Practical Computation of Graph VC-Dimension

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Abstract –

For any set system $\mathcal{H}=(V,\mathcal{R}),\ \mathcal{R}\subseteq 2^V$, a subset $S\subseteq V$ is called *shattered* if every $S'\subseteq S$ results from the intersection of S with some set in \mathcal{R} . The VC-dimension of \mathcal{H} is the size of a largest shattered set in V. In this paper, we focus on the problem of computing the VC-dimension of graphs. In particular, given a graph G=(V,E), the VC-dimension of G is defined as the VC-dimension of G of G contains each subset of G that can be obtained as the closed neighborhood of some vertex G in G. Our main contribution is an algorithm for computing the VC-dimension of any graph, whose effectiveness is shown through experiments on various types of practical graphs, including graphs with millions of vertices. A key aspect of its efficiency resides in the fact that practical graphs have small VC-dimension, up to 8 in our experiments. As a side-product, we present several new bounds relating the graph VC-dimension to other classical graph theoretical notions. We also establish the G in our experiments of the graph VC-dimension problem by extending a previous result for arbitrary set systems.

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1 Introduction

Since the seminal work of Vapnik and Chervonenkis [35], VC-dimension is one of the basic quantities describing the complexity of a set system. As such, VC-dimension is the foundation of many results in mathematics and theoretical computer science: it plays a central role in uniform sampling guarantees and is often required as an input parameter of algorithms. For instance, one of the fundamental results in computational learning theory states that a set system is PAC-learnable if and only if it has bounded VC-dimension [2]. The Sample Compression Conjecture, which has been called in [6] one of the oldest open problems in theoretical machine learning, is also related to VC-dimension [17]. VC-dimension has become a key concept in other fields as well. In computational geometry, bounds on the VC-dimension of various geometric set systems (e.g., ones induced by half-spaces, balls or axis-parallel boxes) are essential parameters in methods developed for approximating and processing geometric data [12, 37]. Indeed, set systems with bounded VC-dimension admit structures such as small-size ε -nets [20] and ε -approximations [35, 24, 10], matchings and spanning paths of low crossing numbers [36, 7], colorings with low discrepancy [28] to name a few. We refer to the survey [27] for more details.

Practical applications of this parameter (e.g. in the design of PAC-learning algorithms), require bounds on the VC-dimension of the considered set systems. However, it was observed that the general bounds found in the literature are of limited use in practice [21]. Thus, the problem of computing the VC-dimension of set systems has attracted some attention. It is proved to be logNP-hard [31], and W[1]-hard for the natural parameterization by the VC-dimension [13]. Furthermore, under plausible complexity hypotheses, the VC-dimension is hard to be approximated within a sub-logarithmic factor in polynomial time [26].

There is a fast growing body of literature demonstrating the strong potential of using VC-dimension as a graph parameter. In recent studies, researchers have made significant progress in improving results in extremal and algorithmic graph theory by limiting the problems to objects with bounded VC-dimension [25, 18, 14, 16, 4, 3], including an EPTAS for the MAXIMUM CLIQUE problem [3] and subquadratic-time algorithms for diameter computation [14, 16]. The VC-dimension of a graph was also linked to the complexity of approximating a minimum-cardinality identifying code on hereditary graph classes [4].

The most commonly used notion of VC-dimension for graphs is defined as the VCdimension of its neighbourhood set system (see Section 2 for a formal definition). More specifically, in this paper we consider the *closed* neighbourhoods of vertices (i.e., each vertex is included in the set of its adjacent vertices). However, we could instead consider their open neighbourhoods. Both notions result in different but comparable values for the VCdimension. The algorithmic applications listed in [4, 14, 16] are proved using the set systems of closed neighbourhoods in a graph, while those listed in [3, 18] are proved using the set systems of open neighbourhoods. Note that the VC-dimension of other graph-related set systems has been considered in [8, 9, 16, 22], with different combinatorial and algorithmic implications. These alternative VC-dimension parameters are not considered in our paper. However, it is noteworthy that several of these parameters can be lower bounded by the graph VC-dimension. More generally, it was observed in [5] that every set system \mathcal{H} can be represented as a split graph $G_{\mathcal{H}}$, in such a way that the VC-dimension of \mathcal{H} is equal to the VC-dimension of the closed neighbourhoods of vertices in the stable set of $G_{\mathcal{H}}$. Therefore, graph VC-dimension is as general as the VC-dimension of arbitrary set systems, if we allow ourselves to only consider the closed neighbourhoods of a restricted subset of vertices.

Just like for general set systems, the problem of computing the VC-dimension of a

graph is known to be logNP-hard [22]. However, we are not aware of any previous study on the parameterized complexity of the problem. Similarly, very few is known about the VC-dimension of complex networks. The closest such related work would be [11], where the stronger property of bounded expansion is considered. The VC-dimension of random graphs has been studied in [1], where for any fixed value d, a density threshold for the property of having VC-dimension at most d is derived.

Our Contributions.

While developing improved methods for graphs with bounded VC-dimension is a fruitful direction, it is just as important to provide efficient algorithms and conditions to help computing or approximating the VC-dimension of the input graph. We address this problem both from a theoretical and practical point of view.

The main contribution of this paper is a practical algorithm for computing the VCdimension of any graph (Algorithm 1). Note that a naive algorithm for this problem would consider all vertex subsets of size at most the VC-dimension of the input. By contrast, our algorithm repeatedly updates a lower bound on the VC-dimension, so that most unexplored vertex subsets below this bound can be discarded. Furthermore, while exploring for larger shattered subsets, we use our degree-based upper bounds on the VC-dimension in order to discard at once all vertices of too small degree (at most exponential in the current lower bound). Similarly, we show that while growing a shattered subset by iteratively adding new vertices, some branches can be ignored using a simple, but surprisingly powerful, upper bound on the size of a largest shattered superset (Lemma 2). By doing so, we considerably reduce the search space, as evidenced by our experiments on some real-life networks. We implemented a few more tricks, based on a combination of bit masks and partition refinement techniques, in order to speed up some important routine tasks in the algorithm, such as: the test of whether a given subset is shattered, that of whether a search branch can be pruned, and reduction schemes for the graph to be considered. Overall, we were able to compute the VC-dimension of graphs with millions of nodes in less than 40 minutes, providing the first practical algorithm for computing graph VC-dimension. We demonstrate the efficiency of our algorithm on various practical graphs. To the best of our knowledge, this is the first analysis of the VC-dimension in real networks. Interestingly, we observe that for all graphs considered in our experiments, the VC-dimension ranges between 3 and 8.

Our next contribution is proving that computing the VC-dimension of a graph is a W[1]-hard problem for the natural parameterization by the VC-dimension (Theorem 1). For that, we revisit a previous W[1]-hardness proof for arbitrary set systems [13], which we combine with some insights on shattered subsets in graphs from [15].

Finally, we note that we obtain a series of new bounds on the VC-dimension with respect to classical graph parameters, such as maximum degree, degeneracy and matching number. Some of these parameters are included in the setup of [4] who show that the VC-dimension of a graph can be functionally upper bounded by any hereditary graph parameter that stays unbounded on the following graph classes: split graphs, bipartite graphs and co-bipartite graphs. However, the bounds that can be derived from [4] are rather rough, due to the use of Ramsey's theory. By contrast, we give linear and sharp bounds for all the considered parameters.

