# Package 'Icens'

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Title NPMLE for Censored and Truncated Data
<b>Description</b> Many functions for computing the NPMLE for censored and truncated data.
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# Description

Bisect

Most of the optimizations in *Icens* have a one dimensional root-finding component. Since the quantities involved are generally restricted to a subset of [0,1] we use bisection to find the roots.

An implementation of the bisection algorithm for root finding.

# Usage

```
Bisect(tA, pvec, ndir, Meps, tolbis=1e-07)
```

# **Arguments**

tA The transpose of the clique matrix.

pvec The current estimate of the probability vector.

ndir The direction to explore.

Meps Machine epsilon, elements of pvec that are less than this are assumed to be zero.

tolbis The tolerance used to determine if the algorithm has converged.

#### **Details**

We search from pvec in the direction ndir to obtain the new value of pvec that maximizes the likelihood.

## Value

The new estimate of pvec.

## Author(s)

Alain Vandal and Robert Gentleman.

BVcliques 3

# References

Any book on optimization.

BVcliques	Find the bivariate cliques from the marginal data.	

# **Description**

The maximal cliques of the intersection graph are obtained by first finding the cliques for the marginal data and then combining them using the algorithm in Gentleman and Vandal (1999).

# Usage

```
BVcliques(intvlx, intvly, Lxopen=TRUE, Rxopen=FALSE, Lyopen=TRUE, Ryopen=FALSE)
```

#### **Arguments**

intvlx	The cliques for one marginal component, alternatively the marginal intervals can be supplied.
intvly	The cliques for the other marginal component, alternatively the marginal intervals can be supplied.
Lxopen	Boolean indicating whether the left end point in the x coordinate is open.
Rxopen	Boolean indicating whether the right end point in the x coordinate is open.
Lyopen	Boolean indicating whether the left end point in the y coordinate is open.
Ryopen	Boolean indicating whether the right end point in the y coordinate is open.

# Value

A list of the maximal cliques of the intersection graph of the data.

# Author(s)

A. Vandal and R. Gentleman

# References

*Graph–Theoretical Aspects of Bivariate Censored Data*, R. Gentleman and A. Vandal, 1999, submitted.

# See Also

```
BVclmat, BVsupport
```

```
data(cmv)
cmv.cl <- BVcliques(cmv[,1:2], cmv[,3:4], Lxopen=FALSE, Lyopen=FALSE )</pre>
```

4 BVcImat

BVclmat

Compute the clique matrix from the clique list.

# **Description**

Given the clique list, obtained from BVcliques, the clique matrix is obtained. This is the m (number of cliques) by n (number of observations) matrix. A[i,j] is one if individual j is in maximal clique i.

#### Usage

```
BVclmat(cliques)
```

# **Arguments**

cliques

The clique list.

#### Value

The m by n clique matrix.

# Author(s)

A. Vandal and R. Gentleman

#### References

Graph-Theoretical Aspects of Bivariate Censored Data, R. Gentleman and A. Vandal, 1999, submitted.

# See Also

```
BVcliques, BVsupport
```

```
data(cmv)
bcl <- BVcliques(cmv[,1:2], cmv[,3:4])
A <- BVclmat(bcl)</pre>
```

BVsupport 5

BVsupport	Compute the support for the cliques of a bivariate intersection graph.

# **Description**

Given the regions where the events occurred and the cliques of the intersection graph the support of the cliques is computed. For each clique it is the intersection of the event time regions for all observations in that clique.

# Usage

```
BVsupport(intvlx, intvly, cliques=BVcliques(intvlx, intvly))
```

# **Arguments**

intvlx	The event time intervals for one dimension.
intvly	The event time intervals for the other dimension.
cliques	The list of maximal cliques of the intersection graph, optionally.

#### Value

An m by 4 matrix containing the corners of the intervals of support for the maximal cliques of the intersection graph corresponding to the first two arguments to the function.

# Author(s)

A. Vandal and R. Gentleman

## References

Graph-Theoretical Aspects of Bivariate Censored Data, R. Gentleman and A. Vandal, 1999, submitted.

