

Package: **condvis2** (via **r-universe**)

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Title Interactive Conditional Visualization for Supervised and Unsupervised Models in Shiny

Version 0.1.2

Description Constructs a shiny app function with interactive displays for conditional visualization of models, data and density functions. An extended version of package 'condvis'. Catherine B. Hurley, Mark O'Connell, Katarina Domijan (2021) <10.1080/10618600.2021.1983439>.

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Encoding UTF-8

LazyData true

Imports shiny, RColorBrewer, ggplot2, scales, cluster, DendSer, methods, plyr, colorspace, gower

RoxygenNote 7.2.1

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VignetteBuilder knitr

BugReports <https://github.com/cbhurley/condvis2/issues>

URL <https://cbhurley.github.io/condvis2/>,
<https://github.com/cbhurley/condvis2>

Repository <https://cbhurley.r-universe.dev>

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clusPath	<i>Constructs tours of data space based on centers of clusters</i>
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Description

Constructs tours of data space based on centers of clusters

Usage

```
kmeansPath(data, length = 10, reorder = TRUE, conditionvars = NULL, ...)
```

```
pamPath(
  data,
  length = 10,
  reorder = TRUE,
  conditionvars = NULL,
  maxn = 4000,
  ...
)
```

```
claraPath(data, length = 10, reorder = TRUE, conditionvars = NULL, ...)
```

```
medoidPath(data, cl, reorder = FALSE)
```

```
centroidPath(data, cl, reorder = FALSE)
```

Arguments

data	A dataframe
length	Path length, defaults to 10
reorder	If TRUE uses DendSer to reorder the path dser

conditionvars A vector of variable names. The returned tour is for this subset of variables.
 ... ignored
 maxn (pamPath only) For datasets with more than maxn rows, use maxn randomly selected rows.
 cl A vector specifying cluster membership for rows of data.

Value

A dataframe with the path

Functions

- kmeansPath(): Constructs a tour of data space following length k-means centroids
- pamPath(): Constructs a tour of data space following length pam medoids
- claraPath(): Constructs a tour of data space following length clara medoids
- medoidPath(): Returns a path visiting cluster medoids
- centroidPath(): Returns a path visiting cluster centroids

Examples

```

kmeansPath(mtcars,length=4)
pamPath(mtcars,length=4)
claraPath(mtcars,length=4)
medoidPath(mtcars,cl=rep(1:3, length.out=nrow(mtcars)))
  
```

conditionPlot	<i>Plots a conditionPlot.</i>
---------------	-------------------------------

Description

Plots a conditionPlot, showing one, two or many predictors. The predictor setting in varVal is drawn in magenta.

Usage

```

conditionPlot(
  CVdata,
  var,
  varVal,
  pointColor = "steelblue",
  sim = NULL,
  resetpar = TRUE,
  plotrows = NULL
)
  
```

Arguments

CVdata	the dataset used for the fit
var	one more condition vars. Draws a parallel coordinate plot for more than two.
varVal	the current setting of the conditionvars, shown in magenta.
pointColor	a color, vector of colors, or the name of variable to be used for coloring
sim	If non-NULL should be a vector of similarity weights.
resetpar	For use withing shiny app.
plotrows	If non-NULL should be a vector of case indices

Examples

```
conditionPlot(mtcars, c("wt","hp"), c("wt"=3, "hp"=200), pointColor="am")

conditionPlot(mtcars, c("wt","hp"), mtcars[1,], pointColor="am")

#Calculate similarity using wt, hp observations from first case
sim <- similarityweight(mtcars[1, c("wt","hp")], mtcars[, c("wt","hp")], threshold=1)

# Marks points with black border with positive sim values. These are points within 1 (threshold) sd
#of pink cross.
conditionPlot(mtcars, c("wt","hp"), mtcars[1,], pointColor="am", sim=sim)

sim <- similarityweight(mtcars[1, ], mtcars, threshold=2)
conditionPlot(mtcars, names(mtcars), mtcars[1,], sim=sim)
```

