SuperNoder: a tool to discover over-represented modular structures in networks
Danilo Dessì¹*, Jacopo Cirrone², Diego Reforgiato Recupero¹ and Dennis Shasha²

Abstract
Background: Networks whose nodes have labels can seem complex. Fortunately, many have substructures that occur often (“motifs”). A societal example of a motif might be a household. Replacing such motifs by named supernodes reduces the complexity of the network and can bring out insightful features. Doing so repeatedly may give hints about higher level structures of the network. We call this recursive process Recursive Supernode Extraction.

Results: This paper describes algorithms and a tool to discover disjoint (i.e. non-overlapping) motifs in a network, replacing those motifs by new nodes, and then recursing. We show applications in food-web and protein-protein interaction (PPI) networks where our methods reduce the complexity of the network and yield insights.

Conclusions: SuperNoder is a web-based and standalone tool which enables the simplification of big graphs based on the reduction of high frequency motifs. It applies various strategies for identifying disjoint motifs with the goal of enhancing the understandability of networks.

Keywords: Motifs discovery, PPI interaction network, Food-web network, Computational complexity, Network compression

Background
Imagine describing a road map with words alone. The task would be difficult and unclear to most people. Networks provide a far better representation of any data representing interrelationships. However, because the size of modern networks (for example, in social science) can extend to thousands, millions, or even billions of nodes, networks themselves need to be abstracted for the sake of intelligibility and insight.

As in other disciplines, a way to reduce the size of the problem is to discover similar components and give them a common name. Linguists do this when they categorize parts of speech (noun, verb, adverb etc). Biologists do this when they group animals into species and families. In networks, we will do this by finding connected labeled sub-components that are isomorphic in label and topology. Formally, this entails finding common subgraphs or motifs that occur with a certain frequency.

Much research has proposed algorithms that aim at finding frequent motifs [1–5]. The motivation is usually to gain insights about metabolic and protein-protein interactions, ecological food-webs, social networks, collaboration networks, information networks of interlinked documents and products [6–14].

Most of this work does not distinguish between motifs that overlap and motifs that do not. However, this distinction can be critical for understandability. For example, households are a convenient abstraction in social graphs because they are disjoint whereas friendship motifs do not tend to be. For networks whose motifs are not naturally disjoint, identifying disjoint motifs may help to understand network structure (e.g. cliques in friendship networks). One work that has done this is [15] which showed algorithms to find edge-disjoint motifs in unlabelled networks. Our work focuses on node-disjoint motifs (which share neither nodes nor edges) in labeled networks. The usefulness of labels is intuitive as we will see in our examples and node-disjoint motifs are readily decomposable. We also present promising algorithms to make this process reasonably fast even for sizeable networks.
Once disjoint motifs of a certain size $k$ have been identified, each such motif can be collapsed into a **supernode**, which is a single node that inherits all the connections and properties of the motifs. This procedure can be performed recursively in order to find motifs on graphs consisting of a combination of nodes and super-nodes. Figure 1 shows an example where motifs have been collapsed into supernodes.

Thus, our tool SuperNoder finds disjoint motifs on a base graph $G_1$, reducing $G_1$ to a new graph $G_2$, and then recursively repeats the procedure to find $G_3$, $G_4$, and so on. SuperNoder attempts to find the most possible disjoint frequent motifs of a given size in a target network in each stage of the process. We present several techniques to achieve this goal.

Orthogonally, the SuperNoder tool can take input nodes at different layers in a label hierarchy. For example in phylogeny, there is a hierarchy of species, genus, family, kingdom. Relationships that may be obscure at a low level may be clearer at a high level (e.g. felines eat rodents).

This paper makes three contributions:

- Efficient algorithms to find disjoint supernodes in labeled networks, including networks already containing supernodes, yielding a recursive algorithm.
- A tool incorporating these algorithms that is free to the community.
- Example applications to show the usefulness of the approach.

Frequent (based on the possibly overlapping F1 measure) motifs have been shown to give insights in regulatory [16], food-web [17–19], and social science [20, 21] networks. Reduction methods aim at minimizing the loss of information while maximizing the understandability, often establishing which components are less interesting for the behavior of networks. Recent studies have focused on finding high-order clusterings [22, 23]. However, most of this research has focused on modeling graphs without considering node labels, despite the fact that many networks have them. Moreover, they usually consider overlapping motifs, therefore, a single node can belong to several patterns, making further analysis (and understandability) difficult.

An early compression graph method was proposed by [24] where the authors show how finding substructures and merging them in vertexes for compressing data. Our approach builds on theirs, but their approach does not find all substructures that occur nor does it attempt to find the most highly repetitive subgraphs which are the best candidates for capturing subgraph regularities.

