Package ‘arakno’

October 12, 2022

Title ARAchnid KNowledge Online
Version 1.3.0
Date 2022-08-23
Description Allows the user to connect with the World Spider Cata-
logue (WSC; <https://wsc.nmbe.ch/>) and the World Spi-
der Trait (WST; <https://spidertraits.sci.muni.cz/>) databases. Also performs sev-
eral basic functions such as checking names validity, retrieving coordi-
nate data from the Global Biodiversity Information Facil-
itly (GBIF; <https://www.gbif.org/), and mapping.
Depends R (>= 3.5.0)
Imports ape, graphics, httr, jsonlite, phytools, rgbif, rworldmap,
rworldxtra, stats, utils
Encoding UTF-8
License GPL-3
LazyData true
RoxygenNote 7.2.1
NeedsCompilation no
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Repository CRAN
Date/Publication 2022-08-23 19:30:02 UTC

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authors

Get species authors from WSC.

Description

Get species authority from the World Spider Catalogue.

Usage

authors(tax, order = FALSE)

Arguments

tax A taxon name or vector with taxa names.
order Order taxa alphabetically or keep as in tax.

Details

This function will get species authorities from the World Spider Catalogue (2022). Higher taxa will be converted to species names.

Value

A data.frame with species and authority names.

References


Examples

```r
## Not run:
authors("Amphiledorus")
authors(tax = c("Iberesia machadoi", "Nemesia bacelarae", "Amphiledorus ungoliantae"), order = TRUE)

## End(Not run)
```
buildtree

Create phylogenetic tree.

Description
Create a phylogenetic tree based on the backbone from Macias-Hernandez et al. (2020) and the species taxonomical hierarchy.

Usage
buildtree(tax, update = FALSE)

Arguments
tax
A taxon name or vector with taxa names. Should be in the general form "Family_sp" or "Genus speciesname", with family or genus name plus anything to uniquely identify the species separated by "_" or " ".

update
Whether to update the taxonomy of the backbone tree according to the WSC (2022).

Details
Based on the backbone phylogeny of Macias-Hernandez et al. (2020). All species in tax present in the backbone are included in the output tree. If the species is not in the backbone, or if only family or genus are known, species are inserted at the level of the most recent common ancestor of confamiliar or congenerics respectively. If only one congeneric or confamiliar are in the backbone, the species is inserted at half the length of the corresponding edge.

Value
A phylo object with a phylogenetic tree for the community.

References


Examples
## Not run:
spp = c("Atypus affinis", "Tenuiphantes tenuis", "Zodarion sp1", "Araneus diadematus")
spp = c(spp, "Zodarion sp2", "Atypus_nsp", "Nemesia ungoliant", "Linyphiidae sp1")
spp = c(spp, "Zoropsis spinimana", "Pardosa sp1", "Pardosa acorensis", "Liphistius nsp")
tree = buildtree(spp)
plot(tree)
## End(Not run)
checknames  

Check taxa names in WSC.

Description

Check taxa names against the World Spider Catalogue.

Usage

checknames(tax, full = FALSE, order = FALSE)

Arguments

tax  A taxon name or vector with taxa names.
full  returns the full list of names.
order  Order taxa alphabetically or keep as in tax.

Details

This function will check if all species, genera and family names in tax are updated according to the World Spider Catalogue (2022). If not, it returns a matrix with nomenclature changes, valid synonyms or possible misspellings using fuzzy matching (Levenshtein edit distance).

Value

If any mismatches, a matrix with taxa not found in WSC or, if full = TRUE, the full list of names.

References


Examples

```r
## Not run:
tax = c("Nemesia", "Nemesia brauni", "Iberesia machadoi", "Nemesia bacelari")
checknames(tax)
checknames(tax, full = TRUE, order = TRUE)
## End(Not run)
```
countries

Get taxon countries from WSC.

Description
Get countries of taxa from the World Spider Catalogue textual descriptions.

Usage
countries(tax)

Arguments

tax A taxon name or vector with taxa names.

