

CACHE PERFORMANCE OF PORTFOLIO-APPROACH-BASED PARALLEL SAT SOLVERS

por

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Abstract

A SAT solver is a computer program designed to solve instances of the SAT problem. As it is known, the SAT problem is NP-complete so there is no guarantee that any algorithm will solve a given instance of the SAT problem in a reasonable amount of time. There is a wide range of SAT solvers with different solving strategies which lead to very different implementations. We can compare them by having a benchmark set of SAT problems and count how many a determined SAT solver can solve within a limited time frame. The more problems it manages to solves, the better the SAT solver.

Because SAT solver programs run on a computer, their performance is also bound to the architecture of computers. This is the reason why SAT solvers' designs take into account modern computer features such as memory hierarchy and multicore architecture. To take advantage of multicore architectures, SAT solver developers are now designing parallel SAT solvers, which can run multiple processes in parallel to help speed up the total solving time of a problem. One of the most successful design approaches of parallel SAT solver is known as the portfolio-approach. The main idea is the fact that different SAT solvers will perform differently for different SAT problems. Some will perform better for some problems and others will perform better for another set of problems. So what a portfolio-approach parallel SAT solver will do is run different SAT solvers in parallel on the same problem, and wait for one of them to solve it. Also, all SAT solvers running in parallel can share information between them, because they are solving the same problem and thus might find useful information about the problem that could help the other solvers.

One of the most novel approaches to share information between solvers of a portfolio-approach parallel SAT solver is sharing it physically, which means that all solvers running in parallel will have access to the same memory locations to share data. In this work we made an empirical study of such approach and concluded that the current implementations of parallel SAT solvers which share information physically do not add any considerable advantage in performance. Moreover, only a

limited amount of physical information sharing has shown to bring small performance improvements.

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Chapter 1

Introduction

One of the most well-known problems in computer science is the satisfiability (SAT) problem. The SAT problem consists in determining whether a logical formula can be evaluated to true or not. For example, the logical formula $(a \vee b \vee c) \wedge (\neg a \vee \neg d) \wedge (c \vee d)$ can be satisfied with the variable assignments $a = \text{true}$, $b = \text{true}$, $c = \text{true}$ and $d = \text{false}$. So this formula is satisfiable since there is a variable assignment that satisfies the formula. SAT was the first problem proven to be NP-complete [4], proof that derived the Cook-Levin theorem. One year later, in 1972, Karp proved that many common combinatorial problems can be reduced in polynomial time to instances of the SAT problem [11], thus drawing even more attention to SAT problems by the scientific community. Since many combinatorial problems can be reduced to SAT, it is not strange to find many practical problems with useful applications (such as circuit design and automatic theorem proving) that could be solved if there was an efficient algorithm to solve the SAT problem. Unfortunately, because of the NP-completeness of SAT, unless $P = NP$, no polynomial algorithm can be found, but instead, many researchers have improved the current SAT solving algorithms that remain being exponential. Over the years, SAT solvers have shown impressive improvement, the first complete algorithm, the Davis Putnam algorithm [19], was very limited and could only handle small problems. Today, modern SAT solvers can handle instances with millions of variables, making such solvers suitable even for industrial application. In the next chapter we will point out the main features that have improved SAT solvers significantly in the past.

In the last decade parallel computing has become increasingly popular. As CPU manufacturers have found difficult and expensive to keep increasing the clock speed of processors, they have instead turned to increase the number of cores each chip has. Unfortunately, if the algorithms are not thought to be run in parallel, more cores

will bring small improvements. This is the reason why there is a growing concern to parallelize algorithms so that they can take advantage of many-cores architectures of today's computers. In SAT solving it is no different. The annual SAT competition¹, an event to determine which is the fastest SAT solver, has two main categories: sequential SAT solvers and parallel SAT solvers. In the last years parallel SAT solvers have outperformed sequential solvers in total wall clock time, so the interest in parallel solvers has grown with new designs and approaches explored for this kind of solvers. One of the most successful approaches to implement a parallel SAT solver is the *portfolio approach* with no *physical sharing* of information among cores. This approach basically runs different solvers in parallel, with each core keeping its own copy of the whole problem in memory, and wait for one of them to solve it up. No physical sharing refers to the fact that each core keeps its own copy of the problem's information, they do not access the same memory locations. This is a very simple and straight forward approach of parallelization, but we have also encountered one drawback to it: as we add more solvers to different cores of a single chip, the overall performance of the parallel solver decreases in around 20-40%. Preliminary experiments strongly suggest that this decrease in performance is caused by an increment in *cache misses*. A computer's processor keeps a small, but fast, memory hardware, called cache, from where it fetches data to make calculations. If a data requested is already in this cache, fetching the data will as a result be a fast process. On the other hand, if the data requested is not in the cache, but in main memory, fetching the data will take a longer time and thus make the whole process expensive, time wise. Because all cores in a single chip share the same cache², and because each thread holds a copy of the original problem in memory, the more threads we add, the bigger the amount of data we have to handle. Since there is only one cache shared among all cores, the amount of total accesses from the processor to main memory will increase, since now there is a bigger volume of data to handle.

We plan to research the impact on cache performance of physically sharing clauses. Physical sharing obviously involves a more complex implementation, because you need to ensure data integrity when modifying the same memory locations from different

¹www.satcompetition.org

²In fact there are different levels of cache, but we will refer to the last level cache which is shared.

threads, and also ensure that the correctness of the algorithms is kept. The mechanisms to accomplish these requirements add an overhead to the solver which is not present when threads do not share information. On the other hand, as we will demonstrate in our work, it is known that sharing information physically between threads usually improves cache performance, so there is a trade-off between both kind of approaches. The outcome of this trade-off is yet not clear and there are no serious studies about the cache performance of different parallel SAT solver implementations. So the goal of our work will be to study the impact of physically sharing clauses in cache performance of portfolio approach SAT solvers.

In the next chapter we will introduce the background work which is the base of our own. We will mostly discuss SAT solvers and modern computer's memory architecture. Chapter 3 will cover all experiments done, first with cache and then with SAT solvers. Finally, we will draw conclusions and discuss possible future work.

Chapter 2

Background and Related Work

For readers which are nor familiar with SAT solving, we need to explain some concepts, algorithms and works on which our own work is based. This chapter will explain how modern SAT solvers have become what they are today.

2.1 SAT problem

Given a set of boolean variables Σ , a literal L is either a variable or the negation of a variable in Σ , and a *clause* is a disjunction of literals over distinct variables¹ of Σ . A propositional sentence is in *conjunctive normal form (CNF)* if it has the form $\alpha_1 \wedge \alpha_2 \wedge \dots \wedge \alpha_n$, where each α_i is a clause. The notation of sentences in CNF we will be using are sets. A clause $l_1 \vee l_2 \vee \dots \vee l_m$, where l_i is a literal, can be expressed as the set $\{l_1, l_2, \dots, l_m\}$. Furthermore, the CNF $\alpha_1 \wedge \alpha_2 \wedge \dots \wedge \alpha_n$ can be expressed as the set of clauses $\{\alpha_1, \alpha_2, \dots, \alpha_n\}$. Given a CNF Δ , the SAT problem is answering the question: Is there an assignment of values for variables in Σ , such that Δ evaluates to true? A formula is said to be inconsistent if no such assignment exists. On the contrary, a formula is said to be satisfiable, or valid, if there is an assignment such that it evaluates to true. In particular, a CNF Δ is valid if Δ is the empty set: $\Delta = \emptyset$. A CNF Δ will be inconsistent if it contains the empty set: $\emptyset \in \Delta$.

Example 2.1.1.

$$\Delta = \{\{A, B, \neg C\}, \{\neg A, D\}, \{B, C, D\}\}$$

if $A = \text{true}$, $B = \text{true}$, $C = \text{true}$ and $D = \text{true}$, then

$$\Delta = \emptyset$$

Because a satisfied clause becomes irrelevant in the formula and thus removed. \square

¹That all literals in a clause have to be over distinct variables is not standard.

2.2 Algorithms for solving the SAT problem

In this section we will discuss the main complete algorithms that solve the SAT problem, their evolution and how they have become the modern SAT solvers that are used today.

2.2.1 Resolution

This is one of the first and most simple algorithms used to solve a SAT problem. The *resolution inference rule* [20] is defined as follows. Let P be a Boolean variable, and suppose that Δ is a CNF which contains clauses C_i and C_j , where $P \in C_i$ and $\neg P \in C_j$. The resolution inference rule allows us to derive the clause $(C_i - \{P\}) \cup (C_j - \{\neg P\})$, which is called a *resolvent* that is obtained by *resolving* C_i and C_j . A simple example is the CNF $\{\{A, \neg B\}, \{B, C\}\}$. Applying resolution to these two clauses would derive the clause $\{A, C\}$, which would be called a *B-resolvent*. Resolution is incomplete in the sense that it is not guaranteed to derive every clause that is implied by the CNF, but it is *refutation complete* on CNFs. It is guaranteed that resolution will derive the empty clause if the given CNF is unsatisfiable.

Unit resolution is an important special case of resolution. It is a resolution strategy which requires that at least one of the resolved clauses has only one literal. Such clause is called a *unit clause*. The importance of unit resolution does not rely on its completeness (it is actually not refutation complete, as resolution is), but one can apply all possible unit resolution steps in time linear to the size of a given CNF. Its efficiency makes it a key technique employed by modern solvers.

2.2.2 The depth-first search algorithm

One way to picture the search for a possible assignment of variables that satisfies the CNF formula is to use a tree. For example, given $\Sigma = \{A, B, C\}$ and $\Delta = \{\{\neg A, B\}, \{\neg B, C\}\}$, Figure 2.1 shows the search tree for this CNF. Each node n_i of the tree represents a variable, for each level we have a different variable. The branches are the different truth values the variable can be assigned: a **t** for true and an **f** for false. Each w_i represents a possible truth assignment of all the variables

in Σ . Note that w_1, w_5, w_7 and w_8 are all assignments that satisfy the CNF, while w_2, w_3, w_4 and w_6 do not. We can observe that the leaves of this tree are in one-to-one correspondence with all the possible true assignments of variables involved, so testing satisfiability can be viewed as searching for a leaf node that satisfies the CNF. Another important observation is that the depth of the tree is equal to the number of boolean variables, so performing a depth-first-search would be best to explore the tree.

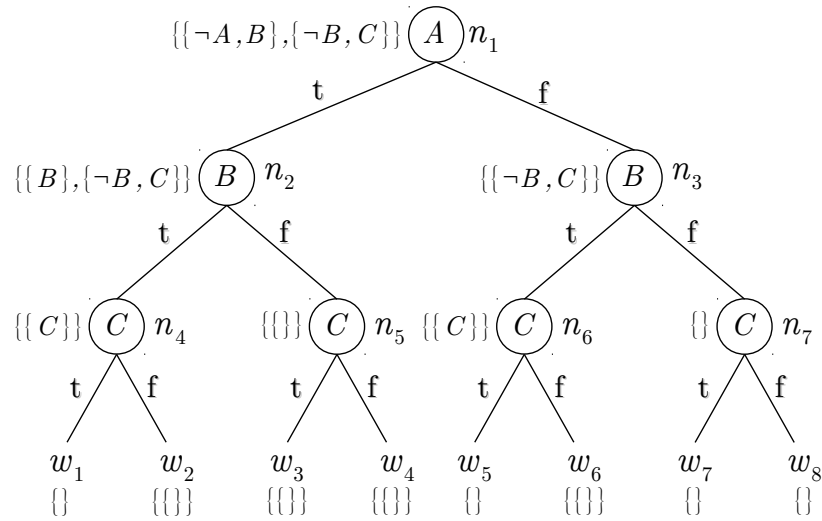


Figure 2.1: A search tree for the CNF $\Delta = \{\{\neg A, B\}, \{\neg B, C\}\}$.

The depth-first search algorithm also makes use of *conditioning*. Conditioning a CNF Δ on a literal L consists on replacing every occurrence of L by the constant **true**, replacing $\neg L$ with **false**, and simplifying accordingly. The result of conditioning Δ on L will be denoted by $\Delta|L$ and can be defined as follows:

$$\Delta|L = \{\alpha - \{\neg L\} | \alpha \in \Delta, L \notin \alpha\}$$

This means that the new set of clauses $\Delta|L$ will be all the clauses in Δ that do not contain L , but with the literal $\neg L$ removed. The clauses that contain L are removed because they are now satisfied, since we made L **true**. $\neg L$ is removed from the remaining clauses because it was set to **false** and no longer has any effect.

The definition of conditioning can be extended to multiple literals. For example,

the CNF:

$$\Delta = \{\{A, B, \neg C\}, \{\neg A, D\}, \{B, C, D\}\}$$

can be conditioned as $\Delta|CA\neg D = \{\emptyset\}$ (an inconsistent CNF). Moreover, $\Delta|\neg CD = \emptyset$ (a valid CNF). Intuitively, conditioning a CNF is just to assume that a certain variable is assigned a value and then simplify the clauses that include said variable accordingly. Algorithm 1 performs a depth-first-search using the conditioning operator on CNFs to remove clauses or reduce their size.

