Systematic identification of statistically significant network measures

Etay Ziv,1,2 Robin Koytcheff,3 Manuel Middendorf,4 and Chris Wiggins3,5

1College of Physicians and Surgeons, Columbia University, New York, New York 10027, USA
2Department of Biomedical Engineering, Columbia University, New York, New York 10027, USA
3Department of Applied Physics and Applied Mathematics, Columbia University, New York, New York 10027, USA
4Department of Physics, Columbia University, New York, New York 10027, USA
5Center for Computational Biology and Bioinformatics, Columbia University, New York, New York 10027, USA

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We present a graph embedding space (i.e., a set of measures on graphs) for performing statistical analyses of networks. Key improvements over existing approaches include discovery of “motif hubs” (multiple overlapping significant subgraphs), computational efficiency relative to subgraph census, and flexibility (the method is easily generalizable to weighted and signed graphs). The embedding space is based on scalars, functionals of the adjacency matrix representing the network. Scalars are global, involving all nodes; although they can be related to subgraph enumeration, there is not a one-to-one mapping between scalars and subgraphs. Improvements in network randomization and significance testing—we learn the distribution rather than assuming Gaussianity—are also presented. The resulting algorithm establishes a systematic approach to the identification of the most significant scalars and suggests machine-learning techniques for network classification.

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BACKGROUND

Recent studies of real-world biological, social, and technological networks have catalyzed an explosion of research from a broad range of disciplines. Much of the effort in this emerging field has focused on characterizing the structure of networks using various statistical properties that are local (analysis relies on subset of nodes) or global (relying on all nodes) in scope. The former analysis includes subgraph census (comparing frequency of subgraph occurrences in a given graph with those over a distribution of graphs [1,2]), while examples of the latter include path lengths and degree distributions (see citations in [3]).

To study local structure statistics, sociologists developed the k-subgraph census, an enumeration of all possible subgraphs of k nodes appearing in networks. For example, sociologists used the three-subgraph census, compared with three-subgraph distributions in randomized graphs, to quantify network transitivity [4–6] (in the context of a social network, high transitivity means that many of your friends are friends with each other). Applying such techniques first to the Escherichia coli genetic network [2] and later to various biological and physical networks [7], Milo et al. showed that different networks have different “most significant” subgraphs. Major limitations of these subgraph approaches include computational cost and generalizability. The number of isomorphism classes of digraphs grows rapidly with graph size [5,8] and subgraph isomorphism is an NP-complete problem [9].1 These computational limitations bias results, since structures with more than three or four nodes would not be counted. Moreover, it is not readily obvious how to extend subgraph census to weighted and/or signed graphs. This is particularly relevant for genetic regulatory networks in which the interactions can be described quantitatively via binding affinities and qualitatively as activating or repressing, or similarly neuronal networks, in which the interactions are often weighted by the number of synapses between neurons and can also exhibit excitatory and inhibitory behaviors.

In their ground-breaking work, Shen-Orr et al. identified three significant motifs in the E. coli genetic network. However, rather than counting all structures up to a given size, the authors had to resort to posing putative significant structures, thus making prior assumptions about which subgraphs are important. One topology was found by enumerating all three-node subgraphs in the network; a second by searching for single regulator genes regulating at least 13 distinct operons; and a third by presenting a clustering algorithm based on several new parameters. Similarly, in [10] six different subgraphs were defined using six different algorithms. Rather than finding subsets of motifs via tailored, parametrized, and thresholded algorithms, a single, generalizable method for identifying motifs is needed.

In this paper, we first present an embedding space for networks, which is (i) computationally efficient as compared to subgraph census for naturally occurring networks and (ii) easily applicable to weighted and signed graphs. We then employ this space in a single, generalizable algorithm to discover arbitrarily large, statistically significant network measures in the E. coli and the Saccharomyces cerevisiae genetic regulatory networks. Results are presented about the structure of these networks, including the presence of overlapping significant subgraphs. We also introduce a randomization scheme that generates independent identically distributed samples rather than a Markov chain, and we integrate density estimation into our significance testing rather than asserting Gaussianity.

