gene2pathway

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R topics documented:

Index		12
	test.overrepresentation	11
	retrain.signaltrans	
	retrain	8
	ORF2Entrez	7
	internal	7
	getComponents	6
	gene2pathway.signaltrans	4
	gene2pathway	3
	classificationModelSignalTrans	2
	classificationModel	1

classificationModel

Hierarchical Classification Model

Description

This file contains the hierarchical classification model to predict KEGG pathway branches for genes. The model uses a pruned KEGG hierarchy, where metabolic pathways are not distinguished further, and the KEGG hierarchy for "cellular processes" and "genetic information processing" is pruned at the 2nd level. By default the model uses bagging to improve prediction accuracy. Important: There exists one separate model file for each organism.

Format

List of class "model", where each model has the following entries:

- W learned decision hyperplane normal vector
- C dictionary of label vectors, which can be predicted individually or which can be used to predict combinations of them
- **detectors** SVM models trained to separate one specific pathway branch from the rest of the hierarchy
- **used_domains** InterPro domains used by the classifier to separate the specific branch from the rest of the hierarchy

alldomains all InterPro domains used to build feature vectors

allpathways hierarchy branches, which can be predicted

treesizes relative size of hierarchy below the corresponding branch

kegg_hierarchy a nested list with information (parent branches, pathway names, pathway IDs, hierarchy level) on all higher hierarchy branches for each pathway

Author(s)

Holger Froehlich

See Also

classificationModelSignalTrans

classificationModelSignalTrans

Hierarchical Classification Model for Signaling Transduction Pathways and Pathway Components

Description

This file contains the hierarchical classification model to predict KEGG signaling pathways and pathway components for genes. The model contains only pathway components, to which a specified minimum number of genes could be mapped in the training phase (see retrain.signaltrans). Important: There exists one separate model file for each organism.

Format

List of class "model", where each model has the following entries:

W learned decision hyperplane normal vector

C dictionary of label vectors, which can be predicted individually or which can be used to predict combinations of them

detectors SVM models trained to separate one specific pathway branch from the rest of the hierarchy

used_domains InterPro domains used by the classifier to separate the specific branch from the rest of the hierarchy

alldomains all InterPro domains used to build feature vectors

allpathways hierarchy branches, which can be predicted

treesizes" relative size of hierarchy below the corresponding branch

kegg_hierarchy a nested list with information (parent branches, pathway names, pathway IDs, hierarchy level) on all higher hierarchy branches for each pathway

elemIDs a list of KEGG element IDs mapping to each pathway component - may be used to high-light pathway components with color.pathway.by.elements.

Author(s)

Holger Froehlich

See Also

classificationModel

gene2pathway 3

Description

Predicts a gene's membership to a branch in the KEGG hierarchy via the contained InterPro domains.

Usage

```
gene2pathway(geneIDs=NULL, flyBase=FALSE, gene2Domains=NULL, organism="hsa", use
```

Arguments

geneids	a character vector of Entrez gene IDs or FlyBase identifiers (not necessary, if
	the argument gene2Domains is provided)

flyBase Are FlyBase identifiers provided? Default: No

gene2Domains By default associations between genes and InterPro domains are retrieved via

biomaRt from Ensembl. Alternatively, the user can provide its own mapping of

genes to InterPro domains in form of a list here (see details).

organism KEGG letter code describing an organism. Please refer to <URL:http://www.genome.jp/kegg-

bin/create_kegg_menu> for a complete list of organisms (and their letter codes)

supported by KEGG.

useKEGG Should KEGG information instead of a prediction be used when possible?

KEGG. package If useKEGG=TRUE: Instead of retrieving information directly from KEGG, one

can use the KEGG.db package instead, which is significantly faster. However, the KEGG.db package only supports a fraction of organisms so far. Please refer to the manual pages of the KEGG.db package for further information. Default:

use KEGG.db package, if useKEGG=TRUE

Details

A hierarchical classification model based on SVMs and a ranking perceptron is used. This model is usually additionally bagged to improve prediction qualitiy. The model is stored in the package data directory and is recommended to be retrained from time to time.

