

# Using **plantR** to Manage Taxonomy

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## Contents

<b>1</b>	<b>Introduction</b>	<b>1</b>
<b>2</b>	<b>Installing plantR</b>	<b>2</b>
<b>3</b>	<b>A practical example</b>	<b>2</b>
3.1	Preparing names using <code>fixSpecies()</code> . . . . .	3
3.1.1	Internal functions . . . . .	3
3.2	Validating taxon names using <code>prepSpecies()</code> . . . . .	4
3.2.1	Internal functions . . . . .	5
3.3	Validating family names using <code>prepFamily()</code> . . . . .	6
<b>4</b>	<b>Brief code summary</b>	<b>6</b>
<b>5</b>	<b>Citation</b>	<b>6</b>
<b>6</b>	<b>References</b>	<b>7</b>
<b>7</b>	<b>Bug report and suggestions</b>	<b>7</b>

## 1 Introduction

One of the most important steps in managing and using information from biological datasets is the management of taxonomic nomenclature. Taxon names need to be correctly spelled and we should use their currently accepted names based on a given taxonomic backbone (Lima et al. in prep.). However, there are many small corrections and standardizations that need to be made before one can cross-validate taxonomic nomenclature, which can become quite burdensome as the size of the dataset increases.

**plantR** provides tools to format, spell-check and validate taxon names at the family and species levels and using different taxonomic backbones. The default backbone used by **plantR** is the Flora e Funga do Brasil. However, any taxonomic backbone can be used, as long as it has a specific content and format. The companion R package **plantRdata** provides other backbones already in this specific format from the World Flora Online, the World Checklist of Vascular Plants and GBIF.

To simplify the process, the management of taxonomic nomenclature in **plantR** can be simply done using the wrapper function `formatTax()` (see the section Brief code summary below). But first, we will explain in detail each step and function to manage taxonomy within **plantR**. These functions can

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be applied individually, as we will show below, but the management of taxonomic nomenclature is potentialised when they are executed in a specific order.

## 2 Installing plantR

The package **plantR** can be installed and loaded from GitHub with the following codes (this may take a minute or two):

```
if (!requireNamespace("remotes"))
  install.packages("remotes")
library("remotes")

if (!requireNamespace("plantR"))
  install_github("LimaRAF/plantR")
library("plantR")
```

For this tutorial, we will also need the global taxonomic backbones available from the companion R package **plantRdata**. Installing this package takes a couple of minutes as well because it stores some large files (>10 MB).

```
install_github("LimaRAF/plantRdata")
```

## 3 A practical example

Let's create a list of names with many common issues in biological datasets. This list includes issues related to format, casing, spelling, synonyms, typos, names with/without authors, incomplete identifications, etc.

```
names <- c(
  "Lindsaea sp.", "Lindsaeaceae sp.",
  "Lindsaea lancea", "Lindsaea lancia", "Lindsaea pumila",
  "Lindsaea lancea (L.) Bedd.",
  "lindsaea lancea", "Lindsaea Lancea", "LINDSAEA LANCEA",
  "Lindsaea lancea var. Falcata",
  "Lindsaea lancea var falcata",
  "Lindsaea Aff. lancea",
  "Lindsaea Aff.lancea",
  "Lindsaea aff. lancea (L.) Bedd.",
  "Lindsaea ×improvisa K.U.Kramer",
  "Parablechnum C.Presl",
  "Blechnum spannagelii Rosenst.",
  "Blechnum cf. spannagelii", "Blechnumcf.spannagelii",
  "Blechnum austrobrasillianum de la Sota",
  "Casearia sylvestris var. angustifolia",
  "Casearia sylvestris var. angustifolia Uittien",
  "Casearia sylvestris angustifolia Uittien",
  "Casearia sylvestris Sw. var. sylvestris",
  "Blechnaceae1",
  "Blechnum sp.2", "Blechnum sp. 2", "Blechnum sp 2", "Blechnum sp",
  "indet", "Indeterminada1"
)
```

### 3.1 Preparing names using `fixSpecies()`

This function accepts both a vector of names or a data frame. In the latter case, the data frame should contain the taxon names in a column called, by default, 'scientificName'. The name of this column can be defined by the user through the argument `tax.name`. And if taxon names and authorities are available in separate columns, which is advised, please make sure that author names are stored in a column called 'scientificNameAuthorship' or alter this column name using the argument `author.name`.