# See Also

```
BVcliques, BVclmat
```

```
data(cmv)
cmv.cl <- BVcliques(cmv[,1:2], cmv[,3:4])
boxes <- BVsupport(cmv[,1:2], cmv[,3:4], cmv.cl)</pre>
```

6 cosmesis

Data on times to shedding of cytomegalovirus and to colonization of mycobacterium avium complex.

# **Description**

The cmv data frame has 204 rows and 4 columns. The intervals should be treated as closed at both ends to replicate the analysis in Betensky and Finkelstein.

#### **Format**

This data frame contains the following columns:

cmvL The left end of the CMV shedding interval.

cmvR The right end of the CMV shedding interval.

macL The left end of the MAC colonization interval.

macR The right end of the MAC colonization interval.

#### **Details**

Betensky and Finkelstein, 1999 present data from the AIDS Clinical Trials Group protocol ACTG 181. This was a natural history substudy of a comparative trial. Patients were scheduled for clinic visits during follow–up and data was collected on the time until two events; shedding of cytomegalovirus (CMV) in the urine and blood and for colonization of mycobacterium avium complex (MAC) in the sputum or stool.

## Source

Betensky, R. A. and Finkelstein, D. M., 1999, A nonparametric maximum likelihood estimator for bivariate interval censored data, Statistics in Medicine,

# **Examples**

data(cmv)

cosmesis

The time taken until cosmetic deterioration of breast cosmesis.

#### Description

The cosmesis data frame has 95 rows and 3 columns.

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#### **Format**

This data frame contains the following columns:

L The left end point of the cosmetic deterioration interval.

**R** The right end point of the cosmetic deterioration interval.

**Trt** The treatment indicator. It is zero for those that received radiotherapy.

#### Source

A semiparametric model for regression analysis of interval-censored failure time data, D. M. Finkelstein and R. A. Wolfe, 1985, Biometrics.

# **Examples**

```
data(cosmesis)
```

EM	A function to compute the NPMLE of p based on the incidence matrix
	$\Delta$

# Description

The incidence matrix, A is the m by n matrix that represents the data. There are m probabilities that must be estimated. The EM, or expectation maximization, method is applied to these data.

# Usage

```
EM(A, pvec, maxiter=500, tol=1e-12)
```

#### **Arguments**

A The incidence matrix.

pvec The probability vector.

maxiter The maximum number of iterations.
tol The tolerance used to judge convergence.

# **Details**

Lots.

# Value

An object of class icsurv containing the following components:

pf The NPMLE of the probability vector.

numiter The number of iterations used.

converge A boolean indicating whether the algorithm converged.

intmap If present indicates the real representation of the support for the values in pf.

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#### Author(s)

Alain Vandal and Robert Gentleman.

#### References

The EM algorithm applied to the maximal cliques of the intersection graph of the censored data. *The empirical distribution function with arbitrarily grouped, censored and truncated data*, B. W. Turnbull, 1976, JRSS;B.

#### See Also

```
VEM, ISDM, EMICM, PGM
```

#### **Examples**

```
data(cosmesis)
csub1 <- subset(cosmesis, subset= Trt==0, select=c(L,R))
EM(csub1)
data(pruitt)
EM(pruitt)</pre>
```

**EMICM** 

Compute the NPMLE for censored data using the EMICM.

# Description

An implementation of the hybrid EM ICM (Iterative convex minorant) estimator of the distribution function proposed by Wellner and Zahn (1997).

# Usage

```
EMICM(A, EMstep=TRUE, ICMstep=TRUE, keepiter=FALSE, tol=1e-07,
maxiter=1000)
```

## **Arguments**

,	4	Either the m by n clique matrix or the n by 2 matrix containing the event time intervals.
E	EMstep	Boolean, indicating whether to take an EM step in the iteration.
	ICMstep	Boolean, indicating whether to take an ICM step.
ļ	keepiter	Boolean determining whether to keep the iteration states.
1	tol	The maximal L1 distance between successive estimates before stopping iteration.
r	maxiter	The maximal number of iterations to perform before stopping.

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#### **Details**

Lots, and they're complicated too!

#### Value

An object of class icsurv containing the following components:

pf The estimated probabilities.

sigma The NPMLE of the survival function on the maximal antichains.

