

Package ‘metabomxtr’

April 10, 2015

Type Package

Title A package to run mixture models for truncated metabolomics data with normal or lognormal distributions.

Version 1.0.0

Date 2014-07-12

Author Michael Nodzinski, Anna Reisetter, Denise Scholtens

Maintainer Michael Nodzinski <michael.nodzinski@northwestern.edu>

Depends methods, Biobase

Imports optimx, Formula, plyr, multtest

Suggests xtable

Description The functions in this package return optimized parameter estimates and log likelihoods for mixture models of truncated data with normal or lognormal distributions.

License GPL-2

biocViews Metabolomics, MassSpectrometry

R topics documented:

metabomxtr-package	2
metabdata	3
mxtrmod	5
mxtrmodLL	8
mxtrmodLRT	9
mxtrmodstart	11
xdesign-methods	12
yvals-methods	13
zdesign-methods	13

Index	14
--------------	-----------

metabomxtr-package *A package to run mixture models on truncated normal or lognormal data.*

Description

The functions in this package return optimized parameter estimates and negative log-likelihoods for mixture models of truncated normal or lognormal data.

Details

Package: metabomxtr
Type: Package
Version: 0.99.1
Date: 2014-07-12
License: GPL-2

The function `mxtrmodLL` calculates the negative log-likelihood of mixture models. The function `mxtrmodstart` returns starting parameter estimates to be used when optimizing the mixture model parameters. The function `mxtrmod` returns optimized mixture model parameter estimates and the negative log-likelihood of the model. The function `mxtrmodLRT` performs likelihood ratio tests of full vs. reduced mixture models.

Author(s)

Michael Nodzenski, Anna Reisetter, Denise Scholtens

Maintainer: Michael Nodzenski <michael.nodzenski@northwestern.edu>

References

Moulton LH, Halsey NA. A mixture model with detection limits for regression analyses of antibody response to vaccine. *Biometrics*. 1995 Dec;51(4):1570-8.

Examples

```
#Create sample data
set.seed(123)
yvar<-rlnorm(200)
these<-sample(1:100,20)
yvar[these]<-NA
logyvar<-log(yvar)
y2var<-rlnorm(200)
those<-sample(1:200,25)
y2var[those]<-NA
logy2var<-log(y2var)
pred1<-sample(0:1,200,replace=TRUE)
```

```

pred2<-sample(1:10,200,replace=TRUE)
pred3<-sample(0:1,200,replace=TRUE)
pred3miss<-sample(1:200,50)
pred3[pred3miss]<-NA
testdata<-data.frame(cbind(yvar,y2var,logyvar,logy2var,pred1,pred2,pred3))

#Get the names of the response variables
ynames<-names(testdata)[3:4]

#Run a full mixture model on each response variable
fullMod<-~pred1+pred2+pred3|pred1+pred2+pred3
fullModRes<-mxtrmod(ynames=ynames,mxtrModel=fullMod,data=testdata)
fullModRes

#Run a reduced mixture model on each response variable
redMod<-~pred2|pred2
redModRes<-mxtrmod(ynames=ynames,mxtrModel=redMod,data=testdata,fullModel=fullMod)
redModRes

#Compare models using likelihood ratio test
mxtrmodLRT(fullModRes,redModRes)

```

metabdata

A sample data set of truncated metabolomics data.

Description

This data set contains log transformed metabolite levels and phenotype data of sample 115 of pregnant women. Metabolite levels are contained in columns 11-59, and missing values are indicated by NA. All metabolites included contain at least 5 missing values. Columns 1:10 represent phenotypic data.

Usage

```
data(metabdata)
```

Format

A data frame with 115 observations on the following 59 variables.

`sample` a numeric vector representing sample number.

`PHENO` a factor representing high and low fasting blood glucose, with levels `MomHighFPG` `MomLowFPG`.

`age_ogtt` a numeric vector representing the woman's age in years when the oral glucose tolerance test was performed.

