Package 'taxlist'

July 15, 2021

Version 0.2.2

Encoding UTF-8

Title Handling Taxonomic Lists

Depends R(>= 3.5.0)

Imports foreign, methods, stats, taxa, taxize, stringdist, utils, vegdata

Suggests ape, knitr, testthat, rmarkdown, roxygen2

LazyData true

Description Handling taxonomic lists through objects of class 'taxlist'.

This package provides functions to import species lists from 'Turboveg' (<<u>https://www.synbiosys.alterra.nl/turboveg</u>/>) and the possibility to create backups from resulting R-objects. Also quick displays are implemented as summary-methods.

License GPL (≥ 2)

URL https://cran.r-project.org/package=taxlist,

https://github.com/ropensci/taxlist,

https://docs.ropensci.org/taxlist/

BugReports https://github.com/ropensci/taxlist/issues

Collate 'imports.R' 'internal.R' 'deprecated-functions.R' 'replace_x.R' 'dissect_name.R' 'clean_strings.R' 'taxlist-class.R' 'clean.R' 'as.list.R' 'taxon_views.R' 'count_taxa.R' 'taxon_names.R' 'taxon_relations.R' 'taxon_traits.R' 'levels.R' 'accepted_name.R' 'get_children.R' 'merge_taxa.R' 'Extract.R' 'subset.R' 'backup_object.R' 'summary.R' 'df2taxlist.R' 'tv2taxlist.R' 'tnrs.R' 'tax2traits.R' 'match_names.R' 'print_name.R' 'indented_list.R' 'Easplist-data.R' 'taxlist-package.R'

VignetteBuilder knitr

RoxygenNote 7.1.1.9000

NeedsCompilation no

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Repository CRAN

Date/Publication 2021-07-15 16:20:02 UTC

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taxlist-package

Description

The class taxlist is defined in this package using the S4 language. The main task of taxlist objects is to keep the taxonomic coherence of information included in taxonomic lists and to implement functions (methods) for a proper data handling. Objects of class taxlist can be included in further objects, for instance in biodiversity records as done in the package vegtable.

Details

The class taxlist is defined in this package using the S4 language. The main task of taxlist objects is to keep the taxonomic coherence of information included in taxonomic lists and to implement functions (methods) for a proper data handling. Objects of class taxlist can be included in further objects, for instance in biodiversity records as done in the package vegtable.

For a more detailed description of this package, see Alvarez & Luebert (2018).

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

References

Alvarez M, Luebert F (2018). The taxlist package: managing plant taxonomic lists in R. *Biodiversity Data Journal* 6: e23635. doi: 10.3897/bdj.6.e23635

accepted_name

Manage accepted names, synonyms and basionyms

Description

Taxon usage names for a taxon concept can be divided into three categories: accepted names, basionyms and synonyms. Each single taxon concept may at least have an accepted name, while basionym and synonyms are optional. The functions accepted_name, basionym and synonyms can be used either to display the respective usage names or to set usage names in one of those categories.

Usage

```
accepted_name(taxlist, ConceptID, ...)
## S4 method for signature 'taxlist,numeric'
accepted_name(taxlist, ConceptID, show_traits = FALSE, ...)
## S4 method for signature 'taxlist,missing'
```

```
accepted_name(taxlist, ConceptID, ...)
accepted_name(taxlist, ConceptID) <- value</pre>
## S4 replacement method for signature 'taxlist,numeric,numeric'
accepted_name(taxlist, ConceptID) <- value</pre>
synonyms(taxlist, ConceptID, ...)
## S4 method for signature 'taxlist, numeric'
synonyms(taxlist, ConceptID, ...)
## S4 method for signature 'taxlist,missing'
synonyms(taxlist, ConceptID, ...)
basionym(taxlist, ConceptID, ...)
## S4 method for signature 'taxlist,numeric'
basionym(taxlist, ConceptID, ...)
## S4 method for signature 'taxlist,missing'
basionym(taxlist, ConceptID, ...)
basionym(taxlist, ConceptID) <- value</pre>
## S4 replacement method for signature 'taxlist,numeric,numeric'
basionym(taxlist, ConceptID) <- value</pre>
```

Arguments

taxlist	An object of class taxlist.
ConceptID	Integer containing concept IDs where to request or set names for one category.
	Further arguments passed among methods.
show_traits	Logical value, whether traits should be included in the output of accepted_name or not.
value	Integer containing usage IDs to be set to the respective category in the respective taxon concept.

Details

The function accepted_name retrieves the accepted names for the indicated taxon concepts or for the whole taxlist object. By using show_traits=TRUE, the respective taxon traits will be displayed as well, providing an overview of taxa included in the object. The replacement method for this function will set the respective usage name IDs as accepted names for the respective taxon concept, provided that these names are already set as synonyms in the respective concepts.

The function synonyms is working in a similar way as accepted_name, but this function does not include taxon traits in the output and there is no replacing method for synonyms. Alternatives for

as.list

inserting new synonyms into a taxon concept are either moving synonyms from other taxa by using change_concept<- or inserting new names in the object by using add_synonym().

The function basionym is retrieving and setting basionyms in the respective taxon concepts similarly to accepted_name, but this function does not retrieve any information on taxon traits, either.

Value

Most of the methods return information in data frames, while replacement methods do it as taxlist objects.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

See Also

add_synonym() change_concept<-</pre>

Examples

```
## Set a different accepted name for Cyclosorus interruptus
summary(Easplist, "Cyclosorus interruptus")
accepted_name(Easplist, 50074) <- 53097
summary(Easplist, 50074)
## Inserting a new name first
summary(Easplist, "Basella alba")
```

```
Easplist <- add_synonym(taxlist=Easplist, ConceptID=68,
    TaxonName="Basella cordifolia", AuthorName="Lam.")
summary(Easplist, 68)
accepted_name(Easplist, 68) <- 56139
summary(Easplist, 68)
```

as.list

Coerce an S4 object to a list.

