Package ‘Benchmarking’

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Title Benchmark and Frontier Analysis Using DEA and SFA
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Description Methods for frontier analysis, Data Envelopment Analysis (DEA), under different technology assumptions (fdh, vrs, drs, crs, irs, add/frh, and fdh+), and using different efficiency measures (input based, output based, hyperbolic graph, additive, super, and directional efficiency). Peers and slacks are available, partial price information can be included, and optimal cost, revenue and profit can be calculated. Evaluation of mergers is also supported. Methods for graphing the technology sets are also included. There is also support comparative methods based on Stochastic Frontier Analyses (SFA). In general, the methods can be used to solve not only standard models, but also many other model variants. It complements the book, Bogetoft and Otto, Benchmarking with DEA, SFA, and R, Springer-Verlag, 2011, but can of course also be used as a stand-alone package.
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The Benchmarking package contains methods to estimate technologies and measure efficiencies using DEA and SFA. Data Envelopment Analysis (DEA) are supported under different technology assumptions (fdh, vrs, drs, crs, irs, add), and using different efficiency measures (input based, output based, hyperbolic graph, additive, super, directional). Peers are available, partial price information can be included, and optimal cost, revenue and profit can be calculated. Evaluation of mergers are also supported. A comparative method for estimating stochastic frontier function (SFA) efficiencies is included. The methods can solve not only standard models, but also many other model variants, and they can be modified to solve new models.
The package also support simple plots of DEA technologies with two goods; either as a transformation curve (2 outputs), an isoquant (2 inputs), or a production function (1 input and 1 output). When more inputs and outputs are available they are aggregated using weights (prices, relative prices).

The package complements the book, Bogetoft and Otto, *Benchmarking with DEA, SFA, and R*, Springer-Verlag 2011, but can of course also be used as a stand-alone package.

Details

| Package: | Benchmarking |
| Type:    | Package      |
| Version: | 0.27 ($Revision: 157 $) |
| Date:    | $Date: 2015-07-09 11:40:20 +0200 (to, 09 jul 2015) $ |
| License: | Copyright    |

dea | DEA input or output efficiency measures, peers, lambdas and slacks |
dea.dual | Dual weights (prices), including restrictions on weights |
dea.direct | Directional efficiency |
sdea | Super efficiency. |
dea.add | Additive efficiency; sum of slacks in DEA technology. |
mea | Multidirectional efficiency analysis or potential improvements, |
eff | Efficiency from an object returned from any of the dea or sfa functions. |
slack | Slacks in DEA models |
excess | Calculates excess input or output compared to DEA frontier. |
peers | get the peers for each firm. |
dea.boot | Bootstrap DEA models |
cost.opt | Optimal input for given output and prices. |
revenue.opt | Optimal output for given input and prices. |
profit.opt | Optimal input and output for given input and output prices. |
dea.plot | Graphs of DEA technologies under alternative technology assumptions. |
dea.plot.frontier | Specialized for 1 input and 1 output. |
dea.plot.isoquant | Specialized for 2 inputs. |
dea.plot.transform | Specialized for 2 outputs. |
eladder | Efficiency ladder for a single firm. |
eladder.plot | Plot efficiency ladder for a single firm. |
make.merge | Make an aggregation matrix to perform mergers. |
dea.merge | Decompose efficiency from a merger of firms |
sfa | Stochastic frontier analysis, production, distance, and cost function (SFA) |
outlier.ap | Detection of outliers |
eff.dens | Estimate and plot kernel density of efficiencies |
critValue | Critical values calculated from bootstrap DEA models. |
typeIerror | Probability of a type I error for a test in bootstrap DEA models. |
Note

The interface for the methods are very much like the interface to the methods in the package FEAR (Wilson 2008). One change is that the data now are transposed to reflect how data is usually available in applications, i.e. we have firms on rows, and inputs and output in the columns. Also, the argument for the options RTS and ORIENTATION can be given as memotechnical strings, and there are more options to control output.

The input and output matrices can contain negative numbers, and the methods can thereby manage restricted or fixed input or output.

The return is not just the efficiency, but also slacks, dual values (shadow prices), peers, and lambdas (weights).

Author(s)

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References

Bogetoft and Otto; Benchmarking with DEA, SFA, and R; Springer 2011

Examples

# Plot of different technologies
x <- matrix(c(100,200,300,500),ncol=1,dimnames=list(LETTERS[1:4],"x"))
y <- matrix(c(75,100,300,400),ncol=1,dimnames=list(LETTERS[1:4],"y"))
dea.plot(x,y,RTS="vrs",ORIENTATION="in-out",txt=rownames(x))
dea.plot(x,y,RTS="drs",ORIENTATION="in-out",add=TRUE,lty="dashed",lwd=2)
dea.plot(x,y,RTS="crs",ORIENTATION="in-out",add=TRUE,lty="dotted")
dea.plot(x,y,RTS="fdh",ORIENTATION="in-out",txt=rownames(x),main="fdh")
dea.plot(x,y,RTS="irs",ORIENTATION="in-out",txt=TRUE,main="irs")
dea.plot(x,y,RTS="irs2",ORIENTATION="in-out",txt=rownames(x),main="irs2")
dea.plot(x,y,RTS="add",ORIENTATION="in-out",txt=rownames(x),main="add")

# A quick frontier with 1 input and 1 output
dea.plot(x,y, main="Basic plot of frontier")

de(x,y, RTS="vrs", ORIENTATION="in")
e <- dea(x,y, RTS="vrs", ORIENTATION="in")
e
eff(e)
peers(e)
peers(e, NAMES=TRUE)
print(peers(e, NAMES=TRUE), quote=FALSE)
lambda(e)
summary(e)
# Calculating super efficiency
esuper <- sdea(x,y, RTS="vrs", ORIENTATION="in")
esuper
print(peers(esuper,NAMES=TRUE),quote=FALSE)

# Technology for super efficiency for firm number 3/C
# Note that drop=FALSE is necessary for XREF and YREF to be matrices
# when one of the dimensions is or is reduced to 1.
e3 <- dea(x,y, XREF=x[-3,,drop=FALSE], YREF=y[-3,,drop=FALSE])
dea.plot(x[-3],y[-3],RTS="vrs",ORIENTATION="in-out",txt=LETTERS[c(1,2,4)])
points(x[3],y[3],cex=2)
text(x[3],y[3],LETTERS[3],adj=c(-.75,.75))
e3 <- dea(x,y, XREF=x[-3,,drop=FALSE], YREF=y[-3,,drop=FALSE])
eff(e3)
peers(e3)
print(peers(e3,NAMES=TRUE),quote=FALSE)
lambda(e3)
e3$lambda

# Taking care of slacks
x <- matrix(c(100,200,300,500,100,600),ncol=1,
dimnames=list(LETTERS[1:6],"x"))
y <- matrix(c(75,100,300,400,50,400),ncol=1,
dimnames=list(LETTERS[1:6],"y"))

# Phase one, calculate efficiency
e <- dea(x,y)
print(e)
peers(e)
lambda(e)

# Phase two, calculate slacks (maximize sum of slacks)
sl <- slack(x,y,e)
data.frame(sl$sx,sl$sy)
peers(sl)
lambda(sl)
sl$lambda
summary(sl)

# The two phases in one function call
e2 <- dea(x,y,SLACK=TRUE)
print(e2)
data.frame(eff(e2),e2$slack,e2$sx,e2$sy,lambda(e2))
peers(e2)
lambda(e2)
e2$lambda
Description
The data set is from an US federally sponsored program for providing remedial assistance to disad-
vantaged primary school students. The firms are 70 school sites, and data are from entire sites. The
variables consists of results from three different kind of tests, a reading score, $y_1$, a math score, $y_2$,
and a self–esteem score, $y_3$, which are considered outputs in the model, and five different variables
considered to be inputs, the education level of the mother, $x_1$, the highest occupation of a family
member, $x_2$, parental visits to school, $x_3$, time spent with children in school-related topics, $x_4$, and
the number of teachers at the site, $x_5$.

Usage
data(charnes1981)

Format
A data frame with 70 school sites with the following variables.

- firm: school site number
- $x_1$: education level of the mother
- $x_2$: highest occupation of a family member
- $x_3$: parental visits to school
- $x_4$: time spent with children in school-related topics
- $x_5$: the number of teachers at the site
- $y_1$: reading score
- $y_2$: math score
- $y_3$: self–esteem score
- pft: =1 if in program (program follow through) and =0 if not in program
- name: Site name

Details
The command data(charnes1981) will create a data frame named charnes1981 with the above
data.
Beside input and output varianles there is further information in the data set, that the first 50 school
sites followed the program and that the last 20 are the results for sites not following the program.
This is showed by the variable pft.

Note
Data as .csv are loaded by the command data using read.table(...,header=TRUE,sep=";")
such that this file is a semicolon separated file and not a comma separated file.
Therefore, to read the file from a script the command must be read.csv("charnes1981.csv",sep=";")
or read.csv2("charnes1981.csv").
Thus the data can be read either as charnes1981 <-
read.csv2(paste(.Library,"Benchmarking/data","charnes1981.csv",sep="/"))
or as data(charnes1981) if the package **Benchmarking** is loaded. In both cases the data will be
in the data frame charnes1981.
Source


Examples

data(charnes1981)
x <- with(charnes1981, cbind(x1,x2,x3,x4,x5))
y <- with(charnes1981, cbind(y1,y2,y3))

# Farrell input efficiency; vrs technology
e <- dea(x,y)
# The number of times each peer is a peer
np <- get.number.peers(e)
# Peers that are peers for more than 20 schools, and the number of
# times they are peers
np[which(np[,2]>20),]

# Plot first input against first output and emphasize the peers that
# are peers for more than 20 schools in the model with five inputs and
# three outputs
inp <- np[which(np[,2]>20),1]
dea.plot(x[,1],y[,1])
points(x[inp,1], y[inp,1], pch=16, col="red")

cost.opt

*DEA optimal cost, revenue, and profit*

Description

Estimates the input and/or output vector(s) that minimize cost, maximize revenue or maximize profit in the context of a DEA technology

Usage

cost.opt(XREF, YREF, W, YOBS=NULL, RTS="vrs", param=NULL, TRANSPOSE=FALSE, LP=FALSE, LPK = NULL)

revenue.opt(XREF, YREF, P, XOBS=NULL, RTS="vrs", param=NULL, TRANSPOSE = FALSE, LP = FALSE, LPK = NULL)

profit.opt(XREF, YREF, W, P, RTS = "vrs", param=NULL, TRANSPOSE = FALSE, LP = FALSE, LPK = NULL)
Arguments

Input and output matrices are in the same form as for the method dea.

XREF
Input of the firms defining the technology, a K x m matrix of observations of K firms with m inputs (firm x input). In case TRANSPOSE=TRUE the input matrix is transposed as input x firm.

YREF
output of the firms defining the technology, a K x n matrix of observations of K firms with n outputs (firm x input). In case TRANSPOSE=TRUE the output matrix is transposed as output x firm.

W
Input prices as a matrix. Either same prices for all firms or individual prices for all firms; i.e. either a 1 x m or a K x m matrix for K firms and m inputs

P
Output prices as a matrix. Either same prices for all firms or individual prices for all firms; i.e. either a 1 x n or K x n matrix for K firms and n outputs

XOBS
The input for which an optimal, revenue maximizing, output vector is to be calculated. Defaults is XREF. Same form as XREF

YOBS
The output for which an optimal, cost minimizing input vector is to be calculated. Defaults is YREF. Same form as YREF

RTS
A text string or a number defining the underlying DEA technology / returns to scale assumption.

0 fdh Free disposability hull, no convexity assumption
1 vrs Variable returns to scale, convexity and free disposability
2 drs Decreasing returns to scale, convexity, down-scaling and free disposability
3 crs Constant returns to scale, convexity and free disposability
4 irs Increasing returns to scale, (up-scaling, but not down-scaling), convexity and free disposability
5 add Additivity (scaling up and down, but only with integers), and free disposability
6 fdh+ A combination of free disposability and restricted or local constant return to scale

param
Possible parameters. At the moment only used for RTS=“fdh+” to set low and high values for restrictions on lambda; see the section details and examples in dea for its use. Future versions might also use param for other purposes.

TRANSPOSE
Input and output matrices are treated as firms times goods for the default value TRANSPOSE=FALSE corresponding to the standard in R for statistical models. When TRUE data matrices, quantities and prices, are transposed to goods times firms matrices.