`age_ogtt_mc` a numeric vector representing mean centered `age_ogtt`.

`ga_ogtt_wks` a numeric vector representing gestational age in weeks when the oral glucose tolerance test was performed.

ga_ogtt_wks_mc a numeric vector representing mean centered ga_ogtt_wks.

FCg a factor representing field center where the data were collected with levels BCD E F P.

parity12 a numeric vector with 1 indicative of previous pregnancy and 0 otherwise.

storageTimesYears a numeric vector representing the number of years the metabolite sample had been stored prior to assay.

storageTimesYears_mc a numeric vector representing mean centered storageTimesYears.

ketovaline a numeric vector representing log2 transformed metabolite abundance.

alpha.ketoglutaric.acid a numeric vector representing log2 transformed metabolite abundance.

ketoleucine a numeric vector representing log2 transformed metabolite abundance.

acetoacetate a numeric vector representing log2 transformed metabolite abundance.

aldohexose a numeric vector representing log2 transformed metabolite abundance.

beta.alanine a numeric vector representing log2 transformed metabolite abundance.

methylmalonic.acid a numeric vector representing log2 transformed metabolite abundance.

creatinine a numeric vector representing log2 transformed metabolite abundance.

hexuronic.acid a numeric vector representing log2 transformed metabolite abundance.

ethanolamine a numeric vector representing log2 transformed metabolite abundance.

glutamine a numeric vector representing log2 transformed metabolite abundance.

glycolic.acid a numeric vector representing log2 transformed metabolite abundance.

isoleucine a numeric vector representing log2 transformed metabolite abundance.

malonic.acid a numeric vector representing log2 transformed metabolite abundance.

ribose a numeric vector representing log2 transformed metabolite abundance.

phenylalanine a numeric vector representing log2 transformed metabolite abundance.

pyruvic.acid a numeric vector representing log2 transformed metabolite abundance.

hexitol.1 a numeric vector representing log2 transformed metabolite abundance.

lysine a numeric vector representing log2 transformed metabolite abundance.

disaccharide.1 a numeric vector representing log2 transformed metabolite abundance.

tyrosine a numeric vector representing log2 transformed metabolite abundance.

leucine a numeric vector representing log2 transformed metabolite abundance.

hexitol.2 a numeric vector representing log2 transformed metabolite abundance.

disaccharide.2 a numeric vector representing log2 transformed metabolite abundance.

ornithine a numeric vector representing log2 transformed metabolite abundance.

disaccharide.3 a numeric vector representing log2 transformed metabolite abundance.

beta.tocopherol a numeric vector representing log2 transformed metabolite abundance.

hexitol.3 a numeric vector representing log2 transformed metabolite abundance.

benzene.1.2.4.triol a numeric vector representing log2 transformed metabolite abundance.

heptadecane a numeric vector representing log2 transformed metabolite abundance.

nonadecane a numeric vector representing log2 transformed metabolite abundance.

tetradecanedioic.acid a numeric vector representing log2 transformed metabolite abundance.
 pentadecanoic.acid a numeric vector representing log2 transformed metabolite abundance.
 undecane a numeric vector representing log2 transformed metabolite abundance.
 methyl.heptadecanoate a numeric vector representing log2 transformed metabolite abundance.
 hydrocarbon a numeric vector representing log2 transformed metabolite abundance.
 deoxyhexose a numeric vector representing log2 transformed metabolite abundance.
 glucose a numeric vector representing log2 transformed metabolite abundance.
 pentose.sugar a numeric vector representing log2 transformed metabolite abundance.
 hexitol.4 a numeric vector representing log2 transformed metabolite abundance.
 beta.sitosterol a numeric vector representing log2 transformed metabolite abundance.
 X1.5.anhydroglucitol a numeric vector representing log2 transformed metabolite abundance.
 threonine a numeric vector representing log2 transformed metabolite abundance.
 proline a numeric vector representing log2 transformed metabolite abundance.
 campesterol a numeric vector representing log2 transformed metabolite abundance.
 X6.deoxy.glucose a numeric vector representing log2 transformed metabolite abundance.
 erythronic.acid a numeric vector representing log2 transformed metabolite abundance.
 methyl.myristate a numeric vector representing log2 transformed metabolite abundance.
 methyl.eicosanoate a numeric vector representing log2 transformed metabolite abundance.