condvis

Creates Condvis Shiny app

Description

Creates Condvis Shiny app

Usage

```
condvis(
  data,
  model = NULL,
  response = NULL,
  sectionvars = NULL,
  conditionvars = NULL,
  predsInit = NULL,
  pointColor = c("steelblue", "grey0"),
  cPlotPCP = FALSE,
  cPlotn = 1000,
  orderConditionVars = "default",
```

```

    threshold = 1,
    thresholdmax = NULL,
    linecols = NULL,
    showsim = NULL,
    theta3d = 45,
    phi3d = 20,
    dataplot = "pcp",
    tours = NULL,
    predictArgs = NULL,
    xlim = NULL,
    ylim = NULL,
    zlim = NULL,
    density = FALSE,
    showdata = density == FALSE,
    displayHeight = 950
  )

```

Arguments

<code>data</code>	the dataset used for the fit. Should not have NAs for response, sectionvars or conditionvars.
<code>model</code>	A fitted model or list of models. May be NULL.
<code>response</code>	name of response variable. If null, tries to extract from model.
<code>sectionvars</code>	names of sectionvars. If null, extracts from data.
<code>conditionvars</code>	names of condition vars. If null, extracts from data.
<code>predsInit</code>	Optionally provide starting value for some predictors. Defaults to medoid.
<code>pointColor</code>	a color, or the name of variable to be used for coloring. If the named variable is numeric, it is first converted to a factor with 3 levels.
<code>cPlotPCP</code>	if TRUE, conditionplots are drawn as a single PCP (for more than two conditionvars)
<code>cPlotn</code>	Defaults to 1000. Shows a sample of this number of points in conditionplots. Non-numeric values are ignored.
<code>orderConditionVars</code>	If supplied, a function to order the Condition Vars
<code>threshold</code>	used for similarity weights, defaults to 1.
<code>thresholdmax</code>	maximum value allowed of threshold.
<code>linecols</code>	vector of colors to be used for fits
<code>showsim</code>	if TRUE, shows sim in conditionplots with points/lines. Defaults to TRUE with 150 or fewer cases.
<code>theta3d, phi3d</code>	Angles defining the viewing direction for 3d surface. <code>theta3d</code> gives the azimuthal direction and <code>phi3d</code> the colatitude. See persp .
<code>dataplot</code>	"pcp" or "pairs". Used when there is no response, or more than two sectionvars.
<code>tours</code>	A list of pre-calculated tours
<code>predictArgs</code>	a list with one entry per fit, giving arguments for CVpredict

xlim	passed on to sectionplot
ylim	passed on to sectionplot
zlim	passed on to sectionplot
density	default FALSE. Use TRUE if model is a density function.
showdata	defaults to density==TRUE. If FALSE, data on section not shown.
displayHeight	supply a value for the display height

Examples

```
fit <- lm(mpg ~ wt+hp+am, data=mtcars)
if(interactive()){
  condvis(mtcars,fit, response="mpg",sectionvars="wt", conditionvars=c("am", "hp"), pointColor="red")
}
```

CVpredict

A predict generic function for condvis

Description

A predict generic function for condvis

Usage

```
CVpredict(
  fit,
  newdata,
  ...,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL,
  pinterval = NULL,
  pinterval_level = 0.95
)
```

