

DynamiTE: Parallel Materialization of Dynamic RDF Data

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Abstract. One of the main advantages of using semantically annotated data is that machines can *reason* on it, deriving implicit knowledge from explicit information. In this context, *materializing* every possible implicit derivation from a given input can be computationally expensive, especially when considering large data volumes.

Most of the solutions that address this problem rely on the assumption that the information is static, i.e., that it does not change, or changes very infrequently. However, the Web is extremely dynamic: online newspapers, blogs, social networks, etc., are frequently changed so that outdated information is removed and replaced with fresh data. This demands for a materialization that is not only *scalable*, but also *reactive* to changes.

In this paper, we consider the problem of *incremental materialization*, that is, how to update the materialized derivations when new data is added or removed. To this purpose, we consider the ρ df RDFS fragment [11], and present a parallel system that implements a number of algorithms to quickly recalculate the derivation. In case new data is added, our system uses a parallel version of the well-known semi-naive evaluation of Datalog. In case of removals, we have implemented two algorithms, one based on previous theoretical work, and another one that is more efficient since it does not require a complete scan of the input for every update.

We have evaluated the performance using a prototype system called *DynamiTE*, which organizes the knowledge bases with a number of indices to facilitate the query process and exploits parallelism to improve the performance. The results show that our methods are indeed capable to recalculate the derivation in a short time, opening the door to reasoning on much more dynamic data than is currently possible.

1 Introduction

One of the main advantages of using semantically annotated data is that machines can *reason* on it, deriving new, implicit, knowledge from existing information. To this end, several systems have been developed over the last few years to make all possible conclusions from a given input explicit, so that no reasoning is needed when the user queries the knowledge base.

This task, also called *materialization*, can be computationally expensive, especially if the input is large. In fact, while some of these systems can perform the materialization of several billions of statements [9, 15, 19], they still demand for significant computational resources, and the process may require up to a few days to complete. Because of this, most of these systems work under the assumption that the input data is static, i.e., that it does not change, or changes very infrequently. This assumption does not match with the current Web, which is extremely dynamic: online newspapers, blogs, social networks, are frequently changed and updated. This demands for a materialization process that is not only scalable, but also *reactive* to changes. A new research area, called *stream reasoning*, has recently emerged to address this specific problem [3].

With this paper, we contribute to this area by considering the problem of *incrementally maintaining* a large materialized knowledge base in the presence of frequent changes, using monotonic rule-based reasoning as the method to derive new information. More specifically, we propose *DynamiTE*, a parallel system capable of efficiently generating the complete materialization of large RDF knowledge bases, and maintaining it after the knowledge base is updated.

We consider two types of updates: (i) the *addition* of new information, which requires a re-computation of the materialization to add new derivations, and (ii) the *removal* of existing information, which requires the deletion of the explicit knowledge, and also of all the implicit information that is no longer valid. For the addition updates, *DynamiTE* applies in parallel the well-known Datalog semi-naive evaluation. For the removal updates, it implements two algorithms: one that was presented in the literature but only from a theoretical perspective [10], and another one, which is more efficient since it does not require a complete scan of the input for every update.

To evaluate the performance of *DynamiTE*, we consider the minimal RDFS fragment (ρ df [11] ruleset), which captures the main semantic functionality of RDFS [8], and can be expressed with Datalog [1]. We designed some experiments to study the behavior of our system over large amounts of data, trying to understand, from a system point of view, what are the main factors that drive the performance. The results show that our system is capable of efficiently computing the materialization of large knowledge bases up to one billion statements, and can alter them in a range from hundred of milliseconds to less than two minutes when considering substantial updates of two hundred thousand triples.

The remaining of this paper is organized as follows: Section 2 contains some background information to make the reader familiar with the concepts we use throughout the paper. Next, Section 3 reports an overview of our system while Sections 4 and 5 focus on the crucial task of the system, i.e., the incremental maintenance of the materialization. After this, we present an evaluation of our system in Section 6. Finally, Section 7 discusses related work and Section 8 concludes the paper, also reporting some indications for future research.

