# Package: DMRnet (via r-universe)

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Type Package

**Title** Delete or Merge Regressors Algorithms for Linear and Logistic Model Selection and High-Dimensional Data

Version 0.4.0.9000

Description Model selection algorithms for regression and classification, where the predictors can be continuous or categorical and the number of regressors may exceed the number of observations. The selected model consists of a subset of numerical regressors and partitions of levels of factors. Szymon Nowakowski, Piotr Pokarowski, Wojciech Rejchel and Agnieszka Sołtys, 2023. Improving Group Lasso for High-Dimensional Categorical Data. In: Computational Science -ICCS 2023. Lecture Notes in Computer Science, vol 14074, p. 455-470. Springer, Cham. <doi:10.1007/978-3-031-36021-3\_47>. Aleksandra Maj-Kańska, Piotr Pokarowski and Agnieszka Prochenka, 2015. Delete or merge regressors for linear model selection. Electronic Journal of Statistics 9(2): 1749-1778. <doi:10.1214/15-EJS1050>. Piotr Pokarowski and Jan Mielniczuk, 2015. Combined 11 and greedy 10 penalized least squares for linear model selection. Journal of Machine Learning Research 16(29): 961-992. <https://www.jmlr.org/papers/volume16/pokarowski15a/pokarowski15a.pdf>. Piotr Pokarowski, Wojciech Rejchel, Agnieszka Sołtys, Michał Frej and Jan Mielniczuk, 2022. Improving Lasso for model selection and prediction. Scandinavian Journal of Statistics, 49(2): 831-863. <doi:10.1111/sjos.12546>.

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**Encoding** UTF-8

LazyData true

Imports hclust1d, glmnet, grpreg, stats, graphics, utils

Suggests knitr

URL https://github.com/SzymonNowakowski/DMRnet

BugReports https://github.com/SzymonNowakowski/DMRnet/issues

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# Contents

DMRnet-package	2
coef.cv.DMR	4
coef.DMR	5
coef.gic.DMR	5
cv.DMR	6
cv.DMRnet	8
DMR	10
DMRnet	12
gic.DMR	15
miete	16
plot.cv.DMR	17
plot.DMR	18
plot.gic.DMR	18
predict.cv.DMR	19
predict.DMR	20
predict.gic.DMR	22
print.DMR	23
promoter	23
	26

# Index

DMRnet-package

DMRnet-package

# Description

Model selection algorithms for regression and classification, where the predictors can be continuous or categorical and the number of regressors may exceed the number of observations. The selected model consists of a subset of numerical regressors and partitions of levels of factors.

#### DMRnet-package

#### **DMRnet Functions**

Similar in use to **glmnet**. It consists of the following functions:

DMR - Model selection algorithm for p<n; produces a path of models.

DMRnet - Model selection algorithm both for p<n and for p>=n; produces a path of models.

print.DMR, coef.DMR, plot.DMR, predict.DMR - Functions for inspection of the models on the path.

gic.DMR, cv.DMR, cv.DMRnet - Functions for final model selection, resulting with one model from the path.

coef.gic.DMR, coef.cv.DMR, plot.gic.DMR, plot.cv.DMR, predict.gic.DMR, predict.cv.DMR
- Functions for inspection of the final model.

miete, promoter - Two data sets used for vignettes, examples, etc.

For more information see a friendly "Getting started" vignette:

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#### References

Aleksandra Maj-Kańska, Piotr Pokarowski and Agnieszka Prochenka, 2015. Delete or merge regressors for linear model selection. Electronic Journal of Statistics 9(2): 1749-1778. doi:10.1214/ 15EJS1050

Piotr Pokarowski and Jan Mielniczuk, 2015. Combined 11 and greedy 10 penalized least squares for linear model selection. Journal of Machine Learning Research 16(29): 961-992. https://www.jmlr.org/papers/volume16/pokarowski15a/pokarowski15a.pdf

Piotr Pokarowski, Wojciech Rejchel, Agnieszka Sołtys, Michał Frej and Jan Mielniczuk, 2022. Improving Lasso for model selection and prediction. Scandinavian Journal of Statistics, 49(2): 831–863. doi:10.1111/sjos.12546

# Examples

```
## Not run:
vignette("getting-started", package="DMRnet")
```

## End(Not run)

coef.cv.DMR

# Description

Extracts coefficients from a cv.DMR object (for the model with minimal cross-validated error /the default/ or the smallest model falling under the upper curve of a prediction error plus one standard deviation).

