

Package ‘umiAnalyzer’

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Maintainer Stefan Filges <stefan.filges@gu.se>

Description Tools for analyzing sequencing data containing unique molecular identifiers generated by 'UMIErrorCorrect' (<<https://github.com/stahlberggroup/umierrorcorrect>>).

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URL <https://github.com/sfilges/umiAnalyzer>

BugReports <https://github.com/sfilges/umiAnalyzer/issues>

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Author Stefan Filges [aut, cre] (ORCID: <<https://orcid.org/0000-0002-5994-6699>>),
Gustav Johansson [ctb]

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addMetaData

Add metaData

Description

Add metaData

Usage

```
addMetaData(object, attributeName, attributeValue)
```

Arguments

object R object to which meta data should be added
 attributeName Name of the meta data attribute.
 attributeValue Meta data to be saved.

Value

A UMIexperiment object

Examples

```
library(umiAnalyzer)

main <- system.file("extdata", package = "umiAnalyzer")

simSen <- createUmiExperiment(main)

metaData <- system.file("extdata", "metadata.txt", package = "umiAnalyzer")

simSen <- addMetaData(simSen, 'metaData', metaData)
```

addUmiSample

Add UMI sample to an existing experiment object

Description

Add UMI sample to an existing experiment object

Usage

```
addUmiSample(object, sampleName, sampleDir, clearData = FALSE)
```

Arguments

object UMIexperiment object
 sampleName Name of new sample
 sampleDir Directory to new sample
 clearData Should other data in UMIexperiment be cleared

Value

A UMIexperiment object

AmpliconHeatmap	<i>Amplicon heatmap</i>
-----------------	-------------------------

Description

Generates a heatmap of mutations with sample clustering using pheatmap.

Usage

```
AmpliconHeatmap(
  object,
  filter.name = "default",
  cut.off = 5,
  left.side = "columns",
  amplicons = NULL,
  samples = NULL,
  abs.count = FALSE,
  font.size = 10
)
```

Arguments

object	Requires a UMI sample or UMI experiment object
filter.name	Name of the filter to be plotted.
cut.off	How many variant reads are necessary to consider a variant above background? Default is 5 reads.
left.side	Show assays or sample on the left side of the heatmap. Default is assays
amplicons	(Optional) character vector of amplicons to be plotted.
samples	(Optional) character vector of samples to be plotted.
abs.count	Logical. Should absolute counts be used instead of frequencies?
font.size	Font size to use for sample labels

Value

A graphics object

Examples

```
## Not run:
library(umiAnalyzer)

main = system.file('extdata', package = 'umiAnalyzer')
samples <- list.dirs(path = main, full.names = FALSE, recursive = FALSE)
simsten <- createUmiExperiment(experimentName = 'example', mainDir = main, sampleNames = samples)
simsten <- filterUmiObject(simsten)
```

```
hmap <- AmpliconHeatmap(simsen)

## End(Not run)
```

AmpliconPlot	<i>Generate Amplicon plots</i>
--------------	--------------------------------

Description

Plots variant allele frequencies or alternate allele counts for chosen samples and assays.

Usage

```
AmpliconPlot(
  object,
  filter.name = "default",
  cut.off = 5,
  min.count = 0,
  min.vaf = 0,
  amplicons = NULL,
  samples = NULL,
  abs.count = FALSE,
  y_min = 0,
  y_max = NULL,
  theme = "classic",
  option = "default",
  direction = "default",
  plot.text = FALSE,
  plot.ref = TRUE,
  stack.plot = FALSE,
  classic.plot = FALSE,
  fdr = 0.05,
  font.size = 6,
  angle = 45,
  use.caller = FALSE,
  use.plotly = TRUE
)
```

Arguments

<code>object</code>	Requires a UMI sample or UMI experiment object
<code>filter.name</code>	Name of the filter to be plotted.
<code>cut.off</code>	How many variant reads are necessary to consider a variant above background? Default is 5 reads.
<code>min.count</code>	Minimum variants counts to plot, default is 0.

