Package ‘igraph’

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Version 1.3.5

Title Network Analysis and Visualization

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Description Routines for simple graphs and network analysis. It can handle large graphs very well and provides functions for generating random and regular graphs, graph visualization, centrality methods and much more.

Depends methods

Imports graphics, grDevices, magrittr, Matrix, pkgconfig (>= 2.0.0), rlang, stats, utils

Suggests ape, graph, igraphdata, rgl, scales, stats4, tcltk, testthat, withr, digest

License GPL (>= 2)


SystemRequirements gmp (optional), libxml2 (optional), glpk (>= 4.57, optional), C++11

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Encoding UTF-8

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igraph-package

The igraph package

Description

igraph is a library and R package for network analysis.

Introduction

The main goals of the igraph library is to provide a set of data types and functions for 1) pain-free implementation of graph algorithms, 2) fast handling of large graphs, with millions of vertices and edges, 3) allowing rapid prototyping via high level languages like R.

igraph graphs

igraph graphs have a class ‘igraph’. They are printed to the screen in a special format, here is an example, a ring graph created using make_ring:

```
IGRAPH U--- 10 10 -- Ring graph
+ attr: name (g/c), mutual (g/x), circular (g/x)
```

‘IGRAPH’ denotes that this is an igraph graph. Then come four bits that denote the kind of the graph: the first is ‘U’ for undirected and ‘D’ for directed graphs. The second is ‘N’ for named graph (i.e. if the graph has the ‘name’ vertex attribute set). The third is ‘W’ for weighted graphs (i.e. if the ‘weight’ edge attribute is set). The fourth is ‘B’ for bipartite graphs (i.e. if the ‘type’ vertex attribute is set).

Then come two numbers, the number of vertices and the number of edges in the graph, and after a double dash, the name of the graph (the ‘name’ graph attribute) is printed if present. The second line is optional and it contains all the attributes of the graph. This graph has a ‘name’ graph attribute, of type character, and two other graph attributes called ‘mutual’ and ‘circular’, of a complex type. A complex type is simply anything that is not numeric or character. See the documentation of print.igraph for details.

If you want to see the edges of the graph as well, then use the print_all function:
> print_all(g)
IGRAPH badcafe U--- 10 10 -- Ring graph
+ attr: name (g/c), mutual (g/x), circular (g/x)
+ edges:
[1] 1-- 2 2-- 3 3-- 4 4-- 5 5-- 6 6-- 7 7-- 8 8-- 9 9--10 1--10

Creating graphs

There are many functions in igraph for creating graphs, both deterministic and stochastic; stochastic graph constructors are called ‘games’.

To create small graphs with a given structure probably the graph_from_literal function is easiest. It uses R’s formula interface, its manual page contains many examples. Another option is graph, which takes numeric vertex ids directly. graph_from_atlas creates graph from the Graph Atlas, make_graph can create some special graphs.

To create graphs from field data, graph_from_edgelist, graph_from_data_frame and graph_from_adjacency_matrix are probably the best choices.

The igraph package includes some classic random graphs like the Erdos-Renyi GNP and GNM graphs (sample_gnp, sample_gnm) and some recent popular models, like preferential attachment (sample_pa) and the small-world model (sample_smallworld).

Vertex and edge IDs

Vertices and edges have numerical vertex ids in igraph. Vertex ids are always consecutive and they start with one. I.e. for a graph with \( n \) vertices the vertex ids are between 1 and \( n \). If some operation changes the number of vertices in the graphs, e.g. a subgraph is created via induced_subgraph, then the vertices are renumbered to satisfy this criteria.

The same is true for the edges as well, edge ids are always between one and \( m \), the total number of edges in the graph.

It is often desirable to follow vertices along a number of graph operations, and vertex ids don’t allow this because of the renumbering. The solution is to assign attributes to the vertices. These are kept by all operations, if possible. See more about attributes in the next section.

Attributes

In igraph it is possible to assign attributes to the vertices or edges of a graph, or to the graph itself. igraph provides flexible constructs for selecting a set of vertices or edges based on their attribute values, see vertex_attr, V and E for details.

Some vertex/edge/graph attributes are treated specially. One of them is the ‘name’ attribute. This is used for printing the graph instead of the numerical ids, if it exists. Vertex names can also be used to specify a vector or set of vertices, in all igraph functions. E.g. degree has a \( v \) argument that gives the vertices for which the degree is calculated. This argument can be given as a character vector of vertex names.

Edges can also have a ‘name’ attribute, and this is treated specially as well. Just like for vertices, edges can also be selected based on their names, e.g. in the delete_edges and other functions.

We note here, that vertex names can also be used to select edges. The form ‘from|to’, where ‘from’ and ‘to’ are vertex names, select a single, possibly directed, edge going from ‘from’ to ‘to’. The two forms can also be mixed in the same edge selector.

Other attributes define visualization parameters, see igraph.plotting for details.

Attribute values can be set to any R object, but note that storing the graph in some file formats might result the loss of complex attribute values. All attribute values are preserved if you use save and load to store/retrieve your graphs.
Visualization

igraph provides three different ways for visualization. The first is the `plot.igraph` function. (Actually you don’t need to write `plot.igraph`, `plot` is enough. This function uses regular R graphics and can be used with any R device.

The second function is `tkplot`, which uses a Tk GUI for basic interactive graph manipulation. (Tk is quite resource hungry, so don’t try this for very large graphs.)

The third way requires the rgl package and uses OpenGL. See the `rglplot` function for the details.

Make sure you read `igraph.plotting` before you start plotting your graphs.

File formats

igraph can handle various graph file formats, usually both for reading and writing. We suggest that you use the GraphML file format for your graphs, except if the graphs are too big. For big graphs a simpler format is recommended. See `read_graph` and `write_graph` for details.

Further information

The igraph homepage is at https://igraph.org. See especially the documentation section. Join the discussion forum at https://igraph.discourse.group if you have questions or comments.

---

`.apply_modifiers` Applies a set of constructor modifiers to an already constructed graph.

Description

This is a helper function for the common parts of `make_` and `sample_`.

Usage

```
.apply_modifiers(graph, mods)
```

Arguments

- `graph` The graph to apply the modifiers to
- `mods` The modifiers to apply

Value

The modified graph
.extract_constructor_and_modifiers

Takes an argument list and extracts the constructor specification and constructor modifiers from it.

Description

This is a helper function for the common parts of make_ and sample_.

Usage

.extract_constructor_and_modifiers(..., .operation, .variant)

Arguments

... Parameters to extract from
.operation Human-readable description of the operation that this helper is a part of
.variant Constructor variant; must be one of ‘make’, ‘graph’ or ‘sample’. Used in cases when the same constructor specification has deterministic and random variants.

Value

A named list with three items: ‘cons’ for the constructor function, ‘mods’ for the modifiers and ‘args’ for the remaining, unparsed arguments.

[.igraph

Query and manipulate a graph as it were an adjacency matrix

Description

Query and manipulate a graph as it were an adjacency matrix

Usage

## S3 method for class ‘igraph’

x[
  i,
  j,
  ..., from,
  to,
  sparse = igraph_opt("sparsematrices"),
  edges = FALSE,
  drop = TRUE,
  attr = if (is_weighted(x)) "weight" else NULL
]
Arguments

- **x**: The graph.
- **i**: Index. Vertex ids or names or logical vectors. See details below.
- **j**: Index. Vertex ids or names or logical vectors. See details below.
- **from**: A numeric or character vector giving vertex ids or names. Together with the `to` argument, it can be used to query/set a sequence of edges. See details below. This argument cannot be present together with any of the `i` and `j` arguments and if it is present, then the `to` argument must be present as well.
- **to**: A numeric or character vector giving vertex ids or names. Together with the `from` argument, it can be used to query/set a sequence of edges. See details below. This argument cannot be present together with any of the `i` and `j` arguments and if it is present, then the `from` argument must be present as well.
- **sparse**: Logical scalar, whether to return sparse matrices.
- **edges**: Logical scalar, whether to return edge ids.
- **drop**: Ignored.
- **attr**: If not `NULL`, then it should be the name of an edge attribute. This attribute is queried and returned.

Details

The single bracket indexes the (possibly weighted) adjacency matrix of the graph. Here is what you can do with it:

1. Check whether there is an edge between two vertices \((v \text{ and } w)\) in the graph:
   \[
   \text{graph}[v, w]
   \]
   A numeric scalar is returned, one if the edge exists, zero otherwise.

2. Extract the (sparse) adjacency matrix of the graph, or part of it:
   \[
   \text{graph}[,]
   \]
   \[
   \text{graph}[1:3, 5:6]
   \]
   \[
   \text{graph}[c(1,3,5),]
   \]
   The first variants returns the full adjacency matrix, the other two return part of it.

3. The `from` and `to` arguments can be used to check the existence of many edges. In this case, both `from` and `to` must be present and they must have the same length. They must contain vertex ids or names. A numeric vector is returned, of the same length as the `from` argument, it contains ones for existing edges and zeros for non-existing ones. Example:
   \[
   \text{graph[from=1:3, to=c(2,3,5)]
   \]

4. For weighted graphs, the `[ operator returns the edge weights. For non-existent edges zero weights are returned. Other edge attributes can be queried as well, by giving the `attr` argument.

5. Querying edge ids instead of the existence of edges or edge attributes. E.g.
   \[
   \text{graph}[1, 2, \text{ edges=TRUE}]
   \]
   returns the id of the edge between vertices 1 and 2, or zero if there is no such edge.
6. Adding one or more edges to a graph. For this the element(s) of the imaginary adjacency matrix must be set to a non-zero numeric value (or TRUE):

```r
graph[1, 2] <- 1
graph[1:3, 1] <- 1
graph[from=1:3, to=c(2,3,5)] <- TRUE
```

This does not affect edges that are already present in the graph, i.e. no multiple edges are created.

7. Adding weighted edges to a graph. The `attr` argument contains the name of the edge attribute to set, so it does not have to be "weight":

```r
graph[1, 2, attr="weight"] <- 5
graph[from=1:3, to=c(2,3,5)] <- c(1,-1,4)
```

If an edge is already present in the network, then only its weights or other attribute are updated. If the graph is already weighted, then the `attr="weight"` setting is implicit, and one does not need to give it explicitly.

8. Deleting edges. The replacement syntax allow the deletion of edges, by specifying `FALSE` or `NULL` as the replacement value:

```r
graph[v, w] <- FALSE
```

removes the edge from vertex `v` to vertex `w`. As this can be used to delete edges between two sets of vertices, either pairwise:

```r
graph[from=v, to=w] <- FALSE
```

or not:

```r
graph[v, w] <- FALSE
```

if `v` and `w` are vectors of edge ids or names.

`[]` allows logical indices and negative indices as well, with the usual R semantics. E.g.

```r
graph[degree(graph)==0, 1] <- 1
```

adds an edge from every isolate vertex to vertex one, and

```r
G <- make_empty_graph(10)
G[-1,1] <- TRUE
```

creates a star graph.

Of course, the indexing operators support vertex names, so instead of a numeric vertex id a vertex can also be given to `[` and `[[`.

**Value**

A scalar or matrix. See details below.

**See Also**

Other structural queries: `[[.igraph()`, `adjacent_vertices()`, `are_adjacent()`, `ends()`, `get.edge.ids()`, `gorder()`, `gsize()`, `head_of()`, `incident_edges()`, `incident()`, `is_directed()`, `neighbors()`, `tail_of()`
Query and manipulate a graph as it were an adjacency list

Description
Query and manipulate a graph as it were an adjacency list

Usage
## S3 method for class 'igraph'
x[[i, j, from, to, ..., directed = TRUE, edges = FALSE, exact = TRUE]]

Arguments
- x: The graph.
- i: Index, integer, character or logical, see details below.
- j: Index, integer, character or logical, see details below.
- from: A numeric or character vector giving vertex ids or names. Together with the to argument, it can be used to query/set a sequence of edges. See details below. This argument cannot be present together with any of the i and j arguments and if it is present, then the to argument must be present as well.
- to: A numeric or character vector giving vertex ids or names. Together with the from argument, it can be used to query/set a sequence of edges. See details below. This argument cannot be present together with any of the i and j arguments and if it is present, then the from argument must be present as well.
- ...: Additional arguments are not used currently.
- directed: Logical scalar, whether to consider edge directions in directed graphs. It is ignored for undirected graphs.
- edges: Logical scalar, whether to return edge ids.
- exact: Ignored.

Details
The double bracket operator indexes the (imaginary) adjacency list of the graph. This can used for the following operations:

1. Querying the adjacent vertices for one or more vertices:
   ```r
graph[[1:3,]]
graph[,1:3]]
   ```
   The first form gives the successors, the second the predecessors or the 1:3 vertices. (For undirected graphs they are equivalent.)

2. Querying the incident edges for one or more vertices, if the edges argument is set to TRUE:
   ```r
graph[[1:3, , edges=TRUE]]
graph[, 1:3, edges=TRUE]]
   ```

3. Querying the edge ids between two sets or vertices, if both indices are used. E.g.
   ```r
graph[[v, w, edges=TRUE]]
   ```
gives the edge ids of all the edges that exist from vertices \( v \) to vertices \( w \).

The alternative argument names \( \text{from} \) and \( \text{to} \) can be used instead of the usual \( i \) and \( j \), to make the code more readable:

```r
graph[[from = 1:3]]
graph[[from = v, to = w, edges = TRUE]]
```

`
` operators allows logical indices and negative indices as well, with the usual R semantics.

Vertex names are also supported, so instead of a numeric vertex id a vertex can also be given to ‘\[\]’ and ‘\[[\]’.

**See Also**

Other structural queries: \(.\text{igraph}()\), \.\text{adjacent_vertices}()\), \.\text{are_adjacent}()\), \.\text{ends}()\), \.\text{get.edge.ids}()\), \.\text{gorder}()\), \.\text{gsize}()\), \.\text{head_of}()\), \.\text{incident_edges}()\), \.\text{incident}()\), \.\text{is_directed}()\), \.\text{neighbors}()\), \.\text{tail_of}()

---

### Magrittr’s pipes

**Description**

igraph re-exports the %>% operator of magrittr, because we find it very useful. Please see the documentation in the magrittr package.

**Arguments**

- `lhs`: Left hand side of the pipe.
- `rhs`: Right hand side of the pipe.

**Value**

Result of applying the right hand side to the result of the left hand side.

**Examples**

```r
make_ring(10) %>%
add_edges(c(1,6)) %>%
plot()
```
+igraph

Add vertices, edges or another graph to a graph

Description

Add vertices, edges or another graph to a graph

Usage

## S3 method for class 'igraph'
e1 + e2

Arguments

e1  First argument, probably an igraph graph, but see details below.
e2  Second argument, see details below.

Details

The plus operator can be used to add vertices or edges to graph. The actual operation that is performed depends on the type of the right hand side argument:

- If is is another igraph graph object and they are both named graphs, then the union of the two graphs are calculated, see union.
- If it is another igraph graph object, but either of the two are not named, then the disjoint union of the two graphs is calculated, see disjoint_union.
- If it is a numeric scalar, then the specified number of vertices are added to the graph.
- If it is a character scalar or vector, then it is interpreted as the names of the vertices to add to the graph.
- If it is an object created with the vertex or vertices function, then new vertices are added to the graph. This form is appropriate when one wants to add some vertex attributes as well. The operands of the vertices function specifies the number of vertices to add and their attributes as well. The unnamed arguments of vertices are concatenated and used as the 'name' vertex attribute (i.e. vertex names), the named arguments will be added as additional vertex attributes. Examples:

  g <- g +
  vertex(shape="circle", color= "red")
  g <- g + vertex("foo", color="blue")
  g <- g + vertex("bar", "foobar")
  g <- g + vertices("bar2", "foobar2", color=1:2, shape="rectangle")

  vertex is just an alias to vertices, and it is provided for readability. The user should use it if a single vertex is added to the graph.
- If it is an object created with the edge or edges function, then new edges will be added to the graph. The new edges and possibly their attributes can be specified as the arguments of the edges function. The unnamed arguments of edges are concatenated and used as vertex ids of the end points of the new edges. The named arguments will be added as edge attributes. Examples:
g <- make_empty_graph() +
   vertices(letters[1:10]) +
   vertices("foo", "bar", "bar2", "foobar2")
g <- g + edge("a", "b")
g <- g + edges("foo", "bar", "bar2", "foobar2")
g <- g + edges(c("bar", "foo", "foobar2", "bar2"), color="red", weight=1:2)

See more examples below.

edge is just an alias to edges and it is provided for readability. The user should use it if a single edge is added to the graph.

• If it is an object created with the path function, then new edges that form a path are added.
The edges and possibly their attributes are specified as the arguments to the path function.
The non-named arguments are concatenated and interpreted as the vertex ids along the path.
The remaining arguments are added as edge attributes.
Examples:

g <- make_empty_graph() + vertices(letters[1:10])
g <- g + path("a", "b", "c", "d")
g <- g + path("e", "f", "g", weight=1:2, color="red")
g <- g + path(c("f", "c", "j", "d"), width=1:3, color="green")

It is important to note that, although the plus operator is commutative, i.e. is possible to write

graph <- "foo" + make_empty_graph()

it is not associative, e.g.

graph <- "foo" + "bar" + make_empty_graph()

results a syntax error, unless parentheses are used:

graph <- "foo" + ("bar" + make_empty_graph())

For clarity, we suggest to always put the graph object on the left hand side of the operator:

graph <- make_empty_graph() + "foo" + "bar"

See Also

Other functions for manipulating graph structure: add_edges(), add_vertices(), delete_edges(), delete_vertices(), edge(), igraph-minus, path(), vertex()

Examples

# 10 vertices named a,b,c,... and no edges
g <- make_empty_graph() + vertices(letters[1:10])

# Add edges to make it a ring
g <- g + path(letters[1:10], letters[1], color = "grey")

# Add some extra random edges

g <- g + edges(sample(V(g), 10, replace = TRUE), color = "red")
g$layout <- layout_in_circle
plot(g)
**Description**

The new edges are given as a vertex sequence, e.g. internal numeric vertex ids, or vertex names. The first edge points from edges[1] to edges[2], the second from edges[3] to edges[4], etc.

**Usage**

```r
add_edges(graph, edges, ..., attr = list())
```

**Arguments**

- `graph`: The input graph
- `edges`: The edges to add, a vertex sequence with even number of vertices.
- `...`: Additional arguments, they must be named, and they will be added as edge attributes, for the newly added edges. See also details below.
- `attr`: A named list, its elements will be added as edge attributes, for the newly added edges. See also details below.

**Details**

If attributes are supplied, and they are not present in the graph, their values for the original edges of the graph are set to `NA`.

**Value**

The graph, with the edges (and attributes) added.

**See Also**

Other functions for manipulating graph structure: `+`, `.igraph`, `add_vertices`, `delete_edges`, `delete_vertices`, `edge`, `igraph-minus`, `path`, `vertex`

**Examples**

```r
g <- make_empty_graph(n = 5) %>%
  add_edges(c(1,2, 2,3, 3,4, 4,5)) %>%
  set_edge_attr("color", value = "red") %>%
  add_edges(c(5,1), color = "green")
E(g)[[1]]
plot(g)
```
add_layout_  Add layout to graph

Description
Add layout to graph

Usage
add_layout_(graph, ..., overwrite = TRUE)

Arguments
graph       The input graph.
...         Additional arguments are passed to layout_.
overwrite   Whether to overwrite the layout of the graph, if it already has one.

Value
The input graph, with the layout added.

See Also
layout_ for a description of the layout API.
Other graph layouts: component_wise(), layout_as_bipartite(), layout_as_star(), layout_as_tree(),
layout_in_circle(), layout_nicely(), layout_on_grid(), layout_on_sphere(), layout_randomly(),
layout_with_dh(), layout_with_fr(), layout_with_gem(), layout_with_graphopt(), layout_with_kk(),
layout_with_lgl(), layout_with_mds(), layout_with_sugiyama(), layout_(), merge_coords(),
norm_coords(), normalize()

Examples
(make_star(11) + make_star(11)) %>%
  add_layout_(as_star(), component_wise()) %>%
  plot()

add_vertices  Add vertices to a graph

Description
If attributes are supplied, and they are not present in the graph, their values for the original vertices
of the graph are set to NA.

Usage
add_vertices(graph, nv, ..., attr = list())
adjacent_vertices

Arguments

graph The input graph.

nv The number of vertices to add.

... Additional arguments, they must be named, and they will be added as vertex attributes, for the newly added vertices. See also details below.

attr A named list, its elements will be added as vertex attributes, for the newly added vertices. See also details below.

Value

The graph, with the vertices (and attributes) added.

See Also

Other functions for manipulating graph structure: +.igraph(), add_edges(), delete_edges(), delete_vertices(), edge(), igraph-minus, path(), vertex()

Examples

g <- make_empty_graph() %>%
  add_vertices(3, color = "red") %>%
  add_vertices(2, color = "green") %>%
  add_edges(c(1, 2, 3, 4, 5))

g V(g)[[3]]
plot(g)

adjacent_vertices

Adjacent vertices of multiple vertices in a graph

Description

This function is similar to neighbors, but it queries the adjacent vertices for multiple vertices at once.

Usage

adjacent_vertices(graph, v, mode = c("out", "in", "all", "total"))

Arguments

graph Input graph.

v The vertices to query.

mode Whether to query outgoing (‘out’), incoming (‘in’) edges, or both types (‘all’). This is ignored for undirected graphs.

Value

A list of vertex sequences.
See Also

Other structural queries: [.igraph(), [[.igraph(), are_adjacent(), ends(), get.edge.ids(), gorder(), gsize(), head.of(), incident_edges(), incident(), is.directed(), neighbors(), tail.of()]

Examples

```r
g <- make_graph("Zachary")
adjacent_vertices(g, c(1, 34))
```

---

**all_simple_paths**

List all simple paths from one source

**Description**

This function lists all simple paths from one source vertex to another vertex or vertices. A path is simple if the vertices it visits are not visited more than once.

**Usage**

```r
all_simple_paths(
  graph,
  from,
  to = V(graph),
  mode = c("out", "in", "all", "total"),
  cutoff = -1
)
```

**Arguments**

- `graph`: The input graph.
- `from`: The source vertex.
- `to`: The target vertex of vertices. Defaults to all vertices.
- `mode`: Character constant, gives whether the shortest paths to or from the given vertices should be calculated for directed graphs. If `out` then the shortest paths from the vertex, if `in` then `to` it will be considered. If `all`, the default, then the corresponding undirected graph will be used, i.e. not directed paths are searched. This argument is ignored for undirected graphs.
- `cutoff`: Maximum length of path that is considered. If negative, paths of all lengths are considered.

**Details**

Note that potentially there are exponentially many paths between two vertices of a graph, and you may run out of memory when using this function, if your graph is lattice-like.

This function currently ignores multiple and loop edges.

**Value**

A list of integer vectors, each integer vector is a path from the source vertex to one of the target vertices. A path is given by its vertex ids.
Examples

```r
  g <- make_ring(10)
  all_simple_paths(g, 1, 5)
  all_simple_paths(g, 1, c(3,5))
```

### alpha_centrality

Find Bonacich alpha centrality scores of network positions

#### Description

alpha_centrality calculates the alpha centrality of some (or all) vertices in a graph.

#### Usage

```r
alpha_centrality(
  graph, 
  nodes = V(graph), 
  alpha = 1, 
  loops = FALSE, 
  exo = 1, 
  weights = NULL, 
  tol = 1e-07, 
  sparse = TRUE
)
```

#### Arguments

- **graph**  
The input graph, can be directed or undirected
- **nodes**  
Vertex sequence, the vertices for which the alpha centrality values are returned. (For technical reasons they will be calculated for all vertices, anyway.)
- **alpha**  
Parameter specifying the relative importance of endogenous versus exogenous factors in the determination of centrality. See details below.
- **loops**  
Whether to eliminate loop edges from the graph before the calculation.
- **exo**  
The exogenous factors, in most cases this is either a constant – the same factor for every node, or a vector giving the factor for every vertex. Note that too long vectors will be truncated and too short vectors will be replicated to match the number of vertices.
- **weights**  
A character scalar that gives the name of the edge attribute to use in the adjacency matrix. If it is NULL, then the ‘weight’ edge attribute of the graph is used, if there is one. Otherwise, or if it is NA, then the calculation uses the standard adjacency matrix.
- **tol**  
Tolerance for near-singularities during matrix inversion, see solve.
- **sparse**  
Logical scalar, whether to use sparse matrices for the calculation. The ‘Matrix’ package is required for sparse matrix support
Details

The alpha centrality measure can be considered as a generalization of eigenvector centrality to directed graphs. It was proposed by Bonacich in 2001 (see reference below).

The alpha centrality of the vertices in a graph is defined as the solution of the following matrix equation:

\[ x = \alpha A^T x + e, \]

where \( A \) is the (not necessarily symmetric) adjacency matrix of the graph, \( e \) is the vector of exogenous sources of status of the vertices and \( \alpha \) is the relative importance of the endogenous versus exogenous factors.

Value

A numeric vector containing the centrality scores for the selected vertices.

Warning

Singular adjacency matrices cause problems for this algorithm, the routine may fail in certain cases.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

eigen_centrality and power_centrality

Examples

# The examples from Bonacich's paper
g.1 <- graph(c(1,3,2,3,4,5))
g.2 <- graph(c(2,1,3,1,4,1))
g.3 <- graph(c(1,2,3,4,1,5,1))
alpha_centrality(g.1)
alpa_centrality(g.2)
alpa_centrality(g.3,alpha=0.5)

Description

Are two vertices adjacent?

The order of the vertices only matters in directed graphs, where the existence of a directed \((v1, v2)\) edge is queried.
Usage

are_adjacent(graph, v1, v2)

Arguments

graph The graph.
v1 The first vertex, tail in directed graphs.
v2 The second vertex, head in directed graphs.

Value

A logical scalar, TRUE is a (v1, v2) exists in the graph.

See Also

Other structural queries: [[igraph(). adjacent_vertices(). ends(). get.edge.ids(), gorder(). gsize(). head_of(). incident_edges(). incident(). is_directed(). neighbors(). tail_of()]

Examples

ug <- make_ring(10)
ug
are_adjacent(ug, 1, 2)
are_adjacent(ug, 2, 1)

dg <- make_ring(10, directed = TRUE)
dg
are_adjacent(ug, 1, 2)
are_adjacent(ug, 2, 1)

arpack_defaults ARPACK eigenvector calculation

Description

Interface to the ARPACK library for calculating eigenvectors of sparse matrices

Usage

arpack_defaults

arpack(  
  func,  
  extra = NULL,  
  sym = FALSE,  
  options = arpack_defaults,  
  env = parent.frame(),  
  complex = !sym  
)

Arguments

func  The function to perform the matrix-vector multiplication. ARPACK requires to perform these by the user. The function gets the vector \( x \) as the first argument, and it should return \( Ax \), where \( A \) is the “input matrix”. (The input matrix is never given explicitly.) The second argument is extra.

eextra  Extra argument to supply to func.

sym  Logical scalar, whether the input matrix is symmetric. Always supply TRUE here if it is, since it can speed up the computation.

options  Options to ARPACK, a named list to overwrite some of the default option values. See details below.

eenv  The environment in which func will be evaluated.

ccomplex  Whether to convert the eigenvectors returned by ARPACK into R complex vectors. By default this is not done for symmetric problems (these only have real eigenvectors/values), but only non-symmetric ones. If you have a non-symmetric problem, but you’re sure that the results will be real, then supply FALSE here.

Format

An object of class list of length 14.

Details

ARPACK is a library for solving large scale eigenvalue problems. The package is designed to compute a few eigenvalues and corresponding eigenvectors of a general \( n \) by \( n \) matrix \( A \). It is most appropriate for large sparse or structured matrices \( A \) where structured means that a matrix-vector product \( w \leftarrow Av \) requires order \( n \) rather than the usual order \( n^2 \) floating point operations.

This function is an interface to ARPACK. igraph does not contain all ARPACK routines, only the ones dealing with symmetric and non-symmetric eigenvalue problems using double precision real numbers.

The eigenvalue calculation in ARPACK (in the simplest case) involves the calculation of the \( Av \) product where \( A \) is the matrix we work with and \( v \) is an arbitrary vector. The function supplied in the fun argument is expected to perform this product. If the product can be done efficiently, e.g. if the matrix is sparse, then arpack is usually able to calculate the eigenvalues very quickly.

The options argument specifies what kind of calculation to perform. It is a list with the following members, they correspond directly to ARPACK parameters. On input it has the following fields:

bmat  Character constant, possible values: ‘T’, standard eigenvalue problem, \( Ax = \lambda x \); and ‘G’, generalized eigenvalue problem, \( Ax = \lambda Bx \). Currently only ‘T’ is supported.

n  Numeric scalar. The dimension of the eigenproblem. You only need to set this if you call arpack directly. (i.e. not needed for eigen_centrality, page_rank, etc.)

which  Specify which eigenvalues/vectors to compute, character constant with exactly two characters.

Possible values for symmetric input matrices:

"LA"  Compute nev largest (algebraic) eigenvalues.

"SA"  Compute nev smallest (algebraic) eigenvalues.

"LM"  Compute nev largest (in magnitude) eigenvalues.

"SM"  Compute nev smallest (in magnitude) eigenvalues.
"BE" Compute nev eigenvalues, half from each end of the spectrum. When nev is odd, compute one more from the high end than from the low end.

Possible values for non-symmetric input matrices:

"LM" Compute nev eigenvalues of largest magnitude.
"SM" Compute nev eigenvalues of smallest magnitude.
"LR" Compute nev eigenvalues of largest real part.
"SR" Compute nev eigenvalues of smallest real part.
"LI" Compute nev eigenvalues of largest imaginary part.
"SI" Compute nev eigenvalues of smallest imaginary part.

This parameter is sometimes overwritten by the various functions, e.g. page_rank always sets 'LM'.

nev Numeric scalar. The number of eigenvalues to be computed.

tol Numeric scalar. Stopping criterion: the relative accuracy of the Ritz value is considered acceptable if its error is less than tol times its estimated value. If this is set to zero then machine precision is used.

cnev Number of Lanczos vectors to be generated.

ldv Numeric scalar. It should be set to zero in the current implementation.

ishift Either zero or one. If zero then the shifts are provided by the user via reverse communication. If one then exact shifts with respect to the reduced tridiagonal matrix T. Please always set this to one.

maxiter Maximum number of Arnoldi update iterations allowed.

nb Blocksize to be used in the recurrence. Please always leave this on the default value, one.

mode The type of the eigenproblem to be solved. Possible values if the input matrix is symmetric:

1 $Ax = \lambda x$, $A$ is symmetric.
2 $Ax = \lambda Mx$, $A$ is symmetric, $M$ is symmetric positive definite.
3 $Kx = \lambda Mx$, $K$ is symmetric, $M$ is symmetric positive semi-definite.
4 $Kx = \lambda KGx$, $K$ is symmetric positive semi-definite, $KG$ is symmetric indefinite.
5 $Ax = \lambda Mx$, $A$ is symmetric, $M$ is symmetric positive semi-definite. (Cayley transformed mode.)

Please note that only mode==1 was tested and other values might not work properly.

Possible values if the input matrix is not symmetric:

1 $Ax = \lambda x$.
2 $Ax = \lambda Mx$, $M$ is symmetric positive definite.
3 $Ax = \lambda Mx$, $M$ is symmetric semi-definite.
4 $Ax = \lambda Mx$, $M$ is symmetric semi-definite.

Please note that only mode==1 was tested and other values might not work properly.

start Not used currently. Later it be used to set a starting vector.

sigma Not used currently.

sigmai Not use currently.

On output the following additional fields are added:

info Error flag of ARPACK. Possible values:
0 Normal exit.
1 Maximum number of iterations taken.
No shifts could be applied during a cycle of the Implicitly restarted Arnoldi iteration. One possibility is to increase the size of \texttt{ncv} relative to \texttt{nev}. ARPACK can return more error conditions than these, but they are converted to regular igraph errors.

- **\texttt{iter}**: Number of Arnoldi iterations taken.
- **\texttt{nconv}**: Number of “converged” Ritz values. This represents the number of Ritz values that satisfy the convergence criterion.
- **\texttt{numop}**: Total number of matrix-vector multiplications.
- **\texttt{numopb}**: Not used currently.
- **\texttt{numreo}**: Total number of steps of re-orthogonalization.

Please see the ARPACK documentation for additional details.

**Value**

A named list with the following members:

- **values**: Numeric vector, the desired eigenvalues.
- **vectors**: Numeric matrix, the desired eigenvectors as columns. If \texttt{complex=TRUE} (the default for non-symmetric problems), then the matrix is complex.
- **options**: A named list with the supplied options and some information about the performed calculation, including an ARPACK exit code. See the details above.

**Author(s)**

Rich Lehoucq, Kristi Maschhoff, Danny Sorensen, Chao Yang for ARPACK, Gabor Csardi \texttt{<csardi.gabor@gmail.com>} for the R interface.

**References**


**See Also**

- `eigen_centrality`
- `page_rank`
- `hub_score`
- `cluster_leading_eigen` are some of the functions in igraph that use ARPACK.

**Examples**

```r
# Identity matrix
f <- function(x, extra=NULL) x
arpack(f, options=list(n=10, nev=2, ncv=4), sym=TRUE)

# Graph laplacian of a star graph (undirected), n>=2
# Note that this is a linear operation
f <- function(x, extra=NULL) {
  y <- x
  y[1] <- (length(x)-1)*x[1] - sum(x[-1])
}
for (i in 2:length(x)) {
  y[i] <- x[i] - x[1]
}
y
arpack(f, options=list(n=10, nev=1, ncv=3), sym=TRUE)

# double check
eigen(laplacian_matrix(make_star(10, mode="undirected")))

## First three eigenvalues of the adjacency matrix of a graph
## We need the 'Matrix' package for this
if (require(Matrix)) {
  set.seed(42)
  g <- sample_gnp(1000, 5/1000)
  M <- as_adj(g, sparse=TRUE)
  f2 <- function(x, extra=NULL) { cat("."); as.vector(M %*% x) }
  baev <- arpack(f2, sym=TRUE, options=list(n=vcount(g), nev=3, ncv=8, which="LM", maxiter=2000))
}

articulation_points

articulation_points  Articulation points and bridges of a graph

Description

articulation_points finds the articulation points (or cut vertices)

Usage

articulation_points(graph)

bridges(graph)

Arguments

graph The input graph. It is treated as an undirected graph, even if it is directed.

Details

Articulation points or cut vertices are vertices whose removal increases the number of connected components in a graph. Similarly, bridges or cut-edges are edges whose removal increases the number of connected components in a graph. If the original graph was connected, then the removal of a single articulation point or a single bridge makes it undirected. If a graph contains no articulation points, then its vertex connectivity is at least two.

Value

For articulation_points, a numeric vector giving the vertex IDs of the articulation points of the input graph. For bridges, a numeric vector giving the edge IDs of the bridges of the input graph.
as.directed

Convert between directed and undirected graphs

Description

as.directed converts an undirected graph to directed, as.undirected does the opposite, it converts a directed graph to undirected.

Usage

as.directed(graph, mode = c("mutual", "arbitrary", "random", "acyclic"))

as.undirected(
  graph,
  mode = c("collapse", "each", "mutual"),
  edge.attr.comb = igraph_opt("edge.attr.comb")
)

Arguments

graph The graph to convert.
mode Character constant, defines the conversion algorithm. For as.directed it can be mutual or arbitrary. For as.undirected it can be each, collapse or mutual. See details below.
edge.attr.comb Specifies what to do with edge attributes, if mode="collapse" or mode="mutual". In these cases many edges might be mapped to a single one in the new graph, and their attributes are combined. Please see attribute.combination for details on this.

Examples

g <- disjoint_union( make_full_graph(5), make_full_graph(5) )
clu <- components(g)$membership
g <- add_edges(g, c(match(1, clu), match(2, clu)) )
articulation_points(g)

g <- make_graph("krackhardt_kite")
bridges(g)
Details
Conversion algorithms for \texttt{as.directed}:

- **"arbitrary"** The number of edges in the graph stays the same, an arbitrarily directed edge is created for each undirected edge, but the direction of the edge is deterministic (i.e. it always points the same way if you call the function multiple times).

- **"mutual"** Two directed edges are created for each undirected edge, one in each direction.

- **"random"** The number of edges in the graph stays the same, and a randomly directed edge is created for each undirected edge. You will get different results if you call the function multiple times with the same graph.

- **"acyclic"** The number of edges in the graph stays the same, and a directed edge is created for each undirected edge such that the resulting graph is guaranteed to be acyclic. This is achieved by ensuring that edges always point from a lower index vertex to a higher index. Note that the graph may include cycles of length 1 if the original graph contained loop edges.

Conversion algorithms for \texttt{as.undirected}:

- **"each"** The number of edges remains constant, an undirected edge is created for each directed one, this version might create graphs with multiple edges.

- **"collapse"** One undirected edge will be created for each pair of vertices which are connected with at least one directed edge, no multiple edges will be created.

- **"mutual"** One undirected edge will be created for each pair of mutual edges. Non-mutual edges are ignored. This mode might create multiple edges if there are more than one mutual edge pairs between the same pair of vertices.

Value
A new graph object.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

See Also
\texttt{simplify} for removing multiple and/or loop edges from a graph.

Examples

```r
# Creating a directed graph
g <- make_ring(10)
as.directed(g, "mutual")
g2 <- make_star(10)
as.undirected(g)

# Combining edge attributes
g3 <- make_ring(10, directed=TRUE, mutual=TRUE)
E(g3)$weight <- seq_len(ecount(g3))
ug3 <- as.undirected(g3)
print(ug3, e=TRUE)
```

## Not run:
```r
x11(width=10, height=5)
layout(rbind(1:2))
plot(g3, layout=layout_in_circle, edge.label=E(g3)$weight)
plot(ug3, layout=layout_in_circle, edge.label=E(ug3)$weight)
```
```r
## End(Not run)

g4 <- graph(c(1,2, 3,2,3,4, 5,4,5, 6,7, 7,6,7,8, 8,7,8,9, 9,8,9,9, 10,10,10))
E(g4)$weight <- seq_len(ecount(g4))
ug4 <- as.undirected(g4, mode="mutual",
                     edge.attr.comb=list(weight=length))
print(ug4, e=TRUE)
```

### as.igraph

#### Conversion to igraph

**Description**

These functions convert various objects to igraph graphs.

**Usage**

```r
as.igraph(x, ...)
```

**Arguments**

- `x`  The object to convert.
- `...` Additional arguments. None currently.

**Details**

You can use `as.igraph` to convert various objects to igraph graphs. Right now the following objects are supported:

- codeigraphHRG These objects are created by the `fit_hrg` and `consensus_tree` functions.

**Value**

All these functions return an igraph graph.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>.

**Examples**

```r
g <- make_full_graph(5) + make_full_graph(5)
hrg <- fit_hrg(g)
as.igraph(hrg)
```
as.matrix.igraph | Convert igraph objects to adjacency or edge list matrices

Description

Get adjacency or edgelist representation of the network stored as an igraph object.

Usage

```r
## S3 method for class 'igraph'
as.matrix(x, matrix.type = c("adjacency", "edgelist"), ...)  
```

Arguments

- `x`: object of class igraph, the network
- `matrix.type`: character, type of matrix to return, currently "adjacency" or "edgelist" are supported
- `...`: other arguments to/from other methods

Details

If `matrix.type` is "edgelist", then a two-column numeric edge list matrix is returned. The value of `attrname` is ignored.

If `matrix.type` is "adjacency", then a square adjacency matrix is returned. For adjacency matrices, you can use the `attr` keyword argument to use the values of an edge attribute in the matrix cells. See the documentation of `as_adjacency_matrix` for more details.

Other arguments passed through `...` are passed to either `as_adjacency_matrix` or `as_edgelist` depending on the value of `matrix.type`.

Value

Depending on the value of `matrix.type` type either a square adjacency matrix or a two-column numeric matrix representing the edgelist.

Author(s)

Michal Bojanowski, originally from the intergraph package

See Also

`as_adjacency_matrix`, `as_edgelist`

Examples

```r
g <- make_graph("zachary")
as.matrix(g, "adjacency")
as.matrix(g, "edgelist")
# use edge attribute "weight"
E(g)$weight <- rep(1:10, each=ecount(g))
as.matrix(g, "adjacency", sparse=FALSE, attr="weight")
```
as_adj_list

Adjacency lists

Description
Create adjacency lists from a graph, either for adjacent edges or for neighboring vertices

Usage
as_adj_list(graph, mode = c("all", "out", "in", "total"))
as_adj_edge_list(graph, mode = c("all", "out", "in", "total"))

Arguments
- graph: The input graph.
- mode: Character scalar, it gives what kind of adjacent edges/vertices to include in the lists. 'out' is for outgoing edges/vertices, 'in' is for incoming edges/vertices, 'all' is for both. This argument is ignored for undirected graphs.

Details
as_adj_list returns a list of numeric vectors, which include the ids of neighbor vertices (according to the mode argument) of all vertices.
as_adj_edge_list returns a list of numeric vectors, which include the ids of adjacent edges (according to the mode argument) of all vertices.
If igraph_opt("return.vs.es") is true (default), the numeric vectors of the adjacency lists are coerced to igraph.vs, this can be a very expensive operation on large graphs.

Value
A list of igraph.vs or a list of numeric vectors depending on the value of igraph_opt("return.vs.es"), see details for performance characteristics.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

See Also
as_edgelist, as_adj

Examples
```r
  g <- make_ring(10)
as_adj_list(g)
as_adj_edge_list(g)
```
as_adjacency_matrix

Convert a graph to an adjacency matrix

Description

Sometimes it is useful to work with a standard representation of a graph, like an adjacency matrix.

Usage

as_adjacency_matrix(
  graph,
  type = c("both", "upper", "lower"),
  attr = NULL,
  edges = FALSE,
  names = TRUE,
  sparse = igraph_opt("sparsematrices")
)

as_adj(
  graph,
  type = c("both", "upper", "lower"),
  attr = NULL,
  edges = FALSE,
  names = TRUE,
  sparse = igraph_opt("sparsematrices")
)

Arguments

graph The graph to convert.

type Gives how to create the adjacency matrix for undirected graphs. It is ignored for directed graphs. Possible values: upper: the upper right triangle of the matrix is used, lower: the lower left triangle of the matrix is used, both: the whole matrix is used, a symmetric matrix is returned.

attr Either NULL or a character string giving an edge attribute name. If NULL a traditional adjacency matrix is returned. If not NULL then the values of the given edge attribute are included in the adjacency matrix. If the graph has multiple edges, the edge attribute of an arbitrarily chosen edge (for the multiple edges) is included. This argument is ignored if edges is TRUE.

Note that this works only for certain attribute types. If the sparse argument is TRUE, then the attribute must be either logical or numeric. If the sparse argument is FALSE, then character is also allowed. The reason for the difference is that the Matrix package does not support character sparse matrices yet.

edges Logical scalar, whether to return the edge ids in the matrix. For non-existant edges zero is returned.

names Logical constant, whether to assign row and column names to the matrix. These are only assigned if the name vertex attribute is present in the graph.

sparse Logical scalar, whether to create a sparse matrix. The 'Matrix' package must be installed for creating sparse matrices.
Details

as_adjacency_matrix returns the adjacency matrix of a graph, a regular matrix if sparse is FALSE, or a sparse matrix, as defined in the `Matrix` package, if sparse if TRUE.

Value

A vcount(graph) by vcount(graph) (usually) numeric matrix.

See Also

`graph_from_adjacency_matrix`, `read_graph`

Examples

```r
g <- sample_gnp(10, 2/10)
as_adjacency_matrix(g)
V(g)$name <- letters[1:vcount(g)]
as_adjacency_matrix(g)
E(g)$weight <- runif(ecount(g))
as_adjacency_matrix(g, attr="weight")
```

---

**as_data_frame**

Creating igraph graphs from data frames or vice-versa

Description

This function creates an igraph graph from one or two data frames containing the (symbolic) edge list and edge/vertex attributes.

Usage

```r
as_data_frame(x, what = c("edges", "vertices", "both"))
graph_from_data_frame(d, directed = TRUE, vertices = NULL)
from_data_frame(...)```

Arguments

- `x` An igraph object.
- `what` Character constant, whether to return info about vertices, edges, or both. The default is ‘edges’.
- `d` A data frame containing a symbolic edge list in the first two columns. Additional columns are considered as edge attributes. Since version 0.7 this argument is coerced to a data frame with `as.data.frame`.
- `directed` Logical scalar, whether or not to create a directed graph.
- `vertices` A data frame with vertex metadata, or NULL. See details below. Since version 0.7 this argument is coerced to a data frame with `as.data.frame`, if not NULL.
- `...` Passed to `graph_from_data_frame`.
**Details**

`graph_from_data_frame` creates igraph graphs from one or two data frames. It has two modes of operation, depending whether the `vertices` argument is `NULL` or not.

If `vertices` is `NULL`, then the first two columns of `d` are used as a symbolic edge list and additional columns as edge attributes. The names of the attributes are taken from the names of the columns.

If `vertices` is not `NULL`, then it must be a data frame giving vertex metadata. The first column of `vertices` is assumed to contain symbolic vertex names, this will be added to the graphs as the ‘name’ vertex attribute. Other columns will be added as additional vertex attributes. If `vertices` is not `NULL` then the symbolic edge list given in `d` is checked to contain only vertex names listed in `vertices`.

Typically, the data frames are exported from some spreadsheet software like Excel and are imported into R via `read.table`, `read.delim` or `read.csv`.

All edges in the data frame are included in the graph, which may include multiple parallel edges and loops.

`as_data_frame` converts the igraph graph into one or more data frames, depending on the `what` argument.

If the `what` argument is `edges` (the default), then the edges of the graph and also the edge attributes are returned. The edges will be in the first two columns, named `from` and `to`. (This also denotes edge direction for directed graphs.) For named graphs, the vertex names will be included in these columns, for other graphs, the numeric vertex ids. The edge attributes will be in the other columns. It is not a good idea to have an edge attribute named `from` or `to`, because then the column named in the data frame will not be unique. The edges are listed in the order of their numeric ids.

If the `what` argument is `vertices`, then vertex attributes are returned. Vertices are listed in the order of their numeric vertex ids.

If the `what` argument is `both`, then both vertex and edge data is returned, in a list with named entries `vertices` and `edges`.

**Value**

An igraph graph object for `graph_from_data_frame`, and either a data frame or a list of two data frames named `edges` and `vertices` for `as.data.frame`.

**Note**

For `graph_from_data_frame` `NA` elements in the first two columns ‘d’ are replaced by the string “NA” before creating the graph. This means that all NAs will correspond to a single vertex.

`NA` elements in the first column of ‘vertices’ are also replaced by the string “NA”, but the rest of ‘vertices’ is not touched. In other words, vertex names (=the first column) cannot be `NA`, but other vertex attributes can.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**See Also**

`graph_from_literal` for another way to create graphs, `read.table` to read in tables from files.
Examples

```r
## A simple example with a couple of actors
## The typical case is that these tables are read in from files....
actors <- data.frame(name=c("Alice", "Bob", "Cecil", "David", "Esmeralda"),
                     age=c(48,33,45,34,21),
                     gender=c("F","M","F","M","F"))
relations <- data.frame(from=c("Bob", "Cecil", "Cecil", "David", "David", "Esmeralda"),
                        to=c("Alice", "Bob", "Alice", "Alice", "Bob", "Alice"),
                        same.dept=c(FALSE,FALSE,TRUE,FALSE,FALSE,TRUE),
                        friendship=c(4,5,5,2,1,1), advice=c(4,5,5,4,2,3))
g <- graph_from_data_frame(relations, directed=TRUE, vertices=actors)
print(g, e=TRUE, v=TRUE)

## The opposite operation
as_data_frame(g, what="vertices")
as_data_frame(g, what="edges")
```

---

**as_edgelist**

*Convert a graph to an edge list*

**Description**

Sometimes it is useful to work with a standard representation of a graph, like an edge list.

**Usage**

```r
as_edgelist(graph, names = TRUE)
```

**Arguments**

- `graph` The graph to convert.
- `names` Whether to return a character matrix containing vertex names (ie. the name vertex attribute) if they exist or numeric vertex ids.

**Details**

`as_edgelist` returns the list of edges in a graph.

**Value**

A `gsize(graph)` by 2 numeric matrix.

**See Also**

- `graph_from_adjacency_matrix`, `read_graph`
as_graphnel

Examples

```r
# Undirected
# Not run:
g <- make_ring(10)
V(g)$name <- letters[1:10]
GNEL <- as_graphnel(g)
g2 <- graph_from_graphnel(GNEL)
g2

# Directed
g3 <- make_star(10, mode="in")
V(g3)$name <- letters[1:10]
GNEL2 <- as_graphnel(g3)
```

as_graphnel

Convert igraph graphs to graphNEL objects from the graph package

Description

The graphNEL class is defined in the `graph` package, it is another way to represent graphs. These functions are provided to convert between the igraph and the graphNEL objects.

Usage

```r
as_graphnel(graph)
```

Arguments

graph

An igraph graph object.

Details

as_graphnel converts an igraph graph to a graphNEL graph. It converts all graph/vertex/edge attributes. If the igraph graph has a vertex attribute ‘name’, then it will be used to assign vertex names in the graphNEL graph. Otherwise numeric igraph vertex ids will be used for this purpose.

Value

as_graphnel returns a graphNEL graph object.

See Also

`graph_from_graphnel` for the other direction, `as_adj`, `graph_from_adjacency_matrix`, `as_adj_list` and `graph.adjlist` for other graph representations.

Examples

```r
## Undirected
## Not run:
g <- sample_gnp(10, 2/10)
as_edgelist(g)
V(g)$name <- LETTERS[seq_len(gorder(g))]
as_edgelist(g)
```
as_ids

Convert a vertex or edge sequence to an ordinary vector

Description

Convert a vertex or edge sequence to an ordinary vector

Usage

as_ids(seq)

## S3 method for class 'igraph.vs'
as_ids(seq)

## S3 method for class 'igraph.es'
as_ids(seq)

Arguments

seq The vertex or edge sequence.

Details

For graphs without names, a numeric vector is returned, containing the internal numeric vertex or edge ids.

For graphs with names, and vertex sequences, the vertex names are returned in a character vector.

For graphs with names and edge sequences, a character vector is returned, with the ‘bar’ notation: a|b means an edge from vertex a to vertex b.

Value

A character or numeric vector, see details below.

Examples

g <- make_ring(10)
as_ids(V(g))
as_ids(E(g))

V(g)$name <- letters[1:10]
as_ids(V(g))
as_ids(E(g))
as_incidence_matrix

Incidence matrix of a bipartite graph

Description

This function can return a sparse or dense incidence matrix of a bipartite network. The incidence matrix is an $n \times m$ matrix, $n$ and $m$ are the number of vertices of the two kinds.

Usage

as_incidence_matrix(
  graph,
  types = NULL,
  attr = NULL,
  names = TRUE,
  sparse = FALSE
)

Arguments

graph The input graph. The direction of the edges is ignored in directed graphs.
types An optional vertex type vector to use instead of the type vertex attribute. You must supply this argument if the graph has no type vertex attribute.
attr Either NULL or a character string giving an edge attribute name. If NULL, then a traditional incidence matrix is returned. If not NULL then the values of the given edge attribute are included in the incidence matrix. If the graph has multiple edges, the edge attribute of an arbitrarily chosen edge (for the multiple edges) is included.
names Logical scalar, if TRUE and the vertices in the graph are named (i.e. the graph has a vertex attribute called name), then vertex names will be added to the result as row and column names. Otherwise the ids of the vertices are used as row and column names.
sparse Logical scalar, if it is TRUE then a sparse matrix is created, you will need the Matrix package for this.

Details

Bipartite graphs have a type vertex attribute in igraph, this is boolean and FALSE for the vertices of the first kind and TRUE for vertices of the second kind.

Value

A sparse or dense matrix.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

graph_from_incidence_matrix for the opposite operation.
as_membership

Examples

```r
# Example 1
g <- make_bipartite_graph( c(0,1,0,1,0,0), c(1,2,2,3,3,4) )
as_membership(g)
```

```r
# Example 2
as_long_data_frame(g)
```

as_membership

Declare a numeric vector as a membership vector

Description

This is useful if you want to use functions defined on membership vectors, but your membership vector does not come from an igraph clustering method.

Usage

```r
as_membership(x)
```

Arguments

- `x`: The input vector.
assortativity

Value

The input vector, with the membership class added.

Examples

```r
## Compare to the correct clustering

g <- (make_full_graph(10) + make_full_graph(10)) %>%
  rewire(each_edge(p = 0.2))
correct <- rep(1:2, each = 10) %>% as_membership
fc <- cluster_fast_greedy(g)
compare(correct, fc)
compare(correct, membership(fc))
```

assortativity  Assortativity coefficient

Description

The assortativity coefficient is positive is similar vertices (based on some external property) tend to connect to each, and negative otherwise.

Usage

```r
assortativity(graph, types1, types2 = NULL, directed = TRUE)
assortativity_nominal(graph, types, directed = TRUE)
assortativity_degree(graph, directed = TRUE)
```

Arguments

- `graph`: The input graph, it can be directed or undirected.
- `types1`: The vertex values, these can be arbitrary numeric values.
- `types2`: A second value vector to be using for the incoming edges when calculating assortativity for a directed graph. Supply NULL here if you want to use the same values for outgoing and incoming edges. This argument is ignored (with a warning) if it is not NULL and undirected assortativity coefficient is being calculated.
- `directed`: Logical scalar, whether to consider edge directions for directed graphs. This argument is ignored for undirected graphs. Supply TRUE here to do the natural thing, i.e. use directed version of the measure for directed graphs and the undirected version for undirected graphs.
- `types`: Vector giving the vertex types. They as assumed to be integer numbers, starting with one. Non-integer values are converted to integers with `as.integer`.

Details

The assortativity coefficient measures the level of homophily of the graph, based on some vertex labeling or values assigned to vertices. If the coefficient is high, that means that connected vertices tend to have the same labels or similar assigned values.
M.E.J. Newman defined two kinds of assortativity coefficients, the first one is for categorical labels of vertices. `assortativity_nominal` calculates this measure. It is defined as

$$ r = \frac{\sum_i e_{ii} - \sum_i a_i b_i}{1 - \sum_i a_i b_i} $$

where $e_{ij}$ is the fraction of edges connecting vertices of type $i$ and $j$, $a_i = \sum_j e_{ij}$ and $b_j = \sum_i e_{ij}$.

The second assortativity variant is based on values assigned to the vertices. `assortativity` calculates this measure. It is defined as

$$ r = \frac{1}{\sigma_q^2} \sum_{jk} jk(e_{jk} - q_j q_k) $$

for undirected graphs ($q_i = \sum_j e_{ij}$) and as

$$ r = \frac{1}{\sigma_o \sigma_i} \sum_{jk} jk(e_{jk} - q_o^i q_k^i) $$

for directed ones. Here $\sigma_q^2 = \sum_j e_{ij}$, $q_i^i = \sum_j e_{ji}$, moreover, $\sigma_q$, $\sigma_o$, and $\sigma_i$ are the standard deviations of $q$, $q_o$ and $q_i$, respectively.

The reason of the difference is that in directed networks the relationship is not symmetric, so it is possible to assign different values to the outgoing and the incoming end of the edges.

`assortativity_degree` uses vertex degree (minus one) as vertex values and calls `assortativity`.

**Value**

A single real number.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**References**


**Examples**

```python
# random network, close to zero
assortativity_degree(sample_gnp(10000, 3/10000))

# BA model, tends to be dissortative
assortativity_degree(sample_pa(10000, m=4))
```
authority_score  

Kleinberg’s authority centrality scores.

Description

The authority scores of the vertices are defined as the principal eigenvector of $A^T A$, where $A$ is the adjacency matrix of the graph.

Usage

authority_score(graph, scale = TRUE, weights = NULL, options = arpack_defaults)

Arguments

- **graph**: The input graph.
- **scale**: Logical scalar, whether to scale the result to have a maximum score of one. If no scaling is used then the result vector has unit length in the Euclidean norm.
- **weights**: Optional positive weight vector for calculating weighted scores. If the graph has a weight edge attribute, then this is used by default. This function interprets edge weights as connection strengths. In the random surfer model, an edge with a larger weight is more likely to be selected by the surfer.
- **options**: A named list, to override some ARPACK options. See arpack for details.

Details

For undirected matrices the adjacency matrix is symmetric and the authority scores are the same as hub scores, see hub_score.

Value

A named list with members:

- **vector**: The authority/hub scores of the vertices.
- **value**: The corresponding eigenvalue of the calculated principal eigenvector.
- **options**: Some information about the ARPACK computation, it has the same members as the options member returned by arpack, see that for documentation.

References


See Also

hub_score, eigen_centrality for eigenvector centrality, page_rank for the Page Rank scores. arpack for the underlining machinery of the computation.
Examples

```r
## An in-star
g <- make_star(10)
hub_score(g)$vector
authority_score(g)$vector

## A ring
g2 <- make_ring(10)
hub_score(g2)$vector
authority_score(g2)$vector
```

### automorphism_group

Generating set of the automorphism group of a graph

Description

Compute the generating set of the automorphism group of a graph.

Usage

```r
automorphism_group(
  graph,
  colors,
  sh = c("fm", "f", "fs", "fl", "flm", "fsm"),
  details = FALSE
)
```

Arguments

- **graph**: The input graph, it is treated as undirected.
- **colors**: The colors of the individual vertices of the graph; only vertices having the same color are allowed to match each other in an automorphism. When omitted, igraph uses the color attribute of the vertices, or, if there is no such vertex attribute, it simply assumes that all vertices have the same color. Pass NULL explicitly if the graph has a color vertex attribute but you do not want to use it.
- **sh**: The splitting heuristics for the BLISS algorithm. Possible values are: 'f': first non-singleton cell, 'fl': first largest non-singleton cell, 'fs': first smallest non-singleton cell, 'fm': first maximally non-trivially connected non-singleton cell, 'flm': first largest maximally non-trivially connected non-singleton cell, 'fsm': first smallest maximally non-trivially connected non-singleton cell.
- **details**: Specifies whether to provide additional details about the BLISS internals in the result.

Details

An automorphism of a graph is a permutation of its vertices which brings the graph into itself. The automorphisms of a graph form a group and there exists a subset of this group (i.e. a set of permutations) such that every other permutation can be expressed as a combination of these permutations. These permutations are called the generating set of the automorphism group.

This function calculates a possible generating set of the automorphism of a graph using the BLISS algorithm. See also the BLISS homepage at [http://www.tcs.hut.fi/Software/bliss/index](http://www.tcs.hut.fi/Software/bliss/index).
Calculate the number of automorphisms of a graph, i.e. the number of isomorphisms to itself.

Usage

```r
automorphisms(graph, colors, sh = c("fm", "f", "fs", "f1", "f1m", "fsm"))
```
Arguments

- **graph**: The input graph, it is treated as undirected.
- **colors**: The colors of the individual vertices of the graph; only vertices having the same color are allowed to match each other in an automorphism. When omitted, igraph uses the color attribute of the vertices, or, if there is no such vertex attribute, it simply assumes that all vertices have the same color. Pass NULL explicitly if the graph has a color vertex attribute but you do not want to use it.

Details

An automorphism of a graph is a permutation of its vertices which brings the graph into itself.

This function calculates the number of automorphism of a graph using the BLISS algorithm. See also the BLISS homepage at [http://www.tcs.hut.fi/Software/bliss/index.html](http://www.tcs.hut.fi/Software/bliss/index.html). If you need the automorphisms themselves, use `automorphism_group` to obtain a compact representation of the automorphism group.

Value

A named list with the following members:

- **group.size**: The size of the automorphism group of the input graph, as a string. This number is exact if igraph was compiled with the GMP library, and approximate otherwise.
- **nof.nodes**: The number of nodes in the search tree.
- **nof.leaf.nodes**: The number of leaf nodes in the search tree.
- **nof.bad.nodes**: Number of bad nodes.
- **nof.canupdates**: Number of canrep updates.
- **max.level**: Maximum level.

Author(s)

Tommi Junttila ([http://users.ics.aalto.fi/tjunttil/](http://users.ics.aalto.fi/tjunttil/)) for BLISS and Gabor Csardi <csardi.gabor@gmail.com> for the igraph glue code and this manual page.

References


See Also

`canonical_permutation`, `permute`, and `automorphism_group` for a compact representation of all automorphisms
Examples

```r
## A ring has n*2 automorphisms, you can "turn" it by 0-9 vertices
## and each of these graphs can be "flipped"
g <- make_ring(10)
automorphisms(g)

## A full graph has n! automorphisms; however, we restrict the vertex
## matching by colors, leading to only 4 automorphisms
h <- make_full_graph(4)
automorphisms(h, colors=c(1,2,1,2))
```

---

**bfs**

*Breadth-first search*

Description

Breadth-first search is an algorithm to traverse a graph. We start from a root vertex and spread along every edge “simultaneously”.

Usage

```r
bfs(
  graph,
  root,
  mode = c("out", "in", "all", "total"),
  unreachable = TRUE,
  restricted = NULL,
  order = TRUE,
  rank = FALSE,
  father = FALSE,
  pred = FALSE,
  succ = FALSE,
  dist = FALSE,
  callback = NULL,
  extra = NULL,
  rho = parent.frame(),
  neimode
)
```

Arguments

- **graph**: The input graph.
- **root**: Numeric vector, usually of length one. The root vertex, or root vertices to start the search from.
- **mode**: For directed graphs specifies the type of edges to follow. ‘out’ follows outgoing, ‘in’ incoming edges. ‘all’ ignores edge directions completely. ‘total’ is a synonym for ‘all’. This argument is ignored for undirected graphs.
- **unreachable**: Logical scalar, whether the search should visit the vertices that are unreachable from the given root vertex (or vertices). If TRUE, then additional searches are performed until all vertices are visited.
bfs

restricted NULL (=no restriction), or a vector of vertices (ids or symbolic names). In the latter case, the search is restricted to the given vertices.

order Logical scalar, whether to return the ordering of the vertices.

rank Logical scalar, whether to return the rank of the vertices.

father Logical scalar, whether to return the father of the vertices.

pred Logical scalar, whether to return the predecessors of the vertices.

succ Logical scalar, whether to return the successors of the vertices.

dist Logical scalar, whether to return the distance from the root of the search tree.

callback If not NULL, then it must be callback function. This is called whenever a vertex is visited. See details below.

extra Additional argument to supply to the callback function.

rho The environment in which the callback function is evaluated.

neimode This argument is deprecated from igraph 1.3.0; use mode instead.

Details

The callback function must have the following arguments:

**graph** The input graph is passed to the callback function here.

**data** A named numeric vector, with the following entries: ‘vid’, the vertex that was just visited, ‘pred’, its predecessor (zero if this is the first vertex), ‘succ’, its successor (zero if this is the last vertex), ‘rank’, the rank of the current vertex, ‘dist’, its distance from the root of the search tree.

**extra** The extra argument.

The callback must return FALSE to continue the search or TRUE to terminate it. See examples below on how to use the callback function.

Value

A named list with the following entries:

**root** Numeric scalar. The root vertex that was used as the starting point of the search.

**neimode** Character scalar. The mode argument of the function call. Note that for undirected graphs this is always ‘all’, irrespectively of the supplied value.

**order** Numeric vector. The vertex ids, in the order in which they were visited by the search.

**rank** Numeric vector. The rank for each vertex.

**father** Numeric vector. The father of each vertex, i.e. the vertex it was discovered from.

**pred** Numeric vector. The previously visited vertex for each vertex, or 0 if there was no such vertex.

**succ** Numeric vector. The next vertex that was visited after the current one, or 0 if there was no such vertex.

**dist** Numeric vector, for each vertex its distance from the root of the search tree.

Note that **order**, **rank**, **father**, **pred**, **succ** and **dist** might be NULL if their corresponding argument is FALSE, i.e. if their calculation is not requested.
**biconnected_components**

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**See Also**

dfs for depth-first search.

**Examples**

```r
## Two rings
bfs(make_ring(10) %du% make_ring(10), root=1, "out",
    order=TRUE, rank=TRUE, father=TRUE, pred=TRUE,
    succ=TRUE, dist=TRUE)
```

```c
## How to use a callback
f <- function(graph, data, extra) {
  print(data)
  FALSE
}

## How to use a callback to stop the search
## We stop after visiting all vertices in the initial component
f <- function(graph, data, extra) {
  data['succ'] == -1
}

bfs(make_ring(10) %du% make_ring(10), root=1, "out",
    callback=f)
```

**Description**

Finding the biconnected components of a graph

**Usage**

`biconnected_components(graph)`

**Arguments**

- `graph` The input graph. It is treated as an undirected graph, even if it is directed.
A graph is biconnected if the removal of any single vertex (and its adjacent edges) does not disconnect it.

A biconnected component of a graph is a maximal biconnected subgraph of it. The biconnected components of a graph can be given by the partition of its edges: every edge is a member of exactly one biconnected component. Note that this is not true for vertices: the same vertex can be part of many biconnected components.

Value

A named list with three components:

- **no**
  - Numeric scalar, an integer giving the number of biconnected components in the graph.

- **tree_edges**
  - The components themselves, a list of numeric vectors. Each vector is a set of edge ids giving the edges in a biconnected component. These edges define a spanning tree of the component.

- **component_edges**
  - A list of numeric vectors. It gives all edges in the components.

- **components**
  - A list of numeric vectors, the vertices of the components.

- **articulation_points**
  - The articulation points of the graph. See `articulation_points`.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

- `articulation_points`, `components`, `is_connected`, `vertex_connectivity`

Examples

```r
  g <- disjoint_union( make_full_graph(5), make_full_graph(5) )
  clu <- components(g)$membership
  g <- add_edges(g, c(which(clu==1), which(clu==2)))
  bc <- biconnected_components(g)
```

---

**bipartite_mapping**

*Decide whether a graph is bipartite*

This function decides whether the vertices of a network can be mapped to two vertex types in a way that no vertices of the same type are connected.

**Usage**

*bipartite_mapping(graph)*
bipartite_projection

Arguments

graph The input graph.

Details

A bipartite graph in igraph has a ‘type’ vertex attribute giving the two vertex types. This function simply checks whether a graph could be bipartite. It tries to find a mapping that gives a possible division of the vertices into two classes, such that no two vertices of the same class are connected by an edge.

The existence of such a mapping is equivalent of having no circuits of odd length in the graph. A graph with loop edges cannot bipartite.

Note that the mapping is not necessarily unique, e.g. if the graph has at least two components, then the vertices in the separate components can be mapped independently.

Value

A named list with two elements:

res A logical scalar, TRUE if the can be bipartite, FALSE otherwise.

type A possible vertex type mapping, a logical vector. If no such mapping exists, then an empty vector.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

## Rings with an even number of vertices are bipartite

g <- make_ring(10)
bipartite_mapping(g)

## All star graphs are bipartite

g2 <- make_star(10)
bipartite_mapping(g2)

## A graph containing a triangle is not bipartite

g3 <- make_ring(10)
g3 <- add_edges(g3, c(1,3))
bipartite_mapping(g3)

bipartite_projection Project a bipartite graph

Description

A bipartite graph is projected into two one-mode networks
Usage

```r
bipartite_projection(
  graph,
  types = NULL,
  multiplicity = TRUE,
  probe1 = NULL,
  which = c("both", "true", "false"),
  remove.type = TRUE
)
```

Arguments

- `graph`: The input graph. It can be directed, but edge directions are ignored during the computation.
- `types`: An optional vertex type vector to use instead of the 'type' vertex attribute. You must supply this argument if the graph has no 'type' vertex attribute.
- `multiplicity`: If `TRUE`, then igraph keeps the multiplicity of the edges as an edge attribute called 'weight'. E.g. if there is an A-C-B and also an A-D-B triple in the bipartite graph (but no more X, such that A-X-B is also in the graph), then the multiplicity of the A-B edge in the projection will be 2.
- `probe1`: This argument can be used to specify the order of the projections in the resulting list. If given, then it is considered as a vertex id (or a symbolic vertex name); the projection containing this vertex will be the first one in the result list. This argument is ignored if only one projection is requested in argument `which`.
- `which`: A character scalar to specify which projection(s) to calculate. The default is to calculate both.
- `remove.type`: Logical scalar, whether to remove the type vertex attribute from the projections. This makes sense because these graphs are not bipartite any more. However if you want to combine them with each other (or other bipartite graphs), then it is worth keeping this attribute. By default it will be removed.

Details

Bipartite graphs have a type vertex attribute in igraph, this is boolean and `FALSE` for the vertices of the first kind and `TRUE` for vertices of the second kind.

`bipartite_projection_size` calculates the number of vertices and edges in the two projections of the bipartite graphs, without calculating the projections themselves. This is useful to check how much memory the projections would need if you have a large bipartite graph.

`bipartite_projection` calculates the actual projections. You can use the `probe1` argument to specify the order of the projections in the result. By default vertex type `FALSE` is the first and `TRUE` is the second.

`bipartite_projection` keeps vertex attributes.

Value

A list of two undirected graphs. See details above.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>
Examples

```r
## Projection of a full bipartite graph is a full graph
G <- make_full_bipartite_graph(10, 5)
proj <- bipartite_projection(G)
graph.isomorphic(proj[[1]], make_full_graph(10))
graph.isomorphic(proj[[2]], make_full_graph(5))

## The projection keeps the vertex attributes
M <- matrix(0, nrow=5, ncol=3)ownames(M) <- c("Alice", "Bob", "Cecil", "Dan", "Ethel")
colnames(M) <- c("Party", "Skiing", "Badminton")
M[] <- sample(0:1, length(M), replace=TRUE)
G2 <- graph_from_incidence_matrix(M)
G2$name <- "Event network"
proj2 <- bipartite_projection(G2)
print(proj2[[1]], g=TRUE, e=TRUE)
print(proj2[[2]], g=TRUE, e=TRUE)
```

---

### c.igraph.es

**Concatenate edge sequences**

**Description**

Concatenate edge sequences

**Usage**

```r
## S3 method for class 'igraph.es'
c(..., recursive = FALSE)
```

**Arguments**

- `...` The edge sequences to concatenate. They must all refer to the same graph.
- `recursive` Ignored, included for S3 compatibility with the base `c` function.

**Value**

An edge sequence, the input sequences concatenated.

**See Also**


**Examples**

```r
g <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
c(E(g)[[1]], E(g)[['A'|'B']], E(g)[1:4])
```
c.igraph.vs  Concatenate vertex sequences

Description

Concatenate vertex sequences

Usage

```r
## S3 method for class 'igraph.vs'
c(..., recursive = FALSE)
```

Arguments

- `...` - The vertex sequences to concatenate. They must refer to the same graph.
- `recursive` - Ignored, included for S3 compatibility with the base `c` function.

Value

A vertex sequence, the input sequences concatenated.

See Also


Examples

```r
g <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
c(V(g)[1], V(g)['A'], V(g)[1:4])
```

canonical_permutation  Canonical permutation of a graph

Description

The canonical permutation brings every isomorphic graphs into the same (labeled) graph.

Usage

```r
canonical_permutation(
  graph,
  colors,
  sh = c("fm", "f", "fs", "fl", "flm", "fsm")
)
```
canonical_permutation

Arguments

graph
The input graph, treated as undirected.

colors
The colors of the individual vertices of the graph; only vertices having the same color are allowed to match each other in an automorphism. When omitted, igraph uses the color attribute of the vertices, or, if there is no such vertex attribute, it simply assumes that all vertices have the same color. Pass NULL explicitly if the graph has a color vertex attribute but you do not want to use it.

sh
Type of the heuristics to use for the BLISS algorithm. See details for possible values.

Details

canonical_permutation computes a permutation which brings the graph into canonical form, as defined by the BLISS algorithm. All isomorphic graphs have the same canonical form.

See the paper below for the details about BLISS. This and more information is available at http://www.tcs.hut.fi/Software/bliss/index.html.

The possible values for the sh argument are:

"f" First non-singleton cell.
"fl" First largest non-singleton cell.
"fs" First smallest non-singleton cell.
"fm" First maximally non-trivially connectec non-singleton cell.
"flm" Largest maximally non-trivially connected non-singleton cell.
"fsm" Smallest maximally non-trivially connected non-singleton cell.

See the paper in references for details about these.

Value

A list with the following members:

labeling
The canonical permutation which takes the input graph into canonical form. A numeric vector, the first element is the new label of vertex 0, the second element for vertex 1, etc.

info
Some information about the BLISS computation. A named list with the following members:

"nof_nodes" The number of nodes in the search tree.
"nof_leaf_nodes" The number of leaf nodes in the search tree.
"nof_bad_nodes" Number of bad nodes.
"nof_canupdates" Number of canrep updates.
"max_level" Maximum level.
"group_size" The size of the automorphism group of the input graph, as a string. The string representation is necessary because the group size can easily exceed values that are exactly representable in floating point.

Author(s)

Tommi Junttila for BLISS, Gabor Csardi <csardi.gabor@gmail.com> for the igraph and R interfaces.
References


See Also

`permute` to apply a permutation to a graph, `graph.isomorphic` for deciding graph isomorphism, possibly based on canonical labels.

Examples

```r
## Calculate the canonical form of a random graph
g1 <- sample_gnm(10, 20)
cp1 <- canonical_permutation(g1)
cf1 <- permute(g1, cp1$labeling)

## Do the same with a random permutation of it
g2 <- permute(g1, sample(vcount(g1)))
cp2 <- canonical_permutation(g2)
cf2 <- permute(g2, cp2$labeling)

## Check that they are the same
el1 <- as_edgelist(cf1)
el2 <- as_edgelist(cf2)
el1 <- el1[ order(el1[,1], el1[,2]), ]
el2 <- el2[ order(el2[,1], el2[,2]), ]
all(el1 == el2)
```

categorical_pal

Palette for categories

Description

This is a color blind friendly palette from [https://jfly.uni-koeln.de/color/](https://jfly.uni-koeln.de/color/). It has 8 colors.

Usage

`categorical_pal(n)`

Arguments

- `n` The number of colors in the palette. We simply take the first `n` colors from the total 8.

Details

This is the suggested palette for visualizations where vertex colors mark categories, e.g. community membership.

Value

A character vector of RGB color codes.
Examples

```r
library(igraphdata)
data(karate)
karate <- karate
  add_layout_(with_fr())
  set_vertex_attr("size", value = 10)

cl_k <- cluster_optimal(karate)
V(karate)$color <- membership(cl_k)
karate$palette <- categorical_pal(length(cl_k))
plot(karate)
```

See Also

Other palettes: `diverging_pal()`, `r_pal()`, `sequential_pal()`

---

### centr_betw

Centralize a graph according to the betweenness of vertices

#### Description

See `centralize` for a summary of graph centralization.

#### Usage

```r
centr_betw(graph, directed = TRUE, nobigint = TRUE, normalized = TRUE)
```

#### Arguments

- `graph` The input graph.
- `directed` Logical scalar, whether to use directed shortest paths for calculating betweenness.
- `nobigint` Logical scalar, whether to use big integers for the betweenness calculation. This argument is deprecated in igraph 1.3 and will be removed in igraph 1.4.
- `normalized` Logical scalar. Whether to normalize the graph level centrality score by dividing by the theoretical maximum.

#### Value

A named list with the following components:

- `res` The node-level centrality scores.
- `centralization` The graph level centrality index.
- `theoretical_max` The maximum theoretical graph level centralization score for a graph with the given number of vertices, using the same parameters. If the `normalized` argument was TRUE, then the result was divided by this number.
See Also

Other centralization related: `centr_betw_tmax()`, `centr_clo_tmax()`, `centr_clo()`, `centr_degree_tmax()`, `centr_degree()`, `centr_eigen_tmax()`, `centr_eigen()`, `centralize()

Examples

# A BA graph is quite centralized

g <- sample_pa(1000, m = 4)

centr_degree(g)$centralization

centr_clo(g, mode = "all")$centralization

centr_betw(g, directed = FALSE)$centralization

centr_eigen(g, directed = FALSE)$centralization


---

`centr_betw_tmax`  
*Theoretical maximum for betweenness centralization*

Description

See `centralize` for a summary of graph centralization.

Usage

`centr_betw_tmax(graph = NULL, nodes = 0, directed = TRUE)`

Arguments

- `graph`: The input graph. It can also be `NULL`, if `nodes` is given.
- `nodes`: The number of vertices. This is ignored if the graph is given.
- `directed`: logical scalar, whether to use directed shortest paths for calculating betweenness.

Value

Real scalar, the theoretical maximum (unnormalized) graph betweenness centrality score for graphs with given order and other parameters.

See Also

Other centralization related: `centr_betw()`, `centr_clo_tmax()`, `centr_clo()`, `centr_degree_tmax()`, `centr_degree()`, `centr_eigen_tmax()`, `centr_eigen()`, `centralize()`

Examples

# A BA graph is quite centralized

g <- sample_pa(1000, m = 4)

centr_betw(g, normalized = FALSE)$centralization %>%
  `/` (centr_betw_tmax(g))

centr_betw(g, normalized = TRUE)$centralization
Centralize a graph according to the closeness of vertices

Description

See `centralize` for a summary of graph centralization.

Usage

```r
centr_clo(graph, mode = c("out", "in", "all", "total"), normalized = TRUE)
```

Arguments

- `graph`: The input graph.
- `mode`: This is the same as the `mode` argument of `closeness`.
- `normalized`: Logical scalar. Whether to normalize the graph level centrality score by dividing by the theoretical maximum.

Value

A named list with the following components:

- `res`: The node-level centrality scores.
- `centralization`: The graph level centrality index.
- `theoretical_max`: The maximum theoretical graph level centralization score for a graph with the given number of vertices, using the same parameters. If the `normalized` argument was TRUE, then the result was divided by this number.

See Also

Other centralization related: `centr_betw_tmax()`, `centr_betw()`, `centr_clo_tmax()`, `centr_degree_tmax()`, `centr_degree()`, `centr_eigen_tmax()`, `centr_eigen()`, `centralize()`

Examples

```r
# A BA graph is quite centralized
g <- sample_pa(1000, m = 4)
centr_degree(g)$centralization
centr_clo(g, mode = "all")$centralization
centr_betw(g, directed = FALSE)$centralization
centr_eigen(g, directed = FALSE)$centralization
```
centr_clo_tmax  Theoretical maximum for closeness centralization

Description

See centralize for a summary of graph centralization.

Usage

```r
centr_clo_tmax(graph = NULL, nodes = 0, mode = c("out", "in", "all", "total"))
```

Arguments

- **graph**: The input graph. It can also be `NULL`, if `nodes` is given.
- **nodes**: The number of vertices. This is ignored if the graph is given.
- **mode**: This is the same as the `mode` argument of `closeness`.

Value

Real scalar, the theoretical maximum (unnormalized) graph closeness centrality score for graphs with given order and other parameters.

See Also

Other centralization related: `centr_betw_tmax()`, `centr_betw()`, `centr_clo()`, `centr_degree_tmax()`, `centr_degree()`, `centr_eigen_tmax()`, `centr_eigen()`, `centralize()`

Examples

```r
# A BA graph is quite centralized
g <- sample_pa(1000, m = 4)
centr_clo(g, normalized = FALSE)$centralization %>%
  `/`(centr_clo_tmax(g))
centr_clo(g, normalized = TRUE)$centralization
```

centr_degree  Centralize a graph according to the degrees of vertices

Description

See centralize for a summary of graph centralization.

Usage

```r
centr_degree(
  graph,
  mode = c("all", "out", "in", "total"),
  loops = TRUE,
  normalized = TRUE
)
```
Arguments

- **graph**: The input graph.
- **mode**: This is the same as the mode argument of degree.
- **loops**: Logical scalar, whether to consider loops edges when calculating the degree.
- **normalized**: Logical scalar. Whether to normalize the graph level centrality score by dividing by the theoretical maximum.

