# Package 'staRdom'

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Type Package

Title PARAFAC Analysis of EEMs from DOM

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- **Depends** R (>= 4.3), ggplot2 (>= 3.4.2), eemR (>= 1.0.1), parallel (>= 4.3)
- Description This is a user-friendly way to run a parallel factor (PARAFAC) analysis (Harshman, 1971) <doi:10.1121/1.1977523> on excitation emission matrix (EEM) data from dissolved organic matter (DOM) samples (Murphy et al., 2013) <doi:10.1039/c3ay41160e>. The analysis includes profound methods for model validation. Some additional functions allow the calculation of absorbance slope parameters and create beautiful plots.

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### LazyData true

```
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```

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# **R** topics documented:

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.eem\_csv

Import EEMs from generic csv files.

# Description

Import EEMs from generic csv files.

# Usage

.eem\_csv(file, col = "ex")

# Arguments

file	path to file
col	either "ex" or "em", whatever wavelength is arranged in columns

# Value

list with EEM data

.trans\_parafac

# Description

Add data of a PARAFAC model derived from multiway from EEMs

# Usage

.trans\_parafac(parafac, em, ex, samples, comp, const, norm\_factors)

### Arguments

parafac	parafac model
em	emission wavelengths
ex	excitation wavelengths
samples	sample names
comp	number of components
const	constraints
norm_factors	factors to invert normalisation

#### Value

parafac model

absorbance\_read *Reading absorbance data from txt and csv files.* 

# Description

Reading absorbance data from txt and csv files.

# Usage

```
absorbance_read(
  absorbance_path,
  order = TRUE,
  recursive = TRUE,
  dec = NULL,
  sep = NULL,
  verbose = FALSE,
  cores = parallel::detectCores(logical = FALSE),
  ...
)
```

#### Arguments

absorbance_path		
	directory containing absorbance data files or path to single file. See details for format of absorbance data.	
order	logical, data is ordered according to wavelength	
recursive	read files recursive, include subfolders	
dec	optional, either you set a decimal separator or the table is tested for . and ,	
sep	optional, either you set a field separator or it is tried to be determined automati- cally	
verbose	logical, provide more information	
cores	number of CPU cores to be used simultanuously	
	additional arguments that are passed on to fread.	

# Details

If absorbance\_path is a directory, contained files that end on "csv" or "txt" are passed on to read.table. If the path is a file, this file is read. Tables can either contain data from one sample or from several samples in columns. The first column is considered the wavelength column. A multi-sample file must have sample names as column names. All tables are combined to one with one wavelength column and one column for each sample containing the absorbance data. Column and decimal separators are guessed from the supplied data. In some cases, this can lead to strange results. Plaese set 'sep' and 'dec' manually if you encounter any problems.

# Value

A data frame containing absorbance data. An attribute "location" contains the filenames where each sample was taken from.

#### See Also

fread

#### Examples

```
absorbance_path <- system.file("extdata", "absorbance", package = "staRdom")
absorbance <- absorbance_read(absorbance_path, verbose = TRUE, cores = 2)</pre>
```

abs\_blcor

Baseline correction for absorbance data

#### Description

Baseline correction for absorbance data

# abs\_fit\_slope

# Usage

abs\_blcor(abs\_data, wlrange = c(680, 700))

# Arguments

abs_data	data.frame containing samples in columns, the column containing wavelengths must be named "wavelength"
wlrange	range of wavelengths that should be used for correction, absorbance mean in that range is subtracted from each value (sample-wise)

# Value

data.frame

# Examples

data(absorbance)
abs\_data\_cor <- abs\_blcor(absorbance)</pre>

abs\_data\_cor1 <- abs\_blcor(absorbance[1:2])</pre>

abs_fit_slope	Fit absorbance data to exponential curve.	drm is used for the fitting
	process.	

# Description

Fit absorbance data to exponential curve. drm is used for the fitting process.

# Usage

```
abs_fit_slope(
  wl,
  abs,
  lim,
  l_ref = 350,
  control = drmc(errorm = FALSE, noMessage = TRUE),
  ...
)
```

#### Arguments

wl	vector containing wavelengths
abs	vector containing absorption in m^-1
lim	vector containing lower and upper limits for wavelengths to use

#### abs\_parms

#### Value

numeric exponential slope coefficient

### See Also

drm

# Examples

```
data(absorbance)
abs_fit_slope(absorbance$wavelength,absorbance$sample1,lim=c(350,400),l_ref=350)
```

abs_parms	Calculating slopes and slope ratios of a data frame of absorbance
	data.

# Description

Calculating slopes and slope ratios of a data frame of absorbance data.

### Usage

```
abs_parms(
  abs_data,
  cuvle = NULL,
  unit = c("absorbance", "absorption"),
  add_as = NULL,
  limits = list(c(275, 295), c(350, 400), c(300, 700)),
  l_ref = list(275, 350, 300),
  S = TRUE,
  lref = FALSE,
  p = FALSE,
 model = FALSE,
  Sint = FALSE,
  interval = 21,
  r2threshold = 0.8,
 cores = parallel::detectCores(logical = FALSE),
  verbose = FALSE
)
```

#### abs\_parms

#### Arguments

abs_data	data frame containing absorbance data.
cuvle	cuvette (path) length in cm, ignored if unit is absorption
unit	unit of absorbance data: if "absorbance", absorbance data is multiplied by log(10) = 2.303 for slope calculations
add_as	additionally to a254 and a300, absorbance at certain wavelengths can be added to the table
limits	list with vectors containig upper and lower bounds of wavelengeth ranges to be fitted
l_ref	list with reference wavelengths, same length as limits
S	logical, include slope indices in the table
lref	logical, include reference wavelength in the table
р	logical, include ps of the coefficients in the table
model	logical, include complete model in data frame
Sint	logical, wether the spectral curve is calculated interval-wise (cdom_spectral_curve)
interval	passed on to cdom_spectral_curve
r2threshold	passed on to cdom_spectral_curve
cores	number of cores to be used for parallel processing
verbose	logical, additional information is provided

# Details

The absorbance data is a data frame with the first column called "wavelength" containg the wavelength. Each other column contains the data from one sample. You can use absorbance\_read to read in appropriate data.

The following spectral parameters are calculated:

- \$S\_275-295\$ slope between 275 and 295 nm calculated with nonlinear regression
- \$S\_350-400\$ slope between 350 and 400 nm calculated with nonlinear regression
- \$S\_300-700\$ slope between 275 and 295 nm calculated with nonlinear regression
- SR slope ratio, calculated by \$S\_275-295\$/\$S\_350-400\$
- E2:E3 ratio \$a\_250\$/\$a\_365\$
- E4:E6 ratio \$a\_465\$/\$a\_665\$
- \$a\_254\$ absorbance at 254 nm
- \$a\_300\$ absorbance at 300 nm

Depending on available wavelength range, values might be NA. Additionally other wavelength limits can be defined. The slope ratio might fail in this case. For further details please refer to Helm et al. (2008).

### Value

A data frame containing the adsorption slopes and slope ratios in column, one line for each sample.

#### References

Helms, J., Kieber, D., Mopper, K. 2008. Absorption spectral slopes and slope ratios as indicators of molecular weight, source, and photobleaching of chromophoric dissolved organic matter. Limnol. Oceanogr., 53(3), 955–969 https://aslopubs.onlinelibrary.wiley.com/doi/10.4319/lo. 2008.53.3.0955

#### Examples

data(absorbance)

```
a1 <- abs_parms(absorbance, cuvle = 5, verbose = TRUE, cores = 2)
a2 <- abs_parms(absorbance, cuvle = 5,1_ref=list(NA,NA,NA), lref=TRUE, cores = 2) # fit lref as well</pre>
```

as.data.frame.eem Converting EEM data from class eem to data.frame.

#### Description

Converting EEM data from class eem to data.frame.

### Usage

```
## S3 method for class 'eem'
as.data.frame(x, row.names = NULL, optional = FALSE, gather = TRUE, ...)
```

#### Arguments

Х	abc
row.names	abc
optional	ignored
gather	logical, says whether data.frame is returned with excitation wavelength as col- umn names or as values of a column. If the data is gathered, the sample name is added as value in a calumn
	ignored

#### Value

A data frame containing the EEM data.

# Examples

```
data(eem_list)
```

as.data.frame(eem\_list[[1]])
as.data.frame(eem\_list[[1]],gather=FALSE)

A\_missing

### Description

Samples from an eemlist that were not used in the modelling process are added as entries in the A-modes. Values are calculated using fixed B and C modes in the PARAFAC algorithm. B and C modes can be provided via a previously calculated model or as matrices manually.

#### Usage

```
A_missing(
    eem_list,
    pfmodel = NULL,
    cores = parallel::detectCores(logical = FALSE),
    components = NULL,
    const = NULL,
    control = NULL,
    ...
)
```

#### Arguments

eem_list	object of class eemlist with sample data
pfmodel	object of class parafac
cores	number of cores to use for parallel processing
components	optionally supply components to use manually, either as a variable of class parafac_components or as a list of variables of class parafac_components, if you do so,
const	optional constraints for model, just used, when components are supplied
control	optional constraint control parameters for model, just used, when components are supplied
	additional arguments passed to eem_parafac

# Details

This function can be used to calculate A modes (sample loadings) for samples that were previously excluded from the modelling process (e.g. outliers). Another way to use it would be a recombination of components from different models and calculating the according sample loadings. Expecially the later application is experimental and results have to be seen critically! Nevertheless, I decided to supply this function to stimulate some experiments on that and would be interested in your findings and feedback.

# Value

object of class parafac

# Examples

data(eem\_list) data(pf\_models)

```
A_missing(eem_list, pf4[[1]], cores = 2)
```

eem2array

Data from an eemlist is transformed into an array

# Description

Data matrices from EEM are combined to an array that is needed for a PARAFAC analysis.

# Usage

eem2array(eem\_list)

# Arguments

eem\_list object of class eemlist

# Value

object of class array

# Examples

data(eem\_list)

X <- eem2array(eem\_list)</pre>

eempf4analysis

Create table of PARAFAC components and (optionally) EEM peaks and indices as well as absorbance slope parameters.

# Description

Please refer to eem\_biological\_index, eem\_coble\_peaks, eem\_fluorescence\_index, eem\_biological\_index and abs\_parms for details on the certain values

#### Usage

```
eempf4analysis(
   pfmodel,
   eem_list = NULL,
   absorbance = NULL,
   cuvl = NULL,
   n = 4,
   export = NULL,
   cores = parallel::detectCores(logical = FALSE),
   ...
)
```

# Arguments

pfmodel	PARAFAC model where loadings of the components are extracted
eem_list	optional eemlist used for peak and indices calculation
absorbance	optional absorbance table used for absorbance slope parameter calculation
cuvl	optional cuvette length of absorbance data in cm
n	optional size of moving window in nm for data smoothing in advance of peak picking
export	optional file path of csv or txt table where data is exported
cores	number of parallel calculations (e.g. number of physical cores in CPU)
	additional parameters passed to write.table

#### Value

data frame

# Examples

