# Package 'CRTgeeDR'

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# Description

The CRTgeeDR package allows you to estimates parameters in a regression model (with possibly a link function). It allows treatment augmentation and IPW for missing data alone.

#### **Details**

The only function you're likely to need from **CRTgeeDR** is geeDREstimation. Otherwise refer to the help documentation.

data.sim

The data.sim Dataset.

# Description

HIV risk of infection after STI/HIV intervention in a cluster randomized trial.

#### **Format**

A data frame with 10000 rows and 8 variables

#### **Details**

A dataset containing the HIV risk scores and presence of risky behaviors (yes/no) and other covarites of 10000 subjects among 100 communities. The variables are as follows:

- · IDPAT subject id
- CLUSTER cluster id
- TRT treatment status, 1 is received STI/HIV intervention
- X1 A covariate following a N(0,1)
- JOB employement status
- MARRIED marital status

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- AGE age
- HIV.KNOW Score for HIV knowlege
- RELIGION religiosity score
- OUTCOME Binary outcome 1 if the subject is at high risk of HIV infection, 0 otherwise. NA if missing.
- MISSING 1 if the ouctome is missing 0 otherwise.

fitted.CRTgeeDR

Fit CRTgeeDR object.

#### **Description**

Fit CRTgeeDR object to a dataset

#### Usage

```
## S3 method for class 'CRTgeeDR'
fitted(object, ...)
```

#### **Arguments**

```
object CRTgeeDR object ignored
```

geeDREstimation

Doubly Robust Inverse Probability Weighted Augmented GEE Estimator

## Description

This function implements a GEE estimator. It implements classical GEE, IPW-GEE, augmented GEE and IPW-Augmented GEE (Doubly robust).

#### Usage

```
geeDREstimation(formula, id, data = parent.frame(), family = gaussian,
  corstr = "independence", Mv = 1, weights = NULL, aug = NULL,
  pi.a = 1/2, corr.mat = NULL, init.beta = NULL, init.alpha = NULL,
  init.phi = 1, scale.fix = FALSE, sandwich = TRUE, maxit = 20,
  tol = 1e-05, print.log = FALSE, typeweights = "VW", nameTRT = "TRT",
  model.weights = NULL, model.augmentation.trt = NULL,
  model.augmentation.ctrl = NULL, stepwise.augmentation = FALSE,
  stepwise.weights = FALSE, nameMISS = "MISSING", nameY = "OUTCOME",
  sandwich.nuisance = FALSE, fay.adjustment = FALSE, fay.bound = 0.75)
```

Arguments		

id

formula an object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted.

a vector which identifies the clusters. The length of "id" should be the same as the number of observations. Data are assumed to be sorted so that observations

on a cluster are contiguous rows for all entities in the formula.

an optional data frame, list or environment (or object coercible by as.data.frame to a data frame) containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from

which CRTgeeDR is called.

family a description of the error distribution and link function to be used in the model.

This can be a character string naming a family function, a family function or the result of a call to a family function. (See family for details of family functions.)

corstr a character string specifying the correlation structure. The following are permit-

ted: '"independence"', '"exchangeable"', '"ar1"', '"unstructured"' and '"userde-

fined"

Mv for "m-dependent", the value for m

weights A vector of weights for each observation. If an observation has weight 0, it

is excluded from the calculations of any parameters. Observations with a NA anywhere (even in variables not included in the model) will be assigned a weight

of 0.

aug A list of vector (one for A=1 treated, one for A=0 control) for each observation

representing E(Y|X,A=a).

pi.a A number, Probability of treatment attribution P(A=1)

corr . mat The correlation matrix for "fixed". Matrix should be symmetric with dimensions

>= the maximum cluster size. If the correlation structure is "userdefined", then

this is a matrix describing which correlations are the same.

init.beta an optional vector with the initial values of beta. If not specified, then the in-

tercept will be set to InvLink(mean(response)). init.beta must be specified if not

using an intercept.

init.alpha an optional scalar or vector giving the initial values for the correlation. If pro-

vided along with Mv>1 or unstructured correlation, then the user must ensure

that the vector is of the appropriate length.