Value

A named list with the following components:

- **res**: The node-level centrality scores.
- **centralization**: The graph level centrality index.
- **theoretical_max**: The maximum theoretical graph level centralization score for a graph with the given number of vertices, using the same parameters. If the normalized argument was TRUE, then the result was divided by this number.

See Also

Other centralization related: `centr_betw_tmax()`, `centr_betw()`, `centr_clo_tmax()`, `centr_clo()`, `centr_degree_tmax()`, `centr_eigen_tmax()`, `centr_eigen()`, `centralize()`

Examples

```r
# A BA graph is quite centralized
g <- sample_pa(1000, m = 4)
centr_degree(g)$centralization
centr_clo(g, mode = "all")$centralization
centr_betw(g, directed = FALSE)$centralization
centr_eigen(g, directed = FALSE)$centralization
```

centr_degree_tmax

**Theoretical maximum for degree centralization**

Description

See `centralize` for a summary of graph centralization.

Usage

```r
centr_degree_tmax(
  graph = NULL,
  nodes = 0,
  mode = c("all", "out", "in", "total"),
  loops = FALSE
)
```
Arguments

- **graph**: The input graph. It can also be `NULL`, if `nodes`, `mode` and `loops` are all given.
- **nodes**: The number of vertices. This is ignored if the graph is given.
- **mode**: This is the same as the `mode` argument of `degree`.
- **loops**: Logical scalar, whether to consider loops edges when calculating the degree. Currently the default value is `FALSE`, but this argument will be required from igraph 1.4.0.

Value

Real scalar, the theoretical maximum (unnormalized) graph degree centrality score for graphs with given order and other parameters.

See Also

Other centralization related: `centr_betw_tmax()`, `centr_betw()`, `centr_clo_tmax()`, `centr_clo()`, `centr_degree()`, `centr_eigen_tmax()`, `centr_eigen()`, `centralize()`

Examples

```r
# A BA graph is quite centralized
library(igraph)
g <- sample_pa(1000, m = 4)
centr_degree(g, normalized = FALSE)$centralization %>%
  `/`(centr_degree_tmax(g, loops = FALSE))
```

```
centr_degree(g, normalized = TRUE)$centralization
```

### centr_eigen

Centralize a graph according to the eigenvector centrality of vertices

Description

See `centralize` for a summary of graph centralization.

Usage

```r
centr_eigen(  
  graph,  
  directed = FALSE,  
  scale = TRUE,  
  options = arpack_defaults,  
  normalized = TRUE  
)
```

Arguments

- **graph**: The input graph.
- **directed**: logical scalar, whether to use directed shortest paths for calculating eigenvector centrality.
- **scale**: Whether to rescale the eigenvector centrality scores, such that the maximum score is one.
options
This is passed to `eigen_centrality`, the options for the ARPACK eigensolver.

normalized
Logical scalar. Whether to normalize the graph level centrality score by dividing by the theoretical maximum.

Value
A named list with the following components:

- **vector**: The node-level centrality scores.
- **value**: The corresponding eigenvalue.
- **options**: ARPACK options, see the return value of `eigen_centrality` for details.
- **centralization**: The graph level centrality index.
- **theoretical_max**: The same as above, the theoretical maximum centralization score for a graph with the same number of vertices.

See Also
Other centralization related: `centr_betw_tmax()`, `centr_betw()`, `centr_clo_tmax()`, `centr_clo()`, `centr_degree_tmax()`, `centr_degree()`, `centr_eigen_tmax()`, `centralize()`

Examples
```r
# A BA graph is quite centralized
g <- sample_pa(1000, m = 4)
centr_degree(g)$centralization
centr_clo(g, mode = "all")$centralization
centr_betw(g, directed = FALSE)$centralization
centr_eigen(g, directed = FALSE)$centralization

# The most centralized graph according to eigenvector centrality
G0 <- make_graph(c(2,1), n = 10, dir = FALSE)
G1 <- make_star(10, mode = "undirected")
centr_eigen(G0)$centralization
centr_eigen(G1)$centralization
```

---

### Description
See `centralize` for a summary of graph centralization.

#### Usage
```
centr_eigen_tmax(graph = NULL, nodes = 0, directed = FALSE, scale = TRUE)
```
Centralization of a graph

Centralization is a method for creating a graph level centralization measure from the centrality scores of the vertices.

Usage

```r
centralize(scores, theoretical.max = 0, normalized = TRUE)
```

Arguments

- `scores`: The vertex level centrality scores.
- `theoretical.max`: Real scalar. The graph-level centralization measure of the most centralized graph with the same number of vertices as the graph under study. This is only used if the `normalized` argument is set to `TRUE`.
- `normalized`: Logical scalar. Whether to normalize the graph level centrality score by dividing by the supplied theoretical maximum.
Centralization is a general method for calculating a graph-level centrality score based on node-level centrality measure. The formula for this is

\[ C(G) = \sum_v (\max_w c_w - c_v), \]

where \( c_v \) is the centrality of vertex \( v \).

The graph-level centralization measure can be normalized by dividing by the maximum theoretical score for a graph with the same number of vertices, using the same parameters, e.g. directedness, whether we consider loop edges, etc.

For degree, closeness and betweenness the most centralized structure is some version of the star graph, in-star, out-star or undirected star.

For eigenvector centrality the most centralized structure is the graph with a single edge (and potentially many isolates).

centralize implements general centralization formula to calculate a graph-level score from vertex-level scores.

**Value**

A real scalar, the centralization of the graph from which scores were derived.

**References**


**See Also**

Other centralization related: centr_betw_tmax(), centr_betw(), centr_clo_tmax(), centr_clo(), centr_degree_tmax(), centr_degree(), centr_eigen_tmax(), centr_eigen()
cliques

Functions to find cliques, ie. complete subgraphs in a graph

Description

These functions find all, the largest or all the maximal cliques in an undirected graph. The size of the largest clique can also be calculated.

Usage

cliques(graph, min = 0, max = 0)

max_cliques(graph, min = NULL, max = NULL, subset = NULL, file = NULL)

Arguments

graph The input graph, directed graphs will be considered as undirected ones, multiple edges and loops are ignored.

min Numeric constant, lower limit on the size of the cliques to find. NULL means no limit, ie. it is the same as 0.

max Numeric constant, upper limit on the size of the cliques to find. NULL means no limit.

subset If not NULL, then it must be a vector of vertex ids, numeric or symbolic if the graph is named. The algorithm is run from these vertices only, so only a subset of all maximal cliques is returned. See the Eppstein paper for details. This argument makes it possible to easily parallelize the finding of maximal cliques.

file If not NULL, then it must be a file name, i.e. a character scalar. The output of the algorithm is written to this file. (If it exists, then it will be overwritten.) Each clique will be a separate line in the file, given with the numeric ids of its vertices, separated by whitespace.

Details

cliques find all complete subgraphs in the input graph, obeying the size limitations given in the min and max arguments.

largest_cliques finds all largest cliques in the input graph. A clique is largest if there is no other clique including more vertices.

max_cliques finds all maximal cliques in the input graph. A clique is maximal if it cannot be extended to a larger clique. The largest cliques are always maximal, but a maximal clique is not necessarily the largest.

count_max_cliques counts the maximal cliques.

clique_num calculates the size of the largest clique(s).

clique_size_counts returns a numeric vector representing a histogram of clique sizes, between the given minimum and maximum clique size.
closeness

Value

cliques, largest_cliques and clique_num return a list containing numeric vectors of vertex ids. Each list element is a clique, i.e. a vertex sequence of class igraph.vs.

max_cliques returns NULL, invisibly, if its file argument is not NULL. The output is written to the specified file in this case.

clique_num and count_max_cliques return an integer scalar.

clique_size_counts returns a numeric vector with the clique sizes such that the i-th item belongs to cliques of size i. Trailing zeros are currently truncated, but this might change in future versions.

Author(s)

Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

ivs

Examples

# this usually contains cliques of size six
g <- sample_gnp(100, 0.3)
clique_num(g)
cliques(g, min=6)
largest_cliques(g)

# To have a bit less maximal cliques, about 100-200 usually

g <- sample_gnp(100, 0.03)
max_cliques(g)

closeness

Closeness centrality of vertices

Description

Closeness centrality measures how many steps is required to access every other vertex from a given vertex.

Usage

closeness(
  graph,
  vids = V(graph),
  mode = c("out", "in", "all", "total"),
  weights = NULL,
  normalized = FALSE,
  cutoff = -1
)
closeness

Arguments

- **graph**: The graph to analyze.
- **vids**: The vertices for which closeness will be calculated.
- **mode**: Character string, defined the types of the paths used for measuring the distance in directed graphs. “in” measures the paths to a vertex, “out” measures paths from a vertex, *all* uses undirected paths. This argument is ignored for undirected graphs.
- **weights**: Optional positive weight vector for calculating weighted closeness. If the graph has a weight edge attribute, then this is used by default. Weights are used for calculating weighted shortest paths, so they are interpreted as distances.
- **normalized**: Logical scalar, whether to calculate the normalized closeness, i.e. the inverse average distance to all reachable vertices. The non-normalized closeness is the inverse of the sum of distances to all reachable vertices.
- **cutoff**: The maximum path length to consider when calculating the closeness. If zero or negative then there is no such limit.

Details

The closeness centrality of a vertex is defined as the inverse of the sum of distances to all the other vertices in the graph:

\[ \frac{1}{\sum_{i \neq v} d_{vi}} \]

If there is no (directed) path between vertex v and i, then i is omitted from the calculation. If no other vertices are reachable from v, then its closeness is returned as NaN.

cutoff or smaller. This can be run for larger graphs, as the running time is not quadratic (if cutoff is small). If cutoff is zero or negative (which is the default), then the function calculates the exact closeness scores. Using zero as a cutoff is deprecated and future versions (from 1.4.0) will treat zero cutoff literally (i.e. no paths considered at all). If you want no cutoff, use a negative number. 

estimate_closeness is an alias for closeness with a different argument order, for sake of compatibility with older versions of igraph.

Closeness centrality is meaningful only for connected graphs. In disconnected graphs, consider using the harmonic centrality with harmonic_centrality.

Value

Numeric vector with the closeness values of all the vertices in v.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

betweenness, degree, harmonic_centrality
cluster_edge_betweenness

Examples

```r
  g <- make_ring(10)
  g2 <- make_star(10)
  closeness(g)
  closeness(g2, mode="in")
  closeness(g2, mode="out")
  closeness(g2, mode="all")
```

cluster_edge_betweenness

Community structure detection based on edge betweenness

Description

Many networks consist of modules which are densely connected themselves but sparsely connected to other modules.

Usage

```r
cluster_edge_betweenness(
  graph,
  weights = NULL,
  directed = TRUE,
  edge.betweenness = TRUE,
  merges = TRUE,
  bridges = TRUE,
  modularity = TRUE,
  membership = TRUE
)
```

Arguments

graph The graph to analyze.
weights The weights of the edges. It must be a positive numeric vector, NULL or NA. If it is NULL and the input graph has a ‘weight’ edge attribute, then that attribute will be used. If NULL and no such attribute is present, then the edges will have equal weights. Set this to NA if the graph was a ‘weight’ edge attribute, but you don’t want to use it for community detection. Edge weights are used to calculate weighted edge betweenness. This means that edges are interpreted as distances, not as connection strengths.
directed Logical constant, whether to calculate directed edge betweenness for directed graphs. It is ignored for undirected graphs.
edge.betweenness Logical constant, whether to return the edge betweenness of the edges at the time of their removal.
merges Logical constant, whether to return the merge matrix representing the hierarchical community structure of the network. This argument is called merges, even if the community structure algorithm itself is divisive and not agglomerative: it builds the tree from top to bottom. There is one line for each merge (i.e. split) in
matrix, the first line is the first merge (last split). The communities are identified by integer number starting from one. Community ids smaller than or equal to \( N \), the number of vertices in the graph, belong to singleton communities, i.e. individual vertices. Before the first merge we have \( N \) communities numbered from one to \( N \). The first merge, the first line of the matrix creates community \( N + 1 \), the second merge creates community \( N + 2 \), etc.

### bridges
Logical constant, whether to return a list the edge removals which actually split a component of the graph.

### modularity
Logical constant, whether to calculate the maximum modularity score, considering all possibly community structures along the edge-betweenness based edge removals.

### membership
Logical constant, whether to calculate the membership vector corresponding to the highest possible modularity score.

### Details
The edge betweenness score of an edge measures the number of shortest paths through it, see `edge_betweenness` for details. The idea of the edge betweenness based community structure detection is that it is likely that edges connecting separate modules have high edge betweenness as all the shortest paths from one module to another must traverse through them. So if we gradually remove the edge with the highest edge betweenness score we will get a hierarchical map, a rooted tree, called a dendrogram of the graph. The leaves of the tree are the individual vertices and the root of the tree represents the whole graph.

`cluster_edge_betweenness` performs this algorithm by calculating the edge betweenness of the graph, removing the edge with the highest edge betweenness score, then recalculating edge betweenness of the edges and again removing the one with the highest score, etc.

`edge.betweeness.community` returns various information collected through the run of the algorithm. See the return value down here.

### Value
`cluster_edge_betweenness` returns a `communities` object, please see the `communities` manual page for details.

### Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

### References

### See Also
* `edge_betweenness` for the definition and calculation of the edge betweenness, `cluster_walktrap`, `cluster_fast_greedy`, `cluster_leading_eigen` for other community detection methods.

See `communities` for extracting the results of the community detection.
Examples

```r
g <- sample_pa(100, m = 2, directed = FALSE)
ec <- cluster_edge_betweenness(g)

g <- make_full_graph(10) %du% make_full_graph(10)
g <- add_edges(g, c(1,11))
ec <- cluster_edge_betweenness(g)
ec
```

cluster_fast_greedy  
Community structure via greedy optimization of modularity

Description

This function tries to find dense subgraph, also called communities in graphs via directly optimizing a modularity score.

Usage

```r
cluster_fast_greedy(
  graph,
  merges = TRUE,
  modularity = TRUE,
  membership = TRUE,
  weights = NULL
)
```

Arguments

- **graph**: The input graph
- **merges**: Logical scalar, whether to return the merge matrix.
- **modularity**: Logical scalar, whether to return a vector containing the modularity after each merge.
- **membership**: Logical scalar, whether to calculate the membership vector corresponding to the maximum modularity score, considering all possible community structures along the merges.
- **weights**: The weights of the edges. It must be a positive numeric vector, NULL or NA. If it is NULL and the input graph has a ‘weight’ edge attribute, then that attribute will be used. If NULL and no such attribute is present, then the edges will have equal weights. Set this to NA if the graph was a ‘weight’ edge attribute, but you don’t want to use it for community detection. A larger edge weight means a stronger connection for this function.

Details

cluster_fluid_communities

Value

cluster_fast_greedy returns a communities object, please see the communities manual page for details.

Author(s)

Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com> for the R interface.

References


See Also

communities for extracting the results.

See also cluster_walktrap, cluster_spinglass, cluster_leading_eigen and cluster_edge_betweenness, cluster_louvain cluster_leiden for other methods.

Examples

g <- make_full_graph(5) %du% make_full_graph(5) %du% make_full_graph(5)
g <- add_edges(g, c(1,6, 1,11, 6, 11))
fc <- cluster_fast_greedy(g)
membership(fc)
sizes(fc)

Description

The algorithm detects communities based on the simple idea of several fluids interacting in a non-homogeneous environment (the graph topology), expanding and contracting based on their interaction and density.

Usage

cluster_fluid_communities(graph, no.of.communities)

Arguments

graph The input graph. The graph must be simple and connected. Empty graphs are not supported as well as single vertex graphs. Edge directions are ignored. Weights are not considered.

no.of.communities The number of communities to be found. Must be greater than 0 and fewer than number of vertices in the graph.
Value

cluster_fluid_communities returns a communities object, please see the communities manual page for details.

Author(s)

Ferran Parés

References


See Also

See communities for extracting the membership, modularity scores, etc. from the results.

Other community detection algorithms: cluster_walktrap, cluster_spinglass, cluster_leading_eigen, cluster_edge_betweenness, cluster_fast_greedy, cluster_label_prop, cluster_louvain, cluster_leiden

Examples

g <- graph.famous("Zachary")
comms <- cluster_fluid_communities(g, 2)

Description

Find community structure that minimizes the expected description length of a random walker trajectory

Usage

cluster_infomap(
  graph,
  e.weights = NULL,
  v.weights = NULL,
  nb.trials = 10,
  modularity = TRUE
)

Arguments

- **graph**: The input graph.
- **e.weights**: If not `NULL`, then a numeric vector of edge weights. The length must match the number of edges in the graph. By default the ‘weight’ edge attribute is used as weights. If it is not present, then all edges are considered to have the same weight. Larger edge weights correspond to stronger connections.
- **v.weights**: If not `NULL`, then a numeric vector of vertex weights. The length must match the number of vertices in the graph. By default the ‘weight’ vertex attribute is used as weights. If it is not present, then all vertices are considered to have the same weight. A larger vertex weight means a larger probability that the random surfer jumps to that vertex.
- **nb.trials**: The number of attempts to partition the network (can be any integer value equal or larger than 1).
- **modularity**: Logical scalar, whether to calculate the modularity score of the detected community structure.

Details

Please see the details of this method in the references given below.

Value

`cluster_infomap` returns a `communities` object, please see the `communities` manual page for details.

Author(s)

Martin Rosvall wrote the original C++ code. This was ported to be more igraph-like by Emmanuel Navarro. The R interface and some cosmetics was done by Gabor Csardi `<csardi.gabor@gmail.com>`.

References


See Also

Other community finding methods and `communities`.

Examples

```r
## Zachary's karate club
g <- make_graph("Zachary")
imc <- cluster_infomap(g)
membership(imc)
communities(imc)
```
cluster_label_prop  Finding communities based on propagating labels

Description

This is a fast, nearly linear time algorithm for detecting community structure in networks. In works by labeling the vertices with unique labels and then updating the labels by majority voting in the neighborhood of the vertex.

Usage

cluster_label_prop(graph, weights = NULL, initial = NULL, fixed = NULL)

Arguments

graph               The input graph, should be undirected to make sense.
weights             The weights of the edges. It must be a positive numeric vector, NULL or NA. If it is NULL and the input graph has a ‘weight’ edge attribute, then that attribute will be used. If NULL and no such attribute is present, then the edges will have equal weights. Set this to NA if the graph was a ‘weight’ edge attribute, but you don’t want to use it for community detection. A larger edge weight means a stronger connection for this function.
initial             The initial state. If NULL, every vertex will have a different label at the beginning. Otherwise it must be a vector with an entry for each vertex. Non-negative values denote different labels, negative entries denote vertices without labels.
fixed               Logical vector denoting which labels are fixed. Of course this makes sense only if you provided an initial state, otherwise this element will be ignored. Also note that vertices without labels cannot be fixed.

Details

This function implements the community detection method described in: Raghavan, U.N. and Albert, R. and Kumara, S.: Near linear time algorithm to detect community structures in large-scale networks. Phys Rev E 76, 036106. (2007). This version extends the original method by the ability to take edge weights into consideration and also by allowing some labels to be fixed.

From the abstract of the paper: “In our algorithm every node is initialized with a unique label and at every step each node adopts the label that most of its neighbors currently have. In this iterative process densely connected groups of nodes form a consensus on a unique label to form communities.”

Value

cluster_label_prop returns a communities object, please see the communities manual page for details.

Author(s)

Tamas Nepusz <ntamas@gmail.com> for the C implementation, Gabor Csardi <csardi.gabor@gmail.com> for this manual page.
References


See Also

`communities` for extracting the actual results.
`cluster_fast_greedy`, `cluster_walktrap`, `cluster_spinglass`, `cluster_louvain` and `cluster_leiden` for other community detection methods.

Examples

```r
  g <- sample_gnp(10, 5/10) %du% sample_gnp(9, 5/9)
  g <- add_edges(g, c(1, 12))
  cluster_label_prop(g)
```

---

**cluster_leading_eigen**  Community structure detecting based on the leading eigenvector of the community matrix

Description

This function tries to find densely connected subgraphs in a graph by calculating the leading non-negative eigenvector of the modularity matrix of the graph.

Usage

```r
  cluster_leading_eigen(
    graph, steps = -1, weights = NULL, start = NULL, options = arpack_defaults, callback = NULL, extra = NULL, env = parent.frame()
  )
```

Arguments

- **graph**  The input graph. Should be undirected as the method needs a symmetric matrix.
- **steps**  The number of steps to take, this is actually the number of tries to make a step. It is not a particularly useful parameter.
- **weights**  The weights of the edges. It must be a positive numeric vector, NULL or NA. If it is NULL and the input graph has a 'weight' edge attribute, then that attribute will be used. If NULL and no such attribute is present, then the edges will have equal weights. Set this to NA if the graph was a 'weight' edge attribute, but you don’t want to use it for community detection. A larger edge weight means a stronger connection for this function.
cluster_leading_eigen

start NULL, or a numeric membership vector, giving the start configuration of the algorithm.

options A named list to override some ARPACK options.

callback If not NULL, then it must be callback function. This is called after each iteration, after calculating the leading eigenvector of the modularity matrix. See details below.

extra Additional argument to supply to the callback function.

eval The environment in which the callback function is evaluated.

Details

The function documented in these section implements the ‘leading eigenvector’ method developed by Mark Newman, see the reference below.

The heart of the method is the definition of the modularity matrix, B, which is B=A-P, A being the adjacency matrix of the (undirected) network, and P contains the probability that certain edges are present according to the ‘configuration model’. In other words, a P[i, j] element of P is the probability that there is an edge between vertices i and j in a random network in which the degrees of all vertices are the same as in the input graph.

The leading eigenvector method works by calculating the eigenvector of the modularity matrix for the largest positive eigenvalue and then separating vertices into two community based on the sign of the corresponding element in the eigenvector. If all elements in the eigenvector are of the same sign that means that the network has no underlying community structure. Check Newman’s paper to understand why this is a good method for detecting community structure.

Value

cluster_leading_eigen returns a named list with the following members:

membership The membership vector at the end of the algorithm, when no more splits are possible.

merges The merges matrix starting from the state described by the membership member. This is a two-column matrix and each line describes a merge of two communities, the first line is the first merge and it creates community 'N', N is the number of initial communities in the graph, the second line creates community N+1, etc.

options Information about the underlying ARPACK computation, see arpack for details.

Callback functions

The callback argument can be used to supply a function that is called after each eigenvector calculation. The following arguments are supplied to this function:

membership The actual membership vector, with zero-based indexing.

community The community that the algorithm just tried to split, community numbering starts with zero here.

value The eigenvalue belonging to the leading eigenvector the algorithm just found.

vector The leading eigenvector the algorithm just found.

multiplier An R function that can be used to multiply the actual modularity matrix with an arbitrary vector. Supply the vector as an argument to perform this multiplication. This function can be used with ARPACK.

extra The extra argument that was passed to cluster_leading_eigen. The callback function should return a scalar number. If this number is non-zero, then the clustering is terminated.
cluster_leiden

Finding community structure of a graph using the Leiden algorithm of Traag, van Eck & Waltman.

Description

The Leiden algorithm is similar to the Louvain algorithm, cluster_louvain, but it is faster and yields higher quality solutions. It can optimize both modularity and the Constant Potts Model, which does not suffer from the resolution-limit (see preprint http://arxiv.org/abs/1104.3083).

Usage

cluster_leiden(
  graph, 
  objective_function = c("CPM", "modularity"), 
  weights = NULL, 
  resolution_parameter = 1, 
  beta = 0.01, 
  initial_membership = NULL, 
  n_iterations = 2, 
  vertex_weights = NULL 
)

Arguments

graph The input graph, only undirected graphs are supported.
objective_function Whether to use the Constant Potts Model (CPM) or modularity. Must be either "CPM" or "modularity".
The weights of the edges. It must be a positive numeric vector, `NULL` or `NA`. If it is `NULL` and the input graph has a `‘weight’` edge attribute, then that attribute will be used. If `NULL` and no such attribute is present, then the edges will have equal weights. Set this to `NA` if the graph was a `‘weight’` edge attribute, but you don’t want to use it for community detection. A larger edge weight means a stronger connection for this function.

The resolution parameter to use. Higher resolutions lead to more smaller communities, while lower resolutions lead to fewer larger communities.

Parameter affecting the randomness in the Leiden algorithm. This affects only the refinement step of the algorithm.

If provided, the Leiden algorithm will try to improve this provided membership. If no argument is provided, the algorithm simply starts from the singleton partition.

the number of iterations to iterate the Leiden algorithm. Each iteration may improve the partition further.

the vertex weights used in the Leiden algorithm. If this is not provided, it will be automatically determined on the basis of the `objective_function`. Please see the details of this function how to interpret the vertex weights.

The Leiden algorithm consists of three phases: (1) local moving of nodes, (2) refinement of the partition and (3) aggregation of the network based on the refined partition, using the non-refined partition to create an initial partition for the aggregate network. In the local move procedure in the Leiden algorithm, only nodes whose neighborhood has changed are visited. The refinement is done by restarting from a singleton partition within each cluster and gradually merging the subclusters. When aggregating, a single cluster may then be represented by several nodes (which are the subclusters identified in the refinement).

The Leiden algorithm provides several guarantees. The Leiden algorithm is typically iterated: the output of one iteration is used as the input for the next iteration. At each iteration all clusters are guaranteed to be connected and well-separated. After an iteration in which nothing has changed, all nodes and some parts are guaranteed to be locally optimally assigned. Finally, asymptotically, all subsets of all clusters are guaranteed to be locally optimally assigned. For more details, please see Traag, Waltman & van Eck (2019).

The objective function being optimized is

$$
\frac{1}{2m} \sum_{ij} (A_{ij} - \gamma n_i n_j) \delta(\sigma_i, \sigma_j)
$$

where \(m\) is the total edge weight, \(A_{ij}\) is the weight of edge \((i, j)\), \(\gamma\) is the so-called resolution parameter, \(n_i\) is the node weight of node \(i\), \(\sigma_i\) is the cluster of node \(i\) and \(\delta(x, y) = 1\) if and only if \(x = y\) and 0 otherwise. By setting \(n_i = k_i\), the degree of node \(i\), and dividing \(\gamma\) by \(2m\), you effectively obtain an expression for modularity.

Hence, the standard modularity will be optimized when you supply the degrees as `vertex_weights` and by supplying as a resolution parameter \(\frac{\gamma}{2m}\), with \(m\) the number of edges. If you do not specify any `vertex_weights`, the correct vertex weights and scaling of \(\gamma\) is determined automatically by the `objective_function` argument.
cluster_louvain

Value

cluster_leiden returns a communities object, please see the communities manual page for details.

Author(s)

Vincent Traag

References


See Also

See communities for extracting the membership, modularity scores, etc. from the results.

Other community detection algorithms: cluster_walktrap, cluster_spinglass, cluster_leading_eigen, cluster_edge_betweenness, cluster_fast_greedy, cluster_label_prop cluster_louvain cluster_fluid_communities cluster_infomap cluster_optimal cluster_walktrap

Examples

g <- make_graph("Zachary")
# By default CPM is used
r <- quantile(strength(g))[2] / (gorder(g) - 1)
# Set seed for sake of reproducibility
set.seed(1)
ldc <- cluster_leiden(g, resolution_parameter=r)
print(ldc)
plot(ldc, g)

cluster_louvain

Finding community structure by multi-level optimization of modularity

Description

This function implements the multi-level modularity optimization algorithm for finding community structure, see references below. It is based on the modularity measure and a hierarchical approach.

Usage

custom_louvain(graph, weights = NULL, resolution = 1)

Arguments

graph

The input graph.

weights

The weights of the edges. It must be a positive numeric vector, NULL or NA. If it is NULL and the input graph has a ‘weight’ edge attribute, then that attribute will be used. If NULL and no such attribute is present, then the edges will have equal weights. Set this to NA if the graph was a ‘weight’ edge attribute, but you don’t want to use it for community detection. A larger edge weight means a stronger connection for this function.
Optional resolution parameter that allows the user to adjust the resolution parameter of the modularity function that the algorithm uses internally. Lower values typically yield fewer, larger clusters. The original definition of modularity is recovered when the resolution parameter is set to 1.

Details


It is based on the modularity measure and a hierarchical approach. Initially, each vertex is assigned to a community on its own. In every step, vertices are re-assigned to communities in a local, greedy way: each vertex is moved to the community with which it achieves the highest contribution to modularity. When no vertices can be reassigned, each community is considered a vertex on its own, and the process starts again with the merged communities. The process stops when there is only a single vertex left or when the modularity cannot be increased any more in a step. Since igraph 1.3, vertices are processed in a random order.

This function was contributed by Tom Gregorovic.

Value

cluster_louvain returns a communities object, please see the communities manual page for details.

Author(s)

Tom Gregorovic, Tamas Nepusz <ntamas@gmail.com>

References


See Also

See communities for extracting the membership, modularity scores, etc. from the results.

Other community detection algorithms: cluster_walktrap, cluster_spinglass, cluster_leading_eigen, cluster_edge_betweenness, cluster_fast_greedy, cluster_label_prop, cluster_leiden

Examples

# This is so simple that we will have only one level
g <- make_full_graph(5) %du% make_full_graph(5) %du% make_full_graph(5)
g <- add_edges(g, c(1,6, 1,11, 6, 11))
cluster_louvain(g)
cluster_optimal

Description

This function calculates the optimal community structure of a graph, by maximizing the modularity measure over all possible partitions.

Usage

cluster_optimal(graph, weights = NULL)

Arguments

- **graph**: The input graph. Edge directions are ignored for directed graphs.
- **weights**: The weights of the edges. It must be a positive numeric vector, NULL or NA. If it is NULL and the input graph has a ‘weight’ edge attribute, then that attribute will be used. If NULL and no such attribute is present, then the edges will have equal weights. Set this to NA if the graph was a ‘weight’ edge attribute, but you don’t want to use it for community detection. A larger edge weight means a stronger connection for this function.

Details

This function calculates the optimal community structure for a graph, in terms of maximal modularity score.

The calculation is done by transforming the modularity maximization into an integer programming problem, and then calling the GLPK library to solve that. Please the reference below for details.

Note that modularity optimization is an NP-complete problem, and all known algorithms for it have exponential time complexity. This means that you probably don’t want to run this function on larger graphs. Graphs with up to fifty vertices should be fine, graphs with a couple of hundred vertices might be possible.

Value

cluster_optimal returns a communities object, please see the communities manual page for details.

Examples

```r
## Zachary's karate club
g <- make_graph("Zachary")

## We put everything into a big 'try' block, in case
## igraph was compiled without GLPK support

## The calculation only takes a couple of seconds
oc <- cluster_optimal(g)

## Double check the result
```

```r
print(modularity(oc))
print(modularity(g, membership(oc)))

## Compare to the greedy optimizer
fc <- cluster_fast_greedy(g)
print(modularity(fc))
```

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**References**


**See Also**

- `communities` for the documentation of the result, `modularity`. See also `cluster_fast_greedy` for a fast greedy optimizer.

---

**cluster_spinglass**

*Finding communities in graphs based on statistical mechanics*

**Description**

This function tries to find communities in graphs via a spin-glass model and simulated annealing.

**Usage**

```r
cluster_spinglass(
  graph,
  weights = NULL,
  vertex = NULL,
  spins = 25,
  parupdate = FALSE,
  start.temp = 1,
  stop.temp = 0.01,
  cool.fact = 0.99,
  update.rule = c("config", "random", "simple"),
  gamma = 1,
  implementation = c("orig", "neg"),
  gamma.minus = 1
)
```

**Arguments**

- `graph` The input graph, can be directed but the direction of the edges is neglected.
weights: The weights of the edges. It must be a positive numeric vector, NULL or NA. If it is NULL and the input graph has a ‘weight’ edge attribute, then that attribute will be used. If NULL and no such attribute is present, then the edges will have equal weights. Set this to NA if the graph was a ‘weight’ edge attribute, but you don’t want to use it for community detection. A larger edge weight means a stronger connection for this function.

vertex: This parameter can be used to calculate the community of a given vertex without calculating all communities. Note that if this argument is present then some other arguments are ignored.

spins: Integer constant, the number of spins to use. This is the upper limit for the number of communities. It is not a problem to supply a (reasonably) big number here, in which case some spin states will be unpopulated.

parupdate: Logical constant, whether to update the spins of the vertices in parallel (synchronously) or not. This argument is ignored if the second form of the function is used (i.e. the ‘vertex’ argument is present). It is also not implemented in the “neg” implementation.

start.temp: Real constant, the start temperature. This argument is ignored if the second form of the function is used (i.e. the ‘vertex’ argument is present).

stop.temp: Real constant, the stop temperature. The simulation terminates if the temperature lowers below this level. This argument is ignored if the second form of the function is used (i.e. the ‘vertex’ argument is present).

cool.fact: Cooling factor for the simulated annealing. This argument is ignored if the second form of the function is used (i.e. the ‘vertex’ argument is present).

update.rule: Character constant giving the ‘null-model’ of the simulation. Possible values: “simple” and “config”. “simple” uses a random graph with the same number of edges as the baseline probability and “config” uses a random graph with the same vertex degrees as the input graph.

gamma: Real constant, the gamma argument of the algorithm. This specifies the balance between the importance of present and non-present edges in a community. Roughly, a community is a set of vertices having many edges inside the community and few edges outside the community. The default 1.0 value makes existing and non-existing links equally important. Smaller values make the existing links, greater values make the missing links more important.

implementation: Character scalar. Currently igraph contains two implementations for the Spinglass community finding algorithm. The faster original implementation is the default. The other implementation, that takes into account negative weights, can be chosen by supplying ‘neg’ here.

gamma.minus: Real constant, the gamma.minus parameter of the algorithm. This specifies the balance between the importance of present and non-present negative weighted edges in a community. Smaller values of gamma.minus, leads to communities with lesser negative intra-connectivity. If this argument is set to zero, the algorithm reduces to a graph coloring algorithm, using the number of spins as the number of colors. This argument is ignored if the ‘orig’ implementation is chosen.

Details

This function tries to find communities in a graph. A community is a set of nodes with many edges inside the community and few edges between outside it (i.e. between the community itself and the rest of the graph.)
This idea is reversed for edges having a negative weight, i.e. few negative edges inside a community and many negative edges between communities. Note that only the ‘neg’ implementation supports negative edge weights.

The `spinglass.community` function can solve two problems related to community detection. If the `vertex` argument is not given (or it is `NULL`), then the regular community detection problem is solved (approximately), i.e. partitioning the vertices into communities, by optimizing the an energy function.

If the `vertex` argument is given and it is not `NULL`, then it must be a vertex id, and the same energy function is used to find the community of the the given vertex. See also the examples below.

**Value**

If the `vertex` argument is not given, i.e. the first form is used then a `cluster_spinglass` returns a `communities` object.

If the `vertex` argument is present, i.e. the second form is used then a named list is returned with the following components:

- `community` Numeric vector giving the ids of the vertices in the same community as `vertex`.
- `cohesion` The cohesion score of the result, see references.
- `adhesion` The adhesion score of the result, see references.
- `inner.links` The number of edges within the community of `vertex`.
- `outer.links` The number of edges between the community of `vertex` and the rest of the graph.

**Author(s)**

Jorg Reichardt for the original code and Gabor Csardi (<csardi.gabor@gmail.com>) for the igraph glue code.

Changes to the original function for including the possibility of negative ties were implemented by Vincent Traag (http://www.traag.net/).

**References**


**See Also**

`communities`, `components`

**Examples**

```r
g <- sample_gnp(10, 5/10) %du% sample_gnp(9, 5/9)
g <- add_edges(g, c(1, 12))
g <- induced_subgraph(g, subcomponent(g, 1))
cluster_spinglass(g, spins=2)
cluster_spinglass(g, vertex=1)
```
cluster_walktrap  Community structure via short random walks

Description

This function tries to find densely connected subgraphs, also called communities in a graph via random walks. The idea is that short random walks tend to stay in the same community.

Usage

```r
cluster_walktrap(
  graph,
  weights = NULL,
  steps = 4,
  merges = TRUE,
  modularity = TRUE,
  membership = TRUE
)
```

Arguments

- **graph**: The input graph, edge directions are ignored in directed graphs.
- **weights**: The weights of the edges. It must be a positive numeric vector, NULL or NA. If it is NULL and the input graph has a ‘weight’ edge attribute, then that attribute will be used. If NULL and no such attribute is present, then the edges will have equal weights. Set this to NA if the graph was a ‘weight’ edge attribute, but you don’t want to use it for community detection. Larger edge weights increase the probability that an edge is selected by the random walker. In other words, larger edge weights correspond to stronger connections.
- **steps**: The length of the random walks to perform.
- **merges**: Logical scalar, whether to include the merge matrix in the result.
- **modularity**: Logical scalar, whether to include the vector of the modularity scores in the result. If the `membership` argument is true, then it will always be calculated.
- **membership**: Logical scalar, whether to calculate the membership vector for the split corresponding to the highest modularity value.

Details

This function is the implementation of the Walktrap community finding algorithm, see Pascal Pons, Matthieu Latapy: Computing communities in large networks using random walks, https://arxiv.org/abs/physics/0512106

Value

`cluster_walktrap` returns a `communities` object, please see the `communities` manual page for details.

Author(s)

Pascal Pons (http://psl.ons.free.fr/) and Gabor Csardi <csardi.gabor@gmail.com> for the R and igraph interface.
References


See Also

See communities on getting the actual membership vector, merge matrix, modularity score, etc. 
modularity and cluster_fast_greedy, cluster_spinglass, cluster_leading_eigen, cluster_edge_betweenness, cluster_louvain, and cluster_leiden for other community detection methods.

Examples

g <- make_full_graph(5) %du% make_full_graph(5) %du% make_full_graph(5)
g <- add_edges(g, c(1,6, 1,11, 6, 11))
cluster_walktrap(g)

cocitation

Cocitation coupling

Description

Two vertices are cocited if there is another vertex citing both of them. cocitation simply counts how many types two vertices are cocited. The bibliographic coupling of two vertices is the number of other vertices they both cite, bibcoupling calculates this.

Usage

cocitation(graph, v = V(graph))

Arguments

graph The graph object to analyze
v Vertex sequence or numeric vector, the vertex ids for which the cocitation or bibliographic coupling values we want to calculate. The default is all vertices.

Details

cocitation calculates the cocitation counts for the vertices in the v argument and all vertices in the graph.
bibcoupling calculates the bibliographic coupling for vertices in v and all vertices in the graph.
Calculating the cocitation or bibliographic coupling for only one vertex costs the same amount of computation as for all vertices. This might change in the future.

Value

A numeric matrix with length(v) lines and vcount(graph) columns. Element (i,j) contains the cocitation or bibliographic coupling for vertices v[i] and j.
Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

Examples

```
g <- make_kautz_graph(2,3)
cocitation(g)
bibcoupling(g)
```

---

**cohesive_blocks**

*Calculate Cohesive Blocks*

**Description**

Calculates cohesive blocks for objects of class `igraph`.

**Usage**

```r
cohesive_blocks(graph, labels = TRUE)
```
plot_hierarchy(
  blocks,
  layout = layout_as_tree(hierarchy(blocks), root = 1),
  ...
)

export_pajek(blocks, graph, file, project.file = TRUE)

max_cohesion(blocks)

Arguments

graph For cohesive_blocks a graph object of class igraph. It must be undirected and simple. (See is_simple.)

For graphs_from_cohesive_blocks and export_pajek the same graph must be supplied whose cohesive block structure is given in the blocks argument.

labels Logical scalar, whether to add the vertex labels to the result object. These labels can be then used when reporting and plotting the cohesive blocks.

blocks, x, object A cohesiveBlocks object, created with the cohesive_blocks function.

... Additional arguments. plot_hierarchy and plot pass them to plot.igraph. print and summary ignore them.

y The graph whose cohesive blocks are supplied in the x argument.

colbar Color bar for the vertex colors. Its length should be at least \( m + 1 \), where \( m \) is the maximum cohesion in the graph. Alternatively, the vertex colors can also be directly specified via the col argument.

col A vector of vertex colors, in any of the usual formats. (Symbolic color names (e.g. ‘red’, ‘blue’, etc.), RGB colors (e.g. '#FF9900FF'), integer numbers referring to the current palette. By default the given colbar is used and vertices with the same maximal cohesion will have the same color.

mark.groups A list of vertex sets to mark on the plot by circling them. By default all cohesive blocks are marked, except the one corresponding to the all vertices.

layout The layout of a plot, it is simply passed on to plot.igraph, see the possible formats there. By default the Reingold-Tilford layout generator is used.

file Defines the file (or connection) the Pajek file is written to.

If the project.file argument is TRUE, then it can be a filename (with extension), a file object, or in general any kind of connection object. The file/connection will be opened if it wasn’t already.

If the project.file argument is FALSE, then several files are created and file must be a character scalar containing the base name of the files, without extension. (But it can contain the path to the files.)

See also details below.

project.file Logical scalar, whether to create a single Pajek project file containing all the data, or to create separated files for each item. See details below.

Details

Cohesive blocking is a method of determining hierarchical subsets of graph vertices based on their structural cohesion (or vertex connectivity). For a given graph \( G \), a subset of its vertices \( S \subset V(G) \)
is said to be maximally \( k \)-cohesive if there is no superset of \( S \) with vertex connectivity greater than or equal to \( k \). Cohesive blocking is a process through which, given a \( k \)-cohesive set of vertices, maximally \( l \)-cohesive subsets are recursively identified with \( l > k \). Thus a hierarchy of vertex subsets is found, with the entire graph \( G \) at its root.

The function `cohesive_blocks` implements cohesive blocking. It returns a cohesiveBlocks object. cohesiveBlocks should be handled as an opaque class, i.e. its internal structure should not be accessed directly, but through the functions listed here.

The function `length` can be used on cohesiveBlocks objects and it gives the number of blocks.

The function `blocks` returns the actual blocks stored in the cohesiveBlocks object. They are returned in a list of numeric vectors, each containing vertex ids.

The function `graphs_from_cohesive_blocks` is similar, but returns the blocks as (induced) subgraphs of the input graph. The various (graph, vertex and edge) attributes are kept in the subgraph.

The function `cohesion` returns a numeric vector, the cohesion of the different blocks. The order of the blocks is the same as for the blocks and graphs_from_cohesive_blocks functions.

The block hierarchy can be queried using the hierarchy function. It returns an igraph graph, its vertex ids are ordered according the order of the blocks in the blocks and graphs_from_cohesive_blocks, cohesion, etc. functions.

`parent` gives the parent vertex of each block, in the block hierarchy, for the root vertex it gives 0.

`plot_hierarchy` plots the hierarchy tree of the cohesive blocks on the active graphics device, by calling `igraph.plot`.

The export_pajek function can be used to export the graph and its cohesive blocks in Pajek format. It can either export a single Pajek project file with all the information, or a set of files, depending on its project.file argument. If project.file is TRUE, then the following information is written to the file (or connection) given in the file argument: (1) the input graph, together with its attributes, see `write_graph` for details; (2) the hierarchy graph; and (3) one binary partition for each cohesive block. If project.file is FALSE, then the file argument must be a character scalar and it is used as the base name for the generated files. If file is ‘basename’, then the following files are created: (1) ‘basename.net’ for the original graph; (2) ‘basename_hierarchy.net’ for the hierarchy graph; (3) ‘basename_block_x.net’ for each cohesive block, where ‘x’ is the number of the block, starting with one.

`max_cohesion` returns the maximal cohesion of each vertex, i.e. the cohesion of the most cohesive block of the vertex.

The generic function `summary` works on cohesiveBlocks objects and it prints a one line summary to the terminal.

The generic function print is also defined on cohesiveBlocks objects and it is invoked automatically if the name of the cohesiveBlocks object is typed in. It produces an output like this:

```
Cohesive block structure:
B-1 c 1, n 23
  ' - B-2 c 2, n 14 ooooooo.. .o.......oo ooo
  ' - B-4 c 5, n 7 ooooooo... ............ ...
  ' - B-3 c 2, n 10 ......o.o 0.oooooo... ...
  ' - B-5 c 3, n 4 ......o.o oo o...........
```

The left part shows the block structure, in this case for five blocks. The first block always corresponds to the whole graph, even if its cohesion is zero. Then cohesion of the block and the number of vertices in the block are shown. The last part is only printed if the display is wide enough and shows the vertices in the blocks, ordered by vertex ids. ‘o’ means that the vertex is included, a dot means that it is not, and the vertices are shown in groups of ten.

The generic function `plot` plots the graph, showing one or more cohesive blocks in it.
cohesive_blocks

Value

cohesive_blocks returns a cohesiveBlocks object.
blocks returns a list of numeric vectors, containing vertex ids.
graphs_from_cohesive_blocks returns a list of igraph graphs, corresponding to the cohesive blocks.
cohesion returns a numeric vector, the cohesion of each block.
hierarchy returns an igraph graph, the representation of the cohesive block hierarchy.
parent returns a numeric vector giving the parent block of each cohesive block, in the block hierarchy. The block at the root of the hierarchy has no parent and 0 is returned for it.

Author(s)

Gabor Csardi (<csardi.gabor@gmail.com>) for the current implementation, Peter McMahan (https://socialsciences.uchicago.edu/news/alumni-profile-peter-mcmahan-phd17-sociology) wrote the first version in R.

References


See Also

cohesion

Examples

```r
## The graph from the Moody-White paper

mwBlocks <- cohesive_blocks(mw)

# Inspect block membership and cohesion
mwBlocks
do(mwBlocks)
do(cohesion(mwBlocks))

# Save results in a Pajek file
## Not run:
export_pajek(mwBlocks, mw, file="/tmp/mwBlocks.paj")

## End(Not run)
```
# Plot the results
plot(mwBlocks, mw)

## The science camp network

camp <- graph_from_literal(Harry:Steve:Don:Bert - Harry:Steve:Don:Bert,
Pam:Brazey:Carol:Pat - Pam:Brazey:Carol:Pat,
Holly - Carol:Pat:Pam:Jennie:Bill,
Bill - Pauline:Michael:Lee:Holly,
Pauline - Bill:Jennie:Ann,
Jennie - Holly:Michael:Lee:Ann:Pauline,
Michael - Bill:Jennie:Ann:Lee:John,
Ann - Michael:Jennie:Pauline,
Lee - Michael:Bill:Jennie,
Gery - Pat:Steve:Russ:John,
Russ - Steve: Bert: Gery: John,
John - Gery: Russ: Michael)

campBlocks <- cohesive_blocks(camp)
campBlocks

plot(campBlocks, camp, vertex.label=V(camp)$name, margin=-0.2,
vertex.shape="rectangle", vertex.size=24, vertex.size2=8,
mark.border=1, colbar=c(NA, NA, "cyan", "orange") )

---

**compare**  

Compares community structures using various metrics

Description

This function assesses the distance between two community structures.

Usage

```r
compare(
  comm1,
  comm2,
  method = c("vi", "nmi", "split.join", "rand", "adjusted.rand")
)
```

Arguments

- **comm1**: A `communities` object containing a community structure; or a numeric vector, the membership vector of the first community structure. The membership vector should contain the community id of each vertex, the numbering of the communities starts with one.

- **comm2**: A `communities` object containing a community structure; or a numeric vector, the membership vector of the second community structure, in the same format as for the previous argument.

- **method**: Character scalar, the comparison method to use. Possible values: ‘vi’ is the variation of information (VI) metric of Meila (2003), ‘nmi’ is the normalized mutual information measure proposed by Danon et al. (2005), ‘split.join’ is the split-join distance of can Dongen (2000), ‘rand’ is the Rand index of Rand (1971), ‘adjusted.rand’ is the adjusted Rand index by Hubert and Arabie (1985).
**complementer**

**Value**

A real number.

**Author(s)**

Tamas Nepusz <ntamas@gmail.com>

**References**


**See Also**

See `cluster_walktrap, cluster_spinglass, cluster_leading_eigen, cluster_edge_betweenness, cluster_fast_greedy, cluster_label_prop` for various community detection methods.

**Examples**

```r
g <- make_graph("Zachary")
sg <- cluster_spinglass(g)
le <- cluster_leading_eigen(g)
compare(sg, le, method="rand")
compare(membership(sg), membership(le))
```

---

**complementer Complementer of a graph**

**Description**

A complementer graph contains all edges that were not present in the input graph.

**Usage**

`complementer(graph, loops = FALSE)`
component_distribution

Connected components of a graph

Description

Calculate the maximal (weakly or strongly) connected components of a graph

Usage

component_distribution(graph, cumulative = FALSE, mul.size = FALSE, ...)
components(graph, mode = c("weak", "strong"))
is_connected(graph, mode = c("weak", "strong"))
count_components(graph, mode = c("weak", "strong"))

Arguments

graph The input graph, can be directed or undirected.
loops Logical constant, whether to generate loop edges.

Details

complementer creates the complementer of a graph. Only edges which are not present in the original graph will be included in the new graph.
complementer keeps graph and vertex attributes, edge attributes are lost.

Value

A new graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

## Complementer of a ring

```r
g <- make_ring(10)
complementer(g)
```

## A graph and its complementer give together the full graph

g <- make_ring(10)
gc <- complementer(g)
gu <- union(g, gc)
gu
graph.isomorphic(gu, make_full_graph(vcount(g)))
```
Arguments

- **graph**: The graph to analyze.
- **cumulative**: Logical, if TRUE the cumulative distribution (relative frequency) is calculated.
- **mul.size**: Logical. If TRUE the relative frequencies will be multiplied by the cluster sizes.
- **mode**: Character string, either “weak” or “strong”. For directed graphs “weak” implies weakly, “strong” strongly connected components to search. It is ignored for undirected graphs.

Details

- **is_connected**: decides whether the graph is weakly or strongly connected. The null graph is considered disconnected.
- **components**: finds the maximal (weakly or strongly) connected components of a graph.
- **count_components**: does almost the same as **components** but returns only the number of clusters found instead of returning the actual clusters.
- **component_distribution**: creates a histogram for the maximal connected component sizes.

The weakly connected components are found by a simple breadth-first search. The strongly connected components are implemented by two consecutive depth-first searches.

Value

For **is_connected** a logical constant.

For **components** a named list with three components:

- **membership**: numeric vector giving the cluster id to which each vertex belongs.
- **csize**: numeric vector giving the sizes of the clusters.
- **no**: numeric constant, the number of clusters.

For **count_components** an integer constant is returned.

For **component_distribution** a numeric vector with the relative frequencies. The length of the vector is the size of the largest component plus one. Note that (for currently unknown reasons) the first element of the vector is the number of clusters of size zero, so this is always zero.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

decompose, subcomponent, groups

Examples

```r
g <- sample_gnp(20, 1/20)
clu <- components(g)
groups(clu)
```
Description

This is a layout modifier function, and it can be used to calculate the layout separately for each component of the graph.

Usage

\[
\text{component\_wise(merge\_method = "dla")}
\]

Arguments

merge\_method Merging algorithm, the method argument of \text{merge\_coords}.

See Also

merge\_coords, layout\_.

Other layout modifiers: \text{normalize()}.

Other graph layouts: \text{add\_layout()}, \text{layout\_as\_bipartite()}, \text{layout\_as\_star()}, \text{layout\_as\_tree()}, \text{layout\_in\_circle()}, \text{layout\_nicely()}, \text{layout\_on\_grid()}, \text{layout\_on\_sphere()}, \text{layout\_randomly()}, \text{layout\_with\_dh()}, \text{layout\_with\_fr()}, \text{layout\_with\_gem()}, \text{layout\_with\_graphopt()}, \text{layout\_with\_kk()}, \text{layout\_with\_lgl()}, \text{layout\_with\_mds()}, \text{layout\_with\_sugiyama()}, \text{layout\_}, \text{merge\_coords()}, \text{norm\_coords()}, \text{normalize()}.

Examples

\[
g <- \text{make\_ring(10)} + \text{make\_ring(10)}
g \\%\% 
\text{add\_layout\_(in\_circle()), component\_wise()} \\%\% 
\text{plot()}
\]

compose

Compose two graphs as binary relations

Description

Relational composition of two graph.

Usage

\[
\text{compose(g1, g2, byname = "auto")}
\]

Arguments

\begin{itemize}
  \item \text{g1} The first input graph.
  \item \text{g2} The second input graph.
  \item \text{byname} A logical scalar, or the character scalar auto. Whether to perform the operation based on symbolic vertex names. If it is auto, that means TRUE if both graphs are named and FALSE otherwise. A warning is generated if auto and one graph, but not both graphs are named.
\end{itemize}
Details

compose creates the relational composition of two graphs. The new graph will contain an (a,b) edge only if there is a vertex c, such that edge (a,c) is included in the first graph and (c,b) is included in the second graph. The corresponding operator is `%c%`.

The function gives an error if one of the input graphs is directed and the other is undirected.

If the `byname` argument is `TRUE` (or `auto` and the graphs are all named), then the operation is performed based on symbolic vertex names. Otherwise numeric vertex ids are used.

compose keeps the attributes of both graphs. All graph, vertex and edge attributes are copied to the result. If an attribute is present in multiple graphs and would result a name clash, then this attribute is renamed by adding suffixes: `_1`, `_2`, etc.

The `name` vertex attribute is treated specially if the operation is performed based on symbolic vertex names. In this case `name` must be present in both graphs, and it is not renamed in the result graph.

Note that an edge in the result graph corresponds to two edges in the input, one in the first graph, one in the second. This mapping is not injective and several edges in the result might correspond to the same edge in the first (and/or the second) graph. The edge attributes in the result graph are updated accordingly.

Also note that the function may generate multigraphs, if there are more than one way to find edges (a,b) in g1 and (b,c) in g2 for an edge (a,c) in the result. See `simplify` if you want to get rid of the multiple edges.

The function may create loop edges, if edges (a,b) and (b,a) are present in g1 and g2, respectively, then (a,a) is included in the result. See `simplify` if you want to get rid of the self-loops.

Value

A new graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

```r
g1 <- make_ring(10)
g2 <- make_star(10, mode="undirected")
gc <- compose(g1, g2)
print_all(gc)
print_all(simplify(gc))
```

Description

These functions find the vertices not farther than a given limit from another fixed vertex, these are called the neighborhood of the vertex.
Usage

```r
connect(graph, order, mode = c("all", "out", "in", "total"))
```

```r
ego_size(
  graph,
  order = 1,
  nodes = V(graph),
  mode = c("all", "out", "in"),
  mindist = 0
)
```

```r
ego(
  graph,
  order = 1,
  nodes = V(graph),
  mode = c("all", "out", "in"),
  mindist = 0
)
```

```r
make_ego_graph(
  graph,
  order = 1,
  nodes = V(graph),
  mode = c("all", "out", "in"),
  mindist = 0
)
```

Arguments

- **graph**: The input graph.
- **order**: Integer giving the order of the neighborhood.
- **mode**: Character constant, it specifies how to use the direction of the edges if a directed graph is analyzed. For 'out' only the outgoing edges are followed, so all vertices reachable from the source vertex in at most `order` steps are counted. For 'in' all vertices from which the source vertex is reachable in at most `order` steps are counted. ‘all’ ignores the direction of the edges. This argument is ignored for undirected graphs.
- **nodes**: The vertices for which the calculation is performed.
- **mindist**: The minimum distance to include the vertex in the result.

Details

The neighborhood of a given order `o` of a vertex `v` includes all vertices which are closer to `v` than the order. I.e. order 0 is always `v` itself, order 1 is `v` plus its immediate neighbors, order 2 is order 1 plus the immediate neighbors of the vertices in order 1, etc.

- **ego_size** calculates the size of the neighborhoods for the given vertices with the given order.
- **ego** calculates the neighborhoods of the given vertices with the given order parameter.
- **make_ego_graph** is creates (sub)graphs from all neighborhoods of the given vertices with the given order parameter. This function preserves the vertex, edge and graph attributes.
- **connect** creates a new graph by connecting each vertex to all other vertices in its neighborhood.
consensus_tree

Value

- `ego_size` returns with an integer vector.
- `ego` returns a list of `igraph.vs` or a list of numeric vectors depending on the value of `igraph_opt("return.vs.es")`, see details for performance characteristics.
- `make_ego_graph` returns a list of graphs.
- `connect` returns with a new graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>, the first version was done by Vincent Matossian

Examples

```r
# Create a consensus tree from several HRG models

# Create a ring graph

g <- make_ring(10)

# Calculate ego sizes for orders 0, 1, 2

ego_size(g, order = 0, 1:3)
ego_size(g, order = 1, 1:3)
ego_size(g, order = 2, 1:3)

ego(g, order = 0, 1:3)
ego(g, order = 1, 1:3)
ego(g, order = 2, 1:3)

# Preserving attributes

V(g)$name <- c("a", "b", "c", "d", "e", "f", "g", "h", "i", "j")
make_ego_graph(g, order = 2, 1:3)

# Connecting to the neighborhood

g <- make_ring(10)
g <- connect(g, 2)
```

Description

`consensus_tree` creates a consensus tree from several fitted hierarchical random graph models, using phylogeny methods. If the `hrg` argument is given and `start` is set to TRUE, then it starts sampling from the given HRG. Otherwise it optimizes the HRG log-likelihood first, and then samples starting from the optimum.

Usage

```r
consensus_tree(graph, hrg = NULL, start = FALSE, num.samples = 10000)
```

Arguments

- **graph**
  The graph the models were fitted to.

- **hrg**
  A hierarchical random graph model, in the form of an `igraphHRG` object. `consensus_tree` allows this to be `NULL` as well, then a HRG is fitted to the graph first, from a random starting point.
Logical, whether to start the fitting/sampling from the supplied igraphHRG object, or from a random starting point.

Number of samples to use for consensus generation or missing edge prediction.

consensus_tree returns a list of two objects. The first is an igraphHRGConsensus object, the second is an igraphHRG object. The igraphHRGConsensus object has the following members:

- parents: For each vertex, the id of its parent vertex is stored, or zero, if the vertex is the root vertex in the tree. The first n vertex ids (from 0) refer to the original vertices of the graph, the other ids refer to vertex groups.

- weights: Numeric vector, counts the number of times a given tree split occurred in the generated network samples, for each internal vertices. The order is the same as in the parents vector.

---

**See Also**

- Other hierarchical random graph functions: `fit_hrg()`, `hrg-methods`, `hrg()`, `predict_edges()`, `print.igraphHRGConsensus()`, `print.igraphHRG()`, `sample_hrg()`

**console**

*The igraph console*

---

**Description**

The igraph console is a GUI windows that shows what the currently running igraph function is doing.

**Usage**

```r
console()
```

**Details**

The console can be started by calling the console function. Then it stays open, until the user closes it.

Another way to start it to set the verbose igraph option to "tkconsole" via `igraph_options`. Then the console (re)opens each time an igraph function supporting it starts; to close it, set the verbose option to another value.

The console is written in Tcl/Tk and required the tcltk package.

**Value**

NULL, invisibly.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**See Also**

`igraph_options` and the verbose option.
**Description**

Given a graph, `constraint` calculates Burt’s constraint for each vertex.

**Usage**

```r
constraint(graph, nodes = V(graph), weights = NULL)
```

**Arguments**

- `graph`: A graph object, the input graph.
- `nodes`: The vertices for which the constraint will be calculated. Defaults to all vertices.
- `weights`: The weights of the edges. If this is `NULL` and there is a `weight` edge attribute this is used. If there is no such edge attribute all edges will have the same weight.

**Details**

Burt’s constraint is higher if ego has less, or mutually stronger related (i.e. more redundant) contacts. Burt’s measure of constraint, \( C_i \), of vertex \( i \)’s ego network \( V_i \), is defined for directed and valued graphs,

\[
C_i = \sum_{j \in V_i \setminus \{i\}} (p_{ij} + \sum_{q \in V_i \setminus \{i,j\}} p_{iq} p_{qj})^2
\]

for a graph of order (ie. number of vertices) \( N \), where proportional tie strengths are defined as

\[
p_{ij} = \frac{a_{ij} + a_{ji}}{\sum_{k \in V_i \setminus \{i\}} (a_{ik} + a_{ki})},
\]

\( a_{i,j} \) are elements of \( A \) and the latter being the graph adjacency matrix. For isolated vertices, constraint is undefined.

**Value**

A numeric vector of constraint scores

**Author(s)**

Jeroen Bruggeman (https://sites.google.com/site/jebrug/jeroen-bruggeman-social-science) and Gabor Csardi <csardi.gabor@gmail.com>

**References**


**Examples**

```r
g <- sample_gnp(20, 5/20)
constraint(g)
```
contract  

**Contract several vertices into a single one**

**Description**

This function creates a new graph, by merging several vertices into one. The vertices in the new graph correspond to sets of vertices in the input graph.

**Usage**

```
contract(graph, mapping, vertex.attr.comb = igraph_opt("vertex.attr.comb"))
```

**Arguments**

- `graph`: The input graph, it can be directed or undirected.
- `mapping`: A numeric vector that specifies the mapping. Its elements correspond to the vertices, and for each element the id in the new graph is given.
- `vertex.attr.comb`: Specifies how to combine the vertex attributes in the new graph. Please see `attribute.combination` for details.

**Details**

The attributes of the graph are kept. Graph and edge attributes are unchanged, vertex attributes are combined, according to the `vertex.attr.comb` parameter.

**Value**

A new graph object.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**Examples**

```
g <- make_ring(10)
g$name <- "Ring"
V(g)$name <- letters[1:vcount(g)]
E(g)$weight <- runif(ecount(g))

g2 <- contract(g, rep(1:5, each=2),
               vertex.attr.comb=toString)

## graph and edge attributes are kept, vertex attributes are
## combined using the 'toString' function.
print(g2, g=TRUE, v=TRUE, e=TRUE)
```
convex_hull  

*Convex hull of a set of vertices*

**Description**

Calculate the convex hull of a set of points, i.e. the covering polygon that has the smallest area.

**Usage**

```r
convex_hull(data)
```

**Arguments**

data  
The data points, a numeric matrix with two columns.

**Value**

A named list with components:

resverts  
The indices of the input vertices that constitute the convex hull.

rescoords  
The coordinates of the corners of the convex hull.

**Author(s)**

Tamas Nepusz <ntamas@gmail.com>

**References**


**Examples**

```r
M <- cbind( runif(100), runif(100) )
convex_hull(M)
```

coreness  

*K-core decomposition of graphs*

**Description**

The k-core of graph is a maximal subgraph in which each vertex has at least degree k. The coreness of a vertex is k if it belongs to the k-core but not to the (k+1)-core.

**Usage**

```r
coreness(graph, mode = c("all", "out", "in"))
```
count_isomorphisms

Arguments

graph The input graph, it can be directed or undirected
mode The type of the core in directed graphs. Character constant, possible values: in: in-cores are computed, out: out-cores are computed, all: the corresponding undirected graph is considered. This argument is ignored for undirected graphs.

Details

The k-core of a graph is the maximal subgraph in which every vertex has at least degree k. The cores of a graph form layers: the (k+1)-core is always a subgraph of the k-core.

This function calculates the coreness for each vertex.

Value

Numeric vector of integer numbers giving the coreness of each vertex.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References

Vladimir Batagelj, Matjaz Zaversnik: An O(m) Algorithm for Cores Decomposition of Networks, 2002

See Also
degree

Examples

g <- make_ring(10)
g <- add_edges(g, c(1,2, 2,3, 1,3))
coreness(g) # small core triangle in a ring

---

count_isomorphisms Count the number of isomorphic mappings between two graphs

Description

Count the number of isomorphic mappings between two graphs

Usage

count_isomorphisms(graph1, graph2, method = "vf2", ...)

count_motifs

Description

Graph motifs are small connected subgraphs with a well-defined structure. These functions search a graph for various motifs.

Usage

```
count_motifs(graph, size = 3, cut.prob = rep(0, size))
```
count_subgraph_isomorphisms

Arguments

- graph: Graph object, the input graph.
- size: The size of the motif.
- cut.prob: Numeric vector giving the probabilities that the search graph is cut at a certain level. Its length should be the same as the size of the motif (the size argument). By default no cuts are made.

Details

count_motifs calculates the total number of motifs of a given size in graph.

Value

count_motifs returns a numeric scalar.

See Also

- isomorphism_class

Other graph motifs: motifs(), sample_motifs()

Examples

```r
# Example 1
pattern <- barabasi.game(100)
print(count_subgraph_isomorphisms(pattern, target, method = "lad"))
```

```r
# Example 2
print(count_subgraph_isomorphisms(pattern, target, method = "vf2"))
```

Description

Count the isomorphic mappings between a graph and the subgraphs of another graph

Usage

count_subgraph_isomorphisms(pattern, target, method = c("lad", "vf2"), ...)
Value

Logical scalar, TRUE if the pattern is isomorphic to a (possibly induced) subgraph of target.

‘lad’ method

This is the LAD algorithm by Solnon, see the reference below. It has the following extra arguments:

domains If not NULL, then it specifies matching restrictions. It must be a list of target vertex sets, given as numeric vertex ids or symbolic vertex names. The length of the list must be vcount(pattern) and for each vertex in pattern it gives the allowed matching vertices in target. Defaults to NULL.

induced Logical scalar, whether to search for an induced subgraph. It is FALSE by default.

time.limit The processor time limit for the computation, in seconds. It defaults to Inf, which means no limit.

‘vf2’ method

This method uses the VF2 algorithm by Cordella, Foggia et al., see references below. It supports vertex and edge colors and have the following extra arguments:

vertex.color1, vertex.color2 Optional integer vectors giving the colors of the vertices for colored graph isomorphism. If they are not given, but the graph has a “color” vertex attribute, then it will be used. If you want to ignore these attributes, then supply NULL for both of these arguments. See also examples below.

domain.color1, domain.color2 Optional integer vectors giving the colors of the edges for edge-colored (sub)graph isomorphism. If they are not given, but the graph has a “color” edge attribute, then it will be used. If you want to ignore these attributes, then supply NULL for both of these arguments.

References


See Also

Other graph isomorphism: count_isomorphisms(), graph_from_isomorphism_class(), isomorphic(), isomorphism_class(), isomorphisms(), subgraph_isomorphic(), subgraph_isomorphisms()

count_triangles | Find triangles in graphs

Description

Count how many triangles a vertex is part of, in a graph, or just list the triangles of a graph.
count_triangles

Usage

count_triangles(graph, vids = V(graph))

Arguments

graph  The input graph. It might be directed, but edge directions are ignored.
vids  The vertices to query, all of them by default. This might be a vector of numeric
       ids, or a character vector of symbolic vertex names for named graphs.

Details

triangles lists all triangles of a graph. For efficiency, all triangles are returned in a single vector.
The first three vertices belong to the first triangle, etc.
count_triangles counts how many triangles a vertex is part of.

Value

For triangles a numeric vector of vertex ids, the first three vertices belong to the first triangle
found, etc.
For count_triangles a numeric vector, the number of triangles for all vertices queried.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

transitivity

Examples

## A small graph
kite <- make_graph("Krackhardt_Kite")
plot(kite)
matrix(triangles(kite), nrow=3)

## Adjacent triangles
atri <- count_triangles(kite)
plot(kite, vertex.label=atri)

## Always true
sum(count_triangles(kite)) == length(triangles(kite))

## Should match, local transitivity is the
## number of adjacent triangles divided by the number
## of adjacency triples
transitivity(kite, type="local")
count_triangles(kite) / (degree(kite) * (degree(kite)-1)/2)
curve_multiple

Optimal edge curvature when plotting graphs

Description
If graphs have multiple edges, then drawing them as straight lines does not show them when plotting the graphs; they will be on top of each other. One solution is to bend the edges, with different curvature, so that all of them are visible.

Usage
curve_multiple(graph, start = 0.5)

Arguments
graph The input graph.
start The curvature at the two extreme edges. All edges will have a curvature between -start and start, spaced equally.

Details
curve_multiple calculates the optimal edge.curved vector for plotting a graph with multiple edges, so that all edges are visible.

Value
A numeric vector, its length is the number of edges in the graph.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

See Also
igraph.plotting for all plotting parameters, plot.igraph, tkplot and rglplot for plotting functions.

Examples
g <- graph( c(0,1,0,1,2,1,3,1,3,1,3,2,3,2,3,2,3,0,1)+1 )

curve_multiple(g)

## Not run:
set.seed(42)
plot(g)

## End(Not run)
decompose

Decompose a graph into components

Description

Creates a separate graph for each component of a graph.

Usage

decompose(graph, mode = c("weak", "strong"), max.comps = NA, min.vertices = 0)

Arguments

graph The original graph.
mode Character constant giving the type of the components, either weak for weakly connected components or strong for strongly connected components.
max.comps The maximum number of components to return. The first max.comps components will be returned (which hold at least min.vertices vertices, see the next parameter), the others will be ignored. Supply NA here if you don’t want to limit the number of components.
min.vertices The minimum number of vertices a component should contain in order to place it in the result list. Eg. supply 2 here to ignore isolate vertices.

Value

A list of graph objects.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

is_connected to decide whether a graph is connected, components to calculate the connected components of a graph.

Examples

# the diameter of each component in a random graph
g <- sample_gnp(1000, 1/1000)
components <- decompose(g, min.vertices=2)
sapply(components, diameter)
**degree**

*Degree and degree distribution of the vertices*

**Description**

The degree of a vertex is its most basic structural property, the number of its adjacent edges.

**Usage**

```r
degree(
  graph,
  v = V(graph),
  mode = c("all", "out", "in", "total"),
  loops = TRUE,
  normalized = FALSE
)
```

```r
degree_distribution(graph, cumulative = FALSE, ...)
```

**Arguments**

- `graph` The graph to analyze.
- `v` The ids of vertices of which the degree will be calculated.
- `mode` Character string, “out” for out-degree, “in” for in-degree or “total” for the sum of the two. For undirected graphs this argument is ignored. “all” is a synonym of “total”.
- `loops` Logical; whether the loop edges are also counted.
- `normalized` Logical scalar, whether to normalize the degree. If TRUE then the result is divided by \( n - 1 \), where \( n \) is the number of vertices in the graph.
- `cumulative` Logical; whether the cumulative degree distribution is to be calculated.
- `...` Additional arguments to pass to `degree`, eg. `mode` is useful but also `v` and `loops` make sense.

**Value**

For `degree` a numeric vector of the same length as argument `v`.

For `degree_distribution` a numeric vector of the same length as the maximum degree plus one. The first element is the relative frequency zero degree vertices, the second vertices with degree one, etc.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**Examples**

```r
g <- make_ring(10)
degree(g)
g2 <- sample_gnp(1000, 10/1000)
degree_distribution(g2)
```
delete_edge_attr  
Delete an edge attribute

Description
Delete an edge attribute

Usage
delete_edge_attr(graph, name)

Arguments
- **graph**: The graph
- **name**: The name of the edge attribute to delete.

Value
The graph, with the specified edge attribute removed.

See Also
Other graph attributes: delete_graph_attr(), delete_vertex_attr(), edge_attr<-(), edge_attr_names(), edge_attr(), graph_attr<-(), graph_attr_names(), graph_attr(), igraph-dollar, igraph-vs-attributes, set_edge_attr(), set_graph_attr(), set_vertex_attr(), vertex_attr<-(), vertex_attr_names(), vertex_attr()

Examples
```r
g <- make_ring(10) %>%
  set_edge_attr("name", value = LETTERS[1:10])
edge_attr_names(g)
g2 <- delete_edge_attr(g, "name")
edge_attr_names(g2)
```

delete_edges  
Delete edges from a graph

Description
Delete edges from a graph

Usage
delete_edges(graph, edges)

Arguments
- **graph**: The input graph.
- **edges**: The edges to remove, specified as an edge sequence. Typically this is either a numeric vector containing edge IDs, or a character vector containing the IDs or names of the source and target vertices, separated by |
delete_graph_attr

Value
The graph, with the edges removed.

See Also
Other functions for manipulating graph structure: \+.igraph(), add_edges(), add_vertices(), delete_vertices(), edge(), igraph-minus, path(), vertex()

Examples

```r
# A directed ring graph
g <- make_ring(10) %>%
  delete_edges(seq(1, 9, by = 2))
g

g <- make_ring(10) %>%
  delete_edges("10\|1")
g

g <- make_ring(5)
g <- delete_edges(g, get.edge.ids(g, c(1,5, 4,5)))
g
```

---

delete_graph_attr  Delete a graph attribute

Description
Delete a graph attribute

Usage
delete_graph_attr(graph, name)

Arguments
graph The graph.
name Name of the attribute to delete.

Value
The graph, with the specified attribute removed.

See Also
Other graph attributes: delete_edge_attr(), delete_vertex_attr(), edge_attr<-(), edge_attr_names(),
edge_attr(), graph_attr<-(), graph_attr_names(), graph_attr(), igraph-dollar, igraph-vs-attributes,
set_edge_attr(), set_graph_attr(), set_vertex_attr(), vertex_attr<-(), vertex_attr_names(),
vertex_attr()
Examples

```r
g <- make_ring(10)
graph_attr_names(g)
g2 <- delete_graph_attr(g, "name")
graph_attr_names(g2)
```

### Description

Delete a vertex attribute

### Usage

```r
delete_vertex_attr(graph, name)
```

### Arguments

- **graph**: The graph
- **name**: The name of the vertex attribute to delete.

### Value

The graph, with the specified vertex attribute removed.

### See Also

Other graph attributes: `delete_edge_attr()`, `delete_graph_attr()`, `edge_attr<-()`, `edge_attr_names()`, `edge_attr()`, `graph_attr<-()`, `graph_attr_names()`, `graph_attr()`, `igraph-dollar`, `igraph-vs-attributes`, `set_edge_attr()`, `set_graph_attr()`, `set_vertex_attr()`, `vertex_attr<-()`, `vertex_attr_names()`, `vertex_attr()`

### Examples

```r
g2 <- make_ring(10)
set_vertex_attr("name", value = LETTERS[1:10])
vertex_attr_names(g)
g2 <- delete_vertex_attr(g, "name")
vertex_attr_names(g2)
```
**delete_vertices**  
*Delete vertices from a graph*

**Description**
Delete vertices from a graph

**Usage**
```r
delete_vertices(graph, v)
```

**Arguments**
- `graph`  
The input graph.
- `v`  
The vertices to remove, a vertex sequence.

**Value**
The graph, with the vertices removed.

**See Also**
Other functions for manipulating graph structure: `+.igraph()`, `add_edges()`, `add_vertices()`, `delete_edges()`, `edge()`, `igraph-minus`, `path()`, `vertex()`

**Examples**
```r
g <- make_ring(10) %>%
  set_vertex_attr("name", value = LETTERS[1:10])
g
V(g)

g2 <- delete_vertices(g, c(1,5)) %>%
delete_vertices("B")
g2
V(g2)
```

---

**dfs**  
*Depth-first search*

**Description**
Depth-first search is an algorithm to traverse a graph. It starts from a root vertex and tries to go quickly as far from as possible.
Usage

dfs(
  graph,
  root,
  mode = c("out", "in", "all", "total"),
  unreachable = TRUE,
  order = TRUE,
  order.out = FALSE,
  father = FALSE,
  dist = FALSE,
  in.callback = NULL,
  out.callback = NULL,
  extra = NULL,
  rho = parent.frame(),
  neimode
)

Arguments

graph The input graph.
root The single root vertex to start the search from.
mode For directed graphs specifies the type of edges to follow. ‘out’ follows outgoing, ‘in’ incoming edges. ‘all’ ignores edge directions completely. ‘total’ is a synonym for ‘all’. This argument is ignored for undirected graphs.
unreachable Logical scalar, whether the search should visit the vertices that are unreachable from the given root vertex (or vertices). If TRUE, then additional searches are performed until all vertices are visited.
order Logical scalar, whether to return the DFS ordering of the vertices.
order.out Logical scalar, whether to return the ordering based on leaving the subtree of the vertex.
father Logical scalar, whether to return the father of the vertices.
dist Logical scalar, whether to return the distance from the root of the search tree.
in.callback If not NULL, then it must be callback function. This is called whenever a vertex is visited. See details below.
out.callback If not NULL, then it must be callback function. This is called whenever the subtree of a vertex is completed by the algorithm. See details below.
extra Additional argument to supply to the callback function.
rho The environment in which the callback function is evaluated.
neimode This argument is deprecated from igraph 1.3.0; use mode instead.

Details

The callback functions must have the following arguments:

graph The input graph is passed to the callback function here.
data A named numeric vector, with the following entries: ‘vid’, the vertex that was just visited and ‘dist’, its distance from the root of the search tree.
extra The extra argument.

The callback must return FALSE to continue the search or TRUE to terminate it. See examples below on how to use the callback functions.
**Value**

A named list with the following entries:

- **root**
  Numeric scalar. The root vertex that was used as the starting point of the search.

- **neimode**
  Character scalar. The mode argument of the function call. Note that for undirected graphs this is always ‘all’, irrespectively of the supplied value.

- **order**
  Numeric vector. The vertex ids, in the order in which they were visited by the search.

- **order.out**
  Numeric vector, the vertex ids, in the order of the completion of their subtree.

- **father**
  Numeric vector. The father of each vertex, i.e. the vertex it was discovered from.

- **dist**
  Numeric vector, for each vertex its distance from the root of the search tree.

Note that order, order.out, father, and dist might be NULL if their corresponding argument is FALSE, i.e. if their calculation is not requested.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**See Also**

- bfs for breadth-first search.

**Examples**

```r
## A graph with two separate trees
dfs(make_tree(10) %du% make_tree(10), root=1, "out",
     TRUE, TRUE, TRUE, TRUE)

## How to use a callback
f.in <- function(graph, data, extra) {
  cat("in:", paste(collapse="", data), "\n")
  FALSE
}
f.out <- function(graph, data, extra) {
  cat("out:", paste(collapse="", data), "\n")
  FALSE
}
tmp <- dfs(make_tree(10), root=1, "out",
            in.callback=f.in, out.callback=f.out)

## Terminate after the first component, using a callback
f.out <- function(graph, data, extra) {
  data["vid"] == 1
}
tmp <- dfs(make_tree(10) %du% make_tree(10), root=1,
            out.callback=f.out)
```

---

**dfs**

Value

A named list with the following entries:

- **root**
  Numeric scalar. The root vertex that was used as the starting point of the search.

- **neimode**
  Character scalar. The mode argument of the function call. Note that for undirected graphs this is always ‘all’, irrespectively of the supplied value.

- **order**
  Numeric vector. The vertex ids, in the order in which they were visited by the search.

- **order.out**
  Numeric vector, the vertex ids, in the order of the completion of their subtree.

- **father**
  Numeric vector. The father of each vertex, i.e. the vertex it was discovered from.

- **dist**
  Numeric vector, for each vertex its distance from the root of the search tree.

Note that order, order.out, father, and dist might be NULL if their corresponding argument is FALSE, i.e. if their calculation is not requested.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

- bfs for breadth-first search.

Examples

```r
## A graph with two separate trees
dfs(make_tree(10) %du% make_tree(10), root=1, "out",
     TRUE, TRUE, TRUE, TRUE)

## How to use a callback
f.in <- function(graph, data, extra) {
  cat("in:", paste(collapse="", data), "\n")
  FALSE
}
f.out <- function(graph, data, extra) {
  cat("out:", paste(collapse="", data), "\n")
  FALSE
}
tmp <- dfs(make_tree(10), root=1, "out",
            in.callback=f.in, out.callback=f.out)

