Exploiting Database Management Systems and Treewidth for Counting*

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Abstract Bounded treewidth is one of the most cited combinatorial invariants, which was applied in the literature for solving several counting problems efficiently. A canonical counting problem is #Sat, which asks to count the satisfying assignments of a Boolean formula. Recent work shows that benchmarking instances for #Sat often have reasonably small treewidth. This paper deals with counting problems for instances of small treewidth. We introduce a general framework to solve counting questions based on state-of-the-art database management systems (DBMS). Our framework takes explicitly advantage of small treewidth by solving instances using dynamic programming (DP) on tree decompositions (TD). Therefore, we implement the concept of DP into a DBMS (PostgreSQL), since DP algorithms are already often given in terms of table manipulations in theory. This allows for elegant specifications of DP algorithms and the use of SQL to manipulate records and tables, which gives us a natural approach to bring DP algorithms into practice. To the best of our knowledge, we present the first approach to employ a DBMS for algorithms on TDs. A key advantage of our approach is that DBMS naturally allow to deal with huge tables with a limited amount of main memory (RAM), parallelization, as well as suspending computation.

 $\label{eq:Keywords: Dynamic Programming Parameterized Algorithmics Bounded Treewidth \cdot Database Management Systems \cdot SQL \cdot Relational Algebra \cdot Counting \cdot Model Counting$

1 Introduction

Counting solutions is a well-known task in mathematics, computer science, and other areas [9,19,28,43]. In combinatorics, for instance, one characterizes the number of solutions to problems by means of mathematical expressions, e.g., generating functions [20]. One particular counting problem, namely *model counting* (#SAT) asks to output the number of solutions of a given Boolean formula. Model counting and variants thereof have already been applied for solving a variety of real-world applications [9,11,22,48]. Such problems are typically considered rather

^{*} Our system dpdb is available under GPL3 license at github.com/hmarkus/dp_on_dbs.

hard, since #SAT is complete for the class #P [3,40], i.e., one can simulate any problem of the polynomial hierarchy with polynomially many calls [45] to a #SAT solver. However, recently it was shown that the better part [26] of the publicly available #SAT instances, techniques from parameterized complexity [27,21,38,13] might come to the rescue. In particular, after using regular preprocessors, e.g., pmc [33], in this area, more than 80% of these practically relevant instances (graph representations thereof) have reasonably small treewidth, where small treewidth is one of the most cited combinatorial invariants. To be more concrete, the measure treewidth is a structural parameter of graphs, which models the closeness of the graph to being a tree. This, gives rise to a general framework for counting problems that leverage treewidth. The general idea to develop such frameworks is indeed not new, since there are both, specialized solvers [10,26,29], as well as general systems like D-FLAT [5], Jatatosk [4], and sequoia [35], that exploit treewidth (to name a few). Some of these systems explicitly use dynamic programming (DP) to directly exploit treewidth by means of so-called tree decompositions (TDs), whereas others provide some kind of declarative layer to model the problem. However, for solving problems, most of the general systems [4,35] require a descriptive model of the problem, where an abstract view or for example certain logics are used internally, but problems in these systems are not directly described by means of dynamic programming algorithms. In this work, we solve (counting) problems by means of DP, where the algorithm is specified by giving essential parts of the DP algorithm in form of SQL SELECT queries. The whole DP algorithm is done by our system dpdb, which employs database management systems (DBMS) [47] for solving. This has not only the advantage of naturally describing and manipulating the tables that are obtained during DP, but also allows us to leverage from decades of database technology in form of the capability to deal with huge tables using limited amount of main memory (RAM), dedicated implementations of database joins, as well as query optimization and data-dependent execution plans.

Contribution. We implement a system dpdb for solving counting problems based on dynamic programming on tree decompositions, and present the following contributions. (i) Our system dpdb uses database management systems to efficiently handle table operations needed for performing dynamic programming efficiently. The system dpdb is written in Python and employs PostgreSQL as DBMS, but can work with other DBMSs easily. (ii) The architecture of dpdb allows to solve general problems of bounded treewidth that can be solved by by means of table operations (in form of relational algebra and SQL) on tree decompositions. As a result, dpdb is a generalized framework for dynamic programming on tree decompositions, where one only needs to specify the essential and problem-specific parts of dynamic programming in order to solve (counting) problems. (iii) Finally, we show how to solve the canonical problem #SAT with the help of dpdb, where it seems that the architecture of dpdb is particularly well-suited. Concretely, we

⁴ On October 17, 2019, Google Scholar shows 19,200 results on treewidth. Treewidth was also subject of recent competitions [16,17].

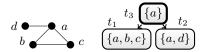


Figure 1: Graph G (left) with a TD \mathcal{T} of graph G (right).

compare the runtime of our system with state-of-the-art model counters, where we observe competitive behavior and promising indications for future work.

2 Preliminaries

We assume familiarity with terminology of graphs and trees. For details, we refer to the literature and standard textbooks [6,18].

Boolean Satisfiability. We define Boolean formulas and their evaluation in the usual way, cf., [28,30]. A literal is a Boolean variable x or its negation $\neg x$. A CNF formula φ is a set of clauses, interpreted as conjunction, which are sets of literals interpreted as disjunction. For a formula or clause X, we abbreviate by var(X) the variables that occur in X. An assignment of φ is a mapping $I : var(\varphi) \to \{0, 1\}$. The formula $\varphi(I)$ under assignment I is obtained by removing every clause c from φ that contains a literal set to 1 by I, and removing from every remaining clause of φ all literals set to 0 by I. An assignment I is satisfying if $\varphi(I) = \emptyset$. Problem #SAT asks to output the number of satisfying assignments of a formula.