2 Background

To describe our system, we use the notions and notations of the Datalog language. Because of space constraints, we cannot present a complete overview of this language. Therefore, we only present some basic concepts that we use in the paper, and we refer the reader to existing literature, e.g. [1], for more details.

First of all, a Datalog *program* P is defined as a finite set of rules in the form $R_1(w_1) \leftarrow R_2(w_2), R_3(w_3), \dots, R_n(w_n)$, where each component $R_i(w_i)$ is called a *literal*. A literal is composed of a *predicate* (e.g. R_i) and a tuple of terms $w_i := t_1, \dots, t_m$. A term t_i can be either a variable from a finite domain V or a constant term from another disjoint finite domain C . We denote with $t_j \in R_i(w_i)$ the term that appears at the j^{th} position of w_i .

We define $\text{var}(R_i(w_i))$ as the set of all variables in the literal $R_i(w_i)$, and $\text{const}(R_i(w_i))$ as the set of all its constants. The left side of a rule r is called the *head* of r ($\text{head}(r)$), while the right side is defined as its *body* ($\text{body}(r)$). Datalog imposes that each variable that appears in the head of the rule must also appear in its body. This means that $\forall v \in \text{var}(R_1(w_1))$ there must be an $i \in (2..n)$ so that $v \in \text{var}(R_i(w_i))$. Furthermore, Datalog makes a distinction between *edb* predicates, which never appear in the head of a rule, and *idb* predicates, which appear in the head of some rule.

A literal containing only constants is called a *fact*. We say that a fact f *instantiates* the literal l if they share the same predicate, and every $c_i \in \text{const}(l)$ that appears in l at position i is equal to the corresponding term t_i of f . We call $f_1 \leftarrow f_2, f_3, \dots, f_n$ an *instantiation* of a rule $R_1(w_1) \leftarrow R_2(w_2), R_3(w_3), \dots, R_n(w_n)$ if every $f_{i \in \{1..n\}}$ is an instantiation of the corresponding $R_i(w_i)$, and that any term $t_j \in f_i$ is equal to another term $t_m \in f_{j \in \{1..n\} \wedge i \neq j}$ if $v_i \in \text{var}(R_i(w_i))$, $v_j \in \text{var}(R_j(w_j))$, and $v_i = v_j$.

In Datalog, the operator T_P is called the *immediate consequence operator* of P . This operator maps a generic database \mathfrak{J} to another database $T_P(\mathfrak{J})$ that contains all facts that are *direct consequences* for \mathfrak{J} and P . A fact f is a direct consequence for \mathfrak{J} and P if either $f \in \mathfrak{J}(R)$ for some *edb* predicate R in P or $f \leftarrow f_1, f_2, \dots, f_n$ is an instantiation of a rule in P and each $f_{i \in \{1..n\}} \in \mathfrak{J}$. Intuitively, T_P can be seen as the abstract operator that applies the rules in P over \mathfrak{J} to derive new conclusions.

T_P is monotonic: given two databases $\mathfrak{J}, \mathfrak{J}'$, if $\mathfrak{J} \subseteq \mathfrak{J}'$, then $T_P(\mathfrak{J}) \subseteq T_P(\mathfrak{J}')$. Because of this, repeated executions of the T_P operator over augmented versions of a database \mathfrak{J} will lead to a fix-point. This means that if we define $T_P^n(\mathfrak{J})$ as

$$T_P^n(\mathfrak{J}) = \begin{cases} \mathfrak{J} & n = 0 \\ T_P(T_P^{n-1}(\mathfrak{J})) & n > 0 \end{cases}$$

there is a n' such that $T_P^{n'+1}(\mathfrak{J}) = T_P^{n'}(\mathfrak{J})$. In this case, $T_P^{n'}(\mathfrak{J})$ is named as the fix-point of T_P , and denoted with $T_P^\omega(\mathfrak{J})$. $T_P^\omega(\mathfrak{J})$ is the *materialization* of \mathfrak{J} , since it contains all the possible derivations that can be obtained from \mathfrak{J} and P .

