Experimental Evaluation of Parameterized Algorithms for Feedback Vertex Set*

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Abstract

Feedback Vertex Set is a classic combinatorial optimization problem that asks for a minimum set of vertices in a given graph whose deletion makes the graph acyclic. From the point of view of parameterized algorithms and fixed-parameter tractability, Feedback Vertex Set is one of the landmark problems: a long line of study resulted in multiple algorithmic approaches and deep understanding of the combinatorics of the problem. Because of its central role in parameterized complexity, the first edition of the Parameterized Algorithms and Computational Experiments Challenge (PACE) in 2016 featured Feedback Vertex Set as the problem of choice in one of its tracks. The results of PACE 2016 on one hand showed large discrepancy between performance of different classic approaches to the problem, and on the other hand indicated a new approach based on half-integral relaxations of the problem as probably the most efficient approach to the problem. In this paper we provide an exhaustive experimental evaluation of fixed-parameter and branching algorithms for Feedback Vertex Set.

1 Introduction

The Feedback Vertex Set problem asks to delete from a given graph a minimum number of vertices to make it acyclic. It is one of the classic graph optimization problems, appearing already on Karp’s list of 21 NP-hard problems [24]. In this work, we are mostly focusing on fixed-parameter algorithms for the problem, that is, exact (and thus exponential-time, as we are dealing with an NP-hard problem) algorithms whose exponential blow-up in the running time bound is confined by a proper parameterization. More formally, a fixed-parameter algorithm on an instance of size $n$ with parameter value $k$ runs in time bounded by $f(k) \cdot n^c$ for some computable (usually exponential) function $f$ and a constant $c$ independent of $k$.

Feedback Vertex Set is one of the most-studied problems from the point of view of parameterized algorithms. A long line of research [4, 14, 15, 29, 23, 11, 18, 9, 8, 3, 10] lead to a very good understanding of the combinatorics of the problem, multiple known algorithmic approaches, and a long “race” for the fastest parameterized algorithm under the parameterization of the solution size (i.e., the parameter $k$ equals the size of the solution we are looking for). Among many approaches, one can find the following:

1. A classic randomized algorithm of Becker et al. [3] with expected running time $4^k n^{O(1)}$.
   
   This algorithm is based on the following observation: after performing a set of simple reductions that reduce the graph to minimum degree at least 3, at least half of the edges of the graph are incident with solution vertices, so a random endpoint of an edge chosen uniformly at random is in the solution with probability at least $1/4$.

2. A line of research of branching algorithms based on iterative compression and an intricate measure to bound the size of the branching tree [9, 8, 26], leading to a simple $3.62^k n^{O(1)}$-time deterministic

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algorithm [20]. These algorithms solve a “disjoint” version of the problem where, given a set $U \subseteq V(G)$ such that $G - U$ is a forest, one seeks for a solution disjoint with $U$ of size at most $k$. A branching algorithm picks a vertex $v \in V(G) \setminus U$ and includes it in the solution (deletes and decreases $k$ by one) or moves to $U$. The crucial observation is that here, apart from $k$, the number of connected components of $G[U]$ is a useful potential to measure the progress of a branching algorithm: if the branching pivot $v$ has many neighbors in $U$, then moving it to $U$ decreases the number of connected components significantly.

3. A polynomial-time algorithm for Feedback Vertex Set in subcubic graphs [8] (see also [26] for a simpler proof), used heavily in other approaches as a subroutine. The algorithm builds on a reduction from Feedback Vertex Set in cubic graphs to the matroid matching problem in graphic matroids, which is polynomial-time solvable.

4. A Monte Carlo algorithm running in time $3^k n^{O(1)}$ via the Cut&Count technique [10]. The Cut&Count technique is a generic framework for turning algorithms with running time $2^{O(t \log t)} n^{O(1)}$ for connectivity problems in graphs of treewidth bounded by $t$ into Monte-Carlo algorithms with running time bounds $c^t n^{O(1)}$ by changing the “connectivity” requirement into a modulo-2 counting requirement. For Feedback Vertex Set, the base of the exponent has been optimized to $c = 3$, and has been proven to be optimal under the Strong Exponential Time Hypothesis.

5. A surprisingly simple branching algorithm by Cao [7] with a running time bound $8^k n^{O(1)}$. Cao [7] shown that the following algorithm has running time bounded by $8^k n^{O(1)}$: after applying simple reduction rules, one either branch on maximum-degree vertex or, if the maximum degree is at most 3, solve the instance in polynomial time.

6. A branching algorithm based on intricate half-integral relaxation due to Imanishi and Iwata [19, 20] with running time bound $4^k k^{O(1)} n$.

