Package ‘DAMisc’

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Description Miscellaneous set of functions I use in my teaching either at the University of Western Ontario or the Inter-university Consortium for Political and Social Research (ICPSR) Summer Program in Quantitative Methods. Broadly, the functions help with presentation and interpretation of LMs and GLMs, but also implement some new tools like Alternating Least Squares Optimal Scaling for dependent variables, a Bayesian analog to the ALSOS algorithm. There are also tools to help understand interactions in both LMs and binary GLMs.
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Description

Functions to aid in the presentation of linear model results

Details

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</tbody>
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These are functions that help present linear model results. Largely, the represent alternatives in presentation to other R packages. For example, the factorplot function offers an alternative to David Firth's qvcalc package. This function calculates and presents exact variances of all simple contrasts. Both DAintfun and DAintfun2 are alternative ways of presenting interactions between two continuous variables. DAintfun2 gives results in line with the suggestions in Brambor, Clark and Golder (2006).

Author(s)

Dave Armstrong
Maintainer: Dave Armstrong <davearmstrong.ps@gmail.com>

References


Description

A subset of data from Alvarez et. al. (1996).
Format

A data frame with 4126 observations on the following 7 variables.

- **cname** Country name
- **country** Numeric country identifier
- **year** Year of observation
- **reg** A dichotomous variable coded 1 for dictatorship, 0 for democracy
- **gdpw** GDP/worker, 1985 prices
- **popg** Population growth
- **democ** A dichotomous variable coded 1 for democracy, 0 for dictatorship, (1-reg)

References


alsos

Alternating Least Squares Optimal Scaling

Description

Estimates the Alternating Least Squares Optimal Scaling (ALSOS) solution for qualitative variables.

Usage

```r
alsos(
  os_form,
  raw_form = ~1,
  data,
  scale_dv = FALSE,
  maxit = 30,
  level = 2,
  process = 1,
  starts = NULL,
  ...
)
```

Arguments

- **os_form** A two-sided formula including the independent variables to be scaled on the left-hand side. Optionally, the dependent variable can also be scaled.
- **raw_form** A right-sided formula with covariates that will not be scaled.
- **data** A data frame.
alsosDV

Description

This is a wrapper for the newer alsos function which allows optimal scaling of both dependent and independent variables. I retain the old operationalization of alsosDV for backward compatibility purposes.

Usage

alsosDV(form, data, maxit = 30, level = 2, process = 1, starts = NULL, ...)
Arguments

form A formula for a linear model where the dependent variable will be optimally
      scaled relative to the model.
data A data frame.
maxit Maximum number of iterations of the optimal scaling algorithm.
level Measurement level of the dependent variable 1=Nominal, 2=Ordinal
process Nature of the measurement process: 1=discrete, 2=continuous. Basically iden-
      tifies whether tied observations will continue to be tied in the optimally scaled
      variable (1) or whether the algorithm can untie the points (2) subject to the overall
      measurement constraints in the model.
starts Optional starting values for the optimal scaling algorithm.
... Other arguments to be passed down to lm.

Value

A list with the following elements:

result The result of the optimal scaling process
data The original data frame with additional columns adding the optimally scaled DV
iterations The iteration history of the algorithm
form Original formula

Author(s)

Dave Armstrong and Bill Jacoby

References


Young, Forrest, Jan de Leeuw and Yoshio Takane. 1976. ‘Regression with Qualitative and Quantitative Variables: An Alternating Least Squares Method with Optimal Scaling Features’ Psychometrika, 41:502-529.
aveEffPlot

Average Effect Plot for Generalized Linear Models

Description

For objects of class glm, it calculates the change the average predicted probability (like the one calculated by glmChange2) for a hypothetical candidate set of values of a covariate.

Usage

aveEffPlot(
obj, varname, data, R = 1500, nvals = 25, level = 0.95, ciType = c("percent", "normal"),
return = c("ci", "plot", "sim"),
...)

Arguments

obj A model object of class glm.
varname Character string giving the variable name for which average effects are to be calculated.
data Data frame used to fit object.
R Number of simulations to perform.
nvals Number of evaluation points at which the average probability will be calculated.
level Scalar giving the confidence level of the point-wise confidence intervals.

ciType Type of confidence interval to be created. If "perc", a percentile interval will be created from the distribution of effects. If "normal" a normal-theory interval will be calculated using the standard deviation of the fitted response from the simulation.
return Character string indicating what should be returned. Multiple entries are supported.
...

Other arguments to be passed down to xyplot.

Details

The function plots the average effect of a model covariate, for objects of class glm. The function does not work with poly unless the coefficients are provided as arguments to the command in the model (see example below).
Value

A plot or a data frame

Author(s)

Dave Armstrong

Examples

data(france)
p <- poly(france$lrself, 2)
left.mod <- glm(voteleft ~ male + age + retnat +
poly(lrself, 2, coefs=attr(p, "coefs")), data=france, family=binomial)
aveEffPlot(left.mod, "age", data=france, plot=FALSE)

balsos

Bayesian Alternating Least Squares Optimal Scaling

Description

Estimates a Bayesian analog to the Alternating Least Squares Optimal Scaling (ALSOS) solution for qualitative dependent variables.

Usage

balsos(
  formula,
  data,
  iter = 2500,
  chains = 1,
  alg = c("NUTS", "HMC", "Fixed_param"),
  ...
)

Arguments

formula A formula with a dependent variable that will be optimally scaled
data A data frame.
iter Number of samples for the MCMC sampler.
chains Number of parallel chains to be run.
alg Algorithm used to do sampling. See stan for more details.
... Other arguments to be passed down to stanfit.
Details

balsos estimates a Bayesian analog to the Alternating Least Squares Optimal Scaling solution on the dependent variable. This permits testing linearity assumptions on the original scale of the dependent variable.

Value

A list with the following elements:

- fit: The fitted stan output
- y: The dependent variable values used in the regression.
- X: The design matrix for the regression

Author(s)

Dave Armstrong

References


Young, Forrest, Jan de Leeuw and Yoshio Takane. 1976. ‘Regression with Qualitative and Quantitative Variables: An Alternating Least Squares Method with Optimal Scaling Features’ Psychometrika, 41:502-529.

BGMtest (obj, vars, digits = 3, level = 0.05, two.sided = TRUE)

Arguments

- obj: An object of class `lm`.
- vars: A vector of two variable names giving the two quantitative variables involved in the interaction. These variables must be involved in one, and only one, interaction.
- digits: Number of digits to be printed in the summary.
- level: Type I error rate for the tests.
- two.sided: Logical indicating whether the tests should be two-sided (if TRUE, the default) or one-sided (if FALSE).

Tests the five Berry, Golder and Milton (2012) Interactive Hypothesis

Description

This function tests the five hypotheses that Berry, Golder and Milton identify as important when two quantitative variables are interacted in a linear model.
binfit  

**Value**

A matrix giving five t-tests.

**Author(s)**

Dave Armstrong

**Examples**

```r
data(Duncan, package="carData")
mod <- lm(prestige ~ income*education + type, data=Duncan)
BGMtest(mod, c("income", "education"))
```

---

**binfit**  

*Scalar Measures of Fit for Binary Variable Models*

**Description**

Calculates scalar measures of fit for models with binary dependent variables along the lines described in Long (1997) and Long and Freese (2005).

**Usage**

```r
binfit(mod)
```

**Arguments**

- `mod`  
  
  A model of class `glm` with `family=binomial`.

**Details**

`binfit` calculates scalar measures of fit (many of which are pseudo-R-squared measures) to describe how well a model fits data with a binary dependent variable.

**Value**

A named vector of scalar measures of fit

**Author(s)**

Dave Armstrong
References


Examples

data(france)
left.mod <- glm(voteleft ~ male + age + retnat + poly(lrself, 2), data=france, family=binomial)
binfit(left.mod)

binVar

Bin a Variable

Description

Bins a continuous variable into a categorical variable

Usage

binVar(
  x, 
  bins = 4,
  method = c("intervals", "proportions"),
  labels = FALSE,
  include.lowest = TRUE,
  right = FALSE,
  ...
)

Arguments

x  Continuous variable to be binninged
bins  Number of groups in new variable
method  Method for generating "intervals" for fixed-width intervals and "proportions" for cut-points based on quantiles of the distribution.
labels  An optional vector of labels to apply to the groups
include.lowest  Logical indicating whether a value equal to the lowest (if right=TRUE) or highest (if right=FALSE) should be included.
right  Logical indicating Whether the intervals should be closed on the right and open on the left (if TRUE) or vice versa (if FALSE). Open intervals are those that do not include the end-point of the range and closed intervals do.
...  Other arguments to be passed down to cut

Function adapted from binVariable from the RcmdrMisc pakcage.
Value
A factor

Author(s)
John Fox

---

**boot.alsos**

*Bootstrapping function for the ALSOS algorithm*

**Description**

Executes a non-parametric bootstrap for the alsos algorithm to get uncertainty estimates for the optimally scaled values of the variables.

**Usage**

```r
boot.alsos(
  os_form,
  raw_form = ~1,
  data,
  scale_dv = TRUE,
  maxit = 30,
  level = 1,
  process = 1,
  starts = NULL,
  R = 50,
  conf.level = 0.95,
  return = c("data", "boot.obj", "both"),
  ...
)
```

**Arguments**

- `os_form` A two-sided formula including the independent variables to be scaled on the left-hand side. Optionally, the dependent variable can also be scaled.
- `raw_form` A right-sided formula with covariates that will not be scaled.
- `data` A data frame.
- `scale_dv` Logical indicating whether the dependent variable should be optimally scaled.
- `maxit` Maximum number of iterations of the optimal scaling algorithm.
- `level` Measurement level of the dependent variable: 1=Nominal, 2=Ordinal
- `process` Nature of the measurement process: 1=discrete, 2=continuous. Basically identifies whether tied observations will continue to be tied in the optimally scaled variable (1) or whether the algorithm can untie the points (2) subject to the overall measurement constraints in the model.
Optional starting values for the optimal scaling algorithm.

Number of bootstrap samples to be calculated

Level of confidence for the confidence intervals.

Whether the aggregated result with percentile confidence intervals, the bootstrap object or both should be returned.

Other arguments to be passed down to `lm`.

A list with either `data` and/or `boot.obj` entries.

Beck et. al. (1998) identified that binary time-series cross-section data are discrete-time duration data and time dependence can be modeled in a logistic regression by including a flexible function (e.g., cubic spline) of time since the last event as a covariate. This function creates the variable identifying time since last event.

btscs(data, event, tvar, csunit, pad.ts = FALSE)

A data frame.

Character string giving the name of the dichotomous variable identifying the event (where an event is coded 1 and the absence of an event is coded 0).

Character string giving the name of the time variable.

Character string giving the name of the cross-sectional unit.

Logical indicating whether the time-series should be filled in, when panels are unbalanced.

The original data frame with one additional variable. The `spell` variable identifies the number of observed periods since the last event.

Dave Armstrong
```r
library(splines)
## Data from Alvarez et al. (1996)
data(aclp)
newdat <- btscs(aclp, "democ", "year", "country")

# Estimate Model with and without spell
full.mod <- glm(democ ~ log(gdpw) + popg + bs(spell, df=4), data=newdat, family=binomial)
restricted.mod <- glm(democ ~ log(gdpw) + popg, data=newdat, family=binomial)

# Incremental F-test of time dependence
anova(restricted.mod, full.mod, test="Chisq")
```

---

**References**


**Examples**

```r
library(splines)
## Data from Alvarez et al. (1996)
data(aclp)

newdat <- btscs(aclp, "democ", "year", "country")

# Estimate Model with and without spell
full.mod <- glm(democ ~ log(gdpw) + popg + bs(spell, df=4), data=newdat, family=binomial)
restricted.mod <- glm(democ ~ log(gdpw) + popg, data=newdat, family=binomial)

# Incremental F-test of time dependence
anova(restricted.mod, full.mod, test="Chisq")
```

---

**cat2Table**

*Fitted Values and CIs for 2-Categorical Interactions*

**Description**

This function makes a table of fitted values and confidence intervals for all of the combinations of two categorical variables in an interaction.

**Usage**

```r
cat2Table(eff.obj, digits = 2, rownames = NULL, colnames = NULL)
```

**Arguments**

- `eff.obj`: An object generated by `effect` from the `effects` package where the effect is calculated for two factors involved in an interaction.
- `digits`: Number of digits of the fitted values and confidence intervals to print.
- `rownames`: An optional vector of row names for the table, if `NULL`, the levels of the factor will be used.
- `colnames`: An optional vector of column names for the table, if `NULL`, the levels of the factor will be used.

