2,
  min_resources = 1,
  min_references = 0,
  min_provenances = 1,
  consensus_percentile = 50,
  loc_consensus_percentile = 30,
  ligand_receptor = FALSE,
  simplify = FALSE,
  unique_pairs = FALSE,
  omnipath = TRUE,
  ligrecrextra = TRUE,
  kinaseextra = FALSE,
  pathwayextra = FALSE,
  ...
)
```

Arguments

| | |
|-----------------------------------|---|
| <code>network</code> | An intercell network data frame, as provided by <code>import_intercell_network</code> , without <code>simplify</code> . |
| <code>transmitter_topology</code> | Character vector: topologies allowed for the entities in transmitter role. Abbreviations allowed: "sec", "pmtm" and "pmp". |
| <code>receiver_topology</code> | Same as <code>transmitter_topology</code> for the entities in the receiver role. |
| <code>min_curation_effort</code> | Numeric: a minimum value of curation effort (resource-reference pairs) for network interactions. Use zero to disable filtering. |
| <code>min_resources</code> | Numeric: minimum number of resources for interactions. The value 1 means no filtering. |
| <code>min_references</code> | Numeric: minimum number of references for interactions. Use zero to disable filtering. |
| <code>min_provenances</code> | Numeric: minimum number of provenances (either resources or references) for interactions. Use zero or one to disable filtering. |
| <code>consensus_percentile</code> | Numeric: percentile threshold for the consensus score of generic categories in intercell annotations. The consensus score is the number of resources supporting the classification of an entity into a category based on combined information of many resources. Here you can apply a cut-off, keeping only the annotations supported by a higher number of resources than a certain percentile of each |

category. If NULL no filtering will be performed. The value is either in the 0-1 range, or will be divided by 100 if greater than 1. The percentiles will be calculated against the generic composite categories and then will be applied to their resource specific annotations and specific child categories.

loc_consensus_percentile

Numeric: similar to `consensus_percentile` for major localizations. For example, with a value of 50, the secreted, plasma membrane transmembrane or peripheral attributes will be TRUE only where at least 50 percent of the resources support these.

ligand_receptor

Logical. If TRUE, only *ligand* and *receptor* annotations will be used instead of the more generic *transmitter* and *receiver* categories.

simplify

Logical: keep only the most often used columns. This function combines a network data frame with two copies of the intercell annotation data frames, all of them already having quite some columns. With this option we keep only the names of the interacting pair, their intercellular communication roles, and the minimal information of the origin of both the interaction and the annotations.

unique_pairs

Logical: instead of having separate rows for each pair of annotations, drop the annotations and reduce the data frame to unique interacting pairs. See [unique_intercell_network](#) for details.

omnipath

Logical: shortcut to include the *omnipath* dataset in the interactions query.

ligrecextra

Logical: shortcut to include the *ligrecextra* dataset in the interactions query.

kinaseextra

Logical: shortcut to include the *kinaseextra* dataset in the interactions query.

pathwayextra

Logical: shortcut to include the *pathwayextra* dataset in the interactions query.

...

If `simplify` or `unique_pairs` is TRUE, additional column names can be passed here to `dplyr::select` on the final data frame. Otherwise ignored.

Value

An intercell network data frame filtered.

See Also

- [import_intercell_network](#)
- [unique_intercell_network](#)
- [simplify_intercell_network](#)

Examples

```
icn <- import_intercell_network()
icn_f <- filter_intercell_network(
  icn,
  consensus_percentile = 75,
  min_provenances = 3,
  simplify = TRUE
)
```

| | |
|----------------|---|
| find_all_paths | <i>All paths between two groups of vertices</i> |
|----------------|---|

Description

Finds all paths up to length ‘maxlen’ between specified groups of vertices. This function is needed only because igraph’s ‘all_shortest_paths’ finds only the shortest, not any path up to a defined length.

Usage

```
find_all_paths(  
  graph,  
  start,  
  end,  
  attr = NULL,  
  mode = 'OUT',  
  maxlen = 2,  
  progress = TRUE  
)
```

Arguments

| | |
|----------|---|
| graph | An igraph graph object. |
| start | Integer or character vector with the indices or names of one or more start vertices. |
| end | Integer or character vector with the indices or names of one or more end vertices. |
| attr | Character: name of the vertex attribute to identify the vertices by. Necessary if ‘start’ and ‘end’ are not igraph vertex ids but for example vertex names or labels. |
| mode | Character: IN, OUT or ALL. Default is OUT. |
| maxlen | Integer: maximum length of paths in steps, i.e. if maxlen = 3, then the longest path may consist of 3 edges and 4 nodes. |
| progress | Logical: show a progress bar. |

Value

List of vertex paths, each path is a character or integer vector.

See Also

- [interaction_graph](#)
- [enzsub_graph](#)
- [giant_component](#)

Examples

```
interactions <- import_omnipath_interactions()
graph <- interaction_graph(interactions)
paths <- find_all_paths(
  graph = graph,
  start = c('EGFR', 'STAT3'),
  end = c('AKT1', 'ULK1'),
  attr = 'name'
)
```

get_annotation_resources

Retrieves a list of available resources in the annotations database of OmniPath

Description

Get the names of the resources from <https://omnipath.org/annotations>.

Usage

```
get_annotation_resources(dataset = NULL, ...)
```

Arguments

- | | |
|---------|-------------------------------|
| dataset | ignored for this query type |
| ... | optional additional arguments |

Value

character vector with the names of the annotation resources

See Also

- [get_resources](#)
- [import_omnipath_annotations](#)

Examples

```
get_annotation_resources()
```

get_complex_genes *Get all the molecular complexes for a given gene(s)*

Description

This function returns all the molecular complexes where an input set of genes participate. User can choose to retrieve every complex where any of the input genes participate or just retrieve these complexes where all the genes in input set participate together.

Usage

```
get_complex_genes(  
  complexes = import_omnipath_complexes(),  
  select_genes,  
  total_match = FALSE  
)
```

Arguments

| | |
|--------------|--|
| complexes | complexes data frame (obtained using import_omnipath_complexes) |
| select_genes | vector containing the genes for whom complexes will be retrieved (hgnc format). |
| total_match | [default=FALSE] logical indicating if the user wants to get all the complexes where any of the input genes participate (FALSE) or to get only the complexes where all the input genes participate together (TRUE). |

Value

Data frame of complexes

See Also

[import_omnipath_complexes](#)

Examples

```
complexes <- import_omnipath_complexes(  
  filter_databases = c("CORUM", "hu.MAP")  
)  
query_genes <- c("LMNA", "BANF1")  
complexes_query_genes <- get_complex_genes(complexes, query_genes)
```

`get_complex_resources` *Retrieve a list of complex resources available in Omnipath*

Description

Get the names of the resources from <https://omnipath.org/complexes>

Usage

```
get_complex_resources(dataset = NULL)
```

Arguments

| | |
|---------|-----------------------------|
| dataset | ignored for this query type |
|---------|-----------------------------|

Value

character vector with the names of the databases

See Also

- [get_resources](#)
- [import_omnipath_complexes](#)

Examples

```
get_complex_resources()
```

`get_db` *Access a built in database*

Description

Databases are resources which might be costly to load but can be used many times by functions which usually automatically load and retrieve them from the database manager. Each database has a lifetime and will be unloaded automatically upon expiry.

Usage

```
get_db(key, param = NULL, reload = FALSE)
```

Arguments

| | |
|--------|--|
| key | Character: the key of the database to load. For a list of available keys see omnipath_show_db . |
| param | List: override the defaults or pass further parameters to the database loader function. See the loader functions and their default parameters in omnipath_show_db . If the database is already loaded with different parameters it will be reloaded with the new parameters only if the reload option is TRUE. |
| reload | Reload the database if param passed here is different from the parameters used the last time the database was loaded. If different functions with different parameters access the database repeatedly and request reload the frequent reloads might cost substantial time and resource use. |

Value

An object with the database contents. The exact format depends on the database, most often it is a data frame or a list.

See Also

[omnipath_show_db](#).

Examples

```
goslim <- get_db('go_slim')
```

get_enzsub_resources *Retrieves a list of enzyme-substrate resources available in OmniPath*

Description

Get the names of the enzyme-substrate relationship resources available in <https://omnipath.org/enzsub>

Usage

```
get_enzsub_resources(dataset = NULL)
```

Arguments

| | |
|---------|-----------------------------|
| dataset | ignored for this query type |
|---------|-----------------------------|

Value

character vector with the names of the enzyme-substrate resources

See Also

- [get_resources](#)
- [import_omnipath_enzsub](#)

Examples

```
get_enzsub_resources()
```

get_interaction_resources

Retrieve a list of interaction resources available in Omnipath

Description

Gets the names of the resources from <https://omnipath.org/interactions>.

Usage

```
get_interaction_resources(dataset = NULL)
```

Arguments

- | | |
|---------|--|
| dataset | a dataset within the interactions query type. Currently available datasets are ‘omnipath’, ‘kinaseextra’, ‘pathwayextra’, ‘ligrecrextra’, ‘dorothea’, ‘tf_target’, ‘tf_mirna’, ‘mirnatarget’ and ‘lncrna_mrna’ |
|---------|--|

Value

character vector with the names of the interaction databases

See Also

- [get_resources](#)
- [import_all_interactions](#)
- [import_omnipath_interactions](#)
- [import_pathwayextra_interactions](#)
- [import_kinaseextra_interactions](#)
- [import_ligrecrextra_interactions](#)
- [import_mirnatarget_interactions](#)
- [import_dorothea_interactions](#)

Examples

```
get_interaction_resources()
```

get_intercell_categories

Categories in the intercell database of OmniPath

Description

Retrieves a list of categories from <https://omnipath.org/intercell>.

Usage

```
get_intercell_categories()
```

Value

character vector with the different intercell categories

See Also

- [import_omnipath_intercell](#)
- [get_intercell_generic_categories](#)

Examples

```
get_intercell_categories()
```

get_intercell_generic_categories

Retrieves a list of the generic categories in the intercell database of OmniPath

Description

Retrieves a list of the generic categories from <https://omnipath.org/intercell>.

Usage

```
get_intercell_generic_categories()
```

Value

character vector with the different intercell main classes

See Also

- [import_omnipath_intercell](#)
- [get_intercell_categories](#)

Examples

```
get_intercell_generic_categories()
```

get_intercell_resources

Retrieves a list of intercellular communication resources available in OmniPath

Description

Retrieves a list of the databases from <https://omnipath.org/intercell>.

Usage

```
get_intercell_resources(dataset = NULL)
```

Arguments

| | |
|---------|----------------------------|
| dataset | ignored at this query type |
|---------|----------------------------|

Value

character vector with the names of the databases

See Also

- [get_resources](#)
- [import_omnipath_intercell](#)

Examples

```
get_intercell_resources()
```

get_ontology_db *Access an ontology database*

Description

Retrieves an ontology database with relations in the desired data structure. The database is automatically loaded and the requested data structure is constructed if necessary. The databases stay loaded up to a certain time period (see the option `omnipath.db_lifetime`). Hence the first one of repeated calls to this function might take long and the subsequent ones should be really quick.

Usage

```
get_ontology_db(key, rel_fmt = "tbl", child_parents = TRUE)
```

Arguments

- | | |
|---------------|--|
| key | Character: key of the ontology database. For the available keys see omnipath_show_db . |
| rel_fmt | Character: the data structure of the ontology relations. Possible values are 1) "tbl" a data frame, 2) "lst" a list or 3) "gra" a graph. |
| child_parents | Logical: whether the ontology relations should point from child to parents (TRUE) or from parent to children (FALSE). |

Value

A list with the following elements: 1) "names" a table with term IDs and names; 2) "namespaces" a table to connect term IDs and namespaces they belong to; 3) "relations" a table with relations between terms and their parent terms; 4) "subsets" a table with terms and the subsets they are part of; 5) "obsolete" character vector with all the terms labeled as obsolete.

See Also

- [omnipath_show_db](#)
- [get_db](#)

Examples

```
go <- get_ontology_db('go_basic', child_parents = FALSE)
```

| | |
|---------------|--|
| get_resources | <i>Retrieve the available resources for a given query type</i> |
|---------------|--|

Description

Collects the names of the resources available in OmniPath for a certain query type and optionally for a dataset within that.

Usage

```
get_resources(query_type, datasets = NULL, generic_categories = NULL)
```

Arguments

| | |
|--------------------|--|
| query_type | one of the query types ‘interactions’, ‘enz_sub’, ‘complexes’, ‘annotations’ or ‘intercell’ |
| datasets | currently within the ‘interactions’ query type only, multiple datasets are available: ‘omnipath’, ‘kinaseextra’, ‘pathwayextra’, ‘ligrecrextra’, ‘dorothea’, ‘tf_target’, ‘tf_mirna’, ‘mirnatarget’ and ‘lncrna_mrna’. |
| generic_categories | for the ‘intercell’ query type, restrict the search for some generic categories e.g. ‘ligand’ or ‘receptor’. |

Value

a character vector with resource names

Examples

```
get_resources(query_type = 'interactions')
```

| | |
|-----------------|--|
| get_signed_ptms | <i>Signs for enzyme-substrate interactions</i> |
|-----------------|--|

Description

Enzyme-substrate data does not contain sign (activation/inhibition), we generate this information based on the interaction network.

Usage

```
get_signed_ptms(
  enzsub = import_omnipath_enzsub(),
  interactions = import_omnipath_interactions()
)
```

Arguments

- | | |
|--------------|--|
| enzsub | Enzyme-substrate data frame generated by import_omnipath_enzsub |
| interactions | interaction data frame generated by import_omnipath_interactions |

Value

Data frame of enzyme-substrate relationships with is_inhibition and is_stimulation columns.

See Also

- [import_omnipath_enzsub](#)
- [import_omnipath_interactions](#)

Examples

```
enzsub <- import_omnipath_enzsub(resources = c('PhosphoSite', 'SIGNOR'))
interactions <- import_omnipath_interactions()
enzsub <- get_signed_ptms(enzsub, interactions)
```

| | |
|-----------------|-----------------------------------|
| giant_component | <i>Giant component of a graph</i> |
|-----------------|-----------------------------------|

Description

For an igraph graph object returns its giant component.

Usage

```
giant_component(graph)
```

Arguments

- | | |
|-------|-------------------------|
| graph | An igraph graph object. |
|-------|-------------------------|

Value

An igraph graph object containing only the giant component.

Examples

```
interactions <- import_post_translational_interactions()
graph <- interaction_graph(interactions)
graph_gc <- giant_component(graph)
```

go_annot_download *Gene annotations from Gene Ontology*

Description

Gene Ontology is an ontology of gene subcellular localizations, molecular functions and involvement in biological processes. Gene products across many organisms are annotated with the ontology terms. This function downloads the gene-ontology term associations for certain model organisms or all organisms. For a description of the columns see <http://geneontology.org/docs/go-annotation-file-gaf-format-2.2/>.

Usage

```
go_annot_download(organism = "human", aspects = c("C", "F", "P"), slim = NULL)
```

Arguments

| | |
|----------|--|
| organism | Character: either "chicken", "cow", "dog", "human", "pig" or "uniprot_all". |
| aspects | Character vector with some of the following elements: "C" (cellular component), "F" (molecular function) and "P" (biological process). Gene Ontology is three separate ontologies called as three aspects. By this parameter you can control which aspects to include in the output. |
| slim | Character: if not NULL, the name of a GOsubset (slim). instead of the full GO annotation, the slim annotation will be returned. See details at go_annot_slim . If TRUE, the "generic" slim will be used. |

Value

A tibble (data frame) of annotations as it is provided by the database

Examples

```
goa_data <- go_annot_download()
goa_data
# # A tibble: 606,840 x 17
#   db      db_object_id db_object_symbol qualifier go_id    db_ref
#   <fct>   <chr>       <chr>           <fct>    <chr>   <chr>
# 1 UniProt A0A024RBG1  NUDT4B          NA        GO:000... GO_REF:00.
# 2 UniProt A0A024RBG1  NUDT4B          NA        GO:000... GO_REF:00.
# 3 UniProt A0A024RBG1  NUDT4B          NA        GO:004... GO_REF:00.
# 4 UniProt A0A024RBG1  NUDT4B          NA        GO:005... GO_REF:00.
# 5 UniProt A0A024RBG1  NUDT4B          NA        GO:005... GO_REF:00.
# # . with 606,830 more rows, and 11 more variables:
# #   evidence_code <fct>, with_or_from <chr>, aspect <fct>,
# #   db_object_name <chr>, db_object_synonym <chr>,
# #   db_object_type <fct>, taxon <fct>, date <date>,
# #   assigned_by <fct>, annotation_extension <chr>,
# #   gene_product_from_id <chr>
```

go_annot_slim *GO slim gene annotations*

Description

GO slims are subsets of the full GO which "give a broad overview of the ontology content without the detail of the specific fine grained terms". In order to annotate genes with GO slim terms, we take the annotations and search all ancestors of the terms up to the root of the ontology tree. From the ancestors we select the terms which are part of the slim subset.

Usage

```
go_annot_slim(  
  organism = "human",  
  slim = "generic",  
  aspects = c("C", "F", "P"),  
  cache = TRUE  
)
```

Arguments

| | |
|----------|--|
| organism | Character: either "chicken", "cow", "dog", "human", "pig" or "uniprot_all". |
| slim | Character: the GO subset (GO slim) name. Available GO slims are: "agr" (Alliance for Genomics Resources), "generic", "aspergillus", "candida", "drosophila", "chembl", "metagenomic", "mouse", "plant", "pir" (Protein Information Resource), "pombe" and "yeast". |
| aspects | Character vector with some of the following elements: "C" (cellular component), "F" (molecular function) and "P" (biological process). Gene Ontology is three separate ontologies called as three aspects. By this parameter you can control which aspects to include in the output. |
| cache | Logical: Load the result from cache if available. |

Details

Building the GO slim is resource intensive in its current implementation. For human annotation and generic GO slim it might take around 20 minutes. The result is saved into the cache so next time loading the data from there is really quick. If the cache option is FALSE the data will be built fresh (the annotation and ontology files still might come from cache), and the newly build GO slim will overwrite the cache instance.

Value

A tibble (data frame) of genes annotated with ontology terms in in the GO slim (subset).

See Also

- [go_annot_download](#)
- [go_ontology_download](#)
- [get_db](#)

Examples

```
goslim <- go_annot_slim(organism = 'human', slim = 'generic')
goslim
# # A tibble: 276,371 x 8
#   db      db_object_id db_object_symbol go_id aspect db_object_name
#   <fct>  <chr>        <chr>          <chr> <fct>  <chr>
# 1 UniPr. A0A024RBG1  NUDT4B           GO:0. F    Diphosphoinosito.
# 2 UniPr. A0A024RBG1  NUDT4B           GO:0. F    Diphosphoinosito.
# 3 UniPr. A0A024RBG1  NUDT4B           GO:0. C    Diphosphoinosito.
# 4 UniPr. A0A024RBG1  NUDT4B           GO:0. C    Diphosphoinosito.
# 5 UniPr. A0A024RBG1  NUDT4B           GO:0. C    Diphosphoinosito.
# # . with 276,366 more rows, and 2 more variables:
# #   db_object_synonym <chr>, db_object_type <fct>
```

[go_ontology_download](#) *The Gene Ontology tree*

Description

The Gene Ontology tree

Usage

```
go_ontology_download(
  basic = TRUE,
  tables = TRUE,
  subset = NULL,
  relations = c("is_a", "part_of", "occurs_in", "regulates", "positively_regulates",
               "negatively_regulates")
)
```

Arguments

- basic** Logical: use the basic or the full version of GO. As written on the GO home page: "the basic version of the GO is filtered such that the graph is guaranteed to be acyclic and annotations can be propagated up the graph. The relations included are is a, part of, regulates, negatively regulates and positively regulates. This version excludes relationships that cross the 3 GO hierarchies. This version should be used with most GO-based annotation tools."

| | |
|------------------|--|
| tables | In the result return data frames or nested lists. These later can be converted to each other if necessary. However converting from table to list is faster. |
| subset | Character: the GO subset (GO slim) name. GO slims are subsets of the full GO which "give a broad overview of the ontology content without the detail of the specific fine grained terms". This option, if not NULL, overrides the basic parameter. Available GO slims are: "agr" (Alliance for Genomics Resources), "generic", "aspergillus", "candida", "drosophila", "chembl", "metagenomic", "mouse", "plant", "pir" (Protein Information Resource), "pombe" and "yeast". |
| relations | Character vector: the relations to include in the processed data. |

Value

A list with the following elements: 1) "names" a list with terms as names and names as values; 2) "namespaces" a list with terms as names and namespaces as values; 3) "relations" a list with relations between terms: terms are keys, values are lists with relations as names and character vectors of related terms as values; 4) "subsets" a list with terms as keys and character vectors of subset names as values (or NULL if the term does not belong to any subset); 5) "obsolete" character vector with all the terms labeled as obsolete. If the **tables** parameter is TRUE, "names", "namespaces", "relations" and "subsets" will be data frames (tibbles).

Examples

```
# retrieve the generic GO slim, a small subset of the full ontology
go <- go_ontology_download(subset = 'generic')
```

guide2pharma_download *Downloads interactions from the Guide to Pharmacology database*

Description

Downloads ligand-receptor interactions from the Guide to Pharmacology (IUPHAR/BPS) database (<https://www.guidetopharmacology.org/>).

Usage

```
guide2pharma_download()
```

Value

A tibble (data frame) of interactions as it is provided by the database

Examples

```
g2p_data <- guide2pharma_download()
g2p_data
# # A tibble: 21,586 x 38
#   target target_id target_gene_sym. target_uniprot target_ensembl_.
#   <chr>     <dbl> <chr>           <chr>           <chr>
# 1 12S-L.      1387 ALOX12        P18054        ENSG00000108839
# 2 15-L0.      1388 ALOX15        P16050        ENSG00000161905
# 3 15-L0.      1388 ALOX15        P16050        ENSG00000161905
# 4 15-L0.      1388 ALOX15        P16050        ENSG00000161905
# # . with 21,576 more rows, and 33 more variables: target_ligand <chr>,
# #   target_ligand_id <chr>, target_ligand_gene_symbol <chr>,
# ... (truncated)
```

harmonizome_download *Downloads a Harmonizome network dataset*

Description

Downloads a single network dataset from Harmonizome <https://maayanlab.cloud/Harmonizome>.

Usage

```
harmonizome_download(dataset)
```

Arguments

| | |
|---------|---|
| dataset | The dataset part of the URL. Please refer to the download section of the Harmonizome webpage. |
|---------|---|

Value

Data frame (tibble) with interactions.

Examples

```
harmonizome_data <- harmonizome_download('phosphositeplus')
harmonizome_data
# # A tibble: 6,013 x 7
#   source  source_desc source_id target target_desc target_id weight
#   <chr>    <chr>       <dbl> <chr>    <chr>       <dbl> <dbl>
# 1 TP53     na          7157 STK17A na        9263     1
# 2 TP53     na          7157 TP53RK na       112858    1
# 3 TP53     na          7157 SMG1    na        23049    1
# 4 UPF1     na          5976 SMG1    na        23049    1
# # . with 6,003 more rows
```

hpo_download*Downloads protein annotations from Human Phenotype Ontology*

Description

Human Phenotype Ontology (HPO) provides a standardized vocabulary of phenotypic abnormalities encountered in human disease. Each term in the HPO describes a phenotypic abnormality. HPO currently contains over 13,000 terms and over 156,000 annotations to hereditary diseases. See more at <https://hpo.jax.org/app/>.

Usage

```
hpo_download()
```

Value

A tibble (data frame) of annotations as it is provided by the database

Examples

