# Package 'directPA'

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<b>Description</b> Direction analysis is a set of tools designed to identify combinatorial effects of multiple treatments/conditions on pathways and kinases profiled by microarray, RNA-seq, proteomics, or phosphoproteomics data. See Yang P et al (2014) <doi:10.1093 bioinformatics="" btt616="">; and Yang P et al. (2016) <doi:10.1002 pmic.2016000<="" td=""></doi:10.1002></doi:10.1093>
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Contents

directPA-package	 	 	 	 		 	•		•		2
directExplorer2d	 	 	 	 		 					3
directPA											
geneStats	 	 	 	 		 					5
HEK											
kinasePA	 	 	 	 		 					6
Pathways.KEGG	 	 	 	 		 					8
Pathways.reactome											
pathwayStats	 	 	 	 		 					8
perturbPlot2d	 	 	 	 		 					9
perturbPlot3d	 	 	 	 		 					10

2 directPA-package

dire	ctPA-package	Dir	ecti	on.	Pat	hw	ay	Αı	nai	lys	sis	Pα	ıck	kag	ge										
Index																									14
	rotate3d	• •				•		•	•	•			•	•	•	 •	•	•	 •	•	•	 •	 •	•	12
	rotate2d																								
	PM																								
	PhosphoSite.mouse																								
	PhosphoSite.human	١																							11
	PhosphoELM.mous	se .																							11
	PhosphoELM.huma	ın .																							10

# Description

The directPA-package is designed for analysing pathways in experiments with multiple perturbations. The combination effects of different treatments are tested by rotating polar coordinates in two-dimentional space when the experiment contains two perturbations and corresponding controls, or spherical coordinates in three-dimensional space when the experiment contains three perturbations and corresponding controls.

#### **Details**

Package: directPA
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#### References

Pengyi Yang, Ellis Patrick, Shi-Xiong Tan, Daniel J. Fazakerley, James Burchfield, Christopher Gribben, Matthew J. Prior, David E. James, Yee Hwa Yang, Direction pathway analysis of large-scale proteomics data reveals novel features of the insulin action pathway, Bioinformatics, 30(6), 808-814, 2014.

directExplorer2d 3

directExplorer2d	Batch Direction Analysis in 2-dimentional space	
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# Description

Rotate to the direction of interest in polar coordinates by degree (e.g. pi/4).

# Usage

```
directExplorer2d(Tc, annotation=NULL, gene.method="OSP",
path.method="Stouffer", top=10, nd=8, ...)
```

#### **Arguments**

Тс	a numeric matrix with 2 columns. The rows are genes or phosphorylation sites and the columns are treatments vs control statistics.
annotation	a list with names correspond to pathways or kinases and elements correspond to genes or substrates belong to each pathway or kinase, respectively.
gene.method	the method to be used for integrating statistics across treatments for each gene or phosphorylation site. Available methods are Stouffer, OSP, Fisher, and maxP. Default method is OSP.
path.method	the method to be used for integrating statistics of all genes or phosphorylation sites that belongs to a pathway or kinase. Available methods are Stouffer, OSP, Fisher, and maxP. Default method is Stouffer.
top	the number of entries to be highlighted in the plot.
nd	the number of directions to plot (4 or 8)
	parameters for controlling the plot.

#### Value

The the list of enrichment analysis in tables.

# **Examples**

```
# load the phosphoproteomics dataset
data(HEK)

# load the kinase-substrate annoations
data(PhosphoSite)

# test enrichment on 8 directions in polar coordinate system.
bda <- directExplorer2d(Tc=HEK, annotation=PhosphoSite.mouse)

# the direction are denoted as follow for the two treatments vs control:
# ++: up-regulated in both treatments
# +*: up-regulated in the first treatment and unchanged in the second treatment</pre>
```

