

# Package ‘brendaDb’

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**Type** Package

**Title** The BRENDA Enzyme Database

**Version** 1.8.0

**Description** R interface for importing and analyzing enzyme information from the BRENDA database.

**License** MIT + file LICENSE

**Encoding** UTF-8

**biocViews** ThirdPartyClient, Annotation, DataImport

**URL** <https://github.com/y1zhou/brendaDb>

**BugReports** <https://github.com/y1zhou/brendaDb/issues>

**Suggests** testthat, BiocStyle, knitr, rmarkdown, devtools

**Imports** dplyr, Rcpp, tibble, stringr, magrittr, purrr, BiocParallel, crayon, utils, tidyr, curl, xml2, grDevices, rlang, BiocFileCache, rappdirs

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**Roxygen** list(markdown = TRUE)

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## R topics documented:

brendaDb-package	2
BiocycPathwayEnzymes	3
BiocycPathwayGenes	3
DownloadBrenda	4
ExtractField	5
ID2Enzyme	5
print.brenda.entries	6
print.brenda.entry	7
QueryBrenda	7
ReadBrenda	8
ReadBrendaFile	9
SeparateEntries	9
ShowFields	10
<b>Index</b>	<b>11</b>

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brendaDb-package	<i>brendaDb: the BRENDA enzyme database.</i>
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### Description

brendaDb provides an R interface to download, clean and extract enzyme information from the BRENDA database.

### Details

The main aims of brendaDb include:

- Read text file downloaded from BRENDA into an R tibble
- Retrieve information for specific enzymes
- Query enzymes using their synonyms, gene symbols, etc.
- Query enzyme information for specific **BioCyc** pathways

To learn more about brendaDb, please refer to the vignette. `browseVignettes(package = "brendaDb")`

### Author(s)

**Maintainer:** Yi Zhou <yi.zhou@uga.edu> ([ORCID](#))

### See Also

Useful links:

- <https://github.com/y1zhou/brendaDb>
- Report bugs at <https://github.com/y1zhou/brendaDb/issues>

---

BiocycPathwayEnzymes *Get all EC numbers involved in a BioCyc pathway.*

---

**Description**

Get all EC numbers involved in a BioCyc pathway.

**Usage**

```
BiocycPathwayEnzymes(org.id = "HUMAN", pathway, sleep = 0)
```

**Arguments**

org.id	The identifier for the organism database in BioCyc, e.g. ECOLI, HUMAN, META, AFER243159
pathway	A case-sensitive pathway object identifier, e.g. PWY66-400, LYSINE-DEG1-PWY.
sleep	Optional, default is 0. If set to a non-zero value, pause for the specified seconds after retrieving each reaction. Increase this if the default fails (probably limited on the BioCyc API side).

**Value**

If the pathway is found, returns a tibble with columns - RxnID: reaction IDs found in the pathway. - EC: EC number of the enzyme catalyzing the reaction. - ReactionDirection: direction of the reaction. - LHS: left-hand-side of the reaction. - RHS: right-hand-side of the reaction. Different compounds in the reactions are separated by " + " in columns LHS and RHS. The function returns NULL if the pathway ID is not found.

**Examples**

```
BiocycPathwayEnzymes("HUMAN", "PWY66-400")
```

---

BiocycPathwayGenes *Get all genes involved in a BioCyc pathway.*

---

**Description**

Get all genes involved in a BioCyc pathway.

**Usage**

```
BiocycPathwayGenes(org.id = "HUMAN", pathway)
```

**Arguments**

org.id	The identifier for the organism database in BioCyc, e.g. ECOLI, HUMAN, META, AFER243159
pathway	A case-sensitive pathway object identifier, e.g. PWY66-400, LYSINE-DEG1-PWY.

**Value**

If the pathway is found, returns a tibble with columns BiocycGene, BiocycProtein, Symbol and Ensembl, where BiocycGene and BiocycProtein are the gene and protein IDs in the BioCyc database, respectively. Returns NULL if the pathway ID is not found.

**Examples**

```
BiocycPathwayGenes("HUMAN", "PWY66-400")  
BiocycPathwayGenes("HUMAN", "TRYPTOPHAN-DEGRADATION-1")
```

---

DownloadBrenda

*Download and unzip the BRENDA text file.*

---

**Description**

By default, the function downloads a zipped BRENDA text file to a local cache directory, and extracts a brenda\_download.txt file.