**Value**

A matrix of fitted values and confidence intervals.
**Examples**

```r
library(effects)
data(Duncan, package="carData")
Duncan$inc.cat <- cut(Duncan$income, 3)
mod <- lm(prestige ~ inc.cat*type + income, data=Duncan)
e1 <- effect("inc.cat*type", mod)
cat2Table(e1)
```

---

**Description**

Calculates the regions of statistical significance in interaction and identifies the points at which the statistical significance of conditional coefficients changes.

**Usage**

`changeSig(obj, vars, alpha = 0.05)`

**Arguments**

- `obj` A model of class `glm` or class `lm`.
- `vars` A character vector of the names of the two variables involved in the interaction.
- `alpha` Critical p-value of the test.

**Value**

Printed output that identifies the change-points in statistical significance.

**Author(s)**

Dave Armstrong
combTest

Test for Combining Categories in Multinomial Logistic Regression Models.

Description

Tests the null hypothesis that categories can be combined in Multinomial Logistic Regression Models.

Usage

combTest(obj)

Arguments

obj

An object of class multinom.

Value

A matrix of test statistics and p-values.

Author(s)

Dave Armstrong

Examples

library(nnet)
data(france)
mnl.mod <- multinom(vote ~ age + male + retnat + lrself, data=france)
combTest(mnl.mod)

crSpanTest

Test of Span Parameter in linearity for Component + Residual Plots

Description

This function performs crTest for a user-defined range of span parameters, optionally allowing for multiple testing corrections in the p-values.
Usage

```
crSpanTest(
    model,  # A model object of class lm
    spfromto,  # A vector of two values across which a range of n span values will be generated and tested.
    n = 10,  # Number of span parameters to test.
    adjust.method = "none",  # Adjustment method for multiple-testing procedure, using p.adjust from stats.
    adjust.type = c("none", "across", "within", "both")
)
```

Arguments

- `model`: A model object of class `lm`.
- `spfromto`: A vector of two values across which a range of `n` span values will be generated and tested.
- `n`: Number of span parameters to test.
- `adjust.type`: String giving the values over which the multiple testing correction will be performed. Here, ‘both’ refers to a multiple testing correction done over all span parameters and all variables in the model. ‘within’ means the multiple testing correction should be done within each model, but not across the span parameters and ‘across’ means that the multiple testing correction should be for each variable across the various span parameters, but not across variables within the same model. ‘none’ refers to a pass-through option of no multiple testing procedure.

Value

A list with two elements:

- `x`: Sequence of span values used in testing
- `y`: p-values for each variable for each span parameter

Author(s)

- Dave Armstrong

Examples

```
data(Prestige, package="carData")
mod <- lm(prestige ~ income + education + women, data=Prestige)
tmp <- crSpanTest(mod, c(.1, .9), adjust.method="holm",
                  adjust.type="both")
matplot(tmp$x, tmp$y, type="l")
```
Description

This function estimates a linear model and a loess model on the component-plus-residual plot (i.e., a partial residual plot) for each quantitative variable in the model. The residual sums of squares for each are used to calculate an F-test for each quantitative variable.

Usage

crTest(
  model,
  adjust.method = "none",
  cat = 5,
  var = NULL,
  span.as = TRUE,
  span = 0.75,
  ...
)

Arguments

model A model object of class lm
adjust.method Adjustment method for multiple-testing procedure, using p.adjust from stats.
cat Number of unique values below which numeric variables are considered categorical for the purposes of the smooth.
var Character string indicating the term desired for testing. If left NULL, the default value, all numeric variables will be tested.
span.as Logical indicating whether the span should be automatically selected through AICC or GCV
span Span to be passed down to the loess function if span.as=FALSE.
... Other arguments to be passed down to the call to loess.

Value

A matrix with the following columns for each variable:

RSSp Residual sum-of-squares for the parametric (linear) model.
RSSnp Residual sum-of-squares for the non-parametric (loess) model.
DFnum Numerator degrees of freedom for the F-test: tr(S)-(k+1).
DFdenom Denominator degrees of freedom for the F-test: n-tr(S)
F F-statistic
p p-value, potentially adjusted for multiple comparisons.
Author(s)
Dave Armstrong

Examples

data(Prestige, package="carData")
mod <- lm(prestige ~ income + education + women, data=Prestige)
crTest(mod)

---

**cv.lo2**

Cross-validating Loess curve

Description

Function provides the cross-validation error for the loess curve in a manner that is amenable to optimization of the span.

Usage

```r
## S3 method for class 'lo2'
cv(span, form, data, cost = function(y, yhat) mean((y - yhat)^2, na.rm = TRUE),
   K = n,
   numiter = 100,
   which = c("corrected", "raw")
)
```

Arguments

- `span`: The span of the loess smoother.
- `form`: The formula that identifies the model
- `data`: A data frame containing the required variables.
- `cost`: Cost function to be passed down to loess.
- `K`: Number of folds for the cross-validation
- `numiter`: Number of times over which the cv error will be aggregated
- `which`: Return raw or corrected cv error

Value

The cross-validation error from the loess curve.
DAintfun

Author(s)

Dave Armstrong

DAintfun (Surface Plots for Two-Way Interactions)

Description

Makes surface plots to display interactions between two continuous variables

Usage

DAintfun(
  obj,
  varnames,
  theta = 45,
  phi = 10,
  xlab = NULL,
  ylab = NULL,
  zlab = NULL,
  adjustY = 0,
  plot = TRUE,
  hcols = NULL,
  ...
)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>obj</td>
<td>A model object of class lm</td>
</tr>
<tr>
<td>varnames</td>
<td>A two-element character vector where each element is the name of a variable involved in a two-way interaction.</td>
</tr>
<tr>
<td>theta</td>
<td>Angle defining the azimuthal viewing direction to be passed to persp</td>
</tr>
<tr>
<td>phi</td>
<td>Angle defining the colatitude viewing direction to be passed to persp</td>
</tr>
<tr>
<td>xlab</td>
<td>Optional label to put on the x-axis, otherwise if NULL, it will take the first element of varnames</td>
</tr>
<tr>
<td>ylab</td>
<td>Optional label to put on the y-axis, otherwise if NULL, it will take the second element of varnames</td>
</tr>
<tr>
<td>zlab</td>
<td>Optional label to put on the z-axis, otherwise if NULL, it will be ‘Predictions’.</td>
</tr>
<tr>
<td>adjustY</td>
<td>Scalar indicating a constant that should be added to all of fitted values. Defaults to 0.</td>
</tr>
<tr>
<td>plot</td>
<td>Logical indicating whether the plot should be returned. If FALSE, the data are returned instead.</td>
</tr>
<tr>
<td>hcols</td>
<td>Vector of four colors to color increasingly high density areas</td>
</tr>
<tr>
<td>...</td>
<td>Other arguments to be passed down to the initial call to persp</td>
</tr>
</tbody>
</table>
Details
This function makes a surface plot of an interaction between two continuous covariates. If the model is
\[ y_i = b_0 + b_1 x_{i1} + b_2 x_{i2} + b_3 x_{i1} \times x_{i2} + \ldots + e_i, \]
this function plots \( b_1 x_{i1} + b_2 x_{i2} + b_3 x_{i1} \times x_{i2} \) for values over the range of \( X_1 \) and \( X_2 \). The highest 75%, 50% and 25% of the bivariate density of \( X_1 \) and \( X_2 \) (as calculated by \texttt{sm.density} from the \texttt{sm} package) are colored in with colors of increasing gray-scale.

Value
- \( x1 \): Values of the first element of \texttt{varnames} used to make predictions.
- \( x2 \): Values of the second element of \texttt{varnames} used to make predictions.
- \( \text{pred} \): The predictions based on the values \( x1 \) and \( x2 \).
- \( \text{graph} \): A graph is produced, but no other information is returned.

Author(s)
Dave Armstrong

Examples
```r
data(InteractionEx)
mod <- lm(y ~ x1*x2 + z, data=InteractionEx)
DAintfun(mod, c("x1", "x2"))
```

Description
Generates two conditional effects plots for two interacted continuous covariates in linear models.

Usage
```r
DAintfun2(
  obj,
  varnames,
  varcov = NULL,
  rug = TRUE,
  ticksize = -0.03,
  hist = FALSE,
  level = 0.95,
```
hist.col = "gray75",
nclass = c(10, 10),
scale.hist = 0.5,
border = NA,
name.stem = "cond_eff",
xlab = NULL,
ylab = NULL,
plot.type = c("screen", "pdf", "png", "eps", "none")
)

Arguments

obj A model object of class \texttt{lm}

varnames A two-element character vector where each element is the name of a variable involved in a two-way interaction.

varcov A variance-covariance matrix with which to calculate the conditional standard errors. If \texttt{NULL}, it is calculated with \texttt{vcov(obj)}.

rug Logical indicating whether a rug plot should be included.

ticksize A scalar indicating the size of ticks in the rug plot (if included) positive values put the rug inside the plotting region and negative values put it outside the plotting region.

hist Logical indicating whether a histogram of the x-variable should be included in the plotting region.

level Level for the confidence bounds.

hist.col Argument to be passed to \texttt{polygon} indicating the color of the histogram bins.

nclass vector of two integers indicating the number of bins in the two histograms, which will be passed to \texttt{hist}.

scale.hist A scalar in the range (0,1] indicating how much vertical space in the plotting region the histogram should take up.

border Argument passed to \texttt{polygon} indicating how the border of the histogram bins should be printed (\texttt{NA} for no border).

name.stem A character string giving filename to which the appropriate extension will be appended.

xlab Optional vector of length two giving the x-labels for the two plots that are generated. The first element of the vector corresponds to the figure plotting the conditional effect of the first variable in \texttt{varnames} given the second and the second element of the vector corresponds to the figure plotting the conditional effect of the second variable in \texttt{varnames} conditional on the first.

ylab Optional vector of length two giving the y-labels for the two plots that are generated. The first element of the vector corresponds to the figure plotting the conditional effect of the first variable in \texttt{varnames} given the second and the second element of the vector corresponds to the figure plotting the conditional effect of the second variable in \texttt{varnames} conditional on the first.

plot.type One of \texttt{pdf}, \texttt{png}, \texttt{eps} or \texttt{screen}, where the one of the first three will produce two graphs starting with \texttt{name.stem} written to the appropriate file type and the third will produce graphical output on the screen.
Details

This function produces graphs along the lines suggested by Brambor, Clark and Golder (2006) and Berry, Golder and Milton (2012), that show the conditional effect of one variable in an interaction given the values of the conditioning variable. This is an alternative to the methods proposed by John Fox in his effects package, upon which this function depends heavily.

Specifically, if the model is

\[ y_i = b_0 + b_1 x_{i1} + b_2 x_{i2} + b_3 x_{i1} \times x_{i2} + \ldots + e_i, \]

this function plots calculates the conditional effect of \( X_1 \) given \( X_2 \)

\[ \frac{\partial y}{\partial X_1} = b_1 + b_3 X_2 \]

and the variances of the conditional effects

\[ V(b_1 + b_3 X_2) = V(b_1 + X_2^2 V(b_3) + 2(1)(X_2) V(b_1, b_3)) \]

for different values of \( X_2 \) and then switches the places of \( X_1 \) and \( X_2 \), calculating the conditional effect of \( X_2 \) given a range of values of \( X_1 \). 95% confidence bounds are then calculated and plotted for each conditional effects along with a horizontal reference line at 0.

Value
graphs

Either a single graph is printed on the screen (using `par(mfrow=c(1,2))`) or two figures starting with `name.stem` are produced where each gives the conditional effect of one variable based on the values of another.

Author(s)

Dave Armstrong

References


Examples

data(InteractionEx)
mod <- lm(y ~ x1*x2 + z, data=InteractionEx)
DAintfun2(mod, c("x1", "x2"), hist=TRUE, scale.hist=.3)
DAintfun3  Conditional Effects Plots for Interactions in Linear Models

Description
Generates two conditional effects plots for two interacted continuous covariates in linear models.

