# Package 'taxa'

February 20, 2024

Title Classes for Storing and Manipulating Taxonomic Data

Type Package

RoxygenNote 7.3.1

```
Description Provides classes for storing and manipulating taxonomic data.
      Most of the classes can be treated like base R vectors (e.g. can be used
      in tables as columns and can be named). Vectorized classes can store taxon names
      and authorities, taxon IDs from databases, taxon ranks, and other types of
      information. More complex classes are provided to store taxonomic trees and
      user-defined data associated with them.
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      vctrs
LazyLoad yes
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License MIT + file LICENSE
URL https://docs.ropensci.org/taxa/, https://github.com/ropensci/taxa
BugReports https://github.com/ropensci/taxa/issues
Roxygen list(markdown = TRUE)
Imports dplyr,
     magrittr,
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      rlang,
      stringr,
      crayon,
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      pillar,
      methods,
      viridisLite,
      cli
Suggests roxygen2 (>= 6.0.1),
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```

## $\textbf{X-schema.org-applicationCategory} \ \ \textbf{Taxonomy}$

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# **R** topics documented:

as_data_frame
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classification
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as\_data\_frame

as\_data\_frame

Convert a taxa object to a data.frame

#### **Description**

Convert the information in a taxa object to a data.frame using base R vectors as columns. Use as tibble to convert to tibbles.

#### Usage

```
as_data_frame(
    x,
    row.names = NULL,
    optional = FALSE,
    ...,
    stringsAsFactors = FALSE
)
```

#### **Arguments**

An object defined by taxa, such as taxon or taxon\_id

NULL or a character vector giving the row names for the data frame. Missing values are not allowed.

optional logical. If TRUE, setting row names and converting column names (to syntactic names: see make.names) is optional. Note that all of R's base package as.data.frame() methods use optional only for column names treatment, basically with the meaning of data.frame(\*, check.names = !optional). See also the make.names argument of the matrix method.

... additional arguments to be passed to or from methods.

stringsAsFactors

logical: should the character vector be converted to a factor?

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as\_taxon

Convert to a taxon vector

#### **Description**

Convert other objects to taxon vectors. Compatible base R vectors can also be converted using the taxon constructor.

#### Usage

```
as_taxon(x, ...)
```

#### **Arguments**

x An object to be converted to a taxon vector

... Additional parameters.

#### **Examples**

```
# Convert a taxonomy object to a taxon vector
x <- taxonomy(taxon(name = c('Carnivora', 'Felidae', 'Panthera', 'Panthera leo',
                              'Panthera tigris', 'Ursidae', 'Ursus', 'Ursus arctos'),
                    rank = c('order', 'family', 'genus', 'species',
                              'species', 'family', 'genus', 'species'),
                    id = taxon_id(c('33554', '9681', '9688', '9689',
                                     '9694', '9632', '9639', '9644'),
                                   db = 'ncbi'),
             auth = c('Bowdich, 1821', 'Fischer de Waldheim, 1817', 'Oken, 1816', 'L., 1758',
                        'L., 1758', 'Fischer de Waldheim, 1817', 'L., 1758', 'L., 1758')),
              supertaxa = c(NA, 1, 2, 3, 3, 1, 6, 7))
names(x) <- letters[1:8]</pre>
as_taxon(x)
# Convert base R vectors
as_taxon(c('Carnivora', 'Felidae', 'Panthera', 'Panthera leo'))
as_taxon(factor(c('Carnivora', 'Felidae', 'Panthera', 'Panthera leo')))
```

classification

Taxon class

#### **Description**

**Experimental** Used to store classifications in reference to a taxonomic tree.

```
classification(x = NULL, taxonomy = NULL, .names = NULL)
```

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#### **Arguments**

x One of:

• A list where each item represents a series of nested taxa. The contents of the list can be in any form that can be converted to a taxon vector.

• The indexes/names of each instance of a taxon in a taxonomy object specified by the taxonomy option. Can be any length, but must consist of valid indexes for taxa in the taxonomy object.

taxonomy

A taxonomy object. Only needed if taxon indexes are supplied as the first argu-

ment.

. names The names of the vector.

#### Value

An S3 object of class taxa\_classification

#### See Also

```
Other classes: [.taxa_classification(), taxon(), taxon_authority(), taxon_db(), taxon_id(), taxon_rank()
```