Algorithm 1: Depth-first-search algorithm. Variables are named P_1, P_2, \dots . Initial depth is 0.

```

1 DEPTH-FIRST-SEARCH(CNF  $\Delta$ , depth  $d$ ):
2 if  $\Delta = \{\}$  then
3   return  $\{\}$ .
4 else if  $\{\} \in \Delta$  then
5   return UNSATISFIABLE
6 else if  $L = \text{DEPTH-FIRST-SEARCH}(\Delta|P_{d+1}, d+1) \neq \text{UNSATISFIABLE}$  then
7   return  $L \cup \{P_{d+1}\}$ 
8 else if  $L = \text{DEPTH-FIRST-SEARCH}(\Delta|\neg P_{d+1}, d+1) \neq \text{UNSATISFIABLE}$  then
9   return  $L \cup \{\neg P_{d+1}\}$ 
10 else
11   return UNSATISFIABLE

```

Example 2.2.1. Consider the CNF of Figure 2.1 and the node labelled n_5 . At this node, Algorithm 1 will condition Δ on literals $A, \neg B$, which according to the definition will lead to:

$$\Delta|A, \neg B = \{\{\}\}.$$

and would be the same as writing:

$$\{\{\text{false}, \text{false}\}, \{\text{true}, C\}\}$$

The algorithm will detect that there is a contradiction at this internal node, hence concluding that neither w_3 or w_4 are models of Δ , without having to visit them explicitly. The algorithm can also detect success at internal nodes, implying that all assignments from that particular node are models of the CNF. Node n_7 , for example,

is already a valid CNF, so we do not have to check w_7 and w_8 . \square

2.2.3 The Davis-Putnam-Logemann-Loveland (DPLL) algorithm.

In this section we will discuss the DPLL [5] algorithm, which performs a systematic search in the space of truth assignments. The importance of the DPLL algorithm is that it is the foundation of modern SAT solvers.

One of the downsides of Algorithm 1 is that we cannot perform any kind of pruning to the search tree. We will have to systematically visit all nodes of the tree until we find a solution or until we have visited them all. The *unit resolution* (also called *unit propagation*) technique allows us to prune the search tree at each decision level. The unit resolution technique is very simple: Before we check for success or contradictions, we first collect all unit clauses. Then we assume that each variable which makes a clause unit is set to satisfy its clause. If new clauses become unit we keep repeating this process until we reach success or a contradiction.

Example 2.2.2. Consider the tree of Figure 2.2 and the CNF:

$$\Delta = \{\{\neg A, B\}, \{\neg B, \neg C\}, \{C, \neg D\}\}.$$

Consider also the node n_2 , which results from setting variable A to **true**, and its corresponding CNF:

$$\Delta|A = \{\{B\}, \{\neg B, \neg C\}, \{C, \neg D\}\}.$$

Algorithm 1 cannot declare early success or early failure, because the CNF is neither empty, nor contains the empty clause, reason why it keeps searching below Level 1 as shown in Figure 2.2. However, by using unit resolution, we can declare success and end the search at Level 1. Because the clause $\{B\}$ has become unit, we can assume that variable B is **true**. Now the clause $\{\neg B, \neg C\}$ is simplified to $\{\neg C\}$ (because $\neg B$ is **false** and cannot satisfy the clause) which is a unit clause. This will cause C to be **false** and clause $\{C, \neg D\}$ to be simplified to $\{\neg D\}$. This clause is also unit now so D must be assigned **false** and hence all clauses in this CNF have been satisfied through unit resolution. We can then declare success without having to go any further in

the search tree. Remember that the unit resolution process is very fast and does not require to make any decisions. \square

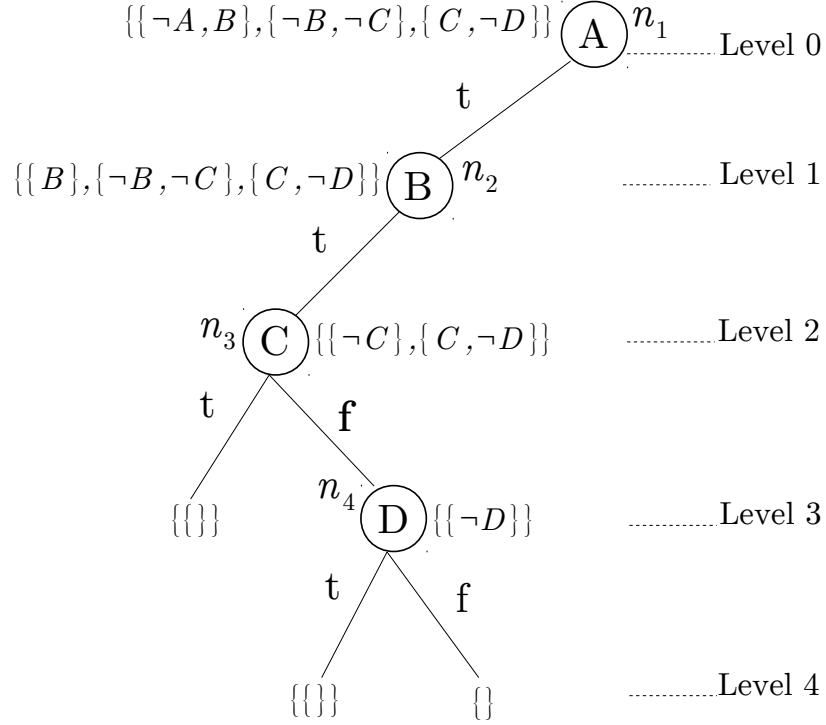


Figure 2.2: A search tree which only shows the nodes visited by the depth-first search algorithm. Note that this algorithm can declare success at level 4, after visiting the leafs of node n_4 . On the other hand, DPLL algorithm can declare success at node n_2 at level 1.

To incorporate the advantages of unit resolution into Algorithm 1, we will introduce a function **UNIT-RESOLUTION**, which applies to a CNF Δ and returns two results:

- **I**: a set of literals that are either present as unit clauses in Δ , or were derived from Δ by unit resolution.
- Γ : a new CNF which results from conditioning Δ on literals **I**.

Example 2.2.3. If the argument of **UNIT-RESOLUTION** was the CNF

$$\Delta = \{\{\neg A, \neg B\}, \{B, C\}, \{\neg C, D\}, \{A\}\},$$

then $\mathbf{I} = \{A, \neg B, C, D\}$ and $\Gamma = \{\}$. Moreover, if

$$\Delta = \{\{\neg A, \neg B\}, \{B, C\}, \{\neg C, D\}, \{C\}\},$$

then $\mathbf{I} = \{C, D\}$ and $\Gamma = \{\{\neg A, \neg B\}\}$. \square

The DPLL algorithm (Algorithm 2) is a refinement of Algorithm 1. The first change made is that it adds unit resolution in line 1. Also, it does not longer assume that variables are examined in a strict order as we go down the search tree and it does not longer assume that a variable is set to **true** and then to **false**. The choice of a literal L on line 7 can have a dramatic impact on the running time of DPLL. This is where inference comes into play when solving a SAT instance with a DPLL based algorithm, heuristics and random factors are commonly used at this point.

Note that DPLL algorithm will make sure that all nodes are discarded if no solution is found, either by visiting them explicitly or by pruning them. Furthermore, it will never visit the same node of the search tree more than once (it will not make the same decision sequence twice). Therefore, algorithm DPLL is sound and complete.

Algorithm 2: DPLL(CNF Δ): returns a set of literals or UNSATISFIABLE

```

1   $(\mathbf{I}, \Gamma) = \text{UNIT-RESOLUTION}(\Delta)$ 
2  if  $\Gamma = \{\}$  then
3    return  $\mathbf{I}$ .
4  else if  $\{\} \in \Gamma$  then
5    return UNSATISFIABLE
6  else
7    choose a literal  $L$  in  $\Gamma$ 
8    if  $\mathbf{L} = \text{DPLL}(\Gamma|L) \neq \text{UNSATISFIABLE}$  then
9      return  $\mathbf{L} \cup \mathbf{I} \cup \{L\}$ 
10   else if  $\mathbf{L} = \text{DPLL}(\Gamma|\neg L) \neq \text{UNSATISFIABLE}$  then
11     return  $\mathbf{L} \cup \mathbf{I} \cup \{\neg L\}$ 
12   else
13     return UNSATISFIABLE

```

2.2.4 Conflict-Driven Clause-Learning Solvers

There are several disadvantages that the DPLL algorithm still suffers from. The most important one has to do with the fact that the DPLL algorithm performs a *chronological backtracking*. In this section we will discuss this limitation and the improvements that have been made to refine the DPLL algorithm into a Conflict-Driven Clause-Learning (CDCL) algorithm.

Chronological backtracking. To explain chronological backtracking, we will first introduce the concept of *conflicts*. A conflict occurs when the assumption of a variable value leads to an unsatisfiable CNF. In the DPLL algorithm this happens when we make a literal choice (in line 7 of Algorithm 1) and then the algorithm realizes that the new CNF that includes such assumption is unsatisfiable. In this case the algorithm will *backtrack* the decision (undo the assignment) and try with another value. Because of the recursive nature of the DPLL algorithm, we should notice that the backtracking done is always the exact inverse order as the order in which we made the assignments. For example, if an assignment $\Delta|A, B, C, D$, made in that same particular order, leads to a contradiction, then the algorithm would backtrack the last decision made and undo the D assignment. It will now try the $\neg D$ assignment, but if it also proves to lead to a contradiction, then it will also backtrack this decision. At this point it is implied that $\Delta|A, B, C$ is also a contradiction, but note that the DPLL algorithm did not detect the contradiction here (could only imply it after making the D and $\neg D$ assignments). Now the algorithm will also backtrack the C assignment and try $\neg C$. As we have seen in this brief example, the backtracking is always done in the same order that the assignments were done. In general, when we detect a conflict at level l of the search tree, we have to rewind back to level $l - 1$, undoing all assignments made in the process. Then we try another value at level $l - 1$, if none remains to be tried, we then go back further to level $l - 2$, and so on. If we ever reach level 0 and each value there leads to a contradiction, we will know that the CNF is inconsistent. This type of backtracking in the DPLL algorithm is called chronological backtracking, because we move back in the same order we got there: one level at a time. A more efficient way to backtrack conflicts is to use *non-chronological*

backtracking, but first we need to introduce a new feature called *clause learning* in order to understand how non-chronological backtracking works.

Clause learning. When a conflict is detected, a *conflict-analysis* procedure is used. Such procedure consists of analyzing the conflict structure through *implication graphs* and then learning a clause from it. The following example will explain how a new clause can be derived from a conflict.

Example 2.2.4. Let's consider the following CNF:

$$\Delta = \left\{ \begin{array}{l} 1. \{A, B\} \\ 2. \{B, C\} \\ 3. \{\neg A, \neg X, Y\} \\ 4. \{\neg A, X, Z\} \\ 5. \{\neg A, \neg Y, Z\} \\ 6. \{\neg A, X, \neg Z\} \\ 7. \{\neg A, \neg Y, \neg Z\} \end{array} \right. \quad (2.1)$$

The implication graph on Figure 2.3 shows a possible search path of a CDCL solver. The nodes of this graph are variable assignments made, with the notation $l/V = v$, where l is a decision level, V a variable and v a truth value. Remember that a variable might be assigned a value in two cases: by a branching decision or by an implication as a result of the propagation process. The directed edges in the graph indicate which assignments implicated other assignments, so an edge from node α to node β would mean that the variable assignment in node α implies the assignment shown in node β . Nodes that do not have any incident edge are decision assignments, because they weren't implied by other assignments. The numeric label on the arcs correspond to the number of the clause that was used for implying that assignment.

A *cut* in an implication graph (dashed lines in Figure 2.3) is a set of edges that separate the decision assignments (root nodes) from the contradiction (the leaf node containing $\{\}$). A *conflict set* is a set of variable assignments such that each has an outgoing edge belonging to the same cut, in other words, a conflict set contains enough variable assignments to cause the conflict [16, 26, 22]. Figure 2.3 shows three possible

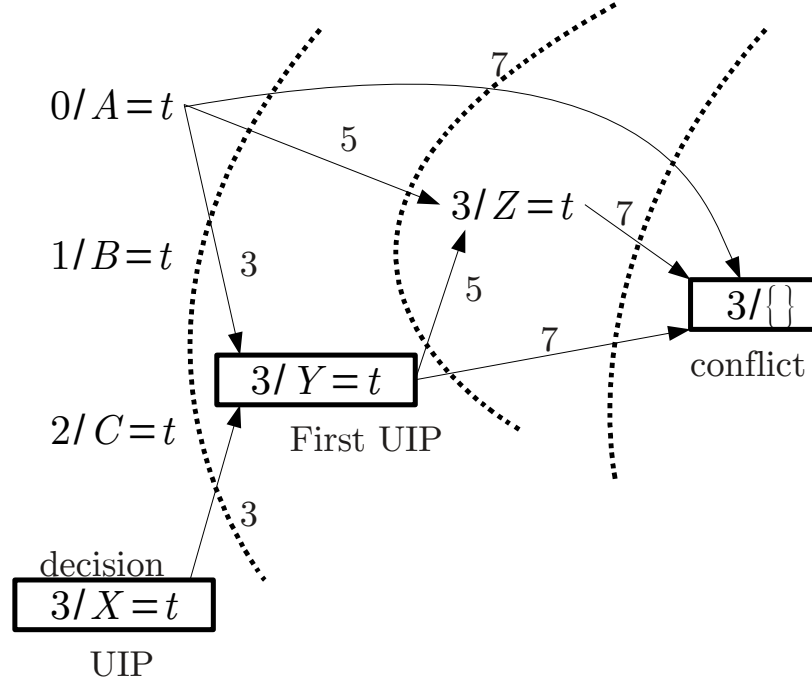


Figure 2.3: An implication graph of a conflict.

cuts for that implication graph, they lead to the conflict sets $\{A = \mathbf{true}, X = \mathbf{true}\}$, $\{A = \mathbf{true}, Y = \mathbf{true}\}$ and $\{A = \mathbf{true}, Y = \mathbf{true}, Z = \mathbf{true}\}$. The UIP label of some nodes will be explained later in this section.

