# Package 'mplot'

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```
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      (2010, <doi:10.1111/j.1751-5823.2010.00108.x>); Murray, Heritier and Mueller
      (2013, <doi:10.1002/sim.5855>)] as well as the adaptive fence [Jiang et al.
      (2008, <doi:10.1214/07-AOS517>); Jiang et al.
      (2009, <doi:10.1016/j.spl.2008.10.014>)] for linear and generalised linear models.
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2 mplot-package

# **R** topics documented:

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# Description

Graphical model stability and model selection procedures

#### References

Tarr G, Mueller S and Welsh AH (2018). mplot: An R Package for Graphical Model Stability and Variable Selection Procedures. Journal of Statistical Software, 83(9), pp. 1-28. doi: 10.18637/jss.v083.i09

af 3

af

The adaptive fence procedure

# Description

This function implements the adaptive fence procedure to first find the optimal cstar value and then finds the corresponding best model as described in Jiang et. al. (2009) with some practical modifications.

# Usage

```
af(
   mf,
   B = 60,
   n.c = 20,
   initial.stepwise = FALSE,
   force.in = NULL,
   cores,
   nvmax,
   c.max,
   screen = FALSE,
   seed = NULL,
   ...
)
```

## **Arguments**

mf	a fitted 'full' model, the result of a call to lm or glm (and in the future lme or lmer).			
В	number of bootstrap replications at each fence boundary value			
n.c	number of boundary values to be considered			
initial.stepwise				
	logical. Performs an initial stepwise procedure to look for the range of model sizes where attention should be focussed. See details for implementation.			
force.in	the names of variables that should be forced into all estimated models			
cores	number of cores to be used when parallel processing the bootstrap			
nvmax	size of the largest model that can still be considered as a viable candidate. Included for performance reasons but if it is an active constraint it could lead to misleading results.			
c.max	manually specify the upper boundary limit. Only applies when initial.stepwise=FALSE.			
screen	logical, whether or not to perform an initial screen for outliers. Highly experimental, use at own risk. Default = FALSE.			
seed	random seed for reproducible results			
	further arguments (currently unused)			

4 af

#### **Details**

The initial stepwise procedure performs forward stepwise model selection using the AIC and backward stepwise model selection using BIC. In general the backwise selection via the more conservative BIC will tend to select a smaller model than that of the forward selection AIC approach. The size of these two models is found, and we go two dimensions smaller and larger to estimate a sensible range of c values over which to perform a parametric bootstrap.

This procedure can take some time. It is recommended that you start with a relatively small number of bootstrap samples (B) and grid of boundary values (n.c) and increase both as required.

If you use initial.stepwise=TRUE then in general you will need a smaller grid of boundary values than if you select initial.stepwise=FALSE. It can be useful to check initial.stepwise=FALSE with a small number of bootstrap replications over a sparse grid to ensure that the initial.stepwise=TRUE has landed you in a reasonable region.

The best.only=FALSE option when plotting the results of the adaptive fence is a modification to the adaptive fence procedure which considers all models at a particular size that pass the fence hurdle when calculating the p\* values. In particular, for each value of c and at each bootstrap replication, if a candidate model is found that passes the fence, then we look to see if there are any other models of the same size that also pass the fence. If no other models of the same size pass the fence, then that model is allocated a weight of 1. If there are two models that pass the fence, then the best model is allocated a weight of 1/2. If three models pass the fence, the best model gets a weight of 1/3, and so on. After B bootstrap replications, we aggregate the weights by summing over the various models. The p\* value is the maximum aggregated weight divided by the number of bootstrap replications. This correction penalises the probability associated with the best model if there were other models of the same size that also passed the fence hurdle. The rationale being that if a model has no redundant variables then it will be the only model at that size that passes the fence over a range of values of c. The result is more pronounced peaks which can help to determine the location of the correct peak and identify the optimal c\*.

See ?plot.af or help("plot.af") for details of the plot method associated with the result.

#### References

Jiang J., Nguyen T., Sunil Rao J. (2009), A simplified adaptive fence procedure, Statistics & Probability Letters, 79(5):625-629. doi: 10.1016/j.spl.2008.10.014

Jiang J., Sunil Rao J., Gu Z, Nguyen T. (2008), Fence methods for mixed model selection, Annals of Statistics, 36(4):1669-1692. doi: 10.1214/07-AOS517

#### See Also

```
plot.af
Other fence: glmfence(), lmfence()
```

#### **Examples**

```
n = 100
set.seed(11)
e = rnorm(n)
x1 = rnorm(n)
x2 = rnorm(n)
```

artificialeg 5

```
x3 = x1^2
x4 = x2^2
x5 = x1*x2
y = 1 + x1 + x2 + e
dat = data.frame(y,x1,x2,x3,x4,x5)
lm1 = lm(y ~ ., data = dat)

## Not run:
af1 = af(lm1, initial.stepwise = TRUE, seed = 1)
summary(af1)
plot(af1)

## End(Not run)
```

artificialeg

Artificial example

## **Description**

An artificial data set which causes stepwise regression procedures to select a non-parsimonious model. The true model is a simple linear regression of y against x8.