The current version of the KEGG hierarchy is always retrieved directly from KEGG via FTP. By default associations between genes and InterPro domains are retrieved automatically via biomaRt from Ensembl. Please refer to <URL:http://www.ebi.ac.uk/ensembl/> for a list of organisms supported by Ensembl. Alternatively to using Ensembl and biomaRt, the user can provide its own mapping of genes to InterPro domains in form of a list. This especially allows for using organisms, which are supported by KEGG, but not by Ensembl so far. The list has the form genes -> InterPro domains, and each list entry is named by a gene identifier of the corresponding gene. If useKEGG=TRUE, Entrez gene IDs or FlyBase identifiers have to be used. Otherwise, arbitrary identifiers are allowed.

Value

gene2Path mapping of gene IDs to corresponding KEGG pathway IDs gene2Pathname

mapping of gene IDs to corresponding KEGG pathway names

byKEGG inticates by TRUE/FALSE for each gene whether the mapping information was obtained directly from KEGG or whether it was predicted

confidence scores for the prediction (0, if no prediction was performed): see scores

notes for details

Note

By default a bagged model prediction is used, i.e. each of the individual sub-models is giving a vote for a specific output. The final output is determined by the majority of the votes for each hierarchy branch separately. The corresponding fraction voting for a specific branch may be interpreted as its probability. In the ideal case all individual branch probabilites should always be close to 1, if the gene maps to that part of the KEGG hierarchy, and close to 0 otherwise. A cumulative measure of confindence is thus the average over all probabilities > 0.5 and one minus the average over all probabilities < 0.5. We combine both measure by taking the average of both and report it as a reliability score.

If the user decides to retrain a model WITHOUT using bagging, then the reliability score is simply the margin between the highest and the second highest ranked solution. This margin should be larger 2 for good confindence.

Author(s)

Holger Froehlich

See Also

retrain, classificationModel

Examples

```
## Not run:
gene2pathway("10327")
## End(Not run)
```

gene2pathway.signaltrans

Pathway membership prediction

Description

Predicts a gene's membership to a KEGG signaling pathway and/or pathway component via the contained InterPro domains.

Usage

```
gene2pathway.signaltrans(geneIDs=NULL, flyBase=FALSE, gene2Domains=NULL, organis
```

Arguments

geneIDs a character vector of Entrez gene IDs or FlyBase identifiers (not necessary, if

the argument gene2Domains is provided)

flyBase Are FlyBase identifiers provided? Default: No

gene2Domains By default associations between genes and InterPro domains are retrieved via

biomaRt from Ensembl. Alternatively, the user can provide its own mapping of

genes to InterPro domains in form of a list here (see details).

organism KEGG letter code describing an organism. Please refer to <URL:http://www.genome.jp/kegg-

bin/create_kegg_menu> for a complete list of organisms (and their letter codes)

supported by KEGG.

useKEGG Should KEGG information instead of a prediction be used when possible?

Details

A hierarchical classification model based on SVMs and a ranking perceptron is used. This model is usually additionally bagged to improve prediction qualitiy. The model is stored in the package data directory and is recommended to be retrained from time to time.

The current version of the KEGG hierarchy is always retrieved directly from KEGG via FTP. By default associations between genes and InterPro domains are retrieved automatically via biomaRt from Ensembl. Please refer to <URL:http://www.ebi.ac.uk/ensembl/> for a list of organisms supported by Ensembl. Alternatively to using Ensembl and biomaRt, the user can provide its own mapping of genes to InterPro domains in form of a list. This especially allows for using organisms, which are supported by KEGG, but not by Ensembl so far. The list has the form genes -> InterPro domains, and each list entry is named by a gene identifier of the corresponding gene. If useKEGG=TRUE, Entrez gene IDs or FlyBase identifiers have to be used. Otherwise, arbitrary identifiers can are allowed.

