

Package ‘ER’

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Author Kristian Hovde Liland [aut, cre],
Ellen Færgestad Mosleth [ctb]

Maintainer Kristian Hovde Liland <kristian.liland@nmbu.no>

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 confints

Confidence Intervals of Effect Differences

Description

Confidence Intervals of Effect Differences

Usage

```
confints(X1, X2, confidence = 0.95, df.used = 0)
```

```
## S3 method for class 'confints'
plot(
  x,
  y,
  xlab = "",
  ylab = "normalised log2",
  sorted = TRUE,
  labels = FALSE,
  nonZero = FALSE,
  xlim = NULL,
  ylim = NULL,
  text.pt = 12,
  ...
)
```

Arguments

X1	data.frame containing first effect.
X2	data.frame containing second effect.
confidence	Level of confidence, default = 0.95.
df.used	Optional argument indicating how many degrees of freedom have been consumed during deflation. Default = 0.
x	Object of class confint.
y	Not used.
xlab	X label (character)
ylab	Y label (character)
sorted	Logical indicating if intervals should be sorted according to their mean values, or a vector of indices/labels to sort by.
labels	Logical indicating if sample labels should be used on x axis.
nonZero	Logical indicating if intervals are required not to include zero.
xlim	Limits of the horizontal scale.
ylim	Limits of the vertical scale.
text.pt	Size scaling of text in the plot (default = 16).
...	Further arguments to qplot.

Value

An object of class `confints`, which holds the information needed to perform statistics or plot the confidence intervals is returned from `confints`. The plotting routine returns a `ggplot` structure for plotting.

See Also

[ER](#), [elastic](#) and [pls](#).

Examples

```
data(MS)
# Compare MS and non-MS patients within cluster 1
conf <- with(MS, confints(proteins[MS == "yes" & cluster == 1,],
                        proteins[MS == "no" & cluster == 1,]))

p1 <- plot(conf)
p2 <- plot(conf, nonZero = TRUE) # Only intervals without 0.
grid.arrange(p1,p2)

# Shorter plot with labels
confShort <- conf[1:10,]
p1 <- plot(confShort, labels = TRUE)
p2 <- plot(confShort, labels = TRUE, nonZero = TRUE)
grid.arrange(p1,p2)
```

Diabetes

Diabetes data

Description

A data.frame with a design and transcriptomic data.

Usage

```
data(Diabetes)
```

Details

Clinical study on humans was performed as a 2-way factorial design with two factors both on two levels: bariatric surgery on two levels (before and after the bariatric surgery) and type 2 diabetes (T2D) on two levels (with and without T2D). There were 8 patients without T2D and 7 with T2D. It was discovered that the patients with T2D would be separated in two groups: 3 patients in the group called T2D1 and 4 patients in the group called T2D2. The experiment can therefore also analysed as 2 way factorial design where the disease factor is on three levels. All patients were obese before bariatric surgery (BMI >45). Transcriptome in the subcutaneous adipose tissue were obtained before and one year after bariatric surgery.

Author(s)

Ellen Færgestad Mosleth

References

Dankel et al. 2010. Switch from Stress Response to Homeobox Transcription Factors in Adipose Tissue After Profound Fat Loss. Plos One 5.

Examples

```
data(Diabetes)
str(Diabetes)
```

elastic

Elastic-net modeling of ER objects.

Description

Elastic-net modeling of ER objects.

Usage

```
elastic(er, ...)

## S3 method for class 'ER'
elastic(
  er,
  effect,
  alpha = 0.5,
  newdata = NULL,
  validation,
  segments = NULL,
  measure = measure,
  family = family,
  ...
)
```

Arguments

er	Object of class ER.
...	Additional arguments for <code>pls::cvsegments</code> .
effect	The effect to be used as response.
alpha	The elasticnet mixing parameter.
newdata	Optional new data matrix for prediction.
validation	Optional validation parameters.


```

  what = "raw",
  col = NULL,
  pch = NULL,
  model.line = (what %in% c("raw")),
  ylim = NULL,
  ylab = "",
  xlab = "",
  main = NULL,
  ...
)

```

```
tableER(object, variable)
```

Arguments

formula	a model formula specifying features and effects.
data	a data.frame containing response variables (features) and design factors or other groupings/continuous variables.
x	Object of class ER.
...	Additional arguments to plot
y	Response name or number.
what	What part of ER to plot; raw data (default), fits, residuals or a named model effect (can be combined with 'effect', see Examples).
col	Color of points, defaults to grouping. Usually set to a factor name.
pch	Plot character of points, defaults to 1. Usually set to a factor name.
model.line	Include line indicating estimates, default = TRUE. Can be an effect name.
ylim	Y axis limits (numeric, but defaults to NULL)
ylab	Y label (character)
xlab	X label (character)
main	Main title, defaults to y with description from what.
object	ER object.
variable	Numeric for selecting a variable for extraction.