1Digraphs (directed graphs, or graphs whose edges have directionality) are isomorphic if there exists a relabeling of their vertices such that the two graphs are identical. The NP class consists of decision problems whose solution can be found in polynomial time on a nondeterministic Turing machine. NP-complete problems are a subset of NP which are particularly hard. Given two graphs G1 and G2, subgraph isomorphism asks if G1 is isomorphic to a subgraph of G2.
MOTIVATION

As a motivating example, we consider the three-subgraph \( T_{030} \) defined as the triad of nodes \( i, j, \) and \( k \) and edges \( i \to j, j \to k, i \to k \) [see Fig. 1(a)], which we represent by its adjacency matrix \( A \) (\( A_{ij} = 1 \) if the \( j \)th node is the parent of the \( i \)th node, and 0 otherwise). This nomenclature references the earliest work in subgraph census [1,5]. We observe that the number of \( T_{030} \) subgraphs in this graph is trivially found by simple matrix manipulation of its adjacency matrix as follows. The trace of the square of \( A \) is constructed from the letter \( A T \). We define a "word" to be the matrix built from the letters \( A, A^T, D, U \) and \( U \) as the integer obtained from the adjacency matrix \( A \) as follows. The diagonal projection operator \( D \) and its complement \( U \) are defined for any matrix \( Q \) by \( D(Q)_{ij} = Q_{ij} I_{ij} \) and \( U(Q) = Q - D(Q) \), respectively. Note that we do not use Einstein’s summation convention and \( D \) is not the trace.

\[ A, A^T, D, U \] can be visualized as motion on the digraph: \( A \) and \( A^T \) represent moving one step forward or backward, respectively; \( D \) represents restriction to closed paths; \( U \) represents open paths. In terms of the adjacency matrix and the functions that act on it, we can enumerate \( T_{030} \) as \( \Sigma \{DA^TAA\} \). Reading this expression from right to left, we start at a node, move two steps forward \( (A^T) \), then one step backward \( (A) \), and arrive at the original starting node \( (D) \). By summing all \( n^2 \) elements of the resulting matrix, we obtain a count of \( T_{030} \). Instead of summing, we could also count the number of nonzero elements, \( N \). These operations on the resulting matrix, \( \Sigma \) or \( N \), yield the number of distinct paths between all pairs of end points or the number of distinct pairs of end points, respectively.

We define a "word" to be the matrix built from the letters \( A, A^T, D, U \) and \( U \) as the integer obtained from the operations \( \Sigma \) or \( N \) on a word. An enumeration of words and sub-sequent evaluation of scalars allows us to embed a given network in an infinite-dimensional space. To enumerate words we systematically combine letters. Obvious redundancies can be eliminated (e.g., \( U^2 = U, D^2 = D, UD = DU = 0 \)). To enumerate words we combine letters such that each letter acts on everything to its right. As an example, the word \( D(A^T(A(A))) \) is constructed from the letter \( D \) acting on \( A^T \) acting on \( A \) acting on \( A \). The scalar is obtained by evaluating either \( \Sigma \) (the sum over) or \( N \) (the number of nonzero elements in) the word. Other choices for construction of words are possible [e.g., using different combinations of parentheses, \( D(A^T(A(A))) \)]. Our method can easily be generalized to include these words. For simplicity, herein we will assume parentheses are implicit and write words without the parentheses. Thus, \( D(A^T(A(A))) \) will be written \( DA^TAA \).

PROPOSAL

Given this embedding space for networks, essentially a set of measures on a network, one can then employ standard tools from statistics and machine learning to characterize a network of interest. For the specific application of identifying statistically significant features of a network, this tantalizing observation motivates the technique presented here: (1) systematically enumerate words; (2) evaluate the scalars obtained from these words for a graph of interest; (3) compare scalars with the distribution obtained by evaluating scalars over a randomly generated distribution of matrices, thus finding statistically significant scalars. The fact that scalars are based on combinations of functions of the adjacency matrix makes our method easily extendable to weighted and signed graphs. For example, in the former one could simply use the weight matrix in place of the adjacency matrix; in the latter, one could use two adjacency matrices representing the two types of interactions.

