

# Package: geeCRT (via r-universe)

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**Type** Package

**Title** Bias-Corrected GEE for Cluster Randomized Trials

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**Description** Population-averaged models have been increasingly used in the design and analysis of cluster randomized trials (CRTs). To facilitate the applications of population-averaged models in CRTs, the package implements the generalized estimating equations (GEE) and matrix-adjusted estimating equations (MAEE) approaches to jointly estimate the marginal mean models correlation models both for general CRTs and stepped wedge CRTs. Despite the general GEE/MAEE approach, the package also implements a fast cluster-period GEE method by Li et al. (2022) <doi:10.1093/biostatistics/kxaa056> specifically for stepped wedge CRTs with large and variable cluster-period sizes and gives a simple and efficient estimating equations approach based on the cluster-period means to estimate the intervention effects as well as correlation parameters. In addition, the package also provides functions for generating correlated binary data with specific mean vector and correlation matrix based on the multivariate probit method in Emrich and Piedmonte (1991) <doi:10.1080/00031305.1991.10475828> or the conditional linear family method in Qaqish (2003) <doi:10.1093/biomet/90.2.455>.

**License** GPL (>= 2)

**LazyData** TRUE

**Depends** R (>= 3.6.0)

**Imports** MASS, rootSolve, mvtnorm

**Suggests** knitr, rmarkdown

**VignetteBuilder** knitr

**RoxygenNote** 7.2.3

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Elizabeth L. Turner [aut], John Preisser [aut]**Date/Publication** 2021-10-10 04:10:02 UTC**Repository** <https://hengshiyu.r-universe.dev>**RemoteUrl** <https://github.com/hengshiyu/geecrt>**RemoteRef** HEAD**RemoteSha** 882b13f581313748cd3522c9602e55891d2ee864

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cpgeeSWD	<i>Cluster-Period GEE for Estimating the Mean and Correlation Parameters in Cross-Sectional SW-CRTs</i>
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## Description

cpgeeSWD implements the cluster-period GEE developed for cross-sectional stepped wedge cluster randomized trials (SW-CRTs). It provides valid estimation and inference for the treatment effect and intraclass correlation parameters within the GEE framework, and is computationally efficient for SW-CRTs with large cluster sizes. The program currently only allows for a marginal mean model with discrete period effects and the intervention indicator without additional covariates. The program offers bias-corrected ICC estimates as well as bias-corrected sandwich variances for both the treatment effect parameter and the ICC parameters. The technical details of the cluster-period GEE approach are provided in Li et al. (2020+).

## Usage

```
cpgeeSWD(
  y,
  X,
  id,
  m,
```

```

  corstr,
  family = "binomial",
  maxiter = 500,
  epsilon = 0.001,
  printrange = TRUE,
  alpadj = FALSE,
  rho.init = NULL
)

```

### Arguments

<code>y</code>	a vector specifying the cluster-period means (proportions)
<code>X</code>	design matrix for the marginal mean model, including period indicator and intervention indicator
<code>id</code>	a vector specifying cluster identifier
<code>m</code>	a vector of the cluster-period sizes
<code>corstr</code>	correlation structure specified for the individual-level outcomes, could be 'exchangeable', 'nest_exch' or 'exp_decay'
<code>family</code>	See corresponding documentation to <code>glm</code> . The current version only supports <code>family = 'binomial'</code>
<code>maxiter</code>	maximum number of iterations for Fisher scoring updates
<code>epsilon</code>	tolerance for convergence
<code>printrange</code>	print details of range violations when <code>family = 'binomial'</code> . The default is <code>TRUE</code>
<code>alpadj</code>	if <code>TRUE</code> , performs bias adjustment for the alpha estimating equations. The default is <code>FALSE</code>
<code>rho.init</code>	user-specified initial value for the decay parameter when <code>corstr = 'exp_decay'</code>

### Value

`outbeta` estimates of marginal mean model parameters and standard errors with different finite-sample bias corrections. The current version supports model-based standard error (MB), the sandwich standard error (BC0) extending Zhao and Prentice (2001), the sandwich standard errors (BC1) extending Kauermann and Carroll (2001), the sandwich standard errors (BC2) extending Mancl and DeRouen (2001), and the sandwich standard errors (BC3) extending the Fay and Graubard (2001). A summary of these bias-corrections can also be found in Lu et al. (2007), and Li et al. (2018).

