# Package 'progeny'

April 16, 2019

April 10, 2017	
<b>Title</b> Pathway RespOnsive GENes for activity inference from gene expression	
Version 1.4.1	
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<b>Description</b> This package provides a function to infer pathway activity from gene expression using PROGENy. It contains the linear model we inferred in the publication ``Perturbation-response genes reveal signaling footprints in cancer gene expression".	
<pre>URL https://github.com/saezlab/progeny</pre>	
<pre>BugReports https://github.com/saezlab/progeny/issues</pre>	
<b>Depends</b> R (>= 3.4.0)	
Imports Biobase	
<b>biocViews</b> SystemsBiology, GeneExpression, FunctionalPrediction, GeneRegulation	
License Apache License (== 2.0)   file LICENSE	
LazyData true	
Encoding UTF-8	
<b>Suggests</b> airway, biomaRt, BiocFileCache, broom, DESeq2, dplyr, knitr, readr, readxl	
VignetteBuilder knitr	
RoxygenNote 6.0.1	
git_url https://git.bioconductor.org/packages/progeny	
git_branch RELEASE_3_8	
git_last_commit e5551a6	
git_last_commit_date 2019-04-10	
Date/Publication 2019-04-15	
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model

The linear model underlying PROGENy

# Description

HGNC gene symbols in rows, pathways in columns. Pathway activity inference works by a matrix multiplication of gene expression with the model.

### Usage

model

#### **Format**

An object of class matrix with 1059 rows and 11 columns.

#### **Source**

http://biorxiv.org/content/early/2016/08/28/065672

progeny

Calculate PROGENy pathway scores from gene expression

# **Description**

This function uses the linear model of pathway-responsive genes underlying the PROGENy method. It transforms a gene expression matrix with HGNC gene symbols in rows and sample names in columns into a pathway score matrix with samples and in rows and pathways in columns.

This function uses the linear model of pathway-responsive genes underlying the PROGENy method. It transforms a gene expression matrix with HGNC gene symbols in rows and sample names in columns into a pathway score matrix with samples and in rows and pathways in columns.

# Usage

```
progeny(expr, scale = TRUE)
```

## **Arguments**

expr A gene expression object with HGNC symbols in rows and samples in columns

scale Logical value indicating whether to scale the scores of each pathway to have a

mean of zero and standard deviation of one

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#### **Details**

The publication of the method is available at: https://www.biorxiv.org/content/early/2016/08/28/065672

The supplied expression object has to contain HGNC symbols in rows. This will, in most cases (and how we originally used it), be either normalized gene expression of a microarray experiment or log-transformed (and possible variance-stabilized) counts from an RNA-seq experiment.

The model matrix itself consists of 11 pathways and 1059 genes. Its coefficients are non-zero if the gene-pathway pair corresponds to the top 100 genes that were up-regulated upon stimulation of the pathway in a wide range of experiments. The value corresponds to the fitted z-score across experiments in our model fit. Only rows with at least one non-zero coefficient were included, as the rest is not used to infer pathway activity.

### Value

A matrix with samples in columns and pathways in rows

## **Examples**

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