Since there are many possible cuts, and thus conflict sets, we are only interested in choosing the most useful ones. An *asserting clause* [26] is a clause derived from a conflict set (containing the negation of the assignments in the conflict set) which contains exactly one variable assigned at the level of the conflict. From the previous conflict sets, $\{\neg A, \neg X\}$ and $\{\neg A, \neg Y\}$ are asserting clauses, because only X and Y were decided at the conflict level.

A refinement to the definition of an asserting clause is the notion of a *unique implication point* (UIP) [16, 26]. A UIP of a decision level is a variable which was set at that level and that lies in every path from the decision variable at that level to the conflict. For example, the assignments $3/Y = \mathbf{true}$ and $3/X = \mathbf{true}$ are both UIPs of level 3, and $0/A = \mathbf{true}$ is a UIP of level 0. UIPs of a decision level can also be given an order, the first UIP is the one closest to the contradiction. In Figure 2.3 the first UIP of level 3 is $3/Y = \mathbf{true}$. The *1UIP scheme*, used in modern solvers,

learns asserting conflict-driven clauses which have the first UIP of the decision level of the conflict. \square

Non-chronological backtracking. The problem with chronological backtracking, which is the one used in the DPLL algorithm, is that contradictions that trigger the backtrack often have valuable information. Such information could lead us to learn new clauses and backtrack more levels than just one, thus saving time in the search. For example, let us say that we have a CNF Δ with variables A, B, C, D , such that Δ will always be inconsistent when $A = \mathbf{true}$. Unfortunately, the DPLL algorithm might not realise this beforehand, because it uses unit resolution which is not refutation complete. So the algorithm might only conclude that $A = \mathbf{true}$ will not satisfy Δ after assigning all possible values to B, C, D . If we could detect that A is not part of any solution, we could avoid going any further in that branch of the search tree and learn that information as a new clause (in our example we would learn the clause $\{\neg A\}$ to enforce A to be **false**). Learning such clauses is called *conflict-driven clause-learning* (CDCL) [26, 15].

Adding CDCL features to the DPLL algorithm does not affect its completeness. It is not so obvious that completeness is maintained, because now the algorithm, unlike the basic DPLL version, can make the same decisions more than once (visit a node in the search tree various times), since it does not maintain memory of the previous decisions made. However, it does so in a different context, because it has learned clauses since its previous visits to the same node. This guarantees the completeness of this new algorithm. A completeness proof of the DPPL algorithm with CDCL features can be found in [27, 22]. In the next section we will discuss in detail how such new clauses are found or how can we decide which level to backtrack after a contradiction is found.

2.2.5 Modern CDCL solvers

Most modern solvers today are CDCL SAT solvers. They apply the same structure of the DPLL algorithm, but also include clause-learning features in them. In the last years most improvements have been focused on data structures and heuristic refinements. In this section we will explain the main CDCL algorithm and also the main

features which make modern CDCL SAT solvers.

CDCL algorithm. Given a CNF φ , a partial assignment of variables ν , Algorithm 3 outlines the general structure of a CDCL SAT solver, where x is a variable, v a truth value and β a decision level. We will shortly explain the main functions of this algorithm.

Algorithm 3: Typical CDCL algorithm

```

Input: A CNF  $\varphi$  and a variable assignment  $\nu$ 
1 if (UNITPROPAGATION( $\varphi, \nu$ ) == CONFLICT) then
2   return UNSAT.
3  $dl \leftarrow 0$ 
4 while (not ALLVARIABLESASSIGNED( $\varphi, \nu$ )) do
5    $(x, v) = \text{PICKBRANCHINGVARIABLE}(\varphi, \nu)$ 
6    $dl \leftarrow dl + 1$ 
7    $\nu \leftarrow \nu \cup \{(x, v)\}$ 
8   if (UNITPROPAGATION( $\varphi, \nu$ ) == CONFLICT) then
9      $\beta = \text{CONFLICTANALYSIS}(\varphi, \nu)$ 
10    if ( $\beta < 0$ ) then
11      return UNSAT
12    else
13      BACKTRACK( $\varphi, \nu, \beta$ )
14       $dl \leftarrow \beta$ 
15 return SAT

```

- UNITPROPAGATION consists of iteratively deducing the truth value of variables. The values are deduced by logical reasoning on φ and ν . We already discussed this function in the previous sections.
- PICKBRANCHINGVARIABLE consists of selecting a variable to assign, and the respective value. Heavily relies in heuristics/random factors to pick variables.
- CONFLICTANALYSIS consists of analyzing the most recent contradiction and learning a new clause from it. It returns the decision level to backtrack to (non-chronological backtracking).

- **BACKTRACK** undoes variable assignments and backtracks to a previous decision level as computed by **CONFLICTANALYSIS**.
- **ALLVARIABLESASSIGNED** tests whether all variables have been assigned a truth value.

Search restarts. SAT solvers may exhibit high runtime variability on some problems. This means that some decision heuristics and random factors make the solver take a really long time in solving a problem, but others can help solve the same problem in a very short runtime. This phenomena exhibited by SAT solvers and these kind of problems is known as a *heavy tailed distribution*. A heavy tailed distribution has infinite variance and infinite mean, so a SAT solver might get trapped on a very long run with a problem, while a different instance of the solver with other parameters could solve it fast. To address this problem modern SAT solvers have included search restarts in their solving algorithms. To avoid keeping the solver trapped in a very long search path, search restarts are issued over time and the solver starts the search all over again. In practice this strategy has shown dramatic improvements on problems which exhibit the heavy tailed behaviour [10]. It is also important to mention that learnt clauses can be kept between each restart, technique that still ensures the completeness of the algorithm (only one learnt clause in each iteration suffices to guarantee completeness[14]).

Variable decision heuristics. There are several heuristics that can be employed when choosing a decision variable, even making this decision completely random, but the most popular one was introduced by the solver **Chaff** and it is called Variable State Independent Decaying Sum (VSIDS). This heuristic consists of the following:

- Each literal has a counter, initialized to 0.
- The counter of a literal is only incremented in 1 when a new clause that contains that literal is learnt.
- When choosing a decision literal, the unassigned literal with the highest counter is chosen.

- Ties are broken by random.
- All counters are periodically divided by a constant.

MiniSAT [8] later made some refinements to this heuristic and instead of keeping counters for each literal, it keeps it for each variable. It also uses a dynamic increment factor (not just by 1) and does not use a periodic decay constant, but just scales all values down when one of them has become too large. This refinement of VSIDS is known as Exponential VSIDS (EVSIDS).

Clause cleanup. Adding clauses with no restrictions can be impractical in some cases. We could eventually exhaust all the available memory by learning too many clauses and it is also known that large clauses are not really useful in the search process [16]. Large clauses also add some considerable overhead to the search process and it is sometimes better to get rid of them than to keep them. Because of this, there are mainly 3 techniques employed to keep the clause database manageable:

1. Only keep clauses with n or less literals [7].
2. Clauses are kept only if they imply variable assignments or are unit clauses. They are discarded if the number of unassigned literals is greater than a number m [2].
3. Clauses with a number of literals greater than a threshold k are discarded as soon as the number of unassigned literals is greater than one [16].

Lately, some solvers have started using more advanced heuristics as clause deletion policies, such as keeping information about the activity of a clause in conflicts. One of the best performing heuristics is employed in the SAT solver **glucose** [1], which implements a particular metric of clause activity for its deletion heuristics.

2.2.6 Data structures

Data structures play a fundamental role in the performance of CDCL SAT solvers. Many improvements have been achieved over the last years, but one of the most noticeable ones is use of the so called *lazy data structures* to implement propagation.

There are different types of lazy data structures used in modern SAT solvers, they mainly try to address cache performance problems when the solver is performing unit resolution. In this section we will explain two important data structures that are used by almost all modern SAT solvers: One is the two watched literals and the other are the binary implication lists.

The two watched literals lazy data structure. The two main approaches to implement lazy data structures are the *head/tail lists* used in Sato [25] and the *two watched literals* used in Chaff [17]. We will explain the later in detail, as it is the one implemented in most modern CDCL SAT solvers.

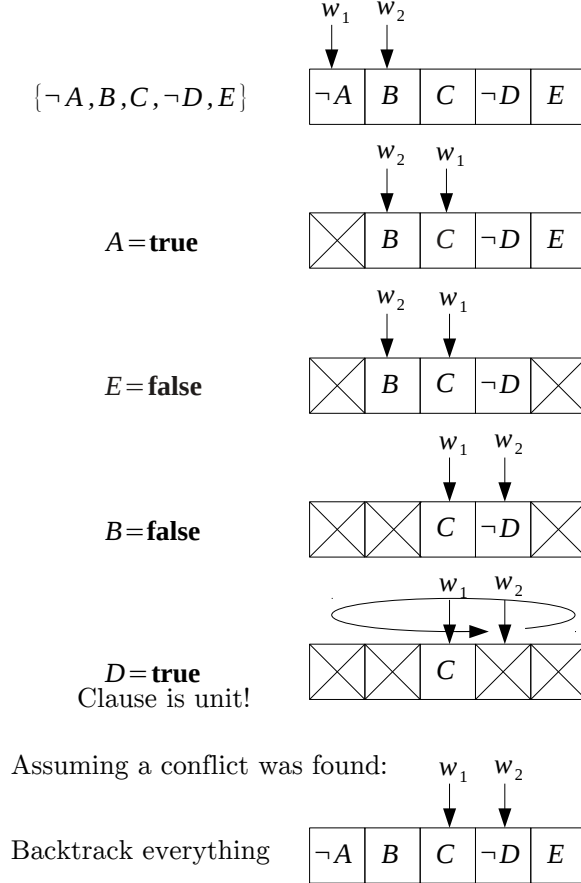


Figure 2.4: Operation of the two watched literal data structure.

As we have seen, unit propagation is a key function to speed up SAT solvers. We need fast ways of identifying which clauses become unit when assigning variables. For

example, if we have the clause

$$\{\neg A, B, C, \neg D, E\}$$

and the variable assignment

$$A = \mathbf{true}, E = \mathbf{false}, B = \mathbf{false}, D = \mathbf{true},$$

we would need to identify somehow that this clause is unit on variable C . The two watched literal data structure addresses this problem and makes backtracking very efficient. Instead of trying to keep precise information on the complete state of the clause, the two watched literal strategy only keeps track of two literals in each clause. Such two literals are the watched literals of a clause.

Example 2.2.5. The following example considers the clause $\{\neg A, B, C, \neg D, E\}$. Figure 2.4 shows how the two watched literal scheme would work on such a clause. The context of this example is a CDCL algorithm trying to solve a SAT problem. As we showed earlier, this algorithm will make variable assignments (according to the chosen heuristic), and after each assignment, it will call the `UNITPROPAGATION` function. It is in this function that the two watched literals scheme comes into play. The algorithm needs to check all clauses to identify all the ones that have become unit. For this example we will only be examining one particular clause and assume that the solver is making variable assignments and then checking the clause. First we will assume that no variable assignments have been done yet and that the two watched literals are $\neg A$ and B , which in Figure 2.4 correspond to w_1 and w_2 . Suppose that on the next decision level variable A is assigned as **true**. We would check the watched literals and realize that this clause might be unit, because now there is only one watched literal which is unassigned and the other does not satisfy the clause. Remember that we are only aware of the watched literals and not the rest of them. To check if the clause is really unit, we will attempt to change the first watched literal to an unassigned variable in the clause. We could search to the right and find that literal B is already being watched (the two watched literals must be different), so we keep searching to the right and find that literal C is unassigned, concluding that the clause is not yet

unit. Now our two watched literals for this clause are B and C . Let us suppose now that propagation gives the value **false** to variable E and then **false** to B , then we would again wonder if the clause is unit or not. Searching to the right for another unassigned literal to watch will find literal $\neg D$, so now our two watched literals will be C and $\neg D$. Finally, if variable D is assigned to **true**, then we would search the clause for an available literal, but will not find any (after checking all literals). Only now we can declare that this clause is unit, because there is only one unassigned watched literal and the other can not find any unassigned position. Propagation would assign **true** to variable C , because it is the only way to satisfy that clause. \square

If a conflict is found, while propagating across the different variables, we have to undo all variable assignments after the backtracking point, but the key advantage of the two watched literal structure is that no backtracking has to be done to any watched literal reference of any clause. The clear drawback to this advantage is that, as we saw in the previous example, we have to check all literals in the clause before declaring it unit, since we only keep track of two literals at a time.

Binary implication lists. To speed up the propagation of SAT solvers, *binary clauses*, clauses with two literals, are sometimes stored in special data structures called *binary implication lists*. Binary clauses are frequently used in propagation because they can imply other variable values in just one step. There is also no point on distinguishing two watched literals in these clauses for the simple reason that there are only two literals in them, so knowing when a binary clause is unit becomes trivial. Figure 2.5 represents a set of binary clauses stored in a binary implication list. We have a literal index vector to identify each literal (a variable and its negation), each position of this vector points to a list of literals which are implied by the literal corresponding to the origin of the pointer. Hence, if the $\neg A$ position in the index vector points to $\neg B$ and D , this means that if A were to be **false**, it would imply that B must be **false** and D **true**.