```
hpo_data <- hpo_download()
hpo_data
# # A tibble: 231,738 x 9
#   entrez_gene_id entrez_gene_symb. hpo_term_id hpo_term_name
#   <dbl> <chr>           <chr>          <chr>
# 1       8192 CLPP          HP:0000013 Hypoplasia of the ute.
# 2       8192 CLPP          HP:0004322 Short stature
# 3       8192 CLPP          HP:0000786 Primary amenorrhea
# 4       8192 CLPP          HP:0000007 Autosomal recessive i.
# 5       8192 CLPP          HP:0000815 Hypergonadotropic hyp.
# # . with 231,733 more rows, and 5 more variables:
# #   frequency_raw <chr>, frequency_hpo <chr>, info_gd_source <chr>,
# #   gd_source <chr>, disease_id <chr>
```

htridb_download*Downloads TF-target interactions from HTRIdb*

Description

HTRIdb (<https://www.lbhc.ibb.unesp.br/htri/>) is a database of literature curated human TF-target interactions. As the database is recently offline, the data is distributed by the OmniPath rescued data repository (<https://rescued.omnipathdb.org/>).

Usage

```
htridb_download()
```

Value

Data frame (tibble) with interactions.

Examples

```
htridb_data <- htridb_download()
htridb_data
# # A tibble: 18,630 x 7
#   OID GENEID_TF SYMBOL_TF GENEID_TG SYMBOL_TG TECHNIQUE
#   <dbl> <dbl> <chr>     <dbl> <chr>      <chr>
# 1 32399    142 PARP1      675 BRCA2 Electrophoretic Mobi.
# 2 32399    142 PARP1      675 BRCA2 Chromatin Immunoprec.
# 3 28907     196 AHR       1543 CYP1A1 Chromatin Immunoprec.
# 4 29466     196 AHR       1543 CYP1A1 Electrophoretic Mobi.
# 5 28911     196 AHR       1543 CYP1A1 Chromatin Immunoprec.
# # . with 18,620 more rows, and 1 more variable: PUBMED_ID <chr>
```

import_all_interactions

Imports all interaction datasets available in OmniPath

Description

The interaction datasets currently available in OmniPath:

Usage

```
import_all_interactions(
  resources = NULL,
  organism = 9606,
  dorothaea_levels = c("A", "B"),
  exclude = NULL,
  fields = NULL,
  default_fields = TRUE,
  references_by_resource = TRUE,
  ...
)
import_AllInteractions(...)
```

Arguments

| | |
|------------------------|---|
| <code>resources</code> | interactions not reported in these databases are removed. See get_interaction_resources for more information. |
| <code>organism</code> | Interactions are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse |

| | |
|------------------------|---|
| dorothea_levels | The confidence levels of the dorothea interactions (TF-target) which range from A to D. Set to A and B by default. |
| exclude | Character: datasets or resources to exclude. |
| fields | The user can define here the fields to be added. If used, set the next argument, ‘default_fields’, to FALSE. |
| default_fields | whether to include the default fields (columns) for the query type. If FALSE, only the fields defined by the user in the ‘fields’ argument will be added. |
| references_by_resource | if FALSE, removes the resource name prefixes from the references (PubMed IDs); this way the information which reference comes from which resource will be lost and the PubMed IDs will be unique. |
| ... | Passed to <code>import_all_interactions</code> . |

Details

omnipath: the OmniPath data as defined in the paper, an arbitrary optimum between coverage and quality pathwayextra: activity flow interactions without literature reference kinaseextra: enzyme-substrate interactions without literature reference ligrecextra: ligand-receptor interactions without literature reference dorothea: transcription factor (TF)-target interactions from DoRothEA tf_target: transcription factor (TF)-target interactions from other resources mirnatarget: miRNA-mRNA interactions tf_mirna: TF-miRNA interactions lncrna_mrna: lncRNA-mRNA interactions

Value

A dataframe containing all the datasets in the interactions query

See Also

- [get_interaction_resources](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```
interactions <- import_all_interactions(
  resources = c('HPRD', 'BioGRID'),
  organism = 9606
)
```

```
import_dorothea_interactions
```

*From the OmniPath webservice imports interactions from the
DoRothEA dataset*

Description

Imports the dataset from: <https://omnipathdb.org/interactions?datasets=dorothea> which contains transcription factor (TF)-target interactions from DoRothEA <https://github.com/saezlab/DoRothEA>

Usage

```
import_dorothea_interactions(  
  resources = NULL,  
  organism = 9606,  
  dorothea_levels = c("A", "B"),  
  fields = NULL,  
  default_fields = TRUE,  
  references_by_resource = TRUE,  
  exclude = NULL,  
  ...  
)
```

Arguments

| | |
|------------------------|--|
| resources | interactions not reported in these databases are removed. See get_interaction_resources for more information. |
| organism | Interactions are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse |
| dorothea_levels | Vector detailing the confidence levels of the interactions to be downloaded. In dorothea, every TF-target interaction has a confidence score ranging from A to E, being A the most reliable interactions. By default we take A and B level interactions (c(A, B)). It is to note that E interactions are not available in OmnipathR. |
| fields | The user can define here the fields to be added. If used, set the next argument, ‘default_fields’, to FALSE. |
| default_fields | whether to include the default fields (columns) for the query type. If FALSE, only the fields defined by the user in the ‘fields’ argument will be added. |
| references_by_resource | if FALSE, removes the resource name prefixes from the references (PubMed IDs); this way the information which reference comes from which resource will be lost and the PubMed IDs will be unique. |
| exclude | Character: datasets or resources to exclude. |
| ... | optional additional arguments |

Value

A dataframe containing TF-target interactions from DoRothEA

See Also

- [get_interaction_resources](#)
- [import_all_interactions](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```
interactions <- import_dorothea_interactions(  
    resources = c('DoRothEA', 'ARACNe-GTEx_DoRothEA'),  
    organism = 9606,  
    dorothea_levels = c('A', 'B', 'C')  
)
```

import_intercell_network

Intercellular communication network

Description

Imports an intercellular network by combining intercellular annotations and protein interactions. First imports a network of protein-protein interactions. Then, it retrieves annotations about the proteins intercellular communication roles, once for the transmitter (delivering information from the expressing cell) and second, the receiver (receiving signal and relaying it towards the expressing cell) side. These 3 queries can be customized by providing parameters in lists which will be passed to the respective methods ([import_omnipath_interactions](#) for the network and [import_omnipath_intercell](#) for the annotations). Finally the 3 data frames combined in a way that the source proteins in each interaction annotated by the transmitter, and the target proteins by the receiver categories. If undirected interactions present (these are disabled by default) they will be duplicated, i.e. both partners can be both receiver and transmitter.

Usage

```
import_intercell_network(  
    interactions_param = list(),  
    transmitter_param = list(),  
    receiver_param = list(),  
    resources = NULL,  
    entity_types = NULL,  
    ligand_receptor = FALSE,  
    high_confidence = FALSE,
```

```

simplify = FALSE,
unique_pairs = FALSE,
consensus_percentile = NULL,
loc_consensus_percentile = NULL,
omnipath = TRUE,
ligrecextra = TRUE,
kinaseextra = !high_confidence,
pathwayextra = !high_confidence,
...
)

```

Arguments

`interactions_param`

a list with arguments for an interactions query: `import_omnipath_interactions`, `import_pathwayextra_interactions`, `import_kinaseextra_interactions`, `import_ligrecextra_interactions`

`transmitter_param`

a list with arguments for `import_omnipath_intercell`, to define the transmitter side of intercellular connections

`receiver_param` a list with arguments for `import_omnipath_intercell`, to define the receiver side of intercellular connections

`resources` A character vector of resources to be applied to both the interactions and the annotations. For example, `resources = 'CellChatDB'` will download the transmitters and receivers defined by CellChatDB, connected by connections from CellChatDB.

`entity_types` Character, possible values are "protein", "complex" or both.

`ligand_receptor`

Logical. If TRUE, only *ligand* and *receptor* annotations will be used instead of the more generic *transmitter* and *receiver* categories.

`high_confidence`

Logical: shortcut to do some filtering in order to include only higher confidence interactions. The intercell database of OmniPath covers a very broad range of possible ways of cell to cell communication, and the pieces of information, such as localization, topology, function and interaction, are combined from many, often independent sources. This unavoidably result some weird and unexpected combinations which are false positives in the context of intercellular communication. This option sets some minimum criteria to remove most (but definitely not all!) of the wrong connections. These criteria are the followings: 1) the receiver must be plasma membrane transmembrane; 2) the curation effort for interactions must be larger than one; 3) the consensus score for annotations must be larger than the 50 percentile within the generic category (you can override this by `consensus_percentile`). 4) the transmitter must be secreted or exposed on the plasma membrane. 5) The major localizations have to be supported by at least 30 percent of the relevant resources (you can override this by `loc_consensus_percentile`). 6) The datasets with lower level of curation (`kinaseextra` and `pathwayextra`) will be disabled. These criteria are of medium

| | |
|--------------------------|---|
| | stringency, you can always tune them to be more relaxed or stringent by filtering manually, using filter_intercell_network . |
| simplify | Logical: keep only the most often used columns. This function combines a network data frame with two copies of the intercell annotation data frames, all of them already having quite some columns. With this option we keep only the names of the interacting pair, their intercellular communication roles, and the minimal information of the origin of both the interaction and the annotations. |
| unique_pairs | Logical: instead of having separate rows for each pair of annotations, drop the annotations and reduce the data frame to unique interacting pairs. See unique_intercell_network for details. |
| consensus_percentile | Numeric: a percentile cut off for the consensus score of generic categories in intercell annotations. The consensus score is the number of resources supporting the classification of an entity into a category based on combined information of many resources. Here you can apply a cut-off, keeping only the annotations supported by a higher number of resources than a certain percentile of each category. If NULL no filtering will be performed. The value is either in the 0-1 range, or will be divided by 100 if greater than 1. The percentiles will be calculated against the generic composite categories and then will be applied to their resource specific annotations and specific child categories. |
| loc_consensus_percentile | Numeric: similar to consensus_percentile for major localizations. For example, with a value of 50, the secreted, plasma membrane transmembrane or peripheral attributes will be TRUE only where at least 50 percent of the resources support these. |
| omnipath | Logical: shortcut to include the <i>omnipath</i> dataset in the interactions query. |
| ligrecrextra | Logical: shortcut to include the <i>ligrecrextra</i> dataset in the interactions query. |
| kinaseextra | Logical: shortcut to include the <i>kinaseextra</i> dataset in the interactions query. |
| pathwayextra | Logical: shortcut to include the <i>pathwayextra</i> dataset in the interactions query. |
| ... | If simplify or unique_pairs is TRUE, additional column names can be passed here to dplyr::select on the final data frame. Otherwise ignored. |

Details

By default this function creates almost the largest possible network of intercellular interactions. However, this might contain a large number of false positives. Please refer to the documentation of the arguments, especially `high_confidence`, and the [filter_intercell_network](#) function. Note: if you restrict the query to certain intercell annotation resources or small categories, it's not recommended to use the `consensus_percentile` or `high_confidence` options, instead filter the network with [filter_intercell_network](#) for more consistent results.

Value

A data frame containing information about protein-protein interactions and the inter-cellular roles of the proteins involved in those interactions.

See Also

- `get_intercell_categories`
- `get_intercell_generic_categories`
- `import_omnipath_intercell`
- `import_omnipath_interactions`
- `import_pathwayextra_interactions`
- `import_kinaseextra_interactions`
- `import_ligrecextra_interactions`
- `unique_intercell_network`
- `simplify_intercell_network`
- `filter_intercell_network`

Examples

```
intercell_network <- import_intercell_network(
  interactions_param = list(datasets = 'ligrecextra'),
  receiver_param = list(categories = c('receptor', 'transporter')),
  transmitter_param = list(categories = c('ligand', 'secreted_enzyme'))
)
```

import_kinaseextra_interactions

Imports interactions from the ‘kinase extra’ dataset of OmniPath

Description

Imports the dataset from: <https://omnipathdb.org/interactions?datasets=kinaseextra>, which contains enzyme-substrate interactions without literature reference. The enzyme-substrate interactions supported by literature references are part of the ‘omnipath’ dataset.

Usage

```
import_kinaseextra_interactions(
  resources = NULL,
  organism = 9606,
  fields = NULL,
  default_fields = TRUE,
  references_by_resource = TRUE,
  exclude = NULL,
  ...
)
```

Arguments

| | |
|------------------------|---|
| resources | interactions not reported in these databases are removed. See get_interaction_resources for more information. |
| organism | Interactions are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse |
| fields | The user can define here the fields to be added. If used, set the next argument, ‘default_fields’, to FALSE. |
| default_fields | whether to include the default fields (columns) for the query type. If FALSE, only the fields defined by the user in the ‘fields’ argument will be added. |
| references_by_resource | if FALSE, removes the resource name prefixes from the references (PubMed IDs); this way the information which reference comes from which resource will be lost and the PubMed IDs will be unique. |
| exclude | Character: datasets or resources to exclude. |
| ... | Optional additional arguments. |

Value

A dataframe containing enzyme-substrate interactions without literature reference

See Also

- [get_interaction_resources](#)
- [import_all_interactions](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```
interactions <-
  import_kinaseextra_interactions(
    resources = c('PhosphoPoint', 'PhosphoSite'),
    organism = 9606
  )
```

`import_ligrecrextra_interactions`

Imports interactions from the ‘ligrec extra’ dataset of OmniPath

Description

Imports the dataset from: <https://omnipathdb.org/interactions?datasets=ligrecrextra>, which contains ligand-receptor interactions without literature reference. The ligand-receptor interactions supported by literature references are part of the ‘omnipath’ dataset.

Usage

```
import_ligrecrextra_interactions(
  resources = NULL,
  organism = 9606,
  fields = NULL,
  default_fields = TRUE,
  references_by_resource = TRUE,
  exclude = NULL,
  ...
)
```

Arguments

| | |
|-------------------------------------|---|
| <code>resources</code> | interactions not reported in these databases are removed. See get_interaction_resources for more information. |
| <code>organism</code> | Interactions are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse |
| <code>fields</code> | The user can define here the fields to be added. If used, set the next argument, ‘ <code>default_fields</code> ’, to FALSE. |
| <code>default_fields</code> | whether to include the default fields (columns) for the query type. If FALSE, only the fields defined by the user in the ‘ <code>fields</code> ’ argument will be added. |
| <code>references_by_resource</code> | if FALSE, removes the resource name prefixes from the references (PubMed IDs); this way the information which reference comes from which resource will be lost and the PubMed IDs will be unique. |
| <code>exclude</code> | Character: datasets or resources to exclude. |
| ... | optional additional arguments |

Value

A dataframe containing ligand-receptor interactions including the ones without literature references

See Also

- [get_interaction_resources](#)
- [import_all_interactions](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```
interactions <- import_ligrecrextra_interactions(
  resources = c('HPRD', 'Guide2Pharma'),
  organism = 9606
)
```

```
import_lncrna_mrna_interactions
```

Imports interactions from the lncRNA-mRNA dataset of OmniPath

Description

Imports the dataset from: https://omnipathdb.org/interactions?datasets=lncrna_mrna, which contains lncRNA-mRNA interactions

Usage

```
import_lncrna_mrna_interactions(  
  resources = NULL,  
  organism = 9606,  
  fields = NULL,  
  default_fields = TRUE,  
  references_by_resource = TRUE,  
  exclude = NULL,  
  ...  
)
```

Arguments

| | |
|------------------------|---|
| resources | interactions not reported in these databases are removed. See get_interaction_resources for more information. |
| organism | Interactions are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse |
| fields | The user can define here the fields to be added. If used, set the next argument, ‘default_fields’, to FALSE. |
| default_fields | whether to include the default fields (columns) for the query type. If FALSE, only the fields defined by the user in the ‘fields’ argument will be added. |
| references_by_resource | if FALSE, removes the resource name prefixes from the references (PubMed IDs); this way the information which reference comes from which resource will be lost and the PubMed IDs will be unique. |
| exclude | Character: datasets or resources to exclude. |
| ... | optional additional arguments |

Value

A dataframe containing lncRNA-mRNA interactions

See Also

- [get_interaction_resources](#)
- [import_all_interactions](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```
interactions <-
  import_lncrna_mrna_interactions(
    resources = c('ncRDeathDB')
  )
```

import_mirnatarget_interactions

Imports interactions from the miRNA-target dataset of OmniPath

Description

Imports the dataset from: <https://omnipathdb.org/interactions?datasets=mirnatarget>, which contains miRNA-mRNA interactions.

Usage

```
import_mirnatarget_interactions(
  resources = NULL,
  organism = 9606,
  fields = NULL,
  default_fields = TRUE,
  references_by_resource = TRUE,
  exclude = NULL,
  ...
)
```

Arguments

| | |
|-----------------------|---|
| resources | interactions not reported in these databases are removed. See get_interaction_resources for more information. |
| organism | Interactions are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse |
| fields | The user can define here the fields to be added. If used, set the next argument, 'default_fields', to FALSE. |
| default_fields | whether to include the default fields (columns) for the query type. If FALSE, only the fields defined by the user in the 'fields' argument will be added. |

`import_omnipath_annotations`

51

```
references_by_resource  
    if FALSE, removes the resource name prefixes from the references (PubMed  
    IDs); this way the information which reference comes from which resource will  
    be lost and the PubMed IDs will be unique.  
exclude      Character: datasets or resources to exclude.  
...           optional additional arguments
```

Value

A dataframe containing miRNA-mRNA interactions

See Also

- [get_interaction_resources](#)
- [import_all_interactions](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```
interactions <-  
  import_mirnatarget_interactions(  
    resources = c('miRTarBase', 'miRecords')  
)
```

```
import_omnipath_annotations  
Imports annotations from OmniPath
```

Description

Imports protein annotations about function, localization, expression, structure and other properties of proteins from OmniPath <https://omnipathdb.org/annotations>. Note: there might be also a few miRNAs annotated; a vast majority of protein complex annotations are inferred from the annotations of the members: if all members carry the same annotation the complex inherits.

Usage

```
import_omnipath_annotations(  
  proteins = NULL,  
  resources = NULL,  
  wide = FALSE,  
  ...  
)
```

Arguments

| | |
|------------------------|---|
| <code>proteins</code> | Vector containing the genes or proteins for whom annotations will be retrieved (UniProt IDs or HGNC Gene Symbols or miRBase IDs). It is also possible to download annotations for protein complexes. To do so, write 'COMPLEX:' right before the genesymbols of the genes integrating the complex. Check the vignette for examples. |
| <code>resources</code> | Load the annotations only from these databases. See get_annotation_resources for possible values. |
| <code>wide</code> | Convert the annotation table to wide format, which corresponds more or less to the original resource. If the data comes from more than one resource a list of wide tables will be returned. See examples at pivot_annotations . |
| <code>...</code> | Additional arguments. |

Details

Downloading the full annotations dataset is disabled by default because the size of this data is around 1GB. We recommend to retrieve the annotations for a set of proteins or only from a few resources, depending on your interest. You can always download the full database from https://archive.omnipathdb.org/omnipath_webservice_annotations__recent.tsv using any standard R or `readr` method.

Value

A data frame containing different gene and complex annotations.

See Also

- [get_annotation_databases](#)
- [pivot_annotations](#)

Examples

```
annotations <- import_omnipath_annotations(
  proteins = c('TP53', 'LMNA'),
  resources = c('HPA_subcellular')
)
```

import_omnipath_complexes
Imports protein complexes from OmniPath

Description

Imports the complexes stored in Omnipath database from <https://omnipathdb.org/complexes>.

Usage

```
import_omnipath_complexes(resources = NULL, ...)
```

Arguments

| | |
|-----------|--|
| resources | complexes not reported in these databases are removed. See get_complexes_databases for more information. |
| ... | optional additional arguments |

Value

A dataframe containing information about complexes

See Also

- [get_complexes_databases](#)

Examples

```
complexes = import_omnipath_complexes(  
    resources = c('CORUM', 'hu.MAP')  
)
```

import_omnipath_enzsub

Imports enzyme-substrate relationships from OmniPath

Description

Imports the enzyme-substrate (more exactly, enzyme-PTM) relationship database from <https://omnipathdb.org/enzsub>

Usage

```
import_omnipath_enzsub(  
    resources = NULL,  
    organism = 9606,  
    fields = NULL,  
    default_fields = TRUE,  
    references_by_resource = TRUE,  
    exclude = NULL,  
    ...  
)
```

Arguments

| | |
|-------------------------------------|---|
| <code>resources</code> | PTMs not reported in these databases are removed. See get_ptms_databases for more information. |
| <code>organism</code> | PTMs are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse |
| <code>fields</code> | You can define here additional fields to be added to the result. If used, set the next argument, <code>default_fields</code> , to FALSE. |
| <code>default_fields</code> | Whether to include the default fields (columns) for the query type. If FALSE, only the fields defined by the user in the <code>fields</code> argument will be added. |
| <code>references_by_resource</code> | if FALSE, removes the resource name prefixes from the references (PubMed IDs); this way the information which reference comes from which resource will be lost and the PubMed IDs will be unique. |
| <code>exclude</code> | Character: datasets or resources to exclude. |
| ... | Optional additional arguments. |

Value

A data frame containing the information about ptms

See Also

- [get_enzsub_resources](#)
- [import_omnipath_interactions](#)
- [enzsub_graph](#)
- [print_interactions](#)

Examples

```
enzsub <- import_omnipath_enzsub(
  resources = c('PhosphoSite', 'SIGNOR'),
  organism = 9606
)
```

`import_omnipath_interactions`

Imports interactions from the ‘omnipath’ dataset of Omnipath

Description

Imports the database from <https://omnipathdb.org/interactions>, which contains only interactions supported by literature references. This part of the interaction database compiled a similar way as it has been presented in the first paper describing OmniPath (Turei et al. 2016).

Usage

```
import_omnipath_interactions(  
  resources = NULL,  
  organism = 9606,  
  datasets = "omnipath",  
  fields = NULL,  
  default_fields = TRUE,  
  references_by_resource = TRUE,  
  exclude = NULL,  
  ...  
)
```

Arguments

| | |
|------------------------|--|
| resources | interactions not reported in these databases are removed. See get_interaction_resources for more information. |
| organism | Interactions are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse |
| datasets | Names of the interaction datasets to download: omnipath (by default). Other possiblites are: pathwayextra, kinaseextra, ligrecrextra, dorothaea, tf_target, mirnatarget, tf_mirna, lncrna_mrna. The user can select multiple datasets as for example: c('omnipath', 'pathwayextra', 'kinaseextra') |
| fields | The user can define here the fields to be added. If used, set the next argument, 'default_fields', to FALSE. |
| default_fields | whether to include the default fields (columns) for the query type. If FALSE, only the fields defined by the user in the 'fields' argument will be added. |
| references_by_resource | if FALSE, removes the resource name prefixes from the references (PubMed IDs); this way the information which reference comes from which resource will be lost and the PubMed IDs will be unique. |
| exclude | Character: datasets or resources to exclude. |
| ... | optional additional arguments |

Value

A dataframe of protein-protein interactions

See Also

- [get_interaction_resources](#)
- [import_all_interactions](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```
interactions = import_omnipath_interactions(
  resources = c('SignaLink3'),
  organism = 9606
)
```

import_omnipath_intercell

Imports OmniPath intercell annotations

Description

Imports the OmniPath intercellular communication role annotation database from <https://omnipathdb.org/intercell>. It provides information on the roles in inter-cellular signaling. E.g. if a protein is a ligand, a receptor, an extracellular matrix (ECM) component, etc.

