

# Package ‘spm2’

March 10, 2022

**Title** Spatial Predictive Modeling

**Version** 1.1.2

**Date** 2022-03-10

**Description** An updated and extended version of 'spm' package, by introducing some further novel functions for modern statistical methods (i.e., generalised linear models, glmnet, generalised least squares), thin plate splines, support vector machine, kriging methods (i.e., simple kriging, universal kriging, block kriging, kriging with an external drift), and novel hybrid methods (228 hybrids plus numerous variants) of modern statistical methods or machine learning methods with mathematical and/or univariate geostatistical methods for spatial predictive modelling. For each method, two functions are provided, with one function for assessing the predictive errors and accuracy of the method based on cross-validation, and the other for generating spatial predictions. It also contains a couple of functions for data preparation and predictive accuracy assessment.

**Depends** R (>= 2.10)

**Imports** spm, gstat, sp, randomForest, psy, gbm, stats, ranger, MASS, fields, nlme, glmnet, e1071

**License** GPL (>= 2)

**LazyData** true

**RoxygenNote** 7.1.1

**Encoding** UTF-8

**Suggests** knitr, rmarkdown

**NeedsCompilation** no

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**Repository** CRAN

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bees	<i>A dataset of bees count data and relevant information in oilseed Brassica fields in an Australian temperate landscape.</i>
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### Description

This dataset contains 212 samples of 61 variables including three bee species, inflorescence, temperature, wind speed and various derived landscape variables.

**Usage**

```
data("bees")
```

**Format**

A data frame with 212 observations on the following 61 variables.

**transid** a factor with levels G1 G10 G11 G12 G13 G14 G15 G16 G1D1 G1D10 G1D11 G1D12 G1D13 G1D14 G1D15 G1D16 G1D17 G1D2 G1D3 G1D4 G1D5 G1D6 G1D7 G1D8 G1D9 G1G1 G1G10 G1G11 G1G12 G1G13 G1G14 G1G15 G1G16 G1G17 G1G2 G1G3 G1G4 G1G5 G1G6 G1G7 G1G8 G1G9 G2 G2H1 G2H10 G2H11 G2H2 G2H3 G2H4 G2H5 G2H6 G2H7 G2H8 G2H9 G2S1 G2S10 G2S11 G2S2 G2S3 G2S4 G2S5 G2S6 G2S7 G2S8 G2S9 G3 G4 G5 G6 G7 G8 G9 GCH1 GCH10 GCH11 GCH12 GCH13 GCH14 GCH15 GCH16 GCH17 GCH18 GCH2 GCH3 GCH4 GCH5 GCH6 GCH7 GCH8 GCH9 GCS1 GCS10 GCS11 GCS12 GCS13 GCS14 GCS15 GCS16 GCS17 GCS18 GCS2 GCS3 GCS4 GCS5 GCS6 GCS7 GCS8 GCS9 H1 H10 H11 H12 H13 H14 H15 H16 H2 H3 H4 H5 H6 H7 H8 H9 MC1G1 MC1G2 MC1G3 MC1G4 MC1G5 MC1G6 MC1G7 MC1G8 MC1G9 MC1H1 MC1H2 MC1H3 MC1H4 MC1H5 MC1H6 MC1H7 MC1H8 MC1H9 MC2AH1 MC2AH10 MC2AH11 MC2AH12 MC2AH13 MC2AH14 MC2AH15 MC2AH16 MC2AH17 MC2AH18 MC2AH2 MC2AH3 MC2AH4 MC2AH5 MC2AH6 MC2AH7 MC2AH8 MC2AH9 MC2AS1 MC2AS10 MC2AS11 MC2AS12 MC2AS13 MC2AS14 MC2AS15 MC2AS16 MC2AS17 MC2AS18 MC2AS2 MC2AS3 MC2AS4 MC2AS5 MC2AS6 MC2AS7 MC2AS8 MC2AS9 MC2BB1 MC2BB10 MC2BB11 MC2BB12 MC2BB13 MC2BB14 MC2BB15 MC2BB16 MC2BB17 MC2BB2 MC2BB3 MC2BB4 MC2BB5 MC2BB6 MC2BB7 MC2BB8 MC2BB9 MC2BG1 MC2BG10 MC2BG11 MC2BG12 MC2BG13 MC2BG14 MC2BG15 MC2BG16 MC2BG17 MC2BG2 MC2BG3 MC2BG4 MC2BG5 MC2BG6 MC2BG7 MC2BG8 MC2BG9

**transsurv** a numeric vector

**plotsurv** a numeric vector

**paddock** a numeric vector

**plot** a factor with levels G1-1 G1-10 G1-11 G1-12 G1-13 G1-14 G1-15 G1-16 G1-17 G1-2 G1-3 G1-4 G1-5 G1-6 G1-7 G1-8 G1-9 G21 G210 G211 G22 G23 G24 G25 G26 G27 G28 G29 GC1 GC10 GC11 GC12 GC13 GC14 GC15 GC16 GC17 GC18 GC2 GC3 GC4 GC5 GC6 GC7 GC8 GC9 MC-1 MC-10 MC-11 MC-12 MC-13 MC-14 MC-15 MC-16 MC-2 MC-3 MC-4 MC-5 MC-6 MC-7 MC-8 MC-9 MC1-1 MC1-2 MC1-3 MC1-4 MC1-5 MC1-6 MC1-7 MC1-8 MC1-9 MC2-A1 MC2-A10 MC2-A11 MC2-A12 MC2-A13 MC2-A14 MC2-A15 MC2-A16 MC2-A17 MC2-A18 MC2-A2 MC2-A3 MC2-A4 MC2-A5 MC2-A6 MC2-A7 MC2-A8 MC2-A9 MC2-B1 MC2-B10 MC2-B11 MC2-B12 MC2-B13 MC2-B14 MC2-B15 MC2-B16 MC2-B17 MC2-B2 MC2-B3 MC2-B4 MC2-B5 MC2-B6 MC2-B7 MC2-B8 MC2-B9

**obs** a factor with levels Andrew Barbara Micah Sonia Steve

**hbee** a numeric vector

**nbee** a numeric vector

**hover** a numeric vector

**date** a numeric vector

**sx** a numeric vector

**fx** a numeric vector

**sy** a numeric vector

**fy** a numeric vector

**loc** a factor with levels 100 150 200 400 450 edge

pair a factor with levels I 0  
inf a numeric vector  
rankinf a numeric vector  
dupl a numeric vector  
temp a numeric vector  
windspeed a numeric vector  
winddir a factor with levels N NE NNE NNW NW  
cloudc a numeric vector  
disttoedgecal a numeric vector  
disttoedgemeasured a numeric vector  
w100 a numeric vector  
w200 a numeric vector  
w300 a numeric vector  
w400 a numeric vector  
w500 a numeric vector  
w600 a numeric vector  
w700 a numeric vector  
w800 a numeric vector  
w900 a numeric vector  
w1000 a numeric vector  
w1500 a numeric vector  
w2000 a numeric vector  
c100 a numeric vector  
c200 a numeric vector  
c300 a numeric vector  
c400 a numeric vector  
c500 a numeric vector  
c1000 a numeric vector  
c1500 a numeric vector  
area a numeric vector  
perimeter a numeric vector  
gyration a numeric vector  
paratio a numeric vector  
shape a numeric vector  
fractaldimention a numeric vector  
circumscircle a numeric vector  
contiguity a numeric vector

links100 a numeric vector  
links200 a numeric vector  
links300 a numeric vector  
links400 a numeric vector  
links500 a numeric vector  
links1000 a numeric vector  
links1500 a numeric vector  
links2000 a numeric vector  
windspeed2 a numeric vector

### Details

For details, please see the source. This dataset was published as an appendix of the paper listed in the source. Where the long and lat were reprojected to easting and northing

### Source

The data source is [<https://doi.org/10.25919/5f17b34638cca>] or [<https://data.csiro.au/collections/collection/CIcsiro:45533>], which provides bees count data and relevant predictive variables along with a brief description of the data. The detailed descriptions of the data are available in: "Arthur, A. D., et al. (2010). "Influence of woody vegetation on pollinator densities in oilseed Brassica fields in an Australian temperate landscape." *Basic and Applied Ecology* 11: 406-414."

### References

Arthur, A. D., Li, J., Henry, S., Cunningham, S.A. (2010). "Influence of woody vegetation on pollinator densities in oilseed Brassica fields in an Australian temperate landscape." *Basic and Applied Ecology* 11: 406-414.

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ccr	<i>Correct classification rate for predictive models based on cross - validation</i>
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### Description

This function is to calculate correct classification rate (ccr) for categorical data with the observed (obs) data specified as factor. It is based on the differences between the predicted values for and the observed values of validation samples for cross-validation. For 0 and 1 data, the observed values need to be specified as factor in order to use this accuracy measure. It is modified from the function 'pred.acc' in 'spm' package.

### Usage

```
ccr(obs, pred)
```

**Arguments**

obs            a vector of observation values of validation samples.  
pred           a vector of prediction values of predictive models for validation samples.

**Value**

A list with the following component: ccr (correct classification rate) for categorical data.

**Author(s)**

Jin Li

**References**

Jin Li (2019). spm: Spatial Predictive Modeling. R package version 1.2.0. <https://CRAN.R-project.org/package=spm>.

**Examples**

```
set.seed(1234)
x <- as.factor(sample(letters[1:2], 30, TRUE))
y <- sample(x, 30)
ccr(x, y)
```

---

cran-comments

*Note on notes*

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**Description**

This is an updated and extended version of 'spm' package. The change in package name from 'spm' to 'spm2' is due to the change in Author's support from Geoscience Australia to Data2Action Australia.

## R CMD check results 0 errors | 0 warnings | 0 notes

**Author(s)**

Jin Li

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`datasplit`*Split data for k-fold cross-validation*

---

**Description**

This function is a data splitting function for k-fold cross-validation and uses a stratified random sampling technique. It resamples the training data based on sample quantiles.

**Usage**

```
datasplit(trainy, k.fold = 10)
```

**Arguments**

<code>trainy</code>	a vector of response, must have a length equal to sample size.
<code>k.fold</code>	integer; number of folds in the cross-validation. if > 1, then apply k-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.

**Value**

A list of samples each with an index of k-fold number.

**Note**

This function is largely based on `rfcv` in `randomForest`.

**Author(s)**

Jin Li

**References**

A. Liaw and M. Wiener (2002). Classification and Regression by `randomForest`. *R News* 2(3), 18-22.

**Examples**

```
library(spm)
data(petrel)
idx1 <- datasplit(petrel[, 3], k.fold = 10)
table(idx1)
```

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decimaldigit	<i>Digit number after decimal point for a numeric variable</i>
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**Description**

This function is to derive the digit number after decimal point for a numeric variable (e.g., lat and long).

**Usage**

```
decimaldigit(x, dechar = ".", nint = NA, ndec = NA, pad.left = TRUE)
```

**Arguments**

x	one or more decimal numbers.
dechar	The character used to separate the decimal part of a number.
nint	The number of characters to which the integer part of the numbers should be padded.
ndec	The number of characters to which the decimal part of the numbers should be padded.
pad.left	Whether the left (integer) side of the numbers should be padded as well as the right.

**Value**

A list of integer number to show digit number after decimal point of x.

**Note**

This function is modified from decimal.align in 'prettyR' package.

**Author(s)**

Jin Li

**References**

Jim Lemon and Philippe Grosjean (2019). 'prettyR': Pretty Descriptive Stats. R package version 2.1.1. <https://CRAN.R-project.org/package=prettyR>.

**Examples**

```
x<-c(0.1, 2.2, 3.03, 44.444, 555.0005, 6666.66666)
decimaldigit(x)
```



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gbmkrigeidwcv	<i>Cross validation, n-fold and leave-one-out for the hybrid methods of generalized boosted regression modeling ('gbm'), 'kriging' and inverse distance weighted ('IDW').</i>
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---

## Description

This function is a cross validation function for 38 hybrid methods of 'gbm', 'kriging' and 'IDW', including the average of 'gbmkrige' and 'gbmidw' ('gbmkrige**gbmidw**') and the average of 'gbm', 'gbmkrige' and 'gbmidw' ('gbm**gbmkrige****gbmidw**'), where 'kriging' methods include ordinary kriging ('OK'), simple kriging ('SK'), block 'OK' ('BOK') and block 'SK' ('BSK') and 'IDW' also covers 'NN' and 'KNN'. The data splitting is based on a stratified random sampling method (see the 'datasplit' function for details).

## Usage

```
gbmkrigeidwcv(
  longlat,
  trainx,
  trainy,
  var.monotone = rep(0, ncol(trainx)),
  family = "gaussian",
  n.trees = 3000,
  learning.rate = 0.001,
  interaction.depth = 2,
  bag.fraction = 0.5,
  train.fraction = 1,
  n.minobsinnode = 10,
  transformation = "none",
  weights = rep(1, nrow(trainx)),
  keep.data = FALSE,
  verbose = TRUE,
  delta = 1,
  formula = res1 ~ 1,
  vgm.args = "Sph",
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  idp = 2,
  nmaxidw = 12,
  hybrid.parameter = 2,
  lambda = 1,
  validation = "CV",
  cv.fold = 10,
  predacc = "VEcv",
```

```

    n.cores = 8,
    ...
  )

```

### Arguments

<code>longlat</code>	a dataframe contains longitude and latitude of point samples.
<code>trainx</code>	a dataframe or matrix contains columns of predictive variables.
<code>trainy</code>	a vector of the response variable.
<code>var.monotone</code>	an optional vector, the same length as the number of predictors, indicating which variables have a monotone increasing (+1), decreasing (-1), or arbitrary (0) relationship with the outcome. By default, a vector of 0 is used.
<code>family</code>	either a character string specifying the name of the distribution to use or a list with a component name specifying the distribution and any additional parameters needed. See <code>gbm</code> for details. By default, "gaussian" is used.
<code>n.trees</code>	the total number of trees to fit. This is equivalent to the number of iterations and the number of basis functions in the additive expansion. By default, 3000 is used.
<code>learning.rate</code>	a shrinkage parameter applied to each tree in the expansion. Also known as step-size reduction.
<code>interaction.depth</code>	the maximum depth of variable interactions. 1 implies an additive model, 2 implies a model with up to 2-way interactions, etc. By default, 2 is used.
<code>bag.fraction</code>	the fraction of the training set observations randomly selected to propose the next tree in the expansion. By default, 0.5 is used.
<code>train.fraction</code>	The first <code>train.fraction * nrow(data)</code> observations are used to fit the <code>gbm</code> and the remainder are used for computing out-of-sample estimates of the loss function.
<code>n.minobsinnode</code>	minimum number of observations in the trees terminal nodes. Note that this is the actual number of observations not the total weight. By default, 10 is used.
<code>transformation</code>	transform the residuals of 'gbm' to normalize the data for 'krige'; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
<code>weights</code>	an optional vector of weights to be used in the fitting process. Must be positive but do not need to be normalized. If <code>keep.data = FALSE</code> in the initial call to <code>gbm</code> then it is the user's responsibility to resupply the weights to <code>gbm.more</code> . By default, a vector of 1 is used.
<code>keep.data</code>	a logical variable indicating whether to keep the data and an index of the data stored with the object. Keeping the data and index makes subsequent calls to <code>gbm.more</code> faster at the cost of storing an extra copy of the dataset. By default, 'FALSE' is used.
<code>verbose</code>	If TRUE, <code>gbm</code> will print out progress and performance indicators. By default, 'TRUE' is used.
<code>delta</code>	numeric; to avoid $\log(0)$ in the log transformation. The default is 1.

formula	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details. The default is 'formula = res1 ~ 1'.
vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
hybrid.parameter	the default is 2 that is for 'gbmkrigeidw'; for 'gbmgbmkrigeidw', it needs to be 3.
lambda,	ranging from 0 to 2; the default is 1 for 'gbmkrigeidw' and 'gbmgbmkrigeidw'; and if it is < 1, more weight is placed on 'krige', otherwise more weight is placed on 'idw'; and if it is 0, 'idw' is not considered and the resultant methods is 'gbmkrige' when the default 'hybrid.parameter' is used; and if it is 2, then the resultant method is 'gbmidw' when the default 'hybrid.parameter' is used.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
n.cores	The number of CPU cores to use. See gbm for details. By default, 8 is used.
...	other arguments passed on to 'randomForest', 'krige' and 'gstat'.

**Value**

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

**Note**

This function is largely based on 'gbm' in 'spm', and 'krige' in 'spm2'.