4 directPA

```
# +-: up-regulated in the first treatment and down-regulated in the second treatment
# *-: unchanged in the first treatment and down-regulated in the second treatment
# --: down-regulated in both treatments
# -*: down-regulated in the first treatment and unchanged in the second treatment
# -+: down-regulated in the first treatment and up-regulated in the second treatment
# *+: unchanged in the first treatment and up-regulated in the second treatment
# sort the most enriched phosphorylation sites and kinases on down-regulation from both
# treatments (i.e. "--") and displa the top-10 entries
bda$gene.tab[order(bda$gene.tab[,"--"]),][1:10,]
bda$path.tab[order(bda$path.tab[,"--"]),][1:10,]
```

directPA

Direction Analysis for Pathways

#### **Description**

The main function of direction Analysis. This function takes in a matrix of test statistics with two (2-dimensional space) or three (3-dimensional space) columns, the direction of interests, and the annotation list such as pathway annotation, and test for enrichment of pathways on the specified direction.

#### Usage

```
directPA(Tc, direction, annotation, minSize=5, gene.method="OSP",
path.method="Stouffer", visualize=TRUE, ...)
```

#### **Arguments**

Tc	a numeric matrix. Rows are genes and columns are treatments vs control statistics.
direction	the direction to be tested for enrichment. Either specified as a degree for two-dimensional analysis or as contrast (in a triplet) for three-dimensional analysis.
annotation	a list with names correspond to pathways and elements correspond to genes belong to each pathway, respectively.
minSize	the size of annotation groups to be considered for calculating enrichment. Groups that are smaller than the minSize will be removed from the analysis.
gene.method	the method to be used for integrating statistics across treatments for each gene. Available methods are Stouffer, OSP, Fisher, and maxP. Default method is OSP.
path.method	the method to be used for integrating statistics of all genes that belongs to a pathway. Available methods are Stouffer, OSP, Fisher, and maxP. Default method is Stouffer.
visualize	whether to visualize the plot.
	other visualization parameters to pass on.

geneStats 5

#### Value

a list that contains directional p-values for each gene and directional enrichment for each pathway.

#### **Examples**

```
# load the proteomics dataset
data(PM)
# load pathway annotations
data(Pathways)
# display reactome pathways. Could be replaced by any other pathway databases
Pathways.reactome[1:5]
# direction pathway analysis in 3-dimensional space. Implemnted as rotating by contrast
# (1) test combined effect of all 3 treatments (stimulation and inhibitions) vs control (basal)
# on the original direction.
dPA <- directPA(Tc=PM, direction=c(1,1,1), annotation=Pathways.reactome)</pre>
dPA$gst[order(unlist(dPA$gst[,1])),][1:20,]
# rank substrates on the direciton of interest
sort(dPA$gene.pvalues)[1:20]
# (2) test combined effect of all 3 treatments vs controls on direction c(1,-1, ∅)
# this rotates Ins by 0 degree, Wmn by 90 degree, and MK by 45 degree.
dPA <- directPA(Tc=PM, direction=c(1,-1,0), annotation=Pathways.reactome)
dPA$gst[order(unlist(dPA$gst[,1])),][1:20,]
# (3) test combined effect of all 3 perturbations vs controls on direction c(1,-1, 1)
# this rotates Ins by 0 degree, Wmn by 90 degree, and MK by 0 degree.
dPA <- directPA(Tc=PM, direction=c(1,-1,1), annotation=Pathways.reactome)
dPA$gst[order(unlist(dPA$gst[,1])),][1:20,]
```

geneStats

Molecule Level Statistics

# Description

Takes a vector of statistics with each element corresponds to a treatment vs control comparison, and calculates a combined statistics across multiple treatments.

#### Usage

```
geneStats(T, method="OSP")
```

#### **Arguments**

Τ

a vector of statistics (z-scores converted) with each element correspond to a treatment vs control comparison.

6 kinasePA

method

the p-value integration method for combining accross multiple treatments. Available methods are Stouffer, OSP, Fisher, and maxP. The default method is OSP.

#### Value

a p-value after integration across treatments.