Source

Scholtens DM, Muehlbauer MJ, Daya NR, Stevens RD, Dyer AR, Lowe LP, Metzger BE, Newgard CB, Bain JR, Lowe WL Jr; HAPO Study Cooperative Research Group. Metabolomics reveals broad-scale metabolic perturbations in hyperglycemic mothers during pregnancy. *Diabetes Care*. 2014 Jan; 37(1):158-66.

mxtrmod

A function to return optimized parameter estimates and the negative log-likelihood of mixture models for truncated normal or lognormal data.

Description

This function returns optimized parameter estimates and the negative log-likelihood of mixture models for truncated normal or lognormal data.

Usage

```
mxtrmod(ynames, mxtrModel, Tvals=NULL, nNA=5, minProp=0.2, method="BFGS", data, fullModel=NULL)
```

Arguments

yname	A character vector of the mixture model outcome names, e.g. metabolites. If the input data object is a matrix or data frame, these should be column names. If the input data object is an expression set, these should be row names. Response variables should have normal or lognormal distributions. If lognormal, log transformed variables should be input. Missing values should be denoted by NA.
mxtrModel	A formula of the form $\sim x_1 + x_2 \dots z_1 + z_2 \dots$, where x's are the names of covariates included in the discrete portion of the model and z's are names of covariates included in the continuous portion. For intercept only models, enter 1 instead of covariate names on the appropriate side of the .
Tvals	A vector of thresholds below which continuous variables are not observable. By default, this parameter will be set to the minimum of the response variable.
nNA	The minimum number of unobserved values needed to be present for the discrete portion of the model likelihood to be calculated. Models for variables with fewer than nNA missing values will include only the continuous portion. The default value is 5.
minProp	The minimum proportion of non-missing data in the response variable necessary to run the model. The default value is 0.2. Models will not be run if more than 80% of response variable values are missing.
method	The method used to optimize the parameter estimates of the mixture model. "BFGS" is the default method. Other options are documented in the manual for the function 'optimx' in package optimx.
data	The input data object. Matrices, data frames, and expression sets are all acceptable classes. If a data frame or matrix, rows are subjects and columns are metabolites or outcomes.
fullModel	A formula of the form $\sim x_1 + x_2 \dots z_1 + z_2 \dots$, where x's are the names of covariates included in the discrete portion of the full model and z's are names of covariates included in the continuous portion. Input if the mxtrModel parameter represents a reduced model.

Value

Returns a data frame containing optimized estimates for all parameters in the mixture model, the negative log likelihood of the model, the optimization method used, whether the algorithm converged, and the total number of observations used.

Note

This function may generate warning messages about production of NaNs, but the function is still operating normally.

Author(s)

Michael Nodzinski, Anna Reisetter, Denise Scholtens

References

Moulton LH, Halsey NA. A mixture model with detection limits for regression analyses of antibody response to vaccine. *Biometrics*. 1995 Dec;51(4):1570-8.