```
cuvl = 5, n = 4, cores = 2)
```

eempf\_bindxc Combining extracted components of PARAFAC models

# Description

Combining extracted components of PARAFAC models

### Usage

```
eempf_bindxc(components)
```

# Arguments

components list of parafac\_components

### Value

parafac\_components

# Examples

```
data(pf_models)
pfmodel <- pf4[[1]]
comps <- eempf_excomp(pfmodel,c(1,3))
comps2 <- eempf_excomp(pfmodel,c(4,6))
comps3 <- eempf_bindxc(list(comps, comps2))</pre>
```

eempf\_compare

Plot a set of PARAFAC models to compare the single components

# Description

Three plots are returned:

- 1. plot of numer of components vs. model fit
- 2. plot of different components as colour maps
- 3. plot of different components as peak lines

The plots are intended to help with a suitable number of components.

# eempf\_comps3D

# Usage

eempf\_compare(pfres, ...)

# Arguments

pfres	list of several objects of class parafac
	arguments passed on to <code>eempf_fits</code> and <code>eempf_plot_comps</code>

# Value

3 objects of class ggplot

### See Also

eempf\_fits, eempf\_plot\_comps

# Examples

```
data(pf_models)
```

eempf\_compare(pf4)

eempf\_comps3D 3L

# 3D plots of PARAFAC components

# Description

Interactive 3D plots are created using plotly.

# Usage

```
eempf_comps3D(pfmodel, which = NULL)
```

# Arguments

pfmodel	object of class parafac
which	optional, if numeric selects certain component

# Value

plotly plot

# Examples

```
## Not run:
data(pf_models)
eempf_comps3D(pf4[[1]])
## End(Not run)
```

eempf\_comp\_load\_plot Plot components from a PARAFAC model

# Description

Additionally a bar plot with the amounts of each component in each sample is produced.

# Usage

```
eempf_comp_load_plot(pfmodel, ...)
```

# Arguments

pfmodel	object of class parafac
	attributes passe don to ggeem

# Value

ggplot

# See Also

ggeem, eempf\_load\_plot

# Examples

```
data(pf_models)
```

eempf\_comp\_load\_plot(pf4[[1]])

 $\texttt{eempf\_comp\_mat}$ 

### Description

The components of a PARAFAC analysis are extracted as a data frame

#### Usage

```
eempf_comp_mat(pfmodel, gather = TRUE)
```

# Arguments

pfmodel	object of class parafac
gather	logical value whether excitation wavelengths are a column, otherwise excitation wavelengths are column names

# Value

a list of class data frames

### Examples

```
data(pf_models)
```

eempf\_comp\_mat(pf4[[1]])

eempf\_comp\_names Extract names from PARAFAC model components

# Description

Extract names from PARAFAC model components

#### Usage

eempf\_comp\_names(pfmodel)

# Arguments

pfmodel parafac model

### Value

vector of names or list of vecters of names

# Examples

```
data(pf_models)
eempf_comp_names(pf4)
eempf_comp_names(pf4) <- c("A","B","C","D","E","F","G")
value <- list(c("A1","B1","C1","D","E","F","G"),
c("A2","B2","C","D","E","F","G"),
c("A3","B3","C","D","E","F","G"),
c("A4","B4","C","D","E","F","G"),
c("A5","B5","C","D","E","F","G5")
)
eempf_comp_names(pf4) <- value
eempf_comp_names(pf4)
ggeem(pf4[[1]])
```

eempf\_comp\_names<- Set names of PARAFAC components

# Description

Set names of PARAFAC components

# Usage

```
eempf_comp_names(pfmodel) <- value</pre>
```

#### Arguments

pfmodel	model of class parafac
value	character vector containing the new names for the components

#### Value

parafac model

# Examples

data(pf\_models)

eempf\_comp\_names(pf4) <- c("A", "B", "C", "D", "E", "F", "G")</pre>

eempf\_convergence *Extract modelling information from a PARAFAC model.* 

# Description

The convergence behaviour of all initialisations in a PARAFAC model is shown by printing the numbers

# Usage

```
eempf_convergence(pfmodel, print = TRUE)
```

# Arguments

pfmodel	PARAFAC model created with staRdom using output = "all"
print	logical, whether you want console output or just a list with results

### Value

list with numbers of converging models, cflags and SSEs

#### Examples

```
data("pf_models")
pfmodel <- pf4[[1]]
conv_beh <- eempf_convergence(pfmodel)</pre>
```

eempf\_corcondia Calculate the core consistancy of an EEM PARAFAC model

#### Description

This is basically a wrapper for corcondia that deals with the normalisation of the original data., Other than corcondia, the default dicisor = "core".

### Usage

```
eempf_corcondia(pfmodel, eem_list, divisor = "core")
```

# Arguments

pfmodel	PARAFAC model
eem_list	eemlist
divisor	divisor, please refer to corcondia

#### Value

numeric

# Examples

```
## Not run:
# due to data limitation in package, example does not work with that data!
```

```
# eempf_corcondia(pfmodel,eem_list)
```

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

eempf\_corplot Plot correlations of components in samples

# Description

A pair plot showing correlations between samples is created.

# Usage

```
eempf_corplot(
   pfmodel,
   normalisation = FALSE,
   lower = list(continuous = "smooth"),
   mapping = aes(alpha = 0.2),
   ...
)
```

#### Arguments

pfmodel	object of class parafac
normalisation	logical, whether normalisation is undone or not
lower	style of lower plots, see ggpairs
mapping	aesthetic mapping, see ggpairs
	passed on to ggpairs

# Value

object of class ggplot

### See Also

ggpairs

# eempf\_cortable

# Examples

```
data(pf_models)
eempf_corplot(pf4[[1]])
```

eempf\_cortable

Calculating correlations between the component loadings in all samples (C-Modes).

# Description

Calculating correlations between the component loadings in all samples (C-Modes).

# Usage

eempf\_cortable(pfmodel, normalisation = FALSE, method = "pearson", ...)

# Arguments

pfmodel	results from a PARAFAC analysis, class parafac
normalisation	logical, whether normalisation is undone or not
method	method of correlation, passed to cor
	passed on to cor

# Value

matrix

# Examples

```
data(pf_models)
eempf_cortable(pf4[[1]])
```

 ${\tt eempf\_eemqual}$ 

# Description

Calculating EEMqual which is an indicator of a PARAFAC model's quality

#### Usage

```
eempf_eemqual(pfmodel, eem_list, splithalf = NULL, ...)
```

### Arguments

pfmodel	PARAFAC model
eem_list	EEM data as eemlist
splithalf	optionally, you can supplie available splithalf results from model to decrease computation time
	additional arguments passed to splithalf

# Value

data frame containing fit, corcondia, product of best TCCs from splithalf analysis, eemqual and splithalf models

# References

Rasmus Bro, Maider Vidal, EEMizer: Automated modeling of fluorescence EEM data, Chemometrics and Intelligent Laboratory Systems, Volume 106, Issue 1, 2011, Pages 86-92, ISSN 0169-7439

# Examples

```
# data(eem_list)
# data(pf_models)
# pfmodel <- pf4[[1]]
# eempf_eemqual(eem_list,pfmodel) # insuficient example data to run!</pre>
```

eempf\_excomp

# Description

Extracting components of a PARAFAC model

# Usage

```
eempf_excomp(pfmodel, comps)
```

# Arguments

pfmodel	parafac model
comps	vector with numbers of components to extract

# Value

list

# Examples

data(pf\_models)
pfmodel <- pf4[[1]]
comps <- eempf\_excomp(pfmodel,c(1,3))</pre>

eempf_export	Create one table containing the PARAFAC models factors and option-
	ally exporting it to csv or txt

# Description

Create one table containing the PARAFAC models factors and optionally exporting it to csv or txt

# Usage

```
eempf_export(pfmodel, export = NULL, Fmax = TRUE, ...)
```

# Arguments

pfmodel	PARAFAC model
export	file path to export table
Fmax	rescale modes so the A mode shows the maximum fluorescence
	additional parameters passed to write.table

# Value

data frame

# Examples

```
data(pf_models)
```

```
factor_table <- eempf_export(pf4[[1]])</pre>
```

eempf\_fits

# Fits vs. components of PARAFAC models are plotted

# Description

Fits vs. components of PARAFAC models are plotted

# Usage

eempf\_fits(pfres, ...)

# Arguments

pfres	list of objects of class parafac
	arguments passed on to ggplot

# Value

object of class ggplot

# Examples

data(pf\_models)

eempf\_fits(pf4)

eempf\_leverage

Calculate the leverage of each emission and excitation wavelength and each sample from a single PARAFAC model

### Description

Calculate the leverage of each emission and excitation wavelength and each sample from a single PARAFAC model

# Usage

```
eempf_leverage(pfmodel)
```

# Arguments

pfmodel object of class parafac

### Value

list of 3 named vectors (emission, excitation wavelengths and samples)

#### Examples

data(pf\_models)

```
eempf_leverage(pf4[[1]])
```

eempf\_leverage\_data Combine leverages into one data frame and add optional labels.

# Description

Combine leverages into one data frame and add optional labels.

#### Usage

```
eempf_leverage_data(cpl, qlabel = 0.1)
```

#### Arguments

cpl	leverage, outpout from eempf_leverage
qlabel	optional, quantile of which labels are shown $(1 = all, 0 = no labels)$

# Value

data frame

### Examples

```
data(pf_models)
```

```
leverage <- eempf_leverage(pf4[[1]])
lev_data <- eempf_leverage_data(leverage)</pre>
```

eempf\_leverage\_ident Plot leverage of emission wavelengths, excitation wavelengths and samples.

# Description

Plot is interactive where you can select values with your mouse. A list of vectors is returned to remove this outliers in a further step from your samples. The labels to be shown can be selected by adding the quatile of samples with highest leverages to be labeled.

# Usage

```
eempf_leverage_ident(cpl, qlabel = 0.1)
```

#### Arguments

cpl	leverage, outpout from eempf_leverage
qlabel	optional, quantile of which labels are shown $(1 = all, 0 = no labels)$

#### Value

list of three vectors containing the names of selected samples

#### See Also

eempf\_leverage\_plot

### Examples

```
data(pf_models)
```

leverage <- eempf\_leverage(pf4[[1]])
outliers <- eempf\_leverage\_ident(leverage)</pre>

eempf\_leverage\_plot Plot leverage of emission wavelengths, excitation wavelengths and samples.

# Description

The labels to be shown can be selected by adding the quatile of samples with highest leverages to be labeled.

# Usage