init.phi an optional initial overdispersion parameter. If not supplied, initialized to 1.

scale.fix if set to TRUE, then the scale parameter is fixed at the value of init.phi.

sandwich if set to TRUE, the sandwich variance is provided together with the naive esti-

mator of variance.

maxit maximum number of iterations.

tol tolerance in calculation of coefficients.

print.log if set to TRUE, a report is printed.

typeweights a character string specifying the weights implementation. The following are

permitted: "GENMOD" for  $W^{1/2}V^{-1}W^{1/2}$ , "WV" for  $V^{-1}W$ 

nameTRT Name of the variable containing information for the treatment

an object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted for the propensity score. Must model the probability of being observed.

model.augmentation.trt

an object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted for the ouctome model for the treated group (A=1).

model.augmentation.ctrl

an object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted for the ouctome model for the control group (A=0).

stepwise.augmentation

if set to TRUE, a stepwise for the augmentation model is performed during the fit of the augmentation model for the OM

stepwise.weights

if set to TRUE, a stepwise for the propensity score is performed during the fit of the augmentation model for the OM

nameMISS Name of the variable containing information for the Missing indicator

nameY Name of the variable containing information for the outcome

sandwich.nuisance

if set to TRUE, the nuisance adjusted sandwich variance is provided.

fay.adjustment if set to TRUE, the small-sample nuisance adjusted sandwich variance with Fay's adjustement is provided.

fay. bound if set to 0.75 by default, bound value used for Fay's adjustement.

#### **Details**

The estimator is founds by solving:

$$0 = \sum_{i=1}^{M} \left[ \boldsymbol{D}_{i}^{T} \boldsymbol{V}_{i}^{-1} \boldsymbol{W}_{i}(\boldsymbol{X}_{i}, A_{i}, \boldsymbol{\eta}_{W}) \left( \boldsymbol{Y}_{i} - \boldsymbol{B}(\boldsymbol{X}_{i}, A_{i}, \boldsymbol{\eta}_{B}) \right) \right.$$

$$+ \sum_{a=0,1} p^{a} (1-p)^{1-a} \boldsymbol{D}_{i}^{T} \boldsymbol{V}_{i}^{-1} \Big( \boldsymbol{B}(\boldsymbol{X}_{i}, A_{i} = a, \boldsymbol{\eta}_{B}) - \boldsymbol{\mu}_{i}(\boldsymbol{\beta}, A_{i} = a) \Big) \Bigg]$$

where  $D_i = \frac{\partial \mu_i(\beta,A_i)}{\partial \beta^T}$  is the design matrix,  $V_i$  is the covariance matrix equal to  $U_i^{1/2}C(\alpha)U_i^{1/2}$  with  $U_i$  a diagonal matrix with elements  $\text{var}(y_{ij})$  and  $C(\alpha)$  is the working correlation structure with non-diagonal terms  $\alpha$ . Parameters  $\alpha$  are estimated using simple moment estimators from the Pearson residuals. The matrix of weights  $W_i(X_i,A_i,\eta_W) = diag\left[R_{ij}/\pi_{ij}(X_i,A_i,\eta_W)\right]_{j=1,\dots,n_i}$ , where  $\pi_{ij}(X_i,A_i,\eta_W) = P(R_{ij}|X_i,A_i)$  is the Propensity score (PS). The function  $B(X_i,A_i=a,\eta_B)$ , which is called the Outcome Model (OM), is a function linking  $Y_{ij}$  with  $X_i$  and  $A_i$ . The  $\eta_B$  are nuisance parameters that are estimated. The estimator is most efficient if the OM is equal to  $E(Y_i|X_i,A_i=a)$  The estimator denoted  $\hat{\beta}_{aug}$  is found by solving the estimating equation. Although analytic solutions sometimes exist, coefficient estimates are generally obtained using an

iterative procedure such as the Newton-Raphson method. Automatic implementation is such that,  $\hat{\eta}_W$  in  $W_i(X_i, A_i, \hat{\eta}_W)$  are obtained using a logistic regression and  $\hat{\eta}_B$  in  $B(X_i, A_i, \hat{\eta}_B)$  are obtained using a linear regression.

