## Package 'oem'

October 13, 2022

Type Package

Title Orthogonalizing EM: Penalized Regression for Big Tall Data

Version 2.0.11

Maintainer Jared Huling <jaredhuling@gmail.com>

Description Solves penalized least squares problems for big tall data using the orthogonalizing EM algorithm of Xiong et al. (2016) <doi:10.1080/00401706.2015.1054436>. The main fitting function is oem() and the functions cv.oem() and xval.oem() are for cross validation, the latter being an accelerated cross validation function for linear models. The big.oem() function allows for out of memory fitting. A description of the underlying methods and code interface is described in Huling and Chien (2022) <doi:10.18637/jss.v104.i06>.

URL https://arxiv.org/abs/1801.09661,

https://github.com/jaredhuling/oem, https://jaredhuling.org/oem/

BugReports https://github.com/jaredhuling/oem/issues

**License** GPL ( $\geq 2$ )

Encoding UTF-8

Depends R (>= 3.2.0), bigmemory

**Imports** Rcpp (>= 0.11.0), Matrix, foreach, methods

LinkingTo Rcpp, RcppEigen, BH, bigmemory, RcppArmadillo

RoxygenNote 7.2.0

Suggests knitr, rmarkdown

VignetteBuilder knitr

NeedsCompilation yes

Author Bin Dai [aut], Jared Huling [aut, cre] (<https://orcid.org/0000-0003-0670-4845>), Yixuan Qiu [ctb], Gael Guennebaud [cph], Jitse Niesen [cph]

**Repository** CRAN

Date/Publication 2022-10-13 08:20:08 UTC

### **R** topics documented:

big.oem	2
cv.oem	6
logLik.oem	8
oem	9
oem.xtx	3
oemfit	7
plot.oem	9
predict.cv.oem	
predict.oem	3
predict.xval.oem	5
print.summary.cv.oem	
summary.cv.oem	
xval.oem	7
3	1

### Index

big.oem

Orthogonalizing EM for big.matrix objects

### Description

Orthogonalizing EM for big.matrix objects

### Usage

```
big.oem(
 х,
 у,
 family = c("gaussian", "binomial"),
 penalty = c("elastic.net", "lasso", "ols", "mcp", "scad", "mcp.net", "scad.net",
  "grp.lasso", "grp.lasso.net", "grp.mcp", "grp.scad", "grp.mcp.net", "grp.scad.net",
    "sparse.grp.lasso"),
 weights = numeric(0),
 lambda = numeric(0),
 nlambda = 100L,
  lambda.min.ratio = NULL,
  alpha = 1,
  gamma = 3,
  tau = 0.5,
  groups = numeric(0),
 penalty.factor = NULL,
  group.weights = NULL,
  standardize = TRUE,
  intercept = TRUE,
 maxit = 500L,
  tol = 1e-07,
```

### big.oem

```
irls.maxit = 100L,
irls.tol = 0.001,
compute.loss = FALSE,
gigs = 4,
hessian.type = c("full", "upper.bound")
)
```

### Arguments

x	input big.matrix object pointing to design matrix Each row is an observation, each column corresponds to a covariate
У	numeric response vector of length nobs.
family	"gaussian" for least squares problems, "binomial" for binary response. "binomial" currently not available.
penalty	Specification of penalty type. Choices include:
	<ul> <li>"elastic.net" - elastic net penalty, extra parameters: "alpha"</li> <li>"lasso" - lasso penalty</li> <li>"ols" - ordinary least squares</li> <li>"mcp" - minimax concave penalty, extra parameters: "gamma"</li> <li>"scad" - smoothly clipped absolute deviation, extra parameters: "gamma"</li> <li>"mcp.net" - minimax concave penalty + 12 penalty, extra parameters: "gamma", "alpha"</li> <li>"scad.net" - smoothly clipped absolute deviation + 12 penalty, extra pa-</li> </ul>
	<ul> <li>rameters: "gamma", "alpha"</li> <li>"grp.lasso" - group lasso penalty</li> <li>"grp.lasso.net" - group lasso penalty + 12 penalty, extra parameters: "alpha"</li> <li>"grp.mcp" - group minimax concave penalty, extra parameters: "gamma"</li> <li>"grp.scad" - group smoothly clipped absolute deviation, extra parameters: "gamma"</li> <li>"grp.mcp.net" - group minimax concave penalty + 12 penalty, extra parameters: "gamma"</li> <li>"grp.scad.net" - group smoothly clipped absolute deviation + 12 penalty, extra parameters: "gamma", "alpha"</li> </ul>
	<ul> <li>"sparse.grp.lasso" - sparse group lasso penalty (group lasso + lasso), extra parameters: "tau"</li> <li>Careful consideration is required for the group lasso, group MCP, and group SCAD penalties. Groups as specified by the groups argument should be chosen</li> </ul>
	in a sensible manner.
weights	observation weights. Not implemented yet. Defaults to 1 for each observation (setting weight vector to length 0 will default all weights to 1)
lambda	A user supplied lambda sequence. By default, the program computes its own lambda sequence based on nlambda and lambda.min.ratio. Supplying a value of lambda overrides this.

nlambda lambda.min.rati	The number of lambda values - default is 100.
	Smallest value for lambda, as a fraction of lambda.max, the (data derived) entry value (i.e. the smallest value for which all coefficients are zero). The default depends on the sample size nobs relative to the number of variables nvars. If nobs > nvars, the default is 0.0001, close to zero. If nobs < nvars, the default is 0.0001, close to zero. If nobs < nvars, the default is 0.001. A very small value of lambda.min.ratio will lead to a saturated fit when nobs < nvars.
alpha	<pre>mixing value for elastic.net, mcp.net, scad.net, grp.mcp.net, grp.scad.net. penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp/grp.lasso penalty)</pre>
gamma	tuning parameter for SCAD and MCP penalties. must be $\geq 1$
tau	mixing value for sparse.grp.lasso. penalty applied is (1 - tau) * (group lasso penalty) + tau * (lasso penalty)
groups	A vector of describing the grouping of the coefficients. See the example below. All unpenalized variables should be put in group 0
penalty.factor	Separate penalty factors can be applied to each coefficient. This is a number that multiplies lambda to allow differential shrinkage. Can be 0 for some variables, which implies no shrinkage, and that variable is always included in the model. Default is 1 for all variables.
group.weights	penalty factors applied to each group for the group lasso. Similar to penalty.factor, this is a number that multiplies lambda to allow differential shrinkage. Can be 0 for some groups, which implies no shrinkage, and that group is always included in the model. Default is sqrt(group size) for all groups.
standardize	Logical flag for x variable standardization, prior to fitting the models. The co- efficients are always returned on the original scale. Default is standardize = TRUE. If variables are in the same units already, you might not wish to standard- ize. Keep in mind that standardization is done differently for sparse matrices, so results (when standardized) may be slightly different for a sparse matrix object and a dense matrix object
intercept	Should intercept(s) be fitted (default = TRUE) or set to zero (FALSE)
maxit	integer. Maximum number of OEM iterations
tol	convergence tolerance for OEM iterations
irls.maxit	integer. Maximum number of IRLS iterations
irls.tol	convergence tolerance for IRLS iterations. Only used if family != "gaussian"
compute.loss	should the loss be computed for each estimated tuning parameter? Defaults to FALSE. Setting to TRUE will dramatically increase computational time
gigs	maximum number of gigs of memory available. Used to figure out how to break up calculations involving the design matrix $x$
hessian.type	only for logistic regression. if hessian.type = "full", then the full hessian is used. If hessian.type = "upper.bound", then an upper bound of the hessian is used. The upper bound can be dramatically faster in certain situations, ie when $n \gg p$

### big.oem

### Value

An object with S3 class "oem"