```
# Create classification vector with a list
x <- classification(list(</pre>
  c('Carnivora', 'Felidae', 'Panthera', 'Panthera leo'),
  c('Carnivora', 'Felidae', 'Panthera', 'Panthera tigris'),
  c('Carnivora', 'Ursidae', 'Ursus', 'Ursus arctos'),
  c('Carnivora', 'Ursidae', 'Ursus', 'Ursus arctos'),
  c('Carnivora', 'Felidae', 'Panthera', 'Panthera tigris')
))
# Create classification vector with indexes and a taxonomy
x \leftarrow classification(c(3, 4, 4, 5, 5, 6, 8, 8, 2, 5, 6, 2),
                     taxonomy(c('Carnivora', 'Felidae', 'Panthera', 'Panthera leo',
                                'Panthera tigris', 'Ursidae', 'Ursus', 'Ursus arctos'),
                              supertaxa = c(NA, 1, 2, 3, 3, 1, 6, 7))
x \leftarrow classification(c(3, 4, 4, 5, 5, 6, 8, 8, 2, 5, 6, 2),
               taxonomy(taxon(name = c('Carnivora', 'Felidae', 'Panthera', 'Panthera leo',
                                   'Panthera tigris', 'Ursidae', 'Ursus', 'Ursus arctos'),
                                    rank = c('order', 'family', 'genus', 'species',
                                              'species', 'family', 'genus', 'species'),
                                    id = taxon_id(c('33554', '9681', '9688', '9689',
                                                     '9694', '9632', '9639', '9644'),
                                                  db = 'ncbi'),
                                    auth = c('Bowdich, 1821', 'Fischer, 1817',
                                              'Oken, 1816', 'L., 1758',
                                              'L., 1758', 'Fischer, 1817',
                                              'L., 1758', 'L., 1758')),
```

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```
supertaxa = c(NA, 1, 2, 3, 3, 1, 6, 7)))
names(x) \leftarrow letters[1:12]
# Get parts of the classification vector
tax_name(x)
tax_rank(x)
tax_id(x)
tax_db(x)
tax_auth(x)
tax_author(x)
tax_date(x)
tax_cite(x)
# Manipulate classification vectors
x[1:3]
x[tax_rank(x) > 'family']
\# c(x, x)
# x['b'] <- NA
is.na(x)
# as.data.frame(x)
# tibble::as_tibble(x)
# Use as columns in tables
tibble::tibble(x = x, y = 1:12)
data.frame(x = x, y = 1:12)
```

db\_ref

Valid taxonomy databases

## Description

This defines the valid taxonomic databases that can be used in taxon\_db objects and objects that use taxon\_db objects, such as taxon\_id and taxon. db\_ref\$get can be used to see information for the databases. Users can add their own custom databases to the list using db\_ref\$set. For each database the following information is included:

- The URL for the website associated with the database
- · A short description
- The regular expression that defines valid taxon IDs
- The ranks used in the database if specified

#### Usage

db\_ref

#### Format

An object of class list of length 3.

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#### Attribution

This code is based on the code handling options in the knitr package.

#### **Examples**

```
# List all database definitions
db_ref$get()

# Get a specific database definition
db_ref$get('ncbi')

# Add or overwrite a database definition
db_ref$set(
    name = "my_new_database",
    url = "http://www.my_tax_database.com",
    desc = "I just made this up",
    id_regex = ".*"
)

# Reset definitions to default values
db_ref$reset()
```

internodes

Get internodes

## **Description**

Get internodes indexes for each taxon or another per-taxon value. An internode is a taxon with exactly one supertaxon and one subtaxon. These taxa can be removed without losing information on the relationships of the remaining taxa.

#### **Usage**

```
internodes(x)
```

#### **Arguments**

Χ

The object to get internodes for, such as a taxonomy object.

## See Also

```
Other taxonomy functions: leaves(), roots(), stems(), subtaxa(), supertaxa()
Other internode functions: is_internode()
```

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#### **Examples**

is\_classification

Check if is a classification

#### **Description**

Check if an object is the classification class

#### Usage

```
is_classification(x)
```

#### **Arguments**

Х

An object to test

is\_internode

Check if taxa are internodes

## Description

Check if each taxon is an internode. An internode is a taxon with exactly one supertaxon and one subtaxon. These taxa can be removed without losing information on the relationships of the remaining taxa.