A trivial way to compute $T_P^\omega(\mathfrak{J})$ is to start from $n = 0$, execute T_P , and increase n until the fix-point is reached. This approach, known as the *naive*

Head	Body
$T_i(A, P, B)$	$\Leftarrow T_e(A, P, B)$
$T_i(A, \text{SPO}, C)$	$\Leftarrow T_i(A, \text{SPO}, B), T_i(B, \text{SPO}, C)$
$T_i(A, P, B)$	$\Leftarrow T_i(Q, \text{SPO}, P), T_i(A, Q, B)$
$T_i(A, \text{TYPE}, C)$	$\Leftarrow T_i(B, \text{TYPE}, C), T_i(A, \text{TYPE}, B)$
$T_i(A, \text{SCO}, C)$	$\Leftarrow T_i(A, \text{SCO}, B), T_i(B, \text{SCO}, C)$
$T_i(A, \text{TYPE}, D)$	$\Leftarrow T_i(P, \text{DOMAIN}, D), T_i(A, P, B)$
$T_i(A, \text{TYPE}, R)$	$\Leftarrow T_i(P, \text{RANGE}, R), T_i(B, P, A)$

Table 1: Supported ruleset. Datalog variables are in italics; constants are in fixed-width characters. The abbreviations **SPO**, **SCO**, **TYPE**, **RANGE** and **DOMAIN** stand for the URIs *rdfs:subProperty*, *rdfs:subClassOf*, *rdf:type*, *rdfs:range*, and *rdfs:domain*. The first rule is only used to map the *edb* predicate T_e to the *idb* predicate T_i that is used in the other rules.

evaluation, is very inefficient since, at each iteration, an application of T_P will recompute all the derivations already computed in the previous iterations.

A more efficient algorithm, called *semi-naive* evaluation [1], optimizes this process by instantiating a rule r only if at least one fact that instantiates a literal in $body(r)$ was derived in the previous iteration. In this way, the algorithm is able to significantly decrease the number of duplicates.

The semi-naive evaluation can be implemented by annotating each fact in the database with a numeric step that indicates at which stage of the derivation that information was derived (facts in the original input are marked with a step of zero). Then, at every n^{th} iteration of the evaluation, the operator T_P accepts only instantiations if at least one fact that instantiates a literal has a step that is equal or greater than $n - 1$. This significantly reduces the number of rules execution and consequently the number of duplicates that are generated.

3 DynamiTE: System Overview

The purpose of *DynamiTE* is to efficiently compute and incrementally maintain the materialization of a database, which consists of RDF triples. We consider the minimal RDFS fragment ρdf [11], and execute the rules presented in Table 1.

To formalize our problem in Datalog, let P be a program consisting of the rules in Table 1, and \mathcal{J} a given database, which represents the initial RDF knowledge base expressed as a set of Datalog facts $T_e(s, p, o)$ where T_e is an *edb* predicate, and s, p, o are respectively the subject, predicate, and object of a RDF triple that are mapped to constant terms in Datalog.

DynamiTE implements three main tasks: (i) First, it computes the complete materialization of \mathcal{J} . Then, it maintains it after a set of triples δ is either (ii) added, or (iii) removed. More formally, in (i) the system calculates $T_P^\omega(\mathcal{J})$, in (ii), $T_P^\omega(\mathcal{J} \cup \delta)$, and in (iii) $T_P^\omega(\mathcal{J} \setminus \delta)$ is computed.

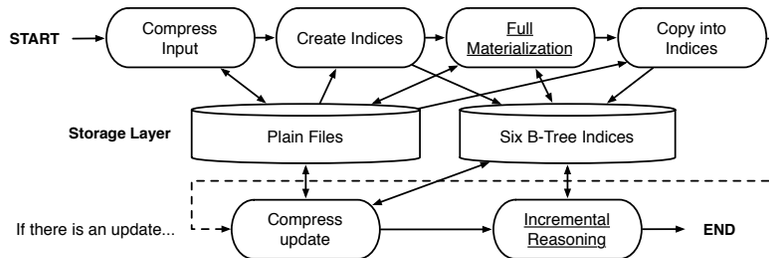


Fig. 1: *DynamiTE*: General System Workflow

3.1 System Workflow

Fig. 1 shows the general workflow of *DynamiTE*. The initial input consists of a collection of triples encoded with the N-Triples format. First, *DynamiTE* performs the *Compress Input* operation, which converts the textual terms into numbers using the technique of dictionary encoding. The algorithm used for this phase is an adaptation of the distributed MapReduce version presented in [16].

