Package 'glmfitmiss'

April 22, 2025

Title Fitting GLMs with Missing Data in Both Responses and Covariates

Description

Fits generalized linear models (GLMs) when there is missing data in both the response and categorical covariates. The functions implement likelihood-based methods using the Expectation and Maximization (EM) algorithm and optionally apply Firth's bias correction for improved inference. See Pradhan, Nychka, and Bandyopadhyay (2025) https://www.englistand.com proved inference. See Pradhan, Nychka, and Bandyopadhyay (2025) https://www.englistand.com han (2009) <doi:10.1111/j.1541-0420.2008.01186.x>, Maity, Pradhan, and Das (2019) <doi:10.1080/00031305.2017.1407359> for further methodological details.

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Contents

glmfitmiss-package						•		•	 •						•	•		 	•						2
emBinRegMAR				•	•			•	 •			•		•		•		 	•		•	•	•		4
emBinRegMixedMAR			•	•	•	•	•	•	 •		•	•	•	•		•	•	 	•	•	•	•	•	•	7

emBinRegNonIG	9
emforbeta	11
emil	14
emyxmiss	16
est	19
est45	20
felinedata	22
ibrahim	23
incontinence	24
llkmiss	25
logRegMAR	26
meningitis	27
meningitis60ymis	28
metastmelanoma	29
simulateCovariateData	30
simulateData	31
simulateMissDfYorX	32
sixcitydata	33
testyxm	34
	36

Index

glmfitmiss-package glmfitmiss: Fitting Binary Regression Models with Missing Data

Description

The glmfitmiss package provides functions for fitting binary regression models in the presence of missing data in both response variable level and covariate levels. The package includes likelihood-based methods, primarily based on the EM algorithm by Ibrahim (1990) for handling missing data mechanisms. Bias-reducing adjusted score approaches introduced by Firth (1993) are also incorporated in all the supported methods.

Details

This package enhances the accuracy of binary regression modeling in the presence of missing data by incorporating Ibrahim (1990) EM algorithm and Firth (1993) bias-reducing adjusted score methods.

The main functions in this package are:

- emBinRegMAR: Fits a binary regression model with missing categorical covariates. Assumes missing data are Missing at Random (MAR).
- emBinRegNonIG: Fits a binary regression model with missing responses that are nonignorable based on Ibrahim and Lipsitz (1996).
- emBinRegMixedMAR: Fits a binary regression model with missing responses and covariates, accounting for the non-ignorable missing responses assumption and Missing at Random (MAR) missing covariates.

glmfitmiss-package

• logRegMAR: Fits a logistic regression model (binary regression with a link=logit) with missing categorical covariates that are Missing at Random (MAR).

The other functions and data included in this package are

- emforbeta: The function to fit binary regression models with missing categorical covariates is implemented using a likelihood-based method, specifically the EM algorithm proposed by Ibrahim (1990).
- est: Example using Eastern Cooperative Oncology Group clinical trial.
- meningitis: Example using Meningococcal Disease Data.
- metastmelanoma: Example from a cancer clinical trial metastatic melanoma Kirkwood et al. (1996).
- emyxmiss: The main function fits binary regression models while accounting for missing responses and missing categorical covariates. This function implements a novel likelihoodbased method using the EM algorithm. For more information, refer to the work by Pradhan, Nychka, and Bandyopadhyay (2025).
- meningitis60ymis: Meningococcal Disease Data with missing response variable.
- 11kmiss: Log-likelihood function for models with missing data with out using EM-algorithms.
- est45: Example using Eastern Cooperative Oncology Group clinical trial a subset of the 'est' data.
- simulateData: Function to simulate response data.
- simulateCovariateData: Function to simulate covariate data.
- felinedata: Sykes et al. (1999) data, the risk factors for Chlamy, a chlamydial infection in cats.
- simulateMissDfYorX: This function generates missing covariate or missing responses data. The missing data generation in the last two supplied covariates will be generated based on a predefined mechanisms. Missing data generation in the response variable will be based on the supplied true alpha.
- sixcitydata: Longitudinal study of health effects of air pollution using data from six cities Ware et al. (1984).
- ibrahim: Example dataset used in Ibrahim (1990, JASA).
- testyxm: Function for testing models with missing data.

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References

Firth, D. (1993). Bias reduction of maximum likelihood estimates, Biometrika, 80, 27-38. doi:10.2307/2336755.

Ibrahim, J. G. (1990). Incomplete data in generalized linear models. Journal of the American Statistical Association 85, 765–769.

Ibrahim, J. G., and Lipsitz, S. R. (1996). Parameter Estimation from Incomplete Data in Binomial Regression when the Missing Data Mechanism is Nonignorable, Biometrics, 52, 1071–1078.

Kosmidis, I., Firth, D. (2021). Jeffreys-prior penalty, finiteness and shrinkage in binomial-response generalized linear models. Biometrika, 108, 71-82. doi:10.1093/biomet/asaa052.

Louis, T. A. (1982). Finding the observed information when using the EM algorithm. Proceedings of the Royal Statistical Society, Ser B, 44, 226-233.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Pradhan, V., Nychka, D., and Bandyopadhyay, S. (2025). Addressing Missing Responses and Categorical Covariates in Binary Regression Modeling: An Integrated Framework (to be submitted).

Pradhan, V., Nychka, D., and Bandyopadhyay, S. (2025). Bridging Gaps in Logistic Regression: Tackling Missing Categorical Covariates with a New Likelihood Method (to be submitted).

Pradhan, V., Nychka, D., and Bandyopadhyay, S. (2025). glmFitMiss: Binary Regression with Missing Data in R (to be submitted).

See Also

emBinRegMAR, emBinRegMixedMAR, logRegMAR, meningitis, emforbeta, meningitis60ymis, emyxmiss, est, metastmelanoma, simulateCovariateData, est45, simulateData, felinedata, sixcitydata, ibrahim, testyxm, llkmiss

emBinRegMAR

Fitting binary regression with missing categorical covariates using Expectation-Maximisation (EM) based method

Description

This function allows users to fit generalized linear models with incomplete predictors that are categorical. The model is fitted using a likelihood-based method, which ensures reliable parameter estimation even when dealing with missing data. For more information on the underlying methodology, please refer to Pradhan, Nychka, and Bandyopadhyay (2025).

emBinRegMAR

Usage

```
emBinRegMAR(
   formula,
   data,
   conflev = 0.95,
   vcorctn = TRUE,
   family = binomial(link = "logit"),
   biascorrectn = TRUE,
   verbose = TRUE
)
```

Arguments

ial
10

Details

The family parameter in the emBinRegMAR function allows you to specify the probability distribution and link function for the response variable in the linear model. It determines the nature of the relationship between the predictors and the response variable. The family argument is particularly important when working with binary data, where the response variable has only two possible outcomes. In such cases, you typically want to fit a logistic regression model.