Description

Coercion of S4 objects to lists can be applied to explore their content, avoiding errors caused by their validation.

Usage

```
S4_to_list(x)
## S4 method for signature 'taxlist'
as.list(x, ...)
```

Arguments

х	An object of class taxlist or any S4 class.
	further arguments passed to or from other methods.

Details

The function S4_to_list transforms any S4 object to a list setting slots to elements of the list and it is running internally in the method as.list for taxlist objects.

Value

An object of class list.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

Examples

```
Easplist <- as.list(Easplist)
class(Easplist)</pre>
```

backup_object

Make and load backups of R objects

Description

When work with data becomes risky, the best practice is to produce backup files. The function of backup_object is a wrapper of save(), adding a time stamp and a suffix to the name of the resulting file (an R image file with extension *.rda). The function load_last is adapted to this style, loading the newest version to the session.

Usage

```
backup_object(
    ...,
    objects = character(),
    file,
    stamp = TRUE,
    overwrite = FALSE
)
load_last(file, fext = ".rda")
```

backup_object

Arguments

	Names of the objects to be saved (either symbols or character strings).
objects	A character vector indicating the names of objects to be included in the backup file.
file	A character value indicating the name of the backup file, without the extension.
stamp	A logical value indicating whether time should be stamped in the backup name or not.
overwrite	A logical value indicating whether existing files must be overwritten or not.
fext	A character value indicating the file extension (including the dot symbol).

Details

In both functions the argument file may include either the path relative to the working directory or the absolute path to the file, excluding stamps and extension. For overwrite=FALSE (the default), a numeric suffix will be added to the backup's name, if another backup was produced at the same day. For overwrite=TRUE no suffix will be included in the file and existing files will be overwritten.

The function load_last() will load the newest version among backups stored in the same folder, provided that the backup name includes a time stamp.

Value

An R image with extension *.rda.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

See Also

save load.

Examples

```
## Not run:
## A subset with Pseudognaphalium and relatives
Pseudognaphalium <- subset(x=Easplist, subset=grepl("Pseudognaphalium",
    TaxonName), slot="names")
Pseudognaphalium <- get_parents(Easplist, Pseudognaphalium)
## Create a backup with date stamp
backup_object(Pseudognaphalium, file="Pseudonaphalium")
## The same
backup_object(objects="Pseudognaphalium", file="Pseudonaphalium")
## To load the last backup into a session
```

```
## End(Not run)
```

```
## Load pre-installed backup
load_last(file.path(path.package("taxlist"), "extdata", "Podocarpus"))
```

```
clean
```

Delete orphaned records

Description

Manipulation of slots may generate orphaned entries in taxlist objects. The function clean deletes such entries and restores the consistency of the objects.

Usage

clean(object, ...)

S4 method for signature 'taxlist'
clean(object, times = 2, ...)

Arguments

object	A taxlist object.
	Further arguments passed from or to other methods.
times	An integer indicating how many times the cleaning should be repeated.

Details

Cleaning of objects will follow the deletion of orphaned names, orphaned taxon trait entries, and orphaned parent entries.

Value

A clean taxlist object.

Author(s)

Miguel Alvarez.

Examples

```
## Direct manipulation of slot taxonRelations generates an invalid object
Easplist@taxonRelations <- Easplist@taxonRelations[1:5, ]
## Not run:
summary(Easplist)</pre>
```

End(Not run)

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```
## Now apply cleaning
Easplist <- clean(Easplist)
summary(Easplist)</pre>
```

clean_strings Cleaning character strings.

Description

Multiple, leading and trailing white spaces as well as wrong encodings may cause serious problems in information dealing with taxonomic names. The function clean_strings get rid of them.

Usage

```
clean_strings(x, ...)
## S4 method for signature 'character'
clean_strings(x, from = "utf8", to = "utf8", ...)
## S4 method for signature 'factor'
clean_strings(x, from = "utf8", to = "utf8", ...)
## S4 method for signature 'data.frame'
clean_strings(x, from = "utf8", to = "utf8", ...)
```

Arguments

Х	Object to be cleaned.
	Further arguments passed among methods (not yet in use).
from, to	Arguments passed to iconv().

Details

This function automatically deletes leading, trailing and multiple white spaces, either in strings (method character), levels (method factor or in single columns (method data.frame).

Value

The same as input x.

Author(s)

Miguel Alvarez.

Examples

```
library(taxlist)
clean_strings(" Cyperus papyrus L. ")
```

count_taxa

Count taxa within a taxlist object.

Description

Counting number of taxa within taxlist objects or character vectors containing taxon names.

Usage

```
count_taxa(object, data, ...)
## S4 method for signature 'character,missing'
count_taxa(object, na.rm = TRUE, ...)
## S4 method for signature 'factor,missing'
count_taxa(object, na.rm = TRUE, ...)
## S4 method for signature 'taxlist,missing'
count_taxa(object, level, ...)
## S4 method for signature 'formula,taxlist'
count_taxa(object, data, include_na = FALSE, suffix = "_count", ...)
```

Arguments

object	An object containing a taxonomic list or a formula.
data	An object of class taxlist in the formula method.
	further arguments passed among methods.
na.rm	Logical value, whether NAs have to be removed from the input vector or not.
level	Character value indicating the taxonomic rank of counted taxa.
include_na	Logical value indicating whether NA values in a taxon trait should be considered for counting taxa or just ignored (only used in formula method).
suffix	Character value used as suffix for the counted rank in the output data frame (only used in formula method).

Details

This function is written by convenience in order to reduce code for counting taxa within taxlist objects and it is just a wrapper of length().

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Value

An integer with the number of taxa.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

Examples

