

Package ‘Compositional’

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mixture, nnet, parallel, RANN, Rfast, Rfast2, sn, stats

Description Regression, classification, contour plots, hypothesis testing and fitting of distributions for compositional data are some of the functions included. The standard textbook for such data is John Aitchison's (1986) "The statistical analysis of compositional data". Relevant papers include a) Tsagris M.T., Preston S. and Wood A.T.A. (2011) A data-based power transformation for compositional data. Fourth International International Workshop on Compositional Data Analysis. b) Tsagris M. (2014). The k-NN algorithm for compositional data: a revised approach with and without zero values present. *Journal of Data Science*, 12(3):519--534. c) Tsagris M. (2015). A novel, divergence based, regression for compositional data. *Proceedings of the 28th Panhellenic Statistics Conference*, 15-18 April 2015, Athens, Greece, 430--444. d) Tsagris M. (2015). Regression analysis with compositional data containing zero values. *Chilean Journal of Statistics*, 6(2):47--57. e) Tsagris M., Preston S. and Wood A.T.A. (2016). Improved supervised classification for compositional data using the alpha-transformation. *Journal of Classification*, 33(2):243--261. <doi:10.1007/s00357-016-9207-5>. f) Tsagris M., Preston S. and Wood A.T.A. (2017). Nonparametric hypothesis testing for equality of means on the simplex. *Journal of Statistical Computation and Simulation*, 87(2): 406--422. <doi:10.1080/00949655.2016.1216554> g) Tsagris M. and Stewart C. (2018). A Dirichlet regression model for compositional data with zeros. *Lobachevskii Journal of Mathematics*, 39(3): 398--412. <doi:10.1134/S1995080218030198>. h) Alenazi A. (2019). Regression for compositional data with compositional data as predictor variables with or without zero values. *Journal of Data Science*, 17(1): 219--238. <doi:10.6339/JDS.201901_17(1).0010>. i) Tsagris M. and Stewart C. (2019). A Dirichlet regression model for compositional data with zeros. *Lobachevskii Journal of Mathematics*, 40(3): 398--412. <doi:10.1134/S1995080219030198>.

art C. (2020). A folded model for compositional data analysis. Australian and New Zealand Journal of Statistics, 62(2):249--277. <doi:10.1111/anzs.12289>. j) Tsagris M., Alenazi A. and Stewart C. (2020). The alpha-k-NN regression for compositional data. <arXiv:2002.05137>. We further include functions for percentages (or proportions).

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Compositional-package *Compositional Data Analysis*

Description

A collection of functions for compositional data analysis.

Details

Package: Compositional
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Maintainers

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Note

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Author(s)

Michail Tsagris <mtsagris@uoc.gr>, Giorgos Athineou <gioathineou@gmail.com> and Abdulaziz Alenazi <a.alenazi@nbu.edu.sa>.

References

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

Aithison's simple zero replacement strategy
Aithison's simple zero replacement strategy

Description

Aithison's simple zero replacement strategy.

Usage

```
zeroreplace(x, a = 2/3)
```

Arguments

x	A matrix with the compositional data.
a	The replacement value will be "a" times the minimum value observed in the compositional data.

Details

This is the simple zero replacement strategy suggested in Aitchison (1986, pg. 269).

Value

A matrix with the zero replaced compositional data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[perturbation, alfa](#)

Examples

```
x <- as.matrix(iris[1:20, 1:4])
x <- x/ rowSums(x)
x[ sample(1:20, 4), sample(1:4, 1) ] <- 0
zeroreplace(x)
```

All pairwise additive log-ratio transformations

All pairwise additive log-ratio transformations

Description

All pairwise additive log-ratio transformations.

Usage

```
alr.all(x)
```

Arguments

x A numerical matrix with the compositional data.

Details

The additive log-ratio transformation with the first component being the common divisor is applied. Then all the other pairwise log-ratios are computed and added next to each column. For example, divide by the first component, then divide by the second component and so on.

Value

A matrix with all pairwise alr transformed data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[alr](#), `\link{alfa}`

Examples

```
x <- as.matrix(iris[, 2:4])
x <- x / rowSums(x)
y <- alr.all(x)
```

Beta regression

Beta regression

Description

Beta regression.

Usage

```
beta.reg(y, x, xnew = NULL)
```

Arguments

y	The response variable. It must be a numerical vector with proportions excluding 0 and 1.
x	The independent variable(s). It can be a vector, a matrix or a dataframe with continuous only variables, a data frame with mixed or only categorical variables.
xnew	If you have new values for the predictor variables (dataset) whose response values you want to predict insert them here.

Details

Beta regression is fitted.

Value

A list including:

phi	The estimated precision parameter.
info	A matrix with the estimated regression parameters, their standard errors, Wald statistics and associated p-values.
loglik	The log-likelihood of the regression model.
est	The estimated values if xnew is not NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

Ferrari S.L.P. and Cribari-Neto F. (2004). Beta Regression for Modelling Rates and Proportions. Journal of Applied Statistics, 31(7): 799-815.

See Also

[beta.est](#), [prop.reg](#), [diri.reg](#)

Examples

```
y <- rbeta(300, 3, 5)
x <- matrix( rnorm(300 * 2), ncol = 2)
beta.reg(y, x)
```

Contour plot of a Dirichlet distribution in S^2

Contour plot of a Dirichlet distribution in S^2

Description

Contour plot of a Dirichlet distribution in S^2 .

Usage

```
diri.contour(a, n = 100, x = NULL)
```

Arguments

a	A vector with three elements corresponding to the 3 (estimated) parameters.
n	The number of grid points to consider over which the density is calculated.
x	This is either NULL (no data) or contains a 3 column matrix with compositional data.

Details

The user can plot only the contour lines of a Dirichlet with a given vector of parameters, or can also add the relevant data should he/she wish to.

Value

A ternary diagram with the points and the Dirichlet contour lines.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Ng Kai Wang, Guo-Liang Tian and Man-Lai Tang (2011). Dirichlet and related distributions: Theory, methods and applications. John Wiley & Sons.

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[norm.contour](#), [bivt.contour](#), [comp.kerncontour](#), [mixnorm.contour](#)

Examples

```
x <- as.matrix( iris[, 1:3] )
x <- x / rowSums(x)
diri.contour( a = c(3, 4, 2) )
```

Contour plot of a Gaussian mixture model in S^2

Contour plot of a Gaussian mixture model in S^2

Description

Contour plot of a Gaussian mixture model in S^2 .

Usage

```
mixnorm.contour(x, mod)
```

Arguments

x	A matrix with the compositional data.
mod	An object containing the output of a mix.compnorm model.

Details

The contour plot of a Gaussian mixture model is plotted. For this you need the data and the fitted model.

Value

A ternary plot with the data and the contour lines of the fitted Gaussian mixture model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Ryan P. Browne, Aisha ElSherbiny and Paul D. McNicholas (2015). R package mixture: Mixture Models for Clustering and Classification

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[mix.compnorm](#), [bic.mixcompnorm](#), [diri.contour](#)

Examples

```
## Not run:
x <- as.matrix(iris[, 1:3])
x <- x / rowSums(x)
mod <- mix.compnorm(x, 3, model = "EII")
mixnorm.contour(x, mod)

## End(Not run)
```

Contour plot of the kernel density estimate in S^2

Contour plot of the kernel density estimate in S^2

Description

Contour plot of the kernel density estimate in S^2 .

Usage

```
comp.kerncontour(x, type = "alr", n = 100)
```

Arguments

x	A matrix with the compositional data. It has to be a 3 column matrix.
type	This is either "alr" or "ilr", corresponding to the additive and the isometric log-ratio transformation respectively.
n	The number of grid points to consider, over which the density is calculated.

Details

The alr or the ilr transformation are applied to the compositional data. Then, the optimal bandwidth using maximum likelihood cross-validation is chosen. The multivariate normal kernel density is calculated for a grid of points. Those points are the points on the 2-dimensional simplex. Finally the contours are plotted.

Value

A ternary diagram with the points and the kernel contour lines.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

M.P. Wand and M.C. Jones (1995). Kernel smoothing, CrC Press.

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[diri.contour](#), [mixnorm.contour](#), [bivt.contour](#), [norm.contour](#)

Examples

```
x <- as.matrix(iris[, 1:3])
x <- x / rowSums(x)
comp.kerncontour(x, type = "alr", n = 20)
comp.kerncontour(x, type = "ilr", n = 20)
```

Contour plot of the normal distribution in S^2

Contour plot of the normal distribution in S^2

Description

Contour plot of the normal distribution in S^2 .

Usage

```
norm.contour(x, type = "alr", n = 100, appear = TRUE)
```

Arguments

x	A matrix with the compositional data. It has to be a 3 column matrix.
type	This is either "alr" or "ilr", corresponding to the additive and the isometric log-ratio transformation respectively.
n	The number of grid points to consider over which the density is calculated.
appear	Should the available data appear on the ternary plot (TRUE) or not (FALSE)?

Details

The alr or the ilr transformation is applied to the compositional data at first. Then for a grid of points within the 2-dimensional simplex the bivariate normal density is calculated and the contours are plotted along with the points.

Value

A ternary diagram with the points (if appear = TRUE) and the bivariate normal contour lines.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[diri.contour](#), [mixnorm.contour](#), [bivt.contour](#), [skewnorm.contour](#)

Examples

```
x <- as.matrix(iris[, 1:3])
x <- x / rowSums(x)
norm.contour(x)
norm.contour(x, type = "ilr")
```

Contour plot of the skew skew-normal distribution in S^2

Contour plot of the skew skew-normal distribution in S^2

Description

Contour plot of the skew skew-normal distribution in S^2 .

Usage

```
skewnorm.contour(x, type = "alr", n = 100, appear = TRUE)
```

Arguments

x	A matrix with the compositional data. It has to be a 3 column matrix.
type	This is either "alr" or "ilr", corresponding to the additive and the isometric log-ratio transformation respectively.
n	The number of grid points to consider over which the density is calculated.
appear	Should the available data appear on the ternary plot (TRUE) or not (FALSE)?

Details

The alr or the ilr transformation is applied to the compositional data at first. Then for a grid of points within the 2-dimensional simplex the bivariate skew skew-normal density is calculated and the contours are plotted along with the points.

Value

A ternary diagram with the points (if appear = TRUE) and the bivariate skew skew-normal contour lines.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Azzalini A. and Valle A. D. (1996). The multivariate skew-skewnormal distribution. *Biometrika* 83(4):715-726.

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[diri.contour](#), [mixnorm.contour](#), [bivt.contour](#), [norm.contour](#)

Examples

```
x <- as.matrix(iris[51:100, 1:3])
x <- x / rowSums(x)
skewnorm.contour(x)
```

Contour plot of the t distribution in S^2

Contour plot of the t distribution in S^2

Description

Contour plot of the t distribution in S^2 .

Usage

```
bivt.contour(x, type = "alr", n = 100, appear = TRUE)
```

Arguments

<code>x</code>	A matrix with compositional data. It has to be a 3 column matrix.
<code>type</code>	This is either "alr" or "ilr", corresponding to the additive and the isometric log-ratio transformation respectively.
<code>n</code>	The number of grid points to consider over which the density is calculated.
<code>appear</code>	Should the available data appear on the ternary plot (TRUE) or not (FALSE)?

Details

The alr or the ilr transformation is applied to the compositional data at first and the location, scatter and degrees of freedom of the bivariate t distribution are computed. Then for a grid of points within the 2-dimensional simplex the bivariate t density is calculated and the contours are plotted along with the points.

Value

A ternary diagram with the points (if `appear = TRUE`) and the bivariate t contour lines.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[diri.contour](#), [mixnorm.contour](#), [norm.contour](#), [skewnorm.contour](#)

Examples

```
x <- as.matrix( iris[, 1:3] )
x <- x / rowSums(x)
bivt.contour(x)
bivt.contour(x, type = "ilr")
```

Cross validation for some compositional regression models

Cross validation for some compositional regression models

Description

Cross validation for some compositional regression models.

Usage

```
cv.comp.reg(y, x, type = "comp.reg", nfolds = 10, folds = NULL, seed = FALSE)
```

Arguments

<code>y</code>	A matrix with compositional data. Zero values are allowed for some regression models.
<code>x</code>	The predictor variable(s).
<code>type</code>	This can be one of the following: "comp.reg", "robust", "kl.compreg", "js.compreg", "diri.reg" or "zadr".
<code>nfolds</code>	The number of folds to be used. This is taken into consideration only if the folds argument is not supplied.
<code>folds</code>	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
<code>seed</code>	If seed is TRUE the results will always be the same.

Details

A k-fold cross validation for a compositional regression model is performed.

Value

A list including:

runtime	The runtime of the cross-validation procedure.
kl	The Kullback-Leibler divergences for all runs.
js	The Jensen-Shannon divergences for all runs.
perf	The average Kullback-Leibler divergence and average Jensen-Shannon divergence.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

See Also

[comp.reg](#), [kl.compreg](#), [compppr.tune](#), [aknreg.tune](#)

Examples

```
y <- as.matrix( iris[, 1:3] )
y <- y / rowSums(y)
x <- iris[, 4]
mod <- cv.comp.reg(y, x)
```

Cross validation for the alpha-k-NN regression for compositional response data
Cross validation for the α -k-NN regression for compositional response data

Description

Cross validation for the α -k-NN regression for compositional response data.

Usage

```
aknreg.tune(y, x, a = seq(0.1, 1, by = 0.1), k = 2:10, apostasi = "euclidean",
n folds = 10, folds = NULL, seed = FALSE, B = 1, rann = FALSE)
```

Arguments

y	A matrix with the compositional response data. Zeros are allowed.
x	A matrix with the available predictor variables.
a	A vector with a grid of values of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied.

<code>k</code>	The number of nearest neighbours to consider. It can be a single number or a vector.
<code>apostasi</code>	The type of distance to use, either "euclidean" or "manhattan".
<code>nfolds</code>	The number of folds. Set to 10 by default.
<code>folds</code>	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
<code>seed</code>	If seed is TRUE the results will always be the same.
<code>B</code>	If you want to correct for the optimistic bias set this to more than 1, otherwise no bootstrap bias correction takes place. If you have large sample sizes, say 1000 or more, bootstrap bias correction may not be really necessary.
<code>rann</code>	If you have large scale datasets and want a faster k-NN search, you can use kd-trees implemented in the R package "RANN". In this case you must set this argument equal to TRUE. Note however, that in this case, the only available distance is by default "euclidean".

Details

A k-fold cross validation for the α -k-NN regression for compositional response data is performed.

Value

A list including:

<code>kl</code>	The Kullback-Leibler divergence for all combinations of α and k.
<code>js</code>	The Jensen-Shannon divergence for all combinations of α and k.
<code>klmin</code>	The minimum Kullback-Leibler divergence.
<code>jsmin</code>	The minimum Jensen-Shannon divergence.
<code>bc.kl</code>	The bootstrap bias corrected minimum Kullback-Leibler divergence.
<code>bs.js</code>	The bootstrap bias corrected minimum Jensen-Shannon divergence.
<code>kl.alpha</code>	The optimum α that leads to the minimum Kullback-Leibler divergence.
<code>kl.k</code>	The optimum k that leads to the minimum Kullback-Leibler divergence.
<code>js.alpha</code>	The optimum α that leads to the minimum Jensen-Shannon divergence.
<code>js.k</code>	The optimum k that leads to the minimum Jensen-Shannon divergence.
<code>runtime</code>	The runtime of the cross-validation procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Michail Tsagris, Abdulaziz Alenazi and Connie Stewart (2020). The α -k-NN regression for compositional data. <https://arxiv.org/pdf/2002.05137.pdf>

See Also

[alfa.rda](#), [alfa.fda](#), [rda.tune](#)

Examples

```
y <- as.matrix( iris[, 1:3] )
y <- y / rowSums(y)
x <- iris[, 4]
mod <- aknnreg.tune(y, x, a = c(0.4, 0.6), k = 2:4, nolds = 5)
```

Cross validation for the alpha-k-NN regression with compositional predictor variables
Cross validation for the α -k-NN regression with compositional predictor variables

Description

Cross validation for the α -k-NN regression with compositional predictor variables.

Usage

```
alfaknnreg.tune(y, x, a = seq(-1, 1, by = 0.1), k = 2:10, nolds = 10,
apostasi = "euclidean", method = "average", folds = NULL, seed = FALSE, graph = FALSE)
```

Arguments

y	The response variable, a numerical vector.
x	A matrix with the available compositional data. Zeros are allowed.
a	A vector with a grid of values of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied.
k	The number of nearest neighbours to consider. It can be a single number or a vector.
nolds	The number of folds. Set to 10 by default.
apostasi	The type of distance to use, either "euclidean" or "manhattan".
method	If you want to take the average of the reponses of the k closest observations, type "average". For the median, type "median" and for the harmonic mean, type "harmonic".
folds	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
seed	If seed is TRUE the results will always be the same.
graph	If graph is TRUE (default value) a filled contour plot will appear.

Details

A k-fold cross validation for the α -k-NN regression for compositional response data is performed.

Value

A list including:

mspe	The mean square error of prediction.
performance	The minimum mean square error of prediction.
opt_a	The optimal value of α .
opt_k	The optimal value of k.
runtime	The runtime of the cross-validation procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Michail Tsagris, Abdulaziz Alenazi and Connie Stewart (2020). The alpha-k-NN- regression for compositional data. <https://arxiv.org/pdf/2002.05137.pdf>

See Also

[alfa.rda](#), [alfa.fda](#), [rda.tune](#)

Examples

```
library(MASS)
x <- as.matrix(fgl[, 2:9])
x <- x / rowSums(x)
y <- fgl[, 1]
mod <- alfaknnreg.tune(y, x, a = seq(0.2, 0.4, by = 0.1), k = 2:4, nfolds = 5)
```

Cross validation for the regularised and flexible discriminant analysis with compositional data using
*Cross validation for the regularised and flexible discriminant analysis
with compositional data using the α -transformation*

Description

Cross validation for the regularised and flexible discriminant analysis with compositional data using the α -transformation.

Usage

```
alfarda.tune(x, ina, a = seq(-1, 1, by = 0.1), n folds = 10,  
gam = seq(0, 1, by = 0.1), del = seq(0, 1, by = 0.1),  
ncores = 1, folds = NULL, stratified = TRUE, seed = FALSE)
```

```
alfafda.tune(x, ina, a = seq(-1, 1, by = 0.1), n folds = 10,  
folds = NULL, stratified = TRUE, seed = FALSE, graph = FALSE)
```

Arguments

x	A matrix with the available compositional data. Zeros are allowed.
ina	A group indicator variable for the available data.
a	A vector with a grid of values of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied.
n folds	The number of folds. Set to 10 by default.
gam	A vector of values between 0 and 1. It is the weight of the pooled covariance and the diagonal matrix.
del	A vector of values between 0 and 1. It is the weight of the LDA and QDA.
ncores	The number of cores to use. If it is more than 1 parallel computing is performed. It is advisable to use it if you have many observations and or many variables, otherwise it will slow down the process.
folds	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
stratified	Do you want the folds to be created in a stratified way? TRUE or FALSE.
seed	If seed is TRUE the results will always be the same.
graph	If graph is TRUE (default value) a filled contour plot will appear.

Details

A k-fold cross validation is performed.

Value

For the `alfa.rda` a list including:

res	The estimated optimal rate and the best values of α , γ and δ .
percent	For the best value of α the averaged over all folds best rates of correct classification. It is a matrix, where rows correspond to the γ values and columns correspond to δ values.
se	The estimated standard errors of the "percent" matrix.
runtime	The runtime of the cross-validation procedure.

For the `alfa.fda` a list including:

per	The performance of the fda in each fold for each value of α .
-----	--

performance	The average performance for each value of α .
opt_a	The optimal value of α .
runtime	The runtime of the cross-validation procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <gioathineou@gmail.com> and Michail Tsagris <mtsagris@uoc.gr>

References

Friedman Jerome, Trevor Hastie and Robert Tibshirani (2009). The elements of statistical learning, 2nd edition. Springer, Berlin

Tsagris M.T., Preston S. and Wood A.T.A. (2016). Improved classification for compositional data using the α -transformation. *Journal of Classification*, 33(2):243-261.

Hastie, Tibshirani and Buja (1994). Flexible Discriminant Analysis by Optimal Scoring. *Journal of the American Statistical Association*, 89(428):1255-1270.

See Also

[alfa.rda](#), [alfa.fda](#), [rda.tune](#)

Examples

```
library(MASS)
x <- as.matrix(fgl[, 2:9])
x <- x / rowSums(x)
ina <- fgl[, 10]
moda <- alfarda.tune(x, ina, a = seq(0.7, 1, by = 0.1), nolds = 10,
gam = seq(0.1, 0.3, by = 0.1), del = seq(0.1, 0.3, by = 0.1) )
```

Cross validation for the ridge regression

Cross validation for the ridge regression

Description

Cross validation for the ridge regression is performed. There is an option for the GCV criterion which is automatic.

