# Package 'CNORfeeder'

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Type Package

Title Integration of CellNOptR to add missing links

**Version** 1.49.0

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**Description** This package integrates literature-constrained and data-driven methods to infer signalling networks from perturbation experiments. It permits to extends a given network with links derived from the data via various inference methods and uses information on physical interactions of proteins to guide and validate the integration of links.

License GPL-3

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## Description

CNORfeeder permits to extend a network derived from literature with links derived strictly from the data via various inference methods using information on physical interactions of proteins to guide and validate the integration of links. The package is designed to be integrated with CellNOptR.

## **Details**

Package: CNORfeeder
Type: Package
Version: 1.0.0.
Date: 2012-11-22
License: GPLv2
LazyLoad: yes

## Author(s)

F. Eduati Maintainer: F. Eduati <eduati@ebi.ac.uk>

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#### References

F. Eduati, J. De Las Rivas, B. Di Camillo, G. Toffolo, J. Saez-Rodriguez. Integrating literature-constrained and data-driven inference of signalling networks. Bioinformatics, 28(18):2311-2317, 2012.

## **Examples**

```
library(CNORfeeder)
# this is an example of the main steps of the integrated CellNOptR - CNORfeeder pipeline
# load the data already formatted as CNOlist
data(CNOlistDREAM,package="CellNOptR")
# load the model (PKN) already in the CNO format
data(DreamModel,package="CellNOptR")
# see CellNOptR documentation to import other data/PKNs)
# A. INFERENCE - CNORfeeder
# FEED inference: codified in Boolean Tables
BTable <- makeBTables(CNOlist=CNOlistDREAM, k=2, measErr=c(0.1, 0))
# B. COMPRESSION - CellNOptR
# preprocessing step
model<-CellNOptR::preprocessing(data=CNOlistDREAM, model=DreamModel)</pre>
# C. INTEGRATION - CNORfeeder
# integration with the compressed model
modelIntegr <- mapBTables2model(BTable=BTable,model=model,allInter=TRUE)</pre>
\mbox{\tt\#} see example in ?MapDDN2Model to use other reverse-engineering methods
# D. WEGHTING - CNORfeeder
# integrated links are weighted more according to the integratin factor integrFac
\verb|modelIntegrWeight| <- \verb|weighting(modelIntegr=modelIntegr)| PKNmodel=DreamModel, \\
                               CNOlist=CNOlistDREAM, integrFac=10)
# E. TRAINING - CellNOptR
# initBstring<-rep(1,length(modelIntegr$reacID))</pre>
# training to data using genetic algorithm (run longer to obtain better results)
DreamT1opt<-gaBinaryT1W(</pre>
CNOlist=CNOlistDREAM,
model=modelIntegrWeight,
maxGens=2,
 popSize=5,
 verbose=FALSE)
```

Binference

Bayesian network inference

## Description

This function uses data (CNOlist) to infer a Bayesian network using the catnet package.

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## Usage

## **Arguments**

CN0list a CN0list structure, as produced by makeCN0list

mode a character, optimization network selection criterion such as "AIC" and "BIC",

to be used in cnSearchSA

tempCheckOrders

an integer, the number of iteration, orders to be searched, with constant temper-

ature, to be used in cnSearchSA

maxIter an integer, the total number of iterations, thus orders, to be processed, to be used

in cnSearchSA

filename name of the sif file saved, default BAYESIAN

#### **Details**

This function transforms the data in a format compatible with catnet package, infers the network using the Stochastic Network Search as implemented in catnet (see cnSearchSA), computes the consensus model of the models returned by cnSearchSA considering only links that have a frequency of appearence greater than 0.1 and returns the model in the sif format.

## Value

sif the inferred data-driven network in sif format

## Author(s)

F.Eduati

#### See Also

mapDDN2model

## **Examples**

buildFeederObjectDynamic

Building Feeder-Object for the integration to the PKN

## Description

This function estimates the possible mechanisms of interactions to be added to the PKN from a database of interactions for improving the fitting cost.