LP
Only for debugging. If LP=TRUE then input and output for the LP program are written to standard output for each unit.

LPK
When LPK=k then a mps file is written for firm k; it can be used as input to an alternative LP solver to check the results.

Details

The LP optimization problem is formulated in Bogetoft and Otto (2011, pp 35 and 102) and is solved by the LP method in the package lpSolveAPI.

The methods print and summary are working for cost.opt, revenue.opt, and profit.opt
The values returned are the optimal input, and/or optimal output. When saved in an object the following components are available:

- **xopt**: The optimal input, returned as a matrix by `cost.opt` and `profit.cost`.
- **yopt**: The optimal output, returned as a matrix by `revenue.opt` and `profit.cost`.
- **cost**: The optimal/minimal cost.
- **revenue**: The optimal/maximal revenue
- **profit**: The optimal/maximal profit
- **lambda**: The peer weights that determines the technology, a matrix. Each row is the lambdas for the firm corresponding to that row; for the vrs technology the rows sum to 1. A column shows for a given firm how other firms are compared to this firm; i.e. peers are firms with a positive element in their columns.

Note

The index for peer units can be returned by the method `peers` and the weights are returned in `lambda`. Note that the peers now are the firms for the optimal input and/or output allocation, not just the technical efficient firms.

Author(s)

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References

Bogetoft and Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011

See Also


Examples

```r
x <- matrix(c(2,12, 2,8, 5,5, 10,4, 10,6, 3,13), ncol=2, byrow=TRUE)
y <- matrix(1,nrow=dim(x)[1],ncol=1)
w <- matrix(c(1.5, 1),ncol=2)

txt <- LETTERS[1:dim(x)[1]]
dea.plot(x[,1],x[,2], ORIENTATION="in", cex=1.25)
text(x[,1],x[,2],txt,adj=c(-.7,-.2),cex=1.25)

# technical efficiency
te <- dea(x,y,RTS="vrs")
xopt <- cost.opt(x,y,w,RTS=1)
cobs <- x %*% t(w)
copt <- xopt$x %*% t(w)
```
# cost efficiency
ce <- copt/cobs
# allocativce efficiency
ae <- ce/te$eff
data.frame("ce"=ce,"te"=te$eff,"ae"=ae)
print(cbind("ce"=c(ce),"te"=te$eff,"ae"=c(ae)),digits=2)

# isocost line in the technology plot
abline(a=copt[1]/w[2], b=-w[1]/w[2], lty="dashed")

critValue

Description

Calculates critical value for test using bootstrap output in DEA models

Usage

critValue(s, alpha=0.05)

Arguments

s Vector with calculated values of the statistic for each of the NREP bootstraps;
NREP is from boot.sw98

alpha The size of the test

Details

Needs bootstrapped values of the test statistic

Value

Returns the critical value

Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

See Also


Examples

# The critical value for two-sided test in normal distribution found
# by simulation.
x <- rnorm(1000000)
critValue(x,.975)
DEA efficiency

Description
Estimates a DEA frontier and calculates efficiency measures a la Farrell.

Usage
dea(X, Y, RTS="vrs", ORIENTATION="in", XREF=NULL, YREF=NULL, FRONT.IDX=NULL, SLACK=FALSE, DUAL=FALSE, DIRECT=NULL, param=NULL, TRANSPOSE=FALSE, FAST=FALSE, LP=FALSE, CONTROL=NULL, LPK=NULL)
## S3 method for class 'Farrell'
print(x, digits=4, ...)
## S3 method for class 'Farrell'
summary(object, digits=4, ...)

Arguments
X Inputs of firms to be evaluated, a K x m matrix of observations of K firms with m inputs (firm x input). In case TRANSPOSE=TRUE the input matrix is transposed to input x firm.
Y Outputs of firms to be evaluated, a K x n matrix of observations of K firms with n outputs (firm x input). In case TRANSPOSE=TRUE the output matrix is transposed to output x firm.
RTS Text string or a number defining the underlying DEA technology / returns to scale assumption.
0 fdh Free disposability hull, no convexity assumption
1 vrs Variable returns to scale, convexity and free disposability
2 drs Decreasing returns to scale, convexity, down-scaling and free disposability
3 crs Constant returns to scale, convexity and free disposability
4 irs Increasing returns to scale, (up-scaling, but not down-scaling), convexity and free disposability
5 irs2 Increasing returns to scale (up-scaling, but not down-scaling), additivity, and free disposability
6 add Additivity (scaling up and down, but only with integers), and free disposability; also known as replicability and free disposability
7 fdh+ A combination of free disposability and restricted or local constant return to scale
10 vrs+ As vrs, but with restrictions on the individual lambdas via param
ORIENTATION Input efficiency "in" (1), output efficiency "out" (2), and graph efficiency "graph" (3). For use with DIRECT, an additional option is "in-out" (0).
XREF Inputs of the firms determining the technology, defaults to X
YREF Outputs of the firms determining the technology, defaults to Y
FRONT.IDX Index for firms determining the technology
SLACK Calculate slack in a phase II calculation by an intern call of the function slack. Note that the precision for calculating slacks for orientation graph is low.
DUAL Calculate dual variables, i.e. shadow prices; not calculated for orientation graph as that is not an LP problem.

DIRECT Directional efficiency, DIRECT is either a scalar, an array, or a matrix with non-negative elements.

If the argument is a scalar, the direction is \((1,1,...,1)\) times the scalar; the value of the efficiency depends on the scalar as well as on the unit of measurements.

If the argument is an array, this is used for the direction for every firm; the length of the array must correspond to the number of inputs and/or outputs depending on the ORIENTATION.

If the argument is a matrix then different directions are used for each firm. The dimensions depend on the ORIENTATION (and TRANSPOSE), the number of firms must correspond to the number of firms in \(X\) and \(Y\).

DIRECT must not be used in connection with DIRECTION="graph".

param Possible parameters. At the moment only used for RTS="fdh+" to set low and high values for restrictions on lambda; see the section details and examples for its use. Future versions might also use param for other purposes.

TRANSPOSE Input and output matrices are treated as firms times goods matrices for the default value TRANSPOSE=FALSE corresponding to the standard in R for statistical models. When TRUE data matrices are transposed to good times firms matrices as is normally used in LP formulation of the problem.

LP Only for debugging. If LP=TRUE then input and output for the LP program are written to standard output for each unit.

FAST Only calculate efficiencies and just return them as a vector, i.e. no lambda or other output. The return when using FAST cannot be used as input for slack and peers.

CONTROL Possible controls to lpSolveAPI, see the documentation for that package.

... Optional parameters for the print and summary methods.

object, x An object of class Farrell (returned by the function dea) – R code uses ‘object’ and ‘x’ alternating for generic methods.

digits digits in printed output, handled by format in print.

LPK when LPK=k then a mps file is written for firm k; it can be used as input to an alternative LP solver to check the results.

Details

The return from dea and sdea is an object of class Farrell. The efficiency in dea is calculated by the LP method in the package lpSolveAPI. Slacks can be calculated either in the call of dea using the option SLACK=TRUE or in a following call to the function slack.

The directional efficiency when the argument DIRECT is used, depends on the unit of measurement and is not restricted to be less than 1 (or greater than 1 for output efficiency) and is therefore completely different from the Farrell efficiency.

The crs factor in RTS="fdh+" that sets the lower and upper bound can be changed by the argument param that will set the lower and upper bound to 1-param and 1+param; the default value is param=.15. The value must be greater than or equal to 0 and strictly less than 1. A value of 0 corresponds to RTS="fdh". To get an asymmetric interval set param to a 2 dimensionial array with values...
for the low and high end for interval, for instance \texttt{param=c(.8,1.15)}. The FDH+ technology set is described in Bogetoft and Otto (2011) pages 73–74.

The technology RTS="vrs+" uses the parameter \texttt{param} to set restrictions on lambda, the convexity parameters. The elements of \texttt{param} are \texttt{param=(low,high,sum_low,sum_high)} where "low" and "high" are restrictions on the individual lambda and "sum_low" and "sum_high" are restrictions on the sum of lambdas. The individual lambda must be in the interval from low to high or be zero. With one parameter the restrictions set are \texttt{(param,1+1-(param),1,1)}, with two parameters \texttt{(param[1],param[2],1,1)}, and with four parameters \texttt{(param[1],param[2],param[3],param[4])}. The resulting technology set is not necessarily convex.

The graph orientated efficiency is calculated by bisection between feasible and infeasible values of \textit{G}. The precision in the result is less than for the other orientations.

When the argument \texttt{DIRECT=d} is used then the returned value \textit{e} for input orientation is the excess input measured in \textit{d} units of measurements, i.e. \texttt{x - ed}, and for output orientation \texttt{y + ed}. The directional efficiency can be restricted to inputs (\texttt{ORIENTATION="in"}), restricted to outputs (\texttt{ORIENTATION="out"}), or both include inputs and output directions (\texttt{ORIENTATION="in-out"}). Directional efficiency is discussed on pages 31–35 and 121–127 in Bogetoft and Otto (2011).

\textbf{Value}

The results are returned in a Farrell object with the following components. The last three components in the list are only part of the object when \texttt{SLACK=TRUE}.

- \texttt{eff} The efficiencies. Note when \texttt{DIRECT} is used then the efficiencies are not Farrell efficiencies but rather excess values in \texttt{DIRECT} units of measurement
- \texttt{lambda} The lambdas, i.e. the weight of the peers, for each firm
- \texttt{objval} The objective value as returned from the LP program; normally the same as \texttt{eff}, but for \texttt{slack} it is the the sum of the slacks.
- \texttt{RTS} The return to scale assumption as in the option \texttt{RTS} in the call
- \texttt{ORIENTATION} The efficiency orientation as in the call
- \texttt{TRANSPOSE} As in the call
- \texttt{slack} A logical vector where the component for a firm is \texttt{TRUE} if the sums of slacks for the corresponding firm is positive. Only calculated in \texttt{dea} when option \texttt{SLACK=TRUE}
- \texttt{sum} A vector with sums of the slacks for each firm. Only calculated in \texttt{dea} when option \texttt{SLACK=TRUE}
- \texttt{sx} A matrix for input slacks for each firm, only calculated if the option \texttt{SLACK} is \texttt{TRUE} or returned from the method \texttt{slack}
- \texttt{sy} A matrix for output slack, see \texttt{sx}
- \texttt{ux} Dual variable for input, only calculated if \texttt{DUAL} is \texttt{TRUE}
- \texttt{vy} Dual variable for output, only calculated if \texttt{DUAL} is \texttt{TRUE}. 
**Note**

The arguments `X`, `Y`, `XREF`, and `YREF` are supposed to be matrices or numerical data frames that in the function will be converted to matrices. When subsetting a matrix or data frame to just one column then the class of the resulting object/variable is no longer a matrix or a data frame, but just a numeric (array, vector). Therefore, in this case a numeric input that is not a matrix nor a data frame is transformed to a 1 column matrix, and here the use of the argument `TRANSPOSE=TRUE` gives an error.

The dual values are not unique for extreme points (firms on the boundary with an efficiency of 1) and therefore the calculated dual values for these firms can depend on the order of firms in the reference technology. The same lack of uniqueness also make the peers for some firms depend on the order of firms in the reference technology.

To calculate slack use the argument `SLACK=TRUE` or use the function `slack` directly.

When there is slack, and slack is not taken into consideration, then the peers for a firm with slack might depend on the order of firms in the data set; this is a property of the LP algorithm used to solve the problem.

To handle fixed, non-discretionary inputs, one can let it appear as negative output in an input-based mode, and inversely for fixed, non-discretionary outputs. Fixed inputs (outputs) can also be handled by directional efficiency; set the direction, the argument `DIRECT`, equal to the variable, discretionary inputs (outputs) and 0 for the fixed inputs (outputs).

When the the argument `DIRECT=X` is used the then the returned efficiency is equal to 1 minus the Farrell efficiency for input orientation and to the Farrell efficiency minus 1 for output orientation.

To use matrices `X` and `Y` prepared for the methods in the package `FEAR` (Wilson 2008) set the options `TRANSPOSE=TRUE`; for consistency with `FEAR` the options `RTS` and `ORIENTATION` also accepts numbers as in `FEAR`.

The tolerance that lambda is zero or one is 1e-7, the default value of 'epsint' in the package `lpSolveAPI`, i.e. values closer than 1e-7 from zero or one are set to respective integer value. The 'epsint' is the tolerance that is used to determine whether a floating-point number is in fact an integer. The same tolerance is used for efficiency value near one.