#### Usage

```
data(artificialeg)
```

### **Format**

A data frame with 50 observations on 10 variables.

#### **Details**

Inspired by the pathoeg data set in the MPV pacakge.

# Examples

```
data(artificialeg)
full.mod = lm(y~.,data=artificialeg)
step(full.mod)
# generating model
n=50
set.seed(8) # a seed of 2 also works
x1 = rnorm(n,0.22,2)
x7 = 0.5*x1 + rnorm(n,0,sd=2)
x6 = -0.75*x1 + rnorm(n,0,3)
x3 = -0.5-0.5*x6 + rnorm(n,0,2)
x9 = rnorm(n,0.6,3.5)
x4 = 0.5*x9 + rnorm(n,0,sd=3)
x2 = -0.5 + 0.5*x9 + rnorm(n,0,sd=2)
x5 = -0.5*x2+0.5*x3+0.5*x6-0.5*x9+rnorm(n,0,1.5)
```

6 bglmnet

```
x8 = x1 + x2 - 2*x3 - 0.3*x4 + x5 - 1.6*x6 - 1*x7 + x9 + rnorm(n,0,0.5)

y = 0.6*x8 + rnorm(n,0,2)

artificialeg = round(data.frame(x1,x2,x3,x4,x5,x6,x7,x8,x9,y),1)
```

bglmnet

Model stability and variable importance plots for glmnet

#### Description

Model stability and variable importance plots for glmnet

# Usage

```
bglmnet(
   mf,
   nlambda = 100,
   lambda = NULL,
   B = 100,
   penalty.factor,
   screen = FALSE,
   redundant = TRUE,
   cores = NULL,
   force.in = NULL,
   seed = NULL
)
```

# Arguments

a fitted 'full' model, the result of a call to lm or glm.

how many penalty values to consider. Default = 100.

lambda manually specify the penalty values (optional).

B number of bootstrap replications

penalty.factor Separate penalty factors can be applied to each coefficient. This is a number that

multiplies lambda to allow differential shrinkage. Can be 0 for some variables, which implies no shrinkage, and that variable is always included in the model. Default is 1 for all variables (and implicitly infinity for variables listed in exclude). Note: the penalty factors are internally rescaled to sum to nvars, and the

lambda sequence will reflect this change.

screen logical, whether or not to perform an initial screen for outliers. Highly experi-

mental, use at own risk. Default = FALSE.

redundant logical, whether or not to add a redundant variable. Default = TRUE.

cores number of cores to be used when parallel processing the bootstrap (Not yet

implemented.)

force.in the names of variables that should be forced into all estimated models. (Not yet

implemented.)

seed random seed for reproducible results

bodyfat 7

### **Details**

The result of this function is essentially just a list. The supplied plot method provides a way to visualise the results.

#### See Also

```
plot.bglmnet
```

### **Examples**

```
n = 100
set.seed(11)
e = rnorm(n)
x1 = rnorm(n)
x2 = rnorm(n)
x3 = x1^2
x4 = x2^2
x5 = x1*x2
y = 1 + x1 + x2 + e
dat = data.frame(y, x1, x2, x3, x4, x5)
lm1 = lm(y \sim ., data = dat)
## Not run:
bg1 = bglmnet(lm1, seed = 1)
# plot(bg1, which = "boot_size", interactive = TRUE)
plot(bg1, which = "boot_size", interactive = FALSE)
# plot(bg1, which = "vip", interactive = TRUE)
plot(bg1, which = "vip", interactive = FALSE)
## End(Not run)
```

bodyfat

Body fat data set

## **Description**

A data frame with 128 observations on 15 variables.

#### Usage

```
data(bodyfat)
```

# **Format**

A data frame with 128 observations on 15 variables.