7. A formulation of a problem as an integer linear program that can be solved using a general ILP solver. For every vertex $v \in V(G)$ one can create a variable $x_v$ denoting whether we put the given vertex in the solution ($x_v = 1$), or we leave the vertex in the graph ($x_v = 0$). This special case of integer linear programming, where every variable is not only an integer, but also binary, is also on Karp’s list of 21 NP-hard problems [24]. However, there exist several general integer linear program solvers that often work very efficiently in practice.

Having defined variables for the set of vertices, one can create constraints twofold. The first approach is to create a constraint for every cycle, stating that the sum of variables in it has to be at least equal to 1. It means the same as the definition of the problem — we need to delete at least one vertex in every cycle.

Second way to define the problem in terms of integer linear programming is to create constraints for every subset of vertices. For those subsets we want to delete as few vertices as possible to make it a tree. In this approach we need more variables than only for vertices, but the worst-case number of constraints is smaller.

In both of these approaches there is a small drawback – it is possible that there will be exponential number of constraints. For this reason, the algorithms based on these ideas need to add the constraints dynamically and check whether the solution using only a part of constraints implies the solution for the whole graph.
Another line of research concerns so-called polynomial kernels for the problem \([6, 5, 30, 20]\).

Parameterized Algorithms and Computational Experiments challenge is an annual programming challenge started in 2016 that aims to “investigate the applicability of algorithmic ideas studied and developed in the subfields of multivariate, fine-grained, parameterized, or fixed-parameter tractable algorithms”. With three successful editions so far \([12, 13, 16]\) and the fourth one just announced, PACE continues to bring together theory and practice in parameterized algorithms community.

The first edition of PACE in 2016 featured Feedback Vertex Set as the problem of choice in Track B. The winning entry by Imanishi and Iwata, implementing the aforementioned algorithm based on half-integral relaxation, turned out to outperform the second entry by the second author of this work \([28]\), based on the algorithm of Cao \([7]\). The winning margin has been substantial: out of 130 test cases, the winning entry solved 84, while the second entry solved 66.

These results indicated the branching algorithms based on half-integral relaxation of the problem as potentially most efficient approach to Feedback Vertex Set in practice. Furthermore, experimental results of Akiba and Iwata \([2]\) showed big potential in a branching algorithm based on the same principle for the Vertex Cover problem.

In the light of the above, we see the need to rigorously experimentally evaluate different approaches to Feedback Vertex Set. While the results of PACE 2016 indicate algorithms based on half-integral relaxation as potentially fastest, a lot of differences may come from the use of different preprocessing routines, different choice of lower bounding or pruning techniques, or even simply different data structures handling basic graph operations.\(^1\)

In this work, we offer such a comparison. We implement a number of branching strategies mentioned above. Our implementations use the same data structures for handling graphs, the same implementations of basic graph operations, the same basic reduction rules such as suppressing degree-2 vertices, and the same branching framework. Most of the tested approaches differ only at a small fraction of code: they usually differ by the choice of the branching pivot, and some use one or more approach-specific reduction rules.

In our experiments, we follow up the set up of PACE 2016: we take their 230 instances (100 public and 130 hidden, on which evaluation took place) as our benchmark set, allow each algorithm to run for 30 minutes on each test instance.

The paper is organized as follows. In Section 2 we discuss the studied approaches and some technical details of the implementations. Section 3 discusses the setup of the experiment. Section 4 presents results, while Section 5 concludes the paper.

## 2 Studied algorithms

Most of the known branching algorithm for Feedback Vertex Set, in particular all algorithms studied in our work, follow the following general framework.

Every instance to be solved by a recursive branching algorithm consists of a multigraph \(G\), a set \(U \subseteq V(G)\) of undeletable vertices and allowed budget \(k\) for solution size. The goal is to find a set \(X \subseteq V(G) \setminus U\) of size at most \(k\) such that \(G - X\) is a forest. Each branching step consists of picking a vertex \(v \in V(G) \setminus U\) and branching into two cases: either \(v\) gets picked to a solution (and the algorithm recurses on the instance \((G - \{v\}, U, k - 1)\)) or moved to set \(U\) (and the algorithm recurses on the instance \((G, U \cup \{v\}, k)\)).

The intuition of the progress of the algorithm is as follows. In the first branch, the budget \(k\) gets decreased. For the second branch, note that the algorithm can safely terminate for instances with \(G[U]\) not being acyclic. Thus, if the branching pivot \(v\) has \(d\) neighbors in \(U\), then the number of connected components of \(G[U \cup \{v\}]\) decreases by \((d - 1)\) as compared to \(G[U]\).

