# Package 'fenr'

November 26, 2024

Title Fast functional enrichment for interactive applications

Version 1.5.0

**Description** Perform fast functional enrichment on feature lists (like genes or proteins) using the hypergeometric distribution. Tailored for speed, this package is ideal for interactive platforms such as Shiny. It supports the retrieval of functional data from sources like GO, KEGG, Reactome, Bioplanet and WikiPathways. By downloading and preparing data first, it allows for rapid successive tests on various feature selections without the need for repetitive, time-consuming preparatory steps typical of other packages.

URL https://github.com/bartongroup/fenr

BugReports https://github.com/bartongroup/fenr/issues

**Depends** R (>= 4.1.0)

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**Encoding** UTF-8

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Author Marek Gierlinski [aut, cre] (ORCID:

<https://orcid.org/0000-0001-9149-3514>)

Maintainer Marek Gierlinski < M. Gierlinski@dundee.ac.uk >

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fenr-package

fenr: Fast functional enrichment for interactive applications

# Description

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This R package provides a fast and efficient method for functional enrichment analysis, optimized for speed and designed for use in interactive applications, such as \*Shiny\* apps.

To learn more about fenr, start with the vignette: vignette("fenr").

#### Value

Package fenr.

## Author(s)

Maintainer: Marek Gierlinski < M. Gierlinski@dundee.ac.uk > (ORCID)

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#### See Also

Useful links:

- https://github.com/bartongroup/fenr
- Report bugs at https://github.com/bartongroup/fenr/issues

enrichment\_interactive

Small Shiny app serving as example for fast enrichment

## Description

Small Shiny app serving as example for fast enrichment

## Usage

```
enrichment_interactive(de, term_data)
```

# Arguments

de Differential expression results, yeast\_de attached to this package can be used.

term\_data A list of fenr\_terms objects containing functional data for various ontologies.

fetch\_terms\_for\_example can be used to create this object.

#### Value

An interactive Shiny app

```
## Not run:
data(yeast_de)
term_data <- fetch_terms_for_example(yeast_de)
    enrichment_interactive(yeast_de, term_data)
## End(Not run)</pre>
```

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exmpl\_all

Example set of background genes.

# Description

A set of gene names for proteins that could be detected in a typical proteomics experiment on yeast samples.

## Usage

```
data(exmpl_all)
```

#### **Format**

A character vector with 6985 elements.

## Value

A vector with background gene names.

exmpl\_sel

Example set of selected genes.

## Description

A set of gene names manually selected to illustrate functional enrichment.

## Usage

```
data(exmpl_sel)
```

#### **Format**

A character vector with 21 elements.

## Value

A vector with selected gene names.

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fetch\_bp

Get functional term data from BioPlanet

## Description

Download term information (term ID and name) and gene-pathway mapping (NCBI gene ID, gene symbol and pathway ID) from BioPlanet.

#### Usage

```
fetch_bp(use_cache = TRUE, on_error = c("stop", "warn", "ignore"))
```

## **Arguments**

use\_cache Logical, if TRUE, the remote file will be cached locally.

on\_error A character string indicating the error handling strategy: either "stop" to halt

execution, "warn" to issue a warning and return 'NULL' or "ignore" to return

'NULL' without warnings. Defaults to "stop".

#### Value

A list with terms and mapping tibbles.

#### **Examples**

```
bioplanet_data <- fetch_bp(on_error = "warn")</pre>
```

fetch\_go

Get Gene Ontology (GO) data

## Description

This function downloads term information (GO term ID and name) and gene-term mapping (gene ID, symbol, and GO term ID) from either the Ensembl database (using BioMart) or the Gene Ontology database (using GAF files), depending on the provided argument.

## Usage

```
fetch_go(
  species = NULL,
  dataset = NULL,
  use_cache = TRUE,
  on_error = c("stop", "warn", "ignore")
)
```

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#### **Arguments**

species (Optional) Species designation. Examples are goa\_human for human, mgi for mouse, or sgd for yeast. Full list of available species can be obtained using fetch\_go\_species - column designation. This argument is used when fetching data from the Gene Ontology database.

dataset (Optional) A string representing the dataset passed to Ensebml's Biomart, e.g. 'scerevisiae\_gene\_ensembl'. To see the different datasets available within a biomaRt you can e.g. do: mart <- biomaRt::useEnsembl(biomart = "ensembl"), followed by biomaRt::listDatasets(mart).

use\_cache Logical, if TRUE, the remote data will be cached locally.

A character string indicating the error handling strategy: either "stop" to halt execution, "warn" to issue a warning and return 'NULL' or "ignore" to return

'NULL' without warnings. Defaults to "stop".

