

Package ‘msQC’

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Type Package

Title An R package for proteomics data quality control

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Description This package creates a HTML format QC report for MS/MS-based proteomics data. The report is intended to allow the user to quickly assess the quality of proteomics data.

Depends R (>= 3.0.0), XML, VennDiagram

Imports rTANDEM, plyr, seqinr, Nozzle.R1, ggplot2, reshape2, parallel

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Suggests RforProteomics (>= 1.0.16), knitr, BiocStyle, rpx, R.utils,RUnit,BiocGenerics

Collate 'msQC.R' 'report.R' 'visualization.R'

VignetteBuilder knitr

biocViews Proteomics, MassSpectrometry, QualityControl, Visualization,ReportWriting

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addSummaryChart	<i>Add PRIDE summary charts</i>
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Description

Add PRIDE summary charts in the technical replicate level

Usage

```
addSummaryChart(res)
```

Arguments

res	An object returned by msQCpipe function
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cntStat	<i>contaminants stat</i>
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Description

Common Contaminants in Proteomics Mass Spectrometry Experiments

Usage

```
cntStat(res)
```

Arguments

res	An object of msQCres
-----	----------------------

Value

A data.frame will be shown in HTML report

combineRun	<i>Combine multiple results</i>
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Description

Combine multiple results

Usage

```
combineRun(pepFiles, fasta, outputPathFile, outdir, prefix)
```

Arguments

pepFiles	peptideSummary files
fasta	database file
outputPathFile	out file
outdir	output directory
prefix	output prefix

Value

A data.frame

Author(s)

Bo Wen <wenbo@genomics.cn>

createTargetDecoyDB	<i>Create target-decoy database</i>
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Description

Create target-decoy database

Usage

```
createTargetDecoyDB(fa, outdb)
```

Arguments

fa	target database
outdb	output target-decoy database

Value

target-decoy database file name

Author(s)

Bo Wen <wenbo@genomics.cn>

getEnzyme

Get the enzymes list

Description

Get the enzymes list

Usage

getEnzyme()

Value

A data frame which contains all of the enzymes

Author(s)

Bo Wen <wenbo@genomics.cn>

getMods

Get the modification list

Description

Get the modification list

Usage

getMods()

Value

A data frame which contains all of the modifications

Author(s)

Bo Wen <wenbo@genomics.cn>

loadmsQCres	<i>Load the result of msQCpipe</i>
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Description

Load the result of [msQCpipe](#)

Usage

```
loadmsQCres(outdir)
```

Arguments

outdir The output directory of [msQCpipe](#)

Author(s)

Laurent Gatto <lg390@cam.ac.uk>, Bo Wen <wenbo@genomics.cn>

Examples

```
zpqc <- system.file("extdata/qc.zip", package = "msQC")
unzip(zpqc)
qcres <- loadmsQCres("./qc")
```

msQCpipe	<i>The main function of msQC pipeline</i>
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Description

This function is designed to automate generating of target-decoy database, database searching and post-processing.

Usage

```
msQCpipe(spectralist = NULL, fasta = "", outdir = "./", mode = "",
  miss = 2, enzyme = 1, varmod = c(2, 3, 4), fixmod = c(1), tol = 10,
  tolu = "ppm", itol = 0.6, itolu = "Daltons", threshold = 0.01,
  cpu = 0, xmx = 2, ...)
```

Arguments

spectralist	A file contains the experiment design
fasta	database file, must contain decoy sequences
outdir	output directory
mode	identification or quantification
miss	max miss cleavage
enzyme	enzyme
varmod	Variable modifications are those which may or may not be present.
fixmod	Fixed modifications are applied universally, to every instance of the specified residue(s) or terminus.
tol	The error window on experimental peptide mass values
tolu	Units can be selected from: ppm, Daltons.
itol	Error window for MS/MS fragment ion mass values.
itolu	Units can be selected from: Daltons
threshold	FDR value for PSM
cpu	Max number of cpu used
xmx	JAVA -Xmx
...	Additional parameters passed to read.table used to read the experimental design.

Value

A list which contains all of the information for data quality report generating

Author(s)

Bo Wen <wenbo@genomics.cn>

Examples

```
## Not run:
library("rpx")
px <- PXDataset("PXD000864")
mgfs <- grep("mgf", pxfiles(px), value = TRUE)
mgfs <- grep("-0[5-6]-[1|2]", mgfs, value=TRUE)
mgffiles <- pxget(px, mgfs)
library("R.utils")
mgffiles <- sapply(mgffiles, gunzip)
## Generate the lightweight qc report,
## trim the mgf files to 1/10 of their size.
trimMgf <- function(f, m = 1/10, overwrite = FALSE) {
  message("Reading ", f)
  x <- readLines(f)
  beg <- grep("BEGIN IONS", x)
  end <- grep("END IONS", x)
```

```
n <- length(beg)
message("Sub-setting to ", m)
i <- sort(sample(n, floor(n * m)))
k <- unlist(mapply(seq, from = beg[i], to = end[i]))
if (overwrite) {
  unlink(f)
  message("Writing ", f)
  writeLines(x[k], con = f)
  return(f)
} else {
  g <- sub(".mgf", "_small.mgf", f)
  message("Writing ", g)
  writeLines(x[k], con = g)
  return(g)
}
}
set.seed(1)
mgffiles <- sapply(mgffiles, trimMgf, overwrite = TRUE)
fas <- pxget(px, "TTE2010.zip")
fas <- unzip(fas)
design <- system.file("extdata/PXD000864-design.txt", package = "msQC")
read.table(design, header = TRUE)
qcres <- msQCpipe(spectralist = design,
                 fasta = fas,
                 outdir = "./qc",
                 miss = 0,
                 enzyme = 1, varmod = 2, fixmod = 1,
                 tol = 10, itol = 0.6, cpu = 2,
                 mode = "identification")
html <- reportHTML(qcres)

## End(Not run)
```