Id Identifier

Bodyfat Bodyfat percentage

8 diabetes

```
Age Age (years)
```

Weight Weight (kg)

**Height** Height (inches)

Neck Neck circumference (cm)

Chest Chest circumference (cm)

**Abdo** Abdomen circumference (cm) "at the umbilicus and level with the iliac crest"

**Hip** Hip circumference (cm)

Thigh Thigh circumference (cm)

Knee Knee circumference (cm)

**Ankle** Ankle circumference (cm)

**Bic** Extended biceps circumference (cm)

**Fore** Forearm circumference (cm)

Wrist Wrist circumference (cm) "distal to the styloid processes"

#### **Details**

A subset of the 252 observations available in the mfp package. The selected observations avoid known high leverage points and outliers. The unused points from the data set could be used to validate selected models.

#### References

Johnson W (1996, Vol 4). Fitting percentage of body fat to simple body measurements. Journal of Statistics Education. Bodyfat data retrieved from http://www.amstat.org/publications/jse/v4n1/datasets.johnson.html An expanded version is included in the mfp R package.

#### **Examples**

```
data(bodyfat)
full.mod = lm(Bodyfat~.,data=subset(bodyfat,select=-Id))
```

diabetes

Blood and other measurements in diabetics

#### **Description**

The diabetes data frame has 442 rows and 11 columns. These are the data used in Efron et al. (2004).

```
data(diabetes)
```

fev 9

#### **Format**

A data frame with 442 observations on 11 variables.

age Age

sex Gender

bmi Body mass index

map Mean arterial pressure (average blood pressure)

tc Total cholesterol (mg/dL)? Desirable range: below 200 mg/dL

ldl Low-density lipoprotein ("bad" cholesterol)? Desirable range: below 130 mg/dL

hdl High-density lipoprotein ("good" cholesterol)? Desirable range: above 40 mg/dL

tch Blood serum measurement

ltg Blood serum measurement

glu Blood serum measurement (glucose?)

y A quantitative measure of disease progression one year after baseline

#### **Details**

Data sourced from http://web.stanford.edu/~hastie/Papers/LARS

#### References

Efron, B., Hastie, T., Johnstone, I., Tibshirani, R., (2004). Least angle regression. The Annals of Statistics 32(2) 407-499. DOI: 10.1214/009053604000000067

# **Examples**

```
data(diabetes)
full.mod = lm(y~.,data=diabetes)
```

fev

Forced Expiratory Volume

# Description

This data set consists of 654 observations on youths aged 3 to 19 from East Boston recorded duing the middle to late 1970's. Forced expiratory volume (FEV), a measure of lung capacity, is the variable of interest. Age and height are two continuous predictors. Sex and smoke are two categorical predictors.

```
data(fev)
```

10 glmfence

#### **Format**

A data frame with 654 observations on 5 variables.

```
age Age (years)
```

**fev** Forced expiratory volume (liters). Roughly the amount of air an individual can exhale in the first second of a forceful breath.

```
height Height (inches).
```

```
sex Female is 0. Male is 1.
```

**smoke** A binary variable indicating whether or not the youth smokes. Nonsmoker is 0. Smoker is 1.

#### **Details**

Copies of this data set can also be found in the coneproj and tmle packages.

#### References

Tager, I. B., Weiss, S. T., Rosner, B., and Speizer, F. E. (1979). Effect of parental cigarette smoking on pulmonary function in children. *American Journal of Epidemiology*, **110**, 15-26.

Rosner, B. (1999). Fundamentals of Biostatistics, 5th Ed., Pacific Grove, CA: Duxbury.

Kahn, M.J. (2005). An Exhalent Problem for Teaching Statistics. *Journal of Statistics Education*, **13**(2). http://www.amstat.org/publications/jse/v13n2/datasets.kahn.html

# **Examples**

```
data(fev)
full.mod = lm(fev~.,data=fev)
step(full.mod)
```

glmfence

The fence procedure for generalised linear models

#### **Description**

This function implements the fence procedure to find the best generalised linear model.

```
glmfence(mf, cstar, nvmax, adaptive = TRUE, trace = TRUE, ...)
```

Imfence 11

#### **Arguments**

mf an object of class glm specifying the full model.

cstar the boundary of the fence, typically found through bootstrapping.

nvmax the maximum number of variables that will be be considered in the model.

adaptive logical. If TRUE the boundary of the fence is given by cstar. Otherwise, it the

original (non-adaptive) fence is performed where the boundary is cstar\*hat(sigma)\_M,tildeM.

trace logical. If TRUE the function prints out its progress as it iterates up through the

dimensions.

.. further arguments (currently unused)

#### References

Jiming Jiang, Thuan Nguyen, J. Sunil Rao, A simplified adaptive fence procedure, Statistics & Probability Letters, Volume 79, Issue 5, 1 March 2009, Pages 625-629, http://dx.doi.org/10.1016/j.spl.2008.10.014.