#### Usage

```
is_internode(x)
```

#### **Arguments**

Х

The object to get internodes for, such as a taxonomy object.

#### See Also

Other internode functions: internodes()

is\_leaf 9

#### **Examples**

is\_leaf

Check if taxa are leaves

## Description

Check if each taxon is a leaf. A leaf is a taxon with no subtaxa. subtaxa.

#### Usage

```
is_leaf(x)
```

#### **Arguments**

Х

The object to get leaves for, such as a taxonomy object

## See Also

```
Other leaf functions: leaves(), n_leaves()
```

#### **Examples**

is\_root

Test if taxa are roots

## Description

Check if each taxon is a root. A root is a taxon with no supertaxon.

```
is_root(x, subset = NULL)
```

is\_stem

## **Arguments**

x An object containing taxonomic relationships, such as taxonomy objects.

subset The subset of the tree to search for roots to that subset. Can be indexes or names.

#### See Also

Other root functions: roots()

#### **Examples**

is\_stem

Check if taxa are stems

#### **Description**

Check if each taxon is a stem. A stem is any taxa from a root to the first taxon with multiple subtaxa.

#### Usage

```
is_stem(x)
```

#### **Arguments**

Х

An object with taxonomic relationships, like taxonomy objects.

#### See Also

Other stem functions: stems()

is\_taxon 11

is\_taxon

Check if something is a taxon object

## Description

Check if an object is of the taxon class

#### Usage

```
is_taxon(x)
```

#### **Arguments**

Х

An object to test

#### **Examples**

```
x <- taxon(c('A', 'B', 'C'))
is_taxon(x)
is_taxon(1:2)</pre>
```

is\_taxonomy

Check if something is a taxonomy

## Description

Check if an object is of the taxonomy class

## Usage

```
is_taxonomy(x)
```

#### **Arguments**

Х

An object to test

is\_taxon\_db

is\_taxon\_authority

Check if is a taxon\_authority

## Description

Check if an object is of the taxon\_authority class

## Usage

```
is_taxon_authority(x)
```

## **Arguments**

Χ

An object to test

## **Examples**

is\_taxon\_db

Check if something is a taxon\_db

## Description

Check if an object is of the taxon\_db class

## Usage

```
is_taxon_db(x)
```

#### **Arguments**

Χ

An object to test

```
x <- taxon_db(c('ncbi', 'ncbi', 'itis'))
is_taxon_db(x)
is_taxon_db(1:3)</pre>
```

is\_taxon\_id 13

is\_taxon\_id

Check if something is a taxon\_id object

## Description

Check if an object is of the taxon\_id class

## Usage

```
is_taxon_id(x)
```

## Arguments

Х

An object to test

## **Examples**

```
x <- taxon_id(c('9606', '1386', '4890', '4345'), db = 'ncbi')
is_taxon_id(x)
is_taxon_id(1:3)</pre>
```

is\_taxon\_rank

Check if something is a taxon\_rank

## Description

Check if an object is of the taxon\_rank class

## Usage

```
is_taxon_rank(x)
```

## Arguments

Х

An object to test

```
x <- taxon_rank(c('species', 'species', 'phylum', 'family'))
is_taxon_rank(x)
is_taxon_rank(1:3)</pre>
```

n\_leaves

leaves

Get leaves

#### **Description**

Get leaves indexes for each taxon or another per-taxon value. Leaves are taxa with no subtaxa.

## Usage

```
leaves(x, value = NULL, ...)
```

#### **Arguments**

x The object to get leaves for, such as a taxonomy object

value Something to return instead of indexes. Must be the same length as the number

of taxa.

... Additional arguments.

#### See Also

```
Other taxonomy functions: internodes(), roots(), stems(), subtaxa(), supertaxa()
Other leaf functions: is_leaf(), n_leaves()
```

#### **Examples**

n\_leaves

Number of leaves per taxon

## **Description**

Get the number of leaves per taxon. A leaf is a taxon with no subtaxa.

#### Usage

```
n_leaves(x)
```

#### **Arguments**

Χ

The object to get leaves for, such as a taxonomy object

n\_subtaxa 15

#### See Also

```
Other leaf functions: is_leaf(), leaves()
```

#### **Examples**

n\_subtaxa

Number of subtaxa per taxon

#### **Description**

Get the number of subtaxa per taxon.

