

Query Transformations for Improving the Efficiency of ILP Systems

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Editors: James Cussens and Alan M. Frisch

Abstract

Relatively simple transformations can speed up the execution of queries for data mining considerably. While some ILP systems use such transformations, relatively little is known about them or how they relate to each other. This paper describes a number of such transformations. Not all of them are novel, but there have been no studies comparing their efficacy. The main contributions of the paper are: (a) it clarifies the relationship between the transformations; (b) it contains an empirical study of what can be gained by applying the transformations; and (c) it provides some guidance on the kinds of problems that are likely to benefit from the transformations.

Keywords: Inductive Logic Programming, Program Analysis

1. Introduction

Since 1991 Inductive Logic Programming (ILP) systems have been used to construct predictive models for data drawn from diverse domains. These include the sciences (King et al., 1992, 1996; Muggleton et al., 1992), engineering (Feng, 1992; Dolšak and Muggleton, 1992), language processing (Zelle and Mooney, 1993; Cussens, 1997), environment monitoring (Džeroski et al., 1994), and software analysis (Bratko and Grobelnik, 1993).

Like many other algorithms in the field of machine learning, ILP algorithms construct “hypotheses” for data by performing a search through a large space. Such a search typically involves generating and then testing the quality of candidates. Testing a candidate hypothesis involves executing a query in first-order logic against the data. Improvements in efficiency thus result from (A) minimizing, to the best extent possible, the number of candidates generated; and (B) maximizing, to the best extent possible, the efficiency of testing each candidate.

Problem (A)—minimizing the search space—has received a lot of attention in ILP research. This has resulted in the development of powerful language bias specification mechanisms that limit the size of the search space (Nédellec *et al.*, 1996) the study of refinement operators that allow one to efficiently navigate through a hypothesis space (van der Laag and Nienhuys-Cheng, 1998), etc.

Problem (B)—efficient testing of candidate hypotheses—does not appear to have received as much attention. It is the less pressing problem: however, in the light of the advances that have been made on techniques for minimizing the search space, the issue does become important. There has been some previous work of a stochastic nature (Srinivasan, 1999; Sebag and Rouveirol, 1997). These reduce the evaluation effort at the cost of being correct only with high probability. This paper is concerned with exact transformations, extending the work first reported by Santos Costa *et al.* (2000). In that paper, the authors illustrated that query execution was a very high percentage of total running time. They also proposed two simple transformations that converted a first order query into an equivalent one which was more efficient to execute. Empirical evidence presented suggested that under some conditions the transformations could significantly improve the efficiency of an ILP system—there, the system Aleph (Srinivasan, 2001). Here, we extend this work in the following ways: (1) we consider two further transformations; (2) we describe how the four transformations relate to each other; (3) we present a theoretical analysis of the transformations; and (4) we provide empirical evidence of the behaviour of the transformations on artificial and real-world problems.

The paper is organised as follows: in Section 2 we outline a simple procedure that serves as a model for many ILP algorithms. Section 3 describes the four transformations. Section 4 provides an analysis of the computational complexity of the transformations as well as the execution of the original and transformed queries. An empirical evaluation of the approach is presented in Section 5. Section 6 concludes the paper. The paper is accompanied by two appendices containing material relevant to Sections 3 and 5.

2. A Simple ILP Implementation

We adopt the following informal prescription for an ILP algorithm, based on Muggleton (1994). We are given: (a) background knowledge B in the form of a Prolog program; (b) some language specification \mathcal{L} describing the hypothesis space; (c) an optional set of constraints I on acceptable hypotheses; and (d) a finite set of examples E such that none of the elements of set E are derivable from B . The set is divided into a set of positive examples E^+ and a set of negative examples E^- . The task is to find some hypothesis $H \in \mathcal{L}$ such that (1) H is consistent with the I ; (2) the E^+ are derivable from $B \cup H$; and (3) the E^- are not derivable from $B \cup H$.

Several prominent ILP systems such as CProgol (Muggleton, 1995), Golem (Muggleton and Feng, 1990), and FOIL (Quinlan, 1990) use a simple iterative procedure to construct the hypothesis H one clause at a time. Figure 1 shows one such procedure, reproduced with minor changes from Srinivasan (1999). We refer the reader to this paper for proofs of correctness and complexity arguments. Here we will assume the following:

generalise(B, I, \mathcal{L}, E): Given background knowledge B , hypothesis constraints I , a finite training set $E = E^+ \cup E^-$, returns a hypothesis H in \mathcal{L} such that $B \cup H$ derives the E^+ .

1. $i = 0$
2. $E_i^+ = E^+, H_i = \emptyset$
3. while $E_i^+ \neq \emptyset$ do
 - (a) increment i
 - (b) $Train_i = E_{i-1}^+ \cup E^-$
 - (c) $D_i = search(B, H_{i-1}, I, \mathcal{L}, Train_i)$
 - (d) $H_i = H_{i-1} \cup \{D_i\}$
 - (e) $E_p = \{e_p : e_p \in E_{i-1}^+ .s.t. B \cup H_i \models \{e_p\}\}$.
 - (f) $E_i^+ = E_{i-1}^+ \setminus E_p$
4. return H_i

Figure 1: A simple ILP implementation. The function *search*() is some search procedure that returns one clause after evaluating alternatives in some search-space of clauses, all of which are in \mathcal{L} .

- A1. On each iteration, *search*() will examine a finite number of clauses;
- A2. For each clause examined, *search*() will check (at least) the derivability of each example in $Train_i$;
- A3. The check for derivability always terminates; and
- A4. The actual proofs of derivability of an example are irrelevant to evaluating the ‘goodness’ of a clause.

These assumptions hold not only for ILP systems that use an algorithm similar to the one in Figure 1, but for a much broader range of relational learning systems, including systems that induce trees, e.g. SRT (Kramer, 1996), instance based learners e.g. RIBL (Emde and Wettschereck, 1996) etc. Our results are equally relevant for those approaches. The transformations we propose in the next section are concerned with improving the efficiency of derivation checks arising from Assumption A2.

3. Four Query Transformations

All the transformations we study are ‘correct’ in the following sense: if a clause D_i is transformed into a clause D'_i , then $B \cup H_{i-1} \cup \{D'_i\}$ derives exactly the same examples in $Train_i$ as $B \cup H_{i-1} \cup \{D_i\}$. If Assumption A4 holds, the identification of the best clause by *search*() will then be unaffected.

An implementation (in Prolog) of all the transformations described here is available on request from any of the authors. The following sections will use terminology from logic programming (Lloyd, 1987) and from global analysis (Cousot and Cousot, 1992).

3.1 The Theta-transformation t_θ

We are concerned here with clauses of the form $Head : -Body$ where $Body$ is a conjunction of goals G_1, \dots, G_n (in the Prolog sense, or literals in the more traditional logical sense). Each of the G_i are meta-calls to user-defined or built-in predicates. In general, let N be an upper-bound on the number of solutions to any of the G_i . It follows straightforwardly that in the worst-case, the number of meta-calls is $O(N^n)$. Lowering N requires a re-appraisal of predicate definitions used by clauses (either by re-encoding of some predicates, or by removing them entirely). We do not pursue this here, and concentrate instead on lowering the value of n .

A straightforward reduction in the number of goals being tested (thus lowering n) is achieved by eliminating obviously redundant ones. A simple logical check for this during the execution of the steps described in Figure 1 is as follows. On iteration i of the procedure in Figure 1, let P denote $B \cup H_{i-1}$. Let C be a clause being examined in a search. Then, a literal l in C is redundant iff $P \cup \{C\}$ is logically equivalent with $P \cup \{C'\}$ where $C' = C - \{l\}$.

From this, it is easy to show that l is redundant iff $P \cup \{C\} \models \{C'\}$. This constraint, while providing an exact test for literal-redundancy, is expensive to implement (with a resolution-based theorem prover like that in YAP, it requires checking for the proof of inconsistency from the set $P \cup \{C\} \cup \{\neg C'\}$). Instead, we will employ a simpler test based on the subsumption relation (Nienhuys-Cheng and De Wolf, 1997). Recall that C subsumes C' iff there is some substitution θ such that $C\theta \subseteq C'$ and if C subsumes C' then $\{C\} \models \{C'\}$. Clearly therefore if C subsumes C' then $P \cup \{C\} \models \{C'\}$ and l is redundant (the reverse is not true, and hence we call it a “weak” test for redundancy).

