pypath Documentation

Release 0.7.98

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pypath is a Python package built around igraph to work with molecular network representations e.g. protein, miRNA and drug compound interaction networks.

note pypath supports both Python 2.7 and Python 3.6+. In the beginning, pypath has been developed only for Python 2.7. Then the code have been adjusted to Py3 however we can not guarantee no incompatibilities remained. If you find any method does not work please submit an issue on github. For few years I develop and test pypath in Python 3. Therefore this is the better supported Python variant.

contributions turei.denes@gmail.com

documentation http://pypath.omnipathdb.org/ **issues** https://github.com/saezlab/pypath/issues

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CHAPTER

ONE

INSTALLATION

1.1 Linux

In almost any up-to-date Linux distribution the dependencies of **pypath** are built-in, or provided by the distributors. You only need to install a couple of things in your package manager (cairo, py(2)cairo, igraph, python(2)-igraph, graphviz, pygraphviz), and after install **pypath** by *pip* (see below). If any module still missing, you can install them the usual way by *pip* or your package manager.

1.1.1 igraph C library, cairo and pycairo

python(2)-igraph is a Python interface to use the igraph C library. The C library must be installed. The same goes for cairo, py(2)cairo and graphviz.

1.1.2 Directly from git

pip install git+https://github.com/saezlab/pypath.git

1.1.3 With pip

Download the package from /dist, and install with pip:

pip install pypath-x.y.z.tar.gz

1.1.4 Build source distribution

Clone the git repo, and run setup.py:

python setup.py sdist

1.2 Mac OS X

On OS X installation is not straightforward primarily because cairo needs to be compiled from source. We provide 2 scripts here: the **mac-install-brew.sh** installs everything with HomeBrew, and **mac-install-conda.sh** installs from Anaconda distribution. With these scripts installation of igraph, cairo and graphviz goes smoothly most of the time, and options are available for omitting the 2 latter. To know more see the description in the script header. There is a

third script **mac-install-source.sh** which compiles everything from source and presumes only Python 2.7 and Xcode installed. We do not recommend this as it is time consuming and troubleshooting requires expertise.

1.2.1 Troubleshooting

• no module named ... when you try to load a module in Python. Did

theinstallation of the module run without error? Try to run again the specific part from the mac install shell script to see if any error comes up. Is the path where the module has been installed in your \$PYTHONPATH? Try echo \$PYTHONPATH to see the current paths. Add your local install directories if those are not there, e.g. export PYTHONPATH="/Users/me/local/python2.7/site-packages:\$PYTHONPATH". If it works afterwards, don't forget to append these export path statements to your ~/.bash_profile, so these will be set every time you launch a new shell.

• pkgconfig not found. Check if the \$PKG_CONFIG_PATH variable is

set correctly, and pointing on a directory where pkgconfig really can be found.

• Error while trying to install py(2)cairo by pip. py(2)cairo could not be

installed by pip, but only by waf. Please set the \$PKG_CONFIG_PATH before. See mac-install-source.sh on how to install with waf.

• Error at pygraphviz build: graphviz/cgraph.h file not found. This is

because the directory of graphviz detected wrong by pkgconfig. See **mac-install-source.sh** how to set include dirs and library dirs by --global-option parameters.

• Can not install bioservices, because installation of jurko-suds fails. Ok,

this fails because pip is not able to install the recent version of setuptools, because a very old version present in the system path. The development version of jurko-suds does not require setuptools, so you can install it directly from git as it is done in **mac-install-source.sh**.

• In **Anaconda**, pypath can be imported, but the modules and classes are

missing. Apparently Anaconda has some built-in stuff called *pypath*. This has nothing to do with this module. Please be aware that Anaconda installs a completely separated Python distribution, and does not detect modules in the main Python installation. You need to install all modules within Anaconda's directory. **mac-install-conda.sh** does exactly this. If you still experience issues, please contact us.

1.3 Microsoft Windows

Not many people have used *pypath* on Microsoft computers so far. Please share your experiences and contact us if you encounter any issue. We appreciate your feedback, and it would be nice to have better support for other computer systems.

1.3.1 With Anaconda

The same workflow like you see in mac-install-conda.sh should work for Anaconda on Windows. The only problem you certainly will encounter is that not all the channels have packages for all platforms. If certain channel provides no package for Windows, or for your Python version, you just need to find an other one. For this, do a search:

anaconda search -t conda <package name>

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For example, if you search for *pycairo*, you will find out that *vgauther* provides it for osx-64, but only for Python 3.4, while *richlewis* provides also for Python 3.5. And for win-64 platform, there is the channel of *KristanAmstrong*. Go along all the commands in mac-install-conda.sh, and modify the channel if necessary, until all packages install successfully.

1.3.2 With other Python distributions

Here the basic principles are the same as everywhere: first try to install all external dependencies, after *pip* install should work. On Windows certain packages can not be installed by compiled from source by *pip*, instead the easiest to install them precompiled. These are in our case *fisher*, *lxml*, *numpy* (*mkl* version), *pycairo*, *igraph*, *pygraphviz*, *scipy and statsmodels*. The precompiled packages are available here: http://www.lfd.uci.edu/~gohlke/pythonlibs/. We tested the setup with Python 3.4.3 and Python 2.7.11. The former should just work fine, while with the latter we have issues to be resolved.

1.3.3 Known issues

• "No module fabric available." – or pysftp missing: this is not

important, only certain data download methods rely on these modules, but likely you won't call those at all. * Progress indicator floods terminal: sorry about that, will be fixed soon. * Encoding related exceptions in Python2: these might occur at some points in the module, please send the traceback if you encounter one, and we will fix as soon as possible.

Special thanks to Jorge Ferreira for testing pypath on Windows!

1.3. Microsoft Windows

CHAPTER

TWO

REFERENCE

Main network object.

Parameters

- ncbi_tax_id (int) Optional, 9606 (Homo sapiens) by default. NCBI Taxonomic identifier of the organism from which the data will be downloaded.
- default_name_type (dict) Optional, {'protein': 'uniprot', 'mirna': 'mirbase', 'drug': 'chembl', 'lncrna': 'lncrna-genesymbol'} by default. Contains the default identifier types to which the downloaded data will be converted. If others are used, user may need to provide the format definitions for the conversion tables.
- **copy** (pypath.main.PyPath) Optional, None by default. Other pypath.main. PyPath instance from which the data will be copied.
- mysql (tuple) Optional, (None, 'mapping') by default. Contains the MySQL parameters used by the pypath.mapping module to load the ID conversion tables.
- chembl_mysql (tuple) Optional, (None, 'chembl') by default. Contains the MySQL parameters used by the pypath.mapping module to load the ChEMBL ID conversion tables.
- name (str) Optional, 'unnamed' by default. Session or project name (custom).
- outdir (str) Optional, 'results' by default. Output directory where to store all output files.
- loglevel (str) Optional, 'INFO' by default. Sets the level of the logger. Possible levels are: 'DEBUG', 'INFO', 'WARNING', 'ERROR' or 'CRITICAL'.
- **loops** (bool) Optional, False by default. Determines if self-loop edges are allowed in the graph.

Variables

- adjlist (list) List of [set] containing the adjacency of each node. See PyPath. update_adjlist() method for more information.
- **chemb1** (pypath.chemb1.Chemb1) Contains the ChEMBL data. See pypath. chemb1 module documentation for more information.
- **chembl_mysql** (tuple) Contains the MySQL parameters used by the pypath. mapping module to load the ChEMBL ID conversion tables.

- data (dict) Stores custom loaded interaction and attribute table. See PyPath. read data file() method for more information.
- **db_dict** (*dict*) Dictionary of dictionaries. Outer-level keys are 'nodes' and 'edges', corresponding values are [dict] whose keys are the database sources with values of type [set] containing the edge/node indexes for which that database provided some information.
- **dgraph** (*igraph* . *Graph*) Directed network graph object.
- **disclaimer** (str) Disclaimer text.
- dlabDct (dict) Maps the directed graph node labels [str] (keys) to their indices [int] (values).
- dnodDct (dict) Maps the directed graph node names [str] (keys) to their indices [int] (values).
- **dnodInd** (set) Stores the directed graph node names [str].
- dnodLab (dict) Maps the directed graph node indices [int] (keys) to their labels [str] (values).
- dnodNam (dict) Maps the directed graph node indices [int] (keys) to their names [str] (values).
- **edgeAttrs** (*dict*) Stores the edge attribute names [str] as keys and their corresponding types (e.g.: set, list, str, ...) as values.
- **exp** (pandas.DataFrame) Stores the expression data for the nodes (if loaded).
- **exp_prod** (pandas.DataFrame) Stores the edge expression data (as the product of the normalized expression between the pair of nodes by default). For more details see PyPath.edges_expression().
- **exp_samples** (*set*) Contains a list of tissues as downloaded by ProteomicsDB. See *PyPath.get_proteomicsdb*() for more information.
- **failed_edges** (list) List of lists containing information about the failed edges. Each failed edge sublist contains (in this order): [tuple] with the node IDs, [str] names of nodes A and B, [int] IDs of nodes A and B and [int] IDs of the edges in both directions.
- go (dict) Contains the organism(s) NCBI taxonomy ID as key [int] and pypath.go. GOAnnotation object as value, which contains the GO annotations for the nodes in the graph. See pypath.go.GOAnnotation for more information.
- graph (igraph. Graph) Undirected network graph object.
- **gsea** (*pypath.gsea.GSEA*) Contains the loaded gene-sets from MSigDB. See pypath.gsea.GSEA for more information.
- has_cats (set) Contains the categories (e.g.: resources) [str] loaded in the current network.
- htp (dict) Contains information about high-throughput data of the network for different thresholds [int] (keys). Values are [dict] containing the number of references ('rnum') [int], number of edges ('enum') [int], number of sources ('snum') [int] and list of PMIDs of the most common references above the given threshold ('htrefs') [set].
- labDct (dict) Maps the undirected graph node labels [str] (keys) to their indices [int] (values).

