Mining Tree-Query Associations in Graphs

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Abstract

New applications of data mining, such as in biology, bioinformatics, or sociology, are faced with large datasets structured as graphs. We introduce a novel class of tree-shaped patterns called tree queries, and present algorithms for mining tree queries and tree-query associations in a large data graph. Novel about our class of patterns is that they can contain constants, and can contain existential nodes which are not counted when determining the number of occurrences of the pattern in the data graph. Our algorithms have a number of provable optimality properties, which are based on the theory of conjunctive database queries. We propose a practical, database-oriented implementation in SQL, and show that the approach works in practice through experiments on data about food webs, protein interactions, and citation analysis.

1 Introduction

The problem of mining patterns in graph-structured data has received considerable attention in recent years, as it has many interesting applications in such diverse areas as biology, the life sciences, the World Wide Web, or social sciences. In the present work we introduce a novel class of patterns, called tree queries, and we present algorithms for mining these tree queries and tree-query associations in a large data graph. This article is based on two earlier conference papers [17, 20].

Tree queries are powerful tree-shaped patterns, inspired by conjunctive database queries [18]. In comparison to the kinds of patterns used in most other graph mining approaches, tree queries have some extra features:

- Patterns may have “existential” nodes: any occurrence of the pattern must have a copy of such a node, but existential nodes are not counted when determining the number of occurrences.

- Moreover, patterns may have “parameterized” nodes, labeled by constants, which must map to fixed designated nodes of the data graph.

- An “occurrence” of the pattern in a data graph \( G \) is defined as any homomorphism from the pattern in \( G \). When counting the number of occurrences, two occurrences that differ only on existential nodes are identified.

Past work in graph mining has dealt with node labels, but only with non-unique ones: such labels are easily simulated by constants, but the converse
is not obvious. It is also possible to simulate edge labels using constants. To simulate a node label $a$, add a special node $a$, and express that node $x$ has label $a$ by drawing an edge from $x$ to $a$. For an edge $x \rightarrow y$ labeled $b$, introduce an intermediate node $x.y$ with $x \rightarrow x.y \rightarrow y$, and label node $x.y$ by $b$.

A simple example of a tree query is shown in Figure 1(a) when applied to a food web: a data graph of organisms, where there is an edge $x \rightarrow y$ if $y$ feeds on $x$, it describes all organisms $x$ that compete with organism #8 for some organism as food, that itself feeds on organism #0. This pattern has one existential node, two parameters, and one distinguished node $x$. Figure 1(b) shows another example of a tree query; when applied to a food web, it describes all organisms $x$ that have a path of length four beneath them that ends in organism #8.

Effectively, tree queries are what is known in database research as conjunctive queries [9, 40, 2]; these are the queries we could pose to the data graph (stored as a two-column table) in the core fragment of SQL where we do not use aggregates or subqueries, and use only conjunctions of equality comparisons as where-conditions. For example, the pattern of Figure 1(a) amounts to the following SQL query on a table $G$(from,to):

\[
\text{select distinct G3.to as x}
\text{from G G1, G G2, G G3}
\text{where G1.from=0 and G1.to=G2.from}
\text{and G2.to=8 and G3.from=G2.from}
\]

In the present work we also introduce association rules over tree queries. By mining for tree-query associations we can discover quite subtle properties of the data graph. Figure 2(a) shows a very simple example of an association that our algorithm might find in a social network: a data graph of persons where there is an edge $x \rightarrow y$ if $x$ considers $y$ to be a close friend. The tree query on the left matches all pairs $(x_1, x_2)$ of “co-friends”: persons that are friends of a common person (represented by an existential variable). The query on the right matches all co-friends $x_1$ of person #5 (represented by a parameterized node), and pairs all those co-friends to person #5. Now were the association from the left to the right to be discovered with a confidence of $c$, with $0 \leq c \leq 1$, then this would mean that the pairs retrieved by the right query actually constitute a fraction
\[
\begin{array}{c}
(x_1, x_2) \\
\exists \\
\downarrow x_1 \\
\Rightarrow \\
\exists \quad x_2 \\
\downarrow 5
\end{array}
\quad \quad \quad \quad \quad \quad
\begin{array}{c}
(x_1, 5) \\
\exists \\
\downarrow x_1 \\
\Rightarrow \\
\exists \quad 5
\end{array}
\quad \quad \quad \quad \quad \quad
\begin{array}{c}
(x) \\
\exists \\
\downarrow x \\
\Rightarrow \\
\exists \quad 5
\end{array}
\end{array}
\]

(a) (b) (c)

Figure 2: Simple examples of association rules over tree queries.

of \( c \) of all pairs retrieved by the left query, which indicates (for nonnegligible \( c \)) that 5 plays a special role in the network.

Figure 2(b) shows quite a different, but again simple, example of a tree-query association that our algorithm might discover in a food web. With confidence \( c \), this association means that of all organisms that are not on top of the food chain (i.e., they are fed upon by some other organism), a fraction of \( c \) is actually at least two down in the food chain.

The examples of tree queries and associations we just saw are didactical examples, but in Section 7 we will see more complicated examples of tree queries and associations mined in real-life datasets.

In this paper we present algorithms for mining tree queries and associations rules over tree queries in a large data graph. Some important features of these algorithms are the following:

1. Our algorithms belong to the group of graph mining algorithms where the input is a single large graph, and the task is to discover patterns that occur sufficiently often in the single data graph. We will refer to this group of algorithms as the single graph category. There is also a second category of graph mining algorithms, called the transactional category, which is explained in Section 2.

2. We restrict to patterns that are trees, such as the example in Figure 1. Tree patterns have formed an important special case in the transactional category (Section 2), but have not yet received special attention in the single-graph literature. Note that the data graph that is being mined is not restricted in any way.

3. The tree-query-mining algorithm is incremental in the number of nodes of the pattern. So, our algorithm systematically considers ever larger trees, and can be stopped any time it has run long enough or has produced enough results. Our algorithm does not need any space beyond what is needed to store the mining results. Thanks to the restriction to tree shapes the duplicate-free generation of trees can be done efficiently.

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Note that this does not just mean that 5 has many co-friends; if we only wanted to express that, just a frequent pattern in the form of the right query would suffice. For instance, imagine a data graph consisting of \( n \) disjoint 2-cliques (pairs of persons who have each other as a friend), where additionally all these persons also consider 5 to be an extra friend (but not vice versa). In such a data graph, 5 is a co-friend of everybody, and the association has a rather high confidence of more than \( 2/7 \). If, however, we would now add to the data graph a separate \( n \)-clique, then still \( 2/3 \) of all persons are a co-friend of 5, which is still a lot, but the confidence drops to below \( 2/n \).
4. For each tree, all conjunctive queries based on that tree are generated in the tree-query-mining algorithm. Here, we work in a levelwise fashion in the sense of Mannila and Toivonen [31].

5. As in classical association rules over itemsets [3], our association rule generation phase comes after the generation of frequent patterns and does not require access to the original dataset.

6. We apply the theory of conjunctive database queries [9, 40, 2] to formally define and to correctly generate association rules over tree queries. The conjunctive-query approach to pattern matching allows for an efficiently checkable notion of frequency, whereas in the subgraph-based approach, determining whether a pattern is frequent is NP-complete (in that approach the frequency of a pattern is the maximal number of disjoint subgraphs isomorphic to the pattern [29]).

7. There is a notion of equivalence among tree queries and association rules over tree queries. We carefully and efficiently avoid the generation of equivalent tree queries and associations, by using and adapting what is known from the theory of conjunctive database queries. Due to the restriction to tree shapes, equivalence and redundancy (which are normally NP-complete) are efficiently checkable.

8. Last but not least, our algorithms naturally suggest a database-oriented implementation in SQL. This is useful for several reasons. First, the number of discovered patterns can be quite large, and it is important to keep them available in a persistent and structured manner, so that they can be browsed easily, and so that association rules can be derived efficiently. Moreover, we will show how the use of SQL allows us to generate and check large numbers of similar patterns in parallel, taking advantage of the query processing optimizations provided by modern relational database systems. Third, a database-oriented implementation does not require us to move the dataset out of the database before it can be mined. In classical itemset mining, database-oriented implementations have received serious attention [39, 36], but less so in graph mining, a recent exception being an implementation in SQL of the seminal SUBDUE algorithm [8].

The purpose of this paper is to introduce tree queries and tree-query associations and to present algorithms for mining tree queries and tree-query associations. Concrete applications to discover new knowledge about scientific datasets are the topic of current research. Yet, the algorithms are fully implemented and we can already show that our approach works in practice, by showing some concrete results mined from a food web, a protein interactions graph, and a citation graph. We will also give performance results on random data graphs (as a worst-case scenario).

2 Related Work

Approaches to graph mining, especially mining for frequent patterns or association rules, can be divided in two major categories which are not to be confused.
1. In transactional graph mining, e.g., [12, 21, 22, 23, 28, 41, 42], the dataset consists of many small data graphs which we call transactions, and the task is to discover patterns that occur at least once in a sufficient number of transactions. (Approaches from machine learning or inductive logic programming usually call the small data graphs “examples” instead of transactions.)

2. In single-graph mining the dataset is a single large data graph, and the task is to discover patterns that occur sufficiently often in the dataset.

Note that single-graph mining is more difficult than transactional mining, in the sense that transactional graph mining can be simulated by single-graph mining, but the converse is not obvious.

Since our approach falls squarely within the single-graph category, we will focus on that category in this section. Most work in this category has been done on frequent pattern mining, and less attention has been spend on association rules. We briefly review the work in this category next:

- Cook and Holder [11] apply in their SUBDUE system the minimum description length (MDL) principle to discover substructures in a labeled data graph. The MDL principle states that the best pattern is that pattern that minimizes the description length of the complete data graph. Hence, in SUBDUE a pattern is evaluated on how well it can compress the entire dataset. The input for the SUBDUE system is a labeled data graph; nodes and edges are labeled with non-unique labels. This is in contrast with the unique labels (‘constants’) in our system. But as we already noted, non-unique node labels and edge-labels can easily be simulated by constants, but the converse is not obvious. The SUBDUE system only mines patterns, no association rules.

- Ghazizadeh and Chawathe [16] mine in their SEuS system for connected subgraphs in a labeled, directed data graph, as in the SUBDUE system. Instead of generating candidate patterns using the input data graph, SEuS uses a summary of the data graph. This summary gives an upper bound for the support of the patterns, and the user can then select those patterns of which he wants to know the exact support. SEuS also only mines for frequent patterns and not for associations.

- Vanetik, Gudes, and Shimony [19] propose an Apriori-like [3] algorithm for mining subgraphs from a labeled data graph. The support of a graph pattern is defined as the maximal number of edge-disjoint instances of the pattern in the data graph. By reducing the support counting problem to the maximal independent set problem on graphs, they show that in worst case, computing the support of a graph pattern is NP-hard. They propose an Apriori-like algorithm to minimize the number of patterns for which the support needs to be computed. The major idea of their approach is using edge-disjoint paths as building blocks instead of items in classical itemset mining. Vanetik, Gudes, and Shimony also only mine for frequent patterns in the data graph.

- Kuramochi en Karypis [29] use the same support measure for graph patterns as Vanetik, Gudes and Shimony [19]. They also note that computing
the support of a graph pattern is NP-hard in worst case, since it can be reduced to finding the maximum independent set (MIS) in a graph. Karamochi and Karypis quickly compute the support of a graph pattern using approximate MIS-algorithms. The number of candidate patterns is restricted using canonical labeling. As the majority of algorithms, Karamochi and Karypis only mine for frequent patterns.

- Jeh and Widom [24] consider patterns that are, like our tree queries, inspired by conjunctive database queries, and they also emphasize the tree-shaped case. A severe restriction, however, is that their patterns can be matched by single nodes only, rather than by tuples of nodes. Still their work is interesting in that it presents a rather nonstandard approach to graph mining, quite different from the standard incremental, levelwise approach, and in that it incorporates ranking. Jeh and Widom mention association rules as an example of an application of their mining framework.

The related work that was most influential for us is Warmr [12, 13], although it belongs to the transactional category. Based on inductive logic programming, patterns in Warmr also feature existential variables and parameters. While not restricted to tree shapes, the queries in Warmr are restricted in another sense so that only transactional mining can be supported. Association rules in Warmr are defined in a naive manner through pattern extension, rather than being founded upon the theory of conjunctive query containment. The Warmr system is also Prolog-oriented, rather than database-oriented, which we believe is fundamental to mining of single large data graphs, and which allows a more uniform and parallel treatment of parameter instantiations, as we will show in this paper. Finally, Warmr does not seriously attempt to avoid the generation of duplicates. Yet, Warmr remains a pathbreaking work, which did not receive sufficient follow-up in the data mining community at large. We hope our present work represents an improvement in this respect. Many of the improvements we make to Warmr were already envisaged (but without concrete algorithms) in 2002 by Goethals and the second author [13].

Finally, we note that parameterized conjunctive database queries have been used in data mining quite early, e.g., [39, 38], but then in the setting of “data mining query languages”, where a single such query serves to specify a family of patterns to be mined or queried for, rather than the mining for such queries themselves, let alone associations among them.

3 Problem Statement

In this section we define some concepts formally. In the appendix an overview of all notations used in this paper is given.

We basically assume a set $U$ of data constants from which the nodes of the data graph to be mined will be taken.

3.1 Graph-theoretic concepts

Let $N \subseteq U$ be any finite set of nodes; nodes can be any data objects such as numbers or strings. For our purposes, we define a (directed) graph on $N$ as
3.2 Tree Pattern

Tree Patterns A parameterized tree pattern $P$ is a tree whose nodes are called variables, and where additionally:

- Some variables may be marked as being existential;
- Some other variables may be marked as parameters;
- The variables of $P$ that are neither existential nor parameters are called distinguished.

We will denote the set of existential variables by $\Pi$, the set of parameters by $\Sigma$, and the set of distinguished variables by $\Delta$. To make clear that these sets belong to some parameterized tree pattern $P$ we will use a subscript as in $\Pi_P$ or $\Sigma_P$.

A parameter assignment $\alpha$, for a parameterized tree pattern $P$, is a mapping $\Sigma \rightarrow U$ which assigns data constants to the parameters.

An instantiated tree pattern is a pair $(P, \alpha)$, with $P$ a parameterized tree pattern and $\alpha$ a parameter assignment for $P$. We will also denote this by $P^\alpha$.

When depicting parameterized tree patterns, existential nodes are indicated by labeling them with the symbol ‘$\exists$’ and parameters are indicated by labeling them with the symbol ‘$\sigma$’. When depicting instantiated tree patterns, parameters are indicated by directly writing down their parameter assignment.

Figure 3 shows an illustration.

Matching Recall that a homomorphism from a graph $G_1$ to a graph $G_2$ is a mapping $\mu$ from the nodes of $G_1$ to the nodes of $G_2$ that preserves edges, i.e., if $(i, j) \in G_1$ then $(\mu(i), \mu(j)) \in G_2$. We now define a matching of an instantiated tree pattern $P^\alpha$ in a data graph $G$ as a homomorphism $\mu$ from the underlying tree of $P$ to $G$, with the constraint that for any parameter $\sigma$, if $\alpha(\sigma) = a$, then $\mu(\exists) = a$. 

Figure 3: (a) is a parameterized tree pattern, and (b) is an instantiation of (a).
\( \mu(\sigma) \) must be the node \( a \). We denote the set \( \{ \mu | \Delta : \mu \text{ is a matching of } P^{\alpha} \text{ in } G \} \) by \( P^{\alpha}(G) \).

**Frequency of a tree pattern** The frequency of an instantiated tree pattern \( P^{\alpha} \) in a data graph \( G \), is formally defined as the cardinality of \( P^{\alpha}(G) \). So, we count the number of matchings of \( P^{\alpha} \) in \( G \), with the important provision that we identify any two matchings that agree on the distinguished variables. Indeed, two matchings that differ only on the existential nodes need not be distinguished, as this is precisely the intended semantics of existential nodes. Note that we do not need to worry about selected nodes, as all matchings will agree on those by definition. For a given threshold \( k \) (a natural number) we say that \( P^{\alpha} \) is \( k \)-frequent if its frequency is at least \( k \). Often the threshold is understood implicitly, and then we talk simply about “frequent” patterns and denote the threshold by \( \minsup \).

**Example.** Take again the instantiated tree pattern \( P^{\alpha} \) shown in Figure 3(b). Let us name the existential node by \( y \); let us name the parameter labeled 0 by \( z_1 \); the parameter labeled 8 by \( z_2 \); and the parameter labeled 6 by \( z_3 \). The distinguished node already has the name \( x \). Now let us apply \( P^{\alpha} \) to the simple example data graph \( G \) shown in Figure 4(a). The following table lists all matchings of \( P^{\alpha} \) in \( G \):

| \( h_1 \) | 0 | 1 | 4 | 8 | 6 |
| \( h_2 \) | 0 | 2 | 4 | 8 | 6 |
| \( h_3 \) | 0 | 2 | 5 | 8 | 6 |

As required by the definition, all matchings match \( z_1 \) to 0, \( z_2 \) to 8, and \( z_3 \) to 6. Although there are three matchings, when determining the frequency of \( P^{\alpha} \) in \( G \), we only look at their value on \( x \) to distinguish them, as \( y \) is existential. So, \( h_2 \) and \( h_3 \) are identified as identical matchings when counting the number of matchings. In conclusion, the frequency of \( P^{\alpha} \) in \( G \) is two, as \( x \) can be matched to the two different nodes 1 and 2.

### 3.3 Tree Query

**Tree Queries** A parameterized tree query \( Q \) is a pair \((H, P)\) where:

1. \( P \) is a parameterized tree pattern, called the **body** of \( Q \);
2. \( H \) is a tuple of distinguished variables and parameters coming from \( P \). All distinguished variables of \( P \) must appear at least once in \( H \). We call \( H \) the head of \( Q \).

A parameter assignment for \( Q \) is simply a parameter assignment for its body, and an instantiated tree query is then again a pair \((Q, \alpha)\) with \( Q \) a parameterized tree query and \( \alpha \) a parameter assignment for \( Q \). We will again also denote this by \( Q^\alpha \).

When depicting tree queries, the head is given above a horizontal line, and the body below it. Two illustrations are given in Figure 5.

**Frequency of a tree query** The frequency of an instantiated tree query \( Q^\alpha = ((H,P), \alpha) \) in a data graph \( G \), is defined as the frequency of the body \( P^\alpha \) in \( G \). When \( G \) is understood, we denote the frequency by \( \text{Freq}(P^\alpha) \). For a given threshold \( k \) (a natural number) we say that \( Q^\alpha \) is \( k \)-frequent if its frequency is at least \( k \). Again, this threshold is often understood implicitly, and then we talk simply about “frequent” queries and denote the threshold by \( \text{minsup} \).

**Containment of tree queries** An important step towards our formal definition of tree-query association is the notion of containment among queries. Since queries are parameterized, a variation of the classical notion of containment [9, 40, 2] is needed in that we now need to specify a parameter correspondence.

First, we define the answer set of an instantiated tree query \( Q^\alpha \), with \( Q = (H,P) \), in a data graph \( G \) as follows:

\[
Q^\alpha(G) := \{ \mu(H) \mid \mu \text{ is a matching of } P^\alpha \text{ in } G \}
\]

Consider two parameterized tree queries \( Q_1 \) and \( Q_2 \), with \( Q_i = (H_i, P_i) \) for \( i = 1, 2 \). A parameter correspondence from \( Q_1 \) to \( Q_2 \) is any mapping \( \rho : \Sigma_1 \to \Sigma_2 \). We then say that a parameterized tree query \( Q_2 \) is \( \rho \)-contained in a parameterized tree query \( Q_1 \), if for every \( \alpha_2 \), a parameter assignment for \( Q_2 \), \( Q_2^{\alpha_2}(G) \subseteq Q_1^{\rho \circ \alpha_2}(G) \) for all data graphs \( G \). In shorthand notation we write this as \( Q_2 \preceq_\rho Q_1 \).