**Author(s)**

Jin Li

## References

- Li, J., Potter, A., Huang, Z., and Heap, A. (2012). Predicting Seabed Sand Content across the Australian Margin Using Machine Learning and Geostatistical Methods, *Geoscience Australia, Record 2012/48*, 115pp.
- Li, J., Heap, A., Potter, A., and Danilel, J.J. (2011). Predicting Seabed Mud Content across the Australian Margin II: Performance of Machine Learning Methods and Their Combination with Ordinary Kriging and Inverse Distance Squared, *Geoscience Australia, Record 2011/07*, 69pp.
- Greg Ridgeway with contributions from others (2015). *gbm: Generalized Boosted Regression Models*. R package version 2.1.1. <https://CRAN.R-project.org/package=gbm>
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

## Examples

```
library(spm)
# gbmgbmkgbmidw
data(sponge)
longlat <- sponge[, 1:2]
set.seed(1234)
gbmgbmkrigeidwcv1 <- gbmkrigeidwcv(longlat = longlat,
  trainx = sponge[, -3], trainy = sponge[, 3], family = "poisson", interaction.depth = 3,
  transformation = "none", formula = res1 ~ 1, vgm.args = "Sph",
  nmaxkrige = 12, idp = 2, nmaxidw = 12, hybrid.parameter = 3, validation = "CV",
  predacc = "ALL", n.cores = 8)
gbmgbmkrigeidwcv1

# gbmkgbmidw for count data
data(sponge)
longlat <- sponge2[, 1:2]
y = sponge[, 3]
trainx = sponge[, -3]
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  gbmkrigeidwcv1 <- gbmkrigeidwcv(longlat = longlat,
    trainx = trainx, trainy = y, family = "poisson", interaction.depth = 3,
    transformation = "none", formula = res1 ~ 1, vgm.args = "Sph",
    nmaxkrige = 12, idp = 2, nmaxidw = 12, hybrid.parameter = 2, validation = "CV",
    predacc = "VEcv", n.cores = 8)
  VEcv [i] <- gbmkrigeidwcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for gbmkgbmidw", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)
```

---

gbmkrigeidwpred	<i>Generate spatial predictions using the hybrid methods of generalized boosted regression modeling ('gbm'), 'kriging' and inverse distance weighted ('IDW').</i>
-----------------	---

---

## Description

This function is for generating spatial predictions using the hybrid methods of 'gbm', 'kriging' and 'IDW', including all methods implemented in 'gbmkrigeidwcv'.

## Usage

```
gbmkrigeidwpred(  
  longlat,  
  trainx,  
  predx,  
  trainy,  
  longlatpredx,  
  var.monotone = rep(0, ncol(trainx)),  
  family = "gaussian",  
  n.trees = 3000,  
  learning.rate = 0.001,  
  interaction.depth = 2,  
  bag.fraction = 0.5,  
  train.fraction = 1,  
  n.minobsinnode = 10,  
  transformation = "none",  
  weights = rep(1, nrow(trainx)),  
  keep.data = FALSE,  
  verbose = TRUE,  
  delta = 1,  
  formula = res1 ~ 1,  
  vgm.args = "Sph",  
  anis = c(0, 1),  
  alpha = 0,  
  block = 0,  
  beta,  
  nmaxkrige = 12,  
  idp = 2,  
  nmaxidw = 12,  
  hybrid.parameter = 2,  
  lambda = 1,  
  cv.fold = 10,  
  n.cores = 8,  
  ...  
)
```

**Arguments**

<code>longlat</code>	a dataframe contains longitude and latitude of point samples.
<code>trainx</code>	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples.
<code>predx</code>	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
<code>trainy</code>	a vector of the response variable in the formula, that is, the left part of the formula.
<code>longlatpredx</code>	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted.
<code>var.monotone</code>	an optional vector, the same length as the number of predictors, indicating which variables have a monotone increasing (+1), decreasing (-1), or arbitrary (0) relationship with the outcome. By default, a vector of 0 is used.
<code>family</code>	either a character string specifying the name of the distribution to use or a list with a component name specifying the distribution and any additional parameters needed. See <code>gbm</code> for details. By default, "gaussian" is used.
<code>n.trees</code>	the total number of trees to fit. This is equivalent to the number of iterations and the number of basis functions in the additive expansion. By default, 3000 is used.
<code>learning.rate</code>	a shrinkage parameter applied to each tree in the expansion. Also known as step-size reduction.
<code>interaction.depth</code>	the maximum depth of variable interactions. 1 implies an additive model, 2 implies a model with up to 2-way interactions, etc. By default, 2 is used.
<code>bag.fraction</code>	the fraction of the training set observations randomly selected to propose the next tree in the expansion. By default, 0.5 is used.
<code>train.fraction</code>	The first <code>'train.fraction * nrows(data)'</code> observations are used to fit the <code>gbm</code> and the remainder are used for computing out-of-sample estimates of the loss function.
<code>n.minobsinnode</code>	minimum number of observations in the trees terminal nodes. Note that this is the actual number of observations not the total weight. By default, 10 is used.
<code>transformation</code>	transform the residuals of <code>'gbm'</code> to normalise the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
<code>weights</code>	an optional vector of weights to be used in the fitting process. Must be positive but do not need to be normalized. If <code>keep.data = FALSE</code> in the initial call to <code>gbm</code> then it is the user's responsibility to resupply the weights to <code>gbm.more</code> . By default, a vector of 1 is used.
<code>keep.data</code>	a logical variable indicating whether to keep the data and an index of the data stored with the object. Keeping the data and index makes subsequent calls to <code>gbm.more</code> faster at the cost of storing an extra copy of the dataset. By default, 'FALSE' is used.
<code>verbose</code>	If TRUE, <code>gbm</code> will print out progress and performance indicators. By default, 'TRUE' is used.

delta	numeric; to avoid $\log(0)$ in the log transformation. The default is 1.
formula	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details. The default is 'formula = res1 ~ 1'.
vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
hybrid.parameter	the default is 2 that is for 'gbmkrigeidw'; for 'gbmgbmkrigeidw', it needs to be 3.
lambda,	ranging from 0 to 2; the default is 1 for 'gbmkrigeidw' and 'gbmgbmkrigeidw'; and if it is $< 1$ , more weight is placed on 'krige', otherwise more weight is placed on 'idw'; and if it is 0, 'idw' is not considered and the resultant method is 'gbmkrige' when the default 'hybrid.parameter' is used; and if it is 2, then the resultant method is 'gbmidw' when the default 'hybrid.parameter' is used.
cv.fold	integer; number of folds in the cross-validation. if $> 1$ , then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
n.cores	The number of CPU cores to use. See gbm for details. By default, 6 is used.
...	other arguments passed on to 'gbm', 'krige' and 'gstat'.

**Value**

A dataframe of longitude, latitude, and predictions.

**Author(s)**

Jin Li

**References**

Li, J., Potter, A., Huang, Z., and Heap, A. (2012). Predicting Seabed Sand Content across the Australian Margin Using Machine Learning and Geostatistical Methods, Geoscience Australia, Record 2012/48, 115pp.

Li, J., Heap, A., Potter, A., and Danilel, J.J. (2011). Predicting Seabed Mud Content across the Australian Margin II: Performance of Machine Learning Methods and Their Combination with Ordinary Kriging and Inverse Distance Squared, Geoscience Australia, Record 2011/07, 69pp.

Greg Ridgeway with contributions from others (2015). gbm: Generalized Boosted Regression Models. R package version 2.1.1. <https://CRAN.R-project.org/package=gbm>

Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. Computers & Geosciences, 30: 683-691.

## Examples

```
library(spm)

data(sponge)
data(sponge.grid)
longlat <- sponge[, 1:2]

set.seed(1234)

gbmkrigeidwpred1 <- gbmkrigeidwpred(longlat = longlat, trainx = sponge[, -3],
  predx = sponge.grid, trainy = sponge[, 3], longlatpredx = sponge.grid[, c(1:2)],
  family = "poisson", interaction.depth = 3, transformation = "none", formula = res1 ~ 1,
  vgm.args = "Sph", nmaxkrige = 12, idp = 2, nmaxidw = 12, hybrid.parameter = 3,
  n.cores = 8)

names(gbmkrigeidwpred1)

range(gbmkrigeidwpred1$predictions)
```

---

glmcv

*Cross validation, n-fold and leave-one-out for generalised linear models ('glm')*

---

## Description

This function is a cross validation function for 'glm' method in 'stats' package.

## Usage

```
glmcv(
  formula = NULL,
  trainxy,
  y,
  family = "gaussian",
  validation = "CV",
  cv.fold = 10,
```



```

    predacc = "VEcv",
    ...
  )

```

### Arguments

formula	a formula defining the response variable and predictive variables.
trainxy	a dataframe contains predictive variables and the response variable of point samples. The location information, longitude (long), latitude (lat), need to be included in the 'trainx' for spatial predictive modeling.
y	a vector of the response variable in the formula, that is, the left part of the formula.
family	a description of the error distribution and link function to be used in the model. See '?glm' for details.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'glm'.

### Value

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

### Note

This function is largely based on 'rfcv' in 'randomForest' and 'glm' in 'stats'.

### Author(s)

Jin Li

### References

A. Liaw and M. Wiener (2002). Classification and Regression by randomForest. R News 2(3), 18-22.

### Examples

```

library(spm)

data(petrel)
gravel <- petrel[, c(1, 2, 6:9, 5)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
set.seed(1234)
glmcv1 <- glmcv(formula = model, gravel, log(gravel[, 7] +1), validation = "CV",

```

```

predacc = "ALL")
glmcv1 # Since the default 'family' is used, it is actually a 'lm' model.

data(sponge)
model <- sponge ~ easting + I(easting^2)
set.seed(1234)
glmcv1 <- glmcv(formula = model, sponge, sponge[, 3], family = poisson,
validation = "CV", predacc = "ALL")
glmcv1

# For glm
model <- gravel / 100 ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
glmcv1 <- glmcv(formula = model, gravel, gravel[, 7] / 100, family =
binomial(link=logit), validation = "CV", predacc = "VEcv")
VEcv [i] <- glmcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLM", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

```

---

glmidwcv

*Cross validation, n-fold and leave-one-out for the hybrid method of generalised linear models ('glm') and inverse distance weighted ('IDW') ('glmldw')*

---

## Description

This function is a cross validation function for the hybrid method of 'glm' and 'idw' using 'gstat' (glmldw) (see reference #1), where the data splitting is based on a stratified random sampling method (see the 'datasplit' function for details).

## Usage

```

glmldwcv(
  formula = NULL,
  longlat,
  trainxy,
  y,
  family = "gaussian",
  idp = 2,
  nmaxidw = 12,
  validation = "CV",
  cv.fold = 10,

```

```

    predacc = "VEcv",
    ...
)

```

### Arguments

formula	a formula defining the response variable and predictive variables for 'glm'.
longlat	a dataframe contains longitude and latitude of point samples.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples.
y	a vector of the response variable in the formula, that is, the left part of the formula.
family	a description of the error distribution and link function to be used in the model. See '?glm' for details.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'glm' and 'gstat'.

### Value

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only.

### Note

This function is largely based on 'rfcv' in 'randomForest', 'idwcv' in 'spm' and 'glm' in 'stats'.

### Author(s)

Jin Li

### References

- Li, J., Alvarez, B., Siwabessy, J., Tran, M., Huang, Z., Przeslawski, R., Radke, L., Howard, F. and Nichol, S. (2017). "Application of random forest, generalised linear model and their hybrid methods with geostatistical techniques to count data: Predicting sponge species richness." *Environmental Modelling & Software* 97: 112-129.
- A. Liaw and M. Wiener (2002). Classification and Regression by randomForest. *R News* 2(3), 18-22.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

**Examples**

```

library(spm)

data(petrel)
gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)

set.seed(1234)
glmidwcv1 <- glmidwcv(formula = model, longlat = longlat, trainxy = gravel,
y = y, idp = 2, nmaxidw = 12, validation = "CV", predacc = "ALL")
glmidwcv1 # Since the default 'family' is used, actually a 'lm' model is used.

data(spongelonglat)
longlat <- spongelonglat[, 7:8]
model <- sponge ~ long + I(long^2)
y = spongelonglat[, 1]
set.seed(1234)
glmidwcv1 <- glmidwcv(formula = model, longlat = longlat, trainxy = spongelonglat,
y = y, family = poisson, idp = 2, nmaxidw = 12, validation = "CV",
predacc = "ALL")
glmidwcv1

# glmidw for count data
data(spongelonglat)
longlat <- spongelonglat[, 7:8]
model <- sponge ~ . # use all predictive variables in the dataset
y = spongelonglat[, 1]
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  glmidwcv1 <- glmidwcv(formula = model, longlat = longlat, trainxy = spongelonglat,
y = y, family = poisson, idp = 2, nmaxidw = 12, validation = "CV",
predacc = "VEcv")
  VEcv [i] <- glmidwcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLM", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

# glmidw for percentage data
library(MASS)
longlat <- petrel[, c(1, 2)]
model <- gravel / 100 ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  glmidwcv1 <- glmcv(formula = model, longlat = longlat, trainxy = gravel,

```

```

y = gravel[, 7] / 100, family = binomial(link=logit), idp = 2, nmaxidw = 12,
validation = "CV", predacc = "VEcv")
VEcv [i] <- glmidwcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLM", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

```

---

glmidwpred	<i>Generate spatial predictions using the hybrid method of generalised linear models ('glm') and inverse distance weighted ('IDW') ('glmidw')</i>
------------	---

---

### Description

This function is for generating spatial predictions using the hybrid method of 'glm' and 'idw' ('glmidw') (see reference #1).

### Usage

```

glmidwpred(
  formula = NULL,
  longlat,
  trainxy,
  y,
  longlatpredx,
  predx,
  family = "gaussian",
  idp = 2,
  nmaxidw = 12,
  ...
)

```

### Arguments

formula	a formula defining the response variable and predictive variables for 'glm'.
longlat	a dataframe contains longitude and latitude of point samples. The location information must be named as 'long' and 'lat'.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be named as 'long' and 'lat'.
y	a vector of the response variable in the formula, that is, the left part of the formula.
longlatpredx	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted.

predx	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
family	a description of the error distribution and link function to be used in the model. See '?glm' for details.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
...	other arguments passed on to 'glm'.

**Value**

A dataframe of longitude, latitude, and predictions.

**Author(s)**

Jin Li

**References**

Li, J., Alvarez, B., Siwabessy, J., Tran, M., Huang, Z., Przeslawski, R., Radke, L., Howard, F. and Nichol, S. (2017). "Application of random forest, generalised linear model and their hybrid methods with geostatistical techniques to count data: Predicting sponge species richness." *Environmental Modelling & Software* 97: 112-129.

Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

**Examples**

```
library(spm)
data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)

glmidwpred1 <- glmidwpred(formula = model, longlat = longlat, trainxy = gravel,
y = y, longlatpredx = petrel.grid[, c(1:2)], predx = petrel.grid, idp = 2,
nmaxidw = 12)
# Since the default 'family' is used, actually a 'lm' model is used.

names(glmidwpred1)

# Back transform 'glmidwpred1$predictions' to generate the final predictions
glmidwpred1$predictions.bt <- exp(glmidwpred1$predictions) - 1
range(glmidwpred1$predictions.bt)
```

---

glmkrigecv	<i>Cross validation, n-fold and leave-one-out for the hybrid method of generalised linear models ('glm') and 'krige' ('glmkrige')</i>
------------	---

---

## Description

This function is a cross validation function for the hybrid method of 'glm' and 'krige' (glmkrige), where 'krige' methods include ordinary kriging ('OK'), simple kriging ('SK'), block 'OK' ('BOK') and block 'SK' ('BSK') (see reference #1 for further info).

## Usage

```
glmkrigecv(
  formula.glm = NULL,
  longlat,
  trainxy,
  y,
  family = "gaussian",
  transformation = "none",
  delta = 1,
  formula.krige = res1 ~ 1,
  vgm.args = c("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  validation = "CV",
  cv.fold = 10,
  predacc = "VEcv",
  ...
)
```

## Arguments

formula.glm	a formula defining the response variable and predictive variables for 'glm'.
longlat	a dataframe contains longitude and latitude of point samples.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples.
y	a vector of the response variable in the formula, that is, the left part of the formula.
family	a description of the error distribution and link function to be used in the model. See '?glm' for details.

transformation	transform the residuals of 'glm' to normalise the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
delta	numeric; to avoid log(0) in the log transformation. The default is 1.
formula.krige	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.
vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'glm' and 'krige'.

**Value**

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

**Note**

This function is largely based on 'rfcv' in 'randomForest', 'krigecv' in 'spm2' and 'glm' in 'stats'.

**Author(s)**

Jin Li

**References**

Li, J., Alvarez, B., Siwabessy, J., Tran, M., Huang, Z., Przeslawski, R., Radke, L., Howard, F. and Nichol, S. (2017). "Application of random forest, generalised linear model and their hybrid methods with geostatistical techniques to count data: Predicting sponge species richness." *Environmental Modelling & Software* 97: 112-129.

Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.