### References

Huling. J.D. and Chien, P. (2022), Fast Penalized Regression and Cross Validation for Tall Data with the oem Package. Journal of Statistical Software 104(6), 1-24. doi:10.18637/jss.v104.i06

```
## Not run:
set.seed(123)
nrows <- 50000
ncols <- 100
bkFile <- "bigmat.bk"</pre>
descFile <- "bigmatk.desc"</pre>
bigmat <- filebacked.big.matrix(nrow=nrows, ncol=ncols, type="double",</pre>
                                 backingfile=bkFile, backingpath=".",
                                  descriptorfile=descFile,
                                  dimnames=c(NULL,NULL))
# Each column value with be the column number multiplied by
# samples from a standard normal distribution.
set.seed(123)
for (i in 1:ncols) bigmat[,i] = rnorm(nrows)*i
y <- rnorm(nrows) + bigmat[,1] - bigmat[,2]</pre>
fit <- big.oem(x = bigmat, y = y,</pre>
               penalty = c("lasso", "elastic.net",
                            "ols",
                            "mcp",
                                          "scad",
                            "mcp.net",
                                          "scad.net",
                            "grp.lasso", "grp.lasso.net",
                            "grp.mcp", "grp.scad",
                            "sparse.grp.lasso"),
               groups = rep(1:20, each = 5))
fit2 <- oem(x = bigmat[,], y = y,</pre>
            penalty = c("lasso", "grp.lasso"),
            groups = rep(1:20, each = 5))
max(abs(fit$beta[[1]] - fit2$beta[[1]]))
layout(matrix(1:2, ncol = 2))
plot(fit)
plot(fit, which.model = 2)
## End(Not run)
```

cv.oem

### Description

Cross validation for Orthogonalizing EM

### Usage

```
cv.oem(
 х,
 у,
 penalty = c("elastic.net", "lasso", "ols", "mcp", "scad", "mcp.net", "scad.net",
  "grp.lasso", "grp.lasso.net", "grp.mcp", "grp.scad", "grp.mcp.net", "grp.scad.net",
    "sparse.grp.lasso"),
 weights = numeric(0),
  lambda = NULL,
  type.measure = c("mse", "deviance", "class", "auc", "mae"),
  nfolds = 10,
  foldid = NULL,
  grouped = TRUE,
 keep = FALSE,
 parallel = FALSE,
 ncores = -1,
  . . .
)
```

### Arguments

x	input matrix of dimension n x p or CsparseMatrix objects of the <b>Matrix</b> (sparse not yet implemented. Each row is an observation, each column corresponds to a covariate. The cv.oem() function is optimized for n » p settings and may be very slow when $p > n$ , so please use other packages such as glmnet, ncvreg, grpreg, or gglasso when $p > n$ or p approx n.
У	numeric response vector of length nobs.
penalty	Specification of penalty type in lowercase letters. Choices include "lasso", "ols" (Ordinary least squares, no penaly), "elastic.net", "scad", "mcp", "grp.lasso"
weights	observation weights. defaults to 1 for each observation (setting weight vector to length 0 will default all weights to 1)
lambda	A user supplied lambda sequence. By default, the program computes its own lambda sequence based on nlambda and lambda.min.ratio. Supplying a value of lambda overrides this.
type.measure	measure to evaluate for cross-validation. The default is type.measure = "deviance", which uses squared-error for gaussian models (a.k.a type.measure = "mse"

	there), deviance for logistic regression. type.measure = "class" applies to binomial only. type.measure = "auc" is for two-class logistic regression only. type.measure = "mse" or type.measure = "mae" (mean absolute error) can be used by all models; they measure the deviation from the fitted mean to the re- sponse.
nfolds	number of folds for cross-validation. default is 10. 3 is smallest value allowed.
foldid	an optional vector of values between 1 and nfold specifying which fold each observation belongs to.
grouped	Like in <b>glmnet</b> , this is an experimental argument, with default TRUE, and can be ignored by most users. For all models, this refers to computing nfolds separate statistics, and then using their mean and estimated standard error to describe the CV curve. If grouped = FALSE, an error matrix is built up at the observation level from the predictions from the nfold fits, and then summarized (does not apply to type.measure = "auc").
keep	If keep = TRUE, a prevalidated list of arrasy is returned containing fitted values for each observation and each value of lambda for each model. This means these fits are computed with this observation and the rest of its fold omitted. The folid vector is also returned. Default is keep = FALSE
parallel	If TRUE, use parallel foreach to fit each fold. Must register parallel before hand, such as <b>doMC</b> .
ncores	Number of cores to use. If parallel = TRUE, then neores will be automatically set to 1 to prevent conflicts
	other parameters to be passed to "oem" function

### Value

An object with S3 class "cv.oem"

### References

Huling. J.D. and Chien, P. (2022), Fast Penalized Regression and Cross Validation for Tall Data with the oem Package. Journal of Statistical Software 104(6), 1-24. doi:10.18637/jss.v104.i06

```
layout(matrix(1:2, ncol = 2))
plot(fit)
plot(fit, which.model = 2)
```

logLik.oem

log likelihood function for fitted oem objects

### Description

log likelihood function for fitted oem objects

log likelihood function for fitted cross validation oem objects

log likelihood function for fitted cross validation oem objects

### Usage

```
## S3 method for class 'oem'
logLik(object, which.model = 1, ...)
## S3 method for class 'cv.oem'
logLik(object, which.model = 1, ...)
## S3 method for class 'xval.oem'
logLik(object, which.model = 1, ...)
```

#### Arguments

object	fitted "oem" model object.
which.model	If multiple penalties are fit and returned in the same oem object, the which.model argument is used to specify which model to plot. For example, if the oem object "oemobj" was fit with argument penalty = $c("lasso", "grp.lasso")$ , then which.model = 2 provides a plot for the group lasso model.
	not used

### Examples

```
set.seed(123)
n.obs <- 2000
n.vars <- 50
true.beta <- c(runif(15, -0.25, 0.25), rep(0, n.vars - 15))
x <- matrix(rnorm(n.obs * n.vars), n.obs, n.vars)
y <- rnorm(n.obs, sd = 3) + x %*% true.beta
fit <- oem(x = x, y = y, penalty = c("lasso", "mcp"), compute.loss = TRUE)
logLik(fit)</pre>
```

oem

### Orthogonalizing EM

### Description

Orthogonalizing EM

### Usage

```
oem(
  х,
  у,
  family = c("gaussian", "binomial"),
 penalty = c("elastic.net", "lasso", "ols", "mcp", "scad", "mcp.net", "scad.net",
   "grp.lasso", "grp.lasso.net", "grp.mcp", "grp.scad", "grp.mcp.net", "grp.scad.net",
    "sparse.grp.lasso"),
  weights = numeric(0),
  lambda = numeric(0),
  nlambda = 100L,
  lambda.min.ratio = NULL,
  alpha = 1,
  gamma = 3,
  tau = 0.5,
  groups = numeric(\emptyset),
  penalty.factor = NULL,
  group.weights = NULL,
  standardize = TRUE,
  intercept = TRUE,
  maxit = 500L,
```

```
tol = 1e-07,
irls.maxit = 100L,
irls.tol = 0.001,
accelerate = FALSE,
ncores = -1,
compute.loss = FALSE,
hessian.type = c("upper.bound", "full")
)
```

### Arguments

X	input matrix of dimension n x p or CsparseMatrix object of the <b>Matrix</b> package. Each row is an observation, each column corresponds to a covariate. The oem() function is optimized for n » p settings and may be very slow when p > n, so please use other packages such as glmnet, ncvreg, grpreg, or gglasso when $p > n$ or p approx n.
У	numeric response vector of length nobs.
family	"gaussian" for least squares problems, "binomial" for binary response.
penalty	Specification of penalty type. Choices include:
	<ul> <li>"elastic.net" - elastic net penalty, extra parameters: "alpha"</li> <li>"lasso" - lasso penalty</li> <li>"ols" - ordinary least squares</li> </ul>
	<ul> <li>"mcp" - minimax concave penalty, extra parameters: "gamma"</li> </ul>
	<ul> <li>"scad" - smoothly clipped absolute deviation, extra parameters: "gamma"</li> </ul>
	<ul> <li>"mcp.net" - minimax concave penalty + 12 penalty, extra parameters: "gamma", "alpha"</li> </ul>
	<ul> <li>"scad.net" - smoothly clipped absolute deviation + 12 penalty, extra parameters: "gamma", "alpha"</li> </ul>
	<ul> <li>"grp.lasso" - group lasso penalty</li> </ul>
	<ul> <li>"grp.lasso.net" - group lasso penalty + 12 penalty, extra parameters: "alpha"</li> </ul>
	• "grp.mcp" - group minimax concave penalty, extra parameters: "gamma"
	<ul> <li>"grp.scad" - group smoothly clipped absolute deviation, extra parameters: "gamma"</li> </ul>
	<ul> <li>"grp.mcp.net" - group minimax concave penalty + 12 penalty, extra parameters: "gamma", "alpha"</li> </ul>
	<ul> <li>"grp.scad.net" - group smoothly clipped absolute deviation + 12 penalty, extra parameters: "gamma", "alpha"</li> </ul>
	<ul> <li>"sparse.grp.lasso" - sparse group lasso penalty (group lasso + lasso), extra parameters: "tau"</li> </ul>
	Careful consideration is required for the group lasso, group MCP, and group SCAD penalties. Groups as specified by the groups argument should be chosen in a sensible manner.
weights	observation weights. Not implemented yet. Defaults to 1 for each observation (setting weight vector to length 0 will default all weights to 1)

