

Package ‘decoupleR’

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Type Package

Title Package to decouple gene sets from statistics

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Description Transcriptome profiling followed by differential gene expression analysis often leads to lists of genes that are hard to analyze and interpret. Downstream analysis tools can be used to summarize deregulation events into a smaller set of biologically interpretable features. In particular, methods that estimate the activity of transcription factors (TFs) from gene expression are commonly used. It has been shown that the transcriptional targets of a TF yield a much more robust estimation of the TF activity than observing the expression of the TF itself. Consequently, for the estimation of transcription factor activities, a network of transcriptional regulation is required in combination with a statistical algorithm that summarizes the expression of the target genes into a single activity score. Over the years, many different regulatory networks and statistical algorithms have been developed, mostly in a fixed combination of one network and one algorithm. To systematically evaluate both networks and algorithms, we developed decoupleR , an R package that allows users to apply efficiently any combination provided.

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URL <https://saezlab.github.io/decoupleR/>

BugReports <https://github.com/saezlab/decoupleR/issues>

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R topics documented:

convert_f_defaults	2
convert_to_	4
decouple	5
filter_regulons	7
run_gsva	7
run_mean	8
run_ora	10
run_pscira	12
run_scira	13
run_viper	15
Index	17

convert_f_defaults	<i>Rename columns and add defaults values if column not present</i>
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Description

convert_f_defaults() combine the `dplyr::rename()` way of working and with the `tibble::add_column()` to add columns with default values in case they don't exist after renaming data.

Usage

```
convert_f_defaults(.data, ..., .def_col_val = c(), .use_dots = TRUE)
```

Arguments

<code>.data</code>	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from dbplyr or dtplyr). See <i>Methods</i> , below, for more details.
<code>...</code>	For <code>rename()</code> : <tidy-select> Use <code>new_name = old_name</code> to rename selected variables. For <code>rename_with()</code> : additional arguments passed onto <code>.fn</code> .
<code>.def_col_val</code>	Named vector with columns with default values if none exist after rename.
<code>.use_dots</code>	Should a dot prefix be added to renamed variables? This will allow swapping of columns.

Details

The objective of using `.use_dots` is to be able to swap columns which, by default, is not allowed by the `dplyr::rename()` function. The same behavior can be replicated by simply using the `dplyr::select()`, however, the select evaluation allows much more flexibility so that unexpected results could be obtained. Despite this, a future implementation will consider this form of execution to allow renaming the same column to multiple ones (i.e. extend dataframe extension).

Value

An object of the same type as `.data`. The output has the following properties:

- Rows are not affected.
- Column names are changed.
- Column order is the same as that of the function call.

Examples

```
df <- tibble::tibble(x = 1, y = 2, z = 3)

# Rename columns
df <- tibble::tibble(x = 1, y = 2)
convert_f_defaults(
  .data = df,
  new_x = x,
  new_y = y,
  new_z = NULL,
  .def_col_val = c(new_z = 3)
)
```

convert_to_	<i>Convert a network to run under the method of interest.</i>
-------------	---

Description

Convert a long-format network to the suggested standard for the specified `run_{statistic}()`. If the default parameters are not modified, then the function sets its own null values for those columns.

Usage

```
convert_to_(network)

convert_to_scira(network, .source, .target, .mor = NULL)

convert_to_pscira(network, .source, .target, .mor = NULL)

convert_to_mean(network, .source, .target, .mor = NULL, .likelihood = NULL)

convert_to_viper(network, .source, .target, .mor = NULL, .likelihood = NULL)

convert_to_gsva(network, .source, .target)

convert_to_ora(network, .source, .target)
```

Arguments

<code>network</code>	Tibble or dataframe with edges and it's associated metadata.
<code>.source</code>	Column with source nodes.
<code>.target</code>	Column with target nodes.
<code>.mor</code>	Column with edge mode of regulation (i.e. mor).
<code>.likelihood</code>	Column with edge likelihood.