**Author(s)**

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**References**

Peter Bogetoft and Lars Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011

**See Also**


**Examples**

```r
x <- matrix(c(100,200,300,500,100,200,600),ncol=1)
y <- matrix(c(75,100,300,400,25,50,400),ncol=1)
dea.plot.frontier(x,y,txt=TRUE)
```


```r
e <- dea(x, y)
eff(e)
print(e)
summary(e)
lambda(e)

# Input savings potential for each firm
(1-eff(e)) * x
(1-e$eff) * x

# calculate slacks
el <- dea(x, y, SLACK=TRUE)
data.frame(e$eff, el$eff, el$slack, el$sx, el$sy)

# Fully efficient units, eff==1 and no slack
which(eff(e) == 1 & !el$slack)

# fdh+ with limits in the interval [.7, 1.2]
dea(x, y, RTS="fdh+", param=c(.7, 1.2))
```

---

**dea.add**

*Additive DEA model*

**Description**

Calculates additive efficiency as sum of input and output slacks within different DEA models

**Usage**

```r
dea.add(X, Y, RTS="vrs", XREF=NULL, YREF=NULL,
FRONT.IDX=NULL, param=NULL, TRANSPOSE=FALSE, LP=FALSE)
```

**Arguments**

- **X**
  - Inputs of firms to be evaluated, a K x m matrix of observations of K firms with m inputs (firm x input). In case TRANSPOSE=TRUE the input matrix is transposed to input x firm.

- **Y**
  - Outputs of firms to be evaluated, a K x n matrix of observations of K firms with n outputs (firm x input). In case TRANSPOSE=TRUE the output matrix is transposed to output x firm.

- **RTS**
  - Text string or a number defining the underlying DEA technology / returns to scale assumption.

- **0**
  - `fdh` Free disposability hull, no convexity assumption

- **1**
  - `vrs` Variable returns to scale, convexity and free disposability
2  drs  Decreasing returns to scale, convexity, down-scaling and free disposability
3  crs  Constant returns to scale, convexity and free disposability
4  irs  Increasing returns to scale, (up-scaling, but not down-scaling), convexity and free disposability
5  add  Additivity (scaling up and down, but only with integers), and free disposability

XREF  Inputs of the firms determining the technology, defaults to X
YREF  Outputs of the firms determining the technology, defaults to Y
FRONT_IDX  Index for firms determining the technology
param  Possible parameters. At the moment only used for RTS="fdh+" to set low and high values for restrictions on lambda; see the section details and examples for its use. Future versions might also use param for other purposes.
TRANSPOSE  Input and output matrices are treated as firms times goods matrices for the default value TRANSPOSE=FALSE corresponding to the standard in R for statistical models. When TRUE data matrices are transposed to good times firms matrices as is normally used in LP formulation of the problem.
LP  Only for debugging. If LP=TRUE then input and output for the LP program are written to standard output for each unit.

Details

The sum of the slacks is maximized in a LP formulation of the DEA technology. The sum of the slacks can be seen as distance to the frontier when you only move parallel to the axes of inputs and outputs, i.e. not a usual Euclidean distance, but what is also known as an L1 norm.

Since it is the sum of slacks that is calculated, there is no exogenous ORIENTATION in the problem. Rather, there is generally both an input and an output direction in the slacks. The model considers the input excess and output shortfall simultaneously and finds a point on the frontier that is most distant to the point being evaluated.

Value

sum  Sum of all slacks for each firm, sum=sum(sx)+sum(sy).
slack  A non-NULL vector of logical variables, TRUE if there is slack for the corresponding firm, and FALSE if the there is no slack, i.e. the sum of slacks is zero.
sx  A matrix of input slacks for each firm
sy  A matrix of output slack for each firm
lambda  The lambdas, i.e. the weights of the peers for each firm

Note

This is neither a Farrell nor a Shephard like efficiency.

The value of the slacks depends on the scaling of the different inputs and outputs. Therefore the values are not independent of how the input and output are measured.

Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>
Source

Corresponds to Eqs. 4.34-4.38 in Cooper et al. (2007)

References

Bogetoft and Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011

Examples

```r
x <- matrix(c(2,3,2,4,6,5,6,8),ncol=1)
y <- matrix(c(1,3,2,3,5,2,3,5),ncol=1)
dea.plot.frontier(x,y,txt=1:dim(x)[1])

sb <- dea.add(x,y,RTS="vrs")
data.frame("sx"=sb$sx,"sy"=sb$sy,"sum"=sb$sum,"slack"=sb$slack)
```

de.dea.boot

*Bootstrap DEA models*

Description

The function `dea.boot` bootstrap DEA models and returns bootstrap of Farrell efficiencies. This function is slower than the `boot.sw89` from the package `FEAR`. The faster function `boot.fear` is a wrapper for `boot.sw89` from the package `FEAR` returning results directly as Farrell measures.

Usage

```r
dea.boot(X, Y, NREP = 200, EFF = NULL, RTS = "vrs", ORIENTATION="in",
alpha = 0.05, XREF = NULL, YREF = NULL, FRONT.IDX=NULL, EREF = NULL,
DIRECT = NULL, TRANSPOSE = FALSE, SHEPHARD.INPUT = TRUE, LP)

boot.fear(X, Y, NREP = 200, EFF = NULL, RTS = "vrs", ORIENTATION = "in",
alpha = 0.05, XREF = NULL, YREF = NULL, EREF = NULL)
```

Arguments

- **X**  
  Inputs of firms to be evaluated, a K x m matrix of observations of K firms with m inputs (firm x input)
- **Y**  
  Outputs of firms to be evaluated, a K x n matrix of observations of K firms with n outputs (firm x input).
- **NREP**  
  Number of bootstrap replicats
- **EFF**  
  Efficiencies for (X,Y) relative to the technology generated from (XREF,YREF).
- **RTS**  
  The returns to scale assumptions as in `dea`, only works for "vrs", "drs", and "crs"; more to come.
Input efficiency "in" (1), output efficiency "out" (2), and graph efficiency "graph" (3).

One minus the size of the confidence interval for the bias corrected efficiencies

Inputs of the firms determining the technology, defaults to \( X \).

Outputs of the firms determining the technology, defaults to \( Y \).

Index for firms determining the technology.

Efficiencies for the firms in XREF, YREF.

Does not yet work and is therefore not used.

Input and output matrices are \( K \times m \) and \( K \times n \) for the default value \( \text{TRANSPOSE} = \text{FALSE} \); this is standard in R for statistical models. When \( \text{TRANSPOSE} = \text{TRUE} \) data matrices are \( m \times K \) and \( n \times K \).

The bootstrap of the Farrell input efficiencies is done as a Shephard input distance function, the inverse Farrell input efficiency. The option is only relevant for input and graph oriented directions.

Only for debugging purposes.


The bootstrap at the moment does not work for any kind of directional efficiency.

The returned confidence intervals are for the bias corrected efficiencies; to get confidence intervals for the uncorrected efficiencies add the biases to both upper and lower values for the intervals.

Under the default option \( \text{SHEPHARD\_INPUT=TRUE} \) bias and bias corrected efficiencies are calculated for Shephard input distance function and then transformed to Farrell input efficiencies to avoid possible negative biased corrected input efficiencies. If this is not wanted use the option \( \text{SHEPHARD\_INPUT=FALSE} \), This option is only relevant for input and graph oriented directions.

The returned values from both functions are as follows:

- **eff**: Efficiencies
- **eff.bc**: Bias-corrected efficiencies
- **bias**: An array of bootstrap bias estimates for the K firms
- **conf.int**: \( K \times 2 \) matrix with confidence interval for the estimated efficiencies
- **var**: An array of bootstrap variance estimates for the K firms
- **boot**: The replica bootstrap estimates of the Farrell efficiencies, a \( K \times \text{NREP} \) matrix
**Note**

The function `dea.boot` does not depend on the FEAR package and can therefore be used on computers where the package FEAR is not available. This, however, comes with a time penalty as it takes around 4 times longer to run compared to using FEAR directly.

The returned bootstrap estimates from `FEAR::boot.sw98` of efficiencies are sorted for each firm individually.

Unfortunately, this means that the component of replicas is not the efficiencies for the same bootstrap replica, but could easily be from different bootstrap replicas. This also means that this function can **not** be used to bootstrap tests for statistical hypotheses where the statistics involves summing of firms efficiencies.

**Author(s)**

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

**References**

Peter Bogetoft and Lars Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011.

Cinzia Dario and L. Simar; *Advanced Robust and Nonparametric Methods in Efficiency Analysis.* Methodology and Applications; Springer 2007.


**See Also**

The documentation for `boot.sw98` in the package `FEAR`.

**Examples**

```r
x <- matrix(c(100,200,300,500,100,200,600),ncol=1)
y <- matrix(c( 75,100,300,400, 25, 50,400),ncol=1)

e <- dea(x,y)

eff(e)

dea.plot.frontier(x,y,txt=TRUE)
```

# To bootstrap for real, NREP should be at least 2000. Run the
# following lines a couple of times with nrep=100 and see how the
# bootstrap frontier changes from one run to the next. Try the same
# with NREP=2000 even though is does take a longer time to run,
# especially for dea.boot.

nrep <- 5
nrep <- 2000

# if ( "FEAR" %in% .packages(TRUE) ) {
## The following only works if the package FEAR is installed; it does
## not have to be loaded.

```r
# b <- boot.fear(x, y, NREP=nrep)
# } else {
  b <- dea.boot(x, y, NREP=nrep)
# }
```

# bias corrected frontier
dea.plot.frontier(b$eff.bc*x, y, add=TRUE, lty="dashed")
# outer 95% confidence interval frontier for uncorrected frontier
dea.plot.frontier((b$conf.int[,1]+b$bias)*x, y, add=TRUE, lty="dotted")

## Test of hypothesis in DEA model
# Null hypothesis is that technology is CRS and the alternative is VRS
ec <- dea(x, y, RTS="crs")
Ec <- eff(ec)
ev <- dea(x, y, RTS="vrs")
Ev <- eff(ev)
# The test statistic; equation (6.1)
S <- sum(Ec)/sum(Ev)

# To calculate CRS and VRS efficiencies in the same bootstrap replicas
# we reset the random number generator before each call of the
# function dea.boot.

# To get the an initial value for the random number generating process
# we save its state (seed)
save.seed <- sample.int(1e9,1)

# The bootstrap and calculate CRS and VRS under the assumption that
# the true technology is CRS (the null hypothesis) and such that the
# results corresponds to the case where CRS and VRS are calculated for
# the same reference set of firms; to make this happen we set the
# random number generator to the same state before the calls.
set.seed(save.seed)
bc <- dea.boot(x, y, nrep, RTS="crs")
set.seed(save.seed)

# Calculate the statistic for each bootstrap replica
bs <- colSums(bc$boot)/colSums(bv$boot)
# The critical value for the test (default size \code{alpha} of test is 5%)
critValue(bs, alpha=.1)
S
# Accept the hypothesis at 10% level?
critValue(bs, alpha=.1) <= S

# The probability of observering a smaller value of S when the
# hypothesis is true; the p--value.
typeIerror(S, bs)
# Accept the hypothesis at size level 10%?
typeIerror(S, bs) >= .10
**dea.direct**  

*Directional efficiency*

**Description**

Directional efficiency rescaled to an interpretation à la Farrell efficiency and the corresponding peer importance (lambda).