Tree Decomposition and Treewidth. A tree decomposition (TD) of a given graph G is a pair $\mathcal{T}=(T,\chi)$ where T is a rooted tree and χ is a mapping which assigns to each node $t\in V(T)$ a set $\chi(t)\subseteq V(G)$, called bag, such that (i) $V(G)=\bigcup_{t\in V(T)}\chi(t)$ and $E(G)\subseteq \{\{u,v\}\mid t\in V(T),\{u,v\}\subseteq \chi(t)\};$ and (ii) for each $r,s,t\in V(T)$, such that s lies on the path from r to t, we have $\chi(r)\cap\chi(t)\subseteq\chi(s)$. We let width $(T):=\max_{t\in V(T)}|\chi(t)|-1$. The treewidth $\mathrm{tw}(G)$ of G is the minimum width (T) over all TDs T of G. For a node $t\in V(T)$, we say that type(t) is leaf if t has no children and $\chi(t)=\emptyset$; join if t has children t' and t'' with $t'\neq t''$ and $\chi(t)=\chi(t')=\chi(t'')$; intr ("introduce") if t has a single child $t',\chi(t')\subseteq\chi(t)$ and $|\chi(t)|=|\chi(t)|+1$; rem ("removal") if t has a single child $t',\chi(t')\supseteq\chi(t)$ and $|\chi(t')|=|\chi(t)|+1$. If for every node $t\in N$, type(t) $\in \{leaf, join, intr, rem\}$, then the TD is called nice.

Example 1. Figure 1 depicts a graph G and a TD \mathcal{T} of G of width 2. The treewidth of G is also 2 since G contains [31] a complete graph with 3 vertices.

Relational Algebra. We use relational algebra [12] for manipulation of relations, which forms the theoretical basis of its the well-known implementation database standard Structured Query Language (SQL) [47] on tables. An attribute a is of a certain finite domain dom(a). Then, a tuple r over set att(r) of attributes, is a set of pairs of the form (a, v) with $a \in \text{att}(r)$, $v \in \text{dom}(a)$ s.t. for each $a \in \text{att}(r)$, there is exactly one $v \in \text{dom}(a)$ with (a, v) $\in r$. A relation R is a finite set of tuples r

over set att(R) := att(r) of attributes. Given a relation R over att(R). Then, we let $dom(R) := \bigcup_{a \in att(R)} dom(a)$, and let relation R projected to $A \subseteq att(R)$ be given by $\Pi_A(R) := \{r_A \mid r \in R\}$, where $r_A := \{(a,v) \mid (a,v) \in r, a \in A\}$. This concept can be lifted to extended projection $\dot{\Pi}_{A,S}$, where we assume in addition to $A \subseteq \operatorname{att}(R)$, a set S of expressions of the form $a \leftarrow f$, such that $a \in \operatorname{att}(R) \setminus A$, and f is an arithmetic function that takes a tuple $r \in R$, such that there is at most one expression in S for each $a \in \text{att}(R) \setminus A$. Formally, we define $\dot{\Pi}_{A,S}(R) :=$ $\{r_A \cup r^S \mid r \in R\}$ with $r^S := \{(a, f(r)) \mid a \in \operatorname{att}(r), (a \leftarrow f) \in S\}$. Later, we use aggregation by grouping ${}_{A}G_{(a \leftarrow q)}$, where we assume $A \subseteq \operatorname{att}(R), a \in \operatorname{att}(R) \setminus A$ and a so-called aggregate function g, which takes a relation $R' \subseteq R$ and returns a value of domain dom(a). Therefore, we let ${}_AG_{(a\leftarrow g)}(R) := \{r \cup \{(a,g(R[r]))\} \mid r \in \{r \in A, r \in B\}\}$ $\Pi_A(R)$, where $R[r] := \{r' \mid r' \in R, r \subseteq r'\}$. We define renaming of R given set A of attributes, and a bijective mapping $m: \operatorname{att}(R) \to A$ s.t. $\operatorname{dom}(a) = \operatorname{dom}(m(a))$ for $a \in \text{att}(R)$, by $\rho_m(R) := \{(m(a), v) \mid (a, v) \in R\}$. Selection of rows in R according to a given Boolean formula φ with equality⁵ is defined by $\sigma_{\varphi}(R) :=$ $\{r \mid r \in R, \varphi(r^E) = \emptyset\}$, where r^E is a truth assignment over $var(\varphi)$ such that for each $v, v', v'' \in \text{dom}(R) \cup \text{att}(R)$ (1) $r^E(v \approx v') = 1$ if $(v, v') \in r$, (2) $r^E(v \approx v) = 1$, (3) $r^E(v \approx v') = r^E(v' \approx v)$, and (4) if $r^E(v \approx v') = 1$, and $r^E(v' \approx v'') = 1$, then $r^{E}(v \approx v'') = 1$. Given a relation R' with $\operatorname{att}(R') \cap \operatorname{att}(R) = \emptyset$. Then, we refer to the cross-join by $R \times R' := \{r \cup r' \mid r \in R, r' \in R'\}$. Further, wet let θ -join correspond to $R \bowtie_{\varphi} R' := \sigma_{\varphi}(R \times R')$.

