

# Package ‘epicontacts’

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**Type** Package

**Title** Handling, Visualisation and Analysis of Epidemiological Contacts

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**Description** A collection of tools for representing epidemiological contact data, composed of case line lists and contacts between cases. Also contains procedures for data handling, interactive graphics, and statistics.

**License** GPL (>= 2)

**RoxygenNote** 7.1.1

**Imports** grDevices, dplyr, igraph, visNetwork, threejs, colorspace, methods

**Suggests** outbreaks, testthat, covr, shiny, readr, knitr, rmarkdown

**VignetteBuilder** knitr

**URL** <https://www.repidemicsconsortium.org/epicontacts/>

**BugReports** <https://github.com/reconhub/epicontacts/issues>

**NeedsCompilation** no

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---

as.igraph.epicontacts *Create igraph object from contact data*

---

### Description

This function creates an igraph object from a given `epicontacts` object containing a 'contacts' dataframe.

### Usage

```
## S3 method for class 'epicontacts'
as.igraph(x)
```

### Arguments

x An `epicontacts` object.

### Value

An igraph object (from the `igraph` package). Note: any column called "name" in the original linelist will be stored as a new vertex attribute in the igraph object named 'epicontacts\_name'. This is due to the inherent behaviour of igraph creating its own 'name' vertex attribute.

**Author(s)**

Nistara Randhawa (<nrandhawa@ucdavis.edu>)

**Examples**

```
if (require(outbreaks) && require(igraph)) {
  ## build data

  x <- make_epicontacts(ebola_sim$linelist, ebola_sim$contacts,
                       id = "case_id", to = "case_id", from = "infector",
                       directed = TRUE)

  ## subset data - keep 50 cases from linelist with contacts

  ids <- get_id(x, "common")[1:50]
  ids
  x <- x[ids, ids]

  ## make igraph object with associated attributes from epicontacts object

  net <- as.igraph(x)
  net
  plot(net, vertex.label = "", vertex.size = 10,
       vertex.color = cases_pal(50))
}
```

---

codeawesome

*Reference codes for fontawesome*

---

**Description**

The object codeawesome is a character vector of fontawesome codes, named after their aliases.

**Usage**

```
codeawesome
```

**Format**

An object of class character of length 519.

**Author(s)**

Thibaut Jombart

---

get\_clusters                      *Assign cluster IDs to epicontacts data*

---

### Description

This function identifies transitive clusters (i.e. connected components) as well as the number of members in each cluster, and adds this information to the linelist data.

### Usage

```
get_clusters(  
  x,  
  output = c("epicontacts", "data.frame"),  
  member_col = "cluster_member",  
  size_col = "cluster_size",  
  override = FALSE  
)
```

### Arguments

x	An <a href="#">epicontacts</a> object.
output	A character string indicating the type of output: either an <a href="#">epicontacts</a> object (default) or a <a href="#">data.frame</a> containing cluster memberships to which members of <a href="#">epicontacts</a> linelist belong to.
member_col	Name of column to which cluster membership is assigned to in the linelist. Default name is 'cluster_member'.
size_col	Name of column to which cluster sizes are assigned to in the linelist. Default name is 'cluster_size'.
override	Logical value indicating whether cluster member and size columns should be overwritten if they already exist in the linelist. Default is 'FALSE'.

### Value

An [epicontacts](#) object whose 'linelist' dataframe contains new columns corresponding to cluster membership and size, or a [data.frame](#) containing member ids, cluster memberships as factors, and associated cluster sizes. All ids that were originally in the 'contacts' dataframe but not in the linelist will also be added to the linelist.

### Author(s)

Nistara Randhawa (<nrandhawa@ucdavis.edu>)

## Examples

```
if (require(outbreaks)) {
  ## build data
  x <- make_epicontacts(ebola_sim$linelist, ebola_sim$contacts,
                       id = "case_id",
                       to = "case_id",
                       from = "infectior",
                       directed = TRUE)

  ## add cluster membership and sizes to epicontacts 'linelist'
  y <- get_clusters(x, output = "epicontacts")
  y

  ## return a data.frame with linelist member ids and cluster memberships as
  ## factors
  z <- get_clusters(x, output = "data.frame")
  head(z)
}
```

---

get\_degree

*Find node degree for epicontacts objects*

---

## Description

This function computes the number of contacts per cases in a [epicontacts](#) dataset. Whenever contacts are directed, the argument 'type' can be used to specify which kind of contact should be considered: 'in' (towards the case), 'out' (from the case), or 'both'.