Containment as just defined is a semantical property, referring to all possible data graphs, and it is not immediately clear how one could decide this property syntactically. The required syntactical notion for this is that of \( \rho \)-containment.

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**Figure 5:**

- (a) and (b) are parameterized tree queries; (c) is an instantiation of (a); (d) is an instantiation of (b); and query (b) is \( \rho \)-contained in query (a).

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\[\begin{array}{cccc}
(x_1, x_2, x_3) & (x, x, \sigma_3) & (x_1, x_2, x_3) & (x, x, 6) \\
\sigma_1 & \sigma_1 & 0 & 0 \\
x_1 & x & x_1 & x \\
\exists_1 & \exists & \exists_1 & \exists \\
\sigma_2 & \sigma_2 & \exists & 8 \\
x_2 & \exists & \exists_2 & x_3 \\
\sigma_3 & \exists & \sigma_3 & 6 \\
x_3 & & & \\
(a) & (b) & (c) & (d)
\end{array}\]
Hence, we can conclude that \( f \) contains mapping, which we next define in two steps. For the tree queries \( Q_1 \) and \( Q_2 \) as above, and \( \rho \) a parameter correspondence from \( Q_1 \) to \( Q_2 \):

1. A \( \rho \)-containment mapping from \( P_1 \) to \( P_2 \) is a homomorphism \( f \) from the underlying tree of \( P_1 \) to the underlying tree of \( P_2 \), with the properties:
   
   (a) \( f \) maps the distinguished nodes of \( P_1 \) to distinguished nodes or parameters of \( P_2 \); and
   
   (b) \( f|_{\Sigma_1} = \rho \), i.e., for each \( z \in \Sigma_1 \) we have \( f(z) = \rho(z) \).

2. Finally, a \( \rho \)-containment mapping from \( Q_1 \) to \( Q_2 \) is a \( \rho \)-containment mapping \( f \) from \( P_1 \) to \( P_2 \) such that \( f(H_1) = H_2 \).

For later use, we note:

**Lemma 1.** Consider three parameterized tree patterns \( P_1 \), \( P_2 \), and \( P_3 \), a parameter correspondence \( \rho_1 : \Sigma_1 \to \Sigma_2 \), a parameter correspondence \( \rho_2 : \Sigma_2 \to \Sigma_3 \), a \( \rho_1 \)-containment mapping \( f_1 \) from \( P_1 \) to \( P_2 \), and a \( \rho_2 \)-containment mapping \( f_2 \) from \( P_2 \) to \( P_3 \). Then \( f_2 \circ f_1 \) is a \( (\rho_2 \circ \rho_1) \)-containment mapping from \( P_1 \) to \( P_3 \).

**Proof.** We will show that:

1. \( f_2 \circ f_1 \) is homomorphism;

2. \( f_2 \circ f_1 \) maps distinguished nodes of \( P_1 \) to distinguished nodes or parameters of \( P_3 \); and

3. \( (f_2 \circ f_1)|_{\Sigma_1} = \rho_2 \circ \rho_1 \).

   (1) Clearly \( f_2 \circ f_1 \) is a homomorphism since both \( f_1 \) and \( f_2 \) are homomorphisms, and it is already known that a composition of homomorphisms is a homomorphism.

   (2) Consider a \( x_1 \in \Delta_1 \), then there are two possibilities for \( f_1(x_1) \):

   1. \( f_1(x_1) = x_2 \), with \( x_2 \in \Delta_2 \). Then we know, since \( f_2 \) is a \( \rho_2 \)-containment mapping, that \( f_2(x_2) \) is either a distinguished node \( x_3 \in \Delta_3 \), or a parameter \( z_3 \in \Sigma_3 \).

   2. \( f_1(x_1) = z_2 \), with \( z_2 \in \Sigma_2 \). Then we know, since \( f_2|_{\Sigma_2} = \rho_2 \), that \( f_2(z_2) = z_3 \), with \( z_3 \in \Sigma_3 \).

Hence, we can conclude that \( f_2 \circ f_1 \) maps distinguished nodes of \( P_1 \) to distinguished nodes or parameters of \( P_3 \).

   (3) For each \( z_1 \in \Sigma_1 \), we have \( f_2(f_1(z_1)) = \rho_2(\rho_1(z_1)) \). Hence, \( (f_2 \circ f_1)|_{\Sigma_1} = \rho_2 \circ \rho_1 \).

From the theory of conjunctive database queries \[ \text{[9]} \] \[ \text{[10]} \] \[ \text{[2]} \] we can derive the following:

**Lemma 2.** Consider two parameterized tree queries \( Q_1 \) and \( Q_2 \), with \( Q_1 = (H_1, P_1) \) and \( Q_2 = (H_2, P_2) \) and a parameter correspondence \( \rho : \Sigma_1 \to \Sigma_2 \). Then \( Q_2 \) is \( \rho \)-contained in \( Q_1 \) \( Q_2 \subseteq_\rho Q_1 \), if and only if there exists a \( \rho \)-containment mapping from \( Q_1 \) to \( Q_2 \).
Proof. Let us start with the ‘only if’ direction. We first introduce the concept of a freezing of a parameterized tree query \( Q = (H, P) \). Recall that \( U \) is the set of data constants from which the nodes of the data graph to be mined will be taken. A freezing \( \beta \) of \( P \) is then a one-to-one mapping from the nodes of \( P \) to \( U \). We denote by freeze\(_\beta\)(\( P \)) the data graph constructed from \( P \) by replacing each node \( n \) of \( P \) by \( \beta(n) \), and we denote by freeze\(_\beta\)(\( H \)) the tuple constructed from \( H \) by replacing each node \( n \) in \( H \) by the data constant \( \beta(n) \).

For example, consider the parameterized tree query \( Q = (H, P) \) in Figure 6(a). Figure 6(b) shows freeze\(_\beta\)(\( P \)) and freeze\(_\beta\)(\( H \)) for the freezing \( \beta \) given as follows: \( x_1 \rightarrow c_1; x_2 \rightarrow c_2; z_1 \rightarrow c_3; x_4 \rightarrow c_4; x_5 \rightarrow c_5; \sigma_6 \rightarrow c_6 \).

We can now continue with the proof of the ‘only if’ direction. Consider a freezing \( \beta \) from the nodes of \( P_2 \) to \( U \). Note that \( \beta|_{\Sigma_2} \) is a parameter assignment for \( Q_2 \), and freeze\(_\beta\)(\( H_2 \)) \( \in Q_2^{\beta|_{\Sigma_2}=\emptyset}(\)freeze\(_\beta\)(\( P_2 \))\). Since \( Q_2 \subseteq \rho \), also freeze\(_\beta\)(\( H_2 \)) \( \in Q_1^{\beta|_{\Sigma_2}=\emptyset}(\)freeze\(_\beta\)(\( P_2 \))\). Hence, there must be a matching \( \mu \) from \( P_1^{\beta|_{\Sigma_2}=\emptyset} \) in freeze\(_\beta\)(\( P_2 \)) such that \( \mu(H_1) = \) freeze\(_\beta\)(\( H_2 \)). Now consider the function \( g : \beta^{-1} \circ \mu \). We show that \( g \) is \( \rho \)-containment mapping from \( Q_1 \) to \( Q_2 \):

1. Clearly, \( g \) is a homomorphism from \( P_1 \) to \( P_2 \) since \( \mu \) is a homomorphism and \( \beta^{-1} \) is an isomorphism. Also the following properties hold for \( g \):
   \[
   (a) \quad g \text{ maps distinguished nodes of } P_1 \text{ to distinguished nodes or parameters of } P_2 \text{ since } g(H_1) = H_2 \text{ (as shown in (2)); and}
   \[
   (b) \quad \text{ for each } z \in \Sigma_1: \quad g(z) = \beta^{-1}(\mu(z)) = \beta^{-1}(\beta(\rho(z))) = \rho(z), \text{ hence } g|_{\Sigma_1} = \rho
   \]

2. \( g(H_1) = \beta^{-1}(\mu(H_1)) = \beta^{-1}(\text{freeze\(_\beta\)(\( H_2 \))) = H_2.\)

Hence, we conclude that \( g \) is a \( \rho \)-containment mapping from \( Q_1 \) to \( Q_2 \).

Let us then look at the ‘if’ direction. Let \( h \) be the \( \rho \)-containment mapping from \( Q_1 \) to \( Q_2 \). Consider an arbitrary parameter assignment \( \alpha_2 \) for \( Q_2 \). We must prove that for every data graph \( G \), if \( a \in Q_2^{\alpha_2}(G) \), then also \( a \in Q_1^{\beta|_{\Sigma_2}=\emptyset}(G) \). Consider such an arbitrary data graph \( G \). Since, \( a \in Q_2^{\alpha_2}(G) \), we know that there exists a matching \( \mu \) of \( P_2^{\alpha_2} \) in \( G \) such that \( a = \mu(H_2) \). Now consider the function \( g = \mu \circ h \). We show that \( g \) is a matching from \( P_1^{\beta|_{\Sigma_2}=\emptyset} \) in \( G \) and \( a = g(H_1) \):

1. \( g \) is a homomorphism since both \( \mu \) and \( g \) are homomorphisms; and
2. for each \( z \in \Sigma_1 \) we have \( g(z) = \mu(h(z)) = \mu(\rho(z)) = \alpha_2(\rho(z)) \).

So, \( g \) is indeed a matching of \( P_1^{\beta|_{\Sigma_2}=\emptyset} \) in \( G \). Finally, we observe that \( g(H_1) = \mu(h(H_1)) = \mu(H_2) = a \), as desired.

Checking for a containment mapping is evidently computable, and although the problem for general database conjunctive queries is NP-complete, our restriction to tree shapes allows for efficient checking, as we will see later.

Example. Consider the parameterized and instantiated tree queries shown in Figure 4. In the example data graph in Figure 4(a) the frequency of query (c) is 10 and that of query (d) is 2. Let \( \Sigma_a \) be the set of parameters of query (a), and let \( \Sigma_b \) be the set of parameters of query (b); then let the parameter correspondence \( \rho : \Sigma_a \rightarrow \Sigma_b \) be as follows: \( \sigma_1 \rightarrow \sigma_1; \sigma_2 \rightarrow \sigma_2 \). A moment’s
reflection should convince the reader that (b) is ρ-contained in (a), and indeed a ρ-containment mapping \( f \) from (a) to (b) can be found as follows:

| \( f \) | \( \sigma_1 \) | \( \sigma_1 \) |
|-----|-----|-----|
| \( x_1 \) | \( x \) |
| \( x_2 \) | \( x \) |
| \( \exists_1 \) | \( \exists \) |
| \( \exists_2 \) | \( \exists \) |
| \( \sigma_2 \) | \( \sigma_2 \) |
| \( x_3 \) | \( \sigma_3 \) |

### 3.4 Tree-Query Association

**Association Rules** A parameterized association rule (pAR) is of the form \( Q_1 \Rightarrow \rho Q_2 \), with \( Q_1 \) and \( Q_2 \) parameterized tree queries and \( \rho \) a parameter correspondence from \( \Sigma_1 \) to \( \Sigma_2 \). We call a pAR *legal* if \( Q_2 \subseteq \rho Q_1 \). We call \( Q_1 \) the left-hand side (lhs), and \( Q_2 \) the right-hand side (rhs). A parameter assignment \( \alpha \), for a pAR, is a mapping \( \Sigma_2 \rightarrow U \) which assigns data constants to the parameters. An instantiated association rule (iAR) is a pair \( (Q_1 \Rightarrow \rho Q_2, \alpha) \), with \( Q_1 \Rightarrow \rho Q_2 \) a pAR and \( \alpha \) a parameter assignment for \( Q_1 \Rightarrow \rho Q_2 \). Note that while \( \alpha \) is only defined on the rhs, we can also apply it to the lhs by using \( \rho \) first.

**Confidence** The confidence of an iAR in a data graph \( G \) is defined as the frequency of \( Q_2^{\alpha_2} \) divided by the frequency of \( Q_1^{\alpha_2 \rho} \). If the AR is legal, we know that the answer set of \( Q_2^{\alpha_2} \) is a subset of the answer set of \( Q_1^{\alpha_2 \rho} \), and hence the confidence equals precisely the proportion that the \( Q_2^{\alpha_2} \) answer set takes up in the \( Q_1^{\alpha_2 \rho} \) answer set. Thus, our notions of a legal pAR and confidence are very intuitive and natural.

For a given threshold \( c \) (a rational number, \( 0 \leq c \leq 1 \)) we say that the iAR is *c-confident* in \( G \) if its confidence in \( G \) is at least \( c \). Often the threshold is understood implicitly, and then we talk simply about “confident” iARs and denote the threshold by \( \text{minconf} \).

Furthermore, the iAR is called *frequent* in \( G \) if \( Q_2^{\alpha_2} \) is frequent in \( G \). Note that if the iAR is legal and frequent, then also \( Q_1^{\alpha_2 \rho} \) is frequent, since the rhs is ρ-contained in the lhs.
Example. Continuing the previous example, we can see that we can form a legal pAR from the queries of Figure 5 with (a) the lhs and (b) the rhs and $\rho$ as follows: $\sigma_1 \rightarrow \sigma_1; \sigma_2 \rightarrow \sigma_2$. We can also form an iAR with the tree queries in Figure 5(c) and Figure 5(d); the confidence of this iAR in the data graph of Figure 4(a) is $2/10$. Many more examples of ARs are given in Section 5.

3.5 Mining Problems

We are now finally ready to define the graph mining problems we want to solve.

3.5.1 Mining Tree Queries

**Input:** A data graph $G$; a threshold $minsup$.

**Output:** All frequent instantiated tree queries $Q = ((H, P), \alpha)$.

In theory, however, there are infinitely many $k$-frequent tree queries, and even if we set an upper bound on the size of the patterns, there may be exponentially many. As an extreme example, if $G$ is the complete graph on the set of nodes $\{1, \ldots, n\}$, and $k \leq n$, then any instantiated pattern with all parameters assigned to values in $\{1, \ldots, n\}$, and with at least one distinguished variable, is frequent.

Hence, in practice, we want an algorithm that runs incrementally, and that can be stopped any time it has run long enough or has produced enough results. We introduce such an algorithm in Section 4.

3.5.2 Association Rule Mining

**Input:** A data graph $G$; a threshold $minsup$; a parameterized tree query $Q_{\text{left}}$; and a threshold $minconf$.

**Output:** All iARs ($Q_{\text{left}} \Rightarrow \rho Q_{\text{right}}, \alpha$) that are legal, frequent and confident in $G$.

In theory, however, there are infinitely many legal, frequent and confident association rules for a fixed lhs, and even if we set an upper bound on the size of the rhs, there may be exponentially many. Hence, in practice, we want an algorithm that runs incrementally, and that can be stopped any time it has run long enough or has produced enough results. We introduce such an algorithm in Section 5.

4 Mining Tree Queries

In this Section we present an algorithm for mining frequent instantiated tree queries in a large data graph. But first we show that we do not need to tackle the problem in its full generality.

4.1 Problem Reduction

In this subsection we show that, without loss of generality, we can focus on parameterized tree queries that are ‘pure’.
Figure 7: (a) is an impure parameterized tree query. The parameterized tree query in (b) is the purification of the parameterized tree query (a), and expresses precisely the same information.

Figure 8: (b) is the pure instantiated tree query constructed from the instantiated tree pattern in (a)

**Pure Tree Queries** To define this formally, assume that all possible variables (nodes of tree patterns) have been arranged in some fixed but arbitrary order. We then call a parameterized tree query \( Q = (H, P) \) pure when \( H \) consists of the enumeration, in order and without repetitions, of all the distinguished variables of \( P \). In particular \( H \) cannot contain parameters. We call \( H \) the *pure head* for \( P \). As an illustration, the parameterized tree query in Figure 5(a) is pure, while the parameterized tree query in Figure 5(b) is not pure.

A parameterized tree query that is not pure can always be rewritten to a parameterized tree query that is pure, in such a way that all instantiations of the impure query correspond to instantiations of the pure query, with the same frequency. Indeed, take a parameterized tree query \( Q = (H, P) \). We can purify \( Q \) by removing all parameters and repetitions of distinguished variables from \( H \), and sort \( H \) by the order on the variables. An illustration of this is given in Figure 7.

We can conclude that it is sufficient to only consider pure instantiated tree queries. As a consequence, rather than mining tree queries, it suffices to mine for tree patterns, because the frequency of a query is nothing else than the frequency of his body, i.e., a pattern. An illustration is given in Figure 8.
4.2 Overall Approach

An overall outline of our tree-query mining algorithm is the following:

**Outer loop:** Generate, incrementally, all possible trees $T$ of increasing sizes. Avoid trees that are isomorphic to previously generated ones.

**Inner loop:** For each $T$, generate all instantiated tree patterns $P^*$ based on $T$, and test their frequency.

The algorithm is incremental in the number of nodes of the pattern. We generate canonically ordered rooted trees of increasing sizes, avoiding the generation of isomorphic duplicates. It is well known how to do this efficiently [37, 30, 42, 10]. Note that this generation of trees is in no way “levelwise” [31]. Indeed, under the way we count pattern occurrences, a subgraph of a pattern might be less frequent than the pattern itself (this was already pointed out by Kuramochi and Karypis [29]). So, our algorithm systematically considers ever larger trees, and can be stopped any time it has run long enough or has produced enough results. Our algorithm does not need any space beyond what is needed to store the mining results. The outer loop of our algorithm will be explained in more detail in Section 4.3.

For each tree, all conjunctive queries based on that tree are generated. Here, we do work in a levelwise fashion. This aspect of our algorithm has clear similarities with “query flocks” [39]. A query flock is a user-specified conjunctive query, in which some constants are left unspecified and viewed as parameters. A levelwise algorithm was proposed for mining all instantiations of the parameters under which the resulting query returns enough answers. We push that approach further by also mining the query flocks themselves. Consequently, the specialization relation on queries used to guide the levelwise search is quite different in our approach. The inner loop of our algorithm will be explained in more detail in Section 4.4.

A query based on some tree may be equivalent to a query based on a previously seen tree. Furthermore, two queries based on the same tree may be equivalent. We carefully and efficiently avoid the counting of equivalent queries, by using and adapting what is known from the theory of conjunctive database queries. This will be discussed in Section 4.5.

4.3 Outer Loop

In the outer loop we generate all possible trees of increasing sizes and we avoid trees that are isomorphic to previously generated ones. In fact, it is well known how to do this [37, 30, 42, 10]. What these procedures typically do is generating trees that are canonically ordered in the following sense. Given an (unordered) tree $T$, we can order the children of every node in some way, and call this an ordering of $T$. For example, Figure 4 shows two orderings of the same tree. From the different orderings of a tree $T$, we want to uniquely select one, to be the canonical ordering of $T$. For each such possible ordering of $T$, we can write down the level sequence of the resulting tree. This is actually a string representation of the resulting tree. This level sequence is as follows: if the tree has $n$ nodes then this is a sequence of $n$ numbers, where the $i$th number is the depth of the $i$th node in preorder. Here, the depth of the root is 0, the depth of
its children is 1, and so on. The canonical ordering of $T$ is then the ordering of $T$ that yields the lexicographically maximal level sequence among all possible orderings of $T$.

