R documentation

of all in '.'

June 6, 2024

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mjolnir1 RAN

RAN: Reads Allotment in N samples

Description

RAN function will prepare the FASTQ raw data for each sample with the sample tags and primers removed. The R1 output files will contain all forward sequences and R2 the reverse sequences. Please, read the Details section before running.

Usage

```
\label{eq:mjolnir1_RAN} $$ mjolnir1_RAN($ R1_filenames = "", $$ lib_prefix = "", $$ experiment = NULL, $$ primer_F = "GGWACWRGWTGRACWNTNTAYCCYCC", $$ primer_R = "TANACYTCNGGRTGNCCRAARAAYCA", $$ cores = 1, $$
```

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```
\begin{array}{l} R1\_motif = "\_R1", \\ R2\_motif = "\_R2", \\ multilane = FALSE, \\ tag\_error = 0, \\ primer\_error = 0.1, \\ \dots \\ ) \end{array}
```

Arguments

$R1_filenames$	Character vector with the names of the forward fastq or fastq.gz files. Only needed for multiplexed libraries.
lib_prefix	Character vector. Acronym for each sequencing library. This acronym must be of 4 characters in capital letters. Do not mix up library and experiment acronyms. The latter will be required in following steps. However they can be the same.
$\exp { m eriment}$	Character string. Acronym for the experiment. This acronym must be of 4 characters in capital letters. Do not mix up library and experiment acronyms. However they can be the same.
$primer_F$	Character string of the Forward primer. Necessary when samples are already demultiplexed.
$primer_R$	Character string of the Reverse primer. Necessary when samples are already demultiplexed.
cores	Numeric. Number of parts into which the input files will be split for the parallel processing of the FREYJA function.
$R1_motif$	Character string that distinguishes the forward line file from the reverse.
$R2_motif$	Character string that distinguishes the reverse line file from the forward.
$\operatorname{multilane}$	Logical. If FALSE, the function will consider that the each library has only one sequencing lane. If TRUE, only one library is processed at a time.
${ m tag_error}$	Numeric. From 0 (no errors allowed) to 1. will determine the proportion of nucleotides that can be different in the sample tags.
primer_error	Numeric. From 0 (no errors allowed) to 1. will determine the proportion of nucleotides that can be different in the primers.

Details

RAN considers the following scenarios:

- **Scenario 1**. The samples are multiplexed in a library/libraries (it is possible to run multiple libraries at once from a single experiment). Each library has only two raw .fastq(.gz), R1 and R2. For each library there must be only one ngsfile (see below) and for the whole experiment only one metadata file (see below). mjolnir_agnomens (the standard name of the sample during the pipeline) must be unique, it must not be the same in different ngsfiles. *lib_prefix* and *R1_filenames* must be the same length.
- **Scenario 2**. The samples are multiplexed but for each library we have more than two raw data files. For example, when your library has been sequenced across multiple lanes on a Novaseq. In

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this case, RAN has to be run separately for each library and the option *multilane* must be set to TRUE. In this case, *lib_prefix* can only be of length one (only one library), but the *R1_filenames* can be longer.

- **Scenario 3**. The samples are demultiplexed but the primer remains. In this case RAN will trimm the primers and separate the sequences into fwd and rev files.

Starting with multiplexed libraries:

Files required:

- ngsfilter file. Needed only for multiplexed libraries For each library, a ngsfilter file is needed and must be named ngsfilter_library identifier>.tsv. This must contain five tab-separated columns and no header. The first column with the library identifier (four charachter identifier), the second with the mjolnir_agnomens, the third with the sample tags, the fourth with the forward primers and the fifth with the reverse primers.
- *metadata file*. A metadata file containing at least two columns required, "original_samples" and "mjolnir_agnomens" ("fastq_name_R1" for demultiplexed libraries, see below), and named as **<ex-periment identifier> metadata.tsv**.

Important: when the same library has different sequencing lanes (NovaSeq), RAN has to be run separately for each library and the option *multilane* must be set to TRUE.

Starting with demultiplexed samples:

If samples are already demultiplexed, primers need to be set (*primer_F* & *primer_R*) or LERAY_XT primers for COI will be used by default.

Files required:

- metadata file. RAN will read the names of each individual R1 fastq files (full name including extension) from a column in the **<experiment identifier>_metadata.tsv** file, called "fastq_name_R1". In the metadata table, each sample in the "original_samples" column must have a their corresponding fastq_name_R1 and "mjolnir_agnomen" (LIBX_sample_XXX, i.e LIBA_sample_001). If the "fastq_name_R1" column is not in the metadata, RAN will use the lib_prefix.

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mjolnir2_FREYJA FREYJA: Filtering of Reads, Enrollment, Yoke-reads Joining and Alignment

Description

FREYJA will use OBITools3 commands to merge paired-end reads, trim primer sequences, filter by length, split sequences per sample and dereplicate within each sample.

Usage