#### Usage

```
n_subtaxa(x, subset = NULL, max_depth = NULL, include = FALSE)
```

#### **Arguments**

x The object to get subtaxa for, such as a taxonomy object. subset The subset of the tree to search. Can be indexes or names.

max\_depth The number of ranks to traverse. For example, max\_depth = 1 returns only im-

mediate subtaxa. By default (NULL) information for all subtaxa is returned (i.e.

subtaxa of subtaxa, etc).

include If TRUE, include information for each taxon in the output.

#### See Also

Other subtaxa functions: subtaxa()

n\_supertaxa

```
\# Only return data for some taxa (faster than subsetting the whole result) n\_subtaxa(x, subset = 1:3)
```

n\_supertaxa

Number of supertaxa per taxon

## Description

Get the number of supertaxa each taxon is contained in.

#### Usage

```
n_supertaxa(x, subset = NULL, max_depth = NULL, include = FALSE)
```

## Arguments

x	The object to get supertaxa for, such as a taxonomy object.
subset	The subset of the tree to search for roots to that subset. Can be indexes or names.
max_depth	The number of levels to traverse. For example, max_depth = 1 returns only immediate supertaxa. By default (NULL) information for all supertaxa is returned.
include	If TRUE, include information for each taxon in the output.

#### See Also

Other supertaxa functions: supertaxa()

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roots

Get root taxa

#### Description

Get the indexes of root taxa in a taxonomy.

#### Usage

```
roots(x, subset = NULL)
```

## **Arguments**

x An object containing taxonomic relationships, such as taxonomy objects.

subset The subset of the tree to search for roots to that subset. Can be indexes or names.

#### See Also

```
Other taxonomy functions: internodes(), leaves(), stems(), subtaxa(), supertaxa()
Other root functions: is_root()
```

#### **Examples**

stems

Get stems

#### **Description**

Get stem indexes for each taxon or another per-taxon value.

#### Usage

```
stems(x, value = NULL, ...)
```

#### **Arguments**

x An object with taxonomic relationships, like taxonomy objects.

value Something to return instead of indexes. Must be the same length as the number

of taxa.

... Additional arguments.

18 subtaxa

#### See Also

```
Other taxonomy functions: internodes(), leaves(), roots(), subtaxa(), supertaxa()
Other stem functions: is_stem()
```

## **Examples**

subtaxa

Get subtaxa

## **Description**

Get subtaxa indexes for each taxon or another per-taxon value. Subtaxa are taxa contained within a taxon.

#### Usage

```
subtaxa(x, subset = NULL, max_depth = NULL, include = FALSE, value = NULL, ...)
```

#### **Arguments**

Χ	The object to get subtaxa for, such as a taxonomy object.
subset	The subset of the tree to search. Can be indexes or names.
max_depth	The number of ranks to traverse. For example, max_depth = 1 returns only immediate subtaxa. By default (NULL) information for all subtaxa is returned (i.e. subtaxa of subtaxa, etc).
include	If TRUE, include information for each taxon in the output.
value	Something to return instead of indexes. Must be the same length as the number of taxa.
	Additional arguments.

#### See Also

```
Other taxonomy functions: internodes(), leaves(), roots(), stems(), supertaxa()
Other subtaxa functions: n_subtaxa()
```

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#### **Examples**

supertaxa

Get supertaxa

#### **Description**

Get supertaxa indexes for each taxon or another per-taxon value. Supertaxa are taxa a taxon is contained in.

#### Usage

```
supertaxa(
   x,
   subset = NULL,
   max_depth = NULL,
   include = FALSE,
   value = NULL,
   use_na = FALSE,
   ...
)
```

#### **Arguments**

x The object to get supertaxa for, such as a taxonomy object.

subset The subset of the tree to search for roots to that subset. Can be indexes or names.

max\_depth The number of levels to traverse. For example, max\_depth = 1 returns only immediate supertaxa. By default (NULL) information for all supertaxa is returned.

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include If TRUE, include information for each taxon in the output.

value Something to return instead of indexes. Must be the same length as the number

of taxa.

use\_na Add a NA to represent the root of the taxonomy (i.e. no supertaxon)

.. Additional arguments.

