# Package 'EMMAgeo'

June 29, 2025

Type Package
Title End-Member Modelling of Grain-Size Data
Version 0.9.9
<b>Date</b> 2025-06-28
Maintainer Michael Dietze <michael.dietze@uni-goettingen.de></michael.dietze@uni-goettingen.de>
<b>Description</b> End-member modelling analysis of grain-size data is an approach to unmix a data set's underlying distributions and their contribution to the data set. EMMAgeo provides deterministic and robust protocols for that purpose.
License GPL-3
Encoding UTF-8
<b>Depends</b> R (>= $4.5$ )
Imports GPArotation, nnls, caTools, shiny
RoxygenNote 7.3.2
NeedsCompilation no
Author Michael Dietze [cre, aut, trl], Elisabeth Dietze [ctb]
Repository CRAN
<b>Date/Publication</b> 2025-06-28 22:20:19 UTC
Contents
EMMAgeo-package       2         check.data       3         click.limits       4         convert.units       5         create.EM       6         EMMA       1         EMpot       16         EMrob       10         get.l       1

2 EMMAgeo-package

EMMA	geo-package			~	 porting functi mpositional a	ions for unmix lata.	;-
Index							38
	X	 	 		 		36
	test.robustness						
	test.parameters						
	test.l.max	 	 		 		30
	test.1	 	 		 		29
	test.factors						
	robust.scores						
	robust.loadings						
	robust.EM						
	residual.EM						
	model.EM						
	mix.EM						
	GUI interpolate.classes .						
	get.q						
	get.limits						
	get.l.opt	 	 		 		12

#### **Description**

EMMAgeo provides a set of functions for end-member modelling analysis (EMMA) of grain-size data and other cases of compositional data. EMMA describes a multivariate data set of m samples, each comprising n parameters (e.g. grain-size classes), as a linear combination of end-member loadings (the underlying distributions) and end-member scores (the contribution of each loading to each sample).

EMMA can be run in two principal ways, a deterministic and a robust, including modelling the uncertainties. The deterministic way can be accessed simply with the function EMMA(). For the robust way there are two protocols that need to be respected. There is a compact protocol, which is mainly automated but needs adjustments by the user, and there is an extended protocol, which allows access to all parameterisation steps of robust EMMA.

The package contains further auxiliary functions to check and prepare input data, test parameters and use a graphic user interface for deterministic EMMA. The package also contains an example data set, comprising meaured grain-size distributions of real world sediment end-members.

#### **Details**

Package: EMMAgeo Type: Package Version: 0.9.8 Date: 2025-03-24 License: GPL-3 check.data 3

# Author(s)

Michael Dietze, Elisabeth Dietze

check.data

Check correctness and consistency of input data

## **Description**

The input data matrix (X), number of end-members (q), weight transformation limits (l) and constant sum scaling parameter (c) are checked. This includes checking for absence of missing values, columns containing only zero-values and for numeric data type of all variables. A further check tests if l is below the maximum possible value, preventing numerical instability prior to factor rotation.

## Usage

```
check.data(X, q, 1, c, ...)
```

## **Arguments**

Χ	Numeric matrix, input data set with m samples (rows) and n variables (columns).

q Numeric scalar, number of end-members to be modelled.

1 Numeric scalar or vector, weight transformation limit, i.e. quantile.

c Numeric scalar, constant sum scaling parameter, e.g. 1, 100, 1000.

. . . Further arguments passed to the function.

## Value

Character vector, verbose test results.

## Author(s)

Michael Dietze, Elisabeth Dietze

#### See Also

EMMA

4 click.limits

## **Examples**

click.limits

Define mode limits by mouse clicks.

## Description

This function allows defining limits for robust end-members by mouse clicks on a combined plot output, showing a histogram and all end-members together. Clicks must be placed in the order lower limit, upper limit - for each end-member successively.

## Usage

```
click.limits(data, n, classunits)
```

## **Arguments**

data List object, output of test.robustness.

n Numeric scalar, number of target end-members.

classunits Numeric vector, optional class units (e.g. micrometers or phi-units).

## Value

Numeric matrix, limit classes. The first row contains lower limits, the second row upper limits for each end-member.

#### Author(s)

Michael Dietze, Elisabeth Dietze

#### See Also

```
test.robustness, robust.EM
```

convert.units 5

## **Examples**

```
## load example data set
data(example_X)

## Test robustness
q <- 4:7
1 <- seq(from = 0, to = 0.1, by = 0.02)
TR <- test.robustness(X = X, q = q, l = 1)

## define 2 limits by mouse clicks (uncomment to use).
# limits <- click.limits(data = TR, n = 2)
# limits</pre>
```

convert.units

Convert between phi and micrometers.

## **Description**

The function converts values from the phi-scale to the micrometer-scale and vice versa.

## Usage

```
convert.units(phi, mu)
```

# Arguments

phi Numeric vector, grain-size class values in phi to be converted.

mu Numeric vector, grain-size class values in micrometres to be converted.

## Value

Numeric vector, converted grain-size class values.

## Author(s)

Michael Dietze, Elisabeth Dietze

## See Also

```
interpolate.classes
```

6 create.EM

## **Examples**

```
## generate phi-values
phi <- -2:5

## convert and show phi to mu
mu <- convert.units(phi = phi)
mu

## convert and show mu to phi
convert.units(mu = mu)</pre>
```

create.EM

Create grain-size-distributions.

## **Description**

This function allows creating artificial grain-size end-members. One such "artificial end-member loading" may be composed of one or more superimposed normal distributions.

## Usage

```
create.EM(p1, p2, s, boundaries, n)
```

## **Arguments**

p1	Numeric vector, means of normal distributions, i.e. mode positions.
p2	Numeric vector, standard deviations of normal distributions, i.e. mode width.
S	Numeric vector, relative proportions of each mode, i.e. relative mode height.
boundaries	Numeric vector of length two with class boundaries (i.e. $c(lower boundary, upper boundary))$ .
n	Numeric scalar with number of classes, i.e. resolution of the end-member.