Value

ER returns an object of class ER containing effects, ER values, fitted values, residuals, features, coefficients, dummy design, symbolic design, dimensions, highest level interaction and feature names.

References

- * Mosleth et al. (2021) Cerebrospinal fluid proteome shows disrupted neuronal development in multiple sclerosis. Scientific Report, 11,4087. <doi:10.1038/s41598-021-82388-w>
- * E.F. Mosleth et al. (2020). Comprehensive Chemometrics, 2nd edition; Brown, S., Tauler, R., & Walczak, B. (Eds.). Chapter 4.22. Analysis of Megavariate Data in Functional Omics. Elsevier. <doi:10.1016/B978-0-12-409547-2.14882-6>

See Also

Analyses using ER: [elastic](#) and [pls](#). Confidence interval plots [confints](#).

Examples

```
## Multiple Sclerosis
data(MS, package = "ER")
er <- ER(proteins ~ MS * cluster, data = MS)
print(er)
plot(er) # Raw data, first feature
plot(er,2) # Raw data, numbered feature
plot(er,'Q76L83', col='MS', pch='cluster') # Selected colour and plot character
plot(er,'Q76L83', what='effect MS',
      model.line='effect cluster') # Comparison of factors (points and lines)

# Example compound plot
old.par <- par(c("mfrow", "mar"))
# on.exit(par(old.par))
par(mfrow = c(3,3), mar = c(2,4,4,1))
plot(er,'Q76L83') # Raw data, named feature
plot(er,'Q76L83', what='fits') # Fitted values
plot(er,'Q76L83', what='residuals') # Residuals
plot(er,'Q76L83', what='effect MS') # Effect levels
plot(er,'Q76L83', what='effect cluster') # ----||----
plot(er,'Q76L83', what='effect MS:cluster') # ----||----
plot(er,'Q76L83', what='MS') # ER values
plot(er,'Q76L83', what='cluster') # -----||-----
plot(er,'Q76L83', what='MS:cluster') # -----||-----
par(old.par)

# Complete overview of ER
tab <- tableER(er, 1)

# In general there can be more than two, effects, more than two levels, and continuous effects:
# MS$three <- factor(c(rep(1:3,33),1:2))
# er3 <- ER(proteins ~ MS * cluster + three, data = MS)

## Lactobacillus
data(Lactobacillus, package = "ER")
erLac <- ER(proteome ~ strain * growthrate, data = Lactobacillus)
print(erLac)
plot(erLac) # Raw data, first feature
plot(erLac,2) # Raw data, numbered feature
plot(erLac,'P.LSA0316', col='strain',
      pch='growthrate') # Selected colour and plot character
plot(erLac,'P.LSA0316', what='strain',
      model.line='growthrate') # Selected model.line

## Diabetes
```

```
data(Diabetes, package = "ER")
erDia <- ER(transcriptome ~ surgery * T2D, data = Diabetes)
print(erDia)
plot(erDia) # Raw data, first feature
plot(erDia,2) # Raw data, numbered feature
plot(erDia,'ILMN_1720829', col='surgery',
      pch='T2D') # Selected colour and plot character
```

Lactobacillus

Lactobacillus data

Description

A data.frame with a design and proteomic data, transcriptomic data and phenotypic data.

Usage

```
data(Lactobacillus)
```

Details

Experiment on *Lactobacillus sakei* was performed as a 2-way factorial design with two factors both on two levels: strain (*L. sakei* strains LS25 and 23K) (factor A) and growth condition (high and low glucose availability) (factor B) both on two levels, and their interaction term (factor AB). There were three biological replicates within each group. Transcriptome, proteome and end product profile (lactate, formate, acetate and ethanol) were observed.

Author(s)

Ellen Færgestad Mosleth

References

McLeod et al. 2017. Effects of glucose availability in *Lactobacillus sakei*; metabolic change and regulation of the proteome and transcriptome. Plos One 12, e0187542.

Examples

```
data(Lactobacillus)
str(Lactobacillus)
```

MS

Multiple Sclerosis data

Description

A data.frame with a design and proteomic data.