Usage

```
import_omnipath_intercell(
  categories = NULL,
  resources = NULL,
  parent = NULL,
  scope = NULL,
  aspect = NULL,
  source = NULL,
  transmitter = NULL,
  receiver = NULL,
  secreted = NULL,
  plasma_membrane_peripheral = NULL,
  plasma_membrane_transmembrane = NULL,
  proteins = NULL,
  topology = NULL,
  causality = NULL,
  consensus_percentile = NULL,
  loc_consensus_percentile = NULL,
  ...
)
```

Arguments

| | |
|------------|---|
| categories | vector containing the categories to be retrieved. All the genes belonging to those categories will be returned. For further information about the categories see codeget_intercell_categories . |
| resources | limit the query to certain resources; see the available resources by get_intercell_resources . |

| | |
|-------------------------------|--|
| parent | vector containing the parent classes to be retrieved. All the genes belonging to those classes will be returned. For further information about the main classes see get_intercell_categories . |
| scope | either ‘specific’ or ‘generic’ |
| aspect | either ‘locational’ or ‘functional’ |
| source | either ‘resource_specific’ or ‘composite’ |
| transmitter | logical, include only transmitters i.e. proteins delivering signal from a cell to its environment. |
| receiver | logical, include only receivers i.e. proteins delivering signal to the cell from its environment. |
| secreted | logical, include only secreted proteins |
| plasma_membrane_peripheral | logical, include only plasma membrane peripheral membrane proteins. |
| plasma_membrane_transmembrane | logical, include only plasma membrane transmembrane proteins. |
| proteins | limit the query to certain proteins |
| topology | topology categories: one or more of ‘secreted’ (sec), ‘plasma_membrane_peripheral’ (pmp), ‘plasma_membrane_transmembrane’ (pmtm) (both short or long notation can be used). |
| causality | ‘transmitter’ (trans), ‘receiver’ (rec) or ‘both’ (both short or long notation can be used). |
| consensus_percentile | Numeric: a percentile cut off for the consensus score of generic categories. The consensus score is the number of resources supporting the classification of an entity into a category based on combined information of many resources. Here you can apply a cut-off, keeping only the annotations supported by a higher number of resources than a certain percentile of each category. If NULL no filtering will be performed. The value is either in the 0-1 range, or will be divided by 100 if greater than 1. The percentiles will be calculated against the generic composite categories and then will be applied to their resource specific annotations and specific child categories. |
| loc_consensus_percentile | Numeric: similar to codeconsensus_percentile for major localizations. For example, with a value of 50, the secreted, plasma membrane transmembrane or peripheral attributes will be true only where at least 50 percent of the resources support these. |
| ... | Additional optional arguments, ignored. |

Value

A dataframe containing information about roles in intercellular signaling.

See Also

- [get_intercell_categories](#)
- [get_intercell_generic_categories](#)
- [import_intercell_network](#)
- [intercell_consensus_filter](#)

Examples

```
intercell <- import_omnipath_intercell(categories = 'ecm')
```

import_pathwayextra_interactions

Imports interactions from the ‘pathway extra’ dataset of Omnipath

Description

Imports the dataset from: <https://omnipathdb.org/interactions?datasets=pathwayextra>, which contains activity flow interactions without literature reference. The activity flow interactions supported by literature references are part of the ‘omnipath’ dataset.

Usage

```
import_pathwayextra_interactions(
  resources = NULL,
  organism = 9606,
  fields = NULL,
  default_fields = TRUE,
  references_by_resource = TRUE,
  exclude = NULL,
  ...
)
```

Arguments

| | |
|-----------------------------|--|
| <code>resources</code> | interactions not reported in these databases are removed. See get_interaction_resources for more information. |
| <code>organism</code> | Interactions are available for human, mouse and rat. Choose one of those: 9606 human (default), 10116 rat or 10090 Mouse. |
| <code>fields</code> | The user can define here the fields to be added. If used, set the next argument, ‘ <code>default_fields</code> ’, to FALSE. |
| <code>default_fields</code> | whether to include the default fields (columns) for the query type. If FALSE, only the fields defined by the user in the ‘ <code>fields</code> ’ argument will be added. |

```
references_by_resource  
    if FALSE, removes the resource name prefixes from the references (PubMed  
    IDs); this way the information which reference comes from which resource will  
    be lost and the PubMed IDs will be unique.  
exclude      Character: datasets or resources to exclude.  
...           optional additional arguments
```

Value

A dataframe containing activity flow interactions between proteins without literature reference

See Also

- [get_interaction_resources](#)
- [import_all_interactions](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```
interactions <-  
  import_pathwayextra_interactions(  
    resources = c('BioGRID', 'IntAct'),  
    organism = 9606  
)
```

import_post_translational_interactions

All post-translational interactions from OmniPath

Description

Imports interactions from all post-translational datasets of OmniPath. The datasets are "omnipath", "kinaseextra", "pathwayextra" and "ligrecrextra".

Usage

```
import_post_translational_interactions(  
  resources = NULL,  
  organism = 9606,  
  exclude = NULL,  
  references_by_resource = TRUE,  
  ...  
)
```

Arguments

- `resources` interactions not reported in these databases are removed. See [get_interaction_resources](#) for more information.
- `organism` Interactions are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse
- `exclude` Character: datasets or resources to exclude.
- `references_by_resource` if FALSE, removes the resource name prefixes from the references (PubMed IDs); this way the information which reference comes from which resource will be lost and the PubMed IDs will be unique.
- `...` optional additional arguments

Value

A dataframe containing post-translational interactions

See Also

- [get_interaction_resources](#)
- [import_all_interactions](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```
interactions <-  
  import_post_translational_interactions(  
    resources = c('BioGRID')  
  )
```

import_tf_mirna_interactions

Imports interactions from the TF-miRNA dataset of OmniPath

Description

Imports the dataset from: https://omnipathdb.org/interactions?datasets=tf_mirna, which contains transcription factor-miRNA gene interactions

Usage

```
import_tf_mirna_interactions(  
  resources = NULL,  
  organism = 9606,  
  fields = NULL,  
  default_fields = TRUE,  
  references_by_resource = TRUE,  
  exclude = NULL,  
  ...  
)
```

Arguments

| | |
|------------------------|---|
| resources | interactions not reported in these databases are removed. See get_interaction_resources for more information. |
| organism | Interactions are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse |
| fields | The user can define here the fields to be added. If used, set the next argument, 'default_fields', to FALSE. |
| default_fields | whether to include the default fields (columns) for the query type. If FALSE, only the fields defined by the user in the 'fields' argument will be added. |
| references_by_resource | if FALSE, removes the resource name prefixes from the references (PubMed IDs); this way the information which reference comes from which resource will be lost and the PubMed IDs will be unique. |
| exclude | Character: datasets or resources to exclude. |
| ... | optional additional arguments |

Value

A dataframe containing TF-miRNA interactions

See Also

- [get_interaction_resources](#)
- [import_all_interactions](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```
interactions <-  
  import_tf_mirna_interactions(  
    resources = c('TransmiR')  
)
```

import_tf_target_interactions

Imports interactions from the TF-target dataset of OmniPath

Description

Imports the dataset from: https://omnipathdb.org/interactions?datasets=tf_target, which contains transcription factor-target protein coding gene interactions. Note: this is not the only TF-target dataset in OmniPath, ‘dorothea’ is the other one and the ‘tf_mirna’ dataset provides TF-miRNA gene interactions.

Usage

```
import_tf_target_interactions(  
  resources = NULL,  
  organism = 9606,  
  fields = NULL,  
  default_fields = TRUE,  
  references_by_resource = TRUE,  
  exclude = NULL,  
  ...  
)
```

Arguments

| | |
|-------------------------------------|---|
| <code>resources</code> | interactions not reported in these databases are removed. See get_interaction_resources for more information. |
| <code>organism</code> | Interactions are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse |
| <code>fields</code> | The user can define here the fields to be added. If used, set the next argument, ‘ <code>default_fields</code> ’, to FALSE. |
| <code>default_fields</code> | whether to include the default fields (columns) for the query type. If FALSE, only the fields defined by the user in the ‘ <code>fields</code> ’ argument will be added. |
| <code>references_by_resource</code> | if FALSE, removes the resource name prefixes from the references (PubMed IDs); this way the information which reference comes from which resource will be lost and the PubMed IDs will be unique. |
| <code>exclude</code> | Character: datasets or resources to exclude. |
| ... | Optional additional arguments |

Value

A dataframe containing TF-target interactions

See Also

- [get_interaction_resources](#)
- [import_all_interactions](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```
interactions <-  
  import_tf_target_interactions(  
    resources = c('DoRothEA', 'SIGNOR')  
)
```

import_transcriptional_interactions

Imports all TF-target interactions from OmniPath

Description

Imports the dataset from: https://omnipathdb.org/interactions?datasets=tf_target,dorothea, which contains transcription factor-target protein coding gene interactions.

Usage

```
import_transcriptional_interactions(  
  resources = NULL,  
  organism = 9606,  
  dorothea_levels = c("A", "B"),  
  references_by_resource = TRUE,  
  exclude = NULL,  
  ...  
)
```

Arguments

| | |
|-----------------|---|
| resources | interactions not reported in these databases are removed. See get_interaction_resources for more information. |
| organism | Interactions are available for human, mouse and rat. Choose among: 9606 human (default), 10116 rat and 10090 Mouse |
| dorothea_levels | Vector detailing the confidence levels of the interactions to be downloaded. In dorothea, every TF-target interaction has a confidence score ranging from A to E, being A the most reliable interactions. By default we take A and B level interactions (c(A,B)). It is to note that E interactions are not available in OmnipathR. |

```

references_by_resource
    if FALSE, removes the resource name prefixes from the references (PubMed
    IDs); this way the information which reference comes from which resource will
    be lost and the PubMed IDs will be unique.

exclude          Character: datasets or resources to exclude.

...              Optional additional arguments.

```

Value

A data frame containing TF-target interactions.

See Also

- [get_interaction_resources](#)
- [import_all_interactions](#)
- [interaction_graph](#)
- [print_interactions](#)

Examples

```

interactions <- 
  import_transcriptional_interactions(
    resources = c('PAZAR', 'ORegAnno', 'DoRothEA')
  )

```

| | |
|--------------------------|--|
| inbiomap_download | <i>Downloads and preprocesses network data from InWeb InBioMap</i> |
|--------------------------|--|

Description

Downloads the data by [inbiomap_raw](#), extracts the UniProt IDs, Gene Symbols and scores and removes the irrelevant columns.

Usage

```
inbiomap_download(...)
```

Arguments

| | |
|-----|--|
| ... | Passed to inbiomap_raw . |
|-----|--|

Value

A data frame (tibble) of interactions.

See Also[inbiomap_raw](#)**Examples**

```
inbiomap_interactions <- inbiomap_download()
inbiomap_interactions

# # A tibble: 625,641 x 7
#   uniprot_a uniprot_b genesymbol_a genesymbol_b inferred score1 score2
#   <chr>     <chr>      <chr>      <chr>      <lgl>    <dbl>   <dbl>
# 1 A0A5B9    P01892    TRBC2      HLA-A      FALSE    0.417  0.458
# 2 A0AUZ9    Q96CV9    KANSL1L    OPTN       FALSE    0.155  0.0761
# 3 A0AV02    P24941    SLC12A8   CDK2       TRUE     0.156  0.0783
# 4 A0AV02    Q00526    SLC12A8   CDK3       TRUE     0.157  0.0821
# 5 A0AV96    P0CG48    RBM47      UBC        FALSE    0.144  0.0494
# # . with 625,631 more rows
```

inbiomap_raw*Downloads network data from InWeb InBioMap*

Description

Downloads the data from <https://inbio-discover.com/map.html#downloads> in tar.gz format, extracts the PSI MITAB table and returns it as a data frame.

Usage

```
inbiomap_raw(curl_verbose = FALSE)
```

Arguments

`curl_verbose` Logical. Perform CURL requests in verbose mode for debugging purposes.

Value

A data frame (tibble) with the extracted interaction table.

See Also[inbiomap_download](#)**Examples**

```
inbiomap_psimitab <- inbiomap_raw()
```

`interaction_graph` *Build Omnipath interaction graph*

Description

Transforms the interactions data frame to an igraph graph object.

Usage

```
interaction_graph(interactions = interactions)
```

Arguments

- `interactions` data.frame created by
- [import_omnipath_enzsub](#)
 - [import_omnipath_interactions](#)
 - [import_pathwayextra_interactions](#)
 - [import_kinaseextra_interactions](#)
 - [import_ligrecextra_interactions](#)
 - [import_post_translational_interactions](#)
 - [import_dorothea_interactions](#)
 - [import_tf_target_interactions](#)
 - [import_transcriptional_interactions](#)
 - [import_mirnatarget_interactions](#)
 - [import_all_interactions](#)

Value

An igraph graph object.

See Also

- [import_omnipath_interactions](#)
- [import_pathwayextra_interactions](#)
- [import_kinaseextra_interactions](#)
- [import_ligrecextra_interactions](#)
- [import_dorothea_interactions](#)
- [import_mirnatarget_interactions](#)
- [import_all_interactions](#)
- [giant_component](#)
- [find_all_paths](#)

Examples

```
interactions <- import_omnipath_interactions(resources = c('SignalLink3'))
g <- interaction_graph(interactions)
```

intercell_categories *Full list of intercell categories and resources*

Description

Full list of intercell categories and resources

Usage

```
intercell_categories()
```

Value

A data frame of categories and resources.

Examples

```
ic_cat <- intercell_categories()
ic_cat
# # A tibble: 1,125 x 3
#   category          parent      database
#   <chr>            <chr>       <chr>
# 1 transmembrane    transmembrane UniProt_location
# 2 transmembrane    transmembrane UniProt_topology
# 3 transmembrane    transmembrane UniProt_keyword
# 4 transmembrane    transmembrane_predicted Phobius
# 5 transmembrane_phobius transmembrane_predicted Almen2009
# # . with 1,120 more rows
```

intercell_consensus_filter
Quality filter for intercell annotations

Description

Quality filter for intercell annotations

Usage

```
intercell_consensus_filter(data, percentile = NULL, loc_percentile = NULL)
```

Arguments

| | |
|-----------------------------|--|
| <code>data</code> | A data frame with intercell annotations, as provided by import_omnipath_intercell . |
| <code>percentile</code> | Numeric: a percentile cut off for the consensus score of composite categories. The consensus score is the number of resources supporting the classification of an entity into a category based on combined information of many resources. Here you can apply a cut-off, keeping only the annotations supported by a higher number of resources than a certain percentile of each category. If NULL no filtering will be performed. The value is either in the 0-1 range, or will be divided by 100 if greater than 1. The percentiles will be calculated against the generic composite categories and then will be applied to their resource specific annotations and specific child categories. |
| <code>loc_percentile</code> | Numeric: similar to percentile for major localizations. For example, with a value of 50, the secreted, plasma membrane transmembrane or peripheral attributes will be TRUE only where at least 50 percent of the resources support these. |

Value

The data frame in `data` filtered by the consensus scores.

Examples

```
intercell <- import_omnipath_intercell(parent = c('ligand', 'receptor'))
nrow(intercell)
# [1] 50174
intercell_q50 <- intercell_consensus_filter(intercell, 50)
nrow(intercell_q50)
# [1] 42863
```

| | |
|-----------------------------|----------------------------------|
| <code>is_ontology_id</code> | <i>Looks like an ontology ID</i> |
|-----------------------------|----------------------------------|

Description

Tells if the input has the typical format of ontology IDs, i.e. a code of capital letters, a colon, followed by a numeric code.

Usage

```
is_ontology_id(terms)
```

Arguments

| | |
|--------------------|---|
| <code>terms</code> | Character vector with strings to check. |
|--------------------|---|

Value

A logical vector with the same length as the input.

Examples

```
is_ontology_id(c('GO:0000001', 'reproduction'))  
# [1] TRUE FALSE
```

kegg_info

*Information about a KEGG Pathway***Description**

Information about a KEGG Pathway

Usage

```
kegg_info(pathway_id)
```

Arguments

pathway_id Character: a KEGG Pathway identifier, e.g. "hsa04710". For a complete list of IDs see [kegg_pathway_list](#).

Value

List with the pathway information.

See Also

- [kegg_pathway_list](#)
- [kegg_picture](#)
- [kegg_open](#)

Examples

```
kegg_info('map00563')
```

kegg_open*Open a KEGG Pathway diagram in the browser*

Description

Open a KEGG Pathway diagram in the browser

Usage

```
kegg_open(pathway_id)
```

Arguments

pathway_id Character: a KEGG Pathway identifier, e.g. "hsa04710". For a complete list of IDs see [kegg_pathway_list](#).

Details

To open URLs in the web browser the "browser" option must be set to a valid executable. You can check the value of this option by `getOption("browser")`. If your browser is firefox and the executable is located in the system path, you can set the option to point to it: `options(browser = "firefox")`. To make it a permanent setting, you can also include this in your `.Rprofile` file.

Value

Returns NULL.

See Also

- [kegg_pathway_list](#)
- [kegg_picture](#)
- [kegg_info](#)

Examples

```
if(any(getOption('browser') != '')) kegg_open('hsa04710')
```

kegg_pathways_download
Download the KEGG Pathways database

Description

Downloads all pathway diagrams in the KEGG Pathways database in KGML format and processes the XML to extract the interactions.

Usage

```
kegg_pathways_download(max_expansion = NULL, simplify = FALSE)
```

Arguments

- max_expansion Numeric: the maximum number of relations derived from a single relation record. As one entry might represent more than one molecular entities, one relation might yield a large number of relations in the processing. This happens in a combinatorial way, e.g. if the two entries represent 3 and 4 entities, that results 12 relations. If NULL, all relations will be expanded.
- simplify Logical: remove KEGG's internal identifiers and the pathway annotations, keep only unique interactions with direction and effect sign.

Value

A data frame (tibble) of interactions.

See Also

- [kegg_pathway_list](#)
- [kegg_process](#)
- [kegg_pathway_download](#)

Examples

```
kegg_pw <- kegg_pathways_download(simplify = TRUE)
kegg_pw
# # A tibble: 6,765 x 6
#   uniprot_source uniprot_target type  effect genesymbol_source
#   <chr>          <chr>        <chr> <chr>  <chr>
# 1 Q03113        Q15283      PPrel activ. GNA12
# 2 Q9Y4G8        P62070      PPrel activ. RAPGEF2
# 3 Q13972        P62070      PPrel activ. RASGRF1
# 4 O95267        P62070      PPrel activ. RASGRP1
# 5 P62834        P15056      PPrel activ. RAP1A
# # . with 6,760 more rows, and 1 more variable: genesymbol_target <chr>
```

kegg_pathway_annotations
Protein pathway annotations

Description

Downloads all KEGG pathways and creates a table of protein-pathway annotations.

Usage

```
kegg_pathway_annotations(pathways = NULL)
```

Arguments

pathways A table of KEGG pathways as produced by [kegg_pathways_download](#).

Value

A data frame (tibble) with UniProt IDs and pathway names.

See Also

[kegg_pathways_download](#)

Examples

```
kegg_pw_annot <- kegg_pathway_annotations()  
kegg_pw_annot  
# # A tibble: 7,341 x 4  
#   uniprot genesymbol pathway           pathway_id  
#   <chr>    <chr>    <chr>           <chr>  
# 1 Q03113  GNA12    MAPK signaling pathway hsa04010  
# 2 Q9Y4G8  RAPGEF2   MAPK signaling pathway hsa04010  
# 3 Q13972  RASGRF1   MAPK signaling pathway hsa04010  
# 4 O95267  RASGRP1   MAPK signaling pathway hsa04010  
# 5 P62834  RAP1A     MAPK signaling pathway hsa04010  
# # . with 7,336 more rows
```

kegg_pathway_download *Download one KEGG pathway*

Description

Downloads one pathway diagram from the KEGG Pathways database in KGML format and processes the XML to extract the interactions.

Usage

```
kegg_pathway_download(  
  pathway_id,  
  process = TRUE,  
  max_expansion = NULL,  
  simplify = FALSE  
)
```

Arguments

| | |
|---------------|---|
| pathway_id | Character: a KEGG pathway identifier, for example "hsa04350". |
| process | Logical: process the data or return it in raw format. processing means joining the entries and relations into a single data frame and adding UniProt IDs. |
| max_expansion | Numeric: the maximum number of relations derived from a single relation record. As one entry might represent more than one molecular entities, one relation might yield a large number of relations in the processing. This happens in a combinatorial way, e.g. if the two entries represent 3 and 4 entities, that results 12 relations. If NULL, all relations will be expanded. |
| simplify | Logical: remove KEGG's internal identifiers and the pathway annotations, keep only unique interactions with direction and effect sign. |

Value

A data frame (tibble) of interactions if process is TRUE, otherwise a list with two data frames: "entries" is a raw table of the entries while "relations" is a table of relations extracted from the KGML file.

See Also

- [kegg_process](#)
- [kegg_pathways_download](#)
- [kegg_pathway_list](#)

Examples

```
tgf_pathway <- kegg_pathway_download('hsa04350')
tgf_pathway
# # A tibble: 50 x 12
#   source target type effect relation_id kegg_id_source
#   <chr>  <chr>  <chr> <chr> <chr>      <chr>
# 1 51     49     PPrel activ. --> hsa04350:1 hsa:7040 hsa:.
# 2 57     55     PPrel activ. --> hsa04350:2 hsa:151449 hs.
# 3 34     32     PPrel activ. --> hsa04350:3 hsa:3624 hsa:.
# 4 20     17     PPrel activ. --> hsa04350:4 hsa:4838
# 5 60     46     PPrel activ. --> hsa04350:5 hsa:4086 hsa:.
# # . with 45 more rows, and 5 more variables: genesymbol_source <chr>,
# #   uniprot_source <chr>, kegg_id_target <chr>,
# #   genesymbol_target <chr>, uniprot_target <chr>
```

kegg_pathway_list *List of KEGG pathways*

Description

Retrieves a list of available KEGG pathways.

Usage

```
kegg_pathway_list()
```

Value

Data frame of pathway names and identifiers.

See Also

- [kegg_process](#)
- [kegg_pathway_download](#)
- [kegg_pathways_download](#)
- [kegg_open](#)
- [kegg_picture](#)
- [kegg_info](#)

Examples

```
kegg_pws <- kegg_pathway_list()
kegg_pws
# # A tibble: 521 x 2
#   id      name
#   <chr>   <chr>
```

```
# 1 map01100 Metabolic pathways
# 2 map01110 Biosynthesis of secondary metabolites
# 3 map01120 Microbial metabolism in diverse environments
# 4 map01200 Carbon metabolism
# 5 map01210 2-Oxocarboxylic acid metabolism
# 6 map01212 Fatty acid metabolism
# 7 map01230 Biosynthesis of amino acids
# # . with 514 more rows
```

kegg_picture*Download a pathway diagram as a picture*

Description

Downloads a KEGG Pathway diagram as a PNG image.

Usage

```
kegg_picture(pathway_id, path = NULL)
```

Arguments

| | |
|------------|--|
| pathway_id | Character: a KEGG Pathway identifier, e.g. "hsa04710". For a complete list of IDs see kegg_pathway_list . |
| path | Character: save the image to this path. If NULL, the image will be saved in the current directory under the name <pathway_id>.png. |

Value

Invisibly returns the path to the downloaded file.

See Also

[kegg_pathway_list](#)

- [kegg_pathway_list](#)
- [kegg_open](#)
- [kegg_info](#)

Examples

```
kegg_picture('hsa04710')
kegg_picture('hsa04710', path = 'foo/bar')
kegg_picture('hsa04710', path = 'foo/bar/circadian.png')
```

Description

Processes KEGG Pathways data extracted from a KGML file. Joins the entries and relations into a single data frame and translates the Gene Symbols to UniProt IDs.

Usage

```
kegg_process(entries, relations, max_expansion = NULL, simplify = FALSE)
```

Arguments

| | |
|---------------|---|
| entries | A data frames with entries extracted from a KGML file by kegg_pathway_download . |
| relations | A data frames with relations extracted from a KGML file by kegg_pathway_download . |
| max_expansion | Numeric: the maximum number of relations derived from a single relation record. As one entry might represent more than one molecular entities, one relation might yield a large number of relations in the processing. This happens in a combinatorial way, e.g. if the two entries represent 3 and 4 entities, that results 12 relations. If NULL, all relations will be expanded. |
| simplify | Logical: remove KEGG's internal identifiers and the pathway annotations, keep only unique interactions with direction and effect sign. |

Value

A data frame (tibble) of interactions. In rare cases when a pathway doesn't contain any relation, returns NULL.

