Package 'ggscidca'

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Type Package

Title Plotting Decision Curve Analysis with Coloured Bars
Version 0.2.3
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Description Decision curve analysis is a method for evaluating and comparing prediction models that incorporates clinical consequences, requires only the data set on which the models are tested, and can be applied to models that have either continuous or dichotomous results. The 'ggscidca' package adds coloured bars of discriminant relevance to the traditional decision curve. Improved practicality and aesthetics. This method was described by Balachandran VP (2015) <doi:10.1016 s1470-2045(14)71116-7="">.</doi:10.1016>
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Breastcancer

A survival data on breast cancer.

Description

A survival data on breast cancer.

Usage

data(Breastcancer)

Format

An object of class data. frame with 660 rows and 12 columns.

Examples

data(Breastcancer)

cmprskstdca

cmprskstdca

Description

Generate data for plotting survival analysis decision curves.

cmprskstdca 3

Arguments

data	a data frame containing the variables in the model.
outcome	the outcome, response variable. Must be a variable contained within the data frame specified in data=.
predictors	the predictor variable(s). Must be a variable(s) contained within the data frame specified in data=.
probability	specifies whether or not each of the independent variables are probabilities. The default is TRUE.
xstart	starting value for x-axis (threshold probability) between 0 and 1. The default is 0.01 .
xstop	stopping value for x-axis (threshold probability) between 0 and 1. The default is 0.99 .
xby	increment for threshold probability. The default is 0.01.
ymin	minimum bound for graph.
harm	specifies the harm(s) associated with the independent variable(s). The default is none.
graph	specifies whether or not to display graph of net benefits. The default is TRUE.
intervention	plot net reduction in interventions
interventionpe	
	number of net reduction in interventions per interger. The default is 100
loess.span	specifies the degree of smoothing. The default is 0.10.
timepoint	specifies the time point at which the decision curve analysis is performed.
cmprsk	if evaluating outcome in presence of a competing risk. The default is FALSE
smooth	specifies whether or not to smooth net benefit curve. The default is FALSE.
ttoutcome	Enter the time variable in your data.
legend.position	
	Set the position of the legend.

Details

This function was created and written by Dr Andrew Vickers to generate decision curve data.

Value

Returns a data for plotting a decision curve.

4 cox.tcdca

Description

You can use it to plot decision curves for multiple generative analysis or competitive risk models.

Arguments

	•••	Fill in multiple survival analysis or competitive risk models. You cannot mix and match.
	newdata	If the decision curve of the validation set is to be analysed. Fill in the validation set data here.
	timepoint	If it is a survival analysis, fill in the point in time you need to study. The default is the median time.
	cmprsk	If it is a competitive risk model, select TRUE here.
	modelnames	Defines the name of the generated image model.
	y.min	The maximum value of the negative part of the picture. Generally defaults to positive values multiplied by 0.4 .
	xstop	The maximum value of the X-axis of the picture.
	y.max	The maximum value of the Y-axis. The default value is the maximum net benefit.
	pyh	The height at which the bars are plotted cannot exceed y.min.
	relcol	The colour of the relevant part of the bar. The default is red.
	irrelcol	The colour of the irrelevant part of the bar. The default is blue.
	relabel	Relevance Tags.
	irrellabel	No relevant tags.
	text.size	Font size.
	text.col	The colour of the font.
	colbar	The default is true, and if false is selected, bar plotting is cancelled.
	merge	If true is selected it will merge the two long zones.
	threshold.text	The default is FALSE, if TRUE is selected, a text message for the threshold will be added.
	threshold.line	The default is FALSE, and if TRUE is selected, lines for the threshold will be added.
	nudge_x	Used to adjust the x-axis position of the point where the threshold is located.
	nudge_y	Used to adjust the y-axis position of the point where the threshold is located.
threshold.linetype		
		The line shape of the threshold line.

The line width of the threshold line.

threshold.linewidth

dca 5

threshold.linecol

The colour of the threshold line.

liftpec Threshold point left displacement.
rightpec Threshold point right displacement.

legend.position

Set the position of the legend.

Splitface Name the faceted image.

Value

A picture.

dca dca

Description

Generate data for plotting survival analysis decision curves.