Usage
DAintfun3(
  obj,
  varnames,
  varcov = NULL,
  name.stem = "cond_eff",
  xlab = NULL,
  ylab = NULL,
  plot.type = "screen"
)

Arguments
obj A model object of class lm
varnames A two-element character vector where each element is the name of a variable involved in a two-way interaction.
varcov A variance-covariance matrix with which to calculate the conditional standard errors. If NULL, it is calculated with vcov(obj).
name.stem A character string giving filename to which the appropriate extension will be appended
xlab Optional vector of length two giving the x-labels for the two plots that are generated. The first element of the vector corresponds to the figure plotting the conditional effect of the first variable in varnames given the second and the second element of the vector corresponds to the figure plotting the conditional effect of the second variable in varnames conditional on the first.
ylab Optional vector of length two giving the y-labels for the two plots that are generated. The first element of the vector corresponds to the figure plotting the conditional effect of the first variable in varnames given the second and the second element of the vector corresponds to the figure plotting the conditional effect of the second variable in varnames conditional on the first.
plot.type One of ‘pdf’, ‘png’, ‘eps’ or ‘screen’, where the one of the first three will produce two graphs starting with name.stem written to the appropriate file type and the third will produce graphical output on the screen.

Details
This function does the same thing as DAintfun2, but presents effects only at the mean of the conditioning variable and the mean +/- 1 standard deviation.
Value

graphs Either a single graph is printed on the screen (using `par(mfrow=c(1,2))`) or two figures starting with `name.stem` are produced where each gives the conditional effect of one variable based on the values of another.

Author(s)

Dave Armstrong

References


Examples

data(InteractionEx)
mod <- lm(y ~ x1*x2 + z, data=InteractionEx)
DAintfun3(mod, c("x1", "x2"))

---

**describe_data**

*Interactive List of Variables*

Description

Makes an interactive list of variables and their descriptive labels. Uses the **DT** package to generate the table.

Usage

`describe_data(data)`

Arguments

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>data</td>
<td>A dataset to be described.</td>
</tr>
</tbody>
</table>
Description

Plots the effect of a variable in a model estimated with Firth Logit.

Usage

effect_logistf(var, obj, data, ...)

Arguments

var A character string giving the name of the variable whose effect is to be generated.
obj An object of class logistf.
data A data frame.
... Other arguments to be passed down to the Effect function.

Details

The effect.logistf function calculates the effect (predicted probabilities) of a variable in a Firth logit model estimated with the logistf function. The function estimates the analogous glm. It then replaces the coefficient vector in that model object with the Frith logit coefficients. It also puts the variance-covariance matrix from the Firth logit in the model object and uses a custom extractor function in the Effect function to extract that variance-covariance matrix rather than the one usually extracted with vcov. Note that variability and confidence intervals for the effects will not be calculated using profile likelihood as they are in the Firth logit, but will be calculated using the appropriate variance-covariance matrix.

Value

An object of class eff that can be used with other functions from the effects package.

Author(s)

Dave Armstrong
france  

Example data for factorplot function

Description
A subset of data from the 1994 Eurobarometer for France

Format
A data frame with 542 observations on the following 5 variables.

- lrself: respondent’s left-right self-placement on a 1(left)-10(right) scale
- male: a dummy variable coded 1 for males and 0 for females
- age: respondent’s age
- vote: a factor indicating vote choice with levels PCF, PS, Green, RPR and UDF
- retnat: a factor indicating the respondent’s retrospective national economic evaluation with levels Better, Same and Worse
- voteleft: a dichotomous variable where 1 indicates a vote for a left party, 0 otherwise

References

ggpie  

Pie Charts with ggplot2

Description
This function produces pie charts with ggplot2.

Usage

```r
ggpie(data, variable, addPct = c("pie", "none", "legend"))
```

Arguments

- data: A data frame to pass to the ggplot function
- variable: A variable to be plotted.
- addPct: Where labels should be added - "none" gives no labels, "legend" adds percentages to the color legend, "pie" adds the legends to the pie pieces.

Value

a ggplot
**glmChange**  
*Maximal First Differences for Generalized Linear Models*

**Description**

For objects of class `glm`, it calculates the change in predicted responses, for maximal discrete changes in all covariates holding all other variables constant at typical values.

**Usage**

```r
glmChange(  
  obj,  
  data,  
  V = NULL,  
  typical.dat = NULL,  
  change.dat = NULL,  
  diffchange = c("range", "sd", "unit"),  
  outcome = c("diff", "maxdiff"),  
  n = 1,  
  catdiff = c("biggest", "all"),  
  sim = FALSE,  
  R = 1000,  
  qtiles = c(0.025, 0.975),  
  adjust = c("none", "shift", "trim")  
)
```

**Arguments**

- `obj`: A model object of class `glm`.
- `data`: Data frame used to fit `object`.
- `V`: An optional variance-covariance matrix for the coefficients, if NULL, will be obtained through a call to `vcov`.
- `typical.dat`: Data frame with a single row containing values at which to hold variables constant when calculating first differences. These values will be passed to `predict`, so factors must take on a single value, but have all possible levels as their levels attribute.
- `change.dat`: A named list of values over which the variables of interest will be changed. If NULL or partially specified, unspecified variables will use `diffchange`. If a categorical variable is specified in here, this overrides the `catdiff` argument and only the specified contrast will be generated.
- `diffchange`: A string indicating the difference in predictor values to calculate the discrete change. `range` gives the difference between the minimum and maximum, `sd` gives plus and minus one-half standard deviation change around the median and `unit` gives a plus and minus one-half unit change around the median.
outcome

For quantitative variables, should the difference over the range of chosen values be calculated (the default) or should the maximum probability difference over the range be calculated. These will be the same for single-term quantitative variables, but could be different for multi-term variables, like splines and polynomials.

n

number of units of diffchange to move. Only active for unit or sd.

catdiff

String identifying how differences in factor variables is handled. Options are "all" in which case all pairwise differences are returned, or "biggest" in which case the biggest difference is returned.

sim

Logical indicating whether simulated confidence bounds on the difference should be calculated and presented.

R

Number of simulations to perform if sim is TRUE

qtiles

Quantiles to calculate if sim=TRUE.

adjust

String identifying how range should be changed if it goes out of the bounds of the observed data. Trimming will simply truncate the size of the change to make it fit in bounds. Shifting will shift the interval so both ends are in bounds. If the shifted interval is wider than the range of the data, the change will be truncated to the range of the data.

Details

The function calculates the changes in predicted responses for maximal discrete changes in the covariates, for objects of class glm. This function works with polynomials specified with the poly function. It also works with multiplicative interactions of the covariates by virtue of the fact that it holds all other variables at typical values. By default, typical values are the median for quantitative variables and the mode for factors. The way the function works with factors is a bit different. The function identifies the two most different levels of the factor and calculates the change in predictions for a change from the level with the smallest prediction to the level with the largest prediction.

Value

A list with the following elements:

diffs A matrix of calculated first differences

minmax A matrix of values that were used to calculate the predicted changes

Author(s)

Dave Armstrong

Examples

data(france)
left.mod <- glm(voteleft ~ male + age + retnat + poly(lrself, 2), data=france, family=binomial)
typical.france <- data.frame(  
  retnat = factor(1, levels=1:3, labels=levels(france$retnat)),  
)
For objects of class `glm`, it calculates the change in predicted responses, for discrete changes in a covariate holding all other variables at their observed values.

Usage

```r
glmChange2(
  obj, 
  varname, 
  data, 
  V = NULL, 
  diffchange = c("unit", "sd"), 
  outcome = c("diff", "maxdiff"), 
  baseline = c("obs", "median"), 
  catdiff = c("biggest", "all"), 
  n = 1, 
  R = 1500, 
  adjust = c("none", "shift", "trim"), 
  ...
)
```

Arguments

- **obj**: A model object of class `glm`.
- **varname**: Character string giving the variable name for which average effects are to be calculated.
- **data**: Data frame used to fit `object`.
- **V**: An optional variance-covariance matrix for the coefficients, if `NULL`, will be obtained through a call to `vcov`.
- **diffchange**: A string indicating the difference in predictor values to calculate the discrete change. `sd` gives plus and minus one-half standard deviation change around the median and `unit` gives a plus and minus one-half unit change around the median.
- **outcome**: For quantitative variables, should the difference over the range of chosen values be calculated (the default) or should the maximum probability difference over the range be calculated. These will be the same for single-term quantitative variables, but could be different for multi-term variables, like splines and polynomials.
baseline Character string representing the baseline to use for the change. It can be one of "obs", in which case each observations value is used as the baseline or "median", in which case the median is used as a common baseline for all observations.

catdiff String identifying how differences in factor variables is handled. Options are "all" in which case all pairwise differences are returned, or "biggest" in which case the biggest difference is returned.

n Number of diffchange to move.

R Number of simulations to perform.

adjust String identifying how range should be changed if it goes out of the bounds of the observed data. Trimming will simply truncate the size of the change to make it fit in bounds. Shifting will shift the interval so both ends are in bounds. If the shifted interval is wider than the range of the data, the change will be truncated to the range of the data.

... Allows user to specify legacy argument change

Details

The function calculates the average change in predicted probability for a discrete change in a single covariate with all other variables at their observed values, for objects of class glm. This function works with polynomials specified with the poly function.

Value

res A vector of values giving the average and 95 percent confidence bounds

ames The average change in predicted probability (across all N observations) for each of the R simulations.

avesamp The average change in predicted probability for each of the N observation (across all of the R simulations).

Author(s)

Dave Armstrong

Examples

data(france)
left.mod <- glm(voteleft ~ male + age + retnat + poly(lrself, 2), data=france, family=binomial)
glmChange2(left.mod, "age", data=france, diffchange="sd")
**impCoef**

*Plot Variable Importance.*

**Description**

Builds a plot of importance based on Silber, Rosenbaum and Ross’s (1995) idea of importance as the variance of the predicted model terms.

**Usage**

```r
impCoef(obj, pct = FALSE, names = NULL, orderSize = TRUE)
```

**Arguments**

- `obj`: Any object that permits prediction of model terms using the `predict(obj, type="terms")` syntax.
- `pct`: Logical indicating whether the entries should be percentagized by the total sum of squares in the predictions.
- `names`: An optional vector of names for the coefficients.
- `orderSize`: Logical indicating whether the terms are ordered by importance in the graph.

**Value**

Returns an initialized ggplot, but geometries need to be added to produce meaningful output (see examples).

**References**


**Examples**

```r
data(aclp)
library(ggplot2)
mod <- glm(democ ~ log(gdpw) + popg + year, data=aclp, family=binomial)
impCoef(mod, pct=TRUE, names=c("GDP", "Population", "Year")) +
  geom_point(size=2) +
  labs(x="Importance", y="")
```
inspect  
*Inspect a Variable in a Data Frame*

**Description**

Shows the variable label, factor levels (i.e., value labels) and frequency distribution for the specified variable.

**Usage**

```r
inspect(data, x, includeLabels = FALSE, ...)
```

**Arguments**

- `data`: A data frame of class `data.frame` or `tbl_df`.
- `x`: A string identifying the name of the variable to be inspected.
- `includeLabels`: Logical indicating whether value labels should also be included.
- `...`: Other arguments to be passed down, currently unimplemented.

**Value**

A list with a variable label (if present), factor levels/value labels (if present) and a frequency distribution

**Author(s)**

Dave Armstrong

**Examples**

```r
data(france)
inspect(france, "vote")
```

---

**intEff  
*Functions for Estimating Interaction Effects in Logit and Probit Models***

**Description**

Usage

\texttt{intEff(obj, vars, data)}

Arguments

- \texttt{obj} A binary logit or probit model estimated with \texttt{glm}.
- \texttt{vars} A vector of the two variables involved in the interaction.
- \texttt{data} A data frame used in the call to \texttt{obj}.

Value

A list is returned with two elements - byobs and atmean. The byobs result gives the interaction effect evaluated at each observation. The atmean element has the marginal effect evaluated at the mean. Each element contains an element \texttt{int} which is a data frame with the following variable:

- \texttt{int_eff} The correctly calculated marginal effect.
- \texttt{linear} The incorrectly calculated marginal effect following the linear model analogy.
- \texttt{phat} Predicted \Pr(Y=1|X).
- \texttt{se_int_eff} Standard error of \texttt{int_eff}.
- \texttt{zstat} The interaction effect divided by its standard error

The \texttt{X} element of each returned result is the \texttt{X}-matrix used to generate the result.