We call the transformation that removes a set of redundant goals from a clause the *Theta-transformation*. It is evident that a transformation that removes redundant goals is correct. We use an approximation of the clausal-subsumption test (see Section 4) that is relatively inexpensive to perform and retains the property of correctness.¹ While this is a simple transformation technique, it is not clear from descriptions of existing ILP systems whether they actually employ it during the search.² In general, while the efficacy of the transformation will depend on the enumeration operator employed by the search function (that is, whether it is prone to introduction of redundant literals), we expect it to be more useful when the background predicates are non-determinate.

3.2 The Cut-transformation t_l

We observe that when executing a conjunction of goals G_1, \dots, G_n , failure of a goal G_i will result in attempting to generate more solutions for goals earlier in the sequence. This effort is useless if these solutions do not alter the computation of G_i . The *cut-transformation*, t_l , exploits the notion of goal independence by partitioning the set of goals in a clause into classes such that goals in different classes are independent. We will execute each class in sequence, and use the pruning operator ! to avoid any backtracking between classes.

In pure logic programs, goals depend on each other because they share variables. Given a function $vars(T)$ that returns all variables in the term T , two goals G_i and G_j are said to *share*, that is the relation $Shares(G_i, G_j)$ holds, when:

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1. Significant work has been spent on correct and efficient implementations of subsumption tests (Kietz and Lübke, 1994; Scheffer et al., 1996), but for the goals of this paper our simple implementation is good enough.
 2. Programs like WARMR (Dehaspe and Toivonen, 1999) perform extensive checks to ensure that clauses subsumed by others that have previously been shown to be “useless” are not examined further. This is more elaborate than the proposal here, which simply checks for redundant literals within a clause.

$$i = j \vee \text{vars}(G_i) \cap \text{vars}(G_j) \neq \emptyset$$

The definition ensures that the relation *Shares* is reflexive and symmetric. Its transitive closure *Linked*, defined as the smallest transitive relation that is a superset of *Shares*, is reflexive, symmetric and transitive, and therefore an equivalence relation.

Given a set of goals $\mathcal{G} = \{G_1, \dots, G_n\}$, we shall name the equivalence classes established by *Linked* as I_1, \dots, I_m . If two goals are in the same equivalence class, we will say they may be *dependent*, otherwise we will say they are *independent*. We would like to divide the original clause into several conjunctions of independent goals and execute them separately.³ To do so, we place the goals in each class I_i in a conjunction G_i . Our notion of dependence is a safe approximation of the dependencies that can possibly occur at run-time, because as soon as two goals have a variable in common they belong to the same equivalence class. Moreover, the computation of the equivalence classes is efficient.

In this approximation all goals that include a head variable or that share variables with one such goal will belong to the same independence class. Often, we know beforehand that head variables will be grounded before calling the body. Clearly, such variables cannot introduce sharing. To take advantage of this extra information, we classify variables as either *Grounded* or *PossiblySharing*, and define $\text{vars}(T)$ to return all *PossiblySharing* variables in T . More sophisticated analysis is possible, as discussed in Appendix A.

To effect t_1 it is sufficient to implement the following procedure:

1. Given the original clause:

$$H :- G_1, \dots, G_i, \dots, G_n.$$

classify all variables in H, G_1, \dots, G_n as *Grounded* or *PossiblySharing* and compute the equivalence classes for the (approximated) sharing relation.

2. Number the goal literals according to the equivalence class they belong to:

$$H :- G_{1j}, \dots, G_{ik}, \dots, G_{nl}.$$

3. Reorder the literals in the clause according to the class they belong to:

$$H :- G_{a1}, \dots, G_{b1}, \dots, G_{cm}, \dots, G_{dm}.$$

4. We are interested in any solution, if one exists. Thus we need to compute every class once and if a class has no solution, the computation should wholly fail. The following program transformation uses cut to guarantee such a computation:

$$H :- G_{a1}, \dots, G_{b1}, !, \dots, !, G_{cm}, \dots, G_{dm}.$$

3. McCreath's LIME system (McCreath, 1999) works by putting together conjunctions of what he calls 'simple clauses'. These correspond to our equivalence classes. LIME is however restricted to dealing with determinate ground back-ground knowledge.

The transformation is correct in the sense that the examples derivable before and after the transformation are the same. Step (3) is correct because the switching lemma allows us to reorder goals. Step (4) is correct because whenever we introduce a new cut, all goals before the cut are independent of all goals after the cut. Backtracking to before the cut therefore could never result in new solutions for the goals after the cut. A more complete discussion is given by Santos Costa *et al.* (2000).

3.3 The Once-transformation t_o

The cut transformation just described ensures that the search for each independent set of subgoals always stops at finding the first solution. Notice that the transformed body goal $q_1,!,q_2,!,\dots,q_n$ can also be written as $once(q_1),once(q_2),\dots,once(q_n)$ where the meta-predicate *once* is defined as

$$once(X):- X, !.$$

While at the clause level the *once* constructs have the same effect as the cuts, they have the advantage that they can be nested, whereas cuts cannot.

The objective is to fine-tune t_1 by processing each set of independent goals in more detail. We start from the observation that the data guiding the successful application of the cut-transformation was that certain variables are known not to share at runtime. We can improve the partitioning process further by using extra data for each non-head variable and by taking advantage of Prolog's left-to-right selection function. More precisely, we shall use prior knowledge on whether a literal grounds some of its arguments, and on whether a literal cannot cause its arguments to share by considering the status of each variable, as we move from goal to goal. This will allow us to transform each independent set of goals by considering the variables grounded by the first literal just like the ground head variables in the cut transformation. We then apply the transformation recursively and refine each independent set of subgoals into subsets.

Example 1 Consider the query $p(X,Y,Z), q(X,Y,U), q(X,Z,V)$. If p grounds X and cannot cause Y and Z to share, then $q(X,Y,U)$ and $q(X,Z,V)$ do not share, and after the call to p they can be executed independently.

We therefore 'look inside' each equivalence class returned by the t_1 . First, we find a prefix of one or more literals such that each literal will ground some of its arguments and it is known not to cause sharing among the others, then we apply the cut transformation on the rest of this subgoal, treating the grounded variables just like ground head variables.

We call this transformation the *once-transformation* or t_o . Blockeel *et al.* (2000b) describe two different versions of t_o . The dynamic version transforms queries during their execution, which results in an overhead but which also makes it possible to check groundness and sharing during execution instead of having to pre-compute a safe approximation. However, we only present the static version here.

For the static version, there is the open issue of how to estimate which literals ground or cause sharing between which arguments. It is reasonable to assume that such information is provided either by the user or through analysis. For instance, in many ILP data sets most predicates are defined by ground facts or range-restricted clauses only; such predicates always ground all their arguments (and hence do not cause sharing). However, a more sophisticated analysis is also possible (see Appendix A).

once-transform(q):
 let $q = l_1, l_2, \dots, l_n$
 find the smallest k s.t. there exists a partition \mathcal{P} of $\{l_{k+1}, \dots, l_n\}$ s. t. $\forall q_i, q_j \in \mathcal{P}$:
 $V(q_i) \cap V(q_j) \subseteq \text{GroundedVars}(\{l_1, \dots, l_k\})$ and
 $\forall v \in V(q_i), w \in V(q_j) : \{v, w\} \notin \text{PossiblySharing}(\{l_1, \dots, l_k\})$
 for all $q_i \in \mathcal{P}$: once-transform(q_i)

Figure 2: The once-transformation algorithm. $\text{GroundedVars}(G)$ contains all variables grounded by a call to G . $\text{PossiblySharing}(G)$ is the set of all pairs of variables that may share after a call to G .

A high level description of the once-transformation algorithm is shown in Figure 2. Essentially, the algorithm finds a prefix of a conjunction such that this prefix grounds enough variables for the rest of the conjunction to contain independent subgoals; then it is called recursively on these subgoals. Note that, similar to the cut transformation, we assume that the order of independent groups of literals may be switched but the relative order of literals that are dependent of each other does not change.

