# igraph library

# API Documentation

# October 8, 2020

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# 1 Package igraph

IGraph library.

Version: 0.8.3

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#### 1.1 Modules

- \_igraph: Low-level Python interface for the igraph library. (Section 2, p. 226)
- app: User interfaces of igraph

(Section 3, p. 231)

- shell: Command-line user interface of igraph (Section 4, p. 232)
- clustering: Classes related to graph clustering.

(Section 5, p. 241)

- configuration: Configuration framework for igraph. (Section 6, p. 269)
- cut: Classes representing cuts and flows on graphs. (Section 7, p. 274)
- datatypes: Additional auxiliary data types (Section 8, p. 281)
- drawing: Drawing and plotting routines for IGraph.

(Section 9, p. 292)

- baseclasses: Abstract base classes for the drawing routines.
  - (Section 10, p. 312)
- colors: Color handling functions.

(Section 11, p. 315)

- coord: Coordinate systems and related plotting routines (Section 12, p. 328)
- edge: Drawers for various edge styles in graph plots.
   (Section 13, p. 331)
- **graph**: Drawing routines to draw graphs. (Section 14, p. 339)
- metamagic: Auxiliary classes for the default graph drawer in igraph.
   (Section 15, p. 346)
- shapes: Shape drawing classes for igraph (Section 16, p. 349)
- text: Drawers for labels on plots.

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```
(Section 17, p. 352)
```

- utils: Utility classes for drawing routines.

(Section 18, p. 357)

vertex: Drawing routines to draw the vertices of graphs.

(Section 19, p. 368)

- formula (Section ??, p. ??)
- layout: Layout-related code in the IGraph library. (Section 20, p. 373)
- matching: Classes representing matchings on graphs. (Section 21, p. 380)
- operators: Implementation of union, disjoint union and intersection operators. (Section 22, p. 383)
- remote: Classes that help igraph communicate with remote applications.

(Section 23, p. 386)

- gephi: Classes that help igraph communicate with Gephi (http://www.gephi.org).
   (Section 24, p. 387)
- statistics: Statistics related stuff in igraph (Section 25, p. 394)
- summary (Section 1.2, p. 4)
- summary': Summary representation of a graph. (Section 26, p. 405)
- utils: Utility functions that cannot be categorised anywhere else. (Section 27, p. 408)
- version (Section 28, p. 415)

#### 1.2 Functions

## autocurve(graph, attribute='curved', default=0)

Calculates curvature values for each of the edges in the graph to make sure that multiple edges are shown properly on a graph plot.

This function checks the multiplicity of each edge in the graph and assigns curvature values (numbers between -1 and 1, corresponding to CCW (-1), straight (0) and CW (1) curved edges) to them. The assigned values are either stored in an edge attribute or returned as a list, depending on the value of the *attribute* argument.

#### **Parameters**

graph: the graph on which the calculation will be run

attribute: the name of the edge attribute to save the curvature values to. The

default value is curved, which is the name of the edge attribute the default graph plotter checks to decide whether an edge should be curved on the plot or not. If *attribute* is None, the result will not be

stored.

default: the default curvature for single edges. Zero means that single edges

will be straight. If you want single edges to be curved as well, try

passing 0.5 or -0.5 here.

#### Return Value

the list of curvature values if attribute is None, otherwise None.

Functions Package igraph

#### get include()

Returns the folder that contains the C API headers of the Python interface of igraph.

read(filename, \*args, \*\*kwds)

Loads a graph from the given filename.

This is just a convenience function, calls Graph.Read directly. All arguments are passed unchanged to Graph.Read

#### **Parameters**

filename: the name of the file to be loaded

# load(filename, \*args, \*\*kwds)

Loads a graph from the given filename.

This is just a convenience function, calls Graph.Read directly. All arguments are passed unchanged to Graph.Read

#### **Parameters**

filename: the name of the file to be loaded

## write(graph, filename, \*args, \*\*kwds)

Saves a graph to the given file.

This is just a convenience function, calls Graph.write directly. All arguments are passed unchanged to Graph.write

#### **Parameters**

graph: the graph to be saved

filename: the name of the file to be written

## **save**(graph, filename, \*args, \*\*kwds)

Saves a graph to the given file.

This is just a convenience function, calls Graph.write directly. All arguments are passed unchanged to Graph.write

#### **Parameters**

graph: the graph to be saved

filename: the name of the file to be written

## summary(obj, stream=None, \*args, \*\*kwds)

Prints a summary of object o to a given stream

Positional and keyword arguments not explicitly mentioned here are passed on to the underlying summary() method of the object if it has any.

#### **Parameters**

obj: the object about which a human-readable summary is requested.

stream: the stream to be used. If None, the standard output will be used.

# 1.3 Variables

Variables Name	Description
config	Value: None
ADJ DIRECTED	Value: 0
ADJ LOWER	Value: 3
ADJ MAX	Value: 1
ADJ MIN	Value: 4
ADJ PLUS	Value: 5
ADJ UNDIRECTED	Value: 1
ADJ UPPER	Value: 2
ALL	Value: 3
BLISS F	Value: 0
BLISS_FL	Value: 1
BLISS_FLM	Value: 4
BLISS_FM	Value: 3
BLISS_FS	Value: 2
BLISS_FSM	Value: 5
GET_ADJACENCY_BOTH	Value: 2
GET_ADJACENCY_LOWE-	Value: 1
R	
GET_ADJACENCY_UPPER	Value: 0
IN	Value: 2
OUT	Value: 1
REWIRING_SIMPLE	Value: 0
REWIRING_SIMPLE_LOOP-	Value: 1
S	
STAR_IN	Value: 1
STAR_MUTUAL	Value: 3
STAR_OUT	Value: 0
STAR_UNDIRECTED	Value: 2
STRONG	Value: 2
TRANSITIVITY_NAN	Value: 0
TRANSITIVITY_ZERO	Value: 1
TREE_IN	Value: 1
TREE_OUT	Value: 0
TREE_UNDIRECTED	Value: 2
WEAK	Value: 1
package	Value: 'igraph'
arpack_options	Value: <igraph.arpackoptions 0x10fde9b90="" at="" object=""></igraph.arpackoptions>
ma ma a	
name	Value: 'write_svg'

# 1.4 Class Vertex

object — igraph.Vertex

Class representing a single vertex in a graph.

The vertex is referenced by its index, so if the underlying graph changes, the semantics of the vertex object

might change as well (if the vertex indices are altered in the original graph).

The attributes of the vertex can be accessed by using the vertex as a hash:

## 1.4.1 Methods

 $\frac{\_\_delitem\_\_(x, y)}{\text{del x[y]}}$ 

 $\underbrace{\frac{-\mathbf{eq}_{-}(x, y)}{\mathbf{x} == \mathbf{y}}}$ 

 $\frac{- ge_{-}(x, y)}{x >= y}$ 

 $\frac{\text{\__getitem}_{\_(x, y)}}{\mathbf{x}[\mathbf{y}]}$ 

 $\begin{bmatrix} \underline{-gt}\underline{-}(x, y) \\ x > y \end{bmatrix}$ 

 $\frac{-\mathbf{le}_{-}(x, y)}{x <= y}$ 

 $\frac{-\operatorname{len}_{-}(x)}{\operatorname{len}(x)}$ 

 $\frac{-\mathbf{lt}_{-}(x, y)}{x < y}$ 

 $\begin{bmatrix} -\mathbf{ne}_{-}(x, y) \\ x! = y \end{bmatrix}$ 

```
repr__(x)

repr(x)

Overrides: object.__repr__
```

```
\frac{\__{\text{setitem}}_{\_(x, i, y)}}{x[i]=y}
```

```
all edges(...)
```

Proxy method to Graph.incident(..., mode="all")

This method calls the incident() method of the Graph class with this vertex as the first argument and "all" as the mode argument, and returns the result.

See Also: Graph.incident() for details.

# attribute names()

Returns the list of vertex attribute names

#### Return Value

list

#### attributes()

Returns a dict of attribute names and values for the vertex

#### Return Value

dict

## betweenness(...)

Proxy method to Graph.betweenness()

This method calls the betweenness method of the **Graph** class with this vertex as the first argument, and returns the result.

See Also: Graph.betweenness() for details.

#### closeness(...)

Proxy method to Graph.closeness()

This method calls the closeness method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.closeness() for details.

#### constraint(...)

Proxy method to Graph.constraint()

This method calls the constraint method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.constraint() for details.

## degree(...)

Proxy method to Graph.degree()

This method calls the degree method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.degree() for details.

#### delete(...)

Proxy method to Graph.delete\_vertices()

This method calls the delete\_vertices method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.delete vertices() for details.

#### diversity(...)

Proxy method to Graph.diversity()

This method calls the diversity method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.diversity() for details.

# eccentricity(...)

Proxy method to Graph.eccentricity()

This method calls the eccentricity method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.eccentricity() for details.

# ${\tt get\_shortest\_paths}(...)$

Proxy method to Graph.get\_shortest\_paths()

This method calls the get\_shortest\_paths method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.get shortest paths() for details.

#### in edges(...)

Proxy method to Graph.incident(..., mode="in")

This method calls the incident() method of the Graph class with this vertex as the first argument and "in" as the mode argument, and returns the result.

See Also: Graph.incident() for details.

## incident(...)

Proxy method to Graph.incident()

This method calls the incident method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.incident() for details.

#### indegree(...)

Proxy method to Graph.indegree()

This method calls the indegree method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.indegree() for details.

#### is minimal separator(...)

Proxy method to Graph.is\_minimal\_separator()

This method calls the is\_minimal\_separator method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.is minimal separator() for details.

# $\mathbf{is}_{-}\mathbf{separator}(...)$

Proxy method to Graph.is\_separator()

This method calls the is\_separator method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.is separator() for details.

#### neighbors(...)

Proxy method to Graph.neighbors()

This method calls the neighbors method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.neighbors() for details.

#### out edges(...)

Proxy method to Graph.incident(..., mode="out")

This method calls the incident() method of the Graph class with this vertex as the first argument and "out" as the mode argument, and returns the result.

See Also: Graph.incident() for details.

# outdegree(...)

Proxy method to Graph.outdegree()

This method calls the outdegree method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.outdegree() for details.

#### pagerank(...)

Proxy method to Graph.pagerank()

This method calls the pagerank method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.pagerank() for details.

#### personalized pagerank(...)

Proxy method to Graph.personalized\_pagerank()

This method calls the personalized\_pagerank method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.personalized pagerank() for details.

#### predecessors(...)

Proxy method to Graph.predecessors()

This method calls the predecessors method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.predecessors() for details.

#### shortest paths(...)

Proxy method to Graph.shortest\_paths()

This method calls the shortest\_paths method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.shortest paths() for details.

## strength(...)

Proxy method to Graph.strength()

This method calls the strength method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.strength() for details.

## successors(...)

Proxy method to Graph.successors()

This method calls the successors method of the Graph class with this vertex as the first argument, and returns the result.

See Also: Graph.successors() for details.

# update attributes (E, \*\*F)

Updates the attributes of the vertex from dict/iterable E and F.

If E has a keys() method, it does: for k in E: self[k] = E[k]. If E lacks a keys() method, it does: for (k, v) in E: self[k] = v. In either case, this is followed by: for k in F: self[k] = F[k].

This method thus behaves similarly to the update() method of Python dictionaries.

## Return Value

None

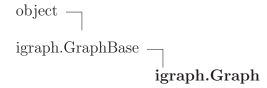
# Inherited from object

delattr(), _	$\_format\_\_$	(),ge	tattribute	(), _	$\_\_\mathrm{init}\_$	_(), _	_new_	()
reduce(), _	_reduce_ex	(), _	_setattr_	_(),	_sizeof_	_(), _	_str_	_(),
subclasshook	()							

#### 1.4.2 Properties

Name	Description
graph	The graph the vertex belongs to
index	Index of the vertex
Inherited from object	
class	

# 1.5 Class Graph



# Generic graph.

This class is built on top of GraphBase, so the order of the methods in the Epydoc documentation is a little bit obscure: inherited methods come after the ones implemented directly in the subclass. Graph provides many functions that GraphBase does not, mostly because these functions are not speed critical and they were easier to implement in Python than in pure C. An example is the attribute handling in the constructor: the constructor of Graph accepts three dictionaries corresponding to the graph, vertex and edge attributes while the constructor of GraphBase does not. This extension was needed to make Graph serializable through the pickle module. Graph also overrides some functions from GraphBase to provide a more convenient interface; e.g., layout functions return a Layout instance from Graph instead of a list of coordinate pairs.

Graphs can also be indexed by strings or pairs of vertex indices or vertex names. When a graph is indexed by a string, the operation translates to the retrieval, creation, modification or deletion of a graph attribute:

```
>>> g = Graph.Full(3)
>>> g["name"] = "Triangle graph"
>>> g["name"]
'Triangle graph'
>>> del g["name"]
```

When a graph is indexed by a pair of vertex indices or names, the graph itself is treated as an adjacency matrix and the corresponding cell of the matrix is returned:

```
>>> g = Graph.Full(3)
>>> g.vs["name"] = ["A", "B", "C"]
>>> g[1, 2]
1
>>> g["A", "B"]
1
>>> g["A", "B"] = 0
>>> g.ecount()
2
```

Assigning values different from zero or one to the adjacency matrix will be translated to one, unless the graph is weighted, in which case the numbers will be treated as weights:

```
>>> g.is_weighted()
False
>>> g["A", "B"] = 2
>>> g["A", "B"]
1
>>> g.es["weight"] = 1.0
>>> g.is_weighted()
True
```

```
>>> g["A", "B"] = 2
>>> g["A", "B"]
2
>>> g.es["weight"]
[1.0, 1.0, 2]
```

#### 1.5.1 Methods

# omega()

Returns the clique number of the graph.

The clique number of the graph is the size of the largest clique.

See Also: largest\_cliques() for the largest cliques.

# alpha()

Returns the independence number of the graph.

The independence number of the graph is the size of the largest independent vertex set.

See Also: largest\_independent\_vertex\_sets() for the largest independent vertex sets

# shell index(mode=ALL)

Finds the coreness (shell index) of the vertices of the network.

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

## **Parameters**

mode: whether to compute the in-corenesses (IN), the out-corenesses (OUT) or the undirected corenesses (ALL). Ignored and assumed to be ALL for undirected graphs.

## Return Value

the corenesses for each vertex.

**Reference:** Vladimir Batagelj, Matjaz Zaversnik: An O(m) Algorithm for Core Decomposition of Networks.

# cut vertices()

Returns the list of articulation points in the graph.

A vertex is an articulation point if its removal increases the number of connected components in the graph.

Calculates the eigenvector centralities of the vertices in a graph.

Eigenvector centrality is a measure of the importance of a node in a network. It assigns relative scores to all nodes in the network based on the principle that connections from high-scoring nodes contribute more to the score of the node in question than equal connections from low-scoring nodes. In practice, the centralities are determined by calculating eigenvector corresponding to the largest positive eigenvalue of the adjacency matrix. In the undirected case, this function considers the diagonal entries of the adjacency matrix to be twice the number of self-loops on the corresponding vertex.

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.

#### **Parameters**

directed: whether to consider edge directions in a

directed graph. Ignored for undirected graphs.

scale: whether to normalize the centralities so the

largest one will always be 1.

weights: edge weights given as a list or an edge

attribute. If None, all edges have equal weight.

return\_eigenvalue: whether to return the actual largest

eigenvalue along with the centralities

arpack\_options: an ARPACKOptions object that can be used to

fine-tune the calculation. If it is omitted, the module-level variable called arpack\_options

is used.

## Return Value

the eigenvector centralities in a list and optionally the largest eigenvalue (as a second member of a tuple)

vertex\_disjoint\_paths(source=-1, target=-1, checks=True,
neighbors="error")

Calculates the vertex connectivity of the graph or between some vertices.

The vertex connectivity between two given vertices is the number of vertices that have to be removed in order to disconnect the two vertices into two separate components. This is also the number of vertex disjoint directed paths between the vertices (apart from the source and target vertices of course). The vertex connectivity of the graph is the minimal vertex connectivity over all vertex pairs.

This method calculates the vertex connectivity of a given vertex pair if both the source and target vertices are given. If none of them is given (or they are both negative), the overall vertex connectivity is returned.

# **Parameters**

source: the source vertex involved in the calculation.

target: the target vertex involved in the calculation.

checks: if the whole graph connectivity is calculated and this is

True, igraph performs some basic checks before

calculation. If the graph is not strongly connected, then the connectivity is obviously zero. If the minimum degree is one, then the connectivity is also one. These simple checks are much faster than checking the entire graph, therefore it is advised to set this to True. The parameter is ignored if the connectivity between two

given vertices is computed.

neighbors: tells igraph what to do when the two vertices are

connected. "error" raises an exception, "infinity"

returns infinity, "ignore" ignores the edge.

#### Return Value

the vertex connectivity

# edge disjoint paths(source=-1, target=-1, checks=True)

Calculates the edge connectivity of the graph or between some vertices.

The edge connectivity between two given vertices is the number of edges that have to be removed in order to disconnect the two vertices into two separate components. This is also the number of edge disjoint directed paths between the vertices. The edge connectivity of the graph is the minimal edge connectivity over all vertex pairs.

This method calculates the edge connectivity of a given vertex pair if both the source and target vertices are given. If none of them is given (or they are both negative), the overall edge connectivity is returned.

#### Parameters

source: the source vertex involved in the calculation.

target: the target vertex involved in the calculation.

checks: if the whole graph connectivity is calculated and this is

True, igraph performs some basic checks before calculation.

If the graph is not strongly connected, then the

connectivity is obviously zero. If the minimum degree is one, then the connectivity is also one. These simple checks are much faster than checking the entire graph, therefore it is advised to set this to True. The parameter is ignored if the connectivity between two given vertices is computed.

## Return Value

the edge connectivity

**cohesion**(source=-1, target=-1, checks=True, neighbors="error")

Calculates the vertex connectivity of the graph or between some vertices.

The vertex connectivity between two given vertices is the number of vertices that have to be removed in order to disconnect the two vertices into two separate components. This is also the number of vertex disjoint directed paths between the vertices (apart from the source and target vertices of course). The vertex connectivity of the graph is the minimal vertex connectivity over all vertex pairs.

This method calculates the vertex connectivity of a given vertex pair if both the source and target vertices are given. If none of them is given (or they are both negative), the overall vertex connectivity is returned.

#### **Parameters**

source: the source vertex involved in the calculation.

target: the target vertex involved in the calculation.

checks: if the whole graph connectivity is calculated and this is

True, igraph performs some basic checks before

calculation. If the graph is not strongly connected, then the connectivity is obviously zero. If the minimum degree is one, then the connectivity is also one. These simple checks are much faster than checking the entire graph, therefore it is advised to set this to True. The

given vertices is computed.

neighbors: tells igraph what to do when the two vertices are

connected. "error" raises an exception, "infinity"

parameter is ignored if the connectivity between two

returns infinity, "ignore" ignores the edge.

## Return Value

the vertex connectivity

# adhesion(source=-1, target=-1, checks=True)

Calculates the edge connectivity of the graph or between some vertices.

The edge connectivity between two given vertices is the number of edges that have to be removed in order to disconnect the two vertices into two separate components. This is also the number of edge disjoint directed paths between the vertices. The edge connectivity of the graph is the minimal edge connectivity over all vertex pairs.

This method calculates the edge connectivity of a given vertex pair if both the source and target vertices are given. If none of them is given (or they are both negative), the overall edge connectivity is returned.

# Parameters

source: the source vertex involved in the calculation.

target: the target vertex involved in the calculation.

checks: if the whole graph connectivity is calculated and this is

True, igraph performs some basic checks before calculation.

If the graph is not strongly connected, then the

connectivity is obviously zero. If the minimum degree is one, then the connectivity is also one. These simple checks are much faster than checking the entire graph, therefore it is advised to set this to True. The parameter is ignored if the connectivity between two given vertices is computed.

## Return Value

the edge connectivity

 $shortest\_paths\_dijkstra(source=None, target=None, weights=None, mode=OUT)$ 

Calculates shortest path lengths for given vertices in a graph.

The algorithm used for the calculations is selected automatically: a simple BFS is used for unweighted graphs, Dijkstra's algorithm is used when all the weights are positive. Otherwise, the Bellman-Ford algorithm is used if the number of requested source vertices is larger than 100 and Johnson's algorithm is used otherwise.

#### **Parameters**

source: a list containing the source vertex IDs which should be

included in the result. If None, all vertices will be

considered.

target: a list containing the target vertex IDs which should be

included in the result. If None, all vertices will be

considered.

weights: a list containing the edge weights. It can also be an

attribute name (edge weights are retrieved from the given

attribute) or None (all edges have equal weight).

mode: the type of shortest paths to be used for the calculation in

directed graphs. OUT means only outgoing, IN means only incoming paths. ALL means to consider the directed graph

as an undirected one.

# Return Value

the shortest path lengths for given vertices in a matrix

# subgraph(vertices, implementation="auto")

Returns a subgraph spanned by the given vertices.

## **Parameters**

vertices: a list containing the vertex IDs which should be

included in the result.

implementation: the implementation to use when constructing the

new subgraph. igraph includes two implementations at the moment.

"copy\_and\_delete" copies the original graph and removes those vertices that are not in the given set. This is more efficient if the size of the

subgraph is comparable to the original graph. The other implementation ("create\_from\_scratch") constructs the result graph from scratch and then copies the attributes accordingly. This is a better solution if the subgraph is relatively small, compared to the original graph. "auto" selects

between the two implementations automatically, based on the ratio of the size of the subgraph and

the size of the original graph.

## Return Value

the subgraph

<u>\_\_\_init\_\_\_</u>(n=0, edges=None, directed=False, graph\_attrs=None, vertex attrs=None, edge attrs=None)

Constructs a graph from scratch.

#### **Parameters**

n: the number of vertices. Can be omitted, the default

is zero. Note that if the edge list contains vertices with indexes larger than or equal to m, then the number of vertices will be adjusted accordingly.

edges: the edge list where every list item is a pair of

integers. If any of the integers is larger than n-1, the number of vertices is adjusted accordingly. None

means no edges.

directed: whether the graph should be directed

graph\_attrs: the attributes of the graph as a dictionary.

vertex\_attrs: the attributes of the vertices as a dictionary. Every

dictionary value must be an iterable with exactly n

items.

edge\_attrs: the attributes of the edges as a dictionary. Every

dictionary value must be an iterable with exactly m

items where m is the number of edges.

Overrides: object. init

# add edge(source, target, \*\*kwds)

Adds a single edge to the graph.

Keyword arguments (except the source and target arguments) will be assigned to the edge as attributes.

#### **Parameters**

source: the source vertex of the edge or its name.

target: the target vertex of the edge or its name.

#### Return Value

the newly added edge as an Edge object. Use add\_edges([(source, target)]) if you don't need the Edge object and want to avoid the overhead of creating t.

# add edges(es)

Adds some edges to the graph.

#### Parameters

es: the list of edges to be added. Every edge is represented

with a tuple containing the vertex IDs or names of the two endpoints. Vertices are enumerated from zero.

attributes: dict of sequences, all of length equal to the number of

edges to be added, containing the attributes of the

new edges.

Overrides: igraph.GraphBase.add edges

# add vertex(name=None, \*\*kwds)

Adds a single vertex to the graph. Keyword arguments will be assigned as vertex attributes. Note that name as a keyword argument is treated specially; if a graph has name as a vertex attribute, it allows one to refer to vertices by their names in most places where igraph expects a vertex ID.

#### Return Value

the newly added vertex as a Vertex object. Use add\_vertices(1) if you don't need the Vertex object and want to avoid the overhead of creating t.

# add vertices(n, attributes=None)

Adds some vertices to the graph.

Note that if n is a sequence of strings, indicating the names of the new vertices, and attributes has a key name, the two conflict. In that case the attribute will be applied.

#### **Parameters**

n: the number of vertices to be added, or the name of a

single vertex to be added, or a sequence of strings, each corresponding to the name of a vertex to be added. Names will be assigned to the name vertex attribute.

attributes: dict of sequences, all of length equal to the number of

vertices to be added, containing the attributes of the new vertices. If n is a string (so a single vertex is added), then the values of this dict are the attributes themselves, but if n=1 then they have to be lists of

length 1.

Overrides: igraph.GraphBase.add vertices

# adjacent(vertex, mode=0UT)

Returns the edges a given vertex is incident on.

Deprecated: replaced by Graph.incident() since igraph 0.6

# as directed(\*args, \*\*kwds)

Returns a directed copy of this graph. Arguments are passed on to Graph.to\_directed() that is invoked on the copy.

# as undirected(\*args, \*\*kwds)

Returns an undirected copy of this graph. Arguments are passed on to Graph.to\_undirected() that is invoked on the copy.

# **delete** edges(self, \*args, \*\*kwds)

Deletes some edges from the graph.

The set of edges to be deleted is determined by the positional and keyword arguments. If the function is called without any arguments, all edges are deleted. If any keyword argument is present, or the first positional argument is callable, an edge sequence is derived by calling EdgeSeq.select with the same positional and keyword arguments. Edges in the derived edge sequence will be removed. Otherwise the first positional argument is considered as follows:

- None deletes all edges (deprecated since 0.8.3)
- a single integer deletes the edge with the given ID
- a list of integers deletes the edges denoted by the given IDs
- a list of 2-tuples deletes the edges denoted by the given source-target vertex pairs. When multiple edges are present between a given source-target vertex pair, only one is removed.

#### **Parameters**

es: the list of edges to be removed. Edges are identifed by edge IDs. EdgeSeq objects are also accepted here. No argument deletes all edges.

Overrides: igraph.GraphBase.delete edges

**Deprecated:** Graph.delete\_edges(None) has been replaced by Graph.delete\_edges() - with no arguments - since igraph 0.8.3.

indegree(self, \*args, \*\*kwds)

Returns the in-degrees in a list.

See degree for possible arguments.

outdegree(self, \*args, \*\*kwds)

Returns the out-degrees in a list.

See degree for possible arguments.

# all st cuts(self, source, target)

Returns all the cuts between the source and target vertices in a directed graph.

This function lists all edge-cuts between a source and a target vertex. Every cut is listed exactly once.

## **Parameters**

source: the source vertex ID
target: the target vertex ID

# Return Value

a list of Cut objects.

 $Overrides: igraph.GraphBase.all\_st\_cuts$ 

Reference: JS Provan and DR Shier: A paradigm for listing (s,t)-cuts in

graphs. Algorithmica 15, 351–372, 1996.

# all st mincuts(self, source, target, capacity=None)

Returns all the mincuts between the source and target vertices in a directed graph.

This function lists all minimum edge-cuts between a source and a target vertex.

#### Parameters

source: the source vertex ID

target: the target vertex ID

capacity: the edge capacities (weights). If None, all edges have

equal weight. May also be an attribute name.

## Return Value

a list of Cut objects.

Overrides: igraph.GraphBase.all st mincuts

Reference: JS Provan and DR Shier: A paradigm for listing (s,t)-cuts in

graphs. Algorithmica 15, 351–372, 1996.

# **biconnected** components(self, return articulation points=False)

Calculates the biconnected components of the graph.

#### **Parameters**

return\_articulation\_points: whether to return the articulation points as well

## Return Value

a VertexCover object describing the biconnected components, and optionally the list of articulation points as well

Overrides: igraph.GraphBase.biconnected components

# blocks(self, return\_articulation\_points=False)

Calculates the biconnected components of the graph.

#### **Parameters**

return\_articulation\_points: whether to return the articulation points as well

#### Return Value

a VertexCover object describing the biconnected components, and optionally the list of articulation points as well

# clear()

Clears the graph, deleting all vertices, edges, and attributes.

See Also: Graph.delete\_vertices and Graph.delete\_edges.

# cohesive blocks()

Calculates the cohesive block structure of the graph.

Cohesive blocking is a method of determining hierarchical subsets of graph vertices based on their structural cohesion (i.e. vertex connectivity). For a given graph G, a subset of its vertices S 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 obtained in the end, with the entire graph G at its root.

#### Return Value

an instance of CohesiveBlocks. See the documentation of CohesiveBlocks for more information.

Overrides: igraph.GraphBase.cohesive blocks

See Also: CohesiveBlocks

# clusters(mode = STRONG)

Calculates the (strong or weak) clusters (connected components) for a given graph.

#### **Parameters**

mode: must be either STRONG or WEAK, depending on the clusters being sought. Optional, defaults to STRONG.

## Return Value

a VertexClustering object

Overrides: igraph.GraphBase.clusters

# components(mode = STRONG)

Calculates the (strong or weak) clusters (connected components) for a given graph.

# **Parameters**

mode: must be either STRONG or WEAK, depending on the clusters being sought. Optional, defaults to STRONG.

# Return Value

a VertexClustering object

# **degree** distribution(bin\_width=1, ...)

Calculates the degree distribution of the graph.

Unknown keyword arguments are directly passed to degree().

#### **Parameters**

bin\_width: the bin width of the histogram

# Return Value

a histogram representing the degree distribution of the graph.

# dyad census()

Calculates the dyad census of the graph.

Dyad census means classifying each pair of vertices of a directed graph into three categories: mutual (there is an edge from a to b and also from b to a), asymmetric (there is an edge from a to b or from b to a but not the other way round) and null (there is no connection between a and b).

## Return Value

a DyadCensus object.

Overrides: igraph.GraphBase.dyad census

Reference: Holland, P.W. and Leinhardt, S. (1970). A Method for Detecting Structure in Sociometric Data. American Journal of Sociology, 70, 492-513.

# get adjacency(self, type=2, attribute=None, default=0, eids=False)

Returns the adjacency matrix of a graph.

#### **Parameters**

type: either GET\_ADJACENCY\_LOWER (uses the lower triangle of

the matrix) or GET\_ADJACENCY\_UPPER (uses the upper triangle) or GET\_ADJACENCY\_BOTH (uses both parts).

Ignored for directed graphs.

attribute: if None, returns the ordinary adjacency matrix. When

the name of a valid edge attribute is given here, the matrix returned will contain the default value at the places where there is no edge or the value of the given attribute where there is an edge. Multiple edges are not supported, the value written in the matrix in this case will be unpredictable. This parameter is ignored if *eids* 

is True

default: the default value written to the cells in the case of

adjacency matrices with attributes.

eids: specifies whether the edge IDs should be returned in the

adjacency matrix. Since zero is a valid edge ID, the cells in the matrix that correspond to unconnected vertex pairs will contain -1 instead of 0 if *eids* is True. If *eids* is False, the number of edges will be returned in

the matrix for each vertex pair.

#### Return Value

the adjacency matrix as a Matrix.

Overrides: igraph.GraphBase.get adjacency

## get adjacency sparse(self, attribute=None)

Returns the adjacency matrix of a graph as a SciPy CSR matrix.

## **Parameters**

attribute: if None, returns the ordinary adjacency matrix. When

the name of a valid edge attribute is given here, the matrix returned will contain the default value at the places where there is no edge or the value of the given

attribute where there is an edge.

#### Return Value

the adjacency matrix as a scipy.sparse.csr\_matrix.

# get adjlist(mode=OUT)

Returns the adjacency list representation of the graph.

The adjacency list representation is a list of lists. Each item of the outer list belongs to a single vertex of the graph. The inner list contains the neighbors of the given vertex.

## Parameters

mode: if OUT, returns the successors of the vertex. If IN, returns the predecessors of the vertex. If ALL, both the predecessors and the successors will be returned. Ignored for undirected graphs.

# get adjedgelist(mode=OUT)

Returns the incidence list representation of the graph.

Deprecated: replaced by Graph.get\_inclist() since igraph 0.6

See Also: Graph.get inclist()

# get all simple paths(v, to=None, cutoff=-1, mode=OUT)

Calculates all the simple paths from a given node to some other nodes (or all of them) in a graph.

A path is simple if its vertices are unique, i.e. no vertex is visited more than once.

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

### **Parameters**

v: the source for the calculated paths

to: a vertex selector describing the destination for the calculated paths. This can be a single vertex ID, a list of vertex IDs, a single vertex name, a list of vertex names or a

VertexSeq object. None means all the vertices.

cutoff: maximum length of path that is considered. If negative,

paths of all lengths are considered.

mode: the directionality of the paths. IN means to calculate

incoming paths, OUT means to calculate outgoing paths,

ALL means to calculate both ones.

### Return Value

all of the simple paths from the given node to every other reachable node in the graph in a list. Note that in case of mode=IN, the vertices in a path are returned in reversed order!

# **get** inclist(mode=OUT)

Returns the incidence list representation of the graph.

The incidence list representation is a list of lists. Each item of the outer list belongs to a single vertex of the graph. The inner list contains the IDs of the incident edges of the given vertex.

### **Parameters**

mode: if OUT, returns the successors of the vertex. If IN, returns the predecessors of the vertex. If ALL, both the predecessors and the successors will be returned. Ignored for undirected graphs.

# gomory hu tree(capacity=None, flow="flow")

Calculates the Gomory-Hu tree of an undirected graph with optional edge capacities.

The Gomory-Hu tree is a concise representation of the value of all the maximum flows (or minimum cuts) in a graph. The vertices of the tree correspond exactly to the vertices of the original graph in the same order. Edges of the Gomory-Hu tree are annotated by flow values. The value of the maximum flow (or minimum cut) between an arbitrary (u,v) vertex pair in the original graph is then given by the minimum flow value (i.e. edge annotation) along the shortest path between u and v in the Gomory-Hu tree.

# **Parameters**

capacity: the edge capacities (weights). If None, all edges have

equal weight. May also be an attribute name.

flow: the name of the edge attribute in the returned graph in

which the flow values will be stored.

### Return Value

the Gomory-Hu tree as a Graph object.

Overrides: igraph.GraphBase.gomory\_hu\_tree

# is named()

Returns whether the graph is named, i.e., whether it has a "name" vertex attribute.

# is weighted()

Returns whether the graph is weighted, i.e., whether it has a "weight" edge attribute.

# maxflow(source, target, capacity=None)

Returns a maximum flow between the given source and target vertices in a graph.

A maximum 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 (i.e. the assigned number) is not more than the capacity of the edge (see the *capacity* argument)
- 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 or the outgoing flow of the source (which are equal). The maximum flow is the maximum possible such value.

# **Parameters**

capacity: the edge capacities (weights). If None, all edges have equal weight. May also be an attribute name.

### Return Value

a Flow object describing the maximum flow

Overrides: igraph.GraphBase.maxflow

# mincut(source=None, target=None, capacity=None)

Calculates the minimum cut between the given source and target vertices or within the whole graph.

The minimum cut is the minimum set of edges that needs to be removed to separate the source and the target (if they are given) or to disconnect the graph (if neither the source nor the target are given). The minimum is calculated using the weights (capacities) of the edges, so the cut with the minimum total capacity is calculated.

For undirected graphs and no source and target, the method uses the Stoer-Wagner algorithm. For a given source and target, the method uses the push-relabel algorithm; see the references below.

### **Parameters**

source: the source vertex ID. If None, the target must also be

None and the calculation will be done for the entire

graph (i.e. all possible vertex pairs).

target: the target vertex ID. If None, the source must also be

None and the calculation will be done for the entire

graph (i.e. all possible vertex pairs).

capacity: the edge capacities (weights). If None, all edges have

equal weight. May also be an attribute name.

# Return Value

a Cut object describing the minimum cut

Overrides: igraph.GraphBase.mincut

### st mincut(source, target, capacity=None)

Calculates the minimum cut between the source and target vertices in a graph.

#### **Parameters**

source: the source vertex ID
target: the target vertex ID

capacity: the capacity of the edges. It must be a list or a valid

attribute name or None. In the latter case, every edge

will have the same capacity.

### Return Value

the value of the minimum cut, the IDs of vertices in the first and second partition, and the IDs of edges in the cut, packed in a 4-tuple

Overrides: igraph.GraphBase.st mincut

# modularity(membership, weights=None)

Calculates the modularity score of the graph with respect to a given clustering.

The modularity of a graph w.r.t. some division measures how good the division is, or how separated are the different vertex types from each other. It's defined as Q=1/(2m)\*sum(Aij-ki\*kj/(2m)delta(ci,cj),i,j). m is the number of edges, Aij is the element of the A adjacency matrix in row i and column j, ki is the degree of node i, kj is the degree of node j, and Ci and cj are the types of the two vertices (i and j). delta(x,y) is one iff x=y, 0 otherwise.

If edge weights are given, the definition of modularity is modified as follows: Aij becomes the weight of the corresponding edge, ki is the total weight of edges adjacent to vertex i, kj is the total weight of edges adjacent to vertex j and m is the total edge weight in the graph.

### **Parameters**

membership: a membership list or a VertexClustering object

weights: optional edge weights or None if all edges are weighed

equally. Attribute names are also allowed.

### Return Value

the modularity score

Overrides: igraph.GraphBase.modularity

**Reference:** MEJ Newman and M Girvan: Finding and evaluating community structure in networks. Phys Rev E 69 026113, 2004.

# path length hist(directed=True)

Returns the path length histogram of the graph

#### Parameters

directed: whether to consider directed paths. Ignored for undirected graphs.

### Return Value

a Histogram object. The object will also have an unconnected attribute that stores the number of unconnected vertex pairs (where the second vertex can not be reached from the first one). The latter one will be of type long (and not a simple integer), since this can be very large.

Overrides: igraph.GraphBase.path length hist

pagerank(self, vertices=None, directed=True, damping=0.85, weights=None, arpack\_options=None, implementation='prpack', niter=1000, eps=0.001)

Calculates the Google PageRank values of a graph.

### **Parameters**

vertices: the indices of the vertices being queried. None

means all of the vertices.

directed: whether to consider directed paths.

damping: the damping factor. 1-damping is the PageRank

value for nodes with no incoming links. It is also the probability of resetting the random walk to a

uniform distribution in each step.

weights: edge weights to be used. Can be a sequence or

iterable or even an edge attribute name.

arpack\_options: an ARPACKOptions object used to fine-tune the

ARPACK eigenvector calculation. If omitted, the module-level variable called arpack\_options is used. This argument is ignored if not the ARPACK implementation is used, see the

implementation argument.

implementation: which implementation to use to solve the

PageRank eigenproblem. Possible values are:

• "prpack": use the PRPACK library. This is a new implementation in igraph 0.7

• "arpack": use the ARPACK library. This implementation was used from version 0.5, until version 0.7.

• "power": use a simple power method. This is the implementation that was used before igraph version 0.5.

niter: The number of iterations to use in the power

method implementation. It is ignored in the other

implementations

eps: The power method implementation will consider

the calculation as complete if the difference of PageRank values between iterations change less than this value for every node. It is ignored by

the other implementations.

# Return Value

a list with the Google PageRank values of the specified vertices.

# **spanning** tree(self, weights=None, return tree=True)

Calculates a minimum spanning tree for a graph.

### **Parameters**

weights: a vector containing weights for every edge in the

graph. None means that the graph is unweighted.

return\_tree: whether to return the minimum spanning tree (when

return\_tree is True) or to return the IDs of the edges in the minimum spanning tree instead (when return\_tree is False). The default is True for historical reasons as this argument was introduced in

igraph 0.6.

### Return Value

the spanning tree as a Graph object if return\_tree is True or the IDs of the edges that constitute the spanning tree if return\_tree is False.

**Reference:** Prim, R.C.: Shortest connection networks and some generalizations. Bell System Technical Journal 36:1389-1401, 1957.

# transitivity avglocal undirected(self, mode='nan', weights=None)

Calculates the average of the vertex transitivities of the graph.

In the unweighted case, the transitivity measures the probability that two neighbors of a vertex are connected. In case of the average local transitivity, this probability is calculated for each vertex and then the average is taken. Vertices with less than two neighbors require special treatment, they will either be left out from the calculation or they will be considered as having zero transitivity, depending on the *mode* parameter. The calculation is slightly more involved for weighted graphs; in this case, weights are taken into account according to the formula of Barrat et al (see the references).

Note that this measure is different from the global transitivity measure (see transitivity\_undirected()) as it simply takes the average local transitivity across the whole network.

### Parameters

mode:

defines how to treat vertices with degree less than two. If TRANSITIVITY\_ZERO or "zero", these vertices will have zero transitivity. If TRANSITIVITY\_NAN or "nan", these vertices will be excluded from the average.

weights: edge weights to be used. Can be a sequence or iterable or even an edge attribute name.

Overrides: igraph.GraphBase.transitivity avglocal undirected

See Also: transitivity\_undirected(), transitivity\_local\_undirected()

### Reference:

- Watts DJ and Strogatz S: Collective dynamics of small-world networks. Nature 393(6884):440-442, 1998.
- Barrat A, Barthelemy M, Pastor-Satorras R and Vespignani A: The architecture of complex weighted networks. PNAS 101, 3747 (2004). http://arxiv.org/abs/cond-mat/0311416.

# triad census()

Calculates the triad census of the graph.

# Return Value

a TriadCensus object.

Overrides: igraph.GraphBase.triad census

**Reference:** Davis, J.A. and Leinhardt, S. (1972). The Structure of Positive Interpersonal Relations in Small Groups. In: J. Berger (Ed.), Sociological Theories in Progress, Volume 2, 218-251. Boston: Houghton Mifflin.

 $count\_automorphisms\_vf2(self, color=None, edge\_color=None, node\_compat\_fn=None, edge\_compat\_fn=None)$ 

Returns the number of automorphisms of the graph.

This function simply calls count\_isomorphisms\_vf2 with the graph itself. See count\_isomorphisms\_vf2 for an explanation of the parameters.

### Return Value

the number of automorphisms of the graph

See Also: Graph.count isomorphisms vf2