## **Details**

When building a data set of many artificial end member loadings, these should all have the same boundaries and n. The function builds composites of individual normal distributions. Each distribution is scaled according to s. Finally the distribution is scaled to 100 %.

## Value

Numeric vector with normalised end-member loadings, consisting of the mixed normal distributions according to the input parameters.

# Author(s)

Michael Dietze, Elisabeth Dietze

EMMA 7

## See Also

mix.EM

## **Examples**

**EMMA** 

End-member modelling analysis algorithm.

## **Description**

A multivariate data set (m samples composed of n variables) is decomposed by eigenspace analysis and modelled with a given number of end-members (q). Several steps of scaling, transformation, normalisation, eigenspace decomposition, factor rotation, data modelling and evaluation are performed.

# Usage

```
EMMA(
    X,
    q,
    1,
    c,
    Vqn,
    classunits,
    ID,
    EM.ID,
    rotation = "Varimax",
    plot = FALSE,
    ...
)
```

8 EMMA

#### **Arguments**

Χ	Numeric matrix, input data set with m samples (rows) and n variables (columns).
q	Numeric scalar, number of end-members to be modelled.
1	Numeric scalar or vector, weight transformation limit, i.e. quantile. Set to zero if omitted.
С	Numeric scalar, constant sum scaling parameter, e.g. $1,100,1000$ . Set to $100$ if omitted.
Vqn	Numeric matrix, optional unscaled user-defined end-member loadings. If provided, these are used instead of model-derived ones. See details.
classunits	Numeric vector, optional class units (e.g. micrometers or phi-units) of the same length as columns of $X$ .
ID	Numeric or character vector, optional sample IDs of the same length as rows of $\boldsymbol{X}$ .
EM.ID	Character vector, end-member names. If present, these will be set as rownames of the output data set and used in the legend text.
rotation	Character scalar, rotation type, default is "Varimax". See details.
plot	Logical scalar, optional graphical output of the results, default is FALSE. If set to TRUE, end-member loadings and end-member scores are plotted.
	Additional arguments passed to the plot function. Since the function returns two plots some additional graphical parameters must be specified as vector with the first element for the first plot and the second element for the second plot.

#### **Details**

The parameter Vqn is useful when EMMA shall be performed with a set of prior unscaled endmembers, e.g. from other data sets that are to be used as reference or when modelling a data set with mean end-members, as in the output of robust.loadings.

The rotation type Varimax was used by Dietze et al. (2012). In this R package, one out of the rotations provided by the package GPArotation is possible, as well. However, tests showed that the rotation type has no dramatic consequences for the result.

The function values \$loadings and \$scores are redundant. They are essentially the same as \$Vqsn and \$Mqs. However, they are included for user convenience.

# Value

A list with numeric matrix objects.

loadings Normalised rescaled end-member loadings.

scores Rescaled end-member scores.

Vqn Normalised end-member loadings.

Vqsn Normalised rescaled end-member loadings.

Mqs Rescaled end-member scores.

Xm Modelled data.

modes Mode class of end-member loadings.

EMMA 9

Mqs.var	Explained variance of end-members
Em	Absolute row-wise model error.
En	Absolute column-wise model error.
RMSEm	row-wise root mean square erroe
RMSEn	column-wise root mean square erroe
Rm	Row-wise (sample-wise) explained variance.
Rn	Column-wise (variable-wise) explained variance.
ol	Number of overlapping end-members.

## Author(s)

Michael Dietze, Elisabeth Dietze

#### References

Dietze E, Hartmann K, Diekmann B, IJmker J, Lehmkuhl F, Opitz S, Stauch G, Wuennemann B, Borchers A. 2012. An end-member algorithm for deciphering modern detrital processes from lake sediments of Lake Donggi Cona, NE Tibetan Plateau, China. Sedimentary Geology 243-244: 169-180.

Klovan JE, Imbrie J. 1971. An Algorithm and FORTRAN-IV Program for Large-Scale Q-Mode Factor Analysis and Calculation of Factor Scores. Mathematical Geology 3: 61-77. Miesch AT. 1976. Q-Mode factor analysis of geochemical and petrologic data matrices with constant row sums. U.S. Geological Survey Professional Papers 574.

10 EMrob

**EMpot** 

example data

# Description

A list with output of the function test.robustness()

#### **Format**

```
The format is: List of 8 \ q: num [1:90] 4 \ 4 \ 4 \ 4 \ 4 \ 4 \ 4 \ 4 \ 4 \ \dots \  lw: num [1:90] 0 \ 0 \ 0 \ 0.05 \ 0.05 \ 0.05 \ 0.05 \ 0.1 \ 0.1 \ \dots \  modes: num [1:90] 12 \ 32 \ 61 \ 80 \ 12 \ 32 \ \dots
```

## **Details**

The dataset is the result of the function test.robustness() of the R-package EMMAgeo.

# **Examples**

```
## load example data set
data(example_EMpot)
```

**EMrob** 

example data

## **Description**

Robust end-members, a list with output of the function robust.EM()

## **Format**

The format is: List of 12  $\$  Vqsn.data :List of 4 ... : num [1:15, 1:80] 0.18929 0.184 0.18304 0.00698 0.02033 ...

# **Details**

The dataset is the result of the function robust.EM() of the R-package EMMAgeo.

```
## load example data set
data(example_EMrob)
```

get.l 11

get.l

Generate a vector of weight transformation values from l.min to l.max.

## **Description**

This function generates a sequence of weight transformation values that range from l\_min (by default zero) to l\_max (by default 95 % of the maximum possible value). It uses the function test.l.max().