For example, in Figure 9, the left one is the canonical one.

4.4 Inner Loop

Let $G$ be the data graph being mined, and let $U$ be its set of nodes. In this section, we fix a tree $T$, and we want to find all instantiated tree patterns $P^\alpha$ based on $T$ whose frequency in $G$ is at least $\text{minsup}$.

This task lends itself naturally to a levelwise approach [31]. A natural choice for the specialization relation is suggested by an alternative notation for the patterns under consideration. Concretely, since the underlying tree $T$ is fixed, any parameterized tree pattern $P$ based on $T$ is characterized by two parameters:

1. The set $\Pi$ of existential nodes;
2. The set $\Sigma$ of parameters.

Note that $\Pi$ and $\Sigma$ are disjoint.

Thus, a parameterized tree pattern $P$ is completely characterized by the pair $(\Pi, \Sigma)$. An instantiation $P^\alpha$ of $P$ is then represented by the triple $(\Pi, \Sigma, \alpha)$. For two parameterized tree patterns $P_1 = (\Pi_1, \Sigma_1)$ and $P_2 = (\Pi_2, \Sigma_2)$ we now say that $P_1$ specializes $P_2$ if $\Pi_1 \supseteq \Pi_2$ and $\Sigma_1 \supseteq \Sigma_2$; and $\alpha_2 = \alpha_1|_{\Sigma_2}$. We also say that $P_2$ generalizes $P_1$.

Parent An immediate generalization of a tree pattern is called a parent. Formally, let $P = (\Pi, \Sigma)$ and $P' = (\Pi', \Sigma')$ be parameterized tree patterns based on $T$. We say that $P'$ is a parent of $P$ if:

(i) $\Sigma = \Sigma'$ and $\Pi = \Pi' \cup \{y\}$ for some node $y \notin \Pi'$; or

(ii) $\Pi = \Pi'$ and $\Sigma = \Sigma' \cup \{z\}$ for some node $z \notin \Sigma'$.

From the following lemma, it follows that specialized patterns have a lower frequency, as expected for a specialization relation:

Lemma 3. Let $P$ and $P'$ be parameterized tree patterns such that $P'$ is a parent of $P$. Let $P^\alpha$ be an instantiation of $P$, and let $\alpha' = \alpha|_{\Sigma'}$. Then $\text{Freq}(P^\alpha) \leq \text{Freq}(P'^{\alpha'})$.

Proof. We will show that $\#P^\alpha(G) \leq \#P'^{\alpha'}(G)$ by defining an injection $I : P^\alpha(G) \rightarrow P'^{\alpha'}(G)$. 

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Since \( P' \) is a parent of \( P \), we know that \( \Delta' = \Delta \cup \{ u \} \) where \( u \) is either an existential node or a parameter of \( P \). Note that each \( \mu \in P'^{\alpha}(G) \) is of the form \( \overline{\mu}_{|\Delta} \) for some matching \( \mu \) of \( P^{\alpha} \in G \). For each \( \mu \) in \( P^{\alpha}(G) \), we fix arbitrarily \( \overline{\mu} \).

Now we define \( I(\mu) := \overline{\mu}_{|\Delta} \). To see that \( I \) is an injection, let \( \mu_1, \mu_2 \in P'^{\alpha}(G) \) and suppose that \( I(\mu_1) = I(\mu_2) \). In other words, \( \overline{\mu}_1|_{\Delta'} = \overline{\mu}_2|_{\Delta'} \). In particular, \( \mu_1 = \overline{\mu}_1|_{\Delta} = \overline{\mu}_2|_{\Delta} = \mu_2 \), as desired.

Hence, we can conclude that \( \#P^{\alpha}(G) \leq \#P'^{\alpha'}(G) \) and that \( \text{Freq}(P^{\alpha}) \leq \text{Freq}(P'^{\alpha'}) \).

The above lemma suggests the following definition of specialization among instantiated tree patterns: we say that \((\Pi_1, \Sigma_1, \alpha_1)\) is a specialization of \((\Pi_2, \Sigma_2, \alpha_2)\) if the parameterized tree pattern \((\Pi_1, \Sigma_1)\) is a specialization of the parameterized tree pattern \((\Pi_2, \Sigma_2)\), and \( \alpha_2 = \alpha_1|_{\Sigma_2} \).

Intuitively, the previous lemma then expresses that the frequency of an instantiated tree pattern is always at most the frequency of any of its instantiated parents.

### 4.4.1 Candidate generation

#### Candidate pattern

A candidate pattern is an instantiated tree pattern whose frequency is not yet determined, but all whose generalizations are known to be frequent.

Using the specialization relation and the definition for a candidate pattern we explain how the levelwise search for frequent instantiated tree patterns will go.

#### Levelwise search

We start with the most general instantiated tree pattern \( P = (\emptyset, \emptyset, \emptyset) \), and we progressively consider more specific patterns. The search has the typical property that, in each new iteration, new candidate patterns are generated; the frequency of all newly discovered candidate patterns is determined, and the process repeats.

There are many different instantiations to consider for each parameterized tree pattern. Hence, to generate candidate patterns in an efficient manner, we propose the use of candidacy tables and frequency tables. These candidacy and frequency tables allow us to generate all frequent instantiations for a particular parameterized tree pattern in parallel. A frequency table contains all frequent instantiations for a particular parameterized tree pattern.

Formally, for any parameterized pattern \( P = (\Pi, \Sigma) \), we define:

- \( \text{CanTab}_{\Pi, \Sigma} = \{ \alpha \mid P^{\alpha} \text{ is a candidate instantiated tree pattern} \} \)
- \( \text{FreqTab}_{\Pi, \Sigma} = \{ \alpha \mid P^{\alpha} \text{ is a frequent instantiated tree pattern} \} \)

Technically, the table has columns for the different parameters, plus a column \( \text{freq} \). Note that when \( \Sigma = \emptyset \), i.e., \( P \) has no parameters, this is a single-column, single-row table containing just the frequency of \( P \). This still makes sense and can be interpreted as boolean values; for example, if \( \text{FreqTab}_{\Pi, \emptyset} \) contains the empty tuple, then the pattern \((\Pi, \emptyset, \emptyset)\) is frequent; if the table is empty, the pattern is not frequent. Of course in practice, all frequency tables for parameterless patterns can be combined into a single table. All frequency tables are kept in a relational database.

The following crucial lemma shows these tables can be populated efficiently.
Join Lemma. A parameter assignment \( \alpha \) is in \( \text{CanTab}_{\Pi,\Sigma} \) if and only if the following conditions are satisfied for every parent \((\Pi',\Sigma')\) of \((\Pi,\Sigma)\):

(i) If \( \Pi = \Pi' \), then \( \alpha|_{\Sigma'} \in \text{FreqTab}_{\Pi',\Sigma'} \);
(ii) If \( \Sigma = \Sigma' \), then \( \alpha \in \text{FreqTab}_{\Pi',\Sigma'} \).

Proof. For the ‘only-if’ direction: By definition of a candidacy table, if \( \alpha \in \text{CanTab}_{\Pi,\Sigma} \), then all generalizations of \((\Pi,\Sigma,\alpha)\) are frequent. In particular, for all parents \((\Pi',\Sigma')\) of \((\Pi,\Sigma)\), we know that \((\Pi',\Sigma',\alpha|_{\Sigma'})\) is frequent, since parents are generalizations.

For the ‘if’ direction, we must show that all generalizations of \((\Pi,\Sigma,\alpha)\) are frequent. Consider such a generalization \((\Pi_{g1},\Sigma_{g1},\alpha|_{\Sigma_{g1}})\). Let us denote the parent relation by \( \geq_p \). Then there is a sequence of parent patterns:

\[
(\Pi_{g1},\Sigma_{g1}) \geq_p (\Pi_{g2},\Sigma_{g2}) \geq_p ... \geq_p (\Pi',\Sigma').
\]

And we have: \( \text{Freq}(\Pi_{g1},\Sigma_{g1},\alpha|_{\Sigma_{g1}}) \geq \text{Freq}(\Pi_{g2},\Sigma_{g2},\alpha|_{\Sigma_{g2}}) \geq ... \geq \text{Freq}(\Pi',\Sigma',\alpha|_{\Sigma'}) \geq \minsup. \) The last inequality is given by (i) or (ii), the other inequalities are given by Lemma 3.

The Join Lemma has its name because, viewing the tables as relational database tables, it can be phrased as follows:

Each candidacy table can be computed by taking the natural join of its parent frequency tables.

The only exception is when \( \Pi = \emptyset \) and \( \Sigma = \{z\} \) is a singleton; this is the initial iteration of the search process, when there are no constants in the parent tables to start from. In that case, we define \( \text{CanTab}_{\emptyset,\{z\}} \) as the table with a single column \( z \), holding all nodes of the data graph \( G \) being mined.

4.4.2 Frequency counting using SQL

The search process starts by determining the frequency of the underlying tree \( T = (\emptyset,\emptyset) \); indeed, formally this amounts to computing \( \text{FreqTab}_{\emptyset,\emptyset} \). Similarly, for each parameterized tree pattern \( P = (\Pi,\emptyset) \) with \( \Pi \neq \emptyset \), all we can do is determine its frequency, except that here, we do this only on condition that its parent patterns are frequent.

We have seen above that, if the frequency tables are viewed as relational database tables, we can compute each candidacy table by a single database query, using the Join Lemma. Now suppose the data graph \( G \) that is being mined is stored in the relational database system as well, in the form of a table \( G(\text{from},\text{to}) \). Then also each frequency table can be computed by a single SQL query.

Indeed, in the cases where \( \Sigma = \emptyset \) this simply amounts to formulating the pattern in SQL, and determining its count (eliminating duplicates). Since our patterns are in fact conjunctive queries (or datalog rules) known from database research \[2, 40\]. They can easily be translated in SQL:

- The FROM-clause consists of all table references of the form \( G \) as \( Gij \), for all edges \( x_i \rightarrow x_j \) in \( T \).
- The WHERE-clause consists of all equalities of the form \( Gij.\text{from} = Gik.\text{from} \) as well of equalities of the form \( Gij.\text{to} = Gjh.\text{from} \).
The SELECT-clause is of the from SELECT DISTINCT and consists of all column references of the form $G_{ij}.to$ when $x_{ij}$ is a distinguished node in $P$, plus one reference of the form $G_{1k}.from$ if the root node is distinguished.

The SQL query for the tree in Figure 10 with $\Pi = \{x_2\}$ and $\Sigma = \emptyset$ is as follows:

$$E = \text{SELECT } G_{12}.from, G_{23}.to, G_{24}.to$$
$$\text{FROM } G \text{ as } G_{12}, G \text{ as } G_{23}, G \text{ as } G_{24}$$
$$\text{WHERE } G_{12}.to = G_{23}.from \text{ AND } G_{12}.to = G_{24}.from$$

But also when $\Sigma \neq \emptyset$, we can compute $\text{FreqTab}_{\Pi, \Sigma}$ by a single SQL query. Note that we thus compute the frequency of a large number of instantiated tree patterns in parallel! We proceed as follows:

1. we formulate the pattern $(\Pi, \emptyset)$ in SQL; call the resulting expression $E$

2. We then take the natural join of $E$ and $\text{CanTab}_{\Pi, \Sigma}$, group by $\Sigma$, and count each group.

The join with the candidacy table ensures that only candidate patterns are counted.

For instance, the SQL query to compute the frequency table for the tree in Figure 10, with $\Pi = \{x_2\}$ and $\Sigma = \{x_1, x_3\}$, with $E$ as above, is as follows:

$$\text{SELECT } E.x1, E.x3, \text{COUNT(*)}$$
$$\text{FROM } E, \text{ CanTab}_{\{x_2\}, \{x_1, x_3\}} \text{ CT}$$
$$\text{WHERE } E.x1 = \text{CT}.x1 \text{ AND } E.x3 = \text{CT}.x3$$
$$\text{GROUP BY } E.x1, E.x3 \text{ HAVING COUNT(*) } \geq \text{minsup}$$

It goes without saying that, whenever the frequency table of a tree pattern is found to be empty, the search for more specialized patterns is pruned at that point.

### 4.4.3 The algorithm

Putting everything together so far, the algorithm is given in Algorithm 1. In outline it is a double Apriori algorithm [3], where the sets $\Pi$ form one dimension of itemsets, and the sets $\Sigma$ another. A graphical illustration of the algorithm is given in Figure 11. In this illustration we use tries (or prefix-trees) to store the itemsets. A trie [5, 7, 26] is commonly used in implementations of the Apriori algorithm.
Algorithm 1 Levelwise search for frequent tree patterns.

1: for each unordered, rooted tree $T$ do
2: $X :=$ set of nodes of $T$
3: $p := 0; \mathcal{P}_0 := \{\emptyset\}$
4: repeat
5: for each $\Pi \in \mathcal{P}_p$ do
6: Compute $\text{FreqTab}_{\Pi,\emptyset}$ in SQL
7: if $\text{FreqTab}_{\Pi,\emptyset} \neq \emptyset$ then
8: $s := 1$
9: $\mathcal{S}_1 := \{\{z\} \mid z \in X - \Pi\}$
10: repeat
11: for each $\Sigma \in \mathcal{S}_s$ do
12: if $p = 0$ and $s = 1$ then
13: $\text{CanTab}_{\Pi,\Sigma} :=$ set of nodes of $G$
14: else
15: $\text{CanTab}_{\Pi,\Sigma} := \bowtie \{\text{FreqTab}_{\Pi',\Sigma'} \mid (\Pi', \Sigma')$ parent of $(\Pi, \Sigma)\}$
16: end if
17: Compute $\text{FreqTab}_{\Pi,\Sigma}$ in SQL
18: if $\text{FreqTab}_{\Pi,\Sigma} = \emptyset$ then
19: remove $\Sigma$ from $\mathcal{S}_s$ \{ $\Sigma$ is pruned away \}
20: end if
21: end for
22: $\mathcal{S}_{s+1} := \{\Sigma \subseteq X - \Pi \mid \#\Sigma = s + 1$
23: and each $s$-subset of $\Sigma$ is in $\mathcal{S}_s\}$
24: $s := s + 1$
25: until $\mathcal{S}_s = \emptyset$
26: else
27: remove $\Pi$ from $\mathcal{P}_p$ \{ $\Pi$ is pruned away \}
28: end if
29: end for
30: $\mathcal{P}_{p+1} := \{\Pi \subseteq X \mid \#\Pi = p + 1 \text{ and each } p \text{-subset of } \Pi \text{ is in } \mathcal{P}_p\}$
31: $p := p + 1$
32: until $\mathcal{P}_p = \emptyset$
33: end for
Figure 11: Illustration of the algorithm in Algorithm 1

Figure 12: Data graph $G$ and unordered rooted tree $T$ for the example run in Section 4.4.4.
4.4.4 Example run

In this Section we give an example run of the proposed algorithm in Algorithm 1. Consider the example data graph $G$ in Figure 12(a) the unordered rooted tree $T$ in Figure 12(b) and let the minimum support threshold be 3.

The example run then looks as follows:

| $\Pi$ | $\Sigma$ | $P$ | $CanTab$ | $FreqTab$ |
|-------|---------|-----|----------|---------|
| $\emptyset$ | $\emptyset$ | $x_1 \xrightarrow{} x_2 \xrightarrow{} x_3$ | $\emptyset$ | $\emptyset$ |
| $\emptyset$ | $\{x_1\}$ | $x_2 \xrightarrow{} x_1 \xrightarrow{}$ | $\emptyset$ | $\{x_1\}$ |
| $\emptyset$ | $\{x_2\}$ | $x_3 \xrightarrow{} x_1 \xrightarrow{}$ | $\emptyset$ | $\emptyset$ |
| $\emptyset$ | $\{x_3\}$ | $x_1 \xrightarrow{} x_3 \xrightarrow{}$ | $\emptyset$ | $\emptyset$ |
| $\emptyset$ | $\{x_1, x_2\}$ | $x_2 \xrightarrow{} x_1 \xrightarrow{}$ | $\emptyset$ | $\emptyset$ |
| $\emptyset$ | $\{x_1, x_3\}$ | $x_3 \xrightarrow{} x_1 \xrightarrow{}$ | $\emptyset$ | $\emptyset$ |
| $\emptyset$ | $\{x_2, x_3\}$ | $x_3 \xrightarrow{} x_2 \xrightarrow{}$ | $\emptyset$ | $\emptyset$ |
| $\emptyset$ | $\{x_1, x_2, x_3\}$ | $x_3 \xrightarrow{} x_2 \xrightarrow{}$ | $\emptyset$ | $\emptyset$ |
| $\{x_1\}$ | $\emptyset$ | $x_2 \xrightarrow{} x_3 \xrightarrow{}$ | $\emptyset$ | $\emptyset$ |
| $\{x_1\}$ | $\{x_2\}$ | $x_3 \xrightarrow{} x_2 \xrightarrow{}$ | $\emptyset$ | $\emptyset$ |
| $\{x_1\}$ | $\{x_3\}$ | $x_3 \xrightarrow{} x_1 \xrightarrow{}$ | $\emptyset$ | $\emptyset$ |
| $\{x_1\}$ | $\{x_2, x_3\}$ | $\exists \ \sigma_2 \ \sigma_3$ | $\text{FreqTab}_{0, \{x_2, x_3\}} \ \ni \ \text{FreqTab}_{\{x_1\}, \{x_2\}} \ \ni \ \text{FreqTab}_{\{x_1\}, \{x_3\}}$ | 0 |
|---|---|---|---|---|
| $\{x_2\}$ | $\emptyset$ | $x_1 \ \exists \ x_3$ | $\sigma_1$ | $\sigma_3$ |
| | | | $\text{Pruned}$ | 0 |
| $\{x_2\}$ | $\{x_1\}$ | $\exists \ \sigma_1 \ x_3$ | $\text{FreqTab}_{0, \{x_1\}} \ \ni \ \text{FreqTab}_{\{x_2\}, \emptyset}$ | $\sigma_1$ |
| | | | $\text{Freq}$ | 3 |
| $\{x_2\}$ | $\{x_3\}$ | $\exists \ x_1 \ \sigma_3$ | $\text{FreqTab}_{0, \{x_3\}}$ | 0 |
| $\{x_2\}$ | $\{x_1, x_3\}$ | $\exists \ \sigma_1 \ \sigma_3$ | $\text{Pruned}$ | 0 |
| $\{x_3\}$ | $\emptyset$ | $x_1 \ \exists \ x_2$ | $\sigma_1$ | 0 |
| | | | $\sigma_3$ | 3 |
| $\{x_3\}$ | $\{x_1\}$ | $\exists \ x_1 \ \sigma_3$ | $\text{FreqTab}_{0, \{x_1\}} \ \ni \ \text{FreqTab}_{\{x_3\}, \emptyset}$ | $\sigma_1$ |
| | | | $\text{Freq}$ | 3 |
| $\{x_3\}$ | $\{x_2\}$ | $\exists \ x_1 \ \sigma_2$ | $\text{FreqTab}_{0, \{x_2\}} \ \ni \ \text{FreqTab}_{\{x_3\}, \emptyset}$ | 0 |
| $\{x_3\}$ | $\{x_1, x_2\}$ | $\exists \ \sigma_1 \ \sigma_2$ | $\text{Pruned}$ | 0 |
| $\{x_1, x_2\}$ | $\emptyset$ | $\exists \ \sigma_2 \ \sigma_3$ | $\text{Freq}$ | 6 |
| | | | 0 |
| $\{x_1, x_2\}$ | $\{x_3\}$ | $\exists \ \sigma_3$ | $\text{FreqTab}_{\{x_1\}, \{x_3\}} \ \ni \ \text{FreqTab}_{\{x_1\}, \{x_3\}} \ \ni \ \text{FreqTab}_{\{x_2\}, \emptyset}$ | 0 |
| | | | $\text{Freq}$ | 6 |
4.5 Equivalence among Tree Patterns

In this Section we make a number of modifications to the algorithm described so far, so as to avoid duplicate work.

