EFFICIENT TOLERANT PATTERN MATCHING WITH CONSTRAINT ABSTRACTIONS IN DESCRIPTION LOGIC

Carsten Elfers, Stefan Edelkamp and Otthein Herzog
Center for Computing and Communication Technologies, University of Bremen, Am Fallturm 1, 28359 Bremen, Germany

Keywords: Tolerant pattern matching, Constraint abstraction, Description logic, Constraint satisfaction.

Abstract: In this paper we consider efficiently matching logical constraint compositions called patterns by introducing a degree of satisfaction. The major advantage of our approach to other soft pattern matching methods is to exploit existing domain knowledge represented in Description Logic to handle imprecision in the data and to overcome the problem of an insufficient number of patterns. The matching is defined in a probabilistic framework to support post-processing with probabilistic models. Additionally, we propose an efficient complete algorithm for this kind of pattern matching, which reduces the number of inference calls to the reasoner. We analyze its worst-case complexity and compare it to a simple and to a theoretical optimal algorithm.

1 INTRODUCTION

Conventional pattern matching methods determine if a given pattern is satisfied or not. In real-world domains these approaches suffer heavily from uncertainty in the environment in form of typical noise or imprecision in the data. This leads to the natural conclusion that matching patterns must become a matter of degree (Dubois and Prade, 1993).

Our application area is improved network security within Security Information and Event Management (SIEM) systems that collect events from several intrusion detection sensors distributed in a computer network. The typical huge amount of events collected by a SIEM system calls for pattern matching correlation techniques to be reduced to the most relevant ones. Some well-known examples of rule-based approaches are NADIR (Hochberg et al., 1993), STAT (Porras, 1993) and IDIOT (Kumar and Spafford, 1995). However, due to constantly varying attacks and varying network configurations the pattern matching method must deal with changing conditions. Variations can be learned, e.g., by Pattern Mining approaches like in MADAM ID (Lee and Stolfo, 2000). In real world domains, the amount of reference data with detected professional successful attacks is sparse. Therefore, modeling patterns have been a dominating strategy in enterprise SIEM systems like in the market-leading system ArcSight (cf. (Nicolett and Kavanagh, 2010)).

In this paper an efficient tolerant pattern matching algorithm is proposed, which handles variations by exploiting ontological (or categorical) background knowledge in Description Logics (DL). A semantically well-defined and intuitive method is presented to calculate the degree of matching from patterns and data. In contrast to the similarity measurement to handle noise in the matchmaking process suggested by He et al. (2004), our approach does not require similarity weights. Additionally, we extend their work by also describing how to handle disjunctions and negations of conditions (or constraints). Previous work that uses DL for pattern matching in the security domain (Undercoffer et al., 2003; Li and Tian, 2010) do not handle variations that deviate from the modeled patterns. We present an efficient algorithm for finding the best tolerant matching; a critical requirement in real-world domains.

2 PATTERN MATCHING

Our tolerant pattern matching approach is based on constraint satisfaction in ontologies specified in DL. It consist of concepts (classes of objects), roles (binary relations between concepts) and individuals (instances of classes) (cf., (Gomez-Perez et al., 2004, p. 17)). The semantics are defined by the interpretation (an interpretation can be regarded as the corresponding domain) of the set of all concept names, the set of all role names and the set of all names of the individuals (cf., (Baader et al., 2008, p. 140)). A concept C
is subsumed by a concept \( D \) if for all interpretations \( I \) we have \( C^I \subseteq D^I \). In this work, tolerant pattern matching is realized by successively generalizing the pattern, and determining a residual degree of satisfaction.

**Definition 1** (Entity, Constraint, Satisfaction). An entity \( E \) is either (a) an individual, (b) a concept, or (c) a variable. A constraint \( \gamma \in \Gamma \) is defined as \( \gamma = eRe' \) of a left hand side entity \( e \), a right hand side entity \( e' \) and a relation \( R \) between these entities. It is assumed that either \( e \) or \( e' \) is an individual (fixed by an observation), or a variable. A constraint \( \gamma \) is satisfied if there exists an interpretation \( I \) with \((eRe')^I\).

