# Package 'NetCoupler'

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**Title** Inference of Causal Links Between a Network and an External Variable **Version** 0.1.0

**Description** The 'NetCoupler' algorithm identifies potential direct effects of correlated, high-dimensional variables formed as a network with an external variable. The external variable may act as the dependent/response variable or as an independent/predictor variable to the network.

```
License MIT + file LICENSE
URL https://github.com/NetCoupler/NetCoupler
BugReports https://github.com/NetCoupler/NetCoupler/issues
Depends R (>= 3.5.0)
Imports checkmate,
     dplyr,
     ids,
     igraph,
     lifecycle,
     magrittr,
     pcalg,
     ppcor,
     purrr,
     rlang (>= 0.4.6),
     stats,
     tibble,
     tidyselect,
     utils,
     tidygraph
Suggests broom,
     furrr,
     knitr,
     rmarkdown,
     spelling,
     testthat (>= 2.1.0)
VignetteBuilder knitr
```

RdMacros lifecycle

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### biocViews

ByteCompile true

**Encoding** UTF-8

LazyData true

**Roxygen** list(markdown = TRUE)

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Language en-US

# R topics documented:

as_edge_tbl																	
classify_options															 		
nc_estimate_links .																	
nc_estimate_networ	k																
nc_standardize																	
simulated_data															 		

as\_edge\_tbl

Convert network graphs to edge tables as tibbles/data.frames.

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#### **Description**

#### [Experimental]

#### Usage

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```
as_edge_tbl(network_object)
```

#### **Arguments**

network\_object Network graph from nc\_estimate\_network().

#### Value

A tibble, with at least two columns:

- source\_node: The starting node (variable).
- target\_node: The ending node (variable) that links with the source node.
- adjacency\_weight: (Optional) The "weight" given to the edge, which represents the strength of the link between two nodes.

#### See Also

See nc\_estimate\_links for examples on using NetCoupler.

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classify\_options

Classification options for direct, ambigious, and no effect.

# Description

Classification options for direct, ambigious, and no effect.

#### Usage

```
classify_options(
  single_metabolite_threshold = 0.05,
  network_threshold = 0.1,
  direct_effect_adjustment = NA
)
```

# **Arguments**

single\_metabolite\_threshold, network\_threshold, direct\_effect\_adjustment

See the classify\_option\_list argument in nc\_estimate\_links for details.

#### Value

List with options for the classification.

nc\_estimate\_links

Compute model estimates between an external (exposure or outcome) variable and a network.

# Description

#### [Experimental]

This is the main function that identifies potential links between external factors and the network. There are two functions to estimate and classify links:

- nc\_estimate\_exposure\_links(): Computes the model estimates for the exposure side.
- nc\_estimate\_outcome\_links(): Computes the model estimates for the exposure side.

#### Usage

```
nc_estimate_exposure_links(
  data,
  edge_tbl,
  exposure,
  adjustment_vars = NA,
  model_function,
  model_arg_list = NULL,
```

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```
exponentiate = FALSE,
  classify_option_list = classify_options()
)
nc_estimate_outcome_links(
  data,
  edge_tbl,
  outcome,
  adjustment_vars = NA,
 model_function,
 model_arg_list = NULL,
  exponentiate = FALSE,
  classify_option_list = classify_options()
)
```

### **Arguments**

data

The data frame or tibble that contains the variables of interest, including the variables used to make the network.

edge\_tbl

Output graph object from nc\_estimate\_network(), converted to an edge table using as\_edge\_tbl().

exposure, outcome

Character. The exposure or outcome variable of interest.

adjustment\_vars

Optional. Variables to adjust for in the models.

model\_function A function for the model to use (e.g. stats::lm(), stats::glm(), survival::coxph()). Can be any model as long as the function has the arguments formula and data. Type in the model function as a bare object (without (), for instance as 1m).

model\_arg\_list Optional. A list containing the named arguments that will be passed to the model function. A simple example would be list(family = binomial(link = "logit")) to specify that the glm model is a logistic model and not a linear one. See the examples for more on the usage.

exponentiate

Logical. Whether to exponentiate the log estimates, as computed with e.g. logistic regression models.

classify\_option\_list

A list with classification options for direct, ambigious, or no effects. Used with the classify\_options() function with the arguments:

- single\_metabolite\_threshold: Default of 0.05. P-values from models with only the index metabolite (no neighbour adjustment) are classified as effects if below this threshold. For larger sample sizes and networks, we recommend lowering the threshold to reduce risk of false positives.
- network\_threshold: Default of 0.1. P-values from any models that have direct neighbour adjustments are classified as effects if below this threshold. This is assumed as a one-sided p-value threshold. Like the threshold above, a lower value should be used for larger sample sizes and networks.