Usage

```
ridge.tune(y, x, nolds = 10, lambda = seq(0, 2, by = 0.1), folds = NULL,
ncores = 1, seed = FALSE, graph = FALSE)
```

Arguments

y	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using the logit transformation.
x	A numeric matrix containing the variables.
nfolds	The number of folds in the cross validation.
lambda	A vector with the a grid of values of λ to be used.
folds	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
ncores	The number of cores to use. If it is more than 1 parallel computing is performed.
seed	If seed is TRUE the results will always be the same.
graph	If graph is set to TRUE the performances for each fold as a function of the λ values will appear.

Details

A k-fold cross validation is performed. This function is used by [alfaridge.tune](#).

Value

A list including:

msep	The performance of the ridge regression for every fold.
mspe	The values of the mean prediction error for each value of λ .
lambda	The value of λ which corresponds to the minimum MSPE.
performance	The minimum MSPE.
runtime	The time required by the cross-validation procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <gioathineou@gmail.com> and Michail Tsagris <mtsagris@uoc.gr>

References

Hoerl A.E. and R.W. Kennard (1970). Ridge regression: Biased estimation for nonorthogonal problems. *Technometrics*, 12(1):55-67.

Brown P. J. (1994). *Measurement, Regression and Calibration*. Oxford Science Publications.

See Also

[ridge.reg](#), [alfaridge.tune](#)

Examples

```
y <- as.vector(iris[, 1])
x <- as.matrix(iris[, 2:4])
ridge.tune( y, x, nfolds = 10, lambda = seq(0, 2, by = 0.1), graph = TRUE )
```

Cross validation for the ridge regression with compositional data as predictor using the alpha-transformation
Cross validation for the ridge regression with compositional data as predictor using the α -transformation

Description

Cross validation for the ridge regression is performed. There is an option for the GCV criterion which is automatic. The predictor variables are compositional data and the α -transformation is applied first.

Usage

```
alfaridge.tune(y, x, nfolds = 10, a = seq(-1, 1, by = 0.1),
lambda = seq(0, 2, by = 0.1), folds = NULL, ncores = 1,
graph = TRUE, col.nu = 15, seed = FALSE)
```

Arguments

y	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using the logit transformation.
x	A numeric matrix containing the compositional data, i.e. the predictor variables. Zero values are allowed.
nfolds	The number of folds in the cross validation.
a	A vector with the a grid of values of α to be used.
lambda	A vector with the a grid of values of λ to be used.
folds	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
ncores	The number of cores to use. If it is more than 1 parallel computing is performed. It is advisable to use it if you have many observations and or many variables, otherwise it will slow down the process.
graph	If graph is TRUE (default value) a filled contour plot will appear.
col.nu	A number parameter for the filled contour plot, taken into account only if graph is TRUE.
seed	If seed is TRUE the results will always be the same.

Details

A k-fold cross validation is performed.

Value

If graph is TRUE a field contour a filled contour will appear. A list including:

mspe	The MSPE where rows correspond to the α values and the columns to the number of principal components.
best.par	The best pair of α and λ .
performance	The minimum mean squared error of prediction.
runtime	The run time of the cross-validation procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <gioathineou@gmail.com> and Michail Tsagris <mtsagris@uoc.gr>

References

Hoerl A.E. and R.W. Kennard (1970). Ridge regression: Biased estimation for nonorthogonal problems. *Technometrics*, 12(1):55-67.

Brown P. J. (1994). *Measurement, Regression and Calibration*. Oxford Science Publications.

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In *Proceedings of the 4th Compositional Data Analysis Workshop*, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[alfa.ridge](#), [ridge.tune](#)

Examples

```
library(MASS)
y <- as.vector(fgl[, 1])
x <- as.matrix(fgl[, 2:9])
x <- x / rowSums(x)
alfaridge.tune( y, x, nfolds = 10, a = seq(0.1, 1, by = 0.1),
lambda = seq(0, 1, by = 0.1) )
```

Cross validation for the transformation-free linear regression for compositional responses and predictors

Cross validation for the transformation-free linear regression for compositional responses and predictors

Description

Cross validation for the transformation-free linear regression for compositional responses and predictors.

Usage

```
cv.tflr(y, x, nfolds = 10, folds = NULL, seed = FALSE)
```

Arguments

y	A matrix with compositional response data. Zero values are allowed.
x	A matrix with compositional predictors. Zero values are allowed.
nfolds	The number of folds to be used. This is taken into consideration only if the folds argument is not supplied.
folds	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
seed	If seed is TRUE the results will always be the same.

Details

A k-fold cross validation for the transformation-free linear regression for compositional responses and predictors is performed.

Value

A list including:

runtime	The runtime of the cross-validation procedure.
kl	The Kullback-Leibler divergences for all runs.
js	The Jensen-Shannon divergences for all runs.
perf	The average Kullback-Leibler divergence and average Jensen-Shannon divergence.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

See Also

[tflr](#), [klalfapcr](#), [tune](#)

Examples

```
## Not run:
library(MASS)
y <- rdiri(214, runif(3, 1, 3))
x <- as.matrix(fgl[, 2:9])
x <- x / rowSums(x)
mod <- cv.tflr(y, x)
mod

## End(Not run)
```

Density of the Dirichlet distribution

Density of the Dirichlet distribution

Description

Density of the Dirichlet distribution.

Usage

```
diri.density(x, a, logged = FALSE)
```

Arguments

x	A vector or a matrix with compositional data.
a	A vector of the non-negative alpha parameters.
logged	Do you want the logarithm of the density values? TRUE or FALSE.

Value

The density value(s).

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

See Also

[fd.density](#), [rdiri](#), [diri.nr](#)

Examples

```
a <- runif(3, 1, 5)
x <- rdiri(100, a)
a <- diri.nr(x)$param
x <- diri.density(x, a)
```

Density of the Flexible Dirichlet distribution
Density of the Flexible Dirichlet distribution

Description

Density of the Flexible Dirichlet distribution

Usage

```
fd.density(x, alpha, prob, tau)
```

Arguments

x	A vector or a matrix with compositional data.
alpha	A vector of the non-negative alpha parameters.
prob	A vector of the clusters' probabilities. It must sum to one.
tau	The non-negative scalar tau parameter.

Details

For more information see the references.

Value

The density value(s).

Author(s)

Michail Tsagris ported from the R package FlexDir. <mtsagris@uoc.gr>.

References

Ongaro, A. and Migliorati, S. (2013) A generalization of the Dirichlet distribution. *Journal of Multivariate Analysis*, 114, 412–426.

Migliorati, S., Ongaro, A. and Monti, G. S. (2016) A structured Dirichlet mixture model for compositional data: inferential and applicative issues. *Statistics and Computing*, 1–21.

See Also

[fd.est](#), [rfd](#)

Examples

```
alpha <- c(12, 11, 10)
prob <- c(0.25, 0.25, 0.5)
tau <- 8
x <- rfd(20, alpha, prob, tau)
fd.density(x, alpha, prob, tau)
```

Density values of a Dirichlet distribution

Density values of a Dirichlet distribution

Description

Density values of a Dirichlet distribution.

Usage

```
ddiri(x, a, logged = TRUE)
```

Arguments

x	A matrix containing compositional data. This can be a vector or a matrix with the data.
a	A vector of parameters. Its length must be equal to the number of components, or columns of the matrix with the compositional data and all values must be greater than zero.
logged	A boolean variable specifying whether the logarithm of the density values to be returned. It is set to TRUE by default.

Details

The density of the Dirichlet distribution for a vector or a matrix of compositional data is returned.

Value

A vector with the density values.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Ng Kai Wang, Guo-Liang Tian and Man-Lai Tang (2011). Dirichlet and related distributions: Theory, methods and applications. John Wiley & Sons.

See Also

[diri.nr](#), [diri.est](#), [diri.contour](#), [rdiri](#)

Examples

```
x <- rdiri( 100, c(5, 7, 4, 8, 10, 6, 4) )
a <- diri.est(x)
f <- ddiri(x, a$param)
sum(f)
a
```

Dirichlet random values simulation

Dirichlet random values simulation

Description

Dirichlet random values simulation.

Usage

```
rdiri(n, a)
```

Arguments

n	The sample size, a numerical value.
a	A numerical vector with the parameter values.

Details

The algorithm is straightforward, for each vector, independent gamma values are generated and then divided by their total sum.

Value

A matrix with the simulated data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Ng Kai Wang, Guo-Liang Tian and Man-Lai Tang (2011). Dirichlet and related distributions: Theory, methods and applications. John Wiley & Sons.

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[diri.est](#), [diri.nr](#), [diri.contour](#)

Examples

```
x <- rdiri( 100, c(5, 7, 1, 3, 10, 2, 4) )
diri.est(x)
```

Dirichlet regression *Dirichlet regression*

Description

Dirichlet regression.

Usage

```
diri.reg(y, x, plot = TRUE, xnew = NULL)

diri.reg2(y, x, xnew = NULL)
```

Arguments

y	A matrix with the compositional data (dependent variable). Zero values are not allowed.
x	The predictor variable(s), they can be either continuous or categorical or both.
plot	A boolean variable specifying whether to plot the leverage values of the observations or not. This is taken into account only when xnew = NULL.
xnew	If you have new data use it, otherwise leave it NULL.

Details

A Dirichlet distribution is assumed for the regression. This involves numerical optimization. The function "diri.reg2" allows for the covariates to be linked with the precision parameter ϕ via the exponential link function $\phi = e^{x*b}$.

Value

A list including:

runtime	The time required by the regression.
loglik	The value of the log-likelihood.
phi	The precision parameter. If covariates are linked with it (function "diri.reg2"), this will be a vector.
hipar	The coefficients of the phi parameter if it is linked to the covariates.

<code>std.phi</code>	The standard errors of the coefficients of the phi parameter is it linked to the covariates.
<code>log.phi</code>	The logarithm of the precision parameter.
<code>std.logphi</code>	The standard error of the logarithm of the precision parameter.
<code>be</code>	The beta coefficients.
<code>seb</code>	The standard error of the beta coefficients.
<code>sigma</code>	The covariance matrix of the regression parameters (for the mean vector and the phi parameter) in the function "diri.reg2".
<code>lev</code>	The leverage values.
<code>est</code>	For the "diri.reg" this contains the fitted or the predicted values (if <code>xnew</code> is not NULL). For the "diri.reg2" if <code>xnew</code> is NULL, this is also NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Maier, Marco J. (2014) DirichletReg: Dirichlet Regression for Compositional Data in R. Research Report Series/Department of Statistics and Mathematics, 125. WU Vienna University of Economics and Business, Vienna. <http://epub.wu.ac.at/4077/1/Report125.pdf>

Gueorguieva, Ralitzia, Robert Rosenheck, and Daniel Zelterman (2008). Dirichlet component regression and its applications to psychiatric data. *Computational statistics & data analysis* 52(12): 5344-5355.

See Also

[js.compreg](#), [kl.compreg](#), [ols.compreg](#), [comp.reg](#), [alfa.reg](#)

Examples

```
x <- as.vector(iris[, 4])
y <- as.matrix(iris[, 1:3])
y <- y / rowSums(y)
mod1 <- diri.reg(y, x)
mod2 <- diri.reg2(y, x)
mod3 <- comp.reg(y, x)
```

Divergence based regression for compositional data

Divergence based regression for compositional data

Description

Regression for compositional data based on the Kullback-Leibler the Jensen-Shannon divergence and the symmetric Kullback-Leibler divergence.

Usage

```
kl.compreg(y, x, B = 1, ncores = 1, xnew = NULL, tol = 1e-07, maxiters = 50)
js.compreg(y, x, B = 1, ncores = 1, xnew = NULL)
symkl.compreg(y, x, B = 1, ncores = 1, xnew = NULL)
```

Arguments

y	A matrix with the compositional data (dependent variable). Zero values are allowed.
x	The predictor variable(s), they can be either continuous or categorical or both.
B	If B is greater than 1 bootstrap estimates of the standard error are returned. If B=1, no standard errors are returned.
ncores	If ncores is 2 or more parallel computing is performed. This is to be used for the case of bootstrap. If B=1, this is not taken into consideration.
xnew	If you have new data use it, otherwise leave it NULL.
tol	The tolerance value to terminate the Newton-Raphson procedure.
maxiters	The maximum number of Newton-Raphson iterations.

Details

In the `kl.compreg` the Kullback-Leibler divergence is adopted as the objective function. In case of problematic convergence the "multinom" function by the "nnet" package is employed. This will obviously be slower. The `js.compreg` uses the Jensen-Shannon divergence and the `symkl.compreg` uses the symmetric Kullback-Leibler divergence. There is no actual log-likelihood for neither regression.

Value

A list including:

runtime	The time required by the regression.
iters	The number of iterations required by the Newton-Raphson in the <code>kl.compreg</code> function.
loglik	The log-likelihood. This is actually a quasi multinomial regression. This is basically minus the half deviance, or $-\sum_{i=1}^n y_i \log y_i / \hat{y}_i$.

be The beta coefficients.
seb The standard error of the beta coefficients, if bootstrap is chosen, i.e. if $B > 1$.
est The fitted values of x_{new} if x_{new} is not NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Murteira, Jose MR, and Joaquim JS Ramalho 2016. Regression analysis of multivariate fractional data. *Econometric Reviews* 35(4): 515-552.

Tsagris, Michail (2015). A novel, divergence based, regression for compositional data. Proceedings of the 28th Panhellenic Statistics Conference, 15-18/4/2015, Athens, Greece. <https://arxiv.org/pdf/1511.07600.pdf>

Endres, D. M. and Schindelin, J. E. (2003). A new metric for probability distributions. *Information Theory, IEEE Transactions on* 49, 1858-1860.

Osterreicher, F. and Vajda, I. (2003). A new class of metric divergences on probability spaces and its applicability in statistics. *Annals of the Institute of Statistical Mathematics* 55, 639-653.

See Also

[diri.reg](#), [js.compreg](#), [ols.compreg](#), [comp.reg](#)

Examples

```
library(MASS)
x <- as.vector(fg1[, 1])
y <- as.matrix(fg1[, 2:9])
y <- y / rowSums(y)
mod1<- kl.compreg(y, x, B = 1, ncores = 1)
mod2 <- js.compreg(y, x, B = 1, ncores = 1)
```

Divergence based regression for compositional data with compositional data in the covariates side using the α -transformation

Divergence based regression for compositional data with compositional data in the covariates side using the α -transformation

Description

Divergence based regression for compositional data with compositional data in the covariates side using the α -transformation.

Usage

```
kl.alfapcr(y, x, covar = NULL, a, k, xnew = NULL, B = 1, ncores = 1, tol = 1e-07,
maxiters = 50)
```

Arguments

y	A numerical matrix with compositional data with or without zeros.
x	A matrix with the predictor variables, the compositional data. Zero values are allowed.
covar	If you have other covariates as well put them here.
a	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied.
k	A number at least equal to 1. How many principal components to use.
xnew	A matrix containing the new compositional data whose response is to be predicted. If you have no new data, leave this NULL as is by default.
B	If B is greater than 1 bootstrap estimates of the standard error are returned. If B=1, no standard errors are returned.
ncores	If ncores is 2 or more parallel computing is performed. This is to be used for the case of bootstrap. If B=1, this is not taken into consideration.
tol	The tolerance value to terminate the Newton-Raphson procedure.
maxiters	The maximum number of Newton-Raphson iterations.

Details

The α -transformation is applied to the compositional data first, the first k principal component scores are calculated and used as predictor variables for the Kullback-Leibler divergence based regression model.

Value

A list including:

runtime	The time required by the regression.
iters	The number of iterations required by the Newton-Raphson in the kl.compreg function.
loglik	The log-likelihood. This is actually a quasi multinomial regression. This is basically minus the half deviance, or $-\sum_{i=1}^n y_i \log y_i / \hat{y}_i$.
be	The beta coefficients.
seb	The standard error of the beta coefficients, if bootstrap is chosen, i.e. if B > 1.
est	The fitted values of xnew if xnew is not NULL.

Author(s)

Initial code by Abdulaziz Alenazi. Modifications by Michail Tsagris.

R implementation and documentation: Abdulaziz Alenazi <a.alenazi@nbu.edu.sa> Michail Tsagris <mtsagris@uoc.gr>

References

Alenazi A. (2019). Regression for compositional data with compositional data as predictor variables with or without zero values. *Journal of Data Science*, 17(1): 219-238. http://www.jds-online.com/file_download/688/01+No.10+315+REGRESSION+FOR+COMPOSITIONAL+DATA+WITH+COMPOSITIONAL

Tsagris M. (2015). Regression analysis with compositional data containing zero values. *Chilean Journal of Statistics*, 6(2): 47-57. <http://arxiv.org/pdf/1508.01913v1.pdf>

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In *Proceedings of the 4th Compositional Data Analysis Workshop*, Girona, Spain. <http://arxiv.org/pdf/1106.1451.pdf>

See Also

[klalfapcr.tune](#), [tflr](#), [pcr](#), [glm.pcr](#), [alfapcr.tune](#)

Examples

```
library(MASS)
y <- rdirich(214, runif(4, 1, 3))
x <- as.matrix(fgl[, 2:9])
x <- x / rowSums(x)
mod <- alfa.pcr(y = y, x = x, 0.7, 1)
mod
```

Divergence matrix of compositional data

Divergence matrix of compositional data

Description

Divergence matrix of compositional data.

Usage

```
divergence(x, type = "kullback_leibler", vector = FALSE)
```

Arguments

x	A matrix with the compositional data.
type	This is either "kullback_leibler" (Kullback-Leibler, which computes the symmetric Kullback-Leibler divergence) or "jensen_shannon" (Jensen-Shannon) divergence.
vector	For return a vector instead a matrix.

Details

The function produces the distance matrix either using the Kullback-Leibler (distance) or the Jensen-Shannon (metric) divergence. The Kullback-Leibler refers to the symmetric Kullback-Leibler divergence.

Value

if the vector argument is `FALSE` a symmetric matrix with the divergences, otherwise a vector with the divergences.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

Endres, D. M. and Schindelin, J. E. (2003). A new metric for probability distributions. *Information Theory, IEEE Transactions on* 49, 1858-1860.

Osterreicher, F. and Vajda, I. (2003). A new class of metric divergences on probability spaces and its applicability in statistics. *Annals of the Institute of Statistical Mathematics* 55, 639-653.

See Also

[comp.knn](#), [js.compreg](#)

Examples

```
x <- as.matrix(iris[1:20, 1:4])
x <- x / rowSums(x)
divergence(x)
```

Empirical likelihood for a one sample mean vector hypothesis testing
Empirical likelihood for a one sample mean vector hypothesis testing

Description

Empirical likelihood for a one sample mean vector hypothesis testing.

Usage

```
el.test1(x, mu, R = 1, ncores = 1, graph = FALSE)
```

Arguments

x	A matrix containing Euclidean data.
mu	The hypothesized mean vector.
R	If R is 1 no bootstrap calibration is performed and the classical p-value via the χ^2 distribution is returned. If R is greater than 1, the bootstrap p-value is returned.
ncores	The number of cores to use, set to 1 by default.
graph	A boolean variable which is taken into consideration only when bootstrap calibration is performed. IF TRUE the histogram of the bootstrap test statistic values is plotted.

Details

Multivariate hypothesis test for a one sample mean vector. This is a non parametric test and it works for univariate and multivariate data.

Value

A list with the outcome of the function `el.test` which includes the -2 log-likelihood ratio, the observed P-value by chi-square approximation, the final value of Lagrange multiplier λ , the gradient at the maximum, the Hessian matrix, the weights on the observations (probabilities multiplied by the sample size) and the number of iteration performed. In addition the runtime of the procedure is reported. In the case of bootstrap, the bootstrap p-value is also returned.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Owen, A. (1990). Empirical likelihood ratio confidence regions. *Annals of Statistics*, 18, 90-120.

Owen A. B. (2001). *Empirical likelihood*. Chapman and Hall/CRC Press.

See Also

`eel.test1`, `hotel1T2`, `james`, `hotel2T2`, `maov`, `el.test2`, `comp.test`

Examples

```
x <- Rfast::rmvnorm(100, numeric(10), diag( rexp(10, 0.5) ) )
el.test1(x, mu = numeric(10) )
eel.test1(x, mu = numeric(10) )
```

Empirical likelihood hypothesis testing for two mean vectors

Empirical likelihood hypothesis testing for two mean vectors

Description

Empirical likelihood hypothesis testing for two mean vectors.

Usage

```
el.test2(y1, y2, R = 0, ncores = 1, graph = FALSE)
```

Arguments

y1	A matrix containing the Euclidean data of the first group.
y2	A matrix containing the Euclidean data of the second group.
R	If R is 0, the classical chi-square distribution is used, if R = 1, the corrected chi-square distribution (James, 1954) is used and if R = 2, the modified F distribution (Krishnamoorthy and Yanping, 2006) is used. If R is greater than 3 bootstrap calibration is performed.
ncores	How many to cores to use.
graph	A boolean variable which is taken into consideration only when bootstrap calibration is performed. IF TRUE the histogram of the bootstrap test statistic values is plotted.