See Also

- [kegg_pathway_download](#)
- [kegg_pathways_download](#)
- [kegg_pathway_list](#)

Examples

```
hsa04350 <- kegg_pathway_download('hsa04350', process = FALSE)
tgf_pathway <- kegg_process(hsa04350$entries, hsa04350$relations)
tgf_pathway
# # A tibble: 50 x 12
#   source target type  effect arrow relation_id kegg_id_source
#   <chr>  <chr>  <chr> <chr>  <chr> <chr>      <chr>
# 1 51     49     PPrel activ. -->  hsa04350:1  hsa:7040 hsa:.
# 2 57     55     PPrel activ. -->  hsa04350:2  hsa:151449 hs.
# 3 34     32     PPrel activ. -->  hsa04350:3  hsa:3624 hsa:.
# 4 20     17     PPrel activ. -->  hsa04350:4  hsa:4838
```

```
# 5 60      46      PPrel activ. --> hsa04350:5 hsa:4086 hsa:.
# # . with 45 more rows, and 5 more variables: genesymbol_source <chr>,
# #   uniprot_source <chr>, kegg_id_target <chr>,
# #   genesymbol_target <chr>, uniprot_target <chr>
```

| | |
|---------|---------------------------------|
| load_db | <i>Load a built in database</i> |
|---------|---------------------------------|

Description

Load a built in database

Usage

```
load_db(key, param = list())
```

Arguments

- | | |
|-------|---|
| key | Character: the key of the database to load. For a list of available keys see omnipath_show_db . |
| param | List: override the defaults or pass further parameters to the database loader function. See the loader functions and their default parameters in omnipath_show_db . |

Details

This function loads a database which is stored within the package namespace until its expiry. The loaded database is accessible by [get_db](#) and the loading process is typically initiated by [get_db](#), not by the users directly.

Value

Returns NULL.

See Also

[omnipath_show_db](#).

Examples

```
load_db('go_slim')
omnipath_show_db()
```

nichenet_build_model *Construct a NicheNet ligand-target model*

Description

Construct a NicheNet ligand-target model

Usage

```
nichenet_build_model(optimization_results, networks, use_weights = TRUE)
```

Arguments

| | |
|----------------------|--|
| optimization_results | The outcome of NicheNet parameter optimization as produced by nichenet_optimization . |
| networks | A list with NicheNet format signaling, ligand-receptor and gene regulatory networks as produced by nichenet_networks . |
| use_weights | Logical: whether to use the optimized weights. |

Value

A named list with two elements: ‘weighted_networks’ and ‘optimized_parameters’.

Examples

```
expression <- nichenet_expression_data()

networks <- nichenet_networks()
optimization_results <- nichenet_optimization(networks, expression)
nichenet_model <- nichenet_build_model(optimization_results, networks)
```

nichenet_expression_data

Expression data from ligand-receptor perturbation experiments used by NicheNet

Description

NicheNet uses expression data from a collection of published ligand or receptor KO or perturbation experiments to build its model. This function retrieves the original expression data, deposited in Zenodo (<https://zenodo.org/record/3260758>).

Usage

```
nichenet_expression_data()
```

Value

Nested list, each element contains a data frame of processed expression data and key variables about the experiment.

Examples

```
exp_data <- nichenet_expression_data()
head(names(exp_data))
# [1] "bmp4_tgfb"      "tgfb_bmp4"      "nodal_Nodal"    "spectrum_Il4"
# [5] "spectrum_Tnf"   "spectrum_Ifng"
purrr::map_chr(head(exp_data), 'from')
#     bmp4_tgfb      tgfb_bmp4  nodal_Nodal  spectrum_Il4  spectrum_Tnf
#     "BMP4"        "TGFB1"      "NODAL"       "IL4"        "TNF"
# spectrum_Ifng
#     "IFNG"
```

nichenet_gr_network *Builds a NicheNet gene regulatory network*

Description

Builds gene regulatory network prior knowledge for NicheNet using multiple resources.

Usage

```
nichenet_gr_network(
  omnipath = list(),
  harmonizome = list(),
  regnetwork = list(),
  htridb = list(),
  remap = list(),
  evex = list(),
  pathwaycommons = list(),
  trrust = list(),
  only_omnipath = FALSE
)
```

Arguments

| | |
|-------------|--|
| omnipath | List with paramaters to be passed to nichenet_gr_network_omnipath . |
| harmonizome | List with paramaters to be passed to nichenet_gr_network_harmonizome . |
| regnetwork | List with paramaters to be passed to nichenet_gr_network_regnetwork . |
| htridb | List with paramaters to be passed to nichenet_gr_network_htridb . |
| remap | List with paramaters to be passed to nichenet_gr_network_remap . |
| evex | List with paramaters to be passed to nichenet_gr_network_evex . |

pathwaycommons List with paramaters to be passed to `nichenet_gr_network_pathwaycommons`.
trrust List with paramaters to be passed to `nichenet_gr_network_trrust`.
only_omnipath Logical: a shortcut to use only OmniPath as network resource.

Value

A network data frame (tibble) with gene regulatory interactions suitable for use with NicheNet.

See Also

- `nichenet_gr_network_evex`
- `nichenet_gr_network_harmonizome`
- `nichenet_gr_network_htridb`
- `nichenet_gr_network_omnipath`
- `nichenet_gr_network_pathwaycommons`
- `nichenet_gr_network_regnetwork`
- `nichenet_gr_network_remap`
- `nichenet_gr_network_trrust`

Examples

```
# load everything with the default parameters:
gr_network <- nichenet_gr_network()

# less targets from ReMap, not using RegNetwork:
gr_network <- nichenet_gr_network(
  # I needed to disable ReMap here due to some issues
  # of one of the Bioconductor build servers
  # remap = list(top_targets = 200),
  remap = NULL,
  regnetwork = NULL,
)

# use only OmniPath:
gr_network_omnipath <- nichenet_gr_network(only_omnipath = TRUE)
```

`nichenet_gr_network_evex`

NicheNet gene regulatory network from EVEX

Description

Builds a gene regulatory network using data from the EVEX database and converts it to a format suitable for NicheNet.

Usage

```
nichenet_gr_network_evex(
  top_confidence = 0.75,
  indirect = FALSE,
  regulation_of_expression = FALSE
)
```

Arguments

`top_confidence` Double, between 0 and 1. Threshold based on the quantile of the confidence score.

`indirect` Logical: whether to include indirect interactions.

`regulation_of_expression` Logical: whether to include also the "regulation of expression" type interactions.

Value

Data frame of interactions in NicheNet format.

Data frame with gene regulatory interactions in NicheNet format.

See Also

- [nichenet_gr_network](#)
- [evex_download](#)

Examples

```
# use only the 10% with the highest confidence:
evex_gr_network <- nichenet_gr_network_evex(top_confidence = .9)
```

`nichenet_gr_network_harmonizome`

NicheNet gene regulatory network from Harmonizome

Description

Builds gene regulatory network prior knowledge for NicheNet using Harmonizome

Usage

```
nichenet_gr_network_harmonizome(
  datasets = c("cheappi", "encodetfppi", "jasparpwm", "transfac", "transfacpwm",
  "motifmap", "geotf", "geokinase", "geogene"),
  ...
)
```

Arguments

- `datasets` The datasets to use. For possible values please refer to default value and the Harmonizome webpage.
`...` Ignored.

Value

Data frame with gene regulatory interactions in NicheNet format.

See Also

- [nichenet_gr_network](#)
- [harmonizome_download](#)

Examples

```
# use only JASPAR and TRANSFAC:  
hz_gr_network <- nichenet_gr_network_harmonizome(  
  datasets = c('jasparpwm', 'transfac', 'transfacpwm')  
)
```

nichenet_gr_network_htridb

NicheNet gene regulatory network from HTTRIdb

Description

Builds a gene regulatory network using data from the HTTRIdb database and converts it to a format suitable for NicheNet.

Usage

`nichenet_gr_network_htridb()`

Value

Data frame with gene regulatory interactions in NicheNet format.

See Also

- [htridb_download](#), [nichenet_gr_network](#)

Examples

```
htri_gr_network <- nichenet_gr_network_htridb()
```

nichenet_gr_network_omnipath

Builds gene regulatory network for NicheNet using OmniPath

Description

Retrieves network prior knowledge from OmniPath and provides it in a format suitable for NicheNet. This method never downloads the ‘ligrecrextra’ dataset because the ligand-receptor interactions are supposed to come from [nichenet_lr_network_omnipath](#).

Usage

```
nichenet_gr_network_omnipath(min_curation_effort = 0, ...)
```

Arguments

| | |
|---------------------|---|
| min_curation_effort | Lower threshold for curation effort |
| ... | Passed to import_transcriptional_interactions |

Value

A network data frame (tibble) with gene regulatory interactions suitable for use with NicheNet.

See Also

- [nichenet_gr_network_evex](#)
- [nichenet_gr_network_harmonizome](#)
- [nichenet_gr_network_htridb](#)
- [nichenet_gr_network_omnipath](#)
- [nichenet_gr_network_pathwaycommons](#)
- [nichenet_gr_network_RegNetwork](#)
- [nichenet_gr_network_remap](#)
- [nichenet_gr_network_trrust](#)

Examples

```
# use interactions up to confidence level "C" from DoRothEA:  
op_gr_network <- nichenet_gr_network_omnipath(  
    dorothaea_levels = c('A', 'B', 'C')  
)
```

nichenet_gr_network_pathwaycommons

NicheNet gene regulatory network from PathwayCommons

Description

Builds gene regulation prior knowledge for NicheNet using PathwayCommons.

Usage

```
nichenet_gr_network_pathwaycommons(  
  interaction_types = "controls-expression-of",  
  ...  
)
```

Arguments

| | |
|-------------------|---|
| interaction_types | Character vector with PathwayCommons interaction types. Please refer to the default value and the PathwayCommons webpage. |
| ... | Ignored. |

Value

Data frame with gene regulatory interactions in NicheNet format.

See Also

- [nichenet_gr_network](#)
- [pathwaycommons_download](#)

Examples

```
pc_gr_network <- nichenet_gr_network_pathwaycommons()
```

nichenet_gr_network_regnetwork

NicheNet gene regulatory network from RegNetwork

Description

Builds a gene regulatory network using data from the RegNetwork database and converts it to a format suitable for NicheNet.

Usage

```
nichenet_gr_network_remap()
```

Value

Data frame with gene regulatory interactions in NicheNet format.

See Also

- [regnetwork_download](#)
- [nichenet_gr_network](#)

Examples

```
regn_gr_network <- nichenet_gr_network_remap()
```

nichenet_gr_network_remap

NicheNet gene regulatory network from ReMap

Description

Builds a gene regulatory network using data from the ReMap database and converts it to a format suitable for NicheNet.

Usage

```
nichenet_gr_network_remap(  
  score = 100,  
  top_targets = 500,  
  only_known_tfs = TRUE  
)
```

Arguments

- score** Numeric: a minimum score between 0 and 1000, records with lower scores will be excluded. If NULL no filtering performed.
- top_targets** Numeric: the number of top scoring targets for each TF. Essentially the maximum number of targets per TF. If NULL the number of targets is not restricted.
- only_known_tfs** Logical: whether to exclude TFs which are not in TF census.

Value

Data frame with gene regulatory interactions in NicheNet format.

See Also

- [remap_filtered](#)
- [nichenet_gr_network](#)

Examples

```
# use only max. top 100 targets for each TF:  
remap_gr_network <- nichenet_gr_network_remap(top_targets = 100)
```

nichenet_gr_network_trrust

NicheNet gene regulatory network from TRRUST

Description

Builds a gene regulatory network using data from the TRRUST database and converts it to a format suitable for NicheNet.

Usage

```
nichenet_gr_network_trrust()
```

Value

Data frame with gene regulatory interactions in NicheNet format.

See Also

- [trrust_download](#)
- [nichenet_gr_network](#)

Examples

```
trrust_gr_network <- nichenet_gr_network_trrust()
```

nichenet_ligand_activities

Calls the NicheNet ligand activity analysis

Description

Calls the NicheNet ligand activity analysis

Usage

```
nichenet_ligand_activities(  
  ligand_target_matrix,  
  lr_network,  
  expressed_genes_transmitter,  
  expressed_genes_receiver,  
  genes_of_interest,  
  background_genes = NULL,  
  n_top_ligands = 42,  
  n_top_targets = 250  
)
```

Arguments

| | |
|-----------------------------|--|
| ligand_target_matrix | A matrix with rows and columns corresponding to ligands and targets, respectively. Produced by nichenet_ligand_target_matrix or nichenetr::construct_ligand_target_matrix . |
| lr_network | A data frame with ligand-receptor interactions, as produced by nichenet_lr_network . |
| expressed_genes_transmitter | Character vector with the gene symbols of the genes expressed in the cells transmitting the signal. |
| expressed_genes_receiver | Character vector with the gene symbols of the genes expressed in the cells receiving the signal. |
| genes_of_interest | Character vector with the gene symbols of the genes of interest. These are the genes in the receiver cell population that are potentially affected by ligands expressed by interacting cells (e.g. genes differentially expressed upon cell-cell interaction). |
| background_genes | Character vector with the gene symbols of the genes to be used as background. |
| n_top_ligands | How many of the top ligands to include in the ligand-target table. |
| n_top_targets | For each ligand, how many of the top targets to include in the ligand-target table. |

Value

A named list with ‘ligand_activities‘ (a tibble giving several ligand activity scores; following columns in the tibble: \$test_ligand, \$auroc, \$aupr and \$pearson) and ‘ligand_target_links‘ (a tibble with columns ligand, target and weight (i.e. regulatory potential score)).

Examples

```
networks <- nichenet_networks()
expression <- nichenet_expression_data()
optimization_results <- nichenet_optimization(networks, expression)
nichenet_model <- nichenet_build_model(optimization_results, networks)
lt_matrix <- nichenet_ligand_target_matrix(
  nichenet_model$weighted_networks,
  networks$lr_network,
  nichenet_model$optimized_parameters
)
ligand_activities <- nichenet_ligand_activities(
  ligand_target_matrix = lt_matrix,
  lr_network = networks$lr_network,
  # the rest of the parameters should come
  # from your transcriptomics data:
  expressed_genes_transmitter = expressed_genes_transmitter,
  expressed_genes_receiver = expressed_genes_receiver,
  genes_of_interest = genes_of_interest
)
```

nichenet_ligand_target_links

Compiles a table with weighted ligand-target links

Description

A wrapper around `nichenetr::get_weighted_ligand_target_links` to compile a data frame with weighted links from the top ligands to their top targets.

Usage

```
nichenet_ligand_target_links(
  ligand_activities,
  ligand_target_matrix,
  genes_of_interest,
  n_top_ligands = 42,
  n_top_targets = 250
)
```

Arguments

| | |
|-----------------------------------|--|
| <code>ligand_activities</code> | Ligand activity table as produced by <code>nichenetr::predict_ligand_activities</code> . |
| <code>ligand_target_matrix</code> | Ligand-target matrix as produced by <code>nichenetr::construct_ligand_target_matrix</code> or the wrapper around it in the current package: <code>nichenet_ligand_target_matrix</code> . |
| <code>genes_of_interest</code> | Character vector with the gene symbols of the genes of interest. These are the genes in the receiver cell population that are potentially affected by ligands expressed by interacting cells (e.g. genes differentially expressed upon cell-cell interaction). |
| <code>n_top_ligands</code> | How many of the top ligands to include in the ligand-target table. |
| <code>n_top_targets</code> | For each ligand, how many of the top targets to include in the ligand-target table. |

Value

A tibble with columns ligand, target and weight (i.e. regulatory potential score).

Examples

```

networks <- nichenet_networks()
expression <- nichenet_expression_data()
optimization_results <- nichenet_optimization(networks, expression)
nichenet_model <- nichenet_build_model(optimization_results, networks)
lt_matrix <- nichenet_ligand_target_matrix(
  nichenet_model$weighted_networks,
  networks$lr_network,
  nichenet_model$optimized_parameters
)
ligand_activities <- nichenet_ligand_activities(
  ligand_target_matrix = lt_matrix,
  lr_network = networks$lr_network,
  # the rest of the parameters should come
  # from your transcriptomics data:
  expressed_genes_transmitter = expressed_genes_transmitter,
  expressed_genes_receiver = expressed_genes_receiver,
  genes_of_interest = genes_of_interest
)
lt_links <- nichenet_ligand_target_links(
  ligand_activities = ligand_activities,
  ligand_target_matrix = lt_matrix,
  genes_of_interest = genes_of_interest,
  n_top_ligands = 20,
  n_top_targets = 100
)

```

nichenet_ligand_target_matrix*Creates a NicheNet ligand-target matrix***Description**

Creates a NicheNet ligand-target matrix

Usage

```
nichenet_ligand_target_matrix(
  weighted_networks,
  lr_network,
  optimized_parameters,
  use_weights = TRUE,
  construct_ligand_target_matrix_param = list()
)
```

Arguments

`weighted_networks`

Weighted networks as provided by [nichenet_build_model](#).

`lr_network` A data frame with ligand-receptor interactions, as produced by [nichenet_lr_network](#).

`optimized_parameters`

The outcome of NicheNet parameter optimization as produced by [nichenet_build_model](#).

`use_weights` Logical: whether the network sources are weighted. In this function it only affects the output file name.

`construct_ligand_target_matrix_param`

Override parameters for `nichenetr::construct_ligand_target_matrix`.

Value

A matrix containing ligand-target probability scores.

Examples

```
networks <- nichenet_networks()
expression <- nichenet_expression_data()
optimization_results <- nichenet_optimization(networks, expression)
nichenet_model <- nichenet_build_model(optimization_results, networks)
lt_matrix <- nichenet_ligand_target_matrix(
  nichenet_model$weighted_networks,
  networks$lr_network,
  nichenet_model$optimized_parameters
)
```

nichenet_lr_network *Builds a NicheNet ligand-receptor network*

Description

Builds ligand-receptor network prior knowledge for NicheNet using multiple resources.

Usage

```
nichenet_lr_network(  
  omnipath = list(),  
  guide2pharma = list(),  
  ramilowski = list(),  
  only_omnipath = FALSE,  
  quality_filter_param = list()  
)
```

Arguments

- | | |
|----------------------|--|
| omnipath | List with paramaters to be passed to nichenet_lr_network_omnipath . |
| guide2pharma | List with paramaters to be passed to nichenet_lr_network_guide2pharma . |
| ramilowski | List with paramaters to be passed to nichenet_lr_network_ramilowski . |
| only_omnipath | Logical: a shortcut to use only OmniPath as network resource. |
| quality_filter_param | Arguments for filter_intercell_network (quality filtering of the OmniPath ligand-receptor network). It is recommended to check these parameters and apply some quality filtering. The defaults already ensure certain filtering, but you might want more relaxed or stringent options. |

Value

A network data frame (tibble) with ligand-receptor interactions suitable for use with NicheNet.

See Also

- [nichenet_lr_network_omnipath](#)
- [nichenet_lr_network_guide2pharma](#)
- [nichenet_lr_network_ramilowski](#)
- [filter_intercell_network](#)

Examples

```
# load everything with the default parameters:  
lr_network <- nichenet_lr_network()  
  
# don't use Ramilowski:  
lr_network <- nichenet_lr_network(ramilowski = NULL)  
  
# use only OmniPath:  
lr_network_omnipath <- nichenet_lr_network(only_omnipath = TRUE)
```

nichenet_lr_network_guide2pharma

Ligand-receptor network from Guide to Pharmacology

Description

Downloads ligand-receptor interactions from the Guide to Pharmacology database and converts it to a format suitable for NicheNet.

Usage

```
nichenet_lr_network_guide2pharma()
```

Value

Data frame with ligand-receptor interactions in NicheNet format.

See Also

[nichenet_lr_network](#), [guide2pharma_download](#)

Examples

```
g2p_lr_network <- nichenet_lr_network_guide2pharma()
```

nichenet_lr_network_omnipath

Builds ligand-receptor network for NicheNet using OmniPath

Description

Retrieves network prior knowledge from OmniPath and provides it in a format suitable for NicheNet. This method never downloads the ‘ligrecrextra’ dataset because the ligand-receptor interactions are supposed to come from [nichenet_lr_network_omnipath](#).

Usage

```
nichenet_lr_network_omnipath(quality_filter_param = list(), ...)
```

Arguments

`quality_filter_param`

List with arguments for [filter_intercell_network](#). It is recommended to check these parameters and apply some quality filtering. The defaults already ensure certain filtering, but you might want more relaxed or stringent options.

`...` Passed to [import_intercell_network](#)

Value

A network data frame (tibble) with ligand-receptor interactions suitable for use with NicheNet.

See Also

- [nichenet_lr_network](#)
- [import_intercell_network](#)

Examples

```
# use only ligand-receptor interactions (not for example ECM-adhesion):
op_lr_network <- nichenet_lr_network_omnipath(ligand_receptor = TRUE)

# use only CellPhoneDB and Guide to Pharmacology:
op_lr_network <- nichenet_lr_network_omnipath(
  resources = c('CellPhoneDB', 'Guide2Pharma')
)

# only interactions where the receiver is a transporter:
op_lr_network <- nichenet_lr_network_omnipath(
  receiver_param = list(parent = 'transporter')
)
```

nichenet_lr_network_ramilowski

Ligand-receptor network from Ramiłowski 2015

Description

Downloads ligand-receptor interactions from Supplementary Table 2 of the paper 'A draft network of ligand–receptor-mediated multicellular signalling in human' (Ramilowski et al. 2015, <https://www.nature.com/articles/ncomms8866>). It converts the downloaded table to a format suitable for NicheNet.

Usage

```
nichenet_lr_network_ramilowski(  
  evidences = c("literature supported", "putative")  
)
```

Arguments

evidences Character: evidence types, "literature supported", "putative" or both.

Value

Data frame with ligand-receptor interactions in NicheNet format.

See Also

- [nichenet_lr_network](#)
- [ramilowski_download](#)

Examples

```
# use only the literature supported data:  
rami_lr_network <- nichenet_lr_network_ramilowski(  
  evidences = 'literature supported'  
)
```

| | |
|---------------|--|
| nichenet_main | <i>Executes the full NicheNet pipeline</i> |
|---------------|--|

Description

Builds all prior knowledge data required by NicheNet. For this it calls a multitude of methods to download and combine data from various databases according to the settings. The content of the prior knowledge data is highly customizable, see the documentation of the related functions. After the prior knowledge is ready, it performs parameter optimization to build a NicheNet model. This results a weighted ligand- target matrix. Then, considering the expressed genes from user provided data, a gene set of interest and background genes, it executes the NicheNet ligand activity analysis.

Usage

```
nichenet_main(  
  only_omnipath = FALSE,  
  expressed_genes_transmitter = NULL,  
  expressed_genes_receiver = NULL,  
  genes_of_interest = NULL,  
  background_genes = NULL,  
  use_weights = TRUE,  
  n_top_ligands = 42,  
  n_top_targets = 250,  
  signaling_network = list(),  
  lr_network = list(),  
  gr_network = list(),  
  small = FALSE,  
  tiny = FALSE,  
  make_multi_objective_function_param = list(),  
  objective_function_param = list(),  
  mlrmbo_optimization_param = list(),  
  construct_ligand_target_matrix_param = list(),  
  results_dir = NULL,  
  quality_filter_param = list()  
)
```

Arguments

- `only_omnipath` Logical: use only OmniPath for network knowledge. This is a simple switch for convenience, further options are available by the other arguments. By default we use all available resources. The networks can be customized on a resource by resource basis, as well as providing custom parameters for individual resources, using the parameters ‘signaling_network’, ‘lr_network’ and ‘gr_network’.
- `expressed_genes_transmitter` Character vector with the gene symbols of the genes expressed in the cells transmitting the signal.

| | |
|---|---|
| <code>expressed_genes_receiver</code> | Character vector with the gene symbols of the genes expressed in the cells receiving the signal. |
| <code>genes_of_interest</code> | Character vector with the gene symbols of the genes of interest. These are the genes in the receiver cell population that are potentially affected by ligands expressed by interacting cells (e.g. genes differentially expressed upon cell-cell interaction). |
| <code>background_genes</code> | Character vector with the gene symbols of the genes to be used as background. |
| <code>use_weights</code> | Logical: calculate and use optimized weights for resources (i.e. one resource seems to be better than another, hence the former is considered with a higher weight). |
| <code>n_top_ligands</code> | How many of the top ligands to include in the ligand-target table. |
| <code>n_top_targets</code> | How many of the top targets (for each of the top ligands) to consider in the ligand-target table. |
| <code>signaling_network</code> | A list of parameters for building the signaling network, passed to nichenet_signaling_network . |
| <code>lr_network</code> | A list of parameters for building the ligand-receptor network, passed to nichenet_lr_network . |
| <code>gr_network</code> | A list of parameters for building the gene regulatory network, passed to nichenet_gr_network . |
| <code>small</code> | Logical: build a small network for testing purposes, using only OmniPath data. It is also a high quality network, it is reasonable to try the analysis with this small network. |
| <code>tiny</code> | Logical: build an even smaller network for testing purposes. As this involves random subsetting, it's not recommended to use this network for analysis. |
| <code>make_multi_objective_function_param</code> | Override parameters for <code>smoof::makeMultiObjectiveFunction</code> . |
| <code>objective_function_param</code> | Override additional arguments passed to the objective function. |
| <code>mlrmbo_optimization_param</code> | Override arguments for <code>nichenetr::mlrmbo_optimization</code> . |
| <code>construct_ligand_target_matrix_param</code> | Override parameters for <code>nichenetr::construct_ligand_target_matrix</code> . |
| <code>results_dir</code> | Character: path to the directory to save intermediate and final outputs from NicheNet methods. |
| <code>quality_filter_param</code> | Arguments for <code>filter_intercell_network</code> (quality filtering of the OmniPath ligand-receptor network). It is recommended to check these parameters and apply some quality filtering. The defaults already ensure certain filtering, but you might want more relaxed or stringent options. |

Details

About *small* and *tiny* networks: Building a NicheNet model is computationally demanding, taking several hours to run. As this is related to the enormous size of the networks, to speed up testing we

can use smaller networks, around 1,000 times smaller, with few thousands of interactions instead of few millions. Random subsetting of the whole network would result disjunct fragments, instead we load only a few resources. To run the whole pipeline with tiny networks use [nichenet_test](#).