Our work also draws inspiration from [15] where the authors propose two methods to find disjoint motifs under the F2 frequency measure (where two graphs are disjoint if they do not share a common edge). First, they propose a method to find motifs based on a small set of patterns, and then give methods to find non-overlapping motifs solving the Maximum Independent Set (MIS) problem. They invented their own method for finding frequent motifs and did not choose to compare their method with state-of-the-art motif-finding techniques [25–30]. By contrast, we have chosen to base our approach on the motif-finding algorithm of [25] because of its simple implementation and promising results [31]. As in [15], the second phase of our algorithm uses an overlap graph.

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**Fig. 1** Example of motifs collapsed into supernodes in a Protein-Protein Interaction network. **a** The original nodes of the network. **b** The new nodes of the network after two motifs of size three have been collapsed.
and we have explored some heuristics to deal with larger overlap graphs beyond what they used.

While we do contribute algorithms for finding disjoint motifs given a collection of already found motifs, we do not advance the state of the art in finding the motifs themselves. Instead, our work builds on top of an existing overlapping motif finding algorithm which has been compared and studied many times in literature [31].

The remainder of this paper is organized as follows. “Implementation” section describes the proposed approach. “Results” section describes the biological datasets we have used, shows an example application of SuperNoder to the yeast network, and analyzes both the performance and quality of SuperNoder on real networks. “Conclusions” section gives perspectives on the problem and future directions. “Availability and requirements” section lists abbreviations we use in the paper.

Implementation

Labeled networks or graphs are formally characterized by a triple $G = (N,E,L)$ where $N$ denotes a set of nodes, $E$ denotes a set of edges (pairs) $e = (n_i,n_j) \in N$, and $L$ is a mapping from $N$ to some set of labels. Edges represent an application-dependent relationship. For instance, an edge may connect two nodes representing people if the people are friends.

We say that a graph is undirected if every edge from $n$ to $n'$ implies the existence of an edge from $n'$ to $n$. Otherwise the graph is said to be directed. A subgraph is a connected component $G_S = (N_S,E_S)$ such that $N_S \subseteq N$ and $E_S \subseteq E$ if there exists a path from each $n_i \in N_S$ to each $n_j \in N_S$. A $k$-subgraph is a subgraph with $k$ nodes.

Two subgraphs $S_1, S_2$ are isomorphic if (i) there exists a bijective function $f : N_{S1} \rightarrow N_{S2}$ such that for each pair $(n_i,n_j) \in E_{S1}$ if and only if $(f(n_i),f(n_j)) \in E_{S2}$ and (ii) for all $k$, the label of $n_k$, $L(n_k)$ is the same as $L(f(n_k))$. To count the number of occurrences of a given subgraph, three different measures can be used [32]. The first measure, named F1, is the count of each subgraph regardless of whether it overlaps with others. The second one, named F2, avoids overlaps of subgraphs if they share at least an edge (or equivalently a connected pair of nodes). The last one, named F3, requires that two subgraphs share no nodes. F3 is therefore, the most strict criterion of disjointness (and is the one used in this paper). We define the frequency of a subgraph $S_1$ in $G$ as the number of occurrences of $S_1$ in $G$. We call subgraphs $k$-motifs if they have $k$ nodes and occur over a threshold $t$ using the F1 measure.

The SuperNoder pipeline consists of the following steps:

1. Solicit a size $s$ from the user corresponding to the number of nodes each motif should have.
2. Solicit a threshold $t$ from the user corresponding to the number of times that a motif should be present to be considered. (In the future, we may add specific shapes of motifs or specific motifs labels, as further filters in addition to threshold.)
3. Search for all possible motifs in the input network meeting threshold $t$, using the F1 measure (i.e. allowing overlaps). Call that set $M$.
4. Search for the maximum number of non-overlapping motifs from $M$.
5. Collapse non-overlapping motifs into supernodes.
6. Repeat steps 2 through 5 until satisfied.

In this section we provide details of our tool for accomplishing these tasks.

Input network and motifs finding

SuperNoder requires two series of data as an input:

- A list of node rows, where each row represents a node by means of a unique ID and a label separated by a blank space.
- A list of edge rows, where each row consists of two node IDs separated by a blank space.

SuperNoder uses the Randomized Enumeration algorithm [25] for the purpose of motif finding. The result of the algorithm is a set of all possible undirected motifs in the network, allowing overlaps.