lambda	A user supplied lambda sequence. By default, the program computes its own lambda sequence based on nlambda and lambda.min.ratio. Supplying a value of lambda overrides this.
nlambda lambda.min.rati	The number of lambda values. The default is 100.
	Smallest value for lambda, as a fraction of lambda.max, the (data derived) entry value (i.e. the smallest value for which all coefficients are zero). The default depends on the sample size nobs relative to the number of variables nvars. If nobs > nvars, the default is 0.0001, close to zero. If nobs < nvars, the default is 0.001. A very small value of lambda.min.ratio will lead to a saturated fit when nobs < nvars.
alpha	<pre>mixing value for elastic.net, mcp.net, scad.net, grp.mcp.net, grp.scad.net. penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp/grp.lasso penalty)</pre>
gamma	tuning parameter for SCAD and MCP penalties. must be $>= 1$
tau	mixing value for sparse.grp.lasso. penalty applied is (1 - tau) * (group lasso penalty) + tau * (lasso penalty)
groups	A vector of describing the grouping of the coefficients. See the example below. All unpenalized variables should be put in group 0
penalty.factor	Separate penalty factors can be applied to each coefficient. This is a number that multiplies lambda to allow differential shrinkage. Can be 0 for some variables, which implies no shrinkage, and that variable is always included in the model. Default is 1 for all variables.
group.weights	penalty factors applied to each group for the group lasso. Similar to penalty.factor, this is a number that multiplies lambda to allow differential shrinkage. Can be 0 for some groups, which implies no shrinkage, and that group is always included in the model. Default is sqrt(group size) for all groups.
standardize	Logical flag for x variable standardization, prior to fitting the models. The co- efficients are always returned on the original scale. Default is standardize = TRUE. If variables are in the same units already, you might not wish to standard- ize. Keep in mind that standardization is done differently for sparse matrices, so results (when standardized) may be slightly different for a sparse matrix object and a dense matrix object
intercept	Should intercept(s) be fitted (default = TRUE) or set to zero (FALSE)
maxit	integer. Maximum number of OEM iterations
tol	convergence tolerance for OEM iterations
irls.maxit	integer. Maximum number of IRLS iterations
irls.tol	convergence tolerance for IRLS iterations. Only used if family != "gaussian"
accelerate	boolean argument. Whether or not to use Nesterov acceleration with adaptive restarting
ncores	Integer scalar that specifies the number of threads to be used
compute.loss	should the loss be computed for each estimated tuning parameter? Defaults to FALSE. Setting to TRUE will dramatically increase computational time

```
hessian.type only for logistic regression. if hessian.type = "full", then the full hessian is
used. If hessian.type = "upper.bound", then an upper bound of the hessian is
used. The upper bound can be dramatically faster in certain situations, ie when
n » p
```

### Value

An object with S3 class "oem"

### References

Shifeng Xiong, Bin Dai, Jared Huling, and Peter Z. G. Qian. Orthogonalizing EM: A design-based least squares algorithm. Technometrics, 58(3):285-293, 2016. https://amstat.tandfonline. com/doi/abs/10.1080/00401706.2015.1054436

Huling. J.D. and Chien, P. (2022), Fast Penalized Regression and Cross Validation for Tall Data with the oem Package. Journal of Statistical Software 104(6), 1-24. doi:10.18637/jss.v104.i06

```
set.seed(123)
n.obs <- 1e4
n.vars <- 50
true.beta <- c(runif(15, -0.25, 0.25), rep(0, n.vars - 15))</pre>
x <- matrix(rnorm(n.obs * n.vars), n.obs, n.vars)</pre>
y <- rnorm(n.obs, sd = 3) + x \% true.beta
fit <- oem(x = x, y = y,
           penalty = c("lasso", "grp.lasso", "sparse.grp.lasso"),
           groups = rep(1:10, each = 5))
layout(matrix(1:3, ncol = 3))
plot(fit)
plot(fit, which.model = 2)
plot(fit, which.model = "sparse.grp.lasso")
# the oem package has support for
# sparse design matrices
library(Matrix)
xs <- rsparsematrix(n.obs * 25, n.vars * 2, density = 0.01)</pre>
ys <- rnorm(n.obs * 25, sd = 3) + as.vector(xs %*% c(true.beta, rep(0, n.vars)) )</pre>
x.dense <- as.matrix(xs)</pre>
system.time(fit <- oem(x = x.dense, y = ys,</pre>
                        penalty = c("lasso", "grp.lasso"),
                        groups = rep(1:20, each = 5), intercept = FALSE,
                        standardize = FALSE))
system.time(fits <- oem(x = xs, y = ys,</pre>
```

```
penalty = c("lasso", "grp.lasso"),
                         groups = rep(1:20, each = 5), intercept = FALSE,
                         standardize = FALSE, lambda = fit$lambda))
max(abs(fit$beta[[1]] - fits$beta[[1]]))
max(abs(fit$beta[[2]] - fits$beta[[2]]))
# logistic
y <- rbinom(n.obs, 1, prob = 1 / (1 + exp(-x %*% true.beta)))</pre>
system.time(res <- oem(x, y, intercept = FALSE,</pre>
                        penalty = c("lasso", "sparse.grp.lasso", "mcp"),
                        family = "binomial",
                        groups = rep(1:10, each = 5),
                        nlambda = 10,
                        irls.tol = 1e-3, tol = 1e-8))
layout(matrix(1:3, ncol = 3))
plot(res)
plot(res, which.model = 2)
plot(res, which.model = "mcp")
# sparse design matrix
xs <- rsparsematrix(n.obs * 2, n.vars, density = 0.01)</pre>
x.dense <- as.matrix(xs)</pre>
ys <- rbinom(n.obs * 2, 1, prob = 1 / (1 + exp(-x %*% true.beta)))</pre>
system.time(res.gr <- oem(x.dense, ys, intercept = FALSE,</pre>
                           penalty = "grp.lasso",
                           family = "binomial",
                           nlambda = 10,
                           groups = rep(1:5, each = 10),
                           irls.tol = 1e-3, tol = 1e-8))
system.time(res.gr.s <- oem(xs, ys, intercept = FALSE,</pre>
                             penalty = "grp.lasso",
                             family = "binomial",
                             nlambda = 10,
                             groups = rep(1:5, each = 10),
                             irls.tol = 1e-3, tol = 1e-8))
max(abs(res.gr$beta[[1]] - res.gr.s$beta[[1]]))
```

oem.xtx

Orthogonalizing EM with precomputed XtX

### Description

Orthogonalizing EM with precomputed XtX

### Usage

```
oem.xtx(
  xtx,
  xty,
  family = c("gaussian", "binomial"),
 penalty = c("elastic.net", "lasso", "ols", "mcp", "scad", "mcp.net", "scad.net",
   "grp.lasso", "grp.lasso.net", "grp.mcp", "grp.scad", "grp.mcp.net", "grp.scad.net",
    "sparse.grp.lasso"),
  lambda = numeric(0),
  nlambda = 100L,
  lambda.min.ratio = NULL,
  alpha = 1,
  gamma = 3,
  tau = 0.5,
  groups = numeric(0),
  scale.factor = numeric(0),
  penalty.factor = NULL,
  group.weights = NULL,
  maxit = 500L,
  tol = 1e-07,
  irls.maxit = 100L,
  irls.tol = 0.001
)
```