Organization. After defining the main notions and notation of this paper in Section 2, we prove that computing the VC-dimension of graphs is W[1]-hard in Section 3. Then in Section 4, we summarise our bounds on the graph VC-dimension. In Section 5, we give a

new exact algorithm for computing the VC-dimension of graphs and discuss several possible optimizations. In Section 6, we report our experimental results and discuss the advantages of the different optimizations methods.

2 Definitions and notation

Throughout this note, we use lowercase letters u, v, x, y, \ldots for vertices, uppercase letters X, Y, Z, \ldots for sets of vertices, calligraphic letters as \mathcal{R} for collections of sets, and we let log denote the base 2 logarithm. Given an undirected graph G=(V,E) with |V|=n vertices and |E|=m edges, let $N_G[v]$ denote the closed neighborhood of v defined as $N_G[v]=\{u\in V\mid uv\in E \text{ or } u=v\}$. We define the degree $\deg(v)=|N_G[v]|$ of a vertex as its closed neighborhood cardinality. We use this unusual convention of counting a vertex in its own degree for the sake of simplicity when considering closed neighborhood sizes. We also define the ball $B_G[v,r]$ centered at a vertex v and with radius r as the set of nodes at distance at most r from v. In particular, we have $B_G[v,1]=N_G[v]$. We omit the G subscript when G is clear from the context.

A set system $\mathcal{H} = (V, \mathcal{R})$ (or hypergraph) is defined by a ground set V and a collection \mathcal{R} of subsets of V called ranges. Recall that a set $X \subseteq V$ is said to be shattered by \mathcal{R} (or simply shattered if \mathcal{R} is clear from the context) if for every $Y \subseteq X$ there exists a range $R \in \mathcal{R}$ such that $Y = R \cap X$. For any $R \in \mathcal{R}$, the intersection $Y = R \cap X$ is called the trace of R on X. The VC-dimension of a graph G, denoted by VCdim(G), is defined as the VC-dimension of its closed neighborhood set system $(V, \{N_G[v] \mid v \in V\})$. A subset $X \subseteq V$ is thus shattered if for every $Y \subseteq X$ there exists a vertex v_Y such that its trace $N_G[v_Y] \cap X$ on X equals Y. When considering such a set system, we say that a vertex v has trace Y on a set X when its closed neighborhood has trace $Y = N_G[v] \cap X$.

3 W[1]-hardness

As we have mentioned in the introduction, computing the VC-dimension of graphs is known to be logNP-hard [22]. We show that it is also W[1]-hard for the natural parameterization by the VC-dimension by showing the following statement in Appendix A.

▶ Theorem 1. For any graph G and parameter $k \leq |V(G)|$, there exists a graph H_G such that G contains a k-clique if and only if the VC-dimension of H_G is at least k. Furthermore, we can construct H_G from G in $\mathcal{O}(k2^kn^2)$ time.

4 Simple bounds

The following are upper bounds on the size of shattered subsets in graphs, with respect to various graph parameters. We emphasize on Lemma 2, which is a cornerstone of our practical algorithm for computing the VC-dimension of graphs (presented in the next Section 5).

▶ **Lemma 2.** Consider a shattered set X and $Y \subseteq X$. Let Y' be the set of vertices with trace Y on X. Any shattered set Z containing X satisfies $2^{|Z|-|X|} \le |Y'|$.

Proof. For any subset $X' \subseteq Z \setminus X$, there must exist a vertex $v_{X'}$ with trace $N[v_{X'}] \cap Z = Y \cup X'$. As $v_{X'}$ has trace Y on X, it is included in Y'. All vertices $v_{X'}$ are pairwise distinct since they have pairwise distinct traces on Z. Hence, Y' has size at least $2^{|Z \setminus X|}$.

Setting $X = Y = \{x\}$ for any vertex x, we have Y' = N[x] and obtain the following bound.

Corollary 3. Any shattered set Z containing a vertex x has size at most $|\log \deg(x)| + 1$.

In Appendix B, we additionally relate the VC-dimension of a graph with its degeneracy k and its matching number ν (recall that a graph G is called k-degenerate if every subgraph of G contains a vertex with at most k neighbours, and that ν is the size of a maximum matching in G). We summarize the obtained upper-bounds in the next lemma.

▶ **Lemma 4.** Let G be a non-empty graph on n vertices with maximum degree Δ , matching number ν , and degeneracy k, then

$$VCdim(G) \le min \{ |\log n|, |\log \Delta| + 1, k + 1, \nu + 1 \}.$$

The following lemma allows to restrict the search of a shattered set containing a given node x to its ball B[x,2] of radius 2.

▶ **Lemma 5.** For any shattered set X and $x \in X$, we have $X \subseteq B[x,2]$.

Proof. Since X is shattered, for any vertex $y \in X \setminus \{x\}$, there exists a vertex $v \in V$ such that $N[v] \cap X = \{x, y\}$. That is, v is a common neighbor of x and y and so $\text{dist}_G(x, y) \leq 2$.

5 Algorithm

The exact algorithm is given in Algorithm 1. Apart from the graph, it receives a lower bound lb on its VC-dimension. Alternatively, one can start the algorithm with input value lb = 0. In Section 5.2, we describe a way to obtain a better starting value for lb.

5.1 Outline of the method

Given a graph G and a lower bound lb of its VC-dimension, our algorithm consists in exploring all possible shattered sets of size at least lb+1 according to Algorithm 1. If one is found, lb is updated and the search is continued on larger sets. When no shattered set of size lb+1 is found, we conclude that lb is equal to the VC-dimension. The most technical part—checking for shattered supersets—is contained in the function Exploreshattered (see Algorithm 2). We encode a subset $Y \subset X = \{x_1, \ldots, x_k\}$ by the integer with binary representation $y = y_k \cdots y_1$ where $y_i = 1$ if $x_i \in Y$ and $y_i = 0$ otherwise. The trace $N[v] \cap X$ of each vertex v is therefore stored in a mask M[v] which is updated as we visit subsets X of H: the ith bit of M[v] indicates whether the ith vertex of X is in N[v].

Algorithm 1 VCDIMCOMPUTATION (G, n, lb)

```
Input: A graph G=(V,E) with n=|V| vertices, lower bound lb on VCdim(G)
Output: The VC-dimension of G

1 Let H be an array containing all vertices of degree at least 2^{lb}
2 Sort H (optional). // We consider 3 different ways of sorting, see Section 5.3
3 Initialize a mask M[v] \leftarrow 0 for all v \in H. // Trace of N[v] on X=\emptyset
4 Initialize T=[T[0]] \leftarrow [n]
// T[y] counts the number of vertices v \in V with trace M[v]=y on X=\emptyset
5 For i=1 to |H| do
6 |Lb \leftarrow \text{EXPLORESHATTERED}(H,i,\emptyset,T,lb)
7 Return lb
```

To make the algorithm more efficient, we incorporated the following key ideas:

Algorithm 2 ExploreShattered (H, i, X, T, lb)

```
1 Set x \leftarrow H[i], s \leftarrow |X \cup \{x\}|, and m \leftarrow 2^{s-1}.
                                                                     \ensuremath{/\!/}\ m is the bit mask for x
 2 T' \leftarrow \text{TRACECOUNTADD}(T, x, m) // T'[y] = \# \text{ vertices with trace } y \text{ on } X \cup \{x\}
 s prune \leftarrow \texttt{False}
 4 For y = 0 to 2^s - 1 do
    If T'[y] < 2^{lb+1-s} then prune \leftarrow \text{True}.
                                                                                        // by Lemma 2
 6 If not prune then
       For v \in N_G[x] do
                                      // Update M(v) to be the trace of N[v] on X \cup \{x\}
         M[v] \leftarrow M[v] + m
       If s > lb then lb \leftarrow s
                                                                          // X \cup \{x\} is shattered
       For j = i + 1 to |H| do
10
        lb \leftarrow \text{ExploreShattered}(H, j, X \cup \{x\}, T', lb)
11
       For v \in N_G[x] do M[v] \leftarrow M[v] - m // Restore the trace of N[v] on X
12
13 Return lb
14 Function TRACECOUNTADD(T, x, m)
       Let T' be a copy of T resized to 2^s and padded with zeros.
15
       For v \in N_G[x] do
16
                                                                      // The trace of N[v] on X
17
            Consider y \leftarrow M[v]
            T'[y] \leftarrow T'[y] - 1
18
           T'[y+m] \leftarrow T'[y] + 1
19
       Return T'
20
```

- According to Corollary 3, we can restrict the search to the set H of vertices with degree 2^{lb} at least.
- We fix an ordering \prec of H and scan subsets of H in a depth first search manner. More precisely, ExploreShattered performs a DFS of the inclusion graph of subsets of H by following arcs $X \to Z$ for $X, Z \subseteq H$ such that $Z = X \cup \{z\}$ and $x \prec z$ for all $x \in X$. Starting from the empty set, any set $X = \{x_1, \ldots, x_k\}$ with $x_1 \prec \cdots \prec x_k$ is thus reachable through $\emptyset \to \{x_1\} \to \{x_1, x_2\} \to \cdots \to X$.
- For each visited set X, we compute a table T counting for each $Y \subseteq X$ the number of vertices v with trace $N[v] \cap X = Y$. If $T[Y] < 2^{lb+1-|X|}$ for some Y, then Lemma 2 implies that there exists no shattered set $Z \supseteq X$ of size lb+1 or more, so we do not explore the supersets of X. Note that this test is not satisfied when X is not shattered as we then have T[Y] = 0 for some $Y \subseteq X$. The argument X of Exploreshattered is thus always a shattered set.
- When considering $Z = X \cup \{x\}$, the table T' for Z can be obtained from T in time $\mathcal{O}(|T| + \Delta) = \mathcal{O}(2^d + \Delta)$ where Δ is the maximum degree in G and d is its VC-dimension.

The overall worst case complexity of the algorithm is thus $\mathcal{O}(n^d(2^d + \Delta))$ as we visit only shattered sets. Moreover, Corollary 3 implies $2^d = \mathcal{O}(\Delta)$ and the complexity is thus $\mathcal{O}(n^d\Delta)$. This pessimistic bound assumes that a constant fraction of sets of size d are shattered. Empirically we observed much better running times, as reported in Section 6 with the evaluation on practical graphs. Section 5.3 describes several optimizations, including how to sort H.

5.2 Lower-bound computation

We compute a lower bound lb by a method similar to VCDIMCOMPUTATION with lb = 0 (thus H = V). We make the search faster by only performing a partial scan of subsets of H. In order to find large shattered sets, we sort H by non-increasing degrees. We restrict the search in two ways: we fix a maximum number maxvisits = 64 of times a vertex x can be added to the current shattered set; we also restrict the for loop of Line 10 of ExploreShattered to the first maxvisits/2 elements. As each vertex is visited a constant number of times (at most maxvisits), this modified version takes linear time.

5.3 Optimizations

Vertex ordering.

We have considered the following options for sorting the set H of high degree vertices:

- \blacksquare non-increasing degrees (D^-) ,
- \blacksquare non-decreasing degrees (D^+) ,
- \blacksquare k-core ordering (K): an ordering of $G_{|H}$, the subgraph restricted to H, obtained by repeatedly removing a vertex with lowest degree, vertices removed first are ordered first,
- \blacksquare random ordering (R).

The intuition for choosing non-increasing degrees follows that of the lower bound computation: higher degree vertices tend to participate to larger shattered set and exploring them first can improve the lower bound earlier, allowing to restrict the rest of the search more severely thanks to Lemma 2. Conversely, if we start with a good enough lower bound, lower degree vertices tend to participate to smaller shattered sets and the exploration from these vertices tends to be faster. Exploring them first then speeds up the exploration from high degree vertices as we do not need to consider adding already scanned vertices anymore. Using a k-core ordering follows a similar intuition with the refinement of taking into account the degree after removing previous vertices rather than the degree in the full graph. Using a random ordering seems a basic choice for comparison.

Ball restriction.

Lemma 5 implies that when starting an exploration from x = H[i] (Line 6 of VCDIMCOMPUTATION) we can restrict the search to consider only vertices in $B[x,2] \cap \{H[i], H[i+1], \ldots\}$.

Graph reduction.

As we focus on shattered sets included in H, we can restrict the graph while preserving all possible traces on H. For that purpose, for each possible trace $Y\subseteq H$ which can be obtained as $H\cap N[v]$ for $v\in V$, we keep at most one vertex $v\in V\setminus H$ with trace $N[v]\cap H=Y$. Such a selection can be obtained in linear time using partition refinement [30, 19] as follows. Starting from the partition $\mathcal{P}=\{V\}$ we iteratively refine it by sets N[x] for $x\in H$: each refinement step consists in splitting each part $P\in \mathcal{P}$ into $P\cap N[X]$ and $P\setminus N[X]$ (if one of the two sets is empty, P remains unchanged). Each refinement step clearly maintains the invariant that all vertices in a part have the same trace on the set of vertices of H processed so far. At the end of the process, all vertices in a part must have the same trace on H. We thus keep one vertex per final part not intersecting H and all vertices in H to obtain a set $V'\supseteq H$ of vertices providing the same traces on H as V and proceed on $G_{|V'|}$ instead of G.

Table 1 The graphs we use with their main parameters and the time (in seconds) required by our reference implementation *KBG* to compute their VC-dimension.

Graph	#nodes	#edges	max.deg.	$\operatorname{VC-dim}d$	time (s)
BIO-MV-Physical-3.5	21 866	68 123	1 117	5	0.11
BIO-SYS-Aff-Cap-MS-3.5	38 926	299855	2193	7	4.41
BIO-SYS-Aff-Cap-RNA-3.5	12 716	40541	3571	7	0.03
dip20170205	27 029	73762	289	5	0.14
oregon2-010331	10 900	31 180	2 343	6	0.20
CAIDA-as-20130601	44 611	151434	3962	7	11.50
DIMES-201204	25 367	75004	3781	7	0.16
as-skitter	1 696 415	11095298	35455	8	2116.90
p2p-Gnutella09	8 1 1 4	26 013	102	5	0.01
gnutella31	62 586	147892	95	4	0.29
notreDame	325 729	1090108	10721	6	2.94
y-BerkStan	685 231	6649470	84230	7	11.44
ca-HepPh	12 008	118 489	491	5	46.53
com-dblp	317 080	1049866	343	5	2.30
epinions1	75 888	405740	3044	7	58.69
facebook-combined	4 039	88234	1045	6	8.15
twitter-combined	81 306	1342296	3383	7	456.22
t.CAL	1890816	2315222	7	3	4.56
t.FLA	1 070 377	1343951	8	3	2.53
buddha	543 652	1631574	17	4	3.21
froz	754 197	2902525	8	3	4.70
z-alue7065	34 047	54841	4	3	0.03
grid300-10	90 601	162535	4	3	0.21
xgrid500-10	251 001	450766	4	3	0.61
powerlaw2.5	1 000 000	1916356	9447	6	3.79