```
## factor method
count_taxa(iris$Species)
## taxlist method
count_taxa(Easplist)
count_taxa(Easplist, level="species")
## using a formula
count_taxa(~ lf_behn_2018, Easplist)
```

Deprecated-functions Deprecated functions

Description

Most of those functions have been replaced by alternative 'update' ones.

Usage

add_parent()
add_trait()
add_level()
replace_view()
taxlist2taxmap()
taxmap2taxlist()

```
df2taxlist
```

Description

Taxon lists may be provided in data frame format, which will be converted to a taxlist object.

Usage

```
df2taxlist(x, AcceptedName, ...)
## S4 method for signature 'data.frame,logical'
df2taxlist(x, AcceptedName, levels, ...)
## S4 method for signature 'data.frame,missing'
df2taxlist(x, AcceptedName, ...)
## S4 method for signature 'character,missing'
df2taxlist(x, AcceptedName, ...)
```

Arguments

х	A data frame or a character vector with taxon names.
AcceptedName	A logical vector indicating accepted names with value TRUE.
	Additional vectors to be added as columns in slottaxonNames.
levels	A vector with the names of the taxonomic ranks. This argument is passed to levels().

Details

In the method data.frame, the input data frame must have following columns:

TaxonUsageID Numeric code for the name.

TaxonConceptID Numeric code for the concept.

TaxonName Full name (usage), excluding author name.

AuthorName Author of the combination (taxon name).

If the argument AcceptedName is missing, all names will be assumed as accepted names. In the alternative character method, author names have to be added as additional vectors.

Be aware that the resulting object misses any information on taxon views, basionyms, parent concepts, hierarchical levels and taxon traits. All those elements can be added *a posteriori* by further functions provided in this package.

Value

A taxlist object.

dissect_name

Author(s)

Miguel Alvarez <kamapu78@gmail.com>.

Examples

```
## Read the table with names of Cyperus species
Cyperus <- read.csv(file=file.path(path.package("taxlist"), "cyperus",
        "names.csv"), stringsAsFactors=FALSE)
head(Cyperus)
## Convert to 'taxlist' object
Cyperus <- df2taxlist(Cyperus, AcceptedName =! Cyperus$SYNONYM)
summary(Cyperus)
## Create a 'taxlist' object from character vectors
Plants <- df2taxlist(c("Triticum aestivum","Zea mays"), AuthorName="L.")
summary(Plants, "all")
```

dissect_name

Dissect Scientific Names into their Elements

Description

Depending the degree of resolution and specific roles of nomenclature, strings containing taxon usage names (scientific names) are constructed with different parts. A string with names can be consequently split into those elements, meanwhile the number of elements may suggest the taxonomic ranks.

This function is a wrapper of strsplit(), while name element can be re-pasted if indicated in argument repaste.

Usage

```
dissect_name(x, split = " ", fixed = TRUE, repaste, ...)
```

Arguments

х	A character vector containing taxon names.
split, fixed,	
	Arguments passed to strsplit().
repaste	An integer vector indicating the elements of the name selected for the output.

Value

A character matrix with as many rows as names in the input vector. If repaste is indicated, then the output will be a character vector.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

See Also

strsplit()

Examples

```
Easplist <- subset(x=Easplist, subset=Level == "variety", slot="relations")
Easplist <- accepted_name(Easplist)[c(1:10),"TaxonName"]</pre>
```

split name
dissect_name(Easplist)

re-paste the two first words
dissect_name(Easplist, repaste=c(1:2))

Easplist-data List of vascular plants from East Africa

Description

Example of an incomplete taxonomic list including taxa recorded in East Africa.

Usage

Easplist

Format

An object of class taxlist.

Details

This list is a subset of the taxonomic list implemented in the database SWEA-Dataveg. Since this list is being complemented regarding stored vegetation plots, it is an incomplete list.

Source

African Plant Database, SWEA-Dataveg.

Examples

summary(Easplist)

Extract

Description

Quick access to slots taxonTraits and taxonRelations within taxlist objects.

Usage

```
## S4 method for signature 'taxlist'
x[i, j, drop = FALSE]
## S4 method for signature 'taxlist'
x$name
```

Arguments

х	Object of class taxlist.
i	Integer or logical vector used as index for access to taxon concepts, referring to the rows in slot 'taxonRelations'. These indices can be used to produce a object with a subset of taxon concepts. It is not recommended to use character values for this index.
j	Integer, logical or character vector used as index for access to variables in slot 'taxonTraits'. These indices can be used to reduce the number of variables in the mentioned slot.
drop	A logical value passed to Extract.
name	A symbol or character value for the method \$, corresponding to a variable either at slot 'taxonTraits' or slot 'taxonRelations'.

Value

The method \$ retrieves a vector, while [retrieves a subset of the input taxlist object.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>.

See Also

taxlist subset

Examples

```
## Statistics on life forms
summary(as.factor(Easplist$lf_behn_2018))
## First ten concepts in this list
summary(Easplist[1:10, ], "all")
```

get_children	Retrieve children or parents of taxon concepts
--------------	--

Description

Retrieve all children or all parents of a queried taxon concept.

Usage

```
get_children(taxlist, ConceptID, ...)
```

```
## S4 method for signature 'taxlist,numeric'
get_children(taxlist, ConceptID, ...)
```

```
## S4 method for signature 'taxlist,taxlist'
get_children(taxlist, ConceptID, ...)
```

get_parents(taxlist, ConceptID, ...)

