# Package 'SQMtools'

July 3, 2019

Title Analyze results generated by the SqueezeMeta pipeline

Version 0.3.2

Description SqueezeMeta is a versatile pipeline for the automated analysis of metagenomics/metatranscriptomics data (http://github.com/jtamames/SqueezeMeta). This package provides functions loading SqueezeMeta results into R, filtering them based on different criteria, and visualizing the results using basic plots. The SqueezeMeta project (and any subsets of it generated by the different filtering functions) is parsed into a single object, whose different components (e.g. tables with the taxonomic or functional composition across samples, contig/gene abundance profiles) can be easily analyzed using other R packages such as vegan or DESeq2

Author Fernando Puente-Sánchez, Natalia García-García

Maintainer Fernando Puente-Sánchez <fpuente@cnb.csic.es>

**Depends** R (>= 3.4.0)

Imports reshape2, ggplot2

Suggests vegan, DESeq2

License GPLv3

**Encoding** UTF-8

LazyData true

RoxygenNote 6.1.1.9000

BugReports https://github.com/jtamames/SqueezeMeta/issues

URL https://github.com/jtamames/SqueezeMeta

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combineSQM

Combine several SQM objects

## **Description**

Combine an arbitrary number of SQM objects into a single SQM object.

#### Usage

```
combineSQM(..., tax_source = "orfs", trusted_functions_only = F,
  ignore_unclassified_functions = F, rescale_copy_number = T)
```

## Arguments

... an a

an arbitrary number of SQM objects

tax\_source

character. Features used for calculating aggregated abundances at the different taxonomic ranks. Either "orfs" or "contigs" (default "orfs"). If the objects being combined contain a subset of taxa or bins, this parameter can be set to TRUE.

 ${\tt trusted\_functions\_only}$ 

logical. If TRUE, only highly trusted functional annotations (best hit + best average) will be considered when generating aggregated function tables. If FALSE (default), best hit annotations will be used.

 $ignore\_unclassified\_functions$ 

logical. If FALSE, ORFs with no functional classification will be aggregated together into an "Unclassified" category. If TRUE, they will be ignored.

rescale\_copy\_number

logical. If TRUE, copy numbers with be recalculated using the RecA/RadA coverages in the subset. Otherwise, RecA/RadA coverages will be taken from the original object. By default it is set to TRUE, which means that the returned copy numbers will represent the average copy number per function *in the genomes of* 

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the selected bins or contigs. If any SQM objects that are being combined contain a functional subset rather than a contig/bins subset, this parameter should be set to FALSE.

#### Value

A SQM object

#### See Also

```
subsetFun, subsetTax
```

## **Examples**

```
data(Hadza)
# Select Carbohydrate metabolism ORFs in Bacteroidetes, and Amino acid metabolism ORFs in Proteobacteria
bact = subsetTax(Hadza, 'phylum', 'Bacteroidetes <phylum>')
bact.carb = subsetFun(bact, 'Carbohydrate metabolism')
proteo = subsetTax(Hadza, 'phylum', 'Proteobacteria')
proteo.amins = subsetFun(proteo, 'Amino acid metabolism')
bact.carb_proteo.amins = combineSQM(bact.carb, proteo.amins, rescale_copy_number=F)
```

exportTable

Export results in tabular format

## Description

This function is a wrapper for R's write.table function.

## Usage

```
exportTable(table, output_name)
```

## **Arguments**

table vector, matrix or data.frame. The table to be written.logical.

output\_name character. Name of the output file.

```
data(Hadza)
Hadza.iron = subsetFun(Hadza, 'iron')
# Write the taxonomic distribution at the genus level of all the genes related to iron.
exportTable(Hadza.iron$taxa$genus$percent, 'Hadza.ironGenes.genus.tsv')
# Now write the distribution of the different iron-related COGs (Clusters of Orthologous Groups) across samples.
exportTable(Hadza.iron$functions$COG$tpm, 'Hadza.ironGenes.COG.tsv')
# Now write all the information contained in the ORF table.
exportTable(Hadza.iron$orfs$table, 'Hadza.ironGenes.orftable.tsv')
```

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Hadza

Hadza hunter-gatherer gut metagenomes

## **Description**

Subset of 5 bins (and the associated contigs and genes) obtained from running SqueezeMeta on two gut metagenomic samples obtained from two hunter-gatherers of the Hadza ethnic group.