6 Experiments

6.1 Dataset

We evaluate the performance of our algorithm on various types of practical graphs. We use graphs from the BioGRID interaction database (BIO-*) [29]; a protein interactions network (dip20170205) [32]; and graphs of the autonomous systems from the Internet (oregon2, CAIDA_as and DIMES) [23, 34, 33]. We also test computer networks (Gnutella, Skitter), web graphs (notreDame and BerkStan), social networks (Epinions, Facebook, Twitter), co-author graphs (ca-HepPh, dblp), road networks (t.CAL, t.FLA), a 3D triangular mesh (buddha), a graph from a computer game (FrozenSea), and grid-like graphs from VLSI applications (z-alue7065). The data is available from snap.stanford.edu, webgraph.di.unimi.it, www.dis.uniroma1.it/challenge9, graphics.stanford.edu, steinlib.zib.de, and movingai.com. Furthermore, we use synthetic inputs: grid300-10 and grid500-10 are square grids with respective sizes 301×301 , and 501×501 where 10% of the edges were randomly deleted; and powerlaw2.5 is a random graph generated according to the configuration model with a degree sequence following a power law with exponent 2.5.

All experiments were performed on a cluster of 20 nodes equipped with two Cacade Lake

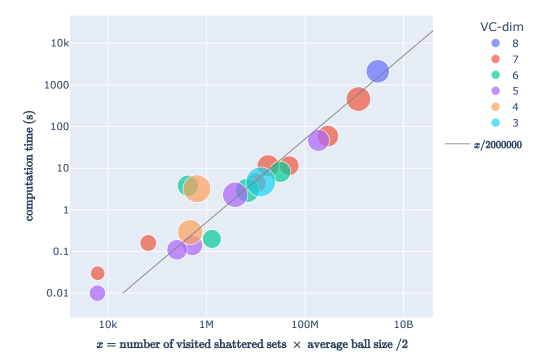


Figure 1 Computation time t in seconds versus the estimated number x of tentative shattered sets considered by KBG: each network in the dataset is represented by a disk with coordinates (x,t), whose color indicates the VC-dimension d of the network, while the size is proportional to the logarithm of the number of high degree nodes.

Intel Xeon 5218 16 cores processors at 2.4GHz and 192GB of memory each. Sixteen processes were run in parallel on four nodes of the cluster. Our computation times are thus pessimistic compared to running each process on a fully reserved node (for example, the computation of the VC-dimension of the twitter graph of our dataset takes 431s on a fully reserved node compared to 456s in our experiment even though we report user times). The code is available at gitlab.inria.fr/viennot/graph-vcdim.

6.2 Graphs and reference time

We first present our dataset graphs in Table 1 together with their VC-dimension and time required to compute it with our reference implementation KBG: which uses k-core ordering (K), ball restriction (B) and graph reduction (G). We have chosen KBG as our reference choice of optimizations as it provides the minimum sum of running times over all graphs of our dataset. We list also their size in terms of number of nodes (not counting isolated nodes) and number of edges (duplicate edges are removed).

We observe that the VC-dimension of all graphs in the dataset is rather small: at most 8, even for graphs with millions of nodes and over 10M edges (M stands for million). Moreover, its computation with our KBG implementation takes at most a few seconds for most of the graphs, and less than a few minutes for all but one: as-skitter for which it takes around 35 minutes. Memory usage (not reported here) grows with the input graph size up to roughly 600 megabytes for as-skitter. Not surprisingly, this most difficult graph is both the largest in terms of number of edges and the most complex in terms of VC-dimension. We analyze the dependency of the computation time with respect to some graph parameters in Section 6.3.

Graph	d	all	$deg \ge 2^d$	Lem.2	Lem.2G	$\# \deg \ge 2^d$	KBG	$\# \deg \ge 2^{lb}$	bsize
BIO-MV-Physical-3.5	5	6 970 780	1618308	4 471	4 242	582	4 242	582	118
BIO-SYS-Aff-Cap-MS-3.5	7	_	419882850	133927	133546	455	133546	455	152
BIO-SYS-Aff-Cap-RNA-3.5	7	1552654200	9438	1388	1246	19	1292	152	9
dip20170205	5	9050178	2823068	14503	14227	586	14227	586	72
oregon2-010331	6	18 763 142	1774582	2 932	2610	105	21 750	258	118
CAIDA-as-20130601	7	_	134557021	179232	174320	168	441741	421	217
DIMES-201204	7	345796843	1982414	4206	3870	67	3870	67	33
as-skitter	8	_	_	6671019	6658159	4182	6658159	4182	907
p2p-Gnutella09	5	1367927	39 096	433	420	58	420	58	29
gnutella31	4	1781293	131652	3958	2135	1938	19417	16089	47
notreDame	6	294870046	46413353	5022	2883	2063	26733	18807	507
y-BerkStan	7	_	33179823	108641	90394	1540	90394	1540	395
ca-HepPh	5	150 884 864	137719903	1046942	1037751	1 529	1037751	1 529	363
com-dblp	5	17874027	2489073	24748	14329	4280	65024	20566	117
epinions1	7	_	_	1026033	1025863	1162	1025863	1162	570
facebook-combined	6	722273307	568585300	389816	389471	882	389471	882	162
twitter-combined	7	_	_	1709219	1707213	3283	2562770	10483	950
t.CAL	3	8 107 636	12	12	0	0	0	0	0
t.FLA	3	4805712	6	6	0	0	0	0	0
buddha	4	7958071	21	17	0	0	270936	119239	4
froz	3	17402814	16370600	2033860	2026837	691850	2026837	691850	12
z-alue7065	3	250178	0	0	0	0	0	0	0
grid300-10	3	800 788	0	0	0	0	0	0	0
xgrid500-10	3	2222193	0	0	0	0	0	0	0
powerlaw2 5	6	236 286 923	31 023 430	4 653	2.674	886	2.674	886	314

Table 2 VC-dimension d, counts of shattered sets and high degree nodes, average ball size.

To appreciate these running times, recall that our algorithm for computing the VCdimension consists in first computing a lower-bound lb, then identifying the set H of highdegree nodes, that is those with degree at least 2^{lb} , and then ordering this set for exploring shattered sets included in H. Figure 1 shows that running times are basically proportional to the number x of shattered sets considered by KBG, which is estimated as follows. For each visited shattered set $X = \{v_1, \ldots, v_k\}$ with $v_1 \prec \cdots \prec v_k$, our algorithm tries to add high degree nodes of $B[v_1, 2]$ coming after v_k in the ordering \prec used for H. We can thus estimate x as the product of the number s of visited shattered sets multiplied by half of the average ball size $bsize = \frac{1}{|H|} \sum_{v \in H} ||B[v, 2] \cap H||$. These numbers are reported in Table 2. We see in Figure 1 that most networks are close to the black line that corresponds to a rate of 2 millions tentative shattered sets per second. For low values of x five networks appear significantly above the line: p2p-gnutella09, BIO-SYS-Aff-Cap-RNA-3.5, DIMES-201204, powerlaw 2.5 and buddha (from left to right). This is due to the overhead of reading the input file, computing the lower-bound, the k-core ordering, and the graph reduction which appears to be higher than the time for exploring shattered sets in these networks (we get comparable times when running all these phases and stopping before exploring).