<code>min.vaf</code>	Minimum variants allele frequency to plot, default is 0.
<code>amplicons</code>	(Optional) character vector of amplicons to be plotted.
<code>samples</code>	(Optional) character vector of samples to be plotted.
<code>abs.count</code>	Should absolute counts be plotted instead of frequencies? Default is FALSE.
<code>y_min</code>	Minimum y-axis value, default is 0
<code>y_max</code>	Maximum y-axis value, default is NULL (autoscale)
<code>theme</code>	Plotting theme to use, default is classic.
<code>option</code>	Color palette to use.
<code>direction</code>	Orientation of the color palette.
<code>plot.text</code>	Should non-references bases be indicated above the bar?
<code>plot.ref</code>	If true show reference base instead of position on x-axis.
<code>stack.plot</code>	Show all variant alleles in a stacked bar plot.
<code>classic.plot</code>	Show classical debarcer amplicon plot with raw error.
<code>fdr</code>	False-discovery-rate cut-off for variants.
<code>font.size</code>	Font size
<code>angle</code>	Font angle
<code>use.caller</code>	Should data from variant caller be used? Default is FALSE
<code>use.plotly</code>	Should plotly be used instead of the regular ggplot device? Default is TRUE

Value

A UMIexperiment object containing a ggplot object with the amplicon plot.

Examples

```
library(umiAnalyzer)

main = system.file('extdata', package = 'umiAnalyzer')
samples <- list.dirs(path = main, full.names = FALSE, recursive = FALSE)
simsten <- createUmiExperiment(experimentName = 'example', mainDir = main, sampleNames = samples)
simsten <- filterUmiObject(simsten)

amplicon_plot <- AmpliconPlot(simsten)
```

`BarcodeFamilyHistogram`*Consensus depth histograms*

Description

Generate histograms for the frequency of barcode family depths.

Usage

```
BarcodeFamilyHistogram(  
  object,  
  xMin = 0,  
  xMax = 100,  
  samples = NULL,  
  option = "viridis",  
  direction = 1,  
  theme = "classic"  
)
```

Arguments

<code>object</code>	Requires a UMI sample or UMI experiment object
<code>xMin</code>	Minimum consensus family size to plot, default is 0.
<code>xMax</code>	Maximum consensus family size to plot. Default is 100.
<code>samples</code>	List of samples to be shown.
<code>option</code>	Color scheme to use
<code>direction</code>	If using viridis colors sets the orientation of color scale.
<code>theme</code>	ggplot theme to use. Defaults to classic.

Value

A ggplot object

Examples

```
library(umiAnalyzer)  
main = system.file('extdata', package = 'umiAnalyzer')  
simSen <- createUmiExperiment(main, importBam = TRUE)  
barcode_dist <- BarcodeFamilyHistogram(simSen)
```

beta_binom	<i>Beta binomial model</i>
------------	----------------------------

Description

Code was obtained from VGAM package function VGAM::rbetabinom.ab. The VGAM package is available under the GPL-3 license and maintained by Thomas Yee <t.yee at auckland.ac.nz>. Source code of the function is identical to rbetabinom.ab, but the function name was changed to beta_binom.

Usage

```
beta_binom(n, size, shape1, shape2, limit.prob = 0.5, .dontuse.prob = NULL)
```

Arguments

n	n
size	size
shape1	alpha
shape2	beta
limit.prob	0.5
.dontuse.prob	NULL

Value

Numeric

References

Yee TW (2015). Vector Generalized Linear and Additive Models: With an Implementation in R. Springer, New York, USA.

Examples

```
beta_binom(10,5, 0.5, 1)
beta_binom(10,2, 0.5, 1)
```

callVariants	<i>callVariants using beta binomial distribution</i>
--------------	--

Description

Calculate variant p-values using permutation-based testing. A prior is fitted to model the background error using maximum likelihood estimation of a beta distribution. The maximum likelihood estimate of the beta distribution is then used to define the shape of a beta-binomial distribution used to estimate variant P-Values. This can be interpreted as a probability for a variant to not have arisen by chance.

Usage

```
callVariants(object, minDepth = 3, minCoverage = 100, computePrior = FALSE)
```

Arguments

object	A UMIErrorCorrect object.
minDepth	Minimum consensus depth required default is 3
minCoverage	Minimum Coverage to use, default is 100 reads.
computePrior	Should a new distribution be derived from data? Default is FALSE.

Value

Object containing raw and FDR-adjusted P-Values

See Also

[filterVariants](#) on how to filter variants.