**Usage**

```r
dea.direct(X, Y, DIRECT, RTS = "vrs", ORIENTATION = "in", XREF = NULL, YREF = NULL, FRONT.IDX = NULL, SLACK = FALSE, param=NULL, TRANSPOSE = FALSE)
```

**Arguments**

- **X**: Inputs of firms to be evaluated, a K x m matrix of observations of K firms with m inputs (firm x input).
- **Y**: Outputs of firms to be evaluated, a K x n matrix of observations of K firms with n outputs (firm x input).
- **DIRECT**: Directional efficiency, DIRECT is either a scalar, an array, or a matrix with non-negative elements.
  - If the argument is a scalar, the direction is (1,1,...,1) times the scalar; the value of the efficiency depends on the scalar as well as on the unit of measurements.
  - If the argument an array, this is used for the direction for every firm; the length of the array must correspond to the number of inputs and/or outputs depending on the ORIENTATION.
  - If the argument is a matrix then different directions are used for each firm. The dimensions depend on the ORIENTATION (and TRANSPOSE), the number of firms must correspond to the number of firms in X and Y.
  - DIRECT must not be used in connection with DIRECTION="graph".
- **RTS**: Text string or a number defining the underlying DEA technology / returns to scale assumption.
  - 0: fdh, Free disposability hull, no convexity assumption
  - 1: vrs, Variable returns to scale, convexity and free disposability
  - 2: drs, Decreasing returns to scale (down-scaling, but not up-scaling), convexity, and free disposability
  - 3: crs, Constant returns to scale, convexity and free disposability
  - 4: irs, Increasing returns to scale (up-scaling, but not down-scaling), convexity, and free disposability
  - 6: add, Additivity (scaling up and down, but only with integers), and free disposability
  - 7: fdh+, A combination of free disposability and restricted or local constant return to scale
- **ORIENTATION**: Input efficiency "in" (1), output efficiency "out" (2), and graph efficiency "graph" (3). For use with DIRECT, an additional option is "in-out" (0).
- **XREF**: Inputs of the firms determining the technology, defaults to X.
Outputs of the firms determining the technology, defaults to $Y$.

Index for firms determining the technology.

See `dea` and `slack`.

Possible parameters. At the moment only used for `RTS=“fdh+”` to set low and high values for restrictions on lambda; see the section details and examples in `dea` for its use. Future versions might also use `param` for other purposes.

see `dea`

Details

When the argument `DIRECT=d` is used then component `objval` of the returned object for input orientation is the maximum value of $e$ where for input orientation $x - ed$, and for output orientation $y + ed$ are in the generated technology set. The returned component `eff` is for input $1 - ed/X$ and for output $1 + ed/Y$ to make the interpretation as for a Farrell efficiency. Note that when the direction is not proportional to $X$ or $Y$ the returned `eff` are different for different inputs or outputs and `eff` is a matrix and not just an array. The directional efficiency can be restricted to inputs (ORIENTATION=“in”), restricted to outputs (ORIENTATION=“out”), or both include inputs and output directions (ORIENTATION=“in-out”). Directional efficiency is discussed on pages 31–35 and 121–127 in Bogetoft and Otto (2011).

The Farrell efficiency interpretation is the ratio by which a firm can proportionally reduce all inputs (or expand all outputs) without producing less outputs (using more inputs). The directional efficiencies have the same interpretation expect that the direction is not proportional to the inputs (or outputs) and therefore the different inputs may have different reduction ratios, the efficiency is an array and not just a number.

Value

The results are returned in a Farrell object with the following components. The method `slack` only returns the three components in the list relevant for slacks.

- `eff`: The Farrell efficiencies. Note that the efficiencies are calculated to have the same interpretations as Farrell efficiencies. `eff` is a matrix if there are more than 1 good.
- `lambda`: The lambdas, i.e. the weight of the peers, for each firm
- `objval`: The objective value as returned from the LP program; the `objval` are excess values in DIRECT units of measurement.
- `RTS`: The return to scale assumption as in the option `RTS` in the call
- `ORIENTATION`: The efficiency orientation as in the call
- `TRANSPOSE`: As in the call
- `slack`: A vector with sums of the slacks for each firm. Only calculated in `dea` when option `SLACK=TRUE`
- `sx`: A matrix for input slacks for each firm, only calculated if the option `SLACK` is `TRUE` or returned from the method `slack`
- `sy`: A matrix for output slack, see `sx`
**Note**

To handle fixed, non-discretionary inputs, one can let it appear as negative output in an input-based mode, and reversely for fixed, non-discretionary outputs. Fixed inputs (outputs) can also be handled by directional efficiency; set the direction, the argument `DIRECT`, equal to the variable, discretionary inputs (outputs) and 0 for the fixed inputs (outputs).

When the the argument `DIRECT=X` is used the then the returned efficiency is equal to 1 minus the Farrell efficiency for input orientation and equal to the Farrell efficiency minus 1 for output orientation.

**Author(s)**

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

**References**


Peter Bogetoft and Lars Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011

**See Also**

dea

dea.direct

**Examples**

```r
# Directional efficiency
x <- matrix(c(2, 5, 1, 2, 2, 2, 3, 2, 3, 1, 4, 1), ncol=2,byrow=TRUE)
y <- matrix(1,nrow=dim(x)[1])
dea.plot.isoquant(x[,1], x[,2],txt=1:dim(x)[1])

t <- dea(x,y)
z <- c(1,1)
e <- dea.direct(x,y,DIRECT=z)
data.frame(Farrell=t$eff, Perform=e$eff, objval=e$objval)

# The direction
arrows(x[,1], x[,2], (x-z)[,1], (x-z)[,2], lty="dashed")
# The efficiency (e$objval) along the direction
segments(x[,1], x[,2], (x-e$objval*z)[,1], (x-e$objval*z)[,2], lwd=2)

# Different directions
x1 <- c(.5, 1, 2, 4, 3, 1)
x2 <- c(4, 2, 1,.5, 2, 4)
x <- cbind(x1,x2)
y <- matrix(1,nrow=dim(x)[1])
dir1 <- c(1,.25)
dir2 <- c(.25, 4)
dir3 <- c(1,4)
e <- dea(x,y)
e1 <- dea.direct(x,y,DIRECT=dir1)
```

### dea.dual

**Dual DEA models and assurance regions**

#### Description

Solution of dual DEA models, possibly with partial value information given as restrictions on the ratios (assurance regions)

#### Usage

```r
dea.dual(X, Y, RTS = "vrs", ORIENTATION = "in",
          XREF = NULL, YREF = NULL,
          FRONT.IDX = NULL, DUAL = NULL, DIRECT=NULL,
          TRANSPOSE = FALSE, LP = FALSE, CONTROL=NULL, LPK=NULL)
```

#### Arguments

- **X**: Inputs of firms to be evaluated, a K x m matrix of observations of K firms with m inputs (firm x input). In case TRANSPOSE=TRUE the input matrix is transposed to input x firm.
Outputs of firms to be evaluated, a K x n matrix of observations of K firms with n outputs (firm x input). In case TRANSPOSE=TRUE the output matrix is transposed to output x firm.

RTS
A text string or a number defining the underlying DEA technology / returns to scale assumption.

1 vrs Variable returns to scale, convexity and free disposability
2 drs Decreasing returns to scale, convexity, down-scaling and free disposability
3 crs Constant returns to scale, convexity and free disposability
4 irs Increasing returns to scale, (up-scaling, but not down-scaling), convexity and free disposability.

ORIENTATION
Input efficiency "in" (1), output efficiency "out" (2), and graph efficiency "graph" (3) (not yet implemented). For use with DIRECT an additional option is "in-out" (0). In this case, "graph" is not feasible

XREF
Input of the firms determining the technology, defaults to X

YREF
Output of the firms determining the technology, defaults to Y

FRONT.IDX
Index for firms determining the technology

DUAL
Matrix of order “number of input plus number of outputs minus 2” times 2. The first column is the lower bound and the second column is the upper bound for the restrictions on the multiplier ratios. The ratios are relative to the first input and the first output, respectively. This implies that there is no restriction for neither the first input nor the first output so that the number of restrictions is two less than the total number of inputs and outputs.

DIRECT
Directional efficiency, DIRECT is either a scalar, an array, or a matrix with non-negative elements.

NB Not yet implemented

TRANSPOSE
Input and output matrices are treated as firms times goods for the default value TRANSPOSE=FALSE corresponding to the standard in R for statistical models. When TRUE data matrices shall be transposed to good times firms matrices as is normally used in LP formulation of the problem.

LP
Only for debugging. If LP=TRUE then input and output for the LP program are written to standard output for each unit.

CONTROL
Possible controls to lpSolveAPI, see the documentation for that package.

LPK
When LPK=k then a mps file is written for firm k; it can be used as input to an alternative LP solver just to check the our results.

Details
Solved as an LP program using the package lpSolveAPI. The method dea.dual.dea calls the method dea with the option DUAL=TRUE.

Value

eff
The efficiencies

objval
The objective value as returned from the LP problem, normally the same as eff
RTS

The return to scale assumption as in the option RTS in the call

ORIENTATION

The efficiency orientation as in the call

TRANSPOSE

As in the call

u

Dual values, prices, for inputs

v

Dual values, prices, for outputs

gamma

The values of gamma, the shadow price(s) for returns to scale restriction

sol

Solution of all variables as one component, sol=c(u,v,gamma).

Note

Note that the dual values are not unique for extreme points in the technology set. In this case the value of the calculated dual variable can depend on the order of the complete efficient firms.

Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

References

Bogetoft and Otto; Benchmarking with DEA, SFA, and R; Springer 2011. Sect. 5.10: Partial value information

See Also

dea

Examples

\begin{verbatim}
x <- matrix(c(2,5 , 1,2 , 2,2 , 3,2 , 3,1 , 4,1), ncol=2,byrow=TRUE)
y <- matrix(1,nrow=dim(x)[1])
dea.plot.isoquant(x[,1],x[,2],txt=1:dim(x)[1])
segments(0,0, x[,1], x[,2], lty="dotted")

e <- dea(x,y,RTS="crs",SLACK=TRUE)
ed <- dea.dual(x,y,RTS="crs")
print(cbind("e"=e$eff,"ed"=ed$eff, peers(e), lambda(e),
e$sx, e$sy, ed$u, ed$v), digits=3)

dual <- matrix(c(.5, 2.5), nrow=dim(x)[2]+dim(y)[2]-2, ncol=2, byrow=TRUE)
er <- dea.dual(x,y,RTS="crs", DUAL=dual)
print(cbind("e"=e$eff,"ar"=er$eff, lambda(e), e$sx, e$sy, er$u,
   "ratio"=er$u[,2]/er$u[,1],er$v),digits=3)
\end{verbatim}
dea.merge

Estimate potential merger gains and their decompositions

Description

Calculate and decompose potential gains from mergers of similar firms (horizontal integration).

Usage

dea.merge(X, Y, M, RTS = "vrs", ORIENTATION = "in",
XREF = NULL, YREF = NULL, FRONT.IDX = NULL, TRANSPOSE=FALSE)

Arguments

Most of the arguments correspond to the arguments in dea, with K firms, m inputs, and n outputs.

X K times m matrix as in dea
Y K times n matrix as in dea
M Kg times K matrix where each row defines a merger by the firms (columns) included; matrix as returned from method make.merge
RTS as in dea
ORIENTATION as in dea
XREF as in dea
YREF as in dea
FRONT.IDX as in dea
TRANSPOSE as in dea

Details

The K firms are merged into Kg new, merged firms.
The decomposition is summarized on page 275 and in table 9.1 page 276 in Bogetoft and Otto (2011) and is based on Bogetoft and Wang (2005)

Value

Eff Overall efficiencies of mergers, Kg vector
Estar Adjusted overall efficiencies of mergers after the removal of individual learning, Kg vector
learning Learning effects, Kg vector
harmony Harmony (scope) effects, Kg vector
size Size (scale) effects, Kg vector
dea.plot

Plot of DEA technologies

Description

Draw a graph of a DEA technology. Designed for two goods illustrations, either isoquant (2 inputs), transformation curve (2 outputs), or a production function (1 input and 1 output). If the number of good is larger than 2 then aggregation occur, either simple or weighted.

Usage

dea.plot(x, y, RTS="vrs", ORIENTATION="in-out", txt=NULL, add=FALSE, wx=NULL, wy=NULL, TRANSPOSE=FALSE, fex=1, GRID=FALSE, RANGE=FALSE, param=NULL, ..., xlim, ylim, xlab, ylab)
**Arguments**

- **x**  
The good illustrated on the first axis. If there are more than 1 input then inputs are just summed or, if wx is present, a weighted sum of inputs is used.

- **y**  
The good illustrated on the second axis. If there are more than 1 output then outputs are just summed or, if wy is present, a weighted sum of outputs is used.

- **x1, y1**  
The good illustrated on the first axis

- **x2, y2**  
The good illustrated on the second axis

- **RTS**  
Underlying DEA model / assumptions about returns to scale: "fdh" (0), "vrs" (1), "drs" (2), "crs" (3), "irs" (4), "irs2" (5) (irs without convexity), "add" (6), and "fdh+" (7). Numbers in parenthesis can also be used as values for RTS

- **ORIENTATION**  
Input-output graph of 1 input and 1 output is "in-out" (0), graph of 2 inputs is "in" (1), and graph of 2 outputs is "out" (2).