#### Usage

## S3 method for class 'cv.DMR'
coef(object, md = "df.min", ...)

# Arguments

object	Fitted cv.DMR object.
md	Value of the model dimension parameter at which predictions are required. The default is md="df.min" value indicating the model minimizing the cross validation error. Alternatively, md="df.1se" can be used, indicating the smallest model falling under the upper curve of a prediction error plus one standard deviation.
	Further arguments passed to or from other methods.

# Details

Similar to other coef methods, this function extracts coefficients from a fitted cv.DMR object.

#### Value

Vector of coefficients.

```
## cv.DMR for linear regression
set.seed(13)
data(miete)
y <- miete$rent
X <- miete$area
cv = cv.DMR(X,y)
coef(cv)</pre>
```

coef.DMR

# Description

Extracts coefficients from a DMR object.

# Usage

## S3 method for class 'DMR'
coef(object, df = NULL, ...)

# Arguments

object	Fitted DMR object.
df	Number of parameters in the model for which coefficients are required. Default is the entire path of models.
	Further arguments passed to or from other methods.

# Details

Similar to other coef methods, this function extracts coefficients from a fitted DMR object.

#### Value

Vector or matrix of coefficients.

# Examples

```
data(miete)
y <- miete[,1]
X <- miete[,-1]
m <- DMR(X, y)
coef(m, df = 12)</pre>
```

coef.gic.DMR

# coef.gic.DMR

#### Description

Extracts coefficients from a gic.DMR object (for the model with minimal gic).

#### Usage

```
## S3 method for class 'gic.DMR'
coef(object, ...)
```

#### Arguments

object	Fitted gic.DMR object.
	Further arguments passed to or from other methods.

#### Details

Similar to other coef methods, this function extracts coefficients from a fitted gic.DMR object for the model with minimal gic.

# Value

Vector of coefficients.

# Examples

```
data(miete)
y <- miete[,1]
X <- miete[,-1]
m <- DMR(X, y)
g <- gic.DMR(m, c = 2.5)
coef(g)</pre>
```

cv.DMR

cross-validation for DMR

# Description

Executes k-fold cross-validation for DMR and returns a value for df.

#### Usage

```
cv.DMR(
   X,
   y,
   family = "gaussian",
   clust.method = "complete",
   lam = 10^(-7),
   nfolds = 10,
   indexation.mode = "GIC"
)
```

#### Arguments

```
Х
```

Input data frame, of dimension n x p; DMR works only if p<n, for p>=n see DMRnet; each row is an observation vector. Columns can be numerical or integer for continuous predictors or factors for categorical predictors.

у	Response variable. Numerical for family="gaussian" or a factor with two levels for family="binomial". For family="binomial" the last level in alphabetical order is the target class.
family	Response type; one of: "gaussian", "binomial".
clust.method	Clustering method used for partitioning levels of factors; see function hclust in package <b>stats</b> for details. clust.method="complete" is the default.
lam	The amount of penalization in ridge regression (used for logistic regression in or- der to allow for parameter estimation in linearly separable setups) or the amount of matrix regularization in case of linear regression. Used only for numerical reasons. The default is 1e-7.
nfolds indexation.mode	Number of folds in cross-validation. The default value is 10.
	How the cross validation algorithm should index the models for internal qual- ity comparisons; one of: "GIC" (the default) for GIC-indexed cross validation, "dimension", for model dimension-indexed cross validation.

#### Details

cv.DMR algorithm does cross-validation for DMR with nfolds folds. The df for the minimal estimated prediction error is returned.

#### Value

An object with S3 class "cv.DMR" is returned, which is a list with the ingredients of the cross-validation fit.

df.min df (number of parameters) of the model with minimal cross-validated error.

**df.1se** df (number of parameters) of the smallest model falling under the upper curve of a prediction error plus one standard deviation.

dmr.fit Fitted DMR object for the full data.

cvm The mean cross-validated error for the entire sequence of models.

foldid The fold assignments used.

#### See Also

plot.cv.DMR for plotting, coef.cv.DMR for extracting coefficients and predict.cv.DMR for prediction.

```
## cv.DMR for linear regression
set.seed(13)
data(miete)
ytr <- miete$rent[1:1500]
Xtr <- miete$area[1:1500]
Xte <- miete$area[1501:2053]
cv <- cv.DMR(Xtr, ytr)
print(cv)</pre>
```

```
plot(cv)
coef(cv)
ypr <- predict(cv, newx = Xte)</pre>
```

cv.DMRnet

#### cross-validation for DMRnet

# Description

Executes k-fold cross-validation for DMR and returns a value for df.

