Package 'kyotil'

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```
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LazyData yes
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      doParallel, Exact, survey, magick
Description
      Helper functions for creating formatted summary of regression models, writing publication-
      ready tables to latex files, and running Monte Carlo experiments.
VignetteBuilder R.rsp
License GPL (>= 2)
NeedsCompilation yes
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```

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Description

Index

Calculate age, by Jason P Becker, modified very slightly in how arguments are passed to the function.

```
age_calc(dob, enddate = Sys.Date(), units = c("days", "months", "years"), precise = TRUE)
```

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Arguments

dob POSIXIt or Date. Birthday

enddate POSIXIt or Date. Date to compute age

units string. Choose a unit.

precise Boolean.

Author(s)

Jason P Becker

References

http://blog.jsonbecker.com/2013/12/calculating-age-with-precision-in-r.html

Examples

```
age_calc (dob=strptime("290CT2002", format="%d%b%Y"),
    enddate=strptime("300CT2003", format="%d%b%Y"), units='years', precise=TRUE)
age_calc (dob=strptime("290CT2002", format="%d%b%Y"),
    enddate=strptime("30DEC2003", format="%d%b%Y"), units='years', precise=FALSE)
```

auc

AUC

Description

AUC methods.

```
## S3 method for class 'auc'
coef(object, ...)
## S3 method for class 'auc'
predict(object, newdata, case.percentage = NULL, ...)
## S3 method for class 'auc'
print(x, ...)
## S3 method for class 'auc'
summary(object, ...)
## S3 method for class 'auc'
trainauc(fit, training.data = NULL, ...)
## S3 method for class 'auc'
ratio(fit)
## S3 method for class 'glm'
trainauc(fit, ...)
## S3 method for class 'glm'
ratio(fit)
```

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Arguments

fit an object that inherits from class 'auc' such as 'rauc' or 'sauc' object an object that inherits from class 'auc' such as 'rauc' or 'sauc' x an object that inherits from class 'auc' such as rauc, sauc or sauc.dca.

newdata data at which to predict

case.percentage

used for class prediction, defaults to NULL

training.data data frame used to compute auc based on a fit obtained by a call to rauc, sauc

or sauc.dca

... arguments passed to or from methods

Author(s)

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Krisztian Sebestyen <>

base.functions

Some Base Functions

Description

cbinduneven binds together a list of matrixes/dataframes of different lengths, rows are matched by names binary returns binary representation of an integer. binary2 returns binary representation of an integer with leading 0, the length of string is n. mysystem can call any exe file that is in the PATH f2c convert temperature from f to c/

```
mytable (..., exclude = if (useNA == "no") c(NA, NaN), useNA = "ifany",
    dnn = list.names(...), deparse.level = 1)

cbinduneven(li)

binary(i)

multi.outer (f, ...)

myreshapelong(dat, cols.to.be.stacked, label.cols.to.be.stacked, new.col.name)
binary2(i, n)

f2c(f)

ftoi(f)
```

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```
keepWarnings(expr)
meanmed(x, na.rm = FALSE)
myaggregate(x, by, FUN, new.col.name = "aggregate.value", ...)
myreshapewide(formula, dat, idvar, keep.extra.col=FALSE)
mysapply(X, FUN, ..., simplify = TRUE, USE.NAMES = TRUE, ret.mat = TRUE)
myscale(x)
mysystem(cmd, ...)
mytapply(X, INDEX, FUN = NULL, ..., simplify = TRUE)
read.sv(file, header = TRUE, ...)
read.tsv(file, header = TRUE, sep = "\t", ...)
table.prop(x,y=NULL,digit=1,style=2,whole.table.add.to.1=FALSE,useNA="ifany",
   add.perc=FALSE, add.total.column = FALSE)
table.cases (case,group,include.all=TRUE,desc="cases")
table.cases.3(case,group1,group2)
unix()
mycor (x, use = "everything", method = c("pearson", "kendall", "spearman"),
   alternative = c("two.sided", "less", "greater"), exact = NULL,
   conf.level = 0.95, continuity = FALSE,
   digits.coef=2, digits.pval=3,
    ...)
```

Arguments

```
exclude exclude

dnn dnn

deparse.level deparse.level

add.total.column
tbdi

use tbdi

method tbdi

alternative tbdi
```

base.functions

exact tbdi conf.level tbdi continuity tbdi digits.coef tbdi digits.pval tbdi cols.to.be.stacked tbdi label.cols.to.be.stacked tbdi li a list i tbdi tbdn n f In multi.out, f is a function. vector of 0/1 case vector of multi-group indicators group a formula object. formula tbdexpr expr tbdxХ tbdna.rm na.rm desc tbdby tbdby by whole.table.add.to.1 Boolean tbdnew.col.name new.col.name tbd... . . . tbddat dat tbdidvar idvar tbdXΧ simplify tbdsimplify USE.NAMES tbdUSE.NAMES ret.mat tbdret.mat cmdtbdcmd INDEX tbdINDEX file tbdfile tbdheader header sep tbdsep tbdy

tbddigit

digit

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```
style tbdstyle
FUN tbdFUN
keep.extra.col tbdFUN
useNA tbdFUN
add.perc tbdFUN
include.all tbdFUN
group1 tbdFUN
group2 tbdFUN
```

Examples

binaryloess

Using loess to Check Functional Form for Logistic Regression

Description

This function plots a smoothed line of how the average value of Y changes with X in order to check functional form for logistic regression.

```
binaryloess(x, y, scale = c("logit", "linear"), span = 0.7, weights = NULL, ...)
```

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Arguments

Х	tbdx
у	tbdy
scale	tbdscale
span	smoothing parameter, passed to loess. If less than 1, the neighbourhood includes proportion a of the points. If greater than 1, all points are used, with the maximum distance assumed to be a^(1/p) times the actual maximum distance for p explanatory variables. Missing records are removed first.
weights	sampling weights, passed to loess
	passed to plotting function

Details

This function comes from Jonathan Bartlett ()https://thestatsgeek.com/2014/09/13/checking-functional-form-in-logistic-regression-using-loess/).

Examples

```
set.seed(1234)
n <- 1000
x <- rnorm(n)
xb <- -2+x
pr <- exp(xb)/(1+exp(xb))
y=rbern(n, pr)

par(mfrow=c(1,2))
binaryloess(x, y, scale = "logit", span = 0.7, weights = NULL, ylab="logit(p)")
binaryloess(x, y, scale = "linear", span = 0.7, weights = NULL, ylab="prob")</pre>
```

cox.zph.2 Test the Proportional Hazards Assumption of a Cox Regression (a slightly modified version)

Description

A slightly modified test of the proportional hazards assumption for a Cox regression model fit (coxph). This version corrects some conservativeness of the test.

```
cox.zph.2(fit, transform = "km", global = TRUE, exact=TRUE)
```

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Arguments

fit fit transform transform

global global

exact Boolean. If FALSE, this function is an identical copy of cox.zph. If TRUE, it

computes the variance of the test statistic exactly, instead of approximately.

Details

When the model uses time-dependent covariates, the approximation used in Grambsch and Therneau resulted in conservativeness of the test. This is "fixed" here at a cost of up to 2.5 times longer execution time.

References

Fong, Y. and Halloran, M Elizabeth and Gilbert, P. Using Time-Dependent Age Group in Cox Regression Analysis of Vaccine Efficacy Trials, Just Another Epi Journal, in prep.

