# Package: addhaz (via r-universe)

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Title Binomial and Multinomial Additive Hazard Models
Version 0.5
<b>Description</b> Functions to fit the binomial and multinomial additive hazard models and to estimate the contribution of diseases/conditions to the disability prevalence, as proposed by Nusselder and Looman (2004) and extended by Yokota et al (2017).
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Fit Binomial Additive Hazard Models

## **Description**

This function fits binomial additive hazard models subject to linear inequality constraints using the function constrOptim in the stats package for binary outcomes. Additionally, it calculates the cause-specific contributions to the disability prevalence based on the attribution method, as proposed by Nusselder and Looman (2004).

## Usage

## Arguments

formula	a formula expression of the form response ~ predictors, similar to other regression models. In case of attrib = TRUE, the first predictor in the formula should be the attrib.var. See example.
data	an optional data frame or matrix containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which BinAddHaz is called.
subset	an optional vector specifying a subset of observations to be used in the fitting process. All observations are included by default.
weights	an optional vector of survey weights to be used in the fitting process.
na.action	a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options, and is na.fail if that is unset. The 'factory-fresh' default is na.omit.
mode1	logical. If TRUE, the model frame is included as a component of the returned object.
contrasts	an optional list to be used for some or all of the factors appearing as variables in the model formula.
start	an optional vector of starting values. If not provided by the user, it is automatically generated using glm, family = poisson.
attrib	logical. Should the attribution of disability to chronic diseases/conditions be estimated? Default is TRUE.
attrib.var	character indicating the name of the attribution variable to be used if attrib = TRUE. If missing, the attribution results will not be stratified by the levels of the attribution variable. The attribution variable must be the first variable included in the linear predictor in formula. See example.

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collapse.background

logical. Should the background be collapsed across the levels of the attrib.var? If FALSE, the background will be estimated for each level of the attrib.var. If TRUE, only one background will be estimated. If TRUE, attrib.var must be specified. Default is FALSE.

attrib.disease logical. Should the attribution of diseases be stratified by the levels of the attri-

bution variable? If FALSE, the attribution of diseases will not be stratified by the levels of the attrib.var. If TRUE, the attribution of diseases will be estimated for each level of the attrib.var. If TRUE, interaction between diseases and the

attribution variable must be provided in the formula. Default is FALSE.

type of attribution to be estimated. The options are "abs" for absolute contritype.attrib

bution, "rel" for relative contribution, or "both" for both absolute and relative

contributions. Default is "abs".

seed an optional integer indicating the random seed.

logical. Should bootstrap percentile confidence intervals be estimated for the bootstrap

model parameters and attributions? Default is FALSE. See details.

conf.level scalar containing the confidence level of the bootstrap percentile confidence in-

tervals. Default is 0.95.

nbootstrap integer. Number of bootstrap replicates.

parallel logical. Should parallel calculations be used to obtain the bootstrap percentile

confidence intervals? Only valid if bootstrap = TRUE. Default is FALSE.

type of parallel operation to be used (if parallel = TRUE), with options: "multicore" type.parallel

and "snow". Default is "snow". See details.

ncpus integer. Number of processes to be used in the parallel operation: typically one

would choose this to be the number of available CPUs. Default is 4.

other arguments passed to or from the other functions.

## **Details**

The model is a generalized linear model with a non-canonical link function, which requires a restriction on the linear predictor ( $\eta \geq 0$ ) to produce valid probabilities. This restriction is implemented in the optimization procedure, with an adaptive barrier algorithm, using the function constrOptim in the stats package.

The variance-covariance matrix is based on the observed information matrix.

This version of the package only allows the calculation of non-parametric bootstrap percentile confidence intervals (CI). Also, the function gives the user the option to do parallel calculation of the bootstrap CI. The snow parallel option is available for all operating systems (Windows, Linux, and Mac OS) while the multicore option is only available for Linux and Mac OS systems. These two calculations are done by calling the boot function in the boot package. For more details, see the documentation of the boot package.

More information about the binomial additive hazard model and the calculation of the contribution of chronic conditions to the disability prevalence can be found in the references.

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#### Value

A list with arguments:

coefficients numerical vector with the regression coefficients.

ci confidence intervals calculated via bootstraping (if bootstrap = TRUE) or as the

inverse of the observed information matrix.

resDeviance residual deviance.
df degrees of freedom.

pvalue numerical vector of p-values based on the Wald test. Only provided if bootstrap

= FALSE.

stdError numerical vector with the standard errors for the parameter estimates based on

the inverse of the observed information matrix. Only provided if bootstrap =

FALSE.

vcov variance-covariance (inverse of the observed information matrix). Only pro-

vided if bootstrap = FALSE.

contribution for type.attrib = "abs" or "rel", a matrix is provided. For type.attrib =

"both", a list with two matrices ("abs" and "rel") is provided. This represents the contribution of each predictor in the model (usually diseases) to the disability prevalence. Percentile bootstrap confidence intervals are provided if bootstrap

= TRUE.

bootsRep matrix with the bootstrap replicates of the regression coefficients and contribu-

tions (if attrib = TRUE). If attrib = FALSE, it returns a logical, FALSE.

conf.level confidence level of the bootstrap CI. Only provided if bootstrap = TRUE.

bootstrap logical. Was bootstrap CI requested?

fitted.values the fitted mean values, obtained by transforming the linear predictor by the in-

verse of the link function.

residuals difference between the observed response and the fitted values.

call the matched call.