2.2.7 Parallel SAT solvers

As mentioned before, some parallel SAT solvers have performed at the top of the last SAT competitions, but even though they all fall into the parallel solvers category,

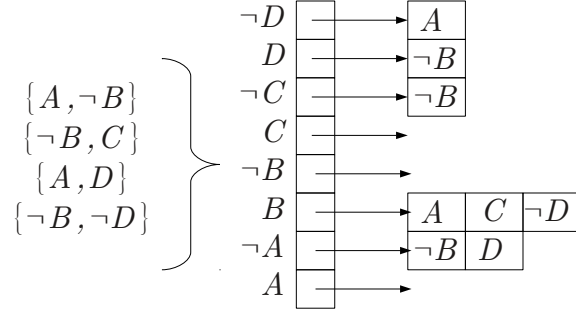


Figure 2.5: A group of binary clauses and their corresponding binary list.

their parallel strategies and implementations vastly differ from each other. We mainly classify parallel SAT solvers into two categories: portfolio approach solvers and divide-and-conquer ones.

The main idea behind portfolio approach solvers is that different kinds of sequential solvers will perform differently for different kinds of SAT problems. The portfolio approach is a very straight forward strategy: They run a group of sequential solvers in parallel, each with different heuristic values and/or different search strategies. The time they take to solve the problem will be the time of the fastest sequential solver in the group of solvers running in parallel. Although all portfolio approach solvers share this same principle, they also have quite different kinds of implementations. We identify in this group the solvers that do not share information, the ones that share clauses only logically, and the ones that share clauses physically and logically.

Solvers that do not share information have the most simple design. They run completely independent solvers in parallel and wait for one of them to give an answer. Despite their simplicity, the solver `ppfolio` [21], a portfolio approach solver that does not share any information among its parallel solvers, was the winner of the crafted and random categories of the 2011 SAT competition of parallel solvers, and second place in the application category.

On the other hand, we have more elaborated portfolio approach solvers, which can also share clauses logically between their different solvers. One of the characteristics of CDCL solvers is the fact that they can learn new lemmas as they solve a SAT problem. These new lemmas will provide additional information during the solution search, so that the solver doesn't fall into previous fruitless search paths.

The idea is that different solvers running in parallel can share their learned lemmas so that they all benefit from what other solvers have learned and improve their own search. An example of these kind of solvers is **ManySAT** [9], which won the 2009 SAT competition in the parallel solver application category. **ManySAT** has its own sequential state-of-the-art SAT solver and runs different instances of it in parallel, using different VSIDS [24] heuristics (branching heuristics) and restart policies for each of it. The difference with pure portfolio approach solvers, is that **ManySAT** also shares learned lemmas between solving threads. It is called logical sharing of clauses, because the lemmas are passed as messages between threads and they never share the same physical information in memory. The advantage of logical sharing is that it is easier to implement message passing between threads, than having threads reading and modifying the same memory locations, which often requires locks that could hinder the overall solver performance. One of the best parallel performing solvers, **Plingeling** [3], also shares clauses logically. It is a very weak sharing though, since it only shares unit lemmas and it does so through message passing, using a master thread to coordinate messages between worker threads.

Portfolio approach solvers that share clauses physically have the same strategy as mentioned before, but they share clauses by allowing threads to access the same memory locations, instead of message passing. One solver in this category is **SArTagnan** [12], which shares clauses logically and physically. This solver did not perform as well as **ppfolio**, it only achieved the 39th place in the last SAT solving competition of 2011, in the application category.

Divide-and-conquer solvers do not try to run different solvers in parallel, they run one solving instance, but try to parallelize the search and divide it between the different threads. A common strategy to divide the search space is to use *guiding paths*. A guiding path is a partial assignment of variables which restricts the search space of the SAT problem. A solver that divides its search space with guiding paths will assign threads to solve the CNF with the given partial assignment from the guiding path the thread was assigned with. Once a thread finishes searching a guiding path with no success, it can request another to keep searching. **MiraXT** [23] is a divide-and-conquer SAT solver which uses guiding paths. Moreover, different threads solving

different guiding paths also share a common clause database, in which they store their learned lemmas. This is another example of physical clause sharing. Figure 2.6

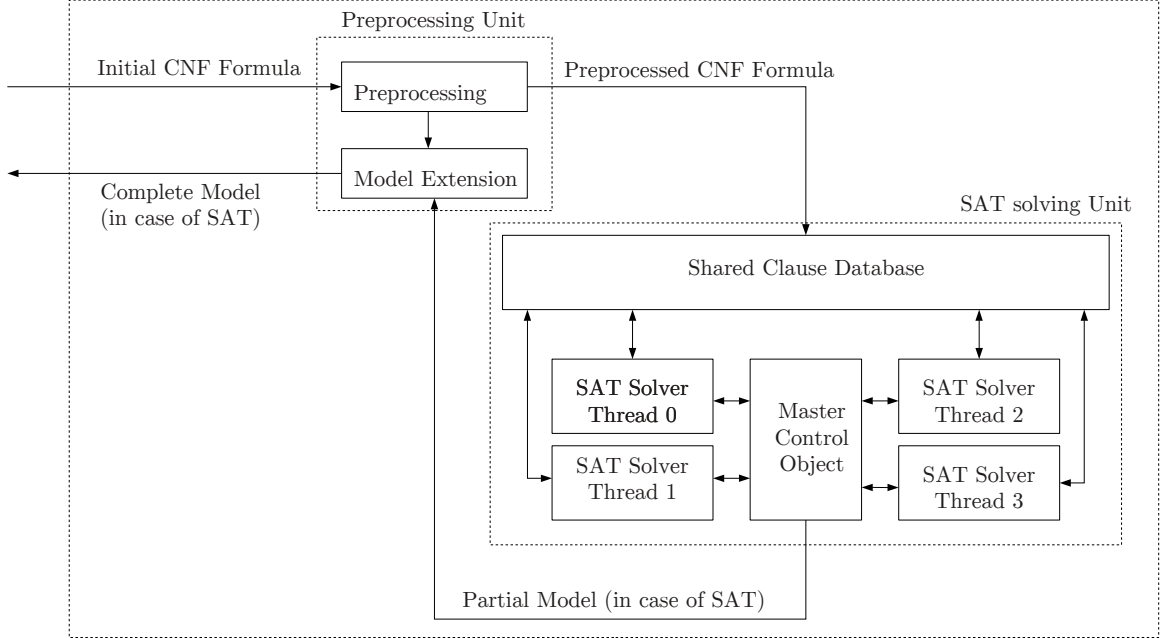


Figure 2.6: Overview of MiraXT design.

shows the general structure of the MiraXT solver and its multi-threaded organization. The *Preprocessing Unit* simplifies the initial CNF, a common process in all modern SAT solvers, but which is not relevant for this work. The *SAT Solving Unit* will be of high interest for our work though. The key structure in this design is the *Shared Clause Database*, which is a database of clauses shared among all SAT solving threads. Notice in this design that all threads are using a common pool of clauses called Shared Clause Database. There is also a *Master Control Object* (MCO) structure, which is responsible for the coordination of different threads. The reason for this is that, as we mentioned earlier, MiraXT is a divide-and-conquer solver, so different threads will contribute to one solution and hence will need to be coordinated between each other in this task. On the other hand, portfolio approach solvers do not need such component, because different threads will not need to interact with each other directly and no coordination between them is needed.

Figure 2.7 shows a representation of the shared clause database. Each thread keeps track of the two-watched-literals for the clauses it uses, and it also keeps two

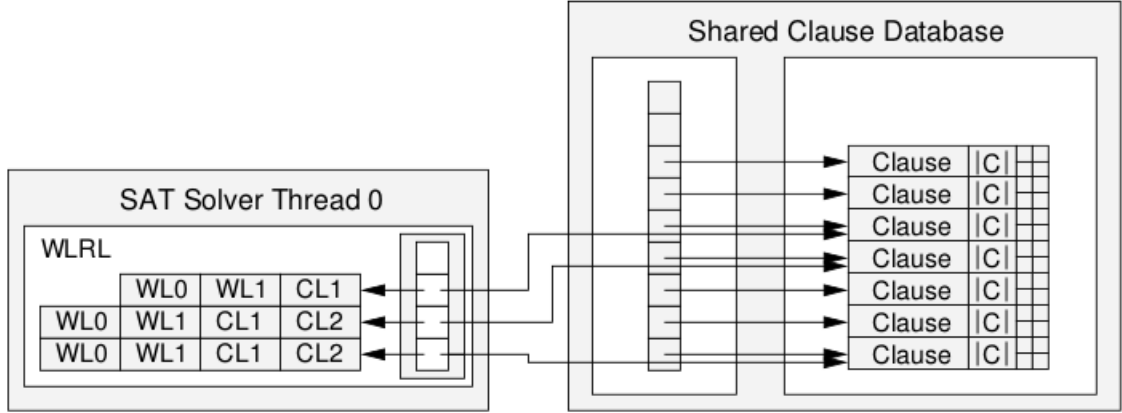


Figure 2.7: MiraXT shared clause database.

Cache Literals, which serve to replace any of the watched literals that need to be updated, so that we do not need to load them from the shared clause database. If none of the Cache Literals would serve, then they would need to be updated too from the original clauses in the Shared Clause Database. Each clause in each thread has a pointer to the original clause in the Shared Clause Database, so that it knows where to update the Cache Literals from. The Shared Clause Database is read only, so locks are only used when a new clause is inserted, authors mention that lock contention only account for less than 1% of total solving time.

To solve a problem MiraXT will start an initial thread solving the entire problem and depending on whether or not there are idle threads waiting, it will divide the problem into subproblems, with the guiding path technique, and share it with any idle thread. All communication and coordination between threads is done through the MCO with message passing, as shown in Figure 2.6.

2.3 Cache

Computers today usually have three levels of memory cache and a main memory. The processor can have multiple cores in it and cores can run multiple threads in them. The difference between a core and a thread is that cores have separate copies of almost all the hardware resources. The cores can run independently unless they are using the same resource (for example the connection to the outside) at the same

time. Threads, on the other hand, share almost all of the processor resources.

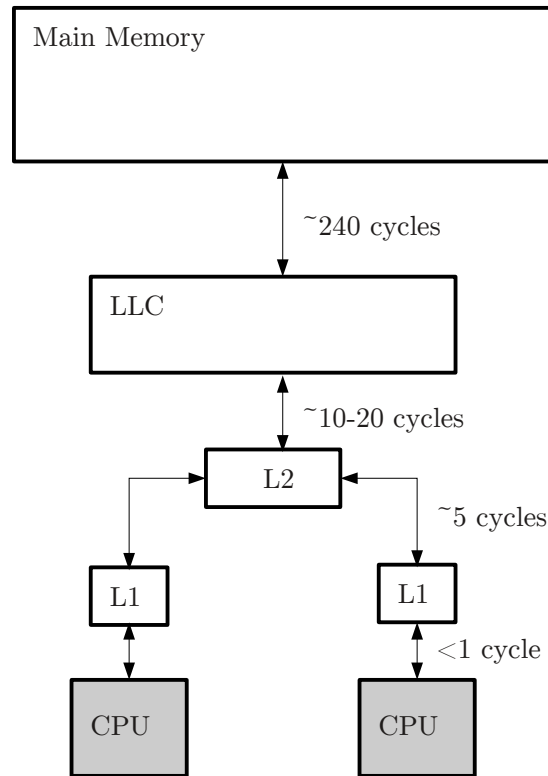


Figure 2.8: Typical memory hierarchy of modern computers.

When a thread, which is running on a core, needs to fetch data, it first tries to look for it in the first level cache (the L1 cache¹), if the data is not there, then it tries to find it in the level 2 cache (L2 cache). If the data is still not there, it then tries to fetch it from the level 3 cache (L3 cache) and if that fails too, it goes up to main memory to get the data. If it still isn't in main memory, then it has to retrieve it from the hard disk. We should notice that this hierarchy involves increasing fetch times as we go up. Getting data from the L1 cache is much faster than getting it from L2, and getting data from L2 is much faster than getting it from main memory. The problem with lower level memory is that, because of the technology and costs involved, they are much smaller. So the big picture is that at lower levels we have faster and smaller memories, and at higher levels we have massive and slower memory storages. Figure 2.8 is a schematic of today's computer memory hierarchy. To get an idea of the times

¹There is L1 data cache and also L1 instruction cache, we will be referring to L1 data cache

involved in accessing data from different memory storages, we present the following table of costs associated with hits and misses, for an Intel Pentium M:

To where	Cycles
Register	≤ 1
L1	3
LL	14
Main Memory	240

Chapter 3

Experiments and results

First we will present some simple experiments about cache behaviour, so we can have a better understanding of how cache can change its behaviour depending on the algorithms. Then we will run experiments with existing state-of-the-art SAT solvers and study their cache performance. Finally we will introduce our own proposed SAT solver, explain its components and features, and show the results we obtained from experimenting with it.

3.1 Cache and algorithms

When an algorithm requests data which is not found in any of the three caches, it has to be loaded from main memory. Data is not transferred individually, instead, a fixed amount of bytes containing the data (or part of it if it's large) is fetched. This fixed amount of bytes is called a *word* or *cache line*. Intel uses an *inclusive* memory cache protocol, which loads a requested word from main memory into all cache levels. Because transferring data from main memory is so costly (compared to any cache level), we would like to transfer the highest amount of useful data from main memory to cache every time. Since the access times to any cache are so little compared to main memory, we will assume the bottleneck of data-fetching performance is main memory.