```
## S4 method for signature 'taxlist,numeric'
get_parents(taxlist, ConceptID, ...)
```

```
## S4 method for signature 'taxlist,taxlist'
get_parents(taxlist, ConceptID, ...)
```

Arguments

taxlist	A taxlist object.
ConceptID	Concept IDs for selecting parents or children or a subset of taxlist.
	Further arguments passed among methods.

Details

This function produces subsets of taxlist objects including all children or parents of queried taxon concepts. Multiple concepts can be queried in these function. The argument ConceptID can be a vector of concept IDs or a subset of the input taxlist object.

Value

A taxlist object with a subset including requested concepts with children or parents.

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indented_list

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

Examples

```
## Subset with family Ebenaceae and children
Ebenaceae <- subset(Easplist, charmatch("Ebenaceae", TaxonName))
Ebenaceae <- get_children(Easplist, Ebenaceae)
summary(Ebenaceae)
summary(object=Ebenaceae, ConceptID="all", maxsum=100)
```

indented_list Print hierarchical structure in indented lists

Description

Print taxonomic hierarchies (ranks and parent-child relationships) from taxlist objects in an indented list.

Usage

```
indented_list(object, ...)
## S4 method for signature 'taxlist'
indented_list(
 object,
  filter,
  keep_children = TRUE,
  keep_parents = TRUE,
  rankless_as,
  indent = " ",
  lead_br = "",
  print = TRUE,
  author = TRUE,
  level = FALSE,
  synonyms = FALSE,
  syn_encl = c("= ", ""),
  secundum,
```

```
alphabetical = FALSE,
...
```

Arguments

object	A taxlist object containing taxonomic concepts.
	Further arguments (not used yet).
filter	A character value (optional) that will be matched with the taxon usage names to produce a subset of 'object'. Note that this filter will be also applied to synonyms, independent of the argument applied in parameter 'synonyms'.
keep_children	A logical value indicating whether children of matched concept should be in- cluded in the result.
keep_parents	A logical value indicating whether parents of matched concept should be in- cluded in the result.
rankless_as	A character vector indicating a level (taxonomic rank) to which rankless taxa may be set before doing the list.
indent	Symbol used for indentation. This symbol will be multiplied by the depth of the taxonomic rank. The default is a blank space. This can be also provided as a named vector, with a different indentation symbol for the respective taxonomic ranks.
lead_br	Optional line break symbol leading before the indentation. It may be required for r-markdown documents.
print	A logical value indicating whether the indented list should be printed in the console or not (default = TRUE).
author	A logical value indicating whether the author should be printed with the name (default = TRUE).
level	A logical value indicating whether the name of the level (taxonomic rank) should be included before the name or not (default = FALSE).
synonyms	A logical value indicating whether the synonyms should be included after accepted names or not (default = FALSE).
syn_encl	A character vector of length 2 including the symbols used to enclose synonyms. First value will be set before the synonyms and second value, after the synonyms.
secundum	A character value matching a name in slot 'taxonViews', which will be printed as secundum (taxon view). It is not printed by default.
alphabetical	A logical value indicating whether taxa may be sorted by names or by IDs. The default is FALSE, thus taxa are sorted by IDs. Note that argument TRUE may not work properly if the object contains homonymous taxa.

Value

If 'print = TRUE', the indented list is printed in the console. The result, which is a data frame with the elements used to format the names, can be also assigned to an object.

levels

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

Examples

```
## Show taxonomy of papyrus
indented_list(Easplist, "papyrus")
```

```
## Include synonyms and taxon views
indented_list(Easplist, "papyrus", level = TRUE, synonyms = TRUE,
    secundum = "secundum")
```

levels

Set and retrieves hierarchical levels

Description

Taxonomic hierarchies can be set as levels in taxlist objects, ordered from lower to higher levels.

Add taxonomic levels for specific taxon concepts in a taxlist object. Also changes in concept circumscription may implicate changes in its taxonomic hierarchy.

Usage

```
## S4 method for signature 'taxlist'
levels(x)
```

S4 replacement method for signature 'taxlist'
levels(x) <- value</pre>

Arguments

Х	A taxlist object.
value	A character vector with replacement values for levels o x.

Details

Taxonomic levels will be handled as factors in the taxlist objects. Those levels are useful for creating subsets of related groups (e.g. by functions get_children() or get_parents()).

Levels in combination to parent-child relationships will be further used for checking consistency of taxonomic lists.

A replacement method of the form levels(x) < -value it is also implemented.

Value

A character vector or a taxlist object with added or modified taxonomic levels.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

Examples

```
## Get levels of species list
taxlist::levels(Easplist)
## Add aggregate as new taxonomic level
levels(Easplist) <- c("form", "variety", "subspecies", "species",
        "complex", "aggregate", "genus", "family")
summary(Easplist)
```

match_names

```
Search matchings between character and taxlist objects
```

Description

Names provided in a character vector will be compared with names stored in slot taxonNames within an object of class taxlist by using the function stringsim().

Usage

```
match_names(x, object, ...)
## S4 method for signature 'character,character'
match_names(x, object, best = 5, clean = TRUE, decreasing = TRUE, ...)
## S4 method for signature 'character,taxlist'
match_names(
  х,
  object,
  clean = TRUE,
  output = "data.frame",
  best = 5,
  show_concepts = FALSE,
  accepted_only = FALSE,
 method = "lcs",
  decreasing,
  . . .
)
```

Arguments

Х	A character vector with names to be compared.
object	An object of class taxlist to be compared with.

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best	Integer value indicating how many from the best matches have to be displayed (only working for output="list").
clean	Logical value, whether leading, tailing and double blanks should be deleted from x.
decreasing	Logical value indicating whether retrieved names should be sorted by decreasing or increasing similarity value. In the character method, the sorting corresponds to similarities between the queried value and the reference vector (argument object). In the taxlist method using output="data.frame", the order corre- sponds to the similarity of the best match (by default, no sorting is done). This argument is passed to order().
output	Character value indicating the type of output. Alternative values are "list" (taxon concepts ID's sorted by similarity for each queried name) or "data.frame" (a table including the best match for every queried name).
show_concepts	Logical value, whether respective concepts should be displayed in output or not.
accepted_only	Logical value, whether only accepted names should be matched or all usage names (including synonyms).
method,	Further arguments passed to stringsim().

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

See Also

stringsim()

Examples