Default S3 method:

```
CVpredict(
  fit,
  newdata,
  ...,
  ptype = "pred",
  pthreshold = NULL,
  pinterval = NULL,
  pinterval_level = 0.95,
  ylevels = NULL,
  ptrans = NULL
)
```

```
)

## S3 method for class 'lm'
CVpredict(
  fit,
  newdata,
  ...,
  ptype = "pred",
  pthreshold = NULL,
  pinterval = NULL,
  pinterval_level = 0.95,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'glm'
CVpredict(
  fit,
  ...,
  type = "response",
  ptype = "pred",
  pthreshold = NULL,
  pinterval = NULL,
  pinterval_level = 0.95,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'lda'
CVpredict(
  fit,
  ...,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'qda'
CVpredict(
  fit,
  ...,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)
```

```
## S3 method for class 'nnet'
CVpredict(
  fit,
  ...,
  type = NULL,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'randomForest'
CVpredict(
  fit,
  ...,
  type = NULL,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'ranger'
CVpredict(
  fit,
  ...,
  type = NULL,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'rpart'
CVpredict(
  fit,
  ...,
  type = NULL,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'tree'
CVpredict(
  fit,
  ...,
```



```
    type = NULL,
    ptype = "pred",
    pthreshold = NULL,
    ylevels = NULL,
    ptrans = NULL
  )

## S3 method for class 'C5.0'
CVpredict(
  fit,
  ...,
  type = NULL,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'svm'
CVpredict(
  fit,
  ...,
  type = NULL,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'gbm'
CVpredict(
  fit,
  ...,
  type = NULL,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  n.trees = fit$n.trees,
  ptrans = NULL
)

## S3 method for class 'loess'
CVpredict(fit, newdata = NULL, ...)

## S3 method for class 'ksvm'
CVpredict(
  fit,
  newdata,
```

```
    ...,
    type = NULL,
    ptype = "pred",
    pthreshold = NULL,
    ylevels = NULL,
    ptrans = NULL
  )

## S3 method for class 'glmnet'
CVpredict(
  fit,
  newdata,
  ...,
  type = "response",
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL,
  s = NULL,
  makex = NULL
)

## S3 method for class 'cv.glmnet'
CVpredict(
  fit,
  newdata,
  ...,
  type = "response",
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL,
  makex = NULL
)

## S3 method for class 'glmnet.formula'
CVpredict(
  fit,
  newdata,
  ...,
  type = "response",
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL,
  s = NULL
)
```

```
## S3 method for class 'cv.glmnet.formula'
CVpredict(
  fit,
  newdata,
  ...,
  type = "response",
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'keras.engine.training.Model'
CVpredict(
  fit,
  newdata,
  ...,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL,
  batch_size = 32,
  response = NULL,
  predictors = NULL
)

## S3 method for class 'kde'
CVpredict(fit, newdata = fit$x, ..., scale = TRUE)

## S3 method for class 'densityMclust'
CVpredict(
  fit,
  newdata = NULL,
  ...,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL,
  scale = TRUE
)

## S3 method for class 'MclustDA'
CVpredict(
  fit,
  newdata,
  ...,
  ptype = "pred",
  pthreshold = NULL,
```

```
    ylevels = NULL,  
    ptrans = NULL  
  )  
  
## S3 method for class 'MclustDR'  
CVpredict(  
  fit,  
  newdata,  
  ...,  
  ptype = "pred",  
  pthreshold = NULL,  
  ylevels = NULL,  
  ptrans = NULL  
)  
  
## S3 method for class 'Mclust'  
CVpredict(  
  fit,  
  newdata,  
  ...,  
  ptype = "pred",  
  pthreshold = NULL,  
  ylevels = NULL,  
  ptrans = NULL  
)  
  
## S3 method for class 'train'  
CVpredict(  
  fit,  
  newdata,  
  ...,  
  type = "response",  
  ptype = "pred",  
  pthreshold = NULL,  
  ylevels = NULL,  
  ptrans = NULL  
)  
  
## S3 method for class 'bartMachine'  
CVpredict(  
  fit,  
  newdata,  
  ...,  
  type = NULL,  
  ptype = "pred",  
  pthreshold = NULL,  
  ylevels = NULL,  
  ptrans = NULL
```

```
)

## S3 method for class 'wbart'
CVpredict(
  fit,
  newdata,
  ...,
  type = NULL,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'lbart'
CVpredict(
  fit,
  newdata,
  ...,
  type = NULL,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'pbart'
CVpredict(
  fit,
  newdata,
  ...,
  type = NULL,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)

## S3 method for class 'bart'
CVpredict(
  fit,
  newdata,
  ...,
  type = NULL,
  ptype = "pred",
  pthreshold = NULL,
  ylevels = NULL,
  ptrans = NULL
)
```

```
)  
  
## S3 method for class 'model_fit'  
CVpredict(  
  fit,  
  ...,  
  type = NULL,  
  ptype = "pred",  
  pthreshold = NULL,  
  ylevels = NULL,  
  ptrans = NULL,  
  pinterval = NULL,  
  pinterval_level = 0.95  
)  
  
## S3 method for class 'WrappedModel'  
CVpredict(  
  fit,  
  newdata,  
  ...,  
  type = NULL,  
  ptype = "pred",  
  pthreshold = NULL,  
  ylevels = NULL,  
  ptrans = NULL,  
  pinterval = NULL,  
  pinterval_level = 0.95  
)  
  
## S3 method for class 'Learner'  
CVpredict(  
  fit,  
  newdata,  
  ...,  
  type = NULL,  
  ptype = "pred",  
  pthreshold = NULL,  
  ylevels = NULL,  
  ptrans = NULL,  
  pinterval = NULL,  
  pinterval_level = 0.95  
)
```