## Usage

```
get_degree(x, type = c("in", "out", "both"), only_linelist = FALSE)
```

## Arguments

x	an <a href="#">epicontacts</a> object
type	the type of degree to be computed (see description); if contacts are not directed, this will be forced to 'both'
only_linelist	a logical indicating if cases whose degree is computed should be from the linelist exclusively

## Author(s)

Thibaut Jombart (<[thibautjombart@gmail.com](mailto:thibautjombart@gmail.com)>)

## Examples

```
## make epicontacts object
if (require(outbreaks)) {
  x <- make_epicontacts(ebola_sim$linelist, ebola_sim$contacts,
                        id="case_id", to="case_id", from="infector",
                        directed=TRUE)

  x

  ## compute in-degree
  deg_in <- get_degree(x)
  table(deg_in)

  ## compute out-degree
  deg_out <- get_degree(x, "out")
  barplot(table(deg_out), main = "Reproduction number distribution")
  mtext(side = 3, "(based on case out-degree)")

}
```

---

get\_id

*Access unique identifiers in epicontacts objects*

---

## Description

This accessor is used to extract unique identifiers from `epicontacts` objects. The argument `'which'` can be used to specify if IDs should include: `linelist` only (`'linelist'`), `contacts` only (`'contacts'`), the union of both (`'all'`), or the intersection of both (`'common'`); two additional options are `'from'` (ID `'giving'` contacts) and `'to'` (ID `'receiving'` contacts).

## Usage

```
get_id(
  x,
  which = c("linelist", "contacts", "all", "common", "from", "to"),
  na.rm = TRUE
)
```

## Arguments

<code>x</code>	an <code>epicontacts</code> object
<code>which</code>	the type of ID to return (see description); value can be <code>'linelist'</code> , <code>'contacts'</code> , <code>'all'</code> , <code>'common'</code> , <code>'from'</code> or <code>'to'</code> .
<code>na.rm</code>	a <code>'logical'</code> indicating if <code>'NA'</code> should be removed from the output ( <code>'TRUE'</code> , default) or not.

## Value

`x` a character vector of unique identifiers

**Author(s)**

Thibaut Jombart (<thibautjombart@gmail.com>)

**Examples**

```
if (require(outbreaks)) {
  ## build data
  x <- make_epicontacts(ebola_sim$linelist, ebola_sim$contacts,
                       id="case_id", to="case_id", from="infectior",
                       directed=TRUE)

  ## get identifiers
  id1 <- get_id(x, "linelist")
  id2 <- get_id(x, "contacts")
  id3 <- get_id(x, "all")
  id4 <- get_id(x, "common")

  ## check intersections and unions
  all.equal(union(id1, id2), id3)
  all.equal(intersect(id1, id2), id4)

}
```

---

get\_pairwise

*Characterise contacts by comparing case attributes*

---

**Description**

This function extract attributes of cases involved in contacts using case information provided in the linelist of an [epicontacts](#) dataset. If not provided, the function used to process attributes will adjust to the type of attribute selected (see details).

**Usage**

```
get_pairwise(x, attribute, f = NULL, hard_NA = TRUE)
```

**Arguments**

x	an <a href="#">epicontacts</a> object
attribute	the attribute to be examined between contact pairs
f	a function processing the attributes of 'from' and 'to'
hard_NA	a logical indicating if the output should be NA whenever one of the paired values is NA (TRUE, default); otherwise, 'NA' may be treated as another character (e.g. when pasting paired values)

**Author(s)**

Thibaut Jombart (<thibautjombart@gmail.com>) Tom Crellen (<tomcrellen@gmail.com>)

