

Package ‘BgeeCall’

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Type Package

Title Automatic RNA-Seq present/absent gene expression calls generation

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Description BgeeCall allows to generate present/absent gene expression calls without using an arbitrary cutoff like TPM<1.

Calls are generated based on reference intergenic sequences. These sequences are generated based on expression of all RNA-Seq libraries of each species integrated in Bgee (<https://bgee.org>).

Depends R (>= 3.6)

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URL <https://github.com/BgeeDB/BgeeCall>

BugReports <https://github.com/BgeeDB/BgeeCall/issues>

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AbundanceMetadata-class

AbundanceMetadata S4 class

Description

An S4 class that is the parent class of all abundance tool Classes. It contains information needed to all abundance tools. This class can be seen as an abstract class, you should never instantiate it.

Slots

- txOut** Similar to tximport txOut parameter. Allows to keep abundance at transcript level if TRUE (default = FALSE)
- ignoreTxVersion** logical used to remove transcript version in transcript ID if TRUE (default = FALSE)
- cutoff_type** Defines the approach used to generate present/absent calls. default value is 'pValue', allowing calls to be generated using a pValue. Other possible values are 'intergenic' allowing to use a ratio of intergenic sequences considered as present as a threshold, or use qValue allowing calls to be generated from a qValue.
- cutoff** numeric value of the cutoff used to generate the present/absent calls. If value of the slot cutoff_type is 'pValue' this cutoff will correspond to the highest pValue allowing to define a gene as present. If value of the slot cutoff_type is 'intergenic' this cutoff will correspond to the proportion of intergenic present divided by proportion of protein coding present. If value of the slot cutoff_type is 'qValue' this cutoff will correspond to the highest qValue allowing to define a gene as present. The qValue is calculated based on the proportion of intergenic/(intergenic + genic) at each unique abundance value (TPM). The default value is 0.05. Be careful when modifying this value as it could have a huge impact on present/absent calls.
- full_transcriptome_file** Name of the fasta file containing both transcriptomic and intergenic regions. This file is created by the pipeline. You should edit this slot only if you already have such a file with a different name.
- tx2gene_file** Name of the file containing the mapping between transcript IDs and gene IDs (See the tximport package vignette for more details). This file is created by the pipeline. You should edit this slot only if you already have such a file with a different name. This file must be store at get_species_path()

`tx2gene_file_without_version` Name of the file containing the mapping between transcript IDs and gene IDs if `ignoreTxVersion == TRUE` (See the `tximport` package vignette for more details). This file is created by the pipeline. You should edit this slot only if you already have such a file with a different name. This file must be store at `get_species_path()`

`gene2biotype_file` Name of the file containing the mapping between gene IDs and biotypes. This file is created by the pipeline. You should edit this slot only if you already have such a file with a different name.

`tool_name` Name of the tool that will be use to generate transcript abundance estimation. All descendant of this class have to define a value for this slot (in the prototype section)

`abundance_file` Name of the transcript-level abundance file. All descendant of this class have to define a value for this slot (in the prototype section)

`read_size_kmer_threshold` read size of the library below which transcript index is created using a smaller kmer size

`transcript_id_header` Name of the header of the column that contains transcript ID

`count_header` Name of the header of the column that contains count

`abundance_header` Name of the header of the column that contains abundance

`eff_length_header` Name of the header of the column that contains effective length

`transcript_calls_file_name` default name of file containing all transcript ids and calls (if calls created at transcript level)

`gene_calls_file_name` default name of file containing all gene ids and calls (if calls created at gene level)

`transcript_cutoff_file_name` default name of file containing summary of cutoff used to generate transcript expression calls (if calls created at transcript level)

`gene_cutoff_file_name` default name of file containing summary of cutoff used to generate gene expression calls (if calls created at gene level)

`transcript_distribution_file_name` default name of density plot file containing TPM distribution of all transcripts (if calls created at transcript level)

`gene_distribution_file_name` default name of density plot file containing TPM distribution of all genes (if calls created at gene level)

BgeeCall

generate gene expression calls with BgeeCall

Description

BgeeCall allows to generate present/absent gene expression calls without using an arbitrary cut-off like $TPM < 1$. Calls are generated based on reference intergenic sequences. These sequences are generated based on expression of all RNA-Seq libraries of each species integrated in Bgee (<https://bgee.org>).

Details

Thes most important functions are :

- `generate_calls_workflow` : generate present/absent calls on a computer
- `generate_slurm_indexes` : generate kallisto indexes for a list of libraries on a cluster with slurm queuing system.
- `generate_slurm_calls` : generate present/absent calls for a list of libraries on a cluster with slurm queuing system. Indexes have to be generated first with the function '`generate_slurm_indexes`'
- `merging_libraries` : merge calls from different libraries corresponding to the same condition. Extremely useful if different libraries correspond to same condition (e.g. same anatomical entity from same species)

For more details please have a look at the vignette with the command **`vignette("BgeeCall")`**

Author(s)

Julien Wollbrett

See Also

<https://github.com/BgeeDB/BgeeCall>

BgeeMetadata-class	<i>BgeeMetadata S4 class</i>
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Description

An S4 class that contains all information to retrieve intergenic regions generated by Bgee.

Slots

`intergenic_release` Bgee intergenic release that will be used

`all_releases` list of all reference intergenic releases that can be used to generate your present/absent expression calls.

`intergenic_prefix` String used to generate an intergenic release specific output directory

create_kallisto_index *Create kallisto indexes.*

Description

This function creates kallisto indexes. Two indexes can be created depending on the reads size (see 'myKallistoMetadata@read_size_kmer_threshold' and 'UserMetadata@reads_size' for more information). One with default kmer value (31 nt) and one with kmer size of 15 nt. In order to generate.