3.4 The Smartcall-transformation t_s

The smartcall-transformation t_s , called thus for historical reasons, was originally described by Bloekel (1998). The basic idea is as follows. ILP systems typically generate a clause by adding literals to (or *refining*) a clause encountered earlier (its *parent*). Assume a clause $c = h \leftarrow p, q$ has as parent $h \leftarrow p$. The parent clause has been evaluated before, which means at some point it was computed which facts $h\theta$ are derived by the clause. If these θ are remembered, we can exploit this in two ways. Firstly, for the refined clause we need to check derivability of these facts only, because the facts derivable by the new clause are necessarily a subset of the examples derivable by its parent. Secondly, each derivability test can itself be simplified, and this is exactly what the smartcall-transformation does. Consider the equivalence classes I_i of c , as mentioned above. Each class can be checked independently of the others. For those I_i for which $I_i \subseteq p$, a solution must exist because there is one for the whole p . Hence, when testing the new clause with a head substitution θ , we only need to check those I_i that contain literals that were added during refinement.

Example 2 Consider a clause $P = p(X) :- a(X, Y), b(X, Z)$ to which during refinement a literal $c(X, U)$ is added, resulting in

$$Q = p(X) :- a(X, Y), b(X, Z), c(X, U).$$

We have

$$t_!(Q) = p(X) :- a(X, Y), !, b(X, Z), !, c(X, U).$$

If P is known to derive a specific fact $p(e)$, we know that the goals $a(e, Y)$ and $b(e, Z)$ must have solutions. Moreover, $c(e, U)$ can be checked independently from these goals. Hence, t_s can drop the first two equivalence classes, yielding $t_s(Q) = p(X) :- c(X, U)$.

If instead of $c(X, U)$ we add $c(X, Y)$, then we get

$$t_1(Q') = p(X) \text{ :- } a(X,Y), c(X,Y), !, b(X,Z).$$

Now $b(X,Z)$ can still be dropped, but $a(X,Y)$ cannot, because even when we know the goal $a(e,Y)$ has a solution, the conjunction $(a(e,Y), c(e,Y))$ may not have any.

The smartcall-transformation is correct in the following sense : *given that an example was derived by the parent clause*, it can be derived by the transformed clause if and only if it can be derived by the original clause. The emphasized condition need not be fulfilled for the other transformations. It also implies that in order to use the smartcall-transformation, a list of all the facts derived by the parent clause (its “coverage list”) needs to be available. This may not be feasible in all ILP systems; more specifically, when using a greedy (beam) search or iterative deepening, the number of clauses for which the coverage list needs to be kept would typically be small; but for exhaustive searches (best-first, A^* , ...) that keep a long queue of clauses to be refined, the number of clauses multiplied with the number of examples might become prohibitively large.

3.5 Composition of Transformations

One can combine transformations by applying them one after another to a query. For brevity we introduce the notation t_{ab} where $t_{ab}(c) = t_b(t_a(c))$.

Composition of transformations is not commutative, therefore we respectively have 12, 24 and 24 possibilities for composing 2, 3 or 4 of the transformations. We can reduce the number of possibilities by remarking that

- t_o implicitly performs t_1 , hence a combination of both is equivalent to t_o ;
- Since both t_θ and t_s remove literals, they should always be applied before t_1 or t_o . Indeed, removal of literals may cause equivalence classes to “fall apart”. Thus, $t_1(t_\theta(c))$ may have more cuts than $t_\theta(t_1(c))$. For an example, consider $p(X,U)$, $q(X,Y)$, $p(Y,V)$, $p(a,U)$, $q(a,b)$, $p(b,V)$.
- For the same reason, t_θ should come before t_s .

Hence when applying multiple transformations the optimal order is: first t_θ , next t_s , and then either t_1 or t_o . Figure 3 shows a lattice that contains all the ordered combinations of transformations that may be useful; the lattice is constructed according to the “... implicitly performs ...” relation.

4. Analysis of Transformations

The time that we take to test each candidate clause will have two components: the time we take to apply the transformation plus the time to execute the transformed query.

4.1 Transformation Complexity

We will use the following symbols in analysing the complexity of transforming a clause: v , the maximum number of variables in a literal; and N , the number of literals in the query.

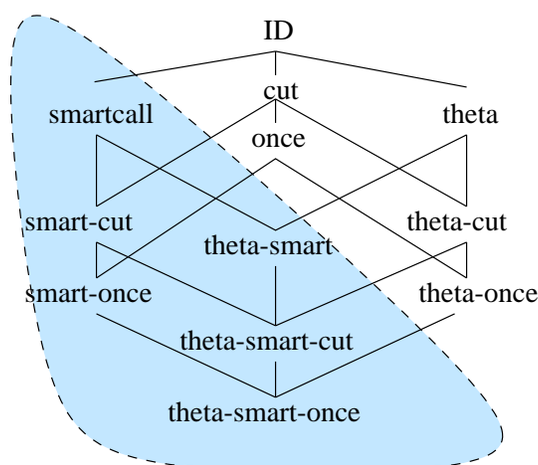


Figure 3: Lattice generated by the “...implicitly performs ...” relation among transformations. Transformations in the shaded area can only be applied under certain conditions.

4.1.1 COMPLEXITY OF t_θ

Our implementation of t_θ is a more efficient but safe approximation of the computation of a reduced clause under θ -subsumption (Nienhuys-Cheng and De Wolf, 1997). The transformation tries to remove each literal l_i in turn by doing a subsumption test to see whether the query without the given literal is equivalent to the original query q . This subsumption test is a polynomial time approximation of θ -subsumption. The algorithm computes the most general unifier θ of l_i and a literal of the remaining part q' . For each unification it skolemizes the original query $q\theta$ and q' . If $q\theta$ is a subset of q' then l_i is redundant and can be removed. The subset test in our implementation is $O(N^2)$ (but it could be done in $O(N \log N)$). The overall complexity of t_θ is then $O(N^4)$ (optimally $O(N^3 \log N)$) because the subset test is repeated for each possible θ and for each l_i .

4.1.2 COMPLEXITY OF t_l

The dominant component of t_l stems from calculating groups of dependent literals. The algorithm starts with an empty set of groups. In the i -th iteration it creates a new group containing the i -th literal in the clause, $\{l_i\}$, and merges this group with all existing groups that share variables with l_i . Testing if a group j shares variables with l_i is $O(n_j v)$ with n_j the number of literals in the group. This linear intersection test is possible because the algorithm keeps a sorted list of variables for each group. Performing the intersection test on all groups is $O(iv)$ because $\sum_j n_j = i - 1$. Now assume g_i groups have been identified that share with $\{l_i\}$; these have to be merged. Merging one group is $O(iv)$, so merging g_i groups is $O(g_i \cdot iv)$. The total complexity of t_l is $T(N) = O(\sum_{i=1}^N iv + g_i \cdot iv) = O(N^2 v + N v \cdot \sum_{i=1}^N g_i) = O(N^2 v)$. The last step is possible because $\sum_{i=1}^N g_i < N$.

4.1.3 COMPLEXITY OF t_o

The implementation of t_o first finds independent groups of literals using the same algorithm as the cut-transformation. If all literals are in the same group then the algorithm grounds the variables of

the first literal and calls itself recursively for the other literals. If the literals are partitioned into several groups then the algorithm does a recursive call for each group. In the first case the time complexity $T(N) = O(N^2\nu) + T(N - 1)$ with ν the maximum number of variables in a literal and N the number of literals in the query. If each recursive call is of this type, then the total complexity is $O(N^3\nu)$. In the second case $T(N) = O(N^2\nu) + \sum_{i=1}^{g_N} T(n_i)$ with $g_N > 1$ the number of subgroups and n_i the number of literals in subgroup i . It can be shown that the worst case is $g_N = 2$, $n_1 = 1$ and $n_2 = N - 1$.⁴ The overall complexity is $O(N^3\nu)$ for both cases.

4.1.4 COMPLEXITY OF t_s

The t_s transformation splits the query $q = q^{prev} \wedge q^{new}$ in a part q^{prev} that is known to succeed and the new part q^{new} . It first partitions q^{prev} into independent groups using the same algorithm as the cut transformation and then removes all groups that do not share variables with q^{new} . The partitioning into dependent groups is dominant, hence giving Smartcall the same complexity as the cut transformation.

All transformations are thus polynomial in the length of the clause. Note that while the transformation overhead can grow large for long clauses, evaluation time for the untransformed clause can be exponential in N , so that the net gain obtained by transforming the clause can be expected to increase with N .