**Examples**

```

library(spm)

data(petrel)

gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)
set.seed(1234)
glmkrigecv1 <- glmkrigecv(formula.glm = model, longlat = longlat, trainxy = gravel,
y = y, transformation = "none", formula.krige = res1 ~ 1, vgm.args = "Sph",
nmaxkrige = 12, validation = "CV", predacc = "ALL")
glmkrigecv1 # Since the default 'family' is used, actually a 'lm' model is used.

data(spongelonglat)
longlat <- spongelonglat[, 7:8]
model <- sponge ~ long + I(long^2)
y = spongelonglat[, 1]
set.seed(1234)
glmkrigecv1 <- glmkrigecv(formula.glm = model, longlat = longlat, trainxy =
spongelonglat, y = y, family = poisson, transformation = "arcsine",
formula.krige = res1 ~ 1, vgm.args = ("Sph"), nmaxkrige = 12,
validation = "CV", predacc = "ALL")
glmkrigecv1

# glmok for count data
data(spongelonglat)
longlat <- spongelonglat[, 7:8]
model <- sponge ~ . # use all predictive variables in the dataset
y = spongelonglat[, 1]
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  glmkrigecv1 <- glmkrigecv(formula.glm = model, longlat = longlat, trainxy = spongelonglat,
y = y, family = poisson, formula.krige = res1 ~ 1, vgm.args = ("Sph"), nmaxkrige = 12,
validation = "CV", predacc = "VEcv")
  VEcv [i] <- glmkrigecv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLM", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

# glmok for percentage data
library(MASS)
longlat <- petrel[, c(1, 2)]
model <- gravel / 100 ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL

```

```

for (i in 1:n) {
  glmkrige1 <- glmkrigeidwcv(formula.glm = model, longlat = longlat, trainxy = gravel,
  y = gravel[, 7] / 100, family = binomial(link=logit), formula.krige = res1 ~ 1,
  vgm.args = ("Sph"), nmaxkrige = 12, validation = "CV", predacc = "VEcv")
  VEcv [i] <- glmkrige1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLM", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

```

---

glmkrigeidwcv

*Cross validation, n-fold and leave-one-out for the hybrid methods of generalised linear models ('glm'), 'kriging' and inverse distance weighted ('IDW').*

---

## Description

This function is a cross validation function for 38 hybrid methods of 'glm', 'kriging' and 'IDW', including the average of 'glmkrige' and 'glmkrigeidw' ('glmkrigeidw') and the average of 'glm', 'glmkrige' and 'glmkrigeidw' ('glmkrigeidw'), where 'kriging' methods include ordinary kriging ('OK'), simple kriging ('SK'), block 'OK' ('BOK') and block 'SK' ('BSK') and 'IDW' also covers 'NN' and 'KNN' (for details, see reference #1). This function can also be used for 38 hybrid methods of 'lm', 'kriging' and 'IDW'.

## Usage

```

glmkrigeidwcv(
  formula.glm = NULL,
  longlat,
  trainxy,
  y,
  family = "gaussian",
  transformation = "none",
  delta = 1,
  formula.krige = res1 ~ 1,
  vgm.args = c("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  idp = 2,
  nmaxidw = 12,
  hybrid.parameter = 2,
  lambda = 1,
  validation = "CV",

```

```

    cv.fold = 10,
    predacc = "VEcv",
    ...
)

```

## Arguments

<code>formula.glm</code>	a formula defining the response variable and predictive variables for 'glm'.
<code>longlat</code>	a dataframe contains longitude and latitude of point samples.
<code>trainxy</code>	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples.
<code>y</code>	a vector of the response variable in the formula, that is, the left part of the formula.
<code>family</code>	a description of the error distribution and link function to be used in the model. See '?glm' for details.
<code>transformation</code>	transform the residuals of 'glm' to normalise the data for 'krige'; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
<code>delta</code>	numeric; to avoid log(0) in the log transformation. The default is 1.
<code>formula.krige</code>	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.
<code>vgm.args</code>	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
<code>anis</code>	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
<code>alpha</code>	direction in plane (x,y). see variogram in 'gstat' for details.
<code>block</code>	block size. see 'krige' in 'gstat' for details.
<code>beta</code>	for simple kriging. see 'krige' in 'gstat' for details.
<code>nmaxkrige</code>	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
<code>idp</code>	a numeric number specifying the inverse distance weighting power.
<code>nmaxidw</code>	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
<code>hybrid.parameter</code>	the default is 2 that is for 'glmkrigeidw'; for 'glmglmkrigeidw', it needs to be 3.
<code>lambda</code> ,	ranging from 0 to 2; the default is 1 for 'glmkrigeidw' and 'glmglmkrigeidw'; and if it is < 1, more weight is placed on 'krige', otherwise more weight is placed on 'idw'; and if it is 0, 'idw' is not considered and the resultant methods is 'glmkrige' when the default 'hybrid.parameter' is used; and if it is 2, then the resultant method is 'glmidw' when the default 'hybrid.parameter' is used.

validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'glm', 'krige' and 'gstat'.

**Value**

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

**Note**

This function is largely based on 'rfcv' in 'randomForest', 'krige' in 'spmg' and 'glm' in 'stats'.

**Author(s)**

Jin Li

**References**

Li, J., Alvarez, B., Siwabessy, J., Tran, M., Huang, Z., Przeslawski, R., Radke, L., Howard, F. and Nichol, S. (2017). "Application of random forest, generalised linear model and their hybrid methods with geostatistical techniques to count data: Predicting sponge species richness." *Environmental Modelling & Software* 97: 112-129.

A. Liaw and M. Wiener (2002). Classification and Regression by randomForest. *R News* 2(3), 18-22.

Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

**Examples**

```
library(spm)
# glmokglidw
data(petrel)
gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)
set.seed(1234)
glmkrigeidwcv1 <- glmkrigeidwcv(formula.glm = model, longlat = longlat,
trainxy = gravel, y = y, transformation = "none", formula.krige = res1 ~ 1,
vgm.args = "Sph", nmaxkrige = 12, idp = 2, nmaxidw = 12, validation = "CV",
predacc = "ALL")
glmkrigeidwcv1 # Since the default 'family' is used, actually a 'lm' model is used.

# glmokglmidw
data(spongelonglat)
longlat <- spongelonglat[, 7:8]
```

```

model <- sponge ~ long + I(long^2)
y = spongelonglat[, 1]
set.seed(1234)
glmkrigeidwcv1 <- glmkrigeidwcv(formula.glm = model, longlat = longlat,
trainxy = spongelonglat, y = y, family = poisson, transformation = "arcsine",
formula.krige = res1 ~ 1, vgm.args = ("Sph"), nmaxkrige = 12, idp = 2,
nmaxidw = 12, validation = "CV", predacc = "ALL")
glmkrigeidwcv1

# glmokglmidw
data(spongelonglat)
longlat <- spongelonglat[, 7:8]
model <- sponge ~ long + I(long^2)
y = spongelonglat[, 1]
set.seed(1234)
glmokglmidwcv1 <- glmkrigeidwcv(formula.glm = model, longlat = longlat,
trainxy = spongelonglat, y = y, family = poisson, transformation = "arcsine",
formula.krige = res1 ~ 1, vgm.args = ("Sph"), nmaxkrige = 12, idp = 2,
nmaxidw = 12, hybrid.parameter = 3, validation = "CV", predacc = "ALL")
glmokglmidwcv1

# glmokglidw for count data
data(spongelonglat)
longlat <- spongelonglat[, 7:8]
model <- sponge ~ . # use all predictive variables in the dataset
y = spongelonglat[, 1]
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  glmkrigeidwcv1 <- glmkrigeidwcv(formula.glm = model, longlat = longlat,
trainxy = spongelonglat, y = y, family = poisson, formula.krige = res1 ~ 1,
vgm.args = ("Sph"), nmaxkrige = 12, idp = 2, nmaxidw = 12, validation = "CV",
predacc = "VEcv")
  VEcv [i] <- glmkrigeidwcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLM", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

# glmokglmidw for percentage data
library(MASS)
longlat <- petrel[, c(1, 2)]
model <- gravel / 100 ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  glmkrigeidwcv1 <- glmkrigeidwcv(formula.glm = model, longlat = longlat,
trainxy = gravel, y = gravel[, 7] / 100, family = binomial(link=logit),
formula.krige = res1 ~ 1, vgm.args = ("Sph"), nmaxkrige = 12, idp = 2,
nmaxidw = 12, validation = "CV", predacc = "VEcv")
  VEcv [i] <- glmkrigeidwcv1
}

```

```

}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLM", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

```

---

glmkrigeidwpred	<i>Generate spatial predictions using the hybrid methods of generalised linear models ('glm'), 'kriging' and inverse distance weighted ('IDW').</i>
-----------------	---

---

### Description

This function is for generating spatial predictions using the hybrid methods of 'glm', 'kriging' and 'IDW', including all methods implemented in 'glmkrigeidwcv'.

### Usage

```

glmkrigeidwpred(
  formula.glm = NULL,
  longlat,
  trainxy,
  predx,
  y,
  longlatpredx,
  family = "gaussian",
  transformation = "none",
  delta = 1,
  formula.krige = res1 ~ 1,
  vgm.args = c("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  idp = 2,
  nmaxidw = 12,
  hybrid.parameter = 2,
  lambda = 1,
  ...
)

```

### Arguments

formula.glm	a formula defining the response variable and predictive variables for 'glm'.
longlat	a dataframe contains longitude and latitude of point samples.

trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples.
predx	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
y	a vector of the response variable in the formula, that is, the left part of the formula.
longlatpredx	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted.
family	a description of the error distribution and link function to be used in the model. See '?glm' for details.
transformation	transform the residuals of 'glm' to normalise the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
delta	numeric; to avoid log(0) in the log transformation. The default is 1.
formula.krige	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.
vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
hybrid.parameter	the default is 2 that is for 'glmkrigeidw'; for 'glmglmkrigeidw', it needs to be 3.
lambda,	ranging from 0 to 2; the default is 1 for 'glmkrigeidw' and 'glmglmkrigeidw'; and if it is < 1, more weight is placed on 'krige', otherwise more weight is placed on 'idw'; and if it is 0, 'idw' is not considered and the resultant methods is 'glmkrige' when the default 'hybrid.parameter' is used; and if it is 2, then the resultant method is 'glmidw' when the default 'hybrid.parameter' is used.
...	other arguments passed on to 'glm', 'krige' and 'gstat'.

**Value**

A dataframe of longitude, latitude, and predictions.

**Author(s)**

Jin Li

**References**

Li, J., Alvarez, B., Siwabessy, J., Tran, M., Huang, Z., Przeslawski, R., Radke, L., Howard, F. and Nichol, S. (2017). "Application of random forest, generalised linear model and their hybrid methods with geostatistical techniques to count data: Predicting sponge species richness." *Environmental Modelling & Software* 97: 112-129.

Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

**Examples**

```
library(spm)

data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)

glmkrigeidwpred1 <- glmkrigeidwpred(formula.glm = model, longlat = longlat, trainxy = gravel,
  predx = petrel.grid, y = y, longlatpredx = petrel.grid[, c(1:2)],
  formula.krige = res1 ~ 1, vgm.args = "Sph", nmaxkrige = 12, idp = 2, nmaxidw = 12)
# Since the default 'family' is used, actually a 'lm' model is used.

names(glmkrigeidwpred1)

# Back transform 'glmkrigeidwpred$predictions' to generate the final predictions
glmkrigeidw.predictions <- exp(glmkrigeidwpred1$predictions) - 1
range(glmkrigeidw.predictions)
```

---

 glmkrigepred

*Generate spatial predictions using the hybrid method of generalised linear models ('glm') and 'krige'*

---

**Description**

This function is for generating spatial predictions using the hybrid method of 'glm' and 'krige', including all methods implemented in 'glmkrigecv'. (see reference #1 for further info).



**Usage**

```

glmkrigepred(
  formula.glm = NULL,
  longlat,
  trainxy,
  predx,
  y,
  longlatpredx,
  family = "gaussian",
  transformation = "none",
  delta = 1,
  formula.krige = res1 ~ 1,
  vgm.args = c("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  ...
)

```

**Arguments**

<code>formula.glm</code>	a formula defining the response variable and predictive variables for 'glm'.
<code>longlat</code>	a dataframe contains longitude and latitude of point samples.
<code>trainxy</code>	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples.
<code>predx</code>	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
<code>y</code>	a vector of the response variable in the formula, that is, the left part of the formula.
<code>longlatpredx</code>	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted.
<code>family</code>	a description of the error distribution and link function to be used in the model. See '?glm' for details.
<code>transformation</code>	transform the residuals of 'glm' to normalise the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
<code>delta</code>	numeric; to avoid log(0) in the log transformation. The default is 1.
<code>formula.krige</code>	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.
<code>vgm.args</code>	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
<code>anis</code>	anisotropy parameters: see notes 'vgm' in 'gstat' for details.

alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
...	other arguments passed on to 'glm' and 'krige'.

**Value**

A dataframe of longitude, latitude, and predictions.

**Author(s)**

Jin Li

**References**

Li, J., Alvarez, B., Siwabessy, J., Tran, M., Huang, Z., Przeslawski, R., Radke, L., Howard, F. and Nichol, S. (2017). "Application of random forest, generalised linear model and their hybrid methods with geostatistical techniques to count data: Predicting sponge species richness." *Environmental Modelling & Software* 97: 112-129.

Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

**Examples**

```
library(spm)

data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)

glmkrigepred1 <- glmkrigepred(formula.glm = model, longlat = longlat, trainxy = gravel,
  predx = petrel.grid, y = y, longlatpredx = petrel.grid[, c(1:2)],
  transformation = "none", formula.krige = res1 ~ 1, vgm.args = "Sph", nmaxkrige = 12)
# Since the default 'family' is used, actually a 'lm' model is used.

names(glmkrigepred1)

# Back transform 'glmkrigepred$predictions' to generate the final predictions
glmkrige.predictions <- exp(glmkrigepred1$predictions) - 1
range(glmkrige.predictions)
```

---

glmnetcv	<i>Cross validation, n-fold and leave-one-out, for 'glmnet' in 'glmnet' package</i>
----------	---

---

## Description

This function is a cross validation function for 'glmnet' method in 'glmnet' package.

## Usage

```
glmnetcv(
  trainx,
  y,
  family = "gaussian",
  alpha = 0.5,
  relax = FALSE,
  type.measure = "mse",
  validation = "CV",
  cv.fold = 10,
  predacc = "VEcv",
  ...
)
```

## Arguments

trainx	a matrix contains predictive variables of point samples. The location information, longitude (long), latitude (lat), need to be included in the 'trainx' for spatial predictive modelling.
y	a vector of the response variable in the formula, that is, the left part of the formula.
family	a description of the error distribution and link function to be used in the model. See '?glmnet' for details.
alpha,	an elasticnet mixing parameter, with $0 \leq \alpha \leq 1$ . See '?glmnet' for details.
relax,	if TRUE then for each active set in the path of solutions, the model is refit without any regularization. See '?glmnet' for more information.
type.measure,	loss to use for cross-validation. The default is type.measure="mse". See '?cv.glmnet' for more information.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'fields'.

**Value**

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

**Note**

This function is largely based on 'glmcv' in this 'spm2' package.

**Author(s)**

Jin Li

**References**

A. Liaw and M. Wiener (2002). Classification and Regression by randomForest. R News 2(3), 18-22.

**Examples**

```
library(spm)

data(petrel)
x <- as.matrix(petrel[, c(1, 2, 6:9)])
y <- log(petrel[, 5] + 1)
set.seed(1234)
glmnetcv1 <- glmnetcv(x, y, validation = "CV", predacc = "ALL")
glmnetcv1

data(sponge)
x <- as.matrix(cbind(sponge$easting, sponge$easting^2))
set.seed(1234)
glmnetcv1 <- glmnetcv(x, sponge[, 3], family = poisson, validation = "CV",
predacc = "ALL")
glmnetcv1

# For glmnet with gaussian
x <- as.matrix(petrel[, c(1, 2, 6:9)])
y <- log(petrel[, 5] + 1)
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  glmnetcv1 <- glmnetcv(x, y, validation = "CV", predacc = "VEcv")
  VEcv [i] <- glmnetcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for glmnet", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

# For glmnet with binomial
x <- as.matrix(cbind(petrel[, c(2, 6)], petrel$long^3, petrel$lat^2, petrel$lat^3))
```

```

set.seed(1234)
n <- 20 # number of iterations,60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  glmnetcv1 <- glmnetcv(x, petrel[, 5] / 100, family = binomial(link=logit),
  validation = "CV", predacc = "VEcv")
  VEcv [i] <- glmnetcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for glmnet", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

```

---

glmpred

*Generate spatial predictions using generalised linear models ('glm')*


---

### Description

This function is for generating spatial predictions using 'glm' method in 'stats' package.