Motif count and thresholding

To count motifs, we implemented a function to compute isomorphisms between subgraphs similar to the one of Cordella and colleagues [33]. First, the algorithm takes the labels of subgraph nodes and counts how many nodes have the same label. Second, for each label it computes the sum of in-degrees and the sum of out-degrees (i.e. for each node label, it computes $l_{n,i,o}$, where $n$ is the number of nodes with label $l$, $i$ is the sum of in-degree of nodes with label $l$, and $o$ is the sum of out-degree of nodes with label $l$). Finally, it sorts these labels using the lexicographic order and computes their hash. If the number of subgraphs having hash value $h$ is greater than the user-given threshold $t$, then all such subgraphs are checked to see how many are in fact isomorphic. If, after the check, the number is greater than $t$, then those subgraphs pass the initial filter to be a motif and thus belong to the “frequent motif set”. Thus the frequent motif set may contain different topologies, e.g. at least $t$ stars of size $s$, at least $t$ paths of length $s$, and so on.

Finding disjoint motifs

Our methods to find disjoint motifs, given the potentially overlapping frequent motif set, uses the concept of an overlap graph. An overlap graph is a pair $(M,E)$ where $M$
is the set of motifs and there is an edge between motif $m_1$ and motif $m_2$ if they share at least one node in the original graph. (In the case of recursive reduction, the original graph at reduction $i$ is the one produced from the graph at reduction $i-1$, containing both normal nodes and supernodes.)

We briefly present an overview of our heuristics for finding disjoint motifs here (Table 1), but the full pseudocode is available in the github site containing the SuperNoder source code as well.

**H1 (Greedy Elimination).** This simple but effective heuristic finds disjoint motifs by using a Maximal Independent Set technique. Given the frequent motif set $M$ and a user-given parameter $n$, randomly shuffle the potentially overlapping motif instances from the frequent motif set $M$. For each motif instance $m$, if the motif instance overlaps no other motif instances of $M$, then output it. Otherwise remove it and all its edges from the overlap graph. Because this approach is naively greedy, SuperNoder tries $n$ (parameter given by the user) different random shufflings to try to obtain the greatest number of disjoint motifs.

**H2 (Ramsey).** Heuristic-2 exploits both sampling and the Ramsey method whose functions can be seen in [34]. Given the list of motif instances $M$ and a number $k$, the heuristic (i) takes disjoint subsets of size $k$ from $M$ and constructs the induced subgraph of the overlap network from each subset. (ii) On each subgraph, it performs the Ramsey algorithm obtaining a $MIS_{subgraph}$. (iii) Then, it merges all $MIS_{subgraphs}$ into a reduced list of motif instances which takes the role of $M$. The algorithm repeats steps (i) through (iii) until there are no more overlaps and outputs the resulting set of motifs.

**H3 (Ranked Elimination).** Heuristic-3 assigns to each (possibly overlapping) motif instance $m$ a degree equal to the sum of degrees of the nodes in $m$ ignoring the edges between nodes in $m$ (i.e. the sum of the degrees of the nodes in $m$ pertaining to edges that connect to nodes outside $m$). The algorithm then orders the motif instances in ascending order of degree so calculated, forming a list called $MotifDegree$. For each node $n$ in the original graph, find the first motif instance in $MotifDegree$ and discard all other motifs in $MotifDegree$ containing $n$. This process yields a new list called $PotentialSuperNodes$. Then traverse this $PotentialSuperNodes$ list, preserving motif instances having no overlaps and deleting motif instances that have higher degrees when there are overlaps.

**H4 (Repeated Ranked Elimination).** This approach is an improvement over H3, because H3 misses some motif instances when one or more overlapping motif instances are removed and the nodes of the removed motif instances then have no chance to be included in any other motif instances. Given as input the list of motif instances $M$ found using the Randomized Enumeration method seen above, build the $MotifDegree$ list as in Heuristic-3. For each node $n$, the motif instance $m \in MotifDegree$ with the lowest degree that contains $n$ is copied to a list of potential supernodes, called $PotentialSuperNodes$. All the motif instances in $PotentialSuperNodes$ with no overlaps are considered valid. Then, for each pair $\{m', m''\}$ of overlapping motif instances in $PotentialSuperNodes$, discard the motif instance with the higher degree. Continue until there are no more motif instances. Now consider all the nodes $N_{orphan}$ that are not in any disjoint motif instance found so far and consider motif instances based on the F1 measure that apply to nodes of $N_{orphan}$. Repeat the above procedure to generate more disjoint motif instances. Repeat until there are no more nodes in $N_{orphan}$.