## Terminate after the first component, using a callback
f.out <- function(graph, data, extra) {
  data["vid"] == 1
}
tmp <- dfs(make_tree(10) %du% make_tree(10), root=1,
            out.callback=f.out)
```
**Description**

The diameter of a graph is the length of the longest geodesic.

**Usage**

\[
diameter(graph, directed = TRUE, unconnected = TRUE, weights = NULL)
\]

**Arguments**

- **graph**: The graph to analyze.
- **directed**: Logical, whether directed or undirected paths are to be considered. This is ignored for undirected graphs.
- **unconnected**: Logical, what to do if the graph is unconnected. If FALSE, the function will return a number that is one larger the largest possible diameter, which is always the number of vertices. If TRUE, the diameters of the connected components will be calculated and the largest one will be returned.
- **weights**: Optional positive weight vector for calculating weighted distances. If the graph has a weight edge attribute, then this is used by default.

**Details**

The diameter is calculated by using a breadth-first search like method.

- **get_diameter**: returns a path with the actual diameter. If there are many shortest paths of the length of the diameter, then it returns the first one found.
- **farthest_vertices**: returns two vertex ids, the vertices which are connected by the diameter path.

**Value**

A numeric constant for `diameter`, a numeric vector for `get_diameter`. `farthest_vertices` returns a list with two entries:

- **vertices**: The two vertices that are the farthest.
- **distance**: Their distance.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**See Also**

- `distances`
**Examples**

```r
g <- make_ring(10)
g2 <- delete_edges(g, c(1,2,1,10))
diameter(g2, unconnected=TRUE)
diameter(g2, unconnected=FALSE)
```

```r
## Weighted diameter
set.seed(1)
g <- make_ring(10)
E(g)$weight <- sample(seq_len(ecount(g)))
diameter(g)
get_diameter(g)
diameter(g, weights=NA)
get_diameter(g, weights=NA)
```

---

### difference

**Difference of two sets**

**Description**

This is an S3 generic function. See `methods("difference")` for the actual implementations for various S3 classes. Initially it is implemented for igraph graphs (difference of edges in two graphs), and igraph vertex and edge sequences. See `difference.igraph`, and `difference.igraph.vs`.

**Usage**

`difference(...)`

**Arguments**

... Arguments, their number and interpretation depends on the function that implements `difference`.

**Value**

Depends on the function that implements this method.

---

### difference.igraph

**Difference of graphs**

**Description**

The difference of two graphs are created.

**Usage**

```r
## S3 method for class 'igraph'
difference(big, small, byname = "auto", ...)
```
difference.igraph

Arguments

big The left hand side argument of the minus operator. A directed or undirected graph.
small The right hand side argument of the minus operator. A directed or undirected graph.
bbyname A logical scalar, or the character scalar auto. Whether to perform the operation based on symbolic vertex names. If it is auto, that means TRUE if both graphs are named and FALSE otherwise. A warning is generated if auto and one graph, but not both graphs are named.

... Ignored, included for S3 compatibility.

Details
difference creates the difference of two graphs. Only edges present in the first graph but not in the second will be included in the new graph. The corresponding operator is %m%.

If the byname argument is TRUE (or auto and the graphs are all named), then the operation is performed based on symbolic vertex names. Otherwise numeric vertex ids are used.

difference keeps all attributes (graph, vertex and edge) of the first graph.

Note that big and small must both be directed or both be undirected, otherwise an error message is given.

Value

A new graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

## Create a wheel graph
wheel <- union(make_ring(10),
               make_star(11, center=11, mode="undirected"))
V(wheel)$name <- letters[seq_len(vcount(wheel))]

## Subtract a star graph from it
sstar <- make_star(6, center=6, mode="undirected")
V(sstar)$name <- letters[c(1,3,5,7,9,11)]
G <- wheel %m% sstar
print_all(G)
plot(G, layout=layout_nicely(wheel))
difference.igraph.es  Difference of edge sequences

Description

Difference of edge sequences

Usage

## S3 method for class 'igraph.es'
difference(big, small, ...)

Arguments

big

The ‘big’ edge sequence.

small

The ‘small’ edge sequence.

...  Ignored, included for S3 signature compatibility.

Details

They must belong to the same graph. Note that this function has ‘set’ semantics and the multiplicity of edges is lost in the result.

Value

An edge sequence that contains only edges that are part of big, but not part of small.

See Also

Other vertex and edge sequence operations: c.igraph.es(), c.igraph.vs(), difference.igraph.es(), igraph-es-indexing2, igraph-es-indexing, igraph-vs-indexing2, igraph-vs-indexing, intersection.igraph.es(), intersection.igraph.vs(), rev.igraph.es(), rev.igraph.vs(), union.igraph.es(), union.igraph.vs(), unique.igraph.es(), unique.igraph.vs()

Examples

```r
g <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
difference(V(g), V(g)[6:10])
```

difference.igraph.vs  Difference of vertex sequences

Description

Difference of vertex sequences

Usage

## S3 method for class 'igraph.vs'
difference(big, small, ...)

### Examples

```r
g <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
difference(V(g), V(g)[6:10])
```
Arguments

big The ‘big’ vertex sequence.
small The ‘small’ vertex sequence.
... Ignored, included for S3 signature compatibility.

Details

They must belong to the same graph. Note that this function has ‘set’ semantics and the multiplicity of vertices is lost in the result.

Value

A vertex sequence that contains only vertices that are part of big, but not part of small.

See Also


Examples

g <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
difference(V(g), V(g)[6:10])

dim_select

Dimensionality selection for singular values using profile likelihood.

Description

Select the number of significant singular values, by finding the ‘elbow’ of the scree plot, in a principled way.

Usage

dim_select(sv)

Arguments

sv A numeric vector, the ordered singular values.

Details

The input of the function is a numeric vector which contains the measure of ‘importance’ for each dimension.

For spectral embedding, these are the singular values of the adjacency matrix. The singular values are assumed to be generated from a Gaussian mixture distribution with two components that have different means and same variance. The dimensionality $d$ is chosen to maximize the likelihood when the $d$ largest singular values are assigned to one component of the mixture and the rest of the singular values assigned to the other component.
This function can also be used for the general separation problem, where we assume that the left and the right of the vector are coming from two Normal distributions, with different means, and we want to know their border. See examples below.

Value

A numeric scalar, the estimate of $d$.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

embed_adjacency_matrix

Examples

# Generate the two groups of singular values with
# Gaussian mixture of two components that have different means
sing.vals <- c( rnorm (10, mean=1, sd=1), rnorm(10, mean=3, sd=1) )
dim.chosen <- dim_select(sing.vals)
dim.chosen

# Sample random vectors with multivariate normal distribution
# and normalize to unit length
lpvs <- matrix(rnorm(200), 10, 20)
lpvs <- apply(lpvs, 2, function(x) { (abs(x) / sqrt(sum(x^2))) })
RDP.graph <- sample_dot_product(lpvs)
dim_select( embed_adjacency_matrix(RDP.graph, 10)$D )

# Sample random vectors with the Dirichlet distribution
lpvs.dir <- sample_dirichlet(n=20, rep(1, 10))
RDP.graph.2 <- sample_dot_product(lpvs.dir)
dim_select( embed_adjacency_matrix(RDP.graph.2, 10)$D )

# Sample random vectors from hypersphere with radius 1.
lpvs.sph <- sample_sphere_surface(dim=10, n=20, radius=1)
RDP.graph.3 <- sample_dot_product(lpvs.sph)
dim_select( embed_adjacency_matrix(RDP.graph.3, 10)$D )
Usage

\texttt{disjoint\_union(...)}

\texttt{x \%du\% y}

Arguments

... \hspace{1cm} \text{Graph objects or lists of graph objects.}
\texttt{x, y} \hspace{1cm} \text{Graph objects.}

Details

disjoint\_union creates a union of two or more disjoint graphs. Thus first the vertices in the second, third, etc. graphs are relabeled to have completely disjoint graphs. Then a simple union is created. This function can also be used via the \%du\% operator.

graph\_disjoint\_union handles graph, vertex and edge attributes. In particular, it merges vertex and edge attributes using the basic c() function. For graphs that lack some vertex/edge attribute, the corresponding values in the new graph are set to NA. Graph attributes are simply copied to the result. If this would result a name clash, then they are renamed by adding suffixes: _1, _2, etc.

Note that if both graphs have vertex names (ie. a name vertex attribute), then the concatenated vertex names might be non-unique in the result. A warning is given if this happens.

An error is generated if some input graphs are directed and others are undirected.

Value

A new graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

### A star and a ring
\texttt{g1 <- make\_star(10, mode=\"undirected\")}
\texttt{V(g1)$name <- letters[1:10]}
\texttt{g2 <- make\_ring(10)}
\texttt{V(g2)$name <- letters[11:20]}
\texttt{print\_all(g1 \%du\% g2)}

---

distance\_table \hspace{1cm} \textit{Shortest (directed or undirected) paths between vertices}

Description

distances calculates the length of all the shortest paths from or to the vertices in the network. shortest\_paths calculates one shortest path (the path itself, and not just its length) from or to the given vertex.
Usage

distance_table(graph, directed = TRUE)

mean_distance(
  graph,
  weights = NULL,
  directed = TRUE,
  unconnected = TRUE,
  details = FALSE
)

distances(
  graph,
  v = V(graph),
  to = V(graph),
  mode = c("all", "out", "in"),
  weights = NULL,
  algorithm = c("automatic", "unweighted", "dijkstra", "bellman-ford", "johnson")
)

shortest_paths(
  graph,
  from,
  to = V(graph),
  mode = c("out", "all", "in"),
  weights = NULL,
  output = c("vpath", "epath", "both"),
  predecessors = FALSE,
  inbound.edges = FALSE,
  algorithm = c("automatic", "unweighted", "dijkstra", "bellman-ford")
)

all_shortest_paths(
  graph,
  from,
  to = V(graph),
  mode = c("out", "all", "in"),
  weights = NULL
)

Arguments

graph The graph to work on.
directed Whether to consider directed paths in directed graphs, this argument is ignored for undirected graphs.
weights Possibly a numeric vector giving edge weights. If this is NULL and the graph has a weight edge attribute, then the attribute is used. If this is NA then no weights are used (even if the graph has a weight attribute).
unconnected What to do if the graph is unconnected (not strongly connected if directed paths are considered). If TRUE, only the lengths of the existing paths are considered and averaged; if FALSE, the length of the missing paths are considered as having infinite length, making the mean distance infinite as well.
Whether to provide additional details in the result. Functions accepting this argument (like `mean_distance`) return additional information like the number of disconnected vertex pairs in the result when this parameter is set to `TRUE`.

**v**
Numeric vector, the vertices from which the shortest paths will be calculated.

**to**
Numeric vector, the vertices to which the shortest paths will be calculated. By default it includes all vertices. Note that for distances every vertex must be included here at most once. (This is not required for `shortest_paths`.

**mode**
Character constant, gives whether the shortest paths to or from the given vertices should be calculated for directed graphs. If `out` then the shortest paths from the vertex, if `in` then `to` it will be considered. If `all`, the default, then the corresponding undirected graph will be used, i.e. not directed paths are searched. This argument is ignored for undirected graphs.

**algorithm**
Which algorithm to use for the calculation. By default igraph tries to select the fastest suitable algorithm. If there are no weights, then an unweighted breadth-first search is used, otherwise if all weights are positive, then Dijkstra’s algorithm is used. If there are negative weights and we do the calculation for more than 100 sources, then Johnson’s algorithm is used. Otherwise the Bellman-Ford algorithm is used. You can override igraph’s choice by explicitly giving this parameter. Note that the igraph C core might still override your choice in obvious cases, i.e. if there are no edge weights, then the unweighted algorithm will be used, regardless of this argument.

**from**
Numeric constant, the vertex from or to the shortest paths will be calculated. Note that right now this is not a vector of vertex ids, but only a single vertex.

**output**
Character scalar, defines how to report the shortest paths. “vpath” means that the vertices along the paths are reported, this form was used prior to igraph version 0.6. “epath” means that the edges along the paths are reported. “both” means that both forms are returned, in a named list with components “vpath” and “epath”.

**predecessors**
Logical scalar, whether to return the predecessor vertex for each vertex. The predecessor of vertex i in the tree is the vertex from which vertex i was reached. The predecessor of the start vertex (in the `from` argument) is itself by definition. If the predecessor is zero, it means that the given vertex was not reached from the source during the search. Note that the search terminates if all the vertices in `to` are reached.

**inbound.edges**
Logical scalar, whether to return the inbound edge for each vertex. The inbound edge of vertex i in the tree is the edge via which vertex i was reached. The start vertex and vertices that were not reached during the search will have zero in the corresponding entry of the vector. Note that the search terminates if all the vertices in `to` are reached.

**Details**

The shortest path, or geodesic between two pair of vertices is a path with the minimal number of vertices. The functions documented in this manual page all calculate shortest paths between vertex pairs.

`distances` calculates the lengths of pairwise shortest paths from a set of vertices (`from`) to another set of vertices (`to`). It uses different algorithms, depending on the `algorithm` argument and the weight edge attribute of the graph. The implemented algorithms are breadth-first search (`'unweighted'`), this only works for unweighted graphs; the Dijkstra algorithm (`'dijkstra'`), this works for graphs with non-negative edge weights; the Bellman-Ford algorithm (`'bellman-ford'`),
and Johnson's algorithm ("johnson"). The latter two algorithms work with arbitrary edge weights, but (naturally) only for graphs that don't have a negative cycle.

igraph can choose automatically between algorithms, and chooses the most efficient one that is appropriate for the supplied weights (if any). For automatic algorithm selection, supply 'automatic' as the algorithm argument. (This is also the default.)

shortest_paths calculates a single shortest path (i.e. the path itself, not just its length) between the source vertex given in from, to the target vertices given in to. shortest_paths uses breadth-first search for unweighted graphs and Dijkstra's algorithm for weighted graphs. The latter only works if the edge weights are non-negative.

all_shortest_paths calculates all shortest paths between pairs of vertices. More precisely, between the from vertex to the vertices given in to. It uses a breadth-first search for unweighted graphs and Dijkstra's algorithm for weighted ones. The latter only supports non-negative edge weights.

mean_distance calculates the average path length in a graph, by calculating the shortest paths between all pairs of vertices (both ways for directed graphs). It uses a breadth-first search for unweighted graphs and Dijkstra's algorithm for weighted ones. The latter only supports non-negative edge weights.

distance_table calculates a histogram, by calculating the shortest path length between each pair of vertices. For directed graphs both directions are considered, so every pair of vertices appears twice in the histogram.

**Value**

For distances a numeric matrix with length(to) columns and length(v) rows. The shortest path length from a vertex to itself is always zero. For unreachable vertices Inf is included.

For shortest_paths a named list with four entries is returned:

- **vpath**
  This itself is a list, of length length(to); list element i contains the vertex ids on the path from vertex from to vertex to[i] (or the other way for directed graphs depending on the mode argument). The vector also contains from and i as the first and last elements. If from is the same as i then it is only included once. If there is no path between two vertices then a numeric vector of length zero is returned as the list element. If this output is not requested in the output argument, then it will be NULL.

- **epath**
  This is a list similar to vpath, but the vectors of the list contain the edge ids along the shortest paths, instead of the vertex ids. This entry is set to NULL if it is not requested in the output argument.

- **predecessors**
  Numeric vector, the predecessor of each vertex in the to argument, or NULL if it was not requested.

- **inbound_edges**
  Numeric vector, the inbound edge for each vertex, or NULL, if it was not requested.

For all_shortest_paths a list is returned, each list element contains a shortest path from from to a vertex in to. The shortest paths to the same vertex are collected into consecutive elements of the list.

For mean_distance a single number is returned if details=FALSE, or a named list with two entries: res is the mean distance as a numeric scalar and unconnected is the number of unconnected vertex pairs, also as a numeric scalar.

distance_table returns a named list with two entries: res is a numeric vector, the histogram of distances, unconnected is a numeric scalar, the number of pairs for which the first vertex is not reachable from the second. The sum of the two entries is always \( n(n - 1) \) for directed graphs and \( n(n - 1)/2 \) for undirected graphs.
Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

References

Examples
```r
g <- make_ring(10)
distances(g)
shortest_paths(g, 5)
all_shortest_paths(g, 1, 6:8)
mean_distance(g)
## Weighted shortest paths
el <- matrix(ncol=3, byrow=TRUE,
c(1,2,0, 1,3,2, 1,4,1, 2,3,0, 2,5,5, 2,6,2, 3,2,1, 3,4,1, 3,7,1, 4,3,0, 4,7,2, 5,6,2, 5,8,8, 6,3,2, 6,7,1, 6,9,1, 6,10,3, 8,6,1, 8,9,1, 9,10,4) )
g2 <- add_edges(make_empty_graph(10), t(el[,1:2]), weight=el[,3])
distances(g2, mode="out")
```

---

diverging_pal

* Diverging palette

Description
This is the ‘PuOr’ palette from [https://colorbrewer2.org/](https://colorbrewer2.org/). It has at most eleven colors.

Usage
diverging_pal(n)

Arguments

- **n**
  The number of colors in the palette. The maximum is eleven currently.

Details
This is similar to `sequential_pal`, but it also puts emphasis on the mid-range values, plus the two extreme ends. Use this palette, if you have such a quantity to mark with vertex colors.

Value
A character vector of RGB color codes.

See Also
Other palettes: `categorical_pal()`, `r_pal()`, `sequential_pal()`
Examples

```r
## Not run:
library(igraphdata)
data(foodwebs)
fw <- foodwebs[[1]] %>%
  induced_subgraph(V(.)[ECO == 1]) %>%
  add_layout_(with_fr()) %>%
  set_vertex_attr("label", value = seq_len(gorder(.))) %>%
  set_vertex_attr("size", value = 10) %>%
  set_edge_attr("arrow.size", value = 0.3)
V(fw)$color <- scales::dscale(V(fw)$Biomass %>% cut(10), diverging_pal)
plot(fw)
data(karate)
karate <- karate %>%
  add_layout_(with_kk()) %>%
  set_vertex_attr("size", value = 10)
V(karate)$color <- scales::dscale(degree(karate) %>% cut(5), diverging_pal)
plot(karate)
## End(Not run)
```

### diversity

**Graph diversity**

Calculates a measure of diversity for all vertices.

#### Usage

```r
diversity(graph, weights = NULL, vids = V(graph))
```

#### Arguments

- **graph**: The input graph. Edge directions are ignored.
- **weights**: NULL, or the vector of edge weights to use for the computation. If NULL, then the ‘weight’ attribute is used. Note that this measure is not defined for unweighted graphs.
- **vids**: The vertex ids for which to calculate the measure.

#### Details

The diversity of a vertex is defined as the (scaled) Shannon entropy of the weights of its incident edges:

\[
D(i) = \frac{H(i)}{\log k_i}
\]

and

\[
H(i) = -\sum_{j=1}^{k_i} p_{ij} \log p_{ij},
\]
where
\[ p_{ij} = \frac{w_{ij}}{\sum_{l=1}^{k_i} V_{il}}, \]
and \( k_i \) is the (total) degree of vertex \( i \), \( w_{ij} \) is the weight of the edge(s) between vertices \( i \) and \( j \).
For vertices with degree less than two the function returns NaN.

**Value**

A numeric vector, its length is the number of vertices.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**References**


**Examples**

```r
G1 <- sample_gnp(20, 2/20)
G2 <- sample_gnp(20, 2/20)
G3 <- sample_gnp(20, 5/20)
E(G1)$weight <- 1
E(G2)$weight <- runif(ecount(G2))
E(G3)$weight <- runif(ecount(G3))
diversity(G1)
diversity(G2)
diversity(G3)
```

---

**dominator_tree**

**Dominator tree**

**Description**

Dominator tree of a directed graph.

**Usage**

```r
dominator_tree(graph, root, mode = c("out", "in", "all", "total"))
```

**Arguments**

- `graph`: A directed graph. If it is not a flowgraph, and it contains some vertices not reachable from the root vertex, then these vertices will be collected and returned as part of the result.
- `root`: The id of the root (or source) vertex, this will be the root of the tree.
- `mode`: Constant, must be ‘in’ or ‘out’. If it is ‘in’, then all directions are considered as opposite to the original one in the input graph.
Details

A flowgraph is a directed graph with a distinguished start (or root) vertex \( r \), such that for any vertex \( v \), there is a path from \( r \) to \( v \). A vertex \( v \) dominates another vertex \( w \) (not equal to \( v \)), if every path from \( r \) to \( w \) contains \( v \). Vertex \( v \) is the immediate dominator or \( w \), \( v = \text{idom}(w) \), if \( v \) dominates \( w \) and every other dominator of \( w \) dominates \( v \). The edges \((\text{idom}(w), w) | w \neq r \) form a directed tree, rooted at \( r \), called the dominator tree of the graph. Vertex \( v \) dominates vertex \( w \) if and only if \( v \) is an ancestor of \( w \) in the dominator tree.

This function implements the Lengauer-Tarjan algorithm to construct the dominator tree of a directed graph. For details see the reference below.

Value

A list with components:
- **dom**: A numeric vector giving the immediate dominators for each vertex. For vertices that are unreachable from the root, it contains \( \text{NaN} \). For the root vertex itself it contains minus one.
- **domtree**: A graph object, the dominator tree. Its vertex ids are the as the vertex ids of the input graph. Isolate vertices are the ones that are unreachable from the root.
- **leftout**: A numeric vector containing the vertex ids that are unreachable from the root.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


Examples

```r
## The example from the paper

g <- graph_from_literal(R-+A:B:C, A-+D, B-+A:D:E, C-+F:G, D-+L, 
                        E-+H, F-+I, G-+I:J, H-+E:K, I-+K, J-+I, 
                        K-+I:R, L-+H)
dtree <- dominator_tree(g, root="R")
layout <- layout_as_tree(dtree$domtree, root="R")
layout[,2] <- -layout[,2]
plot(dtree$domtree, layout=layout, vertex.label=V(dtree$domtree)$name)
```

---

**Description**

The `.data` and `.env` pronouns make it explicit where to look up attribute names when indexing \( V(g) \) or \( E(g) \), i.e. the vertex or edge sequence of a graph. These pronouns are inspired by `.data` and `.env` in `rlang` - thanks to Michał Bojanowski for bringing these to our attention.

The rules are simple:
• \texttt{.data} retrieves attributes from the graph whose vertex or edge sequence is being evaluated.
• \texttt{.env} retrieves variables from the calling environment.

Note that \texttt{.data} and \texttt{.env} are injected dynamically into the environment where the indexing expressions are evaluated; you cannot get access to these objects outside the context of an indexing expression. To avoid warnings printed by \texttt{R CMD check} when code containing \texttt{.data} and \texttt{.env} is checked, you can import \texttt{.data} and \texttt{.env} from \texttt{igraph} if needed. Alternatively, you can declare them explicitly with \texttt{utils::globalVariables()} to silence the warnings.

---

**Description**

The common bits of the three plotting functions \texttt{plot.igraph}, \texttt{tkplot} and \texttt{rglplot} are discussed in this manual page.

**Details**

There are currently three different functions in the \texttt{igraph} package which can draw graph in various ways:

\texttt{plot.igraph} does simple non-interactive 2D plotting to R devices. Actually it is an implementation of the \texttt{plot} generic function, so you can write \texttt{plot(graph)} instead of \texttt{plot.igraph(graph)}. As it used the standard R devices it supports every output format for which R has an output device. The list is quite impressive: PostScript, PDF files, XFig files, SVG files, JPG, PNG and of course you can plot to the screen as well using the default devices, or the good-looking anti-aliased Cairo device. See \texttt{plot.igraph} for some more information.

\texttt{tkplot} does interactive 2D plotting using the \texttt{tcltk} package. It can only handle graphs of moderate size, a thousand vertices is probably already too many. Some parameters of the plotted graph can be changed interactively after issuing the \texttt{tkplot} command: the position, color and size of the vertices and the color and width of the edges. See \texttt{tkplot} for details.

\texttt{rglplot} is an experimental function to draw graphs in 3D using OpenGL. See \texttt{rglplot} for some more information.

Please also check the examples below.

**How to specify graphical parameters**

There are three ways to give values to the parameters described below, in section `Parameters`. We give these three ways here in the order of their precedence.

The first method is to supply named arguments to the plotting commands: \texttt{plot.igraph}, \texttt{tkplot} or \texttt{rglplot}. Parameters for vertices start with prefix `vertex.', parameters for edges have prefix `edge.', and global parameters have no prefix. Eg. the color of the vertices can be given via argument `vertex.color', whereas `edge.color' sets the color of the edges. `layout' gives the layout of the graphs.

The second way is to assign vertex, edge and graph attributes to the graph. These attributes have no prefix, ie. the color of the vertices is taken from the `color' vertex attribute and the color of the edges from the `color' edge attribute. The layout of the graph is given by the `layout' graph attribute. (Always assuming that the corresponding command argument is not present.) Setting vertex and edge attributes are handy if you want to assign a given `look' to a graph, attributes are saved with
the graph is saved with `save` or in GraphML format with `write_graph`, so the graph will have the same look after loading it again.

If a parameter is not given in the command line, and the corresponding vertex/edge/graph attribute is also missing then the general igraph parameters handled by `igraph_options` are also checked. Vertex parameters have prefix `vertex.`, edge parameters are prefixed with `edge.`, general parameters like `layout` are prefixed with `plot`. These parameters are useful if you want all or most of your graphs to have the same look, vertex size, vertex color, etc. Then you don’t need to set these at every plotting, and you also don’t need to assign vertex/edge attributes to every graph.

If the value of a parameter is not specified by any of the three ways described here, its default valued is used, as given in the source code.

Different parameters can have different type, e.g. vertex colors can be given as a character vector with color names, or as an integer vector with the color numbers from the current palette. Different types are valid for different parameters, this is discussed in detail in the next section. It is however always true that the parameter can always be a function object in which it will be called with the graph as its single argument to get the “proper” value of the parameter. (If the function returns another function object that will not be called again…)

The list of parameters

Vertex parameters first, note that the `vertex.` prefix needs to be added if they are used as an argument or when setting via `igraph_options`. The value of the parameter may be scalar valid for every vertex or a vector with a separate value for each vertex. (Shorter vectors are recycled.)

- **size** The size of the vertex, a numeric scalar or vector, in the latter case each vertex size may differ. This vertex sizes are scaled in order have about the same size of vertices for a given value for all three plotting commands. It does not need to be an integer number.
  - The default value is 15. This is big enough to place short labels on vertices.

- **size2** The “other” size of the vertex, for some vertex shapes. For the various rectangle shapes this gives the height of the vertices, whereas `size` gives the width. It is ignored by shapes for which the size can be specified with a single number.
  - The default is 15.

- **color** The fill color of the vertex. If it is numeric then the current palette is used, see `palette`. If it is a character vector then it may either contain integer values, named colors or RGB specified colors with three or four bytes. All strings starting with `#` are assumed to be RGB color specifications. It is possible to mix named color and RGB colors. Note that `tkplot` ignores the fourth byte (alpha channel) in the RGB color specification.
  - For `plot.igraph` and integer values, the default igraph palette is used (see the ‘palette’ parameter below. Note that this is different from the R palette.
  - If you don’t want (some) vertices to have any color, supply `NA` as the color name.
  - The default value is “SkyBlue2”.

- **frame.color** The color of the frame of the vertices, the same formats are allowed as for the fill color.
  - If you don’t want vertices to have a frame, supply `NA` as the color name.
  - By default it is “black”.

- **frame.width** The width of the frame of the vertices.
  - The default value is 1.

- **shape** The shape of the vertex, currently “circle”, “square”, “csquare”, “rectangle”, “crectangle”, “vrectangle”, “pie” (see `vertex.shape`), ‘sphere’, and “none” are supported, and only by the `plot.igraph` command. “none” does not draw the vertices at all, although vertex label
are plotted (if given). See *shapes* for details about vertex shapes and *vertex.shape.pie* for using pie charts as vertices.

The “sphere” vertex shape plots vertices as 3D ray-traced spheres, in the given color and size. This produces a raster image and it is only supported with some graphics devices. On some devices raster transparency is not supported and the spheres do not have a transparent background. See *dev.capabilities* and the `rasterImage` capability to check that your device is supported.

By default vertices are drawn as circles.

**label**
The vertex labels. They will be converted to character. Specify NA to omit vertex labels.

The default vertex labels are the vertex ids.

**label.family**
The font family to be used for vertex labels. As different plotting commands can use different fonts, they interpret this parameter different ways. The basic notation is, however, understood by both `plot.igraph` and `tkplot`. `rglplot` does not support fonts at all right now, it ignores this parameter completely.

For `plot.igraph` this parameter is simply passed to `text` as argument `family`.

For `tkplot` some conversion is performed. If this parameter is the name of an existing Tk font, then that font is used and the `label.font` and `label.cex` parameters are ignored completely. If it is one of the base families (serif, sans, mono) then Times, Helvetica or Courier fonts are used, there are guaranteed to exist on all systems. For the ‘symbol’ base family we used the symbol font is available, otherwise the first font which has ‘symbol’ in its name. If the parameter is not a name of the base families and it is also not a named Tk font then we pass it to `tkfont.create` and hope the user knows what she is doing. The `label.font` and `label.cex` parameters are also passed to `tkfont.create` in this case.

The default value is ‘serif’.

**label.font**
The font within the font family to use for the vertex labels. It is interpreted the same way as the the *font* graphical parameter: 1 is plain text, 2 is bold face, 3 is italic, 4 is bold and italic and 5 specifies the symbol font.

For `plot.igraph` this parameter is simply passed to `text`.

For `tkplot`, if the `label.family` parameter is not the name of a Tk font then this parameter is used to set whether the newly created font should be italic and/or boldface. Otherwise it is ignored.

For `rglplot` it is ignored.

The default value is 1.

**label.cex**
The font size for vertex labels. It is interpreted as a multiplication factor of some device-dependent base font size.

For `plot.igraph` it is simply passed to `text` as argument `cex`.

For `tkplot` it is multiplied by 12 and then used as the size argument for `tkfont.create`. The base font is thus 12 for tkplot.

For `rglplot` it is ignored.

The default value is 1.

**label.dist**
The distance of the label from the center of the vertex. If it is 0 then the label is centered on the vertex. If it is 1 then the label is displayed beside the vertex.

The default value is 0.

**label.degree**
It defines the position of the vertex labels, relative to the center of the vertices. It is interpreted as an angle in radian, zero means ‘to the right’, and ‘pi’ means to the left, up is –pi/2 and down is pi/2.

The default value is –pi/4.
**Drawing graphs**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>label.color</td>
<td>The color of the labels, see the color vertex parameter discussed earlier for possible values. The default value is black.</td>
</tr>
</tbody>
</table>

Edge parameters require to add the `edge.` prefix when used as arguments or set by `igraph_options`. The edge parameters:

**color** The color of the edges, see the color vertex parameter for the possible values. By default this parameter is darkgrey.

**width** The width of the edges. The default value is 1.

**arrow.size** The size of the arrows. Currently this is a constant, so it is the same for every edge. If a vector is submitted then only the first element is used, i.e. if this is taken from an edge attribute then only the attribute of the first edge is used for all arrows. This will likely change in the future. The default value is 1.

**arrow.width** The width of the arrows. Currently this is a constant, so it is the same for every edge. If a vector is submitted then only the first element is used, i.e. if this is taken from an edge attribute then only the attribute of the first edge is used for all arrows. This will likely change in the future. This argument is currently only used by `plot.igraph`. The default value is 1, which gives the same width as before this option appeared in igraph.

**lty** The line type for the edges. Almost the same format is accepted as for the standard graphics `par`, 0 and “blank” mean no edges, 1 and “solid” are for solid lines, the other possible values are: 2 (“dashed”), 3 (“dotted”), 4 (“dotdash”), 5 (“longdash”), 6 (“twodash”). `tkplot` also accepts standard Tk line type strings, it does not however support “blank” lines, instead of type ‘0’ type ‘1’, i.e. solid lines will be drawn. This argument is ignored for `rglplot`. The default value is type 1, a solid line.

**label** The edge labels. They will be converted to character. Specify NA to omit edge labels. Edge labels are omitted by default.

**label.family** Font family of the edge labels. See the vertex parameter with the same name for the details.

**label.font** The font for the edge labels. See the corresponding vertex parameter discussed earlier for details.

**label.cex** The font size for the edge labels, see the corresponding vertex parameter for details.

**label.color** The color of the edge labels, see the color vertex parameters on how to specify colors.

**label.x** The horizontal coordinates of the edge labels might be given here, explicitly. The NA elements will be replaced by automatically calculated coordinates. If NULL, then all edge horizontal coordinates are calculated automatically. This parameter is only supported by `plot.igraph`.

**label.y** The same as `label.x`, but for vertical coordinates.

**curved** Specifies whether to draw curved edges, or not. This can be a logical or a numeric vector or scalar.

First the vector is replicated to have the same length as the number of edges in the graph. Then it is interpreted for each edge separately. A numeric value specifies the curvature of the edge: zero curvature means straight edges, negative values mean the edge bends clockwise, positive values the opposite. TRUE means curvature 0.5, FALSE means curvature zero.
By default the vector specifying the curvature is calculated via a call to the `curve_multiple` function. This function makes sure that multiple edges are curved and are all visible. This parameter is ignored for loop edges.

The default value is `FALSE`.

This parameter is currently ignored by `rglplot`.

`arrow.mode` This parameter can be used to specify for which edges should arrows be drawn. If this parameter is given by the user (in either of the three ways) then it specifies which edges will have forward, backward arrows, or both, or no arrows at all. As usual, this parameter can be a vector or a scalar value. It can be an integer or character type. If it is integer then 0 means no arrows, 1 means backward arrows, 2 is for forward arrows and 3 for both. If it is a character vector then “<” and “<.” specify backward, “>” and “>.” forward arrows and “<>” and “<->” stands for both arrows. All other values mean no arrows, perhaps you should use “-” or “–” to specify no arrows.

Hint: this parameter can be used as a ‘cheap’ solution for drawing “mixed” graphs: graphs in which some edges are directed some are not. If you want do this, then please create a directed graph, because as of version 0.4 the vertex pairs in the edge lists can be swapped in undirected graphs.

By default, no arrows will be drawn for undirected graphs, and for directed graphs, an arrow will be drawn for each edge, according to its direction. This is not very surprising, it is the expected behavior.

`loop.angle` Gives the angle in radian for plotting loop edges. See the `label.dist` vertex parameter to see how this is interpreted.

The default value is 0.

`loop.angle2` Gives the second angle in radian for plotting loop edges. This is only used in 3D, `loop.angle` is enough in 2D.

The default value is 0.

Other parameters:

`layout` Either a function or a numeric matrix. It specifies how the vertices will be placed on the plot.

If it is a numeric matrix, then the matrix has to have one line for each vertex, specifying its coordinates. The matrix should have at least two columns, for the x and y coordinates, and it can also have third column, this will be the z coordinate for 3D plots and it is ignored for 2D plots.

If a two column matrix is given for the 3D plotting function `rglplot` then the third column is assumed to be 1 for each vertex.

If `layout` is a function, this function will be called with the graph as the single parameter to determine the actual coordinates. The function should return a matrix with two or three columns. For the 2D plots the third column is ignored.

The default value is `layout_nicely`, a smart function that chooses a layouter based on the graph.

`margin` The amount of empty space below, over, at the left and right of the plot, it is a numeric vector of length four. Usually values between 0 and 0.5 are meaningful, but negative values are also possible, that will make the plot zoom in to a part of the graph. If it is shorter than four then it is recycled.

`rglplot` does not support this parameter, as it can zoom in and out the graph in a more flexible way.

Its default value is 0.
palettes The color palette to use for vertex color. The default is `categorical_pal`, which is a color-blind friendly categorical palette. See its manual page for details and other palettes. This parameters is only supported by `plot`, and not by `tkplot` and `rglplot`.

rescale Logical constant, whether to rescale the coordinates to the [-1,1]x[-1,1] interval. This parameter is not implemented for `tkplot`.

Defaults to TRUE, the layout will be rescaled.

asp A numeric constant, it gives the `asp` parameter for `plot`, the aspect ratio. Supply 0 here if you don’t want to give an aspect ratio. It is ignored by `tkplot` and `rglplot`.

Defaults to 1.

frame Boolean, whether to plot a frame around the graph. It is ignored by `tkplot` and `rglplot`.

Defaults to FALSE.

main Overall title for the main plot. The default is empty if the `annotate.plot` igraph option is FALSE, and the graph’s name attribute otherwise. See the same argument of the base `plot` function. Only supported by `plot`.

sub Subtitle of the main plot, the default is empty. Only supported by `plot`.

xlab Title for the x axis, the default is empty if the `annotate.plot` igraph option is FALSE, and the number of vertices and edges, if it is TRUE. Only supported by `plot`.

ylab Title for the y axis, the default is empty. Only supported by `plot`.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

`plot.igraph`, `tkplot`, `rglplot`, `igraph_options`

Examples

```r
## Not run:
# plotting a simple ring graph, all default parameters, except the layout
g <- make_ring(10)
g$layout <- layout_in_circle
plot(g)
tkplot(g)
rglplot(g)

# plotting a random graph, set the parameters in the command arguments
g <- barabasi.game(100)
plot(g, layout=layout_with_fr, vertex.size=4,
     vertex.label.dist=0.5, vertex.color="red", edge.arrow.size=0.5)

# plot a random graph, different color for each component
g <- sample_gnp(100, 1/100)
comps <- components(g)$membership
colbar <- rainbow(max(comps)+1)
V(g)$color <- colbar[comps+1]
plot(g, layout=layout_with_fr, vertex.size=5, vertex.label=NA)

# plot communities in a graph
g <- make_full_graph(5) %du% make_full_graph(5) %du% make_full_graph(5)
g <- add_edges(g, c(1,6, 1,11, 6,11))
```
Com <- cluster_spinglass(g, spins=5)
V(g)$color <- com$membership+1
g <- set_graph_attr(g, "layout", layout_with_kk(g))
plot(g, vertex.label.dist=1.5)

# draw a bunch of trees, fix layout
igraph_options(plot.layout="layout_as_tree")
plot(make_tree(20, 2))
plot(make_tree(50, 3), vertex.size=3, vertex.label=NA)
tkplot(make_tree(50, 2, mode="undirected"), vertex.size=10,
        vertex.color="green")

## End(Not run)

---

**dyad_census**  

**Dyad census of a graph**

**Description**

Classify dyads in a directed graph. The relationship between each pair of vertices is measured. It can be in three states: mutual, asymmetric or non-existent.

**Usage**

```r
dyad_census(graph)
```

**Arguments**

- `graph`  
The input graph. A warning is given if it is not directed.

**Value**

A named numeric vector with three elements:

- `mut`  
The number of pairs with mutual connections.
- `asym`  
The number of pairs with non-mutual connections.
- `null`  
The number of pairs with no connection between them.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**References**


**See Also**

- `triad_census` for the same classification, but with triples.
Examples

```r
g <- sample_pa(100)
dyad_census(g)
```

### Description

An edge sequence is a vector containing numeric edge ids, with a special class attribute that allows custom operations: selecting subsets of edges based on attributes, or graph structure, creating the intersection, union of edges, etc.

### Usage

```r
E(graph, P = NULL, path = NULL, directed = TRUE)
```

### Arguments

- **graph**: The graph.
- **P**: A list of vertices to select edges via pairs of vertices. The first and second vertices select the first edge, the third and fourth the second, etc.
- **path**: A list of vertices, to select edges along a path. Note that this only works reliably for simple graphs. If the graph has multiple edges, one of them will be chosen arbitrarily to be included in the edge sequence.
- **directed**: Whether to consider edge directions in the `P` argument, for directed graphs.

### Details

Edge sequences are usually used as igraph function arguments that refer to edges of a graph.

An edge sequence is tied to the graph it refers to: it really denoted the specific edges of that graph, and cannot be used together with another graph.

An edge sequence is most often created by the `E()` function. The result includes edges in increasing edge id order by default (if none of the `P` and `path` arguments are used). An edge sequence can be indexed by a numeric vector, just like a regular R vector. See links to other edge sequence operations below.

### Value

An edge sequence of the graph.

### Indexing edge sequences

Edge sequences mostly behave like regular vectors, but there are some additional indexing operations that are specific for them; e.g. selecting edges based on graph structure, or based on edge attributes. See `.igraph.es` for details.

### Querying or setting attributes

Edge sequences can be used to query or set attributes for the edges in the sequence. See `$igraph.es` for details.
each_edge

See Also
Other vertex and edge sequences: \( V() \), \( \text{igraph-es-attributes} \), \( \text{igraph-es-indexing2} \), \( \text{igraph-es-indexing} \), \( \text{igraph-vs-attributes} \), \( \text{igraph-vs-indexing2} \), \( \text{igraph-vs-indexing} \), \( \text{print.igraph.es()} \), \( \text{print.igraph.vs()} \)

Examples

```
# Edges of an unnamed graph
g <- make_ring(10)
E(g)

# Edges of a named graph
g2 <- make_ring(10) %>%
  set_vertex_attr("name", value = letters[1:10])
E(g2)
```

---

**each_edge**

Rewires the endpoints of the edges of a graph to a random vertex

Description

This function can be used together with \text{rewire}. This method rewires the endpoints of the edges with a constant probability uniformly randomly to a new vertex in a graph.

Usage

```
each_edge(
  prob,
  loops = FALSE,
  multiple = FALSE,
  mode = c("all", "out", "in", "total")
)
```

Arguments

- **prob** The rewiring probability, a real number between zero and one.
- **loops** Logical scalar, whether loop edges are allowed in the rewired graph.
- **multiple** Logical scalar, whether multiple edges are allowed in the generated graph.
- **mode** Character string, specifies which endpoint of the edges to rewire in directed graphs. ‘all’ rewires both endpoints, ‘in’ rewires the start (tail) of each directed edge, ‘out’ rewires the end (head) of each directed edge. Ignored for undirected graphs.

Details

Note that this method might create graphs with multiple and/or loop edges.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>
eccentricity

Description

The eccentricity of a vertex is its shortest path distance from the farthest other node in the graph.

Usage

eccentricity(graph, vids = V(graph), mode = c("all", "out", "in", "total"))

Arguments

graph The input graph, it can be directed or undirected.
vids The vertices for which the eccentricity is calculated.
mode Character constant, gives whether the shortest paths to or from the given vertices should be calculated for directed graphs. If out then the shortest paths from the vertex, if in then to it will be considered. If all, the default, then the corresponding undirected graph will be used, edge directions will be ignored. This argument is ignored for undirected graphs.

Details

The eccentricity of a vertex is calculated by measuring the shortest distance from (or to) the vertex, to (or from) all vertices in the graph, and taking the maximum.

This implementation ignores vertex pairs that are in different components. Isolate vertices have eccentricity zero.

Value

eccentricity returns a numeric vector, containing the eccentricity score of each given vertex.

References

**See Also**

`radius` for a related concept, `distances` for general shortest path calculations.

**Examples**

```r
g <- make_star(10, mode="undirected")
eccentricity(g)
```

---

**Description**

This is a helper function that simplifies adding and deleting edges to/from graphs.

**Usage**

```r
dummy_function()  
edummy_function(...)
```

**Arguments**

... See details below.

**Details**

`edges` is an alias for `edge`.

When adding edges via `+`, all unnamed arguments of `edge` (or `edges`) are concatenated, and then passed to `add_edges`. They are interpreted as pairs of vertex ids, and an edge will added between each pair. Named arguments will be used as edge attributes for the new edges.

When deleting edges via `-`, all arguments of `edge` (or `edges`) are concatenated via `c()` and passed to `delete_edges`.

**Value**

A special object that can be used with together with igraph graphs and the plus and minus operators.

**See Also**

Other functions for manipulating graph structure: `.igraph()`, `add_edges()`, `add_vertices()`, `delete_edges()`, `delete_vertices()`, `igraph-minus`, `path()`, `vertex()`
edge_attr

Examples

\[
g \leftarrow \text{make\_ring}(10) \ %\% \\
\quad \text{set\_edge\_attr("color", \ value = "red")}
\]

\[
g \leftarrow g + \text{edge}(1, 5, \ \text{color = "green"}) + \\
\quad \text{edge}(2, 6, \ \text{color = "blue"}) - \\
\quad \text{edge("8|9")}
\]

\[
E(g)[[]]
\]

\[
g %>% \\
\quad \text{add\_layout\_in\_circle()} %>% \\
\quad \text{plot()}
\]

\[
g \leftarrow \text{make\_ring}(10) + \text{edges}(1:10)
\]

\[
\text{plot(g)}
\]

Description

Query edge attributes of a graph

Usage

\[
\text{edge\_attr(graph, name, index = E(graph))}
\]

Arguments

- **graph**: The graph
- **name**: The name of the attribute to query. If missing, then all edge attributes are returned in a list.
- **index**: An optional edge sequence, to query edge attributes for a subset of edges.

Value

The value of the edge attribute, or the list of all edge attributes if name is missing.

See Also

Other graph attributes: `delete_edge_attr()`, `delete_graph_attr()`, `delete_vertex_attr()`, `edge_attr<-()`, `edge_attr_names()`, `graph_attr<-()`, `graph_attr_names()`, `graph_attr()`, `igraph-dollar`, `igraph-vs-attributes`, `set_edge_attr()`, `set_graph_attr()`, `set_vertex_attr()`, `vertex_attr<-()`, `vertex_attr_names()`, `vertex_attr()`

Examples

\[
g \leftarrow \text{make\_ring}(10) \ %\% \\
\quad \text{set\_edge\_attr("weight", \ value = 1:10) %\%} \\
\quad \text{set\_edge\_attr("color", \ value = "red")}
\]

\[
g \leftarrow \text{plot(g, \ edge\_width = E(g)$weight)}
\]
### edge_attr_names

**List names of edge attributes**

**Description**

List names of edge attributes

**Usage**

```r
edge_attr_names(graph)
```

**Arguments**

- `graph` The graph.

**Value**

Character vector, the names of the edge attributes.

**See Also**

Other graph attributes: `delete_edge_attr()`, `delete_graph_attr()`, `delete_vertex_attr()`, `edge_attr<-()`, `edge_attr()`, `graph_attr<-()`, `graph_attr()`, `igraph-dollar`, `igraph-vs-attributes`, `set_edge_attr()`, `set_graph_attr()`, `set_vertex_attr()`, `vertex_attr<-()`, `vertex_attr_names()`.

**Examples**

```r
g <- make_ring(10) %>%
  set_edge_attr("label", value = letters[1:10])
edge_attr_names(g)
plot(g)
```

### edge_attr<-

**Set one or more edge attributes**

**Description**

Set one or more edge attributes

**Usage**

```r
edge_attr(graph, name, index = E(graph)) <- value
```

**Arguments**

- `graph` The graph.
- `name` The name of the edge attribute to set. If missing, then value must be a named list, and its entries are set as edge attributes.
- `index` An optional edge sequence to set the attributes of a subset of edges.
- `value` The new value of the attribute(s) for all (or index) edges.
value

The graph, with the edge attribute(s) added or set.

see also

Other graph attributes: delete_edge_attr(), delete_graph_attr(), delete_vertex_attr(), edge_attr_names(), edge_attr(), graph_attr<-(), graph_attr_names(), graph_attr(), igraph-dollar, igraph-vs-attributes, set_edge_attr(), set_graph_attr(), set_vertex_attr(), vertex_attr<-(), vertex_attr_names(), vertex_attr()

examples

g <- make_ring(10)
edge_attr(g) <- list(name = LETTERS[1:10],
  color = rep("green", gsize(g)))
edge_attr(g, "label") <- E(g)$name
g
plot(g)

edge_connectivity

Edge connectivity.

description

The edge connectivity of a graph or two vertices, this is recently also called group adhesion.

usage

edge_connectivity(graph, source = NULL, target = NULL, checks = TRUE)

arguments

graph The input graph.
source The id of the source vertex, for edge_connectivity it can be NULL, see details below.
target The id of the target vertex, for edge_connectivity it can be NULL, see details below.
checks Logical constant. Whether to check that the graph is connected and also the degree of the vertices. If the graph is not (strongly) connected then the connectivity is obviously zero. Otherwise if the minimum degree is one then the edge connectivity is also one. It is a good idea to perform these checks, as they can be done quickly compared to the connectivity calculation itself. They were suggested by Peter McMahan, thanks Peter.
The edge connectivity of a pair of vertices (source and target) is the minimum number of edges needed to remove to eliminate all (directed) paths from source to target. edge_connectivity calculates this quantity if both the source and target arguments are given (and not NULL).

The edge connectivity of a graph is the minimum of the edge connectivity of every (ordered) pair of vertices in the graph. edge_connectivity calculates this quantity if neither the source nor the target arguments are given (i.e., they are both NULL).

A set of edge disjoint paths between two vertices is a set of paths between them containing no common edges. The maximum number of edge disjoint paths between two vertices is the same as their edge connectivity.

The adhesion of a graph is the minimum number of edges needed to remove to obtain a graph which is not strongly connected. This is the same as the edge connectivity of the graph.

The three functions documented on this page calculate similar properties, more precisely the most general is edge_connectivity, the others are included only for having more descriptive function names.

Value

A scalar real value.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References

Douglas R. White and Frank Harary: The cohesiveness of blocks in social networks: node connectivity and conditional density, TODO: citation

See Also

max_flow, vertex_connectivity, vertex_disjoint_paths, cohesion

Examples

g <- barabasi.game(100, m=1)
g2 <- barabasi.game(100, m=5)
edge_connectivity(g, 100, 1)
edge_connectivity(g2, 100, 1)
edge_disjoint_paths(g2, 100, 1)

g <- sample_gnp(50, 5/50)
g <- as.directed(g)
g <- induced_subgraph(g, subcomponent(g, 1))
adhesion(g)
edge_density

Graph density

Description

The density of a graph is the ratio of the number of edges and the number of possible edges.

Usage

edge_density(graph, loops = FALSE)

Arguments

graph The input graph.
loops Logical constant, whether to allow loop edges in the graph. If this is TRUE then self loops are considered to be possible. If this is FALSE then we assume that the graph does not contain any loop edges and that loop edges are not meaningful.

Details

Note that this function may return strange results for graph with multiple edges, density is ill-defined for graphs with multiple edges.

Value

A real constant. This function returns NaN (=0.0/0.0) for an empty graph with zero vertices.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

vcount, ecount, simplify to get rid of the multiple and/or loop edges.

Examples

g1 <- make_empty_graph(n=10)
g2 <- make_full_graph(n=10)
g3 <- sample_gnp(n=10, 0.4)

# loop edges
g <- graph( c(1,2, 2,2, 2,3) )
edge_density(g, loops=FALSE) # this is wrong!!!
edge_density(g, loops=TRUE)  # this is right!!!
edge_density(simplify(g), loops=FALSE) # this is also right, but different
**Description**

eigen_centrality takes a graph (graph) and returns the eigenvector centralities of positions \( v \) within it.

**Usage**

eigen_centrality(graph, directed = FALSE, scale = TRUE, weights = NULL, options = arpack_defaults)

**Arguments**

- **graph**: Graph to be analyzed.
- **directed**: Logical scalar, whether to consider direction of the edges in directed graphs. It is ignored for undirected graphs.
- **scale**: Logical scalar, whether to scale the result to have a maximum score of one. If no scaling is used then the result vector has unit length in the Euclidean norm.
- **weights**: A numerical vector or NULL. This argument can be used to give edge weights for calculating the weighted eigenvector centrality of vertices. If this is NULL and the graph has a weight edge attribute then that is used. If **weights** is a numerical vector then it used, even if the graph has a weight edge attribute. If this is NA, then no edge weights are used (even if the graph has a weight edge attribute). Note that if there are negative edge weights and the direction of the edges is considered, then the eigenvector might be complex. In this case only the real part is reported. This function interprets weights as connection strength. Higher weights spread the centrality better.
- **options**: A named list, to override some ARPACK options. See arpack for details.

**Details**

Eigenvector centrality scores correspond to the values of the first eigenvector of the graph adjacency matrix; these scores may, in turn, be interpreted as arising from a reciprocal process in which the centrality of each actor is proportional to the sum of the centralities of those actors to whom he or she is connected. In general, vertices with high eigenvector centralities are those which are connected to many other vertices which are, in turn, connected to many others (and so on). (The perceptive may realize that this implies that the largest values will be obtained by individuals in large cliques (or high-density substructures). This is also intelligible from an algebraic point of view, with the first eigenvector being closely related to the best rank-1 approximation of the adjacency matrix (a relationship which is easy to see in the special case of a diagonalizable symmetric real matrix via the \( SLS^{-1} \) decomposition).)
The adjacency matrix used in the eigenvector centrality calculation assumes that loop edges are counted twice; this is because each loop edge has two endpoints that are both connected to the same vertex, and you could traverse the loop edge via either endpoint.

In the directed case, the left eigenvector of the adjacency matrix is calculated. In other words, the centrality of a vertex is proportional to the sum of centralities of vertices pointing to it.

Eigenvector centrality is meaningful only for connected graphs. Graphs that are not connected should be decomposed into connected components, and the eigenvector centrality calculated for each separately. This function does not verify that the graph is connected. If it is not, in the undirected case the scores of all but one component will be zeros.

Also note that the adjacency matrix of a directed acyclic graph or the adjacency matrix of an empty graph does not possess positive eigenvalues, therefore the eigenvector centrality is not defined for these graphs. igraph will return an eigenvalue of zero in such cases. The eigenvector centralities will all be equal for an empty graph and will all be zeros for a directed acyclic graph. Such pathological cases can be detected by checking whether the eigenvalue is very close to zero.

From igraph version 0.5 this function uses ARPACK for the underlying computation, see arpack for more about ARPACK in igraph.

**Value**

A named list with components:

- `vector` A vector containing the centrality scores.
- `value` The eigenvalue corresponding to the calculated eigenvector, i.e. the centrality scores.
- `options` A named list, information about the underlying ARPACK computation. See arpack for the details.

**Author(s)**


**References**


**Examples**

```r
#Generate some test data
g <- make_ring(10, directed=FALSE)
#Compute eigenvector centrality scores
eigen_centrality(g)
```
**embed_adjacency_matrix**

*Spectral Embedding of Adjacency Matrices*

**Description**

Spectral decomposition of the adjacency matrices of graphs.

**Usage**

```r
embed_adjacency_matrix(
  graph,  
  no,    
  weights = NULL, 
  which = c("lm", "la", "sa"), 
  scaled = TRUE, 
  cvec = graph.strength(graph, weights = weights)/(vcount(graph) - 1), 
  options = igraph.arpack.default
)
```

**Arguments**

- `graph` The input graph, directed or undirected.
- `no` An integer scalar. This value is the embedding dimension of the spectral embedding. Should be smaller than the number of vertices. The largest no-dimensional non-zero singular values are used for the spectral embedding.
- `weights` Optional positive weight vector for calculating a weighted embedding. If the graph has a weight edge attribute, then this is used by default. In a weighted embedding, the edge weights are used instead of the binary adjacency matrix.
- `which` Which eigenvalues (or singular values, for directed graphs) to use. 'lm' means the ones with the largest magnitude, 'la' is the ones (algebraic) largest, and 'sa' is the (algebraic) smallest eigenvalues. The default is 'lm'. Note that for directed graphs 'la' and 'lm' are the equivalent, because the singular values are used for the ordering.
- `scaled` Logical scalar, if FALSE, then U and V are returned instead of X and Y.
- `cvec` A numeric vector, its length is the number vertices in the graph. This vector is added to the diagonal of the adjacency matrix.
- `options` A named list containing the parameters for the SVD computation algorithm in ARPACK. By default, the list of values is assigned the values given by `igraph.arpack.default`.

**Details**

This function computes a no-dimensional Euclidean representation of the graph based on its adjacency matrix, $A$. This representation is computed via the singular value decomposition of the adjacency matrix, $A = UDV^T$. In the case, where the graph is a random dot product graph generated using latent position vectors in $R^{no}$ for each vertex, the embedding will provide an estimate of these latent vectors.

For undirected graphs the latent positions are calculated as $X = U^{no}D^{1/2}$, where $U^{no}$ equals to the first no columns of $U$, and $D^{1/2}$ is a diagonal matrix containing the top no singular values on the diagonal.
For directed graphs the embedding is defined as the pair $X = U^{no} D^{1/2}$ and $Y = V^{no} D^{1/2}$. (For undirected graphs $U = V$, so it is enough to keep one of them.)

Value

A list containing with entries:

- $X$: Estimated latent positions, an $n$ times $no$ matrix, $n$ is the number of vertices.
- $Y$: NULL for undirected graphs, the second half of the latent positions for directed graphs, an $n$ times $no$ matrix, $n$ is the number of vertices.
- $D$: The eigenvalues (for undirected graphs) or the singular values (for directed graphs) calculated by the algorithm.
- options: A named list, information about the underlying ARPACK computation. See `arpack` for the details.

References


See Also

`sample_dot_product`

Examples

```r
## A small graph
lpvs <- matrix(rnorm(200), 20, 10)
lpvs <- apply(lpvs, 2, function(x) { return (abs(x)/sqrt(sum(x^2))) })
RDP <- sample_dot_product(lpvs)
embed <- embed_adjacency_matrix(RDP, 5)
```

```
embed_laplacian_matrix
Spectral Embedding of the Laplacian of a Graph

Description

Spectral decomposition of Laplacian matrices of graphs.

Usage

`embed_laplacian_matrix(`
  graph,
  no,
  weights = NULL,
  which = c("lm", "la", "sa"),
  type = c("default", "D-A", "DAD", "I-DAD", "OAP"),
  scaled = TRUE,
  options = igraph.arpack.default
)`
embed_laplacian_matrix

Arguments

graph
The input graph, directed or undirected.

no An integer scalar. This value is the embedding dimension of the spectral embedding. Should be smaller than the number of vertices. The largest no-dimensional non-zero singular values are used for the spectral embedding.

weights Optional positive weight vector for calculating a weighted embedding. If the graph has a weight edge attribute, then this is used by default. For weighted embedding, edge weights are used instead of the binary adjacency matrix, and vertex strength (see strength) is used instead of the degrees.

which Which eigenvalues (or singular values, for directed graphs) to use. ‘lm’ means the ones with the largest magnitude, ‘la’ is the ones (algebraic) largest, and ‘sa’ is the (algebraic) smallest eigenvalues. The default is ‘lm’. Note that for directed graphs ‘la’ and ‘lm’ are the equivalent, because the singular values are used for the ordering.

type The type of the Laplacian to use. Various definitions exist for the Laplacian of a graph, and one can choose between them with this argument.

Possible values: D−A means $D - A$ where $D$ is the degree matrix and $A$ is the adjacency matrix; DAD means $D^{1/2} A D^{1/2}$, $D^{1/2}$ is the inverse of the square root of the degree matrix; I−DAD means $I - D^{1/2}$, where $I$ is the identity matrix. OAP is $O^{1/2} A P^{1/2}$, where $O^{1/2}$ is the inverse of the square root of the out-degree matrix and $P^{1/2}$ is the same for the in-degree matrix. OAP is not defined for undirected graphs, and is the only defined type for directed graphs.

The default (i.e. type default) is to use D−A for undirected graphs and OAP for directed graphs.

scaled Logical scalar, if FALSE, then $U$ and $V$ are returned instead of $X$ and $Y$.

options A named list containing the parameters for the SVD computation algorithm in ARPACK. By default, the list of values is assigned the values given by igraph.arpack.default.

Details

This function computes a no-dimensional Euclidean representation of the graph based on its Laplacian matrix, $L$. This representation is computed via the singular value decomposition of the Laplacian matrix.

They are essentially doing the same as embed_adjacency_matrix, but work on the Laplacian matrix, instead of the adjacency matrix.

Value

A list containing with entries:

X Estimated latent positions, an n times no matrix, n is the number of vertices.

Y NULL for undirected graphs, the second half of the latent positions for directed graphs, an n times no matrix, n is the number of vertices.

D The eigenvalues (for undirected graphs) or the singular values (for directed graphs) calculated by the algorithm.

options A named list, information about the underlying ARPACK computation. See arpack for the details.
Incident vertices of some graph edges

Description
Incident vertices of some graph edges

Usage
ends(graph, es, names = TRUE)

Arguments

- graph: The input graph
- es: The sequence of edges to query
- names: Whether to return vertex names or numeric vertex ids. By default vertex names are used.

Value
A two column matrix of vertex names or vertex ids.

See Also
Other structural queries: [.igraph(), [[.igraph(), adjacent_vertices(), are_adjacent(), get.edge.id(), gorder(), gsize(), head_of(), incident_edges(), incident(), is.directed(), neighbors(), tail_of()]

Examples
```
g <- make_ring(5)
ends(g, E(g))
```
erdos.renyi.game

Generate random graphs according to the Erdos-Renyi model

Description
This model is very simple, every possible edge is created with the same constant probability.

Usage

```r
erdos.renyi.game(
  n,
  p.or.m,
  type = c("gnp", "gnm"),
  directed = FALSE,
  loops = FALSE
)
```

Arguments

- `n` The number of vertices in the graph.
- `p.or.m` Either the probability for drawing an edge between two arbitrary vertices (G(n,p) graph), or the number of edges in the graph (for G(n,m) graphs).
- `type` The type of the random graph to create, either `gnp` (G(n,p) graph) or `gnm` (G(n,m) graph).
- `directed` Logical, whether the graph will be directed, defaults to FALSE.
- `loops` Logical, whether to add loop edges, defaults to FALSE.

Details

In G(n,p) graphs, the graph has `n` vertices and for each edge the probability that it is present in the graph is `p`.

In G(n,m) graphs, the graph has `n` vertices and `m` edges, and the `m` edges are chosen uniformly randomly from the set of all possible edges. This set includes loop edges as well if the `loops` parameter is TRUE.

`random.graph.game` is an alias to this function.

Value

A graph object.

Deprecated

Since igraph version 0.8.0, both `erdos.renyi.game` and `random.graph.game` are deprecated, and `sample_gnp` and `sample_gnm` should be used instead.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>
estimate_betweenness

References


See Also

.sample_pa

Examples

```r
g <- erdos.renyi.game(1000, 1/1000)
degree_distribution(g)
```

---

**estimate_betweenness**  
Vertex and edge betweenness centrality

**Description**

The vertex and edge betweenness are (roughly) defined by the number of geodesics (shortest paths) going through a vertex or an edge.

**Usage**

```r
estimate_betweenness(
  graph,
  vids = V(graph),
  directed = TRUE,
  cutoff,
  weights = NULL,
  nobigint = TRUE
)

betweenness(
  graph,
  v = V(graph),
  directed = TRUE,
  weights = NULL,
  nobigint = TRUE,
  normalized = FALSE,
  cutoff = -1
)

edge_betweenness(
  graph,
  e = E(graph),
  directed = TRUE,
  weights = NULL,
  cutoff = -1
)
```
estimate_betweenness

**Arguments**

- `graph`: The graph to analyze.
- `vids`: The vertices for which the vertex betweenness estimation will be calculated.
- `directed`: Logical, whether directed paths should be considered while determining the shortest paths.
- `cutoff`: The maximum path length to consider when calculating the betweenness. If zero or negative then there is no such limit.
- `weights`: Optional positive weight vector for calculating weighted betweenness. If the graph has a weight edge attribute, then this is used by default. Weights are used to calculate weighted shortest paths, so they are interpreted as distances.
- `nobigint`: Logical scalar, whether to use big integers during the calculation. Deprecated since igraph 1.3 and will be removed in igraph 1.4.
- `v`: The vertices for which the vertex betweenness will be calculated.
- `normalized`: Logical scalar, whether to normalize the betweenness scores. If TRUE, then the results are normalized by the number of ordered or unordered vertex pairs in directed and undirected graphs, respectively. In an undirected graph,

  \[ B^n = \frac{2B}{(n - 1)(n - 2)} \]

  where \( B^n \) is the normalized, \( B \) the raw betweenness, and \( n \) is the number of vertices in the graph.

- `e`: The edges for which the edge betweenness will be calculated.

**Details**

The vertex betweenness of vertex \( v \) is defined by

\[
\sum_{i \neq j, i \neq v, j \neq v} g_{ij}/g_{ij}.
\]

The edge betweenness of edge \( e \) is defined by

\[
\sum_{i \neq j} g_{iej}/g_{ij}.
\]

`betweenness` calculates vertex betweenness, `edge_betweenness` calculates edge betweenness.

Here \( g_{ij} \) is the total number of shortest paths between vertices \( i \) and \( j \) while \( g_{iej} \) is the number of those shortest paths which pass though vertex \( v \).

Both functions allow you to consider only paths of length `cutoff` or smaller; this can be run for larger graphs, as the running time is not quadratic (if `cutoff` is small). If `cutoff` is zero or negative, then the function calculates the exact betweenness scores. Using zero as a cutoff is deprecated and future versions (from 1.4.0) will treat zero cutoff literally (i.e. no paths considered at all). If you want no cutoff, use a negative number.

`estimate_betweenness` and `estimate_edge_betweenness` are aliases for `betweenness` and `edge_betweenness`, with a different argument order, for sake of compatibility with older versions of igraph.

For calculating the betweenness a similar algorithm to the one proposed by Brandes (see References) is used.
Value

A numeric vector with the betweenness score for each vertex in \( v \) for `betweenness`.

A numeric vector with the edge betweenness score for each edge in \( e \) for `edge_betweenness`.

Note

`edge_betweenness` might give false values for graphs with multiple edges.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

closeness, degree, harmonic_centrality

Examples

g <- sample_gnp(10, 3/10)
betweenness(g)
edge_betweenness(g)

feedback_arc_set

Finding a feedback arc set in a graph

Description

A feedback arc set of a graph is a subset of edges whose removal breaks all cycles in the graph.

Usage

`feedback_arc_set(graph, weights = NULL, algo = c("approx_eades", "exact_ip"))`

Arguments

graph The input graph

weights Potential edge weights. If the graph has an edge attribute called `weight`, and this argument is `NULL`, then the edge attribute is used automatically. The goal of the feedback arc set problem is to find a feedback arc set with the smallest total weight.

algo Specifies the algorithm to use. "exact_ip" solves the feedback arc set problem with an exact integer programming algorithm that guarantees that the total weight of the removed edges is as small as possible. "approx_eades" uses a fast (linear-time) approximation algorithm from Eades, Lin and Smyth. "exact" is an alias to "exact_ip" while "approx" is an alias to "approx_eades".
Details

Feedback arc sets are typically used in directed graphs. The removal of a feedback arc set of a directed graph ensures that the remaining graph is a directed acyclic graph (DAG). For undirected graphs, the removal of a feedback arc set ensures that the remaining graph is a forest (i.e. every connected component is a tree).

Value

An edge sequence (by default, but see the return.vs.es option of igraph_options) containing the feedback arc set.

References


Examples

g <- sample_gnm(20, 40, directed=TRUE)
feedback_arc_set(g)
feedback_arc_set(g, algo="approx")

fit_hrg  
Fit a hierarchical random graph model

Description

fit_hrg fits a HRG to a given graph. It takes the specified steps number of MCMC steps to perform the fitting, or a convergence criteria if the specified number of steps is zero. fit_hrg can start from a given HRG, if this is given in the hrg argument and the start argument is TRUE.

Usage

fit_hrg(graph, hrg = NULL, start = FALSE, steps = 0)

Arguments

graph  The graph to fit the model to. Edge directions are ignored in directed graphs.

hrg  A hierarchical random graph model, in the form of an igraphHRG object. fit_hrg allows this to be NULL, in which case a random starting point is used for the fitting.

start  Logical, whether to start the fitting/sampling from the supplied igraphHRG object, or from a random starting point.

steps  The number of MCMC steps to make. If this is zero, then the MCMC procedure is performed until convergence.
Value

`fit_hrg` returns an `igraphHRG` object. This is a list with the following members:

- **left**: Vector that contains the left children of the internal tree vertices. The first vertex is always the root vertex, so the first element of the vector is the left child of the root vertex. Internal vertices are denoted with negative numbers, starting from -1 and going down, i.e. the root vertex is -1. Leaf vertices are denoted by non-negative number, starting from zero and up.

- **right**: Vector that contains the right children of the vertices, with the same encoding as the `left` vector.

- **prob**: The connection probabilities attached to the internal vertices, the first number belongs to the root vertex (i.e. internal vertex -1), the second to internal vertex -2, etc.

- **edges**: The number of edges in the subtree below the given internal vertex.

- **vertices**: The number of vertices in the subtree below the given internal vertex, including itself.

References


See Also

Other hierarchical random graph functions: `consensus_tree()`, `hrg-methods`, `hrg_tree()`, `hrg()`, `predict_edges()`, `print.igraphHRGConsensus()`, `print.igraphHRG()`, `sample_hrg()`

Examples

```r
## We are not running these examples any more, because they
## take a long time (~15 seconds) to run and this is against the CRAN
## repository policy. Copy and paste them by hand to your R prompt if
## you want to run them.

## Not run:
## A graph with two dense groups
g <- sample_gnp(10, p=1/2) + sample_gnp(10, p=1/2)
hrg <- fit_hrg(g)
hrg

## The consensus tree for it
consensus_tree(g, hrg=hrg, start=TRUE)

## Prediction of missing edges
g2 <- make_full_graph(4) + (make_full_graph(4) - path(1,2))
predict_edges(g2)

## End(Not run)
```
Description

`fit_power_law` fits a power-law distribution to a data set.

Usage

```r
fit_power_law(
  x,
  xmin = NULL,
  start = 2,
  force.continuous = FALSE,
  implementation = c("plfit", "R.mle"),
  ...)
```

Arguments

- `x` The data to fit, a numeric vector. For implementation `R.mle` the data must be integer values. For the `plfit` implementation non-integer values might be present and then a continuous power-law distribution is fitted.
- `xmin` Numeric scalar, or `NULL`. The lower bound for fitting the power-law. If `NULL`, the smallest value in `x` will be used for the `R.mle` implementation, and its value will be automatically determined for the `plfit` implementation. This argument makes it possible to fit only the tail of the distribution.
- `start` Numeric scalar. The initial value of the exponent for the minimizing function, for the `R.mle` implementation. Usually it is safe to leave this untouched.
- `force.continuous` Logical scalar. Whether to force a continuous distribution for the `plfit` implementation, even if the sample vector contains integer values only (by chance). If this argument is false, igraph will assume a continuous distribution if at least one sample is non-integer and assume a discrete distribution otherwise.
- `implementation` Character scalar. Which implementation to use. See details below.
- `...` Additional arguments, passed to the maximum likelihood optimizing function, `mle`, if the `R.mle` implementation is chosen. It is ignored by the `plfit` implementation.

Details

This function fits a power-law distribution to a vector containing samples from a distribution (that is assumed to follow a power-law of course). In a power-law distribution, it is generally assumed that \( P(X = x) \) is proportional to \( x^{-\alpha} \), where \( x \) is a positive number and \( \alpha \) is greater than 1. In many real-world cases, the power-law behaviour kicks in only above a threshold value \( x_{\text{min}} \). The goal of this function is to determine \( \alpha \) if \( x_{\text{min}} \) is given, or to determine \( x_{\text{min}} \) and the corresponding value of \( \alpha \).

`fit_power_law` provides two maximum likelihood implementations. If the implementation argument is `R.mle`, then the BFGS optimization (see `mle`) algorithm is applied. The additional
arguments are passed to the mle function, so it is possible to change the optimization method and/or its parameters. This implementation can not fit the $x_{min}$ argument, so use the ‘plfit’ implementation if you want to do that.

The ‘plfit’ implementation also uses the maximum likelihood principle to determine $\alpha$ for a given $x_{min}$; When $x_{min}$ is not given in advance, the algorithm will attempt to find its optimal value for which the p-value of a Kolmogorov-Smirnov test between the fitted distribution and the original sample is the largest. The function uses the method of Clauset, Shalizi and Newman to calculate the parameters of the fitted distribution. See references below for the details.

**Value**

Depends on the implementation argument. If it is ‘R.mle’, then an object with class ‘mle’. It can be used to calculate confidence intervals and log-likelihood. See mle-class for details.

If implementation is ‘plfit’, then the result is a named list with entries:

- **continuous** Logical scalar, whether the fitted power-law distribution was continuous or discrete.
- **alpha** Numeric scalar, the exponent of the fitted power-law distribution.
- **xmin** Numeric scalar, the minimum value from which the power-law distribution was fitted. In other words, only the values larger than $x_{min}$ were used from the input vector.
- **logLik** Numeric scalar, the log-likelihood of the fitted parameters.
- **KS.stat** Numeric scalar, the test statistic of a Kolmogorov-Smirnov test that compares the fitted distribution with the input vector. Smaller scores denote better fit.
- **KS.p** Numeric scalar, the p-value of the Kolmogorov-Smirnov test. Small p-values (less than 0.05) indicate that the test rejected the hypothesis that the original data could have been drawn from the fitted power-law distribution.

**Author(s)**

Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com>

**References**


**See Also**

mle

**Examples**

```r
# This should approximately yield the correct exponent 3
g <- barabasi.game(1000) # increase this number to have a better estimate
d <- degree(g, mode="in")
fit1 <- fit_power_law(d+1, 10)
fit2 <- fit_power_law(d+1, 10, implementation="R.mle")
fit1$alpha
```
get.edge.ids

Find the edge ids based on the incident vertices of the edges

Description

Find the edges in an igraph graph that have the specified end points. This function handles multigraph (graphs with multiple edges) and can consider or ignore the edge directions in directed graphs.

Usage

get.edge.ids(graph, vp, directed = TRUE, error = FALSE, multi = FALSE)

Arguments

- **graph**: The input graph.
- **vp**: The incident vertices, given as vertex ids or symbolic vertex names. They are interpreted pairwise, i.e. the first and second are used for the first edge, the third and fourth for the second, etc.
- **directed**: Logical scalar, whether to consider edge directions in directed graphs. This argument is ignored for undirected graphs.
- **error**: Logical scalar, whether to report an error if an edge is not found in the graph. If FALSE, then no error is reported, and zero is returned for the non-existant edge(s).
- **multi**: Logical scalar, whether to handle multiple edges properly. If FALSE, and a pair of vertices are given twice (or more), then always the same edge id is reported back for them. If TRUE, then the edge ids of multiple edges are correctly reported.

Details

igraph vertex ids are natural numbers, starting from one, up to the number of vertices in the graph. Similarly, edges are also numbered from one, up to the number of edges.

This function allows finding the edges of the graph, via their incident vertices.

Value

A numeric vector of edge ids, one for each pair of input vertices. If there is no edge in the input graph for a given pair of vertices, then zero is reported. (If the error argument is FALSE.)

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

Other structural queries: [.igraph(), [[.igraph(). adjacent_vertices(). are_adjacent(). ends(). gorder(). gsize(). head_of(). incident_edges(). incident(). is_directed(). neighbors(). tail_of()
girth

Examples

g <- make_ring(10)
ei <- get.edge.ids(g, c(1,2, 4,5))
E(g)[ei]

## non-existant edge
get.edge.ids(g, c(2,1, 1,4, 5,4))

## multiple edges
## multi = FALSE, a single edge id is returned,
## as many times as corresponding pairs in the vertex series.
g <- make_graph(rep(c(1,2), 5))
eis <- get.edge.ids(g, c(1,2, 1,2), multi=FALSE)
eis
E(g)[eis]

## multi = TRUE, as many different edges, if any,
## are returned as pairs in the vertex series.
eim <- get.edge.ids(g, c(1,2, 1,2, 1,2), multi=TRUE)
eim
E(g)[eim]

________________________
girth  Girth of a graph

description

The girth of a graph is the length of the shortest circle in it.

Usage

girth(graph, circle = TRUE)

Arguments

graph  The input graph. It may be directed, but the algorithm searches for undirected circles anyway.
circle Logical scalar, whether to return the shortest circle itself.

details

The current implementation works for undirected graphs only, directed graphs are treated as undirected graphs. Loop edges and multiple edges are ignored. If the graph is a forest (ie. acyclic), then zero is returned.

This implementation is based on Alon Itai and Michael Rodeh: Finding a minimum circuit in a graph Proceedings of the ninth annual ACM symposium on Theory of computing, 1-10, 1977. The first implementation of this function was done by Keith Briggs, thanks Keith.
global_efficiency

Value
A named list with two components:

- `girth`: Integer constant, the girth of the graph, or 0 if the graph is acyclic.
- `circle`: Numeric vector with the vertex ids in the shortest circle.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

References
Alon Itai and Michael Rodeh: Finding a minimum circuit in a graph Proceedings of the ninth annual ACM symposium on Theory of computing, 1-10, 1977

Examples

# No circle in a tree
g <- make_tree(1000, 3)
girth(g)

# The worst case running time is for a ring
g <- make_ring(100)
girth(g)

# What about a random graph?
g <- sample_gnp(1000, 1/1000)
girth(g)

global_efficiency  Efficiency of a graph

Description
These functions calculate the global or average local efficiency of a network, or the local efficiency of every vertex in the network. See below for definitions.

Usage

- `global_efficiency(graph, weights = NULL, directed = TRUE)`
- `local_efficiency(graph, vids = V(graph), weights = NULL, directed = TRUE, mode = c("all", "out", "in", "total")
)
- `average_local_efficiency(graph,`
global_efficiency

weights = NULL,
directed = TRUE,
mode = c("all", "out", "in", "total")
)

Arguments

graph
The graph to analyze.

weights
The edge weights. All edge weights must be non-negative; additionally, no edge weight may be NaN. If it is NULL (the default) and the graph has a weight edge attribute, then it is used automatically.

directed
Logical scalar, whether to consider directed paths. Ignored for undirected graphs.

vids
The vertex ids of the vertices for which the calculation will be done. Applies to the local efficiency calculation only.

mode
Specifies how to define the local neighborhood of a vertex in directed graphs. "out" considers out-neighbors only, "in" considers in-neighbors only, "all" considers both.

Value

For global_efficiency, the global efficiency of the graph as a single number. For average_local_efficiency, the average local efficiency of the graph as a single number. For local_efficiency, the local efficiency of each vertex in a vector.

Global efficiency

The global efficiency of a network is defined as the average of inverse distances between all pairs of vertices.

More precisely:

$$E_g = \frac{1}{n(n-1)} \sum_{i \neq j} \frac{1}{d_{ij}}$$

where $n$ is the number of vertices.

The inverse distance between pairs that are not reachable from each other is considered to be zero. For graphs with fewer than 2 vertices, NaN is returned.

Local efficiency

The local efficiency of a network around a vertex is defined as follows: We remove the vertex and compute the distances (shortest path lengths) between its neighbours through the rest of the network. The local efficiency around the removed vertex is the average of the inverse of these distances.

The inverse distance between two vertices which are not reachable from each other is considered to be zero. The local efficiency around a vertex with fewer than two neighbours is taken to be zero by convention.

Average local efficiency

The average local efficiency of a network is simply the arithmetic mean of the local efficiencies of all the vertices; see the definition for local efficiency above.
References


Examples

```r
g <- make_graph("zachary")
global_efficiency(g)
average_local_efficiency(g)
```

---

**gorder**

Order (number of vertices) of a graph

**Description**

vcount is an alias of this function.

**Usage**

`gorder(graph)`

**Arguments**

- `graph` The graph

**Value**

Number of vertices, numeric scalar.

**See Also**

Other structural queries: ```igraph()```, ```[.igraph()```, ```adjacent_vertices()```, ```are_adjacent()```, ```ends()```, ```get.edge.ids()```, ```gsize()```, ```head_of()```, ```incident_edges()```, ```incident()```, ```is_directed()```, ```neighbors()```, ```tail_of()```

**Examples**

```r
g <- make_ring(10)
gorder(g)
vcount(g)
```
Convert object to a graph

Description
This is a generic function to convert R objects to igraph graphs.

