Package 'ZIPG'

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Type Package				
Title Zero-Inflated Poisson-Gamma Regression				
Version 1.1				
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Description We provide a flexible Zero-inflated Poisson-Gamma Model (ZIPG) by connecting both the mean abundance and the variability to different covariates, and build valid statistical inference procedures for both parameter estimation and hypothesis testing. These functions can be used to analyze microbiome count data with zero-inflation and overdispersion. The model is discussed in Jiang et al (2023) <doi:10.1080 01621459.2022.2151447="">.</doi:10.1080>				
License GPL (>= 2)				
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ZIPG_CI

Dietary

A diet-microbiome data

Description

A diet-microbiome data with shotgun metagenomic sequencing results of fecal samples and daily dietary records of 34 subjects in 17 consecutive days. See original data from article "Daily Sampling Reveals Personalized Diet-Microbiome Associations in Humans" (Johnson et al. 2019).

Usage

Dietary

Format

A list with OTU table and other covariates

OTU OTU table

COV Major covariates extracted from original data

M Sequencing depth

Source

https://github.com/knights-lab/dietstudy_analyses

ZIPG_CI

Get confidence interval from ZIPG model

Description

Get confidence interval from ZIPG model

Usage

```
ZIPG_CI(ZIPG_res, type = "Wald", CI_type = "normal", alpha = 0.05)
```

Arguments

ZIPG_res	Result from ZIPG_main()
type	Type of hypothesis testing method, 'Wald' or 'bWald'.
CI_type	Type of confidence interval, 'Wald', 'bWald' or 'pbWald'.

alpha We construct (1- alpha)% confidence interval by alpha/2 and (1-alpha/2).

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Value

Table of confidence interval

Examples

```
data(Dietary)
dat = Dietary
ZIPG_res <- ZIPG_main(data = dat$COV,
X = ~ALC01+nutrPC1+nutrPC2, X_star = ~ ALC01,
W = dat$OTU[,100], M = dat$M )
ZIPG_CI(ZIPG_res)</pre>
```

ZIPG_main

Fit zero-inflated poisson-gamma model via EM Algorithm

Description

Fit zero-inflated poisson-gamma model via EM Algorithm

Usage

```
ZIPG_main(
  data,
  W,
  M,
  X,
  X_star,
  return_model = TRUE,
  pbWald_list = NULL,
  bWald_list = NULL
)
```

Arguments

data	Data.frame for covariates of interest
W	Count data
М	Sequencing depth, ZIPG use log(M) as offset by default
Χ	Formula of covariates of differential abundance
X_star	Formula of covariates of differential varibility
return_model	whether return full complete imfomation for fitted model
pbWald_list	A list of arguments for parameteric bootstrap Wald test, B for bootstrap sample size, X0 and X_star0 for formula of covariates included in H0
bWald_list	A list of arguments for non-parameteric bootstrap Wald test, B for bootstrap sample size,

ZIPG_simulate

Value

A list of ZIPG fitted model. Use ZIPG_summary() for a quick look at the results.

Examples

```
data(Dietary)
dat = Dietary
ZIPG_res <- ZIPG_main(data = dat$COV,
X = ~ALC01+nutrPC1+nutrPC2, X_star = ~ ALC01,
W = dat$OTU[,100], M = dat$M )
ZIPG_summary(ZIPG_res)</pre>
```

ZIPG_simulate

Simulate W from ZIPG model

Description

Simulate W from ZIPG model

Usage

```
ZIPG_simulate(
   M,
   X,
   X_star,
   A = 1,
   d,
   d_star,
   parms,
   N,
   zi = TRUE,
   returnU = FALSE
)
```

Arguments

М	Sequencing depth
Χ	Covariates matrix with intercept, n * (d+1)
X_star	Covariates matrix with intercept, n * (d_star+1)
A	no use, reserved for multi-taxa
d	number of covariates in X
d_star	number of covariates in X_star
parms	model paraneters, input c(beta,beta*,gamma)
N	repetition times
zi	whether generate zero-inflated distribution
returnU	whether return fluctuation factor U

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Value

A list of W generated from ZIPG model with input parameter

Examples

```
data(Dietary)
dat = Dietary
sim_M = sample(dat$M,100,replace = TRUE)
sim_pre = rep(sample(rep(c(0,1),each = 10)),each = 5)
sim_PC1_mean = rep(rnorm(20, mean = 0, sd = 1), each = 5)
sim_PC1_error = rnorm(100,0,0.1)
sim_PC1 = sim_PC1_mean + sim_PC1_error
X = as.matrix(cbind(1,data.frame(X1 = sim_pre, X2 = sim_PC1)))
parms = c(-4.23,1,0.45,0.6,1,0,0) #p = 0.5
W_sim <- ZIPG_simulate(M = sim_M, X=X, X_star=X, d=2, d_star=2, parms = parms, N=100)
hist(W_sim$W_list[[1]])
ZIPG_res <- ZIPG_main(data = data.frame(X1 = sim_pre, X2 = sim_PC1),</pre>
X = \sim X1+X2, X_star = \sim X1, W = W_sim$W_list[[2]], M = sim_M)
ZIPG_summary(ZIPG_res)
```

ZIPG_summary

Summary for ZIPG_main() result.

Description

Summary for ZIPG_main() result.

Usage

```
ZIPG_summary(ZIPG_res, type = "Wald")
```

Arguments

Result from ZIPG_main() ZIPG_res type

Type of hypothesis testing method, 'Wald', 'bWald' or 'pbWald'.

Value

pvalue

Examples

```
data(Dietary)
dat = Dietary
ZIPG_res <- ZIPG_main(data = dat$COV,</pre>
X = ~ALC01+nutrPC1+nutrPC2, X_star = ~ ALC01,
W = dat\$OTU[,100], M = dat\$M)
ZIPG_summary(ZIPG_res)
```

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