4.2 Execution Complexity

We now examine the complexity arising from executing a transformed query with a view to estimating the gain in efficiency obtained from performing a transformation. We further consider estimates of average gains only, as some of the transformations cannot always guarantee efficiency gain. We will use the following notation:

- q represents a query with literals l_1, l_2, \dots, l_N
- N is the number of literals in the query (as before)
- e is the number of examples
- b is the average branching factor of the SLD tree of the query and can be seen as a measure of non-determinacy of the data
- d is the depth of the SLD tree
- q_i and m are defined by $t!(q) = q_1, !, q_2, !, \dots, q_m, !$; we call the q_i independent subgoals
- d_i is the depth of the SLD tree generated by q_i
- N_i is the number of literals in q_i

4. To see this, note that $T(N - i) + T(i) < T(N - 1) + T(1) < T(N)$ when T is a superlinear function and $1 < i < N$. In our case T has to be superlinear because it contains an N^2 term. The demonstrandum follows by recursively applying these inequalities on the terms in $\sum_i T(n_i)$.

4.2.1 COMPLEXITY OF t_θ

The effect of t_θ is just to reduce the length of a query. Since the cost of evaluating a query is exponential in its length, it may yield significant results, even when only a single literal is removed. If the removed literal is non-determinate and may succeed k times, then the SLD-tree for the query will be reduced by a factor k . If it is determinate, we still save a number of meta-calls during the execution of the query. Profiling results reported by Santos Costa et al. (2000) suggest that this will be relevant in practice, although the effect will be most noticeable if the removed literal was at the end of the clause.

4.2.2 COMPLEXITY OF t_l

We approach the efficiency gain yielded by t_l from two points of view: first, looking at SLD-trees and non-determinacy; second, at a higher abstraction level, looking at the execution times the subgoals consume.

Let us assume for now that a literal l_i succeeds $k_i > 0$ times. Then without the transformation the number of nodes in the SLD tree is $\prod_i k_i$. After the transformation it becomes $\sum_i \prod_{l_j \in q_i} k_j$. If we simplify this by assuming a constant branching factor b , execution of the original query takes time $O(b^N)$ while execution of the transformed query takes time $O(b^{\max N_i})$. Thus, the execution time of the query is reduced from “exponential in its length” to “exponential in the length of the longest conjunction between two cuts”.

We can obtain some more insights from a second stance, describing efficiency in terms of times rather than SLD tree sizes. We introduce

- t_i : the time needed for exhausting the search space of q_i
- s_i : the number of times q_i succeeds
- $\bar{t}_i = t_i / (s_i + 1)$: the average time until success or failure for subgoal q_i
- $k = \min\{i | s_i = 0\}$ (i.e. q_k is the first subgoal that has no solutions)

Without cuts the time needed to confirm failure of the clause on a single example is

$$t_1 + (s_1(t_2 + s_2(\dots))) = t_1 + s_1 t_2 + \dots + s_1 \dots s_{k-1} t_k$$

and after applying the cut-transformation this becomes

$$\bar{t}_1 + \bar{t}_2 + \dots + \bar{t}_{k-1} + t_k$$

If the last term in both formulae is dominant (which happens if t_k is large), a speedup of at most $s_1 s_2 \dots s_{k-1}$ can be obtained. If an earlier t_j dominates, this results in a speedup of roughly $s_1 s_2 \dots s_{j-1}$.

4.2.3 COMPLEXITY OF t_o

The once-transformation, t_o , is the recursive application of t_l , and thus has a similar effect on efficiency; only now the original equivalence classes may be subdivided further. The execution time of the transformed query is most easily described by introducing the concept of a once-tree. A once-tree is constructed from a once-transformed query as follows: the root of the tree contains the

Transformation	Evaluation Complexity
ID	$O(eb^d)$
t_θ	$O(eb^{d'})$ with $d' \leq d$
$t_!$	$O(eb^{\max d_i})$
t_o	$O(eb^{\max d'_i})$ with $\forall i : d'_i \leq d_i$
t_s	$O(e'b^{d_m})$ with $e' \leq e$

Table 1: Summary of efficiency of evaluation of transformed queries.

part of the query that is not inside once-constructs, and for each once construct it has a child which is the once-tree of the subgoal inside the once. The execution time of the transformed query is then exponential in the length of the longest path from the root of the tree to a leaf, where length is interpreted as the total number of literals on the path.

4.2.4 COMPLEXITY OF t_s

The smartcall transformation consists of dropping all subgoals that are independent of the last added literals. This makes the execution time exponential in the length of the remaining subgoals. Table 1 presents an overview of the results in terms of non-determinacy (represented by the average branching factor b).

The effect of combining transformations is a possible further reduction of the d parameters. When $t_!$ or t_o are applied after t_θ , the resulting d_i and d'_i parameters may be less (but not greater) than if t_θ had not been called in advance. Applying t_o after t_s reduces the b^{d_m} factor to $b^{\max d'_i}$ with d'_i the maximal number of literals on a path from root to leaf in the once-tree of $t_s(q)$.

4.3 Summary

The results on computational complexity can be summarized as follows:

- As the transformation time itself does not depend on the size of the data set, for large data sets it will become negligible compared to the execution time gained by performing the transformation.
- Concerning the evaluation of clauses, the efficiency gain the transformations yield depends on the size of the SLD tree of the original and transformed clause. This size is affected by:
 - the number of examples, which determines the number of head variable substitutions for the clauses
 - the non-determinacy of the literals in the clause, which measures the branching factor in the SLD tree
 - the depth of the SLD tree, which can be considered proportional to the length of the clause for the untransformed query; for the transformed query it is proportional to the length of the longest path in its once-tree.

This yields four parameters that influence efficiency gain: number of examples, non-determinacy, length of clauses, and complexity of the most complex subgoals (as measured by the once-tree). A

	a	b	c
Size	100–500	501–2000	2001–10000
Non-determinacy]1, 2]]2, 3]]3, 6]
Length	1-4	5–8	9–12

Table 2: Discretization of controlled parameters into three categories.

maximal gain can be expected when the first three parameters have large values and the last one is small.

5. Empirical Evaluation

The analysis above gives us some insights into the behaviour of the transformations. In practice, we would like to know the context in which each transformation can be expected to work best. Although theory does not answer this question, it does provide us with the basis on which we can design experiments to provide guidance.

Our empirical evaluation is of two types: **(1)** controlled experiments using artificial data (Section 5.1) and **(2)** uncontrolled experiments using real-world data (Section 5.2). All experiments were run on a Linux PC (Pentium III, 850 MHz, 256 MB). We used Aleph 3 running on Yap 4.2.0; coverage lists were enabled throughout.

5.1 Controlled Experiments

The purpose of our controlled experiments is to investigate the influence of parameters on the efficiency of the different transformations. We estimate, as a consequence, the set of transformations best suited for each setting of the parameters.

5.1.1 MATERIALS AND METHOD

We control the following three parameters: number of examples, length of clauses and non-determinacy. The fourth parameter, complexity of the hardest independent subgoal, is more difficult to control, but can be measured.

We discretize the values of the controlled parameters into three categories: low, medium, high. The thresholds are shown in Table 2. The three categories for three different parameters give rise to 27 combinations. For each combination we generate a corresponding artificial data set and random clauses, then we run the transformations on the clauses and measure the relative performance of each transformation, to determine which one works best for that combination.

5.1.2 THE DATASETS

The artificial datasets are all directed graphs. Each graph contains a number of nodes e and each node has b_0 outgoing edges to a randomly chosen node of the graph. Each individual node serves as an example in the data set. By controlling b_0 and e we control size and nondeterminacy, respectively. A more precise description of how the data sets were generated is given in Appendix B.