**Usage**

```
DownloadBrenda(force.download = FALSE)
```

**Arguments**

force.download Boolean value. If TRUE, ignore the cache and force re-download of the BRENDA text file. Default is FALSE.

**Value**

A string of the path to the downloaded BRENDA text file.

**Examples**

```
## Not run: DownloadBrenda()
```

---

ExtractField	<i>Extract a specific field from a <code>brenda.entries</code> object.</i>
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---

### Description

Retrieve one field from all the `brenda.entry` objects. A column of EC numbers will be added to distinguish different enzymes.

### Usage

```
ExtractField(res, field, entries = NULL)
```

### Arguments

<code>res</code>	A <code>brenda.entries</code> object from <code>QueryBrenda()</code> .
<code>field</code>	A string indicating the field to extract. Nested fields should be separated by \$, e.g. <code>parameters\$ph.optimum</code> .
<code>entries</code>	A character vector with values of EC numbers. This should be a subset of <code>names(res)</code> .

### Value

A tibble with all columns from the tibble in the given field, and extra columns containing the EC numbers and organisms.

### Examples

```
df <- ReadBrenda(system.file("extdata", "brenda_download_test.txt",
                             package = "brendaDb"))
res <- QueryBrenda(brenda = df, EC = c("1.1.1.1", "6.3.5.8"),
                  n.core = 2)
ExtractField(res, field = "molecular$stability$general.stability")
ExtractField(res, field = "structure$subunits")
```

---

ID2Enzyme	<i>A helper function for converting names/synonyms to EC numbers.</i>
-----------	---

---

### Description

A helper function for converting names/synonyms to EC numbers.

### Usage

```
ID2Enzyme(brenda, ids)
```

**Arguments**

brenda	A tibble generated from <a href="#">ReadBrenda()</a> .
ids	A character vector of IDs to be converted.

**Details**

The function goes through "RECOMMENDED\_NAME", "SYSTEMATIC\_NAME", and "SYNONYMS" in the BRENDA file, and uses regexes to look for the given IDs. Values in the three columns are kept if the regex had a hit, otherwise NA is filled. The function can take in IDs of multiple sources, e.g. `c("ADH4", "CD38", "pyruvate dehydrogenase")`. Note that using aliases instead of symbols could lead to false positives in the output table.

**Value**

A tibble with columns ID, EC, and at least one of (RECOMMENDED\_NAME, SYSTEMATIC\_NAME and SYNONYMS).

**Examples**

```
df <- ReadBrenda(system.file("extdata", "brenda_download_test.txt",
                             package = "brendaDb"))
ID2Enzyme(df, c("CD38", "ADH4", "pyruvate dehydrogenase"))
```

---

`print.brenda.entries` *Show the number of regular and transferred/deleted brenda.entry objects in the brenda.entries list.*

---

**Description**

Show the number of regular and transferred/deleted brenda.entry objects in the brenda.entries list.

**Usage**

```
## S3 method for class 'brenda.entries'
print(x, ..., verbose = FALSE)
```

**Arguments**

x	A brenda.entries list returned by <a href="#">QueryBrenda()</a> .
...	Other arguments passed to the generic function.
verbose	Boolean; if TRUE, print tree views of brenda.query objects.

**Value**

Nothing; print summary information to the terminal.

---

```
print.brenda.entry
```

*Show the non-empty fields in the query result.*

---

### Description

For details, see [PrettyPrintBrendaEntry\(\)](#).

### Usage

```
## S3 method for class 'brenda.entry'
print(x, full.output = FALSE, ...)
```

### Arguments

x	A brenda.entry object (elements in the list returned by the function <a href="#">QueryBrenda()</a> ).
full.output	A boolean default to FALSE. If TRUE, include all entries even if they are empty (NA or 0 rows).
...	Other arguments passed to the generic function.

### Value

Nothing; print object information to the terminal.

---

```
QueryBrenda
```

*Query for multiple enzymes.*

---

### Description

Use a vector of EC numbers to retrieve information from the BRENDA tibble read in by [ReadBrenda\(\)](#). Invalid EC numbers will be removed and a message will be generated.