Examples

crossvalidation

Cross Validation Functions

Description

Cross validation utility functions.

```
sample.for.cv (dat, v, seed)
get.kfold.splits (dat, k, seed)
kfold.split (k, n1, n0)
ran.kfold.split(k, n1, n0, replicates)
lpo.split(n1, n0)
get.splits (dat, cv.scheme=c("LPO","5fold","50xrandom4:1"), seed)
```

Deming Deming

Arguments

dat	a data frame. One of the columns must be named y and y should be $0/1$ with 1 for case and 0 for control
V	v-fold cross validation
seed	seed for random number generators
k	var.equal
n1	var.equal
n0	var.equal
replicates	var.equal
cv.scheme	var.equal

Details

sample.for.cv: case and controls are sampled separately.

Value

sample.for.cv returns a list of two vector of integers: train and test, which refer to the rows of dat

Deming	Fit Deming regression.

Description

Deming regression fit. Assume x and y variances are the same. Slightly modified from MethComp R package.

Usage

```
Deming(x, y, vr = sdr^2, sdr = sqrt(vr), boot = TRUE, keep.boot = FALSE, alpha = 0.05)
```

Arguments

X	tbd
у	tbdy
vr	tbdvr
sdr	tbdsdr
boot	tbdboot
keep.boot	tbdkeep.boot
alpha	tbdalpha

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Examples

```
## Not run:
set.seed(1)
x=rnorm(100,0,1)
y=x+rnorm(100,0,.5)
x=x+rnorm(100,0,.5)
fit=Deming(x,y, boot=TRUE)
summary(fit)
plot(x,y)
abline(fit)
# compare with lm fit
fit.1=lm(y~x, data.frame(x,y))
summary(fit.1)
abline(fit.1, col=2)
## End(Not run)
```

DMHeatMap

Better Heatmap Function

Description

Makes a heatmap representation of correaltion coefficients easier.

```
DMHeatMap(x, Rowv = TRUE, Colv = if (symm) "Rowv" else TRUE,
distfun = dist, hclustfun = hclust, dendrogram =
c("both", "row", "column", "none"), symm = FALSE,
scale = c("none", "row", "column"), na.rm = TRUE, revC
= identical(Colv, "Rowv"), add.expr, breaks, symbreaks
= min(x < 0, na.rm = TRUE) || scale != "none", col =
"heat.colors", colsep, rowsep, sepcolor = "white",
sepwidth = c(0.05, 0.05), cellnote, notecex = 1,
notecol = "cyan", na.color = par("bg"), trace =
c("column", "row", "both", "none"), tracecol = "cyan",
hline = median(breaks), vline = median(breaks),
linecol = tracecol, margins = c(5, 5), ColSideColors,
RowSideColors, cexRow = 0.2 + 1/log10(nr), cexCol =
0.2 + 1/log10(nc), labRow = NULL, labCol = NULL,
labColor = NULL, axis = TRUE, heatmapOnly = FALSE, key
= TRUE, keysize = 1.5, density.info = c("histogram",
"density", "none"), denscol = tracecol, symkey = min(x
< 0, na.rm = TRUE) || symbreaks, densadj = 0.25, main
= NULL, xlab = NULL, ylab = NULL, lmat = NULL, lhei =
NULL, lwid = NULL, lower.left.only = TRUE, legend =
TRUE, legend.x = "topright", verbose = FALSE, ...)
```

DMHeatMap

Arguments

x tbd
axis tbd
heatmapOnly tbd
verbose tbd
legend.x tbd
legend tbd

Rowv tbdRowv
Colv tbdColv
distfun tbddistfun
hclustfun tbdhclustfun
dendrogram tbddendrogram

symm tbdsymm
scale tbdscale
na.rm tbdna.rm
revC tbdrevC
add.expr tbdadd.expr
breaks tbdsymbreaks

col tbdcol colsep tbdcolsep rowsep tbdrowsep sepcolor tbdsepcolor sepwidth tbdsepwidth cellnote tbdcellnote notecex tbdnotecex notecol tbdnotecol na.color tbdna.color trace tbdtrace tracecol tbdtracecol tbdhline hline vline tbdvline linecol tbdlinecol margins tbdmargins

ColSideColors tbdColSideColors RowSideColors tbdRowSideColors

cexRow tbdcexRow

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cexCol	tbdcexCol
labRow	tbdlabRow
labCol	tbdlabCol
labColor	tbdlabColor
key	tbdkey
keysize	tbdkeysize
density.info	tbddensity.info
denscol	tbddenscol
symkey	tbdsymkey
densadj	tbddensadj
main	tbdmain
xlab	tbdxlab
ylab	tbdylab
lmat	tbdlmat
lhei	tbdlhei
lwid	tbdlwid
lower.left.only	y
	tbdlower.left.only
	tbd

Examples

```
cor=matrix(runif(15),5,3)
breaks=c(-1,-.7,-.5,-.3,-.1,.1,.3,.5,.7,1)
hU=DMHeatMap(cor, trace="none", symm=FALSE,dendrogram="none", col=RColorBrewer::brewer.pal(
    length(breaks)-1,"RdYlGn"), distfun = function(c) as.dist(1 - c), cexRow =1.5, cexCol =1.5,
    lmat=rbind(c(2, 1), c(4,3)), lhei=c(4, 1), breaks=breaks, margins=c(2,2), key = FALSE,
    Rowv=NA, lower.left.only=FALSE)
```

get.sim.res

Read simulation results

Description

Go through a folder and read all files and combine the results into a multidimensional array.

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Usage

Arguments

dir directory of MC result files partial path to the directory of MC result files path res.name name of the R object saved in the files, default is res, but may be others verbose Boolean a string to denote simulation setting sim nn a vector of sample sizes fit.method a string to denote fitting method. sim, nn and fit.method together forms the name of the directory containing MC result files exclude.col column number exclude.some whether to exclude MC results that are extreme coef.0 simulation truth digit1 digits sum.est use mean or median as location estimate summary use mean or median as sd estimate summary sum.sd style integer

Details

Depends on package abind to combine arrays from files.

keep.intercept whether to include intercept in the table

Value

A multidimensional array.

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|--|

Description

getK calculates the kernel matrix between X and itself and returns a n by n matrix. Alternatively, it calculates the kernel matrix between X and X2 and returns a n by n2 matrix.

Usage

```
getK (X,kernel,para=NULL,X2=NULL,C = NULL)
```

Arguments

X covariate matrix with dimension n by d. Note this is not the paired difference of

covariate matrix.

kernel string specifying type of kernel: polynomial or p $(1 + \langle x,y \rangle)^{\wedge}$ para,

rbf or r exp(-para* $||x-y||^2$),

linear or 1 < x,y>,

ibs or i 0.5*mean(2.0 - |x-y|) or sum(w*(2.0 - |x-y|))/sum(w), with x[i],y[i] in

 $\{0,1,2\}$ and weights 'w' given in 'para'.

hamming or h for sum(x == y) with x[i],y[i] binary,

no default.

para parameter of the kernel fucntion. for ibs or hamming, para can be a vector of

weights.

X2 optional second covariate matrix with dimension n2 by d

C logical. If TRUE, kernels are computed by custom routines in C, which may be

more memory efficient, and faster too for ibs and hamming kernels.

Details

IBS stands for 'Identical By State'. If 'x','y' are in in {0,1,2} then

 $IBS(x,y) = 0 \text{ if } |x-y| = 2, \ 1 \text{ if } |x-y| = 1, \ 2 \text{ if } |x-y| = 0, \text{ or } IBS(x,y) = 2.0 - |x-y|.$

K(u,v) = sum(IBS(u[i],v[i])) / 2K where K = length(u).

The 'hamming' kernel is the equivalent of the 'ibs' kernel for binary data. Note that 'hamming' kernel is based on hamming similarity(!), not on dissimilarity distance.