Value

- `convert_to_` Return same as input.
- `convert_to_gsva()` Return a list of regulons suitable for [GSVA::gsva\(\)](#).
- `convert_to_mean()` Return a tibble with four columns: `tf`, `target`, `mor` and `likelihood`.
- `convert_to_ora()` Return a named list of regulons; `tf` with associated targets.
- `convert_to_pscira()` Returns a tibble with three columns: `tf`, `target` and `mor`.
- `convert_to_scira()` Returns a tibble with three columns: `tf`, `target` and `mor`.
- `convert_to_viper()` Return a list of regulons suitable for [viper::viper\(\)](#)

See Also

[convert_f_defaults\(\)](#)

Examples

```
inputs_dir <- system.file("testdata", "inputs", package = "decoupleR")

network <- readRDS(file.path(inputs_dir, "input-dorothea_genesets.rds"))

convert_to_(network)
convert_to_gsva(network, tf, target)
convert_to_mean(network, tf, target, mor, likelihood)
convert_to_ora(network, tf, target)
convert_to_pscira(network, tf, target, mor)
convert_to_scira(network, tf, target, mor)
convert_to_viper(network, tf, target, mor, likelihood)
```

decouple

Evaluate multiple statistics with same input data

Description

Calculate the TF activity per sample out of a gene expression matrix by coupling a regulon network with a variety of statistics.

Usage

```
decouple(
  mat,
  network,
  .source,
  .target,
  statistics,
  args = list(NULL),
  include_time = FALSE,
  show_toy_call = FALSE
)
```

Arguments

<code>mat</code>	Matrix to evaluate (e.g. expression matrix). Target nodes in rows and conditions in columns. <code>rownames(mat)</code> must have at least one intersection with the elements in <code>network .target</code> column.
<code>network</code>	Tibble or dataframe with edges and it's associated metadata.
<code>.source</code>	Column with source nodes.
<code>.target</code>	Column with target nodes.
<code>statistics</code>	Statistical methods to be coupled.
<code>args</code>	A list of argument-lists the same length as <code>statistics</code> (or length 1). The default argument, <code>list(NULL)</code> , will be recycled to the same length as <code>statistics</code> , and will call each function with no arguments (apart from <code>mat</code> , <code>network</code> , <code>.source</code> and, <code>.target</code>).

include_time Should the time per statistic evaluated be informed?
 show_toy_call The call of each statistic must be informed?

Value

A long format tibble of the enrichment scores for each tf across the samples. Resulting tibble contains the following columns:

1. statistic: Indicates which method is associated with which score.
2. tf: Source nodes of network.
3. condition: Condition representing each column of mat.
4. score: Regulatory activity (enrichment score).
5. statistic_time: If requested, internal execution time indicator.
6. ...: Columns of metadata generated by certain statistics.

See Also

Other decoupleR statistics: [run_gsva\(\)](#), [run_mean\(\)](#), [run_ora\(\)](#), [run_pscira\(\)](#), [run_scira\(\)](#), [run_viper\(\)](#)

Examples

```
if (FALSE) {
  inputs_dir <- system.file("testdata", "inputs", package = "decoupleR")

  mat <- readRDS(file.path(inputs_dir, "input-expr_matrix.rds"))
  network <- readRDS(file.path(inputs_dir, "input-dorothea_genesets.rds"))

  decouple(
    mat = mat,
    network = network,
    .source = "tf",
    .target = "target",
    statistics = c("gsva", "mean", "pscira", "scira", "viper"),
    args = list(
      gsva = list(verbose = FALSE),
      mean = list(.mor = "mor", .likelihood = "likelihood"),
      pscira = list(.mor = "mor"),
      scira = list(.mor = "mor"),
      viper = list(
        .mor = "mor",
        .likelihood = "likelihood",
        verbose = FALSE
      )
    )
  )
}
```

filter_regulons	<i>Filter network by size of regulons</i>
-----------------	---

Description

Keep only sources which satisfied the condition $\text{min_size} \geq n \leq \text{max_size}$, where n denotes the number of targets per source.

Usage

```
filter_regulons(network, .source, min_size = 1, max_size = Inf)
```

Arguments

network	Tibble or dataframe with edges and it's associated metadata.
.source	Column with source nodes.
min_size	Minimum number of targets allowed per regulon.
max_size	Maximum number of targets allowed per regulon.

Value

Filtered tibble.