Author(s)

Dave Armstrong

References


Examples

\begin{verbatim}
data(france)
mod <- glm(voteleft ~ age*lrself + retnat + male, data=france, family=binomial)
out <- intEff(obj=mod, vars=c("age", "lrself"), data=france)
out <- out$byobs$int
plot(out$phat, out$int_eff, xlab="Predicted \Pr(Y=1|X)", ylab = "Interaction Effect")
ag <- aggregate(out$linear, list(out$phat), mean)
lines(ag[,1], ag[,2], lty=2, col="red", lwd=2)
\end{verbatim}
InteractionEx

Example Data for DAintfun

Description
Data to execute example code for DAintfun

Format
A data frame with 500 observations on the following 4 variables.

- y a numeric vector
- x1 a numeric vector
- x2 a numeric vector
- z a numeric vector

Details
These are randomly generated data to highlight the functionality of DAintfun

intQualQuant

Predictions for Factor-Numeric Interactions in Linear Models

Description
This function works on linear models with a single interaction between a continuous (numeric) variable and a factor. The output is a data frame that gives the predicted effect of moving from each category to each other category of the factor over the range of values of the continuous conditioning variable.

Usage
intQualQuant(
  obj,
  vars,
  level = 0.95,
  varcov = NULL,
  labs = NULL,
  n = 10,
  onlySig = FALSE,
  type = c("facs", "slopes"),
);
\begin{verbatim}
plot = TRUE,
vals = NULL,
rug = TRUE,
ci = TRUE,
digits = 3,
...

Arguments

obj An object of class \texttt{lm}.
vars A vector of two variable names giving the two quantitative variables involved in the interaction. These variables must be involved in one, and only one, interaction.
level Confidence level desired for lower and upper bounds of confidence interval.
varcov A potentially clustered or robust variance-covariance matrix of parameters used to calculate standard errors. If \texttt{NULL}, the \texttt{vcov} function will be used.
labs An optional vector of labels that will be used to identify the effects, if \texttt{NULL}, the factor levels will be used.
n Number of values of the conditioning variable to use.
onlySig Logical indicating whether only contrasts with significant differences should be returned. Significance is determined to exist if the largest lower bound is greater than zero or the smallest upper bound is smaller than zero.
type String indicating whether the conditional partial effect of the factors is plotted (if ‘facs’), or the conditional partial effect of the quantitative variable (if ‘slopes’) is produced.
plot Logical indicating whether graphical results (if \texttt{TRUE}) or numerical results (if \texttt{FALSE}) are produced.
vals A vector of values at which the continuous variable will be held constant. If \texttt{NULL}, a sequence of length \texttt{n} across the variable’s range will be used.
rug Logical indicating whether rug plots should be plotted in the panels.
cl Logical indicating whether confidence bounds should be drawn.
digits Number indicating how many decimal places to round the numeric output.
...
Other arguments to be passed down to \texttt{effect} if \texttt{plot.type = ‘slopes’}.

Value

For \texttt{type = ‘facs’} and \texttt{plot = FALSE}, a data frame with the following values:

\begin{verbatim}
fit The expected difference between the two factor levels at the specified value of the conditioning variable.
se.fit The standard error of the expected differences.
x The value of the continuous conditioning variable
contrast A factor giving the two values of the factor being evaluated.
\end{verbatim}
\end{verbatim}
lower The lower 95% confidence interval for fit
upper The upper 95% confidence interval for fit

For type = 'facs' and plot = TRUE, a lattice display is returned For type = 'slopes' and plot = FALSE, A character matrix with the following columns:

B The conditional effect of the quantitative variable for each level of the factor.
SE(B) The standard error of the conditional effect.
t-stat The t-statistic of the conditional effect.
Pr(>|t|) The two-sided p-value.

For type = 'slopes' and plot = TRUE, a lattice display is returned

Author(s)
Dave Armstrong

Examples

data(Prestige, package="carData")
Prestige$income <- Prestige$income/1000
mod <- lm(pre prestige - income * type + education, data=Prestige)
intQualQuant(mod, c("income", "type"), n=10, plot.type="none")
intQualQuant(mod, c("income", "type"), n=10, plot.type="facs")
intQualQuant(mod, c("income", "type"), n=10, plot.type="slopes")

loess.as

Fit a local polynomial regression with automatic smoothing parameter selection

Description
Fit a local polynomial regression with automatic smoothing parameter selection. Two methods are available for the selection of the smoothing parameter: bias-corrected Akaike information criterion (aicc); and generalized cross-validation (gcv).

Usage

loess.as(
  x,
  y,
  degree = 1,
  criterion = c("aicc", "gcv"),
)
```r
family = c("gaussian", "symmetric"),
user.span = NULL,
plot = FALSE,
...
)

as.crit(x)

opt.span(model, criterion = c("aicc", "gcv"), span.range = c(0.05, 0.95))
```

**Arguments**

- `x` An object of class `loess`.
- `y` a vector of response values
- `degree` the degree of the local polynomials to be used. It can be 0, 1 or 2.
- `criterion` The criterion used to find the optimal span
- `family` if "gaussian" fitting is by least-squares, and if "symmetric" a re-descending M estimator is used with Tukey's biweight function.
- `user.span` the user-defined parameter which controls the degree of smoothing.
- `plot` if TRUE, the fitted curve or surface will be generated.
- `...` control parameters.

**Value**

An object of class `loess`.

**Author(s)**

X.F. Wang

**Examples**

```r
## Fit Local Polynomial Regression with Automatic Smoothing Parameter Selection
n1 <- 100
x1 <- runif(n1, min=0, max=3)
sd1 <- 0.2
e1 <- rnorm(n1, sd=sd1)
y1 <- sin(2*x1) + e1
(y1.fit <- loess.as(x1, y1, plot=TRUE))
```
loessDeriv  

Estimate Derivatives of LOESS Curve.

Description

Estimates the first derivatives of the LOESS curve.

Usage

loessDeriv(obj, delta = 1e-05)

Arguments

obj An object of class loess.
delta Small change to be induced to estimate derivative.

Value

A vector of first derivative values evaluated at each original x-value.

Author(s)

Dave Armstrong

logit_cc  

Functions for Estimating Interaction Effects in Logit and Probit Models

Description

Norton and Ai (2003) and Norton, Wang and Ai (2004) discuss methods for calculating the appropriate marginal effects for interactions in binary logit/probit models. These functions are direct translations of the Norton, Wang and Ai (2004) Stata code. These functions are not intended to be called by the user directly, rather they are called as needed by intEff.

Usage

logit_cc(obj = obj, int.var = int.var, vars = vars, b = b, X = X)

Arguments

obj A binary logit or probit model estimated with glm.
int.var The name of the interaction variable.
vars A vector of the two variables involved in the interaction.
b Coefficients from the glm object.
X Model matrix from the glm object.
**Value**

A data frame with the following variable:

- `int_eff`: The correctly calculated marginal effect.
- `linear`: The incorrectly calculated marginal effect following the linear model analogy.
- `phat`: Predicted Pr(Y=1|X).
- `se_int_eff`: Standard error of `int_eff`.
- `zstat`: The interaction effect divided by its standard error.

**Author(s)**

Dave Armstrong

**References**


---

**makeHypSurv**

*Make Hypothetical Predictions for Survey Data*

**Description**

Calculates survival probabilities for hypothetical data.

**Usage**

`makeHypSurv(l, obj, ...)`

**Arguments**

- `l`: A named list where variable names are the names and values of the variables are the values. Combinations will be made with `expand.grid`.
- `obj`: A model object estimated with `survreg`.
- `...`: currently not implemented.

**Value**

A data frame.
Author(s)
Dave Armstrong

make_assoc_stats  Make Categorical Association Statistics

Description
Makes several common measures of association for contingency tables. The p-values are obtained through simulation.

Usage
make_assoc_stats(
  x, 
  y, 
  chisq = FALSE, 
  phi = FALSE, 
  cramersV = TRUE, 
  lambda = FALSE, 
  gamma = TRUE, 
  d = FALSE, 
  taub = TRUE, 
  n = 1000, 
  weight = NULL
)

Arguments
x  The row-variable in a contingency table
y  The column-variable in a contingency table
chisq Logical indicating whether the chi-squared statistic should be produced.
phi Logical indicating whether the phi statistic should be produced.
cramersV Logical indicating whether the Cramer’s V statistic should be produced.
lambda Logical indicating whether the lambda statistic should be produced.
gamma Logical indicating whether the gamma statistic for ordinal data should be produced.
d Logical indicating whether Somer’s D for ordinal data should be produced.
taub Logical indicating whether Kendall’s Tau-b statistic should be produced.
n Number of iterations in the simulation.
weight Vector of weights used to generate the table.

Value
A matrix of statistics and p-values.
**mnlAveEffPlot**  
*Average Effects Plot for Multinomial Logistic Regression*

**Description**

Produces a plot of average effects for one variable while holding the others constant at observed values.

**Usage**

```r
mnlAveEffPlot(obj, varname, data, R = 1500, nvals = 25, plot = TRUE, ...)
```

**Arguments**

- `obj`: An object of class `multinom`.
- `varname`: A string indicating the variable for which the plot is desired.
- `data`: The data used to estimate `obj`.
- `R`: Number of simulations used to generate confidence bounds.
- `nvals`: Number of evaluation points for the predicted probabilities.
- `plot`: Logical indicating whether a plot should be produced (if `TRUE`) or numerical results should be returned (if `FALSE`).
- `...`: Other arguments to be passed down to `xyplot`.

**Value**

Either a plot or a data frame with variables

- `mean`: The average effect (i.e., predicted probability)
- `lower`: The lower 95% confidence bound
- `upper`: The upper 95% confidence bound
- `y`: The values of the dependent variable being predicted
- `x`: The values of the independent variable being manipulated

**Author(s)**

Dave Armstrong

**References**

Examples

library(nnet)
data(france)
mnl.mod <- multinom(vote ~ age + male + retnat + lrself, data=france)
## Not run: mnlAveEffPlot(mnl.mod, "lrself", data=france)

mnlChange

Maximal First Differences for Multinomial Logistic Regression Models

Description

For objects of class multinom, it calculates the change in predicted probabilities, for maximal discrete changes in all covariates holding all other variables constant at typical values.

Usage

mnlChange(
  obj,
  data,
  typical.dat = NULL,
  diffchange = c("range", "sd", "unit"),
  n = 1,
  sim = TRUE,
  R = 1500
)

Arguments

obj A model object of class multinom.
data Data frame used to fit object.
typical.dat Data frame with a single row containing values at which to hold variables constant when calculating first differences. These values will be passed to predict, so factors must take on a single value, but have all possible levels as their levels attribute.
diffchange A string indicating the difference in predictor values to calculate the discrete change. range gives the difference between the minimum and maximum, sd gives plus and minus one-half standard deviation change around the median and unit gives a plus and minus one-half unit change around the median.
n Number of diffchange units to change.
sim Logical indicating whether simulated confidence bounds should be produced.
R Number of simulations to perform if sim = TRUE
Details

The function calculates the changes in predicted probabilities for maximal discrete changes in the covariates for objects of class `multinom`. This function works with polynomials specified with the `poly` function. It also works with multiplicative interactions of the covariates by virtue of the fact that it holds all other variables at typical values. By default, typical values are the median for quantitative variables and the mode for factors. The way the function works with factors is a bit different. The function identifies the two most different levels of the factor and calculates the change in predictions for a change from the level with the smallest prediction to the level with the largest prediction.

Value

A list with the following elements:

- `difs`: A matrix of calculated first differences
- `minmax`: A matrix of values that were used to calculate the predicted changes
- `minPred`: A matrix of predicted probabilities when each variable is held at its minimum value, in turn.
- `maxPred`: A matrix of predicted probabilities when each variable is held at its maximum value, in turn.

Author(s)

Dave Armstrong

Examples

```r
library(nnet)
data(france)
mnl.mod <- multinom(vote ~ age + male + retnat + lrself, data=france)
typical.france <- data.frame(
age = 35,
retnat = factor(1, levels=1:3, labels=levels(france$retnat)),
stringsAsFactors=TRUE)
mnlChange(mnl.mod, data=france, typical.dat=typical.france)
```

Description

Calculates average effects of a variable in multinomial logistic regression holding all other variables at observed values.
Usage

mnlChange2(obj, varnames, data, diffchange = c("unit", "sd"), n = 1, R = 1500)

Arguments

- **obj**: An object of class `multinom`
- **varnames**: A string identifying the variable to be manipulated.
- **data**: Data frame used to fit `object`.
- **diffchange**: A string indicating the difference in predictor values to calculate the discrete change. `sd` gives plus and minus one-half standard deviation change around the median and `unit` gives a plus and minus one-half unit change around the median.
- **n**: Number of `diffchange` units to change.
- **R**: Number of simulations.