\(^1\)A good example here is as follows. In Feedback Vertex Set, it is natural to keep the graph in the form of adjacency lists, as the considered graphs are usually of constant average degree. However, given an edge \(uv\), it is not clear whether the vertex \(u\) in its adjacency list should only store the vertex \(v\), or also a pointer to the position where \(v\) keeps \(u\) in its adjacency list. On one hand, such pointers greatly simplify the operations of deleting a vertex or contracting an edge. On the other hand, they effectively double the size of the graph data structure, increasing the cost of copying the graph in the branching step.
In the literature, a number of simple reduction steps are known that are performed by all our algorithms:

1. If \( k \) gets negative or \( G[U] \) is not acyclic, stop.
2. Remove all vertices of degree at most 1.
3. If two vertices are connected by more than 2 parallel edges, reduce their multiplicity to 2.
4. If there exists a vertex \( v \) that has a self-loop, or there exists a single connected component \( D \) of \( G[U] \) more than one edge incident with \( v \) and a vertex of \( D \) (i.e., there exists a cycle \( C \) in \( G \) with \( v \) being the only vertex of \( V(C) \setminus U \)), then greedily include it in the solution (i.e., delete it and decrease \( k \) by one).
5. Suppress vertices of degree 2. That is, if a vertex \( v \) is of degree 2 with incident edges \( vu \) and \( vw \) present, delete \( v \) and replace it with an edge \( uw \).
6. If there exists a vertex \( v \) with two neighbors \( u \) and \( w \), such that \( vu \) is a single edge and \( vw \) is a double edge, greedily include \( w \) in the solution.

For efficiency, the above reduction rules are implemented in the form of a queue of vertices to reduce: when

- split into connected components
- solving subcubic instances
- solution lower bound

Between branching step, the algorithm is allowed to perform a number of reduction (preprocessing) steps. Here, we observed that it is often the case that if the current graph is disconnected, the algorithm is allowed to perform a number of reduction (preprocessing) steps.

Other preprocessing steps used by some of our algorithms are:

- split into connected components
- solving subcubic instances
- solution lower bound

Consider an instance \((G, U, k)\) and let \(v_1, v_2, \ldots\) be the vertices of \(V(G) \setminus U\) in the nonincreasing order of degrees in \(G\). If \((G, U, k)\) admits a solution \(X\) of size \(j\), then \(G - X\) has at least \(|E(G)| - \sum_{i=1}^{j} \deg_G(v_i)\) edges. On the other hand, if \(G - X\) is a forest, it has less than \(|V(G)| - j\) edges. Consequently, we can safely stop if for all \(0 \leq j \leq k\) we have that

\[
|E(G)| - \sum_{i=1}^{j} \deg_G(v_i) \geq |V(G)| - j.
\]

The above pruning strategy has been used in the entry of Imanishi and Iwata [19].
Unless otherwise noted, all our implementations split instances into connected components. We also compare a number of selected approaches without this preprocessing step to see its impact on performance.

The algorithm of Cao [7] uses all aforementioned simple reduction rules as well as the solver of subcubic instances. On a branching step, it simply chooses the vertex of highest degree. As shown by Cao, such an algorithm has running time bound $8^k n^{O(1)}$. We also test a variant of the algorithm of Cao that first branches on vertices incident with double edges, a variant that does not use the subcubic instance solver, and a variant that prunes the search tree via the aforementioned lower bound.

2.1 Approximation and iterative compression

The algorithms of [9, 8, 26] operate in the framework of iterative compression. That is, their central subroutine solves a seemingly simpler problem, where additionally a slightly too large solution $Y$ is given, and the algorithm first branches on the vertices of $Y$ (putting each $y \in Y$ into the solution or into set $U$). Since at the beginning $G - Y$ is a forest, and every vertex of $Y$ is either deleted or put into $U$, we obtain the property that $G - U$ is a forest. This greatly helps in the analysis.

In the literature, the set $Y$ is traditionally taken from the iterative compression step. One picks an order $V(G) = \{v_1, v_2, \ldots, v_n\}$ and solves iteratively Feedback Vertex Set on graphs $G_i = G[\{v_1, v_2, \ldots, v_i\}]$. Given a solution $X_{i-1}$ to $G_{i-1}$, one can set $Y = X_{i-1} \cup \{v_i\}$ for $G_i$.

However, such an approach leads to multiple invocation of the same algorithm, and a substantial multiplicative overhead in the running time bound. In our algorithms, we instead find $Y$ via a simple heuristic: reduce the graph via simple reductions as long as possible and, when impossible, delete the vertex of highest degree. In Section 4 we discuss the performance of this heuristic on our test data.

Our branching framework keeps a queue of branching hints and, if nonempty, the algorithm always branches on a vertex from the queue. For algorithms based on iterative compression, the queue is initiated by an approximate solution found by our heuristic. This corresponds to passing the set $Y$ to the algorithms based on iterative compression, but allows to reduce some of the vertices of the set $Y$ by reductions after a number of branching steps. If an algorithm does not use iterative compression, the queue is empty through the entire run of the algorithm.

