Package 'EEM'

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(EEM) Data
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Description Read raw EEM data and prepares them for further analysis.
Depends R (>= $3.0.0$)
Imports tools, reshape2, graphics, colorRamps, utils, R.utils, sp, ggplot2
Suggests stats, pls, knitr, testthat
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applejuice

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Apple juice

Description

Apples of each of six types (Aomori–Fuji, Aomori–Jona, Aomori–Ohrin, NZ–Envy, NZ–Jazz, NZ–Fuji) were blended and filtered using a gauze. Fluorescence profiles of complete excitation–emission matrix of filtered solutions (diluted with water to 147 times) were measured using fluorescence spectroscopy machines. The sample name refers to "type–fruit number–replicate". To save space, only two apples of each types were given in the dataset.

Usage

```
data("applejuice")
```

```
data(applejuice)
summary(applejuice)
```

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commonizeEEM

Smooth out the different dimensions of EEM data

Description

Smooth out the difference dimensions of EEM data by finding the common variables of all data and subset those data.

Usage

```
commonizeEEM(EEM)
```

Arguments

EEM

a list containing EEM data as created by readEEM function.

Value

EEM class object with only common variables

Examples

```
data(gluten)
data(applejuice)
data <- c(gluten, applejuice)
summary(data) # different dimensions
data_combined <- commonizeEEM(data)
summary(data_combined) # same dimension, ready for unfold</pre>
```

cutEEM

Cut portions of EEM

Description

Cut portions of EEM

Usage

```
cutEEM(x, cutEX = NULL, cutEM = NULL)
## S3 method for class 'EEM'
cutEEM(x, cutEX = NULL, cutEM = NULL)
## S3 method for class 'EEMweight'
cutEEM(x, cutEX = NULL, cutEM = NULL)
```

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Arguments

Х	a list of EEM data generated by readEEM function or EEMweight object generated by extract-related functions.
cutEX	Numeric or sequential data specifying regions to be cut for excitation wavelength. Examples, 200 or 200:500
cutEM	Numeric or sequential data specifying regions to be cut for emission wavelength. Examples, 200 or 200:500

Value

A list similar to input EEM is returned but with specified portions cut.

Examples

```
data(applejuice)
applejuice_cut <- cutEEM(applejuice, cutEX = 300:450)
drawEEM(applejuice_cut, 1)</pre>
```

delScattering

Delete scattering rays

Description

This function deletes two regions that are not related to fluorescence emission: (1) regions where emission wavelength is shorten than excitation light, (2) scattering rays and their second, third and forth order lights.

Usage

```
delScattering(EEM, rep = 0, first = 30, second = 40, third = 40,
    forth = 40)
```

Arguments

EEM	A list containing EEM data as created by readEEM function.
rep	(optional) Regions to be deleted are to be replaced with rep: 0 or NA
first	(optional) Width of region to be deleted for first order scattering rays [nm]
second	(optional) Width of region to be deleted for second order scattering rays [nm]
third	(optional) Width of region to be deleted for third order scattering rays [nm]
forth	(optional) Width of region to be deleted for forth order scattering rays [nm]

Value

A list similar to input EEM is returned but with all scattering rays deleted.

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References

Fujita, K., Tsuta, M., Kokawa, M., and Sugiyama, J. (2010). Detection of deoxynivalenol using fluorescence excitation–emission matrix. Food and Bioprocess Technology, 3(6), 922–927.

Examples

```
data(applejuice)
drawEEM(delScattering(applejuice, NA), 1)
```

delScattering2

Delete scattering rays

Description

This function deletes three regions that are not related to fluorescence emission: (1) regions where emission wavelength is shorten than excitation light ($Em \le Ex$), (2) scattering rays and their second order light, (3) regions above second-order scattering ($EM \ge 2*EX$)

Usage

```
delScattering2(EEM, rep = 0, first = 30, second = 40)
```

Arguments

EEM	A list containing EEM data as created by readEEM function.	
rep	(optional) Regions to be deleted are to be replaced with rep: 0 or NA	
first	(optional) Width of region to be deleted for first order scattering rays [nm]	
second	(optional) Width of region to be deleted for second order scattering rays [nm]	

Value

A list similar to input EEM is returned but with all scattering rays deleted.

```
data(applejuice)
drawEEM(delScattering2(applejuice, NA), 1)
```

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drawEEM

Draw contour for EEM data

Description

This function is a wrapper function for filled.contour to draw contour for EEM data.