Examples

```
library(umiAnalyzer)
main <- system.file("extdata", package = "umiAnalyzer")

simSen <- createUmiExperiment(main)
simSen <- filterUmiObject(simSen)
simSen <- callVariants(simSen, computePrior = FALSE)
```

createUmiExperiment	<i>Method for creating a UMI experiment object</i>
---------------------	--

Description

Method for creating a UMI experiment object

Usage

```
createUmiExperiment(  
  mainDir,  
  experimentName = NULL,  
  sampleNames = NULL,  
  importBam = FALSE,  
  as.shiny = FALSE  
)
```

Arguments

mainDir	Main experiment directory
experimentName	Name of the experiment
sampleNames	List of sample names. Can be either NULL or list. If NULL all subdirectories of mainDir will be searched.
importBam	Logical. Should bam files be imported on creation? Default is False.
as.shiny	Set to TRUE if run within a shiny::withProgress environment

Value

An object of class UMIexperiment

Examples

```
library(umiAnalyzer)  
  
main = system.file('extdata', package = 'umiAnalyzer')  
  
samples <- list.dirs(path = main, full.names = FALSE, recursive = FALSE)  
  
exp1 <- createUmiExperiment(experimentName = 'exp1', mainDir = main, sampleNames = samples)
```

`createUMIexperiment_Debarcer`*Method for creating a UMI experiment object*

Description

Method for creating a UMI experiment object

Usage

```
createUMIexperiment_Debarcer(experiment.name, main.dir, dir.names)
```

Arguments

<code>experiment.name</code>	Name of the experiment
<code>main.dir</code>	Main experiment directory
<code>dir.names</code>	List of sample names

Value

A UMIexperiment object

`createUmiSample`*createUmiSample*

Description

Method for creating a UMI sample from UMIErrorCorrect output.

Usage

```
createUmiSample(sampleName, sampleDir, importBam = FALSE)
```

Arguments

<code>sampleName</code>	UMI sample object name
<code>sampleDir</code>	Path to UMI sample folders. Must be a folder generated by UMIErrorCorrect
<code>importBam</code>	Logical. Should BAM files be imported at object initialization? Default is False.

Value

An object of class UMiSample

Examples

```
library(umiAnalyzer)

main = system.file('extdata', package = 'umiAnalyzer')
samples <- list.dirs(path = main, full.names = FALSE, recursive = FALSE)
s1 <- createUmiSample('s1', sampleDir = paste(main, "/", samples[1], sep = ""))
```

```
createUMIsample_Debarcoder
```

Method for creating a UMIsample object

Description

Method for creating a UMIsample object

Usage

```
createUMIsample_Debarcoder(sample.name, sample.dir, cons = "10")
```

Arguments

sample.name	UMI sample object name
sample.dir	Path to UMI sample
cons	Consensus depth. Needs to be string; default is 10.

Value

A UMIsample object

```
download_template
```

Download meta data template

Description

Function for downloading a template file containing metadata.

Usage

```
download_template(object)
```

Arguments

object	A UMIexperiment object
--------	------------------------

Value

A tibble containing a metadata template

Examples

```
library(umiAnalyzer)

main <- system.file("extdata", package = "umiAnalyzer")

simsen <- createUmiExperiment(main)

download_template(simsen)
```

filterUmiObject	<i>Method for filtering UMIexperiment and sample objects</i>
-----------------	--

Description

Method for filtering UMIexperiment and sample objects

Usage

```
filterUmiObject(
  object,
  name = "default",
  minDepth = 3,
  minCoverage = 100,
  minFreq = 0,
  minCount = 0
)
```

Arguments

object	Requires a UMI sample or UMI experiment object.
name	String. Name of the filter. Default is "default".
minDepth	Consensus depth to analyze. Default is 3.
minCoverage	Minimum coverage required for amplicons. Default is 1.
minFreq	Minimum variant allele frequency to keep. Default is 0.
minCount	Minimum variant allele count to keep. Default is 3.

Value

A UMI sample or UMI experiment object.

Examples

```
library(umiAnalyzer)

main = system.file('extdata', package = 'umiAnalyzer')

samples <- list.dirs(path = main, full.names = FALSE, recursive = FALSE)

simSen <- createUmiExperiment(experimentName = 'simSen', mainDir = main, sampleNames = samples)

simSen <- filterUmiObject(simSen)
```

filterVariants	<i>Filter variants based on p values or depth</i>
----------------	---

Description

You can filter variants called with the the "callVariants" function based on adjusted p-value, minimum variant allele count and supply a list of assays and samples to plot.

Usage

```
filterVariants(
  object,
  p.adjust = 0.2,
  minVarCount = 5,
  amplicons = NULL,
  samples = NULL
)
```

Arguments

object	A UMIexperiment object
p.adjust	Numeric. Adjusted p value (FDR). Default is 0.2.
minVarCount	Integer. Minimum variant allele count. Default is 5.
amplicons	NULL or list of assays to plot. NULL uses all.
samples	NULL or list of samples to plot. NULL uses all.