```
eempf_leverage_plot(cpl, qlabel = 0.1)
```

# Arguments

cpl	leverage, outpout from eempf_leverage
qlabel	optional, quantile of which labels are shown $(1 = all, 0 = no labels)$

#### Value

ggplot

#### See Also

eempf\_leverage\_ident

# Examples

data(pf\_models)

leverage <- eempf\_leverage(pf4[[1]])
eempf\_leverage\_plot(leverage)</pre>

eempf\_load\_plot Plot amount of each component in each sample as bar plot

### Description

Plot amount of each component in each sample as bar plot

# Usage

```
eempf_load_plot(pfmodel)
```

# Arguments

pfmodel parafac model

#### Value

ggplot

# Examples

data(pf\_models)

eempf\_load\_plot(pf4[[1]])

eempf\_mleverage

Calculate the leverage of each emission and excitation wavelength and each sample from a list of PARAFAC models

# Description

Calculate the leverage of each emission and excitation wavelength and each sample from a list of PARAFAC models

# Usage

```
eempf_mleverage(pfres_comps, ecdf = FALSE, stats = FALSE)
```

# Arguments

pfres_comps	object of class parafac
ecdf	logical, transforme leverages to according empirical quantiles (ecdf)
stats	logical, whether means and standard deviations are calculated from leverages

# Value

data frame containing leverages of wavelengths and samples for each model

# Examples

data(pf\_models)

eempf\_mleverage(pf3)

eempf\_OF\_upload

# Description

This function uploads a PARAFAC model to openfluor.org from within R. You need to have an account at openfluor.org and supply the email used for the account to the function. Your password is then asked in a secure way and only used within one execution of this function.

#### Usage

eempf\_OF\_upload(email, file)

#### Arguments

email	email address you use to login at openfluor.org as string
file	the file containing a PARAFAC model in openfluor format

### Value

HTTP status code from the upload POST

### Examples

```
## due to the need of a valid account, this function cannot be
## tested with generic data.
## Please use your own account to do so.
## Not run:
data(pf_models)
file <- file.path(tempdir(),"openfluor_example.txt")
eempf_openfluor(pf4[[1]],file)
eempf_OF_upload("helena.glory@rur.play", file)</pre>
```

## End(Not run)

eempf\_openfluor Write out PARAFAC components to submit to openfluor.org.

#### Description

openfluor.org offers the possibility to compare your results to others, that were uploaded to the database. This functions writes out a txt containing the header lines and your components. Please open the file in an editor and fill in further information that cannot be covered by this function.

# Usage

```
eempf_openfluor(
   pfmodel,
   file,
   Fmax = TRUE,
   upload = FALSE,
   email = NULL,
   model_details = list()
)
```

#### Arguments

pfmodel	PARAFAC model
file	string, path to outputfile. The directory must exist, the file will be created or overwritten if already present.
Fmax	rescale modes so the A mode shows the maximum fluorescence. As openfluor does not accept values above 1, this is a way of scaling the B and C modes to a range between 0 and 1.
upload	logical, whether model is directly uploaded to openfluor.org
email	optional email address to log into openfluor.org
<pre>model_details</pre>	optional named list with strings to be added in the openfluor file in the fields corresponding to the list names

# Value

txt file

# Examples

```
data(pf_models)
model_details <- list(name = "River", creator = "Helena Glory",
constraints = "non-negative", validation = "split-half", unit= "RU")
eempf_openfluor(pf4[[1]],file.path(tempdir(),"openfluor_example.txt"),
upload = FALSE, model_details = model_details)</pre>
```

eempf\_plot\_comps Plot all components of PARAFAC models

# Description

The components can be plottet in two ways: either as a colour map or as two lines (emission, excitation wavelengths) intersecting at the component maximum. If the list of provided models is named, these names are shown in the plot. Otherwise, the models are automatically named by "model#".

eempf\_plot\_ssccheck

# Usage

```
eempf_plot_comps(
   pfres,
   type = 1,
   names = TRUE,
   contour = FALSE,
   colpal = "default",
   ...
)
```

# Arguments

pfres	list of PARAFAC models
type	1 for a colour map and 2 for em and ex wavelength loadings
names	logical, whether names of components should be written into the plot
contour	in case of 3 dimensional component plots, contours are added
colpal	"default" to use the viridis colour palette, "rainbow" to use a subset of the rain- bow palette, any custom vector of colors or a colour palette. A gradient will be produced from this vector. Larger vectors (e.g. 50 elements) can produce smoother gradients.
	arguments passed on to other functions, e.g.

# Value

object of class ggplot

# Examples

```
data(pf_models)
eempf_plot_comps(pf4, type = 1)
# use a different colour scheme:
# eempf_plot_comps(pf4, type = 1, colpal = heat.colors(50))
eempf_plot_comps(pf4, type = 2)
eempf_plot_comps(list(pf4[[1]],pf4[[1]]), type=1)
```

# Description

Plot results from an SSC check

# Usage

eempf\_plot\_ssccheck(ssccheck)

# Arguments

ssccheck outpout from eempf\_ssccheck

# Value

ggplot element

# Examples

data(pf\_models)

```
ssccheck <- eempf_ssccheck(pfmodels = pf3[1:3], cores = 2)
eempf_plot_ssccheck(ssccheck)</pre>
```

eempf\_reorder

# Reorder PARAFAC components

# Description

Reorder PARAFAC components

# Usage

eempf\_reorder(pfmodel, order, decreasing = FALSE)

# Arguments

pfmodel	model of class parafac
order	vector containing desired new order or "em" or "ex" to reorder according to emission or excitation wavelengths of the peaks
decreasing	logical, whether components are reordered according to peak wvalengths in a decreasing direction

### Value

parafac model

# eempf\_report

# Examples

```
data(pf_models)
ggeem(pf4[[1]])
pf4r <- eempf_reorder(pf4[[1]], "ex")</pre>
```

ggeem(pf4r)

```
eempf_report
```

Create a html report of a PARAFAC analysis

# Description

Create a html report of a PARAFAC analysis

# Usage

```
eempf_report(
    pfmodel,
    export,
    eem_list = NULL,
    absorbance = NULL,
    meta = NULL,
    metacolumns = NULL,
    splithalf = FALSE,
    shmodel = NULL,
    performance = FALSE,
    residuals = FALSE,
    spp = 5,
    cores = parallel::detectCores(logical = FALSE),
    ...
)
```

# Arguments

pfmodel	PARAFAC model
export	path to exported html file
eem_list	optional EEM data
absorbance	optional absorbance data
meta	optional meta data table
metacolumns	optional column names of metadata table
splithalf	optional logical, states whether split-half analysis should be included
shmodel	optional results from split-half analysis. If this data is not supplied but EEM data is available the split-half analysis is calculated on the creation of the report. Calculating the split-half analysis takes some time!

performance	calculating model performance: eempf_eemqual
residuals	logical, whether residuals are plotted in the report
spp	plots per page for loadgins and residuals plot
cores	cores to be used for the calculation
	arguments to or from other functions

#### Value

TRUE if report was created

#### Examples

eempf\_rescaleBC Rescale B and C modes of PARAFAC model

#### Description

B and C modes (emission and excitation wavelengths) are rescaled to RMS of value newscale. This is compensated in A mode (sample loadings).

#### Usage

```
eempf_rescaleBC(pfmodel, newscale = "Fmax")
```

# Arguments

pfmodel	object of class parafac
newscale	If (default) newscale = "Fmax", each component will be scaled so the maximum of each component is 1. It is also possible to set a desired root mean-square for each column of the rescaled mode. Can input a scalar or a vector with length
	equal to the number of factors for the given mode.

# Value

object of class parafac

#### See Also

rescale

# Examples

```
data(pf_models)
```

new\_pf <- eempf\_rescaleBC(pf4[[1]])</pre>

eempf\_residuals Calculate residuals of EEM data according to a certain model

# Description

Calculate residuals of EEM data according to a certain model

# Usage

```
eempf_residuals(
   pfmodel,
   eem_list,
   select = NULL,
   cores = parallel::detectCores(logical = FALSE)/2
)
```

# Arguments

pfmodel	PARAFAC model of class parafac
eem_list	eemlist containing EEM data
select	character vector containing the names of the desired samples
cores	number of cores to use for parallel processing

# Value

data frame with EEM residuals

# Examples

```
data(eem_list)
data(pf_models)
residuals <- eempf_residuals(pf4[[1]], eem_list, cores = 2)</pre>
```

eempf\_residuals\_metrics

Calculate residual metrics from a PARAFAC model

# Description

The metrics calculated with this function are:

- RSS: residual sum of squares
- MAE: mean absolute error
- SAE: sum of absolute errors
- RSAE: sum of absolute error in relation to the sum of fluorescence and
- LEV: the leverage as described in eempf\_leverage The example contains a way to plot these numbers.

#### Usage

```
eempf_residuals_metrics(residuals, leverage)
```

#### Arguments

residuals	data.frame as derived from eempf_residuals
leverage	list of data.frames as derived from eempf_leverage

#### Value

a list of data.frames containing residuals metrics for each sample, emission and excitation wavelength

# Examples

```
data(eem_list)
data(pf_models)
residuals <- eempf_residuals(pf4[[1]], eem_list, cores = 2)
leverage <- eempf_leverage(pf4[[1]])
metrics <- eempf_residuals_metrics(residuals, leverage)</pre>
```
```
metrics$sample
```

```
## plot different residual metrics
require(dplyr)
require(tidyr)
require(ggplot2)
lapply(names(metrics), function(name){
 metrics[[name]] %>%
 mutate(mode = name, element = !!sym(name))
}) %>%
 bind_rows() %>%
 pivot_longer(cols = RSS:LEV, names_to = "metric", values_to = "value") %>%
 # uncomment the following line to select certain metrics
 # filter(metric %in% c("RSS","LEV")) %>%
 ggplot(aes(x = element, y = value, colour = metric))+
 geom_point()+
 facet_wrap(mode ~ ., ncol = 3, scales = "free")+
 theme(axis.text.x = element_text(angle = 90))+
 scale_y_continuous(trans="log")
```

<pre>eempf_residuals_plot</pre>	Plot samples by means of whole sample, each single component and
	residuum

## Description

A raster of plots is created. Each column shows one sample. The top n rows show the n components from the model according their occurance in the certain samples. The second last row shows the residual, not covered by any component in the model and the last row shows the whole sample.

#### Usage