Arguments

data a data frame containing the variables in the model.

outcome the outcome, response variable. Must be a variable contained within the data

frame specified in data=.

predictors the predictor variable(s). Must be a variable(s) contained within the data frame

specified in data=.

probability specifies whether or not each of the independent variables are probabilities. The

default is TRUE.

xstart starting value for x-axis (threshold probability) between 0 and 1. The default is

0.01.

xstop stopping value for x-axis (threshold probability) between 0 and 1. The default

is 0.99.

xby increment for threshold probability. The default is 0.01.

ymin minimum bound for graph.

harm specifies the harm(s) associated with the independent variable(s). The default is

none.

graph specifies whether or not to display graph of net benefits. The default is TRUE.

intervention plot net reduction in interventions

interventionper

number of net reduction in interventions per interger. The default is 100

loess.span specifies the degree of smoothing. The default is 0.10.

smooth specifies whether or not to smooth net benefit curve. The default is FALSE.

6 df_surv

Details

This function was created and written by Dr Andrew Vickers to generate decision curve data.

Value

Returns a data for plotting a decision curve.

demo

A medical examination related data.

Description

A medical examination related data.

Usage

```
data(demo)
```

Format

An object of class data. frame with 832 rows and 34 columns.

Examples

data(demo)

df_surv

A data for competitive risk modelling.

Description

A data for competitive risk modelling.

Usage

```
data(df_surv)
```

Format

An object of class tbl_df (inherits from tbl, data.frame) with 750 rows and 9 columns.

Examples

```
data(df_surv)
```

LIRI 7

LIRI

A data for random forest analysis.

Description

A data for random forest analysis.

Usage

```
data(LIRI)
```

Format

An object of class data. frame with 232 rows and 6 columns.

Examples

```
data(LIRI)
```

netdata

netdata

Description

netdata

Usage

```
netdata(
  fit,
  newdata = NULL,
  timepoint = "median",
  cmprsk = FALSE,
 modelnames = NULL,
 merge = FALSE,
 y.min = NULL,
  xstop = NULL,
 y.max = NULL,
 pyh = NULL,
  relcol = "#c01e35",
  irrelcol = "#0151a2",
  relabel = "Nomogram relevant",
  irrellabel = "Nomogram irrelevant",
  text.size = 4.5,
  text.col = "green",
```

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```
colbar = TRUE,
  threshold.text = FALSE,
  threshold.line = FALSE,
  nudge_x = 0,
  nudge_y = 0,
  threshold.linetype = 2,
  threshold.linewidth = 1.2,
  threshold.linecol = "black",
  po.text.size = 4,
  po.text.col = "black",
  po.text.fill = "white",
  liftpec = NULL,
  rightpec = NULL,
  legend.position = c(0.85, 0.75)
)
```

Arguments

fit Fill in the model you want to analyze. Support survival analysis and logistic

regression.

newdata If the decision curve of the validation set is to be analysed. Fill in the validation

set data here.

timepoint If it is a survival analysis, fill in the point in time you need to study. The default

is the median time.

cmprsk If it is a competitive risk model, select TRUE here.

modelnames Defines the name of the generated image model.

merge If true is selected it will merge the two long zones.

y.min The maximum value of the negative part of the picture. Generally defaults to

positive values multiplied by 0.4.

xstop The maximum value of the X-axis of the picture.

y.max The maximum value of the Y-axis. The default value is the maximum net benefit.

pyh The height at which the bars are plotted cannot exceed y.min.

relcol The colour of the relevant part of the bar. The default is red.

irrelcol The colour of the irrelevant part of the bar. The default is blue.

relabel Relevance Tags.
irrellabel No relevant tags.

text.size Font size.

text.col The colour of the font.

colbar The default is true, and if false is selected, bar plotting is cancelled.

threshold.text The default is FALSE, if TRUE is selected, a text message for the threshold will

be added.

threshold.line The default is FALSE, and if TRUE is selected, lines for the threshold will be

added.

netdata.ksvm 9

Used to adjust the x-axis position of the point where the threshold is located. nudge_x nudge_y Used to adjust the y-axis position of the point where the threshold is located. threshold.linetype The line shape of the threshold line. threshold.linewidth The line width of the threshold line. threshold.linecol The colour of the threshold line. po.text.size The size of the threshold point text. po.text.col The colour of the threshold point text. The background of the threshold point text. po.text.fill liftpec Threshold point left displacement. rightpec Threshold point right displacement. legend.position Set the position of the legend.

Value

A data used for plotting.

netdata.ksvm

netdata.ksvm

Description

netdata.ksvm

Usage