Here we provide to the function the vector of names defined above:

```
names_fixed <- fixSpecies(names)
head(names_fixed[, -c(2,4)], 7)
```

```
#>      scientificName scientificName.new scientificNameStatus
#> 1      Lindsaea sp.      Lindsaea sp.      indet
#> 2      Lindsaeaceae sp. Lindsaeaceae sp.      family_as_genus
#> 3      Lindsaea lancea Lindsaea lancea      possibly_ok
#> 4      Lindsaea lancia Lindsaea lancia      possibly_ok
#> 5      Lindsaea pumila Lindsaea pumila      possibly_ok
#> 6 Lindsaea lancea (L.) Bedd. Lindsaea lancea      name_w_authors
#> 7      lindsaea lancea Lindsaea lancea      name_w_wrong_case
```

The output of `fixSpecies()` is a data frame that contains the columns necessary for the name validation step below. For each name, it is returned a new column containing a suggestion of a more standardised name (i.e. 'scientificName.new'), with the isolation of the name modifiers (e.g. cf. or aff.). Also, if the name contains author names (and if those author names are in a detectable format), they are split into two different columns ('scientificName.new' and 'scientificNameAuthorship.new').

The new column 'scientificNameStatus' stores all possible flags detected in the original name. These flags include the detection and standardization of open nomenclature (e.g. aff., cf.), infra-specific levels (e.g. var., subsp., f.), hybrids, incomplete identifications and undeterminations. It also flags and solves issues related to name casing and the notation of morphotypes.

#### 3.1.1 Internal functions

Besides the flagging of particular cases of taxon names, the function `fixSpecies()` contains different internal functions that standardise the:

- notation of open nomenclature abbreviations and name modifiers (function `fixAnnotation()`);
- notation of incomplete identifications and undeterminations (function `fixIndet()`);
- the casing of taxon names (function `fixCase()`);
- the separation between taxon names and taxon name authorships (function `fixAuthors()`).

These internal functions may also be useful in itself depending on the user's goals. Below, some examples of their isolate use:

```
plantR:::fixAnnotation(c("Lindsaea lancea var falcata", "Lindsaea Aff.lancea"))
#> [1] "Lindsaea lancea var. falcata" "Lindsaea aff. lancea"
plantR:::fixIndet(c("Indet1", "Blechnum sp. 2", "Blechnum sp 2", "Blechnum sp"))
#> [1] "Indet. sp.1" "Blechnum sp.2" "Blechnum sp.2" "Blechnum sp."
plantR:::fixCase(c("lindsaea lancea", "Lindsaea Lancea", "LINDSAEA LANCEA"))
#> lindsaea lancea Lindsaea Lancea LINDSAEA LANCEA
#> "Lindsaea lancea" "Lindsaea lancea" "Lindsaea lancea"
plantR:::fixAuthors(c("Lindsaea lancea (L.) Bedd.", "Parablechnum C.Presl"))
#> orig.name tax.name tax.author
```

```
#> 1 Lindsaea lancea (L.) Bedd. Lindsaea lancea (L.) Bedd.
#> 2 Parablechnum C.Presl Parablechnum C.Presl
```

### 3.2 Validating taxon names using prepSpecies()

After the standardization of notation and format of taxon names, it is possible to validate them against a taxonomic backbone, aiming at finding synonyms, orthographic variants and/or typos.