3 Towards Relational Algebra for Dynamic Programming

A solver based on dynamic programming (DP) evaluates the input \mathcal{I} in parts along a given TD of a graph representation G of the input. Thereby, for each node t of the TD, intermediate results are stored in a table τ_t . This is achieved by running a so-called table algorithm A, which is designed for a certain graph representation, and stores in τ_t results of problem parts of \mathcal{I} , thereby considering tables $\tau_{t'}$ for child nodes t' of t. The DP approach works for many problems \mathcal{P} as follows.

- 1. Construct a graph representation G of the given input instance \mathcal{I} .
- 2. Heuristically compute a tree decomposition $\mathcal{T} = (T, \chi)$ of G.
- 3. Traverse the nodes in V(T) in post-order, i.e., perform a bottom-up traversal of T. At every node t during post-order traversal, execute a table algorithm A that takes as input t, bag $\chi(t)$, a local problem $\mathcal{P}(t,\mathcal{I}) = \mathcal{I}_t$ depending on \mathcal{P} , as well as previously computed child tables of t and stores the result in τ_t .
- 4. Interpret table τ_n for the root n of T in order to output the solution of \mathcal{I} .

For solving problem $\mathcal{P} = \# \mathrm{SAT}$, we need the following graph representation. The primal graph G_{φ} [41] of a formula φ has as vertices its variables, where two variables are joined by an edge if they occur together in a clause of φ .

⁵ We allow for φ to contain expressions $v \approx v'$ as variables for $v, v' \in \text{dom}(R) \cup \text{att}(R)$, and we abbreviate for $v \in \text{att}(R)$ with $\text{dom}(v) = \{0, 1\}$, $v \approx 1$ by v and $v \approx 0$ by $\neg v$.

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Listing 2: Table algorithm S(t, \chi(t), \varphi_t, \langle \tau_1, \dots, \tau_\ell \rangle) for \#SAT [41] using nice TD.
     In: Node t, bag \chi(t), clauses \varphi_t, sequence \langle \tau_1, \dots \tau_\ell \rangle of child tables. Out: Table \tau_t.
1 if type(t) = leaf then \tau_t := \{\langle \emptyset, 1 \rangle \}
2 else if type(t) = intr, and a \in \chi(t) is introduced then
           \tau_t := \{\langle \boldsymbol{J}, c \rangle
                                                     |\langle I, c \rangle \in \tau_1, J \in \{I_{a \mapsto 0}^+, I_{a \mapsto 1}^+\}, \varphi_t(J) = \emptyset\}
4 else if type(t) = rem, and a \notin \chi(t) is removed then
           \tau_t := \{ \langle I_a^-, \varSigma_{\langle J, c \rangle \in \tau_1 : I_a^- = J_a^-} c \rangle
                                                                                  |\langle I, \cdot \rangle \in \tau_1 \}
6 else if type(t) = join then
           \tau_t := \{ \langle I, c_1 \cdot c_2 \rangle
                                                      |\langle \mathbf{I}, c_1 \rangle \in \tau_1, \langle \mathbf{I}, c_2 \rangle \in \tau_2 \}
S_e^- := S \setminus \{e \mapsto 0, e \mapsto 1\}, S_s^+ := S \cup \{s\}.
                                                                                                                   \{a\}
             \tau_5
                                                                                              \{a,b\}
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Figure 2: Selected tables obtained by DP on \mathcal{T}' for φ of Example 2 using algorithm S.

2

3

 $1\}, 1\rangle$

 $\underbrace{\{a,b,c\}}_{\{a,c\}}t$

 $\{a\}$

 $\langle {m i}, c_{1.i}
angle$

Given formula φ , a TD $\mathcal{T} = (T, \chi)$ of G_{φ} and a node t of T. Then, we let local problem $\#Sat(t, \varphi) = \varphi_t$ be $\varphi_t := \{c \mid c \in \varphi, var(c) \subseteq \chi(t)\}$, which are the clauses entirely covered by $\chi(t)$.

Table algorithm S as presented in Listing 2 shows all the cases that are needed to solve #SAT by means of DP of nice TDs. Each table τ_t consist of rows of the form $\langle I,c\rangle$, where I is an assignment of φ_t and c is a counter. Nodes t with type(t) = leaf consist of the empty assignment and counter 1, cf., Line 1. For a node t with introduced variable $a \in \chi(t)$, we guess in Line 3 for each assignment β of the child table, whether a is set to true or to false, and ensure that φ_t is satisfied. When an atom a is removed in node t, we project assignments of child tables to $\chi(t)$, cf., Line 5, and counters of the same assignments are summed up. For join nodes t, counters of common assignments in the child tables are multiplied as in Line 7.

Example 2. Consider formula $\varphi := \{\{\neg a, b, c\}, \{a, \neg b, \neg c\}, \{a, d\}, \{a, \neg d\}\}\}$. Satisfying assignments of formula φ are, e.g., $\{a \mapsto 1, b \mapsto 1, c \mapsto 0, d \mapsto 0\}$, $\{a \mapsto 1, b \mapsto 0, c \mapsto 1, d \mapsto 0\}$ or $\{a \mapsto 1, b \mapsto 1, c \mapsto 1, d \mapsto 1\}$. In total, there are 6 satisfying assignments of φ . Observe that graph G of Figure 1 actually depicts the primal graph G_{φ} of φ . Intuitively, \mathcal{T} of Figure 1 allows to evaluate formula φ in parts. Figure 2 illustrates a nice TD $\mathcal{T}' = (\cdot, \chi)$ of the primal graph G_{φ} and tables $\tau_1, \ldots, \tau_{12}$ that are obtained during the execution of S on nodes t_1, \ldots, t_{12} .

