Rule Ordering in Bottom-Up Fixpoint Evaluation of Logic Programs

Raghu Ramakrishnan Divesh Srivastava S. Sudarshan

Abstract

Logic programs can be evaluated bottom-up by repeatedly applying all rules, in "iterations", until the fixpoint is reached. However, it is often desirable—and in some cases, e.g. programs with stratified negation, even necessary to guarantee the semantics—to apply the rules in some order. We present two algorithms that apply rules in a specified order without repeating inferences. One of them (GSN) is capable of dealing with a wide range of rule orderings but with a little more overhead than the well-known *semi-naive* algorithm (which we call BSN). The other (PSN) handles a smaller class of rule orderings, but with no overheads beyond those in BSN.

We also demonstrate that by choosing a good ordering, we can reduce the number of rule applications (and thus joins). We present a theoretical analysis of rule orderings and identify orderings that minimize the number of rule applications (for all possible instances of the base relations) with respect to a class of orderings called fair orderings. We also show that while non-fair orderings may do a little better on some data sets, they can do much worse on others. The analysis is supplemented by performance results.

Index Terms: Bottom-up Evaluation, Control Expression, Cyclic Ordering, Deductive Database, Query Evaluation, Logic Programming, Rule Ordering, Semi-naive Evaluation.

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The authors are with the Computer Sciences Department, University of Wisconsin-Madison, WI 53706, U.S.A. The authors' e-mail addresses are {raghu,divesh,sudarsha}@cs.wisc.edu.

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1 Introduction

Bottom-up evaluation of logic programs is an important issue in deductive database applications. Algorithms that do not repeat inferences are considered desirable and said to have the *non-repetition* property. Essentially the same algorithm (called the "Semi-Naive" algorithm in the literature, and which we call Basic Semi-Naive or BSN) has been proposed independently by several researchers. It evaluates the fixpoint in an iterative fashion, with every rule applied once in each iteration. Facts generated in an iteration can be used to generate other facts only in subsequent iterations. The issue of how to apply rules in a specified order while retaining the non-repetition property is not addressed. Rule orderings are significant for several reasons.

• First, they are sometimes required to compute the answers correctly.

For example, in programs with stratified negation, lower strata must be evaluated before higher strata, and this simple ordering can become much more complex once we rewrite the program using Magic Sets [22, 3, 7, 20] (which is an important technique used widely to avoid inferences that are not "relevant" to the query).

• Second, rule ordering can result in increased efficiency.

For example, in an SCC-by-SCC evaluation of a program, rules in lower SCCs do not need to be considered while applying rules in higher SCCs; this can improve the efficiency of evaluation.

An important contribution of this paper is to demonstrate that ordering rules *within* an SCC can also improve efficiency by further reducing the number of rule applications. Although the orderings we consider for this purpose do not affect the number of inferences made, the processing becomes more set-oriented, with each rule application generating more tuples.

• Finally, rule orderings have been proposed to prune redundant derivations and to allow the user to specify a desired semantics [12, 11, 13].

In this paper, we use regular expressions over rules to specify orderings of the application of rules; we call these *control expressions*.

In the first part of our paper, we present two fixpoint algorithms that address the issue of how to apply rules in a specified order while retaining the non-repetition property. One of them, General Semi-Naive (GSN), applies a rule to produce new facts, and then immediately makes these facts available to subsequent applications of other rules (possibly in the same iteration). The GSN algorithm can deal with a large set of control expressions, and is described in Section 3. The other algorithm we present, Predicate Semi-Naive (PSN), can utilize facts produced for a predicate p in the same iteration they have been derived in, although not always in the immediately following rule application; this is described in Section 4. It handles a more restricted set of control expressions compared to GSN, but is cheaper than GSN. Indeed, it has no additional overheads compared to BSN.

The algorithms we describe in this paper are independent of program rewriting techniques such as Magic Sets that also seek to reduce the number of derivations. We can apply such techniques (and other techniques that generate control expressions on rule applications [12, 11]) on a given program, and then evaluate the resultant program using our algorithms.

In the second part of our paper, in Section 5, we study rule orderings in detail, and establish a close connection between cycles in rule graphs (which are a variant of rule/goal graphs defined in [4, 25]) and orderings that minimize the number of iterations and rule applications. We define what it means for a rule ordering to preserve a simple cycle, and show that a rule ordering that preserves all simple cycles in the rule graph (if such an ordering exists) is optimal within a certain class of rule orderings in minimizing the number of iterations, and hence the number of rule applications and joins.

In the third part of our paper, in Section 6, we present a performance study that underscores the importance of utilizing facts early, and choosing a good rule ordering. The terminology used in the rest of this paper is introduced in Section 2, and related work is presented in Section 7.

Due to space limitations, several proofs and details have been omitted. The performance study is also presented briefly. We direct the interested reader to [21] for a version of this paper with all proofs and additional discussion of several points.

2 Background

2.1 Definitions

The language considered in this paper is that of Horn logic. Such a language has a countably infinite set of variables and countable sets of function and predicate symbols, these sets being pairwise disjoint. We adopt the Prolog convention of denoting variables by strings of characters starting with an upper case letter (e.g. X, Y1); function and predicate symbols are strings of characters starting with a lower case letter. It is assumed, without loss of generality, that with each function symbol f and each predicate symbol p is associated a unique natural number n, referred to as the *arity* of the symbol; f and p are then said to be *n*-ary symbols. A 0-ary function symbol is referred to as a *constant*.

A term in such a language is a variable, a constant, or a compound term $f(t_1, \ldots, t_n)$ where f is an *n*-ary function symbol and the t_i are terms. A tuple of terms is sometimes denoted simply by the use of an overbar, e.g. \overline{t} . A literal (or predicate occurrence) in such a language is of the form $p(t_1, \ldots, t_n)$ (a positive literal) or $\neg p(t_1, \ldots, t_n)$ (a negative literal), where p is an *n*-ary predicate symbol and the t_i are terms. Predicates in such a language are also referred to as relations.

A substitution is a mapping from the set of variables of the language under consideration to the set of terms. Substitutions are denoted by lower case Greek letters θ, σ, φ , etc. A substitution σ is more general than a substitution θ if there is a substitution φ such that $\theta = \sigma[\varphi]$. Two terms t_1 and t_2 are said to be *unifiable* if there is a substitution σ such that $t_1[\sigma] = t_2[\sigma]$; σ is said to be a *unifier* of t_1 and t_2 . Note that if two terms have a unifier, they have a most general unifier (*mgu*) that is unique up to renaming of variables.

A clause is the disjunction of a finite number of literals, and is said to be a Horn clause if it has at most one positive literal. A Horn clause with exactly one positive literal is referred to as a *definite clause*. The positive literal in a definite clause is its *head*, and the remaining literals, if any, constitute its *body*. Following the syntax of Prolog, definite clauses (or *rules*) are written as: " $p: -q_1, \ldots, q_n$." This is read declaratively as q_1 and \ldots and q_n implies p. A predicate definition consists of a set of definite clauses, whose heads all have the same predicate symbol; a goal is a set of negative literals. We consider a logic program to be a pair $\langle P, Q \rangle$ where P is a set of predicate definitions and Q is the input, consisting of a query, or goal, and a (possibly empty) set of facts for "database predicates" appearing in the program.¹

We follow the convention in deductive database literature of separating the set of rules with non-empty bodies from the set of facts, or unit clauses, which appears in P. The set of facts is called the *database*. Predicates appearing in the heads of rules with non-empty bodies are referred to as *derived* predicates, and predicates appearing in database facts are referred to as *base* predicates. The program can be normalized to make the set of base predicates disjoint from the set of derived predicates, and we assume that this has been done. The motivation for separating the program from the database is that optimizations are applied only to the program, and not to the database. This is important in the database context since the set of facts can be very large. However, the distinction is artificial, and we may choose to consider (a subset of) facts to be rules.

The meaning of a logic program is given by its least Herbrand model [26]. From [26], this is equivalent to the least fixpoint semantics. A derived predicate p in a program P is said to be *safe* if, given any finite extension for each of the base predicates, p has a finite extension in the least Herbrand model of P.

We use the notion of derivation trees and derivation steps in several proofs.

Definition 2.1 Derivation tree : Given a program P with database D, derivation trees in $\langle P, D \rangle$ are defined as follows:

- Every fact h in D is a derivation tree for itself; a single node with label h.
- Let R be a rule: $p: -q_1, \ldots, q_n$ in P, let $d_i, 1 \le i \le n$, be facts with derivation trees T_i , and let θ be the mgu of (q_1, \ldots, q_n) and (d_1, \ldots, d_n) . Then the following is a derivation tree for $p[\theta]$: the root is a node labeled with $p[\theta]$ and R, and each $T_i, 1 \le i \le n$, is a child of the root.

A derivation step for fact $p[\theta]$ consists of a rule $R : p: -q_1, \ldots, q_n$, and facts $d_i, 1 \le i \le n$, with derivation trees, such that θ is the mgu of (q_1, \ldots, q_n) and (d_1, \ldots, d_n) . Thus, a derivation step consists of a non-leaf node and all its children in a derivation tree. \Box

¹Our definitions and results can be extended to handle features such as negation, set grouping and aggregation [6].

Note that the substitution θ is not applied to the children of $p[\theta]$ in the second part of the above definition. Thus, a derivation tree records which set of (previously generated) facts is used to generate a new fact using a rule, rather than the set of substitution instances of these facts that instantiated the rule. Derivation trees are important due to the following well-known property (see for example [18]):

Proposition 2.1 For every fact t in the least Herbrand model, there is a derivation tree with root r, and a substitution σ such that $t = r[\sigma]$, and for every fact r' that is the root of a derivation tree, each fact t' that is a ground instance of r' is in the least Herbrand model. \Box

Definition 2.2 Derivation height : The *height* of a derivation tree is defined to be the number of nodes in the longest path in the tree (which is always from the root to a leaf). The *derivation height* of a fact is defined as the minimum among the heights of its derivation trees. \Box

Definition 2.3 Rule application, Evaluation : The *application* of a rule R, using a given set of facts D, denoted by R(D), produces the set of *all* facts that can be derived in a single derivation step, using R and only the facts in D.

We define an *evaluation* as a sequence of rule applications. \Box

Definition 2.4 Non-repetition property : An evaluation is said to have the *non-repetition* property if no derivation step is repeated in the evaluation. \Box

A fact is said to have been seen by a rule R if the fact was available to an application of R. The *independent* application of a set of rules $\{R_1, \ldots, R_n\}$ on a set of facts D is defined as $R_1(D) \cup \ldots \cup R_n(D)$; i.e. each rule is applied once but the facts produced using a particular rule application are not seen by any of the other rules in the set. The *closure* of a set of rules $\mathcal{R} = \{R_1, \ldots, R_n\}$ using a given set of facts D refers to the derivation of all facts that can be computed using the given facts, and any number of applications of the rules, i.e.

$$closure(\mathcal{R}, D) = D \cup F(D) \cup F^2(D) \cup F^3(D) \cup \dots$$

where $F(D) = R_1(D) \cup \ldots \cup R_n(D)$.