Details
Countries based on the interpretation of the textual descriptions available at the World Spider Catalogue (2022). These might be only approximations to country level and should be taken with caution.

Value
A vector with country ISO codes. Family and genera names will be converted to species.

References

Examples

```r
## Not run:
countries("Iberesia machadoi")
countries(c("Iberesia machadoi", "Nemesia"))

## End(Not run)
```
distribution

Get species distributions from WSC.

Description

Get species distribution from the World Spider Catalogue.

Usage

distribution(tax, order = FALSE)

Arguments

tax A taxon name or vector with taxa names.
order Order taxa alphabetically or keep as in tax.

Details

This function will get species distributions from the World Spider Catalogue (2022).

Value

A data.frame with species and distribution. Family and genera names will be converted to species.

References


Examples

## Not run:
distribution("Nemesia")
distribution(tax = c("Iberesia machadoi", "Amphildorus ungoliantea"), order = TRUE)

## End(Not run)
endemics

Description

Get endemic species in any country or region from the World Spider Catalogue textual descriptions.

Usage

dermics(country)

Arguments

country 

The country/region name or ISO3 code.

Details

Species list based on the interpretation of the textual descriptions available at the World Spider Catalogue (2022). These might be only approximations to country level and should be taken with caution.

Value

A vector with species names.

References


Examples

## Not run:
endemics("Portugal")
endemics("Azores")
endemics("FIN")

## End(Not run)
globalTree

*Global spider backbone tree.*

Description

From Macias-Hernandez et al. (2020) with nomenclature updated.

Usage

```
data(globalTree)
```

Format

A phylo object with 132 families and 800+ genera, 1400+ species.

lsid

*Get species LSID from WSC.*

Description

Get species LSID from the World Spider Catalogue.

Usage

```
lsid(tax, order = FALSE)
```

Arguments

- `tax` A taxon name or vector with taxa names.
- `order` Order taxa names alphabetically or keep as in tax.

Details

This function will get species LSID from the World Spider Catalogue (2022). Family and genera names will be converted to species.

Value

A data.frame with species and LSID.

References

### Not run:
```r
lsid("Anapistula")
lsid(tax = c("Iberesia machadoi", "Nemesia bacelarae", "Amphiledorus ungoliantea"), order = TRUE)
```
## End(Not run)

---

**map**  
*Map species ranges.*

---

**Description**

Maps species range according to the World Spider Catalogue and records according to GBIF and the World Spider Trait database.

**Usage**

```r
map(
  tax,  
  countries = TRUE,  
  records = TRUE,  
  hires = FALSE,  
  zoom = FALSE,  
  order = FALSE,  
  verbose = TRUE
)
```

**Arguments**

- `tax` A taxon name or vector with taxa names.
- `countries` Maps countries according to WSC.
- `records` Maps records according to GBIF and WST.
- `hires` Provides high resolution maps. Beware it might take longer to render.
- `zoom` If records is TRUE, the map will be zoomed to the region with records.
- `order` Order taxa names alphabetically or keep as in tax.
- `verbose` Display information as data are retrieved.

**Details**

Countries based on the interpretation of the textual descriptions available at the World Spider Catalogue (2022). These might be only approximations to country level and should be taken with caution.

**Value**

A world map with countries and records highlighted.
## records

Get coordinate data from GBIF and WST.

### Description

Downloads coordinate data from records in GBIF and the World Spider Trait database.

### Usage

```r
records(tax, order = FALSE, verbose = TRUE)
```

### Arguments

- **tax**: A taxon name or vector with taxa names.
- **order**: Order taxa names alphabetically or keep as in tax.
- **verbose**: Display information as data are retrieved.

### Details

Outputs non-duplicate records with geographical (long, lat) coordinates. As always when using data from multiple sources the user should be careful and check if records "make sense" before using them.

### Value

A data.frame with species name, longitude, latitude, source database and reference.