Examples

```
#Create sample data frame
set.seed(123)
yvar<-rlnorm(200)
these<-sample(1:100,20)
yvar[these]<-NA
logyvar<-log(yvar)
y2var<-rlnorm(200)
those<-sample(1:200,25)
y2var[those]<-NA
logy2var<-log(y2var)
pred1<-sample(0:1,200,replace=TRUE)
pred2<-sample(1:10,200,replace=TRUE)
pred3<-sample(0:1,200,replace=TRUE)
pred3miss<-sample(1:200,50)
pred3[pred3miss]<-NA
testdata<-data.frame(cbind(yvar,y2var,logyvar,logy2var,pred1,pred2,pred3))

#Get the names of the response variables
ynames<-names(testdata)[3:4]

#Run a mixture model on each response variable
mod<-~pred1+pred2+pred3|pred1+pred2+pred3
mxtrmod(ynames=ynames,mxtrModel=mod,data=testdata)

#Create example expression set
#Specify the response variables
exprsobs<-t(testdata[,3:4])

#Specify the phenotype data
exprspheno<-testdata[,5:7]

#make phenotype data an annotated data frame
phenoData <- new("AnnotatedDataFrame",data=exprspheno)

#combine into example expression set
testexpr<-ExpressionSet(assayData=exprsobs,phenoData=phenoData)

#Get the names of the response variables
ynames<-rownames(exprs(testexpr))

#Run the mixture model on each response variable
mxtrmod(ynames=ynames,mxtrModel=mod,data=testexpr)

#Load the data set from the package
data(metabdata)
```

```
#Select the response variables
ynames<-names(metabdata)[11:17]

#Run the mixture models
mod2<-~PHENO|PHENO+age_ogtt_mc+parity12+ga_ogtt_wks_mc
mxtrmod(ynames,mxtrModel=mod2,data=metabdata)
```

mxtrmodLL	<i>A function to return the negative log-likelihood of mixture models of truncated, normal or lognormal data.</i>
-----------	---

Description

This function returns the negative log-likelihood of the specified mixture model.

Usage

```
mxtrmodLL(params, obsY, xVars, zVars, Tvals, includeDiscrete)
```

Arguments

params	A vector of parameter estimates for all paramters in the mixture model.
obsY	A vector containing the response variable, which must be normally or log normally distributed. If lognormal, log transformed Y's should be input as the response variable. Missing Y values should be indicated by NA.
xVars	The design matrix for the covariates included in the discrete portion of the model.
zVars	The design matrix for the covariates included in the continuous portion of the model.
Tvals	A vector of thresholds below which continuous variables are not observable.
includeDiscrete	A logical indicator for whether or not to include the discrete portion of the model.

Value

Returns the negative log-likelihood of the specified mixture model.

Author(s)

Michael Nodzenski, Anna Reisetter, Denise Scholtens

References

Moulton LH, Halsey NA. A mixture model with detection limits for regression analyses of antibody response to vaccine. *Biometrics*. 1995 Dec;51(4):1570-8.

Examples

```

#Create sample data
set.seed(123)
yvar<-rlnorm(200)
these<-sample(1:100,20)
yvar[these]<-NA
logyvar<-log(yvar)
y2var<-rlnorm(200)
those<-sample(1:200,25)
y2var[those]<-NA
logy2var<-log(y2var)
pred1<-sample(0:1,200,replace=TRUE)
pred2<-sample(1:10,200,replace=TRUE)
testdata<-data.frame(cbind(yvar,y2var,logyvar,logy2var,pred1,pred2))

#Create a vector of starting values for the function
startvals<-c(2,0,0,1.5,0,0,2)

#Create a vector of response variables
obsY<-testdata$logyvar

#Create the design matrix for the discrete portion of the model
xVars<-model.matrix(~pred1+pred2,data=testdata)

#Create the design matrix for the continuous portion of the model
zVars<-model.matrix(~pred1+pred2,data=testdata)

#Create the Tvals vector
Tvals<-rep(min(obsY,na.rm=TRUE),length(obsY))

#Determine if the discrete portion should be included in the model
includeDiscrete<-sum(is.na(obsY))>5

#Calculate the negative log-likelihood
mxtrmodLL(params=startvals,obsY=obsY,xVars=xVars,zVars=zVars,Tvals=Tvals,
           includeDiscrete=includeDiscrete)

```

mxtrmodLRT

A function to run likelihood ratio tests on full vs. reduced mixture models.