**Examples**

```

if (require(outbreaks)) {
## example using MERS outbreak in Korea, 2014
head(mers_korea_2015[[1]])
head(mers_korea_2015[[2]])

x <- make_epicontacts(linelist=mers_korea_2015[[1]],
contacts=mers_korea_2015[[2]], directed=TRUE)

## estimate serial interval (onset->onset)
SI <- get_pairwise(x, "dt_onset")
SI
summary(SI)
hist(SI, col="grey", border="white", xlab="Days after symptoms",
main="MERS Korea 2014 - Serial Interval")

## check gender mixing:
get_pairwise(x, "sex") # not good, we want 2-way table

get_pairwise(x, "sex", f=table) # use custom function
fisher.test(get_pairwise(x, "sex", f=table)) # test association
}

```

---

graph3D

---

*Interactive 3D Force-directed graph from epicontacts object*


---

**Description**

This function creates a 3D graph from an epicontacts object

**Usage**

```

graph3D(
  x,
  node_color = "id",
  annot = TRUE,
  col_pal = cases_pal,
  NA_col = "lightgrey",
  g_title = "",
  bg_col = "white",
  label_col = "darkgrey",
  node_size = 1,
  edge_size = 0.5
)

```



**Arguments**

x	An <code>epicontacts</code> object
node_color	An index or character string indicating which field of the linelist should be used to color the nodes. Default is <code>id</code>
annot	An index, logical, or character string indicating which fields of the linelist should be used for annotating the nodes upon mouseover. The default <code>TRUE</code> shows the <code>'id'</code> and <code>'node_color'</code> (if the grouping column is different from <code>'id'</code> ).
col_pal	A color palette for the <code>node_colors</code> .
NA_col	The color used for unknown <code>node_color</code> .
g_title	The title of the graph.
bg_col	The background color of graph.
label_col	The color of the graph title and labels of groups.
node_size	The sizes of graph nodes.
edge_size	The width of graph edges.

**Value**

An `htmlwidget` object that is displayed using the object's `show` or `print` method. (If you don't see your widget plot, try printing it with the `print` function.)

**Note**

All colors must be specified as color names like `"red"`, `"blue"`, etc. or as hexadecimal color values without opacity channel, for example `"#FF0000"`, `"#0a3e55"` (upper or lower case hex digits are allowed).

Double-click or tap on the plot to reset the view.

**Author(s)**

Nistara Randhawa (<nrandhawa@ucdavis.edu>) Thibaut Jombart (<thibautjombart@gmail.com>)  
VP Nagraj (<vpnagraj@virginia.edu>)

**References**

Original `rthreejs` code by B. W. Lewis: <https://github.com/bwlewis/rthreejs>.

**Examples**

```
if (require(outbreaks)) {

  ## example using MERS outbreak in Korea, 2014
  head(mers_korea_2015[[1]])
  head(mers_korea_2015[[2]])

  x <- make_epicontacts(linelist = mers_korea_2015$linelist,
                       contacts = mers_korea_2015$contacts,
                       directed = FALSE)
```

```
## Not run:
graph3D(x)
graph3D(x, annot = FALSE)
graph3D(x, node_color = "sex", g_title = "MERS Korea 2014")
graph3D(x, node_color = "sex", annot = c("sex", "age"),
        g_title = "MERS Korea 2014")

## End(Not run)
}
```

---

make\_epicontacts      *Read linelist and contact data*

---

## Description

This function reads data stored as `data.frame` containing `linelist` (case information, where each row corresponds to a unique patient), and `contacts` between patients. Common identifiers should be used in the two data sources for matching to be achieved.