Usage

```
create_kallisto_index(  
  myKallistoMetadata,  
  myBgeeMetadata,  
  myUserMetadata,  
  transcriptome_path = ""  
)
```

Arguments

myKallistoMetadata A Reference Class KallistoMetadata object.

myBgeeMetadata A Reference Class BgeeMetadata object.

myUserMetadata A Reference Class UserMetadata object.

transcriptome_path path to the transcriptome fasta file. If no path is provided the default path created using BgeeCall will be used. **IMPORTANT** : in BgeeCall the transcriptome used to generate present/absent calls contains both intergenic sequences downloaded from Bgee and the reference transcriptome. If this function is run to generate present/absent then 'transcriptome_path' has to be empty

Value

create kallisto index and save it on the hard drive

Author(s)

Julien Wollbrett.

Examples

```
## Not run:  
# first a transcriptome is needed. Here it is downloaded from AnnotationHub  
library(AnnotationHub)  
ah <- AnnotationHub()  
ah_resources <- query(ah, c('Ensembl', 'Caenorhabditis elegans', '84'))
```

```

# kallisto can not deal with S4 objects. A Path to a transcriptome file is
# required
transcriptome_object <- rtracklayer::import.2bit(ah_resources[['AH50453']])
transcriptome_path <- file.path(getwd(), 'transcriptome.fa')
Biostrings::writeXStringSet(transcriptome_object, transcriptome_path)

# initialize objects needed to create destination folder
bgee <- new('BgeeMetadata')
user <- new('UserMetadata', species_id = '6239')
kallisto <- new('KallistoMetadata')

# generate transcriptome index
create_kallisto_index(kallisto, bgee, user, transcriptome_path)

## End(Not run)

```

download_fasta_intergenic

Download fasta intergenic

Description

Check if reference intergenic fasta file has already been downloaded. If not the file is downloaded from Bgee FTP or from the community repository depending on myBgeeMetadata@intergenic_release. if myBgeeMetadata@intergenic_release == "community" then reference intergenic will be downloaded from the Zenodo community repository. Otherwise Reference intergenic sequences will be downloaded from the official Bgee FTP. Be careful when using reference intergenic sequences generated by the community as the Bgee team do not deeply review them.

Usage

```

download_fasta_intergenic(
  myBgeeMetadata = new("BgeeMetadata"),
  myUserMetadata,
  intergenic_file
)

```

Arguments

myBgeeMetadata A Reference Class BgeeMetadata object (optional)
 myUserMetadata A Reference Class UserMetadata object.
 intergenic_file
 path where intergenic file will be saved

Value

download fasta intergenic from Bgee FTP or from the Zenodo community and save it locally

Examples

```
{
  bgee_intergenic_file <- file.path(getwd(), 'intergenic.fasta')
  userMetadata <- new('UserMetadata', species_id = '7227')
}
```

download_kallisto	<i>Download binary version of kallisto.</i>
-------------------	---

Description

Check your OS and download correct binary version of kallisto.

Usage

```
download_kallisto(myKallistoMetadata, myUserMetadata)
```

Arguments

myKallistoMetadata
A Reference Class KallistoMetadata object.

myUserMetadata A Reference Class UserMetadata object.

Value

save uncompressed executable of kallisto on the hard drive

Author(s)

Julien Wollbrett.

Examples

```
{
  kallisto <- new('KallistoMetadata')
  user <- new('UserMetadata')
  download_kallisto(kallisto, user)
}
```

galaxy_calls	<i>generate calls for galaxy</i>
--------------	----------------------------------

Description

Function generating calls for galaxy. It is a simple adaptor of the function “generate_calls_workflow” adapted to galaxy requirements.

Usage

```
galaxy_calls(  
  species_id = NULL,  
  run_ids = NULL,  
  rnaseq_lib_path = NULL,  
  reads_size = 51,  
  transcriptome_path = NULL,  
  annotation_path = NULL,  
  output_file = NULL,  
  custom_intergenic_path = NULL  
)
```

Arguments

species_id	The NCBI taxon ID of the species
run_ids	The subset of runs from the library used to generate the calls. By default all runs are used.
rnaseq_lib_path	Path to the directory containing the fastq files of the library
reads_size	mean size of the reads. (Default = 51)
transcriptome_path	Path to the transcriptome fastq file
annotation_path	path to the gtf file containing gene annotation
output_file	file containing the path to all result files generated
custom_intergenic_path	(optional) To use if the "custom" reference intergenic release has been selected. Provides the path to the reference intergenic file

```
generate_calls_workflow
```

```
generate present/absent calls
```

Description

Main function running the workflow that generates present/absent calls from a file, a data.frame, or objects of the classe UserMetadata (please choose only 1 out of the 3). This workflow is highly tunable by editing default values of the slots of S4 objects. For more information on how to tune the workflow please have a look at the vignette and the documentation of the classes KallistoMetadata, AbundanceMetadata, UserMetadata and BgeeMetadata

Usage

```
generate_calls_workflow(
  abundanceMetadata = new("KallistoMetadata"),
  bgeeMetadata = new("BgeeMetadata"),
  userMetadata = NULL,
  userDataFrame = NULL,
  userFile = NULL,
  checkTxVersion = FALSE
)
```

Arguments

- | | |
|-------------------|--|
| abundanceMetadata | A Class AbundanceMetadata object (optional) allowing to tune your gene quantification abundance analyze |
| bgeeMetadata | A Class BgeeMetadata object (optional) allowing to choose the version of reference intergenic sequences |
| userMetadata | A Class UserMetadata object (optional). generate present/absent calls using slots of the UserMetadata class. |
| userDataFrame | <p>a data.frame containing all information to generate present/absent calls. Each line of this data.frame will generate calls for one RNA-Seq library. This data.frame must contains between 4 and 8 columns :</p> <ul style="list-style-type: none"> • species_id : The ensembl species ID • run_ids : (optional) allows to generate calls for a subpart of all runs of the library. must be a character or a list of characters • reads_size (optional) the size of the reads of the library (Default = 51) if the reads size is lower than 51 abundance quantification will be run from an index generated with a smaller kmer size • rnaseq_lib_path : path to RNA-Seq library directory • transcriptome_path : path to transcriptome file • annotation_path : path to annotation file • output_dir : (optional)root of the directory where results will be written |