Algorithm 4: Row sum of elements

Input: A matrix M of size $m \times n$, which elements are integers

```
1  $sum \leftarrow 0$ 
2 for  $i \leftarrow 0$  to  $m$  do
3   for  $j \leftarrow 0$  to  $n$  do
4      $sum = sum + M[i][j]$ 
5 return  $sum$ 
```

Algorithm 5: Column sum of elements

Input: A matrix M of size $m \times n$, which elements are integers

```

1  $sum \leftarrow 0$ 
2 for  $i \leftarrow 0$  to  $n$  do
3   for  $j \leftarrow 0$  to  $m$  do
4      $sum = sum + M[j][i]$ 
5 return  $sum$ 

```

Data-fetching performance is not only an issue that concerns hardware designers, but also programmers. Depending on how the program is written, the same task (input-output wise) might take much longer if one is not aware about how memory behaves. Since we want the highest amount of useful data in each word transferred, the data structures in which the programmer decides to store data will have a dramatic impact in cache performance. Consider, for example, the following problem: Write a program that sums all elements in a matrix size $m \times n$ of integers. Algorithm 4 and 5 both solve the problem, and apparently with the same level of efficiency, from an algorithmic point of view at least. However, the actual results of both algorithms in practice have a vast difference in total solving time. The following table shows the results for both algorithms implemented in C, for a matrix of size 31000×31000 :

Algorithm	Elapsed time (seconds)
Row sum	2.8
Column sum	28.3

Figure 3.1 shows a representation of how matrix M could be stored in main memory. As you can see, elements that are in the same row are stored in contiguous memory locations, while elements that are in different columns might be far away from each other. As we mentioned earlier, when an instruction requests data, it will bring a whole word that contains all or some of the data, from main memory to the cache. If we assume that the word size for our example is five integers, then when we fetch element a of the matrix, we will also be fetching the whole row together. If Algorithm 4 is being executed, then after using element a , we will need element b , but since we already loaded the whole row into cache, there is no need to fetch b from main memory! Same would happen with values b , c , d and e . If, on the other

hand, we were executing Algorithm 5, after using element a we would need to fetch element f , but element f was not loaded in cache, so we will need to fetch another word from main memory that contains the f element. Let's suppose now that we only have one level cache and that it can only hold one word at a time (remember cache is much smaller than main memory). Since the cache is full with the word that contains element a , we will need to *evict* this word. Evicting a word means pushing it back to main memory and marking the cache line as usable. Once the word containing element a is evicted from cache, we can then proceed to fetch the other cache line containing element f . After we have used element f , we will then need element b (because we are performing a column sum) and hence produce another eviction and access to main memory. In this simple example, the final count of main memory accesses for Algorithm 4 is two, while for Algorithm 5 is ten.

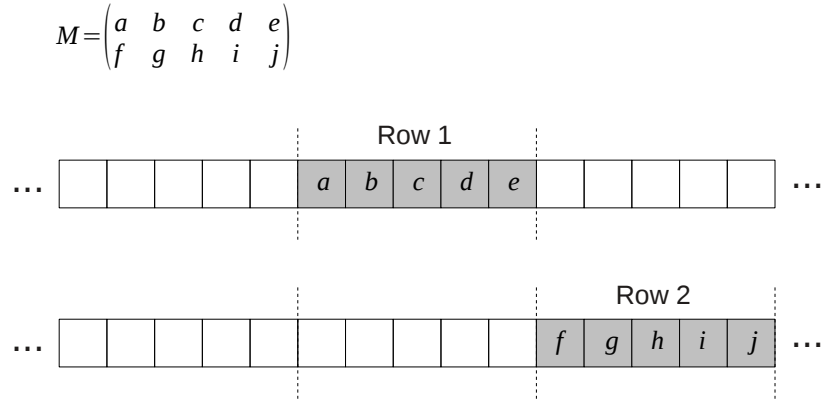


Figure 3.1: A representation of matrix M in main memory.

If we generalize our previous example to any matrix size, any word size and any cache size, we have the following formula to calculate the maximum amount of main memory accesses in Algorithm 4:

$$t = \left(\frac{n}{w} + 1 \right) \cdot m \quad (3.1)$$

Where t is the number of accesses to main memory, w the word size, n the number of rows of the matrix and m the number of columns. Equation (3.1) does not take into account the cache size since it will always need to transfer each word containing the rows once. In the big-oh notation, we would have that:

$$O(t) = \frac{n \cdot m}{w}$$

For Algorithm 5, the formula to calculate the amount of transfers could be:

$$t = (n \cdot m) - \left(\left(\frac{c}{w} - 1 \right) \cdot (w - 1) \cdot n \right), \quad (3.2)$$

assuming that $m \cdot n \gg c$, where c is the cache size.

Equation (3.2) also depends on other factors though, such as cache eviction policy. For that formula we are assuming a *FIFO* (*First In First Out*) policy, which means that the newest cache line fetched will be evicted when it's necessary. For this case, in big-oh notation, we would have that:

$$O(t) = n \cdot m$$

So Algorithm 4 will perform better for this problem and that is the reason why, in practice, we see such a big difference between the run time of both algorithms.

3.1.1 Perf: A cache performance tool

Perf [6] is a linux tool specialized in collecting and analyzing performance data. The advantage of **Perf** over other performance tools is that it uses special hardware registers dedicated to counting events associated with performance. Other popular tools, such as **Valgrind** [18], are software based and emulate the program being monitored to get performance statistics, process that could change the behaviour of the program being studied. The following hardware cache events can be viewed with **Perf** and will be used throughout this work:

- **L1-dcache-loads**: The amount of loads done from the L1 data cache to the CPU.
- **L1-dcache-load-misses**: The amount of *load misses* from the L1 data cache. A load miss is produced when data is requested from a particular memory level and is not found there, so a load from a higher level in the memory hierarchy is requested.

- **LLC-loads:** The amount of loads done from the last level cache (LLC) to the lower level cache (L1 or L2). On modern computers the LLC is usually the L3 cache, which is shared among different cores.
- **LLC-load-misses:** The amount of load misses from the LLC.

We will be using the terms cache misses and load misses indistinctively throughout this work. We will also assume that when we talk about cache we are referring to the last level cache (LLC). With all the previous information we can draw important results of the cache performance of any program. As an example of the use of `Perf`, we will make a small cache performance analysis of our C program of Algorithm 4 and Algorithm 5. The `Perf` statistics of cache events for Algorithm 4 will yield the following output:

Performance counter stats for './row_sum':

```
1,682,744,050 L1-dcache-loads
132,425,239 L1-dcache-load-misses # 7.87% of all L1-dcache hits
4,010,868 LLC-loads
3,607,332 LLC-load-misses # 89.94% of all LL-cache hits

2.787773717 seconds time elapsed
```

It should come to our attention that there is a low percentage of L1 cache misses and a high percentage of LL cache misses. The reason why L1 cache misses are so low is because Algorithm 4 sums elements by row, so almost all data that is loaded in L1 is used by the CPU, making the hit percentage very high, as shown in the results. On the contrary, we can also observe that the miss percentage of the LL cache is really high. The reason for this is because when summing elements by rows, we will never need to load the same cache line from the LL cache to L1 (when we load the cache line into L1 we will use all of its elements), hence every time we request data from the LL cache, it will not be there since it's the first time we are requesting that line.

The next output corresponds to the `Perf` stats of the cache performance of Algorithm 5:

Performance counter stats for './column_sum':

```

2,652,844,624 L1-dcache-loads
2,127,243,131 L1-dcache-load-misses  # 80.19% of all L1-dcache hits
2,037,794,644 LLC-loads
    506,598,124 LLC-load-misses      # 24.86% of all LL-cache hits

27.936225613 seconds time elapsed

```

This time the percentage of L1 load misses is very high, this is because we only use one element of the cache line loaded into L1, the next element will be in a different cache line (different columns are not contiguous in memory) so L1 cache hit percentage will be low. On the other hand, the percentage of LL cache misses is much lower than for the row sum algorithm, since we need to load the same cache line more than once, there is a good chance that a line requested will already be in the LL cache. It is important to note that we are mostly interested on what happens in the LL cache, when analyzing cache performance, because every load miss from the LL cache means a load from main memory, which are much more expensive than any other cache load. Under this criteria, one could argue that column sum is better than row sum, because it has a much lower load-miss percentage, but that's why it is also important to take into account the total number of loads and not only percentages. Row sum, despite having a much higher percentage of load-misses, has only a small fraction of the total load-misses on the LL cache that column sum has, hence a much lower number of total loads from main memory.

3.1.2 Size does matter

Aside from the code itself, the amount of data a program handles is also an important factor in cache performance. Let's say we have a square matrix M of size n and that we will perform a fixed number of sums of random elements in the matrix. From an algorithmic point of view, the size of M makes no difference, because the number of sums we will perform is fixed. But again, if we take cache performance into account, the size of matrix M will have a big impact in the time of a program performing this

Matrix size	Time (s)	L1 loads	L2 loads	LLC loads	MM Loads
16 kB	20.8	32673	8	< 1	< 1
130 kB	20.5	32675	745	8	< 1
4 MB	27.6	32653	1227	910	15
3000 MB	126.5	33278	3379	2570	1446

Table 3.1: Different times and number of memory loads for different size of matrices. The number of loads are in millions of loads.

task. Table 3.1 shows the different times and cache loads to perform a fixed amount of sums of random elements of different sized matrices. As expected, the number of loads from L1 is kept constant (the small variations are due to other system process running), because we are performing the same amount of operations in all tests. We can also observe that as we increase the size of the matrix, the number of loads from higher level memories also starts increasing. The reason for this is because as the matrix gets bigger, the lower level caches are not big enough to hold all of it, so it uses higher level cache/memory. Only when we start making loads from main memory is when we start noticing differences in time, these results can give us an insight on how expensive loads from main memory are compared to cache ones. The machine has a 256 kB L1 cache, a 1024 kB L2 cache, a 8 MB LL cache and 6 GB of main memory.

3.1.3 Cache performance in simple parallel-shared-memory programs

The main motivation behind this work is our experimental evidence that when threads in different cores share memory, the cache performance improves compared to having threads using their own memory. This experiment will consist on four threads in four different cores of a same chip performing sums over random elements of a matrix. At first, every thread will have its own copy of the same matrix and perform the operations over it. Then we will repeat the experience, but now with all threads performing operations over the same physical matrix. We will also vary the size of the matrix and only use read operations over the matrix, to keep concurrency problems out of the equation.

To measure cache performance we will use **Perf**. Table 3.2 shows the results of

Matrix size	Sharing	Time (s)	L1 loads	L2 loads	LLC loads	MM Loads
128 kB	no	4.37	3199	1353	1	< 1
128 kB	yes	4.39	3200	1365	17	< 1
512 kB	no	4.33	3201	1642	792	< 1
512 kB	yes	4.40	3201	1720	834	< 1
4 MB	no	11.09	3226	2312	1513	776
4 MB	yes	4.88	3205	2175	1512	8
600 MB	no	24.33	4720	5271	3679	2190
600 MB	yes	17.10	3484	5208	3691	1597
1.12 GB	no	29.09	6045	5607	3953	2594
1.12 GB	yes	19.28	3711	5371	3945	1738

Table 3.2: Times and memory loads vary depending on size of the matrix and if it is being shared.

the experiment. Notice that when we share the matrix among threads the cache performance increases significantly. This is because all cores in a single chip share the same LL cache, which has a limited size, so not sharing the matrix means handling a higher volume of data, which hinders cache performance as we saw in the previous section. On the other hand, if we share the matrix, then there is a higher chance that the requested matrix element will be found in cache, thus reducing the number of total main memory accesses. For the 128kB and 512kB sized matrices, we can easily fit the four copies, one for each thread, in the LLC, so that is why we barely notice any time difference. When the size gets to 4MB though, a single copy of the matrix is small enough to fit in the LLC, but four copies will make up to 16MB of memory and would not fit in the LLC. That is why sharing a 4MB matrix keeps a similar time to the smaller matrices, but not sharing it presents a 56% increase in time for this case, because of all the LLC loads the machine has to perform as 16MB is much bigger than what the LLC can hold. Remember that the LLC of the machine has a size of 8MB.

The relation of this experiment to our work is that SAT solvers have a clause database, which would be analogous to the matrix, and different solving threads can also share this clause database, as we do with the matrix. So from this small experiment we can get a hint that sharing the clause database should improve the cache performance of a portfolio approach SAT solver. No doubt that a SAT solver

is a much more complex program than the experiment we just did (sharing a clause database is not as simple as sharing the matrix in this experiment) so it is still not clear if the same results will be obtained when this strategy is applied to a parallel SAT solver.

3.2 Cache performance of the portfolio-approach SAT solver `plingeling`

Our first step will be to measure and identify the performance problems that afflict portfolio-approach parallel SAT solvers. The state-of-the-art portfolio approach parallel SAT solver `plingeling` is one of the best performing parallel solvers to this day. This solver will not share clauses in any way¹ and we will measure the performance behavior as we add more solving threads.