### Arguments

xtx	input matrix equal to crossprod(x) / nrow(x). where x is the design matrix. It is highly recommended to scale by the number of rows in x. If xtx is scaled, xty must also be scaled or else results may be meaningless!
xty	numeric vector of length nvars. Equal to crosprod(x, y) / nobs. It is highly recommended to scale by the number of rows in x.
family	"gaussian" for least squares problems, "binomial" for binary response. (only gaussian implemented currently)
penalty	Specification of penalty type. Choices include:
	• "elastic.net" - elastic net penalty, extra parameters: "alpha"
	• "lasso" - lasso penalty
	<ul> <li>"ols" - ordinary least squares</li> </ul>
	<ul> <li>"mcp" - minimax concave penalty, extra parameters: "gamma"</li> </ul>
	• "scad" - smoothly clipped absolute deviation, extra parameters: "gamma"
	<ul> <li>"mcp.net" - minimax concave penalty + 12 penalty, extra parameters: "gamma", "alpha"</li> </ul>
	<ul> <li>"scad.net" - smoothly clipped absolute deviation + 12 penalty, extra parameters: "gamma", "alpha"</li> </ul>
	• "grp.lasso" - group lasso penalty
	<ul> <li>"grp.lasso.net" - group lasso penalty + 12 penalty, extra parameters: "alpha"</li> </ul>

