Package 'KSEAapp'

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KSData

Kinase-Substrate (K-S) Relationship Dataset

Description

K-S annotations from PhosphoSitePlus and NetworKIN predictions; This is an abbreviated version of the full dataset used purely for demonstration; please go to the GitHub page for access to the complete file: github.com/casecpb/KSEA/

Usage

data(KSData)

Format

abbreviated dataframe containing the kinase-substrate annotations and source

References

Hornbeck et al. (2015) Nucleic Acids Res. 43:D512-20 Horn et al. (2014) Nature Methods 11(6):603-4

KSEA.Barplot

The KSEA App Analysis (KSEA Bar Plot Only)

Description

Takes a formatted phoshoproteomics data input and returns just the summary bar plot of kinase scores

Usage

```
KSEA.Barplot(KSData, PX, NetworKIN, NetworKIN.cutoff, m.cutoff, p.cutoff,
   export)
```

Arguments

KSData the Kinase-Substrate dataset uploaded from the file prefaced with "PSP&NetworKIN_"

available from github.com/casecpb/KSEA/

PX the experimental data file formatted as described in the KSEA.Complete() doc-

umentation

NetworkIN a binary input of TRUE or FALSE, indicating whether or not to include Net-

worKIN predictions; NetworKIN = TRUE means inclusion of NetworKIN pre-

dictions

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m.cutoff

| a numeric value between 1 and infinity setting the minimum NetworKIN score |
|--|
| (can be left out if NetworKIN = FALSE) |
| a numeric value between 0 and infinity indicating the min. # of substrates a |
| kinase must have to be included in the bar plot output |

p.cutoff a numeric value between 0 and 1 indicating the p-value cutoff for indicating

significant kinases in the bar plot

export a binary input of TRUE or FALSE, indicating whether or not to export the bar

plot as a .tiff image into the working directory

Value

creates the bar plot output highlighting key kinase results

References

```
Casado et al. (2013) Sci Signal. 6(268):rs6
Hornbeck et al. (2015) Nucleic Acids Res. 43:D512-20
Horn et al. (2014) Nature Methods 11(6):603-4
```

Examples

KSEA.Complete

The Complete KSEA App Analysis

Description

Takes a formatted phoshoproteomics data input and performs KSEA calculations to infer relative kinase activities

Usage

```
KSEA.Complete(KSData, PX, NetworKIN, NetworKIN.cutoff, m.cutoff, p.cutoff)
```

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Arguments

KSData

the Kinase-Substrate dataset uploaded from the file prefaced with "PSP&NetworKIN_" available from github.com/casecpb/KSEA/

PX

the experimental data file formatted exactly as described below; must have 6 columns in the exact order: Protein, Gene, Peptide, Residue.Both, p, FC; cannot have NA values, or else the entire peptide row is deleted; Description of each column in PX:

- "Protein" the Uniprot ID for the parent protein
- "Gene" the HUGO gene name for the parent protein
- "Peptide" the peptide sequence
- "Residue.Both" all phosphosites from that peptide, separated by semicolons if applicable; must be formatted as the single amino acid abbrev. with the residue position (e.g. S102)
- "p" the p-value of that peptide (if none calculated, please write "NULL", cannot be NA)
- "FC" the fold change (not log-transformed); usually the control sample is the denominator

NetworKIN

a binary input of TRUE or FALSE, indicating whether or not to include NetworKIN predictions; NetworKIN = TRUE means inclusion of NetworKIN predictions

NetworKIN.cutoff

a numeric value between 1 and infinity setting the minimum NetworKIN score (can be left out if NetworKIN = FALSE)

m.cutoff

a numeric value between 0 and infinity indicating the min. # of substrates a kinase must have to be included in the bar plot output

p.cutoff

a numeric value between 0 and 1 indicating the p-value cutoff for indicating significant kinases in the bar plot

Value

creates the following outputs that are deposited into your working directory: a bar plot highlighting key kinase results, a .csv file of all KSEA kinase scores, and a .csv file listing all kinase-substrate relationships used for the calculations

References

Casado et al. (2013) Sci Signal. 6(268):rs6 Hornbeck et al. (2015) Nucleic Acids Res. 43:D512-20 Horn et al. (2014) Nature Methods 11(6):603-4