Listing 3: Alternative table algorithm $S_{RAlg}(t, \chi(t), \varphi_t, \langle \tau_1, \dots, \tau_\ell \rangle)$ for #SAT.

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In: Node t, bag \chi(t), clauses \varphi_t, sequence \langle \tau_1, \dots \tau_\ell \rangle of child tables. Out: Table \tau_t.

1 if \operatorname{type}(t) = \operatorname{leaf} then \tau_t := \{\{(\operatorname{cnt}, 1)\}\}

2 else if \operatorname{type}(t) = \operatorname{intr}, and a \in \chi(t) is introduced then

3 | \tau_t := \tau_1 \bowtie_{\varphi_t} \{\{(\llbracket a \rrbracket, 0)\}, \{(\llbracket a \rrbracket, 1)\}\}

4 else if \operatorname{type}(t) = \operatorname{rem}, and a \notin \chi(t) is removed then

5 | \tau_t := \chi_{(t)} G_{\operatorname{cnt} \leftarrow \operatorname{SUM}(\operatorname{cnt})}(\Pi_{\operatorname{att}(\tau_1) \setminus \{\llbracket a \rrbracket\}} \tau_1)

6 else if \operatorname{type}(t) = \operatorname{join} then

7 | \tau_t := \dot{\Pi}_{\chi(t), \{\operatorname{cnt} \leftarrow \operatorname{cnt} \cdot \operatorname{cnt}'\}}(\tau_1 \bowtie_{\Lambda_{a \in \chi(t)}} \llbracket a \rrbracket \approx_{\llbracket a \rrbracket'} \rho \bigcup_{a \in \operatorname{att}(\tau_2)} \llbracket a \rrbracket \mapsto_{\llbracket a \rrbracket'} \tau_2)
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We assume that each row in a table τ_t is identified by a number, i.e., row i corresponds to $u_{t,i} = \langle I_{t,i}, c_{t,i} \rangle$.

Table $\tau_1 = \{ \langle \emptyset, 1 \rangle \}$ has type $(t_1) = leaf$. Since type $(t_2) = intr$, we construct table τ_2 from τ_1 by taking $I_{1.i} \cup \{a \mapsto 0\}$ and $I_{1.i} \cup \{a \mapsto 1\}$ for each $\langle I_{1.i}, c_{1.i} \rangle \in \tau_1$. Then, t_3 introduces c and t_4 introduces b. $\varphi_{t_1} = \varphi_{t_2} = \varphi_{t_3} = \emptyset$, but since $\chi(t_4) \subseteq \text{var}(c_1)$ we have $\varphi_{t_4} = \{c_1, c_2\}$ for t_4 . In consequence, for each $I_{4,i}$ of table τ_4 , we have $\{c_1, c_2\}(I_{4.i}) = \emptyset$ since S enforces satisfiability of φ_t in node t. Since type $(t_5) = rem$, we remove variable c from all elements in τ_4 and sum up counters accordingly to construct τ_5 . Note that we have already seen all rules where c occurs and hence c can no longer affect interpretations during the remaining traversal. We similarly create $\tau_6 = \{\langle \{a \mapsto 0\}, 3\rangle, \langle \{a \mapsto 1\}, 3\rangle \}$ and $\tau_{10} = \{\langle \{a \mapsto 1\}, 2 \rangle \}$. Since type $(t_{11}) = join$, we build table τ_{11} by taking the intersection of τ_6 and τ_{10} . Intuitively, this combines assignments agreeing on a, where counters are multiplied accordingly. By definition (primal graph and TDs), for every $c \in \varphi$, variables var(c) occur together in at least one common bag. Hence, since $\tau_{12} = \{\langle \emptyset, 6 \rangle\}$, we can reconstruct for example model $\{a \mapsto$ $1, b \mapsto 1, c \mapsto 0, d \mapsto 1$ = $I_{11.1} \cup I_{5.4} \cup I_{9.2}$ of φ using highlighted (yellow) rows in Figure 2. On the other hand, if φ was unsatisfiable, τ_{12} would be empty (\emptyset) .

Alternative: Relational Algebra. Instead of using set theory to describe how tables are obtained during dynamic programming are performed, one could alternatively use relational algebra. There, tables τ_t for each TD node t are pictured as relations, where τ_t distinguishes a unique column (attribute) $[\![x]\!]$ for each $x \in \chi(t)$. Further, there might be additional attributes required depending on the problem at hand, e.g., we need an attribute cnt for counting in #SAT, or an attribute for modeling costs or weights in case of optimization problems. Listing 3 presents a table algorithm for problem #SAT that is equivalent to Listing 2, but relies on relational algebra only for computing tables. This step from set notation to relational algebra is driven by the observation that in these table algorithms one can identify recurring patterns, and one mainly has to adjust problem-specific parts of it (highlighted by coloring in Listing 2). In particular, one typically derives for nodes t with type(t) = t0 Listing 3. Then, whenever an atom t0 is introduced, such algorithms often use t0-joins with a fresh initial table for the introduced variable t1 that

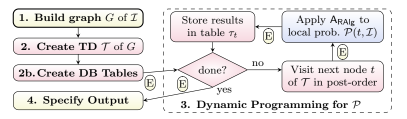


Figure 3: Architecture of Dynamic Programming with Databases. Steps highlighted in red are provided by the system depending on specification of yellow and blue parts, which is given by the user for specific problems \mathcal{P} . The yellow "E"s represent events that can be intercepted and handled by the user. The blue part concentrates on table algorithm A_{RAlg} , where the user specifies how SQL code is generated in a modular way.

represent potential values a can have. In Line 3 the selection of the θ -join is performed by ensuring φ_t , corresponding to the local problem of #SAT. Further, for nodes t with type(t) = rem, these table algorithms oftentimes need projection. In case of Listing 3, Line 5 also needs grouping in order to maintain the counter, as several rows of τ_1 might collapse in τ_t . Finally, for a node t with type(t) = join, in Line 7 we use again θ -joins (and extended projection for maintaining counters), which allows us later to leverage database technology of the last decades.