## Usage

```
make_epicontacts(
  linelist,
  contacts,
  id = 1L,
  from = 1L,
  to = 2L,
  directed = FALSE
)
```

## Arguments

<code>linelist</code>	a <code>data.frame</code> with at least one column providing unique patient identifiers
<code>contacts</code>	a <code>data.frame</code> that needs at least two columns indicating patients between which cases take place; these need not be referenced in the <code>linelist</code>
<code>id</code>	an index or name indicating which column in <code>linelist</code> contains unique identifiers; default is first column in <code>linelist</code> data frame
<code>from</code>	an index or name indicating which column in <code>contacts</code> contains the first case of a contact
<code>to</code>	an index or name indicating which column in <code>contacts</code> contains the second case of a contact
<code>directed</code>	a logical indicating if contacts are directed or not; default is <code>FALSE</code> but note that contacts will be indicated as 'from' and 'to' even in non-directed contacts

## Details

An epicontacts object can be created from two components:

- a linelist provided as a `data.frame` where columns are different variables describing cases, and where each row is a different case. and a contact list.
- a contact list provided as a `data.frame` where each row contains unique pairs of contacts with unique features of contact in columns. The line list and contact list should share an identification scheme for individuals.

## Value

An epicontacts object in list format with three elements:

- `linelist`: `data.frame` of cases with first column 'id' containing character vector of unique identifiers
- `contacts`: `data.frame` of contacts with first two columns named 'from' and 'to' indicating unique pairs of contact between individuals
- `directed`: indicator as to whether or not the contacts are to be considered directed or not

## Author(s)

Thibaut Jombart (<thibautjombart@gmail.com>)

## References

<https://foodborne.unl.edu/public/role/epidemiologist/lineLists.html>

## Examples

```
if (require(outbreaks)) {
  ## make epicontacts object from simulated Ebola data
  x <- make_epicontacts(ebola_sim$linelist, ebola_sim$contacts)

  ## test reordering of columns
  linelist <- ebola_sim$linelist[,rev(seq_len(ncol(ebola_sim$linelist)))]
  contacts <- ebola_sim$contacts[,rev(seq_len(ncol(ebola_sim$contacts)))]
  head(linelist)
  head(contacts)

  ## make object
  x <- make_epicontacts(linelist, contacts, id = "case_id",
                       to = "case_id", from = "infectors")
  head(x$linelist)
  head(x$contacts)
}
```

---

plot.epicontacts      *Plot epicontacts objects*

---

### Description

This function plots [epicontacts](#) objects using various approaches. The default method uses [vis\\_epicontacts](#).

### Usage

```
## S3 method for class 'epicontacts'  
plot(  
  x,  
  node_color = "id",  
  method = c("visNetwork", "graph3D"),  
  thin = TRUE,  
  ...  
)
```

### Arguments

x	An <a href="#">epicontacts</a> object
node_color	An integer or a character string indicating which attribute column in the linelist should be used to color the nodes.
method	A character string indicating the plotting method to be used; available values are "visNetwork" and "graph3D"; see details.
thin	A logical indicating if the data should be thinned with <a href="#">thin</a> so that only cases with contacts should be plotted.
...	Further arguments passed to the plotting methods.

### Details

This function is merely a wrapper for other plotting functions in the package, depending on the value of method:

- visNetwork: calls the function [vis\\_epicontacts](#)
- graph3D: calls the function [graph3D](#)

### Author(s)

Thibaut Jombart (<[thibautjombart@gmail.com](mailto:thibautjombart@gmail.com)>)

### See Also

[vis\\_epicontacts](#), which uses the package visNetwork, and [codeawesome](#) for icon codes.

## Examples

```
if (require(outbreaks)) {
  ## example using MERS outbreak in Korea, 2014
  head(mers_korea_2015[[1]])
  head(mers_korea_2015[[2]])

  x <- make_epicontacts(linelist = mers_korea_2015[[1]],
                       contacts = mers_korea_2015[[2]], directed=TRUE)

  ## Not run:
  plot(x)
  plot(x, "place_infect")
  plot(x, "loc_hosp", legend_max = 20, annot = TRUE)
  plot(x, "place_infect", node_shape = "sex",
       shapes = c(M = "male", F = "female"))
  plot(x, 4)
  plot(x, 4, method = "graph3D")

  ## End(Not run)
}
```

---

print.epicontacts      *Print method for epicontacts objects*

---

## Description

This method prints the content of epicontacts objects, giving a brief summary of the reported cases and contacts.

