# TR8: Extract traits data for plant species

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## 1 Rationale

The TR8 package has been built in order to provide the user with the possibility of easily retrieving traits data for plant species from the following publicly available databases:

Biolflor http://www.ufz.de/biolflor/index.jsp [7]

Ecological Flora of the British Isles http://www.ecoflora.co.uk/ [3]

LEDA traitbase http://www.leda-traitbase.org/LEDAportal/ [6]

Ellenberg values for Italian Flora [10]

Flowering period for Italian Flora [10] (data retrieved from http://luirig. altervista.org/)

Mycorrhizal intensity database [2]

MycoFlor database [4]

Catminat database [5]

**BROT** [8][9]

Please note that not all the traits available on the listed databases are downloaded by the package: this may change in future versions of the package (ie. some functionalities may be added and more traits will be made available).

## 2 Installation

The TR8 package is available on CRAN, thus it can be easily installed through:

```
> install.packages("TR8",dependencies = TRUE)
```

The option dependencies = TRUE takes care of installing those packages which are needed by TR8 to work properly.

Once the package is installed, you can load it with:

> library(TR8)

Please note that:

The user is asked to always cite the data sources: the development of traits databases is a long and costly process, thus all the users of the TR8 package are asked (and reminded every time they load the package) to always cite the original sources of the data (see paragraph 6).

#### 2.1 Using the development version

The devel version of the package is hosted on github at https://github.com/ GioBo/TR8: to use this version (instead of the stable one, released from CRAN), you'll need the devtools package (https://github.com/hadley/devtools):

```
> ## install the package
> install.packages("devtools")
> ## load it
> library(devtools)
> ## activate dev_mode
> dev_mode(on=T)
> ## install TR8
> install_github("GioBo/TR8",ref="master")
> ## load it
> library(TR8)
> ## you can now work with TR8 functions
>
> ## if you want to go back and use the CRAN version
> ## already installed, simply deactivate dev_mode
> dev_mode(on=F)
```

### 3 Simple usage

Using the TR8 package is fairly simple: users just need to call the tr8 function passing, as arguments, a vector of plant species names (without authors' names!<sup>1</sup>) and a vector containing the codes corresponding to the traits which are to be downloaded:

<sup>&</sup>lt;sup>1</sup>This is needed since some traitbases do not include authors' names in the species' names.

```
> ## a vector containing a list of plant species names
> my_species<-c("Apium graveolens","Holcus mollis","Lathyrus sylvestris")
> ## a vector of traits
> to_be_downloaded<-c("reprod_B","strategy")
> ## now run tr8 and store the results in the my_traits object
> my_traits<-tr8(species_list = my_species,download_list = to_be_downloaded)</pre>
```

The codes which are accepted by  ${\tt TR8}$  are listed in the <code>available\_tr8</code> database:

> ## see the firs lines of available_tr8 database			
> head(available_tr8)			
	<pre>short_code</pre>	description	db
1	h_max	Maximum height	Ecoflora
2	h_min	Minimum height	Ecoflora
3	le_area	Leaf area	Ecoflora
4	le_long	Leaf longevity	Ecoflora
5	phot_path	Photosynthetic pathway	Ecoflora
6	li_form	Life form	Ecoflora

The database is composed of three columns:

- short\_code contains the codes that should be passed to the download\_list
  argument of the tr8 function.
- **description** contains short description of each trait (please refer to the original sources for detailed descriptions).
- db refers to the databases from which are providing traits data

Suppose the user is interested in downloading the *maximum height*, the *leaf* area and the *life form* (which are available through the *Ecolfora* database) for Salix alba and Populus nigra and store the resulting data in the my\_Data object; the command should be:

```
> my_species<-c("Salix alba","Populus nigra")
> my_traits<-c("h_max","le_area","li_form")
> my_Data<-tr8(species_list = my_species, download_list = my_traits)</pre>
```

The tr8 function will take care of downloading the data and store them in the my\_Data object; you can see the results by printing them:

> ## see the downloaded data
> print(my\_Data)

Or you can convert them to a data frame using the extract\_traits function:

```
> traits_dataframe<-extract_traits(my_Data)</pre>
```

All the traits are now contained in a data frame with species as rows and columns as traits; where no trait data were available, you will see a NA.

In order to make the dataframe more readable, traits' names (i.e. columns' names) are converted to shorter codes: to see a brief explanation of the codes used to identify the traits, use the lookup function:

```
> lookup(my_Data)
```

The object returned by the **lookup** function can also be stored in order to be available for further elaborations:

```
> my_lookup<-lookup(my_Data)
> head(my_lookup)
```

#### 3.1 Checking retrieved data

Several steps can go wrong during the data retrieval process (e.g. a database may contain two entries for the same species); tr8() will keep track of some of this problematic cases and the issues() function can be used to see whether any problem was faced during the process.

```
> my_species<-c("Salix alba","Populus nigra")</pre>
```

```
> my_traits<-c("h_max","le_area","li_form")</pre>
```

```
> my_Data<-tr8(species_list = my_species, download_list = my_traits)</pre>
```

```
> issues(my_Data)
```

#### 3.2 Interactive use of tr8

Up to now we've been using the tr8 function in a non-interactive way. In order to help those user which are more familiar with a GUI approach, the function can also be run setting the gui\_config parameter to TRUE (without providing any trait to the download\_list parameter) and a multi-panel window will appear: the user is asked to choose those traits which are to be downloaded from the various databases <sup>2</sup>.