- lists (dict) Contains specific lists of nodes (values) for different categories [str] (keys). These can to be loaded from a file or a resource. Some methods include PyPath. receptor_list() ('rec'), PyPath.druggability_list() ('dgb'), PyPath.kinases_list() ('kin'), PyPath.tfs_list() ('tf'), PyPath.disease_genes_list() ('dis'), PyPath.signaling_proteins_list() ('sig'), PyPath.proteome_list() ('proteome') and PyPath.cancer drivers list() ('cdv').
- loglevel (str) The level of the logger.
- **loops** (bool) Whether if self-loop edges are allowed in the graph.
- mapper (pypath.mapping.Mapper) pypath.mapper.Mapper object for ID conversion and other ID-related operations across resources.
- mutation_samples (list) DEPRECATED
- mysql_conf (tuple) Contains the MySQL parameters used by the pypath. mapping module to load the ID conversion tables.
- name (str) Session or project name (custom).
- ncbi_tax_id (int) NCBI Taxonomic identifier of the organism from which the data will be downloaded.
- negatives (dict) -
- nodDct (dict) Maps the undirected graph node names [str] (keys) to their indices [int] (values).
- **nodInd** (set) Stores the undirected graph node names [str].
- nodLab (dict) Maps the undirected graph node indices [int] (keys) to their labels [str] (values).
- nodNam (dict) Maps the directed graph node indices [int] (keys) to their names [str] (values).
- **outdir** (*str*) Output directory where to store all output files.
- ownlog (pypath.logn.logw) Logger class instance, see pypath.logn.logw for more information.
- palette -
- pathway_types -
- pathways -
- plots -
- proteomicsdb -
- raw_data -
- reflists -
- seq-
- session -
- session_name -
- sourceNetEdges -
- sourceNetNodes -

```
• sources -
```

- u pfam-
- uniprot_mapped -
- unmapped -
- vertexAttrs -

```
acsn effects(graph=None)
```

```
add_genesets (genesets)
```

add_list_eattr(edge, attr, value)

add_update_vertex (defAttrs, originalName, originalNameType, extraAttrs={}, add=False)

Updates the attributes of one node in the network. Optionally it creates a new node and sets the attributes, but it is not efficient as igraph needs to reindex vertices after this operation, so better to create new nodes and edges in batch.

```
affects (identifier)
```

all between (nameA, nameB)

Returns all edges between two given vertex names. Similar to straight_between(), but checks both directions, and returns list of edge ids in [undirected, straight, reversed] format, for both nameA -> nameB and nameB -> nameA edges.

all_neighbours (indices=False)

```
apply_list (name, node_or_edge='node')
```

Creates vertex or edge attribute based on a list.

```
apply_negative (settings)
```

```
attach_network (edgeList=False, regulator=False)
```

Adds edges to the network from edgeList obtained from file or other input method.

```
basic_stats (latex=False, caption=", latex_hdr=True, fontsize=8, font='HelveticaNeueLTStd-
LtCn', fname=None, header_format='%s', row_order=None, by_category=True,
use_cats=['p', 'm', 'i', 'r'], urls=True, annots=False)
```

Returns basic numbers about the network resources, e.g. edge and node counts.

latex Return table in a LaTeX document. This can be compiled by PDFLaTeX: latex stats.tex

```
basic_stats_intergroup(groupA, groupB, header=None)
```

```
cancer_gene_census_list()
```

Loads the list of cancer driver proteins from the COSMIC Cancer Gene Census.

clean_graph()

Removes multiple edges, unknown molecules and those from wrong taxon. Multiple edges will be combined by *combine_attr()* method. Loops will be deleted unless the *loops* attribute set to *True*.

collapse_by_name (graph=None)

Collapses nodes with the same name with copying and merging all edges and attributes.

```
combine_attr (lst, num_method=<built-in function max>)
```

Combines multiple attributes into one. This method attempts to find out which is the best way to combine attributes.

• if there is only one value or one of them is None, then returns the one available

- lists: concatenates unique values of lists
- numbers: returns the greater by default or calls *num_method()* if given.
- sets: returns the union
- dicts: calls *common.merge_dicts()*
- Direction: calls their special merge() method

Works on more than 2 attributes recursively.

Parameters

- 1st (list) List of one or two attribute values.
- num_method (callable) Method to merge numeric attributes.

```
communities (method, **kwargs)
complex_comembership_network (graph=None, resources=None)
complexes (methods=['3dcomplexes', 'havugimana', 'corum', 'complexportal', 'compleat'])
complexes_in_network (csource='corum', graph=None)
compounds_from_chembl (chembl_mysql=None,
                                                   nodes=None.
                                                                   crit=None,
                                                                                 andor='or'.
                                                  F'],
                                                            relationship_types=['D',
                                                                                       'H'],
                            assay\_types=['B']
                            multi_query=False, **kwargs)
consistency()
copy (other)
copy_edges (sources, target, move=False, graph=None)
    Copies edges of one node to another, keeping attributes and directions.
```

Parameters

- **sources** (*list*) Vertex IDs to copy from.
- target (int) Vertex ID to copy for.
- move (bool) Whether perform copy or move, i.e. remove or keep the source edges.

```
count_sol()
```

Counts nodes with zero degree.

```
curation_effort (sum_by_source=False)
```

Returns the total number of reference-interactions pairs.

@sum_by_source [bool] If True, counts the refrence-interaction pairs by sources, and returns the sum of these values.

Removes the proteins of all organisms which are not listed.

```
Parameters tax (list) - List of NCBI Taxonomy IDs of the organisms. E.g. [7227, 9606]
```

```
delete_unknown (tax, typ='protein', defaultNameType=None)
```

Removes those proteins which are not in the list of all default IDs of the organisms. By default, it means to remove all protein nodes not having a human SwissProt ID.

@tax [list] List of NCBI Taxonomy IDs of the organisms of interest. E.g. [7227, 9606]

@typ [str] Molecule type. E.g. 'protein' or 'mirna'

@defaultNameType [str] The default name type of the given molecular species. For proteins it's 'uniprot' by default.

delete_unmapped()

dgenesymbol (genesymbol)

Returns igraph. Vertex() object if the GeneSymbol can be found in the default directed network, otherwise None.

@genesymbol [str] GeneSymbol.

dgenesymbols (genesymbols)

dgs (genesymbol)

Returns igraph. Vertex() object if the GeneSymbol can be found in the default directed network, otherwise None.

@genesymbol [str] GeneSymbol.

dgss (genesymbols)

disease_genes_list (dataset='curated')

Loads the list of all disease related genes from DisGeNet. This resource is human only.

```
dneighbors (identifier, mode='ALL')
```

dp (*identifier*)

Same as PyPath.get_node, just for the directed graph. Returns igraph.Vertex() object if the identifier is a valid vertex index in the default directed graph, or a UniProt ID or GeneSymbol which can be found in the default directed network, otherwise None.

@identifier [int, str] Vertex index (int) or GeneSymbol (str) or UniProt ID (str) or igraph. Vertex object.

dproteins (identifiers)

dps (identifiers)

druggability_list()

Loads the list of druggable proteins from DgiDB. This resource is human only.

duniprot (uniprot)

Same as PyPath.uniprot(), just for directed graph. Returns ``igraph. Vertex() object if the UniProt can be found in the default directed network, otherwise None.

@uniprot [str] UniProt ID.

duniprots (uniprots)

Returns list of igraph. Vertex() object for a list of UniProt IDs omitting those could not be found in the default directed graph.

dup (uniprot)

Same as PyPath.uniprot(), just for directed graph. Returns ``igraph. Vertex() object if the UniProt can be found in the default directed network, otherwise None.

```
@uniprot [str] UniProt ID.
```

```
dups (uniprots)
```

Returns list of igraph. Vertex() object for a list of UniProt IDs omitting those could not be found in the default directed graph.

```
dv (identifier)
```

Same as PyPath.get_node, just for the directed graph. Returns igraph.Vertex() object if the identifier is a valid vertex index in the default directed graph, or a UniProt ID or GeneSymbol which can be found in the default directed network, otherwise None.

@identifier [int, str] Vertex index (int) or GeneSymbol (str) or UniProt ID (str) or igraph. Vertex object.

```
dvs (identifiers)
```

```
edge_exists (nameA, nameB)
```

Returns a tuple of vertice indices if edge doesn't exists, otherwise edge id. Not sensitive to direction.

```
edge_loc (graph=None, topn=2)
edge_names (e)
edges_3d (methods=['dataio.get_instruct', 'dataio.get_i3d'])
edges_expression (func=<function <lambda>>)
```

Executes function *func* for each pairs of connected proteins in the network, for every expression dataset. By default, *func* simply gives the product the (normalized) expression values.

func [callable] Function to handle 2 vectors (pandas.Series() objects), should return one vector of the same length.

```
edges_in_comlexes (csources=['corum'], graph=None)
```

Creates edge attributes <code>complexes</code> and <code>in_complex</code>. These are both dicts where the keys are complex resources. The values in <code>complexes</code> are the list of complex names both the source and the target vertices belong to. The values <code>in_complex</code> are boolean values whether there is at least one complex in the given resources both the source and the target vertex of the edge belong to.

@csources [list] List of complex resources. Should be already loaded.

@graph [igraph.Graph()] The graph object to do the calculations on.

```
edges_ptms()
edgeseq inverse(edges)
```

Builds a pygraphviz.AGraph() object with filtering the edges and vertices along arbitrary criteria. Returns the Agraph object if requested, or exports the dot file, or saves the graphics.

