Package ‘NetworkInference’

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Description This is an R implementation of the netinf algorithm (Gomez Rodriguez, Leskovec, and Krause, 2010)<doi:10.1145/1835804.1835933>. Given a set of events that spread between a set of nodes the algorithm infers the most likely stable diffusion network that is underlying the diffusion process.
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as.data.frame.cascade

Convert a cascade object to a data frame

Description
Generates a data frame containing the cascade information in the cascade object.

Usage
```r
## S3 method for class 'cascade'
as.data.frame(x, row.names = NULL, optional = FALSE, ...)
```

Arguments
- `x` Cascade object to convert.
- `row.names` NULL or a character vector giving the row names for the data frame. Missing values are not allowed.
- `optional` logical. If TRUE, setting row names and converting column names (to syntactic names: see make.names) is optional. (Not supported)
- `...` Additional arguments passed to `data.frame`.
as.matrix.cascade

Value
A data frame with three columns. Containing 1) The names of the nodes ("node_name") that experience an event in each cascade, 2) the event time ("event_time") of the corresponding node, 3) the cascade identifier "cascade_id".

Examples

```r
data(cascades)
as.data.frame(cascades)
```

as.matrix.cascade  Convert a cascade object to a matrix

Description
Generates a matrix containing the cascade information in the cascade object in wide format. Missing values are used for nodes that do not experience an event in a cascade.

Usage

```r
## S3 method for class 'cascade'
as.matrix(x, ...)
```

Arguments

- `x` cascade object to convert.
- `...` additional arguments to be passed to or from methods. (Currently not supported.)

Value
A matrix containing all cascade information in wide format. That is, each row of the matrix corresponds to a node and each column to a cascade. Cell entries are event times. Censored nodes have NA for their entry.

Examples

```r
data(cascades)
as.matrix(cascades)
```
as_cascade_long  Transform long data to cascade

Description
Create a cascade object from data in long format.

Usage
as_cascade_long(data, cascade_node_name = "node_name",
    event_time = "event_time", cascade_id = "cascade_id",
    node_names = NULL)

Arguments
data  data.frame, containing the cascade data with column names corresponding to the
    arguments provided to cascade_node_names, event_time and cascade_id.
cascade_node_name  character, column name of data that specifies the node names in the cascade.
event_time  character, column name of data that specifies the event times for each node
    involved in a cascade.
cascade_id  character, column name of the cascade identifier.
node_names  character, factor or numeric vector containing the names for each node. Optional. If not provided, node names are inferred from the cascade data.

Details
Each row of the data describes one event in the cascade. The data must contain at least three
    columns:
    1. Cascade node name: The identifier of the node that experiences the event.
    2. Event time: The time when the node experiences the event. Note that if the time column is of
class date or any other special time class, it will be converted to an integer with 'as.numeric()'.
    3. Cascade id: The identifier of the cascade that the event pertains to.
The default names for these columns are node_name, event_time and cascade_id. If other names
are used in the data object the names have to be specified in the corresponding arguments (see
argument documentation)

Value
An object of class cascade. This is a list containing three (named) elements:
    1. "node_names" A character vector of node names.
    2. "cascade_nodes" A list with one character vector per cascade containing the node names in
order of the events.
    3. "cascade_times" A list with one element per cascade containing the event times for the
nodes in "cascade_names".
Examples

```r
df <- simulate_rnd_cascades(10, n_nodes = 20)
cascades <- as_cascade_long(df)
is.cascade(cascades)
```

---

as_cascade_wide

**Transform wide data to cascade**

Description

Create a cascade object from data in wide format.

Usage

```r
as_cascade_wide(data, node_names = NULL)
```

Arguments

data: data.frame or matrix, rows corresponding to nodes, columns to cascades. Matrix entries are the event times for each node, cascade pair. Missing values indicate censored observations, that is, nodes that did not have an event). Specify column and row names if cascade and node ids other than integer sequences are desired. Note that, if the time column is of class date or any other special time class, it will be converted to an integer with `as.numeric()`.

node_names: character, factor or numeric vector, containing names for each node. Optional. If not provided, node names are inferred from the provided data.