The next operation, *Create Indices*, stores the compressed data into B-Tree indices (along with the dictionary tables to allow quick decompression). Since the size of the database can easily become too large to fit in main memory, we need to consider data structures that can be off-loaded to disk. To this end, we use six on-disk B-Trees to store all the possible permutations of the input triples. We chose the B-Tree data structure because we want our system to support generic querying, and storing six indices has proven to be ideal for SPARQL [13] querying, allowing an efficient retrieval for all possible atomic queries [20]. As we will show in the evaluation, the *Create Indices* operation is the most expensive of the entire workflow. To reduce its cost, *DynamiTE* sorts the triples before insertion (in our tests, this improves the performance by at least 10%).

Next, *DynamiTE* performs the *Full Materialization*. We describe this process in detail in Section 4.1. This operation implements a parallel version of the semi-naive evaluation, which iteratively reads the entire input and augments it with new derivations. The B-Trees are not particularly efficient in supporting these operations. Therefore, during the full materialization, the system writes the new derivation on plain files, and copies them on the B-Trees in the next operation, *Copy into Indices*. According to some tests we performed, using files makes the entire process at least 30% faster.

At this point, *DynamiTE* is ready to receive updates (see Fig. 1, bottom). For each update δ , it first compresses the content of δ , and then performs incremental reasoning. We describe this last operation in Sections 4 and 5.

3.2 Physical Rules Instantiation

In order to physically instantiate the rules, we make a fundamental distinction between *schema* and *generic* triples. We denote as *schema* all those triples having

SPO, SCO, DOM, or RANGE as predicate. We call all the others *generic* triples. The design and implementation of our algorithm for rules instantiation relies on the assumption that the number of schema triples in the input is significantly smaller than the rest, so that all of them (explicit and inferred) will fit in main memory. This assumption holds for the vast majority of web data [15], but there can be scenarios where this is no longer true.

First, the system assigns all the rules to three disjoint subsets, named *Type 1*, *Type 2*, and *Type 3*, depending on the number and type of their literals. The assignment criteria for a rule r are defined as follows:

- *Type 1*. All the literals in r can be instantiated only by schema triples.
- *Type 2*. The body of r consists of only one literal and can be instantiated by generic triples.
- *Type 3*. The body of r contains exactly two literals, one of them can be instantiated only by schema triples while the other is instantiated by generic triples.

Note that there are no rules of *Type 2* in Table 1. Nevertheless, we included their execution for generality, so that our system can also be used with other RDFS rules that are not in the *pdf* fragment.

DynamiTE implements a different rules instantiation strategy depending on the type of the rules. Rules of *Type 1* are instantiated by first loading all schema triples, i.e., all the triples that can instantiate the literals in $body(r)$, in memory. Then, in case there is only one literal in $body(r)$ the instantiation becomes trivial since the system only needs to generate a new triple that instantiates $head(r)$ and copy the values of the variable in the body to the corresponding position in the head. If there are multiple literals in the body, then the system must join the triples having common terms. Consider for example the second rule in Table 1: its application needs to find all the pairs of triples s, p, o , and s', p', o' , such that $p = p' = \text{SPO}$ and $o = s'$. The system performs this operation in memory, computing a hash join between the two sets of triples.

Rules of *Type 2* are similar to rules of *Type 1* with only one literal in their body. The only difference is that here the system cannot assume that they fit into the main memory and thus it needs to retrieve them from the disk.