Usage
```
graph_(...)  
```

Arguments
```
...  Parameters, see details below.  
```

Details
TODO

Examples
```
## These are equivalent
graph_(cbind(1:5,2:6), from_edgelist(directed = FALSE))
graph_(cbind(1:5,2:6), from_edgelist(), directed = FALSE)
```

Graph attributes of a graph

Description
Graph attributes of a graph

Usage
```
graph_attr(graph, name)  
```

Arguments
```
graph  Input graph.  
name  The name of attribute to query. If missing, then all attributes are returned in a list.  
```

Value
A list of graph attributes, or a single graph attribute.

See Also
Other graph attributes: delete_edge_attr(), delete_graph_attr(), delete_vertex_attr(), edge_attr<-(), edge_attr_names(), edge_attr(), graph_attr<-(), graph_attr_names(), igraph-dollar, igraph-vs-attributes, set_edge_attr(), set_graph_attr(), set_vertex_attr(), vertex_attr<-(), vertex_attr_names(), vertex_attr()
graph_attr<- Examples

    g <- make_ring(10)
    graph_attr(g)
    graph_attr(g, "name")

graph_attr_names List names of graph attributes

Description
List names of graph attributes

Usage
graph_attr_names(graph)

Arguments

graph The graph.

Value
Character vector, the names of the graph attributes.

See Also
Other graph attributes: delete_edge_attr(), delete_graph_attr(), delete_vertex_attr(),
edge_attr<-(), edge_attr_names().edge_attr(), graph_attr<-(), graph_attr().igraph-dollar,
igraph-vs-attributes.set_edge_attr(), set_graph_attr(), set_vertex_attr(), vertex_attr<-(),
vertex_attr_names(), vertex_attr()

Examples
g <- make_ring(10)
graph_attr_names(g)

graph_attr<- Set all or some graph attributes

Description
Set all or some graph attributes

Usage

graph_attr(graph, name) <- value
Arguments

- **graph**  The graph.
- **name**  The name of the attribute to set. If missing, then **value** should be a named list, and all list members are set as attributes.
- **value**  The value of the attribute to set

Value

The graph, with the attribute(s) added.

See Also

Other graph attributes: `delete_edge_attr()`, `delete_graph_attr()`, `delete_vertex_attr()`, `edge_attr<-()`, `edge_attr_names()`, `edge_attr()`, `graph_attr_names()`, `graph_attr()`, `igraph-dollar`, `igraph-vs-attributes`, `set_edge_attr()`, `set_graph_attr()`, `set_vertex_attr()`, `vertex_attr<-()`, `vertex_attr_names()`, `vertex_attr()`

Examples

```r
# Create a graph from an adjacency list
adjlist <- list(
  c(1, 2, 3),  # Adjacency list
  c(4, 5),
  c(6, 7)
)

# Set attributes
graph_attr(g, "name") <- "4-star"

# Plot the graph
plot(g)
```

Description

An adjacency list is a list of numeric vectors, containing the neighbor vertices for each vertex. This function creates an igraph graph object from such a list.

Usage

```r
graph_from_adj_list(
  adjlist,
  mode = c("out", "in", "all", "total"),
  duplicate = TRUE
)
```

Arguments

- **adjlist**  The adjacency list. It should be consistent, i.e. the maximum throughout all vectors in the list must be less than the number of vectors (i.e., the number of vertices in the graph). Note that the list is expected to be 0-indexed.
- **mode**  Character scalar, it specifies whether the graph to create is undirected ("all" or "total") or directed; and in the latter case, whether it contains the outgoing ("out") or the incoming ("in") neighbors of the vertices.
graph_from_adjacency_matrix

duplicate Logical scalar. For undirected graphs it gives whether edges are included in the list twice. E.g. if it is TRUE then for an undirected \{A, B\} edge graph_from_adj_list expects A included in the neighbors of B and B to be included in the neighbors of A.

This argument is ignored if mode is out or in.

Details

Adjacency lists are handy if you intend to do many (small) modifications to a graph. In this case adjacency lists are more efficient than igraph graphs.

The idea is that you convert your graph to an adjacency list by \texttt{as_adj_list}, do your modifications to the graphs and finally create again an igraph graph by calling \texttt{graph_from_adj_list}.

Value

An igraph graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

\texttt{as_edgelist}

Examples

```r
## Directed
g <- make_ring(10, directed=TRUE)
al <- as_adj_list(g, mode="out")
g2 <- graph_from_adj_list(al)
graph.isomorphic(g, g2)

## Undirected
g <- make_ring(10)
al <- as_adj_list(g)
g2 <- graph_from_adj_list(al, mode="all")
graph.isomorphic(g, g2)
ecount(g2)
g3 <- graph_from_adj_list(al, mode="all", duplicate=FALSE)
ecount(g3)
which_multiple(g3)
```

graph_from_adjacency_matrix

Create graphs from adjacency matrices

Description

\texttt{graph_from_adjacency_matrix} is a flexible function for creating \texttt{igraph} graphs from adjacency matrices.
Usage

```r
graph_from_adjacency_matrix(
  adjmatrix, 
  mode = c("directed", "undirected", "max", "min", "upper", "lower", "plus"), 
  weighted = NULL, 
  diag = TRUE, 
  add.colnames = NULL, 
  add.rownames = NA 
)
```

`from_adjacency(...)`

Arguments

- **adjmatrix**: A square adjacency matrix. From igraph version 0.5.1 this can be a sparse matrix created with the `Matrix` package.
- **mode**: Character scalar, specifies how igraph should interpret the supplied matrix. See also the `weighted` argument, the interpretation depends on that too. Possible values are: `directed`, `undirected`, `upper`, `lower`, `max`, `min`, `plus`. See details below.
- **weighted**: This argument specifies whether to create a weighted graph from an adjacency matrix. If it is `NULL` then an unweighted graph is created and the elements of the adjacency matrix gives the number of edges between the vertices. If it is a character constant then for every non-zero matrix entry an edge is created and the value of the entry is added as an edge attribute named by the `weighted` argument. If it is `TRUE` then a weighted graph is created and the name of the edge attribute will be `weight`. See also details below.
- **diag**: Logical scalar, whether to include the diagonal of the matrix in the calculation. If this is `FALSE` then the diagonal is zeroed out first.
- **add.colnames**: Character scalar, whether to add the column names as vertex attributes. If it is `NULL` (the default) then, if present, column names are added as vertex attribute `name`. If `NA` then they will not be added. If a character constant, then it gives the name of the vertex attribute to add.
- **add.rownames**: Character scalar, whether to add the row names as vertex attributes. Possible values the same as the previous argument. By default row names are not added. If `add.rownames` and `add.colnames` specify the same vertex attribute, then the former is ignored.
- **...**: Passed to `graph_from_adjacency_matrix`.

Details

The order of the vertices are preserved, i.e. the vertex corresponding to the first row will be vertex 0 in the graph, etc.

`graph_from_adjacency_matrix` operates in two main modes, depending on the `weighted` argument.

If this argument is `NULL` then an unweighted graph is created and an element of the adjacency matrix gives the number of edges to create between the two corresponding vertices. The details depend on the value of the `mode` argument:

"directed" The graph will be directed and a matrix element gives the number of edges between two vertices.
"undirected" This is exactly the same as max, for convenience. Note that it is not checked whether the matrix is symmetric.
"max" An undirected graph will be created and \( \max(A(i,j), A(j,i)) \) gives the number of edges.
"upper" An undirected graph will be created, only the upper right triangle (including the diagonal) is used for the number of edges.
"lower" An undirected graph will be created, only the lower left triangle (including the diagonal) is used for creating the edges.
"min" An undirected graph will be created with \( \min(A(i,j), A(j,i)) \) edges between vertex i and j.
"plus" An undirected graph will be created with \( A(i,j) + A(j,i) \) edges between vertex i and j.

If the weighted argument is not NULL then the elements of the matrix give the weights of the edges (if they are not zero). The details depend on the value of the mode argument:

"directed" The graph will be directed and a matrix element gives the edge weights.
"undirected" First we check that the matrix is symmetric. It is an error if not. Then only the upper triangle is used to create a weighted undirected graph.
"max" An undirected graph will be created and \( \max(A(i,j), A(j,i)) \) gives the edge weights.
"upper" An undirected graph will be created, only the upper right triangle (including the diagonal) is used (for the edge weights).
"lower" An undirected graph will be created, only the lower left triangle (including the diagonal) is used for creating the edges.
"min" An undirected graph will be created, \( \min(A(i,j), A(j,i)) \) gives the edge weights.
"plus" An undirected graph will be created, \( A(i,j) + A(j,i) \) gives the edge weights.

Value
An igraph graph object.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

See Also
graph and graph_from_literal for other ways to create graphs.

Examples
```
adjm <- matrix(sample(0:1, 100, replace=TRUE, prob=c(0.9,0.1)), ncol=10)
g1 <- graph_from_adjacency_matrix(adjm)
adjm <- matrix(sample(0:5, 100, replace=TRUE, prob=c(0.9,0.02,0.02,0.02,0.02,0.02)), ncol=10)
g2 <- graph_from_adjacency_matrix(adjm, weighted=TRUE, mode="undirected")
E(g2)$weight
```

```
nzs <- function(x) sort(x[!x==0])
adjm <- matrix(runif(100), 10)
adjm[ adjm<0.5 ] <- 0
g3 <- graph_from_adjacency_matrix((adjm + t(adjm))/2, weighted=TRUE, mode="undirected")
```
g4 <- graph_from_adjacency_matrix(adjm, weighted=TRUE, mode="max")
all(nzs(pmax(adjm, t(adjm))[upper.tri(adjm)]) == sort(E(g4)$weight))

g5 <- graph_from_adjacency_matrix(adjm, weighted=TRUE, mode="min")
all(nzs(pmin(adjm, t(adjm))[upper.tri(adjm)]) == sort(E(g5)$weight))

g6 <- graph_from_adjacency_matrix(adjm, weighted=TRUE, mode="upper")
all(nzs(adjm[upper.tri(adjm)]) == sort(E(g6)$weight))

g7 <- graph_from_adjacency_matrix(adjm, weighted=TRUE, mode="lower")
all(nzs(adjm[lower.tri(adjm)]) == sort(E(g7)$weight))

g8 <- graph_from_adjacency_matrix(adjm, weighted=TRUE, mode="plus")
d2 <- function(x) { diag(x) <- diag(x)/2; x }
all(nzs((d2(adjm+t(adjm)))[lower.tri(adjm)]) == sort(E(g8)$weight))

g9 <- graph_from_adjacency_matrix(adjm, weighted=TRUE, mode="plus", diag=FALSE)
d0 <- function(x) { diag(x) <- 0 }
all(nzs((d0(adjm+t(adjm)))[lower.tri(adjm)]) == sort(E(g9)$weight))

## row/column names
rownames(adjm) <- sample(letters, nrow(adjm))
colnames(adjm) <- seq(ncol(adjm))
g10 <- graph_from_adjacency_matrix(adjm, weighted=TRUE, add.rownames="code")
summary(g10)

---

**graph_from_atlas**

Create a graph from the Graph Atlas

**Description**

`graph_from_atlas` creates graphs from the book ‘An Atlas of Graphs’ by Roland C. Read and Robin J. Wilson. The atlas contains all undirected graphs with up to seven vertices, numbered from 0 up to 1252. The graphs are listed:

1. in increasing order of number of nodes;
2. for a fixed number of nodes, in increasing order of the number of edges;
3. for fixed numbers of nodes and edges, in increasing order of the degree sequence, for example 111223 < 112222;
4. for fixed degree sequence, in increasing number of automorphisms.

**Usage**

`graph_from_atlas(n)`

`atlas(...)`

**Arguments**

- `n` The id of the graph to create.
- `...` Passed to `graph_from_atlas`. 
Value

An igraph graph.

See Also

Other deterministic constructors: graph_from_edgelist(), graph_from_literal(), make_chordal_ring(), make_empty_graph(), make_full_citation_graph(), make_full_graph(), make_graph(), make_lattice(), make_ring(), make_star(), make_tree()

Examples

```r
## Some randomly picked graphs from the atlas
graph_from_atlas(sample(0:1252, 1))
graph_from_atlas(sample(0:1252, 1))
```

graph_from_edgelist Create a graph from an edge list matrix

Description

graph_from_edgelist creates a graph from an edge list. Its argument is a two-column matrix, each row defines one edge. If it is a numeric matrix then its elements are interpreted as vertex ids. If it is a character matrix then it is interpreted as symbolic vertex names and a vertex id will be assigned to each name, and also a name vertex attribute will be added.

Usage

```r
graph_from_edgelist(el, directed = TRUE)
from_edgelist(...)```

Arguments

el The edge list, a two column matrix, character or numeric.
directed Whether to create a directed graph.
... Passed to graph_from_edgelist.

Value

An igraph graph.

See Also

Other deterministic constructors: graph_from_atlas(), graph_from_literal(), make_chordal_ring(), make_empty_graph(), make_full_citation_graph(), make_full_graph(), make_graph(), make_lattice(), make_ring(), make_star(), make_tree()
Examples

```r
e1 <- matrix(c("foo", "bar", "bar", "foobar"), nc = 2, byrow = TRUE)
graph_from_edgelist(e1)

# Create a ring by hand
graph_from_edgelist(cbind(1:10, c(2:10, 1)))
```

Description

This function downloads a graph from a database created for the evaluation of graph isomorphism testing algorithms.

Usage

```r
graph_from_graphdb(
  url = NULL,
  prefix = "iso",
  type = "r001",
  nodes = NULL,
  pair = "A",
  which = 0,
  base = "http://cneurocvs.rmki.kfki.hu/graphdb/gzip",
  compressed = TRUE,
  directed = TRUE
)
```

Arguments

- `url`  If not NULL it is a complete URL with the file to import.
- `prefix`  Gives the prefix. See details below. Possible values: iso, i2, si4, si6, mcs10, mcs30, mcs50, mcs70, mcs90.
- `type`  Gives the graph type identifier. See details below. Possible values: r001, r005, r01, r02, m2D, m2Dr2, m2Dr4, m2Dr6, m3D, m3Dr2, m3Dr4, m3Dr6, m4D, m4Dr2, m4Dr4, m4Dr6, b03, b03m, b06, b06m, b09, b09m.
- `nodes`  The number of vertices in the graph.
- `pair`  Specifies which graph of the pair to read. Possible values: A and B.
- `which`  Gives the number of the graph to read. For every graph type there are a number of actual graphs in the database. This argument specifies which one to read.
- `base`  The base address of the database. See details below.
- `compressed`  Logical constant, if TRUE than the file is expected to be compressed by gzip. If `url` is NULL then a ‘.gz’ suffix is added to the filename.
- `directed`  Logical constant, whether to create a directed graph.
graph_from_graphdb reads a graph from the graph database from an FTP or HTTP server or from a local copy. It has two modes of operation:

If the url argument is specified then it should the complete path to a local or remote graph database file. In this case we simply call read_graph with the proper arguments to read the file.

If url is NULL, and this is the default, then the filename is assembled from the base, prefix, type, nodes, pair and which arguments.

Unfortunately the original graph database homepage is now defunct, but see its old version at http://web.archive.org/web/20090215182331/http://amalfi.dis.unina.it/graph/db/doc/graphdbat.html for the actual format of a graph database file and other information.

Value

A new graph object.

Examples

g <- graph_from_graphdb(prefix="iso", type="r001", nodes=20, pair="A", which=10, compressed=TRUE)
g2 <- graph_from_graphdb(prefix="iso", type="r001", nodes=20, pair="B", which=10, compressed=TRUE)
graph.isomorphic.vf2(g, g2) % should be TRUE
g3 <- graph_from_graphdb(url=paste(sep="/",
  "http://cneurocvs.rmki.kfki.hu",
  "graphdb/gzip/iso/bvg/b06m",
  "iso_b06m_m200.A09.gz"))

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

read_graph, graph.isomorphic.vf2

graph_from_graphnel

Convert graphNEL objects from the graph package to igraph

Description

The graphNEL class is defined in the graph package, it is another way to represent graphs. graph_from_graphnel takes a graphNEL graph and converts it to an igraph graph. It handles all graph/vertex/edge attributes. If the graphNEL graph has a vertex attribute called ‘name’ it will be used as igraph vertex attribute ‘name’ and the graphNEL vertex names will be ignored.
Usage

\texttt{graph_from_graphnel(graphNEL, name = TRUE, weight = TRUE, unlist.attrs = TRUE)}

Arguments

\begin{description}
\item \texttt{graphNEL} The graphNEL graph.
\item \texttt{name} Logical scalar, whether to add graphNEL vertex names as an igraph vertex attribute called \texttt{‘name’}.
\item \texttt{weight} Logical scalar, whether to add graphNEL edge weights as an igraph edge attribute called \texttt{‘weight’}. (graphNEL graphs are always weighted.)
\item \texttt{unlist.attrs} Logical scalar. graphNEL attribute query functions return the values of the attributes in R lists, if this argument is \texttt{TRUE} (the default) these will be converted to atomic vectors, whenever possible, before adding them to the igraph graph.
\end{description}

Details

Because graphNEL graphs poorly support multiple edges, the edge attributes of the multiple edges are lost: they are all replaced by the attributes of the first of the multiple edges.

Value

\texttt{graph_from_graphnel} returns an igraph graph object.

See Also

\texttt{as_graphnel} for the other direction, \texttt{as_adj, graph_from_adjacency_matrix}, \texttt{as_adj_list} and \texttt{graph.adjlist} for other graph representations.

Examples

\begin{verbatim}
## Not run:
## Undirected
g <- make_ring(10)
V(g)$name <- letters[1:10]
GNEL <- as_graphnel(g)
g2 <- graph_from_graphnel(GNEL)
g2

## Directed
g3 <- make_star(10, mode="in")
V(g3)$name <- letters[1:10]
GNEL2 <- as_graphnel(g3)
g4 <- graph_from_graphnel(GNEL2)
g4

## End(Not run)
\end{verbatim}
graph_from_incidence_matrix

Create graphs from an incidence matrix

Description
graph_from_incidence_matrix creates a bipartite igraph graph from an incidence matrix.

Usage
graph_from_incidence_matrix(
  incidence, 
  directed = FALSE, 
  mode = c("all", "out", "in", "total"), 
  multiple = FALSE, 
  weighted = NULL, 
  add.names = NULL 
)

deprecated
from_incidence_matrix(....)

Arguments

incidence The input incidence matrix. It can also be a sparse matrix from the Matrix package.
directed Logical scalar, whether to create a directed graph.
mode A character constant, defines the direction of the edges in directed graphs, ignored for undirected graphs. If ‘out’, then edges go from vertices of the first kind (corresponding to rows in the incidence matrix) to vertices of the second kind (columns in the incidence matrix). If ‘in’, then the opposite direction is used. If ‘all’ or ‘total’, then mutual edges are created.
multiple Logical scalar, specifies how to interpret the matrix elements. See details below.
weighted This argument specifies whether to create a weighted graph from the incidence matrix. If it is NULL then an unweighted graph is created and the multiple argument is used to determine the edges of the graph. If it is a character constant then for every non-zero matrix entry an edge is created and the value of the entry is added as an edge attribute named by the weighted argument. If it is TRUE then a weighted graph is created and the name of the edge attribute will be ‘weight’.
add.names A character constant, NA or NULL. graph_from_incidence_matrix can add the row and column names of the incidence matrix as vertex attributes. If this argument is NULL (the default) and the incidence matrix has both row and column names, then these are added as the ‘name’ vertex attribute. If you want a different vertex attribute for this, then give the name of the attributes as a character string. If this argument is NA, then no vertex attributes (other than type) will be added.

... Passed to graph_from_incidence_matrix.
Bipartite graphs have a ‘type’ vertex attribute in igraph, this is boolean and FALSE for the vertices of the first kind and TRUE for vertices of the second kind.

graph_from_incidence_matrix can operate in two modes, depending on the multiple argument. If it is FALSE then a single edge is created for every non-zero element in the incidence matrix. If multiple is TRUE, then the matrix elements are rounded up to the closest non-negative integer to get the number of edges to create between a pair of vertices.

A bipartite igraph graph. In other words, an igraph graph that has a vertex attribute type.

Gabor Csardi <csardi.gabor@gmail.com>

make_bipartite_graph for another way to create bipartite graphs

inc <- matrix(sample(0:1, 15, repl=TRUE), 3, 5)
colnames(inc) <- letters[1:5]
rownames(inc) <- LETTERS[1:3]
graph_from_incidence_matrix(inc)

description The isomorphism class is a non-negative integer number. Graphs (with the same number of vertices) having the same isomorphism class are isomorphic and isomorphic graphs always have the same isomorphism class. Currently it can handle directed graphs with 3 or 4 vertices and undirected graphs with 3 to 6 vertices.

Usage

graph_from_isomorphism_class(size, number, directed = TRUE)

Arguments

size The number of vertices in the graph.
number The isomorphism class.
directed Whether to create a directed graph (the default).

Value

An igraph object, the graph of the given size, directedness and isomorphism class.
Creating a graph from LCF notation

Description

LCF is short for Lederberg-Coxeter-Frucht, it is a concise notation for 3-regular Hamiltonian graphs. It consists of three parameters, the number of vertices in the graph, a list of shifts giving additional edges to a cycle backbone and another integer giving how many times the shifts should be performed. See http://mathworld.wolfram.com/LCFNotation.html for details.

Usage

graph_from_lcf(n, shifts, repeats = 1)

Arguments

n Integer, the number of vertices in the graph.
shifts Integer vector, the shifts.
repeats Integer constant, how many times to repeat the shifts.

Value

A graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

graph can create arbitrary graphs, see also the other functions on the its manual page for creating special graphs.

Examples

# This is the Franklin graph:
g1 <- graph_from_lcf(12, c(5,-5), 6)
g2 <- make_graph("Franklin")
isomorphic(g1, g2)
Creating (small) graphs via a simple interface

This function is useful if you want to create a small (named) graph quickly, it works for both directed and undirected graphs.

Usage

\[
\text{graph_from_literal}(\ldots, \text{simplify} = \text{TRUE})
\]
\[
\text{from_literal}(\ldots)
\]

Arguments

\ldots 
For \text{graph_from_literal} the formulae giving the structure of the graph, see details below. For \text{from_literal} all arguments are passed to \text{graph_from_literal}.

\text{simplify} 
Logical scalar, whether to call \text{simplify} on the created graph. By default the graph is simplified, loop and multiple edges are removed.

Details

\text{graph_from_literal} is very handy for creating small graphs quickly. You need to supply one or more R expressions giving the structure of the graph. The expressions consist of vertex names and edge operators. An edge operator is a sequence of ‘-’ and ‘+’ characters, the former is for the edges and the latter is used for arrow heads. The edges can be arbitrarily long, ie. you may use as many ‘-’ characters to “draw” them as you like.

If all edge operators consist of only ‘-’ characters then the graph will be undirected, whereas a single ‘+’ character implies a directed graph.

Let us see some simple examples. Without arguments the function creates an empty graph:

\[
\text{graph_from_literal}()
\]

A simple undirected graph with two vertices called 'A' and 'B' and one edge only:

\[
\text{graph_from_literal}(A\text{-}B)
\]

Remember that the length of the edges does not matter, so we could have written the following, this creates the same graph:

\[
\text{graph_from_literal}( A\text{-----}B )
\]

If you have many disconnected components in the graph, separate them with commas. You can also give isolate vertices.

\[
\text{graph_from_literal}( A\text{--}B, C\text{--}D, E\text{--}F, G\text{--}H, I, J, K )
\]

The ‘:’ operator can be used to define vertex sets. If an edge operator connects two vertex sets then every vertex from the first set will be connected to every vertex in the second set. The following form creates a full graph, including loop edges:
`graph_from_literal( A:B:C:D -- A:B:C:D )`

In directed graphs, edges will be created only if the edge operator includes a arrow head (`'+'`) at the end of the edge:

`graph_from_literal( A --> B --> C )`
`graph_from_literal( A --> B + C )`
`graph_from_literal( A + B -- C )`

Thus in the third example no edge is created between vertices `B` and `C`.
Mutual edges can be also created with a simple edge operator:

`graph_from_literal( A +-- B +-- C +-- D + E)`

Note again that the length of the edge operators is arbitrary. `'+', '++' and '+++++'` have exactly the same meaning.
If the vertex names include spaces or other special characters then you need to quote them:

`graph_from_literal( "this is" + "a silly" + "graph here" )`

You can include any character in the vertex names this way, even `'+` and `'-` characters.
See more examples below.

**Value**

An igraph graph

**See Also**

Other deterministic constructors: `graph_from_atlas()`, `graph_from_edgelist()`, `make_chordal_ring()`, `make_empty_graph()`, `make_full_citation_graph()`, `make_full_graph()`, `make_graph()`, `make_lattice()`, `make_ring()`, `make_star()`, `make_tree()`

**Examples**

# A simple undirected graph
g <- graph_from_literal( Alice-Bob-Cecil-Alice, Daniel-Cecil-Eugene, Cecil-Gordon )
g

# Another undirected graph, "":" notation
g2 <- graph_from_literal( Alice-Bob:Cecil:Daniel, Cecil:Daniel-Eugene:Gordon )
g2

# A directed graph
g3 <- graph_from_literal( Alice --> Bob --> Cecil --> Daniel, Eugene --> Gordon:Helen )
g3

# A graph with isolate vertices
g4 <- graph_from_literal( Alice -- Bob -- Daniel, Cecil:Gordon, Helen )
g4 V(g4)$name
**graph_id**

Get the id of a graph

---

**Description**

Graph ids are used to check that a vertex or edge sequence belongs to a graph. If you create a new graph by changing the structure of a graph, the new graph will have a new id. Changing the attributes will not change the id.

**Usage**

```r
graph_id(x, ...)```

**Arguments**

- `x`: A graph or a vertex sequence or an edge sequence.
- `...`: Not used currently.

**Value**

The id of the graph, a character scalar. For vertex and edge sequences the id of the graph they were created from.

**Examples**

```r
g <- make_ring(10)
graph_id(g)
graph_id(V(g))
graph_id(E(g))
g2 <- g + 1
graph_id(g2)
```
**graph_version**

**igraph data structure versions**

**Description**

igraph’s internal data representation changes sometimes between versions. This means that it is not possible to use igraph objects that were created (and possibly saved to a file) with an older igraph version.

**Usage**

```r
graph_version(graph)
```

**Arguments**

- `graph` The input graph. If it is missing, then the version number of the current data format is returned.

**Details**

`graph_version` queries the current data format, or the data format of a possibly older igraph graph.

`upgrade_graph` can convert an older data format to the current one.

**Value**

A character scalar.

**See Also**

`upgrade_graph` to convert the data format of a graph.

**graphlet_basis**

**Graphlet decomposition of a graph**

**Description**

Graphlet decomposition models a weighted undirected graph via the union of potentially overlapping dense social groups. This is done by a two-step algorithm. In the first step a candidate set of groups (a candidate basis) is created by finding cliques if the thresholded input graph. In the second step these the graph is projected on the candidate basis, resulting a weight coefficient for each clique in the candidate basis.
Usage

graphlet_basis(graph, weights = NULL)

graphlet_proj(
  graph,
  weights = NULL,
  cliques,
  niter = 1000,
  Mu = rep(1, length(cliques))
)

Arguments

- **graph**: The input graph, edge directions are ignored. Only simple graph (i.e. graphs without self-loops and multiple edges) are supported.
- **weights**: Edge weights. If the graph has a weight edge attribute and this argument is NULL (the default), then the weight edge attribute is used.
- **cliques**: A list of vertex ids, the graphlet basis to use for the projection.
- **niter**: Integer scalar, the number of iterations to perform.
- **Mu**: Starting weights for the projection.

Details

igraph contains three functions for performing the graph decomposition of a graph. The first is `graphlets`, which performed both steps on the method and returns a list of subgraphs, with their corresponding weights. The second and third functions correspond to the first and second steps of the algorithm, and they are useful if the user wishes to perform them individually: `graphlet_basis` and `graphlet_proj`.

Value

- **graphlets**: returns a list with two members:
  - **cliques**: A list of subgraphs, the candidate graphlet basis. Each subgraph is give by a vector of vertex ids.
  - **Mu**: The weights of the subgraphs in graphlet basis.
- **graphlet_basis**: returns a list of two elements:
  - **cliques**: A list of subgraphs, the candidate graphlet basis. Each subgraph is give by a vector of vertex ids.
  - **thresholds**: The weight thresholds used for finding the subgraphs.
- **graphlet_proj**: return a numeric vector, the weights of the graphlet basis subgraphs.

Examples

```r
## Create an example graph first
D1 <- matrix(0, 5, 5)
D2 <- matrix(0, 5, 5)
D3 <- matrix(0, 5, 5)
D1[1:3, 1:3] <- 2
D2[3:5, 3:5] <- 3
```
D3[2:5, 2:5] <- 1

g <- simplify(graph_from_adjacency_matrix(D1 + D2 + D3, mode="undirected", weighted=TRUE))

V(g)$color <- "white"
E(g)$label <- E(g)$weight
E(g)$label.cex <- 2
E(g)$color <- "black"

layout(matrix(1:6, nrow=2, byrow=TRUE))

c <- layout_with_kk(g)

par(mar=c(1,1,1,1))

plot(g, layout=co)

## Calculate graphlets

gl <- graphlets(g, niter=1000)

## Plot graphlets

for (i in 1:length(gl$cliques)) {
  sel <- gl$cliques[[i]]
  V(g)$color <- "white"
  V(g)[sel]$color <- "#E495A5"
  E(g)$width <- 1
  E(g)[ V(g)[sel] %--% V(g)[sel] ]$width <- 2
  E(g)$label <- ""
  E(g)[ width == 2 ]$label <- round(gl$Mu[i], 2)
  E(g)$color <- "black"
  E(g)[ width == 2 ]$color <- "#E495A5"

  plot(g, layout=co)
}

---

greedy_vertex_coloring

Greedy vertex coloring

Description

greedy_vertex_coloring finds a coloring for the vertices of a graph based on a simple greedy algorithm.

Usage

greedy_vertex_coloring(graph, heuristic = c("colored_neighbors"))

Arguments

graph The graph object to color
heuristic The selection heuristic for the next vertex to consider. Currently only one heuristic is supported: "colored_neighbors" selects the vertex with the largest number of already colored neighbors.
groups

Details

The goal of vertex coloring is to assign a "color" (i.e. a positive integer index) to each vertex of the graph such that neighboring vertices never have the same color. This function solves the problem by considering the vertices one by one according to a heuristic, always choosing the smallest color index that differs from that of already colored neighbors. The coloring obtained this way is not necessarily minimal but it can be calculated in linear time.

Value

A numeric vector where item i contains the color index associated to vertex i

Examples

```r
g <- make_graph("petersen")
col <- greedy_vertex_coloring(g)
plot(g, vertex.color=col)
```

Description

Create a list of vertex groups from some graph clustering or community structure.

Usage

`groups(x)`

Arguments

- `x` Some object that represents a grouping of the vertices. See details below.

Details

Currently two methods are defined for this function. The default method works on the output of `components`. (In fact it works on any object that is a list with an entry called `membership`.) The second method works on `communities` objects.

Value

A named list of numeric or character vectors. The names are just numbers that refer to the groups. The vectors themselves are numeric or symbolic vertex ids.

See Also

`components` and the various community finding functions.
Examples

```r
g <- make_graph("Zachary")
fgc <- cluster_fast_greedy(g)
groups(fgc)

g2 <- make_ring(10) + make_full_graph(5)
groups(components(g2))
```

---

gsize

The size of the graph (number of edges)

Description

ecount is an alias of this function.

Usage

gsize(graph)

Arguments

graph The graph.

Value

Numeric scalar, the number of edges.

See Also

Other structural queries: \[\text{.igraph()}.\][\[\text{.igraph()}.\][\text{.igraph()}.\]. adjacent_vertices(). are_adjacent().
ends(). get.edge.ids(). gorder(). head_of(). incident_edges(). incident(). is_directed().
neighbors().tail_of()

Examples

```r
g <- sample_gnp(100, 2/100)
gsize(g)
ecount(g)

# Number of edges in a G(n,p) graph
replicate(100, sample_gnp(10, 1/2), simplify = FALSE) %>%
vapply(gsize, 0) %>%
hist()
```
Harmonic centrality

Description

The harmonic centrality of a vertex is the mean inverse distance to all other vertices. The inverse distance to an unreachable vertex is considered to be zero.

Usage

harmonic_centrality(
  graph,
  vids = V(graph),
  mode = c("out", "in", "all", "total"),
  weights = NULL,
  normalized = FALSE,
  cutoff = -1
)

Arguments

graph The graph to analyze.
vids The vertices for which harmonic centrality will be calculated.
mode Character string, defining the types of the paths used for measuring the distance in directed graphs. "out" follows paths along the edge directions only, "in" traverses the edges in reverse, while "all" ignores edge directions. This argument is ignored for undirected graphs.
weights Optional positive weight vector for calculating weighted harmonic centrality. If the graph has a weight edge attribute, then this is used by default. Weights are used for calculating weighted shortest paths, so they are interpreted as distances.
normalized Logical scalar, whether to calculate the normalized harmonic centrality. If true, the result is the mean inverse path length to other vertices, i.e. it is normalized by the number of vertices minus one. If false, the result is the sum of inverse path lengths to other vertices.
cutoff The maximum path length to consider when calculating the harmonic centrality. There is no such limit when the cutoff is negative. Note that zero cutoff means that only paths of at most length 0 are considered.

details

The cutoff argument can be used to restrict the calculation to paths of length cutoff or smaller only; this can be used for larger graphs to speed up the calculation. If cutoff is negative (which is the default), then the function calculates the exact harmonic centrality scores.

Value

Numeric vector with the harmonic centrality scores of all the vertices in v.

References

has_eulerian_path

See Also

betweenness, closeness

Examples

g <- make_ring(10)
g2 <- make_star(10)
harmonic_centrality(g)
harmonic_centrality(g2, mode="in")
harmonic_centrality(g2, mode="out")
harmonic_centrality(g %du% make_full_graph(5), mode="all")

Description

has_eulerian_path and has_eulerian_cycle checks whether there is an Eulerian path or cycle in the input graph. eulerian_path and eulerian_cycle return such a path or cycle if it exists, and throws an error otherwise.

Usage

has_eulerian_path(graph)
has_eulerian_cycle(graph)
eulerian_path(graph)
eulerian_cycle(graph)

Arguments

graph An igraph graph object

Details

has_eulerian_path decides whether the input graph has an Eulerian path, i.e. a path that passes through every edge of the graph exactly once, and returns a logical value as a result. eulerian_path returns a possible Eulerian path, described with its edge and vertex sequence, or throws an error if no such path exists.

has_eulerian_cycle decides whether the input graph has an Eulerian cycle, i.e. a path that passes through every edge of the graph exactly once and that returns to its starting point, and returns a logical value as a result. eulerian_cycle returns a possible Eulerian cycle, described with its edge and vertex sequence, or throws an error if no such cycle exists.
**has_eulerian_path and has_eulerian_cycle**

For `has_eulerian_path` and `has_eulerian_cycle`, a logical value that indicates whether the graph contains an Eulerian path or cycle. For `eulerian_path` and `eulerian_cycle`, a named list with two entries:

- `epath`: A vector containing the edge ids along the Eulerian path or cycle.
- `vpath`: A vector containing the vertex ids along the Eulerian path or cycle.

**Examples**

```r
library(igraph)

# Create an Eulerian graph
g <- make_graph(~ A-B-C-D-E-A-F-D-B-F-E)

# Check for Eulerian path and cycle
has_eulerian_path(g)
eulerian_path(g)

# Check for Eulerian cycle
has_eulerian_cycle(g)
# Not run: eulerian_cycle(g)
```

---

**head_of**

**Head of the edge(s) in a graph**

**Description**

For undirected graphs, head and tail is not defined. In this case `head_of` returns vertices incident to the supplied edges, and `tail_of` returns the other end(s) of the edge(s).

**Usage**

`head_of(graph, es)`

**Arguments**

- `graph`: The input graph.
- `es`: The edges to query.

**Value**

A vertex sequence with the head(s) of the edge(s).

**See Also**

Other structural queries: `.[igraph]`, `[[.igraph]`, `adjacent_vertices`, `are_adjacent`, `ends`, `get.edge.ids`, `gorder`, `gsize`, `incident_edges`, `incident`, `is_directed`, `neighbors`, `tail_of`
head_print  \hspace{1cm} \textit{Print the only the head of an R object}

\textbf{Description}

Print the only the head of an R object

\textbf{Usage}

\begin{verbatim}
head_print(
x, 
  max_lines = 20,
  header = "",
  footer = "",
  omitted_footer = "",
  ...
)
\end{verbatim}

\textbf{Arguments}

- \texttt{x} \hspace{0.5cm} The object to print, or a callback function. See \texttt{printer_callback} for details.
- \texttt{max_lines} \hspace{0.5cm} Maximum number of lines to print, \textit{not} including the header and the footer.
- \texttt{header} \hspace{0.5cm} The header, if a function, then it will be called, otherwise printed using \texttt{cat}.
- \texttt{footer} \hspace{0.5cm} The footer, if a function, then it will be called, otherwise printed using \texttt{cat}.
- \texttt{omitted_footer} \hspace{0.5cm} Footer that is only printed if anything is omitted from the printout. If a function, then it will be called, otherwise printed using \texttt{cat}.
- \texttt{...} \hspace{0.5cm} Extra arguments to pass to \texttt{print()}. 

\textbf{Value}

\begin{verbatim}
x, invisibly.
\end{verbatim}

hrg \hspace{1cm} \textit{Create a hierarchical random graph from an igraph graph}

\textbf{Description}

\texttt{hrg} creates a HRG from an igraph graph. The igraph graph must be a directed binary tree, with \( n - 1 \) internal and \( n \) leaf vertices. The \texttt{prob} argument contains the HRG probability labels for each vertex; these are ignored for leaf vertices.

\textbf{Usage}

\begin{verbatim}
hrg(graph, prob)
\end{verbatim}

\textbf{Arguments}

- \texttt{graph} \hspace{0.5cm} The igraph graph to create the HRG from.
- \texttt{prob} \hspace{0.5cm} A vector of probabilities, one for each vertex, in the order of vertex ids.
hrg-methods

Value

hrg returns an igraphHRG object.

See Also

Other hierarchical random graph functions: consensus_tree(), fit_hrg(), hrg-methods, hrg_tree(), predict_edges(), print.igraphHRGConsensus(), print.igraphHRG(), sample_hrg()

hrg-methods

Hierarchical random graphs

Description

Fitting and sampling hierarchical random graph models.

Details

A hierarchical random graph is an ensemble of undirected graphs with \( n \) vertices. It is defined via a binary tree with \( n \) leaf and \( n-1 \) internal vertices, where the internal vertices are labeled with probabilities. The probability that two vertices are connected in the random graph is given by the probability label at their closest common ancestor.

Please see references below for more about hierarchical random graphs.

igraph contains functions for fitting HRG models to a given network (fit_hrg, for generating networks from a given HRG ensemble (sample_hrg), converting an igraph graph to a HRG and back (hrg, hrg_tree), for calculating a consensus tree from a set of sampled HRGs (consensus_tree) and for predicting missing edges in a network based on its HRG models (predict_edges).

The igraph HRG implementation is heavily based on the code published by Aaron Clauset, at his website (not functional any more).

See Also

Other hierarchical random graph functions: consensus_tree(), fit_hrg(), hrg_tree(), hrg(), predict_edges(), print.igraphHRGConsensus(), print.igraphHRG(), sample_hrg()

hrg_tree

Create an igraph graph from a hierarchical random graph model

Description

hrg_tree creates the corresponding igraph tree of a hierarchical random graph model.

Usage

hrg_tree(hrg)

Arguments

hrg A hierarchical random graph model.
**hub_score**

**Value**
An igraph graph.

**See Also**
Other hierarchical random graph functions: `consensus_tree()`, `fit_hrg()`, `hrg-methods`, `hrg()`, `predict_edges()`, `print.igraphHRGConsensus()`, `print.igraphHRG()`, `sample_hrg()`

---

**hub_score**

Kleinberg’s hub centrality scores.

**Description**

The hub scores of the vertices are defined as the principal eigenvector of $AA^T$, where $A$ is the adjacency matrix of the graph.

**Usage**

```r
hub_score(graph, scale = TRUE, weights = NULL, options = arpack_defaults)
```

**Arguments**

- `graph` The input graph.
- `scale` Logical scalar, whether to scale the result to have a maximum score of one. If no scaling is used then the result vector has unit length in the Euclidean norm.
- `weights` Optional positive weight vector for calculating weighted scores. If the graph has a weight edge attribute, then this is used by default. This function interprets edge weights as connection strengths. In the random surfer model, an edge with a larger weight is more likely to be selected by the surfer.
- `options` A named list, to override some ARPACK options. See `arpack` for details.

**Details**

For undirected matrices the adjacency matrix is symmetric and the hub scores are the same as authority scores, see `authority_score`.

**Value**

A named list with members:

- `vector` The authority/hub scores of the vertices.
- `value` The corresponding eigenvalue of the calculated principal eigenvector.
- `options` Some information about the ARPACK computation, it has the same members as the `options` member returned by `arpack`, see that for documentation.

**References**

identical_graphs

See Also

authority_score, eigen_centrality for eigenvector centrality, page_rank for the Page Rank scores. arpack for the underlining machinery of the computation.

Examples

```r
## An in-star
g <- make_star(10)
hub_score(g)$vector

## A ring
g2 <- make_ring(10)
hub_score(g2)$vector
```

Description

Two graphs are considered identical by this function if and only if they are represented in exactly the same way in the internal R representation. This means that the two graphs must have the same list of vertices and edges, in exactly the same order, with same directedness, and the two graphs must also have identical graph, vertex and edge attributes.

Usage

```r
identical_graphs(g1, g2, attrs = TRUE)
```

Arguments

- `g1, g2`: The two graphs
- `attrs`: Whether to compare the attributes of the graphs

Details

This is similar to identical in the base package, but it ignores the mutable piece of igraph objects; those might be different even if the two graphs are identical.

Attribute comparison can be turned off with the `attrs` parameter if the attributes of the two graphs are allowed to be different.

Value

Logical scalar
Description

Many times, when the structure of a graph is modified, vertices/edges map of the original graph map to vertices/edges in the newly created (modified) graph. For example `simplify` maps multiple edges to single edges. `igraph` provides a flexible mechanism to specify what to do with the vertex/edge attributes in these cases.

Details

The functions that support the combination of attributes have one or two extra arguments called `vertex.attr.comb` and/or `edge.attr.comb` that specify how to perform the mapping of the attributes. E.g. `contract` contracts many vertices into a single one, the attributes of the vertices can be combined and stores as the vertex attributes of the new graph.

The specification of the combination of (vertex or edge) attributes can be given as

1. a character scalar,
2. a function object or
3. a list of character scalars and/or function objects.

If it is a character scalar, then it refers to one of the predefined combinations, see their list below.

If it is a function, then the given function is expected to perform the combination. It will be called once for each new vertex/edge in the graph, with a single argument: the attribute values of the vertices that map to that single vertex.

The third option, a list can be used to specify different combination methods for different attributes.

A named entry of the list corresponds to the attribute with the same name. An unnamed entry (i.e. if the name is the empty string) of the list specifies the default combination method. I.e.

```
list(weight="sum", "ignore")
```

specifies that the weight of the new edge should be sum of the weights of the corresponding edges in the old graph; and that the rest of the attributes should be ignored (=dropped).

Predefined combination functions

The following combination behaviors are predefined:

"ignore" The attribute is ignored and dropped.

"sum" The sum of the attributes is calculated. This does not work for character attributes and works for complex attributes only if they have a `sum` generic defined. (E.g. it works for sparse matrices from the `Matrix` package, because they have a `sum` method.)

"prod" The product of the attributes is calculated. This does not work for character attributes and works for complex attributes only if they have a `prod` function defined.

"min" The minimum of the attributes is calculated and returned. For character and complex attributes the standard R `min` function is used.

"max" The maximum of the attributes is calculated and returned. For character and complex attributes the standard R `max` function is used.
"random" Chooses one of the supplied attribute values, uniformly randomly. For character and complex attributes this is implemented by calling sample.

"first" Always chooses the first attribute value. It is implemented by calling the head function.

"last" Always chooses the last attribute value. It is implemented by calling the tail function.

"mean" The mean of the attributes is calculated and returned. For character and complex attributes this simply calls the mean function.

"median" The median of the attributes is selected. Calls the R median function for all attribute types.

"concat" Concatenate the attributes, using the c function. This results almost always a complex attribute.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

graph_attr, vertex_attr, edge_attr on how to use graph/vertex/edge attributes in general.
igraph_options on igraph parameters.

Examples

```r
# create a graph with some edge attributes
E(g)weight <- 1:5
g <- graph( c(1,2, 1,2, 1,2, 2,3, 3,4) )

## print attribute values with the graph
igraph_options(print.graph.attributes=TRUE)
igraph_options(print.vertex.attributes=TRUE)
igraph_options(print.edge.attributes=TRUE)

## new attribute is the sum of the old ones
simplify(g, edge.attr.comb="sum")

## collect attributes into a string
simplify(g, edge.attr.comb=toString)

## concatenate them into a vector, this creates a complex attribute
simplify(g, edge.attr.comb="concat")

E(g)$name <- letters[seq_len(ecount(g))]

## both attributes are collected into strings
simplify(g, edge.attr.comb=toString)

## harmonic average of weights, names are dropped
simplify(g, edge.attr.comb=list(weight=function(x) length(x)/sum(1/x),
                            name="ignore"))
```
igraph-dollar

Getting and setting graph attributes, shortcut

Description

The $ operator is a shortcut to get and set graph attributes. It is shorter and just as readable as graph_attr and set_graph_attr.

Usage

## S3 method for class 'igraph'
x$name

## S3 replacement method for class 'igraph'
x$name <- value

Arguments

x An igraph graph
name Name of the attribute to get/set.
value New value of the graph attribute.

See Also

Other graph attributes: delete_edge_attr(), delete_graph_attr(), delete_vertex_attr(), edge_attr<()$, edge_attr_names(), edge_attr(), graph_attr<()$, graph_attr_names(),
graph_attr(), igraph-vs-attributes, set_edge_attr(), set_graph_attr(), set_vertex_attr(),
vertex_attr<()$, vertex_attr_names(), vertex_attr()

Examples

g <- make_ring(10)
g$name
g$name <- "10-ring"
g$name

igraph-es-attributes Query or set attributes of the edges in an edge sequence

Description

The $ operator is a syntactic sugar to query and set edge attributes, for edges in an edge sequence.
## S3 replacement method for class 'igraph.es'

```r
x[[i]] <- value
```

## S3 replacement method for class 'igraph.es'

```r
x[i] <- value
```

## S3 method for class 'igraph.es'

```r
x$name
```

## S3 replacement method for class 'igraph.es'

```r
x$name <- value
```

```r
E(x, path = NULL, P = NULL, directed = NULL) <- value
```

### Arguments

- **x**: An edge sequence. For `E<-` it is a graph.
- **i**: Index.
- **value**: New value of the attribute, for the edges in the edge sequence.
- **name**: Name of the edge attribute to query or set.
- **path**: Select edges along a path, given by a vertex sequence. See `E`.
- **P**: Select edges via pairs of vertices. See `E`.
- **directed**: Whether to use edge directions for the `path` or `P` arguments.

### Details

The query form of `[[` is a shortcut for `edge_attr`, e.g. `E(g)[idx]$attr` is equivalent to `edge_attr(g, attr, E(g)[idx])`.

The assignment form of `[[` is a shortcut for `set_edge_attr`, e.g. `E(g)[idx]$attr <- value` is equivalent to `g <- set_edge_attr(g, attr, E(g)[idx], value)`.

### Value

A vector or list, containing the values of the attribute `name` for the edges in the sequence. For numeric, character or logical attributes, it is a vector of the appropriate type, otherwise it is a list.

### See Also

Other vertex and edge sequences: `E()`, `V()`, `igraph-es-indexing2`, `igraph-es-indexing`, `igraph-vs-attributes`, `igraph-vs-indexing2`, `igraph-vs-indexing`, `print.igraph.es()`, `print.igraph.vs()`

Other vertex and edge sequences: `E()`, `V()`, `igraph-es-indexing2`, `igraph-es-indexing`, `igraph-vs-attributes`, `igraph-vs-indexing2`, `igraph-vs-indexing`, `print.igraph.es()`, `print.igraph.vs()`

### Examples

```r
# color edges of the largest component
largest_comp <- function(graph) {
  cl <- components(graph)
  V(graph)[which.max(cl$csize) == cl$membership]
}
```
g <- sample_(gnp(100, 1/100),
  with_vertex_(size = 3, label = ""),
  with_graph_(layout = layout_with_fr)
)
giant_v <- largest_comp(g)
E(g)$color <- "orange"
E(g)[giant_v %--% giant_v]$color <- "blue"
plot(g)

igraph-es-indexing  Indexing edge sequences

Description

Edge sequences can be indexed very much like a plain numeric R vector, with some extras.

Usage

## S3 method for class 'igraph.es'
x[...]

Arguments

x          An edge sequence

...        Indices, see details below.

Value

Another edge sequence, referring to the same graph.

Multiple indices

When using multiple indices within the bracket, all of them are evaluated independently, and then
the results are concatenated using the c() function. E.g. E(g)[1, 2, .inc(1)] is equivalent to
c(E(g)[1], E(g)[2], E(g)[.inc(1)]).

Index types

Edge sequences can be indexed with positive numeric vectors, negative numeric vectors, logical
vectors, character vectors:

- When indexed with positive numeric vectors, the edges at the given positions in the sequence
  are selected. This is the same as indexing a regular R atomic vector with positive numeric
  vectors.
- When indexed with negative numeric vectors, the edges at the given positions in the sequence
  are omitted. Again, this is the same as indexing a regular R atomic vector.
- When indexed with a logical vector, the lengths of the edge sequence and the index must
  match, and the edges for which the index is TRUE are selected.
- Named graphs can be indexed with character vectors, to select edges with the given names.
  Note that a graph may have edge names and vertex names, and both can be used to select
  edges. Edge names are simply used as names of the numeric edge id vector. Vertex names
effectively only work in graphs without multiple edges, and must be separated with a | bar
character to select an edges that incident to the two given vertices. See examples below.
Edge attributes

When indexing edge sequences, edge attributes can be referred to simply by using their names. E.g. if a graph has a `weight` edge attribute, then \( E(G)[\text{weight} > 1] \) selects all edges with a weight larger than one. See more examples below. Note that attribute names mask the names of variables present in the calling environment; if you need to look up a variable and you do not want a similarly named edge attribute to mask it, use the `.env` pronoun to perform the name lookup in the calling environment. In other words, use \( E(g)[.env\$weight > 1] \) to make sure that `weight` is looked up from the calling environment even if there is an edge attribute with the same name. Similarly, you can use `.data` to match attribute names only.

Special functions

There are some special igraph functions that can be used only in expressions indexing edge sequences:

- `.inc` takes a vertex sequence, and selects all edges that have at least one incident vertex in the vertex sequence.
- `.from` similar to `.inc()`, but only the tails of the edges are considered.
- `.to` is similar to `.inc()`, but only the heads of the edges are considered.
- `%--%` a special operator that can be used to select all edges between two sets of vertices. It ignores the edge directions in directed graphs.
- `%->%` similar to `%--%`, but edges from the left hand side argument, pointing to the right hand side argument, are selected, in directed graphs.
- `%<-%` similar to `%--%`, but edges to the left hand side argument, pointing from the right hand side argument, are selected, in directed graphs.

Note that multiple special functions can be used together, or with regular indices, and then their results are concatenated. See more examples below.

See Also

Other vertex and edge sequences: `E()`, `V()`, `igraph-es-attributes`, `igraph-es-indexing2`, `igraph-vs-attributes`, `igraph-vs-indexing2`, `igraph-vs-indexing`, `print.igraph.es()`, `print.igraph.vs()`

Other vertex and edge sequence operations: `c.igraph.es()`, `c.igraph.vs()`, `difference.igraph.es()`, `difference.igraph.vs()`, `igraph-es-indexing2`, `igraph-vs-indexing2`, `igraph-vs-indexing`, `intersection.igraph.es()`, `intersection.igraph.vs()`, `rev.igraph.es()`, `rev.igraph.vs()`, `union.igraph.es()`, `union.igraph.vs()`, `unique.igraph.es()`, `unique.igraph.vs()`

Examples

```r
# Special operators for indexing based on graph structure

# The edges along the diameter

g <- sample_pa(100, power = 0.3)
d <- get_diameter(g)
E(g, path = d)
```

# Select edges based on attributes

```r
g <- sample_gnp(20, 3/20) 
  set_edge_attr("weight", value = rnorm(gsize(.))) 
E(g)[[ weight < 0 ]] 
```

# Indexing with a variable whose name matches the name of an attribute 
# may fail; use .env to force the name lookup in the parent environment

```r
E(g)$x <- E(g)$weight
x <- 2
E(g)[.env$x]
```

---

**igraph-es-indexing2**  
Select edges and show their metadata

---

### Description

The double bracket operator can be used on edge sequences, to print the meta-data (edge attributes) of the edges in the sequence.

### Usage

```r
## S3 method for class 'igraph.es'
x[[...]]
```

### Arguments

- `x`: An edge sequence.
- `...`: Additional arguments, passed to `.[]`.  

### Details

Technically, when used with edge sequences, the double bracket operator does exactly the same as the single bracket operator, but the resulting edge sequence is printed differently: all attributes of the edges in the sequence are printed as well.

See `[].igraph.es` for more about indexing edge sequences.

### Value

Another edge sequence, with metadata printing turned on. See details below.

### See Also

Other vertex and edge sequences: `E()`, `V()`, `igraph-es-attributes`, `igraph-es-indexing`, `igraph-vs-attributes`, `igraph-vs-indexing`, `print.igraph.es()`, `print.igraph.vs()`  
Other vertex and edge sequence operations: `c.igraph.es()`, `c.igraph.vs()`, `difference.igraph.es()`, `difference.igraph.vs()`, `igraph-es-indexing`, `igraph-vs-indexing2`, `igraph-vs-indexing`, `intersection.igraph.es()`, `intersection.igraph.vs()`, `rev.igraph.es()`, `rev.igraph.vs()`, `union.igraph.es()`, `union.igraph.vs()`, `unique.igraph.es()`, `unique.igraph.vs()`
igraph-minus

Examples

```r
g <- make_(ring(10),
    with_vertex_(name = LETTERS[1:10]),
    with_edge_(weight = 1:10, color = "green"))
E(g)
E(g)[[]]
E(g)[[.inc('A')]]
```

igraph-minus Delete vertices or edges from a graph

Description

Delete vertices or edges from a graph

Usage

```r
## S3 method for class 'igraph'
e1 - e2
```

Arguments

- `e1` Left argument, see details below.
- `e2` Right argument, see details below.

Details

The minus operator (`-`) can be used to remove vertices or edges from the graph. The operation performed is selected based on the type of the right hand side argument:

- If it is an igraph graph object, then the difference of the two graphs is calculated, see `difference`.
- If it is a numeric or character vector, then it is interpreted as a vector of vertex ids and the specified vertices will be deleted from the graph. Example:

  ```r
g <- make_ring(10)
V(g)$name <- letters[1:10]
g <- g - c("a", "b")
```

- If `e2` is a vertex sequence (e.g. created by the `V` function), then these vertices will be deleted from the graph.
- If it is an edge sequence (e.g. created by the `E` function), then these edges will be deleted from the graph.
- If it is an object created with the `vertex` (or the `vertices`) function, then all arguments of `vertices` are concatenated and the result is interpreted as a vector of vertex ids. These vertices will be removed from the graph.
- If it is an object created with the `edge` (or the `edges`) function, then all arguments of `edges` are concatenated and then interpreted as edges to be removed from the graph. Example:

  ```r
g <- make_ring(10)
V(g)$name <- letters[1:10]
E(g)$name <- LETTERS[1:10]
g <- g - edge("e|f")
g <- g - edge("H")
```
• If it is an object created with the \texttt{path} function, then all \texttt{path} arguments are concatenated and then interpreted as a path along which edges will be removed from the graph. Example:

\begin{verbatim}
g <- make_ring(10)
V(g)$name <- letters[1:10]
g <- g - path("a", "b", "c", "d")
\end{verbatim}

\textbf{Value}

An igraph graph.

\textbf{See Also}

Other functions for manipulating graph structure: \texttt{+.igraph()}, \texttt{add_edges()}, \texttt{add_vertices()}, \texttt{delete_edges()}, \texttt{delete_vertices()}, \texttt{edge()}, \texttt{path()}, \texttt{vertex()}

\begin{verbatim}
igraph-vs-attributes  Query or set attributes of the vertices in a vertex sequence
\end{verbatim}

\textbf{Description}

The \$ operator is a syntactic sugar to query and set the attributes of the vertices in a vertex sequence.

\textbf{Usage}

\begin{verbatim}
### S3 replacement method for class 'igraph.vs'
x[[i]] <- value

### S3 replacement method for class 'igraph.vs'
x[i] <- value

### S3 method for class 'igraph.vs'
x$name

### S3 replacement method for class 'igraph.vs'
x$name <- value

V(x) <- value
\end{verbatim}

\textbf{Arguments}

\begin{itemize}
  \item \texttt{x} A vertex sequence. For \texttt{V<-} it is a graph.
  \item \texttt{i} Index.
  \item \texttt{value} New value of the attribute, for the vertices in the vertex sequence.
  \item \texttt{name} Name of the vertex attribute to query or set.
\end{itemize}

\textbf{Details}

The query form of \$ is a shortcut for \texttt{vertex_attr}, e.g. \texttt{V(g)[idx]$attr} is equivalent to \texttt{vertex_attr(g, attr, V(g)[idx])}.

The assignment form of \$ is a shortcut for \texttt{set_vertex_attr}, e.g. \texttt{V(g)[idx]$attr <- value} is equivalent to \texttt{g <- set_vertex_attr(g, attr, V(g)[idx], value)}. 


Value

A vector or list, containing the values of attribute name for the vertices in the vertex sequence. For numeric, character or logical attributes, it is a vector of the appropriate type, otherwise it is a list.

See Also

Other vertex and edge sequences: E(), V(), igraph-es-attributes, igraph-es-indexing2, igraph-es-indexing, igraph-vs-indexing2, igraph-vs-indexing, print.igraph.es(), print.igraph.vs()

Other graph attributes: delete_edge_attr(), delete_graph_attr(), delete_vertex_attr(), edge_attr<-(), edge_attr_names(), edge_attr(), graph_attr<-(), graph_attr_names(), graph_attr(), igraph-dollar, set_edge_attr(), set_graph_attr(), set_vertex_attr(), vertex_attr<-(), vertex_attr_names(), vertex_attr()

Examples

g <- make_(ring(10),
  with_vertex_(
    name = LETTERS[1:10],
    color = sample(1:2, 10, replace=TRUE)
  )
)
V(g)$name
V(g)$color
V(g)$frame.color <- V(g)$color

# color vertices of the largest component
largest_comp <- function(graph) {
  cl <- components(graph)
  V(graph)[which.max(cl$csize) == cl$membership]
}
g <- sample_(gnp(100, 2/100),
  with_vertex_(size = 3, label = ""),
  with_graph_(layout = layout_with_fr)
)
giant_v <- largest_comp(g)
V(g)$color <- "blue"
V(g)[giant_v]$color <- "orange"
plot(g)

igraph-vs-indexing     Indexing vertex sequences

Description

Vertex sequences can be indexed very much like a plain numeric R vector, with some extras.

Usage

## S3 method for class 'igraph.vs'
x[...], na_ok = FALSE]
Arguments

- **x**: A vertex sequence.
- **...**: Indices, see details below.
- **na_ok**: Whether it is OK to have NAs in the vertex sequence.

Details

Vertex sequences can be indexed using both the single bracket and the double bracket operators, and they both work the same way. The only difference between them is that the double bracket operator marks the result for printing vertex attributes.

Value

Another vertex sequence, referring to the same graph.

Multiple indices

When using multiple indices within the bracket, all of them are evaluated independently, and then the results are concatenated using the `c()` function (except for the `na_ok` argument, which is special and must be named. E.g. `V(g)[1, 2, .nei(1)]` is equivalent to `c(V(g)[1], V(g)[2], V(g)[.nei(1)])`.

Index types

Vertex sequences can be indexed with positive numeric vectors, negative numeric vectors, logical vectors, character vectors:

- When indexed with positive numeric vectors, the vertices at the given positions in the sequence are selected. This is the same as indexing a regular R atomic vector with positive numeric vectors.
- When indexed with negative numeric vectors, the vertices at the given positions in the sequence are omitted. Again, this is the same as indexing a regular R atomic vector.
- When indexed with a logical vector, the lengths of the vertex sequence and the index must match, and the vertices for which the index is `TRUE` are selected.
- Named graphs can be indexed with character vectors, to select vertices with the given names.

Vertex attributes

When indexing vertex sequences, vertex attributes can be referred to simply by using their names. E.g. if a graph has a `name` vertex attribute, then `V(g)[name == "foo"]` is equivalent to `V(g)[V(g)$name == "foo"]`. See more examples below. Note that attribute names mask the names of variables present in the calling environment; if you need to look up a variable and you do not want a similarly named vertex attribute to mask it, use the `.env` pronoun to perform the name lookup in the calling environment. In other words, use `V(g)[.env$name == "foo"]` to make sure that name is looked up from the calling environment even if there is a vertex attribute with the same name. Similarly, you can use `.data` to match attribute names only.

Special functions

There are some special igraph functions that can be used only in expressions indexing vertex sequences:
.nei  takes a vertex sequence as its argument and selects neighbors of these vertices. An optional
  mode argument can be used to select successors (mode="out"), or predecessors (mode="in")
  in directed graphs.

.inc Takes an edge sequence as an argument, and selects vertices that have at least one incident
  edge in this edge sequence.

.from Similar to .inc, but only considers the tails of the edges.

.to Similar to .inc, but only considers the heads of the edges.

.innei, outnei .innei(v) is a shorthand for .nei(v, mode = "in"), and .outnei(v) is a short-
  hand for .nei(v, mode = "out").

Note that multiple special functions can be used together, or with regular indices, and then their
results are concatenated. See more examples below.

See Also

Other vertex and edge sequences: E(), V(), igraph-es-attributes, igraph-es-indexing2,
igraph-es-indexing, igraph-vs-attributes, igraph-vs-indexing2, print.igraph.es(), print.igraph.vs()

Other vertex and edge sequence operations: c.igraph.es(), c.igraph.vs(), difference.igraph.es(),
difference.igraph.vs(), intersection.igraph.es(), intersection.igraph.vs(), rev.igraph.es(), rev.igraph.vs(),
union.igraph.es(), union.igraph.vs(), unique.igraph.es(), unique.igraph.vs()

Examples

# -----------------------------------------------------------------
# Setting attributes for subsets of vertices
largest_comp <- function(graph) {
  cl <- components(graph)
  V(graph)[which.max(cl$csize) == cl$membership]
}
g <- sample_(gnp(100, 2/100),
  with_vertex_(size = 3, label = ""),
  with_graph_(layout = layout_with_fr)
}
giant_v <- largest_comp(g)
V(g)$color <- "green"
V(g)[giant_v]$color <- "red"
plot(g)

# -----------------------------------------------------------------
# nei() special function

# The same with vertex names
\begin{verbatim}
g <- graph(~ A -> B, B -> C:D, D -> B)
V(g)[.nei( c('B', 'D') ) ]
V(g)[.nei( c('B', 'D'), "in") ]
V(g)[.nei( c('B', 'D'), "out") ]
\end{verbatim}

# -----------------------------------------------------------------
# Resolving attributes

g <- graph(~ A -> B, B -> C:D, D -> B)
V(g)$color <- c("red", "red", "green", "green")
V(g)[color == "red"]

# Indexing with a variable whose name matches the name of an attribute
# may fail; use .env to force the name lookup in the parent environment
V(g)$x <- 10:13
x <- 2
V(g)[.env$x]

---

igraph-vs-indexing2

Select vertices and show their metadata

Description

The double bracket operator can be used on vertex sequences, to print the meta-data (vertex attributes) of the vertices in the sequence.

Usage

```r
## S3 method for class 'igraph.vs'
x[[...]]
```

Arguments

- `x` A vertex sequence.
- `...` Additional arguments, passed to `[`. 

Details

Technically, when used with vertex sequences, the double bracket operator does exactly the same as the single bracket operator, but the resulting vertex sequence is printed differently: all attributes of the vertices in the sequence are printed as well.

See `.[.igraph.vs` for more about indexing vertex sequences.

Value

The double bracket operator returns another vertex sequence, with meta-data (attribute) printing turned on. See details below.

See Also


Examples

```r
library(igraph)

# Create a ring graph with 10 vertices
g <- make_ring(10)
# Set vertex attributes
set_vertex_attr("color", value = "red")
set_vertex_attr("name", value = LETTERS[1:10])

V(g)
V(g)[[]]
V(g)[1:5]
V(g)[[1:5]]
```

igraph_demo
Run igraph demos, step by step

Description
Run one of the accompanying igraph demos, somewhat interactively, using a Tk window.

Usage
`igraph_demo(which)`

Arguments
- `which`: If not given, then the names of the available demos are listed. Otherwise it should be either a filename or the name of an igraph demo.

Details
This function provides a somewhat nicer interface to igraph demos that come with the package, than the standard `demo` function. igraph demos are divided into chunks and `igraph_demo` runs them chunk by chunk, with the possibility of inspecting the workspace between two chunks.

The `tcltk` package is needed for `igraph_demo`.

Value
Returns NULL, invisibly.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

See Also
demo

Examples
```r
igraph_demo()
if (interactive()) {
  igraph_demo("centrality")
}
```
igraph_options  Parameters for the igraph package

Description

igraph has some parameters which (usually) affect the behavior of many functions. These can be set for the whole session via `igraph_options`.

Usage

`igraph_options(...)`

`igraph_opt(x, default = NULL)`

Arguments

`...`  A list may be given as the only argument, or any number of arguments may be in the `name=value` form, or no argument at all may be given. See the Value and Details sections for explanation.

`x`  A character string holding an option name.

`default`  If the specified option is not set in the options list, this value is returned. This facilitates retrieving an option and checking whether it is set and setting it separately if not.

Details

The parameter values set via a call to the `igraph_options` function will remain in effect for the rest of the session, affecting the subsequent behaviour of the other functions of the igraph package for which the given parameters are relevant.

This offers the possibility of customizing the functioning of the igraph package, for instance by insertions of appropriate calls to `igraph_options` in a load hook for package igraph.

The currently used parameters in alphabetical order:

- **add.params** Logical scalar, whether to add model parameter to the graphs that are created by the various graph constructors. By default it is TRUE.
- **add.vertex.names** Logical scalar, whether to add vertex names to node level indices, like degree, betweenness scores, etc. By default it is TRUE.
- **annotate.plot** Logical scalar, whether to annotate igraph plots with the graph’s name (name graph attribute, if present) as main, and with the number of vertices and edges as xlab. Defaults to FALSE.
- **dend.plot.type** The plotting function to use when plotting community structure dendrograms via `plot_dendrogram`. Possible values are ‘auto’ (the default), ‘phylo’, ‘hclust’ and ‘dendrogram’. See `plot_dendrogram` for details.
- **edge.attr.comb** Specifies what to do with the edge attributes if the graph is modified. The default value is `list(weight="sum", name="concat", "ignore")`. See `attribute.combination` for details on this.
- **print.edge.attributes** Logical constant, whether to print edge attributes when printing graphs. Defaults to FALSE.
igraph_options

print.full Logical scalar, whether print.igraph should show the graph structure as well, or only a summary of the graph.

print.graph.attributes Logical constant, whether to print graph attributes when printing graphs. Defaults to FALSE.

print.vertex.attributes Logical constant, whether to print vertex attributes when printing graphs. Defaults to FALSE.

return.vs.es Whether functions that return a set or sequence of vertices/edges should return formal vertex/edge sequence objects. This option was introduced in igraph version 1.0.0 and defaults to TRUE. If your package requires the old behavior, you can set it to FALSE in the .onLoad function of your package, without affecting other packages.

sparsematrices Whether to use the Matrix package for (sparse) matrices. It is recommended, if the user works with larger graphs.

verbose Logical constant, whether igraph functions should talk more than minimal. Eg. if TRUE then some functions will use progress bars while computing. Defaults to FALSE.

vertex.attr.comb Specifies what to do with the vertex attributes if the graph is modified. The default value is list(name="concat", "ignore") See attribute.combination for details on this.

Value

igraph_options returns a list with the old values of the updated parameters, invisibly. Without any arguments, it returns the values of all options.

For igraph_opt, the current value set for option x, or NULL if the option is unset.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

igraph_options is similar to options and igraph_opt is similar to getOption.

Other igraph options: with_igraph_opt()

Examples

oldval <- igraph_opt("verbose")
igraph_options(verbos TRUE)
layout_with_kk(make_ring(10))
igraph_options(verbos = oldval)

oldval <- igraph_options(verbos = TRUE, sparsematrices = FALSE)
make_ring(10)[]
igraph_options(oldval)
igraph_opt("verbose")
igraph_test  

Description

Runs all package tests.

Usage

igraph_test()

Details

The testthat package is needed to run all tests. The location tests themselves can be extracted from the package via system.file("tests",package="igraph").

This function simply calls the test_dir function from the testthat package on the test directory.

Value

Whatever is returned by test_dir from the testthat package.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

igraph_version  

Description

Queries igraph's original version string. See details below.

Usage

igraph_version()

Details

The igraph version string is the same as the version of the R package for all released igraph versions. For development versions and nightly builds, they might differ however.

The reason for this is, that R package version numbers are not flexible enough to cover in-between releases versions, e.g. alpha and beta versions, release candidates, etc.

Value

A character scalar, the igraph version string.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>
## Incident edges of a vertex in a graph

**incident**

**Description**

Incident edges of a vertex in a graph

**Usage**

```r
incident(graph, v, mode = c("all", "out", "in", "total"))
```

**Arguments**

- `graph`: The input graph.
- `v`: The vertex of which the incident edges are queried.
- `mode`: Whether to query outgoing ("out"), incoming ("in") edges, or both types ("all"). This is ignored for undirected graphs.

**Value**

An edge sequence containing the incident edges of the input vertex.

**See Also**

Other structural queries: `[.igraph()`, `[[.igraph()`, `adjacent_vertices()`, `are_adjacent()`, `ends()`, `get.edge.ids()`, `gorder()`, `gsize()`, `head_of()`, `incident_edges()`, `is_directed()`, `neighbors()`, `tail_of()`

**Examples**

```r
g <- make_graph("Zachary")
incident(g, 1)
incident(g, 34)
```
incident_edges  Incident edges of multiple vertices in a graph

Description

This function is similar to `incident`, but it queries multiple vertices at once.

Usage

```r
incident_edges(graph, v, mode = c("out", "in", "all", "total"))
```

Arguments

- `graph`: Input graph.
- `v`: The vertices to query.
- `mode`: Whether to query outgoing (‘out’), incoming (‘in’) edges, or both types (‘all’). This is ignored for undirected graphs.

Value

A list of edge sequences.

See Also

Other structural queries: `.[igraph()`, `[[.igraph()`, `adjacent_vertices()`, `are_adjacent()`, `ends()`, `get.edge.ids()`, `gorder()`, `gsize()`, `head_of()`, `incident()`, `is_directed()`, `neighbors()`, `ends()`, `tail_of()`

Examples

```r
g <- make_graph("Zachary")
incident_edges(g, c(1, 34))
```

indent_print  Indent a printout

Description

Indent a printout

Usage

```r
indent_print(..., .indent = " ", .printer = print)
```

Arguments

- `...`: Passed to the printing function.
- `.indent`: Character scalar, indent the printout with this.
- `.printer`: The printing function, defaults to `print`.
intersection

Value

The first element in . . . , invisibly.

intersection  Intersection of two or more sets

Description

This is an S3 generic function. See methods("intersection") for the actual implementations for various S3 classes. Initially it is implemented for igraph graphs and igraph vertex and edge sequences. See intersection.igraph, and intersection.igraph.vs.

Usage

intersection(...)

Arguments

...  Arguments, their number and interpretation depends on the function that implements intersection.

Value

Depends on the function that implements this method.

intersection.igraph  Intersection of graphs

Description

The intersection of two or more graphs are created. The graphs may have identical or overlapping vertex sets.

Usage

## S3 method for class 'igraph'
intersection(..., byname = "auto", keep.all.vertices = TRUE)

Arguments

...  Graph objects or lists of graph objects.
byname  A logical scalar, or the character scalar auto. Whether to perform the operation based on symbolic vertex names. If it is auto, that means TRUE if all graphs are named and FALSE otherwise. A warning is generated if auto and some (but not all) graphs are named.
keep.all.vertices  Logical scalar, whether to keep vertices that only appear in a subset of the input graphs.
Details

intersection creates the intersection of two or more graphs: only edges present in all graphs will
be included. The corresponding operator is %s%.

If the byname argument is TRUE (or auto and all graphs are named), then the operation is performed
on symbolic vertex names instead of the internal numeric vertex ids.

intersection keeps the attributes of all graphs. All graph, vertex and edge attributes are copied
to the result. If an attribute is present in multiple graphs and would result a name clash, then this
attribute is renamed by adding suffixes: _1, _2, etc.

The name vertex attribute is treated specially if the operation is performed based on symbolic vertex
names. In this case name must be present in all graphs, and it is not renamed in the result graph.

An error is generated if some input graphs are directed and others are undirected.

Value

A new graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

## Common part of two social networks
net1 <- graph_from_literal(D-A:B:F:G, A-C-F-A, B-E-G-B, A-B, F-G,
H-F:G, H-I-J)
print_all(net1 %s% net2)

---

intersection.igraph.es

Intersection of edge sequences

Description

Intersection of edge sequences

Usage

## S3 method for class 'igraph.es'
intersection(...)

Arguments

... The edge sequences to take the intersection of.

Details

They must belong to the same graph. Note that this function has ‘set’ semantics and the multiplicity
of edges is lost in the result.


**Value**

An edge sequence that contains edges that appear in all given sequences, each edge exactly once.

**See Also**


**Examples**

```r
g <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
intersection(E(g)[1:6], E(g)[5:9])
```

---

**Description**

Intersection of vertex sequences

**Usage**

```r
## S3 method for class 'igraph.vs'
intersection(...)
```

**Arguments**

... 

The vertex sequences to take the intersection of.

**Details**

They must belong to the same graph. Note that this function has ‘set’ semantics and the multiplicity of vertices is lost in the result.

**Value**

A vertex sequence that contains vertices that appear in all given sequences, each vertex exactly once.

**See Also**


**Examples**

```r
g <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
intersection(E(g)[1:6], E(g)[5:9])
```
**is_bipartite**

Create a bipartite graph

**Description**

A bipartite graph has two kinds of vertices and connections are only allowed between different kinds.

**Usage**

is_bipartite(graph)

make_bipartite_graph(types, edges, directed = FALSE)

bipartite_graph(...)

**Arguments**

- **graph**
  - The input graph.

- **types**
  - A vector giving the vertex types. It will be coerced into boolean. The length of the vector gives the number of vertices in the graph. When the vector is a named vector, the names will be attached to the graph as the name vertex attribute.

- **edges**
  - A vector giving the edges of the graph, the same way as for the regular `graph` function. It is checked that the edges indeed connect vertices of different kind, according to the supplied `types` vector. The vector may be a string vector if `types` is a named vector.

- **directed**
  - Whether to create a directed graph, boolean constant. Note that by default undirected graphs are created, as this is more common for bipartite graphs.

- **...**
  - Passed to `make_bipartite_graph`.

**Details**

Bipartite graphs have a type vertex attribute in igraph, this is boolean and `FALSE` for the vertices of the first kind and `TRUE` for vertices of the second kind.

`make_bipartite_graph` basically does three things. First it checks the `edges` vector against the `types` vector. Then it creates a graph using the `edges` vector and finally it adds the `types` vector as a vertex attribute called `type`. `edges` may contain strings as vertex names; in this case, `types` must be a named vector that specifies the type for each vertex name that occurs in `edges`.

`is_bipartite` checks whether the graph is bipartite or not. It just checks whether the graph has a vertex attribute called `type`.

**Value**

- `make_bipartite_graph` returns a bipartite igraph graph. In other words, an igraph graph that has a vertex attribute named `type`.

- `is_bipartite` returns a logical scalar.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>
**is_chordal**

See Also

- `graph` to create one-mode networks

Examples

```r
g <- make_bipartite_graph(rep(0:1, length.out=10), c(1:10))
print(g, v=TRUE)
```

---

**is_chordal**

Chordality of a graph

Description

A graph is chordal (or triangulated) if each of its cycles of four or more nodes has a chord, which is an edge joining two nodes that are not adjacent in the cycle. An equivalent definition is that any chordless cycles have at most three nodes.

Usage

```r
is_chordal(
  graph,
  alpha = NULL,
  alpham1 = NULL,
  fillin = FALSE,
  newgraph = FALSE
)
```

Arguments

- **graph**: The input graph. It may be directed, but edge directions are ignored, as the algorithm is defined for undirected graphs.
- **alpha**: Numeric vector, the maximal cardinality ordering of the vertices. If it is `NULL`, then it is automatically calculated by calling `max_cardinality`, or from `alpham1` if that is given.
- **alpham1**: Numeric vector, the inverse of `alpha`. If it is `NULL`, then it is automatically calculated by calling `max_cardinality`, or from `alpha`.
- **fillin**: Logical scalar, whether to calculate the fill-in edges.
- **newgraph**: Logical scalar, whether to calculate the triangulated graph.

Details

The chordality of the graph is decided by first performing maximum cardinality search on it (if the `alpha` and `alpham1` arguments are `NULL`), and then calculating the set of fill-in edges.

The set of fill-in edges is empty if and only if the graph is chordal.

It is also true that adding the fill-in edges to the graph makes it chordal.
is_dag

Value

A list with three members:

- **chordal**: Logical scalar, it is TRUE iff the input graph is chordal.
- **fillin**: If requested, then a numeric vector giving the fill-in edges. NULL otherwise.
- **newgraph**: If requested, then the triangulated graph, an igraph object. NULL otherwise.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

max_cardinality

Examples

```r
## The examples from the Tarjan-Yannakakis paper
gl <- graph_from_literal(A-B:C:I, B-A:C:D, C-A:B:E:H, D-B:E:F,  
                          I-A:H)
max_cardinality(gl)
is_chordal(gl, fillin=TRUE)

g2 <- graph_from_literal(A-B:E, B-A:E:F:D, C-E:D:G, D-B:F:E:C:G,  
max_cardinality(g2)
is_chordal(g2, fillin=TRUE)
```

is_dag

*Directed acyclic graphs*

Description

This function tests whether the given graph is a DAG, a directed acyclic graph.

Usage

is_dag(graph)

Arguments

- **graph**: The input graph. It may be undirected, in which case FALSE is reported.
is_degseq

Details

is_dag checks whether there is a directed cycle in the graph. If not, the graph is a DAG.

Value

A logical vector of length one.

Author(s)

Tamas Nepusz <ntamas@gmail.com> for the C code, Gabor Csardi <csardi.gabor@gmail.com> for the R interface.

Examples

g <- make_tree(10)
is_dag(g)
g2 <- g + edge(5,1)
is_dag(g2)

is_degseq

Check if a degree sequence is valid for a multi-graph

Description

is_degseq checks whether the given vertex degrees (in- and out-degrees for directed graphs) can be realized by a graph. Note that the graph does not have to be simple, it may contain loop and multiple edges. For undirected graphs, it also checks whether the sum of degrees is even. For directed graphs, the function checks whether the lengths of the two degree vectors are equal and whether their sums are also equal. These are known sufficient and necessary conditions for a degree sequence to be valid.