Examples

```
inputs_dir <- system.file("testdata", "inputs", package = "decoupleR")
network <- readRDS(file.path(inputs_dir, "input-dorothea_genesets.rds"))
filter_regulons(network, .source = tf, min_size = 30, max_size = 50)
```

run_gsva	<i>Gsva wrapper</i>
----------	---------------------

Description

This function is a convenient wrapper for the [Gsva::gsva\(\)](#) function.

Usage

```
run_gsva(mat, network, .source = .data$tf, .target = .data$target, ...)
```

Arguments

mat	Matrix to evaluate (e.g. expression matrix). Target nodes in rows and conditions in columns. rownames(mat) must have at least one intersection with the elements in network .target column.
network	Tibble or dataframe with edges and it's associated metadata.
.source	Column with source nodes.
.target	Column with target nodes.
...	Arguments passed on to GSVA: :gsva

Value

A long format tibble of the enrichment scores for each tf across the samples. Resulting tibble contains the following columns:

- 1. statistic: Indicates which method is associated with which score.
- 2. tf: Source nodes of network.
- 3. condition: Condition representing each column of mat.
- 4. score: Regulatory activity (enrichment score).

See Also

Other decoupleR statistics: [decouple\(\)](#), [run_mean\(\)](#), [run_oracle\(\)](#), [run_pscira\(\)](#), [run_scira\(\)](#), [run_viper\(\)](#)

Examples

```
inputs_dir <- system.file("testdata", "inputs", package = "decoupleR")

mat <- readRDS(file.path(inputs_dir, "input-expr_matrix.rds"))
network <- readRDS(file.path(inputs_dir, "input-dorothea_genesets.rds"))

run_gsva(mat, network, tf, target, verbose = FALSE)
```

run_mean	<i>Weighted mean</i>
----------	----------------------

Description

Calculate the activity of all regulons in network through the conditions in the mat matrix by calculating the mean over the expression of all genes.

Usage

```
run_mean(
  mat,
  network,
  .source = .data$tf,
  .target = .data$target,
  .mor = .data$mor,
  .likelihood = .data$likelihood,
  times = 2,
  seed = 42,
  sparse = TRUE,
  randomize_type = "rows"
)
```

Arguments

mat	Matrix to evaluate (e.g. expression matrix). Target nodes in rows and conditions in columns. rownames(mat) must have at least one intersection with the elements in network .target column.
network	Tibble or dataframe with edges and it's associated metadata.
.source	Column with source nodes.
.target	Column with target nodes.
.mor	Column with edge mode of regulation (i.e. mor).
.likelihood	Column with edge likelihood.
times	How many permutations to do?
seed	A single value, interpreted as an integer, or NULL for random number generation.
sparse	Should the matrices used for the calculation be sparse?
randomize_type	How to randomize the expression matrix.

Details

run_mean() calculates the activity score, but in addition, it takes advantage of the permutations used to calculate the p-value, to provide the normalized activity score. This is represented in the statistic column which will contain two values for each call to run_mean(); **mean** and **normalized_mean**.

Value

A long format tibble of the enrichment scores for each tf across the samples. Resulting tibble contains the following columns:

1. statistic: Indicates which method is associated with which score.
2. tf: Source nodes of network.
3. condition: Condition representing each column of mat.
4. score: Regulatory activity (enrichment score).
5. p_value: p-value for the score of mean method.

See Also

Other decoupleR statistics: [decouple\(\)](#), [run_gsva\(\)](#), [run_ora\(\)](#), [run_pscira\(\)](#), [run_scira\(\)](#), [run_viper\(\)](#)

Examples

```
inputs_dir <- system.file("testdata", "inputs", package = "decoupleR")

mat <- readRDS(file.path(inputs_dir, "input-expr_matrix.rds"))
network <- readRDS(file.path(inputs_dir, "input-dorothea_genesets.rds"))

run_mean(mat, network, tf, target, mor, likelihood)
```

run_ora

*Over Representation Analysis - Fisher Exact Test***Description**

Performs an over-representation analysis using [stats::fisher.test\(\)](#).