Rules of *Type 3* are the most challenging, since they require a join between two sets of triples, schema and generic triples, where the second set can be quite large. In previous work, we tackled this problem proposing a distributed execution over multiple processing nodes using the MapReduce model and the Hadoop framework [15]. Schema triples were replicated on every node, and a hash join was performed against the generic triples that were being read from the input. While this approach proved to be scalable, it introduced high latency (due to the usage of Hadoop), which conflicts with our need for reactivity. In *DynamiTE*, we re-implemented similar algorithms to replicate the MapReduce programming model without using a resilient and distributed architecture such as Hadoop, but instead exploiting the parallelism offered by modern multi-core hardware to reduce the processing time.

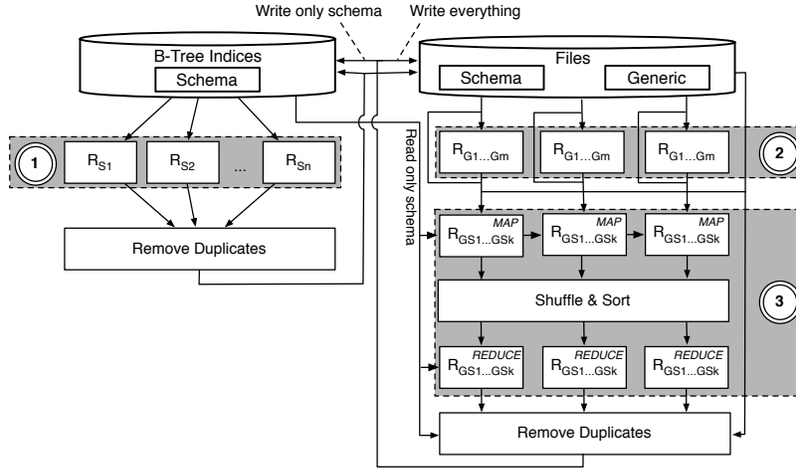


Fig. 2: *DynamiTE*: One iteration of the full materialization. In this figure, there are n rules of *Type 1*, $R_{S1} \dots R_{Sn}$, m rules of *Type 2*, $R_{G1} \dots R_{Gm}$, and k rules of *Type 3*, $R_{GS1} \dots R_{GSk}$.

4 Materialization After Data Additions

We distinguish two types of updates, depending on whether the initial database is empty. In the first case we perform a full materialization, while in the second an incremental materialization is done.

4.1 Full Materialization

For this task, *DynamiTE* provides a parallel implementation of the semi-naive evaluation to exploit the parallelism of modern computer architectures. As we described in Section 2, the semi-naive evaluation performs multiple iterations, until it reaches a fix-point. Fig. 2 shows a single iteration in our system. On top of the figure there is the database, physically stored both on six B-Trees and on a number of files. During the full materialization, we read the database and write the derivation only to files, except for the schema triples that are always being replicated on the B-Trees to improve their retrieval in the next iterations.

First, *DynamiTE* applies all the rules of *Type 1*. This step is shown in the gray box marked with a “1”. The execution is parallel, with each rule r being instantiated in a separate thread t . Each thread t retrieves from the B-Trees the schema triples that instantiate the literals in $body(r)$, joins them, if needed, and generates the triples that instantiate $head(r)$. Finally, it stores all derivations both in the B-Trees and on files.

Next, *DynamiTE* instantiates rules of *Type 2* and *Type 3*. They require a complete scan over the input, which can potentially be large. *DynamiTE* optimizes this step by partitioning the input files into smaller blocks, with each

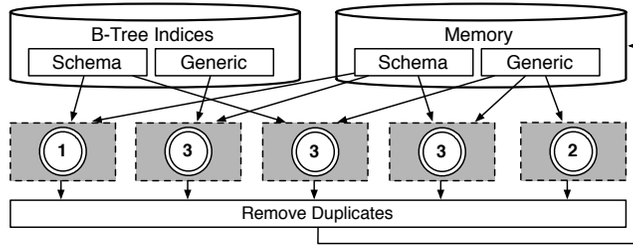


Fig. 3: *DynamiTE*: One iteration of the incremental materialization.

block b being read by a different thread t . First, each thread t applies the rules of *Type 2* on the triples in b . Then, it applies rules of *Type 3*, considering both the original input and the output of the rules of *Type 2*. Rules of *Type 3* are instantiated using the MapReduce-like algorithm outlined before. Notice that, for executing rules of *Type 3*, *DynamiTE* also accesses the B-Trees for retrieving schema triples. After execution, all the derivations are stored on files, while schema triples are also replicated in the B-Trees.