Value

 $\begin{tabular}{ll} gene 2 Path & mapping of gene IDs to corresponding KEGG pathway IDs \\ gene 2 Pathname \\ \end{tabular}$

mapping of gene IDs to corresponding KEGG pathway names

by KEGG inticates by TRUE/FALSE for each gene whether the mapping information was

obtained directly from KEGG or whether it was predicted

scores confindence scores for the prediction (0, if no prediction was performed): see

notes for details

elemIDs KEGG elements mapping to the corresponding predicted pathway components,

if there are any, otherwise NULL. May be used to highlight pathway components

with color.pathway.by.elements.

Note

By default a bagged model prediction is used, i.e. each of the individual sub-models is giving a vote for a specific output. The final output is determined by the majority of the votes for each hierarchy branch separately. The corresponding fraction voting for a specific branch may be interpreted as its probability. In the ideal case all individual branch probabilites should always be close to 1, if the gene maps to that part of the KEGG hierarchy, and close to 0 otherwise. A cumulative measure of confindence is thus the average over all probabilities > 0.5 and one minus the average over all probabilites < 0.5. We combine both measure by taking the average of both and report it as a reliability score.

6 getComponents

If the user decides to retrain a model WITHOUT using bagging, then the reliability score is simply the margin between the highest and the second highest ranked solution. This margin should be larger 2 for good confindence.

Author(s)

Holger Froehlich

See Also

```
retrain.signaltrans, classificationModelSignalTrans
```

Examples

```
## Not run:
  gene2pathway.signaltrans("1443")
## End(Not run)
```

getComponents

KEGG pathway information

Description

1. get connected pathway components; 2. get all elements of a given pathway; 3. color certain elements in a pathway.

Usage

```
getComponents(pathway.id)
get.elements.by.pathway(pathway.id)
color.pathway.by.elements(pathway.id, elements)
```

Arguments

```
pathway.id KEGG pathway ID, e.g. "path:hsa04012" elements KEGG element IDs: character vector of numbers
```

Details

All functions use the KEGG SOAP service.

Value

getComponents: a list with the entries

geneIDs Entrez gene IDs mapping to each pathway component
elemIDs KEGG element IDs mapping to each pathway component

get.elements.by.pathway: list, see <URL http://www.genome.jp/kegg/soap/doc/keggapi_manual.html> for details

color.pathway.by.elements: an URL of a colored gif file, see <URL http://www.genome.jp/kegg/soap/doc/keggapi_manua for details

internal 7

Author(s)

Holger Froehlich

Examples

```
## Not run:
   comp = getComponents("path:hsa04020") # get all connected components
   color.pathway.by.elements("path:hsa04020", comp$elemIDs[[1]]) # mark first component
## End(Not run)
```

internal

internal functions

Description

internal functions: do not call these functions directly.

Usage

various

Arguments

various

Value

various

Author(s)

Holger Froehlich

ORF2Entrez

Conversion locus tag -> Entrez ID and Entrez ID -> locus tag according to KEGG

Description

Conversion locus tag -> Entrez ID and Entrez ID -> locus tag according to KEGG (see <URL:ftp://ftp.genome.jp/pub/keg

Usage

```
ORF2Entrez(ORFIDs, organism="dme")
Entrez2ORF(entrezIDs, organism="dme")
```

8 retrain

Arguments

ORFIDs character vector of locus tags / ORF identifiers

organism KEGG letter code describing an organism. Please refer to <URL:http://www.genome.jp/kegg-

bin/create_kegg_menu> for a complete list of organisms (and their letter codes)

supported by KEGG.

entrezIDs character vector of Entrez gene IDs

Value

character vector

Examples

```
entrez=ORF2Entrez("Dmel_CG4942", organism="dme")
Entrez2ORF(entrez, organism="dme") # should be "Dmel_CG4942"
```

retrain

Retrain classification model

Description

Retrains the hierarchical classification model. This way new information from InterPro and KEGG databases can be incorporated to give better predictions. Retraining should be done on a regular basis from time to time.