## Usage

```
## S3 method for class 'epicontacts'
print(x, ...)
```

## Arguments

x                    an [epicontacts](#) object  
...                    further parameters to be passed to other methods (currently not used)

## Author(s)

Thibaut Jombart (<[thibautjombart@gmail.com](mailto:thibautjombart@gmail.com)>)

---

```
print.summary_epicontacts
```

*Print method for summary\_epicontacts objects*

---

### Description

This method outputs a printed summary of the content of `summary_epicontacts` objects.

### Usage

```
## S3 method for class 'summary_epicontacts'  
print(x, ...)
```

### Arguments

`x` a `summary_epicontacts` object  
`...` further parameters to be passed to other methods (currently not used)

### Author(s)

VP Nagraj (<vpnagraj@virginia.edu>)

---

```
subset.epicontacts
```

*Subset an epicontact object by factors*

---

### Description

This function subsets an `epicontacts` object based on node, edge and/or cluster attributes. Specifying node attributes will return an `epicontacts` object containing only individuals with these given attributes in the `linelist`. Specifying edge attributes will return contacts with the attributes provided. Specifying cluster attributes will return clusters of connected cases, and can be defined by `ids` (returning clusters of cases connected to specified cases) or cluster sizes (returning cluster of a specific, minimum or maximum size).

### Usage

```
## S3 method for class 'epicontacts'  
subset(  
  x,  
  node_attribute = NULL,  
  edge_attribute = NULL,  
  cluster_id = NULL,  
  cs = NULL,  
  cs_min = NULL,  
  cs_max = NULL,  
  ...  
)
```

**Arguments**

<code>x</code>	an epi_contact object to be subsetted
<code>node_attribute</code>	a named list defining the node attribute name and node attribute value (as a single value or vector of values). Dates must be provided as a vector of date objects, defining the range of dates included in the subset. If only one date is provided, only node attributes with that date will be returned.
<code>edge_attribute</code>	a named list defining the edge attribute name and edge attribute value (as a single value or vector of values). Dates must be provided as a vector of date objects, defining the range of dates included in the subset. If only one date is provided, only edge attributes with that date will be returned.
<code>cluster_id</code>	a character vector of case identifiers; the connected components attached to these cases will be retained in the output object.
<code>cs</code>	cluster size to be used for subsetting
<code>cs_min</code>	minimum cluster size for subsetting
<code>cs_max</code>	maximum cluster size for subsetting
<code>...</code>	further arguments passed on to other methods

**Author(s)**

Finlay Campbell (<f.campbell15@imperial.ac.uk>), Nistara Randhawa (<nrandhawa@ucdavis.edu>)

**Examples**

```

if (require(outbreaks)) {
## build data

x <- make_epicontacts(ebola_sim$linelist, ebola_sim$contacts,
                     id = "case_id", to = "case_id",
                     from = "infectior", directed = FALSE)

## subset based on node and edge attributes
x_subset <- subset(x, node_attribute = list("gender" = "f"),
                  edge_attribute = list("source" = "funeral"))

## subset a cluster connected to a given id
## (can be a vector of ids as well)
## here picking node with highest out-degree

id <- names(which.max(get_degree(x, "out")))
x_subset <- thin(subset(x, cluster_id = id), 2)
x_subset
plot(x_subset)

## subset based on cluster size range

x_subset <- subset(x, cs_min = 12, cs_max = 15)

```

```
## subset based on single cluster size
x_subset <- subset(x, cs = 12)

## subset based on minimum cluster size
x_subset <- subset(x, cs_min = 10)

## subset based on maximum cluster size
x_subset <- subset(x, cs_max = 9)

}
```

---

subset\_clusters\_by\_id *Subset epicontacts by case-specified clusters*

---

### Description

This function subsets an [epicontacts](#) object by identifying clusters of cases connected to specified cases.

### Usage

```
subset_clusters_by_id(x, id)
```

### Arguments

x	an <a href="#">epicontacts</a> object
id	a character vector of case identifiers; the connected components attached to these cases will be retained in the output object.

### Value

An [epicontacts](#) object whose contact dataframe corresponds to all clusters containing specified case id.