```
\begin{tabular}{ll} {\bf get\_automorphisms\_vf2} (self,\ color={\tt None},\ edge\_color={\tt None},\ node\_compat\_fn={\tt None},\ edge\_compat\_fn={\tt None}) \\ \end{tabular}
```

Returns all the automorphisms of the graph

This function simply calls get\_isomorphisms\_vf2 with the graph itself. See get\_isomorphisms\_vf2 for an explanation of the parameters.

### Return Value

a list of lists, each item containing a possible mapping of the graph vertices to itself according to the automorphism

See Also: Graph.get\_isomorphisms\_vf2

# **community** fastgreedy(self, weights=None)

Community structure based on the greedy optimization of modularity.

This algorithm merges individual nodes into communities in a way that greedily maximizes the modularity score of the graph. It can be proven that if no merge can increase the current modularity score, the algorithm can be stopped since no further increase can be achieved.

This algorithm is said to run almost in linear time on sparse graphs.

#### Parameters

weights: edge attribute name or a list containing edge weights

#### Return Value

an appropriate VertexDendrogram object.

Overrides: igraph.GraphBase.community fastgreedy

**Reference:** A Clauset, MEJ Newman and C Moore: Finding community structure in very large networks. Phys Rev E 70, 066111 (2004).

 $community\_infomap(self, edge\_weights=None, vertex\_weights=None, trials=10)$ 

Finds the community structure of the network according to the Infomap method of Martin Rosvall and Carl T. Bergstrom.

### **Parameters**

edge\_weights: name of an edge attribute or a list containing

edge weights.

vertex\_weights: name of an vertex attribute or a list containing

vertex weights.

trials: the number of attempts to partition the network.

### Return Value

an appropriate VertexClustering object with an extra attribute called codelength that stores the code length determined by the algorithm.

Overrides: igraph.GraphBase.community\_infomap

#### Reference:

- M. Rosvall and C. T. Bergstrom: Maps of information flow reveal community structure in complex networks, PNAS 105, 1118 (2008). http://dx.doi.org/10.1073/pnas.0706851105, http://arxiv.org/abs/0707.0609.
- M. Rosvall, D. Axelsson, and C. T. Bergstrom: The map equation, Eur. Phys. J. Special Topics 178, 13 (2009). http://dx.doi.org/10.1140/epjst/e2010-01179-1, http://arxiv.org/abs/0906.1405.

# community\_leading\_eigenvector\_naive(clusters=None, return merges=False)

A naive implementation of Newman's eigenvector community structure detection. This function splits the network into two components according to the leading eigenvector of the modularity matrix and then recursively takes the given number of steps by splitting the communities as individual networks. This is not the correct way, however, see the reference for explanation. Consider using the correct community\_leading\_eigenvector method instead.

### **Parameters**

clusters: the desired number of communities. If None, the

algorithm tries to do as many splits as possible. Note that the algorithm won't split a community further if the signs of the leading eigenvector are all the same, so the actual number of discovered communities can be less than the desired one.

return\_merges: whether the returned object should be a

dendrogram instead of a single clustering.

### Return Value

an appropriate VertexClustering or VertexDendrogram object.

**Reference:** MEJ Newman: Finding community structure in networks using the eigenvectors of matrices, arXiv:physics/0605087

 $\begin{array}{lll} \textbf{community\_leading\_eigenvector}(\textit{clusters} = \texttt{None}, \textit{weights} = \texttt{None}, \\ \textit{arpack\_options} = \texttt{None}) \end{array}$ 

Newman's leading eigenvector method for detecting community structure. This is the proper implementation of the recursive, divisive algorithm: each split is done by maximizing the modularity regarding the original network.

# **Parameters**

clusters: the desired number of communities. If None, the

algorithm tries to do as many splits as possible. Note that the algorithm won't split a community further if the signs of the leading eigenvector are all the same, so the actual number of discovered communities can be less than the desired one.

weights: name of an edge attribute or a list containing

edge weights.

arpack\_options: an ARPACKOptions object used to fine-tune the

ARPACK eigenvector calculation. If omitted, the module-level variable called arpack\_options is

used.

### Return Value

an appropriate VertexClustering object.

Overrides: igraph.GraphBase.community leading eigenvector

Reference: MEJ Newman: Finding community structure in networks using

the eigenvectors of matrices, arXiv:physics/0605087

# community label propagation(weights=None, initial=None, fixed=None)

Finds the community structure of the graph according to the label propagation method of Raghavan et al. Initially, each vertex is assigned a different label. After that, each vertex chooses the dominant label in its neighbourhood in each iteration. Ties are broken randomly and the order in which the vertices are updated is randomized before every iteration. The algorithm ends when vertices reach a consensus. Note that since ties are broken randomly, there is no guarantee that the algorithm returns the same community structure after each run. In fact, they frequently differ. See the paper of Raghavan et al on how to come up with an aggregated community structure.

### **Parameters**

weights: name of an edge attribute or a list containing edge weights

initial: name of a vertex attribute or a list containing the initial

vertex labels. Labels are identified by integers from zero to n-1 where n is the number of vertices. Negative numbers may also be present in this vector, they represent

unlabeled vertices.

fixed: a list of booleans for each vertex. True corresponds to

vertices whose labeling should not change during the algorithm. It only makes sense if initial labels are also

given. Unlabeled vertices cannot be fixed.

### Return Value

an appropriate VertexClustering object.

Overrides: igraph.GraphBase.community label propagation

**Reference:** 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. http://arxiv.org/abs/0709.2938.

# **community** multilevel(self, weights=None, return levels=False)

Community structure based on the multilevel algorithm of Blondel et al.

This is a bottom-up algorithm: initially every vertex belongs to a separate community, and vertices are moved between communities iteratively in a way that maximizes the vertices' local contribution to the overall modularity score. When a consensus is reached (i.e. no single move would increase the modularity score), every community in the original graph is shrank to a single vertex (while keeping the total weight of the adjacent edges) and the process continues on the next level. The algorithm stops when it is not possible to increase the modularity any more after shrinking the communities to vertices.

This algorithm is said to run almost in linear time on sparse graphs.

#### Parameters

weights: edge attribute name or a list containing edge

weights

return\_levels: if True, the communities at each level are returned

in a list. If False, only the community structure

with the best modularity is returned.

### Return Value

a list of VertexClustering objects, one corresponding to each level (if return\_levels is True), or a VertexClustering corresponding to the best modularity.

Overrides: igraph.GraphBase.community multilevel

Reference: VD Blondel, J-L Guillaume, R Lambiotte and E Lefebvre: Fast unfolding of community hierarchies in large networks, J Stat Mech P10008 (2008), http://arxiv.org/abs/0803.0476

# community optimal modularity(self, \*args, \*\*kwds)

Calculates the optimal modularity score of the graph and the corresponding community structure.

This function uses the GNU Linear Programming Kit to solve a large integer optimization problem in order to find the optimal modularity score and the corresponding community structure, therefore it is unlikely to work for graphs larger than a few (less than a hundred) vertices. Consider using one of the heuristic approaches instead if you have such a large graph.

# **Parameters**

weights: name of an edge attribute or a list containing edge weights.

### Return Value

the calculated membership vector and the corresponding modularity in a tuple.

Overrides: igraph.GraphBase.community optimal modularity

community edge betweenness(self, clusters=None, directed=True, weights = None

Community structure based on the betweenness of the edges in the network.

The idea is that the betweenness of the edges connecting two communities is typically high, as many of the shortest paths between nodes in separate communities go through them. So we gradually remove the edge with the highest betweenness and recalculate the betweennesses after every removal. This way sooner or later the network falls of to separate components. The result of the clustering will be represented by a dendrogram.

### **Parameters**

clusters: the number of clusters we would like to see. This practically defines the "level" where we "cut" the dendrogram to get the membership vector of the vertices. If None, the dendrogram is cut at the level which maximizes the modularity when the graph is unweighted; otherwise the dendrogram is cut at at a single cluster (because cluster count selection based on modularities does not make sense for this method if not all the weights are equal).

directed: whether the directionality of the edges should be taken

into account or not.

name of an edge attribute or a list containing edge

weights.

# Return Value

a VertexDendrogram object, initally cut at the maximum modularity or at the desired number of clusters.

Overrides: igraph.GraphBase.community edge betweenness

 $\begin{array}{l} \textbf{community\_spinglass}(weights=\texttt{None},\ spins=\texttt{25},\ parupdate=\texttt{False},\\ start\_temp=\texttt{1},\ stop\_temp=\texttt{0.01},\ cool\_fact=\texttt{0.99},\ update\_rule=\texttt{"config"},\\ gamma=\texttt{1},\ implementation=\texttt{"orig"},\ lambda\_=\texttt{1}) \end{array}$ 

Finds the community structure of the graph according to the spinglass community detection method of Reichardt & Bornholdt.

### **Parameters**

weights: edge weights to be used. Can be a sequence or

iterable or even an edge attribute name.

spins: integer, 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: whether to update the spins of the vertices in

parallel (synchronously) or not

start\_temp: the starting temperature

stop\_temp: the stop temperature

cool\_fact: cooling factor for the simulated annealing

update\_rule: specifies the null model of the simulation.

Possible values are "config" (a random graph with the same vertex degrees as the input graph) or "simple" (a random graph with the same

number of edges)

gamma: the gamma argument of the algorithm, specifying

the balance between the importance of present and missing edges within a community. The default value of 1.0 assigns equal importance to

both of them.

implementation: currently igraph contains two implementations of

the spinglass community detection algorithm. The faster original implementation is the default. The other implementation is able to take into account negative weights, this can be chosen by

setting implementation to "neg"

lambda\_: the lambda argument of the algorithm, which

specifies the balance between the importance of present and missing negatively weighted edges within a community. Smaller values of lambda

lead to communities with less negative

intra-connectivity. If the argument is zero, the algorithm reduces to a graph coloring algorithm,

using the number of spins as colors. This

argument is ignored if the original

implementation is used. Note the underscore at the end of the argument name; this is due to the fact that lambda is a reserved keyword in Python.

# **community** walktrap(self, weights=None, steps=4)

Community detection algorithm of Latapy & Pons, based on random walks.

The basic idea of the algorithm is that short random walks tend to stay in the same community. The result of the clustering will be represented as a dendrogram.

### **Parameters**

weights: name of an edge attribute or a list containing edge weights

steps: length of random walks to perform

# Return Value

a VertexDendrogram object, initially cut at the maximum modularity.

Overrides: igraph.GraphBase.community walktrap

**Reference:** Pascal Pons, Matthieu Latapy: Computing communities in large networks using random walks, <a href="http://arxiv.org/abs/physics/0512106">http://arxiv.org/abs/physics/0512106</a>.

# k core(self, \*arqs)

Returns some k-cores of the graph.

The method accepts an arbitrary number of arguments representing the desired indices of the k-cores to be returned. The arguments can also be lists or tuples. The result is a single Graph object if an only integer argument was given, otherwise the result is a list of Graph objects representing the desired k-cores in the order the arguments were specified. If no argument is given, returns all k-cores in increasing order of k.

 $\begin{array}{l} \textbf{community\_leiden}(objective\_function=\texttt{CPM},\ weights=\texttt{None},\\ resolution\_parameter=\texttt{1.0},\ beta=\texttt{0.01},\ initial\_membership=\texttt{None},\\ n\_iterations=\texttt{2},\ node\_weights=\texttt{None}) \end{array}$ 

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

### Parameters

objective\_function: whether to use the Constant Potts Model

(CPM) or modularity. Must be either

"CPM" or "modularity".

weights: edge weights to be used. Can be a

sequence or iterable or even an edge

attribute name.

resolution\_parameter: the resolution parameter to use. Higher

resolutions lead to more smaller

communities, while lower resolutions lead

to fewer larger communities.

beta: parameter affecting the randomness in the

Leiden algorithm. This affects only the

refinement step of the algorithm.

initial\_membership: if provided, the Leiden algorithm will try

to improve this provided membership. If no argument is provided, the aglorithm simply starts from the singleton partition.

n\_iterations: the number of iterations to iterate the

Leiden algorithm. Each iteration may improve the partition further. Using a negative number of iterations will run until a stable iteration is encountered (i.e. the quality was not increased during that

iteration).

node\_weights: the node weights used in the Leiden

algorithm. If this is not provided, it will be automatically determined on the basis of whether you want to use CPM or modularity. If you do provide this, please make sure that you understand what you

are doing.

# Return Value

an appropriate VertexClustering object.

Overrides: igraph.GraphBase.community leiden

Reference: Traag, V. A., Waltman, L., & van Eck, N. J. (2019). From Louvain to Leiden: guaranteeing well-connected communities. Scientific

reports, 9(1), 5233. doi: 10.1038/s41598-019-41695-z

# layout(self, layout=None, \*args, \*\*kwds)

Returns the layout of the graph according to a layout algorithm.

Parameters and keyword arguments not specified here are passed to the layout algorithm directly. See the documentation of the layout algorithms for the explanation of these parameters.

Registered layout names understood by this method are:

- auto, automatic: automatic layout (see Graph.layout\_auto)
- bipartite: bipartite layout (see Graph.layout\_bipartite)
- circle, circular: circular layout (see Graph.layout\_circle)
- dh, davidson\_harel: Davidson-Harel layout (see Graph.layout\_davidson\_harel)
- drl: DrL layout for large graphs (see Graph.layout\_drl)
- drl\_3d: 3D DrL layout for large graphs (see Graph.layout\_drl)
- fr, fruchterman\_reingold: Fruchterman-Reingold layout (see Graph.layout\_fruchterman\_reingold).
- fr\_3d, fr3d, fruchterman\_reingold\_3d: 3D Fruchterman-Reingold layout (see Graph.layout\_fruchterman\_reingold).
- grid: regular grid layout in 2D (see Graph.layout\_grid)
- grid\_3d: regular grid layout in 3D (see Graph.layout\_grid\_3d)
- graphopt: the graphopt algorithm (see Graph.layout\_graphopt)
- kk, kamada\_kawai: Kamada-Kawai layout (see Graph.layout\_kamada\_kawai)
- kk\_3d, kk3d, kamada\_kawai\_3d: 3D Kamada-Kawai layout (see Graph.layout\_kamada\_kawai)
- lgl, large, large\_graph: Large Graph Layout (see Graph.layout\_lgl)
- mds: multidimensional scaling layout (see Graph.layout\_mds)
- random: random layout (see Graph.layout\_random)
- random\_3d: random 3D layout (see Graph.layout\_random)
- rt, tree, reingold\_tilford: Reingold-Tilford tree layout (see Graph.layout\_reingold\_tilford)
- rt\_circular, reingold\_tilford\_circular: circular Reingold-Tilford tree layout (see Graph.layout\_reingold\_tilford\_circular)
- sphere, spherical, circle\_3d, circular\_3d: spherical layout (see Graph.layout\_circle)
- star: star layout (see Graph.layout\_star)
- sugiyama: Sugiyama layout (see Graph.layout\_sugiyama)

### **Parameters**

layout: the layout to use. This can be one of the registered layout names or a callable which feturns either a Layout object or a list of lists containing the coordinates. If None, uses the value of the plotting.layout configuration key.

# Return Value

# layout auto(self, \*args, \*\*kwds)

Chooses and runs a suitable layout function based on simple topological properties of the graph.

This function tries to choose an appropriate layout function for the graph using the following rules:

- 1. If the graph has an attribute called layout, it will be used. It may either be a Layout instance, a list of coordinate pairs, the name of a layout function, or a callable function which generates the layout when called with the graph as a parameter.
- 2. Otherwise, if the graph has vertex attributes called x and y, these will be used as coordinates in the layout. When a 3D layout is requested (by setting dim to 3), a vertex attribute named z will also be needed.
- 3. Otherwise, if the graph is connected and has at most 100 vertices, the Kamada-Kawai layout will be used (see Graph.layout\_kamada\_kawai()).
- 4. Otherwise, if the graph has at most 1000 vertices, the Fruchterman-Reingold layout will be used (see Graph.layout\_fruchterman\_reingold()).
- 5. If everything else above failed, the DrL layout algorithm will be used (see Graph.layout\_drl()).

All the arguments of this function except dim are passed on to the chosen layout function (in case we have to call some layout function).

#### **Parameters**

dim: specifies whether we would like to obtain a 2D or a 3D layout.

# Return Value

a Layout object.

# layout grid fruchterman reingold(\*args, \*\*kwds)

Compatibility alias to the Fruchterman-Reingold layout with the grid option turned on.

See Also: Graph.layout fruchterman reingold()

 $\label{layout_sugiyama} \begin{tabular}{ll} layout\_sugiyama (layers=None, weights=None, hgap=1, vgap=1, maxiter=100, return\_extended\_graph=False) \end{tabular}$ 

Places the vertices using a layered Sugiyama layout.

This is a layered layout that is most suitable for directed acyclic graphs, although it works on undirected or cyclic graphs as well.

Each vertex is assigned to a layer and each layer is placed on a horizontal line. Vertices within the same layer are then permuted using the barycenter heuristic that tries to minimize edge crossings.

Dummy vertices will be added on edges that span more than one layer. The returned layout therefore contains more rows than the number of nodes in the original graph; the extra rows correspond to the dummy vertices.

### **Parameters**

layers: a vector specifying a non-negative integer

layer index for each vertex, or the name of a numeric vertex attribute that contains the layer indices. If None, a layering will be determined automatically. For undirected graphs, a spanning tree will be extracted and vertices will be assigned to layers using a breadth first search from the node with the largest degree. For directed graphs, cycles are broken by reversing the direction of edges in an approximate feedback arc set using the heuristic of Eades, Lin and Smyth, and then using longest path layering to

place the vertices in layers.

weights: edge weights to be used. Can be a

sequence or iterable or even an edge

attribute name.

hgap: minimum horizontal gap between vertices

in the same layer.

vgap: vertical gap between layers. The layer

index will be multiplied by vgap to

obtain the Y coordinate.

maxiter: maximum number of iterations to take in

the crossing reduction step. Increase this if you feel that you are getting too many

edge crossings.

return\_extended\_graph: specifies that the extended graph with

the added dummy vertices should also be returned. When this is True, the result will be a tuple containing the layout and the extended graph. The first |V| nodes of the extended graph will correspond to

maximum\_bipartite\_matching(self, types='type', weights=None, eps=None)

Finds a maximum matching in a bipartite graph.

A maximum matching is a set of edges such that each vertex is incident on at most one matched edge and the number (or weight) of such edges in the set is as large as possible.

### **Parameters**

types: vertex types in a list or the name of a vertex attribute

holding vertex types. Types should be denoted by zeros and ones (or False and True) for the two sides of the bipartite graph. If omitted, it defaults to type, which is the default vertex type attribute for bipartite graphs.

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name.

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 this value. This is required to avoid the accumulation of numerical errors. If you pass None here,

igraph will try to determine an appropriate value

automatically.

### Return Value

an instance of Matching.

# to networkx(self)

Converts the graph to network format

### from networkx(klass, q)

Converts the graph from networkx

Vertex names will be converted to "\_nx\_name" attribute and the vertices will get new ids from 0 up (as standard in igraph).

### **Parameters**

g: networkx Graph or DiGraph

 $to\_graph\_tool(self, graph\_attributes=None, vertex\_attributes=None, edge\_attributes=None)$ 

Converts the graph to graph-tool

Data types: graph-tool only accepts specific data types. See the following web page for a list:

https://graph-tool.skewed.de/static/doc/quickstart.html

Note: because of the restricted data types in graph-tool, vertex and edge attributes require to be type-consistent across all vertices or edges. If you set the property for only some vertices/edges, the other will be tagged as None in python-igraph, so they can only be converted to graph-tool with the type 'object' and any other conversion will fail.

### **Parameters**

graph\_attributes: dictionary of graph attributes to transfer.

Keys are attributes from the graph, values are data types (see below). None means no graph

attributes are transferred.

vertex\_attributes: dictionary of vertex attributes to transfer.

Keys are attributes from the vertices, values are data types (see below). None means no

vertex attributes are transferred.

edge\_attributes: dictionary of edge attributes to transfer. Keys

are attributes from the edges, values are data

types (see below). None means no vertex

attributes are transferred.

# from graph tool(klass, g)

Converts the graph from graph-tool

# Parameters

g: graph-tool Graph

# write\_adjacency(self, f, sep=' ', eol='\n', \*args, \*\*kwds)

Writes the adjacency matrix of the graph to the given file

All the remaining arguments not mentioned here are passed intact to Graph.get\_adjacency.

### **Parameters**

f: the name of the file to be written.

sep: the string that separates the matrix elements in a row

eol: the string that separates the rows of the matrix. Please note that igraph is able to read back the written adjacency matrix if and only if this is a single newline character

 $\begin{array}{l} \mathbf{Read\_Adjacency}(klass,\,f,\,sep = \mathtt{None},\,comment\_\,char = \verb|'#|',\,attribute = \mathtt{None},\\ *args,\, \boxed{**}kwds) \end{array}$ 

Constructs a graph based on an adjacency matrix from the given file

Additional positional and keyword arguments not mentioned here are passed intact to Graph. Adjacency.

#### **Parameters**

f: the name of the file to be read or a file object

sep: the string that separates the matrix elements in a

row. None means an arbitrary sequence of

whitespace characters.

comment\_char: lines starting with this string are treated as

comments.

attribute: an edge attribute name where the edge weights are

stored in the case of a weighted adjacency matrix. If None, no weights are stored, values larger than 1 are

considered as edge multiplicities.

### Return Value

the created graph

write dimacs(self, f, source=None, target=None, capacity='capacity')

Writes the graph in DIMACS format to the given file.

# **Parameters**

f: the name of the file to be written or a Python file handle.

source: the source vertex ID. If None, igraph will try to infer it

from the source graph attribute.

target: the target vertex ID. If None, igraph will try to infer it

from the target graph attribute.

capacity: the capacities of the edges in a list or the name of an

edge attribute that holds the capacities. If there is no such edge attribute, every edge will have a capacity of 1.

Overrides: igraph.GraphBase.write\_dimacs

# write graphmlz(self, f, compresslevel=9)

Writes the graph to a zipped GraphML file.

The library uses the gzip compression algorithm, so the resulting file can be unzipped with regular gzip uncompression (like gunzip or zcat from Unix command line) or the Python gzip module.

Uses a temporary file to store intermediate GraphML data, so make sure you have enough free space to store the unzipped GraphML file as well.

### **Parameters**

f: the name of the file to be written.

compresslevel: the level of compression. 1 is fastest and produces

the least compression, and 9 is slowest and

produces the most compression.

# **Read DIMACS**(*f*, *directed*=False)

Reads a graph from a file conforming to the DIMACS minimum-cost flow file format.

For the exact definition of the format, see http://lpsolve.sourceforge.net/5.5/DIMACS.htm.

Restrictions compared to the official description of the format are as follows:

- igraph's DIMACS reader requires only three fields in an arc definition, describing the edge's source and target node and its capacity.
- Source vertices are identified by 's' in the FLOW field, target vertices are identified by 't'.
- Node indices start from 1. Only a single source and target node is allowed.

### **Parameters**

f: the name of the file or a Python file handle

directed: whether the generated graph should be directed.

### Return Value

the generated graph. The indices of the source and target vertices are attached as graph attributes source and target, the edge capacities are stored in the capacity edge attribute.

Overrides: igraph.GraphBase.Read DIMACS

### Read GraphMLz(f, directed=True, index=0)

Reads a graph from a zipped GraphML file.

### **Parameters**

f: the name of the file

index: if the GraphML file contains multiple graphs, specified the one that should be loaded. Graph indices start from zero, so if you want to load the first graph, specify 0 here.

# Return Value

the loaded graph object

# write pickle(self, fname=None, version=-1)

Saves the graph in Python pickled format

### **Parameters**

fname: the name of the file or a stream to save to. If None, saves

the graph to a string and returns the string.

version: pickle protocol version to be used. If -1, uses the highest

protocol available

### Return Value

None if the graph was saved successfully to the given file, or a string if fname was None.

# write picklez(self, fname=None, version=-1)

Saves the graph in Python pickled format, compressed with gzip.

Saving in this format is a bit slower than saving in a Python pickle without compression, but the final file takes up much less space on the hard drive.

#### Parameters

fname: the name of the file or a stream to save to.

version: pickle protocol version to be used. If -1, uses the highest

protocol available

# Return Value

None if the graph was saved successfully to the given file.

# Read Pickle(klass, fname=None)

Reads a graph from Python pickled format

# **Parameters**

fname: the name of the file, a stream to read from, or a string containing the pickled data.

# Return Value

the created graph object.

# Read Picklez(klass, fname, \*args, \*\*kwds)

Reads a graph from compressed Python pickled format, uncompressing it on-the-fly.

### **Parameters**

fname: the name of the file or a stream to read from.

# Return Value

the created graph object.

write\_svg(self, fname, layout='auto', width=None, height=None,
labels='label', colors='color', shapes='shape', vertex\_size=10,
edge\_colors='color', edge\_stroke\_widths='width', font\_size=16, \*args,
\*\*kwds)

Saves the graph as an SVG (Scalable Vector Graphics) file

The file will be Inkscape (http://inkscape.org) compatible. In Inkscape, as nodes are rearranged, the edges auto-update.

### **Parameters**

fname: the name of the file or a Python file handle

layout: the layout of the graph. Can be either an

explicitly specified layout (using a list of coordinate pairs) or the name of a layout algorithm (which should refer to a method in the Graph object, but without the layout\_

prefix.

width: the preferred width in pixels (default: 400)

height: the preferred height in pixels (default: 400)

labels: the vertex labels. Either it is the name of a

vertex attribute to use, or a list explicitly specifying the labels. It can also be None.

colors: the vertex colors. Either it is the name of a

vertex attribute to use, or a list explicitly specifying the colors. A color can be anything acceptable in an SVG file.

shapes: the vertex shapes. Either it is the name of a

vertex attribute to use, or a list explicitly specifying the shapes as integers. Shape 0 means hidden (nothing is drawn), shape 1 is a circle, shape 2 is a rectangle and shape 3 is a rectangle that automatically sizes to the

inner text.

vertex\_size: vertex size in pixels

edge\_colors: the edge colors. Either it is the name of an

edge attribute to use, or a list explicitly specifying the colors. A color can be anything acceptable in an SVG file.

edge\_stroke\_widths: the stroke widths of the edges. Either it is

the name of an edge attribute to use, or a list explicitly specifying the stroke widths. The stroke width can be anything acceptable

in an SVG file.

font\_size: font size. If it is a string, it is written into

the SVG file as-is (so you can specify anything which is valid as the value of the font-size style). If it is a number, it is

# Read(klass, f, format=None, \*args, \*\*kwds)

Unified reading function for graphs.

This method tries to identify the format of the graph given in the first parameter and calls the corresponding reader method.

The remaining arguments are passed to the reader method without any changes.

### **Parameters**

f: the file containing the graph to be loaded

format: the format of the file (if known in advance). None means

auto-detection. Possible values are: "ncol" (NCOL format), "lgl" (LGL format), "graphdb" (GraphDB format), "graphml", "graphmlz" (GraphML and gzipped GraphML format), "gml" (GML format), "net", "pajek" (Pajek format), "dimacs" (DIMACS format), "edgelist", "edges" or "edge" (edge list), "adjacency" (adjacency matrix), "dl" (DL format used by UCINET), "pickle" (Python pickled format), "picklez" (gzipped Python

pickled format)

### Raises

# Load(klass, f, format=None, \*args, \*\*kwds)

Unified reading function for graphs.

This method tries to identify the format of the graph given in the first parameter and calls the corresponding reader method.

The remaining arguments are passed to the reader method without any changes.

# **Parameters**

f: the file containing the graph to be loaded

format: the format of the file (if known in advance). None means auto-detection. Possible values are: "ncol" (NCOL format), "lgl" (LGL format), "graphdb" (GraphDB format), "graphml", "graphmlz" (GraphML and gzipped GraphML format), "gml" (GML format), "net", "pajek" (Pajek format), "dimacs" (DIMACS format), "edgelist", "edges" or "edge" (edge list), "adjacency" (adjacency matrix), "dl" (DL format used by UCINET), "pickle" (Python pickled format), "picklez" (gzipped Python

pickled format)

### Raises

# write(self, f, format=None, \*args, \*\*kwds)

Unified writing function for graphs.

This method tries to identify the format of the graph given in the first parameter (based on extension) and calls the corresponding writer method.

The remaining arguments are passed to the writer method without any changes.

# **Parameters**

f: the file containing the graph to be saved

- "adjacency": adjacency matrix format
- "dimacs": DIMACS format
- "dot", "graphviz": GraphViz DOT format
- "edgelist", "edges" or "edge": numeric edge list format
- "gml": GML format
- "graphml" and "graphmlz": standard and gzipped GraphML format
- "gw", "leda", "lgr": LEDA native format
- "lgl": LGL format
- "ncol": NCOL format
- "net", "pajek": Pajek format
- "pickle", "picklez": standard and gzipped Python pickled format
- "svg": SVG format

### Raises

# **save**(self, f, format=None, \*args, \*\*kwds)

Unified writing function for graphs.

This method tries to identify the format of the graph given in the first parameter (based on extension) and calls the corresponding writer method.

The remaining arguments are passed to the writer method without any changes.

# **Parameters**

f: the file containing the graph to be saved

- "adjacency": adjacency matrix format
- "dimacs": DIMACS format
- "dot", "graphviz": GraphViz DOT format
- "edgelist", "edges" or "edge": numeric edge list format
- "gml": GML format
- "graphml" and "graphmlz": standard and gzipped GraphML format
- "gw", "leda", "lgr": LEDA native format
- "lgl": LGL format
- "ncol": NCOL format
- "net", "pajek": Pajek format
- "pickle", "picklez": standard and gzipped Python pickled format
- "svg": SVG format

### Raises

DictList(klass, vertices, edges, directed=False, vertex\_name\_attr='name',
edge\_foreign\_keys=('source', 'target'), iterative=False)

Constructs a graph from a list-of-dictionaries representation.

This representation assumes that vertices and edges are encoded in two lists, each list containing a Python dict for each vertex and each edge, respectively. A distinguished element of the vertex dicts contain a vertex ID which is used in the edge dicts to refer to source and target vertices. All the remaining elements of the dict are considered vertex and edge attributes. Note that the implementation does not assume that the objects passed to this method are indeed lists of dicts, but they should be iterable and they should yield objects that behave as dicts. So, for instance, a database query result is likely to be fit as long as it's iterable and yields dict-like objects with every iteration.

# **Parameters**

vertices: the data source for the vertices or None if

there are no special attributes assigned to vertices and we should simply use the edge list

of dicts to infer vertex names.

edges: the data source for the edges.

directed: whether the constructed graph will be directed

vertex\_name\_attr: the name of the distinguished key in the dicts

in the vertex data source that contains the vertex names. Ignored if vertices is None.

edge\_foreign\_keys: the name of the attributes in the dicts in the

edge data source that contain the source and

target vertex names.

iterative: whether to add the edges to the graph one by

one, iteratively, or to build a large edge list first and use that to construct the graph. The latter approach is faster but it may not be suitable if your dataset is large. The default is to add the edges in a batch from an edge list.

### Return Value

the graph that was constructed

TupleList(klass, edges, directed=False, vertex\_name\_attr='name',
edge\_attrs=None, weights=False)

Constructs a graph from a list-of-tuples representation.

This representation assumes that the edges of the graph are encoded in a list of tuples (or lists). Each item in the list must have at least two elements, which specify the source and the target vertices of the edge. The remaining elements (if any) specify the edge attributes of that edge, where the names of the edge attributes originate from the edge\_attrs list. The names of the vertices will be stored in the vertex attribute given by vertex\_name\_attr.

The default parameters of this function are suitable for creating unweighted graphs from lists where each item contains the source vertex and the target vertex. If you have a weighted graph, you can use items where the third item contains the weight of the edge by setting edge\_attrs to "weight" or ["weight"]. If you have even more edge attributes, add them to the end of each item in the edges list and also specify the corresponding edge attribute names in edge\_attrs as a list.

### Parameters

edges: the data source for the edges. This must be a

list where each item is a tuple (or list)

containing at least two items: the name of the source and the target vertex. Note that names will be assigned to the name vertex attribute

(or another vertex attribute if

vertex\_name\_attr is specified), even if all the vertex names in the list are in fact numbers.

directed: whether the constructed graph will be directed

vertex\_name\_attr: the name of the vertex attribute that will

contain the vertex names.

edge\_attrs: the names of the edge attributes that are filled

with the extra items in the edge list (starting from index 2, since the first two items are the source and target vertices). None means that only the source and target vertices will be extracted from each item. If you pass a string

here, it will be wrapped in a list for

convenience.

weights: alternative way to specify that the graph is

weighted. If you set weights to true and edge\_attrs is not given, it will be assumed that edge\_attrs is ["weight"] and igraph will parse the third element from each item into an edge weight. If you set weights to a string, it will be assumed that edge\_attrs contains that

string only, and igraph will store the edge

weights in that attribute.

# Return Value

# Formula(Graph, formula=None, attr="name", simplify=True)

Generates a graph from a graph formula

A graph formula is a simple string representation of a graph. It is very handy for creating small graphs quickly. The string consists of vertex names separated by edge operators. An edge operator is a sequence of dashes (-) that may or may not start with an arrowhead (< at the beginning of the sequence or > at the end of the sequence). The edge operators can be arbitrarily long, i.e., you may use as many dashes to draw them as you like. This makes a total of four different edge operators:

- --- makes an undirected edge
- <-- makes a directed edge pointing from the vertex on the right hand side of the operator to the vertex on the left hand side
- --> is the opposite of <--
- <--> creates a mutual directed edge pair between the two vertices

If you only use the undirected edge operator (---), the graph will be undirected. Otherwise it will be directed. Vertex names used in the formula will be assigned to the name vertex attribute of the graph.

Some simple examples:

```
>>> from igraph import Graph
>>> print Graph.Formula()
                                        # empty graph
IGRAPH UN-- O O --
+ attr: name (v)
>>> g = Graph.Formula("A-B")
                                   # undirected graph
>>> g.vs["name"]
['A', 'B']
>>> print g
IGRAPH UN-- 2 1 --
+ attr: name (v)
+ edges (vertex names):
A--B
>>> g.get_edgelist()
[(0, 1)]
>>> g2 = Graph.Formula("A-----B")
>>> g2.isomorphic(g)
True
>>> g = Graph.Formula("A ---> B") # directed graph
>>> g.vs["name"]
['A', 'B']
>>> print g
IGRAPH DN-- 2 1 --
+ attr: name (v)
                                       72
+ edges (vertex names):
A->B
```

If you have may disconnected components, you can separate them with commas. You can also specify isolated vertices:

## **Bipartite**(types, edges, directed=False)

Creates a bipartite graph with the given vertex types and edges. This is similar to the default constructor of the graph, the only difference is that it checks whether all the edges go between the two vertex classes and it assigns the type vector to a type attribute afterwards.

# Examples:

```
>>> g = Graph.Bipartite([0, 1, 0, 1], [(0, 1), (2, 3), (0, 3)])
>>> g.is_bipartite()
True
>>> g.vs["type"]
[False, True, False, True]
```

#### **Parameters**

types: the vertex types as a boolean list. Anything that

evaluates to False will denote a vertex of the first kind, anything that evaluates to True will denote a vertex of

the second kind.

edges: the edges as a list of tuples.

directed: whether to create a directed graph. Bipartite networks

are usually undirected, so the default is False

#### Return Value

the graph with a binary vertex attribute named "type" that stores the vertex classes.

# Full Bipartite(n1, n2, directed=False, mode=ALL)

Generates a full bipartite graph (directed or undirected, with or without loops).

```
>>> g = Graph.Full_Bipartite(2, 3)
>>> g.is_bipartite()
True
>>> g.vs["type"]
[False, False, True, True, True]
```

#### **Parameters**

n1: the number of vertices of the first kind.

n2: the number of vertices of the second kind.

directed: whether tp generate a directed graph.

mode: if OUT, then all vertices of the first kind are connected to

the others; IN specifies the opposite direction, ALL creates mutual edges. Ignored for undirected graphs.

### Return Value

the graph with a binary vertex attribute named "type" that stores the vertex classes.

 $\label{eq:andom_Bipartite} \textbf{Random\_Bipartite}(\textit{n1}, \textit{n2}, \textit{p}=\texttt{None}, \textit{m}=\texttt{None}, \textit{directed}=\texttt{False}, \\ \textit{neimode}=\texttt{ALL})$ 

Generates a random bipartite graph with the given number of vertices and edges (if m is given), or with the given number of vertices and the given connection probability (if p is given).

If m is given but p is not, the generated graph will have n1 vertices of type 1, n2 vertices of type 2 and m randomly selected edges between them. If p is given but m is not, the generated graph will have n1 vertices of type 1 and n2 vertices of type 2, and each edge will exist between them with probability p.

### **Parameters**

n1: the number of vertices of type 1.

n2: the number of vertices of type 2.

p: the probability of edges. If given, m must be missing.

m: the number of edges. If given, p must be missing.

directed: whether to generate a directed graph.

neimode: if the graph is directed, specifies how the edges will be

generated. If it is "all", edges will be generated in both

directions (from type 1 to type 2 and vice versa)

independently. If it is "out" edges will always point from type 1 to type 2. If it is "in", edges will always point from type 2 to type 1. This argument is ignored for

undirected graphs.

### GRG(n, radius, torus=False, return coordinates=False)

Generates a random geometric graph.

The algorithm drops the vertices randomly on the 2D unit square and connects them if they are closer to each other than the given radius. The coordinates of the vertices are stored in the vertex attributes x and y.

### **Parameters**

n: The number of vertices in the graph

radius: The given radius

torus: This should be True if we want to use a torus instead of a

square.

 $\begin{aligned} &\textbf{Incidence}(\textit{matrix}, \textit{directed} = \texttt{False}, \textit{mode} = \texttt{ALL}, \textit{multiple} = \texttt{False}, \\ &\textit{weighted} = \texttt{None}) \end{aligned}$ 

Creates a bipartite graph from an incidence matrix.

## Example:

>>> g = Graph.Incidence([[0, 1, 1], [1, 1, 0]])

### **Parameters**

matrix: the incidence matrix.

directed: whether to create a directed graph.

mode: defines the direction of edges in the graph. If OUT, then

edges go from vertices of the first kind (corresponding to rows of the matrix) to vertices of the second kind (the columns of the matrix). If IN, the opposite direction is used. ALL creates mutual edges. Ignored for undirected

graphs.

multiple: defines what to do with non-zero entries in the matrix. If

False, non-zero entries will create an edge no matter what the value is. If True, non-zero entries are rounded up to the nearest integer and this will be the number of

multiple edges created.

weighted: defines whether to create a weighted graph from the

incidence matrix. If it is  $c\{None\}$  then an unweighted graph is created and the multiple argument is used to determine the edges of the graph. If it is a string 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'.

### Return Value

the graph with a binary vertex attribute named "type" that stores the vertex classes.

#### Raises

ValueError if the weighted and multiple are passed together.

# DataFrame(edges, directed=True, vertices=None)

Generates a graph from one or two dataframes.

## **Parameters**

edges: pandas DataFrame containing edges and metadata

directed: bool setting whether the graph is directed

vertices: None (default) or pandas DataFrame containing vertex

metadata. The first column must contain the unique ids of the vertices and will be set as attribute 'name'. All other columns will be added as vertex attributes by

column name.

## Return Value

the graph

bipartite\_projection(self, types='type', multiplicity=True, probe1=-1,
which='both')

Projects a bipartite graph into two one-mode graphs. Edge directions are ignored while projecting.

## Examples:

```
>>> g = Graph.Full_Bipartite(10, 5)
>>> g1, g2 = g.bipartite_projection()
>>> g1.isomorphic(Graph.Full(10))
True
>>> g2.isomorphic(Graph.Full(5))
```

### **Parameters**

types: an igraph vector containing the vertex types, or an

attribute name. Anything that evalulates to False corresponds to vertices of the first kind, everything

else to the second kind.

multiplicity: if True, then igraph keeps the multiplicity of the

edges in the projection in an edge attribute called "weight". E.g., if there is an A-C-B and an A-D-B triplet in the bipartite graph and there is no other X (apart from X=B and X=D) for which an A-X-B triplet would exist in the bipartite graph, 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 and non-negative, then it is considered as a vertex ID; the projection containing the vertex will be the first

one in the result.

which: this argument can be used to specify which of the

two projections should be returned if only one of them is needed. Passing 0 here means that only the first projection is returned, while 1 means that only the second projection is returned. (Note that we use 0 and 1 because Python indexing is zero-based). False is equivalent to 0 and True is equivalent to 1. Any other value means that both projections will be

returned in a tuple.

### Return Value

a tuple containing the two projected one-mode graphs if which is not 1 or 2, or the projected one-mode graph specified by the which argument if its value is 0, 1, False or True.

Overrides: igraph.GraphBase.bipartite\_projection

# bipartite projection size(types="type")

Calculates the number of vertices and edges in the bipartite projections of this graph according to the specified vertex types. This is useful if you have a bipartite graph and you want to estimate the amount of memory you would need to calculate the projections themselves.

### Parameters

types: an igraph vector containing the vertex types, or an attribute name. Anything that evalulates to False corresponds to vertices of the first kind, everything else to the second kind.

## Return Value

a 4-tuple containing the number of vertices and edges in the first projection, followed by the number of vertices and edges in the second projection.

Overrides: igraph.GraphBase.bipartite projection size

## get incidence(self, types="type")

Returns the incidence matrix of a bipartite graph. The incidence matrix is an n times m matrix, where n and m are the number of vertices in the two vertex classes.

## **Parameters**

types: an igraph vector containing the vertex types, or an attribute name. Anything that evalulates to False corresponds to vertices of the first kind, everything else to the second kind.

#### Return Value

the incidence matrix and two lists in a triplet. The first list defines the mapping between row indices of the matrix and the original vertex IDs. The second list is the same for the column indices.

Overrides: igraph.GraphBase.get incidence

#### $\mathbf{dfs}(self, vid, mode=1)$

Conducts a depth first search (DFS) on the graph.

### **Parameters**

vid: the root vertex ID

mode: either IN or OUT or ALL, ignored for undirected graphs.

## Return Value

a tuple with the following items:

- The vertex IDs visited (in order)
- The parent of every vertex in the DFS

iadd (self, other)

In-place addition (disjoint union).

See Also: \_\_add\_\_

add (self, other)

Copies the graph and extends the copy depending on the type of the other object given.

### **Parameters**

other: if it is an integer, the copy is extended by the given number of vertices. If it is a string, the copy is extended by a single vertex whose name attribute will be equal to the given string. If it is a tuple with two elements, the copy is extended by a single edge. If it is a list of tuples, the copy is extended by multiple edges. If it is a Graph, a disjoint union is performed.

and (self, other)

Graph intersection operator.

## Parameters

other: the other graph to take the intersection with.

#### Return Value

the intersected graph.

\_\_isub\_\_\_(self, other)

In-place subtraction (difference).

See Also: \_\_sub\_\_

**sub** (self, other)

Removes the given object(s) from the graph

#### **Parameters**

other: if it is an integer, removes the vertex with the given ID from the graph (note that the remaining vertices will get re-indexed!). If it is a tuple, removes the given edge. If it is a graph, takes the difference of the two graphs. Accepts lists of integers or lists of tuples as well, but they can't be mixed! Also accepts Edge and EdgeSeq objects.

$\operatorname{mul}$	(self,	other)

Copies exact replicas of the original graph an arbitrary number of times.

### **Parameters**

other: if it is an integer, multiplies the graph by creating the given number of identical copies and taking the disjoint union of them.

```
\_\_nonzero\_\_(self)
```

Returns True if the graph has at least one vertex, False otherwise.

```
\mathbf{or} (self, other)
```

Graph union operator.

## Parameters

other: the other graph to take the union with.

# Return Value

the union graph.

\_\_coerce\_\_(self, other)

Coercion rules.

This method is needed to allow the graph to react to additions with lists, tuples, integers, strings, vertices, edges and so on.

 $\_\_reduce\_\_(self)$ 

Support for pickling.

Overrides: object.\_\_reduce\_\_

plot (self, context, bbox, palette, \*args, \*\*kwds)

Plots the graph to the given Cairo context in the given bounding box

The visual style of vertices and edges can be modified at three places in the following order of precedence (lower indices override higher indices):

- 1. Keyword arguments of this function (or of plot() which is passed intact to Graph.\_\_plot\_\_().
- 2. Vertex or edge attributes, specified later in the list of keyword arguments.
- 3. igraph-wide plotting defaults (see igraph.config.Configuration)
- 4. Built-in defaults.

E.g., if the vertex\_size keyword attribute is not present, but there exists a vertex attribute named size, the sizes of the vertices will be specified by that attribute.

Besides the usual self-explanatory plotting parameters (context, bbox, palette), it accepts the following keyword arguments:

- autocurve: whether to use curves instead of straight lines for multiple edges on the graph plot. This argument may be True or False; when omitted, True is assumed for graphs with less than 10.000 edges and False otherwise.
- drawer\_factory: a subclass of AbstractCairoGraphDrawer which will be used to draw the graph. You may also provide a function here which takes two arguments: the Cairo context to draw on and a bounding box (an instance of BoundingBox). If this keyword argument is missing, igraph will use the default graph drawer which should be suitable for most purposes. It is safe to omit this keyword argument unless you need to use a specific graph drawer.
- keep\_aspect\_ratio: whether to keep the aspect ratio of the layout that igraph calculates to place the nodes. True means that the layout will be scaled proportionally to fit into the bounding box where the graph is to be drawn but the aspect ratio will be kept the same (potentially leaving empty space next to, below or above the graph). False means that the layout will be scaled independently along the X and Y axis in order to fill the entire bounding box. The default is False.
- layout: the layout to be used. If not an instance of Layout, it will be passed to Graph.layout to calculate the layout. Note that if you want a deterministic layout that does not change with every plot, you must either use a deterministic layout function (like Graph.layout\_circle) or calculate the layout in advance and pass a Layout object here.
- margin: the top, right, bottom, left margins as a 4-tuple. If it has less than 4 elements or is a single float, the elements will be re-used until the length is at least 4.
- mark\_groups: whether to highlighter some of the vertex groups by colored polygons. This argument can be one of the following:
  - False: no groups will be highlighted
  - A dict mapping tuples of vertex indices to color names. The given

# str (self)

Returns a string representation of the graph.

Behind the scenes, this method constructs a **GraphSummary** instance and invokes its \_\_str\_\_ method with a verbosity of 1 and attribute printing turned off.

See the documentation of GraphSummary for more details about the output.

Overrides: object.\_\_str\_\_

# summary(self, verbosity=0, width=None, \*arqs, \*\*kwds)

Returns the summary of the graph.

The output of this method is similar to the output of the <code>\_\_str\_\_</code> method. If *verbosity* is zero, only the header line is returned (see <code>\_\_str\_\_</code> for more details), otherwise the header line and the edge list is printed.

Behind the scenes, this method constructs a GraphSummary object and invokes its \_\_str\_\_ method.

#### **Parameters**

verbosity: if zero, only the header line is returned (see \_\_str\_\_ for

more details), otherwise the header line and the full

edge list is printed.

width: the number of characters to use in one line. If None, no

limit will be enforced on the line lengths.

### Return Value

the summary of the graph.

# **disjoint** union(self, other)

Creates the disjoint union of two (or more) graphs.

#### Parameters

other: graph or list of graphs to be united with the current one.

#### Return Value

the disjoint union graph

## union(self, other, byname="auto")

Creates the union of two (or more) graphs.

### **Parameters**

other: graph or list of graphs to be united with the current one.

byname: whether to use vertex names instead of ids. See

igraph.union for details.

## Return Value

the union graph

# intersection(self, other, byname="auto")

Creates the intersection of two (or more) graphs.

### **Parameters**

other: graph or list of graphs to be intersected with the current

one.

byname: whether to use vertex names instead of ids. See

igraph.intersection for details.

### Return Value

the intersection graph

# layout fruchterman reingold 3d(\*args, \*\*kwds)

Alias for layout\_fruchterman\_reingold() with dim=3.

See Also: Graph.layout fruchterman reingold()

# layout kamada kawai 3d(\*args, \*\*kwds)

Alias for layout\_kamada\_kawai() with dim=3.

See Also: Graph.layout\_kamada\_kawai()

# layout random 3d(\*args, \*\*kwds)

Alias for layout\_random() with dim=3.

See Also: Graph.layout random()

# layout grid 3d(\*args, \*\*kwds)

Alias for layout\_grid() with dim=3.

See Also: Graph.layout grid()

layout sphere(\*args, \*\*kwds)

Alias for layout\_circle() with dim=3.

See Also: Graph.layout\_circle()

layout bipartite(types="type", hgap=1, vgap=1, maxiter=100)

Place the vertices of a bipartite graph in two layers.

The layout is created by placing the vertices in two rows, according to their types. The positions of the vertices within the rows are then optimized to minimize the number of edge crossings using the heuristic used by the Sugiyama layout algorithm.

### **Parameters**

types: an igraph vector containing the vertex types, or an

attribute name. Anything that evalulates to False

corresponds to vertices of the first kind, everything else to

the second kind.

hgap: minimum horizontal gap between vertices in the same

layer.

vgap: vertical gap between the two layers.

maxiter: maximum number of iterations to take in the crossing

reduction step. Increase this if you feel that you are

getting too many edge crossings.

### Return Value

the calculated layout.

Overrides: igraph.GraphBase.layout bipartite

# layout circle(dim=2, order=None)

Places the vertices of the graph uniformly on a circle or a sphere.

## **Parameters**

dim: the desired number of dimensions for the layout. dim=2

means a 2D layout, dim=3 means a 3D layout.

order: the order in which the vertices are placed along the circle.

Not supported when dim is not equal to 2.

#### Return Value

the calculated layout.

Overrides: igraph.GraphBase.layout circle

```
layout davidson harel(seed=None, maxiter=10, fineiter=-1,
cool fact=0.75, weight node dist=1.0, weight border=0.0,
weight edge lengths=-1, weight edge crossings=-1,
weight node edge dist=-1)
```

Places the vertices on a 2D plane according to the Davidson-Harel layout algorithm.

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.

#### Parameters

seed: if None, uses a random starting layout for

> the algorithm. If a matrix (list of lists), uses the given matrix as the starting

position.

Number of iterations to perform in the maxiter:

annealing phase.

fineiter: Number of iterations to perform in the

> fine-tuning phase. Negative numbers set up a reasonable default from the base-2 logarithm of the vertex count, bounded

by 10 from above.

Cooling factor of the simulated annealing cool\_fact:

phase.

Weight for the node-node distances in the weight\_node\_dist:

energy function.

Weight for the distance from the border weight\_border:

> component of the energy function. Zero means that vertices are allowed to sit on the border of the area designated for the

layout.

weight\_edge\_lengths: Weight for the edge length component of

> the energy function. Negative numbers are replaced by the density of the graph

divided by 10.

weight\_edge\_crossings: Weight for the edge crossing component

of the energy function. Negative numbers are replaced by one minus the square root of the density of the graph.

weight\_node\_edge\_dist: Weight for the node-edge distance

component of the energy function. Negative numbers are replaced by 0.2

minus 0.2 times the density of the graph.

layout drl(weights=None, fixed=None, seed=None, options=None, dim=2)

Places the vertices on a 2D plane or in the 3D space ccording to the DrL layout algorithm.

This is an algorithm suitable for quite large graphs, but it can be surprisingly slow for small ones (where the simpler force-based layouts like layout\_kamada\_kawai() or layout\_fruchterman\_reingold() are more useful.

# Parameters

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name.

seed: if None, uses a random starting layout for the algorithm.

If a matrix (list of lists), uses the given matrix as the

starting position.

fixed: if a seed is given, you can specify some vertices to be kept

fixed at their original position in the seed by passing an appropriate list here. The list must have exactly as many items as the number of vertices in the graph. Items of the list that evaluate to **True** denote vertices that will not be

moved.

options: if you give a string argument here, you can select from five default preset parameterisations: default, coarsen for a coarser layout, coarsest for an even coarser layout,

refine for refining an existing layout and final for finalizing a layout. If you supply an object that is not a string, the DrL layout parameters are retrieved from the respective keys of the object (so it should be a dict or something else that supports the mapping protocol). The

following keys can be used:

- 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 initialization phase
- init\_temperature: start temperature during initialization
- init\_attraction: attraction during initialization
- init\_damping\_mult: damping multiplier during initialization
- liquid\_iterations, liquid\_temperature, liquid\_attraction, liquid\_damping\_mult: same parameters for the liquid phase
- expansion\_iterations, expansion\_temperature,

 $\begin{array}{l} \textbf{layout\_fruchterman\_reingold}(weights=\texttt{None},\ niter=\texttt{500},\ seed=\texttt{None},\ start\_temp=\texttt{None},\ minx=\texttt{None},\ maxx=\texttt{None},\ miny=\texttt{None},\ maxy=\texttt{None},\ minz=\texttt{None},\ maxz=\texttt{None},\ grid=\texttt{"auto"}) \end{array}$ 

Places the vertices on a 2D plane according to the Fruchterman-Reingold algorithm.

This is a force directed layout, see Fruchterman, T. M. J. and Reingold, E. M.: Graph Drawing by Force-directed Placement. Software – Practice and Experience, 21/11, 1129–1164, 1991

#### **Parameters**

weights: edge weights to be used. Can be a sequence or iterable

or even an edge attribute name.

niter: the number of iterations to perform. The default is

500.

start\_temp: Real scalar, the start temperature. This is the

maximum amount of movement alloved along one axis, within one step, for a vertex. Currently it is decreased linearly to zero during the iteration. The default is the square root of the number of vertices divided by 10.

minx: if not None, it must be a vector with exactly as many

elements as there are vertices in the graph. Each element is a minimum constraint on the X value of the

vertex in the layout.

maxx: similar to minx, but with maximum constraints

miny: similar to minx, but with the Y coordinates

maxy: similar to maxx, but with the Y coordinates

minz: similar to minx, but with the Z coordinates. Use only

for 3D layouts (dim=3).

maxz: similar to maxx, but with the Z coordinates. Use only

for 3D layouts (dim=3).

seed: if None, uses a random starting layout for the

algorithm. If a matrix (list of lists), uses the given

matrix as the starting position.

grid: whether to use a faster, but less accurate grid-based

implementation of the algorithm. "auto" decides based on the number of vertices in the graph; a grid will be used if there are at least 1000 vertices. "grid" is equivalent to True, "nogrid" is equivalent to False.

### Return Value

the calculated layout.

Overrides: igraph.GraphBase.layout\_fruchterman\_reingold

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$$\label{layout_graphopt} \begin{split} &\textbf{layout\_graphopt}(niter = \texttt{500}, \ node\_charge = \texttt{0.001}, \ node\_mass = \texttt{30}, \\ &spring\_length = \texttt{0}, \ spring\_constant = \texttt{1}, \ max\_sa\_movement = \texttt{5}, \ seed = \texttt{None}) \end{split}$$

This 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.

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 or the maximal number of iterations is reached.

See http://www.schmuhl.org/graphopt/ for the original graphopt.

## **Parameters**

niter: the number of iterations to perform. Should be a

couple of hundred in general.

node\_charge: the charge of the vertices, used to calculate

electric repulsion.

node\_mass: the mass of the vertices, used for the spring

forces

spring\_length: the length of the springs

spring\_constant: the spring constant

max\_sa\_movement: the maximum amount of movement allowed in a

single step along a single axis.

seed: a matrix containing a seed layout from which

the algorithm will be started. If None, a random

layout will be used.

### Return Value

the calculated layout.

Overrides: igraph.GraphBase.layout graphopt

# **layout grid**(*width*=0, *height*=0, *dim*=2)

Places the vertices of a graph in a 2D or 3D grid.

## **Parameters**

width: the number of vertices in a single row of the layout. Zero or

negative numbers mean that the width should be

determined automatically.

height: the number of vertices in a single column of the layout.

Zero or negative numbers mean that the height should be determined automatically. It must not be given if the

number of dimensions is 2.

dim: the desired number of dimensions for the layout. dim=2

means a 2D layout, dim=3 means a 3D layout.

## Return Value

the calculated layout.

Overrides: igraph.GraphBase.layout\_grid

 $\begin{array}{l} \textbf{layout\_kamada\_kawai}(maxiter=1000,\,seed=\texttt{None},\,maxiter=1000,\\epsilon=\texttt{0},\,kkconst=\texttt{None},\,minx=\texttt{None},\,maxx=\texttt{None},\,miny=\texttt{None},\\maxy=\texttt{None},\,minz=\texttt{None},\,maxz=\texttt{None},\,dim=\texttt{2}) \end{array}$ 

Places the vertices on a plane according to the Kamada-Kawai algorithm.

This is a force directed layout, see Kamada, T. and Kawai, S.: An Algorithm for Drawing General Undirected Graphs. Information Processing Letters, 31/1, 7–15, 1989.

#### **Parameters**

maxiter: the maximum number of iterations to perform.

seed: if None, uses a random starting layout for the algorithm.

If a matrix (list of lists), uses the given matrix as the

starting position.

epsilon: quit if the energy of the system changes less than epsilon.

See the original paper for details.

kkconst: the Kamada-Kawai vertex attraction constant. None

means the square of the number of vertices.

minx: if not None, it must be a vector with exactly as many

elements as there are vertices in the graph. Each element is a minimum constraint on the X value of the vertex in

the layout.

maxx: similar to minx, but with maximum constraints

miny: similar to minx, but with the Y coordinates

maxy: similar to maxx, but with the Y coordinates

minz: similar to minx, but with the Z coordinates. Use only for

3D layouts (dim=3).

maxz: similar to maxx, but with the Z coordinates. Use only for

3D layouts (dim=3).

dim: the desired number of dimensions for the layout. dim=2

means a 2D layout, dim=3 means a 3D layout.

#### Return Value

the calculated layout.

Overrides: igraph.GraphBase.layout kamada kawai

layout\_lgl(maxiter=150, maxdelta=-1, area=-1, coolexp=1.5,
repulserad=-1, cellsize=-1, root=None)

Places the vertices on a 2D plane according to the Large Graph Layout.

#### **Parameters**

maxiter: the number of iterations to perform.

maxdelta: the maximum distance to move a vertex in an

iteration. If negative, defaults to the number of

vertices.

area: the area of the square on which the vertices will be

placed. If negative, defaults to the number of vertices

squared.

coolexp: the cooling exponent of the simulated annealing.

repulserad: determines the radius at which vertex-vertex repulsion

cancels out attraction of adjacent vertices. If negative,

defaults to area times the number of vertices.

cellsize: the size of the grid cells. When calculating the

repulsion forces, only vertices in the same or

neighboring grid cells are taken into account. Defaults

to the fourth root of area.

root: the root vertex, this is placed first, its neighbors in the

first iteration, second neighbors in the second, etc.

None means that a random vertex will be chosen.

## Return Value

the calculated layout.

Overrides: igraph.GraphBase.layout lgl

# layout mds(dist=None, dim=2, arpack options=None)

Places the vertices in an Euclidean space with the given number of dimensions using multidimensional scaling.

This layout requires a distance matrix, where the intersection of row i and column j specifies the desired distance between vertex i and vertex j. The algorithm will try to place the vertices in a way that approximates the distance relations prescribed in the distance matrix. igraph uses the classical multidimensional scaling by Torgerson (see reference below).

For unconnected graphs, the method will decompose the graph into weakly connected components and then lay out the components individually using the appropriate parts of the distance matrix.

#### **Parameters**

dist: the distance matrix. It must be symmetric and

the symmetry is not checked – results are unspecified when a non-symmetric distance matrix is used. If this parameter is None, the shortest path lengths will be used as distances. Directed graphs are treated as undirected when calculating the shortest path lengths to ensure

symmetry.

dim: the number of dimensions. For 2D layouts, supply

2 here; for 3D layouts, supply 3.

arpack\_options: an ARPACKOptions object used to fine-tune the

ARPACK eigenvector calculation. If omitted, the module-level variable called arpack\_options is

used.

### Return Value

the calculated layout.

Overrides: igraph.GraphBase.layout mds

Reference: Cox & Cox: Multidimensional Scaling (1994), Chapman and Hall,

London.

## layout random(dim=2)

Places the vertices of the graph randomly.

#### **Parameters**

dim: the desired number of dimensions for the layout. dim=2 means a 2D layout, dim=3 means a 3D layout.

### Return Value

the coordinate pairs in a list.

Overrides: igraph.GraphBase.layout random

# layout reingold tilford(mode="out", root=None, rootlevel=None)

Places the vertices on a 2D plane according to the Reingold-Tilford layout algorithm.

This is a tree layout. If the given graph is not a tree, a breadth-first search is executed first to obtain a possible spanning tree.

#### **Parameters**

mode: specifies which edges to consider when builing 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 behaviour in igraph 0.5 and before). This parameter also influences how the root vertices are calculated if they are not

given. See the *root* parameter.

root: the index of the root vertex or root vertices. if this is a

non-empty vector then the supplied vertex IDs are used as the roots of the trees (or a single tree if the graph is connected. If this is None or an empty list, the root

vertices are automatically calculated based on

topological sorting, performed with the opposite of the

mode argument.

rootlevel: this argument is useful when drawing forests which are

not trees. It specifies the level of the root vertices for

every tree in the forest.

#### Return Value

the calculated layout.

Overrides: igraph.GraphBase.layout reingold tilford

See Also: layout reingold tilford circular

Reference: EM Reingold, JS Tilford: Tidier Drawings of Trees. IEEE

Transactions on Software Engineering 7:22, 223-228, 1981.

# 

Circular Reingold-Tilford layout for trees.

This layout is similar to the Reingold-Tilford layout, but the vertices are placed in a circular way, with the root vertex in the center.

See layout\_reingold\_tilford for the explanation of the parameters.

#### Return Value

the calculated layout.

Overrides: igraph.GraphBase.layout reingold tilford circular

See Also: layout reingold tilford

Reference: EM Reingold, JS Tilford: Tidier Drawings of Trees. IEEE

Transactions on Software Engineering 7:22, 223-228, 1981.

## layout star(center=0, order=None)

Calculates a star-like layout for the graph.

### **Parameters**

center: the ID of the vertex to put in the center

order: a numeric vector giving the order of the vertices (including

the center vertex!). If it is None, the vertices will be placed

in increasing vertex ID order.

## Return Value

the calculated layout.

Overrides: igraph.GraphBase.layout star

# $Inherited\ from\ igraph.GraphBase(Section\ 1.12)$

Adjacency(), Asymmetric\_Preference(), Atlas(), Barabasi(), De\_Bruijn(), De-gree\_Sequence(), Erdos\_Renyi(), Establishment(), Famous(), Forest\_Fire(), Full(), Full\_Citation(), Growing\_Random(), Isoclass(), K\_Regular(), Kautz(), LCF(), Lattice(), Preference(), Read\_DL(), Read\_Edgelist(), Read\_GML(), Read\_GraphDB(), Read\_GraphML(), Read\_Lgl(), Read\_Ncol(), Read\_Pajek(), Recent\_Degree(), Ring(), SBM(), Star(), Static\_Fitness(), Static\_Power\_Law(), Tree(), Watts\_Strogatz(), Weighted\_Adjacency(), \_\_delitem\_\_(), \_\_getitem\_\_(), \_\_invert\_\_(), \_\_new\_\_(), \_\_setitem\_\_(), all\_minimal\_st\_separators(), are\_connected(), articulation\_points(), assortativity(), assortativity\_degree(), assortativity\_nominal(), attributes(), authority\_score(), average\_path\_length(), betweenness(), bfs(), bfsiter(), bibcoupling(), bridges(), canonical\_permutation(), clique\_number(), cliques(), closeness(), cocitation(), complementer(), compose(), constraint(), contract\_vertices(), convergence\_degree(), convergence\_field\_size(), copy(), coreness(), count\_isomorphisms\_vf2(),

count multiple(), count subisomorphisms vf2(), decompose(), degree(), delete vertices(), density(), dfsiter(), diameter(), difference(), diversity(), dominator(), eccentricity(), ecount(), edge attributes(), edge betweenness(), edge connectivity(), eigen adjacency(), eigenvector centrality(), farthest points(), feedback arc set(), get all shortest paths(), get diameter(), get edgelist(), get eid(), get eids(), get isomorphisms vf2(), get shortest paths(), get subisomorphisms lad(), get subisomorphisms vf2(), girth(), has multiple(), hub score(), incident(), independence number(), independent vertex sets(), induced subgraph(), is bipartite(), is connected(), is dag(), is directed(), is loop(), is minimal separator(), is multiple(), is mutual(), is separator(), is simple(), isoclass(), isomorphic(), isomorphic bliss(), isomorphic vf2(), knn(), laplacian(), largest cliques(), largest independent vertex sets(), linegraph(), maxdegree(), maxflow value(), maximal cliques(), maximal independent vertex sets(), mincut value(), minimum size separators(), motifs randesu(), motifs randesu estimate(), motifs randesu no(), neighborhood(), neighborhood size(), neighbors(), permute vertices(), personalized pagerank(), predecessors(), radius(), random walk(), reciprocity(), rewire(), rewire edges(), shortest paths(), similarity dice(), similarity inverse log weighted(), similarity jaccard(), simplify(), strength(), subcomponent(), subgraph edges(), subisomorphic lad(), subisomorphic vf2(), successors(), to directed(), to prufer(), to undirected(), topological sorting(), transitivity local undirected(), transitivity undirected(), unfold tree(), vcount(), vertex attributes(), vertex connectivity(), write dot(), write edgelist(), write gml(), write graphml(), write leda(), write lgl(), write ncol(), write pajek()

## Inherited from object

### 1.5.2 Properties

Name	Description
VS	The vertex sequence of the graph
es	The edge sequence of the graph
_as_parameter_	
Inherited from object	
class	

#### 1.5.3 Class Variables

Name	Description
iter	Value: None
hash	Value: None

## 1.6 Class VertexSeq

```
object — igraph.drawing.graph.VertexSeq — igraph.VertexSeq
```

Class representing a sequence of vertices in the graph.

This class is most easily accessed by the vs field of the Graph object, which returns an ordered sequence of all vertices in the graph. The vertex sequence can be refined by invoking the VertexSeq.select() method. VertexSeq.select() can also be accessed by simply calling the VertexSeq object.

An alternative way to create a vertex sequence referring to a given graph is to use the constructor directly:

```
>>> g = Graph.Full(3)
>>> vs = VertexSeq(g)
>>> restricted_vs = VertexSeq(g, [0, 1])
```

The individual vertices can be accessed by indexing the vertex sequence object. It can be used as an iterable as well, or even in a list comprehension:

```
>>> g=Graph.Full(3)
>>> for v in g.vs:
... v["value"] = v.index ** 2
...
>>> [v["value"] ** 0.5 for v in g.vs]
[0.0, 1.0, 2.0]
```

The vertex set can also be used as a dictionary where the keys are the attribute names. The values corresponding to the keys are the values of the given attribute for every vertex selected by the sequence.

```
>>> g=Graph.Full(3)
>>> for idx, v in enumerate(g.vs):
... v["weight"] = idx*(idx+1)
...
>>> g.vs["weight"]
[0, 2, 6]
>>> g.vs.select(1,2)["weight"] = [10, 20]
>>> g.vs["weight"]
[0, 10, 20]
```

If you specify a sequence that is shorter than the number of vertices in the VertexSeq, the sequence is reused:

```
>>> g = Graph.Tree(7, 2)
>>> g.vs["color"] = ["red", "green"]
>>> g.vs["color"]
['red', 'green', 'red', 'green', 'red', 'green', 'red']
```

You can even pass a single string or integer, it will be considered as a sequence of length 1:

```
>>> g.vs["color"] = "red"
>>> g.vs["color"]
['red', 'red', 'red', 'red', 'red', 'red']
```

Some methods of the vertex sequences are simply proxy methods to the corresponding methods in the Graph object. One such example is VertexSeq.degree():

```
>>> g=Graph.Tree(7, 2)
>>> g.vs.degree()
[2, 3, 3, 1, 1, 1, 1]
>>> g.vs.degree() == g.degree()
```

#### 1.6.1 Methods

# attributes(self)

Returns the list of all the vertex attributes in the graph associated to this vertex sequence.

```
find(self, *args, **kwds)
```

Returns the first vertex of the vertex sequence that matches some criteria.

The selection criteria are equal to the ones allowed by VertexSeq.select. See VertexSeq.select for more details.

For instance, to find the first vertex with name foo in graph g:

```
>>> g.vs.find(name="foo") #doctest:+SKIP
```

To find an arbitrary isolated vertex:

```
>>> g.vs.find(_degree=0) #doctest:+SKIP
```

## Return Value

Vertex

Overrides: igraph.drawing.graph.VertexSeq.find

# **select**(self, \*args, \*\*kwds)

Selects a subset of the vertex sequence based on some criteria

The selection criteria can be specified by the positional and the keyword arguments. Positional arguments are always processed before keyword arguments.

- If the first positional argument is None, an empty sequence is returned.
- If the first positional argument is a callable object, the object will be called for every vertex in the sequence. If it returns True, the vertex will be included, otherwise it will be excluded.
- If the first positional argument is an iterable, it must return integers and they will be considered as indices of the current vertex set (NOT the whole vertex set of the graph the difference matters when one filters a vertex set that has already been filtered by a previous invocation of VertexSeq.select(). In this case, the indices do not refer directly to the vertices of the graph but to the elements of the filtered vertex sequence.
- If the first positional argument is an integer, all remaining arguments are expected to be integers. They are considered as indices of the current vertex set again.

Keyword arguments can be used to filter the vertices based on their attributes. The name of the keyword specifies the name of the attribute and the filtering operator, they should be concatenated by an underscore (\_) character. Attribute names can also contain underscores, but operator names don't, so the operator is always the largest trailing substring of the keyword name that does not contain an underscore. Possible operators are:

- eq: equal to
- ne: not equal to
- 1t: less than
- gt: greater than
- le: less than or equal to
- ge: greater than or equal to
- in: checks if the value of an attribute is in a given list
- notin: checks if the value of an attribute is not in a given list

For instance, if you want to filter vertices with a numeric age property larger than 200, you have to write:

```
>>> g.vs.select(age_gt=200) #doctest: +SKIP
```

Similarly, to filter vertices whose type is in a list of predefined types:

```
>>> list_of_types = ["HR", "Finance", "Management"]
>>> g.vs.select(type_in=list_of_types) #doctest: +SKIP
```

If the operator is omitted, it defaults to eq. For instance, the following selector selects vertices whose cluster property equals to 2:

**call** (self, \*args, \*\*kwds)

Shorthand notation to select()

This method simply passes all its arguments to VertexSeq.select().

betweenness(\*args, \*\*kwds)

Proxy method to Graph.betweenness()

This method calls the betweenness() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.betweenness() for details.

bibcoupling(\*args, \*\*kwds)

Proxy method to Graph.bibcoupling()

This method calls the bibcoupling() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.bibcoupling() for details.

closeness(\*args, \*\*kwds)

Proxy method to Graph.closeness()

This method calls the closeness() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.closeness() for details.

cocitation(\*args, \*\*kwds)

Proxy method to Graph.cocitation()

This method calls the cocitation() method of the Graph class restricted to this sequence, and returns the result.

**See Also:** Graph.cocitation() for details.

constraint(\*args, \*\*kwds)

Proxy method to Graph.constraint()

This method calls the constraint() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.constraint() for details.

## degree(\*args, \*\*kwds)

Proxy method to Graph.degree()

This method calls the degree() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.degree() for details.

# $\mathbf{delete}(*args, **kwds)$

Proxy method to Graph.delete\_vertices()

This method calls the delete\_vertices() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.delete\_vertices() for details.

# diversity(\*args, \*\*kwds)

Proxy method to Graph.diversity()

This method calls the diversity() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.diversity() for details.

## eccentricity(\*args, \*\*kwds)

Proxy method to Graph.eccentricity()

This method calls the eccentricity() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.eccentricity() for details.

# get shortest paths(\*args, \*\*kwds)

Proxy method to Graph.get\_shortest\_paths()

This method calls the get\_shortest\_paths() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.get\_shortest\_paths() for details.

## indegree(\*args, \*\*kwds)

Proxy method to Graph.indegree()

This method calls the indegree() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.indegree() for details.

# is minimal separator(\*args, \*\*kwds)

Proxy method to Graph.is\_minimal\_separator()

This method calls the is\_minimal\_separator() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.is\_minimal\_separator() for details.

# is separator(\*args, \*\*kwds)

Proxy method to Graph.is\_separator()

This method calls the is\_separator() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.is separator() for details.

# isoclass(\*args, \*\*kwds)

Proxy method to Graph.isoclass()

This method calls the isoclass() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.isoclass() for details.

# maxdegree(\*args, \*\*kwds)

Proxy method to Graph.maxdegree()

This method calls the maxdegree() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.maxdegree() for details.

## outdegree(\*args, \*\*kwds)

Proxy method to Graph.outdegree()

This method calls the outdegree() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.outdegree() for details.

# pagerank(\*args, \*\*kwds)

Proxy method to Graph.pagerank()

This method calls the pagerank() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.pagerank() for details.

# personalized pagerank(\*args, \*\*kwds)

Proxy method to Graph.personalized\_pagerank()

This method calls the personalized\_pagerank() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.personalized\_pagerank() for details.

## shortest paths(\*args, \*\*kwds)

Proxy method to Graph.shortest\_paths()

This method calls the shortest\_paths() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.shortest paths() for details.

## similarity dice(\*args, \*\*kwds)

Proxy method to Graph.similarity\_dice()

This method calls the similarity\_dice() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.similarity\_dice() for details.

similarity jaccard(\*args, \*\*kwds)

Proxy method to Graph.similarity\_jaccard()

This method calls the similarity\_jaccard() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.similarity jaccard() for details.