Description

This function runs likelihood ratio tests on full vs. reduced mixture models. Input arguments are data frame outputs from the mxtrmod function.

Usage

```
mxtrmodLRT(fullmod, redmod, adj = NULL)
```

Arguments

fullmod	The output data frame from the mxtrmod function on the full mixture model.
redmod	The output data frame from the mxtrmod function on the reduced mixture model.
adj	The adjustment method for multiple comparisons. The default is set to NULL. Options for adjustment methods are described in the documentation for the function <code>mt.rawp2adjp</code> in the <code>multtest</code> package.

Value

A data frame containing the response variables (i.e. metabolites), negative log likelihoods of full and reduced models, chi square statistics, degrees of freedom, p-values, and, if requested, adjusted p-values.

Author(s)

Michael Nodzenski, Anna Reisetter, Denise Scholtens

References

Moulton LH, Halsey NA. A mixture model with detection limits for regression analyses of antibody response to vaccine. *Biometrics*. 1995 Dec;51(4):1570-8.

Examples

```
#Create sample data
set.seed(123)
yvar<-rlnorm(200)
these<-sample(1:100,20)
yvar[these]<-NA
logyvar<-log(yvar)
y2var<-rlnorm(200)
those<-sample(1:200,25)
y2var[those]<-NA
logy2var<-log(y2var)
pred1<-sample(0:1,200,replace=TRUE)
pred2<-sample(1:10,200,replace=TRUE)
pred3<-sample(0:1,200,replace=TRUE)
pred3miss<-sample(1:200,50)
pred3[pred3miss]<-NA
testdata<-data.frame(cbind(yvar,y2var,logyvar,logy2var,pred1,pred2,pred3))

#Get the names of the response variables
ynames<-names(testdata)[3:4]

#Run a full mixture model on each response variable
fullMod<-~pred1+pred2+pred3|pred1+pred2+pred3
fullModRes<-mxtrmod(ynames=ynames,mxtrModel=fullMod,data=testdata)
fullModRes

#Run a reduced mixture model on each response variable
```

```

redMod<-~pred2|pred2
redModRes<-mxtrmod(ynames=ynames,mxtrModel=redMod,data=testdata,fullModel=fullMod)
redModRes

#Compare models using likelihood ratio test
mxtrmodLRT(fullModRes,redModRes)

```

mxtrmodstart	<i>A function to generate starting parameter estimates for the optimization of mixture model parameters.</i>
--------------	--

Description

This function returns starting parameter estimates for the optimization of the mixture model parameters. The intercept of the continuous portion is set to the mean of the observed responses and the intercept of the discrete portion is set to the log odds of having observed a response. All other parameter starting values are set to zero.

Usage

```
mxtrmodstart(obsY, xVars, zVars, includeDiscrete)
```

Arguments

obsY	A vector containing the response variable, which must be normally or log normally distributed. If lognormal, log transformed Y's should be input as the response variable. Missing Y values should be indicated by NA.
xVars	The design matrix for the covariates included in the discrete portion of the model.
zVars	The design matrix for the covariates included in the continuous portion of the model.
includeDiscrete	A logical indicator for whether or not to include the discrete portion of the model.

Value

A vector containing the starting values for each parameter in the mixture model function, to be used as starting points when optimizing the parameter estimates.

Author(s)

Michael Nodzenski, Anna Reisetter, Denise Scholtens

References

Moulton LH, Halsey NA. A mixture model with detection limits for regression analyses of antibody response to vaccine. *Biometrics*. 1995 Dec;51(4):1570-8.