- custom_intergenic_path : (optional) To use if the "custom" reference intergenic release has been selected. Provide the path to the reference intergenic file
- userFile path to a tsv file containing between 4 and 8 columns. these columns are the same than for userDataFrame (see above). a template of this file is available at the root of the package and accessible with the command `system.file('userMetadataTemplate.tsv', package = 'BgeeCall')`
- checkTxVersion boolean used to define if BgeeCall check rather transcript version should be removed. Default value is FALSE

Value

paths to the 5 results files (see vignette for more details)

Author(s)

Julien Wollbrett

See Also

AbundanceMetadata, KallistoMetadata, BgeeMetadata, UserMetadata

Examples

```
## Not run:
# import gene annotation and transcriptome from AnnotationHub
library(AnnotationHub)
ah <- AnnotationHub()
ah_resources <- query(ah, c('Ensembl', 'Caenorhabditis elegans', '84'))
annotation_object <- ah_resources[['AH50789']]
transcriptome_object <- rtracklayer::import.2bit(ah_resources[['AH50453']])

# instanciate BgeeCall object
# add annotation and transcriptome in the user_BgeeCall object
# it is possible to import them using an S4 object (GRanges, DNASTringSet)
# or a file (gtf, fasta) with methods setAnnotationFromFile() and
# setTranscriptomeFromFile()
user_BgeeCall <- setAnnotationFromObject(user_BgeeCall,
                                         annotation_object,
                                         'WBcel235_84')
user_BgeeCall <- setTranscriptomeFromObject(user_BgeeCall,
                                             transcriptome_object,
                                             'WBcel235')

# provide path to the directory of your RNA-Seq library
user_BgeeCall <- setRNASeqLibPath(user_BgeeCall,
                                  system.file('extdata', 'SRX099901_subset',
                                               package = 'BgeeCall'))

# run the full BgeeCall workflow
calls_output <- generate_calls_workflow(
  userMetadata = user_BgeeCall)
```

```
## End(Not run)
```

```
generate_initial_intergenic_regions
```

generates the initial intergenic regions

Description

generates intergenic regions from the species GTF and FASTA file by taking intergenic regions upstream and downstream of each gene

Usage

```
generate_initial_intergenic_regions(  
  gene_gtf_path = "./genomes/Homo_sapiens.GRCh38.gtf.gz",  
  genome_fasta_path = "./genomes/Homo_sapiens.GRCh38.genome.fa",  
  N_block_size = 31,  
  N_proportion = 0.05,  
  output_gtf_path = "./intergenic_regions/",  
  Rout_path = "./"  
)
```

Arguments

gene_gtf_path	full path to input gene gtf file
genome_fasta_path	full path to input genome fasta file
N_block_size	number of successive N from which it is considered as a block of N (and removed)
N_proportion	higher proportion of N than this threshold results in removing the sequence
output_gtf_path	full path to output folder + base name for output files
Rout_path	path to output .Rout file (optional)

Author(s)

Alessandro Brandulas Cammarata
Julien Wollbrett
Julien Roux
Marta Rosikiewicz

generate_presence_absence
Generate presence absence

Description

Generate presence absence calls. It corresponds to the last part of the generation of the expression calls workflow. It runs the last part of the workflow generating present/absent expression calls. This function should only be used by advanced user who already manually run all previous parts of the pipeline. If you are not an advanced user it is safer to run the function “generate_calls_workflow” that run all steps of the workflow

Usage

```
generate_presence_absence(  
  myAbundanceMetadata = new("KallistoMetadata"),  
  myBgeeMetadata = new("BgeeMetadata"),  
  myUserMetadata,  
  pvalueCorrection = "None"  
)
```

Arguments

myAbundanceMetadata A descendant object of the Class myAbundanceMetadata (optional).
myBgeeMetadata A Class BgeeMetadata object (optional).
myUserMetadata A Class UserMetadata object.
pvalueCorrection A string indicating the method to use to correct the pValue.

Value

path to the 4 output files

Author(s)

Julien Wollbrett
Julien Roux
Sara Fonseca Costa
Alessandro Brandulas Cammarata

See Also

generate_calls_workflow

Examples

```
{
# this example reuse data present in the directory 'extdata' of the package.
user <- new('UserMetadata', working_path = system.file('extdata',
package = 'BgeeCall'), species_id = '6239', rnaseq_lib_path = system.file(
'extdata', 'SRX099901_subset', package = 'BgeeCall'),
annotation_name = 'WBcel235_84', simple_arborescence = TRUE)
bgee <- new('BgeeMetadata', intergenic_release = '1.0')
calls_output <- generate_presence_absence(myUserMetadata = user, myBgeeMetadata = bgee)

#
}
```

```
generate_reference_intergenic_regions
```

Generates reference intergenic regions by removing outlier intergenics

Description

We sum the TPM values of intergenic regions in specified directory containing multiple libraries and find a threshold to remove outlier intergenic regions based on protein coding genes we then filter the fasta file containing all intergenic regions to the reference intergenic regions by removing outliers

Usage

```
generate_reference_intergenic_regions(
  intergenic_regions_path,
  tx2gene_path,
  gene2biotype_path,
  TPM_path,
  output_path = "./reference_intergenic_regions",
  species = "my_species"
)
```

Arguments

intergenic_regions_path	path to the intergenic regions
tx2gene_path	path to the tx2gene file of the species
gene2biotype_path	path to the gene2biotype file of the species
TPM_path	path to the abundance files of the libraries
output_path	path to the output folder
species	species name or ID to be used in the output files

Author(s)

Alessandro Brandulas Cammarata

generate_slurm_calls *Generate present/absent calls on slurm queuing system*

Description

This function is meant to be used with a cluster where the Slurm queuing system is installed. It processes all steps to generate present/absent calls at RNA-Seq library level. This function does not generate the kallisto indexes. If they are not already generated please run function “generate_slurm_indexes” first. Steps of present/absent gene expression calls generation are :