```
subgraph(*args, **kwds)
```

Proxy method to Graph.subgraph()

This method calls the subgraph() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.subgraph() for details.

# $Inherited\ from\ igraph.drawing.graph.VertexSeq$

```
__delitem__(), __getitem__(), __init__(), __len__(), __new__(), __setitem__(), attribute names(), get attribute values(), set attribute values()
```

# Inherited from object

#### 1.6.2 Properties

Name	Description	
Inherited from igraph.drawing.graph.VertexSeq		
graph, indices		
Inherited from object		
class		

### 1.7 Class EdgeSeq

```
object — igraph.EdgeSeq — igraph.EdgeSeq
```

Class representing a sequence of edges in the graph.

This class is most easily accessed by the es field of the Graph object, which returns an ordered sequence of all edges in the graph. The edge sequence can be refined by invoking

the EdgeSeq.select() method. EdgeSeq.select() can also be accessed by simply calling the EdgeSeq object.

An alternative way to create an edge sequence referring to a given graph is to use the constructor directly:

```
>>> g = Graph.Full(3)
>>> es = EdgeSeq(g)
>>> restricted_es = EdgeSeq(g, [0, 1])
```

The individual edges can be accessed by indexing the edge sequence object. It can be used as an iterable as well, or even in a list comprehension:

```
>>> g=Graph.Full(3)
>>> for e in g.es:
...    print e.tuple
...
(0, 1)
(0, 2)
(1, 2)
>>> [max(e.tuple) for e in g.es]
[1, 2, 2]
```

The edge sequence can also be used as a dictionary where the keys are the attribute names. The values corresponding to the keys are the values of the given attribute of every edge in the graph:

```
>>> g=Graph.Full(3)
>>> for idx, e in enumerate(g.es):
... e["weight"] = idx*(idx+1)
...
>>> g.es["weight"]
[0, 2, 6]
>>> g.es["weight"] = range(3)
>>> g.es["weight"]
[0, 1, 2]
```

If you specify a sequence that is shorter than the number of edges in the EdgeSeq, the sequence is reused:

```
>>> g = Graph.Tree(7, 2)
>>> g.es["color"] = ["red", "green"]
>>> g.es["color"]
['red', 'green', 'red', 'green', 'red', 'green']
```

You can even pass a single string or integer, it will be considered as a sequence of length 1:

```
>>> g.es["color"] = "red"
```

```
>>> g.es["color"]
['red', 'red', 'red', 'red', 'red']
```

Some methods of the edge sequences are simply proxy methods to the corresponding methods in the Graph object. One such example is EdgeSeq.is\_multiple():

```
>>> g=Graph(3, [(0,1), (1,0), (1,2)])
>>> g.es.is_multiple()
[False, True, False]
>>> g.es.is_multiple() == g.is_multiple()
True
```

### 1.7.1 Methods

# attributes(self)

Returns the list of all the edge attributes in the graph associated to this edge sequence.

```
\mathbf{find}(self, *args, **kwds)
```

Returns the first edge of the edge sequence that matches some criteria.

The selection criteria are equal to the ones allowed by VertexSeq.select. See VertexSeq.select for more details.

For instance, to find the first edge with weight larger than 5 in graph g:

```
>>> g.es.find(weight_gt=5) #doctest:+SKIP
```

## Return Value

Edge

Overrides: igraph. igraph.EdgeSeq.find

```
select(self, *args, **kwds)
```

Selects a subset of the edge sequence based on some criteria

The selection criteria can be specified by the positional and the keyword arguments. Positional arguments are always processed before keyword arguments.

- If the first positional argument is None, an empty sequence is returned.
- If the first positional argument is a callable object, the object will be called for every edge in the sequence. If it returns True, the edge will be included, otherwise it will be excluded.
- If the first positional argument is an iterable, it must return integers and they will be considered as indices of the current edge set (NOT the whole edge set of the graph the difference matters when one filters an edge set that has already been filtered by a previous invocation of EdgeSeq.select(). In this case, the indices do not refer directly to the edges of the graph but to the elements of the filtered edge sequence.
- If the first positional argument is an integer, all remaining arguments are expected to be integers. They are considered as indices of the current edge set again.

Keyword arguments can be used to filter the edges based on their attributes and properties. The name of the keyword specifies the name of the attribute and the filtering operator, they should be concatenated by an underscore (\_) character. Attribute names can also contain underscores, but operator names don't, so the operator is always the largest trailing substring of the keyword name that does not contain an underscore. Possible operators are:

- eq: equal to
- ne: not equal to
- 1t: less than
- gt: greater than
- le: less than or equal to
- ge: greater than or equal to
- in: checks if the value of an attribute is in a given list
- notin: checks if the value of an attribute is not in a given list

For instance, if you want to filter edges with a numeric weight property larger than 50, you have to write:

```
>>> g.es.select(weight_gt=50) #doctest: +SKIP
```

Similarly, to filter edges whose type is in a list of predefined types:

```
>>> list_of_types = ["inhibitory", "excitatory"]
>>> g.es.select(type_in=list_of_types) #doctest: +SKIP
```

If the operator is omitted, it defaults to eq. For instance, the following selector selects edges whose type property is intracluster:

**call** (self, \*args, \*\*kwds)

Shorthand notation to select()

This method simply passes all its arguments to EdgeSeq.select().

count multiple(\*args, \*\*kwds)

Proxy method to Graph.count\_multiple()

This method calls the count\_multiple() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.count\_multiple() for details.

 $\mathbf{delete}(*args, **kwds)$ 

Proxy method to Graph.delete\_edges()

This method calls the delete\_edges() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.delete edges() for details.

edge betweenness(\*args, \*\*kwds)

Proxy method to Graph.edge\_betweenness()

This method calls the edge\_betweenness() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.edge betweenness() for details.

 $is\_loop(*args, **kwds)$ 

Proxy method to Graph.is\_loop()

This method calls the is\_loop() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.is\_loop() for details.

is multiple(\*args, \*\*kwds)

Proxy method to Graph.is\_multiple()

This method calls the is\_multiple() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.is multiple() for details.

is mutual(\*args, \*\*kwds)

Proxy method to Graph.is\_mutual()

This method calls the is\_mutual() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.is\_mutual() for details.

# subgraph(\*args, \*\*kwds)

Proxy method to Graph.subgraph\_edges()

This method calls the subgraph\_edges() method of the Graph class restricted to this sequence, and returns the result.

See Also: Graph.subgraph edges() for details.

# Inherited from igraph. igraph. EdgeSeq

```
__delitem__(), __getitem__(), __init__(), __len__(), __new__(), __setitem__(), attribute names(), get attribute values(), is all(), set attribute values()
```

# Inherited from object

#### 1.7.2 Properties

Name Description		
Inherited from igraphigrap	h.EdgeSeq	
graph, indices		
Inherited from object		
class		

# 1.8 Class ARPACKOptions

Class representing the parameters of the ARPACK module.

ARPACK is a Fortran implementation of the implicitly restarted Arnoldi method, an algorithm for calculating some of the eigenvalues and eigenvectors of a given matrix. igraph uses this package occasionally, and this class can be used to fine-tune the behaviour of ARPACK in such cases.

The class has several attributes which are not documented here, since they are usually of marginal use to the ordinary user. See the source code of the original ARPACK Fortran package (especially the file dsaupd.f) for a detailed explanation of the parameters. Only the most basic attributes are explained here. Most of them are read only unless stated otherwise.

- bmat: type of the eigenproblem solved. 'I' means standard eigenproblem  $(A^*x = lambda^*x)$ , 'G' means generalized eigenproblem  $(A^*x = lambda^*B^*x)$ .
- n: dimension of the eigenproblem
- tol: precision. If less than or equal to zero, the standard machine precision is used as computed by the LAPACK utility called dlamch. This can be modified.
- mxiter: maximum number of update iterations to take. This can be modified. You can also use maxiter.
- iter: actual number of update iterations taken
- numop: total number of OP\*x operations
- numopb: total number of B\*x operations if bmat is 'G'
- numreo: total number of steps of re-orthogonalization

#### 1.8.1 Methods

$\_\_\mathbf{new}\_\_(T, S,)$
Return Value
a new object with type S, a subtype of T
Overrides: objectnew

$\_\_\mathbf{str}\_\_(x)$	
$\operatorname{str}(\mathrm{x})$	
Overrides: objectstr	

# Inherited from object

```
\label{eq:continuous} $$\_\_delattr\_(), \_\_format\_(), \_\_getattribute\_(), \_\_hash\_(), \_\_init\_(), \_\_reduce\_(), \_\_reduce\_ex\_(), \_\_repr\_(), \_\_setattr\_(), \_\_sizeof\_(), \_\_subclasshook\_()
```

#### 1.8.2 Properties

Name	Description
Inherited from object	
class	

Class DFSIter Package igraph

1	0	$O_{1}$	DECL
т.	.9	Class	BFSIter



igraph BFS iterator object

### 1.9.1 Methods

$\_\_$ iter $\_\_(x)$	
iter(x)	

 $\mathbf{next}(x)$ 

## Return Value

the next value, or raise StopIteration

## Inherited from object

### 1.9.2 Properties

Name	Description
Inherited from object	
class	

### 1.10 Class DFSIter

object igraph.DFSIter

igraph DFS iterator object

# 1.10.1 Methods

iter(x)	
ter(x)	

 $\mathbf{next}(x)$ 

Return Value

the next value, or raise StopIteration

Class Edge Package igraph

# Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __init__(), __new__(), __reduce__ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

#### 1.10.2 Properties

Name	Description
Inherited from object	
class	

# 1.11 Class Edge

Class representing a single edge in a graph.

The edge is referenced by its index, so if the underlying graph changes, the semantics of the edge object might change as well (if the edge indices are altered in the original graph).

The attributes of the edge can be accessed by using the edge as a hash:

```
>>> e ["weight"] = 2 #doctest: +SKIP
>>> print e ["weight"] #doctest: +SKIP
```

### 1.11.1 Methods

```
\underline{\phantom{a}}_{\text{del x[y]}}^{\text{del tem}}\underline{\phantom{a}}_{(x, y)}
```

```
 \frac{\mathbf{eq}_{-}(x, y)}{\mathbf{x} = -\mathbf{y}}
```

```
 \frac{\mathbf{ge}_{-}(x, y)}{x > = y}
```

Class Edge Package igraph

$\begin{bmatrix} \_\_\mathbf{gt}\_\_(x, y) \end{bmatrix}$	z, y)	
x>y		

 $\frac{\_\_hash\_\_(x)}{hash(x)}$ Overrides: object. \_\_hash\_\_

 $\frac{\mathbf{le}_{-}(x, y)}{\mathbf{x} < = \mathbf{y}}$ 

 $\frac{-\operatorname{len}_{-}(x)}{\operatorname{len}(x)}$ 

 $\frac{-\mathbf{lt}_{-}(x, y)}{x < y}$ 

 $\frac{\mathbf{ne}_{-}(x, y)}{x! = y}$ 

 $\frac{\_\_setitem}{x[i]=y} (x, i, y)$ 

# attribute names()

Returns the list of edge attribute names

Return Value

list

# attributes()

Returns a dict of attribute names and values for the edge

Return Value

dict

Class Edge Package igraph

# count multiple(...)

Proxy method to Graph.count\_multiple()

This method calls the count\_multiple method of the Graph class with this edge as the first argument, and returns the result.

See Also: Graph.count multiple() for details.

### delete(...)

Proxy method to Graph.delete\_edges()

This method calls the delete\_edges method of the Graph class with this edge as the first argument, and returns the result.

See Also: Graph.delete\_edges() for details.

# is loop(...)

Proxy method to Graph.is\_loop()

This method calls the is\_loop method of the Graph class with this edge as the first argument, and returns the result.

See Also: Graph.is\_loop() for details.

### is multiple(...)

Proxy method to Graph.is\_multiple()

This method calls the is\_multiple method of the Graph class with this edge as the first argument, and returns the result.

See Also: Graph.is multiple() for details.

### is mutual(...)

Proxy method to Graph.is\_mutual()

This method calls the is\_mutual method of the Graph class with this edge as the first argument, and returns the result.

See Also: Graph.is\_mutual() for details.

# update attributes(E, \*\*F)

Updates the attributes of the edge from dict/iterable E and F.

If E has a keys() method, it does: for k in E: self[k] = E[k]. If E lacks a keys() method, it does: for (k, v) in E: self[k] = v. In either case, this is followed by: for k in F: self[k] = F[k].

This method thus behaves similarly to the update() method of Python dictionaries.

# Return Value

None

# Inherited from object

$\_$ _delattr $\_$ _(),	format();	,getattribute	(),i	$\operatorname{nit}\_\_(),$ $\_$	_new()
reduce(),	reduce_ex_	_(),setattr_	_(),siz	eof(), _	$_{-}$ str $_{-}(),$
subclasshook	()				

### 1.11.2 Properties

Name	Description
graph	The graph the edge belongs to
index	Index of this edge
source	Source vertex index of this edge
source_vertex	Source vertex of this edge
target	Target vertex index of this edge
target_vertex	Target vertex of this edge
tuple	Source and target vertex index of this edge as a
	tuple
vertex_tuple	Source and target vertex of this edge as a tuple
Inherited from object	
class	

## 1.12 Class GraphBase

object igraph.GraphBase

Known Subclasses: igraph.Graph

Low-level representation of a graph.

Don't use it directly, use igraph.Graph instead.

### 1.12.1 Methods

# $Adjacency(matrix, mode = ADJ_DIRECTED)$

Generates a graph from its adjacency matrix.

### **Parameters**

matrix: the adjacency matrix

mode: the mode to be used. Possible values are:

• ADJ\_DIRECTED - the graph will be directed and a matrix element gives the number of edges between two vertex.

- ADJ\_UNDIRECTED alias to ADJ\_MAX for convenience.
- ADJ\_MAX undirected graph will be created and the number of edges between vertex i and j is max(A(i,j), A(j,i))
- ADJ\_MIN like ADJ\_MAX, but with min(A(i,j), A(j,i))
- ADJ\_PLUS like ADJ\_MAX, but with A(i,j) + A(j,i)
- ADJ\_UPPER undirected graph with the upper right triangle of the matrix (including the diagonal)
- ADJ\_LOWER undirected graph with the lower left triangle of the matrix (including the diagonal)

These values can also be given as strings without the ADJ prefix.

Generates a graph based on asymmetric vertex types and connection probabilities.

This is the asymmetric variant of Graph.Preference. A given number of vertices are generated. Every vertex is assigned to an "incoming" and an "outgoing" vertex type according to the given joint type probabilities. Finally, every vertex pair is evaluated and a directed edge is created between them with a probability depending on the "outgoing" type of the source vertex and the "incoming" type of the target vertex.

### **Parameters**

n: the number of vertices in the graph

type\_dist\_matrix: matrix giving the joint distribution of vertex

types

pref\_matrix:
matrix giving the connection probabilities for

different vertex types.

attribute: the vertex attribute name used to store the

vertex types. If None, vertex types are not

stored.

loops: whether loop edges are allowed.

### Atlas(idx)

Generates a graph from the Graph Atlas.

#### **Parameters**

idx: The index of the graph to be generated. Indices start from zero, graphs are listed:

- 1. in increasing order of number of vertices;
- 2. for a fixed number of vertices, in increasing order of the number of edges;
- 3. for fixed numbers of vertices and edges, in increasing order of the degree sequence, for example 111223 < 112222;
- 4. for fixed degree sequence, in increasing number of automorphisms.

**Reference:** An Atlas of Graphs by Ronald C. Read and Robin J. Wilson, Oxford University Press, 1998.

 ${f Barabasi}(n,\ m,\ outpref={\sf False},\ directed={\sf False},\ power=1,\ zero\_appeal=1,\ implementation="psumtree",\ start\ from={\sf None})$ 

Generates a graph based on the Barabasi-Albert model.

#### **Parameters**

n: the number of vertices

m: either the number of outgoing edges generated for

each vertex or a list containing the number of outgoing edges for each vertex explicitly.

outpref: True if the out-degree of a given vertex should

also increase its citation probability (as well as its

in-degree), but it defaults to False.

directed: True if the generated graph should be directed

(default: False).

power: the power constant of the nonlinear model. It can

be omitted, and in this case the usual linear

model will be used.

zero\_appeal: the attractivity of vertices with degree zero.

implementation: the algorithm to use to generate the network.

Possible values are:

- "bag": the algorithm that was the default in igraph before 0.6. It works by putting the ids of the vertices into a bag (multiset) exactly as many times as their in-degree, plus once more. The required number of cited vertices are then drawn from the bag with replacement. It works only for power=1 and zero appeal=1.
- "psumtree": this algorithm uses a partial prefix-sum tree to generate the graph. It does not generate multiple edges and it works for any values of *power* and *zero\_appeal*.
- "psumtree\_multiple": similar to "psumtree", but it will generate multiple edges as well. igraph before 0.6 used this algorithm for *powers* other than 1.

start\_from: if given and not None, this must be another Graph

object. igraph will use this graph as a starting point for the preferential attachment model.

**Reference:** Barabasi, A-L and Albert, R. 1999. Emergence of scaling in random networks. Science, 286 509-512.

# De Bruijn(m, n)

Generates a de Bruijn graph with parameters (m, n)

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  $m \hat{n}$  vertices and even more edges, so probably you don't want to supply too big numbers for m and n.

### **Parameters**

 $\mathtt{m}\colon$  the size of the alphabet

n: the length of the strings

# Degree Sequence(out, in=None, method="simple")

Generates a graph with a given degree sequence.

#### **Parameters**

out:

the out-degree sequence for a directed graph. If the in-degree sequence is omitted, the generated graph will be undirected, so this will be the in-degree sequence as well

in:

the in-degree sequence for a directed graph. If omitted, the generated graph will be undirected.

method: the generation method to be used. One of the following:

- "simple" simple generator that sometimes generates loop edges and multiple edges. The generated graph is not guaranteed to be connected.
- "no\_multiple" similar to "simple" but avoids the generation of multiple and loop edges at the expense of increased time complexity. The method will re-start the generation every time it gets stuck in a configuration where it is not possible to insert any more edges without creating loops or multiple edges, and there is no upper bound on the number of iterations, but it will succeed eventually if the input degree sequence is graphical and throw an exception if the input degree sequence is not graphical.
- "v1" a more sophisticated generator that can sample undirected, connected simple graphs uniformly. It uses Monte-Carlo methods to randomize the graphs. This generator should be favoured if undirected and connected graphs are to be generated and execution time is not a concern. igraph uses the original implementation of Fabien Viger; see the following URL and the paper cited on it for the details of the algorithm:

http://www-rp.lip6.fr/~latapy/FV/generation.html.

# Erdos Renyi(n, p, m, directed=False, loops=False)

Generates a graph based on the Erdos-Renyi model.

#### **Parameters**

n: the number of vertices.

p: the probability of edges. If given, m must be missing.

m: the number of edges. If given, p must be missing.

directed: whether to generate a directed graph.

loops: whether self-loops are allowed.

## $\mathbf{Establishment}(n, k, type\_dist, pref\_matrix, directed = \mathbf{False})$

Generates a graph based on a simple growing model with vertex types.

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.

#### **Parameters**

n: the number of vertices in the graph

k: the number of connections tried in each step

type\_dist: list giving the distribution of vertex types

pref\_matrix: matrix (list of lists) giving the connection

probabilities for different vertex types

directed: whether to generate a directed graph.

### Famous(name)

Generates a famous graph based on its name.

Several famous graphs are known to igraph including (but not limited to) the Chvatal graph, the Petersen graph or the Tutte graph. This method generates one of them based on its name (case insensitive). See the documentation of the C interface of igraph for the names available: http://igraph.org/doc/c.

### **Parameters**

name: the name of the graph to be generated.

# Forest Fire(n, fw prob, bw factor=0.0, ambs=1, directed=False)

Generates a graph based on the forest fire model

The forest fire model is a growing graph model. In every time step, a new vertex is added to the graph. The new vertex chooses an ambassador (or more than one if ambs>1) and starts a simulated forest fire at its ambassador(s). The fire spreads through the edges. The spreading probability along an edge is given by  $fw\_prob$ . The fire may also spread backwards on an edge by probability  $fw\_prob * bw\_factor$ . When the fire ended, the newly added vertex connects to the vertices "burned" in the previous fire.

### **Parameters**

n: the number of vertices in the graph

fw\_prob: forward burning probability

bw\_factor: ratio of backward and forward burning probability

ambs: number of ambassadors chosen in each step

directed: whether the graph will be directed

## Full(n, directed=False, loops=False)

Generates a full graph (directed or undirected, with or without loops).

## **Parameters**

n: the number of vertices.

directed: whether to generate a directed graph.

loops: whether self-loops are allowed.

### Full Citation(n, directed = False)

Generates a full citation graph

A full citation graph is a graph where the vertices are indexed from 0 to n-1 and vertex i has a directed edge towards all vertices with an index less than i.

### **Parameters**

n: the number of vertices.

directed: whether to generate a directed graph.

# **Growing** Random(n, m, directed=False, citation=False)

Generates a growing random graph.

## **Parameters**

n: The number of vertices in the graph

m: The number of edges to add in each step (after adding a

new vertex)

directed: whether the graph should be directed.

citation: whether the new edges should originate from the most

recently added vertex.

## Isoclass(n, class, directed=False)

Generates a graph with a given isomorphism class.

#### **Parameters**

n: the number of vertices in the graph (3 or 4)

class: the isomorphism class

directed: whether the graph should be directed.

# K Regular(n, k, directed=False, multiple=False)

Generates a k-regular random graph

A k-regular random graph is a random graph where each vertex has degree k. If the graph is directed, both the in-degree and the out-degree of each vertex will be k.

# Parameters

n: The number of vertices in the graph

k: The degree of each vertex if the graph is undirected, or

the in-degree and out-degree of each vertex if the graph

is directed

directed: whether the graph should be directed.

multiple: whether it is allowed to create multiple edges.

# $\mathbf{Kautz}(m, n)$

Generates a Kautz graph with parameters (m, n)

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.

#### **Parameters**

m: the size of the alphabet minus one

n: the length of the strings minus one

# LCF(n, shifts, repeats)

Generates a graph from LCF notation.

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 <a href="http://mathworld.wolfram.com/LCFNotation.html">http://mathworld.wolfram.com/LCFNotation.html</a> for details.

#### **Parameters**

n: the number of vertices

shifts: the shifts in a list or tuple

repeats: the number of repeats

### Lattice(dim, nei=1, directed=False, mutual=True, circular=True)

Generates a regular lattice.

### **Parameters**

dim: list with the dimensions of the lattice

nei: value giving the distance (number of steps) within which

two vertices will be connected.

directed: whether to create a directed graph.

mutual: whether to create all connections as mutual in case of a

directed graph.

circular: whether the generated lattice is periodic.

 $\begin{aligned} \mathbf{Preference}(n, \ type\_\ dist, \ pref\_\ matrix, \ attribute = \mathtt{None}, \ directed = \mathtt{False}, \\ loops = \mathtt{False}) \end{aligned}$ 

Generates a graph based on vertex types and connection probabilities.

This is practically the nongrowing variant of Graph.Establishment. A given number of vertices are generated. Every vertex is assigned to a vertex type according to the given type probabilities. Finally, every vertex pair is evaluated and an edge is created between them with a probability depending on the types of the vertices involved.

#### **Parameters**

n: the number of vertices in the graph

type\_dist: list giving the distribution of vertex types

pref\_matrix: matrix giving the connection probabilities for

different vertex types.

attribute: the vertex attribute name used to store the vertex

types. If None, vertex types are not stored.

directed: whether to generate a directed graph.

loops: whether loop edges are allowed.

# Read DIMACS(f, directed=False)

Reads a graph from a file conforming to the DIMACS minimum-cost flow file format.

For the exact description of the format, see http://lpsolve.sourceforge.net/5.5/DIMACS.htm

Restrictions compared to the official description of the format:

- igraph's DIMACS reader requires only three fields in an arc definition, describing the edge's source and target node and its capacity.
- Source vertices are identified by 's' in the FLOW field, target vertices are identified by 't'.
- Node indices start from 1. Only a single source and target node is allowed.

### **Parameters**

f: the name of the file or a Python file handle

directed: whether the generated graph should be directed.

# Return Value

the generated graph, the source and the target of the flow and the edge capacities in a tuple

## Read DL(f, directed=True)

Reads an UCINET DL file and creates a graph based on it.

#### **Parameters**

f: the name of the file or a Python file handle

directed: whether the generated graph should be directed.

## **Read** Edgelist(f, directed=True)

Reads an edge list from a file and creates a graph based on it.

Please note that the vertex indices are zero-based. A vertex of zero degree will be created for every integer that is in range but does not appear in the edgelist.

#### **Parameters**

f: the name of the file or a Python file handle

directed: whether the generated graph should be directed.

# Read GML(f)

Reads a GML file and creates a graph based on it.

## **Parameters**

f: the name of the file or a Python file handle

# Read GraphDB(f, directed=False)

Reads a GraphDB format file and creates a graph based on it.

GraphDB is a binary format, used in the graph database for isomorphism testing (see http://amalfi.dis.unina.it/graph/).

## **Parameters**

f: the name of the file or a Python file handle

directed: whether the generated graph should be directed.

# Read GraphML(f, directed=True, index=0)

Reads a GraphML format file and creates a graph based on it.

#### **Parameters**

f: the name of the file or a Python file handle

index: if the GraphML file contains multiple graphs, specifies the one that should be loaded. Graph indices start from zero, so if you want to load the first graph, specify 0 here.

Read Lgl(f, names=True, weights="if\_present", directed=True)

Reads an .lgl file used by LGL.

It is also useful for creating graphs from "named" (and optionally weighted) edge lists.

This format is used by the Large Graph Layout program. See the documentation of  $LGL^a$  regarding the exact format description.

LGL originally cannot deal with graphs containing multiple or loop edges, but this condition is not checked here, as igraph is happy with these.

### **Parameters**

f: the name of the file or a Python file handle

names: If True, the vertex names are added as a vertex attribute

called 'name'.

weights: If True, the edge weights are added as an edge attribute

called 'weight', even if there are no weights in the file. If False, the edge weights are never added, even if they are present. "auto" or "if\_present" means that weights are added if there is at least one weighted edge in the

input file, but they are not added otherwise.

directed: whether the graph being created should be directed

 $<sup>^</sup>a$ http://bioinformatics.icmb.utexas.edu/lgl/

# Read Ncol(f, names=True, weights="if\_present", directed=True)

Reads an .ncol file used by LGL.

It is also useful for creating graphs from "named" (and optionally weighted) edge lists.

This format is used by the Large Graph Layout program. See the documentation of  $LGL^a$  regarding the exact format description.

LGL originally cannot deal with graphs containing multiple or loop edges, but this condition is not checked here, as igraph is happy with these.

### **Parameters**

f: the name of the file or a Python file handle

names: If True, the vertex names are added as a vertex attribute

called 'name'.

weights: If True, the edge weights are added as an edge attribute

called 'weight', even if there are no weights in the file. If False, the edge weights are never added, even if they are present. "auto" or "if\_present" means that weights are added if there is at least one weighted edge in the

input file, but they are not added otherwise.

directed: whether the graph being created should be directed

### Read Pajek(f)

Reads a Pajek format file and creates a graph based on it.

### **Parameters**

f: the name of the file or a Python file handle

 $<sup>^</sup>a$ http://bioinformatics.icmb.utexas.edu/lgl/

# Recent Degree(n, m, window, outpref=False, directed=False, power=1)

Generates a graph based on a stochastic model where the probability of an edge gaining a new node is proportional to the edges gained in a given time window.

#### **Parameters**

n: the number of vertices

m: either the number of outgoing edges generated for each

vertex or a list containing the number of outgoing edges

for each vertex explicitly.

window: size of the window in time steps

outpref: True if the out-degree of a given vertex should also

increase its citation probability (as well as its in-degree),

but it defaults to False.

directed: True if the generated graph should be directed (default:

False).

power: the power constant of the nonlinear model. It can be

omitted, and in this case the usual linear model will be

used.

## Ring(n, directed=False, mutual=False, circular=True)

Generates a ring graph.

## **Parameters**

n: the number of vertices in the ring

directed: whether to create a directed ring.

mutual: whether to create mutual edges in a directed ring.

circular: whether to create a closed ring.

# SBM(n, pref matrix, block sizes, directed=False, loops=False)

Generates a graph based on a stochastic blockmodel.

A given number of vertices are generated. Every vertex is assigned to a vertex type according to the given block sizes. Vertices of the same type will be assigned consecutive vertex IDs. Finally, every vertex pair is evaluated and an edge is created between them with a probability depending on the types of the vertices involved. The probabilities are taken from the preference matrix.

### **Parameters**

n: the number of vertices in the graph

pref\_matrix: matrix giving the connection probabilities for

different vertex types.

block\_sizes: list giving the number of vertices in each block; must

sum up to n.

directed: whether to generate a directed graph.

loops: whether loop edges are allowed.

# Star(n, mode="undirected", center=0)

Generates a star graph.

## **Parameters**

n: the number of vertices in the graph

mode: Gives the type of the star graph to create. Should be either

"in", "out", "mutual" or "undirected"

center: Vertex ID for the central vertex in the star.

 $Static\_Fitness(m, fitness\_out, fitness\_in=None, loops=False, multiple=False)$ 

Generates a non-growing graph with edge probabilities proportional to node fitnesses.

The algorithm randomly selects vertex pairs and connects them until the given number of edges are created. Each vertex is selected with a probability proportional to its fitness; for directed graphs, a vertex is selected as a source proportional to its out-fitness and as a target proportional to its in-fitness.

#### **Parameters**

m: the number of edges in the graph

fitness\_out: a numeric vector with non-negative entries, one for

each vertex. These values represent the fitness scores (out-fitness scores for directed graphs). fitness is an

alias of this keyword argument.

fitness\_in: a numeric vector with non-negative entries, one for

each vertex. These values represent the in-fitness scores for directed graphs. For undirected graphs,

this argument must be None.

loops: whether loop edges are allowed.

multiple: whether multiple edges are allowed.

# Return Value

a directed or undirected graph with the prescribed power-law degree distributions.

Static\_Power\_Law(n, m, exponent\_out, exponent\_in=-1, loops=False, multiple=False, finite size correction=True)

Generates a non-growing graph with prescribed power-law degree distributions.

#### **Parameters**

n: the number of vertices in the graph

m: the number of edges in the graph

exponent\_out: the exponent of the out-degree

distribution, which must be between 2

and infinity (inclusive). When

exponent\_in is not given or negative,
the graph will be undirected and this

parameter specifies the degree

distribution. exponent is an alias to this

keyword argument.

exponent\_in: the exponent of the in-degree

distribution, which must be between 2 and infinity (inclusive) It can also be negative, in which case an undirected

graph will be generated.

loops: whether loop edges are allowed.

multiple: whether multiple edges are allowed.

finite\_size\_correction: whether to apply a finite-size correction

to the generated fitness values for exponents less than 3. See the paper of

Cho et al for more details.

### Return Value

a directed or undirected graph with the prescribed power-law degree distributions.

### Reference:

- Goh K-I, Kahng B, Kim D: Universal behaviour of load distribution in scale-free networks. Phys Rev Lett 87(27):278701, 2001.
- Cho YS, Kim JS, Park J, Kahng B, Kim D: Percolation transitions in scale-free networks under the Achlioptas process. Phys Rev Lett 103:135702, 2009.

## $\mathbf{Tree}(n, children, type = \mathsf{TREE\_UNDIRECTED})$

Generates a tree in which almost all vertices have the same number of children.

#### **Parameters**

n: the number of vertices in the graph

children: the number of children of a vertex in the graph

type: determines whether the tree should be directed, and if

this is the case, also its orientation. Must be one of

TREE\_IN, TREE\_OUT and TREE\_UNDIRECTED.

# Watts Strogatz(dim, size, nei, p, loops=False, multiple=False)

### **Parameters**

dim: the dimension of the lattice

size: the size of the lattice along all dimensions

nei: value giving the distance (number of steps) within which

two vertices will be connected.

p: rewiring probability

loops: specifies whether loop edges are allowed

multiple: specifies whether multiple edges are allowed

See Also: Lattice(), rewire(), rewire\_edges() if more flexibility is needed

Reference: Duncan J Watts and Steven H Strogatz: Collective dynamics of

small world networks, Nature 393, 440-442, 1998

Generates a graph from its adjacency matrix.

#### **Parameters**

matrix: the adjacency matrix

mode: the mode to be used. Possible values are:

- ADJ\_DIRECTED the graph will be directed and a matrix element gives the number of edges between two vertex.
- ADJ\_UNDIRECTED alias to ADJ\_MAX for convenience.
- ADJ\_MAX undirected graph will be created and the number of edges between vertex i and j is max(A(i,j), A(j,i))
- ADJ\_MIN like ADJ\_MAX, but with min(A(i,j), A(j,i))
- ADJ\_PLUS like ADJ\_MAX, but with A(i,j) + A(j,i)
- ADJ\_UPPER undirected graph with the upper right triangle of the matrix (including the diagonal)
- ADJ\_LOWER undirected graph with the lower left triangle of the matrix (including the diagonal)

These values can also be given as strings without the ADJ prefix.

attr: the name of the edge attribute that stores the edge weights.

loops: whether to include loop edges. When False, the diagonal of the adjacency matrix will be ignored.

delitem_	(x, y)		
del x[y]			

```
__init__(...)

x.__init__(...) initializes x; see help(type(x)) for signature

Overrides: object.__init__
```

```
\_\_ new\_\_ ( T, \overline{S}, ...)
```

## Return Value

a new object with type S, a subtype of T

Overrides: object.\_\_new\_\_