**Definition 2** (Partially Matching Pattern, Degree of Matching). A pattern \( p \) consists of a set of constraints and logical compositions among them. A partially matching pattern \( p \) - given the data \( x \) - is a real valued function with range \([0,1]\). The value of such a function is called degree of matching or matching degree.\(^{1}\)

\[
p(x) = \begin{cases} 
1 & \text{if } p \text{ fully matches} \\
\alpha \in [0,1] & \text{if } p \text{ matches to degree } \alpha \\
0 & \text{otherwise}
\end{cases}
\]

Each constraint in a pattern can be expressed as a query triple in DL. This allows an easy transformation of patterns into a query language like SPARQL, which can be interpreted by DL reasoners. Next, a relation \( \geq_\gamma \) (adapted from Defourneaux and Peltier (1996)) describes that a constraint is an abstraction of another constraint. We say \( \gamma_1 \) is more general or equal than \( \gamma_2 \) (noted \( \gamma_1 \geq_\gamma \gamma_2 \)) if for all interpretations \( p \) of \( \gamma_2 \) there exists an interpretation \( \gamma_1' \) of \( \gamma_1 \) such that \( \gamma_2^p \subseteq \gamma_1'^p \).

Fig. 1 shows an example of a pattern \( p \) with three constraints \( \gamma_1, \gamma_2, \gamma_1' \) and their direct abstractions \( \gamma_1', \gamma_2', \gamma_1'' \). (Superscripts enumerate different levels of specialization. A zero denotes the most abstracted case, while a larger number indicates an increasing specialization, e.g., \( p^0 \) is the direct abstraction of \( p^1 \).)

To abstract a pattern it is first necessary to propagate all negations to the leaves (i.e., to the constraint triples \( \gamma \)) by applying De Morgan’s rules. The construction of this negational normal form can be done before abstracting a pattern or be applied to all patterns in advance (even to the most specific ones). Moreover, each constraint must be abstracted by the following rules:

- A negated constraint is abstracted to a tautology, since the current model of such a constraint includes all individuals except of the negated one (e.g., \( \top^p \) in Fig. 1 has changed to a tautology).
- If the entity of the abstracted constraint is a concept or an individual this is replaced by a more general concept due to the definition of \( \geq_\gamma \).
- The relation of the abstracted constraint might have to be replaced to ensure that the set of interpretations of the constraint increases, e.g., the identity relation must be exchanged to an appropriate (transitive) subclass relation.

A similarity measure \( \theta(\gamma_1, \gamma_2) \) for constraints \( \gamma_1 \) and \( \gamma_2 \) is assumed to quantify the similarity of an abstracted constraint \( \gamma \) from the original level \( j \) to an abstract level \( k \). A simple example of such a measure is \( \theta(\gamma_1, \gamma_2) = 1/(|j-k|+1) \). We write \( \gamma_1^j \) for the original constraint on the most specific level \( \top \), and \( \theta(\gamma_1, \gamma_2) \) for \( \theta(\gamma_1^j, \gamma_2^j) \). Independent of a concrete realization, such a measurement is assumed to be 1 if the constraint is not abstracted, and decreases, if the constraint is getting more abstract; by still being greater than or equal to 0. This measurement can also be regarded as a similarity function, which says how exactly \( \gamma_1 \) describes \( \gamma_2 \), or how similar they are.

**Definition 3** (Similarity, extended from Fanizzi and d’Amato (2006) and Batagelj and Bren (1995)). A similarity measure \( \theta \) is a real-valued function into \([0,1]\) defined by the following properties:

- \( \forall \gamma_1, \gamma_2: \theta(\gamma_1, \gamma_2) \geq 0 \) (positive definiteness)
- \( \forall \gamma_1, \gamma_2: \theta(\gamma_1, \gamma_2) = \theta(\gamma_2, \gamma_1) \) (symmetry)
- \( \forall \gamma_1, \gamma_2: \theta(\gamma_1, \gamma_2) \leq \theta(\gamma_1, \gamma_2') \) (identity)
- \( \forall j < k: \theta(\gamma_1, \gamma_2^j) < \theta(\gamma_1, \gamma_2^k) \) (monotonicity)

Such similarity function values of the constraints are combined to a matching degree of the pattern by applying some fusion operator \( F(\theta_1, \ldots, \theta_n) \) similar to fuzzy pattern matching (cf., (Cadenas et al., 2005)) to consider the semantics of the logical operators while abstracting the pattern. A probabilistic fusion approach is suggested by using a Bayesian interpretation of the tree of logical operators in each pattern as follows.\(^2\)

---

\(^{1}\) As we will see later, the degree of matching is determined by a fusion function \( F \).