`outalpha` estimates of correlation parameters and standard errors with different finite-sample bias corrections. The current version supports the sandwich standard error (BC0) extending Zhao and Prentice (2001), the sandwich standard errors (BC1) extending Kauermann and Carroll (2001), the sandwich standard errors (BC2) extending Mancl and DeRouen (2001), and the sandwich standard errors (BC3) extending the Fay and Graubard (2001). A summary of these bias-corrections can also be found in Preisser et al. (2008).

`beta` a vector of estimates for marginal mean model parameters

`alpha` a vector of estimates of correlation parameters

`MB` model-based covariance estimate for the marginal mean model parameters

BC0 robust sandwich covariance estimate of the marginal mean model and correlation parameters  
 BC1 robust sandwich covariance estimate of the marginal mean model and correlation parameters with the Kauermann and Carroll (2001) correction  
 BC2 robust sandwich covariance estimate of the marginal mean model and correlation parameters with the Mancl and DeRouen (2001) correction  
 BC3 robust sandwich covariance estimate of the marginal mean model and correlation parameters with the Fay and Graubard (2001) correction  
 ni ter number of iterations used in the Fisher scoring updates for model fitting

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### References

Zhao, L. P., Prentice, R. L. (1990). Correlated binary regression using a quadratic exponential model. *Biometrika*, 77(3), 642-648.

Mancl, L. A., DeRouen, T. A. (2001). A covariance estimator for GEE with improved small sample properties. *Biometrics*, 57(1), 126-134.

Kauermann, G., Carroll, R. J. (2001). A note on the efficiency of sandwich covariance matrix estimation. *Journal of the American Statistical Association*, 96(456), 1387-1396.

Fay, M. P., Graubard, B. I. (2001). Small sample adjustments for Wald type tests using sandwich estimators. *Biometrics*, 57(4), 1198-1206.

Lu, B., Preisser, J. S., Qaqish, B. F., Suchindran, C., Bangdiwala, S. I., Wolfson, M. (2007). A comparison of two bias corrected covariance estimators for generalized estimating equations. *Biometrics*, 63(3), 935-941.

Preisser, J. S., Lu, B., Qaqish, B. F. (2008). Finite sample adjustments in estimating equations and covariance estimators for intracluster correlations. *Statistics in Medicine*, 27(27), 5764-5785.

Li, F., Turner, E. L., Preisser, J. S. (2018). Sample size determination for GEE analyses of stepped wedge cluster randomized trials. *Biometrics*, 74(4), 1450-1458.

Li, F. (2020). Design and analysis considerations for cohort stepped wedge cluster randomized trials with a decay correlation structure. *Statistics in Medicine*, 39(4), 438-455.

Li, F., Yu, H., Rathouz, P., Turner, E. L., Preisser, J. S. (2021). Marginal modeling of cluster-period means and intraclass correlations in stepped wedge designs with binary outcomes. *Biostatistics*, kxaa056.