As an example of duplicate work, consider the parameterized tree pattern $P_1$ from the example run in Section 4.4.4 (\(\Pi = \{x_2\}\) and \(\Sigma = \emptyset\)):

\[
\exists \sigma_2 x_3
\]

and the parameterized tree pattern $P_2$:

\[
\exists x_3 x_2
\]

Clearly, $P_1$ and $P_2$ have the same answer set for all data graphs $G$, up to renaming of the distinguished variables ($x_2$ by $x_3$). However, these patterns have different underlying trees, and hence Algorithm 1 will compute the answer set for both patterns (line 6). The answer set of $P_1$ is computed before the answer set of $P_2$, since our algorithm is incremental in the number of nodes of $T$. Hence, we can conclude that our algorithm performs some duplicate work which we want to avoid.

Another example of duplicate work our algorithm performs: Consider the parameterized tree pattern $P_3$ from the example run in Section 4.4.4 (\(\Pi = \{x_1\}\) and \(\Sigma = \{x_2\}\)):

\[
\exists \sigma_2 x_3
\]

and the parameterized tree pattern $P_4$ also from the example run in Section 4.4.4 (\(\Pi = \{x_1\}\) and \(\Sigma = \{x_3\}\)):

\[
\exists x_2 \sigma_3
\]
As one can see in Section 4.4.4, these two parameterized patterns have the same instantiations for all data graphs $G$, up to renaming of the parameters ($\sigma_2$ by $\sigma_3$), and for each instantiation, the same answer set for all data graphs $G$, up to renaming of the distinguished variables ($x_3$ by $x_2$). However, when we look at the outline of our algorithm in Algorithm 1 we see that for both patterns the candidacy and frequency tables are computed between line 11 and line 17. Hence, we can conclude again that our algorithm performs duplicate work that we want to avoid.

In the rest of this Section we formalize the duplicate work our algorithm performs, and we make a number of modifications to the algorithm described so far, so as to avoid the duplicate work.

4.5.1 Equivalency

Intuitively we call two parameterized tree patterns equivalent if they have the same answer sets and the same parameter assignments for all data graphs $G$, up to renaming of the parameters and the distinguished variables. For instance, the parameterized tree patterns $P_1$ and $P_2$ from above we call equivalent, as the tree patterns $P_3$ and $P_4$ from above.

To define equivalent parameterized tree patterns formally we introduce the notion of $(\delta, \rho)$-equivalence.

$(\delta, \rho)$-Equivalence Let $P_1$ and $P_2$ be two parameterized tree patterns and $\rho$ a parameter correspondence from $P_1$ to $P_2$ (recall Section 3.3). We define an answer set correspondence from $P_1$ to $P_2$ as any mapping $\delta$: $\Delta_1 \rightarrow \Delta_2$. Furthermore, assume that $\delta$ and $\rho$ are bijections. We then say that $P_1$ and $P_2$ are $(\delta, \rho)$-equivalent, denoted by $P_1 \equiv_{\delta, \rho} P_2$, if for all data graphs $G$, and all parameter assignments $\alpha_2: \Sigma_2 \rightarrow U$, we have $P_2^{\alpha_2}(G) \circ \delta = P_1^{\alpha_2 \circ \rho}(G)$, where $P_2^{\alpha_2}(G) \circ \delta$ denotes the set $\{ f \circ \delta : f \in P_2^{\alpha_2}(G) \}$.

For example, consider the two parameterized tree patterns in Figure 13, and let $\rho: \Sigma_a \rightarrow \Sigma_b$ be as follows:

| $\rho$ |
|---|
| $\sigma_1$ | $\sigma_1$ |
| $\sigma_2$ | $\sigma_3$ |
| $\sigma_3$ | $\sigma_2$ |

and let $\delta: \Delta_a \rightarrow \Delta_b$ be as follows:

| $\delta$ |
|---|
| $x_1$ | $x_3$ |
| $x_2$ | $x_1$ |
| $x_3$ | $x_2$ |

The two parameterized tree patterns are clearly $(\delta, \rho)$-equivalent, as are the three parameterized tree patterns shown in Figure 14 with an empty parameter correspondence $\rho$ and $\delta$ the identity.

The parameter correspondence $\rho$ is a bijection in the definition of $\rho$-equivalence, since intuitively we want equivalent parameterized tree patterns to have essentially the same set of instantiations. Hence it is necessary that the tree patterns have the same number of parameters. Intuitively we also want equivalent
tree patterns to have the same answer sets up to renaming of the distinguished variables. That is the reason why an answer set correspondence is introduced that is a bijection.

We then define equivalent parameterized tree patterns as follows:

**Equivalent parameterized tree patterns** We call two parameterized tree patterns $P_1$ and $P_2$ equivalent if $P_1$ is $(\delta, \rho)$-equivalent with $P_2$ for some bijective parameter correspondence $\rho$ and some bijective answer set correspondence $\delta$.

Note that there can exist more than one parameter correspondence $\rho$ and more than one answer set correspondence $\delta$ for which the two parameterized tree patterns are $(\delta, \rho)$-equivalent. An illustration of this is given in Figure 15.

Let $\rho_1 : \Sigma_a \rightarrow \Sigma_b$, $\delta_1 : \Delta_a \rightarrow \Delta_b$, $\rho_2 : \Sigma_a \rightarrow \Sigma_b$ and $\delta_2 : \Delta_a \rightarrow \Delta_b$ be as follows: $\rho_1$ is the identity; $\delta_1$ is the identity and

| $\rho_2$ | $\delta_2$ |
|-------|-------|
| $\sigma_1$ | $\sigma_1$ |
| $\sigma_2$ | $\sigma_2$ |
| $x_1$ | $x_1$ |
| $x_2$ | $x_4$ |
| $x_3$ | $x_5$ |
| $x_4$ | $x_2$ |
| $x_5$ | $x_3$ |

Then the two tree patterns in Figure 15 are clearly $(\delta_1, \rho_1)$-equivalent and $(\delta_2, \rho_2)$-equivalent.

Equivalence as just defined is a semantical property, referring to all possible data graphs, and it is not immediately clear how one could decide this property syntactically. The required syntactical notion is given by the following Lemma and Corollary.

**Lemma 4.** Consider two parameterized tree patterns $P_1$ and $P_2$, $\delta : \Delta_1 \rightarrow \Delta_2$ a bijective answer set correspondence, and $\rho : \Sigma_1 \rightarrow \Sigma_2$ a bijective parameter correspondence. Then $P_1$ and $P_2$ are $(\delta, \rho)$-equivalent if there exist bijections $\rho_1 : \Delta_a \rightarrow \Delta_b$ and $\delta_1 : \Sigma_a \rightarrow \Sigma_b$ such that $P_1$ and $P_2$ are $(\delta_1, \rho_1)$-equivalent.
correspondence. Then $P_1 \equiv_{P}^\delta P_2$ if and only if we have the following containment relations among the tree queries $(H_1, P_1)$ and $(\delta(H_1), P_2)$, with $H_1$ the pure head of $P_1$ (cfr. Section [7]):

1. $(\delta(H_1), P_2) \subseteq_{\rho} (H_1, P_1)$; and
2. $(H_1, P_1) \subseteq_{\rho^{-1}} (\delta(H_1), P_2)$

Proof. Let us start with the if direction. We need to prove that for every parameter assignment $\alpha_2$ for $P_2$, and every data graph $G$ that $P_2^{\alpha_2}(G) \circ \delta = P_1^{\alpha_2 \circ \rho}(G)$.

We know that $(\delta(H_1), P_2)^{\alpha_2}(G) \subseteq (H_1, P_1)^{\alpha_2 \circ \rho}(G)$ since $(\delta(H_1), P_2) \subseteq_{\rho} (H_1, P_1)$. We may rewrite this as: $P_2^{\alpha_2}(G) \circ \delta \subseteq P_1^{\alpha_2 \circ \rho}(G)$ since $H_1$ is an enumeration of $\Delta_1$.

We also know that $(H_1, P_1)^{\alpha_1}(G) \subseteq (\delta(H_1), P_2)^{\alpha_1 \circ \rho^{-1}}(G)$ for every parameter assignment $\alpha_1$ for $P_1$ since $(H_1, P_1) \subseteq_{\rho^{-1}} (\delta(H_1), P_2)$. Now take $\alpha_1 = \alpha_2 \circ \rho$. We then have $(H_1, P_1)^{\alpha_2 \circ \rho}(G) \subseteq (\delta(H_1), P_2)^{\alpha_2}(G)$. Again since $H_1$ is an enumeration of $\Delta_1$ we may rewrite this as: $P_2^{\alpha_2}(G) \subseteq P_1^{\alpha_2 \circ \rho}(G) \circ \delta$. Hence we can conclude that $P_2^{\alpha_2}(G) \circ \delta = P_1^{\alpha_2 \circ \rho}(G)$.

Let us then look at the only-if direction. To prove that $(\delta(H_1), P_2) \subseteq_{\rho} (H_1, P_1)$, we will show that for every $\alpha_2$ parameter assignment for $P_2$, and every data graph $G$, we have $(\delta(H_1), P_2)^{\alpha_2}(G) \subseteq (H_1, P_1)^{\alpha_2 \circ \rho}(G)$. Since $P_2^{\alpha_2}(G) \circ \delta = P_1^{\alpha_2 \circ \rho}(G)$, we have $(\delta(H_1), P_2)^{\alpha_2}(G) = (H_1, P_1)^{\alpha_2 \circ \rho}(G)$, and hence clearly $(\delta(H_1), P_2)^{\alpha_2}(G) \subseteq (H_1, P_1)^{\alpha_2 \circ \rho}(G)$.

To prove that $(H_1, P_1) \subseteq_{\rho^{-1}} (\delta(H_1), P_2)$, we will show that for every $\alpha_1$ parameter assignment for $P_1$, and every data graph $G$, we have $(H_1, P_1)^{\alpha_1}(G) \subseteq (\delta(H_1), P_2)^{\alpha_1 \circ \rho^{-1}}(G)$. We know that for every $\alpha_2$ parameter assignment for $P_2$, we have $P_2^{\alpha_2}(G) \circ \delta = P_1^{\alpha_2 \circ \rho}(G)$. Now take $\alpha_2 = \alpha_1 \circ \rho^{-1}$. We then have $P_2^{\alpha_2}(G) \circ \delta = P_1^{\alpha_1}(G)$, hence $(\delta(H_1), P_2)^{\alpha_1 \circ \rho^{-1}}(G) = (H_1, P_1)^{\alpha_1}(G)$, hence clearly $(H_1, P_1)^{\alpha_1}(G) \subseteq (\delta(H_1), P_2)^{\alpha_1 \circ \rho^{-1}}(G)$. \qed

**Corollary 1.** Consider two parameterized tree patterns $P_1$ and $P_2$. Then $P_1$ is equivalent with $P_2$ if and only if there exist:

1. a bijective answer set correspondence $\delta : \Delta_1 \rightarrow \Delta_2$;
2. a bijective parameter correspondence $\rho : \Sigma_1 \rightarrow \Sigma_2$;
3. a $p$-containment mapping $f_1 : (H_1, P_1) \rightarrow (\delta(H_1), P_2)$; and
4. a $p^{-1}$-containment mapping $f_2 : (\delta(H_1), P_2) \rightarrow (H_1, P_1)$.

with $H_1$ the pure head for $P_1$. 27
Of course, we want to avoid that our algorithm considers some parameterized tree pattern \( P_2 \) if it is equivalent to an earlier considered parameterized tree pattern \( P_1 \). Since our algorithm generates trees in increasing sizes, there are two cases to consider:

**Case A:** \( P_1 \) has fewer nodes than \( P_2 \).

**Case B:** \( P_1 \) and \( P_2 \) have the same number of nodes.

Armed with the above Lemma and Corollary, we can now analyze the above two cases.

### 4.5.2 Case A: Redundancy checking

Let us start by defining the notion of a redundancy.

**Redundant subtree** A redundant subtree \( R \), is a subtree of a parameterized tree pattern \( P \), such that removing \( R \) from \( P \) yields a parameterized tree pattern \( P' \) that is equivalent with \( P \).

For example, the first two parameterized tree patterns in Figure 14 indeed contain a redundant subtree.

The following lemma shows that two parameterized tree patterns with different numbers of nodes can only be equivalent if the largest one contains redundant subtrees:

**Lemma 5.** Consider two parameterized tree patterns \( P \) and \( P' \), and the number of nodes of \( P' \) is smaller than the number of nodes of \( P \). Then \( P \) can only be equivalent with \( P' \) if \( P \) contains redundant subtrees.

**Proof.** Since \( P \) and \( P' \) are equivalent we know from Corollary 1 that the following exist:

1. an answer set correspondence \( \delta : \Delta \to \Delta' \) that is a bijection;
2. a parameter correspondence \( \rho : \Sigma \to \Sigma' \) that is a bijection;
3. a \( \rho \)-containment mapping \( f_1 : (H, P) \to (\delta(H), P') \); and
4. a \( \rho^{-1} \)-containment mapping \( f_2 : (\delta(H), P') \to (H, P) \).

with \( H \) the pure head for \( P \). Since the number of nodes of \( P' \) is smaller than the number of nodes of \( P \), we know that some subtrees \( R \) of \( P \) are not in the range of \( f_2 \). We will prove that these subtrees \( R \) are redundant subtrees, by showing that \( P \) and \( P - R \) are equivalent.

Since the containment mappings \( f_1 \) and \( f_2 \) exist, we know that in particular the following containment mappings will exist:

1. \( g_1 = f_1|_{P-R} \), a \( \rho \)-containment mapping from \( (H, P - R) \) to \( (\delta(H), P') \), and
2. \( g_2 = f_2 \), a \( \rho^{-1} \)-containment mapping from \( (\delta(H), P') \) to \( (H, P - R) \).
with \( \delta \) and \( \rho \) as above.

Let us now look at the following mappings: \( h_1 = g_2 \circ f_1 \) and \( h_2 = f_2 \circ g_1 \).

By Lemma 1, \( h_1 = g_2 \circ f_1 \) and \( h_2 = f_2 \circ g_1 \) are identity-containment mappings.

Using Corollary 1 we can now conclude that \( P \) and \( P - R \) are (identity, identity)-equivalent and thus \( R \) is a redundant subtree.

From this lemma follows that Case A can only happen if \( P_2 \) contains redundant subtrees. Hence, if we can avoid redundancies, Case A will never occur.

The following lemma provides us with an efficient check for redundancies.

**Redundancy Lemma.** Let \( P \) be a parameterized tree pattern. Then \( P \) has a redundancy if and only if it contains a subtree \( C \) in the form of a linear chain of existential nodes (possibly just a single node), such that the parent of \( C \) has another subtree that is at least as deep as \( C \).

Before we prove this Lemma, let us see some examples. For instance the parameterized tree patterns in Figure 14(a) and Figure 14(b) contain a linear chain of existential nodes that is redundant. In both tree patterns this linear chain is rooted in \( \exists_1 \). Another example of such a redundancy is given in Figure 16. Here the linear chain is rooted in \( \exists_3 \). Note that when we remove the linear chain rooted in \( \exists_3 \), we have a new linear chain rooted in \( \exists_1 \) that is redundant.

**Proof.** Let us refer to a subtree \( C \) as described in the lemma as an “eliminable path”. An eliminable path is clearly redundant, so we only need to prove the only-if direction. Let \( T \) be a redundant subtree of \( P \) that is maximal, in the sense that it is not the subtree of another redundant subtree. Then following Corollary 1 there must be a \( \rho \)-containment mapping \( h \) from \((H, P)\) to \((\delta(H), P - T)\) with \( \rho \) and \( \delta \) bijections and \( H \) the pure head for \( P \). All distinguished variables of \( P \) must be in \( P - T \), since \( \delta \) is a bijection. Also all parameters of \( P \) must be in \( P - T \), since \( \rho \) is also a bijection. So \( T \) consists entirely of existential nodes.

Furthermore, note that \( h \) must fix the root of \( P \), since the height of \( P \) is at least that of \( P - T \).

Any iteration \( h^\alpha \) of \( h \) is a \( \rho^\alpha \)-containment mapping from \((H, P)\) to \((\delta^\alpha(H), P - T)\) by Lemma 1. Moreover, each \( h^\alpha|_{\Delta \cup \Sigma} \) induces a permutation on the set \( \Delta \cup \Sigma \) of distinguished variables and parameters. Since \( \Delta \cup \Sigma \) is finite, there are only a finite number of possible permutations of \( \Delta \cup \Sigma \), namely \(|\Delta \cup \Sigma|\). Hence, there will be an iteration \( h^k|_{\Delta \cup \Sigma} \) and an iteration \( h^{(k+1)}|_{\Delta \cup \Sigma} \) such that
$h^k|_{\Delta \cup \Sigma} = h^{(k+1)}|_{\Delta \cup \Sigma}$. Hence, $h^i|_{\Delta \cup \Sigma}$ is the identity on $\Delta \cup \Sigma$, because

\[
\text{id}|_{\Delta \cup \Sigma} = (h^{-1})^k|_{\Delta \cup \Sigma} \circ h^k|_{\Delta \cup \Sigma} = (h^{-1})^k|_{\Delta \cup \Sigma} \circ h^{(k+1)}|_{\Delta \cup \Sigma} = h^i|_{\Delta \cup \Sigma}
\]

There are now two possible cases.

1. $T$ itself is a linear chain. Let us then look at the parent $p$ of $T$ in $P$. Again there are two possibilities:
   
   (a) $h^i(p) = p$: Since $h^i$ is a $\rho'$-containment mapping from $(H, P)$ to $(\delta'(H), P - T)$ and $T$ is redundant, we know that $T$ must be mapped to another subtree of $p$, $T'$, that is at least as deep as $T$. Hence, $T$ is an eliminable path. An illustration is given in Figure 17(a).
   
   (b) $h^i(p) \neq p$: Then $p$ can only be an existential node. We now have two possibilities:
      
      i. $T$ is the only subtree of $p$. We will show that the subtree $T'$, rooted in $p$ is redundant as well. Clearly we have the following containment relations:
         
         - $h_1 = h^i$, a $\rho'$-containment mapping from $(H, P)$ to $(\delta'(H), P - T')$; and
         - $h_2 = h^{-1}$, a $\rho^\perp$-containment mapping from $(\delta'(H), P - T')$ to $(H, P)$.
      
      with $\delta$ and $\rho$ as above. By Corollary 1, $T'$ is then a redundant subtree. This is in contradiction with the assumption that $T$ is maximal. Hence, it is impossible that $p$ has only one subtree and $p$ is existential. An illustration of this is given in Figure 17(b).
      
      ii. $p$ has more than one subtree. Consider such another subtree $T'$. We will show that all subtrees $T'$ of $p$ consist entirely of existential nodes. Suppose a node $n \in T'$ is not an existential node. We then know that $h^i(n) = n$. However, since $h^i$ is a homomorphism and $p$ is an ancestor of $n$, $h^i(p)$ must be $p$. But this is in contradiction with the assumption that $h^i(p) \neq p$. So $T'$ must consist entirely of existential nodes. Hence this brings us to the second case where $T$ is not a linear chain. An illustration is given in Figure 17(c).