\(^{2}\) These equations naturally result from a Bayesian net-
\textbf{Definition 4} (Fusion Function). The fusion function \( F(p) \) of pattern \( p \) is recursively defined with respect to some similarity function \( \theta \) of constraints \( \gamma \) composed by logical operators.

\[
F(\gamma_1 \land \gamma_2) = F(\gamma_1) \cdot F(\gamma_2)
\]
\[
F(\gamma_1 \lor \gamma_2) = 1 - \left( 1 - F(\gamma_1) \right) \cdot \left( 1 - F(\gamma_2) \right)
\]
\[
F(\neg \gamma) = \begin{cases} 
\beta \cdot F(\gamma), & \text{for } i = \bot \\
1 - F(\gamma), & \text{otherwise}
\end{cases}
\]
\[
F(\gamma) = \theta(\gamma),
\]

where \( \beta \) is a penalty factor to additionally penalize the abstraction of negations, since these may have a greater impact on the result.

The factor \( \beta \) may depend on the used similarity function and the depth of the ontology. The fusion function’s conjunction/disjunction can be regarded as deterministic AND/OR nodes. Therefore, the conditional probability table is fully specified.

The negation is interpreted differently to a corresponding Bayesian conditional probability table to ensure that an increasing abstraction leads to a decreasing fusion function. This leads to the monotonicity of \( F \) wrt. \( \theta \), which is very useful for finding the most specific abstraction as we will see later.

With these properties a partial order of patterns with respect to the generality of their containing constraints is defined. From this basis it is necessary to find the best matching pattern, i.e., the pattern with the biggest \( F \). This problem can be postulated for a pattern \( p \) containing \( d \) constraints \( \gamma^1, \ldots, \gamma^d \) to find a combination of \( x_1, \ldots, x_d \) which satisfies the pattern and maximizes \( F \). The solution of interest is in the Pareto front of maximum \( x \) due to the monotonicity of \( F \) with respect to \( \theta \) and the monotonicity of \( \theta \) itself (cf. Def. 3). If the level of specialization increases, \( F \) increases as well, or – in other words – if any constraint of the pattern is abstracted, \( F \) decreases.

\section{Algorithm Pareto}

In this section a divide-and-conquer algorithm is proposed to efficiently search for the most specific satisfied patterns, building the Pareto front of the constraint abstractions. Each level of abstraction of a constraint is represented as one dimension of the search space. The search space \( X = \{0, \ldots, n-1\}^d \) is divided into satisfied elements (satisfied constraint combinations) \( X^+ \subseteq X \) and unsatisfied elements \( X^- \subseteq X \) with \( X^+ \cap X^- = \emptyset \).

3 ALGORITHM PARETO

\begin{algorithm}
\caption{Algorithm Pareto}
\end{algorithm}

Fig. 2 gives an example how the algorithm works for the 2D case (i.e., for \( \gamma_1 \) and \( \gamma_2 \)). At first the middle of the search space is determined, i.e., point (4,4). Around this point the search space is divided into (in the 2D case) four equal sized areas each including the middle and the corresponding border elements. Two of these areas are marked with a gray background the others are area A and area B. The minus sign at (4,4) indicates that the pattern with \( \gamma_1 \) and \( \gamma_2 \) is unsatisfied, the circle indicates an inference call to test this satisfaction. Therefore, all more specific pattern combinations are omitted in the further recursion, i.e., area A. This method is continued for the gray areas but at first for area B. Area B is divided into four equal sized areas around the middle (2,2). This is a match, therefore, we know that each more abstract pattern than \( \gamma_1 \) and \( \gamma_2 \) is also matching (or satisfied), marked as area C which can be omitted in the following. The recursion is continued for the new middle (3,3). At this point an unsatisfied area can be determined which also affects the currently not investigated gray areas due to we know that from (3,3) to (8,8) every solution must be unsatisfied because they are more or equal specific. This temporary results are stored in a list and checked before investigating the gray areas in subsequent recursion steps to omit inference calls for these points.

The algorithm can be limited in the search space (by limiting the search depth) to give approximate results. By increasing the search depth the solution is more and more appropriately approximated.