```
\begin{split} & \text{mjolnir2\_FREYJA(} \\ & \text{experiment} = \text{NULL,} \\ & \text{cores} = 1, \\ & \text{Lmin} = 299, \\ & \text{Lmax} = 320, \\ & \text{score\_obialign} = 40, \\ & \text{R1\_motif} = "\_\text{R1",} \\ & \text{R2\_motif} = "\_\text{R2",} \\ & \text{remove\_DMS} = \text{T,} \\ & \text{run\_on\_tmp} = \text{F,} \\ & \dots \\ & ) \end{split}
```

Arguments

$\operatorname{exp}\operatorname{eriment}$	Character string. Acronym for the experiment. This acronym must be of 4 characters in capital letters. Do not mix up library and experiment acronyms. However they can be the same.
cores	Numeric. Number of threads for parallel processing.
Lmin	Numeric. Minimum bp length for a sequence to be accepted.
Lmax	Numeric. Maximum bp length for a sequence to be accepted.
$R1_motif$	Character string that distinguish the forward line file from the reverse.
$ m R2_motif$	Character string that distinguish the reverse line file from the forward.
${\rm remove_DMS}$	Logical. If TRUE, it will delete all obidms objects that are created during the process. This can save a lot of hard disk space. The FALSE option is useful for developing and debugging.
$\operatorname{run_on_tmp}$	Logical. If TRUE, the obidms objects will be created in the /tmp location. This increases the speed as the communication within the processor and the object that is being edited all the time is faster. However, this method will consume much of the /tmp memory and it is recommended to have three to four times the memory available in the /tmp directory than the original forward files and remove_DMS=T
$score_obilign$	Numeric. Minimum quality threshold to retain a sequence in the quality filtering after pairalignment.

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Details

Input file fastq files are expected to be without primers sequence and all forward sequences in the R1 file and all reverse sequences in the R2 file.

Examples

```
library(mjolnir)
# Define input fastq files (only names of R1 files are needed)
R1 filenames <- c("ULO1 R1.fastq.gz", "ULO2 R1.fastq.gz", "ULO3 R1.fastq.gz",
             "ULO4 R1.fastq.gz")
# Input identifiers for the individual libraries to be used.
# It should be a 4-character name, matching the information in the
# ngsfilter files.
lib prefixes <- c("ULO1", "ULO2", "ULO3", "ULO4")
# experiment identifier
experiment <- 'ULOY'
# Enter number of cores to be used in parallel.
cores < -7
mjolnir1 RAN(R1 filenames, lib prefix = lib prefixes, experiment = experiment,
         cores = cores, R1 motif = "R1", R2 motif = "R2")
# Run FREYJA
mjolnir2 FREYJA(experiment = experiment, cores = cores, Lmin=299, Lmax=320)
```

mjolnir3 HELA

HELA: Hierarchical Elimination of Lurking Artifacts

Description

This function uses the uchime_denovo algorithm implemented in VSEARCH to remove chimaeric sequences from the dataset.

Usage

```
mjolnir3 HELA(experiment = NULL, cores = 1, ...)
```

Arguments

experiment Character string. Acronym for the experiment. This acronym must be of 4

characters in capital letters. Do not mix up library and experiment acronyms.

However they can be the same.

cores Numeric. Number of threads for parallel processing.

Details

HELA works in a sample-by-sample basis. HELA will process all individual fasta files in the current folder matching the pattern EXPX_XXXX_sample_XXX.fasta being EXPX the acronym set by experiment parameter. This allows for parallel computing, significantly decreasing calculation times.

Examples

