Package ‘Cascade’

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https://github.com/fbertran/Cascade/

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NeedsCompilation no
Cascade-package

The Cascade Package: Selection, Reverse-Engineering and Prediction in Cascade Networks

Description

A modeling tool allowing gene selection, reverse engineering, and prediction in cascade networks.

Author(s)

This package has been written by Frédéric Bertrand, Myriam Maumy-Bertrand and Nicolas Jung with biological insights from Laurent Vallat. Maintainer: Frédéric Bertrand <frederic.bertrand@math.unistra.fr>
References


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**analyze_network**

### Analysing the network

**Description**

Calculates some indicators for each node in the network.

**Usage**

```r
## S4 method for signature 'network'
analyze_network(Omega, nv, label_v = NULL)
```

**Arguments**

- **Omega**: a network object
- **nv**: the level of cutoff at which the analysis should be done
- **label_v**: (optional) the name of the genes

**Value**

A matrix containing, for each node, its betweenness, its degree, its output, its closeness.

**Author(s)**

Nicolas Jung, Frédéric Bertrand, Myriam Maumy-Bertrand.

**References**


Examples

```r
data(network)
analyze_network(network, nv=0)
```

**as.micro_array**  
Coerce a matrix into a micro_array object.

**Description**

Coerce a matrix into a micro_array object.

**Usage**

```r
as.micro_array(M, time, subject)
```

**Arguments**

- **M**  
  A matrix. Contains the microarray measurements. Should of size N * K, with N the number of genes and K=T*P with T the number of time points, and P the number of individuals. This matrix should be created using `cbind(M1,M2,...)` with M1 a N*T matrix with the measurements for individual 1, M2 a N*T matrix with the measurements for individual 2.

- **time**  
  A vector. The time points measurements.

- **subject**  
  The number of subjects.

**Value**

A micro_array object.

**Author(s)**

Nicolas Jung, Frédéric Bertrand, Myriam Maumy-Bertrand.

**References**


compare-methods

Examples

```r
if(require(CascadeData)){
  data(micro_US)
  micro_US<-as.micro_array(micro_US,time=c(60,90,210,390),subject=6)
}
```

Some basic criteria of comparison between actual and inferred network.

Description

Allows comparison between actual and inferred network.

Usage

```r
## S4 method for signature 'network, network, numeric'
compare(Net, Net_inf, nv = 1)
```

Arguments

- `Net` A network object containing the actual network.
- `Net_inf` A network object containing the inferred network.
- `nv` A number that indicates at which level of cutoff the comparison should be done.

Value

A vector containing: sensibility, predictive positive value, and the F-score

Methods

```r
list("signature(Net = \"network\", Net_inf = \"network\", nv = \"numeric\")")
```

Author(s)

Nicolas Jung, Frédéric Bertrand, Myriam Maumy-Bertrand.

References


Examples

data(Net)
data(Net_inf)

#Comparing true and inferred networks
F_score=NULL

#Here are the cutoff level tested
test.seq<-seq(0,max(abs(Net_inf@network*0.9)),length.out=200)
for(u in test.seq){
  F_score<-rbind(F_score,Cascade::compare(Net,Net_inf,u))
}
matplot(test.seq,F_score,type="l",ylab="criterion value",xlab="cutoff level",lwd=2)


cutoff.network-method  Choose the best cutoff

Description

Allows estimating the best cutoff, in function of the scale-freeness of the network. For a sequence of cutoff, the corresponding p-value is then calculated.

Usage

## S4 method for signature 'network'
cutoff(Omega, sequence = NULL, x_min = 0)

Arguments

Omega  a network object
sequence (optional) a vector corresponding to the sequence of cutoffs that will be tested.
x_min (optional) an integer ; only values over x_min are further retained for performing the test.

Value

A list containing two objects :

  p.value  the p values corresponding to the sequence of cutoff
  p.value.inter  the smoothed p value vector, using the loess function

Author(s)

Nicolas Jung, Frédéric Bertrand, Myriam Maumy-Bertrand.
dim

References


Examples

data(network)
cutoff(network)
#See vignette for more details

---

dim

*Dimension of the data*

Description

Dimension of the data

Usage

```r
## S4 method for signature 'micro_array'
dim(x)
```

Arguments

- `x` an object of class "micro_array"

Methods

- `list(signature(x = "micro_array"))` Gives the dimension of the matrix of measurements.