The algorithm of [9] implements the iterative compression framework and branches on a vertex of $V(G) \setminus U$ that is incident to maximum number of edges leading to $U$. As shown in [9], such an algorithm has $5^k n^{O(1)}$ time bound guarantee.

The algorithm of [26] is arguably a simplification of the arguments of [8], so we implement only the first one. It modifies the algorithm of [9] in the following way:

- It leaves alone vertices $v \in V(G) \setminus U$ that are of degree 3 and all their incident edges lead to $U$ (such vertices are called henceforth tents). The crux is that if every vertex $v \in V(G) \setminus U$ is a tent, then we can apply the polynomial-time algorithm of [8, 26] to the instance. In other words, tents form a “polynomial-time solvable” part of the instance.
- Given a vertex $v \in V(G) \setminus U$ of degree 3 with exactly one neighbor $u$ in $V(G) \setminus U$ (and other 2 neighbors in $U$), it proceeds as follows:
  - subdivide the edge $uv$ with a new vertex $w \in U$; note that this does not change the set of feasible solutions to the Feedback Vertex Set problem;
  - marks $u$ irreducible for the reduction suppressing degree-2 vertices.

Note that this operation turns $v$ into a tent, while reducing the number of vertices of $V(G) \setminus U$ that are not tents.
- It applies the solver for subcubic instances if every vertex of $V(G) \setminus U$ is a tent.

As shown in [26], such an algorithm has $3.62^k n^{O(1)}$ time bound guarantee.
Inspired by the methods of choice of branching pivots of the algorithms \cite{9,8,20}, we also test a variant of Cao’s algorithm where the choice of the branching pivot is as in \cite{9}: vertex with maximum number of neighbors in $U$ (but, contrary to \cite{9}, no iterative compression).

Additionally, we check how much the algorithms can be sped up by adding pruning via the aforementioned lower bound and if the Cao’s algorithm can benefit from the use of iterative compression.

2.2 Branching based on half-integral relaxation

Iwata, Wahlström, and Yoshida showed a generic approach to numerous transversal problems via half-integral relaxations \cite{21}. They are all based on the following principle: a half-integral relaxation of a variant of the problem is defined and shown to be polynomial-time solvable. Furthermore, the solution to the half-integral relaxation has some persistency property: it either indicates some greedy choice for the integral problem, or indicates a good branching pivot. In this approach, the time needed to find a solution of the half-integral relaxation is critical.

The algorithms of \cite{21} run in linear time for edge deletion problems, but unfortunately for vertex-deletion problems the subroutine that finds the half-integral relaxation requires solving linear programs. The main contribution of Iwata \cite{20} (implemented in the PACE 2016 entry by Iminishi and Iwata \cite{19}) is a combinatorial linear-time solver for the half-integral relaxation in the special case of \textsc{Feedback Vertex Set}. We reimplement this solver in our branching framework.

Iwata \cite{20} also observed that the half-integral relaxation can be used to find a polynomial kernel for the problem, improving the previous seminal kernel of Thomassé \cite{30}. Apart from the reduction rules mentioned before, this kernel employs another involved reduction rule, applicable on vertices of degree more than $2k$. All our implementations based on a half-integral relaxation implement this preprocessing step as well.

The half-integral relaxation of \cite{21,20} does not solve \textsc{Feedback Vertex Set} directly, but rather given an undeletable vertex $u \in U$, tries to separate an acyclic connected component (i.e., a tree) containing $u$ from the rest of the graph. On high level, the branching strategy of the algorithm is as follows. If $U = \emptyset$, we branch on any vertex; we choose one of highest degree here for efficiency. Otherwise, we use a vertex $u \in U$ as a root for the half-integral relaxation. The persistency properties of the relaxation ensure that we can perform a greedy step unless the relaxation put values 0.5 on all neighbors of $u$. If this is the case, we branch on a neighbor of $u$; we choose highest-degree neighbor here for efficiency. Note that once a tree component with $u$ gets separated, simple reduction rules delete it from the graph.

Since the kernelization routine of \cite{20} is computationally expensive, it is not obvious if one should apply it at every step. We experiment with two variants: when we run the kernelization step at every step, or only at steps with $U = \emptyset$. Furthermore, we also check how much pruning with the lower bound heuristic or solver of subcubic instances helps.