<ul> <li>"grp.mcp" - group minimax concave penalty, extra parameters: "         "grp.mcp.scad" - group smoothly clipped absolute deviation, extra pa "gamma"         "grp.mcp.net" - group minimax concave penalty + 12 penalty, rameters: "gamma", "alpha"         "grp.mcp.net" - group smoothly clipped absolute deviation + 12 extra parameters: "gamma", "alpha"         "grp.scad.net" - group smoothly clipped absolute deviation + 12 extra parameters: "gamma", "alpha"         "grp.scad.net" - group smoothly clipped absolute deviation + 12 extra parameters: "tau"         Careful consideration is required for the group lasso, group MCP, a SCAD penalties. Groups as specified by the groups argument should be in a sensible manner.         lambda         A user supplied lambda sequence. By default, the program compute lambda sequence based on nlambda and lambda.min.ratio. Supplyin of lambda overrides this.         nlambda         The number of lambda, as a fraction of lambda.max, the (data deriv value (i.e. the smallest value for which all coefficients are zero). Th depends on the sample size nobs relative to the number of variables m default is 0.0001         alpha         mixing value for elastic.net, mcp.net, scad.net, grp.mcp.net, gr penalty applied is (1 - alpha)* (ridge penalty) + alpha* (lasso/mcp/mcp penalty)         groups         A vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0         scale.factor         of length nvars === ncol(xtx) == length(xty) for scaling columns- standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling,         penalty factors can be applied to each coefficient. This is a nu multiplies lambda to allow differential shrinkage, for some row which implies no shrinkage, and that variable is always in inte model. Default is sqr(group size) for all groups.         maxit integer. Maximum number of OEM iterations         row some groups. whic</li></ul>	
<ul> <li>"gamma"</li> <li>"grp.nncp.net" - group minimax concave penalty + 12 penalty, rameters: "gamma", "alpha"</li> <li>"grp.scad.net" - group smoothly clipped absolute deviation + 12 extra parameters: "gamma", "alpha"</li> <li>"sparse.grp.lasso" - sparse group lasso penalty (group lasso extra parameters: "tau"</li> <li>Careful consideration is required for the group lasso, group MCP, a SCAD penalties. Groups as specified by the groups argument should b in a sensible manner.</li> <li>lambda A user supplied lambda sequence. By default, the program compute lambda sequence based on nlambda and lambda.min.ratio. Supplyin of lambda overrides this.</li> <li>nlambda The number of lambda, as a fraction of lambda.max, the (data deriv value (i.e. the smallest value for which all coefficients are zero). Th depends on the sample size nobs relative to the number of variables m default is 0.0001</li> <li>alpha mixing value for elastic.net, mcp.net, scad.net, grp.mcp.net, gr penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp penalty) + tau * (lasso penalty)</li> <li>groups A vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0</li> <li>scale.factor Separate penalty factors can be applied to each coefficient. See the examp All unpenalized variables should be put in group 0</li> <li>scale.factor Separate penalty factors can be applied to each coefficient. This is a nu multiplies lambda to allow differential shrinkage. and that variable is always included in the Default is 1 or all variables.</li> </ul>	-
<ul> <li>rameters: "gamma", "alpha"</li> <li>"grp.scad.net" - group smoothly clipped absolute deviation + 12 extra parameters: "gamma", "alpha"</li> <li>"sparse.grp.lasso" - sparse group lasso penalty (group lasso extra parameters: "tau"</li> <li>Careful consideration is required for the group lasso, group MCP, a SCAD penalties. Groups as specified by the groups argument should be in a sensible manner.</li> <li>lambda A user supplied lambda sequence. By default, the program compute lambda sequence based on nlambda and lambda.min.ratio. Supplyin of lambda vortides this.</li> <li>nlambda The number of lambda values - default is 100.</li> <li>lambda the example size nobs relative to the number of variables walue for which all coefficients are zero). Th depends on the sample size nobs relative to the number of variables m default is 0.0001</li> <li>alpha mixing value for elastic.net, mcp.net, scad.net, grp.mcp.net, grp penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp penalty)</li> <li>gamma tuning parameter for SCAD and MCP penalties. must be &gt;= 1</li> <li>tau mixing value for sparse.grp.lasso. penalty applied is (1 - tau) * (group penalty) + tau * (lasso penalty)</li> <li>groups A vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0</li> <li>scale.factor of length nvars === ncol(xtx) == length(xty) for scaling columns-standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling.</li> <li>penalty.factors applied to each coefficient. This is a num untiplies lambda to allow differential shrinkage. for some which implies no shrinkage, and that variable is always included in the Default is 1 for all variables.</li> <li>group.weights penalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage. for some groups, wh</li></ul>	
extra parameters: "gamma", "alpha"• "sparse.grp.lasso" - sparse group lasso penalty (group lasso extra parameters: "tau"Careful consideration is required for the group lasso, group MCP, a SCAD penalties. Groups as specified by the groups argument should b in a sensible manner.lambdaA user supplied lambda sequence. By default, the program compute lambda overrides this.nlambdaThe number of lambda values - default is 100.lambda.min.ratioSmallest value for lambda, as a fraction of lambda.max, the (data deriv value (i.e. the smallest value for which all coefficients are zero). Th depends on the sample size nobs relative to the number of variables m default is 0.0001alphamixing value for elastic.net,mcp.net, scad.net,grp.mcp.net,grp penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp penalty)groupsA vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0scale.factorof length nvars === ncol(xtx) == length(xty) for scaling columns standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling.penalty.factorSeparate penalty factors can be applied to each coefficient. This is a nu multiplies lambda to allow differential shrinkage. for some groups, which implies no shrinkage, and that variable is always included in th Default is 1 for all variables.group.weightspenalty factors applied to each coefficient. This is a nu multiplies lambda to allow differential shrinkage. for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups.	extra pa-
extra parameters: "tau" Careful consideration is required for the group lasso, group MCP, a SCAD penalties. Groups as specified by the groups argument should b in a sensible manner. lambda A user supplied lambda sequence. By default, the program compute lambda sequence based on nlambda and lambda.min.ratio. Supplyin of lambda overrides this. nlambda The number of lambda, as a fraction of lambda.max, the (data deriv value (i.e. the smallest value for which all coefficients are zero). Th depends on the sample size nobs relative to the number of variables m default is 0.0001 alpha mixing value for elastic.net,mcp.net, scad.net,grp.mcp.net,grp penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp penalty) gamma tuning parameter for SCAD and MCP penalties. must be >= 1 tau mixing value for sparse.grp.lasso.penalty applied is (1 - tau) * (grp penalty) + tau * (lasso penalty) groups A vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0 scale.factor of length nvars === ncol(xtx) == length(xty) for scaling columns- standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling. penalty.factor separate penalty factors can be applied to each coefficient. This is a num multiplies lambda to allow differential shrinkage. Can be 0 for some v which implies no shrinkage, and that variable is always included in th Default is 1 for all variables. group .weights penalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage, for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups. maxit integer. Maximum number of OEM iterations	2 penalty,
SCAD penalties. Groups as specified by the groups argument should be in a sensible manner.lambdaA user supplied lambda sequence. By default, the program compute lambda sequence based on nlambda and lambda.min.ratio. Supplyin of lambda overrides this.nlambdaThe number of lambda values - default is 100. lambda.min.ratiolambda overrides this.Smallest value for lambda, as a fraction of lambda.max, the (data deriv value (i.e. the smallest value for which all coefficients are zero). Th depends on the sample size nobs relative to the number of variables m default is 0.0001alphamixing value for elastic.net,mcp.net, scad.net,grp.mcp.net,grp penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp penalty)gammatuning parameter for SCAD and MCP penalties. must be >= 1taumixing value for sparse.grp.lasso. penalty applied is (1 - tau) * (gr penalty) + tau * (lasso penalty)groupsA vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0scale.factorof length nvars === ncol(xtx) == length(xty) for scaling columns standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling.penalty.factorsSeparate penalty factors can be applied to each coefficient. This is a num multiplies lambda to allow differential shrinkage.group.weightspenalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage. for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups.	) + lasso),
<ul> <li>lambda sequence based on nlambda and lambda.min.ratio. Supplyin of lambda overrides this.</li> <li>nlambda The number of lambda values - default is 100.</li> <li>lambda.min.ratio Smallest value for lambda, as a fraction of lambda.max, the (data derivivalue (i.e. the smallest value for which all coefficients are zero). The depends on the sample size nobs relative to the number of variables midefault is 0.0001</li> <li>alpha mixing value for elastic.net, mcp.net, scad.net, grp.mcp.net, grp penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp penalty)</li> <li>gamma tuning parameter for SCAD and MCP penalties. must be &gt;= 1</li> <li>tau mixing value for sparse.grp.lasso.penalty applied is (1 - tau) * (grp penalty) + tau * (lasso penalty)</li> <li>groups A vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0</li> <li>scale.factor of length nvars === ncol(xtx) == length(xty) for scaling columns standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling.</li> <li>penalty.factor</li> <li>group.weights penalty factors applied to each coefficient. This is a numultiplies lambda to allow differential shrinkage. Can be 0 for some which implies no shrinkage, and that variable is always included in the Default is 1 for all variables.</li> <li>group.weights penalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage. for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups.</li> </ul>	
<pre>lambda.min.ratio Smallest value for lambda, as a fraction of lambda.max, the (data deriv value (i.e. the smallest value for which all coefficients are zero). Th depends on the sample size nobs relative to the number of variables m default is 0.0001 alpha mixing value for elastic.net, mcp.net, scad.net, grp.mcp.net, grp penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp penalty) gamma tuning parameter for SCAD and MCP penalties. must be &gt;= 1 tau mixing value for sparse.grp.lasso.penalty applied is (1 - tau) * (grp penalty) + tau * (lasso penalty) groups A vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0 scale.factor of length nvars === ncol(xtx) == length(xty) for scaling columns standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling. penalty.factor Separate penalty factors can be applied to each coefficient. This is a num multiplies lambda to allow differential shrinkage. Can be 0 for some v which implies no shrinkage, and that variable is always included in th Default is 1 for all variables. group.weights penalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage, for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups. maxit integer. Maximum number of OEM iterations</pre>	
Smallest value for lambda, as a fraction of lambda.max, the (data deriv value (i.e. the smallest value for which all coefficients are zero). Th depends on the sample size nobs relative to the number of variables m default is 0.0001alphamixing value for elastic.net, mcp.net, scad.net, grp.mcp.net, grp penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp penalty)gammatuning parameter for SCAD and MCP penalties. must be >= 1taumixing value for sparse.grp.lasso.penalty applied is (1 - tau) * (gr penalty) + tau * (lasso penalty)groupsA vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0scale.factorof length nvars === ncol(xtx) == length(xty) for scaling columns - standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling.penalty.factorSeparate penalty factors can be applied to each coefficient. This is a num multiplies lambda to allow differential shrinkage. Can be 0 for some v which implies no shrinkage, and that variable is always included in th Default is 1 for all variables.group.weightspenalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage, for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups.maxitinteger. Maximum number of OEM iterations	
<pre>penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp penalty) gamma tuning parameter for SCAD and MCP penalties. must be &gt;= 1 tau mixing value for sparse.grp.lasso.penalty applied is (1 - tau) * (group penalty) + tau * (lasso penalty) groups A vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0 scale.factor of length nvars === ncol(xtx) == length(xty) for scaling columns standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling. penalty.factor Separate penalty factors can be applied to each coefficient. This is a num multiplies lambda to allow differential shrinkage. Can be 0 for some which implies no shrinkage, and that variable is always included in th Default is 1 for all variables. group.weights penalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage. for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups. maxit integer. Maximum number of OEM iterations</pre>	he default
<ul> <li>tau mixing value for sparse.grp.lasso. penalty applied is (1 - tau) * (group penalty) + tau * (lasso penalty)</li> <li>groups A vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0</li> <li>scale.factor of length nvars === ncol(xtx) == length(xty) for scaling columns standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling.</li> <li>penalty.factor Separate penalty factors can be applied to each coefficient. This is a numultiplies lambda to allow differential shrinkage. Can be 0 for some which implies no shrinkage, and that variable is always included in the Default is 1 for all variables.</li> <li>group.weights penalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage, for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups.</li> </ul>	
penalty) + tau * (lasso penalty)groupsA vector of describing the grouping of the coefficients. See the examp All unpenalized variables should be put in group 0scale.factorof length nvars === ncol(xtx) == length(xty) for scaling columns standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling.penalty.factorSeparate penalty factors can be applied to each coefficient. This is a num multiplies lambda to allow differential shrinkage. Can be 0 for some v which implies no shrinkage, and that variable is always included in th Default is 1 for all variables.group.weightspenalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage, for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups.maxitinteger. Maximum number of OEM iterations	
All unpenalized variables should be put in group 0scale.factorof length nvars === ncol(xtx) == length(xty) for scaling columns standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling.penalty.factorSeparate penalty factors can be applied to each coefficient. This is a num multiplies lambda to allow differential shrinkage. Can be 0 for some y which implies no shrinkage, and that variable is always included in th Default is 1 for all variables.group.weightspenalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage, for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups.maxitinteger. Maximum number of OEM iterations	roup lasso
<ul> <li>standard deviation for each column of x is a common choice for scale Coefficients will be returned on original scale. Default is no scaling.</li> <li>penalty.factor</li> <li>Separate penalty factors can be applied to each coefficient. This is a num multiplies lambda to allow differential shrinkage. Can be 0 for some which implies no shrinkage, and that variable is always included in the Default is 1 for all variables.</li> <li>group.weights</li> <li>penalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage. for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups.</li> <li>maxit</li> </ul>	ple below.
<ul> <li>multiplies lambda to allow differential shrinkage. Can be 0 for some which implies no shrinkage, and that variable is always included in the Default is 1 for all variables.</li> <li>group.weights penalty factors applied to each group for the group lasso. Similar to pen this is a number that multiplies lambda to allow differential shrinkage. for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups.</li> <li>maxit integer. Maximum number of OEM iterations</li> </ul>	
<ul> <li>this is a number that multiplies lambda to allow differential shrinkage.</li> <li>for some groups, which implies no shrinkage, and that group is always in the model. Default is sqrt(group size) for all groups.</li> <li>maxit integer. Maximum number of OEM iterations</li> </ul>	variables,
-	. Can be 0
tol convergence tolerance for OEM iterations	
irls.maxit integer. Maximum number of IRLS iterations	
irls.tol convergence tolerance for IRLS iterations. Only used if family != "ga	aussian"

### Value

An object with S3 class "oem"