2. $T$ is not a linear chain. An easy induction on the height of $T$, shows that any non-linear tree consisting entirely of existential nodes must contain an eliminable path. If the height of $T$ is 1, there is an eliminable path of a single node: just choose one of the children of the root. If the height of $T$ is $n > 1$, consider the subtree $S$ of the root of $T$ with the smallest height, at most $n - 1$. Then we have two possibilities: If $S$ is a linear chain, we found our eliminable path. And if $S$ is a non-linear chain we know by induction that $S$ will contain an eliminable path. Hence, $T$, and thus also $P$, contains an eliminable path as desired.
Figure 17: Figures to illustrate the proof of the Redundancy Lemma.
As we have seen in Section 4.4, our algorithm introduces existential nodes levelwise, one by one. This makes the redundancy test provided by the redundancy lemma particularly easy to perform. Indeed, if \((\Pi, \Sigma)\) is a parameterized tree patterns of which we already know it has no redundancies, and we make one additional node \(n\) existential, then it suffices to test whether \(n\) thus becomes part of a subtree \(C\) as in the Redundancy Lemma. If so, we will prune the entire search at \(\Pi \cup \{n\}\).

### 4.5.3 Case B: Canonical forms

We may now assume that \(P_1\) and \(P_2\) do not contain redundancies, for if they would, they would have been dismissed already.

Let us start by defining isomorphic parameterized tree patterns.

**Isomorphic Parametrized Tree Patterns** We call two parameterized tree patterns \(P_1\) and \(P_2\) isomorphic if there exists a homomorphism \(f: P_1 \rightarrow P_2\) that is a bijection and that maps distinguished nodes to distinguished nodes, parameters to parameters and existential nodes to existential nodes. We call \(f\) an isomorphism. Since we are working with trees, \(f^{-1}\) is also a homomorphism.

For example, the two parameterized tree patterns in Figure 13 are indeed isomorphic with \(f\) as follows:

|   | \(\sigma_1\) | \(\sigma_2\) | \(\sigma_3\) | \(x_1\) | \(x_2\) | \(x_3\) |
|---|-----------|-----------|-----------|-------|-------|-------|
| \(\exists\) | \(\exists\) | \(\sigma_2\) | \(\sigma_3\) | \(x_1\) | \(x_2\) | \(x_3\) |

Clearly, we have the following:

**Property 1.** Any two isomorphic parameterized tree patterns \(P_1\) and \(P_2\) are equivalent.

**Proof.** Using Corollary 4.4 we have to show that the following exists:

1. a bijective answer set correspondence \(\delta: \Delta_1 \rightarrow \Delta_2\);
2. a bijective parameter correspondence \(\rho: \Sigma_1 \rightarrow \Sigma_2\);
3. a \(\rho\)-containment mapping \(f_1: (H_1, P_1) \rightarrow (\delta(H_1), P_2)\); and
4. a \(\rho^{-1}\)-containment mapping \(f_2: (\delta(H_1), P_2) \rightarrow (H_1, P_1)\).

with \(H_1\) the pure head for \(P_1\).

Since \(P_1\) and \(P_2\) are isomorphic, there exists a homomorphism \(f: P_1 \rightarrow P_2\) that is a bijection and that maps distinguished nodes to distinguished nodes, parameters to parameters and existential nodes to existential nodes.

We now take \(\delta = f|_{\Delta_1}\) and \(\rho = f|_{\Sigma_1}\). Then \(\delta\) and \(\rho\) are bijections since \(f\) is a bijection.

For (3) we will show that \(f\) is \(\rho\)-containment mapping from \((H_1, P_1)\) to \((\delta(H_1), P_2)\), with \(H_1\) the pure head for \(P_1\):
• $f$ is a homomorphism;
• $f$ maps distinguished nodes to distinguished nodes and $f|_{\Delta_1} = \delta$;
• $f$ maps parameters to parameters and $f|_{\Sigma_1} = \rho$; and
• $f(H_1) = \delta(H_1)$.

For (4) we will show that $f^{-1}$ is $\rho^{-1}$-containment mapping from $(\delta(H_1), P_2)$ to $(H_1, P_1)$, with $H_1$ the pure head for $P_1$:

• $f^{-1}$ is a homomorphism since $f$ is a bijection and we are working with trees;
• $f^{-1}$ maps distinguished nodes to distinguished nodes and $f^{-1}|_{\Delta_2} = \delta^{-1}$;
• $f^{-1}$ maps parameters to parameters and $f^{-1}|_{\Sigma_2} = \rho^{-1}$; and
• $f^{-1}(\delta(H_1)) = H_1$.

The following lemma shows that two parameterized tree patterns without redundancies and with the same number of nodes can only be equivalent if they are isomorphic.

**Isomorphism Lemma.** Consider two parameterized tree patterns $P_1$ and $P_2$ without redundancies, and with the same number of nodes. Then $P_1$ and $P_2$ are equivalent if and only if $P_1$ and $P_2$ are isomorphic.

**Proof.** We only need to show the only-if direction.

Since $P_1$ and $P_2$ are equivalent we know that the following exists by Corollary 1:

1. a bijective answer set correspondence $\delta : \Delta_1 \rightarrow \Delta_2$;
2. a bijective parameter correspondence $\rho : \Sigma_1 \rightarrow \Sigma_2$;
3. a $\rho$-containment mapping $f_1 : (H_1, P_1) \rightarrow (\delta(H_1), P_2)$; and
4. a $\rho^{-1}$-containment mapping $f_2 : (\delta(H_1), P_2) \rightarrow (H_1, P_1)$.

with $H_1$ a pure head for $P_1$.

We also know that $P_1$ and $P_2$ have the same number of existential nodes since $P_1$ and $P_2$ have the same number of nodes and $\rho$ and $\delta$ are bijections.

Hence to prove that $P_1$ and $P_2$ are isomorphic, we only need to show that:

1. $f_1$ maps existential nodes to existential nodes; and
2. $f_1|_{\Pi_1}$ is a bijection.

Theoreto, it suffices to prove that $f_1$ is surjective on the existential nodes of $P_2$, because $f_1$ is already a bijection from $\Delta_1 \cup \Sigma_1$ to $\Delta_2 \cup \Sigma_2$.

Assume that $f_1|_{\Pi_1}$ is not surjective. Hence, there will be some existential nodes $p \in \Pi_2$ that are not in the range of $f_1$. Note that these existential nodes $p$ can never have descendants that are parameters or distinguished nodes since $f_1$ is a homomorphism and $\delta$ and $\rho$ bijections. Now fix some $p$ as high as possible
in the tree. Then the entire subtree $R$ rooted in $p$ consists entirely of existential nodes, that are not in the range of $f_1$. We will now show that this subtree $R$ is a redundant subtree in $P_2$. We then have a contradiction since we assume that $P_2$ is redundancy free.

Since the containment mappings $f_1$ and $f_2$ exist, we know that in particular the following containment mappings will exist:

1. $g_1 = f_1$ a $\rho$-containment mapping from $(H_1, P_1)$ to $(\delta(H_1), P_2 - R)$, and
2. $g_2 = f_2|_{P_2 - R}$ a $\rho^{-1}$-containment mapping from $(\delta(H_1), P_2 - R)$ to $(H_1, P_1)$.

with $\delta$ and $\rho$ as above.

Let us now look at the following mappings $h_1 = f_1 \circ g_2$ and $h_2 = g_1 \circ f_2$. By Lemma 1, $h_1$ and $h_2$ are identity-containment mappings. Using Corollary 1, we can now conclude that $P_2$ and $P_2 - R$ are equivalent, and thus $R$ is a redundant subtree.

From the above lemma it follows that Case B can only happen if $P_1$ and $P_2$ are actually isomorphic. In particular, $P_1$ and $P_2$ have the same underlying tree.

So, in our algorithm, we need an efficient way to avoid isomorphic parameterized tree patterns based on the same tree $T$.

Fortunately, there is a standard way to do this, by working with canonical forms of parameterized tree patterns. Consider a pair $(\Pi, \Sigma)$, as in Section 4.4. We can view this pair as a labeling of $T$: all nodes in $\Pi$ get the same generic label ‘$\exists$’; all nodes in $\Sigma$ get ‘$\sigma$’; and all distinguished nodes get ‘$x$’. We then observe that the patterns $(\Pi_1, \Sigma_1)$ and $(\Pi_2, \Sigma_2)$ are isomorphic iff there is a tree isomorphism between the corresponding labeled versions of $T$ that respects the labels.

In order to represent each pair $(\Pi, \Sigma)$ uniquely up to isomorphism, we can rather straightforwardly refine the canonical ordering of the underlying unlabeled tree $T$, which we already have (Section 4.3), to take into account the node labels. Furthermore, the classical linear-time algorithm to canonize a tree [4] generalizes straightforwardly to labeled trees. A nice review of these generalizations has been given by Chi, Yang and Muntz [10].

We will omit the details of the canonical form; in fact, there are several ways to realize it. All that is important is that we can check in linear time whether a pair is canonical; that a pair can be canonized in linear time; and that two pairs are isomorphic if and only if their canonical forms are identical.

**Example.** We can refine the level sequence introduced in Section 4.3 to a refined level sequence for parameterized tree patterns $P$ as follows: if the tree pattern $P$ consists of $n$ nodes, then the refined level sequence is now a sequence of $n$ elements, where the $i$th element is the depth of the $i$th node in preorder in the pattern, followed by a ‘$d$’ if the node is distinguished; followed by a ‘$e$’ if the node is existential and followed by a ‘$p$’ if the node is a parameter. The canonical ordering of a parameterized tree pattern $P$, is then the ordering of $P$ that yields the lexicographically maximal refined level sequence, among all orderings of $P$. Then the refined level sequences for the parameterized tree patterns in Figure 13 are:

(a) 0p1e2p2d1d2p2d

34
Figure 18: The tree pattern in (b) is a parent of a tree pattern in (a), and (c) is the canonical version of (b).

(b) 0p1d2d2p1e2d2p and (a) is the canonical one.

Armed by the canonical form, we are now in a position to describe how the algorithm of Section 4.4 must be modified to avoid equivalent parameterized tree patterns. First of all, we only work with patterns \((\Pi, \Sigma)\) in canonical form; the others are dismissed. However, the problem then arises that a parent pattern \((\Pi', \Sigma')\), where we omit a variable from either \(\Pi\) or \(\Sigma\) as described in Section 4.4, might be non-canonical. In that case the frequency table for \((\Pi', \Sigma')\) will not exist. We can solve this by canonizing \((\Pi', \Sigma')\) to its canonical version \((\Pi'', \Sigma'')\), and remembering the renaming of variables this entails. The table \(FreqTab_{\Pi'', \Sigma''}\) can then serve in place of \(FreqTab_{\Pi', \Sigma'}\), after we have applied the inverse renaming to its column headings.

This does not completely solve the problem, however. Indeed the frequency table of \((\Pi'', \Sigma'')\) might not yet have been computed. For example, consider the parameterized tree pattern in Figure 18(a) and one of its parents in Figure 18(b). The canonical version of this parent, using the canonical ordering from the previous example, is shown in Figure 18(c). Using the current order for computing the frequency tables as in Algorithm 1, the frequency table for the pattern in Figure 18(c) is not yet computed, when we want to compute the frequency for the pattern in Figure 18(a).

We can solve this by changing the order in which we compute the frequency tables. We work with increasing levels: in level \(i\) we compute the frequency tables for all pairs \((\Pi, \Sigma)\), where \(#\Pi + #\Sigma = i\). If we use this order, we are sure that when we compute the frequency table of a pair \((\Pi, \Sigma)\), all frequency tables of pairs \((\Pi', \Sigma')\) with \(#\Pi' + #\Sigma' < (\#\Pi + #\Sigma)\), have been computed.

### 4.5.4 The Algorithm

The final algorithm is now given in Algorithm 2. The outline for canonizing a parameterized tree pattern in given in Function 3.

### 4.5.5 Example run

In this Section we give an example run of the final algorithm in Algorithm 2. We use the same data graph \(G\); unordered rooted tree \(T\); and minimum support threshold, 3, as in the example in Section 4.4.4.
Algorithm 2 Levelwise search for frequent tree patterns with equivalence checking.

1: for each unordered, rooted tree $T$ do
2:   level := number of nodes of $T$
3:   $i := 0$
4:   $C_0 := \{ (\emptyset, \emptyset) \}; \quad F := \emptyset$
5:   while $i \leq$ level AND $C_i \neq \emptyset$ do
6:     {Candidate evaluation}
7:     for each pattern $(\Pi, \Sigma)$ in $C_i$ do
8:       if $\Sigma = \emptyset$ then
9:         Compute $\text{FreqTab}_{\Pi, \Sigma}$ in SQL
10:       else
11:         if $(\#\Sigma = 1$ AND $\#\Pi = 0)$ then
12:            $\text{CanTab}_{\Pi, \Sigma} := \text{set of nodes of } G$
13:         else
14:            $\text{CanTab}_{\Pi, \Sigma} := \{ f^{-1}(\text{FreqTab}_{\Pi', \Sigma'}) \mid (\Pi', \Sigma') \text{ parent of } (\Pi, \Sigma) \}$
15:       end if
16:     end if
17:     end if
18:     Compute $\text{FreqTab}_{\Pi, \Sigma}$ in SQL
19:     if $(\text{FreqTab}_{\Pi, \Sigma} \neq \emptyset)$ then
20:       $F = F \cup \{ (\Pi, \Sigma) \}$
21:     end if
22:   end for
23:   {Candidate generation}
24:   $C_{i+1} = \{ (\Pi, \Sigma) \mid \text{all parents } (\Pi', \Sigma') \text{ of } (\Pi, \Sigma) \text{ are in } F \}$
25:   {Equivalence Check}
26:   for each pattern $(\Pi, \Sigma)$ in $C_{i+1}$ do
27:     if $(\Pi, \Sigma)$ contains a redundancy) then
28:       remove $(\Pi, \Sigma)$ from $C_{i+1}$
29:     else if $(\Pi, \Sigma)$ is not canonical) then
30:       remove $(\Pi, \Sigma)$ from $C_{i+1}$
31:     end if
32:   end for
33:   $i := i + 1$
34: end while

Function 3 Canonize $(\Pi', \Sigma')$ based on $T$

1: $(\Pi_c, \Sigma_c) := \text{canonization of } (\Pi', \Sigma') \text{ based on } T$
2: $f := \text{isomorphism from } (\Pi_c, \Sigma_c) \text{ to } (\Pi', \Sigma')$
3: return $(f, (\Pi_c, \Sigma_c))$
Note that there are two important differences between this run and the run in Section 4.4.4:

1. duplicate work is avoided: equivalent tree patterns are not generated; and
2. the order for computing the tree patterns is different in the sense that here, the tree patterns are generated in levels, as explained in Section 4.5.3.

The example run then looks as follows:

| Π | Σ | P | CanTab | FreqTab |
|---|---|---|--------|---------|
| | | | Level 0 | |
| ∅ | ∅ | x₁ \(\xrightarrow{σ_1}\) x₂ \(\xrightarrow{σ_2}\) x₃ | | Freq 15 |
| | | | Level 1 | |
| ∅ | \{x₁\} | x₁ \(\xrightarrow{σ_1}\) x₂ \(\xrightarrow{σ_2}\) x₃ nodes of G | | σ₁ Freq \begin{tabular}{ll} 0 & 9 \\ 2 & 4 \end{tabular} |
| ∅ | \{x₂\} | x₁ \(\xrightarrow{σ_1}\) x₂ \(\xrightarrow{σ_2}\) x₃ nodes of G | | σ₂ Freq \begin{tabular}{ll} 1 & 3 \\ 2 & 3 \\ 3 & 3 \\ 4 & 3 \end{tabular} |
| ∅ | \{x₃\} | x₁ \(\xrightarrow{σ_1}\) x₂ \(\xrightarrow{σ_2}\) x₃ | | Equivalent with (∅, \{x₂\}) |
| \{x₁\} | ∅ | x₁ \(\xrightarrow{σ_1}\) x₂ \(\xrightarrow{σ_2}\) x₃ | | Freq 14 |
| \{x₂\} | ∅ | x₁ \(\xrightarrow{σ_1}\) x₂ \(\xrightarrow{σ_2}\) x₃ | | Redundancy |
| \{x₃\} | ∅ | x₁ \(\xrightarrow{σ_1}\) x₂ \(\xrightarrow{σ_2}\) x₃ | | Equivalent with (\{x₂\}, ∅) |
| | | | Level 2 | |
| ∅ | \{x₁, x₂\} | x₁ \(\xrightarrow{σ_1}\) x₂ \(\xrightarrow{σ_2}\) x₃ | | \begin{tabular}{lll} σ₁ & σ₂ & Freq \\ 0 & 1 & 3 \\ 0 & 2 & 3 \\ 0 & 3 & 3 \end{tabular} |
| ∅ | \{x₁, x₃\} | x₁ \(\xrightarrow{σ_1}\) x₂ \(\xrightarrow{σ_2}\) x₃ | | Equivalent with (∅, \{x₁, x₂\}) |
| ∅ | \{x₂, x₃\} | x₁ \(\xrightarrow{σ_1}\) x₂ \(\xrightarrow{σ_2}\) x₃ | | \begin{tabular}{ll} σ₁ & σ₂ \\ 0 & 1 \\ 0 & 2 \\ 0 & 3 \end{tabular} \begin{tabular}{l} FreqTab₀,\{x₂\} \\ \& \ \ FreqTab₀,\{x₃\} \end{tabular} |
\[
\begin{array}{|c|c|c|c|}
\hline
\{x_1\} & \{x_2\} & \exists & \sigma_2 \xrightarrow{x_3} \sigma_3 \\
\hline
{FreqTab}_{\emptyset,(x_2)} & {FreqTab}_{\{x_1\},(\emptyset)} & \sigma_2 & \begin{array}{c}
\begin{array}{c}
1 \\
2 \\
3
\end{array}
\end{array} \begin{array}{c}
\begin{array}{c}
Freq \\
3 \\
3 \\
3
\end{array}
\end{array}
\hline
{FreqTab}_{\emptyset,(x_1)} & {FreqTab}_{\{x_1\},(x_2)} & \sigma_2 & \begin{array}{c}
\begin{array}{c}
1 \\
2 \\
3
\end{array}
\end{array} \begin{array}{c}
\begin{array}{c}
Freq \\
3 \\
3 \\
3
\end{array}
\end{array}
\hline
\end{array}
\]

1. `{x_1}` `{x_3}` \(\exists\) Equivalent with \((\{x_1\}, \{x_2\})\)
2. `{x_2}` `{x_1}` \(\exists\) Redundancy
3. `{x_2}` `{x_3}` \(\exists\) Redundancy
4. `{x_3}` `{x_1}` \(\exists\) Equivalent with \((\{x_2\}, \{x_1\})\)
5. `{x_3}` `{x_2}` \(\exists\) Equivalent with \((\{x_2\}, \{x_3\})\)
6. `{x_1, x_2}` `{\emptyset}` \(\exists\) Redundancy
7. `{x_1, x_3}` `{\emptyset}` \(\exists\) Equivalent with \((\{x_1, x_2\}, \emptyset)\)
8. `{x_2, x_3}` `{\emptyset}` \(\exists\) Redundancy

**Level 3**

\[
\begin{array}{|c|c|c|c|}
\hline
\emptyset & \{x_1, x_2, x_3\} & \exists & \sigma_2 \xrightarrow{x_1} \sigma_3 \\
\hline
\sigma_1 & Pruned & \sigma_2 & \sigma_3 \\
\hline
{FreqTab}_{\emptyset,(x_2,x_3)} & {FreqTab}_{\{x_1\},(x_2)} & {FreqTab}_{\{x_1\},(x_3)} & \emptyset \\
\hline
{FreqTab}_{\emptyset,(x_1)} & {FreqTab}_{\{x_1\},(x_2)} & {FreqTab}_{\{x_1\},(x_3)} & \emptyset \\
\hline
\end{array}
\]

9. `{x_1}` `{x_2, x_3}` \(\exists\)
10. `{x_2}` `{x_1, x_3}` \(\exists\) Pruned
11. `{x_3}` `{x_1, x_2}` \(\exists\) Equivalent with \((\{x_2\}, \{x_1, x_3\})\)
4.6 Result Management: Pattern Database

When the algorithm is terminated, its final output consists of a set of frequency tables for each tree $T$ that was investigated. All frequency tables are kept in a relational database that we call the pattern database. The pattern database is an ideal platform for an interactive tool for browsing the frequent queries. We developed such a browser called Certhia and discuss it in Section 6.