Algorithm 1 shows the implementation of the approach. This is initialized with an empty set of solutions (representing the most specific satisfied patterns) \( S^+ \) and \( S^- \) (representing the most abstract unsatisfied patterns). The individual search spaces are specified by a most specific bound (msb) and a most abstract bound (mab), where msb and mab are coordinates of the search space. Initially, for all \( i \) we have msb, \( i = 0 \) and mab, \( i = n \) (to ensure completeness mab is located outside of the actual search space). For
Algorithm 1: Pareto.

Input: msb, mab ∈ X = \{(x_1, \ldots, x_d)\}^d
1: \begin{align*}
  m &= \lfloor (\text{mab} + \text{msb})/2 \rfloor \\
  \text{if } \exists x \in S \text{ with } (m \geq s) \text{ then } \\
  \text{s} &= \text{true} \\
  \text{else } &
\end{align*}
2: \begin{align*}
  s &= \text{false} \\
  \text{else } &
\end{align*}
3: \begin{align*}
  m &= \text{true} \\
  \text{else } &
\end{align*}
4: \begin{align*}
  \text{if } \exists x \in S \text{ with } (m \leq s) \text{ then } \\
  s &= \text{true} \\
  \text{else } &
\end{align*}
5: \begin{align*}
  s &= \text{Eval}(m) \\
  \text{if } s &= \text{true then } \\
  S' &= \{x \in S' \cup (m) \mid \forall x' \in S' \cup (m) : x' \geq x\} \\
  \text{else } &
\end{align*}
6: \begin{align*}
  S' &= \{x \in S' \cup (m) \mid \forall x' \in S' \cup (m) : x' \leq x\} \\
  \text{if } mab &= m \text{ then } \\
  \text{return } &
\end{align*}
7: \begin{align*}
  \text{if } s &= \text{true then } \\
  \text{Pareto}(m, m) &
\end{align*}
8: \begin{align*}
  \text{Pareto}(m, mab) &
\end{align*}
9: \begin{align*}
  \text{for each } h \in \text{Hypercubenodes}(m, mab) &
\end{align*}
10: \begin{align*}
  \text{for } e = 1 \text{ to } d &
\end{align*}
11: \begin{align*}
  \text{if } mab &= \text{max}(h, m) &
\end{align*}
12: \begin{align*}
  \text{if } mab &= \text{min}(h, m) &
\end{align*}
13: \begin{align*}
  \text{Pareto}(m, mab) &
\end{align*}

reasons of simplicity, each constraint is assumed to have an equal amount of specializations/abstractions, however, the algorithm is also capable of differing amounts.

Besides Eval, the call to the reasoner, Hypercubenodes(msb, mab) enumerates the sublattices (without msb, mab themselves); formally defined as

\[ \bigcup_{i=1}^{2d-2} \text{msb} \odot \text{bin}(i) + \text{mab} \odot \overline{\text{bin}(i)}, \]

where bin(i) denotes the binary representation of a number i, bin(i) denotes its (bitwise) complement, and \( \odot \) the bitwise multiplication of two vectors.

The definition for \( \geq_g \) in the d dimensional search space \( X \) is given by:

**Definition 5 (Domination).** Let

\[ \geq_g = \{(x, x') \in X^2 \mid \forall i (x_i \leq_g x'_i)\}. \]

We say that \( x \in X^+ \) dominates \( x' \in X \) if \( x' \leq_g x \) and \( x \in X^+ \) dominates \( x' \in X \) if \( x' \geq_g x \).

All patterns more specific than an unsatisfied one are still unsatisfied and all patterns more general than a satisfied one are still satisfied. In other words, we have \( \forall x' \in X^-, x' \in X \Rightarrow x' \leq_g x \) and \( \forall x \in X^+, x' \in X \Rightarrow x' \geq_g x \).

**Definition 6 (Pareto Frontier).** The Pareto frontier is the set of extreme points \( E^+ = E^+ \cup E^- \) with \( E^+ \cap E^- = \emptyset \) containing each element of \( X^+ \) with no element in \( X^+ \) being more general

\[ E^+ = \{x \in X^+ \mid \forall x' \in X^+ (x' \geq_g x)\} \]

and each element of \( X^- \) with no element in \( X^- \) being more specific

\[ E^- = \{x \in X^- \mid \forall x' \in X^- (x' \leq_g x)\}. \]

No element in \( E \) is dominated by another element in this set, i.e., the compactest representation of the set of satisfied/unsatisfied solutions.

Next, we show that Algorithm 1 computes \( E^+ \).