### References

Huling. J.D. and Chien, P. (2022), Fast Penalized Regression and Cross Validation for Tall Data with the oem Package. Journal of Statistical Software 104(6), 1-24. doi:10.18637/jss.v104.i06

```
set.seed(123)
n.obs <- 1e4
n.vars <- 100
true.beta <- c(runif(15, -0.25, 0.25), rep(0, n.vars - 15))</pre>
x <- matrix(rnorm(n.obs * n.vars), n.obs, n.vars)</pre>
y <- rnorm(n.obs, sd = 3) + x %*% true.beta</pre>
fit <- \text{ oem}(x = x, y = y,
           penalty = c("lasso", "elastic.net",
                         "ols",
                         "mcp",
                                       "scad",
                                       "scad.net",
                         "mcp.net",
                         "grp.lasso", "grp.lasso.net",
                         "grp.mcp", "grp.scad",
                         "sparse.grp.lasso"),
           standardize = FALSE, intercept = FALSE,
           groups = rep(1:20, each = 5))
xtx <- crossprod(x) / n.obs</pre>
xty <- crossprod(x, y) / n.obs</pre>
fit.xtx <- oem.xtx(xtx = xtx, xty = xty,</pre>
                    penalty = c("lasso", "elastic.net",
                                 "ols",
                                 "mcp",
                                              "scad",
                                              "scad.net",
                                 "mcp.net",
                                 "grp.lasso", "grp.lasso.net",
                                 "grp.mcp", "grp.scad",
                                 "sparse.grp.lasso"),
                    groups = rep(1:20, each = 5))
max(abs(fit$beta[[1]][-1,] - fit.xtx$beta[[1]]))
max(abs(fit$beta[[2]][-1,] - fit.xtx$beta[[2]]))
layout(matrix(1:2, ncol = 2))
plot(fit.xtx)
plot(fit.xtx, which.model = 2)
```

oemfit

### Description

These functions have been renamed and deprecated in **oem**: oemfit() (use oem()), cv.oemfit() (use cv.oem()), print.oemfit(), plot.oemfit(), predict.oemfit(), and coef.oemfit().

### Usage

```
oemfit(
  formula,
  data = list(),
  lambda = NULL,
  nlambda = 100,
  lambda.min.ratio = NULL,
  tolerance = 0.001,
 maxIter = 1000,
  standardized = TRUE,
  numGroup = 1,
  penalty = c("lasso", "scad", "ols", "elastic.net", "ngarrote", "mcp"),
  alpha = 3,
  evaluate = 0,
  condition = -1
)
cv.oemfit(
  formula,
  data = list(),
  lambda = NULL,
  type.measure = c("mse", "mae"),
  ...,
  nfolds = 10,
  foldid,
  penalty = c("lasso", "scad", "elastic.net", "ngarrote", "mcp")
)
## S3 method for class 'oemfit'
plot(
  х,
 xvar = c("norm", "lambda", "loglambda", "dev"),
  xlab = iname,
 ylab = "Coefficients",
  . . .
)
## S3 method for class 'oemfit'
```

```
predict(
   object,
   newx,
   s = NULL,
   type = c("response", "coefficients", "nonzero"),
   ...
)
## S3 method for class 'oemfit'
print(x, digits = max(3, getOption("digits") - 3), ...)
```

### Arguments

formula	an object of 'formula' (or one that can be coerced to that class): a symbolic description of the model to be fitted. The details of model specification are given under 'Details'	
data	an optional data frame, list or environment (or object coercible by 'as.data.frame' to a data frame) containing the variables in the model. If not found in 'data', the variables are taken from 'environment(formula)', typically the environment from which 'oemfit' is called.	
lambda	A user supplied lambda sequence. Typical usage is to have the program compute its own lambda sequence based on nlambda and lambda.min.ratio. Supplying a value of lambda overrides this. WARNING: use with care. Do not supply a single value for lambda (for predictions after CV use predict() instead). Supply instead a decreasing sequence of lambda values. oemfit relies on its warms starts for speed, and its often faster to fit a whole path than compute a single fit.	
nlambda The number of lambda values - default is 100.		
lambda.min.rati		
	Smallest value for lambda, as a fraction of lambda.max, the (data derived) entry value (i.e. the smallest value for which all coefficients are zero). The default depends on the sample size nobs relative to the number of variables nvars. If nobs > nvars, the default is $0.0001$ , close to zero. If nobs < nvars, the default is $0.001$ . A very small value of lambda.min.ratio will lead to a saturated fit in the nobs < nvars case.	
tolerance	Convergence tolerance for OEM. Each inner OEM loop continues until the max- imum change in the objective after any coefficient update is less than tolerance. Defaults value is 1E-3.	
maxIter	Maximum number of passes over the data for all lambda values; default is 1000.	
standardized	Logical flag for x variable standardization, prior to fitting the model sequence. The coefficients are always returned on the original scale. Default is standardize=TRUE. If variables are in the same units already, you might not wish to standardize.	
numGroup	Integer value for the number of groups to use for OEM fitting. Default is 1.	
penalty	type in lower letters. Different types include 'lasso', 'scad', 'ols' (ordinary least square), 'elastic-net', 'ngarrote' (non-negative garrote) and 'mcp'.	
alpha	alpha value for scad and mcp.	

### plot.oem

evaluate	debugging argument
condition	Debugging for different ways of calculating OEM.
type.measure	<pre>type.measure measure to evaluate for cross-validation. type.measure = "mse" (mean squared error) or type.measure = "mae" (mean absolute error)</pre>
	arguments to be passed to oemfit()
nfolds	number of folds for cross-validation. default is 10.
foldid	an optional vector of values between 1 and nfold specifying which fold each observation belongs to.
x	fitted oemfit object
xvar	what is on the X-axis. "norm" plots against the L1-norm of the coefficients, "lambda" against the log-lambda sequence, and "dev" against the percent deviance explained.
xlab	x-axis label
ylab	y-axis label
object	fitted oemfit object
newx	matrix of new values for x at which predictions are to be made. Must be a matrix.
S	Value(s) of the penalty parameter lambda at which predictions are required. Default is the entire sequence used to create the model.
type	not used.
digits	significant digits in print out.

### Details

The sequence of models implied by 'lambda' is fit by OEM algorithm.

### Author(s)

Bin Dai

plot.oem

Plot method for Orthogonalizing EM fitted objects

### Description

Plot method for Orthogonalizing EM fitted objects

Plot method for Orthogonalizing EM fitted objects

### Usage

```
## S3 method for class 'oem'
plot(
  х,
 which.model = 1,
 xvar = c("norm", "lambda", "loglambda", "dev"),
  labsize = 0.6,
 xlab = iname,
 ylab = NULL,
 main = x$penalty[which.model],
  . . .
)
## S3 method for class 'cv.oem'
plot(x, which.model = 1, sign.lambda = 1, ...)
## S3 method for class 'xval.oem'
plot(
 х,
 which.model = 1,
  type = c("cv", "coefficients"),
 xvar = c("norm", "lambda", "loglambda", "dev"),
 labsize = 0.6,
 xlab = iname,
 ylab = NULL,
 main = x$penalty[which.model],
 sign.lambda = 1,
  . . .
)
```

### Arguments

Х	fitted "oem" model object
which.model	If multiple penalties are fit and returned in the same oem object, the which.model argument is used to specify which model to plot. For example, if the oem object "oemobj" was fit with argument penalty = c("lasso", "grp.lasso"), then which.model = 2 provides a plot for the group lasso model.
xvar	What is on the X-axis. "norm" plots against the L1-norm of the coefficients, "lambda" against the log-lambda sequence, and "dev" against the percent deviance explained.
labsize	size of labels for variable names. If labsize = 0, then no variable names will be plotted
xlab	label for x-axis
ylab	label for y-axis
main	main title for plot
	other graphical parameters for the plot