## Usage

```
get.1(X, n = 10, max = 0.95, min = 0)
```

# **Arguments**

Χ	Numeric matrix, input data set with m samples (rows) and n variables (columns).
n	Numeric scalar, length of the output vector (by default 10).
max	Numeric scalar, fraction of the maximum value (by default 0.95).
min	Numeric scalar, minimum value (by default zero).

## Value

Numeric vector of class "EMMAgeo\_1", weight transformation values.

## Author(s)

Michael Dietze, Elisabeth Dietze

## See Also

```
test.l.max
```

12 get.l.opt

ge	t.	1.	opt
5~	٠.		OPC

Identify optimum weight transformation value

# Description

This function returns for a series of input vaules the weight transformation value, which yielded the highest measure of model quality.

## Usage

```
get.l.opt(X, 1, quality = "mRt", Vqn, rotation, plot = TRUE, ...)
```

# Arguments

Χ	Numeric matrix, input data set with m samples (rows) and n variables (columns).
1	Numeric vector, weight transformation values to test.
quality	Character scalar, quality measure for against which to test the influence of 1. See details for a list of the a vailable keywords. Default is "mRt".
Vqn	Numeric matrix specifying optional unscaled user-defined end-member loadings.
rotation	Character scalar, rotation type, default is "Varimax" (cf. EMMA for further information).
plot	Logical scalar, optional graphical output of the result.
	Further arguments passed to the function.

# **Details**

The parameter quality can be one out of the following keywords: "mRm", "mRn", "mRn", "mEm", "mEn" and "mEt". See EMMA for definition of these keywords.

## Value

Numeric scalar, weight tranformation value with optimal EMMA result.

# Author(s)

Michael Dietze, Elisabeth Dietze

#### See Also

 ${\sf EMMA}$ 

get.limits 13

## **Examples**

get.limits

Infer lower and upper mode position limits to define robust endmembers.

# Description

This function identifies the lower and upper limits within which robust end-members have clustered mode positions. It uses a kernel density estimate of the mode positions of all input end-member loadings, clips it at a user-defined minimum density and returns the resulting rising and falling shoulders of the kde peaks as limits.

#### Usage

```
get.limits(loadings, classunits, bw, threshold = 0.7)
```

## **Arguments**

loadings R object, output of function model.EM.

classunits Numeric vector, optional class units (e.g. micrometers or phi-units) of the same

length as columns of X.

bw Numeric scalar, bandwidth of the kernel, moved over the data set. If omitted,

the default value of 1 used.

threshold Numeric scalar, threshold quantile which is used to identify mode clusters. Only

kde densities above this values are kept and used to derieve mode cluster limits.

## Details

Note that the threshold above which a mode cluster is identified is an arbitrary, user-defined value and probably needs to be adjusted iteratively to get reasonable results. The default value may or may not be adequate!

#### Value

Numeric matrix with lower and upper mode limits.

14 get.q

## Author(s)

Michael Dietze, Elisabeth Dietze

#### See Also

```
EMMA, model.EM
```

## **Examples**

```
## load example data set
data(example_EMpot)

## infer mode cluster limits
limits <- get.limits(loadings = EMpot)</pre>
```

get.q

Generate a parameter matrix with q.min and q.max values for robust EMMA.

# Description

This function uses the input data matrix X and a vector of weight transformation limits to generate a matrix of minimum and maximum likely numbers of end-members to be used to model and extract robust end-members.

# Usage

```
get.q(
    X,
    1 = 0,
    q.min = 2,
    q.max = 10,
    criteria.min = 0.5,
    criteria.max = "local_max",
    correct.output = TRUE,
    ...
)
```

## **Arguments**

Χ	Numeric matrix, input data set with m samples (rows) and n variables (columns).
1	Numeric vector, weight transformation limits, default is zero.
q.min	Numeric scalar, minimum number of end-members to use, default is 2.
q.max	Numeric scalar, maximum number of end-members to use, default is 10.
criteria.min	Numeric scalar, minimum value of explained variance reached to be a valid model realisation, default is 0.5.

get.q

criteria.max Character or numeric scalar, either keyword "local\_max" to use first local maximum or any numeric value of explained variance, default is "local\_max".

correct.output Logical scalar, option to correct the output for twisted values and remove combinations with NA-values. See details.

... Further arguments, passed to the function.

#### **Details**

The parameter q.min should be at least 2 because otherwise the entire dataset would consist of one end-member and there would be no variability at all. The parameter q.max is set to 10 by default, based on practical issues. In natural systems, there are only rarely occasions when such a high number of sediment transport regimes may be preserved in and can be resolved from sedimentary deposits. The parameter 1 should be a vector between the minimum possible (zero) and maximum possible value (by definition the median, 0.5, but usually a lower value). Whensubmitting only a scalar, the variability can be only due to the range of possible endmembers (between q.min and q.max). If the parameter correct.output is enabled, this can decrease the number of valid values for 1, i.e. the number of rows of the output matrix may no longer be the same as the length of the input vector of 1. In such a case the vector 1 must be replaced by the rownames of the output matrix (1 <- as.numeric(rownames(get.q()))).

#### Value

Numeric matrix of class "EMMAgeo\_q", minimum and maximum numbers of end-members as well as corresponding weight transformation values as rownames.

## Author(s)

Michael Dietze, Elisabeth Dietze

#### References

Dietze E, Hartmann K, Diekmann B, IJmker J, Lehmkuhl F, Opitz S, Stauch G, Wuennemann B, Borchers A. 2012. An end-member algorithm for deciphering modern detrital processes from lake sediments of Lake Donggi Cona, NE Tibetan Plateau, China. Sedimentary Geology 243-244: 169-180.

#### See Also

```
EMMA, test.parameters, test.robustness
```

```
## load example data set
data("example_X")

## create parameter matrix
get.q(X = X, 1 = c(0, 0.05, 0.10))
```

16 GUI

GUI

Start GUI for EMMA

# Description

This function starts a browser-based graphic user interface for EMMA. The GUI has so far been tested on a Linux system, both with the browser of RStudio and Mozilla Firefox. It permits basic access to import, display and model a user-provided data set.

