Package 'BAREB'

December 18, 2024

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Type Package
Title A Bayesian Repulsive Biclustering Model for Periodontal Data
Version 0.1.2
Description Simultaneously clusters the Periodontal diseases (PD) patients and their tooth sites after taking the patient- and site-level covariates into consideration. 'BAREB' uses the determinantal point process (DPP) prior to induce diversity among different biclusters to facilitate parsimony and interpretability. Essentially, 'BAREB' is a clusterwise linear model based on Yuliang (2020) <doi:10.1002 sim.8536="">.</doi:10.1002>
License GPL-3
Encoding UTF-8
LazyData true
RoxygenNote 7.3.2
LinkingTo Rcpp, RcppArmadillo
Imports Rcpp
NeedsCompilation yes
Depends R (>= $3.5.0$)
Repository CRAN
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Date/Publication 2024-12-18 16:10:08 UTC
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BAREI	B The BAREB package: summary information	

Description

This package implements BAREB (A Bayesian Repulsive Biclustering Model for Periodontal Data)

Value

This is the summary page. No return value.

Main Features

The functions in the pacakge implement BAREB. BAREB can simultaneously cluster periodontal disease patients and their tooth site after taking the patient- and site-level covariates into consideration. BAREB uses the determinantal point process prior to induce diversity among different biclusters to facilitate parsimony and interpretability. In addition, since periodontal diseases are the leading cause for tooth loss, the missing data mechanism is non-ignorable. Such nonrandom missingness is incorporated into BAREB.

const 3

Functions

The main functions are updateBeta, update.theta.beta, update.theta.gamma, updatec, updateE, updateGamma, updatemu, updatemustar, updateR, updateZstar, update_RJ, update_sigma_square, update_w, and update_w_beta; other functions intended for direct access by the user are: kernelC and updateC. There are undocumented functions which are called by these.

Requirements

R version \geq 3.4.3. Packages Rcpp and RcppArmadillo are used so that complicated functions are implemented in C++ to speed up.

Version

This is version 0.1.2.

Licence

This package and its documentation are usable under the terms of the "GNU General Public License", a copy of which is distributed with the package.

Author(s)

Yuliang Li (Dept Applied Mathematics and Statistics, Johns Hopkins University, USA) and Yanxun Xu (Dept Applied Mathematics and Statistics, Johns Hopkins University, USA) and Dipankar Bandyopadhyay (Dept Biostatistics, Virginia Commonwealth University, USA) . Please send comments, error reports, etc. to the maintainer (Yuliang) via email.

Description

The function to compute the normalizing constant for Determinantal Point Process (DPP)

Usage

```
const(theta, tau)
```

Arguments

theta The parameter that controls how repulsive the DPP is

tau The parameter we fixed at a large number

Value

The computed normalizing constant

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Init

The initial value of patient- and site-level covariates for simulation

Description

These data record the initial value of patient- and site-level covariates in simulation in the paper "BAREB: A Bayesian Repulsive Biclustering Model for Periodontal Data". It is obtained by simple linear regression.

The variables are:

Init It has two parts: the initial value of patient-level covariates, Beta; the initial value of site-level covariates, Gamma

Usage

```
data("Init")
```

Examples

```
# output patient level covariates
data("Init")
Init$Beta
Init$Gamma
```

kernelC

The function to get the kernel function value

Description

This function take two configurations x and y, two parameters of the kernel function and returns its kernel function value.

Usage

```
kernelC(x,y,theta,tau)
```

Arguments

x a numeric vector, representing one configuration
y a numeric vector, representing one configuration
theta a parameter of the DPP's kernel function
tau a parameter of the DPP's kernel function

Value

kernelC(x,y,theta,tau) returns the value of the kernel function

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Author(s)

Yuliang Li

See Also

update_RJ for a complete example for all functions in this package.

Examples

```
x <- rnorm(5)
y <- rnorm(5) + 1
kernelC(x,y,1,1)</pre>
```

likeli_theta

likeli_theta

Description

The function computes the likelihood of the hyperparameter

Usage

```
likeli_theta(theta, tau, Beta)
```

Arguments

theta The parameter that controls how repulsive the DPP is

The parameter we fixed at a large number

Beta The matrix that records the samples

Value

The likelihood

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mulation observation
r

Description

These data record the simulation observation (observed value and missing indicator) and covariates in the paper "BAREB: A Bayesian Repulsive Biclustering Model for Periodontal Data".