The pattern database is also an ideal platform for tree query association mining as will be described in Section 5.

5 Mining Tree-Query Associations

In this Section we present an algorithm for mining confident tree-query associations in a large data graph. Recall from Section 5.4 that a parameterized association rule (pAR) is something of the form $Q_1 \Rightarrow \rho Q_2$, with $Q_1$ and $Q_2$ parameterized tree queries, $\rho : \Sigma_1 \rightarrow \Sigma_2$ a parameter correspondence, and $Q_2 \subseteq \rho Q_1$. An instantiated association rule (iAR) is a pair $(Q_1 \Rightarrow \rho Q_2, \alpha)$, with $Q_1 \Rightarrow \rho Q_2$ a pAR and $\alpha : \Sigma_2 \rightarrow U$ a parameter assignment for $Q_1 \Rightarrow \rho Q_2$. Also recall that the confidence of an iAR in a data graph $G$ is defined as $\frac{\text{Freq}(Q_2^\alpha)}{\text{Freq}(Q_1 \Rightarrow \rho Q_2)}$.

The algorithm presented in this Section finds all iARs of the form $(Q_{\text{left}} \Rightarrow \rho Q_{\text{right}}, \alpha)$ that are confident and frequent in a given data graph $G$ for a given $\text{lhs} Q_{\text{left}}$. Before presenting the algorithm we first show that we do not need to tackle the problem in its full generality.

5.1 Problem Reduction

In this section we show that, without loss of generality, we can focus on the case where the given lhs tree query $Q_{\text{left}}$ is pure in the sense that was defined in Section 4.1. We will also show that this restriction can not be imposed on the rhs tree queries to be output. We also make a remark regarding “free constants” in the head of a tree query.

Pure lhs's Assume that all possible variables (nodes of tree patterns) have been arranged in some fixed but arbitrary order. Recall then from Section 4.1 that we call a parameterized tree query $Q = (H, P)$ pure when $H$ consists of the enumeration in order and without repetitions of all distinguished variables of $P$. In particular $H$ can not contain parameters. We call $H$ the pure head for
As an illustration, the lhs of rule (a) of Figure 19 is impure, while the lhs of rule (b) is pure.

Consider the pARs in Figure 19(a) and Figure 19(b) and their instantiations in Figure 19(c) and Figure 19(d). The rules in Figure 19(a) and Figure 19(c) have an impure lhs. If we apply the iARs in Figure 19(c) and Figure 19(d) to the data graph $G$ in Figure 4(a), both have the same frequency, namely 2, and the same confidence, namely 33%. Indeed, since the frequency of a tree query is in fact the frequency of its body, repetitions of distinguished variables in the head and the occurrence of parameters in the head do not change the frequency of a tree query. In fact the pAR in Figure 19(b) is the purification of the pAR in Figure 19(a): the repetition of the distinguished variable $x_2$ is removed from the heads, and the parameter $\sigma_1$ is removed from the heads.

Hence, a pAR with an impure lhs can always be rewritten to an equivalent pAR with a pure lhs, in such a way that all instantiations of the pAR with the impure lhs correspond to instantiations of the pAR with pure lhs, with the same confidence and frequency. Indeed, take a legal pAR $Q_1 \Rightarrow_\rho Q_2$ with $Q_1$ not pure. We know that $Q_1$’s head is mapped to $Q_2$’s head by some $\rho$-containment mapping. Hence, we can purify $Q_1$ by removing all parameters and repetitions of distinguished variables from $Q_1$’s head, sort the head by the order on the variables, and perform the corresponding actions on $Q_2$’s head as prescribed by the $\rho$-containment mapping.

We can conclude that it is sufficient to only consider pARs with pure lhs’s. The rhs, however, need not be pure; impure rhs’s are in fact interesting, as we will demonstrate next.

**Impure rhs’s** Consider the pAR in Figure 20(a). The rhs is impure since $x_2$ appears twice in the head. The pAR expresses that a sufficient proportion of the matchings of the lhs pattern, are also matchings of the rhs pattern, which
Figure 20: (a) and (b) are pARs with impure rhs. (c) is an ill-advised attempt to purify (b) on the rhs.

Figure 21: A pAR with a pure rhs.

is the same as the lhs pattern except that \( x_2 \) is equal to \( x_3 \). Since the pAR has no parameters, we can identify it with its instantiation by the empty parameter assignment. The confidence is then:

\[
\frac{m}{\sum_x \text{deg}^2 x}
\]

where \( m \) is the number of edges, \( x \) ranges over the nodes in the graph, and \( \text{deg} x \) is the outdegree of (number of edges leaving) \( x \). Since \( m = \sum_x \text{deg} x \), we show by an easy calculation that this confidence is much larger than \( 1/m \):

\[
\frac{m}{\sum_x \text{deg}^2 x} = \frac{\sum_x \text{deg} x}{\sum_x \text{deg}^2 x} \\
\geq \frac{\sum_x \text{deg} x}{(\sum_x \text{deg} x)^2} \\
= \frac{1}{\sum_x \text{deg} x} \\
= \frac{1}{m}
\]

Hence, the sparser the graph (with the number of nodes remaining the same), the higher the confidence, and thus the pAR is interesting in that it tells us something about the sparsity of the graph. As an illustration, on the graph of Figure 4(a) the confidence is 0.4, but on the the graph of Figure 4(b) it is 0.6.
Also consider the pAR in Figure 20(b). Again the rhs is impure since its head contains a parameter. Create an iAR for this pAR with $\alpha = \sigma_2 \mapsto 8$. With confidence $c$, this IAR then expresses that a fraction of $c$ of all edges point to node 8, which again would be an interesting property of the graph.

The knowledge expressed by the above two example pARs cannot be expressed using pARs with pure rhs’s. To illustrate, the pAR of Figure 20(c) may at first seem equivalent (and has a pure rhs) to that of Figure 20(b). On second thought, however, it says nothing about the proportion of edges pointing to $\sigma_2$, but only about the proportion of nodes with an edge to $\sigma_2$.

Of course, we are not implying that pARs with pure rhs’s are uninteresting. But all they can express are statements about the proportion of matchings of the lhs that can be specialized or extended to a matching of the rhs (another example is in Figure 21, which says something about the proportion of edges that can be extended); they cannot say anything about the proportion of matchings of the lhs that satisfy certain equalities in the distinguished variables.

**Free Constants** Most treatments of conjunctive database queries [9, 2, 40] allow arbitrary constants in the head. In our treatment, a constant can only appear in the head as the value of a parameter. Fortunately this is enough. We do not need to consider “free” constants, i.e., constants not corresponding to a parameter value. To see this, first consider the possibility of free constants in the lhs. The same argument we already gave to assume that the lhs is pure can be used to dismiss this possibility. Next consider a constant in the rhs of an iAR $(Q_1 \Rightarrow \rho, Q_2, \alpha)$, with $Q_1 = (H_1, P_1)$ and $Q_2 = (H_2, P_2)$ and $Q_1$ already pure. Then there must be a $\rho$-containment mapping $f : Q_1 \to Q_2$, with $f(H_1) = H_2$, for the iAR to be legal. Hence, a constant $a$ can only appear in $H_2$ by one of the following two possibilities:

1. $a = \alpha(f(\sigma)) = \alpha(\rho(\sigma))$, with $\sigma \in H_1$; or
2. $a = \alpha(f(x))$, with $x$ a distinguished variable in $H_1$.

However, in both cases $a$ is not actually free, being equal to a parameter value.

**5.2 Overall Approach**

Given the inputs: $G$; $Q_{left} = (H_{left}, P_{left})$; minconf; and minsup, an outline of our algorithm for the association rule mining problem is that of four nested loops:

1. Generate, incrementally, all possible trees of increasing sizes. Avoid trees that are isomorphic to previously generated ones. The height of the generated trees must be at least the height of the tree underlying $P_{left}$. (When enough trees have been generated, this loop can be terminated.)

2. For each new generated tree $T$, generate all frequent instantiated tree patterns $P^\alpha$ based on that tree.

These first two loops are nothing but our algorithm for mining frequent tree queries as presented in Section 4.
3. For each parameterized tree pattern $P$, generate all containment mappings $f$ from $P_{\text{left}}$ to $P$. Here, a plain “containment mapping” is a $\rho$-containment mapping, as defined in Section 3.3, for some $\rho$. Note that $\rho$ then equals $f|_{\Sigma_{\text{left}}}$. 

4. For each $f$, generate the parameterized tree query $Q_{\text{right}} = (f(H_{\text{left}}), P)$, and all parameter assignments $\alpha$ such that $(Q_{\text{left}} \Rightarrow_{\rho} Q_{\text{right}}, \alpha)$ is frequent; and the confidence exceeds $\minconf$. The generation of all these $\alpha$’s happens in a parallel fashion.

This approach is complete, i.e., it will output everything that must be output. In proof, consider a legal, frequent and confident iAR $(Q_{\text{left}} \Rightarrow_{\rho_0} Q_{\text{right}}, \alpha_0)$, with $Q_{\text{right}} = (H_{\text{right}}, P_{\text{right}})$. The tree $T$ is the underlying tree of $P_{\text{right}}$; $P_{\text{right}}$ is a tree pattern $P$ in loop 2; the containment mapping $f$ in loop 3 is the $\rho_0$-containment mapping that exists since the iAR is legal; $H_{\text{right}}$ is $f(H_{\text{left}})$; and $\alpha$ in loop 4 is $\alpha_0$.

The reader may wonder whether loop 3 cannot be organized in a levelwise fashion. This is not obvious, however, since any two queries of the form $((f_1(H_{\text{left}}), P), \alpha)$ and $((f_2(H_{\text{left}}), P), \alpha)$ have exactly the same frequency, namely that of $P^\alpha$. Loop 4, however, is levelwise because it is based on loop 2 which is levelwise.

As already mentioned, these first two loops are nothing but our algorithm for mining frequent tree queries as presented in Section 4. As already explained in Section 4.6 in loop 2 we build up a structured database containing all frequency tables for all trees in loop 1. We call this database the pattern database. In fact these two loops should be regarded as a preprocessing step; once built up, this pattern database can be used to generate association rules.

Hence, in practice an outline for our rule-mining algorithm is the following:

1. Preprocessing step: Generate a pattern database $D$ using the algorithm discussed in Section 4. Halt this algorithm when enough patterns are generated.

2. Consider, in a levelwise order, each parameterized tree pattern $P$ that has frequent instantiations in $D$, and such that the height of the underlying tree of $P$ is at least the height of the underlying tree of $P_{\text{left}}$.

3. For each parameterized tree pattern $P$, generate all containment mappings $f$ from $P_{\text{left}}$ to $P$ and let $\rho$ be $f|_{\Sigma_{\text{left}}}$.

4. For each $f$, generate the parameterized tree query $Q = (f(H_{\text{left}}), P)$, and all parameter assignments $\alpha$ such that $(Q_{\text{left}} \Rightarrow_{\rho} Q, \alpha)$ is frequent; and the confidence exceeds $\minconf$. The generation of all these $\alpha$’s happens in a parallel fashion.

We present loops 3 and 4 in detail in Sections 5.3 and 5.4. In Section 5.6 we will show how our overall approach must be refined so that the generation of equivalent association rules is avoided.

### 5.3 Generation of Containment Mappings

In this section, we discuss loop 3, the generation of all containment mappings $f$ from $P_{\text{left}}$ to $P$. So, we need to solve the following problem: Given two
parameterized tree patterns $P_1$ and $P_2$, find all containment mappings $f$ from $P_1$ to $P_2$.

Since the patterns are typically small, a naive algorithm suffices. For a node $x_1$ of $P_1$ and a node $x_2$ of $P_2$, we say that $x_1$ “matches” $x_2$ if there is a containment mapping $f$ from the subpattern of $P_1$ rooted at $x_1$ to the subpattern of $P_2$ rooted at $x_2$ such that $f(x_1) = x_2$. In a first phase, we determine for every node $y$ of $P_2$ separately whether the root $r_1$ of $P_1$ matches $y$. While doing so, we also determine for every other node $x_1$ of $P_1$, and every node $x_2$ below $y$ at the same distance as $x_1$ is from $r_1$, whether $x_1$ matches $x_2$. We store all these boolean values in a two-dimensional matrix $Map$. The function for filling in $Map$ is given in function 4. In line 2 of this function we mean by “$x_1 \rightarrow x_2$ is legal”, that if $x_1$ is a distinguished variable, then $x_2$ is a distinguished variable or a parameter; and if $x_1$ is a parameter then $x_2$ is a parameter, as prescribed by the definition of a $\rho$-containment mapping in Section 3.3.

Function 4 Function for filling in Map

1: bool FillIn($x_1 \in P_1$, $x_2 \in P_2$)
2: if $x_1 \rightarrow x_2$ is legal then
3: Match := true;
4: for each child $c_1$ of $x_1$ from left to right do
5: MatchChild := false;
6: for each child $c_2$ of $x_2$ from left to right do
7: MatchChild := MatchChild OR FillIn($c_1$, $c_2$)
8: end for
9: Match := Match AND MatchChild;
10: end for
11: Map[$x_1$, $x_2$] := Match;
12: return Match;
13: else
14: Map[$x_1$, $x_2$] := false;
15: return false
16: end if

This first phase compares every possible pair $(x_1, x_2)$, with $x_1$ a node in $P_1$ and $x_2$ a node in $P_2$, at most once. Indeed, if $x_1$ is at distance $d$ from $r_1$, then $x_1$ will be compared to $x_2$ only during the matching of $r_1$ with the node $y$ that is $d$ steps above $x_2$ in $P_2$ (if existing). We thus have an $O(n_1 \times n_2)$ algorithm, where $n_1$ ($n_2$) is the number of nodes in $P_1$ ($P_2$).

In a second phase, we output all containment mappings. Initially, by a synchronous preorder traversal of $P_1$ and $P_2$, we map each node of $P_1$ to the first matching node of $P_2$. We store this first mapping in a one-dimensional matrix $C_m$. In function 5 an outline for finding the initial containment mapping is given.

In each subsequent step, we look for the last node $x_1$ (in preorder) of $P_1$, currently matched to some node $x_2$, with the property that $x_1$ can also be matched to a right sibling $x_3$ of $x_2$, and now map $x_1$ to the first such $x_3$. The mappings of all nodes of $P_1$ coming after $x_1$ are reinitialized. Every such step takes time that is linear in $n_1$ and $n_2$. Of course, the total number of different containment mappings may well be exponential in $n_1$. An outline of this step is given in Function 6.
Function 5 Function for finding the initial containment mapping

1: $\text{Init}(x_1 \in P_1, x_2 \in P_2)$
2: $\text{Cm}[x_1] := x_2$;
3: for each child $c_1$ of $x_1$ from left to right do
4: for each child $c_2$ of $x_2$ from left to right do
5: if $\text{Map}[c_1, c_2]$ then
6: $\text{Init}(c_1, c_2)$;
7: Break;
8: end if
9: end for
10: end for

Function 6 Function for finding the other containment mappings

1: $\text{bool } \text{Step}(x \in P_1)$
2: $\text{Found} := \text{false}$;
3: for each child $c$ from $x$ from right to left do
4: if $\text{Step}(c)$ then
5: $\text{Found} := \text{true}$;
6: Break;
7: end if
8: end for
9: if $\text{Found}$ then
10: for each right-sibling $z$ of $c$ from left to right do
11: $p_2 := \text{Cm}[x]$;
12: for each child $c_2$ of $p_2$ from left to right do
13: if $\text{Map}[z, c_2]$ then
14: $\text{Init}(z, c_2)$
15: end if
16: end for
17: end for
18: return true;
19: else
20: if $x$ is the root of $P_1$ then
21: return false;
22: else
23: $m := \text{Cm}[x]$;
24: for each right-sibling $s$ of $m$ from left to right do
25: if $\text{Map}[x, s]$ then
26: $\text{Init}(x, s)$
27: Break;
28: end if
29: end for
30: return true;
31: end if
32: end if
The complete outline for the generation of all containment mappings is given in Function 7.

**Function 7** Function for generating all containment mappings from \( P_1 \) to \( P_2 \)

1: \( \text{GenerateCm}(P_1, P_2) \)
2: Initialize Map;
3: \( r_1 := \text{root of } P_1 \);
4: \text{for each } \( x_2 \in P_2 \) in preorder \text{ do}
5: \( \text{FillIn}(r_1, x_2) \);
6: \text{end for}
7: \text{for each node } \( x_2 \in P_2 \) in preorder \text{ do}
8: \text{if } \text{Map}[r_1, x_2] \text{ then}
9: \( \text{Initialize Cm} \);
10: \( \text{Init}(r_1, x_2) \);
11: \text{repeat}
12: \text{Output Cm} \text{; }
13: \text{until not Step}(r_1) \text{; }
14: \text{end if}
15: \text{end for}

We can thus easily generate all containment mappings \( f \) from \( P_{\text{left}} \) to \( P \) as required for loop 3 of our overall algorithm. Note, however, that in loop 4 these mappings are used to produce the head \( f(H_{\text{left}}) \) of query \( Q_{\text{right}} \). For \( Q_{\text{right}} \) to be a legal query, this head must contain all distinguished variables of \( P \). Hence, we only pass to loop 4 those \( f \) whose image contains all distinguished variables of \( P \).

### 5.4 Generation of Parameter Assignments

In loop 4, our task is the following. Given a containment mapping \( f : P_{\text{left}} \rightarrow P \), let \( \rho = f|_{\Sigma_{\text{left}}} \), and generate all parameter assignments \( \alpha \) such that \((Q_{\text{left}} \Rightarrow \rho (f(H_{\text{left}}), P), \alpha)\) is frequent and confident in \( G \). We show how this can be done in a parallel database-oriented fashion.

Recall from Section 4.6 that the frequency tables for \( P_{\text{left}} \) and \( P \) are available in a relational database. Our crucial observation is that we can compute precisely the required set of parameter assignments \( \alpha \), together with the frequency and confidence of the corresponding association rules, by a single relational algebra expression. This expression has the following form:

\[
\pi_{\text{plist}} \sigma \left( \text{FreqTab}_{P_{\text{left}}}.\text{freq} \geq \minconf \right) \left( \text{FreqTab}_{P}.\text{freq} \right)
\]

Here, \( \pi \) denotes projection, \( \sigma \) denotes selection, and \( \bowtie \) denotes join. The join condition \( \theta \) and the projection list \( \text{plist} \) are defined as follows. For \( \theta \), we take the conjunction:

\[
\bigwedge_{\sigma \in \Sigma_{\text{left}}} \text{FreqTab}_{P_{\text{left}}}.\sigma = \text{FreqTab}_{P}.\rho(\sigma)
\]

Furthermore, \( \text{plist} \) consists of all attributes \( P_{\text{left}}.\sigma_{\text{left}} \), with \( \sigma_{\text{left}} \in \Sigma_{\text{left}} \); all attributes \( P.\sigma \), with \( \sigma \in \Sigma \); together with the attributes \( \text{FreqTab}_{P}.\text{freq} \) and \( \text{FreqTab}_{P_{\text{left}}}.\text{freq} \).
Referring back to our overall algorithm (Section 5.2), we thus generate, for each pattern $P$ from loop 2 and each containment mapping $f$ in loop 3, all association rules with the given $Q_{\text{left}}$ as lhs in parallel, by one relational database query (which can be implemented by a simple SQL select-statement).