```
\frac{-\operatorname{str}_{-}(x)}{\operatorname{str}(x)} Overrides: object.__str__
```

# add edges(es)

Adds edges to the graph.

### **Parameters**

es: the list of edges to be added. Every edge is represented with a tuple, containing the vertex IDs of the two endpoints. Vertices are enumerated from zero.

## add vertices(n)

Adds vertices to the graph.

#### **Parameters**

n: the number of vertices to be added

# all minimal st separators()

Returns a list containing all the minimal s-t separators of a graph.

A minimal separator is a set of vertices whose removal disconnects the graph, while the removal of any subset of the set keeps the graph connected.

# Return Value

a list where each item lists the vertex indices of a given minimal s-t separator.

**Reference:** Anne Berry, Jean-Paul Bordat and Olivier Cogis: Generating all the minimal separators of a graph. In: Peter Widmayer, Gabriele Neyer and Stephan Eidenbenz (eds.): Graph-theoretic concepts in computer science, 1665, 167–172, 1999. Springer.

## all st cuts(source, target)

Returns all the cuts between the source and target vertices in a directed graph.

This function lists all edge-cuts between a source and a target vertex. Every cut is listed exactly once.

### Parameters

source: the source vertex ID
target: the target vertex ID

### Return Value

a tuple where the first element is a list of lists of edge IDs representing a cut and the second element is a list of lists of vertex IDs representing the sets of vertices that were separated by the cuts.

Attention: this function has a more convenient interface in class Graph which wraps the result in a list of Cut objects. It is advised to use that.

# all st mincuts(source, target)

Returns all minimum cuts between the source and target vertices in a directed graph.

### **Parameters**

source: the source vertex ID
target: the target vertex ID

Attention: this function has a more convenient interface in class Graph which wraps the result in a list of Cut objects. It is advised to use that.

#### are connected (v1, v2)

Decides whether two given vertices are directly connected.

### Parameters

v1: the ID or name of the first vertex

v2: the ID or name of the second vertex

### Return Value

True if there exists an edge from v1 to v2, False otherwise.

### articulation points()

Returns the list of articulation points in the graph.

A vertex is an articulation point if its removal increases the number of connected components in the graph.

## assortativity(types1, types2=None, directed=True)

Returns the assortativity of the graph based on numeric properties of the vertices.

This coefficient is basically the correlation between the actual connectivity patterns of the vertices and the pattern expected from the disribution of the vertex types.

See equation (21) in Newman MEJ: Mixing patterns in networks, Phys Rev E 67:026126 (2003) for the proper definition. The actual calculation is performed using equation (26) in the same paper for directed graphs, and equation (4) in Newman MEJ: Assortative mixing in networks, Phys Rev Lett 89:208701 (2002) for undirected graphs.

#### **Parameters**

types1: vertex types in a list or the name of a vertex attribute

holding vertex types. Types are ideally denoted by

numeric values.

types2: in directed assortativity calculations, each vertex can

have an out-type and an in-type. In this case, types1 contains the out-types and this parameter contains the in-types in a list or the name of a vertex attribute. If

None, it is assumed to be equal to types1.

directed: whether to consider edge directions or not.

#### Return Value

the assortativity coefficient

#### Reference:

- Newman MEJ: Mixing patterns in networks, Phys Rev E 67:026126, 2003.
- Newman MEJ: Assortative mixing in networks, Phys Rev Lett 89:208701,

1.

See Also: assortativity\_degree() when the types are the vertex degrees

# assortativity degree(directed=True)

Returns the assortativity of a graph based on vertex degrees.

See assortativity() for the details. assortativity\_degree() simply calls assortativity() with the vertex degrees as types.

#### Parameters

directed: whether to consider edge directions for directed graphs or not. This argument is ignored for undirected graphs.

#### Return Value

the assortativity coefficient

See Also: assortativity()

# **assortativity nominal**(*types*, *directed*=True)

Returns the assortativity of the graph based on vertex categories.

Assuming that the vertices belong to different categories, this function calculates the assortativity coefficient, which specifies the extent to which the connections stay within categories. The assortativity coefficient is one if all the connections stay within categories and minus one if all the connections join vertices of different categories. For a randomly connected network, it is asymptotically zero.

See equation (2) in Newman MEJ: Mixing patterns in networks, Phys Rev E 67:026126 (2003) for the proper definition.

# Parameters

types: vertex types in a list or the name of a vertex attribute

holding vertex types. Types should be denoted by

numeric values.

directed: whether to consider edge directions or not.

### Return Value

the assortativity coefficient

**Reference:** Newman MEJ: Mixing patterns in networks, Phys Rev E 67:026126, 2003.

### attributes()

#### Return Value

the attribute name list of the graph

 $\begin{array}{lll} \textbf{authority\_score}(weights=\texttt{None},\ scale=\texttt{True},\ arpack\_options=\texttt{None},\\ return\_eigenvalue=\texttt{False}) \end{array}$ 

Calculates Kleinberg's authority score for the vertices of the graph

#### **Parameters**

weights: edge weights to be used. Can be a sequence or

iterable or even an edge attribute name.

scale: whether to normalize the scores so that the

largest one is 1.

arpack\_options:
an ARPACKOptions object used to fine-tune

the ARPACK eigenvector calculation. If omitted, the module-level variable called

arpack\_options is used.

return\_eigenvalue: whether to return the largest eigenvalue

## Return Value

the authority scores in a list and optionally the largest eigenvalue as a second member of a tuple

See Also: hub score()

# average path length(directed=True, unconn=True)

Calculates the average path length in a graph.

#### **Parameters**

directed: whether to consider directed paths in case of a directed

graph. Ignored for undirected graphs.

unconn: what to do when the graph is unconnected. If True, the

average of the geodesic lengths in the components is calculated. Otherwise for all unconnected vertex pairs, a path length equal to the number of vertices is used.

#### Return Value

the average path length in the graph

 $\betweenness(vertices=\verb|None|, directed=\verb|True|, cutoff=\verb|None|, weights=\verb|None|, nobigint=\verb|True|)$ 

Calculates or estimates the betweenness of vertices in a graph.

Keyword arguments:

### **Parameters**

vertices: the vertices for which the betweennesses must be

returned. If None, assumes all of the vertices in the

graph.

directed: whether to consider directed paths.

cutoff: if it is an integer, only paths less than or equal to this

length are considered, effectively resulting in an

estimation of the betweenness for the given vertices. If

None, the exact betweenness is returned.

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name.

nobigint: if True, igraph uses the longest available integer type on

the current platform to count shortest paths. For some large networks that have a specific structure, the counters may overflow. To prevent this, use nobigint=False, which forces igraph to use arbitrary precision integers at

the expense of increased computation time.

#### Return Value

the (possibly estimated) betweenness of the given vertices in a list

#### $\mathbf{bfs}(vid, mode = \mathsf{OUT})$

Conducts a breadth first search (BFS) on the graph.

## **Parameters**

vid: the root vertex ID

mode: either IN or OUT or ALL, ignored for undirected graphs.

### Return Value

a tuple with the following items:

- The vertex IDs visited (in order)
- The start indices of the layers in the vertex list
- The parent of every vertex in the BFS

## **bfsiter**(vid, mode=OUT, advanced=False)

Constructs a breadth first search (BFS) iterator of the graph.

#### **Parameters**

vid: the root vertex ID

mode: either IN or OUT or ALL.

advanced: if False, the iterator returns the next vertex in BFS

order in every step. If True, the iterator returns the distance of the vertex from the root and the parent of

the vertex in the BFS tree as well.

### Return Value

the BFS iterator as an igraph.BFSIter object.

# bibcoupling(vertices=None)

Calculates bibliographic coupling scores for given vertices in a graph.

#### **Parameters**

vertices: the vertices to be analysed. If None, all vertices will be

considered.

### Return Value

bibliographic coupling scores for all given vertices in a matrix.

# biconnected components(return articulation points=True)

Calculates the biconnected components of the graph.

Components containing a single vertex only are not considered as being biconnected.

## **Parameters**

return\_articulation\_points: whether to return the articulation points as well

## Return Value

a list of lists containing edge indices making up spanning trees of the biconnected components (one spanning tree for each component) and optionally the list of articulation points

# **bipartite** projection(types, multiplicity=True, probe1=-1, which=-1)

Internal function, undocumented.

See Also: Graph.bipartite projection()

# bipartite projection size(types)

Internal function, undocumented.

See Also: Graph.bipartite projection size()

# bridges()

Returns the list of bridges in the graph.

An edge is a bridge if its removal increases the number of (weakly) connected components in the graph.

# canonical permutation(sh="fm", color=None)

Calculates the canonical permutation of a graph using the BLISS isomorphism algorithm.

Passing the permutation returned here to Graph.permute\_vertices() will transform the graph into its canonical form.

See http://www.tcs.hut.fi/Software/bliss/index.html for more information about the BLISS algorithm and canonical permutations.

#### **Parameters**

sh: splitting heuristics for graph as a case-insensitive string, with the following possible values:

- "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": largest maximally non-trivially connected non-singleton cell
- "fsm": smallest maximally non-trivially connected non-singleton cell

color: optional vector storing a coloring of the vertices with respect to which the isomorphism is computed. If None, all vertices have the same color.

## Return Value

a permutation vector containing vertex IDs. Vertex 0 in the original graph will be mapped to an ID contained in the first element of this vector; vertex 1 will be mapped to the second and so on.

# clique number()

Returns the clique number of the graph.

The clique number of the graph is the size of the largest clique.

See Also: largest\_cliques() for the largest cliques.

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

Returns some or all cliques of the graph as a list of tuples.

A clique is a complete subgraph - a set of vertices where an edge is present between any two of them (excluding loops)

## **Parameters**

min: the minimum size of cliques to be returned. If zero or negative, no lower bound will be used.

max: the maximum size of cliques to be returned. If zero or negative, no upper bound will be used.

 $\begin{calculation} {\bf closeness}(vertices = {\tt None},\ mode = {\tt ALL},\ cutoff = {\tt None},\ weights = {\tt None},\ normalized = {\tt True}) \end{calculation}$ 

Calculates the closeness centralities of given vertices in a graph.

The closeness centerality of a vertex measures how easily other vertices can be reached from it (or the other way: how easily it can be reached from the other vertices). It is defined as the number of the number of vertices minus one divided by the sum of the lengths of all geodesics from/to the given vertex.

If the graph is not connected, and there is no path between two vertices, the number of vertices is used instead the length of the geodesic. This is always longer than the longest possible geodesic.

## Parameters

vertices: the vertices for which the closenesses must be

returned. If None, uses all of the vertices in the graph.

mode: must be one of IN, OUT and ALL. IN means that the

length of the incoming paths, OUT means that the length of the outgoing paths must be calculated. ALL

means that both of them must be calculated.

cutoff: if it is an integer, only paths less than or equal to this

length are considered, effectively resulting in an estimation of the closeness for the given vertices (which is always an underestimation of the real closeness, since some vertex pairs will appear as disconnected even though they are connected).. If

None, the exact closeness is returned.

weights: edge weights to be used. Can be a sequence or iterable

or even an edge attribute name.

normalized: Whether to normalize the raw closeness scores by

multiplying by the number of vertices minus one.

### Return Value

the calculated closenesses in a list

## clusters(mode = STRONG)

Calculates the (strong or weak) clusters for a given graph.

#### **Parameters**

mode: must be either STRONG or WEAK, depending on the clusters being sought. Optional, defaults to STRONG.

### Return Value

the component index for every node in the graph.

**Attention:** this function has a more convenient interface in class **Graph** which wraps the result in a **VertexClustering** object. It is advised to use that.

# cocitation(vertices=None)

Calculates cocitation scores for given vertices in a graph.

#### **Parameters**

vertices: the vertices to be analysed. If None, all vertices will be considered.

### Return Value

cocitation scores for all given vertices in a matrix.

# cohesive blocks()

Calculates the cohesive block structure of the graph.

Attention: this function has a more convenient interface in class Graph which wraps the result in a CohesiveBlocks object. It is advised to use that.

## community edge betweenness(directed=True, weights=None)

Community structure detection based on the betweenness of the edges in the network. This algorithm was invented by M Girvan and MEJ Newman, see: M Girvan and MEJ Newman: Community structure in social and biological networks, Proc. Nat. Acad. Sci. USA 99, 7821-7826 (2002).

The idea is that the betweenness of the edges connecting two communities is typically high. So we gradually remove the edge with the highest betweenness from the network and recalculate edge betweenness after every removal, as long as all edges are removed.

### **Parameters**

directed: whether to take into account the directedness of the

edges when we calculate the betweenness values.

weights: name of an edge attribute or a list containing edge

weights.

#### Return Value

a tuple with the merge matrix that describes the dendrogram and the modularity scores before each merge. The modularity scores use the weights if the original graph was weighted.

Attention: this function is wrapped in a more convenient syntax in the derived class Graph. It is advised to use that instead of this version.

## **community** fastgreedy(weights=None)

Finds the community structure of the graph according to the algorithm of Clauset et al based on the greedy optimization of modularity.

This is a bottom-up algorithm: initially every vertex belongs to a separate community, and communities are merged one by one. In every step, the two communities being merged are the ones which result in the maximal increase in modularity.

#### **Parameters**

weights: name of an edge attribute or a list containing edge weights

### Return Value

- a tuple with the following elements:
  - 1. The list of merges
  - 2. The modularity scores before each merge

Attention: this function is wrapped in a more convenient syntax in the derived class Graph. It is advised to use that instead of this version.

**Reference:** A. Clauset, M. E. J. Newman and C. Moore: *Finding community structure in very large networks*. Phys Rev E 70, 066111 (2004).

See Also: modularity()

 $\begin{array}{l} \textbf{community\_infomap}(\textit{edge\_weights} = \texttt{None}, \textit{vertex\_weights} = \texttt{None}, \\ \textit{trials} = \texttt{10}) \end{array}$ 

Finds the community structure of the network according to the Infomap method of Martin Rosvall and Carl T. Bergstrom.

See http://www.mapequation.org for a visualization of the algorithm or one of the references provided below.

# **Parameters**

edge\_weights: name of an edge attribute or a list containing

edge weights.

vertex\_weights: name of an vertex attribute or a list containing

vertex weights.

trials: the number of attempts to partition the network.

### Return Value

the calculated membership vector and the corresponding codelength in a tuple.

### Reference:

- M. Rosvall and C. T. Bergstrom: Maps of information flow reveal community structure in complex networks. PNAS 105, 1118 (2008). http://arxiv.org/abs/0707.0609
- M. Rosvall, D. Axelsson and C. T. Bergstrom: *The map equation*. Eur Phys J Special Topics 178, 13 (2009). http://arxiv.org/abs/0906.1405

# **community** label propagation(weights=None, initial=None, fixed=None)

Finds the community structure of the graph according to the label propagation method of Raghavan et al.

Initially, each vertex is assigned a different label. After that, each vertex chooses the dominant label in its neighbourhood in each iteration. Ties are broken randomly and the order in which the vertices are updated is randomized before every iteration. The algorithm ends when vertices reach a consensus.

Note that since ties are broken randomly, there is no guarantee that the algorithm returns the same community structure after each run. In fact, they frequently differ. See the paper of Raghavan et al on how to come up with an aggregated community structure.

## **Parameters**

weights: name of an edge attribute or a list containing edge weights

initial: name of a vertex attribute or a list containing the initial

vertex labels. Labels are identified by integers from zero to n-1 where n is the number of vertices. Negative

numbers may also be present in this vector, they represent

unlabeled vertices.

fixed: a list of booleans for each vertex. True corresponds to

vertices whose labeling should not change during the algorithm. It only makes sense if initial labels are also given. Unlabeled vertices cannot be fixed. Note that

vertex attribute names are not accepted here.

### Return Value

the resulting membership vector

**Reference:** 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. http://arxiv.org/abs/0709.2938.

 $\begin{array}{l} \mathbf{community\_leading\_eigenvector}(n{=}\text{-1},\ arpack\_options}{=}\mathbb{N} \texttt{one},\\ weights}{=}\mathbb{N} \texttt{one}) \end{array}$ 

A proper implementation of Newman's eigenvector community structure detection. Each split is done by maximizing the modularity regarding the original network. See the reference for details.

### **Parameters**

n: the desired number of communities. If negative,

the algorithm tries to do as many splits as possible. Note that the algorithm won't split a community further if the signs of the leading

eigenvector are all the same.

arpack\_options: an ARPACKOptions object used to fine-tune the

ARPACK eigenvector calculation. If omitted, the module-level variable called arpack\_options is

used.

weights: name of an edge attribute or a list containing

edge weights

#### Return Value

a tuple where the first element is the membership vector of the clustering and the second element is the merge matrix.

**Attention:** this function is wrapped in a more convenient syntax in the derived class **Graph**. It is advised to use that instead of this version.

Reference: MEJ Newman: Finding community structure in networks using

the eigenvectors of matrices, arXiv:physics/0605087

community\_leiden(edge\_weights=None, node\_weights=None,
resolution\_parameter=1.0, normalize\_resolution=False, beta=0.01,
initial\_membership=None, n\_iterations=2)

Finds the community structure of the graph using the Leiden algorithm of Traag, van Eck & Waltman

#### **Parameters**

edge\_weights: edge weights to be used. Can be a

sequence or iterable or even an edge

attribute name.

node\_weights: the node weights used in the Leiden

algorithm.

resolution\_parameter: the resolution parameter to use. Higher

resolutions lead to more smaller

communities, while lower resolutions lead

to fewer larger communities.

normalize\_resolution: if set to true, the resolution parameter will

be divided by the sum of the node weights. If this is not supplied, it will default to the node degree, or weighted degree in case

edge weights are supplied.

node\_weights: the node weights used in the Leiden

algorithm.

beta: parameter affecting the randomness in the

Leiden algorithm. This affects only the

refinement step of the algorithm.

initial\_membership: if provided, the Leiden algorithm will try

to improve this provided membership. If no argument is provided, the aglorithm simply starts from the singleton partition.

n\_iterations: the number of iterations to iterate the

Leiden algorithm. Each iteration may

improve the partition further.

### Return Value

the community membership vector.

## **community** multilevel(weights=None, return levels=True)

Finds the community structure of the graph according to the multilevel algorithm of Blondel et al. This is a bottom-up algorithm: initially every vertex belongs to a separate community, and vertices are moved between communities iteratively in a way that maximizes the vertices' local contribution to the overall modularity score. When a consensus is reached (i.e. no single move would increase the modularity score), every community in the original graph is shrank to a single vertex (while keeping the total weight of the incident edges) and the process continues on the next level. The algorithm stops when it is not possible to increase the modularity any more after shrinking the communities to vertices.

#### **Parameters**

weights: name of an edge attribute or a list containing edge

weights

return\_levels: if True, returns the multilevel result. If False,

only the best level (corresponding to the best

modularity) is returned.

#### Return Value

either a single list describing the community membership of each vertex (if return\_levels is False), or a list of community membership vectors, one corresponding to each level and a list of corresponding modularities (if return\_levels is True).

**Attention:** this function is wrapped in a more convenient syntax in the derived class **Graph**. It is advised to use that instead of this version.

**Reference:** VD Blondel, J-L Guillaume, R Lambiotte and E Lefebvre: Fast unfolding of community hierarchies in large networks. J Stat Mech P10008 (2008), http://arxiv.org/abs/0803.0476

See Also: modularity()

# community optimal modularity(weights=None)

Calculates the optimal modularity score of the graph and the corresponding community structure.

This function uses the GNU Linear Programming Kit to solve a large integer optimization problem in order to find the optimal modularity score and the corresponding community structure, therefore it is unlikely to work for graphs larger than a few (less than a hundred) vertices. Consider using one of the heuristic approaches instead if you have such a large graph.

# **Parameters**

weights: name of an edge attribute or a list containing edge weights.

### Return Value

the calculated membership vector and the corresponding modularity in a tuple.

 $\begin{array}{l} \textbf{community\_spinglass}(weights=\texttt{None},\ spins=\texttt{25},\ parupdate=\texttt{False},\\ start\_temp=\texttt{1},\ stop\_temp=\texttt{0.01},\ cool\_fact=\texttt{0.99},\ update\_rule=\texttt{"config"},\\ gamma=\texttt{1},\ implementation=\texttt{"orig"},\ lambda=\texttt{1}) \end{array}$ 

Finds the community structure of the graph according to the spinglass community detection method of Reichardt & Bornholdt.

### Parameters

weights: edge weights to be used. Can be a sequence or

iterable or even an edge attribute name.

spins: integer, 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: whether to update the spins of the vertices in

parallel (synchronously) or not

start\_temp: the starting temperature

stop\_temp: the stop temperature

cool\_fact: cooling factor for the simulated annealing

update\_rule: specifies the null model of the simulation.

Possible values are "config" (a random graph with the same vertex degrees as the input graph) or "simple" (a random graph with the same

number of edges)

gamma: the gamma argument of the algorithm, specifying

the balance between the importance of present and missing edges within a community. The default value of 1.0 assigns equal importance to

both of them.

implementation: currently igraph contains two implementations for

the spinglass community detection algorithm.

The faster original implementation is the default.

The other implementation is able to take into account negative weights, this can be chosen by

setting implementation to "neg".

lambda: the lambda argument of the algorithm, which

specifies the balance between the importance of present and missing negatively weighted edges within a community. Smaller values of lambda

lead to communities with less negative

intra-connectivity. If the argument is zero, the algorithm reduces to a graph coloring algorithm,

using the number of spins as colors. This

argument is ignored if the original

implementation is used.

#### Return Value

the community membership vector.

# **community** walktrap(weights=None, steps=None)

Finds the community structure of the graph according to the random walk method of Latapy & Pons.

The basic idea of the algorithm is that short random walks tend to stay in the same community. The method provides a dendrogram.

#### **Parameters**

weights: name of an edge attribute or a list containing edge weights

#### Return Value

a tuple with the list of merges and the modularity scores corresponding to each merge

Attention: this function is wrapped in a more convenient syntax in the derived class Graph. It is advised to use that instead of this version.

**Reference:** Pascal Pons, Matthieu Latapy: Computing communities in large networks using random walks, <a href="http://arxiv.org/abs/physics/0512106">http://arxiv.org/abs/physics/0512106</a>.

See Also: modularity()

# complementer(loops = False)

Returns the complementer of the graph

#### **Parameters**

loops: whether to include loop edges in the complementer.

### Return Value

the complementer of the graph

# **compose**(other)

Returns the composition of two graphs.

# constraint(vertices = None, weights = None)

Calculates Burt's constraint scores for given vertices in a graph.

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 as follows:

$$C[i] = sum( sum( (p[i,q] p[q,j])^2, q in V[i], q != i,j ), j in V[], j != i)$$

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

p[i,j]=(a[i,j]+a[j,i]) / sum(a[i,k]+a[k,i], k in V[i], k != i), a[i,j] are elements of A and the latter being the graph adjacency matrix.

For isolated vertices, constraint is undefined.

### **Parameters**

vertices: the vertices to be analysed or None for all vertices.

weights: weights associated to the edges. Can be an attribute

name as well. If None, every edge will have the same

weight.

#### Return Value

constraint scores for all given vertices in a matrix.

# **contract** vertices(mapping, combine attrs=None)

Contracts some vertices in the graph, i.e. replaces groups of vertices with single vertices. Edges are not affected.

#### **Parameters**

mapping:

numeric vector which gives the mapping between old and new vertex IDs. Vertices having the same new vertex ID in this vector will be remapped into a single new vertex. It is safe to pass the membership vector of a VertexClustering object here.

combine\_attrs: specifies how to combine the attributes of the vertices being collapsed into a single one. If it is None, all the attributes will be lost. If it is a function, the attributes of the vertices will be collected and passed on to that function which will return the new attribute value that has to be assigned to the single collapsed vertex. It can also be one of the following string constants which define built-in collapsing functions: sum, prod, mean, median, max, min, first, last, random. You can also specify different combination functions for different attributes by passing a dict here which maps attribute names to functions. See Graph.simplify() for more details.

### Return Value

None.

See Also: Graph.simplify()

# convergence degree()

Undocumented (yet).

# convergence field size()

Undocumented (yet).

# copy()

Creates a copy of the graph.

Attributes are copied by reference; in other words, if you use mutable Python objects as attribute values, these objects will still be shared between the old and new graph. You can use 'deepcopy()' from the 'copy' module if you need a truly deep copy of the graph.

# coreness(mode=ALL)

Finds the coreness (shell index) of the vertices of the network.

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

#### **Parameters**

mode: whether to compute the in-corenesses (IN), the out-corenesses (OUT) or the undirected corenesses (ALL). Ignored and assumed to be ALL for undirected graphs.

### Return Value

the corenesses for each vertex.

**Reference:** Vladimir Batagelj, Matjaz Zaversnik: An O(m) Algorithm for Core Decomposition of Networks.

 $\begin{array}{l} \mathbf{count\_isomorphisms\_vf2}(other = \mathtt{None},\ color1 = \mathtt{None},\ color2 = \mathtt{None},\ edge\_color1 = \mathtt{None},\ edge\_color2 = \mathtt{None},\ node\_compat\_fn = \mathtt{None},\ edge\_compat\_fn = \mathtt{None}) \end{array}$ 

Determines the number of isomorphisms between the graph and another one

Vertex and edge colors may be used to restrict the isomorphisms, as only vertices and edges with the same color will be allowed to match each other.

#### **Parameters**

other: the other graph. If None, the number of

automorphisms will be returned.

color1: optional vector storing the coloring of the vertices

of the first graph. If None, all vertices have the

same color.

color2: optional vector storing the coloring of the vertices

of the second graph. If None, all vertices have the

same color.

edge\_color1: optional vector storing the coloring of the edges of

the first graph. If None, all edges have the same

color.

edge\_color2: optional vector storing the coloring of the edges of

the second graph. If None, all edges have the

same color.

node\_compat\_fn: a function that receives the two graphs and two

node indices (one from the first graph, one from the second graph) and returns True if the nodes given by the two indices are compatible (i.e. they could be matched to each other) or False

otherwise. This can be used to restrict the set of isomorphisms based on node-specific criteria that are too complicated to be represented by node color vectors (i.e. the color1 and color2 parameters). None means that every node is

compatible with every other node.

edge\_compat\_fn: a function that receives the two graphs and two

edge indices (one from the first graph, one from the second graph) and returns True if the edges given by the two indices are compatible (i.e. they could be matched to each other) or False otherwise. This can be used to restrict the set of isomorphisms based on edge-specific criteria that

are too complicated to be represented by edge color vectors (i.e. the edge\_color1 and

edge\_color2 parameters). None means that every edge is compatible with every other node.

# Return Value

the number of isomorphisms between the two given graphs (or the number of automorphisms if other is None.

# $\mathbf{count} \quad \mathbf{multiple}(\mathit{edges} {=} \mathtt{None})$

Counts the multiplicities of the given edges.

# **Parameters**

edges: edge indices for which we want to count their multiplicity. If None, all edges are counted.

# Return Value

the multiplicities of the given edges as a list.

 $\begin{array}{l} \textbf{count\_subisomorphisms\_vf2}(\textit{other}, \textit{color1} = \texttt{None}, \textit{color2} = \texttt{None}, \\ \textit{edge\_color1} = \texttt{None}, \textit{edge\_color2} = \texttt{None}, \textit{node\_compat\_fn} = \texttt{None}, \\ \textit{edge\_compat\_fn} = \texttt{None}) \end{array}$ 

Determines the number of subisomorphisms between the graph and another one

Vertex and edge colors may be used to restrict the isomorphisms, as only vertices and edges with the same color will be allowed to match each other.

#### Parameters

other: the other graph.

color1: optional vector storing the coloring of the vertices

of the first graph. If None, all vertices have the

same color.

color2: optional vector storing the coloring of the vertices

of the second graph. If None, all vertices have the

same color.

edge\_color1: optional vector storing the coloring of the edges of

the first graph. If None, all edges have the same

color.

edge\_color2: optional vector storing the coloring of the edges of

the second graph. If None, all edges have the

same color.

node\_compat\_fn: a function that receives the two graphs and two

node indices (one from the first graph, one from the second graph) and returns True if the nodes given by the two indices are compatible (i.e. they could be matched to each other) or False

otherwise. This can be used to restrict the set of isomorphisms based on node-specific criteria that are too complicated to be represented by node color vectors (i.e. the color1 and color2 parameters). None means that every node is

compatible with every other node.

edge\_compat\_fn: a function that receives the two graphs and two

edge indices (one from the first graph, one from the second graph) and returns True if the edges given by the two indices are compatible (i.e. they

could be matched to each other) or False

otherwise. This can be used to restrict the set of isomorphisms based on edge-specific criteria that are too complicated to be represented by edge

color vectors (i.e. the edge\_color1 and edge\_color2 parameters). None means that every edge is compatible with every other node.

## Return Value

the number of subisomorphisms between the two given graphs

## decompose(mode=STRONG, maxcompno=None, minelements=1)

Decomposes the graph into subgraphs.

### **Parameters**

mode: must be either STRONG or WEAK, depending on

the clusters being sought.

maxcompno: maximum number of components to return. None

means all possible components.

minelements: minimum number of vertices in a component. By

setting this to 2, isolated vertices are not returned as

separate components.

#### Return Value

a list of the subgraphs. Every returned subgraph is a copy of the original.

## degree(vertices, mode=ALL, loops=True)

Returns some vertex degrees from the graph.

This method accepts a single vertex ID or a list of vertex IDs as a parameter, and returns the degree of the given vertices (in the form of a single integer or a list, depending on the input parameter).

### **Parameters**

vertices: a single vertex ID or a list of vertex IDs

mode: the type of degree to be returned (OUT for out-degrees,

IN IN for in-degrees or ALL for the sum of them).

loops: whether self-loops should be counted.

# delete edges(es)

Removes edges from the graph.

All vertices will be kept, even if they lose all their edges. Nonexistent edges will be silently ignored.

#### **Parameters**

es: the list of edges to be removed. Edges are identifed by edge IDs. EdgeSeq objects are also accepted here. No argument deletes all edges.

## delete vertices(vs)

Deletes vertices and all its edges from the graph.

#### **Parameters**

vs: a single vertex ID or the list of vertex IDs to be deleted. No argument deletes all vertices.

# density(loops = False)

Calculates the density of the graph.

#### **Parameters**

loops: whether to take loops into consideration. If True, the algorithm assumes that there might be some loops in the graph and calculates the density accordingly. If False, the algorithm assumes that there can't be any loops.

#### Return Value

the density of the graph.

### dfsiter(vid, mode=OUT, advanced=False)

Constructs a depth first search (DFS) iterator of the graph.

#### Parameters

vid: the root vertex ID

mode: either IN or OUT or ALL.

advanced: if False, the iterator returns the next vertex in DFS

order in every step. If True, the iterator returns the distance of the vertex from the root and the parent of

the vertex in the DFS tree as well.

## Return Value

the DFS iterator as an igraph.DFSIter object.

## diameter(directed=True, unconn=True, weights=None)

Calculates the diameter of the graph.

#### **Parameters**

directed: whether to consider directed paths.

unconn: if True and the graph is unconnected, the longest

geodesic within a component will be returned. If False and the graph is unconnected, the result is the number of vertices if there are no weights or infinity if there are

weights.

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name.

#### Return Value

the diameter

# **difference**(other)

Subtracts the given graph from the original

# diversity(vertices=None, weights=None)

Calculates the structural diversity index of the vertices.

The structural diversity index of a vertex is simply the (normalized) Shannon entropy of the weights of the edges incident on the vertex.

The measure is defined for undirected graphs only; edge directions are ignored.

## Parameters

vertices: the vertices for which the diversity indices must be

returned. If None, uses all of the vertices in the graph.

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name.

#### Return Value

the calculated diversity indices in a list, or a single number if a single vertex was supplied.

Reference: Eagle N, Macy M and Claxton R: Network diversity and economic development, Science 328, 1029–1031, 2010.

## dominator(...)

dominator(vid, mode=)

Returns the dominator tree from the given root node@param vid: the root vertex ID

### Parameters

mode: either IN or OUT

### Return Value

a list containing the dominator tree for the current graph.

# dyad census()

Dyad census, as defined by Holland and Leinhardt

Dyad census means classifying each pair of vertices of a directed graph into three categories: mutual, there is an edge from a to b and also from b to a; asymmetric, there is an edge either from a to b or from b to a but not the other way and null, no edges between a and b.

## Return Value

the number of mutual, asymmetric and null connections in a 3-tuple.

Attention: this function has a more convenient interface in class Graph which wraps the result in a DyadCensus object. It is advised to use that.

### eccentricity(vertices=None, mode=ALL)

Calculates the eccentricities of given vertices in a graph.

The eccentricity of a vertex is calculated by measuring the shortest distance from (or to) the vertex, to (or from) all other vertices in the graph, and taking the maximum.

# **Parameters**

vertices: the vertices for which the eccentricity scores must be

returned. If None, uses all of the vertices in the graph.

mode: must be one of IN, OUT and ALL. IN means that edge

directions are followed; OUT means that edge directions are followed the opposite direction; ALL means that directions are ignored. The argument has no effect for

undirected graphs.

### Return Value

the calculated eccentricities in a list, or a single number if a single vertex was supplied.

## ecount()

Counts the number of edges.

# Return Value

the number of edges in the graph.

(type=integer)

# edge attributes()

# Return Value

the attribute name list of the graph's edges

# edge betweenness(directed=True, cutoff=None, weights=None)

Calculates or estimates the edge betweennesses in a graph.

#### **Parameters**

directed: whether to consider directed paths.

cutoff: if it is an integer, only paths less than or equal to this

length are considered, effectively resulting in an

estimation of the betweenness values. If None, the exact

betweennesses are returned.

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name.

## Return Value

a list with the (exact or estimated) edge betweennesses of all edges.

## edge connectivity(source=-1, target=-1, checks=True)

Calculates the edge connectivity of the graph or between some vertices.

The edge connectivity between two given vertices is the number of edges that have to be removed in order to disconnect the two vertices into two separate components. This is also the number of edge disjoint directed paths between the vertices. The edge connectivity of the graph is the minimal edge connectivity over all vertex pairs.

This method calculates the edge connectivity of a given vertex pair if both the source and target vertices are given. If none of them is given (or they are both negative), the overall edge connectivity is returned.

#### Parameters

source: the source vertex involved in the calculation.

target: the target vertex involved in the calculation.

checks: if the whole graph connectivity is calculated and this is

True, igraph performs some basic checks before calculation.

If the graph is not strongly connected, then the

connectivity is obviously zero. If the minimum degree is one, then the connectivity is also one. These simple checks are much faster than checking the entire graph, therefore it is advised to set this to True. The parameter is ignored if the connectivity between two given vertices is computed.

### Return Value

the edge connectivity

### eigen adjacency(...)

eigenvector\_centrality(directed=True, scale=True, weights=None,
return eigenvalue=False, arpack options=None)

Calculates the eigenvector centralities of the vertices in a graph.

Eigenvector centrality is a measure of the importance of a node in a network. It assigns relative scores to all nodes in the network based on the principle that connections from high-scoring nodes contribute more to the score of the node in question than equal connections from low-scoring nodes. In practice, the centralities are determined by calculating eigenvector corresponding to the largest positive eigenvalue of the adjacency matrix. In the undirected case, this function considers the diagonal entries of the adjacency matrix to be twice the number of self-loops on the corresponding vertex.

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.

#### **Parameters**

directed: whether to consider edge directions in a

directed graph. Ignored for undirected graphs.

scale: whether to normalize the centralities so the

largest one will always be 1.

weights: edge weights given as a list or an edge

attribute. If None, all edges have equal weight.

return\_eigenvalue: whether to return the actual largest

eigenvalue along with the centralities

arpack\_options: an ARPACKOptions object that can be used to

fine-tune the calculation. If it is omitted, the module-level variable called arpack\_options

is used.

### Return Value

the eigenvector centralities in a list and optionally the largest eigenvalue (as a second member of a tuple)

# farthest points(directed=True, unconn=True, weights=None)

Returns two vertex IDs whose distance equals the actual diameter of the graph.

If there are many shortest paths with the length of the diameter, it returns the first one it found.

#### **Parameters**

directed: whether to consider directed paths.

unconn: if True and the graph is unconnected, the longest

geodesic within a component will be returned. If False and the graph is unconnected, the result contains the number of vertices if there are no weights or infinity if

there are weights.

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name.

### Return Value

a triplet containing the two vertex IDs and their distance. The IDs are None if the graph is unconnected and unconn is False.

## feedback arc set(weights=None, method="eades")

Calculates an approximately or exactly minimal feedback arc set.

A feedback arc set is a set of edges whose removal makes the graph acyclic. Since this is always possible by removing all the edges, we are in general interested in removing the smallest possible number of edges, or an edge set with as small total weight as possible. This method calculates one such edge set. Note that the task is trivial for an undirected graph as it is enough to find a spanning tree and then remove all the edges not in the spanning tree. Of course it is more complicated for directed graphs.

#### **Parameters**

weights: edge weights to be used. Can be a sequence or iterable or

minimize the total weight of the feedback arc set.

even an edge attribute name. When given, the algorithm will strive to remove lightweight edges in order to

method.

the algorithm to use. "eades" uses the greedy cycle breaking heuristic of Eades, Lin and Smyth, which is linear in the number of edges but not necessarily optimal; however, it guarantees that the number of edges to be removed is smaller than  $|\mathbf{E}|/2 - |\mathbf{V}|/6$ . "ip" uses an integer programming formulation which is guaranteed to yield an optimal result, but is too slow for large graphs.

### Return Value

the IDs of the edges to be removed, in a list.

**Reference:** Eades P, Lin X and Smyth WF: A fast and effective heuristic for the feedback arc set problem. In: Proc Inf Process Lett 319-323, 1993.

# $get adjacency(type=GET\_ADJACENCY\_BOTH, eids=False)$

Returns the adjacency matrix of a graph.

### **Parameters**

type: either GET\_ADJACENCY\_LOWER (uses the lower triangle of the matrix) or GET\_ADJACENCY\_UPPER (uses the upper triangle) or GET\_ADJACENCY\_BOTH (uses both parts). Ignored for directed graphs.

eids: if True, the result matrix will contain zeros for non-edges and the ID of the edge plus one for edges in the appropriate cell.If False, the result matrix will contain the number of edges for each vertex pair.

#### Return Value

the adjacency matrix.

# get all shortest paths(v, to=None, weights=None, mode=OUT)

Calculates all of the shortest paths from/to a given node in a graph.

#### **Parameters**

v: the source for the calculated paths

to: a vertex selector describing the destination for the

calculated paths. This can be a single vertex ID, a list of vertex IDs, a single vertex name, a list of vertex names or

a VertexSeq object. None means all the vertices.

weights: edge weights in a list or the name of an edge attribute

holding edge weights. If None, all edges are assumed to

have equal weight.

mode: the directionality of the paths. IN means to calculate

incoming paths, OUT means to calculate outgoing paths,

ALL means to calculate both ones.

#### Return Value

all of the shortest path from the given node to every other reachable node in the graph in a list. Note that in case of mode=IN, the vertices in a path are returned in reversed order!

# get diameter(directed=True, unconn=True, weights=None)

Returns a path with the actual diameter of the graph.

If there are many shortest paths with the length of the diameter, it returns the first one it founds.

# **Parameters**

directed: whether to consider directed paths.

unconn: if True and the graph is unconnected, the longest

geodesic within a component will be returned. If False and the graph is unconnected, the result is the number of vertices if there are no weights or infinity if there are

weights.

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name.

#### Return Value

the vertices in the path in order.

# get edgelist()

Returns the edge list of a graph.

# get eid(v1, v2, directed=True, error=True)

Returns the edge ID of an arbitrary edge between vertices v1 and v2

#### **Parameters**

v1: the ID or name of the first vertex

v2: the ID or name of the second vertex

directed: whether edge directions should be considered in directed

graphs. The default is True. Ignored for undirected

graphs.

error: if True, an exception will be raised when the given edge

does not exist. If False, -1 will be returned in that case.

### Return Value

the edge ID of an arbitrary edge between vertices v1 and v2

# get eids(pairs=None, path=None, directed=True, error=True)

Returns the edge IDs of some edges between some vertices.

This method can operate in two different modes, depending on which of the keyword arguments pairs and path are given.

The method does not consider multiple edges; if there are multiple edges between a pair of vertices, only the ID of one of the edges is returned.

## **Parameters**

pairs: a list of integer pairs. Each integer pair is considered as a

source-target vertex pair; the corresponding edge is looked up in the graph and the edge ID is returned for

each pair.

path: a list of vertex IDs. The list is considered as a continuous

path from the first vertex to the last, passing through the intermediate vertices. The corresponding edge IDs between the first and the second, the second and the third and so on are looked up in the graph and the edge IDs are returned. If both path and pairs are given, the

two lists are concatenated.

directed: whether edge directions should be considered in directed

graphs. The default is True. Ignored for undirected

graphs.

error: if True, an exception will be raised if a given edge does

not exist. If False, -1 will be returned in that case.

### Return Value

the edge IDs in a list

Internal function, undocumented.

 ${\bf See \ Also: \ Graph.get\_incidence()}$ 

Returns all isomorphisms between the graph and another one

Vertex and edge colors may be used to restrict the isomorphisms, as only vertices and edges with the same color will be allowed to match each other.

#### **Parameters**

other: the other graph. If None, the automorphisms will

be returned.

color1: optional vector storing the coloring of the vertices

of the first graph. If None, all vertices have the

same color.

color2: optional vector storing the coloring of the vertices

of the second graph. If None, all vertices have the

same color.

edge\_color1: optional vector storing the coloring of the edges of

the first graph. If None, all edges have the same

color.

edge\_color2: optional vector storing the coloring of the edges of

the second graph. If None, all edges have the

same color.

node\_compat\_fn: a function that receives the two graphs and two

node indices (one from the first graph, one from the second graph) and returns True if the nodes given by the two indices are compatible (i.e. they could be matched to each other) or False otherwise. This can be used to restrict the set of

isomorphisms based on node-specific criteria that are too complicated to be represented by node color vectors (i.e. the color1 and color2 parameters). None means that every node is

compatible with every other node.

edge\_compat\_fn: a function that receives the two graphs and two

edge indices (one from the first graph, one from the second graph) and returns True if the edges given by the two indices are compatible (i.e. they could be matched to each other) or False otherwise. This can be used to restrict the set of isomorphisms based on edge-specific criteria that are too complicated to be represented by edge

color vectors (i.e. the edge\_color1 and edge\_color2 parameters). None means that

every edge is compatible with every other node.

## Return Value

a list of lists, each item of the list containing the mapping from vertices of the second graph to the vertices of the first one

 $get\_shortest\_paths(v, to=None, weights=None, mode=OUT, output="vpath")$ 

Calculates the shortest paths from/to a given node in a graph.

#### Parameters

v: the source/destination for the calculated paths

to: a vertex selector describing the destination/source for the

calculated paths. This can be a single vertex ID, a list of vertex IDs, a single vertex name, a list of vertex names or

a VertexSeq object. None means all the vertices.

weights: edge weights in a list or the name of an edge attribute

holding edge weights. If None, all edges are assumed to

have equal weight.

mode: the directionality of the paths. IN means to calculate

incoming paths, OUT means to calculate outgoing paths,

ALL means to calculate both ones.

output: determines what should be returned. If this is "vpath", a

list of vertex IDs will be returned, one path for each target vertex. For unconnected graphs, some of the list elements may be empty. Note that in case of mode=IN, the vertices in a path are returned in reversed order. If output="epath", edge IDs are returned instead of vertex

IDs.

#### Return Value

see the documentation of the output parameter.

 ${f get\_subisomorphisms\_lad}(other,\ domains={f None},\ induced={f False},\ time\ limit=0)$ 

Returns all subisomorphisms between the graph and another one using the LAD algorithm.

The optional domains argument may be used to restrict vertices that may match each other. You can also specify whether you are interested in induced subgraphs only or not.

#### **Parameters**

other: the pattern graph we are looking for in the graph.

domains: a list of lists, one sublist belonging to each vertex in

the template graph. Sublist i contains the indices of the vertices in the original graph that may match vertex i in the template graph. None means that every

vertex may match every other vertex.

induced: whether to consider induced subgraphs only.

time\_limit: an optimal time limit in seconds. Only the integral

part of this number is taken into account. If the time limit is exceeded, the method will throw an exception.

### Return Value

a list of lists, each item of the list containing the mapping from vertices of the second graph to the vertices of the first one

Returns all subisomorphisms between the graph and another one

Vertex and edge colors may be used to restrict the isomorphisms, as only vertices and edges with the same color will be allowed to match each other.

#### **Parameters**

other: the other graph.

color1: optional vector storing the coloring of the vertices

of the first graph. If None, all vertices have the

same color.

color2: optional vector storing the coloring of the vertices

of the second graph. If None, all vertices have the

same color.

edge\_color1: optional vector storing the coloring of the edges of

the first graph. If None, all edges have the same

color.

edge\_color2: optional vector storing the coloring of the edges of

the second graph. If None, all edges have the

same color.

node\_compat\_fn: a function that receives the two graphs and two

node indices (one from the first graph, one from the second graph) and returns True if the nodes given by the two indices are compatible (i.e. they

could be matched to each other) or False

otherwise. This can be used to restrict the set of isomorphisms based on node-specific criteria that are too complicated to be represented by node color vectors (i.e. the color1 and color2 parameters). None means that every node is

compatible with every other node.

edge\_compat\_fn: a function that receives the two graphs and two

edge indices (one from the first graph, one from the second graph) and returns True if the edges given by the two indices are compatible (i.e. they

could be matched to each other) or False

otherwise. This can be used to restrict the set of isomorphisms based on edge-specific criteria that are too complicated to be represented by edge

color vectors (i.e. the edge\_color1 and edge\_color2 parameters). None means that

every edge is compatible with every other node.

#### Return Value

a list of lists, each item of the list containing the mapping from vertices of the second graph to the vertices of the first one

# girth(return\_shortest\_circle=False)

Returns the girth of the graph.

The girth of a graph is the length of the shortest circle in it.

## **Parameters**

return\_shortest\_circle: whether to return one of the shortest circles found in the graph.

# Return Value

the length of the shortest circle or (if return\_shortest\_circle) is true, the shortest circle itself as a list

# gomory hu tree(capacity=None)

Internal function, undocumented.

See Also: Graph.gomory\_hu\_tree()

# has multiple()

Checks whether the graph has multiple edges.

# Return Value

True if the graph has at least one multiple edge, False otherwise.

(type=boolean)

hub\_score(weights=None, scale=True, arpack\_options=None,
return eigenvalue=False)

Calculates Kleinberg's hub score for the vertices of the graph

#### **Parameters**

weights: edge weights to be used. Can be a sequence or

iterable or even an edge attribute name.

scale: whether to normalize the scores so that the

largest one is 1.

arpack\_options: an ARPACKOptions object used to fine-tune

the ARPACK eigenvector calculation. If omitted, the module-level variable called

arpack\_options is used.

return\_eigenvalue: whether to return the largest eigenvalue

#### Return Value

the hub scores in a list and optionally the largest eigenvalue as a second member of a tuple

See Also: authority score()

# incident(vertex, mode=OUT)

Returns the edges a given vertex is incident on.

#### **Parameters**

vertex: a vertex ID

mode: whether to return only successors (OUT), predecessors (IN)

or both (ALL). Ignored for undirected graphs.

# independence number()

Returns the independence number of the graph.

The independence number of the graph is the size of the largest independent vertex set.

See Also: largest\_independent\_vertex\_sets() for the largest independent vertex sets

## independent vertex sets(min=0, max=0)

Returns some or all independent vertex sets of the graph as a list of tuples.

Two vertices are independent if there is no edge between them. Members of an independent vertex set are mutually independent.

#### **Parameters**

min: the minimum size of sets to be returned. If zero or negative, no lower bound will be used.

max: the maximum size of sets to be returned. If zero or negative, no upper bound will be used.

# induced subgraph(vertices, implementation="auto")

Returns a subgraph spanned by the given vertices.

#### **Parameters**

vertices: a list containing the vertex IDs which should be

included in the result.

implementation: the implementation to use when constructing the

new subgraph. igraph includes two implementations at the moment.

"copy\_and\_delete" copies the original graph and removes those vertices that are not in the given set. This is more efficient if the size of the subgraph is comparable to the original graph. The

other implementation ("create\_from\_scratch") constructs the result graph from scratch and then copies the attributes accordingly. This is a better solution if the subgraph is relatively small, compared to the original graph. "auto" selects between the two implementations automatically, based on the ratio of the size of the subgraph and

the size of the original graph.

# Return Value

the subgraph

# $\mathbf{bipartite}(\mathit{return\_types} = \mathsf{False})$

Decides whether the graph is bipartite or not.

Vertices of a bipartite graph can be partitioned into two groups A and B in a way that all edges go between the two groups.

## **Parameters**

return\_types: if False, the method will simply return True or False depending on whether the graph is bipartite or not. If True, the actual group assignments are also returned as a list of boolean values. (Note that the group assignment is not unique, especially if the graph consists of multiple components, since the assignments of components are independent from each other).

## Return Value

True if the graph is bipartite, False if not. If return\_types is True, the group assignment is also returned.

## is connected(mode = STRONG)

Decides whether the graph is connected.

### **Parameters**

mode: whether we should calculate strong or weak connectivity.

## Return Value

True if the graph is connected, False otherwise.

## is dag()

Checks whether the graph is a DAG (directed acyclic graph).

A DAG is a directed graph with no directed cycles.

### Return Value

True if it is a DAG, False otherwise.

(type=boolean)

### is directed()

Checks whether the graph is directed.

## Return Value

True if it is directed, False otherwise.

(type=boolean)

## is loop(edges=None)

Checks whether a specific set of edges contain loop edges

#### **Parameters**

edges: edge indices which we want to check. If None, all edges are checked.

### Return Value

a list of booleans, one for every edge given

## is minimal separator(vertices)

Decides whether the given vertex set is a minimal separator.

A minimal separator is a set of vertices whose removal disconnects the graph, while the removal of any subset of the set keeps the graph connected.

#### **Parameters**

vertices: a single vertex ID or a list of vertex IDs

### Return Value

True is the given vertex set is a minimal separator, False otherwise.

## is multiple(edges=None)

Checks whether an edge is a multiple edge.

Also works for a set of edges – in this case, every edge is checked one by one. Note that if there are multiple edges going between a pair of vertices, there is always one of them that is *not* reported as multiple (only the others). This allows one to easily detect the edges that have to be deleted in order to make the graph free of multiple edges.

### **Parameters**

edges: edge indices which we want to check. If None, all edges are checked.

### Return Value

a list of booleans, one for every edge given

## is mutual(edges=None)

Checks whether an edge has an opposite pair.

Also works for a set of edges – in this case, every edge is checked one by one. The result will be a list of booleans (or a single boolean if only an edge index is supplied), every boolean corresponding to an edge in the edge set supplied. True is returned for a given edge  $a \rightarrow b$  if there exists another edge  $b \rightarrow a$  in the original graph (not the given edge set!). All edges in an undirected graph are mutual. In case there are multiple edges between a and b, it is enough to have at least one edge in either direction to report all edges between them as mutual, so the multiplicity of edges do not matter.

### **Parameters**

edges: edge indices which we want to check. If None, all edges are checked.

### Return Value

a list of booleans, one for every edge given

## is separator(vertices)

Decides whether the removal of the given vertices disconnects the graph.

## **Parameters**

vertices: a single vertex ID or a list of vertex IDs

### Return Value

True is the given vertex set is a separator, False if not.

## is simple()

Checks whether the graph is simple (no loop or multiple edges).

## Return Value

True if it is simple, False otherwise.

(type=boolean)

## isoclass(vertices)

Returns the isomorphism class of the graph or its subgraph.

Isomorphy class calculations are implemented only for graphs with 3 or 4 vertices.

#### **Parameters**

vertices: a list of vertices if we want to calculate the isomorphism class for only a subset of vertices. None means to use the full graph.

### Return Value

the isomorphism class of the (sub)graph

# isomorphic(other)

Checks whether the graph is isomorphic to another graph.

The algorithm being used is selected using a simple heuristic:

- If one graph is directed and the other undirected, an exception is thrown.
- If the two graphs does not have the same number of vertices and edges, it returns with False
- If the graphs have three or four vertices, then an O(1) algorithm is used with precomputed data.
- Otherwise if the graphs are directed, then the VF2 isomorphism algorithm is used (see Graph.isomorphic\_vf2).
- Otherwise the BLISS isomorphism algorithm is used, see Graph.isomorphic\_bliss.

### Return Value

True if the graphs are isomorphic, False otherwise.

isomorphic bliss(other, return mapping 12=False, return mapping 21=False, sh1="fm", sh2=None, color1=None, color2 = None

Checks whether the graph is isomorphic to another graph, using the BLISS isomorphism algorithm.

See http://www.tcs.hut.fi/Software/bliss/index.html for more information about the BLISS algorithm.

## **Parameters**

the other graph with which we want to other:

compare the graph.

optional vector storing the coloring of the color1:

vertices of the first graph. If None, all vertices

have the same color.

color2: optional vector storing the coloring of the

vertices of the second graph. If None, all

vertices have the same color.

return\_mapping\_12: if True, calculates the mapping which maps

the vertices of the first graph to the second.

return\_mapping\_21: if True, calculates the mapping which maps

the vertices of the second graph to the first.

sh1: splitting heuristics for the first graph as a

case-insensitive string, with the following

possible values:

• "f": first non-singleton cell

• "f1": first largest non-singleton cell

• "fs": first smallest non-singleton cell

• "fm": first maximally non-trivially connected non-singleton cell

• "flm": largest maximally non-trivially connected non-singleton cell

• "fsm": smallest maximally non-trivially connected non-singleton cell

sh2: splitting heuristics to be used for the second

> graph. This must be the same as sh1; alternatively, it can be None, in which case it will automatically use the same value as sh1.

Currently it is present for backwards

compatibility only.

### Return Value

if no mapping is calculated, the result is True if the graphs are isomorphic, False otherwise. If any 5 or both mappings are calculated, the result is a 3-tuple, the first element being the above mentioned boolean, the second element being the 1 -> 2 mapping and the third element being the 2 -> 1 mapping. If the corresponding mapping was not calculated. None is returned in the

**isomorphic** vf2(other=None, color1=None, color2=None, edge color1=None, edge color2=None, return mapping 12=False, return mapping 21=False, node compat fn=None, edge compat fn=None, callback=None)

Checks whether the graph is isomorphic to another graph, using the VF2 isomorphism algorithm.

Vertex and edge colors may be used to restrict the isomorphisms, as only vertices and edges with the same color will be allowed to match each other.

#### **Parameters**

other: the other graph with which we want to

compare the graph. If None, the

automorphisms of the graph will be tested.

optional vector storing the coloring of the color1:

vertices of the first graph. If None, all vertices

have the same color.

color2: optional vector storing the coloring of the

vertices of the second graph. If None, all

vertices have the same color.

optional vector storing the coloring of the edge\_color1:

edges of the first graph. If None, all edges

have the same color.

optional vector storing the coloring of the edge\_color2:

edges of the second graph. If None, all edges

have the same color.

return\_mapping\_12: if True, calculates the mapping which maps

the vertices of the first graph to the second.

return\_mapping\_21: if True, calculates the mapping which maps

the vertices of the second graph to the first.

if not None, the isomorphism search will not callback:

> stop at the first match; it will call this callback function instead for every

isomorphism found. The callback function must accept four arguments: the first graph,

the second graph, a mapping from the nodes of the first graph to the second, and a

mapping from the nodes of the second graph to the first. The function must return True if

the search should continue or False otherwise.

a function that receives the two graphs and node\_compat\_fn:

> two node indices (one from the first graph, one from the second graph) and returns True if the nodes186 ven by the two indices are compatible (i.e. they could be matched to each other) or False otherwise. This can be used to restrict the set of isomorphisms based

on node-specific criteria that are too

# knn(vids=None, weights=None)

Calculates the average degree of the neighbors for each vertex, and the same quantity as the function of vertex degree.

### **Parameters**

vids: the vertices for which the calculation is performed. None

means all vertices.

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name. If this is given, the vertex strength will be used instead of the vertex degree in the calculations, but the "ordinary" vertex degree will be used

for the second (degree- dependent) list in the result.

#### Return Value

two lists in a tuple. The first list contains the average degree of neighbors for each vertex, the second contains the average degree of neighbors as a function of vertex degree. The zeroth element of this list corresponds to vertices of degree 1.

## laplacian(weights=None, normalized=False)

Returns the Laplacian matrix of a graph.

The Laplacian matrix is similar to the adjacency matrix, but the edges are denoted with -1 and the diagonal contains the node degrees.

Normalized Laplacian matrices have 1 or 0 in their diagonals (0 for vertices with no edges), edges are denoted by  $1 / \operatorname{sqrt}(d_i * d_j)$  where  $d_i$  is the degree of node i.

Multiple edges and self-loops are silently ignored. Although it is possible to calculate the Laplacian matrix of a directed graph, it does not make much sense.

# **Parameters**

weights: edge weights to be used. Can be a sequence or iterable

or even an edge attribute name. When edge weights are used, the degree of a node is considered to be the

weight of its incident edges.

normalized: whether to return the normalized Laplacian matrix.

### Return Value

the Laplacian matrix.

## largest cliques()

Returns the largest cliques of the graph as a list of tuples.

Quite intuitively a clique is considered largest if there is no clique with more vertices in the whole graph. All largest cliques are maximal (i.e. nonextendable) but not all maximal cliques are largest.

See Also: clique\_number() for the size of the largest cliques or maximal\_cliques() for the maximal cliques

## largest independent vertex sets()

Returns the largest independent vertex sets of the graph as a list of tuples.

Quite intuitively an independent vertex set is considered largest if there is no other set with more vertices in the whole graph. All largest sets are maximal (i.e. nonextendable) but not all maximal sets are largest.

See Also: independence\_number() for the size of the largest independent vertex sets or maximal\_independent\_vertex\_sets() for the maximal (nonextendable) independent vertex sets

# layout bipartite(types="type", hgap=1, vgap=1, maxiter=100)

Place the vertices of a bipartite graph in two layers.

The layout is created by placing the vertices in two rows, according to their types. The positions of the vertices within the rows are then optimized to minimize the number of edge crossings using the heuristic used by the Sugiyama layout algorithm.

#### **Parameters**

types: an igraph vector containing the vertex types, or an

attribute name. Anything that evalulates to False

corresponds to vertices of the first kind, everything else to

the second kind.

hgap: minimum horizontal gap between vertices in the same

laver.

vgap: vertical gap between the two layers.

maxiter: maximum number of iterations to take in the crossing

reduction step. Increase this if you feel that you are

getting too many edge crossings.

### Return Value

# $\mathbf{layout} \quad \mathbf{circle}(\mathit{dim}{=}2, \mathit{order}{=}\mathtt{None})$

Places the vertices of the graph uniformly on a circle or a sphere.

# **Parameters**

dim: the desired number of dimensions for the layout. dim=2

means a 2D layout, dim=3 means a 3D layout.

order: the order in which the vertices are placed along the circle.

Not supported when dim is not equal to 2.

# Return Value