## Usage

buildFeederObjectDynamic(model = model, cnolist = cnolist, indices = indices, database = NULL, DDN =

## **Arguments**

model	a model as returned by readSIF. Alternatively, the filename can also be provided
cnolist	a cnolist structure, as produced by makeCNOlist
indices	a list of indices of poorly fitted measurements as returned from $identifyMisfitIndices$
database	a database of interactions which can be optionally provided as an interaction matrix with 3 or 4 colums (source of interaction, sign of interaction, target of interaction and optionally a weight value from 0 to 1 indicating the significance of that interaction in the database). Default: database=NULL
DDN	a parameter indicating whether integrating links inferred from the Data-Driven FEED approach. Default: $DDN = TRUE$
pathLength	a path length parameter for the maximal path length of additional interactions to search for in the database. Default: $pathLength = 2$
k	a parameter that determine the threshold of significancy of the effect of stimuli and inhibitors, default to $2$
measErr	a 2 value vector (err1, err2) defining the error model of the data as $sd^2 = err1^2 + (err2*data)^2$ , default to $c(0.1, 0)$
timePoint	time-points to be considered. By default set to NA, which means that the function will search for poorly fitted measurements at each time-point.

## **Details**

The function identifies and proposes the new links to integrate in the PKN either either by means of the data-driven method from the FEED algorithm or from the provided database of interactions or from both of them.

## Value

this function returns a list with fields:

Original PKN the original PKN

Feed mechanisms

the list of proposed interactions to integrate to the PKN (if both the database and the data-driven method are considered by the user, the last mechanism corresponds to the data-driven approach)

6 database

## Author(s)

E.Gjerga

## **Examples**

cnolist

**CNOlist** 

## **Description**

CNOlist object containing the perturbation experimental data.

## Usage

cnolist

## **Format**

cnolist is the object which contains the data in the MIDAS file.

### Source

This object is generated from the dynamic-feeder example

database

OmniPath PPI

## Description

Data-frame containing signed and directed interactions from Omnipath.

## Usage

database

### **Format**

database is the object which contains new interactions which can potentially be integrated.

feederObject 7

### **Source**

This object is generated from the dynamic-feeder example

Feeder Object
---------------

## **Description**

Object list as obtained from the buildFeederObjectDynamic() function.

### Usage

feederObject

### **Format**

feederObject is a list containing interactions suggested to be added in the PKN.

### **Source**

This object is generated from the dynamic-feeder example

gaBinaryT1W	Genetic algorithm used to optimise a model differently weighting links

## Description

This function is the genetic algorithm to be used to optimise a model by fitting to data containing one time point. It is the function gaBinaryT1 of CellNOptR modified in orter to differently weights for the integrated links

## Usage

```
gaBinaryT1W(CNOlist, model, initBstring=NULL, sizeFac = 1e-04,
   NAFac = 1, popSize = 50, pMutation = 0.5, maxTime = 60, maxGens = 500,
   stallGenMax = 100, selPress = 1.2, elitism = 5, relTol = 0.1, verbose=TRUE,
   priorBitString=NULL, maxSizeHashTable=5000)
```

## **Arguments**

CNOlist	a CNOlist on which the score is based (based on valueSignals[[2]], i.e. data at time 1)
model	a model structure, as created by readSIF, normally pre-processed but that is not a requirement of this function. If the linksWeight field is provided in model structure, all links are weighted according to that.
initBstring	an initial bitstring to be tested, should be of the same size as the number of reactions in the model above (model\$reacID). Default is all ones.
sizeFac	the scaling factor for the size term in the objective function, default to 0.0001

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NAFac the scaling factor for the NA term in the objective function, default to 1

popSize the population size for the genetic algorithm, default set to 50

pMutation the mutation probability for the genetic algorithm, default set to 0.5

maxTime the maximum optimisation time in seconds, default set to 60

maxGens the maximum number of generations in the genetic algorithm, default set to 500

stallGenMax the maximum number of stall generations in the genetic algorithm, default to