Examples

```

#Create sample data
set.seed(123)
yvar<-rlnorm(200)
these<-sample(1:100,20)
yvar[these]<-NA
logyvar<-log(yvar)
y2var<-rlnorm(200)
those<-sample(1:200,25)
y2var[those]<-NA
logy2var<-log(y2var)
pred1<-sample(0:1,200,replace=TRUE)
pred2<-sample(1:10,200,replace=TRUE)
testdata<-data.frame(cbind(yvar,y2var,logyvar,logy2var,pred1,pred2))

#Create a vector of response variables
obsY<-testdata$logyvar

#Create the design matrix for the discrete portion of the model
xVars<-model.matrix(~pred1+pred2,data=testdata)

#Create the design matrix for the continuous portion of the model
zVars<-model.matrix(~pred1+pred2,data=testdata)

#Determine if the discrete portion should be included in the model
includeDiscrete<-sum(is.na(obsY))>5

#Calculate starting values
mxtrmodstart(obsY=obsY,xVars=xVars,zVars=zVars,includeDiscrete=includeDiscrete)

```

xdesign-methods

~~ Methods for Function xdesign ~~

Description

~~ Methods for function xdesign ~~

Methods

signature(x="data.frame",m="ANY") The columns of data frame 'x' specified in the input Formula object 'm' are converted to the design matrix for the discrete portion of the mixture model.

signature(x="ExpressionSet",m="ANY") The columns of the phenoData section of expression set 'x' specified in the input Formula object 'm' are converted to the design matrix for the discrete portion of the mixture model.

signature(x="matrix",m="ANY") The columns of matrix 'x' specified in the input Formula object 'm' are converted to the design matrix for the discrete portion of the mixture model.

yvals-methods

~~ Methods for Function yvals ~~

Description

~~ Methods for function yvals ~~

Methods

signature(y="data.frame",n="character") The columns of data frame 'y' with names 'n' are converted to a data frame, whose columns are to be used as the response variables in the specified mixture model.

signature(y="ExpressionSet",n="character") The rows of the assayData section of expression set 'y' with names 'n' are converted to a data frame, whose columns are to be used as the response variables in the specified mixture model.

signature(y="matrix",n="character") The columns of matrix 'y' with names 'n' are converted to a data frame, whose columns are to be used as the response variables in the specified mixture model.

zdesign-methods

~~ Methods for Function zdesign ~~

Description

~~ Methods for function zdesign ~~

Methods

signature(x="data.frame",m="ANY") The columns of data frame 'x' specified in the input Formula object 'm' are converted to the design matrix for the continuous portion of the mixture model.

signature(x="ExpressionSet",m="ANY") The columns of the phenoData section of expression set 'x' specified in the input Formula object 'm' are converted to the design matrix for the continuous portion of the mixture model.

signature(x="matrix",m="ANY") The columns of matrix 'x' specified in the input Formula object 'm' are converted to the design matrix for the continuous portion of the mixture model.

Index

*Topic **datasets**

metabdata, [3](#)

*Topic **methods**

xdesign-methods, [12](#)

yvals-methods, [13](#)

zdesign-methods, [13](#)

*Topic **package**

metabomxtr-package, [2](#)

metabdata, [3](#)

metabomxtr (metabomxtr-package), [2](#)

metabomxtr-package, [2](#)

mxtrmod, [5](#)

mxtrmodLL, [8](#)

mxtrmodLRT, [9](#)

mxtrmodstart, [11](#)

xdesign (xdesign-methods), [12](#)

xdesign,data.frame-method
(xdesign-methods), [12](#)

xdesign,ExpressionSet-method
(xdesign-methods), [12](#)

xdesign,matrix-method
(xdesign-methods), [12](#)

xdesign-methods, [12](#)

yvals (yvals-methods), [13](#)

yvals,data.frame,character-method
(yvals-methods), [13](#)

yvals,ExpressionSet,character-method
(yvals-methods), [13](#)

yvals,matrix,character-method
(yvals-methods), [13](#)

yvals-methods, [13](#)

zdesign (zdesign-methods), [13](#)

zdesign,data.frame-method
(zdesign-methods), [13](#)

zdesign,ExpressionSet-method
(zdesign-methods), [13](#)

zdesign,matrix-method
(zdesign-methods), [13](#)

zdesign-methods, [13](#)