Value

A UMIexperiment object with filtered variants. Can be used to generate VCF files.

See Also

[callVariants](#) on how to call variants.

Examples

```
## Not run:
library(umiAnalyzer)
main <- system.file("extdata", package = "umiAnalyzer")

simSen <- createUmiExperiment(main)
simSen <- filterUmiObject(simSen)
simSen <- callVariants(simSen, computePrior = FALSE)
simSen <- filterVariants(simSen, p.adjust = 0.05)

## End(Not run)
```

findConsensusReads	<i>Find consensus reads A function to analyze consensus read tables generated with parseBamFiles or a UMIexperiment object containing reads.</i>
--------------------	--

Description

Find consensus reads A function to analyze consensus read tables generated with parseBamFiles or a UMIexperiment object containing reads.

Usage

```
findConsensusReads(
  object,
  consDepth = 0,
  groupBy = c("none", "sample", "position", "both"),
  pattern = NULL
)
```

Arguments

object	Either a tibble generated with parseBamFiles or a UMIexperiment object
consDepth	Minimum consensus depth to keep. Default is 0.
groupBy	Should data be grouped by position, sample, both or not at all.
pattern	Regular expression

Value

A data table

Examples

```
library(umiAnalyzer)
main <- system.file("extdata", package = "umiAnalyzer")
simssen <- createUmiExperiment(main, importBam = TRUE)

reads <- findConsensusReads(simssen)
reads
```

generateVCF

*Generate VCF file from UMI sample or UMI experiment object***Description**

Generate VCF file from UMI sample or UMI experiment object

Usage

```
generateVCF(object, outDir = getwd(), outFile, printAll = FALSE)
```

Arguments

object	Requires a UMI sample or UMI experiment object
outDir	String. Output directory, defaults to working directory.
outFile	String. Name of the output file
printAll	Logical. Should all or only trusted variant be printed?

Value

A VCF file

Examples

```
## Not run:
library(umiAnalyzer)

main <- system.file("extdata", package = "umiAnalyzer")

simssen <- createUmiExperiment(main)

simssen <- filterUmiObject(simssen)

generateVCF(simssen, 'simssen.vcf', printAll = FALSE, save = FALSE)

## End(Not run)
```

getFilteredData	<i>Method for retrieving filtered data</i>
-----------------	--

Description

Method for retrieving filtered data

Usage

```
getFilteredData(  
  object,  
  name = "default",  
  save = FALSE,  
  outDir = getwd(),  
  fileName = NULL,  
  delim = ";"  
)
```

Arguments

object	Requires a UMI sample or UMI experiment object.
name	String. Name of the filter. Default is "default".
save	Logical, should data be saved as csv file? Default is FALSE.
outDir	Output directory
fileName	Filename to be used, default is the same as 'name'
delim	Character string denoting delimiter to be used, default is ';'.

Value

A filtered consensus table, as a tibble.

Examples

```
library(umiAnalyzer)  
  
main = system.file('extdata', package = 'umiAnalyzer')  
  
samples <- list.dirs(path = main, full.names = FALSE, recursive = FALSE)  
  
simсен <- createUmiExperiment(experimentName = 'simсен',mainDir = main,sampleNames = samples)  
simсен <- filterUmiObject(simсен)  
  
myfilter <- getFilteredData(simсен)  
myfilter
```

getMetaData	<i>Retrieve meta data by name.</i>
-------------	------------------------------------

Description

Retrieve meta data by name.

Usage

```
getMetaData(object, attributeName)
```

Arguments

object	R object from which to get meta data.
attributeName	Name of the meta data attribute.

Value

Metadata

Examples

```
library(umiAnalyzer)

main <- system.file("extdata", package = "umiAnalyzer")

simSen <- createUmiExperiment(main)

metaData <- system.file("extdata", "metadata.txt", package = "umiAnalyzer")

simSen <- importDesign(object = simSen, file = metaData)
design <- getMetaData(object = simSen, attributeName = "design")
design
```

importBedFile	<i>Import bed file</i>
---------------	------------------------

Description

Import bed file

Usage

```
importBedFile(path)
```

Arguments

path path to bed file

Value

A table containing genome positions

importDesign	<i>Import experimental design meta data such as replicates, treatments, categorical variables.</i>
--------------	--

Description

Import experimental design meta data such as replicates, treatments, categorical variables.

