# Package 'RNASeqAnalysis'

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Title Analysis of RNA-Seq data: QC filtering, expression analysis, differential expression pipeline
Version 1.0.0
Author Nicky Thrupp [aut, cre]
Maintainer Nicky Thrupp <nixthrupp@gmail.com></nixthrupp@gmail.com>
<b>Description</b> The package implements a pipeline for the analysis of RNA-Seq data. The pipeline is designed to simplify the analysis as much as possible, thus requiring minimal user input.
Depends R (>= 3.2.1), ShortRead, lattice
Imports grDevices, graphics, stats
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R topics documented:
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2 availableReferences

addKallistoResToTable Adds the output kallisto directory to the original datafile.

## Description

**GOES HERE** 

## Usage

addKallistoResToTable(dataFile, countDat)

## **Arguments**

dataFile MORE countDat MORE

## Value

dataframe MORE

### See Also

collectKallistoCounts

availableReferences

A number of transcriptome indices have already been built, for use with kallistoQuant

## Description

A number of transcriptome index files (.idx files) have been pre-built and provided with the package. These files may be used as input to the kallsitoQuant pseudoalignment function (or the user may provide his/her own transcriptome file). The availableReferences function provides a list of these ready-available transcriptomes.

## Usage

availableReferences()

## **Arguments**

none No parameters.

## Value

list A list of available transcriptome indices (.idx files).

## **Examples**

availableReferences()

checkDataFile 3

checkDataFile	Checks that the format of the "data file" (which is used as input in many functions) is correct.

### **Description**

The function takes as input a data file (see datafileTemplate), and checks whether it is in a format acceptable to be used as input to other functions.

## Usage

```
checkDataFile(x)
```

## **Arguments**

х.

A data frame containing: the name of the file to be processed, whether it is single or paried-end data, the sample and replicate ID, and (optional) the name of an output file which results from the runQAandFilter function will be written to.

#### Value

A data frame which has been modified - the column names and classes have been verified (and corrected if necessary), all white space has been removed, observations relating to files which do not exist in the directory have been removed. The function also prints out any duplicated files - these should be removed by the user to prevent complications downstream.

## See Also

```
datafileTemplate
```

## Examples

```
dirPath <- system.file("extdata", "RNA-Seq-data", package = "RNASeqAnalysis")
setwd(dirPath)
dataF <- read.csv("data3.csv")
checkDataFile(dataF)</pre>
```

 $collect Kall is to {\tt Counts}\ \ \textit{Collects raw counts of gene (or \textit{transcript}) data\ \textit{generated by kallisto}.$ 

## **Description**

Extracts raw counts from each of the "abundance.tsv" files generated by kallisto (https://pachterlab.github.io/kallisto/about) expression analysis. The counts are put into a matrix (genes by sample), and rounded. This step collects the raw count data ready for further expression analysis, or for differential expression, using DESeq2.

## Usage

```
collectKallistoCounts(baseDir = "./kallistoResults")
```

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## **Arguments**

baseDir A character string of the directory where kallisto results are stored. For example,

baseDir = "kallistoResults" will search "kallistoResults/\*/abundance.tsv".

### Value

matrix. A matrix of (rounded) raw counts of genes (or transcripts) for each sample.

#### See Also

kallisto (https://pachterlab.github.io/kallisto/about)

compareQA	Compares quality before and after filtering fastq files. Porvides a
	graphic of the change in quality of fastq files, before and after run-
	ning the filterBadSeqs function.

## **Description**

Compares quality before and after filtering fastq files. Porvides a graphic of the change in quality of fastq files, before and after running the filterBadSeqs function.

## Usage

```
compareQA(dataFile, prefiltData, postfiltData)
```

#### **Arguments**

dataFile	An R data frame with the data to be processed. The R object is a standard
	format and must contain the following headings: File PF Sample Penlicate

format, and must contain the following headings: File, PE, Sample, Replicate, FilteredFile. More information about the file is available at datafileTemplate.

The checkDataFile function should be run on the file before use.

prefiltData A FastqQA object resulting from runQA. The object contains QA data on fastq

files before the filtering procedure (filterBadSeqs).

 $postfilt Data \qquad A \ Fastq QA \ object \ resulting \ from \ run QA. \ The \ object \ contains \ QA \ data \ on \ fastq$ 

files after the filtering procedure (filterBadSeqs).