```
eempf_residuals_plot(
   pfmodel,
   eem_list,
   res_data = NULL,
   spp = 5,
   select = NULL,
   residuals_only = FALSE,
   cores = parallel::detectCores(logical = FALSE),
   contour = FALSE,
   colpal = "default"
)
```

## Arguments

pfmodel	object of class parafac containing the generated model
eem_list	object of class eemlist with all the samples that should be plotted
res_data	optional, data of sample residuals related to the model, output from ${\tt eempf\_residuals}$
spp	optional, samples per plot
select	optional, character vector of samples you want to plot
residuals_only	plot only residuals
cores	number of cores to use for parallel processing
contour	logical, states whether contours should be plotted
colpal	"default" to use the viridis colour palette, "rainbow" to use a subset of the rain- bow palette, any custom vector of colors or a colour palette. A gradient will be produced from this vector. Larger vectors (e.g. 50 elements) can produce smoother gradients.

### Details

eem\_list may contain samples not used for modelling. Calculation is done by A\_missing. This especially interesting if outliers are excluded prior modelling and should be evaluated again afterwards. Usually, residuals contain negative values, while these is the exception in samples and PARAFAC components. Therefore, we decided to use a similar colour palette as in the other plot functions but adding a purple tone for negative values.

### Value

several ggplot objects

## Examples

```
data(eem_list)
data(pf_models)
eem_list <- eem_extract(eem_list, 1:10)</pre>
eem_list <- eem_rem_scat(eem_list, rep(TRUE, 4), c(15,10,16,12))</pre>
eempf_residuals_plot(pf4[[1]], eem_list, cores = 2)
# use other colour schemes:
# eempf_residuals_plot(pf4[[1]], eem_list, colpal = c("blue",heat.colors(50)))
# plots <- eempf_residuals_plot(pf4[[1]], eem_list)</pre>
# lapply(plots, function(pl){
#
   pl +
#
      scale_fill_viridis_c() +
      scale_colour_viridis_c()
#
# })
```

eempf\_ssc

Calculate the shift-and shape-sensitive congruence (SSC) between model components

### Description

The data variable pf\_models can be supplied as list of PARAFAC models, output from a splithalf analysis or list of matrices Please see details of calculation in: U.J. Wünsch, R. Bro, C.A. Stedmon, P. Wenig, K.R. Murphy, Emerging patterns in the global distribution of dissolved matter fluorescence, Anal. Methods, 11 (2019), pp. 888-893

### Usage

```
eempf_ssc(
   pfmodels,
   tcc = FALSE,
   m = FALSE,
   cores = parallel::detectCores(logical = FALSE)
)
```

#### Arguments

pfmodels	list of either PARAFAC models or component matrices
tcc	if set TRUE, TCC is returned instead
m	logical, if TRUE, emission and excitation SSCs or TCCs are combined by cal- culating the geometric mean
cores	number of CPU cores to be used

## Value

(list of) tables containing SCCs between components

## Examples

```
pf_models <- pf3[1:3]
sscs <- eempf_ssc(pf_models, cores = 2)
sscs
tcc <- eempf_ssc(pf_models, tcc = TRUE, cores = 2)
tcc
## mixed tcc (combine em and ex)
mtcc <- eempf_ssc(pf_models, tcc = TRUE, m = TRUE, cores = 2)
mtcc</pre>
```

```
sh_sscs <- eempf_ssc(sh, cores = 2)
sh_sscs
## view diagonals only (components with similar numbers only)
lapply(sh_sscs, lapply, diag)</pre>
```

eempf\_ssccheck Check SSCs between different models or initialisations of one model

#### Description

Check SSCs between different models or initialisations of one model

### Usage

```
eempf_ssccheck(
  pfmodels,
  best = length(pfmodels),
  tcc = FALSE,
  cores = parallel::detectCores(logical = FALSE)
)
```

## Arguments

pfmodels	list of parafac models
best	number of models with the highest R^2 to be used, default is all models
tcc	logical, if TRUE, TCC instead of SSC is calculated
cores	number of CPU cores to be used

## Value

data.frame containing SSCs

## Examples

```
data(pf_models)
```

eempf\_ssccheck(pf3[1:2], cores = 2)

# SSCs of split-half models, models need to be unlisted data(sh) eempf\_ssccheck(unlist(sh, recursive = FALSE), cores = 2)

eempf\_varimp

## Description

Calculate the importance of each component.

#### Usage

```
eempf_varimp(
   pfmodel,
   eem_list,
   cores = parallel::detectCores(logical = FALSE),
   ...
)
```

#### Arguments

pfmodel	model of class parafac
eem_list	eemlist used to calculate that model
cores	cores to be used for the calculation
	other aruments passed to eem_parafac

## Details

The importance of each variable is calculated by means of creating a model without a specific component and calculating the difference between the original R-squared and the one with the left out component. The derived values state the loss in model fit if one component is not used in the modeling process. For the creation of the new models, the exact components of the original model are used.

## Value

numeric vector, values are in the same order of the components in the supplied model.

### Examples

```
data(pfmodel)
data(eem_list)
eempf_varimp(pf4[[1]], eem_list, cores = 2)
```

eem\_absdil

Multiply absorbance data according to the dilution and remove absorbance from samples where undiluted data is used.

# Description

According to dilution data absorbance is either multiplied by the according factor or the undiluted absorbance data is deleted. You can either specify the cor\_data data table coming from eem\_dilcorr or supply an eemlist, and the dilution data to created on the fly.

### Usage

```
eem_absdil(
   abs_data,
   eem_list = NULL,
   dilution = NULL,
   cor_data = NULL,
   auto = TRUE,
   verbose = FALSE
)
```

## Arguments

abs_data	absorbance data
eem_list	optional eemlist
dilution	optional dilution data as data frame
cor_data	optional output from eem_dilcorr as data frame
auto	optional, see eem_dilcorr
verbose	optional, see eem_dilcorr

# Value

data frame

## Examples

# no appropriate exmaple data available yet

eem\_apply

### Description

Applying functions on EEMs

#### Usage

eem\_apply(data, func, return = c("eemlist", "value"), ...)

#### Arguments

data	eemlist to be modified
func	a function to be applied on the data.
return	either "eemlist" or "value"
	additional arguments passed on to func

## Details

The EEMs are passed on as first argument to func. Additionally, the vector of excitation wavelengths is passed on as ex and the emission wavelengths as em. Therefore, the supplied function has to allow these arguments. The easiest way would be . . . (see example).

#### Value

eemlist or list

## Examples

```
## define a function, that would divide a matrix by its maximum
# more general, if you want to return a valid eemlist (see below),
# a matrix of the same size has to be returned
# ... is used as a placeholder for any argument, important: em and
# ex wavelengths are passed on, so the function needs to take them as arguments,
# even if they are not used
norm_max <- function(x, ...){
    x/max(x)
}
# load example data
data("eem_list")
# normalise eems by the function defined above
norm_eems <- eem_apply(eem_list,norm_max,"eemlist")
# plot the results to see the difference
ggeem(norm_eems)</pre>
```

```
# define another function. what values were used to
# multiply the eems with?
norm_fac <- function(x, ...){</pre>
  1/max(x)
}
# return a list of factors used for normalisation
norm_factors <- eem_apply(eem_list,norm_fac,"value")</pre>
unlist(norm_factors)
# return list of em vectors.
# important: x needs to be in the first position, but
# is not used later!
extr_em <- function(x,em,...){</pre>
  em
}
em_vectors <- eem_apply(eem_list,extr_em,"value")</pre>
em_vectors
```

eem\_checkdata Check your EEM, absorption and metadata before processing

# Description

The function tries to lead you to possible problems in your data.

## Usage

```
eem_checkdata(
    eem_list,
    absorbance,
    metadata = NULL,
    metacolumns = NULL,
    correction = FALSE,
    error = TRUE
)
```

# Arguments

eem_list	eemlist continaing EEM data.
absorbance	data.frame containing absorbance data.
metadata	optional data.frame containing metadata.
metacolumns	character vector of columns that are checkt for complete data sets

correction	logical, whether EEMs should be checked for applied corrections
error	logical, whether a problem should cause an error or not.

#### Details

The returned list contains character vectors with sample names where possible problems were found: problem (logical, whether a severe problem was found), nas (sample names with NAs in EEM data), missing\_correction (correction of EEM samples was not done or not done success-fully),eem\_no\_abs (EEM samples with no absorbance data), abs\_no\_eem (samples with present absorbance but no EEM data), duplse (duplicate sample names in EEM data), duplsa (duplicate sample names in absorbance data), invalid\_eem (invalid EEM sample name), invalid\_abs (invalid absorbance sample name), range\_mismatch (wavelength ranges of EEM and absorbance data are mismatching), metadupls (duplicate sample names in metadata), metamissing (EEM samples where metadata is missing), metaadd (samples in metadata without EEM data)

#### Value

writes out possible porblems to command line, additionally list with sample names where possible problems were found, see details.

#### Examples

```
folder <- system.file("extdata/EEMs", package = "staRdom") # load example data
eem_list <- eem_read(folder, recursive = TRUE, import_function = eem_csv)
abs_folder <- system.file("extdata/absorbance", package = "staRdom") # load example data
absorbance <- absorbance_read(abs_folder, cores = 2)
metatable <- system.file("extdata/metatable_dreem.csv",package = "staRdom")
meta <- read.table(metatable, header = TRUE, sep = ",", dec = ".", row.names = 1)
checked <- eem_checkdata(eem_list, absorbance, metadata = meta,
metacolumns = "dilution", error = FALSE)
# This example returns a message, that absorbance data for the
# blank samples are missing. As absorbance is supposed to be 0 over
# the whole spectrum when you measure blanks, there is no need
# to supply the data and do an inner-filter effect correction.</pre>
```

eem\_checksize Check size of EEMs

#### Description

The size of EEMs in an eemlist is checked and the sample names of samples with more data than the sample with the smallest range are returned.

#### Usage

```
eem_checksize(eem_list)
```

## Arguments

eem\_list eemlist

# Value

character vector

## Examples

```
data(eem_list)
eem_checksize(eem_list)
```

and corrections	Roturn namo	ot campl	ac whare	cortain	corrections	are miccina
eem_corrections	Return names	on summer	es where	cenum		uve mussine.
		- J				

# Description

Return names of samples where certain corrections are missing.

## Usage

```
eem_corrections(eem_list)
```

# Arguments

eem\_list eemlist to be checked

#### Value

prints out sample names

## Examples

data(eem\_list)

eem\_corrections(eem\_list)

eem\_csv

## Description

This function can be used to import generic csv files containing EEM data using eem\_read. Excitation wavelengths are assumed column-wise and emission wavelengths row-wise. If your data is arranged the other way round, please use eem\_csv2

#### Usage

eem\_csv(file)

#### Arguments

file path to file passed from eem\_read

### Value

list with EEM data

#### Examples