4 Dynamic Programming on TDs using Databases & SQL

In this section, we present a general architecture to model table algorithms by means of database management systems. The architecture is influenced by the DP approach of the previous section and works as depicted in Figure 3, where the steps highlighted in yellow and blue need to be specified depending on the problem \mathcal{P} . Steps outside Step 3 are mainly setup tasks, the yellow "E"s indicate events that might be needed to solve more complex problems on the polynomial hierarchy. For example, one could create and drop auxiliary sub-tables for each node during Step 3 within such events. Observe that after the generation of a TD $\mathcal{T}=(T,\chi)$, Step 2b automatically creates tables τ_t for each node t of T, where the corresponding table schema of τ_t is specified in the blue part, i.e., within A_{RAlg} . The default schema of such a table τ_t that is assumed in this section foresees one column for each element of the bag $\chi(t)$, where additional columns such as counters or costs can be added.

Actually, the core of this architecture is focused on the table algorithm A_{RAlg} executed for each node t of TD $\mathcal{T} = (T, \chi)$. Besides the definition of table schemes, the blue part concerns specification of the table algorithm by means of a procedural generator template that describes how to dynamically obtain SQL code for each node t thereby oftentimes depending on $\chi(t)$. This generated SQL code is then used internally for manipulation of tables τ_t during the tree decomposition traversal in Step 3 of dynamic programming. Listing 4 presents a general template, where parts of table algorithms for problems that are typically

⁶ Recall that SQL is a specific implementation standard (set) of relational algebra.

Listing 4: Template of $A_{\mathsf{RAlg}}(t, \chi(t), \mathcal{I}_t, \langle \tau_1, \dots, \tau_\ell \rangle)$ of Figure 3 for problem \mathcal{P} .

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In: Node t, bag \chi(t), instance \mathcal{I}_t, sequence \langle \tau_1, \dots \tau_\ell \rangle of child tables. Out: Table \tau_t.

1 if \operatorname{type}(t) = \operatorname{leaf} \operatorname{then} \ \tau_t := \#\varepsilon \operatorname{\mathsf{Tab}} \#

2 else if \operatorname{type}(t) = \operatorname{intr}, \ \operatorname{and} \ a \in \chi(t) \ is \ \operatorname{introduced} \ \operatorname{then}

3 \mid \tau_t := \dot{I}_{\chi(t), \#\operatorname{extProj} \#}(\tau_1 \bowtie_{\#\operatorname{localProbFilter} \#} \#\operatorname{intr} \operatorname{\mathsf{Tab}} \#)

4 else if \operatorname{type}(t) = \operatorname{rem}, \ \operatorname{and} \ a \notin \chi(t) \ is \ \operatorname{removed} \ \operatorname{then}

5 \mid \tau_t := \chi_{(t)} G_{\#\operatorname{aggrExp} \#}(\Pi_{\operatorname{att}(\tau_1) \setminus \{\llbracket a \rrbracket\}} \tau_1)

6 else if \operatorname{type}(t) = \operatorname{join} \ \operatorname{then}

7 \mid \tau_t := \dot{I}_{\chi(t), \#\operatorname{extProj} \#}(\tau_1 \bowtie_{\Lambda_a \in \chi(t)} \llbracket a \rrbracket \approx \llbracket a \rrbracket' \rho \bigcup_{a \in \operatorname{att}(\tau_2)} \llbracket a \rrbracket \mapsto \llbracket a \rrbracket' \} \tau_2)
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problem-specific are replaced by colored placeholders of the form #placeHolder#, cf., Listing 3. Observe that Line 3 of Listing 4 uses extended projection as in Line 7. This is needed for some problems, where additional columns require updates, whenever vertices get introduced.

Note, however, that the whole architecture does not depend on certain normalization or forms of TDs, e.g., whether it is nice or not. Instead, a table algorithm of any TD is simply specified by handling problem-specific implementations of the placeholders of Listing 4, where the system following this architecture is responsible for interleaving and overlapping these cases within a node t. In fact, we discuss an implementation of a system according to this architecture next, where for efficiency it is crucial to implement non-nice TDs.

4.1 System dpdb: Dynamic Programming with Databases

We implemented the proposed architecture of the previous section in the prototypical dpdb system. The system is open-source⁷, written in Python 3 and uses PostgreSQL as DBMS. We are convinced though that one can easily replace PostgreSQL by any other state-of-the-art relational database that uses SQL. In the following, we discuss implementation specifics that are crucial for a performant system that is still extendable and flexible.