We also use some terminology from graph theory. A directed graph G = (V, E) (where V is the vertex set and E is the edge set) is said to be *strongly connected* if every vertex in V is reachable (using the directed edges in E) from every other vertex in V. Given a directed graph G, a subgraph G1 of G is said to be a *strongly connected component* (SCC) of G if G1 is a

maximal subgraph in G that is strongly connected. Note that the SCCs of a directed graph G partition the vertices of G. Define the *reduction* of G wrt the SCCs as the graph G' = (V', E') obtained with vertex set V' being the set of SCCs $\{S_1, \ldots, S_k\}$ of G. An edge $(S_i, S_j), i \neq j, \in E'$ if there is an edge $(V_k, V_l) \in E$, where V_k is a vertex in S_i and V_l is a vertex in S_j . The reduction G' of G reflects the SCC structure of G, and is acyclic. An SCC S_1 is said to be *lower* than SCC S_2 in G if there is a (non-trivial) path from S_1 to S_2 in G'.

Definition 2.5 Rule (predicate) graph : Given a program P with rules $\mathcal{R} = \{R_1, R_2, ..., R_n\}$, we define the *rule graph* of P as the directed graph $G = (\mathcal{R}, E)$, where $(R_i, R_j) \in E$ iff the head of R_i unifies with a predicate occurrence in the body of R_j . We refer to SCCs in the rule graph as *Rule-SCCs*.

Given a program P with predicates $Pred = \{p_1, p_2, \ldots, p_n\}$, we define the *predicate graph* of P as the directed graph G = (Pred, E), where $(p_i, p_j) \in E$ iff p_i occurs in the body of a rule defining p_j . We refer to SCCs in the predicate graph as *Pred-SCCs*. \Box

Where the distinction is not relevant, we refer to Rule-SCCs and Pred-SCCs as SCCs.

Given a program P, Pred-SCCs S_1, \ldots, S_m are said to be in *topological order*, if whenever S_i contains a predicate used in S_j , then $i \leq j$.

2.2 Basic Semi-Naive Evaluation

Given a program P with an input database D, the Naive evaluation ([2]) of $\langle P, D \rangle$ proceeds in iterations. In each iteration, each of the rules in the program is independently applied on the set of available facts, and the set of available facts is updated at the end of the iteration. The Naive evaluation terminates when no new facts can be computed for any of the derived predicates of the program. The database D constitutes the initial set of available facts. This evaluation strategy can be refined by evaluating one strongly connected component (SCC) of the rule graph at a time, in a topological ordering of the Rule-SCCs. The Naive evaluation of a program on a database is extremely inefficient since it repeats a lot of derivations; each derivation step made in an application of a rule is repeated in every subsequent application of the same rule. Example 2.1 below illustrates this.

Iteration	Derivations made
1	$\{\mathbf{R1}:\mathbf{anc}(1,2),\mathbf{R1}:\mathbf{anc}(2,3),\mathbf{R1}:\mathbf{anc}(4,5)\}$
2	$\{\mathbf{R2}:\mathbf{anc}(1,3),\mathbf{R3}:\mathbf{anc}(1,3),R1:anc(1,2),R1:anc(2,3),R1:anc(4,5)\}$
3	$\{R2: anc(1,3), R3: anc(1,3), R1: anc(1,2), R1: anc(2,3), R1: anc(4,5)\}$

Table 1: Derivations in a Naive Evaluation of P

Example 2.1 Consider the following program $\langle P, Q \rangle$:

$$\begin{split} &R1: anc(X,Y): - par(X,Y).\\ &R2: anc(X,Y): - par(X,Z), anc(Z,Y).\\ &R3: anc(X,Y): - anc(X,Z), anc(Z,Y).\\ & par(1,2). \quad par(2,3). \quad par(4,5)..\\ & \text{Query: $?-anc(1,X).} \end{split}$$

A naive bottom-up evaluation of $\langle P, Q \rangle$ would result in the derivations of *anc* facts as shown in Table 1. (The rule used to derive a fact is also indicated, and new derivations are shown in bold-face.) Only anc(1,2) and anc(1,3) are answers to the query. Note that each derivation made in an iteration of naive evaluation is repeated in subsequent iterations. \Box

Recall that an evaluation is said to have the non-repetition property if no derivation step is repeated in the evaluation. A number of researchers have independently proposed an evaluation technique that we call *Basic Semi-Naive*, or BSN, with the non-repetition property ([9, 19, 5, 2, 1]). Given a program P, the *Basic Semi-Naive* (BSN) evaluation of P proceeds a Pred-SCC at a time in a topological ordering of the Pred-SCCs. There are essentially two components to the BSN evaluation of a Pred-SCC.

1. The first is a rewriting of the Pred-SCC S that defines "differential" versions of predicates, in order to distinguish facts that have been newly generated (and not yet used in inferences) from older facts. We present below a variant of the version described in [1].

For each predicate p defined in S, we have four predicates p, p^{old} , δp^{old} and δp^{new} . For each rule in S of the form $R: p: -q_1, \ldots, q_m$, where q_1, \ldots, q_m are all non-recursive to p, the following semi-naive rewritten rule is obtained ([1]) from $R: R_1 \ \delta p^{new}: -q_1, \ldots, q_m$. We call such rules *non-recursive* semi-naive rules.

For each rule in S of the form: $R: p: -p_1, \ldots, p_n, q_1, \ldots, q_m$, where p_1, \ldots, p_n (n > 0) are mutually recursive to p and q_1, \ldots, q_m are not, the following n semi-naive rewritten rules

are obtained from R:

$$R_{1}: \delta p^{new}: - \delta p_{1}^{old}, p_{2}, \dots, p_{n}, q_{1}, \dots, q_{m}.$$

$$R_{2}: \delta p^{new}: - p_{1}^{old}, \delta p_{2}^{old}, p_{3}, \dots, p_{n}, q_{1}, \dots, q_{m}.$$

$$\vdots$$

$$R_{n}: \delta p^{new}: - p_{1}^{old}, p_{2}^{old}, \dots, p_{n-1}^{old}, \delta p_{n}^{old}, q_{1}, \dots, q_{m}.$$

We call such rules *recursive* semi-naive rules.

2. The second component is a technique to apply the rewritten rules and update these differentials, ensuring that all derivations are made exactly once.

In evaluating S, the first iteration consists of applying each of the semi-naive (both nonrecursive and recursive) rewritten rules in S. Subsequent iterations consist of applying only the recursive semi-naive rules. The evaluation of S proceeds by iterating until no new facts are computed for any of the predicates defined in S. After applying the semi-naive rules in an iteration, the extensions of the semi-naive relations for each p_i are updated using Procedure SN_Update below. (Note that the operators "—" and "U" involve subsumption checks if non-ground facts are generated.)

procedure SN_Update(p_i) $p_i^{old} := p_i^{old} \cup \delta p_i^{old}.$ $\delta p_i^{old} := \delta p_i^{new} - p_i^{old}.$ $p_i := p_i^{old} \cup \delta p_i^{old}.$ $\delta p_i^{new} := \phi.$

At every stage of the evaluation, the set of relations p_i^{old} , for all *i*, has the property that every derivation that uses only these facts has been made. This can be seen from the nature of the δ terms in rule bodies, and the order of updates of the various semi-naive relations.

Example 2.2 Consider again the program P of Example 2.1. Using BSN, the sequence of derivations made is shown in Table 2. Note that each derivation made in the naive evaluation is also made in the BSN evaluation. However, the BSN evaluation does not repeat any derivations. But if a fact (anc(1,3) in this case) is derived by two *different* derivations, each of these derivations is made in the BSN evaluation.

Iteration	Derivations made
1	$\{\mathbf{R1}: anc(1,2), \mathbf{R1}: anc(2,3), \mathbf{R1}: anc(4,5)\}$
2	$\{\mathbf{R2}:\mathbf{anc}(1,3),\mathbf{R3}:\mathbf{anc}(1,3)\}$
3	{}

Table 2: Derivations in a BSN Evaluation of P

Such "redundant" derivations can be avoided by recognizing that every fact derived using rule r2 is also derived using r3 and vice versa. Consequently, a scheme that prunes redundant derivations ([11], for instance) could recognize this and never apply rule R3; the derivation of R3 : anc(1,3) is not made in such a case. Given a program that has redundant derivations, its evaluation using BSN does *not* eliminate such derivations.

Rewriting the program P of Example 2.1 using Magic Sets, and evaluating the rewritten program also results in avoiding the derivation of anc(4, 5), since this is "irrelevant" to computing the answers to the query. Refer to [7, 20] for more details. \Box

2.3 Control Expressions

Helm [12, 11] introduced the notion of control on the bottom-up evaluation of logic programs using control expressions, and also looked at control as a way of increasing the efficiency of evaluation by eliminating some redundant derivations. Control expressions have also been proposed to let the user specify a desired semantics in the presence of negation ([13]), as well as evaluate the Magic Sets transformation of stratified logic programs ([8]). While applications of control expressions have been considered, not much attention has been given to efficient implementation of control expressions.

Let R_1, \ldots, R_n denote the rules of a program. The syntax of our control expressions is given by the following grammar.

$$S \rightarrow T$$

$$T \rightarrow F | F + T | F \oplus T | F \cdot T$$

$$F \rightarrow R_i | (T) | F^* | F^{\bullet}$$

where S is the start symbol of the grammar.

A control expression α is a (non-deterministic) mapping $\mathcal{D} \to \mathcal{D}$, where \mathcal{D} is the set of all

database states. (A database state is a set of facts for the base and derived predicates.) The initial database state consists of the set of all given facts for the base predicates. The following equations recursively define the semantics of control expressions.

- 1. If $\alpha = R_i$, $\alpha(D) = D \cup R_i(D)$.
- 2. If $\alpha = \alpha_1 \cdot \alpha_2$, $\alpha(D) = \alpha_2(\alpha_1(D))$.
- 3. If $\alpha = \alpha_1 \oplus \alpha_2$, non-deterministically choose α_1 or α_2 and call it β_1 and call the other β_2 . if $\beta_1(D) \neq D$, $\alpha(D) = \beta_1(D)$, else $\alpha(D) = \beta_2(D)$.
- 4. If $\alpha = \alpha_1^{\bullet}$, $\alpha(D) = \alpha_1^i(D)$, for some arbitrary choice of $i \ge 0$, where $\alpha_1^0(D) = D$ and $\alpha_1^{j+1}(D) = \alpha_1(\alpha_1^j(D)), j \ge 0.$
- 5. If $\alpha = \alpha_1^*$, $\alpha(D) = \alpha_1^i(D), i > 0$, such that $\alpha_1^{i+1}(D) = \alpha_1^i(D)$, where $\alpha_1^0(D) = D$ and $\alpha_1^{j+1}(D) = \alpha_1(\alpha_1^j(D)), j \ge 0$.
- 6. If $\alpha = \alpha_1 + \alpha_2$, $\alpha(D) = \alpha_1(D) \cup \alpha_2(D)$.

The semantics of the control operators that we consider are different from those considered by Helm and we do not discuss the evaluation of Helm's control operators in this paper.

The control expressions we use form a superset of those used by Imielinski and Naqvi [13]. It follows from [13] that our control expressions can be used to specify the inflationary semantics for negation ([14]). Our control expressions also form a superset of those used by Beeri et al. [8].