#### See Also

```
af, 1mfence
```

Other fence: af(), lmfence()

1mfence The fence procedure for linear models

## **Description**

This function implements the fence procedure to find the best linear model.

# Usage

```
lmfence(mf, cstar, nvmax, adaptive = TRUE, trace = TRUE, force.in = NULL, ...)
```

#### **Arguments**

mf an object of class 1m specifying the full model.

cstar the boundary of the fence, typically found through bootstrapping.

nvmax the maximum number of variables that will be be considered in the model.

adaptive logical. If TRUE the boundary of the fence is given by cstar. Otherwise, it the

original (non-adaptive) fence is performed where the boundary is cstar\*hat(sigma)\_M,tildeM.

trace logical. If TRUE the function prints out its progress as it iterates up through the

dimensions.

force.in the names of variables that should be forced into all estimated models.

. . . further arguments (currently unused)

12 mplot

#### References

Jiming Jiang, Thuan Nguyen, J. Sunil Rao, A simplified adaptive fence procedure, Statistics & Probability Letters, Volume 79, Issue 5, 1 March 2009, Pages 625-629, http://dx.doi.org/10.1016/j.spl.2008.10.014.

#### See Also

```
af, glmfence
Other fence: af(), glmfence()
```

#### **Examples**

```
n = 40 # sample size
beta = c(1,2,3,0,0)
K=length(beta)
set.seed(198)
X = cbind(1,matrix(rnorm(n*(K-1)),ncol=K-1))
e = rnorm(n)
y = X%*%beta + e
dat = data.frame(y,X[,-1])
# Non-adaptive approach (not recommended)
lm1 = lm(y~.,data=dat)
lmfence(lm1,cstar=log(n),adaptive=FALSE)
```

mplot

Model selection and stability curves

### Description

Opens a shiny GUI to investigate a range of model selection and stability issues

### Usage

```
mplot(mf, ...)
```

### **Arguments**

```
mf a fitted model.... objects of type vis or af or bglmnet.
```

## References

Tarr G, Mueller S and Welsh AH (2018). mplot: An R Package for Graphical Model Stability and Variable Selection Procedures. Journal of Statistical Software, 83(9), pp. 1-28. doi: 10.18637/jss.v083.i09

plot.af

#### **Examples**

```
n = 100
set.seed(11)
e = rnorm(n)
x1 = rnorm(n)
x2 = rnorm(n)
x3 = x1^2
x4 = x2^2
x5 = x1*x2
y = 1 + x1 + x2 + e
dat = round(data.frame(y,x1,x2,x3,x4,x5),2)
lm1 = lm(y \sim ., data = dat)
## Not run:
v1 = vis(lm1)
af1 = af(lm1)
bg1 = bglmnet(lm1)
mplot(lm1, v1, af1, bg1)
## End(Not run)
```

plot.af

Plot diagnostics for an af object

## **Description**

Summary plot of the bootstrap results of an af object.

```
## S3 method for class 'af'
plot(
  х,
 pch,
  interactive = FALSE,
  classic = NULL,
  tag = NULL,
  shiny = FALSE,
  best.only = FALSE,
 width = 800,
 height = 400,
  fontSize = 12,
  left = 50,
  top = 30,
  chartWidth = "60%",
  chartHeight = "80%",
  backgroundColor = "transparent",
  legend.position = "top",
```

14 plot.af

```
model.wrap = NULL,
legend.space = NULL,
options = NULL,
...
)
```

#### **Arguments**

x af object, the result of af

pch plotting character, i.e., symbol to use

interactive logical. If interactive=TRUE a googleVis plot is provided instead of the base

graphics plot. Default is interactive=FALSE.

classic logical. Depricated. If classic=TRUE a base graphics plot is provided instead

of a googleVis plot. For now specifying classic will overwrite the default interactive behaviour, though this is likely to be removed in the future.

tag Default NULL. Name tag of the objects to be extracted from a gvis (googleVis)

object.

The default tag for is NULL, which will result in R opening a browser window. Setting tag='chart' or setting options(gvis.plot.tag='chart') is useful

when googleVis is used in scripts, like knitr or rmarkdown.

shiny Default FALSE. Set to TRUE when using in a shiny interface.

best.only logical determining whether the output used the standard fence approach of only

considering the best models that pass the fence (TRUE) or if it should take into

account all models that pass the fence at each boundary value (FALSE).

width Width of the googleVis chart canvas area, in pixels. Default: 800. height Height of the googleVis chart canvas area, in pixels. Default: 400.

fontSize font size used in googleVis chart. Default: 12.

left space at left of chart (pixels?). Default: "50".

top space at top of chart (pixels?). Default: "30".