**H5 (Sampled Ranked Elimination).** This heuristic unifies sampling with the overlap graph approach. After the sampling is done as for the Ramsey algorithm, the heuristic constructs an overlap graph on the surviving motif instances. The heuristic considers the motif instances in ascending order by degree in the overlap graph. If a motif instance has no edges, then put it in the result. If a motif instance $m_1$ has an edge with another motif instance $m_2$, then remove the motif instance with the largest degree.

### Network reduction

After the non-overlapping motif instances have been found, each one is collapsed into a supernode, preserving the external connections of the original nodes of motifs. The label of each supernode is the concatenation of labels of its member nodes in alphabetical order. The new network can be saved as an output using the same format as the input network and the whole pipeline can be iterated on it.

### Results

**The test networks**

We demonstrate SuperNoder on three different labeled biological networks:

| Table 1 | Summary of the characteristics of the heuristics |
|---------|-----------------------------------------------|
| Heuristic ID | Overlap graph | Ramsey | Order by degree | Random approach | Sampling approach |
| H1 | - | - | - | V | - |
| H2 | V | V | - | - | V |
| H3 | - | - | V | V | - |
| H4 | - | - | V | V | - |
| H5 | V | V | V | V | V |

The symbol V indicates that the heuristic exploits that characteristic. - if not. H1 = Greedy Elimination, H2 = Ramsey, H3 = Ranked Elimination, H4 = Ranked Replacement, H5 = Sampled Ranked Elimination
Fig. 2 An example of four supernodes built using SuperNoder with motifs of size three on the yeast network. From left to right, labels of original nodes, labels of the fifth level hierarchy, labels of the third level hierarchy. On the third level, many proteins share the same pattern and these patterns are often disjoint.

- A food-web subnetwork of Florida bay network\(^1\) [35] with 93 nodes and 960 edges.
- A Protein-Protein Interaction (PPI) network of yeast\(^2\) [36] with 2361 nodes and 7182 edges.
- A PPI network of Arabidopsis\(^3\) [37] with 18167 nodes and 10928 edges.

**Food-web network.** The original nodes have labels that represent animals or plants (e.g. predatory chanodichthys, dinoflagellates, coral bryaninops, etc.). We have mapped the network using a taxonomy\(^4\), retrieving for each node genus, family, order, class, phylum, and kingdom. From the original network we have removed species that did not have higher phylogenetic categories.

**Protein-Protein Interaction networks.** In a Protein-Protein Interaction (PPI) network, each node represents a different protein. For the higher-level categorization of PPI networks, we have employed the ontology annotations available at this link\(^5\). First, we have retrieved the Gene Ontology (GO) term that belongs to Biological Processes (BPs) and that has the lowest (i.e. most empirically based) evidence code for each protein. Second, we have traversed the ontology go-basic\(^6\) starting from each GO term in our network to the GO term which represents all Biological Processes. Since each GO term can have more than one parent, we have chosen the GO term with the lowest (i.e., most conclusive) evidence code going up in the hierarchy. More precisely, given a label of a node \(l\), we retrieve a GO term \(g\) with the lowest evidence code. Let \(\{g_1, g_2, \ldots, g_n\}\) be the parents of \(g\), then we choose the \(g_i\) with \(1 \leq i \leq n\) with the lowest evidence code, building a hierarchy \(l, g, g_i\). Then, we repeat the same operation as long as the GO term which represents all Biological Processes (BPs) has not been yet reached. In doing so, we have built a taxonomy that can enable the analysis of protein functions.

**Use case**

In the analysis of biological networks, interactions often occur between proteins of the same class [38]. SuperNoder can find these relations when high level functional classes are considered, highlighting frequent related processes and simplifying their identification.

To show how SuperNoder may help to simplify networks, we focus on the yeast network, and explain how higher levels of the Gene Ontology (GO) terms enable the abstraction of protein functions allowing SuperNoder to reduce the network complexity. The motivation is simple: at a lower level in the hierarchy of GO terms there may be no motifs that occur more than \(t\) times for a moderately large \(t\). At higher levels, there might be. In the example, the yeast network has been mapped onto five levels of the GO terms hierarchy. To be considered a motif, a subgraph has to occur at least 50 times, i.e. with threshold \(t = 50\).