Currently family=binomial is supported for binary data:

You can also specify different link functions within binomial family. The default link function is the logit function, which models the log-odds of success. Other available link functions include:

- "probit" for the probit link function, which models the cumulative standard normal distribution.
- "cloglog" for the complementary log-log link function, which models the complementary log-log of the survival function.

It is important to choose the appropriate link function based on the specific characteristics and assumptions of your binary data. The default "binomial" family with the logit link function is often a good starting point, but alternative link functions might be more appropriate depending on the

emBinRegMAR

research question and the nature of the data. Note that, this function uses the function 'emforbeta' function. For more details of the function and corresponding different output objects, review the 'emforbeta' function.

Value

return the glm estimates

References

Firth, D. (1993). Bias reduction of maximum likelihood estimates, Biometrika, 80, 27-38. doi:10.2307/2336755.

Ibrahim, J. G. (1990). Incomplete data in generalized linear models. Journal of the American Statistical Association 85, 765–769.

Kosmidis, I., Firth, D. (2021). Jeffreys-prior penalty, finiteness and shrinkage in binomial-response generalized linear models. Biometrika, 108, 71-82. doi:10.1093/biomet/asaa052.

Louis, T. A. (1982). Finding the observed information when using the EM algorithm. Proceedings of the Royal Statistical Society, Ser B, 44, 226-233.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Examples

f_fit1

```
data(ibrahim)
#Fits a logistic regression mode with missing categorical covariates using Ibrahim (1990)
fit <- emBinRegMAR(y~x1+x2+x3, data=ibrahim)
fit
data(est45)
f_fit <- emBinRegMAR (resp ~ Fetoprtn + Antigen + Jaundice + Age, data = est45, biascorrectn=FALSE)
f_fit
data(est45)
f_fit <- emBinRegMAR (resp ~ Fetoprtn + Antigen + Jaundice + Age, data = est45, biascorrectn=FALSE)
f_fit
# ------Bias reduced estimates due to Firth (1993) -------
f_fit1 <- emBinRegMAR (resp ~ Fetoprtn + Antigen + Jaundice + Age, data = est45, biascorrectn=TRUE)</pre>
```

6

emBinRegMixedMAR

Fits binary regression models with both nonignorable missing responses and missing categorical covariates.

Description

This function allows users to fit generalized linear models with presence of both missing responses that are nonignorable and incomplete predictors that are categorical. The model is fitted using an EM-based method, which ensures reliable parameter estimation even when dealing with missing data. For more information on the underlying methodology, please refer to Pradhan, Nychka, and Bandyopadhyay (2025).

Usage

```
emBinRegMixedMAR(
   formula,
   data,
   conflev = 0.95,
   adtnlCovforR = NULL,
   vcorctn = TRUE,
   family = binomial(link = "logit"),
   biascorrectn = TRUE,
   verbose = TRUE
)
```

Arguments

formula	a formula expression as for regression models, of the form response ~ predic- tors. The response should be a numeric binary variable with missing values, and predictors can be any variables. A predictor with categorical values with missing can be used in the model. See the documentation of formula for other details.
data	Input data for fitting the model.
conflev	a value for the confidence interval, the default is 0.95
adtnlCovforR	an optional list of covariates to be used to fit the logistic regression logit(R) ~ response+predictors+adtnlCovforR. adtnlCovforR has to be supplied as a vector. Default is NULL.
vcorctn	a TRUE or FALSE value, by default it is FALSE. If TRUE, it calculates a vari- ance and standard error using Louis (1982). The default is vcorctn= TRUE.
family	A character string specifying the type of model family. The default is family=binomial (lin=logit).
biascorrectn	a TRUE or FALSE value, an option for bias reduced estimates due to Firth (1993). The default is biascorrectn=TRUE.
verbose	a TRUE or FALSE value, default is verbose = TRUE

Details

The family parameter in the emBinRegMixedMAR function allows you to specify the probability distribution and link function for the response variable in the linear model. It determines the nature of the relationship between the predictors and the response variable. The family argument is particularly important when working with binary data, where the response variable has only two possible outcomes. In such cases, you typically want to fit a logistic regression model.

Currently family=binomial is supported for binary data:

You can also specify different link functions within binomial family. The default link function is the logit function, which models the log-odds of success. Other available link functions include:

- "probit" for the probit link function, which models the cumulative standard normal distribution.
- "cloglog" for the complementary log-log link function, which models the complementary log-log of the survival function.

It is important to choose the appropriate link function based on the specific characteristics and assumptions of your binary data. The default "binomial" family with the logit link function is often a good starting point, but alternative link functions might be more appropriate depending on the research question and the nature of the data.

Value

return the glm estimates

References

Firth, D. (1993). Bias reduction of maximum likelihood estimates, Biometrika, 80, 27-38. doi:10.2307/2336755.

Ibrahim, J. G. (1990). Incomplete data in generalized linear models. Journal of the American Statistical Association 85, 765–769.

Kosmidis, I., Firth, D. (2021). Jeffreys-prior penalty, finiteness and shrinkage in binomial-response generalized linear models. Biometrika, 108, 71-82. doi:10.1093/biomet/asaa052.

Louis, T. A. (1982). Finding the observed information when using the EM algorithm. Proceedings of the Royal Statistical Society, Ser B, 44, 226-233.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Addressing Missing Responses and Categorical Covariates in Binary Regression Modeling: An Integrated Framework (submitted).

Examples

8

emBinRegNonIG

fit\$beta

emBinRegNonIG	Fitting binary regression with missing
	based on Ibrahim and Lipsitz (1996)

Description

This function allows users to fit binary regression models with nonignorable missing responses. The model is fitted using a likelihood-based method, which ensures reliable parameter estimation even when dealing with missing data. For more information on the underlying methodology, please refer to Pradhan, Nychka, and Bandyopadhyay (2025).