# Usage

```
cv.DMRnet(
    X,
    y,
    family = "gaussian",
    o = 5,
    nlambda = 100,
    lam = 10^(-7),
    interc = TRUE,
    maxp = ifelse(family == "gaussian", ceiling(length(y)/2), ceiling(length(y)/4)),
    nfolds = 10,
    indexation.mode = "GIC",
    algorithm = "DMRnet",
    clust.method = ifelse(algorithm == "glamer", "single", "complete")
)
```

# Arguments

Х	Input data frame, of dimension n x p; each row is an observation vector. Columns can be numerical or integer for continuous predictors or factors for categorical predictors.
У	Response variable. Numerical for family="gaussian" or a factor with two levels for family="binomial". For family="binomial" the last level in alphabetical order is the target class.
family	Response type; one of: "gaussian", "binomial".
0	Parameter of the group lasso screening step, described in DMRnet.
nlambda	Parameter of the group lasso screening step, described in DMRnet. The default value is 100.
lam	The amount of penalization in ridge regression (used for logistic regression in or- der to allow for parameter estimation in linearly separable setups) or the amount of matrix regularization in case of linear regression. Used only for numerical reasons. The default value is 1e-7.

8

#### cv.DMRnet

interc	Should intercept(s) be fitted (the default, interc=TRUE) or set to zero (interc=FALSE). If in X there are any categorical variables, interc=TRUE must be set.
maxp	Maximal number of parameters of the model, smaller values result in quicker computation.
nfolds indexation.mode	Number of folds in cross-validation. The default value is 10.
	How the cross validation algorithm should index the models for internal qual- ity comparisons; one of: "GIC" (the default) for GIC-indexed cross validation, "dimension", for model dimension-indexed cross validation.
algorithm	The algorithm to be used; for partition selection (merging levels) use one of: "DMRnet" (the default), "glamer" or "PDMR". Alternatively, use "var_sel" for variable (group) selection with no partition selection.
clust.method	Clustering method used for partitioning levels of factors; see function hclust in package <b>stats</b> for details. clust.method="complete" is the default for all algorithms except algorithm="glamer", for which clust.method="single" is the default.

# Details

cv.DMRnet algorithm does nfold-fold cross-validation for DMRnet. The df for the minimal estimated prediction error is returned.

#### Value

An object with S3 class "cv.DMR" is returned, which is a list with the ingredients of the cross-validation fit.

df.min df (number of parameters) of the model with minimal cross-validated error.

**df.1se** df (number of parameters) of the smallest model falling under the upper curve of a prediction error plus one standard deviation.

dmr.fit Fitted DMR object for the full data.

cvm The mean cross-validated error for the entire sequence of models.

foldid The fold assignments used.

#### See Also

plot.cv.DMR for plotting, coef.cv.DMR for extracting coefficients and predict.cv.DMR for prediction.

```
## cv.DMRnet for linear regression
set.seed(13)
data(miete)
ytr <- miete$rent[1:1500]
Xtr <- miete$area[1:1500]
Xte <- miete$area[1501:2053]
cv <- cv.DMRnet(Xtr, ytr)</pre>
```

```
print(cv)
plot(cv)
coef(cv)
ypr <- predict(cv, newx = Xte)</pre>
```

DMR

#### Delete or Merge Regressors

# Description

Fits a path of linear (family="gaussian") or logistic (family="binomial") regression models, where the number of parameters changes from 1 to p (p is the number of columns in the model matrix). Models are subsets of continuous predictors and partitions of levels of factors in X.

# Usage

```
DMR(
   X,
   y,
   family = "gaussian",
   clust.method = "complete",
   lam = 10^(-7),
   lambda = NULL
)
```

#### Arguments

x	Input data frame; each row is an observation vector; each column can be numer- ical or integer for a continuous predictor or a factor for a categorical predictor; DMR works only if p <n (n="" is="" number="" observations,="" of="" of<br="" p="" the="">columns in the model matrix), for p&gt;=n see DMRnet.</n>
у	Response variable; Numerical for family="gaussian" or a factor with two levels for family="binomial". For family="binomial" the last level in alphabetical order is the target class.
family	Response type; one of: "gaussian", "binomial".
clust.method	Clustering method used for partitioning levels of factors; see function hclust in package <b>stats</b> for details. clust.method="complete" is the default.
lam	The amount of penalization in ridge regression (used for logistic regression in or- der to allow for parameter estimation in linearly separable setups) or the amount of matrix regularization in case of linear regression. Used only for numerical reasons. The default is 1e-7.
lambda	The net of lambda values. It is optional and serves only for consistency with DMRnet. It is not used in DMR.