As stated above, a major limitation of subgraph census is computational efficiency. Here we present analytic and numerical comparisons between subgraph census and our scalar technique. Traditional algorithms count subgraphs by performing walks [7,11]. Given a graph with \( N \) nodes and \( M \) edges, the computational cost of subgraph counting grows exponentially in the size of the subgraph \( n \), worse than exponentially in the density \( M/N \), and is traditionally infeasible for \( n > 4 \), especially in scale-free networks [7,11,12]. In scalar calculation, computational complexity is upper bounded by \( N^2 \Sigma \ell \ell \), where \( \ell \) is the number of letters in scalar \( i \) and the sum is performed over all scalars. While complexity grows exponentially in the number of letters, the exponential term is independent of the density and the degree distribution. Thus feature selection using scalars is especially suited for dense, clustered, or scale-free networks.

This observation is particularly relevant as many naturally occurring networks have heterogenous degree distributions [13]. To quantify the extent of degree distributions on the performance of the two algorithms, we benchmark the subgraph census against our scalars method using randomly generated networks. We generate multiple graphs of the same size and density as the \( E. \) coli genetic regulatory network, but with different degree distributions, using the class of growing random network (GRN) models with tunable parameter \( \gamma \), first proposed by Krapivsky et al. [14] as a generalization of the cumulative advantage or preferential attachment models [15,13]. In the GRN model, at every time step a new node is added, and with probability \( A_k \) an edge is created between the new node and an existing node with \( k \) edges, where \( A_k = k^\gamma \). The preferential attachment parameter \( \gamma \) acts to tune the degree of heterogeneity in the degree distribution. As \( \gamma \) approaches 1, or linear preferential attachment, the degree distribution becomes more heavy tailed.
and thus more similar to naturally occurring networks. In Fig. 2 we show degree distributions for graphs generated at three different values of $\gamma$. In Fig. 3, we demonstrate how the scalars method significantly improves computational time for these types of degree distributions, which many biological (as well as technological and sociological) networks evidence.

FIG. 2. Degree distributions of networks generated using the Barabasi and Albert preferential attachment model with the tunable parameter $\gamma$ such that $A_k = k^\gamma$, where $A_k$ is the probability of a new vertex attaching to an existing vertex with $k$ links. All of these networks are the same size and have the same density (423 nodes, 519 edges), but differ in their degree distributions.

FIG. 3. A numerical experiment comparing efficiency of “traditional” subgraph counting algorithm (circles) and the proposed “scalars” algorithm (triangles), as a function of $\gamma$, a parameter which tunes the degree of scale invariance in the network (see Fig. 2). The number of nodes and the density in the networks were kept constant and equal to those of the E. coli network tested in the paper. Scale-free properties similar to naturally occurring networks emerge with linear preferential attachment, where $\gamma=1$ (e.g., at $\gamma \sim 1$ the network contains hubs whose degree is similar to the degree of hubs in the E. coli network). We see here that the scalars algorithm becomes more efficient at $\gamma>0.7$.

FIG. 4. A histogram of the kurtosis (a measure of the degree of peakedness of a distribution) for scalars demonstrates many non-Gaussian distributions (i.e., distributions with kurtosis greater than or less than 3). This is also the case for subgraph distributions and hence we employ density estimation rather than assuming Gaussianity.

BACKGROUND ENSEMBLE

A vast literature discusses different randomly generated network models [3,5,16–21]. In [2] a random model was used which preserved $N(k, k_s)$: the in degree and out degree of each node (random matching of a given in- and out-degree sequence is also known as the configuration model [3,20]). This can be done efficiently by representing the graph as ordered lists of parents and children. The number of times the node occurs in the parent (child) list is the node’s out (in) degree. Permuting one of the two lists, one attains the configuration model. Pathological permutations give rise to multiple edges and self-interactions. Individual pathologies can be corrected at little additional computational expense (see FIXPATH in [22] for details). In this case, we preserve $N(k_s, k_{s0})$, the joint distribution for in and out degree and self-interactions. In [23,12] a similar ensemble is used where multiple edges are disallowed; however, our approach differs in the following respects: (i) our algorithm is a more efficient single shuffle rather than multiple swaps; (ii) iteratively rewiring requires the introduction of another cutoff parameter, defining how many rewiring steps are needed; shuffling obviates the need for this additional parameter; (iii) iterative swapping generates Markov chain realizations whereas shuffling generates independent, identically distributed samples; and (iv) we preserve self-interactions.