Usage

```
emBinRegNonIG(
  formula,
  data,
  conflev = 0.95,
  vcorctn = TRUE,
  biascorrectn = TRUE,
  verbose = TRUE
)
```

Arguments

formula	a formula expression as for regression models, of the form response ~ predic- tors. The response should be a numeric binary variable with missing values, and predictors can be any variables. A predictor with categorical values with missing can be used in the model. See the documentation of formula for other details.
data	Input data for fitting the model

responses that are nonignorable

conflev	a value for the confidence interval, the default is 0.95
vcorctn	a variance-covariance matrix computation using Louis (1982). Default is TRUE.
biascorrectn	a TRUE or FALSE value, an option for bias reduced estimates due to Firth (1993). The default is TRUE
verbose	a TRUE or FALSE value, default is verbose = TRUE

Details

The family parameter in the emBinRegNonIG function allows you to specify the binomial distribution and link function for the response variable in the linear model. It determines the nature of the relationship between the predictors and the response variable. The family argument is particularly important when working with binary data, where the response variable has only two possible outcomes. In such cases, you typically want to fit a binary regression model.

You can also specify different link functions within binomial family. The default link function is the logit function, which models the log-odds of success. Other available link functions include:

- "probit" for the probit link function, which models the cumulative standard normal distribution.
- "cloglog" for the complementary log-log link function, which models the complementary log-log of the survival function.

It is important to choose the appropriate link function based on the specific characteristics and assumptions of your binary data. The default "binomial" family with the logit link function is often a good starting point, but alternative link functions might be more appropriate depending on the research question and the nature of the data. Note that, this function uses the function 'emforbeta' function. For more details of the function and corresponding different output objects, review the 'emforbeta' function.

Value

return the glm estimates

References

Firth, D. (1993). Bias reduction of maximum likelihood estimates, Biometrika, 80, 27-38. doi:10.2307/2336755.

Ibrahim, J. G. (1990). Incomplete data in generalized linear models. Journal of the American Statistical Association 85, 765–769.

Ibrahim, J. G., and Lipsitz, S. R. (1996). Parameter Estimation from Incomplete Data in Binomial Regression when the Missing Data Mechanism is Nonignorable, Biometrics, 52, 1071–1078.

Kosmidis, I., Firth, D. (2021). Jeffreys-prior penalty, finiteness and shrinkage in binomial-response generalized linear models. Biometrika, 108, 71-82. doi:10.1093/biomet/asaa052.

Louis, T. A. (1982). Finding the observed information when using the EM algorithm. Proceedings of the Royal Statistical Society, Ser B, 44, 226-233.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

emforbeta

Examples

```
data(incontinence)
#Fits a binary regression mode with nonignorable missing responses using Ibrahim and Lipsitz (1996)
#biascorrectn=TRUE enables Firth type bias correction of the parameter estimates
fit <- emBinRegNonIG(y~x1+x2+x3, data=incontinence, biascorrectn=TRUE)
fit$beta
#prints the nonignorable missing mechanism
summary(fit$alpha)</pre>
```

emforbeta

Fitting binary regression with missing categorical covariates using likelihood based method

Description

This function allows users to fit generalized linear models with incomplete predictors that are categorical. The model is fitted using a likelihood-based method, which ensures reliable parameter estimation even when dealing with missing data. For more information on the underlying methodology, please refer to Pradhan, Nychka, and Bandyopadhyay (2025).

Usage

```
emforbeta(
   formula,
   data,
   family = "binomial",
   vcorctn = FALSE,
   method = "glm.fit",
   NIterations = 50,
   verbose = FALSE,
   theta = NULL,
   convergenceCriterion = 1e-04,
   augmented = NULL,
   VarWithMissingVal = NULL
)
```

Arguments

formula	a formula expression as for regression models, of the form response ~ predic- tors. The response should be a numeric binary variable with missing values, and predictors can be any variables. A predictor with categorical values with missing can be used in the model. See the documentation of formula for other details.
data	Input data for fitting the model
family	a character string specifying the type of model family. The default is fam- ily=binomial (lin=logit)

vcorctn	a TRUE or FALSE value, by default it is FALSE. If TRUE, it calculates a variance and standard error using Louis (1982)
method	a method="brglmFit" or method="glm.fit" will be used for fitting model. The method="brglmFit" fits generalized linear models using bias reduction methods (Kosmidis, 2014), and other penalized maximum likelihood methods.The deaf- ult option method="glm.fit" fits regression with generalized linear models.
NIterations	is the number of iterations to be used for convergence. The default is NIterations= 50
verbose	a TRUE or FALSE value, by default it is FALSE. A value TRUE prints all intermediate computational details
theta	a vector containing multinomial parameters that sums to 1, default is NULL
convergenceCri	terion
	Convergence criteria to be used for convergence. The default is 1e-4
augmented	is the name of an augmented data. The default is NULL
VarWithMissing ^v	Val
	is a vector of variables including missing values. The default is NULL

is a vector of variables including missing values. The default is NULL

Details

The family parameter in the emforbeta function allows you to specify the probability distribution and link function for the response variable in the linear model. It determines the nature of the relationship between the predictors and the response variable. The family argument is particularly important when working with binary data, where the response variable has only two possible outcomes. In such cases, you typically want to fit a logistic regression model.

The following commonly used families are supported for binary data:

"binomial" for a binomial distribution, suitable for binary or dichotomous response variables.

You can also specify different link functions within binomial family. The default link function is the logit function, which models the log-odds of success. Other available link functions include:

- "probit" for the probit link function, which models the cumulative standard normal distribution.
- "cloglog" for the complementary log-log link function, which models the complementary log-log of the survival function.

It is important to choose the appropriate link function based on the specific characteristics and assumptions of your binary data. The default "binomial" family with the logit link function is often a good starting point, but alternative link functions might be more appropriate depending on the research question and the nature of the data. See also the function 'emBinRegMAR' function.

Value

return the glm estimates

emforbeta

References

Firth, D. (1993). Bias reduction of maximum likelihood estimates, Biometrika, 80, 27-38. doi:10.2307/2336755.

Ibrahim, J. G. (1990). Incomplete data in generalized linear models. Journal of the American Statistical Association 85, 765–769.

Kosmidis, I., Firth, D. (2021). Jeffreys-prior penalty, finiteness and shrinkage in binomial-response generalized linear models. Biometrika, 108, 71-82. doi:10.1093/biomet/asaa052.