# DMR

#### Details

DMR algorithm is based on a traditional stepwise method. A nested family of models is built based on the values of squared Wald statistics:

1. For each continuous variable the squared Wald statistic is calculated for a hypothesis that the variable is equal to zero (it should be deleted).

2. For each factor a dissimilarity matrix is constructed using squared Wald statistics for hypotheses that two parameters are equal (the two levels of factor should be merged). Next, hierarchical clustering is preformed using the dissimilarity matrix. All cutting heights are recorded.

3. Squared Wald statistics and cutting heights and values of from steps 2 and 3 are concatenated and sorted, resulting in vector h.

4. Nested family of models of size 1 to p is built by accepting hypotheses according to increasing values in vector h.

# Value

An object with S3 class "DMR", which is a list with the ingredients:

beta	Matrix p times p of estimated parameters; each column corresponds to a model on the nested path having from p to 1 parameter (denoted as df).
df	Vector of degrees of freedom; from p to 1.
rss/loglik	Measure of fit for the nested models: rss (residual sum of squares) is returned for family="gaussian" and loglik (loglikelihood) is returned for family="binomial".
n	Number of observations.
levels.listed	Minimal set of levels of respective factors present in data.
lambda	The net of lambda values used in the screening step, empty vector in case of DMR.
arguments	List of the chosen arguments from the function call.
interc	If the intercept was fitted: for DMR always equal to TRUE.

#### See Also

print.DMR for printing, plot.DMR for plotting, coef.DMR for extracting coefficients and predict.DMR for prediction.

```
## DMR for linear regression
data(miete)
ytr <- miete[1:1500,1]
Xtr <- miete[1:1500,-1]
Xte <- miete[1501:2053,-1]
m1 <- DMR(Xtr, ytr)
print(m1)
plot(m1)
g <- gic.DMR(m1, c = 2.5)
plot(g)</pre>
```

```
coef(m1, df = g$df.min)
ypr <- predict(m1, newx = Xte, df = g$df.min)
## DMR for logistic regression
# notice that only part of dataset promoter was used: DMR works only if p<n, for p>=n use DMRnet
data(promoter)
ytr <- factor(promoter[1:80,1])
Xtr <- promoter[1:80,2:11]
Xte <- promoter[81:106,2:11]
m2 <- DMR(Xtr, ytr, family = "binomial")
print(m2)
plot(m2)
g <- gic.DMR(m2, c = 2)
plot(g)
coef(m2, df = g$df.min)
ypr <- predict(m2, newx = Xte, df = g$df.min)</pre>
```

DMRnet

Delete or Merge Regressors net

#### Description

Fits a path of linear (family="gaussian") or logistic (family="binomial") regression models, where models are subsets of continuous predictors and partitions of levels of factors in X. Works even if p>=n (the number of observations is greater than the number of columns in the model matrix).

#### Usage

```
DMRnet(
    X,
    y,
    family = "gaussian",
    o = 5,
    nlambda = 100,
    lam = 10^(-7),
    interc = TRUE,
    maxp = ifelse(family == "gaussian", ceiling(length(y)/2), ceiling(length(y)/4)),
    lambda = NULL,
    algorithm = "DMRnet",
    clust.method = ifelse(algorithm == "glamer", "single", "complete")
)
```

#### Arguments

Х

Input data frame; each row is an observation vector; each column can be numerical or integer for a continuous predictor or a factor for a categorical predictor.

12

У	Response variable; Numerical for family="gaussian" or a factor with two levels for family="binomial". For family="binomial" the last level in alphabetical order is the target class.
family	Response type; one of: "gaussian", "binomial".
0	Parameter of the group lasso screening step, described in Details, the default value is 5.
nlambda	Parameter of the group lasso screening step, described in Details, the default value is 100.
lam	The amount of penalization in ridge regression (used for logistic regression in or- der to allow for parameter estimation in linearly separable setups) or the amount of matrix regularization in case of linear regression. Used only for numerical reasons. The default is 1e-7.
interc	Should intercept(s) be fitted (the default, interc=TRUE) or set to zero (interc=FALSE). If in X there are any categorical variables, interc=TRUE must be set.
maxp	Maximal number of parameters of the model, smaller values result in quicker computation
lambda	Explicitly provided net of lambda values for the group lasso screening step, described in Details. If provided, it overrides the value of nlambda parameter.
algorithm	The algorithm to be used; for partition selection (merging levels) use one of: "DMRnet" (the default), "glamer" or "PDMR". Alternatively, use "var_sel" for variable (group) selection with no partition selection.
clust.method	Clustering method used for partitioning levels of factors; see function hclust in package stats for details. clust.method="complete" is the default for all algorithms except algorithm="glamer", for which clust.method="single" is the default.