3 General Semi-Naive Evaluation

We now present a technique, *General Semi-Naive*, or GSN, evaluation, that makes facts computed by an application of a rule R available to *all* other rule applications immediately after the application of R, while maintaining the non-repetition property. We first look at how to apply a single rule using the GSN technique. Then, in Section 3.3, we look at how to evaluate general control expressions using this technique. Finally, in Section 3.4, we look at a specific form of control expression to evaluate a program Pred-SCC by Pred-SCC, in a topological ordering of the Pred-SCCs.

3.1 GSN for a Rule

Consider any rule R_i in the program:

$$R_i: p_h(\overline{t}): -p_1(\overline{t_1}), \ldots, p_n(\overline{t_n}).$$

where p_h, p_1, \ldots, p_n are not necessarily distinct. Similar to BSN evaluation, the GSN evaluation of a rule consists of two components.

1. The first is the semi-naive rewriting that defines the "differential" versions of predicates.

Associated with each rule R_i of the program, and each (non-recursive as well as recursive) predicate p_j that occurs in the body of R_i , we maintain a relation p_{j,R_i}^{old} . (Even if a predicate occurs more than once in the body of R_i , only one copy of the relation needs to be maintained per rule.) The set of relations p_{j,R_i}^{old} , $1 \le j \le n$, has the property that every derivation that can be made by an application of R_i using only these facts has already been made. (In this respect, it is similar to the p^{old} relations maintained by BSN.)

Associated with each predicate p_j , we also maintain its complete extension; only one copy of p_j needs to be maintained independent of the number of rules p_j occurs in. We also have "temporary" relations δp_j^{new} and δp_j^{old} associated with each derived predicate p_j that appears in the program. These δ relations are used to keep track of "new" facts; facts which have not been seen by a rule application.

The semi-naive rewriting for GSN is obtained as follows. Each predicate p_j in the body of R_i is treated as being recursive to the head. The semi-naive rewriting described for BSN (in Section 2.2) is modified by replacing each predicate occurrence p_j^{old} in the body of the semi-naive rewritten versions of rule R_i by p_{j,R_i}^{old} .

2. The second is a technique to apply the rewritten rules and update these differentials in the order specified for rule applications, ensuring that all derivations are made exactly once. Procedure GSN_Rule below describes the application of a rule R_i on a set of facts D.

procedure $GSN_Rule(R_i, D)$

/* All relations mentioned below are part of D. */

Let p_h be the predicate defined by rule R_i .

(1) For every predicate p_j in the body of R_i , $\delta p_j^{old} := p_j - p_{j,R_i}^{old}$.

- (2) $\delta p_h^{new} := \phi$.
- (3) Apply each semi-naive rewritten version of R_i

(modified as described earlier) independently.

- (4) For every predicate p_j in the body of R_i , $p_{j,R_i}^{old} := p_j$.
- (5) $p_h := p_h \cup \delta p_h^{new}$.
- (6) return δp_h^{new} .

Example 3.1 Consider rule R3 from P in Example 2.1.

R3: anc(X, Y): - anc(X, Z), anc(Z, Y).

The semi-naive rewriting of this rule for GSN evaluation is given by:

 $R3': \delta anc^{new}(X,Y): -\delta anc^{old}(X,Z), anc(Z,Y).$ $R3'': \delta anc^{new}(X,Y): -anc^{old}_{R_3}(X,Z), \delta anc^{old}(Z,Y).$

Prior to the application of these rules, δanc^{old} is initialized to $anc - anc_{R_3}^{old}$, i.e. the tuples that have not been seen as yet by an application of R_3 . The relation δanc^{new} is initialized to ϕ . Application of rules R3' and R3'' (potentially) results in adding facts to the δanc^{new} relation. After the application of this rule, each fact in δanc^{old} also has been seen by R_3 . Consequently, the $anc_{R_3}^{old}$ relation is updated to reflect this. The newly generated facts are also added to the *anc* relation.

Example 3.2 describes the order in which rules are applied (and relations updated) in the GSN evaluation of program P. \Box

3.2 Efficient Implementation of GSN

The description of General Semi-Naive above suggests that each rule has to separately maintain an extension for each predicate that occurs in its body and would thus appear to be inefficient in terms of the storage used. However, it has a simple implementation in which each relation is maintained as a list of tuples. The new facts produced by a rule application are appended to the extension of the predicate. (In the case of generalized tuples, if a new fact subsumes an existing fact then the existing fact may have to be deleted.)

In step (4) of $\mathsf{GSN_Rule}(R_i, D)$, every fact in a relation p_j becomes part of p_{j,R_i}^{old} . Since the tuples in each relation p_j are ordered by "time" of insertion, a pointer to the end of the extension

of p_j at step (4) has the following property: every p_j fact preceding the pointer is part of p_{j,R_i}^{old} , and every p_j fact occurring after the pointer has not been seen by rule R_i yet. Hence, the extension of each p_{j,R_i}^{old} is replaced by a pointer into the extension of p_j . Similarly, we replace the extension of each δp_j^{old} by another pointer into the extension of p_j such that the set of facts between the pointers for p_{j,R_i}^{old} and δp_j^{old} constitutes the extension for δp_j^{old} . The set of facts beyond the pointer for δp_j^{old} constitutes the extension of δp_j^{new} . Thus, separate extensions of a predicate do not have to be maintained for each rule.

Indices are important for efficiently accessing a relation during evaluation. The pointers into the extent of a relation p_j partition it based on the order of insertion of tuples. Each of the relations p_{j,R_i}^{old} , $\delta p_{j,R_i}^{old}$ and δp_j^{new} is the union of a contiguous set of these partitions. To index any of these relations, we index in turn each partition that is contained in it. Since there is at most one pointer per rule into p_j , the number of partitions is bounded by the number of rules in the program. There are two operations on pointers into the relation. The first is the creation of a new pointer to the end of a relation, and the second is the deletion of a pointer into a relation. In the first case a new partition is created (and is initially empty), and in the second case two existing partitions get merged. This involves moving tuples from the indices of one partition to the indices of the other. However, each tuple is moved in this fashion only a constant number of times, and hence this will not increase the time complexity of evaluation. Thus, this indexing technique can be expected to provide efficient access to tuples. (Alternatively, we could index each of the relations p_{j,R_i}^{old} , $\delta p_{j,R_i}^{old}$ and δp_j^{new} separately. This could result in each tuple being indexed multiple times.)

3.3 Implementing Control Expressions Using GSN

The semantics of control expressions was described (using recursive equations) in Section 2.3. This description also suggests a straightforward way of evaluating control expressions, as outlined in Procedure Simple_CE(α , D) below. In this evaluation, every fact produced by a rule application on a given database is immediately added to the database, and made available to subsequent rule applications. Unfortunately, such an evaluation does not have the non-repetition property, and can be extremely inefficient.

procedure Simple_CE(α , D)

/* We need to evaluate control expression α on database D. */

case

(1) $\alpha = R_i$: return $D \cup R_i(D)$. /* the exit case */ return Simple_CE(α_2 , Simple_CE(α_1 , D)). (2) $\alpha = \alpha_1 \cdot \alpha_2$: (3) $\alpha = \alpha_1 \oplus \alpha_2$: Non-deterministically choose α_1 or α_2 and call it β_1 and call the other β_2 . if Simple_CE(β_1, D) $\neq D$, return Simple_CE(β_1, D) else return Simple_ $CE(\beta_2, D)$. (4) $\alpha = \alpha_1^{\bullet}$: Choose $n \ge 0$ non-deterministically. for i = 1 to n do Let D denote the result of evaluating Simple_CE(α_1, D). return D. (5) $\alpha = \alpha_1^*$: repeat Let D' denote D at this stage. Let D denote the result of evaluating Simple_CE(α_1 , D). until (D = D')return D. (6) $\alpha = \alpha_1 + \alpha_2$: return Simple_CE(α_1, D) \cup Simple_CE(α_2, D). end case

Note that Procedure Simple_CE is non-deterministic in that the final database state may not be uniquely determined if the control expression contains \oplus or \bullet .

Since a control expression is a non-deterministic mapping, we use {Semantics(α , D)} to denote the set of all possible database states that are results of $\alpha(D)$. Similarly, we use {Simple_CE(α , D)} to denote the set of all possible results of the procedure Simple_CE(α , D). Theorem 3.1 For all databases D, and control expressions α , {Simple_CE(α , D)} = {Semantics

$$(\alpha, D)\} \square$$

The proof follows in a straightforward manner from the direct correspondence between each case in the semantics of control expressions and Procedure Simple_CE.

Consider the restricted set of control expressions generated by the grammar described in Section 2.3 without the production that uses the "+" operator. Procedure GSN_CE (α , D) below describes how we can evaluate such a control expression while preserving the non-repetition property.

```
procedure GSN_CE(\alpha, D)

\alpha is a control expression and D is an initial database.

/* D contains the p_j as well as all p_{j,R_i}^{old} relations. */

case

(1) \alpha = R_i: return D \cup GSN_Rule(R_i, D).

....

/* Cases (2)-(5) remain unchanged from Simple_CE(\alpha, D). */

end case
```

The updates to the various differential relations (corresponding to the predicates occurring in the head and body of rule R_i) performed by **GSN_Rule** maintain the set of facts that have been used by the rule R_i in previous applications of R_i . This ensures that in subsequent applications of R_i , none of the previous derivations is repeated. Intuitively, each rule in the program individually ensures that its applications have the non-repetition property; we prove this claim formally in Theorem 3.2. First we need the following proposition and its corollary.

Proposition 3.1 Suppose a control expression α does not use the operator "+". Given a database D, any evaluation of Simple_CE (α, D) performs a sequence of rule applications; each rule application in the sequence takes as input the set of all facts derived up to the previous rule application in the sequence. \Box

The intuition behind the proof of this proposition is that each call to Simple_CE either performs a rule application, or performs a sequence of recursive calls to Simple_CE, each of which takes as input all facts derived in earlier calls. The structure of GSN_CE is similar to that of Simple_CE, and hence we have the following corollary.

Corollary 3.1 Suppose a control expression α does not use the operator "+". Given a database D, any evaluation of GSN_CE(α , D) performs a sequence of calls to GSN_Rule; each call takes as input the set of all facts derived up to the previous call in the sequence. \Box

Since GSN_CE is a non-deterministic procedure, we use $\{GSN_CE(\alpha, D)\}$ to denote the set of all possible results of $GSN_CE(\alpha, D)$.

Theorem 3.2 For all databases D, and control expressions α that have no occurrence of the "+" operator, $\{\mathsf{GSN_CE}(\alpha, D)\} = \{\mathsf{Simple_CE}(\alpha, D)\}$. Further, evaluations of $\mathsf{GSN_CE}$ have the non-repetition property.

Proof: By Proposition 3.1 and Corollary 3.1, Simple_CE (resp. GSN_CE) evaluations of control expressions that do not contain "+" perform a sequence of rule applications (resp. calls to GSN_Rule).

We first prove three properties of a sequence of calls to $\mathsf{GSN_Rule}$. Consider a particular call to $\mathsf{GSN_Rule}(R_i, D)$ in this sequence of calls.