### Author(s)

Nistara Randhawa (<nrandhawa@ucdavis.edu>)



### Examples

```
if (require(outbreaks)) {  
  ## build data  
  x <- make_epicontacts(ebola_sim$linelist, ebola_sim$contacts,  
                        id="case_id", to="case_id", from="infector",  
                        directed=TRUE)  
  
  id <- "cac51e" ## it can be a vector of id as well  
  
  ## subset based on cluster to which "cac51e" belongs  
  x_subset <- subset_clusters_by_id(x, id)  
  
}
```

---

subset\_clusters\_by\_size

*Subset clusters from epicontacts object by cluster size*

---

### Description

This function subsets an `epicontacts` object based on defined size(s) of clusters (clusters being groups of connected individuals/nodes). Subsetting may be done by specifying a particular cluster size of interest, minimum cluster size, maximum cluster size, or a range (minimum and maximum) of cluster sizes.

### Usage

```
subset_clusters_by_size(x, cs = NULL, cs_min = NULL, cs_max = NULL)
```

### Arguments

x	an <code>epicontacts</code> object
cs	cluster size to be used for subsetting
cs_min	minimum cluster size for subsetting
cs_max	maximum cluster size for subsetting

### Value

An `epicontacts` object whose contact dataframe corresponds to all clusters of specified cluster sizes.

### Author(s)

Nistara Randhawa (<nrandhawa@ucdavis.edu>)

## Examples

```
if (require(outbreaks)) {  
  ## build data  
  x <- make_epicontacts(ebola_sim$linelist, ebola_sim$contacts,  
                        id="case_id", to="case_id", from="infectior",  
                        directed=TRUE)  
  
  ## subset based on cluster size range  
  x_subset <- subset_clusters_by_size(x, cs_min = 12, cs_max = 15)  
  
  ## subset based on single cluster size  
  x_subset <- subset_clusters_by_size(x, cs = 12)  
  
  ## subset based on minimum cluster size  
  x_subset <- subset_clusters_by_size(x, cs_min = 10)  
  
  ## subset based on maximum cluster size  
  x_subset <- subset_clusters_by_size(x, cs_max = 9)  
}
```

---

summary.epicontacts    *Summary method for epicontacts objects*

---

## Description

This method outputs a summary of the content of epicontacts objects.

## Usage

```
## S3 method for class 'epicontacts'  
summary(object, ...)
```

## Arguments

object            an [epicontacts](#) object  
...                further parameters to be passed to other methods (currently not used)

## Author(s)

VP Nagraj (<vpnagraj@virginia.edu>)

---

`thin`*Thin data to retain matching linelist / contacts*

---

## Description

This function can be used to remove ('thin') data from `epicontacts` objects to ensure stricter matching of linelists and contacts. It has two behaviours, triggered by the argument `what`: either it thins data from `$linelist`, keeping only cases that are in `$contacts` (`thin = "linelist"`, default), or the converse, i.e. removing contacts which are not fully documented in the linelist.

## Usage

```
thin(x, what = "linelist")
```

## Arguments

<code>x</code>	An <code>epicontacts</code> object.
<code>what</code>	A character string or integer determining which type of data is removed ('thinned'). "linelist" / 1 indicates that only cases appearing in <code>\$contacts</code> are kept in <code>\$linelist</code> . "contacts / 2" indicates that only cases appearing in <code>\$linelist</code> are kept in <code>\$contacts</code> .

## Author(s)

Thibaut Jombart (<thibautjombart@gmail.com>)

## Examples

```
if (require(outbreaks)) {  
  ## build data  
  x <- make_epicontacts(ebola_sim$linelist, ebola_sim$contacts,  
                        id = "case_id", to = "case_id", from = "infectior",  
                        directed = TRUE)  
  
  ## keep contacts from a specific case '916d0a'  
  x <- x[j = "916d0a", contacts = "from"]  
}
```

---

`transp`*Color tools and palettes for epicontacts*

---

**Description**

These functions are used for defining palettes or colors in the `epicontacts` package. They include:

**Usage**

```
transp(col, alpha = 0.5)
```

```
edges_pal(n)
```

```
cases_pal(n)
```

```
spectral(n)
```

```
fac2col(x, pal = cases_pal, NA_col = "lightgrey", legend = FALSE)
```

**Arguments**

<code>col</code>	A color vector to which transparency should be added.
<code>alpha</code>	The threshold to be used for transparency: 0 for full transparency, and 1 for full opacity.
<code>n</code>	An integer indicating the number of colors.
<code>x</code>	A character or a factor to be converted to colors.
<code>pal</code>	A color palette.
<code>NA_col</code>	The color to be used for NA values.
<code>legend</code>	A logical indicating if legend info should be added to the output. If TRUE, the output will be a list, with colors in the <code>\$color</code> component.