Details

If data is in wide format, each row corresponds to a node and each column to a cascade. Each cell indicates the event time for a node - cascade combination. If a node did not experience an event for a cascade (the node is censored) the cell entry must be `NA`.

Value

An object of class cascade. This is a list containing three (named) elements:

1. "node_names" A character vector of node names.
2. "cascade_nodes" A list with one character vector per cascade containing the node names in order of the events.
3. "cascade_times" A list with one element per cascade containing the event times for the nodes in "cascade_names".
Examples

```r
data("policies")
cascades <- as_cascade_long(policies, cascade_node_name = 'statenam',
                           event_time = 'adopt_year', cascade_id = 'policy')
wide_policies = as.matrix(cascades)
cascades <- as_cascade_wide(wide_policies)
is.cascade(cascades)
```

---

Example cascades

---

Description

An example dataset of 31 nodes and 54 cascades. From the original netinf implementation in SNAP.

Usage

```r
data(cascades)
```

Format

An object of class cascade containing 4 objects

- **node_names**  Character node names
- **cascade_nodes**  A list of integer vectors. Each containing the names of the nodes infected in this cascades in the order of infection
- **cascade_times**  A list of numeric vectors. Each containing the infection times for the corresponding nodes in cascade_nodes

Source

### count_possible_edges

**Count the number of possible edges in the dataset**

**Description**
Across all cascades, count the edges that are possible. An edge from node $u$ to node $v$ is only possible if in at least one cascade $u$ experienced an event before $v$.

**Usage**

```r
count_possible_edges(cascades)
```

**Arguments**

- `cascades`: Object of class cascade containing the data.

**Value**
An integer count.

**Examples**

```r
data(cascades)
count_possible_edges(cascades)
```

---

### drop_nodes

**Drop nodes from a cascade object**

**Description**
Drop nodes from a cascade object

**Usage**

```r
drop_nodes(cascades, nodes, drop = TRUE)
```

**Arguments**

- `cascades`: cascade, object to drop nodes from.
- `nodes`: character or integer, vector of node_ids to drop.
- `drop`: logical, Should empty cascades be dropped.

**Value**
An object of class cascade containing the cascades without the dropped nodes.
Examples

data(policies)
cascades <- as_cascade_long(policies, cascade_node_name = 'statenam',
    event_time = 'adopt_year', cascade_id = 'policy')
new_cascades <- drop_nodes(cascades, c("California", "New York"))

is.cascade

Description

Is the object of class cascade?

Usage

is.cascade(object)

Arguments

object the object to be tested.

Value

TRUE if object is a cascade, FALSE otherwise.

Examples

data(cascades)
is.cascade(cascades)
# > TRUE
is.cascade(1)
# > FALSE

is.diffnet

Description

Tests if an object is of class diffnet. The class diffnet is appended to the object returned by netinf
for dispatch of appropriate plotting methods.

Usage

is.diffnet(object)
Arguments

object  the object to be tested.

Value

TRUE if object is a diffnet, FALSE otherwise.

Examples

data(cascades)
result <- netinf(cascades, n_edges = 6, params = 1)
is.diffnet(result)

Description

Infer a network of diffusion ties from a set of cascades. Each cascade is defined by pairs of node ids and infection times.

Usage

netinf(cascades, trans_mod = "exponential", n_edges = NULL, p_value_cutoff = NULL, params = NULL, quiet = FALSE, trees = FALSE)

Arguments

cascades  an object of class cascade containing node and cascade information. See as_cascade_long and as_cascade_wide for details.

trans_mod  character, indicating the choice of model: "exponential", "rayleigh" or "log-normal".

n_edges  integer, number of edges to infer. Leave unspecified if using p_value_cutoff.

p_value_cutoff  numeric, in the interval (0, 1). If specified, edges are inferred in each iteration until the Vuong test for edge addition reaches the p-value cutoff or when the maximum possible number of edges is reached. Leave unspecified if using n_edges to explicitly specify number of edges to infer.

params  numeric, Parameters for diffusion model. If left unspecified reasonable parameters are inferred from the data. See details for how to specify parameters for the different distributions.

quiet  logical, Should output on progress by suppressed.

trees  logical, Should the inferred cascade trees be returned. Note, that this will lead to a different the structure of the function output. See section Value for details.
Details

The algorithm is described in detail in Gomez-Rodriguez et al. (2010). Additional information can be found on the netinf website (http://snap.stanford.edu/netinf/).