Details

Empirical likelihood is a non parametric hypothesis testing procedure for one sample. The generalization to two (or more samples) is via searching for the mean vector that minimizes the sum of the two test statistics.

Value

A list including:

test	The empirical likelihood test statistic value.
modif.test	The modified test statistic, either via the chi-square or the F distribution.
dof	Three degrees of freedom of the chi-square or the F distribution.
pvalue	The asymptotic or the bootstrap p-value.
mu	The estimated common mean vector.
runtime	The runtime of the bootstrap calibration.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

- G.S. James (1954). Tests of Linear Hypotheses in Univariate and Multivariate Analysis when the Ratios of the Population Variances are Unknown. *Biometrika*, 41(1/2): 19-43
- Krishnamoorthy K. and Yanping Xia (2006). On Selecting Tests for Equality of Two Normal Mean Vectors. *Multivariate Behavioral Research* 41(4): 533-548.
- Owen A. B. (2001). Empirical likelihood. Chapman and Hall/CRC Press.
- Owen A.B. (1988). Empirical likelihood ratio confidence intervals for a single functional. *Biometrika* 75(2): 237-249.
- Amaral G.J.A., Dryden I.L. and Wood A.T.A. (2007). Pivotal bootstrap methods for k-sample problems in directional statistics and shape analysis. *Journal of the American Statistical Association* 102(478): 695-707.
- Preston S.P. and Wood A.T.A. (2010). Two-Sample Bootstrap Hypothesis Tests for Three-Dimensional Labelled Landmark Data. *Scandinavian Journal of Statistics* 37(4): 568-587.

See Also

[eel.test2](#), [maovjames](#), [maov](#), [hotel2T2](#), [james](#), [comp.test](#)

Examples

```
e1.test2( y1 = as.matrix(iris[1:25, 1:4]), y2 = as.matrix(iris[26:50, 1:4]), R = 0 )
e1.test2( y1 = as.matrix(iris[1:25, 1:4]), y2 = as.matrix(iris[26:50, 1:4]), R = 1 )
e1.test2( y1 = as.matrix(iris[1:25, 1:4]), y2 = as.matrix(iris[26:50, 1:4]), R = 2 )
```

Estimating location and scatter parameters for compositional data

Estimating location and scatter parameters for compositional data

Description

Estimating location and scatter parameters for compositional data in a robust and non robust way.

Usage

```
comp.den(x, type = "alr", dist = "normal", tol = 1e-09)
```

Arguments

x	A matrix containing compositional data. No zero values are allowed.
type	A boolean variable indicating the transformation to be used. Either "alr" or "ilr" corresponding to the additive or the isometric log-ratio transformation respectively.
dist	Takes values "normal", "t", "skewnorm", "rob" and "spatial". They first three options correspond to the parameters of the normal, t and skew normal distribution respectively. If it set to "rob" the MCD estimates are computed and if set to "spatial" the spatial median and spatial sign covariance matrix are computed.
tol	A tolerance level to terminate the process of finding the spatial median when dist = "spatial". This is set to 1e-09 by default.

Details

This function calculates robust and non robust estimates of location and scatter.

Value

A list including: The mean vector and covariance matrix mainly. Other parameters are also returned depending on the value of the argument "dist".

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

- P. J. Rousseeuw and K. van Driessen (1999) A fast algorithm for the minimum covariance determinant estimator. *Technometrics* 41, 212-223.
- Mardia K.V., Kent J.T., and Bibby J.M. (1979). *Multivariate analysis*. Academic press.
- Aitchison J. (1986). *The statistical analysis of compositional data*. Chapman & Hall.
- T. Karkkainen and S. Ayramo (2005). On computation of spatial median for robust data mining. *Evolutionary and Deterministic Methods for Design, Optimization and Control with Applications to Industrial and Societal Problems EUROGEN 2005*.
- A Durre, D Vogel, DE Tyler (2014). The spatial sign covariance matrix with unknown location. *Journal of Multivariate Analysis*, 130: 107-117.
- J. T. Kent, D. E. Tyler and Y. Vardi (1994) A curious likelihood identity for the multivariate t-distribution. *Communications in Statistics-Simulation and Computation* 23, 441-453.
- Azzalini A. and Dalla Valle A. (1996). The multivariate skew-normal distribution. *Biometrika* 83(4): 715-726.

See Also

[spatmed.reg](#), [multivt](#)

Examples

```
library(MASS)
x <- as.matrix(iris[, 1:4])
x <- x / rowSums(x)
comp.den(x)
comp.den(x, type = "alr", dist = "t")
comp.den(x, type = "alr", dist = "spatial")
```

Estimation of the value of alpha in the folded model

Estimation of the value of α in the folded model

Description

Estimation of the value of α in the folded model.

Usage

```
a.est(x)
```

Arguments

x A matrix with the compositional data. No zero values are allowed.

Details

This is a function for choosing or estimating the value of α in the folded model (Tsagris and Stewart, 2019).

Value

A list including:

runtime	The runtime of the algorithm.
best	The estimated optimal α of the folded model.
loglik	The maximised log-likelihood of the folded model.
p	The estimated probability inside the simplex of the folded model.
mu	The estimated mean vector of the folded model.
su	The estimated covariance matrix of the folded model.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

Tsagris Michail and Stewart Connie, (2020). A folded model for compositional data analysis. Australian and New Zealand Journal of Statistics (to appear). <https://arxiv.org/pdf/1802.07330.pdf>

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[alfa.profile](#), [alfa](#), [alfainv](#), [alpha.mle](#)

Examples

```
x <- as.matrix(iris[, 1:4])
x <- x / rowSums(x)
alfa.tune(x)
a.est(x)
```

Estimation of the value of alpha via the profile log-likelihood
Estimation of the value of α via the alfa profile log-likelihood

Description

Estimation of the value of α via the alfa profile log-likelihood.

Usage

```
alfa.profile(x, a = seq(-1, 1, by = 0.01))
```

Arguments

x	A matrix with the compositional data. Zero values are not allowed.
a	A grid of values of α .

Details

For every value of α the normal likelihood (see the refernece) is computed. At the end, the plot of the values is constructed.

Value

A list including:

res	The chosen value of α , the corresponding log-likelihood value and the log-likelihood when $\alpha = 0$.
ci	An asymptotic 95% confidence interval computed from the log-likelihood ratio test.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[alfa.tune](#), [alfa](#), [alfainv](#)

Examples

```
x <- as.matrix(iris[, 1:4])
x <- x / rowSums(x)
alfa.tune(x)
alfa.profile(x)
```

Exponential empirical likelihood for a one sample mean vector hypothesis testing
Exponential empirical likelihood for a one sample mean vector hypothesis testing

Description

Exponential empirical likelihood for a one sample mean vector hypothesis testing.

Usage

```
eel.test1(x, mu, tol = 1e-06, R = 1)
```

Arguments

x	A matrix containing Euclidean data.
mu	The hypothesized mean vector.
tol	The tolerance value used to stop the Newton-Raphson algorithm.
R	The number of bootstrap samples used to calculate the p-value. If R = 1 (default value), no bootstrap calibration is performed

Details

Multivariate hypothesis test for a one sample mean vector. This is a non parametric test and it works for univariate and multivariate data.

Value

A list including:

<code>p</code>	The estimated probabilities.
<code>lambda</code>	The value of the Lagrangian parameter λ .
<code>iter</code>	The number of iterations required by the newton-Raphson algorithm.
<code>info</code>	The value of the log-likelihood ratio test statistic along with its corresponding p-value.
<code>runtime</code>	The runtime of the process.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Jing Bing-Yi and Andrew TA Wood (1996). Exponential empirical likelihood is not Bartlett correctable. *Annals of Statistics* 24(1): 365-369.

Owen A. B. (2001). *Empirical likelihood*. Chapman and Hall/CRC Press.

See Also

[el.test1](#), [hotel1T2](#), [james](#), [hotel2T2](#), [maov](#), [el.test2](#), [comp.test](#)

Examples

```
x <- Rfast::rmvnorm(100, numeric(10), diag( rexp(10, 0.5) ) )
eel.test1(x, numeric(10) )
el.test1(x, numeric(10) )
```

Exponential empirical likelihood hypothesis testing for two mean vectors
Exponential empirical likelihood hypothesis testing for two mean vectors

Description

Exponential empirical likelihood hypothesis testing for two mean vectors.

Usage

```
eel.test2(y1, y2, tol = 1e-07, R = 0, graph = FALSE)
```

Arguments

y1	A matrix containing the Euclidean data of the first group.
y2	A matrix containing the Euclidean data of the second group.
tol	The tolerance level used to terminate the Newton-Raphson algorithm.
R	If R is 0, the classical chi-square distribution is used, if R = 1, the corrected chi-square distribution (James, 1954) is used and if R = 2, the modified F distribution (Krishnamoorthy and Yanping, 2006) is used. If R is greater than 3 bootstrap calibration is performed.
graph	A boolean variable which is taken into consideration only when bootstrap calibration is performed. IF TRUE the histogram of the bootstrap test statistic values is plotted.

Details

Exponential empirical likelihood is a non parametric hypothesis testing procedure for one sample. The generalization to two (or more samples) is via searching for the mean vector that minimises the sum of the two test statistics.

Value

A list including:

test	The empirical likelihood test statistic value.
modif.test	The modified test statistic, either via the chi-square or the F distribution.
dof	The degrees of freedom of the chi-square or the F distribution.
pvalue	The asymptotic or the bootstrap p-value.
mu	The estimated common mean vector.
runtime	The runtime of the bootstrap calibration.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

- Jing Bing-Yi and Andrew TA Wood (1996). Exponential empirical likelihood is not Bartlett correctable. *Annals of Statistics* 24(1): 365-369.
- G.S. James (1954). Tests of Linear Hypotheses in Univariate and Multivariate Analysis when the Ratios of the Population Variances are Unknown. *Biometrika*, 41(1/2): 19-43
- Krishnamoorthy K. and Yanping Xia (2006). On Selecting Tests for Equality of Two Normal Mean Vectors. *Multivariate Behavioral Research* 41(4): 533-548.
- Owen A. B. (2001). *Empirical likelihood*. Chapman and Hall/CRC Press.

Amaral G.J.A., Dryden I.L. and Wood A.T.A. (2007). Pivotal bootstrap methods for k-sample problems in directional statistics and shape analysis. *Journal of the American Statistical Association* 102(478): 695-707.

Preston S.P. and Wood A.T.A. (2010). Two-Sample Bootstrap Hypothesis Tests for Three-Dimensional Labelled Landmark Data. *Scandinavian Journal of Statistics* 37(4): 568-587.

Tsagris M., Preston S. and Wood A.T.A. (2017). Nonparametric hypothesis testing for equality of means on the simplex. *Journal of Statistical Computation and Simulation*, 87(2): 406-422.

See Also

[e1.test2](#), [maovjames](#), [maov](#), [hotel2T2](#), [james](#), [comp.test](#)

Examples

```
y1 = as.matrix(iris[1:25, 1:4])
y2 = as.matrix(iris[26:50, 1:4])
eel.test2(y1, y2)
eel.test2(y1, y2 )
eel.test2( y1, y2 )
```

Fast estimation of the value of α

Fast estimation of the value of α

Description

Fast estimation of the value of α .

Usage

```
alfa.tune(x, B = 1, ncores = 1)
```

Arguments

x	A matrix with the compositional data. No zero values are allowed.
B	If no (bootstrap based) confidence intervals should be returned this should be 1 and more than 1 otherwise.
ncores	If ncores is greater than 1 parallel computing is performed. It is advisable to use it if you have many observations and or many variables, otherwise it will slow down the process.

Details

This is a faster function than [alfa.profile](#) for choosing the value of α .

Value

A vector with the best alpha, the maximised log-likelihood and the log-likelihood at $\alpha = 0$, when $B = 1$ (no bootstrap). If $B > 1$ a list including:

param	The best alpha and the value of the log-likelihood, along with the 95% bootstrap based confidence intervals.
message	A message with some information about the histogram.
runtime	The time (in seconds) of the process.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[alfa.profile](#), [alfa](#), [alfainv](#)

Examples

```
library(MASS)
x <- as.matrix(iris[, 1:4])
x <- x / rowSums(x)
alfa.tune(x)
alfa.profile(x)
```

Fitting a Dirichlet distribution

Fitting a Dirichlet distribution

Description

Estimation of the parameters of a fitted Dirichlet distribution.

Usage

```
diri.est(x, type = "mle")
```

Arguments

x	A matrix containing compositional data.
type	If you want to estimate the parameters use type="mle". If you want to estimate the mean vector along with the precision parameter, the second parametrisation of the Dirichlet, use type="prec".

Details

Maximum likelihood estimation of the parameters of a Dirichlet distribution is performed.

Value

A list including:

loglik	The value of the log-likelihood.
param	The estimated parameters.
phi	The estimated precision parameter, if type = "prec".
a	The estimated mean vector, if type = "prec".
runtime	The run time of the maximisation procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Ng Kai Wang, Guo-Liang Tian and Man-Lai Tang (2011). Dirichlet and related distributions: Theory, methods and applications. John Wiley & Sons.

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[diri.nr](#), [diri.contour](#), [rdiri](#), [ddiri](#)

Examples

```
x <- rdiri( 100, c(5, 7, 1, 3, 10, 2, 4) )
diri.est(x)
diri.est(x, type = "prec")
```

Fitting a Dirichlet distribution via Newton-Rapshon

Fitting a Dirichlet distribution via Newton-Rapshon

Description

Fitting a Dirichlet distribution via Newton-Rapshon.

Usage

```
diri.nr(x, type = 1, tol = 1e-07)
```

Arguments

x	A matrix containing compositional data. Zeros are not allowed.
type	Type can either be 1, so that the Newton-Rapshon is used for the maximisation of the log-likelihood, as Minka (2012) suggested or it can be 1. In the latter case the Newton-Raphson algorithm is implemented involving matrix inversions. In addition an even faster implementation has been implemented (in C++) in the package Rfast and is used here.
tol	The tolerance level indicating no further increase in the log-likelihood.

Details

Maximum likelihood estimation of the parameters of a Dirichlet distribution is performed via Newton-Raphson. Initial values suggested by Minka (2003) are used. The estimation is super faster than "diri.est" and the difference becomes really apparent when the sample size and or the dimensions increase. In fact this will work with millions of observations. So in general, I trust this one more than "diri.est".

The only problem I have seen with this method is that if the data are concentrated around a point, say the center of the simplex, it will be hard for this and the previous methods to give estimates of the parameters. In this extremely difficult scenario I would suggest the use of the previous function with the precision parametrization "diri.est(x, type = "prec")". It will be extremely fast and accurate.

Value

A list including:

iter	The number of iterations required. If the argument "type" is set to 2 this is not returned.
loglik	The value of the log-likelihood.
param	The estimated parameters.
runtime	The run time of the procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Thomas P. Minka (2003). Estimating a Dirichlet distribution. <http://research.microsoft.com/en-us/um/people/minka/papers/dirichlet/minka-dirichlet.pdf>

See Also

[diri.est](#), [diri.contour](#) [rdiri](#), [ddiri](#)

Examples

```
x <- rdiri( 100, c(5, 7, 5, 8, 10, 6, 4) )
diri.nr(x)
diri.nr(x, type = 2)
diri.est(x)
```

Fitting a Flexible Dirichlet distribution

Fitting a Flexible Dirichlet distribution

Description

Fitting a Flexible Dirichlet distribution.

Usage

```
fd.est(x, ini.iter = 50, final.iter = 100)
```

Arguments

<code>x</code>	A matrix or a dataframe containing the compositional data.
<code>ini.iter</code>	Number of iterations for the initial SEM step. Default value is 50.
<code>final.iter</code>	Number of iterations for the final EM step. Default value is 100.

Details

For more information see the references.

Value

A list including:

alpha	Estimated values of the parameter vector Alpha.
prob	Estimated values of the parameter vector P.
tau	Estimated value of the parameter Tau.
loglik	The Log-likelihood value.

Author(s)

Michail Tsagris ported from the R package FlexDir. <mtsagris@uoc.gr>.

References

- Ongaro, A. and Migliorati, S. (2013) A generalization of the Dirichlet distribution. *Journal of Multivariate Analysis*, 114, 412–426.
- Migliorati, S., Ongaro, A. and Monti, G. S. (2016) A structured Dirichlet mixture model for compositional data: inferential and applicative issues. *Statistics and Computing*.

See Also

[rfd](#), [fd.density](#)

Examples

```
## Not run:
x <- rfd(n = 50, a = c(12, 11, 10), p = c(0.25, 0.25, 0.5), tau = 4 )
ela <- fd.est(x, ini.iter = 10, final.iter = 20)
ela

## End(Not run)
```

Gaussian mixture models for compositional data
Gaussian mixture models for compositional data

Description

Gaussian mixture models for compositional data.

Usage

```
mix.compnorm(x, g, model, type = "alr")
```

Arguments

x	A matrix with the compositional data.
g	How many clusters to create.
model	The type of model to be used. <ol style="list-style-type: none"> 1. "EII": All groups have the same diagonal covariance matrix, with the same variance for all variables. 2. "VII": Different diagonal covariance matrices, with the same variance for all variables within each group. 3. "EEI": All groups have the same diagonal covariance matrix. 4. "VEI": Different diagonal covariance matrices. If we make all covariance matrices have determinant 1, (divide the matrix with the p-th root of its determinant) then all covariance matrices will be the same. 5. "EVI": Different diagonal covariance matrices with the same determinant. 6. "VVI": Different diagonal covariance matrices, with nothing in common. 7. "EEE": All covariance matrices are the same. 8. "EEV": Different covariance matrices, but with the same determinant and in addition, if we make them have determinant 1, they will have the same trace. 9. "VEV": Different covariance matrices but if we make the matrices have determinant 1, then they will have the same trace. 10. "VVV": Different covariance matrices with nothing in common. 11. "EVE": Different covariance matrices, but with the same determinant. In addition, calculate the eigenvectors for each covariance matrix and you will see the extra similarities. 12. "VVE": Different covariance matrices, but they have something in common with their directions. Calculate the eigenvectors of each covariance matrix and you will see the similarities. 13. "VEE": Different covariance matrices, but if we make the matrices have determinant 1, then they will have the same trace. In addition, calculate the eigenvectors for each covariance matrix and you will see the extra similarities. 14. "EVV": Different covariance matrices, but with the same determinant.
type	Either the additive ("alr") or the isometric ("ilr") log-ratio transformation is to be used..

Details

A log-ratio transformation is applied and then a Gaussian mixture model is constructed.

Value

A list including:

mu	A matrix where each row corresponds to the mean vector of each cluster.
su	An array containing the covariance matrix of each cluster.
prob	The estimated mixing probabilities.
est	The estimated cluster membership values.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Ryan P. Browne, Aisha ElSherbiny and Paul D. McNicholas (2015). R package mixture: Mixture Models for Clustering and Classification.

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[bic.mixcompnorm](#), [rmixcomp](#), [mixnorm.contour](#)

Examples

```
## Not run:
x <- as.matrix(iris[, 1:4])
x <- x/ rowSums(x)
mod1 <- mix.compnorm(x, 3, model = "EII" )
mod2 <- mix.compnorm(x, 4, model = "VII")

## End(Not run)
```

Generate random folds for cross-validation

Generate random folds for cross-validation

Description

Random folds for use in a cross validation are generated. There is the option for stratified splitting as well.

Usage

```
makefolds(ina, nfolds = 10, stratified = TRUE, seed = FALSE)
```

Arguments

<code>ina</code>	A variable indicating the groupings.
<code>nfolds</code>	The number of folds to produce.
<code>stratified</code>	A boolean variable specifying whether stratified random (TRUE) or simple random (FALSE) sampling is to be used when producing the folds.
<code>seed</code>	A boolean variable. If set to TRUE, the folds will always be the same.

Details

I was inspired by the command in the package **TunePareto** in order to do the stratified version.

Value

A list with `nfolds` elements where each element is a fold containing the indices of the data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

See Also

[rda.tune](#)

Examples

```
a <- makefolds(iris[, 5], nfolds = 5, stratified = TRUE)
table(iris[a[[1]], 5]) ## 10 values from each group
```

Greenacre's power transformation

Greenacre's power transformation

Description

Greenacre's power transformation.

Usage

```
green(x, theta)
```

Arguments

<code>x</code>	A matrix with the compositional data.
<code>theta</code>	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\theta = 0$ the log transformation is applied.

Details

Greenacre's transformation is applied to the compositional data.