#### See Also

```
Other taxonomy functions: internodes(), leaves(), roots(), stems(), subtaxa()
Other supertaxa functions: n_supertaxa()
```

#### **Examples**

taxa\_taxon-class

Taxon class

## Description

Taxon class. See taxon for more information

## Description

Taxonomy class. See taxonomy for more information

 ${\tt taxa\_taxon\_authority\_class}$ 

Taxon authority class

## Description

Taxon authority class. See taxon\_authority for more information

taxa\_taxon\_db-class

Taxon database class

## Description

Taxon database class. See taxon\_db for more information

taxa\_taxon\_id-class

Taxon ID class

## Description

Taxon ID class. See taxon\_id for more information

## Description

Taxon rank class. See taxon\_rank for more information

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	taxon	Taxon class	
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## Description

Maturing Used to store information about taxa, such as names, ranks, and IDs.

#### Usage

```
taxon(name = character(0), rank = NA, id = NA, auth = NA, .names = NA, ...)
```

#### **Arguments**

name	The names of taxa. Inputs with be coerced into a character vector if anything else is given.
rank	The ranks of taxa. Inputs with be coerced into a taxon_rank vector if anything else is given.
id	The ids of taxa. These should be unique identifier and are usually associated with a database. Inputs with be coerced into a taxon_id vector if anything else is given.
auth	The authority of the taxon. Inputs with be coerced into a taxon_authority vector if anything else is given.
.names	The names of the vector.
	Additional arguments.

#### Value

An S3 object of class taxa\_taxon

#### See Also

```
Other classes: [.taxa_classification(), classification(), taxon_authority(), taxon_db(), taxon_id(), taxon_rank()
```

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```
tax_id(x)
tax_db(x)
tax_auth(x)
tax_author(x)
tax_date(x)
tax_cite(x)
# Set parts of the taxon name vector
tax_name(x) <- tolower(tax_name(x))</pre>
tax_rank(x)[1] \leftarrow NA
tax_name(x)['b'] <- 'Billy'</pre>
tax_id(x) <- '9999'
tax_db(x) <- 'itis'
tax_auth(x) <- NA
tax_author(x)[2:3] <- c('Joe', 'Billy')</pre>
tax_date(x) <- c('1999', '2013', '1796', '1899')
tax\_cite(x)[1] \leftarrow Linnaeus, C. (1771). Mantissa plantarum altera generum.'
# Manipulate taxon name vectors
x[1:3]
x[tax_rank(x) > 'family']
x['b'] <- NA
x[c('c', 'd')] \leftarrow 'unknown'
is.na(x)
# Use as columns in tables
tibble::tibble(x = x, y = 1:4)
data.frame(x = x, y = 1:4)
# Converting to tables
tibble::as_tibble(x)
as_data_frame(x)
```

taxon\_authority

Taxon authority class

#### **Description**

Maturing Used to store information on taxon authorities, such as author names, date, and citation.

```
taxon_authority(
  author = character(),
  date = "",
  citation = "",
  .names = "",
  extract_date = TRUE
)
```

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#### **Arguments**

author Zero or more author names.

date Zero or more dates.

citation Zero or more literature citations.

. names The names of the vector.

input is given, then the date is separated from the author input.

#### Value

An S3 object of class taxa\_taxon\_authority

#### See Also

```
Other classes: [.taxa_classification(), classification(), taxon(), taxon_db(), taxon_id(), taxon_rank()
```

```
# Making new objects
x <- taxon_authority(c('A', 'B', 'C'))</pre>
x <- taxon_authority(c('Cham. & Schldl.', 'L.'),</pre>
                      date = c('1827', '1753'))
# Manipulating objects
as.character(x)
x[2]
x[2] \leftarrow 'ABC'
names(x) \leftarrow c('a', 'b')
x['b'] <- 'David Bowie'
tax_author(x)[1] <- tolower(tax_author(x)[1])</pre>
tax_author(x)
tax_date(x) <- c('2000', '1234')
tax_cite(x)[2] <- c('Linnaeus, C. (1771). Mantissa plantarum altera generum.')</pre>
tax_cite(x)
# Using as columns in tables
tibble::tibble(x = x, y = 1:2)
data.frame(x = x, y = 1:2)
# Converting to tables
tibble::as_tibble(x)
as_data_frame(x)
```

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taxon\_db

Taxon database class

#### **Description**

**Maturing** Used to store the names of taxon databases defined in db\_ref. Primarily used in other classes like taxon\_id to define databases for each item.

## Usage