Examples

```r
if(require(CascadeData)){
data(micro_US)
micro_US<-as.micro_array(micro_US,time=c(60,90,210,390),subject=6)
dim(micro_US)
}
```
See the evolution of the network with change of cutoff

Description

See the evolution of the network with change of cutoff. This function may be useful to see if the global topology is changed while increasing the cutoff.

Usage

```r
## S4 method for signature 'network'
 evolution(
   net,
   list_nv,
   gr = NULL,
   color.vertex = NULL,
   fix = TRUE,
   gif = TRUE,
   taille = c(2000, 1000),
   label_v = 1:dim(net@network)[1],
   legend.position = "topleft",
   frame.color = "black",
   label.hub = FALSE
)
```

Arguments

**net**
- a network object

**list_nv**
- a vector of cutoff at which the network should be shown

**gr**
- a vector giving the group of each gene

**color.vertex**
- a vector giving the color of each node

**fix**
- logical, should the position of the node in the network be calculated once at the beginning? Defaults to TRUE.

**gif**
- logical, TRUE

**taille**
- vector giving the size of the plot. Default to c(2000,1000)

**label_v**
- (optional) the name of the genes

**legend.position**
- (optional) the position of the legend, defaults to "topleft"

**frame.color**
- (optional) the color of the frame, defaults to "black"

**label.hub**
- (optional) boolean, defaults to FALSE

Value

A HTML page with the evolution of the network.
Author(s)
Nicolas Jung, Frédéric Bertrand, Myriam Maumy-Bertrand.

References


Examples

```r
data(network)
sequence<-seq(0,0.2,length.out=20)
#setwd("inst/animation")
#evolution(network,sequence)
```

### geneNeighborhood.network-method

*Find the neighborhood of a set of nodes.*

Description
Find the neighborhood of a set of nodes.

Usage
```r
## S4 method for signature 'network'
geneNeighborhood(
  net, targets,
  nv = 0, order = length(net@time.pt) - 1, label_v = NULL,
  ini = NULL, frame.color = "white", label.hub = FALSE,
  graph = TRUE, names = FALSE)
```
geneNeighborhood.network-method

Arguments

- **net**: a network object
- **targets**: a vector containing the set of nodes
- **nv**: the level of cutoff. Default to 0.
- **order**: of the neighborhood. Default to `length(net@time_pt)-1`.
- **label_v**: vector defining the vertex labels.
- **ini**: using the “position” function, you can fix the position of the nodes.
- **frame.color**: color of the frames.
- **label.hub**: logical; if TRUE only the hubs are labeled.
- **graph**: plot graph of the network. Defaults to ‘TRUE’.
- **names**: return names of the neighbors. Defaults to ‘FALSE’.

Value

The neighborhood of the targeted genes.

Author(s)

Nicolas Jung, Frédéric Bertrand, Myriam Maumy-Bertrand.

References


Examples

data(Selection)
data(network)
# A nv value can be chosen using the cutoff function
nv=.11
EGR1<-which(match(Selection@name,"EGR1")==1)
P<-position(network,nv=nv)
geneNeighborhood(network,targets=EGR1,nv=nv,ini=P,
  label_v=network@name)
Description

Selection of differentially expressed genes.

Usage

## S4 method for signature 'micro_array,micro_array,numeric'
geneSelection(
  x,
  y,
  tot.number,
  data_log = TRUE,
  wanted.patterns = NULL,
  forbidden.patterns = NULL,
  peak = NULL,
  alpha = 0.05,
  Design = NULL,
  lfc = 0
)

## S4 method for signature 'list,list,numeric'
geneSelection(
  x,
  y,
  tot.number,
  data_log = TRUE,
  alpha = 0.05,
  cont = FALSE,
  lfc = 0,
  f.asso = NULL
)

## S4 method for signature 'micro_array,numeric'
genePeakSelection(
  x,
  peak,
  y = NULL,
  data_log = TRUE,
  durPeak = c(1, 1),
  abs_val = TRUE,
  alpha_diff = 0.05
)
Arguments

x

either a micro_array object or a list of micro_array objects. In the first case, the micro_array object represents the stimulated measurements. In the second case, the control unstimulated data (if present) should be the first element of the list.

y

either a micro_array object or a list of strings. In the first case, the micro_array object represents the stimulated measurements. In the second case, the list is the way to specify the contrast:

First element: condition, condition&time or pattern. The condition specification is used when the overall is to compare two conditions. The condition&time specification is used when comparing two conditions at two precise time points. The pattern specification allows to decide which time point should be differentially expressed.