```
library(mjolnir)
# Define input fastq files (only names of R1 files are needed)
R1 filenames <- c("ULO1 R1.fastq.gz", "ULO2 R1.fastq.gz", "ULO3 R1.fastq.gz",
             "ULO4 R1.fastq.gz")
# Input identifiers for the individual libraries to be used.
# It should be a 4-character name, matching the information in the
# ngsfilter files.
lib prefixes <- c("ULO1", "ULO2", "ULO3", "ULO4")
# experiment identifier
experiment <- 'ULOY'
# Enter number of cores to be used in parallel.
cores < -7
mjolnir1 RAN(R1 filenames, lib prefix = lib prefixes, experiment = experiment,
         cores = cores, R1\_motif = "\_R1", R2\_motif = "\_R2")
# Run FREYJA
mjolnir2 FREYJA(experiment = experiment, cores = cores, Lmin=299, Lmax=320)
# Run HELA
mjolnir3 HELA(experiment = experiment, cores = cores)
```

mjolnir4 ODIN

ODIN: OTU Delimitation Inferred by Networks

Description

ODIN performs MOTU clustering and/or denoising. It is one of the main steps of MJOLNIR.

Usage

```
\begin{split} & \text{mjolnir4\_ODIN(} \\ & \text{experiment} = \text{NULL,} \\ & \text{cores} = 1, \\ & \text{d} = 13, \\ & \text{min\_reads\_MOTU} = 2, \\ & \text{min\_reads\_ESV} = 2, \\ & \text{min\_relative} = 1/50000, \end{split}
```

