# Package 'PADOG'

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Version 1.50.0 Date 2020-1-31 Title Pathway Analysis with Down-weighting of Overlapping Genes (PADOG) Author Adi Laurentiu Tarca <atarca@med.wayne.edu>; Zhonghui Xu <zhonghui.xu@gmail.com> **Depends** R (>= 3.0.0), KEGGdzPathwaysGEO, methods, Biobase Suggests doParallel, parallel Maintainer Adi L. Tarca <atarca@med.wayne.edu> Description This package implements a general purpose gene set analysis method called PADOG that downplays the importance of genes that apear often accross the sets of genes to be analyzed. The package provides also a benchmark for gene set analysis methods in terms of sensitivity and ranking using 24 public datasets from KEGGdzPathwaysGEO package. License GPL (>= 2) Collate padog.R compPADOG.R filteranot.R Imports limma, AnnotationDbi, GSA, foreach, doRNG, hgu133plus2.db, hgu133a.db, KEGGREST, nlme LazyLoad yes biocViews Microarray, OneChannel, TwoChannel git\_url https://git.bioconductor.org/packages/PADOG git\_branch RELEASE\_3\_21 git\_last\_commit 4f7fb8c git\_last\_commit\_date 2025-04-15 **Repository** Bioconductor 3.21 Date/Publication 2025-07-23

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#### compPADOG

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compPADOG

#### Description

This is a general purpose function to compare a given gene set analysis method in terms of sensitivity and ranking agains PADOG and GSA (if installed) using 24 public datasets.

#### Usage

#### Arguments

| datasets        | A character vector with valid names of datasets to use from the PADOGsets package. If left NULL all datasets availbale in PADOGsets will be used.   |
|-----------------|---|
| existingMethods | S   |
|                 | A character vector with one or more of the predefined methods c("GSA","PADOG"). The first is used as reference method.  |
| mymethods       | A list whose elements are valid functions implementing gene set analysis meth-<br>ods. See the example to see what arguments the functions have to take in and<br>what kind of output they need to produce.   |
| gslist          | Either the value "KEGGRESTpathway" or a list with the gene sets. If set to "KEGGRESTpathway", then gene sets will be made of all KEGG pathways for human since all datasets available in PADOG are for human. |
| organism        | A three letter string giving the name of the organism supported by the "KEG-GRESTpathway" package.  |
| gs.names        | A character vector giving additional information about each gene set. For in-<br>stance when gene seta are pathways, the full name of the pathway would be a<br>meaningful gene set name.                     |
| Nmin            | The minimum size of gene sets to be included in the analysis for all methods.   |
| NI              | Number of iterations to determine the gene set score significance p-values in PADOG and GSA methods.  |
| parallel        | Should paralell be used if multiple cores are available and the package parallel is available. If se to TRUE one dataset will be run on on multiple CPU at a time (Not available on Windows).                 |
| ncr             | The number of CPU cores used when use.parallel set to TRUE. Default is to use all CPU cores detected.   |
| pkgs            | Character vector of packages that the existingMethods and mymethods depend on (NULL for "PADOG" and "GSA"). Consult the .packages argument in foreach function from foreach package.                          |

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#### compPADOG

| expVars | Character vector of variables to export. Consult the .export argument in foreach function from foreach package.                 |
|---------|---|
| dseed   | Optional initial seed for random number generator (integer) used in padog.  |
| plots   | If set to TRUE will plot the ranks of the target genesets and the ranks differences between a methods and the reference method. |
| verbose | This argument will be passed to PADOG and AbsmT methods. If set to TRUE it will show the interations performed so far.          |

#### Details

See cited documents for more details.

#### Value

#### Author(s)

Adi Laurentiu Tarca <atarca@med.wayne.edu>

#### References

Adi L. Tarca, Sorin Draghici, Gaurav Bhatti, Roberto Romero, Down-weighting overlapping genes improves gene set analysis, BMC Bioinformatics, 13(136), 2012.

Adi L. Tarca, Gaurav Bhatti, Roberto Romero, A Comparison of Gene Set Analysis Methods in Terms of Sensitivity, Prioritization and Specificity, PLoS One. 8(11), 2013.