### Examples

```
# Simulated SW-CRT example with binary outcome

#####
### Example 1): simulated SW-CRT with smaller cluster-period sizes (5~10)
#####

sampleSWCRT <- sampleSWCRTsmall
```

```
#####
### cluster-period id, period, outcome, and design matrix ###
#####

### id, period, outcome
id <- sampleSWCRT$id
period <- sampleSWCRT$period
y <- sampleSWCRT$y_bin
X <- as.matrix(sampleSWCRT[, c("period1", "period2", "period3", "period4", "treatment")])

m <- as.matrix(table(id, period))
n <- dim(m)[1]
t <- dim(m)[2]
clp_mu <- tapply(y, list(id, period), FUN = mean)
y_cp <- c(t(clp_mu))

### design matrix for correlation parameters
trt <- tapply(X[, t + 1], list(id, period), FUN = mean)
trt <- c(t(trt))

time <- tapply(period, list(id, period), FUN = mean)
time <- c(t(time))
X_cp <- matrix(0, n * t, t)

s <- 1
for (i in 1:n) {
  for (j in 1:t) {
    X_cp[s, time[s]] <- 1
    s <- s + 1
  }
}
X_cp <- cbind(X_cp, trt)
id_cp <- rep(1:n, each = t)
m_cp <- c(t(m))

#####
### cluster-period matrix-adjusted estimating equations (MAEE)
### with exchangeable, nested exchangeable and exponential decay correlation structures ###
#####

# exchangeable
est_mae_exc <- cpgeeSWD(
  y = y_cp, X = X_cp, id = id_cp,
  m = m_cp, corstr = "exchangeable",
  alpadj = TRUE
)
print(est_mae_exc)

# nested exchangeable
est_mae_nex <- cpgeeSWD(
  y = y_cp, X = X_cp, id = id_cp,
  m = m_cp, corstr = "nest_exch",
```

```

    alpadj = TRUE
  )
  print(est_mae_nex)

  # exponential decay
  est_mae_ed <- cpgeeSWD(
    y = y_cp, X = X_cp, id = id_cp,
    m = m_cp, corstr = "exp_decay",
    alpadj = TRUE
  )
  print(est_mae_ed)

#####
### cluster-period GEE
### with exchangeable, nested exchangeable and exponential decay correlation structures ###
#####

# exchangeable
est_uee_exc <- cpgeeSWD(
  y = y_cp, X = X_cp, id = id_cp,
  m = m_cp, corstr = "exchangeable",
  alpadj = FALSE
)
print(est_uee_exc)

# nested exchangeable
est_uee_nex <- cpgeeSWD(
  y = y_cp, X = X_cp, id = id_cp,
  m = m_cp, corstr = "nest_exch",
  alpadj = FALSE
)
print(est_uee_nex)

# exponential decay
est_uee_ed <- cpgeeSWD(
  y = y_cp, X = X_cp, id = id_cp,
  m = m_cp, corstr = "exp_decay",
  alpadj = FALSE
)
print(est_uee_ed)

#####
### Example 2): simulated SW-CRT with larger cluster-period sizes (20~30)
#####

sampleSWCRT <- sampleSWCRTLarge

#####
### cluster-period id, period, outcome, and design matrix ###
#####

### id, period, outcome
id <- sampleSWCRT$id

```

```

period <- sampleSWCRT$period
y <- sampleSWCRT$y_bin
X <- as.matrix(sampleSWCRT[, c("period1", "period2", "period3", "period4", "period5", "treatment")])

m <- as.matrix(table(id, period))
n <- dim(m)[1]
t <- dim(m)[2]
clp_mu <- tapply(y, list(id, period), FUN = mean)
y_cp <- c(t(clp_mu))

### design matrix for correlation parameters
trt <- tapply(X[, t + 1], list(id, period), FUN = mean)
trt <- c(t(trt))

time <- tapply(period, list(id, period), FUN = mean)
time <- c(t(time))
X_cp <- matrix(0, n * t, t)

s <- 1
for (i in 1:n) {
  for (j in 1:t) {
    X_cp[s, time[s]] <- 1
    s <- s + 1
  }
}
X_cp <- cbind(X_cp, trt)
id_cp <- rep(1:n, each = t)
m_cp <- c(t(m))

#####
### cluster-period matrix-adjusted estimating equations (MAEE)
### with exchangeable, nested exchangeable and exponential decay correlation structures ###
#####

# exchangeable
est_mae_exc <- cpgeeSWD(
  y = y_cp, X = X_cp, id = id_cp,
  m = m_cp, corstr = "exchangeable",
  alpadj = TRUE
)
print(est_mae_exc)

# nested exchangeable
est_mae_nex <- cpgeeSWD(
  y = y_cp, X = X_cp, id = id_cp,
  m = m_cp, corstr = "nest_exch",
  alpadj = TRUE
)
print(est_mae_nex)

# exponential decay
est_mae_ed <- cpgeeSWD(
  y = y_cp, X = X_cp, id = id_cp,

```

```

    m = m_cp, corstr = "exp_decay",
    alpadj = TRUE
  )
print(est_mae_ed)

#####
### cluster-period GEE
### with exchangeable, nested exchangeable and exponential decay correlation structures ###
#####

# exchangeable
est_uee_exc <- cpgeeSWD(
  y = y_cp, X = X_cp, id = id_cp,
  m = m_cp, corstr = "exchangeable",
  alpadj = FALSE
)
print(est_uee_exc)

# nested exchangeable
est_uee_nex <- cpgeeSWD(
  y = y_cp, X = X_cp, id = id_cp,
  m = m_cp, corstr = "nest_exch",
  alpadj = FALSE
)
print(est_uee_nex)

# exponential decay
est_uee_ed <- cpgeeSWD(
  y = y_cp, X = X_cp, id = id_cp,
  m = m_cp, corstr = "exp_decay",
  alpadj = FALSE
)
print(est_uee_ed)