### Usage

```
glmpred(formula = NULL, trainxy, longlatpredx, predx, family = "gaussian", ...)
```

### Arguments

formula	a formula defining the response variable and predictive variables.
trainxy	a dataframe contains predictive variables and the response variable of point samples. The location information, longitude (long), latitude (lat), need to be included in the 'trainx' for spatial predictive modeling, need to be named as 'long' and 'lat'.
longlatpredx	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted.
predx	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
family	a description of the error distribution and link function to be used in the model. See '?glm' for details.
...	other arguments passed on to 'glm'.

### Value

A dataframe of longitude, latitude and predictions.

### Author(s)

Jin Li

**Examples**

```

library(spm)
data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)

glm1 <- glm(formula = model, trainxy = gravel,
            longlatpredx = petrel.grid[, c(1:2)], predx = petrel.grid)

names(glm1)

# Back transform 'glm1$pred.glm1' to generate the final predictions
glm.predictions <- exp(glm1$pred.glm1) - 1
range(glm.predictions)

```

---

glscv	<i>Cross validation, n-fold and leave-one-out for generalized least squares ('gls')</i>
-------	---

---

**Description**

This function is a cross validation function for 'gls' method in 'nlme' package.

**Usage**

```

glscv(
  model = var1 ~ 1,
  trainxy,
  y,
  corr.args = NULL,
  weights = NULL,
  validation = "CV",
  cv.fold = 10,
  predacc = "VEcv",
  ...
)

```

**Arguments**

model	a formula defining the response variable and predictive variables.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be names as 'long' and 'lat'.

<code>y</code>	a vector of the response variable in the formula, that is, the left part of the formula.
<code>corr.args</code>	arguments for 'correlation' in 'gls'. See '?corClasses' in 'nlme' for details. By default, "NULL" is used. When "NULL" is used, then 'gls' is actually performing 'lm'.
<code>weights</code>	describing the within-group heteroscedasticity structure. Defaults to "NULL", corresponding to homoscedastic errors. See '?gls' in 'nlme' for details.
<code>validation</code>	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
<code>cv.fold</code>	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
<code>predacc</code>	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
<code>...</code>	other arguments passed on to 'gls'.

**Value**

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

**Note**

This function is largely based on rfcv in 'randomForest' and 'gls' in 'library(nlme)'.

**Author(s)**

Jin Li

**References**

Pinheiro, J. C. and D. M. Bates (2000). Mixed-Effects Models in S and S-PLUS. New York, Springer.

**Examples**

```
library(spm)
library(nlme)

data(petrel)
gravel <- petrel[, c(1, 2, 6:9, 5)]
range1 <- 0.8
nugget1 <- 0.5

model <- log(gravel + 1) ~ long + lat + bathy + dist + I(long^2) + I(lat^2) +
I(lat^3) + I(bathy^2) + I(bathy^3) + I(dist^2) + I(dist^3) + I(relief^2) + I(relief^3)

glscv1 <- glscv(model = model, gravel, log(gravel[, 7] + 1), validation = "CV",
  corr.args = corSpher(c(range1, nugget1), form = ~ long + lat, nugget = T),
  predacc = "ALL")
```

```

gls cv1

#For gls
set.seed(1234)
n <- 20 # number of iterations,60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
gls cv1 <- gls cv(model = model, gravel, log(gravel[, 7] +1), validation = "CV",
  corr.args = corSpher(c(range1, nugget1), form = ~ long + lat,
  nugget = T), predacc = "VEcv")
VEcv [i] <- gls cv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLS", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

# For lm, that is, gls with 'correlation = NULL'
n <- 20 # number of iterations,60 to 100 is recommended.
VEcv <- NULL
set.seed(1234)
for (i in 1:n) {
gls cv1 <- gls cv(model = model, gravel, log(gravel[, 7] +1),
  validation = "CV", predacc = "VEcv")
VEcv [i] <- gls cv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLS", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

```

---

glsidwcv

*Cross validation, n-fold and leave-one-out for the hybrid method of generalized least squares ('gls') and inverse distance weighted ('idw') (glsidw)*

---

## Description

This function is a cross validation function for the hybrid method of 'gls' and 'idw', where the data splitting is based on a stratified random sampling method (see the 'datasplit' function for details)

## Usage

```

glsidwcv(
  model = var1 ~ 1,
  longlat,
  trainxy,
  y,
  corr.args = NULL,

```



```

weights = NULL,
idp = 2,
nmaxidw = 12,
validation = "CV",
cv.fold = 10,
predacc = "VEcv",
...
)

```

### Arguments

model	a formula defining the response variable and predictive variables.
longlat	a dataframe contains longitude and latitude of point samples.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be names as 'long' and 'lat'.
y	a vector of the response variable in the formula, that is, the left part of the formula.
corr.args	arguments for 'correlation' in 'gls'. See '?corClasses' in 'nlme' for details. By default, "NULL" is used. When "NULL" is used, then 'gls' is actually performing 'lm'.
weights	describing the within-group heteroscedasticity structure. Defaults to "NULL", corresponding to homoscedastic errors. See '?gls' in 'nlme' for details.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'gls' and 'gstat'.

### Value

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only.

### Note

This function is largely based on rfcv in 'randomForest' and 'gls' in 'library(nlme)'.

### Author(s)

Jin Li

## References

- Pinheiro, J. C. and D. M. Bates (2000). *Mixed-Effects Models in S and S-PLUS*. New York, Springer.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

## Examples

```
library(spm)
library(nlme)

data(petrel)
gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
range1 <- 0.8
nugget1 <- 0.5
model <- log(gravel + 1) ~ long + lat + bathy + dist + I(long^2) + I(lat^2) +
I(lat^3) + I(bathy^2) + I(bathy^3) + I(dist^2) + I(dist^3) + I(relief^2) + I(relief^3)

glsidwcv1 <- glsidwcv(model = model, longlat = longlat, trainxy = gravel,
y = log(gravel[, 7] + 1), idp = 2, nmaxidw = 12, validation = "CV",
corr.args = corSpher(c(range1, nugget1), form = ~ lat + long, nugget = T),
predacc = "ALL")
glsidwcv1

# For glsidw
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
glsidwcv1 <- glsidwcv(model = model, longlat = longlat, trainxy = gravel,
y = log(gravel[, 7] + 1), idp = 2, nmaxidw = 12, validation = "CV",
corr.args = corSpher(c(range1, nugget1), form = ~ lat + long, nugget = T),
predacc = "VEcv")
VEcv [i] <- glsidwcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLSIDW", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)
```

**Description**

This function is for generating spatial predictions using the hybrid method of 'gls' and 'idw' ('glsidw') (see reference #1).

**Usage**

```
glsidwpred(
  model = var1 ~ 1,
  longlat,
  trainxy,
  y,
  longlatpredx,
  predx,
  corr.args = NULL,
  weights = NULL,
  idp = 2,
  nmaxidw = 12,
  ...
)
```

**Arguments**

model	a formula defining the response variable and predictive variables.
longlat	a dataframe contains longitude and latitude of point samples.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be names as 'long' and 'lat'.
y	a vector of the response variable in the formula, that is, the left part of the formula.
longlatpredx	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted. The location information must be named as 'long' and 'lat'.
predx	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
corr.args	arguments for 'correlation' in 'gls'. See '?corClasses' in 'nlme' for details. By default, "NULL" is used. When "NULL" is used, then 'gls' is actually performing 'lm'.
weights	describing the within-group heteroscedasticity structure. Defaults to "NULL", corresponding to homoscedastic errors. See '?gls' in 'nlme' for details.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
...	other arguments passed on to 'gls' and 'gstat'.

**Value**

A dataframe of longitude, latitude, and predictions.

**Author(s)**

Jin Li

**References**

- Pinheiro, J. C. and D. M. Bates (2000). *Mixed-Effects Models in S and S-PLUS*. New York, Springer.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

**Examples**

```
library(spm)
library(nlme)

data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
range1 <- 0.8
nugget1 <- 0.5
model <- log(gravel + 1) ~ long + lat + bathy + dist + I(long^2) + I(lat^2) +
I(lat^3) + I(bathy^2) + I(bathy^3) + I(dist^2) + I(dist^3) + I(relief^2) + I(relief^3)
y <- log(gravel[, 7] + 1)

glsidwpred1 <- glsidwpred(model = model, longlat = longlat, trainxy = gravel,
y = y, longlatpredx = petrel.grid[, c(1:2)], predx = petrel.grid,
idp = 2, nmaxidw = 12, corr.args = corSpher(c(range1, nugget1),
form = ~ lat + long, nugget = T))

names(glsidwpred1)

# Back transform 'glsidwpred1$predictions' to generate the final predictions
glsidw.predictions <- exp(glsidwpred1$predictions) - 1
range(glsidw.predictions)
```

**Description**

This function is a cross validation function for the hybrid method of 'gls' and 'krige' ('glskrige'), where the data splitting is based on a stratified random sampling method (see the 'datasplit' function for details)

**Usage**

```
glskrige cv(
  model = var1 ~ 1,
  longlat,
  trainxy,
  y,
  corr.args = NULL,
  weights = NULL,
  transformation = "none",
  delta = 1,
  formula.krige = res1 ~ 1,
  vgm.args = c("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  validation = "CV",
  cv.fold = 10,
  predacc = "VEcv",
  ...
)
```

**Arguments**

model	a formula defining the response variable and predictive variables.
longlat	a dataframe contains longitude and latitude of point samples.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be names as 'long' and 'lat'.
y	a vector of the response variable in the formula, that is, the left part of the formula.
corr.args	arguments for 'correlation' in 'gls'. See '?corClasses' in 'nlme' for details. By default, "NULL" is used. When "NULL" is used, then 'gls' is actually performing 'lm'.
weights	describing the within-group heteroscedasticity structure. Defaults to "NULL", corresponding to homoscedastic errors. See '?gls' in 'nlme' for details.
transformation	transform the residuals of 'gls' to normalize the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.

<code>delta</code>	numeric; to avoid $\log(0)$ in the log transformation. The default is 1.
<code>formula.krige</code>	formula defining the response vector and (possible) regressor. an object (i.e., <code>'variogram.formula'</code> ) for <code>'variogram'</code> or a formula for <code>'krige'</code> . see <code>'variogram'</code> and <code>'krige'</code> in <code>'gstat'</code> for details.
<code>vgm.args</code>	arguments for <code>'vgm'</code> , e.g. variogram model of response variable and anisotropy parameters. see <code>'vgm'</code> in <code>'gstat'</code> for details. By default, "Sph" is used.
<code>anis</code>	anisotropy parameters: see notes <code>'vgm'</code> in <code>'gstat'</code> for details.
<code>alpha</code>	direction in plane (x,y). see <code>variogram</code> in <code>'gstat'</code> for details.
<code>block</code>	block size. see <code>'krige'</code> in <code>'gstat'</code> for details.
<code>beta</code>	for simple kriging. see <code>'krige'</code> in <code>'gstat'</code> for details.
<code>nmaxkrige</code>	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
<code>validation</code>	validation methods, include <code>'LOO'</code> : leave-one-out, and <code>'CV'</code> : cross-validation.
<code>cv.fold</code>	integer; number of folds in the cross-validation. if $> 1$ , then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
<code>predacc</code>	can be either <code>"VEcv"</code> for <code>vecv</code> or <code>"ALL"</code> for all measures in function <code>pred.acc</code> .
<code>...</code>	other arguments passed on to <code>'gls'</code> and <code>'krige'</code> .

**Value**

A list with the following components: `me`, `rme`, `mae`, `rmae`, `mse`, `rmse`, `rmse`, `vecv` and `e1`; or `vecv` only.

**Note**

This function is largely based on `rfcv` in `'randomForest'`, `'krigecv'` in `'spm2'` and `'gls'` in `'library(nlme)'`.

**Author(s)**

Jin Li

**References**

- Pinheiro, J. C. and D. M. Bates (2000). *Mixed-Effects Models in S and S-PLUS*. New York, Springer.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the `gstat` package. *Computers & Geosciences*, 30: 683-691.

**Examples**

```
library(spm)
library(nlme)
```

```

data(petrel)
gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
range1 <- 0.8
nugget1 <- 0.5
model <- log(gravel + 1) ~ long + lat + bathy + dist + I(long^2) + I(lat^2) +
I(lat^3) + I(bathy^2) + I(bathy^3) + I(dist^2) + I(dist^3) + I(relief^2) + I(relief^3)

glskrigevc1 <- glskrigevcv(model = model, longlat = longlat, trainxy = gravel,
y = log(gravel[, 7] + 1), transformation = "none", formula.krige = res1 ~ 1,
vgm.args = "Sph", nmaxkrige = 12, validation = "CV",
corr.args = corSpher(c(range1, nugget1), form = ~ lat + long, nugget = T),
predacc = "ALL")
glskrigevc1

# For glskrige
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
glskrigevc1 <- glskrigevcv(model = model, longlat = longlat, trainxy = gravel,
y = log(gravel[, 7] + 1), transformation = "none", formula.krige = res1 ~ 1,
vgm.args = "Sph", nmaxok = 12, validation = "CV",
corr.args = corSpher(c(range1, nugget1), form = ~ lat + long, nugget = T),
predacc = "VEcv")
VEcv [i] <- glskrigevc1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLSOK", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

```

---

glskrigeidwcv

*Cross validation, n-fold and leave-one-out for the hybrid methods of generalised least squares ('gls'), 'kriging' and inverse distance weighted ('IDW')*

---

## Description

This function is a cross validation function for 38 hybrid methods of 'gls', 'kriging' and 'IDW', including the average of 'glskrige' and 'glsidw' ('glskrigeidw') and the average of 'gls', 'glskrige' and 'glsidw' ('glsidw'), where 'kriging' methods include ordinary kriging ('OK'), simple kriging ('SK'), block 'OK' ('BOK') and block 'SK' ('BSK') and 'IDW' also covers 'NN' and 'KNN'.. The data splitting is based on a stratified random sampling method (see the 'datasplit' function for details).

## Usage

```
glskrigeidwcv(
```

```

model = var1 ~ 1,
longlat,
trainxy,
y,
corr.args = NULL,
weights = NULL,
transformation = "none",
delta = 1,
formula.krige = res1 ~ 1,
vgm.args = c("Sph"),
anis = c(0, 1),
alpha = 0,
block = 0,
beta,
nmaxkrige = 12,
idp = 2,
nmaxidw = 12,
hybrid.parameter = 2,
lambda = 1,
validation = "CV",
cv.fold = 10,
predacc = "VEcv",
...
)

```

### Arguments

<code>model</code>	a formula defining the response variable and predictive variables.
<code>longlat</code>	a dataframe contains longitude and latitude of point samples.
<code>trainxy</code>	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be names as 'long' and 'lat'.
<code>y</code>	a vector of the response variable in the formula, that is, the left part of the formula.
<code>corr.args</code>	arguments for 'correlation' in 'gls'. See '?corClasses' in 'nlme' for details. By default, "NULL" is used. When "NULL" is used, then 'gls' is actually performing 'lm'.
<code>weights</code>	describing the within-group heteroscedasticity structure. Defaults to "NULL", corresponding to homoscedastic errors. See '?gls' in 'nlme' for details.
<code>transformation</code>	transform the residuals of 'gls' to normalise the data for 'krige'; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
<code>delta</code>	numeric; to avoid log(0) in the log transformation. The default is 1.
<code>formula.krige</code>	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.



vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
hybrid.parameter	the default is 2 that is for 'glskrigeidw'; for 'glskrigeidw', it needs to be 3.
lambda,	ranging from 0 to 2; the default is 1 for 'glskrigeidw' and 'glskrigeidw'; and if it is < 1, more weight is placed on 'krige', otherwise more weight is placed on 'idw'; and if it is 0, 'idw' is not considered and the resultant methods is 'glskrige' when the default 'hybrid.parameter' is used; and if it is 2, then the resultant method is 'idw' when the default 'hybrid.parameter' is used.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'gls', 'krige' and 'gstat'.

**Value**

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only.

**Note**

This function is largely based on rfcv in 'randomForest', 'krige' in 'spm2' and 'gls' in 'library(nlme)'.

**Author(s)**

Jin Li

**References**

- Pinheiro, J. C. and D. M. Bates (2000). Mixed-Effects Models in S and S-PLUS. New York, Springer.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. Computers & Geosciences, 30: 683-691.