```
## Names to be compared
species <- c("Cperus papyrus", "Typha australis", "Luke skywalker")
## Comparing character vectors
match_names("Cyperus paper", species)
## Retrieve taxon usage names
match_names(species, Easplist)
## Display accepted names in output
match_names(x=species, object=Easplist, show_concepts=TRUE)
```

merge_taxa

Description

Merge taxon concepts form a taxlist object into single ones.

Usage

```
merge_taxa(object, concepts, level, ...)
## S4 method for signature 'taxlist,numeric,missing'
merge_taxa(object, concepts, print_output = FALSE, ...)
## S4 method for signature 'taxlist,missing,character'
merge_taxa(object, concepts, level, ...)
change_concept(taxlist, UsageID) <- value
## S4 replacement method for signature 'taxlist'</pre>
```

change_concept(taxlist, UsageID) <- value</pre>

Arguments

object, taxlist

	Object of class taxlist.
concepts	Numeric (integer) vector including taxon concepts to be merged.
level	Character vector indicating the lowest level for merging.
	Further arguments to be passed to or from other methods.
print_output	Logical value indicating whether the merged concept should be displayed in the console.
UsageID	Numeric vector with taxon usage IDs to be changed from concept.
value	Numeric vector with taxon concept IDs to be assigned to the names.

Details

Taxon concepts indicated in argument concepts will be merged into a single concept. The new concept inherits the ID and respective attributes from slots taxonRelations and taxonTraits from the first taxon concept indicated in argument concepts.

For convenience the resulting concept can be displayed by setting print_output=TRUE but only when using argument concepts.

An alternative application of this function is implemented through the argument level, where all lower rank taxa will be merged to the indicated level or higher (if parent of merged taxa are at a higher rank).

print_name

Value

An object of class taxlist.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

Examples

```
## Merge Cyperus papyrus and Cyperus dives
summary(Easplist, c(206, 197))
Easplist <- merge_taxa(object=Easplist, concepts=c(206, 197),
    print_output=TRUE)
## Move the name Typha aethiopica to concept 573 (T. latifolia)
change_concept(Easplist, 53130) <- 573
summary(Easplist, c(50105,573))
```

print_name

Format usage names for publications

Description

When writing on bio-diversity, usage names could be automatically inserted in documents including the typical italic format for different elements of a scientific name. The function print_name can be applied either in markdown documents or for graphics.

Usage

```
print_name(object, id, ...)
## S4 method for signature 'taxlist,numeric'
print_name(
    object,
    id,
    concept = TRUE,
    second_mention = FALSE,
    include_author = TRUE,
    secundum,
    style = "markdown",
    ...
)
```

Arguments

object	An object of class taxlist.
id	Integer containing either a concept or a name ID.
	Further arguments passed among methods.
concept	Logical value, whether id corresponds to a concept ID or a taxon usage name ID.
second_mention	Logical value, whether the genus name should be abbreviated or not.
include_author	Logical value, whether authors of the name should be mentioned or not.
secundum	Character value indicating the column in slot taxonViews that will be mentioned as <i>secundum</i> (according to).
style	Character value indicating the alternative format for italics (at the moment only markdown and html implemented).

Details

In Rmarkdown documents use *Cyperus papyrus* L. for inserting a formatted a species name.

Value

A character value including format to italic font.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

See Also

ape::mixedFontLabel().

Examples

```
summary(Easplist, 363, secundum="secundum")
## Empty plot
plot(x=NA, xlim=c(0,5), ylim=c(7,1), bty="n", xaxt="n", xlab="",
ylab="options")
## Accepted name with author
text(x=0, y=1, labels=print_name(Easplist, 363, style="expression"), pos=4)
## Including taxon view
text(x=0, y=2, labels=print_name(Easplist, 363, style="expression",
    secundum="secundum"), pos=4)
## Second mention in text
text(x=0, y=3, labels=print_name(Easplist, 363, style="expression",
    second_mention=TRUE), pos=4)
```

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replace_x

replace_x

Data manipulation.

Description

This is a series of functions designed for a fast coding of replacements both, as internal functions and in workflows dealing with information stored in vectors and data frames. Such functions are especially useful when handling with functional traits stored in taxlist objects.

replace_x() is used to exchange values in vectors. replace_idx() changes values in vectors by matching indices or conditions. The function replace_na() works in the same way as replace_idx() but will only insert values in empty elements (NAs).

The function insert_rows() will add rows and columns at the same time. This function will be used when a new table is appended to another but sharing only part of the columns.

Usage

```
replace_x(x, old, new)
replace_idx(x, idx1 = x, idx2 = idx1, new)
replace_na(x, idx1, idx2 = idx1, new)
insert_rows(x, y)
```

Arguments

х	A vector to be modified. In the case of insert_rows(), x is a data frame.
old	A vector with values to be replaced by replace_x() in a vector.
new	A vector containing values to be inserted, either comparing values or using in- dices.
idx1, idx2	Indices applied for value replacements to match x with new, respectively. If idx^2 is not provided, it will be assumed as equivalent to idx^1 .
У	A data frame including rows (and columns) to be inserted in x.

A vector or data frame with the modified values.

Author(s)

Miguel Alvarez.

Examples