```

---

geemaee

---

*GEE and Matrix-adjusted Estimating Equations (MAEE) for Estimating the Marginal Mean and Correlation Parameters in CRTs*


---

## Description

geemaee implements the GEE and matrix-adjusted estimating equations (MAEE) for analyzing cluster randomized trials (CRTs). It supports estimation and inference for the marginal mean and intraclass correlation parameters within the population-averaged modeling framework. With suitable choice of the design matrices, the function can be used to analyze parallel, crossover and stepped wedge cluster randomized trials. The program also offers bias-corrected intraclass correlation estimates, as well as bias-corrected sandwich variances for both the marginal mean and correlation parameters. The technical details of the GEE and MAEE approach are provided in Preisser (2008) and Li et al. (2018, 2019).



**Usage**

```
geemae(
  y,
  X,
  id,
  Z,
  family,
  link = NULL,
  maxiter = 500,
  epsilon = 0.001,
  printrange = TRUE,
  alpadj = FALSE,
  shrink = "ALPHA",
  makevone = TRUE
)
```

**Arguments**

y	a vector specifying the outcome variable across all clusters
X	design matrix for the marginal mean model, including the intercept
id	a vector specifying cluster identifier
Z	design matrix for the correlation model, should be all pairs $j < k$ for each cluster
family	See corresponding documentation to <code>glm</code> . The current version supports 'continuous', 'binomial', 'poisson' and 'quasipoisson'
link	a specification for the model link function name
maxiter	maximum number of iterations for Fisher scoring updates
epsilon	tolerance for convergence. The default is 0.001
printrange	print details of range violations. The default is TRUE
alpadj	if TRUE, performs bias adjustment for the correlation estimating equations. The default is FALSE
shrink	method to tune step sizes in case of non-convergence including 'THETA' or 'ALPHA'. The default is 'ALPHA'
makevone	if TRUE, it assumes unit variances for the correlation parameters in the correlation estimating equations. The default is TRUE

**Value**

outbeta estimates of marginal mean model parameters and standard errors with different finite-sample bias corrections. The current version supports model-based standard error (MB), the sandwich standard error (BC0) extending Liang and Zeger (1986), the sandwich standard errors (BC1) extending Kauermann and Carroll (2001), the sandwich standard errors (BC2) extending Mancl and DeRouen (2001), and the sandwich standard errors (BC3) extending the Fay and Graubard (2001). A summary of these bias-corrections can also be found in Lu et al. (2007), and Li et al. (2018).

outalpha estimates of intraclass correlation parameters and standard errors with different finite-sample bias corrections. The current version supports the sandwich standard error (BC0) extending

Zhao and Prentice (2001), the sandwich standard errors (BC1) extending Kauermann and Carroll (2001), the sandwich standard errors (BC2) extending Mancl and DeRouen (2001), and the sandwich standard errors (BC3) extending the Fay and Graubard (2001). A summary of these bias-corrections can also be found in Preisser et al. (2008).

beta a vector of estimates for marginal mean model parameters

alpha a vector of estimates of correlation parameters

MB model-based covariance estimate for the marginal mean model parameters

BC0 robust sandwich covariance estimate of the marginal mean model and correlation parameters

BC1 robust sandwich covariance estimate of the marginal mean model and correlation parameters with the Kauermann and Carroll (2001) correction

BC2 robust sandwich covariance estimate of the marginal mean model and correlation parameters with the Mancl and DeRouen (2001) correction

BC3 robust sandwich covariance estimate of the marginal mean model and correlation parameters with the Fay and Graubard (2001) correction

niter number of iterations used in the Fisher scoring updates for model fitting

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- Fay, M. P., Graubard, B. I. (2001). Small sample adjustments for Wald type tests using sandwich estimators. *Biometrics*, 57(4), 1198-1206.
- Lu, B., Preisser, J. S., Qaqish, B. F., Suchindran, C., Bangdiwala, S. I., Wolfson, M. (2007). A comparison of two bias corrected covariance estimators for generalized estimating equations. *Biometrics*, 63(3), 935-941.