### References


species

Examples

## Not run:
records("Pardosa hyperborea")
records(tax = c("Pardosa hyperborea", "Anapistula"), order = TRUE)

## End(Not run)

---

species  Get species from higher taxa.

Description

Get species within given families or genera from the World Spider Catalogue.

Usage

species(tax, order = FALSE)

Arguments

tax  A taxon name or vector with taxa names.
order  Order species names alphabetically.

Details

This function will get all species currently listed for given families or genera from the World Spider Catalogue (2022).

Value

A vector with species names.

References


Examples

## Not run:
species("Amphiledorus")
species(tax = c("Amphiledorus", "Nemesiidae"), order = TRUE)

## End(Not run)
taxonomy

Get taxonomy from species.

Description

Get species sub/infraclass, family and genus from the World Spider Catalogue.

Usage

taxonomy(tax, check = FALSE, aut = FALSE, id = FALSE, order = FALSE)

Arguments

tax A taxon name or vector with taxa names.
check species names should be replaced by possible matches in the WSC if outdated.
aut add species authorities.
id the lsid should be returned.
order Order taxa names alphabetically or keep as in tax.

Details

This function will get species sub/infraclass, family and genus from the World Spider Catalogue (2022). Optionally, it will correct the species names (using function checknames) and provide the lsid and authors from the WSC (using functions lsid and authors).

Value

A data.frame with species and taxonomy.

References


Examples

## Not run:
taxonomy("Symphytognathidae", order = TRUE, aut = TRUE)
taxonomy(c("Nemesia machadoi", "Nemesia bacelari"), check = TRUE, aut = TRUE, id = TRUE)

## End(Not run)
traits

Get trait data from WST.

Description

Downloads the most recent data from the World Spider Trait database.

Usage

traits(
  tax,
  trait = NULL,
  sex = NULL,
  life = NULL,
  country = NULL,
  habitat = NULL,
  user = "",
  key = "",
  order = FALSE,
  verbose = TRUE
)

Arguments

tax A taxon name or vector with taxa names.
trait A vector with required trait(s) as abbreviations. Valid values can be found at:
https://spidertraits.sci.muni.cz/traits
sex A vector with required sex(es).
life A vector with required life stage(s).
country A vector with required country(ies) ISO3 code(s).
habitat A vector with required habitat(s).
user To obtain restricted data get a user name from https://spidertraits.sci.muni.cz/api.
key To obtain restricted data get an api key from https://spidertraits.sci.muni.cz/api.
order Order taxa names alphabetically or keep as in tax.
verbose Display information as data are retrieved.

Details

The World Spider Trait database (Pekar et al. 2021) has been designed to contain trait data in
a broad sense, from morphological traits to ecological characteristics, ecophysiology, behavioural
habits, and more (Lowe et al. 2020). This function will download everything available for the taxa
given, possibly filtered to the traits given in parameter trait. Some data might be restricted access,
in which case a user name and api key are needed (https://spidertraits.sci.muni.cz/api), otherwise
the value will show as NA.
Value

A matrix with trait data.

References


Examples

```r
## Not run:
traits("Atypus affinis")
traits("Atypus", order = TRUE)
traits("Atypidae", country = c("PRT", "CZE"), order = TRUE)
traits(c("Zodarion costapratae", "Zodarion alacre"))
traits(c("Iberesia machadoi", "Zodarion costapratae"), trait = c("balo", "bole"))
## End(Not run)
```

Description

Downloads the most recent data from the World Spider Catalogue.

Usage

```r
wsc()
```

Details

The World Spider Catalog (2022) lists all currently valid species of spiders, from Clerck to date. Updated daily.

Value

A matrix with all current species names and distribution. This should be used for other functions using wsc data.
**wscmap**

**References**


**Examples**

```r
## Not run:
wsc()

## End(Not run)
```

---

| wscmap | Matrix matching WSC and ISO3 country codes. |

**Description**

A dataset that links species distribution descriptions with the map using the ISO3 code

**Usage**

```r
data(wscmap)
```

**Format**

A matrix with regions and corresponding ISO3 codes.
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