Biological network motif detection: principles and practice

Elisabeth Wong, Brittany Baur, Saad Quader and Chun-Hsi Huang

Submitted: 28th February 20II; Received (in revised form): 13th May 20II

Abstract

Network motifs are statistically overrepresented sub-structures (sub-graphs) in a network, and have been recognized as 'the simple building blocks of complex networks'. Study of biological network motifs may reveal answers to many important biological questions. The main difficulty in detecting larger network motifs in biological networks lies in the facts that the number of possible sub-graphs increases exponentially with the network or motif size (node counts, in general), and that no known polynomial-time algorithm exists in deciding if two graphs are topologically equivalent. This article discusses the biological significance of network motifs, the motivation behind solving the motif-finding problem, and strategies to solve the various aspects of this problem. A simple classification scheme is designed to analyze the strengths and weaknesses of several existing algorithms. Experimental results derived from a few comparative studies in the literature are discussed, with conclusions that lead to future research directions.

Keywords: Network motifs; biological networks; graph isomorphism

INTRODUCTION

Biology of a living cell involves many intricate networks of interdependent events and interactions among biomolecules. Examples of such networks include transcriptional or gene regulation networks, protein-protein interaction (PPI) networks, metabolic pathways, neural networks, etc. In 2002, Milo et al. [1, 2] showed that networks from diverse fields-biological and non-biological-contain several small topological patterns that are so frequent that it is unlikely to occur by chance. Different networks tend to have different sets of such frequent local structures [1]. These patterns, referred to as 'network motifs', are recognized as 'the simple building blocks of complex networks' [1]. The discovery spawned a multitude of research efforts in the past decade and the area is fertile to this day. Network motifs are also studied in such other networks as the electronic circuits and power distribution networks, ecological networks (food web), software engineering diagrams, molecular structures, World Wide Web (the Internet), and social networks, etc [1, 3].

Biological network motifs are shown to perform various computational tasks in the network. For example, consider the motif called 'Feedforward Loop' or FFL [1, 4–7]. This motif is commonly found in many gene systems and organisms [1, 2, 8–11]. The motif consists of three genes and three regulatory interactions. The target gene Z is regulated by two transcription factors (TFs) X and Y and in addition TF Y is also regulated by TF X. Since each of the regulatory interactions may either be positive or negative, there are possibly eight types of FFL motifs [6]. Two of those eight types: the coherent

Corresponding author. Chun-Hsi Huang. Department of Computer Science and Engineering, University of Connecticut, Storrs, CT 06269, USA. Tel: +1-860-486-5472; Fax: +1-860-486-4817; E-mail: huang@engr.uconn.edu

Elisabeth A. Wong is currently a Biology/Mathematics major at the Bowdoin College, Brunswick, Maine. She was a recipient of the 2010 NSF Bio-Grid REU fellowship.

Brittany Baur is currently a Biology/Mathematics major at the Manhattanville College, Purchase, New York. She was a recipient of the 2010 NSF Bio-Grid REU fellowship.

Saad Quader is currently a graduate student at the Department of Computer Science and Engineering, University of Connecticut. He received his BSc in Computer Science and Engineering from Bangladesh University of Engineering and Technology in 2007. **Chun-Hsi Huang** received his PhD from the State University of New York at Buffalo in Computer Science in 2001. He is currently an Associate Professor at the Department of Computer Science and Engineering of the University of Connecticut. He is affiliated with the International Society of Computational Biology (ISCB) and the American Medical Informatics Association (AMIA).

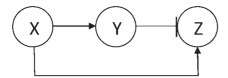


Figure I: Incoherent Type I Feedforward Loop (II-FFL) motif can act as a pulse generator. FFLs, in general, can work as an AND gate, OR gate, or even perform the SUM function (addition) in biological networks.

Type 1 FFL (C1-FFL) (where all interactions are positive) and the incoherent Type 1 FFL (I1-FFL) (X activates Z and also activates Y which represses Z, Figure 1) are found much more frequently in the transcription network of *Escherichia coli* and yeast than the other six types [6, 7]. In addition to the structure of the circuitry, the way in which the signals from X and Y are integrated by the Z promoter should also be considered. In most of the cases the FFL is either an AND gate (X and Y are required for Z activation) or OR gate (either X or Y are sufficient for Z activation) but other input functions are also possible.

There are many other prominent motifs like autoregulation, single input module, dense overlapping regulons and feedback loops, etc [4, 5]. Each motif has specific functions in the network. (The readers are referred to the references [4] and [5] for the functions and topologies of these motifs.) In many occasions, multiple motifs are found to be working in conjunction with one another, where output nodes of one motif are used as input to other motifs [12–15]. It is also shown that interactions in biological networks can be broken down into different levels of modularity, where each level shows a different distribution of network motifs [16, 17]. This is one way that network motifs can act as building blocks of biological networks, as proposed by Milo *et al.* [1].

Biologists are interested in knowing whether the functional behavior of a motif can be predicted from its structural topology as well as whether the abundance in appearance of such a motif necessarily implies biological significance. Some studies also investigated how network motifs might be shaped by evolution. For example, Kashtan *et al.* [17] demonstrated that when a network is placed under fixed environmental conditions, evolution optimizes the network topology for some specific functions, and no motifs form in this process. But, when the same network is placed under varying environmental

conditions where each condition demands different functional behavior from the network, several network motifs emerge. This happens since motifs—although having the same topology—are able to perform different tasks in different input conditions [5]. Yet other studies have argued that overabundance of a network substructure might be a secondary result of some other phenomena [18], or that network motifs might not have evolutionary traces [19].

All in all, the ability to computationally determine motifs in a given network is an essential step in furthering these research efforts. Given a network G and a set of random graphs, we need to identify all k-node (equivalently, size-k, throughout the article) sub-graphs that are statistically overrepresented in G. However, one of the difficulties is that determining if two graphs are topologically equivalent requires 'graph isomorphism' checking, a highly computation-intensive problem with no known polynomial-time solution. This problem is compounded by the fact that the number of sub-graphs of a given size in a network is exponential in both the network size and the sub-graph size. Moreover, real-life networks tend to be large and dense in many cases. The computations also need to be carried out on a large number of random graphs, typically ranging from hundreds to one thousand. By far, even the best-known algorithms cannot find motifs with more than 10 nodes in a large, dense network within a practical time frame without doing heuristics [20].