```
## S3 method for class 'ksvm'
netdata(
  fit,
  newdata = NULL,
  timepoint = "median",
  cmprsk = FALSE,
  modelnames = NULL,
 merge = FALSE,
  y.min = NULL,
  xstop = NULL,
  y.max = NULL,
  pyh = NULL,
  relcol = "#c01e35",
  irrelcol = "#0151a2",
  relabel = "Nomogram relevant",
  irrellabel = "Nomogram irrelevant",
```

10 netdata.ksvm

```
text.size = 4.5,
  text.col = "green",
  colbar = TRUE,
  threshold.text = FALSE,
  threshold.line = FALSE,
  nudge_x = 0,
  nudge_y = 0,
  threshold.linetype = 2,
  threshold.linewidth = 1.2,
  threshold.linecol = "black",
  po.text.size = 4,
  po.text.col = "black",
  po.text.fill = "white",
  liftpec = NULL,
  rightpec = NULL,
  legend.position = c(0.85, 0.75)
)
```

Arguments

fit Fill in the model you want to analyze. Support survival analysis and logistic

regression.

newdata If the decision curve of the validation set is to be analysed. Fill in the validation

set data here.

timepoint If it is a survival analysis, fill in the point in time you need to study. The default

is the median time.

cmprsk If it is a competitive risk model, select TRUE here.

modelnames Defines the name of the generated image model.

merge If true is selected it will merge the two long zones.

y.min The maximum value of the negative part of the picture. Generally defaults to

positive values multiplied by 0.4.

xstop The maximum value of the X-axis of the picture.

y.max The maximum value of the Y-axis. The default value is the maximum net benefit.

pyh The height at which the bars are plotted cannot exceed y.min.

relcol The colour of the relevant part of the bar. The default is red.

irrelcol The colour of the irrelevant part of the bar. The default is blue.

relabel Relevance Tags.
irrellabel No relevant tags.

text.size Font size.

text.col The colour of the font.

colbar The default is true, and if false is selected, bar plotting is cancelled.

threshold.text The default is FALSE, if TRUE is selected, a text message for the threshold will

be added.

newcrr 11

threshold.line The default is FALSE, and if TRUE is selected, lines for the threshold will be

added.

nudge_x Used to adjust the x-axis position of the point where the threshold is located.

nudge_y Used to adjust the y-axis position of the point where the threshold is located.

threshold.linetype

The line shape of the threshold line.

threshold.linewidth

The line width of the threshold line.

threshold.linecol

The colour of the threshold line.

po.text.size The size of the threshold point text.

po.text.col The colour of the threshold point text.

po.text.fill The background of the threshold point text.

liftpec Threshold point left displacement.
rightpec Threshold point right displacement.

legend.position

Set the position of the legend.

Value

A data used for plotting.

newcrr newcrr

Description

Types of transformation of survival analysis models into competitive risk models.

Usage

```
newcrr(fit, cencode = 0, failcode = 1)
```

Arguments

fit Modelling for Survival Analysis.

cencode Censor status, default is 0.
failcode Events of interest, default is 1.

Value

A list of competing risk model formats.

12 scidca

scidca scidca

Description

You can use it to generate a decision curve with coloured bars.

Usage