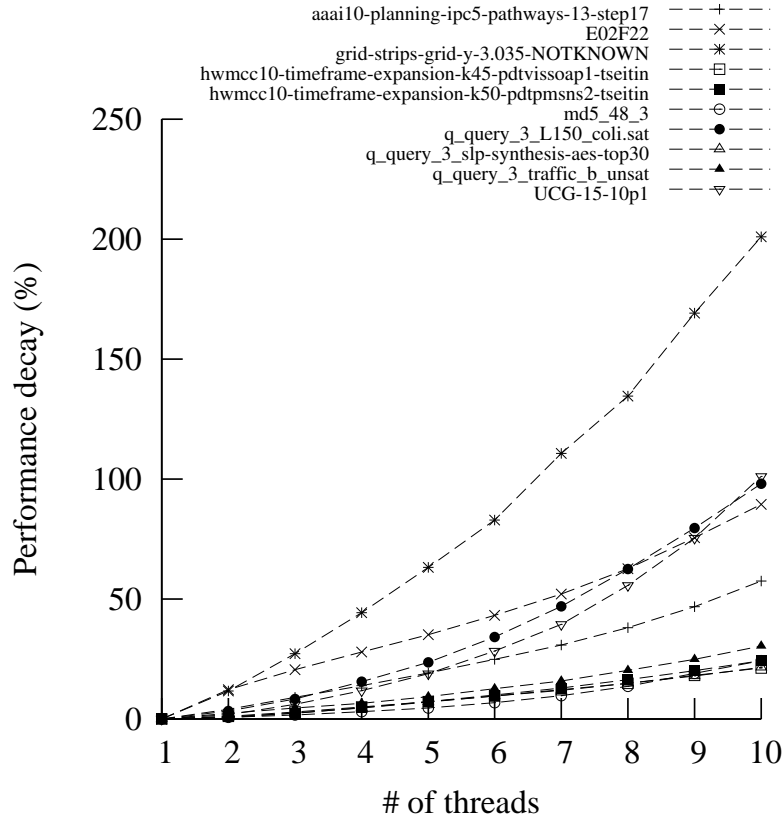


Figure 3.2: Decay of `plingeling` solver performance when adding threads.

¹It does share unary clauses through message-passing, but we have deactivated this feature for our experiments.

Because portfolio approach SAT solvers implement different search strategies among their solving threads, it is hard to measure the real impact on performance of adding an extra thread. The new thread might implement a successful search strategy, which will make the total solving time improve drastically and hide the negative impact on the cache performance when adding a new thread. This is why we have modified **plingeling** to make the exact same search in all of its threads. In theory, we would expect the solver to finish at the same time with one thread than with n threads, because all threads are performing the exact same search and thus should finish at the same time. Nonetheless, we know that each solver thread also keeps its own clause database, so basing our reasoning on the previous experiment, we should expect that adding threads will hinder the overall solver performance. This modified **plingeling** solver with n threads should perform worse as n grows, assuming that we have enough cores to run n threads and that all cores belong to a same chip. Figure 3.2 shows how the **plingeling** solver decreases its performance as we add more threads. Each line represents a different benchmark SAT problem (same problem set as used in Figure 3.2), the X axis is the number of threads being used, and the Y axis is the performance decay %, which is calculated as:

$$P_d = \left(\left(\frac{T_1}{T_n} \right) - 1 \right) \cdot 100$$

Where P_d is performance decrease, T_1 the solving time with one thread, and T_n the solving time with n threads. The decrease in performance is very noticeable as we add threads and it can even reach up to 200%! (which intuitively means that it takes three times more than its one threaded version). Our first hypothesis is that this decrease in performance is mostly due to cache issues.

To prove that the problem in scalability lies in cache, we have also performed the experiment on four different chips, running each different thread in a different chip. As chips have their own last level cache, we expect the performance decrease to be minimal as we add threads, because different threads will not be sharing cache. Figure 3.3(b) shows the decrease in performance as we add threads in separate chips. We can observe that the performance decrease in this case is minimal for most problems (only around 1-2%), contrary to the situation where threads are in different cores of a

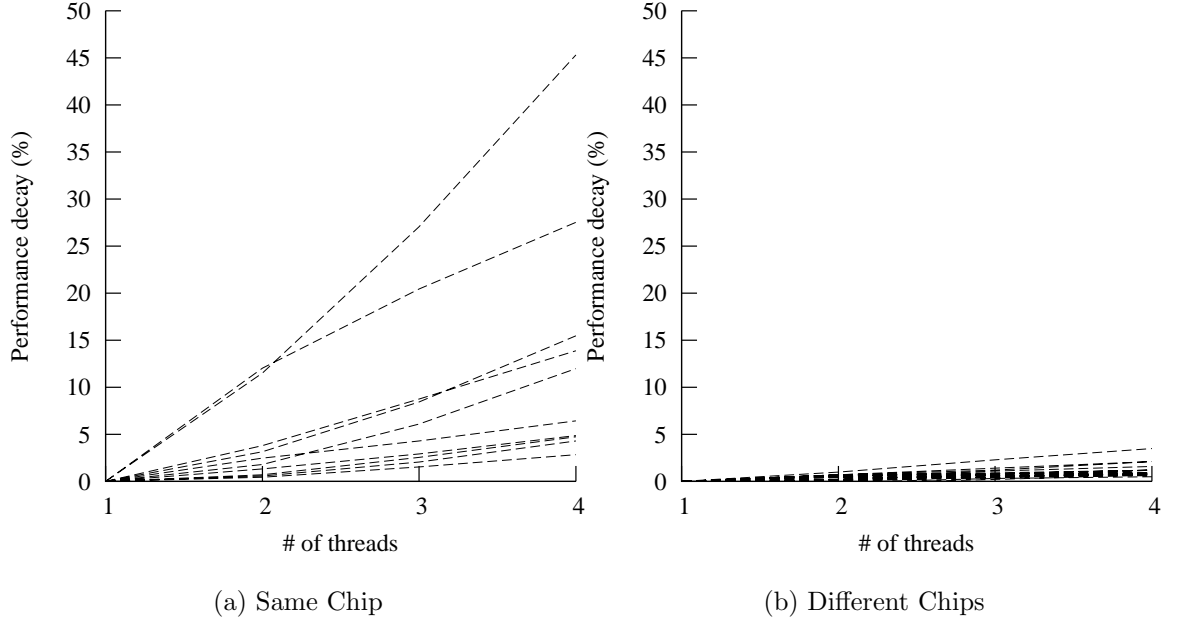


Figure 3.3: `plingeling` performance on different machine set ups.

same chip (where many problems exceed a 4% decrease), as shown in Figure 3.3(a). The fundamental difference between the two set ups used is that cores in the same chip share the LLC, while cores in different chips have their own LLC and only share main memory. Each solver has its own database of clauses, so the more LLC memory it has at its disposal, the better it will perform individually. These results are an indirect proof of our initial hypothesis, we have not measured the cache performance directly, but the main hardware resource being shared by cores of a same chip is the LLC.

To take our experimental claims even further, we will also measure the cache performance of the `plingeling` solver in a machine with a chip of six physical cores. As we can observe in Figure 3.4, the miss rate of the LLC highly increases as we add more threads. This means that there will be an important performance decay caused by cache, which is what we observe in previous performance experiments.

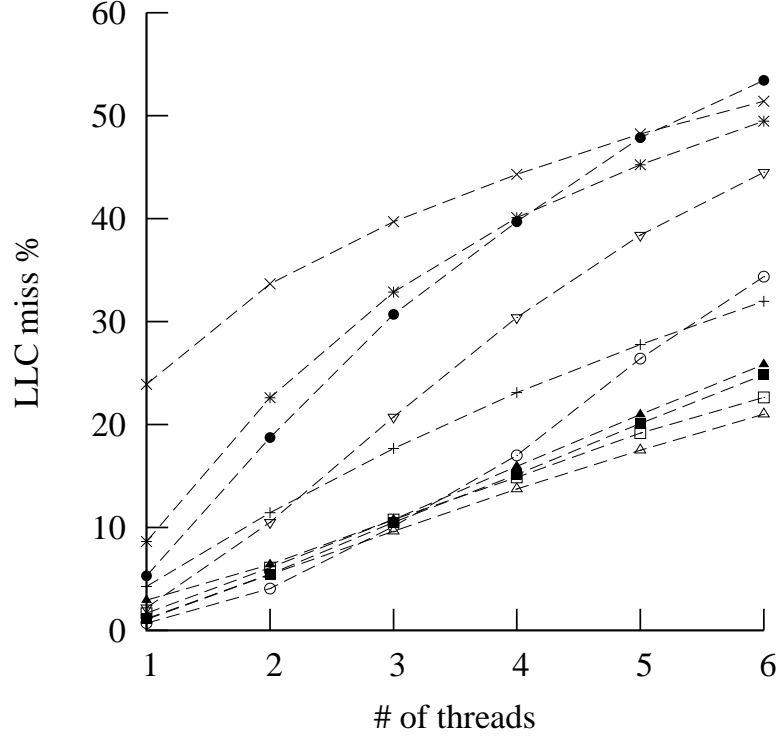


Figure 3.4: LLC miss % when adding threads. Same legend as Figure 3.2

3.3 AzuDICI: A parallel portfolio approach SAT solver with physically shared clause database

It is now clear that as we add more data to a SAT solver clause database, we make its performance decrease because of the cache misses involved in handling a bigger volume of data. We will now perform an analysis on how well does sharing data in SAT solvers help reduce the cache misses as we add threads. To be able to do such analysis, we implemented AzuDICI: a portfolio-approach SAT solver that shares clauses physically. As we mentioned earlier, there are other solvers which implement physically shared clause databases, but they do not make a detailed analysis of the cache performance of such solvers, which will be the contribution of our experiments with AzuDICI.

AzuDICI² is a standard CDCL solver mostly based on `barcelogic` and `miraXT`.

²You can find the latest implementation of AzuDICI at <https://github.com/leoferres/AzuDICI>

In particular, AzuDICI implements binary implication lists for the propagation with binary clauses, and the two-watched literals scheme for unit propagation with clauses of more than two literals. For the multithreaded versions that share clauses, the two-watched literals scheme has been modified and it will be explained later in this chapter. For conflict detection and clause learning, AzuDICI implements the 1-UIP algorithm. A lemma simplification algorithm is used for new learnt clauses, it's the same used in `PicoSAT`. For search restarts, it uses Luby restarts [13]. For database database clean up it keeps binary and ternary lemmas, and more than four-literal lemmas that have participated in a conflict since the last cleanup. Finally, AzuDICI also incorporates the EVSIDS heuristic for branching literal decisions.

To allow a a comparison of clause sharing in these kind of SAT solvers, AzuDICI comes in three flavours: AzuDICI-SharedAll, AzuDICI-SharedBinaries and AzuDICI-SharedNone.

3.3.1 AzuDICI-SharedNone

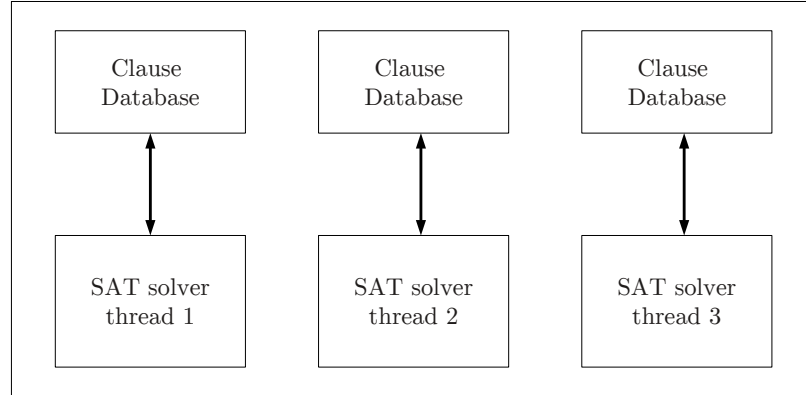


Figure 3.5: Three threads in an AzuDICI-SharedNone solver.

This solver is a pure portfolio approach CDCL SAT solver implementation that doesn't share any kind of clause. Each thread has its own AzuDICI sequential solver and its own clause database, Figure 3.5 is a schematization of the solver's internal structure. No locks or special data structures are required for this version, since we do not share any clauses (logically or physically), there are only completely independent solvers working on each thread. Each solver will have a different search, because they

will use different random values for its heuristics.

3.3.2 AzuDICI-SharedBinaries

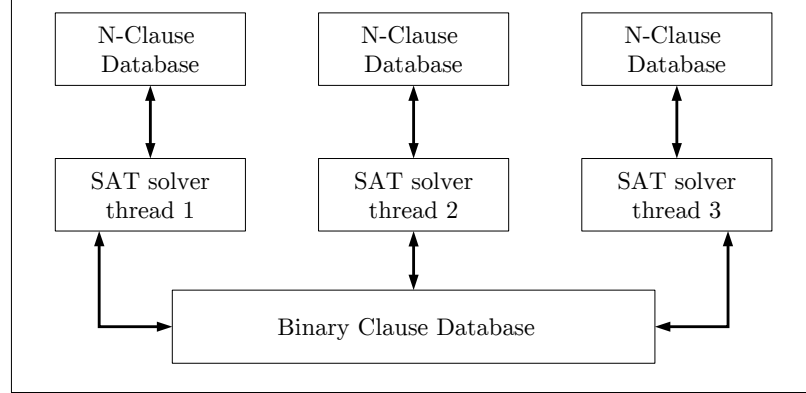


Figure 3.6: Three threads in an AzuDICI-SharedBinaries solver.

This solver shares binary clauses between its threads physically. All threads have access to the same binary implication lists and they can read and add binary clauses when necessary. Figure 3.6 shows the structure of this solver. Each thread has its own n-clause database, but they all share the same source of binary clauses. The advantage of this approach over sharing everything, is that a shared implication list of binary clauses is simple to implement and does not add significant overhead to the overall performance of the solver. This is because we do not need to keep additional information for each thread and propagation works the same way as it would without sharing the binary clause database.