#### Author(s)

Renata T C Yokota. This function is based on the R code developed by Caspar W N Looman and Wilma J Nusselder for non R-users, with modifications. Original R code is available upon request to Wilma J Nusselder (w.nusselder@erasmusmc.nl).

#### References

Nusselder, W.J., Looman, C.W.N. (2004). Decomposition of differences in health expectancy by cause. Demography, 41(2), 315-334.

Nusselder, W.J., Looman, C.W.N. (2010). WP7: Decomposition tools: technical report on attribution tool. European Health Expectancy Monitoring Unit (EHEMU). Available at <a href="http://www.europex.eu/pdf/Reports\_2010/2010TR7.2\_TR%20on%20attribution%20tool.pdf">http://www.europex.eu/pdf/Reports\_2010/2010TR7.2\_TR%20on%20attribution%20tool.pdf</a>.

Yokota, R.T.C., Van Oyen, H., Looman, C.W.N., Nusselder, W.J., Otava, M., Kifle, Y.W., Molenberghs, G. (2017). Multinomial additive hazard model to assess the disability burden using cross-sectional data. Biometrical Journal, 59(5), 901-917.

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#### See Also

MultAddHaz

## **Examples**

```
data(disabData)
 ## Model without bootstrap CI and no attribution
 fit1 <- BinAddHaz(dis.bin ~ diab + arth + stro , data = disabData, weights = wgt,</pre>
                    attrib = FALSE)
 summary(fit1)
 ## Model with bootstrap CI and attribution without stratification, no parallel calculation
 # Warning message due to the low number of bootstrap replicates
## Not run:
 fit2 <- BinAddHaz(dis.bin ~ diab + arth + stro , data = disabData, weights = wgt,</pre>
                    attrib = TRUE, collapse.background = FALSE, attrib.disease = FALSE,
                  type.attrib = "both", seed = 111, bootstrap = TRUE, conf.level = 0.95,
                    nbootstrap = 5)
 summary(fit2)
 ## Model with bootstrap CI and attribution of diseases and background stratified by
 ## age, with parallel calculation of bootstrap CI
 # Warning message due to the low number of bootstrap replicates
 diseases <- as.matrix(disabData[,c("diab", "arth", "stro")])</pre>
 fit3 <- BinAddHaz(dis.bin ~ factor(age) -1 + diseases:factor(age), data = disabData,
                    weights = wgt, attrib = TRUE, attrib.var = age,
                collapse.background = FALSE, attrib.disease = TRUE, type.attrib = "both",
                    seed = 111, bootstrap = TRUE, conf.level = 0.95, nbootstrap = 10,
                    parallel = TRUE, type.parallel = "snow", ncpus = 4)
 summary(fit3)
## End(Not run)
```

disabData

Example of disability data

## **Description**

The disabData is a subset of the data from the 2013 National Health Survey in Brazil ("Pesquisa Nacional de Saude, 2013"). The data are restricted to women aged 60 years or older, resulting in 6294 individuals.

## Usage

```
data(disabData)
```

LRTest

#### **Format**

This dataset has information about disability and chronic conditions. The disability outcomes were defined as limitations on instrumental activities of daily living (IADL). Individuals with missing data were excluded. The data frame contains 7 variables:

- dis.bin: disability as a binary variable, with 2 categories: 0 (no disability), 1 (disability).
- **dis.mult:** disability as a multinomial variable, with 3 categories: 0 (no disability), 1 (mild disability), and 2 (severe disability).
- wgt: survey weights.
- age: binary variable for age: 0 (60-79 years) or 1 (80+ years).
- diab: binary variable for diabetes: 0 (no) or 1 (yes).
- arth: binary variable for arthritis: 0 (no) or 1 (yes).
- stro: binary variable for stroke: 0 (no) or 1 (yes).

#### Source

The data were obtained from the National Health Survey 2013, Brazil. For more information about the survey, see references.

#### References

Szwarcwald, C.L., Malta, D.C., Pereira, C.A., Vieira, M.L., Conde, W.L., Souza Junior, P.R., et al. (2013). National Health Survey in Brazil: design and methodology of application. Cien Saude Colet., 19(2): 333:42 [Article in Portuguese].

