Package 'EHRmuse'

January 20, 2025

Type Package		
Title Multi-Cohort Selection Bias Correction using IPW and AIPW Methods		
Version 0.0.2.0		
Description Comprehensive toolkit for addressing selection bias in binary disease models across diverse non-probability samples, each with unique selection mechanisms. It utilizes Inverse Probability Weighting (IPW) and Augmented Inverse Probability Weighting (AIPW) methods to reduce selection bias effectively in multiple non-probability cohorts by integrating data from either individual-level or summary-level external sources. The package also provides a variety of variance estimation techniques. Please refer to Kundu et al. <doi:10.48550 arxiv.2412.00228="">.</doi:10.48550>		
License GPL (>= 2)		
<pre>URL https://github.com/Ritoban1/EHRmuse</pre>		
BugReports https://github.com/Ritoban1/EHRmuse/issues		
Depends R (>= $4.0.0$)		
Imports dplyr (>= 1.0.0), magrittr, MASS, nleqslv (>= 3.3.2), xgboost (>= 1.4.1), survey (>= 4.1.0), stats, nnet (>= 7.3-17), simplexreg (>= 0.1.6)		
Encoding UTF-8		
RoxygenNote 7.3.2		
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Repository CRAN		
Date/Publication 2025-01-20 16:20:13 UTC		
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EHRmuse

IPW and AIPW Methods for Multi-cohort Selection Bias in Non-probability Samples

Description

IPW and AIPW Methods for Multi-cohort Selection Bias in Non-probability Samples

Usage

```
EHRmuse(
 Κ,
  Z_names,
  intdata_list,
 N = NULL,
 UW_CS = FALSE,
  IPW = FALSE,
 weights_user = NULL,
 AIPW = FALSE,
  ipw_method = "PL",
  extdata = NULL,
 marginals_list = NULL,
  select_var_list = NULL,
  aux_var_list = NULL,
 Weights_e = NULL,
  aux_model = "XGBoost",
 variance = FALSE,
  type_var = "approx"
)
```

Arguments

K	Necessary Input. Number of cohorts. Should be a numeric positive integer.
Z_names	Necessary Input. A character vector containing the names of the Z variables or disease model covariates.
intdata_list	Necessary Input. A list of size K where each element of list corresponds to the data for each of the K multiple cohorts including the disease indicator D, the Z variables and the selection variables in the disease model. Each data should be of the form of a data frame. Please include a column named "id" to indicate unique identifiers to the units.
N	Target Population Size.
UW_CS	An indicator variable (TRUE or FALSE) for using the unweighted logistic regression model with cohort-specific intercepts.
IPW	An indicator variable (TRUE or FALSE) for using the Inverse Probability Weighted Methods (IPW) methods.

weights_user User specified weights. A numeric vector of weights for the combined data (not

duplicated).

AIPW An indicator variable (TRUE or FALSE) for using the Joint Augmented IPW

method. If AIPW is TRUE, please also input IPW to be TRUE.

ipw_method If IPW is TRUE, specify the IPW method to be used. A character variable.

Default is PL (Pseudolikelihood). Other options are Simplex Regression (SR),

Calibration (CL) or User Specified (US).

frame containing individual level external data which is a probability sample like NHANES. The external data should contain all the selection variables and if AIPW is TRUE, then also all the auxiliary score model variables. Please

include a column named "id" to indicate unique identifiers to the units.

marginals_list If IPW method is set to CL, please provide a list of size K in which each element

is a numeric vector containing the marginal sums of the selection variables of each of the K cohorts. Please ensure the first element for each of the K numeric

vector should be the population size, N.

select_var_list

If IPW is set to be TRUE, please provide a list of size K in which each element is a character vector corresponding to the selection variables' names for each of

the K cohorts.

aux_var_list If AIPW is set to be TRUE, please provide a list of size K in which each element

is a character vector corresponding to the auxiliary score model variables' names

for each of the K cohorts.

Weights_e If IPW method is set to PL or SR or AIPW is TRUE, please provide the known

selection weights for the external probability sample. The input should be a

numeric vector.

aux_model If AIPW is true, please provide the auxiliary score model. Default is XGboost.

variance An indicator variable (TRUE or FALSE) whether variance should be computed

or not, along with the point estimate.

type_var If variance is true, indicate the method type to be used for computing the vari-

ance. For the unweighted method, do not provide any type. For IPW methods, PL and CL, we have two options, asy (asymptotic variance incorporating the variance from nuisance parameters) and "approx" ignoring the variance from nuisance parameters. For SR and AIPW, we have only the approx method. The

default for IPW and AIPW methods is approx.

Value

If variance=TRUE, it will return a list of estimate vector and variance vector. If variance=FALSE, it will return an estimate vector.

