# Package 'isolateR'

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Type Package

**Title** Automated processing of Sanger sequencing data, taxonomic profiling, and generation of microbial strain libraries

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Description isolateR aims to enhance microbial isolation workflows and support the identification of novel taxa. It addresses the challenges of manual Sanger sequencing data processing and limitations of conventional BLAST searches, crucial for identifying microorganisms and creating strain libraries. The package offers a streamlined three-step process that automates quality trimming Sanger sequence files, taxonomic classification via global alignment against type strain databases, and efficient strain library creation based on customizable sequence similarity thresholds. It features interactive HTML output tables for easy data exploration and optional tools for generating phylogenetic trees to visualize microbial diversity. Citation: Daisley et al. (2024). isolateR: an R package for generating microbial libraries from Sanger sequencing data. Bioinformatics/btae448. https://doi.org/10.1093/bioinformatics/btae448

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LazyData true

**Depends** R (>= 4.0)

Imports ape, BiocManager, Biostrings, crosstalk, cowplot, dataui, dplyr, ggtree, ggbeeswarm, ggiraph, IRanges, plotly, htmltools, patchwork, LPSN, methods, msa, pander, R.utils, reactable, reactablefmtr, rentrez, S4Vectors, sangeranalyseR, sangerseqR, scales, seqinr, shiny, stringr, svMisc, xmlconvert, getPass

Remotes timelyportfolio/dataui, glin/reactable, thomasp85/patchwork

**Roxygen** list(markdown = TRUE)

RoxygenNote 7.3.2

Suggests knitr, rmarkdown

VignetteBuilder knitr

Additional\_repositories

http://R-Forge.R-project.org/https://bioconductor.org/packages/3.18/bioc

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# Description

S4 wrapper for isoLIB function. Access data via S4 slot functions.

### Value

Returns an class-isoLIB object.

# **Slots**

input Character string containing input directory information.

sequence\_group Character string containing list of group representative filenames.

date Character string containing run date from each of the input Sanger sequence .ab1 files ("YYYY\_MM\_DD" format).

filename Character string containing input filenames.

phred\_trim Numeric string containing mean Phred scores after trimming.

Ns\_trim Numeric string containing count of N's after trimming.

length\_trim Numeric string containing sequence length after trimming.

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- seqs\_trim Character string containing sequence after trimming.
- closest\_match Character string containing species + type strain no. of closest match from reference database.
- NCBI\_acc Character string containing NCBI accession number associated with closest match from reference database.
- ID Numeric string containing pairwise similarity value for query vs database reference sequence. Calculation of ID is determined by isoTAX 'iddef' parameter (0-4, Default=2). See VSEARCH documentation for more details.
  - (0) CD-HIT definition: (matching columns) / (shortest sequence length).
  - (1) Edit distance: (matching columns) / (alignment length).
  - (2) Edit distance excluding terminal gaps (default definition).
  - (3) Marine Biological Lab definition counting each gap opening (internal or terminal) as a single mismatch, whether or not the gap was extended: 1.0- ((mismatches + gap openings)/(longest sequence length)).
  - (4) BLAST definition, equivalent to –iddef 1 for global pairwise alignments.

rank\_phylum Character string containing Phylum rank taxonomy

rank\_class Character string containing Class rank taxonomy

rank\_order Character string containing Order rank taxonomy

rank\_family Character string containing Family rank taxonomy

rank\_genus Character string containing Genus rank taxonomy

rank\_species Character string containing Species rank taxonomy

- phylum\_threshold Numeric string containing Phylum-level sequence similarity threshold for rank demarcation
- class\_threshold Numeric string containing Class-level sequence similarity threshold for rank demarcation
- order\_threshold Numeric string containing Order-level sequence similarity threshold for rank demarcation
- family\_threshold Numeric string containing Family-level sequence similarity threshold for rank demarcation
- genus\_threshold Numeric string containing Genus-level sequence similarity threshold for rank demarcation
- species\_threshold Numeric string containing Species-level sequence similarity threshold for rank demarcation

#### See Also

isoLIB

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class-isoOC

isoQC Class Object

### **Description**

S4 wrapper for isoQC function. Access data via S4 slot functions.

#### Value

Returns an class-isoQC object.

### **Slots**

date Character string containing run date from each of the input Sanger sequence .ab1 files ("YYYY\_MM\_DD" format).

filename Character string containing input filenames.

trim.start.pos Numeric string containing trimming position start point.

trim.end.pos Numeric string containing trimming position end point.

phred\_spark\_raw List containing per nucleotide Phred score values for each sequence

phred\_raw Numeric string containing mean Phred scores before trimming.

phred\_trim Numeric string containing mean Phred scores after trimming.

Ns\_raw Numeric string containing count of N's before trimming.

Ns\_trim Numeric string containing count of N's after trimming.

length\_raw Numeric string containing sequence length before trimming.

length\_trim Numeric string containing sequence length after trimming.

seqs\_raw Character string containing sequences before trimming.

seqs\_trim Character string containing sequence after trimming.

decision Character string containing decision (PASS/FAIL) information based on isoQC 'min\_phred\_score' and 'min\_length cutoffs'.

input Character string containing input directory information.