```
eems <- system.file("extdata/EEMs",package="staRdom")
eem_list <- eem_read(eems, recursive = TRUE, import_function = eem_csv)</pre>
```

eem\_list

eem\_csv2

Importer function for generic csv files to be used with eem\_read().

# Description

This function can be used to import generic csv files containing EEM data using eem\_read. Excitation wavelengths are assumed row-wise and emission wavelengths column-wise If your data is arranged the other way round, please use eem\_csv

## Usage

eem\_csv2(file)

#### Arguments

file path to file passed from eem\_read

#### Value

list with EEM data

## Examples

```
## no example data provided with the package
## below is an example how this could like like
# eems <- "C:/some/path/to/eem.csv"
# eem_list <- eem_read(eems, recursive = TRUE, import_function = eem_csv2)
# eem_list
```

eem\_dilcorr

Create table how samples should be corrected because of dilution

#### Description

Due to dilution absorbance spectra need to be multiplied by the dilution factor and names of EEM samples can be adjusted to be similar to their undiluted absorbance sample. The table contains information about these two steps. Undiluted samples are suggested by finding absorbance samples match the beginning of EEM sample name (see details).

#### Usage

```
eem_dilcorr(eem_list, abs_data, dilution, auto = FALSE, verbose = TRUE)
```

### Arguments

eem_list	eemlist
abs_data	absorbance data as data frame
dilution	dilution data as data frame with rownames
auto	way how to deal with dilution is chosen automatically. See details.
verbose	print out more information

### Details

If you choose an automatic analysis EEMs are renamed if there is only one matching undiluted absorbance sample. Matching samples is done by comparing the beginning of the sample name (e.g. "sample3\_1to10" fits "sample3").

### Value

data frame

## eem\_dilution

#### Examples

# no appropriate exmaple data available yet

eem\_dilution

Modifying fluorescence data according to dilution.

#### Description

If samples were diluted before measuring, a dilution factor has to be added to the measured data. This function can do that by either multilpying each sample with the same value or using a data frame with different values for each sample.

### Usage

```
eem_dilution(data, dilution = 1)
```

## Arguments

data	fluorescence data with class eemlist
dilution	dilution factor(s), either numeric value or data frame. Row names of data frame
	have to be similar to sample names in eemlist.

# Value

fluorescence data with class eemlist

## Examples

```
data(eem_list)
```

eem\_list2 <- eem\_dilution(eem\_list, dilution = 5)</pre>

dilutionT <- data.frame(dilution = rep(5, length(eem\_list)))
row.names(dilutionT) <- eem\_names(eem\_list)
dilutionT</pre>

eem\_list3 <- eem\_dilution(eem\_list, dilution = dilutionT)</pre>

eem\_duplicates

### Description

Check for duplicate sample names

#### Usage

```
eem_duplicates(data)
## Default S3 method:
eem_duplicates(data)
## S3 method for class 'eemlist'
eem_duplicates(data)
```

## S3 method for class 'data.frame'
eem\_duplicates(data)

### Arguments

data eemlist or data.frame containing absorbance data

### Value

named character vector with duplicate sample names

#### Examples

### check

eem\_easy

Opens an R markdown template for an easy and userfriendly analysis of EEM data.

#### Description

In your default editor (e.g. RStudio), a Rmd file is opened. It consists of bloacks gathering the parameters and information needed and continues with a series of data corrections, peak picking and plots. Finally you get a report of your analysis, a table with the peaks and optional pngs of your fluorescence data. To continue working and keeping your settings, the file can be sa ved anywhere and reused anytime.

#### Usage

eem\_easy()

## eem\_eemdil

# Details

Function does not work well in Windows. You might try file.edit(system.file("EEM\_simple\_analysis.Rmd", package = "staRdom"))

#### Value

A pdf report, a peak picking table and optional plots.

#### Examples

```
## Not run:
#
eem_easy()
# this function fails very often, so you might use that:
file.edit(system.file("EEM_simple_analysis.Rmd", package = "staRdom"))
## End(Not run)
```

```
eem_eemdil
```

Correct names of EEM samples to match undiluted absorbance data.

#### Description

Correct names of EEM samples to match undiluted absorbance data.

# Usage

```
eem_eemdil(
    eem_list,
    abs_data = NULL,
    dilution = NULL,
    cor_data = NULL,
    auto = TRUE,
    verbose = FALSE
)
```

### Arguments

eem_list	eemlist
abs_data	optinal absorbance data as data frame
dilution	optinal dilution data as data frame
cor_data	optional output from eem_dilcorr as data frame
auto	optional, see eem_dilcorr
verbose	optional, see eem_dilcorr

### Value

eemlist

### Examples

# no appropriate exmaple data available yet

eem\_exclude

Exclude complete wavelengths or samples form data set

# Description

Outliers in all modes should be avoided. With this functions excitation or emission wavelengths as well as samples can be removed completely from your sample set.

### Usage

```
eem_exclude(eem_list, exclude = list, verbose = FALSE)
```

#### Arguments

eem_list	object of class eemlist
exclude	list of three vectors, see details
verbose	states whether additional information is given in the command line

### Details

The argument exclude is a named list of three vectors. The names must be "ex", "em" and "sample". Each element contains a vector of wavelengths or sample names that are to be excluded from the data set.

#### Value

object of class eemlist

## Examples

```
data(eem_list)
exclude <- list("ex" = c(280,285,290,295),
    "em" = c(),
    "sample" = c("667sf", "494sf")
)</pre>
```

eem\_list\_ex <- eem\_exclude(eem\_list, exclude)</pre>

eem\_export

### Description

Export all samples of an eem\_list

### Usage

```
eem_export(file, format = c("csv", "mat"), ...)
```

## Arguments

file	path to directory (csv format) or file (Matlab format)
format	either "csv" or "mat" to specify export format
	one or more eem_list objects

#### Value

0 on successful export

### Examples

```
# create temporary directory to write out
file <- paste0(tempdir(),"/eem_export/")
dir.create(file)
# run eem_export to write one csv file for each sample
eem_export(file, format = "csv", eem_list)
# show content of output directory
dir(file)
```

eem\_extend2largest EEM sample data is extended to include all wavelengths in all samples

## Description

Compared to the whole sample set, wavelengths missing in some samples are added and set NA or interpolated. This can be especially helpful, if you want to combine data measured with different wavelength intervals in a given range.

#### Usage

```
eem_extend2largest(eem_list, interpolation = FALSE, ...)
```

eem\_getextreme

### Arguments

eem_list	eemlist
interpolation	logical, whether added NAs should be interpolated
	arguments passed to eem_interp

## Value

eemlist

## Examples

```
library(dplyr)
data(eem_list)
eem_list <- eem_exclude(eem_list[1:5] %>%
`class<-`("eemlist"), exclude = list(em = c(318,322,326,550,438), ex = c(270,275))) %>%
eem_bind(eem_list[6:15] %>% `class<-`("eemlist"))
ggeem(eem_list)
eem_extend2largest(eem_list) %>%
ggeem()
```

eem_getextreme	Determines the the biggest range of EEM spectrum where data is
	available from each sample.

## Description

Determines the the biggest range of EEM spectrum where data is available from each sample.

# Usage

eem\_getextreme(data)

### Arguments

data eemlist

#### Value

list of numeric vector containing the biggest available range

## Examples

```
data(eem_list)
eem_getextreme(eem_list)
eem_list <- eem_range(eem_list,ex = c(250,Inf),em = c(280,500))
eem_getextreme(eem_list)</pre>
```

eem\_hitachi

## Description

This function can be used to import txt files from Hitachi F-7000 containing EEM data using eem\_read.

## Usage

```
eem_hitachi(file)
```

# Arguments

file

path to file passed from eem\_read

# Value

list with EEM data

## Examples

```
## no example data provided with the package
## below is an example how this could like like
# eems <- "C:/some/path/to/hitachi.TXT"
# eem_list <- eem_read(eems, recursive = TRUE, import_function = eem_hitachi)
# eem_list</pre>
```

eem\_ife\_correction Wrapper function to allow eem\_inner\_filter\_effect (eemR) handling different cuvette lengths.

## Description

Calls eem\_inner\_filter\_effect for each sample to use different cuvette lengths.

## Usage

```
eem_ife_correction(
   data,
   abs_data,
   cuvl = NULL,
   unit = c("absorbance", "absorption")
)
```

### Arguments

data	fluorescence data of class eemlist
abs_data	absorbance data
cuvl	length of cuvette of absorption measurment in cm. Either a number or a data frame. Row names of data frame have to be similar to sample names in data. This is ignored, if unit is "absorption".
unit	unit of absorbance data. Either "absorbance" or "absorption".

# Value

fluorescence data of class eemlist

## Examples

```
folder <- system.file("extdata/cary/scans_day_1", package = "eemR") # load example data
eem_list <- eem_read(folder, import_function = "cary")
data(absorbance)</pre>
```

eem\_import\_dir Load all eemlist obects saved in different Rdata or RDa files in a folder.

## Description

Reads Rdata and RDa files with one eemlist each. The eemlists are combined into one and returned.

## Usage

```
eem_import_dir(dir, verbose = FALSE)
```

### Arguments

dir	folder where RData files are saved
verbose	logical, set TRUE to show more information during import

#### Value

eemlist

# Examples

```
## Not run:
```

```
\ensuremath{\texttt{\#}} due to package size issues no example data is provided for this function
```

```
# eem_import_dir("C:/some_folder/with_EEMS/only_Rdata_files")
```

## End(Not run)

eem\_interp

# Description

Missing EEM data can be interpolated. Usually it is the result of removing scatter or other parts where noise is presumed. Different interpolation algorithms can be used (see details).

#### Usage

```
eem_interp(
   data,
   cores = parallel::detectCores(logical = FALSE),
   type = TRUE,
   verbose = FALSE,
   nonneg = TRUE,
   extend = FALSE,
   ...
)
```

#### Arguments

data	object of class eemlist with spectra containing missing values
cores	specify number of cores for parallel computation
type	numeric 0 to 4 or TRUE which resembles type 1
verbose	logical, whether more information on calculation should be provided
nonneg	logical, whether negative values should be replaced by 0
extend	logical, whether data is extrapolated using type 1
	arguments passed on to other functions (pchip, na.approx, mba.points)

## Details

The types of interpolation are (0) setting all NAs to 0, (1) spline interpolation with mba.points, (2) excitation and emission wavelength-wise interpolation with pchip and subsequent mean, (3) excitation wavelength-wise interpolation with pchip and (4) linear interpolation in 2 dimensions with na.approx and again subsequent mean calculation. Calculating the mean is a way of ensuring NAs are also interpolated where missing boundary values would make that impossible. Using type = 1, extrapolation can be suppressed by adding the argument extend = FALSE.

#### Value

object of class eemlist with interpoleted spectra.

#### References

Elcoroaristizabal, S., Bro, R., García, J., Alonso, L. 2015. PARAFAC models of fluorescence data with scattering: A comparative study. Chemometrics and Intelligent Laboratory Systems, 142, 124-130 doi:10.1016/j.chemolab.2015.01.017

# See Also

pchip, mba.points, na.approx

## Examples