Second element: a vector of length 2. The two conditions which should be compared. If a condition is used as control, it should be the first element of the vector. However, if this control is not measured through time, the option cont=TRUE should be used.

Third element: depends on the first element. It is no needed if condition has been specified. If condition&time has been specified, then this is a vector containing the time point at which the comparison should be done. If pattern has been specified, then this is a vector of 0 and 1 of length T, where T is the number of time points. The time points with desired differential expression are provided with 1.

tot.number

an integer. The number of selected genes. If tot.number < 0 all differentially genes are selected. If tot.number > 1, tot.number is the maximum of differentially genes that will be selected. If 0<tot.number<1, tot.number represents the proportion of differentially genes that are selected.

data_log

logical (default to TRUE); should data be logged ?

wanted.patterns

a matrix with wanted patterns [only for geneSelection].

forbidden.patterns

a matrix with forbidden patterns [only for geneSelection].

peak

interger. At which time points measurements should the genes be selected [optionnal for geneSelection].

alpha

float; the risk level. Default to ‘alpha=0.05’

Design

the design matrix of the experiment. Defaults to ‘NULL’.

lfc

log fold change value used in limma’s ‘topTable’. Defaults to 0.

cont

use contrasts. Defaults to ‘FALSE’.

f.asso

function used to assess the association between the genes. The default value ‘NULL’ implies the use of the usual ‘mean’ function.

durPeak

vector of size 2 (default to c(1,1)) ; the first elements gives the length of the peak at the left, the second at the right. [only for genePeakSelection]

abs_val

logical (default to TRUE); should genes be selected on the basis of their absolute value expression ? [only for genePeakSelection]

alpha_diff

float; the risk level
**Value**

A micro_array object.

**Author(s)**

Nicolas Jung, Frédéric Bertrand, Myriam Maumy-Bertrand.

**References**


**Examples**

```r
if(require(CascadeData)){
data(micro_US)
micro_US<-as.micro_array(micro_US,time=c(60,90,210,390),subject=6)
data(micro_S)
micro_S<-as.micro_array(micro_S,time=c(60,90,210,390),subject=6)

#Basically, to find the 50 more significant expressed genes you will use:
Selection_1<-geneSelection(x=micro_S,y=micro_US,
tot.number=50,data_log=TRUE)
summary(Selection_1)

#If we want to select genes that are differentially
#at time t60 or t90 :
Selection_2<-geneSelection(x=micro_S,y=micro_US,tot.number=30,
wanted.patterns= rbind(c(0,1,0,0),c(1,0,0,0),c(1,1,0,0)))
summary(Selection_2)

#To select genes that have a differential maximum of expression at a specific time point.
Selection_3<-genePeakSelection(x=micro_S,y=micro_US,peak=1,
abs_val=FALSE,alpha_diff=0.01)
summary(Selection_3)
}
```

if(require(CascadeData)){
data(micro_US)
micro_US<-as.micro_array(micro_US,time=c(60,90,210,390),subject=6)
data(micro_S)
micro_S<-as.micro_array(micro_S,time=c(60,90,210,390),subject=6)

#Genes with differential expression at t1
Selection1<-geneSelection(x=micro_S, y=micro_US, 20, wanted.patterns= rbind(c(1,0,0,0)))
#Genes with differential expression at t2
Selection2<-geneSelection(x=micro_S, y=micro_US, 20, wanted.patterns= rbind(c(0,1,0,0)))
#Genes with differential expression at t3
Selection3<-geneSelection(x=micro_S, y=micro_US, 20, wanted.patterns= rbind(c(0,0,1,0)))
#Genes with differential expression at t4
Selection4<-geneSelection(x=micro_S, y=micro_US, 20, wanted.patterns= rbind(c(0,0,0,1)))
#Genes with global differential expression
Selection5<-geneSelection(x=micro_S, y=micro_US, 20)