100

selPress the selective pressure in the genetic algorithm, default set to 1.2

elitism the number of best individuals that are propagated to the next generation in the

genetic algorithm, default set to 5

relTol the relative tolerance for the best bitstring reported by the genetic algorithm, i.e.,

how different from the best solution, default set to 0.1

verbose logical (default to TRUE) do you want the statistics of each generation to be

printed on the screen?

priorBitString At each generation, the GA algorithm creates a population of bitstrings that

will be used to perform the optimisation. If the user knows the values of some bits, they can be used to overwrite bit values proposed by the GA algorithm. If provided, the priorBitString must have the same length as the initial bitstring and be made of 0, 1 or NA (by default, this bitstring is set to NULL, which is equivalent to setting all bits to NA). Bits that are set to 0 or 1 are used to replace

the bits created by the GA itself (see example).

maxSizeHashTable

a hash table is use to store bitstring and related score. This allows the GA to be very efficient is the case of small models. The size of the hash table is 5000 by

default, which may be too large for large models.

### **Details**

The whole procedure is described in details in Saez-Rodriguez et al. (2009). The basic principle is that at each generation, the algorithm evaluates a population of models based on excluding or including some gates in the initial pre-processed model (this is encoded in a bitstring with contains 0/1 entries for each gate). The population is then evolved based on the results of the evaluation of these networks, where the evaluation is obtained by simulating the model (to steady state) under the various conditions present in the data, and then computing the squared deviation from the data, to which a penalty is added for size of the model and for species in the model that do not reach steady state.

#### Value

This function returns a list with elements:

bString the best bitstring

results a matrix with columns "Generation", "Best\_score", "Best\_bitString", "Stall\_Generation",

"Avg\_Score\_Gen", "Best\_score\_Gen", "Best\_bit\_Gen", "Iter\_time"

stringsTol the bitstrings whose scores are within the tolerance

stringsTolScores

the scores of the above-mentioned strings

identifyMisfitIndices 9

#### Author(s)

```
C. Terfve, T. Cokelaer, F.Eduati
```

#### References

J. Saez-Rodriguez, L. G. Alexopoulos, J. Epperlein, R. Samaga, D. A. Lauffenburger, S. Klamt and P. K. Sorger. Discrete logic modeling as a means to link protein signaling networks with functional analysis of mammalian signal transduction, Molecular Systems Biology, 5:331, 2009.

## See Also

```
gaBinaryT1
```

## **Examples**

```
data(CNOlistDREAM,package="CellNOptR")
data(DreamModel,package="CellNOptR")
model<-CellNOptR::preprocessing(data=CNOlistDREAM, model=DreamModel)</pre>
BTable <- makeBTables(CNOlist=CNOlistDREAM, k=2, measErr=c(0.1, 0))
modelIntegr <- mapBTables2model(BTable=BTable,model=model,allInter=TRUE)</pre>
modelIntegrWeight <- weighting(modelIntegr=modelIntegr, PKNmodel=DreamModel,</pre>
                         CNOlist=CNOlistDREAM, integrFac=10)
initBstring<-rep(1,length(modelIntegr$reacID))</pre>
# training to data using genetic algorithm (run longer to obtain better results)
DreamT1opt<-gaBinaryT1W(</pre>
CNOlist=CNOlistDREAM,
model=modelIntegrWeight,
 initBstring=initBstring,
 maxGens=2,
 popSize=5,
 verbose=FALSE)
```

## **Description**

This function identifies poorly fitted measurements for specific experimental conditions. It returns a list of possible indices and mse's pointing to possible connections to be added during the feeding process.