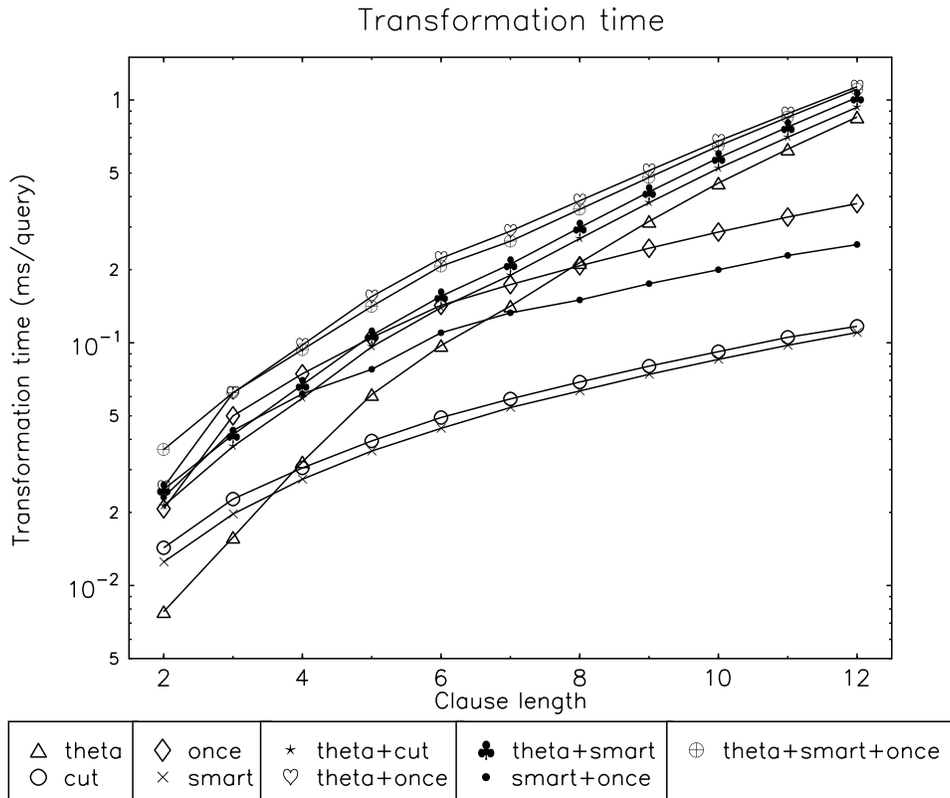


Figure 4: Relative efficiency of the transformations.

5.1.3 QUERY GENERATION

The clauses generated in this experiment can have two kinds of literals in the body: *edge/2* and *label/2* literals. The first argument of an edge literal is an input argument; the second argument is then non-determinate as it can take b_0 different values. *label/2* literals are always added with a constant as second argument and serve as tests (i.e., they may succeed or fail).

The random queries are generated in a levelwise top-down manner, as follows:

```

 $C_0 := \{ p(X) \leftarrow \}$ 
 $l := 1$ 
while  $l < \text{maxclauselength}$ 
   $C_l := n$  random refinements of random clauses from  $C$ 
  increment  $l$ 

```

Clauses are then selected at random from the C_l .

5.1.4 RESULTS

Some results on transformation times are shown in Figure 4. The results not only confirm that the different transformations have complexity N^2 , N^3 and N^4 (these are clearly distinguishable on the

figure), they also give an indication of the actual time they consume. If $T(t, l)$ is the time needed to execute transformation t on a clause of length l , and e is the number of examples, then $T(t, l)/e$ is the time we should at least gain when evaluating a query on a single example for the transformation to be useful.

Figure 5 summarizes results on evaluation times. For each individual clause nondeterminacy was estimated as the average branching factor $b = b_0^{e/l}$ with b_0 the branching factor in the graph; e the number of *edge* literals in the clause, l the length of the clause (since only *edge* literals have a nondeterminacy of b , *label* literals are determinate).

The continuous lines show the actual running times of the slowest and the fastest approach. The graph clearly shows that the longest running times are needed when nondeterminacy as well as clause length are high. The average time for querying one example does not depend much on the size of the data set (which means the total running time is linear in the number of examples). It is also clear that the transformations yield the highest efficiency gain where it counts most, and thus greatly reduce the variance of the timings: execution times for transformed clauses vary over a factor of about 20 instead of a factor of 1000.

Below the absolute measurements, relative speedups are shown; the 1.0 line corresponds to “no transformation”. In almost all cases, all transformations yield a speedup, sometimes a large one. The most complex composed transformation ($t_{\theta_{so}}$) consistently yields the greatest speedup.

As all three parameters vary along the same axis in this graph, the effect of each single parameter is a bit obscured. For instance, it is not apparent whether moving from a simple to a more complex setting always increases the speedup. The settings are partially ordered according to complexity. Figure 6 visualizes this partial ordering in a lattice. To keep the lattice simple the effect of data set size is not included; from Figure 5 it is quite clear that data set size does not significantly influence the speedup ratios. The numbers in the nodes of the lattice show the average speedup (averaged over data sets of varying size) for the $t_{\theta_{so}}$ transformation. It clearly indicates that the speedup indeed increases monotonically with complexity.

5.2 Uncontrolled Experiments

The purpose of the uncontrolled experiments is to examine the performance of the best transformations on particular data sets.

5.2.1 MATERIALS AND METHOD

Data here are from real-world problems or benchmarks. With these problems, the procedure in Figure 7 was adopted. In practice, adopting this procedure is trivial in the sense that for all categories, our experiments on artificial data indicate $t_{\theta_{so}}$ as the best transformation.

5.2.2 NOTE ON DATASETS

We used three datasets for these experiments, which have the following properties:

- Bongard (De Raedt and Van Laer, 1995): 1352 examples; nondeterminacy estimate: low
- Carcinogenesis (Srinivasan et al., 1999): 330 examples; nondeterminacy estimate: high
- Mutagenesis (Srinivasan et al., 1996): 188 examples; nondeterminacy estimate: high

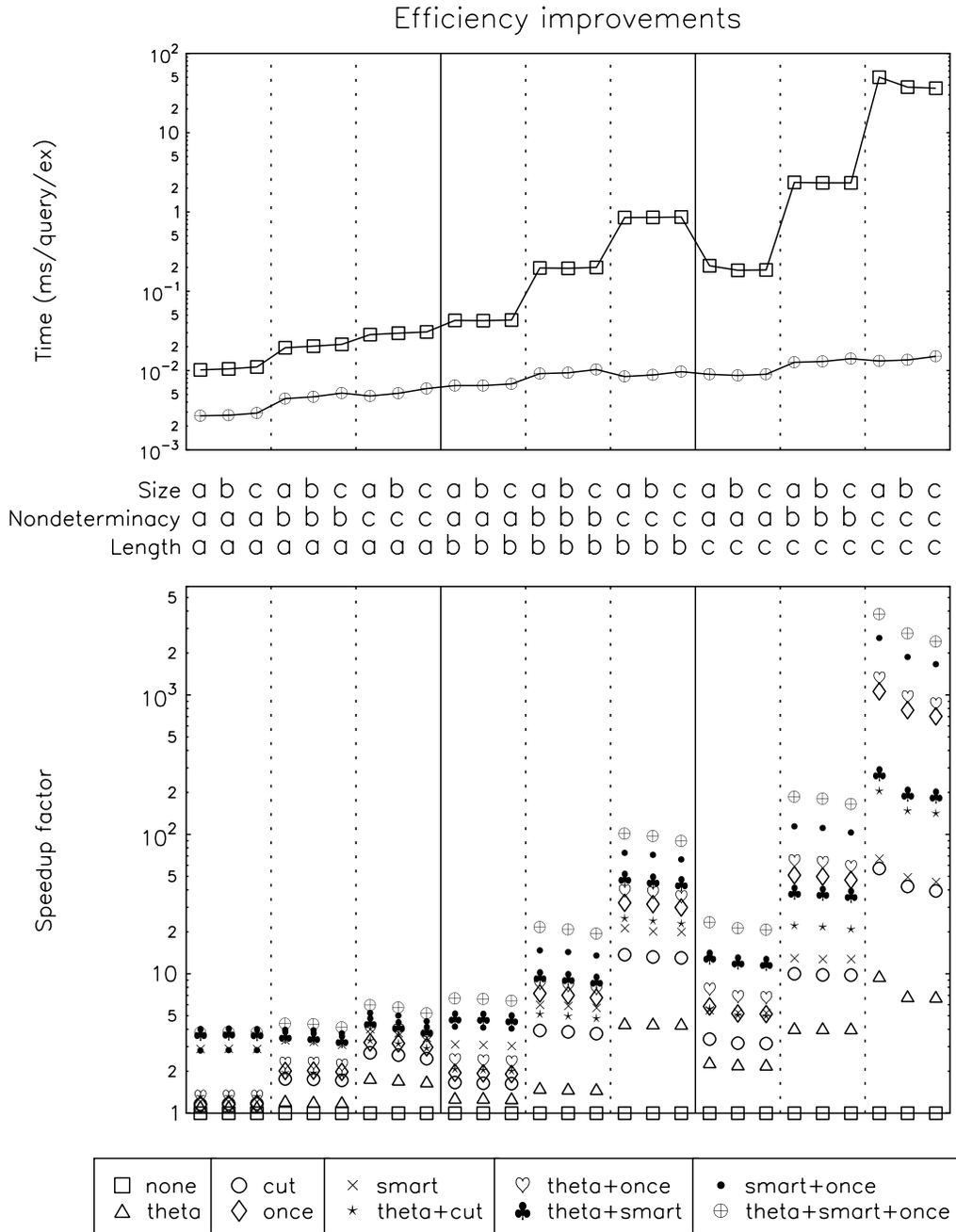


Figure 5: Efficiency improvements obtained with the different transformations. The top graph compares absolute timings for the untransformed clause with those obtained by applying the best transformation. The bottom graph shows for each transformation the relative speedups obtained with it in the different settings.