#### See Also

isoQC

class-isoTAX 5

class-isoTAX

isoTAX Class Object

#### **Description**

S4 wrapper for isoTAX function. Access data via S4 slot functions.

#### Value

Returns an class-isoTAX object.

### **Slots**

input Character string containing input directory information.

warning Character string containing list filenames of sequences that had poor alignment during taxonomic classification step.

date Character string containing run date from each of the input Sanger sequence .ab1 files ("YYYY\_MM\_DD" format).

filename Character string containing input filenames.

phred\_spark\_raw List containing per nucleotide Phred score values for each sequence

phred\_raw Numeric string containing mean Phred scores before trimming.

phred\_trim Numeric string containing mean Phred scores after trimming.

Ns\_raw Numeric string containing count of N's before trimming.

Ns\_trim Numeric string containing count of N's after trimming.

length\_raw Numeric string containing sequence length before trimming.

length\_trim Numeric string containing sequence length after trimming.

seqs\_raw Character string containing sequences before trimming.

seqs\_trim Character string containing sequence after trimming.

closest\_match Character string containing species + type strain no. of closest match from reference database.

NCBI\_acc Character string containing NCBI accession number associated with closest match from reference database.

- ID Numeric string containing pairwise similarity value for query vs database reference sequence. Calculation of ID is determined by isoTAX 'iddef' parameter (0-4, Default=2). See VSEARCH documentation for more details.
  - (0) CD-HIT definition: (matching columns) / (shortest sequence length).
  - (1) Edit distance: (matching columns) / (alignment length).
  - (2) Edit distance excluding terminal gaps (default definition).
  - (3) Marine Biological Lab definition counting each gap opening (internal or terminal) as a single mismatch, whether or not the gap was extended: 1.0- ((mismatches + gap openings)/(longest sequence length)).
  - (4) BLAST definition, equivalent to –iddef 1 for global pairwise alignments.

rank\_phylum Character string containing Phylum rank taxonomy

rank\_class Character string containing Class rank taxonomy

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rank\_order Character string containing Order rank taxonomy

rank\_family Character string containing Family rank taxonomy

rank\_genus Character string containing Genus rank taxonomy

rank\_species Character string containing Species rank taxonomy

phylum\_threshold Numeric string containing Phylum-level sequence similarity threshold for rank demarcation

class\_threshold Numeric string containing Class-level sequence similarity threshold for rank demarcation

order\_threshold Numeric string containing Order-level sequence similarity threshold for rank demarcation

family\_threshold Numeric string containing Family-level sequence similarity threshold for rank demarcation

genus\_threshold Numeric string containing Genus-level sequence similarity threshold for rank demarcation

species\_threshold Numeric string containing Species-level sequence similarity threshold for rank demarcation

### See Also

isoTAX

df\_to\_isoLIB

Convert isoLIB .CSV output to isoLIB class object

### **Description**

Helper function to convert isoLIB .CSV output to a class-isoLIB class object.

### Usage

```
df_to_isoLIB(df)
```

# **Arguments**

df

Dataframe in same format as .CSV output file from isoLIB step.

# Value

Returns an S4 class-isoLIB object that can be used to generate interactive HTML output tables.

df\_to\_isoTAX

df\_to\_isoTAX

Convert isoTAX .CSV output to isoTAX class object

# **Description**

Helper function to convert isoTAX .CSV output to a class-isoTAX class object.

# Usage

```
df_to_isoTAX(df)
```

# **Arguments**

df

Dataframe in same format as .CSV output file from isoTAX step.

### Value

Returns an S4 class-isoTAX object that can be used to generate interactive HTML output tables.

export\_html

Export HTML for isoQC > isoTAX > isoLIB class objects

### **Description**

S4 wrapper functions to export interactive HTML tables from isoQC, isoTAX, or isoLIB class objects. Saves to HTML to current working directory and automatically opens.

# Usage

```
## S4 method for signature 'isoQC'
export_html(
  obj,
  min_phred_score = NULL,
  min_length = NULL,
  sliding_window_cutoff = NULL,
  sliding_window_size = NULL
)

## S4 method for signature 'isoTAX'
export_html(obj, quick_search = NULL, db = NULL)

## S4 method for signature 'isoLIB'
export_html(obj, method = NULL, group_cutoff = NULL)
```

# **Arguments**

obj

An S4 class object generated from one of isoQC, isoTAX, or isoLIB steps

### Value

HTML output file saved to working directory.

get\_os

get\_db

Download taxonomic reference database

# **Description**

This function downloads taxonomic reference database and formats them for use.

# Usage

```
get_db(db = "16S_bac", force_update = FALSE, add_taxonomy = FALSE)
```

# **Arguments**

db Database selection. One of "16S", "16S\_arc", "18S", "ITS", or "cpn60"

force\_update Forces new databases to be downloaded.

is primarily for cases of operation with no internet or when computing within an HPC cluster without administrator access. Results in a semicolon (;) delimited

FASTA header as follows: Accession\_no;d\_\_Domain;p\_\_Phylum;c\_\_Class;o\_\_Order;f\_\_Family;g\_

# Value

Returns file path for database of interest

# **Examples**

```
db.path <- get_db(db="16S", force_update=FALSE)</pre>
```

get\_os

Determine user operating system.

# Description

Determines the type of operating system being used.