Motif finding algorithms use various strategies in order to overcome these difficulties. One of the notable strategies is the use of 'sub-graph sampling' through the target network instead of 'exact enumeration' to acquire an acceptable turn-around time. Another strategy is to generate all possible sub-graphs of a fixed size, and for each sub-graph count its frequency in the target network. The latter strategy, called 'motif-centric approach', can lead to reduction in isomorphism-related computations when coupled with other strategies, namely 'symmetry breaking' and 'mapping' [21] ('Mapping' and 'Symmetry breaking' sections). However, this strategy suffers when looking for larger motifs as the number of sub-graphs of a given size grows exponentially [16, 22-24].

Reviews for network motif finding algorithms exist in the literature [25, 26]. The review by Ciriello *et al.* [25] is specifically focused on PPI networks. On the other hand, the review by Ribeiro

Wong et al.

et al. [26] centers on algorithmic details of three tools. Our survey complements both with the following: (i) We discuss, with sufficient generality, various strategies that are applied in seven motif finding tools including two important algorithms, Kavosh [20] and MOtif Discovery Algorithm (MODA) [27], which have not been covered in those reviews. (ii) We present a simple classification scheme of motif finding algorithms based on the strategies these algorithms use. (iii) We discuss various performance-comparison experiments conducted in different published studies, and thereby conclude the strengths and weaknesses of different strategies. In addition, a few research directions are pointed out for future work.

NETWORK-MOTIF FINDING PROBLEM

Tasks involved in finding network motifs typically include the definition of frequency concepts, random graph generation, determining statistical significance of the frequency of a sub-graph and deciding subgraph isomorphism, etc. A few definitions follow.

Sub-graph frequency

Frequency here refers to the number of matches of a query sub-graph in a network [27]. Three different frequency concepts were discussed in Schreiber and Schwöbbermeyer [28, 29], F_1 , F_2 and F_3 , where F_1 allows arbitrary overlapping of nodes and edges between two sub-graphs; F_2 only allows node overlapping; and F_3 does not allow any overlapping of nodes or edges. Figure 2, Figure 3, and Table 1 illustrate variations in sub-graph frequency based on different frequency concepts.

Graph isomorphism

Two graphs are 'isomorphic' if there exists a one-to-one mapping between their nodes such that each edge in one graph can be mapped to an edge in the other graph [30]. The graph isomorphism problem—checking if two graphs are isomorphic—is one of a very small number of problems belonging to 'NP' which are neither known to be 'NP-Complete' nor known to be solvable in polynomial time [31]. The best known algorithm has a run-time of $2^{O(\sqrt{n\log n})}$ for graphs with *n* vertices [32, 33]. However, the sub-graph isomorphism problem checking if a network G contains an isomorph of another graph g—is NP-Complete

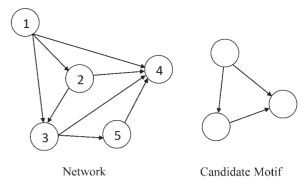


Figure 2: An example directed graph and a candidate motif.

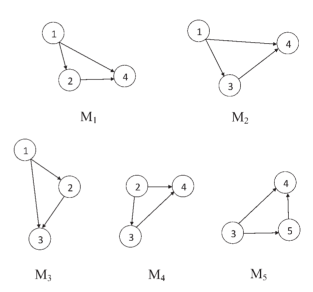


Figure 3: Instances of a candidate motif in the network given in Figure 2 using frequency concept Fl.

Isomorphism can be solved by using 'Canonical Labeling' of its nodes [35]. Practical algorithms such as the NAUTY [36] have been used in several motif finding tools for isomorphism testing.

Statistical significance

When frequencies of all size-k sub-graphs are known in both the input and random networks, different thresholds and metrics are applied to determine which sub-graphs are significantly frequent.

Thresholds

'Frequency threshold' requires that a motif occur at least a certain number of times in the input network. Sub-graphs with a lower frequency than the frequency threshold cannot be a motif. MAVisto, NeMoFinder and Kavosh ('Network-centric

Table I: Motif frequencies under different frequency concepts applied to the network shown in Figures 2 and 3

Concept		Edge overlap	Frequency	Matches
F _I	Yes	Yes	5	$\{M_1, M_2, M_3, M_4, M_5\}$
F_2	Yes	No	2	Either $\{M_1, M_4\}$ or $\{M_3, M_4\}$
F ₃	No	No	I	One of $\{M_1, M_2, M_3, M_4, M_5\}$

algorithms' section) utilize this metric. A sub-graph is 'frequent' if

$$f_{\text{input}} \ge F,$$
 (1)

where F is the frequency threshold and $f_{\rm input}$ is the sub-graph frequency in the input network. On the other hand, the 'uniqueness threshold' requires that the frequency of a candidate motif in the input network be at least a certain level higher than its mean frequency in the random graph ensemble. Let a size-k sub-graph g_k occur $f_{\rm input}$ times in the input network. Let $\overline{f}_{\rm random}$ be the mean of frequencies of g_k in the random networks. Then, g_k is 'unique' if

$$(f_{\text{input}} - \bar{f}_{\text{random}}) > U \times \bar{f}_{\text{random}},$$
 (2)

where U is the uniqueness threshold.

Significance metrics

Let a size-k motif g_k occur $f_{\rm input}$ times in the input network. Let $\overline{f}_{\rm random}$ and $\sigma^2_{\rm random}$ be the mean and variance of frequencies of g_k in a sufficiently large set of random networks, respectively. The z-score is defined as the difference of $f_{\rm input}$ and $\overline{f}_{\rm random}$, divided by the standard deviation $\sigma^2_{\rm random}$. Namely, the z-score is calculated as,

$$z(g_k) = \frac{f_{\text{input}} - \bar{f}_{\text{random}}}{\sqrt{\sigma_{\text{random}}^2}}.$$
 (3)

The P-value represents the probability of a motif to appear an equal or greater number of times in a random network than in the given input network. A motif is usually regarded as 'statistically significant' if the associated P-value is less than 0.01 or z > 2.0.

If the probability distribution of $f_{\rm random}$ (which, in many cases, is assumed to be Gaussian) is known, the probability of getting any particular z-value can be determined. If this probability is less than a given threshold P, the deviation is considered to be

significant with confidence. Specifically, a value of z is significant if,

$$\Pr\left(z\right) \le P \tag{4}$$

If Equation (4) is true with P = 0.01, frequency of g_k in G is significant with 99% confidence. Note, it has been pointed out that many properties of biological networks are non-Gaussian [25, 37], and probability distribution of f_{random} should be estimated from the input network through various estimation techniques [38].