Value

A matrix with the power transformed data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Greenacre, M. (2009). Power transformations in correspondence analysis. *Computational Statistics & Data Analysis*, 53(8): 3107-3116. <http://www.econ.upf.edu/~michael/work/PowerCA.pdf>

See Also

[alfa](#)

Examples

```
library(MASS)
x <- as.matrix(fg1[, 2:9])
x <- x / rowSums(x)
y1 <- green(x, 0.1)
y2 <- green(x, 0.2)
rbind( colMeans(y1), colMeans(y2) )
```

Helper Frechet mean for compositional data

Helper Frechet mean for compositional data

Description

Helper Frechet mean for compositional data.

Usage

```
frechet2(x, di, a, k1)
```

Arguments

x	A matrix with the compositional data.
di	A matrix with indices as produced by the function "dista" of the package "Rfast" or the function "nn2" of the package "RANN". Better see the details section.
a	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied and the closed geometric mean is calculated.
k1	The number of columns of "di" to exclude from the computation of the Frechet means.

Details

The power transformation is applied to the compositional data and the mean vector is calculated. Then the inverse of it is calculated and the inverse of the power transformation applied to the last vector is the Frechet mean.

What this helper function do is to speed up the Frechet mean when used in the α -k-NN regression. The α -k-NN regression computes the Frechet mean of the k nearest neighbours for a value of α and this function does exactly that. Suppose you want to predict the compositional value of some new predictors. For each predictor value you must use the Frechet mean computed at various nearest neighbours. This function performs these computations in a fast way. It is not the fastest way, yet it is a pretty fast way. This function is being called inside the function [aknn.reg](#).

Value

A list where each element contains a matrix. Each matrix contains the Frechet means computed at various nearest neighbours.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[alfa](#), [alfainv](#), [profile](#)

Examples

```
## Not run:
library(MASS)
library(Rfast)
x <- as.matrix(fg1[, 2:9])
x <- x / rowSums(x)
xnew <- x[1:10, ]
x <- x[-c(1:10), ]
k <- 2:5
di <- Rfast::dista( xnew, x, k = max(k), index = TRUE, square = TRUE )
est <- frechet2(x, di, 0.2)

## End(Not run)
```

Helper functions for the Kullback-Leibler regression
Helper functions for the Kullback-Leibler regression

Description

Helper functions for the Kullback-Leibler regression.

Usage

```
kl.compreg2(y, x, xnew = NULL, tol = 1e-07, maxiters = 50)
klcompreg.boot(y, x, der, der2, id, b1, n, p, d, tol = 1e-07, maxiters = 50)
```

Arguments

y	A matrix with the compositional data (dependent variable). Zero values are allowed. For the <code>klcompreg.boot</code> the first column is removed.
x	The predictor variable(s), they can be either continuous or categorical or both. In the <code>klcompreg.boot</code> this is the design matrix, with the ones in the first column.
xnew	If you have new data use it, otherwise leave it NULL.
tol	The tolerance value to terminate the Newton-Raphson procedure.
maxiters	The maximum number of Newton-Raphson iterations.
der	An vector to put the first derivative there.
der2	An empty matrix to put the second derivatives there, the Hessian matrix will be put here.
id	A help vector with indices.
b1	The matrix with the initial estimated coefficients.
n	The sample size
p	The number of columns of the design matrix.
d	The dimensionality of the simplex, that is the number of columns of the compositional data minus 1.

Details

These are help functions for the `kl.compreg` function. They are not to be called directly by the user.

Value

For `kl.compreg2` a list including:

iters	The nubmer of iterations required by the Newton-Raphson.
loglik	The loglikelihood.
be	The beta coefficients.

est The fitted or the predicted values (if xnew is not NULL).

For `klcompreg.boot` a list including:

loglik The loglikelihood.

be The beta coefficients.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

Murteira, Jose MR, and Joaquim JS Ramalho 2016. Regression analysis of multivariate fractional data. *Econometric Reviews* 35(4): 515-552.

See Also

[diri.reg](#), [js.compreg](#), [ols.compreg](#), [comp.reg](#)

Examples

```
library(MASS)
x <- as.vector(fgl[, 1])
y <- as.matrix(fgl[, 2:9])
y <- y / rowSums(y)
mod1 <- kl.compreg(y, x, B = 1, ncores = 1)
mod2 <- js.compreg(y, x, B = 1, ncores = 1)
```

Hotelling's multivariate version of the 1 sample t-test for Euclidean data
Hotelling's multivariate version of the 1 sample t-test for Euclidean data

Description

Hotelling's test for testing one Euclidean population mean vector.

Usage

```
hotel1T2(x, M, a = 0.05, R = 999, graph = FALSE)
```

Arguments

x	A matrix containing Euclidean data.
a	The significance level, set to 0.05 by default.
M	The hypothesized mean vector.
R	If R is 1 no bootstrap calibration is performed and the classical p-value via the F distribution is returned. If R is greater than 1, the bootstrap p-value is returned.
graph	A boolean variable which is taken into consideration only when bootstrap calibration is performed. IF TRUE the histogram of the bootstrap test statistic values is plotted.

Details

Multivariate hypothesis test for a one sample mean vector. This is the multivariate analogue of the one sample t-test. The p-value can be calculated either asymptotically or via bootstrap.

Value

A list including:

m	The sample mean vector.
info	The test statistic, the p-value, the critical value and the degrees of freedom of the F distribution (numerator and denominator). This is given if no bootstrap calibration is employed.
pvalue	The bootstrap p-value is bootstrap is employed.
runtime	The runtime of the bootstrap calibration.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

K.V. Mardia, J.T. Kent and J.M. Bibby (1979). Multivariate analysis.

See Also

[eel.test1](#), [el.test1](#), [james](#), [hotel2T2](#), [maov](#), [el.test2](#), [comp.test](#)

Examples

```
x <- Rfast::rmvnorm(100, numeric(10), diag( rep(10,0.5) ) )
hotel1T2(x, numeric(10), R = 1)
hotel1T2(x, numeric(10), R = 999, graph = TRUE)
```

Hotelling's multivariate version of the 2 sample t-test for Euclidean data
Hotelling's multivariate version of the 2 sample t-test for Euclidean data

Description

Hotelling's test for testing the equality of two Euclidean population mean vectors.

Usage

```
hotel2T2(x1, x2, a = 0.05, R = 999, graph = FALSE)
```

Arguments

x1	A matrix containing the Euclidean data of the first group.
x2	A matrix containing the Euclidean data of the second group.
a	The significance level, set to 0.05 by default.
R	If R is 1 no bootstrap calibration is performed and the classical p-value via the F distribution is returned. If R is greater than 1, the bootstrap p-value is returned.
graph	A boolean variable which is taken into consideration only when bootstrap calibration is performed. IF TRUE the histogram of the bootstrap test statistic values is plotted.

Details

Multivariate analysis of variance assuming equality of the covariance matrices. The p-value can be calculated either asymptotically or via bootstrap.

Value

A list including:

mesoi	The two mean vectors.
info	The test statistic, the p-value, the critical value and the degrees of freedom of the F distribution (numerator and denominator). This is given if no bootstrap calibration is employed.
pvalue	The bootstrap p-value is bootstrap is employed.
note	A message informing the user that bootstrap calibration has been employed.
runtime	The runtime of the bootstrap calibration.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Everitt B. (2005). An R and S-Plus Companion to Multivariate Analysis p. 139-140. Springer.

See Also

[james](#), [maov](#), [el.test2](#), [comp.test](#)

Examples

```
hotel2T2( as.matrix(iris[1:25, 1:4]), as.matrix(iris[26:50, 1:4]) )
hotel2T2( as.matrix(iris[1:25, 1:4]), as.matrix(iris[26:50, 1:4]), R = 1 )
james( as.matrix(iris[1:25, 1:4]), as.matrix(iris[26:50, 1:4]) )
james( as.matrix(iris[1:25, 1:4]), as.matrix(iris[26:50, 1:4]), R = 1 )
```

Hypothesis testing for two or more compositional mean vectors

Hypothesis testing for two or more compositional mean vectors

Description

Hypothesis testing for two or more compositional mean vectors.

Usage

```
comp.test(x, ina, test = "james", R = 0, ncores = 1, graph = FALSE)
```

Arguments

x	A matrix containing compositional data.
ina	A numerical or factor variable indicating the groups of the data.
test	This can take the values of "james" for James' test, "hotel" for Hotelling's test, "maov" for multivariate analysis of variance assuming equality of the covariance matrices, "maovjames" for multivariate analysis of variance without assuming equality of the covariance matrices. "el" for empirical likelihood or "eel" for exponential empirical likelihood.
R	This depends upon the value of the argument "test". If the test is "maov" or "maovjames", R is not taken into consideration. If test is "hotel", then R denotes the number of bootstrap resamples. If test is "james", then R can be 1 (chi-square distribution), 2 (F distribution), or more for bootstrap calibration. If test is "el", then R can be 0 (chi-square), 1 (corrected chi-sqaure), 2 (F distribution) or more for bootstrap calibration. See the help page of each test for more information.
ncores	How many to cores to use. This is taken into consideration only if test is "el" and R is more than 2.
graph	A boolean variable which is taken into consideration only when bootstrap calibration is performed. IF TRUE the histogram of the bootstrap test statistic values is plotted. This is taken into account only when R is greater than 2.

Details

The idea is to apply the α -transformation, with $\alpha = 1$, to the compositional data and then use a test to compare their mean vectors. See the help page of each test for more information. The function is visible so you can see exactly what is going on.

Value

A list including:

`result` The outcome of each test.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

- Tsagris M., Preston S. and Wood A.T.A. (2017). Nonparametric hypothesis testing for equality of means on the simplex. *Journal of Statistical Computation and Simulation*, 87(2): 406-422.
- G.S. James (1954). Tests of Linear Hypotheses in Univariate and Multivariate Analysis when the Ratios of the Population Variances are Unknown. *Biometrika*, 41(1/2): 19-43
- Krishnamoorthy K. and Yanping Xia (2006). On Selecting Tests for Equality of Two Normal Mean Vectors. *Multivariate Behavioral Research* 41(4): 533-548.
- Owen A. B. (2001). Empirical likelihood. Chapman and Hall/CRC Press.
- Owen A.B. (1988). Empirical likelihood ratio confidence intervals for a single functional. *Biometrika* 75(2): 237-249.
- Amaral G.J.A., Dryden I.L. and Wood A.T.A. (2007). Pivotal bootstrap methods for k-sample problems in directional statistics and shape analysis. *Journal of the American Statistical Association* 102(478): 695-707.
- Preston S.P. and Wood A.T.A. (2010). Two-Sample Bootstrap Hypothesis Tests for Three-Dimensional Labelled Landmark Data. *Scandinavian Journal of Statistics* 37(4): 568-587.
- Jing Bing-Yi and Andrew TA Wood (1996). Exponential empirical likelihood is not Bartlett correctable. *Annals of Statistics* 24(1): 365-369.

See Also

[maovjames](#), [maov](#), [hotel2T2](#), [el.test2](#)

Examples

```
ina <- rep(1:2, each = 50)
comp.test( as.matrix(iris[1:100, 1:4]), ina, test = "james", R = 0 )
comp.test( as.matrix(iris[1:100, 1:4]), ina, test = "hotel", R = 0 )
comp.test( as.matrix(iris[1:100, 1:4]), ina, test = "el", R = 0 )
comp.test( as.matrix(iris[1:100, 1:4]), ina, test = "eel", R = 0 )
```

Inverse of the alpha-transformation
Inverse of the α -transformation

Description

The inverse of the α -transformation.

Usage

```
alfainv(x, a, h = TRUE)
```

Arguments

- | | |
|---|---|
| x | A matrix with Euclidean data. However, they must lie within the feasible, acceptable space. See references for more information. |
| a | The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$, the inverse of the isometric log-ratio transformation is applied. |
| h | If h = TRUE this means that the multiplication with the Helmer sub-matrix will take place. It is set to TRUE by default. |

Details

The inverse of the α -transformation is applied to the data. If the data lie outside the α -space, NAs will be returned for some values.

Value

A matrix with the pairwise distances.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

- Tsagris M.T., Preston S. and Wood A.T.A. (2016). Improved classification for compositional data using the α -transformation. *Journal of Classification* (to appear). <https://arxiv.org/pdf/1506.04976v2.pdf>
- Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In *Proceedings of the 4th Compositional Data Analysis Workshop*, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also[alfa](#), [alfadist](#)**Examples**

```
library(MASS)
x <- as.matrix(fg1[1:10, 2:9])
x <- x / rowSums(x)
y <- alfa(x, 0.5)$aff
alfainv(y, 0.5)
```

James multivariate version of the t-test

James multivariate version of the t-test

Description

James test for testing the equality of two population mean vectors without assuming equality of the covariance matrices.

Usage

```
james(y1, y2, a = 0.05, R = 999, graph = FALSE)
```

Arguments

y1	A matrix containing the Euclidean data of the first group.
y2	A matrix containing the Euclidean data of the second group.
a	The significance level, set to 0.05 by default.
R	If R is 1 no bootstrap calibration is performed and the classical p-value via the F distribution is returned. If R is greater than 1, the bootstrap p-value is returned.
graph	A boolean variable which is taken into consideration only when bootstrap calibration is performed. IF TRUE the histogram of the bootstrap test statistic values is plotted.

Details

Multivariate analysis of variance without assuming equality of the covariance matrices. The p-value can be calculated either asymptotically or via bootstrap. The James test (1954) or a modification proposed by Krishnamoorthy and Yanping (2006) is implemented. The James test uses a corrected chi-square distribution, whereas the modified version uses an F distribution.

Value

A list including:

note	A message informing the user about the test used.
mesoi	The two mean vectors.
info	The test statistic, the p-value, the correction factor and the corrected critical value of the chi-square distribution if the James test has been used or, the test statistic, the p-value, the critical value and the degrees of freedom (numerator and denominator) of the F distribution if the modified James test has been used.
pvalue	The bootstrap p-value if bootstrap is employed.
runtime	The runtime of the bootstrap calibration.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

G.S. James (1954). Tests of Linear Hypotheses in Univariate and Multivariate Analysis when the Ratios of the Population Variances are Unknown. *Biometrika*, 41(1/2): 19-43

Krishnamoorthy K. and Yanping Xia. On Selecting Tests for Equality of Two Normal Mean Vectors (2006). *Multivariate Behavioral Research* 41(4): 533-548

See Also

[hotel2T2](#), [maovjames](#), [el](#), [comp.test](#)

Examples

```
james( as.matrix(iris[1:25, 1:4]), as.matrix(iris[26:50, 1:4]), R = 1 )
james( as.matrix(iris[1:25, 1:4]), as.matrix(iris[26:50, 1:4]), R = 2 )
james( as.matrix(iris[1:25, 1:4]), as.matrix(iris[26:50, 1:4]) )
hotel2T2( as.matrix(iris[1:25, 1:4]), as.matrix(iris[26:50, 1:4]) )
```

Kullback-Leibler divergence and Bhattacharyya distance between two Dirichlet distributions
*Kullback-Leibler divergence and Bhattacharyya distance between two
Dirichlet distributions*

Description

Kullback-Leibler divergence and Bhattacharyya distance between two Dirichlet distributions.

Usage

```
kl.diri(a, b, type = "KL")
```

Arguments

a	A vector with the parameters of the first Dirichlet distribution.
b	A vector with the parameters of the second Dirichlet distribution.
type	A variable indicating whether the Kullback-Leibler divergence ("KL") or the Bhattacharyya distance ("bhatt") is to be computed.

Details

Note that the order is important in the Kullback-Leibler divergence, since this is asymmetric, but not in the Bhattacharyya distance, since it is a metric.

Value

The value of the Kullback-Leibler divergence or the Bhattacharyya distance.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Ng Kai Wang, Guo-Liang Tian and Man-Lai Tang (2011). *Dirichlet and related distributions: Theory, methods and applications*. John Wiley & Sons.

See Also

[diri.est](#), [diri.nr](#)

Examples

```
library(MASS)
a <- runif(10, 0, 20)
b <- runif(10, 1, 10)
kl.diri(a, b)
kl.diri(b, a)
kl.diri(a, b, type = "bhatt")
kl.diri(b, a, type = "bhatt")
```

Log-likelihood ratio test for a Dirichlet mean vector
Log-likelihood ratio test for a Dirichlet mean vector

Description

Log-likelihood ratio test for a Dirichlet mean vector.

Usage

```
dirimean.test(x, a)
```

Arguments

x	A matrix with the compositional data. No zero values are allowed.
a	A compositional mean vector. The concentration parameter is estimated at first. If the elements do not sum to 1, it is assumed that the Dirichlet parameters are supplied.

Details

Log-likelihood ratio test is performed for the hypothesis the given vector of parameters "a" describes the compositional data well.

Value

If there are no zeros in the data, a list including:

param	A matrix with the estimated parameters under the null and the alternative hypothesis.
loglik	The log-likelihood under the alternative and the null hypothesis.
info	The value of the test statistic and its relevant p-value.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Ng Kai Wang, Guo-Liang Tian and Man-Lai Tang (2011). Dirichlet and related distributions: Theory, methods and applications. John Wiley & Sons.

See Also

[sym.test](#), [diri.nr](#), [diri.est](#), [rdiri](#), [ddiri](#)

Examples

```
x <- rdiri( 100, c(1, 2, 3) )
dirimean.test(x, c(1, 2, 3) )
dirimean.test( x, c(1, 2, 3)/6 )
```

Log-likelihood ratio test for a symmetric Dirichlet distribution
Log-likelihood ratio test for a symmetric Dirichlet distribution

Description

Log-likelihood ratio test for a symmetric Dirichlet distribution.

Usage

```
sym.test(x)
```

Arguments

x A matrix with the compositional data. No zero values are allowed.

Details

Log-likelihood ratio test is performed for the hypothesis that all Dirichlet parameters are equal.

Value

A list including:

est.par	The estimated parameters under the alternative hypothesis.
one.par	The value of the estimated parameter under the null hypothesis.
res	The loglikelihood under the alternative and the null hypothesis, the value of the test statistic, its relevant p-value and the associated degrees of freedom, which are actually the dimensionality of the simplex, $D - 1$, where D is the number of components.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Ng Kai Wang, Guo-Liang Tian and Man-Lai Tang (2011). Dirichlet and related distributions: Theory, methods and applications. John Wiley & Sons.

See Also

[diri.nr](#), [diri.est](#), [rdiri](#), [dirimean.test](#)

Examples

```
x <- rdiri( 100, c(5, 7, 1, 3, 10, 2, 4) )
sym.test(x)
x <- rdiri( 100, c(5, 5, 5, 5, 5) )
sym.test(x)
```

Mixture model selection via BIC

Mixture model selection via BIC

Description

Mixture model selection via BIC.

Usage

```
bic.mixcompnorm(x, G, type = "alr", graph = TRUE)
```

Arguments

x	A matrix with compositional data.
G	A numeric vector with the number of components, clusters, to be considered.
type	The type of transformation to be used, either additive log-ratio ("alr") or the isometric log-ratio ("ilr").
graph	A boolean variable, TRUE or FALSE specifying whether a graph should be drawn or not.

Details

The alr or the ilr-transformation is applied to the compositional data first and then mixtures of multivariate Gaussian distributions are fitted. BIC is used to decide on the optimal model and number of components.

Value

a plot with the BIC of the best model for each number of components versus the number of components. A list including:

mod	A message informing the user about the best model.
BIC	The BIC values for every possible model and number of components.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Ryan P. Browne, Aisha ElSherbiny and Paul D. McNicholas (2018). mixture: Mixture Models for Clustering and Classification. R package version 1.5.

Ryan P. Browne and Paul D. McNicholas (2014). Estimating Common Principal Components in High Dimensions. *Advances in Data Analysis and Classification*, 8(2), 217-226.

Aitchison J. (1986). *The statistical analysis of compositional data*. Chapman & Hall.

See Also

[mix.compnorm](#), [mixnorm.contour](#), [rmixcomp](#)

Examples

```
## Not run:
x <- as.matrix( iris[, 1:4] )
x <- x/ rowSums(x)
bic.mixcompnorm(x, 1:3, type = "alr", graph = FALSE)
bic.mixcompnorm(x, 1:3, type = "ilr", graph = FALSE)

## End(Not run)
```

MLE for the multivariate t distribution

MLE for the multivariate t distribution

Description

MLE of the parameters of a multivariate t distribution.

Usage

```
multivt(y, plot = FALSE)
```

Arguments

<code>y</code>	A matrix with continuous data.
<code>plot</code>	If plot is TRUE the value of the maximum log-likelihood as a function of the degrees of freedom is presented.

Details

The parameters of a multivariate t distribution are estimated. This is used by the functions [comp.den](#) and [bivt.contour](#).

Value

A list including:

center	The location estimate.
scatter	The scatter matrix estimate.
df	The estimated degrees of freedom.
loglik	The log-likelihood value.
mesos	The classical mean vector.
covariance	The classical covariance matrix.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Nadarajah, S. and Kotz, S. (2008). Estimation methods for the multivariate t distribution. Acta Applicandae Mathematicae, 102(1):99-118.