```
scidca(
  fit,
  newdata = NULL,
  timepoint = "median",
  cmprsk = FALSE,
 modelnames = NULL,
 merge = FALSE,
 y.min = NULL,
 xstop = NULL,
 y.max = NULL,
 pyh = NULL,
  relcol = "#c01e35",
  irrelcol = "#0151a2",
  relabel = "Nomogram relevant",
  irrellabel = "Nomogram irrelevant",
  text.size = 4.5,
  text.col = "green",
  colbar = TRUE,
  threshold.text = FALSE,
  threshold.line = FALSE,
  nudge_x = 0,
  nudge_y = 0,
  threshold.linetype = 2,
  threshold.linewidth = 1.2,
  threshold.linecol = "black",
  po.text.size = 4,
  po.text.col = "black",
 po.text.fill = "white",
  liftpec = NULL,
  rightpec = NULL,
  legend.position = c(0.85, 0.75)
)
```

Arguments

fit

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newdata If the decision curve of the validation set is to be analysed. Fill in the validation

set data here.

timepoint If it is a survival analysis, fill in the point in time you need to study. The default

is the median time.

cmprsk If it is a competitive risk model, select TRUE here.

modelnames Defines the name of the generated image model.

merge If true is selected it will merge the two long zones.

y.min The maximum value of the negative part of the picture. Generally defaults to

positive values multiplied by 0.4.

xstop The maximum value of the X-axis of the picture.

y.max The maximum value of the Y-axis. The default value is the maximum net benefit.

pyh The height at which the bars are plotted cannot exceed y.min.

relcol The colour of the relevant part of the bar. The default is red.

irrelcol The colour of the irrelevant part of the bar. The default is blue.

relabel Relevance Tags.
irrellabel No relevant tags.

text.size Font size.

text.col The colour of the font.

colbar The default is true, and if false is selected, bar plotting is cancelled.

threshold.text The default is FALSE, if TRUE is selected, a text message for the threshold will

be added.

threshold.line The default is FALSE, and if TRUE is selected, lines for the threshold will be

added.

nudge_x Used to adjust the x-axis position of the point where the threshold is located.

nudge_y Used to adjust the y-axis position of the point where the threshold is located.

threshold.linetype

The line shape of the threshold line.

threshold.linewidth

The line width of the threshold line.

threshold.linecol

The colour of the threshold line.

po.text.size The size of the threshold point text.

po.text.col The colour of the threshold point text.

po.text.fill The background of the threshold point text.

liftpec Threshold point left displacement.
rightpec Threshold point right displacement.

legend.position

Set the position of the legend.

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Details

Table 1 represents the relationship between the baseline values of the data. This function can be easily done. Creates 'Table 1', i.e., description of baseline patient characteristics, which is essential in every medical research. Supports both continuous and categorical variables, as well as p-values and standardized mean differences.

Value

A picture.

Examples

```
library(survival)
library(reshape2)
library(ggplot2)
##Import the internal data of the R package
bc<-Breastcancer
##Categorical variables converted to factors
bc$histgrad<-as.factor(bc$histgrad)</pre>
bc$er<-as.factor(bc$er)</pre>
bc$pr<-as.factor(bc$pr)</pre>
bc$ln_yesno<-as.factor(bc$ln_yesno)</pre>
##Generate Survival Analysis Model
f1<-coxph(Surv(time, status)~er+histgrad+pr+age+ln_yesno,bc)
##Draw decision curve
scidca(f1)
scidca(f1,threshold.line = TRUE,threshold.text = TRUE)
##logistic regression model
fit<-glm(status~er+histgrad+pr+age+ln_yesno,family = binomial(link = "logit"),data=bc)</pre>
##Draw decision curve
scidca(f1)
scidca(f1,threshold.line = TRUE,threshold.text = TRUE)
##random forest model
library(randomForest)
LIRI<-LIRI
set.seed(1)
index <- sample(2,nrow(LIRI),replace = TRUE,prob=c(0.7,0.3))</pre>
traindata <- LIRI[index==1,]</pre>
testdata <- LIRI[index==2,]</pre>
traindata$status<-as.factor(traindata$status)</pre>
#Modelling random forests
fit<-randomForest(status ~ANLN+CENPA+GPR182+BC02 ,data=traindata,ntree=500,
important=TRUE,proximity=TRUE)
scidca(fit,newdata = traindata)
scidca(fit,newdata = testdata )
scidca(fit,newdata = testdata ,threshold.line = TRUE,threshold.text = TRUE)
```

scidca.coxph 15

scidca.coxph

scidca.coxph

Description

scidca.coxph

Usage