P1. Before step (3) of GSN_Rule, for each predicate p_j occurring in the body of R_i , the set of facts $\delta p_j^{old} = p_j - p_{j,R_i}^{old}$ has not been seen by R_i .

P2. Before step (3) of GSN_Rule, every derivation that could be made by an application of R_i using only the facts in the set of relations p_{j,R_i}^{old} , for all j, has already been made.

P3. After step (3) (but before step (5) where p_h is updated) of GSN_Rule, every derivation that could be made using R_i and only the facts in the set of relations p_j , for all j, has been made.

We prove these by induction on the number of times $\text{GSN}_\text{Rule}(R_i, _)$ is called. Before the first call to $\text{GSN}_\text{Rule}(R_i, _)$, p_{j,R_i}^{old} is empty for all p_j in the body of R_i . The call to $\text{GSN}_\text{Rule}(R_i, _)$ computes all facts that can be derived from the given set of facts for the body predicates. Hence, **P1** and **P2** are true before step (3) and **P3** is true after step (3), and the basis holds.

For the induction step, assume that P1, P2, and P3 hold up to the kth call to GSN_Rule(R_i , _), and consider the k + 1th call to GSN_Rule(R_i , _). Steps (4) and (5) of GSN_Rule in the kth call to GSN_Rule(R_i , _), and step (1) in the k + 1th call to GSN_Rule(R_i , _) ensures P1. Since P3 holds in the kth call to GSN_Rule(R_i , _), step (4) guarantees that P2 holds in the k + 1th call to GSN_Rule(R_i , _). Since P2 holds in the k + 1th call to GSN_Rule(R_i , _), step (1) and the correctness of semi-naive rewriting ensures that P3 holds in the k + 1th call to GSN_Rule(R_i , _). This completes the proof of P1, P2, and P3.

The GSN evaluation of a control expression without any occurrence of the "+" operator proceeds sequentially and does not have any parallel branches; databases never need to be merged and, hence, for each predicate p and each rule R that uses p, the relations p_R^{old} and δp^{old} are well defined. Property **P1** then ensures that no derivations using any rule R are repeated and hence evaluations of **GSN_CE** (for control expressions without the "+" operator) have the non-repetition property.

Claim 1: Consider a particular evaluation of Simple_CE(α , D) (where the choices, if any, due to occurrences of the \oplus or \bullet operators, have been made). Then, there is an equivalent evaluation of GSN_CE(α , D).

Claim 2: Consider a particular evaluation of $\mathsf{GSN_CE}(\alpha, D)$ (where the choices, if any, due to occurrences of the \oplus or \bullet operators, have been made). Then, there is an equivalent evaluation of $\mathsf{Simple_CE}(\alpha, D)$.

The intuition behind Claims 1 and 2 is that procedures GSN_CE and Simple_CE have the same recursive structure, and calls to GSN_Rule make the same inferences as new inferences made by the corresponding naive rule application. For formal proofs see [21].

This ends the proof of the theorem. \Box

Evaluating control expressions that contain the "+" operator using the GSN evaluation technique presents some difficulties. Recall that the evaluation of the "+" operator involved merging the two databases obtained by evaluating the operands of the "+" operator independently on the input database. Merging the p_{j,R_i}^{old} relations produced by the two operands of the "+" cannot in general be done consistently. In the special case that the two operands of a "+" operator do not have any rule in common, we can merge the databases consistently, by merely taking the union of the two. This handles the case of control expressions of the form $(R_1 + R_2 + \ldots + R_m)^*$, which simulate Basic Semi-Naive evaluation, as well as control expressions in the class PCE(P)(described in Section 4.1), which can be evaluated using Predicate Semi-Naive evaluation.

3.4 GSN for a Program

Consider a program P. With each Pred-SCC S in P, we can associate all the (non-recursive and recursive) rules defining the predicates in S. The evaluation of S consists of repeated application of the rules associated with S, and the program P can be evaluated Pred-SCC by Pred-SCC in a topological ordering of the Pred-SCCs of P. Procedure **GSN_Prog** below describes such a GSN evaluation of a program. The GSN evaluation of a single Pred-SCC is described by **GSN_SCC**.

procedure $GSN_Prog(P)$

Let S_1, \ldots, S_m be a topological ordering of the Pred-SCCs of P.

/* S_0 is assumed to contain all the base predicates. */

for j = 1 to m do $\mathsf{GSN_SCC}(S_j)$

procedure $GSN_SCC(S_i)$

Let the ordering of the non-recursive rules for the predicates in S_i be E_i^1, \ldots, E_i^m .

Let the ordering of the recursive rules for the predicates in S_i be R_i^1, \ldots, R_i^n .

if no predicate in S_i has any recursive rule defining it, then

Evaluate the following control expression:

 $(E_i^1 \cdot \ldots \cdot E_i^m)$ using GSN_CE.

else /* at least one of the predicates in S_i has a recursive rule defining it */

Evaluate the control expression

 $(E_i^1 \cdot \ldots \cdot E_i^m) \cdot (R_i^1 \cdot \ldots \cdot R_i^n)^*$ using GSN_CE.

In the evaluation of a Pred-SCC S of program P, the non-recursive rules are applied once, followed by the repeated application of the recursive rules of S. The Pred-SCC structure of Pensures that each non-recursive rule E in S only has body predicates that are in "lower" Pred-SCCs of P. Consequently, the facts produced by the application of any of the non-recursive rules cannot be used by any of the other non-recursive rules in S; thus, the order of application of the non-recursive rules in GSN_SCC is irrelevant. Thus we have the following theorem (for a formal proof, see [21]).

Theorem 3.3 Procedure GSN_Prog has the non-repetition property, is sound, and if all the predicates defined in the program are safe, is complete wrt the least fixpoint semantics. \Box

We illustrate GSN_Prog using an example.

Example 3.2 Consider again program P of Example 2.1.

F1 : par(1,2). F2 : par(2,3). F3 : par(4,5). R1 : anc(X,Y) : - par(X,Y). R2 : anc(X,Y) : - par(X,Z), anc(Z,Y).R3 : anc(X,Y) : - anc(X,Z), anc(Z,Y).

Facts par(1,2), par(2,3), and par(4,5) are in S_0 . Rules R1, R2 and R3 are all in Pred-SCC S_1 . A Pred-SCC by Pred-SCC evaluation of P using **GSN_Prog** would proceed as follows.

Iter/Rule	Relation	Facts in relation at end of rule application (New facts in bold-face)
0/R1	anc	$\{\mathbf{R1}:\mathbf{anc}(1,2),\mathbf{R1}:\mathbf{anc}(2,3),\mathbf{R1}:\mathbf{anc}(4,5)\}$
	par	$\{F1: par(1,2), F2: par(2,3), F3: par(4,5)\}$
	par_{R1}^{old}	$\{{f F1}:{f par}(1,2),{f F2}:{f par}(2,3),{f F3}:{f par}(4,5)\}$
1/R2	anc	$\{R1: anc(1,2), R1: anc(2,3), R1: anc(4,5), \mathbf{R2}: \mathbf{anc}(1,3)\}$
	anc_{R2}^{old}	$\{\mathbf{R1}:\mathbf{anc}(1,2),\mathbf{R1}:\mathbf{anc}(2,3),\mathbf{R1}:\mathbf{anc}(4,5)\}$
	par	$\{F1: par(1,2), F2: par(2,3), F3: par(4,5)\}$
	par_{R2}^{old}	$\{{f F1}:{f par}(1,2),{f F2}:{f par}(2,3),{f F3}:{f par}(4,5)\}$
1/R3	anc	$\{R1: anc(1,2), R1: anc(2,3), R1: anc(4,5), R2: anc(1,3), \mathbf{R3}: \mathbf{anc}(1,3)\}$
	anc_{R3}^{old}	$\{\mathbf{R1}: \mathbf{anc}(1, 2), \mathbf{R1}: \mathbf{anc}(2, 3), \mathbf{R1}: \mathbf{anc}(4, 5), \mathbf{R2}: \mathbf{anc}(1, 3)\}$
2/R2	anc	$\{R1: anc(1,2), R1: anc(2,3), R1: anc(4,5), R2: anc(1,3), R3: anc(1,3)\}$
	anc_{R2}^{old}	$\{R1: anc(1,2), R1: anc(2,3), R1: anc(4,5), \mathbf{R2}: \mathbf{anc}(1,3), \mathbf{R3}: \mathbf{anc}(1,3)\}$
	par	$\{F1: par(1,2), F2: par(2,3), F3: par(4,5)\}$
	par_{R2}^{old}	${F1: par(1,2), F2: par(2,3), F3: par(4,5)}$
2/R3	anc	$\{R1: anc(1,2), R1: anc(2,3), R1: anc(4,5), R2: anc(1,3), R3: anc(1,3)\}$
	anc_{R3}^{old}	$\{R1: anc(1,2), R1: anc(2,3), R1: anc(4,5), R2: anc(1,3), \mathbf{R3}: \mathbf{anc}(1,3)\}$

Table 3: Derivations in a GSN Evaluation of P

 S_1 contains only one non-recursive rule, R1. Assume the ordering of recursive rules in S_1 to be R2, R3. (Alternatively, R3, R2 could be the rule order assumed, and subsequent details worked through.) Consequently, the evaluation of S_1 requires evaluation of the control expression $(R1) \cdot (R2 \cdot R3)^*$. Table 3 describes the facts in the various relations during the GSN_Prog evaluation of P using this control expression.

Only the relevant relations that affect and are affected by the rule application are shown. Other relations remain unchanged from the previous step. Initially, *par* contains {*par*(1, 2), *par*(2, 3), *par*(4, 5)}, and all other relations are empty. Iteration 0 refers to the evaluation of the control expression (*R*1). Subsequent iterations refer to the evaluation of the control expression $(R2 \cdot R3)^*$.

Note that the *same* derivations are made in the GSN evaluation of P as in the BSN evaluation of P (Table 2). \Box

We discuss the issues involved in selecting an ordering of rules in Section 5.

4 Predicate Semi-Naive Evaluation

We now present a technique, *Predicate Semi-Naive*, or PSN, evaluation that has the nonrepetition property and can utilize facts produced by a rule application in the same iteration they have been derived in, though not immediately.² This technique maintains just one version of the extension of each predicate and incurs no additional overheads compared to BSN evaluation. We first look at how to evaluate a restricted set of control expressions for evaluating a program using this technique. Then, in Section 4.2, we look at a specific form of control expression to evaluate a program Pred-SCC by Pred-SCC in a topological ordering of the Pred-SCCs.

Similar to BSN (and GSN) evaluation, there are two components to PSN evaluation.

- 1. The first is the semi-naive rewriting that defines "differential" versions of predicates. Similar to BSN evaluation, we maintain four relations $p, p^{old}, \delta p^{old}$ and δp^{new} for each predicate p defined in the program. The semi-naive rewriting is identical to the rewriting described for BSN evaluation.
- 2. The second component is a technique to apply the rewritten rules and update these differentials, ensuring that all derivations are made exactly once. Procedure PSN_CE describes this component for a restricted class of control expressions for a program.