# Usage

```
data(Hadza)
```

## **Format**

```
A SQM object; see loadSQM.
```

## **Source**

```
SRR1927149, SRR1929485.
```

#### References

Rampelli *et al.*, 2015. Metagenome Sequencing of the Hadza Hunter-Gatherer Gut Microbiota. *Curr. biol.* **25**:1682-93 (PubMed).

# **Examples**

```
data(Hadza)
plotTaxonomy(Hadza, 'genus', rescale=T)
plotFunctions(Hadza, 'COG')
```

loadSQM

Load a SqueezeMeta project into R

## **Description**

This function takes the path to a project directory generated by SqueezeMeta (whose name is specified in the -p parameter of the SqueezeMeta.pl script) and parses the results into a SQM object

```
loadSQM(project_path, tax_mode = "allfilter")
```

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## **Arguments**

character, project directory generated by SqueezeMeta.

tax\_mode character, which taxonomic classification should be loaded? SqueezeMeta applies the identity thresholds described in Luo *et al.*, 2014. Use allfilter for applying the minimum identity threshold to all taxa (default) and prokfilter

## Value

SQM object containing the parsed project.

## **Prerequisites**

 Run SqueezeMeta! An example call for running it would be: /path/to/SqueezeMeta/scripts/SqueezeMeta.pl -m coassembly -f fastq\_dir -s samples\_file -p project\_dir

2. Generate tabular outputs with the sqm2tables.py script included in the path/to/SqueezeMeta/utils directory:

for applying the threshold to Bacteria and Archaea, but not to Eukaryotes.

/path/to/SqueezeMeta/utils/sqm2tables.py project\_dir project\_dir/results/tables

## The SQM object structure

The SQM object is a nested list which contains the following information:

lvl1	lvl2	lvl3	type	rows/names	columns	data
\$orfs	\$table		dataframe	orfs	misc. data	misc. data
	\$abund		numeric matrix	orfs	samples	abundances
	\$tpm		numeric matrix	orfs	samples	tpm
	\$seqs		character vector	orfs	(n/a)	sequences
	\$tax		character matrix	orfs	tax. ranks	taxonomy
\$contigs	\$table		dataframe	contigs	misc. data	misc. data
	\$abund		numeric matrix	contigs	samples	abundances
	\$tpm		numeric matrix	contigs	samples	tpm
	\$seqs		character vector	contigs	(n/a)	sequences
	\$tax		character matrix	contigs	tax. ranks	taxonomies
	\$bins		character matrix	contigs	bin. methods	bins
\$bins	\$table		dataframe	bins	misc. data	misc. data
	\$tpm		numeric matrix	bins	samples	tpm
	\$tax		character matrix	bins	tax. ranks	taxonomy
\$taxa	\$superkingdom	\$abund	numeric matrix	superkingdoms	samples	abundances
		\$percent	numeric matrix	superkingdoms	samples	percentages
	\$phylum	\$abund	numeric matrix	phyla	samples	abundances
		\$percent	numeric matrix	phyla	samples	percentages
	\$class	\$abund	numeric matrix	classes	samples	abundances
		\$percent	numeric matrix	classes	samples	percentages
	\$order	\$abund	numeric matrix	orders	samples	abundances
		\$percent	numeric matrix	orders	samples	percentages
	\$family	\$abund	numeric matrix	families	samples	abundances