```
## S3 method for class 'coxph'
scidca(
  fit,
  newdata = NULL,
  timepoint = "median",
  cmprsk = FALSE,
 modelnames = NULL,
 merge = FALSE,
 y.min = NULL,
 xstop = NULL,
 y.max = NULL,
  pyh = NULL,
  relcol = "#c01e35",
  irrelcol = "#0151a2",
  relabel = "Nomogram relevant",
  irrellabel = "Nomogram irrelevant",
  text.size = 4.5,
  text.col = "green",
  colbar = TRUE,
  threshold.text = FALSE,
  threshold.line = FALSE,
  nudge_x = 0,
  nudge_y = 0,
  threshold.linetype = 2,
  threshold.linewidth = 1.2,
  threshold.linecol = "black",
  po.text.size = 4,
  po.text.col = "black",
  po.text.fill = "white",
  liftpec = NULL,
  rightpec = NULL,
 legend.position = c(0.85, 0.75)
)
```

Arguments

fit

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newdata If the decision curve of the validation set is to be analysed. Fill in the validation

set data here.

timepoint If it is a survival analysis, fill in the point in time you need to study. The default

is the median time.

cmprsk If it is a competitive risk model, select TRUE here.
modelnames Defines the name of the generated image model.
merge If true is selected it will merge the two long zones.

y.min The maximum value of the negative part of the picture. Generally defaults to

positive values multiplied by 0.4.

xstop The maximum value of the X-axis of the picture.

y.max The maximum value of the Y-axis. The default value is the maximum net benefit.

pyh The height at which the bars are plotted cannot exceed y.min.

relcol The colour of the relevant part of the bar. The default is red.

irrelcol The colour of the irrelevant part of the bar. The default is blue.

relabel Relevance Tags.
irrellabel No relevant tags.

text.size Font size.

text.col The colour of the font.

colbar The default is true, and if false is selected, bar plotting is cancelled.

threshold.text The default is FALSE, if TRUE is selected, a text message for the threshold will

be added.

threshold.line The default is FALSE, and if TRUE is selected, lines for the threshold will be

added.

nudge_x Used to adjust the x-axis position of the point where the threshold is located.

nudge_y Used to adjust the y-axis position of the point where the threshold is located.

threshold.linetype

The line shape of the threshold line.

threshold.linewidth

The line width of the threshold line.

threshold.linecol

The colour of the threshold line.

po.text.size The size of the threshold point text.

po.text.col The colour of the threshold point text.

po.text.fill The background of the threshold point text.

liftpec Threshold point left displacement.rightpec Threshold point right displacement.

legend.position

Set the position of the legend.

Value

scidca.crr 17

scidca.crr

scidca.crr

Description

scidca.crr

Usage

```
## S3 method for class 'crr'
scidca(
  fit,
  newdata = NULL,
  timepoint = "median",
  cmprsk = FALSE,
 modelnames = NULL,
 merge = FALSE,
 y.min = NULL,
 xstop = NULL,
 y.max = NULL,
  pyh = NULL,
  relcol = "#c01e35",
  irrelcol = "#0151a2",
  relabel = "Nomogram relevant",
  irrellabel = "Nomogram irrelevant",
  text.size = 4.5,
  text.col = "green",
  colbar = TRUE,
  threshold.text = FALSE,
  threshold.line = FALSE,
  nudge_x = 0,
  nudge_y = 0,
  threshold.linetype = 2,
  threshold.linewidth = 1.2,
  threshold.linecol = "black",
  po.text.size = 4,
  po.text.col = "black",
  po.text.fill = "white",
  liftpec = NULL,
  rightpec = NULL,
  legend.position = c(0.85, 0.75)
)
```

Arguments

fit

18 scidca.crr

newdata If the decision curve of the validation set is to be analysed. Fill in the validation

set data here.

timepoint If it is a survival analysis, fill in the point in time you need to study. The default

is the median time.

cmprsk If it is a competitive risk model, select TRUE here.

modelnames Defines the name of the generated image model.

merge If true is selected it will merge the two long zones.

y.min The maximum value of the negative part of the picture. Generally defaults to

positive values multiplied by 0.4.

xstop The maximum value of the X-axis of the picture.

y.max The maximum value of the Y-axis. The default value is the maximum net benefit.

pyh The height at which the bars are plotted cannot exceed y.min.

relcol The colour of the relevant part of the bar. The default is red.

irrelcol The colour of the irrelevant part of the bar. The default is blue.

relabel Relevance Tags. irrellabel No relevant tags.

text.size Font size.

text.col The colour of the font.

colbar The default is true, and if false is selected, bar plotting is cancelled.

threshold.text The default is FALSE, if TRUE is selected, a text message for the threshold will

be added.

threshold.line The default is FALSE, and if TRUE is selected, lines for the threshold will be

added.

nudge_x
Used to adjust the x-axis position of the point where the threshold is located.

Used to adjust the y-axis position of the point where the threshold is located.

Used to adjust the y-axis position of the point where the threshold is located.

threshold.linetype

The line shape of the threshold line.

threshold.linewidth

The line width of the threshold line.

threshold.linecol

The colour of the threshold line.

liftpec Threshold point left displacement.
rightpec Threshold point right displacement.

legend.position

Set the position of the legend.

Value

scidca.glm

scidca.glm

scidca.glm

Description

scidca.glm

Usage