@nodes: list List of vertex ids to be included. @edges: list List of edge ids to be included. @directed: bool Create a directed or undirected graph. @labels: str Name type to be used as id/label in the dot format. @edges_filter: function Function to filter edges, accepting igraph. Edge as argument. @nodes_filter: function Function to filter vertices, accepting igraph. Vertex as argument. @edge_sources: list Sources to be included. @dir_sources: list Direction and effect sources to be included. @graph: igraph. Graph The graph object to export. @return_object: bool Whether to return the pygraphviz. AGraph object. @save_dot: str Filename to export the dot file to. @save_graphics: str Filename to export the graphics, the extension defines the format. @prog: str The graphviz layout algorithm to use. @format: str The graphics format passed to pygraphviz. AGrapg().draw(). @hide: bool Hide filtered edges instead of omit

them. @hide nodes: list Nodes to hide. List of vertex ids. @auto_edges: str Automatic, built-in style for edges. 'DIRECTIONS' or 'RESOURCE_CATEGORIES' are supported. @font: str Font to use for labels. For using more than one fonts refer to graphviz attributes with constant values or define callbacks or mapping dictionaries. @defaults: dict Default values for graphviz attributes, labeled with the entity, e.g. {'edge_penwidth': 0.2}. @**kwargs: constant, callable or dict Graphviz attributes, labeled by the target entity. E.g. edge_penwidth, 'vertex_shape' or graph_label. If the value is constant, this value will be used. If the value is dict, and has _name as key, for every instance of the given entity, the value of the attribute defined by _name will be looked up in the dict, and the corresponding value will be given to the graphviz attribute. If the key _name is missing from the dict, igraph vertex and edge indices will be looked up among the keys. If the value is callable, it will be called with the current instance of the entity and the returned value will be used for the graphviz attribute. E.g. edge_arrowhead(edge) or vertex_fillcolor(vertex) Example:

import pypath from pypath import data_formats net = pypath.PyPath() net.init_network(pfile = 'cache/default.pickle') #net.init_network({'arn': data_formats.omnipath['arn']}) tgf = [v.index for v in net.graph.vs if 'TGF' in v['slk_pathways']] dot = net.export_dot(nodes = tgf, save_graphics = 'tgf_slk.pdf', prog = 'dot',

main_title = 'TGF-beta pathway', return_object = True, label_font = 'HelveticaNeueLT-Std Med Cn', edge_sources = ['SignaLink3'], dir_sources = ['SignaLink3'], hide = True)

```
export_edgelist (fname, graph=None, names=['name'], edge_attributes=[], sep='\t') Write edge list to text file with attributes
```

@param fname: the name of the file or a stream to read from. @param graph: the igraph object containing the network @param names: list with the vertex attribute names to be printed

for source and target vertices

@param edge_attributes: list with the edge attribute names to be printed

@param sep: string used to separate columns

By default UniProt IDs, Gene Symbols, source databases, literature references, directionality and sign information and interaction type are included.

- **outfile** (*str*) Name of the output file. If *None* a file name "netrowk-<session id>.tab" is used.
- extra_node_attrs (dict) Additional node attributes to be included in the exported table. Keys are column ames used in the header while values are names of vertex attributes. In the header _A and _B suffixes will be appended to the column names so the values can be assigned to A and B side interaction partners.
- **extra_edge_attrs** (dict) Additional edge attributes to be included in the exported table. Keys are column ames used in the header while values are names of edge attributes.

filters (line, positiveFilters=[], negativeFilters=[])

find all paths (*start*, *end*, *mode='OUT'*, *maxlen=2*, *graph=None*, *silent=False*)

Finds all paths up to length *maxlen* between groups of vertices. This function is needed only becaues igraph's get_all_shortest_paths() finds only the shortest, not any path up to a defined length.

@start [int or list] Indices of the starting node(s) of the paths.

@end [int or list] Indices of the target node(s) of the paths.

@mode ['IN', 'OUT', 'ALL'] Passed to igraph.Graph.neighbors()

@maxlen [int] Maximum length of paths in steps, i.e. if maxlen = 3, then the longest path may consist of 3 edges and 4 nodes.

@graph [igraph.Graph object] The graph you want to find paths in. self.graph by default.

find_all_paths2 (graph, start, end, mode='OUT', maxlen=2, psize=100)

find_complex(search)

Finds complexes by their non standard names. E.g. to find DNA polymerases you can use the search term *DNA pol* which will be tested against complex names in CORUM.

first_neighbours (node, indices=False)

geneset_enrichment (proteins, all_proteins=None, geneset_ids=None, alpha=0.05, correction method='hommel')

genesymbol (genesymbol)

Returns igraph. Vertex() object if the GeneSymbol can be found in the default undirected network, otherwise None.

@genesymbol [str] GeneSymbol.

```
genesymbol_labels (graph=None, remap_all=False)
```

Creats vertex attribute label and fills up with Gene Symbols of all proteins where the Gene Symbol can be looked up based on the default name of the protein vertex. If the attribute label had been already initialized, updates this attribute or recreates if remap_all is True.

genesymbols (genesymbols)

get_attrs (line, spec, lnum)

get_directed (graph=False, conv_edges=False, mutual=False, ret=False)

Converts graph undirected igraph. Graph object to a directed one. By default it converts the graph in PyPath.graph and places the directed instance in PyPath.dgraph.

@graph [igraph.Graph] Undirected graph object.

@conv_edges [bool] Whether to convert undirected edges (those without explicit direction information) to an arbitrary direction edge or a pair of opposite edges. Otherwise those will be deleted. Default is False.

@mutual [bool] If conv_edges is True, whether to convert the undirected edges to a single, arbitrary directed edge, or a pair of opposite directed edges. Default is False.

@ret [bool] Return the directed graph instance, or return None. Default is False (returns None).

get_dirs_signs()

get_edge (source, target, directed=True)

Returns igraph. Edge object if an edge exist between the 2 proteins, otherwise None.

- **source** (*int*, *str*) Vertex index or UniProt ID or GeneSymbol or igraph. Vertex object.
- target (int, str) Vertex index or UniProt ID or GeneSymbol or igraph. Vertex object.
- **directed** (bool) To be passed to igraph.Graph.get_eid()

```
get_edges (sources, targets, directed=True)
```

Returns a generator with all edges between source and target vertices.

Parameters

- **sources** (*iterable*) Source vertex IDs, names, labels, or any iterable yielding igraph.Vertex objects.
- targets (*iterable*) Target vertec IDs, names, labels, or any iterable yielding igraph. Vertex objects.
- directed (bool) Passed to igraph.get_eid().

```
get_function(fun)
```

```
get_giant (replace=False, graph=None)
```

Returns the giant component of the graph, or replaces the igraph object with only the giant component.

```
get_max (attrList)
```

```
get_network (crit, andor='or', graph=None)
```

```
get_node (identifier)
```

Returns igraph. Vertex() object if the identifier is a valid vertex index in the default undirected graph, or a UniProt ID or GeneSymbol which can be found in the default undirected network, otherwise None.

@identifier [int, str] Vertex index (int) or GeneSymbol (str) or UniProt ID (str) or igraph. Vertex object.

```
get_node_d (identifier)
```

Same as PyPath.get_node, just for the directed graph. Returns igraph.Vertex() object if the identifier is a valid vertex index in the default directed graph, or a UniProt ID or GeneSymbol which can be found in the default directed network, otherwise None.

@identifier [int, str] Vertex index (int) or GeneSymbol (str) or UniProt ID (str) or igraph. Vertex object.

```
get_node_pair (nameA, nameB, directed=False)
get_nodes (identifiers)
get_nodes_d (identifiers)
get_pathways (source)
get_proteomicsdb (user, passwd, tissues=None, pickle=None)
get_sub (crit, andor='or', graph=None)
get_taxon (tax_dict, fields)
go_annotate (aspects=('C', 'F', 'P'))
```

Annotates protein nodes with GO terms. In the go vertex attribute each node is annotated by a dict of sets where keys are one letter codes of GO aspects and values are sets of GO accessions.

```
go_dict (organism=9606)
```

 $Creates\ a\ {\tt pypath.go.GOAnnotation}\ object\ for\ one\ organism\ in\ the\ dict\ under\ {\tt go\ attribute}.$

```
Parameters organism (int) – NCBI Taxonomy ID of the organism.
go_enrichment (proteins=None,
                                   aspect='P',
                                                 alpha=0.05,
                                                               correction method='hommel',
                 all proteins=None)
gs (genesymbol)
    Returns igraph. Vertex() object if the GeneSymbol can be found in the default undirected network,
    otherwise None.
    @genesymbol [str] GeneSymbol.
gs_affected_by (genesymbol)
gs_affects (genesymbol)
gs_edge (source, target, directed=True)
    Returns igraph. Edge object if an edge exist between the 2 proteins, otherwise None.
    @source [str] GeneSymbol
    @target [str] GeneSymbol
    @directed [bool] To be passed to igraph.Graph.get_eid()
gs_in_directed(genesymbol)
gs_in_undirected(genesymbol)
gs_inhibited_by (genesymbol)
gs_inhibits(genesymbol)
gs_neighborhood(genesymbols, order=1, mode='ALL')
gs neighbors (genesymbol, mode='ALL')
gs_stimulated_by (genesymbol)
gs_stimulates (genesymbol)
gss (genesymbols)
guide2pharma()
having_attr (attr, graph=None, index=True, edges=True)
having_eattr (attr, graph=None, index=True)
having_ptm (index=True, graph=None)
having_vattr (attr, graph=None, index=True)
homology_translation (target, source=None, only_swissprot=True, graph=None)
    Translates the current object to another organism by orthology. Proteins without known ortholog will be
    deleted.
        Parameters target (int) - NCBI Taxonomy ID of the target organism. E.g. 10090 for
            mouse.
htp_stats()
in_complex (csources=['corum'])
in directed(vertex)
in_undirected(vertex)
info(name)
init_complex_attr(graph, name)
```

```
init edge attr(attr)
```

Fills edge attribute with its default values, creates lists if in edgeAttrs the attribute is registered as list.