Usage

is_degseq(out.deg, in.deg = NULL)

Arguments

out.deg  Integer vector, the degree sequence for undirected graphs, or the out-degree sequence for directed graphs.

in.deg   NULL or an integer vector. For undirected graphs, it should be NULL. For directed graphs it specifies the in-degrees.

Value

A logical scalar.

Author(s)

Tamas Nepusz <ntamas@gmail.com> and Szabolcs Horvat <szhorvat@gmail.com>
is_directed

References
G. Cairns and S. Mendan, Degree Sequence for Graphs with Loops (2013).

See Also
Other graphical degree sequences: is_graphical()

Examples
```
g <- sample_gnp(100, 2/100)
is_degseq(degree(g))
is_graphical(degree(g))
```

Check whether a graph is directed

Description
Check whether a graph is directed

Usage
```
is_directed(graph)
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>graph</td>
<td>The input graph</td>
</tr>
</tbody>
</table>

Value
Logical scalar, whether the graph is directed.

See Also
Other structural queries: [.igraph(), [[.igraph(), adjacent_vertices(), are_adjacent(), ends(), get.edge.ids(), gorder(), gsize(), head_of(), incident_edges(), incident(), neighbors(), tail_of()]

Examples
```
g <- make_ring(10)
is_directed(g)

g2 <- make_ring(10, directed = TRUE)
is_directed(g2)
```
Description

Determine whether the given vertex degrees (in- and out-degrees for directed graphs) can be realized in a graph.

Usage

\[
\text{is_graphical}(\text{out.deg}, \text{in.deg} = \text{NULL}, \text{allowed.edge.types} = \text{c("simple", "loops", "multi", "all")})
\]

Arguments

out.deg
- Integer vector, the degree sequence for undirected graphs, or the out-degree sequence for directed graphs.
in.deg
- NULL or an integer vector. For undirected graphs, it should be NULL. For directed graphs it specifies the in-degrees.
allowed.edge.types
- The allowed edge types in the graph. ‘simple’ means that neither loop nor multiple edges are allowed (i.e. the graph must be simple). ‘loops’ means that loop edges are allowed but multiple edges are not. ‘multi’ means that multiple edges are allowed but loop edges are not. ‘all’ means that both loop edges and multiple edges are allowed.

Details

The classical concept of graphicality assumes simple graphs. This function can perform the check also when self-loops, multi-edges, or both are allowed in the graph.

Value

A logical scalar.

Author(s)

Tamas Nepusz <ntamas@gmail.com>

References


See Also

Other graphical degree sequences: \text{is_degseq()}
Examples

g <- sample_gnp(100, 2/100)
is_degseq(degree(g))
is_graphical(degree(g))

is_igraph

Is this object an igraph graph?

Description

Is this object an igraph graph?

Usage

is_igraph(graph)

Arguments

graph An R object.

Value

A logical constant, TRUE if argument graph is a graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

g <- make_ring(10)
is_igraph(g)
is_igraph(numeric(10))

is_matching

Graph matching

Description

A matching in a graph means the selection of a set of edges that are pairwise non-adjacent, i.e. they
have no common incident vertices. A matching is maximal if it is not a proper subset of any other
matching.
is_matching

is_matching(graph, matching, types = NULL)

is_max_matching(graph, matching, types = NULL)

max_bipartite_match(
  graph,
  types = NULL,
  weights = NULL,
  eps = .Machine$double.eps
)

Arguments

graph The input graph. It might be directed, but edge directions will be ignored.
matching A potential matching. An integer vector that gives the pair in the matching for each vertex. For vertices without a pair, supply NA here.
types Vertex types, if the graph is bipartite. By default they are taken from the ‘type’ vertex attribute, if present.
weights Potential edge weights. If the graph has an edge attribute called ‘weight’, and this argument is NULL, then the edge attribute is used automatically. In weighted matching, the weights of the edges must match as much as possible.
eps A small real number used in equality tests in the weighted bipartite matching algorithm. Two real numbers are considered equal in the algorithm if their difference is smaller than eps. This is required to avoid the accumulation of numerical errors. By default it is set to the smallest x, such that 1 + x ≠ 1 holds. If you are running the algorithm with no weights, this argument is ignored.

Details

is_matching checks a matching vector and verifies whether its length matches the number of vertices in the given graph, its values are between zero (inclusive) and the number of vertices (inclusive), and whether there exists a corresponding edge in the graph for every matched vertex pair. For bipartite graphs, it also verifies whether the matched vertices are in different parts of the graph.
is_max_matching checks whether a matching is maximal. A matching is maximal if and only if there exists no unmatched vertex in a graph such that one of its neighbors is also unmatched.
max_bipartite_match calculates a maximum matching in a bipartite graph. A matching in a bipartite graph is a partial assignment of vertices of the first kind to vertices of the second kind such that each vertex of the first kind is matched to at most one vertex of the second kind and vice versa, and matched vertices must be connected by an edge in the graph. The size (or cardinality) of a matching is the number of edges. A matching is a maximum matching if there exists no other matching with larger cardinality. For weighted graphs, a maximum matching is a matching whose edges have the largest possible total weight among all possible matchings.

Maximum matchings in bipartite graphs are found by the push-relabel algorithm with greedy initialization and a global relabeling after every n/2 steps where n is the number of vertices in the graph.

Value

is_matching and is_max_matching return a logical scalar.
max_bipartite_match returns a list with components:


**is_min_separator**

**Description**

Check whether a given set of vertices is a minimal vertex separator.

**Usage**

```
is_min_separator(graph, candidate)
```

**Arguments**

- `graph` The input graph. It may be directed, but edge directions are ignored.
- `candidate` A numeric vector giving the vertex ids of the candidate separator.
Details

\texttt{is\_min\_separator} decides whether the supplied vertex set is a minimal vertex separator. A minimal vertex separator is a vertex separator, such that none of its subsets is a vertex separator.

In the special case of a fully connected graph with \(n\) vertices, each set of \(n-1\) vertices is considered to be a vertex separator.

Value

A logical scalar, whether the supplied vertex set is a (minimal) vertex separator or not.

See Also

\texttt{min\_separators} lists all vertex separator of minimum size.

Examples

# The graph from the Moody-White paper
\begin{verbatim}
5-6:7:21, 6-7, 7-8:11:14:19, 8-9:11:14, 9-10,
10-12:13, 11-12:14, 12-16, 13-16, 14-15, 15-16,
21-22:23, 22-23)\end{verbatim}

# Cohesive subgraphs
mw1 <- induced_subgraph(mw, as.character(c(1:7, 17:23)))
mw2 <- induced_subgraph(mw, as.character(7:16))
mw3 <- induced_subgraph(mw, as.character(17:23))
mw4 <- induced_subgraph(mw, as.character(c(7,8,11,14)))
mw5 <- induced_subgraph(mw, as.character(1:7))

check.sep <- function(G) {
  sep <- min_separators(G)
  sapply(sep, is_min_separator, graph=G)
}

check.sep(mw)
check.sep(mw1)
check.sep(mw2)
check.sep(mw3)
check.sep(mw4)
check.sep(mw5)

---

\textbf{is\_named} \hspace{1cm} \textit{Named graphs}

Description

An igraph graph is named, if there is a symbolic name associated with its vertices.

Usage

\texttt{is\_named(graph)}
Arguments

graph The input graph.

Details

In igraph vertices can always be identified and specified via their numeric vertex ids. This is, however, not always convenient, and in many cases there exist symbolic ids that correspond to the vertices. To allow this more flexible identification of vertices, one can assign a vertex attribute called 'name' to an igraph graph. After doing this, the symbolic vertex names can be used in all igraph functions, instead of the numeric ids.

Note that the uniqueness of vertex names are currently not enforced in igraph, you have to check that for yourself, when assigning the vertex names.

Value

A logical scalar.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

```r
g <- make_ring(10)
is_named(g)
V(g)$name <- letters[1:10]
is_named(g)
neighbors(g, "a")
```

---

**is_printer_callback**  
*Is this a printer callback?*

Description

Is this a printer callback?

Usage

```r
is_printer_callback(x)
```

Arguments

x An R object.

See Also

Other printer callbacks: printer_callback()
is_separator

Vertex separators

Description
Check whether a given set of vertices is a vertex separator.

Usage
is_separator(graph, candidate)

Arguments
- graph: The input graph. It may be directed, but edge directions are ignored.
- candidate: A numeric vector giving the vertex ids of the candidate separator.

Details
is_separator decides whether the supplied vertex set is a vertex separator. A vertex set is a vertex separator if its removal results a disconnected graph.

In the special case of a fully connected graph with \( n \) vertices, each set of \( n - 1 \) vertices is considered to be a vertex separator.

Value
A logical scalar, whether the supplied vertex set is a (minimal) vertex separator or not.

See Also
- is_min_separator, min_separators lists all vertex separator of minimum size.

is_tree

Decide whether a graph is a tree.

Description
is_tree decides whether a graph is a tree, and optionally returns a possible root vertex if the graph is a tree.

Usage
is_tree(graph, mode = c("out", "in", "all", "total"), details = FALSE)

Arguments
- graph: An igraph graph object
- mode: Whether to consider edge directions in a directed graph. ‘all’ ignores edge directions; ‘out’ requires edges to be oriented outwards from the root, ‘in’ requires edges to be oriented towards the root.
- details: Whether to return only whether the graph is a tree (FALSE) or also a possible root (TRUE)
Details

An undirected graph is a tree if it is connected and has no cycles. In the directed case, a possible additional requirement is that all edges are oriented away from a root (out-tree or arborescence) or all edges are oriented towards a root (in-tree or anti-arborescence). This test can be controlled using the mode parameter.

By convention, the null graph (i.e. the graph with no vertices) is considered not to be a tree.

Value

When details is FALSE, a logical value that indicates whether the graph is a tree. When details is TRUE, a named list with two entries:

- res: Logical value that indicates whether the graph is a tree.
- root: The root vertex of the tree; undefined if the graph is not a tree.

Examples

```r
g <- make_tree(7, 2)
is_tree(g)
is_tree(g, details=TRUE)
```

is_weighted

Weighted graphs

Description

In weighted graphs, a real number is assigned to each (directed or undirected) edge.

Usage

`is_weighted(graph)`

Arguments

- graph: The input graph.

Details

In igraph edge weights are represented via an edge attribute, called ‘weight’. The `is_weighted` function only checks that such an attribute exists. (It does not even checks that it is a numeric edge attribute.)

Edge weights are used for different purposes by the different functions. E.g. shortest path functions use it as the cost of the path; community finding methods use it as the strength of the relationship between two vertices, etc. Check the manual pages of the functions working with weighted graphs for details.

Value

A logical scalar.
isomorphic

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

Examples

```r
g <- make_ring(10)
shortest_paths(g, 8, 2)
E(g)$weight <- seq_len(ecount(g))
shortest_paths(g, 8, 2)
```

isomorphic

Decide if two graphs are isomorphic

Description

Decide if two graphs are isomorphic

Usage

```r
isomorphic(graph1, graph2, method = c("auto", "direct", "vf2", "bliss"), ...)
```

is_isomorphic_to(
    graph1,
    graph2,
    method = c("auto", "direct", "vf2", "bliss"),
    ...
)

Arguments

- `graph1`: The first graph.
- `graph2`: The second graph.
- `...`: Additional arguments, passed to the various methods.

Value

Logical scalar, TRUE if the graphs are isomorphic.

‘auto’ method

It tries to select the appropriate method based on the two graphs. This is the algorithm it uses:

1. If the two graphs do not agree on their order and size (i.e. number of vertices and edges), then return FALSE.
2. If the graphs have three or four vertices, then the ‘direct’ method is used.
3. If the graphs are directed, then the ‘vf2’ method is used.
4. Otherwise the ‘bliss’ method is used.
‘direct’ method

This method only works on graphs with three or four vertices, and it is based on a pre-calculated and stored table. It does not have any extra arguments.

‘vf2’ method

This method uses the VF2 algorithm by Cordella, Foggia et al., see references below. It supports vertex and edge colors and have the following extra arguments:

vertex.color1, vertex.color2 Optional integer vectors giving the colors of the vertices for colored graph isomorphism. If they are not given, but the graph has a “color” vertex attribute, then it will be used. If you want to ignore these attributes, then supply NULL for both of these arguments. See also examples below.

edge.color1, edge.color2 Optional integer vectors giving the colors of the edges for edge-colored (sub)graph isomorphism. If they are not given, but the graph has a “color” edge attribute, then it will be used. If you want to ignore these attributes, then supply NULL for both of these arguments.

‘bliss’ method

Uses the BLISS algorithm by Junttila and Kaski, and it works for undirected graphs. For both graphs the canonical_permutation and then the permute function is called to transfer them into canonical form; finally the canonical forms are compared. Extra arguments:

sh Character constant, the heuristics to use in the BLISS algorithm for graph1 and graph2. See the sh argument of canonical_permutation for possible values.

sh defaults to ‘fm’.

References


See Also

Other graph isomorphism: count_isomorphisms(), count_subgraph_isomorphisms(), graph_from_isomorphism_class(), isomorphism_class(), isomorphisms(), subgraph_isomorphic(), subgraph_isomorphisms()

Examples

# create some non-isomorphic graphs
g1 <- graph_from_isomorphism_class(3, 10)
g2 <- graph_from_isomorphism_class(3, 11)
isomorphic(g1, g2)

# create two isomorphic graphs, by permuting the vertices of the first
g1 <- barabasi.game(30, m=2, directed=FALSE)
g2 <- permute(g1, sample(vcount(g1)))
# should be TRUE
"isomorphism_class"

isomorphic(g1, g2)

isomorphic(g1, g2, method = "bliss")

isomorphic(g1, g2, method = "vf2")

# colored graph isomorphism

g1 <- make_ring(10)
g2 <- make_ring(10)

isomorphic(g1, g2)

V(g1)$color <- rep(1:2, length = vcount(g1))

V(g2)$color <- rep(2:1, length = vcount(g2))

# consider colors by default

count_isomorphisms(g1, g2)

# ignore colors

count_isomorphisms(g1, g2, vertex.color1 = NULL,

vertex.color2 = NULL)

---

**Description**

The isomorphism class is a non-negative integer number. Graphs (with the same number of vertices) having the same isomorphism class are isomorphic and isomorphic graphs always have the same isomorphism class. Currently it can handle directed graphs with 3 or 4 vertices and undirected graphs with 3 to 6 vertices.

**Usage**

`isomorphism_class(graph, v)`

**Arguments**

- `graph` The input graph.
- `v` Optionally a vertex sequence. If not missing, then an induced subgraph of the input graph, consisting of this vertices, is used.

**Value**

An integer number.

**See Also**

Other graph isomorphism: `count_isomorphisms()`, `count_subgraph_isomorphisms()`, `graph_from_isomorphism_class()`, `isomorphic()`, `isomorphisms()`, `subgraph_isomorphic()`, `subgraph_isomorphisms()`

**Examples**

# create some non-isomorphic graphs

g1 <- graph_from_isomorphism_class(3, 10)
g2 <- graph_from_isomorphism_class(3, 11)

isomorphism_class(g1)
isomorphism_class(g2)
isomorphic(g1, g2)
isomorphisms

*Calculate all isomorphic mappings between the vertices of two graphs*

**Description**

Calculate all isomorphic mappings between the vertices of two graphs

**Usage**

`isomorphisms(graph1, graph2, method = "vf2", ...)`

**Arguments**

- `graph1`: The first graph.
- `graph2`: The second graph.
- `method`: Currently only `vf2` is supported, see `isomorphic` for details about it and extra arguments.
- `...`: Extra arguments, passed to the various methods.

**Value**

A list of vertex sequences, corresponding to all mappings from the first graph to the second.

**See Also**

Other graph isomorphism: `count_isomorphisms()`, `count_subgraph_isomorphisms()`, `graph_from_isomorphism_class()`, `isomorphic()`, `isomorphism_class()`, `subgraph_isomorphic()`, `subgraph_isomorphisms()`

ivs

*Independent vertex sets*

**Description**

A vertex set is called independent if there are no edges between any two vertices in it. These functions find independent vertex sets in undirected graphs

**Usage**

`ivs(graph, min = NULL, max = NULL)`

**Arguments**

- `graph`: The input graph. Directed graphs are considered as undirected, loop edges and multiple edges are ignored.
- `min`: Numeric constant, limit for the minimum size of the independent vertex sets to find. NULL means no limit.
- `max`: Numeric constant, limit for the maximum size of the independent vertex sets to find. NULL means no limit.
**Details**

`ivs` finds all independent vertex sets in the network, obeying the size limitations given in the `min` and `max` arguments.

`largest_ivs` finds the largest independent vertex sets in the graph. An independent vertex set is largest if there is no independent vertex set with more vertices.

`maximal_ivs` finds the maximal independent vertex sets in the graph. An independent vertex set is maximal if it cannot be extended to a larger independent vertex set. The largest independent vertex sets are maximal, but the opposite is not always true.

`independence.number` calculate the size of the largest independent vertex set(s).

These functions use the algorithm described by Tsukiyama et al., see reference below.

**Value**

`ivs`, `largest_ivs` and `maximal_ivs` return a list containing numeric vertex ids, each list element is an independent vertex set.

`ivs_size` returns an integer constant.

**Author(s)**

Tamas Nepusz <ntamas@gmail.com> ported it from the Very Nauty Graph Library by Keith Briggs ([http://keithbriggs.info/](http://keithbriggs.info/)) and Gabor Csardi <csardi.gabor@gmail.com> wrote the R interface and this manual page.

**References**


**See Also**

`cliques`

**Examples**

# Do not run, takes a couple of seconds
## Not run:

# A quite dense graph
set.seed(42)
g <- sample_gnp(100, 0.9)
ivs_size(g)
ivs(g, min=ivs_size(g))
largest_ivs(g)
# Empty graph
induced_subgraph(g, largest_ivs(g)[[1]])

length(maximal_ivs(g))

## End(Not run)
keeping_degseq

Graph rewiring while preserving the degree distribution

Description

This function can be used together with `rewire` to randomly rewire the edges while preserving the original graph’s degree distribution.

Usage

```r
keeping_degseq(loops = FALSE, niter = 100)
```

Arguments

- `loops` Whether to allow destroying and creating loop edges.
- `niter` Number of rewiring trials to perform.

Details

The rewiring algorithm chooses two arbitrary edges in each step ((a,b) and (c,d)) and substitutes them with (a,d) and (c,b), if they not already exists in the graph. The algorithm does not create multiple edges.

Author(s)

Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com>

See Also

`sample_degseq`

Other rewiring functions: `each_edge()`, `rewire()`

Examples

```r
g <- make_ring(10)
g %>%
  rewire(keeping_degseq(niter = 20)) %>%
  degree()
print_all(rewire(g, with = keeping_degseq(niter = vcount(g) * 10)))
```
**Description**

Calculate the average nearest neighbor degree of the given vertices and the same quantity in the function of vertex degree.

**Usage**

```r
knn(
  graph,
  vids = V(graph),
  mode = c("all", "out", "in", "total"),
  neighbor_degree.mode = c("all", "out", "in", "total"),
  weights = NULL
)
```

**Arguments**

- `graph` The input graph. It may be directed.
- `vids` The vertices for which the calculation is performed. Normally it includes all vertices. Note, that if not all vertices are given here, then both ‘knn’ and ‘knnk’ will be calculated based on the given vertices only.
- `mode` Character constant to indicate the type of neighbors to consider in directed graphs. `out` considers out-neighbors, `in` considers in-neighbors and `all` ignores edge directions.
- `neighbor_degree.mode` The type of degree to average in directed graphs. out averages out-degrees, in averages in-degrees and all ignores edge directions for the degree calculation.
- `weights` Weight vector. If the graph has a weight edge attribute, then this is used by default. If this argument is given, then vertex strength (see `strength`) is used instead of vertex degree. But note that knnk is still given in the function of the normal vertex degree. Weights are used to calculate a weighted degree (also called `strength`) instead of the degree.

**Details**

Note that for zero degree vertices the answer in ‘knn’ is NaN (zero divided by zero), the same is true for ‘knnk’ if a given degree never appears in the network.

The weighted version computes a weighted average of the neighbor degrees as

\[ k_{nn,u} = 1/s_u \sum_v w_{uv} k_v, \]

where \( s_u = \sum_v w_{uv} \) is the sum of the incident edge weights of vertex \( u \), i.e. its strength. The sum runs over the neighbors \( v \) of vertex \( u \) as indicated by mode. \( w_{uv} \) denotes the weighted adjacency matrix and \( k_v \) is the neighbors’ degree, specified by `neighbor_degree.mode`.
Value

A list with two members:

- **knn**: A numeric vector giving the average nearest neighbor degree for all vertices in `vids`.
- **knnk**: A numeric vector, its length is the maximum (total) vertex degree in the graph. The first element is the average nearest neighbor degree of vertices with degree one, etc.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


Examples

```r
# Some trivial ones
g <- make_ring(10)
knn(g)
g2 <- make_star(10)
knn(g2)

# A scale-free one, try to plot 'knnk'
g3 <- sample_pa(1000, m=5)
knn(g3)

# A random graph
g4 <- sample_gnp(1000, p=5/1000)
knn(g4)

# A weighted graph
g5 <- make_star(10)
E(g5)$weight <- seq(ecount(g5))
knn(g5)
```

Description

The Laplacian of a graph.

Usage

```r
laplacian_matrix(  
  graph,  
  normalized = FALSE,  
  weights = NULL,  
  sparse = igraph_opt("sparsematrices")  
)
```
Arguments

- **graph**: The input graph.
- **normalized**: Whether to calculate the normalized Laplacian. See definitions below.
- **weights**: An optional vector giving edge weights for weighted Laplacian matrix. If this is NULL and the graph has an edge attribute called weight, then it will be used automatically. Set this to NA if you want the unweighted Laplacian on a graph that has a weight edge attribute.
- **sparse**: Logical scalar, whether to return the result as a sparse matrix. The Matrix package is required for sparse matrices.

Details

The Laplacian Matrix of a graph is a symmetric matrix having the same number of rows and columns as the number of vertices in the graph and element (i,j) is d[i], the degree of vertex i if if i==j, -1 if i!=j and there is an edge between vertices i and j and 0 otherwise.

A normalized version of the Laplacian Matrix is similar: element (i,j) is 1 if i==j, -1/sqrt(d[i] d[j]) if i!=j and there is an edge between vertices i and j and 0 otherwise.

The weighted version of the Laplacian simply works with the weighted degree instead of the plain degree. I.e. (i,j) is d[i], the weighted degree of vertex i if if i==j, -w if i!=j and there is an edge between vertices i and j with weight w, and 0 otherwise. The weighted degree of a vertex is the sum of the weights of its adjacent edges.

Value

A numeric matrix.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

```
g <- make_ring(10)
laplacian_matrix(g)
laplacian_matrix(g, norm=TRUE)
laplacian_matrix(g, norm=TRUE, sparse=FALSE)
```

Description

Now it calls the Fruchterman-Reingold layout, with a warning.

Usage

```
layout.fruchterman.reingold.grid(graph, ...)
```
Arguments

graph Input graph.
... Extra arguments are ignored.

Value

Layout coordinates, a two column matrix.

layout.reingold.tilford

Deprecated layout functions

Description

Please use the new names, see layout_.

Usage

layout.reingold.tilford(..., params = list())
layout.circle(..., params = list())
layout.sphere(..., params = list())
layout.random(..., params = list())
layout.fruchterman.reingold(..., params = list())
layout.kamada.kawai(..., params = list())
layout.lgl(..., params = list())

Arguments

... Passed to the new layout functions.
params Passed to the new layout functions as arguments.

layout.spring

Spring layout, this was removed from igraph

Description

Now it calls the Fruchterman-Reingold layout, with a warning.

Usage

layout.spring(graph, ...)
Arguments

graph  Input graph.
...
Extra arguments are ignored.

Value

Layout coordinates, a two column matrix.

layout.svd  

SVD layout, this was removed from igraph

Description

Now it calls the Fruchterman-Reingold layout, with a warning.

Usage

layout.svd(graph, ...)

Arguments

graph  Input graph.
...
Extra arguments are ignored.

Value

Layout coordinates, a two column matrix.

layout

Graph layouts

Description

This is a generic function to apply a layout function to a graph.

Usage

layout_(graph, layout, ...)

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

## S3 method for class 'igraph_layout_modifier'
print(x, ...)
Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>graph</td>
<td>The input graph.</td>
</tr>
<tr>
<td>layout</td>
<td>The layout specification. It must be a call to a layout specification function.</td>
</tr>
<tr>
<td>...</td>
<td>Further modifiers, see a complete list below. For the print methods, it is ignored.</td>
</tr>
<tr>
<td>x</td>
<td>The layout specification</td>
</tr>
</tbody>
</table>

Details

There are two ways to calculate graph layouts in igraph. The first way is to call a layout function (they all have prefix layout_ on a graph, to get the vertex coordinates.

The second way (new in igraph 0.8.0), has two steps, and it is more flexible. First you call a layout specification function (the one without the layout_ prefix, and then layout_ (or add_layout_) to perform the layouting.

The second way is preferred, as it is more flexible. It allows operations before and after the layouting. E.g. using the component_wise argument, the layout can be calculated separately for each component, and then merged to get the final results.

Value

The return value of the layout function, usually a two column matrix. For 3D layouts a three column matrix.

Modifiers

Modifiers modify how a layout calculation is performed. Currently implemented modifiers:

- component_wise calculates the layout separately for each component of the graph, and then merges them.
- normalize scales the layout to a square.

See Also

add_layout_ to add the layout to the graph as an attribute.

Other graph layouts: add_layout_(), component_wise(), layout_as_bipartite(), layout_as_star(), layout_as_tree(), layout_in_circle(), layout_nicely(), layout_on_grid(), layout_on_sphere(), layout_randomly(), layout_with_dh(), layout_with_fr(), layout_with_gem(), layout_with_graphopt(), layout_with_kk(), layout_with_lgl(), layout_with_mds(), layout_with_sugiyama(), merge_coords(), norm_coords(), normalize()

Examples

g <- make_ring(10) + make_full_graph(5)
coords <- layout_(g, as_star())
plot(g, layout = coords)
layout_as_bipartite  Simple two-row layout for bipartite graphs

Description

Minimize edge-crossings in a simple two-row (or column) layout for bipartite graphs.

Usage

```r
layout_as_bipartite(graph, types = NULL, hgap = 1, vgap = 1, maxiter = 100)
```

Arguments

- `graph`: The bipartite input graph. It should have a logical 'type' vertex attribute, or the `types` argument must be given.
- `types`: A logical vector, the vertex types. If this argument is `NULL` (the default), then the 'type' vertex attribute is used.
- `hgap`: Real scalar, the minimum horizontal gap between vertices in the same layer.
- `vgap`: Real scalar, the distance between the two layers.
- `maxiter`: Integer scalar, the maximum number of iterations in the crossing minimization stage. 100 is a reasonable default; if you feel that you have too many edge crossings, increase this.

... Arguments to pass to `layout_as_bipartite`.

Details

The layout is created by first placing the vertices in two rows, according to their types. Then the positions within the rows are optimized to minimize edge crossings, using the Sugiyama algorithm (see `layout_with_sugiyama`).

Value

A matrix with two columns and as many rows as the number of vertices in the input graph.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

- `layout_with_sugiyama`
- Other graph layouts: `add_layout_()`, `component_wise()`, `layout_as_star()`, `layout_as_tree()`, `layout_in_circle()`, `layout_nicely()`, `layout_on_grid()`, `layout_on_sphere()`, `layout_randomly()`, `layout_with_dh()`, `layout_with_fr()`, `layout_with_gem()`, `layout_with_graphopt()`, `layout_with_kk()`, `layout_with_lgl()`, `layout_with_mds()`, `layout_with_sugiyama()`, `layout_()`, `merge_coords()`, `norm_coords()`, `normalize()`
Examples

# Random bipartite graph
inc <- matrix(sample(0:1, 50, replace = TRUE, prob=c(2,1)), 10, 5)
g <- graph_from_incidence_matrix(inc)
plot(g, layout = layout_as_bipartite,
    vertex.color=c("green","cyan")[V(g)$type+1])

# Two columns
g %>%
  add_layout_(as_bipartite()) %>%
  plot()

layout_as_star

Generate coordinates to place the vertices of a graph in a star-shape

Description

A simple layout generator, that places one vertex in the center of a circle and the rest of the vertices equidistantly on the perimeter.

Usage

layout_as_star(graph, center = V(graph)[1], order = NULL)

as_star(...)

Arguments

graph The graph to layout.
center The id of the vertex to put in the center. By default it is the first vertex.
order Numeric vector, the order of the vertices along the perimeter. The default ordering is given by the vertex ids.
... Arguments to pass to layout_as_star.

Details

It is possible to choose the vertex that will be in the center, and the order of the vertices can be also given.

Value

A matrix with two columns and as many rows as the number of vertices in the input graph.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>
See Also

layout and layout.drl for other layout algorithms, plot.igraph and tkplot on how to plot graphs and star on how to create ring graphs.

Other graph layouts: add_layout(), component_wise(), layout_as_bipartite(), layout_as_tree(), layout_in_circle(), layout_nicely(), layout_on_grid(), layout_on_sphere(), layout_randomly(), layout_with dh(), layout_with fr(), layout_with_gem(), layout_with_graphopt(), layout_with_kk(), layout_with_lgl(), layout_with_mds(), layout_with_sugiyama(), layout(), merge_coords(), norm_coords(), normalize()
rootlevel

This argument can be useful when drawing forests which are not trees (i.e. they are unconnected and have tree components). It specifies the level of the root vertices for every tree in the forest. It is only considered if the roots argument is not an empty vector.

mode

Specifies which edges to consider when building the tree. If it is 'out', then only the outgoing, if it is 'in', then only the incoming edges of a parent are considered. If it is 'all' then all edges are used (this was the behavior in igraph 0.5 and before). This parameter also influences how the root vertices are calculated, if they are not given. See the roots parameter.

flip.y

Logical scalar, whether to flip the 'y' coordinates. The default is flipping because that puts the root vertex on the top.

... 

Passed to layout_as_tree.

Details

Arranges the nodes in a tree where the given node is used as the root. The tree is directed downwards and the parents are centered above its children. For the exact algorithm, the reference below.

If the given graph is not a tree, a breadth-first search is executed first to obtain a possible spanning tree.

Value

A numeric matrix with two columns, and one row for each vertex.

Author(s)

Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

Other graph layouts: add_layout_(), component_wise(), layout_as_bipartite(), layout_as_star(), layout_in_circle(), layout_nicely(), layout_on_grid(), layout_on_sphere(), layout_randomly(), layout_with_dh(), layout_with_fr(), layout_with_gem(), layout_with_graphtopt(), layout_with_kk(), layout_with_lgl(), layout_with_mds(), layout_with_sugiyama(), layout_, merge_coords(), norm_coords(), normalize()

Examples

tree <- make_tree(20, 3)
plot(tree, layout=layout_as_tree)
plot(tree, layout=layout_as_tree(tree, flip.y=FALSE))
plot(tree, layout=layout_as_tree(tree, circular=TRUE))

```
```

tree2 <- make_tree(10, 3) + make_tree(10, 2)
plot(tree2, layout=layout_as_tree)
plot(tree2, layout=layout_as_tree(tree2, root=c(1,11),
rootlevel=c(2,1)))

```

layout_as_tree
**layout_in_circle**

Graph layout with vertices on a circle.

Description

Place vertices on a circle, in the order of their vertex ids.

Usage

```r
layout_in_circle(graph, order = V(graph))
in_circle(...)```

Arguments

- `graph`: The input graph.
- `order`: The vertices to place on the circle, in the order of their desired placement. Vertices that are not included here will be placed at (0,0).
- `...`: Passed to `layout_in_circle`.

Details

If you want to order the vertices differently, then permute them using the `permute` function.

Value

A numeric matrix with two columns, and one row for each vertex.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

Other graph layouts: `add_layout()`, `component_wise()`, `layout_as_bipartite()`, `layout_as_star()`, `layout_as_tree()`, `layout_nicely()`, `layout_on_grid()`, `layout_on_sphere()`, `layout_randomly()`, `layout_with_dh()`, `layout_with_fr()`, `layout_with_gem()`, `layout_with_graphopt()`, `layout_with_kk()`, `layout_with_lgl()`, `layout_with_mds()`, `layout_with_sugiyama()`, `layout()`, `merge_coords()`, `norm_coords()`, `normalize()`

Examples

```r
## Place vertices on a circle, order them according to their
## community
## Not run:
library(igraphdata)
data(karate)
karate_groups <- cluster_optimal(karate)
coords <- layout_in_circle(karate, order = order(membership(karate_groups)))
V(karate)$label <- sub("Actor ", ", ", V(karate)$name)
V(karate)$label.color <- membership(karate_groups)
```
V(karate)$shape <- "none"
plot(karate, layout = coords)
## End(Not run)

---

**layout_nicely**

Choose an appropriate graph layout algorithm automatically

**Description**

This function tries to choose an appropriate graph layout algorithm for the graph, automatically, based on a simple algorithm. See details below.

**Usage**

```r
layout_nicely(graph, dim = 2, ...)
nicely(...)
```

**Arguments**

- `graph` The input graph
- `dim` Dimensions, should be 2 or 3.
- `...` For `layout_nicely` the extra arguments are passed to the real layout function. For `nicely` all arguments are passed to `layout_nicely`.

**Details**

`layout_nicely` tries to choose an appropriate layout function for the supplied graph, and uses that to generate the layout. The current implementation works like this:

1. If the graph has a graph attribute called 'layout', then this is used. If this attribute is an R function, then it is called, with the graph and any other extra arguments.
2. Otherwise, if the graph has vertex attributes called 'x' and 'y', then these are used as coordinates. If the graph has an additional 'z' vertex attribute, that is also used.
3. Otherwise, if the graph is connected and has less than 1000 vertices, the Fruchterman-Reingold layout is used, by calling `layout_with_fr`.
4. Otherwise the DrL layout is used, `layout_with_drl` is called.

In layout algorithm implementations, an argument named ‘weights’ is typically used to specify the weights of the edges if the layout algorithm supports them. In this case, omitting ‘weights’ or setting it to `NULL` will make `igraph` use the ‘weight’ edge attribute from the graph if it is present. However, most layout algorithms do not support non-positive weights, so `layout_nicely` would fail if you simply called it on your graph without specifying explicit weights and the weights happened to include non-positive numbers. We strive to ensure that `layout_nicely` works out-of-the-box for most graphs, so the rule is that if you omit ‘weights’ or set it to `NULL` and `layout_nicely` would end up calling `layout_with_fr` or `layout_with_drl`, we do not forward the weights to these functions and issue a warning about this. You can use `weights = NA` to silence the warning.

**Value**

A numeric matrix with two or three columns.
layout_on_grid

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

plot.igraph

Other graph layouts: add_layout_(), component_wise(), layout_as_bipartite(), layout_as_star(),
layout_as_tree(), layout_in_circle(), layout_on_grid(), layout_on_sphere(), layout_randomly(),
layout_with_dh(), layout_with_fr(), layout_with_gem(), layout_with_graphopt(), layout_with_kk(),
layout_with_lgl(), layout_with_mds(), layout_with_sugiyama(), layout(), merge_coords(),
norm_coords(), normalize()

layout_on_grid  Simple grid layout

Description

This layout places vertices on a rectangular grid, in two or three dimensions.

Usage

layout_on_grid(graph, width = 0, height = 0, dim = 2)

on_grid(...)  
layout.grid.3d(graph, width = 0, height = 0)

Arguments

graph  The input graph.
width  The number of vertices in a single row of the grid. If this is zero or negative,
then for 2d layouts the width of the grid will be the square root of the number
of vertices in the graph, rounded up to the next integer. Similarly, it will be the
cube root for 3d layouts.
height  The number of vertices in a single column of the grid, for three dimensional
layouts. If this is zero or negative, then it is determined automatically.
dim  Two or three. Whether to make 2d or a 3d layout.
...  Passed to layout_on_grid.

Details

The function places the vertices on a simple rectangular grid, one after the other. If you want to
change the order of the vertices, then see the permute function.

Value

A two-column or three-column matrix.

Author(s)

Tamas Nepusz <ntamas@gmail.com>
layout_on_sphere

Graph layout with vertices on the surface of a sphere

Description

Place vertices on a sphere, approximately uniformly, in the order of their vertex ids.

Usage

layout_on_sphere(graph)

on_sphere(...)  

Arguments

graph            The input graph.

...               Passed to layout_on_sphere.

Details

layout_on_sphere places the vertices (approximately) uniformly on the surface of a sphere, this is thus a 3d layout. It is not clear however what “uniformly on a sphere” means.

If you want to order the vertices differently, then permute them using the permute function.

Value

A numeric matrix with three columns, and one row for each vertex.
**layout_randomly**

Randomly place vertices on a plane or in 3d space

**Description**

This function uniformly randomly places the vertices of the graph in two or three dimensions.

**Usage**

```r
layout_randomly(graph, dim = 2)
randomly(...)```

**Arguments**

- `graph`: The input graph.
- `dim`: Integer scalar, the dimension of the space to use. It must be 2 or 3.
- `...`: Parameters to pass to `layout_randomly`.

**Details**

Randomly places vertices on a [-1,1] square (in 2d) or in a cube (in 3d). It is probably a useless layout, but it can use as a starting point for other layout generators.

**Value**

A numeric matrix with two or three columns.

**Author(s)**

Gabor Csardi `<csardi.gabor@gmail.com>`

**See Also**

Other graph layouts: `add_layout_()`, `component_wise()`, `layout_as_bipartite()`, `layout_as_star()`, `layout_as_tree()`, `layout_in_circle()`, `layout_nicely()`, `layout_on_grid()`, `layout_randomly()`, `layout_with_dh()`, `layout_with_fr()`, `layout_with_gem()`, `layout_with_graphopt()`, `layout_with_kk()`, `layout_with_lgl()`, `layout_with_mds()`, `layout_with_sugiyama()`, `layout_()`, `merge_coords()`, `norm_coords()`, `normalize()`
**layout_with_dh**

The Davidson-Harel layout algorithm

**Description**

Place vertices of a graph on the plane, according to the simulated annealing algorithm by Davidson and Harel.

**Usage**

```
layout_with_dh(
  graph,
  coords = NULL,
  maxiter = 10,
  fineiter = max(10, log2(vcount(graph)));
  cool.fact = 0.75,
  weight.node.dist = 1,
  weight.border = 0,
  weight.edge.lengths = edge_density(graph)/10,
  weight.edge.crossings = 1 - sqrt(edge_density(graph)),
  weight.node.edge.dist = 0.2 * (1 - edge_density(graph))
)
```

with_dh(...)

**Arguments**

- **graph** The graph to lay out. Edge directions are ignored.
- **coords** Optional starting positions for the vertices. If this argument is not NULL then it should be an appropriate matrix of starting coordinates.
- **maxiter** Number of iterations to perform in the first phase.
- **fineiter** Number of iterations in the fine tuning phase.
- **cool.fact** Cooling factor.
- **weight.node.dist** Weight for the node-node distances component of the energy function.
- **weight.border** Weight for the distance from the border component of the energy function. It can be set to zero, if vertices are allowed to sit on the border.
- **weight.edge.lengths** Weight for the edge length component of the energy function.
- **weight.edge.crossings** Weight for the edge crossing component of the energy function.
- **weight.node.edge.dist** Weight for the node-edge distance component of the energy function.
- **...** Passed to `layout_with_dh`.

...
This function implements the algorithm by Davidson and Harel, see Ron Davidson, David Harel: Drawing Graphs Nicely Using Simulated Annealing. ACM Transactions on Graphics 15(4), pp. 301-331, 1996.

The algorithm uses simulated annealing and a sophisticated energy function, which is unfortunately hard to parameterize for different graphs. The original publication did not disclose any parameter values, and the ones below were determined by experimentation.

The algorithm consists of two phases, an annealing phase, and a fine-tuning phase. There is no simulated annealing in the second phase.

Our implementation tries to follow the original publication, as much as possible. The only major difference is that coordinates are explicitly kept within the bounds of the rectangle of the layout.

A two- or three-column matrix, each row giving the coordinates of a vertex, according to the ids of the vertex ids.

Gabor Csardi <csardi.gabor@gmail.com>


layout_with_fr, layout_with_kk for other layout algorithms.

Other graph layouts: add_layout(), component_wise(), layout_as_bipartite(), layout_as_star(), layout_as_tree(), layout_in_circle(), layout_nicely(), layout_on_grid(), layout_on_sphere(), layout_randomly(), layout_with_fr(), layout_with_gem(), layout_with_graphopt(), layout_with_kk(), layout_with_lgl(), layout_with_mds(), layout_with_sugiyama(), layout(), merge.coords(), norm.coords(), normalize()

set.seed(42)
## Figures from the paper
g_1b <- make_star(19, mode="undirected") + path(c(2:19, 2)) +
    path(c(seq(2, 18, by=2), 2))
plot(g_1b, layout=layout_with_dh)

plot(g_2, layout=layout_with_dh)

plot(g_3, layout=layout_with_dh)

plot(g_4, layout=layout_with_dh, vertex.size=5, vertex.label=NA)

plot(g_5a, layout=layout_with_dh, vertex.size=5, vertex.label=NA)

plot(g_5a, layout=layout_with_dh, vertex.size=5, vertex.label=NA)
plot(g_5a, layout=layout_with_dh, vertex.size=5, vertex.label=NA)

g_5b <- make_ring(40)
plot(g_5b, layout=layout_with_dh, vertex.size=5, vertex.label=NA)

g_6 <- make_lattice(c(2,2,2))
plot(g_6, layout=layout_with_dh)

g_7 <- graph_from_literal(1:3:5 -- 2:4:6)
plot(g_7, layout=layout_with_dh, vertex.label=V(g_7)$name)

g_8 <- make_ring(5) + make_ring(10) + make_ring(5) +
    edges(1,6, 2,8, 3, 10, 4,12, 5,14, 7,16, 9,17, 11,18, 13,19, 15,20)
plot(g_8, layout=layout_with_dh, vertex.size=5, vertex.label=NA)

g_9 <- make_lattice(c(3,2,2))
plot(g_9, layout=layout_with_dh, vertex.size=5, vertex.label=NA)

g_10 <- make_lattice(c(6,6))
plot(g_10, layout=layout_with_dh, vertex.size=5, vertex.label=NA)

g_11a <- make_tree(31, 2, mode="undirected")
plot(g_11a, layout=layout_with_dh, vertex.size=5, vertex.label=NA)

g_11b <- make_tree(21, 4, mode="undirected")
plot(g_11b, layout=layout_with_dh, vertex.size=5, vertex.label=NA)

g_12 <- make_empty_graph(n=37, directed=FALSE) +
    path(1:5,10,22,31,37:33,27,16,6,1) + path(6,7,11,9,10) + path(16:22) +
    path(27:31) + path(2,7,18,28,34) + path(3,8,11,19,29,32,35) +
    path(4,9,20,30,36) + path(1,7,12,14,19,24,26,30,37) +
    path(5,9,13,15,19,23,25,28,33) + path(3,12,16,25,35,26,22,13,3)
plot(g_12, layout=layout_with_dh, vertex.size=5, vertex.label=NA)

layout_with_drl

The DrL graph layout generator

Description

DrL is a force-directed graph layout toolbox focused on real-world large-scale graphs, developed by Shawn Martin and colleagues at Sandia National Laboratories.

Usage

layout_with_drl(
  graph,
  use.seed = FALSE,
  seed = matrix(runif(vcount(graph) * 2), ncol = 2),
  options = drl_defaults$default,
  weights = NULL,
  fixed = NULL,
  dim = 2
)
with_drl(...)

Arguments

**graph**

The input graph, in can be directed or undirected.

**use.seed**

Logical scalar, whether to use the coordinates given in the seed argument as a starting point.

**seed**

A matrix with two columns, the starting coordinates for the vertices is use.seed is TRUE. It is ignored otherwise.

**options**

Options for the layout generator, a named list. See details below.

**weights**

The weights of the edges. It must be a positive numeric vector, NULL or NA. If it is NULL and the input graph has a ‘weight’ edge attribute, then that attribute will be used. If NULL and no such attribute is present, then the edges will have equal weights. Set this to NA if the graph was a ‘weight’ edge attribute, but you don’t want to use it for the layout. Larger edge weights correspond to stronger connections.

**fixed**

Logical vector, it can be used to fix some vertices. Unfortunately this has never been implemented in the C core of the igraph library and thus it never worked. The argument is now deprecated and will be removed in igraph 1.4.0.

**dim**

Either ‘2’ or ‘3’, it specifies whether we want a two dimensional or a three dimensional layout. Note that because of the nature of the DrL algorithm, the three dimensional layout takes significantly longer to compute.

... Passed to layout_with_drl.

Details

This function implements the force-directed DrL layout generator.

The generator has the following parameters:

**edge.cut**

Edge cutting is done in the late stages of the algorithm in order to achieve less dense layouts. Edges are cut if there is a lot of stress on them (a large value in the objective function sum). The edge cutting parameter is a value between 0 and 1 with 0 representing no edge cutting and 1 representing maximal edge cutting.

**init.iterations**

Number of iterations in the first phase.

**init.temperature**

Start temperature, first phase.

**init.attraction**

Attraction, first phase.

**init.damping.mult**

Damping, first phase.

**liquid.iterations**

Number of iterations, liquid phase.

**liquid.temperature**

Start temperature, liquid phase.

**liquid.attraction**

Attraction, liquid phase.

**liquid.damping.mult**

Damping, liquid phase.

**expansion.iterations**

Number of iterations, expansion phase.

**expansion.temperature**

Start temperature, expansion phase.

**expansion.attraction**

Attraction, expansion phase.

**expansion.damping.mult**

Damping, expansion phase.

**cooldown.iterations**

Number of iterations, cooldown phase.
cooldown.temperature  Start temperature, cooldown phase.
cooldown.attraction  Attraction, cooldown phase.
cooldown.damping.mult  Damping, cooldown phase.
crunch.iterations  Number of iterations, crunch phase.
crunch.temperature  Start temperature, crunch phase.
crunch.attraction  Attraction, crunch phase.
crunch.damping.mult  Damping, crunch phase.
simmer.iterations  Number of iterations, simmer phase.
simmer.temperature  Start temperature, simmer phase.
simmer.attraction  Attraction, simmer phase.
simmer.damping.mult  Damping, simmer phase.

There are five pre-defined parameter settings as well, these are called drl_defaults$default,
  drl_defaults$coarsen, drl_defaults$coarsest, drl_defaults$refine and drl_defaults$final.

Value

A numeric matrix with two columns.

Author(s)

Shawn Martin (http://www.cs.otago.ac.nz/homepages/smartin/) and Gabor Csardi <csardi.gabor@gmail.com>
for the R/igraph interface and the three dimensional version.

References

See the following technical report: Martin, S., Brown, W.M., Klavans, R., Boyack, K.W., DrL:

See Also

layout for other layout generators.

Examples

g <- as.undirected(sample_pa(100, m=1))
l <- layout_with_drl(g, options=list(simmer.attraction=0))
## Not run:
plot(g, layout=l, vertex.size=3, vertex.label=NA)
## End(Not run)
The Fruchterman-Reingold layout algorithm

Description

Place vertices on the plane using the force-directed layout algorithm by Fruchterman and Reingold.

Usage

```
layout_with_fr(
    graph,
    coords = NULL,
    dim = 2,
    niter = 500,
    start.temp = sqrt(vcount(graph)),
    grid = c("auto", "grid", "nogrid"),
    weights = NULL,
    minx = NULL,
    maxx = NULL,
    miny = NULL,
    maxy = NULL,
    minz = NULL,
    maxz = NULL,
    coolexp,
    maxdelta,
    area,
    repulserad,
    maxiter
)
```

```
with_fr(...)```

Arguments

- **graph**: The graph to lay out. Edge directions are ignored.
- **coords**: Optional starting positions for the vertices. If this argument is not NULL then it should be an appropriate matrix of starting coordinates.
- **dim**: Integer scalar, 2 or 3, the dimension of the layout. Two dimensional layouts are places on a plane, three dimensional ones in the 3d space.
- **niter**: Integer scalar, the number of iterations to perform.
- **start.temp**: Real scalar, the start temperature. This is the maximum amount of movement allowed along one axis, within one step, for a vertex. Currently it is decreased linearly to zero during the iteration.
- **grid**: Character scalar, whether to use the faster, but less accurate grid based implementation of the algorithm. By default ("auto"), the grid-based implementation is used if the graph has more than one thousand vertices.
- **weights**: A vector giving edge weights. The weight edge attribute is used by default, if present. If weights are given, then the attraction along the edges will be multiplied by the given edge weights. This places vertices connected with a highly weighted edge closer to each other. Weights must be positive.
If not NULL, then it must be a numeric vector that gives lower boundaries for the 'x' coordinates of the vertices. The length of the vector must match the number of vertices in the graph.

Similar to minx, but gives the upper boundaries.

Similar to minx, but gives the lower boundaries of the 'y' coordinates.

Similar to minx, but gives the upper boundaries of the 'y' coordinates.

Similar to minx, but gives the lower boundaries of the 'z' coordinates.

Similar to minx, but gives the upper boundaries of the 'z' coordinates.

These arguments are not supported from igraph version 0.8.0 and are ignored (with a warning).

A deprecated synonym of niter, for compatibility.

Passed to layout_with_fr.

Details
See the referenced paper below for the details of the algorithm.
This function was rewritten from scratch in igraph version 0.8.0.

Value
A two- or three-column matrix, each row giving the coordinates of a vertex, according to the ids of the vertex ids.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

References

See Also
layout_with_drl, layout_with_kk for other layout algorithms.

Other graph layouts: add_layout_(), component_wise(), layout_as_bipartite(), layout_as_star(), layout_as_tree(), layout_in_circle(), layout_nicely(), layout_on_grid(), layout_on_sphere(), layout_randomly(), layout_with_dh(), layout_with_gem(), layout_with_graphopt(), layout_with_kk(), layout_with_lgl(), layout_with_mds(), layout_with_sugiyama(), layout(), merge_coords(), norm_coords(), normalize()

Examples
# Fixing ego
g <- sample_pa(20, m=2)
minC <- rep(-Inf, vcount(g))
maxC <- rep(Inf, vcount(g))
minC[1] <- maxC[1] <- 0
t <- layout_with_fr(g, minx=minC, maxx=maxC, 
miny=minC, maxy=maxC)
The GEM layout algorithm

Description

Place vertices on the plane using the GEM force-directed layout algorithm.

Usage

```r
layout_with_gem(
  graph,
  coords = NULL,
  maxiter = 40 * vcount(graph)^2,
  temp.max = max(vcount(graph), 1),
  temp.min = 1/10,
  temp.init = sqrt(max(vcount(graph), 1))
)
```

Arguments

- **graph**: The input graph. Edge directions are ignored.
- **coords**: If not `NULL`, then the starting coordinates should be given here, in a two or three column matrix, depending on the `dim` argument.
- **maxiter**: The maximum number of iterations to perform. Updating a single vertex counts as an iteration. A reasonable default is `40 * n * n`, where `n` is the number of vertices. The original paper suggests `4 * n * n`, but this usually only works if the other parameters are set up carefully.
- **temp.max**: The maximum allowed local temperature. A reasonable default is the number of vertices.
- **temp.min**: The global temperature at which the algorithm terminates (even before reaching `maxiter` iterations). A reasonable default is `1/10`.
- **temp.init**: Initial local temperature of all vertices. A reasonable default is the square root of the number of vertices.
- **...**: Passed to `layout_with_gem`.

Details

See the referenced paper below for the details of the algorithm.
Value

A numeric matrix with two columns, and as many rows as the number of vertices.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

layout_with_fr, plot.igraph, tkplot

Other graph layouts: add_layout_().component_wise(), layout_as_bipartite(), layout_as_star(), layout_as_tree(), layout_in_circle(), layout_nicely(), layout_on_grid(), layout_on_sphere(), layout_randomly(), layout_with_dh(), layout_with_fr(), layout_with_graphopt(), layout_with_kk(), layout_with_lgl(), layout_with_mds(), layout_with_sugiyama(), layout(), merge_coords(), norm_coords(), normalize()

Examples

set.seed(42)
g <- make_ring(10)
plot(g, layout=layout_with_gem)

layout_with_graphopt

The graphopt layout algorithm

Description

A force-directed layout algorithm, that scales relatively well to large graphs.

Usage

layout_with_graphopt(
  graph,
  start = NULL,
  niter = 500,
  charge = 0.001,
  mass = 30,
  spring.length = 0,
  spring_constant = 1,
  max.sa.movement = 5
)

with_graphopt(...)
Arguments

- **graph**: The input graph.
- **start**: If given, then it should be a matrix with two columns and one line for each vertex. This matrix will be used as starting positions for the algorithm. If not given, then a random starting matrix is used.
- **niter**: Integer scalar, the number of iterations to perform. Should be a couple of hundred in general. If you have a large graph then you might want to only do a few iterations and then check the result. If it is not good enough you can feed it in again in the start argument. The default value is 500.
- **charge**: The charge of the vertices, used to calculate electric repulsion. The default is 0.001.
- **mass**: The mass of the vertices, used for the spring forces. The default is 30.
- **spring.length**: The length of the springs, an integer number. The default value is zero.
- **spring.constant**: The spring constant, the default value is one.
- **max.sa.movement**: Real constant, it gives the maximum amount of movement allowed in a single step along a single axis. The default value is 5.
- ... Passed to `layout_with_graphopt`.

Details

`layout_with_graphopt` is a port of the graphopt layout algorithm by Michael Schmuhl. Graphopt version 0.4.1 was rewritten in C and the support for layers was removed (might be added later) and a code was a bit reorganized to avoid some unnecessary steps is the node charge (see below) is zero.

Graphopt uses physical analogies for defining attracting and repelling forces among the vertices and then the physical system is simulated until it reaches an equilibrium. (There is no simulated annealing or anything like that, so a stable fixed point is not guaranteed.)

See also [http://www.schmuhl.org/graphopt/](http://www.schmuhl.org/graphopt/) for the original graphopt.

Value

A numeric matrix with two columns, and a row for each vertex.

Author(s)

Michael Schmuhl for the original graphopt code, rewritten and wrapped by Gabor Csardi <csardi.gabor@gmail.com>.

See Also

Other graph layouts: `add_layout()`, `component_wise()`, `layout_as_bipartite()`, `layout_as_star()`, `layout_as_tree()`, `layout_in_circle()`, `layout_nicely()`, `layout_on_grid()`, `layout_on_sphere()`, `layout_randomly()`, `layout_with_dh()`, `layout_with_fr()`, `layout_with_gem()`, `layout_with_kk()`, `layout_with_lgl()`, `layout_with_mds()`, `layout_with_sugiyama()`, `layout()`, `merge.coords()`, `norm_coords()`, `normalize()`
The Kamada-Kawai layout algorithm

Description

Place the vertices on the plane, or in 3D space, based on a physical model of springs.

Usage

```r
layout_with_kk(
  graph,
  coords = NULL,
  dim = 2,
  maxiter = 50 * vcount(graph),
  epsilon = 0,
  kkconst = max(vcount(graph), 1),
  weights = NULL,
  minx = NULL,
  maxx = NULL,
  miny = NULL,
  maxy = NULL,
  minz = NULL,
  maxz = NULL,
  niter,
  sigma,
  initprop,
  coolprop,
  start
)
```

```r
with_kk(...)```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>graph</td>
<td>The input graph. Edge directions are ignored.</td>
</tr>
<tr>
<td>coords</td>
<td>If not NULL, then the starting coordinates should be given here, in a two or three column matrix, depending on the dim argument.</td>
</tr>
<tr>
<td>dim</td>
<td>Integer scalar, 2 or 3, the dimension of the layout. Two dimensional layouts are places on a plane, three dimensional ones in the 3d space.</td>
</tr>
<tr>
<td>maxiter</td>
<td>The maximum number of iterations to perform. The algorithm might terminate earlier, see the epsilon argument.</td>
</tr>
<tr>
<td>epsilon</td>
<td>Numeric scalar, the algorithm terminates, if the maximal delta is less than this. (See the reference below for what delta means.) If you set this to zero, then the function always performs maxiter iterations.</td>
</tr>
<tr>
<td>kkconst</td>
<td>Numeric scalar, the Kamada-Kawai vertex attraction constant. Typical (and default) value is the number of vertices.</td>
</tr>
<tr>
<td>weights</td>
<td>Edge weights, larger values will result longer edges. Note that this is opposite to <code>layout_with_fr</code>. Weights must be positive.</td>
</tr>
</tbody>
</table>
If not NULL, then it must be a numeric vector that gives lower boundaries for the 'x' coordinates of the vertices. The length of the vector must match the number of vertices in the graph.

Similar to minx, but gives the upper boundaries.

Similar to minx, but gives the lower boundaries of the 'y' coordinates.

Similar to minx, but gives the upper boundaries of the 'y' coordinates.

Similar to minx, but gives the lower boundaries of the 'z' coordinates.

Similar to minx, but gives the upper boundaries of the 'z' coordinates.

These arguments are not supported from igraph version 0.8.0 and are ignored (with a warning).

Depreciated synonym for coords, for compatibility.

Passed to layout_with_kk.

See the referenced paper below for the details of the algorithm.

This function was rewritten from scratch in igraph version 0.8.0 and it follows truthfully the original publication by Kamada and Kawai now.

A numeric matrix with two (dim=2) or three (dim=3) columns, and as many rows as the number of vertices, the x, y and potentially z coordinates of the vertices.

Gabor Csardi &lt;csardi.gabor@gmail.com&gt;


Other graph layouts: add_layout_(), component_wise(), layout_as_bipartite(), layout_as_star(), layout_as_tree(), layout_in_circle(), layout_nicely(), layout_on_grid(), layout_on_sphere(), layout_randomly(), layout_with_dh(), layout_with_fr(), layout_with_gem(), layout_with_graphopt(), layout_with_lgl(), layout_with_mds(), layout_with_sugiyama(), layout(), merge_coords(), norm_coords(), normalize()

g <- make_ring(10)
E(g)$weight <- rep(1:2, length.out=ecount(g))
plot(g, layout=layout_with_kk, edge.label=E(g)$weight)
layout_with_lgl

Large Graph Layout

Description

A layout generator for larger graphs.

Usage

```r
layout_with_lgl(
  graph,
  maxiter = 150,
  maxdelta = vcount(graph),
  area = vcount(graph)^2,
  coolexp = 1.5,
  repulserad = area * vcount(graph),
  cellsize = sqrt(sqrt(area)),
  root = NULL
)
```

with_lgl(...)

Arguments

- `graph`: The input graph
- `maxiter`: The maximum number of iterations to perform (150).
- `maxdelta`: The maximum change for a vertex during an iteration (the number of vertices).
- `area`: The area of the surface on which the vertices are placed (square of the number of vertices).
- `coolexp`: The cooling exponent of the simulated annealing (1.5).
- `repulserad`: Cancellation radius for the repulsion (the `area` times the number of vertices).
- `cellsizie`: The size of the cells for the grid. When calculating the repulsion forces between vertices only vertices in the same or neighboring grid cells are taken into account (the fourth root of the number of `area`.
- `root`: The id of the vertex to place at the middle of the layout. The default value is -1 which means that a random vertex is selected.
- `...`: Passed to `layout_with_lgl`.

Details

`layout_with_lgl` is for large connected graphs, it is similar to the layout generator of the Large Graph Layout software (http://lgl.sourceforge.net/).

Value

A numeric matrix with two columns and as many rows as vertices.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>
See Also

Other graph layouts: add_layout_(), component_wise(), layout_as_bipartite(), layout_as_star(), layout_as_tree(), layout_in_circle(), layout_nicely(), layout_on_grid(), layout_on_sphere(), layout_randomly(), layout_with_dh(), layout_with_fr(), layout_with_gem(), layout_with_graphopt(), layout_with_kk(), layout_with_mds(), layout_with_sugiyama(), layout(), merge_coords(), norm_coords(), normalize()

layout_with_mds

Graph layout by multidimensional scaling

Description

Multidimensional scaling of some distance matrix defined on the vertices of a graph.

Usage

layout_with_mds(graph, dist = NULL, dim = 2, options = arpack_defaults)

Arguments

graph The input graph.
dist The distance matrix for the multidimensional scaling. If NULL (the default), then the unweighted shortest path matrix is used.
dim layout_with_mds supports dimensions up to the number of nodes minus one, but only if the graph is connected; for unconnected graphs, the only possible values is 2. This is because merge_coords only works in 2D.
options This is currently ignored, as ARPACK is not used any more for solving the eigenproblem
... Passed to layout_with_mds.

Details

layout_with_mds uses metric multidimensional scaling for generating the coordinates. Multidimensional scaling aims to place points from a higher dimensional space in a (typically) 2 dimensional plane, so that the distance between the points are kept as much as this is possible.

By default igraph uses the shortest path matrix as the distances between the nodes, but the user can override this via the dist argument.

This function generates the layout separately for each graph component and then merges them via merge_coords.

Value

A numeric matrix with dim columns.

Author(s)

Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com>
References


See Also

`layout, plot.igraph`


Examples

```r
   g <- sample_gnp(100, 2/100)
   l <- layout_with_mds(g)
   plot(g, layout=l, vertex.label=NA, vertex.size=3)
```

---

**layout_with_sugiyama**  
*The Sugiyama graph layout generator*

Description

Sugiyama layout algorithm for layered directed acyclic graphs. The algorithm minimized edge crossings.

Usage

```r
   layout_with_sugiyama(
       graph,  
       layers = NULL,  
       hgap = 1,  
       vgap = 1,  
       maxiter = 100,  
       weights = NULL,  
       attributes = c("default", "all", "none")
   )
```

with_sugiyama(...)

Arguments

- **graph**: The input graph.
- **layers**: A numeric vector or NULL. If not NULL, then it should specify the layer index of the vertices. Layers are numbered from one. If NULL, then igraph calculates the layers automatically.
- **hgap**: Real scalar, the minimum horizontal gap between vertices in the same layer.
- **vgap**: Real scalar, the distance between layers.
maxiter  Integer scalar, the maximum number of iterations in the crossing minimization stage. 100 is a reasonable default; if you feel that you have too many edge crossings, increase this.

weights  Optional edge weight vector. If NULL, then the ‘weight’ edge attribute is used, if there is one. Supply NA here and igraph ignores the edge weights. These are used only if the graph contains cycles; igraph will tend to reverse edges with smaller weights when breaking the cycles.

attributes  Which graph/vertex/edge attributes to keep in the extended graph. ‘default’ keeps the ‘size’, ‘size2’, ‘shape’, ‘label’ and ‘color’ vertex attributes and the ‘arrow.mode’ and ‘arrow.size’ edge attributes. ‘all’ keep all graph, vertex and edge attributes, ‘none’ keeps none of them.

...  Passed to layout_with_sugiyama.

Details

This layout algorithm is designed for directed acyclic graphs where each vertex is assigned to a layer. Layers are indexed from zero, and vertices of the same layer will be placed on the same horizontal line. The X coordinates of vertices within each layer are decided by the heuristic proposed by Sugiyama et al. to minimize edge crossings.

You can also try to lay out undirected graphs, graphs containing cycles, or graphs without an a priori layered assignment with this algorithm. igraph will try to eliminate cycles and assign vertices to layers, but there is no guarantee on the quality of the layout in such cases.

The Sugiyama layout may introduce “bends” on the edges in order to obtain a visually more pleasing layout. This is achieved by adding dummy nodes to edges spanning more than one layer. The resulting layout assigns coordinates not only to the nodes of the original graph but also to the dummy nodes. The layout algorithm will also return the extended graph with the dummy nodes.

For more details, see the reference below.

Value

A list with the components:

- layout  The layout, a two-column matrix, for the original graph vertices.
- layout.dummy  The layout for the dummy vertices, a two column matrix.
- extd_graph  The original graph, extended with dummy vertices. The ‘dummy’ vertex attribute is set on this graph, it is a logical attributes, and it tells you whether the vertex is a dummy vertex. The ‘layout’ graph attribute is also set, and it is the layout matrix for all (original and dummy) vertices.

Author(s)

Tamas Nepusz <ntamas@gmail.com>

References

See Also

Other graph layouts: add_layout_(), component_wise(), layout_as_bipartite(), layout_as_star(), layout_as_tree(), layout_in_circle(), layout_nicely(), layout_on_grid(), layout_on_sphere(), layout_randomly(), layout_with_dh(), layout_with_fr(), layout_with_gem(), layout_with_graphopt(), layout_with_kk(), layout_with_lgl(), layout_with_mds(), layout(), merge_coords(), norm_coords(), normalize()

Examples

DC <- graph_from_literal("DC++" ->
    "LinuxDC++":"EiskaltDC++":"StrongDC++":"DiCe!++",
    "LinuxDC++" -> "FreeDC++", "BCDC++" -> "StrongDC++",
    "FreeDC++" -> "BMDC++":"EiskaltDC++",
    "StrongDC++" -> "AirDC++":"zK++":"ApexDC++":"TkDC++",
    "StrongDC++" -> "StrongDC++ SQLite":"RSX++",
    "ApexDC++" -> "FlylinkDC++ ver <= 4xx",
    "ApexDC++" -> "ApexDC++ Speed-Mod":"DiCe!++",
    "StrongDC++ SQLite" -> "FlylinkDC++ ver >= 5xx",
    "ApexDC++ Speed-Mod" -> "FlylinkDC++ ver <= 4xx",
    "ApexDC++ Speed-Mod" -> "GreylinkDC++",
    "FlylinkDC++ ver <= 4xx" -> "FlylinkDC++ ver >= 5xx",
    "FlylinkDC++ ver <= 4xx" -> AvaLink,
    "GreylinkDC++" -> AvaLink:"RayLinkDC++":"SparkDC++":PeLink)

## Use edge types
E(DC)$lty <- 1
E(DC)["BCDC++" %->% "StrongDC++]$lty <- 2
E(DC)["FreeDC++" %->% "EiskaltDC++]$lty <- 2
E(DC)["ApexDC++" %->% "FlylinkDC++ ver <= 4xx"]$lty <- 2
E(DC)["ApexDC++" %->% "DiCe!++]$lty <- 2
E(DC)["StrongDC++ SQLite" %->% "FlylinkDC++ ver >= 5xx"]$lty <- 2
E(DC)["GreylinkDC++" %->% "AvaLink"]$lty <- 2

## Layers, as on the plot
layers <- list(c("DC++"),
    c("LinuxDC++", "BCDC++"),
    c("FreeDC++", "StrongDC++"),
    c("BMDC++", "EiskaltDC++", "AirDC++":"zK++":"ApexDC++":"TkDC++", "RSX++"),
    c("StrongDC++ SQLite", "ApexDC++ Speed-Mod":"DiCe!++"),
    c("FlylinkDC++ ver <= 4xx", "GreylinkDC++"),
    c("FlylinkDC++ ver >= 5xx", "AvaLink", "RayLinkDC++", "SparkDC++", "PeLink")

## Check that we have all nodes
all(sort(unlist(layers)) == sort(V(DC)$name))

## Add some graphical parameters
V(DC)$color <- "white"
V(DC)$shape <- "rectangle"
V(DC)$size <- 20
V(DC)$size2 <- 10
V(DC)$label <- lapply(V(DC)$name, function(x) paste(strwrap(x, 12), collapse="\n"))
E(DC)$arrow.size <- 0.5
## Create a similar layout using the predefined layers

```r
lay1 <- layout_with_sugiyama(DC, layers=apply(sapply(layers,
    function(x) V(DC)$name %in% x), 1, which))
```

## Simple plot, not very nice

```r
par(mar=rep(.1, 4))
plot(DC, layout=lay1$layout, vertex.label.cex=0.5)
```

## Sugiyama plot

```r
plot(lay1$extd_graph, vertex.label.cex=0.5)
```

## The same with automatic layer calculation

```r
## Keep vertex/edge attributes in the extended graph
lay2 <- layout_with_sugiyama(DC, attributes="all")
plot(lay2$extd_graph, vertex.label.cex=0.5)
```

## Another example, from the following paper:

```r
## Markus Eiglsperger, Martin Siebenhaller, Michael Kaufmann:
## An Efficient Implementation of Sugiyama's Algorithm for
## Layered Graph Drawing, Journal of Graph Algorithms and
## Applications 9, 305--325 (2005).
```

```r
ex <- graph_from_literal( 0 -+ 29: 6: 5:20: 4,
1 -+ 12,
2 -+ 23: 8,
3 -+ 4,
4,
5 -+ 2:10:14:26: 4: 3,
6 -+ 9:29:25:21:13,
7,
8 -+ 20:16,
9 -+ 28: 4,
10 -+ 27,
11 -+ 9:16,
12 -+ 9:19,
13 -+ 20,
14 -+ 10,
15 -+ 16:27,
16 -+ 27,
17 -+ 3,
18 -+ 13,
19 -+ 9,
20 -+ 4,
21 -+ 22,
22 -+ 8: 9,
23 -+ 9:24,
24 -+ 12:15:28,
25 -+ 11,
26 -+ 18,
27 -+ 13:19,
28 -+ 7,
29 -+ 25 )
```

```r
layers <- list( 0, c(5, 17), c(2, 14, 26, 3), c(23, 10, 18), c(1, 24),
12, 6, c(29,21), c(25,22), c(11,8,15), 16, 27, c(13,19),
c(9, 20), c(4, 28), 7 )
```
local_scan <- layout_with_sugiyama(ex, layers=apply(sapply(layers,
    function(x) V(ex)$name %in% as.character(x)),
    1, which))

origvert <- c(rep(TRUE, vcount(ex)), rep(FALSE, nrow(layex$layout.dummy)))
realedge <- as_edgelist(layex$extd_graph)[,2] <= vcount(ex)
plot(layex$extd_graph, vertex.label.cex=0.5,
    edge.arrow.size=.5,
    vertex.size=ifelse(origvert, 5, 0),
    vertex.shape=ifelse(origvert, "square", "none"),
    vertex.label=ifelse(origvert, V(ex)$name, ""),
    edge.arrow.mode=ifelse(realedge, 2, 0))

---

**local_scan**

*Compute local scan statistics on graphs*

**Description**

The scan statistic is a summary of the locality statistics that is computed from the local neighborhood of each vertex. The `local_scan` function computes the local statistics for each vertex for a given neighborhood size and the statistic function.

**Usage**

```r
local_scan(
  graph.us, graph.them = NULL,
  k = 1,
  FUN = NULL,
  weighted = FALSE,
  mode = c("out", "in", "all"),
  neighborhoods = NULL,
  ...
)
```

**Arguments**

- `graph.us`
  - An igraph object, the graph for which the scan statistics will be computed.

- `graph.them`
  - An igraph object or NULL, if not NULL, then the ‘them’ statistics is computed, i.e. the neighborhoods calculated from `graph.us` are evaluated on `graph.them`.

- `k`
  - An integer scalar, the size of the local neighborhood for each vertex. Should be non-negative.

- `FUN`
  - Character, a function name, or a function object itself, for computing the local statistic in each neighborhood. If NULL (the default value), `ecount` is used for unweighted graphs (if `weighted=FALSE`) and a function that computes the sum of edge weights is used for weighted graphs (if `weighted=TRUE`). This argument is ignored if `k` is zero.
weighted Logical scalar, TRUE if the edge weights should be used for computation of the scan statistic. If TRUE, the graph should be weighted. Note that this argument is ignored if FUN is not NULL, "ecount" and "sumweights".

mode Character scalar, the kind of neighborhoods to use for the calculation. One of 'out', 'in', 'all' or 'total'. This argument is ignored for undirected graphs.

neighborhoods A list of neighborhoods, one for each vertex, or NULL. If it is not NULL, then the function is evaluated on the induced subgraphs specified by these neighborhoods.

In theory this could be useful if the same graph.us graph is used for multiple graph.them arguments. Then the neighborhoods can be calculated on graph.us and used with multiple graphs. In practice, this is currently slower than simply using graph.them multiple times.

Arguments passed to FUN, the function that computes the local statistics.

Details
See the given reference below for the details on the local scan statistics.

local_scan calculates exact local scan statistics.

If graph.them is NULL, then local_scan computes the 'us' variant of the scan statistics. Otherwise, graph.them should be an igraph object and the 'them' variant is computed using graph.us to extract the neighborhood information, and applying FUN on these neighborhoods in graph.them.

Value
For local_scan typically a numeric vector containing the computed local statistics for each vertex. In general a list or vector of objects, as returned by FUN.

References

See Also
Other scan statistics: scan_stat()

Examples
pair <- sample_correlated_gnp_pair(n = 10^3, corr = 0.8, p = 0.1)
local_0_us <- local_scan(graph.us = pair$graph1, k = 0)
local_1_us <- local_scan(graph.us = pair$graph1, k = 1)

local_0_them <- local_scan(graph.us = pair$graph1,
                           graph.them = pair$graph2, k = 0)
local_1_them <- local_scan(graph.us = pair$graph1,
                           graph.them = pair$graph2, k = 1)

Neigh_1 <- neighborhood(pair$graph1, order = 1)
local_1_them_nhood <- local_scan(graph.us = pair$graph1,
                                  graph.them = pair$graph2,
                                  neighborhoods = Neigh_1)
make_  

Make a new graph

Description

This is a generic function for creating graphs.

Usage

make_(...)  

Arguments

...  Parameters, see details below.

Details

make_ is a generic function for creating graphs. For every graph constructor in igraph that has a make_ prefix, there is a corresponding function without the prefix: e.g. for make_ring there is also ring, etc.

The same is true for the random graph samplers, i.e. for each constructor with a sample_ prefix, there is a corresponding function without that prefix.

These shorter forms can be used together with make_. The advantage of this form is that the user can specify constructor modifiers which work with all constructors. E.g. the with_vertex_ modifier adds vertex attributes to the newly created graphs.

See the examples and the various constructor modifiers below.

See Also

simplified with_edge_ with_graph_ with_vertex_ without_loops without_multiples

Examples

r <- make_(ring(10))
l <- make_(lattice(c(3, 3, 3))

r2 <- make_(ring(10), with_vertex_(color = "red", name = LETTERS[1:10]))
l2 <- make_(lattice(c(3, 3, 3)), with_edge_(weight = 2))

ran <- sample_(degseq(c(3,3,3,3,3,3), method = "simple"), simplified())
deregree(ran)
is_simple(ran)
**Description**

`make_chordal_ring` creates an extended chordal ring. An extended chordal ring is regular graph, each node has the same degree. It can be obtained from a simple ring by adding some extra edges specified by a matrix. Let \( p \) denote the number of columns in the \( W \) matrix. The extra edges of vertex \( i \) are added according to column \( i \mod p \) in \( W \). The number of extra edges is the number of rows in \( W \): for each row \( j \) an edge \( i -> i + w[i][j] \) is added if \( i + w[i][j] \) is less than the number of total nodes. See also Kotsis, G: Interconnection Topologies for Parallel Processing Systems, PARS Mitteilungen 11, 1-6, 1993.

**Usage**

```r
make_chordal_ring(n, w, directed = FALSE)
chordal_ring(...)```

**Arguments**

- `n` The number of vertices.
- `w` A matrix which specifies the extended chordal ring. See details below.
- `directed` Logical scalar, whether or not to create a directed graph.
- `...` Passed to `make_chordal_ring`.

**Value**

An igraph graph.

**See Also**

Other deterministic constructors: `graph_from_atlas()`, `graph_from_edgelist()`, `graph_from_literal()`, `make_empty_graph()`, `make_full_citation_graph()`, `make_full_graph()`, `make_graph()`, `make_lattice()`, `make_ring()`, `make_star()`, `make_tree()`

**Examples**

```r
chord <- make_chordal_ring(15,
  matrix(c(3, 12, 4, 7, 8, 11), nr = 2))```
make_clusters  Creates a communities object.

Description

This is useful to integrate the results of community finding algorithms that are not included in igraph.

Usage

make_clusters(
  graph,
  membership = NULL,
  algorithm = NULL,
  merges = NULL,
  modularity = TRUE
)

Arguments

- **graph**: The graph of the community structure.
- **membership**: The membership vector of the community structure, a numeric vector denoting the id of the community for each vertex. It might be NULL for hierarchical community structures.
- **algorithm**: Character string, the algorithm that generated the community structure, it can be arbitrary.
- **merges**: A merge matrix, for hierarchical community structures (or NULL otherwise.
- **modularity**: Modularity value of the community structure. If this is TRUE and the membership vector is available, then it the modularity values is calculated automatically.

Value

A communities object.

make_de_bruijn_graph  De Bruijn graphs

Description

De Bruijn graphs are labeled graphs representing the overlap of strings.

Usage

make_de_bruijn_graph(m, n)

de_bruijn_graph(...)
**make_empty_graph**

**Arguments**

- **m**: Integer scalar, the size of the alphabet. See details below.
- **n**: Integer scalar, the length of the labels. See details below.
- **...**: Passed to `make_de_bruijn_graph`.

**Details**

A de Bruijn graph represents relationships between strings. An alphabet of \(m\) letters are used and strings of length \(n\) are considered. A vertex corresponds to every possible string and there is a directed edge from vertex \(v\) to vertex \(w\) if the string of \(v\) can be transformed into the string of \(w\) by removing its first letter and appending a letter to it.

Please note that the graph will have \(m^n\) vertices and even more edges, so probably you don’t want to supply too big numbers for \(m\) and \(n\).

De Bruijn graphs have some interesting properties, please see another source, e.g. Wikipedia for details.

**Value**

A graph object.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**See Also**

`make_kautz_graph`, `make_line_graph`

**Examples**

```r
# de Bruijn graphs can be created recursively by line graphs as well
g <- make_de_bruijn_graph(2,1)
make_de_bruijn_graph(2,2)
make_line_graph(g)
```

---

**make_empty_graph**

**A graph with no edges**

**Description**

A graph with no edges

**Usage**

```r
make_empty_graph(n = 0, directed = TRUE)

empty_graph(...)
```
Arguments

- `n` Number of vertices.
- `directed` Whether to create a directed graph.
- `...` Passed to `make_graph_empty`.

Value

An igraph graph.

See Also

Other deterministic constructors: `graph_from_atlas()`, `graph_from_edgelist()`, `graph_from_literal()`, `make_chordal_ring()`, `make_full_citation_graph()`, `make_full_graph()`, `make_graph()`, `make_lattice()`, `make_ring()`, `make_star()`, `make_tree()`

Examples

```r
make_empty_graph(n = 10)
make_empty_graph(n = 5, directed = FALSE)
```

Description

`make_from_prufer` creates an undirected tree graph from its Prufer sequence.