4.1 Implementing Control Expressions Using PSN

Using PSN, we can evaluate a restricted class of control expressions while obtaining the non-repetition property. For a program P, this class of control expressions is given by PCE(P).

The class PCE(P): Consider a program P. The class of expressions PCE(P) is as follows. Let S_1, \ldots, S_k be any topological ordering of the Pred-SCCs of P. Then, the expression $(PCE(S_1) \cdot \ldots \cdot PCE(S_k))$ is in the class PCE(P).

The class PCE(S): Consider a Pred-SCC S. The class of expressions PCE(S) is as follows. Let p_1, \ldots, p_m be some (not necessarily all) of the predicates in S. Let s_1, \ldots, s_n be any partitioning of the predicates p_1, \ldots, p_m . Let $E_{s_i}^1, \ldots, E_{s_i}^{a_i}$ be the non-recursive rules defining predicates in s_i , and let $R_{s_i}^1, \ldots, R_{s_i}^{b_i}$ be the recursive rules defining predicates in

²Although we describe just two evaluation strategies, a gradation is possible between GSN and PSN evaluation of SCCs, resulting in a range of evaluation strategies. Some set of predicates may be evaluated according to the strategy used by GSN, and other predicates evaluated according to PSN. We do not elaborate further on this.

 $s_i, 1 \leq i \leq n$. Let E_{s_i} stand for the control expression $E_{s_i}^1 + \ldots + E_{s_i}^{a_i}$, and let R_{s_i} stand for the control expression $R_{s_i}^1 + \ldots + R_{s_i}^{b_i}$. If none of p_1, \ldots, p_m has any recursive rule defining it, then the expression $(E_{s_1}) \cdot \ldots \cdot (E_{s_n})$ is in PCE(S). If at least one of p_1, \ldots, p_m has a recursive rule defining it, then $(E_{s_1} + R_{s_1}) \cdot \ldots \cdot (E_{s_n} + R_{s_n}) \cdot (R_{s_1} \cdot \ldots \cdot R_{s_n})^*$ is in PCE(S).

Procedure $\mathsf{PSN_CE}(\alpha, D)$ below describes the PSN evaluation of a restricted set of control expressions, without occurrences of the " \oplus " and the " \bullet " operators. Note that the absence of these operators means that each control expression is a deterministic mapping.

procedure PSN_CE(α , D) /* Evaluate control expression α on database D. */

case

(1) $\alpha = R_1 + \ldots + R_n$: /* the exit case */ for i = 1 to n do

Apply each semi-naive rewritten version of R_i on database D.

Let D denote the resulting database.

for every predicate p_j defined by a rule in α , call SN_Update(p_j).

(2) $\alpha = \alpha_1 \cdot \alpha_2$:

Evaluate PSN_CE(α_1, D).

Let D' denote the resulting database.

Evaluate PSN_CE(α_2, D').

(3) $\alpha = (\alpha_1)^*$: repeat Let D' denote D at this stage. Evaluate PSN_CE(α_1, D). Let D denote the resulting database. until (D = D')

end case

The Simple_CE evaluation of a control expression in PCE(S) first applies the rules in $(E_{s_1} + R_{s_1})$ independently. All facts computed are then added to the database. This is followed by applying the rules in $(E_{s_i} + R_{s_i})$ independently, adding facts computed to the database, for

successive values of i until i = n. The Simple_CE evaluation then applies the rules in R_{s_j} independently, adding facts computed to the database, for $j = 1 \dots n$, and repeats this until nothing new is derived.

The PSN_CE evaluation of a control expression in PCE(S) is very similar. It first applies the (semi-naive versions of the) rules in $(E_{s_i} + R_{s_i})$, followed by an SN_Update of the predicates in the partition s_i , for successive values of i until i = n. The PSN_CE evaluation then applies the rules in R_{s_j} followed by an SN_Update of the predicates in the partition s_j for j = 1...n, and repeats this until nothing new is derived.

Theorem 4.1 Consider a program P, a database D, and a control expression α in the class PCE(P). The resulting database after evaluating $PSN_CE(\alpha, D)$ is the same as $Simple_CE(\alpha, D)$. Further, evaluations of PSN_CE have the non-repetition property.

Proof: (Sketch) It is easy to show that the evaluation of $\mathsf{PSN_CE}(\alpha, D)$ is sound, i.e., its result is contained in the result of Simple_CE(α, D).

To show that it is complete, we do the following. We first sequentially number applications of rules in evaluations of $PSN_CE(\alpha, D)$ as well as evaluations of $Simple_CE(\alpha_1, D)$ such that rule applications made in parallel have the same number. Next we construct an ordering of the facts generated in an evaluation of $Simple_CE(\alpha, D)$ based on the numbering of the first rule application that generated the fact. Any derivation using any rule R that is made in an evaluation of $Simple_CE(\alpha_1, D)$ will be made (for the first time) during the first application of R after the generation of the last (in the ordering) fact used in the derivation. We then show by induction on the numbering of rule applications in $Simple_CE(\alpha, D)$ (and using the seminaive updates performed in the evaluation of $PSN_CE(\alpha, D)$) that this derivation is made in an application of R with the same number, in an evaluation of $PSN_CE(\alpha, D)$.

To prove the non-repetition property, we show that the semi-naive updates are done in such a fashion that a derivation as above can be made *only* in the rule application mentioned above. Thus only one rule application can make any derivation. \Box

For a full proof of this theorem, see [21].

4.2 **PSN** for a Program

Consider a program P. With each Pred-SCC S in P, we can associate all the (non-recursive and recursive) rules defining the predicates in S. The PSN evaluation of S consists of repeated applications of the rules associated with S, and the program P can be evaluated Pred-SCC by Pred-SCC in a topological ordering of the Pred-SCCs of P. Procedure **PSN_Prog** below describes the PSN evaluation of a program P, using a control expression in the class PCE(P) where in each Pred-SCC of P each predicate is in a partition by itself.

procedure PSN_Prog(P) Let S_1, \ldots, S_m be a topological ordering of the Pred-SCCs of P. /* S_0 is assumed to contain all the base predicates. */ for j = 1 to m do PSN_SCC(S_j)

procedure $PSN_SCC(S)$

Let the ordering of predicates in S be p_1, \ldots, p_k .

Let $E_{p_i}^1, \ldots, E_{p_i}^{a_i}$ be the non-recursive rules defining p_i , and

let $R_{p_i}^1, \ldots, R_{p_i}^{b_i}$ be the recursive rules defining $p_i, 1 \le i \le k$.

Let E_{p_i} stand for the control expression $E_{p_i}^1 + \ldots + E_{p_i}^{a_i}$, and

let R_{p_i} stand for the control expression $R_{p_i}^1 + \ldots + R_{p_i}^{b_i}$.

if no predicate in S has any recursive rule defining it, then

Evaluate the following control expression:

 $(E_{p_1}) \cdot \ldots \cdot (E_{p_k})$ using PSN_CE

else /* at least one of the predicates in S has a recursive rule defining it */ Evaluate the following control expression:

 $(E_{p_1}+R_{p_1})$... $(E_{p_k}+R_{p_k}) \cdot (R_{p_1} \cdot \ldots \cdot R_{p_k})^*$ using PSN_CE.

The control expression for a program essentially implements Pred-SCC by Pred-SCC evaluation, with rule ordering within the loop to evaluate a Pred-SCC. It is easy to show that the semantics of the control expression is equivalent to the fixpoint semantics of the program. From Theorem 4.1, it follows that the evaluation of the control expression is complete, and has the non-repetition property. Thus we have the following theorem (see [21] for a formal proof of correctness).

Theorem 4.2 Procedure PSN_Prog has the non-repetition property, is sound, and if all the predicates defined in the program are safe, is complete wrt the least fixpoint semantics. \Box

Just as the ordering of rules could affect the performance of GSN, ordering of predicates could affect the performance of PSN. The results of Section 5 can be used to obtain good predicate orderings for the PSN evaluation of a program.

In a sequential evaluation of a program, PSN evaluation is *always* preferable to BSN evaluation, since it can be implemented with the same overheads per iteration, but can do better in terms of the number of iterations. The results of Leask et al. [16], who did a performance evaluation of BSN, PSN and some parallel implementations of BSN using total elapsed time as a metric on a disk-based system, also indicate that PSN is the evaluation strategy of choice, in the absence of semantic information about the data in the base relations. PSN may not be able to utilize facts as early as GSN can, but the overheads associated with GSN could be higher, and the choice of which strategy to choose is not always obvious.

The following example illustrates the differences between BSN, PSN, and GSN evaluation of a program.

Example 4.1 Consider the following program *P*:

$$\begin{split} R_1 &: p(X,Y) := b_1(X,Y). \\ R_2 &: q(X,Y) := b_2(X,Z), p(Z,Y). \\ R_3 &: p(X,Y) := b_3(X,Z), q(Z,Y). \\ R_4 &: p(X,Y) := b_4(X,Z), p(Z,Y). \\ & b_1(4,5). \quad b_2(3,4). \quad b_3(2,3). \quad b_4(1,2).. \\ \text{Query: } ?-p(1,X). \end{split}$$

For the above program, BSN as well as GSN and PSN would evaluate the non-recursive rule, R_1 exactly once (in the beginning) to produce p(4, 5).

Evaluating the rules (R_2, R_3, R_4) repeatedly in this order, using GSN, would produce q(3, 5)using R_2 in the first iteration, and make it immediately available to applications of R_3 and R_4 in the same iteration. Rule R_3 would then produce a fact p(2, 5) in the first iteration, using which R_4 would produce a fact p(1, 5) in the first iteration. Thus, the answer, p(1, 5), to the query would be produced at the end of the first iteration itself using GSN evaluation of P.

If the predicate ordering chosen for PSN is (q, p), the rule defining $q(R_2)$ is applied first, and the fact (q(3, 5) in the first iteration) produced would be immediately available to applications of the rules defining $p(R_3 \text{ and } R_4)$ in the same iteration. Rules R_3 and R_4 are then applied independently, and hence the fact produced using R_3 (p(2, 5) in the first iteration) would be available to R_4 only in the next iteration. Thus p(1,5), the answer to the query, would be produced at the end of the second iteration using PSN evaluation of P.

If BSN evaluation is used instead, facts produced by a rule application would not be available to any other rule application in the same iteration. Consequently, q(3,5) would be produced in the first iteration, p(2,5) in the second iteration, and p(1,5) (the answer to the query) would be produced only in the third iteration. \Box

5 Rule Orderings that Minimize Rule Applications

Example 5.1 Let q and r be base relations. Consider the following program P:

$$R_{0}: p_{k}(X) := q(X).$$

$$R_{1}: p_{1}(f_{1}(X)) := p_{k}(X).$$

$$\dots$$

$$R_{k-1}: p_{k-1}(f_{k-1}(X)): - p_{k-2}(X).$$

$$R_{k}: p_{k}(f_{k}(X)) := -p_{k-1}(X), r(X).$$

The non-recursive rule R_0 is the first rule to be applied in a BSN evaluation of P. In an iteration of a Basic Semi-Naive evaluation, all the recursive rules (i.e. R_1, \ldots, R_k) of the program are applied independently. Rule R_k will be successfully applied for the first time only in the k^{th} iteration. However, it would be possible to successfully apply rule R_k in the first iteration itself if the rules are applied in the order shown, i.e., (R_1, R_2, \ldots, R_k) , and the facts produced by each rule application are immediately made available to subsequent rule applications. If the computation of a fact using this technique (and the given ordering of rules) took n iterations, then computation of the same fact using the BSN evaluation strategy could take up to k * n iterations.

