# Package 'HierFabs'

# April 8, 2021

Type	Package			
Title	A General Framework tion of genomics data	for Identifying Hierachical Interaction of Genomics Data interac-		
Versi	on 0.1.0			
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<b>Description</b> Provides an efficient procedure for fitting the entire solution path adaptively for iden ing hierarchical interaction of genomics data in linear model or cox model under the strong or weak heredity constraint. Lasso penalty is used to generate a sparse solution. Both gene-gene type interactions and gene-environment interactions are supported.			ify-	
Licen	se GPL (>= 2)			
Impo	rts Matrix			
Repo	sitory github			
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Hie	erFabs	A hierarchical Forward and Backward Stagewise (HierFabs) algorithm for identifying hierarchical interaction of genomics data.		
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# Description

A hierarchical Forward and Backward Stagewise (HierFabs) algorithm for identifying hierarchical interaction of genomics data.

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#### Usage

```
HierFabs(
  G,
  у,
  Ε,
  weight = NULL,
  model = c("square", "cox"),
  back = TRUE,
  stoping = TRUE,
  eps = 0.01,
  xi = 10^{-6},
  iter = 3000,
  lambda.min = NULL,
  hier = c("strong", "weak"),
  max_s = NULL,
  diagonal = FALSE,
  status = NULL,
  gamma = NULL
)
```

#### **Arguments**

$\sim$	Gene matrix, each row is an observation vector.
(1	CIEHE HIAHIX. EACH TOW IS AH ODSELVATION VECTOR.

y Response variable.

E An optional environment matrix. If Z is given, the interactions between envi-

ronment and gene are of interest. Otherwise, the gene-gene interactions are of

interest.

weight An optional weights. Default is 1 for each observation.

model A character string representing one of the built-in models.

back An indicator of whether to take backward steps.

stoping An indicator of whether to stop iteration when lambda is less than lambda.min.

eps Step size. Default is 0.01.

xi A tolerate to improve program stability. Default is 10^-6.

iter Maximum number of outer-loop iterations allowed. Default is 3000.

lambda.min Smallest value for lambda, as a fraction of lambda.max.

hier Whether to enforce strong or weak heredity. Default is 'strong'.

max\_s Limit the maximum number of variables in the model.

diagonal An indicator of whether to include "pure" quadratic terms. Work when gene-

gene interactions are of interest.

status A censoring indicator.

gamma A tuning parameter in EBIC.

#### Value

#### A list.

- theta The coefficients of covariates, each column is a solution.
- beta The optimal solution.

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- lambda Lambda sequence.
- direction Update indicator. 1 means take a forward step, -1 means take a backward step.
- iter Number of iterations.
- EBIC EBIC for each solution.
- loss loss for each solution.
- df Number of nonzero coefficients.
- opt Position of the optimal tuning based on EBIC.
- intercept The intercept term, which appearance is due to standardization.

#### See Also

```
predict.HierFabs
```

#### **Examples**

```
set.seed(1)
n = 500
p = 100
x = matrix(rnorm(n*p),n,p)
eta = x[,1:4] %*% rep(1,4) + 3*x[,1]*x[,2] + 3*x[,1]*x[,4]
y = eta + rnorm(n)
xtest = matrix(rnorm(n*p),n,p)
eta.test = xtest[,1:4] %*% rep(1,4) + 3*xtest[,1]*xtest[,2] + 3*xtest[,1]*xtest[,4]
ytest = eta.test + rnorm(n)
fit.gg.strong = HierFabs(x, y)
y.pred.gg.s = predict(fit.gg.strong, xtest)
mean((y.pred.gg.s-ytest)^2)
## Weak hierarchy
fit.gg.weak = HierFabs(x, y, hier="weak")
y.pred.gg.w = predict(fit.gg.strong, xtest)
mean((y.pred.gg.w-ytest)^2)
## Cox model with Gene-Environment interactions
pz = 10
z = matrix(rnorm(n*pz),n,pz)
eta.ge = x[,1] + x[,2] + z[,1] + z[,2] + 3*x[,1]*z[,1] + 3*x[,2]*z[,2]
err = log(rweibull(n, shape = 1, scale = 1))
y0 = exp(-eta.ge + err)
cens = quantile(y0, 0.9)
y.ge = pmin(y0, cens)
status = 1 * (y0 <= cens)
ztest = matrix(rnorm(n*pz),n,pz)
\verb|eta.ge.test| = xtest[,1] + xtest[,2] + ztest[,1] + ztest[,2]|
eta.ge.test = eta.ge.test + 3*xtest[,1]*ztest[,1] + 3*xtest[,2]*ztest[,2]
fit.ge.strong = HierFabs(x, y.ge, z, model="cox", status=status)
y.pred.gg.s = predict(fit.ge.strong, xtest, ztest)
mean((y.pred.gg.s-eta.ge.test)^2)
## Weak hierarchy
fit.ge.weak = HierFabs(x, y.ge, z, model="cox", status=status, hier="weak")
y.pred.ge.w = predict(fit.ge.weak, xtest, ztest)
mean((y.pred.ge.w-eta.ge.test)^2)
```

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predict.HierFabs	Model prediction based on a fitted HierFabs object.
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## Description

Similar to the usual predict methods, this function returns predictions from a fitted 'HierFabs' object.

## Usage

```
## S3 method for class 'HierFabs'
predict(object, new.G, new.E, allpath = FALSE, ...)
```

## Arguments

object	Fitted 'HierFabs' model object.
new.G	Matrix of new values for genes.
new.E	An optional new values for environment variables.
allpath	allpath = T will output all the predictions on the solution path. allpath = FALSE will only output the optimal solution selected in the 'HierFabs' object.
	Not used. Other arguments to predict.

#### Value

Predictions for each observation.

## See Also

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```