See Also

[bivt.contour](#), [comp.den](#), [rmvt](#)

Examples

```
x <- as.matrix(iris[, 1:4])
multivt(x)
```

MLE of distributions defined in the (0, 1) interval

MLE of distributions defined in the (0, 1) interval

Description

MLE of distributions defined in the (0, 1) interval.

Usage

```
beta.est(x, tol = 1e-07)
ibeta.est(x, tol = 1e-07)
logitnorm.est(x)
hsecant01.est(x, tol = 1e-07)
simplex.est(x, tol = 1e-07)
kumar.est(x, tol = 1e-07)
```

Arguments

x	A numerical vector with proportions, i.e. numbers in (0, 1) (zeros and ones are not allowed).
tol	The tolerance level up to which the maximisation stops.

Details

Maximum likelihood estimation of the parameters of the beta distribution is performed via Newton-Raphson. The distributions and hence the functions does not accept zeros. "logitnorm.mle" fits the logistic normal, hence no nnewton-Raphson is required and the "hypersecant01.mle" and "simplex.est" use the golden ratio search as is it faster than the Newton-Raphson (less computations).

Value

A list including:

iters	The number of iterations required by the Newton-Raphson.
loglik	The value of the log-likelihood.
param	The estimated parameters. In the case of "hypersecant01.est" this is called "theta" as there is only one parameter.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Manos Papadakis <papadakm95@gmail.com>

See Also

[diri.nr2](#),

Examples

```
x <- rbeta(1000, 1, 4)
beta.est(x)
ibeta.est(x)

x <- runif(1000)
hsecant01.est(x)
logitnorm.est(x)
ibeta.est(x)

x <- rbeta(1000, 2, 5)
x[sample(1:1000, 50)] <- 0
ibeta.est(x)
```

MLE of the folded model for a given value of alpha
MLE of the folded model for a given value of α

Description

MLE of the folded model for a given value of α .

Usage

```
alpha.mle(x, a)
a.mle(a, x)
```

Arguments

x	A matrix with the compositional data. No zero values are allowed.
a	A value of α .

Details

This is a function for choosing or estimating the value of α in the folded model (Tsagris and Stewart, 2019). It is called by [a.est](#).

Value

If "alpha.mle" is called, a list including:

iters	The number of iterations the EM algorithm required.
loglik	The maximized log-likelihood of the folded model.
p	The estimated probability inside the simplex of the folded model.
mu	The estimated mean vector of the folded model.
su	The estimated covariance matrix of the folded model.

If "a.mle" is called, the log-likelihood is returned only.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

Tsagris Michail and Stewart Connie, (2020). A folded model for compositional data analysis. Australian and New Zealand Journal of Statistics (to appear). <https://arxiv.org/pdf/1802.07330.pdf>
Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[alfa.profile](#), [alfa](#), [alfainv](#), [a.est](#)

Examples

```
x <- as.matrix(iris[, 1:4])
x <- x / rowSums(x)
mod <- alfa.tune(x)
mod
alpha.mle(x, mod[1])
```

Multivariate analysis of variance
Multivariate analysis of variance

Description

Multivariate analysis of variance with assuming equality of the covariance matrices.

Usage

```
maov(x, ina)
```

Arguments

<code>x</code>	A matrix containing Euclidean data.
<code>ina</code>	A numerical or factor variable indicating the groups of the data.

Details

Multivariate analysis of variance assuming equality of the covariance matrices.

Value

A list including:

<code>note</code>	A message stating whether the F or the chi-square approximation has been used.
<code>result</code>	The test statistic and the p-value.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

- Johnson and Wichern (2007, 6th Edition). Applied Multivariate Statistical Analysis p. 302-303.
- Todorov V. and Filzmoser P. (2010). Robust Statistic for the One-way MANOVA. Computational Statistics & Data Analysis 54(1):37-48.

See Also

[maovjames](#), [hotel2T2](#), [james](#), [comp.test](#)

Examples

```
maov( as.matrix(iris[,1:4]), iris[,5] )
maovjames( as.matrix(iris[,1:4]), iris[,5] )
```

Multivariate analysis of variance (James test)
Multivariate analysis of variance (James test)

Description

Multivariate analysis of variance without assuming equality of the covariance matrices.

Usage

```
maovjames(x, ina, a = 0.05)
```

Arguments

x	A matrix containing Euclidean data.
ina	A numerical or factor variable indicating the groups of the data.
a	The significance level, set to 0.005 by default.

Details

Multivariate analysis of variance without assuming equality of the covariance matrices.

Value

A vector with the next 4 elements:

test	The test statistic.
correction	The value of the correction factor.
corr.critical	The corrected critical value of the chi-square distribution.
p-value	The p-value of the corrected test statistic.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

G.S.James (1954). Tests of Linear Hypotheses in Univariate and Multivariate Analysis when the Ratios of the Population Variances are Unknown. *Biometrika*, 41(1/2): 19-43.

See Also

[maov](#), [hotel2T2](#), [james](#), [comp.test](#)

Examples

```
maov( as.matrix(iris[,1:4]), iris[,5] )
maovjames( as.matrix(iris[,1:4]), iris[,5] )
```

Multivariate kernel density estimation

Multivariate kernel density estimation

Description

Multivariate kernel density estimation.

Usage

```
mkde(x, h, thumb = "silverman")
```

Arguments

x	A matrix with Euclidean (continuous) data.
h	The bandwidth value. It can be a single value, which is turned into a vector and then into a diagonal matrix, or a vector which is turned into a diagonal matrix.
thumb	Do you want to use a rule of thumb for the bandwidth parameter? If no, leave it "none", or else put "estim" for maximum likelihood cross-validation, "scott" or "silverman" for Scott's and Silverman's rules of thumb respectively.

Details

The multivariate kernel density estimate is calculated with a (not necessarily given) bandwidth value. It is used a wrapper for the function [comp.kerncontour](#).

Value

A vector with the density estimates calculated for every vector.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Arsalane Chouaib Guidoum (2015). Kernel Estimator and Bandwidth Selection for Density and its Derivatives. The kedd R package.

M.P. Wand and M.C. Jones (1995). Kernel smoothing, pages 91-92.

B.W. Silverman (1986). Density estimation for statistics and data analysis, pages 76-78.

See Also

[mkde.tune](#), [comp.kerncontour](#)

Examples

```
mkde( as.matrix(iris[, 1:4]), thumb = "scott" )
mkde( as.matrix(iris[, 1:4]), thumb = "silverman" )
```

Multivariate linear regression
Multivariate linear regression

Description

Multivariate linear regression.

Usage

```
multivreg(y, x, plot = TRUE, xnew = NULL)
```

Arguments

y	A matrix with the Euclidean (continuous) data.
x	A matrix with the predictor variable(s), they have to be continuous.
plot	Should a plot appear or not?
xnew	If you have new data use it, otherwise leave it NULL.

Details

The classical multivariate linear regression model is obtained.

Value

A list including:

suma	A summary as produced by <code>lm</code> , which includes the coefficients, their standard error, t-values, p-values.
r.squared	The value of the R^2 for each univariate regression.
resid.out	A vector with number indicating which vectors are potential residual outliers.
x.leverage	A vector with number indicating which vectors are potential outliers in the predictor variables space.
out	A vector with number indicating which vectors are potential outliers in the residuals and in the predictor variables space.
est	The predicted values if <code>xnew</code> is not NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

K.V. Mardia, J.T. Kent and J.M. Bibby (1979). *Multivariate Analysis*. Academic Press.

See Also

[diri.reg](#), [js.compreg](#), [kl.compreg](#), [ols.compreg](#), [comp.reg](#)

Examples

```
library(MASS)
x <- as.matrix(iris[, 1:2])
y <- as.matrix(iris[, 3:4])
multivreg(y, x, plot = TRUE)
```

Multivariate normal random values simulation on the simplex

Multivariate normal random values simulation on the simplex

Description

Multivariate normal random values simulation on the simplex.

Usage

```
rcompnorm(n, m, s, type = "alr")
```

Arguments

n	The sample size, a numerical value.
m	The mean vector in R^d .
s	The covariance matrix in R^d .
type	The alr (type = "alr") or the ilr (type = "ilr") is to be used for closing the Euclidean data onto the simplex.

Details

The algorithm is straightforward, generate random values from a multivariate normal distribution in R^d and brings the values to the simplex S^d using the inverse of a log-ratio transformation.

Value

A matrix with the simulated data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[comp.den](#), [rdiri](#), [rcompt](#), [rcompsn](#)

Examples

```
x <- as.matrix(iris[, 1:2])
m <- colMeans(x)
s <- var(x)
y <- rcompnorm(100, m, s)
comp.den(y)
ternary(y)
```

Multivariate or univariate regression with compositional data in the covariates side using the alpha-
*Multivariate or univariate regression with compositional data in the
covariates side using the α -transformation*

Description

Multivariate or univariate regression with compositional data in the covariates side using the α -transformation.

Usage

```
alfa.pcr(y, x, a, k, model = "gaussian", xnew = NULL)
```

Arguments

y	A numerical vector containing the response variable values. They can be continuous, binary, discrete (counts). This can also be a vector with discrete values or a factor for the multinomial regression (model = "multinomial").
x	A matrix with the predictor variables, the compositional data.
a	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied.
k	A number at least equal to 1. How many principal components to use.
model	The type of regression model to fit. The possible values are "gaussian", "multinomial", "binomial" and "poisson".
xnew	A matrix containing the new compositional data whose response is to be predicted. If you have no new data, leave this NULL as is by default.

Details

The α -transformation is applied to the compositional data first, the first k principal component scores are calculated and used as predictor variables for a regression model. The family of distributions can be either, "normal" for continuous response and hence normal distribution, "binomial" corresponding to binary response and hence logistic regression or "poisson" for count response and poisson regression.

Value

A list tincluding:

be	If linear regression was fitted, the regression coefficients of the k principal component scores on the response variable y.
mod	If another regression model was fitted its outcome as produced in the package Rfast .
per	The percentage of variance explained by the first k principal components.
vec	The first k principal components, loadings or eigenvectors. These are useful for future prediction in the sense that one needs not fit the whole model again.
est	If the argument "xnew" was given these are the predicted or estimated values, otherwise it is NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Tsagris M. (2015). Regression analysis with compositional data containing zero values. Chilean Journal of Statistics, 6(2): 47-57. <https://arxiv.org/pdf/1508.01913v1.pdf>

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[pcr](#), [glm.pcr](#), [alfapcr.tune](#)

Examples

```
library(MASS)
y <- as.vector(fg1[, 1])
x <- as.matrix(fg1[, 2:9])
x <- x / rowSums(x)
mod <- alfa.pcr(y = y, x = x, 0.7, 1)
mod
```

Multivariate regression with compositional data
Multivariate regression with compositional data

Description

Multivariate regression with compositional data.

Usage

```
comp.reg(y, x, type = "classical", xnew = NULL, yb = NULL)
```

Arguments

y	A matrix with compositional data. Zero values are not allowed.
x	The predictor variable(s), they have to be continuous.
type	The type of regression to be used, "classical" for standard multivariate regression, or "spatial" for the robust spatial median regression. Alternatively you can type "lmit" for the fast classical multivariate regression that does not return standard errors whatsoever.
xnew	This is by default set to NULL. If you have new data whose compositional data values you want to predict, put them here.
yb	If you have already transformed the data using the additive log-ratio transformation, put it here. Otherwise leave it NULL. This is intended to be used in the function alfareg.tune in order to speed up the process.

Details

The additive log-ratio transformation is applied and then the chosen multivariate regression is implemented. The alr is easier to explain than the ilr and that is why the latter is avoided here.

Value

A list including:

runtime	The time required by the regression.
be	The beta coefficients.
seb	The standard error of the beta coefficients.
est	The fitted values of xnew if xnew is not NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

- Mardia K.V., Kent J.T., and Bibby J.M. (1979). Multivariate analysis. Academic press.
 Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[multivreg](#), [spatmed.reg](#), [js.compreg](#), [diri.reg](#)

Examples

```
library(MASS)
y <- as.matrix(iris[, 1:3])
y <- y / rowSums(y)
x <- as.vector(iris[, 4])
mod1 <- comp.reg(y, x)
mod2 <- comp.reg(y, x, type = "spatial")
```

Multivariate skew normal random values simulation on the simplex

Multivariate skew normal random values simulation on the simplex

Description

Multivariate skew normal random values simulation on the simplex.

Usage

```
rcompsn(n, xi, Omega, alpha, dp = NULL, type = "alr")
```

Arguments

n	The sample size, a numerical value.
xi	A numeric vector of length d representing the location parameter of the distribution.
Omega	A $d \times d$ symmetric positive-definite matrix of dimension.
alpha	A numeric vector which regulates the slant of the density.
dp	A list with three elements, corresponding to xi, Omega and alpha described above. The default value is FALSE. If dp is assigned, individual parameters must not be specified.
type	The alr (type = "alr") or the ilr (type = "ilr") is to be used for closing the Euclidean data onto the simplex.

Details

The algorithm is straightforward, generate random values from a multivariate t distribution in R^d and brings the values to the simplex S^d using the inverse of a log-ratio transformation.

Value

A matrix with the simulated data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Azzalini, A. and Dalla Valle, A. (1996). The multivariate skew-normal distribution. *Biometrika*, 83(4): 715-726.

Azzalini, A. and Capitanio, A. (1999). Statistical applications of the multivariate skew normal distribution. *Journal of the Royal Statistical Society Series B*, 61(3):579-602. Full-length version available from <http://arXiv.org/abs/0911.2093>

Aitchison J. (1986). *The statistical analysis of compositional data*. Chapman & Hall.

See Also

[comp.den](#), [rdiri](#), [rcompnorm](#), [rmvt](#)

Examples

```
x <- as.matrix(iris[, 1:2])
par <- sn::msn.mle(y = x)$dp
y <- rcompsn(100, dp = par)
comp.den(y, dist = "skewnorm")
ternary(y)
```

Multivariate t random values simulation on the simplex

Multivariate t random values simulation on the simplex

Description

Multivariate t random values simulation on the simplex.

Usage

```
rcompt(n, m, s, dof, type = "alr")
```

Arguments

<code>n</code>	The sample size, a numerical value.
<code>m</code>	The mean vector in R^d .
<code>s</code>	The covariance matrix in R^d .
<code>dof</code>	The degrees of freedom.
<code>type</code>	The alr (type = "alr") or the ilr (type = "ilr") is to be used for closing the Euclidean data onto the simplex.

Details

The algorithm is straightforward, generate random values from a multivariate t distribution in R^d and brings the values to the simplex S^d using the inverse of a log-ratio transformation.

Value

A matrix with the simulated data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[comp.den](#), [rdiri](#), [rcompnorm](#), [rmvt](#)

Examples

```
x <- as.matrix(iris[, 1:2])
m <- Rfast::colmeans(x)
s <- var(x)
y <- rcompt(100, m, s, 10)
comp.den(y, dist = "t")
ternary(y)
```

Non linear least squares regression for compositional data

Non linear least squares regression for compositional data

Description

Non linear least squares regression for compositional data.

Usage

```
ols.compreg(y, x, B = 1, ncores = 1, xnew = NULL)
```

Arguments

y	A matrix with the compositional data (dependent variable). Zero values are allowed.
x	The predictor variable(s), they have to be continuous.
B	If B is greater than 1 bootstrap estimates of the standard error are returned. If B=1, no standard errors are returned.
ncores	If ncores is 2 or more parallel computing is performed. This is to be used for the case of bootstrap. If B=1, this is not taken into consideration.
xnew	If you have new data use it, otherwise leave it NULL.

Details

The ordinary least squares between the observed and the fitted compositional data is adopted as the objective function. This involves numerical optimization since the relationship is non linear. There is no log-likelihood.

Value

A list including:

runtime	The time required by the regression.
beta	The beta coefficients.
seb	The standard error of the beta coefficients, if bootstrap is chosen, i.e. if B > 1.
est	The fitted of xnew if xnew is not NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Murteira, Jose MR, and Joaquim JS Ramalho 2016. Regression analysis of multivariate fractional data. *Econometric Reviews* 35(4): 515-552.

See Also

[diri.reg](#), [js.compreg](#), [kl.compreg](#), [comp.reg](#), [comp.reg](#), [alfa.reg](#)

Examples

```
library(MASS)
x <- as.vector(fgl[, 1])
y <- as.matrix(fgl[, 2:9])
y <- y / rowSums(y)
mod1 <- ols.compreg(y, x, B = 1, ncores = 1)
mod2 <- js.compreg(y, x, B = 1, ncores = 1)
```

Perturbation operation

Perturbation operation

Description

Perturbation operation.

Usage

```
perturbation(x, y, oper = "+")
```

Arguments

x	A matrix with the compositional data.
y	Either a matrix with compositional data or a vector with compositional data. In either case, the data may not be compositional data, as long as they non negative.
oper	For the summation this must be "*" and for the negation it must be "/". According to Aitchison (1986), multiplication is equal to summation in the log-space, and division is equal to negation.

Details

This is the perturbation operation defined by Aitchison (1986).

Value

A matrix with the perturbed compositional data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[power](#)

Examples

```
x <- as.matrix(iris[1:15, 1:4])
y <- as.matrix(iris[21:35, 1:4])
perturbation(x, y)
perturbation(x, y[1, ])
```

Power operation

Power operation

Description

Power operation.

Usage

```
pow(x, a)
```

Arguments

x	A matrix with the compositional data.
a	Either a vector with numbers of a single number.

Details

This is the power operation defined by Aitchison (1986). It is also the starting point of the α -transformation.

Value

A matrix with the power transformed compositional data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

- Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.
- Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <http://arxiv.org/pdf/1106.1451.pdf>

See Also

[perturbation,alfa](#)

Examples

```
x <- as.matrix(iris[1:15, 1:4])
a <- runif(1)
pow(x, a)
```

Principal component generalised linear models

Principal component generalised linear models

Description

Principal component generalised linear models.

Usage

```
pcr(y, x, k = 1, xnew = NULL)
glm.pcr(y, x, k = 1, xnew = NULL)
```

Arguments

y	A numerical vector, a real values vector or a numeric vector with 0 and 1 (binary) or a vector with discrete (count) data.
x	A matrix with the predictor variable(s), they have to be continuous.
k	A number greater than or equal to 1. How many principal components to use. In the case of "pcr" this can be a single number or a vector. In the second case you get results for the sequence of principal components.
xnew	If you have new data use it, otherwise leave it NULL.

Details

Principal component regression is performed with linear, binary logistic or Poisson regression, depending on the nature of the response variable. The principal components of the cross product of the independent variables are obtained and classical regression is performed. This is used in the function [alfa.pcr](#).

Value

A list including:

be	The beta coefficients of the predictor variables computed via the principal components if "pcr" is used.
model	The summary of the logistic or Poisson regression model.
per	The percentage of variance of the predictor variables retained by the k principal components.
vec	The principal components, the loadings.
est	The fitted or the predicted values (if xnew is not NULL).

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Aguilera A.M., Escabias M. and Valderrama M.J. (2006). Using principal components for estimating logistic regression with high-dimensional multicollinear data. *Computational Statistics & Data Analysis* 50(8): 1905-1924.

Jolliffe I.T. (2002). *Principal Component Analysis*.

See Also

[alfa.pcr](#), [alfapcr.tune](#)

Examples

```
library(MASS)
x <- as.matrix(fg1[, 2:9])
y <- as.vector(fg1[, 1])
mod1 <- pcr(y, x, 1)
mod2 <- pcr(y, x, 2)
mod <- pcr(y, x, k = 1:4) ## many results at once

x <- as.matrix(iris[, 1:4])
y <- rbinom(150, 1, 0.6)
mod <- glm.pcr(y, x, k = 1)
```

Projection pursuit regression for compositional data
Projection pursuit regression for compositional data

Description

Projection pursuit regression for compositional data.

Usage

```
comp.ppr(y, x, nterms = 3, type = "alr", xnew = NULL, yb = NULL )
```

Arguments

y	A matrix with the compositional data.
x	A matrix with the continuous predictor variables or a data frame including categorical predictor variables.
nterms	The number of terms to include in the final model.
type	Either "alr" or "ilr" corresponding to the additive or the isometric log-ratio transformation respectively.
xnew	If you have new data use it, otherwise leave it NULL.
yb	If you have already transformed the data using a log-ratio transformation put it here. Othewise leave it NULL.

Details

This is the standard projection pursuit. See the built-in funciton "ppr" for more details.

Value

A list includign:

runtime	The runtime of the regression.
mod	The produced model as returned by the function "ppr".
est	The fitted values of xnew if xnew is not NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Friedman, J. H. and Stuetzle, W. (1981). Projection pursuit regression. Journal of the American Statistical Association, 76, 817-823. doi: 10.2307/2287576.

See Also

[compknn.tune,rda,alfa](#)

Examples

```
x <- as.matrix( iris[, 1:4] )
x <- x/ rowSums(x)
ina <- iris[, 5]
mod <- comp.knn(x, x, ina, a = 1, k = 5)
table(ina, mod)
mod2 <- alfa.knn(x, x, ina, a = 1, k = 5)
table(ina, mod2)
```

Quasi binomial regression for proportions

Quasi binomial regression for proportions

Description

Quasi binomial regression for proportions.