#### **Details**

If species is provided, mapping from a Gene Ontology GAF file will be downloaded. GAF files contain more generic information than gene symbols. In this function, the third column of the GAF file (DB Object Symbol) is returned as gene\_symbol, but, depending on the species argument it can contain other entities, e.g. RNA or protein complex names. Similarly, the eleventh column of the GAF file (DB Object Synonym) is returned as gene\_id. It is up to the user to select the appropriate database.

Alternatively, if dataset is provided, mapping will be downloaded from Ensembl database. It will gene symbol and Ensembl gene ID.

#### Value

A list with terms and mapping tibbles.

#### **Examples**

```
# Fetch GO data from Ensembl
go_data_ensembl <- fetch_go(dataset = "scerevisiae_gene_ensembl", on_error = "warn")
# Fetch GO data from Gene Ontology
go_data_go <- fetch_go(species = "sgd", on_error = "warn")</pre>
```

fetch\_go\_species

Find all species available from geneontology.org

## **Description**

This function attempts to scrape HTML web page containing a table of available species and corresponding file names. If the structure of the page changes one day and the function stops working, go to <a href="http://current.geneontology.org/products/pages/downloads.html">http://current.geneontology.org/products/pages/downloads.html</a> and check file names. The species designation used in this package is the GAF file name without extension (e.g. for a file 'goa\_chicken.gaf' the designation is 'goa\_chicken').

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#### Usage

```
fetch_go_species(on_error = c("stop", "warn", "ignore"))
```

#### **Arguments**

on\_error

A character string indicating the error handling strategy: either "stop" to halt execution, "warn" to issue a warning and return 'NULL' or "ignore" to return 'NULL' without warnings. Defaults to "stop".

#### Value

A tibble with columns species and designation.

## **Examples**

```
go_species <- fetch_go_species(on_error = "warn")</pre>
```

fetch\_kegg

Get functional term data from KEGG

#### **Description**

Download information (pathway ID and name) and gene-pathway mapping (entrez gene ID, gene symbol and pathway ID) from KEGG. Gene symbols are extracted from gene descriptions. For some species (e.g. yeast), gene symbols are returned instead of entrez IDs and not in gene description.

## Usage

```
fetch_kegg(species, batch_size = 10, on_error = c("stop", "warn", "ignore"))
```

# Arguments

species	KEGG species code,	for example	"hsa" for human.	The full list of available
---------	--------------------	-------------	------------------	----------------------------

KEGG species can be found by using fetch\_kegg\_species.

batch\_size Number of pathways sent to KEGG database in one query. The maximum al-

lowed is 10.

on\_error A character string indicating the error handling strategy: either "stop" to halt

execution, "warn" to issue a warning and return 'NULL' or "ignore" to return

'NULL' without warnings. Defaults to "stop".

#### Value

A list with terms and mapping tibbles.

```
kegg_data <- fetch_kegg("mge", on_error = "warn")</pre>
```

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fetch\_kegg\_species

Find all species available from KEGG

## Description

Find all species available from KEGG

## Usage

```
fetch_kegg_species(on_error = c("stop", "warn", "ignore"))
```

# **Arguments**

on\_error

A character string indicating the error handling strategy: either "stop" to halt execution, "warn" to issue a warning and return 'NULL' or "ignore" to return 'NULL' without warnings. Defaults to "stop".

#### Value

A tibble, in which column designation contains species designations used in function fetch\_kegg.

# **Examples**

```
spe <- fetch_kegg_species(on_error = "warn")</pre>
```

fetch\_reactome

Get functional term data from Reactome

## **Description**

Download term information (pathway ID and name) and gene-pathway mapping (Ensembl gene ID or gene symbol and pathway ID) from Reactome.

## Usage

```
fetch_reactome(
   species,
   source = c("ensembl", "api", "gene_association"),
   use_cache = TRUE,
   on_error = c("stop", "warn", "ignore")
)
```

fetch\_reactome\_species

#### **Arguments**

species Reactome species designation, for example "Homo sapiens" for human. Full list

of available species can be found using fetch\_reactome\_species().

source How to download the mapping. If 'ensembl' or 'gene\_association', one mapping

file provided by Reactome will be downloaded, if 'api', then Reactome API will

be used. See details.

use\_cache Logical, if TRUE, the remote file will be cached locally.

on\_error A character string indicating the error handling strategy: either "stop" to halt

execution, "warn" to issue a warning and return 'NULL' or "ignore" to return

'NULL' without warnings. Defaults to "stop".

#### **Details**

Reactome makes mapping between Ensembl ID and pathway ID available in form of one down-loadable file. This mapping contains gene symbols as well. Also, a gene association file with mapping between UniProt accession number, gene symbol and Reactome term is available. If source = "ensembl" or source = "gene\_association" is set, one large file will be downloaded and parsed. If source = "api" is set, then Reactome APIs will be interrogated for each pathway available. This method is considerably slower, especially for large genomes. However, gene association file contains far fewer mappings than can be extracted using API. If gene symbols are needed, we recommend using source = "ensembl".