Statistical significance can be measured in other ways. 'Abundance', Δ , is a metric similar to the z-score [15] and is defined as,

$$\Delta(g_k) = \frac{f_{\text{input}} - \bar{f}_{\text{random}}}{f_{\text{input}} + \bar{f}_{\text{random}} + \varepsilon},$$
(5)

where, ε is a small positive number preventing the ratio from approaching infinity when the frequencies are small. The value of Δ usually ranges between -1 (under-represented) and 1 (overrepresented). Another metric, the motif 'significance profile' (SP), is defined as a vector of z-scores of a particular set of motifs, which is normalized to a length of one [15]. Let n be the number of motifs in the set, and z_i be the z-score of the i-th motif. The motif SP of the i-th motif in the set is thus calculated as,

$$SP_i = \frac{z_i}{\sqrt{\sum_{j=1}^n z_j^2}}.$$
 (6)

SP of the entire network gives a histogram of the normalized z-scores of all possible motifs [15]. The 'concentration' of a candidate motif denotes how frequent it is in the network compared to other sub-graphs of the same size [39, 40]. Specifically, if there are n sub-graphs of size k in the network, the concentration of the i-th size-k sub-graph $g_{k,i}$ is defined as

$$C(g_{k,i}) = \frac{f_{k,i}}{\sum_{j=1}^{n} f_{k,j}},$$
(7)

where $f_{k,i}$ denotes the frequency of $g_{k,i}$ in the network.

Random graphs

Sub-graph frequencies are computed in both the input network and a set of topologically similar random networks to determine if the sub-graph is

Wong et al.

significantly frequent in the input network. Thus generating random networks is essential in network motif discovery. The 'Erdős–Rényi' (ER) model creates a random graph of n nodes by repeatedly and independently placing an edge between two nodes. The probability of picking any pair of nodes is uniform. The 'Barabási–Albert' (BA) model is an algorithm for generating random 'scale-free' networks using a 'preferential attachment' mechanism [41]. A scale-free network is a network whose degree distribution follows a power law in the following form:

$$P(k) = ck^{-r} \tag{8}$$

where P(k) is the fraction of nodes in the network with degree k, c is a normalization constant and r is a parameter usually between two and three. Scale-free networks are noteworthy because many empirically observed networks appear to be scale-free, including the World Wide Web, citation networks, biological networks, airline networks and some social networks. A preferential attachment process ensures that, when constructing a scale-free network, a new edge is more likely to be distributed (adjacent) to nodes that already have a higher degree.

It is essential for the generated random graph to resemble the original graph in terms of global properties such as the average degree, diameter, average path length, degree distribution and frequency of particular topological sub-structures, etc [1]. (Note the terms here follow the definitions in graph theory. For example, the 'average degree' is the average number of adjacent edges of the nodes in a graph; the 'diameter' of a graph is the maximum of the shortest paths of all vertex pairs, etc.)

The most common random graph generation technique used in motif finding algorithms is the 'Switching Method'. It repeatedly selects two random edges, $a \rightarrow b$ and $c \rightarrow d$, in the network, and exchanges the ends to form two new edges $a \rightarrow d$ and $b \rightarrow c$. The randomized graph has the same node and edge counts as in the input network, and the in- and out-degrees of the nodes are preserved. The switching method has also been widely applied in studying such features of biological networks as modularity and degree correlation. This method has a drawback that one cannot be certain when the network is adequately randomized. However, numerical studies have shown that, for many networks, 100*E times of switching appear to be adequate to achieve randomization, where E is the number of edges [3].

An alternative algorithm for random graph generation is the 'Matching Method', where each vertex is assigned a set of 'stubs'—the sawn-off ends of incoming and outgoing edges—according to the desired degree sequence. The in-stubs and out-stubs are then picked randomly in pairs and 'fused' to create the network edges. Although minor adjustments are needed to tend to real-world networks, where only a rather minority of nodes have a high degree, the matching algorithm usually correctly generates random networks with the desired properties [3].

The network motif-finding problem is thus defined as follows. Given the following inputs:

G: a network represented as a directed or undirected graph.

K: the maximum size of motif to search in G.

P: the required confidence level.

F: frequency threshold.

U: uniqueness threshold.

N: number of random networks.

Find all k-node sub-graphs $\{g_k\}$ with $2 \le k \le K$ occurring in G such that the frequency (or other metrics such as the concentration) of g_k in G is:

- (i) Above the given thresholds (Equations 1 and 2).
- (ii) Significantly higher than that in the random graphs with confidence level associated with *P* (Equation 4) in terms of either
 - (a) z-score (Equation 3), or
 - (b) Abundance (Equation 5), or
 - (c) SP (Equation 6).

STRATEGIES FOR MOTIF FINDING ALGORITHMS

In this section, we discuss several solution strategies for different parts of the motif-finding problem. There are three major parts of the problem: performing sub-graph census, solving graph isomorphism and ensuring that one sub-graph gets accounted for exactly once. The 'pattern growth' method is widely used in order to systematically explore or generate possible sub-graph variations. Strategies used in sub-graph census include exact 'enumeration', 'sampling' and 'mapping'. Some strategies for isomorphism checking include 'explicit checking' (using canonical labeling or NAUTY [36]) and 'symmetry breaking' coupled with 'mapping'. We present some representative motif finding algorithms under a

Sym-break, mapping

	Network centric	Sampling	Pattern growth	Mapping	Symmetry breaking	Isomorphism checking
MAVisto	Yes	No	Yes	No	No	Canonical labeling
NeMoFinder	Yes	No	Yes	No	No	Canonical labeling
Kavosh	Yes	No	Yes	No	No	NAUTY
MFinder	Yes	Yes	Yes	No	No	
FANMOD	Yes	Yes	Yes	No	No	NAUTY
Grochow	No	No	No	Yes	Yes	Sym-break, mapping

Yes

Yes

Table 2: Summary of strategies adopted by different algorithms

Yes

Yes

simple classification scheme based on the strategies the different algorithms use. These algorithms are MAVisto [28, 42], NeMoFinder [43], Kavosh [20], MFinder [39, 44], FANMOD [40, 45], Grochow and Kellis [21] and MODA [27]. Table 2 illustrates how different strategies can be combined to solve the motif-finding problem. Note that some comparisons made in this and the following sections are partially based on experimental results reported by articles accompanying the algorithms mentioned above. Web-based tools of some algorithms are freely available for scientific use and can be accessed at the following sites:

- MFinder: http://www.weizmann.ac.il/mcb/Uri Alon/groupNetworkMotifSW.html
- FANMOD: http://theinf1.informatik.uni-jena.de /~wernicke/motifs/index.html
- MAVisto: http://mavisto.ipk-gatersleben.de/

Pattern growth tree

MODA

No

A systematic way of generating a large set of variants from a base sub-graph is to extend it one step at a time by modifying one of its nodes or edges, and use the extended sub-graph to generate further variants. This sequence of action can be viewed as a tree (much like a search tree), where each node of the tree holds one sub-graph, and its children hold sub-graphs extended from that node so that a graph at a parent node is a sub-graph of all nodes in its sub-tree. This strategy, called 'pattern growth tree', can be used to systematically generate all possible size-k graphs starting with a size-k tree ('Generating all sub-graphs of a given size' section). It can also be used to systematically enumerate all occurrences of size-k sub-graphs in the target network.