If, instead, the rules are applied in the opposite order, i.e., $(R_k, R_{k-1}, \ldots, R_1)$, the number of iterations taken is the same as BSN evaluation, even if facts produced by a rule application are made available immediately after the rule is applied. Thus we see the importance of a good ordering of rules. \Box

In this section, we provide a theoretical analysis of how the number of rule applications (and iterations) in fixpoint algorithms, that have the non-repetition property and apply a rule using GSN, can be reduced through the use of rule ordering. Our results are significant in that they indicate how this number can be minimized, independent of the data in base relations, over a significant class of rule orderings (Section 5.3). We also present results which suggest that only this class of rule orderings should be considered in the absence of additional semantic information (Section 5.4).

The techniques described in this section deal with rule orderings, but can be extended, in a straightforward fashion, to deal with predicate orderings.

5.1 Benefits of Rule Ordering

The number of inferences has been widely used as a cost metric in the evaluation of logic programs. However, any evaluation technique that has the non-repetition property makes each inference (that can be made) exactly once, and hence all the techniques we study are equivalent under this criterion.

One of the advantages of bottom-up evaluation of logic programs is the increased degree of set-oriented computation. Given that the total number of inferences made by two different evaluation techniques is identical, the technique that performs more set-oriented computation is expected to perform better in terms of the number of I/O operations.

Our theoretical analysis and performance results show that rule ordering can greatly reduce the number of rule applications, and therefore the number of joins in bottom-up evaluation, without making additional inferences. This has two benefits:

- 1. It reduces the overall cost due to constant per-join overheads such as initialization costs, and per iteration overheads such as updating the various predicate extensions.
- 2. It increases the degree of set-orientedness. The fewer the number of rule applications performed, the greater is the number of inferences made in a single rule application. This increases the degree of set-orientedness, and hence decreases the number of I/O operations.

The reduction in cost due to ordering of rules is orthogonal to other techniques such as efficient join and indexing strategies, and duplicate elimination techniques—none of these is made inapplicable by ordering rules.

5.2 Class of Orderings Considered

Definition 5.1 Fair, static orderings : Let the rules of a Rule-SCC S be R_1, \ldots, R_n . A fair, static ordering is an ordering of the form $(R_{i_1}, \ldots, R_{i_n})$, where i_1, \ldots, i_n is a permutation

of $1, \ldots, n$. \Box

We consider such orderings in Section 5.3. Such an ordering is referred to as a *static* ordering since the same ordering is used in each iteration. In such static orderings no rule is applied more often than other rules. Such orderings are referred to as *fair* orderings since in the absence of any prior knowledge of the frequency with which different rules are used, or other semantic information, we have no basis for applying some rules more often than others. In order to compute the closure of an SCC S, using a fair ordering, we apply the rules in the given order in each iteration, repeatedly, making the facts produced by a rule application available to all subsequent rule applications, till no more facts can be computed. Independent applications of rules are not considered, since making the facts produced by a rule application available to other rule applications may significantly reduce the number of rule applications needed to compute the fixpoint, and can never increase it in a sequential evaluation.

In Section 5.4 we consider static orderings in which some rules can be applied more often than other rules. Such orderings are referred to as *non-fair* orderings. This class includes the class of nested orderings, such as those considered by Kuittinen et al. [15]. Non-fair orderings may perform somewhat better than fair orderings on some data sets, but, as we show in Section 5.4, such orderings may also perform considerably worse on other data sets. Hence, in the absence of any information about the kind of data sets, fair orderings are preferable.

We do not consider orderings where the next rule to be applied is chosen dynamically. Although *dynamic* orderings may be better than static orderings in terms of the cost criteria described in Section 5.1, determining which rule to apply may be difficult and involve considerable overheads. While dynamic orderings may be worth investigating, we do not consider them in this paper.

5.3 Fair Orderings

Definition 5.2 Order sequence : Let O be any ordering $(R_{i_1}, \ldots, R_{i_n})$. An order sequence $S = O^m$ denotes the string formed by repeating O, m times.

We use the notation $S[k], 1 \le k \le m * n$ to denote the k^{th} rule in S. \Box

A sequence S_1 of length n is said to be a *subsequence* of a sequence S_2 of length $m \ge n$, if there exist n numbers $1 \le k_1 < k_2 < \ldots < k_n \le m$, such that $S_1[i] = S_2[k_i], 1 \le i \le n$. For example, (R_1, R_3, R_5) is a subsequence of $(R_1, R_2, R_3, R_4, R_5)$. **Definition 5.3 Cycle preserving fair orderings :** Consider an SCC *S*, and let the rules in *S* be $\mathcal{R} = \{R_1, \ldots, R_n\}$. Let $G = (\mathcal{R}, E)$ be the rule graph for the given SCC. Let *O* be any fair ordering $(R_{i_1}, \ldots, R_{i_n})$ of the rules in *R*. Let *C* be any simple cycle R_{j_1}, \ldots, R_{j_m} in *G*.³

We say that a fair ordering O preserves a cycle C, if there is a cyclic permutation O_1 of O such that C forms a subsequence of O_1 . A fair ordering O on G is a cycle preserving fair ordering if for every simple cycle C in G, O preserves C. \Box

A fair ordering O that does not preserve a cycle C is said to break it. A cycle C is broken by degree B(C, O) = i, by a fair ordering O, if i is the least number such that for some cyclic permutation O_1 of O, C is a subsequence of O_1^i . Thus a fair ordering that preserves a cycle can be said to break it by degree one.

Not every graph has a cycle preserving ordering, and it is not hard to construct graphs that have no cycle preserving ordering. A complete directed graph with three vertices is one such. In order to compare rule orderings for all rule graphs, we define a relation \triangleleft on the class of fair orderings. Given two fair orderings O_1 and O_2 on a rule graph G, $O_1 \triangleleft O_2$ if for every simple cycle C in G, $B(C, O_1) \leq B(C, O_2)$. If we have $O_1 \triangleleft O_2$ and $O_2 \triangleleft O_1$, we say that the two orderings are *equivalent*. The use of this relation will be seen in Section 5.3.1, where we show in Theorem 5.2 that if $O_1 \triangleleft O_2$ and the two are not equivalent, then given any database, O_1 is better than O_2 based on the number of iterations needed to compute the closure of an SCC. We also show that if O_1 and O_2 are equivalent, the number of iterations needed by each to compute the closure of an SCC differ by at most a data-independent constant. Thus, we show that an ordering that preserves all cycles is optimal in the class of fair orderings, under this cost criterion. From the definition of the relation \triangleleft we have:

- Cyclic permutations of a fair ordering are equivalent under \triangleleft , and
- Any two cycle preserving fair orderings are equivalent under \triangleleft .

Example 5.2 Consider the rule graph shown in Figure 1. The simple cycle 1, 3, 2, 6 is preserved by the ordering (2, 6, 4, 1, 3, 5) because the ordering has a cyclic permutation (1, 3, 5, 2, 6, 4) which has the simple cycle as a subsequence. However, this ordering breaks the simple cycle 1, 4, 2, 5 by degree 3. \Box

 $^{^{3}}$ Though cycles have the same initial and final vertex, we omit the final vertex in our representation, for convenience.



Figure 1: An Example Rule Graph

Lemma 5.1 Consider a cycle $C = R_1 \dots R_m$ and a fair ordering O. Let O_1 be the cyclic permutation of O that ends with R_m . Then C forms a subsequence of O_1^i for i = B(C, O), but not for any smaller i.

Proof: (Sketch) By definition of B(C, O), for some cyclic permutation O_r of O, C is a subsequence of O_r^i for i = B(C, O), but not for any smaller i. We now take any arbitrary O_r^i as above, and move elements after the last occurrence of R_m to the head of the sequence, to get O_1^i . \Box

We present proof sketches for many of the results in this section. Full proofs may be found in [21].

For the class of fair orderings, we next show that cycle preserving fair orderings are optimal under the cost criterion of the number of iterations needed to compute the closure of an SCC, with an immediate update strategy. Since the number of rule applications is constant within an iteration in a fair ordering, the optimality result carries over for the cost criterion of number of rule applications.

5.3.1 Optimality of Cycle Preserving Orderings

Definition 5.4 Derivation path : A *derivation path* for a fact is a path in a derivation tree for the fact, starting from a leaf node. We represent such a path concisely by listing the rules labeling the nodes in the derivation path in order, starting from the parent of the leaf. \Box Note that two different paths may have the same representation, but that does not affect our analysis.

Definition 5.5 Derivation index : Let O denote a fair ordering of rules in the program and T denote a particular derivation tree for $p(\overline{c})$. Consider the rule application sequence $\mathcal{O}' = O^j$, for arbitrarily large j. With each node in the derivation tree T, we associate a *derivation index*, which is an index into the sequence \mathcal{O}' ,

Leaf nodes (corresponding to base facts) are associated with the derivation index zero. The derivation index of each internal node n', labeled with a derived fact p' and a rule R', is the minimum possible k such that, $\mathcal{O}'[k] = R'$ and the derivation indices of the children nodes of n' are less than k. \Box

Definition 5.6 Iteration height : Let O denote a fair ordering of rules in the program and T denote a particular derivation tree for $p(\overline{c})$. With each node in the derivation tree we associate an *iteration height* $\lceil k/m \rceil$ where k is the derivation index of that node, and m is the length of O.

A derivation tree is said to be *computed* by O using n iterations if the iteration height of the root of the tree under O is n. \Box

The iteration height of a node is defined syntactically but has the following semantic interpretation. If the iteration height of the root of a derivation tree T is n, then the corresponding fact $p(\overline{c})$ is computed in or before the nth iteration of the application of rules according to the ordering O. If the fact is computed in the nth iteration, there is a derivation tree with iteration height n for the fact.

Definition 5.7 Iteration count : The *iteration count* of a fact, for a given fair ordering, is defined to be the minimum of the iteration heights under the given ordering, of derivation trees for this fact. \Box

This gives us the earliest iteration in which the fact is derived. This link between the semantic notion of the number of iterations needed to compute a fact, and the iteration heights of derivation trees for the fact enables us to argue about the computation of facts using purely syntactic criteria.

Definition 5.8 Iteration length : Consider a derivation tree T for a fact, and a fair ordering O. Given a derivation path s in T, the *iteration length* L(s, O) of the path is the minimum n, such that the path forms a subsequence of O^n .

The minimum length order sequence for s is defined to be $O^{L(s,O)}$. We next show the relationship between the notion of the iteration length of a path, and the iteration height of a tree. For a tree T, if T has no internal nodes, define L(T,O) = 0. Otherwise define $L(T,O) = \max\{L(s,O) \mid s \text{ is a path in } T\}$.

