taxize - taxonomic search and retrieval in R

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I. INTRODUCTION

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II. THE CASE FOR TAXIZE

There are a large suite of applications developed around the problem of searching for, resolving, and getting higher taxonomy for species names. For example, Linnaeus http://linnaeus.sourceforge.net/provides ability to search for taxonomic names in documents and normalize names. In addition, there are many web interfaces to search for and normalize names such as Encyclopedia of Life's Global Names Resolver http://resolver.globalnames.org/, uBio tools http://www.ubio.org/index.php?pagename=sample_tools, and iPlant's Taxonomic Name Resolution Service http://tnrs.iplantcollaborative.org/.

All of these tools provide great ways to search for taxonomic names and resolve them in some cases. However, scientists ideally need a tool that can be used programmatically, and thus be made reproducible, and highly customizeable. The goal of taxize is to make it easy to create reproducible and easy to use workflows for searching for taxonomic names, resolving them, getting higher taxonomic names, and other tasks related to research dealing with species.

III. DATA SOURCES

taxize uses many data sources, and more can easily be added.

Tibble ii batta battata da abat ii tamba				
Source name	Name search	Name resolution	Phylogeny	URL
Encyclopedia of Life	Yes	See GNR below	No	http://eol.org/
Integrated Taxonomic Resolution Service	Yes	Synonyms	No	X
iPlant Taxonomic Name Resolution Service	Yes	Yes	No	X
Phylomatic	No	X	No	X
uBio	Yes	X	No	X
Global Names Resolver (EOL)	Yes	X	No	X

TABLE I. Data sources used in taxize

IV. USE CASES

There are a variety of use cases for which taxize is ideally suited, and few side cases in which taxize can be useful. We discuss five ideal use cases for taxize at length, and highlight the side cases in brief.

A. Installing taxize

First, let's install taxize. There are two versions of taxize, a stable release that can be installed from the R package repository, CRAN, and from GitHub, where the code is developed.

Installing from CRAN or GitHub

```
## From CRAN
install.packages("taxize")

## From GitHub
install_github("taxize_", "ropensci")
```

Loading into your R session

```
library(taxize)
```

B. Resolve taxonomic names

This is a common task in biology. We often have a list of species names and we want to know if a) we have the most up to date names, b) our names are spelled correctly, and c) if we have common names, we likely need the scientific names. One way to resolve names is via the Global Names Resolver service provided by the Encyclopedia of Life (http://resolver.globalnames.org/).

```
# Here, we are searching for two misspelled names
temp <- gnr_resolve(names = c("Helianthos annus", "Homo saapiens"), returndf = TRUE)</pre>
# let's take a peek at the data, excluding the data source ID and score
# columns
temp[, -c(1, 4)]
    submitted_name
                                                                  title
                                   name_string
1 Helianthos annus
                          Helianthus annuus L.
                                                      Catalogue of Life
                              Helianthus annus GBIF Taxonomic Backbone
3 Helianthos annus
4 Helianthos annus
                              Helianthus annus
5 Helianthos annus
                           Helianthus annus L.
                                                                     EOL
6 Helianthos annus
                              Helianthus annus
                                                          uBio NameBank
     Homo saapiens Homo sapiens Linnaeus, 1758
                                                      Catalogue of Life
```

Looks like the correct spellings are *Helianthus annuus* and *Homo sapiens*, cool!

Another approach is using the Taxonomic Name Resolution Service via the Taxosaurus API (http://taxosaurus.org/).

```
# Lets set our list of species names
mynames <- c("Helianthus annuus", "Pinus contort", "Poa anua", "Abis magnifica",
    "Rosa california", "Festuca arundinace", "Sorbus occidentalos", "Madia sateva")
# And we'll call the API with the thrs function
tnrs(query = mynames)[, -c(5:7)]
        submittedName
                             acceptedName
                                             sourceId score
7
   Helianthus annuus
                        Helianthus annuus iPlant_TNRS
                                                      1.00
4
        Pinus contort
                           Pinus contorta iPlant_TNRS
                                                       0.98
5
            Poa anua
                                 Poa alta iPlant_TNRS
                                                       0.77
3
       Abis magnifica
                          Abies magnifica iPlant_TNRS
                                                       0.96
8
     Rosa california
                         Rosa californica iPlant_TNRS
                                                       0.99
  Festuca arundinace Festuca arundinacea iPlant_TNRS
                                                       0.99
1 Sorbus occidentalos Sorbus occidentalis iPlant_TNRS
                                                       0.99
         Madia sateva
                             Madia sativa iPlant_TNRS
                                                       0.97
```