Louis, T. A. (1982). Finding the observed information when using the EM algorithm. Proceedings of the Royal Statistical Society, Ser B, 44, 226-233.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Examples

```
data(sixcitydata)
f_fit <- emforbeta(Wheeze~city+soc+cond,</pre>
                   data=sixcitydata,
                    vcorctn= TRUE,
                    family=binomial(link="logit"),
                   method="glm.fit")
summary(f_fit$mfit) #creates the summary like glm using the return object mfit
vcov_beta<-f_fit$cvcov #creates variance using Louis (1982)</pre>
# Computes the standard error of the estimates
se_beta_em<-sqrt(diag(vcov_beta))</pre>
se_beta_em
# Firth correction
f_fit <- emforbeta(Wheeze~city+soc+cond,</pre>
                    data=sixcitydata,
                    family=binomial(link="logit"),
                   method="brglmFit")
# creates the summary like glm using the return object mfit
data(ibrahim)
f_fit2 <- emforbeta(y~x1+x2+x3,</pre>
                    data=ibrahim,
                    family="binomial")
summary(f_fit2$mfit) #creates the summary like glm using the return object mfit
f_fit2 <- emforbeta(y~x1+x2+x3,</pre>
                     data=ibrahim,
                     family=binomial (link="probit"),
                    method="brglmFit")
# creates the summary like glm using the return object mfit
summary(f_fit2$mfit) #
data(est)
```

```
f_fit <- emforbeta(survive~Fetoprtn+Antigen+Jaundice+Age,</pre>
                    data=est,
                    family=binomial,
                   method="glm.fit")
summary(f_fit$mfit)
f_fit <- emforbeta(survive~Fetoprtn+Antigen+Jaundice+Age,</pre>
                    data=est,
                    family=binomial,
                   method="brglmFit")
# Firth corrected estimates with out Louis (1982) correction (see Maiti and Pradhan (2009))
summary(f_fit$mfit)
data(metastmelanoma)
f_fit <- emforbeta(failcens~size+type+nodal+age+sex+trt,</pre>
                    data=metastmelanoma,
                    family=binomial,
                    method="glm.fit")
summary(f_fit$mfit)
f_fit <- emforbeta(failcens~size+type+nodal+age+sex+trt,</pre>
                    data=metastmelanoma,
                    family=binomial,
                    method="brglmFit")
# Firth corrected estimates with out Louis (1982) correction (see Maiti and Pradhan (2009))
summary(f_fit$mfit)
data(felinedata)
f_fit <- emforbeta(chlamy~Season+Agegrp+Conj+FHV1,</pre>
                    data=felinedata,
                    family=binomial,
                    method="glm.fit")
summary(f_fit$mfit)
f_fit <- emforbeta(chlamy~Season+Agegrp+Conj+FHV1,</pre>
                    data=felinedata,
                    family=binomial,
                   method="brglmFit")
# Firth corrected estimates with out Louis (1982) correction
summary(f_fit$mfit)
```

emil

Fitting binary regression model with missing responses based on Ibrahim and Lipsitz (1996)

Description

This function enables users to fit generalized linear models when handling incomplete data in the response variable. The missing responses are assumed to be nonignorable. The model is fitted using

emil

emil

a novel likelihood-based method proposed by Ibrahim and Lipsitz(1996).

Usage

```
emil(
   formula,
   data,
   adtnlCovforR = NULL,
   eps0 = 1e-05,
   maxit = 75,
   family = "binomial",
   method = "brglmFit"
)
```

Arguments

formula	a formula expression as for regression models, of the form response ~ predictors. The response should be a numeric binary variable with missing values, and pre- dictors can be any variables. A predictor with categorical values with missing can be used in the model. See the documentation of formula for other details.
data	an optional data frame in which to interpret the variables occurring in formula.
adtnlCovforR	an optional list of covariates to be used to fit the logistic regression logit(R) ~ response+predictors+adtnlCovforR. adtnlCovforR has to be supplied as a vector. Default is NULL.
eps0	arguments to be used to for the convergence criteria of the maximum likelihood computation of the joint likelihood function. The default is 1e-3.
maxit	arguments to be used to for the maximization of the joint likelihood function. The default is 50.
family	A character string specifying the type of model family.
method	a method="brglmFit" or method="glm.fit" will be used for fitting model. The method="brglmFit" fits generalized linear models using bias reduction methods (Kosmidis, 2014), and other penalized maximum likelihood methods.

Details

The family parameter in the emil function allows you to specify the probability distribution and link function for the response variable in the linear model. It determines the nature of the relationship between the predictors and the response variable. The family argument is particularly important when working with binary data, where the response variable has only two possible outcomes. In such cases, you typically want to fit a binary regression model with an appropriate link.

Currently the package only supports family=binomial for binary or dichotomous response variables.

You can also specify different link functions within the family=binomial. The default link function is the logit function, which models the log-odds of success. Other available link functions include:

• "probit" for the probit link function, which models the cumulative standard normal distribution. "cloglog" for the complementary log-log link function, which models the complementary loglog of the survival function.

It is important to choose the appropriate family and link function based on the specific characteristics and assumptions of your binary data. The default "binomial" family with the logit link function is often a good starting point, but alternative link functions might be more appropriate depending on the research question and the nature of the data.

Value

return the generalized linear model estimates

References

Firth, D. (1993). Bias reduction of maximum likelihood estimates, Biometrika, 80, 27-38. doi:10.2307/2336755.

Ibrahim, J. G. (1990). Incomplete data in generalized linear models. Journal of the American Statistical Association 85, 765–769.

Ibrahim, J. G., and Lipsitz, S. R. (1996). Parameter Estimation from Incomplete Data in Binomial Regression when the Missing Data Mechanism is Nonignorable, Biometrics, 52, 1071–1078.

Kosmidis, I., Firth, D. (2021). Jeffreys-prior penalty, finiteness and shrinkage in binomial-response generalized linear models. Biometrika, 108, 71-82. doi:10.1093/biomet/asaa052.

Louis, T. A. (1982). Finding the observed information when using the EM algorithm. Proceedings of the Royal Statistical Society, Ser B, 44, 226-233.

Maity, A., Pradhan, V., Das U (2019). Bias reduction in logistic regression with missing responses when the missing data mechanism is nonignorable. The American Statistician, (73) 340-349.

Pradhan V, Nychka DW, Bandyopadhyay S (2025). Addressing Missing Responses and Categorical Covariates in Binary Regression Modeling: An Integrated Framework (to be submitted).