The variance of  $\hat{\beta}_{aug}$  is estimated by the sandwich variance estimator. There are two external sources of variability that need to be accounted for: estimation of  $\eta_W$  for the PS and of  $\eta_B$  for the OM. We denote  $\Omega = (\beta, \eta_W, \eta_B)$  the estimated parameters of interest and nuisance parameters. We can stack estimating functions and score functions for  $\Omega$ :

$$egin{aligned} oldsymbol{U}_i(oldsymbol{\Omega}) = \left(egin{array}{c} oldsymbol{\Phi}_i(oldsymbol{Y}_i, oldsymbol{X}_i, A_i, oldsymbol{eta}, oldsymbol{\eta}_W, oldsymbol{\eta}_B) \ oldsymbol{S}_i^B(oldsymbol{X}_i, A_i, oldsymbol{\eta}_B) \end{array}
ight) \end{aligned}$$

where  $S_i^W$  and  $S_i^B$  represent the score equations for patients in cluster i for the estimation of  $\eta_W$  and  $\eta_B$  in the PS and the OM. A standard Taylor expansion paired with Slutzky's theorem and the central limit theorem give the sandwich estimator adjusted for nuisance parameters estimation in the OM and PS:

$$Var(\mathbf{\Omega}) = E\left[\frac{\partial \boldsymbol{U}_i(\mathbf{\Omega})}{\partial \mathbf{\Omega}}\right]^{-1} \underbrace{E\left[\boldsymbol{U}_i(\mathbf{\Omega})\boldsymbol{U}_i^T(\mathbf{\Omega})\right]}_{\boldsymbol{\Delta}_{adj}} \underbrace{E\left[\frac{\partial \boldsymbol{U}_i(\mathbf{\Omega})}{\partial \mathbf{\Omega}}\right]^{-1}}_{\boldsymbol{\Gamma}_{adi}^{-1}}.$$

#### Value

An object of type 'CRTgeeDR'

\$beta Final values for regressors estimates

- \$phi scale parameter estimate
- \$alpha Final values for association parameters in the working correlation structure when exchangeable
- \$coefnames Name of the regressors in the main regression
- \$niter Number of iteration done by the algorithm before convergence
- · \$converged convergence status
- \$var.naiv Variance of the estimates model based (naive)
- · \$var Variance of the estimates sandwich
- \$var.nuisance Variance of the estimates nuisance adjusted sandwich
- \$var.fay Variance of the estimates nuisance adjusted sandwich with Fay correction for small samples
- \$call Call function

- \$corr Correlation structure used
- \$clusz Number of unit in each cluster
- \$FunList List of function associated with the family
- \$X design matrix for the main regression
- \$offset Offset specified in the regression
- \$eta predicted values
- \$weights Weights vector used in the diagonal term for the IPW
- \$ps.model Summary of the regression fitted for the PS if computed internally
- \$om.model.trt Summary of the regression fitted for the OM for treated if computed internally
- \$om.model.ctrl Summary of the regression fitted for the OM for control if computed internally

#### Author(s)

Melanie Prague [based on R packages 'geeM' L. S. McDaniel, N. C. Henderson, and P. J. Rathouz. Fast Pure R Implementation of GEE: Application of the Matrix Package. The R Journal, 5(1):181-188, June 2013.]

#### References

Details regarding implementation can be found in

- 'Augmented GEE for improving efficiency and validity of estimation in cluster randomized trials by leveraging cluster-and individual-level covariates' - 2012 - Stephens A., Tchetgen Tchetgen E. and De Gruttola V.: Stat Med 31(10) - 915-930.
- 'Accounting for interactions and complex inter-subject dependency for estimating treatment effect in cluster randomized trials with missing at random outcomes' 2015 Prague M., Wang R., Stephens A., Tchetgen Tchetgen E. and De Gruttola V.: in revision.
- 'Fast Pure R Implementation of GEE: Application of the Matrix Package' 2013 McDaniel, Lee S and Henderson, Nicholas C and Rathouz, Paul J: The R Journal 5(1) 181-197.
- 'Small-Sample Adjustments for Wald-Type Tests Using Sandwich Estimators' 2001 Fay, Michael P and Graubard, Barry I: Biometrics 57(4) 1198-1206.