## Usage

```
GUI(...)
```

## **Arguments**

... further arguments to pass to function call

## **Details**

To use own data set, this should be a plain ASCII file with samples organised as rows and grain-size classes organised as columns. The ASCII file can be separated by spaces, commas, semi colons or tab stops. The file may contain a leading column with sample IDs and/or one leading row with grain-size class breaks. To run EMMA make sure that there are no classes that contain only zeros throught all samples (i.e., remove them beforehand, e.g., by X = X[, colSums(X) > 0]).

## Author(s)

Michael Dietze

```
## Not run:
# Start the GUI
GUI()
## End(Not run)
```

interpolate.classes 17

interpolate.classes

Interpolate data between different classes.

## **Description**

This function interpolates grain-size data for different classes, either to higher or to lower resolution.

# Usage

```
interpolate.classes(
   X,
   boundaries.in,
   boundaries.out,
   method = "natural",
   fixed.start = TRUE
)
```

## **Arguments**

X Numeric matrix, input data set with m samples (rows) and n variables (columns).

boundaries.in Numeric vector, class boundaries of the input data.

boundaries.out Numeric vector, class boundaries of the output data.

method Logical scalar, interpolation method, one out of "linear" (linear interpolation),

"fmm" (cubic spline), "natural" (natural spline), "periodic" (periodic spline).

Default is "natural".

fixed.start Logocal scalar, specifying if the outer boundaries should be set to the same

values as in the original matrix, default is TRUE. This may become necessary to

avoid interpolation errors, see example.

## Value

Numeric matrix, interpolated class values.

#### Author(s)

Michael Dietze, Elisabeth Dietze

## See Also

EMMA, approx, spline

18 mix.EM

## **Examples**

```
## load example data
data(example_X)
classes.in \leftarrow seq(from = 1, to = 10, length.out = ncol(X))
## Example 1 - decrease the class numbers
## define number of output classes
classes.out <- seq(1, 10, length.out = 20)
## interpolate the data set
Y <- interpolate.classes(X = X,
                         boundaries.in = classes.in,
                         boundaries.out = classes.out,
                         method = "linear")
## show original vs. interpolation for first 10 samples
plot(NA, xlim = c(1, 10), ylim = c(0, 40))
for(i in 1:10) {
  lines(classes.in, X[i,] * 20 + i)
  lines(classes.out, Y[i,] * 20 + i, col = 2)
## Example 2 - increase the class numbers
## define number of output classes
classes.out <- seq(1, 10, length.out = 200)</pre>
## interpolate the data set
Y <- interpolate.classes(X = X,
                         boundaries.in = classes.in,
                         boundaries.out = classes.out)
## show original vs. interpolation for first 10 samples
plot(NA, xlim = c(1, 10), ylim = c(0, 40))
for(i in 1:10) {
  lines(classes.in, X[i,] * 20 + i)
  lines(classes.out, Y[i,] * 20 + i, col = 2)
}
```

mix.EM

Function to mix sample spectres.

## Description

This functions allows to mix grain-size distributions with specified proportions and defined noise levels, for example to test the goodness of the EMMA algorithm.

## Usage

```
mix.EM(EM, proportion, noise, autocorrelation)
```

mix.EM

## Arguments

EM Numeric matrix, grain-size distribution definitions.

Each definition is in a separate row with variable contributions in columns.

proportion Numeric vector, relative proportions of each distribution per sample.

noise Numeric scalar, optional relative white noise levels.

autocorrelation

Numeric scalar, the degree of autocorrelation among classes. Autocorrelation is realised as running mean of the specified length. Only odd values are allowed.

#### **Details**

The function multiplies each end-member with the respective proportion value, sums the resulting variables, adds uniform noise and normalises the resulting mixed sample to 100 %.

#### Value

Numeric vector, a sample composed of known proportions of end-members.

## Author(s)

Michael Dietze, Elisabeth Dietze

#### See Also

create.EM

```
## define end-member loadings and phi vector
EMa.1 <- create.EM(p1 = c(2, 8), p2 = c(1, 0.8), s = c(0.7, 0.3),
                   boundaries = c(0, 11), n = 80)
EMa.2 <- create.EM(p1 = c(4, 7), p2 = c(1.1, 1.4), s = c(0.5, 0.5),
                   boundaries = c(0, 11), n = 80)
EMa
      <- rbind(EMa.1, EMa.2)
      <- seq(0, 11, length.out = 80)
phi
## mix end-member loadings
sample1 <- mix.EM(EMa, proportion = c(0.3, 0.7))
sample2 <- mix.EM(EMa, proportion = c(0.5, 0.5), noise = 0.1,
                  autocorrelation = 3)
## plot end-member loadings (grey) and resulting samples (black)
plot(phi, EMa.1, type="l", col = "grey")
lines(phi, EMa.2, col = "grey")
lines(phi, sample1)
lines(phi, sample2)
```

20 model.EM

model.EM

Model all possible end-member scenarios

#### **Description**

This function takes a definition of weight transformation limits and corresponding minimum and maximum numbers of end-members to model all end-member scenarios in accordance with these parameters. Based on the output the user can decide on robust end-members.

## Usage

```
model.EM(X, q, l, classunits, plot = TRUE, col.q = TRUE, bw, ...)
```

#### **Arguments**

Χ	Numeric matrix, input data set with m samples (rows) and n variables (columns).
q	Numeric matrix, definitions of minimum and maximum number of end-members (cf. get.q()), required.
1	Numeric vector, weight transformation limit values, corresponding to the matrix q, required.
classunits	Numeric vector, optional class units (e.g. micrometers or phi-units) of the same length as columns of X.
plot	Logical scalar, option to plot the results (cf. details for explanations), default is TRUE.
col.q	Logical scalar, option to colour end-member loadings by the number of end-members which were used to create the model realisation, default is TRUE.
bw	Numeric scalar, optional manual setting of the kde bandwidth. By default, bw is calculated as 1 percent of the number of grain-size classes.
	Further arguments passed to the function.