## Value

a datafrome containing densities and quality scores for each fastq file (both before and after filtering). It also outputs a graph which shows comparisons of quality before and after filtering.

datafileTemplate 5

datafileTemplate

(Empty) template data file which summarises the data to be analysed

### **Description**

This is an empty dataset, the purpose of which is to act as a template for users. In the data file, the name of the file, the name of the sample and the technical replicate, and whether the sample is paired-end or single-end should be included. In addition, if the user is planning to filter the input file (see runQAandFilter), a column is provided for the user to add the name of the output fastq file

This data file acts as input to many of the package's functions. The format of the data file should not be altered, save to add observations (rows).

The data set datafileExample is an example of a data file containing observations.

## Usage

datafileTemplate

#### **Format**

a dataframe with 0 observations of 5 vairables

#### **Details**

- File. The name of the fastq (fastq.gz) file to be processed
- PE. "PE" should be entered into this column if the sample is from a paired-end run, or "SE" if the sample is from a single-end run.
- Sample. The name of the sample (e.g. "control", "lung").
- Replicate. The name of the replicate (e.g. "a", "2").
- FilteredFile. The name of the fastq (fastq.gz) file that the results from runQAandFilter function are written to.

## Author(s)

Nicky Thrupp <nixthrupp@gmail.com>

filterBadSeqs

Performs quality checks, then filters reads for quality

## **Description**

The function trims poor-quality bases and unknown bases from the ends of the sequences. Any reads which are too short, or contain any unknown bases (N), are removed from the file.

## Usage

```
filterBadSeqs(dataFile, minlength = 30, Phred = 25, blockSize = 1e+08,
  readerBlockSize = 1e+05, mc.cores = 1)
```

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#### **Arguments**

dataFile	An R data frame with the data to be processed. The R object is a standard format, and must contain the following headings: File, PE, Sample, Replicate, FilteredFile. More information about the file is available at datafileTemplate.
Phred	An integer which specifies Phred (ascii) quality score. Any two consecutive nucleotides with a quality score lower than this threshold will be discarded. Default score is 30.
blockSize	An integer which specifies the number of reads to be read at a time when processing. Default is 1e8.
mc.cores	The number of cores to use when paralleling. Default is 1 (i.e. no parallelisation)
minLength	An integer which specifies the minimum length for a read. Reads shorter than this length will be discarded. Default is 30 nucleotides.
readBlockSize	An integer which specifies the number of bytes (characters) to be read at one time. Smaller readBlockSize reduces memory requirements, but is less efficient. Default is 1e5.

### **Details**

The function should be run in the working directory, where all fastq files are found.

runQAandFilter iterates over each file specified in the "datafile", and filters and trims the reads for quality. This is done by iterating over chunks of reads in the fastq files at a time. The size of the chunks are decided by the "blockSize" and "readerBlockSize" parameters. More information about how this is done is available in the ShortRead package.

- \* it removes any trailing or leadining N's from each sequence,
- \* it removes any reads wich still contain N's,
- \* it trims the trailing end when it finds a minimum of 2 poor-quality bases in a window of 5. The threshold for poor quality is determined by the parameter "Phred", where the Phred score is logarithmically related to the probability of errors at each base,
- \* it removes any reads shorter than a minimum length (this is specified by the "minLength" parameter).

The function produces a new set of fastq files which have been filtered. The user must specify in the "FILTEREDFILE" column of the data file the output file. The user may specify the same output file for multiple input files - this will append new output to existing files, thereby allowing de-multiplexing of samples which have been run on different lanes. A new R object (QualityFilterResults) is created, which contains pointers to the input and output fastq files, as well as a summary of how many reads have been trimmed or removed.

## Value

An object of class QualityFilterResults.

#### See Also

https://en.wikipedia.org/wiki/Phred\_quality\_score for more about quality scores.

ShortRead for more information about quality reports, blockSize (n) and readerBlockSize.

kallistoIndex 7

kallistoIndex	Builds a transcriptome index (.idx) from a FASTA-formatted transcriptome.
	10.110

## **Description**

kallistoIndex builds an index of the transcriptome. Once built, the transcriptome index (in the form of a De Bruijn graph) may be used for pseudoalignment of the target RNA-Seq data in order to quantify expression (see kallistoQuant). A transcriptome De Bruijn graph (T-DBG) is constructed from k-mers in the transcriptome, where each k-mer in the transcriptome is a node in the T-DBG. Each transcript is represented by a different colour, and whenever a node appears in a transcript, it is given the colour of that transcript. MORE. Kallisto must be installed locally. Kallisto can be downloaded from https://pachterlab.github.io/kallisto/download.html.