Usage

```r
make_from_prufer(prufer)
from_prufer(...)```

Arguments

- `prufer` The Prufer sequence to convert into a graph
- `...` Passed to `make_from_prufer`

Details

The Prufer sequence of a tree graph with `n` labeled vertices is a sequence of `n-2` numbers, constructed as follows. If the graph has more than two vertices, find a vertex with degree one, remove it from the tree and add the label of the vertex that it was connected to to the sequence. Repeat until there are only two vertices in the remaining graph.

Value

A graph object.

See Also

`to_prufer` to convert a graph into its Prufer sequence
make_full_bipartite_graph

Create a full bipartite graph

Description

Bipartite graphs are also called two-mode by some. This function creates a bipartite graph in which every possible edge is present.

Usage

```r
make_full_bipartite_graph(
  n1, n2,
  directed = FALSE,
  mode = c("all", "out", "in")
)
```

Arguments

- `n1`: The number of vertices of the first kind.
- `n2`: The number of vertices of the second kind.
- `directed`: Logical scalar, whether the graphs is directed.
- `mode`: Scalar giving the kind of edges to create for directed graphs. If this is ‘out’ then all vertices of the first kind are connected to the others; ‘in’ specifies the opposite direction; ‘all’ creates mutual edges. This argument is ignored for undirected graphs.
- `...`: Passed to `make_full_bipartite_graph`.

Details

Bipartite graphs have a ‘type’ vertex attribute in igraph, this is boolean and FALSE for the vertices of the first kind and TRUE for vertices of the second kind.

Value

An igraph graph, with the ‘type’ vertex attribute set.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>
make_full_citation_graph

Create a complete (full) citation graph

Description

make_full_citation_graph creates a full citation graph. This is a directed graph, where every \( i \rightarrow j \) edge is present if and only if \( j < i \). If directed=FALSE then the graph is just a full graph.

Usage

make_full_citation_graph(n, directed = TRUE)

full_citation_graph(...)

Arguments

n The number of vertices.
directed Whether to create a directed graph.
... Passed to make_full_citation_graph.

Value

An igraph graph.

See Also

Other deterministic constructors: graph_from_atlas(), graph_from_edgelist(), graph_from_literal(), make_chordal_ring(), make_empty_graph(), make_full_graph(), make_graph(), make_lattice(), make_ring(), make_star(), make_tree()

Examples

print_all(make_full_citation_graph(10))
**make_full_graph**

Create a full graph

### Usage

```
make_full_graph(n, directed = FALSE, loops = FALSE)
full_graph(...)
```

### Arguments

- `n` Number of vertices.
- `directed` Whether to create a directed graph.
- `loops` Whether to add self-loops to the graph.
- `...` Passed to `make_full_graph`.

### Value

An igraph graph

### See Also

Other deterministic constructors: `graph_from_atlas()`, `graph_from_edgelist()`, `graph_from_literal()`, `make_chordal_ring()`, `make_empty_graph()`, `make_full_citation_graph()`, `make_graph()`, `make_lattice()`, `make_ring()`, `make_star()`, `make_tree()`

### Examples

```
make_full_graph(5)
print_all(make_full_graph(4, directed = TRUE))
```

---

**make_graph**

Create an igraph graph from a list of edges, or a notable graph

### Description

Create an igraph graph from a list of edges, or a notable graph
make_graph

Usage

make_graph(
  edges,
  ...,  
  n = max(edges),
  isolates = NULL,
  directed = TRUE,
  dir = directed,
  simplify = TRUE
)

make_directed_graph(edges, n = max(edges))

make_undirected_graph(edges, n = max(edges))

directed_graph(...)  
undirected_graph(...)  

Arguments

dges
  A vector defining the edges, the first edge points from the first element to the second, the second edge from the third to the fourth, etc. For a numeric vector, these are interpreted as internal vertex ids. For character vectors, they are interpreted as vertex names.
  Alternatively, this can be a character scalar, the name of a notable graph. See Notable graphs below. The name is case insensitive.
  Starting from igraph 0.8.0, you can also include literals here, via igraph’s formula notation (see \texttt{graph_from_literal}). In this case, the first term of the formula has to start with a ‘\texttt{~}’ character, just like regular formulae in R. See examples below.

\ldots
  For make\_graph: extra arguments for the case when the graph is given via a literal, see \texttt{graph_from_literal}. For directed\_graph and undirected\_graph: Passed to make\_directed\_graph or make\_undirected\_graph.

n
  The number of vertices in the graph. This argument is ignored (with a warning) if edges are symbolic vertex names. It is also ignored if there is a bigger vertex id in edges. This means that for this function it is safe to supply zero here if the vertex with the largest id is not an isolate.

isolates
  Character vector, names of isolate vertices, for symbolic edge lists. It is ignored for numeric edge lists.

directed
  Whether to create a directed graph.

dir
  It is the same as directed, for compatibility. Do not give both of them.

simplify
  For graph literals, whether to simplify the graph.

Value

An igraph graph.
Notable graphs

`make_graph` can create some notable graphs. The name of the graph (case insensitive), a character scalar must be supplied as the edges argument, and other arguments are ignored. (A warning is given is they are specified.)

`make_graph` knows the following graphs:

- **Bull** The bull graph, 5 vertices, 5 edges, resembles to the head of a bull if drawn properly.
- **Chvatal** This is the smallest triangle-free graph that is both 4-chromatic and 4-regular. According to the Grunbaum conjecture there exists an m-regular, m-chromatic graph with n vertices for every m>1 and n>2. The Chvatal graph is an example for m=4 and n=12. It has 24 edges.
- **Coxeter** A non-Hamiltonian cubic symmetric graph with 28 vertices and 42 edges.
- **Cubical** The Platonic graph of the cube. A convex regular polyhedron with 8 vertices and 12 edges.
- **Diamond** A graph with 4 vertices and 5 edges, resembles to a schematic diamond if drawn properly.
- **Dodecahedral, Dodecahedron** Another Platonic solid with 20 vertices and 30 edges.
- **Folkman** The semisymmetric graph with minimum number of vertices, 20 and 40 edges. A semisymmetric graph is regular, edge transitive and not vertex transitive.
- **Franklin** This is a graph whose embedding to the Klein bottle can be colored with six colors, it is a counterexample to the necessity of the Heawood conjecture on a Klein bottle. It has 12 vertices and 18 edges.
- **Frucht** The Frucht Graph is the smallest cubical graph whose automorphism group consists only of the identity element. It has 12 vertices and 18 edges.
- **Grotzsch** The Groetzsch graph is a triangle-free graph with 11 vertices, 20 edges, and chromatic number 4. It is named after German mathematician Herbert Groetzsch, and its existence demonstrates that the assumption of planarity is necessary in Groetzsch’s theorem that every triangle-free planar graph is 3-colorable.
- **Heawood** The Heawood graph is an undirected graph with 14 vertices and 21 edges. The graph is cubic, and all cycles in the graph have six or more edges. Every smaller cubic graph has shorter cycles, so this graph is the 6-cage, the smallest cubic graph of girth 6.
- **Herschel** The Herschel graph is the smallest nonhamiltonian polyhedral graph. It is the unique such graph on 11 nodes, and has 18 edges.
- **House** The house graph is a 5-vertex, 6-edge graph, the schematic draw of a house if drawn properly, basically a triangle of the top of a square.
- **HouseX** The same as the house graph with an X in the square. 5 vertices and 8 edges.
- **Icosahedral, Icosahedron** A Platonic solid with 12 vertices and 30 edges.
- **Levi** The graph is a 4-arc transitive cubic graph, it has 30 vertices and 45 edges.
- **McGee** The McGee graph is the unique 3-regular 7-cage graph, it has 24 vertices and 36 edges.
- **Meredith** The Meredith graph is a quartic graph on 70 nodes and 140 edges that is a counterexample to the conjecture that every 4-regular 4-connected graph is Hamiltonian.
- **Noperfectmatching** A connected graph with 16 vertices and 27 edges containing no perfect matching. A matching in a graph is a set of pairwise non-adjacent edges; that is, no two edges share a common vertex. A perfect matching is a matching which covers all vertices of the graph.
**Nonline** A graph whose connected components are the 9 graphs whose presence as a vertex-induced subgraph in a graph makes a nonline graph. It has 50 vertices and 72 edges.

**Octahedral, Octahedron** Platonic solid with 6 vertices and 12 edges.

**Petersen** A 3-regular graph with 10 vertices and 15 edges. It is the smallest hypohamiltonian graph, i.e. it is non-hamiltonian but removing any single vertex from it makes it Hamiltonian.

**Robertson** The unique (4,5)-cage graph, i.e. a 4-regular graph of girth 5. It has 19 vertices and 38 edges.

**Smallestcyclicgroup** A smallest nontrivial graph whose automorphism group is cyclic. It has 9 vertices and 15 edges.

**Tetrahedral, Tetrahedron** Platonic solid with 4 vertices and 6 edges.

**Thomassen** The smallest hypotraceable graph, on 34 vertices and 52 edges. A hypotraceable graph does not contain a Hamiltonian path but after removing any single vertex from it the remainder always contains a Hamiltonian path. A graph containing a Hamiltonian path is called traceable.

**Tutte** Tait’s Hamiltonian graph conjecture states that every 3-connected 3-regular planar graph is Hamiltonian. This graph is a counterexample. It has 46 vertices and 69 edges.

**Uniquely3colorable** Returns a 12-vertex, triangle-free graph with chromatic number 3 that is uniquely 3-colorable.

**Walther** An identity graph with 25 vertices and 31 edges. An identity graph has a single graph automorphism, the trivial one.


### See Also

Other deterministic constructors: `graph_from_atlas()`, `graph_from_edgelist()`, `graph_from_literal()`, `make_chordal_ring()`, `make_empty_graph()`, `make_full_citation_graph()`, `make_full_graph()`, `make_lattice()`, `make_ring()`, `make_star()`, `make_tree()`

### Examples

```r
make_graph(c(1, 2, 2, 3, 3, 4, 5, 6), directed = FALSE)
make_graph(c("A", "B", "B", "C", "C", "D"), directed = FALSE)

solids <- list(make_graph("Tetrahedron"),
               make_graph("Cubical"),
               make_graph("Octahedron"),
               make_graph("Dodecahedron"),
               make_graph("Icosahedron"))

graph <- make_graph(~ A-B-C-D-A, E-A:B:C:D,
                     F-G-H-I-F, J-F:G:H:I,
                     K-L-M-N-K, O-K:L:M:N,
                     P-Q-R-S-P, T-P:Q:R:S,
                     B-F, E-J, C-I, L-T, O-T, M-S,
                     C-P, C-L, I-L, I-P)
```

---

**make_graph**

- **Nonline**
- **Octahedral, Octahedron**
- **Petersen**
- **Robertson**
- **Smallestcyclicgroup**
- **Tetrahedral, Tetrahedron**
- **Thomassen**
- **Tutte**
- **Uniquely3colorable**
- **Walther**
- **Zachary**

### See Also

- `graph_from_atlas()`, `graph_from_edgelist()`, `graph_from_literal()`, `make_chordal_ring()`, `make_empty_graph()`, `make_full_citation_graph()`, `make_full_graph()`, `make_lattice()`, `make_ring()`, `make_star()`, `make_tree()`

### Examples

```r
make_graph(c(1, 2, 2, 3, 3, 4, 5, 6), directed = FALSE)
make_graph(c("A", "B", "B", "C", "C", "D"), directed = FALSE)
```

```r
solids <- list(make_graph("Tetrahedron"),
               make_graph("Cubical"),
               make_graph("Octahedron"),
               make_graph("Dodecahedron"),
               make_graph("Icosahedron"))
```

```r
graph <- make_graph(~ A-B-C-D-A, E-A:B:C:D,
                     F-G-H-I-F, J-F:G:H:I,
                     K-L-M-N-K, O-K:L:M:N,
                     P-Q-R-S-P, T-P:Q:R:S,
                     B-F, E-J, C-I, L-T, O-T, M-S,
                     C-P, C-L, I-L, I-P)
```
**Description**

Kautz graphs are labeled graphs representing the overlap of strings.

**Usage**

```r
make_kautz_graph(m, n)
kautz_graph(...)```

**Arguments**

- `m`: Integer scalar, the size of the alphabet. See details below.
- `n`: Integer scalar, the length of the labels. See details below.
- `...`: Passed to `make_kautz_graph`.

**Details**

A Kautz graph is a labeled graph, vertices are labeled by strings of length \( n+1 \) above an alphabet with \( m+1 \) letters, with the restriction that every two consecutive letters in the string must be different. There is a directed edge from a vertex \( v \) to another vertex \( w \) if it is possible to transform the string of \( v \) into the string of \( w \) by removing the first letter and appending a letter to it.

Kautz graphs have some interesting properties, see e.g. Wikipedia for details.

**Value**

A graph object.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>, the first version in R was written by Vincent Matossian.

**See Also**

`make_de_bruijn_graph`, `make_line_graph`

**Examples**

```r
make_line_graph(make_kautz_graph(2,1))
make_kautz_graph(2,2)`
make_lattice

Create a lattice graph

Description

make_lattice is a flexible function, it can create lattices of arbitrary dimensions, periodic or aperiodic ones. It has two forms. In the first form you only supply dimvector, but not length and dim. In the second form you omit dimvector and supply length and dim.

Usage

make_lattice(
  dimvector = NULL,
  length = NULL,
  dim = NULL,
  nei = 1,
  directed = FALSE,
  mutual = FALSE,
  circular = FALSE
)

lattice(...)

Arguments

dimvector  A vector giving the size of the lattice in each dimension.
length     Integer constant, for regular lattices, the size of the lattice in each dimension.
dim        Integer constant, the dimension of the lattice.
nei         The distance within which (inclusive) the neighbors on the lattice will be connected. This parameter is not used right now.
directed   Whether to create a directed lattice.
mutual      Logical, if TRUE directed lattices will be mutually connected.
circular    Logical, if TRUE the lattice or ring will be circular.
...         Passed to make_lattice.

Value

An igraph graph.

See Also

Other deterministic constructors: graph_from_atlas(), graph_from_edgelist(), graph_from_literal(), make_chordal_ring(), make_empty_graph(), make_full_citation_graph(), make_full_graph(), make_graph(), make_ring(), make_star(), make_tree()

Examples

make_lattice(c(5, 5, 5))
make_lattice(length = 5, dim = 3)
Description

This function calculates the line graph of another graph.

Usage

```r
make_line_graph(graph)
```

```r
line_graph(...)```

Arguments

- `graph` The input graph, it can be directed or undirected.
- `...` Passed to `make_line_graph`.

Details

The line graph $L(G)$ of a $G$ undirected graph is defined as follows. $L(G)$ has one vertex for each edge in $G$ and two vertices in $L(G)$ are connected by an edge if their corresponding edges share an end point.

The line graph $L(G)$ of a $G$ directed graph is slightly different, $L(G)$ has one vertex for each edge in $G$ and two vertices in $L(G)$ are connected by a directed edge if the target of the first vertex’s corresponding edge is the same as the source of the second vertex’s corresponding edge.

Value

A new graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>, the first version of the C code was written by Vincent Matossian.

Examples

```r
# generate the first De-Bruijn graphs
g <- make_full_graph(2, directed=TRUE, loops=TRUE)
make_line_graph(g)
make_line_graph(make_line_graph(g))
make_line_graph(make_line_graph(make_line_graph(g)))```
### make_ring

Create a ring graph

#### Description

A ring is a one-dimensional lattice and this function is a special case of `make_lattice`.

#### Usage

```r
make_ring(n, directed = FALSE, mutual = FALSE, circular = TRUE)
ring(...)```

#### Arguments

- `n` Number of vertices.
- `directed` Whether the graph is directed.
- `mutual` Whether directed edges are mutual. It is ignored in undirected graphs.
- `circular` Whether to create a circular ring. A non-circular ring is essentially a “line”: a tree where every non-leaf vertex has one child.
- `...` Passed to `make_ring`.

#### Value

An igraph graph.

#### See Also

Other deterministic constructors: `graph_from_atlas()`, `graph_from_edgelist()`, `graph_from_literal()`, `make_chordal_ring()`, `make_empty_graph()`, `make_full_citation_graph()`, `make_full_graph()`, `make_graph()`, `make_lattice()`, `make_star()`, `make_tree()`

#### Examples

```r
print_all(make_ring(10))
print_all(make_ring(10, directed = TRUE, mutual = TRUE))
```

### make_star

Create a star graph, a tree with `n` vertices and `n - 1` leaves

#### Description

`star` creates a star graph, in this every single vertex is connected to the center vertex and nobody else.

#### Usage

```r
make_star(n, mode = c("in", "out", "mutual", "undirected"), center = 1)
star(...)```

#### See Also

Other deterministic constructors: `graph_from_atlas()`, `graph_from_edgelist()`, `graph_from_literal()`, `make_chordal_ring()`, `make_empty_graph()`, `make_full_citation_graph()`, `make_full_graph()`, `make_graph()`, `make_lattice()`, `make_star()`, `make_tree()`

#### Examples

```r
make_star(10)
```
make_tree

Arguments

n Number of vertices.
mode It defines the direction of the edges, in: the edges point to the center, out: the edges point from the center, mutual: a directed star is created with mutual edges, undirected: the edges are undirected.
center ID of the center vertex.
... Passed to make_star.

Value
An igraph graph.

See Also
Other deterministic constructors: graph_from_atlas(), graph_from_edgelist(), graph_from_literal(), make_chordal_ring(), make_empty_graph(), make_full_citation_graph(), make_full_graph(), make_graph(), make_lattice(), make_ring(), make_tree()

Examples

make_star(10, mode = "out")
make_star(5, mode = "undirected")

make_tree Create tree graphs

Description
Create a k-ary tree graph, where almost all vertices other than the leaves have the same number of children.

Usage

make_tree(n, children = 2, mode = c("out", "in", "undirected"))
tree(...)

Arguments

n Number of vertices.
children Integer scalar, the number of children of a vertex (except for leaves)
mode Defines the direction of the edges. out indicates that the edges point from the parent to the children, in indicates that they point from the children to their parents, while undirected creates an undirected graph.
... Passed to make_tree or sample_tree.

Value
An igraph graph
match_vertices

Description

Given two adjacency matrices A and B of the same size, match the two graphs with the help of m seed vertex pairs which correspond to the first m rows (and columns) of the adjacency matrices.

Usage

match_vertices(A, B, m, start, iteration)

Arguments

A: a numeric matrix, the adjacency matrix of the first graph
B: a numeric matrix, the adjacency matrix of the second graph
m: The number of seeds. The first m vertices of both graphs are matched.
start: a numeric matrix, the permutation matrix estimate is initialized with start
iteration: The number of iterations for the Frank-Wolfe algorithm

Details

The approximate graph matching problem is to find a bijection between the vertices of two graphs, such that the number of edge disagreements between the corresponding vertex pairs is minimized. For seeded graph matching, part of the bijection that consist of known correspondences (the seeds) is known and the problem task is to complete the bijection by estimating the permutation matrix that permutes the rows and columns of the adjacency matrix of the second graph.

It is assumed that for the two supplied adjacency matrices A and B, both of size n x n, the first m rows (and columns) of A and B correspond to the same vertices in both graphs. That is, the n x n permutation matrix that defines the bijection is Im \oplus P for a (n-m) x (n-m) permutation matrix P and m times m identity matrix Im. The function match_vertices estimates the permutation matrix P via an optimization algorithm based on the Frank-Wolfe algorithm.

See references for further details.

Value

A numeric matrix which is the permutation matrix that determines the bijection between the graphs of A and B.

See Also

Other deterministic constructors: graph_from_atlas(), graph_from_edgelist(), graph_from_literal(), make_chordal_ring(), make_empty_graph(), make_full_citation_graph(), make_full_graph(), make_graph(), make_lattice(), make_ring(), make_star()
**max_cardinality**

**Author(s)**

Vince Lyzinski [https://www.ams.jhu.edu/~lyzinski/](https://www.ams.jhu.edu/~lyzinski/)

**References**


**See Also**

`sample_correlated_gnp, sample_correlated_gnp_pair`

**Examples**

```r
#require(Matrix)
g1 <- sample_gnp(10, 0.1)
randperm <- c(1:3, 3+sample(7))
g2 <- sample_correlated_gnp(g1, corr=1, p=g1$p, permutation=randperm)
A <- as.matrix(get.adjacency(g1))
B <- as.matrix(get.adjacency(g2))
P <- match_vertices(A, B, m=3, start=diag(rep(1, nrow(A)-3)), 20)
P
```

**Description**

Maximum cardinality search is a simple ordering of vertices that is useful in determining the chordality of a graph.

**Usage**

`max_cardinality(graph)`

**Arguments**

- `graph` The input graph. It may be directed, but edge directions are ignored, as the algorithm is defined for undirected graphs.

**Details**

Maximum cardinality search visits the vertices in such an order that every time the vertex with the most already visited neighbors is visited. Ties are broken randomly.

The algorithm provides a simple basis for deciding whether a graph is chordal, see References below, and also `is_chordal`.
max_flow

Value

A list with two components:

- **alpha**: Numeric vector. The 1-based rank of each vertex in the graph such that the vertex with rank 1 is visited first, the vertex with rank 2 is visited second and so on.

- **alpham1**: Numeric vector. The inverse of alpha. In other words, the elements of this vector are the vertices in reverse maximum cardinality search order.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

- **is_chordal**

Examples

```r
## The examples from the Tarjan-Yannakakis paper

g1 <- graph_from_literal(A-B:C:I, B-A:C:D, C-A:B:E:H, D-B:E:F,
                           I-A:H)
max_cardinality(g1)
is_chordal(g1, fillin=TRUE)

g2 <- graph_from_literal(A-B:E, B-A:E:F:D, C-E:D:G, D-B:F:E:C:G,
                           I-G:H:J, J-H:J)
max_cardinality(g2)
is_chordal(g2, fillin=TRUE)
```

max_flow

Maximum flow in a graph

Description

In a graph where each edge has a given flow capacity the maximal flow between two vertices is calculated.

Usage

```r
max_flow(graph, source, target, capacity = NULL)
```
max_flow

Arguments

- graph: The input graph.
- source: The id of the source vertex.
- target: The id of the target vertex (sometimes also called sink).
- capacity: Vector giving the capacity of the edges. If this is NULL (the default) then the capacity edge attribute is used. Note that the weight edge attribute is not used by this function.

Details

max_flow calculates the maximum flow between two vertices in a weighted (ie. valued) graph. A flow from source to target is an assignment of non-negative real numbers to the edges of the graph, satisfying two properties: (1) for each edge the flow (ie. the assigned number) is not more than the capacity of the edge (the capacity parameter or edge attribute), (2) for every vertex, except the source and the target the incoming flow is the same as the outgoing flow. The value of the flow is the incoming flow of the target vertex. The maximum flow is the flow of maximum value.

Value

A named list with components:

- value: A numeric scalar, the value of the maximum flow.
- flow: A numeric vector, the flow itself, one entry for each edge. For undirected graphs this entry is bit trickier, since for these the flow direction is not predetermined by the edge direction. For these graphs the elements of the this vector can be negative, this means that the flow goes from the bigger vertex id to the smaller one. Positive values mean that the flow goes from the smaller vertex id to the bigger one.
- cut: A numeric vector of edge ids, the minimum cut corresponding to the maximum flow.
- partition1: A numeric vector of vertex ids, the vertices in the first partition of the minimum cut corresponding to the maximum flow.
- partition2: A numeric vector of vertex ids, the vertices in the second partition of the minimum cut corresponding to the maximum flow.
- stats: A list with some statistics from the push-relabel algorithm. Five integer values currently: nopush is the number of push operations, norelabel the number of relabelings, nogap is the number of times the gap heuristics was used, nogapnodes is the total number of gap nodes omitted because of the gap heuristics and nobfs is the number of times a global breadth-first-search update was performed to assign better height (=distance) values to the vertices.

References


See Also

min_cut for minimum cut calculations, distances, edge_connectivity, vertex_connectivity
Examples

```r
E <- rbind( c(1,3,3), c(3,4,1), c(4,2,2), c(1,5,1), c(5,6,2), c(6,2,10))
colnames(E) <- c("from", "to", "capacity")
g1 <- graph_from_data_frame(as.data.frame(E))
max_flow(g1, source=V(g1)["1"], target=V(g1)["2"])
```

Description

igraph community detection functions return their results as an object from the `communities` class. This manual page describes the operations of this class.

Usage

```r
membership(communities)

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

## S3 method for class 'communities'
modularity(x, ...)

## S3 method for class 'communities'
length(x)

sizes(communities)

algorithm(communities)

merges(communities)

crossing(communities, graph)

code_len(communities)

is_hierarchical(communities)

## S3 method for class 'communities'
as.dendrogram(object, hang = -1, use.modularity = FALSE, ...)

## S3 method for class 'communities'
as.hclust(x, hang = -1, use.modularity = FALSE, ...)

as_phylo(x, ...)

cut_at(communities, no, steps)
```
show_trace(communities)

## S3 method for class 'communities'
plot(
    x,
    y,
    col = membership(x),
    mark.groups = communities(x),
    edge.color = c("black", "red") [crossing(x, y) + 1],
    ...
)

Arguments

communities, x, object

A communities object, the result of an igraph community detection function.

... Additional arguments. plot.communities passes these to plot.igraph. The
other functions silently ignore them.

graph An igraph graph object, corresponding to communities.

hang Numeric scalar indicating how the height of leaves should be computed from
the heights of their parents; see plot.hclust.

use.modularity Logical scalar, whether to use the modularity values to define the height of the
branches.

no Integer scalar, the desired number of communities. If too low or two high, then
an error message is given. Exactly one of no and steps must be supplied.

steps The number of merge operations to perform to produce the communities. Ex-
actly one of no and steps must be supplied.

y An igraph graph object, corresponding to the communities in x.

col A vector of colors, in any format that is accepted by the regular R plotting meth-
ods. This vector gives the colors of the vertices explicitly.

mark.groups A list of numeric vectors. The communities can be highlighted using colored
polygons. The groups for which the polygons are drawn are given here. The
default is to use the groups given by the communities. Supply NULL here if you
do not want to highlight any groups.

dfill The colors of the edges. By default the edges within communities are colored
green and other edges are red.

membership Numeric vector, one value for each vertex, the membership vector of the com-
munity structure. Might also be NULL if the community structure is given in
another way, e.g. by a merge matrix.

algorithm If not NULL (meaning an unknown algorithm), then a character scalar, the name
of the algorithm that produced the community structure.

merges If not NULL, then the merge matrix of the hierarchical community structure. See
merges below for more information on its format.

modularity Numeric scalar or vector, the modularity value of the community structure. It
can also be NULL, if the modularity of the (best) split is not available.
Community structure detection algorithms try to find dense subgraphs in directed or undirected graphs, by optimizing some criteria, and usually using heuristics.

igraph implements a number of community detection methods (see them below), all of which return an object of the class communities. Because the community structure detection algorithms are different, communities objects do not always have the same structure. Nevertheless, they have some common operations, these are documented here.

The print generic function is defined for communities, it prints a short summary.

The length generic function call be called on communities and returns the number of communities.

The sizes function returns the community sizes, in the order of their ids.

membership gives the division of the vertices, into communities. It returns a numeric vector, one value for each vertex, the id of its community. Community ids start from one. Note that some algorithms calculate the complete (or incomplete) hierarchical structure of the communities, and not just a single partitioning. For these algorithms typically the membership for the highest modularity value is returned, but see also the manual pages of the individual algorithms.

communities is also the name of a function, that returns a list of communities, each identified by their vertices. The vertices will have symbolic names if the add.vertex.names igraph option is set, and the graph itself was named. Otherwise numeric vertex ids are used.

modularity gives the modularity score of the partitioning. (See modularity.igraph for details. For algorithms that do not result a single partitioning, the highest modularity value is returned.

algorithm gives the name of the algorithm that was used to calculate the community structure.

crossing returns a logical vector, with one value for each edge, ordered according to the edge ids. The value is TRUE iff the edge connects two different communities, according to the (best) membership vector, as returned by membership().

is_hierarchical checks whether a hierarchical algorithm was used to find the community structure. Some functions only make sense for hierarchical methods (e.g. merges, cut_at and as.dendrogram).

merges returns the merge matrix for hierarchical methods. An error message is given, if a non-hierarchical method was used to find the community structure. You can check this by calling is_hierarchical on the communities object.

cut_at cuts the merge tree of a hierarchical community finding method, at the desired place and returns a membership vector. The desired place can be expressed as the desired number of communities or as the number of merge steps to make. The function gives an error message, if called with a non-hierarchical method.

as.dendrogram converts a hierarchical community structure to a dendrogram object. It only works for hierarchical methods, and gives an error message to others. See dendrogram for details.

as.hclust is similar to as.dendrogram, but converts a hierarchical community structure to a hclust object.

as.phylo converts a hierarchical community structure to a phylo object, you will need the ape package for this.

show_trace works (currently) only for communities found by the leading eigenvector method (cluster_leading_eigen), and returns a character vector that gives the steps performed by the algorithm while finding the communities.

code_len is defined for the InfoMAP method (cluster_infomap and returns the code length of the partition.

It is possibly to call the plot function on communities objects. This will plot the graph (and uses plot.igraph internally), with the communities shown. By default it colores the vertices according
to their communities, and also marks the vertex groups corresponding to the communities. It passes additional arguments to \texttt{plot.igraph}, please see that and also \texttt{igraph.plotting} on how to change the plot.

**Value**

\texttt{print} returns the communities object itself, invisibly.

\texttt{length} returns an integer scalar.

\texttt{sizes} returns a numeric vector.

\texttt{membership} returns a numeric vector, one number for each vertex in the graph that was the input of the community detection.

\texttt{modularity} returns a numeric scalar.

\texttt{algorithm} returns a character scalar.

\texttt{crossing} returns a logical vector.

\texttt{is_hierarchical} returns a logical scalar.

\texttt{merges} returns a two-column numeric matrix.

\texttt{cut_at} returns a numeric vector, the membership vector of the vertices.

\texttt{as.dendrogram} returns a \texttt{dendrogram} object.

\texttt{show_trace} returns a character vector.

\texttt{code_len} returns a numeric scalar for communities found with the InfoMAP method and \texttt{NULL} for other methods.

\texttt{plot} for communities objects returns \texttt{NULL}, invisibly.

# @author Gabor Csardi <csardi.gabor@gmail.com>

**See Also**

See \texttt{plot.dendrogram} for plotting community structure dendrograms.

See \texttt{compare} for comparing two community structures on the same graph.

The different methods for finding communities, they all return a communities object: \texttt{cluster_edge_betweenness, cluster_fast_greedy, cluster_label_prop, cluster_leading_eigen, cluster_louvain, cluster_leiden, cluster_optimal, cluster_spinglass, cluster_walktrap}.

**Examples**

karate <- make_graph("Zachary")
w <- cluster_walktrap(karate)
modularity(w)
membership(w)
plot(w, karate)
**merge_coords**  Merging graph layouts

**Description**
Place several graphs on the same layout

**Usage**
```
merge_coords(graphs, layouts, method = "dla")
layout_components(graph, layout = layout_with_kk, ...)
```

**Arguments**
- **graphs**: A list of graph objects.
- **layouts**: A list of two-column matrices.
- **method**: Character constant giving the method to use. Right now only dla is implemented.
- **graph**: The input graph.
- **layout**: A function object, the layout function to use.
- **...**: Additional arguments to pass to the layout layout function.

**Details**
merge_coords takes a list of graphs and a list of coordinates and places the graphs in a common layout. The method to use is chosen via the method parameter, although right now only the dla method is implemented.

The dla method covers the graph with circles. Then it sorts the graphs based on the number of vertices first and places the largest graph at the center of the layout. Then the other graphs are placed in decreasing order via a DLA (diffusion limited aggregation) algorithm: the graph is placed randomly on a circle far away from the center and a random walk is conducted until the graph walks into the larger graphs already placed or walks too far from the center of the layout.

The layout_components function disassembles the graph first into maximal connected components and calls the supplied layout function for each component separately. Finally it merges the layouts via calling merge_coords.

**Value**
A matrix with two columns and as many lines as the total number of vertices in the graphs.

**Author(s)**
Gabor Csardi <csardi.gabor@gmail.com>
min_cut

See Also

plot.igraph, tkplot, layout.disjoint_union

Other graph layouts: add_layout(), component_wise(), layout_as_bipartite(), layout_as_star(), layout_as_tree(), layout_in_circle(), layout_nicely(), layout_on_grid(), layout_on_sphere(), layout_randomly(), layout_with_dh(), layout_with_fr(), layout_with_gem(), layout_with_graphopt(), layout_with_kk(), layout_with_lgl(), layout_with_mds(), layout_with_sugiyama(), layout(), norm_coords(), normalize

Examples

# create 20 scale-free graphs and place them in a common layout
graphs <- lapply(sample(5:20, 20, replace=TRUE), barabasi.game, directed=FALSE)
layouts <- lapply(graphs, layout_with_kk)
lay <- merge_coords(graphs, layouts)
g <- disjoint_union(graphs)
## Not run: plot(g, layout=lay, vertex.size=3, labels=NA, edge.color="black")

min_cut

Minimum cut in a graph

Description

min_cut calculates the minimum st-cut between two vertices in a graph (if the source and target arguments are given) or the minimum cut of the graph (if both source and target are NULL).

Usage

min_cut(
  graph,
  source = NULL,
  target = NULL,
  capacity = NULL,
  value.only = TRUE
)

Arguments

graph The input graph.
source The id of the source vertex.
target The id of the target vertex (sometimes also called sink).
capacity Vector giving the capacity of the edges. If this is NULL (the default) then the capacity edge attribute is used.
value.only Logical scalar, if TRUE only the minimum cut value is returned, if FALSE the edges in the cut and a the two (or more) partitions are also returned.
Details

The minimum st-cut between source and target is the minimum total weight of edges needed to remove to eliminate all paths from source to target.

The minimum cut of a graph is the minimum total weight of the edges needed to remove to separate the graph into (at least) two components. (Which is to make the graph not strongly connected in the directed case.)

The maximum flow between two vertices in a graph is the same as the minimum st-cut, so max_flow and min_cut essentially calculate the same quantity, the only difference is that min_cut can be invoked without giving the source and target arguments and then minimum of all possible minimum cuts is calculated.

For undirected graphs the Stoer-Wagner algorithm (see reference below) is used to calculate the minimum cut.

Value

For min_cut a numeric constant, the value of the minimum cut, except if value.only = FALSE. In this case a named list with components:

- value: Numeric scalar, the cut value.
- cut: Numeric vector, the edges in the cut.
- partition1: The vertices in the first partition after the cut edges are removed. Note that these vertices might be actually in different components (after the cut edges are removed), as the graph may fall apart into more than two components.
- partition2: The vertices in the second partition after the cut edges are removed. Note that these vertices might be actually in different components (after the cut edges are removed), as the graph may fall apart into more than two components.

References


See Also

max_flow for the related maximum flow problem, distances, edge_connectivity, vertex_connectivity

Examples

g <- make_ring(100)
min_cut(g, capacity=rep(1,vcount(g)))
min_cut(g, value.only=FALSE, capacity=rep(1,vcount(g)))

g2 <- graph( c(1,2,3,4, 1,6,6,5,5,4, 4,1) )
E(g2)$capacity <- c(3,1,2, 10,1,3, 2)
min_cut(g2, value.only=FALSE)
**min_separators**

Minimum size vertex separators

**Description**

Find all vertex sets of minimal size whose removal separates the graph into more components.

**Usage**

`min_separators(graph)`

**Arguments**

- `graph`: The input graph. It may be directed, but edge directions are ignored.

**Details**

This function implements the Kanevsky algorithm for finding all minimal-size vertex separators in an undirected graph. See the reference below for the details.

In the special case of a fully connected input graph with \( n \) vertices, all subsets of size \( n - 1 \) are listed as the result.

**Value**

A list of numeric vectors. Each numeric vector is a vertex separator.

**References**


**See Also**

`is.separator`

**Examples**

# The graph from the Moody-White paper
```r
```

# Cohesive subgraphs
```r
mw1 <- induced.subgraph(mw, as.character(c(1:7, 17:23)))
mw2 <- induced.subgraph(mw, as.character(7:16))
mw3 <- induced.subgraph(mw, as.character(17:23))
```
min_st_separators

Description

List all vertex sets that are minimal (s,t) separators for some s and t, in an undirected graph.

Usage

min_st_separators(graph)

Arguments

graph

The input graph. It may be directed, but edge directions are ignored.

Details

A (s, t) vertex separator is a set of vertices, such that after their removal from the graph, there is no path between s and t in the graph.

A (s, t) vertex separator is minimal if none of its subsets is an (s, t) vertex separator.

Value

A list of numeric vectors. Each vector contains a vertex set (defined by vertex ids), each vector is an (s,t) separator of the input graph, for some s and t.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>
modularity.igraph

References

Anne Berry, Jean-Paul Bordat and Olivier Cogis: Generating All the Minimal Separators of a Graph, In: Peter Widmayer, Gabriele Neyer and Stephan Eidenbenz (editors): Graph-theoretic concepts in computer science, 1665, 167–172, 1999. Springer.

Examples

```r
ring <- make_ring(4)
min_st_separators(ring)

chvatal <- make_graph("chvatal")
min_st_separators(chvatal)
```

modularity.igraph  
Modularity of a community structure of a graph

Description

This function calculates how modular is a given division of a graph into subgraphs.

Usage

```r
## S3 method for class 'igraph'
modularity(x, membership, weights = NULL, resolution = 1, directed = TRUE, ...)

modularity_matrix(
  graph,
  membership,
  weights = NULL,
  resolution = 1,
  directed = TRUE
)
```

Arguments

- `x, graph`  
The input graph.
- `membership`  
Numeric vector, one value for each vertex, the membership vector of the community structure.
- `weights`  
If not NULL then a numeric vector giving edge weights.
- `resolution`  
The resolution parameter. Must be greater than or equal to 0. Set it to 1 to use the classical definition of modularity.
- `directed`  
Whether to use the directed or undirected version of modularity. Ignored for undirected graphs.
- `...`  
Additional arguments, none currently.
modularity calculates the modularity of a graph with respect to the given membership vector.

The modularity of a graph with respect to some division (or vertex types) measures how good the division is, or how separated are the different vertex types from each other. It is defined as

$$Q = \frac{1}{2m} \sum_{i,j} (A_{ij} - \gamma \frac{k_i k_j}{2m}) \delta(c_i, c_j),$$

here $m$ is the number of edges, $A_{ij}$ is the element of the $A$ adjacency matrix in row $i$ and column $j$, $k_i$ is the degree of $i$, $k_j$ is the degree of $j$, $c_i$ is the type (or component) of $i$, $c_j$ that of $j$, the sum goes over all $i$ and $j$ pairs of vertices, and $\delta(x, y)$ is 1 if $x = y$ and 0 otherwise. For directed graphs, it is defined as

$$Q = \frac{1}{m} \sum_{i,j} (A_{ij} - \gamma \frac{k^\text{out}_i k^\text{in}_j}{m}) \delta(c_i, c_j).$$

The resolution parameter $\gamma$ allows weighting the random null model, which might be useful when finding partitions with a high modularity. Maximizing modularity with higher values of the resolution parameter typically results in more, smaller clusters when finding partitions with a high modularity. Lower values typically results in fewer, larger clusters. The original definition of modularity is retrieved when setting $\gamma$ to 1.

If edge weights are given, then these are considered as the element of the $A$ adjacency matrix, and $k_i$ is the sum of weights of adjacent edges for vertex $i$.

modularity_matrix calculates the modularity matrix. This is a dense matrix, and it is defined as the difference of the adjacency matrix and the configuration model null model matrix. In other words, element $M_{ij}$ is given as $A_{ij} - \frac{d_i d_j}{(2m)}$, where $A_{ij}$ is the (possibly weighted) adjacency matrix, $d_i$ is the degree of vertex $i$, and $m$ is the number of edges (or the total weights in the graph, if it is weighed).

Value

For modularity a numeric scalar, the modularity score of the given configuration.

For modularity_matrix a numeric square matrix, its order is the number of vertices in the graph.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

cluster_walktrap, cluster_edge_betweenness, cluster_fast_greedy, cluster_spinglass, cluster_louvain and cluster_leiden for various community detection methods.
motifs

Examples

```r
  g <- make_full_graph(5) %du% make_full_graph(5) %du% make_full_graph(5)
  g <- add_edges(g, c(1,6, 1,11, 6, 11))
  wtc <- cluster_walktrap(g)
  modularity(wtc)
  modularity(g, membership(wtc))
```

motifs

Graph motifs

Description

Graph motifs are small connected subgraphs with a well-defined structure. These functions search a graph for various motifs.

Usage

```r
  motifs(graph, size = 3, cut.prob = rep(0, size))
```

Arguments

- `graph`: Graph object, the input graph.
- `size`: The size of the motif, currently sizes 3 and 4 are supported in directed graphs and sizes 3-6 in undirected graphs.
- `cut.prob`: Numeric vector giving the probabilities that the search graph is cut at a certain level. Its length should be the same as the size of the motif (the size argument). By default no cuts are made.

Details

`motifs` searches a graph for motifs of a given size and returns a numeric vector containing the number of different motifs. The order of the motifs is defined by their isomorphism class, see `isomorphism_class`.

Value

`motifs` returns a numeric vector, the number of occurrences of each motif in the graph. The motifs are ordered by their isomorphism classes. Note that for unconnected subgraphs, which are not considered to be motifs, the result will be NA.

See Also

`isomorphism_class`

Other graph motifs: `count_motifs()`, `sample_motifs()`

Examples

```r
  g <- barabasi.game(100)
  motifs(g, 3)
  count_motifs(g, 3)
  sample_motifs(g, 3)
```
Description

A subgraph of a connected graph is a minimum spanning tree if it is tree, and the sum of its edge weights are the minimal among all tree subgraphs of the graph. A minimum spanning forest of a graph is the graph consisting of the minimum spanning trees of its components.

Usage

\texttt{mst(graph, weights = NULL, algorithm = NULL, ...)}

Arguments

- \texttt{graph}: The graph object to analyze.
- \texttt{weights}: Numeric algorithm giving the weights of the edges in the graph. The order is determined by the edge ids. This is ignored if the unweighted algorithm is chosen. Edge weights are interpreted as distances.
- \texttt{algorithm}: The algorithm to use for calculation. unweighted can be used for unweighted graphs, and \texttt{prim} runs Prim's algorithm for weighted graphs. If this is \texttt{NULL} then igraph tries to select the algorithm automatically: if the graph has an edge attribute called \texttt{weight} or the \texttt{weights} argument is not \texttt{NULL} then Prim's algorithm is chosen, otherwise the unweighted algorithm is performed.
- \texttt{...}: Additional arguments, unused.

Details

If the graph is unconnected a minimum spanning forest is returned.

Value

A graph object with the minimum spanning forest. (To check that it is a tree check that the number of its edges is \texttt{vcount(graph)}-1.) The edge and vertex attributes of the original graph are preserved in the result.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

\texttt{components}
### Description

A vertex is a neighbor of another one (in other words, the two vertices are adjacent), if they are incident to the same edge.

### Usage

```r
eighbors(graph, v, mode = c("out", "in", "all", "total"))
```

### Arguments

- **graph**: The input graph.
- **v**: The vertex of which the adjacent vertices are queried.
- **mode**: Whether to query outgoing ('out'), incoming ('in') edges, or both types ('all'). This is ignored for undirected graphs.

### Value

A vertex sequence containing the neighbors of the input vertex.

### See Also

Other structural queries: `igraph()`, `igraph()`, `adjacent_vertices()`, `are_adjacent()`, `ends()`, `get.edge.ids()`, `gorder()`, `gsizes()`, `head_of()`, `incident_edges()`, `incident()`, `is_directed()`, `tail_of()`

### Examples

```r
g <- sample_gnp(100, 3/100)
g_mst <- mst(g)

# Zachary graph example

# Find neighbors
n1 <- neighbors(g, 1)
n34 <- neighbors(g, 34)
intersection(n1, n34)
```
norm_coords

Normalize coordinates for plotting graphs

Description

Rescale coordinates linearly to be within given bounds.

Usage

```r
norm_coords(
  layout,
  xmin = -1,
  xmax = 1,
  ymin = -1,
  ymax = 1,
  zmin = -1,
  zmax = 1
)
```

Arguments

- `layout`: A matrix with two or three columns, the layout to normalize.
- `xmin, xmax`: The limits for the first coordinate, if one of them or both are NULL then no normalization is performed along this direction.
- `ymin, ymax`: The limits for the second coordinate, if one of them or both are NULL then no normalization is performed along this direction.
- `zmin, zmax`: The limits for the third coordinate, if one of them or both are NULL then no normalization is performed along this direction.

Details

`norm_coords` normalizes a layout, it linearly transforms each coordinate separately to fit into the given limits.

Value

A numeric matrix with at the same dimension as `layout`.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

Other graph layouts: `add_layout_()`, `component_wise()`, `layout_as_bipartite()`, `layout_as_star()`, `layout_as_tree()`, `layout_in_circle()`, `layout_nicely()`, `layout_on_grid()`, `layout_on_sphere()`, `layout_randomly()`, `layout_with_dh()`, `layout_with_fr()`, `layout_with_gem()`, `layout_with_graphopt()`, `layout_with_kk()`, `layout_with_lgl()`, `layout_with_mds()`, `layout_with_sugiyama()`, `layout_()`, `merge_coords()`, `normalize()`
normalize

Normalize layout

Description

Scale coordinates of a layout.

Usage

normalize(
  xmin = -1,
  xmax = 1,
  ymin = xmin,
  ymax = xmax,
  zmin = xmin,
  zmax = xmax
)

Arguments

xmin, xmax Minimum and maximum for x coordinates.

ymin, ymax Minimum and maximum for y coordinates.

zmin, zmax Minimum and maximum for z coordinates.

See Also

merge_coords, layout_

Other layout modifiers: component_wise()

Other graph layouts: add_layout_, component_wise, layout_as_bipartite, layout_as_star, layout_as_tree, layout_in_circle, layout_nicely, layout_on_grid, layout_on_sphere, layout_randomly, layout_with_dh, layout_with_fr, layout_with_gem, layout_with_graphopt, layout_with_kw, layout_with_lgl, layout_with_mds, layout_with_sugiyama, layout, merge_coords, norm_coords

Examples

layout_(make_ring(10), with_fr(), normalize())

---

tabular

page_rank  The Page Rank algorithm

Description

Calculates the Google PageRank for the specified vertices.
Usage

page_rank(
  graph,
  algo = c("prpack", "arpack"),
  vids = V(graph),
  directed = TRUE,
  damping = 0.85,
  personalized = NULL,
  weights = NULL,
  options = NULL
)

Arguments

graph The graph object.
algo Character scalar, which implementation to use to carry out the calculation. The default is "prpack", which uses the PRPACK library (https://github.com/dgleich/prpack). This is a new implementation in igraph version 0.7, and the suggested one, as it is the most stable and the fastest for all but small graphs. "arpack" uses the ARPACK library, the default implementation from igraph version 0.5 until version 0.7.

vids The vertices of interest.
directed Logical, if true directed paths will be considered for directed graphs. It is ignored for undirected graphs.
damping The damping factor (‘d’ in the original paper).
personalized Optional vector giving a probability distribution to calculate personalized PageRank. For personalized PageRank, the probability of jumping to a node when abandoning the random walk is not uniform, but it is given by this vector. The vector should contains an entry for each vertex and it will be rescaled to sum up to one.
weights A numerical vector or NULL. This argument can be used to give edge weights for calculating the weighted PageRank of vertices. If this is NULL and the graph has a weight edge attribute then that is used. If weights is a numerical vector then it used, even if the graph has a weights edge attribute. If this is NA, then no edge weights are used (even if the graph has a weight edge attribute. This function interprets edge weights as connection strengths. In the random surfer model, an edge with a larger weight is more likely to be selected by the surfer.

options A named list, to override some ARPACK options. See arpack for details. This argument is ignored if the PRPACK implementation is used.

Details

For the explanation of the PageRank algorithm, see the following webpage: http://infolab.stanford.edu/~backrub/google.html, or the following reference:


The page_rank function can use either the PRPACK library or ARPACK (see arpack) to perform the calculation.
Please note that the PageRank of a given vertex depends on the PageRank of all other vertices, so even if you want to calculate the PageRank for only some of the vertices, all of them must be calculated. Requesting the PageRank for only some of the vertices does not result in any performance increase at all.

**Value**

A named list with entries:

- **vector**: A numeric vector with the PageRank scores.
- **value**: The eigenvalue corresponding to the eigenvector with the page rank scores. It should be always exactly one.
- **options**: Some information about the underlying ARPACK calculation. See `arpack` for details. This entry is NULL if not the ARPACK implementation was used.

**Author(s)**

Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com>

**References**


**See Also**

Other centrality scores: `closeness`, `betweenness`, `degree`

**Examples**

```r
g <- sample_gnp(20, 5/20, directed=TRUE)
pagerank(g)$vector

g2 <- make_star(10)
pagerank(g2)$vector

# Personalized PageRank
g3 <- make_ring(10)
pagerank(g3)$vector
reset <- seq(vcount(g3))
pagerank(g3, personalized=reset)$vector
```

---

**path**

*Helper function to add or delete edges along a path*

**Description**

This function can be used to add or delete edges that form a path.

**Usage**

`path(...)`
Arguments

... See details below.

Details

When adding edges via +, all unnamed arguments are concatenated, and each element of a final vector is interpreted as a vertex in the graph. For a vector of length \( n + 1 \), \( n \) edges are then added, from vertex 1 to vertex 2, from vertex 2 to vertex 3, etc. Named arguments will be used as edge attributes for the new edges.

When deleting edges, all attributes are concatenated and then passed to delete_edges.

Value

A special object that can be used together with igraph graphs and the plus and minus operators.

See Also

Other functions for manipulating graph structure: \+.igraph(), add_edges(), add_vertices(), delete_edges(), delete_vertices(), edge(), igraph-minus, vertex()

Examples

# Create a (directed) wheel
\n\n```r
\n\n\ng <- make_star(11, center = 1) + path(2:11, 2)
plot(g)

\ng <- make_empty_graph(directed = FALSE, n = 10) %>%
  set_vertex_attr("name", value = letters[1:10])
\ng2 <- g + path("a", "b", "c", "d")
plot(g2)
\ng3 <- g2 + path("e", "f", "g", weight=1:2, color=\"red\")
E(g3)[[]]
\ng4 <- g3 + path(c("f", "c", "j", "d"), width=1:3, color=\"green\")
E(g4)[[]]
```

---

permute  

Permute the vertices of a graph

Description

Create a new graph, by permuting vertex ids.

Usage

`permute(graph, permutation)`
Arguments

- **graph**: The input graph, it can directed or undirected.
- **permutation**: A numeric vector giving the permutation to apply. The first element is the new id of vertex 1, etc. Every number between one and `vcount(graph)` must appear exactly once.

Details

This function creates a new graph from the input graph by permuting its vertices according to the specified mapping. Call this function with the output of `canonical_permutation` to create the canonical form of a graph.

`permute` keeps all graph, vertex and edge attributes of the graph.

Value

A new graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

`canonical_permutation`

Examples

```r
# Random permutation of a random graph
g <- sample_gnm(20, 50)
g2 <- permute(g, sample(vcount(g)))
graph.isomorphic(g, g2)

# Permutation keeps all attributes
g$name <- "Random graph, Gnm, 20, 50"
V(g)$name <- letters[1:vcount(g)]
E(g)$weight <- sample(1:5, ecount(g), replace=TRUE)
g2 <- permute(g, sample(vcount(g)))
graph.isomorphic(g, g2)
g2$name
V(g2)$name
E(g2)$weight
all(sort(E(g2)$weight) == sort(E(g)$weight))
```

---

**Pie charts as vertices**

*Using pie charts as vertices in graph plots*

Description

More complex vertex images can be used to express additional information about vertices. E.g. pie charts can be used as vertices, to denote vertex classes, fuzzy classification of vertices, etc.
Details

The vertex shape ‘pie’ makes igraph draw a pie chart for every vertex. There are some extra graphical vertex parameters that specify how the pie charts will look like:

- **pie** Numeric vector, gives the sizes of the pie slices.
- **pie.color** A list of color vectors to use for the pies. If it is a list of a single vector, then this is used for all pies. If the color vector is shorter than the number of areas in a pie, then it is recycled.
- **pie.border** The color of the border line of the pie charts, in the same format as `pie.color`.
- **pie.angle** The slope of shading lines, given as an angle in degrees (counter-clockwise).
- **pie.density** The density of the shading lines, in lines per inch. Non-positive values inhibit the drawing of shading lines.
- **pie.lty** The line type of the border of the slices.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

`igraph.plotting`, `plot.igraph`

Examples

```r
g <- make_ring(10)
values <- lapply(1:10, function(x) sample(1:10,3))
if (interactive()) {
  plot(g, vertex.shape="pie", vertex.pie=values,
       vertex.pie.color=list(heat.colors(5)),
       vertex.size=seq(10,30,length.out=10), vertex.label=NA)
}
```

plot.igraph

Plotting of graphs

Description

`plot.igraph` is able to plot graphs to any R device. It is the non-interactive companion of the tkplot function.

Usage

```r
## S3 method for class 'igraph'
plot(
  x,
  axes = FALSE,
  add = FALSE,
  xlim = c(-1, 1),
  ylim = c(-1, 1),
  mark.groups = list(),
  mark.shape = 1/2,
```
mark.col = rainbow(length(mark.groups), alpha = 0.3),
mark.border = rainbow(length(mark.groups), alpha = 1),
mark.expand = 15,
...)

Arguments

x The graph to plot.

axes Logical, whether to plot axes, defaults to FALSE.

add Logical scalar, whether to add the plot to the current device, or delete the device’s current contents first.

xlim The limits for the horizontal axis, it is unlikely that you want to modify this.

ylim The limits for the vertical axis, it is unlikely that you want to modify this.

mark.groups A list of vertex id vectors. It is interpreted as a set of vertex groups. Each vertex group is highlighted, by plotting a colored smoothed polygon around and “under” it. See the arguments below to control the look of the polygons.

mark.shape A numeric scalar or vector. Controls the smoothness of the vertex group marking polygons. This is basically the ‘shape’ parameter of the xspline function, its possible values are between -1 and 1. If it is a vector, then a different value is used for the different vertex groups.

mark.col A scalar or vector giving the colors of marking the polygons, in any format accepted by xspline; e.g. numeric color ids, symbolic color names, or colors in RGB.

mark.border A scalar or vector giving the colors of the borders of the vertex group marking polygons. If it is NA, then no border is drawn.

mark.expand A numeric scalar or vector, the size of the border around the marked vertex groups. It is in the same units as the vertex sizes. If a vector is given, then different values are used for the different vertex groups.

... Additional plotting parameters. See igraph.plotting for the complete list.

Details

One convenient way to plot graphs is to plot with tkplot first, handtune the placement of the vertices, query the coordinates by the tk_coords function and use them with plot to plot the graph to any R device.

Value

Returns NULL, invisibly.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

layout for different layouts, igraph.plotting for the detailed description of the plotting parameters and tkplot and rglplot for other graph plotting functions.
Examples

```r
g <- make_ring(10)
plot(g, layout=layout_with_kk, vertex.color="green")
```

Description

This function can conveniently plot the results of multiple SIR model simulations.

Usage

```r
## S3 method for class 'sir'
plot(
x, 
comp = c("NI", "NS", "NR"),
median = TRUE,
quantiles = c(0.1, 0.9),
color = NULL,
median_color = NULL,
quantile_color = NULL,
lwd.median = 2,
lwd.quantile = 2,
lty.quantile = 3,
xlim = NULL,
ylim = NULL,
xlab = "Time",
ylab = NULL,
...)
```

Arguments

- **x**: The output of the SIR simulation, coming from the `sir` function.
- **comp**: Character scalar, which component to plot. Either `‘NI’` (infected, default), `‘NS’` (susceptible) or `‘NR’` (recovered).
- **median**: Logical scalar, whether to plot the (binned) median.
- **quantiles**: A vector of (binned) quantiles to plot.
- **color**: Color of the individual simulation curves.
- **median_color**: Color of the median curve.
- **quantile_color**: Color(s) of the quantile curves. (It is recycled if needed and non-needed entries are ignored if too long.)
- **lwd.median**: Line width of the median.
- **lwd.quantile**: Line width of the quantile curves.
- **lty.quantile**: Line type of the quantile curves.
plot_dendrogram

xlim The x limits, a two-element numeric vector. If NULL, then it is calculated from the data.

ylim The y limits, a two-element numeric vector. If NULL, then it is calculated from the data.

xlab The x label.

ylab The y label. If NULL then it is automatically added based on the comp argument.

... Additional arguments are passed to plot, that is run before any of the curves are added, to create the figure.

Details

The number of susceptible/infected/recovered individuals is plotted over time, for multiple simulations.

Value

Nothing.

Author(s)

Eric Kolaczyk (http://math.bu.edu/people/kolaczyk/) and Gabor Csardi <csardi.gabor@gmail.com>.

References


See Also

sir for running the actual simulation.

Examples

g <- sample_gnm(100, 100)
sm <- sir(g, beta=5, gamma=1)
plot(sm)

plot_dendrogram

Community structure dendrogram plots

Description

Plot a hierarchical community structure as a dendrogram.
plot_dendrogram

Usage

plot_dendrogram(x, mode = igraph_opt("dend.plot.type"), ...)

## S3 method for class 'communities'
plot_dendrogram(
  x,
  mode = igraph_opt("dend.plot.type"),
  ...,  
  use.modularity = FALSE,
  palette = categorical_pal(8)
)

Arguments

x An object containing the community structure of a graph. See communities for details.

mode Which dendrogram plotting function to use. See details below.

... Additional arguments to supply to the dendrogram plotting function.

use.modularity Logical scalar, whether to use the modularity values to define the height of the branches.

palette The color palette to use for colored plots.

Details

plot_dendrogram supports three different plotting functions, selected via the mode argument. By default the plotting function is taken from the dend.plot.type igraph option, and it has for possible values:

- auto Choose automatically between the plotting functions. As plot.phylo is the most sophisticated, that is chosen, whenever the ape package is available. Otherwise plot.hclust is used.
- phylo Use plot.phylo from the ape package.
- hclust Use plot.hclust from the stats package.
- dendrogram Use plot.dendrogram from the stats package.

The different plotting functions take different sets of arguments. When using plot.phylo (mode="phylo"), we have the following syntax:

plot_dendrogram(x, mode="phylo", colbar = palette(),
  edge.color = NULL, use.edge.length = FALSE, \dots)

The extra arguments not documented above:

- colbar Color bar for the edges.
- edge.color Edge colors. If NULL, then the colbar argument is used.
- use.edge.length Passed to plot.phylo.
- dots Additional arguments to pass to plot.phylo.

The syntax for plot.hclust (mode="hclust"):  

plot_dendrogram(x, mode="hclust", rect = 0, colbar = palette(),
    hang = 0.01, ann = FALSE, main = "", sub = "", xlab = "",
    ylab = "", \dots)

The extra arguments not documented above:

• rect  A numeric scalar, the number of groups to mark on the
dendrogram. The dendrogram is
cut into exactly rect groups and they are marked via the rect.hclust command. Set this to
zero if you don’t want to mark any groups.
• colbar  The colors of the rectangles that mark the vertex groups via the rect argument.
• hang  Where to put the leaf nodes, this corresponds to the hang argument of plot.hclust.
• ann  Whether to annotate the plot, the ann argument of plot.hclust.
• main  The main title of the plot, the main argument of plot.hclust.
• sub  The sub-title of the plot, the sub argument of plot.hclust.
• xlab  The label on the horizontal axis, passed to plot.hclust.
• ylab  The label on the vertical axis, passed to plot.hclust.
• dots  Additional arguments to pass to plot.hclust.

The syntax for plot.dendrogram (mode="dendrogram"):

plot_dendrogram(x, \dots)

The extra arguments are simply passed to as.dendrogram.

Value

Returns whatever the return value was from the plotting function, plot.phylo, plot.dendrogram
or plot.hclust.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

karate <- make_graph("Zachary")
fc <- cluster_fast_greedy(karate)
plot_dendrogram(fc)
Usage

```r
## S3 method for class 'igraphHRG'
plot_dendrogram(x, mode = igraph_opt("dend.plot.type"), ...)
```

Arguments

- **x**: An igraphHRG, a hierarchical random graph, as returned by the `fit_hrg` function.
- **mode**: Which dendrogram plotting function to use. See details below.
- **...**: Additional arguments to supply to the dendrogram plotting function.

Details

`plot_dendrogram` supports three different plotting functions, selected via the `mode` argument. By default the plotting function is taken from the `dend.plot.type` igraph option, and it has for possible values:

- **auto**: Choose automatically between the plotting functions. As `plot.phylo` is the most sophisticated, that is chosen, whenever the ape package is available. Otherwise `plot.hclust` is used.
- **phylo**: Use `plot.phylo` from the ape package.
- **hclust**: Use `plot.hclust` from the stats package.
- **dendrogram**: Use `plot.dendrogram` from the stats package.

The different plotting functions take different sets of arguments. When using `plot.phylo` (`mode="phylo"`), we have the following syntax:

```r
plot_dendrogram(x, mode="phylo", colbar = rainbow(11, start=0.7, end=0.1), edge.color = NULL, use.edge.length = FALSE, \dots)
```

The extra arguments not documented above:

- `colbar`: Color bar for the edges.
- `edge.color`: Edge colors. If NULL, then the colbar argument is used.
- `use.edge.length`: Passed to `plot.phylo`.
- `dots`: Additional arguments to pass to `plot.phylo`.

The syntax for `plot.hclust` (`mode="hclust"`):

```r
plot_dendrogram(x, mode="hclust", rect = 0, colbar = rainbow(rect),
    hang = 0.01, ann = FALSE, main = ",", sub = ",", xlab = ",",
    ylab = ",", \dots)
```

The extra arguments not documented above:

- `rect`: A numeric scalar, the number of groups to mark on the dendrogram. The dendrogram is cut into exactly `rect` groups and they are marked via the `rect.hclust` command. Set this to zero if you don’t want to mark any groups.
- `colbar`: The colors of the rectangles that mark the vertex groups via the `rect` argument.
- `hang`: Where to put the leaf nodes, this corresponds to the `hang` argument of `plot.hclust`.
- `ann`: Whether to annotate the plot, the `ann` argument of `plot.hclust`. 
power_centrality

• main The main title of the plot, the main argument of plot.hclust.
• sub The sub-title of the plot, the sub argument of plot.hclust.
• xlab The label on the horizontal axis, passed to plot.hclust.
• ylab The label on the vertical axis, passed to plot.hclust.
• dots Additional arguments to pass to plot.hclust.

The syntax for plot.dendrogram (mode="dendrogram"):

plot_dendrogram(x, \dots)

The extra arguments are simply passed to as.dendrogram.

Value

Returns whatever the return value was from the plotting function, plot.phylo, plot.dendrogram or plot.hclust.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

g <- make_full_graph(5) + make_full_graph(5)
hrg <- fit_hrg(g)
plot_dendrogram(hrg)

---

power_centrality  Find Bonacich Power Centrality Scores of Network Positions

Description

power_centrality takes a graph (dat) and returns the Boncich power centralities of positions (selected by nodes). The decay rate for power contributions is specified by exponent (1 by default).

Usage

power_centrality(  
  graph,  
  nodes = V(graph),  
  loops = FALSE,  
  exponent = 1,  
  rescale = FALSE,  
  tol = 1e-07,  
  sparse = TRUE
)
power_centrality

Arguments

- **graph**: the input graph.
- **nodes**: vertex sequence indicating which vertices are to be included in the calculation. By default, all vertices are included.
- **loops**: boolean indicating whether or not the diagonal should be treated as valid data. Set this true if and only if the data can contain loops. loops is FALSE by default.
- **exponent**: exponent (decay rate) for the Bonacich power centrality score; can be negative.
- **rescale**: if true, centrality scores are rescaled such that they sum to 1.
- **tol**: tolerance for near-singularities during matrix inversion (see solve).
- **sparse**: Logical scalar, whether to use sparse matrices for the calculation. The ‘Matrix’ package is required for sparse matrix support.

Details

Bonacich’s power centrality measure is defined by $C_{BP}(\alpha, \beta) = \alpha (I - \beta A)^{-1} A 1$, where $\beta$ is an attenuation parameter (set here by exponent) and $A$ is the graph adjacency matrix. (The coefficient $\alpha$ acts as a scaling parameter, and is set here (following Bonacich (1987)) such that the sum of squared scores is equal to the number of vertices. This allows 1 to be used as a reference value for the “middle” of the centrality range.) When $\beta \to 1/\lambda_{A1}$ (the reciprocal of the largest eigenvalue of $A$), this is to within a constant multiple of the familiar eigenvector centrality score; for other values of $\beta$, the behavior of the measure is quite different. In particular, $\beta$ gives positive and negative weight to even and odd walks, respectively, as can be seen from the series expansion $C_{BP}(\alpha, \beta) = \alpha \sum_{k=0}^{\infty} \beta^k A^{k+1} 1$ which converges so long as $|\beta| < 1/\lambda_{A1}$. The magnitude of $\beta$ controls the influence of distant actors on ego’s centrality score, with larger magnitudes indicating slower rates of decay. (High rates, hence, imply a greater sensitivity to edge effects.)

Interpretively, the Bonacich power measure corresponds to the notion that the power of a vertex is recursively defined by the sum of the power of its alters. The nature of the recursion involved is then controlled by the power exponent: positive values imply that vertices become more powerful as their alters become more powerful (as occurs in cooperative relations), while negative values imply that vertices become more powerful only as their alters become weaker (as occurs in competitive or antagonistic relations). The magnitude of the exponent indicates the tendency of the effect to decay across long walks; higher magnitudes imply slower decay. One interesting feature of this measure is its relative instability to changes in exponent magnitude (particularly in the negative case). If your theory motivates use of this measure, you should be very careful to choose a decay parameter on a non-ad hoc basis.

Value

A vector, containing the centrality scores.

Warning

Singular adjacency matrices cause no end of headaches for this algorithm; thus, the routine may fail in certain cases. This will be fixed when I get a better algorithm. power_centrality will not symmetrize your data before extracting eigenvectors; don’t send this routine asymmetric matrices unless you really mean to do so.

Note

This function was ported (ie. copied) from the SNA package.
Author(s)

Carter T. Butts (http://www.faculty.uci.edu/profile.cfm?faculty_id=5057), ported to igraph by Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

eigen_centrality and alpha_centrality

Examples

```r
# Generate some test data from Bonacich, 1987:
g.c <- graph( c(1,2,1,3,2,4,3,5), dir=FALSE)
g.d <- graph( c(1,2,1,3,1,4,2,5,3,6,4,7), dir=FALSE)
g.e <- graph( c(1,2,1,3,1,4,2,5,2,6,3,7,3,8,4,9,4,10), dir=FALSE)
g.f <- graph( c(1,2,1,3,1,4,2,5,2,6,2,7,3,8,3,9,3,10,4,11,4,12,4,13), dir=FALSE)

# Compute power centrality scores
for (e in seq(-0.5,.5, by=0.1)) {
  print(round(power_centrality(g.c, exp=e)[c(1,2,4)], 2))
}

for (e in seq(-0.4,.4, by=0.1)) {
  print(round(power_centrality(g.d, exp=e)[c(1,2,5)], 2))
}

for (e in seq(-0.4,.4, by=0.1)) {
  print(round(power_centrality(g.e, exp=e)[c(1,2,5)], 2))
}

for (e in seq(-0.4,.4, by=0.1)) {
  print(round(power_centrality(g.f, exp=e)[c(1,2,5)], 2))
}
```

Description

`predict_edges` *Predict edges based on a hierarchical random graph model*

`predict_edges` uses a hierarchical random graph model to predict missing edges from a network. This is done by sampling hierarchical models around the optimum model, proportionally to their likelihood. The MCMC sampling is stated from `hrg`, if it is given and the `start` argument is set to `TRUE`. Otherwise a HRG is fitted to the graph first.
predict_edges

Usage

predict_edges(
  graph,
  hrg = NULL,
  start = FALSE,
  num.samples = 10000,
  num.bins = 25
)

Arguments

graph The graph to fit the model to. Edge directions are ignored in directed graphs.
hrg A hierarchical random graph model, in the form of an igraphHRG object. predict_edges allow this to be NULL as well, then a HRG is fitted to the graph first, from a random starting point.
start Logical, whether to start the fitting/sampling from the supplied igraphHRG object, or from a random starting point.
num.samples Number of samples to use for consensus generation or missing edge prediction.
num.bins Number of bins for the edge probabilities. Give a higher number for a more accurate prediction.

Value

A list with entries:

edges The predicted edges, in a two-column matrix of vertex ids.
prob Probabilities of these edges, according to the fitted model.
hrg The (supplied or fitted) hierarchical random graph model.

References


See Also

Other hierarchical random graph functions: consensus_tree(), fit_hrg(), hrg-methods, hrg_tree(), hrg(), print.igraphHRGConsensus(), print.igraphHRG(), sample_hrg()

Examples

## We are not running these examples any more, because they
## take a long time (~15 seconds) to run and this is against the CRAN
## repository policy. Copy and paste them by hand to your R prompt if
## you want to run them.

## Not run:
## A graph with two dense groups
g <- sample_gnp(10, p=1/2) + sample_gnp(10, p=1/2)
hrg <- fit_hrg(g)
hrg

## The consensus tree for it
consensus_tree(g, hrg=hrg, start=TRUE)

## Prediction of missing edges
g2 <- make_full_graph(4) + (make_full_graph(4) - path(1,2))
predict_edges(g2)

## End(Not run)

---

**print.igraph**  
*Print graphs to the terminal*

**Description**

These functions attempt to print a graph to the terminal in a human readable form.

**Usage**

```r
## S3 method for class 'igraph'
print(
  x,
  full = igraph_opt("print.full"),
  graph.attributes = igraph_opt("print.graph.attributes"),
  vertex.attributes = igraph_opt("print.vertex.attributes"),
  edge.attributes = igraph_opt("print.edge.attributes"),
  names = TRUE,
  max.lines = igraph_opt("auto.print.lines"),
  ...
)

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

**Arguments**

- `x`: The graph to print.
- `full`: Logical scalar, whether to print the graph structure itself as well.
- `graph.attributes`: Logical constant, whether to print graph attributes.
- `vertex.attributes`: Logical constant, whether to print vertex attributes.
- `edge.attributes`: Logical constant, whether to print edge attributes.
- `names`: Logical constant, whether to print symbolic vertex names (ie. the name vertex attribute) or vertex ids.
- `max.lines`: The maximum number of lines to use. The rest of the output will be truncated.
- `...`: Additional arguments.
- `object`: The graph of which the summary will be printed.
print.igraph

Details

summary.igraph prints the number of vertices, edges and whether the graph is directed.

print_all prints the same information, and also lists the edges, and optionally graph, vertex and/or edge attributes.

print.igraph behaves either as summary.igraph or print_all depending on the full argument. See also the 'print.full' igraph option and igraph_opt.

The graph summary printed by summary.igraph (and print.igraph and print_all) consists one or more lines. The first line contains the basic properties of the graph, and the rest contains its attributes. Here is an example, a small star graph with weighted directed edges and named vertices:

```txt
IGRAPH badcafe DNW- 10 9 -- In-star + attr: name (g/c), mode (g/c), center (g/n), name (v/c), weight (e/n)
```

The first line always starts with IGRAPH, showing you that the object is an igraph graph. Then a seven character code is printed, this the first seven characters of the unique id of the graph. See graph_id for more. Then a four letter long code string is printed. The first letter distinguishes between directed ('D') and undirected ('U') graphs. The second letter is 'N' for named graphs, i.e. graphs with the name vertex attribute set. The third letter is 'W' for weighted graphs, i.e. graphs with the weight edge attribute set. The fourth letter is 'B' for bipartite graphs, i.e. for graphs with the type vertex attribute set.

Then, after two dashes, the name of the graph is printed, if it has one, i.e. if the name graph attribute is set.

From the second line, the attributes of the graph are listed, separated by a comma. After the attribute names, the kind of the attribute – graph ('g'), vertex ('v') or edge ('e') – is denoted, and the type of the attribute as well, character ('c'), numeric ('n'), logical ('l'), or other ('x').

As of igraph 0.4 print_all and print.igraph use the max.print option, see options for details.

As of igraph 1.1.1, the str.igraph function is defunct, use print_all().

Value

All these functions return the graph invisibly.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

```r
g <- make_ring(10)
g summary(g)
```
print.igraph.es: Print an edge sequence to the screen

Description

For long edge sequences, the printing is truncated to fit to the screen. Use `print` explicitly and the `codefull` argument to see the full sequence.

Usage

```r
## S3 method for class 'igraph.es'
print(x, full = igraph_opt("print.full"), ...)
```

Arguments

- `x`: An edge sequence.
- `full`: Whether to show the full sequence, or truncate the output to the screen size.
- `...`: Currently ignored.

Details

Edge sequences created with the double bracket operator are printed differently, together with all attributes of the edges in the sequence, as a table.

Value

The edge sequence, invisibly.

See Also

Other vertex and edge sequences: `E()`, `V()`, `igraph-es-attributes`, `igraph-es-indexing2`, `igraph-es-indexing`, `igraph-vs-attributes`, `igraph-vs-indexing2`, `igraph-vs-indexing`, `print.igraph.vs()`

Examples

```r
# Unnamed graphs
g <- make_ring(10)
E(g)

# Named graphs
g2 <- make_ring(10) %>%
  set_vertex_attr("name", value = LETTERS[1:10])
E(g2)

# All edges in a long sequence
g3 <- make_ring(200)
E(g3)
E(g3) %>% print(full = TRUE)

# Metadata
g4 <- make_ring(10) %>%
  set_vertex_attr("name", value = LETTERS[1:10]) %>%
```
set_edge_attr("weight", value = 1:10) %>%
  set_edge_attr("color", value = "green")
E(g4)
E(g4)[[]]
E(g4)[[1:5]]

print.igraph.vs | Show a vertex sequence on the screen

Description

For long vertex sequences, the printing is truncated to fit to the screen. Use `print` explicitly and the `full` argument to see the full sequence.

Usage

```r
## S3 method for class 'igraph.vs'
print(x, full = igraph_opt("print.full"), ...)
```

Arguments

- `x` A vertex sequence.
- `full` Whether to show the full sequence, or truncate the output to the screen size.
- `...` These arguments are currently ignored.

Details

Vertex sequence created with the double bracket operator are printed differently, together with all attributes of the vertices in the sequence, as a table.

Value

The vertex sequence, invisibly.

See Also


Examples

```r
# Unnamed graphs
g <- make_ring(10)
V(g)

# Named graphs
g2 <- make_ring(10) %>%
  set_vertex_attr("name", value = LETTERS[1:10])
V(g2)

# All vertices in the sequence
g3 <- make_ring(1000)
```
V(g3)
print(V(g3), full = TRUE)

# Metadata
g4 <- make_ring(10) %>%
  set_vertex_attr("name", value = LETTERS[1:10]) %>%
  set_vertex_attr("color", value = "red")
V(g4)[[1]]
V(g4)[[2:5, 7:8]]

print.igraphHRG
Print a hierarchical random graph model to the screen

Description

igraphHRG objects can be printed to the screen in two forms: as a tree or as a list, depending on the
type argument of the print function. By default the auto type is used, which selects tree for small
graphs and simple (=list) for bigger ones. The tree format looks like this:

Hierarchical random graph, at level 3:
g1      p=  0
  |- g15   p=0.33  1
  |     |- g13  p=0.88  6 3 9 4 2 10 7 5 8
  |     |- g8    p= 0.5
  |          |- g16 p= 0.2  20 14 17 19 11 15 16 13
  |          |- g5    p=  0 12 18

This is a graph with 20 vertices, and the top three levels of the fitted hierarchical random graph are
printed. The root node of the HRG is always vertex group #1 (‘g1’ in the printout). Vertex pairs
in the left subtree of g1 connect to vertices in the right subtree with probability zero, according to
the fitted model. g1 has two subgroups, g15 and g8. g15 has a subgroup of a single vertex (vertex
1), and another larger subgroup that contains vertices 6, 3, etc. on lower levels, etc. The plain
printing is simpler and faster to produce, but less visual:

Hierarchical random graph:
g1 p=0.0  ->  g12  g10  g2 p=1.0  ->  7 10  g3 p=1.0  ->  g18  14
  g4 p=1.0  ->  g17  15  g5 p=0.4  ->  g15  17  g6 p=0.0  ->  1 4
  g7 p=1.0  ->  11  16  g8 p=0.1  ->  g9  3  g9 p=0.3  ->  g11  g16
  g10 p=0.2  ->  g4  5  g11 p=1.0  ->  g6  5  g12 p=0.8  ->  g8  8
  g13 p=0.0  ->  g14  9  g14 p=1.0  ->  2 6  g15 p=0.2  ->  g19  18
  g16 p=1.0  ->  g13  g2  g17 p=0.5  ->  g7  13  g18 p=1.0  ->  12 19
  g19 p=0.7  ->  g3  20

It lists the two subgroups of each internal node, in as many columns as the screen width allows.

Usage

## S3 method for class 'igraphHRG'
print(x, type = c("auto", "tree", "plain"), level = 3, ...)
Arguments

- **x**  
  igraphHRG object to print.

- **type**  
  How to print the dendrogram, see details below.

- **level**  
  The number of top levels to print from the dendrogram.

- **...**  
  Additional arguments, not used currently.

Value

The hierarchical random graph model itself, invisibly.

See Also

Other hierarchical random graph functions: `consensus_tree()`, `fit_hrg()`, `hrg-methods`, `hrg_tree()`, `hrg()`, `predict_edges()`, `print.igraphHRGConsensus()`, `sample_hrg()`

---

**Description**

Consensus dendrograms (igraphHRGConsensus objects) are printed simply by listing the children of each internal node of the dendrogram:

HRG consensus tree:

```
g1 -> 11 12 13 14 15 16 17 18 19 20
```

```
g2 -> 1 2 3 4 5 6 7 8 9 10
```

```
g3 -> g1 g2
```

The root of the dendrogram is g3 (because it has no incoming edges), and it has two subgroups, g1 and g2.

**Usage**

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

**Arguments**

- **x**  
  igraphHRGConsensus object to print.

- **...**  
  Ignored.

**Value**

The input object, invisibly, to allow method chaining.

**See Also**

Other hierarchical random graph functions: `consensus_tree()`, `fit_hrg()`, `hrg-methods`, `hrg_tree()`, `hrg()`, `predict_edges()`, `print.igraphHRG()`, `sample_hrg()`
print.nexusDatasetInfo

Query and download from the Nexus network repository

Description

The Nexus network repository is an online collection of network data sets. These functions can be used to query it and download data from it, directly as an igraph graph.

Usage

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

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

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

nexus_list(
  tags = NULL,
  offset = 0,
  limit = 10,
  operator = c("or", "and"),
  order = c("date", "name", "popularity"),
  nexus.url = igraph_opt("nexus.url")
)

nexus_info(id, nexus.url = igraph_opt("nexus.url"))

nexus_get(
  id,
  offset = 0,
  order = c("date", "name", "popularity"),
  nexus.url = igraph_opt("nexus.url")
)

nexus_search(
  q,
  offset = 0,
  limit = 10,
  order = c("date", "name", "popularity"),
  nexus.url = igraph_opt("nexus.url")
)

## S3 method for class 'nexusDatasetInfoList'
x[i]
```
Arguments

- **x, object**
  - The `nexusDatasetInfo` object to print.

- **...**
  - Currently ignored.

- **tags**
  - A character vector, the tags that are searched. If not given (or NULL), then all datasets are listed.

- **offset**
  - An offset to select part of the results. Results are listed from offset+1.

- **limit**
  - The maximum number of results to return.

- **operator**
  - A character scalar. If ‘or’ (the default), then all datasets that have at least one of the given tags, are returned. If it if ‘and’, then only datasets that have all the given tags, are returned.

- **order**
  - The ordering of the results, possible values are: ‘date’, ‘name’, ‘popularity’.

- **nexus.url**
  - The URL of the Nexus server. Don’t change this from the default, unless you set up your own Nexus server.

- **id**
  - The numeric or character id of the data set to query or download. Instead of the data set ids, it is possible to supply a `nexusDatasetInfo` or `nexusDatasetInfoList` object here directly and then the query is done on the corresponding data set(s).

- **q**
  - Nexus search string. See examples below.

- **i**
  - Index.

Details

Nexus is an online repository of networks, with an API that allow programmatic queries against it, and programmatic data download as well.

The `nexus_list` and `nexus_info` functions query the online database. They both return `nexusDatasetInfo` objects. `nexus_info` returns more information than `nexus_list`.

`nexus_search` searches Nexus, and returns a list of data sets, as `nexusDatasetInfo` objects. See below for some search examples.

