Package 'joda'

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Title JODA algorithm for quantifying gene deregulation using knowledge

Type Package

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Description Package 'joda' implements three steps of an algorithm called JODA. The algorithm computes gene deregulation scores. For each gene, its deregulation score reflects how strongly an effect of a certain regulator's perturbation on this gene differs between two different cell populations. The algorithm utilizes regulator knockdown expression data as well as knowledge about signaling pathways in which the regulators are involved (formalized in a simple matrix model).
Reference Szczurek E, Markowetz F, Gat-Viks I, Biecek P, Tiuryn J, Vingron M: Deregulation upon DNA damage revealed by joint analysis of context-specific perturbation data. Submitted.
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R topics documented: damage
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Knockout data for three regulators: ATM, RelA and p53 in two dif-
ferent cell populations: healthy (referred to as normal) cells and cells treated with neocarzinostatin (referred to as damaged cells).

Description

Knockout expression data for the kinase ATM and transcription factors p53 and RelA (Elkon et al., 2005), as well as knowledge about their targets and mutual regulatory relations in two different cell populations: healthy and damaged cells.

Usage

data(damage)

Format

```
data.healthy (data.frame): 8463x3, beliefs.healthy (list): 2, model.healthy (matrix): 3x3, data.damage (data.frame): 8463x3, beliefs.damage (list): 1, model.damage (matrix): 3x3.
```

Details

The data.healthy dataset contains log gene expression ratios for the regulator knockouts versus control. For the genes that are known to be targeted by RelA and genes that are targeted by p53 in normal conditions, beliefs.healthy contains certainties (beliefs) that those targets are differentially expressed upon their regulator's knockdown. The model.healthy matrix represents mutual signaling relations between the regulators in the healthy cells (here the model reflects that no regulator influences others). The data.damage dataset contains log gene expression ratios for the regulator knockouts upon treatment with neocarzinostatin versus treatment with neocarzinostatin alone. For the genes that are known to be targeted by p53 in the damaged cells, beliefs.damage contains certainties (beliefs) that they are differentially expressed upon the knockdown of p53. The model.damage matrix represents mutual signaling relations between the regulators in the damaged cells (here the model reflects ATM signaling down to RelA and p53).

Author(s)

Ewa Szczurek

References

Elkon R, Rashi-Elkeles S, Lerenthal Y, Linhart C, Tenne T, Amariglio N, Rechavi G, Shamir R, Shiloh Y. Dissection of a DNA-damage-induced transcriptional network using a combination of microarrays, RNA interference and computational promoter analysis. Genome Biol. 2005;6(5):R43.

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Examples

```
data(damage)
str(data.damage)
str(beliefs.damage)
print(model.damage)
```

deregulation.p.values Calculating deregulation p-values using resampling method.

Description

Deregulation p-values based on deregulation scores. They are calculated as fraction of permutations that give more extreme deregulation scores than for original data.

Usage

```
deregulation.p.values(data.1, beliefs.1, model.1, data.2, beliefs.2, model.2, N=100, verbose=FALSE)
```

Arguments

Details

The deregulation p-values are calculated as fraction of permutations that give more extreme deregulation scores than for original data.

Value

A list with two matrices. This p-values in the slot deregulation.p.values and with the original deregulation scores in the slot deregulationOrg.

Author(s)

Ewa Szczurek

References

http://joda.molgen.mpg.de

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See Also

```
differential.probs, regulation.scores, regulation.scores
```

Examples

Description

Deregulation scores quantify the extent to which the regulatory effect of each regulator changes between the two compared cell populations.

Usage

```
deregulation.scores(reg.scores1, reg.scores2, verbose)
```

Arguments

reg.scores1 A matrix of regulation scores of the genes (rows) for the regulators (columns),

compued with the regulation. scores function. Given for the first cell popu-

lation.

reg.scores2 The same as reg.scores1 but given for the second cell population.

verbose When TRUE, the execution prints informative messages

Details

The deregulation scores are computed by subtracting reg.scores1 from reg.scores2.

Value

A matrix with columns for the regulators, rows for the genes, and entries giving the deregulation scores.