Pattern growth strategy, when coupled with appropriate constraints, yields several benefits. When searching for size-k sub-graphs in the network, a

pattern growth tree built for size-(k-1) sub-graphs can be reused. This may lead to significant improvements in computational cost, which is exploited by MODA ('Motif-centric algorithms' section). Moreover, through a careful extension process one can guarantee that each sub-graph appears only once in the pattern growth tree, thus saving redundant computation. This is exploited by Kavosh ('Network-centric algorithms' section). Also, suppose the sub-graph at any node of the pattern growth tree violates a constraint. For example, its frequency falls below the frequency threshold, or becomes higher than the frequency of its ancestor (thus breaking downward closure property, if applicable). This information can be used to prune sub-trees rooted at that node, leading to increased computational efficiency. This is exploited by MAVisto ('Network-centric algorithms' section).

Sub-graph census: exhaustive versus sampling

Sub-graph census is the process of scanning the target network (node-by-node or edge-by-edge) and enumerating all occurrences of all sub-graphs of a given size. Usually this is done using a pattern growth tree. In general, the number of sub-graphs in a network increases exponentially when the network size increases: there are up to $|E_G|^{|E_g|}$ matches for a sub-graph g in the target network G under frequency concept F_1 , where $|E_G|$ and $|E_g|$ are edge counts in G and g, respectively [28]. The exhaustive census is therefore a time-consuming process, especially when the network or the sub-graph to be enumerated is large.

One can adopt a probabilistic approach by taking an adequate number of random size-k sub-graph samples from the original network. As is the case with probabilistic algorithms, with a good amount of trials one can expect to get closely accurate yet quick results, but with non-zero probability that 208 Wong et al.

some potential motifs will be missed. This is called the 'sampling strategy' for sub-graph census. Sampling saves much computation, thus allowing us to discover larger motifs. Moreover, sampling makes the algorithm insensitive to the size of the target network.

For each sub-graph in the pattern growth tree, an 'edge-sampling' strategy [39] randomly picks an edge from the network and selects a size-k sub-graph around that edge from the network for comparison. MFinder ('Algorithms with sampling—MODA' section) uses this strategy. However, it turns out that this is 'biased sampling': the probability of picking all size-k sub-graphs is not uniform in this strategy because sub-graphs having more edge are likely to be counted more, forcing the algorithm to assign specific weight to each sub-graph (and remember the sub-graph and the weight) to balance off the recounting, which leads to excessive memory usage. Edge extension in the pattern growth tree also encounters the same sub-graphs multiple times, causing redundant computation.

The 'node-sampling' strategy [40, 45], however, is able to ensure uniform probability of picking any size-k sub-graphs. It assigns to each level of its pattern growth tree a probability that the nodes (of the pattern growth tree) at that level will be further explored. It then probabilistically traverses the pattern growth tree, which guarantees that all the terminal nodes (size-k graphs) will be explored with equal probability, thus eliminating the memory drawback of edge-sampling strategy. Also, careful node extension in the pattern growth tree can ensure that a particular sub-graph will be encountered exactly once, which avoids redundant computation. FANMOD ('Network-centric algorithms' section) uses this strategy, as it is significantly efficient and faster than edge-sampling.

In some networks (termed scale-free) the sub-graph aggregation around high-degree nodes is higher than around low-degree nodes [46]. While not all biological networks are scale-free, this case was verified in empirical studies [27]. Thus, there is a variant of node-sampling strategy which probabilistically picks a node depending on its degree. MODA ('Motif-centric algorithms' section) uses this strategy.

Generating all sub-graphs of a given size

Algorithms for computing frequency of any 'given' sub-graph over the target network (as opposed to

sub-graph census, which computes frequency of all sub-graphs of a given size) must generate all possible non-isomorphic variants of that sub-graph. This can be done using a pattern growth tree, as in MODA [27], or by generating all isomorphic classes of a sub-graph using external tools [46], as in Grochow and Kellis [21]. This strategy has the drawback that when the size of the sub-graph gets >10, computations due to the number of possible variants become intractable [23, 47–49]. Another disadvantage is that this process generates and computes frequency for sub-graphs that might not occur in the target graph. One should note that isomorphism is partly resolved even before enumerating sub-graphs in the target network.

Mapping

The mapping strategy [21] of finding sub-graph instances takes a size-k sub-graph (candidate motif) and maps it onto the network in as many different places as it can. This is the opposite view of enumerating all size-k sub-graphs in the network and then computing frequency of a candidate size-k motif. A mapping strategy ranks the nodes in the target network based on their (and their neighbors') degree properties, and then looks for nodes in the input network that have similar characteristics of nodes in the candidate motif [21, 50]. This strategy is introduced by Grochow and Kellis [21] and is also used by MODA ('Motif-centric algorithms' section). Mapping, when used with symmetry breaking ('Symmetry breaking' section), resolves isomorphism without explicit checking [21].

Symmetry breaking

The symmetries of a graph are known as 'automorphisms', or self-isomorphisms. The group of automorphisms are in the same 'equivalence class'. One can specify a set of symmetry breaking conditions [21] for each equivalence class. Checking if these conditions hold between two graphs is easier than checking the isomorphism. Algorithms that use symmetry breaking while generating candidate motifs can prevent many isomorphism computations, thus increasing efficiency. This strategy is used alongside mapping. It was introduced by Grochow and Kellis, and was also used by MODA ('Motif-centric algorithms' section).