The variables are:

delta the missing indicator matrix
 X the patient level covariates
 Y the observed (CAL) value matrix
 Z the site level covariates

Usage

```
data("obs")
```

Examples

```
# output patient level covariates
data("obs")
obs$X
```

RJi

Function to update the number of site level clusters for one patient level cluster via RJMCMC

Description

This function updates the number of site level clusters for one patient level cluster via RJMCMC. Only used for updateRJ function.

Usage

```
RJi(w, K, Gamma, Beta, X, Y, Z, R, delta, mu, mu_star, c, sigma_square, C, theta, tau, m, n,q,T0, hyper_delta)
```

Arguments

W	The weights for site level clusters. Should be a vector.
K	The number of site level cluster for the patient level cluster.
Gamma	The linear coefficients for site level covariates. Should be a matrix.
Beta	The linear coefficients for patient level covariates. Should be a vector.
Χ	The design matrix for patient level clusters.

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Y The observed CAL value matrix

Z The design matrix for site level clusters.

R The current site level clustering membership

delta The missing indicator matrix

mu The current CAL mean vector

mu_star The latent value vector for missingness model

c The linear coefficients for missingness model

sigma_square The current noise variance

C The DPP related kernel matrices. Should be an array of 3 dimensions

theta The DPP hyper-parameter for site level

tau A fixed DPP hyper-parameter, which we suggest high value, say 10⁵

m The number of sites

n The number of patients

The number of site level covariates

To The number of teeth

hyper_delta The hyper-parameter with default value being 1

Value

RJi(w, K, Gamma, Beta, X, Y, Z, R, delta, mu, mu_star, c, sigma_square, C, theta, tau, m, n,q,T0, hyper_delta) returns a list with following variables:

K The numbers of site level clusters

w The weights for site level clusters

Gamma Linear coefficients for site level covariates

R The site level clusteirng membership

C The DPP related kernel matrice

Author(s)

Yuliang Li

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RJi_empty	Function to update the number of site level clusters for one patient level cluster via RJMCMC, when there is no patient in this cluster
	······································

Description

This function updates the number of site level clusters for one patient level cluster via RJMCMC, when there is no patient in this cluster. Only used for updateRJ function.

Usage

```
RJi_empty(w, K, Gamma, R, C, theta, tau, m, q, hyper_delta)
```

Arguments

W	The weights for site level clusters. Should be a vector.
K	The number of site level cluster for the patient level cluster.
Gamma	The linear coefficients for site level covariates. Should be a matrix.
R	The current site level clustering membership
С	The DPP related kernel matrices. Should be an array of 3 dimensions
theta	The DPP hyper-parameter for site level
tau	A fixed DPP hyper-parameter, which we suggest high value, say 10^5
m	The number of sites
q	The number of site level covariates
hyper_delta	The hyper-parameter with default value being 1

Value

RJi_empty(w, K, Gamma, R, C, theta, tau, m, q, hyper_delta) returns a list with following variables:

K The numbers of site level clusters

Gamma Linear coefficients for site level covariates

R The site level clusteirng membership

C The DPP related kernel matrice

The weights for site level clusters

Author(s)

Yuliang Li

truth 9

truth	The simulation truth	

Description

These data record the simulation truth in the paper "BAREB: A Bayesian Repulsive Biclustering Model for Periodontal Data". It includes the true simulated parameters.

The variables are:

S	the number of patient level clusters
E	the clustering membership of patient level
K	the numbers of site level clusters
R	the site level clustering membership
Beta	the patient level linear coefficients
Gamma	the site level linear coefficients
mu	the underlying mean for CAL values
sigma_square	the variance of noise for CAL values
noise	the noise for CAL values
С	the parameter for missingness model
mu.star	the mean of latent values for missingness model
z.star	the latent values for missingness model

Usage

```
data("truth")
```

Examples

```
# output true patient level clustering membership
data("truth")
truth$E

#get the details of the list
str(truth)
```

updateBeta Function to update patient level linear coefficients in the BAREB model

Description

This function takes current parameters and observed data, gives an updated patient level linear coefficients.