Usage

```
run_ora(
  mat,
  network,
  .source = .data$tf,
  .target = .data$target,
  n_up = nrow(mat),
  n_bottom = 0,
  n_background = NULL,
  with_ties = TRUE,
  ...
)
```

Arguments

mat	Matrix to evaluate (e.g. expression matrix). Target nodes in rows and conditions in columns. <code>rownames(mat)</code> must have at least one intersection with the elements in <code>network .target</code> column.
network	Tibble or dataframe with edges and it's associated metadata.
.source	Column with source nodes.
.target	Column with target nodes.
n_up	Integer indicating the number of top targets to slice from mat.
n_bottom	Integer indicating the number of bottom targets to slice from mat.

n_background	Integer indicating the background size of the sliced targets. If not specified the number of background targets is determined by the total number of unique targets in the union of mat and network.
with_ties	Should ties be kept together? The default, TRUE, may return more rows than you request. Use FALSE to ignore ties, and return the first n rows.
...	Arguments passed on to stats::fisher.test
workspace	an integer specifying the size of the workspace used in the network algorithm. In units of 4 bytes. Only used for non-simulated p-values larger than 2×2 tables. Since R version 3.5.0, this also increases the internal stack size which allows larger problems to be solved, however sometimes needing hours. In such cases, <code>simulate.p.values=TRUE</code> may be more reasonable.
hybrid	a logical. Only used for larger than 2×2 tables, in which cases it indicates whether the exact probabilities (default) or a hybrid approximation thereof should be computed.
hybridPars	a numeric vector of length 3, by default describing “Cochran’s conditions” for the validity of the chisquare approximation, see ‘Details’.
control	a list with named components for low level algorithm control. At present the only one used is “mult”, a positive integer ≥ 2 with default 30 used only for larger than 2×2 tables. This says how many times as much space should be allocated to paths as to keys: see file ‘fexact.c’ in the sources of this package.
or	the hypothesized odds ratio. Only used in the 2×2 case.
alternative	indicates the alternative hypothesis and must be one of “two.sided”, “greater” or “less”. You can specify just the initial letter. Only used in the 2×2 case.
conf.int	logical indicating if a confidence interval for the odds ratio in a 2×2 table should be computed (and returned).
conf.level	confidence level for the returned confidence interval. Only used in the 2×2 case and if <code>conf.int = TRUE</code> .
simulate.p.value	a logical indicating whether to compute p-values by Monte Carlo simulation, in larger than 2×2 tables.
B	an integer specifying the number of replicates used in the Monte Carlo test.

Value

A long format tibble of the enrichment scores for each tf across the samples. Resulting tibble contains the following columns:

1. statistic: Indicates which method is associated with which score.
2. tf: Source nodes of network.
3. condition: Condition representing each column of mat.
4. score: Regulatory activity (enrichment score).

See Also

Other decoupleR statistics: [decouple\(\)](#), [run_gsva\(\)](#), [run_mean\(\)](#), [run_pscira\(\)](#), [run_scira\(\)](#), [run_viper\(\)](#)

Examples

```
inputs_dir <- system.file("testdata", "inputs", package = "decoupleR")

mat <- readRDS(file.path(inputs_dir, "input-expr_matrix.rds"))
network <- readRDS(file.path(inputs_dir, "input-dorothea_genesets.rds"))

run_ora(mat, network, tf, target)
```

run_pscira

PSCIRA (*Permutation Single Cell Inference of Regulatory Activity*)

Description

Calculate the regulatory activity of each tf by multiplying the expression values of its objectives with their corresponding associated profiles for each given condition. The result is equal to the z-score of the found value compared to its null distribution.

Usage

```
run_pscira(
  mat,
  network,
  .source = .data$tf,
  .target = .data$target,
  .mor = .data$mor,
  sparse = TRUE,
  times = 10,
  seed = 42
)
```

Arguments

mat	Matrix to evaluate (e.g. expression matrix). Target nodes in rows and conditions in columns. <code>rownames(mat)</code> must have at least one intersection with the elements in <code>network .target</code> column.
network	Tibble or dataframe with edges and it's associated metadata.
.source	Column with source nodes.
.target	Column with target nodes.
.mor	Column with edge mode of regulation (i.e. mor).
sparse	Logical value indicating if the generated profile matrix should be sparse.
times	Number of replications.
seed	A single value, interpreted as an integer, or NULL.

Details

Estimation of regulatory activity: A linear regression of the expression profile is performed against the "target profile" of the given TF, where in the target profile, any regulon member is assigned a +1 for activating interactions and a -1 for inhibitory interactions. All other genes not members of the TF's regulon are assigned a value of 0. TF activity is then defined as the t-statistic of this linear regression.