2.3 Using a generic Integer Linear Program solver

As mentioned in the introduction section, we can create an integer linear programming formulation of our problem in two ways. For purpose of the tests, we have chosen the cycle-based implementation created by organizers of PACE challenge \cite{27}, which as a linear program looks like:

\[
\begin{align*}
\text{minimize} \quad & x_1 + x_2 + \ldots + x_{|V(G)|} \\
\text{subject to} \quad & x_v \in \{0, 1\} \\
& x_{c_1} + x_{c_2} + \ldots + x_{|C|} \geq 1 \quad \forall C = (x_{c_1}, \ldots, x_{|C|})
\end{align*}
\]

One may notice that the number of constraints on the cycles can be exponential in the size of the graph. For example in a clique on $n$ vertices, there are roughly $n!$ cycles. For this reason we cannot just add all of the constraints at once — we want to make the graph smaller first and then add the constraints dynamically.
The implementation first applies the reductions 2. – 5., mentioned in the beginning of the chapter, to the input graph. Next it removes the bridges between connected components of the reduced graphs. Afterwards it processes every component of the graph separately in the steps described below.

- Put a heuristic solution as the starting points of the ILP \( v \in X \Rightarrow x_v = 1 \). Here we use a slight modification of the heuristic: we take shortest cycles one by one, find the highest degree vertex \( v \) in them, and add \( v \) to \( X \). Repeat this as long as there are any cycles in \( G \).

- Find a set of cycles of shortest length (run BFS from every vertex) and add them to our constraint pool. All other constraints are lazy, which means at first the solver finds a feasible solution without using them. Later if evaluation turns out to violate them, they are being added and the solution is recalculated.

- In the callbacks, we check whether the solver returned a set \( X \) that actually removes all the cycles in \( G \). If it is the case, then we have found a solution for our problem. Otherwise, we return to the point of adding new shortest-cycle constraints. In order to only add new cycles, we delete from the graph vertices, which were part of the latest removal set \( X \), and then compute shortest cycles.

During our tests we have used Gurobi 8.0 optimizer.

3 Experiment setup

3.1 Hardware and code

The experiments have been performed on a cluster of 16 computers at the Institute of Informatics, University of Warsaw. Each machine was equipped with Intel Xeon E3-1240v6 3.70GHz processor and 16 GB RAM. All machines shared the same NFS drive. Since the size of the inputs and outputs to the programs is small, the network communication was negligible during the process.

The code has been written in C++ and is available at [25] or at project’s webpage [1].

3.2 Test cases

As discussed in the introduction, we repeat the setup of the PACE 2016 challenge [12]. At PACE 2016, the organizers gathered 230 graphs from different sources [27]. A subset of 100 of them has been made public prior to the competition deadline, and the final evaluation has been made on the hidden 130 instances. We run every tested algorithm on each of the 230 instances with 30 minutes timeout.

Out of the test instances, we gathered two subsets to compare actual running times of the algorithm. The first set, \( A \), consists of test cases solved by all algorithms, but with substantial running time of some of them. The second set, \( B \), is defined similarly, but with regards only to the algorithms that use pruning via the lower bound. The third set, \( C \), is a subset of \( A \) and \( B \) that has been solved by the ILP solver.

More precisely, set \( A \) consists of the following 14 tests:

- hidden_001
- hidden_007
- hidden_012
- hidden_056
- hidden_056
- hidden_065
- hidden_083
- hidden_099
- hidden_106
- public_011
- public_014
- public_037
- public_069
- public_076
- public_086

The set \( B \) consists of the following 7 tests:

- hidden_022
- hidden_041
- hidden_068
- hidden_088
- public_035
- public_066
- public_067

The set \( C \) consists of the following 18 tests:

- hidden_001
- hidden_007
- hidden_012
- hidden_022
- hidden_056
- hidden_065
- hidden_068
- hidden_083
- hidden_088
- hidden_099
- hidden_106
- public_011
- public_014
- public_035
- public_069
- public_076
- public_086
- public_087

We also gather sizes of approximate solution found by our heuristic and compare it with the optimal size found by the algorithm.
3.3 Reduction rules measures

In order to fully compare the branching algorithms, we have decided to measure how big impact do actually the reduction rules have on performance of each algorithm. For this purpose we introduce the following measures. We computed for every test and for the top 3 algorithms in our benchmark.

**initial reductions** How well the reductions work before the first use of branching. There are both numbers of how many vertices/edges were reduced initially and how big part of the whole graph it was (in percents).

Here most of the algorithms use the same set of initial rules, except for the algorithm by Imanishi and Iwata that uses an extra kernelization step it uses.

**average over recursive subcalls** Measurements of how big impact do the reductions have during the runtime of branching. We count an average number of vertices/edges reduced during one step of branching (averaged over all recursive calls to the branching subroutine). In this group we have also counted how big impact did reduction steps have only in branches where the number of vertices in part of graph remaining to solve is between 20 and 40.

**connected components** In this part we have calculated how big are the connected components separated from the largest connected component of the input graph by our "split into connected components" routine.

One may find exact results of above measurements for each of the tests in our repository [25].

We analyze performance of reductions on each algorithm by grouping tests with following methods.