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		\$percent	numeric matrix	families	samples	percentages
	\$genus	\$abund	numeric matrix	genera	samples	abundances
		\$percent	numeric matrix	genera	samples	percentages
	\$species	\$abund	numeric matrix	species	samples	abundances
		\$percent	numeric matrix	species	samples	percentages
\$functions	\$KEGG	\$abund	numeric matrix	KEGG ids	samples	abundances
		\$tpm	numeric matrix	KEGG ids	samples	tpm
		\$copy_number	numeric matrix	KEGG ids	samples	avg. copies
	\$COG	\$abund	numeric matrix	COG ids	samples	abundances
		\$tpm	numeric matrix	COG ids	samples	tpm
		\$copy_number	numeric matrix	COG ids	samples	avg. copies
	\$PFAM	\$abund	numeric matrix	PFAM ids	samples	abundances
		\$tpm	numeric matrix	PFAM ids	samples	tpm
		\$copy_number	numeric matrix	PFAM ids	samples	avg. copies
\$total_reads			numeric vector	samples	(n/a)	total reads
\$misc	<pre>\$project_name</pre>		character vector	(empty)	(n/a)	project name
	\$samples		character vector	(empty)	(n/a)	samples
	<pre>\$tax_names_long</pre>	\$superkingdom	character vector	short names	(n/a)	full names
		\$phylum	character vector	short names	(n/a)	full names
		\$class	character vector	short names	(n/a)	full names
		\$order	character vector	short names	(n/a)	full names
		\$family	character vector	short names	(n/a)	full names
		\$genus	character vector	short names	(n/a)	full names
		\$species	character vector	short names	(n/a)	full names
	\$tax_names_short		character vector	full names	(n/a)	short names
	<b>\$KEGG_names</b>		character vector	KEGG ids	(n/a)	KEGG names
	<b>\$COG_names</b>		character vector	COG ids	(n/a)	COG names

```
## Not run:
# (outside R)
/path/to/SqueezeMeta/scripts/SqueezeMeta.pl -p Hadza -f raw -m coassembly -s test.samples # Run SqueezeMeta on the
/path/to/SqueezeMeta/utils/sqm2tables.py Hadza Hadza/results/tables # Generate the tabular outputs! They must be
# now go into R
R
library(SQMtools)
Hadza = loadSQM("Hadza") # Where Hadza is the path to the SqueezeMeta output directory
## End(Not run)
data(Hadza)
# Which are the ten most abundant KEGG IDs in our data?
topKEGG = sort(rowSums(Hadza$functions$KEGG$tpm), decreasing=T)[1:10]
# Which functions do those KEGG IDs represent?
Hadza$misc$KEGG_names[topKEGG]
What is the relative abundance of the Gammaproteobacteria class across samples?
Hadza$taxa$class$percent['Gammaproteobacteria',]
# Which information is stored in the orf, contig and bin tables?
```

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```
colnames(Hadza$orfs$table)
colnames(Hadza$contigs$table)
colnames(Hadza$bins$table)
# What is the GC content distribution of my metagenome?
boxplot(Hadza$contigs$table[,'GC perc']) # Not weighted by contig length or abundance!
```

mostAbundant

Get the N most abundant rows from a numeric table

## **Description**

Return a subset of an input matrix or data frame, containing only the N most abundant rows, sorted. Alternatively, a custom set of rows can be returned.

## Usage

```
mostAbundant(data, N = 10, items = NULL, others = F, rescale = F)
```

## **Arguments**

data numeric matrix or data frame

N integer Number of rows to return (default 10).

items Character vector. Custom row names to return. If provided, it will override N

(default NULL).

others logical. If TRUE, an extra row will be returned containing the aggregated abun-

dances of the elements not selected with N or items (default FALSE).

rescale logical. Scale result to percentages column-wise (default FALSE).

### Value

A matrix or data frame (same as input) with the selected rows.

```
data(Hadza)
Hadza.carb = subsetFun(Hadza, 'Carbohydrate metabolism')
# Which are the 20 most abundant KEGG functions in the ORFs related to carbohydrate metabolism?
topCarb = mostAbundant(Hadza.carb$functions$KEGG$tpm, N=20)
# Now print them with nice names
rownames(topCarb) = paste(rownames(topCarb), Hadza.carb$misc$KEGG_names[rownames(topCarb)], sep='; ')
topCarb
We can pass this to any R function
heatmap(topCarb)
But for convenience we provide wrappers for plotting ggplot2 heatmaps and barplots
plotHeatmap(topCarb, label_y='TPM')
plotBars(topCarb, label_y='TPM')
```

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plotBars	Plot a barplot using ggplot2	
----------	------------------------------	--

## **Description**

Plot a ggplot2 barplot from a matrix or data frame. The data should be in tabular format (e.g. features in rows and samples in columns).

## Usage

