asContinuous

converts a boolean regulatory model to a continuous model

# Description

Take a net object of class "BooleanNetwork" as that defined by loadNetwork() and return a continuous regulatory network model in the form of a net object of the class "SQUAD". The continuous network can be used as an object function argument for the deSolve R package.

#### Usage

```
asContinuous(net)
```

#### Example

```
> data(cellCycle)
# setting initial state
> state<-generateState(cellCycle,specs=c("CycA"=1,"CycD"=1))
# setting intervals for numerical integration
> times <- seq(min=0,max=1,seq.out=100)

> cellCycle.s <- asContinuous(cellCycle)
> simulations <- ode(y=state, fun=cellCycle.s$fun, times=times, parameters = "default")
> # note that cellCycle$fun can be given as the function
> # parameter to ode solver method
> simulations
```

#### **Details**

The continuous network object retrieved by the function represent a system

of ordinary equations according to the methodology described in Martínez-Sosa, 2013 [1]. This methodology have the same fixed point attractors of that in the Boolean model but have the advantage that it takes into account kinetic parameters.

graphToModel

transforms a connectivity Matrix to a dynamic regulatory network model

### Description

Takes a connectivity matrix that represent a static regulatory network graph and transforms it to a "BooleanNetwork" or "SQUAD" object that can be used to simulate network transitions and calculating attractors.

# Usage

graphToModel(connectivityMatrix)

# Arguments

connectivityMatrix

A square matrix of the nodes of the network with -1, 0, or 1 integer values that represent negative, null or positive regulation.

# Example

- # generate a sample random connectivity matrix
- > net<-randomMatrix.e(dimensions=20)</pre>
- > plotAttractors(getAttractors(net))

#### Details

Connectivity matrices are defined as that with entries  $A_{ij}$  representing

a regulation from the node i to j. Positive (negative) regulations are represent by 1 (-1). No regulatory dependencies are indicated by zero. The model assumes that node j activation is achieved if and only if there is at least one of their positive regulators is active and all negative regulators inputs are inactive.

loadNetWorkSQUAD

loads a regulatory network given as a SQUAD format

# Description

Load a regulatory network model given as a SQUAD format. It returns an object of the class "SQUAD" that can be used to make simulations using squad().

# Usage

loadNetworkSQUAD("file.txt")

## Arguments

file	Α	text.	file	$\circ f$	the	format	"SO	HAD'	,	See	details
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for the definition of the "SQUAD" format.

fixed A list of of named numerical variables. Variables

belongs to the [0,1] interval. A valid fixed parameter is list("gen1" = 0," gen2" = 0.5, ...) where the length of the list is less that the number of nodes

in the regulatory network model.

# Example

```
# loading a "SQUAD" model from the SQUAD format
```

<sup>&</sup>gt; net <- loadNetworkSQUAD("file.txt")</pre>

<sup>&</sup>gt; # note that cellCycle\$fun can be given as the function

- > # parameter to ode solver method
- > simulations

#### Details

An example of the squad format has the following structure:

targets; factors A; A B; min(A, 1 - C)C; 1 - max(A, B)

Where the first line defines is a header which indicates that the file is in squad format. The next lines defines the names of the nodes followed by a ";". The rest of the line is a diffuse logic expression which defines the  $\omega$  parameter of the squad function, see [1]. It returns a object of the class "SQUAD" that can be used to perform simulations using the squad() function.

# Description

Finds all asynchronous stationary states of a Boolean Regulatory Network model. Takes as arguments an object of the "BoolNet" (or "SQUAD" under construction) and returns a data frame which contains the attractors of the model.

# Usage

getAttractorsAsynchronous(net)

# Arguments

net

An object of the "BoolNet" (or "SQUAD") class.

# Example

- > data(cellCycle)
- > attractors <- getAttractorsAsynchronous(cellCycle)
- > attractors

#### **Details**

The function is straightforward. It calls the ginsim library written by Nadi and colleagues, 2009 [2].

performs simulations of regulatory network models using the SQUAD methodology

squad

### Description

Performs a variety simulations of Regulatory network models using the SQUAD methodology. It can be used to set up different initial states, parameters and also to simulate gain/loss of function mutants. It returns

# Usage

squad(net,initialState,parameters,fixedGenes)

#### Arguments

net

An object of the "BoolNet" or "SQUAD" class.

# Example

```
> data(cellCycle)
> net.sq <- asContinuous(cellCycle)
> initialState <- runif(length(cellCycle$genes),0,1)
> times <- seq(0,10,by=0.5)
> h <- rep(50,length(cellCycle$genes))
> gamma <- rep(1,length(cellCycle$genes))
> parameters <- list(h,gamma)
> squad(net.sq)
```

# Details

It performs simulations according to the updated version of the Standardized Qualitative Analysis of Dynamical Systems (SQUAD) method described in Martínez-Sosa in 2013 [1], as modified from the first version in Di Cara, 2007 [2].

# Bibliography

- [1] P. Martínez-Sosa and L. Mendoza. The regulatory network that controls the differentiation of t lymphocytes. *Biosystems*, 113(2):96–103, 2013.
- [2] A. Naldi, D. Berenguier, A. Fauré, F. Lopez, D. Thieffry, and C. Chaouiya. Logical modelling of regulatory networks with ginsim 2.3. *Biosystems*, 97(2):134–139, 2009.