**Details**

- `cases_pal`: discrete color palette used for cases (comes from the `dibbler` package)
- `spectral`: continuous color palette (comes from the `adegetnet` package)
- `transp`: makes colors transparent (comes from the `adegetnet` package)
- `fac2col`: translates a character or a factor to a color using a palette (comes from the `adegetnet` package)

**Author(s)**

Thibaut Jombart <[thibautjombart@gmail.com](mailto:thibautjombart@gmail.com)>

## Examples

```
barplot(1:5, col = cases_pal(5))
barplot(1:50, col = cases_pal(50))
```

---

vis_epicontacts	<i>Plot epicontacts objects using visNetwork</i>
-----------------	--

---

## Description

This function plots [epicontacts](#) objects using the `visNetwork` package. The produced object is an `htmlwidget` which will need rendering within a web browser.

## Usage

```
vis_epicontacts(  
  x,  
  thin = TRUE,  
  node_color = "id",  
  label = "id",  
  annot = TRUE,  
  node_shape = NULL,  
  shapes = NULL,  
  edge_label = NULL,  
  edge_color = NULL,  
  legend = TRUE,  
  legend_max = 10,  
  x_axis = NULL,  
  col_pal = cases_pal,  
  NA_col = "lightgrey",  
  edge_col_pal = edges_pal,  
  width = "90%",  
  height = "700px",  
  selector = TRUE,  
  editor = FALSE,  
  edge_width = 3,  
  ...  
)
```

## Arguments

<code>x</code>	An <a href="#">epicontacts</a> object.
<code>thin</code>	A logical indicating if the data should be thinned with <code>thin</code> so that only cases with contacts should be plotted.
<code>node_color</code>	An index or character string indicating which field of the <code>linelist</code> should be used to color the nodes.

label	An index, logical, or character string indicating which fields of the linelist should be used for labelling the nodes. Logical will be recycled if necessary, so that the default TRUE effectively uses all columns of the linelist.
annot	An index, logical, or character string indicating which fields of the linelist should be used for annotating the nodes. Logical will be recycled if necessary, so that the default TRUE effectively uses all columns of the linelist.
node_shape	An index or character string indicating which field of the linelist should be used to determine the shapes of the nodes.
shapes	A named vector of characters indicating which icon code should be used for each value node_shape, e.g. <code>c(m = "male", f = "female")</code> if 'm' and 'f' are values from node_shape. See <a href="#">codeawesome</a> for all available codes.
edge_label	An index or character string indicating which field of the contacts data should be used to label the edges of the graph.
edge_color	An index or character string indicating which field of the contacts data should be used to color the edges of the graph.
legend	A logical indicating whether a legend should be added to the plot.
legend_max	The maximum number of groups for a legend to be displayed.
x_axis	A character string indicating which field of the linelist data should be used to specify the x axis position (must be numeric or Date)
col_pal	A color palette for the nodes.
NA_col	The color used for unknown group.
edge_col_pal	A color palette for the edges.
width	The width of the output, in html compatible format (e.g. '90%' or '800px').
height	The height of the output, in html compatible format (e.g. '800px').
selector	A logical indicating if the selector tool should be used; defaults to TRUE.
editor	A logical indicating if the editor tool should be used; defaults to FALSE.
edge_width	An integer indicating the width of the edges. Defaults to 3.
...	Further arguments to be passed to visNetwork.

**Value**

The same output as visNetwork.