Usage

data(MS)

Details

Data from biobank are analysed a study population of 101 patients, 37 were diagnosed with multiple sclerosis, and 64 without multiple sclerosis. Of the patients without multiple sclerosis, 50 were diagnosed with other neurological disorders and 14 were neurologically healthy patients who had undergone spinal anaesthesia for orthopaedic surgery on the knee or ankle, i.e. neurologically healthy controls. Unless otherwise stated, all the patients without multiple sclerosis were considered as controls for this study. All patients with multiple sclerosis had relapsing remitting multiple sclerosis. The proteome were obtained on cerebrospinal fluid samples from all patients prior medical treatment for multiple sclerosis. It was discovered the patients separated into two clusters, called cluster 1 and cluster 2. This is utilised in the data analysis by considering the data as 2-way factorial design with the two factors: MS and clusters both on two levels.

Author(s)

Ellen Færgestad Mosleth

References

* Opsahl, J.A. et al. Label-free analysis of human cerebrospinal fluid addressing various normalization strategies and revealing protein groups affected by multiple sclerosis. *Proteomics* 16, 1154-1165 (2016).

* Ellen Færgestad Mosleth, Christian Alexander Vedeler, Kristian Hovde Liland, Anette McLeod, Gerd Haga Bringland, Liesbeth Kroondijk, Frode Berven, Artem Lysenko, Christopher J. Rawlings, Karim El-Hajj Eid, Jill Anette Opsahl, Bjørn Tore Gjertsen, Kjell-Morten Myhr and Sonia Gavasso, Cerebrospinal fluid proteome shows disrupted neuronal development in multiple sclerosis. *Scientific Reports – Nature* 11(4087), (2021).

Examples

data(MS)

str(MS)

 pls

Partial Least Squares modelling of ER objects.

Description

The output of ER is used as input to a PLS classification with the selected effect as response. It is possible to compare two models using the `er2` argument. Variable selection is available through Jackknifing (from package `pls`) and Shaving (from package `plsVarSel`).

Usage

```
pls(er, ...)

## S3 method for class 'ER'
pls(
  er,
  effect,
  ncomp,
  newdata = NULL,
  er2,
  validation,
  jackknife = NULL,
  shave = NULL,
  df.used = NULL,
  ...
)
```

Arguments

<code>er</code>	Object of class ER.
<code>...</code>	Additional arguments for <code>pls</code> .
<code>effect</code>	The effect to be used as response.
<code>ncomp</code>	Number of PLS components.
<code>newdata</code>	Optional new data matrix for prediction.
<code>er2</code>	Second object of class ER for comparison.
<code>validation</code>	Optional validation parameters for <code>pls</code> .
<code>jackknife</code>	Optional argument specifying if jackknifing should be applied.
<code>shave</code>	Optional argument indicating if variable shaving should be used. <code>shave</code> should be a list with two elements: the PLS filter method and the proportion to remove. <code>shave = TRUE</code> uses defaults: <code>list("sMC", 0.2)</code> .
<code>df.used</code>	Optional argument indicating how many degrees of freedom have been consumed during deflation. Default value from input object.

Details

If using the shave options, the segment type is given as type instead of segment.type (see examples).

See Also

[ER](#), [elastic](#) and [confints](#).

Examples

```
data(MS, package = "ER")
er <- ER(proteins ~ MS * cluster, data = MS[-1,])

# Simple PLS using interleaved cross-validation
plsMod <- pls(er, 'MS', 6, validation = "CV",
             segment.type = "interleaved", length.seg = 25)
scoreplot(plsMod, labels = "names")

# PLS with shaving of variables (mind different variable for cross-validation type)
plsModS <- pls(er, 'MS', 6, validation = "CV",
              type = "interleaved", length.seg=25, shave = TRUE)
# Error as a function of remaining variables
plot(plsModS$shave)
# Selected variables for minimum error
with(plsModS$shave, colnames(X)[variables[[min.red+1]]])

# Time consuming due to leave-one-out cross-validation
plsModJ <- pls(er, 'MS', 5, validation = "LOO",
              jackknife = TRUE)
colSums(plsModJ$classes == as.numeric(MS$MS[-1]))
# Jackknifed coefficient P-values (sorted)
plot(sort(plsModJ$jack[,1,1]), pch = '.', ylab = 'P-value')
abline(h=c(0.01,0.05),col=2:3)

scoreplot(plsModJ)
scoreplot(plsModJ, comps=c(1,3)) # Selected components
# Use MS categories for colouring and clusters for plot characters.
scoreplot(plsModJ, col = er$symbolicDesign[['MS']],
          pch = 20+as.numeric(er$symbolicDesign[['cluster']]))
loadingplot(plsModJ, scatter=TRUE) # scatter=TRUE for scatter plot
```

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