#### See Also

compPADOG

#### Examples

#compare a new geneset analysis method with PADOG and GSA

#define your new gene set analysis method that takes as input: #set- the name of dataset file from the PADOGsetspackage #mygslist - a list with the genesets

```
#minsize- minimum number of genes in a geneset to be considered for analysis
```

```
randomF=function(set,mygslist,minsize){
  set.seed(1)
  #this loads the dataset in an ExpressionSet object called x
  data(list=set,package="KEGGdzPathwaysGE0")
  x=get(set)
```

```
#Extract from the dataset the required info to be passed to padog
exp=experimentData(x);
dat.m=exprs(x)
ano=pData(x)
dataset= exp@name
design= notes(exp)$design
annotation= paste(x@annotation,".db",sep="")
targetGeneSets= notes(exp)$targetGeneSets
```

```
#get rid of duplicates probesets per ENTREZ ID by keeping the probeset
#with smallest p-value (computed using limma)
aT1=filteranot(esetm=dat.m,group=ano$Group,paired=(design=="Paired"),
block=ano$Block.annotation=annotation)
#create an output dataframe for this toy method with random gene set p-values
mygslistSize=unlist(lapply(mygslist,function(x){length(intersect(aT1$ENTREZID,x))}))
res=data.frame(ID=names(mygslist),P=runif(length(mygslist)),
Size=mygslistSize,stringsAsFactors=FALSE)
res$FDR=p.adjust(res$P,"fdr")
#drop genesets with less than minsize genes in the current dataset
res=res[res$Size>=minsize,]
#compute ranks
res$Rank=rank(res$P)/dim(res)[1]*100
#needed to compare ranks between methods; must be the same as given
#in mymethods argument "list(myRand="
res$Method="myRand";
#needed because comparisons of ranks between methods is paired at dataset level
res$Dataset<-dataset:
#output only result for the targetGeneSets
#which are gene sets expected to be relevant in this dataset
return(res[res$ID %in% targetGeneSets,])
}
```

```
#run the analysis on all 24 datasets and compare the new method "myRand" with
#PADOG and GSA (if installed) (chosen as reference since is listed first in the existingMethods)
#if the package parallel is installed datasets are analyzed in parallel.
#out=compPADOG(datasets=NULL,existingMethods=c("GSA","PADOG"),
#mymethods=list(myRand=randomF),
#gslist="KEGGRESTpathway",Nmin=3,NI=1000,plots=TRUE,verbose=FALSE)
```

```
#compare myRand against PADOG on 4 datasets only
#mysets=data(package="PADOGsets")$results[,"Item"]
mysets=c("GSE9348","GSE8671","GSE1297")
out=compPADOG(datasets=mysets,existingMethods=c("PADOG"),
mymethods=list(myRand=randomF),
```

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#### filteranot

gslist="KEGGRESTpathway",Nmin=3,NI=20,plots=FALSE,verbose=FALSE)

| filteranot | Remove duplicate probesets/probes from an gene expression matrix<br>based on p-values from a moderated t-test, in order to apply a gene set<br>analysis. |
|------------|--|
|            | analysis.  |

# Description

This function helps to deal with multiple probesets/probes per gene prior to geneset analysis.

#### Usage

filteranot(esetm=NULL,group=NULL,paired=FALSE,block=NULL,annotation=NULL,include.details=FALSE)

## Arguments

| esetm           | A matrix containing log transformed and normalized gene expression data. Rows correspond to genes and columns to samples. Rownames of esetm need to be valid probeset or probe names. |  |  |
|-----------------|---|--|--|
| group           | A character vector with the class labels of the samples. It can only contain "c" for control samples or "d" for disease samples.  |  |  |
| paired          | A logical value to indicate if the samples in the two groups are paired.  |  |  |
| block           | A character vector indicating the block ids of the samples classified by the group variable, if paired=TRUE. The paired samples must have the same block value.                       |  |  |
| annotation      | A valid chip annotation package name (e.g. "hgu133plus2.db")  |  |  |
| include.details |   |  |  |
|                 |   |  |  |

If set to true, will include all columns from limma's topTable for this dataset.

#### Details

See cited documents for more details.

#### Value

A data frame containing the probeset IDs (and corresponding ENTREZ IDs) of the best probesets for each gene ;

#### Author(s)

Adi Laurentiu Tarca <atarca@med.wayne.edu>

#### References

Adi L. Tarca, Sorin Draghici, Gaurav Bhatti, Roberto Romero, Down-weighting overlapping genes improves gene set analysis, BMC Bioinformatics, 2012, submitted.

Adi L. Tarca, Gaurav Bhatti, Roberto Romero, A Comparison of Gene Set Analysis Methods in Terms of Sensitivity, Prioritization and Specificity, PLoS One. 8(11), 2013.

#### See Also

padog

#### Examples

```
#run padog on a colorectal cancer dataset of the 24 datasets benchmark GSE9348
set="GSE9348"
data(list=set,package="KEGGdzPathwaysGEO")
x=get(set)
#Extract from the dataset the required info
exp=experimentData(x);
dataset= exp@name
dat.m=exprs(x)
ano=pData(x)
design= notes(exp)$design
annotation= paste(x@annotation,".db",sep="")
```

dim(dat.m)

#get rid of duplicates in the same way as is done for PADOG and assign probesets to ENTREZ IDS
#get rid of duplicates by choosing the probe(set) with lowest p-value; get ENTREZIDs for probes
aT1=filteranot(esetm=dat.m,group=ano\$Group,paired=(design=="Paired"),block=ano\$Block,annotation)

```
#filtered expression matrix
filtexpr=dat.m[rownames(dat.m)%in%aT1$ID,]
dim(filtexpr)
```

| р | adog | Pathway<br>(PADOG | • | with | Down-weighting | of | Overlapping | Genes |  |
|---|------|-------------------|---|------|----------------|----|-------------|-------|--|
|   |      | (IADOO            | ) |      |                |    |             |       |  |

#### Description

This is a general purpose gene set analysis method that downplays the importance of genes that apear often accross the sets of genes analyzed. The package provides also a benchmark for gene set analysis in terms of sensitivity and ranking using 24 public datasets.