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 direct\_effect\_adjustment: Default is NA. After running the algorithm once, sometimes it's useful to adjust for the direct effects identified to confirm whether other links exist.

#### Value

Outputs a tibble that contains the model estimates from either the exposure or outcome side of the network as well as the effect classification. Each row represents the "no neighbour node adjusted" model and has the results for the outcome/exposure to index node pathway. Columns for the outcome are:

- outcome or exposure: The name of the variable used as the external variable.
- index\_node: The name of the metabolite used as the index node from the network. In combination with the outcome/exposure variable, they represent the individual model used for the classification.
- estimate: The estimate from the outcome/exposure and index node model.
- std\_error: The standard error from the outcome/exposure and index node model.
- fdr\_p\_value: The False Discovery Rate-adjusted p-value from the outcome/exposure and index node model.
- effect: The NetCoupler classified effect between the index node and the outcome/exposure. Effects are classified as "direct" (there is a probable link based on the given thresholds), "ambigious" (there is a potential link but not all thresholds were passed), and "none" (no potential link seen).

The tibble output also has an attribute that contains all the models generated *before* classification. Access it with attr(output, "all\_models\_df").

#### See Also

vignette("examples") article has more details on how to use NetCoupler with different models.

#### **Examples**

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# Get results of all models used prior to classification

nc\_estimate\_network

Create an estimate of the metabolic network as an undirected graph.

#### **Description**

#### [Experimental]

The main NetCoupler network creator. Uses the input data to estimate the underlying undirected graph. The default uses the PC algorithm, implemented within NetCoupler with pc\_estimate\_undirected\_graph() Defaults to using the PC algorithm to calculate possible edges. Any missing values in the input data are removed by this function, since some computations can't handle missingness.

#### Usage

```
nc_estimate_network(data, cols = everything(), alpha = 0.01)
```

# **Arguments**

data	Data that would form the underlying network.

cols <tidy-select> Variables to include by using dplyr::select() style selec-

tion.

alpha The alpha level to use to test whether an edge exists or not. Default is 0.01.

#### Value

Outputs a tidygraph::tbl\_graph() with the start and end nodes, as well as the edge weights.

#### See Also

See nc\_estimate\_links for examples on using NetCoupler and pc\_estimate\_undirected\_graph for more details on the PC-algorithm network estimation method.

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#### **Description**

#### [Experimental]

Can standardize by either 1) log()-transforming and then applying scale() (mean-center and scaled by standard deviation), or 2) if regressed\_on variables are given, then log-transforming, running a linear regression to obtain the stats::residuals(), and finally scaled. Use regressed\_on to try to remove influence of potential confounding.

#### Usage

```
nc_standardize(data, cols = everything(), regressed_on = NULL)
```

#### **Arguments**

data Data frame.

cols Metabolic variables that will make up the network.

regressed\_on Optional. A character vector of variables to regress the metabolic variables

on. Use if you want to standardize the metabolic variables on variables that are known to influence them, e.g. sex or age. Calculates the residuals from a

linear regression model.

#### Value

Outputs a tibble object, with the original metabolic variables now standardized.

#### See Also

nc\_estimate\_links for more detailed examples or the vignette("NetCoupler").

# Examples

```
# Don't regress on any variable
simulated_data %>%
    nc_standardize(starts_with("metabolite_"))

# Extract residuals by regressing on a variable
simulated_data %>%
    nc_standardize(starts_with("metabolite_"), "age")

# Works with factors too
simulated_data %>%
    dplyr::mutate(Sex = as.factor(sample(rep(c("F", "M"), times = nrow(.) / 2)))) %>%
    nc_standardize(starts_with("metabolite_"), c("age", "Sex"))
```

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simulated_data	Simulated dataset with an underlying Directed Graph structure for the metabolites.

# Description

Simulated dataset with an underlying Directed Graph structure for the metabolites.

# Usage

simulated\_data

#### **Format**

The simulated dataset is a tibble with the following variables:

- Two outcome variables (outcome\_continuous and outcome\_binary) along with survival time (outcome\_event\_time) that is used for the outcome\_binary variable
- A generic exposure variable as continuous
- 12 metabolite\_\* variables
- An age variable used as a confounder

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