# Usage

get\_os()

# Value

Returns sysname as one of windows/osx-mac/linux

get\_sanger\_date 9

### **Examples**

```
#Example 1 on a Windows-based operating system
os.index <- get_os()
print(os.index)

#Example 2 on a Mac operating system
os.index <- get_os()
print(os.index)

#Example 3 on a Linux operating system
os.index <- get_os()
print(os.index)</pre>
```

get\_sanger\_date

get\_sanger\_date function

# **Description**

Helper function to automatically retrieve run date from Sanger sequencing .ab1 files.

# Usage

```
get_sanger_date(file = NULL)
```

# **Arguments**

file

The .ab1 file in from which to retrieve the date information. (Must be in S4 abif format)

### Value

Returns date in "YYYY\_MM\_DD" format

# **Examples**

isoALL isoALL

get\_vsearch

Download VSEARCH software reference database

### **Description**

This function downloads the VSEARCH software used querying sequences against taxonomic databases of interest.

### Usage

```
get_vsearch(os = NULL)
```

### **Arguments**

os

Operating system, one of: "windows", "osx-mac", or "linux". If blank (os=NULL) then will try to automatically determine operating system.

#### Value

Returns path for VSEARCH executable

### **Examples**

#Example for automatically detecting operating system and downloading VSEARCH software
vsearch.path <- get\_vsearch()</pre>

isoALL

Perform all commands in one step.

# **Description**

This function effectively wraps isoQC, isoTAX, and isoLIB steps into a single command for convenience. Input can be a single directory or a list of directories to process at once. If multiple directories are provided, the resultant libraries can be sequentially merged together by toggling the parameter 'merge=TRUE'. All other respective parameters from the wrapped functions can be passed through this command. The The respective input parameters from the wrapped can be passed through this command with exception of the .creates a strain library by grouping closely related strains of interest based on sequence similarity. For adding new sequences to an already-established strain library, specify the .CSV file path of the older strain library using the 'old\_lib\_csv" parameter.

# Usage

```
isoALL(
  input = NULL,
  export_html = TRUE,
  export_csv = TRUE,
  export_fasta = TRUE,
  export_fasta_revcomp = FALSE,
  export_blast_table = FALSE,
```

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```
quick_search = FALSE,
db = "16S",
iddef = 2,
phylum_threshold = 75,
class_threshold = 78.5,
order_threshold = 82,
family_threshold = 86.5,
genus_threshold = 94.5,
species_threshold = 98.7,
include_warnings = FALSE,
method = "dark_mode",
group_cutoff = 0.995,
keep_old_reps = TRUE,
merge = FALSE
```

# **Arguments**

input Directory path(s) containing .ab1 files. If more than one, provivde as list (e.g.

'input=c("/path/to/directory1","/path/to/directory2")')

export\_html (Default=TRUE) Output the results as an HTML file

export\_csv (Default=TRUE) Output the results as a CSV file.

export\_fasta (Default=TRUE) Output the sequences in a FASTA file.

export\_fasta\_revcomp

(Default=FALSE) Output the sequences in reverse complement form in a fasta file. This is useful in cases where sequencing was done using the reverse primer and thus the orientation of input sequences needs reversing.

quick\_search

(Default=FALSE) Whether or not to perform a comprehensive database search (i.e. optimal global alignment). If TRUE, performs quick search equivalent to setting VSEARCH parameters "-maxaccepts 100 -maxrejects 100". If FALSE, performs comprehensive search equivalent to setting VSEARCH parameters "-maxaccepts 0 -maxrejects 0"

db

(Default="16S") Select database option(s) including "16S" (for searching against the NCBI Refseq targeted loci 16S rRNA database), "ITS" (for searching against the NCBI Refseq targeted loci ITS database. For combined databases in cases where input sequences are dervied from bacteria and fungi, select "16SITS".

iddef

Set pairwise identity definition as per VSEARCH definitions (Default=2, and is recommended for highest taxonomic accuracy) (0) CD-HIT definition: (matching columns) / (shortest sequence length). (1) Edit distance: (matching columns) / (alignment length). (2) Edit distance excluding terminal gaps (default definition). (3) Marine Biological Lab definition counting each gap opening (internal or terminal) as a single mismatch, whether or not the gap was extended: 1.0-((mismatches + gap openings)/(longest sequence length)). (4) BLAST definition, equivalent to –iddef 1 for global pairwise alignments.

phylum\_threshold

Percent cutoff for phylum rank demarcation

class\_threshold

Percent cutoff for class rank demarcation

order\_threshold

Percent cutoff for order rank demarcation

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family\_threshold

Percent cutoff for family rank demarcation

genus\_threshold

Percent cutoff for genus rank demarcation

species\_threshold

Percent cutoff for species rank demarcation

include\_warnings

(Default=FALSE) Whether or not to keep sequences with poor alignment warnings from Step 2 'isoTAX' function. Set TRUE to keep warning sequences, and FALSE to remove warning sequences.

method

Method used for grouping sequences. Either 1) "dark\_mode", or 2) "closest\_species" (Default="dark\_mode").

- Method 1 ("dark\_mode") performs agglomerative hierarchical-based clustering to group similar sequences based on pairwise identity (see 'id' parameter) and then within each group, attempts to assign the longest sequence with the most top hits as the group representative. This method is tailored for capturing distinct strains which may represent novel taxa (i.e. microbial dark matter) during isolation workflows. As such, the sequence representatives chosen in each group will not always have the highest % identity to the closest matching type strain. In some cases, sequence members within a group may also have different taxonomic classifications due to them having close to equidistant % identities to different matching type strain material indicative of a potentially novel taxonomic grouping.
- Method 2 ("closest\_species") groups similar sequences based on their closest matching type strain. For each unique grouping, this results in all sequence members having the same taxonomic classification. The longest sequence with the highest % identity to the closest matching type strain will be assigned as the group representative. Note: The "id" parameter is only used for Method 1 ("dark\_mode") and otherwise ignored if using Method 2 ("closest\_species").

group\_cutoff

(Default=0.995) Similarity threshold based on pairwise identity (0-1) for delineating between sequence groups. 1=100% identical/0.995=0.5% difference/0.95=5.0% difference/etc. Used only if method="dark\_mode", otherwise ignored.

keep\_old\_reps

(Default=TRUE) If TRUE, original sequence representatives from old library will be kept when merging with new library. If FALSE, sequence group representatives will be recalculated after combining old and new libraries. Ignored if old\_lib\_csv=NULL.

merge

If TRUE, combines isoLIB output files consecutively in the order they are listed. Default=FALSE performs all the steps (isoQC/isoTAX/isoLIB) on each directory separately.

verbose

(Default=FALSE) Output progress while script is running.

files\_manual

(Default=NULL) For testing purposes only. Specify a list of files to run as filenames without extensions, rather than the whole directory format. Primarily used for testing, use at your own risk.

exclude

(Default=NULL) For testing purposes only. Excludes files of interest from input directory.

min\_phred\_score

(Default=20) Do not accept trimmed sequences with a mean Phred score below this cutoff

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min\_length (Default=200) Do not accept trimmed sequences with sequence length below this number

sliding\_window\_cutoff

(Default=NULL) Quality trimming parameter (M2) for wrapping SangerRead function in sangeranalyseR package. If NULL, implements auto cutoff for Phred score (recommended), otherwise set between 1-60.

sliding\_window\_size

(Default=15) Quality trimming parameter (M2) for wrapping SangerRead function in sangeranalyseR package. Recommended range between 5-30.

date Set date "YYYY\_MM\_DD" format. If NULL, attempts to parse date from .ab1 file

# Value

Returns a list of class-isoLIB class objects.

#### See Also