```
layout davidson harel(seed=None, maxiter=10, fineiter=-1,
cool fact=0.75, weight node dist=1.0, weight border=0.0,
weight edge lengths=-1, weight edge crossings=-1,
weight node edge dist=-1)
```

Places the vertices on a 2D plane according to the Davidson-Harel layout algorithm.

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.

#### Parameters

seed: if None, uses a random starting layout for

> the algorithm. If a matrix (list of lists), uses the given matrix as the starting

position.

Number of iterations to perform in the maxiter:

annealing phase.

fineiter: Number of iterations to perform in the

> fine-tuning phase. Negative numbers set up a reasonable default from the base-2 logarithm of the vertex count, bounded

by 10 from above.

Cooling factor of the simulated annealing cool\_fact:

phase.

Weight for the node-node distances in the weight\_node\_dist:

energy function.

Weight for the distance from the border weight\_border:

> component of the energy function. Zero means that vertices are allowed to sit on the border of the area designated for the

layout.

weight\_edge\_lengths: Weight for the edge length component of

> the energy function. Negative numbers are replaced by the density of the graph

divided by 10.

weight\_edge\_crossings: Weight for the edge crossing component

of the energy function. Negative numbers are replaced by one minus the square root of the density of the graph.

weight\_node\_edge\_dist: Weight for the node-edge distance

component of the energy function. Negative numbers are replaced by 0.2

minus 0.2 times the density of the graph.

drl(weights=None, fixed=None, seed=None, options=None, dim=2)

Places the vertices on a 2D plane or in the 3D space ccording to the DrL layout algorithm.

This is an algorithm suitable for quite large graphs, but it can be surprisingly slow for small ones (where the simpler force-based layouts like layout\_kamada\_kawai() or layout\_fruchterman\_reingold() are more useful.

## **Parameters**

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name.

seed: if None, uses a random starting layout for the algorithm.

If a matrix (list of lists), uses the given matrix as the

starting position.

fixed: if a seed is given, you can specify some vertices to be kept

fixed at their original position in the seed by passing an appropriate list here. The list must have exactly as many items as the number of vertices in the graph. Items of the list that evaluate to **True** denote vertices that will not be

moved.

options: if you give a string argument here, you can select from five default preset parameterisations: default coarsen for a

maximal edge cutting.

default preset parameterisations: default, coarsen for a coarser layout, coarsest for an even coarser layout, refine for refining an existing layout and final for finalizing a layout. If you supply an object that is not a string, the DrL layout parameters are retrieved from the respective keys of the object (so it should be a dict or something else that supports the mapping protocol). The following keys can be used:

- 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
- init\_iterations: number of iterations in the initialization phase
- init\_temperature: start temperature during initialization
- init\_attraction: attraction during initialization
- init\_damping\_mult: damping multiplier during initialization
- liquid\_iterations, liquid\_temperature, liquid\_attraction, liquid\_damping\_mult: same parameters for the liquid phase
- expansion\_iterations, expansion\_temperature,

 $\begin{array}{l} \textbf{layout\_fruchterman\_reingold}(weights=\texttt{None},\ niter=\texttt{500},\ seed=\texttt{None},\ start\_temp=\texttt{None},\ minx=\texttt{None},\ maxx=\texttt{None},\ miny=\texttt{None},\ maxy=\texttt{None},\ minz=\texttt{None},\ maxz=\texttt{None},\ grid=\texttt{"auto"}) \end{array}$ 

Places the vertices on a 2D plane according to the Fruchterman-Reingold algorithm.

This is a force directed layout, see Fruchterman, T. M. J. and Reingold, E. M.: Graph Drawing by Force-directed Placement. Software – Practice and Experience, 21/11, 1129–1164, 1991

#### **Parameters**

weights: edge weights to be used. Can be a sequence or iterable

or even an edge attribute name.

niter: the number of iterations to perform. The default is

500.

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. The default is the square root of the number of vertices divided by 10.

minx: if not None, it must be a vector with exactly as many

elements as there are vertices in the graph. Each element is a minimum constraint on the X value of the

vertex in the layout.

maxx: similar to minx, but with maximum constraints

miny: similar to minx, but with the Y coordinates

maxy: similar to maxx, but with the Y coordinates

minz: similar to minx, but with the Z coordinates. Use only

for 3D layouts (dim=3).

maxz: similar to maxx, but with the Z coordinates. Use only

for 3D layouts (dim=3).

seed: if None, uses a random starting layout for the

algorithm. If a matrix (list of lists), uses the given

matrix as the starting position.

grid: whether to use a faster, but less accurate grid-based

implementation of the algorithm. "auto" decides based on the number of vertices in the graph; a grid will be used if there are at least 1000 vertices. "grid" is equivalent to True, "nogrid" is equivalent to False.

### Return Value

layout\_graphopt(niter=500, node\_charge=0.001, node\_mass=30,
spring length=0, spring constant=1, max sa movement=5, seed=None)

This 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.

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 or the maximal number of iterations is reached.

See http://www.schmuhl.org/graphopt/ for the original graphopt.

## **Parameters**

niter: the number of iterations to perform. Should be a

couple of hundred in general.

node\_charge: the charge of the vertices, used to calculate

electric repulsion.

node\_mass: the mass of the vertices, used for the spring

forces

spring\_length: the length of the springs

spring\_constant: the spring constant

max\_sa\_movement: the maximum amount of movement allowed in a

single step along a single axis.

seed: a matrix containing a seed layout from which

the algorithm will be started. If None, a random

layout will be used.

### Return Value

# layout grid(width=0, height=0, dim=2)

Places the vertices of a graph in a 2D or 3D grid.

### **Parameters**

width: the number of vertices in a single row of the layout. Zero or

negative numbers mean that the width should be

determined automatically.

height: the number of vertices in a single column of the layout.

Zero or negative numbers mean that the height should be determined automatically. It must not be given if the

number of dimensions is 2.

dim: the desired number of dimensions for the layout. dim=2

means a 2D layout, dim=3 means a 3D layout.

### Return Value

 $\begin{array}{l} \textbf{layout\_kamada\_kawai}(maxiter=1000,\,seed=\texttt{None},\,maxiter=1000,\\epsilon=0,\,kkconst=\texttt{None},\,minx=\texttt{None},\,maxx=\texttt{None},\,miny=\texttt{None},\\maxy=\texttt{None},\,minz=\texttt{None},\,maxz=\texttt{None},\,dim=2) \end{array}$ 

Places the vertices on a plane according to the Kamada-Kawai algorithm.

This is a force directed layout, see Kamada, T. and Kawai, S.: An Algorithm for Drawing General Undirected Graphs. Information Processing Letters, 31/1, 7–15, 1989.

#### **Parameters**

maxiter: the maximum number of iterations to perform.

seed: if None, uses a random starting layout for the algorithm.

If a matrix (list of lists), uses the given matrix as the

starting position.

epsilon: quit if the energy of the system changes less than epsilon.

See the original paper for details.

kkconst: the Kamada-Kawai vertex attraction constant. None

means the square of the number of vertices.

minx: if not None, it must be a vector with exactly as many

elements as there are vertices in the graph. Each element is a minimum constraint on the X value of the vertex in

the layout.

maxx: similar to minx, but with maximum constraints

miny: similar to minx, but with the Y coordinates

maxy: similar to maxx, but with the Y coordinates

minz: similar to minx, but with the Z coordinates. Use only for

3D layouts (dim=3).

maxz: similar to maxx, but with the Z coordinates. Use only for

3D layouts (dim=3).

dim: the desired number of dimensions for the layout. dim=2

means a 2D layout, dim=3 means a 3D layout.

#### Return Value

layout\_lgl(maxiter=150, maxdelta=-1, area=-1, coolexp=1.5,
repulserad=-1, cellsize=-1, root=None)

Places the vertices on a 2D plane according to the Large Graph Layout.

#### **Parameters**

maxiter: the number of iterations to perform.

maxdelta: the maximum distance to move a vertex in an

iteration. If negative, defaults to the number of

vertices.

area: the area of the square on which the vertices will be

placed. If negative, defaults to the number of vertices

squared.

coolexp: the cooling exponent of the simulated annealing.

repulserad: determines the radius at which vertex-vertex repulsion

cancels out attraction of adjacent vertices. If negative,

defaults to area times the number of vertices.

cellsize: the size of the grid cells. When calculating the

repulsion forces, only vertices in the same or

neighboring grid cells are taken into account. Defaults

to the fourth root of area.

root: the root vertex, this is placed first, its neighbors in the

first iteration, second neighbors in the second, etc. None means that a random vertex will be chosen.

## Return Value

## layout mds(dist=None, dim=2, arpack options=None)

Places the vertices in an Euclidean space with the given number of dimensions using multidimensional scaling.

This layout requires a distance matrix, where the intersection of row i and column j specifies the desired distance between vertex i and vertex j. The algorithm will try to place the vertices in a way that approximates the distance relations prescribed in the distance matrix. igraph uses the classical multidimensional scaling by Torgerson (see reference below).

For unconnected graphs, the method will decompose the graph into weakly connected components and then lay out the components individually using the appropriate parts of the distance matrix.

#### **Parameters**

dist: the distance matrix. It must be symmetric and

the symmetry is not checked – results are unspecified when a non-symmetric distance matrix is used. If this parameter is None, the shortest path lengths will be used as distances. Directed graphs are treated as undirected when calculating the shortest path lengths to ensure

symmetry.

dim: the number of dimensions. For 2D layouts, supply

2 here; for 3D layouts, supply 3.

arpack\_options: an ARPACKOptions object used to fine-tune the

ARPACK eigenvector calculation. If omitted, the module-level variable called arpack\_options is

used.

## Return Value

the calculated layout.

**Reference:** Cox & Cox: Multidimensional Scaling (1994), Chapman and Hall, London.

## layout random(dim=2)

Places the vertices of the graph randomly.

### **Parameters**

dim: the desired number of dimensions for the layout. dim=2 means a 2D layout, dim=3 means a 3D layout.

#### Return Value

the coordinate pairs in a list.

layout reingold tilford(mode="out", root=None, rootlevel=None)

Places the vertices on a 2D plane according to the Reingold-Tilford layout algorithm.

This is a tree layout. If the given graph is not a tree, a breadth-first search is executed first to obtain a possible spanning tree.

#### **Parameters**

mode: specifies which edges to consider when builing 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 behaviour in igraph 0.5 and before). This parameter also influences how the root vertices are calculated if they are not

given. See the root parameter.

root: the index of the root vertex or root vertices. if this is a

non-empty vector then the supplied vertex IDs are used as the roots of the trees (or a single tree if the graph is connected. If this is None or an empty list, the root

vertices are automatically calculated based on

topological sorting, performed with the opposite of the

mode argument.

rootlevel: this argument is useful when drawing forests which are

not trees. It specifies the level of the root vertices for

every tree in the forest.

# Return Value

the calculated layout.

See Also: layout reingold tilford circular

Reference: EM Reingold, JS Tilford: Tidier Drawings of Trees. IEEE

Transactions on Software Engineering 7:22, 223-228, 1981.

Circular Reingold-Tilford layout for trees.

This layout is similar to the Reingold-Tilford layout, but the vertices are placed in a circular way, with the root vertex in the center.

See layout\_reingold\_tilford for the explanation of the parameters.

#### Return Value

the calculated layout.

See Also: layout reingold tilford

Reference: EM Reingold, JS Tilford: Tidier Drawings of Trees. IEEE

Transactions on Software Engineering 7:22, 223-228, 1981.

## layout star(center=0, order=None)

Calculates a star-like layout for the graph.

#### Parameters

center: the ID of the vertex to put in the center

order: a numeric vector giving the order of the vertices (including

the center vertex!). If it is None, the vertices will be placed

in increasing vertex ID order.

## Return Value

the calculated layout.

### linegraph()

Returns the line graph of the graph.

The line graph L(G) of an undirected graph is defined as follows: L(G) has one vertex for each edge in G and two vertices in L(G) are connected iff their corresponding edges in the original graph share an end point.

The line graph of a directed graph is slightly different: two vertices are connected by a directed edge iff the target of the first vertex's corresponding edge is the same as the source of the second vertex's corresponding edge.

## maxdegree(vertices=None, mode=ALL, loops=False)

Returns the maximum degree of a vertex set in the graph.

This method accepts a single vertex ID or a list of vertex IDs as a parameter, and returns the degree of the given vertices (in the form of a single integer or a list, depending on the input parameter).

#### **Parameters**

vertices: a single vertex ID or a list of vertex IDs, or None

meaning all the vertices in the graph.

mode: the type of degree to be returned (OUT for out-degrees,

IN IN for in-degrees or ALL for the sum of them).

loops: whether self-loops should be counted.

## maxflow(source, target, capacity=None)

Returns the maximum flow between the source and target vertices.

#### **Parameters**

source: the source vertex ID
target: the target vertex ID

capacity: the capacity of the edges. It must be a list or a valid

attribute name or None. In the latter case, every edge

will have the same capacity.

### Return Value

a tuple containing the following: the value of the maximum flow between the given vertices, the flow value on all the edges, the edge IDs that are part of the corresponding minimum cut, and the vertex IDs on one side of the cut. For directed graphs, the flow value vector gives the flow value on each edge. For undirected graphs, the flow value is positive if the flow goes from the smaller vertex ID to the bigger one and negative if the flow goes from the bigger vertex ID to the smaller.

**Attention:** this function has a more convenient interface in class **Graph** which wraps the result in a **Flow** object. It is advised to use that.

## **maxflow** value(source, target, capacity=None)

Returns the value of the maximum flow between the source and target vertices.

#### Parameters

source: the source vertex ID

target: the target vertex ID

capacity: the capacity of the edges. It must be a list or a valid

attribute name or None. In the latter case, every edge

will have the same capacity.

### Return Value

the value of the maximum flow between the given vertices

## maximal cliques(min=0, max=0, file=None)

Returns the maximal cliques of the graph as a list of tuples.

A maximal clique is a clique which can't be extended by adding any other vertex to it. A maximal clique is not necessarily one of the largest cliques in the graph.

### **Parameters**

min: the minimum size of maximal cliques to be returned. If zero or negative, no lower bound will be used.

max: the maximum size of maximal cliques to be returned. If zero or negative, no upper bound will be used. If nonzero, the size of every maximal clique found will be compared to this value and a clique will be returned only if its size is smaller than this limit.

file: a file object or the name of the file to write the results to. When this argument is None, the maximal cliques will be returned as a list of lists.

### Return Value

the maximal cliques of the graph as a list of lists, or None if the file argument was given.@see: largest\_cliques() for the largest cliques.

## maximal independent vertex sets()

Returns the maximal independent vertex sets of the graph as a list of tuples.

A maximal independent vertex set is an independent vertex set which can't be extended by adding any other vertex to it. A maximal independent vertex set is not necessarily one of the largest independent vertex sets in the graph.

See Also: largest\_independent\_vertex\_sets() for the largest independent vertex sets

**Reference:** S. Tsukiyama, M. Ide, H. Ariyoshi and I. Shirawaka: A new algorithm for generating all the maximal independent sets. SIAM J Computing, 6:505–517, 1977.

## **mincut**(source=None, target=None, capacity=None)

Calculates the minimum cut between the source and target vertices or within the whole graph.

The minimum cut is the minimum set of edges that needs to be removed to separate the source and the target (if they are given) or to disconnect the graph (if the source and target are not given). The minimum is calculated using the weights (capacities) of the edges, so the cut with the minimum total capacity is calculated. For undirected graphs and no source and target, the method uses the Stoer-Wagner algorithm. For a given source and target, the method uses the push-relabel algorithm; see the references below.

#### **Parameters**

source: the source vertex ID. If None, target must also be

{None} and the calculation will be done for the entire

graph (i.e. all possible vertex pairs).

target: the target vertex ID. If None, source must also be

{None} and the calculation will be done for the entire

graph (i.e. all possible vertex pairs).

capacity: the capacity of the edges. It must be a list or a valid

attribute name or None. In the latter case, every edge

will have the same capacity.

### Return Value

the value of the minimum cut, the IDs of vertices in the first and second partition, and the IDs of edges in the cut, packed in a 4-tuple

Attention: this function has a more convenient interface in class Graph which wraps the result in a Cut object. It is advised to use that.

#### Reference:

- M. Stoer, F. Wagner: A simple min-cut algorithm. Journal of the ACM 44(4):585-591, 1997.
- A. V. Goldberg, R. E. Tarjan: A new approach to the maximum-flow problem. Journal of the ACM 35(4):921-940, 1988.

## mincut value(source=-1, target=-1, capacity=None)

Returns the minimum cut between the source and target vertices or within the whole graph.

### **Parameters**

source: the source vertex ID. If negative, the calculation is done

for every vertex except the target and the minimum is

returned.

target: the target vertex ID. If negative, the calculation is done

for every vertex except the source and the minimum is

returned.

capacity: the capacity of the edges. It must be a list or a valid

attribute name or None. In the latter case, every edge

will have the same capacity.

#### Return Value

the value of the minimum cut between the given vertices

# minimum size separators()

Returns a list containing all separator vertex sets of minimum size.

A vertex set is a separator if its removal disconnects the graph. This method lists all the separators for which no smaller separator set exists in the given graph.

### Return Value

a list where each item lists the vertex indices of a given separator of minimum size.

**Reference:** Arkady Kanevsky: Finding all minimum-size separating vertex sets in a graph. Networks 23:533–541, 1993.

## modularity(membership, weights=None)

Calculates the modularity of the graph with respect to some vertex types.

The modularity of a graph w.r.t. some division measures how good the division is, or how separated are the different vertex types from each other. It is defined as Q=1/(2m) \* sum(Aij-ki\*kj/(2m)delta(ci,cj),i,j). m is the number of edges, Aij is the element of the A adjacency matrix in row i and column j, ki is the degree of node i, kj is the degree of node j, and Ci and cj are the types of the two vertices (i and j). delta(x,y) is one iff x=y, 0 otherwise.

If edge weights are given, the definition of modularity is modified as follows: Aij becomes the weight of the corresponding edge, ki is the total weight of edges incident on vertex i, kj is the total weight of edges incident on vertex j and m is the total edge weight in the graph.

### **Parameters**

membership: the membership vector, e.g. the vertex type index for

each vertex.

weights: optional edge weights or None if all edges are weighed

equally.

### Return Value

the modularity score. Score larger than 0.3 usually indicates strong community structure.

Attention: method overridden in Graph to allow VertexClustering objects as a parameter. This method is not strictly necessary, since the VertexClustering class provides a variable called modularity.

**Reference:** MEJ Newman and M Girvan: Finding and evaluating community structure in networks. Phys Rev E 69 026113, 2004.

## motifs randesu(size=3, cut prob=None, callback=None)

Counts the number of motifs in the graph

Motifs are small subgraphs of a given structure in a graph. It is argued that the motif profile (ie. the number of different motifs in the graph) is characteristic for different types of networks and network function is related to the motifs in the graph.

This function is able to find the different motifs of size three and four (ie. the number of different subgraphs with three and four vertices) in the network.

In a big network the total number of motifs can be very large, so it takes a lot of time to find all of them. In such cases, a sampling method can be used. This function is capable of doing sampling via the  $cut\_prob$  argument. This argument gives the probability that a branch of the motif search tree will not be explored.

## **Parameters**

size: the size of the motifs (3 or 4).

cut\_prob: the cut probabilities for different levels of the search tree.

This must be a list of length *size* or None to find all

motifs.

callback: None or a callable that will be called for every motif

found in the graph. The callable must accept three parameters: the graph itself, the list of vertices in the motif and the isomorphism class of the motif (see Graph.isoclass()). The search will stop when the callback returns an object with a non-zero truth value or

raises an exception.

### Return Value

the list of motifs if callback is None, or None otherwise

**Reference:** S. Wernicke and F. Rasche: FANMOD: a tool for fast network motif detection, Bioinformatics 22(9), 1152–1153, 2006.

See Also: Graph.motifs randesu no()

# motifs randesu estimate(size=3, cut\_prob=None, sample)

Counts the total number of motifs in the graph

Motifs are small subgraphs of a given structure in a graph. This function estimates the total number of motifs in a graph without assigning isomorphism classes to them by extrapolating from a random sample of vertices.

#### **Parameters**

size: the size of the motifs (3 or 4).

cut\_prob: the cut probabilities for different levels of the search tree.

This must be a list of length size or None to find all

motifs.

sample: the size of the sample or the vertex IDs of the vertices to

be used for sampling.

Reference: S. Wernicke and F. Rasche: FANMOD: a tool for fast network

motif detection, Bioinformatics 22(9), 1152–1153, 2006.

See Also: Graph.motifs randesu()

# motifs randesu no(size=3, cut prob=None)

Counts the total number of motifs in the graph

Motifs are small subgraphs of a given structure in a graph. This function counts the total number of motifs in a graph without assigning isomorphism classes to them.

## **Parameters**

size: the size of the motifs (3 or 4).

cut\_prob: the cut probabilities for different levels of the search tree.

This must be a list of length *size* or None to find all

motifs.

Reference: S. Wernicke and F. Rasche: FANMOD: a tool for fast network

motif detection, Bioinformatics 22(9), 1152–1153, 2006.

See Also: Graph.motifs randesu()

## neighborhood(vertices=None, order=1, mode=ALL, mindist=0)

For each vertex specified by *vertices*, returns the vertices reachable from that vertex in at most *order* steps. If *mindist* is larger than zero, vertices that are reachable in less than *mindist* steps are excluded.

## Parameters

vertices: a single vertex ID or a list of vertex IDs, or None

meaning all the vertices in the graph.

order: the order of the neighborhood, i.e. the maximum number

of steps to take from the seed vertex.

mode: specifies how to take into account the direction of the

edges if a directed graph is analyzed. "out" means that only the outgoing edges are followed, so all vertices reachable from the source vertex in at most *order* steps are counted. "in" means that only the incoming edges are followed (in reverse direction of course), so all vertices from which the source vertex is reachable in at

vertices from which the source vertex is reachable in at most *order* steps are counted. "all" treats directed

edges as undirected.

mindist: the minimum distance required to include a vertex in the

result. If this is one, the seed vertex is not included. If this is two, the direct neighbors of the seed vertex are

not included either, and so on.

## Return Value

a single list specifying the neighborhood if *vertices* was an integer specifying a single vertex index, or a list of lists if *vertices* was a list or None.

## neighborhood size(vertices=None, order=1, mode=ALL, mindist=0)

For each vertex specified by *vertices*, returns the number of vertices reachable from that vertex in at most *order* steps. If *mindist* is larger than zero, vertices that are reachable in less than *mindist* steps are excluded.

## Parameters

vertices: a single vertex ID or a list of vertex IDs, or None

meaning all the vertices in the graph.

order: the order of the neighborhood, i.e. the maximum number

of steps to take from the seed vertex.

mode: specifies how to take into account the direction of the

edges if a directed graph is analyzed. "out" means that only the outgoing edges are followed, so all vertices reachable from the source vertex in at most *order* steps are counted. "in" means that only the incoming edges are followed (in reverse direction of course), so all vertices from which the source vertex is reachable in at

most order steps are counted. "all" treats directed

edges as undirected.

mindist: the minimum distance required to include a vertex in the

result. If this is one, the seed vertex is not counted. If this is two, the direct neighbors of the seed vertex are

not counted either, and so on.

## Return Value

a single number specifying the neighborhood size if *vertices* was an integer specifying a single vertex index, or a list of sizes if *vertices* was a list or None.

## **neighbors**(vertex, mode=ALL)

Returns adjacent vertices to a given vertex.

#### **Parameters**

vertex: a vertex ID

mode: whether to return only successors (OUT), predecessors (IN)

or both (ALL). Ignored for undirected graphs.

## path length hist(directed=True)

Calculates the path length histogram of the graph

### **Parameters**

directed: whether to consider directed paths

#### Return Value

a tuple. The first item of the tuple is a list of path lengths, the *i*th element of the list contains the number of paths with length i+1. The second item contains the number of unconnected vertex pairs as a float (since it might not fit into an integer)

Attention: this function is wrapped in a more convenient syntax in the derived class Graph. It is advised to use that instead of this version.

## permute vertices(permutation)

Permutes the vertices of the graph according to the given permutation and returns the new graph.

Vertex k of the original graph will become vertex permutation[k] in the new graph. No validity checks are performed on the permutation vector.

## Return Value

the new graph

 $\label{eq:personalized_pagerank} \begin{aligned} & \texttt{pagerank}(vertices = \texttt{None}, \ directed = \texttt{True}, \ damping = \texttt{0.85}, \\ & reset = \texttt{None}, \ reset\_vertices = \texttt{None}, \ weights = \texttt{None}, \ arpack\_options = \texttt{None}, \\ & implementation = \texttt{"prpack"}, \ niter = \texttt{1000}, \ eps = \texttt{0.001}) \end{aligned}$ 

Calculates the personalized PageRank values of a graph.

The personalized PageRank calculation is similar to the PageRank calculation, but the random walk is reset to a non-uniform distribution over the vertices in every step with probability 1-damping instead of a uniform distribution.

#### **Parameters**

vertices: the indices of the vertices being queried. None

means all of the vertices.

directed: whether to consider directed paths.

damping: the damping factor. 1-damping is the PageRank

value for vertices with no incoming links.

reset: the distribution over the vertices to be used when

resetting the random walk. Can be a sequence, an iterable or a vertex attribute name as long as they return a list of floats whose length is equal to the number of vertices. If None, a uniform distribution is assumed, which makes the method equivalent to the original PageRank algorithm.

reset\_vertices: an alternative way to specify the distribution over

the vertices to be used when resetting the random walk. Simply supply a list of vertex IDs here, or a VertexSeq or a Vertex. Resetting will take place using a uniform distribution over the specified

vertices.

weights: edge weights to be used. Can be a sequence or

iterable or even an edge attribute name.

arpack\_options: an ARPACKOptions object used to fine-tune the

ARPACK eigenvector calculation. If omitted, the module-level variable called arpack\_options is used. This argument is ignored if not the ARPACK implementation is used, see the

implementation argument.

implementation: which implementation to use to solve the PageRank eigenproblem. Possible values are:

- "prpack": use the PRPACK library. This is a new implementation in igraph 0.7
- "arpack": use the ARPACK library. This implementation was used from version 0.5, until version 0.7.
- "power": use a simple power method. This is the implementation that was used before igraph version 0.5.

niter: The number of iterations to use in the power

## predecessors(vertex)

Returns the predecessors of a given vertex.

Equivalent to calling the Graph.neighbors method with type=IN.

## radius(mode=OUT)

Calculates the radius of the graph.

The radius of a graph is defined as the minimum eccentricity of its vertices (see eccentricity()).

#### **Parameters**

mode: what kind of paths to consider for the calculation in case of directed graphs. OUT considers paths that follow edge directions, IN considers paths that follow the opposite edge directions, ALL ignores edge directions. The argument is ignored for undirected graphs.

### Return Value

the radius

See Also: Graph.eccentricity()

## random walk(start, steps, mode="out", stuck="return")

Performs a random walk of a given length from a given node.

### **Parameters**

start: the starting vertex of the walk

steps: the number of steps that the random walk should take

mode: whether to follow outbound edges only (OUT), inbound edges

only (IN) or both (ALL). Ignored for undirected

graphs.@param stuck: what to do when the random walk gets stuck. "return" returns a partial random walk;

"error" throws an exception.

# Return Value

a random walk that starts from the given vertex and has at most the given length (shorter if the random walk got stuck)

## reciprocity(ignore loops=True, mode="default")

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. This measure is calculated if mode is "default".

Prior to igraph 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) disconnected, (2) non-reciprocally connected and (3) reciprocally connected. The result is the size of group (3), divided by the sum of sizes of groups (2) and (3). This measure is calculated if mode is "ratio".

### Parameters

ignore\_loops: whether loop edges should be ignored.

mode: the algorithm to use to calculate the reciprocity; see

above for more details.

# Return Value

the reciprocity of the graph

## rewire(n=1000, mode="simple")

Randomly rewires the graph while preserving the degree distribution.

Please note that the rewiring is done "in-place", so the original graph will be modified. If you want to preserve the original graph, use the copy method before.

## **Parameters**

n: the number of rewiring trials.

 ${\tt mode:}\ the\ rewiring\ algorithm\ to\ use.$  It can either be  ${\tt "simple"}\ or$ 

"loops"; the former does not create or destroy loop edges

while the latter does.

## rewire edges(prob, loops=False, multiple=False)

Rewires the edges of a graph with constant probability.

Each endpoint of each edge of the graph will be rewired with a constant probability, given in the first argument.

Please note that the rewiring is done "in-place", so the original graph will be modified. If you want to preserve the original graph, use the copy method before.

### **Parameters**

prob: rewiring probability

loops: whether the algorithm is allowed to create loop edgesmultiple: whether the algorithm is allowed to create multiple edges.

## shortest paths(source=None, target=None, weights=None, mode=OUT)

Calculates shortest path lengths for given vertices in a graph.

The algorithm used for the calculations is selected automatically: a simple BFS is used for unweighted graphs, Dijkstra's algorithm is used when all the weights are positive. Otherwise, the Bellman-Ford algorithm is used if the number of requested source vertices is larger than 100 and Johnson's algorithm is used otherwise.

## **Parameters**

source: a list containing the source vertex IDs which should be

included in the result. If None, all vertices will be

considered.

target: a list containing the target vertex IDs which should be

included in the result. If None, all vertices will be

considered.

weights: a list containing the edge weights. It can also be an

attribute name (edge weights are retrieved from the given

attribute) or None (all edges have equal weight).

mode: the type of shortest paths to be used for the calculation in

directed graphs. OUT means only outgoing, IN means only incoming paths. ALL means to consider the directed graph

as an undirected one.

## Return Value

the shortest path lengths for given vertices in a matrix

 $\begin{array}{l} \mathbf{similarity\_dice}(\textit{vertices} = \texttt{None}, \textit{pairs} = \texttt{None}, \textit{mode} = \texttt{IGRAPH\_ALL}, \\ \textit{loops} = \texttt{True}) \end{array}$ 

Dice similarity coefficient of vertices.

The Dice similarity coefficient of two vertices is twice the number of their common neighbors divided by the sum of their degrees. This coefficient is very similar to the Jaccard coefficient, but usually gives higher similarities than its counterpart.

#### **Parameters**

vertices: the vertices to be analysed. If None and pairs is also

None, all vertices will be considered.

pairs: the vertex pairs to be analysed. If this is given, vertices

must be None, and the similarity values will be

calculated only for the given pairs. Vertex pairs must be

specified as tuples of vertex IDs.

mode: which neighbors should be considered for directed graphs.

Can be ALL, IN or OUT, ignored for undirected graphs.

loops: whether vertices should be considered adjacent to

themselves. Setting this to True assumes a loop edge for all vertices even if none is present in the graph. Setting this to False may result in strange results: nonadjacent vertices may have larger similarities compared to the case when an edge is added between them – however, this might be exactly the result you want to get.

## Return Value

the pairwise similarity coefficients for the vertices specified, in the form of a matrix if pairs is None or in the form of a list if pairs is not None.

## similarity inverse log weighted(vertices=None, mode=IGRAPH\_ALL)

Inverse log-weighted similarity coefficient of vertices.

Each vertex is assigned a weight which is  $1 / \log(\text{degree})$ . The log-weighted similarity of two vertices is the sum of the weights of their common neighbors.

### **Parameters**

vertices: the vertices to be analysed. If None, all vertices will be

considered.

mode: which neighbors should be considered for directed

graphs. Can be ALL, IN or OUT, ignored for undirected graphs. IN means that the weights are determined by the out-degrees, OUT means that the weights are determined

by the in-degrees.

### Return Value

the pairwise similarity coefficients for the vertices specified, in the form of a matrix (list of lists).

 $similarity\_jaccard(vertices=None, pairs=None, mode=IGRAPH\_ALL, loops=True)$ 

Jaccard similarity coefficient of vertices.

The Jaccard similarity coefficient of two vertices is the number of their common neighbors divided by the number of vertices that are adjacent to at least one of them.

## **Parameters**

vertices: the vertices to be analysed. If None and pairs is also

None, all vertices will be considered.

pairs: the vertex pairs to be analysed. If this is given, vertices

must be None, and the similarity values will be

calculated only for the given pairs. Vertex pairs must be

specified as tuples of vertex IDs.

mode: which neighbors should be considered for directed graphs.

Can be ALL, IN or OUT, ignored for undirected graphs.

loops: whether vertices should be considered adjacent to

themselves. Setting this to True assumes a loop edge for all vertices even if none is present in the graph. Setting this to False may result in strange results: nonadjacent vertices may have larger similarities compared to the case when an edge is added between them – however, this might be exactly the result you want to get.

## Return Value

the pairwise similarity coefficients for the vertices specified, in the form of a matrix if pairs is None or in the form of a list if pairs is not None.

simplify(multiple=True, loops=True, combine edges=None)

Simplifies a graph by removing self-loops and/or multiple edges.

For example, suppose you have a graph with an edge attribute named weight. graph.simplify(combine\_edges=max) will take the maximum of the weights of multiple edges and assign that weight to the collapsed edge. graph.simplify(combine\_edges=sum) will take the sum of the weights. You can also write graph.simplify(combine\_edges=dict(weight="sum")) or graph.simplify(combine\_edges=dict(weight=sum)), since sum is recognised both as a Python built-in function and as a string constant.

## **Parameters**

multiple: whether to remove multiple edges.

loops: whether to remove loops.

combine\_edges: specifies how to combine the attributes of multiple edges between the same pair of vertices into a single attribute. If it is None, only one of the edges will be kept and all the attributes will be lost. If it is a function, the attributes of multiple edges will be collected and passed on to that function which will return the new attribute value that has to be assigned to the single collapsed edge. It can also be one of the following string constants:

- "ignore": all the edge attributes will be ignored.
- "sum": the sum of the edge attribute values will be used for the new edge.
- "product": the product of the edge attribute values will be used for the new edge.
- "mean": the mean of the edge attribute values will be used for the new edge.
- "median": the median of the edge attribute values will be used for the new edge.
- "min": the minimum of the edge attribute values will be used for the new edge.
- "max": the maximum of the edge attribute values will be used for the new edge.
- "first": the attribute value of the first edge in the collapsed set will be used for the new edge.
- "last": the attribute value of the last edge in the collapsed set will be used for the new edge.
- "random": a randomly selected value will be used for the new edge
- "concat": the attribute values will be concatenated for the new edge.

You can also use a dict mapping edge attribute names to functions or the above string constants if

# st mincut(source, target, capacity=None)

Calculates the minimum cut between the source and target vertices in a graph.

## **Parameters**

source: the source vertex ID
target: the target vertex ID

capacity: the capacity of the edges. It must be a list or a valid

attribute name or None. In the latter case, every edge

will have the same capacity.

## Return Value

the value of the minimum cut, the IDs of vertices in the first and second partition, and the IDs of edges in the cut, packed in a 4-tuple

Attention: this function has a more convenient interface in class Graph which wraps the result in a list of Cut objects. It is advised to use that.

# strength(vertices, mode=ALL, loops=True, weights=None)

Returns the strength (weighted degree) of some vertices from the graph

This method accepts a single vertex ID or a list of vertex IDs as a parameter, and returns the strength (that is, the sum of the weights of all incident edges) of the given vertices (in the form of a single integer or a list, depending on the input parameter).

## **Parameters**

vertices: a single vertex ID or a list of vertex IDs

mode: the type of degree to be returned (OUT for out-degrees,

IN IN for in-degrees or ALL for the sum of them).

loops: whether self-loops should be counted.

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name. "None" means to treat the graph as unweighted, falling back to ordinary degree

calculations.

# subcomponent(v, mode = ALL)

Determines the indices of vertices which are in the same component as a given vertex.

# **Parameters**

v: the index of the vertex used as the source/destination

mode: if equals to IN, returns the vertex IDs from where the given vertex can be reached. If equals to OUT, returns the vertex IDs which are reachable from the given vertex. If equals to ALL, returns all vertices within the same component as the given vertex, ignoring edge directions. Note that this is not equal to calculating the union of the results of IN and OUT.

#### Return Value

the indices of vertices which are in the same component as a given vertex.

# subgraph edges(edges, delete vertices=True)

Returns a subgraph spanned by the given edges.

# **Parameters**

edges: a list containing the edge IDs which should be

included in the result.

delete\_vertices: if True, vertices not incident on any of the

specified edges will be deleted from the result. If

False, all vertices will be kept.

## Return Value

the subgraph

Checks whether a subgraph of the graph is isomorphic to another graph.

The optional domains argument may be used to restrict vertices that may match each other. You can also specify whether you are interested in induced subgraphs only or not.

## **Parameters**

other: the pattern graph we are looking for in the graph.

domains: a list of lists, one sublist belonging to each vertex

in the template graph. Sublist i contains the indices of the vertices in the original graph that may match vertex i in the template graph. None means that every vertex may match every other

vertex.

induced: whether to consider induced subgraphs only.

time\_limit: an optimal time limit in seconds. Only the

integral part of this number is taken into account. If the time limit is exceeded, the method will

throw an exception.

return\_mapping: when True, the function will return a tuple, where

the first element is a boolean denoting whether a subisomorphism has been found or not, and the second element describes the mapping of the vertices from the template graph to the original graph. When False, only the boolean is returned.

## Return Value

if no mapping is calculated, the result is True if the graph contains a subgraph that is isomorphic to the given template, False otherwise. If the mapping is calculated, the result is a tuple, the first element being the above mentioned boolean, and the second element being the mapping from the target to the original graph.

 $\begin{array}{l} \mathbf{subisomorphic\_vf2}(other,\ color1=\mathtt{None},\ color2=\mathtt{None},\ edge\_color1=\mathtt{None},\ edge\_color2=\mathtt{None},\ return\_mapping\_12=\mathtt{False},\ return\_mapping\_21=\mathtt{False},\ callback=\mathtt{None},\ node\_compat\_fn=\mathtt{None},\ edge\_compat\_fn=\mathtt{None}) \end{array}$ 

Checks whether a subgraph of the graph is isomorphic to another graph.

Vertex and edge colors may be used to restrict the isomorphisms, as only vertices and edges with the same color will be allowed to match each other.

#### Parameters

other: the other graph with which we want to

compare the graph.

color1: optional vector storing the coloring of the

vertices of the first graph. If None, all vertices

have the same color.

color2: optional vector storing the coloring of the

vertices of the second graph. If None, all

vertices have the same color.

edge\_color1: optional vector storing the coloring of the

edges of the first graph. If None, all edges

have the same color.

edge\_color2: optional vector storing the coloring of the

edges of the second graph. If None, all edges

have the same color.

return\_mapping\_12: if True, calculates the mapping which maps

the vertices of the first graph to the second. The mapping can contain -1 if a given node is

not mapped.

return\_mapping\_21: if True, calculates the mapping which maps

the vertices of the second graph to the first. The mapping can contain -1 if a given node is

not mapped.

callback: if not None, the subisomorphism search will

not stop at the first match; it will call this

callback function instead for every

subisomorphism found. The callback function must accept four arguments: the first graph, the second graph, a mapping from the nodes

of the first graph to the second, and a

mapping from the nodes of the second graph to the first. The function must return True if the search should continue or False otherwise.

node\_compat\_fn: a function that receives the two graphs and

two node inchizes (one from the first graph, one from the second graph) and returns True if the nodes given by the two indices are compatible (i.e. they could be matched to each other) or False otherwise. This can be

## successors(vertex)

Returns the successors of a given vertex.

Equivalent to calling the Graph.neighbors method with type=OUT.

# to directed(mutual=True)

Converts an undirected graph to directed.

## **Parameters**

mutual: True if mutual directed edges should be created for every undirected edge. If False, a directed edge with arbitrary

direction is created.

# to prufer()

Converts a tree graph into a Prufer sequence.

## Return Value

the Prufer sequence as a list

# to undirected(mode="collapse", combine\_edges=None)

Converts a directed graph to undirected.

## **Parameters**

mode: specifies what to do with multiple directed edges

going between the same vertex pair. True or "collapse" means that only a single edge should be created from multiple directed edges. False or "each" means that every edge will be kept (with the arrowheads removed). "mutual" creates one undirected edge for each mutual directed edge pair.

combine\_edges: specifies how to combine the attributes of multiple

edges between the same pair of vertices into a single attribute. See Graph.simplify() for more

details.

# topological sorting(mode=OUT)

Calculates a possible topological sorting of the graph.

Returns a partial sorting and issues a warning if the graph is not a directed acyclic graph.

#### **Parameters**

mode: if OUT, vertices are returned according to the forward topological order – all vertices come before their successors. If IN, all vertices come before their ancestors.

## Return Value

a possible topological ordering as a list

# transitivity avglocal undirected(mode="nan")

Calculates the average of the vertex transitivities of the graph.

The transitivity measures the probability that two neighbors of a vertex are connected. In case of the average local transitivity, this probability is calculated for each vertex and then the average is taken. Vertices with less than two neighbors require special treatment, they will either be left out from the calculation or they will be considered as having zero transitivity, depending on the *mode* parameter.

Note that this measure is different from the global transitivity measure (see transitivity\_undirected()) as it simply takes the average local transitivity across the whole network.

### **Parameters**

mode: defines how to treat vertices with degree less than two. If
 TRANSITIVITT\_ZERO or "zero", these vertices will have zero
 transitivity. If TRANSITIVITY\_NAN or "nan", these vertices
 will be excluded from the average.

See Also: transitivity\_undirected(), transitivity\_local\_undirected()

**Reference:** D. J. Watts and S. Strogatz: Collective dynamics of small-world networks. Nature 393(6884):440-442, 1998.

 $\begin{array}{l} \textbf{transitivity\_local\_undirected}(\textit{vertices} = \texttt{None}, \textit{mode} = \texttt{"nan"}, \\ \textit{weights} = \texttt{None}) \end{array}$ 

Calculates the local transitivity (clustering coefficient) of the given vertices in the graph.

The transitivity measures the probability that two neighbors of a vertex are connected. In case of the local transitivity, this probability is calculated separately for each vertex.

Note that this measure is different from the global transitivity measure (see transitivity\_undirected()) as it calculates a transitivity value for each vertex individually.

The traditional local transitivity measure applies for unweighted graphs only. When the weights argument is given, this function calculates the weighted local transitivity proposed by Barrat et al (see references).

## **Parameters**

vertices: a list containing the vertex IDs which should be included

in the result. None means all of the vertices.

mode: defines how to treat vertices with degree less than two. If

TRANSITIVITT\_ZERO or "zero", these vertices will have zero transitivity. If TRANSITIVITY\_NAN or "nan", these vertices will have NaN (not a number) as their transitivity.

weights: edge weights to be used. Can be a sequence or iterable or

even an edge attribute name.

## Return Value

the transitivities for the given vertices in a list

See Also: transitivity\_undirected(), transitivity\_avglocal\_undirected()

# Reference:

- Watts DJ and Strogatz S: Collective dynamics of small-world networks. Nature 393(6884):440-442, 1998.
- Barrat A, Barthelemy M, Pastor-Satorras R and Vespignani A: The architecture of complex weighted networks. PNAS 101, 3747 (2004). http://arxiv.org/abs/cond-mat/0311416.

# transitivity undirected(mode="nan")

Calculates the global transitivity (clustering coefficient) of the graph.

The transitivity measures the probability that two neighbors of a vertex are connected. More precisely, this is the ratio of the triangles and connected triplets in the graph. The result is a single real number. Directed graphs are considered as undirected ones.

Note that this measure is different from the local transitivity measure (see transitivity\_local\_undirected()) as it calculates a single value for the whole graph.

## **Parameters**

mode: if TRANSITIVITY\_ZERO or "zero", the result will be zero if the graph does not have any triplets. If "nan" or TRANSITIVITY\_NAN, the result will be NaN (not a number).

## Return Value

the transitivity

See Also: transitivity\_local\_undirected(), transitivity\_avglocal\_undirected()

**Reference:** S. Wasserman and K. Faust: Social Network Analysis: Methods and Applications. Cambridge: Cambridge University Press, 1994.

# triad census()

Triad census, as defined by Davis and Leinhardt

Calculating the triad census means classifying every triplets of vertices in a directed graph. A triplet can be in one of 16 states, these are listed in the documentation of the C interface of igraph.

Attention: this function has a more convenient interface in class Graph which wraps the result in a TriadCensus object. It is advised to use that. The name of the triplet classes are also documented there.

# unfold tree(sources=None, mode=OUT)

Unfolds the graph using a BFS to a tree by duplicating vertices as necessary.

## **Parameters**

sources: the source vertices to start the unfolding from. It should

be a list of vertex indices, preferably one vertex from each

connected component. You can use

Graph.topological\_sorting() to determine a suitable

set. A single vertex index is also accepted.

mode: which edges to follow during the BFS. OUT follows

outgoing edges, IN follows incoming edges, ALL follows

both. Ignored for undirected graphs.

## Return Value

the unfolded tree graph and a mapping from the new vertex indices to the old ones.

# vcount()

Counts the number of vertices.

## Return Value

the number of vertices in the graph.

(type=integer)

## vertex attributes()

## Return Value

the attribute name list of the graph's vertices

vertex\_connectivity(source=-1, target=-1, checks=True,
neighbors="error")

Calculates the vertex connectivity of the graph or between some vertices.

The vertex connectivity between two given vertices is the number of vertices that have to be removed in order to disconnect the two vertices into two separate components. This is also the number of vertex disjoint directed paths between the vertices (apart from the source and target vertices of course). The vertex connectivity of the graph is the minimal vertex connectivity over all vertex pairs.

This method calculates the vertex connectivity of a given vertex pair if both the source and target vertices are given. If none of them is given (or they are both negative), the overall vertex connectivity is returned.

# **Parameters**

source: the source vertex involved in the calculation.

target: the target vertex involved in the calculation.

checks: if the whole graph connectivity is calculated and this is

True, igraph performs some basic checks before

calculation. If the graph is not strongly connected, then the connectivity is obviously zero. If the minimum degree is one, then the connectivity is also one. These simple checks are much faster than checking the entire graph, therefore it is advised to set this to True. The parameter is ignored if the connectivity between two

given vertices is computed.

neighbors: tells igraph what to do when the two vertices are

connected. "error" raises an exception, "infinity"

returns infinity, "ignore" ignores the edge.

## Return Value

the vertex connectivity

# **write dimacs**(*f*, *source*, *target*, *capacity*=None)

Writes the graph in DIMACS format to the given file.

# **Parameters**

f: the name of the file to be written or a Python file handle

source: the source vertex ID
target: the target vertex ID

capacity: the capacities of the edges in a list. If it is not a list, the

corresponding edge attribute will be used to retrieve

capacities.

# write dot(f)

Writes the graph in DOT format to the given file.

DOT is the format used by the GraphViz<sup>a</sup> software package.

## **Parameters**

f: the name of the file to be written or a Python file handle

# write edgelist(f)

Writes the edge list of a graph to a file.

Directed edges are written in (from, to) order.

## **Parameters**

f: the name of the file to be written or a Python file handle

# write gml(f, creator=None, ids=None)

Writes the graph in GML format to the given file.

#### **Parameters**

f: the name of the file to be written or a Python file handle

creator: optional creator information to be written to the file. If

None, the current date and time is added.

ids: optional numeric vertex IDs to use in the file. This must

be a list of integers or None. If None, the id attribute of the vertices are used, or if they don't exist, numeric vertex

IDs will be generated automatically.

<sup>&</sup>lt;sup>a</sup>http://www.graphviz.org

# write graphml(f)

Writes the graph to a GraphML file.

#### **Parameters**

f: the name of the file to be written or a Python file handle

# write leda(f, names="name", weights="weights")

Writes the graph to a file in LEDA native format.

The LEDA format supports at most one attribute per vertex and edge. You can specify which vertex and edge attribute you want to use. Note that the name of the attribute is not saved in the LEDA file.

## Parameters

f: the name of the file to be written or a Python file handle

names: the name of the vertex attribute to be stored along with

the vertices. It is usually used to store the vertex names (hence the name of the keyword argument), but you may also use a numeric attribute. If you don't want to store

any vertex attributes, supply None here.

weights: the name of the edge attribute to be stored along with the

edges. It is usually used to store the edge weights (hence the name of the keyword argument), but you may also use a string attribute. If you don't want to store any edge

attributes, supply None here.

# write lgl(f, names="name", weights="weights", isolates=True)

Writes the edge list of a graph to a file in .lgl format.

Note that multiple edges and/or loops break the LGL software, but igraph does not check for this condition. Unless you know that the graph does not have multiple edges and/or loops, it is wise to call simplify() before saving.

## **Parameters**

f: the name of the file to be written or a Python file handle

names: the name of the vertex attribute containing the name of

the vertices. If you don't want to store vertex names,

supply None here.

weights: the name of the edge attribute containing the weight of

the vertices. If you don't want to store weights, supply

None here.

isolates: whether to include isolated vertices in the output.

# write ncol(f, names="name", weights="weights")

Writes the edge list of a graph to a file in .ncol format.

Note that multiple edges and/or loops break the LGL software, but igraph does not check for this condition. Unless you know that the graph does not have multiple edges and/or loops, it is wise to call simplify() before saving.

## **Parameters**

f: the name of the file to be written or a Python file handle

names: the name of the vertex attribute containing the name of

the vertices. If you don't want to store vertex names,

supply None here.

weights: the name of the edge attribute containing the weight of

the vertices. If you don't want to store weights, supply

None here.

# write pajek(f)

Writes the graph in Pajek format to the given file.

## **Parameters**

f: the name of the file to be written or a Python file handle

# Inherited from object

delattr	$(), \_\_format\_\_($	),getattribute	(),reduce	$(), \_\_reduce\_ex\_\_(),$
$_{-}$ repr $_{-}(),$	setattr(),	sizeof(),su	bclasshook()	

## 1.12.2 Properties

Name		
Inherited from object		
class		

## 1.12.3 Class Variables

Name	Description
hash	Value: None

# 2 Module igraph. igraph

Low-level Python interface for the igraph library. Should not be used directly.

#### 2.1 Functions

# convex hull(vs, coords=False)

Calculates the convex hull of a given point set.

## **Parameters**

vs: the point set as a list of lists

coords: if True, the function returns the coordinates of the corners

of the convex hull polygon, otherwise returns the corner

indices.

#### Return Value

either the hull's corner coordinates or the point indices corresponding to them, depending on the coords parameter.

# is degree sequence(out deg, in deg=None)

Returns whether a list of degrees can be a degree sequence of some graph.

Note that it is not required for the graph to be simple; in other words, this function may return True for degree sequences that can be realized using one or more multiple or loop edges only.

In particular, this function checks whether

- all the degrees are non-negative
- for undirected graphs, the sum of degrees are even
- for directed graphs, the two degree sequences are of the same length and equal sums

## **Parameters**

out\_deg: the list of degrees. For directed graphs, this list must

contain the out-degrees of the vertices.

in\_deg: the list of in-degrees for directed graphs. This parameter

must be None for undirected graphs.

## Return Value

True if there exists some graph that can realize the given degree sequence, False otherwise.@see: is\_graphical\_degree\_sequence() if you do not want to allow multiple or loop edges.

# is graphical degree sequence(out\_deg, in\_deg=None)

Returns whether a list of degrees can be a degree sequence of some simple graph.

Note that it is required for the graph to be simple; in other words, this function will return False for degree sequences that cannot be realized without using one or more multiple or loop edges.

## **Parameters**

out\_deg: the list of degrees. For directed graphs, this list must

contain the out-degrees of the vertices.

in\_deg: the list of in-degrees for directed graphs. This parameter

must be None for undirected graphs.

## Return Value

True if there exists some simple graph that can realize the given degree sequence, False otherwise.

See Also: is\_degree\_sequence() if you want to allow multiple or loop edges.

# set progress handler(handler)

Sets the handler to be called when igraph is performing a long operation.

#### **Parameters**

handler: the progress handler function. It must accept two

arguments, the first is the message informing the user about what igraph is doing right now, the second is the

actual progress information (a percentage).

# set random number generator(generator)

Sets the random number generator used by igraph.

#### Parameters

generator: the generator to be used. It must be a Python object

with at least three attributes: random, randint and gauss. Each of them must be callable and their signature and behaviour must be identical to random.random, random.randint and random.gauss. By default, igraph uses the random module for random number generation, but you can supply your alternative implementation here. If the given generator is None, igraph reverts to the default Mersenne twister generator implemented in the C layer, which might be slightly faster than calling back to Python for random numbers,

but you cannot set its seed or save its state.

# set status handler(handler)

Sets the handler to be called when igraph tries to display a status message.

This is used to communicate the progress of some calculations where no reasonable progress percentage can be given (so it is not possible to use the progress handler).

# **Parameters**

handler: the status handler function. It must accept a single

argument, the message that informs the user about what

igraph is doing right now.

## 2.2 Variables

Name	Description
ADJ_DIRECTED	Value: 0
ADJ_LOWER	Value: 3
ADJ_MAX	Value: 1
ADJ_MIN	Value: 4
ADJ_PLUS	Value: 5
ADJ_UNDIRECTED	Value: 1
ADJ_UPPER	Value: 2
ALL	Value: 3
BLISS_F	Value: 0
BLISS_FL	Value: 1
BLISS_FLM	Value: 4
BLISS_FM	Value: 3
BLISS_FS	Value: 2
BLISS_FSM	Value: 5
GET_ADJACENCY_BO-	Value: 2
TH	
GET_ADJACENCY_LO-	Value: 1
WER	
GET_ADJACENCY_UP-	Value: 0
PER	
IN	Value: 2
OUT	Value: 1
REWIRING_SIMPLE	Value: 0
REWIRING_SIMPLE_L-	Value: 1
OOPS	
STAR_IN	Value: 1
STAR_MUTUAL	Value: 3
STAR_OUT	Value: 0
STAR_UNDIRECTED	Value: 2

continued on next page

Name	Description
STRONG	Value: 2
TRANSITIVITY_NAN	Value: 0
TRANSITIVITY_ZERO	Value: 1
TREE_IN	Value: 1
TREE_OUT	Value: 0
TREE_UNDIRECTED	Value: 2
WEAK	Value: 1
build_date	Value: 'Oct 8 2020'
igraph_version	Value: '0.8.3'
package	Value: None
arpack_options	Value: <igraph.arpackoptions 0x10fde9b90="" at="" object=""></igraph.arpackoptions>

# 2.3 Class InternalError

object —		
exceptions.BaseException —		
exceptions.Exception		
	igraph.	igraph. Internal Error

## 2.3.1 Methods

 $Inherited\ from\ exceptions. Exception$ 

 $Inherited\ from\ exceptions. Base Exception$ 

del	$\operatorname{lattr}_{\_\_}$	_(),	getattr	$\operatorname{ribute}_{-}()$	,g	$etitem_{-}()$	),g	${ m etslice}_{\_}$	(), _	re-
duce_	_(), _	_repr_	_(), _	_setattr_	_(), _	_setstate_	_(), _	_str_	_(), _	_uni-
code	()									

Inherited from object

format	()	, hash	().	reduce ex	()	, sizeof	().	, subclasshook	()

# 2.3.2 Properties

Name	Description
Inherited from exceptions. Bo	iseException
args, message	
Inherited from object	

 $continued\ on\ next\ page$ 

Name	Description
class	

Variables Package igraph.app

# 3 Package igraph.app

User interfaces of igraph

# 3.1 Modules

• shell: Command-line user interface of igraph (Section 4, p. 232)

# 3.2 Variables

Name	Description
package	Value: None

# 4 Module igraph.app.shell

Command-line user interface of igraph

The command-line interface launches a Python shell with the igraph module automatically imported into the main namespace. This is mostly a convenience module and it is used only by the **igraph** command line script which executes a suitable Python shell and automatically imports **igraph**'s classes and functions in the top-level namespace.

Supported Python shells are:

- IDLE shell (class IDLEShell)
- IPython shell (class IPythonShell)
- Classic Python shell (class ClassicPythonShell)

The shells are tried in the above mentioned preference order one by one, unless the global.shells configuration key is set which overrides the default order. IDLE shell is only tried in Windows unless explicitly stated by global.shells, since Linux and Mac OS X users are likely to invoke igraph from the command line.

Version: 0.8.3

# 4.1 Functions

main()
The main entry point for igraph when invoked from the command line shell

# 4.2 Variables

Name	Description
package	Value: 'igraph.app'

## 4.3 Class TerminalController

A class that can be used to portably generate formatted output to a terminal.

'TerminalController' defines a set of instance variables whose values are initialized to the control sequence necessary to perform a given action. These can be simply included in normal output to the terminal:

```
>>> term = TerminalController()
>>> print 'This is '+term.GREEN+'green'+term.NORMAL
This is green
```

Alternatively, the 'render()' method can used, which replaces '\${action}' with the string required to perform 'action':

```
>>> term = TerminalController()
>>> print term.render('This is ${GREEN}green${NORMAL}')
This is green
```

If the terminal doesn't support a given action, then the value of the corresponding instance variable will be set to ". As a result, the above code will still work on terminals that do not support color, except that their output will not be colored. Also, this means that you can test whether the terminal supports a given action by simply testing the truth value of the corresponding instance variable:

```
>>> term = TerminalController()
>>> if term.CLEAR_SCREEN:
... print 'This terminal supports clearning the screen.'
```

Finally, if the width and height of the terminal are known, then they will be stored in the 'COLS' and 'LINES' attributes.

Author: Edward Loper

## 4.3.1 Methods

```
__init__(self, term_stream=sys.stdout)
```

Create a 'TerminalController' and initialize its attributes with appropriate values for the current terminal. 'term\_stream' is the stream that will be used for terminal output; if this stream is not a tty, then the terminal is assumed to be a dumb terminal (i.e., have no capabilities).

## render(self, template)

Replace each \$-substitutions in the given template string with the corresponding terminal control string (if it's defined) or " (if it's not).

#### 4.3.2 Class Variables

Name	Description
BOL	Move the cursor to the beginning of the line
	Value: ''
UP	Move the cursor up one line
	Value: ''
DOWN	Move the cursor down one line
	Value: ''
LEFT	Move the cursor left one char
	Value: ''
RIGHT	Move the cursor right one char
	Value: ''

continued on next page

Name	Description
CLEAR SCREEN	Clear the screen and move to home position
_	Value: ','
CLEAR EOL	Clear to the end of the line.
_	Value: ','
CLEAR_BOL	Clear to the beginning of the line.
_	Value: ','
CLEAR_EOS	Clear to the end of the screen
_	Value: ','
BOLD	Turn on bold mode
	Value: ','
BLINK	Turn on blink mode
	Value: ''
DIM	Turn on half-bright mode
	Value: ''
REVERSE	Turn on reverse-video mode
	Value: ''
NORMAL	Turn off all modes
	Value: ''
HIDE_CURSOR	Make the cursor invisible
	Value: ''
SHOW_CURSOR	Make the cursor visible
	Value: ''
COLS	Width of the terminal (None for unknown)
	Value: None
LINES	Height of the terminal (None for unknown)
	Value: None
WHITE	Value: ''
YELLOW	Value: ''
MAGENTA	Value: ''
RED	Value: ''
CYAN	Value: ','
GREEN	Value: ''
BLUE	Value: ''
BLACK	Value: ''
BG_CYAN	Value: ''
BG_GREEN	Value: ''
BG_BLUE	Value: ''
BG_BLACK	Value: ''
BG_WHITE	Value: ''
BG_YELLOW	Value: ''
BG_MAGENTA	Value: ''
BG_RED	Value: ''

# 4.4 Class ProgressBar

A 2-line progress bar, which looks like:

The progress bar is colored, if the terminal supports color output; and adjusts to the width of the terminal.

## 4.4.1 Methods

# update(self, percent=None, message=None)

Updates the progress bar.

# **Parameters**

percent: the percentage to be shown. If None, the previous value

will be used.

message: the message to be shown above the progress bar. If None,

the previous message will be used.

# update message(self, message)

Updates the message of the progress bar.

# **Parameters**

message: the message to be shown above the progress bar

$\mathbf{clear}(\mathit{self})$	
Clears the progress bar (i.e. removes it from the screen)	

## 4.4.2 Class Variables

Name	Description	
BAR	Value: '%3d%%	
	\${GREEN}[\${BOLD}%s%s\${NORMAL}\${GREEN}]\${N	ORMAL}'
HEADER	Value: '\${BOLD}\${CYAN}%s\${NORMAL}\n'	

#### 4.4.3 Instance Variables

Name	Description
cleared	true if we haven't drawn the bar yet.

# 4.5 Class Shell

Known Subclasses: igraph.app.shell.ClassicPythonShell, igraph.app.shell.IDLEShell, igraph.app.shell.IP Superclass of the embeddable shells supported by igraph

## 4.5.1 Methods

```
__init__(self)

x.__init__(...) initializes x; see help(type(x)) for signature

Overrides: object.__init__ extit(inherited documentation)
```

# supports progress bar(self)

Checks whether the shell supports progress bars.

This is done by checking for the existence of an attribute called \_progress\_handler.

# $supports\_status\_messages(self)$

Checks whether the shell supports status messages.

This is done by checking for the existence of an attribute called \_status\_handler.

```
{\tt get\_progress\_handler}(\mathit{self})
```

Returns the progress handler (if exists) or None (if not).

```
\frac{\mathbf{get\_status\_handler}(\mathit{self})}{\mathbf{Returns}\ \mathsf{the}\ \mathsf{status}\ \mathsf{handler}\ (\mathsf{if}\ \mathsf{exists})\ \mathsf{or}\ \mathsf{None}\ (\mathsf{if}\ \mathsf{not}).}
```

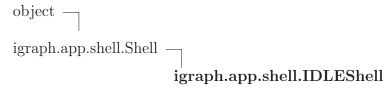
# Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

## 4.5.2 Properties

Name	Description
Inherited from object	
class	

## 4.6 Class IDLEShell



IDLE embedded shell interface.

This class allows igraph to be embedded in IDLE (the Tk Python IDE).

**To Do:** no progress bar support yet. Shell/Restart Shell command should re-import igraph again.

## 4.6.1 Methods

$-\operatorname{init}_{-}(self)$
Constructor.
Imports IDLE's embedded shell. The implementation of this method is ripped from idlelib.PyShell.main() after removing the unnecessary parts.
Overrides: objectinit

$\_\_{call}\_\_(self)$
Starts the shell
Overrides: igraph.app.shell.Shellcall

# $Inherited\ from\ igraph.app.shell.Shell (Section\ 4.5)$

 ${\tt get\_progress\_handler(), get\_status\_handler(), supports\_progress\_bar(), supports\_status\_message} \\ Inherited\ from\ object$ 

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

## 4.6.2 Properties

Name	Description
Inherited from object	
class	

# 4.7 Class ConsoleProgressBarMixin

object — igraph.app.shell.ConsoleProgressBarMixin

Known Subclasses: igraph.app.shell.ClassicPythonShell, igraph.app.shell.IPythonShell Mixin class for console shells that support a progress bar.

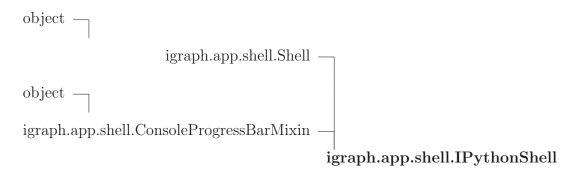
## 4.7.1 Methods

# Inherited from object

## 4.7.2 Properties

Name	Description
Inherited from object	
class	

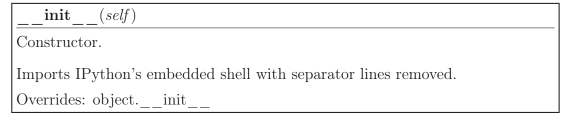
# 4.8 Class IPythonShell



IPython embedded shell interface.

This class allows igraph to be embedded in IPython's interactive shell.

## 4.8.1 Methods



```
__call__(self)
Starts the embedded shell.
Overrides: igraph.app.shell.Shell.__call__
```

# Inherited from igraph.app.shell.Shell(Section 4.5)

```
get\_progress\_handler(), get\_status\_handler(), supports\_progress\_bar(), supports\_status\_message(), supports\_status\_status\_message(), supports\_status\_status\_message(), supports\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_status\_
```

# Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

# 4.8.2 Properties

Name	Description
Inherited from object	
class	

# 4.9 Class ClassicPythonShell

```
object —

igraph.app.shell.Shell —

object —

igraph.app.shell.ConsoleProgressBarMixin —
```

igraph.app.shell.ClassicPythonShell

Classic Python shell interface.

This class allows igraph to be embedded in Python's shell.

## 4.9.1 Methods

$\{init}_{\_(self)}$
Constructor.
Imports Python's classic shell
Overrides: objectinit

```
__call__(self)
Starts the embedded shell.
Overrides: igraph.app.shell.Shell.__call__
```

# Inherited from igraph.app.shell.Shell(Section 4.5)

 ${\tt get\_progress\_handler(), get\_status\_handler(), supports\_progress\_bar(), supports\_status\_message} \\ Inherited\ from\ object$ 

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

## 4.9.2 Properties

Name	Description
Inherited from object	
class	

# 5 Module igraph.clustering

Classes related to graph clustering.

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## 5.1 Functions

compare\_communities(comm1, comm2, method='vi',
remove none=False)

Compares two community structures using various distance measures.

## **Parameters**

comm1: the first community structure as a membership list or

as a Clustering object.

comm2: the second community structure as a membership list

or as a Clustering object.

method: the measure to use. "vi" or "meila" means the

variation of information metric of Meila (2003),
"nmi" or "danon" means the normalized mutual
information as defined by Danon et al (2005),
"split-join" means the split-join distance of van
Dongen (2000), "rand" means the Rand index of
Rand (1971), "adjusted\_rand" means the adjusted

Rand index of Hubert and Arabie (1985).

remove\_none: whether to remove None entries from the membership

lists. This is handy if your Clustering object was

constructed using

VertexClustering.FromAttribute using an attribute which was not defined for all the vertices. If remove\_none is False, a None entry in either comm1 or comm2 will result in an exception. If remove\_none is True, None values are filtered away and only the

remaining lists are compared.

## Return Value

the calculated measure.

# Reference:

- Meila M: Comparing clusterings by the variation of information. In: Scholkopf B, Warmuth MK (eds). Learning Theory and Kernel Machines: 16th Annual Conference on Computational Learning Theory and 7th Kernel Workship, COLT/Kernel 2003, Washington, DC, USA. Lecture Notes in Computer Science, vol. 2777, Springer, 2003. ISBN: 978-3-540-40720-1.
- Danon L, Diaz-Guilera A, Duch J, Arenas A: Comparing community structure identification. J Stat Mech P09008, 2005.
- van Dongen D: Performance criteria for graph clustering and Markov cluster experiments. Technical Report INS-R0012, National Research Institute for Mathematics and Computer Science in the Netherlands, Amsterdam, May 2000.
- Rand WM: Objective criteria for the evaluation of clustering methods. J Am Stat Assoc 66(336):846-850, 1971.
- Hubert L and Arabie P: Comparing partitions. Journal of Classification 2:193-218, 1985.

# split join distance(comm1, comm2, remove none=False)

Calculates the split-join distance between two community structures.

The split-join distance is a distance measure defined on the space of partitions of a given set. It is the sum of the projection distance of one partition from the other and vice versa, where the projection number of A from B is if calculated as follows:

- 1. For each set in A, find the set in B with which it has the maximal overlap, and take note of the size of the overlap.
- 2. Take the sum of the maximal overlap sizes for each set in A.
- 3. Subtract the sum from n, the number of elements in the partition.

Note that the projection distance is asymmetric, that's why it has to be calculated in both directions and then added together. This function returns the projection distance of comm1 from comm2 and the projection distance of comm2 from comm1, and returns them in a pair. The actual split-join distance is the sum of the two distances. The reason why it is presented this way is that one of the elements being zero then implies that one of the partitions is a subpartition of the other (and if it is close to zero, then one of the partitions is close to being a subpartition of the other).

## **Parameters**

comm1: the first community structure as a membership list or

as a Clustering object.

comm2: the second community structure as a membership list

or as a Clustering object.

remove\_none: whether to remove None entries from the membership

lists. This is handy if your Clustering object was

constructed using

 ${\tt VertexClustering.FromAttribute} \ using \ an$ 

attribute which was not defined for all the vertices. If remove\_none is False, a None entry in either comm1 or comm2 will result in an exception. If remove\_none is True, None values are filtered away and only the

remaining lists are compared.

# Return Value

the projection distance of comm1 from comm2 and vice versa in a tuple. The split-join distance is the sum of the two.

**Reference:** van Dongen D: Performance criteria for graph clustering and Markov cluster experiments. Technical Report INS-R0012, National Research Institute for Mathematics and Computer Science in the Netherlands, Amsterdam, May 2000.

See Also: compare\_communities() with method = "split-join" if you are not interested in the individual projection distances but only the sum of them.

# 5.2 Variables

Name	Description
package	Value: 'igraph'

# 5.3 Class Clustering

```
object — igraph.clustering.Clustering
```

Known Subclasses: igraph.clustering.VertexClustering

Class representing a clustering of an arbitrary ordered set.

This is now used as a base for VertexClustering, but it might be useful for other purposes as well.

Members of an individual cluster can be accessed by the [] operator:

```
>>> cl = Clustering([0,0,0,0,1,1,1,2,2,2,2])
>>> cl[0]
[0, 1, 2, 3]
```

The membership vector can be accessed by the membership property:

```
>>> cl.membership
[0, 0, 0, 0, 1, 1, 1, 2, 2, 2, 2]
```

The number of clusters can be retrieved by the len function:

```
>>> len(cl)
```

You can iterate over the clustering object as if it were a regular list of clusters:

If you need all the clusters at once as lists, you can simply convert the clustering object to a list:

```
>>> cluster_list = list(cl)
>>> print cluster_list
[[0, 1, 2, 3], [4, 5, 6], [7, 8, 9, 10]]
```

## 5.3.1 Methods

init (self, membership, params=None)

Constructor.

# **Parameters**

membership: the membership list - that is, the cluster index in

which each element of the set belongs to.

params: additional parameters to be stored in this object's

dictionary.

Overrides: object.\_\_init\_\_

 $\_\_$ getitem $\_\_$ (self, idx)

Returns the members of the specified cluster.

# **Parameters**

idx: the index of the cluster

# Return Value

the members of the specified cluster as a list

# Raises

IndexError if the index is out of bounds

 $\_\_iter\_\_(\mathit{self})$ 

Iterates over the clusters in this clustering.

This method will return a generator that generates the clusters one by one.

 $\_\_len\_\_(\mathit{self})$ 

Returns the number of clusters.

Return Value

the number of clusters

 $\_\_\mathbf{str}\_\_(\mathit{self})$ 

str(x)

Overrides: object.\_\_str\_\_ extit(inherited documentation)

as cover(self)

Returns a Cover that contains the same clusters as this clustering.

# **compare to**(self, other, \*args, \*\*kwds)

Compares this clustering to another one using some similarity or distance metric.

This is a convenience method that simply calls compare\_communities with the two clusterings as arguments. Any extra positional or keyword argument is also forwarded to compare\_communities.

# size(self, idx)

Returns the size of a given cluster.

### Parameters

idx: the cluster in which we are interested.

# sizes(self, \*args)

Returns the size of given clusters.

The indices are given as positional arguments. If there are no positional arguments, the function will return the sizes of all clusters.

# size histogram(self, bin width=1)

Returns the histogram of cluster sizes.

#### **Parameters**

bin\_width: the bin width of the histogram

#### Return Value

a Histogram object

# summary(self, verbosity=0, width=None)

Returns the summary of the clustering.

The summary includes the number of items and clusters, and also the list of members for each of the clusters if the verbosity is nonzero.

## **Parameters**

verbosity: determines whether the cluster members should be printed. Zero verbosity prints the number of items and clusters only.

#### Return Value

the summary of the clustering as a string.

# Inherited from object

 $\_\_delattr\_\_(), \_\_format\_\_(), \_\_getattribute\_\_(), \_\_hash\_\_(), \_\_new\_\_(),$ 

#### 5.3.2 Properties

Name	Description
membership	Returns the membership vector.
n	Returns the number of elements covered by this
	clustering.
Inherited from object	
class	

# 5.4 Class VertexClustering

Known Subclasses: igraph.cut.Cut

The clustering of the vertex set of a graph.

This class extends Clustering by linking it to a specific Graph object and by optionally storing the modularity score of the clustering. It also provides some handy methods like getting the subgraph corresponding to a cluster and such.

Note: since this class is linked to a Graph, destroying the graph by the del operator does not free the memory occupied by the graph if there exists a VertexClustering that references the Graph.

### 5.4.1 Methods

 $\underline{-init}\_(self, graph, membership = \texttt{None}, modularity = \texttt{None}, params = \texttt{None}, modularity\_params = \texttt{None})$ 

Creates a clustering object for a given graph.

**Parameters** 

graph: the graph that will be associated to the

clustering

membership: the membership list. The length of the list

must be equal to the number of vertices in the graph. If None, every vertex is assumed to

belong to the same cluster.

modularity: the modularity score of the clustering. If

None, it will be calculated when needed.

params: additional parameters to be stored in this

object.

modularity\_params: arguments that should be passed to

Graph.modularity when the modularity is (re)calculated. If the original graph was weighted, you should pass a dictionary

containing a weight key with the appropriate

value here.

Overrides: object.\_\_init\_\_

FromAttribute(cls, graph, attribute, intervals=None, params=None)

Creates a vertex clustering based on the value of a vertex attribute.

Vertices having the same attribute will correspond to the same cluster.

## **Parameters**

graph: the graph on which we are working

attribute: name of the attribute on which the clustering is based.

intervals: for numeric attributes, you can either pass a single

number or a list of numbers here. A single number means that the vertices will be put in bins of that width and vertices ending up in the same bin will be in the same cluster. A list of numbers specify the bin positions explicitly; e.g., [10, 20, 30] means that there will be four categories: vertices with the attribute value less than 10, between 10 and 20, between 20 and 30 and over 30. Intervals are closed from the left and open from the right.

params: additional parameters to be stored in this object.

## Return Value

a new VertexClustering object

### as cover(self)

Returns a VertexCover that contains the same clusters as this clustering.

Overrides: igraph.clustering.Clustering.as cover

cluster graph(self, combine vertices=None, combine edges=None)

Returns a graph where each cluster is contracted into a single vertex.

In the resulting graph, vertex i represents cluster i in this clustering. Vertex i and j will be connected if there was at least one connected vertex pair (a, b) in the original graph such that vertex a was in cluster i and vertex b was in cluster j.

## Parameters

combine\_vertices: specifies how to derive the attributes of the

vertices in the new graph from the attributes of the old ones. See Graph.contract\_vertices()

for more details.

combine\_edges: specifies how to derive the attributes of the

edges in the new graph from the attributes of the old ones. See Graph.simplify() for more details. If you specify False here, edges will not be combined, and the number of edges between the vertices representing the original clusters will be equal to the number of edges between the members of those clusters in the

original graph.

## Return Value

the new graph.

### $\mathbf{crossing}(self)$

Returns a boolean vector where element i is True iff edge i lies between clusters, False otherwise.

# recalculate modularity(self)

Recalculates the stored modularity value.

This method must be called before querying the modularity score of the clustering through the class member modularity or q if the graph has been modified (edges have been added or removed) since the creation of the VertexClustering object.

#### Return Value

the new modularity score

### subgraph(self, idx)

Get the subgraph belonging to a given cluster.

#### **Parameters**

idx: the cluster index

#### Return Value

a copy of the subgraph

**Precondition:** the vertex set of the graph hasn't been modified since the moment the clustering was constructed.

## subgraphs(self)

Gets all the subgraphs belonging to each of the clusters.

### Return Value

a list containing copies of the subgraphs

**Precondition:** the vertex set of the graph hasn't been modified since the moment the clustering was constructed.

# $\mathbf{giant}(self)$

Returns the largest cluster of the clustered graph.

The largest cluster is a cluster for which no larger cluster exists in the clustering. It may also be known as the *giant community* if the clustering represents the result of a community detection function.

### Return Value

a copy of the largest cluster.

**Note:** there can be multiple largest clusters, this method will return the copy of an arbitrary one if there are multiple largest clusters.

**Precondition:** the vertex set of the graph hasn't been modified since the moment the clustering was constructed.

```
plot (self, context, bbox, palette, *args, **kwds)
```

Plots the clustering to the given Cairo context in the given bounding box.

This is done by calling Graph.\_\_plot\_\_() with the same arguments, but coloring the graph vertices according to the current clustering (unless overridden by the vertex\_color argument explicitly).

This method understands all the positional and keyword arguments that are understood by Graph.\_\_plot\_\_(), only the differences will be highlighted here:

- mark\_groups: whether to highlight some of the vertex groups by colored polygons. Besides the values accepted by Graph.\_\_plot\_\_ (i.e., a dict mapping colors to vertex indices, a list containing lists of vertex indices, or False), the following are also accepted:
  - True: all the groups will be highlighted, the colors matching the corresponding color indices from the current palette (see the palette keyword argument of Graph.\_\_plot\_\_.
  - A dict mapping cluster indices or tuples of vertex indices to color names. The given clusters or vertex groups will be highlighted by the given colors.
  - A list of cluster indices. This is equivalent to passing a dict mapping numeric color indices from the current palette to cluster indices; therefore, the cluster referred to by element i of the list will be highlighted by color i from the palette.

The value of the plotting.mark\_groups configuration key is also taken into account here; if that configuration key is True and mark\_groups is not given explicitly, it will automatically be set to True.

In place of lists of vertex indices, you may also use VertexSeq instances.

In place of color names, you may also use color indices into the current palette. None as a color name will mean that the corresponding group is ignored.

• palette: the palette used to resolve numeric color indices to RGBA values. By default, this is an instance of ClusterColoringPalette.

See Also:  ${\tt Graph.\_\_plot\_\_()}$  for more supported keyword arguments.

# $Inherited\ from\ igraph.clustering. Clustering (Section\ 5.3)$

```
\__getitem\_\_(), \ \__iter\_\_(), \ \__len\_\_(), \ \__str\_\_(), \ compare\_to(), \ size(), size\_histogram(), sizes(), summary()
```

# Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(),
```

#### 5.4.2 Properties

Name	Description		
modularity	Returns the modularity score		
q	Returns the modularity score		
graph	Returns the graph belonging to this object		
Inherited from igraph.clustering.Clustering (Section 5.3)			
membership, n			
Inherited from object			
class			

# 5.5 Class Dendrogram

Known Subclasses: igraph.clustering.VertexDendrogram

The hierarchical clustering (dendrogram) of some dataset.