### padog

# Usage

# Arguments

| esetm      | A matrix containing log transformed and normalized gene expression data. Rows correspond to genes and columns to samples.  |
|------------|--|
| group      | A character vector with the class labels of the samples. It can only contain "c" for control samples or "d" for disease samples.   |
| paired     | A logical value to indicate if the samples in the two groups are paired.   |
| block      | A character vector indicating the block ids of the samples classified by the group variable, if paired=TRUE. The paired samples must have the same block value.  |
| gslist     | Either the value "KEGGRESTpathway" or a list with the gene sets. If set to "KEGGRESTpathway", then gene sets will be made of all KEGG pathways for the organism specified. If a list is provided, instead, each element of the list should be a character vector with the identifiers for the genes. The identifiers can be probe(sets) ids if the annotation argument is set to a valid annotation package, otherwise the gene identifiers must be of the same kind as the row-names of the matrix esetm. |
| annotation | A valid chip annotation package if the rownames of esetm are probe(set) ids<br>and gslist contains ENTREZ identifiers or gslist is set to "KEGGREST-<br>pathway". If the rownames are other gene identifies, then annotation has tyo<br>be set to NULL, and the row names of esetm needs to be unique and be found<br>among elements of gslist   |
| organism   | A three letter string giving the name of the organism supported by the "KEG-GREST" package.  |
| gs.names   | Character vector with the names of the gene sets. If specified, must have the same length as gslist.   |
| NI         | Number of iterations to determine the gene set score significance p-values.  |
| plots      | If set to TRUE then the distribution of the PADOG scores with and without weighting the genes in raw and standardized form are shown using boxplots. A pdf file will be created in the current directory having the name provided in the targetgs field. The scores for the targetgs gene set will be shown in red.  |
| targetgs   | The identifier of a traget gene set for which the scores will be highlighted in the plots produced if plots=TRUE   |
| Nmin       | The minimum size of gene sets to be included in the analysis.  |
| verbose    | If set to TRUE, displays the number of iterations elapsed is displayed.  |
| parallel   | If set to TRUE, the NI iterations will be executed in parallel if multiple CPU cores are available and foreach and doRNG packages are installed.   |
| dseed      | Optional initial seed for random number generator (integer).   |
| ncr        | The number of CPU cores used when parallel set to TRUE. Default is to use all CPU cores detected.  |

#### Details

See cited documents for more details.

#### Value

A data frame containing the ranked pathways and various statistics: Name is the name of the gene set; ID is the gene set identifier; Size is the number of genes in the geneset; meanAbsT0 is the mean of absolute t-scores; padog0 is the mean of weighted absolute t-scores; PmeanAbsT significance of the meanAbsT0; Ppadog is the significance of the padog0 score;

#### Author(s)

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#### References

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Adi L. Tarca, Gaurav Bhatti, Roberto Romero, A Comparison of Gene Set Analysis Methods in Terms of Sensitivity, Prioritization and Specificity, PLoS One. 8(11), 2013.

#### See Also

padog

#### Examples

```
#run padog on a colorectal cancer dataset of the 24 datasets benchmark GSE9348
#use NI=1000 for accurate results.
set="GSE9348"
data(list=set,package="KEGGdzPathwaysGEO")
x=get(set)
#Extract from the dataset the required info
exp=experimentData(x);
dataset= exp@name
dat.m=exprs(x)
ano=pData(x)
design= notes(exp)$design
annotation= paste(x@annotation,".db",sep="")
targetGeneSets= notes(exp)$targetGeneSets
```

```
myr=padog(
esetm=dat.m,
group=ano$Group,
paired=design=="Paired",
block=ano$Block,
targetgs=targetGeneSets,
annotation=annotation,
gslist="KEGGRESTpathway",
```

# padog

```
organism="hsa",
verbose=TRUE,
Nmin=3,
NI=25,
plots=FALSE,
dseed=1)
```

```
myr2=padog(
esetm=dat.m,
group=ano$Group,
paired=design=="Paired",
block=ano$Block,
targetgs=targetGeneSets,
annotation=annotation,
gslist="KEGGRESTpathway",
organism="hsa",
verbose=TRUE,
Nmin=3,
NI=25,
plots=FALSE,
dseed=1,
paral=TRUE,
ncr=2)
```

myr[1:20,]

all.equal(myr, myr2)

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