Motif finding algorithms

We discuss a simple framework for describing a network motif finding algorithm based on the strategies

 Table 3: Classification of network motif finding algorithms

	Network-centric	Motif-centric
Exact Search	MAVisto [28, 42] NeMoFinder [43] Kavosh [20]	Grochow and Kellis [21]
Sampling	MFinder [39, 44] FANMOD [40, 45]	MODA [27]

it uses. Under this framework, we discuss some representative algorithms (Table 3). Note these algorithms are selected not by their performance, but by the particular strategies they use. These algorithms are classified into two broad classes—'network—centric' and 'motif-centric' algorithms—which can be further classified depending on whether a sampling strategy is used while counting sub-graph frequencies.

Network-centric algorithms

Network-centric algorithms start with the network. They enumerate all sub-graphs with size k that occur in the target network. Network-centric algorithms have the benefit that sub-graphs that do not occur in the target network are never encountered. However, these algorithms cannot compute the frequency of a given sub-graph without doing sub-graph census.

Algorithms with exact census—MAVisto, NeMoFinder and Kavosh

MAVisto stands for 'Motif Analysis and Visualization Tool'. Developed by Schreiber and Schwöbbermeyer [42] in 2005, MAVisto is a network-centric algorithm that uses a pattern growth method called FPF (Flexible Pattern Finder) [28]. It uses the downward closure property commonly used in data mining, as well as a frequency threshold to prune branches of the pattern growth tree.

The 'Network Motif Finder', NeMoFinder, was developed in 2006 by Chen *et al.* [43]. Dedicated to undirected, unlabeled PPI networks, the NeMoFinder finds motifs that are maximal [51] (i.e. not contained in any other motif), are unique sub-graphs, but not necessarily induced. (Note H is an 'induced' sub-graph of G if it has exactly the edges that appear in G over the same vertex set.) It uses frequency and uniqueness thresholds, and the F_1 frequency concept. Notably, the input network is partitioned into smaller sub-networks for

enumeration of lower-level sub-graphs in the pattern growth tree, which makes it less-sensitive to large networks. The 'Canonical Adjacency Matrix' is used to resolve isomorphism through a method called 'Graph Cousins' [27]. This method, however, is likely to produce some redundant isomorphic candidate motifs [27].

Kashani *et al.* [20] developed Kavosh in 2009. It's a network-centric algorithm using the pattern growth tree (with node-extension) to enumerate all size-k sub-graphs in the input network. Kavosh adopts the 'Revolving Door algorithm' [52] for traversing the pattern growth tree, ensuring that every candidate motif is encountered exactly once. The frequency concept F_1 is used, along with a frequency threshold. In addition, Kavosh uses the NAUTY algorithm [36] ('Graph isomorphism' section) for isomorphism testing, which is efficient as the NAUTY does not generate any redundant candidate motif [36].

Algorithms using sampling—Mfinder and Fanmod

MFinder was developed in 2005 by Kashtan *et al.* [44], which showcases their edge-sampling algorithm [39]. The pattern growth tree is used to enumerate all sub-graphs in the network through edge extension. Frequency concept F_1 is used, but motifs are required to be induced sub-graphs. MFinder also uses concentration as the significance metric. MFinder is seriously handicapped with drawbacks of edge-sampling, and is not suitable for finding large motifs [26]. MFinder can also perform exhaustive sub-graph census instead of sampling, as an option. In this regard, the MFinder functions as the enumeration algorithm [1].

FANMOD was developed in 2006 by Wernicke *et al.* [40, 45], which uses a node-sampling strategy and a pattern growth tree using node-extension. FANMOD is fast, and can detect up to size-8 motifs in both directed/undirected networks. It uses F_1 frequency concept, and motifs are required to be induced. FANMOD uses concentration as the significance metric and NAUTY [36] for isomorphism checking. FANMOD's randomized enumeration algorithm is known as Rand-ESU. FANMOD, like MFinder, is also able to do exhaustive census over the target network.

Motif-centric algorithms

Motif-centric algorithms are able to compute frequency of any given sub-graph in the target

2I0 Wong et al.

network. This enables them to directly verify whether the query sub-graph is a motif. In order to compute frequencies of all size-k graphs, a motif-centric algorithm first enumerates all possible size-k graphs. As network-centric algorithms, motif-centric algorithms can be further classified depending on whether sampling is used over the target network while counting sub-graph frequencies.

Algorithms with exact search—Grochow

Devised by Grochow and Kellis [21] in 2007, this algorithm is the first example of a motif-centric algorithm. Grochow uses symmetry breaking with mapping to count the frequency of a query sub-graph. It uses the 'geng' and 'directg' packages by McKay [53–55] to exhaustively generate all non-isomorphic sub-graphs of size-k; it then checks if any of them is a motif. Grochow is able to exhaustively find seven-node motifs in the *Saccharomyces cerevisiae* PPI network efficiently. Although mapping is an exact strategy, it is possible to first use sampling to pick a random (potentially large) sub-graph from the network, and then use symmetry breaking and mapping to determine its frequency.

Algorithms with sampling—MODA

MODA, was developed by Omidi et al. [27] in 2009. This motif centric algorithm uses a pattern growth tree, called 'expansion tree', which uses symmetry breaking to generate unique extensions. It then uses mapping for the first level of the expansion tree. Information from the previous level is then used to efficiently enumerate the extensions for each subsequent level. Note that MODA implements an improved node-sampling strategy in every level—even for mapping, as opposed to Grochow. Moreover, the expansion tree enables it to reuse computations done for smaller-size query motifs, resulting in its high efficiency. Largely due to this combination of efficient strategies, MODA is fast and can handle rather large networks and motifs. It also allows arbitrary overlap between sub-graphs, using frequency concept F_1 .

PERFORMANCE COMPARISONS

Selected results from various comparative studies of motif finding algorithms are presented in Tables 4, 6 and Figures 4–7

Table 4: Execution times (in seconds) of Kavosh, FANMOD, MAVisto and MFinder for sub-graphs of size 3–10 in various networks

Network	k	Kavosh	FANMOD	MAVisto	MFinder (exhaustive)
E. coli	3	1	1	14,000	30
	4	2	3		300
	5	15	16		23,000
	6	140	130		
	7	1400	1200		
	8	13,000	9000		
	9	100,000			
	10	1,000,000			
Social	3	1	1	400	10
	4	1	1	1500	50
	5	2	3		800
	6	10	20		180,000
	7	70	100		
	8	400	800		
	9	2600			
	10	15,000			
S. cerevisiae	3	2	3	16,000	30
	4	30	40		300
	5	1000	1100		34,000
	6	20,000	24,000		
	7	750,000	900,000		
	8	17,000,000	19,000,000		
	9	300,000,000			
	10	7,000,000,000			

Note: Here k denotes a sub-graph size. Note that large numeric values have been rounded to improve visualization (Courtesy of Kashani et al. [20]).