- **Average over all tests finished by all of the three algorithms.**
- **Score function is calculated in two ways.** In the comparison of the initial reductions, since we are comparing only two algorithms, the score is the number of tests on which reductions in one solution have strictly better results than in the others. In the other comparisons, since they compare three approaches, for every test we divide 1 point between all solutions with best result on the test. We report the percentage of total points scored by each algorithm.

While the measurement of initial reductions is a solid comparison of the simple reduction rules with the addition of the sophisticated kernelization rule, the remaining measurements are just our quite crude attempts at measuring how well does the algorithm perform reductions of the graph during the whole branching routine, and should be treated with a grain of salt.

4 Results

A full table with running times of each program at each test can be found at the repository [25].

4.1 Performance of the approximation heuristic

We compared the performance of the approximation heuristic discussed in Section [2.1] (i.e., greedily take the highest-degree vertex after applying the simple reduction rules) with the size of the optimum solution that was known to us on 127 instances. The results are in Table [1]. On only 8 instances, the approximate solution was more than one vertex larger than the optimum one. Consequently, the approximate solution can serve well as the basis for iterative compression.
Table 1: Comparison of the size of the approximate and exact solution found on 127 instances.

| algorithm | optimizations | solved instances | total time (MM:SS.ms) |
|-----------|---------------|-------------------|-----------------------|
|           |               | all public hidden | set A set B set C    |
| Cao       | +             | 100 48 52         | 35:17.21 - -          |
| CFLLV     | +             | 91 44 47          | 35:16.06 - -          |
| KP        | + +           | 101 49 52         | 36:22.48 - -          |
| Cao       | + +           | 101 48 53         | 37:31.83 - -          |
| CFLLV     | + +           | 91 44 47          | 31:23.63 - -          |
| KP        | + + +         | 101 49 52         | 32:00.49 - -          |
| Cao       | +             | 91 43 48          | 38:00.78 - -          |
| II        | +             | 92 46 46          | 34:51.17 - -          |
| II/kernel | +             | 90 45 45          | 75:11.16 - -          |
| Cao       | + + +         | 123 62 61         | 0:19.28 10:38.88 1:54.11 |
| Cao/double| + + +         | 122 62 60         | 3:22.52 21:31.54 15:57.95 |
| Cao/undel | + + +         | 118 58 60         | 0:35.67 8:39.05 8:12.94 |
| Cao       | + + + +       | 123 62 61         | 0:19.99 10:37.88 1:53.48 |
| Cao/double| + + + +       | 122 62 60         | 3:13.71 14:31.66 8:45.33 |
| Cao/undel | + + + +       | 118 58 60         | 0:36.88 8:12.80 7:47.82 |
| CFLLV     | + + +         | 117 57 60         | 0:37.61 8:12.88 7:47.68 |
| KP        | + + + +       | 118 58 60         | 0:38.95 8:28.46 8:03.57 |
| Cao       | + + + +       | 122 61 61         | 0:22.94 10:28.32 1:58.57 |
| II        | + +           | 117 58 59         | 1:36.79 25:31.95 24:29.82 |
| II/kernel | + +           | 109 55 54         | 7:24.42 - -          |
| II        | + +           | 118 59 59         | 1:37.08 25:33.76 24:33.01 |
| ILP       |               | 140 62 78         | - - 3:46.37          |

Table 2: Comparison of different algorithms. The first column indicates the base of the algorithm: Cao’s [7], CFLLV for Chen et al. [9], KP for Kociumaka and Pilipczuk [26], II for Imanishi and Iwata [19, 20] and ILP for Integer Linear Programming. II/kernel stands for the algorithm of [19, 20] that runs the kernelization step more often, namely at every branching step. Cao/double stands for the algorithm of [7] that first branches on double edges. Cao/undel stands for the algorithm of [7] that chooses branching pivot with regards to maximum degree of undeletable vertices. In the optimizations columns, CC stands for splitting into connected components, deg3 for the use of solver of subcubic instances, LB for the use of pruning with the lower bound, and IC stands for iterative compression.

4.2 Comparison

We have run 22 different algorithms on the whole test data. A CSV file with full results is available at the repository [25]. Table 2 contains aggregated values: number of solved test instances within the time limit (30 minutes per instance) and the total running time on sets A, B, and C. Please see the caption of Table 2 for description of the notation used in the table.

The first nine algorithms do not use pruning via the lower bounding technique, and the first three do not use splitting into connected components. They are mostly meant to compare basic approaches.

Without the lower bound pruning, the best approaches are Cao’s [7] and Kociumaka-Pilipczuk [26], and they seem to be rather incomparable. The other algorithm based on iterative compression of Chen et al. [9] is clearly outperformed by the other two, and the same holds for the branching algorithm based on
half-integral relaxation [19, 20].