Preisser, J. S., Lu, B., Qaqish, B. F. (2008). Finite sample adjustments in estimating equations and covariance estimators for intracluster correlations. *Statistics in Medicine*, 27(27), 5764-5785.

Li, F., Turner, E. L., Preisser, J. S. (2018). Sample size determination for GEE analyses of stepped wedge cluster randomized trials. *Biometrics*, 74(4), 1450-1458.

Li, F., Forbes, A. B., Turner, E. L., Preisser, J. S. (2019). Power and sample size requirements for GEE analyses of cluster randomized crossover trials. *Statistics in Medicine*, 38(4), 636-649.

Li, F. (2020). Design and analysis considerations for cohort stepped wedge cluster randomized trials with a decay correlation structure. *Statistics in Medicine*, 39(4), 438-455.

Li, F., Yu, H., Rathouz, P. J., Turner, E. L., & Preisser, J. S. (2022). Marginal modeling of cluster-period means and intraclass correlations in stepped wedge designs with binary outcomes. *Biostatistics*, 23(3), 772-788.

## Examples

```
# Simulated SW-CRT examples

#####
### function to create the design matrix for correlation parameters
### under the nested exchangeable correlation structure
#####
createzCrossSec <- function(m) {
  Z <- NULL
  n <- dim(m)[1]
  for (i in 1:n) {
    alpha_0 <- 1
    alpha_1 <- 2
    n_i <- c(m[i, ])
    n_length <- length(n_i)
    POS <- matrix(alpha_1, sum(n_i), sum(n_i))
    loc1 <- 0
    loc2 <- 0
    for (s in 1:n_length) {
      n_t <- n_i[s]
      loc1 <- loc2 + 1
      loc2 <- loc1 + n_t - 1
      for (k in loc1:loc2) {
        for (j in loc1:loc2) {
          if (k != j) {
            POS[k, j] <- alpha_0
          } else {
            POS[k, j] <- 0
          }
        }
      }
    }
  }
  zrow <- diag(2)
  z_c <- NULL
  for (j in 1:(sum(n_i) - 1)) {
    for (k in (j + 1):sum(n_i)) {
      z_c <- rbind(z_c, zrow[POS[j, k], ])
    }
  }
}
```

```

    }
    Z <- rbind(Z, z_c)
  }
  return(Z)
}

#####
### Example 1): simulated SW-CRT with smaller cluster-period sizes (5~10)
#####

sampleSWCRT <- sampleSWCRTSmall

#####
### Individual-level id, period, outcome, and design matrix ###
#####

id <- sampleSWCRT$id
period <- sampleSWCRT$period
X <- as.matrix(sampleSWCRT[, c("period1", "period2", "period3", "period4", "treatment")])

m <- as.matrix(table(id, period))
n <- dim(m)[1]
t <- dim(m)[2]
### design matrix for correlation parameters
Z <- createzCrossSec(m)

#####
### (1) Matrix-adjusted estimating equations and GEE
### on continous outcome with nested exchangeable correlation structure
#####

est_mae_ind_con <- geemaee(
  y = sampleSWCRT$y_con, X = X, id = id,
  Z = Z, family = "continuous",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = TRUE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_mae_ind_con)

est_uee_ind_con <- geemaee(
  y = sampleSWCRT$y_con, X = X, id = id,
  Z = Z, family = "continuous",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = FALSE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_uee_ind_con)

#####
### (2) Matrix-adjusted estimating equations and GEE

```

```

### on binary outcome with nested exchangeable correlation structure
#####

est_mae_ind_bin <- geemae(
  y = sampleSWCRT$y_bin, X = X, id = id,
  Z = Z, family = "binomial",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = TRUE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_mae_ind_bin)

### GEE
est_uee_ind_bin <- geemae(
  y = sampleSWCRT$y_bin, X = X, id = id,
  Z = Z, family = "binomial",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = FALSE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_uee_ind_bin)

#####
### (3) Matrix-adjusted estimating equations and GEE
### on count outcome with nested exchangeable correlation structure
### using Poisson distribution
#####
### MAEE
est_mae_ind_cnt_poisson = geemae(
  y = sampleSWCRT$y_bin,
  X = X, id = id, Z = Z,
  family = "poisson",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = TRUE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_mae_ind_cnt_poisson)

### GEE
est_uee_ind_cnt_poisson = geemae(
  y = sampleSWCRT$y_bin,
  X = X, id = id, Z = Z,
  family = "poisson",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = FALSE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_uee_ind_cnt_poisson)

#####