#### Details

DMRnet algorithm is a generalization of DMR to high-dimensional data. It uses a screening step in order to decrease the problem to p<n and then uses DMR subsequently. The screening is done with the group lasso algorithm implemented in the grpreg package.

First, the group lasso for the problem is solved for nlambda values of lambda parameter, or for the net of lambda values (if lambda is explicitly provided). Next, for each value of lambda, the scaled nonzero second norms of the groups' coefficients are sorted in decreasing order. Finally, the first i over o fraction of the groups with the largest nonzero values are chosen for further analysis, i = 1,2,...,o-1. E.g., if o=5, first 1/5, first 2/5,..., 4/5 groups with the largest scaled nonzero second norm of coefficients are chosen.

The final path of models is chosen by minimizing the likelihood of the models for the number of parameters df equal to 1,...,l<=maxp for some integer l. Note that, in contrast to DMR, the models on the path need not to be nested.

#### Value

An object with S3 class "DMR", which is a list with the ingredients:

beta	Matrix p times l of estimated parameters; each column corresponds to a model on the nested path having from l to 1 parameter (denoted as df).
df	Vector of degrees of freedom; from l to 1.
rss/loglik	Measure of fit for the nested models: rss (residual sum of squares) is returned for family="gaussian" and loglik (loglikelihood) is returned for family="binomial".
n	Number of observations.
levels.listed	Minimal set of levels of respective factors present in data.
lambda	The net of lambda values used in the screening step.
arguments	List of the chosen arguments from the function call.
interc	If the intercept was fitted: value of parameter interc is returned.

# See Also

print.DMR for printing, plot.DMR for plotting, coef.DMR for extracting coefficients and predict.DMR for prediction.

```
## DMRnet for linear regression
data(miete)
ytr <- miete[1:200,1]</pre>
Xtr <- miete[1:200,-1]
Xte <- miete[201:250,-1]
m1 <- DMRnet(Xtr, ytr)</pre>
print(m1)
plot(m1)
g <- gic.DMR(m1, c = 2.5)
plot(g)
coef(m1, df = g$df.min)
ypr <- predict(m1, newx = Xte, df = g$df.min)</pre>
## DMRnet for logistic regression
data(promoter)
ytr <- factor(promoter[1:70,1])</pre>
Xtr <- promoter[1:70,-1]</pre>
Xte <- promoter[71:106,-1]</pre>
m2 <- DMRnet(Xtr, ytr, family = "binomial")</pre>
print(m2)
plot(m2)
g \leq gic.DMR(m2, c = 2)
plot(g)
coef(m2, df = g$df.min)
ypr <- predict(m2, newx = Xte, df = g$df.min)</pre>
## PDMR for linear regression
data(miete)
ytr <- miete[1:200,1]</pre>
Xtr <- miete[1:200,-1]
Xte <- miete[201:250,-1]
```

# gic.DMR

```
m1 <- DMRnet(Xtr, ytr, algorithm="PDMR")
print(m1)
plot(m1)
g <- gic.DMR(m1, c = 2.5)
plot(g)
coef(m1, df = g$df.min)
ypr <- predict(m1, newx = Xte, df = g$df.min)</pre>
```

gic.DMR

gic.DMR

#### Description

Computes values of Generalized Information Criterion for the entire sequence of models from a DMR object.

# Usage

```
gic.DMR(
    x,
    c = ifelse(x$arguments$family == "gaussian", constants()$RIC_gaussian_constant,
        constants()$RIC_binomial_constant)
)
```

# Arguments

х	Fitted DMR object.
с	Parameter controlling amount of penalization for complexity of the model in the generalized information criterion (GIC). For linear regression GIC for model M is defined as $GIC_M = RSS_M + df_M * c * logp * s^2$ ,
	where $RSS_M$ is the residual sum of squares and $df_M$ is the number of parameters in the model M; $s^2$ is an estimator of $sigma^2$ based on the model in the DMR object with the largest number of parameters. For logistic regression GIC for model M is defined as
	$GIC_M = -2 * loglik_M +  M  * c * logp,$
	where $loglik_M$ is the logarithm of the likelihood function and $df_M$ is the number of parameters in the model M. Recommended values are c=2.5 for linear

# Value

An object of class "gic.DMR" is returned, which is a list with the ingredients of the gic fit.

regression and c=2 for logistic regression.

df.min df (number of parameters) for the model with minimal GIC.

dmr.fit Fitted DMR object.

gic Vector of GIC values for the entire sequence of models.