6.3 Analysis: shattered sets and high degree nodes

The running time of our computation is mostly governed by the number of high degree nodes and the number of shattered sets explored, we thus present a detailed analysis of them. Table 2 lists several measures we could perform as follows. By removing the pruning according to Lemma 2, our algorithm explores all shattered sets. By setting an initial lower-bound of 0, we first tried to compute all shattered sets and report their number in column "all". Note that this computation was not doable within our timeout of 6 hours for six graphs. Unsurprisingly, bigger values are observed for larger VC-dimension. We then tried to compute all shattered sets included in the set H' of nodes with degree at least 2^d where

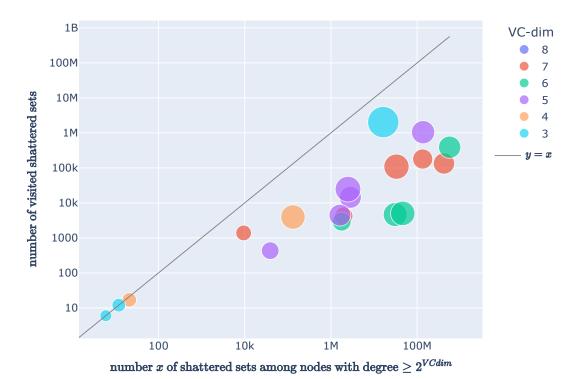


Figure 2 The number y of visited shattered sets versus the number x of shattered sets in H': each network in the dataset is represented by a disk with coordinates (x, y), whose color indicates the VC-dimension d of the network, while its size is proportional to the logarithm of |H'| (H' denotes the set of nodes with degree 2^d at least).

d is the VC-dimension of the graph. Again this computation was out of reach within our time limit for three graphs. We then use KB with d as lower-bound to obtain the number of visited shattered sets in H' according to Lemma 2 in column "Lem.2". We obtain similarly column "Lem.2G" using KBG. Column "KBG" is obtained using KBG with its heuristic lower-bound (instead of the exact value d). We also report the size of H' and H (columns "#deg $\geq 2^{d}$ " and "#deg $\geq 2^{lb}$ " respectively), and average ball size bsize in the last column.

We observe that restricting to H' can reduce the number of shattered sets by a huge factor (e.g. BIO-SYS-Aff-Cap-RNA-3.5). Note that H' can be empty in graphs with low maximum degree such as grids, explaining the zeros in the table. In general, Lemma 2 allows to further reduce the number y of shattered sets to explore as illustrated by Figure 2 (column "deg $\geq 2^d$ " versus "Lem.2" of Table 2). For networks with more than 10k shattered sets in H', we observe a reduction factor varying from roughly 10 for networks with VC-dimension 3-4, to roughly 100 for networks with VC-dimension 5, and from 1000 to 10k for networks with VC-dimension 6-7 except for BIO-SYS-Aff-Cap-RNA-3.5 for which the number x of shattered sets in H' was already quite low. We could not compute the value x for our only network of VC-dimension 8 within our 6 hours limit. Overall, this shows the efficiency of our approach by restricting to high degree nodes and pruning the search by Lemma 2.

6.4 Lower and upper bounds

As detailed in Appendix C.1, the lower bounds we obtain with our heuristic are mostly equal to the true VC-dimension or just one less. The upper bounds presented in Lemma 4, except

Table 3 Comparing different optimization options of our algorithm with our reference selection (*KBG*). A dash (–) indicates that the timeout of 6 hours was reached.

Graph	D^-BG	D^+BG	KBG	RBG	KG	KB	K
BIO-MV-Physical-3.5	0.22	0.09	0.11	0.22	0.17	0.13	0.19
BIO-SYS-Aff-Cap-MS-3.5	18.09	3.06	4.41	5.69	6.50	4.58	6.86
BIO-SYS-Aff-Cap-RNA-3.5	0.03	0.04	0.03	0.06	0.04	0.04	0.06
dip20170205	0.31	0.12	0.14	0.18	0.31	0.17	0.35
oregon2-010331	0.54	0.55	0.20	0.29	0.20	0.25	0.25
CAIDA-as-20130601	11.76	17.26	11.50	6.76	11.85	14.10	14.06
DIMES-201204	0.15	0.08	0.16	0.11	0.09	0.13	0.13
as-skitter	13 830.13	3673.65	2116.90	4022.75	8892.87	2358.56	10102.42
p2p-Gnutella09	0.02	0.01	0.01	0.03	0.02	0.02	0.01
gnutella31	0.17	0.30	0.29	0.32	15.20	0.30	16.77
notreDame	1.87	14.36	2.94	2.85	55.95	4.41	122.59
y-BerkStan	11.83	11.32	11.44	8.33	56.86	38.47	521.76
ca-HepPh	138.93	33.54	46.53	61.00	47.03	47.82	48.18
com-dblp	1.79	2.23	2.30	1.71	82.82	2.68	170.05
epinions1	511.94	31.08	58.69	144.99	58.69	63.67	63.07
facebook-combined	21.37	10.30	8.15	12.09	44.56	8.24	45.04
twitter-combined	1079.24	3132.27	456.22	397.94	1532.12	473.12	1549.07
t.CAL	4.59	4.41	4.56	4.45	4.56	4.85	4.87
t.FLA	2.42	2.35	2.53	2.47	2.46	2.62	2.58
buddha	2.99	2.92	3.21	3.01	2606.40	3.22	4525.09
froz	4.39	4.40	4.70	6.44	_	4.45	_
z-alue7065	0.03	0.03	0.03	0.04	0.03	0.04	0.03
grid300-10	0.22	0.23	0.21	0.21	0.21	0.22	0.21
xgrid500-10	0.61	0.59	0.61	0.63	0.62	0.62	0.65
powerlaw2.5	4.06	3.59	3.79	3.77	3.66	7.88	8.90

for the matching number, are analyzed in Appendix C.2 and are often much larger than the true VC-dimension except for grid-like graphs.

6.5 Optimizations

We now compare our reference implementation with other variants of our algorithm obtained by changing the ordering of the vertices (D^-, D^+, K, R) , using ball restriction (B) or not, and using graph restriction (G) or not (see Table 3).

Concerning the ordering of the nodes used for scanning shattered sets, we first note that non-decreasing degrees (D^+) is almost always faster than non-increasing degrees (D^-) . Notable exceptions are notreDame and twitter-combined where our initial lower bounds are VCdim(notreDame) -2 and VCdim(twitter-combined) -1 respectively. By observing our traces of execution, we explain it by the fact that the non-increasing order allows to find faster a better lower-bound, which then speeds up the rest of the computation. When the starting lower-bound was indeed exact, non-decreasing degrees is always faster or at least very close to non-increasing degrees. Our intuition is that the number of tentative shattered sets inspected is lower in that case. Indeed, for shattered sets constructed from the first nodes of the ordering, we have to consider all possible remaining nodes (that are in their ball of radius two) and try to construct a tentative shattered set by adding each of them. Putting nodes with lower degree first seem to result in a better balance with respect to the number of tentative shattered sets tested for being shattered. In that respect, the k-core ordering (K) seems to work slightly better since it considers the remaining degrees after removing the