### Usage

```
QueryBrenda(brenda, EC, n.core = 0, fields = FALSE, ...)
```

### Arguments

brenda	A tibble containing information from BRENDA.
EC	A string of the EC number.
n.core	Integer specifying the number of cores to use. Default is 0, which would result in using all available cores.
fields	A character vector indicating fields to parse. Default is FALSE, which would be returning all fields.
...	Other parameters passed to <a href="#">QueryBrendaBase()</a> .

**Value**

A list of `brenda.entry` objects.

**See Also**

[QueryBrendaBase\(\)](#) [ConfigBPCores\(\)](#) [SelectOrganism\(\)](#)

**Examples**

```
df <- ReadBrenda(system.file("extdata", "brenda_download_test.txt",
                             package = "brendaDb"))
res <- QueryBrenda(brenda = df, EC = c("1.1.1.1", "1.1.1.10", "6.3.5.8"),
                  n.core = 2, organisms = "Homo sapiens")
```

---

ReadBrenda

*Read BRENDA text file into matrix.*

---

**Description**

For each EC entry, split the annotations into three columns:

- ID: EC number, e.g. 1.1.1.1
- field: the content of the information, e.g. protein, localization
- description: everything else

**Usage**

```
ReadBrenda(filepath, clean = TRUE)
```

**Arguments**

`filepath` A string indicating the path to the text file.  
`clean` Boolean; if TRUE, run [CleanECNumber\(\)](#) after reading the file.

**Value**

A matrix containing information about the EC entries.

**Examples**

```
brenda_txt <- system.file("extdata", "brenda_download_test.txt",
                          package = "brendaDb")
df <- ReadBrenda(brenda_txt)
```



---

ReadBrendaFile	<i>Read raw BRENDA text file.</i>
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---

**Description**

Read file into buffer, and load all non-empty lines. Comment lines (starting with \*) are skipped. The text file should be downloaded from [https://www.brenda-enzymes.org/download\\_brenda\\_without\\_registration.php](https://www.brenda-enzymes.org/download_brenda_without_registration.php)

**Usage**

```
ReadBrendaFile(filepath)
```

**Arguments**

filepath      A string indicating the path to the text file.

**Value**

A vector<string> with each element being a line in the file.

---

SeparateEntries	<i>Convert vector of lines to matrix.</i>
-----------------	---

---

**Description**

For each EC entry, split the annotations into three columns:

- ID: EC number, e.g. 1.1.1.1
- field: the content of the information, e.g. protein, localization
- description: everything else

**Usage**

```
SeparateEntries(lines)
```

**Arguments**

lines      The output vector<string> from read\_brenda\_file.

**Value**

A vector<vector<string>> containing information about the EC entries. In R this is a list of 3 lists.

---

`ShowFields`*Show all unique BRENDA fields and their corresponding acronyms.*

---

**Description**

Show all unique BRENDA fields and their corresponding acronyms.

**Usage**

```
ShowFields(df)
```

**Arguments**

`df` A data.frame with columns "field" and "description"

**Value**

A data.frame with columns "field" and "acronym".

**Examples**

```
df <- ReadBrenda(system.file("extdata", "brenda_download_test.txt",  
                             package = "brendaDb"))  
ShowFields(df)
```

# Index

BiocycPathwayEnzymes, [3](#)  
BiocycPathwayGenes, [3](#)  
brendaDb (brendaDb-package), [2](#)  
brendaDb-package, [2](#)

CleanECNumber(), [8](#)  
ConfigBPCores(), [8](#)

DownloadBrenda, [4](#)

ExtractField, [5](#)

ID2Enzyme, [5](#)

PrettyPrintBrendaEntry(), [7](#)  
print.brenda.entries, [6](#)  
print.brenda.entry, [7](#)

QueryBrenda, [7](#)  
QueryBrenda(), [5–7](#)  
QueryBrendaBase(), [7, 8](#)

ReadBrenda, [8](#)  
ReadBrenda(), [6, 7](#)  
ReadBrendaFile, [9](#)

SelectOrganism(), [8](#)  
SeparateEntries, [9](#)  
ShowFields, [10](#)