Examples

emyxmiss

```
Fitting generalized linear models with Incomplete data
```

Description

This function enables users to fit generalized linear models when handling incomplete data in both the response variable and categorical covariates. The missing responses are assumed to be nonignorable, while missing categorical covariates are assumed to be missing at random. The model is fitted using a novel likelihood-based method proposed by Pradhan, Nychka, and Bandyopadhyay (2025).

emyxmiss

Usage

```
emyxmiss(
  formula,
  data,
  adtnlCovforR = NULL,
  eps0 = 0.001,
  maxit = 75,
  family = "binomial",
  method = "glm.fit"
)
```

Arguments

formula	a formula expression as for regression models, of the form response ~ predictors. The response should be a numeric binary variable with missing values, and pre- dictors can be any variables. A predictor with categorical values with missing can be used in the model. See the documentation of formula for other details.
data	an optional data frame in which to interpret the variables occurring in formula.
adtnlCovforR	an optional list of covariates to be used to fit the logistic regression logit(R) ~ response+predictors+adtnlCovforR. adtnlCovforR has to be supplied as a vector. Default is NULL.
eps0	arguments to be used to for the convergence criteria of the maximum likelihood computation of the joint likelihood function. The default is 1e-3.
maxit	arguments to be used to for the maximization of the joint likelihood function. The default is 50.
family	A character string specifying the type of model family.
method	a method="brglmFit" or method="glm.fit" will be used for fitting model. The method="brglmFit" fits generalized linear models using bias reduction methods (Kosmidis, 2014), and other penalized maximum likelihood methods.

Details

The family parameter in the emyxmmiss function allows you to specify the probability distribution and link function for the response variable in the linear model. It determines the nature of the relationship between the predictors and the response variable. The family argument is particularly important when working with binary data, where the response variable has only two possible outcomes. In such cases, you typically want to fit a logistic regression model.

The following commonly used families are supported for binary data:

• "binomial" for a binomial distribution, suitable for binary or dichotomous response variables.

You can also specify different link functions within each family. For the "binomial" family, the default link function is the logit function, which models the log-odds of success. Other available link functions include:

• "probit" for the probit link function, which models the cumulative standard normal distribution. "cloglog" for the complementary log-log link function, which models the complementary log-log of the survival function.

It is important to choose the appropriate family and link function based on the specific characteristics and assumptions of your binary data. The default "binomial" family with the logit link function is often a good starting point, but alternative link functions might be more appropriate depending on the research question and the nature of the data.

Value

return the generalized linear model estimates

References

Firth, D. (1993). Bias reduction of maximum likelihood estimates, Biometrika, 80, 27-38. doi:10.2307/2336755.

Ibrahim, J. G. (1990). Incomplete data in generalized linear models. Journal of the American Statistical Association 85, 765–769.

Ibrahim, J. G., and Lipsitz, S. R. (1996). Parameter Estimation from Incom- plete Data in Binomial Regression when the Missing Data Mechanism is Nonignorable, Biometrics, 52, 1071–1078.

Kosmidis, I., Firth, D. (2021). Jeffreys-prior penalty, finiteness and shrinkage in binomial-response generalized linear models. Biometrika, 108, 71-82. doi:10.1093/biomet/asaa052.

Louis, T. A. (1982). Finding the observed information when using the EM algorithm. Proceedings of the Royal Statistical Society, Ser B, 44, 226-233.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan V, Nychka DW, Bandyopadhyay S (2025). Addressing Missing Responses and Categorical Covariates in Binary Regression Modeling: An Integrated Framework (to be submitted).

Examples

Firth correction and link=probit

```
fit_yx <- emyxmiss(Wheeze ~ city + soc + cond,</pre>
                    data = dataWithMiss,
                    adtnlCovforR = c("age"),
                     family = binomial(link = "probit"),
                    method = "brglmFit")
fit_yx
# on simulated data
demo_df <- simulateCovariateData(50, nCov=6)</pre>
simulated_df <- simulateData(demo_df)</pre>
testMissData <- simulated_df$dataMissing</pre>
fit_yx <- emyxmiss(y~x2+x3+x4,</pre>
                     data=testMissData,
                    adtnlCovforR=c("x1"),
                     family=binomial,
                    method="glm.fit")
fit_yx
summary(fit_yx$fit_y)
```

est

EST data – Eastern Cooperative Oncology Group clinical trials, EST 2282

Description

The dataset est is from the Eastern Cooperative Oncology Group clinical trials, specifically EST 2282 (Falkson, Cnaan, and Simson, 1990) and EST 1286 (Falkson et al., 1995). The dataset consists of 191 observations. It includes several covariates: Fetoprtn (alpha fetoprotein), Antigen (antihepatitis B antigen), Jaundice (a biochemical marker; coded as 1 if present, 0 otherwise), and Age (age in years). The response variable Y represents the number of cancerous liver cells present at the start of the clinical trial.

To assess the impact of these covariates on the likelihood of survival, a new variable called "survive" is created. "survive" is dichotomized based on Y: it is set to 1 if the number of cancerous liver cells is less than or equal to 8, and 0 otherwise.

Maiti and Pradhan (2009) fitted a logistic regression using the model survive ~ Fetoprtn + Antigen + Jaundice + Age. This model explores the relationship between the covariates and the likelihood of survival for patients in the clinical trials.

Usage

est

Format

A data frame with 191 rows and several variables:

Y Response variable representing the number of cancerous liver cells

Fetoprtn Alpha fetoprotein
BMI Body Mass Index
Antigen Anti-hepatitis B antigen
Jaundice Jaundice indicator, coded as 1 if present, 0 otherwise
Age Age in years
Weeks times in weeks
survive Dichotomized survival variable based on Y

Source

Generated for example purposes

References

Cytel Inc (2010). LogXact 9 User Manual: Discrete Regression Analysis. Cambridge, Massachusetts: Cytel Inc.

Falkson, G., Lipsitz, S., Borden, E., Simson, I., W., and Haller, D. (1995). A ECOG randomized phase II study of beta interferon and Menogoril. American Journal of Clinical Oncology 18, 287–292.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Examples

est45

EST data – Eastern Cooperative Oncology Group clinical trials, EST 2282

est45

Description

The dataset est45 is from the Eastern Cooperative Oncology Group clinical trials, specifically EST 2282 (Falkson, Cnaan, and Simson, 1990) and EST 1286 (Falkson et al., 1995) containing 45 observations. The dataset consists of 191 observations. It includes several covariates: Fetoprtn (alpha fetoprotein), Antigen (antihepatitis B antigen), Jaundice (a biochemical marker; coded as 1 if present, 0 otherwise), and Age (age in years). The response variable Y represents the number of cancerous liver cells present at the start of the clinical trial.