Usage

retrain(minnmap=30, level10nly="Metabolism", level20nly="Genetic Information Pro

Arguments

minnmap	prune hierarchy branches with < minnmap mapping genes
level10nly	for these hierarchy branches only the first level is used
1 1 0 0 - 1	for those higgsplay broughes only the first and the second le

level20nly for these hierarchy branches only the first and the second levels are used

organism KEGG letter code describing an organism. Please refer to <URL:http://www.genome.jp/kegg-

bin/create_kegg_menu> for a complete list of organisms (and their letter codes)

supported by KEGG.

gene2Domains By default associations between genes and InterPro domains are retrieved via

biomaRt from Ensembl. Alternatively, the user can provide its own mapping of

genes to InterPro domains in form of a list here (see details).

KEGG. package Instead of retrieving information directly from KEGG, one can use the KEGG.db

package instead, which is significantly faster. However, the KEGG.db package only supports a fraction of organisms so far. Please refer to the manual pages of the KEGG.db package for further information. Default: Don't use KEGG.db

package

remove.duplicates

remove genes having the same InterPro domains prior training. Default: Don't

do this

use.bagging use bagging

nbag number of models to average over

retrain.signaltrans 9

Details

A hierarchical classification model based on SVMs and a ranking perceptron algorithm is trained. This model is usually additionally bagged to improve prediction quality. The method produces a "classificationModel_[organism].rda" (e.g. "classificationModel_hsa.rda") file, which should be stored in the package data directory. Once a new model has been trained, the complete package should be reloaded.

The current version of the KEGG hierarchy is always retrieved directly from KEGG via FTP. By default associations between genes and InterPro domains are retrieved automatically via biomaRt from Ensembl. Please refer to <URL:http://www.ebi.ac.uk/ensembl/> for a list of organisms supported by Ensembl. Alternatively to using Ensembl and biomaRt, the user can provide its own mapping of genes to InterPro domains in form of a list. This especially allows for using organisms, which are supported by KEGG, but not by Ensembl so far. The list has the form genes -> InterPro domains, and each list entry is named by the Entrez gene ID of the corresponding gene. This is, because KEGG uses Entrez gene IDs for the mapping genes -> KEGG pathways.

Value

The model structure. See classificationModel for details.

Author(s)

Holger Froehlich

See Also

gene2pathway, classificationModel

Examples

retrain.signaltrans

Retrain classification model for signaling pathways

Description

Retrains the hierarchical classification model for signaling pathway components. This way new information from InterPro and KEGG databases can be incorporated to give better predictions. Retraining should be done on a regular basis from time to time.

Usage

```
retrain.signaltrans(minnmap=10, organism="hsa", gene2Domains=NULL, remove.duplic
```

10 retrain.signaltrans

Arguments

minnmap prune hierarchy branches with < minnmap mapping genes

organism KEGG letter code describing an organism. Please refer to <URL:http://www.genome.jp/kegg-

bin/create_kegg_menu> for a complete list of organisms (and their letter codes)

supported by KEGG.

gene2Domains By default associations between genes and InterPro domains are retrieved via

biomaRt from Ensembl. Alternatively, the user can provide its own mapping of

genes to InterPro domains in form of a list here (see details).

remove.duplicates

remove genes having the same InterPro domains prior training

use.bagging use bagging

nbag number of models to average over

Details

A hierarchical classification model based on SVMs and a ranking perceptron algorithm is trained. This model is usually additionally bagged to improve prediction quality. The method produces a "classificationModelSignalTrans_[organism].rda" (e.g. "classificationModelSignalTrans_hsa.rda") file, which should be stored in the package data directory. Once a new model has been trained, the complete package should be reloaded.