```
## S3 method for class 'glm'
scidca(
  fit,
  newdata = NULL,
  timepoint = "median",
  cmprsk = FALSE,
 modelnames = NULL,
 merge = FALSE,
 y.min = NULL,
 xstop = NULL,
 y.max = NULL,
  pyh = NULL,
  relcol = "#c01e35",
  irrelcol = "#0151a2",
  relabel = "Nomogram relevant",
  irrellabel = "Nomogram irrelevant",
  text.size = 4.5,
  text.col = "green",
  colbar = TRUE,
  threshold.text = FALSE,
  threshold.line = FALSE,
  nudge_x = 0,
  nudge_y = 0,
  threshold.linetype = 2,
  threshold.linewidth = 1.2,
  threshold.linecol = "black",
  po.text.size = 4,
  po.text.col = "black",
  po.text.fill = "white",
  liftpec = NULL,
  rightpec = NULL,
 legend.position = c(0.85, 0.75)
)
```

Arguments

fit

20 scidca.glm

newdata If the decision curve of the validation set is to be analysed. Fill in the validation

set data here.

timepoint If it is a survival analysis, fill in the point in time you need to study. The default

is the median time.

cmprsk If it is a competitive risk model, select TRUE here.

modelnames Defines the name of the generated image model.

merge If true is selected it will merge the two long zones.

y.min The maximum value of the negative part of the picture. Generally defaults to

positive values multiplied by 0.4.

xstop The maximum value of the X-axis of the picture.

y.max The maximum value of the Y-axis. The default value is the maximum net benefit.

pyh The height at which the bars are plotted cannot exceed y.min.

relcol The colour of the relevant part of the bar. The default is red.

irrelcol The colour of the irrelevant part of the bar. The default is blue.

relabel Relevance Tags. irrellabel No relevant tags.

text.size Font size.

text.col The colour of the font.

colbar The default is true, and if false is selected, bar plotting is cancelled.

threshold.text The default is FALSE, if TRUE is selected, a text message for the threshold will

be added.

threshold.line The default is FALSE, and if TRUE is selected, lines for the threshold will be

added.

nudge_x Used to adjust the x-axis position of the point where the threshold is located.

nudge_y Used to adjust the y-axis position of the point where the threshold is located.

threshold.linetype

The line shape of the threshold line.

threshold.linewidth

The line width of the threshold line.

threshold.linecol

The colour of the threshold line.

liftpec Threshold point left displacement.
rightpec Threshold point right displacement.

legend.position

Set the position of the legend.

Value

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scidca.ksvm

scidca.ksvm

Description

scidca.ksvm

Usage

```
## S3 method for class 'ksvm'
scidca(
  fit,
  newdata = NULL,
  timepoint = "median",
  cmprsk = FALSE,
 modelnames = NULL,
 merge = FALSE,
 y.min = NULL,
 xstop = NULL,
 y.max = NULL,
 pyh = NULL,
  relcol = "#c01e35",
  irrelcol = "#0151a2",
  relabel = "Nomogram relevant",
  irrellabel = "Nomogram irrelevant",
  text.size = 4.5,
  text.col = "green",
  colbar = TRUE,
  threshold.text = FALSE,
  threshold.line = FALSE,
  nudge_x = 0,
  nudge_y = 0,
  threshold.linetype = 2,
  threshold.linewidth = 1.2,
  threshold.linecol = "black",
  po.text.size = 4,
  po.text.col = "black",
  po.text.fill = "white",
  liftpec = NULL,
  rightpec = NULL,
  legend.position = c(0.85, 0.75)
)
```

Arguments

fit

22 scidca.ksvm

newdata This parameter is indispensable in the random forest decision curve. Fill in your

data.

timepoint If it is a survival analysis, fill in the point in time you need to study. The default

is the median time.

cmprsk If it is a competitive risk model, select TRUE here.

modelnames Defines the name of the generated image model.

merge If true is selected it will merge the two long zones.

y.min The maximum value of the negative part of the picture. Generally defaults to

positive values multiplied by 0.4.

xstop The maximum value of the X-axis of the picture.

y.max The maximum value of the Y-axis. The default value is the maximum net benefit.

pyh The height at which the bars are plotted cannot exceed y.min.

relcol The colour of the relevant part of the bar. The default is red.

irrelcol The colour of the irrelevant part of the bar. The default is blue.

relabel Relevance Tags. irrellabel No relevant tags.

text.size Font size.

text.col The colour of the font.

colbar The default is true, and if false is selected, bar plotting is cancelled.

threshold.text The default is FALSE, if TRUE is selected, a text message for the threshold will

be added.

threshold.line The default is FALSE, and if TRUE is selected, lines for the threshold will be

added.

nudge_x Used to adjust the x-axis position of the point where the threshold is located.

nudge_y Used to adjust the y-axis position of the point where the threshold is located.

threshold.linetype

The line shape of the threshold line.

threshold.linewidth

The line width of the threshold line.

threshold.linecol

The colour of the threshold line.

liftpec Threshold point left displacement.
rightpec Threshold point right displacement.

legend.position

Set the position of the legend.

Value

scidca.randomForest 23

Description

scidca.randomForest

Usage