```
isoQC, isoTAX, isoLIB
```

#### **Examples**

```
#Set path to directory containing example .ab1 files
fpath1 <- system.file("extdata/abif_examples/rocket_salad", package = "isolateR")
#Run isoALL function with default settings
isoALL(input=fpath1)</pre>
```

isoLIB

Generate new strain library or add to existing one.

### **Description**

This function creates a strain library by grouping closely related strains of interest based on sequence similarity. For adding new sequences to an already-established strain library, specify the .CSV file path of the older strain library using the 'old\_lib\_csv" parameter.

### Usage

```
isoLIB(
  input = NULL,
  old_lib_csv = NULL,
  method = "dark_mode",
  group_cutoff = 0.995,
  keep_old_reps = TRUE,
  export_html = TRUE,
  export_csv = TRUE,
  include_warnings = TRUE,
  vsearch_path = NULL,
  phylum_threshold = 75,
  class_threshold = 78.5,
```

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```
order_threshold = 82,
family_threshold = 86.5,
genus_threshold = 94.5,
species_threshold = 98.7
)
```

### **Arguments**

input

Path of CSV output file from isoTAX step.

old\_lib\_csv

Optional: Path of CSV output isoLIB file or combined isoLIB file from previous run(s)

method

Method used for grouping sequences. Either 1) "dark\_mode", or 2) "closest\_species" (Default="dark mode").

- Method 1 ("dark\_mode") performs agglomerative hierarchical-based clustering to group similar sequences based on pairwise identity (see 'id' parameter) and then within each group, attempts to assign the longest sequence with the most top hits as the group representative. This method is tailored for capturing distinct strains which may represent novel taxa (i.e. microbial dark matter) during isolation workflows. As such, the sequence representatives chosen in each group will not always have the highest % identity to the closest matching type strain. In some cases, sequence members within a group may also have different taxonomic classifications due to them having close to equidistant % identities to different matching type strain material indicative of a potentially novel taxonomic grouping.
- Method 2 ("closest\_species") groups similar sequences based on their closest matching type strain. For each unique grouping, this results in all sequence members having the same taxonomic classification. The longest sequence with the highest % identity to the closest matching type strain will be assigned as the group representative. Note: The "id" parameter is only used for Method 1 ("dark\_mode") and otherwise ignored if using Method 2 ("closest\_species").

group\_cutoff

(Default=0.995) Similarity threshold based on pairwise identity (0-1) for delineating between sequence groups. 1=100% identical/0.995=0.5% difference/0.95=5.0% difference/etc. Used only if method="dark\_mode", otherwise ignored.

keep\_old\_reps

(Default=TRUE) If TRUE, original sequence representatives from old library will be kept when merging with new library. If FALSE, sequence group representatives will be recalculated after combining old and new libraries. Ignored if old\_lib\_csv=NULL.