- Quantifying abundances of transcripts from RNA-Seq libraries
- Summarizing abundance at gene level
- generate present/absent expression calls

Usage

```
generate_slurm_calls(
  kallistoMetadata = new("KallistoMetadata"),
  bgeeMetadata = new("BgeeMetadata"),
  userMetadata = new("UserMetadata"),
  userFile,
  submit_sh_template = NULL,
  slurm_options = NULL,
  rscript_path = NULL,
  modules = NULL,
  submit = TRUE,
  nodes = 10,
  checkTxVersion = FALSE
)
```

Arguments

kallistoMetadata	A Reference Class KallistoMetadata object (optional) allowing to tune your gene quantification abundance analyze. If no object is provided a new one will be created with default values.
bgeeMetadata	A Reference Class BgeeMetadata object (optional) allowing to choose the version of reference intergenic sequences. If no object is provided a new one will be created with default values.
userMetadata	A Class UserMetadata object (optional). If no object is provided a new one will be created with default values.

userFile	Path to the file where each line corresponds to one abundance quantification to be run. The structure of the file is the same than the 'userFile' used as input of the 'generate_calls_workflow' function. A template of this file can be loaded with the command : <code>""inputFile <- read.table(system.file("userMetadataTemplate.tsv", package = "BgeeCall"), header = TRUE)""</code> It is important to keep the same column names.
submit_sh_template	A template of the bash script used to submit the jobs. By default the submission script provided by rslurm is used. Modify only if module dependencies have to be added (like kallisto or R)
slurm_options	A named list of options recognized by sbatch. More details in the documentation of the rslurm::slurm_apply function
rscript_path	The location of the Rscript command. If not specified, defaults to the location of Rscript within the R installation being run.
modules	A list of modules you want to load in the environment. Should stay empty except if you need to load R and/or kallisto (e.g module add R)
submit	Whether or not to submit the job to the cluster with sbatch. Default value is TRUE
nodes	The (maximum) number of cluster nodes to spread the calculation over. slurm_apply automatically divides params in chunks of approximately equal size to send to each node. Less nodes are allocated if the parameter set is too small to use all CPUs on the requested nodes. By default this number is 10.
checkTxVersion	boolean used to define if BgeeCall check rather transcript version should be removed. Default value is FALSE

Value

generate calls

Examples

```
## Not run:
# use function with all default values
userFile <- "/path/to/userList.tsv"
sjobs <- generate_slurm_calls(userFile = userFile)

## End(Not run)
```

generate_slurm_indexes

Generate all indexes for the abundance quantification step

Description

Check all unique lines of the input file to check which indexes have to be generated before running all abundance quantification. This function is meant to be used with a cluster where the Slurm queuing system is installed. This step has to be run before the quantification otherwise indexes will be created for each abundance quantification. This will slow down the abundance quantification and can generate errors when writing the same file at the same time from different nodes. This function also generate tx2gene and gene2biotype mapping files.

Usage

```
generate_slurm_indexes(
  kallistoMetadata = new("KallistoMetadata"),
  bgeeMetadata = new("BgeeMetadata"),
  userMetadata = new("UserMetadata"),
  userFile,
  submit_sh_template = NULL,
  slurm_options = NULL,
  rscript_path = NULL,
  modules = NULL,
  submit = TRUE,
  nodes = 10
)
```

Arguments

kallistoMetadata	A Reference Class KallistoMetadata object (optional) allowing to tune your gene quantification abundance analyze. If no object is provided a new one will be created with default values.
bgeeMetadata	A Reference Class BgeeMetadata object (optional) allowing to choose the version of reference intergenic sequences. If no object is provided a new one will be created with default values.
userMetadata	A Class UserMetadata object (optional). If no object is provided a new one will be created with default values.
userFile	Path to the file where each line corresponds to one abundance quantification to be run. The structure of the file is the same than the 'userFile' used as input of the 'generate_calls_workflow' function. A template of this file can be loaded with the command : <code>“inputFile <- read.table(system.file("userMetadataTemplate.tsv", package = "BgeeCall"), header = TRUE)“</code> It is important to keep the same column names.
submit_sh_template	A template of the bash script used to submit the jobs. By default the submission script provided by rslurm is used. Modify only if module dependencies have to be added (like kallisto or R)
slurm_options	A named list of options recognized by sbatch. More details in the documentation of the <code>rslurm::slurm_apply</code> function

rscript_path	The location of the Rscript command. If not specified, defaults to the location of Rscript within the R installation being run.
modules	A list of modules you want to load in the environment. Should stay empty except if you need to load R and/or kallisto (e.g module add R)
submit	Whether or not to submit the job to the cluster with sbatch. Default value is TRUE
nodes	The (maximum) number of cluster nodes to spread the calculation over. slurm_apply automatically divides params in chunks of approximately equal size to send to each node. Less nodes are allocated if the parameter set is too small to use all CPUs on the requested nodes. By default this number is 10.

Value

generate index files

Examples

```
## Not run:
# use function with all default values
userFile <- "/path/to/userList.tsv"
sjobs <- generate_slurm_indexes(userFile = userFile)

## End(Not run)
```

getIntergenicPrefix *'intergenic_prefix' Getter*

Description

Get value of the 'intergenic_prefix' slot

Usage

```
getIntergenicPrefix(bgeeObject)

## S4 method for signature 'BgeeMetadata'
getIntergenicPrefix(bgeeObject)
```

Arguments

bgeeObject The BgeeMetadata object

Value

the value of the 'intergenic_prefix' slot of the object

Examples

```
{
  bgee <- new("BgeeMetadata")
  intergenic_prefix <- getIntergenicPrefix(bgee)
}
```

`getIntergenicRelease` *'intergenic_release' Getter*

Description

Get value of the `'intergenic_release'` slot

Usage

```
getIntergenicRelease(bgeeObject)

## S4 method for signature 'BgeeMetadata'
getIntergenicRelease(bgeeObject)
```