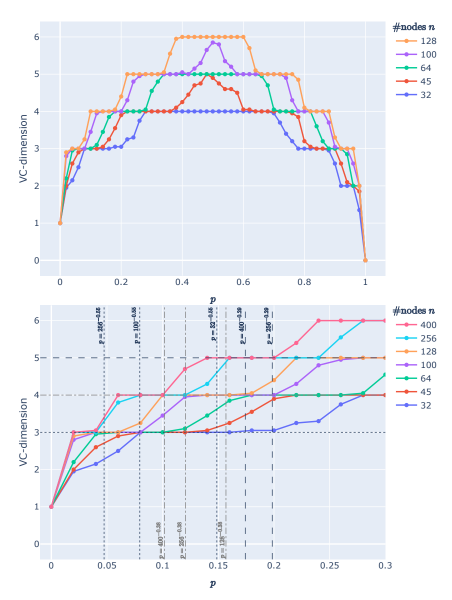


Figure 3 Top: the average VC-dimension of $G_{n,p}$ as a function of p for n = 32, 45, 64, 100, 128. Bottom: a zoom on values $p \in [0, 0.3]$ including additional curves for n = 256, 400.

first nodes rather than degree in the full graph. We note however that a random ordering (R) gives overall good results and RBG can even outperform KBG when our lower-bound is not exact, similarly as D^-BG can outperform D^+BG . This is in particular the case for twitter-combined. Overall there is no clearly better strategy for the ordering and both k-core ordering and random ordering seem legitimate choices.

Recall that the ball optimization (B) consists in restricting the nodes added to a shattered set X to those that are in the ball of radius two centered at the first node of X. It also allows to reduce the number of tentative shattered sets tested and is almost always beneficial. It appears to be mandatory on graphs with very large number of high degree nodes such as froz (the number of high degree nodes is analyzed in Section 6.3).

The graph reduction optimization (G) is not very costly in terms of computation time

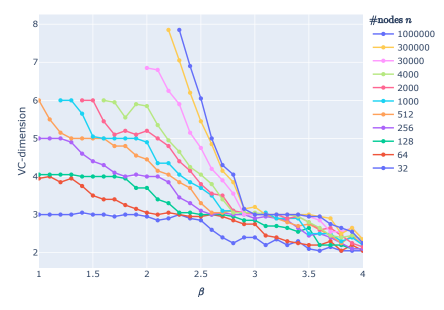


Figure 4 Average VC-dimension of a power-law random graph with respect to the exponent β of the law for various numbers n of nodes. (Curves for large n are truncated for low values of β .)

(KBG is always almost as good as KB) and gives a significant improvement on difficult graphs such as as-skitter and y-BerkStan.

6.6 VC-dimension of random graphs

We computed the average VC-dimension of various Erdős-Rényi random graphs $G_{n,p}$ (where each edge appears independently with probability p) with up to n=400 nodes and compared them to [1] which proved a threshold of $p=n^{-11/20}=n^{-0.55}$ above which the VC-dimension d of $G_{n,p}$ tends to be greater than 3, and similarly $p=n^{-21/55}\approx n^{-0.38}$ for d>4, and $p=n^{-7/24}\approx n^{-0.29}$ for d>5. On the one hand, our results confirm the $p=n^{-0.55}$ and $p=n^{-0.38}$ thresholds which appear to be rather sharp already for n=100 or n=256. On the other hand, observing the $p=n^{-0.29}$ threshold seems to require size n greater than 400 (see Figure 3).

Figure 4 shows the VC-dimension of various power-law random graphs for $n \in \{32, 64, 128, 256, 512, 1\,000, 2\,000, 4\,000, 30\,000, 300\,000, 1\,000\,000\}$ nodes and exponent $\beta \in [1, 4]$. A graph with n nodes and exponent β is obtained by generating a sequence of degrees following a power-law with parameter β (deg(u) gets value d with probability proportional to $1/d^{\beta}$) and generating a graph according to the configuration model (we generate deg(u) half edges for each node u and connect half edges according to a random permutation). For each pair n, β , we have generated 20 random graphs and computed the average VC-dimension of them. Our curves seem to indicate a threshold value near $\beta = 3$ below which the VC-dimension tends to infinity as n grows while the VC-dimension seems to remain constant above (at most three for large n).

7 Conclusion

A first conclusion is that there is room for improvement with respect to the quick estimation of the VC-dimension, especially concerning good upper bounds. Our work can be directly

applied in the setups, where the hyperedges are induced by the open neighborhoods or where only the neighborhoods of a subset of vertices are considered as hyperedges. In future works, besides studying the VC-dimension of power-law random graphs as mentioned above, we will consider extending our algorithm to the practical computation of other VC-dimension parameters on graphs (e.g., the distance VC-dimension, see [8]), and of the VC-dimension of arbitrary set systems where the ranges are given explicitly.

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A Proof of Theorem 1

Proof. Our starting point is the FPT-reduction of Downey et al. [13] who showed that computing the VC-dimension of general set systems is W[1]-hard by the following reduction from k-clique detection.

▶ Theorem 6 (Theorem 3 in [13, Sec. 5]). Let G = (V, E) be a graph and $k \leq |V|$ be a parameter. Then G contains a k-clique if and only if there is a shattered k-subset in the set system $\mathcal{H}_G = (X, \mathcal{R})$, where $X = V \times \{1, 2, ..., k\}$ and $\mathcal{R} = \mathcal{R}_0 \cup \mathcal{R}_1 \cup \mathcal{R}_2 \cup \mathcal{R}_3$ with

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 \begin{split} \mathcal{R}_0 &= \{\emptyset\}; \\ \mathcal{R}_1 &= \{\{(v,i)\} \mid v \in V, \ 1 \leq i \leq k\}; \\ \mathcal{R}_2 &= \{\{(u,i),(v,j)\} \mid uv \in E, \ 1 \leq i,j \leq k\}; \\ \mathcal{R}_3 &= \{\{(v,i) \mid v \in V, \ i \in S\} \mid S \subseteq \{1,2,\ldots,k\} \ and \ |S| \geq 3\}. \end{split}
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Let $\mathcal{H}_G = (X, \mathcal{R})$ be defined as in Theorem 6. Note that the cumulative cardinalities of all sets in \mathcal{R} sum up to $\mathcal{O}(kn + k^2m + k2^kn) = \mathcal{O}(k2^kn^2)$.

We construct a graph H_G from \mathcal{H}_G as follows:

- $V(H_G) = X \cup \mathcal{R};$
- for every $x \in X$ and $R \in \mathcal{R}$ such that $x \in R$, add the edge $\{x, R\}$ to $E(H_G)$;
- if $k \geq 3$, for every distinct $x, y \in X$ add the edge $\{x, y\}$ to $E(H_G)$.

Note that the cost of computing H_G is that of computing \mathcal{H}_G , with an additional cost in $\mathcal{O}(|X|^2) = \mathcal{O}(k^2n^2)$ if $k \geq 3$. It is easy to see that if $Y \subseteq X$ is shattered in \mathcal{H}_G , then it is also shattered by the closed neighborhoods in H_G . Thus, Theorem 6 implies that if G has a k-clique, then the VC-dimension of \mathcal{H}_G , and so the VC-dimension of H_G , is at least k.

It remains to show that if G has no k-clique (or equivalently, the VC-dimension of \mathcal{H}_G is strictly less than k), then the VC-dimension of H_G is at most k-1.

If k = 0, then G must be an empty graph. Then, $\mathcal{R} = \mathcal{R}_0$. It implies that H_G is reduced to one vertex, and so, its VC-dimension equals 0.

If k = 1, then G is a stable set which implies $\mathcal{R}_2 = \emptyset$, and we also have $\mathcal{R}_3 = \emptyset$. In particular, $\mathcal{R} = \mathcal{R}_0 \cup \mathcal{R}_1$. By definition, H_G contains a perfect matching between X and \mathcal{R}_1 , plus an isolated vertex for \mathcal{R}_0 . Therefore, the VC-dimension of H_G equals 1.