 $\label{thm:condition} \mbox{ The diagonal of the likelihood function's second derivative.}$ 

lastchange A vector of differences between the last two iterations.

numiter The total number of iterations performed.

iter Is only present if keepiter is true; states of sigma during the iteration.

The real representation associated with the probabilities reported in pf.

# Author(s)

Alain Vandal and Robert Gentleman

# References

A hybrid algorithm for computation of the nonparametric maximum likelihood estimator from censored data, J. A. Wellner and Y. Zhan, 1997, JASA.

#### See Also

```
EM, VEM, PGM
```

#### **Examples**

```
data(cosmesis)
csub1 <- subset(cosmesis, subset=Trt==0, select=c(L,R))
EMICM(csub1)
data(pruitt)
EMICM(pruitt)</pre>
```

hiv

Intervals for infection time and disease onset for 257 hemophiliac patients.

# **Description**

The hiv data frame has 257 rows and 4 columns.

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#### **Format**

This data frame contains the following columns:

- yL The left end point of the infection time interval.
- **yR** The right end point of the infection time interval.
- **zL** The left end point of the disease onset interval.
- **zR** The right end point of the disease onset interval.
- **Age** Coded as 1 if the estimated age at infection was less than 20 and 2 if the estimated age at infection was greater than 20.
- **Trt** Treatment, Light or Heavy

#### Details

The setting is as follows. Individuals were infected with the HIV virus at some unknown time they subsequently develop AIDS at a second unknown time. The data consist of two intervals,  $(y_L, y_R)$  and  $(z_L, z_R)$ , such that the infection time was in the first interval and the time of disease onset was in the second interval. A quantity of interest is the incubation time of the disease which is T = Z - Y. The authors argue persuasively that this should be considered as bivariate interval censored data. They note that simply forming the differences  $(z_L - y_R, z_R - y_L)$  and analysing the resultant data assumes an incorrect likelihood. DeGruttola and Lagakos transform the problem slightly to study the joint distribution of Y and T = Z - Y. This is equivalent to estimating the joint distribution of Z and Y then transforming. The data, as reported, have been discretized into six month intervals.

We use the data as reported in Table 1 of DeGruttola and Lagakos, 1989. The patients were 257 persons with Type A or B hemophilia treated at two hospitals in France. They were then examined intermittently (as they came in for treatment?) and their HIV and AIDS status was determined. Kim, De Gruttola and Lagakos report some covariate information and their paper is concerned with the modeling of that information. In this paper we concentrate only on the event times and ignore the covariate information; that topic being worthy of separate investigation.

## Source

DeGruttola, V. and Lagakos, S.W., 1989, Analysis of doubly-censored survival data, with application to AIDS, Biometrics.

Kim, Mimi Y. and De Gruttola, Victor G. and Lagakos, Stephen W., 1993, *Analyzing Doubly Censored Data With Covariates, With Application to AIDS*, Biometrics.

#### **Examples**

data(hiv)

Icens-internal

Icens-internal	Internal Icens functions

# **Description**

Internal Icens functions

# **Details**

These are not to be called by the user.

icsurv The class of objects returned by the estimation routines in the Icens library.

# **Description**

An object of class icsurv must contain the following components:

converge A boolean indicating whether the iteration producing pf converged.

**pf** The probability vector.

It can optionally contain any of the following components:

**clmat** The clique matrix used to obtain pf.

intmap The real representations of the support for the components of pf.

iter A matrix containing every iterative estimate of pf, useful for debugging.

**Ival** The value of the **log** likelihood at pf.

numiter The number of iterations taken.

sigma The cumulative sum of pf.

weights Weights used in the EMICM algorithm.

# Author(s)

Alain Vandal and Robert Gentleman.

#### See Also

VEM, ISDM, EMICM, PGM, EM

12 ISDM

ISDM	Estimate the NPMLE of censored data using the ISDM method pro-
	posed in Lesperance and Kalbfleisch (19

# **Description**

ISDM is a method for estimating the NPMLE of censored data.