#### See Also

plot.gic.DMR for plotting, coef.gic.DMR for extracting coefficients and predict.gic.DMR for prediction.

# Examples

data(miete)
y <- miete[,1]
X <- miete[,-1]
m <- DMR(X, y)
(g <- gic.DMR(m, c = 2.5))</pre>

miete

miete dataset

#### Description

The miete data contains the rent index for Munich in 2003.

#### Usage

data(miete)

#### Format

A data frame with 2053 observations on the following 12 variables.

rent Rent in euros.
bathextra Special furniture in bathroom, yes = 1, no = 0.
tiles Bathroom with tiles, yes = 0, no = 1.
area Municipality.
kitchen Upmarket kitchen, yes = 1, no = 0.
rooms Number of rooms.
best Best address, yes = 1, no = 0.
good Good address, yes = 1, no = 0.
warm Warm water, yes = 0, no = 1.
central Central heating, yes = 0, no = 1.
year Year of construction.
size Living space in square meter.

#### References

Fahrmeir, L., Künstler, R., Pigeot, I., Tutz, G. (2004) Statistik: der Weg zur Datenanalyse. 5. Auflage, Berlin: Springer-Verlag.

16

# plot.cv.DMR

#### Examples

data(miete)
summary(miete)

plot.cv.DMR plot.cv.DMR

#### Description

Plots cross-validated error values from a cv. DMR object.

#### Usage

## S3 method for class 'cv.DMR'
plot(x, ...)

# Arguments

х	Fitted cv.DMR object.
	Further arguments passed to or from other methods.

#### Details

Produces a plot of cross-validated error values for the entire sequence of models from the fitted cv.DMR object. The horizontal level indicating separation of one standard deviation from the minimum error is indicated with a blue dashed line. The df.min (the smallest model minimizing the cross-validated error) and df.1se (the smallest model falling under the blue dashed line) are marked with red and blue points, respectively.

```
## cv.DMR for linear regression
set.seed(13)
data(miete)
y <- miete$rent
X <- miete$area
cv = cv.DMR(X,y)
plot(cv)</pre>
```

plot.DMR

# Description

Plots coefficients from a DMR object.

# Usage

## S3 method for class 'DMR'
plot(x, ...)

# Arguments

Х	Fitted DMR object.
	Further arguments passed to or from other methods.

# Details

Produces a coefficient profile plot of the coefficient paths for a fitted DMR object.

# Examples

```
data(miete)
y <- miete[,1]
X <- miete[,-1]
m <- DMR(X, y)
plot(m)</pre>
```

plot.gic.DMR

# Description

Plots gic values from a gic.DMR object.

#### Usage

## S3 method for class 'gic.DMR'
plot(x, ...)

# Arguments

х	Fitted gic.DMR object.
	Further arguments passed to or from other methods

plot.gic.DMR

# predict.cv.DMR

# Details

Produces a plot of Generalized Information Criterion for the entire sequence of models from the fitted gic.DMR object.

# Examples

```
data(miete)
y <- miete[,1]
X <- miete[,-1]
m <- DMR(X, y)
g <- gic.DMR(m, c = 2.5)
plot(g)</pre>
```

predict.cv.DMR predict.cv.DMR

# Description

Makes predictions from a cv.DMR object (for the model with minimal cross-validated error /the default/ or the smallest model falling under the upper curve of a prediction error plus one standard deviation).

# Usage

```
## S3 method for class 'cv.DMR'
predict(
   object,
   newx,
   type = "link",
   md = "df.min",
   unknown.factor.levels = "error",
   ...
)
```

#### Arguments

object	Fitted cv.DMR object.
newx	Data frame of new values for X at which predictions are to be made. The inter- cept column should NOT be passed in a call to predict.
type	One of: "link", "response", "class". For family="gaussian" for all values of type it gives the fitted values. For family="binomial" and type="link" it returns the linear predictors, for type="response" it returns the fitted proba- bilities and for type="class" it produces the class labels corresponding to the maximum probability.

md	Value of the model dimension parameter at which predictions are required. The default is md="df.min" value indicating the model minimizing the cross validation error. Alternatively, md="df.1se" can be used, indicating the smallest model falling under the upper curve of a prediction error plus one standard deviation.
unknown.factor.	levels
	The way of handling factor levels in test data not seen while training a model. One of "error" (the default - throwing an error) or "NA" (returning NA in place of legitimate value for problematic rows).
	Further arguments passed to or from other methods.