As mentioned, *DynamiTE* implements the semi-naive evaluation, where only the last derivations are considered during rule execution. To make this possible, it marks each triple with a *step* attribute, representing the step when it was first derived. For example, at the first iteration all the triples derived by rules of *Type 1* have step one; the derivations from rules of *Type 2* are marked with step two, and so on. After the first iteration, the system accepts a derivation only if at least one of the triples that instantiate a literal in the body of the rule has a step greater or equal than the current one minus three.

The algorithm stops iterating the rules instantiation when none of them derived new triples. As described in the previous section (see Fig. 1), after the materialization is complete, all the derived triples are copied into the B-Tree indices for efficient querying.

4.2 Incremental Materialization

The previous section showed how *DynamiTE* computes the full materialization of a knowledge base and stores it into B-Tree indices. Now, we show how it maintains this materialization in the presence of data additions.

Our system performs this operation incrementally, fully exploiting the existing materialization $T_P^\omega(\mathcal{J})$. The process can be divided into three main steps: (i) Load all the triples in δ into the main memory. (ii) Perform a semi-naive evaluation to derive new triples. Since all derivations generated only from triples in \mathcal{J} are already present in the materialization $T_P^\omega(\mathcal{J})$, in this phase we consider as “new” only triples in δ (or derived from a triple in δ in previous steps). (iii) Add all the new derivations into the B-Tree indices, making them available for querying.

Fig. 3 shows how the semi-naive evaluation in phase (ii) is implemented in *DynamiTE*. For simplicity, we only show the main abstract building blocks. Every gray block in Fig. 3 is implemented as already shown in Fig. 2. All the triples derived in the previous step are stored in memory.

The five blocks shown in Fig. 3 are executed in parallel. They only consider rules in which at least one literal in the body of the rule is instantiated from a triple that is stored in memory, i.e., a triple that is in δ , or that was derived from δ in a previous iteration.

The first block considers rules of *Type 1*, and reads schema triples from both the B-Trees and the memory. We split the execution of rules of *Type 3* into three blocks: one reads both schema and generic triples from the memory; one reads only schema triples from the memory and the generic triples from the B-Trees; the last one reads schema triples from the B-Trees and generic triples from the memory. This division enables us to reduce the amount of information to read from the B-Trees. Indeed, after reading a triple t from the memory, we can extract from the B-Tree only triples that can be joined with t to instantiate some rules.

Finally, the last block considers the generic triples in the in-memory update and executes rules of *Type 2*. After launching the execution of these blocks on different threads, *DynamiTE* waits for them to finish, removes the duplicates, and stores all (new) derivations in memory. It continues to iterate until no new derivation is produced.

5 Materialization After Data Removals

When removing a set of triples δ from a database \mathcal{J} , we also need to remove from the materialized view all the triples that cannot be derived from $\mathcal{J} \setminus \delta$. In this section, we describe the implementation of two algorithms, one already described in [10] (but only from a theoretical perspective) and another one that is based on the idea of counting all the direct derivations.

5.1 DRed Algorithm (and Derivatives)

The first algorithm implemented by *DynamiTE* is known as *Delete and Rederive (DRed)* and was first introduced in [6]. It works in two steps: (i) First, it computes all the triples that can be derived from δ , and removes them from the knowledge base. This process clearly computes an overestimation of the triples to remove, since some of them can have alternative derivations in $\mathcal{J} \setminus \delta$. Therefore, as a second step, (ii) the algorithm re-derives the triples that are still valid, and adds them again to the knowledge base.