## **Details**

The plot output is an overlay of several data. The coloured lines in the background are end-member loadings (number noted in the plot title), resulting from all possible model scenarios. If col.q == TRUE they are coloured according to the number of end-members with which the model was generated. This colour scheme allows to depict end-members that emerge for model realisations with specific number of end-members. The thick black line is a kernel density estimate curve, generated from the mode positions of all end-members. The kernel bandwidth is set to 1 percent of the number of grain-size classes of the input data set, which gave useful results for most of our test data sets. The cumulaitve dot-line-plot is a further visualisation of end-member mode positions. The function is a modified wrapper function for the function test.robustness().

#### Value

List object with all modelled end-members, each described by input parameters, mode position, quality measures and value distributions.

residual.EM 21

## Author(s)

Michael Dietze, Elisabeth Dietze

#### References

Dietze E, Hartmann K, Diekmann B, IJmker J, Lehmkuhl F, Opitz S, Stauch G, Wuennemann B, Borchers A. 2012. An end-member algorithm for deciphering modern detrital processes from lake sediments of Lake Donggi Cona, NE Tibetan Plateau, China. Sedimentary Geology 243-244: 169-180.

#### See Also

```
EMMA, test.1.max
```

## **Examples**

```
## load example data set
data(example_X)

## define input parameters
1 <- c(0, 0.05, 0.10)
q <- cbind(c(2, 2, 3), c(5, 6, 4))

## infer l-vector
em_pot <- model.EM(X = X, q = q, l = l)</pre>
```

residual.EM

Calculate a residual end-member loading.

## **Description**

This function calculates an optional residual end-member loading. It uses the modelled end-member loadings as input and evaluates the root of 1 minus the sum of all squared loadings. The residual end-member can be used to analyse the remaining variance, e.g. if not all (robust) EMs are included (cf. Dietze et al., 2012). Negative values are set to zero.

## Usage

```
residual.EM(Vqn)
```

## **Arguments**

Vqn

Numeric matrix, m unscaled robust end-member loadings.

# Value

Numeric vector, residual end-member loading.

22 robust.EM

## Author(s)

Michael Dietze, Elisabeth Dietze

## References

Dietze E, Hartmann K, Diekmann B, IJmker J, Lehmkuhl F, Opitz S, Stauch G, Wuennemann B, Borchers A. 2012. An end-member algorithm for deciphering modern detrital processes from lake sediments of Lake Donggi Cona, NE Tibetan Plateau, China. Sedimentary Geology 243-244: 169-180.

#### See Also

```
EMMA, robust.EM
```

## **Examples**

robust.EM

Extract robust end-members

# Description

This function takes a list object with potential end-member loadings and extracts those with modes in specified limits to describe them by mean and standard deviation and use these descriptions to propagate the uncertainties to end-member scores.

## Usage

```
robust.EM(
  em,
  limits,
  classunits,
```

robust.EM 23

```
amount,
1,
    mc_n,
    type = "mean",
    qt = c(0.25, 0.75),
    cores = 1,
    plot = FALSE,
    ...
)
```

## **Arguments**

em	List of class "EMMAgeo_empot", i.e. the outout of model.em() or test.robustness(), containing potential end-members, both in unscaled and rescaled version as well as further parameters.
limits	Numeric matrix with two columns, defining the class limits for the robust end- members to calculate. The first column defines the lower limits, the second column the upper limits. End-members are organised in rows.
classunits	Numeric vector, optional class units (e.g. micrometers or phi-units) of the same length as columns of X.
amount	Numeric matrix with two columns, defining the minimum and maximum amount of the modal class for each end-member.
1	Numeric scalar, weight transformation limit for modelling the average end- member output.
mc_n	Numeric scalar, number of Monte Carlo simulations to estimate end-member scores uncertainty. The default setting is ten times the product of number of end-members and number of weight transformation limits. The latter is inherited from model.em(). To disable modelling of scores uncertainty, set mc_n = 0.
type	Character scalar, type of oadings statistics. One out of "mean" and "median". Default is "mean".
qt	Numeric vector of length two, quantiles to describe end-member loadings. Default is c(0.25, 0.75) (i.e., the quartile range).
cores	Numeric scalar, number of CPU cores to be used for calculations. Only useful in multicore architectures. Default is 1 (single core).
plot	Logical scalar, option for plot output. Default is FALSE.
•••	Additional arguments passed to EMMA and plot.

## **Details**

The function is used to extract potential end-member loadings based on their mode positions and, optionally the height of the mode class, and use them to infer mean and stanard deviation of all end-members that match the group criteria defined by limits. These information are then used to model the uncertainty of the corresponding end-member scores. The function uses input from two preceding approaches. In a compact protocol model.em delivers these data in a predefined way. In the extended protocol test.robustness does this.

24 robust.loadings

## Value

List with statistic descriptions of end-member loadings and scores.

## Author(s)

Michael Dietze, Elisabeth Dietze

## See Also

```
robust.loadings, robust.scores
```

```
## Not run:
## load example data set
data(example_X)
## get weight transformation limit vector
1 \leftarrow get.1(X = X)
## get minimum and maximum number of end-members
q \leftarrow get.q(X = X, 1 = 1)
## get all potential model scenarios
EM_pot \leftarrow model.EM(X = X, q = q, plot = TRUE)
## define end-member mode class limits
limits <- cbind(c(61, 74, 95, 102),
                 c(64, 76, 100, 105))
## get robust end-members in the default way, with plot output
rem <- robust.EM(em = EM_pot,</pre>
                  limits = limits,
                  plot = TRUE)
## get robust end-members by only modelling uncertainty in loadings
robust_EM <- robust.EM(em = EM_pot,</pre>
                        limits = limits,
                        plot = TRUE)
## End(Not run)
```

robust.loadings 25

# Description

This function takes a list object with potential end-member loadings and extracts those with modes in specified limits to describe them by mean and standard deviation.