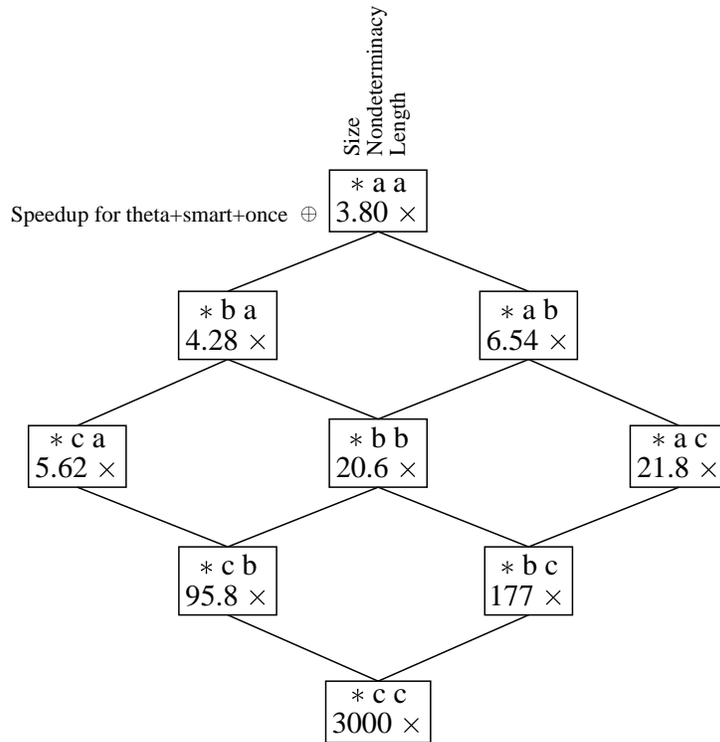


Figure 6: Different settings, partially ordered according to the “is more complex than” relation (excluding the size parameter). In the nodes, speedup factors for $t_{\theta,so}$ are shown.

1. Assign each problem to one of the 27 categories (obtained previously)
2. Choose the best transformation for the category from controlled experiments
3. Employ transformation and record results

Figure 7: Experimental method for uncontrolled experiments

Nondeterminacy estimates are based on the typical use of predicates that occur in the data; for instance, the molecules used in the Carcinogenesis dataset contain on average some 30 atoms; introducing a new *atom* literal with a free variable identifying the atom thus has a nondeterminacy factor of 30. However, there are also e.g. bond literals with lower nondeterminacy; assuming both occur equally frequently in a clause, it makes sense to estimate nondeterminacy as the geometric mean of all these nondeterminacy factors. Based on this reasoning and on previous experience with these datasets, we estimate nondeterminacy to be low for Bongard, and high for Carcinogenesis and Mutagenesis.

Average clause length is the most difficult parameter to estimate, as it does not follow from the data but depends on the complexity of the target theory. For Carcinogenesis, there is reason to believe that there are no good clauses of small length (Srinivasan et al., 1999), so an estimate of medium to high seems appropriate. For Mutagenesis, previous experience suggests that clause length can be expected to be low. For Bongard underlying theories of arbitrary complexity can be generated; here we varied the maximal clause length from low to high.

The categories are then as follows:

- Bongard: Size=medium, Nondeterminacy=low, Clause length = low-high
- Carcinogenesis: Size=low, Nondeterminacy=high, Clause length = medium-high
- Mutagenesis: Size=low, Nondeterminacy=high, Clause length = low

Our previous analysis predicts $t_{\theta_{so}}$ to work best on all three. It also suggests that the speedup factors for query execution that can be obtained with this transformation are: for Bongard, 5 to 20; for Carcinogenesis: 100 to 1000; and for Mutagenesis: 5. Aleph keeps coverage lists for its clauses and the smartcall transformation can be used. However since for other ILP systems it might not be possible to include smartcall in the transformation, it is instructive to examine the results obtained with t_{θ_0} .

5.2.3 RESULTS

Figures 8, 9 and 10 show the efficiency gains obtained on the Bongard, Carcinogenesis and Mutagenesis data sets. We separate the times consumed by the transformation, the execution of the query in the database, and other computations done by the learner. Note that on the artificial data sets we measured only the speedup of the query execution itself.

We now examine each problem in turn. On the Bongard dataset with small nondeterminacy, only little gain is achieved (Figure 8): even for relatively large clause lengths a speedup factor of about two is achieved, and for small clause lengths the overhead of performing the transformation is not compensated by faster query execution. It is also the case, however, that the overhead imposed by the query transformations is small (less than 5%).

The speedup factors are, however, not in line with our expectations. We observe a factor 1 to 2 speedup but 5 to 20 was predicted. To investigate this further, we measured the SLD-tree size for the Bongard experiments with length 6. This suggests a branching factor of 1.54 (solving for b in $n = b^l$ where n is the number of nodes in the SLD-tree and l the length of the query). With controlled data, clauses of maximum length 6 and a branching factor between 1.5 and 1.6 in fact exhibit an average speedup of only 3.8. Interestingly, on the Bongard data set the size reduction of the SLD-tree by the employed transformation was 3.3, which is close to the 3.8 obtained with

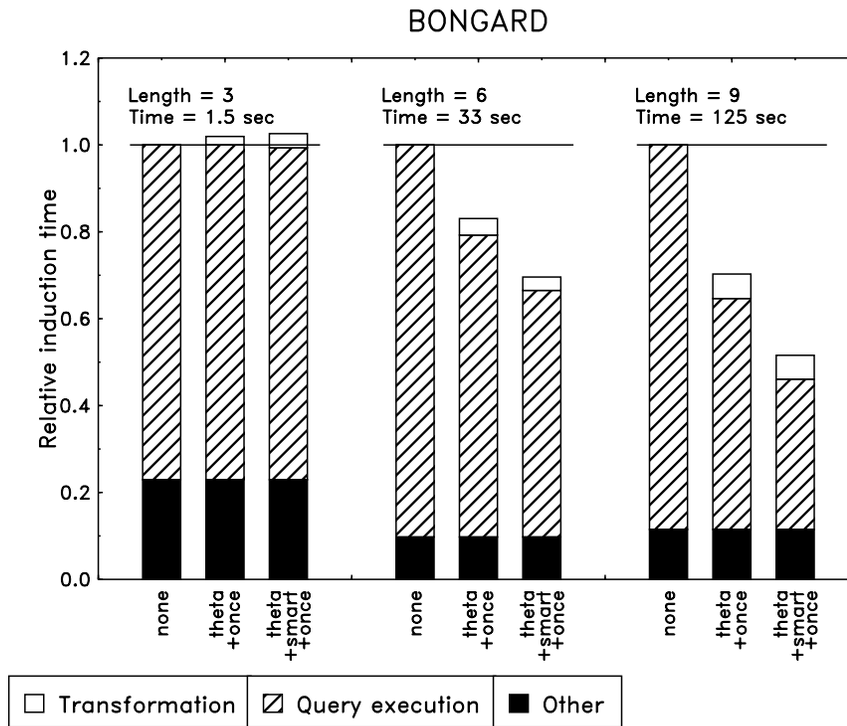


Figure 8: Efficiency improvements obtained on the Bongard data set.

controlled data. Apparently the reduction in SLD-tree size is as expected, but it is not matched by the execution speedup. We have, as yet, been unable to come up with a suitable explanation for this discrepancy.