A hierarchical clustering means that we know not only the way the elements are separated into groups, but also the exact history of how individual elements were joined into larger subgroups.

This class internally represents the hierarchy by a matrix with n rows and 2 columns – or more precisely, a list of lists of size 2. This is exactly the same as the original format used by igraph's C core. The *i*th row of the matrix contains the indices of the two clusters being joined in time step *i*. The joint group will be represented by the ID n+i, with *i* starting from one. The ID of the joint group will be referenced in the upcoming steps instead of any of its individual members. So, IDs less than or equal to n (where n is the number of rows in the matrix) mean the original members of the dataset (with ID from 0 to n), while IDs up from n+1 mean joint groups. As an example, take a look at the dendrogram and the internal representation of a given clustering of five nodes:

```
4 -+---
```

#### 5.5.1 Methods

```
__init__(self, merges)
Creates a hierarchical clustering.
```

## **Parameters**

merges: the merge history either in matrix or tuple format Overrides: object.\_\_init\_\_

```
__str__(self)
str(x)
Overrides: object.__str__ extit(inherited documentation)
```

```
format(self, format='newick')
Formats the dendrogram in a foreign format.

Currently only the Newick format is supported.

Example:

>>> d = Dendrogram([(2, 3), (0, 1), (4, 5)])
>>> d.format()
'((2,3)4,(0,1)5)6;'
>>> d.names = list("ABCDEFG")
>>> d.format()
'((C,D)E,(A,B)F)G;'
```

**summary**(self, verbosity=0, max leaf count=40)

Returns the summary of the dendrogram.

The summary includes the number of leafs and branches, and also an ASCII art representation of the dendrogram unless it is too large.

#### **Parameters**

verbosity: determines whether the ASCII representation of

the dendrogram should be printed. Zero verbosity

prints only the number of leafs and branches.

max\_leaf\_count: the maximal number of leafs to print in the

ASCII representation. If the dendrogram has

more leafs than this limit, the ASCII

representation will not be printed even if the

verbosity is larger than or equal to 1.

## Return Value

the summary of the dendrogram as a string.

```
plot (self, context, bbox, palette, *args, **kwds)
```

Draws the dendrogram on the given Cairo context

Supported keyword arguments are:

• orientation: the orientation of the dendrogram. Must be one of the following values: left-right, bottom-top, right-left or top-bottom. Individual elements are always placed at the former edge and merges are performed towards the latter edge. Possible aliases: horizontal = left-right, vertical = bottom-top, lr = left-right, rl = right-left, tb = top-bottom, bt = bottom-top. The default is left-right.

### Inherited from object

$_{-}$ delattr $_{-}$ (),	$\_\_format\_\_($	),ge	tattribu	$te\_\_()$ ,	hash	(), _	new	_(),
reduce(),	reduce_ex_	(),	_repr	_(),	$setattr\_\_$	_(),	sizeof	$_{-}(),$
subclasshook	()							

#### 5.5.2 Properties

Name	Description
merges	Returns the performed merges in matrix format
names	Returns the names of the nodes in the
	dendrogram
Inherited from object	
class	

continued on next page

	Name	Description
--	------	-------------

# 5.6 Class VertexDendrogram



The dendrogram resulting from the hierarchical clustering of the vertex set of a graph.

## 5.6.1 Methods

init(self, graph, m modularity_params=None)	erges, optimal_count=None, params=None,
Creates a dendrogram obje	ect for a given graph.
Parameters	
graph:	the graph that will be associated to the clustering
merges:	the merges performed given in matrix form.
optimal_count:	the optimal number of clusters where the dendrogram should be cut. This is a hint usually provided by the clustering algorithm that produces the dendrogram. None means that such a hint is not available; the optimal count will then be selected based on the modularity in such a case.
params:	additional parameters to be stored in this object.
modularity_params:	arguments that should be passed to Graph.modularity when the modularity is (re)calculated. If the original graph was weighted, you should pass a dictionary containing a weight key with the appropriate value here.
Overrides: objectinit_	_

# as clustering(self, n=None)

Cuts the dendrogram at the given level and returns a corresponding VertexClustering object.

#### **Parameters**

n: the desired number of clusters. Merges are replayed from the beginning until the membership vector has exactly n distinct elements or until there are no more recorded merges, whichever happens first. If None, the optimal count hint given by the clustering algorithm will be used If the optimal count was not given either, it will be calculated by selecting the level where the modularity is maximal.

### Return Value

a new VertexClustering object.

plot(self, context, bbox, palette, *args, **kwds)			
Draws the vertex dendrogram on the given Cairo context			
See Dendrogramplot for the list of supported keyword arguments.			
Overrides: igraph.clustering.Dendrogramplot			

# $Inherited\ from\ igraph.clustering.Dendrogram (Section\ 5.5)$

$_{-}$ str $_{-}(),$	format(),	summary()	
----------------------	-----------	-----------	--

# $Inherited\ from\ object$

delattr	_(), _	$\_$ format $\_$	(),	ge	tattribı	ite	(),has	h(),	new_	(),
reduce	_(),	_reduce_	_ex	_(),	_repr_	_(), _	$_{\rm setattr}$	(),	_sizeof_	_(),
$_{\rm \_subclassh}$	nook	_()								

### 5.6.2 Properties

Name	Description			
optimal_count	Returns the optimal number of clusters for this			
	dendrogram.			
If an optimal count hint was given at				
construction time, this property simply return				
the hint. If such a count was not given, this				
method calculates the optimal number of				
clusters by maximizing the modularity along				
the possible cuts in the dendrogram.				
Inherited from igraph.clustering.Dendrogram (Section 5.5)				
merges, names				
Inherited from object				

continued on next page

Name	Description
class	

### 5.7 Class Cover

```
object — igraph.clustering.Cover
```

Known Subclasses: igraph.clustering.VertexCover

Class representing a cover of an arbitrary ordered set.

Covers are similar to clusterings, but each element of the set may belong to more than one cluster in a cover, and elements not belonging to any cluster are also allowed.

Cover instances provide a similar API as Clustering instances; for instance, iterating over a Cover will iterate over the clusters just like with a regular Clustering instance. However, they are not derived from each other or from a common superclass, and there might be functions that exist only in one of them or the other.

Clusters of an individual cover can be accessed by the [] operator:

```
>>> cl = Cover([[0,1,2,3], [2,3,4], [0,1,6]])
>>> cl[0]
[0, 1, 2, 3]
```

The membership vector can be accessed by the membership property. Note that contrary to Clustering instances, the membership vector will contain lists that contain the cluster indices each item belongs to:

```
>>> cl.membership
[[0, 2], [0, 2], [0, 1], [0, 1], [1], [], [2]]
```

The number of clusters can be retrieved by the len function:

```
>>> len(cl)
```

You can iterate over the cover as if it were a regular list of clusters:

```
>>> for cluster in cl:
...     print " ".join(str(idx) for idx in cluster)
...
0 1 2 3
2 3 4
0 1 6
```

If you need all the clusters at once as lists, you can simply convert the cover to a list:

```
>>> cluster_list = list(cl)
>>> print cluster_list
[[0, 1, 2, 3], [2, 3, 4], [0, 1, 6]]
```

Clustering objects can readily be converted to Cover objects using the constructor:

```
>>> clustering = Clustering([0, 0, 0, 0, 1, 1, 1, 2, 2, 2])
>>> cover = Cover(clustering)
>>> list(clustering) == list(cover)
True
```

#### 5.7.1 Methods

init (self, clusters, n=0)

Constructs a cover with the given clusters.

#### **Parameters**

clusters: the clusters in this cover, as a list or iterable. Each cluster is specified by a list or tuple that contains the IDs of the items in this cluster. IDs start from zero.

n: the total number of elements in the set that is covered by this cover. If it is less than the number of unique elements found in all the clusters, we will simply use the number of unique elements, so it is safe to leave this at zero. You only have to specify this parameter if there are some elements that are covered by none of the clusters.

Overrides: object.\_\_init\_\_

```
\_\_getitem\_\_(self, index)
```

Returns the cluster with the given index.

```
<u>__iter__(self)</u>
Iterates over the clusters in this cover.
```

```
<u>__len__(self)</u>
Returns the number of clusters in this cover.
```

```
__str__(self)
Returns a string representation of the cover.
Overrides: object.__str__
```

size(self, idx)

Returns the size of a given cluster.

**Parameters** 

idx: the cluster in which we are interested.

sizes(self, \*args)

Returns the size of given clusters.

The indices are given as positional arguments. If there are no positional arguments, the function will return the sizes of all clusters.

size histogram(self, bin\_width=1)

Returns the histogram of cluster sizes.

**Parameters** 

bin\_width: the bin width of the histogram

Return Value

a Histogram object

summary(self, verbosity=0, width=None)

Returns the summary of the cover.

The summary includes the number of items and clusters, and also the list of members for each of the clusters if the verbosity is nonzero.

**Parameters** 

verbosity: determines whether the cluster members should be printed. Zero verbosity prints the number of items and clusters only.

Return Value

the summary of the cover as a string.

# Inherited from object

$\_$ _delattr $\_$ _(), $\_$	$\_{ m format}\_$	(),ge	etattribu	te(), _	hash_	(), _	new_	()
reduce(),	$_{ m reduce}_{ m }$	ex(), _	_repr	_(),se	tattr	(),	sizeof_	_(),
subclasshook	_()							

## 5.7.2 Properties

Name	Description
membership	Returns the membership vector of this cover.
	The membership vector of a cover covering $n$
	elements is a list of length $n$ , where element $i$
	contains the cluster indices of the $i$ th item.
n	Returns the number of elements in the set
	covered by this cover.
Inherited from object	
class	

# 5.8 Class VertexCover

Known Subclasses: igraph.clustering.CohesiveBlocks

The cover of the vertex set of a graph.

This class extends Cover by linking it to a specific Graph object. It also provides some handy methods like getting the subgraph corresponding to a cluster and such.

**Note:** since this class is linked to a **Graph**, destroying the graph by the **del** operator does not free the memory occupied by the graph if there exists a **VertexCover** that references the **Graph**.

#### 5.8.1 Methods

\_\_\_init\_\_\_(self, graph, clusters=None)
Creates a cover object for a given graph.

Parameters
graph: the graph that will be associated to the cover
clusters: the list of clusters. If None, it is assumed that there is
only a single cluster that covers the whole graph.

Overrides: object.\_\_init\_\_

## $\mathbf{crossing}(self)$

Returns a boolean vector where element i is True iff edge i lies between clusters, False otherwise.

# subgraph(self, idx)

Get the subgraph belonging to a given cluster.

## **Parameters**

idx: the cluster index

## Return Value

a copy of the subgraph

**Precondition:** the vertex set of the graph hasn't been modified since the moment the cover was constructed.

# subgraphs(self)

Gets all the subgraphs belonging to each of the clusters.

## Return Value

a list containing copies of the subgraphs

**Precondition:** the vertex set of the graph hasn't been modified since the moment the cover was constructed.

**plot** (self, context, bbox, palette, \*args, \*\*kwds)

Plots the cover to the given Cairo context in the given bounding box.

This is done by calling Graph.\_\_plot\_\_() with the same arguments, but drawing nice colored blobs around the vertex groups.

This method understands all the positional and keyword arguments that are understood by <code>Graph.\_\_plot\_\_()</code>, only the differences will be highlighted here:

- mark\_groups: whether to highlight the vertex clusters by colored polygons. Besides the values accepted by Graph.\_\_plot\_\_ (i.e., a dict mapping colors to vertex indices, a list containing lists of vertex indices, or False), the following are also accepted:
  - True: all the clusters will be highlighted, the colors matching the corresponding color indices from the current palette (see the palette keyword argument of Graph.\_\_plot\_\_.
  - A dict mapping cluster indices or tuples of vertex indices to color names. The given clusters or vertex groups will be highlighted by the given colors.
  - A list of cluster indices. This is equivalent to passing a dict mapping numeric color indices from the current palette to cluster indices; therefore, the cluster referred to by element i of the list will be highlighted by color i from the palette.

The value of the plotting.mark\_groups configuration key is also taken into account here; if that configuration key is True and mark\_groups is not given explicitly, it will automatically be set to True.

In place of lists of vertex indices, you may also use VertexSeq instances. In place of color names, you may also use color indices into the current palette. None as a color name will mean that the corresponding group is ignored.

• palette: the palette used to resolve numeric color indices to RGBA values. By default, this is an instance of ClusterColoringPalette.

See Also: Graph.\_\_plot\_\_() for more supported keyword arguments.

# Inherited from igraph.clustering.Cover(Section 5.7)

 $\__getitem\_\_(), \__iter\_\_(), \__len\_\_(), \__str\_\_(), size(), size\_histogram(), sizes(), summary()$ 

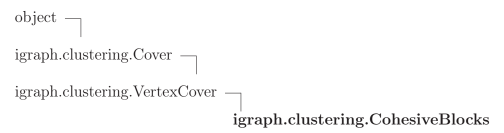
### Inherited from object

```
\_\_subclasshook\_\_()
```

## 5.8.2 Properties

Name	Description			
graph	Returns the graph belonging to this object			
Inherited from igraph.clustering.Cover (Section 5.7)				
membership, n				
Inherited from object				
class				

### 5.9 Class CohesiveBlocks



The cohesive block structure of a graph.

Instances of this type are created by Graph.cohesive\_blocks(). See the documentation of Graph.cohesive\_blocks() for an explanation of what cohesive blocks are.

This class provides a few more methods that make handling of cohesive block structures easier.

#### 5.9.1 Methods

init (self, graph, blocks=None, cohesion=None, parent=None)

Constructs a new cohesive block structure for the given graph.

If any of blocks, cohesion or parent is None, all the arguments will be ignored and Graph.cohesive\_blocks() will be called to calculate the cohesive blocks. Otherwise, these three variables should describe the \*result\* of a cohesive block structure calculation. Chances are that you never have to construct CohesiveBlocks instances directly, just use Graph.cohesive\_blocks().

#### **Parameters**

graph: the graph itself

blocks: a list containing the blocks; each block is described as a

list containing vertex IDs.

cohesion: the cohesion of each block. The length of this list must

be equal to the length of blocks.

parent: the parent block of each block. Negative values or None

mean that there is no parent block for that block. There should be only one parent block, which covers the entire

graph.

Overrides: object. init

**See Also:** Graph.cohesive\_blocks()

#### cohesion(self, idx)

Returns the cohesion of the group with the given index.

## cohesions(self)

Returns the list of cohesion values for each group.

#### hierarchy(self)

Returns a new graph that describes the hierarchical relationships between the groups.

The new graph will be a directed tree; an edge will point from vertex i to vertex j if group i is a superset of group j. In other words, the edges point downwards.

### $\max$ cohesion(self, idx)

Finds the maximum cohesion score among all the groups that contain the given vertex.

# $\max\_cohesions(self)$

For each vertex in the graph, returns the maximum cohesion score among all the groups that contain the vertex.

# parent(self, idx)

Returns the parent group index of the group with the given index or None if the given group is the root.

# parents(self)

Returns the list of parent group indices for each group or None if the given group is the root.

# \_\_plot\_\_(self, context, bbox, palette, \*args, \*\*kwds)

Plots the cohesive block structure to the given Cairo context in the given bounding box.

Since a CohesiveBlocks instance is also a VertexCover, keyword arguments accepted by VertexCover.\_\_plot\_\_() are also accepted here. The only difference is that the vertices are colored according to their maximal cohesions by default, and groups are marked by colored blobs except the last group which encapsulates the whole graph.

See the documentation of VertexCover.\_\_plot\_\_() for more details.

Overrides: igraph.clustering.VertexCover.\_\_plot\_\_

# $Inherited\ from\ igraph.clustering. Vertex Cover (Section\ 5.8)$

 ${\rm crossing}(),\,{\rm subgraph}(),\,{\rm subgraphs}()$ 

# Inherited from igraph.clustering.Cover(Section 5.7)

 $\_getitem\_\_(), \__iter\_\_(), \__len\_\_(), \__str\_\_(), size(), size\_histogram(), sizes(), summary()$ 

# Inherited from object

delattr_	_(), _	$_{\_}$ format	(),	ge	tattribı	$ite\_\_($	(),l	nash	_(), _	_new_	_()
reduce	_(), _	reduce_	_ex	_(),	_repr_	_(), _	_setat	tr()	),	sizeof	_(),
subclassl	nook	()									

#### 5.9.2 Properties

Name	Description			
Inherited from igraph.cluster	ing. VertexCover (Section 5.8)			

continued on next page

Name Description				
graph				
Inherited from igraph.clustering.Cover (Section 5.7)				
membership, n				
Inherited from object				
class				

# 6 Module igraph.configuration

Configuration framework for igraph.

igraph has some parameters which usually affect the behaviour of many functions. This module provides the framework for altering and querying igraph parameters as well as saving them to and retrieving them from disk.

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#### 6.1 Functions

# get platform image viewer()

Returns the path of an image viewer on the given platform

### get user config file()

Returns the path where the user-level configuration file is stored

### init()

Default mechanism to initiate igraph configuration

This method loads the user-specific configuration file from the user's home directory, or if it does not exist, creates a default configuration.

The method is safe to be called multiple times, it will not parse the configuration file twice.

### Return Value

the Configuration object loaded or created.

### 6.2 Variables

Name	Description
package	Value: 'igraph'

# 6.3 Class Configuration

```
object | | igraph.configuration.Configuration
```

Class representing igraph configuration details.

```
(section) General ideas
```

The configuration of igraph is stored in the form of name-value pairs. This object provides an interface to the configuration data using the syntax known from dict:

```
>>> c=Configuration()
>>> c["general.verbose"] = True
>>> print C["general.verbose"]
True
```

Configuration keys are organized into sections, and the name to be used for a given key is always in the form section.keyname, like general.verbose in the example above. In that case, general is the name of the configuration section, and verbose is the name of the key. If the name of the section is omitted, it defaults to general, so general.verbose can be referred to as verbose:

```
>>> c=Configuration()
>>> c["verbose"] = True
>>> print c["general.verbose"]
True
```

User-level configuration is stored in ~/.igraphrc per default on Linux and Mac OS X systems, or in C:\Documents and Settings\username\.igraphrc on Windows systems. However, this configuration is read only when igraph is launched through its shell interface defined in igraph.app.shell. This behaviour might change before version 1.0.

```
(section) Known configuration keys
```

The known configuration keys are presented below, sorted by section. When referring to them in program code, don't forget to add the section name, expect in the case of section general.

```
(section) General settings
```

These settings are all stored in section general.

• shells: the list of preferred Python shells to be used with the command-line igraph

script. The shells in the list are tried one by one until any of them is found on the system. igraph functions are then imported into the main namespace of the shell and the shell is launched. Known shells and their respective class names to be used can be found in igraph.app.shell. Example: IPythonShell, ClassicPythonShell. This is the default, by the way.

• **verbose**: whether **igraph** should talk more than really necessary. For instance, if set to **True**, some functions display progress bars.

# (section) Application settings

These settings specify the external applications that are possibly used by igraph. They are all stored in section apps.

• image\_viewer: image viewer application. If set to an empty string, it will be determined automatically from the platform igraph runs on. On Mac OS X, it defaults to the Preview application. On Linux, it chooses a viewer from several well-known Linux viewers like gthumb, kuickview and so on (see the source code for the full list). On Windows, it defaults to the system's built-in image viewer.

# (section) Plotting settings

These settings specify the default values used by plotting functions. They are all stored in section plotting.

- layout: default graph layout algorithm to be used.
- mark\_groups: whether to mark the clusters by polygons when plotting a clustering object.
- palette: default palette to be used for converting integer numbers to colors. See colors.Palette for more information. Valid palette names are stored in colors.palettes.
- wrap\_labels: whether to try to wrap the labels of the vertices automatically if they don't fit within the vertex. Default: False.

### (section) Shell settings

These settings specify options for external environments in which igraph is embedded (e.g., IPython and its Qt console). These settings are stored in section shell.

• ipython.inlining.Plot: whether to show instances of the Plot class inline in IPython's console if the console supports it. Default: True

### 6.3.1 Methods

 $_{
m init}_{
m \_}(\mathit{self}, \mathit{filename} = {\tt None})$ 

Creates a new configuration instance.

**Parameters** 

filename: file or file pointer to be read. Can be omitted.

Overrides: object.\_\_init\_\_

\_\_contains\_\_(self, item)

Checks whether the given configuration item is set.

**Parameters** 

item: the configuration key to check.

Return Value

True if the key has an associated value, False otherwise.

getitem (self, item)

Returns the given configuration item.

**Parameters** 

item: the configuration key to retrieve.

Return Value

the configuration value

 $\_\_$ setitem $\_\_$ (self, item, value)

Sets the given configuration item.

**Parameters** 

item: the configuration key to set

value: the new value of the configuration key

 $\mathbf{delitem} \qquad (\mathit{self}, \mathit{item})$ 

Deletes the given item from the configuration.

If the item has a default value, the default value is written back instead of the current value. Without a default value, the item is really deleted.

has key(self, item)

Checks if the configuration has a given key.

**Parameters** 

item: the key being sought

load(self, stream=None)

Loads the configuration from the given file.

**Parameters** 

stream: name of a file or a file object. The configuration will be loaded from here. Can be omitted, in this case, the

user-level configuration is loaded.

save(self, stream = None)

Saves the configuration.

**Parameters** 

stream: name of a file or a file object. The configuration will be

saved there. Can be omitted, in this case, the user-level

configuration file will be overwritten.

instance(cls)

Returns the single instance of the configuration object.

# Inherited from object

delattr	r(), _	$\_$ format $\_$	_(),	getattrib	$ute\($	(),has	h(),	new_	()
reduce	e(), _	_reduce_e	ex(),	repr_	(), _	_setattr_	(),	_sizeof_	_(),
$\operatorname{str}$	(), su	ıbclasshook	()						

### 6.3.2 Properties

Name	Description
filename	Returns the filename associated to the object.
	It is usually the name of the configuration file
	that was used when creating the object.
	Configuration.load always overwrites it with
	the filename given to it. If None, the
	configuration was either created from scratch or
	it was updated from a stream without name
	information.
Inherited from object	
class	

Class Cut Module igraph.cut

# 7 Module igraph.cut

Classes representing cuts and flows on graphs.

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#### 7.1 Variables

Name	Description
package	Value: 'igraph'

#### 7.2 Class Cut

object — igraph.clustering.Clustering — igraph.clustering.VertexClustering — igraph.cut.Cut

Known Subclasses: igraph.cut.Flow

A cut of a given graph.

This is a simple class used to represent cuts returned by Graph.mincut(), Graph.all\_st\_cuts() and other functions that calculate cuts.

A cut is a special vertex clustering with only two clusters. Besides the usual VertexClustering methods, it also has the following attributes:

- value the value (capacity) of the cut. It is equal to the number of edges if there are no capacities on the edges.
- partition vertex IDs in the parts created after removing edges in the cut
- cut edge IDs in the cut

Class Cut Module igraph.cut

• es - an edge selector restricted to the edges in the cut.

You can use indexing on this object to obtain lists of vertex IDs for both sides of the partition.

This class is usually not instantiated directly, everything is taken care of by the functions that return cuts.

# Examples:

```
>>> from igraph import Graph
>>> g = Graph.Ring(20)
>>> mc = g.mincut()
>>> print mc.value
2.0
>>> print min(map(len, mc))
1
>>> mc.es["color"] = "red"
```

Class Cut Module igraph.cut

#### 7.2.1 Methods

 $\underline{-nit}\_\_(self, \ graph, \ value = \texttt{None}, \ cut = \texttt{None}, \ partition = \texttt{None}, \\ partition = \underbrace{-\texttt{None}}$ 

Initializes the cut.

This should not be called directly, everything is taken care of by the functions that return cuts.

#### **Parameters**

graph: the graph that will be associated to the

clustering

membership: the membership list. The length of the list

must be equal to the number of vertices in the graph. If None, every vertex is assumed to

belong to the same cluster.

modularity: the modularity score of the clustering. If

None, it will be calculated when needed.

params: additional parameters to be stored in this

object.

modularity\_params: arguments that should be passed to

Graph.modularity when the modularity is (re)calculated. If the original graph was weighted, you should pass a dictionary

containing a weight key with the appropriate

value here.

Overrides: object. init

```
__repr__(self)
repr(x)
Overrides: object.__repr__ extit(inherited documentation)
```

```
-_str__(self)
str(x)
Overrides: object.__str__ extit(inherited documentation)
```

# Inherited from igraph.clustering.VertexClustering(Section 5.4)

FromAttribute(), \_\_plot\_\_(), as\_cover(), cluster\_graph(), crossing(), giant(), recalculate\_modularity(), subgraph(), subgraphs()

# $Inherited\ from\ igraph.clustering.Clustering(Section\ 5.3)$

$$\__getitem\_\_(), \__iter\_\_(), \__len\_\_(), compare\_to(), size(), size\_histogram(), sizes(), summary()$$

# Inherited from object

```
\label{lem:condition} $$\_\_delattr\_(), \_\_format\_\_(), \_\_getattribute\_\_(), \_\_hash\_\_(), \_\_new\_\_(), \_\_reduce\_ex\_\_(), \_\_setattr\_\_(), \_\_sizeof\_\_(), \_\_subclasshook\_\_()
```

## 7.2.2 Properties

Name	Description			
es	Returns an edge selector restricted to the cut			
partition	Returns the vertex IDs partitioned according to			
	the cut			
cut	Returns the edge IDs in the cut			
value	Returns the sum of edge capacities in the cut			
Inherited from igraph.clustering.VertexClustering (Section 5.4)				
graph, modularity, q				
Inherited from igraph.cluster	ring. Clustering (Section 5.3)			
membership, n				
Inherited from object				
class				

### 7.3 Class Flow

object —	
igraph.clustering.Clustering —	
igraph.clustering.VertexClustering —	
igraph.cut.Cut	
	igraph.cut.Flow

A flow of a given graph.

This is a simple class used to represent flows returned by Graph.maxflow. It has the following attributes:

- graph the graph on which this flow is defined
- value the value (capacity) of the flow
- flow the flow values on each edge. For directed graphs, this is simply a list where element i corresponds to the flow on edge i. For undirected graphs, the direction of

the flow is not constrained (since the edges are undirected), hence positive flow always means a flow from the smaller vertex ID to the larger, while negative flow means a flow from the larger vertex ID to the smaller.

- cut edge IDs in the minimal cut corresponding to the flow.
- partition vertex IDs in the parts created after removing edges in the cut
- es an edge selector restricted to the edges in the cut.

This class is usually not instantiated directly, everything is taken care of by Graph.maxflow.

### Examples:

```
>>> from igraph import Graph
>>> g = Graph.Ring(20)
>>> mf = g.maxflow(0, 10)
>>> print mf.value
2.0
>>> mf.es["color"] = "red"
```

#### 7.3.1 Methods

```
(self, graph, value, flow, cut, partition)
    init
Initializes the flow.
This should not be called directly, everything is taken care of by
Graph.maxflow.
Parameters
    graph:
                           the graph that will be associated to the
                           clustering
                           the membership list. The length of the list
    membership:
                           must be equal to the number of vertices in the
                           graph. If None, every vertex is assumed to
                           belong to the same cluster.
    modularity:
                           the modularity score of the clustering. If
                           None, it will be calculated when needed.
    params:
                           additional parameters to be stored in this
                           object.
    modularity_params: arguments that should be passed to
                           Graph.modularity when the modularity is
                           (re)calculated. If the original graph was
                           weighted, you should pass a dictionary
                           containing a weight key with the appropriate
                           value here.
Overrides: object. init
```

```
repr__(self)
repr(x)
Overrides: object.__repr__ extit(inherited documentation)
```

```
str__(self)
str(x)
Overrides: object.__str__ extit(inherited documentation)
```

# $Inherited\ from\ igraph.clustering. VertexClustering (Section\ 5.4)$

FromAttribute(), \_\_plot\_\_(), as\_cover(), cluster\_graph(), crossing(), giant(), recalculate modularity(), subgraph(), subgraphs()

Inherited from igraph.clustering.Clustering(Section 5.3)

 $\__getitem\_\_(), \__iter\_\_(), \__len\_\_(), compare\_to(), size(), size\_histogram(), sizes(), summary()$ 

# Inherited from object

# 7.3.2 Properties

Name	Description			
flow	Returns the flow values for each edge.			
	For directed graphs, this is simply a list where			
	element $i$ corresponds to the flow on edge $i$ . For			
	undirected graphs, the direction of the flow is			
	not constrained (since the edges are			
	undirected), hence positive flow always means a			
	flow from the smaller vertex ID to the larger,			
	while negative flow means a flow from the			
	larger vertex ID to the smaller.			
Inherited from igraph.cut.Cut (Section 7.2)				
cut, es, partition, value				
Inherited from igraph.clustering.VertexClustering (Section 5.4)				
graph, modularity, q				
Inherited from igraph.clustering.Clustering (Section 5.3)				
membership, n				
Inherited from object				
class				

# 8 Module igraph.datatypes

Additional auxiliary data types

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#### 8.1 Variables

Name	Description
package	Value: 'igraph'

## 8.2 Class Matrix

object —

igraph.datatypes.Matrix

Simple matrix data type.

Of course there are much more advanced matrix data types for Python (for instance, the ndarray data type of Numeric Python) and this implementation does not want to compete with them. The only role of this data type is to provide a convenient interface for the matrices returned by the Graph object (for instance, allow indexing with tuples in the case of adjacency matrices and so on).

#### 8.2.1 Methods

\_\_\_init\_\_\_(self, data=None)
Initializes a matrix.

Parameters
 data: the elements of the matrix as a list of lists, or None to create a 0x0 matrix.

Overrides: object.\_\_init\_\_\_

# Fill(cls, value, \*args)

Creates a matrix filled with the given value

### **Parameters**

value: the value to be used

shape: the shape of the matrix. Can be a single integer, two

integers or a tuple. If a single integer is given here, the

matrix is assumed to be square-shaped.

# $\overline{\mathbf{Zero}(cls, *args)}$

Creates a matrix filled with zeros.

#### **Parameters**

shape: the shape of the matrix. Can be a single integer, two integers or a tuple. If a single integer is given here, the matrix is assumed to be square-shaped.

Identity(cls, \*args)

Creates an identity matrix.

# **Parameters**

shape: the shape of the matrix. Can be a single integer, two integers or a tuple. If a single integer is given here, the matrix is assumed to be square-shaped.

# add (self, other)

Adds the given value to the matrix.

### **Parameters**

other: either a scalar or a matrix. Scalars will be added to each element of the matrix. Matrices will be added together elementwise.

#### Return Value

the result matrix

 $\_$  eq $\_$  (self, other)

Checks whether a given matrix is equal to another one



Returns a single item, a row or a column of the matrix

#### **Parameters**

i: if a single integer, returns the *i*th row as a list. If a slice, returns the corresponding rows as another Matrix object. If a 2-tuple, the first element of the tuple is used to select a row and the second is used to select a column.

```
\_\_hash\_\_(self)
```

Returns a hash value for a matrix.

Overrides: object.\_\_hash\_\_

```
\_\_iadd\_\_(self, other)
```

In-place addition of a matrix or scalar.

```
isub (self, other)
```

In-place subtraction of a matrix or scalar.

```
ne (self, other)
```

Checks whether a given matrix is not equal to another one

```
\_\_setitem\_\_(self, i, value)
```

Sets a single item, a row or a column of the matrix

#### **Parameters**

i: if a single integer, sets the *i*th row as a list. If a slice, sets the corresponding rows from another Matrix object. If a 2-tuple, the first element of the tuple is used to select a row and the second is used to select a column.

value: the new value

```
\_sub\_\_(self, other)
```

Subtracts the given value from the matrix.

#### **Parameters**

other: either a scalar or a matrix. Scalars will be subtracted from each element of the matrix. Matrices will be subtracted together elementwise.

## Return Value

the result matrix

```
__repr__(self)
repr(x)
Overrides: object.__repr__ extit(inherited documentation)
```

```
__str__(self)
str(x)
Overrides: object.__str__ extit(inherited documentation)
```

iter (self)

Support for iteration.

This is actually implemented as a generator, so there is no need for a separate iterator class. The generator returns *copies* of the rows in the matrix as lists to avoid messing around with the internals. Feel free to do anything with the copies, the changes won't be reflected in the original matrix.

**plot** (self, context, bbox, palette, \*\*kwds)

Plots the matrix to the given Cairo context in the given box

Besides the usual self-explanatory plotting parameters (context, bbox, palette), it accepts the following keyword arguments:

- style: the style of the plot. boolean is useful for plotting matrices with boolean (True/False or 0/1) values: False will be shown with a white box and True with a black box. palette uses the given palette to represent numbers by colors, the minimum will be assigned to palette color index 0 and the maximum will be assigned to the length of the palette. None draws transparent cell backgrounds only. The default style is boolean (but it may change in the future). None values in the matrix are treated specially in both cases: nothing is drawn in the cell corresponding to None.
- square: whether the cells of the matrix should be square or not. Default is True.
- grid\_width: line width of the grid shown on the matrix. If zero or negative, the grid is turned off. The grid is also turned off if the size of a cell is less than three times the given line width. Default is 1. Fractional widths are also allowed.
- border\_width: line width of the border drawn around the matrix. If zero or negative, the border is turned off. Default is 1.
- row\_names: the names of the rows
- col\_names: the names of the columns.
- values: values to be displayed in the cells. If None or False, no values are displayed. If True, the values come from the matrix being plotted. If it is another matrix, the values of that matrix are shown in the cells. In this case, the shape of the value matrix must match the shape of the matrix being plotted.
- value\_format: a format string or a callable that specifies how the values should be plotted. If it is a callable, it must be a function that expects a single value and returns a string. Example: "%#.2f" for floating-point numbers with always exactly two digits after the decimal point. See the Python documentation of the % operator for details on the format string. If the format string is not given, it defaults to the str function.

If only the row names or the column names are given and the matrix is square-shaped, the same names are used for both column and row names.

### $\min(self, dim = None)$

Returns the minimum of the matrix along the given dimension

## **Parameters**

dim: the dimension. 0 means determining the column minimums, 1 means determining the row minimums. If None, the global minimum is returned.

## $\max(self, dim=None)$

Returns the maximum of the matrix along the given dimension

#### **Parameters**

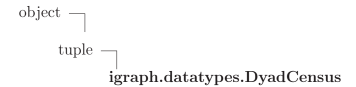
dim: the dimension. 0 means determining the column maximums, 1 means determining the row maximums. If None, the global maximum is returned.

## Inherited from object

#### 8.2.2 Properties

Name	Description
data	Returns the data stored in the matrix as a list
	of lists
shape	Returns the shape of the matrix as a tuple
Inherited from object	
class	

## 8.3 Class DyadCensus



Dyad census of a graph.

This is a pretty simple class - basically it is a tuple, but it allows the user to refer to its individual items by the names mutual (or mut), asymmetric (or asy or asym or asymm) and null.

#### Examples:

```
>>> from igraph import Graph
```

```
>>> g=Graph.Erdos_Renyi(100, 0.2, directed=True)
>>> dc=g.dyad_census()
>>> print dc.mutual
                               #doctest:+SKIP
>>> print dc["asym"]
                              #doctest:+SKIP
1609
>>> print tuple(dc), list(dc) #doctest:+SKIP
(179, 1609, 3162) [179, 1609, 3162]
>>> print sorted(dc.as_dict().items())
                                         #doctest:+ELLIPSIS
[('asymmetric', ...), ('mutual', ...), ('null', ...)]
8.3.1 Methods
     getitem (self, idx)
    x[y]
    Overrides: tuple.__getitem__ extit(inherited documentation)
     \_\_getattr\_\_(\overline{\textit{self, attr}})
    repr (self)
    repr(x)
    Overrides: object. repr extit(inherited documentation)
    \_\_{
m str}\_\_(\overline{self})
    str(x)
    Overrides: object.__str__ extit(inherited documentation)
    as dict(self)
    Converts the dyad census to a dict using the known dyad names.
Inherited from tuple
    __add__(), __contains__(), __eq__(), __ge__(), __getattribute__(), __get-
    newargs__(), __getslice__(), __gt__(), __hash__(), __iter__(), __le__(),
    __len__(), __lt__(), __mul__(), __ne__(), __new__(), __rmul__(),
    count(), index()
Inherited from object
```

### 8.3.2 Properties

Name	Description
Inherited from object	
class	

#### 8.4 Class TriadCensus

$$\begin{array}{c} \text{object} & \frown \\ \text{tuple} & \frown \\ \text{igraph.datatypes.TriadCensus} \end{array}$$

Triad census of a graph.

This is a pretty simple class - basically it is a tuple, but it allows the user to refer to its individual items by the following triad names:

- 003 the empty graph
- 012 a graph with a single directed edge (A -> B, C)
- 102 a graph with a single mutual edge (A <-> B, C)
- 021D the binary out-tree (A <- B -> C)
- 021U the binary in-tree (A -> B <- C)
- 021C the directed line (A -> B -> C)
- 111D A <-> B <- C
- 111U − A <-> B -> C
- $\bullet$  030T A -> B <- C, A -> C
- $\bullet$  030C A <- B <- C, A -> C
- 201 A <-> B <-> C
- $\bullet$  120D A <- B -> C, A <-> C
- 120U − A → B <- C, A <-> C
- $\bullet$  120C A -> B -> C, A <-> C
- $\bullet$  210C A -> B <-> C, A <-> C
- 300 the complete graph (A <-> B <-> C, A <-> C)

Attribute and item accessors are provided. Due to the syntax of Python, attribute names are not allowed to start with a number, therefore the triad names must be prepended with a lowercase t when accessing them as attributes. This is not necessary with the item accessor syntax.

#### Examples:

>>> from igraph import Graph

```
>>> g=Graph.Erdos_Renyi(100, 0.2, directed=True)
>>> tc=g.triad_census()
>>> print tc.t003 #doctest:+SKIP
39864
>>> print tc["030C"] #doctest:+SKIP
1206
```

#### 8.4.1 Methods

```
__getitem__(self, idx)
x[y]
Overrides: tuple.__getitem__ extit(inherited documentation)
```

```
__repr__(self)
repr(x)
Overrides: object.__repr__ extit(inherited documentation)
```

```
__str__(self)
str(x)
Overrides: object.__str__ extit(inherited documentation)
```

## Inherited from tuple

```
__add__(), __contains__(), __eq__(), __ge__(), __getattribute__(), __getnewargs__(), __getslice__(), __gt__(), __hash__(), __iter__(), __le__(), __len__(), __len__(), __new__(), __new__(), __rmul__(), count(), index()
```

# $Inherited\ from\ object$

#### 8.4.2 Properties

Name	Description
Inherited from object	
class	

## 8.5 Class UniqueIdGenerator

```
object | | igraph.datatypes.UniqueIdGenerator
```

A dictionary-like class that can be used to assign unique IDs to names (say, vertex names).

## Usage:

```
>>> gen = UniqueIdGenerator()
>>> gen["A"]
0
>>> gen["B"]
1
>>> gen["C"]
2
>>> gen["A"]  # Retrieving already existing ID
0
>>> gen.add("D")  # Synonym of gen["D"]
3
>>> len(gen)  # Number of already used IDs
4
>>> "C" in gen
True
>>> "E" in gen
False
```

#### 8.5.1 Methods

```
init (self, id\_generator = None, initial = None)
```

Creates a new unique ID generator. 'id\_generator' specifies how do we assign new IDs to elements that do not have an ID yet. If it is 'None', elements will be assigned integer identifiers starting from 0. If it is an integer, elements will be assigned identifiers starting from the given integer. If it is an iterator or generator, its 'next' method will be called every time a new ID is needed.

Overrides: object.\_\_init\_\_

```
__contains__(self, item)
Checks whether 'item' already has an ID or not.
```

```
getitem (self, item)
```

Retrieves the ID corresponding to 'item'. Generates a new ID for 'item' if it is the first time we request an ID for it.

setitem(self, item, value)	
Overrides the ID for 'item'.	

```
__len__(self)
"Returns the number of items
```

## reverse dict(self)

Returns the reverse mapping, i.e., the one that maps from generated IDs to their corresponding objects

## values(self)

Returns the values stored so far. If the generator generates items according to the standard sorting order, the values returned will be exactly in the order they were added. This holds for integer IDs for instance (but for many other ID generators as well).

# add(self, item)

Retrieves the ID corresponding to 'item'. Generates a new ID for 'item' if it is the first time we request an ID for it.

# $Inherited\ from\ object$

#### 8.5.2 Properties

Name	Description
Inherited from object	
class	

# 9 Package igraph.drawing

Drawing and plotting routines for IGraph.

Plotting is dependent on the pycairo or cairocffi libraries that provide Python bindings to the popular Cairo library<sup>1</sup>. This means that if you don't have pycairo<sup>2</sup> or cairocffi<sup>3</sup> installed, you won't be able to use the plotting capabilities. However, you can still use Graph.write\_svg to save the graph to an SVG file and view it from Mozilla Firefox<sup>4</sup> (free) or edit it in Inkscape<sup>5</sup> (free), Skencil<sup>6</sup> (formerly known as Sketch, also free) or Adobe Illustrator.

Whenever the documentation refers to the pycairo library, you can safely replace it with cairocffi as the two are API-compatible.

License: GPL

#### 9.1 Modules

- baseclasses: Abstract base classes for the drawing routines. (Section 10, p. 312)
- colors: Color handling functions. (Section 11, p. 315)
- **coord**: Coordinate systems and related plotting routines (Section 12, p. 328)
- edge: Drawers for various edge styles in graph plots. (Section 13, p. 331)
- graph: Drawing routines to draw graphs. (Section 14, p. 339)
- **metamagic**: Auxiliary classes for the default graph drawer in igraph. (Section 15, p. 346)
- **shapes**: Shape drawing classes for igraph (Section 16, p. 349)
- text: Drawers for labels on plots. (Section 17, p. 352)
- utils: Utility classes for drawing routines. (Section 18, p. 357)
- vertex: Drawing routines to draw the vertices of graphs. (Section 19, p. 368)

<sup>&</sup>lt;sup>1</sup>http://www.cairographics.org

<sup>&</sup>lt;sup>2</sup>http://www.cairographics.org/pycairo

<sup>&</sup>lt;sup>3</sup>http://cairocffi.readthedocs.io

<sup>&</sup>lt;sup>4</sup>http://www.mozilla.org/firefox

<sup>&</sup>lt;sup>5</sup>http://www.inkscape.org

 $<sup>^6 \</sup>mathrm{http://www.skencil.org}$ 

## 9.2 Functions

plot(obj, target=None, bbox=(0, 0, 600, 600), \*args, \*\*kwds)

Plots the given object to the given target.

Positional and keyword arguments not explicitly mentioned here will be passed down to the <code>\_\_plot\_\_</code> method of the object being plotted. Since you are most likely interested in the keyword arguments available for graph plots, see <code>Graph.\_\_plot\_\_</code> as well.

#### **Parameters**

obj: the object to be plotted

target: the target where the object should be plotted. It can be one of the following types:

- None an appropriate surface will be created and the object will be plotted there.
- cairo.Surface the given Cairo surface will be used. This can refer to a PNG image, an arbitrary window, an SVG file, anything that Cairo can handle.
- string a file with the given name will be created and an appropriate Cairo surface will be attached to it. The supported image formats are: PNG, PDF, SVG and PostScript.

bbox:

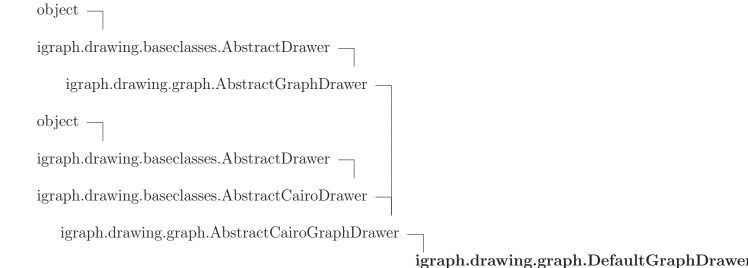
the bounding box of the plot. It must be a tuple with either two or four integers, or a BoundingBox object. If this is a tuple with two integers, it is interpreted as the width and height of the plot (in pixels for PNG images and on-screen plots, or in points for PDF, SVG and PostScript plots, where 72 pt = 1 inch = 2.54 cm). If this is a tuple with four integers, the first two denotes the X and Y coordinates of a corner and the latter two denoting the X and Y coordinates of the opposite corner.

opacity: the opacity of the object being plotted. It can be used to overlap several plots of the same graph if you use the same layout for them – for instance, you might plot a graph with opacity 0.5 and then plot its spanning tree over it with opacity 0.1. To achieve this, you'll need to modify the Plot object returned with Plot.add.

palette: the palette primarily used on the plot if the added objects do not specify a private palette. Must be either an igraph.drawing.colors.Palette object or a string referring to a valid key of igraph.drawing.colors.palettes (see module igraph.drawing.colors) or None. In the latter case, the default palette given by the configuration key plotting.palette is used.

margin: the top, right, bottom, left margins as a 4-tuple. If it has less than 4 elements or is a single float, the elements will be rejused until the length is at least 4. The default

## 9.3 Class DefaultGraphDrawer



Class implementing the default visualisation of a graph.

The default visualisation of a graph draws the nodes on a 2D plane according to a given Layout, then draws a straight or curved edge between nodes connected by edges. This is the visualisation used when one invokes the plot() function on a Graph object.

See Graph.\_\_plot\_\_() for the keyword arguments understood by this drawer.

#### 9.3.1 Methods

\_\_init\_\_\_(self, context, bbox, vertex\_drawer\_factory=<class
'igraph.drawing.vertex.DefaultVertexDrawer'>,
edge\_drawer\_factory=<class 'igraph.drawing.edge.ArrowEdgeDrawer'>,
label\_drawer\_factory=<class 'igraph.drawing.text.TextDrawer'>)

Constructs the graph drawer and associates it to the given Cairo context and the given BoundingBox.

#### Parameters

context: the context on which we will draw

bbox: the bounding box within which we will

draw. Can be anything accepted by the constructor of BoundingBox (i.e., a 2-tuple, a 4-tuple or a BoundingBox

object).

vertex\_drawer\_factory: a factory method that returns an

AbstractCairoVertexDrawer instance bound to a given Cairo context. The factory method must take three parameters: the Cairo context, the bounding box of the drawing area and the palette to be used for drawing colored vertices. The default vertex drawer is

DefaultVertexDrawer.

edge\_drawer\_factory: a factory method that returns an

AbstractEdgeDrawer instance bound to a given Cairo context. The factory method must take two parameters: the Cairo context and the palette to be used for drawing colored edges. You can use any of the actual AbstractEdgeDrawer implementations here to control the style of edges drawn by igraph. The default edge drawer is ArrowEdgeDrawer.

label\_drawer\_factory: a factory method that returns a

TextDrawer instance bound to a given Cairo context. The method must take one parameter: the Cairo context. The default label drawer is TextDrawer.

Overrides: object. init

draw(self, graph, palette, \*args, \*\*kwds)

Abstract method, must be implemented in derived classes.

Overrides: igraph.drawing.baseclasses.AbstractDrawer.draw extit(inherited documentation)

# $Inherited\ from\ igraph.drawing.graph.AbstractGraphDrawer$

```
ensure layout()
```

# $Inherited\ from\ object$

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

#### 9.3.2 Properties

Name	Description
Inherited from $igraph.drawing.baseclasses.AbstractCairoDrawer$ (Section 10.3)	
bbox	
Inherited from object	
class	

# 9.4 Class BoundingBox

object — igraph.drawing.utils.Rectangle —

# igraph. drawing. utils. Bounding Box

Class representing a bounding box (a rectangular area) that encloses some objects.

#### 9.4.1 Methods

```
__ior__(self, other)
Replaces this bounding box with the union of itself and another.

Example:

>>> box1 = BoundingBox(10, 20, 50, 60)
>>> box2 = BoundingBox(70, 40, 100, 90)
>>> box1 |= box2
>>> print(box1)
BoundingBox(10.0, 20.0, 100.0, 90.0)

Overrides: igraph.drawing.utils.Rectangle. ior
```

```
___or__(self, other)

Takes the union of this bounding box with another.

The result is a bounding box which encloses both bounding boxes.

Example:

>>> box1 = BoundingBox(10, 20, 50, 60)
>>> box2 = BoundingBox(70, 40, 100, 90)
>>> box1 | box2
BoundingBox(10.0, 20.0, 100.0, 90.0)

Overrides: igraph.drawing.utils.Rectangle.__or__
```

# Inherited from igraph.drawing.utils.Rectangle(Section 18.1)

## Inherited from object

```
\label{lem:condition} $$\_\_delattr\_(), \_\_format\__(), \_\_getattribute\__(), \_\_new\__(), \_\_reduce\__(), \_\_reduce\_ex\__(), \_\_setattr\_(), \_\_sizeof\__(), \_\_str\__(), \_\_subclasshook\__()
```

#### 9.4.2 Properties

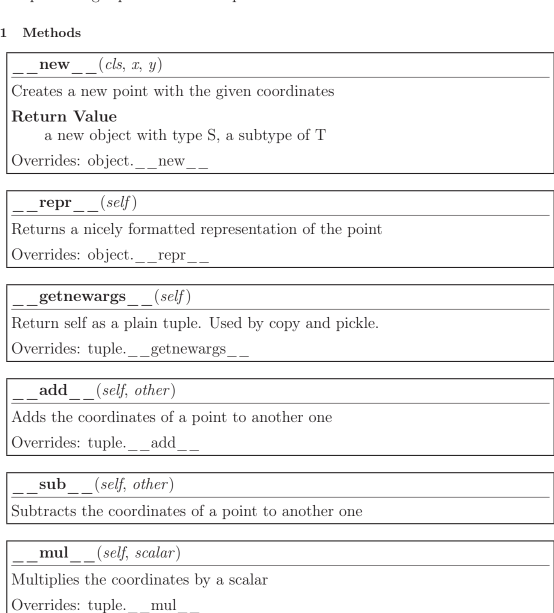
Name	Description
Inherited from igraph.drawing.utils.Rectangle (Section 18.1)	
bottom, coords, height, left, midx, midy, right, shape, top, width	
Inherited from object	
class	

#### 9.5 **Class Point**



Class representing a point on the 2D plane.

#### 9.5.1 Methods



```
\mathbf{rmul} (self, scalar)
```

Multiplies the coordinates by a scalar

Overrides: tuple.\_\_rmul\_\_

```
\_\_ div\_\_ (self, scalar)
```

Divides the coordinates by a scalar

## $as_polar(self)$

Returns the polar coordinate representation of the point.

#### Return Value

the radius and the angle in a tuple.

## **distance**(self, other)

Returns the distance of the point from another one.

# Example:

```
>>> p1 = Point(5, 7)
>>> p2 = Point(8, 3)
>>> p1.distance(p2)
5.0
```

# interpolate(self, other, ratio=0.5)

Linearly interpolates between the coordinates of this point and another one.

#### **Parameters**

other: the other point

ratio: the interpolation ratio between 0 and 1. Zero will return this

point, 1 will return the other point.

#### length(self)

Returns the length of the vector pointing from the origin to this point.

## normalized(self)

Normalizes the coordinates of the point s.t. its length will be 1 after normalization. Returns the normalized point.

## sq length(self)

Returns the squared length of the vector pointing from the origin to this point.

## towards(self, other, distance=0)

Returns the point that is at a given distance from this point towards another one.

# FromPolar(cls, radius, angle)

Constructs a point from polar coordinates.

'radius' is the distance of the point from the origin; 'angle' is the angle between the X axis and the vector pointing to the point from the origin.

## Inherited from tuple

# Inherited from object

#### 9.5.2 Properties

Name	Description
X	Alias for field number 0
У	Alias for field number 1
Inherited from object	
class	

# 9.6 Class Rectangle

object — igraph.drawing.utils.Rectangle

 ${\bf Known~Subclasses:~igraph.drawing.utils.BoundingBox}$ 

Class representing a rectangle.

#### 9.6.1 Methods

```
__init___(self, *args)
```

Creates a rectangle.

The corners of the rectangle can be specified by either a tuple (four items, two for each corner, respectively), four separate numbers (X and Y coordinates for each corner) or two separate numbers (width and height, the upper left corner is assumed to be at (0,0))

Overrides: object.\_\_init\_\_

## **contract**(self, margins)

Contracts the rectangle by the given margins.

#### Return Value

a new Rectangle object.

## expand(self, margins)

Expands the rectangle by the given margins.

#### Return Value

a new Rectangle object.

## **isdisjoint**(self, other)

Returns "True" if the two rectangles have no intersection.

#### Example:

```
>>> r1 = Rectangle(10, 10, 30, 30)
>>> r2 = Rectangle(20, 20, 50, 50)
>>> r3 = Rectangle(70, 70, 90, 90)
>>> r1.isdisjoint(r2)
False
>>> r2.isdisjoint(r1)
False
>>> r1.isdisjoint(r3)
True
>>> r3.isdisjoint(r1)
True
```

```
isempty(self)
Returns "True" if the rectangle is empty (i.e. it has zero width and height).
Example:
   >>> r1 = Rectangle(10, 10, 30, 30)
   >>> r2 = Rectangle(70, 70, 90, 90)
   >>> r1.isempty()
   False
   >>> r2.isempty()
   False
   >>> r1.intersection(r2).isempty()
```

## **intersection**(self, other)

Returns the intersection of this rectangle with another.

```
Example:
  >>> r1 = Rectangle(10, 10, 30, 30)
  >>> r2 = Rectangle(20, 20, 50, 50)
  >>> r3 = Rectangle(70, 70, 90, 90)
  >>> r1.intersection(r2)
  Rectangle(20.0, 20.0, 30.0, 30.0)
  >>> r2 & r1
  Rectangle(20.0, 20.0, 30.0, 30.0)
  >>> r2.intersection(r1) == r1.intersection(r2)
  True
  >>> r1.intersection(r3)
  Rectangle(0.0, 0.0, 0.0, 0.0)
```

```
__and__(self, other)

Returns the intersection of this rectangle with another.

Example:

>>> r1 = Rectangle(10, 10, 30, 30)
>>> r2 = Rectangle(20, 20, 50, 50)
>>> r3 = Rectangle(70, 70, 90, 90)
>>> r1.intersection(r2)
Rectangle(20.0, 20.0, 30.0, 30.0)
>>> r2 & r1
Rectangle(20.0, 20.0, 30.0, 30.0)
>>> r2.intersection(r1) == r1.intersection(r2)
True
>>> r1.intersection(r3)
Rectangle(0.0, 0.0, 0.0, 0.0)
```

```
translate(self, dx, dy)
Translates the rectangle in-place.

Example:
>>> r = Rectangle(10, 20, 50, 70)
>>> r.translate(30, -10)
>>> r
Rectangle(40.0, 10.0, 80.0, 60.0)

Parameters
    dx: the X coordinate of the translation vector
    dy: the Y coordinate of the translation vector
```

### **union**(*self*, *other*)

Returns the union of this rectangle with another.

The resulting rectangle is the smallest rectangle that contains both rectangles.

## Example:

```
>>> r1 = Rectangle(10, 10, 30, 30)
>>> r2 = Rectangle(20, 20, 50, 50)
>>> r3 = Rectangle(70, 70, 90, 90)
>>> r1.union(r2)
Rectangle(10.0, 10.0, 50.0, 50.0)
>>> r2 | r1
Rectangle(10.0, 10.0, 50.0, 50.0)
>>> r2.union(r1) == r1.union(r2)
True
>>> r1.union(r3)
Rectangle(10.0, 10.0, 90.0, 90.0)
```

## or (self, other)

Returns the union of this rectangle with another.

The resulting rectangle is the smallest rectangle that contains both rectangles.