#### **Examples**

```
# load the example data
data(PM)

# convert statistics into z-scores
PM.zscores <- apply(PM, 2, function(x){qnorm(rank(x)/(nrow(PM)+1))})

# Rotate the matrix by contrast 1, -1, -1 (i.e. up-regulation, down-regulation, down-regulation).
PM.rotated <- rotate3d(PM.zscores, contrast = c(1, -1, -1))

# combine rotated statistics across treatments
gene.pvalues <- apply(PM.rotated, 1, geneStats)</pre>
```

HEK

Phosphoproteomics of HEK-293E

# Description

The data object contains all quantified phosphoryaltion sites in HEK-293E cells with Rapamycin or Torin1 treatment vs insulin stimulation. See Hsu et al. Science, 332(6035), 1317-1322, 2011, for more details.

kinasePA

Direction Analysis for Kinases

#### Description

This is a wrapper for runing directPA for kinase perturbation analysis (kinasePA)

# Usage

```
kinasePA(Tc, direction, annotation, minSize=5, substrate.method="OSP",
kinase.method="Stouffer", visualize=TRUE, ...)
```

kinasePA 7

#### **Arguments**

Tc a numeric matrix. The columns are phosphorylation sites and the columns are treatments vs control statistics.

direction the direction to be tested for enrichment. Either specified as a degree for two-dimensional analysis or as contrast (in a triplet) for three-dimensional analysis.

annotation a list with names correspond to kinases and elements correspond to substrates

belong to each kinase, respectively.

minSize the size of annotation groups to be considered for calculating enrichment. Groups

that are smaller than the minSize will be removed from the analysis.

substrate.method

the method to be used for integrating statistics across treatments for each substrate (phosphorylation site). Available methods are Stouffer, OSP, Fisher, and

maxP. Default method is OSP.

kinase.method the method to be used for integrating statistics of all phosphorylation sites that

belongs to a kinase. Available methods are Stouffer, OSP, Fisher, and maxP.

Default method is Stouffer.

visualize whether to visualize the plot.

... other visualization parameters to pass on.

#### Value

a list that contains directional p-values for each substrate and directional enrichment for each kinase.

#### **Examples**

```
# load the phosphoproteomics dataset
data(HEK)
# load the kinase-substrate annoations
data(PhosphoSite)
# direction pathway analysis in 2-dimensional space. Implemented as rotating by degree
# (1) test combined effect of Torin1 and Rapamycin vs insul both on "down-regulation"
# (180 degree to original direction)
kPA <- kinasePA(Tc=HEK, direction=pi, annotation=PhosphoSite.mouse)
kPA$kinase[order(unlist(kPA$kinase[,1])),][1:20,]
# rank substrates on the direciton of interest
sort(kPA$substrate.pvalues)[1:20]
# (2) test combined effect of Torin1 and Rapamycin vs insul on "no change and down-regulation"
# (135 degree to the original direction)
kPA <- kinasePA(Tc=HEK, direction=pi*3/4, annotation=PhosphoSite.mouse)</pre>
kPA$kinase[order(unlist(kPA$kinase[,1])),][1:20,]
# (3) test combined effect of Torin1 and Rapamycin vs insul on "down-regulation and no change"
# (225 degree to the original direction)
kPA <- kinasePA(Tc=HEK, direction=pi*5/4, annotation=PhosphoSite.mouse)
kPA$kinase[order(unlist(kPA$kinase[,1])),][1:20,]
```

8 pathwayStats

Pathways.KEGG	KEGG pathway annotations	
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# Description

The data object contains the annotations of KEGG pathways.

Pathways.reactome	Reactome pathway annotations	

# Description

The data object contains the annotations of reactome pathways.

# Description

Takes a vector of statistics with each element corresponds to a gene or phosphorylation site, and calculates a combined statistics for those that belong to the same pathway or kinase.