The greatest gain is achieved in the highly nondeterminate Carcinogenesis dataset (Figure 9). When restricting the clause length to 6, Aleph’s total run time was reduced by two orders of magnitude, and the query execution time itself by a factor 200. When allowing a maximal clause length of 8, experimentally determining the speedup became impractical. Illustrative examples were still instructive: a certain point in the search space that with transformed clauses took 6 minutes to reach, was only reached after over three days with untransformed clauses. The whole run with transformed clauses took 2.53 hours, and our estimate of the speedup factor is necessarily very approximate. Nevertheless, these results seem consistent with our estimate of 10^2 to 10^3 . The Mutagenesis dataset (Figure 10) can be positioned in between. The query execution speedup is here around 4, which is clearly close to what was expected (5).

6. Conclusions

As ILP systems move from the realm of research to one of technology, implementation issues and concerns of efficiency become increasingly important. Here we have described a number of simple clause-transformation techniques that are directed towards reducing the theorem-proving effort that is at the heart of many ILP systems. Theoretical analysis of the transformations suggests that they can provide significant efficiency gains for problems that require complex theories that use highly

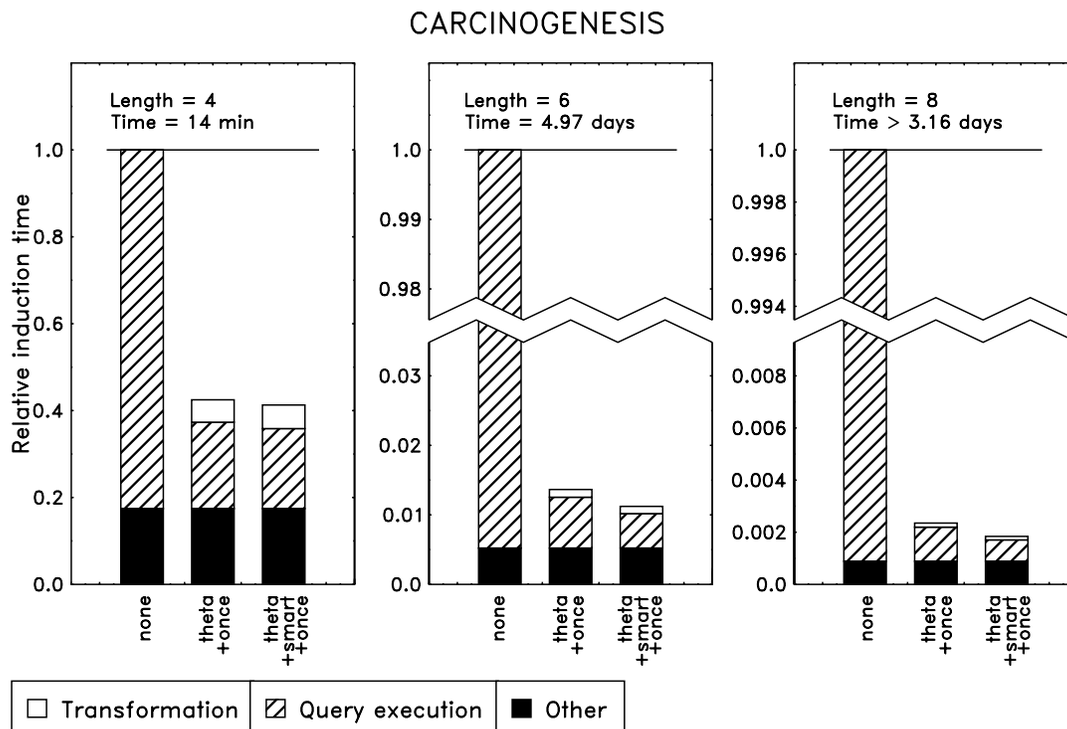


Figure 9: Efficiency improvements obtained on the Carcinogenesis data set.

non-determinate background predicates. We have conducted an empirical study that confirms this and moreover shows that the transformation overhead is sufficiently small so that even on moderately sized data sets with simple queries and low non-determinacy, there is no adverse effect of performing the transformations. It is therefore advisable to always use, in practice, the most sophisticated composition of transformations that is applicable. For systems that refine clauses in a top-down manner and remember the set of examples covered by the parent clause, this is the composition of the theta, smartcall and once transformations; for other systems the composition of the theta and once transformations is the best option.

The approach adopted in this paper is in the long tradition of source-to-source program transformations: changes at the source-level that can improve efficiency without altering correctness (Loveman, 1977). The suggestions here by no means exhaust the transformations of this type. Within ILP, a related approach is described by Blockeel et al. (2000a), where a set of queries is restructured so that they can be executed more efficiently, without changing the individual queries however. The two approaches are obviously complementary, and it would be interesting to see how they can be combined.

Other related work in ILP is that on efficient subsumption testing (Kietz and Lübke, 1994; Scheffer et al., 1996). First, a correct and efficient implementation of the subsumption test could only improve our theta-transformation. Second, some of the methods described in the literature on theta-subsumption are closely related to the methods described here. This is not so surprising given the close relationship between subsumption testing and executing a query; in both cases a mapping of one structure onto another is attempted. While our current approach is to transform clauses and

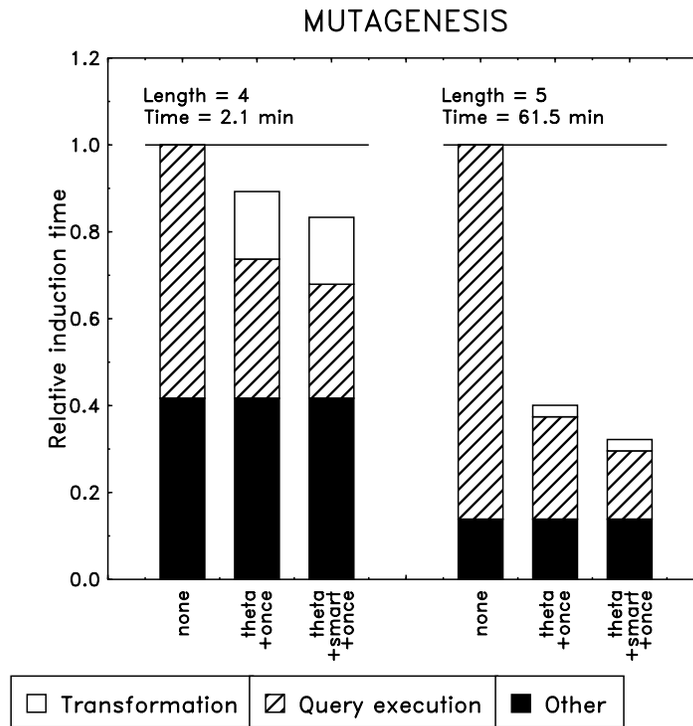


Figure 10: Efficiency improvements obtained on the Mutagenesis data set.

then employ a standard execution mechanism, some of the theta-subsumption optimisations might be useful for improving the execution strategy itself.

Finally, we remark that obtaining efficiency gains by means of transformations such as the ones described here is only one side of the story. Efficiency gains are as much to be made by tailoring compilers of logic programs to account for what ILP systems do. The best results regarding efficiency may well be obtained by combining both views, i.e., designing Prolog engines in conjunction with ILP algorithms.

Acknowledgments

This work has been partly supported by Fundação da Ciência e Tecnologia under the project Dolphin (PRAXIS/2 /2.1/TIT/1577/95), by the PROTEM5 CNPq-NSF Collaboration Project *CLOPⁿ*, and by the FAPERJ Project PLAG. A.S is supported by a Nuffield Trust Fellowship at Green College, Oxford. H.B is a post-doctoral fellow of the Fund for Scientific Research of Flanders (FWO-Vlaanderen). J.S is a research assistant of the Fund for Scientific Research of Flanders. During part of this research, H.V and W.V.L were funded by the FWO-Vlaanderen project G.0246.99, “Query languages for database mining”. We would like to thank reviewers of the paper for their detailed comments.