Usage

```
drawEEM(x, ...)
## S3 method for class 'EEM'
drawEEM(x, n, exlab = "Excitation wavelength [nm]",
 emlab = "Emission wavelength [nm]", color.palette = matlab.like,
 nlevels = 50, main = NULL, flipaxis = FALSE, ...)
## S3 method for class 'EEMweight'
drawEEM(x, ncomp, exlab = "Excitation wavelength [nm]",
 emlab = "Emission wavelength [nm]", color.palette = matlab.like,
 nlevels = 50, main = NULL, flipaxis = FALSE, ...)
## S3 method for class 'matrix'
drawEEM(x, n, exlab = "Excitation wavelength [nm]",
 emlab = "Emission wavelength [nm]", color.palette = matlab.like,
 nlevels = 50, main = NULL, flipaxis = FALSE, ...)
## S3 method for class 'data.frame'
drawEEM(x, n, exlab = "Excitation wavelength [nm]",
 emlab = "Emission wavelength [nm]", color.palette = matlab.like,
  nlevels = 50, main = NULL, flipaxis = FALSE, ...)
## S3 method for class 'numeric'
drawEEM(x, exlab = "Excitation wavelength [nm]",
  emlab = "Emission wavelength [nm]", color.palette = matlab.like,
  nlevels = 50, main = NULL, flipaxis = FALSE, ...)
```

Arguments

```
a list of EEM data generated by readEEM function or EEMweight object generated by extract-related functions.

(optional) further arguments passed to other methods of filled.contour sample number. The number should not exceed length(EEM)

exlab

(optional) excitation-axis label

emlab

(optional) emission-axis label

color.palette

(optional) contour color palette. See palette for more details
```

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nlevels	(optional) number of levels used to separate range of intensity value
main	(optional) plot title
flipaxis	(optional) flip axis
ncomp	number of components

Value

A figure is returned on the graphic device

Methods (by class)

- EEM: draw contour of EEM data created by readEEM function
- EEMweight: draw contours of the output from getLoading and getReg.
- matrix: draw contour of unfolded matrix which have column names in the format of EX...EM...
- data.frame: draw contour of unfolded data.frame which have column names in the format of EX...EM...
- numeric: draw contour of a vector of numeric values which have names in the format of EX...EM...

See Also

drawEEM

Examples

```
# method for class "EEM"
data(applejuice)
drawEEM(applejuice, 1) # draw contour of the first sample
drawEEM(applejuice, 1, flipaxis = TRUE) # flip the axis

# method for class "EEMweight"
applejuice_uf <- unfold(applejuice) # unfold list into matrix
result <- prcomp(applejuice_uf)
drawEEM(getLoading(result), 1) # plot loading of the first PC</pre>
```

drawEEMgg

Draw contour for EEM data using ggplot2

Description

This function draw contour for EEM data using ggplot2. Use 'ggsave' to save the contours.

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Usage

```
drawEEMgg(x, ...)

## S3 method for class 'EEM'
drawEEMgg(x, n, textsize = 20, color.palette = matlab.like,
    nlevels = 20, exlab = "Excitation wavelength [nm]",
    emlab = "Emission wavelength [nm]", main = NULL, has_legend = TRUE,
    zlim = NULL, breaks = waiver(), flipaxis = FALSE, ...)

## S3 method for class 'EEMweight'
drawEEMgg(x, ncomp, textsize = 25,
    color.palette = matlab.like, nlevels = 20,
    exlab = "Excitation wavelength [nm]", emlab = "Emission wavelength [nm]",
    main = NULL, has_legend = TRUE, zlim = NULL, breaks = waiver(),
    flipaxis = FALSE, ...)
```

Arguments

x a list of EEM data generated by readEEM function or EEMweight object generated by extract-related functions.

... arguments for other methods

n sample number. The number should not exceed length(EEM)

textsize (optional) text size

color.palette (optional) contour color palette. See palette for more details nlevels (optional) number of levels used to separate range of intensity value

exlab (optional) excitation-axis label emlab (optional) emission-axis label

main (optional) plot title has_legend logical value for legend zlim zlim = c(min, max)

breaks breaks

flipaxis (optional) flip axis ncomp number of components

Details

drawEEM is faster and should be used.

Value

A figure is returned on the graphic device

Methods (by class)

- EEM: draw EEM of EEM data created by readEEM function
- EEMweight: draw contours of the output from getLoading and getReg.