# Usage

```
ISDM(A, pvec, maxiter=500, tol=1e-07, tolbis=1e-08, verbose=FALSE)
```

# **Arguments**

A The m by n incidence, or clique, matrix. Or the n by 2 matrix containing the

event intervals.

pvec An initial estimate of the probability vector; not required.

maxiter Maximum number of iterations to be made.

tol The tolerance used to determine convergence.

tolbis A second tolerance used for the steps.

verbose Boolean, should verbose output be printed.

#### **Details**

Lots of complicated stuff should go here.

# Value

A list containing:

pf The estimated NPMLE of the probability vector.

numiter The number of iterations performed.

## Author(s)

Alain Vandal and Robert Gentleman

## References

An Algorithm for Computing the Nonparametric MLE of a Mixing Distribution, Lesperance, Mary L. and Kalbfleisch, John D., JASA, 1992

#### See Also

VEM, EMICM, PGM

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## **Examples**

```
data(cosmesis)
  csub1 <- subset(cosmesis, subset=Trt==0, select=c(L,R))
  ISDM(csub1)
# data(pruitt)
# ISDM(pruitt)</pre>
```

Maclist

A function to

#### **Description**

Returns a list of maximal cliques of the intersection graph of the real valued intervals supplied in m. These are one dimensional intervals with one interval for each individual. The algorithm is coded in interpreted code and should be moved to compiled code for speed. How do we handle exact failure times? Which algorithm is used?

#### Usage

```
Maclist(intvls, Lopen=TRUE, Ropen=FALSE)
```

## **Arguments**

intvls A n by 2 matrix, the first column is the left endpoints and the second column contains the right and points of the failure time intervals.

contains the right endpoints of the failure time intervals.

A boolean indicating whether the intervals are open on the left.

A boolean indicating whether the intervals are open on the right.

#### Value

A list of length m. Each element of the list corresponds to one maximal antichain. The row numbers (from m) identify the individuals and all row numbers for the individuals in the maximal clique. Maximal cliques occur in their natural (left to right) order.

# Author(s)

Alain Vandal and Robert Gentleman

## References

Computational Methods for Censored Data using Intersection Graphs, R. Gentleman and A. Vandal, JCGS, 2000.

#### See Also

Macmat

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## **Examples**

```
data(cosmesis)
csub1 <- subset(cosmesis, subset=Trt==0, select=c(L,R))
ml1 <- Maclist(csub1)</pre>
```

Macmat

A function to compute the incidence matrix for an intersection graph.

# **Description**

Returns the Petrie matrix and Petrie pairs of an interval order given its list of maximal antichains. These can be obtained from Maclist.

# Usage

```
Macmat(ml)
```

## **Arguments**

m1

A list containing the maximal cliques of the intersection graph of the data.

# **Details**

Not worth mentioning?

# Value

A list containing two components.

pmat The Petrie or clique matrix of the underlying interval order.

ppairs The Petrie pairs for each observation. These indicate the first and last maximal

clique occupied by the observation.

## Author(s)

Alain Vandal and Robert Gentleman

#### References

Computational Methods for Censored Data using Intersection Graphs, R. Gentleman and A. Vandal, JCGS, 2000.

#### See Also

Maclist

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## **Examples**

```
data(cosmesis)
csub1 <- subset(cosmesis, subset=Trt==0, select=c(L,R))
ml1 <- Maclist(csub1)
mm1 <- Macmat(ml1)</pre>
```

MLEintvl

Compute the real representation for the maximal cliques.

# **Description**

The intervals on the real line that corresponds to the intersections of the maximal cliques are computed and returned.

#### Usage

```
MLEintvl(intvls, ml=Maclist(intvls))
```

# **Arguments**

intvls The n by 2 matrix containing the event time intervals for the individuals under

study.

ml The Maclist computed for the intvls.

#### Value

An m by 2 matrix, where m is the number of maximal cliques. The first column contains the left end point of the real representation for the appropriate maximal clique and the second column contains the right end point.

## Author(s)

Alain Vandal and Robert Gentleman

#### References

Computational Methods for Censored Data using Intersection Graphs, R. Gentleman and A. Vandal, JCGS, 2000.