## Usage

```
identifyMisfitIndices(cnolist = cnolist, model = model, simData = NULL, mseThresh = 0)
```

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## **Arguments**

mseThresh

cnolist a cnolist structure, as produced by makeCNOlist

model as returned by readSIF. Alternatively, the filename can also be provided.

simData a matrix of simulated data values for a specific model as returned by plotLBod-

eFitness (default set to NULL in which case users do not need to do an initial fit of the model and the FEED algorithm will search for new links indiscriminately)

thrreshold parameter for minimal misfit to be considered - if the initial fit (mse)

for a node in a specific condition is larger/wrose than the threshold value, it will

be considered as poorly fitted (mseThresh = 0 by default)

#### Details

This function computes the misfits (MSE values) between the actual measured data points and the data values for a specific set of inferred model parameters. Once the MSE values are calculated for each of the measurements over each experimental condition, the poorly fitted measurements are then identify. A measurement is considered as poorly fitted if the corresponding inferred MSE value is higher than the specified MSE threshold value (mseThresh).

#### Value

this function returns a list with fields:

indices a list of indices pointing to the poorly fitted measurements and the corresponding

ms value

use a matrix of use values indicating the mismatch between model simulations and

data for each measurement at each experimental condition

## Author(s)

E.Gjerga

### **Examples**

indices Mis-fit indices

## Description

Simulation data as obtained from the identifyMisfitIndices() function.

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#### Usage

indices

#### **Format**

indices is a list of poorly predicted measurements.

#### **Source**

This object is generated from the dynamic-feeder example

integratedModel

Integrated Model

## **Description**

PKNlist object as obtained from the integrateLinks() function.

## Usage

integratedModel

#### **Format**

integratedModel is the model we obtain after the integration of the new links.

#### **Source**

This object is generated from the dynamic-feeder example

integrateLinks

Integrating the new links to the PKN

## **Description**

This function integrates the new links inferred via the FEED method or from the database to the original PKN.

## Usage

```
integrateLinks(feederObject = feederObject, cnolist = cnolist, database = NULL)
```

## **Arguments**

feederObject a feederObject structure, as produced by buildFeederObjectDynamic

cnolist a cnolist structure, as produced by makeCNOlist

database a database of interactions which can be optionally provided as an interaction

matrix with 3 or 4 colums (source of interaction, sign of interaction, target of interaction and optionally a weight value from 0 to 1 indicating the significance

of that interaction in the database). Default: database=NULL

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#### **Details**

This function integrates the new links inferred via the FEED method or from the database to the original PKN. Moreover it indicates which are the integrated links and if a weighted database has been used it also shows the weights assigned to each integrated link. Links that are present in the original PKN are assigned a database weight of 0, integrated links that have been inferred via the FEED method and are not present in the database are assigned a database penalty of Inf, while integrated links present in the database take values between 0 and 1.

#### Value

this function returns a list with fields:

model the integrated model

 $integ Links Idx \quad indices \ pointing \ towards \ the \ newly \ integrated \ links \ of \ the \ model$ 

integSpeciesIdx

indices pointing towards the newly integrated species of the model

databaseWeight weights assigned based on the presence of links in the database

#### Author(s)

E.Gjerga

## **Examples**

linksRanking

Ranking of links inferred from data

## **Description**

This function uses data (CNOlist) to rank links based on measurement error model as used by FEED method to reverse-engineer the network.

## Usage

```
linksRanking(CNOlist, measErr=c(0.1, 0), savefile=FALSE)
```

## Arguments

CNOlist a CNOlist structure, as produced by makeCNOlist

measErr a 2 value vector (err1, err2) defining the error model of the data as  $sd^2 = err1^2$ 

+  $(err2*data)^2$ , default to c(0.1, 0)

savefile TRUE to save the file in txt format, FALSE not. Default is FALSE.

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#### **Details**

This function is similar to the fist step of FEED to reverse engineer the network strictly from data, i.e. the inference of Boolean tables, as described in (Eduati et al., PLoS ONE, 2010) and implemented in makeBTables. Links are ranked according to the upper limit value of parameterk allowing the presence of the link, where k is the parameter which is multiplied by the measurement error in order to assess the relevance of a link. The function return link in decreasing order of importance and associate to each link a value (maximum value of k allowing the presence of the link) quantifying its relevance.