```
plotBars(data, label_x = "Samples", label_y = "Abundances",
    label_fill = "Features", color = NULL, base_size = 11)
```

## **Arguments**

data	Numeric matrix or data frame.
label_x	character Label for the x axis (default "Samples").
label_y	character Label for the y axis (default "Abundances").
label_fill	character Label for color categories (default "Features").
color	Vector with custom colors for the different features. If empty, the default ggplot2 palette will be used (default NULL).
base_size	numeric. Base font size (default 11).

## Value

a ggplot2 plot object.

## See Also

plotTaxonomy for plotting the most abundant taxa of a SQM object; plotHeatmap for plotting a heatmap with arbitrary data; mostAbundant for selecting the most abundant rows in a dataframe or matrix. data(Hadza) sk = Hadza\$taxa\$superkingdom\$abund plotBars(sk, label\_y = 'Raw reads', label\_fill = 'Superkingdom')

plotFunctions	Heatmap of the most abundant functions in a SQM object

# Description

This function selects the most abundant functions across all samples in a SQM object and represents their abundances in a heatmap. Alternatively, a custom set of functions can be represented.

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## Usage

```
plotFunctions(SQM, fun_level = "KEGG", count = "tpm", N = 25,
  fun = c(), ignore_unclassified = T, gradient_col = c("ghostwhite",
  "dodgerblue4"), base_size = 11)
```

#### **Arguments**

SQM A SQM object.

fun\_level character. Either "KEGG", "COG" or "PFAM" (default "KEGG").

count character. Either "tpm" for TPM normalized values, "abund" for raw abun-

dances or "copy\_number" for copy numbers (default "tpm").

N integer Plot the N most abundant functions (default 25).

fun character. Custom functions to plot. If provided, it will override N (default

NULL).

ignore\_unclassified

logical. Don't include unclassified ORFs in the plot (default TRUE).

gradient\_col A vector of two colors representing the low and high ends of the color gradient

(default c("ghostwhite", "dodgerblue4")).

base\_size numeric. Base font size (default 11).

#### Value

a ggplot2 plot object.

## See Also

plotTaxonomy for plotting the most abundant taxa of a SQM object; plotBars and plotHeatmap for plotting barplots or heatmaps with arbitrary data.

## **Examples**

```
data(Hadza)
plotFunctions(Hadza)
```

plotHeatmap

Plot a heatmap using ggplot2

## **Description**

Plot a ggplot2 heatmap from a matrix or data frame. The data should be in tabular format (e.g. features in rows and samples in columns).

```
plotHeatmap(data, label_x = "Samples", label_y = "Features",
    label_fill = "Abundance", gradient_col = c("ghostwhite",
    "dodgerblue4"), base_size = 11)
```

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## Arguments

data	numeric matrix or data frame.
label_x	character Label for the x axis (default "Samples").
label_y	character Label for the y axis (default "Features").
label_fill	character Label for color scale (default "Abundance").
gradient_col	A vector of two colors representing the low and high ends of the color gradient (default c("ghostwhite", "dodgerblue4")).
base_size	numeric. Base font size (default 11).

## Value

A ggplot2 plot object.

#### See Also

plotFunctions for plotting the top functional categories of a SQM object; plotBars for plotting a barplot with arbitrary data; mostAbundant for selecting the most abundant rows in a dataframe or matrix.

## **Examples**

```
data(Hadza)
topPFAM = mostAbundant(Hadza$functions$PFAM$tpm)
topPFAM = topPFAM[rownames(topPFAM) != 'Unclassified',] # Take out the Unclassified ORFs.
plotHeatmap(topPFAM, label_x = 'Samples', label_y = 'PFAMs', label_fill = 'TPM')
data(Hadza)
phyla_percent = Hadza$taxa$phylum$percent
plotHeatmap(phyla_percent, label_y = 'Phylum', label_fill = 'Percentage')
```

plotTaxonomy

Barplot of the most abundant taxa in a SQM object

## **Description**

This function selects the most abundant taxa across all samples in a SQM object and represents their abundances in a barplot. Alternatively, a custom set of taxa can be represented.