STATISTICAL SIGNIFICANCE

In the past, statistical analyses of subgraphs have relied on z scores or empirical sample estimates of probabilities. In Fig. 4 we show that many features (both for subgraphs and for scalars) are not Gaussian, so $z$ scores are inappropriate measures of deviation from the background ensemble. Empirical sample estimates are also problematic, for example, if the distribution is undersampled. Instead we apply standard
tools from machine learning, namely, kernel density estimation and cross validation to learn the distribution from the sample data. Cross validation is a model evaluation method where model learning relies on part of the data, while model testing relies on the rest of the data, the holdout set. \( K \)-fold cross validation repeats the holdout method \( k \) times. To quantify a network’s deviation from the background ensemble, we learn the distribution for each scalar and measure deviation as the likelihood that an observation was drawn from the background distribution. Given a graph and our model, we collect \( m \) realizations and estimate the probability density \( p(w) \) for a scalar \( j \) to have a value \( w \) using Gaussian kernel density estimation [24]:

\[
p_{\lambda_*(w)} = \frac{1}{m} \sum_{i=1}^{m} \exp\left(-\frac{|w_i - w|}{2\lambda_*(a)}\right) / \left(2\pi \lambda_*(a)^2\right)^{3/2}
\]

where \( w_i \) are the scalar values of the randomizations, and \( \lambda_*(a) \) is a real-valued smoothing parameter. By partitioning the data into five “folds” and holding out one fold at a time to calculate the average probability of a holdout set according to the other \( 4/5 \) of the data (“fivefold cross validation” [24]), we define the function

\[
Q(\lambda) = \frac{1}{5} \sum_{i=1}^{5} \prod_{j=1}^{4m/5} p_{\lambda}(w_{f_{i}(j)})
\]

where \( \{f_{i}(j)\} \) is the set of indices associated with fold \( i \) (\( i = 1, \ldots, 5 \)). We then determine \( \lambda_*(a) \) as \( \lambda_*(a) = \arg\max \lambda Q(\lambda) \). For a real-world graph of interest, ranking of likelihoods reveals the most significant measures of the network—the scalars which are least like the background ensemble. Figures 5 and 6 show the results of density estimation on the two most significant scalars.

Consider the set of scalars for digraphs, \( \Sigma(B_1B_2\cdots B_n) \times (B_i \in \{A,A^T\}, n \in \mathbb{N}) \). These scalars perform a census which includes all possible walks and therefore all possible subgraphs. The operators \( D \) and \( U \) constrain the set of all subgraphs so that a given scalar only counts a small subset simultaneously. In this way scalars inherit statistical significance from subgraphs. While some scalars count an individual subgraph, other scalars count combinations of subgraphs. The mapping of scalars to subgraphs is thus many to many.

While the analysis proceeds independently of subgraphs, it is possible, given a graph, to find any scalar’s most representative set of subgraphs. We call this process localization. We define a skeleton to be the smallest subgraph with nonzero value of the scalar. As an approximate, greedy algorithm to find a most representative set of skeletons, given a graph \( A \) with nonzero value of a scalar \( W \), we (1) build a subgraph \( s \) by adding nodes from \( A \) until \( W \) evaluated on \( s \) gives a nonzero value (soft localization) or the original value (hard localization); (2) distill this subgraph by removing nodes from \( s \) until we arrive at a subgraph \( s' \) such that removing any additional nodes would cause the value of \( W \) to vanish; and (3) repeat on \( A-s' \) until all nodes have been exhausted.

The resulting algorithm yields a set of representative subgraphs for a given scalar. Each \( s' \) subgraph in the set is labeled according to its isomorphism class. The most representative subgraph is simply the subgraph class that has the highest relative fraction of the total recovered set of subgraphs. Multiple iterations of the localization algorithm should be run since the algorithm depends on the order of the nodes; however, in practice, we do not see differences in results using different orderings.

As an example, hard and soft localization of the scalar \( \Sigma(DAUA^T A) \) from the \( E. \ coli \) genetic network reveals the \( T_{030} \) triad [Fig. 1(a)] familiar from [2]. Arbitrarily large sub-
structures may emerge from a given scalar, highlighting another methodological advantage to the algorithm: the search for significant scalars does not impose any constraints regarding the size of resulting subgraphs. An upper bound on the computational complexity is \((\ell - 1)\Sigma s^3\), where \(s\) is the size of the resulting substructure and \(\ell\) is the length of the scalar. In general, however, the efficiency of the localization algorithm is of less concern, as we localize only on a small set of statistically significant scalars.