#### Value

A list with terms and mapping tibbles

#### **Examples**

```
reactome_data <- fetch_reactome("Saccharomyces cerevisiae", on_error = "warn")
```

fetch\_reactome\_species

List of available Reactome species

#### **Description**

List of available Reactome species

## Usage

```
fetch_reactome_species(on_error = c("stop", "warn", "ignore"))
```

#### **Arguments**

on\_error A character string indicating the error handling strategy: either "stop" to halt

execution, "warn" to issue a warning and return 'NULL' or "ignore" to return

'NULL' without warnings. Defaults to "stop".

## Value

A tibble with species names used by Reactome.

## **Examples**

```
re <- fetch_reactome_species(on_error = "warn")</pre>
```

```
fetch_terms_for_example
```

Create term data for interactive example

## Description

Create term data for interactive example

## Usage

```
fetch_terms_for_example(de)
```

## Arguments

de

Differential expression results, use yeast\_de data attached to this package.

## Value

A list of objects containing functional terms for GO and Reactome.

```
## Not run:
data(yeast_de)
term_data <- fetch_terms_for_example(yeast_de)
## End(Not run)</pre>
```

fetch\_wiki

fetch		
THICH	\\\/ I	ĸι

Get functional term data from WikiPathways

#### **Description**

Download term information (pathway ID and name) and gene-pathway mapping (gene symbol and pathway ID) from WikiPathways.

## Usage

```
fetch_wiki(
  species,
  databases = c("Ensembl", "Entrez Gene", "HGNC", "HGNC Accession number",
    "Uniprot-TrEMBL"),
  types = c("GeneProduct", "Protein", "Rna", "RNA"),
  on_error = c("stop", "warn", "ignore")
)
```

## **Arguments**

species	WikiPathways species designation, for example "Homo sapiens" for human. Full list of available species can be found using fetch_wiki_species().
databases	A character vector with database names to pre-filter mapping data. See details. Full result will be returned if NULL.
types	A character vector with types of entities to pre-filter mapping data. See details. Full result will be returned if NULL.
on_error	A character string indicating the error handling strategy: either "stop" to halt execution, "warn" to issue a warning and return 'NULL' or "ignore" to return 'NULL' without warnings. Defaults to "stop".

## **Details**

WikiPathways contain mapping between pathways and a variety of entities from various databases. Typically a gene symbol is returned in column text\_label and some sort of ID in column id, but this depends on the species and databases used. For gene/protein enrichment, these should be filtered to contain gene symbols only. This can be done by selecting a desired databases and types. The default values for parameters databases and types attempt to select information from generic databases, but there are organism-specific databases not included in the selection. We suggest to run this function with databases = NULL, types = NULL to see what types and databases are available before making selection.

#### Value

A list with terms and mapping tibbles.

functional\_enrichment

#### **Examples**

```
wiki_data <- fetch_wiki("Bacillus subtilis", on_error = "warn")</pre>
```

fetch\_wiki\_species

List of available WikiPathways species

## **Description**

List of available WikiPathways species

# Usage

```
fetch_wiki_species(on_error = c("stop", "warn", "ignore"))
```

## Arguments

on\_error

A character string indicating the error handling strategy: either "stop" to halt execution, "warn" to issue a warning and return 'NULL' or "ignore" to return 'NULL' without warnings. Defaults to "stop".

#### Value

A character vector with species names used by WikiPathways.

#### **Examples**

```
spec <- fetch_wiki_species(on_error = "warn")</pre>
```

functional\_enrichment Fast Functional Enrichment

# Description

Perform fast functional enrichment analysis based on the hypergeometric distribution. Designed for use in interactive applications.

## Usage

```
functional_enrichment(feat_all, feat_sel, term_data, feat2name = NULL)
```

## Arguments

feat_all	A character vector with all feature identifiers, serving as the background for enrichment.
feat_sel	A character vector with feature identifiers in the selection.
term_data	An object of class fenr_terms, created by prepare_for_enrichment.
feat2name	An optional named list to convert feature IDs into feature names.

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#### **Details**

This function carries out functional enrichment analysis on a selection of features (e.g., differentially expressed genes) using the hypergeometric probability distribution (Fisher's exact test). Features can be genes, proteins, etc. The term\_data object contains functional term information and feature-term mapping.