```
plotTaxonomy(SQM, rank = "phylum", count = "percent", N = 15,
  tax = NULL, others = T, ignore_unclassified = F, rescale = F,
  color = NULL, base_size = 11)
```

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#### **Arguments**

SQM A SQM object.

rank Taxonomic rank to plot (default phylum).

count character. Either "percent" for percentages, or "abund" for raw abundances

(default "percent").

N integer Plot the N most abundant taxa (default 15).

tax character. Custom taxa to plot. If provided, it will override N (default NULL).

others logical. Collapse the abundances of least abundant taxa, and include the result

in the plot (default TRUE).

ignore\_unclassified

logical. Don't include unclassified contigs in the plot (default FALSE).

rescale logical. Re-scale results to percentages (default FALSE).

color Vector with custom colors for the different features. If empty, we will use our

own hand-picked pallete if N<=15, and the default ggplot2 palette otherwise

(default NULL).

base\_size numeric. Base font size (default 11).

#### Value

a ggplot2 plot object.

## See Also

plotFunctions for plotting the most abundant functions of a SQM object; plotBars and plotHeatmap for plotting barplots or heatmaps with arbitrary data.

## **Examples**

```
data(Hadza)
Hadza.amin = subsetFun(Hadza, 'Amino acid metabolism')
# Taxonomic distribution of amino acid metabolism ORFs at the family level.
plotTaxonomy(Hadza.amin, 'family')
```

RecA RecA/RadA recombinase

## Description

The recombination protein RecA/RadA is essential for the repair and maintenance of DNA, and has homologs in every bacteria and archaea. By dividing the coverage of functions by the coverage of RecA, abundances can be transformed into copy numbers, which can be used to compare functional profiles in samples with different sequencing depths. RecA-derived copy numbers are available in the SQM object (SQM\$functions\$<annotation\_type>\$copy\_number).

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## Usage

```
data(RecA)
```

#### **Format**

Character vector with the COG identifier for RecA/RadA.

#### **Source**

EggNOG Database.

## **Examples**

```
data(Hadza)
data(RecA)
### Let's calculate the average copy number of each function in our samples.
# We do it for COG annotations here, but we could also do it for KEGG or PFAMs.
COG.coverage = SQMtools::aggregate.fun(Hadza, 'COG', trusted_functions_only=T, ignore_unclassified_functions=F)
COG.copynumber = t(t(COG.coverage) / COG.coverage[RecA,]) # Sample-wise division by RecA tpm.
```

rowMaxs

Return a vector with the row-wise maxima of a matrix or dataframe.

## **Description**

Return a vector with the row-wise maxima of a matrix or dataframe.

# Usage

```
rowMaxs(table)
```

rowMins

Return a vector with the row-wise minima of a matrix or dataframe.

# Description

Return a vector with the row-wise minima of a matrix or dataframe.

```
rowMins(table)
```

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subsetBins	Create a SQM object containing only the requested bins, and the contigs and ORFs contained in them.

## **Description**

Create a SQM object containing only the requested bins, and the contigs and ORFs contained in them.

#### **Usage**

```
subsetBins(SQM, bins, trusted_functions_only = F,
  ignore_unclassified_functions = F, rescale_copy_number = T)
```

## Arguments

SQM object to be subsetted.

bins character. Vector of bins to be selected.

trusted\_functions\_only

logical. If TRUE, only highly trusted functional annotations (best hit + best average) will be considered when generating aggregated function tables. If FALSE, best hit annotations will be used (default FALSE).

ignore\_unclassified\_functions

logical. If FALSE, ORFs with no functional classification will be aggregated together into an "Unclassified" category. If TRUE, they will be ignored (default FALSE).

rescale\_copy\_number

logical. If TRUE, copy numbers with be recalculated using the RecA/RadA coverages in the subset. Otherwise, RecA/RadA coverages will be taken from the original object. By default it is set to TRUE, which means that the returned copy numbers will represent the average copy number per function *in the genomes of the selected bins*.

#### Value

SQM object containing only the requested bins.

# See Also

```
subsetContigs, subsetORFs
```

```
data(Hadza)
# Which are the five most complete bin?
topBinNames = rownames(Hadza$bins$table)[order(Hadza$bins$table[,'Completeness'], decreasing=T)][1:5]
topBins = subsetBins(Hadza, topBinNames)
```

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subsetContigs

Select contigs

## **Description**

Create a SQM object containing only the requested contigs, the ORFs contained in them and the bins that contain them.

## Usage