The following lemma permits us to argue about the number of iterations it takes to compute a derivation tree based on the iteration lengths of the derivation paths in the tree.



Figure 2: Path with Two Alternative Cycle Sequence Lengths

Lemma 5.2 Given a derivation tree T for a fact $p(\overline{c})$, and a fair ordering O, the derivation tree can be computed by a bottom-up fixpoint evaluation using rule ordering O in L(T, O) iterations. L(T, O) is thus also the iteration height of T.

Proof: (Sketch) By definition of iteration height, a tree is computed by a bottom-up fixpoint evaluation in as many iterations as its iteration height. We prove by an induction on the height of trees that the iteration height of a node is the same as the maximum of the iteration lengths of paths from the leaves of the tree to that node. \Box

The above lemma lets us reason in terms of derivation paths rather than derivation trees. We next prove certain properties of derivation paths. The next lemma proves one such property, and is used in turn to prove the result in Theorem 5.1.

Lemma 5.3 For every derivation path s, there exists a sequence s_0, \ldots, s_n of paths in the rule graph G, such that, (1) $s = s_n$. (2) s_0 is an acyclic path in G. (3) For each i > 0, s_i can be constructed from s_{i-1} as follows: Choose a rule R_{j_k} in s_{i-1} , and a simple cycle $C_i = R_{j_1}, \ldots, R_{j_k}$ in G, and insert the cycle just after R_{j_k} .

Proof: (Sketch) The proof is by induction on lengths of paths, and can be easily understood by reversing the process: given a path it is easy to find a simple cycle in it unless it is acyclic. Just start from the beginning, go down the path until any node is repeated; the portion between the repeated nodes, including one repetition of the node, is a simple cycle. By removing the cycle and repeating this process until there are no more cycles, we get a sequence of paths in reverse order, and an acyclic path. \Box

For any derivation path s, such a sequence s_0, \ldots, s_n is called a *construction sequence* for s. A derivation path may not have a unique construction sequence, and different construction sequences can have different lengths, as the following example shows.

Example 5.3 Consider the graph shown in Figure 2. The derivation path 1, 2, 3, 1, 3, 2 has the following two construction sequences associated with it,

- 1. (1,3,2), (1,2,3,1,3,2), which is constructed by inserting the simple cycle 1, 2, 3, 1 into the acyclic path 1, 3, 2 (note that the repeated node 1 in the simple cycle is not inserted).
- 2. (1,2), (1,2,3,2), (1,2,3,1,3,2), which is constructed by inserting the simple cycle 2,3,2 into the acyclic path 1,2, and then inserting the cycle 3, 1, 3 into the result (the repeated nodes in the cycles are not inserted). □

Given a fair ordering O, we now relate the iteration length of a derivation path s with the length of the construction sequence for s and the degree by which the given ordering breaks each of the cycles inserted.

Theorem 5.1 Consider any derivation path s and a construction sequence s_0, \ldots, s_n for s as defined in Lemma 5.3. Let C_i be the cycle inserted in obtaining s_i from s_{i-1} . For every fair ordering O, the iteration length of s under O is given by $L(s, O) = L(s_0, O) + \sum_{i=1}^{n} B(C_i, O)$. Further, $L(s_0, O)$ is bounded by the length of the longest acyclic path in the rule graph of the SCC.

Proof: (Sketch) The proof is by induction on the number of cycles in the construction sequence of the derivation path. The basis case is when the number of cycles is zero, and is trivial. Assume the theorem for all paths with construction sequences having less than k cycles, and consider a path s with a (minimum length) construction sequence of k cycles. Suppose it has a minimal length order sequence $o = O^j$. We remove one cycle c from s to get s', and using the induction hypothesis we get an order sequence o' for s' with the required length. Suppose the cycle started with node n, and the previous node in s was mapped to some instance rule m in o'. We cut o' just after this instance of m. We show using minimality arguments that the initial part of o' must be the same as an initial part of o, and the tail part of o' must be the same as a tail part of o. What remains of o is now the part in between. This must end with rule m, and must be an order sequence for c. By the definition of B(c, O) this must be a cyclic permutation of $O^{B(c,O)}$. This then gives us the iteration length of the minimal length order sequence for s, which completes the induction step. \Box

Note the interesting fact that the above theorem is true for any construction sequence. Since the actual iteration length of a path does not depend on the construction sequence chosen, this tells us that, in a certain sense, all construction sequences are equivalent. For any cycle preserving fair ordering O with a construction sequence t_0, \ldots, t_n , $L(t_n, O) = L(t_0, O) + n$.

Consider a rule graph G, and two fair orderings O_1 and O_2 such that $O_1 \triangleleft O_2$. We define

 $MaxR(O_1, O_2, G) = max\{B(C, O_2)/B(C, O_1) \mid C \text{ is a simple cycle in } G\}.$

This serves as a bound on how much costlier, based on the number of iterations, O_2 can be compared to O_1 .

Given any two fair orderings that are related by the \triangleleft relation, we wish to compare the number of iterations taken to compute the closure of an SCC by the two orderings. To this end, we first compare the iteration lengths of derivation paths. This is used to compare the iteration heights of derivation trees for a fact. We then argue about the number of iterations taken to derive a fact by the two orderings, by comparing the iteration counts of the fact. This leads finally to our main result, stated in Theorem 5.2, that relates the number of iterations taken to compute the closure of the SCC by the two fair orderings.

Lemma 5.4 Consider any two fair orderings O_1 and O_2 for a graph G, such that $O_1 \triangleleft O_2$. For any derivation path s, the iteration lengths of s under the two orderings are related as $L(s, O_1) - k \leq L(s, O_2) \leq MaxR(O_1, O_2, G) \cdot L(s, O_1) + k$, where k is bounded by the length of the longest acyclic path in G.

Proof: (Sketch) We get a construction sequence for s, and use Theorem 5.1 to express the iteration lengths in terms of the length of the construction sequence for s, and the degree to which cycles are broken by the two orderings. The lemma then follows from the definition of MaxR. \Box

Lemma 5.5 Consider any two fair orderings O_1 and O_2 for a graph G, such that $O_1 \triangleleft O_2$. For any derivation tree T, the iteration heights of T under the two orderings are related as $L(T, O_1) - k \leq L(T, O_2) \leq MaxR(O_1, O_2, G) \cdot L(T, O_1) + k$, where k is bounded by the length of the longest acyclic path in G.

Proof: (Sketch) The proof follows from Lemma 5.2, which links iteration heights directly to iteration lengths, and Lemma 5.4, which shows a result similar to that of this lemma, for iteration lengths. \Box

Lemma 5.6 Given any fact $p(\overline{c})$, and any two fair orderings O_1 and O_2 for a graph G, such that $O_1 \triangleleft O_2$, let the iteration counts of $p(\overline{c})$, for O_1 and O_2 , be n_1 and n_2 , respectively. n_1 and

 n_2 are related as $n_1 - k \le n_2 \le MaxR(O_1, O_2, G) \cdot n_1 + k$, where k is bounded by the length of the longest acyclic path in G.

Proof: (Sketch) A fact can have several derivation trees. We choose derivation trees with minimum iteration heights under each of the two orderings. Using Lemma 5.5 with the two derivation trees we directly show the two inequalities in this lemma. \Box

Theorem 5.2 Given an SCC S, any two fair orderings O_1 and O_2 for the rule graph of S, such that $O_1 \triangleleft O_2$, and any set of base facts, let the number of iterations required to compute the closure of S by bottom-up fixpoint evaluations using rule orderings O_1 and O_2 be n_1 and n_2 respectively. n_1 and n_2 are related as $n_1 - k \leq n_2 \leq MaxR(O_1, O_2, G) \cdot n_1 + k$, where k is bounded by the length of the longest acyclic path in the rule graph for the SCC.

Proof: (Sketch) We find facts that are computed last under each of the orderings, and using Lemma 5.6 we show the two inequalities in this theorem. \Box

Corollary 5.1 Given any two cycle preserving fair orderings, the number of iterations required to compute the closure of an SCC by bottom-up fixpoint evaluations using the two orderings differ by at most a (data-independent) constant. Also, the number of rule applications required by the two orderings differ by at most a (data-independent) constant. \Box

5.3.2 Generating Cycle Preserving Fair Orderings

An SCC with n rules could have an exponential number of simple cycles. A naive algorithm for generating cycle preserving orderings would examine all O(n!) orderings and all simple cycles to check if the ordering preserves every simple cycle. Finding an efficient algorithm to construct orderings that are minimal under the \triangleleft relation is an open problem, as is the special case of an algorithm for finding cycle preserving orderings. Finding what complexity classes these problems belong to is also an interesting open problem.

As a heuristic, we suggest using the reverse of a depth-first search pop-out order of the rule graph as a rule ordering. This often generates cycle preserving orderings, but not always. For examples of depth-first search pop-out orders, see [21].

5.4 Non-Fair Orderings

In this section we consider static orderings in which some rules may be applied more often than other rules. We divide this class into the class of flat orderings and the class of non-flat nested orderings.

Definition 5.9 Nested ordering : A *nested ordering* is an ordering O of the form (O1), where O1 is generated by the grammar

$$O1 \rightarrow R_1 \mid \ldots \mid R_n \mid O1 \cdot O1 \mid (O1)^*$$

where R_1, \ldots, R_n are the rules of an SCC S of the rule graph, such that each rule in the SCC occurs at least once in the ordering O. \Box

An example is the ordering $(R_1 \cdot R_2 \cdot (R_3 \cdot R_4)^* \cdot R_5)^*$. Note that a nested ordering can have more than one occurrence of any rule in the SCC.

Definition 5.10 Flat ordering : A *flat ordering* is a nested ordering that has parentheses only at the outermost level. \Box

The *nesting level* of nested orderings is defined in the obvious manner, where a flat ordering is defined to have a nesting level of one. We have the following lemma, whose proof is straightforward.

Lemma 5.7 Consider an SCC S, and any flat ordering O of the rules in S. Let k_1 be the number of rule occurrences in O. Let n_{opt} be the minimum number of rule applications needed to compute the closure of S on a given database, using a sequential evaluation with immediate updates. If n_O denotes the number of rule applications needed to compute the closure of S on the same database, using the ordering O, then $n_O \leq n_{opt} \cdot k_1$. \Box

We now compare flat orderings with nested orderings.

Lemma 5.8 Consider an SCC S, and let O be any nested ordering with k_1 rule occurrences and a nesting level of s. Let $iter_{bsn}$ be the number of iterations needed to compute the closure of S using BSN. If n_O denotes the number of rule applications needed to compute the closure of S using the ordering O, then $n_O \leq (iter_{bsn})^s \cdot k_1$.

Proof: (Sketch) No loop in the nested ordering can iterate more than $iter_{bsn}$ times after it is entered, since $iter_{bsn}$ is the length of the longest derivation path. The result then follows from the nesting level of the ordering. \Box

From the previous two lemmas we have the following theorem, which summarizes our comparison of flat and nested orderings.

