# Package 'eSEM'

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Type Package
Title Enzyme activity inference from Structural Equation Modeling
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<b>Imports</b> lavaan(>= 0.6-11), tidyr(>= 1.2.0), stringr(>= 1.4.0), dplyr(>= 1.0.9), psych(>= 2.2.5), devtools(>= 2.4.5), EFAtools(>= 0.4.1)
<b>Description</b> eSEM is used to infer enzyme activity.
<b>Depends</b> R (>= $3.5.0$ )
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R topics documented:
aceAdjacency catchActRaw eSEM getFa inputTransformPSP mastInput motifEx pspAdjacency pspMotifAdjacency singleEnzymeAff singleHatAct singleKinaseAct singleLigaseAct ubiAdjacency
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eSEM

aceAdjacency

aceAdjacency

#### **Description**

This script is used to build HUMAN HAT-substrate adjacency matrix.

#### Usage

```
aceAdjacency()
```

catchActRaw

catchResults

#### **Description**

Catch Activity and Affinity. Activity contains Raw-Activity, Mean-Center, Z-scaled.

#### Usage

```
catchActRaw(x)
```

## Arguments

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The results from modeling fitting.

rawData

Input raw data (quantitative phospho-proteomics / ubiqutin-proteomics data)

eSEM

eSEM

#### **Description**

Infer enzyme activity from phosphoralation, ubiquitination, acetylation

## Usage

```
eSEM(
  input = NULL,
  datatype = NULL,
  organism = NULL,
  enzyme.organism = NULL,
  cor.off = 0.95,
  kmo.off = 0,
  enzyList = NULL,
  input.log2.norm = FALSE,
  whole.log2.trans = FALSE,
  whole.proteome = NULL,
  motif = NULL,
  output.folder = getwd()
)
```

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#### **Arguments**

input Dataframe. Your quantitative ubiquiti-proteomics, phospho-proteomic and acetyl-

proteomics data.

datatype Factor. Data type: ubiquitination or phosphoralation or acetylation; ("ubi",

"psp", "ace")

organism Vector. Substrate species; ("human", "mouse", "rat")

enzyme.organism

Vector. Enzyme species; ("human", "mouse", "rat")

cor.off Numeric. Set up correlation cutoff value 0-1 to remove high collinear variables.

Default is 0.95.

kmo.off Numeric. Set up KMO cutoff value 0-1. Default is 0.

enzyList Vector. Program only calculate the enzyme in the enzyList. Default is to output

ALL enzyme activities and affinities.

input.log2.norm

Logical. FALSE or TRUE. Need program to do log2 transforming of the input

file or not. Default is FALSE.

whole.log2.trans

Logical. FALSE or TRUE. Need program to do log2 transforming of the whole

proteome or not. (Ignore if -whole.proteome is missing). Default is FALSE.

whole.proteome Dataframe. Set up whole proteome used to normalize phosphor-proteome or

ubiquitin-proteome whole proteome. Default is NULL.

motif Matrix. Added kinase-subatrate relationships from motif discovery.

output.folder Character. Character vector of location to save files if desired. Default is current

directory.

#### **Details**

You can use eSEM package to get enzyme activity.

#### Value

List

#### **Examples**

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```
datatype = "ace",
organism = "human",
input.log2.norm = FALSE)
```

getFa

Build MEME input

## Description

script to extract MEME input .fa file

## Usage

```
getFa(ss = NULL, es = NULL, filePath)
```

## **Arguments**

ss Vector; Substrate species; ("mouse", "rat", "human")
es Vector; Kinase species; ("mouse", "rat", "human")

filePath The path is used to save .fa files. Recommend to create a new folder to save all

.fa.files.

#### **Examples**

```
getFa(ss = "human", es = c("human", "mouse", "rat"), filePath = getwd())
```

input Transform PSP

input Transform

#### **Description**

This script is used to log2 transform AND/OR whole proteome normalize input data.

#### Usage