```
## Replace values in vector
replace_x(x=letters, old=c("b", "p", "f"), new=c("bee", "pork", "fungus"))
## Replace values using indices
replace_idx(x=letters, idx1=1:length(letters), idx2=c(2, 7, 17),
   new=c("second", "seventh", "seventeenth"))
## Replace values if they are NAs
letters[2] <- NA
replace_na(x=letters, idx1=1:length(letters), idx2=c(1:3),
    new=c("alpha", "beta", "zeta"))
## The same applications but this time for functional traits
summary(as.factor(Easplist$lf_behn_2018))
# Merge annuals
Easplist@taxonTraits$lifeform <- replace_x(</pre>
   x=Easplist@taxonTraits$lf_behn_2018,
   old=c("obligate_annual", "facultative_annual"),
   new=c("annual", "annual"))
summary(as.factor(Easplist$lifeform))
# The same effect
Easplist@taxonTraits$lifeform <- replace_idx(</pre>
   x=Easplist@taxonTraits$lf_behn_2018,
    idx1=grepl("annual", Easplist@taxonTraits$lf_behn_2018),
   idx2=TRUE,
   new="annual")
summary(as.factor(Easplist$lifeform))
## Merge data frames including new columns
data(iris)
iris$Species <- paste(iris$Species)</pre>
new_iris <- data.frame(Species=rep("humilis", 2), Height=c(15,20),</pre>
    stringsAsFactors=FALSE)
insert_rows(iris, new_iris)
```

subset

Description

Subset of taxlist objects will be done applying either logical operations or pattern matchings. Subsets can be referred to information contained either in the slot taxonNames, taxonRelations or taxonTraits.

Usage

```
## S4 method for signature 'taxlist'
subset(
    x,
    subset,
    slot = "names",
    keep_children = FALSE,
    keep_parents = FALSE,
    ...
)
```

Arguments

Х	Object of class taxlist.
subset	Logical vector or logical operation to apply as subset.
slot	Character value indicating the slot to be used for the subset.
keep_children	Logical value applied to hierarchical structures.
keep_parents	Logical value applied to hierarchical structures.
	Further arguments to be passed to or from other methods.

Details

The argument subset will be applied to the slot specified in argument slot. This argument also allows partial matchings.

Arguments keep_children and keep_parents are applied to objects including parent-child relationships. When those arguments are set as FALSE (the default), children or parents of selected taxon concepts will not be included in the subset.

Be aware that subset() won't work properly inside of function definitions.

Value

An object of class taxlist.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

Examples

```
summary(as.factor(Easplist$lf_behn_2018))
```

```
summary
```

Print overviews for taxlist Objects and their content

Description

A method to display either an overview of the content of taxlist objects or an overview of selected taxa.

Usage

```
## S4 method for signature 'taxlist'
summary(
   object,
   ConceptID,
   units = "Kb",
   check_validity = TRUE,
   display = "both",
   maxsum = 5,
   secundum = NULL,
   ...
)
## S4 method for signature 'taxlist'
show(object)
## S4 method for signature 'taxlist'
```

print(x, ...)

Arguments

object, x	A taxlist object.
ConceptID	IDs of concepts to be displayed in the summary.
units	Character value indicating the units shown in the object's allocated space.
check_validity	Logical value indicating whether the validity of object should be checked or not.
display	Character value indicating the field to be displayed (see details).
maxsum	Integer indicating the maximum number of displayed taxa.

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summary

secundum	A character value indicating the column from slottaxonViews to be displayed
	in the summary.
	Further arguments passed to or from another methods.

Details

A general overview indicating number of names, concepts and taxon views included in taxlist objects. If argument ConceptID is a vector with concept IDs or names to be matched by grepl(), then a display of all names included in each concept will be produced. Alternative you can use taxon="all" in order to get the listing of names for all concepts included in the object (truncated to the input number of maxsum).

For summaries applied to concepts, there are three alternative displays of names using the argument display. Use display="name" to show the value TaxonName, display="author" to show the value AuthorName or display="both" to show both values. Such values are taken from slot taxonNames.

For big objects it will be recommended to set units="Mb" (see also object.size() for further alternatives).

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

See Also

taxlist

Examples

```
## summary of the object
summary(Easplist, units="Mb")
```

the same output
summary(Easplist)
show(Easplist)
print(Easplist)
Easplist

summary for two taxa summary(Easplist, c(51128,51140))

```
## summary for a name
summary(Easplist, "Acmella")
```

summary for the first 10 taxa
summary(object=Easplist, ConceptID="all", maxsum=10)

```
tax2traits
```

Description

Taxonomic classification can be included in taxlist objects within the information provided at slot taxonRelations. Nevertheless, for statistical analyses it may be more convenient to insert such information in the slot taxonTraits.

Usage

```
tax2traits(object, ...)
```

```
## S4 method for signature 'taxlist'
tax2traits(object, get_names = FALSE, ...)
```

Arguments

object	An object of class taxlist.
	Further arguments to be passed among methods.
get_names	Logical value indicating whether taxon names should be retrieved instead of taxon IDs.

Details

This function can only be applied to objects containing parent-child relationships and information on taxonomic levels.

Value

An object of class taxlist with taxonomy added as traits.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>.

Examples

Description

Class for taxonomic lists including synonyms, hierarchical ranks, parent-child relationships, taxon views and taxon traits.

Note that each taxon becomes an identifier, represented by the column **TaxonConceptID** in the slot **taxonRelations**, analogous to a primary key in a relational database. This identifier is restricted to an integer in taxlist and is specific for the object.

In the same way, each taxon usage name has an identifier in the column **TaxonUsageID**, slot **tax-onNames**. The column **ViewID** in slot **taxonViews** is the identifier of the taxon view.

Slots

taxonNames (data.frame) Table of taxon usage names (accepted names and synonyms).

taxonRelations (data.frame) Relations between concepts, accepted names, basionyms, parents and hierarchical level.

taxonTraits Table of taxon traits.

taxonViews References used to determine the respective concept circumscription.

Author(s)

Miguel Alvarez

References

Alvarez M, Luebert F (2018). The taxlist package: managing plant taxonomic lists in R. *Biodiversity Data Journal* 6: e23635. doi: 10.3897/bdj.6.e23635

Examples

