

# Package: NetGreg (via r-universe)

May 28, 2026

**Type** Package

**Title** Network-Guided Penalized Regression (NetGreg)

**Version** 0.0.4

**Description** A network-guided penalized regression framework that integrates network characteristics from Gaussian graphical models with partial penalization, accounting for both network structure (hubs and non-hubs) and clinical covariates in high-dimensional omics data, including transcriptomics and proteomics. The full methodological details can be found in our publication by Ahn S and Oh EJ (2026)  [<doi:10.1093/bioadv/vbag038>](https://doi.org/10.1093/bioadv/vbag038).

**License** GPL-3

**Encoding** UTF-8

**RoxygenNote** 7.3.2

**Depends** R (>= 3.5.0)

**Imports** huge, glmnet, dplyr, stats, plsgenomics

**Suggests** knitr, rmarkdown, testthat (>= 3.0.0)

**NeedsCompilation** no

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**Repository** <https://cran.r-universe.dev>

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## Contents

identifyHubs . . . . .	2
NetworkGuided . . . . .	3

**Index****4**


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identifyHubs	<i>identifyHubs</i>
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**Description**

A function to identify hub nodes (i.e., genes or proteins) from high-dimensional data using network-based criteria.

**Usage**

```
identifyHubs(X, delta, tau, ebic.gamma = 0.1)
```

**Arguments**

X	A data matrix of dimension $n \times p$ representing samples (rows) by features (columns).
delta	A numeric value indicating the proportion of nodes to considered as hubs in a network.
tau	A user-specified cutoff for the number of hubs.
ebic.gamma	A numeric value specifying the tuning parameter for the extended Bayesian information criterion (eBIC) used in network estimation.

**Value**

A list containing (1) the selected sparse graph structure and model selection results; (2) a data frame of feature names with their associated network characteristics (e.g., degree centrality); and (3) a character vector of top-ranked hub features (e.g., hub genes or proteins).

**Examples**

```
library(plsgenomics)
data(Colon) ## Data from plsgenomics R package
X = data.frame(Colon$X[,1:100]) ## The first 100 genes
Z = data.frame(Colon$X[,101:102]) ## Two clinical covariates
colnames(Z) = c("Z1", "Z2")
Y = as.vector(Colon$X[,1000]) ## Continuous outcome variable

## Apply identifyHubs():
preNG = identifyHubs(X=X, delta=0.05, tau=5, ebic.gamma = 0.1)

## Explore preNG results:
## To display the degree centrality for each node,
## sorted from strongest to weakest.
preNG$assoResults
preNG$hubs ## Returns the names of the identified hub nodes.
```

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NetworkGuided	<i>NetworkGuided</i>
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**Description**

A main function to obtain network-guided penalized regression coefficient estimates.

**Usage**

```
NetworkGuided(Y, X, hubs, Z, nolds = 5)
```

**Arguments**

Y	A continuous outcome variable.
X	A data matrix of dimension $n \times p$ representing samples (rows) by features (columns).
hubs	A vector of hubs identified through identifyHubs function from our package.
Z	A matrix of clinical or demographic covariates.
nolds	A user-specified numeric value for k-fold cross-validation.

**Value**

A vector of network-guided penalized regression coefficients.

**Examples**

```
library(plsgenomics)
data(Colon) ## Data from plsgenomics R package
X = data.frame(Colon$X[,1:100]) ## The first 100 genes
Z = data.frame(Colon$X[,101:102]) ## Two clinical covariates
colnames(Z) = c("Z1", "Z2")
Y = as.vector(Colon$X[,1000]) ## Continuous outcome variable

## Apply identifyHubs():
preNG = identifyHubs(X=X, delta=0.05, tau=5, ebic.gamma = 0.1)

## Explore preNG results:
hubs = preNG$hubs ## Returns the names of the identified hub nodes.

## Use our main NetworkGuided function, to obtain network-guided
## penalized regression coefficient estimates.
NG = NetworkGuided(Y=Y, X=X, hubs=preNG$hubs, Z=Z, nolds=5)
NG$coef
```

# Index

`identifyHubs`, [2](#)

`NetworkGuided`, [3](#)