```

```

### (4) Matrix-adjusted estimating equations and GEE
### on count outcome with nested exchangeable correlation structure
### using Quasi-Poisson distribution
#####
### MAEE
est_mae_ind_cnt_quasipoisson = geemae(
  y = sampleSWCRT$y_bin,
  X = X, id = id, Z = Z,
  family = "quasipoisson",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = TRUE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_mae_ind_cnt_quasipoisson)

### GEE
est_uee_ind_cnt_quasipoisson = geemae(
  y = sampleSWCRT$y_bin,
  X = X, id = id, Z = Z,
  family = "quasipoisson",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = FALSE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_uee_ind_cnt_quasipoisson)

## This will elapse longer.
#####
### Example 2): simulated SW-CRT with larger cluster-period sizes (20~30)
#####

sampleSWCRT <- sampleSWCRTLarge

#####
### Individual-level id, period, outcome, and design matrix ###
#####

id <- sampleSWCRT$id
period <- sampleSWCRT$period
X <- as.matrix(sampleSWCRT[, c("period1", "period2", "period3", "period4", "period5", "treatment")])

m <- as.matrix(table(id, period))
n <- dim(m)[1]
t <- dim(m)[2]
### design matrix for correlation parameters
Z <- createzCrossSec(m)

#####
### (1) Matrix-adjusted estimating equations and GEE
### on continous outcome with nested exchangeable correlation structure
#####

```

```

### MAEE
est_mae_ind_con <- geemae(
  y = sampleSWCRT$y_con, X = X, id = id,
  Z = Z, family = "continuous",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = TRUE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_mae_ind_con)

### GEE
est_uee_ind_con <- geemae(
  y = sampleSWCRT$y_con, X = X, id = id,
  Z = Z, family = "continuous",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = FALSE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_uee_ind_con)

#####
### (2) Matrix-adjusted estimating equations and GEE
### on binary outcome with nested exchangeable correlation structure
#####

### MAEE
est_mae_ind_bin <- geemae(
  y = sampleSWCRT$y_bin, X = X, id = id,
  Z = Z, family = "binomial",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = TRUE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_mae_ind_bin)

### GEE
est_uee_ind_bin <- geemae(
  y = sampleSWCRT$y_bin, X = X, id = id,
  Z = Z, family = "binomial",
  maxiter = 500, epsilon = 0.001,
  printrange = TRUE, alpadj = FALSE,
  shrink = "ALPHA", makevone = FALSE
)
print(est_uee_ind_bin)

```

**Description**

The print format for cpgeeSWD output

**Usage**

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

**Arguments**

x	The object of cpgeeSWD output
...	further arguments passed to or from other methods

**Value**

The output from `print`

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<code>print.geemae</code>	<i>The print format for geemae output</i>
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**Description**

The print format for geemae output

**Usage**

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

**Arguments**

x	The object of geemae output
...	further arguments passed to or from other methods

**Value**

The output from `print`



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sampleSWCRTLARGE	<i>simulated large SW-CRT data</i>
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**Description**

Simulated cross-sectional individual-level SW-CRT data with 12 clusters and 5 periods. The cluster-period size is uniformly distributed between 20 and 30. The correlated binary and continuous outcomes are used for analysis as examples.

**Format**

A data frame with 1508 rows and 10 variables:

**period1** indicator of being at period 1  
**period2** indicator of being at period 2  
**period3** indicator of being at period 3  
**period4** indicator of being at period 4  
**period5** indicator of being at period 5  
**treatment** indicator of being treated  
**id** cluster identification number  
**period** period order number  
**y\_bin** binary outcome variable  
**y\_con** continuous outcome variable

---

sampleSWCRTSMALL	<i>simulated small SW-CRT data</i>
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**Description**

Simulated cross-sectional individual-level SW-CRT data with 12 clusters and 4 periods. The cluster-period size is uniformly distributed between 5 and 10. The correlated binary and continuous outcomes are used for analysis as examples.