Arguments

`bgeeObject` The `BgeeMetadata` object

Value

the value of the `'intergenic_release'` slot of the object

Examples

```
{
  bgee <- new("BgeeMetadata")
  intergenic_release <- getIntergenicRelease(bgee)
}
```

getRunIds	<i>'run_ids' Getter</i>
-----------	-------------------------

Description

Get value of the 'run_ids' slot

Usage

```
getRunIds(userObject)

## S4 method for signature 'UserMetadata'
getRunIds(userObject)
```

Arguments

userObject The UserMetadata object

Value

the value of the 'run_ids' slot of the object

Examples

```
{
  user <- new("UserMetadata")
  run_ids <- getRunIds(user)
}
```

getSimpleArborescence	<i>'simple_arborescence' Getter</i>
-----------------------	-------------------------------------

Description

Get value of the 'simple_arborescence' slot

Usage

```
getSimpleArborescence(userObject)

## S4 method for signature 'UserMetadata'
getSimpleArborescence(userObject)
```

Arguments

userObject The UserMetadata object

Value

the value of the 'simple_arborescence' slot of the object

Examples

```
{
  user <- new("UserMetadata")
  simple_arborescence <- getSimpleArborescence(user)
}
```

getWorkingPath	'working_path' Getter
----------------	-----------------------

Description

Get value of the 'working_path' slot

Usage

```
getWorkingPath(userObject)

## S4 method for signature 'UserMetadata'
getWorkingPath(userObject)
```

Arguments

userObject The UserMetadata object

Value

the value of the 'working_path' slot of the object

Examples

```
{
  user <- new("UserMetadata")
  working_path <- getWorkingPath(user)
}
```

get_summary_stats	<i>Gather statistical information</i>
-------------------	---------------------------------------

Description

Collect the statistics provided by the gene_cutoff_info_file from each individual library, in order to generate a global summary file.

Usage

```
get_summary_stats(userFile, outDir)
```

Arguments

userFile	A data frame containing all information of each library
outDir	Output directory where the generated file should be saved

Value

A tsv file

Author(s)

Sara Fonseca Costa

KallistoMetadata-class	<i>KallistoMetadata S4 class</i>
------------------------	----------------------------------

Description

An S4 class that is the descendant of the AbundanceMetadata class. It contains all metadata needed to run kallisto analysis. All slots of this class have a default value. You do not need to edit them to run the package

Slots

download_kallisto	A logical allowing to use an already installed version of kallisto or to download a version that will be used only by this package
kallisto_windows_url	URL to the binary of kallisto for windows
kallisto_linux_url	URL to the binary of kallisto for linux
kallisto_osx_url	URL to the binary of kallisto for MacOS
kallisto_windows_dir	Name of the directory where kallisto will be installed on windows
kallisto_linux_dir	Name of the directory where kallisto will be installed on linux

kallisto_osx_dir Name of the directory where kallisto will be installed on Mac
unix_kallisto_name Name of the kallisto executable in linux and macOS
windows_kallisto_name Name of the kallisto executable in windows
index_file Name of index file generated by kallisto with default kmer size. It will be generated using the fasta file that contains both transcriptomic and intergenic regions. Do not use an index you generated outside of this package. This file is created by the pipeline. You should edit this slot only if you already have such a file with a different name. This file must be store at `get_tool_path()`
k15_index_file same as `index_file`. This index is generated with smallest kmers and will be used only for libraries containing reads smallest than 50nt.
single_end_parameters kallisto parameters used to run a single end mapping
pair_end_parameters kallisto parameters used to run a pair end mapping
overwrite_index logical allowing to overwrite already existing index. FALSE by default. Then by default already existing index files will not be generated again.
overwrite_quant logical allowing to overwrite already existing abundance.txt files. FALSE by default. Then by default already existing quantification files will not be generated again.
overwrite_calls logical allowing to overwrite already existing present/absent calls. FALSE by default. Then by default already generated calls will not be generated again.

list_bgee_ref_intergenic_species

List species having Bgee reference intergenic sequences

Description

Return information related to species having Bgee reference intergenic sequences available for the selected Bgee intergenic release:

speciesId the NCBI species ID of the species

specieName scientific species name

numberOfLibraries number of libraries used to generate these reference intergenic sequences

genomeVersion version of the genome used to generate the reference intergenic sequences

If a BgeeMetadata object is provided this function retrieve the list of species using `BgeeMetadata@intergenic_release`. If only a 'release' is provided it will use it to retrieve the list of species. If none of them are provided the default Bgee reference intergenic release will be used.

Usage

```
list_bgee_ref_intergenic_species(myBgeeMetadata = NULL, release = NULL)
```

Arguments

myBgeeMetadata A Reference Class BgeeMetadata object

release A Bgee reference intergenic release name

Value

list all species having reference intergenic sequences available in the selected release

Author(s)

Julien Wollbrett

Examples

```
{  
  bgee <- new("BgeeMetadata")  
  list_bgee_ref_intergenic_species(myBgeeMetadata = bgee)  
  list_bgee_ref_intergenic_species(release = '0.2')  
}
```

```
list_community_ref_intergenic_species
```

*List species having reference intergenic sequences created by the
BgeeCall community*

Description

Return information related to species having reference intergenic sequences created by the BgeeCall community - speciesId : the NCBI species ID of the species - url : url to the reference intergenic fasta file - numberOfLibraries : number of libraries used to generate these reference intergenic sequences

Usage

```
list_community_ref_intergenic_species()
```

Value

list all species having reference intergenic sequences created by the community

Author(s)

Julien Wollbrett

Examples

```
{  
  list_community_ref_intergenic_species()  
}
```

`list_intergenic_release`*List reference intergenic releases usable with the BgeeCall package*

Description

Returns information on available Bgee intergenic releases, the access URL for FTP, and the date of release

Usage

```
list_intergenic_release(release = NULL)
```

Arguments

release	A character specifying a targeted release number (e.g., '0.1'). If not specified, all available releases are shown.
---------	---

Value

A data frame with information on Bgee intergenic releases available to use with the BgeeCall package.