Computing TDs. TDs are computed mainly with the library htd version 1.2 with default settings [2], which finds TDs extremely quick also for interesting instances [26] due to heuristics. Note that dpdb directly supports the TD format of recent competitions [16,17], i.e., one could easily replace the TD library. It is crucial though to not enforce htd to compute nice TDs, as this would cause a lot of overhead later in dpdb for copying tables. However, in order to benefit from the implementation of θ -joins, query optimization and state-of-the-art database technology in general, we observed that it is crucial to limit the number of child nodes of every TD node. Then, especially when there are huge tables involved, θ -joins among child node tables cover at most a limited number of child node tables. In consequence, the query optimizer of the database system still has a chance to come up with meaningful execution plans depending on the contents

⁷ Our system dpdb is available under GPL3 license at github.com/hmarkus/dp_on_dbs.

of the table. Note that though one should consider θ -joins with more than just two tables, since such binary θ -joins already fix in which order these tables shall be combined, thereby again limiting the query optimizer. Apart from this trade-off, we tried to outsource the task of joining tables to the DBMS, since the performance of database systems highly depend on query optimization. The actual limit, which is a restriction from experience and practice only, highly depends on the DBMS that is used. For PostgreSQL, we set a limit of at most 5 child nodes for each node of the TD, i.e., each θ -join covers at most 5 child tables.

Towards non-nice TDs. Although this paper presents the algorithms for nice TDs (mainly due to simplicity), the system dpdb interleaves these cases as presented in Listing 4. Concretely, the system executes one query per table τ_t for each node t during the traversal of TD \mathcal{T} . This query consists of serveral parts and we briefly explain its parts from outside to inside. First of all, the inner-most part concerns the row candiates for τ_t consisting of the θ -join as in Line 7 of Listing 4, including parts of Line 3, namely cross-joins for each introduced variable, involving #intrTab# without the filtering on #localProbFilter#. Then, there are different configurations of dpdb concerning these row candidates. For debugging (see below) one could (1) actually materialize the result in a table, whereas for performance runs, one should use (2) common table expressions (CTEs or WITH-queries) or (3) sub-queries (nested queries), which both result in one nested SQL query per table τ_t . On top of these row candidates, projection⁸ and grouping involving #aggrExp# as in Line 5 of Listing 4, as well as selection according to #localProbFilter#, cf., Line 3, is specified. It turns out that PostgreSQL can do better with sub-queries, where the query optimizer oftentimes pushes selection and projection into the sub-query if needed, which is not the case for CTEs, as discussed in the PostgreSQL manual [1, Sec. 7.8.1]. On different DBMS or other vendors, e.g., Oracle, it might be better to use CTEs instead.

Example 3. Consider again Example 2 and Figure 1. If we use table algorithm S_{RAlg} with dpdb on formula φ of TD \mathcal{T} and Option (3): sub-queries, where the row candidates are expressed via a sub-queries. Then, for each node t_i of \mathcal{T} , we dpdb generates a view vi as well as a table τ_i containing in the end the content of vi. Observe that each view only has one column $\llbracket a \rrbracket$ for each variable a of φ since the truth assignment of the other variables are not needed later. Actually, in dpdb the additional columns are kept (empty, null) for readability. This keeps the tables compact, only τ_1 has two rows, τ_2 , and τ_3 have only one row. We obtain the following views.

CREATE VIEW v1 AS SELECT a, sum(cnt) AS cnt FROM

(WITH intrTab AS (SELECT true AS val UNION ALL SELECT false)

SELECT i1.val AS a, i2.val AS b, i3.val AS c, 1 AS cnt

FROM intrTab i1, intrTab i2, intrTab i3)

WHERE (NOT a OR b OR c) AND (a OR NOT b OR NOT c) GROUP BY a

⁸ Actually, dpdb keeps only columns relevant for the table of the parent node of t.

```
CREATE VIEW v2 AS SELECT a, sum(cnt) AS cnt FROM (WITH intrTab AS (SELECT true AS val UNION ALL SELECT false) SELECT i1.val AS a, i2.val AS d, 1 AS cnt FROM intrTab i1, intrTab i2) WHERE (a OR d) AND (a OR NOT d) GROUP BY a 

CREATE VIEW v3 AS SELECT a, sum(cnt) AS cnt FROM (SELECT \tau_1.a, \tau_1.cnt * \tau_2.cnt AS cnt FROM \tau_1, \tau_2 WHERE \tau_1.a = \tau_2.a) GROUP BY a
```

Parallelization. A further reason to not over-restrict the number of child nodes within the TD, lies in parallelization. In dpdb, we compute tables in parallel along the TD, where multiple tables can be computed at the same time, as long as the child tables are computed. Therefore, we tried to keep the number of child nodes in the TD as high as possible. In our system dpdb, we currently allow for at most 24 worker threads for table computations and 24 database connections at the same time (both pooled and configurable). On top of that we have 2 additional threads and database connections for job assignments to workers, as well as one dedicated watcher thread for clean-up and termination of connections, respectively.

Logging, Debugging and Extensions. Currently, we currently have two versions of the dpdb system implemented. One version aims for performance and the other one tries to achieve comprehensive logging and easy debugging of problem (instances), thereby increasing explainability. The former for instance does neither keep intermediate results nor create database tables in advance (Step 2b), as depicted in Figure 3, but creates tables according to an SQL SELECT statement. In the latter we keep all the intermediate results, we record database timestamps before and after certain nodes, provide statistics as, e.g., width, number of rows, etc. Further, since for each table τ_t , exactly one SQL statement is executed for filling this table, we also have a dedicated view of the SQL SELECT statement, whose result is then inserted in τ_t . Together with the power and flexibility of SQL queries, we observed that this really helps in finding errors in the table algorithm specifications.

Besides convient debugging, system dpdb immediately contains an extension for approximation. There, we restrict the table contents to a maximum number of rows. This allows for certain approximations on counting problems or optimization problems, where it is infeasible to compute the full tables. Further, dpdb foresees a dedicated randomization on these restricted number of rows such that if one repeats the computation with a different random seed, we obtain different approximate results.