Within in the code, C is default to TRUE for ibs and hamming kernels and FALSE otherwise.

Value

A kernel matrix.

Author(s)

```
Youyi Fong <youyifong@gmail.com>
Krisztian Sebestyen <ksebestyen@gmail.com>
Shuxin Yin <>
```

Examples

```
X = cbind(x1=rnorm(n=5), x2=rnorm(n=5))
X2 = cbind(x1=rnorm(n=3), x2=rnorm(n=3))
dim(X2)
K = getK(X,"linear")
dim(K)
K = getK(X,"linear",X2=X2)
dim(K)
K1 = getK(X2,"1",X2=X)
dim(K1)
all(K==t(K1))
# RBF kernel
K = getK(X,"rbf",para=1,X2=X2)
K1 = getK(X2,"r",para=1,X2=X)
all(K==t(K1))
# IBS kernel for ternary data
X <- as.matrix(expand.grid(0:2,0:2))</pre>
K = getK(X,kernel = 'ibs')
# add weight
w = runif(ncol(X))
K = getK(X,kernel = 'ibs',para = w)
# IBS kernel for binary data via option 'h' for 'hamming similarity measure'
X <- as.matrix(expand.grid(0:1,0:1))</pre>
K=getK(X,kernel = 'h')
```

```
get_count_from_xy_coor
```

Imaging analysis for spatial region

Description

Counting the number of masks in a rectangular region

```
get_count_from_xy_coor(file, topleft, bottomright, image, plot)
```

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Arguments

file _sizes_coordinates.txt

topleft topleft (x,y) coordiate for a rectangular box

bottomright bottomright (x,y) coordiate for a rectangular box

image image: an image for plotting

plot plot=TRUE shows image with rectangular box

Details

This function counts cells inside of rectangular box made by the topleft and bottomright xy-coordinates.

Value

The number of masks inside of the rectangular box

Author(s)

Sunwoo Han

Examples

```
#get_count_from_xy_coor(file='M926910_Position1_CD3-BUV395_sizes_coordinates.txt',
    #topleft=c(500,0), bottomright=c(1392,500),
    #image='M926910_Position1_CD3-BUV395.tiff', plot=TRUE)
```

iorw

Causal Mediation Analysis of Cowling et al.

Description

Estimate the total, direct, and indirect effects using IORW method (inverse odds ratio weighting) and compute 95

```
iorw(formula.effect, formula.mediators, data, family =
  NULL, nboot = 10000, numCores = 1, save.steps = FALSE,
  verbose = FALSE)

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

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Arguments

formula.effect a formula object for the total and direct effect regression. The first term on the right is assumed to be the binary treatment/exposure variable.

formula.mediators

a formula object for logistic regression. It should be of the form: ~ mediation

marker1 + mediation marker2.

data a data frame.

family if Cox regression, leave as NULL; otherwise, it will be passed to glm().

nboot an integer. Number of bootstrap replicates.

numCores an interger. Number of cores to use for parallel processing.

save. steps boolean. Whether or not to save the fits from the three steps and the weights.

x Object of type iorw

verbose boolean.

. . . Additional arguments passed to the print function.

Details

Code by Cowling and Lim was downloaded from https://datadryad.org/stash/dataset/doi:10.5061/dryad.cv37539 If a bootstrap replicate generates warnings during regression, NA will be returned for that replicate.

The number of such occurrences is recorded in an attribute of boot.perc in the return value.

It does not handle sampling weights yet.

Value

Point estimates and percentile bootstrap confidence intervals.

Author(s)

Youyi Fong, based on code by Cowling and Lim

References

Cowling, B. J., Lim, W. W., Perera, R. A., Fang, V. J., Leung, G. M., Peiris, J. M., & Tchetgen Tchetgen, E. J. (2019). Influenza hemagglutination-inhibition antibody titer as a mediator of vaccine-induced protection for influenza B. Clinical Infectious Diseases, 68(10), 1713-1717.

Nguyen, Q. C., Osypuk, T. L., Schmidt, N. M., Glymour, M. M., & Tchetgen Tchetgen, E. J. (2015). Practical guidance for conducting mediation analysis with multiple mediators using inverse odds ratio weighting. American journal of epidemiology, 181(5), 349-356.

Tchetgen Tchetgen, E. J. (2013). Inverse odds ratio-weighted estimation for causal mediation analysis. Statistics in medicine, 32(26), 4567-4580.

Imai, K., Keele, L., & Tingley, D. (2010). A general approach to causal mediation analysis. Psychological methods, 15(4), 309.

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Examples

```
#### Cox regression
# without adjusting for baseline markers
library(survival)
formula.effect=Surv(surv_time, flu)~vaccine+age
formula.mediators=~log2(postvax.B.Brisbane/5)
res.1=iorw(formula.effect, formula.mediators, kid, nboot=10, numCores=1); res.1
stopifnot(\max(abs(res.1\$boot[1,]-c(0.2029779,0.6070105,0.3039110,0.4283389,0.2124268))) < 1e-6)
# adjust for baseline markers
formula.effect=Surv(surv_time, flu)~vaccine+log2(prevax.B.Brisbane)+age
formula.mediators=~log2(postvax.B.Brisbane/5)
res.2=iorw(formula.effect, formula.mediators, kid, nboot=10, numCores=1); res.2
#### Logistic regression
# without adjusting for baseline markers
formula.effect=flu~vaccine+age
formula.mediators=~log2(postvax.B.Brisbane/5)
res.3=iorw(formula.effect, formula.mediators, kid, family=binomial(), nboot=10, numCores=1); res.3
stopifnot(max(abs(res.3$boot[1,] - c(0.1960024,0.6154349,0.2937164,0.4145470,0.2168644)))<1e-6)
# adjust for baseline markers
formula.effect=flu~vaccine+log2(prevax.B.Brisbane)+age
formula.mediators=~log2(postvax.B.Brisbane/5)
res.4=iorw(formula.effect, formula.mediators, kid, family=binomial(), nboot=10, numCores=1); res.4
```

kid

Dataset from Cowling et al.

Description

Influenza immune response biomarkers dataset.

```
data("kid")
```

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Format

A data frame with 736 observations on the following 10 variables.

```
hhID a numeric vector
age a numeric vector
intervention a character vector
vaccine a numeric vector
vaccine.date a Date
postvax.date a Date
prevax.B.Brisbane a numeric vector
postvax.B.Brisbane a numeric vector
surv_time a numeric vector
flu a numeric vector
```

References

Cowling, B. J., Lim, W. W., Perera, R. A., Fang, V. J., Leung, G. M., Peiris, J. M., & Tchetgen Tchetgen, E. J. (2019). Influenza hemagglutination-inhibition antibody titer as a mediator of vaccine-induced protection for influenza B. Clinical Infectious Diseases, 68(10), 1713-1717.

kyotil kyotil

Description

Utility functions by Youyi Fong and Krisz Sebestyen, and some functions copied from other packages for convenience (acknowledged on their manual pages).

Most useful functions: mypostscript/mypdf, mytex,

See the Index link below for a list of available functions.

The package depends on Hmisc. The main reason for that, besides the usefulness of the package, is Hmisc depends on ggplot2, which also define

make.timedep.dataset 21

make.timedep.dataset Create Dataset for Time-dependent Covariate Proportional Hazard Model Analaysi

Description

Returns a data frame that is suitable for time-dependent covariate Cox model fit.

Usage