For a detailed explanation of each level of a trait, please refer to the original websites (all the databases listed in the references provide the users with very precise and detailed descriptions).

Typically users will have their vegetation data in the form of a *sites*\**species* dataframe (or matrix), thus they may want to extract traits data for the whole dataset (this time using the GUI to select traits), i.e. :

 $<sup>^2</sup>A$  note for Mac users: the GUI relies on the Tcl/Tk toolkit, thus if you want to run the GUI, please make sure that the X11 package is installed - see "Tcl/Tk" issues at http://cran.r-project.org/bin/macosx/RMacOSX-FAQ.html

```
> ## suppose veg_data is our dataframe with
> ## plant species as columns and sites as rows
>
> ## extract species names
> specie_names<-names(veg_data)
> ## use the tr8() function
> ## and tick those traits of interest in the pop-up window
> my_traits<-tr8(species_names,gui_config=TRUE)
> ## print the results
> print(my_traits)
```

### 4 Interpreting retrieved data

Please note that for many traits there is more than one entry in the original databases: in those cases, in order to obtain a single value the following strategy was adopted:

- **Quantitative traits** the mean of all the values was calculated (e.g. when multiple values for "Seed weight mean" are available, the mean of these value is calculated)
- **Qualitative traits** all the values are taken into account and "joined" together in a single string (the values are separated by a score "-")

Nota bene: in some cases some traits are stored as *string* in the original databases, even though they should be treated as numbers (e.g. the number *five* is stored as a string - i.e. "5", not as the numeric value 5): in those case tr8 function is not able to interpret that entry as a numeric, thus, applying the above mentioned criteria to merge multiple traits, strange outputs may result from tr8, e.g. if a species has two entries for the trait height - day 3 and 3.5 meters - the merged value will not be the numeric mean (3.25) but the union of the two strings ("3-3.5").

Note for BROT database: BROT database contains several entries for the same trait for some species: this may be due to the fact that several sources of data were used and/or that data for different regions were reported. In the current version of TR8 these multiple data are treated as follows:

- if there are both numeric (quantitative) and string (qualitative) entries, numeric ones are given the precedence and the mean of all numeric entries is used as the returned trait value
- if several qualitative entries are present (but no quantitative ones), only the most abundant level is reported (in case ties are present, eg. two levels are reported the same number of times, both values are reported, separated by a comma).

## 5 Citing the package

Please use the following citation when using TR8 package:

Bocci, G. (2015). TR8: an R package for easily retrieving plant species traits. Methods in Ecology and Evolution, 6(3):347-350.

#### Or, if you use BibTeX:

```
@Article{,
   author = {Bocci Gionata},
   title = {TR8: an R package for easily retrieving plant species traits},
   journal = {Methods in Ecology and Evolution},
   year = {2015},
   volume = {6},
   number = {3},
   pages = {347--350},
   url = {http://dx.doi.org/10.1111/2041-210X.12327},
}
```

### 6 Citing sources of information

Users of the TR8 package should always cite the sources of information which provided the traits data: the correct citations to be used for the retrieved data can be obtained through the bib method; just use:

> bib(my\_traits)

### 7 Some notes on using tr8

#### A NOTE OF CAUTION

Searching the web is a time (and Internet band) consuming activity, thus the higher the number of your plant species and the traits to be retrieved, the longer it will take to tr8() to complete its job. Moreover, in order not to overflow the remote databases with http requests, the tr8 function will always pause between one search and the following one.

#### A (SECOND) NOTE OF CAUTION

Some users adopt the following workflow for analysing their vegetation data:

1. insert vegetation data into a *spreadsheet file* with species as columns' and sites' as rows

- 2. export the spreadsheet file as a .csv file
- 3. import the .csv file into a R dataframe.

When following these steps, a dot (".") will be inserted between Genus and Species of each plant species name (i.e. column names in the R dataframe will not be in the form c("Abies alba", "Salix alba") but in the form c("Abies.alba", "Salix.alba")). This may cause problems for further processing of plants' species names, thus, in order to avoid this problem, please use the check.names=F option in read.csv. E.g. suppose that my\_veg\_data.csv is the csv file: in the R console, one should use:

```
> My_data<-read.csv("my_veg_data.csv",
+ header=T,row.names=1,check.names=F)
```

### 8 Suggested workflow

We strongly suggest to always check plant species names with the tnrs function (from the taxize package) before using the tr8 function; thus a typical workflow would be the following:

1. Check plant species names (e.g. with something like the following - please refere to the taxize package documentation[1] for further details)

```
> species_names<-names(veg_data)</pre>
```

```
> checked_names<-tnrs(species_names,source="iPlant_TNRS")</pre>
```

> print(checked\_names)

Check which species (rows) in the table have a "score" value lower than 1 and check their names; if needed, correct them before using the tr8() function.