Usage

```
importDesign(object, file, delim = NULL)
```

Arguments

object UMI.experiment to which to add metadata
file File containing meta data
delim Column separator. Default is NULL (automatically determine delimiter)

Value

A UMIexperiment object

Examples

```
library(umiAnalyzer)

main <- system.file("extdata", package = "umiAnalyzer")

simSen <- createUmiExperiment(main)

metaData <- system.file("extdata", "metadata.txt", package = "umiAnalyzer")

simSen <- importDesign(object = simSen, file = metaData)

# Retrieve meta data
design <- getMetaData(object = simSen, attributeName = "design")
design
```

mergeAssays	<i>Merge assays</i>
-------------	---------------------

Description

Merge assays together by name. Requires a name of the new assay and a list of assays that will be merged.

Usage

```
mergeAssays(object, name, assay.list)
```

Arguments

object	A UMIexperiment object
name	Name of the new assay
assay.list	List of assays to merge

Value

merged consensus data

Examples

```
library(umiAnalyzer)

main <- system.file("extdata", package = "umiAnalyzer")

simsen <- createUmiExperiment(main)

simsen <- mergeAssays(object = simsen, name = "new", assay.list = c("PIK3CA_123", "PIK3CA_234"))
```

parseBamFiles	<i>Function to parse bam files</i>
---------------	------------------------------------

Description

Function to parse bam files

Usage

```
parseBamFiles(mainDir, sampleNames = NULL, consDepth = 0, as.shiny = FALSE)
```

Arguments

<code>mainDir</code>	Directory containing UMIErrorCorrect output folders.
<code>sampleNames</code>	A list of sample names.
<code>consDepth</code>	Only retain consensus reads of at least <code>cons.depth</code> . Default is 0.
<code>as.shiny</code>	Set to TRUE if run within a shiny::withProgress environment

Value

A data table

Examples

```
library(umiAnalyzer)
main <- system.file("extdata", package = "umiAnalyzer")
simsen <- createUmiExperiment(main)

reads <- parseBamFiles(main, consDepth = 10)
```

QCplot

Generate QC plots

Description

Visualize the UMI count for each selected assay and sample for a given consensus depth. This is useful to detect differences in coverage, especially for multiplexed assays.

Usage

```
QCplot(
  object,
  group.by = "sample",
  plotDepth = 3,
  assays = NULL,
  samples = NULL,
  theme = "classic",
  option = "viridis",
  direction = "default",
  toggle_mean = TRUE,
  center = "mean",
  line_col = "blue",
  angle = 0,
  plotly = FALSE
)
```

Arguments

object	Requires a UMI sample or UMI experiment object
group.by	String. Which variable should be used as a factor on the x-axis. Default is sample
plotDepth	Which consensus depth to plot
assays	(Optional) user-supplied list of assays to plot. Default is all.
samples	(Optional) user-supplied list of samples to plot. Default is all.
theme	ggplot theme to use.
option	Color palette to use, either ggplot default or viridis colors.
direction	If viridis colors are used, choose orientation of color scale.
toggle_mean	Show mean or median
center	Choose mean or median
line_col	Choose color for mean/median line
angle	Angle of labels on x-axis.
plotly	Should plotly be used for rendering?

Value

A ggplot object

Examples

```
library(umiAnalyzer)

main = system.file('extdata', package = 'umiAnalyzer')
samples <- list.dirs(path = main, full.names = FALSE, recursive = FALSE)
simsen <- createUmiExperiment(experimentName = 'example', mainDir = main, sampleNames = samples)

depth_plot <- QCplot(simsen)
```

runUmiVisualizer

Function to run the umiVisualizer shiny app

Description

Function to run the umiVisualizer shiny app

Usage

```
runUmiVisualizer()
```

Value

Opens the umiVisualizer app

Examples

```
## Not run:
library(umiAnalyzer)

runUmiVisualizer()

## End(Not run)
```

saveConsData	<i>Save consensus data</i>
--------------	----------------------------

Description

If save is set to TRUE data will be written to a csv file otherwise consensus data will be returned as a tibble.