In **plantR** this validation of the taxonomic nomenclature is done using the function `prepSpecies()`:

```
names_valid <- prepSpecies(names_fixed,
                           tax.names = c("scientificName.new",
                                           "scientificNameAuthorship.new"))
head(names_valid[, -c(2,3,4,9,11)], 7)
#>      scientificName scientificNameStatus suggestedFamily suggestedName suggestedAuthors
#> 1      Lindsaea sp.                indet      Lindsaeaceae      Lindsaea      Pic.Se
#> 2      Lindsaeaceae sp.          family_as_genus Lindsaeaceae Lindsaeaceae      C.Pr
#> 3      Lindsaea lancea          possibly_ok    Lindsaeaceae Lindsaea lancea      (L.) Be
#> 4      Lindsaea lancia          possibly_ok    Lindsaeaceae Lindsaea lancea      (L.) Be
#> 5      Lindsaea pumila          possibly_ok    Lindsaeaceae Lindsaea lancea      (L.) Be
#> 6 Lindsaea lancea (L.) Bedd.      name_w_authors Lindsaeaceae Lindsaea lancea      (L.) Be
#> 7      lindsaea lancea          name_w_wrong_case Lindsaeaceae Lindsaea lancea      (L.) Be
#>      scientificNameFull
#> 1      Lindsaea Pic.Serm.
#> 2      Lindsaeaceae C.Presl
#> 3 Lindsaea lancea (L.) Bedd.
#> 4 Lindsaea lancea (L.) Bedd.
#> 5 Lindsaea lancea (L.) Bedd.
#> 6 Lindsaea lancea (L.) Bedd.
#> 7 Lindsaea lancea (L.) Bedd.
```

The output of the function is optimized to ease the user's interpretation and decision-making. The output column 'tax.notes' contains the important comments related to the name validation, while the argument `drop.cols` controls which columns should be returned (set `drop.cols = ""` to get all columns). By default, synonyms and orthographic variants are replaced by the name accepted in the reference backbone.

In addition, `prepSpecies()` has arguments to control the minimum similarity allowed between names in fuzzy matching (argument `sug.dist`) and the editing and cleaning of names before matching (arguments `clean.indet` and `clean.names`).

The function was conceived to validate thousands of names at once. So, to speed up the validation process, users can validate names by their initial letter (argument `split.letters`) or parallelise the computation (arguments `parallel` and `cores`).

By default, `prepSpecies()` uses the internal backbone from the Flora e Funga do Brasil. But any backbone in the right format can be provided by the user via the argument `db` of `prepSpecies()`. The **plantRdata** package, available only on GitHub at this link, provides preformatted objects with backbones from different sources. From **plantRdata** we are going to use two other taxonomic backbones from the World Flora Online and the World Checklist of Vascular Plants (use `?wfoNames` and `?wcvpNames` for details on those backbones).

```
# loading the WFO and WCVp backbones into a temporary environment
temp.env <- new.env(parent = emptyenv())
data(list = c("wfoNames", "wcvpNames"), package = "plantRdata",
```

```

    envir = temp.env)
# using the World Flora Online
names_valid_wfo <- prepSpecies(names_fixed,
                               tax.names = c("scientificName.new",
                                              "scientificNameAuthorship.new"),
                               db = temp.env$wfoNames)
# using the World Checklist of Vascular Plants
names_valid_wcvp <- prepSpecies(names_fixed,
                                tax.names = c("scientificName.new",
                                              "scientificNameAuthorship.new"),
                                db = temp.env$wcvpNames)

# Comparing the results
names_bfo_wfo_wcvp <- cbind.data.frame(names_valid$scientificName.new,
                                         names_valid$scientificNameFull,
                                         names_valid_wfo$scientificNameFull,
                                         names_valid_wcvp$scientificNameFull)
diff <- names_valid$scientificNameFull != names_valid_wfo$scientificNameFull
diff[is.na(diff)] <- FALSE
head(names_bfo_wfo_wcvp[diff, ], 3)
#>   names_valid$scientificName.new names_valid$scientificNameFull names_valid_wfo$scientificNameFull
#> 1          Lindsaea sp.          Lindsaea Pic.Serm.          Lindsaea Dryand. ex Sm.
#> 2          Lindsaeaceae sp.          Lindsaeaceae C.Presl          Lindsaeaceae M.R.Schomb.
#> 5          Lindsaea pumila          Lindsaea lancea (L.) Bedd.          Asplenium dielerectum Viane
#>   names_valid_wcvp$scientificNameFull
#> 1          Lindsaea Dryand. ex Sm.
#> 2          Lindsaeaceae sp.
#> 5          Asplenium dielerectum Viane

```

Note that the computing speed when using larger backbones (over a million names) is very different than when using smaller ones (a few hundred thousand names). This is because the name-matching process is done pairwise between the input and reference names. But please try to play with the arguments `split.letters` and `parallel` and `cores` to investigate how fast it can be.