chartWidth googleVis chart area width. A simple number is a value in pixels; a string con-

taining a number followed by % is a percentage. Default: "60%"

chartHeight googleVis chart area height. A simple number is a value in pixels; a string

containing a number followed by % is a percentage. Default: "80%"

backgroundColor

The background colour for the main area of the chart. A simple HTML color

string, for example: 'red' or '#00cc00'. Default: 'transparent'

legend.position

legend position, e.g. "topleft" or "bottomright"

model.wrap Optional parameter to split the legend names if they are too long for classic plots.

model.wrap=2 means that there will be two variables per line, model.wrap=2 gives three variables per line and model.wrap=4 gives 4 variables per line.

legend.space Optional parameter to add additional space between the legend items for the

classic plot.

options If you want to specify the full set of googleVis options.

... further arguments (currently unused)

plot.bglmnet 15

#### **Details**

For each value of c a parametric bootstrap is performed under the full model. For each bootstrap sample we identify the smallest model inside the fence,  $\hat{\alpha}(c)$ . We calculate the empirical probability of selecting model  $\alpha$  for a given value of c as

$$p^*(c,\alpha) = P^*\{\hat{\alpha}(c) = \alpha\}.$$

Hence, if B bootstrap replications are performed,  $p^*(c, \alpha)$  is the proportion of times that model  $\alpha$  is selected. Finally, define an overall selection probability,

$$p^*(c) = \max_{\alpha \in \mathcal{A}} p^*(c, \alpha)$$

and we plot  $p^*(c)$  against c. The points on the scatter plot are colour coded by the model that yielded the highest inclusion probability.

plot.bglmnet

Plot diagnostics for a bglmnet object

#### Description

A plot method to visualise the results of a bglmnet object.

```
## S3 method for class 'bglmnet'
plot(
 Х,
 highlight,
  interactive = FALSE,
  classic = NULL,
  tag = NULL,
  shiny = FALSE,
 which = c("vip", "boot", "boot_size"),
 width = 800,
 height = 400,
  fontSize = 12,
  left = 50,
  top = 30,
  chartWidth = "60%",
  chartHeight = "80%",
  axisTitlesPosition = "out",
  dataOpacity = 0.5,
  options = NULL,
  hAxis.logScale = TRUE,
  ylim,
  text = FALSE,
 backgroundColor = "transparent",
```

16 plot.bglmnet

```
legend.position = "right",
jitterk = 0.1,
srt = 45,
max.circle = 15,
min.prob = 0.1,
...
)
```

#### **Arguments**

x bglmnet object, the result of bglmnet

highlight the name of a variable that will be highlighted.

interactive logical. If interactive=TRUE a googleVis plot is provided instead of the base

graphics plot. Default is interactive=FALSE.

classic logical. Depricated. If classic=TRUE a base graphics plot is provided instead

of a googleVis plot. For now specifying classic will overwrite the default interactive behaviour, though this is likely to be removed in the future.

tag Default NULL. Name tag of the objects to be extracted from a gvis (googleVis)

object.

The default tag for is NULL, which will result in R opening a browser window. Setting tag='chart' or setting options(gvis.plot.tag='chart') is useful

when googleVis is used in scripts, like knitr or rmarkdown.

shiny Default FALSE. Set to TRUE when using in a shiny interface.

which a vector specifying the plots to be output. Variable inclusion type plots which

= "vip" or plots where the size of the point representing each model is proportional to selection probabilities by model size which = "boot\_size" or by

penalty paramter which = "boot".

width Width of the googleVis chart canvas area, in pixels. Default: 800.

height Height of the googleVis chart canvas area, in pixels. Default: 400.

fontSize font size used in googleVis chart. Default: 12.

left space at left of chart (pixels?). Default: "50".

top space at top of chart (pixels?). Default: "30".

chartWidth googleVis chart area width. A simple number is a value in pixels; a string con-

taining a number followed by % is a percentage. Default: "60%"

chartHeight googleVis chart area height. A simple number is a value in pixels; a string

containing a number followed by % is a percentage. Default: "80%"

axisTitlesPosition

Where to place the googleVis axis titles, compared to the chart area. Supported values: "in" - Draw the axis titles inside the the chart area. "out" - Draw the axis

titles outside the chart area. "none" -  $\mbox{Omit}$  the axis titles.

dataOpacity The transparency of googleVis data points, with 1.0 being completely opaque

and 0.0 fully transparent.

options a list to be passed to the googleVis function giving complete control over the

output. Specifying a value for options overwrites all other plotting variables.

plot.vis 17

hAxis.logScale logical, whether or not to use a log scale on the horizontal axis. Default = TRUE.