### Usage

```
kallistoIndex(refTranscriptome, indexName = "./transcripts.idx")
```

#### Arguments

refTranscriptome

A character string of the transcriptome file (in fasta format) to be indexed.

indexName

A character string of the output index file to be created. Default is "./transcripts.idx").

## Value

An index file (.idx) which can be used to quantify expression data using kallistoQuant pseudoalignment.

## See Also

http://pachterlab.github.io/kallisto/ for more about Kallisto.

## **Examples**

```
kallistoIndex("./transcripts.fasta")
kallistoIndex("./transcripts.fa.gz", "./index.idx")
```

kallistoQuant

Kallisto quantifies transcript abundance from RNA-Seq data.

## **Description**

The data is broken down into k-mers and each k-mer is pseudoaligned to k-mers in the index. Because kallisto doesn't rely on full alignment, it is much quicker than other methods, without losing accuracy.

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#### **Usage**

```
kallistoQuant(dataFile, preFilt = FALSE, refIndex,
 refIndexFromPackage = FALSE, bootstrap = 0, fragmentLength = 200,
 fragmentSD = 10, biasCor = FALSE)
```

#### **Arguments**

refIndex

A character string of the name of the index file (usually, .idx) against which the fastq files are pseudoaligned. This file can be generated from a fasta-formatted transcriptome file using kallistoIndex.

refIndexFromPackage

A logical, false by default. If the user would like to use an index provided by the package (these can be viewed using availableReferences(), he/she should specifiy "true" here. Otherwise, the function will assume the index is provided

by the user.

An integer specifying the number of times to bootstrap. The output will provide bootstrap

a measure of variance.

fragmentLength An integer. Estimated fragment length. Only required for single-end data (for

paired-end data, Kallisto is able to calculate the fragment length). Default is 200

A numeric. The standard deviation of the fragment length. Only required for fragmentSD

single-end data (for paired-end data, Kallisto is able to calculate the standard

deviation). Default is 10 bp.

file1 A character string of the name of the RNA-Seq data file (fastq.gz) to be pro-

file2 A character string of the RNA-Seq data file (fastq.gz) to be processed - in the

case there is paired-end data.

is a logical. If true (default), a paired end protocol is chosen (for this, the file1 pairedEnd

and file2 parameters must be specified). If false, a single end protocol will be

run, and only file1 will be processed.

## Value

A data frame of the estimated abundances of each transcript specified in the input file(s). The data frame is also saved to a folder, which is given the title of the files, exlcuding the extensions. The folder contains these abundances (in text format and compressed format), as well as information about the run.

runQA

Performs quality checks

## **Description**

The function checks each fastq file specified in the "data file" for quality, and writes findings to a

## Usage

```
runQA(dataFile, preFilter = TRUE, mc.cores = NULL)
```

## **Arguments**

An R data frame with the data to be processed. The R object is a standard format, and must contain the following headings: File, PE, Sample, Replicate, FilteredFile. More information about the file is available at datafileTemplate.

PreFilter

A logical - if true (default), the function will select and analyse files which have not yet been processed for quality. If false, the function will select and analyse those files which have been processed for quality, i.e. the "filtered file" in the data file.

Mc.cores

A number specifying the number of cores to be used during parallel processing. Default is null - this will result in the number of cores in the default back-end being used. See BioCParallel::registered.

#### **Details**

The function should be run in the working directory, where all fastq files are found.

runQA iterates over each file specified in the "datafile". It runs a quality assessment from the ShortRead package. The ShortRead package (https://bioconductor.org/packages/release/bioc/html/ShortRead.html) contains more information about this step. The quality assessment may be performed before and after the filtering step, by setting the "pre-filter" parameter to true or to false, respectively. All quality assessment data is output to the "QA" directory. Quality reports are output to the working directory (under the QA directory). R objects of the raw data used to generate the reports are also saved to this directory.

#### Value

FastqQA object. Outputs quality results in the form of raw data (an R FastqQA object) and HTML format (saved to "QA" directory).

summariseFilteringStats

Collects summary statistics on filtering

## Description

**DESCRIPTION GOES HERE** 

#### Usage

summariseFilteringStats(x)

## **Arguments**

x An object of class QualityFilterResults (e.g. returned from runQAandFilter).

#### Value

A dataframe. The dataframe contains the number of reads which have been trimmed (on leading or trailing tails) for poor quality or unknown bases, the number of reads which have been removed entirely due to poor quality, possession of unknown bases, or length too short. Also produces graphs of absolute number of reads output, and relative numbers of reads which have been trimmed or removed (in PDF format).

## See Also

 ${\tt runQAandFilter}$ 

## **Index**

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