10 updateBeta

Usage

```
updateBeta(X, Y, Z,
    delta, Beta, Gamma, E, R,
    S, Ds, mustar, mu,
    sigma, c, C,
    step, runif,
    n, m, T0, p, q, D,
    theta,tau)
```

Arguments

X the patient level covariate matrix

Y the CAL observation matrix, with missing values

Z the site level covariate matrix

delta the missing indicator matrix, with 1 means missing

Beta current patient level linear coefficients matrix

Gamma current site level linear coefficients array

E current patient level clustering vector

R current site level clustering matrix

S number of patient level clusters

Ds a vector recording numbers of site level clusters

mustar current matrix of latent value for missingness model

mu current estimated mean matrix for CAL

sigma current estimated noise variance

c current c for missingness model. It is a vector

C current kernel matrix for DPP step a matrix of steps for M-H

runif a matrix of uniform random variables for deciding whether to accept new pro-

posed point in M-H

n number of patients
m number of sites
T0 number of teeth

p dimension of patient level covariatesq dimension of site level covariates

D the D matrix in the paper

theta parameter for DPP tau parameter for DPP

updateC 11

Value

```
updateBeta(X, Y, Z, delta, Beta, Gamma, E, R, S, Ds, mustar, mu, sigma, c, C, step, runif, n, m, T0, p, q, D, theta, tau) returns a list with following variables:
```

C the updated kernel matrix computed by updated Beta

Beta the updated patient level linear coefficients

mu the updated mu computed by updated Beta

mustar the updated mustar computed by updated Beta

Author(s)

Yuliang Li

See Also

update_RJ for a complete example for all functions in this package.

updateC

Function to obtain the kernel matrix of the determinantal point process

Description

This function takes a matrix and two parameters of kernel function for the determinantal point process (DPP) and gives the kernel matrix for that DPP.

Usage

```
updateC(Z,theta,tau)
```

Arguments

Z a matrix, whose rows stand for configurations of the DPP.

theta a parameter of the DPP's kernel function tau a parameter of the DPP's kernel function

Value

```
updateC(Z, theta, tau) returns the kernel matrix
```

Author(s)

Yuliang Li

See Also

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Examples

```
Z <- matrix(rnorm(15), nrow = 5)
updateC(Z,1,1)</pre>
```

updatec

Function to update c in missingess model

Description

This function takes current parameters, gives updated c in missingess model. Note a double type of value is returned

Usage

```
updatec(Zstar, mu, D, sigmac, sigma_square, n, T0)
```

Arguments

Zstar generated latent value for missingness model

mu current estimated mean matrix for CAL

D the D matrix in the paper

sigmac hyperparamter for c (variance)
sigma_square current estimated noise variance

n number of patientsT0 number of teeth

Value

```
updatec(Zstar, mu, D, sigmac, n, T0) returns the updated c in missingess model.
```

Author(s)

Yuliang Li

See Also

updateE 13

updateE	Function to update patient level clustering in the BAREB model

Description

This function takes current parameters and observed data, gives an updated patient level clustering.

Usage

```
updateE( Beta, Gamma,w,
    X, Y, Z, delta,
    E, R, S, Ds,
    mu, mustar,
    sigma, c,
    n, m, T0, p, q, D)
```

Arguments

Beta	current patient level linear coefficients matrix
Gamma	current site level linear coefficients array
W	current patient level clustering prior prob, a vector
Χ	the patient level covariate matrix
Υ	the CAL observation matrix, with missing values
Z	the site level covariate matrix
delta	the missing indicator matrix, with 1 means missing
E	current patient level clustering vector
R	current site level clustering matrix
S	number of patient level clusters
Ds	a vector recording numbers of site level clusters
mu	current estimated mean matrix for CAL
mustar	current matrix of latent value for missingness model
sigma	current estimated noise variance
С	current c for missingness model. It is a vector
n	number of patients
m	number of sites
Т0	number of teeth
p	dimension of patient level covariates
q	dimension of site level covariates
D	the D matrix in the paper

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Value

```
updateE(Beta, Gamma, w, X, Y, Z, delta, E, R, S, Ds, mu, mustar, sigma, c, n, m, T0, p, q, D) returns a list with following variables:
```

E the updated patient level clustering

Ds new vector recording the numbers of site level clusters

mu the updated mu computed by updated E mustar the updated mustar computed by updated E

Author(s)

Yuliang Li

See Also

update_RJ for a complete example for all functions in this package.

updateGamma

Function to update site level linear coefficients in the BAREB model

Description

This function takes current parameters and observed data, gives an updated site level linear coefficients.

