

# Package ‘ClassifyITS’

April 23, 2026

**Title** Fungal Assignment Pipeline

**Version** 1.0.2

**Description** Fungi are ubiquitous in Earth's wonderfully diverse ecosystems. The 'ClassifyITS' package aids in the taxonomic classification of environmental internal transcribed spacer (ITS) short-read barcoding data. Unlike previous methods, it employs taxon-specific e-value and percent identity cutoffs at each taxonomic rank from kingdom to species. The package takes a conservative approach and outputs both graphics and user-friendly files to help users manually inspect fungal operational taxonomic units (OTUs) that fail classification at relevant levels (e.g., Phylum). 'ClassifyITS' is based on taxonomic cutoff criteria from ``The Global Soil Mycobiome consortium dataset for boosting fungal diversity research" (Fungal Diversity, Tedersoo, 2021, <[doi:10.1007/s13225-021-00493-7](https://doi.org/10.1007/s13225-021-00493-7)>) and ``Best practices in metabarcoding of fungi: From experimental design to results" (Molecular Ecology, Tedersoo, 2022, <[doi:10.1111/mec.16460](https://doi.org/10.1111/mec.16460)>).

**License** GPL-3

**Encoding** UTF-8

**Imports** ggplot2, gridExtra, grid, reshape2, data.table, seqinr

**Suggests** formatR, knitr, rmarkdown

**RoxygenNote** 7.3.3

**VignetteBuilder** knitr, rmarkdown, formatR

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**Author** Quinn Moon [aut, cre]

**Maintainer** Quinn Moon <[qmoon@umich.edu](mailto:qmoon@umich.edu)>

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best\_hit\_taxonomy\_assignment

*Hierarchical best-hit taxonomy assignment with per-rank fallback rule*

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### Description

Pass ONLY those OTUs that haven't been assigned already! For each rank, if the best e-value hit is undefined and the second-best hit is defined and at least 60

### Usage

```
best_hit_taxonomy_assignment(
  blast_qc,
  cutoffs_long,
  genus_cutoff_mode = c("prefer_evalue", "prefer_pident", "both")
)
```

### Arguments

**blast\_qc** A data.frame of BLAST results for query sequences. Must include qseqid, evalue, pident, length, and taxonomy columns: kingdom/phylum/class/order/family/genus/species.

**cutoffs\_long** Long-form cutoffs (parse\_taxonomy\_cutoffs()\$long).

**genus\_cutoff\_mode** One of: "prefer\_evalue", "prefer\_pident", "both".

### Details

Defaults are taken from the cutoffs table itself (Fungi baseline rules), not from a separate defaults list.

### Value

A data.frame containing hierarchical taxonomy assignment for each query sequence.

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check_N	<i>Check proportion of N bases in each sequence.</i>
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**Description**

Calculates the proportion of "N" bases (ambiguous bases) in each sequence and flags if above the given threshold.

**Usage**

```
check_N(rep_seqs, cutoff = 1)
```

**Arguments**

rep_seqs	Character vector, list (e.g., from <code>seqinr::read.fasta(as.string=TRUE)</code> ), or (optionally) a <code>DNAStrngSet</code> .
cutoff	Numeric, percent threshold (default 1).

**Value**

Data frame with columns: `qseqid`, `N_percent`, `N_flag`.

**Examples**

```
seqs <- c(seq1 = "ATGCNNNN", seq2 = "NNNNATGC")
check_N(seqs)
check_N(seqs, cutoff = 10)
```

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consensus_taxonomy_assignment	<i>Per-rank consensus filter for taxonomy assignment</i>
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**Description**

Only confirms or demotes, never promotes Unclassified. FINAL hierarchy check: if any rank is Unclassified, all lower ranks are forced to Unclassified.

**Usage**

```
consensus_taxonomy_assignment(final_table, blast_qc)
```

**Arguments**

final_table	Data frame of taxonomic assignments.
blast_qc	Data frame of filtered BLAST hits for each OTU.

**Value**

Data frame of consensus assignments (same structure as input).

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easy_assignments	<i>Easy taxonomy assignment for OTUs using BLAST QC output &amp; phylum-specific thresholds.</i>
------------------	--

---

**Description**

Easy taxonomy assignment for OTUs using BLAST QC output & phylum-specific thresholds.

**Usage**

```
easy_assignments(blast_filtered, cutoffs_file = NULL, default_cutoff = 98)
```

**Arguments**

`blast_filtered` QC-filtered BLAST dataframe (with parsed taxonomy columns!)

`cutoffs_file` Path to taxonomy cutoffs CSV file. If not supplied or invalid, attempts to locate the default file in the package.

`default_cutoff` Default percent identity cutoff (kept for API compatibility)

**Value**

List with `assigned_otus_df` and `remaining_otus_df`

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ensure_cols	<i>Ensure data frame has all required columns (as character)</i>
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---

**Description**

Ensure data frame has all required columns (as character)

**Usage**

```
ensure_cols(df, all_cols)
```

**Arguments**

`df` Data frame to fix

`all_cols` Vector of required columns

**Value**

Fixed data frame (in correct order, with all columns present)

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ITS\_assignment                      *Complete Fungal Assignment Pipeline*

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**Description**

Runs all steps: QC, filtering, plotting, assignments; optionally writes outputs.