```
data(eem_list)
eem_list <- eem_list[1:6]
class(eem_list) <- "eemlist"
remove_scatter <- c(FALSE, TRUE, TRUE, TRUE)
remove_scatter_width = c(15, 10, 16, 12)
eem_list <- eem_rem_scat(eem_list, remove_scatter, remove_scatter_width)
eem_list <- eem_interp(eem_list, cores = 2)
ggeem(eem_list)
eem_list2 <- eem_setNA(eem_list, ex = 200:280, interpolate=FALSE)
ggeem(eem_list2)
eem_list3 <- eem_interp(eem_list2, type = 1, extend = TRUE, cores = 2)
ggeem(eem_list3)
eem_list3 <- eem_interp(eem_list2, type = 1, extend = FALSE, cores = 2)
ggeem(eem_list3)</pre>
```

eem\_is.na

Check for NAs in EEM data

### Description

Check for NAs in EEM data

## eem\_list

## Usage

eem\_is.na(eem\_list)

# Arguments

eem\_list eemlist to check

## Value

named character vector with sample names where EEM data contains NAs

## Examples

### check

eem\_list

15 fluorescence samples from drEEM used for examples.

# Description

15 fluorescence samples from drEEM used for examples.

# Usage

eem\_list

# Format

eemlist

eem_list_outliers	2 fluorescence samples from drEEM that were excluded as outliers
	from the PARAFAC model.

# Description

2 fluorescence samples from drEEM that were excluded as outliers from the PARAFAC model.

# Usage

eem\_list\_outliers

### Format

eemlist

eem\_load\_dreem

## Description

Load original data from the drEEM tutorial and return it as eemlist

### Usage

```
eem_load_dreem()
```

### Value

eemlist

## Examples

```
# Reading MATLAB files from recent versions like the demo dataset from drEEM
# can cause problems if the R installation lacks UTF8 support in iconv.
# Therefore, we use try() in the example. If you encounter related problems,
# please refer to the help for R.matlab::readMat() for details.
eem_list <- try(eem_load_dreem(), silent = FALSE)</pre>
```

```
eem_list
```

eem\_matmult

Multiply all EEMs with a matrix

## Description

Multiply all EEMs with a matrix

#### Usage

```
eem_matmult(eem_list, matrix = NULL, value = 0)
```

### Arguments

eem_list	EEM data as eemlist
matrix	either a vactor containing "l" and/or "u" or a matrix, see details.
value	in case matrices "l" or "u" are used, this specifies the value to use in this areas. Usually this is 0 (default) or NA but any numeric value can be used.

## Details

All EEMs must be of the same size. If matrix is of type matrix, it is used right away to multiply the EEMs. It has to be of the same size as the EEMs. If matrix is a vector containing "l", values below 1st order Rayleigh scattering are set to 0. If matrix contains "u", values above 2nd order Raman scattering are set to 0. If you want to remove wavelength ranges, take into consideration to use eem\_cut or eem\_range.

### Value

eemlist

### Examples

```
data(eem_list)
eem <- eem_list[1:9]
class(eem) <- "eemlist"
ggeem(eem)
eem_list_cut <- eem_matmult(eem,matrix=c("1"), value= NA)
ggeem(eem_list_cut)</pre>
```

eem_metatemplate	Create table that contains sample names and locations of files.
------------------	---

# Description

You can use this table as an overview of your files and/or as a template for creating a metadata table.

#### Usage

```
eem_metatemplate(eem_list = NULL, absorbance = NULL)
```

### Arguments

eem_list	eemlist
absorbance	data frame with absorbance data

#### Value

data frame

#### Examples

```
folder <- system.file("extdata/EEMs", package = "staRdom") # load example data
eem_list <- eem_read(folder, recursive = TRUE, import_function = eem_csv)
data(absorbance)</pre>
```

```
eem_metatemplate(eem_list,absorbance)
```

eem\_name\_replace Replace matched patterns in sample names

## Description

Sample names in eemlist can be altered.

### Usage

eem\_name\_replace(eem\_list, pattern, replacement)

#### Arguments

eem_list	data of class eemlist
pattern	character vector containing pattern to look for.
replacement	character vector of replacements. Has to have the same length as pattern

# Details

str\_replace\_all from package stringr is used for the replacement. Please read the corresponding help for further options.

# Value

An eemlist.

# See Also

str\_replace\_all

#### Examples

```
eem_names(eem_list)
```

```
eem_list <- eem_name_replace(eem_list,"sample","Sample")
eem_names(eem_list)</pre>
```

eem\_overview\_plot Plot fluorescence data from several samples split into several plots.

#### Description

Plot fluorescence data from several samples split into several plots.

### Usage

```
eem_overview_plot(data, spp = 8, ...)
```

#### Arguments

data	fluorescence data of class eemlist
spp	number of samples per plot or a vector with the numbers of rows and columns in the plot.
	arguments passed on to ggeem

## Value

list of ggplots

## Examples

```
data(eem_list)
eem_overview_plot(eem_list, spp = 9)
# define number of rows and columns in plot
eem_overview_plot(eem_list, spp = c(3, 5))
```

eem\_parafac

```
Runs a PARAFAC analysis on EEM data
```

# Description

One or more PARAFAC models can be calculated depending on the number of components. The idea is to compare the different models to get the most suitable. B-mode is emmission wavelengths, C-mode is excitation wavelengths and, A-mode is the loadings of the samples. The calculation is done with parafac, please see details there.

### Usage

```
eem_parafac(
    eem_list,
    comps,
    maxit = 2500,
    normalise = TRUE,
    const = c("nonneg", "nonneg", "nonneg"),
    nstart = 30,
    ctol = 10^-8,
    strictly_converging = FALSE,
    cores = parallel::detectCores(logical = FALSE),
    verbose = FALSE,
    output = "best",
    ...
)
```

## Arguments

eem_list	object of class eem	
comps	vector containing the desired numbers of components. For each of these numbers one model is calculated	
maxit	maximum iterations for PARAFAC algorithm	
normalise	state whether EEM data should be normalised in advance	
const	constraints of PARAFAC analysis. Default is non-negative ("nonneg"), alter- natively smooth and non-negative ("smonon") might be interesting for an EEM analysis.	
nstart	number of random starts	
ctol	Convergence tolerance (R <sup>2</sup> change)	
strictly_converging		
	calculate nstart converging models and take the best. Please see details!	
cores	number of parallel calculations (e.g. number of physical cores in CPU)	
verbose	print infos	
output	Output the "best" solution (default) only or additionally add "all" nstart solutions to the model as an element named "models".	
	additional parameters that are passed on to parafac	

### Details

PARAFAC models are created based on multiple random starts. In some cases, a model does not converge and the resulting model is then based on less than nstart converging models. In case you want to have nstart converging models, set strictly\_converging TRUE. This calculates models stepwise until the desired number is reached but it takes more calculation time. Increasing the number of models from the beginning is much more time efficient.

eem\_raman\_area

#### Value

object of class parafac

#### See Also

parafac

### Examples

data(eem\_list)

```
dim_min <- 3 # minimum number of components
dim_max <- 7 # maximum number of components</pre>
nstart <- 25 # random starts for PARAFAC analysis, models built simulanuously, best selected
# cores <- parallel::detectCores(logical=FALSE) # use all cores but do not use all threads
cores <- 2 # package checks only run with 2 cores
maxit = 2500
ctol <- 10<sup>-7</sup> # tolerance for parafac
pfres_comps <- eem_parafac(eem_list, comps = seq(dim_min, dim_max),</pre>
    normalise = TRUE, maxit = maxit, nstart = nstart, ctol = ctol, cores = cores)
## with a defined number of converging models
#pfres_comps <- eem_parafac(eem_list, comps = seq(dim_min, dim_max),</pre>
      normalise = TRUE, maxit = maxit, nstart = nstart, ctol = ctol,
#
      output = "all", strictly_converging = TRUE, cores = cores, verbose = TRUE)
#
pfres_comps2 <- eem_parafac(eem_list, comps = seq(dim_min, dim_max),</pre>
  normalise = TRUE, maxit = maxit, nstart = nstart, ctol = ctol, cores = cores, output = "all")
```

eem\_raman\_area Calculate raman area of EEM samples

#### Description

Calculate raman area of EEM samples

#### Usage

```
eem_raman_area(eem_list, blanks_only = TRUE, average = FALSE)
```

## Arguments

eem_list	An object of class eemlist.	
blanks_only	logical. States whether all samples or just blanks will be used.	
average	logical. States whether samples will be averaged before calculating the raman	
	area.	

#### Details

Code based on eem\_raman\_normalisation.

### Value

data frame containing sample names, locations and raman areas

### Examples

```
folder <- system.file("extdata/EEMs",package="staRdom")
eem_list <- eem_read(folder, recursive = TRUE, import_function = eem_csv)
blank <- eem_extract(eem_list,sample ="blank", keep = TRUE)</pre>
```

eem\_raman\_area(blank)

eem\_raman\_normalisation2

Wrapper function to eem\_raman\_normalisation (eemR).

#### Description

Usually Raman normalisation is done with fluorescence data from a blank sample. Sometimes you already know a value for the Raman area. This function can do both.

#### Usage

```
eem_raman_normalisation2(data, blank = "blank")
```

### Arguments

data	fluorescence data of class eemlist
blank	defines how Raman normalisation is done (see 'Details')

## Details

Possible values for blank:

"blank": normalisation is done with a blank sample. Please refer to eem\_raman\_normalisation.

numeric: normalisation is done with one value for all samples.

data frame: normalisation is done with different values for different samples. Values are taken from a data.frame with sample names as rownames and one column containing the raman area values.

#### Value

fluorescence data of class eemlist

#### eem\_range

### Examples

```
data(eem_list)
# correction by blank
eems_bl <- eem_raman_normalisation2(eem_list,blank="blank")
# correction by value</pre>
```

```
eems_num <- eem_raman_normalisation2(eem_list,blank=168)</pre>
```

eem\_range

Cut EEM data matching a given wavelength range

## Description

Cut EEM data matching a given wavelength range

#### Usage

eem\_range(data, ex = c(0, Inf), em = c(0, Inf))

#### Arguments

data	EEM data as eemlist
ex	optional desired range of excitation wavelength
em	optional desired range of emission wavelength

## Value

An eemlist of reduced spectra size.

### Examples