**Examples**

```

library(spm)
library(nlme)

data(petrel)
gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
range1 <- 0.8
nugget1 <- 0.5
model <- log(gravel + 1) ~ long + lat + bathy + dist + I(long^2) + I(lat^2) +
I(lat^3) + I(bathy^2) + I(bathy^3) + I(dist^2) + I(dist^3) + I(relief^2) + I(relief^3)

glskrigeidwcv1 <- glskrigeidwcv(model = model, longlat = longlat, trainxy = gravel,
y = log(gravel[, 7] + 1), transformation = "none", formula.krige = res1 ~ 1,
vgm.args = "Sph", nmaxkrige = 12, idp = 2, nmaxidw = 12, validation = "CV",
corr.args = corSpher(c(range1, nugget1), form = ~ lat + long, nugget = T),
predacc = "ALL")
glskrigeidwcv1

# For glskrigeidwcv
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
glskrigeidwcv1 <- glskrigeidwcv(model = model, longlat = longlat, trainxy = gravel,
y = log(gravel[, 7] + 1), transformation = "none", formula.krige = res1 ~ 1,
vgm.args = "Sph", nmaxkrige = 12, idp = 2, nmaxidw = 12, validation = "CV",
corr.args = corSpher(c(range1, nugget1), form = ~ lat + long, nugget = T),
predacc = "VEcv")
VEcv [i] <- glskrigeidwcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLSOKGLSIDW", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

# For glsglskrigeidwcv
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
glskrigeidwcv1 <- glskrigeidwcv(model = model, longlat = longlat, trainxy = gravel,
y = log(gravel[, 7] + 1), transformation = "none", formula.krige = res1 ~ 1,
vgm.args = "Sph", nmaxkrige = 12, idp = 2, nmaxidw = 12, hybrid.parameter = 3,
validation = "CV", corr.args = corSpher(c(range1, nugget1), form = ~ lat + long,
nugget = T), predacc = "VEcv")
VEcv [i] <- glskrigeidwcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for GLSOKGLSIDW", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

```

---

glskrigewpred	<i>Generate spatial predictions using the hybrid methods of generalised least squares ('gls'), 'kriging' and inverse distance weighted ('IDW')</i>
---------------	--

---

### Description

This function is for generating spatial predictions using the hybrid methods of 'gls', 'kriging' and 'IDW', including all methods implemented in 'glskrigewcv'.

### Usage

```
glskrigewpred(
  model = var1 ~ 1,
  longlat,
  trainxy,
  predx,
  y,
  longlatpredx,
  corr.args = NULL,
  weights = NULL,
  transformation = "none",
  delta = 1,
  formula.krige = res1 ~ 1,
  vgm.args = c("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  idp = 2,
  nmaxidw = 12,
  hybrid.parameter = 2,
  lambda = 1,
  ...
)
```

### Arguments

model	a formula defining the response variable and predictive variables.
longlat	a dataframe contains longitude and latitude of point samples.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be names as 'long' and 'lat'.

predx	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
y	a vector of the response variable in the formula, that is, the left part of the formula.
longlatpredx	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted. The location information must be named as 'long' and 'lat'.
corr.args	arguments for 'correlation' in 'gls'. See '?corClasses' in 'nlme' for details. By default, "NULL" is used. When "NULL" is used, then 'gls' is actually performing 'lm'.
weights	describing the within-group heteroscedasticity structure. Defaults to "NULL", corresponding to homoscedastic errors. See '?gls' in 'nlme' for details.
transformation	transform the residuals of 'gls' to normalise the data for 'krige'; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
delta	numeric; to avoid log(0) in the log transformation. The default is 1.
formula.krige	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.
vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
hybrid.parameter	the default is 2 that is for 'glskrigeidw'; for 'glskrigeidw', it needs to be 3.
lambda,	ranging from 0 to 2; the default is 1 for 'glskrigeidw' and 'glskrigeidw'; and if it is < 1, more weight is placed on 'krige', otherwise more weight is placed on 'idw'; and if it is 0, 'idw' is not considered and the resultant methods is 'glskrige' when the default 'hybrid.parameter' is used; and if it is 2, then the resultant method is 'idw' when the default 'hybrid.parameter' is used.
...	other arguments passed on to 'gls', 'krige' and 'gstat'.

**Value**

A dataframe of longitude, latitude, and predictions.

**Author(s)**

Jin Li

**References**

Pinheiro, J. C. and D. M. Bates (2000). Mixed-Effects Models in S and S-PLUS. New York, Springer.

Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. Computers & Geosciences, 30: 683-691.

**Examples**

```
library(spm)
library(nlme)

data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
range1 <- 0.8
nugget1 <- 0.5
model <- log(gravel + 1) ~ long + lat + bathy + dist + I(long^2) + I(lat^2) +
I(lat^3) + I(bathy^2) + I(bathy^3) + I(dist^2) + I(dist^3) + I(relief^2) + I(relief^3)
y <- log(gravel[, 7] + 1)

glskrigeidwpred1 <- glskrigeidwpred(model = model, longlat = longlat, trainxy = gravel,
predx = petrel.grid, y = y, longlatpredx = petrel.grid[, c(1:2)],
transformation = "none", formula.krige = res1 ~ 1, vgm.args = "Sph", nmaxkrige = 12,
idp = 2, nmaxidw = 12, corr.args = corSpher(c(range1, nugget1),
form = ~ lat + long, nugget = T))

names(glskrigeidwpred1)

# Back transform 'glskrigeidwpred$predictions' to generate the final predictions
glskrigeidw.predictions <- exp(glskrigeidwpred1$predictions) - 1
range(glskrigeidw.predictions)
```

glskrigePred

*Generate spatial predictions using the hybrid method of generalized least squares ('gls') and kriging ('krige') ('glskrige')*

**Description**

This function is for generating spatial predictions using the hybrid method of 'gls' and 'krige' (glskrige).

**Usage**

```

glskrigepred(
  model = var1 ~ 1,
  longlat,
  trainxy,
  predx,
  y,
  longlatpredx,
  corr.args = NULL,
  weights = NULL,
  transformation = "none",
  delta = 1,
  formula.krige = res1 ~ 1,
  vgm.args = c("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  ...
)

```

**Arguments**

model	a formula defining the response variable and predictive variables.
longlat	a dataframe contains longitude and latitude of point samples.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be names as 'long' and 'lat'.
predx	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
y	a vector of the response variable in the formula, that is, the left part of the formula.
longlatpredx	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted. The location information must be named as 'long' and 'lat'.
corr.args	arguments for 'correlation' in 'gls'. See '?corClasses' in 'nlme' for details. By default, "NULL" is used. When "NULL" is used, then 'gls' is actually performing 'lm'.
weights	describing the within-group heteroscedasticity structure. Defaults to "NULL", corresponding to homoscedastic errors. See '?gls' in 'nlme' for details.
transformation	transform the residuals of 'gls' to normalize the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
delta	numeric; to avoid log(0) in the log transformation. The default is 1.

formula.krige	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.
vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
...	other arguments passed on to 'gls' and 'krige'.

**Value**

A dataframe of longitude, latitude, and predictions.

**Author(s)**

Jin Li

**References**

- Pinheiro, J. C. and D. M. Bates (2000). Mixed-Effects Models in S and S-PLUS. New York, Springer.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. Computers & Geosciences, 30: 683-691.

**Examples**

```
library(spm)
library(nlme)

data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
range1 <- 0.8
nugget1 <- 0.5
model <- log(gravel + 1) ~ long + lat + bathy + dist + I(long^2) + I(lat^2) +
I(lat^3) + I(bathy^2) + I(bathy^3) + I(dist^2) + I(dist^3) + I(relief^2) + I(relief^3)
y <- log(gravel[, 7] + 1)

glskrigePred1 <- glskrigePred(model = model, longlat = longlat, trainxy = gravel,
predx = petrel.grid, y = y, longlatpredx = petrel.grid[, c(1:2)],
```

```

transformation = "none", formula.krige = res1 ~ 1,
vgm.args = "Sph", nmaxkrige = 12, corr.args = corSpher(c(range1, nugget1),
form = ~ lat + long, nugget = T))

names(glskrigepred1)

# Back transform 'glskrigepred$predictions' to generate the final predictions
glskrige.predictions <- exp(glskrigepred1$predictions) - 1
range(glskrige.predictions)

```

---

glspred

*Generate spatial predictions using generalized least squares ('gls')*


---

## Description

This function is for generating spatial predictions using 'gls' method in 'nlme' package.

## Usage

```

glspred(
  model = var1 ~ 1,
  trainxy,
  longlatpredx,
  predx,
  corr.args = NULL,
  weights = NULL,
  ...
)

```

## Arguments

model	a formula defining the response variable and predictive variables.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be names as 'long' and 'lat'.
longlatpredx	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted, need to be named as 'long' and 'lat'.
predx	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
corr.args	arguments for 'correlation' in 'gls'. See '?corClasses' in 'nlme' for details. By default, "NULL" is used. When "NULL" is used, then 'gls' is actually performing 'lm'.
weights	describing the within-group heteroscedasticity structure. Defaults to "NULL", corresponding to homoscedastic errors. See '?gls' in 'nlme' for details.
...	other arguments passed on to 'gls'.



**Value**

A dataframe of longitude, latitude and predictions.

**Author(s)**

Jin Li

**References**

Pinheiro, J. C. and D. M. Bates (2000). *Mixed-Effects Models in S and S-PLUS*. New York, Springer.

**Examples**

```
library(spm)
library(nlme)

data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
range1 <- 0.8
nugget1 <- 0.5
model <- log(gravel + 1) ~ long + lat + bathy + dist + I(long^2) + I(lat^2)
+ I(lat^3) + I(bathy^2) + I(bathy^3) + I(dist^2) + I(dist^3) + I(relief^2) + I(relief^3)

glspred1 <- glspred(model = model, trainxy = gravel,
  longlatpredx = petrel.grid[, c(1:2)], predx = petrel.grid,
  corr.args = corSpher(c(range1, nugget1), form = ~ lat + long, nugget = T))

names(glspred1)

# Back transform 'glspred1$predictions' to generate the final predictions
gls.predictions <- exp(glspred1$predictions) - 1
range(gls.predictions)
```

---

krige cv

*Cross validation, n-fold and leave-one-out for kriging methods ('krige')*

---

**Description**

This function is a cross validation function for kriging methods ('krige') in 'gstat'.

**Usage**

```
krigecv(
  longlat,
  trainy,
  trainpredx = NULL,
  validation = "CV",
  cv.fold = 10,
  nmax = 12,
  transformation = "none",
  delta = 1,
  formula = var1 ~ 1,
  vgm.args = ("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  predacc = "VEcv",
  ...
)
```

**Arguments**

longlat	a dataframe contains longitude and latitude of point samples.
trainy	a vector of response, must have length equal to the number of rows in trainx.
trainpredx	a dataframe contains predictive variables of point samples. If longitude and latitude are going to be used as predictive variables, they should also be included but they should be named in names other than 'long' and 'lat'.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
nmax	for local kriging: the number of nearest observations that should be used for a kriging prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
transformation	transform response variable to normalise the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
delta	numeric; to avoid 'log(0)' in "log" transformation. The default is 1.
formula	formula defining response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in the 'gstat' package for details.
vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in the 'gstat' package for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in the 'gstat' package for details.
alpha	direction in plane (x,y). see variogram in the 'gstat' package for details.

block	block size. see 'krige' in the 'gstat' package for details.
beta	for simple kriging. see 'krige' in the 'gstat' package for details.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
...	other arguments passed on to the function 'gstat'.

### Value

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

### Note

This function is largely based on rfcv in 'randomForest' and some functions in 'library(gstat)'. When 'A zero or negative range was fitted to variogram' occurs, to allow 'gstat' running, the range was set to be positive by using 'min(vgm1\$dist)'. In this case, caution should be taken in applying this method. If it still occurs for 'okpred' function, different method may need to be used.

### Author(s)

Jin Li

### References

- Li, J., 2013. Predictive Modelling Using Random Forest and Its Hybrid Methods with Geostatistical Techniques in Marine Environmental Geosciences, In: Christen, P., Kennedy, P., Liu, L., Ong, K.-L., Stranieri, A., Zhao, Y. (Eds.), The proceedings of the Eleventh Australasian Data Mining Conference (AusDM 2013), Canberra, Australia, 13-15 November 2013. Conferences in Research and Practice in Information Technology, Vol. 146.
- A. Liaw and M. Wiener (2002). Classification and Regression by randomForest. R News 2(3), 18-22.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. Computers & Geosciences, 30: 683-691.

### Examples

```
library(sp)
library(spm)
data(swmud)
data(petrel)

set.seed(1234)
okcv1 <- krigevcv(longlat = swmud[, c(1,2)], trainy = swmud[, 3], nmax = 7, transformation =
"arcsine", vgm.args = ("Sph"), predacc = "VEcv")
okcv1

set.seed(1234)
skcv1 <- krigevcv(longlat = swmud[, c(1,2)], trainy = swmud[, 3], nmax = 7, transformation =
"arcsine", vgm.args = ("Sph"), predacc = "VEcv", beta = mean(swmud[, 3]))
```

```

skcv1

set.seed(1234)
ukcv1 <- krigecv(longlat = swmud[, c(1,2)], trainy = swmud[, 3], nmax = 7, transformation =
"arcsine", formula = var1 ~ long + lat, vgm.args = ("Sph"), predacc = "VEcv")
ukcv1

set.seed(1234)
okcv2 <- krigecv(longlat = swmud[, c(1,2)], trainy = swmud[, 3], validation = "L00", nmax = 7,
transformation = "arcsine", vgm.args = ("Sph"), predacc = "ALL")
okcv2

set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
okcv1 <- krigecv(longlat = petrel[, c(1,2)], trainy = petrel[, 5], nmax = 12,
transformation = "arcsine", predacc = "VEcv")
VEcv [i] <- okcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for OK", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
measures <- NULL
for (i in 1:n) {
okcv1 <- krigecv(longlat = petrel[, c(1,2)], trainy = petrel[, 3], nmax = 12, transformation =
"arcsine", predacc = "ALL")
measures <- rbind(measures, okcv1$vecv)
}
plot(measures ~ c(1:n), xlab = "Iteration for OK", ylab = "VEcv (%)")
points(cumsum(measures) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(measures), col = 'blue', lwd = 2)

```

---

krigepred

*Generate spatial predictions using kriging methods ('krige')*


---

## Description

This function is to make spatial predictions using kriging methods ('krige').

## Usage

```

krigepred(
  trainx,
  trainy,

```

```

    trainx2,
    nmax = 12,
    transformation = "none",
    delta = 1,
    formula = var1 ~ 1,
    vgm.args = ("Sph"),
    anis = c(0, 1),
    alpha = 0,
    block = 0,
    beta,
    ...
  )

```

### Arguments

trainx	a dataframe contains longitude (long), latitude (lat) and predictive variables of point samples. The location information must be named as 'long' and 'lat'.
trainy	a vector of response, must have length equal to the number of rows in trainx.
trainx2	a dataframe contains longitude (long), latitude (lat) and predictive variables of point locations (i.e., the centres of grids) to be predicted. The location information must be named as 'long' and 'lat' and in the first two columns respectively..
nmax	for local kriging: the number of nearest observations that should be used for a kriging prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
transformation	transform the response variable to normalise the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
delta	numeric; to avoid log(0) in the log transformation.
formula	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in gstat for details.
vgm.args	arguments for vgm, e.g. variogram model of response variable and anisotropy parameters. see notes vgm in gstat for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes vgm in gstat for details.
alpha	direction in plane (x,y). see variogram in gstat for details.
block	block size. see krige in gstat for details.
beta	for simple kriging. see krige in gstat for details.
...	other arguments passed on to gstat.

### Value

A dataframe of longitude, latitude, predictions and variances.

**Note**

The variances in the output are not transformed back when a transformation is used. This is because kriging variances are not measuring the uncertainty of predictions but they are indicator of the spatial distribution of sample density. The variances are exported only for interested users; and if needed, they can be transformed back from the output.

**Author(s)**

Jin Li

**References**

Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

**Examples**

```
library(sp)
library(spm)
data(swmud)
data(sw)
okpred1 <- krigepred(swmud[, c(1,2)], swmud[, 3], sw, nmax = 7, transformation =
"arcsine", vgm.args = ("Sph"))
names(okpred1)
```

---

rfkrigeidwcv

*Cross validation, n-fold and leave-one-out for the hybrid methods of 'random forest' ('RF'), 'kriging' and inverse distance weighted ('IDW')*

---

**Description**

This function is a cross validation function for 38 hybrid methods of 'RF', 'kriging' and 'IDW', including the average of 'rfkrige' and 'rfidw' ('rfkrigerfidw') and the average of 'rf', 'rfkrige' and 'rfidw' ('rfrfkrigerfidw'), where 'kriging' methods include ordinary kriging ('OK'), simple kriging ('SK'), block 'OK' ('BOK') and block 'SK' ('BSK') and 'IDW' also covers 'NN' and 'KNN'.. The data splitting is based on a stratified random sampling method (see the 'datasplit' function for details).