Our implementation is based on a similar version of DRed, presented in [10], which has the advantage of using only the original set of rules for maintaining the materialization. More precisely, it implements the first phase (*Delete*) as a semi-naive evaluation that only considers rule evaluations in which at least one literal in the body of the rule is instantiated from a triple in δ (or derived from

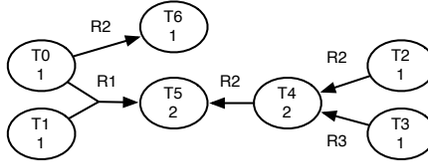


Fig. 4: Counting algorithm. An example of how the count attribute is computed. Nodes represent triples and include their name and their count. Arrows represent applications of rules and are labeled with the name of the rule.

δ in previous iterations). All the triples derived in this phase are removed from the knowledge base. The second phase (*Rederive*) is again implemented as a semi-naive evaluation, which considers all the triples left in the knowledge base.

Since we assume that the size of our update (and the derivations it produces) is small, we first load δ in memory. Then, we start executing the *Delete* phase: we do so by using the implementation of the semi-naive evaluation presented in Fig. 3. This means that we consider only derivations that involve triples in δ (or derived from δ in previous steps) and we store the output of the computation in memory, until we reach the fix-point.

After the first phase completes, we remove all the derivations stored in memory from the knowledge base, and we start the second, *Rederive* phase. Here too, we exploit the implementation of the semi-naive evaluation described above. This step outputs (and stores in main memory) all the triples that were removed from the knowledge base in the *Delete* phase, but that could actually be derived from $\mathcal{J} \setminus \delta$. As a final step, we add them again into our knowledge base.

5.2 Counting Algorithm

During the *Delete* phase, the previous algorithm computes all the triples that can be derived using triples in δ . Let us call this set T_δ . We call $D_\delta \subset T_\delta$ the subset of triples that cannot be derived from $\mathcal{J} \setminus \delta$, and $A_\delta = T_\delta \setminus D_\delta$. While only the triples in D_δ have to be removed from the knowledge base, the previous algorithm does not have enough information for recognizing them. For this reason, it removes all the triples in T_δ and then recalculates the set A_δ during the *Rederive* phase.

To improve the performance, we propose an alternative method, where all the triples are annotated with additional information that allows for immediate discrimination between the two sets A_δ and D_δ . This additional information consists of a new *count* attribute, which represents the number of possible rule instantiations that produced t as a *direct consequence*, plus one if the triple was also present in the original input.

As an example of how the count attribute is computed, consider Fig. 4. The figure shows a simple graph of derivations, where nodes represent triples and arrows represent applications of rules. For instance, T5 can be derived by applying rule R1 to T0 and T1. T0, T1, T2, and T3 are the facts in the original

input. They are stored in the knowledge base with count equal to one. T4 can be derived from rule R2 using T2, and from rule R3 using T3, so its count is two. Similarly, T5 can be derived using R1 from T0 and T1, and from T4 with R2, so its count is two. Finally, T6 can be derived from T0 and has count equal to one. Notice that we consider only *direct* derivations: although T4 can be derived in two ways, it participates in the count of T5 only once.

During the *Delete* phase, the presence of the count attribute enables us to discriminate between triples in D_δ and A_δ . As an example, consider again the graph in Fig. 4, and assume that we want to remove T0, i.e., $\delta = \{T0\}$.

We start a semi-naive evaluation to compute all the triples that can be derived from T0, i.e., T5 and T6, and we decrement their counts by one. All the triples whose count goes to zero (T6, in our example) do not have alternative derivations. They belong to D_δ , and can be removed from the knowledge base. On the other hand, T5 has count two in the materialized database: by removing T0, its count is decreased by one, but still there is one derivation left. This means that T5 is part of A_δ (i.e., it can still be derived from $\mathcal{J} \setminus \delta$), and should not be removed from the materialized database.

In this simple example, the semi-naive evaluation just required one iteration to reach the fix-point. If more iterations are needed the algorithm works as follows: at iteration n , it considers only rule instantiations that involve triples that were actually removed from the knowledge base at iteration $n - 1$, i.e., triples whose count went to zero at iteration $n - 1$.