- **txt**  
txt is an array to label the observations. If txt=TRUE the observations are labeled by the observation number or rownames if there are any.

- **add**  
For add=T the technology is drawn on top of an existing graph. With the default add=F, a new graph is made.

- **wx**  
Weight to aggregate the first axis if there are more than 1 good behind the first axis.

- **wy**  
Weights to aggregate for the second axis if there are more than 1 good behind the second axis.

- **TRANSPOSE**  
Only relevant for more than 1 good for each axis, see dea for a description of this option.

- **GRID**  
If GRIF=TRUE a gray grid is put on the plot.

- **fex**  
Relative size of the text/labels on observations; corresponds to cex, but only changes the size of the text.

- **RANGE**  
A logical variable, if RANGE=TRUE the limits for the graph is the range of the variables; zero is always included. Default is RANGE=FALSE when the range is from zero to the max values. Relevant if some values are negative.

- **param**  
Possible parameters. At the moment only used for RTS="fdh+"; see the section details and examples for its use. Future versions might also use param for other purposes.

- **xlim**  
Possible limits c(x1,x2) for the first axis

- **ylim**  
Possible limits c(y1,y2) for the second axis

- **xlab**  
Possible label for the x-axis

- **ylab**  
Possible label for the y-axis
Details

The method `dea.plot` is the general plotting method. The three others are specialized versions for frontiers (1 input and 1 output), isoquant curves (2 inputs for given outputs), and transformation curves (2 outputs for given inputs) obtained by using the argument `ORIENTATION`.

The `crs` factor in `RTS="fdh+"` that sets the lower and upper bound can be changed by the argument `param` that will set the lower and upper bound to `1-param` and `1+param`; the default value is `param=.15`. The value must be greater than or equal to 0 and strictly less than 1. A value of 0 corresponds to `RTS="fdh"`. The FDH+ technology set is described in Bogetoft and Otto (2011) pages 72–73.

Value

No return, uses the original graphing system.

Note

If there are more than 1 good for the arguments `x` and `y` then the goods are just summed or, if `wx` or `wy` are present, weighted sum of goods are used. In this case the use of the command `identify` must be called as `dea.plot(rowSums(x),rowSums(y))`.

Warning If you use this facility to plot multi input and multi output then the plot may deceive you as fully multi efficient firms are not necessarily placed on the two dimensional frontier.

Note that `RTS="add"` and `RTS="fdh+"` only works for `ORIENTATION="in-out"` (0).

Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

References

Peter Bogetoft and Lars Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011
Paul Murrell; *R Graphics*; Chapman & Hall 2006

See Also

The documentation for the function `plot` and Murrell (2006) for further options and on customizing plots.

Examples

```r
x <- matrix(c(100,200,300,500,600,100),ncol=1)
y <- matrix(c(75,100,300,400,400,50),ncol=1)

dea.plot(x,y,RTS="vrs",ORIENTATION="in-out",txt=LETTERS[1:length(x)])
dea.plot(x,y,RTS="crs",ORIENTATION="in-out",add=TRUE,lty="dashed")

dea.plot.frontier(x,y,txt=1:dim(x)[1])
n <- 10
```
Calculate efficiencies for Farrell and sfa object

Description

Calculate efficiencies for Farrell and sfa object. For a sfa there are several types

Usage

eff( object, ... )
efficiencies( object, ... )
# Default S3 method:
efficiencies( object, ... )
# S3 method for class 'Farrell'
efficiencies(object, type = "Farrell", ...)
# S3 method for class 'Farrell'
eff(object, type = "Farrell", ...)
# S3 method for class 'sfa'
efficiencies(object, type = "BC", ...)
# S3 method for class 'sfa'
eff(object, type = "BC", ...)
Arguments

object A Farrell object returned from a DEA function like `dea`, `sdea`, or `mea` or an sfa object returned from the function `sfa`.

type The type of efficiencies to be calculated. For a Farrell object the possibilities are “Farrell” efficiency or “Shephard” efficiency. For a sfa object the possibilities are “BC”, “Mode”, “J”, or “add”.

Details

The possible types for class `Farrell` (an object returned from `dea` et al. are “Farrell” and “Shephard”.

The possible types for class `sfa` efficiencies are

- **BC** Efficiencies estimated by minimizing the mean square error; Eq. (7.21) in Bogetoft and Otto (2011, 219) and Battese and Coelli (1988, 392)
- **Mode** Efficiencies estimates using the conditional mode approach; Bogetoft and Otto (2011, 219), Jondrow et al. (1982, 235).
- **J** Efficiencies estimates using the conditional mean approach Jondrow et al. (1982, 235).
- **add** Efficiency in the additive model, Bogetoft and Otto (2011, 219)

Value

The efficiencies are returned as an array.

Note

For the Farrell object the orientation is determined by the calculations that led to the object and can not be changed here.

Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

References

Bogetoft and Otto; *Benchmarking with DEA, SFA, and R*, Springer 2011

See Also

- `dea` and `sfa`

Examples

```r
## Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##-- or do help(data=index) for the standard data sets.
```
**Description**

A method to estimate and plot kernel estimate of (Farrell) efficiencies taken into consideration that efficiencies are bounded either above (input direction) or below (output direction).

**Usage**

```r
eff.dens(eff, bw = "nrd0")
eff.dens.plot(obj, bw = "nrd0", ..., xlim, ylim, xlab, ylab)
```

**Arguments**

- `eff`: Either a list of (Farrell) efficiencies or a Farrell object returned from the method `dea`.
- `bw`: Bandwith, look at the documentation of `density` for an explanation.
- `obj`: Either an array of efficiencies or a list returned from `eff.dens`.
- `...`: Further arguments to the `plot` method like line type and line width.
- `xlim`: Range on the x-axis; usually not needed, just use the defaults.
- `ylim`: Range on the x-axis; usually not needed, just use the defaults.
- `xlab`: Label for the x-axis.
- `ylab`: Label for the y-axis.

**Details**

The calculation is based on a reflection method (Silverman 1986, 30) using the default window kernel and default bandwidth (window width) in the method `density`.

The method `eff.dens.plot` plot the density directly, and `eff.dens` just estimate the numerical density, and the result can then either be plotted by `plot`, corresponds to `eff.dens.plot`, or by lines as an overlay on an existing plot.

**Value**

The return from `eff.dens` is a list `list(x,y)` with efficiencies and the corresponding density values.

**Note**

The input efficiency is also bounded below by 0, but for normal firms an efficiency at 0 will not happen, i.e. the boundary is not effective, and therefore this boundary is not taken into consideration.
Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

References


Examples

```r
e <- 1 - rnorm(100)
e[e>1] <- 1
e <- e[e>0]
eff.dens.plot(e)

hist(e, breaks=15, freq=FALSE, xlab="Efficiency", main="")
den <- eff.dens(e)
lines(den, lw=2)
```

eladder

*Efficiency ladder for a single firm*

Description

How the efficiency changes as the most influential peer is removed sequentially one at a time

Usage

```r
eladder(n, X, Y, RTS = "vrs", ORIENTATION = "in",
XREF=NULL, YREF=NULL, DIRECT = NULL, param=NULL, MAXELAD=NULL)
eladder.plot(elad, peer, TRIM = NULL, ...)
```

Arguments

- `n` The number of the firm where the ladder is calculated
- `X` Inputs of firms to be evaluated, a K x m matrix of observations of K firms with m inputs (firm x input). In case TRANSPOSE=TRUE the input matrix is transposed to input x firm.
- `Y` Outputs of firms to be evaluated, a K x n matrix of observations of K firms with n outputs (firm x input). In case TRANSPOSE=TRUE the output matrix is transposed to output x firm.
- `RTS` Text string or a number defining the underlying DEA technology / returns to scale assumption, see the possible values for `dea`
- `ORIENTATION` Input efficiency "in" (1), output efficiency "out" (2), and graph efficiency "graph" (3). For use with DIRECT, an additional option is "in-out" (0).
- `XREF` Inputs of the firms determining the technology, defaults to X
eladder

YREF Outputs of the firms determining the technology, defaults to Y
DIRECT Directional efficiency, DIRECT is either a scalar, an array, or a matrix with non-negative elements. See dea for a further description of this argument.
param Possible parameters. At the moment only used for RTS="fdh+" to set low and high values for restrictions on lambda; see the section details and examples in dea for its use. Future versions might also use param for other purposes.
MAXELAD The maximum number of influential peers to remove.
elad The sequence of efficiencies returned from eladder.
peer The sequence of peers returned from eladder.
TRIM The number of characters for the name of the peers on the axis in the plot.
... Usual options for the method plot.

Details

The function eladder calculates how the efficiency for a firm changes when the most influential peer is removed sequentially one at a time. Somewhere in the sequence the firm becomes efficient and are itself removed from the set of firms generating the technology (or the only firm left) and thereafter the efficiencies are super-efficiencies.

Value

The object returned from eladder is a list with components

eff The sequence of efficiencies when the peer with the largest value of lambda has been removed.
peer The sequence of removed peers corresponding to the largest values of lambda as index in the X rows.

Note

When the number of firms is large then the number of influential peers will also be large and the names or numbers of the peers on the x-axis might be squeeze together and be illegible. In this case restrict the number of influential peers to be removed.


Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

References

Dag Fjeld Edvardsen; Four Essays on the Measurement of Productive Efficiency; University of Gothenburg 2004; http://hdl.handle.net/2077/2923
Examples

```r
data(charnes1981)
x <- with(charnes1981, cbind(x1,x2,x3,x4,x5))
y <- with(charnes1981, cbind(y1,y2,y3))

# Choose the firm for analysis, we choose 'Tacoma'
n <- which(charnes1981$name=="Tacoma")[1]
el <- eladder(n, x, y, RTS="crs")
eladder.plot(el$eff, el$peer)

# Restrict to 20 most influential peers for 'Tacoma' and use names instead of number
eladder.plot(el$eff[1:20], charnes1981$name[el$peer][1:20])

# Truncate the names of the peers and put a title on top
eladder.plot(el$eff[1:20], charnes1981$name[el$peer][1:20], TRIM=5)
title("Eladder for Tacoma")
```

excess

**Excess input compared over frontier input**

### Description

Excess input compared over frontier input and/or less output than frontier/transformation/optimal output.

### Usage

```r
excess(object, X = NULL, Y = NULL)
```

### Arguments

- **object**: A Farrell object as returned from functions like `dea`, `dea.direct`, `linksdea`, and `mea`.
- **X**: Input matrix, only necessary for ordinary input Farrell efficiency
- **Y**: Output matrix, only necessary for ordinary output Farrell efficiency

### Details

For Farrell input efficiency $E$ the excess input is $(1 - E)X$ and for Farrell output efficiency $F$ the missing output is $(F - 1)Y$.

Notice that the excess calculated does not include any slack values. In case slacks are present and calculated it might be more appropriate to add slack; i.e. to use $\text{excess}(\text{object}, X, Y) + \text{slack}(X, Y, \text{object})$.

For directional efficiency $e$ in the direction $D$ the excess input is $eD$.

If a firm is outside the technology set, as could be the case when calculating super-efficiencies, the Farrell input efficiency is larger than 1 and then the excess values are negative.
**Value**

Return a matrix with excess input and/or less output.

**Author(s)**

Peter Bogeroft and Lars Otto <larsot23@gmail.com>

**References**

Peter Bogetoft and Lars Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011

**Examples**

```r
x <- matrix(c(100,200,300,500,100,200,600),ncol=1)
y <- matrix(c(75,100,300,400,25,50,400),ncol=1)

e <- dea(x,y)
excess(e,x)
x - eff(e) * x

e <- dea(x,y, ORIENTATION="graph")
excess(e, x, y)
x - eff(e) * x
1/eff(e) * y -y

me <- mea(x,y)
excess(me)
```

---

### lambda

**Lambdas or the weight of the peers**

**Description**

The lambdas, i.e. the weight of the peers, for each firm.

**Usage**

```r
lambda(object, KEEPREF = FALSE)
lambda.print(x, KEEPREF = FALSE, ...)
```

**Arguments**

- **object, x**
  - A Farrell object as returned from `dea` et al.
- **KEEPREF**
  - if TRUE then all firms are kept as reference firms even though they have all zero weights (lambda); might come handy if one needs to calculate X x lambda such that the firms in X and lambda agree. If FALSE, the default, then only weight for the peers are in the matrix lambda.
- **...**
  - Optional parameters for the print method.
**Details**

Only returns the the lambdas for firms that appear as a peer, i.e. only lambdas for firms where at least one element of the lambda is positive.