### plot.oem

sign.lambda	Either plot against log(lambda) (default) or its negative if sign.lambda = -1.
type	<pre>one of "cv" or "coefficients". type = "cv" will produce a plot of cross val- idation results like plot.cv.oem. type = "coefficients" will produce a coeffi- cient path plot like plot.oem()</pre>

```
set.seed(123)
n.obs <- 1e4
n.vars <- 100
n.obs.test <- 1e3
true.beta <- c(runif(15, -0.5, 0.5), rep(0, n.vars - 15))</pre>
x <- matrix(rnorm(n.obs * n.vars), n.obs, n.vars)</pre>
y <- rnorm(n.obs, sd = 3) + x %*% true.beta</pre>
fit <- oem(x = x, y = y, penalty = c("lasso", "grp.lasso"), groups = rep(1:10, each = 10))</pre>
layout(matrix(1:2, ncol = 2))
plot(fit, which.model = 1)
plot(fit, which.model = 2)
set.seed(123)
n.obs <- 1e4
n.vars <- 100
n.obs.test <- 1e3
true.beta <- c(runif(15, -0.5, 0.5), rep(0, n.vars - 15))</pre>
x <- matrix(rnorm(n.obs * n.vars), n.obs, n.vars)</pre>
y <- rnorm(n.obs, sd = 3) + x %*% true.beta</pre>
fit <- cv.oem(x = x, y = y, penalty = c("lasso", "grp.lasso"), groups = rep(1:10, each = 10))</pre>
layout(matrix(1:2, ncol = 2))
plot(fit, which.model = 1)
plot(fit, which.model = "grp.lasso")
set.seed(123)
n.obs <- 1e4
n.vars <- 100
n.obs.test <- 1e3
true.beta <- c(runif(15, -0.5, 0.5), rep(0, n.vars - 15))</pre>
x <- matrix(rnorm(n.obs * n.vars), n.obs, n.vars)</pre>
y \leq norm(n.obs, sd = 3) + x \% true.beta
fit <- xval.oem(x = x, y = y, penalty = c("lasso", "grp.lasso"), groups = rep(1:10, each = 10))</pre>
layout(matrix(1:4, ncol = 2))
```

```
plot(fit, which.model = 1)
plot(fit, which.model = 2)
plot(fit, which.model = 1, type = "coef")
plot(fit, which.model = 2, type = "coef")
```

```
predict.cv.oem
```

Prediction function for fitted cross validation oem objects

### Description

Prediction function for fitted cross validation oem objects

### Usage

```
## S3 method for class 'cv.oem'
predict(
   object,
   newx,
   which.model = "best.model",
   s = c("lambda.min", "lambda.lse"),
   ...
)
```

### Arguments

object	fitted "cv.oem" model object
newx	Matrix of new values for x at which predictions are to be made. Must be a matrix; can be sparse as in the CsparseMatrix objects of the <b>Matrix</b> package This argument is not used for type = c("coefficients", "nonzero")
which.model	If multiple penalties are fit and returned in the same oem object, the which.model argument is used to specify which model to make predictions for. For example, if the oem object "oemobj" was fit with argument penalty = c("lasso", "grp.lasso"), then which.model = 2 provides predictions for the group lasso model. For predict.cv.oem(), can specify "best.model" to use the best model as estimated by cross-validation
S	Value(s) of the penalty parameter lambda at which predictions are required. Default is the entire sequence used to create the model. For predict.cv.oem(), can also specify "lambda.lse" or "lambda.min" for best lambdas estimated by cross validation
	used to pass the other arguments for predict.oem

### Value

An object depending on the type argument

### predict.oem

### Examples

```
set.seed(123)
n.obs <- 1e4
n.vars <- 100
n.obs.test <- 1e3
true.beta <- c(runif(15, -0.5, 0.5), rep(0, n.vars - 15))</pre>
x <- matrix(rnorm(n.obs * n.vars), n.obs, n.vars)</pre>
y \leq norm(n.obs, sd = 3) + x \% true.beta
x.test <- matrix(rnorm(n.obs.test * n.vars), n.obs.test, n.vars)</pre>
y.test <- rnorm(n.obs.test, sd = 3) + x.test %*% true.beta</pre>
fit <- cv.oem(x = x, y = y,
              penalty = c("lasso", "grp.lasso"),
              groups = rep(1:10, each = 10),
              nlambda = 10)
preds.best <- predict(fit, newx = x.test, type = "response", which.model = "best.model")</pre>
apply(preds.best, 2, function(x) mean((y.test - x) ^ 2))
preds.gl <- predict(fit, newx = x.test, type = "response", which.model = "grp.lasso")</pre>
apply(preds.gl, 2, function(x) mean((y.test - x) ^ 2))
preds.l <- predict(fit, newx = x.test, type = "response", which.model = 1)</pre>
apply(preds.1, 2, function(x) mean((y.test - x) ^ 2))
```

predict.oem

Prediction method for Orthogonalizing EM fitted objects

#### Description

Prediction method for Orthogonalizing EM fitted objects

#### Usage

```
## S3 method for class 'oem'
predict(
   object,
   newx,
   s = NULL,
   which.model = 1,
   type = c("link", "response", "coefficients", "nonzero", "class"),
   ...
)
```

### Arguments

object	fitted "oem" model object
newx	Matrix of new values for x at which predictions are to be made. Must be a matrix; can be sparse as in the CsparseMatrix objects of the <b>Matrix</b> package. This argument is not used for type=c("coefficients", "nonzero")
S	Value(s) of the penalty parameter lambda at which predictions are required. De- fault is the entire sequence used to create the model.
which.model	If multiple penalties are fit and returned in the same oem object, the which.model argument is used to specify which model to make predictions for. For example, if the oem object oemobj was fit with argument penalty = c("lasso", "grp.lasso"), then which.model = 2 provides predictions for the group lasso model.
type	Type of prediction required. type = "link" gives the linear predictors for the "binomial" model; for "gaussian" models it gives the fitted values. type = "response" gives the fitted probabilities for "binomial". type = "coefficients" computes the coefficients at the requested values for s. type = "class" applies only to "binomial" and produces the class label corresponding to the maximum probability.
•••	not used

### Value

An object depending on the type argument

```
set.seed(123)
n.obs <- 1e4
n.vars <- 100
n.obs.test <- 1e3
true.beta <- c(runif(15, -0.5, 0.5), rep(0, n.vars - 15))</pre>
x <- matrix(rnorm(n.obs * n.vars), n.obs, n.vars)</pre>
y \leq rnorm(n.obs, sd = 3) + x \% true.beta
x.test <- matrix(rnorm(n.obs.test * n.vars), n.obs.test, n.vars)</pre>
y.test <- rnorm(n.obs.test, sd = 3) + x.test %*% true.beta</pre>
fit <- oem(x = x, y = y,
           penalty = c("lasso", "grp.lasso"),
           groups = rep(1:10, each = 10),
           nlambda = 10)
preds.lasso <- predict(fit, newx = x.test, type = "response", which.model = 1)</pre>
preds.grp.lasso <- predict(fit, newx = x.test, type = "response", which.model = 2)</pre>
apply(preds.lasso,
                        2, function(x) mean((y.test - x) ^{2})
apply(preds.grp.lasso, 2, function(x) mean((y.test - x) ^ 2))
```

predict.xval.oem Prediction function for fitted cross validation oem objects

### Description

Prediction function for fitted cross validation oem objects

### Usage

```
## S3 method for class 'xval.oem'
predict(
   object,
   newx,
   which.model = "best.model",
   s = c("lambda.min", "lambda.lse"),
   ...
)
```

### Arguments

object	fitted "cv.oem" model object
newx	Matrix of new values for x at which predictions are to be made. Must be a matrix; can be sparse as in the CsparseMatrix objects of the <b>Matrix</b> package This argument is not used for type=c("coefficients","nonzero")
which.model	If multiple penalties are fit and returned in the same oem object, the which.model argument is used to specify which model to make predictions for. For example, if the oem object "oemobj" was fit with argument penalty = c("lasso", "grp.lasso"), then which.model = 2 provides predictions for the group lasso model. For predict.cv.oem(), can specify "best.model" to use the best model as estimated by cross-validation
S	Value(s) of the penalty parameter lambda at which predictions are required. De- fault is the entire sequence used to create the model. For predict.cv.oem, can also specify "lambda.lse" or "lambda.min" for best lambdas estimated by cross validation
	used to pass the other arguments for predict.oem()

### Value

An object depending on the type argument

```
set.seed(123)
n.obs <- 1e4
n.vars <- 100
n.obs.test <- 1e3</pre>
```

print.summary.cv.oem print method for summary.cv.oem objects

### Description

print method for summary.cv.oem objects

### Usage

```
## S3 method for class 'summary.cv.oem'
print(x, digits, ...)
```

### Arguments

Х	a "summary.cv.oem" object
digits	digits to display
	not used

summary.cv.oem

summary method for cross validation Orthogonalizing EM fitted objects

### Description

summary method for cross validation Orthogonalizing EM fitted objects summary method for cross validation Orthogonalizing EM fitted objects