Assume now that $\omega(G) \leq k-1$ with $k \geq 3$ and suppose for the sake of contradiction that there exists a k-element subset $Y \subseteq V(H_G)$ which is shattered by closed neighborhoods in H_G . By definition, H_G is a split graph with clique X and independent set \mathcal{R} .

 \triangleright Claim. We either have $Y \subseteq X$ or $Y \subseteq \mathcal{R}$.

Proof. The proof is based on observations similar to the ones in [15, Proof of Lemma 11]. Assume that Y intersects both X and \mathcal{R} . Since $|Y| \geq 3$, we can take distinct elements $x,y,z \in Y$ with $x \in X$, $z \in \mathcal{R}$ and y belonging to either X or \mathcal{R} . Suppose first that $y \notin N[z]$. Let $v \in V(H_G)$ be such that $N[v] \cap \{x,y,z\} = \{y,z\}$. Since $y \notin N[z]$ and \mathcal{R} is a stable set, necessarily $v \in X$. But then, $x \in N[v]$ because X is a clique, thus contradicting that $N[v] \cap \{x,y,z\} = \{y,z\}$. As a result, we must have $y \in N[z]$. Let $u \in V(H_G)$ be such that $N[u] \cap \{x,y,z\} = \{x,z\}$. Since $u,y \in N[z]$ and N[z] is a clique, necessarily $y \in N[u]$. Again, the latter contradicts our assumption that $N[u] \cap \{x,y,z\} = \{x,z\}$.

First we suppose that $Y \subseteq X$. Since we assumed that the VC-dimension of \mathcal{H}_G is strictly less than k, Y is not shattered in \mathcal{H}_G . Thus

there exists
$$Z \subseteq Y$$
 that is not a trace of any range in \mathcal{R} . (1)

Since Y is shattered by H_G , there is a vertex $v_Z \in V(H_G)$ such that $N[v_Z] \cap Y = Z$. The definition of H_G and (1) imply that there is no $v \in \mathcal{R}$ such that its neighborhood in H_G satisfies $N[v] \cap Y = Z$ and so we get that $v_Z \in X$. Since $H_G[X]$ is a clique, we conclude that Z = Y. However, there exists a range in \mathcal{R}_3 which contains the entire set Y (namely, the one corresponding to the full set $S = \{1, 2, ..., k\}$), a contradiction with (1).

Finally, we consider the case of $Y \subseteq \mathcal{R}$. Let $y \in Y$ be any vertex. Since Y is shattered, each of the 2^{k-1} subsets of Y that contain y is either the trace of N[y] or the trace of N[x] for some neighbor x of y. Therefore, $|N[y]| \ge 2^{k-1} \ge 4$, and so the range in \mathcal{H}_G corresponding to y has size at least 3, in particular, $y \in \mathcal{R}_3$. For any $Y \subseteq \mathcal{R}_3$, the closed neighborhoods of the vertices of H_G can have the following traces on Y:

- if $v \in \mathcal{R}$, then $N[v] \cap Y$ is equal to $\{v\}$ if $v \in Y$ and \emptyset otherwise;
- if $v = (x, i) \in X$, then $N[v] \cap Y$ contains those vertices of Y that correspond to index sets S with $i \in S$ (see the definition of \mathcal{R}_3).

That is, the neighborhoods of vertices in \mathcal{R} can only induce the empty set and the k singleton traces on Y, and for any $x, y \in V(G)$ and $i \in \{1, 2, ..., k\}$, we have $N[(x, i)] \cap Y = N[(y, i)] \cap Y$. This implies that the number of vertices in X that have pairwise different neighborhoods in Y is at most k. On the other hand, since Y is shattered, we need to obtain each of the 2^k subsets of Y as a trace, which implies that $2^k \leq k + k + 1$, and thus $k \leq 2$, a contradiction.

B Simple bounds

Lemma 7. A k-degenerate graph has VC-dimension at most k+1, and this bound is sharp.

Proof. Let G = (V, E) be a k-degenerate graph and consider a shattered set $X \subseteq V$. Let $Z = X \cup \{v_Y \mid Y \subseteq X\}$, where v_Y denotes an arbitrary vertex such that $N[v_Y] \cap X = Y$. We consider the induced subgraph G[Z] and we iteratively remove all vertices of $Z \setminus X$ with at most k neighbours. Let Z' be the set of remaining vertices. Note that $X \subseteq Z'$. Since G[Z'] is also k-degenerate, it has some vertex x with at most k neighbours. Since we iteratively removed all vertices with at most k neighbours in $Z \setminus X$, then necessarily $x \in X$. Furthermore, all vertices of $Z \setminus Z'$ must be of the form v_Y for some $|Y| \le k$. As a result, the number of neighbours of x in G[Z] is no more than $k + \sum_{i=0}^{k-1} {|X|-1 \choose i}$. However, since X is shattered, and there are $2^{|X|-1}$ subsets of X containing vertex x, we must have $|N[x] \cap Z| \ge 2^{|X|-1}$. In particular, the number of neighbours of x in G[Z] must be at least $2^{|X|-1} - 1$. Suppose by contradiction that k < |X| - 1. Then,

$$\begin{split} k + \sum_{i=0}^{k-1} \binom{|X|-1}{i} &= k + 2^{|X|-1} - \sum_{i=k}^{|X|-1} \binom{|X|-1}{i} \\ &\leq k + 2^{|X|-1} - \sum_{i=|X|-2}^{|X|-1} \binom{|X|-1}{i} \\ &= k + 2^{|X|-1} - \binom{|X|-1}{|X|-2} - \binom{|X|-1}{|X|-1} \\ &= k + 2^{|X|-1} - |X| \\ &< 2^{|X|-1} - 1 \end{split}$$

Table 4 VC-dimension lower bounds computed with maxvisits = 16, 32, 64, 128, 256 (bold values indicate that the bound matches the exact value), and the corresponding execution time, the "read" column corresponds to the time for reading the graph.