It looks like there are a few corrections: e.g., *Madia sateva* should be *Madia sativa*, and *Rosa california* should be *Rosa californica*.

C. Retrieve higher taxonomic names

Another task biologists often face is wanting to get higher taxonomic names for their list of taxa. If you have the higher taxonomy you can put in to context the relationships of your list (i.e., Species A and B are in Family X), as opposed to not knowing that Species A and B are closely related. A number of data sources provide this type of capability. First, let's take a look at the Integrated Taxonomic Information Service (ITIS).

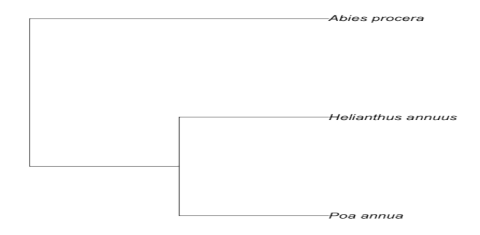
```
specieslist <- c("Abies procera", "Pinus contorta")
classification(get_tsn(specieslist, "sciname"))</pre>
```

```
Retrieving data for species ' Abies procera '
Retrieving data for species ' Pinus contorta '
[[1]]
        parentName parentTsn
                                   rankName
                                                   taxonName
                                                                tsn
1
                                    Kingdom
                                                    Plantae 202422
2
           Plantae
                      202422
                                 Subkingdom
                                             Viridaeplantae 846492
3
    Viridaeplantae
                      846492
                               Infrakingdom
                                               Streptophyta 846494
4
      Streptophyta
                      846494
                                   Division
                                               Tracheophyta 846496
5
      Tracheophyta
                      846496
                                Subdivision Spermatophytina 846504
  Spermatophytina
                      846504 Infradivision
                                               Gymnospermae 846506
6
7
      Gymnospermae
                      846506
                                      Class
                                                   Pinopsida 500009
8
         Pinopsida
                      500009
                                      Order
                                                    Pinales 500028
9
           Pinales
                      500028
                                     Family
                                                   Pinaceae 18030
                                                       Abies 18031
10
          Pinaceae
                      18030
                                     Genus
11
             Abies
                       18031
                                    Species
                                              Abies procera 181835
[[2]]
        parentName parentTsn
                                   rankName
                                                   taxonName
                                                                tsn
                                                    Plantae 202422
1
                                    Kingdom
2
           Plantae
                      202422
                                 Subkingdom
                                             Viridaeplantae 846492
3
    Viridaeplantae
                      846492
                               Infrakingdom
                                               Streptophyta 846494
                                               Tracheophyta 846496
4
                      846494
      Streptophyta
                                   Division
5
      Tracheophyta
                      846496
                                Subdivision Spermatophytina 846504
6
  Spermatophytina
                      846504 Infradivision
                                               Gymnospermae 846506
                                                   Pinopsida 500009
7
      Gymnospermae
                      846506
                                      Class
8
                                      Order
                                                    Pinales 500028
         Pinopsida
                      500009
9
           Pinales
                      500028
                                     Family
                                                   Pinaceae 18030
10
          Pinaceae
                       18030
                                      Genus
                                                       Pinus 18035
                                    Species Pinus contorta 183327
11
             Pinus
                       18035
```

You can also get this type of information from uBio

D. Retrieve a phylogeny

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E. Use ITIS locally for faster searches

text

F. Use ITIS locally for faster searches

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V. CONCLUSION

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VI. FUNDING

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