```
subsetContigs(SQM, contigs, trusted_functions_only = F,
  ignore_unclassified_functions = F, rescale_copy_number = F)
```

## **Arguments**

SQM

SQM object to be subsetted.

contigs

character. Vector of contigs to be selected.

trusted\_functions\_only

logical. If TRUE, only highly trusted functional annotations (best hit + best average) will be considered when generating aggregated function tables. If FALSE, best hit annotations will be used (default FALSE).

ignore\_unclassified\_functions

logical. If FALSE, ORFs with no functional classification will be aggregated together into an "Unclassified" category. If TRUE, they will be ignored (default FALSE).

rescale\_copy\_number

logical. If TRUE, copy numbers with be recalculated using the RecA/RadA coverages in the subset. Otherwise, RecA/RadA coverages will be taken from the original object (default FALSE).

#### Value

SQM object containing only the selected contigs.

#### See Also

subsetORFs

```
data(Hadza)
# Which contigs have a GC content below 40?
lowGCcontigNames = rownames(Hadza$contigs$table[Hadza$contigs$table[,'GC perc']<40,])
lowGCcontigs = subsetContigs(Hadza, lowGCcontigNames)
hist(lowGCcontigs$contigs$table[,'GC perc'])</pre>
```

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Filter results by function

## **Description**

Create a SQM object containing only the ORFs with a given function, and the contigs and bins that contain them.

#### Usage

```
subsetFun(SQM, fun, fixed = F, trusted_functions_only = F,
  ignore_unclassified_functions = F, rescale_copy_number = F)
```

## **Arguments**

SQM object to be subsetted.

fun character, pattern to search for in the different functional classifications.

fixed logical. If TRUE, pattern is a string to be matched as is. If FALSE the pattern is

treated as a regular expression (default FALSE).

trusted\_functions\_only

logical. If TRUE, only highly trusted functional annotations (best hit + best average) will be considered when generating aggregated function tables. If FALSE,

best hit annotations will be used (default FALSE).

ignore\_unclassified\_functions

logical. If FALSE, ORFs with no functional classification will be aggregated together into an "Unclassified" category. If TRUE, they will be ignored (default

FALSE).

rescale\_copy\_number

logical. If TRUE, copy numbers with be recalculated using the RecA/RadA coverages in the subset. Otherwise, RecA/RadA coverages will be taken from the original object (default FALSE).

#### Value

SQM object containing only the requested function.

#### See Also

subsetTax, subsetORFs, combineSQM. The most abundant items of a particular table contained in a SQM object can be eselected with mostAbundant.

```
data(Hadza)
Hadza.iron = subsetFun(Hadza, 'iron')
Hadza.carb = subsetFun(Hadza, 'Carbohydrate metabolism')
```

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subsetORFs

Select ORFs

## **Description**

Create a SQM object containing only the requested ORFs, and the contigs and bins that contain them. Internally, all the other subset functions in this package end up calling subsetORFs to do the work for them.

## Usage

```
subsetORFs(SQM, orfs, tax_source = "orfs", trusted_functions_only = F,
  ignore_unclassified_functions = F, rescale_copy_number = F)
```

# **Arguments**

SQM object to be subsetted.

orfs character. Vector of ORFs to be selected.

tax\_source character. Features used for calculating aggregated abundances at the different

taxonomic ranks. Either "orfs" or "contigs" (default "orfs").

trusted\_functions\_only

logical. If TRUE, only highly trusted functional annotations (best hit + best average) will be considered when generating aggregated function tables. If FALSE,

best hit annotations will be used (default FALSE).

ignore\_unclassified\_functions

logical. If FALSE, ORFs with no functional classification will be aggregated together into an "Unclassified" category. If TRUE, they will be ignored (default

FALSE).

rescale\_copy\_number

logical. If TRUE, copy numbers with be recalculated using the RecA/RadA coverages in the subset. Otherwise, RecA/RadA coverages will be taken from the original object (default FALSE).

#### Value

SQM object containing the requested ORFs.

#### A note on contig/bins subsetting

While this function selects the contigs and bins that contain the desired orfs, it DOES NOT recalculate contig/bin abundance and statistics based on the selected ORFs only. This means that the abundances presented in tables such as SQM\$contig\$abund or SQM\$bins\$tpm will still refer to the complete contigs and bins, regardless of whether only a fraction of their ORFs are actually present in the returned SQM object. This is also true for the statistics presented in SQM\$contigs\$table and SQM\$bins\$table.

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## **Examples**