```
inputTransformPSP(
  input_raw,
  input_log2_trans = FALSE,
  whole_log2_trans = FALSE,
  whole_proteome = NULL
)
```

mastInput 5

#### **Arguments**

input\_raw Input raw data (quantitative phospho-proteomics/ubiquitin-proteomics data) input\_log2\_trans

> FALSE or TRUE. Need program to do log2 transforming of the input file or not. Default is FALSE.

whole\_log2\_trans

Need program to do log2 transforming of the whole proteome or not. Ignore if -whole.proteome is missing. Default is FALSE.

whole\_proteome Set up whole proteome used to normalize phospho-proteomics/ubiquitin-proteomics data by whole proteome. Default is NULL.

mastInput

Build MAST input

#### **Description**

Script to extract MAST input FASTA file.

#### Usage

```
mastInput(input, savePath = getwd())
```

#### **Arguments**

input The same quantitative phosphoproteomics data as eSEM() input file. The path to save built MAST input file. Default is the current directory. savePath

#### **Examples**

```
mastInput(input = input_psp_example)
```

motifEx

motifEx

#### **Description**

This script is used to extract kinase-substrate relationships from motif scanning and build motif\_input file for KSEM().

## Usage

```
motifEx(mastFile, savePath = getwd())
```

#### **Arguments**

mastFile The output file "extract\_result.txt" from Step: 03step\_ExtractMASTresult.sh. savePath Path to save motif\_input.txt file. Default is current directory.

6 pspMotifAdjacency

## Description

This script is used to build kinase-substrate adjacency matrix with only PSP database.

## Usage

```
pspAdjacency(sp, ep)
```

## Arguments

sp	Substrate species; ("mouse", "rat", "human").
ер	Enzyme species: ("mouse", "rat", "human").

pspMotifAdjacency pspMotifAdjacency

## Description

This script is used to build enzyme-substrate adjacency matrix with PSP database + Motif.

## Usage

```
pspMotifAdjacency(sp, ep, motif.ref)
```

## **Arguments**

sp	Substrate species; ("mouse", "rat", "human").
ер	Enzyme species; ("mouse", "rat", "human").
motif.ref	Added enzyme-subatrate relationships from motif discovery. Same as KSEM -motif parameter.

singleEnzymeAff 7

## Description

Get single enzyme affinity.

## Usage

```
singleEnzymeAff(enzyme, input, adj, cor.off, kmo.off)
```

## Arguments

enzyme	Single enzyme name.
input	Normalized and transformed phospho-proteomics / ubiqutin-proteomics data.
adj	Adjacency matrix.
cor.off	Set up correlation cutoff value 0-1 to remove high collinear variables. Default is 0.95.
kmo.off	Set up KMO cutoff value 0-1. Default is 0.

singleHatAct	singleHatAct	

## Description

Get single HAT activity.

## Usage

```
singleHatAct(HAT, input, adj, cor.off, kmo.off)
```

## Arguments

HAT	Single HAT name.
input	Normalized and transformed acetyl-proteomics data.
adj	Adjacency matrix.
cor.off	Set up correlation cutoff value 0-1 to remove high collinear variables. Default is 0.95.
kmo.off	Set up KMO cutoff value 0-1. Default is 0.

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singleKinaseAct		
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#### **Description**

Get single kinase activity.

#### Usage

```
singleKinaseAct(kinase, input, adj, cor.off, kmo.off)
```

#### **Arguments**

kinase Single kinase name.

input Normalized and transformed phosphoproteomics data.

adj Adjacency matrix.

cor.off Set up correlation cutoff value 0-1 to remove high collinear variables. Default is

0.95.

kmo.off Set up KMO cutoff value 0-1. Default is 0.

singleLigaseAct singleLigaseAct

## Description

Get single ligase activity.

## Usage

```
singleLigaseAct(ligase, input, adj, cor.off, kmo.off)
```

## Arguments

input Normalized and transformed ubiquiti-proteomics data.

adj Adjacency matrix.

cor.off Set up correlation cutoff value 0-1 to remove high collinear variables. Default is

0.95.

kmo.off Set up KMO cutoff value 0-1. Default is 0.

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ubiAdjacency

ubiAdjacency

## Description

This script is used to build ligase-substrate adjacency matrix with only PSP database.

## Usage

```
ubiAdjacency(sp)
```

## Arguments

sp

Substrate species; ("mouse", "rat", "human").