# Details

Similar to other predict methods, this function predicts fitted values from a fitted cv.DMR object.

#### Value

Vector of predictions.

# Examples

```
## cv.DMR for linear regression
set.seed(13)
data(miete)
ytr <- miete$rent[1:1500]
Xtr <- miete$area[1:1500]
Xte <- miete$area[1501:2053]
cv <- cv.DMR(Xtr, ytr)
print(cv)
plot(cv)
coef(cv)
ypr <- predict(cv, newx = Xte)</pre>
```

predict.DMR predict.DMR

# Description

Makes predictions from a DMR object.

# Usage

```
## S3 method for class 'DMR'
predict(
    object,
    newx,
    df = NULL,
```

# predict.DMR

```
type = "link",
unknown.factor.levels = "error",
...
```

# Arguments

object	Fitted DMR object.
newx	Data frame of new values for X at which predictions are to be made. The inter- cept column should NOT be passed in a call to predict.
df	Number of parameters in the model for which predictions are required. Default is the entire sequence of models for df=1 to df=p.
type	One of: "link", "response", "class". For family="gaussian" for all values of type it gives the fitted values. For family="binomial" and type="link" it returns the linear predictors, for type="response" it returns the fitted proba- bilities and for type="class" it produces the class labels corresponding to the maximum probability.
unknown.factor	.levels
	The way of handling factor levels in test data not seen while training a model. One of "error" (the default - throwing an error) or "NA" (returning NA in place of legitimate value for problematic rows).
•••	Further arguments passed to or from other methods.

# Details

Similar to other predict methods, this function predicts fitted values from a fitted DMR object.

#### Value

Vector or matrix of predictions.

```
data(miete)
ytr <- miete[1:1500,1]
Xtr <- miete[1:1500,-1]
Xte <- miete[1501:2053,-1]
m <- DMR(Xtr, ytr)
ypr <- predict(m, newx = Xte, df = 11)</pre>
```

predict.gic.DMR predict.gic.DMR

# Description

Makes predictions from a gic.DMR object (for the model with minimal GIC).

# Usage

```
## S3 method for class 'gic.DMR'
predict(object, newx, type = "link", unknown.factor.levels = "error", ...)
```

#### Arguments

object	Fitted gic.DMR object.
newx	Data frame of new values for X at which predictions are to be made. The intercept column should NOT be passed in a call to predict.
type	One of: "link", "response", "class". For family="gaussian" for all values of type it gives the fitted values. For family="binomial" and type="link" it returns the linear predictors, for type="response" it returns the fitted proba- bilities and for type="class" it produces the class labels corresponding to the maximum probability.
unknown.factor.levels	
	The way of handling factor levels in test data not seen while training a model. One of "error" (the default - throwing an error) or "NA" (returning NA in place of legitimate value for problematic rows).
	Further arguments passed to or from other methods.

# Details

Similar to other predict methods, this function predicts fitted values from a fitted gic.DMR object for the model with minimal GIC.

# Value

Vector of predictions.

```
data(miete)
ytr <- miete[1:1500,1]
Xtr <- miete[1:1500,-1]
Xte <- miete[1501:2053,-1]
m <- DMR(Xtr, ytr)
g <- gic.DMR(m, c = 2.5)
ypr <- predict(g, newx = Xte)</pre>
```

print.DMR

print.DMR

#### Description

Prints a DMR object.

# Usage

## S3 method for class 'DMR'
print(x, ...)

#### Arguments

Х	Fitted DMR object.
•••	Further arguments passed to or from other methods.

# Details

Print a summary of the DMR path at each step along the path.

# Value

The summary is silently returned.

# Examples

```
data(miete)
y <- miete[,1]
X <- miete[,-1]
m <- DMR(X, y)
print(m)</pre>
```

promoter

promoter dataset

# Description

It consists of E. coli promoter gene sequences starting at position -50 (p-50) and ending at position +7 (p7). Each of these 57 Fields is filled by one of a, g, t, c. The task is to recognize promoters, which are genetic regions which initiate the first step in the expression of adjacent genes (transcription). There are 53 promoters and 53 non-promoter sequences.

#### Usage

data(promoter)

#### Format

A data frame with 106 observations on the following 58 variables.

**y** One of 1/0, indicating the class (1 = promoter).

**X1** Sequence; filled by one of a, g, t, c.

**X2** Sequence; filled by one of a, g, t, c.

**X3** Sequence; filled by one of a, g, t, c.

**X4** Sequence; filled by one of a, g, t, c.

**X5** Sequence; filled by one of a, g, t, c.

**X6** Sequence; filled by one of a, g, t, c.

X7 Sequence; filled by one of a, g, t, c.