**Value**

The return is a matrix with the firms as rows and the peers as columns.

**Author(s)**

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

**See Also**

dea

**Examples**

```r
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##-- or do help(data=index) for the standard data sets.

make.merge
```

---

**make.merge**

*Make an aggregation matrix to perform mergers*

**Description**

Make an aggregation matrix to perform mergers of firms. The matrix can be post multiplied (matrix multiplication) to input and output matrices to make merged input and output matrices.

**Usage**

```r
make.merge(grp, nFirm = NULL, X = NULL, names = NULL)
```

**Arguments**

- **grp**
  - Either a list of length Kg for Kg firms after mergers; each component of the list is a (named) list with the firm numbers or names going into this merger.
  - Or a factor of length K with Kg levels where each level determines a merger; to exclude firms for mergers set the factor value to NA.

- **nFirm**
  - Number of firms before the mergers

- **X**
  - A matrix of inputs or outputs where the rows corresponds to the number of original (starting) firms

- **names**
  - A list with names of all firms, only needed if the mergers are given as a list of names, i.e. `grp` is a list of names.
Details

Either `nFirm` or `X` must be present; if both are present then `nFirm` must be equal to the number of rows in `X`, the number of firms.

When `X` is an input matrix of dimension `K x m`, `K` firms and `m` inputs, and `M <- make.merge(gr,K)` then `M %*% X` is the input matrix for the merged firms.

Value

Returns an aggregation matrix of dimension `Kg` times `K` where rows corresponds to new merged firms and columns are 1 for firms to be included and 0 for firms to be excluded in the the given merger as defined by the row.

Note

The argument `TRANSPOSE` has not been implemented for this function. If you need transposed matrices you must transpose the merger matrix yourself. If you define mergers via factors there is no need to transpose in the arguments; just do not use `X` in the arguments.

Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

See Also

dea.merge

Examples

# To merge firms 1,3, and 5; and 2 and 4 of 7 firms into 2 new firms
# the aggregation matrix is M; not all firms are involved in a merger.
M <- make.merge(list(c(1,3,5), c(2,4)),7)
print(M)

# Merge 1 and 2, and 4 and 5, and leave 3 alone, total of 5 firms.
# Using a list
M1 <- make.merge(list(c(1,2), c(4,5)), nFirm=5)
print(M1)

# Using a factor
fgr <- factor(c("en","en",NA,"to","to"))
M2 <- make.merge(fgr)
print(M2)

# Name of mergers
M3 <- make.merge(list(AB=c("A","B"), DE=c("D","E")), names=LETTERS[1:5])
print(M3)

# No name of mergers
M4 <- make.merge(list(c("A","B"), c("D","E")), names=LETTERS[1:5])
print(M4)
malmq

Description

Estimates Malmquist indices for productivity and its decomposition between two periods. The units in the two periods do not have to be exactly the same, but the Malmquist index is only calculated for units present in both periods.

Usage

\[ \text{malmq}(X_0, Y_0, \text{ID}_0 = \text{NULL}, X_1, Y_1, \text{ID}_1 = \text{NULL}, \text{RTS} = \text{"vrs"}, \text{ORIENTATION} = \text{"in"}, \text{SLACK} = \text{FALSE}, \text{DUAL} = \text{FALSE}, \text{DIRECT} = \text{NULL}, \text{param} = \text{NULL}, \text{TRANSPOSE} = \text{FALSE}, \text{FAST} = \text{TRUE}, \text{LP} = \text{FALSE}, \text{CONTROL} = \text{NULL}, \text{LPK} = \text{NULL}) \]

Arguments

- **X0**: Inputs of firms in period 0, a K0 x m matrix of observations of K0 firms with m inputs (firm x input).
- **Y0**: Outputs of firms in period 0, a K0 x n matrix of observations of K0 firms with n outputs (firm x input).
- **ID0**: Index for firms in period 0; could be numbers or labels. Length K0.
- **X1**: Inputs of firms in period 1, a K1 x m matrix of observations of K1 firms with m inputs (firm x input).
- **Y1**: Outputs of firms in period 1, a K1 x n matrix of observations of K1 firms with n outputs (firm x input).
- **ID1**: Index for firms in period 0; could be numbers or labels. Length K0.
- **RTS**: Returns to scale assumption as in \text{dea}.
- **ORIENTATION**: Input efficiency "in" (1), output efficiency "out" (2), and graph efficiency "graph" (3) as in \text{dea}.
- **SLACK**: See \text{dea}.
- **DUAL**: See \text{dea}.
- **DIRECT**: See \text{dea}.
- **param**: See \text{dea}.
- **TRANSPOSE**: See \text{dea}.
- **FAST**: See \text{dea}.
- **LP**: See \text{dea}.
- **CONTROL**: See \text{dea}.
- **LPK**: See \text{dea}.
Details

The index for technical changes $tc$ is calculated as $\sqrt{\frac{e_{10}/e_{11} \cdot e_{00}/e_{01}}{}}$ where $e_{s<t}$ is the efficiency for period $s$ when the reference technology is for period $t$, i.e. determined from the observations for period $t$ and XREF=X_t, YREF=Y_t, as is the option for the function dea.

The Malmquist index for productivity $mq$ is calculates as $\sqrt{\frac{e_{10}/e_{00} \cdot e_{11}/e_{01}}{}}$ and the index for change in efficiency $ec$ is $e_{11}/e_{00}$. Note that $mq = tc \cdot ec$.

Value

- $m$: Malmquist index for productivity.
- $tc$: Index for technology change.
- $ec$: Index for efficiency change.
- $mq$: Malmquist index for productivity; same as $m$.
- $id$: Index for firms present in both period 0 and period 1.
- $id0$: Index for firms in period 0 that are also in period 1.
- $id1$: Index for firms in period 1 that are also in period 0.
- $e00$: The efficiencies for period 0 with reference technology from period 0.
- $e10$: The efficiencies for period 1 with reference technology from period 0.
- $e11$: The efficiencies for period 1 with reference technology from period 1.
- $e01$: The efficiencies for period 0 with reference technology from period 1.

Note

The calculations of efficiencies are only done for units present in both periods.

Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

References

Peter Bogetoft and Lars Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011

See Also

dea

Examples

```r
x0 <- matrix(c(10, 28, 30, 60),ncol=1)
y0 <- matrix(c(5, 7, 10, 15),ncol=1)
x1 <- matrix(c(12, 26, 16, 60 ),ncol=1)
y1 <- matrix(c(6, 8, 9, 15 ),ncol=1)

da.plot(x0, y0, RTS="vrs", txt=TRUE)
da.plot(x1, y1, RTS="vrs", add=TRUE, col="red")
```
malmquist

Malmquist index for firms in a panel

Description

Estimate Malmquist index for firms in a panel data set. The data set does not need to be balanced.

Usage

malmquist(X, Y, ID, TIME, RTS = "vrs", ORIENTATION = "in", SLACK = FALSE, DUAL = FALSE,
DIRECT = NULL, param = NULL, TRANSPOSE = FALSE, FAST = TRUE, LP = FALSE,
CONTROL = NULL, LPK = NULL)

Arguments

X Inputs of firms in many periods, a (T*K) x m matrix of observations of K firms
with m outputs (firm x input) in at the most T periods.
Y Outputs of firms in many periods, a (T*K) x n matrix of observations of K0
firms with n outputs (firm x input) in at the most T periods.
ID Identifier for the firms in rows of X and Y.
TIME Array with period number for each row in the input matrix X and output matrix Y
RTS Returns to scale assumption as in dea.
ORIENTATION Input efficiency "in" (1), output efficiency "out" (2), and graph efficiency "graph"
(3) as in dea.
SLACK See dea.
DUAL See dea.
DIRECT See dea.
param See dea.
TRANSPOSE See dea.
FAST See dea.
LP See dea.
CONTROL See dea.
LPK See dea.
**Details**

Malmquist uses \texttt{malmq} for the calculations of the necessary efficiencies, and the returned indeces are as in \texttt{malmq}. The data must be a long data set with regards to \text{TIME} and \text{ID}; see the example below.

Note that the calculated index are index comparing a period and the previous period. To compare the development over time the indices must be turned into a chain index as shown in the example below.

**Value**

- \texttt{m}: Malmquist indicies, an array of length T*K in the order of ID and TIME, i.e. the order of the rows of \texttt{X}.
- \texttt{tc}: Technical change indices, an array of length T*K.
- \texttt{ec}: Efficiency indices, an array of length T*K.
- \texttt{id}: Index for firms as ID
- \texttt{time}: Index for time as TIME
- \texttt{e00}: The efficiencies for period 0 with reference technology from period 0.
- \texttt{e10}: The efficiencies for period 1 with reference technology from period 0.
- \texttt{e11}: The efficiencies for period 1 with reference technology from period 1.
- \texttt{e01}: The efficiencies for period 0 with reference technology from period 1.

**Note**

The lagged values \texttt{e11} are not neccesary equal to values of \texttt{e00} as the reference technology for the two periods could be generated by different units, if the units in different time periods are not the same.

**Author(s)**

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

**References**

Peter Bogetoft and Lars Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011

**See Also**

\texttt{dea, malmq}

**Examples**

```r
x0 <- matrix(c(10, 28, 30, 60), ncol=1)
y0 <- matrix(c(5, 7, 10, 15), ncol=1)
x1 <- matrix(c(12, 26, 16, 60), ncol=1)
y1 <- matrix(c(6, 8, 9, 15), ncol=1)
x2 <- matrix(c(13, 26, 15, 60), ncol=1)
y2 <- matrix(c(7, 9, 10, 15), ncol=1)
```
dea.plot(x0, y0, RTS="vrs", txt=TRUE)
dea.plot(x1, y1, RTS="vrs", add=TRUE, col="red")
dea.plot(x2, y2, RTS="vrs", add=TRUE, col="blue")
points(x1, y1, col="red", pch=16)
# points(x2, y2, col="blue", pch=17)
text(x1, y1, 1:dim(x1)[1], col="red", adj=-1)
text(x2, y2, 1:dim(x1)[1], col="blue", adj=-1)
legend("bottomright", legend=c("Period 0", "Period 1", "Period 2"),
  col=c("black", "red", "blue"), lty=1, pch=c(1,16, 17), bty="n")
X <- rbind(x0, x1, x2)
Y <- rbind(y0, y1, y2)
# Make ID and TIME variables one way or another
ID <- rep(1:dim(x1)[1], 3)
# TIME <- c(rep(0,dim(x1)[1]), rep(1,dim(x1)[1]), rep(2,dim(x1)[1]))
TIME <- gl(3, dim(x1)[1], labels=0:2)
# This is how the data for malmquist must look like
data.frame(TIME, ID, X, Y)
mq <- malmquist(X,Y, ID, TIME)
data.frame(TIME, ID, X, Y, mq$e00, mq$e01, mq$e10, mq$e11, mq$m, mq$tc)[order(ID, TIME),]

# How to make the Malmquist indices to a chain index
# Make data.frame with indices
DM <- data.frame(TIME, ID, m=mq$m, tc=mq$tc, ec=mq$ec)
# Set missing index for first period to 1, the base
DM[DM$TIME==0, c("m", "tc", "ec")]<-1
# Make chain index of the individual indices
AD <- aggregate(cbind(m=DM$m), by=list(ID=DM$ID), cumprod)
# Compare chain index to original index
data.frame(ID, TIME, m=AD$m, DM$m)

---

**mea**

**MEA multi-directional efficiency analysis**

### Description

Potential improvements PI or multi-directional efficiency analysis. The result is an excess value measures by the direction.

The direction is determined by the direction corresponding to the minimum input/maximum direction each good can be changed when they are changed one at a time.

### Usage

```r
mea(X, Y, RTS = "vrs", ORIENTATION = "in", XREF = NULL, YREF = NULL,
    FRONT.IDX = NULL, param=NULL, TRANSPOSE = FALSE,
    LP = FALSE, CONTROL = NULL, LPK = NULL)
mea.lines(N, X, Y, ORIENTATION="in")
```
Arguments

\[ X \]
K times m matrix with K firms and m inputs as in dea

\[ Y \]
K times n matrix with K firms and n outputs as in dea

\[ \text{RTS} \]
Text string or a number defining the underlying DEA technology / returns to scale assumption.