Usage

```
updateGamma(X, Y, Z, delta,
    Beta, Gamma, E, R,
    S, Ds, mu, mustar,
    sigma, c, step, runif,
    n, m, T0, p, q, D, theta,tau)
```

Arguments X

	<u>*</u>
Υ	the CAL observation matrix, with missing values
Z	the site level covariate matrix
delta	the missing indicator matrix, with 1 means missing
Beta	current patient level linear coefficients matrix
Gamma	current site level linear coefficients array
Е	current patient level clustering vector
R	current site level clustering matrix
S	number of patient level clusters

the patient level covariate matrix

updateGamma 15

Ds a vector recording numbers of site level clusters

mu current estimated mean matrix for CAL

mustar current matrix of latent value for missingness model

sigma current estimated noise variance

c current c for missingness model. It is a vector

step an array of steps for M-H

runif an array of uniform random variables for deciding whether to accept new pro-

posed point in M-H

n number of patients

m number of sites
To number of teeth

p dimension of patient level covariates

q dimension of site level covariates

D the D matrix in the paper

theta parameter for DPP tau parameter for DPP

Value

updateGamma(X, Y, Z, delta, Beta, Gamma, E, R, S, Ds, mu, mustar, sigma, c, step, runif, n, m, T0, p, q, D, theta, tau) returns a list with following variables:

Gamma the updated site level linear coefficients

mu the updated mu computed by updated Gamma

mustar the updated mustar computed by updated Gamma

Author(s)

Yuliang Li

See Also

16 updatemu

updatemu	Function to update estimated mean CAL values based on current parameters

Description

This function takes current parameters, gives an estimated mean CAL values.

Usage

```
updatemu(R, Z, X, Gamma, K, Beta, E, m,n,p, q)
```

Arguments

R	current site level clustering matrix
Z	the site level covariate matrix
X	the patient level covariate matrix
Gamma	current site level linear coefficients array
K	a vector recording numbers of site level clusters
Beta	current patient level linear coefficients matrix
E	current patient level clustering vector
m	number of sites
n	number of patients
р	dimension of patient level covariates
q	dimension of site level covariates

Value

```
updatemu(R, Z, X, Gamma, K, Beta, E, m,n,p, q) returns the updated estimated mean CAL matrix.
```

Author(s)

Yuliang Li

See Also

updatemustar 17

updatemustar

Function to update mean latent values for missingness model

Description

This function takes current parameters, gives updated mean latent values for missingness model

Usage

```
updatemustar(mu, c, n, T0, D)
```

Arguments

mu	current estimated mean matrix for CAL
С	current c for missingness model
n	number of patients
TØ	number of teeth
D	the D matrix in the paper

Value

updatemustar(mu, c, n, T0, D) returns the updated mean latent values for missingness model.

Author(s)

Yuliang Li

See Also

update_RJ for a complete example for all functions in this package.

updateR

Function to update site level clustering in the BAREB model

Description

This function takes current parameters and observed data, gives an updated site level clustering.

Usage

```
updateR( w ,Gamma,Beta,
    Y, Z, delta,
    mu, mu_star,
    c, S, sigma_square,
    K, E, X,
    m,n, q, p, T0)
```

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Arguments

W	current site level clustering prior prob, a matrix
Gamma	current site level linear coefficients array

Beta current patient level linear coefficients matrix

Y the CAL observation matrix, with missing values

Z the site level covariate matrix

delta the missing indicator matrix, with 1 means missing

mu current estimated mean matrix for CAL

mu_star current matrix of latent value for missingness model

c current c for missingness model
S number of patient level clusters
sigma_square current estimated noise variance

K a vector recording numbers of site level clusters

E current patient level clustering vector

X the patient level covariate matrix

m number of sitesn number of patients

p dimension of patient level covariates
q dimension of site level covariates

T0 number of teeth

Value

updateR(w,Gamma,Beta,Y,Z,delta,mu,mu_star,c,S,sigma_square,K,E,X,m,n,p,q,T0) returns the updated site level clustering.