Sampling strategies for sub-graph census are faster

FANMOD is efficient in computation and tends to produce results while other exact algorithms (except Kavosh) struggle (Figure 5, Tables 4 and 6). Exact sub-graph census becomes slow as either network or motif size increases, as demonstrated in MAVisto on various occasions (Figures 4 and 5, Table 4). However, NeMoFinder, which is a network-centric tool doing exact census, shows promise in finding large motifs in large PPI networks (Figure 4) as the algorithm works by partitioning the input network and thus effectively reducing the problem size. Similarly, Kavosh, yet another network-centric algorithm with exact census, is sometimes faster than FANMOD and can find larger motifs than FANMOD (Table 4), largely due to the highly efficient sub-graph enumeration strategy of Kavosh and its use of NAUTY [36] for isomorphism checking ('Network-centric algorithms' section). Both MODA and FANMOD use

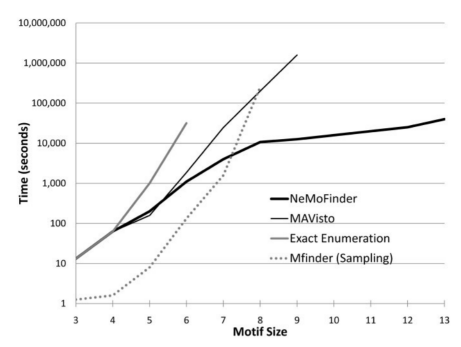


Figure 4: Execution times (in seconds) of NeMoFinder, FPF algorithm (MAVisto), MFinder (sampling) and full enumeration algorithm for sub-graphs of sizes 3–I3 in a PPI network of S. cerevisiae [54] with 1004 nodes and 957 edges, as reported by Chen et al. [43].

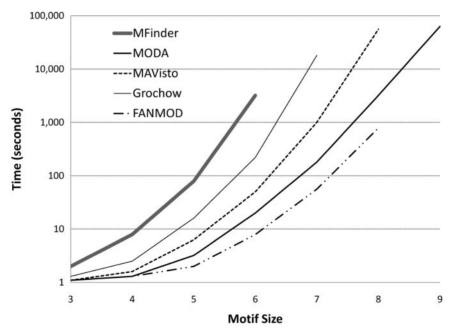


Figure 5: Execution times (in seconds) of MODA, MFinder, Grochow, FPF algorithm and FANMOD for sub-graphs of sizes 3–9 in an *E. coli* transcription network used by Shen-Orr et al. [2], which is a directed graph of II6 nodes and 477 edges (Courtesy of Omidi et al. [27]). Algorithms were executed on the input network only.

the sampling strategy for sub-graph census. FANMOD is faster than MODA (Figure 5) since MODA, a motif-centric tool, considers more sub-graphs than FANMOD. However, MODA

can find larger motifs than FANMOD (Figure 5), partly because of a more efficient isomorphism-checking strategy by mapping and symmetry breaking, and partly because of its ability to reuse

2l2 Wong et al.

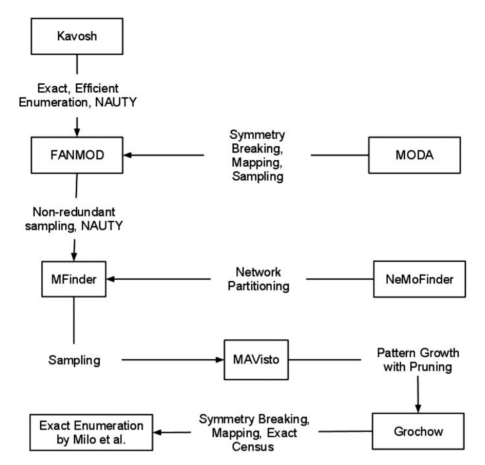


Figure 6: How the combination of strategies affects performance of an algorithm. An arrow from algorithm A to algorithm B implies that A usually outperforms B for larger networks and motifs. Each arrow is labeled with the strategies used in A that contribute to the dominance.

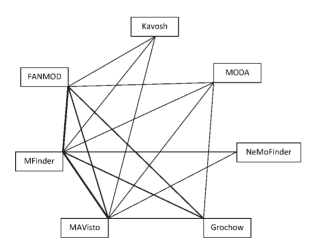


Figure 7: Availability of information on the comparisons of different methods based on Figures 4 and 5, and Tables 4 and 6. The thickness of lines between each pair of algorithms indicates the number of experiments involved to derive Figure 6.

results from earlier computations ('Motif-centric algorithms' section).

Motif-centric algorithms tend to slow down as sub-graph size grows

The effect is observable in all algorithms, but more so in motif-centric algorithms such as the Grochow (Table 6 and Figure 5). Grochow is more efficient than the exact enumeration algorithm [1] due to its symmetry breaking techniques that enables it to reduce redundant counts of sub-graphs. Yet Grochow is found to be slower than MAVisto (Figure 5) and the exhaustive version of MFinder (Table 6), highlighting the limitation of generating all possible size-*k* sub-graphs. However, as mentioned previously, MODA could detect larger motifs than does FANMOD, although FANMOD is faster (Figure 5). Nevertheless, motif size is a bottleneck for motif-centric tools.

Table 5: Input networks used in the comparisons made in Table 6

Network	Nodes	Edges	Description
Circuit	252	399	Electronic circuit S. cerevisiae transcriptional network Social network with heterogeneous communities
Yeast	688	1079	
Social	1000	15541	

Table 6: Execution times (in seconds) of MFinder, FANMOD and Grochow, all with exhaustive census on the input networks

Network	k	MFinder	FANMOD	Grochow
Circuit	4	0.10	0.05	0.36
	5	0.36	0.14	15.79
	6	2.45	0.67	2519.3
	7	17.54	4.34	>4h
	8	130.66	27.46	>4h
	9	MEM	140.66	>4h
Yeast	4	2.42	0.77	2.25
	5	55.67	15.21	107.82
	6	MEM	260.38	>4h
	7	MEM	5023.91	>4h
Social	4	40.44	11.22	16.98
	5	MEM	281.40	1122.74
	6	MEM	7951.47	>4h

Note: Here k indicates the motif size. MEM indicates the out-of-memory situation (Courtesy of Ribeiro et al. [26]).