Author(s)

Julien Wollbrett

Examples

```
{  
  list_intergenic_release()  
}
```

`merge_transcriptome_and_intergenic`*Merge transcriptome file provided by the user with the Bgee intergenic fasta file.*

Description

This function will create a file corresponding to the concatenation of the transcriptome fasta file provided by the user and the corresponding intergenic fasta file created by Bgee.

Usage

```
merge_transcriptome_and_intergenic(  
  myKallistoMetadata,  
  myBgeeMetadata,  
  myUserMetadata  
)
```

Arguments

myKallistoMetadata A Reference Class KallistoMetadata object.
myBgeeMetadata A Reference Class BgeeMetadata object.
myUserMetadata A Reference Class UserMetadata object.

Value

save merged file on the hard drive

Author(s)

Julien Wollbrett.

Examples

```
{  
  bgee <- new('BgeeMetadata', intergenic_release = '0.1')  
  user <- new ('UserMetadata', species_id = '6239')  
  kallisto <- new('KallistoMetadata')  
  user <- setTranscriptomeFromFile(user, system.file("extdata",  
    "transcriptome.fa", package = "BgeeCall"), 'WBcel235')  
  merge_transcriptome_and_intergenic(kallisto, bgee, user)  
}
```

merging_libraries

Calls of expression in combined libraries

Description

Merging/combine libraries based in a condition specified by the user. The merging can be done using the p-values of the libraries, by applying the BH method, or using the q-values of the libraries using the `fdr_inverse` method.

Usage

```
merging_libraries(  
  userFile = NULL,  
  approach = "BH",  
  condition = "species_id",  
  cutoff = 0.05,  
  outDir = NULL,  
  weights = FALSE  
)
```

Arguments

userFile	A file provided by the user with correspondent conditions
approach	Approach used to do the merging of libraries
condition	Condition/s where the merging should be done
cutoff	Cutoff that should be applied to call Present/Absent genes
outDir	Directory where the output files should be saved
weights	Boolean to indicate if the user wants to use weights in the mean/median p-value calculation

Value

A dataframe containing the minimum quantitative value (p-value or q-value) and the calls to each gene id for the referent condition.

Author(s)

Sara Fonseca Costa

Alessandro Brandulas Cammarata

Examples

```
## Not run:  
callsMerging_species <- merging_libraries(userFile = 'PATH_USER_FILE', approach = 'BH',  
condition = 'species_id', cutoff = 0.05, outDir = 'PATH_OUTPUT')  
callsMerging_species_sex <- merging_libraries(userFile = 'PATH_USER_FILE', approach = 'fdr_inverse',  
condition = c(species_id, sex), cutoff = 0.01, outDir = 'PATH_OUTPUT')  
callsMerging_all <- merging_libraries(userFile = 'PATH_USER_FILE', approach = 'fdr_inverse',  
condition = c(species_id, anatEntity, devStage, sex, strain), cutoff = 0.05, outDir = 'PATH_OUTPUT')  
  
## End(Not run)
```

Pvalue_averaging	<i>Pvalue averaging</i>
------------------	-------------------------

Description

Combined p-values for each gene Id using the mean or median averaging method

Usage

```
Pvalue_averaging(pval_collect, w_values = c(), method = "mean")
```

Arguments

pval_collect	A data frame containing the p-values for each gene id of each library
w_values	A vector of weights to be used in the mean p-value calculation
method	Method used to calculate the mean p-value

Value

A dataframe containing the consensus p-value for each gene id

Author(s)

Alessandro Brandulas Cammarata

Examples

```
## Not run:
Pvalue_averaging(pval_collect = pval_collect, w_values = c(0.5, 0.5), method = "mean")

## End(Not run)
```

run_kallisto	<i>Run one kallisto abundance analyse</i>
--------------	---

Description

Run kallisto and all preliminary steps if needed like : - creation of transcriptome with intergenic (if needed) - installation of kallisto (if needed) - index creation (if needed) - run kallisto quantification

Usage

```
run_kallisto(
  myKallistoMetadata,
  myBgeeMetadata,
  myUserMetadata,
  transcriptome_path = ""
)
```

Arguments

myKallistoMetadata A Reference Class KallistoMetadata object.

myBgeeMetadata A Reference Class BgeeMetadata object.

myUserMetadata A Reference Class UserMetadata object. This object has to be edited before running kallisto @seealso UserMetadata.R

transcriptome_path path to the transcriptome fasta file. If no path is provided the default path created using BgeeCall will be used. **IMPORTANT** : in BgeeCall the transcriptome used to generate present/absent calls contains both intergenic sequences downloaded from Bgee and the reference transcriptome.

Value

create kallisto output files and save them on the hard drive

Author(s)

Julien Wollbrett.

Examples

```
## Not run:
# first a transcriptome is needed. Here it is downloaded from AnnotationHub
library(AnnotationHub)
ah <- AnnotationHub()
ah_resources <- query(ah, c('Ensembl', 'Caenorhabditis elegans', '84'))

# kallisto can not deal with S4 objects. Path to transcriptome file is
# required
transcriptome_object <- rtracklayer::import.2bit(ah_resources[['AH50453']])
transcriptome_path <- file.path(getwd(), 'transcriptome.fa')
Biostrings::writeXStringSet(transcriptome_object, transcriptome_path)

# initialize objects needed to create destination folder
bgee <- new('BgeeMetadata')
user <- new('UserMetadata', species_id = '6239')
user <- setRNASeqLibPath(user, system.file(
  'extdata', 'SRX099901_subset',
  package = 'BgeeCall'))
kallisto <- new('KallistoMetadata')
```

```
# generate transcriptome index
run_kallisto(kallisto, bgee, user, transcriptome_path)

## End(Not run)
```

run_tximport	<i>Run tximport</i>
--------------	---------------------

Description

Run tximport. Will summarize abundance estimation from transcript level to gene level if ‘myAbundanceMetadata@txout == FALSE’. Otherwise keep abundance estimation at transcript level.