#### Value

this function returns a list with fields:

Irank

a matrix in which each link is associated with a numerical value, links are ordered in decreasing order of reliability)

#### Author(s)

F.Eduati

#### References

F. Eduati, A. Corradin, B. Di Camillo, G. Toffolo. A Boolean approach to linear prediction for signaling network modeling. PLoS ONE; 5(9): e12789.

### See Also

makeCNOlist, makeBTables

## **Examples**

```
data(CNOlistDREAM,package="CellNOptR")
Lrank <- linksRanking(CNOlist=CNOlistDREAM, measErr=c(0.1, 0))</pre>
```

makeBTables

Make Boolean tables

## **Description**

This function uses data (CNOlist) to infer a Boolean table for each measured protein, codifying if a particular stimulus inhibitor combination affects the protein. A stimulus or an inhibitor significantly affects an output protein if it is able to modify its activity level of a quantity that exceeds the uncertainty associated with its measurement.

## Usage

```
makeBTables(CNOlist, k=2, measErr=c(0.1, 0), timePoint=NA)
```

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## **Arguments**

CN01ist a CN01ist structure, as produced by makeCN01ist

k a parameter that determine the threshold of significancy of the effect of stimuli

and inhibitors, default to 2

measErr a 2 value vector (err1, err2) defining the error model of the data as sd^2 = err1^2

+  $(err2*data)^2$ , default to c(0.1, 0)

timePoint the time point to be considered for the inference of the Boolean tables (i.e. "t1"

or "t2"), if not specified all time points are consideres

#### Details

This function computes the fist step of FEED to reverse engineer the network strictly from data, i.e. the inference of Boolean tables, as described in (Eduati et al., PLoS ONE, 2010). For each protein, a Boolean table is inferred having one columns for each stimulus and one row for each inhibitor. If a stimulus produces a significant effect on the activity level of the protein this is codified with a 1 in the corresponding column, if also the inhibitor affects the protein there is a 2 in the corresponding cell. The sign of the regulation is coded in separate tables.

#### Value

this function returns a list with fields:

namesSignals a vector of names of signals

tables a list with one Boolean table for each protein codifying the effect of stimuli

(columns) and inhibitors (rows), 1 if the stimulus affect the protein, 2 if also the

inhibior does

NotMatStim has the same format as tables but just contains a 1 if the regulation has a negative

effect, and 0 otherwise

NotMatInhib has the same format as tables but just contains a 1 if the regulation has a negative

effect, and 0 otherwise

## Author(s)

F.Eduati

#### References

F. Eduati, A. Corradin, B. Di Camillo, G. Toffolo. A Boolean approach to linear prediction for signaling network modeling. PLoS ONE; 5(9): e12789.

## See Also

makeCNOlist, mapBTables2model

#### **Examples**

```
data(CNOlistDREAM,package="CellNOptR")
BTable <- makeBTables(CNOlist=CNOlistDREAM, k=2, measErr=c(0.1, 0))</pre>
```

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|--|

## **Description**

This function infers the network from the Boolean tables and integrates it with the network encoded in the model (generally derived from prior knowledge), adding links that are missing.

## Usage

mapBTables2model(BTable,model,optimRes=NA,allInter=TRUE,compressed=TRUE)

### **Arguments**

BTable a BTable list, as created by makeBTables
model a model list, as created by readSif

optimRes a bit string with the reaction of the model to be considered, default considers all

reactions

allInter one new link in the network can correspond to more links in the model, set it to

TRUE if you want to add all possible links, FALSE to add only one link, default

is TRUE

compressed this argument is used to decede how to deal with unmeasured and unperturbed

nodes (white nodes). As general guideline, it should be set to TRUE if the PKN has been compressed in the preprocessing step, FALSE otherwise. Default is

TRUE.

### **Details**

The function receive as input the Boolean Tables, infers the data-driven network form them (as descibed in (Eduati et al., PLoS ONE, 2010)) and integrates it with the model, returning a new model with the integrated links. If the Model is not given as input (Model=NULL), the data-driven network is returned as model.