#We then merge these selections:
Selection<-unionMicro(list(Selection1, Selection2, Selection3, Selection4, Selection5))
print(Selection)

#Prints the correlation graphics Figure 4:
summary(Selection, 3)

## Uncomment this code to retrieve geneids.
#library(org.Hs.eg.db)
#
#ff<-function(x){substr(x, 1, nchar(x)-3)}
#ff<-Vectorize(ff)
#
## Here is the function to transform the probeset names to gene ID.
#
#library("hgu133plus2.db")
#
#probe_to_id<-function(n){
#  x <- hgu133plus2$SYMBOL
#  mp<-mappedkeys(x)
#  xx <- unlist(as.list(x[mp]))
#  genes_all = xx[(n)]
#  genes_all[is.na(genes_all)]<="unknown"
#  return(genes_all)
#
#Selection@name<-probe_to_id(Selection@name)
}
Usage

```r
## S4 method for signature 'network'
gene_expr_simulation(network, time_label = 1:4, subject = 5, level_peak = 100)
```

Arguments

- `network`: A network object.
- `time_label`: a vector containing the time labels.
- `subject`: the number of subjects
- `level_peak`: the mean level of peaks.

Value

A `micro_array` object.

Author(s)

Nicolas Jung, Frédéric Bertrand, Myriam Maumy-Bertrand.

References


Examples

```r
data(Net)
set.seed(1)

#We simulate gene expression according to the network Net
Msim<-gene_expr_simulation(
  network=Net,
  time_label=rep(1:4,each=25),
  subject=5,
  level_peak=200)
head(Msim)
```
Overview of a micro_array object

Description

Overview of a micro_array object.

Usage

```r
## S4 method for signature 'micro_array'
head(x, ...)
```

Arguments

- `x`: an object of class `micro_array`.
- `...`: additional parameters

Methods

- `list("signature(x = \"ANY\")")` Gives an overview.
- `list("signature(x = \"micro_array\")")` Gives an overview.

Examples

```r
if(require(CascadeData)){
  data(micro_US)
  micro_US<-as.micro_array(micro_US,time=c(60,90,210,390),subject=6)
  head(micro_US)
}
```

Reverse-engineer the network

Description

Reverse-engineer the network.
Usage

## S4 method for signature 'micro_array'

\texttt{inference(M, tour.max = 30, g = function(x) \{ 1/x \}, conv = 0.001, cv.subjects = TRUE, nb.folds = NULL, eps = 10^-5, type.inf = "iterative")}

Arguments

- \texttt{M} \hspace{1cm} a micro_array object.
- \texttt{tour.max} \hspace{1cm} maximal number of steps. Defaults to ‘tour.max=30’
- \texttt{g} \hspace{1cm} the new solution is choosen as (the old solution + g(x) \times the new solution)/(1+g(x)) where x is the number of steps. Defaults to ‘g=function(x) 1/x’
- \texttt{conv} \hspace{1cm} convergence criterion. Defaults to ‘conv=10e-3’
- \texttt{cv.subjects} \hspace{1cm} should the cross validation be done removing the subject one by one ? Defaults to ‘cv.subjects=TRUE’.
- \texttt{nb.folds} \hspace{1cm} Relevant only if cv.subjects is FALSE. The number of folds in cross validation. Defaults to ‘NULL’.
- \texttt{eps} \hspace{1cm} machine zero. Defaults to ‘10e-5’.
- \texttt{type.inf} \hspace{1cm} “iterative” or “noniterative” : should the algorithm be computed iteratively. Defaults to ‘”iterative”’.

Value

A network object.

Author(s)

Nicolas Jung, Frédéric Bertrand, Myriam Maumy-Bertrand.

References


Examples

```r
# With simulated data
data(M)
infe <- inference(M)
str(infe)

# With selection of genes from GSE39411
data(Selection)
infsel <- inference(Selection)
str(infsel)
```

---

**M**

*Simulated M data for examples.*

---

**Description**

Simulated M microarray.