init_gsea(user)

init_network (lst={'arn': <pypath.input_formats.ReadSettings instance at 0x7f015f22bb48>, 'biogrid': <pypath.input formats.ReadSettings instance at 0x7f015f233cb0>, 'ca1': <pypath.input formats.ReadSettings instance at</pre> 0x7f015f22ba28>, 'ccmap': <pypath.input_formats.ReadSettings instance at 0x7f015f22bfc8>, 'cellphonedb': <pypath.input formats.ReadSettings instance at 0x7f015efb5dd0>, 'dbptm': <pypath.input_formats.ReadSettings instance at 0x7f015f233440>, 'death': <pypath.input_formats.ReadSettings instance at 0x7f015f22bcf8>, 'depod': <pypath.input_formats.ReadSettings instance at 0x7f015f233290>, 'dip': <pypath.input_formats.ReadSettings instance at 0x7f015f233098>, <pypath.input_formats.ReadSettings instance at 0x7f015f2333b0>, 'elm': path.input_formats.ReadSettings instance at 0x7f015f233368>, 'guide2pharma': <pypath.input_formats.ReadSettings instance at 0x7f015f22b950>, 'hpmr': path.input_formats.ReadSettings instance at 0x7f015efb5d40>, 'hprd': <pypath.input formats.ReadSettings instance at 0x7f015f233e18>, 'hprd p': <pypath.input_formats.ReadSettings instance at 0x7f015f233488>, 'innatedb': <pypath.input_formats.ReadSettings instance at 0x7f015f233128>, 'intact': <pypath.input_formats.ReadSettings instance at 0x7f015f233c68>, 'lmpid': <pypath.input_formats.ReadSettings instance at 0x7f015f2332d8>, 'macrophage': <pypath.input_formats.ReadSettings instance at 0x7f015f22bcb0>, 'matrixdb': <pypath.input formats.ReadSettings instance at 0x7f015f233200>, 'mppi': <pypath.input formats.ReadSettings instance at 0x7f015f233050>, 'nrf2': <pypath.input_formats.ReadSettings instance at 0x7f015f22bc20>, 'pdz': <*py*path.input_formats.ReadSettings instance at 0x7f015f22bd40>, 'phelm': <pypath.input_formats.ReadSettings instance at 0x7f015f233320>, 'psite': <pypath.input_formats.ReadSettings instance at 0x7f015f233248>, 'ramilowski2015': <pypath.input_formats.ReadSettings instance at 0x7f015efb5e60>, 'signalink3': <pypath.input_formats.ReadSettings instance at 0x7f015f22b908>, <pypath.input_formats.ReadSettings instance at 0x7f015f22bf38>, 'spike': <pypath.input_formats.ReadSettings instance at 0x7f015f22b8c0>, <pypath.input_formats.ReadSettings instance at 0x7f015f22b7e8>}, exclude=[],cache files={}, pfile=False, save=False, reread=False, redownload=False, **kwargs)

This is a lazy way to start the module, load data and build the high confidence, literature curated part of the signaling network.

```
init_vertex_attr(attr)
```

Fills vertex attribute with its default values, creates lists if in vertexAttrs the attribute is registered as list.

```
intergroup_shortest_paths (groupA, groupB, random=False)
```

```
intogen cancer drivers list (intogen file)
```

Loads the list of cancer driver proteins from IntOGen data.

```
jaccard_edges()
jaccard_meta(jedges, critical)
kegg_directions(graph=None)
kegg_pathways(graph=None)
kinase_stats()
```

kinases_list()

Loads the list of all known kinases in the proteome from kinase.com. This resource is human only.

label_by_go (label, go_terms, **kwargs)

Assigns a boolean vertex attribute to nodes which tells whether the node is annotated by all or any (see method parameter of select by go) the GO terms.

laudanna_directions (graph=None)

laudanna_effects (graph=None)

list_resources()

load_3dcomplexes (graph=None)

load_3did_ddi()

load_3did_ddi2 (ddi=True, interfaces=False)

load_3did_dmi()

load_3did_interfaces()

load_all_pathways (graph=None)

load compleat (graph=None)

Loads complexes from Compleat. Loads data into vertex attribute *graph.vs['complexes']['compleat']*. This resource is human only.

load_complexportal(graph=None)

Loads complexes from ComplexPortal. Loads data into vertex attribute graph.vs['complexes']['complexportal']. This resource is human only.

load_comppi (graph=None)

load_corum(graph=None)

Loads complexes from CORUM database. Loads data into vertex attribute graph.vs['complexes']['corum']. This resource is human only.

load_dbptm(non_matching=False, trace=False, **kwargs)

load ddi(ddi)

ddi is either a list of intera. Domain Domain objects, or a function resulting this list

load_ddis (methods=['dataio.get_3dc_ddi', 'dataio.get_domino_ddi', 'self.load_3did_ddi2'])

load_depod_dmi()

load disgenet (dataset='curated', score=0.0, umls=False, full data=False)

Assigns DisGeNet disease-gene associations to the proteins in the network. Disease annotations will be added to the *dis* vertex attribute.

Parameters

- **score** (*float*) Confidence score from DisGeNet. Only associations above the score provided will be considered.
- ulms (bool) By default we assign a list of disease names to each protein. To use Unified Medical Language System IDs instead set this to *True*.
- **full_data** (bool) By default we load only disease names. Set this to *True* if you wish to load additional annotations like number of PubMed IDs, number of SNPs and original sources.

load_dmi (dmi)

dmi is either a list of intera. Domain Motif objects, or a function resulting this list

```
'self.load dbptm',
load_dmis (methods=['self.pfam_regions',
                                                   'self.load depod dmi',
                                          'self.load_pnetworks_dmi',
              'self.load_mimp_dmi',
                                                                           'self.load domino dmi',
              'self.load pepcyber',
                                    'self.load psite reg', 'self.load psite phos',
                                                                                  'self.load ielm',
              'self.load_phosphoelm', 'self.load_elm', 'self.load_3did_dmi'])
load_domino_dmi (organism=None)
load_elm()
load expression(array=False)
     Expression data can be loaded into vertex attributes, or into a pandas DataFrame – the latter offers faster
     ways to process and use these huge matrices.
load_go (aspects=('C', 'F', 'P'))
     Annotates protein nodes with GO terms. In the go vertex attribute each node is annotated by a dict of sets
     where keys are one letter codes of GO aspects and values are sets of GO accessions.
load_havugimana (graph=None)
     Loads complexes from
                                 Havugimana
                                                 2012.
                                                               Loads
                                                                        data
                                                                               into
                                                                                      vertex
                                                                                               attribute
     graph.vs['complexes']['havugimana']. This resource is human only.
load_hpa (normal=True, pathology=True, cancer=True, summarize_pathology=True, tissues=None,
            quality=set(['Supported', 'Approved']), levels=['High': 3, 'Low': 1, 'Medium': 2, 'Not
            detected': 0}, graph=None, na value=0)
     Loads Human Protein Atlas data into vertex attributes.
load_hprd_ptms (non_matching=False, trace=False, **kwargs)
load ielm()
load_interfaces()
```

load_li2012_ptms (non_matching=False, trace=False, **kwargs)

load ligand receptor network (lig rec resources=True, inference from go=True, sources={'arn': <pypath.input_formats.ReadSettings</pre> instance 'ca1': 0x7f015f22bb48>, <pypath.input formats.ReadSettings instance at 0x7f015f22ba28>, 'cellphonedb': <*py*path.input formats.ReadSettings instance 0x7f015efb5dd0>. 'death': <pypath.input formats.ReadSettings instance 0x7f015f22bcf8>, 'guide2pharma': <*py*path.input_formats.ReadSettings instance 0x7f015f22b950>, 'hpmr': <*py*path.input_formats.ReadSettings instance 0x7f015efb5d40>, 'macrophage': <pypath.input_formats.ReadSettings instance 0x7f015f22bcb0>, 'nrf2': <*py*path.input_formats.ReadSettings instance 0x7f015f22bc20>, *'pdz':* <pypath.input formats.ReadSettings instance at 0x7f015f22bd40>, 'ramilowski2015': <pypath.input formats.ReadSettings instance 0x7f015efb5e60>,'signalink3': <pypath.input formats.ReadSettings instance 0x7f015f22b908>, 'signor': <pypath.input formats.ReadSettings instance 0x7f015f22bf38>, 'spike': <pypath.input formats.ReadSettings instance 0x7f015f22b8c0>, 'trip': <pypath.input_formats.ReadSettings instance at 0x7f015f22b7e8>}, keep_undirected=False, keep_rec_rec=False, keep_lig_lig=False)

Initializes a ligand-receptor network.

load_lmpid(method)

load_mappings()

load_mimp_dmi (non_matching=False, trace=False, **kwargs)

load_mutations (attributes=None, gdsc_datadir=None, mutation_file=None)

Mutations are listed in vertex attributes. Mutation() objects offers methods to identify residues and look up in Ptm(), Motif() and Domain() objects, to check if those residues are modified, or are in some short motif or domain.

load_negatives()

 $\begin{tabular}{ll} \begin{tabular}{ll} \beg$

Loads the OmniPath network as it was before August 2016. Furthermore it gives some more options.

 $\begin{tabular}{ll} \textbf{load_omnipath} & (kinase_substrate_extra=False, & remove_htp=True, & htp_threshold=1, \\ & keep_directed=True, min_refs_undirected=2, old_omnipath_resources=False) \\ & \textbf{Loads the OmniPath network}. \end{tabular}$

load_pathways (source, graph=None)

Generic method to load pathway annotations from a resource. We don't recommend calling this method but either specific methods for a single source e.g. $kegg_pathways()$ or $sirnor_pathways()$ or call $load_all_pathways()$ to load all resources.