Usage

```
propreg(y, x, varb = "quasi", tol = 1e-07, maxiters = 100)
propregs(y, x, varb = "quasi", tol = 1e-07, logged = FALSE, maxiters = 100)
```

Arguments

<code>y</code>	A numerical vector proportions. 0s and 1s are allowed.
<code>x</code>	For the "propreg" a matrix with data, the predictor variables. This can be a matrix or a data frame. For the "propregs" this must be a numerical matrix, where each columns denotes a variable.
<code>tol</code>	The tolerance value to terminate the Newton-Raphson algorithm. This is set to 10^{-9} by default.
<code>varb</code>	The type of estimate to be used in order to estimate the covariance matrix of the regression coefficients. There are two options, either "quasi" (default value) or "glm". See the references for more information.
<code>logged</code>	Should the p-values be returned (FALSE) or their logarithm (TRUE)?
<code>maxiters</code>	The maximum number of iterations before the Newton-Raphson is terminated automatically.

Details

We are using the Newton-Raphson, but unlike R's built-in function "glm" we do no checks and no extra calculations, or whatever. Simply the model. The "propregs" is to be used for very many univariate regressions. The "x" is a matrix in this case and the significance of each variable (column of the matrix) is tested. The function accepts binary responses as well (0 or 1).

Value

For the "propreg" function a list including:

<code>iters</code>	The number of iterations required by the Newton-Raphson.
<code>varb</code>	The covariance matrix of the regression coefficients.
<code>phi</code>	The phi parameter is returned if the input argument "varb" was set to "glm", otherwise this is NULL.
<code>info</code>	A table similar to the one produced by "glm" with the estimated regression coefficients, their standard error, Wald test statistic and p-values.

For the "propregs" a two-column matrix with the test statistics (Wald statistic) and the associated p-values (or their loggarithm).

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Manos Papadakis <papadakm95@gmail.com>.

References

Papke L. E. & Wooldridge J. (1996). Econometric methods for fractional response variables with an application to 401(K) plan participation rates. *Journal of Applied Econometrics*, 11(6): 619–632.

McCullagh, Peter, and John A. Nelder. *Generalized linear models*. CRC press, USA, 2nd edition, 1989.

See Also

[anova_propreg](#), [univglms](#), [score.glms](#), [logistic_only](#)

Examples

```
y <- rbeta(100, 1, 4)
x <- matrix(rnorm(100 * 3), ncol = 3)
a <- propreg(y, x)
y <- rbeta(100, 1, 4)
x <- matrix(rnorm(400 * 100), ncol = 400)
b <- propregs(y, x)
mean(b[, 2] < 0.05)
```

Regression with compositional data using the alpha-transformation

Regression with compositional data using the α -transformation

Description

Regression with compositional data using the α -transformation.

Usage

```
alfa.reg(y, x, a, xnew = NULL, yb = NULL, seb = FALSE)
```

Arguments

y	A matrix with the compositional data.
x	A matrix with the continuous predictor variables or a data frame including categorical predictor variables.
a	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied and the solution exists in a closed form, since it the classical multivariate regression.
xnew	If you have new data use it, otherwise leave it NULL.
yb	If you have already transformed the data using the α -transformation with the same α as given in the argument "a", put it here. Otherwise leave it NULL. This is intended to be used in the function alfareg.tune in order to speed up the process. The time difference in that function is small for small samples. But, if you have a few thousands and or a few more components, there will be bigger differences.
seb	Do you want the standard error of the coefficients to be returned? In the alfareg.tune function this extra computation that is avoided can save some time.

Details

The α -transformation is applied to the compositional data first and then multivariate regression is applied. This involves numerical optimisation.

Value

A list including:

runtime	The time required by the regression.
be	The beta coefficients.
seb	The standard error of the beta coefficients.
est	The fitted values for xnew if xnew is not NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Tsagris M. (2015). Regression analysis with compositional data containing zero values. Chilean Journal of Statistics, 6(2): 47-57. <https://arxiv.org/pdf/1508.01913v1.pdf>

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

Mardia K.V., Kent J.T., and Bibby J.M. (1979). Multivariate analysis. Academic press.

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[alfareg.tune](#), [diri.reg](#), [js.compreg](#), [kl.compreg](#), [ols.compreg](#), [comp.reg](#)

Examples

```
library(MASS)
x <- as.vector(fgl[1:40, 1])
y <- as.matrix(fgl[1:40, 2:9])
y <- y / rowSums(y)
mod <- alfa.reg(y, x, 0.2)
```

Regularised and flexible discriminant analysis for compositional data using the alpha-transformation
*Regularised and flexible discriminant analysis for compositional data
using the α -transformation*

Description

Regularised and flexible discriminant analysis for compositional data using the α -transformation.

Usage

```
alfa.rda(xnew, x, ina, a, gam = 1, del = 0)
alfa.fda(xnew, x, ina, a)
```

Arguments

xnew	A matrix with the new compositional data whose group is to be predicted. Zeros are allowed, but you must be careful to choose strictly positive values of α .
x	A matrix with the available compositional data. Zeros are allowed, but you must be careful to choose strictly positive values of α .
ina	A group indicator variable for the available data.
a	The value of α for the α -transformation.
gam	This is a number between 0 and 1. It is the weight of the pooled covariance and the diagonal matrix.
del	This is a number between 0 and 1. It is the weight of the LDA and QDA.

Details

For the `alfa.rda`, the covariance matrix of each group is calculated and then the pooled covariance matrix. The spherical covariance matrix consists of the average of the pooled variances in its diagonal and zeros in the off-diagonal elements. `gam` is the weight of the pooled covariance matrix and `1-gam` is the weight of the spherical covariance matrix, $S_a = \text{gam} * S_p + (1-\text{gam}) * s_p$. Then it is a compromise between LDA and QDA. `del` is the weight of S_a and `1-del` the weight of each group covariance group. This function is a wrapper for [alfa.rda](#).

For the `alfa.fda` a flexible discriminant analysis is performed. See the R package **fda** for more details.

Value

For the `alfa.rda` a list including:

prob	The estimated probabilities of the new data of belonging to each group.
scores	The estimated scores of the new data of each group.
est	The estimated group membership of the new data.

For the `alfa.fda` a list including:

mod	A fda object as returned by the command <code>fda</code> of the R package <code>mda</code> .
est	The estimated group membership of the new data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Friedman Jerome, Trevor Hastie and Robert Tibshirani (2009). The elements of statistical learning, 2nd edition. Springer, Berlin.

Tsagris Michail, Simon Preston and Andrew T.A. Wood (2016). Improved classification for compositional data using the α -transformation. Journal of classification, 33(2): 243-261. <https://arxiv.org/pdf/1106.1451.pdf>

Hastie, Tibshirani and Buja (1994). Flexible Discriminant Analysis by Optimal Scoring. Journal of the American Statistical Association, 89(428):1255-1270.

See Also

[rda,alfa](#)

Examples

```
x <- as.matrix(iris[, 1:4])
x <- x / rowSums(x)
ina <- iris[, 5]
mod <- alfa.rda(x, x, ina, 0)
table(ina, mod$est)
mod2 <- alfa.fda(x, x, ina, 0)
table(ina, mod2$est)
```

Regularised discriminant analysis for Euclidean data
Regularised discriminant analysis for Euclidean data

Description

Regularised discriminant analysis for Euclidean data.

Usage

```
rda(xnew, x, ina, gam = 1, del = 0)
```

Arguments

xnew	A matrix with the new data whose group is to be predicted. They have to be continuous.
x	A matrix with the available data. They have to be continuous.
ina	A group indicator variable for the available data.
gam	This is a number between 0 and 1. It is the weight of the pooled covariance and the diagonal matrix.
del	This is a number between 0 and 1. It is the weight of the LDA and QDA.

Details

The covariance matrix of each group is calculated and then the pooled covariance matrix. The spherical covariance matrix consists of the average of the pooled variances in its diagonal and zeros in the off-diagonal elements. `gam` is the weight of the pooled covariance matrix and `1-gam` is the weight of the spherical covariance matrix, $S_a = \text{gam} * S_p + (1-\text{gam}) * s_p$. Then it is a compromise between LDA and QDA. `del` is the weight of S_a and `1-del` the weight of each group covariance group. This function is a wrapper for [alfa.rda](#).

Value

A list including:

<code>prob</code>	The estimated probabilities of the new data of belonging to each group.
<code>scores</code>	The estimated scores of the new data of each group.
<code>est</code>	The estimated group membership of the new data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Friedman Jerome, Trevor Hastie and Robert Tibshirani (2009). The elements of statistical learning, 2nd edition. Springer, Berlin

Tsagris Michail, Simon Preston and Andrew TA Wood (2016). Improved classification for compositional data using the α -transformation. Journal of classification, 33(2): 243-261. <http://arxiv.org/pdf/1106.1451.pdf>

See Also

[rda.tune](#), [alfa](#)

Examples

```
x <- as.matrix(iris[, 1:4])
ina <- iris[, 5]
mod <- rda(x, x, ina)
table(ina, mod$est)
```

Ridge regression	<i>Ridge regression</i>
------------------	-------------------------

Description

Ridge regression.

Usage

```
ridge.reg(y, x, lambda, B = 1, xnew = NULL)
```

Arguments

y	A real valued vector. If it contains percentages, the logit transformation is applied.
x	A matrix with the predictor variable(s), they have to be continuous.
lambda	The value of the regularisation parameter λ .
B	If B = 1 (default value) no bootstrap is performed. Otherwise bootstrap standard errors are returned.
xnew	If you have new data whose response value you want to predict put it here, otherwise leave it as is.

Details

This is used in the function [alfa.ridge](#). There is also a built-in function available from the MASS library, called [lm.ridge](#).

Value

A list including:

beta	The beta coefficients.
seb	The standard error of the coefficients. If B > 1 the bootstrap standard errors will be returned.
est	The fitted or the predicted values (if xnew is not NULL).

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Hoerl A.E. and R.W. Kennard (1970). Ridge regression: Biased estimation for nonorthogonal problems. *Technometrics*, 12(1): 55-67.

Brown P. J. (1994). *Measurement, Regression and Calibration*. Oxford Science Publications.

See Also

[ridge.tune](#), [alfa.ridge](#), [ridge.plot](#)

Examples

```
y <- as.vector(iris[, 1])
x <- as.matrix(iris[, 2:4])
mod1 <- ridge.reg(y, x, lambda = 0.1)
mod2 <- ridge.reg(y, x, lambda = 0)
```

Ridge regression plot *Ridge regression plot*

Description

A plot of the regularised regression coefficients is shown.

Usage

```
ridge.plot(y, x, lambda = seq(0, 5, by = 0.1) )
```

Arguments

<code>y</code>	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using the logit transformation. In any case, they must be continuous only.
<code>x</code>	A numeric matrix containing the continuous variables. Rows are samples and columns are features.
<code>lambda</code>	A grid of values of the regularisation parameter λ .

Details

For every value of λ the coefficients are obtained. They are plotted versus the λ values.

Value

A plot with the values of the coefficients as a function of λ .

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <gioathineou@gmail.com> and Michail Tsagris <mtsagris@uoc.gr>

References

- Hoerl A.E. and R.W. Kennard (1970). Ridge regression: Biased estimation for nonorthogonal problems. *Technometrics*, 12(1): 55-67.
- Brown P. J. (1994). *Measurement, Regression and Calibration*. Oxford Science Publications.

See Also

[ridge.reg](#), [ridge.tune](#), [alfa.ridge](#), [alfaridge.plot](#)

Examples

```
y <- as.vector(iris[, 1])
x <- as.matrix(iris[, 2:4])
ridge.plot(y, x, lambda = seq(0, 2, by = 0.1) )
```

Ridge regression with compositional data in the covariates side using the alpha-transformation
*Ridge regression with compositional data in the covariates side using
the α -transformation*

Description

Ridge regression with compositional data in the covariates side using the α -transformation.

Usage

```
alfa.ridge(y, x, a, lambda, B = 1, xnew = NULL)
```

Arguments

- | | |
|--------|--|
| y | A numerical vector containing the response variable values. If they are percentages, they are mapped onto R using the logit transformation. |
| x | A matrix with the predictor variables, the compositional data. Zero values are allowed, but you must be careful to choose strictly positive values of α . |
| a | The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied. |
| lambda | The value of the regularisation parameter, λ . |
| B | If $B > 1$ bootstrap estimation of the standard errors is implemented. |
| xnew | A matrix containing the new compositional data whose response is to be predicted. If you have no new data, leave this NULL as is by default. |

Details

The α -transformation is applied to the compositional data first and then ridge components regression is performed.

Value

The output of the [ridge.reg](#).

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Tsagris M. (2015). Regression analysis with compositional data containing zero values. Chilean Journal of Statistics, 6(2): 47-57. <https://arxiv.org/pdf/1508.01913v1.pdf>

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[ridge.reg](#), [alfaridge.tune](#), [alfaridge.plot](#)

Examples

```
library(MASS)
y <- as.vector(fgl[, 1])
x <- as.matrix(fgl[, 2:9])
x <- x/ rowSums(x)
mod1 <- alfa.ridge(y, x, a = 0.5, lambda = 0.1, B = 1, xnew = NULL)
mod2 <- alfa.ridge(y, x, a = 0.5, lambda = 1, B = 1, xnew = NULL)
```

Ridge regression with the alpha-transformation plot

Ridge regression plot

Description

A plot of the regularised regression coefficients is shown.

Usage

```
alfaridge.plot(y, x, a, lambda = seq(0, 5, by = 0.1) )
```

Arguments

y	A numeric vector containing the values of the target variable. If the values are proportions or percentages, i.e. strictly within 0 and 1 they are mapped into R using the logit transformation. In any case, they must be continuous only.
x	A numeric matrix containing the continuous variables.
a	The value of the α -transformation. It has to be between -1 and 1. If there are zero values in the data, you must use a strictly positive value.
lambda	A grid of values of the regularisation parameter λ .

Details

For every value of λ the coefficients are obtained. They are plotted versus the λ values.

Value

A plot with the values of the coefficients as a function of λ .

Author(s)

Michail Tsagris

R implementation and documentation: Giorgos Athineou <gioathineou@gmail.com> and Michail Tsagris <mtsagris@uoc.gr>

References

Hoerl A.E. and R.W. Kennard (1970). Ridge regression: Biased estimation for nonorthogonal problems. *Technometrics*, 12(1): 55-67.

Brown P. J. (1994). *Measurement, Regression and Calibration*. Oxford Science Publications.

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In *Proceedings of the 4th Compositional Data Analysis Workshop*, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[ridge.plot](#), [alfa.ridge](#)

Examples

```
library(MASS)
y <- as.vector(fgl[, 1])
x <- as.matrix(fgl[, 2:9])
x <- x / rowSums(x)
alfaridge.plot(y, x, a = 0.5, lambda = seq(0, 5, by = 0.1) )
```

Simulation of compositional data from Gaussian mixture models

Simulation of compositional data from Gaussian mixture models

Description

Simulation of compositional data from Gaussian mixture models.

Usage

```
rmixcomp(n, prob, mu, sigma, type = "alr")
```

Arguments

n	The sample size
prob	A vector with mixing probabilities. Its length is equal to the number of clusters.
mu	A matrix where each row corresponds to the mean vector of each cluster.
sigma	An array consisting of the covariance matrix of each cluster.
type	Should the additive ("type=alr") or the isometric (type="ilr") log-ration be used? The default value is for the additive log-ratio transformation.

Details

A sample from a multivariate Gaussian mixture model is generated.

Value

A list including:

id	A numeric variable indicating the cluster of simulated vector.
x	A matrix containing the simulated compositional data. The number of dimensions will be + 1.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Ryan P. Browne, Aisha ElSherbiny and Paul D. McNicholas (2015). R package mixture: Mixture Models for Clustering and Classification.

See Also

[mix.compnorm](#), [bic.mixcompnorm](#)

Examples

```
p <- c(1/3, 1/3, 1/3)
mu <- matrix(nrow = 3, ncol = 4)
s <- array( dim = c(4, 4, 3) )
x <- as.matrix(iris[, 1:4])
ina <- as.numeric(iris[, 5])
mu <- rowsum(x, ina) / 50
s[, , 1] <- cov(x[ina == 1, ])
s[, , 2] <- cov(x[ina == 2, ])
s[, , 3] <- cov(x[ina == 3, ])
y <- rmixcomp(100, p, mu, s, type = "alr")
```

Simulation of compositional data from the Flexible Dirichlet distribution
Simulation of compositional data from the Flexible Dirichlet distribution

Description

Simulation of compositional data from the Flexible Dirichlet distribution.

Usage

```
rfd(n, alpha, prob, tau)
```

Arguments

n	The sample size.
alpha	A vector of the non-negative alpha parameters.
prob	A vector of the clusters' probabilities that must sum to one.
tau	The non-negative scalar tau parameter.

Details

For more information see the references.

Value

A matrix with compositional data.

Author(s)

Michail Tsagris ported from the R package FlexDir. <mtsagris@uoc.gr>.

References

Ongaro, A. and Migliorati, S. (2013) A generalization of the Dirichlet distribution. *Journal of Multivariate Analysis*, 114, 412–426.

Migliorati, S., Ongaro, A. and Monti, G. S. (2016) A structured Dirichlet mixture model for compositional data: inferential and applicative issues. *Statistics and Computing*, 1–21.

See Also

[fd.est](#), [fd.density](#)

Examples

```
alpha <- c(12, 11, 10)
prob <- c(0.25, 0.25, 0.5)
x <- rfd(100, alpha, prob, 7)
```

Simulation of compositional data from the folded normal distribution
Simulation of compositional data from the folded model normal distribution

Description

Simulation of compositional data from the folded model normal distribution.

Usage

```
rfolded(n, mu, su, a)
```

Arguments

n	The sample size.
mu	The mean vector.
su	The covariance matrix.
a	The value of α .

Details

A sample from the folded model is generated.

Value

A matrix with compositional data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

Tsagris Michail and Stewart Connie, (2020). A folded model for compositional data analysis. Australian and New Zealand Journal of Statistics (to appear). <https://arxiv.org/pdf/1802.07330.pdf>

See Also

[alfa](#), [alpha.mle](#), [a.est](#)

Examples

```
s <- c(0.1490676523, -0.4580818209, 0.0020395316, -0.0047446076, -0.4580818209,
1.5227259250, 0.0002596411, 0.0074836251, 0.0020395316, 0.0002596411,
0.0365384838, -0.0471448849, -0.0047446076, 0.0074836251, -0.0471448849,
0.0611442781)
s <- matrix(s, ncol = 4)
m = c(1.715, 0.914, 0.115, 0.167)
x = rfolded(100, m, s, 0.5)
a.est(x)
```

Spatial median regression

Spatial median regression

Description

Spatial median regression with Euclidean data.

Usage

```
spatmed.reg(y, x, xnew = NULL, tol = 1e-07, ses = FALSE)
```

Arguments

<code>y</code>	A matrix with the compositional data. Zero values are not allowed.
<code>x</code>	The predictor variable(s), they have to be continuous.
<code>xnew</code>	If you have new data use it, otherwise leave it NULL.
<code>tol</code>	The threshold upon which to stop the iterations of the Newton-Rapshon algorithm.
<code>ses</code>	If you want to extract the standard errors of the parameters, set this to TRUE. Be careful though as this can slow down the algorithm dramatically. In a run example with 10,000 observations and 10 variables for <code>y</code> and 30 for <code>x</code> , when <code>ses = FALSE</code> the algorithm can take 0.20 seconds, but when <code>ses = TRUE</code> it can go up to 140 seconds.

Details

The objective function is the minimization of the sum of the absolute residuals. It is the multivariate generalization of the median regression. This function is used by [comp.reg](#).

Value

A list including:

<code>iter</code>	The number of iterations that were required.
<code>runtime</code>	The time required by the regression.
<code>be</code>	The beta coefficients.
<code>seb</code>	The standard error of the beta coefficients is returned if <code>ses=TRUE</code> and <code>NULL</code> otherwise.
<code>est</code>	The fitted of <code>xnew</code> if <code>xnew</code> is not <code>NULL</code> .

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Biman Chakraborty (2003) On multivariate quantile regression. Journal of Statistical Planning and Inference http://www.stat.nus.edu.sg/export/sites/dsap/research/documents/tr01_2000.pdf

See Also

[multivreg](#), [comp.reg](#), [alfa.reg](#), [js.compreg](#), [diri.reg](#)

Examples

```
library(MASS)
x <- as.matrix(iris[, 3:4])
y <- as.matrix(iris[, 1:2])
mod1 <- spatmed.reg(y, x)
mod2 <- multivreg(y, x, plot = FALSE)
```

Ternary diagram	<i>Ternary diagram</i>
-----------------	------------------------

Description

Ternary diagram.