```
library(taxlist)
```

```
showClass("taxlist")
```

```
## Create an empty object
Splist <- new("taxlist")</pre>
```

taxon_names

Description

The slot taxonNames in taxlist objects contains taxon usage names for the respective taxon. These functions assist on the access and modification of entries for names.

Usage

```
taxon_names(taxlist, ...)
## S4 method for signature 'taxlist'
taxon_names(taxlist, ...)
taxon_names(taxlist) <- value
## S4 replacement method for signature 'taxlist,data.frame'
taxon_names(taxlist) <- value
add_synonym(taxlist, ConceptID, ...)
## S4 method for signature 'taxlist'
add_synonym(taxlist, ConceptID, TaxonName, AuthorName, ...)
update_name(taxlist, UsageID, ...)
## S4 method for signature 'taxlist,numeric'
update_name(taxlist, UsageID, ...)
## S4 method for signature 'taxlist,numeric'
update_name(taxlist, UsageID, ...)
## S4 method for signature 'taxlist,numeric'
update_name(taxlist, UsageID, ...)
## S4 method for signature 'taxlist,numeric'
delete_name(taxlist, UsageID, ...)</pre>
```

Arguments

taxlist	A taxlist object to be modified.	
	Further arguments passed among methods. In update_name are vectors includ- ing the variables to be updated for the respective taxon usage ID.	
value	A data frame used as new slot taxonNames in taxlist.	
ConceptID	Numeric vector indicating the concept ID to which the synonyms will be added.	
TaxonName, AuthorName		
	Character values used for the new names (synonyms).	
UsageID	Numeric vector indicating the taxon usage IDs to be updated.	

Details

The replacement method taxon_names<- is a quick alternative to include names in empty taxlist objects.

The function add_synonym() works only for adding names to existing taxon concepts. For adding new taxon concepts as well you should use add_concept().

Value

A data frame or, in the case of the replacement method, a taxlist object with modified slot taxonNames.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

See Also

taxlist

Examples

```
## Display of slot 'taxonNames'
Euclea <- subset(x=Easplist, subset=charmatch("Euclea", TaxonName),</pre>
    slot="names", keep_children=TRUE)
Euclea
taxon_names(Euclea)
## Insert a synonym to Diospyros scabra
summary(Easplist, "Diospyros scabra")
Easplist <- add_synonym(taxlist=Easplist, ConceptID=51793,</pre>
    TaxonName="Maba scabra", AuthorName="Chiov.")
summary(Easplist, "Diospyros scabra")
## Delete a synonym of Launaea cornuta
summary(Easplist, "Launaea cornuta")
Easplist <- delete_name(Easplist, 53821)</pre>
summary(Easplist, "Launaea cornuta")
## Hypothetical correction in author name in Launaea cornuta
Easplist <- update_name(taxlist=Easplist, UsageID=355, AuthorName="L.")</pre>
summary(Easplist, "Launaea cornuta")
```

taxon_relations Retrieve or replace slot taxonRelations in taxlist objects

Description

Retrieve the content of slot taxonRelations from a taxlist object or replace it by a new data frame.

Usage

```
taxon_relations(taxlist, ...)
## S4 method for signature 'taxlist'
taxon_relations(taxlist, ...)
taxon_relations(taxlist) <- value
## S4 replacement method for signature 'taxlist,data.frame'
taxon_relations(taxlist) <- value
add_concept(taxlist, TaxonName, ...)
## S4 method for signature 'taxlist,character'
add_concept(taxlist, TaxonName, Level, ...)
## S4 method for signature 'taxlist,taxlist'
add_concept(taxlist, TaxonName, insert_view, ...)
update_concept(taxlist, ConceptID, ...)
## S4 method for signature 'taxlist,numeric'
update_concept(taxlist, ConceptID, ...)</pre>
```

Arguments

taxlist	A taxlist object.
	Further arguments passed among methods.
value	A data.frame object to be set as slot taxonRelations.
TaxonName	Character vector with the accepted name for the new taxon concepts.
Level	Character vector indicating the level of the concept in the list.
insert_view	A numeric (integer) vector, indicating the views to be inserted in taxlist or the value TRUE (see details).
ConceptID	Concept IDs to be updated.

Details

The replacement method taxon_relations<- should be only used when constructing taxlist objects from an empty one (prototype).

New concepts should be first added to a taxlist object using their respective accepted names. Synonyms can be further provided using the function add_synonym().

Additional named vectors can be provided to be included in slot taxonNames, in the cases where those variables already exist, otherwise they will be ignored.

It is recommended also to provide a concept view as ViewID (see taxon_views()). For adding a new view, use add_view().

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taxon_relations

Value

An object of class taxlist with added names and concepts.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

See Also

taxlist

Examples