Arguments

<code>fit</code>	A fitted model
<code>newdata</code>	Where to calculate predictions.
<code>...</code>	extra arguments to predict

<code>ptype</code>	One of "pred", "prob" or "probmatrix"
<code>pthreshold</code>	Used for calculating classes from probs, in the two class case
<code>ylevels</code>	The levels of the response, when it is a factor
<code>ptrans</code>	A function to apply to the result
<code>pinterval</code>	NULL, "confidence" or "prediction". Only for <code>lm</code> , <code>parsnip</code> , <code>mlr</code> (regression, confidence only)
<code>pinterval_level</code>	Defaults to 0.95
<code>type</code>	For some predict methods
<code>n.trees</code>	Used by <code>CVpredict.gbm</code> , passed to <code>predict</code>
<code>s</code>	Used by <code>CVpredict.glmnet</code> and <code>CVpredict.cv.glmnet</code> , passed to <code>predict</code>
<code>makex</code>	Used by <code>CVpredict.glmnet</code> and <code>CVpredict.cv.glmnet</code> . A function to construct <code>xmatrix</code> for <code>predict</code> .
<code>batch_size</code>	Used by <code>CVpredict.keras.engine.training.Model</code> , passed to <code>predict</code>
<code>response</code>	Used by <code>CVpredict.keras.engine.training.Model</code> . Name of response (optional)
<code>predictors</code>	Used by <code>CVpredict.keras.engine.training.Model</code> . Name of predictors
<code>scale</code>	Used by <code>CVpredict</code> for densities. If TRUE (default) rescales the conditional density to integrate to 1.

Details

This is a wrapper for `predict` used by `condvis`. When the model response is numeric, the result is a vector of predictions. When the model response is a factor the result depends on the value of `ptype`. If `ptype="pred"`, the result is a factor. If also `threshold` is numeric, it is used to threshold a numeric prediction to construct the factor when the factor has two levels. For `ptype="prob"`, the result is a vector of probabilities for the last factor level. For `ptype="probmatrix"`, the result is a matrix of probabilities for each factor level.

Value

a vector of predictions, or a matrix when `type` is "probmatrix"

Methods (by class)

- `CVpredict(default)`: `CVpredict` method
- `CVpredict(lm)`: `CVpredict` method
- `CVpredict(glm)`: `CVpredict` method
- `CVpredict(lda)`: `CVpredict` method
- `CVpredict(qda)`: `CVpredict` method
- `CVpredict(nnet)`: `CVpredict` method
- `CVpredict(randomForest)`: `CVpredict` method
- `CVpredict(ranger)`: `CVpredict` method
- `CVpredict(rpart)`: `CVpredict` method

- CVpredict(tree): CVpredict method
- CVpredict(C5.0): CVpredict method
- CVpredict(svm): CVpredict method
- CVpredict(gbm): CVpredict method
- CVpredict(loess): CVpredict method
- CVpredict(ksvm): CVpredict method
- CVpredict(glmnet): CVpredict method
- CVpredict(cv.glmnet): CVpredict method
- CVpredict(glmnet.formula): CVpredict method
- CVpredict(cv.glmnet.formula): CVpredict method
- CVpredict(keras.engine.training.Model): CVpredict method
- CVpredict(kde): CVpredict method
- CVpredict(densityMclust): CVpredict method
- CVpredict(MclustDA): CVpredict method
- CVpredict(MclustDR): CVpredict method
- CVpredict(Mclust): CVpredict method
- CVpredict(train): CVpredict method for caret
- CVpredict(bartMachine): CVpredict method
- CVpredict(wbart): CVpredict method
- CVpredict(lbart): CVpredict method
- CVpredict(pbart): CVpredict method
- CVpredict(bart): CVpredict method
- CVpredict(model_fit): CVpredict method for parsnip
- CVpredict(WrappedModel): CVpredict method for mlr
- CVpredict(Learner): CVpredict method for mlr3