### Usage

```
## S3 method for class 'cv.oem'
summary(object, ...)
```

## S3 method for class 'xval.oem'
summary(object, ...)

### Arguments

object	fitted "cv.oem" object
	not used

xval.oem

Fast cross validation for Orthogonalizing EM

### Description

Fast cross validation for Orthogonalizing EM

#### Usage

```
xval.oem(
    x,
    y,
    nfolds = 10L,
    foldid = NULL,
    type.measure = c("mse", "deviance", "class", "auc", "mae"),
    ncores = -1,
    family = c("gaussian", "binomial"),
    penalty = c("elastic.net", "lasso", "ols", "mcp", "scad", "mcp.net", "scad.net",
    "grp.lasso", "grp.lasso.net", "grp.mcp", "grp.scad", "grp.mcp.net", "grp.scad.net",
    "sparse.grp.lasso"),
    weights = numeric(0),
    lambda = numeric(0),
    nlambda = 100L,
```

```
lambda.min.ratio = NULL,
alpha = 1,
gamma = 3,
tau = 0.5,
groups = numeric(0),
penalty.factor = NULL,
group.weights = NULL,
standardize = TRUE,
intercept = TRUE,
maxit = 500L,
tol = 1e-07,
irls.maxit = 100L,
irls.tol = 0.001,
compute.loss = FALSE
)
```

### Arguments

x	input matrix of dimension n x p (sparse matrices not yet implemented). Each row is an observation, each column corresponds to a covariate. The xval.oem() function is optimized for n » p settings and may be very slow when $p > n$ , so please use other packages such as glmnet, ncvreg, grpreg, or gglasso when p > n or p approx n.
у	numeric response vector of length nobs = nrow(x).
nfolds	integer number of cross validation folds. 3 is the minimum number allowed. defaults to 10
foldid	an optional vector of values between 1 and nfold specifying which fold each observation belongs to.
type.measure	<pre>measure to evaluate for cross-validation. The default is type.measure = "deviance", which uses squared-error for gaussian models (a.k.a type.measure = "mse" there), deviance for logistic regression. type.measure = "class" applies to binomial only. type.measure = "auc" is for two-class logistic regression only. type.measure="mse" or type.measure="mae" (mean absolute error) can be used by all models; they measure the deviation from the fitted mean to the re- sponse.</pre>
ncores	Integer scalar that specifies the number of threads to be used
family	"gaussian" for least squares problems, "binomial" for binary response (not implemented yet).
penalty	Specification of penalty type. Choices include:
	<ul> <li>"elastic.net" - elastic net penalty, extra parameters: "alpha"</li> <li>"lasso" - lasso penalty</li> <li>"ols" - ordinary least squares</li> <li>"mcp" - minimax concave penalty, extra parameters: "gamma"</li> <li>"scad" - smoothly clipped absolute deviation, extra parameters: "gamma"</li> <li>"mcp.net" - minimax concave penalty + 12 penalty, extra parameters: "gamma", "alpha"</li> </ul>

	<ul> <li>"scad.net" - smoothly clipped absolute deviation + 12 penalty, extra parameters: "gamma", "alpha"</li> </ul>
	• "grp.lasso" - group lasso penalty
	<ul> <li>"grp.lasso.net" - group lasso penalty + 12 penalty, extra parameters: "alpha"</li> </ul>
	• "grp.mcp" - group minimax concave penalty, extra parameters: "gamma"
	<ul> <li>"grp.scad" - group smoothly clipped absolute deviation, extra parameters:</li> <li>"gamma"</li> </ul>
	<ul> <li>"grp.mcp.net" - group minimax concave penalty + 12 penalty, extra parameters: "gamma", "alpha"</li> </ul>
	<ul> <li>"grp.scad.net" - group smoothly clipped absolute deviation + 12 penalty, extra parameters: "gamma", "alpha"</li> </ul>
	• "sparse.grp.lasso" - sparse group lasso penalty (group lasso + lasso), extra parameters: "tau"
	Careful consideration is required for the group lasso, group MCP, and group SCAD penalties. Groups as specified by the groups argument should be chosen in a sensible manner.
weights	observation weights. defaults to 1 for each observation (setting weight vector to length 0 will default all weights to 1)
lambda	A user supplied lambda sequence. By default, the program computes its own lambda sequence based on nlambda and lambda.min.ratio. Supplying a value of lambda overrides this.
nlambda lambda.min.rati	The number of lambda values - default is 100.
	Smallest value for lambda, as a fraction of lambda.max, the (data derived) entry value (i.e. the smallest value for which all coefficients are zero). The default depends on the sample size nobs relative to the number of variables nvars. If nobs > nvars, the default is 0.0001, close to zero.
alpha	<pre>mixing value for elastic.net, mcp.net, scad.net, grp.mcp.net, grp.scad.net. penalty applied is (1 - alpha) * (ridge penalty) + alpha * (lasso/mcp/mcp/grp.lasso penalty)</pre>
gamma	tuning parameter for SCAD and MCP penalties. must be $>= 1$
tau	mixing value for sparse.grp.lasso. penalty applied is (1 - tau) * (group lasso penalty) + tau * (lasso penalty)
groups	A vector of describing the grouping of the coefficients. See the example below. All unpenalized variables should be put in group 0
penalty.factor	Separate penalty factors can be applied to each coefficient. This is a number that multiplies lambda to allow differential shrinkage. Can be 0 for some variables, which implies no shrinkage, and that variable is always included in the model. Default is 1 for all variables.
group.weights	penalty factors applied to each group for the group lasso. Similar to penalty.factor, this is a number that multiplies lambda to allow differential shrinkage. Can be 0 for some groups, which implies no shrinkage, and that group is always included in the model. Default is sqrt(group size) for all groups.

standardize	Logical flag for x variable standardization, prior to fitting the models. The co- efficients are always returned on the original scale. Default is standardize = TRUE. If variables are in the same units already, you might not wish to standard- ize.
intercept	Should intercept(s) be fitted (default = TRUE) or set to zero (FALSE)
maxit	integer. Maximum number of OEM iterations
tol	convergence tolerance for OEM iterations
irls.maxit	integer. Maximum number of IRLS iterations
irls.tol	convergence tolerance for IRLS iterations. Only used if family != "gaussian"
compute.loss	should the loss be computed for each estimated tuning parameter? Defaults to FALSE. Setting to TRUE will dramatically increase computational time

### Value

An object with S3 class "xval.oem"

### References

Huling. J.D. and Chien, P. (2022), Fast Penalized Regression and Cross Validation for Tall Data with the oem Package. Journal of Statistical Software 104(6), 1-24. doi:10.18637/jss.v104.i06

```
set.seed(123)
n.obs <- 1e4
n.vars <- 100
true.beta <- c(runif(15, -0.25, 0.25), rep(0, n.vars - 15))</pre>
x <- matrix(rnorm(n.obs * n.vars), n.obs, n.vars)</pre>
y <- rnorm(n.obs, sd = 3) + x %*% true.beta</pre>
system.time(fit <- oem(x = x, y = y,</pre>
                        penalty = c("lasso", "grp.lasso"),
                        groups = rep(1:20, each = 5)))
system.time(xfit <- xval.oem(x = x, y = y,</pre>
                              penalty = c("lasso", "grp.lasso"),
                              groups = rep(1:20, each = 5)))
system.time(xfit2 <- xval.oem(x = x, y = y,</pre>
                               penalty = c("lasso", "grp.lasso",
                                            "mcp", "scad",
                                            "mcp.net",
                                                         "scad.net",
                                            "grp.lasso", "grp.lasso.net",
                                            "grp.mcp", "grp.scad",
                                            "sparse.grp.lasso"),
                               groups = rep(1:20, each = 5)))
```

# Index

big.oem, 2
cv.oem, 6, 17
cv.oemfit (oemfit), 17

logLik.cv.oem(logLik.oem), 8
logLik.oem, 8
logLik.xval.oem(logLik.oem), 8

oem, 9, 17
oem-deprecated (oemfit), 17
oem.xtx, 13
oemfit, 17

plot.cv.oem (plot.oem), 19
plot.oem, 19
plot.oemfit (oemfit), 17
plot.xval.oem (plot.oem), 19
predict.cv.oem, 22
predict.oem, 23
predict.oemfit (oemfit), 17
predict.xval.oem, 25
print.oemfit (oemfit), 17
print.summary.cv.oem, 26

summary.cv.oem, 27
summary.xval.oem(summary.cv.oem), 27

xval.oem, 27