**Theorem 1** (Correctness and Completeness of Algorithm 1). The algorithm determines the whole set of satisfied constraints, i.e., \( E^+ = S^+ \).

**Proof.** (Correctness) To show the correctness of the algorithm we ensure that each element of the expected result set \( E^+ \) is in the solution set \( S^+ \) of the algorithm and, vice versa, i.e., \( e^+ \in E^+ \Rightarrow e^+ \in S^+ \) and \( s^+ \in S^+ \Rightarrow s^+ \in E^+ \).

**Lemma 1.** \((s^+ \in S^+ \Rightarrow s^+ \in E^+)\)

If the search is exhaustive (this is shown later) Line 10 implies that \( s^+ \in S^+ \Rightarrow s^+ \in E^+ \), since it computes \( S' \) as \( \{x \in (S' \cup (m)) \mid \forall x' \in (S' \cup (m)) : x' \geq_g x\} \), which is the same as the expected result \( E \) with \( S^+ \cup (m) \subseteq X^+ \).

**Lemma 2.** \((e^+ \in E^+ \Rightarrow e^+ \in S^+)\)

We investigate four conditions under which an element is inserted into (and kept in) the solution set of the algorithm \( S^+ \). These conditions, directly derived from the algorithm, are as follows

1. Each element from \( S^+ \) must be contained in \( X^+ \) which is exactly the same condition as in definition of \( E^+ \).
2. The following assumption derived from Lines 2 and 4, must hold for \( e^+ \) to be inserted into \( S^+ \)

\[ \neg \exists x' \in S^+. (e^+ \geq_g x'). \]

This condition is not fulfilled if an equivalent solution \( e^+ \) is already in the set \( S^+ \) or if \( e^+ \) dominates another element from \( S^- \). In both cases \( e^+ \) is not inserted into the result set \( S^+ \).
3. The next statement, derived from Line 5 and 7, is

\[ \neg \exists x' \in S^- . (e^+ \geq_g x'). \]

This condition is always fulfilled, since we consider the case that \( e^+ \in X^+ \) which implies that \( x' \in X^+ \) which cannot be the case since \( x' \in S^- \subseteq X^- \).
4. Line 10 does not drop solutions because for all \( m \in X^+ \) we have Eqn. 1.

Analogically, the proof can be made for \( E^- \). □

**Proof.** (Completeness)

Recursion is omitted for
• \( \{ x \in X \mid \text{msb} \leq x \leq m \} \) if \( m \in X^− \) and for
• \( \{ x \in X \mid m \leq x \leq \text{mab} \} \) if \( m \in X^+ \).

This does not affect the set of solutions due to the definition of domination and the definition of \( E \) that there should not be any value in the result set that is dominated by another element. Note that \( m \) has already been checked by the algorithm at this point.

The remaining space under investigation is getting smaller in each recursion path until \( m \) is getting equal to \( \text{mab} \) (termination criterion in Line 13). This is only the case if each edge of the space under investigation is smaller or equal one (Line 1). At some time in the recursion the space of possible solutions is divided into a set of spaces with edges of the length one or less by still covering the whole space of possible solutions as previously shown. Further, if any point of such a smallest area is a possible solution (these are the corners), this point is under investigation in another space due to the recursive call with overlapping borders except of the borders of the whole search space in the specific case due to there is no \( \text{mab} \) of any area including these specific border elements, e.g., there is no \( \text{mab} \) for the one element area \((8,8)\) in the example from Fig. 2. For this border case the algorithm is called with a lifted \( \text{msb} \) to ensure that the unlifted specific bound is included in some smallest (one element) area as \( \text{mab} \) visualized as a light gray border in Fig. 2. Therefore, each element of the search space which is a possible solution is investigated as a \( \text{mab} \) in some recursive path.

After computing the Pareto front, the fusion function \( F \) identifies the most specific matching pattern abstraction in the remaining set of candidates.

We observe that the (worst-case) running time complexity is dominated by the number of calls to the DL reasoner. In the following analysis, we distinguish between the number of recursive calls \( T(n) \) and the number of inference calls \( C(n) \) (for the sake of simplicity, we assume \( n_1 = \ldots = n_k \) and \( n = 2^k \)).

Of course, a trivial algorithm testing all possible elements in \( S \) induces \( C(n) = T(n) = O(n^2) \). We will see that the algorithm \textit{Pareto} is considerably faster.