Examples

```
#Fit a model.
f <- lm(Fertility~ ., data=swiss)
CVpredict(f)

#Fit a model with a factor response
swiss1 <- swiss
swiss1$Fertility <- cut(swiss$Fertility, c(0,80,100))
levels(swiss1$Fertility)<- c("lo", "hi")
f <- glm(Fertility~ ., data=swiss1, family="binomial")
CVpredict(f) # by default gives a factor
CVpredict(f, ptype="prob") # gives prob of level hi
CVpredict(f, ptype="probmatrix") # gives prob of both levels
```

fitPath	<i>Constructs tours of data space based on fits.</i>
---------	--

Description

Constructs tours of data space based on fits.

Usage

```
lofPath(  
  data,  
  fits,  
  length = 10,  
  reorder = TRUE,  
  conditionvars = NULL,  
  predictArgs = NULL,  
  response = NULL,  
  ...  
)
```

```
diffitsPath(  
  data,  
  fits,  
  length = 10,  
  reorder = TRUE,  
  conditionvars = NULL,  
  predictArgs = NULL,  
  ...  
)
```

```
hiresponsePath(  
  data,  
  response = NULL,  
  length = 10,  
  reorder = TRUE,  
  conditionvars = NULL,  
  ...  
)
```

```
loresponsePath(  
  data,  
  response = NULL,  
  length = 10,  
  reorder = TRUE,  
  conditionvars = NULL,  
  ...  
)
```

Arguments

data	A dataframe
fits	A model fit or list of fits
length	Path length, defaults to 10
reorder	If TRUE (default) uses DendSer to reorder the path dser
conditionvars	A vector of variable names. The returned tour is for this subset of variables.
predictArgs	Extra inputs to CVpredict
response	The name of the response variable
...	ignored

Value

A dataframe with the path

Functions

- `lofPath()`: Constructs a tour of data space showing biggest absolute residuals from fits.
- `diffitsPath()`: Constructs a tour of data space showing biggest differences in fits.
- `hiresponsePath()`: Constructs a tour of data space showing high (numeric) response values
- `loresponsePath()`: Constructs a tour of data space showing low (numeric) response values

Examples

```
fit1 <- lm(mpg ~ wt+hp+am, data=mtcars)
fit2 <- lm(mpg ~ wt, data=mtcars)
lofPath(mtcars,fit1, response="mpg")
diffitsPath(mtcars,list(fit1,fit2))
```

medoid

Finds medoid of data

Description

Finds medoid of data

Usage

```
medoid(data, maxn = 4000)
```

Arguments

data	A dataframe
maxn	For datasets with more than maxn rows, use maxn randomly selected rows.

Value

A dataframe with one row, which is the medoid of the data, based on (standardised) daisy dist

Examples

```
medoid(mtcars)
```

pathInterpolate	<i>Interpolation</i>
-----------------	----------------------

Description

Interpolation

Usage

```
pathInterpolate(x, ninterp = 4)

## Default S3 method:
pathInterpolate(x, ninterp = 4L)

## S3 method for class 'factor'
pathInterpolate(x, ninterp = 4L)

## S3 method for class 'data.frame'
pathInterpolate(x, ninterp = 4L)
```

Arguments

x	a numeric or factor vector or dataframe
ninterp	number of interpolated steps

Value

interpolated version of x

Methods (by class)

- pathInterpolate(default): Default interpolate method
- pathInterpolate(factor): pathInterpolate method for factor
- pathInterpolate(data.frame): pathInterpolate method for data.frame

plotTourDiagnostics *Plots diagnostics for the tour supplied*

Description

Plots diagnostics for the tour supplied

Usage

```
plotTourDiagnostics(  
  path,  
  data,  
  pathlen = nrow(path),  
  threshold = 1,  
  which = 1:3,  
  ...  
)
```

Arguments

path	the tour
data	the dataset
pathlen	the pathlength
threshold	used for similarityweight
which	subset of 1:3
...	other args for similarityweight

Details

The first plot shows approximately how much data are visible on each section, the second shows what proportion of data are *visited* by the tour, and the third a density estimate of max similarity values.