# Usage

```
robust.loadings(
  em,
  limits,
  classunits,
  amount,
  type = "mean",
  qt = c(0.25, 0.75),
  plot = FALSE,
  ...
)
```

# Arguments

em	List of class "EMMAgeo_empot", i.e. the outout of model.EM() or test.robustness(), containing potential end-members, both in unscaled and rescaled version as well as further parameters.
limits	Numeric matrix with two columns, defining the class limits for the robust end- members to calculate. The first column defines the lower limits, the second column the upper limits. End-members are organised in rows.
classunits	Numeric vector, optional class units (e.g. micrometers or phi-units) of the same length as the number of (grain-size) classes per sample.
amount	Numeric matrix with two columns, defining the minimum and maximum amount of the modal class for each end-member.
type	Character scalar, type of statistics. One out of "mean" and "median". Default is "mean".
qt	Numeric vector of length two, quantiles to describe end-member loadings Default is $c(0.25, 0.75)$ (i.e., the quartile range).
plot	Logical scalar, option to enable plot output. Default is FALSE.
	Additional arguments passed to EMMA and plot.

## Value

List with statistic descriptions of unscaled and scaled end-member loadings.

# Author(s)

Michael Dietze, Elisabeth Dietze

# See Also

```
robust.EM, robust.scores
```

26 robust.scores

## **Examples**

robust.scores

Extract robust end-member scores.

## **Description**

This function takes a list object with statistics of end-member loadings and propagates these uncertainties to end-member scores using Monte Carlo methods.

## Usage

```
robust.scores(loadings, 1, mc_n, cores = 1, plot = FALSE, ...)
```

## **Arguments**

loadings	List of class "EMMAgeo_robload", i.e. the outout of robust.loadings(), containing statistic descriptions of robust end-member loadings.
1	Numeric scalar, weight transformation limit to use for modelling the average end-member output. Can be output of get.l.opt(). If omitted, it is set to 0.
mc_n	Numeric scalar, number of Monte Carlo simulations to estimate end-member scores uncertainty. The default setting is ten times the product of number of end-members and number of weight transformation limits. The latter is inherited from model.em(). To disable modelling of scores uncertainty, set $mc_n = 0$ .
cores	Numeric scalar, number of CPU cores to be used for calculations. Only useful in multicore architectures. Default is 1 (single core).
plot	Logical scalar, option for plot output. Default is FALSE.
	Additional arguments passed to EMMA and plot.

#### Value

List with statistic descriptions of robust end-member scores.

test.factors 27

## Author(s)

Michael Dietze, Elisabeth Dietze

#### See Also

```
robust.EM, robust.loadings
```

## **Examples**

test.factors

Calculate the initial cumulative explained variance of factors.

# Description

This function performs eigenspace decomposition using the weight-transformed matrix W to determine the explained variance with increasing number of factors. Depending on the number of provided weight transformation limits (1) a vector or a matrix is returned.

## Usage

```
test.factors(X, 1, c, r.min = 0.95, plot = FALSE, legend, ...)
```

# Arguments

Χ	Numeric matrix, input data set with m samples (rows) and n variables (columns).
1	Numeric vector, weight tranformation limits, i.e. quantiles; default is 0.
С	Numeric scalar, constant sum scaling parameter, e.g. $1$ , $100$ , $1000$ ; default is $100$ .
r.min	Numeric scalar, minimum value of explained variance to be reached by the end-members included, default is $0.95$ .
plot	Logical scalar, optional graphical output of the results, default is FALSE.

28 test.factors

legend	Character scalar, specify legend position (cf. legend). If omitted, no legend will be plotted, default is no legend.
	Additional arguments passed to the plot function. Use colour instead of col to create different colours.

## **Details**

The results may be used to define a minimum number of end-members for subsequent modelling steps, e.g. by using the Kaiser criterion, which demands a minimum number of eigenvalues to reach a squared R of 0.95.

#### Value

List with objects

L Vector or matrix of cumulative explained variance.

q.min Vector with number of factors that passed r.min.

## Author(s)

Michael Dietze, Elisabeth Dietze

#### References

Dietze E, Hartmann K, Diekmann B, IJmker J, Lehmkuhl F, Opitz S, Stauch G, Wuennemann B, Borchers A. 2012. An end-member algorithm for deciphering modern detrital processes from lake sediments of Lake Donggi Cona, NE Tibetan Plateau, China. Sedimentary Geology 243-244: 169-180.

test.1 29

test.1

Test a vector of weight transformation limits for mximum value.

## **Description**

This function performs the weight transformation of the data matrix after Klovan & Imbrie (1971) and performs EMMA() with different weight limits to check if valied results are yielded. It returns the maximum value for which the transformation remains stable.

## Usage

```
test.1(X, 1, ...)
```

#### **Arguments**

X Numeric matrix, input data set with m samples (rows) and n variables (columns).

1 Numeric vector, weight transformation limit, i.e. quantile; default is 0.

... Further arguments passed to the function.

## Value

List with objects

step Numeric scalar with position of the last valid value.

1.max Numeric scalar with last valid value of 1.

#### Author(s)

Michael Dietze, Elisabeth Dietze

## References

Dietze E, Hartmann K, Diekmann B, IJmker J, Lehmkuhl F, Opitz S, Stauch G, Wuennemann B, Borchers A. 2012. An end-member algorithm for deciphering modern detrital processes from lake sediments of Lake Donggi Cona, NE Tibetan Plateau, China. Sedimentary Geology 243-244: 169-180

Klovan JE, Imbrie J. 1971. An Algorithm and FORTRAN-IV Program for Large-Scale Q-Mode Factor Analysis and Calculation of Factor Scores. Mathematical Geology 3: 61-77.