Value

A long format tibble of the enrichment scores for each tf across the samples. Resulting tibble contains the following columns:

1. statistic: Indicates which method is associated with which score.
2. tf: Source nodes of network.
3. condition: Condition representing each column of mat.
4. score: Regulatory activity (enrichment score).

See Also

Other decoupleR statistics: [decouple\(\)](#), [run_gsva\(\)](#), [run_mean\(\)](#), [run_oracle\(\)](#), [run_scira\(\)](#), [run_viper\(\)](#)

Examples

```
inputs_dir <- system.file("testdata", "inputs", package = "decoupleR")

mat <- readRDS(file.path(inputs_dir, "input-expr_matrix.rds"))
network <- readRDS(file.path(inputs_dir, "input-dorothea_genesets.rds"))

run_pscira(mat, network, tf, target, mor)
```

run_scira

SCIRA (Single Cell Inference of Regulatory Activity)

Description

Calculates TF activity according to [Improved detection of tumor suppressor events in single-cell RNA-Seq data](#).

Usage

```
run_scira(
  mat,
  network,
  .source = .data$tf,
  .target = .data$target,
  .mor = .data$mor,
```

```

    sparse = FALSE,
    fast = TRUE,
    center = TRUE,
    na.rm = FALSE
  )

```

Arguments

<code>mat</code>	Matrix to evaluate (e.g. expression matrix). Target nodes in rows and conditions in columns. <code>rownames(mat)</code> must have at least one intersection with the elements in <code>network</code> <code>.target</code> column.
<code>network</code>	Tibble or dataframe with edges and it's associated metadata.
<code>.source</code>	Column with source nodes.
<code>.target</code>	Column with target nodes.
<code>.mor</code>	Column with edge mode of regulation (i.e. mor).
<code>sparse</code>	Logical value indicating if the generated profile matrix should be sparse.
<code>fast</code>	Logical value indicating if the lineal model must be calculated with <code>speedglm::speedlm.fit()</code> or with base <code>stats::lm()</code> .
<code>center</code>	Logical value indicating if <code>mat</code> must be centered by <code>base::rowMeans()</code> .
<code>na.rm</code>	Should missing values (including NaN) be omitted from the calculations of <code>base::rowMeans()</code> ?

Details

Estimation of regulatory activity: A linear regression of the expression profile is performed against the "target profile" of the given TF, where in the target profile, any regulon member is assigned a +1 for activating interactions and a -1 for inhibitory interactions. All other genes not members of the TF's regulon are assigned a value of 0. TF activity is then defined as the t-statistic of this linear regression.

Value

A long format tibble of the enrichment scores for each tf across the samples. Resulting tibble contains the following columns:

1. `statistic`: Indicates which method is associated with which score.
2. `tf`: Source nodes of network.
3. `condition`: Condition representing each column of `mat`.
4. `score`: Regulatory activity (enrichment score).

See Also

Other decoupleR statistics: `decouple()`, `run_gsva()`, `run_mean()`, `run_oracle()`, `run_pscira()`, `run_viper()`

Examples

```
inputs_dir <- system.file("testdata", "inputs", package = "decoupleR")

mat <- readRDS(file.path(inputs_dir, "input-expr_matrix.rds"))
network <- readRDS(file.path(inputs_dir, "input-dorothea_genesets.rds"))

run_scira(mat, network, tf, target, mor)
```

run_viper

VIPER wrapper

Description

This function is a convenient wrapper for the `viper::viper()` function.

Usage

```
run_viper(
  mat,
  network,
  .source = .data$tf,
  .target = .data$target,
  .mor = .data$mor,
  .likelihood = .data$likelihood,
  ...
)
```