0  fdh  Free disposability hull, no convexity assumption
1  vrs  Variable returns to scale, convexity and free disposability
2  drs  Decreasing returns to scale, convexity, down-scaling and free disposability
3  crs  Constant returns to scale, convexity and free disposability
4  irs  Increasing returns to scale, (up-scaling, but not down-scaling), convexity and free disposability
6  add  Additivity (scaling up and down, but only with integers), and free disposability
7  fdh+  A combination of free disposability and restricted or local constant return to scale

ORIENTATION
Input efficiency "in" (1) or output efficiency "out" (2), and also the additional option "in-out" (0) for both input and output direction.

XREF
Inputs of the firms determining the technology, defaults to \[ X \]

YREF
Outputs of the firms determining the technology, defaults to \[ Y \]

FRONT.IDX
Index for firms determining the technology

param
Possible parameters. At the moment only used for RTS=“fdh+'' to set low and high values for restrictions on lambda; see the section details and examples in dea for its use. Future versions might also use param for other purposes.

TRANSPOSE
as in dea

LP
as in dea

CONTROL
as in dea

LPK
as in dea

\[ N \]
Number of firms where directional lines are to be drawn on an all ready existing frontier plot (dea.plot.frontier)

Details
Details can be found in Bogetoft and Otto (2011, 121–124).

This method is for input directional efficiency only interesting when there are 2 or more inputs, and for output only when there are 2 or more outputs.

Value
The results are returned in a Farrell object with the following components.

\[ \text{eff} \]
Exces value in DIRECT units of measurement, this is not Farrell efficiency

\[ \lambda \text{mbda} \]
The lambdas, i.e. the weight of the peers, for each firm

\[ \text{objval} \]
The objective value as returned from the LP program; normally the same as eff

\[ \text{RTS} \]
The return to scale assumption as in the option RTS in the call
ORIENTATION  The efficiency orientation as in the call

direct  A K times mlnm+n matrix with directions for each firm: the number of columns
depends on whether it is input, output or in-out orientated.

TRANSPOSE  As in the call

Note

The calculation is done in dea after a calculation of the direction that then is used in the argument
DIRECT. The calculation of the direction is done in a series LP programs, one for each good in the
direction.

Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

References

Peter Bogetoft and Lars Otto; Benchmarking with DEA, SFA, and R; Springer 2011

See Also

dea and the argument DIRECT.

Examples

```r
X <- matrix(c(2, 2, 5, 10, 10, 3, 12, 8, 5, 4, 6, 12), ncol=2)
Y <- matrix(rep(1,dim(X)[1]), ncol=1)

dea.plot.isoquant(X[,1], X[,2], txt=1:dim(X)[1])
mea.lines(c(5,6),X,Y)

me <- mea(X,Y)
me
peers(me)
# MEA potential saving in inputs, excess inputs
eff(me) * me$direct
me$eff * me$direct

# Compare to traditionally Farrell efficiency
e <- dea(X,Y)
e
peers(e)
# Farrell potential saving in inputs, excess inputs
(1-eff(e)) * X
```
**milkProd**

---

**Data: Milk producers**

---

**Description**

Data collected from Danish milk producers.

**Usage**

data(milkProd)

**Format**

A data frame with 108 observations on the following 5 variables.

- **farmNo**: farm number
- **milk**: Output of milk, kg
- **energy**: Energy expenses
- **vet**: Veterinary expenses
- **cows**: Number of cows

**Note**

Data as .csv are loaded by the command `data` using `read.table(..., header = TRUE, sep = ";")` such that this file is a semicolon separated file and not a comma separated file.

**Source**

Accounting and business check data

**Examples**

data(milkProd)
y <- with(milkProd, cbind(milk))
x <- with(milkProd, cbind(energy, vet, cows))
Description

A data set for 113 farmers in forestry in Norway.

Usage

data(norWood2004)

Format

A data frame with 113 observations on the following 7 variables.

- **firm**: firm number
- **m**: Variable cost
- **x**: Woodland, value of forest and land
- **y**: Profit
- **z1**: Secondary income from ordinary farming
- **z3**: Age of forest owner
- **z6**: Whether there is a long-term plan =1 or not =0

Details

Collected from farmers in forestry.

Note

Data as `.csv` are loaded by the command `data` using `read.table(..., header=TRUE, sep=";")` such that this file is a semicolon separated file and not a comma separated file.

Source

Norwegian Agricultural Economics Research Institute.

Examples

data(norWood2004)

```r
# maybe str(norWood2004); plot(norWood2004) ...
```
Description

The functions implements the Wilson (1993) outlier detection method using only R functions.

Usage

```r
outlier.ap(X, Y, NDEL = 3, NLEN = 25, TRANSPOSE = FALSE)

outlier.ap.plot(ratio, NLEN = 25, xlab = "r", ylab = "Log ratio",
                ..., ylim)
```

Arguments

- **X**: Input as a firms times goods matrix, see TRANSPOSE.
- **Y**: Output as a firms times goods matrix, see TRANSPOSE.
- **NDEL**: The maximum number of firms to be considered as a group of outliers, i.e. the maximum number of firms to be deleted.
- **NLEN**: The number of ratios to save for each level or removal, the number of rows in ratio used.
- **TRANSPOSE**: Input and output matrices are treated as firms times goods matrices for the default value TRANSPOSE=FALSE corresponding to the standard in R for statistical models. When TRUE data matrices are transposed to good times firms matrices as is normally used in LP formulation of the problem.
- **ratio**: The ratio component from the list as output from outlier.ap.
- **xlab**: Label for the x-axis.
- **ylab**: Label for the y-axis
- **ylim**: The y limits `(y1,y2)` of the plot, an array/vector of length 2.
- **...**: Usual options for the methods plot and lines.

Details

An implementation of the method in Wilson (1993) using only R functions and especially the function `det` to calculate $R_{\text{min}}^{(i)}$. An elementary presentation of the method is found in Bogetoft and Otto (2011), Sect. 5.13 on outliers.

Value

- **ratio**: A `min(NLEN,K) x NDEL` matrix with the log-ratios to be plotted.
- **imat**: A `NDEL x NDEL` matrix with indicies for deleted firms.
- **r0**: A `NDEL` array with the minimum value $R_i^0$ of the for each number of deleted firms.
Note

The function `outlier.ap` is extremely slow and for `NDEL` larger than 3 or 4 it might be advisable to use the function `ap` from the package **FEAR**.

The name of the returned components are the same as for `ap` in the package **FEAR**.

Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

References

Bogetoft and Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011


See Also

The function `ap` in the package **FEAR**.

Examples

```r
n <- 25
x <- matrix(rnorm(n))
y <- .5 + 2.5*x + 2*rnorm(25)
tap <- outlier.ap(x, y, NDEL=2)
print(cbind(tap$imat, tap$rmin), na.print="", digit=2)
outlier.ap.plot(tap$ratio)
```

peers

Find peer firms and units

Description

The function `peers` finds for each firm its peers, `get.number.peers` finds for each peer the number of times this peer appears as a peer, and `get.which.peers` determines for one or more peers the firms they appear as peers for. Also include a function `get.peers.lambda` to calculate for firms the importances (lambdas) of peers.

Usage

```r
peers(object, NAMES = FALSE, N=1:dim(object$lambda)[1], LAMBDA=0)
get.number.peers(object, NAMES = FALSE, N=1:dim(object$lambda)[2], LAMBDA=0)
get.which.peers(object, N = 1:dim(object$lambda)[2], LAMBDA=0)
get.peers.lambda(object, N=1:dim(object$lambda)[1], LAMBDA=0)
```
**peers**

**Arguments**

- **object**: An object of class Farrell as returned by the functions `dea, dea.direct` et al.
- **NAMES**: If true then names for the peers are returned if names are available otherwise the unit index numbers are used. If NAMES is a list of names with length equal to the number of units then it is used as names for peers.
- **N**: The firm(s) or peer(s) for which to get the results.
- **LAMBDA**: Minimum weight for extracted peers, i.e. the extracted peers have lambda values larger than LAMBDA.

**Details**

The returned values are index of the firms and can be used by itself, but can also by used as an index for a variable with names of the firms.

The `peers` returns a matrix with numbers for the peers for each firm; for firms with efficiency 1 the peers are just the firm itself. If there is slack in the evaluation of a firm with efficiency 1, this can be found with a call to `slack`, either directly or by the argument SLACK when a function `dea` was called to generate the Farrell object.

The `get.number.peers` returns the number of firms that a peer serves as a peer for.

The `get.peers.lambda` returns a list of firms with the peers and corresponding value af lambda.

**Value**

The return values are firm numbers. If the argument NAMES=TRUE is used in the function `peers` the return is a list of names of the peers if names for the firms are available as rownames.

**Note**

Peers are defined as firms where the corresponding lambdas are positive.

Note that peers might change between a Farrell object return from `dea` with SLACK=FALSE and a call with SLACK=TRUE or a following call to the function `slack` because a peer on the frontier with slack might by the call to `dea` be seen as a peer for itself whereas this will not happen when slacks are calculated.

**Author(s)**

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

**References**

Peter Bogetoft and Lars Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011. Sect. 4.6 page 93

**See Also**

`dea`
Examples

```R
x <- matrix(c(100,200,300,500,100,200,600),ncol=1)
y <- matrix(c(75,100,300,400,25,50,400),ncol=1)
e <- dea(x,y)
peers(e)
get.number.peers(e)

# Who are the firms that firm 1 and 4 is peers for
get.which.peers(e, c(1,4))
```

**pigdata**

*Data: Multi-output pig producers*

**Description**

Input and output data for 248 pig producers that also produces crop, i.e. a multi–output data set.

**Usage**

`data(pigdata)`

**Format**

A data frame with 248 observations on the following 16 variables.

- **firm** Serial number for pig producer
- **x1** Input fertilizer
- **x2** Input feedstuf
- **x3** Input land
- **x4** Input labour
- **x5** Input machinery
- **x6** Input other capital
- **y2** Output crop
- **y4** Output pig
- **w1** Price of fertilizer
- **w2** Price of feedstuf
- **w3** Price of land
- **w4** Price of labour
- **w5** Price of machinery
- **w6** Price of other capital
- **p2** Price of pig
- **p4** Price of crop
- **cost** Total cost, \( w_1 x_1 + \ldots + w_6 x_6 \).
- **rev** Total revenue, \( p_2 y_2 + p_4 y_4 \).
**Details**

In raising pigs, most farmers also produce crops to feed the pigs. Labor and capital are used not just directly for pig-raising but also on the field.

**Note**

Data as .csv are loaded by the command `data` using `read.table(..., header = TRUE, sep = "");") such that this file is a semicolon separated file and not a comma separated file.

**Source**

Farmers accounting data converted to index.

**Examples**

```r
data(pigdata)
## maybe str(pigdata); plot(pigdata) ...
```

---

**projekt Data: Milk producers**

**Description**

Accounting and production data for 101 milk producing farmers.

**Usage**

```r
data(projekt)
```

**Format**

A data frame with 101 observations on the following 14 variables.

- `numb` Serial number for the milk producer
- `cows` Number of cows
- `vet` Veterinary expences
- `unitCost` Unit cost, variable cost
- `capCost` Capacity cost
- `fixedCost` Fixed cost
- `milkPerCow` Milk per cow, kg
- `quota` Milk quota
- `fatPct` Fat percent in milk
- `protPct` Protein percent in milk
- `cellCount` Cell count for milk
- `race` Race for cows, a factor with levels jersey, large, and mixed
- `type` Type of production, conventional or organic, a factor with levels conv orga
- `age` Age of the farmer
Details

Data is a mix of accounting data and production controls.

Note

Data as .csv are loaded by the command data using read.table(...,header = TRUE, sep = ",") such that this file is a semicolon separated file and not a comma separated file.

Source

Collected from farmers.

Examples

data(projekt)
## maybe str(projekt) ; plot(projekt) ...

\[
\text{sdea} \hspace{1cm} \text{Super efficiency}
\]

Description

The method sdea calculates super-efficiency and returns the same class of object as dea.

Usage

sdea(X, Y, RTS = "vrs", ORIENTATION = "in", DIRECT = NULL, param = NULL,
TRANPOSE = FALSE, LP = FALSE)

Arguments

X 
Inputs of firms to be evaluated, a K x m matrix of observations of K firms with m inputs (firm x input). In case TRANSPOSE=TRUE the input matrix is transposed to input x firm.

Y 
Outputs of firms to be evaluated, a K x n matrix of observations of K firms with n outputs (firm x input). In case TRANSPOSE=TRUE the output matrix is transposed to output x firm.