```
data(Hadza)
# Select the 100 most abundant ORFs in our dataset.
mostAbundantORFnames = names(sort(rowSums(Hadza$orfs$tpm), decreasing=T))[1:100]
mostAbundantORFs = subsetORFs(Hadza, mostAbundantORFnames)
```

subsetRand

Select random ORFs

# Description

Create a random subset of a SQM object.

# Usage

```
subsetRand(SQM, N)
```

## **Arguments**

SQM object to be subsetted.

N numeric. number of random ORFs to select.

## Value

SQM object containing a random subset of ORFs.

#### See Also

subsetORFs

subsetTax

Filter results by taxonomy

# Description

Create a SQM object containing only the contigs with a given consensus taxonomy, the ORFs contained in them and the bins that contain them.

```
subsetTax(SQM, rank, tax, trusted_functions_only = F,
  ignore_unclassified_functions = F, rescale_copy_number = T)
```

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## **Arguments**

SQM object to be subsetted.

rank character. The taxonomic rank from which to select the desired taxa (superkingdom,

phylum, class, order, family, genus, species)

tax character. The taxon to select.

trusted\_functions\_only

logical. If TRUE, only highly trusted functional annotations (best hit + best average) will be considered when generating aggregated function tables. If FALSE, best hit annotations will be used (default FALSE).

ignore\_unclassified\_functions

logical. If FALSE, ORFs with no functional classification will be aggregated together into an "Unclassified" category. If TRUE, they will be ignored (default FALSE).

rescale\_copy\_number

logical. If TRUE, copy numbers with be recalculated using the RecA/RadA coverages in the subset. Otherwise, RecA/RadA coverages will be taken from the original object. By default it is set to TRUE, which means that the returned copy numbers will represent the average copy number per function *in the genomes of the selected taxa*.

#### Value

SQM object containing only the requested taxon.

## See Also

subsetFun, subsetContigs, combineSQM. The most abundant items of a particular table contained in a SQM object can be eselected with mostAbundant.

## **Examples**

```
data(Hadza)
Hadza.Escherichia = subsetTax(Hadza, 'genus', 'Escherichia')
Hadza.Bacteroidetes = subsetTax(Hadza, 'phylum', 'Bacteroidetes')
```

summary.SQM

summary method for class SQM

## **Description**

Computes different statistics of the data contained in the SQM object.

```
## S3 method for class 'SQM'
summary(SQM)
```

USiCGs 19

#### Value

A list of summary statistics.

USiCGs

Universal Single-Copy Genes

## **Description**

Lists of Universal Single Copy Genes for Bacteria and Archaea. These are useful for transforming coverages or tpms into copy numbers. This is an alternative way of normalizing data in order to be able to compare functional profiles in samples with different sequencing depths.

#### Usage

data(USiCGs)

#### **Format**

Character vector with the KEGG identifiers for 15 Universal Single Copy Genes.

#### Source

Carr et al., 2013. Table S1.

## References

Carr, Shen-Orr & Borenstein (2013). Reconstructing the Genomic Content of Microbiome Taxa through Shotgun Metagenomic Deconvolution *PLoS Comput. Biol.* **9**:e1003292. (PubMed).

# Examples

data(Hadza)

```
data(USiCGs)
### Let's look at the Universal Single Copy Gene distribution in our samples.
KEGG.tpm = Hadza$functions$KEGG$tpm
all(USiCGs %in% rownames(KEGG.tpm)) # Are all the USiCGs present in our dataset?
# Plot a boxplot of USiCGs tpms and calculate median USiCGs tpm.
# This looks weird in the test dataset bc it's only a subset of the metagenomes.
# In a set of complete metagenomes USiCGs should have fairly similar TPM averages and low dispersion across sample boxplot(t(KEGG.tpm[USiCGs,]), names=USiCGs, ylab='TPM', col='slateblue2')

### Now let's calculate the average copy numbers of each function.
# We do it for KEGG annotations here, but we could also do it for COGs or PFAMs.
KEGG.coverage = SQMtools::aggregate.fun(Hadza, 'KEGG', trusted_functions_only=T, ignore_unclassified_functions=USiCGs.cov = apply(KEGG.coverage[USiCGs,], 2, median)
KEGG.copynumber = t(t(KEGG.coverage) / USiCGs.cov) # Sample-wise division by the median USiCG coverage.
```

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