- Exponential distribution: trans_mod = "exponential", params = c(lambda). Parametrization: $\lambda e^{-\lambda x}$.
- Rayleigh distribution: trans_mod = "rayleigh", params = c(alpha). Parametrization: $\frac{x}{\alpha^2} e^{-\frac{x^2}{2\alpha^2}}$.
- Log-normal distribution: trans_mod = "log-normal", params = c(mu, sigma). Parametrization: $\frac{1}{\sqrt{2\pi \sigma^2}} e^{-\frac{(lnx-\mu)^2}{2\sigma^2}}$.

If higher performance is required and for very large data sets, a faster pure C++ implementation is available in the Stanford Network Analysis Project (SNAP). The software can be downloaded at http://snap.stanford.edu/netinf/.

Value

Returns the inferred diffusion network as an edgelist in an object of class `diffnet` and `data.frame`. The first column contains the sender, the second column the receiver node. The third column contains the improvement in fit from adding the edge that is represented by the row. The output additionally has the following attributes:

- "diffusion_model": The diffusion model used to infer the diffusion network.
- "diffusion_model_parameters": The parameters for the model that have been inferred by the approximate profile MLE procedure.

If the argument trees is set to TRUE, the output is a list with the first element being the data.frame described above, and the second element being the trees in edge-list form in a single data.frame.

References


Examples

# Data already in cascades format:
data(cascades)
out <- netinf(cascades, trans_mod = "exponential", n_edges = 5, params = 1)

# Starting with a dataframe
df <- simulate_rnd_cascades(10, n_nodes = 20)
cascades2 <- as_cascade_long(df, node_names = unique(df$node_name))
out <- netinf(cascades2, trans_mod = "exponential", n_edges = 5, params = 1)
NetworkInference: Inferring latent diffusion networks

Description
This package provides an R implementation of the netinf algorithm created by Gomez Rodriguez, Leskovec, and Krause (2010). Given a set of events that spread between a set of nodes the algorithm infers the most likely stable diffusion network that is underlying the diffusion process.

Details
The package provides three groups of functions: 1) data preparation 2) estimation and 3) interpretation.

Data preparation
The core estimation function netinf requires an object of class cascade (see as_cascade_long and as_cascade_wide). Cascade data contains information on the potential nodes in the network as well as on event times for each node in each cascade.

Estimation
Diffusion networks are estimated using the netinf function. It produces a diffusion network in form of an edgelist (of class data.frame).

Interpretation and Visualization
Cascade data can be visualized with the plot method of the cascade class (diffnet, plot.cascade). Results of the estimation process can be visualized using the plotting method of the diffnet class.

Performance
If higher performance is required and for very large data sets, a faster pure C++ implementation is available in the Stanford Network Analysis Project (SNAP). The software can be downloaded at http://snap.stanford.edu/netinf/.

plot.cascade
Plot a cascade object

Description
Allows plotting of one or multiple, labeled or unlabeled cascades.

Usage
## S3 method for class 'cascade'
plot(x, label_nodes = TRUE, selection = NULL, ...)

### References
plot.diffnet

Visualize netinf output

Description

Visualize the inferred diffusion network or the marginal gain in fit obtained by addition of each edge.

Usage

## S3 method for class 'diffnet'
plot(x, type = "network", ...)
Arguments

- **x**: object of class diffnet to be plotted.
- **type**: character, one of c("network", "improvement", "p-value") indicating if the inferred diffusion network, the improvement for each edge or the p-value from the vuong test for each edge should be visualized.
- ... additional arguments.