**Examples**

```r
data(M)
head(M)
```

---

**micropredict-class**

*Class "micropredict"*

---

**Description**

The "micropredict" class

**Objects from the Class**

Objects can be created by calls of the form `new("micropredict", ...)`.  

**Examples**

```r
showClass("micropredict")
```
micro_array-class

Class "micro_array"

Description

The "micro_array" class

Objects from the Class

Objects can be created by calls of the form new("micro_array", ...).

Examples

   showClass("micro_array")

Net

Simulated network data for examples.

Description

Simulated network.

Examples

   data(Net)
   str(Net)

network

A network object data.

Description

A network object. It is the same as the result in the vignette for the inference of the network.

Examples

   data(network)
   plot(network)
   print(network)
network-class  

Class "network"

Description

The "network" class

Objects from the Class

Objects can be created by calls of the form `new("network", ...)`. 

Examples

```r
showClass("network")
```

network_random  

Generates a network.

Description

Generates a network.

Usage

```r
network_random(
  nb,
  time_label,
  exp,
  init,
  regul,
  min_expr,
  max_expr,
  casc.level
)
```

Arguments

- **nb**  
  Integer. The number of genes.
- **time_label**  
  Vector. The time points measurements.
- **exp**  
  The exponential parameter, as in the barabasi.game function in igraph package.
- **init**  
  The attractiveness of the vertices with no adjacent edges. See barabasi.game function.
- **regul**  
  A vector mapping each gene with its number of regulators.
Net_inf

min_expr Minimum of strength of a non-zero link
max_expr Maximum of strength of a non-zero link
casc.level ... 

Value
A network object.

Author(s)
Nicolas Jung, Frédéric Bertrand, Myriam Maumy-Bertrand.

References


Examples

set.seed(1)
Net<-network_random(
  nb=100,
  time_label=rep(1:4,each=25),
  exp=1,
  init=1,
  regul=round(rexp(100,1))+1,
  min_expr=0.1,
  max_expr=2,
  casc.level=0.4
)
plot(Net)

---

Net_inf Reverse-engineered network of the simulated data.

Description
The reverse-engineered network of the simulated data (M and Net).
Examples

```r
data(Net_inf)
str(Net_inf)
```

Description

Considering the class of the argument which is passed to `plot`, the graphical output differs.

Usage

```r
## S4 method for signature 'micro_array,ANY'
plot(x, y, ...)

## S4 method for signature 'network,ANY'
plot(
  x,
  y,
  choice = "network",
  nv = 0,
  gr = NULL,
  ini = NULL,
  color.vertex = NULL,
  video = TRUE,
  weight.node = NULL,
  ani = FALSE,
  taille = c(2000, 1000),
  label_v = 1:dim(x@network)[1],
  horiz = TRUE,
  legend.position = "topleft",
  frame.color = "black",
  label.hub = FALSE,
  ...
)

## S4 method for signature 'micropredict,ANY'
plot(
  x,
  time = NULL,
  label_v = NULL,
  frame.color = "white",
  ini = NULL,
  label.hub = FALSE,
)```
edge.arrow.size = 0.7,
edge.thickness = 1
)

Arguments

x a micro_array object, a network object or a micropredict object
y optional and not used if x is an appropriate structure
... additional parameters
choice what graphic should be plotted: either "F" (for a representation of the matrices F) or "network".
nv the level of cutoff. Default to ‘0’.
gr a vector giving the group of each gene
ini using the “position” function, you can fix the position of the nodes.
color.vertex a vector defining the color of the vertex.
video if ani is TRUE and video is TRUE, the result of the animation is saved as an animated GIF.
weight.node nodes weighting. Defaults to ‘NULL’.
ani animated plot?
taille vector giving the size of the plot. Default to ‘c(2000,1000)’.
label.v vector defining the vertex labels.
horiz landscape? Defaults to ‘TRUE’.
legend.position position of the legend.
frame.color color of the frames.
label.hub logical ; if TRUE only the hubs are labeled.
time sets the time for plot of the prediction. Defaults to ‘NULL’
edge.arrow.size size of the arrows ; default to 0.7.
edge.thickness edge thickness ; default to 1.

Methods

list("signature(x = "micro_array", y = "ANY",...)") x a micro_array object
list_nv a vector of cutoff at which the network should be shown
list("signature(x = "network", y = "ANY",...)") x a network object
list() Optionnal arguments:
gr a vector giving the group of each gene
choice what graphic should be plotted: either "F" (for a representation of the matrices F) or "network".
nv the level of cutoff. Default to 0.
ini using the “position” function, you can fix the position of the nodes
position-methods

Returns the position of edges in the network

Description

Returns the position of edges in the network

Usage

## S4 method for signature 'network'
position(net, nv = 0)

Arguments

net a network object

nv the level of cutoff at which the analysis should be done
Methods