Value

Table of max sims attained.

sectionPlot	<i>Plots the main condvis display</i>
-------------	---------------------------------------

Description

The section plot relates a fit or fits to one or two predictors (sectionvar), for fixed values of other predictors in conditionvals.

Usage

```
sectionPlot(
  CVdata,
  CVfit = NULL,
  response = NULL,
  preds,
  sectionvar,
  conditionvals,
  pointColor = "steelblue",
  sim = NULL,
  threshold = 1,
  linecols = NULL,
  dataplot = "pcp",
  gridsize = 50,
  probs = FALSE,
  view3d = FALSE,
  theta3d = 45,
  phi3d = 20,
  xlim = NULL,
  ylim = NULL,
  zlim = NULL,
  pointSize = 1.5,
  predictArgs = NULL,
  resetpar = TRUE,
  density = FALSE,
  showdata = density == FALSE,
  returnInfo = FALSE,
  pointColorFromResponse = FALSE,
  pcolInfo = NULL
)
```

Arguments

CVdata	the dataset used for the fit
CVfit	a fit or list of fits
response	name of response variable
preds	names of predictors

sectionvar	section variable, or variables.
conditionvals	conditioning values. A vector/list or dataframe with one row
pointColor	a color, vector of colors, or the name of variable to be used for coloring
sim	vector of similarity weights
threshold	used for similarity weights, defaults to 1.
linecols	vector of line colours
dataplot	"pcp" or "pairs". Used when there is no response, or more than two sectionvars.
gridsize	used to construct grid of fitted values.
probs	Logical; if TRUE, shows predicted class probabilities instead of just predicted classes. Only available with two numeric sectionvars and the model's predict method provides this.
view3d	Logical; if TRUE plots a three-dimensional regression surface if possible.
theta3d, phi3d	Angles defining the viewing direction. theta3d gives the azimuthal direction and phi3d the colatitude. See persp .
xlim	passed on to plot
ylim	passed on to plot
zlim	passed on to plot
pointSize	used for points
predictArgs	a list with one entry per fit, giving arguments for predict
resetpar	When TRUE (the default) resets pars after drawing.
density	default FALSE. Use TRUE if model is a density function.
showdata	If FALSE, data on section not shown.
returnInfo	If TRUE, returns coordinates for some plots
pointColorFromResponse	ignore—For interactive use only
pcolInfo	ignore—For interactive use only

Details

The type of plot depends on the fit and the section variables. Observations with non zero values of the similarity weights `sim` are shown. If no fit is provided, the data are shown as a parallel coordinate plot or pairs plot, depending on `dataplot`. The fit could also be a density estimate.

Value

plotted coordinates, for some plots

Examples

```

#Fit a model.
f <- lm(Fertility~ ., data=swiss)
svar <- "Education"
preds <- variable.names(f)[-1]
sectionPlot(swiss,f, "Fertility",preds,svar, swiss[12,])
sectionPlot(swiss,f, "Fertility",preds,svar, apply(swiss,2,median))
sectionPlot(swiss,f, "Fertility",preds,preds[1:2], apply(swiss,2,median))
sectionPlot(swiss,f, "Fertility",preds,preds[1:2], apply(swiss,2,median), view3d=TRUE)

# PCP of swiss data, showing only cases whose percent catholic and infant.mortality are
# similar to those of the first case
sectionPlot(swiss,preds=names(swiss),
            sectionvar= names(swiss)[1:4],conditionvals=swiss[1,] )
# Use dataplot="pairs" to switch to a pairs plot

# A density estimate example
## Not run:
library(ks)
fde <-kde(iris[,1:3])
sectionPlot(iris,list(kde=fde), response=NULL,
            preds=names(iris)[1:3],
            sectionvar=names(iris)[1],
            conditionvals=iris[1,],density=TRUE)

## End(Not run)

```

similarityweight

Calculate the similarity weight for a set of observations

Description

Calculate the similarity weight for a set of observations, based on their distance from some arbitrary points in data space. Observations which are very similar to the point under consideration are given weight 1, while observations which are dissimilar to the point are given weight zero.