## Value

a new model with the integrated links and an additional field:

indexIntegr a vector with the indexes of the integrated links

## Author(s)

F.Eduati

## References

F. Eduati, A. Corradin, B. Di Camillo, G. Toffolo. A Boolean approach to linear prediction for signaling network modeling. PLoS ONE; 5(9): e12789.

### See Also

readSif, readMIDAS, makeBTables

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#### **Examples**

```
data(CNOlistDREAM,package="CellNOptR")
data(DreamModel,package="CellNOptR")
model<-CellNOptR::preprocessing(data=CNOlistDREAM, model=DreamModel)
BTable <- makeBTables(CNOlist=CNOlistDREAM, k=2, measErr=c(0.1, 0))
modelIntegr <- mapBTables2model(BTable=BTable,model=model,allInter=TRUE)
# modelIntegr$reacID[modelIntegr$indexIntegr] to see the integrated links</pre>
```

mapDDN2model

Integrate data-drive network with the model

## **Description**

This function integrates the data-driven network (in sif format) with the network encoded in the model (generally derived from prior knowledge), adding links that are missing.

### Usage

```
mapDDN2model(DDN,model,CN0list,allInter=TRUE)
```

## **Arguments**

DDN a sif file encoding a data-driven network, as created by Binference or MIinfer-

ence

model a model list, as created by readSif

CNOlist a CNOlist, as created by makeCNOlist

allInter one new link in the network can correspond to more links in the model, set it to

TRUE if you want to add all possible links, FALSE to add only one link, default

is TRUE

## **Details**

The function receives as input a sif file with the data-driven network, as created by Binference or MIinference, and integrates it with the model, returning a new model with the integrated links.

### Value

a new Model with the integrated links and an additional field:

indexIntegr a vector with the indexes of the integrated links

## Author(s)

F.Eduati

## See Also

```
readSif, readMIDAS, Binference, MIinference
```

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#### **Examples**

MIinference

Mutual information based network inference

## Description

This function uses data (CNOlist) to infer a data-driven network using the mutual information based appoaches ARACNe and CLR as implemented in the minet package.

## Usage

## **Arguments**

cN0list a CN0list structure, as produced by makeCN0list

method a character, the name of the method to be used: ARACNE or CLR. Default,

ARACNE

PKNgraph a network to be used for comparison to assess the directionality of some links.

Default is NULL.

filename name of the sif file saved, default ARACNE

## **Details**

This function transforms the data in a format compatible with minet package, infers the network using aracne or clr as implemented in the minet package and returns the network in the sif format. It is important to notice that mutual information approaches do not allow for determining the directionality of the links thus both directions are considered. The function allows to give as input a network in graph format (graph package, see sif2graph to convert from sif to graph format) to be used as comparison to assess the directionality of some links, e.g. PKN.

## Value

sif the inferred data-driven network in sif format

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## Author(s)

F.Eduati

### References

P. E. Meyer, F. Lafitte and G. Bontempi (2008). MINET: An open source R/Bioconductor Package for Mutual Information based Network Inference. BMC Bioinformatics, 9(1), 2008

### See Also

```
mapDDN2model, sif2graph, model2sif
```

## **Examples**

model

Prior Knowledge Network

## **Description**

Model object from the dynamic-feeder example.

## Usage

model

#### **Format**

model is an PKNlist with proteins as nodes and undirected links as physical protein interactions.

## **Source**

This object is generated from the dynamic-feeder example

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**PPINigraph** 

Protein-protein interaction netwrok

### **Description**

The human protein-protein interaction network was built using a unified PPI dataset obtained as APID (Prieto, C. and De Las Rivas, J. 2006), by the combination of interactions coming from six source databases. The starting whole dataset was composed by 68488 human physical protein-protein interactions validated at least by one experimental method and reported in one article published in PubMed. From this dataset we obtained two PPI subsets with increasing confidence: a set of 28971 interactions validated by at least one binary experimental method (binary as defined in (De Las Rivas, J. and Fontanillo, C. 2010)); a set 6033 interactions validated by at least two experimental methods, one of them binary.