With \( \ln n \) we refer to the dual logarithm \( \log_2 n \), while \( \ln n \) refers to the natural logarithm \( \log n \).

**Recursive Calls.** For the 2D case, the number of calls of the divide-and-conquer algorithm in a \( 0/1 \) \((n \times n)\) matrix is bounded by

\[
T(k) = \sum_{i=0}^{k} 3^i = \left( 3^{k+1} - 1 \right) / 2
\]

Assuming \( n = 2^k \), we have

\[
T(n) = \left( 3^{\lg n} - 1 \right) / 2 = \left( n^{\lg 3} - 1 \right) / 2 = O(n^{1.5849})
\]

For larger dimensions \( d \) the complexities \( O(n^{g(2^d-1)}) \) rise.

**Inference Calls.** For the 2D case the structure of the recursion corresponds to find a binary search to the SAT/UNSAT boundary. The recursion depth is bounded by \( \lg n \). Therefore, the worst-case number of calls to the reasoner of the algorithm in a \( 0/1 \) \((n \times n)\) matrix is defined by

\[
C(n) = 2C(n/2) + O(\lg n).
\]

The \( O(\lg n) \) term is due to the binary search. In the worst case the boundary between SAT and UNSAT cells is in the middle, where one quarter of SAT and one quarter of UNSAT elements are omitted.

Using the Akra-Bazzi theorem (Akra and Bazzi, 1998), the above recursion can be shown to reduce to \( C(n) = O(n) \) as follows. For \( k = 0 \) it states that for recurrence equation \( T(n) = g(n) + aT(n/b) \) with \( a = b^p \) we have the following closed form

\[
T(n) = O \left( n^p \left( 1 + \int_1^n \frac{g(u)}{u^2} \, du \right) \right).
\]

Here, \( g(n) = \ln n = \ln n / \ln 2 \) and \( a = b = 2 \) so that \( p = 1 \) and

\[
T(n) = O \left( n + n \cdot \int_1^n \ln(u) / u^2 \, du \right)
\]

\[
= O(n + n \cdot [- \ln u / u]^n) = O(n).
\]

For larger dimensions \( d \) the complexities \( O(n^{g(2^d-2)}) \) rise.

### 4 EVALUATION

We have evaluated the efficiency of the algorithm with respect to the number of inference calls. In Fig. 3 two tolerant matching algorithms and the result of a perfect guessing algorithm (a lower bound) are visualized. It is assumed that the lower bound algorithm checks exactly the Pareto border of satisfied and unsatisfied elements. Therefore, the best possible algorithm needs at least \( |S^+| + |S^-| \) inference calls.

The proposed divide-and-conquer algorithm \textit{Pareto} with pruning the recursive calls as in Algorithm 1 – but without using the lists \( S^+ \) and \( S^- \) – is called \textit{Pareto-0}. This is the first efficient algorithm one might think of. The proposed algorithm is visualized as \textit{Pareto} and the lower bound as \textit{LOWERBOUND} for the 2D case in Fig. 3 (no log-scale) and for the 4D case in Fig. 4 (log-scale). The x-axis represents the amount of possible abstractions and the y-axis the amount of reasoner calls.
Both figures show that the inference calls of Pareto is near to the optimal lower bound LOWER-BOUND and considerably better than the typical divide-and-conquer algorithm Pareto-0 in both 2D and 4D. These results are reasonable for the proposed pattern matching algorithm due to the depth of an ontology being typically smaller than 30 and the patterns having typically a small amount of constraints.

We see that the amount of Pareto results, which is around the half of the LOWERBOUND value, is very small. For this set the degree of matching must be computed with respect to the fusion function to find the most optimal solution out of the set of Pareto-optimal solutions. This search can be done without to call the DL reasoner, since we already know that these solutions are satisfied.

5 CONCLUSIONS

We have shown how to use ontological background DL knowledge to overcome the problem of noisy and imprecise data. Our tolerant pattern matching approach can even address erroneous or missing patterns by successively abstracting them. The algorithm substantially reduced the amount of these inference calls to the DL reasoner. It is correct, complete and needs a number of inference calls close to the lower bound. It can be parameterized to have the ability to infer approximate results.

ACKNOWLEDGEMENTS

This work was supported by the German Federal Ministry of Education and Research (BMBF) under the grant 01IS08022A.

REFERENCES