Examples

```
KSEA.Complete(KSData, PX, NetworKIN=TRUE, NetworKIN.cutoff=5, m.cutoff=5, p.cutoff=0.01) KSEA.Complete(KSData, PX, NetworKIN=FALSE, m.cutoff=2, p.cutoff=0.05)
```

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| Notified that The Robert The Robe | KSEA.Heatmap | The KSEA App Analysis (KSEA Heatmap Only) | |
|--|--------------|---|--|
|--|--------------|---|--|

Description

Takes a list of the KSEA kinase score outputs from KSEA.Scores() and creates a merged heatmap (only applicable for multi-treatment studies)

Usage

```
KSEA.Heatmap(score.list, sample.labels, stats, m.cutoff, p.cutoff, sample.cluster)
```

Arguments

| score.list | the data frame outputs from the KSEA. Scores() function, compiled in a list format $$ |
|----------------|---|
| sample.labels | a character vector of all the sample names for heatmap annotation; the names must be in the same order as the data in score.list; please avoid long names, as they may get cropped in the final image |
| stats | character string of either "p.value" or "FDR" indicating the data column to use for marking statistically significant scores $\frac{1}{2}$ |
| m.cutoff | a numeric value between 0 and infinity indicating the min. # of substrates a kinase must have to be included in the heatmap |
| p.cutoff | a numeric value between 0 and 1 indicating the p-value/FDR cutoff for indicating significant kinases in the heatmap |
| sample.cluster | a binary input of TRUE or FALSE, indicating whether or not to perform hierarchical clustering of the sample columns |

Value

exports a .png heatmap image highlighting the merged datasets; heatmap was generated using the heatmap.2() function (gplots package); asterisks mark scores that met the statistical cutoff, as defined by p.cutoff; blue color indicates negative kinase score, and red indicates positive kinase score

References

```
Casado et al. (2013) Sci Signal. 6(268):rs6
Hornbeck et al. (2015) Nucleic Acids Res. 43:D512-20
Horn et al. (2014) Nature Methods 11(6):603-4
```

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Examples

stats="p.value", m.cutoff=3, p.cutoff=0.05, sample.cluster=TRUE)

KSEA.KS_table

The KSEA App Analysis (K-S Dataset Only)

Description

Takes a formatted phoshoproteomics data input and returns just the kinase-subtrate (K-S) annotations used for KSEA calculations

Usage

```
KSEA.KS_table(KSData, PX, NetworKIN, NetworKIN.cutoff)
```

Arguments

KSData the Kinase-Substrate dataset uploaded from the file prefaced with "PSP&NetworKIN_"

available from github.com/casecpb/KSEA/

PX the experimental data file formatted as described in the KSEA.Complete() doc-

umentation

NetworkIN a binary input of TRUE or FALSE, indicating whether or not to include Net-

worKIN predictions; NetworKIN = TRUE means inclusion of NetworKIN pre-

dictions

NetworKIN.cutoff

a numeric value between 1 and infinity setting the minimum NetworKIN score

(can be left out if NetworKIN = FALSE)

Value

creates a new data frame in R with all kinase-substrate relationships used for the KSEA calculations

References

Casado et al. (2013) Sci Signal. 6(268):rs6

Hornbeck et al. (2015) Nucleic Acids Res. 43:D512-20

Horn et al. (2014) Nature Methods 11(6):603-4

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Examples

```
KSData.dataset = KSEA.KS_table(KSData, PX, NetworKIN=TRUE, NetworKIN.cutoff=3)
KSData.dataset = KSEA.KS_table(KSData, PX, NetworKIN=FALSE)
```

KSEA.Scores

The KSEA App Analysis (KSEA Kinase Scores Only)

Description

Takes a formatted phoshoproteomics data input and returns just the KSEA kinase scores and statistics

Usage

```
KSEA.Scores(KSData, PX, NetworKIN, NetworKIN.cutoff)
```

Arguments

KSData the Kinase-Substrate dataset uploaded from the file prefaced with "PSP&NetworKIN_"

available from github.com/casecpb/KSEA/

PX the experimental data file formatted as described in the KSEA.Complete() doc-

umentation

NetworkIN a binary input of TRUE or FALSE, indicating whether or not to include Net-

worKIN predictions; NetworKIN = TRUE means inclusion of NetworKIN pre-

dictions

NetworKIN.cutoff

a numeric value between 1 and infinity setting the minimum NetworKIN score

(can be left out if NetworKIN = FALSE)