The first three rows differ from rows 4-6 by the usage of splitting into connected components. They show that the effect of this improvement is small, and even hurt a bit Cao’s algorithm [7].

The 7th row treats Cao’s algorithm [7] without the solver of the subcubic instances. It indicates that this solver is essential for the performance of Cao’s algorithm.

The 9th row treats the Imanishi-Iwata algorithm [19, 20] with more often application of the kernelization step, namely at every branching step (not only at the ones with \( U = \emptyset \)). It shows that the step is too expensive to execute it that often.

Let us now discuss the algorithms with the lower bound pruning step. First, the results show that the pruning step greatly improves performance for all algorithms.

With regards to the variants of Cao’s algorithm [7], the results show that any mutation of the branching pivot rule here is undesirable. Also, adding the iterative compression step does not seem to have any particular impact on the performance. Interestingly, the negative effect of dropping the solver of the subcubic instances mostly disappear if one adds the pruning step.

For the branching algorithm based on the half-integral relaxation [19, 20], the last three rows of Table 2 covering II again confirm the corollary that more often application of the kernelization step is undesirable. There is little difference with addition of the solver of subcubic instances (the main difference comes from the fact that the test set contains one huge cubic graph, public_84, which is not amenable to any branching technique we tested).

Finally, in our experiments the best variant of Cao’s algorithm [7] with the pruning slightly outperforms the best variant of the branching algorithm based on half-integral relaxations [19, 20]. However, the difference (5 tests more and roughly 3× speed-up on sets \( A \) and \( B \)) is not big enough to decisively conjecture its advantage.

First, it is possible that a 3× speed-up can be gained by low-level optimizations of the solver of the half-integral relaxations. Arguably, Cao’s algorithm [7] is much simpler and thus easier to optimize. Second, it may be also an artifact of the chosen test data: there are 5 tests in the data set that were solved by the penultimate algorithm in Table 2 but not by the 10th (and 10 tests vice-versa). That is, there are types of instances solved significantly faster by one of the approaches but not by another.

We conclude with a remark about comparison with PACE’16 results [12], where the entry of Imanishi and Iwata [19] solved 84 hidden instances while the entry of the second author [28], based on Cao’s algorithm [7], solved 66. While this data seemingly stands in contradiction with the results in Table 2, one should note that the two entries differ significantly in other internals. Most importantly, the first entry used pruning with the lower bound, while the second one did not. Other difference includes: removal of bridges in the first entry vs splitting into 2-connected components in the second, and the use of bounded treewidth subroutine in the second entry. In other words, our results indicate that the big difference in the performance of the first two entries at PACE 2016 were mainly caused by the difference in preprocessing routines and pruning heuristics (and possibly low-level optimizations) rather than in the underlying base branching algorithm.

The last algorithm in the table is the solution based on integer linear programming. It is the solver that the organizers used while preparing the contest. We may notice that at the public tests set it solves pretty much the same amount of tests (62) as the best branching algorithms. On the other hand, the difference can be noticed on the set of hidden tests, where it solves 17 tests more than any other solution we have tested. We expected ILP solution to greatly outperform all of the solutions on both sets of tests, so the results can be found a little bit surprising.

4.3 Impact of the reduction rules

In this section we will analyse the impact of reduction rules on each of the branching algorithms that achieved the best results in our tests, i.e. Cao, KP, and II. The measures were introduced in Section 3.3. In what follows, each average over tests is taken over all tests solved by all three solvers taken into account.
Table 3: Initial reductions: the impact of reduction rules before start of a branching. $\Delta n$ stands for the decrease in the number of vertices, $\Delta m$ for the decrease in the number of edges. Score indicates the number of tests where one algorithm strictly outperformed the second one.

Initial reductions. On average, reduction rules get rid of almost 50% of the graph even before the start of branching. This shows a great impact of preprocessing on further performance of the algorithms.

From the $\Delta n$ score, we may notice that II has at least as high result as the other algorithms in $\Delta n\%$ on every test. Recall that all algorithms except II use the same set of reductions, while the II algorithm uses an extra sophisticated kernelization step for vertices of large degree. The II algorithm achieves better performance on 14 tests (in terms of the number of reduced vertices) due to the extra kernelization step.

Note that II looses in 4 tests when calculating $\Delta m$ score. The reason for this is pretty simple: the kernelization step, while changing structure of the graph is not only deleting vertices and edges, but also adding some edges to preserve the potential solution unchanged. In those tests it happened that more edges were added than deleted.

Table 4: Average report of reduction rules performance in the run time of each branching. For each test, we distribute one point between the best performing algorithms, and then present the percentage of all points got by each of the algorithms.

Average over recursive calls. As mentioned in the previous section, we present the number of reduced vertices and edges averaged over all recursive calls of the branching algorithm and over all calls that are applied to graphs with at least 20 and at most 40 vertices.