MAVisto and MFinder are limited in motif size

The MFinder stores all sub-graphs in memory and considers the same sub-graph many times. Thus MFinder scales poorly with motif size despite its use of sampling for sub-graph census. Table 6 shows that MFinder (exact census) takes more memory and time than FANMOD (exact census), since FANMOD counts a sub-graph only once. Similar observations can be made in Figure 5 and Table 4. MAVisto, on the other hand, performs poorly (Figures 4, 5 and Table 4) mainly because it does exact census in a less sophisticated way than Kavosh does.

Which tool should one use for larger motifs?

Although current motif finding algorithms are able to find only small motifs when the target network is large and/or dense, one may argue that if an exact result is necessary, Kavosh will be a desirable tool. Otherwise, FANMOD or MODA will be a good

choice. In addition, NeMoFinder is a good choice for large PPI networks.

What are the bottlenecks for performance?

The most obvious bottleneck is the number of random graphs used for comparison. While computing over a large number of random graphs will statistically justify the overrepresentation of motifs, it is clearly a time-consuming process. The second bottleneck is the target network size. Real-world networks tend to be large, and doing exact sub-graph census is not always practical. The third bottleneck is the sub-graph size. When looking for motifs with more than 8–10 nodes, the number of possible sub-graphs becomes prohibitively large (depending on the available computational resources). It affects motif-centric algorithms in particular, and all algorithms in general when the target network is large and dense.

Figure 6 illustrates which strategy makes an algorithm better than others. In this figure, rectangles denote algorithms and arrows denote the dominance between two algorithms. Each arrow is labeled with the strategies that result in the superior performance. We placed MODA over FANMOD in Figure 6 since MODA can find larger motifs as shown in Figure 5, although their execution times are comparable. From this figure one can get a quick understanding of how strategies discussed in 'Strategies for motif finding algorithms' section affect the performance of algorithms. There are no arrows among Kavosh, MODA, and NeMoFinder as they have not been comparatively studied in the literature.

CONCLUSIONS AND FUTURE DIRECTIONS

Solving the network motif-finding problem involves several aspects in research—generating random graphs, checking graph isomorphism, estimating sub-graph frequencies in the network (by exact enumeration or sampling) and generating the set of candidate motifs, etc. Each has challenges and possibilities of its own. While several algorithms exist in the literature with experimental results available on selected target networks, the relative performance when varying other topological features of the networks such as clustering coefficient, scale-freeness, etc need to be further studied to conclude their strengths, weaknesses and the practical relevance.

2l4 Wong et al.

One research direction toward improving these algorithms is to devise a robust, parallel algorithm for a distributed computation platform that could simultaneously analyze different parts of the network, thus reducing the execution time and enabling our reach to larger motifs and networks.

While checking isomorphism for general graphs involves work in exponential time, polynomial-time algorithms do exist for special graphs such as the planar graphs, interval graphs, and the two-connected series-parallel graphs, etc [55]. If some biologically verified (large) motifs have similar topological characteristics that enable a fast isomorphism test, one could prioritize expected outputs of a motif finding algorithm to generate those with such topologies first. This will allow us to generate highly relevant (large) motifs early, if they exist, during the computation process.

Another improvement could be made by reducing unnecessary computations over the random graphs. After performing sub-graph census over the input network, the set of possible motifs is already limited to the sub-graphs found in the census. The same computation over hundreds (or thousands) of random networks is then performed to compute frequencies for sub-graphs, many of which may not occur in the target networks. Such computations can be saved by a more efficient data structure to keep track of candidate motifs.

Key Points

- Network motif finding algorithms can be divided into network-centric and motif-centric groups.
- Network motif finding algorithms can be further classified based on their use of probabilistic sampling of the target network.
- Efficient combination of different strategies leads to a better performance.

FUNDING

This work was supported by the National Science Foundation [CCF-0755373 to E.W., B.B., S. Q. and C.-H.H.]; and the National Institutes of Health [R13-LM008619 to C.-H.H.].

References

 Milo R, Shen-Orr S, Itzkovitz S, et al. Network motifs: simple building blocks of complex networks. Science 2002; 298:824–27.

- Shen-Orr S, Milo R, Mangan S, et al. Network motifs in the transcriptional regulation network of Escherichia coli. Nat Genet 2002;31:64–8.
- Milo R, Kashtan N, Itzkovitz S, et al. On the uniform generation of random graphs with prescribed degree sequences. Arxiv preprint, cond-mat/0312028v2, 2004.
- Shoval O, Alon U. SnapShot: Network Motifs. Cell 2010; 143:326.e1.
- 5. Alon U. Network motifs: theory and experimental approaches. *Nat Rev Genet* 2007;**8**:450–61.
- Mangan S, Alon U. Structure and function of the feed-forward loop network motif. Proc Natl Acad Sci USA 2003;100:11980–85.
- Mangan S, Itzkovitz S, Zaslaver A, et al. The incoherent feed-forward loop accelerates the response time of the gal system of Escherichia coli. JMol Biol 2006;356:1073–81.
- Lee TI. Transcriptional regulatory networks in Saccharomyces cerevisiae. Science 2002;298:799–804.
- Ratushny AV, Ramsey SA, Roda O, et al. Control of transcriptional variability by overlapping feed-forward regulatory motifs. Biophys J 2008;95:3715–23.
- Kalir S, Mangan S, Alon U. A coherent feed-forward loop with a SUM input function prolongs flagella expression in Escherichia coli. Mol Syst Biol 2005;1:2005.0006.
- Setty Y, Mayo AE, Surette MG, et al. Detailed map of a cis-regulatory input function. Proc Natl Acad Sci USA 2003; 100:7702–07.
- Dobrin R, Beg QK, Barabási A-L, et al. Aggregation of topological motifs in the Escherichia coli transcriptional regulatory network. BMC Bioinformatics 2004;5:10.
- Kim J-R, Yoon Y, Cho K-H. Coupled feedback loops form dynamic motifs of cellular networks. *Biophys J* 2008; 94:359–65.
- 14. Kashtan N, Itzkovitz S, Milo R, et al. Topological generalizations of network motifs. *Phys Rev E* 2004;**70**:31909.
- Milo R, Itzkovitz S, Kashtan N, et al. Superfamilies of evolved and designed networks. Science 2004;303:1538– 42.
- Itzkovitz S, Levitt R, Kashtan N, et al. Coarse-graining and self-dissimilarity of complex networks. Phys Rev E 2005;71: 016127.
- Kashtan N, Alon U. Spontaneous evolution of modularity and network motifs. Proc Natl Acad Sci USA 2005;102: 13773–78.
- Solé RV, Valverde S. Are network motifs the spandrels of cellular complexity? *Trends Ecol Evol* 2006;21:419–22.
- Bottani S, Vergassola M, Mazurie A. An evolutionary and functional assessment of regulatory network motifs. *Genome Biol* 2005;6:R.35.
- Kashani ZRM, Ahrabian H, Elahi E, et al. Kavosh: a new algorithm for finding network motifs. BMC Bioinformatics 2009;10:318.
- Grochow JA, Kellis M. Network motif discovery using sub-graph enumeration and symmetry-breaking. Res Comp Mol Biol 2007;4456:92–106.
- 22. Kuramochi M, Karypis G. Frequent sub-graph discovery. In: *Proceedings of IEEE International Conference on Data Mining* 2001, San Jose, California, 313–20.
- 23. Sloane NJA, Plouffe S. Number of graphs on n unlabeled nodes. (A000088, Formerly M1253 N0479). The Online Encyclopedia of