To assess the impact of these covariates on the likelihood of survival, a new variable called "survive" is created. "survive" is dichotomized based on Y: it is set to 1 if the number of cancerous liver cells is less than or equal to 8, and 0 otherwise.

Usage

est45

Format

A data frame with 45 rows and 9 variables:

Y Response variable
Weeks Time in weeks
Fetoprtn Alpha fetoprotein
Antigen Anti-hepatitis B antigen
Jaundice Jaundice indicator
BMI Body mass index
Age Age in years
grp Group identifier
resp Response variable dichotomized

Source

Generated for example purposes

References

Cytel Inc (2010). LogXact 9 User Manual: Discrete Regression Analysis. Cambridge, Massachusetts: Cytel Inc.

Falkson, G., Lipsitz, S., Borden, E., Simson, I., W., and Haller, D. (1995). A ECOG randomized phase II study of beta interferon and Menogoril. American Journal of Clinical Oncology 18, 287–292.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Examples

felinedata

felinedata - Chlamydial Infection in Cats

Description

In a study conducted by Sykes et al. (1999), the risk factors for Chlamy, a chlamydial infection in cats, were investigated. The analysis considered important variables such as FHV1 (Herpes virus infection), Season, Conjunctivitis (Conj), and Age group. Season was coded from 1 to 4 to represent the seasons, FHV1 was binary (1 for infected cats, 0 for non-infected cats), Conj was binary (1 if present, 0 if absent), and Age group was categorized into specific ranges. The original dataset had 462 observations, with around 20% missing values. After removing missing values, the analysis was conducted with a sample size of 371. The fitted model included Chlamy as the outcome variable, with FHV1, Season, Conj, and Age group as predictors, treating Age group and Season as class variables with a base value of 1.

Usage

felinedata

Format

An object of class tbl_df (inherits from tbl, data.frame) with 120 rows and 6 columns.

References

Cytel Inc (2010). LogXact 9 User Manual: Discrete Regression Analysis. Cambridge, Massachusetts: Cytel Inc.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2024). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Sykes, J. E., Anderson, G. A., Studdert, V. P., and Browning, G. F. (1999). Prevalence of feline Chlamydia psittaci and feline her- pesvirus 1 in cats with upper respiratory tract disease. Journal of Veterinary Internal Medicine 13, 153–162.

22

ibrahim

Examples

ibrahim

ibrahim data – Ibrahim (1990) JASA

Description

The dataset ibrahim is from Ibrahim, IG (1990, 85, 765–769, JASA). The data contains a response variable y and predictors x1 x2 x3, and the total number of observations is 82.

Usage

ibrahim

Format

An object of class tbl_df (inherits from tbl, data.frame) with 82 rows and 4 columns.

References

Cytel Inc (2010). LogXact 9 User Manual: Discrete Regression Analysis. Cambridge, Massachusetts: Cytel Inc.

Ibrahim, J. G. (1990). Incomplete data in generalized linear models. Journal of the American Statistical Association 85, 765–769.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Examples

```
data(ibrahim)
f_fit <- emBinRegMAR(y ~ x1+x2+x3, data=ibrahim, family="binomial", biascorrectn=FALSE)
f_fit$beta
#Firth type bias correction
f_fit <- emBinRegMAR(y ~ x1+x2+x3, data=ibrahim, family="binomial", biascorrectn=TRUE)
f_fit$beta</pre>
```

incontinence incontinence- incontinence Data taken from brlrmr pacakge

Description

The dataset incontinence is from. The dataset is available in the brlrmr pacakge. Pradhan, Nychka and Bandyopadhyay (2024) fitted the model $y \sim x1 + x2 + x3$.

Usage

incontinence

Format

A data frame with several rows and columns representing various variables:

- y response variable
- x1 is a covariate
- x2 is a covariate
- x3 is a covariate

References

Maity, A., Pradhan, V., Das, U. (2019). Bias reduction in logistic regression with missing responses when the missing data mechanism is nonignorable. The American Statistician, (73) 340-349.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2024). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Examples

```
fit <- emil(y~x1+x2+x3, data=incontinence,family=binomial, method="brglmFit")
# display summary of the beta estimates of the model
summary(fit$fit_y)
# for non-ignorability setting of the missing responses
summary(fit$fit_r)</pre>
```

24

llkmiss

Fitting binary regression with missing categorical covariates using new likelihood based method that does not require EM algorithm

Description

This function allows users to fit logistic regression models with incomplete predictors that are categorical. The model is fitted using a new likelihood-based method, which ensures reliable parameter estimation even when dealing with missing data. For more information on the underlying methodology, please refer to Pradhan, Nychka, and Bandyopadhyay (2025).

Usage

llkmiss(par, data, formula, augData, biasCorr = TRUE)

Arguments

par	A vector including a list of parameters to be estimated. This include the beta (the regression parameters) and theta, the multinomial paraters for observing a missing covaraite pattern.
data	Input data for fitting the model
formula	A formula expression as for regression models, of the form response ~ predictors. The response should be a numeric binary variable with missing values, and pre- dictors can be any variables. A predictor with categorical values with missing can be used in the model. See the documentation of formula for other details.
augData	An augmented data including all possible covarites that could have been observed.
biasCorr	a TRUE or FALSE value, by default it is TRUE.

Value

return the regression estimates

References

Firth, D. (1993). Bias reduction of maximum likelihood estimates, Biometrika, 80, 27-38. doi:10.2307/2336755.

Kosmidis, I., Firth, D. (2021). Jeffreys-prior penalty, finiteness and shrinkage in binomial-response generalized linear models. Biometrika, 108, 71-82. doi:10.1093/biomet/asaa052.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Bridging Gaps in Logistic Regression: Tackling Missing Categorical Covariates with a New Likelihood Method (to be submitted).

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). glmFitMiss: Binary Regression with Missing Data in R (to be submitted)

logRegMAR

Fitting binary regression with missing categorical covariates using new likelihood based method

Description

This function allows users to fit logistic regression models with incomplete predictors that are categorical. The model is fitted using a new likelihood-based method, which ensures reliable parameter estimation even when dealing with missing data. For more information on the underlying methodology, please refer to Pradhan, Nychka, and Bandyopadhyay (2024).