export\_html

(Default=TRUE) Output the results as an HTML file

export\_csv

(Default=TRUE) Output the results as a CSV file.

include\_warnings

(Default=FALSE) Whether or not to keep sequences with poor alignment warnings from Step 2 'isoTAX' function. Set TRUE to keep warning sequences, and FALSE to remove warning sequences.

vsearch\_path

Path of VSEARCH software if manually downloaded in a custom directory. If NULL (Default), will attempt automatic download.

phylum\_threshold

Percent sequence similarity threshold for phylum rank demarcation

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Percent sequence similarity threshold for genus rank demarcation species\_threshold

Percent sequence similarity threshold for species rank demarcation

#### Value

Returns an isoLIB class object. Default taxonomic cutoffs for phylum (75.0), class (78.5), order (82.0), family (86.5), genus (94.5), and species (98.7) demarcation are based on Yarza et al. 2014, Nature Reviews Microbiology (DOI:10.1038/nrmicro3330)

#### See Also

```
isoTAX, isoLIB
```

### **Examples**

```
#Set path to directory containing example .ab1 files
fpath1 <- system.file("extdata/abif_examples/rocket_salad", package = "isolateR")

#Step 1: Run isoQC function with default settings
isoQC.S4 <- isoQC(input=fpath1)

#Step 2: Run isoTAX function with default settings
fpath2 <- file.path(fpath1, "isolateR_output/01_isoQC_trimmed_sequences_PASS.csv")
isoTAX.S4 <- isoTAX(input=fpath2)

#Step 3: Run isoLIB function with default settings
fpath3 <- file.path(fpath1, "isolateR_output/02_isoTAX_results.csv")
isoLIB.S4 <- isoLIB(input=fpath3)

#Show summary statistics
isoLIB.S4</pre>
```

isoQC

Perform automated quality trimming of input .ab1 files

# Description

This function loads in ABIF files (.ab1 extension) and performs automatic quality trimming in batch mode.

isoQC

#### Usage

```
isoQC(
  input = NULL,
  export_html = TRUE,
  export_csv = TRUE,
  export_fasta = TRUE,
  export_fasta_revcomp = FALSE,
  verbose = FALSE,
  exclude = NULL,
  min_phred_score = 20,
  min_length = 200,
  sliding_window_cutoff = NULL,
  sliding_window_size = 15,
  date = NULL,
  files_manual = NULL
)
```

# **Arguments**

input Path to directory with .ab1 files.

export\_html (Default=TRUE) Output the results as an HTML file export\_csv (Default=TRUE) Output the results as a CSV file. export\_fasta (Default=TRUE) Output the sequences in a FASTA file.

export\_fasta\_revcomp

(Default=FALSE) Output the sequences in reverse complement form in a fasta file. This is useful in cases where sequencing was done using the reverse primer and thus the orientation of input assurance needs reversing

and thus the orientation of input sequences needs reversing.

verbose (Default =FALSE) Output progress while script is running, FALSE for simpli-

fied progress, TRUE for file-by-file details

exclude (Default=NULL) For testing purposes only. Excludes files of interest from input

directory.

min\_phred\_score

(Default=20) Do not accept trimmed sequences with a mean Phred score below

this cutoff

min\_length (Default=200) Do not accept trimmed sequences with sequence length below

this number

sliding\_window\_cutoff

(Default=NULL) Quality trimming parameter (M2) for wrapping SangerRead function in sangeranalyseR package. If NULL, implements auto cutoff for Phred score (recommended), otherwise set between 1-60.

sliding\_window\_size

(Default=15) Quality trimming parameter (M2) for wrapping SangerRead function in sangeranalyseR package. Recommended range between 5-30.

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file

files\_manual (Default=NULL) For testing purposes only. Specify a list of files to run as filenames without extensions, rather than the whole directory format. Primarily

used for testing, use at your own risk.

isoTAX

#### Value

Returns quality trimmed Sanger sequences in FASTA format.

### See Also

```
isoTAX, isoLIB
```

# **Examples**

```
#Set path to directory containing example .ab1 files
fpath1 <- system.file("extdata/abif_examples/rocket_salad", package = "isolateR")
#Step 1: Run isoQC function with default settings
isoQC.S4 <- isoQC(input=fpath1)
#Show summary statistics
isoQC.S4</pre>
```

isoTAX

Classify taxonomy of sequences after quality trimming steps.

# Description

This function performs taxonomic classification steps by searching query Sanger sequences against specified database of interest. Takes CSV input files, extracts FASTA-formatted query sequences and performs global alignment against specified database of interest via Needleman-Wunsch algorithm by wrapping the –usearch\_global command implemented in VSEARCH. Default taxonomic rank cutoffs for 16S rRNA gene sequences are based on Yarza et al. 2014, Nat Rev Microbiol.