- **source** (*str*) Name of the source, this need to match a method in the dict in *get_pathways()* method and the edge and vertex attributes with pathway annotations will be called "<source>_pathways".
- graph (igraph. Graph) A graph, by default the default the graph attribute of the current instance.

```
load pdb (graph=None)
load pepcyber()
load_pfam(graph=None)
load_pfam2()
load_pfam3()
load_phospho_dmi (source, trace=False, return_raw=False, **kwargs)
load_phosphoelm(trace=False, **kwargs)
load_pisa(graph=None)
load pnetworks dmi(trace=False, **kwargs)
load_psite_phos (trace=False, **kwargs)
load_psite_reg()
load ptms()
                                                                                homol-
load_ptms2 (input_methods=None,
                                          map by homology from=[9606],
             ogy_only_swissprot=True, ptm_homology_strict=False, nonhuman_direct_lookup=True,
             inputargs = \{\}
```

This is a new method which will replace *load_ptms*. It uses *pypath.ptm.PtmAggregator*, a newly introduced module for combining enzyme-substrate data from multiple resources using homology translation on users demand.

- input_methods (list) Resources to collect enzyme-substrate interactions from. E.g. ['Signor', 'phosphoELM']. By default it contains Signor, PhosphoSitePlus, HPRD, phosphoELM, dbPTM, PhosphoNetworks, Li2012 and MIMP.
- map_by_homology_from (list) List of NCBI Taxonomy IDs of source taxons used for homology translation of enzyme-substrate interactions. If you have a human network and you add here [10090, 10116] then mouse and rat interactions from the source databases will be translated to human.
- homology_only_swissprot (bool) *True* by default which means only SwissProt IDs are accepted at homology translateion, Trembl IDs will be dropped.
- ptm_homology_strict (bool) For homology translation use PhosphoSite's PTM homology table. This guarantees that only truely homologous sites will be included. Otherwise we only check if at the same numeric offset in the homologous sequence the appropriate residue can be find.
- nonhuman_direct_lookup (bool) Fetch also directly nonhuman data from the resources whereever it's available. PhosphoSite contains mouse enzyme-substrate interactions and it is possible to extract these directly beside translating the human ones to mouse.
- inputargs (dict) Additional arguments passed to *PtmProcessor*. A dict can be supplied for each resource, e.g. {'Signor': {...}, 'PhosphoSite': {...}, ...}. Those not used by *PtmProcessor* are forwarded to the *pypath.dataio* methods.

load_reflist (reflist)
load_reflists (reflst=None)
load_resource (settings, clean=True, cache_files={}, reread=False, redownload=False)

load_resources (lst={'arn': <pypath.input_formats.ReadSettings instance at 0x7f015f22bb48>, 'biogrid': <pypath.input formats.ReadSettings instance at 0x7f015f233cb0>, 'ca1': <pypath.input formats.ReadSettings instance at 0x7f015f22ba28>, 'ccmap': <py-</pre> path.input formats.ReadSettings instance at 0x7f015f22bfc8>, 'cellphonedb': <pypath.input formats.ReadSettings instance at 0x7f015efb5dd0>, 'dbptm': <pypath.input_formats.ReadSettings instance at 0x7f015f233440>, 'death': <pypath.input_formats.ReadSettings instance at 0x7f015f22bcf8>, 'depod': <pypath.input formats.ReadSettings instance at 0x7f015f233290>, 'dip': <pypath.input_formats.ReadSettings instance at 0x7f015f233098>, 'domino': <pypath.input formats.ReadSettings instance at 0x7f015f2333b0>, 'elm': <pypath.input_formats.ReadSettings instance at 0x7f015f233368>, 'guide2pharma': <pypath.input_formats.ReadSettings instance at 0x7f015f22b950>, 'hpmr': <py-</pre> path.input_formats.ReadSettings instance at 0x7f015efb5d40>, 'hprd': path.input formats.ReadSettings instance at 0x7f015f233e18>, 'hprd p': <pypath.input formats.ReadSettings instance at 0x7f015f233488>, 'innatedb': <pypath.input formats.ReadSettings instance at 0x7f015f233128>, 'intact': <pypath.input_formats.ReadSettings instance at 0x7f015f233c68>, 'lmpid': <pypath.input formats.ReadSettings instance at 0x7f015f2332d8>, 'macrophage': <pypath.input_formats.ReadSettings instance at 0x7f015f22bcb0>, 'matrixdb': <pypath.input formats.ReadSettings instance at 0x7f015f233200>, 'mppi': <py-</pre> path.input formats.ReadSettings instance at 0x7f015f233050>, 'nrf2': path.input formats.ReadSettings instance at 0x7f015f22bc20>, 'pdz': <*py*path.input_formats.ReadSettings instance at 0x7f015f22bd40>, 'phelm': <pypath.input_formats.ReadSettings instance at 0x7f015f233320>, 'psite': <pypath.input formats.ReadSettings instance at 0x7f015f233248>, 'ramilowski2015': <pypath.input_formats.ReadSettings instance at 0x7f015efb5e60>, 'signalink3': <pypath.input formats.ReadSettings instance at 0x7f015f22b908>, 'signor': <py-</p> path.input_formats.ReadSettings instance at 0x7f015f22bf38>, 'spike': path.input_formats.ReadSettings instance at 0x7f015f22b8c0>, 'trip': <*py*path.input_formats.ReadSettings instance at 0x7f015f22b7e8>}, exclude=[], cache files={}, reread=False, redownload=False)

Loads multiple resources, and cleans up after. Looks up ID types, and loads all ID conversion tables from UniProt if necessary. This is much faster than loading the ID conversion and the resources one by one.

load signor ptms (non matching=False, trace=False, **kwargs)

load_tfregulons (*levels=set*(['A', 'B']), *only_curated=False*) Adds TF-target interactions from TF regulons to the network.

Parameters

- **levels** (*set*) Confidence levels to be used.
- only_curated (bool) Retrieve only literature curated interactions.

TF regulons is a comprehensive resource of TF-target interactions combining multiple lines of evidences: literature curated databases, ChIP-Seq data, PWM based prediction using HOCOMOCO and JASPAR matrices and prediction from GTEx expression data by ARACNe.

For details see https://github.com/saezlab/DoRothEA.

```
>>> import pypath
     >>> pa = pypath.PyPath()
     >>> pa.load_tfregulons(levels = {'A'})
lookup_cache (name, cache_files, int_cache, edges_cache)
loop_edges (index=True, graph=None)
map edge (edge)
     Translates molecule names in dict representing an edge.
map_item(item)
     Translates the name in item representing a molecule.
map_list (lst, singleList=False)
     Only a wrapper for map_edge()
mean_reference_per_interaction()
merge_lists (nameA, nameB, name=None, and_or='and', delete=False, func='max')
    Merges two lists in lists.
merge nodes (nodes, primary=None, graph=None)
     Merges all attributes and all edges of selected nodes and assigns them to the primary node (by default the
     one with lowest ID).
         Parameters
             • nodes (list) - List of edge IDs.
             • primary (int) – ID of the primary edge; if None the lowest ID selected.
mimp_directions (graph=None)
mutated_edges (sample)
     Compares the mutated residues and the modified residues in PTMs. Interactions are marked as mutated if
     the target residue in the underlying PTM is mutated.
names2vids (names)
negative_report (lst=True, outFile=None)
neighborhood (identifiers, order=1, mode='ALL')
neighbors (identifier, mode='ALL')
neighbourhood_network (center, second=False)
network filter (p=2.0)
     This function aims to cut the number of edges in the network, without loosing nodes, to make the network
     less connected, less hairball-like, more usable for analysis.
network stats(outfile=None)
    Calculates basic statistics for the whole network and each of sources. Writes the results in a tab file.
new_edges (edges)
new_nodes (nodes)
node_exists(name)
numof_directed_edges()
numof_reference_interaction_pairs()
numof_references()
```

```
numof undirected edges()
orthology_translation(target, source=None, only_swissprot=True, graph=None)
     Translates the current object to another organism by orthology. Proteins without known ortholog will be
     deleted.
         Parameters target (int) - NCBI Taxonomy ID of the target organism. E.g. 10090 for
            mouse.
p (identifier)
     Returns igraph. Vertex () object if the identifier is a valid vertex index in the default undirected graph,
     or a UniProt ID or GeneSymbol which can be found in the default undirected network, otherwise None.
     @identifier [int, str] Vertex index (int) or GeneSymbol (str) or UniProt ID (str) or igraph. Vertex
         object.
pathway_attributes (graph=None)
pathway_members (pathway, source)
     Returns an iterator with the members of a single pathway. Apart from the pathway name you need to
     supply its source database too.
pathway_names (source, graph=None)
     Returns the names of all pathways having at least one member in the current graph.
pathway_similarity(outfile=None)
pathways table (filename='genes pathways.list', pw sources=['signalink', 'signor', 'netpath',
                    'kegg'], graph=None)
pfam_regions()
phosphonetworks_directions (graph=None)
phosphopoint_directions (graph=None)
phosphorylation_directions()
phosphorylation_signs()
phosphosite_directions (graph=None)
prdb_tissue_expr(tissue, prdb=None, graph=None, occurrence=1, group_function=<function
                       < lambda >> , na_value = 0.0)
process_directions (dirs,
                                         directed=None,
                                                          stimulation=None,
                                                                              inhibition=None,
                                name,
                         graph=None, id_type=None, dirs_only=False)
process_dmi (source, **kwargs)
     This is an universal function for loading domain-motif objects like load_phospho_dmi() for PTMs. TODO
     this will replace load_elm, load_ielm, etc
protein (identifier)
     Same as PyPath.get_node, just for the directed graph. Returns igraph.Vertex() object if the
     identifier is a valid vertex index in the default directed graph, or a UniProt ID or GeneSymbol which can
     be found in the default directed network, otherwise None.
     @identifier [int, str] Vertex index (int) or GeneSymbol (str) or UniProt ID (str) or igraph. Vertex
         object.
protein_edge (source, target, directed=True)
```

Returns igraph. Edge object if an edge exist between the 2 proteins, otherwise None.