```
## S3 method for class 'randomForest'
scidca(
  fit,
  newdata = NULL,
  timepoint = "median",
  cmprsk = FALSE,
 modelnames = NULL,
 merge = FALSE,
 y.min = NULL,
 xstop = NULL,
 y.max = NULL,
 pyh = NULL,
  relcol = "#c01e35",
  irrelcol = "#0151a2",
  relabel = "Nomogram relevant",
  irrellabel = "Nomogram irrelevant",
  text.size = 4.5,
  text.col = "green",
  colbar = TRUE,
  threshold.text = FALSE,
  threshold.line = FALSE,
  nudge_x = 0,
  nudge_y = 0,
  threshold.linetype = 2,
  threshold.linewidth = 1.2,
  threshold.linecol = "black",
  po.text.size = 4,
  po.text.col = "black",
  po.text.fill = "white",
  liftpec = NULL,
  rightpec = NULL,
 legend.position = c(0.85, 0.75)
)
```

Arguments

24 scidca.randomForest

newdata This parameter is indispensable in the random forest decision curve. Fill in your

data.

timepoint If it is a survival analysis, fill in the point in time you need to study. The default

is the median time.

cmprsk If it is a competitive risk model, select TRUE here.

modelnames Defines the name of the generated image model.

merge If true is selected it will merge the two long zones.

y.min The maximum value of the negative part of the picture. Generally defaults to

positive values multiplied by 0.4.

xstop The maximum value of the X-axis of the picture.

y.max The maximum value of the Y-axis. The default value is the maximum net benefit.

pyh The height at which the bars are plotted cannot exceed y.min.

relcol The colour of the relevant part of the bar. The default is red.

irrelcol The colour of the irrelevant part of the bar. The default is blue.

relabel Relevance Tags. irrellabel No relevant tags.

text.size Font size.

text.col The colour of the font.

colbar The default is true, and if false is selected, bar plotting is cancelled.

threshold.text The default is FALSE, if TRUE is selected, a text message for the threshold will

be added.

threshold.line The default is FALSE, and if TRUE is selected, lines for the threshold will be

added.

nudge_x Used to adjust the x-axis position of the point where the threshold is located.

nudge_y Used to adjust the y-axis position of the point where the threshold is located.

threshold.linetype

The line shape of the threshold line.

threshold.linewidth

The line width of the threshold line.

threshold.linecol

The colour of the threshold line.

liftpec Threshold point left displacement.
rightpec Threshold point right displacement.

legend.position

Set the position of the legend.

Value

scidca.svm 25

scidca.svm

scidca.svm

Description

scidca.svm

Usage