Usage

```
ternary(x, means = TRUE, pca = FALSE)
```

Arguments

x	A matrix with the compositional data.
means	A boolean variable. Should the closed geometric mean and the arithmetic mean appear (TRUE) or not (FALSE)?.
pca	Should the first PCA calculated Aitchison (1983) described appear? If yes, then this should be TRUE, or FALSE otherwise.

Details

The first PCA is calculated using the centred log-ratio transformation as Aitchison (1983, 1986) suggested. If the data contain zero values, the first PCA will not be plotted. There are two ways to create a ternary graph. The one I used here, where each edge is equal to 1 and the one Aitchison (1986) uses. For every given point, the sum of the distances from the edges is equal to 1. Zeros in the data appear with green circles in the triangle and you will also see NaN in the closed geometric mean.

Value

The ternary plot and a matrix with the closed geometric and the simple arithmetic mean vector.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Aitchison, J. (1983). Principal component analysis of compositional data. *Biometrika* 70(1):57-65.
Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[comp.den](#), [alfa](#), [diri.contour](#), [comp.kerncontour](#)

Examples

```
library(MASS)
x <- as.matrix(fgl[, 2:4])
ternary(x, means = FALSE)
x <- as.matrix(iris[, 1:3])
ternary(x, pca = TRUE)
```

The additive log-ratio transformation and its inverse

The additive log-ratio transformation and its inverse

Description

The additive log-ratio transformation and its inverse.

Usage

```
alr(x)
alrinv(y)
```

Arguments

x A numerical matrix with the compositional data.
y A numerical matrix with data to be closed into the simplex.

Details

The additive log-ratio transformation with the first component being the common divisor is applied. The inverse of this transformation is also available.

Value

A matrix with the alr transformed data (if alr is used) or with the compositional data (if the alrinv is used).

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[alr](#), [link{alfainv}](#) [alr.profile](#), [alr.tune](#)

Examples

```
library(MASS)
x <- as.matrix(fg1[, 2:9])
x <- x / rowSums(x)
y <- alr(x)
x1 <- alrinv(y)
```

The alpha-distance *The α -distance*

Description

This is the Euclidean (or Manhattan) distance after the α -transformation has been applied.

Usage

```
alfadist(x, a, type = "euclidean", square = FALSE)
alfadista(xnew, x, a, type = "euclidean", square = FALSE)
```

Arguments

xnew	A matrix or a vector with new compositional data.
x	A matrix with the compositional data.
a	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$, the isometric log-ratio transformation is applied.
type	Which type distance do you want to calculate after the α -transformation, "euclidean", or "manhattan".
square	In the case of the Euclidean distance, you can choose to return the squared distance by setting this TRUE.

Details

The α -transformation is applied to the compositional data first and then the Euclidean or the Manhattan distance is calculated.

Value

For "alfadist" a matrix including the pairwise distances of all observations or the distances between xnew and x. For "alfadista" a matrix including the pairwise distances of all observations or the distances between xnew and x.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

- Tsagris M.T., Preston S. and Wood A.T.A. (2016). Improved classification for compositional data using the α -transformation. *Journal of Classification*. 33(2):243–261. <https://arxiv.org/pdf/1506.04976v2.pdf>
- Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In *Proceedings of the 4th Compositional Data Analysis Workshop*, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[alfa](#), [alfainv](#), [alfa.reg](#), [esov](#)

Examples

```
library(MASS)
x <- as.matrix(fgl[1:20, 2:9])
x <- x / rowSums(x)
alfadist(x, 0.1)
alfadist(x, 1)
```

The α -k-NN regression for compositional response data

The α -k-NN regression for compositional response data

Description

The α -k-NN regression for compositional response data.

Usage

```
aknn.reg(xnew, y, x, a = seq(0.1, 1, by = 0.1), k = 2:10,
apostasi = "euclidean", rann = FALSE)
```

Arguments

xnew	A matrix with the new predictor variables whose compositions are to be predicted.
y	A matrix with the compositional response data. Zeros are allowed.
x	A matrix with the available predictor variables.
a	The value of α . As zero values in the compositional data are allowed, you must be careful to choose strictly positive values of α . If negative values are passed, the positive ones are used only.
k	The number of nearest neighbours to consider. It can be a single number or a vector.
apostasi	The type of distance to use, either "euclidean" or "manhattan".
rann	If you have large scale datasets and want a faster k-NN search, you can use kd-trees implemented in the R package "RANN". In this case you must set this argument equal to TRUE. Note however, that in this case, the only available distance is by default "euclidean".

Details

The α -k-NN regression for compositional response variables is applied.

Value

A list with the estimated compositional response data for each value of α and k.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Michail Tsagris, Abdulaziz Alenazi and Connie Stewart (2020). The alpha-k-NN- regression for compositional data. <https://arxiv.org/pdf/2002.05137.pdf>

See Also

[alfa.reg](#), [comp.ppr](#), [comp.reg](#), [kl.compreg](#)

Examples

```
y <- as.matrix( iris[, 1:3] )
y <- y / rowSums(y)
x <- iris[, 4]
mod <- aknn.reg(x, y, x, a = c(0.4, 0.5), k = 2:3, apostasi = "euclidean")
```

The alpha-k-NN regression with compositional predictor variables
The α -k-NN regression with compositional predictor variables

Description

The α -k-NN regression with compositional predictor variables.

Usage

```
alfa.knn.reg(xnew, y, x, a = 1, k = 2:10, apostasi = "euclidean", method = "average")
```

Arguments

xnew	A matrix with the new compositional predictor variables whose response is to be predicted. Zeros are allowed.
y	The response variable, a numerical vector.
x	A matrix with the available compositional predictor variables. Zeros are allowed.
a	A single value of α . As zero values in the compositional data are allowed, you must be careful to choose strictly positive values of α . If negative values are passed, the positive ones are used only. If the data are already alpha-transformed, you can make this NULL.
k	The number of nearest neighbours to consider. It can be a single number or a vector.
apostasi	The type of distance to use, either "euclidean" or "manhattan".
method	If you want to take the average of the responses of the k closest observations, type "average". For the median, type "median" and for the harmonic mean, type "harmonic".

Details

The α -k-NN regression with compositional predictor variables is applied.

Value

A matrix with the estimated response data for each value of k.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Michail Tsagris, Abdulaziz Alenazi and Connie Stewart (2020). The α -k-NN- regression for compositional data. <https://arxiv.org/pdf/2002.05137.pdf>

See Also

[aknn.reg](#), [alfa.knn](#), [alfa.pcr](#), [alfa.ridge](#)

Examples

```
library(MASS)
x <- as.matrix(fg1[, 2:9])
x <- x / rowSums(x)
y <- fg1[, 1]
mod <- alfa.knn.reg(x, y, x, a = 0.5, k = 2:4)
```

The alpha-transformation
The α -transformation

Description

The α -transformation.

Usage

```
alfa(x, a, h = TRUE)
alef(x, a)
```

Arguments

x	A matrix with the compositional data.
a	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied.
h	A boolean variable. If is TRUE (default value) the multiplication with the Helmert sub-matrix will take place. When $\alpha = 0$ and h = FALSE, the result is the centred log-ratio transformation (Aitchison, 1986). In general, when h = FALSE the resulting transformation maps the data onto a singular space. The sum of the vectors is equal to 0. Hence, from the simplex constraint the data go to another constraint.

Details

The α -transformation is applied to the compositional data. The command "alef" is the same as "alfa(x, a, h = FALSE)", but returns a different element as well and is necessary for the functions [a.est](#), [a.mle](#) and [alpha.mle](#).

Value

A list including:

sa	The logarithm of the Jacobian determinant of the α -transformation. This is used in the "profile" function to speed up the computations.
sk	If the "alef" was called, this will return the sum of the α -power transformed data, prior to being normalised to sum to 1. If $\alpha = 0$, this will not be returned.
aff	The α -transformed data.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

Tsagris Michail and Stewart Connie, (2020). A folded model for compositional data analysis. Australian and New Zealand Journal of Statistics (to appear). <https://arxiv.org/pdf/1802.07330.pdf>

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[alfainv](#), [alfa.profile](#), [alfa.tune a.est](#), [alpha.mle](#)

Examples

```
library(MASS)
x <- as.matrix(fgl[, 2:9])
x <- x / rowSums(x)
y1 <- alfa(x, 0.2)$aff
y2 <- alfa(x, 1)$aff
rbind( colMeans(y1), colMeans(y2) )
y3 <- alfa(x, 0.2)$aff
dim(y1) ; dim(y3)
rowSums(y1)
rowSums(y3)
```

The ESOV-distance

The ESOV-distance

Description

The ESOV-distance.

Usage

```
esov(x)
esova(xnew, x)
es(x1, x2)
```

Arguments

x	A matrix with compositional data.
xnew	A matrix or a vector with new compositional data.
x1	A vector with compositional data.
x2	A vector with compositional data.

Details

The ESOV distance is calculated.

Value

For "esov" a matrix including the pairwise distances of all observations or the distances between x_{new} and x . For "esova" a matrix including the pairwise distances of all observations or the distances between x_{new} and x . For "es" a number, the ESOV distance between x_1 and x_2 .

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Tsagris, Michail (2014). The k-NN algorithm for compositional data: a revised approach with and without zero values present. *Journal of Data Science*, 12(3): 519-534.

Endres, D. M. and Schindelin, J. E. (2003). A new metric for probability distributions. *Information Theory, IEEE Transactions on* 49, 1858-1860.

Osterreicher, F. and Vajda, I. (2003). A new class of metric divergences on probability spaces and its applicability in statistics. *Annals of the Institute of Statistical Mathematics* 55, 639-653.

See Also

[alfadist](#), [comp.knn](#), [js.compreg](#)

Examples

```
library(MASS)
x <- as.matrix(fgl[1:20, 2:9])
x <- x / rowSums(x)
esov(x)
```

The Frechet mean for compositional data

The Frechet mean for compositional data

Description

Mean vector or matrix with mean vectors of compositional data using the α -transformation.

Usage

```
frechet(x, a)
```

Arguments

x	A matrix with the compositional data.
a	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied and the closed geometric mean is calculated. You can also provide a sequence of values of alpha and in this case a matrix of Frechet means will be returned.

Details

The power transformation is applied to the compositional data and the mean vector is calculated. Then the inverse of it is calculated and the inverse of the power transformation applied to the last vector is the Frechet mean.

Value

If α is a single value, the function will return a vector with the Frechet mean for the given value of α . Otherwise the function will return a matrix with the Frechet means for each value of α .

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[alfa,alfainv,profile](#)

Examples

```
library(MASS)
x <- as.matrix(fg1[, 2:9])
x <- x / rowSums(x)
frechet(x, 0.2)
frechet(x, 1)
```

The Helmert sub-matrix

The Helmert sub-matrix

Description

The Helmert sub-matrix.

Usage

```
helm(n)
```

Arguments

n A number greater than or equal to 2.

Details

The Helmert sub-matrix is returned. It is an orthogonal matrix without the first row.

Value

A $(n - 1) \times n$ matrix.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>.

References

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

John Aitchison (2003). The Statistical Analysis of Compositional Data p. 99 Blackburn Press.

Lancaster H. O. (1965). The Helmert matrices. The American Mathematical Monthly 72(1): 4-12.

See Also

[alfa](#), [alfainv](#)

Examples

```
helm(3)
helm(5)
```

The k -nearest neighbours using the α -distance

The k -nearest neighbours using the α -distance

Description

The k -nearest neighbours using the α -distance.

Usage

```
alfann(xnew, x, a, k = 10, rann = FALSE)
```

Arguments

xnew	A matrix or a vector with new compositional data.
x	A matrix with the compositional data.
a	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$, the isometric log-ratio transformation is applied.
k	The number of nearest neighbours to search for.
rann	If you have large scale datasets and want a faster k -NN search, you can use kd-trees implemented in the R package "RANN". In this case you must set this argument equal to TRUE. Note however, that in this case, the only available distance is by default "euclidean".

Details

The α -transformation is applied to the compositional data first and the indices of the k -nearest neighbours using the Euclidean distance are returned.

Value

A matrix including the indices of the nearest neighbours of each xnew from x.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Michail Tsagris, Abdulaziz Alenazi and Connie Stewart (2020). The α - k -NN- regression for compositional data. <https://arxiv.org/pdf/2002.05137.pdf>

See Also

[alfa.knn](#), [aknn.reg](#), [alfa](#), [alfainv](#)

Examples

```

library(MASS)
xnew <- as.matrix(fgl[1:20, 2:9])
xnew <- xnew / rowSums(xnew)
x <- as.matrix(fgl[-c(1:20), 2:9])
x <- x / rowSums(x)
b <- alfann(xnew, x, a = 0.1, k = 10)

```

The k-NN algorithm for compositional data

The k-NN algorithm for compositional data

Description

The k-NN algorithm for compositional data with and without using the power transformation.

Usage

```

comp.knn(xnew, x, ina, a = 1, k = 5, type = "S", apostasi = "ESOV", mesos = TRUE)

alfa.knn(xnew, x, ina, a = 1, k = 5, type = "S", mesos = TRUE,
apostasi = "euclidean", rann = FALSE)

```

Arguments

xnew	A matrix with the new compositional data whose group is to be predicted. Zeros are allowed, but you must be careful to choose strictly positive values of α or not to set <code>apostasi = "Ait"</code> .
x	A matrix with the available compositional data. Zeros are allowed, but you must be careful to choose strictly positive values of α or not to set <code>apostasi = "Ait"</code> .
ina	A group indicator variable for the available data.
a	The value of α . As zero values in the compositional data are allowed, you must be careful to choose strictly positive values of α . You have the option to put <code>a = NULL</code> . In this case, the <code>xnew</code> and <code>x</code> are assumed to be the already α -transformed data.
k	The number of nearest neighbours to consider. It can be a single number or a vector.
type	This can be either "S" for the standard k-NN or "NS" for the non standard (see details).
apostasi	The type of distance to use. For the <code>comp.knn</code> this can be one of the following: "ESOV", "taxicab", "Ait", "Hellinger", "angular" or "CS". See the references for them. For the <code>alfa.knn</code> this can be either "euclidean" or "manhattan".
mesos	This is used in the non standard algorithm. If TRUE, the arithmetic mean of the distances is calculated, otherwise the harmonic mean is used (see details).

`rann` If you have large scale datasets and want a faster k-NN search, you can use kd-trees implemented in the R package "RANN". In this case you must set this argument equal to TRUE. Note however, that in this case, the only available distance is by default "euclidean".

Details

The k-NN algorithm is applied for the compositional data. There are many metrics and possibilities to choose from. The standard algorithm finds the k nearest observations to a new observation and allocates it to the class which appears most times in the neighbours. The non standard algorithm is slower but perhaps more accurate. For every group it finds the k nearest neighbours to the new observation. It then computes the arithmetic or the harmonic mean of the distances. The new point is allocated to the class with the minimum distance.

Value

A vector with the estimated groups.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

- Tsagris, Michail (2014). The k-NN algorithm for compositional data: a revised approach with and without zero values present. *Journal of Data Science*, 12(3): 519-534.
- Friedman Jerome, Trevor Hastie and Robert Tibshirani (2009). *The elements of statistical learning*, 2nd edition. Springer, Berlin
- Tsagris Michail, Simon Preston and Andrew T.A. Wood (2016). Improved classification for compositional data using the α -transformation. *Journal of classification* 33(2): 243-261.
- Connie Stewart (2017). An approach to measure distance between compositional diet estimates containing essential zeros. *Journal of Applied Statistics* 44(7): 1137-1152.
- Endres, D. M. and Schindelin, J. E. (2003). A new metric for probability distributions. *Information Theory, IEEE Transactions on* 49, 1858-1860.
- Osterreicher, F. and Vajda, I. (2003). A new class of metric divergences on probability spaces and its applicability in statistics. *Annals of the Institute of Statistical Mathematics* 55, 639-653.

See Also

[compknn](#), [tune](#), [rda](#), [alfa](#), [esov](#)

Examples

```
x <- as.matrix( iris[, 1:4] )
x <- x/ rowSums(x)
ina <- iris[, 5]
```

```
mod <- comp.knn(x, x, ina, a = 1, k = 5)
table(ina, mod)
mod2 <- alfa.knn(x, x, ina, a = 1, k = 5)
table(ina, mod2)
```

Total variability *Total variability*

Description

Total variability.

Usage

```
totvar(x, a = 0)
```

Arguments

x A numerical matrix with the compositional data.

a The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the centred log-ratio transformation is used.

Details

The α -transformation is applied and the sum of the variances of the transformed variables is calculated. This is the total variability. Aitchison (1986) used the centred log-ratio transformation, but we have extended it to cover more geometries, via the α -transformation.

Value

The total variability of the data in a given geometry as dictated by the value of α .

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

Aitchison J. (1986). The statistical analysis of compositional data. Chapman & Hall.

See Also

[alfa](#), [link{alfainv}](#), [alfa.profile](#), [alfa.tune](#)

Examples

```
x <- as.matrix(iris[, 1:4])
x <- x / rowSums(x)
totvar(x)
```

Transformation-free linear regression for compositional responses and predictors
*Transformation-free linear regression for compositional responses
and predictors*

Description

Transformation-free linear regression for compositional responses and predictors.

Usage

```
tflr(y, x, xnew = NULL)
```

Arguments

y	A matrix with the compositional response. Zero values are allowed.
x	A matrix with the compositional predictors. Zero values are allowed.
xnew	If you have new data use it, otherwise leave it NULL.

Details

The transformation-free linear regression for compositional responses and predictors is implemented.

Value

A list including:

runtime	The time required by the regression.
loglik	The log-likelihood $-\sum_{i=1}^n y_i \log y_i / \hat{y}_i$.
be	The beta coefficients.
est	The fitted values of xnew if xnew is not NULL.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Jacob Fiksel, Scott Zeger and Abhirup Datta (2020). A transformation-free linear regression for compositional outcomes and predictors. <https://arxiv.org/pdf/2004.07881.pdf>

See Also

[cv.tflr](#), [kl.alfapcr](#)

Examples

```
library(MASS)
y <- rdirich(214, runif(3, 1, 3))
x <- as.matrix(fgl[, 2:9])
x <- x / rowSums(x)
mod <- tflr(y, x, x)
mod
```

Tuning of the bandwidth h of the kernel using the maximum likelihood cross validation
Tuning of the bandwidth h of the kernel using the maximum likelihood cross validation

Description

Tuning of the bandwidth h of the kernel using the maximum likelihood cross validation.

Usage

```
mkde.tune( x, low = 0.1, up = 3, s = cov(x) )
```

Arguments

<code>x</code>	A matrix with Euclidean (continuous) data.
<code>low</code>	The minimum value to search for the optimal bandwidth value.
<code>up</code>	The maximum value to search for the optimal bandwidth value.
<code>s</code>	A covariance matrix. By default it is equal to the covariance matrix of the data, but can change to a robust covariance matrix, MCD for example.

Details

Maximum likelihood cross validation is applied in order to choose the optimal value of the bandwidth parameter. No plot is produced.

Value

A list including:

<code>hopt</code>	The optimal bandwidth value.
<code>maximum</code>	The value of the pseudo-log-likelihood at that given bandwidth value.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Arsalane Chouaib Guidoum (2015). Kernel Estimator and Bandwidth Selection for Density and its Derivatives. The kedd R package. <http://cran.r-project.org/web/packages/kedd/vignettes/kedd.pdf>

M.P. Wand and M.C. Jones (1995). Kernel smoothing, pages 91-92.

See Also

[mkde](#), [comp.kerncontour](#)

Examples

```
library(MASS)
mkde.tune(as.matrix(iris[, 1:4]), c(0.1, 3) )
```

Tuning of the divergence based regression for compositional data with compositional data in the covariates side using the alpha-transformation

Tuning of the divergence based regression for compositional data with compositional data in the covariates side using the alpha-transformation

Description

Tuning of the divergence based regression for compositional data with compositional data in the covariates side using the α -transformation.

Usage

```
klalfapcr.tune(y, x, covar = NULL, nfolds = 10, maxk = 50, a = seq(-1, 1, by = 0.1),
folds = NULL, graph = FALSE, tol = 1e-07, maxiters = 50, seed = FALSE)
```

Arguments

y	A numerical matrix with compositional data with or without zeros.
x	A matrix with the predictor variables, the compositional data. Zero values are allowed.
covar	If you have other continuous covariates put them here.
nfolds	The number of folds for the K-fold cross validation, set to 10 by default.
maxk	The maximum number of principal components to check.

Tuning of the divergence based regression for compositional data with compositional data in the covariates side using the alpha-tra

a	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied.
folds	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
graph	If graph is TRUE (default value) a filled contour plot will appear.
tol	The tolerance value to terminate the Newton-Raphson procedure.
maxiters	The maximum number of Newton-Raphson iterations.
seed	If seed is TRUE the results will always be the same.

Details

The M-fold cross validation is performed in order to select the optimal values for α and k, the number of principal components. The α -transformation is applied to the compositional data first, the first k principal component scores are calculated and used as predictor variables for the Kullback-Leibler divergence based regression model. This procedure is performed M times during the M-fold cross validation.