#### See Also

Maclist

```
data(cosmesis)
csub1 <- subset(cosmesis, subset=Trt==0, select=c(L,R))
MLEintvl(csub1)</pre>
```

16 PGM

PGM	An implementation of the projected gradient methods for finding the NPMLE.

# **Description**

An estimate of the NPMLE is obtained by using projected gradient methods. This method is a special case of the methods described in Wu (1978).

# Usage

```
PGM(A, pvec, maxiter = 500, tol=1e-07, told=2e-05, tolbis=1e-08, keepiter=FALSE)
```

#### **Arguments**

A A is either the m by n clique matrix or the n by 2 matrix containing the left and

right end points for each event time.

pvec An initial estimate of the probability vector.

maxiter The maximum number of iterations to take.

tol The tolerance for decreases in likelihood.

told told does not seem to be used.

tolbis The tolerance used in the bisection code.

keepiter A boolean indicating whether to return the number of iterations.

#### **Details**

New directions are selected by the projected gradient method. The new optimal pvec is obtained using the bisection algorithm, moving in the selected direction. Convergence requires both the  $L_1$  distance for the improved pvec and the change in likelihood to be below tol.

#### Value

An object of class icsurv containing the following components:

pf The NPMLE of pvec.

sigma The cumulative sum of pvec.

1val The value of the log likelihood at pvec.

clmat The clique matrix.

method The method used, currently only "MPGM" is possible. lastchange The difference between pf and the previous iterate.

numiter The number of iterations carried out.

eps The tolerances used.

converge A boolean indicating whether convergence occurred within maxiter iterations. iter If keepiter is true then this is a matrix containing all iterations - useful for

debugging.

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#### Author(s)

Alain Vandal and Robert Gentleman.

#### References

Some Algorithmic Aspects of the Theory of Optimal Designs, C.-F. Wu, 1978, Annals.

# See Also

```
VEM, ISDM, EMICM, PGM, EM
```

# **Examples**

```
data(cosmesis)
csub1 <- subset(cosmesis, subset=Trt==0, select=c(L,R))
PGM(csub1)
data(pruitt)
PGM(pruitt)</pre>
```

plot.icsurv

A plot method for the estimates produced by the estimation methods in Icens.

# Description

Produces nice plots of the estimated NPMLE.

# Usage

```
## S3 method for class 'icsurv'
plot(x, type="eq", surv=FALSE, bounds=FALSE, shade=3, density=30,
angle=45, lty=1, new=TRUE, xlab="Time", ylab="Probability", main="GMLE",
ltybnds=2, ...)
```

# Arguments

X	The estimate of the NPMLE.
type	Three options, "eq" for equivalence call, "gw" for the Groeneboom-Wellner estimate, and "lc" for the left-continuous estimate.
surv	Logical indicating whether or not to plot the survival curve.
bounds	Logical indicating whether or not to include bounds around the estimate.
shade	An integer in 1, 2, or 3 denoting what component of the plot to shade.
density	The density of shading lines, in lines per inch.
angle	The slope of shading lines, given as an angle in degrees (counter-clockwise).
lty	The line type for the estimates.

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new	Logical indicating whether or not to create a new plot.
xlab	The x-axis label.
ylab	The y-axis label.
main	The main title for the plot.
ltybnds	The line type for the bounds on the estimates.
	Additional arguments passed to the plot function.

#### Value

No value is returned. A plot of the NPMLE is made on the active graphics device.

#### Author(s)

Alain Vandal and Robert Gentleman.

# See Also

```
VEM, ISDM, EMICM, PGM
```

# **Examples**

```
data(cosmesis)
csub1 <- subset(cosmesis, subset=Trt==0, select=c(L,R))
e1 <- VEM(csub1)
par(mfrow=c(2,2))
plot(e1)
data(pruitt)
e2 <- EM(csub1)
plot(e2)
e3 <- PGM(csub1)
plot(e3)
e4 <- EMICM(csub1)
plot(e4)</pre>
```

Plotboxes

Plot the event time regions for bivariate data.

# **Description**

Plot rectangles described by the interval given in the first two arguments.