## Usage

**PPINigraph** 

#### **Format**

PPINigraph is an igraph with proteins as nodes and undirected links as physical protein interactions.

#### Source

This network was bult for the analysis performed in (Eduati,F. et al. 2012)

## References

- 1. F. Eduati, J. De Las Rivas, B. Di Camillo, G. Toffolo, J. Saez-Rodriguez. Integrating literature-constrained and data-driven inference of signalling networks. Bioinformatics, 28(18):2311-2317, 2012.
- 2. C. Prieto, J. De Las Rivas. APID: Agile Protein Interaction DataAnalyzer. Nucleic Acids Res., 34, W298-302, 2006.
- 3. J. De Las Rivas, C. Fontanillo. Protein-protein interactions essentials: key concepts to building and analyzing interactome networks. PLoS Comput.Biol., 6, e1000807, 2010.

runDynamicFeeder

Modelling the integrated PKN with CNORode

## **Description**

This function evaluates the effects of possible feeder mechanisms which are added to the PKN.

## Usage

runDynamicFeeder(cnolist = cnolist, integratedModel = integratedModel, ode\_parameters = ode\_parame

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#### **Arguments**

cnolist a cnolist structure, as produced by makeCNOlist integratedModel the integrated model as returned from integrateLinks ode\_parameters a list with the ODEs parameter information.

penFactor\_k a penalty factor for the new integrated links obtained from the FEED algorithm and which are not present in the database (if the database was given). Default: penFactor\_tau a penalty factor for all the new nodes integrated in the PKN. Default: penFactor\_tau = 1

penFactorPIN\_k a penalty factor for the new integrated links and which are present in the database (for the cases when the database was given). Default: penFactorPIN\_k = 10

paramsSSm a list of SSm parameters. default is the list returned by defaultParametersSSm

#### **Details**

This function evaluates the effects of possible feeder mechanisms which are added to the PKN. The analysis performed is a simple CNORode analysis over the integrated network where the newly integrated links are supposed to be penalised more than the links present in the original PKN. If a database of interactions is also provided by the user, than normally the links inferred from the FEED mechanism and which re not present in the database should be more penalised than the ones that are. There is also the opportunity to weight database interactions based on their relevance (i.e. number of resources, etc.).

#### Value

this function returns a list with fields:

Parameters the inferred optimal ODE parameters

Integrated-Model

the integrated model which was optimised

CNOlist the CNOlist object containing the data

## Author(s)

E.Gjerga

## **Examples**

```
data(integratedModel_toy, package="CNORfeeder")
data(CNOlistToy_Gene, package="CNORfeeder")
data(simData_toy,package="CNORfeeder")

## To be run with the recent version of the CNORode package:
## https://github.com/saezlab/CNORode
#
# library(CNORode)
#
# paramsSSm=defaultParametersSSm()
#
# ode_parameters=createLBodeContPars(integratedModel$model, LB_n = 1, LB_k = 0,
```

simData 21

```
# LB_tau = 0, UB_n = 3, UB_k = 1, UB_tau = 1, default_n = 3,
# default_k = 0.5, default_tau = 0.01, opt_n = FALSE, opt_k = TRUE,
# opt_tau = TRUE, random = TRUE)
# result = runDynamicFeeder(cnolist = CNOlistToy_Gene, integratedModel = integratedModel,
# ode_parameters = ode_parameters, paramsSSm = paramsSSm,
# penFactor_k = 2, penFactorPIN_k = 0.1, penFactor_tau = 1)
#
```

simData

CNORode simuation data

## **Description**

Simulation data as obtained from the plotLBodeFitness() function.

### Usage

simData

### **Format**

simData is a list containing simulated values for a specific set of ode parameters.

### Source

This object is generated from the dynamic-feeder example

UniprotIDdream

Uniprot identifiers for proteins in DreamModel

## **Description**

This data object contains the Uniprot identifiers corresponding to DreamModel of CellNOptR package, in order to associat them with the corresponding nodes in the protein-protein interaction network (PPINigraph).