X8 Sequence; filled by one of a, g, t, c.

**X9** Sequence; filled by one of a, g, t, c.

**X10** Sequence; filled by one of a, g, t, c.

X11 Sequence; filled by one of a, g, t, c.

**X12** Sequence; filled by one of a, g, t, c.

X13 Sequence; filled by one of a, g, t, c.

X14 Sequence; filled by one of a, g, t, c.

**X15** Sequence; filled by one of a, g, t, c.

X16 Sequence; filled by one of a, g, t, c.

X17 Sequence; filled by one of a, g, t, c.

X18 Sequence; filled by one of a, g, t, c.

**X19** Sequence; filled by one of a, g, t, c.

 $\textbf{X20} \hspace{0.1 cm} \text{Sequence; filled by one of } a, \, g, \, t, \, c.$ 

**X21** Sequence; filled by one of a, g, t, c.

**X22** Sequence; filled by one of a, g, t, c.

**X23** Sequence; filled by one of a, g, t, c.

**X24** Sequence; filled by one of a, g, t, c.

**X25** Sequence; filled by one of a, g, t, c.

**X26** Sequence; filled by one of a, g, t, c.

**X27** Sequence; filled by one of a, g, t, c.

 $\textbf{X28} \hspace{0.1 cm} \text{Sequence; filled by one of } a, \, g, \, t, \, c.$ 

X29 Sequence; filled by one of a, g, t, c.

**X30** Sequence; filled by one of a, g, t, c.

X31 Sequence; filled by one of a, g, t, c.

 $\textbf{X32} \hspace{0.1 cm} \text{Sequence; filled by one of a, g, t, c.}$ 

**X33** Sequence; filled by one of a, g, t, c.

X34 Sequence; filled by one of a, g, t, c.

#### promoter

- X35 Sequence; filled by one of a, g, t, c.
- **X36** Sequence; filled by one of a, g, t, c.
- X37 Sequence; filled by one of a, g, t, c.
- **X38** Sequence; filled by one of a, g, t, c.
- **X39** Sequence; filled by one of a, g, t, c.
- X40 Sequence; filled by one of a, g, t, c.
- X41 Sequence; filled by one of a, g, t, c.
- $\textbf{X42} \hspace{0.1 cm} \text{Sequence; filled by one of a, g, t, c.}$
- X43 Sequence; filled by one of a, g, t, c.
- X44 Sequence; filled by one of a, g, t, c.
- X45 Sequence; filled by one of a, g, t, c.
- X46 Sequence; filled by one of a, g, t, c.
- X47 Sequence; filled by one of a, g, t, c.
- X48 Sequence; filled by one of a, g, t, c.
- X49 Sequence; filled by one of a, g, t, c.
- **X50** Sequence; filled by one of a, g, t, c.
- **X51** Sequence; filled by one of a, g, t, c.
- **X52** Sequence; filled by one of a, g, t, c.
- **X53** Sequence; filled by one of a, g, t, c.
- **X54** Sequence; filled by one of a, g, t, c.
- **X55** Sequence; filled by one of a, g, t, c.
- **X56** Sequence; filled by one of a, g, t, c.
- **X57** Sequence; filled by one of a, g, t, c.

#### Source

UCI machine learning repository: promoter

#### References

Towell, G., Shavlik, J., Noordewier, M. Refinement of approximate domain theories by knowledgebased neural networks. In Proceedings of the eighth National conference on Artificial intelligence, pages 861-866. Boston, MA, 1990.

```
data(promoter)
summary(promoter)
```

# Index

\* datasets miete, 16 promoter, 23 coef.cv.DMR, *3*, *4*, *7*, *9* coef.DMR, *3*, *5*, *11*, *14* coef.gic.DMR, *3*, *5*, *16* cv.DMR, 3, 6 cv.DMRnet, 3, 8 DMR, *3*, 10, *13* DMRnet, *3*, *6*, *8*, *10*, 12 DMRnet-package, 2 gic.DMR, *3*, 15 miete, 16 plot.cv.DMR, *3*, *7*, *9*, 17 plot.DMR, *3*, *11*, *14*, 18 plot.gic.DMR, *3*, *16*, 18 predict.cv.DMR, *3*, *7*, *9*, 19

predict.gic.DMR, *3*, *16*, 22 print.DMR, *3*, *11*, *14*, 23 promoter, 23

predict.DMR, *3*, *11*, *14*, 20