Value

creates a new data frame in R with all the KSEA kinase scores, along with each one's statistical assessment

References

```
Casado et al. (2013) Sci Signal. 6(268):rs6
Hornbeck et al. (2015) Nucleic Acids Res. 43:D512-20
Horn et al. (2014) Nature Methods 11(6):603-4
```

Examples

```
scores = KSEA.Scores(KSData, PX, NetworKIN=TRUE, NetworKIN.cutoff=3)
scores = KSEA.Scores(KSData, PX, NetworKIN=FALSE)
```

KSEA.Scores.2

KSEA.Scores.1

One of the 3 datasets for heatmap plotting

Description

A sample KSEA.Scores output generated from the KSEA.Scores() function (or alternatively, the "KSEA Kinase Scores.csv" output from the KSEA.Complete() function, loaded into R)

Usage

```
data(KSEA.Scores.1)
```

Format

dataframe containing 7 columns in the exact order as listed below.

- "KinaseGene" the HUGO gene name of the kinase
- "mS" the mean log2FC of all the kinase's identified substrates
- "Enrichment" the enrichment score (refer to Casado et al. (2013) Sci. Signal., 6, rs6-rs6)
- "m" the number of experimentally-identifed substrates annotating to that kinase
- "z.score" the normalized kinase score
- "p.value" the statistical assessment of the kinase score
- "FDR" the p-value adjusted for multiple hypothesis testing by the Benjamin-Hochberg method

References

unpublished data

KSEA.Scores.2

One of the 3 datasets for heatmap plotting

Description

A sample KSEA.Scores output generated from the KSEA.Scores() function (or alternatively, the "KSEA Kinase Scores.csv" output from the KSEA.Complete() function, loaded into R)

Usage

```
data(KSEA.Scores.2)
```

KSEA.Scores.3

Format

dataframe containing 7 columns in the exact order as listed below.

- "KinaseGene" the HUGO gene name of the kinase
- "mS" the mean log2FC of all the kinase's identified substrates
- "Enrichment" the enrichment score (refer to Casado et al. (2013) Sci. Signal., 6, rs6-rs6)
- "m" the number of experimentally-identifed substrates annotating to that kinase
- "z.score" the normalized kinase score
- "p.value" the statistical assessment of the kinase score
- "FDR" the p-value adjusted for multiple hypothesis testing by the Benjamin-Hochberg method

References

unpublished data

KSEA.Scores.3

One of the 3 datasets for heatmap plotting

Description

A sample KSEA.Scores output generated from the KSEA.Scores() function (or alternatively, the "KSEA Kinase Scores.csv" output from the KSEA.Complete() function, loaded into R)

Usage

```
data(KSEA.Scores.3)
```

Format

dataframe containing 7 columns in the exact order as listed below.

- "KinaseGene" the HUGO gene name of the kinase
- "mS" the mean log2FC of all the kinase's identified substrates
- "Enrichment" the enrichment score (refer to Casado et al. (2013) Sci. Signal., 6, rs6-rs6)
- "m" the number of experimentally-identifed substrates annotating to that kinase
- "z.score" the normalized kinase score
- "p.value" the statistical assessment of the kinase score
- "FDR" the p-value adjusted for multiple hypothesis testing by the Benjamin-Hochberg method

References

unpublished data

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PX

PX dataset for KSEA calculations

Description

A sample PX dataset of the experimental phosphoproteomics input

Usage

data(PX)

Format

the experimental data file must be formatted exactly as described below; must have 6 columns in the exact order: Protein, Gene, Peptide, Residue.Both, p, FC; cannot have NA values, or else the entire peptide row is deleted; Description of each column in PX:

- "Protein" the Uniprot ID for the parent protein
- "Gene" the HUGO gene name for the parent protein
- "Peptide" the peptide sequence
- "Residue.Both" all phosphosites from that peptide, separated by semicolons if applicable; must be formatted as the single amino acid abbrev. with the residue position (e.g. S102)
- "p" the p-value of that peptide (if none calculated, please write "NULL", cannot be NA)
- "FC" the fold change (not log-transformed); usually the control sample is the denominator

References

unpublished data

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