The current version of the KEGG hierarchy is always retrieved directly from KEGG via FTP. Labels for the training set are obtained via the function <code>getComponents</code>, which uses the KEGG SOAP service. By default associations between genes and InterPro domains are retrieved automatically via biomaRt from Ensembl. Please refer to <URL:http://www.ebi.ac.uk/ensembl/> for a list of organisms supported by Ensembl. Alternatively to using Ensembl and biomaRt, the user can provide its own mapping of genes to InterPro domains in form of a list. This especially allows for using organisms, which are supported by KEGG, but not by Ensembl so far. The list has the form genes -> InterPro domains, and each list entry is named by the Entrez gene ID of the corresponding gene. This is, because KEGG uses Entrez gene IDs for the mapping genes -> KEGG pathways.

Value

The model structure. See classificationModelSignalTrans for details.

Author(s)

Holger Froehlich

See Also

```
gene2pathway.signaltrans,classificationModelSignalTrans
```

Examples

```
## Not run:
    retrain.signaltrans() # retrain classification model for signal transduction path
## End(Not run)
```

test.overrepresentation 11

```
test.overrepresentation
```

Test statistical overrepresentation of KEGG pathways in a list of genes

Description

Test the statistical overrepresentation of KEGG pathways in a group of genes using Fisher's exact test. The analysis can either be based on all KEGG pathways predicted by gene2pathway/gene2pathway.signal or on original KEGG annotation only.

Usage

```
test.overrepresentation(genesOfInterest, predpath, KEGGonly=FALSE, cutoff=0.1, m
```

Arguments

genesOfInterest

a character vector of gene identifiers (see gene2pathway, gene2pathway. signaltrans)

for a gene list of interest

predpath predictions of gene2pathway or gene2pathway.signaltrans

KEGGonly use KEGG annotation only cutoff p-value significance cutoff

min.conf filter predictions such that only those with a confidence score > min.conf are

considered

adj.method multiple testing correction method. Default: Benjamini-Yekutieli

Value

Table with two columns: KEGG pathway and adjusted p-value (adjustment according to Benjamini-Yekutieli)

Index

```
*Topic datasets
                                         modelKEGG(classificationModel), 1
   classificationModel, 1
                                         modelSignalTrans
   classificationModelSignalTrans,
                                                 (classificationModelSignalTrans),
*Topic file
                                         ORF2Entrez, 7
   gene2pathway, 3
   gene2pathway.signaltrans,4
                                         predict.gene2pathway(internal), 7
   getComponents, 6
   internal, 7
                                         retrain. 4.8
   ORF2Entrez, 7
                                         retrain.signaltrans, 2, 6, 9
   retrain, 8
   retrain.signaltrans,9
                                         struct_predict(internal),7
   test.overrepresentation, 11
                                         struct_train(internal),7
                                         svmlearn(internal),7
buildTrainingSet (internal), 7
                                         sympredict (internal), 7
classificationModel, 1, 2, 4, 9
                                         test.overrepresentation, 11
classificationModelSignalTrans,
       1, 2, 6, 10
code_test(internal), 7
code_train(internal),7
color.pathway.by.elements, 2, 5
color.pathway.by.elements
       (getComponents), 6
Entrez2ORF (ORF2Entrez), 7
Entrez2ORF.internal(internal), 7
gene2pathway, 3, 9, 11
gene2pathway.signaltrans, 4, 10, 11
get.element.relations.by.pathway
       (internal), 7
get.elements.by.pathway
       (getComponents), 6
getComponents, 6, 10
getComponents.internal
       (internal), 7
getInterProDomains (internal), 7
getKEGGHierarchy (internal), 7
internal, 7
KEGG2Entrez (internal), 7
loss (internal), 7
```