```
blank relative = 0.1,
metadata table = "",
blank col = "BLANK",
blank tag = TRUE,
alpha = 4,
entropy = c(0.47, 0.23, 1.02, 313),
algorithm = "DnoisE SWARM",
run dnoise = TRUE,
remove singletons = NULL,
remove DMS = TRUE,
```

Arguments

Character string. Acronym for the experiment. This acronym must be of 4 experiment

characters in capital letters. Do not mix up library and experiment acronyms.

However they can be the same.

cores Numeric. Number of threads for parallel processing.

d Numeric value for d parameter of SWARM that refers to the maximum number

of differences between two sequences to be linked in the same cluster

 $\min_{\text{reads}} MOTU$

Numeric. Minimum number of reads that a MOTU needs to have to be retained.

min reads ESV

Numeric. Minimum number of reads that an ESV needs to have to be retained. Also works works for non-denoised sequences. In the case of algorithms involv-

ing SWARM, the removal is performed before SWARM.

min relative Number of the minimum relative abundance for a unit in the sample to be re-

tained.

blank relative Relative abundance threshold for a unit to be removed if the total abundance in

the blank/neg/control is higher than this value in terms of relative abundance of

the total reads in all samples (see Details).

metadata table tsv table, if not specified, the file must be named <EXPX> metadata.tsv. This

table must have: a column named "mjolnir_agnomens" with the names given to the samples during the pipeline in FREYJA; a column named "original_samples" with the samples names that will be given to the samples at the end of the pipeline; and a column with the name specified in in the "blank_col" parameter ("BLANK" by default) where blanks, negatives and controls are tagged with a

flag specified in the "blank_tag" parameter (T by default).

blank col Column name of the blank column in the "metadata_table"

blank tag Unique flag to tag the samples that are blank/neg/controls in the "blank_col"

alpha Numeric. Alpha value for DnoisE to run.

entropy Logical, numeric or character vector specifying whether to run DnoisE with

entropy correction and how.

a) c(0.47,0.23,1.02,313) - formulation refers to the entropy values for the first, second and third position in the codon and the length of the main sequence length expected. See Antich et al. 2022 for further details.

b) FALSE - this will disable the entropy correction. Recommended for non coding markers

c) c("auto_sample",313) - this will compute the entropy values for 313 (plus or minus multiple of 3) bp within DnoisE and use them to perform the entropy correction

d) c("auto_dataset") - this will compute the entropy values for all the dataset and all sequence lengths and use the main sequence length's values for the entropy correction

algorithm

Character. It specifies the algorithm to obtain MOTUs and/or ESVs. Ther are four options:

a)"DnoisE_SWARM" - This option will run DnoisE before SWARM. This option is the best choice for highly diverse data sets so it will reduce the computation time of SWARM. It also allows the analysis of metaphylogeographical approaches and retrieves denoised and quality filter fasta files for each sample that can be used for other experiment without previous run of RAN, FREYJA and HELA. It will result on a table of MOTUs and the ESV clustered into them.

b)"SWARM" - This option will run only SWARM to obtain a MOTU table.

c)"SWARM_DnoisE" - This option will run SWARM before DnoisE. This option responds to a philosophical point of view where the denoising of the sequences have to be performed within MOTUs so closer sequences are compared and to avoid that high abundant sequences from different MOTUs absorb sequences (and thus their reads) from different MOTUs. However, there are no major differences between this option and the "DnoisE_SWARM" option. It allows the analysis of metaphylogeographical and will result on a table of MOTUs and the ESV clustered into them.

d)"DnoisE" - This option will run only DnoisE. This option will retrieve quality filter fasta files for each sample that can be used for other experiment without previous run of RAN, FREYJA and HELA.

 run_dnoise

Logical. In the case of the algorithm='DnoisE_SWARM' there is the option of not running the DnoisE if it has been already run for a previous experiment. The denoised and filtered fasta files are needed.

 $remove_DMS$

Logical. If TRUE, it will delete all obidms objects that are created during the process. This can save a lot of hard disk space. The FALSE option is useful for developing and debugging.

Details

The function mjolnir4_ODIN() uses the four different strategies to delimit MOTUs and/or ESVs. This strategies are set with the algorithm parameter:

a)"DnoisE_SWARM", b)"SWARM", c)"SWARM_DnoisE" and d)"DnoisE".

In short, DnoisE refers to the denoising process with DnoisE to obtain ESV and SWARM to a clustering process with SWARM to obtain MOTUs. DnoisE is a software to merge spurious sequences into their "mothers" (see Antich et al. 2022) to obtain Exact (also Amplicon) Sequence variants.

DnoisE is an open source and parallelizable alternative to Unoise that allows to introduce an entropy correction based on the different entropies of each position in the codon of coding genes. This is highly recommended for markers as COI for which this program was intended. However, with the entropy=FALSE parameter, this programs performs the same denoising procedure as described for Unoise3. SWARM is an algorithm to delimit MOTUs, based on linkage-networks created by step-by-step agregation. This clustering algorithm is not based on an arbitrary, constant, absolute identity threshold. Conversely, SWARM is based on an iterative aggregation of sequences that differ less than a given distance d. This strategy results into linkage-networks of different sizes, which can have different effective values for within-MOTU identity threshold, depending on the complexity of the natural variability of the sequences present in the sample. This procedure is very convenient in the case of hypervariable metabarcoding markers, such as COI, which usually feature extremely high levels of natural diversity, in addition to the random sequencing errors. Dereplication step takes place after joining all samples into the same file before SWARM in algorithms a, b and c and after DnoisE in algorithms a and d

NEW: now ODIN performs most of the filters that were applied in RAGNAROC. The idea is to retrieve an output file for each sample that is independent from the rest of the samples (blank and neg corrected; total reads filtered) and with denoised ASV/ESV using the name

"<mjolnir_agnoment>_ODIN_ESV_<original_name>.fasta.".

To do so these are the the different steps of this function:

«D -> "DnoisE";

DS -> "DnoisE_SWARM"

 $S \rightarrow "SWARM";$

SD -> "SWARM_DnoisE";

SaD (SWARM after DnoisE) -> "DnoisE_SWARM" & run_dnoise = F»

0: define variables and load metadata table

1: D and DS -> denoise the fasta files

2: D,DS,SD,S,SaD -> cat all fasta

3: D,DS,SD,S,SaD -> dereplicate

4: D,DS,SD,S,SaD -> annotate new names

filter1A: D,DS -> blank relative abundances filter AS MOTU

filter2A: D,DS -> min relative abundances filter WS ESV

filter3A: D,DS -> remove nletons # improves SWARM for DS

filter1B: SD,S,SaD -> remove nletons # this will improve SWARM performance # apply this to SaD just in case

5: D,DS,SD,S,SaD -> export csv file of sequences (if denoised, they are ESV)

6: D,DS -> export fasta DnoisEd & blank filt

7: D -> create fasta files for taxonomic assignment

8: DS,SD,S,SaD -> do SWARM

filter2B: SD,S -> blank relative abundances filter AS MOTU

filter3B: SD,S -> min relative abundances filter WS MOTU

filter4B: SD,S -> remove nletons AS MOTUs # remove artefacts

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9: DS,SD,S,SaD -> create csv files of MOTUs and ESV or seqs.

10: DS,SD,S,SaD -> Also create fasta files for taxonomic assignment

11: SD -> run DnoisE over the csv files

Blank filter: remove any MOTU for which abundance in the blank or negative controls is higher than "blank_relative" of its total read abundance and remove blank and NEG samples

Minimum relative abundance filter: Apply a minimum relative abundance threshold for each sample, setting to zero any abundance below "min_relative" of the total reads of this sample. It also applies a "min_reads" filter