	lower-bound time (s					e (s)						
Graph	VC-dim	16	32	64	128	256	read	16	32	64	128	256
BIO-MV-Physical-3.5	5	5	5	5	5	5	0.06	0.05	0.05	0.05	0.05	0.10
BIO-SYS-Aff-Cap-MS-3.5	7	6	6	7	7	7	0.15	0.18	0.18	0.18	0.20	0.23
BIO-SYS-Aff-Cap-RNA-3.5	7	6	6	6	6	7	0.02	0.03	0.02	0.03	0.03	0.04
dip20170205	5	5	5	5	5	5	0.04	0.05	0.05	0.05	0.06	0.07
oregon2-010331	6	5	5	5	5	5	0.01	0.02	0.02	0.02	0.02	0.04
CAIDA-as-20130601	7	6	6	6	6	7	0.07	0.09	0.09	0.09	0.11	0.13
DIMES-201204	7	6	7	7	7	7	0.04	0.05	0.05	0.05	0.05	0.06
as-skitter	8	7	7	8	8	8	6.11	7.04	7.42	7.98	8.24	9.92
p2p-Gnutella09	5	5	5	5	5	5	0.01	0.02	0.02	0.02	0.02	0.03
gnutella31	4	3	3	3	3	3	0.12	0.11	0.11	0.13	0.18	0.26
notreDame	6	3	3	4	5	6	0.74	1.05	1.16	1.12	1.19	1.20
y-BerkStan	7	5	7	7	7	7	3.49	4.87	4.88	5.08	5.39	5.88
ca-HepPh	5	5	5	5	5	5	0.06	0.07	0.10	0.08	0.10	0.14
com-dblp	5	4	4	4	4	0	0.58	0.71	0.78	0.81	0.95	0.00
epinions1	7	6	7	7	7	7	0.25	0.33	0.35	0.36	0.40	0.46
facebook-combined	6	5	6	6	6	6	0.04	0.04	0.05	0.06	0.07	0.10
twitter-combined	7	6	7	6	6	6	0.64	0.70	0.70	0.82	1.03	1.45
t.CAL	3	3	3	3	3	3	3.30	4.61	4.53	4.52	4.53	4.60
t.FLA	3	3	3	3	3	3	1.83	2.45	2.50	2.45	2.44	2.43
buddha	4	3	3	3	3	3	2.13	2.73	2.63	3.27	3.16	4.13
froz	3	3	3	3	3	3	1.43	1.98	2.54	3.02	4.46	6.91
z-alue7065	3	3	3	3	3	3	0.03	0.04	0.04	0.03	0.04	0.03
grid300-10	3	3	3	3	3	3	0.16	0.21	0.23	0.22	0.21	0.20
xgrid500-10	3	3	3	3	3	3	0.50	0.61	0.83	0.62	0.62	0.64
powerlaw2.5	6	6	6	6	6	6	1.38	3.33	3.50	3.36	3.49	3.70

A contradiction. Hence, $|X| \le k + 1$. This is sharp for k = 1 because trees are 1-degenerate, and there exist 2-shattered subsets in trees (e.g., any two leaves in a star with at least three leaves).

Finally, we show upper bounds on the VC-dimension in terms of sizes of maximum and maximal matchings.

▶ **Lemma 8.** Let G = (V, E) be a non-empty graph and M be a maximal matching of G. Then the VC-dimension of G is at most 2|M|. Moreover, if $\nu(G)$ is the size of a maximum matching in G, then we have $VCdim(G) \leq \nu(G) + 1$.

Proof. Since G is non-empty, any maximal matching has at least one edge and thus the statements trivially hold if $\operatorname{VCdim}(G) \leq 2$. Let $X \subseteq V$ be a shattered set of size $\operatorname{VCdim}(G)$. If M covers every vertex of X, then we have $|M| \geq \frac{1}{2} \cdot \operatorname{VCdim}(G)$. Assume that there exists $x \in X$ which is not covered by M. Since M is maximal, each of the at least $2^{\operatorname{VCdim}(G)-1}-1$ neighbors of x need to be covered by M, which implies $|M| \geq \frac{1}{2} \cdot \left(2^{\operatorname{VCdim}(G)-1}-1\right) \geq \frac{1}{2}\operatorname{VCdim}(G)$ for any graph with $\operatorname{VCdim}(G) \geq 3$.

To show that $\operatorname{VCdim}(G) \leq \nu(G) + 1$, it is sufficient to construct a matching where |X| - 1 vertices of X are matched to a vertex outside of X. Since X is shattered, for any $x \in X$, there exists a vertex $v_x \in V(G)$ such that $N[v_x] \cap X = \{x\}$. Observe that we have either $v_x \notin X$ or $v_x = x$ and the second option is only possible if x has no neighbors in X. We build a matching M of size |X| - 1 as follows. First we add all edges $\{x, v_x\}$ to M where $x \in X$ is such that it has at least one neighbor in X. After this, if there is a remaining

set $Y \subseteq X$ which is not yet covered by M, then Y has to be a stable set. Since X is shattered, there is a vertex v_Y such that $v_Y \cap X = Y$. As Y is a stable set, $v_Y \notin Y$ if $|Y| \geq 2$. Thus, we can cover each element of Y (except maybe one) by taking any $y \in Y$, adding $\{v_Y, y\}$ to M, and recursing on $Y' = Y \setminus \{y\}$. In the end, we get a matching of size $|M| \geq |X \setminus Y| + |Y| - 1 = \text{VCdim}(G) - 1$.

Note that stars with at least three leaves have matching number one and VC-dimension two, thus the bound of Lemma 8 is sharp.

C Experiments

C.1 Lower-bound computation

Table 4 gives the lower-bounds obtained by our lower-bound heuristic for various values of maxvisits while KBG uses maxvisits = 64. It also provides the corresponding running times for reading the graph and computing the lower-bound. As a reference, column "read" indicates the time spent for just reading the graph.

We observe a tradeof where increasing maxvisits provides generally a better lower-bound at the cost of a longer running time. Note the exception of twitter-combined for which the best bound is obtained only for maxvisits = 32. We also note that the running time stays within a factor 5 of the time taken for reading the graph, even for maxvisits = 256. The exception of twitter-combined let us think that there is room for improvement of the tuning of our heuristic. For example, the choice of maxvisits/2 for limiting the for loop of ExploreShattered was not intensively explored. However, it already provides lower-bounds which are often exact or one less than the true VC-dimension. This is indeed the case for maxvisits = 128, and almost the case for maxvisits = 64 where notreDame is the only exception with a lower-bound which is two less than the VC-dimension.

Table 5 Some upper-bounds of Lemma 4 compared to lower-bounds and true VC-dimension.

graph	lb64	VCdim	$\lfloor \log \Delta \rfloor + 1$	$\lfloor \log n \rfloor$	degen+1
BIO-MV-Physical-3.5	5	5	11	14	37
BIO-SYS-Aff-Cap-MS-3.5	7	7	12	15	52
BIO-SYS-Aff-Cap-RNA-3.5	6	7	12	13	55
dip20170205	5	5	9	14	22
oregon2-010331	5	6	12	13	32
CAIDA-as-20130601	6	7	12	15	69
DIMES-201204	7	7	12	14	35
as-skitter	8	8	16	20	112
p2p-Gnutella09	5	5	7	12	11
gnutella31	3	4	7	15	7
notreDame	4	6	14	18	156
y-BerkStan	7	7	17	19	202
ca-HepPh	5	5	9	13	239
com-dblp	4	5	9	18	114
epinions1	7	7	12	16	68
facebook-combined	6	6	11	11	116
twitter-combined	6	7	12	16	97
t.CAL	3	3	4	20	4
t.FLA	3	3	4	20	4
buddha	3	4	5	19	6
froz	3	3	4	19	5
z-alue7065	3	3	3	15	3
grid300-10	3	3	3	16	3
xgrid500-10	3	3	3	17	3
powerlaw2.5	6	6	14	19	19

C.2 Upper bounds

Table 5 lists the upper-bounds we can quickly compute on our dataset. The degree upper-bound $\lfloor \log \Delta \rfloor + 1$ where Δ is the maximum degree appears to always be the best one. However, it can be as large as twice the true value and gives a poor confidence bound compared to what we obtained for lower-bounds. The node upper-bound $\lfloor \log n \rfloor$ where n is the number of nodes is almost always greater. The only graph where it matches the degree upper-bound is facebook-combined which has high maximum degree Δ compared to its number n of nodes as it satisfies $\Delta > n/4$ (see Table 1). Finally, the degeneracy upper-bound appears to be good on graphs with low degree, that is road networks and grid like graphs, while it can be very high for graphs with many high degree nodes. Note that it can be quite high even for graphs with relatively low VC-dimension such as ca-HepPh.