ylim the y limits of the which="boot" plots.

text logical, whether or not to add text labels to classic boot plot. Default = FALSE.

backgroundColor

The background colour for the main area of the chart. A simple HTML color

string, for example: 'red' or '#00cc00'. Default: 'transparent'

legend.position

the postion of the legend for classic plots. Default legend.position="right" alternatives include legend.position="top" and legend.position="bottom"

jitterk amount of jittering of the model size in the lvk and boot plots. Default = 0.1.

srt when text=TRUE, the angle of rotation for the text labels. Default = 45.

max.circle determines the maximum circle size. Default = 15.

min.prob lower bound on the probability of a model being selected. If a model has a

selection probability lower than min. prob it will not be plotted.

... further arguments (currently unused)

#### See Also

bglmnet

plot.vis

Plot diagnostics for a vis object

### **Description**

A plot method to visualise the results of a vis object.

```
## S3 method for class 'vis'
plot(
  х,
 highlight,
  interactive = FALSE,
  classic = NULL,
  tag = NULL,
  shiny = FALSE,
  nbest = "all",
  which = c("vip", "lvk", "boot"),
 width = 800,
  height = 400,
  fontSize = 12,
  left = 50,
  top = 30,
  chartWidth = "60%",
```

18 plot.vis

```
chartHeight = "80%",
  axisTitlesPosition = "out",
  dataOpacity = 0.5,
  options = NULL,
  ylim,
  legend.position = "right",
  backgroundColor = "transparent",
  text = FALSE,
  min.prob = 0.4,
  srt = 45,
  max.circle = 15,
  print.full.model = FALSE,
  jitterk = 0.1,
  seed = NULL,
  ...
)
```

#### **Arguments**

x vis object, the result of vis
---------------------------------

highlight the name of a variable that will be highlighted

interactive logical. If interactive=TRUE a googleVis plot is provided instead of the base

graphics plot. Default is interactive=FALSE.

classic logical. Depricated. If classic=TRUE a base graphics plot is provided instead

of a googleVis plot. For now specifying classic will overwrite the default

interactive behaviour, though this is likely to be removed in the future.

tag Default NULL. Name tag of the objects to be extracted from a gvis (googleVis)

object.

The default tag for is NULL, which will result in R opening a browser window. Setting tag='chart' or setting options(gvis.plot.tag='chart') is useful

when googleVis is used in scripts, like knitr or rmarkdown.

shiny Default FALSE. Set to TRUE when using in a shiny interface.

nbest maximum number of models at each model size that will be considered for the

lvk plot. Can also take a value of "all" which displays all models (default).

which a vector specifying the plots to be output. Variable inclusion plots which="vip";

description loss against model size which="lvk"; bootstrapped description loss

against model size which="boot".

width Width of the googleVis chart canvas area, in pixels. Default: 800.

height Height of the googleVis chart canvas area, in pixels. Default: 400.

fontSize font size used in googleVis chart. Default: 12.

left space at left of chart (pixels?). Default: "50".

top space at top of chart (pixels?). Default: "30".

chartWidth googleVis chart area width. A simple number is a value in pixels; a string con-

taining a number followed by % is a percentage. Default: "60%"

plot.vis 19

chartHeight googleVis chart area height. A simple number is a value in pixels; a string

containing a number followed by % is a percentage. Default: "80%"

axisTitlesPosition

Where to place the googleVis axis titles, compared to the chart area. Supported values: "in" - Draw the axis titles inside the the chart area. "out" - Draw the axis

titles outside the chart area. "none" - Omit the axis titles.

dataOpacity The transparency of googleVis data points, with 1.0 being completely opaque

and 0.0 fully transparent.

options a list to be passed to the googleVis function giving complete control over the

output. Specifying a value for options overwrites all other plotting variables.

ylim the y limits of the lvk and boot plots.

legend.position

the postion of the legend for classic plots. Default legend.position="right" alternatives include legend.position="top" and legend.position="bottom"

backgroundColor

The background colour for the main area of the chart. A simple HTML color string, for example: 'red' or '#00cc00'. Default: 'null' (there is an issue with GoogleCharts when setting 'transparent' related to the zoom window sticking - area that's corted out, the default will shound back to 'transparent')

once that's sorted out, the default will change back to 'transparent')

text logical, whether or not to add text labels to classic boot plot. Default = FALSE.

min.prob when text=TRUE, a lower bound on the probability of selection before a text

label is shown.

srt when text=TRUE, the angle of rotation for the text labels. Default = 45.

max.circle determines the maximum circle size. Default = 15.

print.full.model

logical, when text=TRUE this determines if the full model gets a label or not.