```
make.timedep.dataset(dat, X, d, baseline.ageyrs, t.1, t.2 = NULL)
```

Arguments

data frame

X string. Name of the followup time column in dat. Unit needs to be years.

d string. Name of the followup time column in dat.

baseline.ageyrs

string. Name of the followup time column in dat.

t.1 numerical. Cutoff for age group

t.2 numerical. Second cutoff for age group

Details

The function assumes that the followup length is such that only one change of age group is possible.

Value

Returns a data frame with the following columns added: tstart, tstop, .timedep.agegrp, .base-line.agegrp

tstart left bound of time interval tstop right bound of time interval

.timedep.agegrp

time-dependent age group

.baseline.agegrp

baseline age group

Author(s)

Youyi Fong

References

Therneau, T. and Crowson, C. Using Time Dependent Covariates and Time Dependent Coefficients in the Cox Model. A vignette from the R package surival.

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Examples

```
library(survival)
n=3000; followup.length=5; incidence.density=0.015; age.sim="continuous"

dat.0=sim.dat.tvarying.two(n, followup.length, incidence.density, age.sim, seed=1)
dat=subset(dat.0, for.non.tvarying.ana, select=c(ptid, X, d, baseline.age, trt))
dat.timedep = make.timedep.dataset (dat, "X", "d", "baseline.age", 6)
coxph(Surv(tstart,tstop,d) ~ trt*.timedep.agegrp, dat.timedep)
```

math.functions

Math Functions

Description

H calculates entropy.

Usage

```
as.binary(n, base = 2, r = FALSE)
binom.coef(n, m)
expit(x)
logDiffExp(logx1, logx2)
logit(x)
logMeanExp(logx, B = NULL)
logSumExp(logx)
logSumExpFor2(logx, logy)
permn(x, fun = NULL, ...)
Stirling2(n, m)
interpolate(pt1, pt2, x)
```

Arguments

n tbdn base tbdbase matrix.array.functions 23

```
r
                 tbdr
                 tbdm
m
pt1
                 a vector of length 2
                 a vector of length 2
pt2
                 tbdx
logx1
                 tbdlogx1
                 tbdlogx2
logx2
                 tbdlogx
logx
                 tbdB
                 tbdlogy
logy
fun
                 tbdfun
                 tbd...
```

Examples

```
H(rep(1/5,5))
H(rep(3,5))
```

matrix.array.functions

Matrix and Array Functions

Description

concatList returns a string that concatenates the elements of the input list or array

```
AR1(p, w)
concatList(lis, sep = "")
EXCH(p, rho)
fill.jagged.array(a)
getMidPoints(x)
getUpperRight(matri, func = NULL)
last(x, n = 1, ...)
mix(a, b)
```

24 matrix.array.functions

```
## S3 method for class 'data.frame'
rep(x, times = 1, ...)

## S3 method for class 'matrix'
rep(x, times = 1, each = 1, by.row = TRUE, ...)

## S3 method for class 'matrix.block'
rep(x, times = 2, ...)

shift.left(x, k = 1)

shift.right(x, k = 1)

thin.rows(dat, thin.factor = 10)

ThinRows(dat, thin.factor = 10)

tr(m)
```

Arguments

tbdp р tbdw W lis list or array tbdsep sep rho tbdrho tbda tbdx tbdmatri matri tbdfunc func tbdn tbd... . . . tbdb b times tbdtimes each tbdeach tbdby.row by.row k tbdk tbddat dat tbdthin.factor thin.factor tbdm m

Examples

```
concatList(1:3,"_")
```

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matrix2

Matrix Functions that May Be Faster than

Description

DXD computes D %*% X %*% D, where D is a diagonal matrix. tXDX computes t(X) %*% D %*% X. symprod computes S %*% X for symmetric S. txSy computes t(x) %*% S %*% y for symmetric S.

Usage

```
DXD(d1, X, d2)

tXDX(X,D)

symprod(S, X)

txSy(x, S, y)

.as.double(x, stripAttributes = FALSE)
```

Arguments

d1	a diagonal matrix or an array
d2	a diagonal matrix or an array
X	array
у	array
S	symmetric matrix
Χ	matix
D	matix
stripAttributes	
	boolean

Details

.as.double does not copying whereas as.double(x) for older versions of R when using .C(DUP = FALSE) make duplicate copy of x. In addition, even if x is a 'double', since x has attributes $(\dim(x))$ as.double(x) duplicates

The functions do not check whether S is symmetric. If it is not symmetric, then the result will be wrong. DXD offers a big gain, while symprod and txSy gains are more incremental.

Author(s)

Krisztian Sebestyen

26 misc

Examples

```
d1=1:3
d2=4:6
X=matrix(1:9,3,3)
all(DXD(d1, X, d2) == diag(d1) %*% X %*% diag(d2))
S=matrix(c(1,2,3,2,4,5,3,5,8),3,3)
X=matrix(1:9,3,3)
all( symprod(S, X) == S %*% X )

x=1:3
y=4:6
S=matrix(c(1,2,3,2,4,5,3,5,8),3,3)
txSy(x, S, y) == drop(t(x)%*%S%*%y)
```

misc

Misc Functions

Description

Misc functions. summ computes iterative sum, sort of like diff.

Usage

```
pava (x, wt = rep(1, length(x)))
summ(x)
empty2na(x)
## S3 method for class 'pcc'
predict(object, newdat, ...)
rank.inv.norm(x)
INT(x)
dec_to_binary (x,d)
```

Arguments

X	tbdx
d	number of digits in the returned binary representation, including leading 0 's
wt	tbdvar.equal
object	tbdvar.equal
newdat	tbdvar.equal
	tbdvar.equal

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Details

rank.inv.norm: rank-based inverse normal/gaussian transformation dec_to_binary covert a decimal number to a binary representation with d digits

Value

summ returns

p.adj.perm

Permutation-based Multitesting P Values Adjustment

Description

An implementation of Westfall and Young

Usage

```
p.adj.perm(p.unadj, p.perms, alpha = 0.05)
```

Arguments

Details

This implementation is not as fast as the implementation from the package multtest. But ususally the step to create p.perms is the rate-limiting step.

The smallest of the Westfall and Young FWER-controlling multitesting adjusted p values coincides with the p value for testing a global null without any assumptions. But for the multitesting adjustment to hold, it requires the subset pivotality condition.

Author(s)

Sue Li, sli@fredhutch.org

References

Westfall, P. H., & Young, S. S. (1993). Resampling-based multiple testing: Examples and methods for p-value adjustment (Vol. 279). John Wiley & Sons.

Westfall, P. H., & Troendle, J. F. (2008). Multiple testing with minimal assumptions. Biometrical Journal: Journal of Mathematical Methods in Biosciences, 50(5), 745-755.

plotting

Plotting Functions

Description

mypostscript and mypdf sets the width and height based on mfrow input.