Value

A list with elements:

- **mean**: Average effect of the variable for each category of the dependent variable.
- **lower**: Lower 95 percent confidence bound
- **upper**: Upper 95 percent confidence bound

Author(s)

Dave Armstrong

Examples

```r
library(nnet)
data(france)
mnl.mod <- multinom(vote ~ age + male + retnat + lrself, data=france)
mnlChange2(mnl.mod, "lrself", data=france, )
```
**mnlfit**

*Fit Statistics and Specification Test for Multinomial Logistic Regression*

**Description**

Provides fit statistics (pseudo R-squared values) and the Fagerland, Hosmer and Bonfi (2008) specification test for Multinomial Logistic Regression models.

**Usage**

```r
mnlfit(obj, permute = FALSE)
```

**Arguments**

- `obj` An object of class `multinom`
- `permute` Logical indicating whether to check all base categories for the Fagerland et. al. specification test.

**Value**

A list with elements:

- `result` Fit statistics.
- `permres` The results of the base category permutation exercise.

**Author(s)**

Dave Armstrong

**References**


**Examples**

```r
library(nnet)
data(france)
mnl.mod <- multinom(vote ~ age + male + retnat + lrself, data=france)
mnlfit(mnl.mod)
```
Description

By default, the summary for objects of class `multinom` is not particularly helpful. It still requires a lot of work on the part of the user to figure out which coefficients are significantly different from zero and which ones are not. `mnlSig` solves this problem by either flagging significant coefficients with an asterisk or only printing significant coefficients, leaving insignificant ones blank.

Usage

```r
mnlSig(
  obj,
  pval = 0.05,
  two.sided = TRUE,
  flag.sig = TRUE,
  insig.blank = FALSE
)
```

Arguments

- `obj`: A model object of class `multinom`.
- `pval`: The desired Type I error rate to identify coefficients as statistically significant.
- `two.sided`: Logical indicating whether calculated p-values should be two-sided (if `TRUE`) or one-sided (if `FALSE`).
- `flag.sig`: Logical indicating whether an asterisk should be placed beside coefficients which are significant at the `pval` level.
- `insig.blank`: Logical indicating whether coefficients which are not significant at the `pval` level should be blank in the output.

Value

A data frame suitable for printing with the (optionally significance-flagged) coefficients from a multinomial logit model.

Author(s)

Dave Armstrong

Examples

```r
library(nnet)
data(france)
mnl.mod <- multinom(vote ~ retnat + male + retnat + lrself, data=france)
mnlSig(mnl.mod)
```
NKnots

**AIC and BIC selection of number of spline knots**

**Description**

Calculates AIC and BIC for the selection of knots in a spline over values (potentially including polynomials) up to a user-defined maximum.

**Usage**

```r
NKnots(
  form,
  var,
  data,
  degree = 3,
  min.knots = 1,
  max.knots = 10,
  includePoly = FALSE,
  plot = FALSE,
  criterion = c("AIC", "BIC", "CV"),
  cvk = 10,
  cviter = 10
)
```

**Arguments**

- **form**: A formula detailing the model for which smoothing is to be evaluated.
- **var**: A character string identifying the variable for which smoothing is to be evaluated.
- **data**: Data frame providing values of all variables in `form`.
- **degree**: Degree of polynomial in B-spline basis functions.
- **min.knots**: Minimum number of internal B-spline knots to be evaluated.
- **max.knots**: Maximum number of internal B-spline knots to be evaluated.
- **includePoly**: Include linear and polynomial models up to, and including `degree`-th order polynomials.
- **plot**: Logical indicating whether a plot should be returned.
- **criterion**: Statistical criterion to minimize in order to find the best number of knots - AIC, BIC or Cross-validation.
- **cvk**: Number of groups for cross-validation.
- **cviter**: Number of iterations of cross-validation to average over. 10 is the default but in real-world applications, this should be somewhere around 200.
Value

A plot, if plot=TRUE, otherwise a data frame with the degrees of freedom and corresponding fit measure.

Author(s)

Dave Armstrong

Examples

data(Prestige, package="carData")
NKnots(prestige ~ education + type, var="income", data=na.omit(Prestige), plot=FALSE)

Description

Estimate hypothesis test of lower- and higher-order non-linear relationships against an assumed target relationship.

Usage

NKnotsTest(
form,
var,
data,
targetdf = 1,
degree = 3,
min.knots = 1,
max.knots = 10,
adjust = "none"
)

Arguments

form A formula detailing the model for which smoothing is to be evaluated.
var A character string identifying the variable for which smoothing is to be evaluated.
data Data frame providing values of all variables in form.
targetdf The assumed degrees of freedom against which the tests will be conducted.
degree Degree of polynomial in B-spline basis functions.
min.knots Minimum number of internal B-spline knots to be evaluated.
max.knots Maximum number of internal B-spline knots to be evaluated.
adjust Method by which p-values will be adjusted (see p.adjust)
oc2plot

Plot First Differences from Ordinal DV Model

Description
Takes the output from ordChange and turns it into a plot.

Usage
oc2plot(ordc, plot = TRUE)

Arguments
ordc The output from ordChange.
plot Logical indicating whether a plot (if TRUE) or data (if FALSE) should be returned.

Value
Either a lattice plot or a data.frame depending on the specification of the plot argument.
Author(s)

Dave Armstrong

Examples

```r
library(MASS)
data(france)
polr.mod <- polr(vote ~ age + male + retnat + lrself, data=france)
typical.france <- data.frame(
  age = 35,
  retnat = factor(1, levels=1:3, labels=levels(france$retnat)),
  stringsAsFactors=TRUE)
oc.res <- ordChange(polr.mod, data=france, typical.dat=typical.france, sim=TRUE)
oc2plot(oc.res)
```

Description

For objects of class `polr` the function plots the average effect of a single variable holding all other variables at their observed values.

Usage

```r
ordAveEffPlot(
  obj, 
  varname, 
  data, 
  R = 1500, 
  nvals = 25, 
  plot = TRUE, 
  returnInd = FALSE, 
  returnMprob = FALSE, 
  ...
)
```

Arguments

- **obj**: An object of class `polr`
- **varname**: A string providing the name of the variable for which you want the plot to be drawn.
- **data**: Data used to estimate `obj`.
- **R**: Number of simulations to generate confidence intervals.
- **nvals**: Number of evaluation points of the function
ordAveEffPlot

plot Logical indicating whether or not the result should be plotted (if TRUE) or returned to the console (if FALSE).

returnInd Logical indicating whether average individual probabilities should be returned.

returnMprob Logical indicating whether marginal probabilities, averaged over individuals, should be returned.

... Arguments passed down to the call to xyplot

Details

Following the advice of Hanmer and Kalkan (2013) the function calculates the average effect of a variable holding all other variables at observed values and then plots the result.

Value

Either a plot or a list with a data frame containing the variables

mean The average effect (i.e., predicted probability)
lower The lower 95% confidence bound
upper The upper 95% confidence bound
y The values of the dependent variable being predicted
x The values of the independent variable being manipulated

and the elements Ind or Mprob, as requested.

Author(s)

Dave Armstrong

References


Examples

library(MASS)
data(france)
polr.mod <- polr(vote ~ age + male + retnat + lrself, data=france)
## Not run: ordAveEffPlot(polr.mod, "lrself", data=france)
Description

For objects of class `polr`, it calculates the change in predicted probabilities, for maximal discrete changes in all covariates holding all other variables constant at typical values.

Usage

```r
ordChange(
  obj,
  data,
  typical.dat = NULL,
  diffchange = c("range", "sd", "unit"),
  n = 1,
  sim = TRUE,
  R = 1500
)
```

Arguments

- `obj`: A model object of class `polr`.
- `data`: Data frame used to fit `obj`.
- `typical.dat`: Data frame with a single row containing values at which to hold variables constant when calculating first differences. These values will be passed to `predict`, so factors must take on a single value, but have all possible levels as their levels attribute.
- `diffchange`: A string indicating the difference in predictor values to calculate the discrete change. `range` gives the difference between the minimum and maximum, `sd` gives plus and minus one-half standard deviation change around the median and `unit` gives a plus and minus one-half unit change around the median.
- `n`: Number of `diffchange` units to change.
- `sim`: Logical indicating whether or not simulations should be done to generate confidence intervals for the difference.
- `R`: Number of simulations.

Details

The function calculates the changes in predicted probabilities for maximal discrete changes in the covariates for objects of class `polr`. This function works with polynomials specified with the `poly` function. It also works with multiplicative interactions of the covariates by virtue of the fact that it holds all other variables at typical values. By default, typical values are the median for quantitative variables and the mode for factors. The way the function works with factors is a bit different. The function identifies the two most different levels of the factor and calculates the change in predictions for a change from the level with the smallest prediction to the level with the largest prediction.
Value

A list with the following elements:

- **diffs**: A matrix of calculated first differences
- **minmax**: A matrix of values that were used to calculate the predicted changes
- **minPred**: A matrix of predicted probabilities when each variable is held at its minimum value, in turn.
- **maxPred**: A matrix of predicted probabilities when each variable is held at its maximum value, in turn.

Author(s)

Dave Armstrong

Examples

```r
library(MASS)
data(france)
polr.mod <- polr(vote ~ age + male + retnat + lrself, data=france)
typical.france <- data.frame(
age = 35,
  retnat = factor(1, levels=1:3, labels=levels(france$retnat)),
  stringsAsFactors=TRUE)
ordChange(polr.mod, data=france, typical.dat=typical.france, sim=FALSE)
```

Description

For objects of class `polr`, it calculates the average change in predicted probabilities, for discrete changes in a covariate holding all other variables at their observed values.

Usage

```r
ordChange2(obj, varnames, data, diffchange = c("sd", "unit"), n = 1, R = 1500)
```

Arguments

- **obj**: A model object of class `polr`.
- **varnames**: A vector of strings identifying the variable to be manipulated.
- **data**: Data frame used to fit `obj`.
ordChange2

- **diffchange**: A string indicating the difference in predictor values to calculate the discrete change. `sd` gives plus and minus one-half standard deviation change around the median and `unit` gives a plus and minus one-half unit change around the median.

- **n**: Number of `diffchange` units to change.

- **R**: Number of simulations.

**Details**

The function calculates the changes in predicted probabilities for maximal discrete changes in the covariates for objects of class `polr`. This function works with polynomials specified with the `polym` function. It also works with multiplicative interactions of the covariates by virtue of the fact that it holds all other variables at typical values. By default, typical values are the median for quantitative variables and the mode for factors. The way the function works with factors is a bit different. The function identifies the two most different levels of the factor and calculates the change in predictions for a change from the level with the smallest prediction to the level with the largest prediction.

**Value**

A list with the following elements:

- **diffs**: A matrix of calculated first differences
- **minmax**: A matrix of values that were used to calculate the predicted changes
- **minPred**: A matrix of predicted probabilities when each variable is held at its minimum value, in turn.
- **maxPred**: A matrix of predicted probabilities when each variable is held at its maximum value, in turn.

**Author(s)**

Dave Armstrong

**Examples**

```r
library(MASS)
data(france)
polr.mod <- polr(vote ~ age + male + retnat + lrself, data=france)
typical.france <- data.frame(
age = 35,
retnat = factor(1, levels=1:3, labels=levels(france$retnat)),
stringsAsFactors=TRUE)
ordChange2(polr.mod, "age", data=france, diffchange="sd")
```
**ordfit**

---

**Fit Statistics for Proportional Odds Logistic Regression Models**

**Description**

For objects of class `polr`, it calculates a number of fit statistics and specification tests.

**Usage**

`ordfit(obj)`

**Arguments**

- `obj` A model object of class `polr`.

**Value**

An object of class `ordfit` which is a matrix containing statistics and specification tests.

**Author(s)**

Dave Armstrong

**References**


**Examples**

```r
library(MASS)
data(france)
polr.mod <- polr(vote ~ age + male + retnat + lrself, data=france)
ordfit(polr.mod)
```
outEff

Plot Effects of Removing Outliers

Description

Plots the effect of a variable sequentially removing outlying observations.