Usage

```
run_tximport(
  myAbundanceMetadata = new("KallistoMetadata"),
  myBgeeMetadata = new("BgeeMetadata"),
  myUserMetadata,
  abundanceFile = ""
)
```

Arguments

myAbundanceMetadata A descendant object of the Class myAbundanceMetadata.

myBgeeMetadata A Reference Class BgeeMetadata object.

myUserMetadata A Reference Class UserMetadata object.

abundanceFile (Optional) Path to the abundance file. NULL by default. If not NULL, the file located at ‘abundanceFile’ will be used to run tximport. Otherwise (Default) the path to the abundance file is deduced from attributes of classes ‘BgeeMetadata’, ‘UserMetadata’ and ‘AbundanceMetadata’

Value

a tximport object

Author(s)

Julien Wollbrett

Examples

```
{
  user <- new("UserMetadata", working_path = system.file("extdata",
    package = "BgeeCall"), species_id = "6239",
    rnaseq_lib_path = system.file("extdata",
      "SRX099901_subset", package = "BgeeCall"),
    annotation_name = "WBcel235_84", simple_arborescence = TRUE)
  abundance_file <- system.file('extdata', 'abundance.tsv', package = 'BgeeCall')
  bgee <- new("BgeeMetadata", intergenic_release = "1.0")
  tx_import <- run_tximport(myUserMetadata = user, myBgeeMetadata = bgee,
    abundanceFile = abundance_file)
}
```

setAnnotationFromFile *Set annotation_object of one UserMetadata object*

Description

Method of the class UserMetadata. Set annotation_object of one UserMetadata object by providing the path to a fasta transcriptome file.

Usage

```
setAnnotationFromFile(userObject, annotationPath, annotationName)

## S4 method for signature 'UserMetadata,character,missing'
setAnnotationFromFile(userObject, annotationPath, annotationName)

## S4 method for signature 'UserMetadata,character,character'
setAnnotationFromFile(userObject, annotationPath, annotationName)
```

Arguments

userObject The UserMetadata object
 annotationPath Absolute path to the annotation file
 annotationName (optional) Name of the annotation. Will be used to create folders.

Details

If no annotationName is provided the name of the annotation file will be used to create folders.

Value

An object of the class UserMetadata

Examples

```
{
# path to gtf annotation file
annotation_file <- system.file("extdata", "annotation.gtf", package = "BgeeCall")
user <- new("UserMetadata")
user <- setAnnotationFromFile(user, annotation_file,
                             "annotation_name")
}
```

```
setAnnotationFromObject
```

```
Set annotation_object of one UserMetadata object
```

Description

Method of the class UserMetadata. Set annotation_object of one UserMetadata object by using one GRanges object as input.

Usage

```
setAnnotationFromObject(userObject, annotationObject, annotationName)

## S4 method for signature 'UserMetadata,GRanges,character'
setAnnotationFromObject(userObject, annotationObject, annotationName = "")
```

Arguments

```
userObject      The UserMetadata object
annotationObject
                 object of thr GRanges S4 class
annotationName  (optional) Name of the annotation. Will be used to create folders.
```

Details

If no annotationName is provided the name of the file is used to create folders.

Value

An object of the class UserMetadata

Examples

```
{
  user <- new("UserMetadata")
  annotation_object <- rtracklayer::import(system.file("extdata",
    "annotation.gtf", package = "BgeeCall"))
  user <- setAnnotationFromObject(user, annotation_object,
                                "annotation_name")
}
```

setIntergenicRelease *'intergenic_release' Setter*

Description

Set value of the 'intergenic_release' slot

Usage

```
setIntergenicRelease(bgeeObject, intergenicRelease)

## S4 method for signature 'BgeeMetadata,character'
setIntergenicRelease(bgeeObject, intergenicRelease)
```

Arguments

bgeeObject The BgeeMetadata object
intergenicRelease
 character corresponding to the 'intergenic_release'

Value

An object of the class BgeeMetadata with new 'intergenic_release' value

Examples

```
{
  bgee <- new("BgeeMetadata")
  bgee <- setIntergenicRelease(bgee, "0.1")
}
```

setOutputDir	<i>'output_dir' Setter</i>
--------------	----------------------------

Description

Set value of the 'output_dir' slot

Usage

```
setOutputDir(userObject, outputDir)

## S4 method for signature 'UserMetadata,character'
setOutputDir(userObject, outputDir)
```

Arguments

userObject	The UserMetadata object
outputDir	path to the directory wanted as 'output_dir'

Value

An object of the class UserMetadata with new 'output_dir' value

Examples

```
{
  user <- new("UserMetadata")
  user <- setOutputDir(user, getwd())
}
```

setRNASeqLibPath	<i>'rnaseq_lib_path' Setter</i>
------------------	---------------------------------

Description

Set value of the 'rnaseq_lib_path' slot

Usage

```
setRNASeqLibPath(userObject, rnaSeqLibPath)

## S4 method for signature 'UserMetadata,character'
setRNASeqLibPath(userObject, rnaSeqLibPath)
```

Arguments

userObject The UserMetadata object
 rnaSeqLibPath path to the directory wanted as 'rnaseq_lib_path'

Value

An object of the class UserMetadata with new 'rnaseq_lib_path' value

Examples

```
{
  user <- new("UserMetadata")
  user <- setRNASeqLibPath(user, getwd())
}
```

setRunIds	<i>'run_ids' Setter</i>
-----------	-------------------------

Description

Method of the class UserMetadata. Set run_ids of one UserMetadata object by providing the id of all wanted runs

Usage

```
setRunIds(userObject, runIds)

## S4 method for signature 'UserMetadata,character'
setRunIds(userObject, runIds)
```

Arguments

userObject The UserMetadata object
 runIds id of all wanted runs

Value

An object of the class UserMetadata

Examples

```
{
  user <- new("UserMetadata")
  user <- setRunIds(user, c("RUN_1", "RUN_2"))
}
```

```
setSimpleArborescence 'simple_arborescence' Setter
```

Description

Set value of the 'simple_arborescence' slot

Usage

```
setSimpleArborescence(userObject, simpleArborescence)

## S4 method for signature 'UserMetadata,logical'
setSimpleArborescence(userObject, simpleArborescence)
```

Arguments

`userObject` The UserMetadata object

`simpleArborescence`
 boolean defining if output files will be created a simple arborescence (TRUE) or
 not (FALSE)

Value

An object of the class UserMetadata with new 'simple_arborescence' value

Examples

```
{
  user <- new("UserMetadata")
  user <- setSimpleArborescence(user, FALSE)
}
```

```
setTranscriptomeFromFile
  Set transcriptome_object of one UserMetadata object
```

Description

Method of the class UserMetadata. Set transcriptome_object of one UserMetadata object by providing the path to a fasta transcriptome file.