```
smoothed.scaled.hist (dat.ls, bin_width, scale.factors=NULL, cols=NULL,
  legend=NULL, cex.legend=1, ...)
myplot (object, ...)
## S3 method for class 'loess'
myplot(object, xlab="x", ylab="fitted", ...)
whiskers (x, s, ...)
abline.pt.slope(pt1, slope, x2=NULL, ...)
abline.pts(pt1, pt2 = NULL)
butterfly.plot(dat, dat2 = NULL, add = FALSE, xaxislabels = rep("", 4), x.ori = 0,
    xlab = "", ylab = "", cex.axis = 1, ...)
empty.plot()
add.mtext.label (text, cex = 1.4, adj = -0.2)
mydev.off(file = "temp", ext = c("pdf"), res = 200, mydev =
                 NULL, silent = TRUE)
getMfrow(len)
myhist (x, add.norm=TRUE, col.norm="blue", ...)
myforestplot(dat, xlim = NULL, xlab = "", main = "", col.1 = "red",
 col.2 = "blue", plot.labels = TRUE, order = FALSE,
 decreasing = FALSE, vline = TRUE, cols = NULL, log =
 "", null.val = NULL)
my.interaction.plot(dat, x.ori = 0, xaxislabels = rep("", 2), cex.axis = 1, add = FALSE,
    xlab = "", ylab = "", pcol = NULL, lcol = NULL, ...)
myboxplot(object, ...)
## S3 method for class 'formula'
```

```
myboxplot(formula, data, cex = 0.5, xlab = "", ylab = NULL, main
                 = "", box = TRUE, at = NULL, na.action = NULL, p.val =
                 NULL, pch = 1, col = "white", col.points = 1, border =
                 1, test = "", friedman.test.formula = NULL,
                 reshape.formula = NULL, reshape.id = NULL, jitter =
                 TRUE, add.interaction = FALSE, drop.unused.levels =
                 TRUE, bg.pt = NULL, add = FALSE, seed = 1,
                 write.p.at.top = FALSE, ...)
## S3 method for class 'data.frame'
myboxplot(object, cex = 0.5, ylab = "", xlab = "", main = "",
box = TRUE, at = NULL, pch = 1, col = 1, test = "",
paired = FALSE, ...)
## S3 method for class 'list'
myboxplot(object, paired = FALSE, ...)
abline.shade.2(x, col=c(0,1,0))
abline.shade(pt, type = 5, col = c(0, 1, 0), alpha = 0.3)
mylegend(legend, x, y=NULL, lty = NULL, bty = "n", ...)
mymatplot(x, y, type = "b", lty = c(1, 2, 1, 2, 1, 2), pch =
                NULL, col = rep(c("darkgray", "black"), each = 3),
                 xlab = NULL, ylab = "", draw.x.axis = TRUE, bg = NA,
                 lwd = 1, at = NULL, make.legend = TRUE, legend = NULL,
                 impute.missing.for.line = TRUE, legend.x = 9,
                 legend.title = NULL, legend.cex = 1, legend.lty = lty,
                 legend.inset = 0, xaxt = "s", y.intersp = 1.5,
                 x.intersp = 0.3, text.width = NULL, add = FALSE, ...
)
mypairs(dat, ladder = FALSE, show.data.cloud = TRUE,
ladder.add.line = T, ladder.add.text = T, ...)
wtd.hist (x, breaks = "Sturges", freq = NULL, probability = !freq,
    include.lowest = TRUE, right = TRUE, density = NULL, angle = 45,
   col = NULL, border = NULL, main = paste("Histogram of", xname),
   xlim = range(breaks), ylim = NULL, xlab = xname, ylab, axes = TRUE,
   plot = TRUE, labels = FALSE, nclass = NULL, weight = NULL,
    ...)
mylines(x, y, type = "1", ...)
myfigure(mfrow = c(1, 1), mfcol = NULL, width = NULL,
   height = NULL, oma = NULL, mar = NULL, main.outer = FALSE, bg=NULL)
```

```
mypdf(...)
mypng(...)
mytiff(...)
mypostscript(file = "temp", mfrow = c(1, 1), mfcol = NULL, width =
                 NULL, height = NULL, ext = c("eps", "pdf", "png",
                 "tiff"), oma = NULL, mar = NULL, main.outer = FALSE,
                 save2file = TRUE, res = 200, silent = TRUE, ...)
panel.cor(x, y, digits = 2, prefix = "", cex.cor, cor., leading0
= FALSE, cex.cor.dep = TRUE, ...)
panel.hist(x, ...)
panel.nothing(x, ...)
corplot(object, ...)
## Default S3 method:
corplot(object, y, ...)
## S3 method for class 'formula'
corplot(formula, data, main = "", method = c("pearson",
                 "spearman"), col = 1, cex = 0.5, add.diagonal.line =
                 TRUE, add.lm.fit = FALSE, add.loess.fit = FALSE,
                 col.lm = 2, add.deming.fit = FALSE, col.deming = 4,
                 add = FALSE, digit.cor = 2, log = "", same.xylim =
                 FALSE, xlim = NULL, ylim = NULL, ...)
```

Arguments

digit.cor	number of digits to print correlation
col.points	color of points
dat.ls	named list of vectors. A histogram is made for each vector.
bin_width	width of bin for histograms
scale.factors	named vector of scale factors to scale the histogram counts by
cex.legend	cex for legend
silent	tbdadd
legend.lty	tbdadd
cex.cor.dep	tbdadd
add.loess.fit	tbdadd

```
leading0
                 tbdadd
null.val
                 tbdadd
write.p.at.top
                 tbdadd
text.width
                 tbdadd
                 tbdadd
text
                 tbdcex
cex
adj
                 tbdpt2
                 tbdfile
file
ext
                 tbdext
                 resolution.
res
                 Boolean, whether to add normal approximation density line
add.norm
                 string, color of added normal density line
col.norm
pt1
                 tbdpt1
                 tbdslope
s
ladder
                 tbdslope
slope
                 tbdslope
friedman.test.formula
                 tbdslope
reshape.id
                 tbdslope
impute.missing.for.line
                 tbdslope
                 tbdslope
cor.
mydev
                 tbdslope
                 Boolean
jitter
add.interaction
                 Boolean
                 tbd...
. . .
                 tbdpt2
xaxt
                 tbdpt2
breaks
                 tbdpt2
freq
bg.pt
                 tbdpt2
probability
                 tbdpt2
include.lowest \ tbdpt2
right
                 tbdpt2
density
                 tbdpt2
angle
                 tbdpt2
border
                 tbdpt2
                 tbdpt2
axes
```

plot tbdpt2 labels tbdpt2 nclass tbdpt2 weight tbdpt2pt2 tbdpt2 pt tbdpt2tbdpt2 alpha dat tbddat line width. lwd x.intersp

x.intersp controls the look of legend.y.intersp controls the look of legend.

legend.inset legend inset tbddat2 dat2 add tbdadd log log add.lm.fit lm fit add.deming.fit add col.lm col col.deming col reshape.formula

a formula object.

xaxislabels tbdxaxislabels

x.ori tbdx.ori
xlab tbdxlab
ylab tbdylab
cex.axis tbdcex.axis
len tbdlen

 $same.xylim \qquad \quad Boolean. \ Whether xlim \ and \ ylim \ should \ be \ the \ same$

tbdxlim xlim ylim tbdxlim main tbdmain col.1 tbdcol.1 tbdcol.2 col.2 pcol tbdpcol lcol tbdlcol object tbdobject formula tbdformula

data tbddata
box tbdbox
at tbdat
pch tbdpch
col tbdcol
test string. I

test string. For example, "t", "w", "f", "k", "tw"

legend tbdlegend x tbdx lty tbdlty bty tbdbty tbdtype

make.legend tbdmake.legend
legend.x tbdlegend.x
legend.title tbdlegend.title
legend.cex tbdlegend.cex
draw.x.axis tbddraw.x.axis

tbdbg bg method tbdmethod tbdmfrow mfrow mfcol tbdmfcol width tbdwidthtbdheight height tbdoma oma tbdmar mar

main.outer tbdmain.outer save2file tbdsave2file

y tbdy
digits tbddigits
prefix tbdprefix
cex.cor cex cor
plot.labels Boolean
order Boolean
decreasing Boolean
add.diagonal.line

tbdadd.diagonal.line

x2 tbdadd.diagonal.line vline tbdadd.diagonal.line cols tbdadd.diagonal.line