# Usage

```
isoTAX(
  input = NULL,
  export_html = TRUE,
 export_csv = TRUE,
 export_blast_table = FALSE,
  quick_search = FALSE,
 db = "16S_bac",
 db_path = NULL,
 vsearch_path = NULL,
  iddef = 2,
 phylum_threshold = 75,
 class_threshold = 78.5,
 order_threshold = 82,
  family_threshold = 86.5,
  genus_threshold = 94.5,
  species_threshold = 98.7
)
```

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#### **Arguments**

input

Path of either 1) CSV output file from isoQC step, or 2) a FASTA formatted file. If input is a FASTA file, the sequence(s) will be converted and saved as an isoQC-formatted output file in the current working directory ("isolateR\_output/01\_isoQC\_mock\_table Sequence date, name, length, and number of ambiguous bases (Ns) will be calculated from the input file and used to populate the relevant columns. Phred quality scores (phred\_trim) will be set to the maximum value (60) and the remaining columns will be populated with mock data to allow compatibility with the isoTAX function. The main purpose of this output file is for flexibility and to allow users to edit/modify the sequence metadata before continuing with subsequent steps.

export\_html (I

(Default=TRUE) Output the results as an HTML file

export\_csv

(Default=TRUE) Output the results as a CSV file.

export\_blast\_table

(Default=FALSE) Output the results as a tab-separated BLAST-like hits table.

quick\_search

(Default=FALSE) Whether or not to perform a comprehensive database search (i.e. optimal global alignment). If TRUE, performs quick search equivalent to setting VSEARCH parameters "-maxaccepts 100 -maxrejects 100". If FALSE, performs comprehensive search equivalent to setting VSEARCH parameters "-maxaccepts 0 -maxrejects 0"

db

(Default="16S\_bac") Select database option(s) including "16S" (for searching against the NCBI Refseq targeted loci 16S rRNA database), "ITS" (for searching against the NCBI Refseq targeted loci ITS database. For combined databases in cases where input sequences are derived from bacteria and fungi, select "16SITS". Setting to anything other than db=NULL or db="custom" causes 'db.path' parameter to be ignored.

db\_path

Path of FASTA-formatted database sequence file. Ignored if 'db' parameter is set to anything other than NULL or "custom".

vsearch\_path

Path of VSEARCH software if manually downloaded in a custom directory. If NULL (Default), will attempt automatic download.

iddef

Set pairwise identity definition as per VSEARCH definitions (Default=2, and is recommended for highest taxonomic accuracy) (0) CD-HIT definition: (matching columns) / (shortest sequence length). (1) Edit distance: (matching columns) / (alignment length). (2) Edit distance excluding terminal gaps (default definition). (3) Marine Biological Lab definition counting each gap opening (internal or terminal) as a single mismatch, whether or not the gap was extended: 1.0-((mismatches + gap openings)/(longest sequence length)). (4) BLAST definition, equivalent to –iddef 1 for global pairwise alignments.

phylum\_threshold

Percent sequence similarity threshold for phylum rank demarcation

class\_threshold

Percent sequence similarity threshold for class rank demarcation

order\_threshold

Percent sequence similarity threshold for order rank demarcation family\_threshold

Percent sequence similarity threshold for family rank demarcation genus\_threshold

Percent sequence similarity threshold for genus rank demarcation species\_threshold

Percent sequence similarity threshold for species rank demarcation

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#### Value

Returns taxonomic classification table of class isoTAX. Default taxonomic cutoffs for phylum (75.0), class (78.5), order (82.0), family (86.5), genus (94.5), and species (98.7) demarcation are based on Yarza et al. 2014, Nature Reviews Microbiology (DOI:10.1038/nrmicro3330)

### See Also

```
isoQC, isoLIB, search_db
```

# **Examples**

```
#Set path to directory containing example .ab1 files
fpath1 <- system.file("extdata/abif_examples/rocket_salad", package = "isolateR")

#Step 1: Run isoQC function with default settings
isoQC.S4 <- isoQC(input=fpath1)

#Step 2: Run isoTAX function with default settings
fpath2 <- file.path(fpath1, "isolateR_output/01_isoQC_trimmed_sequences_PASS.csv")
isoTAX.S4 <- isoTAX(input=fpath2)
#Show summary statistics
isoTAX.S4</pre>
```

make\_fasta

Convert CSV file containing sequences to FASTA format

# Description

This function extracts sequences from a table in CSV format and converts them to FASTA format. Requires two columns, one with sequences and one with sequence names.

### Usage

```
make_fasta(
  csv_file = NULL,
  col_names = "ID",
  col_seqs = "Sequence",
  output = "output.fasta")
```

# **Arguments**

| csv_file  | Filename (or path and filename if not in working directory) of the table from which you would like to generate a FASTA file. |
|-----------|--|
| col_names | Column name with the unique names/identifiers. (Default="ID")  |
| col_seqs  | Column name with the sequences. (Default="Sequence")   |
| output    | Desired filename for output FASTA file (Default = "output.fasta")  |

### Value

Returns sequences in FASTA format.

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#### **Examples**

```
#Set path to directory containing example .ab1 files
fpath1 <- system.file("extdata/abif_examples/rocket_salad", package = "isolateR")

#Run isoQC function with default settings to generate CSV file
isoQC.S4 <- isoQC(input=fpath1)

#Set path of CSV output file from isoQC step
csv.path <- file.path(fpath1, "isolateR_output/01_isoQC_trimmed_sequences_PASS.csv")

#Run make_fasta function
make_fasta(csv_file= csv.path, col_names="filename", col_seqs="seqs_trim", output="output.fasta")</pre>
```

make\_tree

Generate a phylogenetic tree from an isoLIB output file

# **Description**

This script will help the user make a simple phylogenetic tree from a strain library. It will allow the user to colour the tree by taxonomic rank only. See ggtree documentation for more information on customization options available.