Usage

```

setTranscriptomeFromFile(userObject, transcriptomePath, transcriptomeName)

## S4 method for signature 'UserMetadata,character,missing'
setTranscriptomeFromFile(userObject, transcriptomePath, transcriptomeName)

## S4 method for signature 'UserMetadata,character,character'
setTranscriptomeFromFile(userObject, transcriptomePath, transcriptomeName)

```

Arguments

userObject The UserMetadata object

transcriptomePath
 Absolute path to the transcriptome file

transcriptomeName
 (optional) Name of the transcriptome. Will be used to create folders.

Details

If no transcriptomeName is provided the name of the transcriptome file will be used to create folders.

Value

An object of the class UserMetadata

Examples

```

{
  transcriptome_path <- system.file("extdata", "transcriptome.fa", package = "BgeeCall")
  user <- new("UserMetadata")
  user <- setTranscriptomeFromFile(user, transcriptome_path,
                                   "transcriptome_name")
}

```

```
setTranscriptomeFromObject
```

Set transcriptome_object of one UserMetadata object

Description

Method of the class UserMetadata. Set transcriptome_object of one UserMetadata object by using one DNASTringSet object as input.

Usage

```
setTranscriptomeFromObject(userObject, transcriptomeObject, transcriptomeName)

## S4 method for signature 'UserMetadata,DNAStringSet,character'
setTranscriptomeFromObject(userObject, transcriptomeObject, transcriptomeName)
```

Arguments

```
userObject      UserMetadata object
transcriptomeObject
                  Object of the DNAStringSet S4 class
transcriptomeName
                  Name of the transcriptome. Will be used to create transcriptome folders.
```

Details

Please use a DNAStringSet object as input. This class is defined in the Biostrings package

Value

an object of UserMetadata

Examples

```
{
  user <- new("UserMetadata")
  transcriptome_object <- Biostrings::readDNAStringSet(
    system.file("extdata", "transcriptome.fa", package = "BgeeCall"))
  user <- setTranscriptomeFromObject(user,
    transcriptome_object,
    "transcriptome_name")
}
```

setWorkingPath	<i>‘working_path’ Setter</i>
----------------	------------------------------

Description

Set value of the ‘working_path’ slot

Usage

```
setWorkingPath(userObject, workingPath)

## S4 method for signature 'UserMetadata,character'
setWorkingPath(userObject, workingPath)
```

Arguments

userObject The UserMetadata object
 workingPath path to the directory wanted as 'working_path'

Value

An object of the class UserMetadata with new 'working_path' value

Examples

```
{
  user <- new("UserMetadata")
  user <- setWorkingPath(user, getwd())
}
```

UserMetadata-class	<i>UserMetadata S4 class</i>
--------------------	------------------------------

Description

An S4 class containing all metadata that have to be provided by the user. It is mandatory to edit 'species_id', 'rnaseq_lib_path', 'transcriptome_path', 'annotation_name', 'annotation_object' and potentially 'run_ids' before using the package.

Slots

species_id The NCBI Taxon Id of the species
 run_ids A vector of character. Has to be provided only if a subset of runs present in UserMetadata@rnaseq_lib_path has to be run. If empty, all fastq files present in the rnaseq_lib_path will be considered as technical replicates and merged to run one transcript expression estimation analysis.
 reads_size The size of the reads. If smaller than 'KallistoMetadata@read_size_kmer_threshold', an index with a kmer size of 15 bp will be used.
 rnaseq_lib_path Path to the directory of the RNA-Seq library that contains fastq files. The extension of the fastq files name must be .fq, .fastq, .fq.gz, or .fastq.gz
 transcriptome_name Name of the transcriptome used to generate arborescence of output repositories.
 transcriptome_object Object containing transcriptome
 annotation_name Name of the annotation used to generate arborescence of output repositories.
 annotation_object Object containing annotations from GTF or GFF file
 working_path Working directory. By default the working directory is defined with the 'getwd()' function.
 gtf_source The source name from where the gtf file comes from. By default is ensembl.

`simple_arborescence` logical allowing to create a simple arborescence of directory. If 'TRUE' (default), all results will be on the same directory (`working_path/intergenic_release/all_results/libraryId`). Use 'FALSE' if you plan to generate expression calls for the same library using different transcriptomes or gene annotations, otherwise you will overwrite previous results. When 'FALSE' the path to result folder looks like : `working_path/intergenic_release/speciesId/kallisto/transcriptome_name/annotation_`

`output_dir` (optional) Allows to manually define your output directory. By default the path to output directory is created automatically from the `working_path` (`working_path/intergenic_release/all_results/libraryId/`).

`verbose` logical allowing to use the verbose mode. TRUE by default.

`custom_intergenic_path` path to a local version of reference intergenic fasta file. If NULL (by default) the reference intergenic fasta file will be downloaded. If not NULL BgeeCall will merge this local reference intergenic file with the transcriptome. Except if you generated your own intergenic regions always keep it NULL.

`encrypted_pattern` Allows to manage encrypted libraries. If a fastq file with the suffix `.enc` is found for a run, this slot will allow to use a string pattern to decrypt it. . This `encrypted_pattern` needs to contain the string `FASTQ_PATH` that will be transformed to the actual path to the fastq file.

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