Value

A named list with the intermediate and final outputs of the pipeline: ‘networks’, ‘expression’, ‘optimized_parameters’, ‘weighted_networks’ and ‘ligand_target_matrix’.

See Also

- [nichenet_networks](#)
- [nichenet_signaling_network](#)
- [nichenet_lr_network](#)
- [nichenet_gr_network](#)
- [nichenet_test](#)
- [nichenet_workarounds](#)
- [nichenet_results_dir](#)

Examples

```
nichenet_results <- nichenet_main(  
  # altering some network resource parameters, the rest  
  # of the resources will be loaded according to the defaults  
  signaling_network = list(  
    cpdb = NULL, # this resource will be excluded  
    inbiomap = NULL,  
    evex = list(min_confidence = 1.0) # override some parameters  
  ),  
  gr_network = list(only_omnipath = TRUE),  
  n_top_ligands = 20,  
  # override the default number of CPU cores to use  
  mlrmbo_optimization_param = list(ncores = 4)  
)
```

`nichenet_networks` *Builds NicheNet network prior knowledge*

Description

Builds network knowledge required by NicheNet. For this it calls a multitude of methods to download and combine data from various databases according to the settings. The content of the prior knowledge data is highly customizable, see the documentation of the related functions.

Usage

```
nichenet_networks(
  signaling_network = list(),
  lr_network = list(),
  gr_network = list(),
  only_omnipath = FALSE,
  small = FALSE,
  tiny = FALSE,
  quality_filter_param = list()
)
```

Arguments

| | |
|----------------------|--|
| signaling_network | A list of parameters for building the signaling network, passed to nichenet_signaling_network |
| lr_network | A list of parameters for building the ligand-receptor network, passed to nichenet_lr_network |
| gr_network | A list of parameters for building the gene regulatory network, passed to nichenet_gr_network |
| only_omnipath | Logical: a shortcut to use only OmniPath as network resource. |
| small | Logical: build a small network for testing purposes, using only OmniPath data. It is also a high quality network, it is reasonable to try the analysis with this small network. |
| tiny | Logical: build an even smaller network for testing purposes. As this involves random subsetting, it's not recommended to use this network for analysis. |
| quality_filter_param | Arguments for filter_intercell_network (quality filtering of the OmniPath ligand-receptor network). It is recommended to check these parameters and apply some quality filtering. The defaults already ensure certain filtering, but you might want more relaxed or stringent options. |

Value

A named list with three network data frames (tibbles): the signaling, the ligand-receptor (lr) and the gene regulatory (gr) networks.

See Also

- [nichenet_signaling_network](#)
- [nichenet_lr_network](#)
- [nichenet_gr_network](#)

Examples

```
networks <- nichenet_networks()
dplyr::sample_n(networks$gr_network, 10)
# # A tibble: 10 x 4
#   from     to     source      database
#   <chr>    <chr>   <chr>      <chr>
```

```

# 1 MAX    ALG3      harmonizome_ENCODE  harmonizome
# 2 MAX    IMPDH1    harmonizome_ENCODE  harmonizome
# 3 SMAD5   LCP1     Remap_5          Remap
# 4 HNF4A   TNFRSF19 harmonizome_CHEA   harmonizome
# 5 SMC3    FAP       harmonizome_ENCODE  harmonizome
# 6 E2F6    HIST1H1B harmonizome_ENCODE  harmonizome
# 7 TFAP2C  MAT2B    harmonizome_ENCODE  harmonizome
# 8 USF1    TBX4     harmonizome_TRANSFAC harmonizome
# 9 MIR133B FETUB    harmonizome_TRANSFAC harmonizome
# 10 SP4    HNRNPH2  harmonizome_ENCODE  harmonizome

# use only OmniPath:
omnipath_networks <- nichenet_networks(only_omnipath = TRUE)

```

nichenet_optimization *Optimizes NicheNet model parameters*

Description

Optimize NicheNet method parameters, i.e. PageRank parameters and source weights, basedon a collection of experiments where the effect of a ligand on gene expression was measured.

Usage

```

nichenet_optimization(
  networks,
  expression,
  make_multi_objective_function_param = list(),
  objective_function_param = list(),
  mlrmbo_optimization_param = list()
)

```

Arguments

| | |
|--|--|
| <code>networks</code> | A list with NicheNet format signaling, ligand-receptor and gene regulatory networks as produced by nichenet_networks . |
| <code>expression</code> | A list with expression data from ligand perturbation experiments, as produced by nichenet_expression_data . |
| <code>make_multi_objective_function_param</code> | Override parameters for <code>smoof::makeMultiObjectiveFunction</code> . |
| <code>objective_function_param</code> | Override additional arguments passed to the objective function. |
| <code>mlrmbo_optimization_param</code> | Override arguments for <code>nichenetr::mlrmbo_optimization</code> . |

Value

A result object from the function ‘mlrMBO::mbo’. Among other things, this contains the optimal parameter settings, the output corresponding to every input etc.

Examples

```
networks <- nichenet_networks()
expression <- nichenet_expression_data()
optimization_results <- nichenet_optimization(networks, expression)
```

nichenet_remove_orphan_ligands

Removes experiments with orphan ligands

Description

Removes from the expression data the perturbation experiments involving ligands without connections.

Usage

```
nichenet_remove_orphan_ligands(expression, lr_network)
```

Arguments

| | |
|------------|---|
| expression | Expression data as returned by nichenet_expression_data . |
| lr_network | A NicheNet format ligand-receptor network data frame as produced by nichenet_lr_network . |

Value

The same list as ‘expression’ with certain elements removed.

Examples

```
lr_network <- nichenet_lr_network()
expression <- nichenet_expression_data()
expression <- nichenet_remove_orphan_ligands(expression, lr_network)
```

nichenet_results_dir *Path to the current NicheNet results directory*

Description

Path to the directory to save intermediate and final outputs from NicheNet methods.

Usage

```
nichenet_results_dir()
```

Value

Character: path to the NicheNet results directory.

Examples

```
nichenet_results_dir()  
# [1] "nichenet_results"
```

nichenet_signaling_network
Builds a NicheNet signaling network

Description

Builds signaling network prior knowledge for NicheNet using multiple resources.

Usage

```
nichenet_signaling_network(  
  omnipath = list(),  
  pathwaycommons = list(),  
  harmonizome = list(),  
  vinayagam = list(),  
  cpdb = list(),  
  evex = list(),  
  inbiomap = list(),  
  only_omnipath = FALSE  
)
```

Arguments

| | |
|----------------|---|
| omnipath | List with paramaters to be passed to <i>nichenet_signaling_network_omnipath</i> . |
| pathwaycommons | List with paramaters to be passed to <i>nichenet_signaling_network_pathwaycommons</i> . |
| harmonizome | List with paramaters to be passed to <i>nichenet_signaling_network_harmonizome</i> . |
| vinayagam | List with paramaters to be passed to <i>nichenet_signaling_network_vinayagam</i> . |
| cpdb | List with paramaters to be passed to <i>nichenet_signaling_network_cpdb</i> . |
| evex | List with paramaters to be passed to <i>nichenet_signaling_network_evex</i> . |
| inbiomap | List with paramaters to be passed to <i>nichenet_signaling_network_inbiomap</i> . |
| only_omnipath | Logical: a shortcut to use only OmniPath as network resource. |

Value

A network data frame (tibble) with signaling interactions suitable for use with NicheNet.

See Also

- [nichenet_signaling_network_omnipath](#)
- [nichenet_signaling_network_pathwaycommons](#)
- [nichenet_signaling_network_harmonizome](#)
- [nichenet_signaling_network_vinayagam](#)
- [nichenet_signaling_network_cpdb](#)
- [nichenet_signaling_network_evex](#)
- [nichenet_signaling_network_inbiomap](#)

Examples

```
# load everything with the default parameters:
# we don't load inBio Map due to the - hopefully
# temporary - issues of their server
sig_network <- nichenet_signaling_network(inbiomap = NULL, cpdb = NULL)

# override parameters for some resources:
sig_network <- nichenet_signaling_network(
  omnipath = list(resources = c('SIGNOR', 'SignaLink3', 'SPIKE')),
  pathwaycommons = NULL,
  harmonizome = list(datasets = c('phosphositeplus', 'depod')),
  # we can not include this in everyday tests as it takes too long:
  # cpdb = list(complex_max_size = 1, min_score = .98),
  cpdb = NULL,
  evex = list(min_confidence = 1.5),
  inbiomap = NULL
)
# use only OmniPath:
sig_network_omnipath <- nichenet_signaling_network(only_omnipath = TRUE)
```

nichenet_signaling_network_cpdb

Builds signaling network for NicheNet using ConsensusPathDB

Description

Builds signaling network prior knowledge using ConsensusPathDB (CPDB) data. Note, the interactions from CPDB are not directed and many of them comes from complex expansion. Find out more at <http://cpdb.molgen.mpg.de/>.

Usage

```
nichenet_signaling_network_cpdb(...)
```

Arguments

... Passed to [consensuspathdb_download](#).

Value

A network data frame (tibble) with signaling interactions suitable for use with NicheNet.

See Also

- [nichenet_signaling_network](#)
- [consensuspathdb_download](#)

Examples

```
## Not run:  
# use some parameters stricter than default:  
cpdb_signaling_network <- nichenet_signaling_network_cpdb(  
  complex_max_size = 2,  
  min_score = .99  
)  
## End(Not run)
```

nichenet_signaling_network_evex*NicheNet signaling network from EVEX***Description**

Builds signaling network prior knowledge for NicheNet from the EVEX database.

Usage

```
nichenet_signaling_network_evex(top_confidence = 0.75, indirect = FALSE, ...)
```

Arguments

- `top_confidence` Double, between 0 and 1. Threshold based on the quantile of the confidence score.
- `indirect` Logical: whether to include indirect interactions.
- `...` Ignored.

Value

A network data frame (tibble) with signaling interactions suitable for use with NicheNet.

See Also

- [evex_download](#)
- [nichenet_signaling_network](#)

Examples

```
ev_signaling_network <- nichenet_signaling_network_evex(
  top_confidence = .9
)
```

nichenet_signaling_network_harmonizome*NicheNet signaling network from Harmonizome***Description**

Builds signaling network prior knowledge for NicheNet using Harmonizome

Usage

```
nichenet_signaling_network_harmonizome(  
  datasets = c("phosphositeplus", "kea", "depod"),  
  ...  
)
```

Arguments

- | | |
|----------|---|
| datasets | The datasets to use. For possible values please refer to default value and the Harmonizome webpage. |
| ... | Ignored. |

Value

A network data frame (tibble) with signaling interactions suitable for use with NicheNet.

Examples

```
# use only KEA and PhosphoSite:  
hz_signaling_network <- nichenet_signaling_network_harmonizome(  
  datasets = c('kea', 'phosphositeplus')  
)
```

nichenet_signaling_network_inbiomap

NicheNet signaling network from InWeb InBioMap

Description

Builds signaling network prior knowledge for NicheNet from the InWeb InBioMap database.

Usage

```
nichenet_signaling_network_inbiomap(...)
```

Arguments

- | | |
|-----|----------|
| ... | Ignored. |
|-----|----------|

Value

A network data frame (tibble) with signaling interactions suitable for use with NicheNet.

See Also

[nichenet_signaling_network](#), [inbiomap_download](#)

Examples

```
ib_signaling_network <- nichenet_signaling_network_inbiomap()
```

nichenet_signaling_network_omnipath

Builds signaling network for NicheNet using OmniPath

Description

Retrieves network prior knowledge from OmniPath and provides it in a format suitable for Nich-eNet. This method never downloads the ‘ligrecrextra’ dataset because the ligand-receptor interactions are supposed to come from [nichenet_lr_network_omnipath](#).

Usage

```
nichenet_signaling_network_omnipath(min_curation_effort = 0, ...)
```

Arguments

| | |
|---------------------|--|
| min_curation_effort | Lower threshold for curation effort |
| ... | Passed to import_post_translational_interactions |

Value

A network data frame (tibble) with signaling interactions suitable for use with NicheNet.

See Also

- [nichenet_signaling_network](#)

Examples

```
# use interactions with at least 2 evidences (reference or database)
op_signaling_network <- nichenet_signaling_network_omnipath(
  min_curation_effort = 2
)
```

nichenet_signaling_network_pathwaycommons

NicheNet signaling network from PathwayCommons

Description

Builds signaling network prior knowledge for NicheNet using PathwayCommons.

Usage

```
nichenet_signaling_network_pathwaycommons(  
  interaction_types = c("catalysis-precedes", "controls-phosphorylation-of",  
    "controls-state-change-of", "controls-transport-of", "in-complex-with",  
    "interacts-with"),  
  ...  
)
```

Arguments

interaction_types
Character vector with PathwayCommons interaction types. Please refer to the default value and the PathwayCommons webpage.
... Ignored.

Value

A network data frame (tibble) with signaling interactions suitable for use with NicheNet.

Examples

```
# use only the "controls-transport-of" interactions:  
pc_signaling_network <- nichenet_signaling_network_pathwaycommons(  
  interaction_types = 'controls-transport-of'  
)
```

nichenet_signaling_network_vinayagam

NicheNet signaling network from Vinayagam

Description

Builds signaling network prior knowledge for NicheNet using Vinayagam 2011 Supplementary Table S6. Find out more at <https://doi.org/10.1126/scisignal.2001699>.

Usage

```
nichenet_signaling_network_vinayagam(...)
```

Arguments

... Ignored.

Value

A network data frame (tibble) with signaling interactions suitable for use with NicheNet.

Examples

```
vi_signaling_network <- nichenet_signaling_network_vinayagam()
```

nichenet_test

Run the NicheNet pipeline with a little dummy network

Description

Loads a tiny network and runs the NicheNet pipeline with low number of iterations in the optimization process. This way the pipeline runs in a reasonable time in order to test the code. Due to the random subsampling disconnected networks might be produced sometimes. If you see an error like "Error in if (sd(prediction_vector) == 0) ... missing value where TRUE/FALSE needed", the random subsampled input is not appropriate. In this case just interrupt and call again. This test ensures the computational integrity of the pipeline. If it fails during the optimization process, try to start it over several times, even restarting R. The unpredictability is related to codemlrMBO and nichenetr not being prepared to handle certain conditions, and it's also difficult to find out which conditions lead to which errors. At least 3 different errors appear time to time, depending on the input. It also seems like restarting R sometimes helps, suggesting that the entire system might be somehow stateful. You can ignore the Parallelization was not stopped warnings on repeated runs.

Usage

```
nichenet_test(...)
```

Arguments

... Passed to [nichenet_main](#).

Value

A named list with the intermediate and final outputs of the pipeline: ‘networks’, ‘expression’, ‘optimized_parameters’, ‘weighted_networks’ and ‘ligand_target_matrix’.

Examples

```
nnt <- nichenet_test()
```

nichenet_workarounds *Workarounds using NicheNet without attaching the package*

Description

NicheNet requires the availability of some lazy loaded external data which are not available if the package is not loaded and attached. Also, the BBmisc::convertToShortString used for error reporting in mlrMBO::evalTargetFun.OptState is patched here to print longer error messages. Maybe it's a better solution to attach nichenetr before running the NicheNet pipeline. Alternatively you can try to call this function in the beginning. Why we don't call this automatically is just because we don't want to load datasets from another package without the user knowing about it.

Usage

```
nichenet_workarounds()
```

Value

Returns NULL.

Examples

```
nichenet_workarounds()
```

obo_parser *Generic OBO parser*

Description

Reads the contents of an OBO file and processes it into data frames or a list based data structure.

Usage

```
obo_parser(  
  path,  
  relations = c("is_a", "part_of", "occurs_in", "regulates", "positively_regulates",  
              "negatively_regulates"),  
  shorten_namespace = TRUE,  
  tables = TRUE  
)
```

Arguments

| | |
|--------------------------------|---|
| <code>path</code> | Path to the OBO file. |
| <code>relations</code> | Character vector: process only these relations. |
| <code>shorten_namespace</code> | Logical: shorten the namespace to a single letter code (as usual for Gene Ontology, e.g. cellular_component = "C"). |
| <code>tables</code> | Logical: return data frames (tibbles) instead of nested lists. |

Value

A list with the following elements: 1) "names" a list with terms as names and names as values; 2) "namespaces" a list with terms as names and namespaces as values; 3) "relations" a list with relations between terms: terms are keys, values are lists with relations as names and character vectors of related terms as values; 4) "subsets" a list with terms as keys and character vectors of subset names as values (or NULL if the term does not belong to any subset); 5) "obsolete" character vector with all the terms labeled as obsolete. If the `tables` parameter is TRUE, "names", "namespaces", "relations" and "subsets" will be data frames (tibbles).

See Also

- [relations_list_to_table](#)
- [relations_table_to_list](#)
- [swap_relations](#)

Examples

```
goslim_url <-  
  "http://current.geneontology.org/ontology/subsets/goslim_generic.obo"  
path <- tempfile()  
httr::GET(goslim_url, httr::write_disk(path, overwrite = TRUE))  
obo <- obo_parser(path, tables = FALSE)  
unlink(path)  
names(ob)  
# [1] "names"      "namespaces" "relations"   "subsets"    "obsolete"  
head(ob$relations, n = 2)  
# $`GO:0000001`  
# $`GO:0000001`$is_a  
# [1] "GO:0048308" "GO:0048311"  
#  
# $`GO:0000002`  
# $`GO:0000002`$is_a  
# [1] "GO:0007005"
```

Description

OmnipathR is an R package built to provide easy access to the data stored in the OmniPath web service:

<https://omnipathdb.org/>

And a number of other resources, such as BioPlex, ConsensusPathDB, EVEX, Guide to Pharmacology (IUPHAR/BPS), Harmonizome, HTRIdb, InWeb InBioMap, KEGG Pathway, Pathway Commons, Ramiłowski et al. 2015, RegNetwork, ReMap, TF census, TRRUST and Vinayagam et al. 2011.

The OmniPath web service implements a very simple REST style API. This package make requests by the HTTP protocol to retrieve the data. Hence, fast Internet access is required for a proper use of OmnipathR.

The package also provides some utility functions to filter, analyse and visualize the data. Furthermore, OmnipathR features a close integration with the NicheNet method for ligand activity prediction from transcriptomics data, and its R implementation nichenet (available in CRAN).

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and Attila Gabor <gaborattila87@gmail.com>>

Examples

```
# Download post-translational modifications:  
enzsub <- import_omnipath_enzsub(resources = c("PhosphoSite", "SIGNOR"))  
  
# Download protein-protein interactions  
interactions <- import_omnipath_interactions(resources = c("SignalLink3"))  
  
# Convert to igraph objects:  
enzsub_g <- enzsub_graph(enzsub = enzsub)  
OPI_g <- interaction_graph(interactions = interactions )  
  
# Print some interactions:  
print_interactions(head(ptms))  
  
# interactions with references:  
print_interactions(tail(ptms), writeRefs=TRUE)  
  
# find interactions between kinase and substrate:  
print_interactions(dplyr::filter(ptms, enzyme_genesymbol=="MAP2K1",  
    substrate_genesymbol=="MAPK3"))  
  
# find shortest paths on the directed network between proteins
```

```

print_path_es(shortest_paths(OPI_g, from = "TYR03", to = "STAT3",
  output = 'epath')$epath[[1]], OPI_g)

# find all shortest paths between proteins
print_path_vs(
  all_shortest_paths(
    enzsub_g,
    from = "SRC",
    to = "STAT1"
  )$res,
  enzsub_g
)

```

omnipath_cache_autoclean*Keeps only the latest versions of complete downloads***Description**

Removes the old versions, the failed downloads and the files in the cache directory which are missing from the database. For more flexible operations use [omnipath_cache_remove](#) and [omnipath_cache_clean](#).

Usage

```
omnipath_cache_autoclean()
```

Value

Invisibl returns the cache database (list of cache records).

Examples

```
omnipath_cache_autoclean()
```

omnipath_cache_clean *Removes the items from the cache directory which are unknown by the cache database***Description**

Removes the items from the cache directory which are unknown by the cache database

Usage

```
omnipath_cache_clean()
```

Value

Returns ‘NULL’.

Examples

```
omnipath_cache_clean()
```

omnipath_cache_clean_db

Removes the cache database entries without existing files

Description

Removes the cache database entries without existing files

Usage

```
omnipath_cache_clean_db(...)
```

Arguments

... Ignored.

Value

Returns ‘NULL’.

Examples

```
omnipath_cache_clean_db()
```

omnipath_cache_download_ready*Sets the download status to ready for a cache item***Description**

Sets the download status to ready for a cache item

Usage

```
omnipath_cache_download_ready(version, key = NULL)
```

Arguments

| | |
|----------------------|---|
| <code>version</code> | Version of the cache item. If does not exist a new version item will be created |
| <code>key</code> | Key of the cache item |

Value

Character: invisibly returns the version number of the cache version item.

Examples

```
bioc_url <- 'https://bioconductor.org/'  
# request a new version item (or retrieve the latest)  
new_version <- omnipath_cache_latest_or_new(url = bioc_url)  
# check if the version item is not a finished download  
new_version$status  
# [1] "unknown"  
# download the file  
httr::GET(bioc_url, httr::write_disk(new_version$path, overwrite = TRUE))  
# report to the cache database that the download is ready  
omnipath_cache_download_ready(new_version)  
# now the status is ready:  
version <- omnipath_cache_latest_or_new(url = bioc_url)  
version$status  
# "ready"  
version$dl_finished  
# [1] "2021-03-09 16:48:38 CET"  
omnipath_cache_remove(url = bioc_url) # cleaning up
```

omnipath_cache_filter_versions
Filters the versions from one cache record

Description

Filters the versions based on multiple conditions: their age and status

Usage

```
omnipath_cache_filter_versions(
  record,
  latest = FALSE,
  max_age = NULL,
  min_age = NULL,
  status = CACHE_STATUS$READY
)
```

Arguments

| | |
|---------|--|
| record | A cache record |
| latest | Return the most recent version |
| max_age | The maximum age in days (e.g. 5: 5 days old or more recent) |
| min_age | The minimum age in days (e.g. 5: 5 days old or older) |
| status | Character vector with status codes. By default only the versions with ‘ready’ (completed download) status are selected |

Value

Character vector with version IDs, NA if no version satisfies the conditions.