### Usage

```
make_tree(input = NULL)
```

# Arguments

input

Full path to isoLIB strain library output file in .CSV format.

# Value

Returns a ggtree class object

### See Also

isoLIB

### **Examples**

```
#Set path to directory containing example .ab1 files
fpath1 <- system.file("extdata/abif_examples/rocket_salad", package = "isolateR")

#Step 1: Run isoQC function with default settings
isoQC.S4 <- isoQC(input=fpath1)

#Step 2: Run isoTAX function with default settings
fpath2 <- file.path(fpath1, "isolateR_output/01_isoQC_trimmed_sequences_PASS.csv")
isoTAX.S4 <- isoTAX(input=fpath2)

#Step 3: Run isoLIB function with default settings
fpath3 <- file.path(fpath1, "isolateR_output/02_isoTAX_results.csv")
isoLIB.S4 <- isoLIB(input=fpath3)</pre>
```

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```
#Step 4: Make a tree from isoLIB output CSV file
fpath4 <- file.path(fpath1, "isolateR_output/03_isoLIB_results.csv")
make_tree(input= fpath4)</pre>
```

method-isoLIB

setMethod functions for isoLIB

# **Description**

Initiation of isoLIB functions.

# Usage

```
## S4 method for signature 'missing'
isoLIB(
 input = NULL,
 old_lib_csv = NULL,
 method = "dark_mode"
 group_cutoff = 0.995,
 keep_old_reps = TRUE,
 export_html = TRUE,
 export_csv = TRUE,
 include_warnings = TRUE,
 vsearch_path = NULL,
 phylum_threshold = 75,
 class_threshold = 78.5,
 order_threshold = 82,
 family_threshold = 86.5,
 genus_threshold = 94.5,
  species\_threshold = 98.7
```

method-isoQC

setMethod functions for isoQC

# **Description**

Initiation of isoQC functions.

# Usage

```
## S4 method for signature 'missing'
isoQC(
  input = NULL,
  export_html = TRUE,
  export_csv = TRUE,
  export_fasta = TRUE,
  export_fasta_revcomp = FALSE,
  verbose = FALSE,
  exclude = NULL,
```

22 S4\_to\_dataframe

```
min_phred_score = 20,
min_length = 200,
sliding_window_cutoff = NULL,
sliding_window_size = 15,
date = NULL,
files_manual = NULL
)
```

method-isoTAX

 $setMethod\ functions\ for\ isoTAX$ 

### **Description**

Initiation of isoTAX functions.

# Usage

```
## S4 method for signature 'missing'
isoTAX(
  input = NULL,
  export_html = TRUE,
  export_csv = TRUE,
  export_blast_table = FALSE,
  quick_search = FALSE,
  db = "16S_bac",
  db_path = NULL
  vsearch_path = NULL,
  iddef = 2,
  phylum_threshold = 75,
  class_threshold = 78.5,
  order_threshold = 82,
  family_threshold = 86.5,
  genus_threshold = 94.5,
  species_threshold = 98.7
)
```

S4\_to\_dataframe

Converts S4 objects (isoQC, isoTAX, or isoLIB) to dataframe

# Description

 $Helper\ function\ to\ convert\ S4\ class\ objects\ (\verb"isoQC", \verb"isoTAX", or \verb"isoLIB")\ to\ data frame$ 

### Usage

```
S4_to_dataframe(obj)
```

# **Arguments**

obj

S4 object generated from isoQC, isoTAX, or isoLIB steps

23 sanger\_consensus

#### Value

Returns a dataframe containing sequence information in columns.

sanger\_consensus

Overlap multiple paired Sanger sequences in batch.

# **Description**

This function loads in the CSV results table from isoQC and merges related sequences based on user input. Original file names before isoQC step need to have a common prefix and differentiating suffixes. (e.g. SAMPLE\_01\_F.ab1, SAMPLE\_01\_R.ab1). After aligning paired sequences, the consensus sequence is extracted and priority is given to the read with higher quality. Phred quality scores are reassigned in the final output table in a basic way by taking the mean of both input

Note: This function is designed to be used after the isoQC step and before the isoTAX step.

### Usage

```
sanger_consensus(input = NULL, suffix = "_F.ab1|_R.ab1")
```

#### **Arguments**

input

Path of CSV output file from isoQC step.

suffix

Regex-friendly suffix for denoting filename groupings. Default="\_F.ab1|\_R.ab1" for the common scenario of Sanger sequencing a marker gene in forward and reverse. Direction of sequences including reverse complements will be auto-

matically detected.

### Value

Returns merged pairs of Sanger sequences in FASTA format.

#### See Also

isoQC, isoTAX

search\_db

Perform global alignment pairwise identity search using VSEARCH and type strain database of interest.

# **Description**

Performs global alignment between FASTA-formatted query sequences and the specified database of interest. Uses the Needleman-Wunsch algorithm by wrapping the -usearch\_global command implemented in VSEARCH.

24 search\_db

#### **Usage**