# Usage

```
pathwayStats(PGs, T, minSize=5, method="Stouffer")
```

# Arguments

PGs	an array of names indicating genes or substrates that belong to a given pathway or kinase.
Т	a vector of statistics (z-scores converted) with each element correspond to a gene or phosphorylation site that belong to the same pathway or kinase.
minSize	the size of annotation groups to be considered for calculating enrichment. Groups that are smaller than the minSize will be removed from the analysis.
method	the p-value integration method for combining accross multiple treatments. Available methods are Stouffer, OSP, Fisher, and maxP. The default method is Stouffer.

#### Value

a doublet corresponding to the enrichment after integration across all genes or substrates that belong to the same pathway or kinase, and the size of the mapped genes or substrates to that pathway or kinase.

perturbPlot2d 9

#### **Examples**

```
# load the example data
data(PM)

# load pathway annotations
data(Pathways)

# convert statistics into z-scores
PM.zscores <- apply(PM, 2, function(x){qnorm(rank(x)/(nrow(PM)+1))})

# Rotate the matrix by contrast 1, -1, -1 (i.e. up-regulation, down-regulation, down-regulation).
PM.rotated <- rotate3d(PM.zscores, contrast = c(1, -1, -1))

# combine rotated statistics across treatments
gene.pvalues <- apply(PM.rotated, 1, geneStats)

# compute statistics for all reactome pathways
gene.zscores <- qnorm(gene.pvalues, lower.tail = FALSE)
gst <- t(sapply(Pathways.reactome, pathwayStats, gene.zscores))</pre>
```

perturbPlot2d

Perturbation Plot

#### **Description**

This function takes in a matrix of test statistics with two columns (2-dimensional space) and the annotation list such as pathway annotation or kinase-substrate annotation, and visualize the enrichment of pathways or kinases in direction specific manner.

#### Usage

```
perturbPlot2d(Tc, annotation, minSize=5, ...)
```

# Arguments

Tc	a numeric matrix. The columns are genes or phosphorylation sites and the columns are treatments vs control statistics.
annotation	a list with names correspond to pathways or kinases and elements correspond to genes or substrates belong to each pathway or kinase, respectively.
minSize	the size of annotation groups to be considered for calculating enrichment. Groups that are smaller than the minSize will be removed from the analysis.
	parameters for controling the plot.

#### Value

a list of coordinates for pathways or kinases

10 PhosphoELM.human

#### **Examples**

```
# load the phosphoproteomics dataset
data(HEK)
# load the kinase-substrate annoations
data(PhosphoSite)
perturbPlot2d(Tc=HEK, annotation=PhosphoSite.mouse, cex=3)
```

perturbPlot3d

Perturbation Plot 3D

#### **Description**

This function takes in a matrix of test statistics with two columns (3-dimensional space) and the annotation list such as pathway annotation or kinase-substrate annotation, and visualize the enrichment of pathways or kinases in direction specific manner.

#### Usage

```
perturbPlot3d(Tc, annotation, minSize=5, ...)
```

#### **Arguments**

Tc	a numeric matrix. The columns are genes or phosphorylation sites and the columns are treatments vs control statistics.
annotation	a list with names correspond to pathways or kinases and elements correspond to genes or substrates belong to each pathway or kinase, respectively.
minSize	the size of annotation groups to be considered for calculating enrichment. Groups that are smaller than the minSize will be removed from the analysis.
	parameters for controling the plot.

#### Value

a list of coordinates for pathways or kinases

PhosphoELM.human PhosphoELM annotations for human

#### **Description**

The data object contains the annotations of kinases and their conrresponding substrates as phosphorylation sites in human. It is extracted from the PhosphoELM database. For details of PhosphoELM, please refer to the article: Phospho.ELM: a database of phosphorylation sites—update 2008. Diella, F., Gould, C.M., Chica, C., Via, A. & Gibson, T.J. Nucleic Acids Res. 2008 Jan;36(Database issue):D240-4.