```
na.action
                 tbdadd.diagonal.line
drop.unused.levels
                 tbdadd.diagonal.line
p.val
                 tbdx
seed
                 tbdx
paired
                 tbdx
show.data.cloud
                 tbdx
ladder.add.line
                 tbdx
ladder.add.text
                 thdx
```

Details

myboxplot shows data points along with boxes. The data poins are jittered and the pattern of jittering is made reproducible in repeated calls. The test can only take one type of test currently.

myforestplot is modified from code from Allan deCamp/SCHARP. dat should have three columns. first column should be point estimate, second and third lci and uci, fourth p value. col.1 is the color used for CIs that do not include null, col.2 is used for CIs that do include null. If order is TRUE, the rows are ordered by the first column of dat. descreasing can be used to change the behavior of order.

corplot.formula uses MethComp::Deming by Bendix Carstensen to fit Deming regression.

wtd.hist is copied from weights package, author: Josh Pasek.

mymatplot will use na.approx (zoo) to fill in NA before plotting in order to draw continuous lines. The filled-in values will not be shown as points.

smoothed.scaled.hist draws histograms and overlay densities on top.

Examples

```
set.seed(1)
x=1:50+rnorm(50,0,4)
y=1:50+rnorm(50,0,4)
dat=data.frame(x, y)
corplot(y~x,dat,add.lm.fit=TRUE,add.deming.fit=TRUE,col.lm="red",col.deming="blue")

dat=data.frame(y=c(1:10,2:11), x=rep(c("a","b"),each=10), ptid=c(1:10,1:10))
par(mfrow=c(1,2))
myboxplot(y~x, dat, test="w", jitter=FALSE)
myboxplot(y~x, dat, test="w", add.interaction=TRUE, reshape.formula=y~x, reshape.id="ptid")

myboxplot(list(jitter(1:10), jitter(3:12)), test="w")
myboxplot(list(jitter(1:10), jitter(3:12)), test="w", paired=TRUE)

smoothed.scaled.hist(list(A=rnorm(100,0,1)), bin_width=0.1, xlab="x")
smoothed.scaled.hist(list(A=rnorm(100,0,1)), B=rnorm(500,10,2)),
```

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```
bin_width=0.1, xlab="x")

## Not run:
myfigure(mfrow=c(1,2))
    plot(1:10)
    plot(1:10)
mydev.off(ext="png,pdf", file="tmp")

## End(Not run)

#myboxplot x axis may look weird if log="xy"
```

print.functions

Print Functions

Description

roundup prints a specified number of digits after decimal point even if 0s are needed at the end. formatInt prints a specified number of digits before decimal point even if 0s are needed at the beginning.

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```
FALSE, floating = FALSE, lines = TRUE, hline.after =
                 NULL, add.to.row = NULL, sanitize.text.function =
                 NULL, append = FALSE, preamble = "", input.foldername
                 = NULL, save2input.only = NULL, caption = NULL, label
                 = paste("tab", last(strsplit(file.name, "/")[[1]]),
                 sep = " "), table.placement = "h!",
                 add.clear.page.between.tables = FALSE, longtable =
                 FALSE, verbose = FALSE, silent = TRUE, ...)
mytex.begin(file.name, preamble = "")
mytex.end(file.name)
mywrite(x, ...)
mywrite.csv(x, file = "tmp", row.names = FALSE, digits = NULL,
                 silent = TRUE, ...)
roundup(value, digits, na.to.empty = TRUE, remove.leading0 =
                 FALSE)
formatDouble(value, digits, na.to.empty = TRUE, remove.leading0 =
                 FALSE)
digit
               tbddigit
silent
               tbdnewline
input.foldername
               tbdnewline
```

Arguments

object tbdnewline newline tbdnewline print.name tbddigits save2input.only Boolean include.colnames Boolean col.headers string. Column headers comment Boolean, whether to include the version and timestamp comment hline.after vector add.to.row a list sanitize.text.function a function tbdvalue str

print.functions 37

```
remove.leading0
```

tbdvalue

caption tbdvalue longtable tbdvalue

label default to be the same as file.name stem

table.placement

tbdvalue

na.to.empty tbdvalue value tbdvalue digits tbddigits fill tbdfill

models tbdmodels

model.names tbdmodel.names row.major tbdrow.major round.digits tbdround.digits

dat tbddat

file.name tbdfile.name
display tbddisplay
align tbdalign
append tbdappend
preamble tbdpreamble

include.rownames

tbdinclude.rownames

floating tbdfloating lines tbdlines ... tbd... verbose tbd... x tbdx file tbdfile

row.names tbdrow.names add.clear.page.between.tables

tbdrow.names

Examples

```
roundup (3.1, 2) # 3.10
```

formatInt(3, 2) # 03

Not run:

38 random.functions

```
# demo of dimnames
tab=diag(1:4); rownames(tab)<-colnames(tab)<-1:4; names(dimnames(tab))=c("age","height")
# for greek letter in the labels, we need sanitize.text.function=identity
rownames(tab)[1]="$\alpha$"
# note that to use caption, floating needs to be TRUE
mytex (tab, file="tmp1", sanitize.text.function=identity,
         caption="This is a caption .....", caption.placement="top",
         floating=TRUE)
# col.headers has to have the RIGHT number of columns
# but align is more flexible, may not need to include the rownames col
tab=diag(1:4); rownames(tab)<-colnames(tab)<-1:4</pre>
mytex (tab, file="tmp", include.rownames = TRUE,
         align=c("c","c","c|","c","c"), col.headers=
          "\hline\n & \multicolumn\{2\}\{c\}\{Vaccine\} & \multicolumn\{2\}\{c\}\{Control\} \\ \n")
# not include rownames
mytex (tab, file="tmp", include.rownames = FALSE,
         align=c("c","c","c|","c","c"), col.headers=
                                          \mathcal{multicolumn} \ \multicolumn\2}{c|}{Vaccine} & \multicolumn\2}{c}{Control} \\ \n")
# It should work even if some rownames are duplicated
tab=diag(1:4); rownames(tab)=rep(1,4); colnames(tab)<-1:4</pre>
mytex (tab, file="tmp", include.rownames = TRUE,
         align=c("c","c|","c","c"), col.headers=
          "\hline\n & \multicolumn{2}{c|}{Vaccine} & \multicolumn{2}{c}{Control} \\ \n")
# add.to.rows
tab=diag(1:4); rownames(tab)<-1:4; colnames(tab)<-c("a","b","c","d")
mytex (tab, file="tmp",
         add.to.row=list( list(0,2),
                                                 \mbox{\mbox{\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\mbox{$\m
                   c("
                        "\hline\n \multicolumn\{5\}\{1\}\{Heading 2\}\ \n"
        ))
)
## End(Not run)
```

random.functions

Random Functions

Description

Generate samples from random variables.

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Usage