Examples

```
# creating an example cache record
bioc_url <- 'https://bioconductor.org/'
version <- omnipath_cache_latest_or_new(url = bioc_url)
httr::GET(bioc_url, httr::write_disk(version$path, overwrite = TRUE))
omnipath_cache_download_ready(version)
record <- dplyr::first(omnipath_cache_search('biocond'))

# only the versions with status "ready"
version_numbers <- omnipath_cache_filter_versions(record, status = 'ready')
omnipath_cache_remove(url = bioc_url) # cleaning up
```

`omnipath_cache_get` *Retrieves one item from the cache directory*

Description

Retrieves one item from the cache directory

Usage

```
omnipath_cache_get(
  key = NULL,
  url = NULL,
  post = NULL,
  payload = NULL,
  create = TRUE,
  ...
)
```

Arguments

| | |
|---------|--|
| key | The key of the cache record |
| url | URL pointing to the resource |
| post | HTTP POST parameters as a list |
| payload | HTTP data payload |
| create | Create a new entry if doesn't exist yet |
| ... | Passed to <code>omnipath_cache_record</code> (internal function) |

Value

Cache record: an existing record if the entry already exists, otherwise a newly created and inserted record

Examples

```
# create an example cache record
bioc_url <- 'https://bioconductor.org/'
version <- omnipath_cache_latest_or_new(url = bioc_url)
omnipath_cache_remove(url = bioc_url) # cleaning up

# retrieve the cache record
record <- omnipath_cache_get(url = bioc_url)
record$key
# [1] "41346a00fb20d2a9df03aa70cf4d50bf88ab154a"
record$url
# [1] "https://bioconductor.org/"
```

omnipath_cache_key *Generates a hash which identifies an element in the cache database*

Description

Generates a hash which identifies an element in the cache database

Usage

```
omnipath_cache_key(url, post = NULL, payload = NULL)
```

Arguments

| | |
|---------|---|
| url | Character vector with URLs |
| post | List with the HTTP POST parameters or a list of lists if the url vector is longer than 1. NULL for queries without POST parameters. |
| payload | HTTP data payload. List with multiple items if the url vector is longer than 1. NULL for queries without data. |

Value

Character vector of cache record keys.

Examples

```
bioc_url <- 'https://bioconductor.org/'  
omnipath_cache_key(bioc_url)  
# [1] "41346a00fb20d2a9df03aa70cf4d50bf88ab154a"
```

omnipath_cache_latest_or_new
The latest or a new version of a cache record

Description

Looks up a record in the cache and returns its latest valid version. If the record doesn't exist or no valid version available, creates a new one.

Usage

```
omnipath_cache_latest_or_new(
  key = NULL,
  url = NULL,
  post = NULL,
  payload = NULL,
  create = TRUE,
  ...
)
```

Arguments

| | |
|---------|---|
| key | The key of the cache record |
| url | URL pointing to the resource |
| post | HTTP POST parameters as a list |
| payload | HTTP data payload |
| create | Logical: whether to create and return a new version. If FALSE only the latest existing valid version is returned, if available. |
| ... | Passed to omnipath_cache_get |

Value

A cache version item.

Examples

```
# retrieve the latest version of the first cache record
# found by the search keyword "bioplex"
latest_bioplex <-
  omnipath_cache_latest_or_new(
    names(omnipath_cache_search('bioplex'))[1]
  )

latest_bioplex$dl_finished
# [1] "2021-03-09 14:28:50 CET"
latest_bioplex$path
# [1] "/home/denes/.cache/OmnipathR/378e0def2ac97985f629-1.rds"

# create an example cache record
bioc_url <- 'https://bioconductor.org/'
version <- omnipath_cache_latest_or_new(url = bioc_url)
omnipath_cache_remove(url = bioc_url) # cleaning up
```

omnipath_cache_latest_version

Finds the most recent version in a cache record

Description

Finds the most recent version in a cache record

Usage

```
omnipath_cache_latest_version(record)
```

Arguments

record A cache record

Value

Character: the version ID with the most recent download finished time

omnipath_cache_load *Loads an R object from the cache***Description**

Loads the object from RDS format.

Usage

```
omnipath_cache_load(  
  key = NULL,  
  version = NULL,  
  url = NULL,  
  post = NULL,  
  payload = NULL  
)
```

Arguments

| | |
|---------|--|
| key | Key of the cache item |
| version | Version of the cache item. If does not exist or NULL, the latest version will be retrieved |
| url | URL of the downloaded resource |
| post | HTTP POST parameters as a list |
| payload | HTTP data payload |

Value

Object loaded from the cache RDS file.

See Also

[omnipath_cache_save](#)

Examples

```
url <- paste0(
  'https://omnipathdb.org/intercell?resources=Adhesome,Almen2009,',
  'Baccin2019,CSPA,CellChatDB&license=academic'
)
result <- read.delim(url, sep = '\t')
omnipath_cache_save(result, url = url)
# works only if you have already this item in the cache
intercell_data <- omnipath_cache_load(url = url)
class(intercell_data)
# [1] "data.frame"
nrow(intercell_data)
# [1] 16622
attr(intercell_data, 'origin')
# [1] "cache"

# basic example of saving and loading to and from the cache:
bioc_url <- 'https://bioconductor.org/'
bioc_html <- readChar(url(bioc_url), nchars = 99999)
omnipath_cache_save(bioc_html, url = bioc_url)
bioc_html <- omnipath_cache_load(url = bioc_url)
```

omnipath_cache_move_in

Moves an existing file into the cache

Description

Either the key or the URL (with POST and payload) must be provided.

Usage

```
omnipath_cache_move_in(
  path,
  key = NULL,
  version = NULL,
  url = NULL,
  post = NULL,
  payload = NULL,
  keep_original = FALSE
)
```

Arguments

| | |
|---------------|---|
| path | Path to the source file |
| key | Key of the cache item |
| version | Version of the cache item. If does not exist a new version item will be created |
| url | URL of the downloaded resource |
| post | HTTP POST parameters as a list |
| payload | HTTP data payload |
| keep_original | Whether to keep or remove the original file |

Value

Character: invisibly returns the version number of the cache version item.

See Also

[omnipath_cache_save](#)

Examples

```
path <- tempfile()
saveRDS(rnorm(100), file = path)
omnipath_cache_move_in(path, url = 'the_download_address')

# basic example of moving a file to the cache:

bioc_url <- 'https://bioconductor.org/'
html_file <- tempfile(fileext = '.html')
httr::GET(bioc_url, httr::write_disk(html_file, overwrite = TRUE))
omnipath_cache_move_in(path = html_file, url = bioc_url)
omnipath_cache_remove(url = bioc_url) # cleaning up
```

omnipath_cache_remove *Removes contents from the cache directory*

Description

According to the parameters, it can remove contents older than a certain age, or contents having a more recent version, one specific item, or wipe the entire cache.

Usage

```
omnipath_cache_remove(key = NULL, url = NULL, post = NULL,
                      payload = NULL, max_age = NULL, min_age = NULL, status = NULL,
                      only_latest = FALSE, wipe = FALSE, autoclean = TRUE)
```

Arguments

| | |
|-------------|--|
| key | The key of the cache record |
| url | URL pointing to the resource |
| post | HTTP POST parameters as a list |
| payload | HTTP data payload |
| max_age | Age of cache items in days. Remove everything that is older than this age |
| min_age | Age of cache items in days. Remove everything more recent than this age |
| status | Remove items having any of the states listed here |
| only_latest | Keep only the latest version |
| wipe | Logical: if TRUE, removes all files from the cache and the cache database. Same as calling omnipath_cache_wipe . |
| autoclean | Remove the entries about failed downloads, the files in the cache directory which are missing from the cache database, and the entries without existing files in the cache directory |

Value

Invisibly returns the cache database (list of cache records).

See Also

- [omnipath_cache_wipe](#)
- [omnipath_cache_clean](#)
- [omnipath_cache_autoclean](#)

Examples

```
# remove all cache data from the BioPlex database
cache_records <- omnipath_cache_search(
  'bioplex',
  ignore.case = TRUE
)
omnipath_cache_remove(names(cache_records))

# remove a record by its URL
regnetwork_url <- 'http://www.regnetworkweb.org/download/human.zip'
omnipath_cache_remove(url = regnetwork_url)

# remove all records older than 30 days
omnipath_cache_remove(max_age = 30)

# for each record, remove all versions except the latest
omnipath_cache_remove(only_latest = TRUE)

bioc_url <- 'https://bioconductor.org/'
version <- omnipath_cache_latest_or_new(url = bioc_url)
```

```
httr::GET(bioc_url, httr::write_disk(version$path, overwrite = TRUE))
omnipath_cache_download_ready(version)
key <- omnipath_cache_key(bioc_url)
omnipath_cache_remove(key = key)
```

omnipath_cache_save *Saves an R object to the cache*

Description

Exports the object in RDS format, creates new cache record if necessary.

Usage

```
omnipath_cache_save(
  data,
  key = NULL,
  version = NULL,
  url = NULL,
  post = NULL,
  payload = NULL
)
```

Arguments

| | |
|---------|---|
| data | An object |
| key | Key of the cache item |
| version | Version of the cache item. If does not exist a new version item will be created |
| url | URL of the downloaded resource |
| post | HTTP POST parameters as a list |
| payload | HTTP data payload |

Value

Returns invisibly the data itself.

Invisibly returns the ‘data’.

See Also

[omnipath_cache_move_in](#)

Examples

```
mydata <- data.frame(a = c(1, 2, 3), b = c('a', 'b', 'c'))
omnipath_cache_save(mydata, url = 'some_dummy_address')
from_cache <- omnipath_cache_load(url = 'some_dummy_address')
from_cache
#   a b
# 1 1 a
# 2 2 b
# 3 3 c
attr(from_cache, 'origin')
# [1] "cache"

# basic example of saving and loading to and from the cache:
bioc_url <- 'https://bioconductor.org/'
bioc_html <- readChar(url(bioc_url), nchars = 99999)
omnipath_cache_save(bioc_html, url = bioc_url)
bioc_html <- omnipath_cache_load(url = bioc_url)
```

omnipath_cache_search *Searches for cache items*

Description

Searches the cache records by matching the URL against a string or regexp.

Usage

```
omnipath_cache_search(pattern, ...)
```

Arguments

| | |
|---------|-------------------------------|
| pattern | String or regular expression. |
| ... | Passed to grep |

Value

List of cache records matching the pattern.

Examples

```
# find all cache records from the BioPlex database
bioplex_cache_records <- omnipath_cache_search(
  'bioplex',
  ignore.case = TRUE
)
```

omnipath_cache_set_ext

Sets the file extension for a cache record

Description

Sets the file extension for a cache record

Usage

```
omnipath_cache_set_ext(key, ext)
```

Arguments

| | |
|-----|---|
| key | Character: key for a cache item, alternatively a version entry. |
| ext | Character: the file extension, e.g. "zip". |

Value

Returns 'NULL'.

Examples

```
bioc_url <- 'https://bioconductor.org/'  
version <- omnipath_cache_latest_or_new(url = bioc_url)  
version$path  
# [1] "/home/denes/.cache/OmnipathR/41346a00fb20d2a9df03-1"  
httr::GET(bioc_url, httr::write_disk(version$path, overwrite = TRUE))  
key <- omnipath_cache_key(url = bioc_url)  
omnipath_cache_set_ext(key = key, ext = 'html')  
version <- omnipath_cache_latest_or_new(url = bioc_url)  
version$path  
# [1] "/home/denes/.cache/OmnipathR/41346a00fb20d2a9df03-1.html"  
record <- omnipath_cache_get(url = bioc_url)  
record$ext  
# [1] "html"  
omnipath_cache_remove(url = bioc_url) # cleaning up
```

omnipath_cache_update_status*Updates the status of an existing cache record***Description**

Updates the status of an existing cache record

Usage

```
omnipath_cache_update_status(key, version, status,
                             dl_finished = NULL)
```

Arguments

| | |
|--------------------------|--|
| <code>key</code> | Key of the cache item |
| <code>version</code> | Version of the cache item. If does not exist a new version item will be created |
| <code>status</code> | The updated status value |
| <code>dl_finished</code> | Timestamp for the time when download was finished, if ‘NULL’ the value remains unchanged |

Value

Character: invisibly returns the version number of the cache version item.

Examples

```
bioc_url <- 'https://bioconductor.org/'
latest_version <- omnipath_cache_latest_or_new(url = bioc_url)
key <- omnipath_cache_key(bioc_url)
omnipath_cache_update_status(
  key = key,
  version = latest_version$number,
  status = 'ready',
  dl_finished = Sys.time()
)
omnipath_cache_remove(url = bioc_url) # cleaning up
```

omnipath_cache_wipe *Permanently removes all the cache contents*

Description

After this operation the cache directory will be completely empty, except an empty cache database file.

Usage

`omnipath_cache_wipe(...)`

Arguments

... Ignored.

Value

Returns ‘NULL’.

See Also

[omnipath_cache_remove](#)

Examples

```
omnipath_cache_wipe()  
# the cache is completely empty:  
print(omnipath.env$cache)  
# list()  
list.files(omnipath_get_cachedir())  
# [1] "cache.json"
```

omnipath_get_config_path
 Current config file path

Description

Current config file path

Usage

`omnipath_get_config_path(user = FALSE)`

Arguments

| | |
|-------------------|---|
| <code>user</code> | Logical: prioritize the user level config even if a config in the current working directory is available. |
|-------------------|---|

Value

Character: path to the config file.

Examples

```
omnipath_get_config_path()
```

`omnipath_load_config` *Load the package configuration from a config file*

Description

Load the package configuration from a config file

Usage

```
omnipath_load_config(path = NULL, title = "default", user = FALSE, ...)
```

Arguments

| | |
|--------------------|---|
| <code>path</code> | Path to the config file. |
| <code>title</code> | Load the config under this title. One config file might contain multiple configurations, each identified by a title. If the title is not available the first section of the config file will be used. |
| <code>user</code> | Force to use the user level config even if a config file exists in the current directory. By default, the local config files have priority over the user level config. |
| <code>...</code> | Passed to <code>yaml::yaml.load_file</code> . |

Value

Invisibly returns the config as a list.

Examples

```
# load the config from a custom config file:
omnipath_load_config(path = 'my_custom_omnipath_config.yml')
```

| | |
|---------------------------|--|
| <code>omnipath_log</code> | <i>Browse the current OmnipathR log file</i> |
|---------------------------|--|

Description

Browse the current OmnipathR log file

Usage

```
omnipath_log()
```

Value

Returns ‘NULL’.

See Also

[omnipath_logfile](#)

Examples

```
omnipath_log()  
# then you can browse the log file, and exit with `q`
```

| | |
|-------------------------------|---|
| <code>omnipath_logfile</code> | <i>Path to the current OmnipathR log file</i> |
|-------------------------------|---|

Description

Path to the current OmnipathR log file

Usage

```
omnipath_logfile()
```

Value

Character: path to the current logfile, or NULL if no logfile is available.

See Also

[omnipath_log](#)

Examples

```
omnipath_logfile()
# [1] "/home/denes/omnipathr/omnipathr-log/omnipathr-20210309-1642.log"
```

omnipath_msg

Dispatch a message to the OmnipathR logger

Description

Any package or script can easily send log messages and establish a logging facility with the fantastic ‘logger’ package. This function serves the only purpose if you want to inject messages into the logger of OmnipathR. Otherwise we recommend to use the ‘logger’ package directly.

Usage

```
omnipath_msg(level, ...)
```

Arguments

| | |
|-------|---|
| level | Character, numeric or class loglevel. A log level, if character one of the following: "fatal", "error", "warn", "success", "info", "trace". |
| ... | Arguments for string formatting, passed <code>sprintf</code> or <code>str_glue</code> . |

Value

Returns ‘NULL’.

Examples

```
omnipath_msg(
  level = 'success',
  'Talking to you in the name of OmnipathR, my favourite number is %d',
  round(runif(1, 1, 10))
)
```

`omnipath_reset_config` Restores the built-in default values of all config parameters

Description

Restores the built-in default values of all config parameters

Usage

```
omnipath_reset_config(save = NULL, reset_all = FALSE)
```

Arguments

- | | |
|------------------------|---|
| <code>save</code> | If a path, the restored config will be also saved to this file. If TRUE, the config will be saved to the current default config path (see omnipath_get_config_path). |
| <code>reset_all</code> | Reset to their defaults also the options already set in the R options. |

Value

The config as a list.

See Also

[omnipath_load_config](#), [omnipath_save_config](#)

Examples

```
# restore the defaults and write them to the default config file:  
omnipath_reset_config()  
omnipath_save_config()
```

`omnipath_save_config` Save the current package configuration

Description

Save the current package configuration

Usage

```
omnipath_save_config(path = NULL, title = "default", local = FALSE)
```

Arguments

| | |
|--------------------|--|
| <code>path</code> | Path to the config file. Directories and the file will be created if don't exist. |
| <code>title</code> | Save the config under this title. One config file might contain multiple configurations, each identified by a title. |
| <code>local</code> | Save into a config file in the current directory instead of a user level config file. When loading, the config in the current directory has priority over the user level config. |

Value

Returns 'NULL'.

Examples

```
# after this, all downloads will default to commercial licenses
# i.e. the resources that allow only academic use will be excluded:
options(omnipath.license = 'commercial')
omnipath_save_config()
```

`omnipath_set_cachedir` *Change the cache directory*

Description

Change the cache directory

Usage

```
omnipath_set_cachedir(path = NULL)
```

Arguments

| | |
|-------------------|--|
| <code>path</code> | Character: path to the new cache directory. If don't exist, the directories will be created. If the path is an existing cache directory, the package's cache database for the current session will be loaded from the database in the directory. If NULL, the cache directory will be set to its default path. |
|-------------------|--|

Value

Returns NULL.

Examples

```
tmp_cache <- tempdir()  
omnipath_set_cachedir(tmp_cache)  
# restore the default cache directory:  
omnipath_set_cachedir()
```

omnipath_set_console_loglevel

Sets the log level for the console

Description

Use this method to change during a session which messages you want to be printed on the console. Before loading the package, you can set it also by the config file, with the omnipath.console_loglevel key.

Usage

```
omnipath_set_console_loglevel(level)
```

Arguments

level Character or class ‘loglevel’. The desired log level.

Value

Returns ‘NULL’.

See Also

[omnipath_set_logfile_loglevel](#)

Examples

```
omnipath_set_console_loglevel('warn')  
# or:  
omnipath_set_console_loglevel(logger::WARN)
```

omnipath_set_logfile_loglevel
Sets the log level for the logfile

Description

Use this method to change during a session which messages you want to be written into the logfile. Before loading the package, you can set it also by the config file, with the omnipath.loglevel key.

Usage

```
omnipath_set_logfile_loglevel(level)
```

Arguments

| | |
|-------|---|
| level | Character or class ‘loglevel’. The desired log level. |
|-------|---|

Value

Returns ‘NULL’.

See Also

[omnipath_set_console_loglevel](#)

Examples

```
omnipath_set_logfile_loglevel('info')
# or:
omnipath_set_logfile_loglevel(logger::INFO)
```

omnipath_set_loglevel *Sets the log level for the package logger*

Description

Sets the log level for the package logger

Usage

```
omnipath_set_loglevel(level, target = "logfile")
```

Arguments

| | |
|--------|---|
| level | Character or class ‘loglevel’. The desired log level. |
| target | Character, either ‘logfile’ or ‘console’ |

Value

Returns ‘NULL’.

Examples

```
omnipath_set_loglevel(logger::FATAL, target = 'console')
```

omnipath_show_db

Built in database definitions

Description

Databases are resources which might be costly to load but can be used many times by functions which usually automatically load and retrieve them from the database manager. Each database has a lifetime and will be unloaded automatically upon expiry.

Usage

```
omnipath_show_db()
```

Value

A data frame with the built in database definitions.

Examples

```
database_definitions <- omnipath_show_db()
database_definitions
# # A tibble: 14 x 10
#   name      last_used      lifetime package  loader    loader_p.
#   <chr>     <dttm>       <dbl> <chr>    <chr>    <list>
# 1 Gene Onto. 2021-04-04 20:19:15     300 Omnipat. go_ontol. <named l.
# 2 Gene Onto. NA          300 Omnipat. go_ontol. <named l.
# 3 Gene Onto. NA          300 Omnipat. go_ontol. <named l.
# 4 Gene Onto. NA          300 Omnipat. go_ontol. <named l.
# 5 Gene Onto. NA          300 Omnipat. go_ontol. <named l.
# ... (truncated)
# # . with 4 more variables: latest_param <list>, loaded <lgl>, db <list>,
# #   key <chr>
```

omnipath_unlock_cache_db

Removes the lock file from the cache directory

Description

A lock file in the cache directory avoids simultaneous write and read. It's supposed to be removed after each read and write operation. This might not happen if the process crashes during such an operation. In this case you can manually call this function.

Usage

```
omnipath_unlock_cache_db()
```

Value

Logical: returns TRUE if the cache was locked and now is unlocked; FALSE if it was not locked.

Examples

```
omnipath_unlock_cache_db()
```

ontology_ensure_id *Only ontology IDs*

Description

Converts a mixture of ontology IDs and names to only IDs. If an element of the input is missing from the chosen ontology it will be dropped. This can happen if the ontology is a subset (slim) version, but also if the input is not a valid ID or name.

Usage

```
ontology_ensure_id(terms, db_key = "go_basic")
```

Arguments

| | |
|--------|---|
| terms | Character: ontology IDs or term names. |
| db_key | Character: key to identify the ontology database. For the available keys see omnipath_show_db . |

Value

Character vector of ontology IDs.

Examples

```
ontology_ensure_id(c('mitochondrion inheritance', 'GO:0001754'))  
# [1] "GO:0000001" "GO:0001754"
```

ontology_ensure_name *Only ontology term names*

Description

Converts a mixture of ontology IDs and names to only names. If an element of the input is missing from the chosen ontology it will be dropped. This can happen if the ontology is a subset (slim) version, but also if the input is not a valid ID or name.

Usage

```
ontology_ensure_name(terms, db_key = "go_basic")
```

Arguments

| | |
|--------|---|
| terms | Character: ontology IDs or term names. |
| db_key | Character: key to identify the ontology database. For the available keys see omnipath_show_db . |

Value

Character vector of ontology term names.

Examples

```
ontology_ensure_name(c('reproduction', 'GO:0001754', 'foo bar'))  
# [1] "eye photoreceptor cell differentiation" "reproduction"
```

ontology_name_id *Translate between ontology IDs and names*

Description

Makes sure that the output contains only valid IDs or term names. The input can be a mixture of IDs and names. The order of the input won't be preserved in the output.

Usage

```
ontology_name_id(terms, ids = TRUE, db_key = "go_basic")
```

Arguments

| | |
|--------|---|
| terms | Character: ontology IDs or term names. |
| ids | Logical: the output should contain IDs or term names. |
| db_key | Character: key to identify the ontology database. For the available keys see omnipath_show_db . |

Value

Character vector of ontology IDs or term names.

Examples

```
ontology_name_id(c('mitochondrion inheritance', 'reproduction'))
# [1] "GO:0000001" "GO:0000003"
ontology_name_id(c('GO:0000001', 'reproduction'), ids = FALSE)
# [1] "mitochondrion inheritance" "reproduction"
```

pathwaycommons_download

Interactions from PathwayCommons

Description

PathwayCommons (<http://www.pathwaycommons.org/>) provides molecular interactions from a number of databases, in either BioPAX or SIF (simple interaction format). This function retrieves all interactions in SIF format. The data is limited to the interacting pair and the type of the interaction.

Usage

```
pathwaycommons_download()
```

Value

A data frame (tibble) with interactions.

Examples

```
pc_interactions <- pathwaycommons_download()
pc_interactions
# # A tibble: 1,884,849 x 3
#   from      type          to
#   <chr>    <chr>        <chr>
# 1 A1BG    controls-expression-of  A2M
# 2 A1BG    interacts-with     ABCC6
# 3 A1BG    interacts-with     ACE2
# 4 A1BG    interacts-with     ADAM10
# 5 A1BG    interacts-with     ADAM17
```

```
# # . with 1,884,839 more rows
```

pivot_annotations *Converts annotation tables to a wide format*

Description

Use this method to reconstitute the annotation tables into the format of the original resources. With the ‘wide=TRUE’ option [import_omnipath_annotations](#) applies this function to the downloaded data.