Instituto Brasileiro de Geografia e Estatistica (IBGE). Pesquisa Nacional de Saude 2013. Available at <a href="http://www.ibge.gov.br/home/estatistica/populacao/pns/2013/">http://www.ibge.gov.br/home/estatistica/populacao/pns/2013/</a>.

## **Examples**

data(disabData)
str(disabData)

LRTest

Perform likelihood ratio test

## **Description**

This function performs the likelihood ratio test to compare two nested binomial or multinomial additive hazard models. It can be used for model selection.

## Usage

```
LRTest(model1, model2)
```

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## **Arguments**

model1, model2 objects of class "binaddhazmod" or "multaddhazmod" to be compared. See example.

#### **Details**

The likelihood ratio test is defined as -2\*log(likelihood model 1/likelihood model 2). The resulting test statistic is assumed to follow a chi-squared distribution, with degrees of freedom (df) equal to the difference of the df between the models. If the test is statistically significant, the model with more variables fits the data significantly better than the model with less variables.

#### Value

A data frame with columns:

Res.df degrees of freedom for each model.

Res.dev 2\*log-likelihood of each model.

df difference in the degrees of freedom between models 1 and 2.

Deviance difference between the 2\*log-likelihood of models 1 and 2, representing the value of the likelihood ratio test statistic.

Pr(>Chi) p-value, based on the chi-squared distribution.

## **Examples**

```
data(disabData)
 ## Comparing two binomial models
 fit1 <- BinAddHaz(dis.bin ~ diab + arth + stro , data = disabData, weights = wgt,</pre>
                    attrib = FALSE)
 diseases <- as.matrix(disabData[,c("diab", "arth", "stro")])</pre>
 fit2 <- BinAddHaz(dis.bin ~ factor(age) -1 + diseases:factor(age), data = disabData,
                    weights = wgt, attrib = FALSE)
 LRTest(fit2, fit1)
 ## Comparing two multinomial models
## Not run:
 fit3 <- MultAddHaz(dis.mult ~ diab + arth + stro , data = disabData, weights = wgt,
                     attrib = FALSE)
 fit4 <- MultAddHaz(dis.mult ~ factor(age) -1 + diseases: factor(age), data = disabData,
                     weights = wgt, attrib = FALSE)
 LRTest(fit4, fit3)
## End(Not run)
```

MultAddHaz

Fit Multinomial Additive Hazard Models

## **Description**

This function fits multinomial additive hazard models subject to linear inequality constraints using the function constr0ptim in the stats package for multinomial (multi-category) outcomes. It also calculates the cause-specific contributions to the disability prevalence for each category of the response variable based on the extension of the attribution method, as proposed by Yokota et al (2017).

## Usage

## **Arguments**

Ę	guments	
	formula	a formula expression of the form response ~ predictors, similar to other regression models. In case of attrib = TRUE, the first predictor in the formula should be the attrib.var. See example.
	data	an optional data frame or matrix containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which MultAddHaz is called.
	subset	an optional vector specifying a subset of observations to be used in the fitting process. All observations are included by default.
	weights	an optional vector of survey weights to be used in the fitting process.
	na.action	a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options, and is na.fail if that is unset. The 'factory-fresh' default is na.omit.
	model	logical. If TRUE, the model frame is included as a component of the returned object.
	contrasts	an optional list to be used for some or all of the factors appearing as variables in the model formula.
	start	an optional vector of starting values. If not provided by the user, it is randomly generated.
	attrib	logical. Should the attribution of chronic diseases/conditions for each disability level be estimated? Default is TRUE.
	attrib.var	character indicating the name of the attribution variable to be used if attrib = TRUE. If missing, the attribution results will not be stratified by the levels of the attribution variable. The attribution variable must be the first variable included in the linear predictor in formula. See example.

collapse.background

logical. Should the background be collapsed across the levels of the attrib.var for each disability level? If FALSE, the background will be estimated for each level of the attrib.var. If TRUE, only one background will be estimated. If TRUE, attrib. var must be specified. Default is FALSE.

attrib.disease logical. Should the attribution of diseases be stratified by the levels of the attribution variable for each disability level? If FALSE, the attribution of diseases will not be stratified by the levels of the attrib.var. If TRUE, the attribution of diseases will be estimated for each level of the attrib.var. If TRUE, interaction between diseases and the attribution variable must be provided in the formula. Default is FALSE.

type.attrib

type of attribution to be estimated. The options are "abs" for absolute contribution, "rel" for relative contribution, or "both" for both absolute and relative contributions. Default is "abs".

integer indicating the random seed. seed

bootstrap logical. Should bootstrap percentile confidence intervals be estimated for the

model parameters and attributions? Default is FALSE. See details.

conf.level scalar containing the confidence level of the bootstrap percentile confidence in-

tervals. Default is 0.95.

nbootstrap integer. Number of bootstrap replicates.

logical. Should parallel calculations be used to obtain the bootstrap percentile parallel

confidence intervals? Default is FALSE.

type of parallel operation to be used (if parallel = TRUE), with options: "multicore" type.parallel

and "snow". Default is "snow". See details.

ncpus integer. Number of processes to be used in the parallel operation: typically one

would choose this to be the number of available CPUs. Default is 4.

other arguments passed to or from the other functions.