Note that dpdb can be easily extended. Each problem can overwrite existing default behavior and dpdb also supports problem-specific argument parser for each problem individually. Out-of-the-box, we support the formats DIMACS sat and DIMACS graph [36] as well as the format for TDs used in recent competitions [16,17].

4.2 Table algorithms with dpdb for selected problems

The system dpdb allows for easy protyping of DP algorithms on TDs. This covers decision problems, counting problems as well as optimization problems. As a proof of concept, we present the relevant parts of table algorithm specification according to the template in Listing 4 for a selection of problems below⁹. To this end, we assume in this section a not necessarily nice TD $\mathcal{T} = (T, \chi)$ of the corresponding graph representation of our given instance \mathcal{I} . Further, for the following specifications of the table algorithm using the template A_{RAlg} in Listing 3, we assume any node t of T and its child nodes t_1, \ldots, t_ℓ .

Problem #SAT. Given instance formula $\mathcal{I} = \varphi$. Then, specific parts for #SAT for node t with $\varphi_t = \{\{l_{1,1}, \ldots, l_{1,k_1}\}, \ldots, \{l_{n,1}, \ldots, l_{n,k_n}\}\}$.

Observe that for the corresponding decision problem SAT, where the goal is to decide only the existence of a satisfying assignment for given formula φ , $\#\varepsilon \mathsf{Tab}\#$ returns the empty table and parts $\#\mathsf{aggrExp}\#$, $\#\mathsf{extProj}\#$ are just empty since there is no counter needed.

Problem #o-Col. For given input graph $\mathcal{I} = G = (V, E)$, a o-coloring is a mapping $\iota: V \to \{1, \ldots, o\}$ such that for each edge $\{u, v\} \in E$, we have $\iota(u) \neq \iota(v)$. Problem #o-Col asks to count the number of o-colorings of G. Local problem #o-Col(t, G) is defined by the graph $G_t := (V \cap \chi(t), E \cap [\chi(t) \times \chi(t)])$. Specific parts for #o-Col for node t with $E(G_t) = \{\{u_1, v_1\}, \ldots, \{u_n, v_n\}\}$.

Problem MinVC. Given input graph $\mathcal{I} = G = (V, E)$, a vertex cover is a set of vertices $C \subseteq V$ of G such that for each edge $\{u,v\} \in E$, we have $\{u,v\} \cap C \neq \emptyset$. Then, MinVC asks to find the minimum cardinality |C| among all vertex cover C, i.e., C is such that there is no vertex cover C' with |C'| < |C|. Local problem MinVC $(t,G) := G_t$ is defined as above. To this end, we use an additional column card for storing cardinalities.

Problem MINVC for node t with $E(G_t) = \{\{u_1, v_1\}, \dots, \{u_n, v_n\}\}$ and $\chi(t) = \{a_1, \dots, a_k\}$ can be specified as follows.

⁹ Implementation for problems #SAT as well as MINVC is readily available in dpdb.

Observe that #ExtProj# is a bit more involved on non-nice TDs, as, whenever the column for a vertex a is set o 1, i.e., vertex a is in the vertex cover, we have to consider a only with cost 1, also if a appears in several child node bags.

Note that concrete implementations could generate and apply parts of this specification, as for example in #localProbFilter# only edges involving newly introduced vertices need to be checked.

Similar to MINVC and #o-Col one can model several other (graph) problems. One could also think of counting the number of solutions of problem MINVC, where both a column for cardinalities and one for counting is used. There, in addition to grouping with GROUP BY in dpdb, we additionally could use the HAVING construct of SQL, where only rows are kept, whose column card is minimal.

5 Experiments

We conducted a series of experiments using publicly available benchmark sets for #SAT. Our tested benchmarks [25] are publicly available, and our results are also on github at github.com/hmarkus/dp_on_dbs/padl2020.

5.1 Setup

Measure & Resources. We mainly compare wall clock time and number of timeouts. In the time we include, if applicable, preprocessing time as well as decomposition time for computing a TD with a fixed random seed. For parallel solvers we allowed access to 24 physical cores on machines. We set a timeout of 900 seconds and limited available RAM to 14 GB per instance and solver.

Benchmark Instances. We considered a selection of overall 1494 instances from various publicly available benchmark sets #SAT consisting of fre/meel benchmarks¹⁰(1480 instances), and c2d benchmarks¹¹ (14 instances). However, we considered instances preprocessed by regular #SAT preprocessor pmc [33], similar to results of recent work on #SAT [26], where it was also shown that more than 80% of the #SAT instances have primal treewidth below 19 after preprocessing.

Benchmarked system dpdb. We used PostgreSQL 9.5 for our system dpdb, which was available on our benchmark described hardware below. However, we expect major performance increases if higher versions are used, which was not available on our benchmark machines. In particular, parallel queries, where a query is evaluated in parallel, were added and improved in every version greater than 9.6.

¹⁰ See: tinyurl.com/countingbenchmarks

¹¹ See: reasoning.cs.ucla.edu/c2d

Other benchmarked systems. In our experimental work, we present results for the most recent versions of publicly available #SAT solvers, namely, c2d 2.20 [14], d4 1.0 [34], DSHARP 1.0 [37], miniC2D 1.0.0 [39], cnf2eadt 1.0 [32], bdd_minisat_all 1.0.2 [46], and sdd 2.0 [15] (based on knowledge compilation techniques). We also considered rather recent approximate solvers ApproxMC2, ApproxMC3 [8] and sts 1.0 [23], as well as CDCL-based solvers Cachet 1.21 [42], sharpCDCL¹², and sharpSAT 13.02 [44]. Finally, we also included multi-core solvers gpusat 1.0 and gpusat 2.0 [26], as well as countAntom 1.0 [7] on 12 physical CPU cores, which performed better than on 24 cores. We considered also additional solvers, e.g., d-DNNF reasoner 0.4.180625 on top of d4 as underlying knowledge compiler, where detailed results can be found online. All experiments were conducted with default solver options.

