Package 'omicsTools'

July 3, 2023

```
Title Omics Data Process Toolbox
Version 1.0.5
Date 2023-06-29
Description Processing and analyzing omics data from genomics, transcriptomics, pro-
      teomics, and metabolomics platforms. It provides functions for preprocessing, normalization, vi-
      sualization, and statistical analysis, as well as machine learning algorithms for predictive model-
      ing. 'omicsTools' is an essential tool for researchers working with high-
      throughput omics data in fields such as biology, bioinformatics, and medicine. The QC-
      RLSC (quality control-based robust LOESS signal correction) algorithm is used for normaliza-
      tion. Dunn et al. (2011) <doi:10.1038/nprot.2011.335>.
License MIT + file LICENSE
Imports bs4Dash, config (>= 0.3.1), dplyr, DT, golem (>= 0.3.5),
      magrittr, readr, shiny (>= 1.7.2), tibble
Encoding UTF-8
RoxygenNote 7.2.3
Suggests spelling, testthat (>= 3.0.0)
Config/testthat/edition 3
URL https://github.com/YaoxiangLi/omicsTools
BugReports https://github.com/YaoxiangLi/omicsTools/issues
NeedsCompilation no
Author Yaoxiang Li [cre, aut] (<a href="https://orcid.org/0000-0001-9200-1016">https://orcid.org/0000-0001-9200-1016</a>),
      Zihao Zhang [aut],
      Meth Jayatilake [aut],
      Amrita Cheema [aut] (<a href="https://orcid.org/0000-0003-4877-7583">https://orcid.org/0000-0003-4877-7583</a>)
Maintainer Yaoxiang Li < liyaoxiang@outlook.com>
Repository CRAN
Date/Publication 2023-07-03 16:20:02 UTC
```

2 impute

R topics documented:

Index																					5
	run_app	 	٠.		 		 •	 •		 •		 •			•	 •		•			4
	normalize	 			 																3
	impute	 			 	•	 •	 •	•	 •	•	 •	•	•		 •	•	•	•	•	2

impute

Impute function

Description

This function performs data cleaning and imputation on a given data matrix. It removes blank and NIST samples, features with NA values more than the specified threshold, and imputes remaining NA values with half of the smallest non-NA value.

Usage

```
impute(data, percent = 0.2)
```

Arguments

data

A data frame containing the sample data. The first column should contain the

sample identifiers, and the rest of the columns contain the peaks.

percent

A numeric value between 0 and 1 representing the threshold of the percentage of NA values a feature should have for it to be removed from the dataset. Default

value is 0.2.

Value

A data frame with the first column as the sample identifiers and the rest of the columns containing the cleaned and imputed peak intensities.

Author(s)

```
Yaoxiang Li <yl814@georgetown.edu>
Georgetown University, USA
License: GPL (>= 3)
```

Examples

```
# Load the CSV data
data_file <- system.file("extdata", "example1.csv", package = "omicsTools")</pre>
data <- readr::read_csv(data_file)</pre>
# Apply the impute function
imputed_data <- omicsTools::impute(data, percent = 0.2)</pre>
```

normalize 3

```
# Write the imputed data to a new CSV file
readr::write_csv(imputed_data, paste0(tempdir(), "/imputed_data.csv"))
```

normalize

Normalize function

Description

This function performs normalization on the input data matrix using the loess regression method. Normalization is done based on Quality Control (QC) samples in the data.

Usage

```
normalize(data)
```

Arguments

data

A data frame containing the sample data. The first column should contain the sample identifiers, and the rest of the columns contain the peaks to be normalized. QC samples should be indicated in the sample identifiers with 'QC'.

Value

A data frame with the first column as the sample identifiers and the rest of the columns containing the normalized peak intensities.

Author(s)

```
Yaoxiang Li <y1814@georgetown.edu>
Georgetown University, USA
License: GPL (>= 3)
```

Examples

```
# Load the CSV data
data_file <- system.file("extdata", "example2.csv", package = "omicsTools")
data <- readr::read_csv(data_file)
# Apply the normalize function
normalized_data <- omicsTools::normalize(data)

# Write the normalized data to a new CSV file
readr::write_csv(normalized_data, paste0(tempdir(), "/normalized_data.csv"))</pre>
```

run_app

run_app

Run the Shiny Application

Description

Run the Shiny Application

Usage

```
run_app(
  onStart = NULL,
  options = list(),
  enableBookmarking = NULL,
  uiPattern = "/",
  ...
)
```

Arguments

onStart

A function that will be called before the app is actually run. This is only needed for shinyAppObj, since in the shinyAppDir case, a global.R file can be used

for this purpose.

options

Named options that should be passed to the runApp call (these can be any of the following: "port", "launch.browser", "host", "quiet", "display.mode" and "test.mode"). You can also specify width and height parameters which provide a hint to the embedding environment about the ideal height/width for the app.

enableBookmarking

Can be one of "url", "server", or "disable". The default value, NULL, will re-

spect the setting from any previous calls to enableBookmarking(). See enableBookmarking()

for more information on bookmarking your app.

uiPattern

A regular expression that will be applied to each GET request to determine whether the ui should be used to handle the request. Note that the entire request path must match the regular expression in order for the match to be considered suc-

cessful.

arguments to pass to golem_opts. See '?golem::get_golem_options' for more

details.

Value

No return value, called for launch the application.

Index

```
enableBookmarking(), 4
impute, 2
normalize, 3
run_app, 4
```