Usage

outEff(
  obj, 
  var, 
  data, 
  stat = c("cooksD", "hat", "deviance", "pearson"), 
  nOut = 10, 
  whichOut = NULL, 
  cumulative = FALSE 
)

Arguments

obj An object of class glm that will be used to plot the outlier removed lines.
var A character string giving the name of the variable to be used.
data A data frame.
stat Which statistic to use to evaluate ‘outlyingness’.
nOut Number of outliers to be removed.
whichOut If not NULL, a vector of observation numbers to be removed manually, rather than using stat.
cumulative Logical indicating whether the outliers should be removed cumulatively, or each one in turn, replacing the other outlying observations.

Author(s)

Dave Armstrong

outXT

Create LaTeX or CSV versions of an Object Produced by CrossTable

Description

outXT takes the output from CrossTable in the gmodels package and produces either LaTeX code or CSV file that can be imported into word processing software.
outXT

Usage

outXT(
  obj,
  count = TRUE,
  prop.r = TRUE,
  prop.c = TRUE,
  prop.t = TRUE,
  col.marg = TRUE,
  row.marg = TRUE,
  digits = 3,
  type = "word",
  file = NULL
)

Arguments

obj A list returned by CrossTable from the gmodels package.

count Logical indicating whether the cell frequencies should be returned.

prop.r Logical indicating whether the row proportions should be returned.

prop.c Logical indicating whether the column proportions should be returned.

prop.t Logical indicating whether the cell proportions should be returned.

col.marg Logical indicating whether the column marginals should be printed.

row.marg Logical indicating whether the row marginals should be printed.

digits Number of digits to use in printing the proportions.

type String where word indicates a CSV file will be produced and \texttt{latex} indicates \LaTeX\ code will be generated.

file Connection where the file will be written, if \texttt{NULL} the output will only be written to the console

Value

A file containing \LaTeX\ Code or CSV data to make a table

Author(s)

Dave Armstrong
panel.2cat  

Lattice panel function for confidence intervals with capped bars

Description
This panel function is defined to plot confidence intervals in a multi-panel lattice display where the 
x-variable is categorical. Note, both lower and upper must be passed directly to `xyplot` as they will 
be passed down to the panel function.

Usage

```r
panel.2cat(x, y, subscripts, lower, upper, length = 0.2)
```

Arguments

- `x, y`: Data from the call to `xyplot`.
- `subscripts`: Variable used to created the juxtaposed panels.
- `lower, upper`: 95% lower and upper bounds of `y`.
- `length`: Length of the arrow head lines.

Author(s)

Dave Armstrong

Examples

```r
library(lattice)
library(effects)
data(Duncan, package="carData")
Duncan$inc.cat <- cut(Duncan$income, 3)
mod <- lm(prestige ~ inc.cat * type + education,
data=Duncan)
e1 <- effect("inc.cat*type", mod)
update(plot(e1), panel=panel.2cat)
```

panel.ci  

Lattice panel function for confidence intervals

Description
This panel function is defined to plot confidence intervals in a multi-panel lattice display. Note, 
both lower and upper must be passed directly to `xyplot` as they will be passed down to the prepanel 
function.
Usage

panel.ci(x, y, subscripts, lower, upper, zl)

Arguments

x, y Data from the call to xyplot.
subscripts Variable used to create the juxtaposed panels.
lower, upper 95% lower and upper bounds of y.
zl Logical indicating whether or not a horizontal dotted line at zero is desired.

Author(s)

Dave Armstrong

panel.doublerug Lattice panel function for two rug plots

Description

This panel function is defined to plot two rugs, one on top of the other in a multi-panel lattice display.

Usage

panel.doublerug(
  xa = NULL,
  xb = NULL,
  regular = TRUE,
  start = if (regular) 0 else 0.97,
  end = if (regular) 0.03 else 1,
  x.units = rep("npc", 2),
  lty = 1,
  lwd = 1
)

Arguments

xa, xb Numeric vectors to be plotted.
regular Logical flag indicating whether rug is to be drawn on the usual side (bottom/left) as opposed to the other side (top/right).
start, end Start and end points for the rug ticks on the y-axis.
x.units Character vectors, replicated to be of length two. Specifies the (grid) units associated with start and end above. x.units are for the rug on the x-axis and y-axis respectively (and thus are associated with start and end values on the y and x scales respectively). See panel.rug for more details.
lty, lwd Line type and width arguments (see par for more details).
Author(s)

Dave Armstrong

---

**panel.transci**

*Lattice panel function for translucent confidence intervals*

**Description**

This panel function is defined to plot translucent confidence intervals in a single-panel, grouped (i.e., superposed) lattice display. Note, both lower and upper must be passed directly to `xyplot` as they will be passed down to the panel function.

**Usage**

```r
panel.transci(x, y, groups, lower, upper, ca = 0.25, ...)
```

**Arguments**

- `x, y` Data from the call to `xyplot`.
- `groups` Variable used to created the superposed panels.
- `lower, upper` 95% lower and upper bounds of `y`.
- `ca` Value of the alpha channel in [0,1]
- `...` Other arguments to be passed down to the plotting functions.

Author(s)

Dave Armstrong

---

**pgumbel**

*PDF of the Gumbel Distribution*

**Description**

Returns the PDF of the Gumbel distribution.

**Usage**

```r
pgumbel(q, location = 0, scale = 1, lower.tail = TRUE, log.p = FALSE)
```
Arguments

q    Vector of quantiles
location The location parameter, \( \mu \). This is not the mean of the Gumbel distribution.
scale The scale parameter \( \sigma \).
lower.tail Logical, whether lower, if TRUE or upper, if FALSE, tail probabilities should be returned.
log.p Logical, if TRUE probabilities are given in their log.

Details

This code and the details of the help file were taken from the VGAM package.

The Gumbel distribution is a special case of the generalized extreme value (GEV) distribution where the shape parameter \( \xi = 0 \). The latter has 3 parameters, so the Gumbel distribution has two. The Gumbel distribution function is

\[
G(y) = \exp \left( - \exp \left( - \frac{y - \mu}{\sigma} \right) \right)
\]

where \(-\infty < y < \infty\), \(-\infty < \mu < \infty\) and \(\sigma > 0\). Its mean is

\[
\mu - \sigma \gamma
\]

and its variance is

\[
\sigma^2 \pi^2 / 6
\]

where \(\gamma\) is Euler’s constant (which can be obtained as \(-\text{digamma}(1)\)).

Value

A vector of probabilities

Author(s)

Thomas Yee

plot.alsos

Plotting method for ALSOS object

Description

Makes a plot or optionally returns data for user-generated plots from an alsos object

Usage

```r
## S3 method for class 'alsos'
plot(x, which_var = NULL, return_data = FALSE, ...)
```
Arguments

x An object of class alsos.

which_var The name of a raw variable that was scaled in the alsos procedure for which a measurement function should be returned.

return_data Logical indicating whether the data should be returned (if TRUE) or a plot generated (if FALSE)

... arguments to be passed in, currently not implemented.

Value

A plot.

plot.balsos  
Plot Results from BALSOS

Description

Plots the optimally scaled points with posterior 95% credible intervals.

Usage

```r
## S3 method for class 'balsos'
plot(x, ..., freq = TRUE, offset = 0.1, plot = TRUE)
```

Arguments

x Object of class balsos.

... Other arguments to be passed down, currently not implement and may conflict with the lattice figure. To change the figure the best advice would be to save the plot as an object and use the update function to change its features.

freq Logical indicating whether you want the frequentist result plotted alongside the Bayesian result.

offset If freq=T, the Bayesian points will be plotted at x-offset and the frequentist points will be plotted at x+offset.

plot Logical indicating whether the plot should be returned or just the data.

Value

A lattice graph produce by a call to `xyplot`.

Author(s)

Dave Armstrong
**plot.loess**  

*Plot LOESS curve.*

---

**Description**

Plots the loess curve of the fitted values against a focal x-variable.

**Usage**

```r
## S3 method for class 'loess'
plot(
  x, 
  ..., 
  ci = TRUE, 
  level = 0.95, 
  linear = FALSE, 
  addPoints = FALSE, 
  col.alpha = 0.5
)
```

**Arguments**

- `x`  
  An object of class `loess`.

- `...`  
  Other arguments to be passed down to `xyplot`.

- `ci`  
  Logical indicating whether point-wise confidence intervals should be included around the fitted curve.

- `level`  
  The confidence level of the confidence intervals.

- `linear`  
  Logical indicating whether the OLS line should also be included.

- `addPoints`  
  Logical indicating whether or not points should be added to the figure identifying the position of individual observations.

- `col.alpha`  
  Value for alpha channel of the RGB color palette.

**Details**

Plots the fitted loess curve potentially with point-wise confidence bounds.

**Value**

A plot.

**Author(s)**

Dave Armstrong
plot.secdiff  
*Plotting Method for Second Difference Objects*

**Description**
Plots the results of the `secdiff` function.

**Usage**
```r
## S3 method for class 'secdiff'
plot(x, level = 0.95, ...)
```

**Arguments**
- `x` An object of class `secdiff`
- `level` The confidence level of the confidence interval(s)
- `...` Other arguments to be passed down, currently not implemented.

poisfit  
*Scalar Measures of Fit for Poisson GLMs Models*

**Description**
Calculates scalar measures of fit for models with count dependent variables along the lines described in Long (1997) and Long and Freese (2005).

**Usage**
```r
poisfit(obj)
```

**Arguments**
- `obj` A model of class `glm` with `family=poisson`.

**Details**
poisfit calculates scalar measures of fit (many of which are pseudo-R-squared measures) to describe how well a model fits data with a count dependent variable.

**Value**
A named vector of scalar measures of fit

**Author(s)**
Dave Armstrong
Deviance and Chi-squared Goodness-of-Fit Test for Poisson Models

Description

Deviance and Chi-squared goodness-of-fit test of the null hypothesis that poisson variance is appropriate to model the conditional dispersion of the data, given a particular model.

Usage

poisGOF(obj)

Arguments

obj

A model object of class glm (with family=poisson).

Value

A 2x2 data frame with rows representing the different types of statistics (Deviance and Chi-squared) and columns representing the test statistic and p-value.

Author(s)

Dave Armstrong

References


Examples

```r
# Example taken from MASS help file for glm, identified to be
# Dobson (1990) Page 93: Randomized Controlled Trial :
counts <- c(18,17,15,20,10,20,25,13,12)
outcome <- gl(3,1,9)
treatment <- gl(3,3)
print(d.AD <- data.frame(treatment, outcome, counts, stringsAsFactors=TRUE))
glm.D93 <- glm(counts ~ outcome + treatment, family=poisson())
poisGOF(glm.D93)
```
**powerTrans**  
*Power Transformation Function*

**Description**

Power transformation function that treats everything with absolute power transform < .01 as the log transform.

**Usage**

```
powerTrans(x, transPower)
```

**Arguments**

- `x` Vector of values to be transformed
- `transPower` The power of the transformation

**Value**

A vector of transformed values

---

**pre**  
*Proportional and Expected Proportional Reductions in Error*

**Description**

Calculates proportional reduction in error (PRE) and expected proportional reduction in error (ePRE) from Herron (1999).

**Usage**

```
pre(mod1, mod2 = NULL, sim = FALSE, R = 2500)
```

**Arguments**

- `mod1` A model of class `glm` (with family `binomial`), `polr` or `multinom` for which (e)PRE will be calculated.
- `mod2` A model of the same class as `mod1` against which proportional reduction in error will be measured. If NULL, the null model will be used.
- `sim` A logical argument indicating whether a parametric bootstrap should be used to calculate confidence bounds for (e)PRE. See Details for more information.
- `R` Number of bootstrap samples to be drawn if `sim=TRUE`. 
Details

Proportional reduction in error is calculated as a function of correct and incorrect predictions (and the probabilities of correct and incorrect predictions for ePRE). When sim=TRUE, a parametric bootstrap will be used that draws from the multivariate normal distribution centered at the coefficient estimates from the model and using the estimated variance-covariance matrix of the estimators as Sigma. This matrix is used to form R versions of XB and predictions are made for each of the R different versions of XB. Confidence intervals can then be created from the bootstrap sampled (e)PRE values.

Value

An object of class pre, which is a list with the following elements:

- pre: The proportional reduction in error
- epre: The expected proportional reduction in error
- m1form: The formula for model 1
- m2form: The formula for model 2
- pcp: The percent correctly predicted by model 1
- pmc: The percent correctly predicted by model 2
- epcp: The expected percent correctly predicted by model 1
- epmc: The expected percent correctly predicted by model 2
- pre.sim: A vector of bootstrapped PRE values if sim=TRUE
- epre.sim: A vector of bootstrapped ePRE values if sim=TRUE

Author(s)

Dave Armstrong

References


Examples

data(france)
left.mod <- glm(voteleft ~ male + age + retnat + poly(lrself, 2), data=france, family=binomial)
pre(left.mod)
prepanel.ci  \hspace{1em} \textit{Lattice prepanel function for confidence intervals}

**Description**

This prepanel function is defined so as to allow room for all confidence intervals plotted in a lattice display. Note, both lower and upper must be passed directly to `xyplot` as they will be passed down to the prepanel function.