#### See Also

EMMA, check.data, test.parameters

30 test.l.max

## **Examples**

```
## load example data set
data(example_X)

test <- test.l(X = X, l = seq(from = 0, to = 0.6, by = 0.1))</pre>
```

test.1.max

Find maximum possible wight transformation value.

## **Description**

This function approximates the highest possible value for 1 in a nested loop. It uses test.1 and does not need any further parameters. It starts with 1 between zero and 0.5 and iteratively approximates the last possible values for which the weight-transformed matrix of the input data still allows eigenspace extraction.

## Usage

```
test.l.max(X, n = 10, \ldots)
```

## **Arguments**

X Numeric matrix, input data set with m samples (rows) and n variables (columns).

n Numeric scalar, number of loop runs and values per loop.

Further arguments passed to the function.

## Value

Numeric scalar, maximal possible I value.

## Author(s)

Michael Dietze, Elisabeth Dietze

# References

Dietze E, Hartmann K, Diekmann B, IJmker J, Lehmkuhl F, Opitz S, Stauch G, Wuennemann B, Borchers A. 2012. An end-member algorithm for deciphering modern detrital processes from lake sediments of Lake Donggi Cona, NE Tibetan Plateau, China. Sedimentary Geology 243-244: 169-180.

Klovan JE, Imbrie J. 1971. An Algorithm and FORTRAN-IV Program for Large-Scale Q-Mode Factor Analysis and Calculation of Factor Scores. Mathematical Geology 3: 61-77.

## See Also

```
EMMA, test.1
```

test.parameters 31

# **Examples**

```
## load example data set
data(example_X)

## create weight transformation limits vector
1 <- seq(from = 0, to = 0.6, by = 0.05)

## test l.max (uncomment to run, may take more than 10 sec to run)
# l.max <- test.l.max(X = X)</pre>
```

test.parameters

Evaluate influence of model parameters.

# Description

All possible combinations of number of end-members and weight transformation limits are used to perform EMMA and evaluate the absolute and relative measures of individual model performance.

# Usage

```
test.parameters(
   X,
   q,
   l = 0,
   c = 100,
   rotation = "Varimax",
   plot = FALSE,
   legend,
   multicore = FALSE,
   ...
)
```

# Arguments

Χ	Numeric matrix, input data set with m samples (rows) and n variables (columns).
q	Numeric vector, numbers of end-members to be modelled, e.g., 2:10.
1	Numeric vector specifying the weight tranformation limit, i.e. quantile; default is 0.
С	Numeric scalar specifying the constant sum scaling parameter, e.g., $1, 100, 1000$ ; default is $0.$
rotation	Character scalar, rotation type, default is "Varimax".
plot	Character scalar, optional graphical output of the results as keyword (see details). All plots except "ol" are colour-coded bitmaps of q, l and the specified test parameter and line-plots the specified parameter vs. q.

32 test.parameters

legend Character scalar, specifying legend position (cf. legend). If omitted, no legend

will be plotted, default is no legend.

multicore Logical scalar, optionally ditribute calculations to all available cores of the

computer, default is TRUE.

. . . Additional arguments passed to the plot function (see details).

#### **Details**

The mean total explained variance mRt may be used to define a maximum number of meaningful end-members for subsequent modelling, e.g. as the number of end-members, which reaches the first local mRt maximum.

Overlapping is defined as one end-member having its mode within the "area" of any other endmember, which is genetically not explainable.

Keywords to specify, which tested parameter will be plotted: "mEm" (mean absolute row-wise error), "mEn" (mean absolute column-wise error), "mRm" (mean relative row-wise error), "mRn" (mean relative column-wise error), "mRt" (mean relative total error) and "ol" (number of overlapping end-members).

Since the function returns two plots (except for option "ol"), additional graphical parameters must be specified as vector with the first element for the first plot and the second element for the second plot. If graphical parameters are natively vectors (e.g. a sequence of colours), they must be specified as matrices with each vector as a row. A legend can only be added to the second plot. Colours only apply to the second plot as well. If colours are specified, colour should be used instead of col. See example section for further advice.

#### Value

List with result objects

mEm Absolute row-wise model error.

mEn Absolute column-wise model error.

mRm Mean row-wise explained variance.

mRn Mean column-wise explained variance.

mRt Mean total explained variance.

ol Number of overlapping end-member loadings.
q.max Maximum number of meaningful end-members.

## Author(s)

Michael Dietze, Elisabeth Dietze

#### References

Dietze E, Hartmann K, Diekmann B, IJmker J, Lehmkuhl F, Opitz S, Stauch G, Wuennemann B, Borchers A. 2012. An end-member algorithm for deciphering modern detrital processes from lake sediments of Lake Donggi Cona, NE Tibetan Plateau, China. Sedimentary Geology 243-244: 169-180.

test.robustness 33

## See Also

**EMMA** 

## **Examples**

test.robustness

Test model robustness.

## Description

This function takes a definition of weight transformation limits and corresponding minimum and maximum numbers of end-members to model all end-member scenarios in accordance with these parameters. Based on the output the user can decide on robust end-members.