```
taxon_db(db = character(), .names = NULL, ...)
```

#### **Arguments**

db Zero or more taxonomic database names. Should be a name contained in db\_ref.

Inputs will be transformed to a character vector if possible.

. names The names of the vector.. . . Additional arguments.

#### Value

An S3 object of class taxa\_taxon\_db

#### See Also

```
Other classes: [.taxa_classification(), classification(), taxon(), taxon_authority(), taxon_id(), taxon_rank()
```

```
# Making new objects
x <- taxon_db(c('ncbi', 'ncbi', 'itis'))
x

# Manipulating objects
as.character(x)
x[2:3]
x[2:3] <- 'nbn'
names(x) <- c('a', 'b', 'c')
x['b']
x['b'] <- 'nbn'
x[x == 'itis'] <- 'gbif'

# Using as columns in tables
tibble::tibble(x = x, y = 1:3)
data.frame(x = x, y = 1:3)
# Converting to tables
tibble::as_tibble(x)</pre>
```

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```
as_data_frame(x)

# Trying to use an invalid database generates an error
# x <- taxon_db(c('ncbi', 'ncbi', 'my_custom_db'))
# x[x == 'itis'] <- 'my_custom_db'

# Listing known databases and their properties
db_ref$get()

# Adding and using a new database
db_ref$set(name = 'my_custom_db', desc = 'I just made this up')
db_ref$get()
x <- taxon_db(c('ncbi', 'ncbi', 'my_custom_db'))</pre>
```

taxon\_id

Taxon ID class

#### **Description**

**Maturing** Used to store the ID corresponding to taxa, either arbitrary or from a particular taxonomy database. This is typically used to store taxon IDs in taxon objects.

## Usage

```
taxon_id(id = character(), db = "", .names = NULL)
```

#### **Arguments**

id	Zero or more taxonomic ids. Inputs will be transformed to a character vector if possible.
db	The name(s) of the database(s) associated with the IDs. If not NA (the default), the input must consist of names of databases in db_ref\$get().
.names	The names that will be applied to the vector.

#### Value

```
An S3 object of class taxa_taxon_id
```

#### See Also

```
Other classes: [.taxa_classification(), classification(), taxon(), taxon_authority(), taxon_db(), taxon_rank()
```

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#### **Examples**

```
# Making new objects
x \leftarrow taxon_id(c('A', 'B', 'C'))

x \leftarrow taxon_id(c('9606', '1386', '4890', '4345'), db = 'ncbi')
x <- taxon_id(c('9606', '1386', '4890', '4345'),
db = c('ncbi', 'ncbi', 'itis', 'itis'))
names(x) <- c('a', 'b', 'c', 'd')</pre>
# Manipulating objects
as.character(x)
x[2:3]
x[2:3] \leftarrow 'ABC'
x[c('a', 'c')] <- '123'
x[['b']] \leftarrow taxon_id('123423', db = 'ncbi')
tax_db(x)
tax_db(x) <- 'nbn'
c(x, x)
# Using as columns in tables
tibble::tibble(x = x, y = 1:4)
data.frame(x = x, y = 1:4)
# Convert to tables
tibble::as_tibble(x)
as_data_frame(x)
# Trying to use an invalid ID with a specified database causes an error
#taxon_id('NOLETTERS', db = 'ncbi')
```

taxon\_rank

Taxon rank class

#### **Description**

**Maturing** Used to store taxon ranks, possibly associated with a taxonomy database. This is typically used to store taxon ranks in taxon objects.

```
taxon_rank(
  rank = character(),
  .names = NULL,
  levels = NULL,
  guess_order = TRUE
)
```

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#### **Arguments**

rank Zero or more taxonomic rank names. Inputs will be transformed to a character

vector.

levels A named numeric vector indicating the names and orders of possible taxonomic

ranks. Higher numbers indicate for fine-scale groupings. Ranks of unknown

order can be indicated with NA instead of a number.

guess\_order If TRUE and no rank order is given using numbers, try to guess order based on

rank names.