```
data(eem_list)
eem_range(eem_list,ex = c(250,Inf),em = c(280,500))
```

eem_read_csv	Import EEMs from g	generic csv tables (deprecated)
--------------	--------------------	---------------------------------

## Description

This function is deprecate, please use  $eem_read(..., import_function = eem_csv)$  or  $eem_read(..., import_function = eem_csv2)$  instead. EEM data is loaded from generic files. First column and first row contains wavelength values. The other values are to be plain numbers. fread is used to read the table. It offers a lot of helpful functions (e.g. skipping any number n of header lines by adding 'skip = n')

## Usage

```
eem_read_csv(
   path,
   col = "ex",
   recursive = TRUE,
   is_blank_corrected = FALSE,
   is_scatter_corrected = FALSE,
   is_ife_corrected = FALSE,
   is_raman_normalized = FALSE,
   manufacturer = "unknown",
   ...
)
```

# Arguments

path	path to file(s), either a filename or a folder	
col	either "ex" or "em", what wavelengths are in the columns	
recursive	logical, whether directories are loaded recursively	
is_blank_corre	cted	
	logical, whether blank correction was done	
is_scatter_corrected		
	logical, wether scatters were corrected	
is_ife_corrected		
	logical, wether inner-filter effect correction was done	
is_raman_normalized		
	logical, wether raman normalisation applied	
manufacturer	string specifying manufacturer of instrument	
	parameters from other functions, currently not used	

# Examples

```
eems <- system.file("extdata/EEMs",package="staRdom")
eem_list <- eem_read_csv(eems)</pre>
```

eem\_list

eem_red2smallest	Remove wavelengths, that are missing in at least one sample form the
	whole set.

# Description

Remove wavelengths, that are missing in at least one sample form the whole set.

#### eem\_rem\_scat

## Usage

```
eem_red2smallest(data, verbose = FALSE)
```

#### Arguments

data	data of EEM samples as eemlist
verbose	states whether additional information is given in the command line

## Details

This step is neccessary to perform a PARAFAC analysis which can only be calculated with spectra of similar range.

## Value

eemlist with reduced spectral width

## Examples

```
require(dplyr)
data(eem_list)
eem_list_red <- eem_red2smallest(eem_list)
# create an eemlist where data is missing
eem_list2 <- eem_exclude(eem_list,
    list("ex" = c(280,290,350),
        "em" = c(402,510),
        "sample" = c()))
# modify names of samples with missing data
eem_names(eem_list2) <- paste0("x",eem_names(eem_list2))
# combined the lists with and without missing data
eem_list3 <- eem_bind(eem_list,eem_list2)
# ggeem(eem_list3)
# module the data is the whole complete to the merplete to
```

```
# reduce the data in the whole sampleset to the smallest wavelengths that are present in all samples
eem_list4 <- eem_red2smallest(eem_list3)
#ggeem(eem_list4)</pre>
```

eem\_rem\_scat

Remove Raman and Rayleigh scattering in fluorescence data

#### Description

Wrapper function to remove several scatterings in one step using eem\_remove\_scattering.

# Usage

```
eem_rem_scat(
   data,
   remove_scatter,
   remove_scatter_width = 10,
   interpolation = FALSE,
   cores = parallel::detectCores(logical = FALSE),
   verbose = FALSE
)
```

# Arguments

data	object of class eemlist	
remove_scatter	logical vector. The meanings of the vector are "raman1", "raman2", "rayleigh1" and "rayleigh2" scattering. Set TRUE if certain scattering should be removed.	
remove_scatter	_width	
	numeric vector containing width of scattering to remove. If there is only one element in this vector, each this is the width of each removed scattering. If there are 4 values, differnt widths are used ordered by "raman1", "raman2", "rayleigh1" and "rayleigh2".	
interpolation	logical, optionally states whether interpolation is done right away	
cores	optional, CPU cores to use for interpolation	
verbose	logical, provide additional information	

### Value

eemlist

## Examples

```
data(eem_list)
remove_scatter <- c(TRUE, TRUE, TRUE, TRUE)
remove_scatter_width = c(15,10,16,12)
eems <- eem_rem_scat(eem_list,remove_scatter,remove_scatter_width)
ggeem(eems)</pre>
```

eem\_scale\_ext Determine the range of fluorescence values in a set of samples

# Description

Determine the range of fluorescence values in a set of samples

## eem\_setNA

# Usage

eem\_scale\_ext(data)

## Arguments

data eemlist containing the EEM data

# Value

numeric vector

# Examples

```
data(eem_list)
eem_scale_ext(eem_list)
```

eem_setNA	set parts of specific samples to NA and optionally interpolate these
	parts

# Description

set parts of specific samples to NA and optionally interpolate these parts

# Usage

```
eem_setNA(
    eem_list,
    sample = NULL,
    em = NULL,
    ex = NULL,
    interpolate = TRUE,
    ...
)
```

# Arguments

eem_list	EEMs as eemlist
sample	optional, names or indices of samples to process
em	optional, emission wavelengths to set NA
ex	optional, excitation wavelengths to set NA
interpolate	FALSE, 1 or 2, interpolate NAs or not, 2 different methods, see eem_interp
	arguments passed on to eem_interp

# Details

Samples and wavelengths are optional and if not set all of them are considered in setting data to NA. Wavelengths can be set as vectors containing more than the wavelengths present in the data. E.g. 230:250 removes all wavelengths between 230 and 250 if present. Data is best interpolated if it does not reach data boundaries. Please check the results otherwise as in some cases the interpolation might not produce meaningful data.

## Value

eemlist

# Examples

```
data(eem_list)
eem <- eem_list[1:9]
class(eem) <- "eemlist"
```

ggeem(eem)

eem\_list2 <- eem\_setNA(eem,ex=200:280,em=500:600, interpolate=FALSE)
ggeem(eem\_list2)</pre>

eem_smooth	Smooth fluorescence data by calculating rolling mean along excitation
	wavelengths.

## Description

Smooth fluorescence data by calculating rolling mean along excitation wavelengths.

### Usage

```
eem_smooth(data, n = 4, cores = parallel::detectCores(logical = FALSE))
```

## Arguments

data	fluorescence data of class eemlist
n	width of rolling mean window in nm
cores	number of CPU cores to be used

## Value

eemlist with smoothed data
eem\_spectral\_cor

# Examples

data(eem\_list)

```
eem_list <- eem_smooth(eem_list, n = 4, cores = 2)</pre>
```

<pre>eem_spectral_cor</pre>	Multiply EEMs with spectral correction vectors (Emission and Exci-
	tation)

## Description

Multiply EEMs with spectral correction vectors (Emission and Excitation)

#### Usage

```
eem_spectral_cor(eem_list, Excor, Emcor)
```

## Arguments

eem_list	eemlist
Excor	data frame, first column wavelengths, second column excitation correction
Emcor	data frame, first column wavelengths, second column emission correction

#### Value

eemlist

#### Examples

```
eems <- system.file("extdata/EEMs",package="staRdom")
eem_list <- eem_read(eems, recursive = TRUE, import_function = eem_csv)
excorfile <- system.file("extdata/CorrectionFiles/xc06se06n.csv",package="staRdom")
Excor <- data.table::fread(excorfile)
emcorfile <- system.file("extdata/CorrectionFiles/mcorrs_4nm.csv",package="staRdom")
Emcor <- data.table::fread(emcorfile)
# adjust range of EEMs to cover correction vectors
eem_list <- eem_range(eem_list,ex = range(Excor[,1]), em = range(Emcor[,1]))</pre>
```

```
eem_list_sc <- eem_spectral_cor(eem_list,Excor,Emcor)</pre>
```

eem\_write\_csv

# Description

Export samples in an EEM list to a single csv files

#### Usage

```
eem_write_csv(eem_list, output, ...)
```

## Arguments

eem_list	EEM data as eemlist
output	path to folder where csv files are exported to
	additional arguments

#### Value

returns the exported EEMs as a list of data.frames

## Examples

```
data(eem_list)
output <- tempdir()
output
a <- eem_write_csv(eem_list, output)</pre>
```

ggeem

EEM spectra plotted with ggplot2

# Description

Plots from EEM spectra of class ggplot. In case you work with a larger number of EEMs and want to show then in several plots, you can use eem\_overview\_plot.

## Usage

```
ggeem(data, fill_max = FALSE, ...)
## Default S3 method:
ggeem(data, fill_max = FALSE, ...)
## S3 method for class 'eemlist'
ggeem(data, fill_max = FALSE, eemlist_order = TRUE, ...)
```

```
## S3 method for class 'eem'
ggeem(data, fill_max = FALSE, ...)
## S3 method for class 'parafac'
ggeem(data, fill_max = FALSE, ...)
## S3 method for class 'data.frame'
ggeem(
    data,
    fill_max = FALSE,
    colpal = "default",
    contour = FALSE,
    interpolate = FALSE,
    redneg = NULL,
    ...
)
```

#### Arguments

data	eem, eemlist, parafac or data.frame. The details are given under 'Details'.
fill_max	sets the maximum fluorescence value for the colour scale. This is mainly used
	by other functions, and makes different plots visually comparable.
	parameters passed on to ggplot.
eemlist_order	logical, in case of an eemlist, the order of samples in the plot is the same as in the eemlist, alphabetically otherwise
colpal	"default" to use the viridis colour palette, "rainbow" to use a subset of the rain- bow palette, any custom vector of colors or a colour palette. A gradient will be produced from this vector. Larger vectors (e.g. 50 elements) can produce smoother gradients.
contour	logical, whether contours should be plotted (default FALSE), see geom_contour
interpolate	logical, whether fluorescence should be interpolated, see geom_raster
redneg	deprecated! logical, whether negative values should be coloured discreet.

## Details

The data can be of different sources: eem: a single EEM spectrum is plotted eemlist: all spectra of the samples are plotted, arranged in a grid data.frame: a data.frame containing EEM data. Can be created by e.g. as.data.frame.eem parafac: a PARAFAC model, the components are plotted then.

Using redneg you can give negative values a reddish colour. This can help identifying these parts in samples or components. Negative values are physically not possible and can only be the result of measuring errors, model deviations and problems with interpolated values.

Interpolation (interpolate = TRUE) leeds to smoother plots. The default is FALSE because it might cover small scale inconsistencies.

Contours (contour = TRUE)can be added to the EEM plots.

A colour palette can be specified using the argument colpal.

Plotting distinct samples can be done using eem\_extract. Please see example.

# Value

a ggplot object

# Examples

```
## plotting two distinct samples
data(eem_list)
eem_names(eem_list)
eem <- eem_extract(eem_list,c("^d667sf$", "^d661sf$"),keep=TRUE)
ggeem(eem)</pre>
```