## **Details**

The model is a generalized linear model with a non-canonical link function, which requires a restriction on the linear predictor ( $\eta \geq 0$ ) to produce valid probabilities. This restriction is implemented in the optimization procedure, with an adaptive barrier algorithm, using the function constrOptim in the stats package.

The variance-covariance matrix is based on the observed information matrix.

This version of the package only allows the calculation of non-parametric bootstrap percentile confidence intervals (CI). Stratified bootstrap is applied to each category of the outcome. Also, the function gives the user the option to do parallel calculation of the bootstrap CI. The snow parallel option is available for all operating systems (Windows, Linux, and Mac OS) while the multicore option is only available for Linux and Mac OS systems. These two calculations are done by calling the boot function in the boot package. For more details see the documentation of the boot package.

More information about the multinomial additive hazard model and the estimation of the contribution of chronic conditions to the disability prevalence can be found in the references.

#### Value

## A list with arguments:

coefficients column matrix with the regression coefficients.

ci matrix with confidence intervals calculated via bootstraping (if bootstrap =

TRUE) or as the inverse of the observed information matrix.

resDeviance residual deviance.

df degrees of freedom.

pvalue column matrix of p-values based on the Wald test. Only provided if bootstrap

= FALSE.

stdError column matrix with the standard errors for the parameter estimates based on

the inverse of the observed information matrix. Only provided if bootstrap =

FALSE.

vcov variance-covariance matrix (inverse of the observed information matrix). Only

provided if bootstrap = FALSE.

contribution for type.attrib = "abs" or "rel", a matrix is provided. For type.attrib =

"both", a list with two matrices ("abs" and "rel") is provided. This represents the contribution of each predictor in the model (usually diseases) to the disability prevalence. Percentile bootstrap confidence intervals are provided if bootstrap

= TRUE. If attrib = FALSE, it returns a logical, FALSE.

bootsRep matrix with the bootstrap replicates of the regression coefficients and contribu-

tions (if attrib = TRUE).

conf.level confidence level of the bootstrap CI. Only provided if bootstrap = TRUE.

bootstrap logical. Was bootstrap CI requested?

call the matched call.

#### Author(s)

Renata T. C. Yokota. This function is based on the R code developed by Caspar W. N. Looman and Wilma J. Nusselder for the binomial additive hazard model with modifications and adaptations for the multinomial case.

#### References

Yokota, R.T.C., Van Oyen, H., Looman, C.W.N., Nusselder, W.J., Otava, M., Kifle, Y.W., Molenberghs, G. (2017). Multinomial additive hazard model to assess the disability burden using cross-sectional data. Biometrical Journal, 59(5), 901-917.

#### See Also

BinAddHaz

## **Examples**

```
data(disabData)
 ## Model without bootstrap CI and no attribution
 fit1 <- MultAddHaz(dis.mult ~ diab + arth + stro , data = disabData, weights = wgt,</pre>
                     attrib = FALSE)
 summary(fit1)
 ## Model with bootstrap CI and attribution without stratification, no parallel calculation
 # Warning message due to the low number of bootstrap replicates
 fit2 <- MultAddHaz(dis.mult ~ diab + arth + stro , data = disabData, weights = wgt,</pre>
                     attrib = TRUE, collapse.background = FALSE, attrib.disease = FALSE,
                   type.attrib = "both", seed = 111, bootstrap = TRUE, conf.level = 0.95,
                     nbootstrap = 5)
 summary(fit2)
 ## Model with bootstrap CI and attribution of diseases and background stratified by
 ## age, with parallel calculation of bootstrap CI
 # Warning message due to the low number of bootstrap replicates
 diseases <- as.matrix(disabData[,c("diab", "arth", "stro")])</pre>
 fit3 <- MultAddHaz(dis.mult ~ factor(age) -1 + diseases: factor(age), data = disabData,
                     weights = wgt, attrib = TRUE, attrib.var = age,
                collapse.background = FALSE, attrib.disease = TRUE, type.attrib = "both",
                     seed = 111, bootstrap = TRUE, conf.level = 0.95, nbootstrap = 5,
                     parallel = TRUE, type.parallel = "snow", ncpus = 4)
 summary(fit3)
## End(Not run)
```

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