plotBioRepVenn

Venn plot in biological replicate level

Description

Venn plot in biological replicate level

Usage

```
plotBioRepVenn(res)
```

Arguments

res An object of msQCres

Value

The name of the figure

plotFractionIDResult *Barplot in different level for each fraction*

Description

Barplot in different level for each fraction

Usage

```
plotFractionIDResult(res, level = NA)
```

Arguments

res	An object of msQCres
level	1: total spectrum, 2: identified spectrum, 3: identified peptide, 4: identified protein.

Value

The name of the figure

plotMS1Error *plot MS1 mass error*

Description

plot MS1 mass error

Usage

```
plotMS1Error(res, plot.class = "ppm")
```

Arguments

res	An object of msQCres
plot.class	ppm or da

Value

The name of the figure

plotMS2Error	<i>plot MS2 mass error</i>
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Description

plot MS2 mass error

Usage

```
plotMS2Error(res)
```

Arguments

res	An object of msQCres
-----	----------------------

Value

The name of the figure

plotSampleIDResultErrorBar	<i>Error barplot in different level for each fraction</i>
----------------------------	---

Description

Error Barplot in different level for each fraction

Usage

```
plotSampleIDResultErrorBar(res, level = NA)
```

Arguments

res	An object of parser result
level	1: total spectrum, 2: identified spectrum, 3: identified peptide, 4: identified protein.

Value

The name of the figure

plotSampleVenn *Venn plot in sample level*

Description

Venn plot in sample level

Usage

```
plotSampleVenn(res)
```

Arguments

res An object of msQCres

Value

The name of the figure

plotTechRepVenn *Venn plot in technical replicate level*

Description

Venn plot in technical replicate level

Usage

```
plotTechRepVenn(res)
```

Arguments

res An object of msQCres

Value

The name of the figure

print.msQCres	<i>Print the information of msQCres object</i>
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Description

Print the information of msQCres object

Usage

```
## S3 method for class msQCres  
print(x, ...)
```

Arguments

x	A msQCres object
...	Additional parameters

Author(s)

Laurent Gatto <lg390@cam.ac.uk>, Bo Wen <wenbo@genomics.cn>

Examples

```
zpqc <- system.file("extdata/qc.zip", package = "msQC")  
unzip(zpqc)  
qcres <- loadmsQCres("./qc")  
print.msQCres(qcres)
```

reportHTML	<i>HTML format report generator</i>
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Description

HTML format report generator

Usage

```
reportHTML(res)
```

Arguments

res	An object returned by <code>msQCpipe</code> function
-----	--

Value

null

Author(s)

Bo Wen <wenbo@genomics.cn>

Examples

```
zpqc <- system.file("extdata/qc.zip", package = "msQC")
unzip(zpqc)
qcres <- loadmsQCres("./qc")
html <- reportHTML(qcres)
```

runTandem

Run X!Tandem

Description

Run X!Tandem

Usage

```
runTandem(spectra = "", fasta = "", outdir = "./", outprefix = "",
  cpu = 1, enzyme = 1, xmx = 2, varmod = 2, fixmod = 1, tol = 10,
  tolu = "ppm", itol = 0.6, itolu = "Daltons", miss = 1)
```

Arguments

spectra	MS/MS peak list file
fasta	database file
outdir	output directory
outprefix	output file prefix
cpu	The number of CPU used for X!Tandem
enzyme	The ID of enzyme used for database searching. See showEnzyme .
xmx	Set for parameter of "Java -Xmx".
varmod	Variable modifications used for database searching. See showMods .
fixmod	Fixed modifications used for database searching. See showMods .
tol	The error window on experimental peptide mass values
tolu	Units can be selected from: ppm, Daltons.
itol	Error window for MS/MS fragment ion mass values.
itolu	Units can be selected from: Daltons
miss	Max miss cleavage

Value

a file path

Author(s)

Bo Wen <wenbo@genomics.cn>

showEnzyme	<i>Shown all enzymes</i>
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Description

Shown all enzymes

Usage

```
showEnzyme()
```

Value

A data frame which contains all of the enzymes

Author(s)

Bo Wen <wenbo@genomics.cn>

Examples

```
showEnzyme()
```

showMods	<i>Shown all modifications</i>
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Description

Shown all modifications

Usage

```
showMods()
```

Value

A data frame which contains all of the modifications

Author(s)

Bo Wen <wenbo@genomics.cn>

Examples

```
showMods()
```

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