Author(s)

Yuliang Li

See Also

updateZstar 19

updateZstar	Function to generate new latent values for missingness model

Description

This function takes current parameters, gives updated c in missingess model. Note a double type of value is returned

Usage

```
updateZstar(mu_star, delta, n, T0)
```

Arguments

mu_star current matrix of latent value for missingness model
delta the missing indicator matrix, with 1 means missing
n number of patients
To number of teeth

Value

```
updateZstar(mu_star, delta, n, T0) returns a matrix of new generated latent values.
```

Author(s)

Yuliang Li

See Also

update_RJ for a complete example for all functions in this package.

Description

Update the number of site level clusters for each patient level cluster via RJMCMC

Usage

```
update_RJ(
 W,
 Κ,
 Gamma,
 Beta,
 Ε,
 Ζ,
 Χ,
 R,
 mu,
 mu_star,
 Υ,
 delta,
  С,
  sigma_square,
 С,
 S,
  theta,
  tau,
 q,
 m,
 Τ0,
 hyper_delta = 1
)
```

Arguments

sigma_square

C	
W	The weights for site level clusters. Should be a matrix with S rows.
K	The number of site level cluster for each patient level cluster. Should be a vector of length S
Gamma	The linear coefficients for site level covariates. Should be a 3-dimensional array.
Beta	The linear coefficients for patient level covariates. Should be a matrix with S rows
Е	A vector that records the current clustering membership.
Z	The design matrix for site level clusters.
Χ	The design matrix for patient level clusters.
R	The current site level clustering membership
mu	The current CAL mean matrix
mu_star	The latent value matrix for missingness model
Υ	The observed CAL value matrix
delta	The missing indicator matrix
С	The linear coefficients for missingness model

The current noise variance

С	The DPP related kernel matrices. Should be an array of 3 dimensions
S	The number of patient level clusters.
theta	The DPP hyper-parameter for site level
tau	A fixed DPP hyper-parameter, which we suggest high value, say 10^5
q	The number of site level covariates
m	The number of sites
T0	The number of teeth
hyper_delta	The hyper-parameter with default value being 1

Value

A list with following updated parameters:

K	The numbers of site level clusters
W	The weights for site level clusters
Gamma	Linear coefficients for site level covariates
R	The site level clusteirng membership
С	The DPP related kernel matrice