Default=FALSE.

jitterk amount of jittering of the model size in the lvk and boot plots. Default = 0.1.

seed random seed for reproducible results
... further arguments (currently unused)

#### Details

Specifying which = "lvk" generates a scatter plot where the points correspond to description loss is plot against model size for each model considered. The highlight argument is used to differentiate models that contain a particular variable from those that do not.

Specifying which = "boot" generates a scatter plot where each circle represents a model with a non-zero bootstrap probability, that is, each model that was selected as the best model of a particular dimension in at least one bootstrap replication. The area of each circle is proportional to the corresponding model's bootstrapped selection probability.

#### References

Mueller, S. and Welsh, A. H. (2010), On model selection curves. International Statistical Review, 78:240-256. doi: 10.1111/j.1751-5823.2010.00108.x

20 print.af

Murray, K., Heritier, S. and Mueller, S. (2013), Graphical tools for model selection in generalized linear models. Statistics in Medicine, 32:4438-4451. doi: 10.1002/sim.5855

Tarr G, Mueller S and Welsh AH (2018). mplot: An R Package for Graphical Model Stability and Variable Selection Procedures. Journal of Statistical Software, 83(9), pp. 1-28. doi: 10.18637/jss.v083.i09

#### See Also

vis

#### **Examples**

```
n = 100
set.seed(11)
e = rnorm(n)
x1 = rnorm(n)
x2 = rnorm(n)
x3 = x1^2
x4 = x2^2
x5 = x1*x2
y = 1 + x1 + x2 + e
dat = data.frame(y,x1,x2,x3,x4,x5)
lm1 = lm(y^{-}.,data=dat)
## Not run:
v1 = vis(lm1, seed = 1)
plot(v1, highlight = "x1", which = "lvk")
plot(v1, which = "boot")
plot(v1, which = "vip")
## End(Not run)
```

print.af

Print method for an af object

### **Description**

Prints basic output of the bootstrap results of an af object.

```
## S3 method for class 'af'
print(x, best.only = TRUE, ...)
```

print.vis 21

# **Arguments**

X	an af object, the result of af
best.only	logical determining whether the output used the standard fence approach of only considering the best models that pass the fence (TRUE) or if it should take into account all models that pass the fence at each boundary value (FALSE).
• • •	further arguments (currently unused)

print.vis

Print method for a vis object

# **Description**

Prints basic output of the bootstrap results of an vis object.

#### Usage

```
## S3 method for class 'vis'
print(x, min.prob = 0.3, print.full.model = FALSE, ...)
```

# Arguments

```
x a vis object, the result of vis
min.prob a lower bound on the probability of selection before the result is printed
print.full.model
logical, determines if the full model gets printed or not. Default=FALSE.
... further arguments (currently unused)
```

process.fn

Process results within af function

## **Description**

This function is used by the af function to process the results when iterating over different boundary values

# Usage

```
process.fn(fence.mod, fence.rank)
```

# **Arguments**

fence.mod set of fence models
fence.rank set of fence model ranks

22 vis

summary.af

Summary method for an af object

### Description

Provides comprehensive output of the bootstrap results of an af object.

## Usage

```
## S3 method for class 'af'
summary(object, best.only = TRUE, ...)
```

### **Arguments**

object af object, the result of af

best.only logical determining whether the output used the standard fence approach of only

considering the best models that pass the fence (TRUE) or if it should take into

account all models that pass the fence at each boundary value (FALSE).

... further arguments (currently unused)

vis

Model stability and variable inclusion plots

# Description

Calculates and provides the plot methods for standard and bootstrap enhanced model stability plots (lvk and boot) as well as variable inclusion plots (vip).

```
vis(
  mf,
  nvmax,
  B = 100,
  lambda.max,
  nbest = "all",
  use.glmulti = FALSE,
  cores,
  force.in = NULL,
  screen = FALSE,
  redundant = TRUE,
  seed = NULL,
  ...
)
```

vis 23

#### **Arguments**

mf a fitted 'full' model, the result of a call to lm or glm (and in the future lme or lmer) size of the largest model that can still be considered as a viable candidate nvmax В number of bootstrap replications lambda.max maximum penalty value for the vip plot, defaults to 2\*log(n) nbest maximum number of models at each model size that will be considered for the lvk plot. Can also take a value of "all" which displays all models. use.glmulti logical. Whether to use the glmulti package instead of bestglm. Default use.glmulti=FALSE. cores number of cores to be used when parallel processing the bootstrap force.in the names of variables that should be forced into all estimated models. (Not yet implemented.) logical, whether or not to perform an initial screen for outliers. Highly experiscreen mental, use at own risk. Default = FALSE. redundant logical, whether or not to add a redundant variable. Default = TRUE. random seed for reproducible results seed

#### **Details**

The result of this function is essentially just a list. The supplied plot method provides a way to visualise the results.