Examples

#library(MASS)

K=3 ## Number of Cohorts

```
set.seed(100)
mean_w_p=0
mean_z_1=0
mean_z_2=0
mean_z_3=0
corr=0.5
var_z_w_p=matrix(c(1,corr,corr,corr,
                   corr,1,corr,corr,
                    corr,corr,1,corr,
                    corr,corr,corr,1),
                 nrow=4,ncol=4)
theta=c(-2,0.35,0.45,0.25) ## Theta_Z vector
N=5e4 ## Population size
### selection models
dw=1
dwz1=c(1,0.8,0.6)
dwz2=c(0.6,0.8,1)
dwz3=rep(1,3)
gamma_ext=c(-0.6,1.2,0.4,-0.2,0.5)
gamma_int_1=c(-1,1.5,0.2,0.8,-0.3)
gamma_int_2=c(-1,1.25,0.4,0.6)
gamma_int_3=c(-3,0.8,0.5)
## Generation of population level data
simu_popu<-function(N,mean_w_p,mean_z_1,mean_z_2,mean_z_3,</pre>
                    var_z_w_p,theta,dw){
  \verb|cov<- MASS::mvrnorm| (n = N, mu = c(mean\_w\_p, mean\_z\_1, mean\_z\_2, mean\_z\_3), \\ Sigma = var\_z\_w\_p) \\
    data <- data.frame(Z1 = cov[, 2], Z2 = cov[, 3], Z3=cov[,4])
    W_p=cov[,1]
    # Generate random uniforms
    #set.seed(5678)
    U1 <- runif(N)
    #set.seed(4321)
    # Generate Disease Status
    DISEASE <- expit(theta[1] + theta[2] * data$Z1 + theta[3]*data$Z2 +theta[4]*data$Z3)
    data$D <- ifelse(DISEASE > U1, 1, 0)
    # Relate W_p and D
    data$W_1 \leftarrow W_p + dw* data$D + dwz1[1]*data$Z1 +
        dwz2[1]*data$Z2 + dwz3[1]*data$Z3 +\\
        rnorm(n=N,0,1)
    data$W_2 <- W_p + dw* data$D + dwz1[2]*data$Z1 +
        dwz2[2]*data$Z2 + dwz3[2]*data$Z3 +
        rnorm(n=N,0,1)
    data$W_3 <- W_p + dw* data$D + dwz1[3]*data$Z1 +
        dwz2[3]*data$Z2 + dwz3[3]*data$Z3 +\\
        rnorm(n=N,0,1)
    data$id=c(1:N)
```

```
return(data)
}
## Generation of external individual level data
simu_ext<-function(data,gamma_ext){</pre>
   U2e <- runif(N)
    # Generate Sampling Status
    SELECT <-0.75*expit(gamma_ext[1] +</pre>
                             gamma_ext[2]* data$D +
                             gamma_ext[3] * data$Z1 +
                             gamma_ext[4]* data$Z2 +
                             gamma_ext[5] * data$Z3)
    S_e <- ifelse(SELECT > U2e, TRUE, FALSE)
    # Observed Data
    data_e <- data[which(S_e==1),]</pre>
    data_e$Select_Weights = 0.75*expit(gamma_ext[1] +
                                             {\tt gamma\_ext[2]* data\_e\$D +}
                                             gamma_ext[3] * data_e$Z1 +
                                             gamma_ext[4]* data_e$Z2 +
                                             gamma_ext[5] * data_e$Z3)
    return(data_e)
}
## Generation of internal data 1
simu_int_1<-function(data,gamma_int_1){</pre>
   U2i <- runif(N)
    # Generate Sampling Status
    SELECT <- expit(cbind(1,data$D,data$W_1,data$Z2,data$Z3)</pre>
                     %*% gamma_int_1)
    S_i <- ifelse(SELECT > U2i, TRUE, FALSE)
    # Observed Data
    data_i <- data[which(S_i==1),]</pre>
    return(data_i)
}
## Generation of internal data 2
simu_int_2<-function(data,gamma_int_2){</pre>
   U2i <- runif(N)
    # Generate Sampling Status
    SELECT <- expit(cbind(1,data$D,data$W_2,data$Z3)</pre>
                     %*% gamma_int_2)
    S_i <- ifelse(SELECT > U2i, TRUE, FALSE)
    # Observed Data
    data_i \leftarrow data[which(S_i==1),]
    return(data_i)
}
## Generation of internal data 3
simu_int_3<-function(data,gamma_int_3){</pre>
   U2i <- runif(N)
    # Generate Sampling Status
    SELECT <- expit(cbind(1,data$W_3,data$Z2)</pre>
                     %*% gamma_int_3)
    S_i <- ifelse(SELECT > U2i, TRUE, FALSE)
```

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```
# Observed Data
   data_i <- data[which(S_i==1),]</pre>
   return(data_i)
}
data=simu_popu(N,mean_w_p,mean_z_1,mean_z_2,mean_z_3,
               var_z_w_p,theta,dw)
extdata=simu_ext(data,gamma_ext)
intdata1=simu_int_1(data,gamma_int_1)
intdata2=simu_int_2(data,gamma_int_2)
intdata3=simu_int_3(data,gamma_int_3)
## names of selection variables in each cohort
select\_var\_list=list(c("D","W\_1","Z2","Z3"),c("D","W\_2","Z3"),c("W\_3","Z2"))
## names of auxiliary variables in each cohort
aux_var_list=list(c("D","W_1","Z2","Z3"),c("D","W_2","Z3"),c("W_3","Z2"))
## list of internal data
intdata_list=list(intdata1,intdata2,intdata3)
## names of Z variables
Z_names=c("Z1","Z2","Z3")
theta ## actual theta_z
res_uw=EHRmuse(K=K,N=N,Z_names=Z_names,
               intdata_list=intdata_list,variance = TRUE)
```

expit

Expit

Description

Expit

Usage

expit(x)

Arguments

Χ

numeric vector

Value

```
\exp(x)/(1+\exp(x))
```

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Examples

expit(1)

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