- **source** (*int*, *str*) Vertex index or UniProt ID or GeneSymbol or igraph. Vertex object.
- target (int, str) Vertex index or UniProt ID or GeneSymbol or igraph. Vertex object.
- **directed** (bool) To be passed to igraph.Graph.get eid()

```
proteins (identifiers)
```

proteome_list(swissprot=True)

Loads the whole proteome as a list.

ps (identifiers)

```
random_walk_with_return (q, graph=None, c=0.5, niter=1000)
```

Random walk with return (RWR) starting from one or more query nodes. Returns affinity (probability) vector of all nodes in the graph.

param int,list q Vertex IDs of query nodes.

param igraph.Graph graph An igraph.Graph object.

param float c Probability of restart.

param int niter Number of iterations.

$random_walk_with_return2 (q, c=0.5, niter=1000)$

Literally does random walks. Only for testing of the other method, to be deleted later.

```
read_data_file (settings, keep_raw=False, cache_files={}, reread=False, redownload=False)
```

Interaction data with node and edge attributes can be read from simple text based files. This function works not only with files, but with lists as well. Any other function can be written to download a preprocess data, and then give it to this function to finally attach to the network.

@settings [ReadSettings instance] The detailed definition of the input format. Instead of the file name you can give a function name, which will be executed, and the returned data will be used.

@keep_raw [boolean] To keep the raw data read by this function, in order for debugging purposes, or further use.

receptors_list()

Loads the Human Plasma Membrane Receptome as a list. This resource is human only.

```
reference_edge_ratio()
reference_hist (filename=None)
remove_htp (threshold=50, keep_directed=False)
```

```
remove_undirected (min_refs=None)
run batch (methods, toCall=None)
save_network (pfile=None)
save_session()
     Save current state into pickle dump.
search_attr_and(obj, lst)
search_attr_or (obj, lst)
second_neighbours (node, indices=False, with_first=False)
select_by_go (go_terms, go_desc=None, aspects=('C', 'F', 'P'), method='ANY')
     Selects the nodes annotated by certain GO terms.
     Returns set of vertex IDs.
         Parameters method (str) – If ANY nodes annotated with any of the terms returned. If ALL
             nodes annotated with all the terms returned.
separate()
     Separates networks from different sources. Returns dict of igraph objects.
separate_by_category()
     Separate networks based on categories. Returns dict of igraph objects.
sequences (isoforms=True, update=False)
set_boolean_vattr (attr, vids, negate=False)
set_categories()
set_chembl_mysql (title, config_file=None)
     Sets the ChEMBL MySQL config according to title section in config_file ini style config.
     title (str): section title in ini file config_file (str, NoneType): config file name;
         if None, the mysql_config/defaults.mysql will be used
set_disease_genes (dataset='curated')
     Creates a vertex attribute named dis with boolean values True if the protein encoded by a disease related
     gene according to DisGeNet.
         Parameters dataset (str) – Which dataset to use from DisGeNet. Default is curated.
set druggability()
     Creates a vertex attribute dgb with value True if the protein is druggable, otherwise False.
set drugtargets(pchembl=5.0)
     Creates a vertex attribute dtg with value True if the protein has at least one compound binding with affinity
     higher than pchembl, otherwise False.
         Parameters pchembl (float) - Pchembl threshold.
set_kinases()
     Creates a vertex attribute kin with value True if the protein is a kinase, otherwise False.
set_receptors()
     Creates a vertex attribute rec with value True if the protein is a receptor, otherwise False.
set_signaling_proteins()
     Creates a vertex attribute kin with value True if the protein is a kinase, otherwise False.
set tfs(classes=['a', 'b', 'other'])
```

```
set transcription factors (classes=['a', 'b', 'other'])
     Creates a vertex attribute tf with value True if the protein is a transcription factor, otherwise False.
         Parameters classes (list) - Classes to use from TF Census. Default is ['a', 'b', 'other'].
shortest_path_dist(graph=None, subset=None, outfile=None, **kwargs)
     subset is a tuple of two lists if you wish to look for paths between elements of two groups, or a list if you
     wish to look for shortest paths within this group
signaling_proteins_list()
     Compiles a list of signaling proteins (as opposed to other proteins like metabolic enzymes, matrix proteins),
     by looking up a few simple keywords in short description of GO terms.
signor_pathways (graph=None)
similarity_groups (groups, index='simpson')
small_plot (graph, **kwargs)
     This method is deprecated, do not use it.
sorensen_pathways (pwlist=None)
source_diagram(outf=None, **kwargs)
source_network (font='HelveticaNeueLTStd')
     For EMBL branding, use Helvetica Neue Linotype Standard light
source_similarity(outfile=None)
source_stats()
sources_hist()
sources_overlap (diagonal=False)
sources_venn_data (fname=None, return_data=False)
straight_between (nameA, nameB)
     This does actually the same as get_edge(), but by names instead of vertex ids.
string_effects(graph=None)
sum_in_complex (csources=['corum'], graph=None)
     Returns the total number of edges in the network falling between two members of the same complex.
     Returns as a dict by complex resources. Calls :py:func:pypath.pypath.Pypath.edges_in_comlexes() to do
     the calculations.
     @csources [list] List of complex resources. Should be already loaded.
     @graph [igraph.Graph()] The graph object to do the calculations on.
table_latex (fname, header, data, sum_row=True, row_order=None, latex_hdr=True, caption=",
                font='HelveticaNeueLTStd-LtCn', fontsize=8, sum_label='Total', sum_cols=None,
                header_format='%s', by_category=True)
tfs_list()
     Loads the list of all known transcription factors from TF census (Vaquerizas 2009). This resource is human
     only.
third_source_directions (graph=None, use_string_effects=False, use_laudanna_data=False)
     This method calls a series of methods to get additional direction & effect information from sources having
     no literature curated references, but giving sufficient evidence about the directionality for interactions
     already supported by literature evidences from other sources.
```

Returns a network which includes the proteins expressed in certain tissue according to ProteomicsDB.

tissue_network (tissue, graph=None)

Parameters

- **tissue** (*str*) Tissue name as used in ProteomicsDB.
- **graph** (*igraph*. *Graph*) A graph object, by default the *graph* attribute of the current instance.

```
transcription factors()
translate_refsdir(rd, ids)
uniprot (uniprot)
    Returns igraph. Vertex () object if the UniProt can be found in the default undirected network, oth-
    erwise None.
    @uniprot [str] UniProt ID.
uniprots (uniprots)
    Returns list of igraph. Vertex () object for a list of UniProt IDs omitting those could not be found in
    the default undirected graph.
uniq_ptm(ptms)
uniq_ptms()
up (uniprot)
    Returns igraph. Vertex () object if the UniProt can be found in the default undirected network, oth-
    erwise None.
    @uniprot [str] UniProt ID.
up_affected_by (uniprot)
up_affects(uniprot)
up_edge (source, target, directed=True)
    Returns igraph. Edge object if an edge exist between the 2 proteins, otherwise None.
    @source [str] UniProt ID
    @target [str] UniProt ID
    @directed [bool] To be passed to igraph.Graph.get_eid()
up_in_directed(uniprot)
up_in_undirected(uniprot)
up_inhibited_by (uniprot)
up inhibits(uniprot)
up_neighborhood(uniprot, order=1, mode='ALL')
up_neighbors (uniprot, mode='ALL')
up_stimulated_by (uniprot)
up_stimulates(uniprot)
update_adjlist(graph=None, mode='ALL')
    Creates an adjacency list in a list of sets format.
update_attrs()
update_cats()
```

Makes sure that the *has cats* attribute is an up to date set of all categories in the current network.

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```
update_db_dict()
update_pathway_types()
update_pathways()

update_sources()
    Makes sure that the sources attribute is an up to date list of all sources in the current network.

update_vertex_sources()

update_vindex()
    This is deprecated.

update_vname()
    For fast lookup of node names and indexes, these are hold in a list and a dict as well. However, every time
```

ups (uniprots)

affecting node indices.

Returns list of igraph. Vertex() object for a list of UniProt IDs omitting those could not be found in the default undirected graph.

new nodes are added, these should be updated. This function is automatically called after all operations

```
v (identifier)
```

Returns igraph. Vertex () object if the identifier is a valid vertex index in the default undirected graph, or a UniProt ID or GeneSymbol which can be found in the default undirected network, otherwise None.

@identifier [int, str] Vertex index (int) or GeneSymbol (str) or UniProt ID (str) or igraph. Vertex object.

vertex_pathways()

Some resources assignes interactions some others proteins to pathways. This function converts pathway annotations from edge attributes to vertex attributes.

```
vs (identifiers)
vsgs ()
vsup ()
wang_effects (graph=None)
write_table (tbl, outfile, sep="\t", cut=None, colnames=True, rownames=True)
```

Object storing directionality information of an edge. Also includes information about the reverse direction, mode of regulation and sources of that information.

Parameters

class pypath.main.**Direction** (nameA, nameB)

- nameA (str) Name of the source node.
- nameB (str) Name of the target node.

Variables

- dirs (dict) Dictionary containing the presence of directionality of the given edge. Keys are *straight*, *reverse* and 'undirected' and their values denote the presence/absence [bool].
- **negative** (*dict*) Dictionary contianing the presence/absence [bool] of negative interactions for both straight and reverse directions.
- **negative_sources** (*dict*) Contains the resource names [str] supporting a negative interaction on straight and reverse directions.

- **nodes** (list) Contains the node names [str] sorted alphabetically (nameA, nameB).
- **positive** (*dict*) Dictionary contianing the presence/absence [bool] of positive interactions for both straight and reverse directions.
- **positive_sources** (*dict*) Contains the resource names [str] supporting a positive interaction on straight and reverse directions.
- reverse (tuple) Contains the node names [str] in reverse order e.g. (nameB, nameA).
- **sources** (*dict*) Contains the resource names [str] of a given edge for each directionality (straight, reverse and 'undirected'). Values are sets containing the names of those resources supporting such directionality.
- **straight** (tuple) Contains the node names [str] in the original order e.g. (nameA, nameB).

check_nodes (nodes)

Checks if *nodes* is contained in the edge.

Parameters nodes (list) – Or [tuple], contains the names of the nodes to be checked.

Returns (bool) - True if all elements in nodes are contained in the object nodes list.

check param(di)

Checks if di is 'undirected' or contains the nodes of the current edge. Used internally to check that di is a valid key for the object attributes declared on dictionaries.

Parameters di (tuple) – Or [str], key to be tested for validity.

Returns

(bool) - True if di is 'undirected' or a tuple of node names contained in the edge, False otherwise.

consensus_edges()

Infers the consensus edge(s) according to the number of supporting sources. This includes direction and sign.