```
## Subset for the genus Euclea and display of slot 'taxonNames'
Euclea <- subset(x=Easplist, subset=charmatch("Euclea", TaxonName),</pre>
    slot="names")
Euclea <- get_children(Easplist, Euclea)</pre>
Euclea
taxon_relations(Euclea)
## Subset with family Ebenaceae and children
Ebenaceae <- subset(Easplist, charmatch("Ebenaceae", TaxonName))</pre>
Ebenaceae <- get_children(Easplist, Ebenaceae)</pre>
Ebenaceae
summary(object=Ebenaceae, ConceptID="all", maxsum=100)
## Adding a new concept
Ebenaceae <- add_concept(taxlist=Ebenaceae, TaxonName="Euclea acutifolia",</pre>
    AuthorName="E. Mey. ex A. DC.", Level="species", Parent=55707, ViewID=1)
## A summary again
Ebenaceae
summary(Ebenaceae, "all", maxsum=100)
## Display two Typha species
summary(Easplist, c("Typha domingensis","Typha latifolia"))
## Update a concept
summary(Easplist, "Corchorus olitorius")
Easplist <- update_concept(taxlist=Easplist, ConceptID=155,</pre>
    Level="subspecies")
summary(Easplist, "Corchorus olitorius")
```

```
taxon_traits
```

Description

The slot taxonTraits in taxlist objects contains attributes of taxon concepts (e.g. functional traits). These functions are suitable for replacing, retrieving and appending trait information in taxonomic lists.

Usage

```
taxon_traits(taxlist, ...)
## S4 method for signature 'taxlist'
taxon_traits(taxlist, ...)
taxon_traits(taxlist) <- value
## S4 replacement method for signature 'taxlist,data.frame'
taxon_traits(taxlist) <- value
update_trait(taxlist, ConceptID, ...)
## S4 method for signature 'taxlist,numeric'
update_trait(taxlist, ConceptID, ...)</pre>
```

Arguments

taxlist	A taxlist object.
	Further arguments to be passed among methods.
value	Data frame to be set as slot taxonTraits.
ConceptID	A numeric vector with the respective taxon concept IDs.

Details

Taxon traits are contained in a data frame at the slot taxonTraits in taxlist objects. To optimise space, this data frame contain only entries for those concepts with information, while taxa with no information are skipped from this table. Thus appending new variables may also have to include new rows in this slot, which is automatically carried out by this function.

The replacement method taxon_traits<- should be only used when constructing taxlist objects from an empty one.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

taxon_views

See Also

taxlist.

Examples

```
head(taxon_traits(Easplist))
```

```
accepted_name(taxlist=Easplist, ConceptID=355, show_traits=TRUE)
```

```
taxon_views
```

```
Management of concept views in taxonomic lists.
```

Description

Retrieve or replace slot taxonViews in an object of class taxlist

Usage

```
taxon_views(taxlist, ...)
## S4 method for signature 'taxlist'
taxon_views(taxlist, ...)
taxon_views(taxlist) <- value
## S4 replacement method for signature 'taxlist,data.frame'
taxon_views(taxlist) <- value
add_view(taxlist, ...)
## S4 method for signature 'taxlist'
add_view(taxlist, ...)</pre>
```

Arguments

taxlist	A taxlist object.
	Further arguments to be passed among methods.
value	An object of class data.frame containing the references used to define the circumscription of taxon concepts included in taxlist.

Details

Taxon views indicate in taxlist objects the references determining the circumscription of the respective taxon concepts. When adding a new concept (see add_concept()), the respective reference may not yet occur in the input taxlist object.

The term taxon view was introduced by **Zhong et al. (1996)** and corresponds to the reference used for the definition of a concept.

This function retrieves the slot taxonViews from objects of the class taxlist.

The replacement method taxon_views<- replaces the whole content of slot taxonViews and it is only recommended to use when constructing a new taxlist object from an empty prototype.

Value

An object of class taxlist with added views.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

References

Zhong Y, Jung S, Pramanik S, Beaman JH (1996). Data model and comparison and query methods for interacting classifications in a taxonomic database. *Taxon* 45: 223–241. doi: 10.1093/bioinformatics/15.2.149

See Also

taxlist

Examples

```
## See existing views
taxon_views(Easplist)
## Add a new view
Easplist <- add_view(taxlist=Easplist, secundum="Beentje et al. (1952)",
Title="Flora of Tropical East Africa",
URL="http://www.kew.org/science/directory/projects/FloraTropEAfrica.html")</pre>
```

taxon_views(Easplist)

tnrs

Description

Methods of taxize::tnrs() for taxlist objects.

Usage

```
tnrs(query, ...)
```

```
## S4 method for signature 'character'
tnrs(query, ...)
```

S4 method for signature 'taxlist'
tnrs(query, min_score = 0.8, source = "iPlant_TNRS", ...)

Arguments

query	Either a character vector or a taxlist object with names to search.
	Further arguments passed to taxize::tnrs().
min_score	Minimum value of score for considering accepted names as suggested by the output.
source	Source database.

Details

This function checks for matching of taxon names in taxlist objects with the Taxonomic Name Resolution Service (TNRS). Misspelled names as well as author names will be replaced in the the new object and new accepted names will be inserted.

A method for character vectors is defined for the original function.

Value

A data frame or an object of class taxlist.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

See Also

taxize::tnrs()

```
tv2taxlist
```

Description

Importing species lists from Turboveg https://www.synbiosys.alterra.nl/turboveg/ databases into an object of class taxlist.

Usage

tv2taxlist(taxlist, tv_home = tv.home())

Arguments

taxlist	The name of a species list in Turboveg as character value.
tv_home	Character value indicating the path to the main Turboveg folder.

Details

This function imports species lists using the function read.dbf(). When available, also taxon traits will be imported into the output object (usually the file **ecodbase.dbf**). During import of taxon traits, duplicated entries for a same concept will be discarded as well as entries for non-existing concepts.

By default tv_home will be set by the function $tv_home()$ from the package vegdata-package.

By default, the name of the database will be set as concept view for all concepts included in the species list. If this is not correct, consider setting it manually by using the functions taxon_views() and add_view().

Value

An object of class taxlist.

Author(s)

Miguel Alvarez <kamapu78@gmail.com>

See Also

taxlist

Examples

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