**Usage**

```r
prepanel.ci(x, y, subscripts, lower, upper)
```

**Arguments**

- `x, y` Data from the call to `xyplot`.
- `subscripts` Variable used to created the juxtaposed panels.
- `lower, upper` 95\% lower and upper bounds of `y`.

**Value**

A list giving the ranges and differences in ranges of `x` and the lower and upper bounds of `y`.

**Author(s)**

Dave Armstrong

---

print.diffci  \hspace{1em} \textit{Print Confidence Intervals for Predicted Probabilities and Their Differences}

**Description**

Print method for output from the `probci` function.

**Usage**

```
## S3 method for class 'diffci'
print(
  x,
  type = c("pr", "pw"),
  ..., 
  digits = 4,
  filter = NULL,
  const = NULL,
  onlySig = FALSE
)
```
Arguments

- **x**: A object of class `diffci` produced by `probci`.
- **type**: Which kind of result to print - predictions (`pr`) or pairwise differences in predictions (`pw`).
- **...**: Other arguments to be passed down to print, currently unimplemented.
- **digits**: How many digits to round output.
- **filter**: A named list of values where the names indicate the variable to be filtered and the values in the vector indicate the values to include for the filtering variable.
- **const**: A string identifying the name of the variable to be held constant across comparisons. Only applies if `type = "pw"`.
- **onlySig**: Logical indicating whether all differences should be displayed or only those significant at the 95% two-tailed level.

Value

An data frame with the following variables:

- **variables**: The variables and the values at which they are held constant. For example, `tmp1` would be the first value of `tmp` used in the probability calculation and `tmp2` would be the second value of `tmp` used in the probability calculation.
- **pred_prob**: The difference in predicted probability given the following change in `X`: `tmp2-tmp1`.
- **lower, upper**: The lower and upper 95% confidence bounds.

Author(s)

Dave Armstrong

Examples

data(france)
left.mod <- glm(voteleft ~ male + age + retnat + poly(lrself, 2, raw=TRUE), data=france, family=binomial)
data(france)
left.mod <- glm(voteleft ~ male + age + retnat + poly(lrself, 2, raw=TRUE), data=france, family=binomial)
out <- probci(left.mod, france, numQuantVals=3, changeX=c("retnat", "lrself"), calcPW = TRUE)
print(out, filter=list(retnat=c("Better", "Worse")))
print(out, type="pw", filter=list(retnat=c("Better", "Worse")), const="lrself")
**print.glmc2**  
*Print method for glmChange objects*

**Description**

Print method for object of class glmc2.

**Usage**

```r
## S3 method for class 'glmc2'
print(x, ...)  
```

**Arguments**

- `x` Object of class glmc2
- `...` Currently unimplemented.

---

**print.iqq**  
*Print method for intQualQuant objects.*

**Description**

Print method for objects of class iqq calculated with the intQualQuant function.

**Usage**

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

**Arguments**

- `x` Object of class iqq
- `...` Currently unimplemented.
**print.ordChange**

*Print method for ordChange objects*

**Description**

Print methods for objects of class ordChange

**Usage**

```r
## S3 method for class 'ordChange'
print(x, ..., digits = 3)
```

**Arguments**

- `x`: Object of class ordChange
- `...`: Other arguments to be passed down to the function
- `digits`: Number of digits to print

**print.pre**

*Print method for objects of class pre*

**Description**

Prints the output from an object of class pre. The function prints all components of the calculation and optionally simulated confidence bounds.

**Usage**

```r
## S3 method for class 'pre'
print(x, ..., sim.ci = 0.95)
```

**Arguments**

- `x`: An object of class pre.
- `...`: Other arguments passed to print, currently not implemented
- `sim.ci`: Coverage for the simulated confidence interval, if `sim=TRUE` in the call to `pre`.

**Author(s)**

Dave Armstrong

**See Also**

- `pre`
Confidence Intervals for Predicted Probabilities and Their Differences

Description
Calculates predicted probabilities for any combination of x-variable values holding all other variables constant at either typical values (average case approach) or at observed values (average effect approach).

Usage
probci(
  obj,
  data,
  .b = NULL,
  .vcov = NULL,
  changeX = NULL,
  numQuantVals = 5,
  xvals = NULL,
  type = c("aveEff", "aveCase"),
  returnProbs = FALSE,
  calcPW = FALSE
)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>obj</td>
<td>A model of class glm, particularly those with binomial family.</td>
</tr>
<tr>
<td>data</td>
<td>Data frame used to estimate obj.</td>
</tr>
<tr>
<td>.b</td>
<td>A vector of coefficients to be passed down to the simulation. If NULL, coef() will be used to obtain coefficients from obj.</td>
</tr>
<tr>
<td>.vcov</td>
<td>A parameter variance covariance matrix to be passed to the simulation. If NULL, vcov() will be used to obtain the variance-covariance matrix of the parameters.</td>
</tr>
<tr>
<td>changeX</td>
<td>A vector of strings giving the names of variables for which changes are desired.</td>
</tr>
<tr>
<td>numQuantVals</td>
<td>For quantitative variables, if no x-values are specified in xvals, then numQuantVals gives the number of values used across the range of the variable.</td>
</tr>
<tr>
<td>xvals</td>
<td>A named list of values used to make the predictions. The names in the list should correspond with the variable names specified in changeX.</td>
</tr>
<tr>
<td>type</td>
<td>Type of effect to be generated. aveEff produces the average first difference across all observed values of X, while aveCase gives the first difference holding all other variables constant at typical values.</td>
</tr>
<tr>
<td>returnProbs</td>
<td>Whether or not the vector/matrix of predicted probabilities should be returned as well.</td>
</tr>
<tr>
<td>calcPW</td>
<td>Should the pairwise differences be calculated?</td>
</tr>
</tbody>
</table>
**Details**

Calculates predicted probabilities for any combination of x-variable values holding all other variables constant at either typical values (average case approach) or at observed values (average effect approach). The function uses a parametric bootstrap to provide generate confidence bounds for predicted probabilities and their differences. The confidence intervals produced are raw percentile interviews (at the 5% level).

**Value**

An data frame with the following variables:

- **variables**: The variables and the values at which they are held constant. For example, tmp1 would be the first value of tmp used in the probability calculation and tmp2 would be the second value of tmp used in the probability calculation.

- **pred_prob**: The difference in predicted probability given the following change in X: tmp2-tmp1.

- **lower, upper**: The lower and upper 95% confidence bounds.

**Author(s)**

Dave Armstrong

**Examples**

```r
data(france)
left.mod <- glm(voteleft ~ male + age + retnat + poly(lrself, 2, raw=TRUE), data=france, family=binomial)
out <- probci(left.mod, france, changeX="retnat")
out
out2 <- probci(left.mod, france, changeX="lrself",
               xvals = list(lrself = c(1,10)))
out2
out3 <- probci(left.mod, france, changeX=c("lrself", "retnat"),
               xvals = list(lrself = c(1,10)))
out3
```

**probgroup**

*Plog Probabilities by Group*

**Description**

Plots predicted probabilities by value of the dependent variable for proportional odds logistic regression and multinomial logistic regression models

**Usage**

`probgroup(obj, ...)`
Arguments

obj Object of class polr or multinom where appropriate.

Details

Plots the predicted probabilities by value of the dependent variable. Each panel only includes the observations that had the identified value of the dependent variable.

Value

A plot.

Author(s)

Dave Armstrong

Description

Prints pairwise correlation matrix flagging statistically significant correlations using one of a few different methods.

Usage

pwCorrMat(
  formula,
  data,
  method = c("z", "t", "sim"),
  weight = NULL,
  alpha = 0.05,
  ...
)

Arguments

formula A right-sided formula giving the variables to be correlated separated by pluses.
data A data frame where the variables in the formula can be found.
method A method for calculating the significance of the of the correlation - one of "z", "t" or "sim". When correlations are calculated with weights, a bootstrap is used to generate p-values regardless of the method specified. See details for more.
weight A vector of weightings as long as there are rows in data.
alpha Cutoff for identifying significant correlations.
... Other arguments to be passed down to sig.cor.
Details

The significance is found through one of three ways. For correlation $r$, the $z$-transformation is $0.5 \times \log((1+r)/(1-r))$, the $p$-value for which is found using the standard normal distribution. The $t$-transformation is $r \times \sqrt{(n-2)/(1-r^2)}$, the $p$-value for which is found using a $t$-distribution with $n-2$ degrees of freedom. The "sim" method uses a permutation test to build the sampling distribution of the correlation under the null hypothesis and then calculates a $p$-value from that distribution.

Value

An object of class `pwc`, which is a list with elements `rSig` which is a lower-triangular correlation matrix where only significant correlations are printed, `r` which is the raw-data pairwise correlation matrix and `p` which gives the $p$-values of all of the correlations.

scaleDataFrame

Standardize quantitative variables in a data frame

Description

This function standardizes quantitative variables in a data frame while leaving the others untouched. This leaves not only factors, but also binary variables (those with values 0, 1, or NA).

Usage

```
scaleDataFrame(data, numsd = 1, nvals_fac = 11, exclude = NULL)
```

Arguments

- `data`: A data frame.
- `numsd`: Number of standard deviations to divide by - defaults to 1.
- `nvals_fac`: Number of unique values required to standardize - variables with fewer than `nvals_fac` unique values will not be standardized.
- `exclude`: A character vector of names of variables to exclude from the standardization.

Value

A data frame with standardized quantitative variables

Author(s)

Dave Armstrong
searchVarLabels  

**Search Variable Labels Attribute**

**Description**

Data imported from SPSS or Stata comes with the variable labels set (if they were set in the original dataset) as one of the dataframe’s attributes. This allows you to search the variable labels and returns the variable column number, name and label for all variables that have partially match the search term either in their labels or names.

**Usage**

searchVarLabels(dat, str)

**Arguments**

- **dat**
  
a data frame whose variable labels you want to search.

- **str**
  
string used to search variable labels.

**Details**

For an imported Stata dataset, variable labels are in the `var.labels` attribute of the dataset and in an SPSS dataset, they are in the `variable.labels` attribute. These are searched, ignoring case, for the desired string.

**Value**

A matrix of dimensions n-matches x 2 is returned, where the first column is the column number of the matching variable and the second column is the variable label. The row names of the matrix are the variable names.

**Author(s)**

Dave Armstrong

---

**secondDiff**  

*Calculate Cross-Derivative and its Variability*

**Description**

Calculates the cross-derivative required to evaluate interactions in logistic/probit regression models.
Usage

secondDiff(
  obj,
  vars,
  data,
  method = c("AME", "MER"),
  vals = NULL,
  typical = NULL
)

Arguments

obj  An object of class glm that will be used to find the cross-derivative.
vars A vector of two variables to be used in calculating the derivative.
data A data frame.
method Indicate whether you want to use average marginal effects (AME) or marginal
effects at representative values (MER).
vals A named list of length 2 where each element gives the minimum and maximum
values used in the calculation.
typical A named vector of values at which to hold variables constant.

details

The function calculates the second difference as (Pr(Y=1|x1=max, x2=max) - Pr(Y=1|x1=min,
  x2=max)) - (Pr(Y=1|x1=max, x2=min) - Pr(Y=1|x1=min, x2=min)). The function uses a parametric
 bootstrap to calculate the sampling distribution of the second difference.

Value

A list with two elements:

ave The average second difference in each iteration of the bootstrap.
ind If type == 'AME', ind is returned with the second difference and measures of
uncertainty for each individual observation in the original dataset
probs If type == 'MER', probs is returned with the full matrix of simulated predicted
probabilities for the four conditions.

Author(s)

Dave Armstrong
**simPredpolr**  
*Calculate Predictions for Proportional Odds Logistic Regression*

**Description**
Calculates predicted probabilities from models of class `polr` from a model object and a vector of coefficient values. This is an auxiliary function used in `pre` if `sim=TRUE`.