Table 2 An example of a hierarchical exploration of the yeast network

|       | th | Original | L5   | L4   | L3   | L2   | L1   |
|-------|----|----------|------|------|------|------|------|
| Motifs| 25 | 0        | 290  | 292  | 319  | 377  | 389  |
| Nodes | 25 | 2361     | 1781 | 1776 | 1607 | 1583 | 1333 |
| Edges | 25 | 7182     | 5234 | 5305 | 5018 | 5020 | 5322 |
| Motifs| 50 | 0        | 240  | 236  | 304  | 388  | 390  |
| Nodes | 50 | 2361     | 1841 | 1889 | 1585 | 1361 | 1581 |
| Edges | 50 | 7182     | 5339 | 5429 | 5029 | 5347 | 4990 |

The table reports the number of found motifs, the number of nodes and edges, when the network is mapped to different levels of the GO terms hierarchy and then reduced. At higher levels (L1 is higher level than L2 etc) more motifs pass the threshold.
Figure 2 shows a motif of size three in each row that are mapped on the base level (gene labels), the fifth-level (L5) and the third-level (L3) hierarchy labels (i.e. in ascending order of abstraction). More motifs appear at higher levels in the hierarchy (i.e. first on L5 and then on L3 levels). In fact, with L5 labels the triples in row 2 and row 3 are isomorphic. When L3 labels are used, all triples are isomorphic, thus becoming relevant motifs. Those triples are collapsed into supernodes thus forming a new simplified network. Supernodes indicate proteins that belong to the same class helping biologists with the analysis of basic interactions.

As a specific case study, focus on motifs composed of proteins \((\text{YNL306W}, \text{YDR175C}, \text{YBR251W})\) and \((\text{YGR156W}, \text{YKR002W}, \text{YLR115W})\). Analyzing the network on the base labels, there are no supernodes, since they do not show common features in the labeled graph. Already at lower hierarchical levels (i.e. L5), the motifs GO terms are abstracted into functions, viz, \textit{macromolecule biosynthetic process} and \textit{cellular macromolecule metabolic process} respectively. At hierarchical level L3, the proteins in this example have the label GO:0071704 which indicates that their proteins are related to \textit{organic substance metabolic process}. At that level, we find out that \textit{organic substance metabolic process} (GO:0071704) covers an important role into the yeast network, and that is mainly composed of \textit{macromolecule biosynthetic process} (GO:0009059), \textit{cellular macromolecule metabolic process} (GO:0044260) and \textit{protein metabolic process} (GO:0019538). This shows an example of how our tool can help biologists understand the behavior of proteins (with frequent motifs) belonging to the same class.

The higher the hierarchy levels, the larger the number of relevant motifs that can be used to further reduce the current network (an example of this behavior can be observed in Table 2). In addition, higher level labels enable higher thresholds, sometimes leading to the discovery of very frequent motifs. For example, connections of proteins in Fig. 3a do not show functionalities but those become evident at higher hierarchical levels 3b and 3c. For example, the frequent relation between proteins which have GO:0044237, GO:0044237, GO:0044237 as GO terms as showed in Fig. 3c are only detectable at that level of the hierarchy. Finally, images 3b and 3c show that the reduction at a high level of abstraction enables a better understandability of the network.

### Performance

In this section, we report the time performance, the number of disjoint motifs and the reduction ability of our heuristic algorithms. The time performance is based on the wall clock time required for the execution of the heuristics on all relevant motifs. The number of disjoint motifs is the number of motifs found by each algorithm. The reduction ability is the extent of reduction of networks. All experiments have been performed considering motifs with size = 3 and size = 5 (i.e. having three nodes in the original graph and three nodes or supernodes after each step of the recursion). H1 has been performed with

| Table 3 | Rows list the number of all motifs, the threshold applied in our experiments and the number of motifs that meet that threshold when L3 labels are considered and motifs have size 3 |
| --- | --- | --- |
| Network | N motifs | Threshold | N repetitive motifs |
| Food-Web | 20283 | 5 | 5085 |
| Yeast | 96444 | 50 | 49294 |
| Arabidopsis | 268437 | 100 | 155185 |

| Table 4 | Rows list the number of all motifs, the threshold applied in our experiments and the number of motifs that meet that threshold when L3 labels are considered and motifs have size 5 |
| --- | --- | --- |
| Network | N motifs | Threshold | N repetitive motifs |
| Food-Web | 26841 | 5 | 407 |
| Yeast | 188733 | 50 | 11550 |
| Arabidopsis | 425895 | 100 | 14474 |
five shufflings. H2 and H5 adopted subsets of the overlap graphs consisting of 1000 motif nodes. In our simulations, we chose different thresholds in different networks, as shown in Tables 3 and 4. The reason is that certain thresholds make no sense for certain networks. For example, a threshold of 100 for our food-web network is meaningless because no motifs occur that frequently.