**Example.** Consider $Q_{\text{left}}$ and $P = (\Pi, \Sigma)$ as shown in Figures 22(a) and 22(b). We have $\Sigma_{\text{left}} = \{\sigma_1, \sigma_4\}$ and $\Pi_{\text{left}} = \{x_3, x_6\}$, and $\Sigma = \{\sigma_1, \sigma_4, \sigma_5\}$ and $\Pi = \{x_3\}$. Take the following containment mapping $f$ from $P_{\text{left}}$ to $P$:

| $f$ |
|-----|
| $\sigma_1$ | $\sigma_1$ |
| $x_1$ | $x_2$ |
| $\exists_3$ | $\exists_3$ |
| $\sigma_4$ | $\sigma_4$ |
| $x_5$ | $x_6$ |
| $\exists_6$ | $\exists_6$ |
| $x_7$ | $\sigma_4$ |

Then the rhs query $Q_{\text{right}}$ equals ($(x_2, x_2, \sigma_4), P$), and the relational algebra expression for computing all parameter assignments and their corresponding frequencies and confidences looks as follows:

$$
\pi_{\text{plist}} \sigma_{\text{FreqTab}_{\text{P}}.\text{freq} \geq \text{minconf}} (\text{FreqTab}_{\text{P}} \bowtie_0 \text{FreqTab}_{P})
$$

with $\text{plist}$ equal to

$$
\text{FreqTab}_{\text{P}}.\sigma_1, \text{FreqTab}_{\text{P}}.\sigma_4, \text{FreqTab}_{P}.\sigma_1, \text{FreqTab}_{P}.\sigma_4, \text{FreqTab}_{P}.\sigma_5,
\text{FreqTab}_{P}.\text{freq}, \text{FreqTab}_{P}.\text{freq}/\text{FreqTab}_{\text{P}}.\text{freq}
$$

and $\theta$ equal to

$$
\text{FreqTab}_{P}.\sigma_1 = \text{FreqTab}_{\text{P}}.\sigma_1 \land \text{FreqTab}_{P}.\sigma_4 = \text{FreqTab}_{\text{P}}.\sigma_4
$$

In SQL, we get:

```sql
SELECT freqQleft.x1, freqQleft.x4, freqP.x1, freqP.x4, freqP.x5, freqP.freq, freqP.freq/freqQleft.freq
FROM freqP, freqQleft
WHERE freqQleft.x1 = freqP.x1 AND freqQleft.x4 = freqP.x4
AND freqP.freq/freqQleft.freq >= minconf
```

### 5.5 Example Run

In this Section we give an example run of the algorithm discussed in Section 5.3. We use the same data graph $G$, unordered rooted tree $T$, and minimum support threshold, 3, as in the example run in Section 4.5.4. The fixed lhs tree query is given in Figure 23(a), its corresponding frequency table in Figure 23(b) and the minimum confidence threshold is 30%. All frequent tree patterns based on $T$ were already generated in the example run of Section 4.5.3.

The example run then looks as follows:
Figure 22: Example $Q_{\text{left}}$ and $P$.

Figure 23: The fixed lhs and its frequency table for the example run in Section 5.5.
| $P$  | Containment Mapping | $Q_{\text{right}}$ | $ConfTab$ |
|------|---------------------|-------------------|-----------|
|      |                     |                   |           |
| **Level 0** |                      |                   |           |
| $(\emptyset, \emptyset)$ | No Containment Mappings |                   |           |
| **Level 1** |                      |                   |           |
| $(\emptyset, \{x_1\})$ | No Containment Mappings |                   |           |
| $(\emptyset, \{x_1\})$ | ![Diagram](#) | $(x_1, x_3, x_3)$ | P_{\text{left}} \cdot \sigma_2 | P \cdot \sigma_2 | Freq | Conf |
|      |                     |                   | 1 | 1 | 3 | 33% |
|      |                     |                   | 2 | 2 | 3 | 33% |
|      |                     |                   | 3 | 3 | 3 | 33% |
|      |                     |                   | 4 | 4 | 3 | 60% |
| $(\emptyset, \{x_2\})$ | ![Diagram](#) | $(x_1, \sigma_2, x_3)$ | P_{\text{left}} \cdot \sigma_2 | P \cdot \sigma_2 | Freq | Conf |
|      |                     |                   | 1 | 1 | 3 | 33% |
|      |                     |                   | 2 | 2 | 3 | 33% |
|      |                     |                   | 3 | 3 | 3 | 33% |
|      |                     |                   | 4 | 4 | 3 | 60% |
| $(\emptyset, \{x_2\})$ | ![Diagram](#) | $(x_1, x_3, \sigma_2)$ | P_{\text{left}} \cdot \sigma_2 | P \cdot \sigma_2 | Freq | Conf |
|      |                     |                   | 1 | 1 | 3 | 33% |
|      |                     |                   | 2 | 2 | 3 | 33% |
|      |                     |                   | 3 | 3 | 3 | 33% |
|      |                     |                   | 4 | 4 | 3 | 60% |
| $(\{x_1\}, \emptyset)$ | No containment mappings |                   |           |
| **Level 2** |                      |                   |           |
\[
(x_1, x_2, x_3) \quad \frac{\sigma_1}{\sigma_2} \quad (\sigma_1, x_3, x_4) \quad (\sigma_1, x_3, x_4) \quad \frac{\sigma_1}{\sigma_2}
\]

| \( P_{\text{left}} \cdot \sigma_2 \) | \( P \cdot \sigma_1 \) | \( P \cdot \sigma_2 \) | \text{Freq} | \text{Conf} |
|---|---|---|---|---|
| 0 | 1 | 1 | 3 | 33% |
| 0 | 2 | 2 | 3 | 33% |
| 0 | 3 | 3 | 3 | 33% |

\[
(x_1, x_2, x_3) \quad \frac{\sigma_1}{\sigma_2} \quad (\sigma_1, x_3, x_4) \quad (\sigma_1, x_3, x_4) \quad \frac{\sigma_1}{\sigma_2}
\]

| \( P_{\text{left}} \cdot \sigma_2 \) | \( P \cdot \sigma_1 \) | \( P \cdot \sigma_2 \) | \text{Freq} | \text{Conf} |
|---|---|---|---|---|
| 0 | 1 | 1 | 3 | 33% |
| 0 | 2 | 2 | 3 | 33% |
| 0 | 3 | 3 | 3 | 33% |

\[
(x_1, x_2, x_3) \quad \frac{\sigma_1}{\sigma_2} \quad (\sigma_1, x_3, x_4) \quad (\sigma_1, x_3, x_4) \quad \frac{\sigma_1}{\sigma_2}
\]

| \( P_{\text{left}} \cdot \sigma_2 \) | \( P \cdot \sigma_1 \) | \( P \cdot \sigma_2 \) | \text{Freq} | \text{Conf} |
|---|---|---|---|---|
| 0 | 1 | 1 | 3 | 33% |
| 0 | 2 | 2 | 3 | 33% |
| 0 | 3 | 3 | 3 | 33% |

\[
(x_1, x_2) \quad \text{No containment mappings}
\]

**Level 3**

\[
(x_1, x_2, x_3) \quad \text{No containment mappings}
\]
5.6 Equivalent Association Rules

In this section, we make a number of modifications to the algorithm described so far, so as to avoid duplicate work on equivalent rules.

Let us first look at an example of the duplicate work that the algorithm presented until now performs. Consider \( Q_{\text{left}} \), \( Q_1 = (f_1(H_{\text{left}}), P) \), \( Q_2 = (f_2(H_{\text{left}}), P) \); and \( Q_3 = (f_3(H_{\text{left}}), P) \) in Figure 24 with \( f_1 \), \( f_2 \) and \( f_3 \) as follows:

\[
\begin{array}{c|c}
\text{f}_1 & \text{f}_2 & \text{f}_3 \\
\hline
x_1 & u_1 & x_1 & u_1 & x_1 & u_1 \\
x_2 & u_2 & x_2 & u_2 & x_2 & u_2 \\
x_3 & u_2 & x_3 & u_2 & x_3 & u_3 \\
x_4 & u_3 & x_4 & u_2 & x_4 & u_3 \\
\end{array}
\]

Furthermore, consider pAR1: \( Q_{\text{left}} \Rightarrow Q_1 \); pAR2: \( Q_{\text{left}} \Rightarrow Q_2 \) and pAR3: \( Q_{\text{left}} \Rightarrow Q_3 \).

The confidence of the first rule (pAR1) equals the proportion of tuples from the answer set of \( Q_{\text{left}} \) where the values for variables \( x_2 \) and \( x_3 \) are equal (in the rhs those equal variables are represented by variable \( u_2 \), and the lhs variable \( x_4 \) is represented by the rhs variable \( u_3 \)). Similarly, the confidence of the second rule (pAR2) equals the proportion of tuples from the answer set of \( Q_{\text{left}} \) where the values for the variables \( x_3 \) and \( x_4 \) are equal (again the equal lhs variables \( x_3 \) and \( x_4 \) are represented by the rhs variable \( u_2 \), and the lhs variable \( x_2 \) is represented by the rhs variable \( u_3 \)). Since, due to the symmetry in the lhs pattern, the columns for \( x_2 \), \( x_3 \) and \( x_4 \) are fully interchangeable in the answer set of \( Q_{\text{left}} \), both rules convey precisely the same information: their confidences are equal. The third rule (pAR3) is yet another representation of the same association, but now the equal lhs variables \( x_3 \) and \( x_4 \) are represented by the rhs variable \( u_3 \). Again, it has the same confidence as pAR1 and pAR2.

It is important to note that the above pARs only differ in the containment mappings \( f_1 \), \( f_2 \) and \( f_3 \) that generate the rhs head. The algorithm discussed until now generates all these pARs, since we do not perform any check on the containment mappings generated in loop 3 of the overall approach (Section 5.2).

In this Subsection, motivated by the above example, we consider the general problem of when two pARs \( Q_{\text{left}} \Rightarrow \rho_1 \) \( Q_1 \) and \( Q_{\text{left}} \Rightarrow \rho_2 \) \( Q_2 \) are equivalent, where \( Q_1 \) and \( Q_2 \) are of the form \((f_1(H_{\text{left}}), P)\) and \((f_2(H_{\text{left}}), P)\) for some common rhs pattern \( P \), and containment mappings \( f_1 \) and \( f_2 \) from \( P_{\text{left}} \) to \( P \). (Thus \( \rho_1 \) is \( f_1|_{\Sigma_{\text{left}}} \) and \( \rho_2 \) is \( f_2|_{\Sigma_{\text{left}}} \)). Since such two pARs differ only in \( f_1 \) and \( f_2 \) we can actually focus on \( f_1 \) and \( f_2 \).

It is important to remember for the rest of this Subsection that \( P_{\text{left}} \) and \( P \) are arbitrary but fixed. Furthermore, without loss of generality we assume that the nodes of \( P_{\text{left}} \) and \( P \) are disjoint. This assumption greatly simplifies the representation of containment mappings by graphs, as we will see shortly.

**Equivalent Containment Mappings** Recall from Section 4.5.3 that an isomorphism from a parameterized tree pattern \( P_1 \) to a parameterized tree pattern \( P_2 \) is a homomorphism from \( P_1 \) to \( P_2 \) that is a bijection and that maps distinguished nodes to distinguished nodes, parameters to parameters and existential nodes to existential nodes. We now formalize equivalent containment mappings as follows: Two containment mappings \( f_1 \) and \( f_2 \) are equivalent if the structures
Figure 24: Queries to illustrate the duplicate work in the association mining algorithm

Figure 25: The graph representations of $f_1$ and $f_3$.

$(P_{\text{left}}, P, f_1)$ and $(P_{\text{left}}, P, f_2)$ are isomorphic. Specifically, there must exist isomorphisms (actually automorphisms) $g : P_{\text{left}} \to P_{\text{left}}$ and $h : P \to P$ such that $f_2 \circ g = h \circ f_1$.

Consider for instance $f_1$ and $f_3$ from the example above, then $h$ swaps $u_2$ and $u_3$, and $g$ is the cyclic permutation $u_2 \mapsto u_3 \mapsto u_4 \mapsto u_2$.

5.6.1 Testing for equivalence

To test for equivalent containment mappings efficiently, we represent them using graphs.

**Graph representation of a containment mapping** The graph representation of a containment mapping $f : P_{\text{left}} \to P$ is a directed, edge- and vertex-colored graph, with set of vertices $V_f = \text{Vertices}(P_{\text{left}}) \cup \text{Vertices}(P)$ and set of
edges $E_f = \text{Edges}(P_{\text{left}}) \cup \text{Edges}(P) \cup \{(v, w) \mid f(v) = w\}$ (with the understanding that the edges of $P_{\text{left}}$ and $P$ go from parent to child). We use different colors for the edges of $P_{\text{left}}$, the edges of $P$ and the pairs in $f$, and we also use different colors for the distinguished nodes, the existential nodes and the parameters.

As an illustration, Figure $25$ shows the graph representation of $f_1$ and $f_3$ from our example in the introduction above.

**Graph Isomorphism** Two graphs $G_1 = (V_1, E_1)$ and $G_2 = (V_2, E_2)$ are *colored isomorphic* if there exists a bijection $\varphi : V_1 \to V_2$, extended to edges $(v, w) \in E_1$ in a natural way by $\varphi(v, w) = (\varphi(v), \varphi(w))$, such that the colors of vertices and edges are preserved by $\varphi$, and such that $(v, w) \in E_1 \iff (\varphi(v), \varphi(w)) \in E_2$.

The following Lemma shows then the utility of the colored graph representation of containment mappings.

**Lemma 6.** Two containment mappings are equivalent if and only if their colored graph representations are isomorphic.

**Proof.** Let us start with the only-if direction. Consider two equivalent containment mappings $f_1, f_2$ from $P_{\text{left}}$ to $P$. By definition of equivalent containment mappings, there exist isomorphisms $g : P_{\text{left}} \to P_{\text{left}}$ and $h : P \to P$ such that $f_2 \circ g = h \circ f_1$. Now take $\varphi = g \cup h$. Then, $\varphi$ is clearly a bijection from $V_{f_1}$ to $V_{f_2}$, and clearly preserves the colors of vertices and edges of $G_{f_1}$. Let $(v, w) \in E_{f_1}$. We show that $\varphi$ is indeed an isomorphism from $G_{f_1}$ to $G_{f_2}$. There are three possibilities:

1. $(v, w) \in \text{Edges}(P_{\text{left}})$. Note that then also $g(v, w) \in \text{Edges}(P_{\text{left}})$. We have:

   
   \[
   (v, w) \in E_{f_1} \iff (v, w) \in \text{Edges}(P_{\text{left}})
   \]

   $g$ is an automorphism in $P_{\text{left}}$

   \[
   \iff g(v, w) \in \text{Edges}(P_{\text{left}})
   \]

   $\varphi(v, w) = g(v, w)$

   \[
   \iff g(v, w) \in E_{f_2}
   \]

   $\varphi(v, w) \in E_{f_2}$

   \[
   \iff \varphi(v, w) \in E_{f_2}
   \]

2. $(v, w) \in \text{Edges}(P)$. Note that then also $h(v, w) \in \text{Edges}(P)$. We have:

   
   \[
   (v, w) \in E_{f_1} \iff (v, w) \in \text{Edges}(P)
   \]

   $h$ is an automorphism in $P$

   \[
   \iff g(v, w) \in \text{Edges}(P)
   \]

   $\varphi(v, w) = h(v, w)$

   \[
   \iff h(v, w) \in E_{f_2}
   \]

   $\varphi(v, w) \in E_{f_2}$

   \[
   \iff \varphi(v, w) \in E_{f_2}
   \]

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3. $w = f_1(v)$. We have:

$$(v, w) \in E_{f_1} \iff v = f_1(w)$$
$$\iff h(v) = h(f_1(w))$$
$$\iff h(v) = f_2(g(w))$$
$$\iff \varphi(v) = f_2(\varphi(w))$$
$$\iff (\varphi(v), \varphi(w)) \in E_{f_2}$$

So we can conclude that $G_{f_1}$ and $G_{f_2}$ are indeed colored isomorphic.

Let us now look at the if-direction. Let $\varphi$ be the given isomorphism from $G_{f_1}$ to $G_{f_2}$. Now take $g = \varphi|_{\text{Vertices}(P_{\text{left}})}$ and $h = \varphi|_{\text{Vertices}(P)}$. To prove that $f_1$ and $f_2$ are equivalent it suffices to show that:

1. $g$ is an isomorphism from $P_{\text{left}}$ to $P_{\text{left}}$;
2. $h$ is an isomorphism from $P$ to $P$; and
3. $f_2 \circ g = h \circ f_1$.

Items 1 and 2 hold because $\varphi$ preserves the colors. For 3, let $v \in P_{\text{left}}$. Since $\varphi$ is a graph isomorphism $f_2(\varphi(v))$ equals $\varphi(f_1(v))$. We then have:

$$f_2(g(v)) = f_2(\varphi(v))$$
$$= \varphi(f_1(v))$$
$$= h(f_1)(v)$$

So, using graph isomorphism (to be precise, edge and vertex colored directed graph isomorphism), we can test for equivalence. Since our patterns are not very large, fast heuristics for graph isomorphism can be used. We use the program Nauty \[32, 33\], which is considered as the fastest heuristic for graph isomorphism. Nauty is very efficient for small, dense random graphs \[15\]. Since our graph representations are typically small (no more than 20 vertices) and dense, this works well in our case.

Theoretically this situation is not entirely satisfying, as graph isomorphism is not known to be efficiently (polynomial-time) solvable in general. We can show however that equivalence of our containment mappings is really as hard as the general graph isomorphism problem. This hardness argument is presented in the following Section 5.6.2. As special case of the equivalence problem that is solvable in polynomial time is presented in Section 5.6.3

### 5.6.2 Hardness argument

First recall from graph theory that a graph $B = (V, E)$ is bipartite if $V$ can be split in two disjoint parts, $V = V^a \cup V^b$ with $V^a \cap V^b = \emptyset$, such that for each $(v, w) \in E$ then $v \in V^a$ and $w \in V^b$. The vertices in $V^a$ are called lhs vertices and those in $V^b$ rhs vertices (left-hand side, right-hand side).

We first reduce the problem of bipartite graph isomorphism to equivalence of our containment mapping. Let $B_1 = (V_1, E_1)$ and $B_2 = (V_2, E_2)$ be bipartite graphs. We describe an efficient construction that produces from $B_1$ and $B_2$
two association rules \((P_{\text{left}}, P, f_1)\) and \((P_{\text{left}}, P, f_2)\) such that \(B_1\) and \(B_2\) are isomorphic if and only if the association rules are equivalent. This construction reduces the bipartite graph isomorphism problem to equivalence of containment mappings.