Details

If 'type = improvement' a ggplot object is returned. It can be modified like any other ggplot. See the ggplot documentation and the examples in plot.cascade.

Value

A ggplot plot object if type = "improvement" otherwise an igraph plot.

Examples

```r
## Not run:
data(cascades)
res <- netinf(cascades, quiet = TRUE)
plot(res, type = "network")
plot(res, type = "improvement")
plot(res, type = "p-value")
## End(Not run)
```

---

**policies**

**US State Policy Adoption (SPID)**

Description

The SPID data includes information on the year of adoption for over 700 policies in the American states.

Usage

data(policies)

Format

The data comes in two objects of class data.frame. The first object, named policies contains the adoption events. Each row corresponds to an adoption event. Each adoption event is described by the three columns:

- **statenam**: Name of the adopting state.
• **policy**: Name of the policy.
• **adopt_year**: Year when the state adopted the policy.

The second object (policies_metadata) contains more details on each of the policies. It contains these columns:

• **policy**: Name of the policy.
• **source**: Original source of the data.
• **first_year**: First year any state adopted this policy.
• **last_year**: Last year any state adopted this policy.
• **adopt_count**: Number of states that adopted this policy.
• **description**: Description of the policy.
• **majortopic**: Topic group the policy belongs to.

Both data.frame objects can be joined (merged) on the common column policy (see example code).

**Details**

This version 1.0 of the database. For each policy we document the year of first adoption for each state. Adoption dates range from 1691 to 2017 and includes all fifty states. Policies are adopted by anywhere from 1 to 50 states, with an average of 24 adoptions. The data were assembled from a variety of sources, including academic publications and policy advocacy/information groups. Policies were coded according to the Policy Agendas Project major topic code. Additional information on policies is available at the source repository.

**Source**

https://doi.org/10.7910/DVN/CVYSR7

**References**

Boehmke, Frederick J.; Mark Brockway; Bruce A. Desmarais; Jeffrey J. Harden; Scott LaCombe; Fridolin Linder; and Hanna Wallach. 2018. "A New Database for Inferring Public Policy Innovativeness and Diffusion Networks." Working paper.

**Examples**

data('policies')

# Join the adoption events with the metadata
merged_policies <- merge(policies, policies_metadata, by = 'policy')
simulate_cascades  

Simulate cascades from a diffusion network

**Description**

Simulate diffusion cascades based on the generative model underlying netinf and a diffusion network.

**Usage**

```r
simulate_cascades(diffnet, nsim = 1, max_time = Inf, 
                   start_probabilities = NULL, partial_cascade = NULL, 
                   params = NULL, 
                   model = NULL, nodes = NULL)
```

**Arguments**

- **diffnet** object of class `diffnet`.
- **nsim** integer, number of cascades to simulate.
- **max_time** numeric, the maximum time after which observations are censored
- **start_probabilities** a vector of probabilities for each node in `diffnet`, to be the node with the first event. If NULL a node is drawn from a uniform distribution over all nodes.
- **partial_cascade** object of type `cascade`, containing one partial cascades for which further development should be simulated.
- **params** numeric, (optional) parameters for diffusion time distribution. See the details section of `netinf` for specification details. Only use this argument if parameters different from those contained in the `diffnet` object should be used or the network is not an object of class `diffnet`.
- **model** character, diffusion model to use. One of c("exponential","rayleigh", "log-normal"). Only use this argument if parameters different from those contained in the `diffnet` object should be used or the network is not an object of class `diffnet`.
- **nodes** vector of node ids if different from nodes included in `diffnet`.

**Value**

A data frame with three columns. Containing 1) The names of the nodes ("node_name") that experience an event in each cascade, 2) the event time ("event_time") of the corresponding node, 3) the cascade identifier "cascade_id".
Examples

```r
data(cascades)
out <- netinf(cascades, trans_mod = "exponential", n_edges = 5, params = 1)
simulated_cascades <- simulate_cascades(out, nsim = 10)
```

# Simulation from partial cascade

---

**simulate_rnd_cascades**  
*Simulate a set of random cascades*

### Description

Simulate random cascades, for testing and demonstration purposes. No actual diffusion model is underlying these cascades.