**Format**

A data frame with 373 rows and 9 variables:

**period1** indicator of being at period 1  
**period2** indicator of being at period 2  
**period3** indicator of being at period 3  
**period4** indicator of being at period 4  
**treatment** indicator of being treated

**id** cluster identification number  
**period** period order number  
**y\_bin** binary outcome variable  
**y\_con** continuous outcome variable

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simbinCLF	<i>Generating Correlated Binary Data using the Conditional Linear Family Method.</i>
-----------	--

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### Description

simbinCLF generates correlated binary data using the conditional linear family method (Qaqish, 2003). It simulates a vector of binary outcomes according the specified marginal mean vector and correlation structure. Natural constraints and compatibility between the marginal mean and correlation matrix are checked.

### Usage

```
simbinCLF(mu, Sigma, n = 1)
```

### Arguments

mu	a mean vector when n = 1 or is NULL, otherwise a list of mean vectors for the n clusters
Sigma	a correlation matrix when n = 1 or is NULL, otherwise a list of correlation matrices for the n clusters
n	number of clusters. The default is 1

### Value

y a vector of simulated binary outcomes for n clusters.

### Author(s)

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### References

Qaqish, B. F. (2003). A family of multivariate binary distributions for simulating correlated binary variables with specified marginal means and correlations. *Biometrika*, 90(2), 455-463.

Preisser, J. S., Qaqish, B. F. (2014). A comparison of methods for simulating correlated binary variables with specified marginal means and correlations. *Journal of Statistical Computation and Simulation*, 84(11), 2441-2452.

**Examples**

```
#####
# Simulate 2 clusters, 3 periods and cluster-period size of 5 #####
#####

t <- 3
n <- 2
m <- 5

# means of cluster 1
u_c1 <- c(0.4, 0.3, 0.2)
u1 <- rep(u_c1, c(rep(m, t)))
# means of cluster 2
u_c2 <- c(0.35, 0.25, 0.2)
u2 <- rep(u_c2, c(rep(m, t)))

# List of mean vectors
mu <- list()
mu[[1]] <- u1
mu[[2]] <- u2
# List of correlation matrices

## correlation parameters
alpha0 <- 0.03
alpha1 <- 0.015
rho <- 0.8

## (1) exchangeable
Sigma <- list()
Sigma[[1]] <- diag(m * t) * (1 - alpha1) + matrix(alpha1, m * t, m * t)
Sigma[[2]] <- diag(m * t) * (1 - alpha1) + matrix(alpha1, m * t, m * t)

y_exc <- simbinCLF(mu = mu, Sigma = Sigma, n = n)

## (2) nested exchangeable
Sigma <- list()
cor_matrix <- matrix(alpha1, m * t, m * t)
loc1 <- 0
loc2 <- 0
for (t in 1:t) {
  loc1 <- loc2 + 1
  loc2 <- loc1 + m - 1
  for (i in loc1:loc2) {
    for (j in loc1:loc2) {
      if (i != j) {
        cor_matrix[i, j] <- alpha0
      } else {
        cor_matrix[i, j] <- 1
      }
    }
  }
}
}
```

```

Sigma[[1]] <- cor_matrix
Sigma[[2]] <- cor_matrix

y_nex <- simbinCLF(mu = mu, Sigma = Sigma, n = n)

## (3) exponential decay

Sigma <- list()

### function to find the period of the ith index
region_ij <- function(points, i) {
  diff <- i - points
  for (h in 1:(length(diff) - 1)) {
    if (diff[h] > 0 & diff[h + 1] <= 0) {
      find <- h
    }
  }
  return(find)
}

cor_matrix <- matrix(0, m * t, m * t)
useage_m <- cumsum(m * t)
useage_m <- c(0, useage_m)

for (i in 1:(m * t)) {
  i_reg <- region_ij(useage_m, i)
  for (j in 1:(m * t)) {
    j_reg <- region_ij(useage_m, j)
    if (i_reg == j_reg & i != j) {
      cor_matrix[i, j] <- alpha0
    } else if (i == j) {
      cor_matrix[i, j] <- 1
    } else if (i_reg != j_reg) {
      cor_matrix[i, j] <- alpha0 * (rho^(abs(i_reg - j_reg)))
    }
  }
}

Sigma[[1]] <- cor_matrix
Sigma[[2]] <- cor_matrix

y_ed <- simbinCLF(mu = mu, Sigma = Sigma, n = n)

```

**Description**

simbinPROBIT generates correlated binary data using the multivariate Probit method (Emrich and Piedmonte, 1991). It simulates a vector of binary outcomes according the specified marginal mean vector and correlation structure. Constraints and compatibility between the marginal mean and correlation matrix are checked.