# Usage

```
test.robustness(
   X,
   q,
   1,
   P,
   c,
   classunits,
   ID,
   rotation = "Varimax",
   ol.rej,
   mRt.rej,
   plot = FALSE,
   ...
)
```

34 test.robustness

## **Arguments**

<ul> <li>Numeric vector with number of end-members to be modelled.</li> <li>Numeric vector specifying the weight tranformation limits, i.e. quantiles; default is 0.</li> <li>Numeric matrix, optional alternative input parameters for q and 1, either of the form m:3 with m variations in the columns q.min, q.max, 1 or of the form m:2</li> </ul>
fault is 0.  P Numeric matrix, optional alternative input parameters for q and l, either of the
with m variations in the columns q, l.
c Numeric scalar specifying the constant sum scaling parameter, e.g. 1, 100, 1000; default is 100.
classunits Numeric vector, optional class units (e.g. phi classes or micrometers) of the same length as columns of $\boldsymbol{X}$ .
Numeric or character vector, optional sample IDs of the same length as columns of $\mathbf{X}$ .
rotation Character scalar, rotation type, default is "Varimax" (cf. Dietze et al., 2012). One out of the rotations provided in GPArotation is possible.
ol.rej Numeric scalar, optional rejection threshold for overlapping criterion. All model runs with overlapping end-members greater than the specified integer will be removed.
MRt.rej Numeric scalar, optional rejection threshold for mean total explained variance criterion. All modelled end-members below the specified value will be removed.
plot Logical scalar, optional graphical output of the results, default is FALSE. If set to TRUE, end-member loadings and end-member scores are plotted.
Additional arguments passed to the plot function (see details).

## **Details**

The function value \$loadings is redundant but was added for user convenience.

Since the function returns two plots, additional graphical parameters must be specified as vector with the first element for the first plot and the second element for the second plot. If graphical parameters are natively vectors (e.g. a sequence of colours), they must be specified as matrices with each vector as a row. If colours are specified, colour should be used instead of col. ylim can only be modified for the first plot. See example section for further advice.

## Value

## A list with objects

q	Vector with q.
1	Vector with 1.

modes Vector with mode class.

mRt Vector with mean total explained variance.
ol Vector with n overlapping end-members.

loadings Matrix with normalised rescaled end-member loadings.

Vqsn Matrix with rescaled end-member loadings. Vqn Matrix with normalised factor loadings. test.robustness 35

#### Author(s)

Michael Dietze, Elisabeth Dietze

#### References

Dietze E, Hartmann K, Diekmann B, IJmker J, Lehmkuhl F, Opitz S, Stauch G, Wuennemann B, Borchers A. 2012. An end-member algorithm for deciphering modern detrital processes from lake sediments of Lake Donggi Cona, NE Tibetan Plateau, China. Sedimentary Geology 243-244: 169-180.

```
## load example data set
data(example_X)
## Example 1 - perform the most simple test
q <- 4:7
1 < - seq(from = 0, to = 0.1, by = 0.05)
M1 <- test.robustness(X = X, q = q, l = l,
                       ol.rej = 1, mRt.rej = 0.8,
                        plot = TRUE,
                        colour = c(4, 7),
                        xlab = c(expression(paste("Grain size (", phi, ")",
                                                   sep = "")),
                                 expression(paste("Grain size (", phi, ")",
                                                   sep = ""))))
## Example 2 - perform the test without rejection criteria and plots
P <- cbind(rep(q[1], length(l)),</pre>
            rep(q[3], length(1)),
            1)
M2 \leftarrow test.robustness(X = X, P = P)
## Plot 1 - \text{end-member loadings} which do not overlap and yielded mRt > 0.80.
plot(M2$Vqsn[1,], type = "l", ylim = c(0, max(M2$Vqsn, na.rm = TRUE)),
     main = "End-member loadings")
  for (i in 2:nrow(M2$Vqsn)) lines(M2$Vqsn[i,])
# Plot 2 - histogram of mode positions
hist(M2$modes,
     breaks = 1:ncol(X),
     main = "Mode positions",
     xlab = "Class")
# Plot 3 - positions of modelled end-member modes by number of end-members
# Note how scatter in end-member position decreases for the "correct" number
# of modelled end-members (6) and an appropriate weight limit (ca. 0.1).
ii <- order(M2$q, M2$modes)</pre>
modes <- t(rbind(M2$modes, M2$q))[ii,]</pre>
plot(modes[,1],
     seq(1, nrow(modes)),
```

36 X

```
main = "Model overview",
    xlab = "Class",
    ylab = "EM number in model run",
    pch = as.character(modes[,2]),
    cex = 0.7)

# Illustrate mode positions as stem-and-leave-plot, useful as a simple
# check, which mode maxima are consistently fall into which grain-size
# class (useful to define "limits" in robust.EM).
stem(M2$modes, scale = 2)
```

Χ

example data

## **Description**

Synthetic data set created by randomly mixed natural end-members

## **Format**

```
num [1:100, 1:116] 0.000899 0.000516 0.00136 0.000989 0.00102 ...
```

## **Details**

The dataset is the result of four mixed natural end-members.

X 37

col = rainbow(n = 250))

# **Index**

* EMMA  check.data, 3  click.limits, 4  convert.units, 5  create.EM, 6	get.1, 11 get.1.opt, 12 get.1imits, 13 get.q, 14 GUI, 16
EMMA, 7 get.1, 11 get.1.opt, 12 get.limits, 13 get.q, 14 interpolate.classes, 17 mix.EM, 18 model.EM, 20 residual.EM, 21	interpolate.classes, 17  mix.EM, 18 model.EM, 20  residual.EM, 21 robust.EM, 22 robust.loadings, 24 robust.scores, 26
robust.EM, 22 robust.loadings, 24 robust.scores, 26 test.factors, 27 test.1, 29 test.1.max, 30 test.parameters, 31 test.robustness, 33 * datasets EMpot, 10 EMrob, 10 X, 36 * package EMMAgeo-package, 2	test.factors, 27 test.1, 29 test.1.max, 30 test.parameters, 31 test.robustness, 33 X, 36
check.data, 3 click.limits, 4 convert.units, 5 create.EM, 6  EMMA, 7 EMMAgeo (EMMAgeo-package), 2 EMMAgeo-package, 2 EMpot, 10 EMrob, 10	