```
dbern(x, prob, log = FALSE)
dcorbern(x, p, a, log = FALSE)
dmixnorm(x, mix.p, sd1, sd2, log = FALSE)
dnorm.norm.gamma(x, p, same.distr = FALSE, log = FALSE)
rbern(n, prob, generalized = FALSE)
rbigamma(n, shape.1, shape.2, rate.1, rate.2, rho)
rbilogistic(n, loc.1, loc.2, scale.1, scale.2, rho)
rejective.sampling(N, n, pik)
rnorm.ar(n, sd, rho)
rnorm.norm.gamma(n, mu.0, lambda, alpha, beta)
rmixnorm (n, mix.p, mu1, mu2, sd1, sd2)
rdoublexp(n, location=0, scale=1)
ddoublexp(x, location=0, scale=1)
qdoublexp(p, location=0, scale=1)
pdoublexp(q, location=0, scale=1)
rbidoublexp(n, loc.1, loc.2, scale.1, scale.2, rho)
```

tbdx
tbdx
tbdx
tbdx
tbdprob
tbdlog
tbdp
tbda
tbdmix.p
tbdsd1
tbdsd2
tbdsame.distr

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n	tbdn
generalized	tbdgeneralized
N	tbdN
pik	tbdpik
mu	tbdmu
mu1	tbdmu
mu2	tbdmu
sd	tbdsd
alpha	tbdalpha
mu.0	tbdmu.0
lambda	tbdlambda
beta	tbdbeta
loc.1	tbdbeta
loc.2	tbdbeta
scale.1	tbdbeta
scale.2	tbdbeta
rate.1	tbdbeta
rate.2	tbdbeta
shape.1	tbdbeta
shape.2	tbdbeta
rho	tbdbeta

Details

rbern generates Bernoulli random variables.

rbilogistic generates a bivariate logistic distribution for correlation coefficient 0.5, or [-0.271, 0.478]. In the former case it is generated by calling rbilogis, part of the VGAM package; in the latter case it is generated via the AMH copular.

rnorm.ar simulate autoregressive normal random variables, correlation is rho^d between x_1 and x_1+d

Examples

```
set.seed(1)
rbern(n=10, p=1/2)
rbern(n=2, p=c(.999,.001))
## Not run:
tmp=replicate(1e4, rnorm.cor(10, 1, .81))
round(cor(t(tmp)),2)
## End(Not run)
```

```
regression.model.functions
```

Regression Model Functions

Description

getFormattedSummary prints a table of regression coefficient estimates and standard errors.

```
getFormattedSummary(fits, type = 12, est.digits = 2, se.digits = 2,
robust, random = FALSE, VE = FALSE, to.trim = FALSE,
rows = NULL, coef.direct = FALSE, trunc.large.est =
TRUE, scale.factor = 1, p.digits = 3, remove.leading0
= FALSE, p.adj.method = "fdr", ...)
getVarComponent(object, ...)
getFixedEf(object, ...)
risk.cal(risk, binary.outcome, weights = NULL, ngroups = NULL,
    cuts = NULL, main = "", add = FALSE, show.emp.risk = TRUE,
    lcol = 2, ylim = NULL, scale = c("logit", "risk"))
interaction.table(fit, v1, v2, v1.type = "continuous", v2.type = "continuous",
logistic.regression = TRUE)
## S3 method for class 'coxph'
getFixedEf(object, exp=FALSE,robust=FALSE, ...)
## S3 method for class 'gam'
getFixedEf(object, ...)
## S3 method for class 'gee'
getFixedEf(object, exp = FALSE, ...)
## S3 method for class 'geese'
getFixedEf(object, robust = TRUE, ...)
## S3 method for class 'tps'
getFixedEf(object, exp=FALSE, robust=TRUE, ...)
## S3 method for class 'glm'
getFixedEf(object, exp = FALSE, robust = TRUE, ret.robcov = FALSE,
    ...)
## S3 method for class 'svyglm'
getFixedEf(object, exp = FALSE, robust = TRUE, ...)
```

```
## S3 method for class 'svy_vglm'
getFixedEf(object, exp = FALSE, robust = TRUE, ...)
## S3 method for class 'svycoxph'
getFixedEf(object, exp = FALSE, robust = TRUE, ...)
## S3 method for class 'inla'
getFixedEf(object, ...)
## S3 method for class 'lm'
getFixedEf(object, exp = F, ...)
## S3 method for class 'lme'
getFixedEf(object, ...)
## S3 method for class 'logistf'
getFixedEf(object, exp = FALSE, ...)
## S3 method for class 'matrix'
getFixedEf(object, ...)
## S3 method for class 'MIresult'
getFixedEf(object, exp = FALSE, ...)
## S3 method for class 'hyperpar.inla'
getVarComponent(object, transformation = NULL, ...)
## S3 method for class 'matrix'
getVarComponent(object, ...)
## S3 method for class 'geese'
coef(object, ...)
## S3 method for class 'tps'
coef(object, ...)
## S3 method for class 'geese'
predict(object, x, ...)
## S3 method for class 'tps'
predict(object, newdata = NULL, type = c("link", "response"), ...)
## S3 method for class 'geese'
residuals(object, y, x,...)
## S3 method for class 'geese'
vcov(object, ...)
## S3 method for class 'tps'
vcov(object, robust, ...)
```

```
## S3 method for class 'logistf'
vcov(object, ...)
```

```
tbd...
                 tbdobject
object
fit
                 tbdfit
coef.direct
                 tbdfit
                 Boolean, whether to return robust variance estimate
robust
                 tbdexp
exp
remove.leading0
                 tbdexp
p.adj.method
                 tbdexp
cuts
                 tbdfits
                 tbdfits
ret.robcov
                 tbdfits
fits
type
                 tbdtype
est.digits
                 tbdest.digits
se.digits
                 tbdse.digits
                 tbdse.digits
p.digits
                 tbdrandom
random
۷E
                 tbdrandom
transformation tbdtransformation
weights
                 tbdv1
                 tbdv1
v1
v2
                 tbdv2
v1.type
                 tbdv1.type
v2.type
                 tbdv2.type
logistic.regression
                 tbdlogistic.regression
newdata
                 tbdx
                 tbdx
Χ
                 tbdy
У
to.trim
                 tbdy
rows
                 tbdy
risk
                 tbdfit
binary.outcome tbdfit
                 tbdfit
ngroups
```

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```
tbdfit
main
add
                  tbdfit
show.emp.risk
                  tbdfit
lcol
                  tbdfit
ylim
                  tbdfit
scale
                  tbdfit
trunc.large.est
                  tbdfit
                  tbdfit
scale.factor
```

Details

getFormattedSummary: from a list of fits, say lmer, inla fits, return formatted summary controlled by "type". For a matrix, return Monte Carlo variance random=TRUE returns variance components type=1: est type=2: est (se) type=3: est (2.5 percent, 97.5 percent) type=4: est se

getFixedEf returns a matrix, first column coef, second column se,

getFixedEf.matrix used to get mean and sd from a jags or winbugs sample, getVarComponent.matrix and getFixedEf.matrix do the same thing. Each column of samples is a variable

interaction.table expects coef and vcov to work with fit.

Examples

roc

ROC and AUC

Description

ROC/AUC methods. fastauc calculates the AUC using a sort operation, instead of summing over pairwise differences in R.

computeRoc computes an ROC curve.

plotRoc plots an ROC curve.

addRoc adds an ROC curve to a plot.

classification.error computes classification error

roc 45

Usage

Arguments

score a vector. Linear combination or score.

outcome a vector of 0 and 1. Outcome.

t0 a number between 0 and 1 that is the lower boundary of pAUC t1 a number between 0 and 1 that is the upper boundary of pAUC

reverse.sign.if.nece

a boolean. If TRUE, score is multiplied by -1 if AUC is less than 0.5.

x a list of two elements: sensitivity and specificity.
diag.line boolean. If TRUE, a diagonal line is plotted

add boolean. If TRUE, add to existing plot. If FALSE, create a new plot.

quiet boolean cutpoints threshold threshold verbose boolean

type line type for lines

... arguments passed to plot or lines

Details

These functions originally come from Thomas Lumley and Tianxi Cai et al.

Value

computeRoc returns a list of sensitivity and specificity. plotRoc and addRoc plots ROC curves.