# Usage

```
Plotboxes(int1, int2, textp=FALSE, showmac=FALSE, showsupp=FALSE, showmp=FALSE, cliques=NULL, macprod=NULL, density=c(2, 8, 20), col=c(2, 3, 4), offsetx=0.02, offsety=0.03)
```

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# Arguments

int1 The intervals for the x dimension.int2 The intervals for the y dimension.

textp Boolen, if true add text.

showmac Boolean, if true then the maximal cliques are shown in a different color?

showsupp Boolean, if true show support boxes.

showmp Boolean

cliques Maximal cliques.

macprod macprod

density The density of the polygon shading lines, in lines per inch.

col Color for plotting features.

offsety Offset for x-axis.

offsety Offset for y-axis.

#### Value

No value is returned. The event rectangles are plotted on the active graphics device.

#### Author(s)

A. Vandal and R. Gentleman

# References

*Graph–Theoretical Aspects of Bivariate Censored Data*, R. Gentleman and A. Vandal, 1999, submitted.

## See Also

```
BVclmat, BVsupport, BVcliques
```

```
data(cmv)
Plotboxes(cmv[,1:2], cmv[,3:4], showmac=TRUE)
```

20 PMGA

PMGA

Implement the pool monotone groups algorithm.

# **Description**

For isotonization problems some increase in speed and decrease in complexity can be achieved through the use of the pool monotone groups algorithm of Y.L. Zhang and M.A. Newton (1997). It isotonizes a weighted and ordered set of values.

# Usage

```
PMGA(est, ww=rep(1, length(est)))
```

#### **Arguments**

est The vector of values, in the appropriate order.

ww The weight vector.

#### **Details**

To be supplied at some later date.

#### Value

An object containing the following components:

est The isotonized estimates.

ww The weights associated with the isotonized estimates.

poolnum The number of values pooled in the current estimate.

passes The number of passes which were required to isotonize the list.

## Author(s)

Alain Vandal and Robert Gentleman.

#### References

Y.L. Zhang and M.A. Newton (1997), http://www.stat.wisc.edu/~newton/newton.html)

#### See Also

**EMICM** 

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pruitt	A small artificial, bivariate right-censored data set.	
pruitt	A small artificial, bivariate right-censored data set.	

# Description

The pruitt data was given in Pruitt (1993) as an example for testing different methods of estimating the bivariate NPMLE for right censored data. This matrix represents the clique matrix of the intersection graph of the data set given by Pruitt.

# **Format**

This data frame contains 8 columns, labeled A through H that represent the observations. There are seven rows corresponding to the seven maximal cliques in the intersection graph.

#### Source

Small Sample Comparison of Six Bivariate Survival Curve Estimators, Journal of Statistical Computation and Simulation, R. Pruitt, 1993.

## **Examples**

data(pruitt)

VEM

Compute the NPMLE of p via the Vertex Exchange Method.

#### **Description**

The Vertex Exchange Method is used to obtain the NPMLE of p.

# Usage

```
VEM(A, pvec, maxiter=500, tol=1e-07, tolbis=1e-07, keepiter=FALSE)
```

#### **Arguments**

A	The m by n	incidence	matrix	or the n t	y 2	matrix of ii	ntervals.
---	------------	-----------	--------	------------	-----	--------------	-----------

pvec The initial estimate for the probability vector.

maxiter The maximum number of iterations allowed.

tol The tolerance used to determine convergence.

tolbis The tolerance used in the bisection stage of the algorithm. keepiter Should iteration information be retained and returned.

VEM VEM

# **Details**

Lots.

#### Value

An object of class icsurv with the following components.

pf The NPMLE of the probability vector.

numiter The number of iterations used.

1val The value of the logarithm of the likelihood at the NPMLE.

converge Boolean stating whether the iteration converged.

intmap If present it contains the real representations for the maximal cliques. These are

the intervals (on the real line) where the mass in pf is placed.

# Author(s)

Robert Gentleman and Alain Vandal

#### References

A Vertex-exchange-method in \$D\$-optimal Design Theory, D. Bohning, Metrika, 1986.

#### See Also

```
EM, ISDM, EMICM, PGM
```

```
data(cosmesis)
csub1 <- subset(cosmesis, subset=Trt==0, select=c(L,R))
VEM(csub1)
data(pruitt)
VEM(pruitt)</pre>
```

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