#### **Examples**

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```
#### AUGMENTED GEE
augresults<-geeDREstimation(formula=OUTCOME~TRT,</pre>
                                id="CLUSTER" , data = data.sim,
                                family = "binomial", corstr = "independence",
                                model.augmentation.trt=OUTCOME~AGE,
                       model.augmentation.ctrl=OUTCOME~AGE, stepwise.augmentation=FALSE)
summary(augresults)
## End(Not run)
#### DOUBLY ROBUST
drresults<-geeDREstimation(formula=OUTCOME~TRT,</pre>
                                id="CLUSTER" , data = data.sim,
                                family = "binomial", corstr = "independence",
                                model.weights=I(MISSING==0)~TRT*AGE,
                                model.augmentation.trt=OUTCOME~AGE,
                       model.augmentation.ctrl=OUTCOME~AGE, stepwise.augmentation=FALSE)
summary(drresults)
```

getCI

Get Mean, Sd and CI for estimates from CRTgeeDR object.

#### **Description**

Get the estimates, standard deviations and confidence intervals from an CRTgeeDR object associated with a regressor given in argument.

#### Usage

```
getCI(object, nameTRT = "TRT", quantile = 1.96)
```

#### Arguments

object	CRTgeeDR
nameTRT	character including the name of the variable of interest (often the treatment)
quantile	value of the normal quantile for the IC. default is 1.96 for 95%CI.

getOMPlot

Get the observed vs fitted residuals

#### **Description**

Get the histogram and some basic statistics for the weights used in the IPW part.

#### Usage

```
getOMPlot(object, save = FALSE, name = "plotOM", typeplot = 0)
```

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#### **Arguments**

object CRTgeeDR

save logical if TRUE the plot is saved as a pdf in the current directory

name name of the plot saved as pdf

typeplot integer indicating which is the adequation diagnostic plot for the PS. '0', all

available in plot.glm are displayed, '1' Residuals vs Fitted, '2' Normal Q-Q, '3' Scale-Location, '4' Cook's distance, '5' Residuals vs Leverage and '6' Cook's

dist vs Leverage\* h[ii] / (1 - h[ii])

getPSPlot Get the histogram of weights for IPW and adequation for the glm

weights model

#### **Description**

Get the histogram and some basic statistics for the weights used in the IPW part.

# Usage

```
getPSPlot(object, save = FALSE, name = "plotPS", typeplot = NULL)
```

#### **Arguments**

object CRTgeeDR

save logical if TRUE the plot is saved as a pdf in the current directory

name name of the plot saved as pdf

typeplot integer indicating which is the adequation diagnostic plot for the PS. Default is

NULL no output. '0', all available in plot.glm are displayed, '1' Residuals vs Fitted, '2' Normal Q-Q, '3' Scale-Location, '4' Cook's distance, '5' Residuals

vs Leverage and '6' Cook's dist vs Leverage\* h[ii] / (1 - h[ii])

predict.CRTgeeDR Predict CRTgeeDR object.

#### **Description**

Predict CRTgeeDR object to a dataset

#### **Usage**

```
## S3 method for class 'CRTgeeDR'
predict(object, newdata = NULL, ...)
```

#### **Arguments**

object CRTgeeDR object

newdata dataframe, new dataset to which the CRTgeeDRneed to be used for prediction

... ignored

print.CRTgeeDR

Prints CRTgeeDR object.

# Description

Prints CRTgeeDR object

### Usage

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

# Arguments

x CRTgeeDR x ... ignored

print.summary.CRTgeeDR

Print the summarizing CRTgeeDR object.

# Description

Print Summary CRTgeeDR object

#### Usage

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

#### Arguments

x summary.CRTgeeDR x

... ignored

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summary.CRTgeeDR

 $Summarizing\ CRT geeDR\ object.$ 

# Description

Summary CRTgeeDR object

# Usage

```
## S3 method for class 'CRTgeeDR'
summary(object, ...)
```

# Arguments

object CRTgeeDR object

... ignored

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