Author(s)

```
Shuxin Yin <>
Youyi Fong <youyifong@gmail.com>
Krisztian Sebestyen <>
```

46 sim.dat.tvarying.two

Examples

```
n=1e2
score=c(rnorm(n/2,1), rnorm(n/2,0))
outcome=rep(1:0, each=n/2)
# cannot print due to r cmd check
#plotRoc(computeRoc(score, outcome))

# commented out b/c slower on pc and cause note when r cmd check
## test, fastauc2 is a version without all the checking
#score=rnorm(1e5)
#outcome=rbinom(1e5,1,.5)
#system.time(for (i in 1:1e2) fastauc(score,outcome)) # 4.9 sec
#system.time(for (i in 1:1e2) fastauc2(score,outcome)) # 3.8 sec
```

sim.dat.tvarying.two Simulation Functions for Time-dependent Proportional Hazard Model

Description

sim.dat.tvarying.three simulates from a model with time varing age group variale of three levels, sim.dat.tvarying.two two.

Usage

```
sim.dat.tvarying.three(n, followup.length, incidence.density,
    age.sim = c("tvaryinggroup", "baselinegroup", "continuous","bt"),
    random.censoring.rate = 0.05, seed)

sim.dat.tvarying.two(n, followup.length, incidence.density,
    age.sim = c("tvaryinggroup", "baselinegroup", "continuous","bt"),
    random.censoring.rate = 0.05, seed)
```

sim.dat.tvarying.two 47

Details

In sim.dat.tvarying.three, baseline age is uniformly distributed between 2.0 and 16.0, and divivded into three groups at 6 and 12. In sim.dat.tvarying.two, baseline age is uniformly distributed between 2.0 and 12.0, and divivded into two groups at 6.

Value

Return a data frame with the following columns:

ptid subject identifier trt treatment indicator 0/1 for.non.tvarying.ana

Boolean, used to subset dataset for non-time dependent analysis

C censoring time baseline.age age years at baseline

agegrp a factor with levels [0,6) [6,12) [12,100)

baseline.agegrp

a factor with levels [0,6) [6,12) [12,100)

tstart left bound of time interval tstop right bound of time interval

d event indicator

X followup time, in years

Author(s)

Youyi Fong

See Also

```
make.timedep.dataset
```

Examples

```
library(survival)

dat=sim.dat.tvarying.three(n=6000,followup.length=3, incidence.density=0.05,
    age.sim="tvaryinggroup", seed=1)
f.tvarying = Surv(tstart,tstop,d) ~ trt*agegrp
f = Surv(X,d) ~ trt*baseline.agegrp
fits=list()
fits[["tvarying"]]=coxph(f.tvarying, dat)
fits[["baseline"]]=coxph(f, subset(dat, for.non.tvarying.ana))
fits
```

48 stat.functions

 ${\tt stat.functions}$

Stat Functions

Description

H calculates entropy.

Usage

```
H(p, logbase = c("e", "2"))
mutual.info(two.way.table, logbase = c("e", "2"))
cor.mixed(x, ...)
## Default S3 method:
    cor.mixed(x, na.fun, method=c("pearson", "spearman"), ...)
## S3 method for class 'vector'
    cor.mixed(x, y, na.fun, method=c("pearson", "spearman"), ...)
## S3 method for class 'formula'
    cor.mixed(formula, data, na.fun, method=c("pearson", "spearman"), ...)
skew (x, na.rm = FALSE)
info.cor(two.way.table)
yule.y(two.by.two.matrix)
kappacor(two.by.two.matrix, weight = c(1, 1), maximum = FALSE)
1.measure(two.by.two.matrix)
```

p	either a count vector or a probability vector, but can not be a vector of member- ship indicator
logbase	tbdlogbase
na.rm	tbdlogbase
two.way.table	tbdtwo.way.table
x	tbdx
	tbd
na.fun	tbdna.fun

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```
method tbdmethod
y tbdy
formula tbdformula
data tbddata
two.by.two.matrix
tbdtwo.by.two.matrix
weight tbdweight
maximum tbdmaximum
```

Examples

```
H(rep(1/5,5))
H(rep(3,5))
```

string.functions

String Functions

Description

%+% concatenates its arguments and returns a string.

```
a %.% b

contain(s1, s2)
trim (x, trim.trailing=TRUE, trim.leading=TRUE)

escapeUnderline(name)

fileStem(file.name)

firstIndex(s1, s2)

getExt(file.name)

getFileStem(file.name)

getStem(file.name)

lastIndex(s1, s2)

remove.prefix(s, sep = "_")
```

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Arguments

а a b b s1s1 s2 s2name name file.name file.name sep sep sep trim.leading sep trim.trailing sep

Examples

```
x=1
x %.% "b" %.% "c"
```

testing.functions

Testing Functions

Description

Testing functions.

```
hosmerlem(y, yhat, g = 10)
quick.t.test(x, y, var.equal = FALSE)
signtest(x)
tukey.mtest(mu, ms, n)
vector.t.test(mean.x, mean.y, var.x, var.y, n)
myfisher.test(x,y,...)
mycor.test(x, method = c("pearson", "kendall", "spearman"), idx = NULL)
```

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Arguments

```
tbd
. . .
                 tbdy
у
                 tbdyhat
yhat
g
                 tbdg
Χ
                 tbdx
                 tbdvar.equal
var.equal
method
                 tbdmethod
                 tbdmu
mu
                 tbdms
ms
                 tbdn
mean.x
                 tbdmean.x
mean.y
                 tbdmean.y
                 tbdvar.x
var.x
var.y
                 tbdvar.y
idx
                 tbdvar.y
```

Examples

```
signtest(runif(10))
```

VEplot

Vaccine Efficacy Plots

Description

Vaccine efficacy plots.

```
VEplot (object, ...)
## S3 method for class 'cox.zph'
VEplot(object, resid = TRUE, se = TRUE, df = 4, nsmo = 40,
    var, ylab="VE", xlab="Time", xaxt="s", cex.axis=1, ...)
## S3 method for class 'glm'
VEplot(object, X1, X2, x, ...)
## S3 method for class 'cox.zph'
myplot(object, resid = TRUE, se = TRUE, df = 4, nsmo = 40, var,
    coef.transform=NULL,
    ylab=NULL,
    xlab="Time", xaxt="s", cex.axis=1,
    ...)
```

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Arguments

object An object

resid Boolean, whether to plot residuals

se Boolean, whether to plot confidence band

df degrees of freedom

nsmo number of points used to plot the fitted spline

var estimated variance matrix from the Cox model fit

xlab x label xaxt x axis

cex.axis cex for axis

ylab y label

coef.transform a function to transform Cox hazard ratio estimate

X1 a matrix of dimension k by p, where k is the length of x (see below) and p is the

length of coef(object)

X2 a matrix of dimension k by p, where k is the length of x (see below) and p is the

length of coef(object)

x a vector of length k that represents the x coordinate of the VE plot

... additional parameters

Details

VEplot and myplot.cox.zph are extensions of survival::plot.cox.zph to plot VE curve and other transformations.

myplot.cox.zph adds the following parameters to the original list of parameters in plot.cox.zph: coef.transform: a function to transform the coefficients ylab: y axis label xlab: x axis label

VEplot.glm computes a series of k VEs: for i in 1...k, VE[i] = P(Y=1|X1[i,])/P(Y=1|X2[i,]). It returns a 3 by k matrix, whose first row contains VE estimates and the second and third rows contain lower and upper bounds, respectively.

Author(s)

Youyi Fong, Dennis Chao

References

Durham, Longini, Halloran, Clemens, Azhar and Rao (1998) "Estimation of vaccine efficacy in the presence of waning: application to cholera vaccines." American Journal of Epidemiology 147(10): 948-959.

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