```
Returns (list) - Contains the consensus edge(s) along with the consensus sign. If there is no major directionality, both are returned. The structure is as follows: ['<source>', '<target>', '<(un) directed>', '<sign>']
```

get_dir (direction, sources=False)

Returns the state (or *sources* if specified) of the given *direction*.

Parameters

- direction (tuple) Or [str] (if 'undirected'). Pair of nodes from which direction information is to be retrieved.
- **sources** (bool) Optional, 'False' by default. Specifies if the sources information of the given direction is to be retrieved instead.

Returns (*bool* or *set*) – (if sources=True). Presence/absence of the requested direction (or the list of sources if specified). Returns None if *direction* is not valid.

get_dirs (src, tgt, sources=False)

Returns all directions with boolean values or list of sources.

- src(str) Source node.
- $\mathsf{tgt}(str)$ Target node.

• sources (bool) - Optional, False by default. Specifies whether to return the sources attribute instead of dirs.

Returns Contains the dirs (or sources if specified) of the given edge.

get_sign (direction, sign=None, sources=False)

Retrieves the sign information of the edge in the given direction. If specified in *sign*, only that sign's information will be retrieved. If specified in *sources*, the sources of that information will be retrieved instead.

Parameters

- **direction** (tuple) Contains the pair of nodes specifying the directionality of the edge from which th information is to be retrieved.
- **sign** (*str*) Optional, None by default. Denotes whether to retrieve the 'positive' or 'negative' specific information.
- **sources** (bool) Optional, False by default. Specifies whether to return the sources instead of sign.

Returns (*list*) – If sign=None containing [bool] values denoting the presence of positive and negative sign on that direction, if sources=True the [set] of sources for each of them will be returned instead. If *sign* is specified, returns [bool] or [set] (if sources=True) of that specific direction and sign.

has_sign (direction=None)

Checks whether the edge (or for a specific *direction*) has any signed information (about positive/negative interactions).

Parameters direction (tuple) – Optional, None by default. If specified, only the information of that direction is checked for sign.

Returns

(bool) - True if there exist any information on the sign of the interaction, False otherwise.

is_directed()

Checks if edge has any directionality information.

Returns (bool) - Returns True```if any of the :py:attr:`dirs` attribute values is ``True (except 'undirected'), False otherwise.

is inhibition(direction=None)

Checks if any (or for a specific *direction*) interaction is inhibition (negative interaction).

Parameters direction (*tuple*) – Optional, None by default. If specified, checks the negative attribute of that specific directionality. If not specified, checks both.

Returns (bool) – True if any interaction (or the specified *direction*) is inhibitory (negative).

is_stimulation(direction=None)

Checks if any (or for a specific direction) interaction is activation (positive interaction).

Parameters direction (tuple) – Optional, None by default. If specified, checks the positive attribute of that specific directionality. If not specified, checks both.

Returns (bool) – True if any interaction (or the specified direction) is activatory (positive).

majority_dir()

Infers which is the major directionality of the edge by number of supporting sources.

Returns (*tuple*) – Contains the pair of nodes denoting the consensus directionality. If the number of sources on both directions is equal, None is returned. If there is no directionality information, 'undirected'` will be returned.

majority_sign()

Infers which is the major sign (activation/inhibition) of the edge by number of supporting sources on both directions.

Returns (*dict*) – Keys are the node tuples on both directions (straight/reverse) and values can be either None if that direction has no sign information or a list of two [bool] elements corresponding to majority of positive and majority of negative support. In case both elements of the list are True, this means the number of supporting sources for both signs in that direction is equal.

merge (other)

Merges current edge with another (if and only if they are the same class and contain the same nodes). Updates the attributes dirs, sources, positive, negative, positive_sources and negative_sources.

Parameters other (pypath.main.Direction) - The new edge object to be merged with the current one.

negative_reverse()

Checks if the reverse directionality is a negative interaction.

Returns (*bool*) – True if there is supporting information on the reverse direction of the edge as inhibition. False otherwise.

negative sources reverse()

Retrieves the list of sources for the reverse direction and negative sign.

Returns (*set*) – Contains the names of the sources supporting the reverse directionality of the edge with a negative sign.

${\tt negative_sources_straight}\;(\;)$

Retrieves the list of sources for the straight direction and negative sign.

Returns (*set*) – Contains the names of the sources supporting the straight directionality of the edge with a negative sign.

negative_straight()

Checks if the straight directionality is a negative interaction.

Returns (*bool*) – True if there is supporting information on the straight direction of the edge as inhibition. False otherwise.

positive reverse()

Checks if the reverse directionality is a positive interaction.

Returns (*bool*) – True if there is supporting information on the reverse direction of the edge as activation. False otherwise.

positive_sources_reverse()

Retrieves the list of sources for the reverse direction and positive sign.

Returns (*set*) – Contains the names of the sources supporting the reverse directionality of the edge with a positive sign.

positive_sources_straight()

Retrieves the list of sources for the straight direction and positive sign.

Returns (*set*) – Contains the names of the sources supporting the straight directionality of the edge with a positive sign.

positive_straight()

Checks if the straight directionality is a positive interaction.

Returns (*bool*) — True if there is supporting information on the straight direction of the edge as activation. False otherwise.

reload()

Reloads the object from the module level.

set dir(direction, source)

Adds directionality information with the corresponding data source named. Modifies self attributes dirs and sources.

Parameters

- **direction** (tuple) Or [str], the directionality key for which the value on dirs has to be set True.
- **source** (set) Contains the name(s) of the source(s) from which such information was obtained.

set sign (direction, sign, source)

Sets sign and source information on a given direction of the edge. Modifies the attributes positive and positive_sources or negative and negative_sources depending on the sign. Direction is also updated accordingly, which also modifies the attributes dirs and sources.

Parameters

- **direction** (*tuple*) Pair of edge nodes specifying the direction from which the information is to be set/updated.
- **sign** (*str*) Specifies the type of interaction. If 'positive', is considered activation, otherwise, is assumed to be negative (inhibition).
- **source** (set) Contains the name(s) of the source(s) from which the information was obtained.

sources_reverse()

Retrieves the list of sources for the reverse direction.

Returns (*set*) – Contains the names of the sources supporting the reverse directionality of the edge.

sources_straight()

Retrieves the list of sources for the straight direction.

Returns (*set*) – Contains the names of the sources supporting the straight directionality of the edge.

sources_undirected()

Retrieves the list of sources without directed information.

Returns (*set*) – Contains the names of the sources supporting the edge presence but without specific directionality information.

src (undirected=False)

Returns the name(s) of the source node(s) for each existing direction on the interaction.

Parameters undirected (bool) – Optional, False by default.

Returns (*list*) – Contains the name(s) for the source node(s). This means if the interaction is bidirectional, the list will contain both identifiers on the edge. If the interaction is undirected, an empty list will be returned.

src by source(source)

Returns the name(s) of the source node(s) for each existing direction on the interaction for a specific source.

Parameters source (str) – Name of the source according to which the information is to be retrieved.

Returns (*list*) – Contains the name(s) for the source node(s) according to the specified *source*. This means if the interaction is bidirectional, the list will contain both identifiers on the edge. If the specified *source* is not found or invalid, an empty list will be returned.

tgt (undirected=False)

Returns the name(s) of the target node(s) for each existing direction on the interaction.

Parameters undirected (bool) - Optional, False by default.

Returns (*list*) – Contains the name(s) for the target node(s). This means if the interaction is bidirectional, the list will contain both identifiers on the edge. If the interaction is undirected, an empty list will be returned.

tgt_by_source(source)

Returns the name(s) of the target node(s) for each existing direction on the interaction for a specific source.

Parameters source (str) – Name of the source according to which the information is to be retrieved.

Returns (*list*) – Contains the name(s) for the target node(s) according to the specified *source*. This means if the interaction is bidirectional, the list will contain both identifiers on the edge. If the specified *source* is not found or invalid, an empty list will be returned.

translate (ids)

Translates the node names/identifiers according to the dictionary ids.

Parameters ids (dict) – Dictionary containing (at least) the current names of the nodes as keys and their translation as values.

Returns (pypath.main.Direction) – The copy of current edge object with translated node names.

unset_dir(direction, source=None)

Removes directionality and/or source information of the specified *direction*. Modifies attribute dirs and sources.

Parameters

- direction (tuple) Or [str] (if 'undirected') the pair of nodes specifying the directionality from which the information is to be removed.
- **source** (*set*) Optional, None by default. If specified, determines which specific source(s) is(are) to be removed from sources attribute in the specified *direction*.

unset_sign (direction, sign, source=None)

Removes sign and/or source information of the specified *direction* and *sign*. Modifies attribute positive and positive_sources or negative and negative_sources (or positive_attributes/negative_sources only if source=True).

Parameters

- **direction** (tuple) The pair of nodes specifying the directionality from which the information is to be removed.
- sign(str) Sign from which the information is to be removed. Must be either 'positive' or 'negative'.

• **source** (*set*) – Optional, None by default. If specified, determines which source(s) is(are) to be removed from the sources in the specified *direction* and *sign*.

which_dirs()

Returns the pair(s) of nodes for which there is information about their directionality.