Theorem 5.3 Consider an SCC S, and any flat ordering O_f and any nested ordering O_n on S. Let $iter_{bsn}$ be number of iterations needed to compute the closure of S using BSN. If n_f and

 n_n denote the number of rule applications needed to compute the closure of S using O_f and O_n respectively, then $n_f/k \leq n_n \leq (iter_{bsn})^s \cdot k_1$, where k is the number of rule occurrences in O_f , k_1 is the number of rule occurrences in O_n , and s is the nesting level of O_n .

Proof: The number of rule applications using any flat ordering is bounded as $iter_{bsn} \leq n_f$, where $iter_{bsn}$ is the number of iterations needed to compute the closure of S using BSN. Clearly $n_{opt} \leq n_n$, and the theorem follows from Lemma 5.7 and Lemma 5.8. \Box

Since an optimal fair ordering must take at least as many rule applications as an optimal rule ordering, the above theorem also directly bounds how much worse an arbitrary flat ordering can be compared to an optimal fair ordering.

Note that every fair ordering is also a flat ordering, and hence the above theorem applies when we compare fair orderings with nested orderings. The worst case performance of nested orderings is bounded, as shown above. In Section 6 we describe an example where the performance of a nested ordering is indeed as bad (to within a small constant factor) as the above upper bound allows (Program P2, data set S64). Thus, although a nested ordering can perform somewhat better (i.e. $n_f = c * n_n$, $1 < c \leq k$) than fair orderings on some data sets, it is possible for it to perform much worse (when $n_n = (iter_{bsn})^s .k_1$) on other data sets. Note also that $iter_{bsn} \geq n_f/k$.

6 Performance Results

In this section we summarize the results of a performance study of the benefits of immediate availability of facts, and the benefits of ordering rules as described in Section 5. Our performance study draws upon and extends the work of Kuittinen et al. [15].

We consider two programs, referred to as P1 and P2 in this section.⁴ Mostly, we discuss P1, and discuss P2 only when it offers some additional insight. For a listing of these programs, our datasets, and further details of our performance evaluation, see [21].

The above programs were hand-coded for each of the evaluation techniques, and measurements were made by running the resultant programs on several data sets (which are described in [21]. There is a cycle preserving fair ordering for each SCC of P1 and P2, and the column "General 1" of the tables is for a GSN evaluation using such an ordering. The column "General 2" in the tables for P1 corresponds to a GSN evaluation of P1 using a fair ordering that breaks a

 $^{{}^{4}}P1$ is the same program that was used in [15].

Data Set	Basic	Pred	General 1	General 2
A10	3579	1535	1023	2812
<i>B</i> 64	146	73	68	80
F10	23	10	7	18

Table 4: Program P1: Number of Iterations

Data Set	Basic		Pred		General 1		General 2	
	Non-Nl	Null	Non-Nl	Null	Non-Nl	Null	Non-Nl	Null
A10	6126	14835	6126	4615	5230	2693	6126	11000
B64	1066	196	549	88	588	14	596	96
F10	73	78	57	17	45	10	73	53

Table 5: Program P1: Number of Joins (Non-Null and Null)

cycle to degree six. Column "Basic" corresponds to a Basic Semi-Naive evaluation, and column "Pred" is for a Predicate Semi-Naive evaluation using a cycle preserving fair ordering based on the predicate graph.

For P1 we use the data sets A10, B64, and F10. Data set A10 results in no duplicate derivations with P1, but takes a large number of iterations. Data set B64 is large, takes a moderate number of iterations, and results in a moderate number of duplicates. Data set F10results in a large number of duplicate derivations, but a fewer number of iterations.

For P2 we use two data sets, C16 and S64. C16 is designed such that the nested ordering we consider performs well, and S64 is designed such that the nested ordering performs very badly compared to the cycle preserving fair ordering.

Table 4 shows the number of iterations taken by each evaluation strategy on P1. It should be noted that the total number of rule applications (considered) is directly proportional to the number of iterations taken. PSN outperforms BSN on this measure. It improves over the performance of BSN by over 50% on all data sets considered. GSN with a cycle preserving fair ordering outperforms PSN by about 30% (on A10 and F10) and performs about 50% to 70% better than BSN. GSN with a bad fair ordering performs much worse than GSN with a good fair ordering (although it can never be worse than BSN), and this clearly brings out the benefits of good fair orderings.

Data Set	Eval. Tech.	Non-Nl Joins	Hashed	Constant	Logarithmic	Linear
A10	Basic	6126	1	1	1	1
A10	Pred	6126	1.000	1.000	1.000	1.000
A10	General 1	5230	0.856	1.000	1.000	1.000
<i>B</i> 64	Basic	1066	1	1	1	1
<i>B</i> 64	Pred	549	0.525	1.000	0.999	1.000
B64	General 1	588	0.542	1.000	0.997	1.000
<i>F</i> 10	Basic	73	1	1	1	1
F10	Pred	57	0.900	1.000	1.004	1.001
F10	General 1	45	0.822	1.003	1.013	1.025

Table 6: Program P1: Normalized Joins Costs (Without Overheads)

If one of the relations in the join is empty, the result of the join is null, and we call such a join a *null join*. We may be able to detect this condition at run time without incurring much cost. Table 5 shows the number of joins used by each evaluation strategy on P1, divided into the number of null joins, and the number of non-null joins. The total number of joins taken by General 1 is better than PSN, which in turn is better than BSN. If we count only non-null joins, this is not strictly true. PSN always performs no worse than BSN, and on *B*64 performs about 50% fewer non-null joins. General 1 performs 15% to 45% better than BSN on this count. On one data set, *B*64, PSN is slightly better (less than 7%) than General 1. However, on the other two data sets, General 1 outperforms PSN by about 15% to 20%.

The cost of a single join can be modeled in different ways, depending on the join strategy used. In Table 6 we present the portion of the join cost that excludes the per-join overhead. To get the full join cost, we must add the overhead per join times the number of (non-null) joins. In these tables, the column "Hashed" is a measure of the join cost (excluding overheads) using the hash join technique. The column "Constant" corresponds to a nested loop join with indices that can be accessed in constant time (for instance, a hash index), the column "Logarithmic" corresponds to nested loops join with a logarithmic access time index (for instance, a B-tree index), and the column "Linear" corresponds to a nested loop join without indices (which is equivalent to an index with access time linear in the size of the relation). All these costs are normalized (with Basic = 1 for each technique and data set) order of magnitude estimates

derived directly from the sizes of the relations. The size of the result of the join is included in the join cost. Note the following important points:

- I/O costs and overhead costs for joins are not measured in these tables. The number of non-null joins may be used to estimate the total overhead and this should be added to the cost in the table. Since the overheads are implementation dependent and we currently lack an actual system to measure the overheads, we do not present the total cost. Some results comparing PSN with BSN using total elapsed time on a disk based system are presented in [16].
- The only comparisons that may be made from this table are the differences between the different evaluation strategies. These numbers should not be used to directly compare different join techniques (i.e. the numbers in a single row of the table should not be compared).

The performance study of Kuittinen et al. [15] only models the join cost in one way, the hashed join method.

The performance results in Table 6 indicate that for the hash join strategy the total join cost closely parallels the number of non-null joins. PSN and GSN do consistently better than BSN under this metric. The performance results also indicate that under other join cost assumptions (nested loops without indices, and with constant or logarithmic access time indices), the total join costs apart from the I/O costs and other overheads associated with each join, are practically the same for each of the fixpoint evaluation strategies we study (BSN, PSN, and GSN). In these cases the number of non-null joins becomes the significant factor.

The cost of checking for and eliminating duplicates could form a significant part of the total cost of evaluation. We studied duplicate elimination costs under various assumptions about the indexing costs (constant, logarithmic, linear), but without a detailed model of I/O costs. Since the costs for BSN, PSN and GSN varied by less than 5%, we omit these results.

From Table 7, for the number of rule applications, it can be seen that for C16, Nested is better than the fair ordering indicated by General 1. However, for S64 Nested performs much worse than any of the other evaluation strategies. Both PSN and General 1 are about 20% to 65% better than BSN. This pattern continued to hold approximately when we compared Nested and other strategies with respect to the number of joins.

Data Set	Basic	Pred	General 1	Nested
<i>C</i> 16	282	221	207	179
S64	1717	588	583	2536

Table 7: Program P2: Number of Rule Applications

Our performance results underscore the theoretical results described in earlier sections. The benefits of immediate availability of facts is indicated by the fact that GSN is in general better than PSN, which, in turn, is in general better than BSN, under the cost criterion of number of rule applications and iterations. Further, our results clearly bring out the advantages of cycle preserving fair orderings. Our results also indicate that nested orderings may perform better than fair orderings on some data sets, but can perform much worse on others.

7 Related Work

The idea of semi-naive, or *differential*, evaluation has been independently rediscovered many times ([9, 19, 5, 2, 1]).

The most closely related work is that of Kuittinen et al. [15], who propose a fixpoint evaluation algorithm for logic programs based on the immediate utilization of facts. Their algorithm also reduces the number of iterations, and is dependent upon a choice of rule orderings. They present a performance study of the effects of rule orderings, although they do not analyze the effect of rule orderings theoretically. The results presented in our paper and in [15] were obtained independently. (However, we have drawn upon and extended their performance evaluation.) Although the technique of Kuittinen et al. avoids repeating most derivations, it does not have the non-repetition property since it is possible for some derivations to be repeated. This can adversely affect performance. Further, it makes the algorithm inapplicable when the non-repetition property is used to perform further optimizations. (For example, if a program has the *duplicate-freedom* property, a fixpoint algorithm with the non-repetition property can be modified easily to eliminate run-time checks for duplicates [18].) Finally, their technique handles only a subset of the rule orderings that GSN can handle. Schmidt [23] presents another evaluation technique that allows for some ordering of rules; however, this technique also lacks the non-repetition property.

Gonzalez-Rubio, Rohmer and Bradier [10] have independently proposed a parallel evaluation

technique that has some similarity to GSN evaluation. The idea is to run a copy of the BSN algorithm on each processor, and to exchange facts as needed. A fact received from another processor is treated exactly as if it were derived locally in an iteration. It is possible to modify their technique and obtain an algorithm similar to GSN, although they do not address the issue of ordering rules.

Other related work includes a scheme for ordering facts in a semi-naive evaluation proposed by Schmidt et al. [24]. In the context of transitive closure, Lu [17], presents a technique for facts to be (partially) utilized in the same iteration that they were generated. There is also some connection to work on parallelizing transitive closure by using different expansions of the recursive rule. For a more detailed discussion of related work, see [21].

8 Summary

In this paper, we studied several aspects of rule ordering in the bottom-up evaluation of logic programs. Rule orderings are necessary for ensuring a desired semantics, such as the evaluation of the magic rewritten versions of stratified programs. We presented two evaluation algorithms, GSN and PSN, that could be used for evaluating such rule orderings, while preserving the non-repetition property, and discussed cases where each was useful.

We studied rule orderings theoretically, and showed that for the class of fair orderings, cycle preserving orderings were optimal and, in the absence of additional information, fair orderings are to be preferred to non-fair orderings. An important open problem is to find an efficient algorithm that checks whether an SCC has a cycle preserving fair ordering and if so, produces it.

We also presented a performance study to support our theoretical analyses. Rule orderings were also shown to be useful for improving the total cost of sequential evaluation of logic programs.

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