**E. COLI DATA SET**

We implemented our algorithm on the *E. coli* genetic network. The database includes 577 interactions between 423 nodes, combining an existing database [25] with additional nodes and edges included from a literature search as described by Shen-Orr et al. [2]. We exclude self-interactions for a total of 519 edges. Density estimations (see Figs. 5–7) demonstrate how *E. coli* deviates from our background ensemble. Three of the top-ranking statistically significant scalars, \(\Sigma(DAUA^T A)\), \(\Sigma(DA^T AA^T A)\), and \(\Sigma(DA^T AA^T A)\), localize to several structures consistent with Shen-Orr et al.’s earlier findings with this data set (see Fig. 1). However, we highlight that identification of these three significant structures was done using one algorithm without the need to pose thresholds or parameters or to provide tailored algorithms. No property of the network was assumed to be of interest beforehand. Of interest, \(\Sigma(AA^T DA^T AA)\) was the highest-scoring scalar. Upon soft localization, we recovered the two four-node subgraphs (Fig. 8), which we call “FFB” (feed-forward box) and “+FFL” (feed-forward loop with an input). The four-node structures are more significant than the related three-node \(T_{030}\) topology. The methodology thus assigns significance to a scalar without biasing the size of the resulting subgraphs.

Closier inspection of the top-scoring scalars reveals some unexpected architectural features. Hard localization of the significant scalar \(\Sigma(AA^T DA^T AA)\) yields a 14-node topology (Fig. 9). We observe that the \(T_{030}\) topology, defined by the genes = *hns*, *flhDC*, and *fliA*, is a motif shared by five overlapping FFB’s. Inspecting the word *DA^T AA* on the *E. coli* data, we find that there are 42 distinct \(T_{030}\) paths, but only 10 distinct \(T_{030}\) grandparents. That is, the operation \(\Sigma\) evaluates to 42, while the operation \(N\) evaluates to 10. In fact, the gene *crp* appears in 16 “distinct” \(T_{030}\). In this way the network evidences motif hubs—individual nodes that appear in numerous, overlapping identical motifs, a result first noted in [26] using a more primitive significance test and more recently reported in [27]. Importantly, this result is obtained with a single algorithm without posing any prior assumptions about the network.

Scalars that are significantly smaller relative to the background ensemble also reveal interesting topological features.
of the graph. For example, we find the scalar $N(UA^TA)$ is statistically underrepresented in the *E. coli* network (see Fig. 7). Localizations reveal structures with nodes that have two or more incoming edges. This “fan-in” structure, the opposite of the “SIM” topology, thus appears less often in the network, a finding with important ramifications. For example, recently researchers attempting to infer genetic regulatory interactions have imposed priors which restrict the number of edges converging on a node, but leave unrestricted the number of edges leaving a node [28]. This prior on a general “fan-out” topology is thus supported by our findings.

**S. CEREVISIAE DATA SET**

The yeast dataset is based on the Yeast Proteome Database (YPD) [29] and this particular part of the network consists of 688 nodes with 1079 edges [30]. Analysis of this network shows the most significant word $\Sigma(DAAA^TDAA)$ contains a mutual dyad (a term which we borrow from the sociological network literature, referring to a pair of vertices mutually linked, such that $a \iff b$) as the rightmost DAA indicates. Upon hard localization we find that only four nodes in the network contribute to the word; these four nodes make up a dense cluster which includes a mutual dyad and a three-cycle (see Fig. 10). Another significant feature, $\Sigma(DAAAU^TA)$, hard-localizes to a 22-node substructure (Fig. 11) with a fascinating topology which includes two parent genes that have a large and almost identical set of children. In the soft localization of this feature, a minimal subgraph emerges with a compound topology: the “parent” layer of a FFL is itself a FFL (Fig. 12). Obviously this five-node subgraph would not be identified with subgraph census methods which only count up to three- and four-node subgraphs.