Usage

```
pivot_annotations(annotations)
```

Arguments

annotations A data frame of annotations downloaded from the OmniPath web service by [import_omnipath_annotations](#).

Value

A wide format data frame (tibble) if the provided data contains annotations from one resource, otherwise a list of wide format tibbles.

See Also

[import_omnipath_annotations](#)

Examples

```
# single resource: the result is a data frame
disgenet <- import_omnipath_annotations(resources = 'DisGeNet')
disgenet <- pivot_annotations(disgenet)
disgenet
# # A tibble: 126,588 × 11
#   uniprot genesymbol entity_type disease      type  score   dsi   dpi
#   <chr>    <chr>     <chr>       <chr>      <chr> <dbl> <dbl> <dbl>
# 1 P04217  A1BG      protein     Schizophren. dise.  0.3  0.7  0.538
# 2 P04217  A1BG      protein     Hepatomegaly phen.  0.3  0.7  0.538
# 3 P01023  A2M       protein     Fibrosis, L. dise.  0.3  0.529 0.769
# 4 P01023  A2M       protein     Acute kidne. dise.  0.3  0.529 0.769
# 5 P01023  A2M       protein     Mental Depr. dise.  0.3  0.529 0.769
# # . with 126,583 more rows, and 3 more variables: nof_pmids <dbl>,
# #   nof_snps <dbl>, source <chr>

# multiple resources: the result is a list
annotations <- import_omnipath_annotations(
```

```

resources = c('DisGeNet', 'SignaLink_function', 'DGIdb', 'kinase.com')
)
annotations <- pivot_annotations(annotations)
names(annotations)
# [1] "DGIdb"           "DisGeNet"          "kinase.com"
# [4] "SignaLink_function"
annotations$kinase.com
# # A tibble: 825 x 6
#   uniprot genesymbol entity_type group family subfamily
#   <chr>    <chr>      <chr>     <chr> <chr>   <chr>
# 1 P31749  AKT1       protein    AGC   Akt    NA
# 2 P31751  AKT2       protein    AGC   Akt    NA
# 3 Q9Y243  AKT3       protein    AGC   Akt    NA
# 4 O14578  CIT        protein    AGC   DMPK   CRIK
# 5 Q09013  DMPK       protein    AGC   DMPK   GEK
# # . with 815 more rows

```

Description

Retrieves predicted protein-protein interactions from the PrePPI database (<http://honig.c2b2.columbia.edu/preppi>). The interactions in this table are supposed to be correct with a > 0.5 probability.

Usage

```
preppi_download(...)
```

Arguments

... Minimum values for the scores. The available scores are: str, protpep, str_max, red, ort, phy, coexp, go, total, exp and final. Furthermore, an operator can be passed, either `.op = '&'` or `.op = '|'`, which is then used for combined filtering by multiple scores.

Details

PrePPI is a combination of many prediction methods, each resulting a score. For an explanation of the scores see <https://honiglab.c2b2.columbia.edu/hfpd/help/Manual.html>. The minimum, median and maximum values of the scores:

| Score | Minimum | Median | Maximum |
|---------|---------|--------|---------|
| str | 0 | 5.5 | 6,495 |
| protpep | 0 | 3.53 | 38,138 |
| str_max | 0 | 17.9 | 38,138 |

| | | | |
|-------|-----|-------|-----------------|
| red | 0 | 1.25 | 24.4 |
| ort | 0 | 0 | 5,000 |
| phy | 0 | 2.42 | 2.42 |
| coexp | 0 | 2.77 | 45.3 |
| go | 0 | 5.86 | 181 |
| total | 0 | 1,292 | 106,197,000,000 |
| exp | 1 | 958 | 4,626 |
| final | 600 | 1,778 | 4.91e14 |

Value

A data frame (tibble) of interactions with scores, databases and literature references.

See Also

[preppi_filter](#)

Examples

```
preppi <- preppi_download()
preppi
# # A tibble: 1,545,710 x 15
#   prot1 prot2 str_score protpep_score str_max_score red_score ort_score
#   <chr> <chr>    <dbl>        <dbl>      <dbl>    <dbl>    <dbl>
# 1 Q131. P146.    18.6       6.45     18.6     4.25    0.615
# 2 P064. Q96N.    1.83      14.3     14.3     4.25     0
# 3 Q7Z6. Q8NC.    4.57       0       4.57     0       0
# 4 P370. P154.    485.       0       485.     1.77    0.615
# 5 0004. Q9NR.    34.0       0       34.0     0.512    0
# # . with 1,545,700 more rows, and 8 more variables: phy_score <dbl>,
# #   coexp_score <dbl>, go_score <dbl>, total_score <dbl>, dbs <chr>,
# #   pubs <chr>, exp_score <dbl>, final_score <dbl>
```

Description

Filter PrePPI interactions by scores

Usage

```
preppi_filter(data, ..., .op = "&")
```

Arguments

| | |
|-------------------|---|
| <code>data</code> | A data frame of PrePPI interactions as provided by preppi_download . |
| <code>...</code> | Minimum values for the scores. The available scores are: str, protpep, str_max, red, ort, phy, coexp, go, total, exp and final. See more about the scores at preppi_download . |
| <code>.op</code> | The operator to combine the scores with: either '&' or ' '. With the former, only records where all scores are above the threshold will be kept; with the latter, records where at least one score is above its threshold will be kept. |

Value

The input data frame (tibble) filtered by the score thresholds.

See Also

[preppi_download](#)

Examples

```
preppi <- preppi_download()
preppi_filtered <- preppi_filter(preppi, red = 10, str = 4.5, ort = 1)
nrow(preppi_filtered)
# [1] 8443
```

`print_bma_motif_es` *Prints BMA motifs to the screen from a sequence of edges*

Description

The motifs can be copy-pasted into a BMA canvas.

Usage

```
print_bma_motif_es(edge_seq, G, granularity = 2)
```

Arguments

| | |
|--------------------------|-----------------------------|
| <code>edge_seq</code> | An igraph edge sequence. |
| <code>G</code> | An igraph graph object. |
| <code>granularity</code> | Numeric: granularity value. |

Value

Returns ‘NULL’.

Examples

```

interactions <- import_omnipath_interactions(resources = 'ARN')
graph <- interaction_graph(interactions)
print_bma_motif_es(igraph::E(graph)[1], graph)
# {"Model": {
#   "Name": "Omnipath motif",
#   "Variables": [
#     {"Name": "ULK1",
#      "Id": 1,
#      "RangeFrom": 0,
#      "RangeTo": 2,
#      "Formula": ""},
#     ...
#   ],
#   ...
# },
# ... (truncated)
# }}}
```

`print_bma_motif_vs` *Prints BMA motifs to the screen from a sequence of nodes*

Description

The motifs can be copy-pasted into a BMA canvas.

Usage

```
print_bma_motif_vs(node_seq, G)
```

Arguments

| | |
|----------|--------------------------|
| node_seq | An igraph node sequence. |
| G | An igraph graph object. |

Value

Returns ‘NULL’.

Examples

```

interactions <- import_omnipath_interactions(resources = 'ARN')
graph <- interaction_graph(interactions)
print_bma_motif_vs(
  igraph::all_shortest_paths(
    graph,
    from = 'ULK1',
```

```

        to = 'ATG13'
    )$res,
graph
)

```

print_interactions *Print OmniPath interactions*

Description

Prints the interactions or enzyme-substrate relationships in a nice format.

Usage

```
print_interactions(interDF, writeRefs = FALSE)
```

Arguments

- | | |
|-----------|--|
| interDF | data.frame with the interactions generated by any of the following functions: |
| | <ul style="list-style-type: none"> • import_omnipath_enzsub • import_omnipath_interactions • import_pathwayextra_interactions • import_kinaseextra_interactions • import_ligrecextra_interactions • import_post_translational_interactions • import_dorothea_interactions • import_tf_target_interactions • import_transcriptional_interactions • import_mirnatarget_interactions • import_all_interactions |
| writeRefs | [FALSE] writes also the PubMed IDs if available |

Value

Returns ‘NULL’.

Examples

```

enzsub <- import_omnipath_enzsub()
print_interactions(head(enzsub))
print_interactions(tail(enzsub), writeRefs = TRUE)
print_interactions(
  dplyr::filter(
    enzsub,
    enzyme_genesymbol == 'MAP2K1',

```

```

        substrate_genesymbol == 'MAPK3'
    )
)

signor <- import_omnipath_interactions(resources = 'SIGNOR')
print_interactions(head(signor))
#   source interaction      target n_resources
# 6 MAPK14 (Q16539) ==(+)==> MAPKAPK2 (P49137)      23
# 4 TRPM7 (Q96QT4) ==(+)==> ANXA1 (P04083)      10
# 1 PRKG1 (Q13976) ==(-)==> TRPC3 (Q13507)       8
# 2 PTPN1 (P18031) ==(-)==> TRPV6 (Q9H1D0)       6
# 5 PRKACA (P17612) ==(-)==> MCOLN1 (Q9GZU1)       6
# 3 RACK1 (P63244) ==(-)==> TRPM6 (Q9BX84)        2

```

print_path_es*Prints network paths in an edge sequence***Description**

Pretty prints the interactions in a path.

Usage

```
print_path_es(edgeSeq, G)
```

Arguments

| | |
|---------|--|
| edgeSeq | edge sequence |
| G | igraph object (from ptms or any interaction dataset) |

Value

Returns ‘NULL’.

See Also

- [print_path_vs](#)

Examples

```

interactions <- import_omnipath_interactions(resources = c('SignalLink3'))
OPI_g <- interaction_graph(interactions = interactions)
print_path_es(
  suppressWarnings(igraph::shortest_paths(
    OPI_g,
    from = 'TYRO3',
    to = 'STAT3',
    output = 'epath'
  ))$epath[[1]],

```

```
    OPI_g
)
```

print_path_vs*Print networks paths given by node sequence***Description**

Prints the interactions in the path in a nice format.

Usage

```
print_path_vs(nodeSeq, G)
```

Arguments

| | |
|---------|---|
| nodeSeq | node sequence |
| G | igraph object (from ptms or interactions) |

Value

Returns ‘NULL’.

See Also

[print_path_es](#)

Examples

```
interactions <- import_omnipath_interactions(resources=c('SignaLink3'))
OPI_g <- interaction_graph(interactions = interactions)
print_path_vs(
  igraph::all_shortest_paths(
    OPI_g,
    from = 'TYRO3',
    to = 'STAT3'
  )$vpath,
  OPI_g
)
enzsub <- import_omnipath_enzsub(resources=c('PhosphoSite', 'SIGNOR'))
enzsub_g <- enzsub_graph(enzsub)
print_path_vs(
  igraph::all_shortest_paths(
    enzsub_g,
    from = 'SRC',
    to = 'STAT1'
  )$res,
  enzsub_g
)
```

ramilowski_download *Downloads ligand-receptor interactions from Ramilowski et al. 2015*

Description

Curated ligand-receptor pairs from Supplementary Table 2 of the article "A draft network of ligand-receptor mediated multicellular signaling in human" (<https://www.nature.com/articles/ncomms8866>).

Usage

```
ramilowski_download()
```

Value

A data frame (tibble) with interactions.

Examples

```
rami_interactions <- ramilowski_download()
rami_interactions
# # A tibble: 2,557 x 16
#   Pair.Name Ligand.Approved. Ligand.Name Receptor.Approv.
#   <chr>      <chr>          <chr>      <chr>
# 1 A2M_LRP1  A2M            alpha-2-ma. LRP1
# 2 AANAT_MT. AANAT          aralkylami. MTNR1A
# 3 AANAT_MT. AANAT          aralkylami. MTNR1B
# 4 ACE_AGTR2 ACE           angiotensi. AGTR2
# 5 ACE_BDKR. ACE           angiotensi. BDKRB2
# # . with 2,547 more rows, and 12 more variables: Receptor.Name <chr>,
# #   DLRP <chr>, HPMR <chr>, IUPHAR <chr>, HPRD <chr>,
# #   STRING.binding <chr>, STRING.experiment <chr>, HPMR.Ligand <chr>,
# #   HPMR.Receptor <chr>, PMID.Manual <chr>, Pair.Source <chr>,
# #   Pair.Evidence <chr>
```

regnetwork_directions *Transcription factor effects from RegNetwork*

Description

Transcription factor effects from RegNetwork

Usage

```
regnetwork_directions(organism = "human")
```

Arguments

`organism` Character: either human or mouse.

Value

A data frame (tibble) of TF-target interactions with effect signs.

Examples

```
regn_dir <- regnetwork_directions()
regn_dir
# # A tibble: 3,954 x 5
#   source_genesymb. source_entrez target_genesymb. target_entrez
#   <chr>           <chr>        <chr>           <chr>
# 1 AHR             196          CDKN1B          1027
# 2 APLNR           187          PIK3C3          5289
# 3 APLNR           187          PIK3R4          30849
# 4 AR              367          KLK3            354
# 5 ARNT            405          ALDOA           226
# # . with 3,944 more rows, and 1 more variable: effect <dbl>
```

Description

Downloads transcriptional and post-transcriptional regulatory interactions from the RegNetwork database (<http://www.regnetworkweb.org/>). The information about effect signs (stimulation or inhibition), provided by `regnetwork_directions` are included in the result.

Usage

```
regnetwork_download(organism = "human")
```

Arguments

`organism` Character: either human or mouse.

Value

Data frame with interactions.

Examples

```
regn_interactions <- regnetwork_download()
regn_interactions
# # A tibble: 372,778 x 7
#   source_genesymb. source_entrez target_genesymb. target_entrez
#   <chr>           <chr>          <chr>           <chr>
# 1 USF1            7391           S100A6          6277
# 2 USF1            7391           DUSP1           1843
# 3 USF1            7391           C4A             720
# 4 USF1            7391           ABCA1            19
# 5 TP53            7157           TP73            7161
# # . with 372,768 more rows, and 3 more variables: effect <dbl>,
# #   source_type <chr>, target_type <chr>
```

`relations_list_to_table`

Table from a nested list of ontology relations

Description

Converting the nested list to a table is a more costly operation, it takes a few seconds. Best to do it only once, or pass `tables = TRUE` to [obo_parser](#), and convert the data frame to list, if you also need it in list format.

Usage

```
relations_list_to_table(relations, direction = NULL)
```

Arguments

- | | |
|------------------------|--|
| <code>relations</code> | A nested list of ontology relations (the "relations" element of the list returned by obo_parser in case its argument 'tables' is FALSE). |
| <code>direction</code> | Override the direction (i.e. child -> parents or parent -> children). The nested lists produced by functions in the current package add an attribute "direction" thus no need to pass this value. If the attribute and the argument are both missing, the column will be named simply "side2" and it won't be clear whether the relations point from "term" to "side2" or the other way around. The direction should be a character vector of length 2 with the values "parents" and "children". |

Value

The relations converted to a data frame (tibble).

See Also

- [swap_relations](#)
- [relations_table_to_list](#)
- [obo_parser](#)

Examples

```
goslim_url <-
  "http://current.geneontology.org/ontology/subsets/goslim_generic.obo"
path <- tempfile()
httr::GET(goslim_url, httr::write_disk(path, overwrite = TRUE))
obo <- obo_parser(path, tables = FALSE)
unlink(path)
rel_tbl <- relations_list_to_table(obos$relations)
```

relations_table_to_graph

Graph from a table of ontology relations

Description

Graph from a table of ontology relations

Usage

```
relations_table_to_graph(relations)
```

Arguments

| | |
|-----------|--|
| relations | A data frame of ontology relations (the "relations" element of the list returned by obo_parser in case its argument 'tables' is TRUE). |
|-----------|--|

Details

By default the relations point from child to parents, the edges in the graph will be of the same direction. Use [swap_relations](#) on the data frame to reverse the direction.

Value

The relations converted to an igraph graph object.

Examples

```
go <- get_db('go_basic')
go_graph <- relations_table_to_graph(go$relations)
```

relations_table_to_list

Nested list from a table of ontology relations

Description

Nested list from a table of ontology relations

Usage

```
relations_table_to_list(relations)
```

Arguments

relations A data frame of ontology relations (the "relations" element of the list returned by [obo_parser](#) in case its argument 'tables' is TRUE).

Value

The relations converted to a nested list.

See Also

- [relations_list_to_table](#)
- [swap_relations](#)
- [obo_parser](#)

Examples

```
goslim_url <-
  "http://current.geneontology.org/ontology/subsets/goslim_generic.obo"
path <- tempfile()
httr::GET(goslim_url, httr::write_disk(path, overwrite = TRUE))
obo <- obo_parser(path, tables = TRUE)
unlink(path)
rel_list <- relations_table_to_list(obos$relations)
```

remap_dorothea_download*Downloads TF-target interactions from ReMap***Description**

ReMap (<http://remap.univ-amu.fr/>) is a database of ChIP-Seq experiments. It provides raw and merged peaks and CRMs (cis regulatory motifs) with their associations to regulators (TFs). TF-target relationships can be derived as it is written in Garcia-Alonso et al. 2019: "For ChIP-seq, we downloaded the binding peaks from ReMap and scored the interactions between each TF and each gene according to the distance between the TFBSS and the genes' transcription start sites. We evaluated different filtering strategies that consisted of selecting only the top-scoring 100, 200, 500, and 1000 target genes for each TF." (<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC6673718/#s1title>). This function returns the top TF-target relationships as used in DoRothEA: https://github.com/saezlab/dorothea/blob/master/inst/scripts/02_chip_seq.R).

Usage

```
remap_dorothea_download()
```

Value

Data frame with TF-target relationships.

See Also

[remap_tf_target_download](#)

Examples

```
remap_interactions <- remap_dorothea_download()
remap_interactions
# # A tibble: 136,988 x 2
#   tf      target
#   <chr>  <chr>
# 1 ADNP    ABCC1
# 2 ADNP    ABCC6
# 3 ADNP    ABHD5
# 4 ADNP    ABT1
# 5 ADNP    AC002066.1
# # . with 136,978 more rows
```

| | |
|----------------|--|
| remap_filtered | <i>Downloads TF-target interactions from ReMap</i> |
|----------------|--|

Description

Downloads the ReMap TF-target interactions as processed by Garcia-Alonso et al. (<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC6673718/#s1title>) and filters them based on a score threshold, the top targets and whether the TF is included in the TF census (Vaquerizas et al. 2009). The code for filtering is adapted from DoRothEA, written by Christian Holland.

Usage

```
remap_filtered(score = 100, top_targets = 500, only_known_tfs = TRUE)
```

Arguments

- | | |
|----------------|--|
| score | Numeric: a minimum score between 0 and 1000, records with lower scores will be excluded. If NULL no filtering performed. |
| top_targets | Numeric: the number of top scoring targets for each TF. Essentially the maximum number of targets per TF. If NULL the number of targets is not restricted. |
| only_known_tfs | Logical: whether to exclude TFs which are not in TF census. |

Value

Data frame with TF-target relationships.

See Also

- [remap_tf_target_download](#)
- [remap_filtered](#)
- [tfcensus_download](#)

Examples

```
remap_interactions <- remap_filtered()
nrow(remap_interactions)
# [1] 145680

remap_interactions <- remap_filtered(top_targets = 100)
remap_interactions
# # A tibble: 30,330 x 2
#   source_genesymbol target_genesymbol
#   <chr>              <chr>
# 1 ADNP               ABCC1
# 2 ADNP               ABT1
# 3 ADNP               AC006076.1
# 4 ADNP               AC007792.1
```

```
# 5 ADNP          AC011288.2
# # . with 30,320 more rows
```

remap_tf_target_download*Downloads TF-target interactions from ReMap***Description**

ReMap (<http://remap.univ-amu.fr/>) is a database of ChIP-Seq experiments. It provides raw and merged peaks and CRMs (cis regulatory motifs) with their associations to regulators (TFs). TF-target relationships can be derived as it is written in Garcia-Alonso et al. 2019: "For ChIP-seq, we downloaded the binding peaks from ReMap and scored the interactions between each TF and each gene according to the distance between the TFBSS and the genes' transcription start sites. We evaluated different filtering strategies that consisted of selecting only the top-scoring 100, 200, 500, and 1000 target genes for each TF." (<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC6673718/#s1title>). This function retrieves the full processed TF-target list from the data deposited in <https://zenodo.org/record/3713238>.

Usage

```
remap_tf_target_download()
```

Value

Data frame with TF-target relationships.

See Also

- [remap_dorothea_download](#)
- [remap_filtered](#)

Examples

```
remap_interactions <- remap_tf_target_download()
remap_interactions
# # A tibble: 9,546,470 x 4
#   source_genesymbol target_genesymbol target_ensembl      score
#   <chr>            <chr>           <chr>             <dbl>
# 1 ADNP              PTPRS           ENSG00000105426.16 1000
# 2 AFF4              PRKCH           ENSG0000027075.14 1000
# 3 AHR               CTNND2          ENSG00000169862.18 1000
# 4 AR                PDE4D           ENSG00000113448.18 1000
# 5 ARID1A            PLEC            ENSG00000178209.14 1000
# # . with 9,546,460 more rows
```

`resources_colname` *Name of the column with the resources*

Description

Unfortunately the column title is different across the various query types in the OmniPath web service, so we need to guess.

Usage

```
resources_colname(data)
```

Arguments

data A data frame downloaded by any `import_...` function in the current package.

Value

Character: the name of the column, if any of the column names matches.

Examples

```
co <- import_omnipath_complexes()  
resources_colname(co)  
# [1] "sources"
```

simplify_intercell_network

Description

The intercellular communication network data frames, created by `import_intercell_network`, are combinations of a network data frame with two copies of the intercell annotation data frames, all of them already having quite some columns. Here we keep only the names of the interacting pair, their intercellular communication roles, and the minimal information of the origin of both the interaction and the annotations. Optionally further columns can be selected.

Usage

```
simplify_intercell_network(network, ...)
```

Arguments

`network` An intercell network data frame, as provided by [import_intercell_network](#).
`...` Optional, further columns to select.

Value

An intercell network data frame with some columns removed.

See Also

- [import_intercell_network](#)
- [unique_intercell_network](#)
- [filter_intercell_network](#)

Examples

```
icn <- import_intercell_network()
icn_s <- simplify_intercell_network(icn)
```

swap_relations

Reverse the direction of ontology relations

Description

Reverse the direction of ontology relations

Usage

```
swap_relations(relations)
```

Arguments

- | | |
|------------------------|---|
| <code>relations</code> | The ‘relations’ component of the data returned by obo_parser or any ‘...ontology_download’ function such as go_ontology_download . Depending on the <code>tables</code> argument of those functions the ‘relations’ can be a data frame or a nested list. |
|------------------------|---|

Value

Same type as the input, but the relations swapped: if in the input these pointed from each child to the parents, in the output they point from each parent to their children, and vice versa.

See Also

- [relations_list_to_table](#)
- [relations_table_to_list](#)
- [obo_parser](#)

Examples

```
goslim_url <-
  "http://current.geneontology.org/ontology/subsets/goslim_generic.obo"
path <- tempfile()
httr::GET(goslim_url, httr::write_disk(path, overwrite = TRUE))
obo <- obo_parser(path)
unlink(path)
rel_swapped <- swap_relations(oboj$relations)
```

tfcensus_download *Downloads the list of transcription factors from TF census*

Description

Vaquerizas et al. published in 2009 a list of transcription factors. This function retrieves Supplementary Table 2 from the article (<http://www.nature.com/nrg/journal/v10/n4/index.html>).

Usage

```
tfcensus_download()
```

Value

A data frame (tibble) listing transcription factors.

Examples

```
tfcensus <- tfcensus_download()
tfcensus
# # A tibble: 1,987 x 7
#   Class `Ensembl ID` `IPI ID` `Interpro DBD` `Interpro DNA-b.
#   <chr> <chr>     <chr>     <chr>      <chr>
# 1 a     ENSG00000000. IPI0021. NA          IPR001289
# 2 a     ENSG00000000. IPI0004. IPR000047;IPR. NA
# 3 a     ENSG00000000. IPI0001. IPR001356;IPR. NA
# 4 a     ENSG00000000. IPI0029. IPR000910;IPR. NA
# 5 a     ENSG00000000. IPI0001. IPR007087;IPR. IPR006794
# # . with 1,977 more rows, and 2 more variables: `HGNC symbol` <chr>,
# # `Tissue-specificity` <chr>
```

| | |
|---------------|---|
| translate_ids | <i>Translate gene and protein identifiers</i> |
|---------------|---|

Description

Translates a vector of identifiers, resulting a new vector, or a column of identifiers in a data frame by creating another column with the target identifiers.