#### Value

An S3 object of class taxa\_taxon\_rank

#### See Also

```
Other classes: [.taxa_classification(), classification(), taxon(), taxon_authority(), taxon_db(), taxon_id()
```

```
# Making new objects
x <- taxon_rank(c('species', 'species', 'phylum', 'family'))</pre>
# Specifiying level order
taxon_rank(c('A', 'B', 'C', 'D', 'A', 'D', 'D'),
           levels = c('D', 'C', 'B', 'A'))
taxon_rank(c('A', 'B', 'C', 'D', 'A', 'D', 'D'),
           levels = c(D = NA, A = 10, B = 20, C = 30))
names(x) \leftarrow c('a', 'b', 'c', 'd')
# Manipulating objects
as.character(x)
as.factor(x)
as.ordered(x)
x[2:3]
x[x > 'family'] <- taxon_rank('unknown')</pre>
x[1] <- taxon_rank('order')</pre>
x['b']
x['b'] <- 'order'
# Using as columns in tables
tibble::tibble(x = x, y = 1:4)
data.frame(x = x, y = 1:4)
# Converting to tables
tibble::as_tibble(x)
as_data_frame(x)
# Trying to add an unknown level as a character causes an error
```

```
#x[2] <- 'superkingdom'

# But you can add a new level using taxon_rank objects
x[2] <- taxon_rank('superkingdom')</pre>
```

```
tax_auth.taxa_classification

Set and get taxon authorities
```

#### **Description**

Set and get the taxon authorities in objects that have them, such as taxon objects. Note that this sets all the authority information, such as author name, date, and citations. To set or get just one of part of the authorities, use tax\_author, tax\_date, or tax\_cite instead.

#### Usage

```
## S3 method for class 'taxa_classification'
tax_auth(x)

## S3 replacement method for class 'taxa_classification'
tax_auth(x) <- value

tax_auth(x) <- value

## S3 method for class 'taxa_taxon'
tax_auth(x)

## S3 replacement method for class 'taxa_taxon'
tax_auth(x) <- value

## S3 method for class 'taxa_taxonomy'
tax_auth(x)

## S3 replacement method for class 'taxa_taxonomy'
tax_auth(x)</pre>
```

## **Arguments**

x An object with taxon authorities.

value The taxon IDs to set. Inputs will be coerced into a taxon\_id vector.

#### **Examples**

#### **Description**

Set and get taxon authors in objects that have them, such as taxon\_authority objects.

```
## S3 method for class 'taxa_classification'
tax_author(x)
## S3 replacement method for class 'taxa_classification'
tax_author(x) <- value</pre>
tax_author(x)
tax_author(x) <- value
## S3 method for class 'taxa_taxon'
tax_author(x)
## S3 replacement method for class 'taxa_taxon'
tax_author(x) <- value
## S3 replacement method for class 'taxa_taxon_authority'
tax_author(x) <- value</pre>
## S3 method for class 'taxa_taxon_authority'
tax_author(x)
## S3 method for class 'taxa_taxonomy'
tax_author(x)
## S3 replacement method for class 'taxa_taxonomy'
tax_author(x) <- value
```

x An object with taxon authors.value The taxon authors to set. Inputs will be coerced into a character vector.

#### **Examples**

```
tax_cite.taxa_classification

Set and get taxon authority citations
```

#### **Description**

Set and get the taxon authority citations in objects that have them, such as taxon\_authority objects.

```
## S3 method for class 'taxa_classification'
tax_cite(x)
## S3 replacement method for class 'taxa_classification'
tax_cite(x) <- value</pre>
tax_cite(x)
tax_cite(x) <- value</pre>
## S3 method for class 'taxa_taxon'
tax_cite(x)
## S3 replacement method for class 'taxa_taxon'
tax_cite(x) <- value</pre>
## S3 replacement method for class 'taxa_taxon_authority'
tax_cite(x) <- value</pre>
## S3 method for class 'taxa_taxon_authority'
tax_cite(x)
## S3 method for class 'taxa_taxonomy'
tax_cite(x)
```

```
## S3 replacement method for class 'taxa_taxonomy'
tax_cite(x) <- value</pre>
```

x An object with taxon authority dates.

value The taxon citations to set. Inputs will be coerced into a taxon\_authority vector.

#### **Examples**

tax\_date.taxa\_classification

Set and get taxon authority dates

#### **Description**

Set and get the taxon authority dates in objects that have them, such as taxon\_authority objects.

```
## S3 method for class 'taxa_classification'
tax_date(x)

## S3 replacement method for class 'taxa_classification'
tax_date(x) <- value

tax_date(x)

## S3 method for class 'taxa_taxon'
tax_date(x)

## S3 replacement method for class 'taxa_taxon'
tax_date(x) <- value

## S3 replacement method for class 'taxa_taxon'
tax_date(x) <- value</pre>
```

```
## S3 method for class 'taxa_taxon_authority'
tax_date(x)

## S3 method for class 'taxa_taxonomy'
tax_date(x)

## S3 replacement method for class 'taxa_taxonomy'
tax_date(x) <- value</pre>
```

x An object with taxon authority dates.

value The taxon authority dates to set. Inputs will be coerced into a character vector.