```
# the former redneg argument is deprecated, please see a similar looking example below!
#ggeem(eem, redneg = TRUE)
ggeem(eem, colpal = c(rainbow(75)[58],rainbow(75)[53:1]))
```

```
# use any custom colour palette
ggeem(eem, colpal = heat.colors(50))
# needs package matlab to be installed:
# ggeem(eem, colpal = matlab::jet.colors(50))
# or by adding ggplot2 colour and fill functions:
# ggeem(eem)+
# scale_fill_viridis_c()+
# scale_color_viridis_c()
ggeem(eem, interpolate = TRUE)
```

```
ggeem(eem, contour = TRUE)
```

list\_join

#### Full join of a list of data frames.

# Description

Full join of a list of data frames.

## Usage

list\_join(df\_list, by)

## Arguments

df_list	list of data frames to by joined
by	character vector containing information how to join data frames. Format to be according to by in full_join. Each data frame has to contain the column(s) used for joining.

## Value

The joint data frame.

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## maxlines

## See Also

full\_join

# Examples

a <- data.frame(what=letters[1:5],a=c(1:5)) b <- data.frame(what=letters[1:5],b=c(7:11)) c <- data.frame(what=letters[1:5],c=c(20:24)) df\_list <- list(a,b,c)</pre>

list\_join(df\_list,by="what")

maxlines

*Extract data from emission and excitation wavelengths of the components of a PARAFAC model (scaled B- and C-modes)* 

## Description

Data for each wavelengths is returned. For each component the lines intersecting at the component maxima are returned.

## Usage

maxlines(pfmodel)

# Arguments

pfmodel object of class parafac

## Value

data frame

## Examples

data(pf\_models)

ml <- maxlines(pf4[[1]])</pre>

norm2A

# Description

Factors used for normalisation are saved separately in the PARAFAC models. With this function, the normalisation factors are combined with the A-modes of the model and removed as a separate vector. This means former normalisation is accounted for in the amount of each component in each sample. If no normalisation was done, the original model is returned without warning.

# Usage

norm2A(pfmodel)

## Arguments

pfmodel object of class parafac

## Value

object of class parafac

## Examples

```
data(pf_models)
```

```
pf4[[1]] <- norm2A(pf4[[1]])</pre>
```

norm\_array

Normalise 3-dimensional array in first and second dimension

## Description

Normalise 3-dimensional array in first and second dimension

## Usage

norm\_array(eem\_array)

## Arguments

eem\_array 3-dimensional array

## Value

array

## parafac\_conv

# Examples

```
data(eem_list)
```

```
a <- eem2array(eem_list)
an <- norm_array(a)</pre>
```

parafac\_conv

Calculate a PARAFAC model similar to and using parafac.

# Description

Please refer to parafac for input parameters and details. This wrapper function ensures 'nstart' converging models are calculated. On the contrary, parafac calculates 'nstart' models regardless if they are converging.

## Usage

```
parafac_conv(
    X,
    nstart,
    verbose = FALSE,
    output = c("best", "all"),
    cl = NULL,
    ...
)
```

# Arguments

Х	array
nstart	number of converging models to calculate
verbose	logical, whether more information is supplied
output	Output the best solution (default) or output all nstart solutions.
cl	cluster to be used for parallel processing
	arguments passed on to parafac

# Value

either a parafac model or a list of parafac models

## See Also

parafac

## Examples

data(eem\_list)

80

```
dim_min <- 3 # minimum number of components
dim_max <- 4 # maximum number of components
nstart <- 25 # random starts for PARAFAC analysis, models built simulanuously, best selected
# cores <- parallel::detectCores(logical=FALSE) # use all cores but do not use all threads
cores <- 2 # package checks only run with 2 cores
maxit = 2500
ctol <- 10^-7 # tolerance for parafac
pfres_comps <- eem_parafac(eem_list, comps = seq(dim_min, dim_max),
    normalise = TRUE, strictly_converging = TRUE, maxit = maxit, nstart = nstart,
    ctol = ctol, cores = cores)
# keep all calculated models for diagnostics
pfres_comps_all <- eem_parafac(eem_list, comps = seq(dim_min, dim_max),
    normalise = TRUE, strictly_converging = TRUE, output = "all", maxit = maxit,
    nstart = nstart, ctol = ctol, cores = cores)
```

pf1

PARAFAC model, see vignette, unconstrained

## Description

PARAFAC model, see vignette, unconstrained

# Usage

pf1

# Format

list of parafacs

pf1n

PARAFAC model, see vignette, non-negative constraints

## Description

PARAFAC model, see vignette, non-negative constraints

#### Usage

pf1n

pf1n

# pf2

# Format

list of parafacs

pf2

PARAFAC model, see vignette, non-negative constraints, normalised

# Description

PARAFAC model, see vignette, non-negative constraints, normalised

# Usage

pf2

# Format

list of parafacs

pf3	PARAFAC model, see vignette, non-negative constraints, normalised,
	outliers removed

# Description

PARAFAC model, see vignette, non-negative constraints, normalised, outliers removed

# Usage

pf3

# Format

list of parafacs

pf4

PARAFAC model, see vignette, non-negative constraints, normalised, outliers removed, high accuarcy

# Description

PARAFAC model, see vignette, non-negative constraints, normalised, outliers removed, high accuarcy

# Usage

pf4

# Format

list of parafacs

sh

result from PARAFAC split-half analysis, periodic data split

# Description

result from PARAFAC split-half analysis, periodic data split

## Usage

sh

## Format

list of parafacs

splithalf

## Description

The samples are split into four subsamples: A,B,C,D. Subsamples are then combined and compared: AB vs. CD, AC vs. BD, AD vs. BC. The results show graphs from the components of each of the 6 models.

## Usage

```
splithalf(
  eem_list,
  comps,
  splits = NA,
  rand = FALSE,
  normalise = TRUE,
  nstart = 20,
  cores = parallel::detectCores(logical = FALSE),
  maxit = 2500,
  ctol = 10^(-7),
  rescale = TRUE,
  strictly_converging = FALSE,
  verbose = FALSE,
  ....
)
```

# Arguments

eem_list	eemlist containing sample data
comps	number of desired components
splits	optional, list of 4 numerical vectors containing the sample numbers for A,B,C and D sample subsets
rand	logical, splits are randomised
normalise	state whether EEM data should be normalised in advance
nstart	number of random starts
cores	number of parallel calculations (e.g. number of physical cores in CPU)
maxit	maximum iterations for PARAFAC algorithm
ctol	Convergence tolerance (R <sup>2</sup> change)
rescale	rescale splithalf models to Fmax, see eempf_rescaleBC
strictly_conver	ging
	calculate nstart converging models and take the best. Please see eem_parafac.
verbose	states whether you want additional information during calculation
	additional parameters that are passed on to parafac

## Details

Split data sets can be split suboptimal and cause low TCCs. Therefore, subsamples are recombined in 3 different ways and a TCC close to 1 in only one split combination per component is already a positive result. Check the split sets to check for sample independency.

# Value

data frame containing components of the splithalf models

## See Also

splithalf\_plot, splithalf\_tcc

# Examples

```
data(eem_list)
```

```
splithalf <- splithalf(eem_list, comps = 6, verbose = TRUE, cores = 2)
splithalf_plot(splithalf)</pre>
```

```
# Similarity of splits using SSCs
sscs <- splithalf_tcc(splithalf)</pre>
```

splithalf\_plot Plot results from a splithalf analysis

# Description

Graphs of all components of all models are plotted to be compared.

## Usage

```
splithalf_plot(fits)
```

#### Arguments

fits list of components data

# Value

ggplot

# See Also

splithalf

## splithalf\_splits

## Examples

data(sh)

```
splithalf_plot(sh)
str(sh)
```

splithalf\_splits *Extracting a list of sample names in each subsample from a splithalf analysis* 

# Description

Extracting a list of sample names in each subsample from a splithalf analysis

## Usage

```
splithalf_splits(fits)
```

# Arguments

fits list of parafac models (from a splithalf analysis)

#### Value

data frame containing TCC values

# Examples

```
data(sh)
splithalf_splits(sh)
```

splithalf\_tcc Extracting TCC values from a splithalf analysis

## Description

Extracting TCC values from a splithalf analysis

## Usage

```
splithalf_tcc(fits)
```

## Arguments

fits list of parafac models (from a splithalf analysis)

# Value

data frame containing TCC values

## Examples

data(sh)

splithalf\_tcc(sh)

SSC

*Calculate the shift-and shape-sensitive congruence (SSC) between two matrices* 

## Description

Please see details in: U.J. Wünsch, R. Bro, C.A. Stedmon, P. Wenig, K.R. Murphy, Emerging patterns in the global distribution of dissolved matter fluorescence, Anal. Methods, 11 (2019), pp. 888-893

# Usage

ssc(mat1, mat2, tcc = FALSE)

## Arguments

mat1	matrix
mat2	matrix
tcc	if set TRUE, TCC is returned instead

# Value

table containing pairwise SCC of matrices columns

# Examples

```
pf_models <- pf3
mat1 <- pf_models[[1]][[2]]
mat2 <- pf_models[[2]][[2]]
## calculate SSC
ssc(mat1,mat2)
## calculate TCC
ssc(mat1,mat2, tcc = TRUE)</pre>
```

ssc\_max

## Description

Calculate the combination of components giving the maximum of geometric mean of TCCs

## Usage

ssc\_max(mat)

## Arguments

mat matrix

# Value

vector with TCCs having the highest possible geometric mean

## Examples

```
mat <- matrix(c(7,2,13,6,0,7,1,5,5), nrow = 3)
mat
sscs <- ssc_max(mat)
sscs
# order of components:
attr(sscs,"order")</pre>
```

tcc

Caluclate Tucker's Congruence Coefficient of PARAFAC components

## Description

Componets must be passed as modes, see maxlines

## Usage

tcc(maxl\_table, na.action = "na.omit")

## Arguments

<pre>maxl_table</pre>	data frame containing the peak lines of components
na.action	if "na.omit" NA are deleted from prior the test

## Value

data.frame containing the TCCs

# Examples

```
data(pf_models)
ml <- maxlines(pf4[[1]])
tcc(ml)</pre>
```

<pre>tcc_find_pairs</pre>	Reorders components of different PARAFAC models according to best
	fit (TCC)

# Description

When running a splithalf analysis similar components are not necessarily on the same position. This function looks for best fits with Tucker's Congruence Coefficients and returns a list of models with reordered components.

# Usage

```
tcc_find_pairs(fits)
```

## Arguments

fits list of parafac models

## Value

list of parafac models

## See Also

splithalf

## Examples

data(eem\_list)

# function currently only used from within splithalf splithalf(eem\_list, 6, nstart = 2, cores = 2)

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