Appendix A. Program analysis and set sharing for determining (in)dependent calls

Program analysis aims at deriving at compile-time information about the execution of a program. Typically this information could also be given explicitly by the programmer, e.g. a declaration stating that the execution of a predicate grounds its arguments. In (constraint) logic programming researchers have been using abstract interpretation (Cousot and Cousot, 1992) as a general framework for program analysis: the concrete execution of a program is mimicked by using descriptions of the concrete substitutions, so-called *abstractions*. Much research effort has been put into the development of these abstractions, which have to be *safe* approximations of their concrete counterparts, but which also have to be *precise* enough to capture the properties of interest. Again groundness is a very well studied property. Also note that abstract interpretation computes these abstractions at all program points in a program, typically before and after each call such that the abstractions describe all possible concrete substitutions before and after a call.

For this paper we would like to point out the link with the **set sharing** abstraction (Jacobs and Langen, 1992; Bueno et al., 1994) which has been used to identify possibilities for independent AND-parallelism (Codish et al., 1997) and which can also be used in this context to refine the dependencies between the arguments of the calls. It is beyond the scope of this paper to give a detailed explanation of program analysis topics, but we will show in an informal way how program analysis can help in this context.

Taking into account all variables of the calls for the detection of dependency is a safe abstraction of the execution of the called predicate: execution of the predicate can possibly create dependencies between all the variables in the calls. This approach is compatible with the set sharing abstraction for a call for which no additional information is available and in which all variables are initially free independent variables—the latter is known as a goal-independent analysis. For the call $p(X,Y,Z)$ the set sharing abstraction is the powerset of the set of all its variables $\{X,Y,Z\}$, namely $\{\{\},\{X\},\{Y\},\{Z\},\{X,Y\},\{X,Z\},\{Y,Z\},\{X,Y,Z\}\}$. A subset in the abstraction describes the possibility that the variables in the subset have in their values variables in common. The above abstraction then describes for example the concrete substitution $\{X \leftarrow f(A,B), Y \leftarrow f(A,C), Z \leftarrow g(B,A)\}$. This concrete substitution is not described by the abstraction $\{\{X\},\{Y\},\{Z\},\{X,Y\},\{X,Z\},\{Y,Z\}\}$ as the sharing of A by $\{X,Y,Z\}$ is not allowed, nor by the abstraction $\{\{X\},\{Y\},\{Z\}\}$ as now no shared variables are allowed. The power set safely expresses that we have to assume all possible cases, as we do not know anything about what the call actually does to its free arguments. Reasoning with the largest set of dependent variables is safe as it describes the dependencies we are interested in.

Program analysis allows us to refine the dependency relation between the arguments of calls. A first case is the grounding of variables: as soon as a variable becomes ground it should no longer be considered for dependency determination. In the set sharing abstraction the ground variables are removed from the powerset. This is compatible with the treatment of ground variables. The point here is that program analysis derives this grounding behaviour by computing a more precise abstraction for grounding calls. If the call $p(X,Y,Z)$ grounds all its variables, the abstraction is $\{\}$. If only X is grounded, the abstraction is $\{\{Y\},\{Z\},\{Y,Z\}\}$ (which still allows all possible dependencies between Y and Z). Finally, if X and Y are both grounded, the abstraction becomes $\{\{Z\}\}$.

A further refinement is taking into account the (in)dependence of variables appearing in the same call which can be computed by means of the set sharing abstraction. This is what is needed in

the static version of the once-transformation. For a call $p(X,Y,Z)$ that only can create a dependency between Y and Z , the abstraction is $\{\{X\}, \{Y\}, \{Z\}, \{Y, Z\}\}$. Again, taking all variables of the call would be an overestimation: one can make a distinction between the calls depending on $\{X\}$ and the ones depending on $\{Y, Z\}$. Alternatively, one could for determining dependent calls view $p(X,Y,Z)$ as two independent calls $p_1(X)$ and $p_2(Y,Z)$ (or similarly as the conjunction $p(X,-,-), p(-,Y,Z)$ such that each p_3 call can be part of another set of dependent calls).

Our program analysis based approach could be organised along the following lines. First for all predicates (describing the examples and the background knowledge) a goal-independent set sharing analysis is done which can be used to approximate the dependency property of a call (also taking into account the independence of variables). Also for builtins, set sharing can compute an abstraction. Typically, builtins such as comparison impose that the variables are ground, so the set abstraction will be empty. For $X = Y$, the abstraction is $\{\{X,Y\}\}$. Note this has to be done only once.

Then we consider the calls in a query from left to right together with their dependency approximation (as derived from set sharing abstractions). We will also propagate the groundness information from left to right which will remove ground variables from the dependency approximations.⁵ During this left-to-right traversal we can determine the chains of dependent calls by computing transitive closures on the “grounded” dependency approximations.

Let us consider the following example query where $t(T)$ is the extension.

$p(X,Y), q(X,Z), \text{lof}(Z), r(Y,T), t(T).$

1. In the most general setting, all the calls have as dependency approximation their complete set of variables. Thus they all belong to the same (dependency) chain.
2. Suppose $p(X,Y)$ grounds both X and Y , then we have the following dependency approximation after propagating the groundness: $p(X,Y): \{\}, q(X,Z): \{Z\}, \text{lof}(Z): \{Z\}, r(Y,T): \{T\}, t(T): \{T\}$. Thus after the call $p(X,Y)$ there are two chains: $q(X,Z), \text{lof}(Z)$ and $r(Y,T), t(T)$. Note that backtracking over values for Z or T is not necessary as soon as a chain has succeeded once.
3. Suppose $p(X,Y)$ grounds neither X nor Y but X and Y remain independent, then the set abstraction is $\{\{X\}, \{Y\}\}$ and we could replace $p(X,Y)$ by the conjunction $p(X,-), p(-,Y)$. The dependency approximation then becomes, $p(X,-): \{X\}, p(-,Y): \{Y\}, q(X,Z): \{\{X,Z\}\}, \text{lof}(Z): \{\{Z\}\}, r(Y,T): \{\{Y,T\}\}, t(T): \{T\}$. We can identify two chains: $p(X,-), q(X,Z), \text{lof}(Z)$ and $p(-,Y), r(Y,T), t(T)$.
4. Suppose $p(X,Y)$ grounds only one variable, let us assume X . Then we have two chains: $q(X,Z), \text{lof}(Z)$ and $p(-,Y), r(Y,T), t(T)$.

Appendix B. Generation of Artificial Data Sets

Figure 11 describes the method used for generating artificial data sets and clauses for our first set of experiments. As explained in Section 5.1.2, the data sets are directed graphs containing a number of nodes s and for each node b_0 outgoing edges.

5. In abstract interpretation terms, a goal-dependent analysis could be performed based on the goal-independent analysis results.

```

for each length  $l$  from  $\{1 - 4, 5 - 8, 9 - 12\}$ 
   $Q = \text{sample } 30 \text{ clauses of length } l$ 
  for each branching factor  $b_0$  from  $\{1 - 3, 4 - 6, 7 - 9\}$ 
    for each size  $s$  from  $\{100 - 500, 501 - 2000, 2001 - 10000\}$ 
       $D = \text{generate\_problem}(s, b_0)$ 
      for each clause  $q \in Q$ 
        for each transformation  $t \in \mathcal{T}$ 
           $q' = \text{transform}(t, q)$ 
           $T = \text{time}(q', D)$ 
           $b = b_0^{e/l}$ , with  $e$  the number of edges in  $q$ 
           $\text{compute\_average\_time}(s, b', l, T)$ 

```

Figure 11: Experimental method for controlled experiments

The algorithm generates values for three parameters: the clause length l , the branching factor b_0 (i.e. the number of outgoing edges) and the size s . Each parameter can take values from three intervals: low, medium and high. The for-loops are iterated three times, each time selecting a random value from one of the intervals.

A set of clauses Q is generated for each length l . Likewise a random graph dataset D is generated based on the values of s and b_0 . The two most inner loops apply every transformation to each of the clauses in Q and measure the execution time of the transformed clauses.

The parameter $b = b_0^{e/l}$, with e the number of *edge* literals, is an estimate for the nondeterminacy of the clause q . This correction on b_0 is necessary because only *edge* literals have a nondeterminacy of b_0 , *label* literals are determinate. A side effect of this transformation is that the parameter b has continuous values. These continuous values are discretized into three intervals as shown in Table 2.

Average execution times are computed for each of the 27 combinations of s , b and l . To obtain reliable averages, we repeat the entire experiment 100 times. The average times and the corresponding speedups are plotted in Figure 5.

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