Returns (*list*) – List of tuples containing the nodes for which their attribute dirs is True.

class pypath.main.AttrHelper(value, name=None, defaults={})
 Attribute helper class.

• Initialization arguments:

- value [dict/str]?:
- name [str]?: Optional, None by default.
- *defaults* [dict]:

• Attributes:

- value [dict]?:
- *name* [str]?:
- defaults [dict]:
- *id_type* [type]:

• Call arguments:

- instance []:
- thisDir [tuple?]: Optional, None by default.
- thisSign []: Optional, None by default.
- this Dir Sources []: Optional, None by default.
- this Sources []: Optional, None by default.

• Returns:

_

CHAPTER

THREE

WEBSERVICE

New webservice from 14 June 2018: the queries slightly changed, have been largely extended. See the examples below.

One instance of the pypath webservice runs at the domain http://omnipathdb.org/, serving not only the OmniPath data but other datasets: TF-target interactions from TF Regulons, a large collection additional enzyme-substrate interactions, and literature curated miRNA-mRNA interacions combined from 4 databases. The webservice implements a very simple REST style API, you can make requests by HTTP protocol (browser, wget, curl or whatever).

The webservice currently recognizes 3 types of queries: interactions, ptms and info. The query types resources, network and about have not been implemented yet in the new webservice.

3.1 Mouse and rat

Except the miRNA interactions all interactions are available for human, mouse and rat. The rodent data has been translated from human using the NCBI Homologene database. Many human proteins have no known homolog in rodents hence rodent datasets are smaller than their human counterparts. Note, if you work with mouse omics data you might do better to translate your dataset to human (for example using the pypath.homology module) and use human interaction data.

3.2 Examples

A request without any parameter, gives some basic numbers about the actual loaded dataset:

http://omnipathdb.org

The info returns a HTML page with comprehensive information about the resources:

http://omnipathdb.org/info

The interactions query accepts some parameters and returns interactions in tabular format. This example returns all interactions of EGFR (P00533), with sources and references listed.

http://omnipathdb.org/interactions/?partners=P00533&fields=sources,references

By default only the OmniPath dataset used, to query the TF Regulons or add the extra enzyme-substrate interactions you need to set additional parameters. For example to query the transcriptional regulators of EGFR:

http://omnipathdb.org/interactions/?targets=EGFR&types=TF

The TF Regulons database assigns confidence levels to the interactions. You might want to select only the highest confidence, *A* category:

http://omnipathdb.org/interactions/?targets=EGFR&types=TF&tfregulons_levels=A

Show the transcriptional targets of Smad2 homology translated to rat including the confidence levels from TF Regulons:

http://omnipathdb.org/interactions/?genesymbols=1&fields=type,ncbi_tax_id,tfregulons_level& organisms=10116&sources=Smad2&types=TF

Query interactions from PhosphoNetworks which is part of the kinaseextra dataset:

http://omnipathdb.org/interactions/?genesymbols=1&fields=sources&databases=PhosphoNetworks&datasets=kinaseextra

Get the interactions from Signor, SPIKE and SignaLink3:

http://omnipathdb.org/interactions/?genesymbols=1&fields=sources,references&databases=Signor, SPIKE,SignaLink3

All interactions of MAP1LC3B:

http://omnipathdb.org/interactions/?genesymbols=1&partners=MAP1LC3B

By default partners queries the interaction where either the source or the arget is among the partners. If you set the source_target parameter to AND both the source and the target must be in the queried set:

http://omnipathdb.org/interactions/?genesymbols=1&fields=sources,references&sources=ATG3, ATG7,ATG4B,SQSTM1&targets=MAP1LC3B,MAP1LC3A,MAP1LC3C,Q9H0R8,GABARAP, GABARAPL2&source_target=AND

As you see above you can use UniProt IDs and Gene Symbols in the queries and also mix them. Get the miRNA regulating NOTCH1:

http://omnipathdb.org/interactions/?genesymbols=1&fields=sources,references&datasets=mirnatarget&targets=NOTCH1

Note: with the exception of mandatory fields and genesymbols, the columns appear exactly in the order you provided in your query.

Another query type available is ptms which provides enzyme-substrate interactions. It is very similar to the interactions:

http://omnipathdb.org/ptms?genesymbols=1&fields=sources,references,isoforms&enzymes=FYN

Is there any ubiquitination reaction?

http://omnipathdb.org/ptms?genesymbols=1&fields=sources,references&types=ubiquitination

And acetylation in mouse?

http://omnipathdb.org/ptms?genesymbols=1&fields=sources,references&types=acetylation&organisms=10090

Rat interactions, both directly from rat and homology translated from human, from the PhosphoSite database:

http://omnipathdb.org/ptms?genesymbols=1&fields=sources,references&organisms=10116&databases=PhosphoSite_NosphoSite_noref

FOUR

RELEASE HISTORY

Main improvements in the past releases:

4.1 0.1.0

• First release of pypath, for initial testing.

4.2 0.2.0

- Lots of small improvements in almost every module
- Networks can be read from local files, remote files, lists or provided by any function
- · Almost all redistributed data have been removed, every source downloaded from the original provider.

4.3 0.3.0

• First version with partial Python 3 support.

4.4 0.4.0

- pyreact module with BioPaxReader and PyReact classes added
- Process description databases, BioPax and PathwayCommons SIF conversion rules are supported
- Format definitions for 6 process description databases included.

4.5 0.5.0

- Many classes have been added to the **plot** module
- All figures and tables in the manuscript can be generated automatically
- This is supported by a new module, analysis, which implements a generic

workflow in its Workflow class.

4.6 0.7.74

• homology module: finds the homologs of proteins using the NCBI

Homologene database and the homologs of PTM sites using UniProt sequences and PhosphoSitePlus homology table * ptm module: fully integrated way of processing enzyme-substrate interactions from many databases and their translation by homology to other species * export module: creates pandas.DataFrame or exports the network into tabular file * New webservice * TF Regulons database included and provides much more comprehensive transcriptional regulation resources, including literature curated, in silico predicted, ChIP-Seq and expression pattern based approaches * Many network resources added, including miRNA-mRNA and TF-miRNA interactions

4.7 Upcoming

- · New, more flexible network reader class
- Full support for multi-species molecular interaction networks

(e.g. pathogene-host) * Better support for not protein only molecular interaction networks (metabolites, drug compounds, RNA) * ChEMBL webservice interface, interface for PubChem and eventually forDrugBank * Silent mode: a way to suppress messages and progress bars

CHAPTER

FIVE

FEATURES

The primary aim of **pypath** is to build up networks from multiple sources on one igraph object. **pypath** handles ambiguous ID conversion, reads custom edge and node attributes from text files and **MySQL**.

Submodules perform various features, e.g. graph visualization, working with rug compound data, searching drug targets and compounds in **ChEMBL**.

5.1 ID conversion

The ID conversion module mapping can be used independently. It has the feature to translate secondary UniProt IDs to primaries, and Trembl IDs to SwissProt, using primary Gene Symbols to find the connections. This module automatically loads and stores the necessary conversion tables. Many tables are predefined, such as all the IDs in UniProt mapping service, while users are able to load any table from file or MySQL, using the classes provided in the module input_formats.

5.2 Pathways

pypath includes data and predefined format descriptions for more than 25 high quality, literature curated databases. The inut formats are defined in the data_formats module. For some resources data downloaded on the fly, where it is not possible, data is redistributed with the module. Descriptions and comprehensive information about the resources is available in the descriptions module.

5.3 Structural features

One of the modules called intera provides many classes for representing structures and mechanisms behind protein interactions. These are Residue (optionally mutated), Motif, Ptm, Domain, DomainMotif, DomainDomain and Interface. All these classes have __eq__() methods to test equality between instances, and also __contains__() methods to look up easily if a residue is within a short motif or protein domain, or is the target residue of a PTM.

5.4 Sequences

The module seq contains a simple class for quick lookup any residue or segment in **UniProt** protein sequences while being aware of isoforms.

5.5 Tissue expression

For 3 protein expression databases there are functions and modules for downloading and combining the expression data with the network. These are the Human Protein Atlas, the ProteomicsDB and GIANT. The giant and proteomicsdb modules can be used also as stand alone Python clients for these resources.

5.6 Functional annotations

GSEA and **Gene Ontology** are two approaches for annotating genes and gene products, and enrichment analysis technics aims to use these annotations to highlight the biological functions a given set of genes is related to. Here the enrich module gives abstract classes to calculate enrichment statistics, while the go and the gsea modules give access to GO and GSEA data, and make it easy to count enrichment statistics for sets of genes.

5.7 Drug compounds

UniChem submodule provides an interface to effectively query the UniChem service, use connectivity search with custom settings, and translate SMILEs to ChEMBL IDs with ChEMBL web service.

ChEMBL submodule queries directly your own ChEMBL MySQL instance, has the features to search targets and compounds from custom assay types and relationship types, to get activity values, binding domains, and action types. You need to download the ChEMBL MySQL dump, and load into your own server.

5.8 Technical

MySQL submodule helps to manage MySQL connections and track queries. It is able to run queries parallely to optimize CPU and memory usage on the server, handling queues, and serve the result by server side or client side storage. The chembl and potentially the mapping modules rely on this mysql module.

The most important function in module dataio is a very flexible **download manager** built around curl. The function dataio.curl() accepts numerous arguments, tries to deal in a smart way with local **cache**, authentication, redirects, uncompression, character encodings, FTP and HTTP transactions, and many other stuff. Cache can grow to several GBs, and takes place in ./cache by default. Please be aware of this, and use for example symlinks in case of using multiple working directories.

A simple webservice comes with this module: the server module based on twisted.web.server opens a custom port and serves plain text tables over HTTP with REST style querying.

CHAPT	ER
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OMNIPATH IN R

You can download the data from the webservice and load into R. Look here for an example.

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