Examples

```
library(BAREB)
data("obs")
X <- obs$X
Y <- obs$Y
Z <- obs$Z
delta <- obs$delta
data("truth")
set.seed(1)
n<-80
m<-168
T0<-28
q<-3
p<-3
S<-3
theta1 <- theta2 <- 5
tau <- 100000
D<-matrix(0, nrow = T0, ncol = m)</pre>
for(i in 1:T0){
  indi<-1:6
  indi<-indi+6*(i-1)</pre>
  D[i,indi]<-rep(1/6,6)</pre>
}
nu_gamma<-0.05
nu_beta <- 0.05
data("Init")
```

```
Beta0 <- Init$Beta
Gamma0 <- Init$Gamma
Niter<-10
record <- NULL
record$E<-matrix(NA,nrow = Niter, ncol = n)</pre>
record$R<-array(NA, dim = c(Niter, S, m))</pre>
record$Gamma <-array(NA,dim = c(Niter, 10, q, S))</pre>
record$Beta <- array(NA,dim = c(Niter, S, p))</pre>
record$K <- matrix(NA,nrow = Niter, ncol = S)</pre>
record$sigma_square <-rep(NA,Niter)</pre>
record$theta1<-rep(0,Niter)</pre>
record$theta2<-rep(0,Niter)</pre>
record$c<-matrix(0,nrow = Niter,ncol = T0)</pre>
record$mu<-array(NA,dim = c(Niter,n,m))</pre>
record$w_beta <- array(NA, dim = c(Niter, S))</pre>
record$w <- array(NA, dim = c(Niter, S, 10))</pre>
set.seed(1)
E<-sample.int(S,n,TRUE)</pre>
Beta <- matrix(NA,nrow = S, ncol = p)</pre>
Beta[1,] <- Beta[2,] <- Beta[3,] <- Beta0</pre>
Beta<- Beta + matrix(rnorm(S*p, 0, 1) , nrow = S, ncol = p)</pre>
Gamma \leftarrow array(NA, dim = c(10,q,S))
Gamma[1,,1] \leftarrow Gamma[2,,1] \leftarrow Gamma[3,,1] \leftarrow Gamma0
Gamma[,,2] \leftarrow Gamma[,,3] \leftarrow Gamma[,,1]
Gamma \leftarrow Gamma + array(rnorm(10*q*S, 0, 5), dim = c(10, q, S))
K \leftarrow rep(3,S)
R <- matrix(NA,nrow = S, ncol = m)</pre>
R[1,] \leftarrow R[2,] \leftarrow R[3,] \leftarrow sample.int(3,m,TRUE)
mu < -updatemu(R,Z,X,Gamma,K,Beta,E,m,n,p,q)
mu_star<-updatemustar(mu,rep(0.01,2),n,T0,D)</pre>
z_star<-updateZstar(mu_star,delta,n,T0)</pre>
sigma_square <- 10
C<-array(NA,dim = c(10,10,S))
w<-matrix(NA,nrow = S, ncol = 10)</pre>
w_beta<-rep(1/S,S)</pre>
for(i in 1:S){
 C[1:K[i],1:K[i],i] < -updateC(Gamma[1:K[i],,i],theta2,tau)
  w[i, 1:K[i]]<-rep(1/K[i],K[i])</pre>
c < -c(0, 0.01)
start <- Sys.time()</pre>
for(iter in 1:Niter){
  c<-updatec(z_star, mu,D, 100,sigma_square, n, T0)</pre>
  mu_star<-updatemustar(mu,c,n,T0,D)</pre>
  z_star<-updateZstar(mu_star,delta,n, T0)</pre>
  w <- update_w(K, R, S)</pre>
  R <- updateR(w, Gamma, Beta,</pre>
                 Y, Z, delta, mu, mu_star, c[2], S,
                 sigma_square, K, E, X,
                 m, n, q, p, T0)
  for(i in unique(E)){
    ind<-sort(unique(R[i,]))</pre>
```

```
KK<-length(ind)
  Gamma_temp<-Gamma[ind,,i]</pre>
  Gamma[,,i]<-NA
  Gamma[1:KK,,i]<-Gamma_temp</pre>
  w_temp<-w[i,ind]</pre>
  w_temp<-w_temp/sum(w_temp)</pre>
  w[i,]<-NA
  w[i,1:KK] < -w_temp
  for(k in 1:KK){
    R[i, which(R[i, ]==ind[k])] < -k
  K[i] < -KK
mu<-updatemu(R,Z,X,Gamma,K,Beta,E,m,n,p,q)</pre>
mu_star<-updatemustar(mu,c,n,T0,D)</pre>
step <- array(rnorm(max(K) * S *q, 0, nu_gamma),dim=c( max(K), q,S))</pre>
run<- array(runif(max(K) * S *q, 0, 1), dim=c(max(K), q,S))
 A < -update Gamma(X,Y,Z,delta,Beta,Gamma,E,R,S,K,mu,mu\_star,sigma\_square,rep(c,T0), \\
                step, run, n, m, T0, p, q, D, theta2, tau)
Gamma<-A$Gamma
mu<-A$mu
mu_star<-A$mustar</pre>
for(i in 1:S){
  if(K[i]==1){
    Gammai = t(as.matrix(Gamma[1:K[i],,i]))
    C[1:K[i],1:K[i],i]<-updateC(Gammai,theta2,tau)</pre>
  else{
    C[1:K[i],1:K[i],i] \leftarrow updateC(Gamma[1:K[i],i],theta2,tau)
  }
}
A<-update_RJ(w, K, Gamma, Beta, E,
              Z, X, R, mu, mu_star, Y, delta, c,sigma_square, C,
              S, theta2, tau, q, m, T0)
K<-A$K
w<-A$w
Gamma<-A$Gamma
R<-A$R
C<-A$C
mu<-updatemu(R,Z,X,Gamma,K,Beta,E,m,n,p,q)</pre>
mu_star<-updatemustar(mu,rep(c,T0),n,T0,D)</pre>
C_beta<-updateC(Beta, theta1, tau)</pre>
step<-matrix(rnorm(S*p,0,nu_beta),nrow = S)</pre>
runif<-matrix(runif(S*p,0,1),nrow = S)</pre>
A<- updateBeta( X,Y, Z, delta,
            Beta, Gamma, E,R,S,K,mu_star, mu,sigma_square,rep(c,T0), C_beta, step,runif,
                  n, m, T0, p, q, D,
                  theta1, tau)
Beta<-A$Beta
mu<-A$mu
mu_star<-A$mustar</pre>
record$Beta[iter,,]<-Beta</pre>
A<-updateE(Beta,Gamma, w_beta, X,Y,Z,delta,E, R, S,K, mu, mu_star,sigma_square,rep(c,T0),
```