The implication list used in AzuDICI differs in some aspects to the one studied in the previous chapters. Figure 3.7 is a schematization of our binary implication lists structure. We have an array of *binary lists*, one for each literal. A binary list is basically two pointers, one to a first *binary node* and another to the last binary node associated with that list. A binary node is an array of literals that also has a pointer to another binary node. The amount of literals a binary node can hold will depend on the size of the cache line we are working with; it will have as many literals as a cache line can hold. The literals implied by the literal associated to a binary list will be the ones in the binary node referenced by that binary list pointer and the

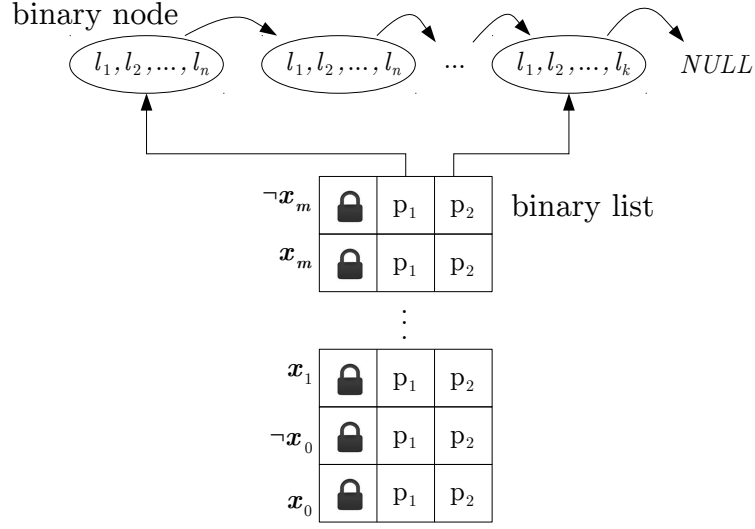


Figure 3.7: Binary clause database

subsequently referenced binary nodes.

When a thread wants to add the clause $\{l_i, l_j\}$, it must look for the binary list associated with $\neg l_i$ and go to the last node linked to that binary list. If there is enough space in that node to add another literal, then it adds l_j . If the node is full, then it must create a new node with the l_j literal, insert it at the end of the linked list of nodes, updating the binary list last node pointer. It does the same for the binary list of $\neg l_j$.

To ensure consistency of data when multiple threads are inserting, each binary list has a lock. If a thread is inserting a new implicated literal, it first locks the binary list where it is inserting and then proceeds to insert. If, by chance, another thread wants to insert in the same binary list, it must wait until the lock is freed. Since adding binary clauses is not a frequent event, and even less frequent the event that it would happen in the same binary list, the contention that these locks generate is unnoticeable in our experiments.

Another feature that sharing binary clauses brings in is that it adds another random factor to the search. Since threads are not synchronized with each other, they can add new binary clauses to the database at any given time, action that would affect the search of other threads (because they now have an extra clause to propagate with). If we were to try and replicate the exact same search as used in a particular

instance of an AzuDICI-SharedNone solver, we would fail, because sharing binary clauses among threads also affects the search path of each individual thread.

3.3.3 AzuDICI-SharedAll

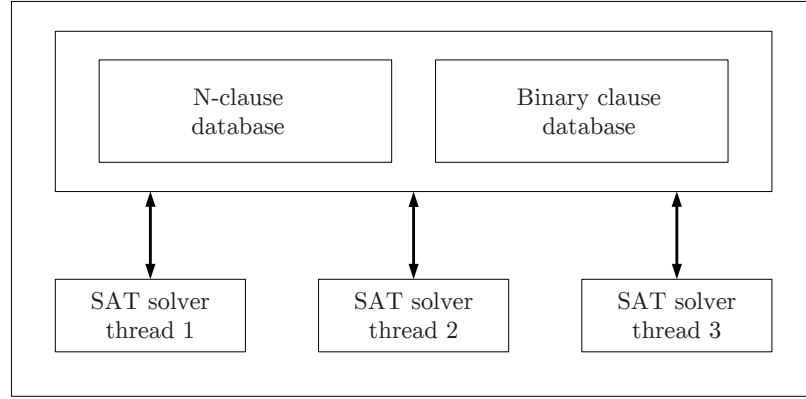


Figure 3.8: Three threads in an AzuDICI-SharedAll solver.

This version physically shares all clauses between threads, as shown in Figure 3.8. The complexity, compared to the previous versions, of implementing such solver lies in the propagation on n-clauses. In a sequential solver, the two-watched-literal scheme is usually implemented in a way that makes changes to the clause in order to identify the literals being watched at one given instant (for example, place both literals first in the clause). On the other hand, these changes to the clause are not feasible when many threads are sharing n-ary clauses, because multiple threads can modify the position of literals. Instead, we have used a similar approach as used in **miraXT**, where each thread keeps track of the literals being watched in each clause. Figure 3.9 is a schematization of how each thread worker relates with the n-ary clause database. Each SAT solver thread has a vector of pointers to *thread clauses* called *watches*, and each literal present in the SAT problem has a position associated in this watches vector. A thread clause is just a reference to an actual n-clause, which keeps extra information the threads needs. It has two watched literals (WL0 and WL1), two pointers to other thread clauses (NW0 and NW1) and a pointer to the actual n-ary clause in the n-ary shared clause database. WL0 and WL1 keep track of the literals being watched by the thread for a given n-ary clause. NW0 and NW1 point

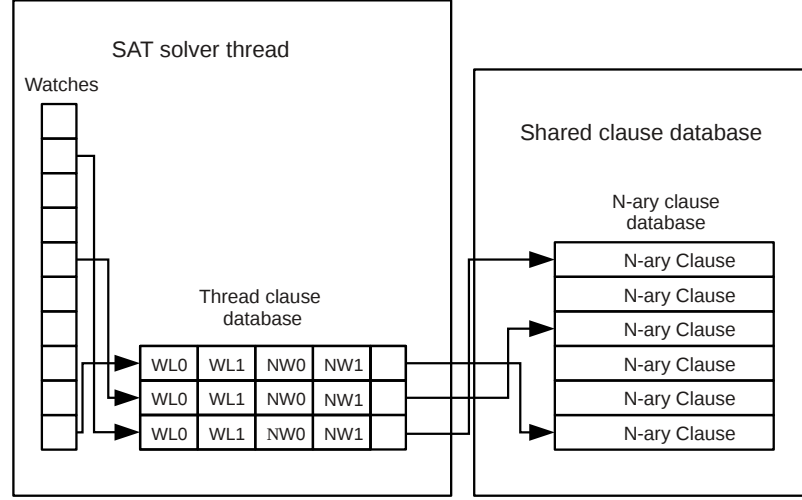


Figure 3.9: The thread clause database and n-ary clause database

to the next thread clauses that are also watching WL0 and WL1. The n-clause also has a set of flags, one for each worker thread for them to identify which ones are using that clause for propagation.

To insert a new n-ary clause, we first make sure that the clause does not exist in the database. If it does not exist, we create the n-clause, set the current thread flag to true and add it to the database. On the other hand, if it does exist, we just toggle the corresponding thread flag of the n-clause to true. The inserting procedure is locked so that two different threads can not insert at the same time. In our experiments we have not noticed any considerable overhead caused by this lock.

3.4 AzuDICI performance

Since we are interested in studying the cache performance of parallel SAT solvers, we will use a benchmark set of SAT problems (all used in previous SAT competitions) which we know that any of the AzuDICI versions can not solve in a defined time frame limit. This time limit was set to five minutes for our experiments. We would not want to use problems that can be solved in our time limit frame mostly because the results would be strongly influenced by the different search paths each version takes. A version that solves a problem faster due to finding a better search path will probably have a better cache performance, not because it is more cache-friendly, but

because it finished before the amount of data to handle was too big.

The experiment will consist on measuring the cache performance of a solver before the timeout of five minutes. For each problem, we will run instances of the solver with one to up to n threads, and see how well cache behaves for that particular problem.

3.4.1 AzuDICI-SharedNone results

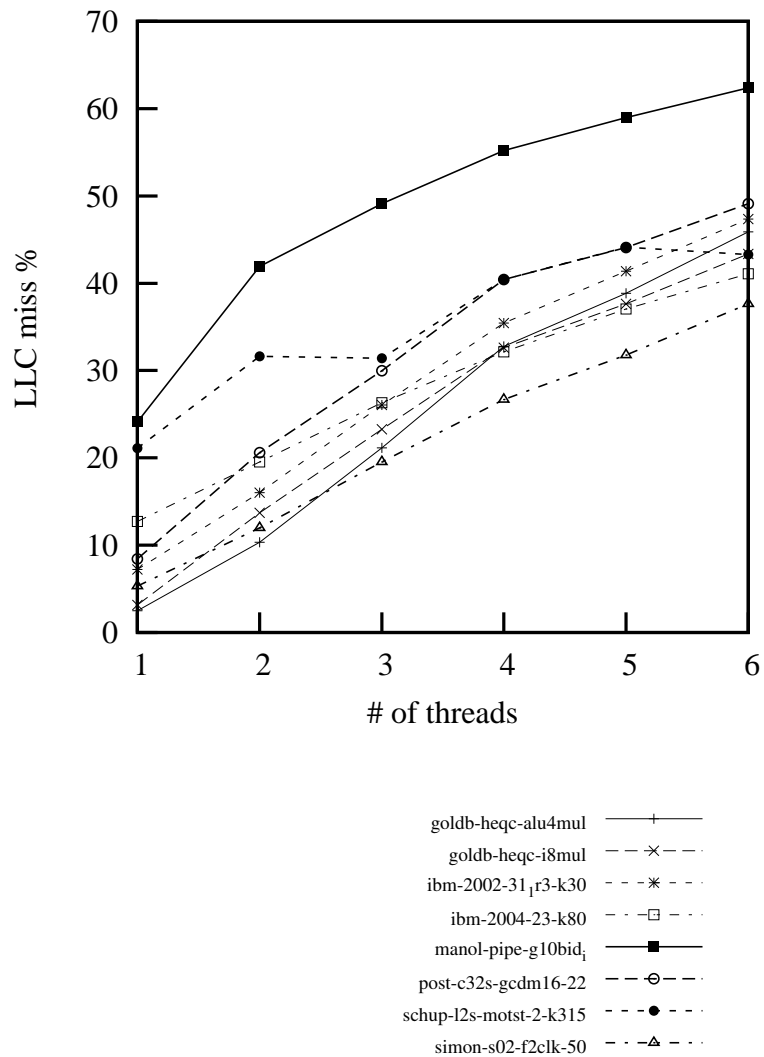


Figure 3.10: SharedNone experiment

In this experiment we ran the SharedNone version with our set of benchmark problems. These benchmark problem names are identified in the legend and correspond to SAT problems used in previous SAT competitions. Figure 3.10 shows the results for this experiment. As we expected, the cache performance decreases as we add more threads, the LLC miss percentage jumps from around 5-10%, for one thread, to about 35-50% when running on six threads. The overall scaling for all problems is poor and the continuous increment in LLC misses suggests that the propagation of clauses is much more slower when working with multiple threads. We can infer from this graph that if we continue adding threads (assuming we had enough cores), the cache performance would be so low that it would not be beneficial to add any more threads, as the propagation of clauses would be too slow to add any improvement. Unfortunately, finding this point, where adding more threads becomes counter productive, should be very difficult, because each problem may show a very different behaviour and each new thread might also behave very differently, yielding different optimal number of threads. We could perform a statistical study over a very large sample of benchmark problems to find that number of optimal threads for a typical SAT problem, but we will leave such experiment for future work. For now, we are only interested in knowing that when we don't share clauses physically, the scaling of cache performance is poor.

3.4.2 AzuDICI-SharedBinaries results

Figure 3.11 shows the results in this experiment and we can observe they are similar to the SharedNone results. This can be expected as we are only sharing a small amount of clauses physically and not making any significant change to the solver. The propagation scheme is still the same for both versions. Only a small improvement, not more than a mere 2% in most cases, in cache performance can be noticed throughout the different benchmark files. This experiment is an intermediate example of clause sharing as we only share binary ones. Once again, we see that the overall scaling as we add threads is poor and that there should be a point when adding more threads is counter productive, just as in the previous version.