**Usage**

```r
simPredpolr(object, coefs, n.coef)
```

**Arguments**
- `object`  
  An object of class `polr`.
- `coefs`  
  A vector of coefficients where elements 1 to `n.coef` give model coefficients and elements `n.coef+1` to `k` have intercepts.
- `n.coef`  
  Number of coefficients (minus intercepts) for the `polr` model.

**Value**
An `n x m`-category matrix of predicted probabilities

**Author(s)**
Dave Armstrong

---

**summary.balsos**  
*Summary method for Bayesian ALSOS*

**Description**
summary method for objects of class `balsos`

**Usage**

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

**Arguments**
- `object`  
  Object of class `balsos`
- `...`  
  Other arguments, currently unimplemented
**summary.secdiff**  

**Summary for Second Difference Objects**

**Description**  
Summary method for objects of class `secdiff`.

**Usage**  
```r  
## S3 method for class 'secdiff'  
summary(object, ..., level = 0.95, digits = 3)  
```

**Arguments**  
- `object`: An object of class `secdiff`
- `...`: Other arguments to be passed down to `summary`.
- `level`: Confidence level for the confidence intervals
- `digits`: Number of digits to print

**sumStats**  

**Summary Statistics**

**Description**  
Provides summary statistics (mean, sd, quartiles, IQR, missing n, valid n) for the variables in a data frame.

**Usage**  
```r  
sumStats(data, vars, byvar = NULL, convertFactors = TRUE)  
```

**Arguments**  
- `data`: A data frame from which variables will be extracted.
- `vars`: A character vector of variable names.
- `byvar`: A character string giving a variable name of a stratifying variable. The summaries of the `vars` will be provided for each level of `byvar`.
- `convertFactors`: Logical indicating whether factors should be converted to numeric first and then summarised.

**Value**  
a vector of summary statistics for each variable or variable-group combination.
test.balsos  

*Testing Measurement Level Assumptions.*

**Description**

Uses the algorithm discussed in Armstrong and Jacoby (2018) to test the intervality of a variable scaled by the Bayesian ALSOS method.

**Usage**

```r
test.balsos(obj, cor.type = c("pearson", "spearman"))
```

**Arguments**

- `obj`: An object of class `balsos`.
- `cor.type`: Type of correlation to be used in the p-value calculation.

**Value**

Printed output giving the Bayesian p-value evaluating the null that there is no interesting difference between the original values and the optimally scaled values.

**Author(s)**

Dave Armstrong

---

testGAMint  

*Simulated F-test for Linear Interactions*

**Description**

Simulates the sampling distribution of the F statistic when comparing a linear interaction model to a generalized additive model with a smooth over the two variables in the interaction.

**Usage**

```r
testGAMint(m1, m2, data, R = 1000, ranCoef = FALSE)
```
Arguments

m1  An object of class gam estimated with the mgcv package. This model should be linear in the interaction of the two x-variables of interest.

m2  An object of class gam estimated with the mgcv package. This model should contain a smooth interaction. For two continuous variables, this should be done with te() unless the variables are measured in the same units (e.g., spatial coordinates) in which case the usual thin-plate regression spline will work. For categorical moderators, you should use the s(x, by=D0) and s(x, by=D) (for a dummy variable moderator, D, where D0=1 when D=0. Remember to include D as a parametric term in the model as well to account for the intercept difference between the two smooth terms.)

data  Data frame used to estimate both models

R  Number of simulated F values to create.

ranCoef  Logical indicating whether the coefficients should be treated as fixed or whether they should be drawn from their implied sampling distribution for each iteration of the simulation.

Details

In simple simulations, an F-test of a linear interaction relative to a smooth interaction with a GAM using a nominal .05 type I error rate, has an actual type I error rate of more than double the nominal rate (this tended to be in the low teens). This function tries to build the F-distribution using simulation. First, it uses the coefficients from the linear interaction model, multiplies them by the coefficients from the linear interaction model and for each iteration of the simulation, it creates the simulated dependent variable by adding a random error to the linear predictor with the same standard deviation as the residual standard deviation from the linear interaction model. All of that is to say that this model has all of the same features as the linear interaction model, except that we are certain that this is the right model. The algorithm then estimates both the linear interaction model and the GAM with a smooth interaction on the original X variables and the new simulated y variable. The F-test is performed and the F-statistic saved for each iteration. The algorithm then calculates the probability of being to the right of the observed F-statistic in the simulated F-distribution.

Value

obsF  The observed F-statistic from the test on the original models.

Fdist  The R different F-statistics calculated at each iteration of the simulation.

Author(s)

Dave Armstrong
**testLoess**

*Significance Test for Loess vs. LM*

**Description**

Calculates an F test to evaluate significant differences between a LOESS model and a parametric alternative estimated with `lm`.

**Usage**

```
testLoess(lmobj, loessobj, alpha = 0.05)
```

**Arguments**

- **lmobj**: An object of class `lm`.
- **loessobj**: An object of class `loess`.
- **alpha**: Desired Type I error rate of test.

**Value**

Printed output describing the results of the test.

**Author(s)**

Dave Armstrong

**Examples**

```r
data(Prestige, package="carData")
linmod <- lm(prestige ~ income, data=Prestige)
lomod <- loess(prestige ~ income, data=Prestige)
testLoess(linmod, lomod)
```

---

**testNL**

*Test Transformations and Polynomials in Non-linear Models*

**Description**

Tests for model improvements for non-linear transformations and polynomials with Clarke’s (2007) distribution-free test for non-nested models.
Usage

testNL(obj, var, transPower, polyOrder, plot = FALSE, ...)

## S3 method for class 'glm'
testNL(obj, var, transPower, polyOrder, plot = FALSE, ...)

## S3 method for class 'lm'
testNL(obj, var, transPower, polyOrder, plot = FALSE, ...)

Arguments

obj Object of a supported class in which non-linear functional forms will be tested.

var String giving name of variable to be tested.

transPower The power used in the transformation. For transformations in the range (-0.01, 0.01), the log transformation is used.

polyOrder The order of the polynomial to be used.

plot Logical indicating whether the effects should be plotted

... Currently not implemented.

Details

Three hypotheses are tested with this function. The first is whether the original specification is preferred to the power transformation. The second is whether the original specification is preferred to the polynomial model. The third is whether the power transformation is preferred to the polynomial model. All tests are done with the Clarke test.

Value

A plot or a data frame giving the results of the tests identified above.

Author(s)

Dave Armstrong

References

tidy_boot_ci  
*Tidy Bootstrap Confidence Intervals*

**Description**

Returns a tibble with confidence intervals for all parameters from a bootstrapping object estimated with the `boot()` function.

**Usage**

```r
 tidy_boot_ci(
    obj,
    indices = NULL,
    type = c("norm", "basic", "stud", "perc", "bca"),
    conf = 0.95,
    term_names = NULL,
    ...
)
```

**Arguments**

- `obj`: An object of class `boot`.
- `indices`: The column numbers of `obj$t` to be used in the calculation. If `NULL`, all columns are used.
- `type`: The type of confidence interval to be produced. Unlike `boot.ci()`, "all" is not an option.
- `conf`: The confidence level to be used for the interval.
- `term_names`: The names of the parameters to be used as identifiers in the tibble.
- `...`: Other arguments to be passed down to `boot.ci()`

**Value**

A tibble with the term name, estimate, lower and upper confidence bounds.

---

tscsflag  
*Lag time-series cross-sectional variables*

**Description**

Lags (or leads) a variable in a time-series cross-sectional dataset.

**Usage**

```r
 tscsflag(dat, x, id, time, lagLength = 1)
```
Arguments

- **dat**: A data frame.
- **x**: A string identifying variable to be lagged.
- **id**: A string identifying the name of the cross-sectional identifier.
- **time**: A string identifying the name of the time variable.
- **lagLength**: The length of the lag, use negative values for leading variables.

Value

A vector giving the lagged values of x.

Author(s)

Dave Armstrong

tTest

**t-Test function**

Description

This function is a wrapper to the t.test function but produces more information in the output.

Usage

tTest(x, y, data, ...)

Arguments

- **x**: The dichotomous variable for the test.
- **y**: The interval/ratio variable for the test.
- **data**: A data frame where x and y can be found.
- **...**: Other arguments to be passed down to t.test.

Value

an object of class tTest
unformulate  Break Apart Model Formula

Description

Works as a sort of inverse to reformulate by breaking apart the formula into response and the term labels. It also returns the variable names of all of the variables implicated in the formula.

Usage

unformulate(form, keep_env = FALSE)

Arguments

form  A formula
keep_env  Logical indicating whether the formula’s environment should be returned with the result

Details

A sort of inverse of the reformulate function.

Value

A list with termlabels giving the rhs terms of the model, response give the lhs of the model, env optionally giving the environment of the formula and vars a vector of the variable names implicated in the formula.

xt  Cross-Tabulation of Weighted or Unweighted Data

Description

Cross-Tabulation of Weighted or Unweighted Data

Usage

xt(data, var, byvar = NULL, controlvar = NULL, weight = NULL, ...)

xt
yeo.johnson

Arguments

data Either a data frame or a survey design object.
var Row variable for the cross-tabular.
byvar Optional column variable for the cross-tabulation. If NULL, a frequency and relative frequency distribution of var will be produced.
controlvar The name of a categorical control variable.
weight A vector of weights to be applied to the table.
... Other arguments to be passed down to make_assoc_stats. You can use this to calculate different statistics. By default, you get Chi-squared, Cramer’s V, Gamma and Kendall’s Tau-b.

Produces a cross-tabulation and Chi-square statistic for weighted or unweighted data.

Value

A list with two elements - table of class tabyl and the returned results from svychisq.

---

yeo.johnson  

Yeo-Johnson Transformation

Description

Computes the normalizing Yeo-Johnson transformation. The code and the details of the help file were taken from the VGAM package.

Usage

yeo.johnson(
  y,
  lambda,
  derivative = 0,
  epsilon = sqrt(.Machine$double.eps),
  inverse = FALSE
)

Arguments

y Numeric, a vector or matrix.
lambda Numeric. It is recycled to the same length as y if necessary.
derivative Non-negative integer. The default is the ordinary function evaluation, otherwise the derivative with respect to lambda.
epsilon Numeric and positive value. The tolerance given to values of lambda when comparing it to 0 or 2.
inverse Logical. Return the inverse transformation?
Details

The Yeo-Johnson transformation can be thought of as an extension of the Box-Cox transformation. It handles both positive and negative values, whereas the Box-Cox transformation only handles positive values. Both can be used to transform the data so as to improve normality. They can be used to perform LMS quantile regression.

Value

A vector of transformed values.

Author(s)

Thomas Yee

---

**Description**

Uses `nlminb` to find the optimal Yeo-Johnson transformation parameters conditional on a parametric model specification.

**Usage**

```
yj_trans(form, data, trans.vars, round.digits = 3, ...)
```

**Arguments**

- `form` A formula with a dependent variable that will be optimally scaled
- `data` A data frame.
- `trans.vars` A character string identifying the variables that should be transformed
- `round.digits` Number of digits to round the transformation parameters.
- `...` Other arguments to be passed down to `lm`.

**Value**

A linear model object that was estimated on the optimally transformed variables.

**Author(s)**

Dave Armstrong
Maximal First Differences for Zero-Inflated Models

Description

Calculates the change in predicted counts or optionally the predicted probability of being in the zero-count group, for maximal discrete changes in all covariates holding all other variables constant at typical values.

Usage

ziChange(obj, data, typical.dat = NULL, type = "count")

Arguments

obj
A model object of class zeroinfl.
data
Data frame used to fit object.
typical.dat
Data frame with a single row containing values at which to hold variables constant when calculating first differences. These values will be passed to predict, so factors must take on a single value, but have all possible levels as their levels attribute.
type
Character string of either ‘count’ (to obtain changes in predicted counts) or ‘zero’ (to obtain changes in the predicted probability of membership in the zero group).

Details

The function calculates the changes in predicted counts, or optionally the predicted probability of being in the zero group, for maximal discrete changes in the covariates. This function works with polynomials specified with the poly function. It also works with multiplicative interactions of the covariates by virtue of the fact that it holds all other variables at typical values. By default, typical values are the median for quantitative variables and the mode for factors. The way the function works with factors is a bit different. The function identifies the two most different levels of the factor and calculates the change in predictions for a change from the level with the smallest prediction to the level with the largest prediction.

Value

A list with the following elements:
diffs
A matrix of calculated first differences
minmax
A matrix of values that were used to calculate the predicted changes

Author(s)

Dave Armstrong
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