# taxize: taxonomic search and retrieval in R



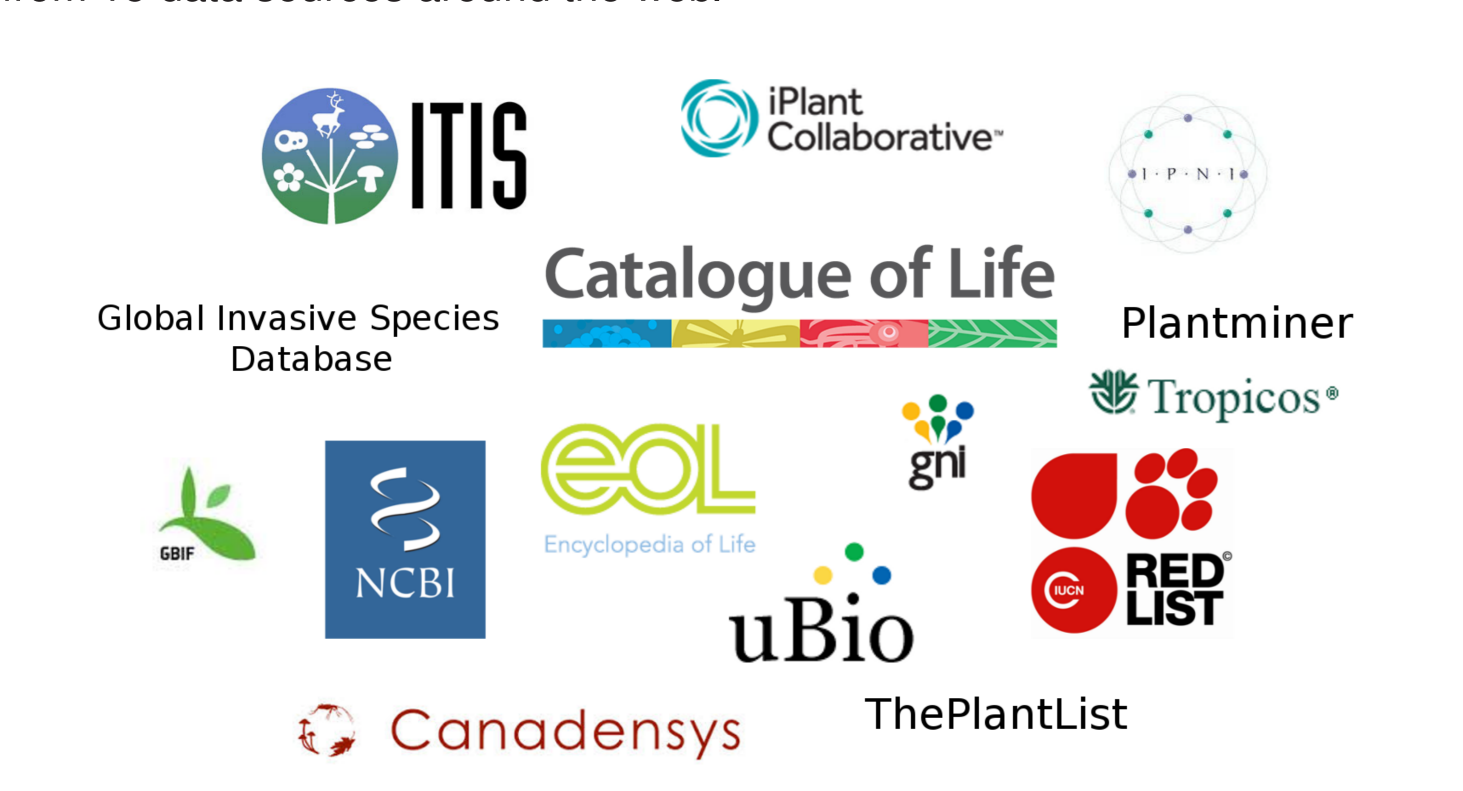
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# Summary

Taxize is a R package that provides an interface to various taxonomic data sources around the web<sup>1</sup>. Data cleaning steps, like fixing taxonomic names, aggregating data to a specific taxonomic level, resolving ambiguous taxa or matching tables with different taxonomic resolution are crucial steps before a statistical analysis<sup>2</sup>. The functionality of taxize simplifies these steps and eases handling of taxonomic data in R.

#### **Data Sources**

Taxize currently provides simple and programmatic access to taxonomic data from 15 data sources around the web.



## **Features**

#### Resolve taxonomic names

We often have a list of species names and we want to know

- a) if we have the most up-to-date names,
- b) if our names are spelled correctly,
- c) and the scientific name for a common name.