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See Also

drawEEM

Examples

```
## Not run:
require(EEM)
require(ggplot2)
data(applejuice)
drawEEMgg(applejuice, 1) # draw EEM of sample no.1
drawEEMgg(applejuice, 1, color.palette = cm.colors) # draw EEM of sample no.31 with different color
drawEEMgg(applejuice, 1, nlevels = 10) # change nlevels
# manually define legend values
drawEEMgg(applejuice, 1, breaks = seq(from = 1000, to = 6000, by = 1000))
# can be combined with other ggplot2 commands
# add point to the plot
drawEEMgg(applejuice, 1) + geom_point(aes(x = 350, y = 500), pch = 17, cex = 10)
# add grid line to the plot
drawEEMgg(applejuice, 1) + theme(panel.grid = element_line(color = "grey"),
panel.grid.major = element_line(colour = "grey"))
# add bg color
drawEEMgg(applejuice, 1, has_legend = FALSE) + geom_raster(aes(fill = value)) +
geom_contour(colour = "white")
## End(Not run)
```

EEM

EEM: A package for reading and preprocessing fluorescence excitation-emission matrix

Description

EEM package can be used to import raw data files, visualizing data and preparing them for multivariate analysis

Details

The latest version and documentation can be found here.

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EEM-misc

Internal functions for EEM package

Description

Internal functions for EEM package

Usage

```
generatePoint(n, pch = NULL)
generateColor(n, color.palette = NULL)
getEX(string, digits = NULL)
getEM(string, digits = NULL)
```

Arguments

n	number
pch	Either an integer specifying a symbol or a single character to be used as the default in plotting points.
color.palette	(optional) contour color palette. See palette for more details
string	string or vector of strings
digits	integer indicating the number of decimal places (round) or significant digits (signif) to be used. Negative values are allowed (see 'Details').

Details

'generatePoint' and 'generateColor' are used to create point and color vector from specified number (n) and palette.

Functions

```
• generateColor: generate colors
```

• getEX: get EX value

• getEM: get EM value

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extract

Extract values from other models

Description

Extract values from other models

Usage

```
getLoading(x)
getReg(x)
```

Arguments

Х

output variable from prcomp or plsr functions

Value

A 'EEMweight' list containing title and value attributes.

Examples

```
data(applejuice)
applejuice_uf <- unfold(applejuice) # unfold list into matrix
result <- prcomp(applejuice_uf)
loading <- getLoading(result)
str(loading)</pre>
```

findLocalMax

Find local maximum peaks

Description

Find local maximum peaks in EEM data

Usage

```
findLocalMax(data, ...)
## S3 method for class 'EEM'
findLocalMax(data, n, threshold = 0.7, showprint = TRUE, ...)
## S3 method for class 'matrix'
findLocalMax(data, n, threshold = 0.7, showprint = TRUE, ...)
```

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```
## S3 method for class 'numeric'
findLocalMax(data, threshold = 0.7, showprint = TRUE, ...)
```

Arguments

data	EEM data generated by readEEM function, unfolded EEM data generated by unfold function or a vector of numeric values which have names in the format of EXEM
	(optional) further arguments passed to other methods
n	sample number. The number should not exceed length (EEM).
threshold	threshold value in between 0 and 1. Lower the value to cover low peaks.
showprint	logical value whether to print out the results or not

Value

return a character vector of peak names. If showprint = TRUE, it will also print a dataframe of indicating the value of local maximum peaks.

Methods (by class)

- EEM: for EEM data created by readEEM function
- matrix: for unfolded EEM data created by unfold function
- numeric: for a vector of numeric values which have names in the format of EX...EM...

Examples

```
data(applejuice)
findLocalMax(applejuice, 1)
applejuice_uf <- unfold(applejuice)
findLocalMax(applejuice_uf, 1)</pre>
```

fold

Fold EEM matrix into a list

Description

Fold EEM matrix into a list

gluten 13

Usage

```
fold(EEM_uf, ...)
## S3 method for class 'matrix'
fold(EEM_uf, ...)
## S3 method for class 'data.frame'
fold(EEM_uf, name = NULL, ...)
## S3 method for class 'numeric'
fold(EEM_uf, ...)
```

Arguments

Unfolded EEM matrix where columns are wavelength condition and rows are

samples. It should have corresponding column names (formatted as EX###EM###)

and row names.

... arguments for other methods

name optional for data.frame input to specify the sample names

Value

EEM a list containing EEM/EEM data

Methods (by class)

• data.frame: fold unfolded data.frame

Examples

```
data(applejuice)
applejuice_uf <- unfold(applejuice) # unfold list into matrix
applejuice_uf_norm <- normalize(applejuice_uf) # normalize matrix
drawEEM(fold(applejuice_uf_norm), 1) # visualize normalized EEM</pre>
```

gluten Gluten

Description

Pure wheat gluten and pure wheat starch were mixed at gluten ratios ranging from 0 to 100 %, in 20 % increments. The samples were set in a cell with a quartz glass window, and the samples were pressed against the glass to obtain a flat surface. This dataset contains fluorescence excitation-emission profiles of each samples with 8 replicates. To save space, only the data with gluten ratios ranging from 0 to 60 % was provided.

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Usage