```
library(mjolnir)
# Define input fastq files (only names of R1 files are needed)
R1_filenames <- c("ULO1_R1.fastq.gz", "ULO2_R1.fastq.gz", "ULO3_R1.fastq.gz",
            "ULO4 R1.fastq.gz")
# Input identifiers for the individual libraries to be used.
# It should be a 4-character name, matching the information in the
# ngsfilter files.
lib prefixes <- c("ULO1", "ULO2", "ULO3", "ULO4")
# experiment identifier
experiment <- 'ULOY'
# Enter number of cores to be used in parallel.
cores < -7
mjolnir1_RAN(R1_filenames, lib_prefix = lib_prefixes, experiment = experiment,
         cores = cores, R1 motif = "R1", R2 motif = "R2")
# Run FREYJA
mjolnir2 FREYJA(experiment = experiment, cores = cores, Lmin=299, Lmax=320)
# Run HELA
mjolnir3 HELA(experiment = experiment, cores = cores)
# Run ODIN
mjolnir4 ODIN(experiment = experiment, cores = cores, d = 13,
          min reads MOTU = 2, min reads ESV = 2,
          min relative = 1 / 50000, blank relative = 0.1,
          metadata_table = "", blank_col = "BLANK", blank_tag = TRUE,
          alpha = 4, entropy = c(0.47, 0.23, 1.02, 313),
          algorithm = "DnoisE SWARM")
```

mjolnir5_THOR

Description

This is a wrapper of ecotag

Usage

```
\begin{split} & \text{mjolnir5\_THOR}(\\ & \text{experiment} = \text{NULL},\\ & \text{cores} = 1,\\ & \text{tax\_db} = \text{NULL},\\ & \text{run\_ecotag} = \text{T},\\ & \text{vsearch} = \text{F},\\ & \text{remove\_DMS} = \text{T},\\ & \text{minimum\_circle} = 0.7,\\ & \dots \\ & ) \end{split}
```

Arguments

experiment Character string. Acronym for the experiment. This acronym must be of 4

characters in capital letters. Do not mix up library and experiment acronyms.

However they can be the same.

cores Numeric. Number of threads for parallel processing.

tax db Character string specifying de PATH to the reference database. In case of using

ecotag, the database must be an .obidms object. Also, when using ecotag it is important to have the following files in the same directory: order.complete.csv;

family_to_order.csv; genus_to_family.csv

run ecotag Logical. Whether to run (TRUE, default) the ecotag taxonomic assignment or

not (FALSE). The latter could take place when alternative taxonomic assigna-

ment software is applied but adding higher taxonomic ranks is desired.

vsearch Logical. Whether to run (TRUE) the vsearch taxonomic assignment or not

(FALSE, default). If vsearch has been selected, even if run ecotag=TRUE,

vsearch will be used instead of ecotag.

remove DMS Logical. If TRUE, it will delete all obidms objects that are created during the

process. This can save a lot of hard disk space. The FALSE option is useful for

developing and debugging.

minimum circle Numeric. For ecotag: Minimum identity considered for the assignment circle

(sequence is assigned to the LCA of all sequences within a similarity circle of the best matches; the threshold for this circle is the highest value between <CIRCLE_THRESHOLD> and the best assignment score found for the query sequence). Give value as a normalized identity, e.g. 0.95 for an identity of 95

for a match. Default: 0.7

Details

After assignment with ecotag, higher taxa at ranks higher than order are added from cust. The database used can be download or build using the NJORDR package (see https://github.com/adriantich/NJORDR-MJOLNIR3)

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For vsearch assignment, the database must be in fasta format and the taxonomy for the output CSV will be only in one single column.

