# Package 'IVGC'

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Type Package

Title Two-step Estimation Using IV Regression and Graph-constrained Regularization Method
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<b>Depends</b> R (>= $3.1.0$ )
Imports mytnorm, glmnet
<b>Description</b> Efficient two-step procedure to achieve variable selection and estimation under an IV regression framework by incorporating gene network information. The method is applicable in both low and high dimensional situations and the cyclical coordinate descent algorithm is used to solve them.
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R topics documented:
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IVGC-package Two-step Estimation Using IV Regression and Graph-constrained Regularization Method

#### **Description**

Efficient two-step procedure to achieve variable selection and estimation under an IV regression framework by incorporating gene network information. The method is applicable in both low and high dimensional situations and the cyclical coordinate descent algorithm is used to solve them.

#### **Details**

Package: IVGC
Type: Package
Version: 1.0.1
Date: 2019-03-12
License: GPL-2

This package use a networkconstrained regularization method under the high-dimensional IV regression framework to address the problem of high correlations between coefficients. The coordinate descent algorithm is applied to fit models. Accepts X,Y,S,G data for regression models, and produces a list including both the Gamma estimator by grouped lasso and the beta estimator by the proposed method and IV method without graph structure. The function is main.

#### Author(s)

Gao Bin, Xu Liu, Hongzhe Li, and Yuehua Cui

#### References

Gao, B., Liu, X., Li, H. and Cui, Y. (2019+) Integrative Analysis of Genetical Genomics Data Incorporating Network Structures. Manuscript.

### **Examples**

```
data(simulatedData)
y <- simulatedData$Y
x <- simulatedData$X
S <- simulatedData$gene
G <- simulatedData$graph
fit <- main(y, x, S, G)</pre>
```

Fit an IV regression model with lasso and graph-constrained regularization

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#### **Description**

This function is a two-step procedure to achieve variable selection and estimation under an IV regression framework by incorporating gene regulatory network structures. Because of high correlations between coefficients, in the first step, the LASSO-type regression is applied to estimate the coefficient matrix. And in the second step, two penalties (LASSO and graph-constrained penalties) are applied to encourage the shrinking coefficients of a pair of connected variable in a network, thus achieving a grouping effect.

### Usage

```
main(y, x, S, G,nlambda=50,nalpha=50,epsilon=1e-6)
```

# Arguments

У	The response, a vector of size $n$
X	The expression data dimension $n \times p$
S	The SNP variable with dimension $n \times q$
G	The graph structure of X with dimension $p \times p$
nlambda	The length of tuning parameter $\lambda$ which is selected automatically. Default is 50.
nalpha	The length of tuning parameter $\alpha$ which is selected automatically. Default is 50.
epsilon	The perturbation when the proposed algorithm is used. Default is epsilon=1e-6.

#### **Details**

The simulated data set is saved in simulatedData.RData, which is list with four values Y, X, gene (S) and graph (G).

# Value

beta_ivgc	the beta estimator by the proposed method
beta_iv	the beta estimator by IV method without graph structure
Gamma_hat	the Gamma estimator by grouped lasso

#### Author(s)

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#### References

Gao, B., Liu, X., Li, H. and Cui, Y. (2019+) Integrative Analysis of Genetical Genomics Data Incorporating Network Structures. Manuscript.

# Examples

```
data(simulatedData)
y <- simulatedData$Y
x <- simulatedData$X
S <- simulatedData$gene
G <- simulatedData$graph
fit <- main(y, x, S, G)</pre>
```

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simulatedData

Genetical Genomics Data Incorporating Network Structures

# **Description**

This is a simulated data on gene expression. Certain correlation structures on genes are also imposed to show the impact of network structures on variable selection and estimation.

#### Usage

```
data("simulatedData")
```

#### **Details**

We simulated a total of p variables of gene expression and considered three group structures on coefficients with five variables in each group. Within each group, the variables are correlated. The strength of the correlation was controlled by the number of effective SNPs they had in common.

Y: the response, a vector of size 600

X: the expression data dimension  $600 \times 100$ 

S: the SNP variable with dimension  $600 \times 100$ 

G: the graph structure of X with dimension  $100 \times 100$ 

#### References

Gao, B., Liu, X., Li, H. and Cui, Y. (2019+) Integrative Analysis of Genetical Genomics Data Incorporating Network Structures. Manuscript.

# **Examples**

data(simulatedData)

y <- simulatedData\$Y

x <- simulatedData\$X

S <- simulatedData\$gene

 ${\tt G} < - {\tt simulatedData\$graph}$ 

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