# Package 'survival666'

November 29, 2021

Version 0.5
Title Eliminate the Influence of Co-Expression Genes on Target Genes
Description Functions can be used for batch survival analysis, but not only for it. Most importantly, it can verify any P-value calculated according to the gene expression level and eliminate the influence of co-expression genes.
<b>Depends</b> R (>= $3.3$ )
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Suggests tidyverse, magrittr, testthat (>= 3.0.0)
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clinic	Clinical records of overall survival	

# Description

This is a document that records the overall survival rate.

# Usage

clinic

## **Format**

An object of class data. frame with 454 rows and 3 columns.

# Source

https://xenabrowser.net/datapages/

|--|

# Description

The super\_survival is the frist function of package survival666. You can use the function to perform batch operations on the gene expression matrix to obtain the survival analysis of the entire transcriptome

# Usage

```
super_survival(exp, time, status, p = 1e-04, title = "survival_rank", path)
```

# Arguments

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)

survivaldata 3

## **Examples**

```
## Not run: super_survival(exp=survivaldata,
time = clinic[,3],
status = clinic[,2],
p = 0.0001,
title = 'survival_rank',
path = '/yourdir/')
## End(Not run)
```

survivaldata

A filtered transcriptome gene expression matrix.

# **Description**

The expression matrix was manipulated to contain less than two thousand genes. Use the symbol as the column name and the sample number as the row name. The reason of this is that to save space.It is best to use the complete transcriptome as the expression matrix when using the fundtion super\_survival.

# Usage

survivaldata

### **Format**

An object of class data. frame with 454 rows and 1712 columns.

#### **Source**

https://xenabrowser.net/datapages/

survivalrank

Rank the significance of survival analysis

# **Description**

This data file is actually the output of super\_survival function. The significance P value of survival analysis for each gene in survival data matrix were recorded

## Usage

survivalrank

#### **Format**

An object of class data. frame with 1712 rows and 9 columns.

4 survival\_pie

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Visualize the factors that influence survival analysis

# **Description**

The checkout\_survival() is the third function of survival666. We use the function to visualize the co-expression interference of a specific gene in survival analysis.

#### Usage

```
survival_pie(exp, survivalrank, target, pcol = 4, symbolcol = 1, path)
```

## **Arguments**

exp This is a transcriptome count matrix, Note that the matrix requires symbol as

column name and sample as row name

survivalrank This is the result calculated by the frist function in the survival666 package.

However, that does not prevent you from getting results that you could get using other methods, as long as three conditions are met. Frist, you need to calculate the p value based on the expression of a particular phenotype and the level of gene expression Second, your results must be large enough, preferably for the entire transcriptome, otherwise filtering accuracy will be reduced Finally, you need to eliminate genes that are not significant enough, and if you are using a regular

PC, it is recommended to have no more than 2000 lines

target gene symbols

pcol Digtal vector,represents the column of P value in survivalrank symbolcol Digtal vector,represents the column of symbol in survivalrank

Path to output file, do not omit '/', example: 'd:/file/'.

# See Also

```
survival_pie super_survival
```

# **Examples**

```
## Not run: survival_pie(exp=survivaldata,
survivalrank=survivalrank[1:300,],
target="EAF2",
pcol=4,
symbolcol=1,
path='/yourdir/')
## End(Not run)
```

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