**Usage**

```
simbinPROBIT(mu, Sigma, n = 1)
```

**Arguments**

mu	a mean vector when n = 1 or is NULL, otherwise a list of mean vectors for the n clusters
Sigma	a correlation matrix when n = 1 or is NULL, otherwise a list of correlation matrices for the n clusters
n	number of clusters. The default is 1

**Value**

y a vector of simulated binary outcomes for n clusters.

**Author(s)**

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**References**

Emrich, L. J., & Piedmonte, M. R. (1991). A method for generating high-dimensional multivariate binary variates. *The American Statistician*, 45(4), 302-304.

Preisser, J. S., Qaqish, B. F. (2014). A comparison of methods for simulating correlated binary variables with specified marginal means and correlations. *Journal of Statistical Computation and Simulation*, 84(11), 2441-2452.

**Examples**

```
#####
# Simulate 2 clusters, 3 periods and cluster-period size of 5 #####
#####

t <- 3
n <- 2
m <- 5

# means of cluster 1
u_c1 <- c(0.4, 0.3, 0.2)
u1 <- rep(u_c1, c(rep(m, t)))
# means of cluster 2
```

```

u_c2 <- c(0.35, 0.25, 0.2)
u2 <- rep(u_c2, c(rep(m, t)))

# List of mean vectors
mu <- list()
mu[[1]] <- u1
mu[[2]] <- u2
# List of correlation matrices

## correlation parameters
alpha0 <- 0.03
alpha1 <- 0.015
rho <- 0.8

## (1) exchangeable
Sigma <- list()
Sigma[[1]] <- diag(m * t) * (1 - alpha1) + matrix(alpha1, m * t, m * t)
Sigma[[2]] <- diag(m * t) * (1 - alpha1) + matrix(alpha1, m * t, m * t)
y_exc <- simbinPROBIT(mu = mu, Sigma = Sigma, n = n)

## (2) nested exchangeable
Sigma <- list()
cor_matrix <- matrix(alpha1, m * t, m * t)
loc1 <- 0
loc2 <- 0
for (t in 1:t) {
  loc1 <- loc2 + 1
  loc2 <- loc1 + m - 1
  for (i in loc1:loc2) {
    for (j in loc1:loc2) {
      if (i != j) {
        cor_matrix[i, j] <- alpha0
      } else {
        cor_matrix[i, j] <- 1
      }
    }
  }
}

Sigma[[1]] <- cor_matrix
Sigma[[2]] <- cor_matrix
y_nex <- simbinPROBIT(mu = mu, Sigma = Sigma, n = n)

## (3) exponential decay

Sigma <- list()

### function to find the period of the ith index
region_ij <- function(points, i) {
  diff <- i - points
  for (h in 1:(length(diff) - 1)) {
    if (diff[h] > 0 & diff[h + 1] <= 0) {
      find <- h
    }
  }
}

```

```
    }
  }
  return(find)
}

cor_matrix <- matrix(0, m * t, m * t)
useage_m <- cumsum(m * t)
useage_m <- c(0, useage_m)

for (i in 1:(m * t)) {
  i_reg <- region_ij(useage_m, i)
  for (j in 1:(m * t)) {
    j_reg <- region_ij(useage_m, j)
    if (i_reg == j_reg & i != j) {
      cor_matrix[i, j] <- alpha0
    } else if (i == j) {
      cor_matrix[i, j] <- 1
    } else if (i_reg != j_reg) {
      cor_matrix[i, j] <- alpha0 * (rho^(abs(i_reg - j_reg)))
    }
  }
}
Sigma[[1]] <- cor_matrix
Sigma[[2]] <- cor_matrix
y_ed <- simbinPROBIT(mu = mu, Sigma = Sigma, n = n)
```

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