```
library(mjolnir)
# Define input fastq files (only names of R1 files are needed)
R1 filenames <- c("ULO1 R1.fastq.gz", "ULO2 R1.fastq.gz", "ULO3 R1.fastq.gz",
            "ULO4 R1.fastq.gz")
# Input identifiers for the individual libraries to be used.
# It should be a 4-character name, matching the information in the
# ngsfilter files.
lib prefixes <- c("ULO1", "ULO2", "ULO3", "ULO4")
# experiment identifier
experiment <- 'ULOY'
# Enter number of cores to be used in parallel.
cores < -7
mjolnir1 RAN(R1 filenames, lib prefix = lib prefixes, experiment = experiment,
         cores = cores, R1 motif = "R1", R2 motif = "R2")
# Run FREYJA
mjolnir2 FREYJA(experiment = experiment, cores = cores, Lmin=299, Lmax=320)
# Run HELA
mjolnir3 HELA(experiment = experiment, cores = cores)
# Run ODIN
mjolnir4 ODIN(experiment = experiment, cores = cores, d = 13,
         \min reads MOTU = 2, \min reads ESV = 2,
         min relative = 1 / 50000, blank relative = 0.1,
         metadata_table = "", blank_col = "BLANK", blank_tag = TRUE,
         alpha = 4, entropy = c(0.47, 0.23, 1.02, 313),
         algorithm = "DnoisE SWARM")
# set the directory where the database is stored
tax db <- "~/taxo NCBI/DUFA COI"
# Run THOR
mjolnir5 THOR(experiment = experiment, cores = cores,
         tax db = tax db, run ecotag = T
```

mjolnir6_FRIGGA

Description

FRIGGA recombined the abundances from ODIN with the taxonomic-annotated TSV files from THOR

Usage

```
mjolnir6 FRIGGA(experiment = NULL, ...)
```

Arguments

experiment

Character string. Acronym for the experiment. This acronym must be of 4 characters in capital letters. Do not mix up library and experiment acronyms. However they can be the same.

Details

Input files must be called <EXPX>_THOR_annotated.tsv for the taxonomic-annotated TSV file and <EXPX>_ODIN_counts.tsv for the read counts of MOTUs or <EXPX>_ODIN_counts.tsv for ESVs. Output file is then called <EXPX>_FRIGGA.tsv

```
library(mjolnir)
# Define input fastq files (only names of R1 files are needed)
R1_filenames <- c("ULO1_R1.fastq.gz", "ULO2_R1.fastq.gz", "ULO3 R1.fastq.gz",
            "ULO4 R1.fastq.gz")
# Input identifiers for the individual libraries to be used.
# It should be a 4-character name, matching the information in the
# ngsfilter files.
lib prefixes <- c("ULO1", "ULO2", "ULO3", "ULO4")
# experiment identifier
experiment <- 'ULOY'
# Enter number of cores to be used in parallel.
cores < -7
mjolnir1 RAN(R1 filenames, lib prefix = lib prefixes, experiment = experiment,
         cores = cores, R1 motif = "R1", R2 motif = "R2")
# Run FREYJA
mjolnir2 FREYJA(experiment = experiment, cores = cores, Lmin=299, Lmax=320)
# Run HELA
mjolnir3 HELA(experiment = experiment, cores = cores)
# Run ODIN
mjolnir4 ODIN(experiment = experiment, cores = cores, d = 13,
         min reads MOTU = 2, min reads ESV = 2,
         min relative = 1 / 50000, blank relative = 0.1,
```

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```
metadata_table = "", blank_col = "BLANK", blank_tag = TRUE, alpha = 4, entropy = c(0.47, 0.23, 1.02, 313), algorithm = "DnoisE_SWARM")

# set the directory where the database is stored tax_db <- "~/taxo_NCBI/DUFA_COI"

# Run THOR
mjolnir5_THOR(experiment = experiment, cores = cores, tax_db = tax_db, run_ecotag = T)

# Run FRIGGA
mjolnir6_FRIGGA(experiment = experiment)
```