**Usage**

```
ITS_assignment(
  blast_file,
  rep_fasta,
  cutoffs_file = NULL,
  cutoff_fraction = 0.6,
  n_cutoff = 1,
  outdir = NULL,
  verbose = FALSE
)
```

**Arguments**

blast_file	Path to BLAST results TSV file
rep_fasta	Path to representative sequences FASTA file
cutoffs_file	Path to taxonomy cutoffs CSV file (optional; defaults to package example if omitted)
cutoff_fraction	Numeric, fraction of median rep-seq length for BLAST filtering (default: 0.6)
n_cutoff	Numeric, N base percentage cutoff (default: 1)
outdir	Output directory for results. If NULL (default), nothing is written.
verbose	Logical; if TRUE emit progress messages. Default FALSE.

**Value**

Named list of results and (if written) output file paths

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load\_and\_check                      *Load and check BLAST results and rep-seq FASTA*

---

**Description**

Load and check BLAST results and rep-seq FASTA

**Usage**

```
load_and_check(blast_file, rep_fasta, taxonomy_col = "stitle", verbose = FALSE)
```

**Arguments**

blast_file	Path to BLAST results TSV file.
rep_fasta	Path to representative sequences FASTA file.
taxonomy_col	The column in BLAST file containing taxonomy strings (default "stitle").
verbose	Logical; if TRUE, emit progress messages. Default FALSE.

**Value**

List with BLAST dataframe (kingdom-filtered) and rep\_seqs as a named list of DNA strings.

---

parse\_taxonomy\_cutoffs  
*Parse taxonomy cutoffs file*

---

**Description**

Reads and processes a taxonomy cutoffs CSV for assignment thresholds at various ranks.

**Usage**

```
parse_taxonomy_cutoffs(cutoffs_file = NULL)
```

**Arguments**

cutoffs_file	Path to a taxonomy cutoffs CSV file. If not supplied or invalid, attempts to locate the default file in the package.
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**Value**

A list with two elements: long, a data frame of parsed cutoffs, and ranks, the vector of taxonomic ranks.

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plot\_alignment\_hist *Create and return alignment length histogram (ggplot object)*

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**Description**

Create and return alignment length histogram (ggplot object)

**Usage**

```
plot_alignment_hist(blast, rep_seqs, cutoff_fraction = 0.6)
```

**Arguments**

blast            BLAST data frame.

rep\_seqs        Named list/character vector of DNA strings (from seqinr::read.fasta(as.string = TRUE)).

cutoff\_fraction    Numeric; fraction of median alignment length for cutoff line. Default 0.6.

**Value**

A ggplot object.

---

safe\_rbind\_list        *Safely rbinds list of data frames, ensuring columns match*

---

**Description**

Safely rbinds list of data frames, ensuring columns match

**Usage**

```
safe_rbind_list(dfs, all_cols = NULL)
```

**Arguments**

dfs            List of data frames

all\_cols       Vector of required columns

**Value**

Combined data frame

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save\_taxonomy\_graphics        *Save taxonomy summary charts and tables to multi-page PDF*

---

**Description**

Save taxonomy summary charts and tables to multi-page PDF

**Usage**

```
save_taxonomy_graphics(
  all_results,
  hist_plot,
  pdf_file = NULL,
  caption_texts = NULL,
  rank_names = c("Phylum", "Class", "Order", "Family", "Genus", "Species"),
  verbose = FALSE
)
```

**Arguments**

all_results	Combined assignments table from write_initial_assignments
hist_plot	ggplot2 object for histogram
pdf_file	Output path for multi-page PDF. If NULL (default), no file is written.
caption_texts	Vector of captions for PDF pages (optional)
rank_names	Vector of rank names (default: c("Phylum",...))
verbose	Logical; if TRUE, emit a message when a PDF is written. Default FALSE.

**Value**

List with plots/tables; includes pdf\_file when written.

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trim_alignments	<i>Trim BLAST alignments by minimum length</i>
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---

**Description**

Trim BLAST alignments by minimum length

**Usage**

```
trim_alignments(blast, rep_seqs, fraction = 0.6)
```

**Arguments**

blast	BLAST data frame.
rep_seqs	Named list/character vector of DNA strings (from seqinr::read.fasta(as.string = TRUE)).
fraction	Numeric; fraction of the median rep-seq length used as the cutoff. Default 0.6.

**Value**

Filtered BLAST data frame.

---

`write_initial_assignments`

*Create and write the initial assignments table including drops at all steps*

---

**Description**

Create and write the initial assignments table including drops at all steps

**Usage**

```
write_initial_assignments(  
  easy_df,  
  consensus_df,  
  rep_seqs,  
  blast,  
  blast_filtered,  
  file = NULL,  
  verbose = FALSE  
)
```

**Arguments**

<code>easy_df</code>	Data frame of easy-assigned OTUs
<code>consensus_df</code>	Data frame of consensus-assigned OTUs (hard ones)
<code>rep_seqs</code>	DNASTringSet or named character vector of rep seqs
<code>blast</code>	Data frame of all BLAST results
<code>blast_filtered</code>	Data frame of filtered BLAST results
<code>file</code>	Path for output CSV. If NULL (default), no file is written.
<code>verbose</code>	Logical; if TRUE emit a message when a file is written. Default FALSE.

**Value**

Data frame of assignments (written if file is not NULL)

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