`nexus_get` downloads a data set from Nexus, based on its numeric id, or based on a Nexus search string. For search strings, only the first search hit is downloaded, but see also the `offset` argument.
(If there are not data sets found, then the function returns an error.)

The `nexusDatasetInfo` objects returned by `nexus_list` have the following fields:

- **id** The numeric id of the dataset.
- **sid** The character id of the dataset.
- **name** Character scalar, the name of the dataset.
- **vertices/edges** Character, the number of vertices and edges in the graph(s). Vertices and edges are separated by a slash, and if the data set consists of multiple networks, then they are separated by spaces.
- **tags** Character vector, the tags of the dataset. Directed graph have the tags ‘directed’. Undirected graphs are tagged as ‘undirected’. Other common tags are: ‘weighted’, ‘bipartite’, ‘social network’, etc.
- **networks** The ids and names of the networks in the data set. The numeric and character id are separated by a slash, and multiple networks are separated by spaces.

`nexusDatasetInfo` objects returned by `nexus_info` have the following additional fields:

- **date** Character scalar, e.g. ‘2011-01-09’, the date when the dataset was added to the database.
formats Character vector, the data formats in which the data set is available. The various formats are separated by semicolons.

licence Character scalar, the licence of the dataset.

licence url Character scalar, the URL of the licence of the dataset. Please make sure you consult this before using a dataset.

summary Character scalar, the short description of the dataset, this is usually a single sentence.

description Character scalar, the full description of the dataset.

citation Character scalar, the paper(s) describing the dataset. Please cite these papers if you are using the dataset in your research, the licence of most datasets requires this.

attributes A list of lists, each list entry is a graph, vertex or edge attribute and has the following entries:

- **type** Type of the attribute, either ‘graph’, ‘vertex’ or ‘edge’.
- **datatype** Data type of the attribute, currently it can be ‘numeric’ and ‘string’.
- **name** Character scalar, the name of the attribute.
- **description** Character scalar, the description of the attribute.

The results of the Nexus queries are printed to the screen in a concise format, similar to the format of igraph graphs. A data set list (typically the result of `nexus_list` and `nexus_search`) looks like this:

```
NEXUS 1-5/18 -- data set list
[1] kaptail.4 39/109-223 #18 Kapferer tailor shop
[2] condmatcollab2003 31163/120029 #17 Condensed matter collaborations+
[3] condmatcollab 16726/47594 #16 Condensed matter collaborations+
[4] powergrid 4941/6594 #15 Western US power grid
[5] celegansneural 297/2359 #14 C. Elegans neural network
```

Each line here represents a data set, and the following information is given about them: the character id of the data set (e.g. `kaptail` or `powergrid`), the number of vertices and number of edges in the graph of the data sets. For data sets with multiple graphs, intervals are given here. Then the numeric id of the data set and the remaining space is filled with the name of the data set. Summary information about an individual Nexus data set is printed as

```
NEXUS B--- 39 109-223 #18 kaptail -- Kapferer tailor shop
+ tags: directed; social network; undirected
+ nets: 1/KAPFTI2; 2/KAPFTS2; 3/KAPFTI1; 4/KAPFTS1
```

This is very similar to the header that is used for printing igraph graphs, but there are some differences as well. The four characters after the NEXUS word give the most important properties of the graph(s): the first is ‘U’ for undirected and ‘D’ for directed graphs, and ‘B’ if the data set contains both directed and undirected graphs. The second is ‘N’ named graphs. The third character is ‘W’ for weighted graphs, the fourth is ‘B’ if the data set contains bipartite graphs. Then the number of vertices and number of edges are printed, for data sets with multiple graphs, the smallest and the largest values are given. Then comes the numeric id, and the string id of the data set. The end of the first line contains the name of the data set. The second row lists the data set tags, and the third row the networks that are included in the data set.

Detailed data set information is printed similarly, but it contains more fields.
Value

nexus_list and nexus_search return a list of nexusDatasetInfo objects. The list also has these attributes:

- **size** The number of data sets returned by the query.
- **totalsize** The total number of data sets found for the query.
- **offset** The offset parameter of the query.
- **limit** The limit parameter of the query.

nexus_info returns a single nexusDatasetInfo object.

nexus_get returns an igraph graph object, or a list of graph objects, if the data set consists of multiple networks.

Examples

```r
nexus_list(tag="weighted")
nexus_list(limit=3, order="name")
nexus_list(limit=3, order="name")[[1]]
nexus_info(2)
g <- nexus_get(2)
summary(g)
```

```
## Data sets related to 'US':
nexus_search("US")

## Search for data sets that have 'network' in their name:
nexus_search("name:network")

## Any word can match
nexus_search("blog or US or karate")
```

printer_callback

Create a printer callback function

Description

A printer callback function is a function can performs the actual printing. It has a number of subcommands, that are called by the printer package, in a form

```r
printer_callback("subcommand", argument1, argument2, ...)
```

See the examples below.

Usage

```r
printer_callback(fun)
```

Arguments

- **fun** The function to use as a printer callback function.
Details

The subcommands:

length The length of the data to print, the number of items, in natural units. E.g. for a list of objects, it is the number of objects.
min_width TODO
width Width of one item, if no items will be printed. TODO
print Argument: no. Do the actual printing, print no items.
done TODO

See Also

Other printer callbacks: is_printer_callback()

---

**printr**  
*Better printing of R packages*

Description

This package provides better printing of R packages.

---

**r_pal**  
*The default R palette*

Description

This is the default R palette, to be able to reproduce the colors of older igraph versions. Its colors are appropriate for categories, but they are not very attractive.

Usage

```r
r_pal(n)
```

Arguments

- `n` The number of colors to use, the maximum is eight.

Value

A character vector of color names.

See Also

Other palettes: categorical_pal(), diverging_pal(), sequential_pal()
radius

Radius of a graph

Description

The eccentricity of a vertex is its shortest path distance from the farthest other node in the graph. The smallest eccentricity in a graph is called its radius.

Usage

radius(graph, mode = c("all", "out", "in", "total"))

Arguments

graph
The input graph, it can be directed or undirected.

mode
Character constant, gives whether the shortest paths to or from the given vertices should be calculated for directed graphs. If `out` then the shortest paths from the vertex, if `in` then to it will be considered. If `all`, the default, then the corresponding undirected graph will be used, edge directions will be ignored. This argument is ignored for undirected graphs.

Details

The eccentricity of a vertex is calculated by measuring the shortest distance from (or to) the vertex, to (or from) all vertices in the graph, and taking the maximum. This implementation ignores vertex pairs that are in different components. Isolate vertices have eccentricity zero.

Value

A numeric scalar, the radius of the graph.

References


See Also

eccentricity for the underlying calculations, codedistances for general shortest path calculations.

Examples

g <- make_star(10, mode="undirected")
eccentricity(g)
radius(g)
random_walk

Description

random_walk performs a random walk on the graph and returns the vertices that the random walk passed through. random_edge_walk is the same but returns the edges that that random walk passed through.

Usage

random_walk(
  graph,     
  start,     
  steps,     
  mode = c("out", "in", "all", "total"),
  stuck = c("return", "error")
)

random_edge_walk(
  graph,     
  start,     
  steps,     
  weights = NULL,
  mode = c("out", "in", "all", "total"),
  stuck = c("return", "error")
)

Arguments

graph The input graph, might be undirected or directed.
start The start vertex.
steps The number of steps to make.
mode How to follow directed edges. "out" steps along the edge direction, "in" is opposite to that. "all" ignores edge directions. This argument is ignored for undirected graphs.
stuck What to do if the random walk gets stuck. "return" returns the partial walk, "error" raises an error.
weights The edge weights. Larger edge weights increase the probability that an edge is selected by the random walker. In other words, larger edge weights correspond to stronger connections. The 'weight' edge attribute is used if present. Supply 'NA' here if you want to ignore the 'weight' edge attribute.

Details

Do a random walk. From the given start vertex, take the given number of steps, choosing an edge from the actual vertex uniformly randomly. Edge directions are observed in directed graphs (see the mode argument as well). Multiple and loop edges are also observed.
For random walk, a vertex sequence containing the vertices along the walk. For random_edge_walk, an edge sequence containing the edges along the walk.

Examples

```r
## Stationary distribution of a Markov chain

g <- make_ring(10, directed = TRUE) %u% make_star(11, center = 11) + edge(11, 1)

ec <- eigen_centrality(g, directed = TRUE)$vector

pg <- page_rank(g, damping = 0.999)$vector

w <- random_walk(g, start = 1, steps = 10000)

## These are similar, but not exactly the same

cor(table(w), ec)

## But these are (almost) the same

cor(table(w), pg)
```

Description

The `read_graph` function is able to read graphs in various representations from a file, or from a http connection. Various formats are supported.

Usage

```r
read_graph(
  file,
  format = c("edgelist", "pajek", "ncol", "lgl", "graphml", "dimacs", "graphdb", "gml", "dl"),
  ...
)
```

Arguments

- **file** The connection to read from. This can be a local file, or a http or ftp connection. It can also be a character string with the file name or URI.
- **format** Character constant giving the file format. Right now edgelist, pajek, ncol, lgl, graphml, dimacs, graphdb, gml and dl are supported, the default is edgelist. As of igraph 0.4 this argument is case insensitive.
- **...** Additional arguments, see below.

Details

The `read_graph` function may have additional arguments depending on the file format (the `format` argument). See the details separately for each file format, below.
Value

A graph object.

Edge list format

This format is a simple text file with numeric vertex ids defining the edges. There is no need to have newline characters between the edges, a simple space will also do.

Additional arguments:

- **n** The number of vertices in the graph. If it is smaller than or equal to the largest integer in the file, then it is ignored; so it is safe to set it to zero (the default).
- **directed** Logical scalar, whether to create a directed graph. The default value is TRUE.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

write_graph

realize_degseq

Creating a graph from a given degree sequence, deterministically

Description

It is often useful to create a graph with given vertex degrees. This function creates such a graph in a deterministic manner.

Usage

realize_degseq(
  out.deg,
  in.deg = NULL,
  allowed.edge.types = c("simple", "loops", "multi", "all"),
  method = c("smallest", "largest", "index")
)

Arguments

- **out.deg** Numeric vector, the sequence of degrees (for undirected graphs) or out-degrees (for directed graphs). For undirected graphs its sum should be even. For directed graphs its sum should be the same as the sum of in.deg.
- **in.deg** For directed graph, the in-degree sequence. By default this is NULL and an undirected graph is created.
- **allowed.edge.types** Character, specifies the types of allowed edges. “simple” allows simple graphs only (no loops, no multiple edges). “multiple” allows multiple edges but disallows loop. “loops” allows loop edges but disallows multiple edges (currently unimplemented). “all” allows all types of edges. The default is “simple”.
- **method** Character, the method for generating the graph; see above.
Details

Simple undirected graphs are constructed using the Havel-Hakimi algorithm (undirected case), or the analogous Kleitman-Wang algorithm (directed case). These algorithms work by choosing an arbitrary vertex and connecting all its stubs to other vertices. This step is repeated until all degrees have been connected up.

The ‘method’ argument controls in which order the vertices are selected during the course of the algorithm.

The “smallest” method selects the vertex with the smallest remaining degree. The result is usually a graph with high negative degree assortativity. In the undirected case, this method is guaranteed to generate a connected graph, regardless of whether multi-edges are allowed, provided that a connected realization exists. In the directed case it tends to generate weakly connected graphs, but this is not guaranteed. This is the default method.

The “largest” method selects the vertex with the largest remaining degree. The result is usually a graph with high positive degree assortativity, and is often disconnected.

The “index” method selects the vertices in order of their index.

Value

The new graph object.

See Also

`sample_degseq` for a randomized variant that samples from graphs with the given degree sequence.

Examples

g <- realize_degseq(rep(2,100))
degree(g)
is_simple(g)

## Exponential degree distribution, with high positive assortativity.
## Loop and multiple edges are explicitly allowed.
## Note that we correct the degree sequence if its sum is odd.
degs <- sample(1:100, 100, replace=TRUE, prob=exp(-0.5*(1:100)))
if (sum(degs) %% 2 != 0) { degs[1] <- degs[1] + 1 }
g4 <- realize_degseq(degs, method="largest", allowed.edge.types="all")
all(degree(g4) == degs)

## Power-law degree distribution, no loops allowed but multiple edges
## are okay.
## Note that we correct the degree sequence if its sum is odd.
degs <- sample(1:100, 100, replace=TRUE, prob=(1:100)^-2)
if (sum(degs) %% 2 != 0) { degs[1] <- degs[1] + 1 }
g5 <- realize_degseq(degs, allowed.edge.types="multi")
all(degree(g5) == degs)
Reciprocity of graphs

Description

Calculates the reciprocity of a directed graph.

Usage

reciprocity(graph, ignore.loops = TRUE, mode = c("default", "ratio"))

Arguments

- graph: The graph object.
- ignore.loops: Logical constant, whether to ignore loop edges.
- mode: See below.

Details

The measure of reciprocity defines the proportion of mutual connections, in a directed graph. It is most commonly defined as the probability that the opposite counterpart of a directed edge is also included in the graph. Or in adjacency matrix notation: \( \sum_{ij} (A \cdot A')_{ij} \), where \( A \cdot A' \) is the element-wise product of matrix \( A \) and its transpose. This measure is calculated if the mode argument is default.

Prior to igraph version 0.6, another measure was implemented, defined as the probability of mutual connection between a vertex pair, if we know that there is a (possibly non-mutual) connection between them. In other words, (unordered) vertex pairs are classified into three groups: (1) not-connected, (2) non-reciprocally connected, (3) reciprocally connected. The result is the size of group (3), divided by the sum of group sizes (2)+(3). This measure is calculated if mode is ratio.

Value

A numeric scalar between zero and one.

Author(s)

Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com>

Examples

```r
g <- sample_gnp(20, 5/20, directed=TRUE)
reciprocity(g)
```
rep.igraph  Replicate a graph multiple times

Description

The new graph will contain the input graph the given number of times, as unconnected components.

Usage

```r
## S3 method for class 'igraph'
rep(x, n, mark = TRUE, ...)
```

```r
## S3 method for class 'igraph'
x * n
```

Arguments

- `x`: The input graph.
- `n`: Number of times to replicate it.
- `mark`: Whether to mark the vertices with a `which` attribute, an integer number denoting which replication the vertex is coming from.
- `...`: Additional arguments to satisfy S3 requirements, currently ignored.

Examples

```r
rings <- make_ring(5) * 5
```

rev.igraph.es  Reverse the order in an edge sequence

Description

Reverse the order in an edge sequence

Usage

```r
## S3 method for class 'igraph.es'
rev(x)
```

Arguments

- `x`: The edge sequence to reverse.

Value

The reversed edge sequence.
See Also

Other vertex and edge sequence operations: \texttt{c.igraph.es()}, \texttt{c.igraph.vs()}, \texttt{difference.igraph.es()}, \texttt{difference.igraph.vs()}, \texttt{igraph-es-indexing2}, \texttt{igraph-es-indexing}, \texttt{igraph-vs-indexing2}, \texttt{igraph-vs-indexing}, \texttt{intersection.igraph.es()}, \texttt{intersection.igraph.vs()}, \texttt{rev.igraph.es()}, \texttt{rev.igraph.vs()}, \texttt{union.igraph.es()}, \texttt{union.igraph.vs()}, \texttt{unique.igraph.es()}, \texttt{unique.igraph.vs()}

Examples

\begin{verbatim}
g <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
E(g)
E(g) %>> rev()
\end{verbatim}

---

\textbf{rev.igraph.vs} \hspace{2cm} Reverse the order in a vertex sequence

Description

Reverse the order in a vertex sequence

Usage

\begin{verbatim}
## S3 method for class 'igraph.vs'
rev(x)
\end{verbatim}

Arguments

\begin{itemize}
\item \textbf{x} \hspace{1cm} The vertex sequence to reverse.
\end{itemize}

Value

The reversed vertex sequence.

See Also

Other vertex and edge sequence operations: \texttt{c.igraph.es()}, \texttt{c.igraph.vs()}, \texttt{difference.igraph.es()}, \texttt{difference.igraph.vs()}, \texttt{igraph-es-indexing2}, \texttt{igraph-es-indexing}, \texttt{igraph-vs-indexing2}, \texttt{igraph-vs-indexing}, \texttt{intersection.igraph.es()}, \texttt{intersection.igraph.vs()}, \texttt{rev.igraph.es()}, \texttt{rev.igraph.vs()}, \texttt{union.igraph.es()}, \texttt{union.igraph.vs()}, \texttt{unique.igraph.es()}, \texttt{unique.igraph.vs()}

Examples

\begin{verbatim}
g <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
V(g) %>> rev()
\end{verbatim}
reverse_edges  

**Reverse edges in a graph**

**Description**

The new graph will contain the same vertices, edges and attributes as the original graph, except that the direction of the edges selected by their edge IDs in the `eids` argument will be reversed. When reversing all edges, this operation is also known as graph transpose.

**Usage**

```r
reverse_edges(graph, eids = E(graph))
```

### S3 method for class 'igraph'

```r
t(x)
```

**Arguments**

- **graph**: The input graph.
- **eids**: The edge IDs of the edges to reverse.
- **x**: The input graph.

**Value**

The result graph where the direction of the edges with the given IDs are reversed.

**Examples**

```r
g <- make_graph( ~ 1+2, 2+3, 3+4 )
reverse_edges(g, 2)
```

rewire  

**Rewiring edges of a graph**

**Description**

See the links below for the implemented rewiring methods.

**Usage**

```r
rewire(graph, with)
```

**Arguments**

- **graph**: The graph to rewire
- **with**: A function call to one of the rewiring methods, see details below.

**Value**

The rewired graph.
rglplot

3D plotting of graphs with OpenGL

Description
Using the `rgl` package, rglplot plots a graph in 3D. The plot can be zoomed, rotated, shifted, etc. but the coordinates of the vertices is fixed.

Usage
```r
rglplot(x, ...)
```

Arguments
- `x` The graph to plot.
- `...` Additional arguments, see `igraph.plotting` for the details

Details
Note that rglplot is considered to be highly experimental. It is not very useful either. See `igraph.plotting` for the possible arguments.

Value
NULL, invisibly.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

See Also
`igraph.plotting`, `plot.igraph` for the 2D version, `tkplot` for interactive graph drawing in 2D.

Examples
```r
## Not run:
g <- make_lattice(c(5,5,5))
coords <- layout_with_fr(g, dim=3)
rglplot(g, layout=coords)
## End(Not run)
```
running_mean  Running mean of a time series

Description

running_mean calculates the running mean in a vector with the given bin width.

Usage

running_mean(v, binwidth)

Arguments

v  The numeric vector.
binwidth  Numeric constant, the size of the bin, should be meaningful, ie. smaller than the length of v.

Details

The running mean of v is a w vector of length length(v)-binwidth+1. The first element of w id the average of the first binwidth elements of v, the second element of w is the average of elements 2:(binwidth+1), etc.

Value

A numeric vector of length length(v)-binwidth+1

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

running_mean(1:100, 10)

sample_  Sample from a random graph model

Description

Generic function for sampling from network models.

Usage

sample_(...)

Arguments

...  Parameters, see details below.
sample_bipartite

**Details**

TODO

**Examples**

```r
def pref_matrix = cbind(c(0.8, 0.1), c(0.1, 0.7))
def blocky = sample_sbm(n = 20, pref.matrix = pref_matrix,
block.sizes = c(10, 10))

def blocky2 = pref_matrix %>%
sample_sbm(n = 20, block.sizes = c(10, 10))

## Arguments are passed on from sample_ to sample_sbm

def blocky3 <- pref_matrix %>%
sample_sbm(n = 20, block.sizes = c(10, 10))
```

---

**Description**

Generate bipartite graphs using the Erdos-Renyi model

**Usage**

```r
sample_bipartite(
  n1,
  n2,
  type = c("gnp", "gnm"),
  p,
  m,
  directed = FALSE,
  mode = c("out", "in", "all")
)
```

**Arguments**

- **n1**: Integer scalar, the number of bottom vertices.
- **n2**: Integer scalar, the number of top vertices.
- **type**: Character scalar, the type of the graph. ‘gnp’ creates a $G(n, p)$ graph, ‘gnm’ creates a $G(n, m)$ graph. See details below.
- **p**: Real scalar, connection probability for $G(n, p)$ graphs. Should not be given for $G(n, m)$ graphs.
- **m**: Integer scalar, the number of edges for $G(n, p)$ graphs. Should not be given for $G(n, m)$ graphs.
- **directed**: Logical scalar, whether to create a directed graph. See also the mode argument.
mode

Character scalar, specifies how to direct the edges in directed graphs. If it is ‘out’, then directed edges point from bottom vertices to top vertices. If it is ‘in’, edges point from top vertices to bottom vertices. ‘out’ and ‘in’ do not generate mutual edges. If this argument is ‘all’, then each edge direction is considered independently and mutual edges might be generated. This argument is ignored for undirected graphs.

Passed to sample_bipartite.

Details

Similarly to unipartite (one-mode) networks, we can define the $G(n, p)$, and $G(n, m)$ graph classes for bipartite graphs, via their generating process. In $G(n, p)$ every possible edge between top and bottom vertices is realized with probability $p$, independently of the rest of the edges. In $G(n, m)$, we uniformly choose $m$ edges to realize.

Value

A bipartite igraph graph.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

sample_gnp for the unipartite version.

Examples

## empty graph
sample_bipartite(10, 5, p=0)

## full graph
sample_bipartite(10, 5, p=1)

## random bipartite graph
sample_bipartite(10, 5, p=.1)

## directed bipartite graph, G(n,m)
sample_bipartite(10, 5, type="Gnm", m=20, directed=TRUE, mode="all")
Usage

```r
sample_correlated_gnp(
  old.graph,
  corr,
  p = edge_density(old.graph),
  permutation = NULL
)
```

Arguments

- `old.graph`: The original graph.
- `corr`: A scalar in the unit interval, the target Pearson correlation between the adjacency matrices of the original and the generated graph (the adjacency matrix being used as a vector).
- `p`: A numeric scalar, the probability of an edge between two vertices, it must in the open (0,1) interval. The default is the empirical edge density of the graph. If you are resampling an Erdos-Renyi graph and you know the original edge probability of the Erdos-Renyi model, you should supply that explicitly.
- `permutation`: A numeric vector, a permutation vector that is applied on the vertices of the first graph, to get the second graph. If NULL, the vertices are not permuted.

Details

Please see the reference given below.

Value

An unweighted graph of the same size as `old.graph` such that the correlation coefficient between the entries of the two adjacency matrices is `corr`. Note each pair of corresponding matrix entries is a pair of correlated Bernoulli random variables.

References


See Also

- `sample_correlated_gnp_pair`
- `sample_gnp`

Examples

```r
g <- sample_gnp(1000, .1)
g2 <- sample_correlated_gnp(g, corr = 0.5)
cor(as.vector(g[]), as.vector(g2[]))
g
g2
```
Sample a pair of correlated \( G(n,p) \) random graphs

**Description**

Sample a new graph by perturbing the adjacency matrix of a given graph and shuffling its vertices.

**Usage**

```r
sample_correlated_gnp_pair(n, corr, p, directed = FALSE, permutation = NULL)
```

**Arguments**

- `n` Numeric scalar, the number of vertices for the sampled graphs.
- `corr` A scalar in the unit interval, the target Pearson correlation between the adjacency matrices of the original the generated graph (the adjacency matrix being used as a vector).
- `p` A numeric scalar, the probability of an edge between two vertices, it must in the open (0,1) interval.
- `directed` Logical scalar, whether to generate directed graphs.
- `permutation` A numeric vector, a permutation vector that is applied on the vertices of the first graph, to get the second graph. If `NULL`, the vertices are not permuted.

**Details**

Please see the reference given below.

**Value**

A list of two igraph objects, named `graph1` and `graph2`, which are two graphs whose adjacency matrix entries are correlated with `corr`.

**References**


**See Also**

`sample_correlated_gnp`, `sample_gnp`.

**Examples**

```r
gg <- sample_correlated_gnp_pair(n = 10, corr = .8, p = .5, directed = FALSE)
gg
cor(as.vector(gg[[1]][,]), as.vector(gg[[2]][,]))
```
Generate random graphs with a given degree sequence

Description

It is often useful to create a graph with given vertex degrees. This function creates such a graph in a randomized manner.

Usage

```r
sample_degseq(
  out.deg,
  in.deg = NULL,
  method = c("simple", "vl", "simple.no.multiple", "simple.no.multiple.uniform")
)
```

degseq(..., deterministic = FALSE)

Arguments

- `out.deg`: Numeric vector, the sequence of degrees (for undirected graphs) or out-degrees (for directed graphs). For undirected graphs its sum should be even. For directed graphs its sum should be the same as the sum of `in.deg`.
- `in.deg`: For directed graph, the in-degree sequence. By default this is `NULL` and an undirected graph is created.
- `method`: Character, the method for generating the graph. Right now the “simple”, “simple.no.multiple” and “vl” methods are implemented.
- `...`: Passed to `realize_degseq` if ‘deterministic’ is true, or to `sample_degseq` otherwise.
- `deterministic`: Whether the construction should be deterministic

Details

The “simple” method connects the out-stubs of the edges (undirected graphs) or the out-stubs and in-stubs (directed graphs) together. This way loop edges and also multiple edges may be generated. This method is not adequate if one needs to generate simple graphs with a given degree sequence. The multiple and loop edges can be deleted, but then the degree sequence is distorted and there is nothing to ensure that the graphs are sampled uniformly.

The “simple.no.multiple” method is similar to “simple”, but tries to avoid multiple and loop edges and restarts the generation from scratch if it gets stuck. It is not guaranteed to sample uniformly from the space of all possible graphs with the given sequence, but it is relatively fast and it will eventually succeed if the provided degree sequence is graphical, but there is no upper bound on the number of iterations.

The “simple.no.multiple.uniform” method is a variant of “simple.no.multiple” with the added benefit of sampling uniformly from the set of all possible simple graphs with the given degree sequence. Ensuring uniformity has some performance implications, though.

The “vl” method is a more sophisticated generator. The algorithm and the implementation was done by Fabien Viger and Matthieu Latapy. This generator always generates undirected, connected simple graphs, it is an error to pass the `in.deg` argument to it. The algorithm relies on first creating
an initial (possibly unconnected) simple undirected graph with the given degree sequence (if this is possible at all). Then some rewiring is done to make the graph connected. Finally a Monte-Carlo algorithm is used to randomize the graph. The “vl” samples from the undirected, connected simple graphs uniformly.

Value

The new graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

sample_gnp, sample_pa, simplify to get rid of the multiple and/or loops edges, realize_degseq for a deterministic variant.

Examples

```r
## The simple generator
g <- sample_degseq(rep(2,100))
degree(g)
is_simple(g)  # sometimes TRUE, but can be FALSE
g2 <- sample_degseq(1:10, 10:1)
degree(g2, mode="out")
degree(g2, mode="in")

## The vl generator
g3 <- sample_degseq(rep(2,100), method="vl")
degree(g3)
is_simple(g3)  # always TRUE

## Exponential degree distribution
## Note, that we correct the degree sequence if its sum is odd
degs <- sample(1:100, 100, replace=TRUE, prob=exp(-0.5*1:100))
if (sum(dgs) %% 2 != 0) { degs[1] <- degs[1] + 1 }
g4 <- sample_degseq(dgs, method="vl")
all(degree(g4) == degs)

## Power-law degree distribution
## Note, that we correct the degree sequence if its sum is odd
degs <- sample(1:100, 100, replace=TRUE, prob=(1:100)^(-2))
if (sum(dgs) %% 2 != 0) { degs[1] <- degs[1] + 1 }
g5 <- sample_degseq(dgs, method="vl")
all(degree(g5) == degs)
```

---

**sample_dirichlet**

Sample from a Dirichlet distribution

**Description**

Sample finite-dimensional vectors to use as latent position vectors in random dot product graphs.
Usage

```r
sample_dirichlet(n, alpha)
```

Arguments

- **n**: Integer scalar, the sample size.
- **alpha**: Numeric vector, the vector of \( \alpha \) parameter for the Dirichlet distribution.

Details

`sample_dirichlet` generates samples from the Dirichlet distribution with given \( \alpha \) parameter. The sample is drawn from \( \text{length}(\alpha)-1 \)-simplex.

Value

A \( \text{dim} \) (length of the alpha vector for `sample_dirichlet`) times \( n \) matrix, whose columns are the sample vectors.

See Also

Other latent position vector samplers: `sample_sphere_surface()`, `sample_sphere_volume()`

Examples

```r
lpvs.dir <- sample_dirichlet(n=20, alpha=rep(1, 10))
RDP.graph.2 <- sample_dot_product(lpvs.dir)
colSums(lpvs.dir)
```

---

**sample_dot_product**

Generate random graphs according to the random dot product graph model

Description

In this model, each vertex is represented by a latent position vector. Probability of an edge between two vertices are given by the dot product of their latent position vectors.

Usage

```r
sample_dot_product(vecs, directed = FALSE)
```

```r
dot_product(...)```

Arguments

- **vecs**: A numeric matrix in which each latent position vector is a column.
- **directed**: A logical scalar, TRUE if the generated graph should be directed.
- **...**: Passed to `sample_dot_product`. 

---

---

---
sample_fitness

Random graphs from vertex fitness scores

Description

This function generates a non-growing random graph with edge probabilities proportional to node fitness scores.

Usage

```
sample_fitness(
  no.of.edges,
  fitness.out,
  fitness.in = NULL,
  loops = FALSE,
  multiple = FALSE
)
```
sample_fitness

Arguments

no.of.edges The number of edges in the generated graph.

fitness.out A numeric vector containing the fitness of each vertex. For directed graphs, this specifies the out-fitness of each vertex.

fitness.in If NULL (the default), the generated graph will be undirected. If not NULL, then it should be a numeric vector and it specifies the in-fitness of each vertex.

If this argument is not NULL, then a directed graph is generated, otherwise an undirected one.

loops Logical scalar, whether to allow loop edges in the graph.

multiple Logical scalar, whether to allow multiple edges in the graph.

Details

This game generates a directed or undirected random graph where the probability of an edge between vertices $i$ and $j$ depends on the fitness scores of the two vertices involved. For undirected graphs, each vertex has a single fitness score. For directed graphs, each vertex has an out- and an in-fitness, and the probability of an edge from $i$ to $j$ depends on the out-fitness of vertex $i$ and the in-fitness of vertex $j$.

The generation process goes as follows. We start from $N$ disconnected nodes (where $N$ is given by the length of the fitness vector). Then we randomly select two vertices $i$ and $j$, with probabilities proportional to their fitnesses. (When the generated graph is directed, $i$ is selected according to the out-fitnesses and $j$ is selected according to the in-fitnesses). If the vertices are not connected yet (or if multiple edges are allowed), we connect them; otherwise we select a new pair. This is repeated until the desired number of links are created.

It can be shown that the expected degree of each vertex will be proportional to its fitness, although the actual, observed degree will not be. If you need to generate a graph with an exact degree sequence, consider sample_degseq instead.

This model is commonly used to generate static scale-free networks. To achieve this, you have to draw the fitness scores from the desired power-law distribution. Alternatively, you may use sample_fitness_pl which generates the fitnesses for you with a given exponent.

Value

An igraph graph, directed or undirected.

Author(s)

Tamas Nepusz <ntamas@gmail.com>

References


Examples

```r
N <- 10000
g <- sample_fitness(5*N, sample((1:50)^-2, N, replace=TRUE))
degree_distribution(g)
```

## Not run:
```r
plot(degree_distribution(g, cumulative=TRUE), log="xy")
```
sample_fitness_pl  Scale-free random graphs, from vertex fitness scores

Description

This function generates a non-growing random graph with expected power-law degree distributions.

Usage

```r
sample_fitness_pl(
  no.of.nodes, 
  no.of.edges, 
  exponent.out, 
  exponent.in = -1, 
  loops = FALSE, 
  multiple = FALSE, 
  finite.size.correction = TRUE
)
```

Arguments

- `no.of.nodes` The number of vertices in the generated graph.
- `no.of.edges` The number of edges in the generated graph.
- `exponent.out` Numeric scalar, the power law exponent of the degree distribution. For directed graphs, this specifies the exponent of the out-degree distribution. It must be greater than or equal to 2. If you pass `Inf` here, you will get back an Erdos-Renyi random network.
- `exponent.in` Numeric scalar. If negative, the generated graph will be undirected. If greater than or equal to 2, this argument specifies the exponent of the in-degree distribution. If non-negative but less than 2, an error will be generated.
- `loops` Logical scalar, whether to allow loop edges in the generated graph.
- `multiple` Logical scalar, whether to allow multiple edges in the generated graph.
- `finite.size.correction` Logical scalar, whether to use the proposed finite size correction of Cho et al., see references below.

Details

This game generates a directed or undirected random graph where the degrees of vertices follow power-law distributions with prescribed exponents. For directed graphs, the exponents of the in- and out-degree distributions may be specified separately.

The game simply uses `sample_fitness` with appropriately constructed fitness vectors. In particular, the fitness of vertex $i$ is $i^{-\alpha}$, where $\alpha = 1/(\gamma - 1)$ and $\gamma$ is the exponent given in the arguments.

To remove correlations between in- and out-degrees in case of directed graphs, the in-fitness vector will be shuffled after it has been set up and before `sample_fitness` is called.

Note that significant finite size effects may be observed for exponents smaller than 3 in the original formulation of the game. This function provides an argument that lets you remove the finite size effects.
effects by assuming that the fitness of vertex \( i \) is \((i + i_0 - 1)^{-\alpha}\) where \( i_0 \) is a constant chosen appropriately to ensure that the maximum degree is less than the square root of the number of edges times the average degree; see the paper of Chung and Lu, and Cho et al for more details.

Value

An igraph graph, directed or undirected.

Author(s)

Tamas Nepusz <ntamas@gmail.com>

References


Examples

```r
g <- sample_fitness_pl(10000, 30000, 2.2, 2.3)
## Not run: plot(degree_distribution(g, cumulative=TRUE, mode="out"), log="xy")
```

---

**sample_forestfire**  
*Forest Fire Network Model*

**Description**

This is a growing network model, which resembles of how the forest fire spreads by igniting trees close by.

**Usage**

```r
sample_forestfire(nodes, fw.prob, bw.factor = 1, ambs = 1, directed = TRUE)
```

**Arguments**

- `nodes`: The number of vertices in the graph.
- `fw.prob`: The forward burning probability, see details below.
- `bw.factor`: The backward burning ratio. The backward burning probability is calculated as `bw.factor*fw.prob`.
- `ambs`: The number of ambassador vertices.
- `directed`: Logical scalar, whether to create a directed graph.
Details

The forest fire model intends to reproduce the following network characteristics, observed in real networks:

- Heavy-tailed in-degree distribution.
- Heavy-tailed out-degree distribution.
- Communities.
- Densification power-law. The network is densifying in time, according to a power-law rule.
- Shrinking diameter. The diameter of the network decreases in time.

The network is generated in the following way. One vertex is added at a time. This vertex connects to (cites) ambs vertices already present in the network, chosen uniformly random. Now, for each cited vertex \( v \) we do the following procedure:

1. We generate two random number, \( x \) and \( y \), that are geometrically distributed with means \( p/(1-p) \) and \( rp(1-rp) \). (\( p \) is \( \text{fw.prob} \), \( r \) is \( \text{bw.factor} \).) The new vertex cites \( x \) outgoing neighbors and \( y \) incoming neighbors of \( v \), from those which are not yet cited by the new vertex. If there are less than \( x \) or \( y \) such vertices available then we cite all of them.

2. The same procedure is applied to all the newly cited vertices.

Value

A simple graph, possibly directed if the `directed` argument is `TRUE`.

Note

The version of the model in the published paper is incorrect in the sense that it cannot generate the kind of graphs the authors claim. A corrected version is available from [http://www.cs.cmu.edu/~jure/pubs/powergrowth-tkdd.pdf](http://www.cs.cmu.edu/~jure/pubs/powergrowth-tkdd.pdf), our implementation is based on this.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

- `barabasi.game` for the basic preferential attachment model.

Examples

```r
g <- sample_forestfire(10000, fw.prob=0.37, bw.factor=0.32/0.37)
dd1 <- degree_distribution(g, mode="in")
dd2 <- degree_distribution(g, mode="out")
plot(seq(along.with=dd1)-1, dd1, log="xy")
points(seq(along.with=dd2)-1, dd2, col=2, pch=2)
```
Generate random graphs according to the G(n,m) Erdos-Renyi model

Description
This model is very simple, every possible edge is created with the same constant probability.

Usage
sample_gnm(n, m, directed = FALSE, loops = FALSE)
gnm(...)

Arguments
n The number of vertices in the graph.
,m The number of edges in the graph.
directed Logical, whether the graph will be directed, defaults to FALSE.
loops Logical, whether to add loop edges, defaults to FALSE.
... Passed to sample_gnm.

Details
The graph has ‘n’ vertices and ‘m’ edges, and the ‘m’ edges are chosen uniformly randomly from the set of all possible edges. This set includes loop edges as well if the loops parameter is TRUE.

Value
A graph object.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

References

See Also
sample_gnp, sample_pa

Examples
```r
  g <- sample_gnm(1000, 1000)
  degree_distribution(g)
```
**sample_gnp**  
Generate random graphs according to the G(n,p) Erdos-Renyi model

**Description**

This model is very simple, every possible edge is created with the same constant probability.

**Usage**

```r
sample_gnp(n, p, directed = FALSE, loops = FALSE)
gnp(...)```

**Arguments**

- `n`  
The number of vertices in the graph.
- `p`  
The probability for drawing an edge between two arbitrary vertices (G(n,p) graph).
- `directed`  
Logical, whether the graph will be directed, defaults to FALSE.
- `loops`  
Logical, whether to add loop edges, defaults to FALSE.
- `...`  
Passed to `sample_gnp`.

**Details**

The graph has ‘n’ vertices and for each edge the probability that it is present in the graph is ‘p’.

**Value**

A graph object.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**References**


**See Also**

`sample_gnm`, `sample_pa`

**Examples**

```r
g <- sample_gnp(1000, 1/1000)
degree_distribution(g)```
Description
Generate a random graph based on the distance of random point on a unit square

Usage
sample_grg(nodes, radius, torus = FALSE, coords = FALSE)
grg(...)  

Arguments
  nodes     The number of vertices in the graph.
  radius    The radius within which the vertices will be connected by an edge.
  torus     Logical constant, whether to use a torus instead of a square.
  coords    Logical scalar, whether to add the positions of the vertices as vertex attributes called ‘x’ and ‘y’.
  ...       Passed to sample_grg.

Details
First a number of points are dropped on a unit square, these points correspond to the vertices of the graph to create. Two points will be connected with an undirected edge if they are closer to each other in Euclidean norm than a given radius. If the torus argument is TRUE then a unit area torus is used instead of a square.

Value
A graph object. If coords is TRUE then with vertex attributes ‘x’ and ‘y’.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>, first version was written by Keith Briggs (http://keithbriggs.info/).

See Also
sample_gnp

Examples
g <- sample_grg(1000, 0.05, torus=FALSE)
g2 <- sample_grg(1000, 0.05, torus=TRUE)
sample_growing

Growing random graph generation

Description
This function creates a random graph by simulating its stochastic evolution.

Usage
sample_growing(n, m = 1, directed = TRUE, citation = FALSE)
growing(...)

Arguments
- n Numeric constant, number of vertices in the graph.
- m Numeric constant, number of edges added in each time step.
- directed Logical, whether to create a directed graph.
- citation Logical. If TRUE a citation graph is created, i.e. in each time step the added edges are originating from the new vertex.
- ... Passed to sample_growing.

Details
This is discrete time step model, in each time step a new vertex is added to the graph and m new edges are created. If citation is FALSE these edges are connecting two uniformly randomly chosen vertices, otherwise the edges are connecting new vertex to uniformly randomly chosen old vertices.

Value
A new graph object.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

See Also
sample_pa, sample_gnp

Examples
```r
g <- sample_growing(500, citation=FALSE)
g2 <- sample_growing(500, citation=TRUE)
```
Sample the hierarchical stochastic block model

Description

Sampling from a hierarchical stochastic block model of networks.

Usage

```r
sample_hierarchical_sbm(n, m, rho, C, p)
```

```r
hierarchical_sbm()
```

Arguments

- `n`: Integer scalar, the number of vertices.
- `m`: Integer scalar, the number of vertices per block. `n / m` must be integer. Alternatively, an integer vector of block sizes, if not all the blocks have equal sizes.
- `rho`: Numeric vector, the fraction of vertices per cluster, within a block. Must sum up to 1, and `rho * m` must be integer for all elements of `rho`. Alternatively a list of `rho` vectors, one for each block, if they are not the same for all blocks.
- `C`: A square, symmetric numeric matrix, the Bernoulli rates for the clusters within a block. Its size must match the size of the `rho` vector. Alternatively, a list of square matrices, if the Bernoulli rates differ in different blocks.
- `p`: Numeric scalar, the Bernoulli rate of connections between vertices in different blocks.
- `...`: Passed to `sample_hierarchical_sbm`.

Details

The function generates a random graph according to the hierarchical stochastic block model.

Value

An igraph graph.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

`sbm.game`
Examples

```r
## Ten blocks with three clusters each
c <- matrix(c(1, 3/4, 0,
            3/4, 0, 3/4,
            0, 3/4, 3/4), nrow=3)
g <- sample_hierarchical_sbm(100, 10, rho=c(3, 3, 4)/10, C=c, p=1/20)
g
if (require(Matrix)) { image(g[]) }
```

Description

`samp;e_hrg` samples a graph from a given hierarchical random graph model.

Usage

`samp;e_hrg(hrg)`

Arguments

- `hrg` A hierarchical random graph model.

Value

An igraph graph.

See Also

Other hierarchical random graph functions: `consensus_tree()`, `fit_hrg()`, `hrg-methods`, `hrg_tree()`, `hrg()`, `predict_edges()`, `print.igraphHRGConsensus()`, `print.igraphHRC`

Sample islands

A graph with subgraphs that are each a random graph.

Description

Create a number of Erdos-Renyi random graphs with identical parameters, and connect them with the specified number of edges.

Usage

`samp;e_islands(islands.n, islands.size, islands.pin, n.inter)`

Arguments

- `islands.n` The number of islands in the graph.
- `islands.size` The size of islands in the graph.
- `islands.pin` The probability to create each possible edge into each island.
- `n.inter` The number of edges to create between two islands.
sample_k_regular

Value
An igraph graph.

Examples

g <- sample_islands(3, 10, 5/10, 1)
oc <- cluster_optimal(g)
oc

Author(s)
Samuel Thiriot

See Also
sample_gnp

sample_k_regular Create a random regular graph

Description
Generate a random graph where each vertex has the same degree.

Usage
sample_k_regular(no.of.nodes, k, directed = FALSE, multiple = FALSE)

Arguments

no.of.nodes  Integer scalar, the number of vertices in the generated graph.
k  Integer scalar, the degree of each vertex in the graph, or the out-degree and in-degree in a directed graph.
directed  Logical scalar, whether to create a directed graph.
multiple  Logical scalar, whether multiple edges are allowed.

Details
This game generates a directed or undirected random graph where the degrees of vertices are equal to a predefined constant k. For undirected graphs, at least one of k and the number of vertices must be even.

The game simply uses sample_degseq with appropriately constructed degree sequences.

Value
An igraph graph.

Author(s)
Tamas Nepusz <ntamas@gmail.com>
See Also

`sample_degseq` for a generator with prescribed degree sequence.

Examples

```r
## A simple ring
ring <- sample_k_regular(10, 2)
plot(ring)

## k-regular graphs on 10 vertices, with k=1:9
k10 <- lapply(1:9, sample_k_regular, no.of.nodes=10)

layout(matrix(1:9, nrow=3, byrow=TRUE))
sapply(k10, plot, vertex.label=NA)
```

---

### Random citation graphs

**Description**

`sample_last_cit` creates a graph, where vertices age, and gain new connections based on how long ago their last citation happened.

**Usage**

```r
sample_last_cit(
  n,
  edges = 1,
  agebins = n/7100,
  pref = (1:(agebins + 1))^-3,
  directed = TRUE
)
```

```r
last_cit(...)
```

```r
sample_cit_types(
  n,
  edges = 1,
  types = rep(0, n),
  pref = rep(1, length(types)),
  directed = TRUE,
  attr = TRUE
)
```

```r
cit_types(...)
```

```r
sample_cit_cit_types(
  n,
  edges = 1,
  types = rep(0, n),
  pref = matrix(1, nrow = length(types), ncol = length(types)),
```
sample_motifs

directed = TRUE,
attr = TRUE
)

cit_cit_types(...)

Arguments

n Number of vertices.
edges Number of edges per step.
agebins Number of aging bins.
pref Vector \( \text{sample\_last\_cit} \) and \( \text{sample\_cit\_types} \) or matrix \( \text{sample\_cit\_cit\_types} \) giving the (unnormalized) citation probabilities for the different vertex types.
directed Logical scalar, whether to generate directed networks.
... Passed to the actual constructor.
types Vector of length 'n', the types of the vertices. Types are numbered from zero.
attr Logical scalar, whether to add the vertex types to the generated graph as a vertex attribute called 'type'.

Details

\( \text{sample\_cit\_cit\_types} \) is a stochastic block model where the graph is growing.
\( \text{sample\_cit\_types} \) is similarly a growing stochastic block model, but the probability of an edge depends on the (potentially) cited vertex only.

Value

A new graph.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Description

Graph motifs are small connected subgraphs with a well-defined structure. These functions search a graph for various motifs.

Usage

sample_motifs(
  graph,
  size = 3,
  cut.prob = rep(0, size),
  sample.size = vcount(graph)/10,
  sample = NULL
)
Arguments

- **graph**: Graph object, the input graph.
- **size**: The size of the motif, currently size 3 and 4 are supported in directed graphs and sizes 3-6 in undirected graphs.
- **cut.prob**: Numeric vector giving the probabilities that the search graph is cut at a certain level. Its length should be the same as the size of the motif (the size argument). By default no cuts are made.
- **sample.size**: The number of vertices to use as a starting point for finding motifs. Only used if the sample argument is NULL.
- **sample**: If not NULL then it specifies the vertices to use as a starting point for finding motifs.

Details

`sample_motifs` estimates the total number of motifs of a given size in a graph based on a sample.

Value

A numeric scalar, an estimate for the total number of motifs in the graph.

See Also

- `isomorphism_class`
- Other graph motifs: `count_motifs()`, `motifs()`

Examples

```r
g <- barabasi.game(100)
motifs(g, 3)
count_motifs(g, 3)
sample_motifs(g, 3)
```

Description

Preferential attachment is a family of simple stochastic algorithms for building a graph. Variants include the Barabási-Abert model and the Price model.

Usage

```r
sample_pa(
  n,
  power = 1,
  m = NULL,
  out.dist = NULL,
  out.seq = NULL,
  out.pref = FALSE,
  zero.appeal = 1,
)```
Arguments

- **n**: Number of vertices.
- **power**: The power of the preferential attachment, the default is one, i.e., linear preferential attachment.
- **m**: Numeric constant, the number of edges to add in each time step. This argument is only used if both `out.dist` and `out.seq` are omitted or NULL.
- **out.dist**: Numeric vector, the distribution of the number of edges to add in each time step. This argument is only used if the `out.seq` argument is omitted or NULL.
- **out.seq**: Numeric vector giving the number of edges to add in each time step. Its first element is ignored as no edges are added in the first time step.
- **out.pref**: Logical, if true the total degree is used for calculating the citation probability, otherwise the in-degree is used.
- **zero.appeal**: The ‘attractiveness’ of the vertices with no adjacent edges. See details below.
- **directed**: Whether to create a directed graph.
- **algorithm**: The algorithm to use for the graph generation. `psumtree` uses a partial prefix-sum tree to generate the graph, this algorithm can handle any power and zero.appeal values and never generates multiple edges. `psumtree-multiple` also uses a partial prefix-sum tree, but the generation of multiple edges is allowed. Before the 0.6 version igraph used this algorithm if `power` was not one, or zero.appeal was not one. `bag` is the algorithm that was previously (before version 0.6) used if `power` was one and zero.appeal was one as well. It works by putting the ids of the vertices into a bag (multiset, really), exactly as many times as their (in-)degree, plus once more. Then the required number of cited vertices are drawn from the bag, with replacement. This method might generate multiple edges. It only works if power and zero.appeal are equal one.
- **start.graph**: NULL or an igraph graph. If a graph, then the supplied graph is used as a starting graph for the preferential attachment algorithm. The graph should have at least one vertex. If a graph is supplied here and the `out.seq` argument is not NULL, then it should contain the out degrees of the new vertices only, not the ones in the `start.graph`.

Details

This is a simple stochastic algorithm to generate a graph. It is a discrete time step model and in each time step a single vertex is added.

We start with a single vertex and no edges in the first time step. Then we add one vertex in each time step and the new vertex initiates some edges to old vertices. The probability that an old vertex is chosen is given by

\[ P[i] \sim k_i^\alpha + a \]
where \(k_i\) is the in-degree of vertex \(i\) in the current time step (more precisely the number of adjacent edges of \(i\) which were not initiated by \(i\) itself) and \(\alpha\) and \(\alpha\) are parameters given by the power and zero argument.

The number of edges initiated in a time step is given by the \(m\), \(out\_dist\) and \(out\_seq\) arguments. If \(out\_seq\) is given and not NULL then it gives the number of edges to add in a vector, the first element is ignored, the second is the number of edges to add in the second time step and so on. If \(out\_seq\) is not given or null and \(out\_dist\) is given and not NULL then it is used as a discrete distribution to generate the number of edges in each time step. Its first element is the probability that no edges will be added, the second is the probability that one edge is added, etc. (\(out\_dist\) does not need to sum up to one, it normalized automatically.) \(out\_dist\) should contain non-negative numbers and at least one element should be positive.

If both \(out\_seq\) and \(out\_dist\) are omitted or NULL then \(m\) will be used, it should be a positive integer constant and \(m\) edges will be added in each time step.

\(sample\_pa\) generates a directed graph by default, set \(directed\) to FALSE to generate an undirected graph. Note that even if an undirected graph is generated \(k_i\) denotes the number of adjacent edges not initiated by the vertex itself and not the total (in- + out-) degree of the vertex, unless the \(out\_pref\) argument is set to TRUE.

**Value**
A graph object.

**Author(s)**
Gabor Csardi <csardi.gabor@gmail.com>

**References**

**See Also**
sample_gnp

**Examples**
```r
  g <- sample_pa(10000)
  degree_distribution(g)
```

---

**sample_pa_age**

Generate an evolving random graph with preferential attachment and aging

**Description**

This function creates a random graph by simulating its evolution. Each time a new vertex is added it creates a number of links to old vertices and the probability that an old vertex is cited depends on its in-degree (preferential attachment) and age.
Usage

```r
sample_pa_age(
  n,
  pa.exp,
  aging.exp,
  m = NULL,
  aging.bin = 300,
  out.dist = NULL,
  out.seq = NULL,
  out.pref = FALSE,
  directed = TRUE,
  zero.deg.appeal = 1,
  zero.age.appeal = 0,
  deg.coef = 1,
  age.coef = 1,
  time.window = NULL
)
```

```r
pa_age(...)```

Arguments

- `n` The number of vertices in the graph.
- `pa.exp` The preferential attachment exponent, see the details below.
- `aging.exp` The exponent of the aging, usually a non-positive number, see details below.
- `m` The number of edges each new vertex creates (except the very first vertex). This argument is used only if both the `out.dist` and `out.seq` arguments are `NULL`.
- `aging.bin` The number of bins to use for measuring the age of vertices, see details below.
- `out.dist` The discrete distribution to generate the number of edges to add in each time step if `out.seq` is `NULL`. See details below.
- `out.seq` The number of edges to add in each time step, a vector containing as many elements as the number of vertices. See details below.
- `out.pref` Logical constant, whether to include edges not initiated by the vertex as a basis of preferential attachment. See details below.
- `directed` Logical constant, whether to generate a directed graph. See details below.
- `zero.deg.appeal` The degree-dependent part of the ‘attractiveness’ of the vertices with no adjacent edges. See also details below.
- `zero.age.appeal` The age-dependent part of the ‘attractiveness’ of the vertices with age zero. It is usually zero, see details below.
- `deg.coef` The coefficient of the degree-dependent ‘attractiveness’. See details below.
- `age.coef` The coefficient of the age-dependent part of the ‘attractiveness’. See details below.
- `time.window` Integer constant, if `NULL` only adjacent added in the last `time.windows` time steps are counted as a basis of the preferential attachment. See also details below.
- `...` Passed to `sample_pa_age`. 
This is a discrete time step model of a growing graph. We start with a network containing a single vertex (and no edges) in the first time step. Then in each time step (starting with the second) a new vertex is added and it initiates a number of edges to the old vertices in the network. The probability that an old vertex is connected to is proportional to

\[ P[i] \sim (c \cdot k_i^\alpha + a)(d \cdot l_i^\beta + b) \]

Here \( k_i \) is the in-degree of vertex \( i \) in the current time step and \( l_i \) is the age of vertex \( i \). The age is simply defined as the number of time steps passed since the vertex is added, with the extension that vertex age is divided to be in aging.bin bins.

c, \( \alpha \), a, d, \( \beta \) and b are parameters and they can be set via the following arguments: pa.exp (\( \alpha \), mandatory argument), aging.exp (\( \beta \), mandatory argument), zero.deg.appeal (a, optional, the default value is 1), zero.age.appeal (b, optional, the default is 0), deg.coef (c, optional, the default is 1), and age.coef (d, optional, the default is 1).

The number of edges initiated in each time step is governed by the m, out.seq and out.pref parameters. If out.seq is given then it is interpreted as a vector giving the number of edges to be added in each time step. It should be of length \( n \) (the number of vertices), and its first element will be ignored. If out.seq is not given (or NULL) and out.dist is given then it will be used as a discrete probability distribution to generate the number of edges. Its first element gives the probability that zero edges are added at a time step, the second element is the probability that one edge is added, etc. (out.seq should contain non-negative numbers, but if they don’t sum up to 1, they will be normalized to sum up to 1. This behavior is similar to the prob argument of the sample command.)

By default a directed graph is generated, but if directed is set to FALSE then an undirected is created. Even if an undirected graph is generated \( k_i \) denotes only the adjacent edges not initiated by the vertex itself except if out.pref is set to TRUE.

If the time.window argument is given (and not NULL) then \( k_i \) means only the adjacent edges added in the previous time.window time steps.

This function might generate graphs with multiple edges.

Value

A new graph.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

sample_pa, sample_gnp

Examples

# The maximum degree for graph with different aging exponents
g1 <- sample_pa_age(10000, pa.exp=1, aging.exp=0, aging.bin=1000)
g2 <- sample_pa_age(10000, pa.exp=1, aging.exp=-1, aging.bin=1000)
g3 <- sample_pa_age(10000, pa.exp=1, aging.exp=-3, aging.bin=1000)
max(degree(g1))
max(degree(g2))
max(degree(g3))
sample_pref

Trait-based random generation

Description

Generation of random graphs based on different vertex types.

Usage

```r
sample_pref(
  nodes,
  types,
  type.dist = rep(1, types),
  fixed.sizes = FALSE,
  pref.matrix = matrix(1, types, types),
  directed = FALSE,
  loops = FALSE
)
```

```r
pref(...)  
sample_asym_pref(
  nodes,
  types,
  type.dist.matrix = matrix(1, types, types),
  pref.matrix = matrix(1, types, types),
  loops = FALSE
)
```

```r
asym_pref(...)  
```

Arguments

- **nodes**: The number of vertices in the graphs.
- **types**: The number of different vertex types.
- **type.dist**: The distribution of the vertex types, a numeric vector of length ‘types’ containing non-negative numbers. The vector will be normed to obtain probabilities.
- **fixed.sizes**: Fix the number of vertices with a given vertex type label. The type.dist argument gives the group sizes (i.e. number of vertices with the different labels) in this case.
- **pref.matrix**: A square matrix giving the preferences of the vertex types. The matrix has ‘types’ rows and columns.
- **directed**: Logical constant, whether to create a directed graph.
- **loops**: Logical constant, whether self-loops are allowed in the graph.
- **...**: Passed to the constructor, sample_pref or sample_asym_pref.
- **type.dist.matrix**: The joint distribution of the in- and out-vertex types.

pref(...)

sample_asym pref(...)

asym pref(...)

Details

Both models generate random graphs with given vertex types. For `sample_pref` the probability that two vertices will be connected depends on their type and is given by the `pref.matrix` argument. This matrix should be symmetric to make sense but this is not checked. The distribution of the different vertex types is given by the `type.dist` vector.

For `sample_asym_pref` each vertex has an in-type and an out-type and a directed graph is created. The probability that a directed edge is realized from a vertex with a given out-type to a vertex with a given in-type is given in the `pref.matrix` argument, which can be asymmetric. The joint distribution for the in- and out-types is given in the `type.dist.matrix` argument.

The types of the generated vertices can be retrieved from the `type` vertex attribute for `sample_pref` and from the `intype` and `outtype` vertex attribute for `sample_asym_pref`.

Value

An igraph graph.

Author(s)

Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com> for the R interface

See Also

`sample_traits`, `sample_traits_callaway`

Examples

```r
pf <- matrix(c(1, 0, 0, 1), nrow=2)
g <- sample_pref(20, 2, pref.matrix=pf)
## Not run: tkplot(g, layout=layout_with_fr)

pf <- matrix(c(0, 1, 0, 0), nrow=2)
g <- sample_asym_pref(20, 2, pref.matrix=pf)
## Not run: tkplot(g, layout=layout_in_circle)
```

Description

Sampling from the stochastic block model of networks

Usage

```r
sample_sbm(n, pref.matrix, block.sizes, directed = FALSE, loops = FALSE)
```

```r
sbm(...)```
Arguments

- **n**: Number of vertices in the graph.
- **pref.matrix**: The matrix giving the Bernoulli rates. This is a \( K \times K \) matrix, where \( K \) is the number of groups. The probability of creating an edge between vertices from groups \( i \) and \( j \) is given by element \((i, j)\). For undirected graphs, this matrix must be symmetric.
- **block.sizes**: Numeric vector giving the number of vertices in each group. The sum of the vector must match the number of vertices.
- **directed**: Logical scalar, whether to generate a directed graph.
- **loops**: Logical scalar, whether self-loops are allowed in the graph.
- **...**: Passed to `sample_sbm`.

Details

This function samples graphs from a stochastic block model by (doing the equivalent of) Bernoulli trials for each potential edge with the probabilities given by the Bernoulli rate matrix, `pref.matrix`. The order of the vertices in the generated graph corresponds to the `block.sizes` argument.

Value

An igraph graph.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

`sample_gnp`, `sample_gnm`

Examples

```r
## Two groups with not only few connection between groups
pm <- cbind( c(.1, .001), c(.001, .05) )
g <- sample_sbm(1000, pref.matrix=pm, block.sizes=c(300,700))
g
```
Sampling a random integer sequence

Description

This function provides a very efficient way to pull an integer random sample sequence from an integer interval.

Usage

```r
sample_seq(low, high, length)
```

Arguments

- `low` The lower limit of the interval (inclusive).
- `high` The higher limit of the interval (inclusive).
- `length` The length of the sample.

Details

The algorithm runs in $O(\text{length})$ expected time, even if `high-low` is big. It is much faster (but of course less general) than the builtin `sample` function of R.

Value

An increasing numeric vector containing integers, the sample.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


Examples

```r
rs <- sample_seq(1, 100000000, 10)
rs
```
The Watts-Strogatz small-world model

Description
Generate a graph according to the Watts-Strogatz network model.

Usage
sample_smallworld(dim, size, nei, p, loops = FALSE, multiple = FALSE)
smallworld(...)

Arguments
dim  Integer constant, the dimension of the starting lattice.
size Integer constant, the size of the lattice along each dimension.
nei  Integer constant, the neighborhood within which the vertices of the lattice will
     be connected.
p    Real constant between zero and one, the rewiring probability.
loops Logical scalar, whether loops edges are allowed in the generated graph.
multiple Logical scalar, whether multiple edges are allowed int the generated graph.
...   Passed to sample_smallworld.

Details
First a lattice is created with the given dim, size and nei arguments. Then the edges of the lattice
are rewired uniformly randomly with probability p.

Note that this function might create graphs with loops and/or multiple edges. You can use simplify
to get rid of these.

Value
A graph object.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

References
Duncan J Watts and Steven H Strogatz: Collective dynamics of ‘small world’ networks, Nature 393,

See Also
make_lattice, rewire
Examples

```r
g <- sample_smallworld(1, 100, 5, 0.05)
mean_distance(g)
transitivity(g, type="average")
```

---

**sample_spanning_tree**  
*Samples from the spanning trees of a graph randomly and uniformly*

**Description**

`sample_spanning_tree` picks a spanning tree of an undirected graph randomly and uniformly, using loop-erased random walks.

**Usage**

```r
sample_spanning_tree(graph, vid = 0)
```

**Arguments**

- `graph`  
The input graph to sample from. Edge directions are ignored if the graph is directed.
- `vid`  
When the graph is disconnected, this argument specifies how to handle the situation. When the argument is zero (the default), the sampling will be performed component-wise, and the result will be a spanning forest. When the argument contains a vertex ID, only the component containing the given vertex will be processed, and the result will be a spanning tree of the component of the graph.

**Value**

An edge sequence containing the edges of the spanning tree. Use `subgraph.edges` to extract the corresponding subgraph.

**See Also**

`subgraph.edges` to extract the tree itself

**Examples**

```r
g <- make_full_graph(10) %du% make_full_graph(5)
edges <- sample_spanning_tree(g)
forest <- subgraph.edges(g, edges)
```
sample_sphere_surface  Sample vectors uniformly from the surface of a sphere

Description

Sample finite-dimensional vectors to use as latent position vectors in random dot product graphs.

Usage

sample_sphere_surface(dim, n = 1, radius = 1, positive = TRUE)

Arguments

dim  Integer scalar, the dimension of the random vectors.
n  Integer scalar, the sample size.
radius  Numeric scalar, the radius of the sphere to sample.
positive  Logical scalar, whether to sample from the positive orthant of the sphere.

Details

sample_sphere_surface generates uniform samples from $S^{dim-1}$ (the (dim-1)-sphere) with radius radius, i.e. the Euclidean norm of the samples equal radius.

Value

A dim (length of the alpha vector for sample_dirichlet) times n matrix, whose columns are the sample vectors.

See Also

Other latent position vector samplers: sample_dirichlet(), sample_sphere_volume()

Examples

lpvs.sph <- sample_sphere_surface(dim=10, n=20, radius=1)
RDP.graph.3 <- sample_dot_product(lpvs.sph)
vec.norm <- apply(lpvs.sph, 2, function(x) { sum(x^2) })
vec.norm
Sample vectors uniformly from the volume of a sphere

Description

Sample finite-dimensional vectors to use as latent position vectors in random dot product graphs.

Usage

```r
sample_sphere_volume(dim, n = 1, radius = 1, positive = TRUE)
```

Arguments

- `dim`: Integer scalar, the dimension of the random vectors.
- `n`: Integer scalar, the sample size.
- `radius`: Numeric scalar, the radius of the sphere to sample.
- `positive`: Logical scalar, whether to sample from the positive orthant of the sphere.

Details

`sample_sphere_volume` generates uniform samples from $S^{dim-1}$ (the (dim-1)-sphere) i.e. the Euclidean norm of the samples is smaller or equal to `radius`.

Value

A `dim` (length of the alpha vector for `sample_dirichlet`) times `n` matrix, whose columns are the sample vectors.

See Also

Other latent position vector samplers: `sample_dirichlet()`, `sample_sphere_surface()`

Examples

```r
lpvs.sph.vol <- sample_sphere_volume(dim=10, n=20, radius=1)
RDP.graph.4 <- sample_dot_product(lpvs.sph.vol)
vec.norm <- apply(lpvs.sph.vol, 2, function(x) { sum(x^2) })
vec.norm
```
**sample_traits_callaway**

*Graph generation based on different vertex types*

**Description**

These functions implement evolving network models based on different vertex types.

**Usage**

```r
sample_traits_callaway(
  nodes,              
  types,              
  edge.per.step = 1,  
  type.dist = rep(1, types),
  pref.matrix = matrix(1, types, types),
  directed = FALSE
)
```

```r
traits_callaway(...)
```

```r
sample_traits(
  nodes,              
  types,              
  k = 1,              
  type.dist = rep(1, types),
  pref.matrix = matrix(1, types, types),
  directed = FALSE
)
```

```r
traits(...)
```

**Arguments**

- **nodes**: The number of vertices in the graph.
- **types**: The number of different vertex types.
- **edge.per.step**: The number of edges to add to the graph per time step.
- **type.dist**: The distribution of the vertex types. This is assumed to be stationary in time.
- **pref.matrix**: A matrix giving the preferences of the given vertex types. These should be probabilities, i.e. numbers between zero and one.
- **directed**: Logical constant, whether to generate directed graphs.
- **k**: Passed to the constructor, `sample_traits` or `sample_traits_callaway`.
- **...**: The number of trials per time step, see details below.

**Details**

For `sample_traits_callaway` the simulation goes like this: in each discrete time step a new vertex is added to the graph. The type of this vertex is generated based on `type.dist`. Then two vertices
are selected uniformly randomly from the graph. The probability that they will be connected de-
pends on the types of these vertices and is taken from `pref.matrix`. Then another two vertices are
selected and this is repeated `edges.per.step` times in each time step.

For `sample_traits` the simulation goes like this: a single vertex is added at each time step. This
new vertex tries to connect to `k` vertices in the graph. The probability that such a connection is
realized depends on the types of the vertices involved and is taken from `pref.matrix`.

**Value**

A new graph object.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**Examples**

```r
# two types of vertices, they like only themselves
g1 <- sample_traits_callaway(1000, 2, pref.matrix=matrix(c(1,0,0,1), ncol=2))
g2 <- sample_traits(1000, 2, k=2, pref.matrix=matrix(c(1,0,0,1), ncol=2))
```

**Description**

`sample_tree` generates a random with a given number of nodes uniform at random from the set of
labelled trees.

**Usage**

```r
sample_tree(n, directed = FALSE, method = c("lerw", "prufer"))
```

**Arguments**

- `n` The number of nodes in the tree
- `directed` Whether to create a directed tree. The edges of the tree are oriented away from
  the root.
- `method` The algorithm to use to generate the tree. ‘prufer’ samples Prufer sequences
  uniformly and then converts the sampled sequence to a tree. ‘lerw’ performs a
  loop-erased random walk on the complete graph to uniformly sample its span-
  ning trees. (This is also known as Wilson’s algorithm). The default is ‘lerw’.
  Note that the method based on Prufer sequences does not support directed trees
  at the moment.

**Details**

In other words, the function generates each possible labelled tree with the given number of nodes
with the same probability.
Value

A graph object.

Examples

g <- sample_tree(100, method="lerw")

Description

Calculate scan statistics on a time series of graphs. This is done by calculating the local scan statistics for each graph and each vertex, and then normalizing across the vertices and across the time steps.

Usage

scan_stat(graphs, tau = 1, ell = 0, locality = c("us", "them"), ...)

Arguments

graphs A list of igraph graph objects. They must be all directed or all undirected and they must have the same number of vertices.

tau The number of previous time steps to consider for the time-dependent normalization for individual vertices. In other words, the current locality statistics of each vertex will be compared to this many previous time steps of the same vertex to decide whether it is significantly larger.

ell The number of previous time steps to consider for the aggregated scan statistics. This is essentially a smoothing parameter.

locality Whether to calculate the ‘us’ or ‘them’ statistics.

... Extra arguments are passed to local_scan.

Value

A list with entries:

stat The scan statistics in each time step. It is NA for the initial \(\tau + \ell\) time steps.

arg_max_v The (numeric) vertex ids for the vertex with the largest locality statistics, at each time step. It is NA for the initial \(\tau + \ell\) time steps.

See Also

Other scan statistics: local_scan()
Examples

```r
## Generate a bunch of SBMs, with the last one being different
num_t <- 20
block_sizes <- c(10, 5, 5)
p_ij <- list(p = 0.1, h = 0.9, q = 0.9)
P0 <- matrix(p_ij$p, 3, 3)
P0[2, 2] <- p_ij$h
PA <- P0
PA[3, 3] <- p_ij$q
num_v <- sum(block_sizes)

tsg <- replicate(num_t - 1, P0, simplify = FALSE) %>%
  append(list(PA)) %>%
  lapply(sample_sbm, n = num_v, block.sizes = block_sizes, directed = TRUE)

scan_stat(graphs = tsg, k = 1, tau = 4, ell = 2)
scan_stat(graphs = tsg, locality = "them", k = 1, tau = 4, ell = 2)
```

---

**scg**

*All-in-one Function for the SCG of Matrices and Graphs*

**Description**

This function handles all the steps involved in the Spectral Coarse Graining (SCG) of some matrices and graphs as described in the reference below.

**Usage**

```r
scg(
X,
ev,
nt,
groups = NULL,
mtype = c("symmetric", "laplacian", "stochastic"),
algo = c("optimum", "interv_km", "interv", "exact_scg"),
norm = c("row", "col"),
direction = c("default", "left", "right"),
evec = NULL,
p = NULL,
use.arpack = FALSE,
maxiter = 300,
sparse = igraph_opt("sparsematrices"),
output = c("default", "matrix", "graph"),
semproj = FALSE,
epairs = FALSE,
stat.prob = FALSE
)
```
Arguments

\(X\)  
The input graph or square matrix. Can be of class igraph, matrix or Matrix.

\(ev\)  
A vector of positive integers giving the indexes of the eigenpairs to be preserved. For real eigenpairs, 1 designates the eigenvalue with largest algebraic value, 2 the one with second largest algebraic value, etc. In the complex case, it is the magnitude that matters.

\(nt\)  
A vector of positive integers of length one or equal to length\((ev)\). When \(algo\) = “optimum”, \(nt\) contains the number of groups used to partition each eigenvector separately. When \(algo\) is equal to “interv_km” or “interv”, \(nt\) contains the number of intervals used to partition each eigenvector. The same partition size or number of intervals is used for each eigenvector if \(nt\) is a single integer. When \(algo\) = “exact_cg” this parameter is ignored.

\(groups\)  
A vector of nrow\((X)\) or vcount\((X)\) integers labeling each group vertex in the partition. If this parameter is supplied most part of the function is bypassed.

\(mtype\)  
Character scalar. The type of semi-projector to be used for the SCG. For now “symmetric”, “laplacian” and “stochastic” are available.

\(algo\)  
Character scalar. The algorithm used to solve the SCG problem. Possible values are “optimum”, “interv_km”, “interv” and “exact_cg”.

\(norm\)  
Character scalar. Either “row” or “col”. If set to “row” the rows of the Laplacian matrix sum up to zero and the rows of the stochastic matrix sum up to one; otherwise it is the columns.

\(direction\)  
Character scalar. When set to “right”, resp. “left”, the parameters \(ev\) and \(evec\) refer to right, resp. left eigenvectors. When passed “default” it is the SCG described in the reference below that is applied (common usage). This argument is currently not implemented, and right eigenvectors are always used.

\(evec\)  
A numeric matrix of (eigen)vectors to be preserved by the coarse graining (the vectors are to be stored column-wise in \(evec\)). If supplied, the eigenvectors should correspond to the indexes in \(ev\) as no cross-check will be done.

\(p\)  
A probability vector of length nrow\((X)\) (or vcount\((X)\)). \(p\) is the stationary probability distribution of a Markov chain when \(mtype\) = “stochastic”. This parameter is ignored in all other cases.

\(use.arpack\)  
Logical scalar. When set to TRUE uses the function arpack to compute eigenpairs. This parameter should be set to TRUE if one deals with large (over a few thousands) AND sparse graphs or matrices. This argument is not implemented currently and LAPACK is used for solving the eigenproblems.

\(maxiter\)  
A positive integer giving the maximum number of iterations for the k-means algorithm when \(algo\) = “interv_km”. This parameter is ignored in all other cases.

\(sparse\)  
Logical scalar. Whether to return sparse matrices in the result, if matrices are requested.

\(output\)  
Character scalar. Set this parameter to “default” to retrieve a coarse-grained object of the same class as \(X\).

\(semproj\)  
Logical scalar. Set this parameter to TRUE to retrieve the semi-projectors of the SCG.

\(epairs\)  
Logical scalar. Set this to TRUE to collect the eigenpairs computed by scg.

\(stat.prob\)  
Logical scalar. This is to collect the stationary probability \(p\) when dealing with stochastic matrices.
Details

Please see `scg-method` for an introduction.

In the following, $V$ is the matrix of eigenvectors for which the SCG is solved. $V$ is calculated from $X$, if it is not given in the `evec` argument.

The algorithm “optimum” solves exactly the SCG problem for each eigenvector in $V$. The running time of this algorithm is $O(\max nt \cdot m^2)$ for the symmetric and laplacian matrix problems (i.e. when `mtype` is “symmetric” or “laplacian”). It is $O(m^3)$ for the stochastic problem. Here $m$ is the number of rows in $V$. In all three cases, the memory usage is $O(m^2)$.

The algorithms “interv” and “interv_km” solve approximately the SCG problem by performing a (for now) constant binning of the components of the eigenvectors, that is $nt[i]$ constant-size bins are used to partition $V[,i]$. When `algo` = “interv_km”, the (Lloyd) k-means algorithm is run on each partition obtained by “interv” to improve accuracy.

Once a minimizing partition (either exact or approximate) has been found for each eigenvector, the final grouping is worked out as follows: two vertices are grouped together in the final partition if they are grouped together in each minimizing partition. In general the size of the final partition is not known in advance when `ncol(V)>1`.

Finally, the algorithm “exact_scg” groups the vertices with equal components in each eigenvector. The last three algorithms essentially have linear running time and memory load.

Value

- `Xt` The coarse-grained graph, or matrix, possibly a sparse matrix.
- `groups` A vector of `nrow(X)` or `vcount(X)` integers giving the group label of each object (vertex) in the partition.
- `L` The semi-projector $L$ if `semproj = TRUE`.
- `R` The semi-projector $R$ if `semproj = TRUE`.
- `values` The computed eigenvalues if `epairs = TRUE`.
- `vectors` The computed or supplied eigenvectors if `epairs = TRUE`.
- `p` The stationary probability vector if `mtype = stochastic` and `stat.prob = TRUE`. For other matrix types this is missing.

Author(s)


References


See Also

`scg-method` for an introduction. `scg_eps`, `scg_group` and `scg_semi_proj`. 
Examples

## We are not running these examples any more, because they
## take a long time (~20 seconds) to run and this is against the CRAN
## repository policy. Copy and paste them by hand to your R prompt if
## you want to run them.

## Not run:
# SCG of a toy network
g <- make_full_graph(5) %du% make_full_graph(5) %du% make_full_graph(5)
g <- add_edges(g, c(1, 6, 1, 11, 6, 11))
cg <- scg(g, 1, 3, algo="exact_scg")

# plot the result
layout <- layout_with_kk(g)
nt <- vcount(cg$Xt)
col <- rainbow(nt)
vsize <- table(cg$groups)
ewidth <- round(E(cg$Xt)$weight, 2)

op <- par(mfrow=c(1,2))
plot(g, vertex.color = col[cg$groups], vertex.size = 20,
     vertex.label = NA, layout = layout)
plot(cg$Xt, edge.width = ewidth, edge.label = ewidth,
     vertex.color = col, vertex.size = 20*vsize/max(vsize),
     vertex.label=NA, layout = layout_with_kk)
par(op)

## SCG of real-world network
library(igraphdata)
data(immuno)
summary(immuno)
n <- vcount(immuno)
interv <- c(100, 100, 50, 25, 12, 6, 3, 2, 2)
cg <- scg(immuno, ev= n-(1:9), nt=interv, mtype="laplacian",
           algo="interv", epairs=TRUE)

## are the eigenvalues well-preserved?
gt <- cg$Xt
nt <- vcount(gt)
Lt <- laplacian_matrix(gt)
evalt <- eigen(Lt, only.values=TRUE)$values[nt-(1:9)]
res <- cbind(interv, cg$values, evalt)
res <- round(res, 5)
colnames(res) <- c("interv","lambda_i","lambda_tilde_i")
rownames(res) <- c("N-1","N-2","N-3","N-4","N-5","N-6","N-7","N-8","N-9")
print(res)

## use SCG to get the communities
com <- scg(laplacian_matrix(immuno), ev=n-c(1,2), nt=2)$groups
col <- rainbow(max(com))
layout <- layout_nicely(immuno)

plot(immuno, layout=layout, vertex.size=3, vertex.color=col[com],
     vertex.label=NA)
## display the coarse-grained graph

gt <- simplify(as.undirected(gt))
layout.cg <- layout_with_kk(gt)
com.cg <- scg(laplacian_matrix(gt), nt=c(1,2), 2)$groups
vsize <- sqrt(as.vector(table(cg$groups)))

op <- par(mfrow=c(1,2))
plot(immuno, layout=layout, vertex.size=3, vertex.color=col[com],
    vertex.label=NA)
plot(gt, layout=layout.cg, vertex.size=15*vsize/max(vsize),
    vertex.color=col[com.cg],vertex.label=NA)
par(op)

## End(Not run)

### scg-method

#### Spectral Coarse Graining

**Description**

Functions to perform the Spectral Coarse Graining (SCG) of matrices and graphs.

**Introduction**

The SCG functions provide a framework, called Spectral Coarse Graining (SCG), for reducing large graphs while preserving their spectral-related features, that is features closely related with the eigenvalues and eigenvectors of a graph matrix (which for now can be the adjacency, the stochastic, or the Laplacian matrix).

Common examples of such features comprise the first-passage-time of random walkers on Markovian graphs, thermodynamic properties of lattice models in statistical physics (e.g. Ising model), and the epidemic threshold of epidemic network models (SIR and SIS models).

SCG differs from traditional clustering schemes by producing a coarse-grained graph (not just a partition of the vertices), representative of the original one. As shown in [1], Principal Component Analysis can be viewed as a particular SCG, called exact SCG, where the matrix to be coarse-grained is the covariance matrix of some data set.

SCG should be of interest to practitioners of various fields dealing with problems where matrix eigenpairs play an important role, as for instance is the case of dynamical processes on networks.

**Author(s)**


**References**


---

**scg_eps**

*Error of the spectral coarse graining (SCG) approximation*

**Description**

`scg_eps` computes $\|v_i - P v_i\|$, where $v_i$ is the $i$th eigenvector in $V$ and $P$ is the projector corresponding to the `mtype` argument.

**Usage**

```r
scg_eps(
  V, 
  groups, 
  mtype = c("symmetric", "laplacian", "stochastic"), 
  p = NULL, 
  norm = c("row", "col")
)
```

**Arguments**

- `V` A numeric matrix of (eigen)vectors assumed normalized. The vectors are to be stored column-wise in $V$.
- `groups` A vector of `nrow(V)` integers labeling each group vertex in the partition.
- `mtype` The type of semi-projector used for the SCG. For now “symmetric”, “laplacian” and “stochastic” are available.
- `p` A probability vector of length `nrow(V)`. $p$ is the stationary probability distribution of a Markov chain when `mtype` = “stochastic”. This parameter is ignored otherwise.
- `norm` Either “row” or “col”. If set to “row” the rows of the Laplacian matrix sum to zero and the rows of the stochastic matrix sum to one; otherwise it is the columns.

**Value**

`scg_eps` returns with a numeric vector whose $i$th component is $\|v_i - P v_i\|$ (see Details).