mjolnir7 LOKI

LOKI: LULU Overseeing with Kinship Identification

Description

LOKI is a convenient wrapper of LULU for the MJOLNIR3 metabarcoding pipeline.

Usage

```
mjolnir7 LOKI(experiment = NULL, min id = 0.84, ...)
```

Arguments

experiment

Character string. Acronym for the experiment. This acronym must be of 4 characters in capital letters. Do not mix up library and experiment acronyms. However they can be the same.

min id

Numeric. Equivalent to the -id option from vsearch -usearch global

From vsearch manual: Reject the sequence match if the pairwise identity is lower than min_id (value ranging from 0.0 to 1.0 included). The search process sorts target sequences by decreasing number of k-mers they have in common with the query sequence, using that information as a proxy for sequence similarity. That efficient pre-filtering also prevents pairwise alignments with weakly matching targets, as there needs to be at least 6 shared kmers to start the pairwise alignment, and at least one out of every 16 k-mers from the query needs to match the target. Consequently, using values lower than –id 0.5 is not likely to capture more weakly matching targets. The pairwise identity is by default defined as the number of (matching columns) / (alignment length - terminal gaps)

Details

LOKI starts from the combined dataset of abundances and taxonomy from the previous step (FRIGGA): <EXPX>_FRIGGA.tsv. A match list of representative MOTU sequences is created using VSEARCH and saved as a txt file. Then Units that are potential errors based on co-occurrence patterns are labelled and removed using LULU. The output is called <EXPX>_LOKI_Curated.tsv

```
library(mjolnir)
# Define input fastq files (only names of R1 files are needed)
R1 filenames <- c("ULO1 R1.fastq.gz", "ULO2 R1.fastq.gz", "ULO3 R1.fastq.gz",
            "ULO4 R1.fastq.gz")
# Input identifiers for the individual libraries to be used.
# It should be a 4-character name, matching the information in the
# ngsfilter files.
lib prefixes <- c("ULO1", "ULO2", "ULO3", "ULO4")
# experiment identifier
experiment <- 'ULOY'
# Enter number of cores to be used in parallel.
cores < -7
mjolnir1 RAN(R1 filenames, lib prefix = lib prefixes, experiment = experiment,
        cores = cores, R1 motif = "R1", R2 motif = "R2")
# Run FREYJA
mjolnir2 FREYJA(experiment = experiment, cores = cores, Lmin=299, Lmax=320)
# Run HELA
mjolnir3 HELA(experiment = experiment, cores = cores)
# Run ODIN
mjolnir4 ODIN(experiment = experiment, cores = cores, d = 13,
         min reads MOTU = 2, min reads ESV = 2,
         min relative = 1 / 50000, blank relative = 0.1,
         metadata table = "", blank col = "BLANK", blank tag = TRUE,
         alpha = 4, entropy = c(0.47, 0.23, 1.02, 313),
         algorithm = "DnoisE SWARM")
# set the directory where the database is stored
tax_db < "^-/taxo NCBI/DUFA COI"
# Run THOR
mjolnir5 THOR(experiment = experiment, cores = cores,
         tax db = tax db, run ecotag = T)
# Run FRIGGA
mjolnir6 FRIGGA(experiment = experiment)
# Run LOKI
mjolnir7 LOKI(experiment = experiment, min id = .84)
```

Description

Final step of the MJOLNIR3 pipeline to apply the last filters to the abundance data

Usage

```
mjolnir8 RAGNAROC(
 experiment = NULL,
 metadata table = "",
 output file = "",
 output file ESV = "",
 \min \text{ reads} = 0,
 remove bacteria = T,
 remove contamination = F,
 contamination file = "contaminants.txt",
 ESV within MOTU = T,
 remove numts = F,
 cores = 1,
```