Taxize provides an interface to the EOL Global Names Resolver and Taxonomic Name Resolution Service, e.g.

```
gnr_resolve('Baetis roodani')
     submitted_name
                     matched_name
## 1 Baetis roodani Baetis rhodani
```

# Retrieve higher taxonomic names

Another common task is to retrieve the complete taxonomic hierarchy for a taxon. Different sources with different coverages can be used.

```
classification('Baetis rhodani', db = 'col')
                           rank
               name
          Animalia
                     Kingdom
## 2
        Arthropoda
                         Phylum
## 3
           Insecta
                          Class
                          Order
      Ephemeroptera
         Baetoidea Superfamily
          Baetidae
                         Family
             Baetis
                          Genus
## 8 Baetis rhodani
                        Species
```

## Retrieve children taxa

One can also search in the opposite direction, i.e. search species within a genus:

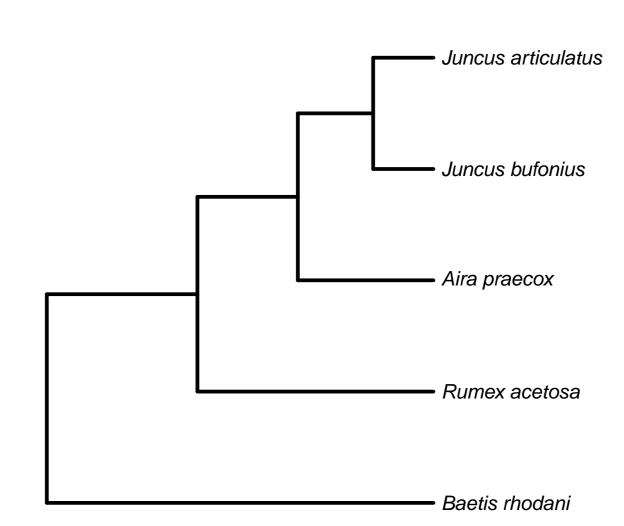
```
downstream('Baetis', db = 'col', downto = 'Species')
        childtaxa_name childtaxa_rank
      Baetis acceptus
                              Species
     Baetis aculeatus
                              Species
    Baetis acuminatus
                              Species
        Baetis adonis
                              Species
                              Species
        Baetis aeneus
        Baetis alius
                              Species
        Baetis alpinus
                              Species
```

#### Features (cont.)

#### Hierarchy trees

The taxonomic relationships between species can be displayed in hierarchy trees. These could be used for example as surrogates when phylogenetic data is scarce<sup>3</sup>.

```
species <- c('Juncus bufonius', 'Juncus articulatus',
    'Aira praecox', 'Rumex acetosa', 'Baetis rhodani')
hier <- classification(species, db = 'ncbi')
plot (class2tree (hier) )
```



## Aggregate taxa

Using the taxonomic information taxa can be easily aggregated to different levels, e.g. to study effects on different taxonomic levels. Taxize provides the tax\_agg() function for this purpose:

```
tax_agg(dune, rank = 'family', db = 'ncbi')
```

## Other functionality

Resolve ambiguous taxa4: Taxonomic information forms the basis for these methods, for details see the restax package<sup>5</sup>.

**IUCN**: Query the IUCN status for taxa.

Invasive species: Check if a taxon is listed in the Global Invasive Species Database (GISD). Integration of the Global Invasive Species Information Network (GISIN) is under development.

### **Under the hood**

taxize grabs data from the internet, formats and returns it to the user. This would not have been possible without the work of others:

### **Calling Servers**

httr and RCurl

### **Parsing**

XML and RJSONIO

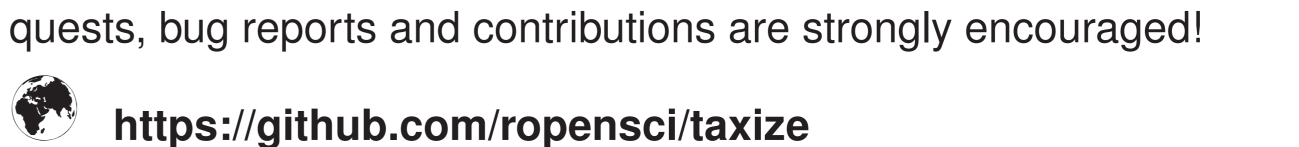
### Data manipulation

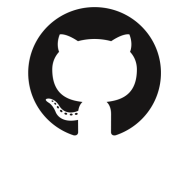
stringr, plyr, reshape2 and vegan

And, of course, base-R;)

# **Get involved!**

taxize is currently developed collaboratively on GitHub. Feature re-





### References

[5] restax-package. https://github.com/EDiLD/restax.

- [1] Scott A. Chamberlain and Eduard Szöcs. taxize: taxonomic search and retrieval in r [v2; ref status: indexed, http://f1000r.es/24v]. F1000Research, 2(191),
- [2] Brad Boyle, Nicole Hopkins, Zhenyuan Lu, et al. The taxonomic name resolution service: an online tool for automated standardization of plant names. BMC Bioinformatics, 14(1):16, January 2013.
- [3] Guillaume Guénard, Peter Carsten von der Ohe, Dick de Zwart, Pierre Legendre, and Sovan Lek. Using phylogenetic information to predict species tolerances to
- toxic chemicals. Ecological Applications, 21(8):3178-3190, 2011. [4] T. F. Cuffney, M. D. Bilger, and A. M. Haigler. Ambiguous taxa: effects on the characterization and interpretation of invertebrate assemblages. Journal of the North
  - American Benthological Society, 26(2):286-307, June 2007.