**Author(s)**


**References**


**See Also**

`scg-method` and `scg`. 
Examples

```r
t <- rexp(20)
k <- kmeans(t, 5)
sum(k$withinss)
scg_eps(cbind(t), km$cluster)^2
```

---

### Description

This function solves the Spectral Coarse Graining (SCG) problem; either exactly, or approximately but faster.

### Usage

```r
scg_group(
  V,
  nt,
  mtype = c("symmetric", "laplacian", "stochastic"),
  algo = c("optimum", "interv_km", "interv", "exact_scg"),
  p = NULL,
  maxiter = 100
)
```

### Arguments

- **V**: A numeric matrix of (eigen)vectors to be preserved by the coarse graining (the vectors are to be stored column-wise in V).
- **nt**: A vector of positive integers of length one or equal to length(ev). When algo = "optimum", nt contains the number of groups used to partition each eigenvector separately. When algo is equal to "interv_km" or "interv", nt contains the number of intervals used to partition each eigenvector. The same partition size or number of intervals is used for each eigenvector if nt is a single integer. When algo = "exact_cg" this parameter is ignored.
- **mtype**: The type of semi-projectors used in the SCG. For now "symmetric", "laplacian" and "stochastic" are available.
- **algo**: The algorithm used to solve the SCG problem. Possible values are "optimum", "interv_km", "interv" and "exact_scg".
- **p**: A probability vector of length equal to nrow(V). p is the stationary probability distribution of a Markov chain when mtype = "stochastic". This parameter is ignored in all other cases.
- **maxiter**: A positive integer giving the maximum number of iterations of the k-means algorithm when algo = "interv_km". This parameter is ignored in all other cases.
Details

The algorithm “optimum” solves exactly the SCG problem for each eigenvector in $V$. The running time of this algorithm is $O(\text{max } nt \cdot m^2)$ for the symmetric and laplacian matrix problems (i.e. when $\text{mtype}$ is “symmetric” or “laplacian”. It is $O(m^3)$ for the stochastic problem. Here $m$ is the number of rows in $V$. In all three cases, the memory usage is $O(m^2)$.

The algorithms “interv” and “interv_km” solve approximately the SCG problem by performing a (for now) constant binning of the components of the eigenvectors, that is $nt[i]$ constant-size bins are used to partition $V[,i]$. When $\text{algo} = \text{“interv_km”}$, the (Lloyd) k-means algorithm is run on each partition obtained by “interv” to improve accuracy.

Once a minimizing partition (either exact or approximate) has been found for each eigenvector, the final grouping is worked out as follows: two vertices are grouped together in the final partition if they are grouped together in each minimizing partition. In general the size of the final partition is not known in advance when $\text{ncol}(V)>1$.

Finally, the algorithm “exact_scg” groups the vertices with equal components in each eigenvector. The last three algorithms essentially have linear running time and memory load.

Value

A vector of $\text{nrow}(V)$ integers giving the group label of each object (vertex) in the partition.

Author(s)

David Morton de Lachapelle <david.morton@epfl.ch>, <david.mortondelachapelle@swissquote.ch>

References


See Also

scg-method for a detailed introduction. scg, scg_eps

Examples

## We are not running these examples any more, because they
## take a long time to run and this is against the CRAN repository
## policy. Copy and paste them by hand to your R prompt if
## you want to run them.

## Not run:
# eigenvectors of a random symmetric matrix
M <- matrix(rexp(10^6), 10^3, 10^3)
M <- (M + t(M))/2
V <- eigen(M, symmetric=TRUE)$vectors[,c(1,2)]

# displays size of the groups in the final partition
gr <- scg_group(V, nt=c(2,3))
col <- rainbow(max(gr))
plot(table(gr), col=col, main="Group size", xlab="group", ylab="size")

## comparison with the grouping obtained by kmeans
## Semi-Projectors

**Description**

A function to compute the $L$ and $R$ semi-projectors for a given partition of the vertices.

**Usage**

```r
scg_semi_proj(
  groups,
  mtype = c("symmetric", "laplacian", "stochastic"),
  p = NULL,
  norm = c("row", "col"),
  sparse = igraph_opt("sparsematrices")
)
```

**Arguments**

- `groups` A vector of `nrow(X)` or `vcount(X)` integers giving the group label of every vertex in the partition.
- `mtype` The type of semi-projectors. For now “symmetric”, “laplacian” and “stochastic” are available.
- `p` A probability vector of length `length(gr)`. $p$ is the stationary probability distribution of a Markov chain when `mtype = "stochastic"`. This parameter is ignored in all other cases.
**norm** Either “row” or “col”. If set to “row” the rows of the Laplacian matrix sum up to zero and the rows of the stochastic sum up to one; otherwise it is the columns.

**sparse** Logical scalar, whether to return sparse matrices.

**Details**

The three types of semi-projectors are defined as follows. Let $\gamma(j)$ label the group of vertex $j$ in a partition of all the vertices.

The symmetric semi-projectors are defined as

$$ L_{\alpha j} = R_{\alpha j} = \frac{1}{\sqrt{\vert \alpha \vert}} \delta_{\alpha \gamma(j)}, $$

the (row) Laplacian semi-projectors as

$$ L_{\alpha j} = \frac{1}{\vert \alpha \vert} \delta_{\alpha \gamma(j)} $$

and $R_{\alpha j} = \delta_{\alpha \gamma(j)}$,

and the (row) stochastic semi-projectors as

$$ L_{\alpha j} = \frac{p_1(j)}{\sum_{k \in \gamma(j)} p_1(k)} $$

and $R_{\alpha j} = \delta_{\alpha \gamma(j)} \delta_{\alpha \gamma(j)}$,

where $p_1$ is the (left) eigenvector associated with the one-eigenvalue of the stochastic matrix. $L$ and $R$ are defined in a symmetric way when norm = col. All these semi-projectors verify various properties described in the reference.

**Value**

- **L** The semi-projector $L$.
- **R** The semi-projector $R$.

**Author(s)**


**References**


**See Also**

[scg-method](#) for a detailed introduction. [scg, scg_eps, scg_group](#)
Examples

```r
library(Matrix)
# compute the semi-projectors and projector for the partition
# provided by a community detection method
g <- sample_pa(20, m = 1.5, directed = FALSE)
eb <- cluster_edge_betweenness(g)
memb <- membership(eb)
lr <- scg_semi_proj(memb)
# In the symmetric case L = R
tcrossprod(lr$R) # same as lr$R %*% t(lr$R)
P <- crossprod(lr$R) # same as t(lr$R) %*% lr$R
# P is an orthogonal projector
isSymmetric(P)
sum((P %*% P-P)^2)

## use L and R to coarse-grain the graph Laplacian
lr <- scg_semi_proj(memb, mtype="laplacian")
L <- laplacian_matrix(g)
Lt <- lr$L %*% L %*% t(lr$R)
## or better lr$L %*% tcrossprod(L,lr$R)
rowSums(Lt)
```

sequential_pal  
Sequential palette

Description

This is the ‘OrRd’ palette from https://colorbrewer2.org/. It has at most nine colors.

Usage

```r
sequential_pal(n)
```

Arguments

- `n` The number of colors in the palette. The maximum is nine currently.

Details

Use this palette, if vertex colors mark some ordinal quantity, e.g. some centrality measure, or some ordinal vertex covariate, like the age of people, or their seniority level.

Value

A character vector of RGB color codes.

See Also

Other palettes: categorical_pal(), diverging_pal(), r_pal()
Examples

```r
## Not run:
library(igraphdata)
data(karate)
karate <- karate %>%
  add_layout_(with_kk()) %>%
  set_vertex_attr("size", value = 10)
V(karate)$color <- scales::dscale(degree(karate) %>% cut(5), sequential_pal)
plot(karate)
## End(Not run)
```

---

**set_edge_attr**

Set edge attributes

**Description**

Set edge attributes

**Usage**

```
set_edge_attr(graph, name, index = E(graph), value)
```

**Arguments**

- `graph`
  - The graph
- `name`
  - The name of the attribute to set.
- `index`
  - An optional edge sequence to set the attributes of a subset of edges.
- `value`
  - The new value of the attribute for all (or index) edges.

**Value**

The graph, with the edge attribute added or set.

**See Also**

Other graph attributes: `delete_edge_attr()`, `delete_graph_attr()`, `delete_vertex_attr()`, `edge_attr<-()`, `edge_attr_names()`, `edge_attr()`, `graph_attr<-()`, `graph_attr_names()`, `graph_attr()`, `igraph-dollar`, `igraph-vs-attributes`, `set_graph_attr()`, `set_vertex_attr()`, `vertex_attr<-()`, `vertex_attr_names()`, `vertex_attr()`

**Examples**

```r
g <- make_ring(10) %>%
  set_edge_attr("label", value = LETTERS[1:10])
g
plot(g)
```
set_graph_attr  Set a graph attribute

Description
An existing attribute with the same name is overwritten.

Usage
set_graph_attr(graph, name, value)

Arguments
graph  The graph.
name  The name of the attribute to set.
value  New value of the attribute.

Value
The graph with the new graph attribute added or set.

See Also
Other graph attributes: delete_edge_attr(), delete_graph_attr(),
delete_vertex_attr(), edge_attr<-(), edge_attr_names(), edge_attr(),
graph_attr<-(), graph_attr_names(),
graph_attr(), igraph-dollar, igraph-vs-attributes, set_edge_attr(),
set_vertex_attr(), vertex_attr<-(), vertex_attr_names(), vertex_attr()

Examples
```r
g <- make_ring(10) %>%
  set_graph_attr("layout", layout_with_fr)
g
plot(g)
```

set_vertex_attr  Set vertex attributes

Description
Set vertex attributes

Usage
set_vertex_attr(graph, name, index = V(graph), value)
**Arguments**

- **graph**
  - The graph.

- **name**
  - The name of the attribute to set.

- **index**
  - An optional vertex sequence to set the attributes of a subset of vertices.

- **value**
  - The new value of the attribute for all (or index) vertices.

**Value**

- The graph, with the vertex attribute added or set.

**See Also**

- Other graph attributes: `delete_edge_attr()`, `delete_graph_attr()`, `delete_vertex_attr()`, `edge_attr<-()`, `edge_attr_names()`, `edge_attr()`, `graph_attr<-()`, `graph_attr_names()`, `graph_attr()`, `igraph-dollar`, `igraph-vs-attributes`, `set_edge_attr()`, `set_graph_attr()`, `vertex_attr<-()`, `vertex_attr_names()`, `vertex_attr()`

**Examples**

```r
  g <- make_ring(10) %>%
    set_vertex_attr("label", value = LETTERS[1:10])
  g
  plot(g)
```

**shapes**

Various vertex shapes when plotting igraph graphs

**Description**

Starting from version 0.5.1 igraph supports different vertex shapes when plotting graphs.

**Usage**

- `shapes(shape = NULL)`
- `shape_noclip(coords, el, params, end = c("both", "from", "to"))`
- `shape_noplot(coords, v = NULL, params)`
- `add_shape(shape, clip = shape_noclip, plot = shape_noplot, parameters = list())`

**Arguments**

- **shape**
  - Character scalar, name of a vertex shape. If it is NULL for shapes, then the names of all defined vertex shapes are returned.

- **coords, el, params, end, v**
  - See parameters of the clipping/plotting functions below.

- **clip**
  - An R function object, the clipping function.

- **plot**
  - An R function object, the plotting function.
Named list, additional plot/vertex/edge parameters. The element named define
the new parameters, and the elements themselves define their default values.
Vertex parameters should have a prefix ‘vertex.’, edge parameters a prefix
‘edge.’. Other general plotting parameters should have a prefix ‘plot.’. See
Details below.

Details

In igraph a vertex shape is defined by two functions: 1) provides information about the size of the
shape for clipping the edges and 2) plots the shape if requested. These functions are called “shape
functions” in the rest of this manual page. The first one is the clipping function and the second is
the plotting function.

The clipping function has the following arguments:

- **coords**: A matrix with four columns, it contains the coordinates of the vertices for the edge list
  supplied in the `el` argument.
- **el**: A matrix with two columns, the edges of which some end points will be clipped. It should have
  the same number of rows as `coords`.
- **params**: This is a function object that can be called to query vertex/edge/plot graphical parameters.
  The first argument of the function is “vertex”, “edge” or “plot” to decide the type of the
  parameter, the second is a character string giving the name of the parameter. E.g.

  ```
  params("vertex", "size")
  ```

- **end**: Character string, it gives which end points will be used. Possible values are “both”, “from”
  and “to”. If “from” the function is expected to clip the first column in the `el` edge list, “to”
  selects the second column, “both” selects both.

The clipping function should return a matrix with the same number of rows as the `el`
arguments. If `end` is both then the matrix must have four columns, otherwise two. The matrix contains the
modified coordinates, with the clipping applied.

The plotting function has the following arguments:

- **coords**: The coordinates of the vertices, a matrix with two columns.
- **v**: The ids of the vertices to plot. It should match the number of rows in the `coords` argument.
- **params**: The same as for the clipping function, see above.

The return value of the plotting function is not used.

shapes can be used to list the names of all installed vertex shapes, by calling it without arguments,
or setting the shape argument to `NULL`. If a shape name is given, then the clipping and plotting
functions of that shape are returned in a named list.

add_shape can be used to add new vertex shapes to igraph. For this one must give the clipping and
plotting functions of the new shape. It is also possible to list the plot/vertex/edge parameters, in
the `parameters` argument, that the clipping and/or plotting functions can make use of. An example
would be a generic regular polygon shape, which can have a parameter for the number of sides.

shape_noclip is a very simple clipping function that the user can use in their own shape definitions.
It does no clipping, the edges will be drawn exactly until the listed vertex position coordinates.

shape_noplot is a very simple (and probably not very useful) plotting function, that does not plot
anything.
Value

shapes returns a character vector if the shape argument is NULL. It returns a named list with entries named ‘clip’ and ‘plot’, both of them R functions.

add_shape returns TRUE, invisibly.

shape_noclip returns the appropriate columns of its coords argument.

Examples

# all vertex shapes, minus "raster", that might not be available
shapes <- setdiff(shapes(), "")
g <- make_ring(length(shapes))
set.seed(42)
plot(g, vertex.shape=shapes, vertex.label=shapes, vertex.label.dist=1,
    vertex.size=15, vertex.size2=15,
    vertex.pie=lapply(shapes, function(x) if (x=="pie") 2:6 else 0),
    vertex.pie.color=list(heat.colors(5)))

# add new vertex shape, plot nothing with no clipping
add_shape("nil")
plot(g, vertex.shape="nil")

#########################################################################
# triangle vertex shape

mytriangle <- function(coords, v=NULL, params) {
  vertex.color <- params("vertex", "color")
  if (length(vertex.color) != 1 && !is.null(v)) {
    vertex.color <- vertex.color[v]
  }
  vertex.size <- 1/200 * params("vertex", "size")
  if (length(vertex.size) != 1 && !is.null(v)) {
    vertex.size <- vertex.size[v]
  }
  symbols(x=coords[,1], y=coords[,2], bg=vertex.color,
    stars=cbind(vertex.size, vertex.size, vertex.size),
    add=TRUE, inches=FALSE)
}
# clips as a circle
add_shape("triangle", clip=shapes("circle")$clip,
  plot=mytriangle)
plot(g, vertex.shape="triangle", vertex.color=rainbow(vcount(g)),
    vertex.size=seq(10,20,length.out=vcount(g)))

#########################################################################
# generic star vertex shape, with a parameter for number of rays

mystar <- function(coords, v=NULL, params) {
  vertex.color <- params("vertex", "color")
  if (length(vertex.color) != 1 && !is.null(v)) {
    vertex.color <- vertex.color[v]
  }
  vertex.size <- 1/200 * params("vertex", "size")
  if (length(vertex.size) != 1 && !is.null(v)) {
    vertex.size <- vertex.size[v]
  }
  norays <- params("vertex", "norays")
  if (length(norays) != 1 && !is.null(v)) {
    norays <- norays[v]
  }
  symbols(x=coords[,1], y=coords[,2], bg=vertex.color,
    vertices=norays, add=TRUE, inches=FALSE)
}
# clips as a circle
add_shape("star", clip=shapes("circle")$clip,
  plot=mystar)
plot(g, vertex.shape="star", vertex.color=rainbow(vcount(g)),
    vertex.size=seq(10,20,length.out=vcount(g)))
similarity <- function(graph, vids = V(graph), mode = c("all", "out", "in", "total"), loops = FALSE, method = c("jaccard", "dice", "invlogweighted")) {
    similarity(
        graph, 
        vids = V(graph), 
        mode = c("all", "out", "in", "total"), 
        loops = FALSE, 
        method = c("jaccard", "dice", "invlogweighted")
    )
}

Arguments

- graph: The input graph.
- vids: The vertex ids for which the similarity is calculated.
- mode: The type of neighboring vertices to use for the calculation, possible values: 'out', 'in', 'all'.
- loops: Whether to include vertices themselves in the neighbor sets.
- method: The method to use.

Details

The Jaccard similarity coefficient of two vertices is the number of common neighbors divided by the number of vertices that are neighbors of at least one of the two vertices being considered. The jaccard method calculates the pairwise Jaccard similarities for some (or all) of the vertices.
The Dice similarity coefficient of two vertices is twice the number of common neighbors divided by the sum of the degrees of the vertices. Method `dice` calculates the pairwise Dice similarities for some or all of the vertices.

The inverse log-weighted similarity of two vertices is the number of their common neighbors, weighted by the inverse logarithm of their degrees. It is based on the assumption that two vertices should be considered more similar if they share a low-degree common neighbor, since high-degree common neighbors are more likely to appear even by pure chance. Isolated vertices will have zero similarity to any other vertex. Self-similarities are not calculated. See the following paper for more details: Lada A. Adamic and Eytan Adar: Friends and neighbors on the Web. Social Networks, 25(3):211-230, 2003.

Value

A `length(vids)` by `length(vids)` numeric matrix containing the similarity scores. This argument is ignored by the `invlogweighted` method.

Author(s)

Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com> for the manual page.

References


See Also

`cocitation` and `bibcoupling`

Examples

```r
  g <- make_ring(5)
  similarity(g, method = "dice")
  similarity(g, method = "jaccard")
```

---

### simplified

Constructor modifier to drop multiple and loop edges

**Description**

Constructor modifier to drop multiple and loop edges

**Usage**

`simplified()`

**See Also**

Other constructor modifiers: `with_edge()`, `with_graph()`, `with_vertex()`, `without_attr()`, `without_loops()`, `without_multiples()`
Examples

```r
sample_(pa(10, m = 3, algorithm = "bag"))
sample_(pa(10, m = 3, algorithm = "bag"), simplified())
```

---

**simplify**  

**Simple graphs**

**Description**

Simple graphs are graphs which do not contain loop and multiple edges.

**Usage**

```r
simplify(
  graph,
  remove.multiple = TRUE,
  remove.loops = TRUE,
  edge.attr.comb = igraph_opt("edge.attr.comb")
)
```

```r
is_simple(graph)
simplify_and_colorize(graph)
```

**Arguments**

- **graph**: The graph to work on.
- **remove.multiple**: Logical, whether the multiple edges are to be removed.
- **remove.loops**: Logical, whether the loop edges are to be removed.
- **edge.attr.comb**: Specifies what to do with edge attributes, if `remove.multiple=TRUE`. In this case many edges might be mapped to a single one in the new graph, and their attributes are combined. Please see `attribute.combination` for details on this.

**Details**

A loop edge is an edge for which the two endpoints are the same vertex. Two edges are multiple edges if they have exactly the same two endpoints (for directed graphs order does matter). A graph is simple is it does not contain loop edges and multiple edges.

`is_simple` checks whether a graph is simple.

`simplify` removes the loop and/or multiple edges from a graph. If both `remove.loops` and `remove.multiple` are `TRUE` the function returns a simple graph.

`simplify_and_colorize` constructs a new, simple graph from a graph and also sets a color attribute on both the vertices and the edges. The colors of the vertices represent the number of self-loops that were originally incident on them, while the colors of the edges represent the multiplicities of the same edges in the original graph. This allows one to take into account the edge multiplicities and the number of loop edges in the VF2 isomorphism algorithm. Other graph, vertex and edge attributes from the original graph are discarded as the primary purpose of this function is to facilitate the usage of multigraphs with the VF2 algorithm.
spectrum

Value

a new graph object with the edges deleted.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

which_loop, which_multiple and count_multiple, delete_edges, delete_vertices

Examples

g <- graph( c(1,2,1,2,3,3) )
is_simple(g)
is_simple(simplify(g, remove.loops=FALSE))
is_simple(simplify(g, remove.multiple=FALSE))
is_simple(simplify(g))

spectrum

Eigenvalues and eigenvectors of the adjacency matrix of a graph

Description

Calculate selected eigenvalues and eigenvectors of a (supposedly sparse) graph.

Usage

spectrum(
  graph,
  algorithm = c("arpack", "auto", "lapack", "comp_auto", "comp_lapack", "comp_arpack"),
  which = list(),
  options = arpack_defaults
)

Arguments

graph
algorithm
which
options

The input graph, can be directed or undirected.
The algorithm to use. Currently only arpack is implemented, which uses the
ARPACK solver. See also arpack.
A list to specify which eigenvalues and eigenvectors to calculate. By default the
leading (i.e. largest magnitude) eigenvalue and the corresponding eigenvector is
calculated.
Options for the ARPACK solver. See arpack_defaults.
Details

The `which` argument is a list and it specifies which eigenvalues and corresponding eigenvectors to calculate. There are eight options:

1. Eigenvalues with the largest magnitude. Set `pos` to `LM`, and `howmany` to the number of eigenvalues you want.
2. Eigenvalues with the smallest magnitude. Set `pos` to `SM` and `howmany` to the number of eigenvalues you want.
3. Largest eigenvalues. Set `pos` to `LA` and `howmany` to the number of eigenvalues you want.
4. Smallest eigenvalues. Set `pos` to `SA` and `howmany` to the number of eigenvalues you want.
5. Eigenvalues from both ends of the spectrum. Set `pos` to `BE` and `howmany` to the number of eigenvalues you want. If `howmany` is odd, then one more eigenvalue is returned from the larger end.
6. Selected eigenvalues. This is not (yet) implemented currently.
7. Eigenvalues in an interval. This is not (yet) implemented.
8. All eigenvalues. This is not implemented yet. The standard `eigen` function does a better job at this, anyway.

Note that ARPACK might be unstable for graphs with multiple components, e.g. graphs with isolate vertices.

Value

Depends on the algorithm used.

For `arpack` a list with three entries is returned:

- `options` See the return value for `arpack` for a complete description.
- `values` Numeric vector, the eigenvalues.
- `vectors` Numeric matrix, with the eigenvectors as columns.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

- `as_adj` to create a (sparse) adjacency matrix.

Examples

```r
## Small example graph, leading eigenvector by default
kite <- make_graph("Krackhardt_kite")
spectrum(kite)[c("values", "vectors")]

## Double check
eigen(as_adj(kite, sparse=FALSE))$vectors[,1]

## Should be the same as 'eigen_centrality' (but rescaled)
cor(eigen_centrality(kite)$vector, spectrum(kite)$vectors)

## Smallest eigenvalues
spectrum(kite, which=list(pos="SM", howmany=2))$values
```
split_join_distance

**Description**

The split-join distance between partitions A and B is the sum of the projection distance of A from B and the projection distance of B from A. The projection distance is an asymmetric measure and it is defined as follows:

**Usage**

```
split_join_distance(comm1, comm2)
```

**Arguments**

- `comm1`: The first community structure.
- `comm2`: The second community structure.

**Details**

First, each set in partition A is evaluated against all sets in partition B. For each set in partition A, the best matching set in partition B is found and the overlap size is calculated. (Matching is quantified by the size of the overlap between the two sets). Then, the maximal overlap sizes for each set in A are summed together and subtracted from the number of elements in A.

The split-join distance will be returned as two numbers, the first is the projection distance of the first partition from the second, while the second number is the projection distance of the second partition from the first. This makes it easier to detect whether a partition is a subpartition of the other, since in this case, the corresponding distance will be zero.

**Value**

Two integer numbers, see details below.

**References**


---

srand

**Description**

Deprecated function, used to set random seed of the C library’s RNG

**Usage**

```
srand(seed)
```
Arguments

seed Ignored.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

st_cuts List all (s,t)-cuts of a graph

Description

List all (s,t)-cuts in a directed graph.

Usage

st_cuts(graph, source, target)

Arguments

graph The input graph. It must be directed.
source The source vertex.
target The target vertex.

Details

Given a G directed graph and two, different and non-adjacent vertices, s and t, an (s, t)-cut is a set of edges, such that after removing these edges from G there is no directed path from s to t.

Value

A list with entries:

cuts A list of numeric vectors containing edge ids. Each vector is an (s, t)-cut.
partition1s A list of numeric vectors containing vertex ids, they correspond to the edge cuts. Each vertex set is a generator of the corresponding cut, i.e. in the graph $G = (V, E)$, the vertex set $X$ and its complementer $V - X$, generates the cut that contains exactly the edges that go from $X$ to $V - X$.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

st_min_cuts to list all minimum cuts.
Examples

# A very simple graph

```r
g <- graph_from_literal(a -- b -- c -- d -- e)
st_cuts(g, source="a", target="e")
```

# A somewhat more difficult graph

```r
g2 <- graph_from_literal(s --+ a:b, a:b --+ t,
                        a --+ 1:2:3, 1:2:3 --+ b)
st_cuts(g2, source="s", target="t")
```

---

**st_min_cuts**  
List all minimum \((s, t)\)-cuts of a graph

Description

Listing all minimum \((s, t)\)-cuts of a directed graph, for given \(s\) and \(t\).

Usage

\[
st_{\text{min}_\text{cuts}}(\text{graph}, \text{source}, \text{target}, \text{capacity} = \text{NULL})
\]

Arguments

- **graph**  
The input graph. It must be directed.
- **source**  
The id of the source vertex.
- **target**  
The id of the target vertex.
- **capacity**  
Numeric vector giving the edge capacities. If this is `NULL` and the graph has a `weight` edge attribute, then this attribute defines the edge capacities. For forcing unit edge capacities, even for graphs that have a `weight` edge attribute, supply `NA` here.

Details

Given a \(G\) directed graph and two, different and non-adjacent vertices, \(s\) and \(t\), an \((s, t)\)-cut is a set of edges, such that after removing these edges from \(G\) there is no directed path from \(s\) to \(t\).

The size of an \((s, t)\)-cut is defined as the sum of the capacities (or weights) in the cut. For unweighted (=equally weighted) graphs, this is simply the number of edges.

An \((s, t)\)-cut is minimum if it is of the smallest possible size.

Value

A list with entries:

- **value**  
  Numeric scalar, the size of the minimum cut(s).
- **cuts**  
  A list of numeric vectors containing edge ids. Each vector is a minimum \((s, t)\)-cut.
- **partition1s**  
  A list of numeric vectors containing vertex ids, they correspond to the edge cuts. Each vertex set is a generator of the corresponding cut, i.e. in the graph \(G = (V, E)\), the vertex set \(X\) and its complementer \(V - X\), generates the cut that contains exactly the edges that go from \(X\) to \(V - X\).
stochastic_matrix

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

References

See Also
st_cuts, min_separators

Examples

```r
# A difficult graph, from the Provan-Shier paper
s <- graph_from_literal(s --+ a:b, a:b --+ t,
a --+ 1:2:3:4:5, 1:2:3:4:5 --+ b)
st_min_cuts(s, source="s", target="t")
```

Description
Retrieves the stochastic matrix of a graph of class igraph.

Usage

```r
stochastic_matrix(
  graph,
  column.wise = FALSE,
  sparse = igraph_opt("sparsematrices")
)
```

Arguments

- **graph**: The input graph. Must be of class igraph.
- **column.wise**: If FALSE, then the rows of the stochastic matrix sum up to one; otherwise it is the columns.
- **sparse**: Logical scalar, whether to return a sparse matrix. The Matrix package is needed for sparse matrices.

Details

Let \( M \) be an \( n \times n \) adjacency matrix with real non-negative entries. Let us define \( D = \text{diag}(\sum_i M_{1i}, \ldots, \sum_i M_{ni}) \). The (row) stochastic matrix is defined as

\[
W = D^{-1} M,
\]

where it is assumed that \( D \) is non-singular. Column stochastic matrices are defined in a symmetric way.
**strength**  

*Value*  

A regular matrix or a matrix of class `Matrix` if a `sparse` argument was `TRUE`.

*Author(s)*  

Gabor Csardi &lt;csardi.gabor@gmail.com&gt;

*See Also*  

`as_adj`

*Examples*  

```r  
library(Matrix)  
## g is a large sparse graph  
g &lt; sample_pa(n = 10^5, power = 2, directed = FALSE)  
W &lt; stochastic_matrix(g, sparse=TRUE)  
  
## a dense matrix here would probably not fit in the memory  
class(W)  
## may not be exactly 1, due to numerical errors  
max(abs(rowSums(W))-1)  
```  

---  

**strength**  

*Strength or weighted vertex degree*

*Description*  

Summing up the edge weights of the adjacent edges for each vertex.

*Usage*  

```r  
strength(  
  graph,  
  vids = V(graph),  
  mode = c("all", "out", "in", "total"),  
  loops = TRUE,  
  weights = NULL  
)  
```  

*Arguments*  

- **graph**: The input graph.  
- **vids**: The vertices for which the strength will be calculated.  
- **mode**: Character string, “out” for out-degree, “in” for in-degree or “all” for the sum of the two. For undirected graphs this argument is ignored.  
- **loops**: Logical; whether the loop edges are also counted.  
- **weights**: Weight vector. If the graph has a weight edge attribute, then this is used by default. If the graph does not have a weight edge attribute and this argument is `NULL`, then a warning is given and `degree` is called.
Value

A numeric vector giving the strength of the vertices.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

degree for the unweighted version.

Examples

g <- make_star(10)
E(g)$weight <- seq(ecount(g))
strength(g)
strength(g, mode="out")
strength(g, mode="in")

# No weights, a warning is given
g <- make_ring(10)
strength(g)

---

subcomponent

In- or out- component of a vertex

Description

Finds all vertices reachable from a given vertex, or the opposite: all vertices from which a given vertex is reachable via a directed path.

Usage

subcomponent(graph, v, mode = c("all", "out", "in"))

Arguments

graph The graph to analyze.
v The vertex to start the search from.
mode Character string, either “in”, “out” or “all”. If “in” all vertices from which v is reachable are listed. If “out” all vertices reachable from v are returned. If “all” returns the union of these. It is ignored for undirected graphs.

Details

A breadth-first search is conducted starting from vertex v.
**Value**

Numeric vector, the ids of the vertices in the same component as \( v \).

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**See Also**

components

**Examples**

```r
g <- sample_gnp(100, 1/200)
subcomponent(g, 1, "in")
subcomponent(g, 1, "out")
subcomponent(g, 1, "all")
```

**Description**

subgraph creates a subgraph of a graph, containing only the specified vertices and all the edges among them.

**Usage**

```r
subgraph(graph, vids)
induced_subgraph(
  graph,
  vids,
  impl = c("auto", "copy_and_delete", "create_from_scratch")
)
```

```r
subgraph.edges(graph, eids, delete.vertices = TRUE)
```

**Arguments**

- **graph**
  The original graph.
- **vids**
  Numeric vector, the vertices of the original graph which will form the subgraph.
- **impl**
  Character scalar, to choose between two implementation of the subgraph calculation. ‘copy_and_delete’ copies the graph first, and then deletes the vertices and edges that are not included in the result graph. ‘create_from_scratch’ searches for all vertices and edges that must be kept and then uses them to create the graph from scratch. ‘auto’ chooses between the two implementations automatically, using heuristics based on the size of the original and the result graph.
- **eids**
  The edge ids of the edges that will be kept in the result graph.
- **delete.vertices**
  Logical scalar, whether to remove vertices that do not have any adjacent edges in eids.
Details

induced_subgraph calculates the induced subgraph of a set of vertices in a graph. This means that exactly the specified vertices and all the edges between them will be kept in the result graph.

subgraph.edges calculates the subgraph of a graph. For this function one can specify the vertices and edges to keep. This function will be renamed to subgraph in the next major version of igraph.

The subgraph function currently does the same as induced_subgraph (assuming 'auto' as the impl argument), but this behaviour is deprecated. In the next major version, subgraph will overtake the functionality of subgraph.edges.

Value
A new graph object.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

Examples

g <- make_ring(10)
g2 <- induced_subgraph(g, 1:7)
g3 <- subgraph.edges(g, 1:5, 1:5)

subgraph_centrality
Find subgraph centrality scores of network positions

Description
Subgraph centrality of a vertex measures the number of subgraphs a vertex participates in, weighting them according to their size.

Usage
subgraph_centrality(graph, diag = FALSE)

Arguments

graph The input graph, it should be undirected, but the implementation does not check this currently.

diag Boolean scalar, whether to include the diagonal of the adjacency matrix in the analysis. Giving FALSE here effectively eliminates the loops edges from the graph before the calculation.

Details
The subgraph centrality of a vertex is defined as the number of closed loops originating at the vertex, where longer loops are exponentially downweighted.

Currently the calculation is performed by explicitly calculating all eigenvalues and eigenvectors of the adjacency matrix of the graph. This effectively means that the measure can only be calculated for small graphs.
subgraph_isomorphic

Value
A numeric vector, the subgraph centrality scores of the vertices.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com> based on the Matlab code by Ernesto Estrada

References

See Also
eigen_centrality, page_rank

Examples
```r
g <- sample_pa(100, m=4, dir=FALSE)
sc <- subgraph_centrality(g)
cor(degree(g), sc)
```

---

subgraph_isomorphic Decide if a graph is subgraph isomorphic to another one

Description
Decide if a graph is subgraph isomorphic to another one

Usage
```
subgraph_isomorphic(pattern, target, method = c("auto", "lad", "vf2"), ...)
```

```
is_subgraph_isomorphic_to(
  pattern,
  target,
  method = c("auto", "lad", "vf2"),
  ...
)
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>pattern</td>
<td>The smaller graph, it might be directed or undirected. Undirected graphs are treated as directed graphs with mutual edges.</td>
</tr>
<tr>
<td>target</td>
<td>The bigger graph, it might be directed or undirected. Undirected graphs are treated as directed graphs with mutual edges.</td>
</tr>
<tr>
<td>...</td>
<td>Additional arguments, passed to the various methods.</td>
</tr>
</tbody>
</table>
Value

Logical scalar, TRUE if the pattern is isomorphic to a (possibly induced) subgraph of target.

‘auto’ method

This method currently selects ‘lad’, always, as it seems to be superior on most graphs.

‘lad’ method

This is the LAD algorithm by Solnon, see the reference below. It has the following extra arguments:

domains If not NULL, then it specifies matching restrictions. It must be a list of target vertex sets, given as numeric vertex ids or symbolic vertex names. The length of the list must be vcount(pattern) and for each vertex in pattern it gives the allowed matching vertices in target. Defaults to NULL.

induced Logical scalar, whether to search for an induced subgraph. It is FALSE by default.

time.limit The processor time limit for the computation, in seconds. It defaults to Inf, which means no limit.

‘vf2’ method

This method uses the VF2 algorithm by Cordella, Foggia et al., see references below. It supports vertex and edge colors and have the following extra arguments:

vertex.color1, vertex.color2 Optional integer vectors giving the colors of the vertices for colored graph isomorphism. If they are not given, but the graph has a “color” vertex attribute, then it will be used. If you want to ignore these attributes, then supply NULL for both of these arguments. See also examples below.

edge.color1, edge.color2 Optional integer vectors giving the colors of the edges for edge-colored (sub)graph isomorphism. If they are not given, but the graph has a “color” edge attribute, then it will be used. If you want to ignore these attributes, then supply NULL for both of these arguments.

References


See Also

Other graph isomorphism: count_isomorphisms(), count_subgraph_isomorphisms(), graph_from_isomorphism_class(), isomorphic(), isomorphism_class(), isomorphisms(), subgraph_isomorphisms()

Examples

# A LAD example
pattern <- make_graph(~ 1:2:3:4:5,
                      1 - 2:5, 2 - 1:5:3, 3 - 2:4, 4 - 3:5, 5 - 4:2:1)
target <- make_graph(~ 1:2:3:4:5:6:7:8:9,
                      1 - 2:5:7, 2 - 1:5:3, 3 - 2:4, 4 - 3:5:6:8:9,
subgraph_isomorphisms

5 - 1:2:4:6:7, 6 - 7:5:4:9, 7 - 1:5:6,
8 - 4:9, 9 - 6:4:8)
domains <- list(`1` = c(1,3,9), `2` = c(5,6,7,8), `3` = c(2,4,6,7,8,9),
 `4` = c(1,3,9), `5` = c(2,4,8,9))
subgraph_isomorphisms(pattern, target)
subgraph_isomorphisms(pattern, target, induced = TRUE)
subgraph_isomorphisms(pattern, target, domains = domains)

# Directed LAD example
pattern <- make_graph(~ 1:2:3, 1 -> 2:3)
dring <- make_ring(10, directed = TRUE)
subgraph_isomorphic(pattern, dring)

| subgraph_isomorphisms | All isomorphic mappings between a graph and subgraphs of another graph |

Description

All isomorphic mappings between a graph and subgraphs of another graph

Usage

subgraph_isomorphisms(pattern, target, method = c("lad", "vf2"), ...)

Arguments

pattern The smaller graph, it might be directed or undirected. Undirected graphs are treated as directed graphs with mutual edges.
target The bigger graph, it might be directed or undirected. Undirected graphs are treated as directed graphs with mutual edges.
...

Value

A list of vertex sequences, corresponding to all mappings from the first graph to the second.

‘lad’ method

This is the LAD algorithm by Solnon, see the reference below. It has the following extra arguments:

domains If not NULL, then it specifies matching restrictions. It must be a list of target vertex sets, given as numeric vertex ids or symbolic vertex names. The length of the list must be vcount(pattern) and for each vertex in pattern it gives the allowed matching vertices in target. Defaults to NULL.
induced Logical scalar, whether to search for an induced subgraph. It is FALSE by default.
time.limit The processor time limit for the computation, in seconds. It defaults to Inf, which means no limit.
‘vf2’ method

This method uses the VF2 algorithm by Cordella, Foggia et al., see references below. It supports vertex and edge colors and have the following extra arguments:

**vertex.color1, vertex.color2** Optional integer vectors giving the colors of the vertices for colored graph isomorphism. If they are not given, but the graph has a “color” vertex attribute, then it will be used. If you want to ignore these attributes, then supply NULL for both of these arguments. See also examples below.

**edge.color1, edge.color2** Optional integer vectors giving the colors of the edges for edge-colored (sub)graph isomorphism. If they are not given, but the graph has a “color” edge attribute, then it will be used. If you want to ignore these attributes, then supply NULL for both of these arguments.

See Also

Other graph isomorphism: `count_isomorphisms()`, `count_subgraph_isomorphisms()`, `graph_from_isomorphism_class()`, `isomorphic()`, `isomorphism_class()`, `isomorphisms()`, `subgraph_isomorphic()`

tail_of

*Tails of the edge(s) in a graph*

**Description**

For undirected graphs, head and tail is not defined. In this case tail_of returns vertices incident to the supplied edges, and head_of returns the other end(s) of the edge(s).

**Usage**

```
tail_of(graph, es)
```

**Arguments**

- **graph**: The input graph.
- **es**: The edges to query.

**Value**

A vertex sequence with the tail(s) of the edge(s).

See Also

Other structural queries: `[.igraph()`, `[[.igraph()`, `adjacent_vertices()`, `are_adjacent()`, `ends()`, `get.edge.ids()`, `gorder()`, `gsize()`, `head_of()`, `incident_edges()`, `incident()`, `is_directed()`, `neighbors()`
Description

Run simulations for an SIR (susceptible-infected-recovered) model, on a graph

Usage

```r
## S3 method for class 'sir'
time_bins(x, middle = TRUE)

## S3 method for class 'sir'
median(x, na.rm = FALSE, ...)

## S3 method for class 'sir'
quantile(x, comp = c("NI", "NS", "NR"), prob, ...)

sir(graph, beta, gamma, no.sim = 100)
```

Arguments

- `x`: A `sir` object, returned by the `sir` function.
- `middle`: Logical scalar, whether to return the middle of the time bins, or the boundaries.
- `na.rm`: Logical scalar, whether to ignore NA values. `sir` objects do not contain any NA values currently, so this argument is effectively ignored.
- `...`: Additional arguments, ignored currently.
- `comp`: Character scalar. The component to calculate the quantile of. `NI` is infected agents, `NS` is susceptibles, `NR` stands for recovered.
- `prob`: Numeric vector of probabilities, in [0,1], they specify the quantiles to calculate.
- `graph`: The graph to run the model on. If directed, then edge directions are ignored and a warning is given.
- `beta`: Non-negative scalar. The rate of infection of an individual that is susceptible and has a single infected neighbor. The infection rate of a susceptible individual with n infected neighbors is n times beta. Formally this is the rate parameter of an exponential distribution.
- `gamma`: Positive scalar. The rate of recovery of an infected individual. Formally, this is the rate parameter of an exponential distribution.
- `no.sim`: Integer scalar, the number simulation runs to perform.

Details

The SIR model is a simple model from epidemiology. The individuals of the population might be in three states: susceptible, infected and recovered. Recovered people are assumed to be immune to the disease. Susceptibles become infected with a rate that depends on their number of infected neighbors. Infected people become recovered with a constant rate.

The function `sir` simulates the model.
Function `time_bins` bins the simulation steps, using the Freedman-Diaconis heuristics to determine the bin width.

Function `median` and `quantile` calculate the median and quantiles of the results, respectively, in bins calculated with `time_bins`.

**Value**

For `sir` the results are returned in an object of class ‘sir’, which is a list, with one element for each simulation. Each simulation is itself a list with the following elements. They are all numeric vectors, with equal length:

- **times** The times of the events.
- **NS** The number of susceptibles in the population, over time.
- **NI** The number of infected individuals in the population, over time.
- **NR** The number of recovered individuals in the population, over time.

Function `time_bins` returns a numeric vector, the middle or the boundaries of the time bins, depending on the `middle` argument.

`median` returns a list of three named numeric vectors, `NS`, `NI` and `NR`. The names within the vectors are created from the time bins.

`quantile` returns the same vector as `median` (but only one, the one requested) if only one quantile is requested. If multiple quantiles are requested, then a list of these vectors is returned, one for each quantile.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>. Eric Kolaczyk ([http://math.bu.edu/people/kolaczyk/](http://math.bu.edu/people/kolaczyk/)) wrote the initial version in R.

**References**


**See Also**

`plot.sir` to conveniently plot the results

**Examples**

```r
  g <- sample_gnm(100, 100)
  sm <- sir(g, beta=5, gamma=1)
  plot(sm)
```
tkigraph

Experimental basic igraph GUI

Description
This function starts an experimental GUI to some igraph functions. The GUI was written in Tcl/Tk, so it is cross platform.

Usage
tkigraph()

Details
tkigraph has its own online help system, please see that for the details about how to use it.

Value
Returns NULL, invisibly.

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

See Also
tkplot for interactive plotting of graphs.

tkplot

Interactive plotting of graphs

Description
tkplot and its companion functions serve as an interactive graph drawing facility. Not all parameters of the plot can be changed interactively right now though, e.g. the colors of vertices, edges, and also others have to be pre-defined.

Usage
tkplot(graph, canvas.width = 450, canvas.height = 450, ...)
tk_close(tkp.id, window.close = TRUE)
tk_off()
tk_fit(tkp.id, width = NULL, height = NULL)
tk_center(tkp.id)
tk_reshape(tkp.id, newlayout, ..., params)
tk_postscript(tkp.id)

tk_coords(tkp.id, norm = FALSE)

tk_set_coords(tkp.id, coords)

tk_rotate(tkp.id, degree = NULL, rad = NULL)

tk_canvas(tkp.id)

Arguments

graph The graph to plot.

canvas.width, canvas.height The size of the tkplot drawing area.

... Additional plotting parameters. See igraph.plotting for the complete list.

tkp.id The id of the tkplot window to close/reshape/etc.

window.close Leave this on the default value.

width The width of the rectangle for generating new coordinates.

height The height of the rectangle for generating new coordinates.

newlayout The new layout, see the layout parameter of tkplot.

params Extra parameters in a list, to pass to the layout function.

norm Logical, should we norm the coordinates.

coords Two-column numeric matrix, the new coordinates of the vertices, in absolute coordinates.

degree The degree to rotate the plot.

rad The degree to rotate the plot, in radian.

Details
tkplot is an interactive graph drawing facility. It is not very well developed at this stage, but it should be still useful.

It's handling should be quite straightforward most of the time, here are some remarks and hints.

There are different popup menus, activated by the right mouse button, for vertices and edges. Both operate on the current selection if the vertex/edge under the cursor is part of the selection and operate on the vertex/edge under the cursor if it is not.

One selection can be active at a time, either a vertex or an edge selection. A vertex/edge can be added to a selection by holding the control key while clicking on it with the left mouse button. Doing this again deselect the vertex/edge.

Selections can be made also from the Select menu. The 'Select some vertices' dialog allows to give an expression for the vertices to be selected: this can be a list of numeric R expressions separated by commas, like '1,2:10,12,14,15' for example. Similarly in the 'Select some edges' dialog two such lists can be given and all edges connecting a vertex in the first list to one in the second list will be selected.

In the color dialog a color name like 'orange' or RGB notation can also be used.
The `tkplot` command creates a new Tk window with the graphical representation of graph. The command returns an integer number, the tkplot id. The other commands utilize this id to be able to query or manipulate the plot.

`tk_close` closes the Tk plot with id `tkp.id`.

`tk_off` closes all Tk plots.

`tk_fit` fits the plot to the given rectangle (`width` and `height`), if some of these are `NULL` the actual physical width od height of the plot window is used.

`tk_reshape` applies a new layout to the plot, its optional parameters will be collected to a list analogous to `layout.par`.

`tk_postscript` creates a dialog window for saving the plot in postscript format.

`tk_canvas` returns the Tk canvas object that belongs to a graph plot. The canvas can be directly manipulated then, eg. labels can be added, it could be saved to a file programmatically, etc. See an example below.

`tk_coords` returns the coordinates of the vertices in a matrix. Each row corresponds to one vertex.

`tk_set_coords` sets the coordinates of the vertices. A two-column matrix specifies the new positions, with each row corresponding to a single vertex.

`tk_center` shifts the figure to the center of its plot window.

`tk_rotate` rotates the figure, its parameter can be given either in degrees or in radians.

### Value

- `tkplot` returns an integer, the id of the plot, this can be used to manipulate it from the command line.
- `tk_canvas` returns `tkwin` object, the Tk canvas.
- `tk_coords` returns a matrix with the coordinates.

`tk_close`, `tk_off`, `tk_fit`, `tk_reshape`, `tk_postscript`, `tk_center` and `tk_rotate` return `NULL` invisibly.

### Examples

```r
g <- make_ring(10)
tkplot(g)

## Saving a tkplot() to a file programmatically
g <- make_star(10, center=10)
E(g)$width <- sample(1:10, ecount(g), replace=TRUE)
lay <- layout_nicely(g)
id <- tkplot(g, layout=lay)
canvas <- tk_canvas(id)
tcltk::tkpostscript(canvas, file="/tmp/output.eps")
tk_close(id)

## Setting the coordinates and adding a title label
g <- make_ring(10)
id <- tkplot(make_ring(10), canvas.width=450, canvas.height=500)
canvas <- tk_canvas(id)
padding <- 20
```
to_prufer

Convert a tree graph to its Prufer sequence

Description

to_prufer converts a tree graph into its Prufer sequence.

Usage

to_prufer(graph)

Arguments

graph The graph to convert to a Prufer sequence

Details

The Prufer sequence of a tree graph with n labeled vertices is a sequence of n-2 numbers, constructed as follows. If the graph has more than two vertices, find a vertex with degree one, remove it from the tree and add the label of the vertex that it was connected to to the sequence. Repeat until there are only two vertices in the remaining graph.

Value

The Prufer sequence of the graph, represented as a numeric vector of vertex IDs in the sequence.

See Also

make_from_prufer to construct a graph from its Prufer sequence

Examples

g <- make_tree(13, 3)
to_prufer(g)
**topo_sort**

*Topological sorting of vertices in a graph*

Description

A topological sorting of a directed acyclic graph is a linear ordering of its nodes where each node comes before all nodes to which it has edges.

Usage

topo_sort(graph, mode = c("out", "all", "in"))

Arguments

g**raph**
The input graph, should be directed

**mode**
Specifies how to use the direction of the edges. For “out”, the sorting order ensures that each node comes before all nodes to which it has edges, so nodes with no incoming edges go first. For “in”, it is quite the opposite: each node comes before all nodes from which it receives edges. Nodes with no outgoing edges go first.

Details

Every DAG has at least one topological sort, and may have many. This function returns a possible topological sort among them. If the graph is not acyclic (it has at least one cycle), a partial topological sort is returned and a warning is issued.

Value

A vertex sequence (by default, but see the return.vs.es option of igraph_options) containing vertices in topologically sorted order.

Author(s)

Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com> for the R interface

Examples

g <- barabasi.game(100)
topo_sort(g)
transitivity  

Transitivity of a graph

Description

Transitivity measures the probability that the adjacent vertices of a vertex are connected. This is sometimes also called the clustering coefficient.

Usage

```r
transitivity(
  graph,
  type = c("undirected", "global", "globalundirected", "localundirected", "local",
           "average", "localaverage", "localaverageundirected", "barrat", "weighted"),
  vids = NULL,
  weights = NULL,
  isolates = c("NaN", "zero")
)
```

Arguments

- **graph**: The graph to analyze.
- **type**: The type of the transitivity to calculate. Possible values:
  - "global": The global transitivity of an undirected graph. This is simply the ratio of the count of triangles and connected triples in the graph. In directed graphs, edge directions are ignored.
  - "local": The local transitivity of an undirected graph. It is calculated for each vertex given in the `vids` argument. The local transitivity of a vertex is the ratio of the count of triangles connected to the vertex and the triples centered on the vertex. In directed graphs, edge directions are ignored.
  - "undirected": This is the same as `global`.
  - "globalundirected": This is the same as `global`.
  - "localundirected": This is the same as `local`.
  - "barrat": The weighted transitivity as defined by A. Barrat. See details below.
  - "weighted": The same as `barrat`.
- **vids**: The vertex ids for the local transitivity will be calculated. This will be ignored for global transitivity types. The default value is NULL, in this case all vertices are considered. It is slightly faster to supply NULL here than `V(graph)`.
- **weights**: Optional weights for weighted transitivity. It is ignored for other transitivity measures. If it is `NULL` (the default) and the graph has a `weight` edge attribute, then it is used automatically.
- **isolates**: Character scalar, defines how to treat vertices with degree zero and one. If it is ‘NaN’ then they local transitivity is reported as NaN and they are not included in the averaging, for the transitivity types that calculate an average. If there are no vertices with degree two or higher, then the averaging will still result NaN. If it is ‘zero’, then we report 0 transitivity for them, and they are included in the averaging, if an average is calculated.
transitivity

Details

Note that there are essentially two classes of transitivity measures, one is a vertex-level, the other a
graph level property.

There are several generalizations of transitivity to weighted graphs, here we use the definition by
A. Barrat, this is a local vertex-level quantity, its formula is

\[ C_i^w = \frac{1}{s_i (k_i - 1)} \sum_{j,h} w_{ij} + w_{ih} \frac{a_{ij} a_{ih} a_{jh}}{2} \]

\( s_i \) is the strength of vertex \( i \), see strength, \( a_{ij} \) are elements of the adjacency matrix, \( k_i \) is the vertex
degree, \( w_{ij} \) are the weights.

This formula gives back the normal not-weighted local transitivity if all the edge weights are the
same.

The \texttt{barrat} type of transitivity does not work for graphs with multiple and/or loop edges. If you
want to calculate it for a directed graph, call \texttt{as.undirected} with the \texttt{collapse} mode first.

Value

For ‘global’ a single number, or \texttt{NaN} if there are no connected triples in the graph.
For ‘local’ a vector of transitivity scores, one for each vertex in ‘vids’.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


Examples

```r
g <- make_ring(10)
transitivity(g)
g2 <- sample_gnp(1000, 10/1000)
transitivity(g2)  # this is about 10/1000

# Weighted version, the figure from the Barrat paper
gw <- graph_from_literal(A-B:C:D:E, B-C:D, C-D)
E(gw)$weight <- 1
E(gw)[ V(gw)[name == "A"] %--% V(gw)[name == "E"] ]$weight <- 5
transitivity(gw, vids="A", type="local")
transitivity(gw, vids="A", type="weighted")

# Weighted reduces to "local" if weights are the same

gw2 <- sample_gnp(1000, 10/1000)
E(gw2)$weight <- 1
t1 <- transitivity(gw2, type="local")
t2 <- transitivity(gw2, type="weighted")
all(is.na(t1) == is.na(t2))
```
triad_census

```r
all(na.omit(t1 == t2))
```

---

**triad_census**

**Triad census, subgraphs with three vertices**

---

**Description**

This function counts the different subgraphs of three vertices in a graph.

**Usage**

```r
triad_census(graph)
```

**Arguments**

- `graph` The input graph, it should be directed. An undirected graph results a warning, and undefined results.

**Details**

Triad census was defined by David and Leinhardt (see References below). Every triple of vertices (A, B, C) are classified into the 16 possible states:

- **003** A,B,C, the empty graph.
- **012** A->B, C, the graph with a single directed edge.
- **102** A<->B, C, the graph with a mutual connection between two vertices.
- **021D** A<-B->C, the out-star.
- **021U** A->B<-C, the in-star.
- **021C** A->B->C, directed line.
- **111D** A<->B<-C.
- **021U** A<-B->C, the in-star.
- **021C** A->B->C, directed line.
- **111D** A<->B<-C.
- **111U** A<-B<->C.
- **030T** A->B<->C, A->C.
- **030C** A<->B<->C, A->C.
- **201** A<->B<->C.
- **120D** A<->B->C, A<->C.
- **120U** A->B<->C, A<->C.
- **120C** A->B->C, A<->C.
- **210** A->B<->C, A<->C.
- **300** A<->B<->C, A<->C, the complete graph.

This functions uses the RANDESU motif finder algorithm to find and count the subgraphs, see `motifs`.

**Value**

A numeric vector, the subgraph counts, in the order given in the above description.
unfold_tree

Author(s)
Gabor Csardi <csardi.gabor@gmail.com>

References

See Also
dyad_census for classifying binary relationships, motifs for the underlying implementation.

Examples
```r
g <- sample_gnm(15, 45, directed = TRUE)
triad_census(g)
```

**Description**
Perform a breadth-first search on a graph and convert it into a tree or forest by replicating vertices that were found more than once.

**Usage**
```r
unfold_tree(graph, mode = c("all", "out", "in", "total"), roots)
```

**Arguments**
- `graph` The input graph, it can be either directed or undirected.
- `mode` Character string, defined the types of the paths used for the breadth-first search. “out” follows the outgoing, “in” the incoming edges, “all” and “total” both of them. This argument is ignored for undirected graphs.
- `roots` A vector giving the vertices from which the breadth-first search is performed. Typically it contains one vertex per component.

**Details**
A forest is a graph, whose components are trees.
The `roots` vector can be calculated by simply doing a topological sort in all components of the graph, see the examples below.

**Value**
A list with two components:
- `tree` The result, an igraph object, a tree or a forest.
- `vertex_index` A numeric vector, it gives a mapping from the vertices of the new graph to the vertices of the old graph.
Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

g <- make_tree(10) %du% make_tree(10)
V(g)$id <- seq_len(vcount(g))-1
roots <- sapply(decompose(g), function(x) {
  V(x)$id[ topo_sort(x)[1]+1 ] })
tree <- unfold_tree(g, roots=roots)

union

Description

This is an S3 generic function. See methods("union") for the actual implementations for various S3 classes. Initially it is implemented for igraph graphs and igraph vertex and edge sequences. See union.igraph and union.igraph.vs.

Usage

union(...)

Arguments

... Arguments, their number and interpretation depends on the function that implements union.

Value

Depends on the function that implements this method.

union.igraph

Description

The union of two or more graphs are created. The graphs may have identical or overlapping vertex sets.

Usage

## S3 method for class 'igraph'
union(..., byname = "auto")
Arguments

...  Graph objects or lists of graph objects.
byname  A logical scalar, or the character scalar auto. Whether to perform the operation based on symbolic vertex names. If it is auto, that means TRUE if all graphs are named and FALSE otherwise. A warning is generated if auto and some (but not all) graphs are named.

details

union creates the union of two or more graphs. Edges which are included in at least one graph will be part of the new graph. This function can be also used via the %u% operator.

If the byname argument is TRUE (or auto and all graphs are named), then the operation is performed on symbolic vertex names instead of the internal numeric vertex ids.
union keeps the attributes of all graphs. All graph, vertex and edge attributes are copied to the result. If an attribute is present in multiple graphs and would result a name clash, then this attribute is renamed by adding suffixes: _1, _2, etc.
The name vertex attribute is treated specially if the operation is performed based on symbolic vertex names. In this case name must be present in all graphs, and it is not renamed in the result graph.
An error is generated if some input graphs are directed and others are undirected.

Value

A new graph object.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

Examples

## Union of two social networks with overlapping sets of actors
print_all(net1 %u% net2)

union.igraph.es  Union of edge sequences

description

Union of edge sequences

Usage

## S3 method for class 'igraph.es'
union(...)

Arguments

...  The edge sequences to take the union of.
Details

They must belong to the same graph. Note that this function has 'set' semantics and the multiplicity of edges is lost in the result. (This is to match the behavior of the based unique function.)

Value

An edge sequence that contains all edges in the given sequences, exactly once.

See Also

Other vertex and edge sequence operations: c.igraph.es(), c.igraph.vs(), difference.igraph.es(), difference.igraph.vs(), igraph-es-indexing2, igraph-es-indexing, igraph-vs-indexing2, igraph-vs-indexing, intersection.igraph.es(), intersection.igraph.vs(), rev.igraph.es(), rev.igraph.vs(), union.igraph.es(), union.igraph.vs(), unique.igraph.es(), unique.igraph.vs()

Examples

g <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
union(E(g)[1:6], E(g)[5:9], E(g)['AIJ'])

union.igraph.vs
Union of vertex sequences

Description

Union of vertex sequences

Usage

## S3 method for class 'igraph.vs'
union(...)

Arguments

... The vertex sequences to take the union of.

Details

They must belong to the same graph. Note that this function has 'set' semantics and the multiplicity of vertices is lost in the result. (This is to match the behavior of the based unique function.)

Value

A vertex sequence that contains all vertices in the given sequences, exactly once.

See Also

Other vertex and edge sequence operations: c.igraph.es(), c.igraph.vs(), difference.igraph.es(), difference.igraph.vs(), igraph-es-indexing2, igraph-es-indexing, igraph-vs-indexing2, igraph-vs-indexing, intersection.igraph.es(), intersection.igraph.vs(), rev.igraph.es(), rev.igraph.vs(), union.igraph.es(), union.igraph.vs(), unique.igraph.es(), unique.igraph.vs()
unique.igraph.es

Examples

```r
# Create a ring graph with vertices named with letters
gr <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
union(V(gr)[1:6], V(gr)[5:10])
```

## unique.igraph.es

**Remove duplicate edges from an edge sequence**

### Description

Remove duplicate edges from an edge sequence

### Usage

```r
## S3 method for class 'igraph.es'
unique(x, incomparables = FALSE, ...)
```

### Arguments

- **x**: An edge sequence.
- **incomparables**: a vector of values that cannot be compared. Passed to base function `duplicated`. See details there.
- **...**: Passed to base function `duplicated()`.

### Value

An edge sequence with the duplicate vertices removed.

### See Also


### Examples

```r
# Create a ring graph with vertices named with letters
gr <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
E(gr)[1, 1:5, 1:10, 5:10]
E(gr)[1, 1:5, 1:10, 5:10] %>% unique()
```
unique.igraph.vs  
Remove duplicate vertices from a vertex sequence

Description
Remove duplicate vertices from a vertex sequence

Usage
```r
## S3 method for class 'igraph.vs'
unique(x, incomparables = FALSE, ...)
```

Arguments
- `x`: A vertex sequence.
- `incomparables`: a vector of values that cannot be compared. Passed to base function `duplicated`. See details there.
- `...`: Passed to base function `duplicated()`.

Value
A vertex sequence with the duplicate vertices removed.

See Also

Examples
```r
g <- make_(ring(10), with_vertex_(name = LETTERS[1:10]))
V(g)[1, 1:5, 1:10, 5:10]
V(g)[1, 1:5, 1:10, 5:10] %>% unique()
```

upgrade_graph  

igraph data structure versions

Description
igraph's internal data representation changes sometimes between versions. This means that it is not possible to use igraph objects that were created (and possibly saved to a file) with an older igraph version.

Usage
```r
upgrade_graph(graph)
```
Arguments

graph The input graph.

Details

graph_version queries the current data format, or the data format of a possibly older igraph graph. upgrade_graph can convert an older data format to the current one.

Value

The graph in the current format.

See Also

graph_version to check the current data format version or the version of a graph.

---

\textbf{V} \hspace{1cm} \textit{Vertices of a graph}

Description

Create a vertex sequence (vs) containing all vertices of a graph.

Usage

\texttt{V(graph)}

Arguments

graph The graph

Details

A vertex sequence is just what the name says it is: a sequence of vertices. Vertex sequences are usually used as igraph function arguments that refer to vertices of a graph.

A vertex sequence is tied to the graph it refers to: it really denoted the specific vertices of that graph, and cannot be used together with another graph.

At the implementation level, a vertex sequence is simply a vector containing numeric vertex ids, but it has a special class attribute which makes it possible to perform graph specific operations on it, like selecting a subset of the vertices based on graph structure, or vertex attributes.

A vertex sequence is most often created by the \texttt{V()} function. The result of this includes all vertices in increasing vertex id order. A vertex sequence can be indexed by a numeric vector, just like a regular R vector. See \texttt{\_igraph\_vs} and additional links to other vertex sequence operations below.

Value

A vertex sequence containing all vertices, in the order of their numeric vertex ids.
Indexing vertex sequences

Vertex sequences mostly behave like regular vectors, but there are some additional indexing operations that are specific for them; e.g. selecting vertices based on graph structure, or based on vertex attributes. See \texttt{\texttt{igraph.vs}} for details.

Querying or setting attributes

Vertex sequences can be used to query or set attributes for the vertices in the sequence. See \texttt{\texttt{\$igraph.vs}} for details.

See Also

Other vertex and edge sequences: \texttt{E()}, \texttt{igraph-es-attributes}, \texttt{igraph-es-indexing}, \texttt{igraph-es-indexing2}, \texttt{igraph-vs-attributes}, \texttt{igraph-vs-indexing2}, \texttt{igraph-vs-indexing}, \texttt{print.igraph.es()}, \texttt{print.igraph.vs()}

Examples

\begin{verbatim}
# Vertex ids of an unnamed graph
g <- make_ring(10)
V(g)

# Vertex ids of a named graph
g2 <- make_ring(10) %>%
  set_vertex_attr("name", value = letters[1:10])
V(g2)
\end{verbatim}

---

\textbf{vertex} \hspace{1cm} \textit{Helper function for adding and deleting vertices}

\section*{Description}

This is a helper function that simplifies adding and deleting vertices to/from graphs.

\section*{Usage}

\begin{verbatim}
vertex(...)
vertices(...)
\end{verbatim}

\section*{Arguments}

\begin{verbatim}
... \hspace{1cm} \textit{See details below.}
\end{verbatim}

\section*{Details}

\texttt{vertices} is an alias for \texttt{vertex}.

When adding vertices via +, all unnamed arguments are interpreted as vertex names of the new vertices. Named arguments are interpreted as vertex attributes for the new vertices.

When deleting vertices via -, all arguments of \texttt{vertex} (or \texttt{vertices}) are concatenated via \texttt{c()} and passed to \texttt{delete_vertices}.  

Value

A special object that can be used with together with igraph graphs and the plus and minus operators.

See Also

Other functions for manipulating graph structure: `.igraph()`, `add_edges()`, `add_vertices()`, `delete_edges()`, `delete_vertices()`, `edge()`, `igraph-minus.path()

Examples

```r
G <- make_(ring(10), with_vertex_(name = LETTERS[1:10])) +
  vertices('X', 'Y')
G
plot(G)
```

---

**vertex_attr**

Query vertex attributes of a graph

Description

Query vertex attributes of a graph

Usage

`vertex_attr(graph, name, index = V(graph))`

Arguments

- `graph` The graph.
- `name` Name of the attribute to query. If missing, then all vertex attributes are returned in a list.
- `index` A vertex sequence, to query the attribute only for these vertices.

Value

The value of the vertex attribute, or the list of all vertex attributes, if `name` is missing.

See Also

Other graph attributes: `delete_edge_attr()`, `delete_graph_attr()`, `delete_vertex_attr()`, `edge_attr<-()`, `edge_attr_names()`, `edge_attr()`, `graph_attr<-()`, `graph_attr_names()`, `graph_attr()`, `igraph-dollar.igraph-vs-attributes.set_edge_attr()`, `set_graph_attr()`, `set_vertex_attr()`

Examples

```r
G <- make_(ring(10)) %>%
  set_vertex_attr("color", value = "red") %>%
  set_vertex_attr("label", value = letters[1:10])
vertex_attr(G, "label")
vertex_attr(G)
plot(G)
```
vertex_attr_names

List names of vertex attributes

Description
List names of vertex attributes

Usage
vertex_attr_names(graph)

Arguments
graph
The graph.

Value
Character vector, the names of the vertex attributes.

See Also
Other graph attributes: delete_edge_attr(), delete_graph_attr(), delete_vertex_attr(), edge_attr<-(,), edge_attr_names(), edge_attr(), graph_attr<-(,), graph_attr_names(), graph_attr(), igraph-dollar, igraph-vs-attributes, set_edge_attr(), set_graph_attr(), set_vertex_attr(), vertex_attr<-(,), vertex_attr()

Examples
```r
g <- make_ring(10) %>%
  set_vertex_attr("name", value = LETTERS[1:10]) %>%
  set_vertex_attr("color", value = rep("green", 10))
vertex_attr_names(g)
plot(g)
```

vertex_attr<-

Set one or more vertex attributes

Description
Set one or more vertex attributes

Usage
vertex_attr(graph, name, index = V(graph)) <- value

Arguments
graph
The graph.
name
The name of the vertex attribute to set. If missing, then value must be a named list, and its entries are set as vertex attributes.
index
An optional vertex sequence to set the attributes of a subset of vertices.
value
The new value of the attribute(s) for all (or index) vertices.
vertex_connectivity

Value

The graph, with the vertex attribute(s) added or set.

See Also

Other graph attributes: delete_edge_attr(), delete_graph_attr(), delete_vertex_attr(), edge_attr<-(), edge_attr_names(), edge_attr(), graph_attr<-(), graph_attr_names(), graph_attr(), igraph-dollar, igraph-vs-attributes, set_edge_attr(), set_graph_attr(), set_vertex_attr(), vertex_attr_names(), vertex_attr()

Examples

```r
g <- make_ring(10)
vertex_attr(g) <- list(name = LETTERS[1:10],
                      color = rep("yellow", gorder(g)))
vertex_attr(g, "label") <- V(g)$name
g
plot(g)
```

Description

The vertex connectivity of a graph or two vertices, this is recently also called group cohesion.

Usage

vertex_connectivity(graph, source = NULL, target = NULL, checks = TRUE)

## S3 method for class 'igraph'
cohesion(x, checks = TRUE, ...)

Arguments

- `graph, x`: The input graph.
- `source`: The id of the source vertex, for `vertex_connectivity` it can be `NULL`, see details below.
- `target`: The id of the target vertex, for `vertex_connectivity` it can be `NULL`, see details below.
- `checks`: Logical constant. Whether to check that the graph is connected and also the degree of the vertices. If the graph is not (strongly) connected then the connectivity is obviously zero. Otherwise if the minimum degree is one then the vertex connectivity is also one. It is a good idea to perform these checks, as they can be done quickly compared to the connectivity calculation itself. They were suggested by Peter McMahan, thanks Peter.
- `...`: Ignored.
Details

The vertex connectivity of two vertices (source and target) in a directed graph is the minimum number of vertices needed to remove from the graph to eliminate all (directed) paths from source to target. `vertex_connectivity` calculates this quantity if both the source and target arguments are given and they're not NULL.

The vertex connectivity of a graph is the minimum vertex connectivity of all (ordered) pairs of vertices in the graph. In other words this is the minimum number of vertices needed to remove to make the graph not strongly connected. (If the graph is not strongly connected then this is zero.) `vertex_connectivity` calculates this quantity if neither the source nor target arguments are given. (i.e. they are both NULL.)

A set of vertex disjoint directed paths from source to vertex is a set of directed paths between them whose vertices do not contain common vertices (apart from source and target). The maximum number of vertex disjoint paths between two vertices is the same as their vertex connectivity in most cases (if the two vertices are not connected by an edge).

The cohesion of a graph (as defined by White and Harary, see references), is the vertex connectivity of the graph. This is calculated by `cohesion`.

These three functions essentially calculate the same measure(s), more precisely `vertex_connectivity` is the most general, the other two are included only for the ease of using more descriptive function names.

Value

A scalar real value.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

`max_flow`, `edge_connectivity`, `edge_disjoint_paths`, `adhesion`

Examples

```r
g <- barabasi.game(100, m=1)
g <- delete_edges(g, E(g)[ 100 %--% 1 ])
g2 <- barabasi.game(100, m=5)
g2 <- delete_edges(g2, E(g2)[ 100 %--% 1 ]) vertex_connectivity(g, 100, 1)
vertex_connectivity(g2, 100, 1)
vertex_disjoint_paths(g2, 100, 1)
```

```r
g <- sample_gnp(50, 5/50)
g <- as.directed(g)
g <- induced_subgraph(g, subcomponent(g, 1)) cohesion(g)
```
weighted_cliques

Functions to find weighted cliques, ie. weighted complete subgraphs in a graph

Description

These functions find all, the largest or all the maximal weighted cliques in an undirected graph. The weight of a clique is the sum of the weights of its edges.

Usage

```r
weighted_cliques(
  graph,
  vertex.weights = NULL,
  min.weight = 0,
  max.weight = 0,
  maximal = FALSE
)
```

Arguments

- **graph**: The input graph, directed graphs will be considered as undirected ones, multiple edges and loops are ignored.
- **vertex.weights**: Vertex weight vector. If the graph has a weight vertex attribute, then this is used by default. If the graph does not have a weight vertex attribute and this argument is NULL, then every vertex is assumed to have a weight of 1. Note that the current implementation of the weighted clique finder supports positive integer weights only.
- **min.weight**: Numeric constant, lower limit on the weight of the cliques to find. NULL means no limit, ie. it is the same as 0.
- **max.weight**: Numeric constant, upper limit on the weight of the cliques to find. NULL means no limit.
- **maximal**: Specifies whether to look for all weighted cliques (FALSE) or only the maximal ones (TRUE).

Details

- **weighted_cliques** find all complete subgraphs in the input graph, obeying the weight limitations given in the `min` and `max` arguments.
- **largest_weighted_cliques** finds all largest weighted cliques in the input graph. A clique is largest if there is no other clique whose total weight is larger than the weight of this clique.
- **max_weighted_cliques** finds all maximal weighted cliques in the input graph. A weighted clique is maximal if it cannot be extended to a clique with larger total weight. The largest weighted cliques are always maximal, but a maximal weighted clique is not necessarily the largest.
- **count_max_weighted_cliques** counts the maximal weighted cliques.
- **weighted_clique_num** calculates the weight of the largest weighted clique(s).
which_multiple

Value
weighted_cliques and largest_weighted_cliques return a list containing numeric vectors of vertex IDs. Each list element is a weighted clique, i.e. a vertex sequence of class igraph.vs.
weighted_clique_num and count_max_weighted_cliques return an integer scalar.

Author(s)
Tamas Nepusz <ntamas@gmail.com> and Gabor Csardi <csardi.gabor@gmail.com>

See Also
ivs

Examples
g <- make_graph("zachary")
V(g)$weight <- 1
V(g)[c(1,2,3,4,14)]$weight <- 3
weighted_cliques(g)
weighted_cliques(g, maximal=TRUE)
largest_weighted_cliques(g)
weighted_clique_num(g)

which_multiple

Find the multiple or loop edges in a graph

Description
A loop edge is an edge from a vertex to itself. An edge is a multiple edge if it has exactly the same head and tail vertices as another edge. A graph without multiple and loop edges is called a simple graph.

Usage
which_multiple(graph, eids = E(graph))

Arguments
    graph        The input graph.
    eids         The edges to which the query is restricted. By default this is all edges in the graph.

Details
any_loop decides whether the graph has any loop edges.
which_loop decides whether the edges of the graph are loop edges.
any_multiple decides whether the graph has any multiple edges.
which_multiple decides whether the edges of the graph are multiple edges.
count_multiple counts the multiplicity of each edge of a graph.
Note that the semantics for `which_multiple` and `count_multiple` is different. `which_multiple` gives TRUE for all occurrences of a multiple edge except for one. I.e. if there are three i-j edges in the graph then `which_multiple` returns TRUE for only two of them while `count_multiple` returns '3' for all three.

See the examples for getting rid of multiple edges while keeping their original multiplicity as an edge attribute.

**Value**

`any_loop` and `any_multiple` return a logical scalar. `which_loop` and `which_multiple` return a logical vector. `count_multiple` returns a numeric vector.

**Author(s)**

Gabor Csardi <csardi.gabor@gmail.com>

**See Also**

`simplify` to eliminate loop and multiple edges.

**Examples**

```r
# Loops
g <- graph(c(1,1,2,2,3,3,4,5) )
any_loop(g)
which_loop(g)

# Multiple edges
g <- barabasi.game(10, m=3, algorithm="bag")
any_multiple(g)
which_multiple(g)
count_multiple(g)
which_multiple(simplify(g))
all(count_multiple(simplify(g)) == 1)

# Direction of the edge is important
which_multiple(graph(c(1,2,2,1) ))
which_multiple(graph(c(1,2,2,1), dir=FALSE ))

# Remove multiple edges but keep multiplicity
g <- barabasi.game(10, m=3, algorithm="bag")
E(g)$weight <- count_multiple(g)
g <- simplify(g, edge.attr.comb=list(weight = "min"))
any(which_multiple(g))
E(g)$weight

which_mutual

Find mutual edges in a directed graph

**Description**

This function checks the reciprocal pair of the supplied edges.
Usage

\texttt{which\_mutual(graph, eids = E(graph))}

Arguments

\begin{itemize}
  \item \texttt{graph} \quad The input graph.
  \item \texttt{eids} \quad Edge sequence, the edges that will be probed. By default is includes all edges in the order of their ids.
\end{itemize}

Details

In a directed graph an (A,B) edge is mutual if the graph also includes a (B,A) directed edge. Note that multi-graphs are not handled properly, i.e. if the graph contains two copies of (A,B) and one copy of (B,A), then these three edges are considered to be mutual. Undirected graphs contain only mutual edges by definition.

Value

A logical vector of the same length as the number of edges supplied.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

See Also

\texttt{reciprocity}, \texttt{dyad\_census} if you just want some statistics about mutual edges.

Examples

\begin{verbatim}
g <- sample_gnm(10, 50, directed=TRUE) reciprocity(g) dyad_census(g) which_mutual(g) sum(which_mutual(g))/2 == dyad_census(g)$mut
\end{verbatim}

\begin{verbatim}
with_edge_ Constructor modifier to add edge attributes
\end{verbatim}

Description

Constructor modifier to add edge attributes

Usage

\texttt{with\_edge\_(...)}

Arguments

\begin{itemize}
  \item \texttt{...} \quad The attributes to add. They must be named.
\end{itemize}
See Also

Other constructor modifiers: simplified(with_graph_(), with_vertex_(), without_attr(), without_loops(), without_multiples()

Examples

```r
make_(ring(10),
  with_edge_(
    color = "red",
    weight = rep(1:2, 5))) %>%
  plot()
```

---

**with_graph_**

Constructor modifier to add graph attributes

Description

Constructor modifier to add graph attributes

Usage

```r
with_graph_(...)
```

Arguments

... The attributes to add. They must be named.

See Also

Other constructor modifiers: simplified(with_graph_(), with_vertex_(), without_attr(), without_loops(), without_multiples()

Examples

```r
make_(ring(10), with_graph_(name = "10-ring"))
```

---

**with_igraph_opt**

Run code with a temporary igraph options setting

Description

Run code with a temporary igraph options setting

Usage

```r
with_igraph_opt(options, code)
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>options</td>
<td>A named list of the options to change.</td>
</tr>
<tr>
<td>code</td>
<td>The code to run.</td>
</tr>
</tbody>
</table>
with_vertex_

Value

The result of the code.

See Also

Other igraph options: igraph_options()

Examples

with_igraph_opt(
    list(sparsematrices = FALSE),
    make_ring(10)[]
)
igraph_opt("sparsematrices")

with_vertex_

Constructor modifier to add vertex attributes

Description

Constructor modifier to add vertex attributes

Usage

with_vertex_(...)

Arguments

...

The attributes to add. They must be named.

See Also

Other constructor modifiers: simplified(), with_edge_(), with_graph_(), without_attr(), without_loops(), without_multiples()

Examples

make_(ring(10),
    with_vertex_
    color = "#7fcdbb",
    frame.color = "#7fcdbb",
    name = LETTERS[1:10])) %>%
plot()
without_attr

Constructor modifier to remove all attributes from a graph

Description

Constructor modifier to remove all attributes from a graph

Usage

without_attr()

See Also

Other constructor modifiers: simplified(), with_edge(), with_graph(), with_vertex(), without_loops(), without_multiples()

Examples

g1 <- make_ring(10)
g1

g2 <- make_(ring(10), without_attr())
g2

without_loops

Constructor modifier to drop loop edges

Description

Constructor modifier to drop loop edges

Usage

without_loops()

See Also

Other constructor modifiers: simplified(), with_edge(), with_graph(), with_vertex(), without_attr(), without_multiples()

Examples

# An artificial example
make_(full_graph(5, loops = TRUE))
make_(full_graph(5, loops = TRUE), without_loops())
without_multiples  Constructor modifier to drop multiple edges

Description
Constructor modifier to drop multiple edges

Usage
without_multiples()

See Also
Other constructor modifiers: simplified(), with_edge(), with_graph(), with_vertex(), without_attr(), without_loops()

Examples
sample_(pa(10, m = 3, algorithm = "bag"))
sample_(pa(10, m = 3, algorithm = "bag"), without_multiples())

write_graph  Writing the graph to a file in some format

Description
write_graph is a general function for exporting graphs to foreign file formats, however not many formats are implemented right now.

Usage
write_graph(
  graph,
  file,
  format = c("edgelist", "pajek", "ncol", "lgl", "graphml", "dimacs", "gml", "dot", "leda"),
  ...
)

Arguments
  graph  The graph to export.
  file   A connection or a string giving the file name to write the graph to.
  format Character string giving the file format. Right now pajek, graphml, dot, gml, edgelist, lgl, ncol and dimacs are implemented. As of igraph 0.4 this argument is case insensitive.
  ...    Other, format specific arguments, see below.
write_graph

Value

A NULL, invisibly.

Edge list format

The edgelist format is a simple text file, with one edge in a line, the two vertex ids separated by a space character. The file is sorted by the first and the second column. This format has no additional arguments.

Author(s)

Gabor Csardi <csardi.gabor@gmail.com>

References


See Also

read_graph

Examples

g <- make_ring(10)
## Not run: write_graph(g, "/tmp/g.txt", "edgelist")
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