## Example:

```
>>> r1 = Rectangle(10, 10, 30, 30)
>>> r2 = Rectangle(20, 20, 50, 50)
>>> r3 = Rectangle(70, 70, 90, 90)
>>> r1.union(r2)
Rectangle(10.0, 10.0, 50.0, 50.0)
>>> r2 | r1
Rectangle(10.0, 10.0, 50.0, 50.0)
>>> r2.union(r1) == r1.union(r2)
True
>>> r1.union(r3)
Rectangle(10.0, 10.0, 90.0, 90.0)
```

9.6.2 Properties

```
ior
                                                            (self, other)
                  Expands this rectangle to include itself and another completely while still
                  being as small as possible.
                  Example:
                                >>> r1 = Rectangle(10, 10, 30, 30)
                                >>> r2 = Rectangle(20, 20, 50, 50)
                                >>> r3 = Rectangle(70, 70, 90, 90)
                                >>> r1 |= r2
                                >>> r1
                               Rectangle(10.0, 10.0, 50.0, 50.0)
                                >>> r1 |= r3
                                >>> r1
                               Rectangle(10.0, 10.0, 90.0, 90.0)
                            \_\mathbf{repr}_- (self)
                  repr(x)
                  Overrides: object.__repr__ extit(inherited documentation)
                                                          (self, other)
                                  eq
                                                           (self, other)
                                  \mathbf{n}\mathbf{e}
                                                                   (self)
                                  bool
                                 nonzero\_\_(self)
                     \mathbf{hash} (self)
                  hash(x)
                  Overrides: object.__hash__ extit(inherited documentation)
Inherited from object
                 \label{eq:continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous
```

Name	Description
coords	The coordinates of the corners.
	The coordinates are returned as a 4-tuple in the
	following order: left edge, top edge, right edge,
	bottom edge.
width	The width of the rectangle
height	The height of the rectangle
left	The X coordinate of the left side of the box
right	The X coordinate of the right side of the box
top	The Y coordinate of the top edge of the box
bottom	The Y coordinate of the bottom edge of the box
midx	The X coordinate of the center of the box
midy	The Y coordinate of the center of the box
shape	The shape of the rectangle (width, height)
Inherited from object	
class	

## 9.7 Class Plot

object — igraph.drawing.Plot

Class representing an arbitrary plot

Every plot has an associated surface object where the plotting is done. The surface is an instance of cairo.Surface, a member of the pycairo library. The surface itself provides a unified API to various plotting targets like SVG files, X11 windows, PostScript files, PNG files and so on. igraph usually does not know on which surface it is plotting right now, since pycairo takes care of the actual drawing. Everything that's supported by pycairo should be supported by this class as well.

Current Cairo surfaces that I'm aware of are:

- cairo.GlitzSurface OpenGL accelerated surface for the X11 Window System.
- cairo. ImageSurface memory buffer surface. Can be written to a PNG image file.
- cairo.PDFSurface PDF document surface.
- cairo.PSSurface PostScript document surface.
- cairo.SVGSurface SVG (Scalable Vector Graphics) document surface.
- cairo.Win32Surface Microsoft Windows screen rendering.
- cairo.XlibSurface X11 Window System screen rendering.

If you create a Plot object with a string given as the target surface, the string will be treated as a filename, and its extension will decide which surface class will be used. Please note that

not all surfaces might be available, depending on your pycairo installation.

A Plot has an assigned default palette (see igraph.drawing.colors.Palette) which is used for plotting objects.

A Plot object also has a list of objects to be plotted with their respective bounding boxes, palettes and opacities. Palettes assigned to an object override the default palette of the plot. Objects can be added by the Plot.add method and removed by the Plot.remove method.

#### 9.7.1 Methods

(self, target = None, bbox = None, palette = None, background = None)init

Creates a new plot.

#### **Parameters**

target:

the target surface to write to. It can be one of the following types:

- None an appropriate surface will be created and the object will be plotted there.
- cairo.Surface the given Cairo surface will be used.
- string a file with the given name will be created and an appropriate Cairo surface will be attached to it.

bbox:

the bounding box of the surface. It is interpreted differently with different surfaces: PDF and PS surfaces will treat it as points (1 point = 1/72 inch). Image surfaces will treat it as pixels. SVG surfaces will treat it as an abstract unit, but it will mostly be interpreted as pixels when viewing the SVG file in Firefox.

palette:

the palette primarily used on the plot if the added objects do not specify a private palette. Must be either an igraph.drawing.colors.Palette object or a string referring to a valid key of igraph.drawing.colors.palettes (see module igraph.drawing.colors) or None. In the latter case, the default palette given by the configuration key plotting.palette is used.

background: the background color. If None, the background will be transparent. You can use any color specification here that is understood by igraph.drawing.colors.color\_name\_to\_rgba.

Overrides: object. init

add(self, obj, bbox=None, palette=None, opacity=1.0, \*args, \*\*kwds)

Adds an object to the plot.

Arguments not specified here are stored and passed to the object's plotting function when necessary. Since you are most likely interested in the arguments acceptable by graphs, see Graph.\_\_plot\_\_ for more details.

#### **Parameters**

obj: the object to be added

bbox: the bounding box of the object. If None, the object will

fill the entire area of the plot.

palette: the color palette used for drawing the object. If the object

tries to get a color assigned to a positive integer, it will use this palette. If None, defaults to the global palette of

the plot.

opacity: the opacity of the object being plotted, in the range

0.0 - 1.0

See Also: Graph.\_\_plot\_\_

## remove(self, obj, bbox=None, idx=1)

Removes an object from the plot.

If the object has been added multiple times and no bounding box was specified, it removes the instance which occurs idxth in the list of identical instances of the object.

#### **Parameters**

obj: the object to be removed

bbox: optional bounding box specification for the object. If given,

only objects with exactly this bounding box will be

considered.

idx: if multiple objects match the specification given by obj and

bbox, only the idxth occurrence will be removed.

#### Return Value

True if the object has been removed successfully, False if the object was not on the plot at all or idx was larger than the count of occurrences

## mark dirty(self)

Marks the plot as dirty (should be redrawn)

 $redraw(\mathit{self}, \mathit{context} = \mathtt{None})$ 

Redraws the plot

 $\mathbf{save}(\mathit{self}, \mathit{fname} = \mathtt{None})$ 

Saves the plot.

#### **Parameters**

fname: the filename to save to. It is ignored if the surface of the plot
 is not an ImageSurface.

 $\mathbf{show}(self)$ 

Saves the plot to a temporary file and shows it.

repr svg (self)

Returns an SVG representation of this plot as a string.

This method is used by IPython to display this plot inline.

## Inherited from object

## 9.7.2 Properties

Name	Description
background	Returns the background color of the plot. None
	means a transparent background.
bounding_box	Returns the bounding box of the Cairo surface
	as a BoundingBox object
height	Returns the height of the Cairo surface on
	which the plot is drawn
surface	Returns the Cairo surface on which the plot is
	drawn
width	Returns the width of the Cairo surface on
	which the plot is drawn
Inherited from object	
class	

# 10 Module igraph.drawing.baseclasses

Abstract base classes for the drawing routines.

#### 10.1 Variables

Name	Description
package	Value: 'igraph.drawing'

#### 10.2 Class AbstractDrawer

object — igraph.drawing.baseclasses.AbstractDrawer

Known Subclasses: igraph.drawing.graph.AbstractGraphDrawer, igraph.drawing.baseclasses.AbstractCaigraph.drawing.vertex.AbstractVertexDrawer, igraph.drawing.baseclasses.AbstractXMLRPCDrawer

Abstract class that serves as a base class for anything that draws an igraph object.

#### 10.2.1 Methods

$\mathbf{draw}(self, *args, **kwds)$
Abstract method, must be implemented in derived classes.

## Inherited from object

#### 10.2.2 Properties

Name	Description
Inherited from object	
class	

#### 10.3 Class AbstractCairoDrawer

igraph. drawing. base classes. Abstract Cairo Drawer

 $\textbf{Known Subclasses:} \ igraph.drawing.graph.AbstractCairoGraphDrawer, igraph.drawing.vertex.AbstractCairoGraphDrawer, igraph.drawer, igr$ 

igraph.drawing.text. Text Drawer, igraph.drawing.shapes. Polygon Drawer, igraph.drawing.coord. Coordinated the property of t

Abstract class that serves as a base class for anything that draws on a Cairo context within a given bounding box.

A subclass of AbstractCairoDrawer is guaranteed to have an attribute named context that represents the Cairo context to draw on, and an attribute named bbox for the BoundingBox of the drawing area.

#### 10.3.1 Methods

${f init}$	(self,	context,	bbox)	)
------------	--------	----------	-------	---

Constructs the drawer and associates it to the given Cairo context and the given BoundingBox.

#### **Parameters**

context: the context on which we will draw

bbox: the bounding box within which we will draw. Can be

anything accepted by the constructor of BoundingBox (i.e., a 2-tuple, a 4-tuple or a BoundingBox object).

Overrides: object.\_\_init\_\_

```
\mathbf{draw}(\mathit{self}, *\mathit{args}, **\mathit{kwds})
```

Abstract method, must be implemented in derived classes.

Overrides: igraph.drawing.baseclasses.AbstractDrawer.draw

## Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

#### 10.3.2 Properties

Name	Description
bbox	The bounding box of the drawing area where
	this drawer will draw.
Inherited from object	
class	

## 10.4 Class AbstractXMLRPCDrawer

object —	
igraph. drawing. base classes. Abstract Drawer	

# igraph. drawing. base classes. Abstract XMLRPCD raw

Known Subclasses: igraph.drawing.graph.CytoscapeGraphDrawer, igraph.drawing.graph.UbiGraphDraw

Abstract drawer that uses a remote service via XML-RPC to draw something on a remote display.

#### 10.4.1 Methods

init(se	elf, url, service=None)
Constructs an abstract drawer using the XML-RPC service at the given URL.	
Parameters	
url:	the URL where the XML-RPC calls for the service should be addressed to.
service: the name of the service at the XML-RPC address. If None, requests will be directed to the server proxy object constructed by xmlrpclib.ServerProxy; if not None, the given attribute will be looked up in the server proxy object.	
Overrides: objectinit	

# $Inherited\ from\ igraph.drawing.base classes. AbstractDrawer (Section\ 10.2)$

draw()

# Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

## 10.4.2 Properties

Name	Description
Inherited from object	
class	

# 11 Module igraph.drawing.colors

Color handling functions.

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#### 11.1 Functions

**color** name to **rgb**(color, palette=None)

Converts a color given in one of the supported color formats to R-G-B values.

This is done by calling color\_name\_to\_rgba and then throwing away the alpha value.

See Also: color\_name\_to\_rgba for more details about what formats are understood by this function.

## color name to rgba(color, palette=None)

Converts a color given in one of the supported color formats to R-G-B-A values.

### Examples:

```
>>> color_name_to_rgba("red")
(1.0, 0.0, 0.0, 1.0)
>>> color_name_to_rgba("#ff8000") == (1.0, 128/255.0, 0.0, 1.0)
>>> color_name_to_rgba("#ff800080") == (1.0, 128/255.0, 0.0, 128/255.0)
True
>>> color_name_to_rgba("#08f") == (0.0, 136/255.0, 1.0, 1.0)
True
>>> color_name_to_rgba("rgb(100%, 50%, 0%)")
(1.0, 0.5, 0.0, 1.0)
>>> color_name_to_rgba("rgba(100%, 50%, 0%, 25%)")
(1.0, 0.5, 0.0, 0.25)
>>> color_name_to_rgba("hsla(120, 100%, 50%, 0.5)")
(0.0, 1.0, 0.0, 0.5)
>>> color_name_to_rgba("hsl(60, 100%, 50%)")
(1.0, 1.0, 0.0, 1.0)
>>> color_name_to_rgba("hsv(60, 100%, 100%)")
(1.0, 1.0, 0.0, 1.0)
```

#### **Parameters**

color: the color to be converted in one of the following formats:

- CSS3 color specification: #rrggbb, #rgb, #rrggbbaa, #rgba, rgb(red, green, blue), rgba(red, green, blue, alpha), hsl(hue, saturation, lightness), hsla(hue, saturation, value) and hsva(hue, saturation, value, alpha) where the components are given as hexadecimal numbers in the first four cases and as decimals or percentages (0%-100%) in the remaining cases. Red, green and blue components are between 0 and 255; hue is between 0 and 360; saturation, lightness and value is between 0 and 100; alpha is between 0 and 1.
- Valid HTML color names, i.e. those that are present in the HTML 4.0 specification
- Valid X11 color names, see http://en.wikipedia.org/wiki/X11 color names
- Red-green-blue components given separately in either a comma-, slash- or whitespace-separated string or a list or a tuple, in the range of 0-255. An alpha value of 255 (maximal opacity) will be assumed.
- Red-green-blue-alpha components given separately in either a comma-, slash- or whitespace-separated string or a list or a tuple, in the

## hsla to rgba(h, s, l, alpha=1.0)

Converts a color given by its HSLA coordinates (hue, saturation, lightness, alpha) to RGBA coordinates.

Each of the HSLA coordinates must be in the range [0, 1].

## hsl to rgb(h, s, l)

Converts a color given by its HSL coordinates (hue, saturation, lightness) to RGB coordinates.

Each of the HSL coordinates must be in the range [0, 1].

# hsva to rgba(h, s, v, alpha=1.0)

Converts a color given by its HSVA coordinates (hue, saturation, value, alpha) to RGB coordinates.

Each of the HSVA coordinates must be in the range [0, 1].

## hsv to $\mathbf{rgb}(h, s, v)$

Converts a color given by its HSV coordinates (hue, saturation, value) to RGB coordinates.

Each of the HSV coordinates must be in the range [0, 1].

## rgba to hsla(r, g, b, alpha=1.0)

Converts a color given by its RGBA coordinates to HSLA coordinates (hue, saturation, lightness, alpha).

Each of the RGBA coordinates must be in the range [0, 1].

## rgba to hsva(r, g, b, alpha=1.0)

Converts a color given by its RGBA coordinates to HSVA coordinates (hue, saturation, value, alpha).

Each of the RGBA coordinates must be in the range [0, 1].

## **rgb** to hsl(r, g, b)

Converts a color given by its RGB coordinates to HSL coordinates (hue, saturation, lightness).

Each of the RGB coordinates must be in the range [0, 1].

## **rgb** to hsv(r, g, b)

Converts a color given by its RGB coordinates to HSV coordinates (hue, saturation, value).

Each of the RGB coordinates must be in the range [0, 1].

#### 11.2 Variables

Name	Description
known_colors	Value: {'alice blue': (0.941176470588,
	0.972549019608, 1.0, 1.0)
palettes	Value: {'gray': <gradientpalette th="" with<=""></gradientpalette>
	256 colors>, 'heat': <adva< th=""></adva<>

#### 11.3 Class Palette

object —

igraph.drawing.colors.Palette

Known Subclasses: igraph.drawing.colors.AdvancedGradientPalette, igraph.drawing.colors.Precalculated igraph.drawing.colors.GradientPalette, igraph.drawing.colors.RainbowPalette

Base class of color palettes.

Color palettes are mappings that assign integers from the range 0..n-1 to colors (4-tuples). n is called the size or length of the palette. igraph comes with a number of predefined palettes, so this class is useful for you only if you want to define your own palette. This can be done by subclassing this class and implementing the Palette.\_get method as necessary.

Palettes can also be used as lists or dicts, for the \_\_getitem\_\_ method is overridden properly to call Palette.get.

#### 11.3.1 Methods

## clear cache(self)

Clears the result cache.

The return values of Palette.get are cached. Use this method to clear the cache.

## get(self, v)

Returns the given color from the palette.

Values are cached: if the specific value given has already been looked up, its value will be returned from the cache instead of calculating it again. Use Palette.clear\_cache to clear the cache if necessary.

#### **Parameters**

v: the color to be retrieved. If it is an integer, it is passed to Palette.\_get to be translated to an RGBA quadruplet.

Otherwise it is passed to color\_name\_to\_rgb() to determine the RGBA values.

#### Return Value

the color as an RGBA quadruplet

**Note:** you shouldn't override this method in subclasses, override \_get instead. If you override this method, lookups in the known\_colors dict won't work, so you won't be able to refer to colors by names or RGBA quadruplets, only by integer indices. The caching functionality will disappear as well. However, feel free to override this method if this is exactly the behaviour you want.

## get many(self, colors)

Returns multiple colors from the palette.

Values are cached: if the specific value given has already been looked upon, its value will be returned from the cache instead of calculating it again. Use Palette.clear\_cache to clear the cache if necessary.

#### **Parameters**

colors: the list of colors to be retrieved. The palette class tries to make an educated guess here: if it is not possible to interpret the value you passed here as a list of colors, the class will simply try to interpret it as a single color by forwarding the value to Palette.get.

### Return Value

the colors as a list of RGBA quadruplets. The result will be a list even if you passed a single color index or color name.

# **getitem** (self, v)

Returns the given color from the palette.

Values are cached: if the specific value given has already been looked up, its value will be returned from the cache instead of calculating it again. Use Palette.clear\_cache to clear the cache if necessary.

#### **Parameters**

v: the color to be retrieved. If it is an integer, it is passed to Palette.\_get to be translated to an RGBA quadruplet.

Otherwise it is passed to color\_name\_to\_rgb() to determine the RGBA values.

#### Return Value

the color as an RGBA quadruplet

**Note:** you shouldn't override this method in subclasses, override \_get instead. If you override this method, lookups in the known\_colors dict won't work, so you won't be able to refer to colors by names or RGBA quadruplets, only by integer indices. The caching functionality will disappear as well. However, feel free to override this method if this is exactly the behaviour you want.

# len (self)

Returns the number of colors in this palette

```
plot (self, context, bbox, palette, *args, **kwds)
```

Plots the colors of the palette on the given Cairo context

Supported keyword arguments are:

- border\_width: line width of the border shown around the palette. If zero or negative, the border is turned off. Default is 1.
- grid\_width: line width of the grid that separates palette cells. If zero or negative, the grid is turned off. The grid is also turned off if the size of a cell is less than three times the given line width. Default is 0. Fractional widths are also allowed.
- orientation: the orientation of the palette. Must be one of the following values: left-right, bottom-top, right-left or top-bottom. Possible aliases: horizontal = left-right, vertical = bottom-top, lr = left-right, rl = right-left, tb = top-bottom, bt = bottom-top. The default is left-right.

```
__repr__(self)
repr(x)
Overrides: object.__repr__ extit(inherited documentation)
```

# Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

#### 11.3.2 Properties

Name	Description
length	Returns the number of colors in this palette
Inherited from object	
class	

### 11.4 Class GradientPalette

Base class for gradient palettes

Gradient palettes contain a gradient between two given colors.

## Example:

```
>>> pal = GradientPalette("red", "blue", 5)
>>> pal.get(0)
(1.0, 0.0, 0.0, 1.0)
>>> pal.get(2)
(0.5, 0.0, 0.5, 1.0)
>>> pal.get(4)
(0.0, 0.0, 1.0, 1.0)
```

#### 11.4.1 Methods

```
___init___(self, color1, color2, n=256)
Creates a gradient palette.

Parameters
    color1: the color where the gradient starts.
    color2: the color where the gradient ends.
    n: the number of colors in the palette.
Overrides: object.__init__
```

# $Inherited\ from\ igraph.drawing.colors.Palette(Section\ 11.3)$

# Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

### 11.4.2 Properties

Name Description		
Inherited from igraph.drawin	g.colors.Palette (Section 11.3)	
length		
Inherited from object		
class		

#### 11.5 Class AdvancedGradientPalette

```
object —
igraph.drawing.colors.Palette —
igraph.drawing.colors.AdvancedGradientPalette
```

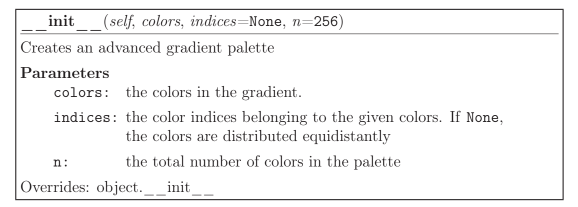
Advanced gradient that consists of more than two base colors.

#### Example:

```
>>> pal = AdvancedGradientPalette(["red", "black", "blue"], n=9)
>>> pal.get(2)
(0.5, 0.0, 0.0, 1.0)
>>> pal.get(7)
```

(0.0, 0.0, 0.75, 1.0)

#### 11.5.1 Methods



# Inherited from igraph.drawing.colors.Palette(Section 11.3)

# Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

## 11.5.2 Properties

Name Description	
Inherited from igraph.drawin	g.colors.Palette (Section 11.3)
length	
Inherited from object	
class	

#### 11.6 Class RainbowPalette

A palette that varies the hue of the colors along a scale.

Colors in a rainbow palette all have the same saturation, value and alpha components, while the hue is varied between two given extremes linearly. This palette has the advantage that it wraps around nicely if the hue is varied between zero and one (which is the default).

## Example:

```
>>> pal = RainbowPalette(n=120)
>>> pal.get(0)
(1.0, 0.0, 0.0, 1.0)
>>> pal.get(20)
(1.0, 1.0, 0.0, 1.0)
>>> pal.get(40)
(0.0, 1.0, 0.0, 1.0)
>>> pal = RainbowPalette(n=120, s=1, v=0.5, alpha=0.75)
>>> pal.get(60)
(0.0, 0.5, 0.5, 0.75)
>>> pal.get(80)
(0.0, 0.0, 0.5, 0.75)
>>> pal.get(100)
(0.5, 0.0, 0.5, 0.75)
>>> pal = RainbowPalette(n=120)
>>> pal2 = RainbowPalette(n=120, start=0.5, end=0.5)
>>> pal.get(60) == pal2.get(0)
>>> pal.get(90) == pal2.get(30)
True
```

This palette was modeled after the rainbow command of R.

#### 11.6.1 Methods

```
___init___(self, n=256, s=1, v=1, start=0, end=1, alpha=1)

Creates a rainbow palette.

Parameters

n: the number of colors in the palette.

s: the saturation of the colors in the palette.

v: the value component of the colors in the palette.

start: the hue at which the rainbow begins (between 0 and 1).

end: the hue at which the rainbow ends (between 0 and 1).

alpha: the alpha component of the colors in the palette.

Overrides: object.__init__
```

# Inherited from igraph.drawing.colors.Palette(Section 11.3)

Inherited	from	object
-----------	------	--------

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

### 11.6.2 Properties

Name Description	
Inherited from igraph.drawing.colors.Palette (Section 11.3)	
length	
Inherited from object	
class	

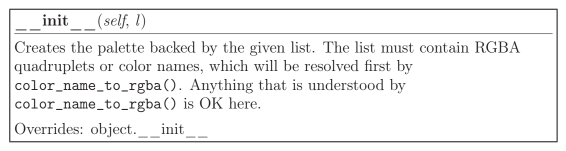
## 11.7 Class PrecalculatedPalette

object —	
igraph.drawing.colors.Palette	
	igraph.drawing.colors.PrecalculatedPalette

Known Subclasses: igraph.drawing.colors.ClusterColoringPalette

A palette that returns colors from a pre-calculated list of colors

### 11.7.1 Methods



# $Inherited\ from\ igraph.drawing.colors.Palette(Section\ 11.3)$

```
__getitem__(), __len__(), __plot__(), __repr__(), clear_cache(), get(), get_many()
```

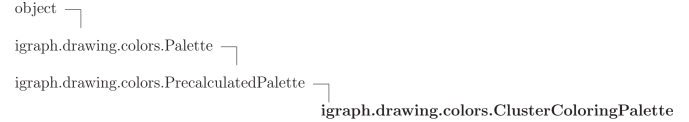
# Inherited from object



## 11.7.2 Properties

Name Description			
Inherited from igraph.drawin	g.colors.Palette (Section 11.3)		
length			
Inherited from object			
class			

# 11.8 Class ClusterColoringPalette



A palette suitable for coloring vertices when plotting a clustering.

This palette tries to make sure that the colors are easily distinguishable. This is achieved by using a set of base colors and their lighter and darker variants, depending on the number of elements in the palette.

When the desired size of the palette is less than or equal to the number of base colors (denoted by n), only the base colors will be used. When the size of the palette is larger than n but less than 2\*n, the base colors and their lighter variants will be used. Between 2\*n and 3\*n, the base colors and their lighter and darker variants will be used. Above 3\*n, more darker and lighter variants will be generated, but this makes the individual colors less and less distinguishable.

#### 11.8.1 Methods

```
__init__(self, n)

Creates the palette backed by the given list. The list must contain RGBA quadruplets or color names, which will be resolved first by color_name_to_rgba(). Anything that is understood by color_name_to_rgba() is OK here.

Overrides: object.__init__ extit(inherited documentation)
```

 $Inherited\ from\ igraph.drawing.colors.Palette(Section\ 11.3)$ 

```
__getitem__(), __len__(), __plot__(), __repr__(), clear_cache(), get(), get_many()
```

Inherited from object

delattr(), _	_format_	(),	$\_$ getattribut $\epsilon$	e(), _	$_{\rm \_hash\_}$	(), _	_new()
reduce(), _	_reduce_	_ex()	,setattr_	_(),	_sizeof_	_(), _	_str(),
subclasshook_	_()						

# 11.8.2 Properties

Name Description		
Inherited from igraph.drawin	g.colors.Palette (Section 11.3)	
length		
Inherited from object		
class		

# 12 Module igraph.drawing.coord

Coordinate systems and related plotting routines

License: GPL

## 12.1 Variables

Name	Description
package	Value: 'igraph.drawing'

# 12.2 Class CoordinateSystem

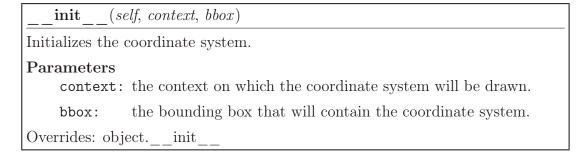
object —	
igraph.drawing.baseclasses.AbstractDrawer —	
$igraph. drawing. base classes. Abstract Cairo Drawer \ -$	
	igraph.drawing.coord.CoordinateSystem

Known Subclasses: igraph.drawing.coord.DescartesCoordinateSystem

Class implementing a coordinate system object.

Coordinate system objects are used when drawing plots which 2D or 3D coordinate system axes. This is an abstract class which must be extended in order to use it. In general, you'll only need the documentation of this class if you intend to implement an own coordinate system not present in igraph yet.

#### 12.2.1 Methods



## draw(self)

Draws the coordinate system.

This method must be overridden in derived classes.

Overrides: igraph.drawing.baseclasses.AbstractDrawer.draw

# local to context(self, x, y)

Converts local coordinates to the context coordinate system (given by the bounding box).

This method must be overridden in derived classes.

## Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

## 12.2.2 Properties

Name	Description
Inherited from igraph.drawin	g.base classes. Abstract Cairo Drawer (Section 10.3)
bbox	
Inherited from object	
class	

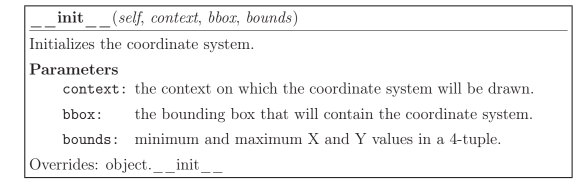
# 12.3 Class DescartesCoordinateSystem

```
object — igraph.drawing.baseclasses.AbstractDrawer — \\igraph.drawing.baseclasses.AbstractCairoDrawer — \\igraph.drawing.coord.CoordinateSystem — .
```

igraph.drawing.coord.DescartesCoordinate

Class implementing a 2D Descartes coordinate system object.

#### 12.3.1 Methods



# draw(self)

Draws the coordinate system.

Overrides: igraph.drawing.baseclasses.AbstractDrawer.draw

# local to context(self, x, y)

Converts local coordinates to the context coordinate system (given by the bounding box).

Overrides: igraph.drawing.coord.CoordinateSystem.local to context

## Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

## 12.3.2 Properties

Name	Description
bbox	Returns the bounding box of the coordinate
	system
bounds	Returns the lower and upper bounds of the X
	and Y values
Inherited from object	
class	

# 13 Module igraph.drawing.edge

Drawers for various edge styles in graph plots.

License: GPL

## 13.1 Class AbstractEdgeDrawer

object — igraph.drawing.edge.AbstractEdgeDrawer

**Known Subclasses:** igraph.drawing.edge.ArrowEdgeDrawer, igraph.drawing.edge.AlphaVaryingEdgeDrawer igraph.drawing.edge.TaperedEdgeDrawer

Abstract edge drawer object from which all concrete edge drawer implementations are derived.

#### 13.1.1 Methods

init (self, context, palette)

Constructs the edge drawer.

**Parameters** 

context: a Cairo context on which the edges will be drawn.

palette: the palette that can be used to map integer color indices

to colors when drawing edges

Overrides: object. init

## draw directed edge(self, edge, src vertex, dest vertex)

Draws a directed edge.

**Parameters** 

edge: the edge to be drawn. Visual properties of the edge

are defined by the attributes of this object.

src\_vertex: the source vertex. Visual properties are given again

as attributes.

dest\_vertex: the target vertex. Visual properties are given again

as attributes.

## draw loop edge(self, edge, vertex)

Draws a loop edge.

The default implementation draws a small circle.

## **Parameters**

edge: the edge to be drawn. Visual properties of the edge are

defined by the attributes of this object.

vertex: the vertex to which the edge is attached. Visual properties

are given again as attributes.

## draw undirected edge(self, edge, src vertex, dest vertex)

Draws an undirected edge.

The default implementation of this method draws undirected edges as straight lines. Loop edges are drawn as small circles.

## **Parameters**

edge: the edge to be drawn. Visual properties of the edge

are defined by the attributes of this object.

src\_vertex: the source vertex. Visual properties are given again

as attributes.

dest\_vertex: the target vertex. Visual properties are given again

as attributes.

# get label position(self, edge, src\_vertex, dest\_vertex)

Returns the position where the label of an edge should be drawn. The default implementation returns the midpoint of the edge and an alignment that tries to avoid overlapping the label with the edge.

## Parameters

edge: the edge to be drawn. Visual properties of the edge

are defined by the attributes of this object.

src\_vertex: the source vertex. Visual properties are given again

as attributes.

dest\_vertex: the target vertex. Visual properties are given again

as attributes.

### Return Value

a tuple containing two more tuples: the desired position of the label and the desired alignment of the label, where the position is given as (x, y) and the alignment is given as (horizontal, vertical). Members of the alignment tuple are taken from constants in the TextAlignment class.

## Inherited from object

$\_\_delattr\_$	_(), _	_format_	(), _	_geta	ttribu	te(	),has	$sh_{}(),$	new_	(),
reduce_	_(), _	_reduce_	ex(	),r	epr	_(),	$_{ m setattr}$	(),	_sizeof_	_(),
str(),	,su	bclasshoo	k()							

### 13.1.2 Properties

Name	Description
Inherited from object	
class	

# 13.2 Class ArrowEdgeDrawer

Edge drawer implementation that draws undirected edges as straight lines and directed edges as arrows.

### 13.2.1 Methods

| draw directed edge(self, edge, src\_vertex, dest\_vertex)|

Draws a directed edge.

## **Parameters**

edge: the edge to be drawn. Visual properties of the edge

are defined by the attributes of this object.

src\_vertex: the source vertex. Visual properties are given again

as attributes.

dest\_vertex: the target vertex. Visual properties are given again

as attributes.

Overrides: igraph.drawing.edge.AbstractEdgeDrawer.draw\_directed\_edge extit(inherited documentation)

# $Inherited\ from\ igraph.drawing.edge.AbstractEdgeDrawer(Section\ 13.1)$

\_\_init\_\_(), draw\_loop\_edge(), draw\_undirected\_edge(), get\_label\_position()

# $Inherited\ from\ object$

$\_\_$ delattr $\_$	_(), _	_format_	(), _	ge	tattribu	ite(	(),ha	$sh_{-}(),$	new_	()
reduce_	_(),	_reduce_	_ex(	$(), \_$	_repr_	_(), _	_setattr	(), _	_sizeof_	_(),
str(),	su	bclasshoo	ok()							

### 13.2.2 Properties

Name	Description
Inherited from object	
class	

# 13.3 Class TaperedEdgeDrawer

object —

 $igraph.drawing.edge. Abstract Edge Drawer \\ -$ 

igraph.drawing.edge.TaperedEdgeDrawer

Edge drawer implementation that draws undirected edges as straight lines and directed edges as tapered lines that are wider at the source and narrow at the destination.

#### 13.3.1 Methods

draw\_directed\_edge(self, edge, src\_vertex, dest\_vertex)

Draws a directed edge.

## **Parameters**

edge: the edge to be drawn. Visual properties of the edge

are defined by the attributes of this object.

src\_vertex: the source vertex. Visual properties are given again

as attributes.

dest\_vertex: the target vertex. Visual properties are given again

as attributes.

Overrides: igraph.drawing.edge.AbstractEdgeDrawer.draw\_directed\_edge extit(inherited documentation)

# $Inherited\ from\ igraph.drawing.edge.AbstractEdgeDrawer(Section\ 13.1)$

\_\_init\_\_(), draw\_loop\_edge(), draw\_undirected\_edge(), get\_label\_position()

# $Inherited\ from\ object$

$\_\_$ delattr $\_$	_(), _	_format_	(), _	ge	tattribu	ite(	(),ha	$sh_{-}(),$	new_	()
reduce_	_(),	_reduce_	_ex(	$(), \_$	_repr_	_(), _	_setattr	(), _	_sizeof_	_(),
str(),	su	bclasshoo	ok()							

### 13.3.2 Properties

Name	Description
Inherited from object	
class	

# 13.4 Class AlphaVaryingEdgeDrawer

object —

igraph.drawing.edge. Abstract Edge Drawer

# igraph.drawing.edge.AlphaVaryingEdgeDrawer

Known Subclasses: igraph.drawing.edge.DarkToLightEdgeDrawer, igraph.drawing.edge.LightToDarkEd

Edge drawer implementation that draws undirected edges as straight lines and directed edges by varying the alpha value of the specified edge color between the source and the destination.

#### 13.4.1 Methods

init (self, context, alpha\_at\_src, alpha\_at\_dest)

Constructs the edge drawer.

## **Parameters**

context: a Cairo context on which the edges will be drawn.

palette: the palette that can be used to map integer color indices

to colors when drawing edges

Overrides: object. init extit(inherited documentation)

# draw directed edge(self, edge, src vertex, dest vertex)

Draws a directed edge.

### **Parameters**

edge: the edge to be drawn. Visual properties of the edge

are defined by the attributes of this object.

src\_vertex: the source vertex. Visual properties are given again

as attributes.

dest\_vertex: the target vertex. Visual properties are given again

as attributes.

Overrides: igraph.drawing.edge.AbstractEdgeDrawer.draw\_directed\_edge extit(inherited documentation)

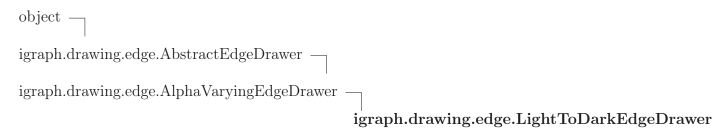
## $Inherited\ from\ igraph.drawing.edge.AbstractEdgeDrawer(Section\ 13.1)$

## Inherited from object

#### 13.4.2 Properties

Name	Description
Inherited from object	
class	

## 13.5 Class LightToDarkEdgeDrawer



Edge drawer implementation that draws undirected edges as straight lines and directed edges by using an alpha value of zero (total transparency) at the source and an alpha value of one (full opacity) at the destination. The alpha value is interpolated in-between.

#### 13.5.1 Methods

```
__init___(self, context)

Constructs the edge drawer.

Parameters
    context: a Cairo context on which the edges will be drawn.

palette: the palette that can be used to map integer color indices to colors when drawing edges

Overrides: object.__init__ extit(inherited documentation)
```

 $Inherited\ from\ igraph. drawing. edge. Alpha Varying Edge Drawer (Section\ 13.4)$ 

```
draw_directed_edge()
```

 $Inherited\ from\ igraph.drawing.edge. Abstract Edge Drawer (Section\ 13.1)$ 

```
{\tt draw\_loop\_edge(),\ draw\_undirected\_edge(),\ get\_label\_position()}
```

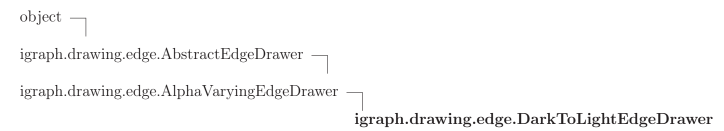
 $Inherited\ from\ object$ 

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

## 13.5.2 Properties

Name	Description
Inherited from object	
class	

## 13.6 Class DarkToLightEdgeDrawer



Edge drawer implementation that draws undirected edges as straight lines and directed edges by using an alpha value of one (full opacity) at the source and an alpha value of zero (total transparency) at the destination. The alpha value is interpolated in-between.

#### 13.6.1 Methods

```
__init__(self, context)

Constructs the edge drawer.

Parameters
    context: a Cairo context on which the edges will be drawn.
    palette: the palette that can be used to map integer color indices to colors when drawing edges

Overrides: object.__init__ extit(inherited documentation)
```

 $Inherited\ from\ igraph. drawing. edge. Alpha Varying Edge Drawer (Section\ 13.4)$ 

```
draw_directed_edge()
```

 $Inherited\ from\ igraph.drawing.edge.AbstractEdgeDrawer(Section\ 13.1)$ 

```
{\tt draw\_loop\_edge(),\ draw\_undirected\_edge(),\ get\_label\_position()}
```

 $Inherited\ from\ object$ 

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

## 13.6.2 Properties

Name	Description
Inherited from object	
class	

# 14 Module igraph.drawing.graph

Drawing routines to draw graphs.

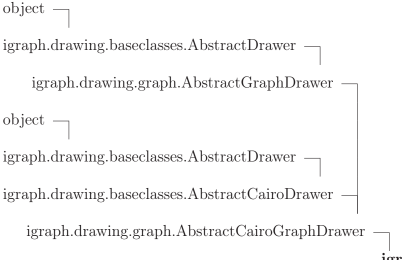
This module contains routines to draw graphs on:

- Cairo surfaces (DefaultGraphDrawer)
- UbiGraph displays (UbiGraphDrawer, see http://ubietylab.net/ubigraph)

It also contains routines to send an igraph graph directly to (Cytoscape<sup>7</sup>) using the (CytoscapeRPC plugin<sup>8</sup>), see CytoscapeGraphDrawer. CytoscapeGraphDrawer can also fetch the current network from Cytoscape and convert it to igraph format.

License: GPL

# 14.1 Class DefaultGraphDrawer



igraph.drawing.graph.DefaultGraphDrawer

Class implementing the default visualisation of a graph.

The default visualisation of a graph draws the nodes on a 2D plane according to a given Layout, then draws a straight or curved edge between nodes connected by edges. This is the visualisation used when one invokes the plot() function on a Graph object.

See Graph.\_\_plot\_\_() for the keyword arguments understood by this drawer.

<sup>&</sup>lt;sup>7</sup>http://www.cytoscape.org

<sup>&</sup>lt;sup>8</sup>http://gforge.nbic.nl/projects/cytoscaperpc/

#### 14.1.1 Methods

 $\label{local_context} \begin{array}{l} \underline{-init}\_\_(self,\; context,\; bbox,\; vertex\_drawer\_factory = < \text{class} \\ \text{'igraph.drawing.vertex.DefaultVertexDrawer'>}, \\ edge\_drawer\_factory = < \text{class 'igraph.drawing.edge.ArrowEdgeDrawer'>}, \\ label\_drawer\_factory = < \text{class 'igraph.drawing.text.TextDrawer'>}) \end{array}$ 

Constructs the graph drawer and associates it to the given Cairo context and the given BoundingBox.

### Parameters

context: the context on which we will draw

bbox: the bounding box within which we will

draw. Can be anything accepted by the constructor of BoundingBox (i.e., a 2-tuple, a 4-tuple or a BoundingBox

object).

vertex\_drawer\_factory: a factory method that returns an

AbstractCairoVertexDrawer instance bound to a given Cairo context. The factory method must take three parameters: the Cairo context, the bounding box of the drawing area and the palette to be used for drawing colored vertices. The default vertex drawer is

DefaultVertexDrawer.

edge\_drawer\_factory: a factory method that returns an

AbstractEdgeDrawer instance bound to a given Cairo context. The factory method must take two parameters: the Cairo context and the palette to be used for drawing colored edges. You can use any of the actual AbstractEdgeDrawer implementations here to control the style of edges drawn by igraph. The default edge drawer is ArrowEdgeDrawer.

label\_drawer\_factory: a factory method that returns a

TextDrawer instance bound to a given Cairo context. The method must take one parameter: the Cairo context. The default label drawer is TextDrawer.

Overrides: object. init

draw(self, graph, palette, \*args, \*\*kwds)

Abstract method, must be implemented in derived classes.

Overrides: igraph.drawing.baseclasses.AbstractDrawer.draw extit(inherited documentation)

# $Inherited\ from\ igraph.drawing.graph.AbstractGraphDrawer$

```
ensure layout()
```

# Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce__ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

### 14.1.2 Properties

Name	Description
Inherited from igraph.drawin	$(g.base classes. Abstract Cairo Drawer\ (Section\ 10.3))$
bbox	
Inherited from object	
class	

# 14.2 Class UbiGraphDrawer

```
object —
igraph.drawing.baseclasses.AbstractDrawer —
igraph.drawing.baseclasses.AbstractXMLRPCDrawer —
object —
igraph.drawing.baseclasses.AbstractDrawer —
igraph.drawing.baseclasses.AbstractDrawer —
```

igraph.drawing.graph.UbiGraphDrawer

Graph drawer that draws a given graph on an UbiGraph display using the XML-RPC API of UbiGraph.

The following vertex attributes are supported: color, label, shape, size. See the Ubigraph documentation for supported shape names. Sizes are relative to the default Ubigraph size.

The following edge attributes are supported: color, label, width. Edge widths are relative

to the default Ubigraph width.

All color specifications supported by igraph (e.g., color names, palette indices, RGB triplets, RGBA quadruplets, HTML format) are understood by the Ubigraph graph drawer.

The drawer also has two attributes, vertex\_defaults and edge\_defaults. These are dictionaries that can be used to set default values for the vertex/edge attributes in Ubigraph.

#### 14.2.1 Methods

\_\_init\_\_(self, url='http://localhost:20738/RPC2')

Constructs an UbiGraph drawer using the display at the given URL.

Parameters

url: the URL where the XML-RPC calls for the service should be addressed to.

service: the name of the service at the XML-RPC address. If

None, requests will be directed to the server proxy object constructed by xmlrpclib.ServerProxy; if not None, the given attribute will be looked up in the server proxy object.

draw(self, graph, \*args, \*\*kwds)

Overrides: object.\_\_init\_\_

Draws the given graph on an UbiGraph display.

### **Parameters**

clear: whether to clear the current UbiGraph display before
 plotting. Default: True.

Overrides: igraph.drawing.baseclasses.AbstractDrawer.draw

# $Inherited\ from\ igraph.drawing.graph.AbstractGraphDrawer$

```
ensure layout()
```

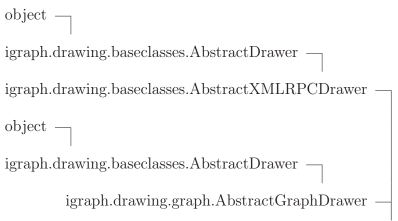
# Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

#### 14.2.2 Properties

Name	Description
Inherited from object	
class	

# 14.3 Class CytoscapeGraphDrawer



igraph.drawing.graph.CytoscapeGraphDra

Graph drawer that sends/receives graphs to/from Cytoscape using CytoscapeRPC.

This graph drawer cooperates with Cytoscape<sup>9</sup> using CytoscapeRPC<sup>10</sup>. You need to install the CytoscapeRPC plugin first and start the XML-RPC server on a given port (port 9000 by default) from the appropriate Plugins submenu in Cytoscape.

Graph, vertex and edge attributes are transferred to Cytoscape whenever possible (i.e. when a suitable mapping exists between a Python type and a Cytoscape type). If there is no suitable Cytoscape type for a Python type, the drawer will use a string attribute on the Cytoscape side and invoke str() on the Python attributes.

If an attribute to be created on the Cytoscape side already exists with a different type, an underscore will be appended to the attribute name to resolve the type conflict.

You can use the network\_id attribute of this class to figure out the network ID of the last graph drawn with this drawer.

<sup>&</sup>lt;sup>9</sup>http://www.cytoscape.org

<sup>&</sup>lt;sup>10</sup>http://wiki.nbic.nl/index.php/CytoscapeRPC

### 14.3.1 Methods

init (self, url='http://localhost:9000/Cytoscape')

Constructs a Cytoscape graph drawer using the XML-RPC interface of Cytoscape at the given URL.

### Parameters

url: the URL where the XML-RPC calls for the service should

be addressed to.

service: the name of the service at the XML-RPC address. If

None, requests will be directed to the server proxy object constructed by xmlrpclib.ServerProxy; if not None, the given attribute will be looked up in the server proxy

object.

Overrides: object. init

 $draw(self, graph, name = 'Network from igraph', create\_view = True, *args, **kwds)$ 

Sends the given graph to Cytoscape as a new network.

## **Parameters**

name: the name of the network in Cytoscape.

create\_view: whether to create a view for the network in

Cytoscape. The default is True.

node\_ids: specifies the identifiers of the nodes to be used in

Cytoscape. This must either be the name of a vertex attribute or a list specifying the identifiers, one for each node in the graph. The default is None, which

simply uses the vertex index for each vertex.

Overrides: igraph.drawing.baseclasses.AbstractDrawer.draw

fetch(self, name=None, directed=False, keep canonical names=False)

Fetches the network with the given name from Cytoscape.

When fetching networks from Cytoscape, the canonicalName attributes of vertices and edges are not converted by default. Use the keep\_canonical\_names parameter to retrieve these attributes as well.

#### **Parameters**

name: the name of the network in Cytoscape.

directed: whether the network is directed.

keep\_canonical\_names: whether to keep the canonicalName

vertex/edge attributes that are added

automatically by Cytoscape

#### Return Value

an appropriately constructed igraph Graph.

## infer cytoscape type(values)

Returns a Cytoscape type that can be used to represent all the values in 'values' and an appropriately converted copy of 'values' that is suitable for an XML-RPC call. Note that the string type in Cytoscape is used as a catch-all type; if no other type fits, attribute values will be converted to string and then posted to Cytoscape.

"None" entries are allowed in 'values', they will be ignored on the Cytoscape side.

# $Inherited\ from\ igraph.drawing.graph.AbstractGraphDrawer$

ensure layout()

# Inherited from object

delattr(	),format(	),ge	etattribu	$te_{-}($	),hash	ı(), ˌ	new_	(),
reduce(	),reduce_ex_	(), _	_repr_	_(),	$_{ m setattr}_{ m }$	_(),	_sizeof	_(),
str (),	subclasshook	()						

## 14.3.2 Properties

Name	Description
Inherited from object	
class	

# 15 Module igraph.drawing.metamagic

Auxiliary classes for the default graph drawer in igraph.

This module contains heavy metaclass magic. If you don't understand the logic behind these classes, probably you don't need them either.

igraph's default graph drawer uses various data sources to determine the visual appearance of vertices and edges. These data sources are the following (in order of precedence):

- The keyword arguments passed to the <code>igraph.plot()</code> function (or to <code>igraph.Graph.\_\_plot\_\_()</code> as a matter of fact, since <code>igraph.plot()</code> just passes these attributes on). For instance, a keyword argument named <code>vertex\_label</code> can be used to set the labels of vertices.
- The attributes of the vertices/edges being drawn. For instance, a vertex that has a label attribute will use that label when drawn by the default graph drawer.
- The global configuration of igraph. For instance, if the global igraph.config.Configuration instance has a key called plotting.vertex\_color, that will be used as a default color for the vertices.
- If all else fails, there is a built-in default; for instance, the default vertex color is "red". This is hard-wired in the source code.

The logic above can be useful in other graph drawers as well, not only in the default one, therefore it is refactored into the classes found in this module. Different graph drawers may inspect different vertex or edge attributes, hence the classes that collect the attributes from the various data sources are generated in run-time using a metaclass called AttributeCollectorMeta. You don't have to use AttributeCollectorMeta directly, just implement a subclass of AttributeCollectorBase and it will ensure that the appropriate metaclass is used. With AttributeCollectorBase, you can use a simple declarative syntax to specify which attributes you are interested in. For example:

```
class VisualEdgeBuilder(AttributeCollectorBase):
    arrow_size = 1.0
    arrow_width = 1.0
    color = ("black", palette.get)
    width = 1.0

for edge in VisualEdgeBuilder(graph.es):
    print edge.color
```

The above class is a visual edge builder — a class that gives the visual attributes of the edges of a graph that is specified at construction time. It specifies that the attributes we are interested in are arrow\_size, arrow\_width, color and width; the default values are also given. For color, we also specify that a method called {palette.get} should be called on every attribute value to translate color names to RGB values. For the other three attributes, float will implicitly be called on all attribute values, this is inferred from the type of the default value itself.

See Also: AttributeCollectorMeta, AttributeCollectorBase

## 15.1 Class AttributeSpecification

## igraph.drawing.metamagic.AttributeSpecification

Class that describes how the value of a given attribute should be retrieved.

The class contains the following members:

- name: the name of the attribute. This is also used when we are trying to get its value from a vertex/edge attribute of a graph.
- alt\_name: alternative name of the attribute. This is used when we are trying to get its value from a Python dict or an igraph.Configuration object. If omitted at construction time, it will be equal to name.
- default: the default value of the attribute when none of the sources we try can provide a meaningful value.
- transform: optional transformation to be performed on the attribute value. If None or omitted, it defaults to the type of the default value.
- func: when given, this function will be called with an index in order to derive the value of the attribute.

#### 15.1.1 Methods

$\boxed{\frac{\text{init}}{\textit{func}} = \text{None}, \ \textit{default} = \text{None}, \ \textit{alt}\_\textit{name} = \text{None}, \ \textit{transform} = \text{None}, \ \textit{func} = \text{None})}$
xinit() initializes x; see help(type(x)) for signature
Overrides: objectinit extit(inherited documentation)

## Inherited from object

dela	ttr()	$, \_\_format_$	(),	_getattrib	$\mathrm{oute}\_\_($	(),hash	n(),	new_	()
redu	.ce()	,reduce_	$ex_{()}$	repr_	(), _	_setattr_	_(),	_sizeof_	_(),
$\operatorname{str}$	(),	subclasshoo	ok ()						

#### 15.1.2 Properties

Name	Description
accessor	
alt_name	
default	
func	
name	

continued on next page

Name	Description
transform	
Inherited from object	
class	

#### 15.2 Class AttributeCollectorBase

object —

## igraph.drawing.metamagic.AttributeCollectorBase

Base class for attribute collector subclasses. Classes that inherit this class may use a declarative syntax to specify which vertex or edge attributes they intend to collect. See AttributeCollectorMeta for the details.

#### 15.2.1 Methods

 $\_\_init\_\_(self, seq, kwds=$ None)

Constructs a new attribute collector that uses the given vertex/edge sequence and the given dict as data sources.

## **Parameters**

seq: an igraph. VertexSeq or igraph. EdgeSeq class that will be

used as a data source for attributes.

kwds: a Python dict that will be used to override the attributes

collected from seq if necessary.

Overrides: object.\_\_init\_\_\_\_

getitem (self, index)

Returns the collected attributes of the vertex/edge with the given index.

\_\_len\_\_(self)

# Inherited from object

\_\_delattr\_\_(), \_\_format\_\_(), \_\_getattribute\_\_(), \_\_hash\_\_(), \_\_new\_\_(), \_\_reduce\_\_(), \_\_reduce\_ex\_\_(), \_\_repr\_\_(), \_\_setattr\_\_(), \_\_sizeof\_\_(), \_\_str\_\_(), \_\_subclasshook\_\_()

### 15.2.2 Properties

Name	Description
Inherited from object	
class	

# 16 Module igraph.drawing.shapes

Shape drawing classes for igraph

Vertex shapes in igraph are usually referred to by short names like "rect" or "circle". This module contains the classes that implement the actual drawing routines for these shapes, and a resolver class that determines the appropriate shape drawer class given the short name.

Classes that are derived from ShapeDrawer in this module are automatically registered by ShapeDrawerDirectory. If you implement a custom shape drawer, you must register it in ShapeDrawerDirectory manually if you wish to refer to it by a name in the shape attribute of vertices.

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### 16.1 Class ShapeDrawerDirectory

object | | igraph.drawing.shapes.ShapeDrawerDirectory

Static class that resolves shape names to their corresponding shape drawer classes.

Classes that are derived from ShapeDrawer in this module are automatically registered by ShapeDrawerDirectory when the module is loaded for the first time.

## 16.1.1 Methods

register(cls, drawer\_class)

Registers the given shape drawer class under the given names.

Parameters

drawer\_class: the shape drawer class to be registered

## register namespace(cls, namespace)

Registers all ShapeDrawer classes in the given namespace

### **Parameters**

namespace: a Python dict mapping names to Python objects.

## **resolve**(cls, shape)

Given a shape name, returns the corresponding shape drawer class

### **Parameters**

shape: the name of the shape

### Return Value

the corresponding shape drawer class

#### Raises

ValueError if the shape is unknown

# ${\bf resolve\_default}(\mathit{cls}, \mathit{shape}, \mathit{default} {=} {\footnotesize {<}} {\tt class}$

'igraph.drawing.shapes.NullDrawer'>)

Given a shape name, returns the corresponding shape drawer class or the given default shape drawer if the shape name is unknown.

### **Parameters**

shape: the name of the shape

default: the default shape drawer to return when the shape is

unknown

#### Return Value

the shape drawer class corresponding to the given name or the default shape drawer class if the name is unknown

## Inherited from object

$\_$ _delattr $\_$ (),	$, \_\_format\_\_$	$_{-}(),  \{\mathrm{getattribu}}$	$te\_\_(), \_\_1$	$\operatorname{hash}\_\_(),$	$_{1}$ init()
new(),	_reduce()	,reduce_ex_	_(),repr_	(),s	$setattr_{-}(),$
sizeof (),	str (),	subclasshook	()		

### 16.1.2 Properties

Name	Description
Inherited from object	
class	

#### 16.1.3 Class Variables

Name	Description
known_shapes	Value: {'': <class< th=""></class<>
	'igraph.drawing.shapes.NullDrawer'>,
	'arrow':

# 17 Module igraph.drawing.text

Drawers for labels on plots.

@undocumented: test License: GPL

## 17.1 Class TextAlignment

Text alignment constants.

### 17.1.1 Methods

## Inherited from object

## 17.1.2 Properties

Name	Description
Inherited from object	
class	

### 17.1.3 Class Variables

Name	Description
BOTTOM	Value: 'bottom'
CENTER	Value: 'center'
LEFT	Value: 'left'
RIGHT	Value: 'right'
TOP	Value: 'top'

## 17.2 Class TextDrawer

object —	
igraph.drawing.baseclasses.AbstractDrawer —	
igraph. drawing. base classes. Abstract Cairo Drawer	
	igraph.drawing.text.TextDrawer

Class that draws text on a Cairo context.

This class supports multi-line text unlike the original Cairo text drawing methods.

#### 17.2.1 Methods

\_\_init\_\_\_(self, context, text='', halign='center', valign='center')

Constructs a new instance that will draw the given text on the given Cairo context. Parameters

context: the context on which we will draw

bbox: the bounding box within which we will draw. Can be

anything accepted by the constructor of BoundingBox (i.e., a 2-tuple, a 4-tuple or a BoundingBox object).

Overrides: object. init

draw(self, wrap=False)

Draws the text in the current bounding box of the drawer.

Since the class itself is an instance of AbstractCairoDrawer, it has an attribute named bbox which will be used as a bounding box. Parameters

wrap: whether to allow re-wrapping of the text if it does not fit within the bounding box horizontally. (type=boolean)

Overrides: igraph.drawing.baseclasses.AbstractDrawer.draw

## get text layout(self, x=None, y=None, width=None, wrap=False)

Calculates the layout of the current text. x and y denote the coordinates where the drawing should start. If they are both None, the current position of the context will be used.

Vertical alignment settings are not taken into account in this method as the text is not drawn within a box. **Parameters** 

x: The X coordinate of the reference point where the layout should start. (type=float or None)

y: The Y coordinate of the reference point where the layout should start. (type=float or None)

width: The width of the box in which the text will be fitted. It matters only when the text is right-aligned or centered. The text will overflow the box if any of the lines is longer than the box width and wrap is False. (type=float or None)

wrap: whether to allow re-wrapping of the text if it does not fit within the given width. (type=boolean)

#### Return Value

a list consisting of (x, y, line) tuples where x and y refer to reference points on the Cairo canvas and line refers to the corresponding text that should be plotted there.

draw at(self, x=None, y=None, width=None, wrap=False)

Draws the text by setting up an appropriate path on the Cairo context and filling it. x and y denote the coordinates where the drawing should start. If they are both None, the current position of the context will be used.

Vertical alignment settings are not taken into account in this method as the text is not drawn within a box. **Parameters** 

- x: The X coordinate of the reference point where the drawing should start. (type=float or None)
- y: The Y coordinate of the reference point where the drawing should start. (type=float or None)
- width: The width of the box in which the text will be fitted. It matters only when the text is right-aligned or centered. The text will overflow the box if any of the lines is longer than the box width. (type=float or None)
- wrap: whether to allow re-wrapping of the text if it does not fit within the given width. (type=boolean)

## text\_extents(self)

Returns the X-bearing, Y-bearing, width, height, X-advance and Y-advance of the text.

For multi-line text, the X-bearing and Y-bearing correspond to the first line, while the X-advance is extracted from the last line. and the Y-advance is the sum of all the Y-advances. The width and height correspond to the entire bounding box of the text.

#### Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

#### 17.2.2 Properties

Name Description			
text	Returns the text to be drawn.		
Inherited from igraph.drawing.baseclasses.AbstractCairoDrawer (Section 10.3)			
bbox			
Inherited from object			

continued on next page

Name	Description		
class			

## 17.2.3 Class Variables

Name	Description
BOTTOM	Value: 'bottom'
CENTER	Value: 'center'
LEFT	Value: 'left'
RIGHT	Value: 'right'
TOP	Value: 'top'

## 18 Module igraph.drawing.utils

Utility classes for drawing routines.

License: GPL

## 18.1 Class Rectangle

object — igraph.drawing.utils.Rectangle

Known Subclasses: igraph.drawing.utils.BoundingBox

Class representing a rectangle.

#### 18.1.1 Methods

init (self, \*args)

Creates a rectangle.

The corners of the rectangle can be specified by either a tuple (four items, two for each corner, respectively), four separate numbers (X and Y coordinates for each corner) or two separate numbers (width and height, the upper left corner is assumed to be at (0,0))

Overrides: object.\_\_init\_\_

**contract**(self, margins)

Contracts the rectangle by the given margins.

Return Value

a new Rectangle object.

**expand**(self, margins)

Expands the rectangle by the given margins.

Return Value

a new Rectangle object.

```
isdisjoint(self, other)
Returns "True" if the two rectangles have no intersection.
Example:
   >>> r1 = Rectangle(10, 10, 30, 30)
   >>> r2 = Rectangle(20, 20, 50, 50)
   >>> r3 = Rectangle(70, 70, 90, 90)
   >>> r1.isdisjoint(r2)
   False
   >>> r2.isdisjoint(r1)
   False
   >>> r1.isdisjoint(r3)
   >>> r3.isdisjoint(r1)
   True
```

## isempty(self)

Returns "True" if the rectangle is empty (i.e. it has zero width and height).

```
Example:
   >>> r1 = Rectangle(10, 10, 30, 30)
  >>> r2 = Rectangle(70, 70, 90, 90)
   >>> r1.isempty()
   False
   >>> r2.isempty()
   False
   >>> r1.intersection(r2).isempty()
   True
```

## intersection(self, other)

Returns the intersection of this rectangle with another.

#### Example:

```
>>> r1 = Rectangle(10, 10, 30, 30)
>>> r2 = Rectangle(20, 20, 50, 50)
>>> r3 = Rectangle(70, 70, 90, 90)
>>> r1.intersection(r2)
Rectangle(20.0, 20.0, 30.0, 30.0)
>>> r2 & r1
Rectangle(20.0, 20.0, 30.0, 30.0)
>>> r2.intersection(r1) == r1.intersection(r2)
True
>>> r1.intersection(r3)
Rectangle(0.0, 0.0, 0.0, 0.0)
```

#### (self, other) and

Returns the intersection of this rectangle with another.

```
Example:
  >>> r1 = Rectangle(10, 10, 30, 30)
  >>> r2 = Rectangle(20, 20, 50, 50)
  >>> r3 = Rectangle(70, 70, 90, 90)
  >>> r1.intersection(r2)
  Rectangle(20.0, 20.0, 30.0, 30.0)
  >>> r2 & r1
  Rectangle(20.0, 20.0, 30.0, 30.0)
  >>> r2.intersection(r1) == r1.intersection(r2)
  True
  >>> r1.intersection(r3)
  Rectangle(0.0, 0.0, 0.0, 0.0)
```

```
translate(self, dx, dy)
Translates the rectangle in-place.

Example:

>>> r = Rectangle(10, 20, 50, 70)
>>> r.translate(30, -10)
>>> r

Rectangle(40.0, 10.0, 80.0, 60.0)

Parameters
    dx: the X coordinate of the translation vector
    dy: the Y coordinate of the translation vector
```

## **union**(self, other)

Returns the union of this rectangle with another.

The resulting rectangle is the smallest rectangle that contains both rectangles.

## Example:

```
>>> r1 = Rectangle(10, 10, 30, 30)
>>> r2 = Rectangle(20, 20, 50, 50)
>>> r3 = Rectangle(70, 70, 90, 90)
>>> r1.union(r2)
Rectangle(10.0, 10.0, 50.0, 50.0)
>>> r2 | r1
Rectangle(10.0, 10.0, 50.0, 50.0)
>>> r2.union(r1) == r1.union(r2)
True
>>> r1.union(r3)
Rectangle(10.0, 10.0, 90.0, 90.0)
```

```
or (self, other)
```

Returns the union of this rectangle with another.

The resulting rectangle is the smallest rectangle that contains both rectangles.

#### Example:

```
>>> r1 = Rectangle(10, 10, 30, 30)

>>> r2 = Rectangle(20, 20, 50, 50)

>>> r3 = Rectangle(70, 70, 90, 90)

>>> r1.union(r2)

Rectangle(10.0, 10.0, 50.0, 50.0)

>>> r2 | r1

Rectangle(10.0, 10.0, 50.0, 50.0)

>>> r2.union(r1) == r1.union(r2)

True

>>> r1.union(r3)

Rectangle(10.0, 10.0, 90.0, 90.0)
```

## ior (self, other)

Expands this rectangle to include itself and another completely while still being as small as possible.

#### Example:

```
>>> r1 = Rectangle(10, 10, 30, 30)

>>> r2 = Rectangle(20, 20, 50, 50)

>>> r3 = Rectangle(70, 70, 90, 90)

>>> r1 |= r2

>>> r1

Rectangle(10.0, 10.0, 50.0, 50.0)

>>> r1 |= r3

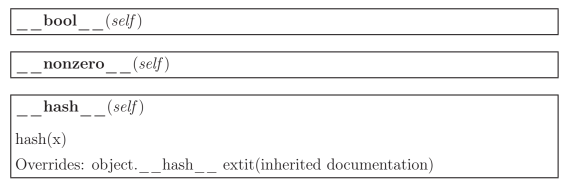
>>> r1

Rectangle(10.0, 10.0, 90.0, 90.0)
```

```
__repr__(self)
repr(x)
Overrides: object.__repr__ extit(inherited documentation)
```

```
__eq__(self, other)
```

```
__ne__(self, other)
```



## $Inherited\ from\ object$

```
\label{lem:condition} $$\_\_delattr\_(), \_\_format\_\_(), \_\_getattribute\_\_(), \_\_new\_\_(), \_\_reduce\_\_(), \_\_reduce\_ex\_\_(), \_\_setattr\_\_(), \_\_sizeof\_\_(), \_\_str\_\_(), \_\_subclasshook\_\_()
```

#### 18.1.2 Properties

Name	Description			
coords	The coordinates of the corners.			
	The coordinates are returned as a 4-tuple in the			
	following order: left edge, top edge, right edge,			
	bottom edge.			
width	The width of the rectangle			
height	The height of the rectangle			
left	The X coordinate of the left side of the box			
right	The X coordinate of the right side of the box			
top	The Y coordinate of the top edge of the box			
bottom	The Y coordinate of the bottom edge of the box			
midx	The X coordinate of the center of the box			
midy	The Y coordinate of the center of the box			
shape	The shape of the rectangle (width, height)			
Inherited from object				
class				

## 18.2 Class BoundingBox

object  $\neg$  igraph.drawing.utils.Rectangle  $\neg$  igraph.drawing.utils.BoundingBox

Class representing a bounding box (a rectangular area) that encloses some objects.

#### 18.2.1 Methods

```
__ior__(self, other)
Replaces this bounding box with the union of itself and another.

Example:

>>> box1 = BoundingBox(10, 20, 50, 60)
>>> box2 = BoundingBox(70, 40, 100, 90)
>>> box1 |= box2
>>> print(box1)
BoundingBox(10.0, 20.0, 100.0, 90.0)

Overrides: igraph.drawing.utils.Rectangle.__ior__
```

```
___or__(self, other)

Takes the union of this bounding box with another.

The result is a bounding box which encloses both bounding boxes.

Example:

>>> box1 = BoundingBox(10, 20, 50, 60)
>>> box2 = BoundingBox(70, 40, 100, 90)
>>> box1 | box2
BoundingBox(10.0, 20.0, 100.0, 90.0)

Overrides: igraph.drawing.utils.Rectangle.__or__
```

## Inherited from igraph.drawing.utils.Rectangle(Section 18.1)

## Inherited from object

```
\label{lem:condition} $$\_\_delattr\_(), \_\_format\__(), \_\_getattribute\__(), \_\_new\__(), \_\_reduce\__(), \_\_reduce\_ex\__(), \_\_setattr\_(), \_\_sizeof\__(), \_\_str\__(), \_\_subclasshook\__()
```

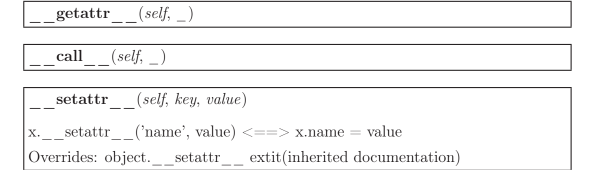
#### 18.2.2 Properties

Name	Description			
Inherited from igraph.drawing.utils.Rectangle (Section 18.1)				
bottom, coords, height, left, midx, midy, right, shape, top, width				
Inherited from object				
class				

#### 18.3 Class FakeModule

Fake module that raises an exception for everything

#### 18.3.1 Methods



## Inherited from object

#### 18.3.2 Properties

Name	Description
Inherited from object	
class	

#### 18.4 Class Point



Class representing a point on the 2D plane.

## 18.4.1 Methods

$\underline{} \mathbf{new} \underline{} (\mathit{cls},  x,  y)$
Creates a new point with the given coordinates
Return Value a new object with type S, a subtype of T
Overrides: objectnew
( 16)
$-$ repr $_{-}$ (self)
Returns a nicely formatted representation of the point
Overrides: objectrepr
$\_\_getnewargs\_\_(self)$
Return self as a plain tuple. Used by copy and pickle.
Overrides: tuplegetnewargs
Overrides. tuplegettlewargs
add(self, other)
Adds the coordinates of a point to another one
Overrides: tupleadd
and (solf other)
$ \frac{-\operatorname{sub}_{-(self, other)}}{\operatorname{sub}_{-(self, other)}} $
Subtracts the coordinates of a point to another one
mul(self, scalar)
Multiplies the coordinates by a scalar
Overrides: tuplemul
rmul(self, scalar)
Multiplies the coordinates by a scalar
Overrides: tuplermul
div (self, scalar)
Divides the coordinates by a scalar

## as polar(self)

Returns the polar coordinate representation of the point.

#### Return Value

the radius and the angle in a tuple.

## **distance**(self, other)

Returns the distance of the point from another one.

## Example:

```
>>> p1 = Point(5, 7)
>>> p2 = Point(8, 3)
>>> p1.distance(p2)
5.0
```

## interpolate(self, other, ratio=0.5)

Linearly interpolates between the coordinates of this point and another one.

#### **Parameters**

other: the other point

ratio: the interpolation ratio between 0 and 1. Zero will return this

point, 1 will return the other point.

#### length(self)

Returns the length of the vector pointing from the origin to this point.

#### normalized(self)

Normalizes the coordinates of the point s.t. its length will be 1 after normalization. Returns the normalized point.

#### sq length(self)

Returns the squared length of the vector pointing from the origin to this point.

## towards(self, other, distance=0)

Returns the point that is at a given distance from this point towards another one.

## FromPolar(cls, radius, angle)

Constructs a point from polar coordinates.

'radius' is the distance of the point from the origin; 'angle' is the angle between the X axis and the vector pointing to the point from the origin.

## Inherited from tuple

## Inherited from object

#### 18.4.2 Properties

Name	Description
X	Alias for field number 0
У	Alias for field number 1
Inherited from object	
class	

## 19 Module igraph.drawing.vertex

Drawing routines to draw the vertices of graphs.

This module provides implementations of vertex drawers, i.e. drawers that the default graph drawer will use to draw vertices.

License: GPL

#### 19.1 Class AbstractVertexDrawer

Known Subclasses: igraph.drawing.vertex.AbstractCairoVertexDrawer

Abstract vertex drawer object from which all concrete vertex drawer implementations are derived.

#### 19.1.1 Methods

init(self, palette, layout)		
Constructs the vertex drawer and associates it to the given palette.		
Parameters		
palette: the palette that can be used to map integer color indices to colors when drawing vertices		
layout: the layout of the vertices in the graph being drawn		
Overrides: objectinit		

draw(self, visual\_vertex, vertex, coords)

Draws the given vertex.

Parameters

visual\_vertex: object specifying the visual properties of the vertex. Its structure is defined by the VisualVertexBuilder of the DefaultGraphDrawer; see its source code.

vertex: the raw igraph vertex being drawn

coords: the X and Y coordinates of the vertex as specified by the layout algorithm, scaled into the bounding box.

Overrides: igraph.drawing.baseclasses.AbstractDrawer.draw

## Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

#### 19.1.2 Properties

Name	Description
Inherited from object	
class	

#### 19.2 Class AbstractCairoVertexDrawer

object —
igraph.drawing.baseclasses.AbstractDrawer $\overline{}$
igraph.drawing.vertex.AbstractVertexDrawer —
object —
igraph.drawing.baseclasses.AbstractDrawer $\overline{}$
igraph.drawing.baseclasses.AbstractCairoDrawer —

igraph.drawing.vertex.AbstractCairoVertexDra

Known Subclasses: igraph.drawing.vertex.DefaultVertexDrawer

Abstract base class for vertex drawers that draw on a Cairo canvas.

#### 19.2.1 Methods

\_\_init\_\_\_(self, context, bbox, palette, layout)

Constructs the vertex drawer and associates it to the given Cairo context and the given BoundingBox.

#### **Parameters**

context: the context on which we will draw

bbox: the bounding box within which we will draw. Can be

anything accepted by the constructor of BoundingBox (i.e., a 2-tuple, a 4-tuple or a BoundingBox object).

palette: the palette that can be used to map integer color indices

to colors when drawing vertices

layout: the layout of the vertices in the graph being drawn

Overrides: object. init

# $Inherited\ from\ igraph.drawing.vertex.AbstractVertexDrawer(Section\ 19.1)$ draw()

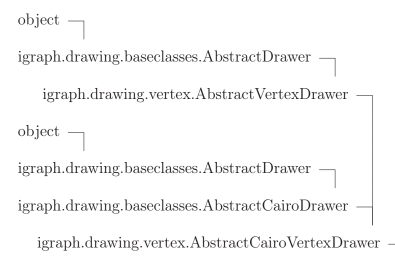
## Inherited from object

delattr(	),format(),	getattrib	$ute_{-}(), _{-}$	hash()	$, \{\text{new}}_{-}(),$
reduce(	),reduce_ex	_(),	(),se	tattr(), _	$\_$ sizeof $\_$ (),
str(), _	_subclasshook(	()			

#### 19.2.2 Properties

Name	Description
Inherited from igraph.drawin	g.base classes. Abstract Cairo Drawer (Section 10.3)
bbox	
Inherited from object	
class	

#### 19.3 Class DefaultVertexDrawer



igraph.drawing.vertex.DefaultVertexDrawe

The default vertex drawer implementation of igraph.

#### 19.3.1 Methods

\_\_init\_\_(self, context, bbox, palette, layout)

Constructs the vertex drawer and associates it to the given Cairo context and the given BoundingBox.

#### **Parameters**

context: the context on which we will draw

bbox: the bounding box within which we will draw. Can be

anything accepted by the constructor of BoundingBox (i.e., a 2-tuple, a 4-tuple or a BoundingBox object).

palette: the palette that can be used to map integer color indices

to colors when drawing vertices

layout: the layout of the vertices in the graph being drawn

Overrides: object.\_\_init\_\_ extit(inherited documentation)

draw(self, visual\_vertex, vertex, coords)

Draws the given vertex.

#### **Parameters**

visual\_vertex: object specifying the visual properties of the

vertex. Its structure is defined by the

VisualVertexBuilder of the DefaultGraphDrawer;

see its source code.

vertex: the raw igraph vertex being drawn

coords: the X and Y coordinates of the vertex as specified

by the layout algorithm, scaled into the bounding

box.

Overrides: igraph.drawing.baseclasses.AbstractDrawer.draw extit(inherited documentation)

## Inherited from object

delattr(	),format(),	$\_\_$ getattrib	$ute_{-}(), _{-}$	$_{-}$ hash $_{-}$ (	$(), \underline{}$ new $\underline{}$ $(),$
reduce(	),reduce_ex	(),repr_	(),se	$etattr_{-}(),$	$\_$ sizeof $\_$ (),
str(), _	_subclasshook(				

#### 19.3.2 Properties

Name	Description
Inherited from igraph.drawin	g.base classes. Abstract Cairo Drawer (Section 10.3)
bbox	
Inherited from object	
class	

## 20 Module igraph.layout

Layout-related code in the IGraph library.

This package contains the implementation of the Layout object.

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#### 20.1 Variables

Name	Description
package	Value: 'igraph'

#### 20.2 Class Layout

Represents the layout of a graph.

A layout is practically a list of coordinates in an n-dimensional space. This class is generic in the sense that it can store coordinates in any n-dimensional space.

Layout objects are not associated directly with a graph. This is deliberate: there were times when I worked with almost identical copies of the same graph, the only difference was that they had different colors assigned to the vertices. It was particularly convenient for me to use the same layout for all of them, especially when I made figures for a paper. However, igraph will of course refuse to draw a graph with a layout that has less coordinates than the node count of the graph.

Layouts behave exactly like lists when they are accessed using the item index operator ([...]). They can even be iterated through. Items returned by the index operator are only copies of the coordinates, but the stored coordinates can be modified by directly assigning to an index.

```
>>> layout = Layout([(0, 1), (0, 2)])
>>> coords = layout[1]
>>> print coords
[0, 2]
>>> coords = (0, 3)
>>> print layout[1]
[0, 2]
>>> layout[1] = coords
>>> print layout[1]
[0, 3]
20.2.1 Methods
         init
                 (self, coords=None, dim=None)
     Constructor.
     Parameters
          coords: the coordinates to be stored in the layout.
         dim:
                   the number of dimensions. If None, the number of
                   dimensions is determined automatically from the length of
                   the first item of the coordinate list. If there are no entries
                   in the coordinate list, the default will be 2. Generally, this
                   should be given if the length of the coordinate list is zero,
                   otherwise it should be left as is.
     Overrides: object.__init__
         len (self)
```

getitem

setitem

(self, idx)

(self, idx, value)

Class Layout Module igraph.layout

## append(self, value)

Appends a new point to the layout

## mirror(self, dim)

Mirrors the layout along the given dimension(s)

#### Parameters

dim: the list of dimensions or a single dimension

## rotate(self, angle, dim1=0, dim2=1, \*\*kwds)

Rotates the layout by the given degrees on the plane defined by the given two dimensions.

#### **Parameters**

angle: the angle of the rotation, specified in degrees.

dim1: the first axis of the plane of the rotation.

dim2: the second axis of the plane of the rotation.

origin: the origin of the rotation. If not specified, the origin will be

the origin of the coordinate system.

## scale(self, \*args, \*\*kwds)

Scales the layout.

Scaling parameters can be provided either through the scale keyword argument or through plain unnamed arguments. If a single integer or float is given, it is interpreted as a uniform multiplier to be applied on all dimensions. If it is a list or tuple, its length must be equal to the number of dimensions in the layout, and each element must be an integer or float describing the scaling coefficient in one of the dimensions.

#### **Parameters**

scale: scaling coefficients (integer, float, list or tuple)

origin: the origin of scaling (this point will stay in place).

Optional, defaults to the origin of the coordinate system

being used.

## translate(self, \*args, \*\*kwds)

Translates the layout.

The translation vector can be provided either through the v keyword argument or through plain unnamed arguments. If unnamed arguments are used, the vector can be supplied as a single list (or tuple) or just as a series of arguments. In all cases, the translation vector must have the same number of dimensions as the layout.

#### **Parameters**

v: the translation vector

```
to_radial(self, min_angle=100, max_angle=80, min_radius=0.0, max_radius=1.0)
```

Converts a planar layout to a radial one

This method applies only to 2D layouts. The X coordinate of the layout is transformed to an angle, with  $\min(x)$  corresponding to the parameter called  $\min\_angle$  and  $\max(y)$  corresponding to  $\max\_angle$ . Angles are given in degrees, zero degree corresponds to the direction pointing upwards. The Y coordinate is interpreted as a radius, with  $\min(y)$  belonging to the minimum and  $\max(y)$  to the maximum radius given in the arguments.

This is not a fully generic polar coordinate transformation, but it is fairly useful in creating radial tree layouts from ordinary top-down ones (that's why the Y coordinate belongs to the radius). It can also be used in conjunction with the Fruchterman-Reingold layout algorithm via its *miny* and *maxy* parameters (see Graph.layout\_fruchterman\_reingold) to produce radial layouts where the radius belongs to some property of the vertices.

#### Parameters

min\_angle: the angle corresponding to the minimum X value max\_angle: the angle corresponding to the maximum X value min\_radius: the radius corresponding to the minimum Y value max\_radius: the radius corresponding to the maximum Y value

## **transform**(self, function, \*args, \*\*kwds)

Performs an arbitrary transformation on the layout

Additional positional and keyword arguments are passed intact to the given function.

#### **Parameters**

function: a function which receives the coordinates as a tuple and returns the transformed tuple.

## **centroid**(self)

Returns the centroid of the layout.

The centroid of the layout is the arithmetic mean of the points in the layout.

## Return Value

the centroid as a list of floats

## **boundaries**(self, border=0)

Returns the boundaries of the layout.

The boundaries are the minimum and maximum coordinates along all dimensions.

#### **Parameters**

border: this value gets subtracted from the minimum bounds and gets added to the maximum bounds before returning the coordinates of the box. Defaults to zero.

#### Return Value

the minimum and maximum coordinates along all dimensions, in a tuple containing two lists, one for the minimum coordinates, the other one for the maximum.

#### Raises

ValueError if the layout contains no layout items

## **bounding box**(*self*, *border*=0)

Returns the bounding box of the layout.

The bounding box of the layout is the smallest box enclosing all the points in the layout.

#### **Parameters**

border: this value gets subtracted from the minimum bounds and gets added to the maximum bounds before returning the coordinates of the box. Defaults to zero.

#### Return Value

the coordinates of the lower left and the upper right corner of the box. "Lower left" means the minimum coordinates and "upper right" means the maximum. These are encapsulated in a BoundingBox object.

# $\mathbf{center}(\mathit{self}, *\mathit{args}, *\overline{*kwds})$

Centers the layout around the given point.

The point itself can be supplied as multiple unnamed arguments, as a simple unnamed list or as a keyword argument. This operation moves the centroid of the layout to the given point. If no point is supplied, defaults to the origin of the coordinate system.

#### **Parameters**

p: the point where the centroid of the layout will be after the operation.

#### copy(self)

Creates an exact copy of the layout.

| fit into(self, bbox, keep\_aspect\_ratio=True)

Fits the layout into the given bounding box.

The layout will be modified in-place.

## **Parameters**

bbox:

the bounding box in which to fit the layout. If the dimension of the layout is d, it can either be a d-tuple (defining the sizes of the box), a 2d-tuple (defining the coordinates of the top left and the bottom right point of the box), or a BoundingBox object (for 2D layouts only).

keep\_aspect\_ratio: whether to keep the aspect ratio of the

current layout. If False, the layout will be rescaled to fit exactly into the bounding box. If True, the original aspect ratio of the layout will be kept and it will be centered within the bounding box.

## Inherited from object

delattr(	$), \_\_$ form	$\operatorname{rat}_{-}(),$	ge	tattribute	=(), _	$\_\_\mathrm{hash}\_$	(), _	_new	()
reduce()	),redu	ce_ex	_(),	$_{ m setattr} _{ m }$	_(),	_sizeof	_(),	$_{ m str}_{ m -}$	_(),
subclasshoo	ok()								

#### 20.2.2 Properties

Name	Description
dim	Returns the number of dimensions
coords	The coordinates as a list of lists
Inherited from object	
class	

## 21 Module igraph.matching

Classes representing matchings on graphs.

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#### 21.1 Variables

Name	Description
package	Value: 'igraph'

## 21.2 Class Matching

```
object — igraph.matching.Matching
```

A matching of vertices in a graph.

A matching of an undirected graph is a set of edges such that each vertex is incident on at most one matched edge. When each vertex is incident on *exactly* one matched edge, the matching called *perfect*. This class is used in **igraph** to represent non-perfect and perfect matchings in undirected graphs.

This class is usually not instantiated directly, everything is taken care of by the functions that return matchings.

#### Examples:

```
>>> from igraph import Graph
>>> g = Graph.Famous("noperfectmatching")
>>> matching = g.maximum_matching()
```

#### 21.2.1 Methods

init (self, graph, matching, types=None)

Initializes the matching.

#### **Parameters**

graph: the graph the matching belongs to

matching: a numeric vector where element i corresponds to vertex i

of the graph. Element i is -1 or if the corresponding vertex is unmatched, otherwise it contains the index of

the vertex to which vertex i is matched.

types: the types of the vertices if the graph is bipartite. It must

either be the name of a vertex attribute (which will be retrieved for all vertices) or a list. Elements in the list will be converted to boolean values True or False, and this will determine which part of the bipartite graph a

given vertex belongs to.

#### Raises

ValueError if the matching vector supplied does not describe a valid matching of the graph.

Overrides: object. init

## $\_\_len\_\_(self)$

 $\_\_\mathbf{repr}\_\_(\mathit{self})$ 

repr(x)

Overrides: object. \_\_repr\_\_ extit(inherited documentation)

\_\_str\_\_(self)

str(x)

Overrides: object.\_\_str\_\_ extit(inherited documentation)

#### edges(self)

Returns an edge sequence that contains the edges in the matching.

If there are multiple edges between a pair of matched vertices, only one of them will be returned.

## is maximal(self)

Returns whether the matching is maximal.

A matching is maximal when it is not possible to extend it any more with extra edges; in other words, unmatched vertices in the graph must be adjacent to matched vertices only.

## is matched(self, vertex)

Returns whether the given vertex is matched to another one.

## $\mathbf{match}$ $\mathbf{of}(self, vertex)$

Returns the vertex a given vertex is matched to.

#### **Parameters**

vertex: the vertex we are interested in; either an integer index or an instance of Vertex.

#### Return Value

the index of the vertex matched to the given vertex, either as an integer index (if *vertex* was integer) or as an instance of Vertex. When the vertex is unmatched, returns None.

## $Inherited\ from\ object$

delattr(	(),	format	()	),g	etattribute	e()	,hash	()	$n, \underline{new}_{-}(),$	
reduce (	),	reduce	ex	(),	setattr	(),	sizeof	(),	subclasshook	()

#### 21.2.2 Properties

Name	Description
graph	Returns the graph corresponding to the
	matching.
matching	Returns the matching vector where element $i$
	contains the ID of the vertex that vertex $i$ is
	matched to.
	The matching vector will contain -1 for
	unmatched vertices.
types	Returns the type vector if the graph is bipartite.
	Element $i$ of the type vector will be False or
	True depending on which side of the bipartite
	graph vertex $i$ belongs to.
	For non-bipartite graphs, this property returns
	None.
Inherited from object	
class	

## 22 Module igraph.operators

Implementation of union, disjoint union and intersection operators. **License:** Copyright (C) 2006-2012 Tamás Nepusz <ntamas@gmail.com> Pázmány Péter sétány 1/a, 1117 Budapest, Hungary

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#### 22.1 Functions

## disjoint union(graphs)

Graph disjoint union.

The disjoint union of two or more graphs is created.

This function 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.

An error is generated if some input graphs are directed and others are undirected.

@param graph: list of graphs. A lazy sequence is not acceptable. @return: the disjoint union graph

#### union(graphs, byname='auto')

Oreturn: the union graph

Graph union.

The union of two or more graphs is created. The graphs may have identical or overlapping vertex sets. Edges which are included in at least one graph will be part of the new graph.

This function 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.

@param graph: list of graphs. A lazy sequence is not acceptable.
@param byname: bool or 'auto' specifying the function behaviour with
 respect to names vertices (i.e. vertices with the 'name' attribute). If
 False, ignore vertex names. If True, merge vertices based on names. If
 'auto', use True if all graphs have named vertices and False otherwise
 (in the latter case, a warning is generated too).

## intersection(graphs, byname='auto', keep\_all\_vertices=True)

Graph intersection.

The intersection of two or more graphs is created. The graphs may have identical or overlapping vertex sets. Edges which are included in all graphs will be part of the new graph.

This function 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.

@param graph: list of graphs. A lazy sequence is not acceptable.
@param byname: bool or 'auto' specifying the function behaviour with
 respect to names vertices (i.e. vertices with the 'name' attribute). If
 False, ignore vertex names. If True, merge vertices based on names. If
 'auto', use True if all graphs have named vertices and False otherwise
 (in the latter case, a warning is generated too).

@param keep\_all\_vertices: bool specifying if vertices that are not in all graphs should be kept in the intersection.

Oreturn: the intersection graph

## 23 Package igraph.remote

Classes that help igraph communicate with remote applications.

## 23.1 Modules

• gephi: Classes that help igraph communicate with Gephi (http://www.gephi.org). (Section 24, p. 387)

## 23.2 Variables

Name	Description
package	Value: None

## 24 Module igraph.remote.gephi

Classes that help igraph communicate with Gephi (http://www.gephi.org). License: Copyright (C) 2006-2012 Tamás Nepusz <ntamas@gmail.com> Pázmány Péter sétány 1/a, 1117 Budapest, Hungary

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## 24.1 Class GephiConnection

Object that represents a connection to a Gephi master server.

#### 24.1.1 Methods

```
\_\_init\_\_(\mathit{self}, \mathit{url} = \texttt{None}, \mathit{host} = \texttt{'127.0.0.1'}, \mathit{port} = \texttt{8080}, \mathit{workspace} = \texttt{1})
```

Constructs a connection to a Gephi master server.

The connection object can be constructed either by specifying the url directly, or by specifying the host, port and workspace arguments. The latter three are evaluated only if url is None; otherwise the url will take precedence.

The url argument does not have to include the operation (e.g., ?operation=updateGraph); the connection will take care of it. E.g., if you wish to connect to workspace 2 in a local Gephi instance on port 7341, the correct form to use for the url is as follows:

```
http://localhost:7341/workspace0
Overrides: object.__init__
```

```
__del__(self)
```

## close(self)

Flushes all the pending operations to the Gephi master server in a single request.

## flush(self)

Flushes all the pending operations to the Gephi master server in a single request.

## write(self, data)

Sends the given raw data to the Gephi streaming master server in an HTTP POST request.

```
repr__(self)
repr(x)
Overrides: object.__repr__ extit(inherited documentation)
```

## Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __new__(), __reduce__(), __reduce_ex__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

#### 24.1.2 Properties

Name	Description
url	The URL of the Gephi workspace where the
	data will be sent.
Inherited from object	
class	

## ${\bf 24.2 \quad Class \; GephiGraph Streaming APIFormat}$

object —

#### igraph.remote.gephi.GephiGraphStreamingAPIFormat

Object that implements the Gephi graph streaming API format and returns Python objects

corresponding to the events defined in the API.

#### **24.2.1** Methods

```
get add node event(self, identifier, attributes={})
```

Generates a Python object corresponding to the event that adds a node with the given identifier and attributes in the Gephi graph streaming API.

## Example:

```
>>> api = GephiGraphStreamingAPIFormat()
>>> api.get_add_node_event("spam")
{'an': {'spam': {}}}
>>> api.get_add_node_event("spam", dict(ham="eggs"))
{'an': {'spam': {'ham': 'eggs'}}}
```

```
{\tt get\_add\_edge\_event}(\mathit{self}, \mathit{identifier}, \mathit{source}, \mathit{target}, \mathit{directed}, \mathit{attributes} {=} \{\})
```

Generates a Python object corresponding to the event that adds an edge with the given source, target, directedness and attributes in the Gephi graph streaming API.

```
{\tt get\_change\_node\_event}(\mathit{self}, \mathit{identifier}, \mathit{attributes})
```

Generates a Python object corresponding to the event that changes the attributes of some node in the Gephi graph streaming API. The given attributes are merged into the existing ones; use C{None} as the attribute value to delete a given attribute.

## Example:

```
>>> api = GephiGraphStreamingAPIFormat()
>>> api.get_change_node_event("spam", dict(ham="eggs"))
{'cn': {'spam': {'ham': 'eggs'}}}
>>> api.get_change_node_event("spam", dict(ham=None))
{'cn': {'spam': {'ham': None}}}
```

## get change edge event(self, identifier, attributes)

Generates a Python object corresponding to the event that changes the attributes of some edge in the Gephi graph streaming API. The given attributes are merged into the existing ones; use C{None} as the attribute value to delete a given attribute.

## Example:

```
>>> api = GephiGraphStreamingAPIFormat()
>>> api.get_change_edge_event("spam", dict(ham="eggs"))
{'ce': {'spam': {'ham': 'eggs'}}}
>>> api.get_change_edge_event("spam", dict(ham=None))
{'ce': {'spam': {'ham': None}}}
```

## get delete node event(self, identifier)

Generates a Python object corresponding to the event that deletes a node with the given identifier in the Gephi graph streaming API.

## Example:

```
>>> api = GephiGraphStreamingAPIFormat()
>>> api.get_delete_node_event("spam")
{'dn': {'spam': {}}}
```

## ${\bf get\_delete\_edge\_event}(\mathit{self}, \mathit{identifier})$

Generates a Python object corresponding to the event that deletes an edge with the given identifier in the Gephi graph streaming API.

## Example:

```
>>> api = GephiGraphStreamingAPIFormat()
>>> api.get_delete_edge_event("spam:ham")
{'de': {'spam:ham': {}}}
```

## Inherited from object

```
__delattr__(), __format__(), __getattribute__(), __hash__(), __init__(), __new__(), __reduce__(), __reduce_ex__(), __repr__(), __setattr__(), __sizeof__(), __str__(), __subclasshook__()
```

#### 24.2.2 Properties

Name	Description
Inherited from object	
class	

## 24.3 Class GephiGraphStreamer

```
\begin{array}{c} \text{object} & \\ \\ \text{igraph.remote.gephi.GephiGraphStreamer} \end{array}
```

Class that produces JSON event objects that stream an igraph graph to Gephi using the Gephi Graph Streaming API.

The Gephi graph streaming format is a simple JSON-based format that can be used to post mutations to a graph (i.e. node and edge additions, removals and updates) to a remote component. For instance, one can open up Gephi (http://www.gephi.org), install the Gephi graph streaming plugin and then send a graph from igraph straight into the Gephi window by using GephiGraphStreamer with the appropriate URL where Gephi is listening.

## Example:

```
>>> from cStringIO import StringIO
>>> from igraph import Graph
>>> buf = StringIO()
>>> streamer = GephiGraphStreamer()
>>> graph = Graph.Formula("A --> B, B --> C")
>>> streamer.post(graph, buf)
>>> print buf.getvalue()  # doctest: +ELLIPSIS, +NORMALIZE_WHITESPACE
{"an": {"igraph:...:v:0": {"name": "A"}}}
{"an": {"igraph:...:v:1": {"name": "B"}}}
{"an": {"igraph:...:v:2": {"name": "C"}}}
{"ae": {"igraph:...:e:0:1": {...}}}
```

#### 24.3.1 Methods

 $_{\rm init}_{\rm \_}(self, encoder={\tt None})$ 

Constructs a Gephi graph streamer that will post graphs to a given file-like object or a Gephi connection.

encoder must either be None or an instance of json. JSONEncoder and it must contain the JSON encoder to be used when posting JSON objects. Overrides: object. init

## iterjsonobj(self, graph)

Iterates over the JSON objects that build up the graph using the Gephi graph streaming API. The objects returned from this function are Python objects; they must be formatted with <code>json.dumps</code> before sending them to the destination.

post(self, graph, destination, encoder=None)

Posts the given graph to the destination of the streamer using the given JSON encoder. When encoder is None, it falls back to the default JSON encoder of the streamer in the encoder property.

destination must be a file-like object or an instance of GephiConnection.

 $\mathbf{send}_{\mathbf{e}}\mathbf{vent}(self,\ event,\ destination,\ encoder = \mathtt{None},\ flush = \mathtt{True})$ 

Sends a single JSON event to the given destination using the given JSON encoder. When encoder is None, it falls back to the default JSON encoder of the streamer in the encoder property.

destination must be a file-like object or an instance of GephiConnection.

The method flushes the destination after sending the event. If you want to avoid this (e.g., because you are sending many events), set flush to False.

#### Inherited from object

\_\_delattr\_\_(), \_\_format\_\_(), \_\_getattribute\_\_(), \_\_hash\_\_(), \_\_new\_\_(), \_\_reduce\_\_(), \_\_reduce\_ex\_\_(), \_\_repr\_\_(), \_\_setattr\_\_(), \_\_sizeof\_\_(),

$$\_\_str\_\_(), \ \_\_subclasshook\_\_()$$

## 24.3.2 Properties

Name	Description
Inherited from object	
class	

## 25 Module igraph.statistics

Statistics related stuff in igraph

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### 25.1 Functions

#### mean(xs)

Returns the mean of an iterable.

Example:

```
>>> mean([1, 4, 7, 11])
5.75
```

#### **Parameters**

xs: an iterable yielding numbers.

#### Return Value

the mean of the numbers provided by the iterable.

See Also: RunningMean() if you also need the variance or the standard deviation

## median(xs, sort=True)

Returns the median of an unsorted or sorted numeric vector.

#### Parameters

xs: the vector itself.

sort: whether to sort the vector. If you know that the vector is sorted already, pass False here.

#### Return Value

the median, which will always be a float, even if the vector contained integers originally.

## percentile(xs, p=(25, 50, 75), sort=True)

Returns the pth percentile of an unsorted or sorted numeric vector.

This is equivalent to calling quantile(xs, p/100.0); see quantile for more details on the calculation.

## Example:

```
>>> round(percentile([15, 20, 40, 35, 50], 40), 2)
26.0
>>> for perc in percentile([15, 20, 40, 35, 50], (0, 25, 50, 75, 100)):
... print "%.2f" % perc
...
15.00
17.50
35.00
45.00
50.00
```

#### **Parameters**

xs: the vector itself.

p: the percentile we are looking for. It may also be a list if you want to calculate multiple quantiles with a single call. The default value calculates the 25th, 50th and 75th percentile.

sort: whether to sort the vector. If you know that the vector is sorted already, pass False here.

#### Return Value

the pth percentile, which will always be a float, even if the vector contained integers originally. If p is a list, the result will also be a list containing the percentiles for each item in the list. power\_law\_fit(data, xmin=None, method='auto',
return alpha only=False)

Fitting a power-law distribution to empirical data

#### **Parameters**

data: the data to fit, a list containing integer values

xmin: the lower bound for fitting the power-law. If None, the optimal xmin value will be estimated as well. Zero means

that the smallest possible xmin value will be used.

method: the fitting method to use. The following methods are implemented so far:

- continuous, hill: exact maximum likelihood estimation when the input data comes from a continuous scale. This is known as the Hill estimator. The statistical error of this estimator is (alpha-1) / sqrt(n), where alpha is the estimated exponent and n is the number of data points above xmin. The estimator is known to exhibit a small finite sample-size bias of order  $O(n^-1)$ , which is small when n > 100. igraph will try to compensate for the finite sample size if n is small.
- discrete: exact maximum likelihood estimation when the input comes from a discrete scale (see Clauset et al among the references).
- auto: exact maximum likelihood estimation where the continuous method is used if the input vector contains at least one fractional value and the discrete method is used if the input vector contains integers only.

### Return Value

a FittedPowerLaw object. The fitted xmin value and the power-law exponent can be queried from the xmin and alpha properties of the returned object.

#### Reference:

- MEJ Newman: Power laws, Pareto distributions and Zipf's law. Contemporary Physics 46, 323-351 (2005)
- A Clauset, CR Shalizi, MEJ Newman: Power-law distributions in empirical data. E-print (2007). arXiv:0706.1062

```
quantile(xs, q=(0.25, 0.5, 0.75), sort=True)
```

Returns the qth quantile of an unsorted or sorted numeric vector.

There are a number of different ways to calculate the sample quantile. The method implemented by igraph is the one recommended by NIST. First we calculate a rank n as q(N+1), where N is the number of items in xs, then we split n into its integer component k and decimal component d. If  $k \le 1$ , we return the first element; if  $k \ge N$ , we return the last element, otherwise we return the linear interpolation between xs[k-1] and xs[k] using a factor d.

### Example:

```
>>> round(quantile([15, 20, 40, 35, 50], 0.4), 2)
26.0
```

#### **Parameters**

xs: the vector itself.

q: the quantile we are looking for. It may also be a list if you want to calculate multiple quantiles with a single call. The default value calculates the 25th, 50th and 75th percentile.

sort: whether to sort the vector. If you know that the vector is sorted already, pass False here.

#### Return Value

the qth quantile, which will always be a float, even if the vector contained integers originally. If q is a list, the result will also be a list containing the quantiles for each item in the list.

## 25.2 Class FittedPowerLaw

```
object — igraph.statistics.FittedPowerLaw
```

Result of fitting a power-law to a vector of samples

### Example:

```
<BLANKLINE>
Log-likelihood = -7.546337
<BLANKLINE>
HO: data was drawn from the fitted distribution
<BLANKLINE>
KS test statistic = 0.213817
p-value = 0.993111
<BLANKLINE>
HO could not be rejected at significance level 0.05
>>> result.alpha
                               # doctest:+ELLIPSIS
2.425828...
>>> result.xmin
3.0
>>> result.continuous
False
25.2.1 Methods
       \_init\_\_(self, continuous, alpha, xmin, L, D, p)
     x. init (...) initializes x; see help(type(x)) for signature
     Overrides: object. init extit(inherited documentation)
      \mathbf{repr} (self)
     repr(x)
     Overrides: object.__repr__ extit(inherited documentation)
     __str__(self)
     str(x)
     Overrides: object.__str__ extit(inherited documentation)
     summary(self, significance=0.05)
     Returns the summary of the power law fit.
     Parameters
         significance: the significance level of the Kolmogorov-Smirnov
                          test used to decide whether the input data could
                          have come from the fitted distribution
     Return Value
```

the summary as a string

## Inherited from object

### 25.2.2 Properties

Name	Description
Inherited from object	
class	

## 25.3 Class Histogram

```
object — igraph.statistics.Histogram
```

Generic histogram class for real numbers

## Example:

```
>>> h = Histogram(5)  # Initializing, bin width = 5

>>> h << [2,3,2,7,8,5,5,0,7,9]  # Adding more items

>>> print h

N = 10, mean +- sd: 4.8000 +- 2.9740

[ 0, 5): **** (4)

[ 5, 10): ****** (6)
```

#### 25.3.1 Methods

```
Initializes the histogram with the given data set.

Parameters
bin_width: the bin width of the histogram.
data: the data set to be used. Must contain real numbers.

Overrides: object.__init__
```

```
add(self, num, repeat=1)
Adds a single number to the histogram.

Parameters
    num: the number to be added
    repeat: number of repeated additions
```

## add many(self, data)

Adds a single number or the elements of an iterable to the histogram.

#### **Parameters**

data: the data to be added

```
\_\_lshift\_\_(self, data)
```

Adds a single number or the elements of an iterable to the histogram.

#### **Parameters**

data: the data to be added

## $\mathbf{clear}(self)$

Clears the collected data

### $\mathbf{bins}(self)$

Generator returning the bins of the histogram in increasing order

#### Return Value

a tuple with the following elements: left bound, right bound, number of elements in the bin

```
plot (self, context, bbox, _, **kwds)
```

Plotting support

 $\textbf{to\_string}(\textit{self}, \textit{max\_width} = \texttt{78}, \textit{show\_bars} = \texttt{True}, \textit{show\_counts} = \texttt{True})$ 

Returns the string representation of the histogram.

#### **Parameters**

max\_width: the maximal width of each line of the string This

value may not be obeyed if it is too small.

show\_bars: specify whether the histogram bars should be shown

show\_counts: specify whether the histogram counts should be

shown. If both *show\_bars* and *show\_counts* are False, only a general descriptive statistics (number of elements, mean and standard deviation) is shown.

```
\_\_\mathbf{str}\_\_(\mathit{self})
```

str(x)

Overrides: object.\_\_str\_\_ extit(inherited documentation)

## Inherited from object

$\_\_$ delattr $\_\_$	$(), \_\_$	$_{ m lormat}$	(), .	$\_\_{ m get}$	tattribu	te(	),has	$\sinh _{-}()$ ,	new_	(),
reduce(	$(), \_\_$	reduce_	_ex	$(), \{-}$	_repr_	_(),	_setattr	(), _	_sizeof_	_(),
subclassho	ok	()								

#### 25.3.2 Properties

Name	Description
n	Returns the number of elements in the
	histogram
mean	Returns the mean of the elements in the
	histogram
sd	Returns the standard deviation of the elements
	in the histogram
var	Returns the variance of the elements in the
	histogram
Inherited from object	
class	

## 25.4 Class RunningMean

$$\begin{array}{c} \text{object} \ \ \, \\ \text{igraph.statistics.} \\ \text{RunningMean} \end{array}$$

Running mean calculator.

This class can be used to calculate the mean of elements from a list, tuple, iterable or any other data source. The mean is calculated on the fly without explicitly summing the values, so it can be used for data sets with arbitrary item count. Also capable of returning the standard deviation (also calculated on the fly)

#### 25.4.1 Methods

```
\_\_init\_\_(items=None, n=0.0, mean=0.0, sd=0.0)
```

Initializes the running mean calculator.

There are two possible ways to initialize the calculator. First, one can provide an iterable of items; alternatively, one can specify the number of items, the mean and the standard deviation if we want to continue an interrupted calculation.

#### **Parameters**

items: the items that are used to initialize the running mean

calcuator. If items is given, n, mean and sd must be zeros.

n: the initial number of elements already processed. If this is

given, items must be None.

mean: the initial mean. If this is given, items must be None.

sd: the initial standard deviation. If this is given, items must

be None.

Overrides: object. init

## add(RunningMean, value, repeat=1)

Adds the given value to the elements from which we calculate the mean and the standard deviation.

#### **Parameters**

value: the element to be added

repeat: number of repeated additions

### add many(RunningMean, values)

Adds the values in the given iterable to the elements from which we calculate the mean. Can also accept a single number. The left shift (<<) operator is aliased to this function, so you can use it to add elements as well:

#### **Parameters**

values: the element(s) to be added

(type=iterable)

```
clear(self)
     Resets the running mean calculator.
      __repr__(self)
     repr(x)
     Overrides: object.__repr__ extit(inherited documentation)
      \_\_\mathbf{str}\_\_(\mathit{self})
     str(x)
     Overrides: object.__str__ extit(inherited documentation)
          lshift
                     (RunningMean, values)
     Adds the values in the given iterable to the elements from which we calculate
     the mean. Can also accept a single number. The left shift (<<) operator is
     aliased to this function, so you can use it to add elements as well:
     >>> rm=RunningMean()
     >>> rm << [1,2,3,4]
     >>> rm.result
                                           # doctest:+ELLIPSIS
      (2.5, 1.290994...)
     Parameters
           values: the element(s) to be added
                      (type=iterable)
          float
                   (self)
          int
                  (self)
                    (self)
          long
          complex
                          (self)
          len
                   (self)
Inherited from object
     \label{lem:condition} $$\_\_delattr\_(), \_\_format\_\_(), \_\_getattribute\_\_(), \_\_hash\_\_(), \_\_new\_\_(), \_\_reduce\_ex\_\_(), \_\_setattr\_\_(), \_\_sizeof\_\_(), \_\_subclasshook\_\_()
```

## 25.4.2 Properties

Name	Description
result	Returns the current mean and standard
	deviation as a tuple
mean	Returns the current mean
sd	Returns the current standard deviation
var	Returns the current variation
Inherited from object	
class	

## 26 Module igraph.summary'

Summary representation of a graph.

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## 26.1 Class GraphSummary

Summary representation of a graph.

The summary representation includes a header line and the list of edges. The header line consists of IGRAPH, followed by a four-character long code, the number of vertices, the number of edges, two dashes (-) and the name of the graph (i.e. the contents of the name attribute, if any). For instance, a header line may look like this:

The four-character code describes some basic properties of the graph. The first character is U if the graph is undirected, D if it is directed. The second letter is W if the graph has a vertex attribute called name, or a dash otherwise. The third letter is W if the graph is weighted (i.e. it has an edge attribute called weight), or a dash otherwise. The fourth letter is B if the graph has a vertex attribute called type; this is usually used for bipartite graphs.

Edges may be presented as an ordinary edge list or an adjacency list. By default, this depends on the number of edges; however, you can control it with the appropriate constructor arguments.

#### 26.1.1 Methods

init (self, graph, verbosity=0, width=78, edge list format='auto', max rows=99999, print graph attributes=False, print vertex attributes=False, print edge attributes=False, full=False) Constructs a summary representation of a graph. **Parameters** the verbosity of the summary. If zero, verbosity: only the header line will be returned. If one, the header line and the list of edges will both be returned. width: the maximal width of each line in the summary. None means that no limit will be enforced. the maximal number of rows to print max\_rows: in a single table (e.g., vertex attribute table or edge attribute table) format of the edge list in the summary. edge\_list\_format: Supported formats are: compressed, adjlist, edgelist, auto, which selects automatically from the other three based on some simple criteria. whether to print graph attributes if print\_graph\_attributes: there are any. print\_vertex\_attributes: whether to print vertex attributes if there are any. print\_edge\_attributes: whether to print edge attributes if there are any. full: False has no effect; True turns on the attribute printing for graph, vertex and edge attributes with verbosity 1. Overrides: object.\_\_init\_\_

\_\_\_str\_\_\_(self)
Returns the summary representation as a string.
Overrides: object.\_\_str\_\_

## Inherited from object

 $\__delattr\_\_(), \_\_format\_\_(), \_\_getattribute\_\_(), \_\_hash\_\_(), \_\_new\_\_(),$ 

reduce	(),	_reduce_	_ex	_(),	_repr_	_(),	_setattr_	(),	$_{ m sizeof}_{ m }$	_()
subclassho	ook	_()								

## 26.1.2 Properties

Name	Description
Inherited from object	
class	

## 27 Module igraph.utils

Utility functions that cannot be categorised anywhere else.

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### 27.1 Functions

named temporary file(\*args, \*\*kwds)

Context manager that creates a named temporary file and returns its name.

All parameters are passed on to tempfile.mkstemp, see its documentation for more info.

## numpy to contiguous memoryview(obj)

Converts a NumPy array or matrix into a contiguous memoryview object that is suitable to be forwarded to the Graph constructor.

This is used internally to allow us to use a NumPy array or matrix directly when constructing a Graph.

Functions Module igraph.utils

```
 \begin{array}{lll} \mathbf{rescale}(values,\ out\_range = \texttt{(0.0,\ 1.0)},\ in\_range = \texttt{None},\ clamp = \texttt{False}, \\ scale = \texttt{None}) \end{array}
```

Rescales a list of numbers into a given range.

out\_range gives the range of the output values; by default, the minimum of the original numbers in the list will be mapped to the first element in the output range and the maximum will be mapped to the second element. Elements between the minimum and maximum values in the input list will be interpolated linearly between the first and second values of the output range.

in\_range may be used to override which numbers are mapped to the first and second values of the output range. This must also be a tuple, where the first element will be mapped to the first element of the output range and the second element to the second.

If clamp is True, elements which are outside the given out\_range after rescaling are clamped to the output range to ensure that no number will be outside out\_range in the result.

If scale is not None, it will be called for every element of values and the rescaling will take place on the results instead. This can be used, for instance, to transform the logarithm of the original values instead of the actual values. A typical use-case is to map a range of values to color identifiers on a logarithmic scale. Scaling also applies to the in\_range parameter if present.

#### Examples:

```
>>> rescale(range(5), (0, 8))
[0.0, 2.0, 4.0, 6.0, 8.0]
>>> rescale(range(5), (2, 10))
[2.0, 4.0, 6.0, 8.0, 10.0]
>>> rescale(range(5), (0, 4), (1, 3))
[-2.0, 0.0, 2.0, 4.0, 6.0]
>>> rescale(range(5), (0, 4), (1, 3), clamp=True)
[0.0, 0.0, 2.0, 4.0, 4.0]
>>> rescale([0]*5, (1, 3))
[2.0, 2.0, 2.0, 2.0, 2.0]
>>> from math import log10
>>> rescale([1, 10, 100, 1000, 10000], (0, 8), scale=log10)
[0.0, 2.0, 4.0, 6.0, 8.0]
>>> rescale([1, 10, 100, 1000, 10000], (0, 4), (10, 1000), scale=log10)
[-2.0, 0.0, 2.0, 4.0, 6.0]
```

Variables Module igraph.utils

```
\mathbf{safemax}(\mathit{iterable},\,\mathit{default}{=}0)
```

Safer variant of max() that returns a default value if the iterable is empty.

## Example:

```
>>> safemax([-5, 6, 4])
6
>>> safemax([])
0
>>> safemax((), 2)
2
```

## safemin(iterable, default=0)

Safer variant of min() that returns a default value if the iterable is empty.

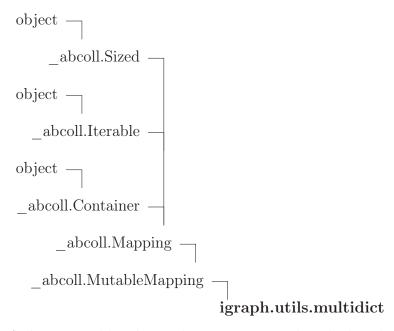
## Example:

```
>>> safemin([-5, 6, 4])
-5
>>> safemin([])
0
>>> safemin((), 2)
```

## 27.2 Variables

Name	Description
dbl_epsilon	Value: 2.22044604925e-16

### 27.3 Class multidict



A dictionary-like object that is customized to deal with multiple values for the same key.

Each value in this dictionary will be a list. Methods which emulate the methods of a standard Python dict object will return or manipulate the first items of the lists only. Special methods are provided to deal with keys having multiple values.

#### **27.3.1** Methods

\_\_contains\_\_(self, key)

Returns whether there are any items associated to the given key. Overrides: abcoll.Container. contains

```
Removes all the items associated to the given key. Overrides:
_abcoll.MutableMapping.__delitem__
```

```
__getitem__(self, key)
```

Returns an arbitrary item associated to the given key. Raises KeyError if no such key exists.

## Example:

```
>>> d = multidict([("spam", "eggs"), ("spam", "bacon")])
>>> d["spam"]
'eggs'
```

Overrides: \_abcoll.Mapping.\_\_getitem\_\_

```
__iter__(self)
```

Iterates over the keys of the multidict. Overrides: \_abcoll.Iterable.\_\_iter\_\_

```
-len_-(self)
```

Returns the number of distinct keys in this multidict. Overrides: \_abcoll.Sized. \_ \_len\_\_

```
__setitem__(self, key, value)
```

Sets the item associated to the given key. Any values associated to the key will be erased and replaced by value.

#### Example:

```
>>> d = multidict([("spam", "eggs"), ("spam", "bacon")])
>>> d["spam"] = "ham"
>>> d["spam"]
'ham'
```

Overrides: \_abcoll.MutableMapping.\_\_setitem\_\_

## add(self, key, value)

Adds value to the list of items associated to key.

## Example:

```
>>> d = multidict()
>>> d.add("spam", "ham")
>>> d["spam"]
'ham'
>>> d.add("spam", "eggs")
>>> d.getlist("spam")
['ham', 'eggs']
```

## clear(self)

Removes all the items from the multidict. Return Value None

Overrides: \_abcoll.MutableMapping.clear

### get(self, key, default=None)

Returns an arbitrary item associated to the given key. If key does not exist or has zero associated items, default will be returned. Return Value D[k] if k in D, else d

Overrides: abcoll.Mapping.get

#### getlist(self, key)

Returns the list of values for the given key. An empty list will be returned if there is no such key.

### iterlists(self)

Iterates over (key, values) pairs where values is the list of values associated with key.

lists	(self)

Returns a list of (key, values) pairs where values is the list of values associated with key.

## update(self, arg, \*\*kwds)

Update D from mapping/iterable E and F. If E present and has a .keys() method, does: for k in E: D[k] = E[k] If E present and lacks .keys() method, does: for (k, v) in E: D[k] = v In either case, this is followed by: for k, v in F.items(): D[k] = v

### Return Value

None

Overrides: \_abcoll.MutableMapping.update extit(inherited documentation)

## $Inherited\ from \quad abcoll. Mutable Mapping$

pop(), popitem(), setdefault()

## $Inherited\ from \quad abcoll. Mapping$

 $_{-eq}(), _{ne}(), _{ne}(), items(), iteritems(), iterkeys(), itervalues(), keys(), values()$ 

## $Inherited\ from\ \_\ abcoll.Sized$

\_\_subclasshook\_\_()

## $Inherited\ from\ object$

#### 27.3.2 Properties

Name	Description			
Inherited from object				
class				

#### 27.3.3 Class Variables

Name	Description			
abstractmethods	Value: frozenset([])			
Inherited from _abcoll.Mapping				
hash				

# 28 Module igraph.version

**Version:** 0.8.3

## 28.1 Variables

Name	Description
version_info	Value: (0, 8, 3)
package	Value: None

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