pymfinder: Tool Guide

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General information

Description: pymfinder is a Python package designed to detect motifs in complex networks and define the roles of nodes and links using these motifs. Both weighted and binary networks can be analyzed. At its core, pymfinder is a combination of Python methods for network-motif analysis as well as a Python wrapper for the original mfinder version 1.2 written in C and available at http://www.weizmann.ac.il/mcb/UriAlon/. This code has been included and modified here with the explicit consent of Nadav Kashtan, the author of mfinder 1.2.

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Platforms: Windows, Linux, Mac OSX. Following recommendations for mfinder, large and dense networks (>10 000 nodes) require a computer with at least 512 Mbyte RAM in order to calculate motif frequencies. Calculating node or link roles will require greater resources.

How to use pymfinder

Download and installation

Download

The *pymfinder* package can be downloaded from Github (url here) or PLOS Computational Biology.

Installation

Installation within a command-line terminal should be straightforward using the function 'setup.py' included in the *pymfinder* package. After navigating to the directory containing the package, run:

```
python setup.py install
```

If an error message of 'Permission denied' or similar is returned, run:

```
python setup.py install --user
```

This will install *pymfinder* locally rather than in the global Python site-packages or dist-packages directory.

Checking installation

After installation, running the included test suite is strongly encouraged. This may be accomplished by running:

```
python setup.py test
```

Basic usage

Input file format

Input network file format should be in simple '.txt' format. Species names may be given as text or integers but should **not** include spaces. Each edge should be represented by a line of the following format:

```
<source node><target node>
```

Example:

```
1 2
3 1
Salmo_trutta midge
Corvus_corax Salmo_trutta
```

If interaction strengths are known, they can be passed to *pymfinder* in the input file. In this case, each edge should be represented by a line with the format:

```
<source node><target node><interaction strength>
```

Example:

```
1 2 1
3 1 2.5
Salmo_trutta midge 0.005
Corvus_corax Salmo_trutta 3
```

Function call and arguments

All of the functions within *pymfinder* can be called using the same framework. Within a Python environment, first import the *pymfinder* package using, for example:

```
import pymfinder as py
```

The motif structure, motif participation, and motif roles for the network can then be calculated simultaneously using:

The *pymfinder* function call includes the following arguments:

- network: Network path. Must be in quotation marks. No default given.
- links: Determines whether or not to calculate statistics for links as well as nodes. If links=True, link participation and roles will be calculated. Defaults to links=False.
- motifsize: Size of motifs to be calculated. Defined for motifsize=3 for unipartite networks and 3\leq motifsize\leq 6 for bipartite networks. Defaults to motifsize=3. There are 13 possible three-species motifs for unipartite networks (Fig. 1). For bipartite networks, there are only four three-species motifs (Fig. 2) and a larger motif size may be necessary.
- **stoufferIDs**: Determines whether to label motifs following Stouffer et al. (1) or based on the representation of the adjacency matrix of the motif as a binary integer, following the original *mfinder*. If **stoufferIDs=True**, labels will be as in Stouffer et al. (1). Defaults to **stoufferIDs=False**.
- allmotifs: If true, displays results for all possible motifs regardless of whether all have been observed. If false, displays only results for motifs observed in the network. Defaults to allmotifs=False.
- **nrandomizations**: Number of random networks with which to compare the observed network. Defaults to 0 (no randomizations performed).
- randomize: Determines whether to conduct the randomization analysis. If false, no randomizations will be conducted regardless of the value given to **nrandomizations**. Defaults to **randomize=False**.
- **usemetropolis**: If randomizations are to be performed, determines whether to use the Metropolis algorithm. If Metropolis is not used, *pymfinder* uses an MCMC algorithm to shuffle the original network while preserving in- and out-degrees of nodes. Defaults to **usemetropolis=False** (MCMC-based randomizations).

• **networktype**: Indicates whether the network is unipartite (all species may interact with all other species) or bipartite (species are divided into two groups and may interact between groups but not within a group). Defaults to **networktype="unipartite"**.

Functions

The pymfinder function call references three subordinate functions: motif_structure, motif_participation, and motif_roles. Each subordinate function may also be called independently if the full output from pymfinder is not required. When pymfinder is called, the subordinate functions are run in order (motif_structure then motif_participation then motif_roles). If a subordinate function is called directly, any preceding function will also be called. That is, calling motif_structure returns only the motif structure output but calling motif_participation returns the motif participation and motif structure output. Note that the three functions differ in the way in which they handle interaction weights.

motif_structure calculates the motif profile of the network. Arguments passable to motif_structure are the same as those for pymfinder, except that the links argument is not relevant. The same motif profile will be returned whether or not links=True. [[how are weights handled?]]

motif_participation calculates the motif participation for each node (and each link, if links=True). All arguments passable to motif_participation are the same as those passed to pymfinder.[[how are weights handled?]]

motif_roles calculates the role of each node (and each link, if links=True). All arguments passable to motif_roles are the same as those passed to pymfinder.[[how are weights handled?]]