Note as well that each taxonomic backbone provides a different opinion on each name. So it is up to the user to select the best backbone for its own purposes.

### 3.2.1 Internal functions

The most important internal function used by `prepSpecies()` is `nameMatching()`, which is the function that actually performs the exact and fuzzy matching between the input names against the reference names from the taxonomic backbone selected. As mentioned above, the name matching can be parallelized and/or performed separately by initial letters to speed up computational time for larger datasets. But basically, it compares a set of input and reference names trying to find exact or fuzzy matches:

```

input_names <- c("Casearia silvestris", "Casearia decandra")
ref_names <- c("Casearia aculeata", "Casearia arborea",
              "Casearia decandra", "Casearia sylvestris")
nameMatching(input_names, ref_names)
#> [1] 4 3

```

### 3.3 Validating family names using `prepFamily()`

**plantR** contains an internal dictionary of valid family names which can be used via the function `prepFamily()`. Currently, valid family names are based on the APG IV (angiosperms) and the PPG I (ferns and lycophytes). And if the family info is missing it can also be found based on genus information.

```
names_valid <- prepFamily(names_valid,
                           fam.name = "suggestedFamily",
                           spp.name = "scientificName.new")
```

It was not the case in this example, but `prepFamily()` returns warnings in the case of conflicts between the original family names and the family name suggested by the APG IV/PPG I family list. In this example, `prepFamily()` just added the missing family names based on the names of the genera, in the new column called 'family.new'.

## 4 Brief code summary

In this tutorial, we described many steps and details related to the functions that manage taxonomy within **plantR**. Here we provide the main codes that you actually need to include in your scripts, assuming that you already created the vector of taxon names called `names` and that you only want to validate it against the Flora e Funga do Brasil:

```
names_fixed <- fixSpecies(names)
names_valid <- prepSpecies(names_fixed,
                           tax.names = c("scientificName.new",
                                           "scientificNameAuthorship.new"))
names_valid <- prepFamily(names_valid,
                           fam.name = "suggestedFamily",
                           spp.name = "scientificName.new")
```

Or, as mentioned above, even simpler using the wrapper `formatTax()`:

```
names_df <- data.frame(scientificName = names)
names_df_valid <- formatTax(names_df)
```

## 5 Citation

If you use **plantR**, please cite it as:

Lima, R.A.F., Sánchez-Tapia, A., Mortara, S.R., ter Steege, H., Siqueira, M.F. (2021). *plantR*: An R package and workflow for managing species records from biological collections. *Methods in Ecology and Evolution* 14(2): 332-339. <https://doi.org/10.1101/2021.04.06.437754>

And please also cite the taxonomic backbones that you used:

Borsch, T., Berendsohn, W., Dalcin, E., et al. (2020). World Flora Online: Placing taxonomists at the heart of a definitive and comprehensive global resource on the world's plants. *Taxon*, 69(6): 1311-1341. <https://doi.org/10.1002/tax.12373>

Govaerts, R., Nic Lughadha, E., Black, N. et al. (2021). The World Checklist of Vascular Plants, a continuously updated resource for exploring global plant diversity. *Sci. Data* 8: 215. <https://doi.org/10.1038/s41597-021-00997-6>

GBIF Secretariat (2023). GBIF Backbone Taxonomy. Checklist dataset. accessed via GBIF.org. <https://doi.org/10.15468/39omei>

Flora e Funga do Brasil (Constantly updated): Flora e Funga do Brasil project. Instituto de Pesquisas Jardim Botânico do Rio de Janeiro. Dataset/Checklist. <https://doi.org/10.15468/1mtkaw>

## **6 References**

Lima, R.A.F. et al. (in prep.) Ten simple rules of taxonomy for ecological datasets.

## **7 Bug report and suggestions**

The plantR project is hosted on GitHub. Please report any bugs and suggestions for improvements to the package [here](#).