**Usage**

```
rfkrigeidwcv(
  longlat,
  trainx,
  trainy,
```

```

mtry = function(p) max(1, floor(sqrt(p))),
ntree = 500,
transformation = "none",
delta = 1,
formula = res1 ~ 1,
vgm.args = c("Sph"),
anis = c(0, 1),
alpha = 0,
block = 0,
beta,
nmaxkrige = 12,
idp = 2,
nmaxidw = 12,
hybrid.parameter = 2,
lambda = 1,
validation = "CV",
cv.fold = 10,
predacc = "VEcv",
...
)

```

### Arguments

longlat	a dataframe contains longitude and latitude of point samples.
trainx	a dataframe or matrix contains columns of predictive variables.
trainy	a vector of the response variable.
mtry	a function of number of remaining predictor variables to use as the 'mtry' parameter in the 'randomForest' call.
ntree	number of trees to grow. This should not be set to too small a number, to ensure that every input row gets predicted at least a few times. By default, 500 is used.
transformation	transform the residuals of 'rf' to normalize the data for 'krige'; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
delta	numeric; to avoid log(0) in the log transformation. The default is 1.
formula	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.
vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.

idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
hybrid.parameter	the default is 2 that is for 'rfkrigerfidw'; for 'rfrkrigerfidw', it needs to be 3.
lambda,	ranging from 0 to 2; the default is 1 for 'rfkrigerfidw' and 'rfrkrigerfidw'; and if it is < 1, more weight is placed on 'krige', otherwise more weight is placed on 'idw'; and if it is 0, 'idw' is not considered and the resultant methods is 'rfkrige' when the default 'hybrid.parameter' is used; and if it is 2, then the resultant method is 'rfidw' when the default 'hybrid.parameter' is used.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'randomForest', 'krige' and 'gstat'.

**Value**

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

**Note**

This function is largely based on 'rfcv' in 'randomForest', and 'krige' in 'spm2'.

**Author(s)**

Jin Li

**References**

- Li, J., Potter, A., Huang, Z., and Heap, A. (2012). Predicting Seabed Sand Content across the Australian Margin Using Machine Learning and Geostatistical Methods, Geoscience Australia, Record 2012/48, 115pp.
- Li, J., Heap, A., Potter, A., and Danilel, J.J. (2011). Predicting Seabed Mud Content across the Australian Margin II: Performance of Machine Learning Methods and Their Combination with Ordinary Kriging and Inverse Distance Squared, Geoscience Australia, Record 2011/07, 69pp.
- Liaw, A. and M. Wiener (2002). Classification and Regression by randomForest. R News 2(3), 18-22.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. Computers & Geosciences, 30: 683-691.



**Examples**

```

library(spm)
# rfrfokrfdw
data(sponge)
longlat <- sponge[, 1:2]
set.seed(1234)
rfrfkrigerfidwcv1 <- rfkrigeidwcv(longlat = longlat,
trainx = sponge[, -3], trainy = sponge[, 3], formula = res1 ~ 1, vgm.args = ("Sph"),
nmaxkrige = 12, idp = 2, nmaxidw = 12, hybrid.parameter = 3, validation = "CV",
predacc = "ALL")
rfrfkrigerfidwcv1

# rfokrfdw for count data
data(sponge)
longlat <- sponge2[, 1:2]
y = sponge[, 3]
trainx = sponge[, -3]
set.seed(1234)
n <- 20 # number of iterations,60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  rfkrigerfidwcv1 <- rfkrigeidwcv(longlat = longlat,
trainx = trainx, trainy = y, formula = res1 ~ 1, vgm.args = ("Sph"),
nmaxkrige = 12, idp = 2, nmaxidw = 12, validation = "CV", predacc = "VEcv")
  VEcv [i] <- rfkrigerfidwcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for rfokrfdw", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

```

---

rfkrigeidwpred

*Generate spatial predictions using the hybrid methods of 'random forest' ('RF'), 'kriging' and inverse distance weighted ('IDW').*

---

**Description**

This function is for generating spatial predictions using the hybrid methods of 'RF', 'kriging' and 'IDW', including all methods implemented in 'rfkrigeidwcv'.

**Usage**

```

rfkrigeidwpred(
  longlat,
  trainx,
  predx,
  trainy,

```

```

longlatpredx,
mtry = function(p) max(1, floor(sqrt(p))),
ntree = 500,
transformation = "none",
delta = 1,
formula.krige = res1 ~ 1,
vgm.args = c("Sph"),
anis = c(0, 1),
alpha = 0,
block = 0,
beta,
nmaxkrige = 12,
idp = 2,
nmaxidw = 12,
hybrid.parameter = 2,
lambda = 1,
...
)

```

### Arguments

longlat	a dataframe contains longitude and latitude of point samples.
trainx	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be named as 'long' and 'lat'.
predx	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
trainy	a vector of the response variable in the formula, that is, the left part of the formula.
longlatpredx	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted.
mtry	a function of number of remaining predictor variables to use as the 'mtry' parameter in the 'randomForest' call.
ntree	number of trees to grow. This should not be set to too small a number, to ensure that every input row gets predicted at least a few times. By default, 500 is used. #' @param longlatpredx a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted. The location information must be named as 'long' and 'lat'.
transformation	transform the residuals of 'rf' to normalise the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
delta	numeric; to avoid log(0) in the log transformation. The default is 1.
formula.krige	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.

vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
hybrid.parameter	the default is 2 that is for 'rfkrigerfidw'; for 'rfrkrigerfidw', it needs to be 3.
lambda,	ranging from 0 to 2; the default is 1 for 'rfkrigerfidw' and 'rfrkrigerfidw'; and if it is < 1, more weight is placed on 'krige', otherwise more weight is placed on 'idw'; and if it is 0, 'idw' is not considered and the resultant methods is 'rfkrige' when the default 'hybrid.parameter' is used; and if it is 2, then the resultant method is 'rfidw' when the default 'hybrid.parameter' is used.
...	other arguments passed on to 'rf', 'krige' and 'gstat'.

**Value**

A dataframe of longitude, latitude, and predictions.

**Author(s)**

Jin Li

**References**

- Li, J., Potter, A., Huang, Z., and Heap, A. (2012). Predicting Seabed Sand Content across the Australian Margin Using Machine Learning and Geostatistical Methods, Geoscience Australia, Record 2012/48, 115pp.
- Li, J., Heap, A., Potter, A., and Danilel, J.J. (2011). Predicting Seabed Mud Content across the Australian Margin II: Performance of Machine Learning Methods and Their Combination with Ordinary Kriging and Inverse Distance Squared, Geoscience Australia, Record 2011/07, 69pp.
- Liaw, A. and M. Wiener (2002). Classification and Regression by randomForest. R News 2(3), 18-22.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. Computers & Geosciences, 30: 683-691.

**Examples**

```

library(spm)

data(sponge)
data(sponge.grid)
longlat <- sponge[, 1:2]
y = sponge[, 3]
trainx = sponge[, -3]

set.seed(1234)

rfkrigeidwpred1 <- rfkrigeidwpred(longlat = longlat, trainx = trainx,
predx = sponge.grid, trainy = y, longlatpredx = sponge.grid[, c(1:2)],
formula.krige = res1 ~ 1, vgm.args = "Sph", nmaxkrige = 12, idp = 2, nmaxidw = 12)

names(rfkrigeidwpred1)

range(rfkrigeidwpred1$predictions)

```

---

sponge2

*A dataset of sponge species richness in the Timor Sea region, northern Australia marine margin*

---

**Description**

This dataset contains 77 samples of 81 variables including easting (longitude), northing (latitude), bathy, backscatter and their derived variables.

**Usage**

```
data("sponge2")
```

**Format**

A data frame with 77 observations on the following 89 variables.

easting a numeric vector, m  
northing a numeric vector, m  
species.richness a numeric vector, no unit  
mud a numeric vector, percentage  
sand a numeric vector, percentage  
gravel a numeric vector, percentage  
bathy a numeric vector, m

bs25 a numeric vector, dB  
bs10 a numeric vector, dB  
bs11 a numeric vector, dB  
bs12 a numeric vector, dB  
bs13 a numeric vector, dB  
bs14 a numeric vector, dB  
bs15 a numeric vector, dB  
bs16 a numeric vector, dB  
bs17 a numeric vector, dB  
bs18 a numeric vector, dB  
bs19 a numeric vector, dB  
bs20 a numeric vector, dB  
bs21 a numeric vector, dB  
bs22 a numeric vector, dB  
bs23 a numeric vector, dB  
bs24 a numeric vector, dB  
bs26 a numeric vector, dB  
bs27 a numeric vector, dB  
bs28 a numeric vector, dB  
bs29 a numeric vector, dB  
bs30 a numeric vector, dB  
bs31 a numeric vector, dB  
bs32 a numeric vector, dB  
bs33 a numeric vector, dB  
bs34 a numeric vector, dB  
bs35 a numeric vector, dB  
bs36 a numeric vector, dB  
bs\_o a numeric vector, dB  
bs\_homo\_o a numeric vector  
bs\_entro\_o a numeric vector, no unit  
bs\_var\_o a numeric vector, dB<sup>2</sup>  
bs\_lmi\_o a numeric vector  
bathy\_o a numeric vector, m  
bathy\_lmi\_o a numeric vector  
tpi\_o a numeric vector, no unit  
slope\_o a numeric vector  
plan\_cur\_o a numeric vector

prof\_cur\_o a numeric vector  
relief\_o a numeric vector  
rugosity\_o a numeric vector  
dist.coast a numeric vector, m  
rugosity3 a numeric vector  
rugosity5 a numeric vector  
rugosity7 a numeric vector  
tpi3 a numeric vector, no unit  
tpi5 a numeric vector, no unit  
tpi7 a numeric vector, no unit  
bathy\_lmi3 a numeric vector  
bathy\_lmi5 a numeric vector  
bathy\_lmi7 a numeric vector  
plan\_curv3 a numeric vector  
plan\_curv5 a numeric vector  
plan\_curv7 a numeric vector  
relief\_3 a numeric vector  
relief\_5 a numeric vector  
relief\_7 a numeric vector  
slope3 a numeric vector  
slope5 a numeric vector  
slope7 a numeric vector  
prof\_cur3 a numeric vector  
prof\_cur5 a numeric vector  
prof\_cur7 a numeric vector  
entro3 a numeric vector, no unit  
entro5 a numeric vector, no unit  
entro7 a numeric vector, no unit  
homo3 a numeric vector  
homo5 a numeric vector  
homo7 a numeric vector  
var3 a numeric vector, dB<sup>2</sup>  
var5 a numeric vector, dB<sup>2</sup>  
var7 a numeric vector, dB<sup>2</sup>  
bs\_lmi3 a numeric vector  
bs\_lmi5 a numeric vector  
bs\_lmi7 a numeric vector

**Details**

For details, please see the source. This dataset was published as an appendix of the paper listed in the source. Where the long and lat were reprojected to easting and northing.

**Source**

see Appendix A-D. Supplementary data at: "<http://dx.doi.org/10.1016/j.envsoft.2017.07.016>."

**References**

Li, J., B. Alvarez, J. Siwabessy, M. Tran, Z. Huang, R. Przeslawski, L. Radke, F. Howard, and S. Nichol. 2017. Application of random forest, generalised linear model and their hybrid methods with geostatistical techniques to count data: Predicting sponge species richness. *Environmental Modelling & Software*, 97: 112-129.

---

spongelonglat	<i>A dataset of sponge species richness in the Timor Sea region, northern Australia marine margin</i>
---------------	---

---

**Description**

This dataset contains 77 samples of 7 predictive variables including longitude, latitude, bathy, backscatter and their derived variables. It is the sponge dataset in 'spm' package, but with long and lat instead of easting and northing.

**Usage**

```
data("spongelonglat")
```

**Format**

A data frame with 77 observations on the following 8 variables.

sponge a numeric vector  
tpi3 a numeric vector  
var7 a numeric vector  
entro7 a numeric vector  
bs34 a numeric vector  
bs11 a numeric vector  
long a numeric vector  
lat a numeric vector

**Details**

For details, please see sponge dataset in library(spm). Where the long and lat were projected to easting and northing.

**Source**

sponge dataset in library(spm)

**References**

Li, J., B. Alvarez, J. Siwabessy, M. Tran, Z. Huang, R. Przeslawski, L. Radke, F. Howard, and S. Nichol. 2017. Application of random forest, generalised linear model and their hybrid methods with geostatistical techniques to count data: Predicting sponge species richness. *Environmental Modelling & Software*, 97: 112-129.

**Examples**

```
data(spongelonglat)
## maybe str(spongelonglat) ; plot(spongelonglat) ...
```

---

svmcv

*Cross validation, n-fold and leave-one-out for support vector machine ('svm')*

---

**Description**

This function is a cross validation function for 'svm' regression in 'e1071' package.

**Usage**

```
svmcv(  
  formula = NULL,  
  trainxy,  
  y,  
  scale = TRUE,  
  type = NULL,  
  kernel = "radial",  
  degree = 3,  
  gamma = if (is.vector(trainxy)) 1 else 1/ncol(trainxy),  
  coef0 = 0,  
  cost = 1,  
  nu = 0.5,  
  tolerance = 0.001,  
  epsilon = 0.1,  
  validation = "CV",  
  cv.fold = 10,  
  predacc = "VEcv",  
  ...  
)
```



**Arguments**

formula	a formula defining the response variable and predictive variables.
trainxy	a dataframe contains predictive variables and the response variable of point samples. The location information, longitude (long), latitude (lat), need to be included in the 'trainx' for spatial predictive modelling, need to be named as 'long' and 'lat'.
y	a vector of the response variable in the formula, that is, the left part of the formula.
scale	A logical vector indicating the variables to be scaled (default: TRUE).
type	the default setting is 'NULL'. See '?svm' for various options.
kernel	the default setting is 'radial'. See '?svm' for other options.
degree	a parameter needed for kernel of type polynomial (default: 3).
gamma	a parameter needed for all 'kernels' except 'linear' (default: 1/(data dimension)).
coef0	a parameter needed for kernels of type 'polynomial' and 'sigmoid' (default: 0).
cost	cost of constraints violation (default: 1).
nu	a parameter needed for 'nu-classification', 'nu-regression', and 'one-classification' (default: 0.5).
tolerance	tolerance of termination criterion (default: 0.001).
epsilon	'epsilon' in the insensitive-loss function (default: 0.1).
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for 'vecv' or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'svm'.

**Value**

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

**Note**

This function is largely based on 'rfcv' in 'randomForest' and 'svm' in 'e1071'.

**Author(s)**

Jin Li

**References**

David Meyer, Evgenia Dimitriadou, Kurt Hornik, Andreas Weingessel and Friedrich Leisch (2020). e1071: Misc Functions of the Department of Statistics, Probability Theory Group (Formerly: E1071), TU Wien. R package version 1.7-4. <https://CRAN.R-project.org/package=e1071>.

## Examples

```

library(spm)

data(petrel)
gravel <- petrel[, c(1, 2, 6:9, 5)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
set.seed(1234)
svmcv1 <- svmcv(formula = model, gravel, log(gravel[, 7] +1), validation = "CV",
  predacc = "ALL")
svmcv1

data(sponge2)
model <- species.richness ~ .
set.seed(1234)
svmcv1 <- svmcv(formula = model, sponge2[, -4], sponge[, 3], gamma = 0.01, cost = 3.5,
  scale = TRUE, validation = "CV", predacc = "VEcv")
svmcv1

# For svm
model <- species.richness ~ .
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  svmcv1 <- svmcv(formula = model, sponge2[, -4], sponge[, 3], gamma = 0.01, cost = 3.5,
    scale = TRUE, validation = "CV", predacc = "VEcv")
  VEcv [i] <- svmcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for svm", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

```

---

 svmidwcv

*Cross validation, n-fold and leave-one-out for the hybrid method of support vector machine ('svm') regression and inverse distance weighted ('IDW') (svmidw)*

---

## Description

This function is a cross validation function for the hybrid method of 'svm' regression and 'idw' using 'gstat' (svmidw), where the data splitting is based on a stratified random sampling method (see the 'datasplit' function for details).

## Usage

```
svmidwcv(
```

```

    formula = NULL,
    longlat,
    trainxy,
    y,
    scale = TRUE,
    type = NULL,
    kernel = "radial",
    degree = 3,
    gamma = if (is.vector(trainxy)) 1 else 1/ncol(trainxy),
    coef0 = 0,
    cost = 1,
    nu = 0.5,
    tolerance = 0.001,
    epsilon = 0.1,
    idp = 2,
    nmaxidw = 12,
    validation = "CV",
    cv.fold = 10,
    predacc = "VEcv",
    ...
)

```

### Arguments

formula	a formula defining the response variable and predictive variables for 'svm'.
longlat	a dataframe contains longitude and latitude of point samples.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be named as 'long' and 'lat'.
y	a vector of the response variable in the formula, that is, the left part of the formula.
scale	A logical vector indicating the variables to be scaled (default: TRUE).
type	the default setting is 'NULL'. See '?svm' for various options.
kernel	the default setting is 'radial'. See '?svm' for other options.
degree	a parameter needed for kernel of type polynomial (default: 3).
gamma	a parameter needed for all 'kernels' except 'linear' (default: 1/(data dimension)).
coef0	a parameter needed for kernels of type 'polynomial' and 'sigmoid'(default: 0).
cost	cost of constraints violation (default: 1).
nu	a parameter needed for 'nu-classification', 'nu-regression', and 'one-classification' (default: 0.5).
tolerance	tolerance of termination criterion (default: 0.001).
epsilon	'epsilon' in the insensitive-loss function (default: 0.1).
idp	a numeric number specifying the inverse distance weighting power.

nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for 'vecv' or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'svm' and 'gstat'.

