Package 'taxonbridge'

January 19, 2022

Title Combined analysis of NCBI taxonomy and GBIF taxonomy

Version 0.0.0.9000

Description The NCBI is the go to resource for taxonomic studies but it only contains data on extant species whereas the GBIF also includes extinct species. `taxonbridge` is useful for the combined analysis of the NCBI taxonomy and GBIF taxonomy.

R topics documented:

LazyData true

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get_lineages Get entries that have lineage information for bedata	both the GBIF and NCBI
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Description

Get entries that have lineage information for both the GBIF and NCBI data

Usage

```
get_lineages(x)
```

Arguments

Х

A tibble created with load_taxonomies() or load_population() or load_sample().

Value

A tibble with complete lineage data

Examples

```
get_lineages(load_sample())
```

get_status

Filter a combined taxonomy by GBIF taxonomic status/synonym

Description

Filter a combined taxonomy by GBIF taxonomic status/synonym

Usage

```
get_status(x, status = "all")
```

Arguments

A tibble created with load_taxonomies() or load_population() or load_sample().

status

Filter on GBIF assigned status (i.e. NA, "doubtful", "accepted", "proparte synonym", "synonym", "homotypic synonym", "heterotypic synonym"). Can be a

string or a vector of strings. Defaults to no filtering.

Value

A filtered tibble

Examples

```
get_status(load_sample(), "synonym")
get_status(load_sample(), c("accepted", "doubtful"))
```

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get_validity	Validate entries of a merged taxonomy	

Description

Validate entries of a merged taxonomy

Usage

```
get_validity(x, rank = "family", valid = TRUE)
```

Arguments

x A tibble created with load_taxonomies() or load_population() or load_sample().

rank A string with GBIF rank that will be used to examine an NCBI lineage for

validation purposes. Must be kingdom, phylum, class, order or family. Defaults to family. Note: If "kingdom" is used, the term_conversion() method should

first be applied.

valid A logical indicating whether the returned data should include valid or invalid

entries (defaults to valid).

Value

A validated tibble

Examples

```
get_validity(load_sample(), valid = TRUE)
```

load_population

Load previously merged GBIF and NCBI taxonomies

Description

load_population() imports a previously merged taxonomy from your file system. An example of an previously merged taxonomy can be downloaded from https://drive.google.com/file/d/1gpvm9QKdOcuGo_cIXPkAgGlB-qfKZZU6/view?usp=sharing.

Usage

load_population(x)

Arguments

Χ

Path to a text file containing previously merged NCBI and GBIF taxonomies (compressed or uncompressed).

Value

A tibble

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Examples

```
## Not run: load_population("path/to/merged_taxonomies")
```

load_sample

Load a sample of previously merged GBIF and NCBI taxonomies

Description

load_sample() returns a small subset of previously merged GBIF and NCBI taxonomies.

Usage

```
load_sample()
```

Value

A tibble containing a sample of merged GBIF and NCBI taxonomic data.

Examples

```
load_sample()
```

load_taxonomies

Load and merge GBIF and NCBI taxonomic data

Description

load_taxonomies() parses and merges a GBIF Taxon.tsv() file (available within the https://hosted-datasets.gbif.org/datasets/backbone/current/backbone.zip archive) and a Taxonkit (https://bioinf.shenwei.me/taxonkit/download/) output file obtained by running: taxonkit list --ids 1 | taxonkit lineage --show-lineage-taxids --show-lineage-ranks --show-rank --show-name > All.lineages.tsv.

Usage

```
load_taxonomies(GBIF_path, NCBI_path)
```

Arguments

GBIF_path Path to the GBIF backbone taxonomy (compressed or uncompressed).

NCBI_path Path to the NCBI taxonomy (compressed or uncompressed).

Value

A tibble containing merged GBIF and NCBI taxonomic data.

Examples

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plot_mdb	Generic for plot_mdb methods	
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Description

A custom generic for methods that plot taxonbridge data types

Usage

```
plot_mdb(x)
```

Arguments

x An object of the class one_rank or the class all_ranks.

Value

A ggplot2 derived plot

Examples

```
plot_mdb(prepare_rank_dist(load_sample(), NCBI = TRUE, GBIF = TRUE))
plot_mdb(prepare_comparable_rank_dist(load_sample()))
plot_mdb(prepare_rank_dist(get_status(load_sample(),status = "synonym"), NCBI = TRUE))
plot_mdb(prepare_comparable_rank_dist(get_validity(get_status(load_sample()), valid = TRUE)))
```

```
prepare_comparable_rank_dist
```

Get comparable NCBI and GBIF taxonomic ranks

Description

prepare_comparable_rank_dist(), like prepare_rank_dist(), returns taxonomic ranks aggregated by frequency for data derived from the NCBI, the GBIF, or both. However, prepare_comparable_rank_dist() only includes taxonomic ranks that have at least one NCBI and one GBIF representative.

Usage

```
prepare_comparable_rank_dist(x, GBIF = TRUE, NCBI = TRUE)
```

Arguments

Χ	A tibble created with load_	_taxonomies() or load_	_population()	orload_sample().
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GBIF A boolean indicating whether GBIF taxonomic ranks are to be retrieved.

NCBI A boolean indicating whether NCBI taxonomic ranks are to be retrieved.

Value

A list of tibble(s) assigned to the S3 class one_rank or to the S3 class all_ranks.

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Examples

```
prepare_comparable_rank_dist(load_sample())
prepare_comparable_rank_dist(get_status(load_sample(), "accepted"), NCBI = FALSE)
```

prepare_rank_dist

Get all NCBI and GBIF taxonomic ranks

Description

prepare_rank_dist() returns taxonomic ranks aggregated by frequency for data derived from the NCBI, the GBIF, or both.

Usage

```
prepare_rank_dist(x, GBIF = FALSE, NCBI = FALSE)
```

Arguments

x A tibble created with load_taxonomies() or load_population() or load_sample().

GBIF A boolean indicating whether GBIF taxonomic ranks are to be retrieved.

NCBI A boolean indicating whether NCBI taxonomic ranks are to be retrieved.

Value

A list of tibble(s) assigned to the S3 class one_rank or to the S3 class all_ranks.

Examples

```
prepare_rank_dist(load_sample(), NCBI=TRUE, GBIF=TRUE)
prepare_rank_dist(load_sample(), NCBI=TRUE)
```

term_conversion

Convert GBIF terms to NCBI terms

Description

term_conversion converts GBIF terminology to NCBI terminology where there is no biological provenance for the difference.

Usage

```
term_conversion(x)
```

Arguments

x A tibble created with load_taxonomies() or load_population() or load_sample().

Value

A tibble with converted terms.

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Examples

term_conversion(load_sample())

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