Usage

```

similarityweight(
  x,
  data,
  threshold = 1,
  distance = "euclidean",
  lambda = NULL,
  scale = TRUE
)

```

Arguments

x	A dataframe describing arbitrary points in the space of the data (i.e., with same colnames as data).
data	A dataframe representing observed data.
threshold	Threshold distance outside which observations will be assigned similarity weight zero. This is numeric and should be > 0. Defaults to 1.
distance	The type of distance measure to be used, currently just three types of Minkowski distance: "euclidean" (default), "maxnorm", "manhattan" and also "gower"
lambda	A constant to multiply by the number of categorical mismatches, before adding to the Minkowski distance, to give a general dissimilarity measure. If left NULL, behaves as though lambda is set larger than threshold, meaning that one factor mismatch guarantees zero weight.
scale	defaults to TRUE, in which case numeric variables are scaled to unit sd.

Details

Similarity weight is assigned to observations based on their distance from a given point. The distance is calculated as Minkowski distance between the numeric elements for the observations whose categorical elements match, or else the Gower distance.

Value

A numeric vector or matrix, with values from 0 to 1. The similarity weights for the observations in data arranged in rows for each row in x.

References

O'Connell M, Hurley CB and Domijan K (2017). "Conditional Visualization for Statistical Models: An Introduction to the **condvis** Package in R." *Journal of Statistical Software*, **81**(5), pp. 1-20. <URL:<http://dx.doi.org/10.18637/jss.v081.i05>>.

Examples

```
## Say we want to find observations similar to the first observation.
## The first observation is identical to itself, so it gets weight 1. The
## second observation is similar, so it gets some weight. The rest are more
## different, and so get zero weight.

data(mtcars)
similarityweight(x = mtcars[1, ], data = mtcars)

## By increasing the threshold, we can find observations which are more
## approximately similar to the first row. Note that the second observation
## now has weight 1, so we lose some ability to discern how similar
## observations are by increasing the threshold.

similarityweight(x = mtcars[1, ], data = mtcars, threshold = 5)

## Can provide a number of points to 'x'. Here we see that the Mazda RX4 Wag
```



```
## is more similar to the Merc 280 than the Mazda RX4 is.
similarityweight(mtcars[1:2, ], mtcars, threshold = 3)
```

tours	<i>Constructs a various tours of data space</i>
-------	---

Description

Constructs a various tours of data space

Usage

```
randomPath(data, length = 10, reorder = TRUE, conditionvars = NULL, ...)
```

```
seqPath(data, length = 10, reorder = FALSE, conditionvars = NULL, ...)
```

```
alongPath(data, var, length = 10, current = NULL, ...)
```

Arguments

data	A dataframe
length	Path length, defaults to 10
reorder	If TRUE (default) uses DendSer to reorder the path dser
conditionvars	A vector of variable names. The returned tour is for this subset of variables.
...	ignored
var	A variable name for alongPath
current	Default value of variables for alongPath

Value

A dataframe with the path

Functions

- `randomPath()`: Constructs a tour of data space following random observations
- `seqPath()`: Constructs a tour of data space following first length observations
- `alongPath()`: Constructs a tour of data space of length equi-spaced values in the range of var. If var is a factor, its levels are used.

Examples

```
randomPath(mtcars,length=5)
seqPath(mtcars,length=5)
alongPath(mtcars,var="mpg", length=5, current=mtcars[1,])
```

weightcolor	<i>Fade colours according to a weight vector</i>
-------------	--

Description

The colours whose weights are less than 1 are diluted. Colours whose weight is zero are returned as white, other weights are grouped in `nlevels` groups and colours diluted proportionally.

Usage

```
weightcolor(col, weights, nlevels = 5)
```

Arguments

<code>col</code>	A vector of colour
<code>weights</code>	A vector of weights, values between 0 and 1
<code>nlevels</code>	The number of groups

Value

A vector of colours

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