### Value

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only.

### Note

This function is largely based on 'rfcv' in 'randomForest', 'idwcv' in 'spm' and 'svm' in 'e1071'.

### Author(s)

Jin Li

### References

- Li, J., Potter, A., Huang, Z., and Heap, A. (2012). Predicting Seabed Sand Content across the Australian Margin Using Machine Learning and Geostatistical Methods, *Geoscience Australia, Record 2012/48*, 115pp.
- Li, J., Heap, A., Potter, A., and Danilel, J.J. (2011). Predicting Seabed Mud Content across the Australian Margin II: Performance of Machine Learning Methods and Their Combination with Ordinary Kriging and Inverse Distance Squared, *Geoscience Australia, Record 2011/07*, 69pp.
- David Meyer, Evgenia Dimitriadou, Kurt Hornik, Andreas Weingessel and Friedrich Leisch (2020). e1071: Misc Functions of the Department of Statistics, Probability Theory Group (Formerly: E1071), TU Wien. R package version 1.7-4. <https://CRAN.R-project.org/package=e1071>.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

### Examples

```
library(spm)

data(petrel)
gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)

set.seed(1234)
```

```
svmidwcv1 <- svmidwcv(formula = model, longlat = longlat, trainxy = gravel,
y = y, idp = 2, nmaxidw = 12, validation = "CV", predacc = "ALL")
svmidwcv1

# svmidw for count data
data(sponge2)
model <- species.richness ~ . # use all predictive variables in the dataset
longlat <- sponge2[, 1:2]
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  svmidwcv1 <- svmidwcv(formula = model, longlat = longlat, trainxy = sponge2[, -4],
y = sponge[, 3], gamma = 0.01, cost = 3.5, scale = TRUE, idp = 2, nmaxidw = 12,
validation = "CV", predacc = "VEcv")
  VEcv [i] <- svmidwcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for svmidw", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)
```

---

svmidwpred

*Generate spatial predictions using the hybrid method of support vector machine ('svm') regression and inverse distance weighted ('IDW') ('svmidw')*

---

## Description

This function is for generating spatial predictions using the hybrid method of 'svm' and 'idw' ('svmidw').

## Usage

```
svmidwpred(
  formula = NULL,
  longlat,
  trainxy,
  y,
  longlatpredx,
  predx,
  scale = TRUE,
  type = NULL,
  kernel = "radial",
  degree = 3,
  gamma = if (is.vector(trainxy)) 1 else 1/ncol(trainxy),
  coef0 = 0,
  cost = 1,
```

```

    nu = 0.5,
    tolerance = 0.001,
    epsilon = 0.1,
    idp = 2,
    nmaxidw = 12,
    ...
)

```

### Arguments

formula	a formula defining the response variable and predictive variables for 'svm'.
longlat	a dataframe contains longitude and latitude of point samples. The location information must be named as 'long' and 'lat'.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples.
y	a vector of the response variable in the formula, that is, the left part of the formula.
longlatpredx	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted.
predx	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
scale	A logical vector indicating the variables to be scaled (default: TRUE).
type	the default setting is 'NULL'. See '?svm' for various options.
kernel	the default setting is 'radial'. See '?svm' for other options.
degree	a parameter needed for kernel of type polynomial (default: 3).
gamma	a parameter needed for all 'kernels' except 'linear' (default: 1/(data dimension)).
coef0	a parameter needed for kernels of type 'polynomial' and 'sigmoid' (default: 0).
cost	cost of constraints violation (default: 1).
nu	a parameter needed for 'nu-classification', 'nu-regression', and 'one-classification' (default: 0.5).
tolerance	tolerance of termination criterion (default: 0.001).
epsilon	'epsilon' in the insensitive-loss function (default: 0.1). See '?svm' for details.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
...	other arguments passed on to 'svm'.

### Value

A dataframe of longitude, latitude, and predictions.

### Author(s)

Jin Li

## References

- Li, J., Potter, A., Huang, Z., and Heap, A. (2012). Predicting Seabed Sand Content across the Australian Margin Using Machine Learning and Geostatistical Methods, *Geoscience Australia, Record 2012/48*, 115pp.
- Li, J., Heap, A., Potter, A., and Danilel, J.J. (2011). Predicting Seabed Mud Content across the Australian Margin II: Performance of Machine Learning Methods and Their Combination with Ordinary Kriging and Inverse Distance Squared, *Geoscience Australia, Record 2011/07*, 69pp.
- David Meyer, Evgenia Dimitriadou, Kurt Hornik, Andreas Weingessel and Friedrich Leisch (2020). e1071: Misc Functions of the Department of Statistics, Probability Theory Group (Formerly: E1071), TU Wien. R package version 1.7-4. <https://CRAN.R-project.org/package=e1071>.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

## Examples

```
library(spm)
data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)

svmidwpred1 <- svmidwpred(formula = model, longlat = longlat, trainxy = gravel,
y = y, longlatpredx = petrel.grid[, c(1:2)], predx = petrel.grid, idp = 2,
nmaxidw = 12)

names(svmidwpred1)

# Back transform 'svmidwpred$predictions' to generate the final predictions
svmidw.predictions <- exp(svmidwpred1$predictions) - 1
range(svmidw.predictions)
```

---

svmkrige cv

---

*Cross validation, n-fold and leave-one-out for the hybrid method of support vector machine ('svm') regression and 'krige' (svmkrige)*


---

## Description

This function is a cross validation function for the hybrid method of 'svm' regression and 'krige' (svmkrige), where the data splitting is based on a stratified random sampling method (see the 'datasplit' function for details).

**Usage**

```
svmkridgecv(
  formula.svm = NULL,
  longlat,
  trainxy,
  y,
  scale = TRUE,
  type = NULL,
  kernel = "radial",
  degree = 3,
  gamma = if (is.vector(trainxy)) 1 else 1/ncol(trainxy),
  coef0 = 0,
  cost = 1,
  nu = 0.5,
  tolerance = 0.001,
  epsilon = 0.1,
  transformation = "none",
  delta = 1,
  formula.krige = res1 ~ 1,
  vgm.args = c("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  validation = "CV",
  cv.fold = 10,
  predacc = "VEcv",
  ...
)
```

**Arguments**

<code>formula.svm</code>	a formula defining the response variable and predictive variables for 'svm'.
<code>longlat</code>	a dataframe contains longitude and latitude of point samples.
<code>trainxy</code>	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be named as 'long' and 'lat'.
<code>y</code>	a vector of the response variable in the formula.svm, that is, the left part of the formula.svm.
<code>scale</code>	A logical vector indicating the variables to be scaled (default: TRUE).
<code>type</code>	the default setting is 'NULL'. See '?svm' for various options.
<code>kernel</code>	the default setting is 'radial'. See '?svm' for other options.
<code>degree</code>	a parameter needed for kernel of type polynomial (default: 3).
<code>gamma</code>	a parameter needed for all 'kernels' except 'linear' (default: 1/(data dimension)).
<code>coef0</code>	a parameter needed for kernels of type 'polynomial' and 'sigmoid'(default: 0).



cost	cost of constraints violation (default: 1).
nu	a parameter needed for 'nu-classification', 'nu-regression', and 'one-classification' (default: 0.5).
tolerance	tolerance of termination criterion (default: 0.001).
epsilon	'epsilon' in the insensitive-loss function (default: 0.1).
transformation	transform the residuals of 'svm' to normalise the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
delta	numeric; to avoid log(0) in the log transformation. The default is 1.
formula.krige	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.
vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for 'vecv' or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'svm' and 'krige'.

**Value**

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

**Note**

This function is largely based on 'rfcv' in 'randomForest', 'krigev' in 'spm2' and 'svm' in 'e1071'.

**Author(s)**

Jin Li

## References

- Li, J., Potter, A., Huang, Z., and Heap, A. (2012). Predicting Seabed Sand Content across the Australian Margin Using Machine Learning and Geostatistical Methods, *Geoscience Australia, Record 2012/48*, 115pp.
- Li, J., Heap, A., Potter, A., and Danilel, J.J. (2011). Predicting Seabed Mud Content across the Australian Margin II: Performance of Machine Learning Methods and Their Combination with Ordinary Kriging and Inverse Distance Squared, *Geoscience Australia, Record 2011/07*, 69pp.
- David Meyer, Evgenia Dimitriadou, Kurt Hornik, Andreas Weingessel and Friedrich Leisch (2020). e1071: Misc Functions of the Department of Statistics, Probability Theory Group (Formerly: E1071), TU Wien. R package version 1.7-4. <https://CRAN.R-project.org/package=e1071>.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

## Examples

```
library(spm)

data(petrel)

gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)
set.seed(1234)
svmkrigecv1 <- svmkrigecv(formula.svm = model, longlat = longlat, trainxy = gravel,
y = y, transformation = "none", formula.krige = res1 ~ 1, vgm.args = "Sph",
nmaxkrige = 12, validation = "CV", predacc = "ALL")
svmkrigecv1

# svmok for count data
data(sponge2)
model <- species.richness ~ . # use all predictive variables in the dataset
longlat <- sponge2[, 1:2]
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  svmkrigecv1 <- svmkrigecv(formula.svm = model, longlat = longlat, trainxy = sponge2[, -4],
y = sponge2[, 3], gamma = 0.01, cost = 3.5, scale = TRUE, formula.krige = res1 ~ 1,
vgm.args = ("Sph"), nmaxkrige = 12, validation = "CV", predacc = "VEcv")
  VEcv [i] <- svmkrigecv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for svm", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)
```

---

svmkridgeidwcv	<i>Cross validation, n-fold and leave-one-out for the hybrid methods of support vector machine ('svm') regression, 'kriging' and inverse distance weighted ('IDW').</i>
----------------	---

---

### Description

This function is a cross validation function for 38 hybrid methods of 'svm', 'kriging' and 'IDW', including the average of 'svmkridge' and 'svmidw' ('svmkridgesvmidw') and the average of 'svm', 'svmkridge' and 'svmidw' ('svmsvmkridgesvmidw'), where 'kriging' methods include ordinary kriging ('OK'), simple kriging ('SK'), block 'OK' ('BOK') and block 'SK' ('BSK') and 'IDW' also covers 'NN' and 'KNN'.. The data splitting is based on a stratified random sampling method (see the 'datasplit' function for details).

### Usage

```
svmkridgeidwcv(
  formula.svm = NULL,
  longlat,
  trainxy,
  y,
  scale = TRUE,
  type = NULL,
  kernel = "radial",
  degree = 3,
  gamma = if (is.vector(trainxy)) 1 else 1/ncol(trainxy),
  coef0 = 0,
  cost = 1,
  nu = 0.5,
  tolerance = 0.001,
  epsilon = 0.1,
  transformation = "none",
  delta = 1,
  formula.krige = res1 ~ 1,
  vgm.args = c("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  idp = 2,
  nmaxidw = 12,
  hybrid.parameter = 2,
  lambda = 1,
  validation = "CV",
  cv.fold = 10,
  predacc = "VEcv",
```

```
    ...
  )
```

### Arguments

<code>formula.svm</code>	a formula defining the response variable and predictive variables for 'svm'.
<code>longlat</code>	a dataframe contains longitude and latitude of point samples.
<code>trainxy</code>	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples. That is, the location information must be named as 'long' and 'lat'.
<code>y</code>	a vector of the response variable in the formula, that is, the left part of the formula.
<code>scale</code>	A logical vector indicating the variables to be scaled (default: TRUE).
<code>type</code>	the default setting is 'NULL'. See '?svm' for various options.
<code>kernel</code>	the default setting is 'radial'. See '?svm' for other options.
<code>degree</code>	a parameter needed for kernel of type polynomial (default: 3).
<code>gamma</code>	a parameter needed for all 'kernels' except 'linear' (default: 1/(data dimension)).
<code>coef0</code>	a parameter needed for kernels of type 'polynomial' and 'sigmoid' (default: 0).
<code>cost</code>	cost of constraints violation (default: 1).
<code>nu</code>	a parameter needed for 'nu-classification', 'nu-regression', and 'one-classification' (default: 0.5).
<code>tolerance</code>	tolerance of termination criterion (default: 0.001).
<code>epsilon</code>	'epsilon' in the insensitive-loss function (default: 0.1).
<code>transformation</code>	transform the residuals of 'svm' to normalise the data for 'krige'; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
<code>delta</code>	numeric; to avoid log(0) in the log transformation. The default is 1.
<code>formula.krige</code>	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.
<code>vgm.args</code>	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
<code>anis</code>	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
<code>alpha</code>	direction in plane (x,y). see variogram in 'gstat' for details.
<code>block</code>	block size. see 'krige' in 'gstat' for details.
<code>beta</code>	for simple kriging. see 'krige' in 'gstat' for details.
<code>nmaxkrige</code>	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
<code>idp</code>	a numeric number specifying the inverse distance weighting power.

nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
hybrid.parameter	the default is 2 that is for 'svmkrigesvmidw'; for 'svmsvmkrigesvmidw', it needs to be 3.
lambda,	ranging from 0 to 2; the default is 1 for 'svmkrigesvmidw' and 'svmsvmkrigesvmidw'; and if it is < 1, more weight is placed on 'krige', otherwise more weight is placed on 'idw'; and if it is 0, 'idw' is not considered and the resultant methods is 'svmkrige' when the default 'hybrid.parameter' is used; and if it is 2, then the resultant method is 'svmidw' when the default 'hybrid.parameter' is used.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'svm', 'krige' and 'gstat'.

**Value**

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

**Note**

This function is largely based on 'rfcv' in 'randomForest', 'krige' in 'spm2' and 'svm' in 'e1071'.

**Author(s)**

Jin Li

**References**

- Li, J., Potter, A., Huang, Z., and Heap, A. (2012). Predicting Seabed Sand Content across the Australian Margin Using Machine Learning and Geostatistical Methods, *Geoscience Australia, Record 2012/48*, 115pp.
- Li, J., Heap, A., Potter, A., and Danilel, J.J. (2011). Predicting Seabed Mud Content across the Australian Margin II: Performance of Machine Learning Methods and Their Combination with Ordinary Kriging and Inverse Distance Squared, *Geoscience Australia, Record 2011/07*, 69pp.
- David Meyer, Evgenia Dimitriadou, Kurt Hornik, Andreas Weingessel and Friedrich Leisch (2020). e1071: Misc Functions of the Department of Statistics, Probability Theory Group (Formerly: E1071), TU Wien. R package version 1.7-4. <https://CRAN.R-project.org/package=e1071>.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

**Examples**

```

library(spm)
# svmokglidw
data(petrel)
gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)
set.seed(1234)
svmkrigesvmidwcv1 <- svmkrigeidwcv(formula.svm = model, longlat = longlat,
trainxy = gravel, y = y, transformation = "none", formula.krige = res1 ~ 1,
vgm.args = "Sph", nmaxkrige = 12, idp = 2, nmaxidw = 12, validation = "CV",
predacc = "ALL")
svmkrigesvmidwcv1

# svmsvmoksvmidw
data(sponge2)
model <- species.richness ~ . # use all predictive variables in the dataset
longlat <- sponge2[, 1:2]
y = sponge[, 3]
set.seed(1234)
svmsvmkrigesvmidwcv1 <- svmkrigeidwcv(formula.svm = model, longlat = longlat,
trainxy = sponge2[, -4], y = y, gamma = 0.01, cost = 3.5, scale = TRUE,
formula.krige = res1 ~ 1, vgm.args = ("Sph"), nmaxkrige = 12, idp = 2,
nmaxidw = 12, hybrid.parameter = 3, validation = "CV", predacc = "ALL")
svmsvmkrigesvmidwcv1

# svmoksvmidw for count data
data(sponge2)
model <- species.richness ~ . # use all predictive variables in the dataset
longlat <- sponge2[, 1:2]
y = sponge2[, 3]
set.seed(1234)
n <- 20 # number of iterations, 60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  svmkrigesvmidwcv1 <- svmkrigeidwcv(formula.svm = model, longlat = longlat,
trainxy = sponge2[, -4], y = y, gamma = 0.01, cost = 3.5, scale = TRUE,
formula.krige = res1 ~ 1, vgm.args = ("Sph"), nmaxkrige = 12, idp = 2,
nmaxidw = 12, validation = "CV", predacc = "VEcv")
  VEcv [i] <- svmkrigesvmidwcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for svm", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

```

---

svmkrideidwpred      *Generate spatial predictions using the hybrid methods of support vector machine ('svm') regression , 'kriging' and inverse distance weighted ('IDW').*

---

### Description

This function is for generating spatial predictions using the hybrid methods of 'svm', 'kriging' and 'IDW', including all methods implemented in 'svmkrideidwcv'.