**Author(s)**

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**See Also**

[visNetwork](#) in the package visNetwork. [edges\\_pal](#) and [cases\\_pal](#) for color palettes used

**Examples**

```

if (require(outbreaks)) {

  ## example using MERS outbreak in Korea, 2014
  head(mers_korea_2015[[1]])
  head(mers_korea_2015[[2]])

  x <- make_epicontacts(linelist=mers_korea_2015[[1]],
                       contacts = mers_korea_2015[[2]],
                       directed=TRUE)

  ## Not run:
  plot(x)
  plot(x, node_color = "place_infect")
  # show transmission tree with time as the horizontal axis, showing all nodes
  vis_epicontacts(x, x_axis = "dt_onset", thin = FALSE)
  plot(x, node_color = "loc_hosp", legend_max=20, annot=TRUE)
  plot(x, node_color = "loc_hosp", legend_max=20, annot=TRUE, x_axis = "dt_onset")
  plot(x, "place_infect", node_shape = "sex",
       shapes = c(M = "male", F = "female"))

  plot(x, "sex", node_shape = "sex", shapes = c(F = "female", M = "male"),
       edge_label = "exposure", edge_color = "exposure")

  ## End(Not run)
}

```

[.epicontacts

*Subset epicontacts objects based on case identifiers***Description**

The "[" operator can be used to subset `epicontacts` objects, retaining a specified set of case identifiers (*i* for the linelist, *j* for contacts). Note that unlike most classical R objects, there is no replacement method for `epicontacts` objects, i.e. no operations such as `foo[i] <- bar`.

**Usage**

```

## S3 method for class 'epicontacts'

x[
  i,
  j,
  k = TRUE,
  l = TRUE,
  contacts = c("both", "either", "from", "to"),
  ...
]

```

**Arguments**

x	An <a href="#">epicontacts</a> object
i	A character vector containing case ID to be retained in the linelist; alternatively, an integer or logical vector used to subset the rows of the <code>\$linelist</code> component.
j	A character vector containing case ID to be retained in the contacts; alternatively, an integer or logical vector used to subset the rows of the <code>\$contacts</code> component.
k	An integer, logical, or character vector subsetting the supplementary columns of <code>x\$linelist</code> , i.e. the columns after 'id'; i.e. <code>k=1</code> refers to the column immediately after 'id'.
l	An integer, logical, or character vector subsetting the supplementary columns of <code>x\$contacts</code> , i.e. the columns after 'from' and 'to'; i.e. <code>l=1</code> refers to the column immediately after 'to'.
contacts	A character string indicating the rules for retaining contacts when <code>j</code> indicates case IDs (see details).
...	Not used (there for compatibility with generic).

**Details**

Details on the 'contacts' argument; possible values are:

- 'both': contacts are retained only if both cases are in `j`
- 'either': contacts are retained if at least one of the cases is in `j`
- 'from': contacts are retained only if the source ('from') is in `j`
- 'to': contacts are retained only if the recipient ('to') is in `j`

**Author(s)**

Thibaut Jombart (<[thibautjombart@gmail.com](mailto:thibautjombart@gmail.com)>)

**See Also**

[thin](#) to retain matching cases in linelist or contacts.

**Examples**

```
if (require(outbreaks)) {
  ## build data
  x <- make_epicontacts(ebola_sim$linelist, ebola_sim$contacts,
                       id = "case_id", to = "case_id", from = "infectior",
                       directed = TRUE)

  ## subset first 10 linelist cases
  x[1:10]

  ## same, remove contacts
  x[1:10, j = FALSE]
```



```
## subset first 10 contacts
x[j = 1:10]

## remove the metadata
x[k = FALSE, j = FALSE]

## keep contacts where both cases are in linelist
x[j = get_id(x, "linelist"), contacts = "both"]

## keep contacts from a specific case '916d0a'
x[j = "916d0a", contacts = "from"]

## more complex: keep all cases and contacts with > 4 secondary contacts
## i) find cases to keep
temp <- table(x$contacts$from)
temp[temp > 4]
to.keep <- names(temp)[temp > 4]
to.keep

## subset the contacts
y <- x[j = to.keep, contacts = "either"]
y

## keep only relevant entries in the linelist
y <- thin(y)

## visualise result
plot(y)
}
```

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