- Integer Sequences 2010. http://oeis.org (15 January 2011, date last accessed).
- Han J, Kamber M. Data Mining: Concepts and Techniques.
 2nd edn. Morgan Kaufmann (an imprint of Elsevier), 2006.
- 25. Ciriello G, Guerra C. A review on models and algorithms for motif discovery in protein–protein interaction networks. *Brief Funct Genomics* 2008;**7**:147–56.
- Ribeiro P, Silva F, Kaiser M. Strategies for network motifs discovery. In: Proceedings of the 5th IEEE International Conference on e-Science 2009, Oxford, UK, 80–87.
- Omidi S, Schreiber F, Masoudi-Nejad A. MODA: an efficient algorithm for network motif discovery in biological networks. *Genes Genetic Syst* 2009;84:385–95.
- Schreiber F, Schwöbbermeyer H. Frequency concepts and pattern detection for the analysis of motifs in networks. *Lect Notes Comput Sci* 2005;3737:89–104.
- 29. Kuramochi M, Karypis G. Finding frequent patterns in a large sparse graph. *Data Min Knowl Disc* 2005;**11**:243–71.
- Fortin S. The Graph isomorphism problem. Technical report 1996, Department of Computing Science, University of Alberta, Edomonton, Alberta, Canada, 1996.
- Garey M, Johnson D. Computers and Intractability: A Guide to the Theory of NP-Completeness. New York: W. H. Freeman and Co., 1979.
- 32. Babai L, Codenotti P. Isomorphism of hypergraphs of low rank in moderately exponential time. In: *Proceedings of the 49th Annual IEEE Symposium on Foundations of Computer Science* 2008, Philadelphia, PA, USA, 667–76.
- 33. Johnson DS. The NP-completeness column. *ACM Trans Algorithms* 2005;1:160–76.
- 34. Cook SA. The complexity of theorem-proving procedures. In: *Proceedings of the third Annual ACM Symposium on Theory of Computing* 1971, Shaker Heights, OH, USA, 151–8.
- 35. Babai L, Luks EM. Canonical labeling of graphs. In: Proceedings of the Fifteenth Annual ACM symposium on Theory of Computing 1983, Boston, MA, USA, 171–83.
- 36. McKay B. *The NAUTY Page*. http://cs.anu.edu.au/~bdm/nauty/ (5 February 2011, date last accessed).
- 37. Ziv E, Koytcheff R, Middendorf M, et al. Systematic identification of statistically significant network measures. *Phys Rev E* 2005;**71**:016110.
- Hastie T, Tibshirani R, Friedman J. The Elements of Statistical Learning: Data Mining, Inference and Prediction. 2nd edn. Springer NYC, NY, USA: Springer Series in Statistics, 2009.
- Kashtan N, Itzkovitz S, Milo R, et al. Efficient sampling algorithm for estimating sub-graph concentrations and detecting network motifs. Bioinformatics 2004;20:1746–58.
- 40. Wernicke S, Rasche F. FANMOD: A tool for fast network motif detection. *Bioinformatics*, 2006;**22**:1152–3.

- 41. Barabási A. Emergence of scaling in random networks. *Science* 1999;**286**:509–12.
- Schreiber F, Schwöbbermeyer H. MAVisto: a tool for the exploration of network motifs. *Bioinformatics* 2005;21: 3572–4.
- 43. Chen J, Hsu W, Le ML, et al. 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 2006, Philadelphia, PA, USA, 106–15.
- Kashtan N, Itzkovitz S, Milo R, et al. Network motif detection tool Mfinder tool guide. Technical report 2005, Departments of Molecular Cell Biology and Computer Science and Applied Mathematics, Weizmann Institute of Science, Rehovot, Israel, 2005.
- Wernicke S. A faster algorithm for detecting network motifs. Algorithms Bioinformatics 2005;3692:165–77.
- Vázquez A, Dobrin R, Sergi D, et al. The topological relationship between the large-scale attributes and local interaction patterns of complex networks. Proc Natl Acad Sci USA 2004;101:17940–45.
- 47. Harary F, Palmer EH. Graphical Enumeration. New York: Academic Press, 1973.
- Inokuchi A, Washio T, Motoda H. An Apriori-based algorithm for mining frequent substructures from graph Data. Principles Data Mining Knowl Disc 2000;1910:13–23.
- Kuramochi M, Karypis G. An efficient algorithm for discovering frequent sub-graphs. IEEE Tran Knowl Data Eng 2004;16:1038–51.
- Ullmann JR. An algorithm for subgraph isomorphism. JACM 1976;23:31–42.
- Huan J, Wang W, Prins J, et al. Spin: mining maximal frequent sub-graphs from graph databases. In: Proceedings of the 10th ACM SIGKDD International Conference on Knowledge Discovery and Data Mining 2004, Seattle, WA, USA, 581–6.
- 52. Kreher DL, Stinson DR. Combinatorial Algorithms: Generation, Enumeration and Search. Boca Raton, FA, USA: CRC Press, 1999;329.
- 53. McKay B. Isomorph-free exhaustive generation. *JAlgorithms* 1998:**26**:306–24.
- 54. Uetz P, Giot L, Cagney G, et al. A comprehensive analysis of protein–protein interactions in Saccharomyces cerevisiae. Nature 2000;403:623–7.
- Lepistö T, Salomaa A, Lingas A, et al. A polynomial-time algorithm for sub-graph isomorphism of two-connected series-parallel graphs. In: Proceedings of the 15th International Colloquium on Automata, Languages and Programming (ICALP) 1988, Tampere, Finland, 394–409.