**Food-web network**

Figure 4 reports the performance of the heuristics applied on the food-web network. In this case, heuristics H1, H2 and H5 which exploit repetitive random approaches (H1), sampled overlap graph (H2 and H5), and H4 show better performance than others in finding disjoint motifs. Heuristics H3 shows a poor reduction factor on this network. The reason is that there are many motifs with the same sums of degrees, so degree-based heuristics do not work well. Heuristic H1 is the fastest. This holds regardless of motif size. In fact, overall, heuristic H1 is both fast and has a good reduction factor.

**Yeast network**

Figure 5 shows the performance on the yeast network. In contrast to the food-web network, heuristics H2 and H5 based on the sampled overlap graph do not obtain the best reduction factor. In this case, heuristic H4 enjoys a greater reduction factor. Although heuristics H2 and H5 can find a large number of disjoint motifs, they require excessive time to find a solution, hence, their use on a network of this size might be avoided. The heuristics H1 and H3 are still the fastest.

**Arabidopsis network**

Experimental results on arabidopsis networks (see Fig. 6) are similar to those on the yeast network and the same
considerations hold. Note that the arabidopsis network is a Protein-Protein Interaction network like the yeast network but is very different in term of size.

**Observations from the Experiments**
Heuristic H1 achieves the best time performance and finds a large number of disjoint motifs though not always the maximum number. Heuristic H4 which is slower can sometimes find more disjoint motifs so should be considered if time is available. The size of motifs and the threshold also matter. Larger motifs entail the processing of more data, but there are fewer repetitive motifs (i.e. motifs that exceed the threshold) so the overall time is sometimes less.
In summary, heuristic H1 shows good performance on all types of network since its greedy approach is fast. The resulting reduction may not however be best. Heuristics H2 and H5 which employ sampling are useful for those networks whose overlap graphs are very large. The size of samples can be chosen according to the available computational resources to balance the execution time and memory use. Heuristic H2 should show better reduction performance than H5 when there are few distinct motifs degree values. By contrast, H3 and H4 should be useful for all those networks that have many distinct motifs degree values, because motifs having less probability to overlap are detected faster.

Reduction
Figures 7 and 8 show the extent of graph reduction on the food-web and yeast networks respectively. Unsurprisingly, lowering the threshold generates more F1 motifs, increasing the number of F3 motifs and reducing the network size. In our example networks, after a few iterations, the networks are no longer reduced. When this plateauing happens depends entirely on the data. In addition, the threshold and the motif size both affect the reduction factor, because a small motif has a higher probability of occurring more often (see Tables 3 and 4). This is well illustrated by our tests where motifs of size 3 show a greater reduction than motifs of size 5. For an illustration of the extent of reduction, consider Fig. 9 where (a) shows the original food web network, (b) after one iteration and (c) after two iterations.

Tool description
Figure 10 shows the graphical interface of SuperNoder that users without programming skills can adopt to analyze networks. On the left, users can use a panel to create nodes, in the center there is one panel to create edges, and, on the right, a list of parameters the user can set. With the first option users can choose the size of motifs they are interested in. The minimum value is 3. The next option is related to the heuristic that should be employed to find disjoint motifs. The user can also choose the type of network: direct or undirect. The fourth parameter is the threshold which represents the minimum value each motif should meet to be considered over-represented (it corresponds to the threshold $t$ of the SuperNoder pipeline algorithm). The last required parameter is the number of iterations. In addition, if the user selects the H1 heuristic, he/she can set the number of repetitions to be executed, specific for H1. If the user selects either the H2 or H5 heuristic, he/she can also

![Fig. 8 Reduction performance on five iterations on the yeast network](image-url)
choose the size of samples. When the Submit network button is clicked, the SuperNoder pipeline will be run and results will be printed and shown online (but not saved anywhere).

The output consists of two sections (nodes and edges) for each chosen iteration using the same input format. Supernodes are indicated by the tag #supernode.

The code has been developed in Python 3.6 using NetworkX\textsuperscript{7} library. SuperNoder functionalities operate on graphs using the standard NetworkX format. The web interface is provided by a python server which runs on a Docker\textsuperscript{8} container. Last but not least, SuperNoder is hosted on a GitHub\textsuperscript{9} page and distributed as a Docker file with the source code freely available under GPLv3 License.

**Conclusions**

SuperNoder enables the simplification and compression of graphs based on high frequency motifs. By identifying disjoint motifs, SuperNoder enhances understandability as the network is reduced. This paper describes and compares various algorithms on real networks, both to show the benefits of the approach and to find high-performing algorithms. SuperNoder has been developed in Python, it can either be installed on local machines or used through its online web interface. Future work includes enhancing performance yet further by using Graphical Processing Units.