See ?plot.vis or help("plot.vis") for details of the plot method associated with the result.

further arguments (currently unused)

#### References

Mueller, S. and Welsh, A. H. (2010), On model selection curves. International Statistical Review, 78:240-256. doi: 10.1111/j.1751-5823.2010.00108.x

Murray, K., Heritier, S. and Mueller, S. (2013), Graphical tools for model selection in generalized linear models. Statistics in Medicine, 32:4438-4451. doi: 10.1002/sim.5855

Tarr G, Mueller S and Welsh AH (2018). mplot: An R Package for Graphical Model Stability and Variable Selection Procedures. Journal of Statistical Software, 83(9), pp. 1-28. doi: 10.18637/jss.v083.i09

#### See Also

```
plot.vis
```

#### **Examples**

```
n = 100
set.seed(11)
e = rnorm(n)
x1 = rnorm(n)
x2 = rnorm(n)
```

24 wallabies

```
x3 = x1^2
x4 = x2^2
x5 = x1*x2
y = 1 + x1 + x2 + e
dat = data.frame(y, x1, x2, x3, x4, x5)
lm1 = lm(y ~ ., data = dat)

## Not run:
v1 = vis(lm1, seed = 1)
plot(v1, highlight = "x1", which = "lvk")
plot(v1, which = "boot")
plot(v1, which = "vip")

## End(Not run)
```

wallabies

Rock-wallabies data set

# Description

On Chalkers Top in the Warrumbungles (NSW, Australia) 200 evenly distributed one metre squared plots were surveyed. Plots were placed at a density of 7-13 per hectare. The presence or absence of fresh (<1 month old) scats of rock-wallabies was recorded for each plot along with location and a selection of predictor variables.

#### Usage

```
data(wallabies)
```

#### **Format**

A data frame with 200 observations on 9 variables.

rw Presence of rock-wallaby scat

edible Percentage cover of edible vegetation

inedible Percentage cover of inedible vegetation

canopy Percentage canopy cover

distance Distance from diurnal refuge

**shelter** Whether or not a plot occurred within a shelter point (large rock or boulder pile)

lat Latitude of the plot location

long Longitude of the plot location

wallabies 25

#### **Details**

Macropods defaecate randomly as they forage and scat (faecal pellet) surveys are a reliable method for detecting the presence of rock-wallabies and other macropods. Scats are used as an indication of spatial foraging patterns of rock-wallabies and sympatric macropods. Scats deposited while foraging were not confused with scats deposited while resting because the daytime refuge areas of rock-wallabies were known in detail for each colony and no samples were taken from those areas. Each of the 200 sites were examined separately to account for the different levels of predation risk and the abundance of rock-wallabies.

#### References

Tuft KD, Crowther MS, Connell K, Mueller S and McArthur C (2011), Predation risk and competitive interactions affect foraging of an endangered refuge-dependent herbivore. Animal Conservation, 14: 447-457. doi: 10.1111/j.1469-1795.2011.00446.x

# **Examples**

```
data(wallabies)
wdat = data.frame(subset(wallabies,select=-c(lat,long)),
    EaD = wallabies$edible*wallabies$distance,
    EaS = wallabies$edible*wallabies$shelter,
    DaS = wallabies$distance*wallabies$shelter)
M1 = glm(rw~., family = binomial(link = "logit"), data = wdat)
```

# **Index**

* Internal	process.fn, 21
glmfence, 10	summary.af, 22
Imfence, 11 process.fn, 21	3diiiiidi y . di , 22
* datasets	vis, 18, 20, 21, 22
artificialeg, 5 bodyfat, 7 diabetes, 8 fev, 9 wallabies, 24	wallabies, 24
* fence	
af, 3 glmfence, 10 lmfence, 11 * package mplot-package, 2	
af, 3, 11, 12, 14, 21, 22 artificialeg, 5	
bglmnet, 6, <i>16</i> , <i>17</i> bodyfat, 7	
diabetes, 8	
fev, 9	
glm, <i>11</i> glmfence, <i>4</i> , 10, <i>12</i>	
lm, 11 lmfence, 4, 11, 11	
<pre>mplot, 12 mplot-package, 2</pre>	
plot.af, 4, 13 plot.bglmnet, 7, 15 plot.vis, 17, 23 print.af, 20 print.vis, 21	

26