### Usage

```
svmkrideidwpred(
  formula.svm = NULL,
  longlat,
  trainxy,
  predx,
  y,
  longlatpredx,
  scale = TRUE,
  type = NULL,
  kernel = "radial",
  degree = 3,
  gamma = if (is.vector(trainxy)) 1 else 1/ncol(trainxy),
  coef0 = 0,
  cost = 1,
  nu = 0.5,
  tolerance = 0.001,
  epsilon = 0.1,
  transformation = "none",
  delta = 1,
  formula.krige = res1 ~ 1,
  vgm.args = c("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  idp = 2,
  nmaxidw = 12,
  hybrid.parameter = 2,
  lambda = 1,
  ...
)
```

### Arguments

formula.svm      a formula defining the response variable and predictive variables for 'svm'.

longlat	a dataframe contains longitude and latitude of point samples.
trainxy	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples.
predx	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
y	a vector of the response variable in the formula, that is, the left part of the formula.
longlatpredx	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted.
scale	A logical vector indicating the variables to be scaled (default: TRUE).
type	the default setting is 'NULL'. See '?svm' for various options.
kernel	the default setting is 'radial'. See '?svm' for other options.
degree	a parameter needed for kernel of type polynomial (default: 3).
gamma	a parameter needed for all 'kernels' except 'linear' (default: 1/(data dimension)).
coef0	a parameter needed for kernels of type 'polynomial' and 'sigmoid' (default: 0).
cost	cost of constraints violation (default: 1).
nu	a parameter needed for 'nu-classification', 'nu-regression', and 'one-classification' (default: 0.5).
tolerance	tolerance of termination criterion (default: 0.001).
epsilon	'epsilon' in the insensitive-loss function (default: 0.1). See '?svm' for details.
transformation	transform the residuals of 'svm' to normalise the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
delta	numeric; to avoid log(0) in the log transformation. The default is 1.
formula.krige	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.
vgm.args	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
anis	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
alpha	direction in plane (x,y). see variogram in 'gstat' for details.
block	block size. see 'krige' in 'gstat' for details.
beta	for simple kriging. see 'krige' in 'gstat' for details.
nmaxkrige	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
idp	a numeric number specifying the inverse distance weighting power.
nmaxidw	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.



hybrid.parameter  
the default is 2 that is for 'svmkrigesvmidw'; for 'svmsvmkrigesvmidw', it needs to be 3.

lambda,  
ranging from 0 to 2; the default is 1 for 'svmkrigesvmidw' and 'svmsvmkrigesvmidw'; and if it is < 1, more weight is placed on 'krige', otherwise more weight is placed on 'idw'; and if it is 0, 'idw' is not considered and the resultant methods is 'svmkrige' when the default 'hybrid.parameter' is used; and if it is 2, then the resultant method is 'svmidw' when the default 'hybrid.parameter' is used.

...  
other arguments passed on to 'svm', 'krige' and 'gstat'.

**Value**

A dataframe of longitude, latitude, and predictions.

**Author(s)**

Jin Li

**References**

- Li, J., Potter, A., Huang, Z., and Heap, A. (2012). Predicting Seabed Sand Content across the Australian Margin Using Machine Learning and Geostatistical Methods, Geoscience Australia, Record 2012/48, 115pp.
- Li, J., Heap, A., Potter, A., and Danilel, J.J. (2011). Predicting Seabed Mud Content across the Australian Margin II: Performance of Machine Learning Methods and Their Combination with Ordinary Kriging and Inverse Distance Squared, Geoscience Australia, Record 2011/07, 69pp.
- David Meyer, Evgenia Dimitriadou, Kurt Hornik, Andreas Weingessel and Friedrich Leisch (2020). e1071: Misc Functions of the Department of Statistics, Probability Theory Group (Formerly: E1071), TU Wien. R package version 1.7-4. <https://CRAN.R-project.org/package=e1071>.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. Computers & Geosciences, 30: 683-691.

**Examples**

```
library(spm)

data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)

svmkridgeidwpred1 <- svmkridgeidwpred(formula.svm = model, longlat = longlat, trainxy = gravel,
predx = petrel.grid, y = y, longlatpredx = petrel.grid[, c(1:2)],
formula.krige = res1 ~ 1, vgm.args = "Sph", nmaxkrige = 12, idp = 2, nmaxidw = 12)
```

```
names(svmkrigeidwpred1)

# Back transform 'svmkridgeidwpred$predictions' to generate the final predictions
svmkridgeidw.predictions <- exp(svmkrigeidwpred1$predictions) - 1
range(svmkrigeidw.predictions)
```

---

svmkridgepred	<i>Generate spatial predictions using the hybrid method of support vector machine ('svm') regression and 'krige' (svmkridge)</i>
---------------	--

---

### Description

This function is for generating spatial predictions using the hybrid method of 'svm' and 'krige' (svmkridge).

### Usage

```
svmkridgepred(
  formula.svm = NULL,
  longlat,
  trainxy,
  predx,
  y,
  longlatpredx,
  scale = TRUE,
  type = NULL,
  kernel = "radial",
  degree = 3,
  gamma = if (is.vector(trainxy)) 1 else 1/ncol(trainxy),
  coef0 = 0,
  cost = 1,
  nu = 0.5,
  tolerance = 0.001,
  epsilon = 0.1,
  transformation = "none",
  delta = 1,
  formula.krige = res1 ~ 1,
  vgm.args = c("Sph"),
  anis = c(0, 1),
  alpha = 0,
  block = 0,
  beta,
  nmaxkrige = 12,
  ...
)
```

**Arguments**

<code>formula.svm</code>	a formula defining the response variable and predictive variables for 'svm'.
<code>longlat</code>	a dataframe contains longitude and latitude of point samples.
<code>trainxy</code>	a dataframe contains longitude (long), latitude (lat), predictive variables and the response variable of point samples.
<code>predx</code>	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
<code>y</code>	a vector of the response variable in the formula, that is, the left part of the formula.
<code>longlatpredx</code>	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted.
<code>scale</code>	A logical vector indicating the variables to be scaled (default: TRUE).
<code>type</code>	the default setting is 'NULL'. See '?svm' for various options.
<code>kernel</code>	the default setting is 'radial'. See '?svm' for other options.
<code>degree</code>	a parameter needed for kernel of type polynomial (default: 3).
<code>gamma</code>	a parameter needed for all 'kernels' except 'linear' (default: 1/(data dimension)).
<code>coef0</code>	a parameter needed for kernels of type 'polynomial' and 'sigmoid'(default: 0).
<code>cost</code>	cost of constraints violation (default: 1).
<code>nu</code>	a parameter needed for 'nu-classification', 'nu-regression', and 'one-classification' (default: 0.5).
<code>tolerance</code>	tolerance of termination criterion (default: 0.001).
<code>epsilon</code>	'epsilon' in the insensitive-loss function (default: 0.1). See '?svm' for details.
<code>transformation</code>	transform the residuals of 'svm' to normalise the data; can be "sqrt" for square root, "arcsine" for arcsine, "log" or "none" for non transformation. By default, "none" is used.
<code>delta</code>	numeric; to avoid log(0) in the log transformation. The default is 1.
<code>formula.krige</code>	formula defining the response vector and (possible) regressor. an object (i.e., 'variogram.formula') for 'variogram' or a formula for 'krige'. see 'variogram' and 'krige' in 'gstat' for details.
<code>vgm.args</code>	arguments for 'vgm', e.g. variogram model of response variable and anisotropy parameters. see 'vgm' in 'gstat' for details. By default, "Sph" is used.
<code>anis</code>	anisotropy parameters: see notes 'vgm' in 'gstat' for details.
<code>alpha</code>	direction in plane (x,y). see variogram in 'gstat' for details.
<code>block</code>	block size. see 'krige' in 'gstat' for details.
<code>beta</code>	for simple kriging. see 'krige' in 'gstat' for details.
<code>nmaxkrige</code>	for a local predicting: the number of nearest observations that should be used for a prediction or simulation, where nearest is defined in terms of the space of the spatial locations. By default, 12 observations are used.
<code>...</code>	other arguments passed on to 'svm' and 'krige'.

**Value**

A dataframe of longitude, latitude, and predictions.

**Author(s)**

Jin Li

**References**

- Li, J., Potter, A., Huang, Z., and Heap, A. (2012). Predicting Seabed Sand Content across the Australian Margin Using Machine Learning and Geostatistical Methods, *Geoscience Australia, Record 2012/48*, 115pp.
- Li, J., Heap, A., Potter, A., and Danilel, J.J. (2011). Predicting Seabed Mud Content across the Australian Margin II: Performance of Machine Learning Methods and Their Combination with Ordinary Kriging and Inverse Distance Squared, *Geoscience Australia, Record 2011/07*, 69pp.
- David Meyer, Evgenia Dimitriadou, Kurt Hornik, Andreas Weingessel and Friedrich Leisch (2020). e1071: Misc Functions of the Department of Statistics, Probability Theory Group (Formerly: E1071), TU Wien. R package version 1.7-4. <https://CRAN.R-project.org/package=e1071>.
- Pebesma, E.J., 2004. Multivariable geostatistics in S: the gstat package. *Computers & Geosciences*, 30: 683-691.

**Examples**

```
library(spm)

data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
longlat <- petrel[, c(1, 2)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)
y <- log(gravel[, 7] + 1)

svmkrigepred1 <- svmkrigepred(formula.svm = model, longlat = longlat, trainxy = gravel,
predx = petrel.grid, y = y, longlatpredx = petrel.grid[, c(1:2)],
transformation = "none", formula.krige = res1 ~ 1, vgm.args = "Sph", nmaxkrige = 12)

names(svmkrigepred1)

# Back transform 'svmkrigepred$predictions' to generate the final predictions
svmkrige.predictions <- exp(svmkrigepred1$predictions) - 1
range(svmkrige.predictions)
```

svmpred

*Generate spatial predictions using support vector machine ('svm')***Description**

This function is for generating spatial predictions using 'svm' method in 'e1071' package.

**Usage**

```
svmpred(
  formula = NULL,
  trainxy,
  longlatpredx,
  predx,
  scale = TRUE,
  type = NULL,
  kernel = "radial",
  degree = 3,
  gamma = if (is.vector(trainxy)) 1 else 1/ncol(trainxy),
  coef0 = 0,
  cost = 1,
  nu = 0.5,
  tolerance = 0.001,
  epsilon = 0.1,
  ...
)
```

**Arguments**

formula	a formula defining the response variable and predictive variables.
trainxy	a dataframe contains predictive variables and the response variable of point samples. The location information, longitude (long), latitude (lat), need to be included in the 'trainx' for spatial predictive modeling, need to be named as 'long' and 'lat'.
longlatpredx	a dataframe contains longitude and latitude of point locations (i.e., the centers of grids) to be predicted, need to be named as 'long' and 'lat'.
predx	a dataframe or matrix contains columns of predictive variables for the grids to be predicted.
scale	A logical vector indicating the variables to be scaled (default: TRUE).
type	the default setting is 'NULL'. See '?svm' for various options.
kernel	the default setting is 'radial'. See '?svm' for other options.
degree	a parameter needed for kernel of type polynomial (default: 3).
gamma	a parameter needed for all 'kernels' except 'linear' (default: 1/(data dimension)).
coef0	a parameter needed for kernels of type 'polynomial' and 'sigmoid'(default: 0).

cost	cost of constraints violation (default: 1).
nu	a parameter needed for 'nu-classification', 'nu-regression', and 'one-classification' (default: 0.5).
tolerance	tolerance of termination criterion (default: 0.001).
epsilon	'epsilon' in the insensitive-loss function (default: 0.1). See '?svm' for details.
...	other arguments passed on to 'svm'.

### Value

A dataframe of longitude, latitude and predictions.

### Author(s)

Jin Li

### References

David Meyer, Evgenia Dimitriadou, Kurt Hornik, Andreas Weingessel and Friedrich Leisch (2020). e1071: Misc Functions of the Department of Statistics, Probability Theory Group (Formerly: E1071), TU Wien. R package version 1.7-4. <https://CRAN.R-project.org/package=e1071>.

### Examples

```
library(spm)
data(petrel)
data(petrel.grid)

gravel <- petrel[, c(1, 2, 6:9, 5)]
model <- log(gravel + 1) ~ lat + bathy + I(long^3) + I(lat^2) + I(lat^3)

svmpred1 <- svmpred(formula = model, trainxy = gravel,
  longlatpredx = petrel.grid[, c(1:2)], predx = petrel.grid)

names(svmpred1)

# Back transform 'svmpred1$pred.svm1' to generate the final predictions
svm.predictions <- exp(svmpred1$pred.svm1) - 1
range(svm.predictions)
```

---

tpscv	<i>Cross validation, n-fold and leave-one-out for thin plate splines ('TPS')</i>
-------	--

---

### Description

This function is a cross validation function for 'Tps' method in 'fields' package.

### Usage

```
tpscv(
  trainx,
  trainy,
  m = NULL,
  p = NULL,
  theta = 3,
  lambda = NULL,
  lon.lat = TRUE,
  validation = "CV",
  cv.fold = 10,
  predacc = "VEcv",
  ...
)
```

### Arguments

trainx	a dataframe contains longitude (long), latitude (lat) and predictive variables of point samples. That is, they must be names as 'long' and 'lat'.
trainy	a vector of response, must have length equal to the number of rows in trainx.
m	A polynomial function of degree (m-1) will be included in the model as the drift (or spatial trend) component. Default is 'm = NULL' that is the value such that 2m-d is greater than zero where d is the dimension of x.
p	polynomial power for Wendland radial basis functions as in 'Tps'. 'p = NULL' that leads to a default value of 2 for spatial predictive modelling based on 'x' containing only the location information.
theta	the tapering range. 'theta = 3' degrees is a very generous taper range. For spatial predictive modeling the taper should be large enough to about 20 non zero nearest neighbors for every location.
lambda	smoothing parameter, the default is 'NULL'. See '?Tps' for further info.
lon.lat	if 'TRUE' locations are interpreted as longitude and latitude and great circle distance is used to find distances among locations.
validation	validation methods, include 'LOO': leave-one-out, and 'CV': cross-validation.
cv.fold	integer; number of folds in the cross-validation. if > 1, then apply n-fold cross validation; the default is 10, i.e., 10-fold cross validation that is recommended.
predacc	can be either "VEcv" for vecv or "ALL" for all measures in function pred.acc.
...	other arguments passed on to 'gstat'.

**Value**

A list with the following components: me, rme, mae, rmae, mse, rmse, rrmse, vecv and e1; or vecv only

**Note**

This function is largely based on 'krigecv' in this package and 'Tps' and 'fastTpsMLE' in 'fields' package.

**Author(s)**

Jin Li

**References**

Douglas Nychka and Reinhard Furrer and John Paige and Stephan Sain and Florian Gerber and Matthew Iverson, 2020. fields: Tools for Spatial Data, R package version 10.3 <https://CRAN.R-project.org/package=fields>.

**Examples**

```
library(fields)
library(spm)
data(petrel)

tpscv1 <- tpscvcv(petrel[, c(1,2)], petrel[, 5], cv.fold = 5, predacc = "VEcv")
tpscv1

tpscv1 <- tpscvcv(petrel[, c(1,2)], petrel[, 5], lambda = 0.9, cv.fold = 5, predacc = "VEcv")
tpscv1

tpscv1 <- tpscvcv(petrel[, c(1,2)], petrel[, 5], validation = "L00", predacc = "VEcv")
tpscv1

set.seed(1234)
n <- 20 # number of iterations,60 to 100 is recommended.
VEcv <- NULL
for (i in 1:n) {
  tpscvcv1 <- tpscvcv(petrel[, c(1,2)], petrel[, 5], cv.fold = 10, lambda = 0.13, predacc = "VEcv")
  VEcv [i] <- tpscvcv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for TPS", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)

n <- 20 # number of iterations,60 to 100 is recommended.
# set.seed(1234)
VEcv <- NULL
for (i in 1:n) {
  set.seed(1234 + i) # set random seed for each iteration. You can remove
```



```
# this line and use above set.seed(1234) and see what you can get.
tpscv1 <- tpscv(petrel[, c(1,2)], petrel[, 5], predacc = "VEcv")
VEcv [i] <- tpscv1
}
plot(VEcv ~ c(1:n), xlab = "Iteration for TPS", ylab = "VEcv (%)")
points(cumsum(VEcv) / c(1:n) ~ c(1:n), col = 2)
abline(h = mean(VEcv), col = 'blue', lwd = 2)
```

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