**Availability and requirements**

**Project name:** SuperNoder  
**Project homepage:** http://glab.sc.unica.it/supernoder/  
**Github link:** https://github.com/danilo-dessi/SuperNoder-v1.0  
**Operating system(s):** Platform independent  
**Programming language:** Python  
**Other requirements:** Docker  
**License:** GPLv3.  
**Any restrictions to use by non-academics:** nothing.
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Author details
1 Department of Mathematics and Computer Science, University of Cagliari, 09124 Cagliari, Italy. 2 Department of Computer Science, Courant Institute of Mathematical Sciences, New York University, 10012 New York City, USA.

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References
1. Blondel VD, Guillaume J-L, Lambiotte R, Lefebvre E. Fast unfolding of communities in large networks. J Stat Mech Theory Experiment. 2008;2008(10):P10008.
2. Newman MEJ, Girvan M. Finding and evaluating community structure in networks. Phys Rev E. 2004;69(2):026113. https://doi.org/10.1103/PhysRevE.69.026113.
3. Mucha P, Richardson T, Macon K, Porter MA, Onnela J-P. Community structure in time-dependent, multiscale, and multiplex networks. Science. 2010;328(5980):876.
4. Granell C, Gómez S, Arenas A. Hierarchical multiresolution method to overcome the resolution limit in complex networks. Int J Bifurcation Chaos. 2012;22(07):1250171. https://doi.org/10.1142/5021872412501714.
5. Ball B, Barrat A, Newman MEJ. Efficient and principled method for detecting communities in networks. Phys Rev E. 2011;84:036103. https://doi.org/10.1103/PhysRevE.84.036103.
6. Benson AR, Gleich DF, Leskovec J. Higher-order organization of complex networks. CoRR. 2016;abs/1612.08447. 1612.08447
7. Yang J, Leskovec J. Overlapping communities explain core–periphery organization of networks. Proc IEEE. 2014;102:1892–1902.
8. Yang Z, Algesheimer R, Tessone CJ. A comparative analysis of community detection algorithms on artificial networks. Sci Rep. 2016;6:30750.
9. Evans TS, Lambiotte R. Line graphs, link partitions, and overlapping communities. Phys Rev E. 2009;80:016105. https://doi.org/10.1103/PhysRevE.80.016105.
10. Palla G, Derenyi I, Farkas I, Vicsek T. Uncovering the overlapping community structure of complex networks in nature and society. Nature. 2005;435(7043):814–8.
11. Shi X, Lu H, Jia G. Adaptive overlapping community detection with bayesian nonnegative matrix factorization. In: International Conference on Database Systems for Advanced Applications. Springer; 2017. p. 339–353.
12. Sales-Pardo M, Guimerà R, Moreira AA, Amaral LAN. Extracting the hierarchical organization of complex systems. Proc Natl Acad Sci. 2007;104(39):15224–9. https://doi.org/10.1073/pnas.0703470104. http://www.pnas.org/content/104/39/15224.full.pdf.
13. Airoldi EM, Blei DM, Fienberg SE, Xing EP. Mixed membership stochastic blockmodels. J Mach Learn Res. 2008;9:1981–2014.
14. Ahn Y-Y, Bagrow JP, Lehmann S. Link communities reveal multiscale complexity in networks. Nature. 2010;466(7307):761–4.
15. Elhesha R, Kahveci T. Identification of large disjoint motifs in biological networks. BMC Bioinformatics. 2016;17(1):408.
16. Legeay M, Duval B, Renou J-P. Differential functional analysis and change motifs in gene networks to explore the role of anti-sense transcription. In: International Symposium on Bioinformatics Research and Applications. Springer; 2016. p. 117–26.
17. Baiser B, Elhesha R, Kahveci T. Motifs in the assembly of food web networks. Oikos. 2016;125(4):480–91.
18. Monteiro AB, Faria LDB. The interplay between population stability and food-web topology predicts the occurrence of motifs in complex food-webs. J Theor Biol. 2016;409:165–71.
19. McDonald-Madden E, Sabadin R, Game E, Baxter P, Chadès I, Possingham H. Using food-web theory to conserve ecosystems. Nat Commun. 2016;7:10245.
20. Rotabi R, Kamath K, Kleinberg J, Sharma A. Detecting strong ties using network motifs. In: Proceedings of the 26th International Conference on World Wide Web Companion. ACM; 2017. p. 983–92. International World Wide Web Conferences Steering Committee. https://doi.org/10.1145/3041021.3055139.
21. Wang S, Zhang Y, Wang H, Huang Z, Wang X, Jiang T. Large scale measurement and analytics on social groups of device-to-device sharing in mobile social networks. Mob Netw Appl. 2017;23:1–13.
22. Yin H, Benson AR, Leskovec J, Gleich DF. Local higher-order graph clustering. In: Proceedings of the 23rd ACM SIGKDD International Conference on Knowledge Discovery and Data Mining. ACM; 2017. p. 555–64.
23. Benson AR, Gleich DF, Leskovec J. Higher-order organization of complex networks. Science. 2016;353(6295):163–6.
24. Holder LB, Cook DJ, Dijkstra S, et al. Substructure discovery in the subdue system. In: KDD Workshop, 1994. p. 169–80. http://dl.acm.org/citation.cfm?id=3008850.3008868.