RTS 
Text string or a number defining the underlying DEA technology / returns to scale assumption; the same values as for dea.

0 fdh Free disposability hull, no convexity assumption
1 vrs Variable returns to scale, convexity and free disposability
2 drs Decreasing returns to scale, convexity, down-scaling and free disposability
3 crs Constant returns to scale, convexity and free disposability
4 irs Increasing returns to scale, (up-scaling, but not down-scaling), convexity and free disposability
5 irs2 Increasing returns to scale (up-scaling, but not down-scaling), additivity, and free disposability
6 add Additivity (scaling up and down, but only with integers), and free disposability
7 fdh+ A combination of free disposability and restricted or local constant return to scale
**ORIENTATION**  
Input efficiency "in" (1), output efficiency "out" (2), and graph efficiency "graph" (3). For use with DIRECT, an additional option is "in-out" (0).

**DIRECT**  
Directional efficiency, DIRECT is either a scalar, an array, or a matrix with non-negative elements.
- If the argument is a scalar, the direction is (1,1,...,1) times the scalar; the value of the efficiency depends on the scalar as well as on the unit of measurements.
- If the argument an array, this is used for the direction for every firm; the length of the array must correspond to the number of inputs and/or outputs depending on the ORIENTATION.
- If the argument is a matrix then different directions are used for each firm. The dimensions depends on the ORIENTATION, the number of firms must correspond to the number of firms in X and Y.

**param**  
Argument is at present only used when RTS="fdh+", see dea for a description.

**TRANSPOSE**  
See the description in dea.

**LP**  
Only for debugging, see the description in dea.

**Details**  
Super-efficiency measures are constructed by avoiding that the evaluated firm can help span the technology; ie. if the firm in question is a firm on the frontier in a normal dea approach then this firm in super efficiency might be outside the technology set.

**Value**  
The object returned is a Farrell object with the component described in dea. The relevant components are

- **eff**  
The efficiencies. Note when DIRECT is used then the efficiencies are not Farrell efficiencies but rather excess values in DIRECT units of measurement

- **lambda**  
The lambdas, i.e. the weight of the peers, for each firm

- **objval**  
The objective value as returned from the LP program; normally the same as eff.

- **RTS**  
The return to scale assumption as in the option RTS in the call

- **ORIENTATION**  
The efficiency orientation as in the call

**Note**  
Calculation of slacks for super efficiency should be done by using the option SLACK=TRUE in the call of the method sdea. If the two phases are done in two steps as first a call to sdea and then a call to slacks the user must make sure to set the reference technology to the one corresponding to super-efficiency in the call to slacks and this requires a loop with calls to slacks.

**Author(s)**  
Peter Bogetoft and Lars Otto <larsot23@gmail.com>
References

Peter Bogetoft and Lars Otto; Benchmarking with DEA, SFA, and R; Springer 2011. Sect. 5.2 page 115


See Also
dea

Examples

```r
x <- matrix(c(100, 200, 300, 500, 100, 200, 600), ncol=1)
y <- matrix(c(75, 100, 300, 400, 25, 50, 400), ncol=1)
se <- sdea(x, y)
se

# Leave out firm 3 as a determining firm of the technology set
n <- 3
dea.plot.frontier(x[-n], y[-n], txt=(1:dim(x)[1])[-n])
# Plot and label firm 3
points(x[n], y[n], cex=1.25, pch=16)
text(x[n], y[n], n, adj=c(-.75,.75))
```

sfa

Stochastic frontier estimation

Description

Estimate a stochastic frontier production or cost function using a maximum likelihood method.

Usage

```r
sfa(x, y, beta0 = NULL, lambda0 = 1, resfun = ebeta,
    TRANSPOSE = FALSE, DEBUG=FALSE,
    control=list(), hessian=2)

sfa.cost(W, Y, COST, beta0 = NULL, lambda0 = 1, resfun = ebeta,
    TRANSPOSE = FALSE, DEBUG=FALSE,
    control=list(), hessian=2)
```

te.sfa(object)
teBC.sfa(object)
teMode.sfa(object)
teJ.sfa(object)
Arguments

- **x**: Input as a K x m matrix of observations on m inputs from K firms; (firm x input); MUST be a matrix. No constant for the intercept should be included in x as it is added by default.
- **y**: Output; K times 1 matrix (one output)
- **Y**: Output; K times n matrix for m outputs; only to be used in cost function estimation.
- **W**: Input prices as a K x m matrix.
- **COST**: Cost as a K array for the K firms
- **beta0**: Optional initial parameter values
- **lambda0**: Optional initial ratio of variances
- **resfun**: Function to calculate the residuals, default is a linear model with an intercept. Must be called as resfun(x,y,parm) where parm=c(beta,lambda) or parm=c(beta), and return the residuals as an array of length corresponding to the length of output y.
- **TRANSPOSE**: If TRUE, data is transposed, i.e. input is now m x K matrix
- **DEBUG**: Set to TRUE to get various debugging information written on the console
- **control**: List of control parameters to ucminf
- **hessian**: How the Hessian is delivered, see the ucminf documentation
- **object**: Object of class ‘sfa’ as output from the function sfa

Details

The optimization is done by the R method ucminf from the package with the same name. The efficiency terms are assumed to be half–normal distributed.

Changing the maximum steplength, the trust region, might be important, and this can be done by the option 'control=list(stepmax=0.1)'. The default value is 0.1 and that value is suitable for parameters around 1; for smaller parameters a lower value should be used. Notice that the step length is updated by the optimizing program and thus must be set for every call of the function sfa if it is to be set.

The generic functions print.sfa, summary.sfa, fitted.sfa, residuals.sfa, logLik.sfa, and coef.sfa all work as expected.

The methods te.sfa, teMode.sfa etc. calculates the efficiency corresponding to different methods
Value

The values returned from sfa is the same as for ucminf, i.e. a list with components plus some especially relevant for sfa:

- **par**: The best set of parameters found \(c(\beta, \lambda)\).
- **value**: The value of minus log-likelihood function corresponding to 'par'.
- **beta**: The parameters for the function
- **sigma2**: The estimate of the total variance
- **lambda**: The estimate of \(\lambda\)
- **N**: The number of observations
- **df**: The degrees of freedom for the model
- **residuals**: The residuals as a \(K\times 1\) matrix/vector, can also be obtained by
  \(\text{residuals(sfa-object)}\)
- **fitted.values**: Fitted values
- **vcov**: The variance-covariance matrix for all estimated parameters incl. \(\lambda\)
- **convergence**: An integer code. '0' indicates successful convergence. Some of the error codes taken from ucminf are
  - '1' Stopped by small gradient (grtol).
  - '2' Stopped by small step (xtol).
  - '3' Stopped by function evaluation limit (maxeval).
  - '4' Stopped by zero step from line search
  More codes are found in \texttt{ucminf}
- **message**: A character string giving any additional information returned by the optimizer, or 'NULL'.
- **o**: The object returned by \texttt{ucminf}, for further information on this see \texttt{ucminf}

Note

Calculation of technical efficiencies for each unit can be done by the method \texttt{te.sfa} as shown in the examples.

- \texttt{te.sfa(sfaObject), teBC.sfa(sfaObject)}: Efficiencies estimated by minimizing the mean square error; Eq. (7.21) in Bogetoft and Otto (2011, 219) and Battese and Coelli (1988, 392)
- \texttt{teJ.sfa(sfaObject), te2.sfa(sfaObject)}: Efficiencies estimates using the conditional mean approach Jondrow et al. (1982, 235).
- \texttt{te.add.sfa(sfaObject)}: Efficiency in the additive model, Bogetoft and Otto (2011, 219)

The variance of the distribution of efficiency can be calculated by \texttt{sigma2u.sfa(sfaObject)}, the variance of the random error by \texttt{sigma2v.sfa(sfaObject)}, and the total variance (sum of variances of efficiency and random noise) by \texttt{sigma2.sfa}.

The ratio of variances of the efficiency and the random noise can be found from the method \texttt{lambda.sfa}

The generic method \texttt{summary} prints the parameters, standard errors, t-values, and a few more statistics from the optimization.
**slack**

Calculate slack in an efficiency analysis

---

**Author(s)**

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

**References**

Bogetoft and Otto; *Benchmarking with DEA, SFA, and R*, Springer 2011; chapters 7 and 8.

**See Also**

See the method `ucminf` for the possible optimization methods and further options to use in the `option control`.

The method `sfa` in the package `frontier` gives another way to estimate stochastic production functions.

**Examples**

```r
# Example from the book by Coelli et al.
# d <- read.csv("c:/0work/rpack/front41Data.csv", header = TRUE, sep = ",")
# x <- cbind(log(d$capital), log(d$labour))
# y <- matrix(log(d$output))

n <- 50
x1 <- 1:50 + rnorm(50,0,10)
x2 <- 100 + rnorm(50,0,10)
x <- cbind(x1,x2)
y <- 0.5 + 1.5*x1 + 2*x2 + rnorm(n,0,1) - pmax(0,rnorm(n,0,1))
sfa(x,y)
summary(sfa(x,y))

# Estimate efficiency for each unit
o <- sfa(x,y)
eff(o)

te <- te.sfa(o)
tem <- teMode.sfa(o)
teJ <- teJ.sfa(o)
cbind(eff(o),te,Mode=eff(o, type="Mode"),tem,teJ)[1:10,]

sigma2.sfa(o)  # Estimated varians
lambda.sfa(o)   # Estimated lambda
```

---

**slack**

Slacks are calculated after taking the efficiency into consideration.
Usage

```r
slack(X, Y, e, XREF = NULL, YREF = NULL, FRONT.IDX = NULL, LP = FALSE)
```

Arguments

- **X**
  - Inputs of firms to be evaluated, a $K \times m$ matrix of observations of $K$ firms with $m$ inputs (firm x input).

- **Y**
  - Outputs of firms to be evaluated, a $K \times n$ matrix of observations of $K$ firms with $n$ outputs (firm x input).

- **e**
  - A Farrell object as returned from `dea` et al.

- **XREF**
  - Inputs of the firms determining the technology, defaults to $X$.

- **YREF**
  - Outputs of the firms determining the technology, defaults to $Y$.

- **FRONT.IDX**
  - Index for firms determining the technology.

- **LP**
  - Set `TRUE` for debugging.

Details

Slacks are calculated in a LP problem where the sum of all slacks are maximized after correction for efficiency. The for calculating slacks for orientation graph is low because of the low precision in the calculated graph efficiency.

Value

The result is returned as the Farrell object used as the argument in the call of the function with the following added components:

- **slack**
  - A logical vector where the component for a firm is `TRUE` if the sums of slacks for the corresponding firm is positive. Only calculated in `dea` when option `SLACK=TRUE`.

- **sum**
  - A vector with sums of the slacks for each firm. Only calculated in `dea` when option `SLACK=TRUE`.

- **sx**
  - A matrix for input slacks for each firm, only calculated if the option `SLACK` is `TRUE` or returned from the method `slack`.

- **sy**
  - A matrix for output slack, see `sx`.

Author(s)

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

References

Peter Bogetoft and Lars Otto; *Benchmarking with DEA, SFA, and R*; Springer 2011. Sect. 5.6 page 127.

Examples

```r
x <- matrix(c(100,200,300,500,100,200,600),ncol=1)
y <- matrix(c(75,100,300,400,25,50,400),ncol=1)
dea.plot.frontier(x,y,txt=1:dim(x)[1])

e <- dea(x,y)
eff(e)

# calculate slacks
sl <- slack(x,y,e)
data.frame(e$eff,sl$slack,sl$sx,sl$sy)
```

---

### typeIerror

**Probability of type I error for test in a bootstrap DEA model**

**Description**

Calculates the probability of a type I error for a test in bootstrapped DEA models.

**Usage**

```r
typeIerror(shat,s)
```

**Arguments**

- `shat`: The value of the statistic for which the probability of a type I error is to be calculated
- `s`: Vector with calculated values of the statistic for each of the NREP bootstraps; NREP is from `dea.boot`

**Details**

Needs bootstrapped values of the test statistic

**Value**

Returns the probability of a type I error

**Author(s)**

Peter Bogetoft and Lars Otto <larsot23@gmail.com>

**See Also**

Examples

# Probability of getting something larger than 1.96 in 10000 random
# standard normal variates.
x <- rnorm(10000)
typeIerror(1.96,x)
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