```r
list("signature(net = \"network\")")
```

Returns a matrix with the position of the node. This matrix can then be used as an argument in the plot function.

Examples

```r
data(Net)
position(Net)
```

DESCRIPTION

Prediction of the gene expressions after a knock-out experience

Usage

```r
## S4 method for signature 'micro_array'
predict(object, Omega, nv = 0, targets = NULL, adapt = TRUE)
```

Arguments

- `object`: a micro_array object
- `Omega`: a network object.
- `nv`: [\(=0\)] numeric; the level of the cutoff
- `targets`: [NULL] vector; which genes are knocked out?
- `adapt`: [TRUE] boolean; do not raise an error if used with vectors instead of one column matrices.

Author(s)

Nicolas Jung, Frédéric Bertrand, Myriam Maumy-Bertrand.

References


Examples

```r
data(Selection)
data(network)
# A nv value can be chosen using the cutoff function
nv = .11
EGR1 <- which(match(Selection@name, "EGR1") == 1)
P <- position(network, nv = nv)

# We predict gene expression modulations within the network if EGR1 is experimentally knocked-out.
prediction_ko5 <- predict(Selection, network, nv = nv, targets = EGR1)

# Then we plot the results. Here for example we see changes at time point t2:
plot(prediction_ko5, time = 2, ini = P, label_v = Selection@name)
```

---

**print-methods**

*Methods for Function print*

**Description**

Methods for function `print`

**Usage**

```r
## S4 method for signature 'micro_array'
print(x, ...)  

## S4 method for signature 'network'
print(x, ...)
```

**Arguments**

- `x` 
  - an object of class micro-array or network
- `...` 
  - additional parameters

**Examples**

```r
data(Net)
print(Net)

data(M)
print(M)
```
Selection

Selection of genes.

Description

20 (at most) genes with differential expression at t1, 20 (at most) genes with differential expression at t2, 20 (at most) genes with differential expression at t3, 20 (at most) genes with differential expression at t4 et 20 (at most) genes with global differential expression were selected.

Examples

data(Selection)
head(Selection)
summary(Selection,3)

Summary-methods

Methods for Function summary

Description

Methods for function summary

Usage

## S4 method for signature 'micro_array'
summary(object, nb.graph = NULL, ...)

Arguments

  object             an object of class micro-array
  nb.graph          (optionnal) choose the graph to plot. Displays all graphs by default.
  ...               additional parameters.

Examples

data(M)
summary(M)
unionMicro-methods

Makes the union between two micro_array objects.

Description

Makes the union between two micro_array objects.

Usage

```r
## S4 method for signature 'micro_array,micro_array'
unionMicro(M1, M2)
```

Arguments

- `M1`: a micro-array or a list of micro-arrays
- `M2`: a micro-array or nothing if `M1` is a list of micro-arrays

Methods

- `list("signature(M1 = "micro_array", M2 = "micro_array")")`: Returns a micro_array object which is the union of `M1` and `M2`.
- `list("signature(M1 = "list", M2 = "ANY")")`: Returns a micro_array object which is the union of the elements of `M1`.

Examples

```r
data(M)
# Create another microarray object with 100 genes
Mbis <- M
# Rename the 100 genes
Mbis@name <- paste(M@name, "bis")
rownames(Mbis@microarray) <- Mbis@name
# Union (merge without duplicated names) of the two microarrays.
str(unionMicro(M, Mbis))
```
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