Author(s)

Ewa Szczurek

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References

http://joda.molgen.mpg.de

See Also

```
differential.probs, regulation.scores
```

Examples

```
data(damage)
 # Step 1
 # Get the probabilities of differential expression
 # for the knockout of ATM in the healthy cells
 probs.healthy.ATM= differential.probs(data.healthy[,"ATM",FALSE], NULL)
 # Get the probabilities of differential expression
 # for the knockout of ATM in the damaged cells
 probs.damage.ATM= differential.probs(data.damage[,"ATM",FALSE], NULL)
 # Step 2
 # Regulation scores for a dataset with only one regulator
 # equal the signed probabilities
 # Step 3
 # Get the deregulation scores
 deregulation.ATM= deregulation.scores(probs.healthy.ATM, probs.damage.ATM, TRUE)
 ## Not run:
 # Step 1
 probs.healthy= differential.probs(data.healthy, beliefs.healthy)
 probs.damage= differential.probs(data.damage, beliefs.damage)
 # Step 2
 regulation.healthy= regulation.scores(probs.healthy, model.healthy)
 regulation.damage= regulation.scores(probs.damage, model.damage)
 # Step 3
 deregulation= deregulation.scores(regulation.healthy, regulation.damage, TRUE)
 ## End(Not run)
differential.probs
                         Calculating probabilities of differential expression in perturbation ex-
                         periments
```

Description

Returns probabilities of differential expression for genes under perturbation of a set of regulators. Takes as input perturbation data and beliefs about known genes.

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Usage

```
differential.probs(data, beliefs, verbose, plot.it)
```

Arguments

data A matrix of log expression ratios perrturbation vs control, for the genes (rows),

in the perturbations of the regulators (columns). The data has to have row and

colnames specified by the user.

beliefs A list with names being a subset of the regulators (i.e., the names of beliefs

have to be a subset of the columns of the data). Each list entry for a given regulator is a matrix with rows corresponding to the genes that are known to respond in some way to the perturbation of this regulator. The rownames of the matrix must be a subset of the rows in the data. The matrix can have either two or three columns. Each row is a distribution over the differential and unchanged cluster (2 columns) or over down, up-regulated and unchanged cluster of genes (3 columns). This distribution reflects the certainties with which a gene that

corresponds to this row belongs to each of those clusters.

verbose When TRUE, for each regulator and its perturbation data, the execution prints

out the parameters of the fitted model(s), indicating which components are dif-

ferential and which are unchanged.

plot.it When TRUE, for each regulator and its perturbation data, the execution plots

the Gaussian components of the fitted model(s), indicating which components

are differential and which are unchanged.

Details

For each regulator, a belief-based mixture model is fitted to the observations in the data. The fitted models have the number of model components equal to the number of columns in the corresponding beliefs. If no beliefs are given, unsupervised two-component mixture modeling is applied.

Value

A matrix with columns for the regulators, rows for the genes, and entries giving the signed probabilities of differential expression.

Author(s)

Ewa Szczurek

References

http://joda.molgen.mpg.de

See Also

regulation.scores, deregulation.scores

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Examples

```
data(damage)
# Get the probabilities of differential expression
# for the knockout of p53 in healthy cells
probs.healthy.p53= differential.probs(data.healthy[,"p53",FALSE],
beliefs.healthy["p53"], TRUE,TRUE)
# Get the probabilities of differential expression
# for the knockout of Ste12 under pheromone treatment
library(bgmm)
data(Ste12)
data=as.matrix(Ste12Data)
colnames(data)="Ste12"
beliefs=list(Ste12=Ste12Beliefs)
diff.p=differential.probs(data,beliefs,TRUE,TRUE)
## Not run:
probs.healthy= differential.probs(data.healthy, beliefs.healthy, TRUE,TRUE)
## End(Not run)
```

regulation.scores

Calculating regulation scores.

Description

Regulation scores reflect the actual regulatory influence of the regulators on the genes. For each regulator, these are probabilities of differential expression averaged over all perturbation experiments that affect the regulator.

Usage

```
regulation.scores(probs, model, verbose)
```

Arguments

probs A matrix of probabilities of differential expression of the genes (rows) under

perturbations of regulators (columns). Obtained with the differential.probs

function

model A pathway model is a matrix with rows and columns equal to the names of the

regulators (i.e., the columns of probs). Each model has an entry 1 where the regulator in the corresponding row influences the regulator in the corresponding column. A transitive closure of the input model is computed to get information

about the experiments affecting each regulator.

verbose When TRUE, the execution prints informative messages

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Value

A matrix with columns for the regulators, rows for the genes, and entries giving the regulation scores.

Author(s)

Ewa Szczurek

References

http://joda.molgen.mpg.de

See Also

differential.probs, deregulation.scores

Examples

```
data(damage)
```

```
# Get the probabilities of differential expression
# for the knockouts of ATM, RelA and p53 in healthy cells
probs.healthy= differential.probs(data.healthy, beliefs.healthy)
#Get the regulation scores for ATM, RelA and for p53
regulation.healthy= regulation.scores(probs.healthy, model.healthy, TRUE)
```

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