Value

A list including:

mspe	A list with the KL divergence for each value of α and k in every fold.
performance	A matrix with the KL divergence for each value of α averaged over all folds. If graph is set to TRUE this matrix is plotted.
best.perf	The minimum KL divergence.
params	The values of α and k corresponding to the minimum KL divergence.

Author(s)

Initial code by Abdulaziz Alenazi. Modifications by Michail Tsagris.

R implementation and documentation: Abdulaziz Alenazi <a.alenazi@nbu.edu.sa> Michail Tsagris <mtsagris@uoc.gr>

References

- Alenazi A. (2019). Regression for compositional data with compositionl data as predictor variables with or without zero values. *Journal of Data Science*, 17(1): 219-238. http://www.jds-online.com/file_download/688/01+No.10+315+REGRESSION+FOR+COMPOSITIONAL+DATA+WITH+COMPOSITIO
- Tsagris M. (2015). Regression analysis with compositional data containing zero values. *Chilean Journal of Statistics*, 6(2): 47-57. <http://arxiv.org/pdf/1508.01913v1.pdf>
- Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In *Proceedings of the 4th Compositional Data Analysis Workshop*, Girona, Spain. <http://arxiv.org/pdf/1106.1451.pdf>

See Also

[kl.alfapcr](#), [cv.tflr](#), [pcr](#), [glm.pcr](#), [alfapcr.tune](#)

Examples

```
library(MASS)
y <- rdirich( 214, runif(4, 1, 3) )
x <- as.matrix( fgl[, 2:9] )
x <- x / rowSums(x)
mod <- klalfapcr.tune(y = y, x = x, a = c(0.7, 0.8) )
mod
```

Tuning of the k-NN algorithm for compositional data

Tuning of the k-NN algorithm for compositional data

Description

Tuning of the k-NN algorithm for compositional data with and without using the power or the α -transformation. In addition, estimation of the rate of correct classification via M-fold cross-validation.

Usage

```
compknn.tune(x, ina, nfolds = 10, k = 2:5, type = "S", mesos = TRUE,
a = seq(-1, 1, by = 0.1), apostasi = "ESOV", folds = NULL,
stratified = FALSE, seed = FALSE, graph = FALSE)
```

```
alfaknn.tune(x, ina, nfolds = 10, k = 2:5, type = "S", mesos = TRUE,
a = seq(-1, 1, by = 0.1), apostasi = "euclidean", rann = FALSE, folds = NULL,
stratified = FALSE, seed = FALSE, graph = FALSE)
```

Arguments

x	A matrix with the available compositional data. Zeros are allowed, but you must be careful to choose strictly positive values of α or not to set <code>apostasi = "Ait"</code> .
ina	A group indicator variable for the available data.
nfolds	The number of folds to be used. This is taken into consideration only if the <code>folds</code> argument is not supplied.
k	A vector with the nearest neighbours to consider.
type	This can be either "S" for the standard k-NN or "NS" for the non standard (see details).
mesos	This is used in the non standard algorithm. If TRUE, the arithmetic mean of the distances is calculated, otherwise the harmonic mean is used (see details).
a	A grid of values of α to be used only if the distance chosen allows for it.

<code>apostasi</code>	The type of distance to use. For the <code>compk.knn</code> this can be one of the following: "ESOV", "taxicab", "Ait", "Hellinger", "angular" or "CS". See the references for them. For the <code>alfa.knn</code> this can be either "euclidean" or "manhattan".
<code>rann</code>	If you have large scale datasets and want a faster k-NN search, you can use kd-trees implemented in the R package "RANN". In this case you must set this argument equal to TRUE. Note however, that in this case, the only available distance is by default "euclidean".
<code>folds</code>	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
<code>stratified</code>	Do you want the folds to be created in a stratified way? TRUE or FALSE.
<code>seed</code>	If seed is TRUE the results will always be the same.
<code>graph</code>	If set to TRUE a graph with the results will appear.

Details

The k-NN algorithm is applied for the compositional data. There are many metrics and possibilities to choose from. The standard algorithm finds the k nearest observations to a new observation and allocates it to the class which appears most times in the neighbours. The non standard algorithm is slower but perhaps more accurate. For every group it finds the k nearest neighbours to the new observation. It then computes the arithmetic or the harmonic mean of the distances. The new point is allocated to the class with the minimum distance.

Value

A list including:

<code>ela</code>	A matrix or a vector (depending on the distance chosen) with the averaged over all folds rates of correct classification for all hyper-parameters (α and k).
<code>performance</code>	The estimated rate of correct classification.
<code>best_a</code>	The best value of α . This is returned for "ESOV" and "taxicab" only.
<code>best_k</code>	The best number of nearest neighbours.
<code>runtime</code>	The run time of the cross-validation procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

- Tsagris, Michail (2014). The k-NN algorithm for compositional data: a revised approach with and without zero values present. *Journal of Data Science*, 12(3): 519-534. <https://arxiv.org/pdf/1506.05216.pdf>
- Friedman Jerome, Trevor Hastie and Robert Tibshirani (2009). *The elements of statistical learning*, 2nd edition. Springer, Berlin

Tsagris Michail, Simon Preston and Andrew T.A. Wood (2016). Improved classification for compositional data using the α -transformation. *Journal of classification* 33(2): 243-261. <https://arxiv.org/pdf/1506.04976v2.pdf>

Connie Stewart (2017). An approach to measure distance between compositional diet estimates containing essential zeros. *Journal of Applied Statistics* 44(7): 1137-1152.

Endres, D. M. and Schindelin, J. E. (2003). A new metric for probability distributions. *Information Theory, IEEE Transactions on* 49, 1858-1860.

Osterreicher, F. and Vajda, I. (2003). A new class of metric divergences on probability spaces and its applicability in statistics. *Annals of the Institute of Statistical Mathematics* 55, 639-653.

See Also

[comp.knn](#), [rda](#), [alfa](#)

Examples

```
x <- as.matrix(iris[, 1:4])
x <- x/ rowSums(x)
ina <- iris[, 5]
mod1 <- compknn.tune(x, ina, a = seq(1, 1, by = 0.1) )
mod2 <- alfa.knn.tune(x, ina, a = seq(-1, 1, by = 0.1) )
```

Tuning of the projection pursuit regression for compositional data

Tuning of the projection pursuit regression for compositional data

Description

Tuning of the projection pursuit regression for compositional data In addition, estimation of the rate of correct classification via K-fold cross-validation.

Usage

```
compppr.tune(y, x, nfolds = 10, folds = NULL, seed = FALSE, nterms = 1:10,
type = "alr", yb = NULL, B = 1000 )
```

Arguments

y	A matrix with the available compositional data, but zeros are not allowed.
x	A matrix with the continuous predictor variables.
nfolds	The number of folds to use.
folds	If you have the list with the folds supply it here.
seed	If seed is TRUE the results will always be the same.
nterms	The number of terms to try in the projection pursuit regression.

type	Either "alr" or "ilr" corresponding to the additive or the isometric log-ratio transformation respectively.
yb	If you have already transformed the data using a log-ratio transformation put it here. Otherwise leave it NULL.
B	The number of bootstrap re-samples to use for the unbiased estimation of the performance of the projection pursuit regression. If B = 1, no bootstrap is applied.

Details

The function performs tuning of the projection pursuit regression algorithm.

Value

A list including:

k1	The average Kullback-Leibler divergence.
bc.perf	The bootstrap bias corrected average Kullback-Leibler divergence. If no bootstrap was performed this is equal to the average Kullback-Leibler divergence.
runtime	The run time of the cross-validation procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Friedman, J. H. and Stuetzle, W. (1981). Projection pursuit regression. *Journal of the American Statistical Association*, 76, 817-823. doi: 10.2307/2287576.

Tsamardinos I., Greasidou E. and Borboudakis G. (2018). Bootstrapping the out-of-sample predictions for efficient and accurate cross-validation. *Machine Learning* 107(12): 1895-1922. <https://link.springer.com/article/10.1007/s11648-018-5714-4>

See Also

[comp.ppr](#), [comp.reg](#), [alfa](#)

Examples

```
x <- as.matrix(iris[, 1:4])
x <- x/ rowSums(x)
ina <- iris[, 5]
mod1 <- compknn.tune(x, ina, a = seq(1, 1, by = 0.1) )
mod2 <- alfaknn.tune(x, ina, a = seq(-1, 1, by = 0.1) )
```

Tuning the number of PCs in the PCR with compositional data using the alpha-transformation
*Tuning the number of PCs in the PCR with compositional data using
the α -transformation*

Description

This is a cross-validation procedure to decide on the number of principal components when using regression with compositional data (as predictor variables) using the α -transformation.

Usage

```
alfapcr.tune(y, x, model = "gaussian", nfolds = 10, maxk = 50, a = seq(-1, 1, by = 0.1),
folds = NULL, ncores = 1, graph = TRUE, col.nu = 15, seed = FALSE)
```

Arguments

y	A vector with either continuous, binary or count data.
x	A matrix with the predictor variables, the compositional data. Zero values are allowed.
model	The type of regression model to fit. The possible values are "gaussian", "binomial" and "poisson".
nfolds	The number of folds for the K-fold cross validation, set to 10 by default.
maxk	The maximum number of principal components to check.
a	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied and the solution exists in a closed form, since it the classical multivariate regression. The estimated bias correction via the (Tibshirani and Tibshirani (2009) criterion is applied.
folds	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
ncores	How many cores to use. If you have heavy computations or do not want to wait for long time more than 1 core (if available) is suggested. It is advisable to use it if you have many observations and or many variables, otherwise it will slow down the process.
graph	If graph is TRUE (default value) a filled contour plot will appear.
col.nu	A number parameter for the filled contour plot, taken into account only if graph is TRUE.
seed	If seed is TRUE the results will always be the same.

Details

The α -transformation is applied to the compositional data first and the function "pcr.tune" or "glm-pcr.tune" is called.

Value

If graph is TRUE a field contour a filled contour will appear. A list including:

mspe	The MSPE where rows correspond to the α values and the columns to the number of principal components.
best.par	The best pair of α and number of principal components.
performance	The minimum mean squared error of prediction.
runtime	The time required by the cross-validation procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Tsagris M. (2015). Regression analysis with compositional data containing zero values. *Chilean Journal of Statistics*, 6(2): 47-57. <https://arxiv.org/pdf/1508.01913v1.pdf>

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In *Proceedings of the 4th Compositional Data Analysis Workshop*, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

Jolliffe I.T. (2002). *Principal Component Analysis*.

See Also

[alfa](#), [profile](#), [alfa.pcr](#), [pcr.tune](#), [glmPCR.tune](#), [glm](#)

Examples

```
library(MASS)
y <- as.vector(fgl[, 1])
x <- as.matrix(fgl[, 2:9])
x <- x/ rowSums(x)
mod <- alfapcr.tune(y, x, nfolds = 10, maxk = 50, a = seq(-1, 1, by = 0.1) )
```

Tuning the parameters of the regularised discriminant analysis

Tuning the parameters of the regularised discriminant analysis

Description

Tuning the parameters of the regularised discriminant analysis for Euclidean data.

Usage

```
rda.tune(x, ina, nfolds = 10, gam = seq(0, 1, by = 0.1), del = seq(0, 1, by = 0.1),
ncores = 1, folds = NULL, stratified = TRUE, seed = FALSE)
```

Arguments

x	A matrix with the data.
ina	A group indicator variable for the available data.
nfolds	The number of folds in the cross validation.
gam	A grid of values for the γ parameter as defined in Tsagris et al. (2016).
del	A grid of values for the δ parameter as defined in Tsagris et al. (2016).
ncores	The number of cores to use. If more than 1, parallel computing will take place. It is advisable to use it if you have many observations and/or many variables, otherwise it will slow down the process.
folds	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
stratified	Do you want the folds to be created in a stratified way? TRUE or FALSE.
seed	If seed is TRUE the results will always be the same.

Details

Cross validation is performed to select the optimal parameters for the regularised discriminant analysis and also estimate the rate of accuracy.

The covariance matrix of each group is calculated and then the pooled covariance matrix. The spherical covariance matrix consists of the average of the pooled variances in its diagonal and zeros in the off-diagonal elements. gam is the weight of the pooled covariance matrix and $1 - \text{gam}$ is the weight of the spherical covariance matrix, $S_a = \text{gam} * S_p + (1 - \text{gam}) * s_p$. Then it is a compromise between LDA and QDA. del is the weight of S_a and $1 - \text{del}$ the weight of each group covariance group. This function is a wrapper for [alfa.rda](#).

Value

A list including: If graph is TRUE a plot of the performance versus the number of principal components will appear.

per	An array with the estimate rate of correct classification for every fold. For each of the M matrices, the row values correspond to gam and the columns to the del parameter.
percent	A matrix with the mean estimated rates of correct classification. The row values correspond to gam and the columns to the del parameter.
se	A matrix with the standard error of the mean estimated rates of correct classification. The row values correspond to gam and the columns to the del parameter.
result	The estimated rate of correct classification along with the best gam and del parameters.
runtime	The time required by the cross-validation procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Friedman Jerome, Trevor Hastie and Robert Tibshirani (2009). The elements of statistical learning, 2nd edition. Springer, Berlin

Tsagris Michail, Simon Preston and Andrew TA Wood (2016). Improved classification for compositional data using the α -transformation. Journal of classification, 33(2):243-261. <http://arxiv.org/pdf/1106.1451.pdf>

See Also

[rda, alfa](#)

Examples

```
mod <- rda.tune(as.matrix(iris[, 1:4]), iris[, 5], gam = seq(0, 1, by = 0.2),
del = seq(0, 1, by = 0.2) )
mod
```

Tuning the principal components with GLMs

Tuning the principal components with GLMs

Description

Tuning the number of principal components in the generalised linear models.

Usage

```
pcr.tune(y, x, nfolds = 10, maxk = 50, folds = NULL, ncores = 1,
seed = FALSE, graph = TRUE)
```

```
glmpcr.tune(y, x, nfolds = 10, maxk = 10, folds = NULL, ncores = 1,
seed = FALSE, graph = TRUE)
```

```
multinompcr.tune(y, x, nfolds = 10, maxk = 10, folds = NULL, ncores = 1,
seed = FALSE, graph = TRUE)
```

Arguments

y	A real valued vector for "pcr.tune". A real valued vector for the "glmpr.tune" with either two numbers, 0 and 1 for example, for the binomial regression or with positive discrete numbers for the poisson. For the "multinompcr.tune" a vector or a factor with more than just two values. This is a multinomial regression.
x	A matrix with the predictor variables, they have to be continuous.
nfolds	The number of folds in the cross validation.
maxk	The maximum number of principal components to check.
folds	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
ncores	The number of cores to use. If more than 1, parallel computing will take place. It is advisable to use it if you have many observations and or many variables, otherwise it will slow down th process.
seed	If seed is TRUE the results will always be the same.
graph	If graph is TRUE a plot of the performance for each fold along the values of α will appear.

Details

Cross validation is performed to select the optimal number of principal components in the GLMs or the multinomial regression. This is used by [alfapcr.tune](#).

Value

If graph is TRUE a plot of the performance versus the number of principal components will appear. A list including:

msp	A matrix with the mean deviance of prediction or mean accuracy for every fold.
mpd	A vector with the mean deviance of prediction or mean accuracy, each value corresponds to a number of principal components.
k	The number of principal components which minimizes the deviance or maximises the accuracy.
performance	The optimal performance, MSE for the linea regression, minimum deviance for the GLMs and maximum accuracy for the multinomial regression.
runtime	The time required by the cross-validation procedure.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>.

References

Aguilera A.M., Escabias M. and Valderrama M.J. (2006). Using principal components for estimating logistic regression with high-dimensional multicollinear data. *Computational Statistics & Data Analysis* 50(8): 1905-1924.

Jolliffe I.T. (2002). *Principal Component Analysis*.

See Also

[pcr.tune](#), [glm.pcr](#), [alfa.pcr](#), [alfapcr.tune](#)

Examples

```
library(MASS)
x <- as.matrix(fg1[, 2:9])
y <- rpois(214, 10)
glmPCR.tune(y, x, nfolds = 10, maxk = 20, folds = NULL, ncores = 1)
```

Tuning the value of alpha in the alpha-regression

Tuning the value of α in the α -regression

Description

Tuning the value of α in the α -regression.

Usage

```
alfareg.tune(y, x, a = seq(0.1, 1, by = 0.1), nfolds = 10,
folds = NULL, nc = 1, seed = FALSE, graph = FALSE)
```

Arguments

y	A matrix with compositional data. zero values are allowed.
x	A matrix with the continuous predictor variables or a data frame including categorical predictor variables.
a	The value of the power transformation, it has to be between -1 and 1. If zero values are present it has to be greater than 0. If $\alpha = 0$ the isometric log-ratio transformation is applied.
nfolds	The number of folds to split the data.
folds	If you have the list with the folds supply it here. You can also leave it NULL and it will create folds.
nc	The number of cores to use. IF you have a multicore computer it is advisable to use more than 1. It makes the procedure faster. It is advisable to use it if you have many observations and or many variables, otherwise it will slow down th process.

seed	If seed is TRUE the results will always be the same.
graph	If graph is TRUE a plot of the performance for each fold along the values of α will appear.

Details

The α -transformation is applied to the compositional data and the numerical optimisation is performed for the regression, unless $\alpha = 0$, where the coefficients are available in closed form.

Value

A plot of the estimated Kullback-Leibler divergences (multiplied by 2) along the values of α (if graph is set to TRUE). A list including:

runtime	The runtime required by the cross-validation.
kula	A matrix with twice the Kullback-Leibler divergence of the observed from the fitted values. Each row corresponds to a fold and each column to a value of α . The average over the columns equal the next argument, "kl".
kl	A vector with twice the Kullback-Leibler divergence of the observed from the fitted values. Every value corresponds to a value of α .
opt	The optimal value of α .
value	The minimum value of twice the Kullback-Leibler.

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr> and Giorgos Athineou <gioathineou@gmail.com>

References

Tsagris Michail (2015). Regression analysis with compositional data containing zero values. Chilean Journal of Statistics, 6(2): 47-57. <https://arxiv.org/pdf/1508.01913v1.pdf>

Tsagris M.T., Preston S. and Wood A.T.A. (2011). A data-based power transformation for compositional data. In Proceedings of the 4th Compositional Data Analysis Workshop, Girona, Spain. <https://arxiv.org/pdf/1106.1451.pdf>

See Also

[alfa.reg](#), [alfa](#)

Examples

```
library(MASS)
y <- as.matrix(fgl[1:40, 2:4])
y <- y / rowSums(y)
x <- as.vector(fgl[1:40, 1])
mod <- alfareg.tune(y, x, a = c(0.2, 0.35, 0.05), nfold = 5)
```

Zero adjusted Dirichlet regression
Zero adjusted Dirichlet regression

Description

Zero adjusted Dirichlet regression.

Usage

```
zadr(y, x, xnew = NULL, tol = 1e-05)
```

```
mixreg(param, z)
```

Arguments

y	A matrix with the compositional data (dependent variable). The number of observations with at least one zero value should not be more than the columns of the predictor variables. Otherwise, the initial values will not be calculated.
x	The predictor variable(s), they can be either continuous or categorical or both.
xnew	If you have new data use it, otherwise leave it NULL.
tol	A tolerance level to terminate the maximisation process.
param	Some arguments passed on to the mixreg helper function.
z	Some arguments passed on to the mixreg helper function.

Details

A zero adjusted Dirichlet regression is being fitted. The likelihood consists of two components. The contributions of the non zero compositional values and the contributions of the compositional vectors with at least one zero value. The second component may have many different sub-categories, one for each pattern of zeros. The function "mixreg" is a helper function and is not intended to be called directly by the user.

Value

A list including:

runtime	The time required by the regression.
loglik	The value of the log-likelihood.
phi	The precision parameter. If covariates are linked with it (function "diri.reg2"), this will be a vector.
be	The beta coefficients.
seb	The standard error of the beta coefficients.
sigma	The covariance matrix of the regression parameters (for the mean vector and the phi parameter) in the function "diri.reg2".
est	The fitted or the predicted values (if xnew is not NULL).

Author(s)

Michail Tsagris

R implementation and documentation: Michail Tsagris <mtsagris@uoc.gr>

References

Tsagris M. and Stewart C. (2018). A Dirichlet Regression Model for Compositional Data with Zeros. Accepted at the Lobachevskii Journal of Mathematics.

Preprint available from <https://arxiv.org/pdf/1410.5011.pdf>

See Also

[diri.reg](#), [kl.compreg](#), [ols.compreg](#), [alfa.reg](#)

Examples

```
x <- as.vector(iris[, 4])
y <- as.matrix(iris[, 1:3])
y <- y / rowSums(y)
mod1 <- diri.reg(y, x)
y[sample(1:450, 15) ] <- 0
mod2 <- zadr(y, x)
```

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