Output

The object 'results' returned by *pymfinder* is a NetworkStats object containing dictionaries of motifs, nodes, and links as well as the network type (unipartite or bipartite) and the size of motifs used. The value for each motif in the .motifs dictionary is an Motif object containing the motif profile for that motif. Similarly, the value for each node or link in the .nodes or .links dictionaries is a NodeLink object containing the motif participation or role of that node or link.

These results can be collected into text-formatted tables and may be seen using:

print results

or written to a file using:

```
f=open('filename','w')
f.write(str(results))
f.close()
```

If links=False, the results include three tables. The first presents the motif profile of the network. Each row gives a motif ID, the count of that motif in the observed network, the mean and standard deviation of the count of that motif in the randomized networks, and the Z-score comparing the real network to the randomized networks. If randomizations=False, the random mean and standard deviations will be reported as 0.000 and the Z-score will be given as 888888.000.

The second table presents the motif participation for each node in the network. Each row gives a node ID and the number of times the node appears in each motif. Motif ID's are given in the first row.

The third table presents the role for each node in the network. Each row gives a node ID and the number of times the node appears in each position in each motif. Node positions are labeled using the following notation: (motif ID, number of predators/out links, number of prey/in links).

If links=True, the results will also contain tables presenting links' motif participation and roles. Links' motif participation follows nodes' motif participation and link roles follow node roles. In both cases, the format of the output table echoes that of the node tables. The only notable difference is in the labeling of link positions. Rather than (motif ID, number of out links, number of in links), link positions are labeled (motif ID, (out links for node 1, in links for node 1), (out links for node 2, in links for node 2)) where nodes 1 and 2 are the two species connected by link (1,2).

Figures

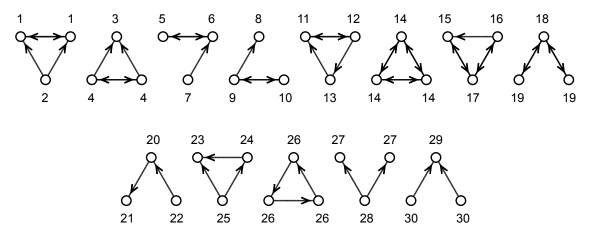


Figure 1: The thirteen three-species motifs in unipartite networks. The 30 unique positions are numbered, but note that positions are not in the same order as in the output provided by motif_roles. [[Recommend we change one or the other, so that they match]] Note also that some motifs contain three unique positions (e.g., positions 5, 6, and 7) while other motifs contain only one or two unique positions.

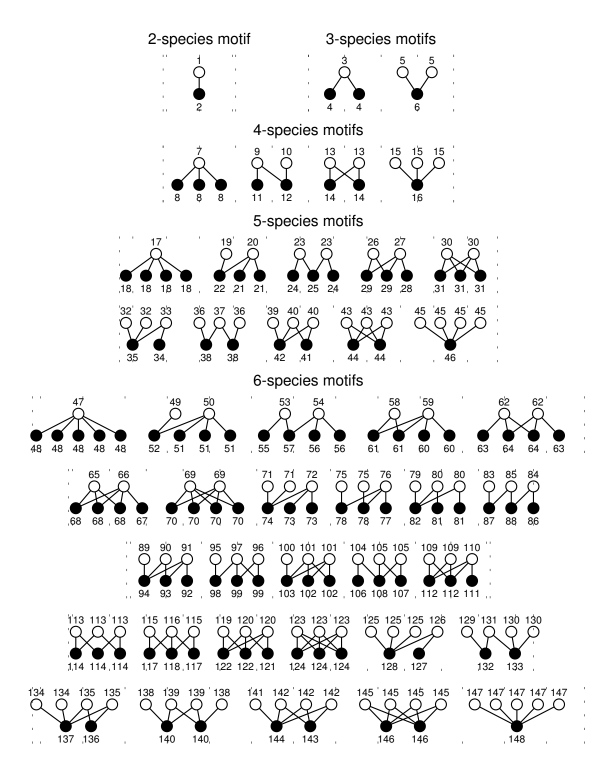


Figure 2: The singe two-species motif, two three-species motifs, four four-species motifs, 10 five-species motifs, and 27 six-species motifs that can be found in bipartite networks. Unique positions are numbered, but note that positions are not in the same order as in the output provided by motif_roles. If the identity of particular motifs and/or positions is important, be sure to note the motif number and position information provided in the output file.

References

[1] Daniel B Stouffer, Juan Camacho, Wenxin Jiang, and Luís A. Nunes Amaral. Evidence for the existence of a robust pattern of prey selection in food webs. *Proceedings of the Royal Society B: Biological Sciences*, 274(1621):1931–1940, August 2007. doi: 10.1098/rspb.2007.0571.