```
data("gluten")
```

References

Kokawa, M., Fujita, K., Sugiyama, J., Tsuta, M., Shibata, M., Araki, T., & Nabetani, H. (2012). Quantification of the distributions of gluten, starch and air bubbles in dough at different mixing stages by fluorescence fingerprint imaging. Journal of Cereal Science, 55(1), 15–21.

Examples

```
data(gluten)
summary(gluten)
```

normalize

Normalize data

Description

Normalize data (area under the curve = 1)

Usage

```
normalize(EEM_uf)
```

Arguments

EEM_uf

Unfolded EEM matrix where columns are wavelength condition and rows are samples

Details

The unfolded EEM data can be normalized by dividing each variable by the sum of the absolute value of all variables in a sample, such that the summation of absolute values of all variables in each sample was equal to 1. This is can be used to reduce the scaling difference, which is common in spectroscopic applications. This difference is usually caused by the scattering effect, source/detector variation and instrumental sensitivity.

Value

A matrix of normalized data

plotLoading 15

Examples

```
data(applejuice)
applejuice_uf <- unfold(applejuice) # unfold list into matrix
applejuice_uf_norm <- normalize(applejuice_uf) # normalize data
rowSums(abs(applejuice_uf_norm), na.rm = TRUE) # the absolute sum of each row equal to 1</pre>
```

plotLoading

Plot loadings for EEM data

Description

Plot loadings for EEM data

Usage

```
plotLoading(x, ncomp = NULL, ...)
```

Arguments

```
x output variable from prcomp or plsr functions
ncomp number of components
... (optional) arguments for drawEEM and filled.contour
```

Value

A figure is returned on the graphic device

```
data(applejuice)
applejuice_uf <- unfold(applejuice) # unfold list into matrix
result <- prcomp(applejuice_uf)
plotLoading(result, ncomp = 1) # plot loading of the first PC</pre>
```

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plotReg

Plot regression coefficients for EEM data

Description

Plot regression coefficients for EEM data

Usage

```
plotReg(x, ncomp = NULL, ...)
```

Arguments

```
    x output variable from plsr function
    ncomp number of components
    ... (optional) arguments for drawEEM and filled.contour
```

Value

A figure is returned on the graphic device

Examples

```
data(gluten)
gluten_uf <- unfold(gluten) # unfold list into matrix

# delete columns with NA values
index <- colSums(is.na(gluten_uf)) == 0
gluten_uf <- gluten_uf[, index]
gluten_ratio <- as.numeric(names(gluten))

require(pls)
model <- plsr(gluten_ratio ~ gluten_uf, ncomp = 3)
plotReg(model)</pre>
```

plotScore

Plot score for prcomp result

Description

Plot score for prcomp (PCA) result

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Usage

```
plotScore(prcompResult, xPC = 1, yPC = 2, group = NULL, group2 = NULL,
  cex = 1.5, cex.legend = 1, label = NULL, pos = 4, col = NULL,
  pch = NULL, legendlocation = "bottomright", legendoutside = FALSE,
  rightwhitespace = 0, ...)
```

Arguments

prcompResult

xPC an integer indicating PC component on x-axis

yPC an integer indicating PC component on y-axis

output object from prcomp function

group a vector of numeric, character or factor class separating the samples into groups.

Correspond to point color.

group2 The second group, can be a vector of numeric, character or factor class separat-

ing the samples into groups. Correspond to point shape.

cex (optional) size of points on graphs
cex.legend (optional) size of fonts in legend

label (optional) a character vector or expression specifying the text to be written.

pos (optional, applicable when label is given) a position specifier for the text. If

specified this overrides any adj value given. Values of 1, 2, 3 and 4, respectively indicate positions below, to the left of, above and to the right of the specified

coordinates.

col point color palette
pch point type palette

legendlocation (optional)location of legend on graph. Look up legend for more details.

legendoutside (optional) set to TRUE if you want to put legend on the outside of the plot. The

legend location is defaulted to topright.

rightwhitespace

(optional) set width for white space for legend. Only applicable if legendoutside

= TRUE

... additional arguments for par

Value

A figure is returned on the graphic device

See Also

plotScorem

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Examples