2. Run tr8 (in this case using the GUI):

```
> my_traits<-tr8(species_names,gui_config = TRUE)
> print(my_traits)
```

Check whether tr8() had any problems in retrieving data:

```
> my_traits<-tr8(species_names,gui_config = TRUE)
> issues(my_traits)
```

3. You may want to have these traits available as a data frame: just use the extract\_traits function which uses the results of tr8 (in this case it's the my\_traits objects) and returns a data frame.

- > traits\_df<-extract\_traits(my\_traits)</pre>
- 4. Observing a big data frame inside R could be difficult, thus users may want to save the traits\_df data frame as a .csv file:

```
> save(traits_df,file="traits_df.csv")
```

and then open that file with a spreadsheet software (e.g. LibreOffice).

#### 8.1 Dealing with synonyms

In some cases the same species may be present in different traitbases under different synonyms. In order to make it easier for the user to deal with this issue, tr8 function has a parameter called synonyms: when set to true it forces the tr8 to call the tnrs function from the taxize package and retrieve possible synonims for the provided species names: in this case the retrieved trait data will have two additional columns, one called synonyms (for the synonyms found) and another named *original\_names* which contains the original names passed by the user with the species\_list parameter<sup>3</sup>.

```
> my_species<-c("Salix alba","Inula viscosa")
> my_traits<-c("h_max","le_area","li_form")
> my_Data<-tr8(species_list = my_species, download_list = my_traits,
+ synonyms=TRUE)</pre>
```

In the Catminat traitbase there are some entries which are in the form "Genus v. subspecies" (e.g. "Myrtus communis v. communis" and "Myrtus communis v. leucocarpa".); in order to allow users to search among those subspecies, the parameter CATMINAT\_ALTERNATIVES has been introduced in the TR8 function: when set to TRUE the function will search for entries which contain, in their names, the names provided SPECIES\_LIST; e.g. if "Myrtus communis" is included in the SPECIES\_LIST, tr8 will query the following existing entries in Catminat: "Myrtus communis", "Myrtus communis v. communis" and "Myrtus communis v. leucocarpa").

```
> my_species<-c("Myrtus communis")
> ## some traits from Catminat
> my_traits<-c("inflorescence_fr","sex_reprod_fr","poll_vect_fr")
> my_Data<-tr8(species_list = my_species, download_list = my_traits,
+ catminat_alternatives=TRUE)</pre>
```

<sup>&</sup>lt;sup>3</sup>**PLEASE NOTE** that when synonyms is set to TRUE, the resulting dataframe will have numbers as row.names: this is mandatory since in a few cases different species names may share a common synonym and R does not allow different rows to share the same name.

#### 9 Further steps

The TR8 package comes with another vignette, called TR8\_workflow, which shows a typical workflow describing all the steps needed for retrieving and analysing traits data with tr8, listing the most common problems that could be faced and the possible solutions to fix them.

The vignette can be opened from within R, using:

```
> vignette("TR8_workflow")
```

Another vignette (called Expanding\_TR8) shows to programmers how sources of data can be added to TR8 (i.e. how functions for retrieving data should be written so that they can be easily integrated in TR8).

### 10 Local storage of remote data

The following databases are stored as files (.txt,.csv or xlsx) on the remote servers:

- LEDA
- Akhmetzhanova
- MycoFlor
- Catminat

These files are (quite) big in size, thus downloading them every time the tr8() function is used is a time consuming activity<sup>4</sup>. In order to make data retrieval more efficient, when tr8 is run AND traits data from the above mentioned databases are requested for the first time, these files are downloaded and an R version (.Rda) copy is stored in a local directory and made available to future requests<sup>5</sup>.

## 11 Update local lookup tables

For some traitbases tr8 uses lookup-tables which contain URLs of the species of interest (i.e. whenever the user search for traits data for some species, tr8 searches for the corresponding URLs in such tables and then retrieve the data from the corresponding webpages); some of the traitbases queried by tr8 are uploaded from time to time thus the function may be unable to retrieve traits

 $<sup>^{4}</sup>$ The text files are not distributed together with the TR8 package - which would save time and memory when executing the tr8() function - in order to avoid possible licensing conflicts between the TR8' GPL license and these datasets.

 $<sup>{}^{5}</sup>$ By default these files will be installed in the directories which are commonly used for storing applications' data (which depends on the underlying operating systems; see https://github.com/hadley/rappdirs for details).

data for the most recently uploaded species. The  ${\tt TR8\_SETUP}$  function allows the user to download the most updated version of such tables; simply run

If the user wants to have available the most updated data,

> tr8\_setup()

**BEWARE**: this function takes a long time to run (nearly 20 minutes are required - on a fast connection - for retrieving lookup tables for BiolFlor); it's very likely that the most recent version of TR8 contains the already up-to-date version of the data, so in most cases will never need to use this function.

### References

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