Arguments

<code>mat</code>	Matrix to evaluate (e.g. expression matrix). Target nodes in rows and conditions in columns. <code>rownames(mat)</code> must have at least one intersection with the elements in <code>network</code> <code>.target</code> column.
<code>network</code>	Tibble or dataframe with edges and it's associated metadata.
<code>.source</code>	Column with source nodes.
<code>.target</code>	Column with target nodes.
<code>.mor</code>	Column with edge mode of regulation (i.e. <code>mor</code>).
<code>.likelihood</code>	Column with edge likelihood.
<code>...</code>	Arguments passed on to <code>viper::viper</code>
<code>dnull</code>	Numeric matrix for the null model, usually generated by <code>nullTtest</code>
<code>pleiotropy</code>	Logical, whether correction for pleiotropic regulation should be performed
<code>nes</code>	Logical, whether the enrichment score reported should be normalized
<code>method</code>	Character string indicating the method for computing the single samples signature, either <code>scale</code> , <code>rank</code> , <code>mad</code> , <code>ttest</code> or <code>none</code>

bootstraps Integer indicating the number of bootstraps iterations to perform. Only the scale method is implemented with bootstraps.

adaptive.size Logical, whether the weighting scores should be taken into account for computing the regulon size

eset.filter Logical, whether the dataset should be limited only to the genes represented in the interactome #' @param mvws Number or vector indicating either the exponent score for the metaViper weights, or the inflection point and trend for the sigmoid function describing the weights in metaViper

pleiotropyArgs list of 5 numbers for the pleiotropy correction indicating: regulators p-value threshold, pleiotropic interaction p-value threshold, minimum number of targets in the overlap between pleiotropic regulators, penalty for the pleiotropic interactions and the method for computing the pleiotropy, either absolute or adaptive

cores Integer indicating the number of cores to use (only 1 in Windows-based systems)

verbose Logical, whether progression messages should be printed in the terminal

Value

A long format tibble of the enrichment scores for each tf across the samples. Resulting tibble contains the following columns:

1. **statistic**: Indicates which method is associated with which score.
2. **tf**: Source nodes of network.
3. **condition**: Condition representing each column of mat.
4. **score**: Regulatory activity (enrichment score).

See Also

Other decoupleR statistics: [decouple\(\)](#), [run_gsva\(\)](#), [run_mean\(\)](#), [run_oracle\(\)](#), [run_pscira\(\)](#), [run_scira\(\)](#)

Examples

```
inputs_dir <- system.file("testdata", "inputs", package = "decoupleR")

mat <- readRDS(file.path(inputs_dir, "input-expr_matrix.rds"))
network <- readRDS(file.path(inputs_dir, "input-dorothea_genesets.rds"))

run_viper(mat, network, tf, target, mor, likelihood, verbose = FALSE)
```


Index

* **convert_to_ variants**

convert_to_, [4](#)

* **decoupleR statistics**

decouple, [5](#)

run_gsva, [7](#)

run_mean, [8](#)

run_ora, [10](#)

run_pscira, [12](#)

run_scira, [13](#)

run_viper, [15](#)

base::rowMeans(), [14](#)

convert_f_defaults, [2](#)

convert_f_defaults(), [4](#)

convert_to_, [4](#)

convert_to_gsva (convert_to_), [4](#)

convert_to_mean (convert_to_), [4](#)

convert_to_ora (convert_to_), [4](#)

convert_to_pscira (convert_to_), [4](#)

convert_to_scira (convert_to_), [4](#)

convert_to_viper (convert_to_), [4](#)

decouple, [5](#), [8](#), [10](#), [11](#), [13](#), [14](#), [16](#)

dplyr::rename(), [2](#), [3](#)

dplyr::select(), [3](#)

filter_regulons, [7](#)

GSVA::gsva, [8](#)

GSVA::gsva(), [4](#), [7](#)

run_gsva, [6](#), [7](#), [10](#), [11](#), [13](#), [14](#), [16](#)

run_mean, [6](#), [8](#), [8](#), [11](#), [13](#), [14](#), [16](#)

run_ora, [6](#), [8](#), [10](#), [10](#), [13](#), [14](#), [16](#)

run_pscira, [6](#), [8](#), [10](#), [11](#), [12](#), [14](#), [16](#)

run_scira, [6](#), [8](#), [10](#), [11](#), [13](#), [13](#), [16](#)

run_viper, [6](#), [8](#), [10](#), [11](#), [13](#), [14](#), [15](#)

speedglm::speedlm.fit(), [14](#)

stats::fisher.test, [11](#)

stats::fisher.test(), [10](#)

stats::lm(), [14](#)

tibble::add_column(), [2](#)

viper::viper, [15](#)

viper::viper(), [4](#), [15](#)