```
data(applejuice)
applejuice_uf <- unfold(applejuice) # unfold list into matrix</pre>
result <- prcomp(applejuice_uf)</pre>
plotScore(result) # plot PC1 vs PC2 score
plotScore(result, pch = 3, col = "blue") # change shape and color
# get country of apple production
country <- sapply(strsplit(names(applejuice), split = "-"), "[", 1)</pre>
plotScore(result, label = country) # add label
# or plot by group
plotScore(result, xPC = 1, yPC = 3, group = country)
# custom point types and color
plotScore(result, xPC = 1, yPC = 3, group = country, pch = c(1,2), col = c("green", "black"))
# move legend outside
plotScore(result, xPC = 1, yPC = 3, group = country, legendoutside = TRUE)
# two groups
cultivar <- sapply(strsplit(names(applejuice), split = "-"), "[", 2)</pre>
plotScore(result, group = country, group2 = cultivar)
# make the points more transparent
## Not run:
require(scales)
plotScore(result, group = country, group2 = country, col = alpha(generateColor(2), 0.7))
## End(Not run)
```

plotScorem

Plot score matrix for prcomp result based on group

Description

Plot score matrix for prcomp (PCA) result based on group

Usage

```
plotScorem(prcompResult, ncomp = 4, group, cex = 1.5, col = NULL,
    pch = NULL, legendtitle = NULL, ...)
```

Arguments

prcompResult output object from prcomp function

ncomp maximum number of PC score to plot
group a vector of numeric, character or factor class separating the samples into groups.

prcompname 19

cex (optional) size of points on graphs

col point color palette
pch point type palette
legendtitle legend title

... additional arguments to be passed on to pairs

Value

A figure is returned on the graphic device

See Also

```
pairs, plotScore
```

Examples

```
data(applejuice)
# country of apple production
country <- sapply(strsplit(names(applejuice), split = "-"), "[", 1)

applejuice_uf <- unfold(applejuice) # unfold list into matrix
result <- prcomp(applejuice_uf)
# plot PC1 vs PC3 score based on country of production
plotScorem(result, ncomp = 4, group = country)
# specify colours
plotScorem(result, ncomp = 4, group = country, col = c("black", "grey"))</pre>
```

prcompname

Create name for prcomp result

Description

Create name for prcomp result

Usage

```
prcompname(prcompResult, PC, explvar = TRUE)
```

Arguments

prcompResult output value from prcomp function

PC PC number

explvar (logical) show explained variance (%) or not

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Value

String

Examples

```
data(applejuice)
applejuice_uf <- unfold(applejuice) # unfold list into matrix
result <- prcomp(applejuice_uf)
prcompname(result, 1)</pre>
```

print.EEM

Print EEM

Description

Print EEM

Usage

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

Arguments

x EEM class object
... arguments for print function

Examples

```
data(applejuice)
print(applejuice)
```

readEEM

Read raw files and return a list

Description

Read raw files from fluorescence spectrometer

Usage

```
readEEM(path = NULL)
```

summary.EEM 21

Arguments

path

path to the files or folders which contains raw files (accept a vector).

Details

The supported format is *.txt, *.csv and *.dat files from FP-8500 (JASCO), F-7000 (Hitachi Hitech), RF-6000 (Shimadzu) and Aqualog (Horiba) fluorescence spectrometer. It is likely that outputs from different machines of the same companies are supported by this function. Please send a word or pull request to add support for other formats.

Value

readEEM returns a list containing each raw files

summary.EEM

SummarizeEEM EEM list

Description

Summarize by listing the sample number, names and their dimensions

Usage

```
## S3 method for class 'EEM'
summary(object, ...)
```

Arguments

object a list containing EEM data as created by readEEM function.

... arguments for summary function

Value

Text on console

```
data(applejuice)
summary(applejuice)
```

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unfold

Unfold EEM list into a matrix

Description

Unfold EEM list into a matrix with columns as variables (wavelength conditions) and rows as samples.

Usage

```
unfold(EEM, replaceNA = TRUE)
```

Arguments

EEM a list containing EEM data as created by readEEM function.

replaceNA logical value whether to replace NA with 0

Value

Unfolded EEM matrix where columns are wavelength condition and rows are samples

Examples

```
data(applejuice)
applejuice_uf <- unfold(applejuice) # unfold list into matrix
dim(applejuice_uf) # dimension of unfolded matrix</pre>
```

[.EEM

Subset EEM list

Description

Subset EEM list

Usage

```
## S3 method for class 'EEM' x[i, ...]
```

Arguments

```
x EEM class object
```

i indices specifying elements to extract

... arguments for subset function

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```
data(applejuice)
selected <- applejuice[1-5]</pre>
```

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