### Usage

```r
simulate_rnd_cascades(n_cascades, n_nodes)
```

### Arguments

- `n_cascades`  
  Number of cascades to generate.

- `n_nodes`  
  Number of nodes in the system.

### Value

A data frame containing (in order of columns) node ids, event time and cascade identifier.

### Examples

```r
df <- simulate_rnd_cascades(10, n_nodes = 20)
head(df)
```
**sim_validation**

Larger simulated validation network.

**Description**
A network from simulated data. For testing purposes.

**Usage**

data(sim_validation)

**Format**
An object of class `data.frame` with 4 columns, containing:

- **origin_node**: Origin of diffusion edge.
- **destination_node**: Destination node of diffusion edge.
- **improvement**: Improvement in score for the edge
- **p-value**: p-value for vuong test

**Source**
See code below.

---

**subset_cascade**

Select a subset of cascades from cascade object

**Description**
Select a subset of cascades from cascade object

**Usage**

subset_cascade(cascade, selection)

**Arguments**

- **cascade**: cascade, object to select from
- **selection**: character or integer, vector of cascade_ids to select

**Value**
An object of class cascade containing just the selected cascades
Examples

data(policies)
cascades <- as_cascade_long(policies, cascade_node_name = 'statenam',
                         event_time = 'adopt_year', cascade_id = 'policy')
cascade_names <- names(cascades$cascade_times)
subset_cascade(cascades, selection = cascade_names[1:10])

subset_cascade_time  Subset a cascade object in time

Description
Remove each all events occurring outside the desired subset for each cascade in a cascade object.

Usage
subset_cascade_time(cascade, start_time, end_time, drop = TRUE)

Arguments
- cascade: cascade, object to subset.
- start_time: numeric, start time of the subset.
- end_time: numeric, end time of the subset.
- drop: logical, should empty sub-cascades be dropped?

Value
An object of class cascade, where only events are included that have times start_time <= t < end_time.

Examples

data(cascades)
sub_cascades <- subset_cascade_time(cascades, 10, 20, drop=TRUE)
summary.cascade

**Summary**

Summarize a cascade object

**Description**

Generates summary statistics for single cascades and across cascades in a collection, contained in a cascades object.

**Usage**

```r
## S3 method for class 'cascade'
summary(object, quiet = FALSE, ...)
```

**Arguments**

- `object` object of class cascade to be summarized.
- `quiet` logical, if FALSE summary stats are printed to std out.
- `...` Additional arguments passed to summary.

**Value**

Prints cascade summary information to the screen (if quiet = FALSE). '# cascades' is the number of cascades in the object, '# nodes' is the number of nodes in the system (nodes that can theoretically experience an event), '# nodes in cascades' is the number of unique nodes of the system that experienced an event and '# possible edges' is the number of edges that are possible given the cascade data (see `count_possible_edges` for details.).

Additional summaries for each cascade are returned invisibly. cascade, length (length of the cascade as an integer of how many nodes experienced and event) and n_ties (number of tied event times per cascade).

**Examples**

```r
data(cascades)
summary(cascades)
```

**validation**

Validation output from netinf source.

**Description**

Contains output from original netinf C++ implementation, executed on cascades. For testing purposes.
Usage

data(validation)

Format

An object of class `data.frame` with 6 columns, containing:

- `origin_node`  Origin of diffusion edge.
- `destination_node`  Destination node of diffusion edge.
- `volume`  ??
- `marginal_gain`  Marginal gain from edge.
- `median_time_difference`  Median time between events in origin and destination
- `mean_time_difference`  Mean time between events in origin and destination

Source

Output from netinf example program ([https://github.com/snap-stanford/snap/tree/master/examples/netinf](https://github.com/snap-stanford/snap/tree/master/examples/netinf)).
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