Usage

```
saveConsData(
  object,
  save = FALSE,
  fileName = "consensus_data.csv",
  outDir = getwd(),
  delim = ";"
)
```

Arguments

object	UMIexperiment object
save	Logical. Should data be saved to file? Default is FALSE.
fileName	String. Name of the file to be saved. Default is 'consensus_data.csv'
outDir	output directory, defaults to working directory
delim	Single character string, either ';' or ',' or tab

Value

A data table

Examples

```
library(umiAnalyzer)

main = system.file('extdata', package = 'umiAnalyzer')

samples <- list.dirs(path = main, full.names = FALSE, recursive = FALSE)
```

```
example <- createUmiExperiment(experimentName = 'example',mainDir = main,sampleNames = samples)

consensus_data <- saveConsData(object = example)
consensus_data
```

simsen

UMIexperiment data generated with SiMSen-Seq

Description

UMIexperiment data generated with SiMSen-Seq

Format

An object of class "UMIexperiment"

timeSeriesGrid

Plot time series data

Description

Function for plotting time series or other meta data. Uses facet wrap to display user-provided categorical variables.

Usage

```
timeSeriesGrid(
  object,
  filter.name = "default",
  cut.off = 5,
  min.count = 0,
  min.vaf = 0,
  amplicons = NULL,
  samples = NULL,
  x_variable = NULL,
  y_variable = "Max Non-ref Allele Frequency",
  columns = "Sample Name",
  rows = "Name",
  color_by = "Name",
  fdr = 0.05,
  use.caller = TRUE,
  bed_positions = NULL
)
```

Arguments

object	A consensus data table
filter.name	"default"
cut.off	5
min.count	0
min.vaf	0
amplicons	NULL
samples	NULL
x_variable	NULL
y_variable	"Max Non-ref Allele Frequency"
columns	"Sample Name"
rows	"Name"
color_by	"Name"
fdr	0.05
use.caller	TRUE
bed_positions	NULL

Value

A ggplot object.

Examples

```
library(umiAnalyzer)

main <- system.file("extdata", package = "umiAnalyzer")
simsen <- createUmiExperiment(main)
simsen <- filterUmiObject(simsen)

metaData <- system.file("extdata", "metadata.txt", package = "umiAnalyzer")
simsen <- importDesign(object = simsen, file = metaData)

bed_dir <- system.file("extdata", "simple.bed", package = "umiAnalyzer")
bed <- importBedFile(path = bed_dir)

time_plot <- timeSeriesGrid(simsen, x_variable = "time", bed_positions = bed)
```

UmiCountsPlot

Plot UMI counts

Description

Visualize the number detected UMI for each consensus depth cut-off. This may be helpful in choosing the right consensus depth for your analysis, by checking the number of reads still available for each assay and sample for your chosen cut-off.

Usage

```
UmiCountsPlot(
  object,
  amplicons = NULL,
  samples = NULL,
  theme = "classic",
  option = "viridis",
  direction = 1
)
```

Arguments

object	Requires a UMI sample or UMI experiment object
amplicons	(Optional) user-supplied list of assays to plot. Default is all.
samples	(Optional) user-supplied list of samples to plot. Default is all.
theme	Plotting theme, default is classic
option	Color palette. Default uses ggplot standard, otherwise viridis options.
direction	If using viridis colors should the scale be inverted or default?

Value

A UMIexperiment object

Examples

```
library(umiAnalyzer)

main = system.file('extdata', package = 'umiAnalyzer')
samples <- list.dirs(path = main, full.names = FALSE, recursive = FALSE)
simSen <- createUmiExperiment(experimentName = 'example', mainDir = main, sampleNames = samples)
simSen <- filterUmiObject(simSen)

count_plot <- UmiCountsPlot(simSen)
```

UMIexperiment-class	<i>UMIexperiment class</i>
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Description

The UMIexperiment is the core data object, storing all data and relevant analysis data associated with your experiment. Each object has number of slots storing raw data, graphs and processed data.

Value

An object of class UMIexperiment

Slots

name Optional project name for record keeping.
cons.data The raw consensus data supplied by the user.
summary.data Summary data from UMIErrorCorrect
raw.error Cons0 error profile
reads Consensus reads imported using the parseBamFiles function.
meta.data Sample data optionally supplied by the user.
filters A list of filtered cons.data, which can be accessed separately.
plots A list of generated plots.
variants Consensus table generated with the umiAnalyzer variant caller.
merged.data Data generated using the mergeTechnicalReplicates function.

UMIsample-class	<i>UMIsample class</i>
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Description

UMIsample class

Value

An object of class UMIsample

Slots

name Sample name
cons.data Raw consensus data
summary.data Summary data from UMIErrorCorrect
reads Consensus reads imported from a bam file.

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