Usage

```
translate_ids(
  d,
  ...,
  uploadlists = FALSE,
  keep_untranslated = TRUE,
  return_df = FALSE,
  organism = 9606,
  reviewed = TRUE
)
```

Arguments

| | |
|--------------------------|---|
| d | Character vector or data frame. |
| ... | At least two arguments, with or without names. The first of these arguments describes the source identifier, the rest of them describe the target identifier(s). The values of all these arguments must be valid identifier types as shown in Details. The names of the arguments are column names. In case of the first (source) ID the column must exist. For the rest of the IDs new columns will be created with the desired names. For ID types provided as arguments without names, the name of the ID type will be used for column name. |
| uploadlists | Force using the uploadlists service from UniProt. By default the plain query interface is used (implemented in uniprot_full_id_mapping_table in this package). If any of the provided ID types is only available in the uploadlists service, it will be automatically selected. The plain query interface is preferred because in the long term, with caching, it requires less download and data storage. |
| keep_untranslated | In case the output is a data frame, keep the records where the source identifier could not be translated. At these records the target identifier will be NA. |
| return_df | Return a data frame even if the input is a vector. |
| organism | Integer, NCBI Taxonomy ID of the organism (by default 9606 for human). Matters only if uploadlists is FALSE. |
| reviewed | Translate only reviewed (TRUE), only unreviewed (FALSE) or both (NULL) UniProt records. Matters only if uploadlists is FALSE. |

Details

This function, depending on the `uploadlists` parameter, uses either the uploadlists service of UniProt or plain UniProt queries to obtain identifier translation tables. The possible values for `from` and `to` are the identifier type abbreviations used in the UniProt API, please refer to the table here: https://www.uniprot.org/help/api_idmapping. In addition, simple synonyms are available which realize a uniform API for the uploadlists and UniProt query based backends. These are the followings:

| OmnipathR | Uploadlists | UniProt query |
|----------------|----------------------|--------------------|
| uniprot | ACC | id |
| uniprot_entry | ID | entry_name |
| genesymbol | GENENAME | genes(PREFERRED) |
| genesymbol_syn | | genes(ALTERNATIVE) |
| hgnc | HGNC_ID | database(HGNC) |
| entrez | P_ENTREZGENEID | database(geneid) |
| ensg | ENSEMBLGENOME_ID | |
| enst | ENSEMBL_TRS_ID | database(ensembl) |
| ensp | ENSEMBL_PRO_ID | |
| ensgt | ENSEMBLGENOME_TRS_ID | |
| ensgp | ENSEMBLGENOME_PRO_ID | |
| ensembl | ENSEMBL_ID | |
| protein_name | | protein_names |
| refseqp | P_REFSEQ_AC | database(refseq) |
| refseqn | REFSEQ_NT_ID | |
| embl | EMBL | database(embl) |
| embl_id | EMBL_ID | |
| gi | P_GI | |
| pir | PIR | |
| pdb | PDB_ID | |

The mapping between identifiers can be ambiguous. In this case one row in the original data frame yields multiple rows or elements in the returned data frame or vector(s).

Value

- Data frame: if the input is a data frame or the input is a vector and `return_df` is TRUE.
- Vector: if the input is a vector, there is only one target ID type and `return_df` is FALSE.
- List of vectors: if the input is a vector, there are more than one target ID types and `return_df` is FALSE. The names of the list will be ID types (as they were column names, see the description of the `...` argument), and the list will also include the source IDs.

See Also

- [uniprot_id_mapping_table](#)
- [uniprot_full_id_mapping_table](#)

Examples

```
d <- data.frame(uniprot_id = c('P00533', 'Q9ULV1', 'P43897', 'Q9Y2P5'))
d <- translate_ids(d, uniprot_id = uniprot, genesymbol)
d
#   uniprot_id genesymbol
# 1 P00533      EGFR
# 2 Q9ULV1      FZD4
# 3 P43897      TSFM
# 4 Q9Y2P5      SLC27A5
```

trrust_download *Downloads TF-target interactions from TRRUST*

Description

TRRUST v2 (<https://www.grnpedia.org/trrust/>) is a database of literature mined TF-target interactions for human and mouse.

Usage

```
ttrrust_download(organism = "human")
```

Arguments

organism Character: either "human" or "mouse".

Value

A data frame of TF-target interactions.

Examples

```
ttrrust_interactions <- ttrrust_download()
ttrrust_interactions
# # A tibble: 11,698 x 4
#   source_genesymbol target_genesymbol effect reference
#   <chr>            <chr>           <dbl> <chr>
# 1 AATF              BAX             -1    22909821
# 2 AATF              CDKN1A          0     17157788
# 3 AATF              KLK3            0     23146908
# 4 AATF              MYC             1     20549547
# 5 AATF              TP53            0     17157788
# 6 ABL1              BAX             1     11753601
# 7 ABL1              BCL2            -1    11753601
# # . with 11,688 more rows
```

uniprot_full_id_mapping_table

Creates an ID translation table from UniProt data

Description

Creates an ID translation table from UniProt data

Usage

```
uniprot_full_id_mapping_table(  
  to,  
  from = "id",  
  reviewed = TRUE,  
  organism = 9606  
)
```

Arguments

| | |
|-----------------------|---|
| <code>to</code> | Character or symbol: target ID type. See Details for possible values. |
| <code>from</code> | Character or symbol: source ID type. See Details for possible values. |
| <code>reviewed</code> | Translate only reviewed (TRUE), only unreviewed (FALSE) or both (NULL) UniProt records. |
| <code>organism</code> | Integer, NCBI Taxonomy ID of the organism (by default 9606 for human). |

Details

For both source and target ID type, this function accepts column codes used by UniProt and some simple shortcuts defined here. For the UniProt codes please refer to <https://www.uniprot.org/help/uniprotkb>. The shortcuts are entrez, genesymbol, genesymbol_syn (synonym gene symbols), hgnc, embl, ref-seq (RefSeq protein), enst (Ensembl transcript), uniprot_entry (UniProtKB AC, e.g. EGFR_HUMAN), protein_name (full name of the protein), uniprot (UniProtKB ID, e.g. P00533). For a complete table please refer to [translate_ids](#).

Value

A data frame (tibble) with columns ‘From’ and ‘To’, UniProt IDs and the corresponding foreign IDs, respectively.

See Also

[translate_ids](#)

Examples

```
uniprot_entrez <- uniprot_full_id_mapping_table(to = 'entrez')
uniprot_entrez
# # A tibble: 20,723 x 2
#   From      To
#   <chr>    <chr>
# 1 Q96R72  NA
# 2 Q9UKL2  23538
# 3 Q9H205  144125
# 4 Q8NGN2  219873
# 5 Q8NGC1  390439
# # . with 20,713 more rows
```

uniprot_id_mapping_table

Retrieves an identifier translation table from the UniProt uploadlists service

Description

Retrieves an identifier translation table from the UniProt uploadlists service

Usage

```
uniprot_id_mapping_table(identifiers, from, to, chunk_size = 5000)
```

Arguments

| | |
|-------------|---|
| identifiers | Character vector of identifiers |
| from | Character or symbol: type of the identifiers provided. See Details for possible values. |
| to | Character or symbol: identifier type to be retrieved from UniProt. See Details for possible values. |
| chunk_size | Integer: query the identifiers in chunks of this size. If you are experiencing download failures, try lower values. |

Details

This function uses the uploadlists service of UniProt to obtain identifier translation tables. The possible values for ‘from’ and ‘to’ are the identifier type abbreviations used in the UniProt API, please refer to the table here: https://www.uniprot.org/help/api_idmapping or the table of synonyms supported by the current package: [translate_ids](#). Note: if the number of identifiers is larger than the chunk size the log message about the cache origin is not guaranteed to be correct (most of the times it is still correct).

Value

A data frame (tibble) with columns ‘From’ and ‘To’, the identifiers provided and the corresponding target IDs, respectively.

See Also

[translate_ids](#)

Examples

```
uniprot_genesymbol <- uniprot_id_mapping_table(  
  c('P00533', 'P23771'), uniprot, genesymbol  
)  
uniprot_genesymbol  
# # A tibble: 2 x 2  
#   From     To  
#   <chr>    <chr>  
# 1 P00533  EGFR  
# 2 P23771  GATA3
```

unique_intercell_network

Unique intercellular interactions

Description

In the intercellular network data frames produced by [import_intercell_network](#), by default each pair of annotations for an interaction is represented in a separate row. This function drops the annotations and keeps only the distinct interacting pairs.

Usage

```
unique_intercell_network(network, ...)
```

Arguments

`network` An intercellular network data frame as produced by [import_intercell_network](#).

`...` Additional columns to keep. Note: if these have multiple values for an interacting pair, only the first row will be preserved.

Value

A data frame with interacting pairs and interaction attributes.

See Also

- `import_intercell_network`
- `simplify_intercell_network`
- `filter_intercell_network`

Examples

```
icn <- import_intercell_network()
icn_unique <- unique_intercell_network(icn)
```

vinayagam_download *Protein-protein interactions from Vinayagam 2011*

Description

Retrieves the Supplementary Table S6 from Vinayagam et al. 2011. Find out more at <https://doi.org/10.1126/scisignal.2001699>.

Usage

```
vinayagam_download()
```

Value

A data frame (tibble) with interactions.

Examples

```
vinayagam_interactions <- vinayagam_download()
vinayagam_interactions
# # A tibble: 34,814 x 5
#   `Input-node` `Gen.` `Output-node` `Ge.` `Output-node` `Ge.
#   <chr>        <dbl> <chr>          <dbl>
# 1 C1orf103    55791 MNAT1           4331
# 2 MAST2       23139 DYNLL1          8655
# 3 RAB22A      57403 APPL2          55198
# 4 TRAP1       10131 EXT2            2132
# 5 STAT2       6773  COPS4           51138
# # . with 34,804 more rows, and 1 more variable:
# # `Edge direction score` <dbl>
```

| | |
|---------------------------------|--|
| <code>walk_ontology_tree</code> | <i>All nodes of a subtree starting from the selected nodes</i> |
|---------------------------------|--|

Description

Starting from the selected nodes, recursively walks the ontology tree until it reaches either the root or leaf nodes. Collects all visited nodes.

Usage

```
walk_ontology_tree(
  terms,
  ancestors = TRUE,
  db_key = "go_basic",
  ids = TRUE,
  method = "gra",
  relations = c("is_a", "part_of", "occurs_in", "regulates", "positively_regulates",
    "negatively_regulates")
)
```

Arguments

| | |
|------------------------|--|
| <code>terms</code> | Character vector of ontology term IDs or names. A mixture of IDs and names can be provided. |
| <code>ancestors</code> | Logical: if FALSE the ontology tree is traversed towards the leaf nodes; if TRUE, the tree is traversed until the root. The former returns the ancestors (parents), the latter the descendants (children). |
| <code>db_key</code> | Character: key to identify the ontology database. For the available keys see omnipath_show_db . |
| <code>ids</code> | Logical: whether to return IDs or term names. |
| <code>method</code> | Character: either "gra" or "lst". The implementation to use for traversing the ontology tree. The graph based implementation is faster than the list based, the latter will be removed in the future. |
| <code>relations</code> | Character vector of ontology relation types. Only these relations will be used. |

Details

Note: this function relies on the database manager, the first call might take long because of the database load process. Subsequent calls within a short period should be faster. See [get_ontology_db](#).

Value

Character vector of ontology IDs. If the input terms are all leaves or roots NULL is returned. The starting nodes won't be included in the result unless they fall onto the traversal path from other nodes.

See Also

- [omnipath_show_db](#)
- [get_ontology_db](#)

Examples

```
walk_ontology_tree(c('GO:0006241', 'GO:0044211'))
# [1] "GO:0006139" "GO:0006220" "GO:0006221" "GO:0006241" "GO:0006725"
# [6] "GO:0006753" "GO:0006793" "GO:0006796" "GO:0006807" "GO:0008150"
# ... (truncated)
walk_ontology_tree(c('GO:0006241', 'GO:0044211'), ancestors = FALSE)
# [1] "GO:0044210" "GO:0044211"
walk_ontology_tree(
  c('GO:0006241', 'GO:0044211'),
  ancestors = FALSE,
  ids = FALSE
)
# [1] "'de novo' CTP biosynthetic process" "CTP salvage"
```

zenodo_download *Retrieves data from Zenodo*

Description

Zenodo is a repository of large scientific datasets. Many projects and publications make their datasets available at Zenodo. This function downloads an archive from Zenodo and extracts the requested file.

Usage

```
zenodo_download(
  path,
  reader = NULL,
  reader_param = list(),
  url_key = NULL,
  zenodo_record = NULL,
  zenodo_fname = NULL,
  url_param = list(),
  url_key_param = list(),
  ...
)
```

Arguments

- | | |
|--------|---|
| path | Character: path to the file within the archive. |
| reader | Optional, a function to read the connection. |

| | |
|---------------|---|
| reader_param | List: arguments for the reader function. |
| url_key | Character: name of the option containing the URL |
| zenodo_record | The Zenodo record ID, either integer or character. |
| zenodo_fname | The file name within the record. |
| url_param | List: variables to insert into the URL string (which is returned from the options). |
| url_key_param | List: variables to insert into the ‘url_key’. |
| ... | Passed to archive_extractor |

Value

A connection

Examples

```
# an example from the OmnipathR::remap_tf_target_download function:
remap_dorothea <- zenodo_download(
  zenodo_record = 3713238,
  zenodo_fname = 'tf_target_sources.zip',
  path = (
    'tf_target_sources/chip_seq/remap/gene_tf_pairs_genesymbol.txt'
  ),
  reader = read_tsv,
  reader_param = list(
    col_names = c(
      'source_genesymbol',
      'target_genesymbol',
      'target_ensembl',
      'score'
    ),
    col_types = cols(),
    progress = FALSE
  ),
  resource = 'ReMap'
)
```

Index

* datasets
 .omnipath_options_defaults, 6
.omnipath_options_defaults, 6

all_uniprots, 6
ancestors, 7
annotated_network, 8
annotation_categories, 9

bioplex1, 10, 11–13
bioplex2, 10, 10, 11–13
bioplex3, 10, 11, 11, 12, 13
bioplex_all, 10, 11, 12, 13
bioplex_hct116_1, 10–12, 13
bma_motif_es, 14
bma_motif_vs, 15

consensuspathdb_download, 15, 103
consensuspathdb_raw_table, 16

descendants, 17

enzsub_graph, 18, 23, 54
evex_download, 19, 81, 104

filter_by_resource, 20
filter_intercell_network, 20, 45, 46, 91,
 93, 96, 98, 156, 164
filter_sources(filter_by_resource), 20
find_all_paths, 18, 23, 66

get_annotation_databases, 52
get_annotation_databases
 (get_annotation_resources), 24
get_annotation_resources, 8, 24, 52
get_complex_genes, 25
get_complex_resources, 26
get_complexes_databases, 53
get_complexes_databases
 (get_complex_resources), 26
get_db, 26, 31, 36, 77

get_enzsub_resources, 27, 54
get_interaction_databases
 (get_interaction_resources), 28
get_interaction_resources, 28, 40–43,
 47–51, 55, 58–64
get_intercell_categories, 29, 29, 46,
 56–58
get_intercell_classes
 (get_intercell_generic_categories),
 29
get_intercell_generic_categories, 29,
 29, 46, 58
get_intercell_resources, 30, 56
get_ontology_db, 7, 17, 31, 165, 166
get_ptms_databases, 54
get_ptms_databases
 (get_enzsub_resources), 27
get_resources, 24, 26, 28, 30, 32
get_signed_ptms, 32
giant_component, 18, 23, 33, 66
go_annot_download, 34, 36
go_annot_slim, 34, 35
go_ontology_download, 36, 36, 156
guide2pharma_download, 37, 92

harmonizome_download, 38, 82
hpo_download, 39
htridb_download, 39, 82

import_all_interactions, 28, 40, 43, 47,
 48, 50, 51, 55, 59–61, 63, 64, 66, 144
import_AllInteractions
 (import_all_interactions), 40
import_dorothea_interactions, 28, 42, 66,
 144
import_intercell_network, 8, 20–22, 43,
 58, 93, 155, 156, 163, 164
import_KinaseExtra_Interactions
 (import_kinaseextra_interactions),
 46

import_kinaseextra_interactions, 28, 44,
 46, 46, 66, 144
import_LigrecExtra_Interactions
 (import_ligrecrextra_interactions),
 47
import_ligrecrextra_interactions, 28, 44,
 46, 47, 66, 144
import_lncrna_mrna_interactions, 49
import_miRNATarget_Interactions
 (import_mirnatarget_interactions),
 50
import_mirnatarget_interactions, 28, 50,
 66, 144
import_OmniPath_annotations
 (import_omnipath_annotations),
 51
import_Omnipath_annotations
 (import_omnipath_annotations),
 51
import_omnipath_annotations, 24, 51, 139
import_OmniPath_complexes
 (import_omnipath_complexes), 52
import_Omnipath_complexes
 (import_omnipath_complexes), 52
import_omnipath_complexes, 25, 26, 52
import_omnipath_enzsub, 18, 28, 33, 53, 66,
 144
import_OmniPath_Interactions
 (import_omnipath_interactions),
 54
import_Omnipath_Interactions
 (import_omnipath_interactions),
 54
import_omnipath_interactions, 8, 28, 33,
 43, 44, 46, 54, 54, 66, 144
import_OmniPath_intercell
 (import_omnipath_intercell), 56
import_Omnipath_intercell
 (import_omnipath_intercell), 56
import_omnipath_intercell, 29, 30, 43, 44,
 46, 56, 68
import_OmniPath_PTMS
 (import_omnipath_enzsub), 53
import_Omnipath_PTMS
 (import_omnipath_enzsub), 53
import_PathwayExtra_Interactions
 (import_pathwayextra_interactions),
 58
import_pathwayextra_interactions, 28,
 44, 46, 58, 66, 144
import_post_translational_interactions,
 59, 66, 106, 144
import_tf_mirna_interactions, 60
import_tf_target_interactions, 62, 66,
 144
import_TFregulons_Interactions
 (import_dorothea_interactions),
 42
import_tfregulons_interactions
 (import_dorothea_interactions),
 42
import_transcriptional_interactions,
 63, 66, 83, 144
inbiomap_download, 64, 65, 105
inbiomap_raw, 64, 65, 65
interaction_graph, 23, 41, 43, 47, 48, 50,
 51, 55, 59–61, 63, 64, 66
intercell_categories, 67
intercell_consensus_filter, 58, 67
is_ontology_id, 68
kegg_info, 69, 70, 74, 75
kegg_open, 69, 70, 74, 75
kegg_pathway_annotations, 72
kegg_pathway_download, 71, 73, 74, 76
kegg_pathway_list, 69–71, 73, 74, 75, 76
kegg_pathways_download, 71, 72–74, 76
kegg_picture, 69, 70, 74, 75
kegg_process, 71, 73, 74, 76
load_db, 77
nichenet_build_model, 78, 90
nichenet_expression_data, 78, 99, 100
nichenet_gr_network, 79, 81, 82, 84–86,
 96–98
nichenet_gr_network_evex, 79, 80, 80, 83
nichenet_gr_network_harmonizome, 79, 80,
 81, 83
nichenet_gr_network_htridb, 79, 80, 82,
 83
nichenet_gr_network_omnipath, 79, 80, 83,
 83
nichenet_gr_network_pathwaycommons, 80,
 83, 84
nichenet_gr_network_regnetwork, 79, 80,
 83, 84

nichenet_gr_network_remap, 79, 80, 83, 85
 nichenet_gr_network_trrust, 80, 83, 86
 nichenet_ligand_activities, 87
 nichenet_ligand_target_links, 88
 nichenet_ligand_target_matrix, 87, 89,
 90
 nichenet_lr_network, 87, 90, 91, 92–94,
 96–98, 100
 nichenet_lr_network_guide2pharma, 91,
 92
 nichenet_lr_network_omnipath, 83, 91, 93,
 93, 106
 nichenet_lr_network_ramilowski, 91, 94
 nichenet_main, 95, 108
 nichenet_networks, 78, 97, 97, 99
 nichenet_optimization, 78, 99
 nichenet_remove_orphan_ligands, 100
 nichenet_results_dir, 97, 101
 nichenet_signaling_network, 96–98, 101,
 103–106
 nichenet_signaling_network_cpdb, 102,
 103
 nichenet_signaling_network_evex, 102,
 104
 nichenet_signaling_network_harmonizome,
 102, 104
 nichenet_signaling_network_inbiomap,
 102, 105
 nichenet_signaling_network_omnipath,
 102, 106
 nichenet_signaling_network_pathwaycommons,
 102, 107
 nichenet_signaling_network_vinayagam,
 102, 107
 nichenet_test, 97, 108
 nichenet_workarounds, 97, 109

 obo_parser, 109, 149–151, 156
 omnipath_cache_autoclean, 112, 122
 omnipath_cache_clean, 112, 112, 122
 omnipath_cache_clean_db, 113
 omnipath_cache_download_ready, 114
 omnipath_cache_filter_versions, 115
 omnipath_cache_get, 116, 118
 omnipath_cache_key, 117
 omnipath_cache_latest_or_new, 117
 omnipath_cache_latest_version, 119
 omnipath_cache_load, 119
 omnipath_cache_move_in, 120, 123

 omnipath_cache_remove, 112, 121, 127
 omnipath_cache_save, 120, 121, 123
 omnipath_cache_search, 124
 omnipath_cache_set_ext, 125
 omnipath_cache_update_status, 126
 omnipath_cache_wipe, 122, 127
 omnipath_get_config_path, 127, 131
 omnipath_load_config, 128, 131
 omnipath_log, 129, 129
 omnipath_logfile, 129, 129
 omnipath_msg, 130
 omnipath_reset_config, 131
 omnipath_save_config, 131, 131
 omnipath_set_cachedir, 132
 omnipath_set_console_loglevel, 133, 134
 omnipath_set_logfile_loglevel, 133, 134
 omnipath_set_loglevel, 134
 omnipath_show_db, 7, 17, 27, 31, 77, 135,
 136–138, 165, 166
 omnipath_unlock_cache_db, 136
 OmnipathR, 111
 ontology_ensure_id, 136
 ontology_ensure_name, 137
 ontology_name_id, 137

 pathwaycommons_download, 84, 138
 pivot_annotations, 52, 139
 preppi_download, 140, 142
 preppi_filter, 141, 141
 print_bma_motif_es, 142
 print_bma_motif_vs, 143
 print_interactions, 41, 43, 47, 48, 50, 51,
 54, 55, 59–61, 63, 64, 144
 print_path_es, 145, 146
 print_path_vs, 145, 146
 printPath_es (print_path_es), 145
 printPath_vs (print_path_vs), 146
 ptms_graph (enzsub_graph), 18

 ramilowski_download, 94, 147
 regnetwork_directions, 147, 148
 regnetwork_download, 85, 148
 relations_list_to_table, 110, 149, 151,
 156
 relations_table_to_graph, 150
 relations_table_to_list, 110, 149, 151,
 156
 remap_dorothea_download, 152, 154
 remap_filtered, 86, 153, 153, 154

remap_tf_target_download, 152, 153, 154
resources_colname, 155

simplify_intercell_network, 22, 46, 155,
164
swap_relations, 110, 149–151, 156

tfcensus_download, 153, 157
translate_ids, 158, 161–163
trrust_download, 86, 160

uniprot_full_id_mapping_table, 158, 159,
161
uniprot_id_mapping_table, 159, 162
unique_intercell_network, 22, 45, 46, 156,
163

vinayagam_download, 164

walk_ontology_tree, 165

zenodo_download, 166