Abbreviations
MIS: Maximum independent set; PPI: Protein-protein interaction

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Availability of data and materials
The tool developed during the study and experimental data are publicly available at https://github.com/danilo-dessi/SuperNoder-v1.0.

Authors’ contributions
All authors were involved in the development of the tool. All authors read and approved the manuscript.

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Not applicable.

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Competing interests
The authors declare that they have no competing interests.

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Endnotes
1 https://snap.stanford.edu/data/Florida-bay.html
2 http://vlado.fmf.uni-lj.si/pub/networks/data/bio/yeast/yeast.htm
3 http://interactome.dfci.harvard.edu/A_thaliana/index.php?page=download
4 https://ftp.ncbi.nlm.nih.gov/pub/taxonomy/
5 http://www.geneontology.org/page/download-annotations
6 http://www.geneontology.org/page/download-ontology
7 https://networkx.github.io/
8 https://www.docker.com/
9 https://github.com/danilo-dessi/SuperNoder-v1.0
25. Wernicke S, Rasche F. Fanmod: a tool for fast network motif detection. Bioinformatics. 2006;22(9):1152–3.
26. Schreiber F, Schwobbermeyer H. Mavisto: a tool for the exploration of network motifs. Bioinformatics. 2005;21(17):3572–4.
27. Chen J, Hsu W, Lee ML, Ng S-K. Nemofinder: Dissecting genome-wide protein-protein interactions with meso-scale network motifs. In: Proceedings of the 12th ACM SIGKDD International Conference on Knowledge Discovery and Data Mining. ACM, 2006. p. 106–15.
28. Kashani ZRM, Ahrabian H, Elahi E, Nowzari-Dalini A, Ansari ES, Asadi S, Mohammadi S, Schreiber F, Masoudi-Nejad A. Kavosh: a new algorithm for finding network motifs. BMC Bioinformatics. 2009;10(1):318.
29. Kashtan N, Itzkovitz S, Milo R, Alon U. Efficient sampling algorithm for estimating subgraph concentrations and detecting network motifs. Bioinformatics. 2004;20(11):1746–58.
30. Omidi S, Schreiber F, Masoudi-Nejad A. Moda: an efficient algorithm for network motif discovery in biological networks. Genes Genet Syst. 2009;84(5):385–95.
31. Masoudi-Nejad A, Schreiber F, Kashani ZRM. Building blocks of biological networks: a review on major network motif discovery algorithms. IET Syst Biol. 2012;6(5):164–74.
32. Schreiber F, Schwobbermeyer H. Frequency concepts and pattern detection for the analysis of motifs in networks. Lect Notes Comput Sci. 2005;3737:89–104.
33. Cordella LP, Foggia P, Sansone C, Vento M. An improved algorithm for matching large graphs. In: 3rd IAPR-TC15 Workshop on Graph-based Representations in Pattern Recognition, 2001. p. 149–59.
34. Boppana R, Halldorsson MM. Approximating maximum independent sets by excluding subgraphs. BIT Numer Math. 1992;32(2):180–96.
35. Robert E Ulanowicz CB, Egnotovich MS. Network Analysis of Trophic Dynamics in South Florida ecosystems–The Florida Bay Ecosystem: Annual Report to the U.S. Geological Survey, 1997.
36. Bu D, Zhao Y, Cai L, Xue H, Zhu X, Lu H, Zhang J, Sun S, Ling L, Zhang N, et al. Topological structure analysis of the protein–protein interaction network in budding yeast. Nucleic Acids Res. 2003;31(9):2443–50.
37. Consortium AIM, et al. Evidence for network evolution in an arabidopsis interactome map. Science. 2011;333(6042):601–7.
38. Wuchty S, Almaas E. Peeling the yeast protein network. Proteomics. 2005;5(2):444–9.