Arguments

experiment

Character string. Acronym for the experiment. This acronym must be of 4 characters in capital letters. Do not mix up library and experiment acronyms. However they can be the same.

metadata table tsv table, if not specified, the file must be named <EXPX> metadata.tsv. This table must have: a column named "mjolnir_agnomens" with the names given to the samples during the pipeline in FREYJA; a column named "original samples" with the samples names that will be given to the samples at the end of the pipeline; and a column with the name specified in in the "blank_col" parameter ("BLANK" by default) where blanks, negatives and controls are tagged with a flag specified in the "blank_tag" parameter (T by default).

output file Character string specifying the outputfile name output file ESV

Character string specifying the outputfile name for ESVs abundances if required.

min reads Number of the minimum number of reads allowed for each MOTU/ESV or ESV within MOTU.

remove bacteria

Logical. If TRUE it will apply the bacteria removal filtering (see Details).

remove contamination

Logical. If TRUE it will apply the contamination removal filtering (see Details). contamination file

Character string specifying the name of the contamination file. (see Details)

```
ESV within MOTU
```

Logical. If TRUE this will take into account the ESV that were clustered into MOTUs in ODIN if algorithm was set to "DnoisE_SWARM" or "SWARM_DnoisE" and apply all filters to both data.

Details

RAGNAROC consists on different contamination removals and filtering steps as follows:

Removal of Bacteria: this removed the Units tagged as "Prokaryota" or "root" in the <EXPX>_LOKI_Cutated.tsv

Removal of contaminations: this step removes the taxa specified in the "contamination file"

NUMT removal: this step is design for the Leray-XT COI marker. It deletes all sequences that do not have a 313 (plus/minus multiple of 3 equivalent to a codon) bp length. Then removes sequences with stop codons and those metazoan sequences that do not translate for 5 specific conservative aminoacids.

```
library(mjolnir)
# Define input fastq files (only names of R1 files are needed)
"ULO4 R1.fastq.gz")
# Input identifiers for the individual libraries to be used.
# It should be a 4-character name, matching the information in the
# ngsfilter files.
lib prefixes <- c("ULO1", "ULO2", "ULO3", "ULO4")
# experiment identifier
experiment <- 'ULOY'
# Enter number of cores to be used in parallel.
cores < -7
mjolnir1 RAN(R1 filenames, lib prefix = lib prefixes, experiment = experiment,
        cores = cores, R1 motif = "R1", R2 motif = "R2")
# Run FREYJA
mjolnir2 FREYJA(experiment = experiment, cores = cores, Lmin=299, Lmax=320)
# Run HELA
mjolnir3 HELA(experiment = experiment, cores = cores)
# Run ODIN
mjolnir4 ODIN(experiment = experiment, cores = cores, d = 13,
         min reads MOTU = 2, min reads ESV = 2,
         min_relative = 1 / 50000, blank_relative = 0.1,
         metadata table = "", blank col = "BLANK", blank tag = TRUE,
         alpha = 4, entropy = c(0.47, 0.23, 1.02, 313),
         algorithm = "DnoisE SWARM")
# set the directory where the database is stored
tax db <- "~/taxo NCBI/DUFA COI"
```

18 numts

numts

numts

Description

Internal function of RAGNAROC to detect numt ESV within MOTUs

Usage

```
numts(datas, is metazoa = FALSE, motu, datas length)
```

Arguments

is metazoa Logical. whether if the MOTU is assigned to metazoa or not

motu Character. Name of the MOTU evaluated

datas length Numeric vector. Vector with the number of characters in each sequence

data Data Frame of the sequences in the MOTU.

Details

numt function will: 1: keep sequences of the same length as the seed 2: remove missaligned sequences 3: search for codon stops using all the mitochondrial genome codes with the Biostrings package and keep the one with less reads with stop codons. 4: if the MOTU is assigned to metazoans and the seed has 313 bp then check it the following aminoacid positions have the conserved ones in metazoans: 20 = "H"; 23 = "G"; 32 = "N"; 81 = "D"; 95 = "G"

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