```
## S3 method for class 'svm'
scidca(
  fit,
  newdata = NULL,
  timepoint = "median",
  cmprsk = FALSE,
 modelnames = NULL,
 merge = FALSE,
 y.min = NULL,
 xstop = NULL,
 y.max = NULL,
 pyh = NULL,
  relcol = "#c01e35",
  irrelcol = "#0151a2",
  relabel = "Nomogram relevant",
  irrellabel = "Nomogram irrelevant",
  text.size = 4.5,
  text.col = "green",
  colbar = TRUE,
  threshold.text = FALSE,
  threshold.line = FALSE,
  nudge_x = 0,
  nudge_y = 0,
  threshold.linetype = 2,
  threshold.linewidth = 1.2,
  threshold.linecol = "black",
  po.text.size = 4,
  po.text.col = "black",
  po.text.fill = "white",
  liftpec = NULL,
  rightpec = NULL,
 legend.position = c(0.85, 0.75)
)
```

Arguments

fit

26 scidca.svm

newdata This parameter is indispensable in the random forest decision curve. Fill in your

data.

timepoint If it is a survival analysis, fill in the point in time you need to study. The default

is the median time.

cmprsk If it is a competitive risk model, select TRUE here.

modelnames Defines the name of the generated image model.

merge If true is selected it will merge the two long zones.

y.min The maximum value of the negative part of the picture. Generally defaults to

positive values multiplied by 0.4.

xstop The maximum value of the X-axis of the picture.

y.max The maximum value of the Y-axis. The default value is the maximum net benefit.

pyh The height at which the bars are plotted cannot exceed y.min.

relcol The colour of the relevant part of the bar. The default is red.

irrelcol The colour of the irrelevant part of the bar. The default is blue.

relabel Relevance Tags. irrellabel No relevant tags.

text.size Font size.

text.col The colour of the font.

colbar The default is true, and if false is selected, bar plotting is cancelled.

threshold.text The default is FALSE, if TRUE is selected, a text message for the threshold will

be added.

threshold.line The default is FALSE, and if TRUE is selected, lines for the threshold will be

added.

nudge_x
Used to adjust the x-axis position of the point where the threshold is located.

Used to adjust the y-axis position of the point where the threshold is located.

Used to adjust the y-axis position of the point where the threshold is located.

threshold.linetype

The line shape of the threshold line.

threshold.linewidth

The line width of the threshold line.

threshold.linecol

The colour of the threshold line.

liftpec Threshold point left displacement.
rightpec Threshold point right displacement.

legend.position

Set the position of the legend.

Value

stdca 27

|--|

Description

Generate data for plotting survival analysis decision curves.

Arguments

data	a data frame containing the variables in the model.
outcome	the outcome, response variable. Must be a variable contained within the data frame specified in data=.
predictors	the predictor variable(s). Must be a variable(s) contained within the data frame specified in data=.
probability	specifies whether or not each of the independent variables are probabilities. The default is TRUE.
xstart	starting value for x-axis (threshold probability) between 0 and 1 . The default is 0.01 .
xstop	stopping value for x-axis (threshold probability) between 0 and 1. The default is 0.99.
xby	increment for threshold probability. The default is 0.01.
ymin	minimum bound for graph.
harm	specifies the harm(s) associated with the independent variable(s). The default is none.
graph	specifies whether or not to display graph of net benefits. The default is TRUE.
intervention	plot net reduction in interventions
interventionpe	
-	number of net reduction in interventions per interger. The default is 100
loess.span	specifies the degree of smoothing. The default is 0.10.
timepoint	specifies the time point at which the decision curve analysis is performed.
cmprsk	if evaluating outcome in presence of a competing risk. The default is FALSE
smooth	specifies whether or not to smooth net benefit curve. The default is FALSE.
ttoutcome	Enter the time variable in your data.

Details

This function was created and written by Dr Andrew Vickers to generate decision curve data.

Value

Returns a data for plotting a decision curve.

28 tcdca

Description

You can use it to plot decision curves for multiple binary classification models.

Arguments

8	
•••	Fill in multiple binary classification models. Cannot populate correlation models with time.
newdata	If the decision curve of the validation set is to be analysed. Fill in the validation set data here.
cmprsk	If it is a competitive risk model, select TRUE here.
modelnames	Defines the name of the generated image model.
y.min	The maximum value of the negative part of the picture. Generally defaults to positive values multiplied by 0.4.
xstop	The maximum value of the X-axis of the picture.
y.max	The maximum value of the Y-axis. The default value is the maximum net benefit.
pyh	The height at which the bars are plotted cannot exceed y.min.
relcol	The colour of the relevant part of the bar. The default is red.
irrelcol	The colour of the irrelevant part of the bar. The default is blue.
relabel	Relevance Tags.
irrellabel	No relevant tags.
text.size	Font size.
text.col	The colour of the font.
colbar	The default is true, and if false is selected, bar plotting is cancelled.
merge	If true is selected it will merge the two long zones.
threshold.text	The default is FALSE, if TRUE is selected, a text message for the threshold will be added.
threshold.line	The default is FALSE, and if TRUE is selected, lines for the threshold will be added.
nudge_x	Used to adjust the x-axis position of the point where the threshold is located.
nudge_y	Used to adjust the y-axis position of the point where the threshold is located.
threshold.linetype	
	The line shape of the threshold line.

threshold.linewidth

The line width of the threshold line.

threshold.linecol

The colour of the threshold line.

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liftpec Threshold point left displacement.
rightpec Threshold point right displacement.

legend.position

Set the position of the legend.

Splitface Name the faceted image.

Value

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