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```
n, m, T0, p, q, D)
E<-A$E
mu<-A$mu
mu_star<-A$mustar
K<-A$Ds
w_beta<- update_w_beta(S, w_beta, E)
theta1<-update_theta_beta(theta1, tau, Beta)
theta2<-update_theta_gamma(theta2, tau, Gamma, S, K)
sigma_square <- update_sigma_squre(Y, mu)
}</pre>
```

update_sigma_squre

update_sigma_squre

Description

Update the noise variance

Usage

```
update_sigma_squre(Y, mu, a = 1, b = 1)
```

Arguments

Υ	The CAL observation matrix, with missing values
mu	Current estimated mean matrix for CAL
a	The hyper-parameter with default value being 1
b	The hyper-parameter with default value being 1

Value

Updated noise variance

See Also

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|--|

Description

Update the DPP hyper-parameter for patient level

Usage

```
update_theta_beta(theta, tau, Beta, sig = 10)
```

Arguments

theta The DPP hyper-parameter for patient leve
--

tau A fixed DPP hyper-parameter, which we suggest high value, say 10^5

Beta The linear coefficients for patient level covariates. Should be a matrix with S

rows

sig The hyper-parameter with default value being 10

Value

updated DPP hyper-parameter for patient level

See Also

update_RJ for a complete example for all functions in this package.

```
update_theta_gamma update_theta_gamma
```

Description

Update the DPP hyper-parameter for site level

Usage

```
update_theta_gamma(theta, tau, Gamma, S, Ds, sig = 10)
```

Arguments

theta	The DPP hyper-parameter for patient level
tau	A fixed DPP hyper-parameter, which we suggest high value, say 10^5
Gamma	The linear coefficients for site level covariates. Should be a 3-dimensional array.
S	The number of patient level clusters
Ds	The number for site level clusters for each patient level cluster
sig	The hyper-parameter with default value being 10

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Value

updated DPP hyper-parameter for site level

See Also

update_RJ for a complete example for all functions in this package.

Description

This function updates the weights for site level clusters

Usage

```
update_w(K, R, S, hyper_delta = 1)
```

Arguments

K	The number of site level cluster for each patient level cluster. Should be a vector of length S
R	The current site level clustering membership. Should be a matrix with S rows.
S	The number of patient level clusters.
hyper_delta	The hyper-parameter with default value being 1

Details

It returns a matrix with 10 columns. For example, first patient cluster has 2 site level clusters. The first row's first 2 elements give the weights for site level clusters in patient cluster 1. Last 8 elements are NA's

Value

The updated weights for site level clusters. Should be a matrix with S rows.

See Also

update_RJ for a complete example for all functions in this package.

Examples

```
#Suppose we know the number of patient level cluster is 2,
#one has 2 site level clusters and one has 3.
#Use the default value, 1, for hyper-parameter
update_w(K=c(2,3),R=matrix(c(1,2,2,1,1,2,3,2),nrow=2,byrow=TRUE), S=2)
#To change the hyper-parameter to, for example 2
update_w(K=c(2,3),R=matrix(c(1,2,2,1,1,2,3,2),nrow=2,byrow=TRUE), S=2, hyper_delta = 2)
```

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Description

This function updates the weights for each patient level cluster

Usage

```
update_w_beta(S, E, hyper_delta = 1)
```

Arguments

S The number of patient level clusters

E A vector that records the current clustering membership.

hyper_delta The hyper-parameter with default value being 1

Value

The updated weights for each patient level cluster

See Also

update_RJ for a complete example for all functions in this package.

Examples

```
#Suppose we know the number of patient level cluster is 4
#Suppose the current clustering membership indicates 3 patients in cluster 1,
#2 patients in cluster 2, 3 patients in cluster 3, 1 patient in cluster 4
#Use the default value, 1, for hyper-parameter
update_w_beta(S=4,E=c(1,1,1,2,2,3,3,3,4))
#To change the hyper-parameter to, for example 2
update_w_beta(S=4,E=c(1,1,1,2,2,3,3,3,4),hyper_delta = 2)
```

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