Usage

```
logRegMAR(formula, data, conflev = 0.95, correctn = TRUE, verbose = TRUE)
```

Arguments

formula	A formula expression as for regression models, of the form response ~ predictors. The response should be a numeric binary variable with missing values, and pre- dictors can be any variables. A predictor with categorical values with missing can be used in the model. See the documentation of formula for other details.
data	Input data for fitting the model
conflev	Confidence level, the default is 0.95
correctn	a TRUE or FALSE value, by default it is TRUE.
verbose	a TRUE or FALSE value, default is verbose = TRUE

Value

return the logistic regression estimates

References

Firth, D. (1993). Bias reduction of maximum likelihood estimates, Biometrika, 80, 27-38. doi:10.2307/2336755.

Kosmidis, I., Firth, D. (2021). Jeffreys-prior penalty, finiteness and shrinkage in binomial-response generalized linear models. Biometrika, 108, 71-82. doi:10.1093/biomet/asaa052.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Bridging Gaps in Logistic Regression: Tackling Missing Categorical Covariates with a New Likelihood Method (to be submitted).

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). glmFitMiss: Binary Regression with Missing Data in R (to be submitted)

meningitis

Examples

meningitis

meningitis- Meningococcal Disease Data with missing data in the response variable

Description

The dataset meningitis is from a brief outbreak of meningococcal disease at the University of Illinois, Urbana-Champaign campus in the years 1991 and 1992. The dataset is available in the LogXact software and also analyzed in Imrey et al. (1996). Maiti and Pradhan (2009) fitted a logistic regression using the model CaseCntrl ~ Numill + Numsleep + Smoke + Set + Reftime.

Usage

meningitis

Format

A data frame with several rows and columns representing various variables:

CaseCntrl Case control status Numnill Number of illnesses Numsleep Number of sleep disturbances Smoke Smoking status Set Set variable Reftime Reference time

References

Cytel Inc (2010). LogXact 9 User Manual: Discrete Regression Analysis. Cambridge, Massachusetts: Cytel Inc.

Imrey, P. B., Jackson, L. A., Ludwinski, P. H., England, A. C. II, Fox, B. C., Isdale, L. B., Reeves, M. W., and Wenger, J. D. (1996). Outbreak of serogroup C meningococcal disease associated with campus bar patronage. American Journal of Epidemiology 143, 624–630.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2024). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Examples

meningitis60ymis meningitis60ymis- Meningococcal Disease Data with missing data in the response variable

Description

The dataset meningitis is from a brief outbreak of meningococcal disease at the University of Illinois, Urbana-Champaign campus in the years 1991 and 1992. The dataset is available in the LogXact software and also analyzed in Imrey et al. (1996). Pradhan, Nychka and Bandyopadhyay (2024) fitted the model resp~Numnill+Numsleep+Smoke+Set+Reftime, where the the response variable resp included missing value.

Usage

```
meningitis60ymis
```

Format

A data frame with several rows and columns representing various variables:

CaseCntrl Case control status

Numnill Number of illnesses

Numsleep Number of sleep disturbances

metastmelanoma

R an indicator variable for missing CaseCntrl

Smoke Smoking status

Set Set variable

m indicator variable for observations with missing values

Reftime Reference time

resp Response variable with missing data for the varaible CaseCntrl

References

Cytel Inc (2010). LogXact 9 User Manual: Discrete Regression Analysis. Cambridge, Massachusetts: Cytel Inc.

Imrey, P. B., Jackson, L. A., Ludwinski, P. H., England, A. C. II, Fox, B. C., Isdale, L. B., Reeves, M. W., and Wenger, J. D. (1996). Outbreak of serogroup C meningococcal disease associated with campus bar patronage. American Journal of Epidemiology 143, 624–630.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Examples

metastmelanoma metastmelanoma - metastatic melanoma trial data

Description

The dataset data from a cancer clinical trial and the results are published in Kirkwood et al. (1996). In this study following surgery for deep primary or metastatic melanoma, the overall survival and the disease-free effect of Interferon alpha-2b (IFN) was investigated on 285 patients. Maiti and Pradhan (2009) fitted a logistic regression considering failcens as the response variable, where failcens is 1 if the subject relapses and 0 otherwise. We fit the model including six important predictors size type nodal age sex trt; where size is the size of primary in cm2, which is dichotomized at the median; type is the type of primary containing two levels— superficial spreading and other; nodal is the presence or absence of microscopic nonpalpable and palpable regional lymph node metastasis—1 for node positive and 0 otherwise; age is the age of a subject in years; sex is a variable indicating the gender (male or female); and finally trt is the treatment containing two levels (1 if treated with IFN and 0 otherwise).

Usage

metastmelanoma

Format

An object of class tbl_df (inherits from tbl, data.frame) with 285 rows and 11 columns.

References

Cytel Inc (2010). LogXact 9 User Manual: Discrete Regression Analysis. Cambridge, Massachusetts: Cytel Inc.

Kirkwood, J. M., Strawderman, M. H., Ernstoff, M. S., Smith, T. J., Borden, E. C., and Blum, R. H. (1996). Interferon alfa-2b ad- juvant therapy of high-risk resected cutaneous melanoma: The Eastern Cooperative Oncology Group trial EST 1684. Journal of Clinical Oncology 14, 7–17.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Examples

```
data(metastmelanoma)
f_fit <- emforbeta(failcens ~ size+type+nodal+age+sex+trt,</pre>
                    data=metastmelanoma,
                    family=binomial, method="glm.fit")
summary(f_fit$mfit)
vcov_beta<-f_fit$cvcov # variance-covariance calculation using Louis (1982)</pre>
vcov_beta
se_beta_em<-sqrt(diag(vcov_beta))</pre>
se_beta_em
# Firth Correction
f_fit <- emforbeta(failcens ~ size+type+nodal+age+sex+trt,</pre>
                    data=metastmelanoma,
                    family=binomial, method="brglmFit")
summary(f_fit$mfit)
vcov_beta<-f_fit$cvcov # variance-covariance calculation using Louis (1982)</pre>
vcov_beta
se_beta_em<-sqrt(diag(vcov_beta))</pre>
se_beta_em
```

simulateCovariateData Simulate data with independent categorical covariates

simulateData

Description

This function generates a simulated data with independent categorical covariates. The first two covariates namely x1 and x2 are generated using random normal rnorm(n, 40, 20) and random poisson rpois(n, lambda = 4). The remaining covariates are generated at random with categories 0,1,2.

Usage

```
simulateCovariateData(n, nCov = 2)
```

Arguments

n	numner of observations to be generated for the data
nCov	4+ number of covariates to be generated for the data, the first 4 covariates gen- erated based on pre-specified distributions

Value

returns a data frame with covariates x1, x2, ...