Benchmark Hardware. Almost all solvers were executed on a cluster of 12 nodes. Each node is equipped with two Intel Xeon E5-2650 CPUs consisting of 12 physical cores each at 2.2 GHz clock speed and 256 GB RAM. The results were gathered on Ubuntu 16.04.1 LTS machines with disabled hyperthreading on kernel 4.4.0-139. As we also took into account solvers using a GPU, for gpusat1 and gpusat2 we used a machine equipped with a consumer GPU: Intel Core i3-3245 CPU operating at 3.4 GHz, 16 GB RAM, and one Sapphire Pulse ITX Radeon RX 570 GPU running at 1.24 GHz with 32 compute units, 2048 shader units, and 4GB VRAM using driver amdgpu-pro-18.30-641594 and OpenCL 1.2. The system operated on Ubuntu 18.04.1 LTS with kernel 4.15.0-34.

5.2 Results

write

Figure 4 illustrates the top five sequential solvers, and all parallel counting solvers with preprocessor pmc in a cactus-like plot. Table 1 presents detailed runtime results for #SAT with preprocessor pmc, respectively. Since the solver sts produced results that varied from the correct result on average more than the value of the correct result, we excluded it from the presented results. If we disallow preprocessing, gpusat2 and gpusat1 perform only slightly better in the overall standing of the solvers. But gpusat2 solves 42 instances more and requires about 10 hours less of wallclock time. Further, we can observe, that the variant gpusat2(A+B) performs particular well, mainly since for instances below width 30, the BST compression seems relatively expensive compared to the array data structure. Interestingly, when considering the results on preprocessing in Table 1 (top, mid) and Figure 4 we observe that the architectural improvements pay off quite well. gpusat2 can solve the vast majority of the instances and ranks second place.

¹² See: tools.computational-logic.org

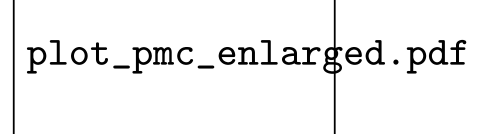


Figure 4: Runtime for the top 5 sequential and all parallel solvers over all the #SAT instances with pmc preprocessor. The x-axis refers to the number of instances and the y-axis depicts the runtime sorted in ascending order for each solver individually.

6 Final Discussion & Conclusions

We presented a generic system dpdb for explicitly exploiting treewidth by means of dynamic programming on databases. The idea of dpdb is to use database management systems (DBMS) for table manipulation, which makes it (1) easy and elegant to perform rapid prototyping for problems, and (2) allows to leverage from decades of database theory and database system tuning. It turned out that all the cases that occur in dynamic programming can be handled quite elegantly with plain SQL queries. Our system dpdb can be used for both decision and counting problems, thereby also considering optimization. We see our system particularly well-suited for counting problems, especially, since it was shown that for model counting (#SAT) instances of practical relevance typically have small treewidth [26]. In consequence, we carried out preliminary experiments on publicly available instances for #SAT, where we see competitive behavior compared to most recent solvers.

Future Work. Our results give rise to several research questions. First of all, we want to push towards PostgreSQL 11, but at the same time also consider other vendors and systems, e.g., Oracle. In particular, the behavior of different systems might change, when we use different strategies on how to write and evaluate our SQL queries, e.g., sub-queries vs. common table expressions. Currently, we do

	solver	0-20	21-30	31-40	41-50	51-60	>60	best	unique	$ \sum $	time[h]
processed by p	miniC2D	1193	29	10	2	1	7	13	0	1242	68.77
	gpusat2	1196	32	1	0	0	0	250	8	1229	71.27
	d4	1163	20	10	2	4	28	52	1	1227	76.86
	gpusat2(A+B)	1187	18	1	0	0	0	120	7	1206	74.56
	countAntom 12	1141	18	10	5	4	13	101	0	1191	84.39
	c2d	1124	31	10	3	3	10	20	0	1181	84.41
	sharpSAT	1029	16	10	2	4	30	253	1	1091	106.88
	gpusat1	1020	16	0	0	0	0	106	7	1036	114.86
	sdd	1014	4	7	1	0	2	0	0	1028	124.23
	solver	0-20	21-30	31-40	41-50	51-60	>60	best	unique		time[h]

Table 1: Number of #SAT instances, grouped by treewidth upper bound intervals, solved by sum of the top five sequential and all parallel counting solvers with preprocessor pmc. time[h] is the cumulated wall clock time in hours, where unsolved instances are counted as 900 seconds.

not create or use any indices, as preliminary tests showed that meaningful B^*tree indices are hard to create and oftentimes cost too much time to create. Further, the exploration of bitmap indices, as available in Oracle enterprise DBMS would be worth trying in our case (and for #SAT), since one can efficiently combine database columns by using extremely efficient bit operations.

It might be worth to rigorously test and explore our extensions on limiting the number of rows per table for *approximating* #SAT or other counting problems, cf., [9,22]. An other interesting research direction is to study whether efficient data representation techniques on DBMS can be combined with dynamic programming in order to lift our solver to quantified Boolean formulas. Finally, we are also interested in extending this work to projected model counting [24].

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