```
search_db(
  query.path = NULL,
 uc.out = "VSEARCH_output.uc",
 b6.out = "VSEARCH_output.b6o",
  path = getwd(),
  quick_search = FALSE,
  db = NULL,
  db_path = NULL,
  vsearch_path = NULL,
  keep_temp_files = FALSE,
  iddef = 2
)
```

#### **Arguments**

Path of FASTA-formatted query sequence file. query.path

uc.out Path of UC-formatted results output table.

b6.out Path of blast6-formatted results output table.

path Working path directory (Default is set to current working directory via 'getwd()'

quick\_search (Default=FALSE) Whether or not to perform a comprehensive database search

(i.e. optimal global alignment). If TRUE, performs quick search equivalent to setting VSEARCH parameters "-maxaccepts 100 -maxrejects 100". If FALSE, performs comprehensive search equivalent to setting VSEARCH parameters "maxaccepts 0 –maxrejects 0" Note: This option is provided for convenience and rough approximation of taxonomy only, set to FALSE for accurate % pairwise

identity results.

Optional: Select any of the standard database option(s) including "16S" (for searching against the NCBI Refseq targeted loci 16S rRNA database), "ITS" (for searching against the NCBI Refseq targeted loci ITS database. For combined databases in cases where input sequences are dervied from bacteria and fungi, select "16SITS". Setting to anything other than db=NULL or db="custom"

causes 'db.path' parameter to be ignored.

Path of VSEARCH software if manually downloaded in a custom directory. If vsearch\_path

NULL (Default), will attempt automatic download.

keep\_temp\_files

Toggle (TRUE/FALSE). If TRUE, temporary .uc and .b6o output files are kept from VSEARCH -uc and -blast6out commands, respectively. If FALSE, tem-

porary files are removed.

iddef Set pairwise identity definition as per VSEARCH definitions (Default=2, and is

> recommended for highest taxonomic accuracy) (0) CD-HIT definition: (matching columns) / (shortest sequence length). (1) Edit distance: (matching columns) / (alignment length). (2) Edit distance excluding terminal gaps (default definition). (3) Marine Biological Lab definition counting each gap opening (internal or terminal) as a single mismatch, whether or not the gap was extended: 1.0-((mismatches + gap openings)/(longest sequence length)). (4) BLAST defini-

tion, equivalent to –iddef 1 for global pairwise alignments.

Path of FASTA-formatted database sequence file. Ignored if 'db' parameter is

set to anything other than "custom"

db

db.path

search\_db 25

#### Value

Returns a dataframe matching the UC-formatted output table from VSEARCH. Query sequences are automatically added to the final column. Summary of column information. See VSEARCH documentation for more details.

- V1 = Record type of hit (H) or no hit (N)
- V2 = Ordinal number of the target sequence (based on input order, starting from zero). Set to '\*' for N.
- V3 = Sequence length. Set to '\*' for N.
- V4 = Percentage of similarity with the target sequence. Set to '\*' for N.
- V5 = Match orientation + or -... Set to '.' for N.
- V6 = Not used, always set to zero for H, or '\*' for N.
- V7 = Not used, always set to zero for H, or '\*' for N.
- V8 = Compact representation of the pairwise alignment using the CIGAR format (Compact Idiosyncratic Gapped Alignment Report): M (match/mismatch), D (deletion) and I (insertion). The equal sign '=' indicates that the query is identical to the centroid sequence. Set to '\*' for N.
- V9 = Label of the query sequence. Equivalent to 'filename' slot of isolateR class objects (e.g. isoQC, isoTAX, isoLIB).
- V10 = Label of the target centroid sequence. Set to '\*' for N.

### See Also

isoTAX

# **Examples**

```
#Set path to directory containing example .ab1 files
fpath1 <- system.file("extdata/abif_examples/rocket_salad", package = "isolateR")

#Run isoQC function with default settings
isoQC.S4 <- isoQC(input=fpath1)

#Set path of CSV output file containing PASS sequences from isoQC step
fasta.path <- "01_isoQC_trimmed_sequences_PASS.fasta"

#Set paths
output.path <- file.path(fpath1, "isolateR_output")

#Run search_db function
uc.df <- search_db(query.path=fasta.path, path=output.path, quick_search=TRUE, db="16S")

#Inspect results
uc.df[1:10,1:10]</pre>
```

26 *valid\_tax\_check* 

show

Generic show method for S4 class objects

# **Description**

Generic show method for S4 class objects.

# Usage

```
## S4 method for signature 'isoQC'
show(object)

## S4 method for signature 'isoTAX'
show(object)

## S4 method for signature 'isoLIB'
show(object)
```

valid\_tax\_check

Validate species name via API client of LPSN

# Description

This function will determine if each species in a CSV file is validly published or not. Result file will be a CSV with the results appended to the input data. This function requires the user to have an LPSN API account setup. For more details and to register, see here: https://api.lpsn.dsmz.de/)

# Usage

```
valid_tax_check(input = NULL, col_species = "rank_species", export_csv = TRUE)
```

# **Arguments**

input CSV file path. Expects full path if CSV file is not in the current working direc-

tory.

col\_species Specify the column containing the binomial species names (e.g. "Akkermansia

muciniphila")

export\_csv Toggle (TRUE/FALSE). Set TRUE to automatically write .CSV file of results to

current directory. (Default=TRUE)

### Value

Returns a CSV saved in working directory

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