#### **Examples**

tax\_db.taxa\_classification

Set and get taxon ID databases

## **Description**

Set and get the taxon ID databases in objects that have them, such as taxon\_id objects.

```
## S3 method for class 'taxa_classification'
tax_db(x)

## S3 replacement method for class 'taxa_classification'
tax_db(x) <- value

tax_db(x)

tax_db(x) <- value

## S3 method for class 'taxa_taxon'
tax_db(x)

## S3 replacement method for class 'taxa_taxon'</pre>
```

```
tax_db(x) <- value

## S3 method for class 'taxa_taxon_id'
tax_db(x)

## S3 replacement method for class 'taxa_taxon_id'
tax_db(x) <- value

## S3 method for class 'taxa_taxonomy'
tax_db(x)

## S3 replacement method for class 'taxa_taxonomy'
tax_db(x) <- value</pre>
```

x An object with taxon authority dates.

value The taxon citations to set. Inputs will be coerced into a taxon\_db vector.

#### **Examples**

```
x \leftarrow taxon_id(c('9606', '1386', '4890', '4345'), db = 'ncbi') tax_db(x) tax_db(x) <- 'nbn' tax_db(x)[2] <- 'itis'
```

```
tax_id.taxa_classification

Set and get taxon IDs
```

#### **Description**

Set and get the taxon IDs in objects that have them, such as taxon objects.

```
## $3 method for class 'taxa_classification'
tax_id(x)

## $3 replacement method for class 'taxa_classification'
tax_id(x) <- value

tax_id(x)

tax_id(x) <- value</pre>
```

```
## S3 method for class 'taxa_taxon'
tax_id(x)

## S3 replacement method for class 'taxa_taxon'
tax_id(x) <- value

## S3 method for class 'taxa_taxonomy'
tax_id(x)

## S3 replacement method for class 'taxa_taxonomy'
tax_id(x) <- value</pre>
```

x An object with taxon IDs.value The taxon IDs to set. Inputs will be coerced into a taxon\_id vector.

#### **Examples**

```
tax_name.taxa_classification

Set and get taxon names
```

#### Description

Set and get the taxon names in objects that have them, such as taxon objects. Note that this is not the same as adding vector names with names.

```
## S3 method for class 'taxa_classification'
tax_name(x)
## S3 replacement method for class 'taxa_classification'
tax_name(x) <- value
tax_name(x)</pre>
```

```
tax_name(x) <- value

## S3 method for class 'taxa_taxon'
tax_name(x)

## S3 replacement method for class 'taxa_taxon'
tax_name(x) <- value

## S3 method for class 'taxa_taxonomy'
tax_name(x)

## S3 replacement method for class 'taxa_taxonomy'
tax_name(x) <- value</pre>
```

x An object with taxon names.

value The taxon names to set. Inputs will be coerced into a character vector.

#### **Examples**

```
tax_rank.taxa_classification

Set and get taxon ranks
```

#### **Description**

Set and get the taxon ranks in objects that have them, such as taxon objects.

```
## S3 method for class 'taxa_classification'
tax_rank(x)
## S3 replacement method for class 'taxa_classification'
tax_rank(x) <- value
tax_rank(x)</pre>
```

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```
tax_rank(x) <- value

## S3 method for class 'taxa_taxon'
tax_rank(x)

## S3 replacement method for class 'taxa_taxon'
tax_rank(x) <- value

## S3 method for class 'taxa_taxonomy'
tax_rank(x)

## S3 replacement method for class 'taxa_taxonomy'
tax_rank(x) <- value</pre>
```

#### **Arguments**

An object with taxon ranks.

value The taxon ranks to set. Inputs will be coerced into a taxon\_rank vector.

## **Examples**

%in%

Value matching for taxa package

## Description

A wrapper for the base value matching %in% that is used to take into consideration features of the taxa package.

#### Usage

```
x %in% table
```

## Arguments

x vector or NULL: the values to be matched. Long vectors are supported.table vector or NULL: the values to be matched against. Long vectors are not supported.

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