**INTERPRETATION**

We have presented a generalizable method for enumerating measures of a network and have demonstrated an application of this method for finding statistically significant features of the *E. coli* and *S. cerevisiae* genetic regulatory networks. The method has the advantages of computational efficiency as compared to subgraph census for naturally occurring networks, particularly clustered or scale-free networks, and flexibility in that it can be easily applied to weighted and signed graphs. For example, many biological networks are published with a “$p$ value” associated with each edge [31,32], i.e., a probability that a certain edge exists (implicit in such publications is the assumption that the existence of each edge is independent of all other edges). In this case, $\Sigma$ refers to the expected value of that scalar, over all realizations of the graph. Alternatively, neuronal networks have weighted edges describing the number of synapses and

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**FIG. 10.** In *S. cerevisiae*, both hard and soft localization of the significant scalar $\Sigma(DAAA^TDAA)$ yields this densely clustered four-node topology which includes a mutual dyad and a three-cycle. Unlike the *E. coli* network, this network contains feedback interactions.

**FIG. 11.** In *S. cerevisiae*, hard localization of the significant scalar $\Sigma(DAAAU^TA)$ yields this interesting 22-node topology. Note again the fan-out structure whereby two genes regulate a very similar set of genes.

**FIG. 12.** In *S. cerevisiae*, soft localization of the significant scalar $\Sigma(DAAAU^TA)$ yields this five-node topology as the most representative subgraph. Interestingly, the structure can be seen as a hierarchical feed-forward loop. For example, if we replace the three-node feed-forward loop with an effective node, that node itself becomes the parent layer of another feed-forward loop.
thus the strength of the interaction. In this case, $\Sigma$ calculates the functionality of a particular word. While the $\mathcal{N}$ operation does not differentiate between weighted and unweighted edges, we could easily imagine other useful quantities of interest that we can also use in our space that would be functions of the weighted edges, such as the standard deviation. Some of our results on *E. coli* confirm earlier findings from previous methods, but unlike those methods, this approach is a single, systematic algorithm which does not require any previous assumptions about the network being analyzed. Moreover, results regarding the structure of the *E. coli* network are presented, including the presence of “motif hubs,” “feed-forward boxes,” and a general “fanout” topology.

It is worth highlighting that under a different randomization scheme with a different set of conditionals, the results may differ substantially. For example, in the case of the yeast data set, if the number of three-cycles or mutual dyads was also preserved, we expect the ranking of scalars to be different. We note, then, that one must take great care in selecting the background ensemble to avoid the possibility that one’s choice of randomization predetermines which scalars are the most significant. While the configuration model and its variants have been used as the appropriate ensemble distribution for networks in the past, many other random network models exist which may be more appropriate. Potentially, the network embedding space we present here will elucidate these issues further. For example, given multiple realizations of a random network model, one can use this space to investigate whether the resulting distributions are separable and which features make them distinguishable.

While motivated by work in which the subgraphs are the primitive degrees of freedom, scalars do not have a one-to-one mapping to subgraphs. However, every subgraph contributes to at least one scalar. Subgraph counting is computationally expensive, particularly for clustered, scale-free, and dense networks, but our method alleviates this issue because its exponential term is independent of these properties. The trade-off is that with localization, we can only find sets of subgraphs that a given scalar counts. A more systematic alphabet could further constrain the set of subgraphs for a scalar.

Closer investigation into the mapping between scalars and subgraphs is needed. The heuristic we develop, localization, appears to work well. The scalars are easily mapped to their most representative subgraphs, and some of these subgraphs confirmed earlier findings on the same data set. However, while in our studies the interpretation of the most representative subgraphs of the significant scalars was straightforward, some scalars may have more difficult interpretations. We note that our focus here was not on subgraphs *per se*, but rather on a data space, a set of measures on a graph from which one can perform various statistical studies. In general, if one is interested in a particular subgraph, then the best approach is to identify that subgraph in the network. If one does not have any preconceptions about which feature of the network is important to study, than the scalar space offers an alternative, systematic, efficient, and effective approach to census and/or listing properties deemed relevant. Indeed, the space may not only be related to subgraphs, but also to more global measures such as various orders of transitivity [3].

Finally, we note additional utilities of the enumeration of words. First, given an algorithm which purports to model a real-world network, one could find statistically significant scalars to identify in what ways the model fails to model the real-world data. Second, given a training set of many graphs of multiple classes, this data space could be used to build a classifier using machine-learning algorithms (e.g., Support Vector Machines (SVMs) [33,34]. Boosting [35]) which could then assign new graphs to one of the classes (see [36,31] for recent work in this direction), providing a modern machine-learning approach for diagnosing networks (e.g., robust versus fragile economies, graphs with different growth laws, etc.).

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