### Usage

UniprotIDdream

### **Format**

UniprotIDdream is a list where each element is a protien of the DreamModel and is associated with the respective Uniprot identifiers.

### **Source**

This data object is manually derived from the Uniprot database.

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#### References

1. F. Eduati, J. De Las Rivas, B. Di Camillo, G. Toffolo, J. Saez-Rodriguez. Integrating literature-constrained and data-driven inference of signalling networks. Bioinformatics, 28(18):2311-2317, 2012.

2. J. Saez-Rodriguez, L. G. Alexopoulos, J. Epperlein, R. Samaga, D. A. Lauffenburger, S. Klamt and P. K. Sorger. Discrete logic modeling as a means to link protein signaling networks with functional analysis of mammalian signal transduction, Molecular Systems Biology, 5:331, 2009.

weighting

Weight integrated links.

#### **Description**

This function weights links integrated in the model using additional penalty and/or information from protien-protein interactions networks (PINs).

#### Usage

weighting(modelIntegr,PKNmodel,CNOlist,integrFac,UniprotID,PPI)

### **Arguments**

modelIntegr the integrated model as created by mapDDN2model or mapBTables2model

PKNmodel the model of the original prior-knowledge network

cNolist a CNolisi, as created by makeCNolist

integrFac a number indicating the penalty for integrated links
UniprotID a list with the Uniprot identifiers of proteins in the PKN

PPI an igraph of the PIN to be used, if no network is provided (=NULL) this infor-

mation is not used. Default is NULL.

#### **Details**

Integrated links are less reliable than links from the PKN, thus should be penalized in the optimization process. This function allows to include a panalty for integrated links (integrFact). Furthermore links can be differently prioritized based on information derived from pritein interaction networks (PIN): the basic idea is that if, for a directed link A -> B integrated in the PKN, there is a corresponding path in the PIN, it is more plausible that there is a molecular pathway A -> B. Because shorter paths are more feasible, as a first approximation the shortest path length between A and B in the PIN can be used as a reliability score for the integrated link. Since the optimization is performed on a compressed version of the PKN, one link integrated in the compressed network generally corresponds to multiple possible links integrated in the PKN and the shortes path of all. The weight for each integrated link in the compressed network is thus computed as (1 + the inverse of the sum of the inverse of the corresponding PKN of the shortest paths in the PIN). A high quality network of known human physical protein-protein interaction assembled from multiple databases is provided with the package: interactions were included only if validated by at least one binary experimental method in a published paper and the number of experimental evidences was reported for each interaction.

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### Value

modelIntegr the input modelIntegr with an additional field: a vector with the weights of the integrated links

### Author(s)

F.Eduati

### See Also

mapDDN2model, mapBTables2model, gaBinaryT1W

## **Examples**

```
data(CNOlistDREAM,package="CellNOptR")
data(DreamModel,package="CellNOptR")
data(UniprotIDdream,package="CNORfeeder")
model<-CellNOptR::preprocessing(data=CNOlistDREAM, model=DreamModel)</pre>
BTable <- makeBTables(CNOlist=CNOlistDREAM, k=2, measErr=c(0.1, 0))
modelIntegr <- mapBTables2model(BTable=BTable,model=model,allInter=TRUE)</pre>
modelIntegrWeight <- weighting(modelIntegr=modelIntegr, PKNmodel=DreamModel,</pre>
      CNOlist=CNOlistDREAM, integrFac=10)
# weighting using PPI might take some minutes
## Not run:
data(UniprotIDdream,package="CNORfeeder")
data(PPINigraph,package="CNORfeeder")
modelIntegrWeight2 <- weighting(modelIntegr=modelIntegr, PKNmodel=DreamModel,</pre>
       CNOlist=CNOlistDREAM, integrFac=10, UniprotID=UniprotIDdream,
       PPI=PPINigraph)
## End(Not run)
```

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