

RefDBdelimiter: An R Package for Refining Molecular Reference Databases with Occurrence Data and Performing Molecular Identification for Metabarcoding Analysis

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OVERVIEW

This study presents an integrated workflow designed for streamlined acquisition and filtering phylogenetic data based on occurrence, and taxonomic assignment of Operational Taxonomic Units (OTUs) generated by metabarcoding approach. The workflow comprises ten sequential steps aimed at ensuring data accuracy and relevance. Initially, occurrence data is acquired from the Global Biodiversity Information Facility (GBIF) database, followed by rigorous filtering to exclude erroneous entries and standardize information. Subsequently, taxa-specific occurrence data are selected based on predefined geographic regions. Concurrently, sequences data are retrieved from the National Center for Biotechnology Information (NCBI) database and undergo stringent filtering procedures, including the exclusion of unverified sequences and standardization of sequence headers. Taxa-specific sequences are then selected to align with the occurrence data. This integrated approach enables the establishment of a local curated database, facilitating efficient data management and analysis. Furthermore, the workflow incorporates the assignment of taxonomic identifications to OTUs generated through metabarcoding workflows, leveraging the curated database for accurate taxonomic inference. The presented workflow offers a comprehensive framework for biodiversity studies, enhancing data reliability and enabling more robust analyses of species distributions and genetic diversity.

• Install and/or Load packages:

• Define the needed directories:

```
Directory <- paste0("C:/Users/Administrador/Downloads")
setwd("C:/Users/Administrador/Downloads")
```

• Install "wsl" command (This command creates a Linux subsystem in any windows machine):

```
# Check if WSL is installed
wsl_path <- Sys.which("wsl")

# If WSL is not installed, run the command to install it
if (nzchar(wsl_path)) {
   print("WSL is installed.")
} else {
   print("WSL is not installed. Installing...")
   system("wsl --install")
}</pre>
```

[1] "WSL is installed."

Get gbif database based on continent and taxa (i.e: Plants from South America):

```
setwd("C:/Users/Administrador/Downloads")
source("C:/Users/Administrador/Downloads/functions.R")
generate_gbif_taxa_dataset_gibi(continent = "SOUTH_AMERICA", scientificName = "Plantae", taxa_n = 300, gbif_taxa_dataset = "C:/Users/Administrador/Downloads/gbif_taxa_dataset.txt")
## [1] "GBIF data acquisition completed successfully."
```

• Format headers of cleaned ncbi database:

```
setwd("C:/Users/Administrador/Downloads/")
source("C:/Users/Administrador/Downloads/functions.R")
format_ncbi_database_gibi(raw_database = "its_database.fasta", database_cleaned = "database_cleaned_formated.fast a", min_sequence_length = 100, pattern = "UNVERIFIED")

## [1] "NCBI data cleaning is completed successfully."
```

Subset NCBI databse based on GBIF:

[1] "NCBI subset is completed successfully."

```
setwd("C:/Users/Administrador/Downloads/")
subset_ncbi_based_on_gbif_gibi(gbif_database = "gbif_taxa_dataset.txt", cleaned_ncbi_database = "database_cleaned
_formated.fasta", ncbi_database_based_on_gbif = "its2_database.fasta")
```

• Database production:

```
setwd("C:/Users/Administrador/Downloads/")
create_blast_db_gibi(database = "its2_database.fasta", parse_seqids = T, title = "ITS2_database", database_type =
"nucl")

## [1] "Database creation completed successfully."
```

• Taxonomic assignment of Operational Taxonomic Units (OTUs) based on a curated database:

```
source("C:/Users/Administrador/Downloads/functions.R")
setwd("C:/Users/Administrador/Downloads/")
Database_File <- paste0("/mnt/c/Users/Administrador/Downloads/its2_database.fasta")
Directory <- "C:/Users/Administrador/Downloads/"

blast_gibi(Directory, Database_File, max_target_seqs = 50, perc_identity = 90, qcov_hsp_perc = 95, num_threads = 6, Specie_Threshold = 98, Genus_Threshold = 95, Family_Threshold = 90)

## Database already exists, returning old file

## [1] "End of Run"</pre>
```

• See the Taxonomic assignment results:

```
taxonomic_assignment <- read.table("taxonomic_assignment.txt", header = TRUE, sep = "")</pre>
head(taxonomic_assignment[,1:10])
                      seqid Phylum
## qseqid
                                             Class
## 1 Otul Epidendrum fulgens Streptophyta Magnoliopsida Asparagales
## 2 Otu2 Talipariti tiliaceum Streptophyta Magnoliopsida
## 3 Otu3 Justicia carnea Streptophyta Magnoliopsida Lamiales
## 4 Otu4 Passiflora caerulea Streptophyta Magnoliopsida Malpighiales
## 5 Otu5 Cymbalaria muralis Streptophyta Magnoliopsida Lamiales
## 6 Otu6 Justicia carnea Streptophyta Magnoliopsida Lamiales
         Family Genus Species pident Sample1
## 1 Orchidaceae Epidendrum Epidendrum fulgens 100 0
## 2 Malvaceae Talipariti Talipariti tiliaceum 100 5
## 3 Acanthaceae Justicia Justicia carnea 100 0
## 4 Passifloraceae Passiflora Passiflora caerulea 100 66
## 5 Plantaginaceae Cymbalaria Cymbalaria muralis 100 0
## 6 Acanthaceae Justicia Justicia carnea 100 12
```