The results of the first part clearly show that Cao reduces most vertices/edges on average during one step of branching. The explanation for this result is the way we choose vertex for branching in implementation of Cao. Choosing vertex of maximum degree (with most edges) for branching favors creation of new reducible vertices.

Even though relative order of algorithms for AVG20-40 vertices score hold, one may wonder why that is not true for score on AVG20-40 edges reductions. It might happen that Cao’s average reduced edges number for instances above 40 is close to some other algorithm (both of the others gain for instances in 20-40). Branching on highest degree vertices when the instance is still big, makes us get rid of huge part of the edges, i.e., the number of edges Cao has left to reduce becomes smaller much faster than for the other algorithms. For this reason, when we get to medium sized instances it may turn out that some other algorithm that had close average number of reduced edges has it higher now and Cao loses point from its score on that test.

Connected components. In the table covering connected components we can see that on average the KP and II algorithms seem to separate the biggest number of vertices by cutting out connected components from
Table 5: Connected components separation performance results. For each test, we distribute one point between the best performing algorithms, and then present as the score the percentage of all points got by each of the algorithms.

|                        | Cao   | KP    | II    |
|------------------------|-------|-------|-------|
| Average total separated vertices | 3201.93 | 20810.23 | 18539.40 |
| Score total separated vertices    | 29.43% | 35.29% | 35.29% |

the graph. If we look at the results for separate tests, it seems that on average sizes of connected components cut out from graphs are small and similar for all of the algorithms. From this we could infer that KP and II cut out connected components much more often than Cao. It stands opposite to the intuition of how the II works: it tries to solve whole connected components on its own during the branching step. The reason for this abnormality in this measure is 1 test on which both KP and II did a huge number of separations (higher than the sum of all the other tests together), while Cao did almost none.

For this reason we find the measure of score much more meaningful this time. It says that every algorithm won on almost the same amount of tests and the impact of connected components separation on KP and II is not that much bigger than on Cao.

4.4 Comparison of solved tests

We have noticed that the 5 test difference between Cao and II algorithms does not mean straightforward that Cao solved 5 tests that II did not. It actually is a little bit different – II solved 5 tests that Cao did not and there were 10 tests the other way. We have decided to note down the tests on which they differ and analyze them in order to find some structural property that makes one algorithm perform better on one set than the other. Below you can find the list of tests solved only by one of the algorithms.

List of tests that Cao solved and II did not:
- hidden_023
- hidden_035
- hidden_064
- hidden_066
- hidden_109
- public_032
- public_034
- public_073
- public_080
- public_087

List of tests that II solved and Cao did not:
- hidden_071
- hidden_079
- hidden_080
- public_023
- public_043

While analyzing the tests, we have first applied the basic reductions on the graphs and than generated statistics like number of vertices, number of edges, number of double edges and the distribution of degrees in the graph. You may find the statistics in our repository. Afterwards we used gephi to draw and look at the graphs.

We have found out that in the tests that Cao solved there is hardly any regularity to be found (with one exception of test hidden_109, which is an almost-regular graph with low degrees). In these tests there usually are some vertices of high degree and many low degree vertices. Such structure of the test makes Cao algorithm work fast: it branches on the vertices of highest degree first. Thanks to that we quickly get rid of huge part of the graph: having decided whether such a vertex is in solution $X$ or in safe $U$ set, degrees of many vertices may decrease. Now having many vertices of low degrees helps: the decrease in degrees affects a lot of them, which at the same time makes the basic reductions quickly become applicable again. At the same time on such tests II is much computationally-heavier and is unable find any structure that it could solve quickly, which is reason for this difference.

On the other hand looking at the tests solved by II and not by Cao we can see a lot of regularity. Here, we have regular graphs with degrees in range from 10 to 16. Because of it, Cao is completely ineffective on these instances. It has to branch through a lot of vertices to achieve anything, which is almost impossible with limited time it is given. II seems to work on such graphs more efficiently. It generates less branches and gets rid of such grid-like structures faster.
5 Conclusions

We have conducted a thorough experimental evaluation of various parameterized algorithms for the Feedback Vertex Set problem, following the setup of Parameterized Algorithms and Computational Experiments Challenge from 2016 [12]. Our results does not confirm greater advantage of the approach based on half-integral relaxations that was suggested by PACE’16 results, but rather suggest that lower bounding techniques and low-level optimizations were decisive at PACE’16.

On the other hand, the approach via half-integral relaxation turned out to be almost on par with the best variant of Cao’s algorithm [7]. This still indicates big potential in this approach, in particular in the light of the recent (theoretical) generalization of this approach to other problems [22]. Experimental evaluation of this approach for other problems such as Multiway Cut and Odd Cycle Transversal is a topic for future work.

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