Examples

simulateCovariateData(10, nCov=15)

simulateData

Simulate data based on an input covariate data

Description

This function generates missing data both in the response variables as well as in the predictors. The missing data generation in the last two supplied covariates will be generated based on a predefined mechanisms. Missing data generation in the response variable will be based on the supplied true alpha.

Usage

```
simulateData(
    dataCov,
    truebeta = c(1, -1, 1, 5),
    truealpha = c(-1, 5, -1, -1, 0.01),
    nsim = 2
)
```

Arguments

dataCov	input data, the default number of covariates is 7 (5+2)
truebeta	the beta parameter to be used to generate binary response values $1/0 \operatorname{slogit}(y=1)=x1+x2+x3$
truealpha	to be used to generate nonignorable missing values based on the model $logit(R=1)=y+x1+x2+x3+x4+$
nsim	number of simulated dataset, default is 2

Value

returns a list with original data called originalData and a data with imputed missing values dataMissing

Examples

```
demo_df <- simulateCovariateData(100, nCov=6)
simulated_df <- simulateData(demo_df, nsim=2)
testMissData <- simulated_df$dataMissing
head(testMissData)</pre>
```

simulateMissDfYorX Simulate missing covariate or missing responses data based on an input covariate data

Description

This function generates missing covariate or missing responses data. The missing data generation in the last two supplied covariates will be generated based on a predefined mechanisms. Missing data generation in the response variable will be based on the supplied true alpha.

Usage

```
simulateMissDfYorX(
    dataCov,
    truebeta = c(1, -1, 1, 5),
    truealpha = c(-1, 5, -1, -1, -1, 0.01),
    x2Mar = c(1, -1, -1),
    ymiss = FALSE,
    nsim = 1
)
```

Arguments

dataCov	input data, the default number of covariates is 7 (5+2)
truebeta	the beta parameter to be used to generate binary responses $1/0 \operatorname{slogit}(y=1)=x1+x2+x3$
truealpha	to be used to generate nonignorable missing values based on the model $logit(R=1)=y+x1+x2+x3+x4+.$
x2Mar	to be used to generate missing values in x2 based on the model logit(x2=missing)=x1+y
ymiss	to be used for missing responses, default is FALSE
nsim	number of simulated dataset, default is 2

32

sixcitydata

Value

returns a list with original data called originalData and a data with imputed missing values dataMissing

Examples

```
demo_df <- simulateCovariateData(100, nCov=6)
simulated_df <- simulateMissDfYorX(demo_df, nsim=2)
testMissData <- simulated_df$dataMissing
head(testMissData)</pre>
```

ery well published Six city data published in many
Ware et al (1984), Ibrahim and Lipsitz (1996). Also
<i>Cact User Manual. The dataset is a longitudinal study</i>
ets of air pollution (ware et al., 1984).

Description

The 'sixcitydata' dataset contains information on wheezing status, city of residence, maternal smoking habits, socioeconomic status, and medical condition of children at age 11.

The dataset includes the following variables:

- Wheeze: Binary response variable indicating wheezing status of children at age 11 (1 for wheeze, 0 for no wheeze).
- city: Categorical variable indicating city of residence (1 for polluted city, 0 for Portage or Wisconsin).
- smoke: Binary variable indicating mother's smoking habits (1 for >20 cigarettes a day, 0 otherwise).
- soc: Binary variable indicating high socioeconomic status of subject (1 for high socioeconomic status, 0 otherwise).
- cond: Binary variable indicating previous medical condition of subject (1 for previous medical condition, 0 otherwise).

Usage

```
sixcitydata
```

Format

An object of class tbl_df (inherits from tbl, data.frame) with 2106 rows and 5 columns.

References

Cytel Inc (2010). LogXact 9 User Manual: Discrete Regression Analysis. Cambridge, Massachusetts: Cytel Inc.

Ibrahim, J. G. (1990). Incomplete data in generalized linear models. Journal of the American Statistical Association 85, 765–769.

Maiti, T., Pradhan, V. (2009). Bias reduction and a solution of separation of logistic regression with missing covariates. Biometrics, 65, 1262-1269.

Pradhan, V., Nychka, D. and Bandyopadhyay, S. (2025). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Ware, JH., Dockery, DW., Spiro, A III., Speizer, FE., and Ferris, BG Jr. (1984). Passive smoking, gas cooking, and respiratory health of children living in sex cities. American Review of Respiratory Disease, 129, 366-374.

Examples

testyxm

Simulated Test Data – testyxm

Description

A test list data that returns a list called testyxm with two components:

- dataOriginal: The original data set without missing values.
- dataMissing: The data set with artificially introduced missing values.

Usage

data(testyxm)

Format

A list with the following components:

dataOriginal A data frame with several rows and columns representing various variables. **dataMissing** A data frame with missing values corresponding to the same structure as dataOriginal.

testyxm

Details

Simulated Test Data

This dataset is a list called testyxm that contains two data frames: dataOriginal and dataMissing.

References

Pradhan, V., Nychka, D., and Bandyopadhyay, S. (2024). Beyond the Odds: Fitting Logistic Regression with Missing Data in Small Samples (submitted).

Examples

data(testyxm) Fulldata <- testyxm\$dataOriginal Missdata <- testyxm\$dataMissing

Index

* datasets est, 19 est45, 20 felinedata, 22 ibrahim, 23 incontinence, 24 meningitis, 27 meningitis60ymis, 28 metastmelanoma, 29 sixcitydata, 33 testyxm, 34 * package glmfitmiss-package, 2 emBinRegMAR, 2, 4, 4 emBinRegMixedMAR, 2, 4, 7 emBinRegNonIG, 2, 9 emforbeta, *3*, *4*, 11 emil, 14 emyxmiss, *3*, *4*, 16 est, 3, 4, 19 est45, *3*, *4*, 20 felinedata, *3*, *4*, 22 glmfitmiss(glmfitmiss-package), 2 glmFitMiss-package (glmfitmiss-package), 2 glmfitmiss-package, 2 ibrahim, *3*, *4*, 23 incontinence, 24 11kmiss, *3*, *4*, 25 logRegMAR, *3*, *4*, 26 meningitis, *3*, *4*, 27

meningitis60ymis, *3*, *4*, 28 metastmelanoma, *3*, *4*, 29

simulateCovariateData, 3, 4, 30

simulateData, 3, 4, 31
simulateMissDfYorX, 3, 32
sixcitydata, 3, 4, 33

testyxm, *3*, *4*, 34