PhosphoELM.mouse 11

PhosphoELM.mouse PhosphoELM annotations for mouse	PhosphoELM annotations for mouse
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# Description

The data object contains the annotations of kinases and their conrresponding substrates as phosphorylation sites in mouse. It is extracted from the PhosphoELM database. For details of PhosphoELM, please refer to the article: Phospho.ELM: a database of phosphorylation sites—update 2008. Diella, F., Gould, C.M., Chica, C., Via, A. & Gibson, T.J. Nucleic Acids Res. 2008 Jan;36(Database issue):D240-4.

PhosphoSite.human Ph	nosphoSitePlus annotations for human
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#### **Description**

The data object contains the annotations of kinases and their conrresponding substrates as phosphorylation sites in human. It is extracted from the PhosphoSitePlus database. For details of PhosphoSitePlus, please refer to the article: Hornbeck et al. Nucleic Acids Res., 40:D261-70, 2012.

PhosphoSite.mouse PhosphoSitePlus annotations for mouse	PhosphoSite.mouse	PhosphoSitePlus annotations for mouse
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#### **Description**

The data object contains the annotations of kinases and their conrresponding substrates as phosphorylation sites in mouse. It is extracted from the PhosphoSitePlus database. For details of PhosphoSitePlus, please refer to the article: Hornbeck et al. Nucleic Acids Res., 40:D261-70, 2012.

PM	Plasma Membrame Protoemics Data

#### **Description**

The data object contains all quantified plasma membrame proteins with insulin simulation with and without prior treatments of MK or Wmn vs basal condition in mouse 3T3L1 cell lines. Please refer to the article: Yang et al. Bioinformatics, 30(6):808-14, 2014.

12 rotate3d

rotate2d

Polar Coordinates Rotation

# **Description**

Rotate to the direction of interest in polar coordinates by degree (e.g. pi/4).

#### Usage

```
rotate2d(T, degree = 0)
```

#### **Arguments**

T a numeric matrix (normally z-score converted) with 2 columns. The rows are

genes or phosphorylation sites and the columns are treatments vs control statis-

tics.

degree the direction to be tested for enrichment. Specified as degree from to the original

direction.

#### Value

A rotated matrix with respect to the direction of interest.

#### **Examples**

```
# load the phosphoproteomics dataset
data(HEK)

# convert statistics into z-scores
HEK.zscores <- apply(HEK, 2, function(x){qnorm(rank(x)/(nrow(HEK)+1))})

# Rotate the matrix by 1/2 pi (i.e. down-regulation, dow-regulation).
HEK.rotated <- rotate2d(HEK.zscores, degree = pi/2)</pre>
```

rotate3d

Spherical Coordinates Rotation

#### **Description**

Rotate to the direction of interest in spherical coordinates by contrasts (e.g. 1, -1, -1).

#### Usage

```
rotate3d(T, contrast)
```

rotate3d 13

#### **Arguments**

T a numeric matrix (normally z-score converted) with 3 columns. The rows are

genes or phosphorylation sites and the columns are treatments vs control statis-

tics.

contrast the direction to be tested for enrichment. Specified as contrast in a triplet (see

example).

#### Value

A rotated matrix with respect to the direction of interest.

# **Examples**

```
# load the example data
data(PM)

# convert statistics into z-scores
PM.zscores <- apply(PM, 2, function(x){qnorm(rank(x)/(nrow(PM)+1))})

# Rotate the matrix by contrast 1, -1, -1 (i.e. up-regulation, down-regulation, down-regulation).
PM.rotated <- rotate3d(PM.zscores, contrast = c(1, -1, -1))</pre>
```

# **Index**

```
directExplorer2d, 3
directPA, 4
directPA-package, 2
geneStats, 5
HEK, 6
kinasePA, 6
{\tt Pathways.KEGG, 8}
Pathways.reactome, 8
pathwayStats, 8
\verb"perturbPlot2d", 9"
perturbPlot3d, 10
{\it PhosphoELM.human}, 10
PhosphoELM.mouse, 11
PhosphoSite.human, 11
PhosphoSite.mouse, 11
PM, 11
rotate2d, 12
rotate3d, 12
```