Without loss of generality, we assume that \(B_1\) and \(B_2\) have precisely the same multiset of outdegrees (for vertices of \(V_1^a\) and \(V_2^a\)), and precisely the same number of vertices in \(V_1^b\) and \(V_2^b\). Indeed, if these conditions are not satisfied, then \(B_1\) and \(B_2\) are never isomorphic and our reduction can output some arbitrary \(P_{\text{left}}, P, f_1\) and \(P_{\text{left}}, P, f_2\) as long as \((P_{\text{left}}, P, f_1)\) and \((P_{\text{left}}, P, f_2)\) are not equivalent.

The construction in now as follows. By the premisses on \(B_1\) and \(B_2\), we may assume, without loss of generality, that \(V_1^a = V_2^a\) and \(V_1^b = V_2^b\). This can be accomplished by sorting the lhs vertices in each graph on their outdegrees an then numbering them arbitrarily (the rhs vertices can simply be numbered arbitrarily).

1. **Construction of \(P_{\text{left}}\).** This is a tree with root called \(r_{\text{left}}\) and as children of the root, all lhs vertices. Moreover, each lhs vertex \(v\) has its own children as follows: if \(v\) has outdegree \(o\), then \(v\) has \(o\) children denoted by \([v, 1], [v, 2], \ldots, [v, o]\).

2. **Construction of \(P\).** This is a tree with root called \(r_{\text{right}}\), and exactly one child of the root, called \(c\). Moreover, \(c\) has as children precisely all rhs vertices.

3. **Construction of \(f_1\).** We define \(f(r_{\text{left}}) := r_{\text{right}}\), and define \(f_1(v) := c\) for each lhs vertex \(v\). Now for each such \(v\), and all outgoing edges \((v, w_1), (v, w_2), \ldots, (v, w_o)\) in \(B_1\), listed in some arbitrary order, we define \(f_1([v, i]) := w_i\), for \(i = 1, 2, \ldots, o\).

4. The construction of \(f_2\) is analogous to that of \(f_1\), but now we look at the outgoing edges in \(B_2\).

The construction is illustrated in Figure 26 for two bipartite graphs \(B_1\) and \(B_2\). We now show the correctness of our reduction.

**Lemma 7.** \(B_1\) and \(B_2\) are isomorphic if and only if \((P_{\text{left}}, P, f_1)\) and \((P_{\text{left}}, P, f_2)\) are isomorphic.

**Proof.** For the only-if direction, let \(\psi\) be an isomorphism from \(B_1\) to \(B_2\). We define an isomorphism from \((P_{\text{left}}, P, f_1)\) to \((P_{\text{left}}, P, f_2)\) as follows:

- \(\varphi(r_{\text{left}}) = r_{\text{left}}, \varphi(r_{\text{right}}) = r_{\text{right}}\) and \(\varphi(c) = c\);
- \(\varphi(v) = \psi(v)\), for any vertex of \(B_1\);
- for any lhs vertex \(v\) of outdegree \(o\), and any \(i = 1, 2, \ldots, o\), let \(w\) be the rhs vertex such that \(f_1([v, i]) = w\). Then we define \(\varphi([v, i]) := [\psi(v), j]\), where \(j\) is such that \(f_2([\psi(v), j]) = \psi(w)\).

To verify that \(\varphi\) is indeed an isomorphism, we only check that \(u = f_1([v, i]) \leftrightarrow \psi(u) = f_2(\psi([v, i]))\). If \(u = f_1([v, i])\) then \((v, u)\) is an edge in \(B_1\) and thus \((\varphi(v), \varphi(u)) = (\psi(v), \psi(w))\) is an edge in \(B_2\). Hence there exists a \(j\) such that,
Figure 26: Illustration of the construction of pARs from bipartite graphs.
ϕ(u) = f_2([ϕ(v), j]), or equivalent, ψ(u) = f_2([ψ(v), j]). By definition of ϕ we have ϕ([v, i]) = [ψ(v), j] and thus ϕ(u) = f_2(ϕ([v, i])) as desired. Conversely, suppose ϕ(u) = f_2(ϕ([v, i])). By definition of ϕ, we have ϕ([v, i]) equals [ψ(v), j] for some unique j, and f_2([ψ(v), j]) equals ψ(f_1([v, i])). Hence, ψ(u) = ϕ(u) = ψ(f_1([v, i])), and thus u = f_1([v, i]) as desired.

For the if-direction, let ϕ be an isomorphism from (P_{left}, P, f_1) to (P_{left}, P, f_2). We define an isomorphism ψ from B_1 to B_2 as follows. Actually, ψ is simple ϕ restricted to the vertices of B_1. Indeed,

\[(v, w) ∈ E_1 ⇔ ∃i : f_1([v, i]) = w\]
\[⇔ ∃i : f_2(ϕ([v, i])) = ϕ(w)\]
\[⇔ ∃j : f_2(ϕ(v), j) = ϕ(w)\]
\[⇔ (ϕ(v), ϕ(w)) ∈ E_2\]

We can already conclude from this reduction that equivalence of pARs is really as hard as isomorphism of bipartite directed graphs. The latter problem, however, is well-known to be as hard as isomorphism of general directed graphs. Indeed, any directed graph \(G = (V, E)\) can be transformed into the bipartite directed graph \(B(G) := (V ∪ E, \{(v, (v, w)) | (v, w) ∈ E \} ∪ \{(v, w), w) | (v, w) ∈ E\})\), and it is easily verified that \(G_1\) and \(G_2\) are isomorphic if and only if \(B(G_1)\) and \(B(G_2)\) are isomorphic.

So, we can now conclude that equivalence of our pARs is really as hard as the general graph isomorphism problem. But as we show next, we can still capture an important special case in polynomial time, so that the general graph isomorphism heuristics only have to be applied on instances not captured by the special case.

### 5.6.3 Polynomial case

The special efficient case is to check whether \((P_{left}, P, f_1)\) and \((P_{left}, P, f_2)\) are already isomorphic with \(g\) the identity, i.e., whether the structures \((P, f_1)\) and \((P, f_2)\) are already isomorphic. So, we look for an automorphism \(h\) of \(P\) such that \(f_2 = h ∘ f_1\). This can be solved efficiently by a reduction to node-labeled tree isomorphism. As explained in Section 5.4.3 if we know the tree \(T\) underlying \(P\), then \(P\) is characterized by the pair \((Π, Σ)\), and thus \((P, f)\) is characterized by \((Π, Σ, f)\). We can view this triple as a labeling of \(T\), as follows. We label every node \(y\) of \(P\) with a triple \((b_1, b_2, f^{-1}(y))\), where \(b_1\) is a bit that is 1 iff \(y ∈ Π\); \(b_2\) is a bit that is defined likewise; and \(f^{-1}(y)\) is the set of nodes of \(P_{left}\) that are mapped by \(f\) to \(y\). Then \((P, f_1)\) and \((P, f_2)\) are isomorphic if and only if the corresponding node-labeled trees are isomorphic, and the latter can be checked in linear time using canonical ordering [4, 10].

### 5.6.4 The Algorithm

We are now in a position to describe how our general algorithm must be modified to avoid equivalent association rules. There is only extra checking to be done in loop 3 (recall Sections 5.2 and 5.3). For each new containment mapping \(f\) from \(P_{left}\) to \(P\), we canonize the corresponding node-labeled tree and we check if
the canonical form is identical to an earlier generated canonical form; if so, \( f \) is dismissed. We can keep track of the canonical forms seen so far efficiently using a trie data structure. If the canonical form was not yet seen, we can either let \( f \) through to loop 4, if the presence of duplicates in the output is tolerable for the application at hand, or we can perform the colored graph isomorphism check of Section 5.6.2 with the containment mappings previously seen, to be absolutely sure we will not generate a duplicate.

6 Certhia: Pattern and Association Browsing

In this Section we introduce an interactive tool, called Certhia, for browsing the frequent tree patterns, and generating association rules.

As already noted in Section 4.6, the result of our tree query mining algorithm in Section 4 is a structured database, called a Pattern Database, containing all frequency tables for each tree \( T \) that was investigated. This pattern database is an ideal platform for an interactive tool for browsing the frequent queries. However, this pattern database is also an ideal platform for generating association rules as explained in Section 5.2, since the first two loops of association rule algorithm are exactly our tree query mining algorithm.

In a typical scenario for Certhia, the user draws a tree shape, marks some nodes as existential, marks some others as parameters, instantiates some parameters by constants, but possibly also leaves some parameters open. The browser then returns, by consulting the appropriate frequency table in the database, all instantiations of the free parameters that make the pattern frequent, together with the frequency. The user can then select one of these instantiations, set a minconf value, and ask the browser to return all rhs’s that form a confident association with the selected pure tree query as lhs. In another scenario the user lets the browser suggest some frequent tree patterns to choose from as an lhs.

Some screenshots of Certhia are given in Figure 27, Figure 28 and Figure 29.

- In Figure 27, the user draws a tree, marks some nodes as existential, some others as parameters, instantiates some parameters with constants, and asks the browser to return all possible instantiations of the remaining parameters and the corresponding frequencies.

- In Figure 28, the user asks the browser to return all association rules for a fixed lhs. The user selects a rhs in the dialog box and asks the browser to return the instantiations and the corresponding frequencies.

- In Figure 29, the browser suggests some frequent tree patterns where the user can choose from.

Efficiency The preprocessing step, i.e., the building up of the pattern database with frequent tree patterns, is of course a hugely intensively task. First because the large datagraph must be accessed intensively, and secondly because the number of frequent patterns is huge. In Section 7.2, we show that this preprocessing step can be implemented with satisfactory performance. Also, in scientific discovery applications it is no problem, indeed typical, if a preprocessing step takes
Figure 27: Screenshot of Certhia: the user draws a pattern and asks for the instantiations.

Figure 28: Screenshot of Certhia: the user asks for all association rules for this lhs.
Figure 29: Screenshot of Certhia: the browser suggests frequent tree patterns.

a few hours, as long as after that the interactive exploration of the found results can happen very fast. And indeed we found that the actual generation of association rules is very fast. This is also shown in Section 7.2.

7 Experimental Results

In this section, we report on some experiments performed using our prototype implementation applied to both real-life and synthetic datasets to show that our approach is indeed workable.

7.1 Real-life datasets

We have worked with a food web, a protein interactions graph, and a citation graph. For each dataset we built up a pattern database using the following parameters:

|           | #nodes | #edges | k | size |
|-----------|--------|--------|---|------|
| food web  | 154    | 370    | 25| 6    |
| proteins  | 2114   | 4480   | 10| 5    |
| citations | 2500   | 350000 | 5 | 4    |

As we set rather generous limits on the maximum size of trees, or on the minimum frequency threshold, each run took several hours.

The food web comprises 154 species that are all directly or indirectly dependent on the Scotch Broom (a kind of shrub). One of the patterns that was mined with frequency 176 is the following:
This is really a rather arbitrary example, just to give an idea of the kind of complex patterns that can be mined. Note also that, thanks to the constant 20 appearing twice, this is really a non-tree shaped pattern: we could equally well draw both arrows to a single node labeled 20.

While we were thus browsing through the results, we quickly noticed that the constant 20 actually occurs quite predominantly, in many different frequent patterns. This constant denotes the species *Orthotylus adenocarpi*, an omnivorous plant bug. To confirm our hypothesis that this species plays a central role in the food web, we asked for all association rules with the following left-hand side:

\[ (x_1, x_2) \quad \Rightarrow \quad (x_1, x_2) \]

Indeed, the rule shown above turned up with 89% confidence! For 89% of all pairs of species that are linked by a path of length four, *Orthotylus adenocarpi* is involved in between.

Two other rules we discovered are:

\[ (x_1, x_2, x_3, x_4, x_5) \quad \Rightarrow \quad (0, x_2, x_3, x_4, x_5) \]

\[ (x_1, x_2, x_3, x_4, x_5) \quad \Rightarrow \quad (0, x_2, x_3, x_4, x_5) \]
Since 45% + 55% = 100%, these rules together say that each path of length 5 either starts in 0, or one beneath 0. This tells us that the depth of the food web equals 6. Constant 0 turns out to denote the Scotch Broom itself, which is the root of the food web.

Another rule we mined, just to give a rather arbitrary example of the kind of rules we find with our algorithm, is the following:

\[
\begin{align*}
(x_1, x_2, x_3, x_4, x_5) & \quad \text{11%} \quad \Rightarrow \\
(x_1, x_2, x_4, x_2, x_5) & \quad 101
\end{align*}
\]

The protein interaction graph \[25\] comprises molecular interactions (symmetric) among 1870 proteins occurring in the yeast \textit{Saccharomyces cerevisiae}. In such interaction networks, typically a small number of highly connected nodes occur. Indeed, we discovered the following association rule with 10% confidence, indicating that protein #224 is highly connected:

\[
\begin{align*}
x_1 & \quad \exists \quad \Rightarrow \quad 224 \\
x_2 &
\end{align*}
\]

We also found the following rule:

\[
\begin{align*}
(x_1, x_2) & \quad 90\% \quad \Rightarrow \\
(x_1, x_2) & \quad 746 \quad 376
\end{align*}
\]

This rule expresses that almost all interactions that link to protein 746 also link to protein 376, which unveils a close relationship between these two proteins.

The citation graph comes from the KDD cup 2003, and contains around 2500 papers about high-energy physics taken from arXiv.org, with around 350 000 cross-references. One of the discovered patterns is the following, with frequency 1655, showing two papers that are frequently cited together (by 6% of all papers).

\[
\begin{align*}
x_1 & \quad 9711200 \quad 9802150
\end{align*}
\]

One of the discovered rules is the following:

\[
\begin{align*}
(x_1, x_2) & \quad 15\% \quad \Rightarrow \\
(x_1, x_2) & \quad 9503124
\end{align*}
\]
7.2 Performance

While our prototype implementations are not tuned for performance, we still conducted some preliminary performance measurements, with encouraging results. The experiments were performed on a Pentium IV (2.8GHz) architecture with 1GB of internal memory, running under Linux 2.6. The program was written in C++ with embedded SQL, with DB2 UDB v8.2 as the relational database system.

We have used two types of synthetic datasets.

Random Web graphs Naturally occurring graphs (as found in biology, sociology, or the WWW) have a number of typical characteristics, such as sparseness and a skewed degree distribution [55]. Various random graph models have been proposed in this respect, of which we have used the “copy model” for Web graphs [27, 6]. We use degree 5 and probability $\alpha = 10\%$ to link to a random node (thus 90\% to copy a link).

On these graphs, we have measured the total running time of the tree query mining algorithm as a function of the size (number of edges) of the graph, where we mine up to tree size 5, with varying minimum frequency thresholds of 4, 10, and 25. The results, depicted in Figure 30, show that the performance of these runs is quite adequate.
Figure 31: Performance in terms of number of discovered patterns.

**Uniform random graphs** We have also experimented with the well-known Erdős–Rényi random graphs, where one specifies a number $n$ of nodes and gives each of the possible $n^2$ edges a uniform probability (we used 10%) of actually belonging to the graph. In contrast to random Web graphs, these graphs are quite dense and uniform, and they serve well as a worst-case scenario to measure the performance of the tree query mining algorithm as a function of the number of discovered patterns, which will be huge.

We have run on graphs with 47, 264, and 997 edges, with minimum frequency thresholds of 10 and 25. The results, depicted in Figure 31, show, first, that huge numbers of patterns are mined within a reasonable time, and second, that the overhead per discovered pattern is constant (all six lines have the same slope).

On these uniform random graphs we also conducted some experiments to check the performance of the association rule mining algorithm. We found the actual generation of association rules (i.e., loops 3 and 4, assuming that a pattern database is already build up) to be very fast. For instance, Figure 32 shows the performance of generating association rules for two different (absolute) values of minconf, against a frequency table database built up for a random graph with 33 nodes and 113 edges, an absolute minsup of 25, and all trees up to size 7. We see that associations are generated with constant overhead, i.e., in linear-output time. The coefficient is larger for the larger minconf, because in this experiment we have counted instantiated rhs's, and per rhs query less instantiations satisfy the confidence threshold for larger such thresholds. Had we simply counted rhs's regardless of the number of confident instantiations, the two lines would have had the same slope.
**Performance issues** One major performance issue that we have not addressed in the present study is that some of the SQL queries that are performed due to pattern generation take a very long time (in order of hours) to answer by the database system. This happens in those cases where the data graph is large (5000 edges or more) with many cycles, and the candidate patterns are large (6 nodes are more). Certainly, some SQL queries can be hand-optimized (or replaced by a combination of simpler queries), to alleviate these performance problems, but we leave this issue to future research.

**Acknowledgment**

We thank Bart Goethals for his contributions in the initial conception of the ideas presented in this work. We also thank Jan Hidders and Dries Van Dyck for their help with our results on graph isomorphism.

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### Appendix

| Notation     | Interpretation                                                                 |
|--------------|-----------------------------------------------------------------------------|
| $U$          | set of data constants                                                       |
| $T$          | ordered rooted tree                                                          |
| $G$          | data graph                                                                  |
| $P$          | parameterized tree pattern                                                   |
| $\Pi$        | set of existential nodes                                                    |
| $\Delta$     | set of distinguished nodes                                                   |
| $\Sigma$     | set of parameters                                                           |
| $\exists$    | existential node                                                            |
| $\sigma$     | parameter                                                                    |
| $x$          | distinguished node                                                           |
| $\alpha$     | parameter assignment                                                        |
| $P^\alpha, (P, \alpha)$ | instantiated tree pattern                                                 |
| $P^\alpha(G)$ | $\{ \mu|_\Delta : \mu$ is a matching of $P^\alpha$ in $G \}$          |
| minsup       | the frequency threshold                                                      |
| $Q = (H, P)$ | parameterized tree query with $H$ the head and $P$ the body                  |
| $Q^\alpha, (Q, \alpha)$ | instantiated tree query                                                    |
| $Q^\alpha(G)$ | answer set of the instantiated tree query $Q^\alpha$ in $G$              |
| $\rho$       | parameter correspondence                                                    |
| $Q_2 \subseteq^\rho Q_1$ | $Q_2$ is $\rho$-contained in $Q_1$                                          |
| freeze$_\beta(P)$ | the freezing of a tree pattern $P$                                           |
| pAR          | parameterized association rule                                               |
| iAR          | instantiated association rule                                               |
| $Q_1 \Rightarrow^\rho Q_2$ | pAR from $Q_1$ to $Q_2$                                                     |
| $(Q_1 \Rightarrow^\rho Q_2, \alpha)$ | iAR from $Q_1$ to $Q_2$                                                     |
| minconf      | the confidence threshold                                                    |
| $Freq(P^\alpha)$ | the frequency of $P^\alpha$ in $G$                                          |
| $(\Pi, \Sigma)$ | a parameterized tree pattern $P$ based on a fixed tree $T$                   |
| $(\Pi, \Sigma, \alpha)$ | an instantiated tree pattern $P$ based on a fixed tree $T$                   |
| CanTab$_{\Pi, \Sigma}$ | $\{ \alpha | P^\alpha$ is a candidate instantiated tree pattern $\}$   |
| FreqTab$_{\Pi, \Sigma}$ | $\{ \alpha | P^\alpha$ is a frequent instantiated tree pattern $\}$  |
| $\delta$     | answer set correspondence                                                   |
| $P_1 \equiv^\rho P_2$ | $P_1$ is $(\delta, \rho)$-equivalent with $P_2$                           |
| $P_2^{\ast}(G) \circ \delta$ | $\{ f \circ \delta : f \in P_2^{\ast}(G) \}$                        |
| $P_1 \simeq P_2$ | $P_1$ and $P_2$ are isomorphic                                              |