#### Value

A tibble with enrichment results, providing the following information for each term:

- N\_with number of features with this term among all features
- n\_with\_sel number of features with this term in the selection
- n\_expect expected number of features with this term in the selection, under the null hypothesis that terms are mapped to features randomly
- enrichment ratio of n\_with\_sel / n\_expect
- odds\_ratio odds ratio for enrichment; is infinite when all features with the given term are in the selection
- p\_value p-value from a single hypergeometric test
- p\_adjust p-value adjusted for multiple tests using the Benjamini-Hochberg approach

## **Examples**

```
## Not run:
data(exmpl_all, exmpl_sel)
go <- fetch_go(species = "sgd")
go_terms <- prepare_for_enrichment(go$terms, go$mapping, exmpl_all, feature_name = "gene_symbol")
enr <- functional_enrichment(exmpl_all, exmpl_sel, go_terms)
## End(Not run)</pre>
```

get\_feature\_terms

Get terms annotating a given feature.

#### **Description**

Get terms annotating a given feature.

## Usage

```
get_feature_terms(term_data, feature_id)
```

#### **Arguments**

```
term_data An object class fenr_terms, created by prepare_for_enrichment.
```

feature\_id A string with a feature ID

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#### Value

A character vector containing functional term IDs annotating given feature.

#### **Examples**

```
## Not run:
go_data <- fetch_go(species = "sgd")
go_terms <- prepare_for_enrichment(go_data$terms, go_data$mapping, feature = "gene_symbol")
trms <- get_feature_terms(go_terms, "GEM1")
## End(Not run)</pre>
```

get\_term\_features

Get features annotated with a given term.

# Description

Get features annotated with a given term.

#### Usage

```
get_term_features(term_data, term_id)
```

## **Arguments**

term\_data An object class fenr\_terms, created by prepare\_for\_enrichment.

term\_id A string with a functional term ID.

#### Value

A character vector containing feature IDs annotated with the term ID.

```
## Not run:
go_data <- fetch_go(species = "sgd")
go_terms <- prepare_for_enrichment(go_data$terms, go_data$mapping, feature = "gene_symbol")
feats <- get_term_features(go_terms, "GO:0000001")
## End(Not run)</pre>
```

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go

GO-terms data downloaded for the vignette.

# Description

```
Downloaded using go <- fetch_go(species = "sgd")
```

# Usage

data(go)

## **Format**

A list of two tibbles

## Value

Contains GO-term descriptions and gene mapping.

go\_species

GO species

# Description

Downloaded using go\_species <- fetch\_go\_species()

## Usage

```
data(go_species)
```

#### **Format**

A tibble

## Value

Contains species available through Gene Ontology

```
prepare_for_enrichment
```

Prepare Term Data for Enrichment Analysis

#### **Description**

Process term data downloaded with the fetch\_\* functions, preparing it for fast enrichment analysis using functional\_enrichment.

#### Usage

```
prepare_for_enrichment(
  terms,
  mapping,
  all_features = NULL,
  feature_name = "gene_id"
)
```

#### Arguments

terms A tibble with at least two columns: term\_id and term\_name. This tibble con-

tains information about functional term names and descriptions.

mapping A tibble with at least two columns, containing the mapping between functional

terms and features. One column must be named term\_id, while the other column should have a name specified by the feature\_name argument. For example, if mapping contains columns term\_id, accession\_number, and gene\_symbol, setting feature\_name = "gene\_symbol" indicates that gene symbols will be

used for enrichment analysis.

all\_features A vector with all feature IDs used as the background for enrichment. If not

specified, all features found in mapping will be used, resulting in a larger object

size.

feature\_name The name of the column in the mapping tibble to be used as the feature identi-

fier. For example, if mapping contains columns term\_id, accession\_number, and gene\_symbol, setting feature\_name = "gene\_symbol" indicates that gene

symbols will be used for enrichment analysis.

#### **Details**

This function takes two tibbles containing functional term information (terms) and feature mapping (mapping), and converts them into an object required by functional\_enrichment for efficient analysis. Terms and mapping can be generated with the database access functions included in this package, such as fetch\_reactome or fetch\_go\_from\_go.

#### Value

An object of class fenr\_terms required by functional\_enrichment.

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#### **Examples**

remove\_cache

Remove all cache

## **Description**

This function will remove all cached data used by 'fenr'. The user will be prompted for confirmation. Use only when you suspect the cache was corrupted. Use with caution!

# Usage

```
remove_cache(ask = TRUE)
```

## **Arguments**

ask

Logical, whether to ask user for confirmation.

#### Value

TRUE if successfully removed.

#### **Examples**

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

yeast\_de

Differential expression results for yeast RNA-seq.

## **Description**

A subset of 6 + 6 replicates was selected from data set reported in https://doi.org/10.1093/bioinformatics/btv425

#### Usage

```
data(yeast_de)
```

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## **Format**

A tibble with 5 columns

# Value

Results for differential expression for yeast RNA-seq.

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