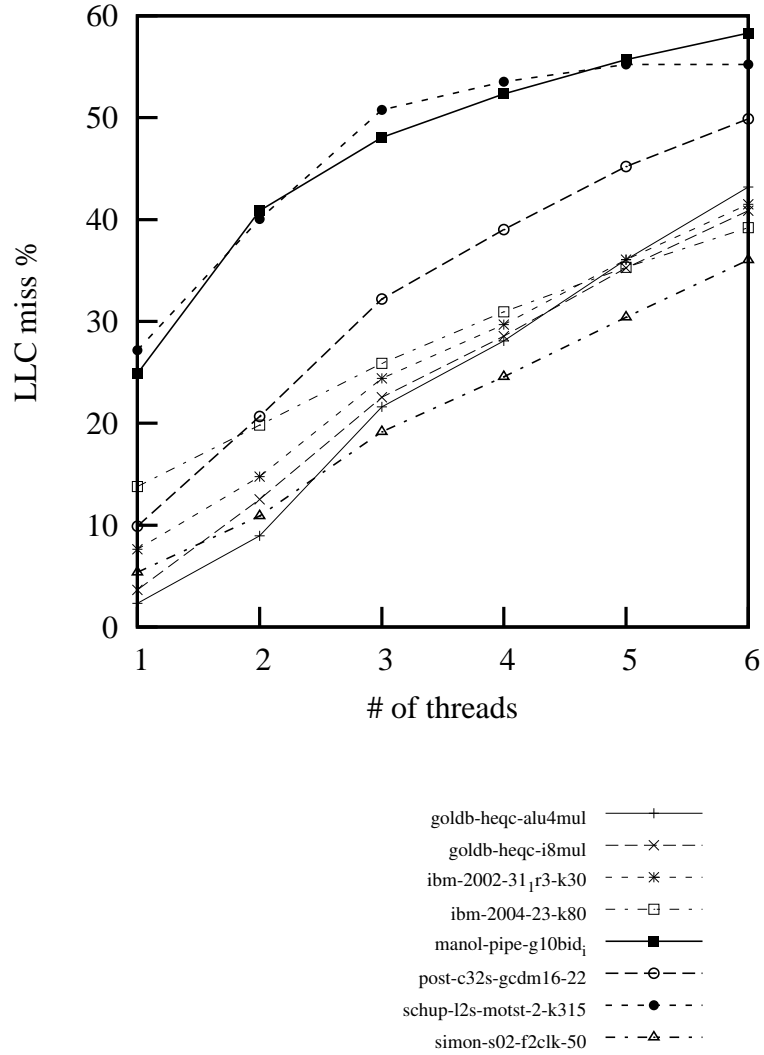


Figure 3.11: SharedBinaries experiment

3.4.3 AzuDICI-SharedAll results

The results for this last version are shown in Figure 3.12. We expected the scaling of this solver to be much better than the previous versions, because sharing all clauses between threads lowers down the amount of total data to handle and should keep the cache misses under control, but our results show otherwise. Not only did this SharedAll version not achieve the desired improved scalability, it also makes the

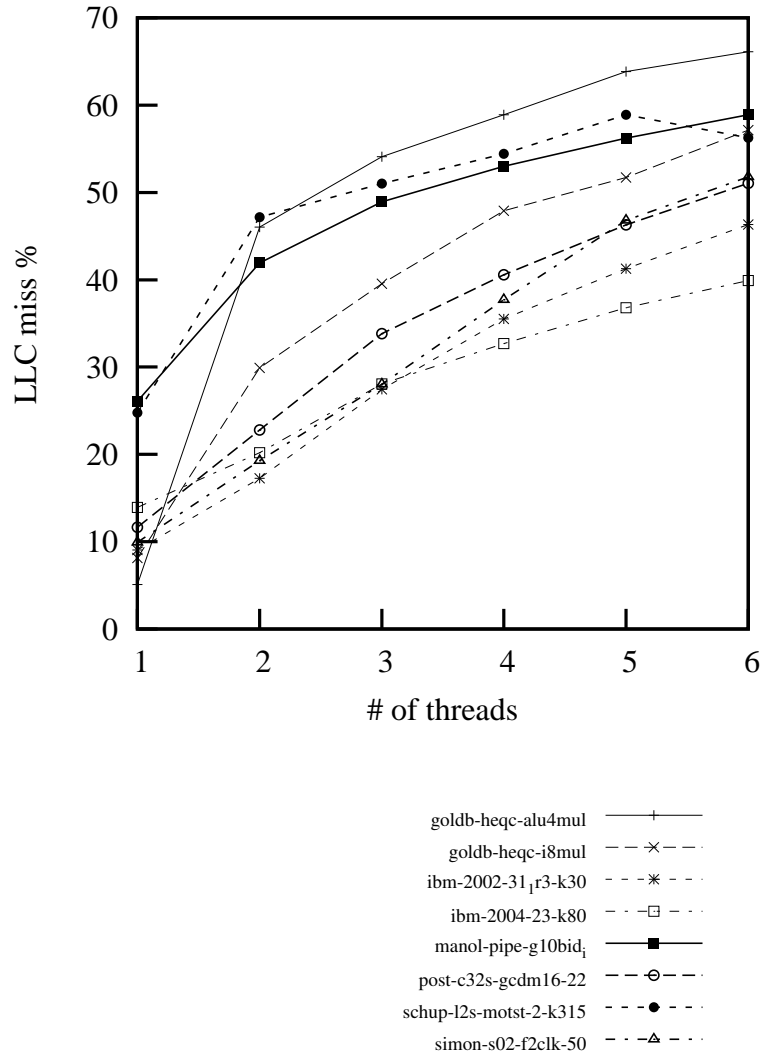


Figure 3.12: SharedAll experiment

amount of cache misses higher for all number of threads. Our explanation for this behaviour is that this solver keeps more information and has to perform additional tasks when propagating. Unlike the SharedNone version, this one needs extra structures for each thread, which keep track of the watched literals of each thread and which clauses are being used. We suspect that this additional data, plus the fact that we have to be constantly checking it during propagation, makes the cache perform even worse and totally shadow the benefits of sharing information. Also, as we described

in an earlier section of this work, the amount of information to handle might be so big that it won't even matter if we share it, the size of it can still be too big to have a better chance of finding data in the LLC.

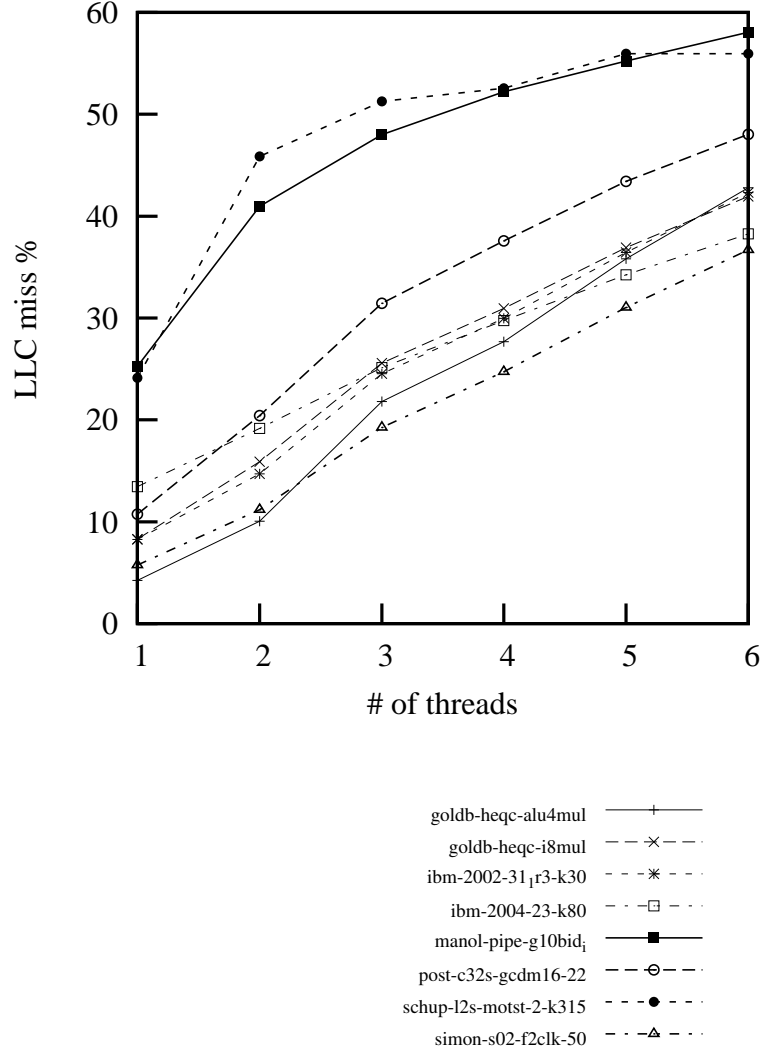


Figure 3.13: SharedAllLimited experiment

We also did an extra experiment with a new solver version, which only shares the initial clauses read from the input file, and then keeps separate clause databases for the new learnt clauses. We called this new version SharedAllLimited and Figure 3.13 shows the results of such experiment. We can observe that the results improve

compared to SharedAll and we end up getting lower LLC miss percentage, but similar numbers to the SharedNone and SharedBinaries versions.

3.4.4 Summing it all up

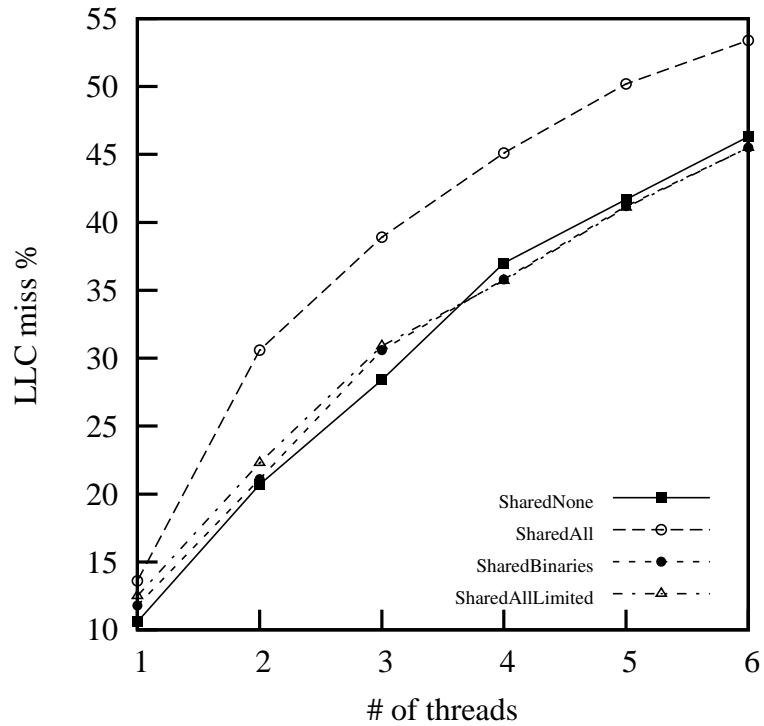


Figure 3.14: Average graph

To get a clearer picture of our results, we have plotted the average LLC miss percentages of all problems for all versions. Figure 3.14 compares all versions averages in a single graph. We can clearly see that SharedAllLimited and SharedBinaries

perform very similar and slightly better than SharedNone. This is because sharing some information, as little as it might be, does help with LLC miss percentages and does not modify the propagation scheme in any way. On the other hand, SharedAll has the worst performance, because, as we mentioned earlier, the gains of sharing clauses physically do not surpass the harms of modifying the propagation scheme and keep additional information for it.

Chapter 4

Conclusions

Throughout this work we have shown that algorithm design might not always be oblivious to modern computer architecture. Memory hierarchy plays an important role in the performance of both single and multi threaded applications. From our small experiments that deal with common cache problems, we can conclude that we must know how modern memory architecture works to get the best performance out of our programs. This is no different in the SAT solving world, where it is already known that cache plays an essential role in the efficiency of state-of-the-art solvers. Although cache performance has been widely studied for sequential SAT solvers (mainly studies about the two-watched-literal scheme [17]), it is yet not clear how it affects the design of parallel SAT solvers, which was the main focus of this work. We were able to prove that it is not possible to just keep adding threads to state-of-the-art parallel SAT solvers and expect to keep performing better. As we add more threads there is a considerable performance degradation of all threads, degradation that might get so high as to prove fruitless to keep adding threads.

It is also clear by now that sharing information between threads in a parallel program does help with cache performance, because of the higher chance to find data in the cache, but it is also equally important that the drawbacks of the modifications, introduced to share information, do not overwhelm the benefits. For instance, sharing information between threads will usually lead to the implementation of some type of lock, specially if the information is being modified, and locks can have a noticeable negative impact in overall performance. In our work this was not the case, but when sharing all clauses we stumbled upon the problem of having to modify the usual scheme of some of the processes involved in SAT solving. This is the reason why making multi-threaded applications is so difficult when the application is complex. There is no general rule of thumb or recipe that will work for all domains, each

program has its own complexities and specific constraints that will require detailed study and experimentation in order to achieve a better parallel version.

In the domain of parallel SAT solving, it is clear that sharing clauses among threads is a good idea and improves the search across all threads, but it is not clear which is the best way to do so. Some solvers, as we mentioned, share them through message passing and others would rather share them physically, but no real analysis of the performance of both approaches has been done. In this work we have used the approach used by **MiraXT** to share clauses physically, but our experiments and cache studies show that this is not a good idea. It is much more simple, straight forward and better to just let each thread manage its own separate clause database and avoid all the difficulties of physical sharing, which at the end only show to harm the overall cache performance of the solver. The only exception would be to physically share the unary and binary clauses, because these kind of clauses can easily be shared with no complications and significant overhead, while improving cache performance by a small margin.

For future work we would like to perform a statistical analysis with state-of-the-art solvers, to measure the optimal number of threads that these solvers should run in one chip, so that degradation in cache performance will not overshadow the benefits of having an extra thread. We would also like to implement versions of **AzuDICI** which share data through message passing, not physically, and compare them to the versions from this work. Another idea we would like to explore is just keeping a fixed amount of running threads, and switch between different solvers among these threads. As we mentioned before, the problem of performance in parallel solvers becomes evident when the amount of data of all threads becomes too big for the LLC performance to be efficient. So to solve this issue we could just keep a low amount of threads and arbitrarily assign running time to each solver among these threads. For example, if we wanted to run 10 worker threads in a 10 core machine, we could infer that 10 threads will generate too much data for cache to be efficient, so we could only run 6 threads with just 5 instances of the solver for a certain period of time, then switch some threads to run the other 5 instances, and keep switching between them. By doing this we would always keep a good cache performance because we would never

have more than 6 clause databases being used at the same time. The drawback would be that the switching of the different solver instances would probably cost some time (because we will need to load different clause databases and solving models each time we switch), but we would need to run experiments to come to a conclusion on whether this is a good idea or not.

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