Using this algorithm, the *Delete* phase computes *only* the triples that actually need to be removed from the knowledge base, i.e., the triples in D_δ . As a consequence, we can skip the *Rederive* phase.

Finally, notice that the counting algorithm requires the *count* attribute to be computed and maintained for each triple in the knowledge base. To do so, we implemented a slightly different version of the algorithms described in the previous sections for computing the (complete or incremental) materialization in case of data additions. In particular, after each derivation step, we never remove duplicates. Instead, every time we add a triple to the knowledge base, we check whether it was already present or not. If it is new, we add it with a count of one; otherwise, we increase its count by one.

6 Evaluation

Our evaluation has two goals: first, we want to test the absolute performance of *DynamiTE* when considering the computation of a full materialization and its maintenance in case of updates. Second, we want to compare the behavior of an existing state of the art algorithm for incremental update, namely DRed, with our own counting algorithm. To perform the experiments, we used one machine in the DAS-4 cluster¹, which is equipped with a dual quad-core Intel E5620

¹ <http://www.cs.vu.nl/das4>

Tasks	Time (sec.)
Compress Input	876
Create Indices	7953
Full Materializ.	886
Copy to Indices	5967

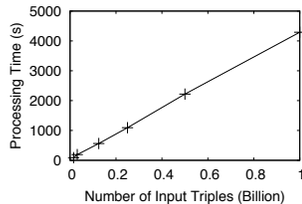


Fig. 5: (Left) Execution time for all the tasks during the loading of the database. (Right) Execution time of the complete closure.

CPU, 24GB of main memory, two hard disks of 1TB connected with RAID-0, and one 500GB SSD disk.²

Dynamite is fully written in Java³, and it uses BerkeleyDB Java Edition [12] as implementation of the on-disk B-Trees. We chose BerkeleyDB since it is among the most widely used databases, and it fully supports many functionalities such as transactions and concurrency. We use the LUBM [4] benchmark test to evaluate the performance of our system. We chose this dataset for two reasons: (i) it is one of the de-facto standard benchmarks to test the performance of reasoning on RDF data; (ii) it allows us to tune the experiments to control the amount of derivation that is produced.

We present three sets of experiments, with the following goals: (i) to evaluate the costs of performing the complete materialization; (ii) to evaluate the cost of the updates, and (iii) to discover the performance bottlenecks of our system.

Complete Materialization. Fig. 5 (left) shows the execution time required by each single task of the initial workflow (see Fig. 1) to materialize the LUBM(1000) dataset, which contains about 138 million triples. From this table, we notice that both compression and materialization take a relatively short time compared to the operation of copying the triples into the B-Tree indices. This last task is not related to reasoning, yet it is a necessary part in our workflow.

To better understand the performance of our system in computing the full materialization, we study this single operation in more detail. To this purpose, Fig. 5 (right) shows how the execution time for the full materialization changes with the number of triples present in the original input, starting from LUBM(125), consisting of 16 million triples, to LUBM(8000), of about one billion triples. We observe that our implementation has good scalability, since the execution time increases linearly with respect to the size of the input. Furthermore, the system has a high throughput: it computes the closure of about one billion LUBM triples in about 4400 seconds, which results in an input processing ratio of about 227K triples/sec. As an informal comparison, the throughput per machine of WebPIE [15] to materialize the same dataset with RDFS was about 55K triples/sec, four times lower than *DynamiTE*.

² In some experiments the machine had only a 256GB SSD disk.

³ The code is available online at <http://github.com/jrbrn/dynamite>

Incremental Updates. After the system has materialized the input knowledge base, the user can update it by removing or adding new triples. To evaluate the performance of this operation, we create six different type of updates:⁴.

- **Update 1.** Add/Remove one triple, which does not trigger any reasoning.
- **Update 2.** Add/Remove $\sim 16k$ triples, which do not trigger any reasoning.
- **Update 3.** Add/Remove $\sim 8k$ triples with the predicate `LUBM:emailAddress`. This update triggers reasoning so that a fixed number of triples is derived.
- **Update 4.** Remove the triple indicating that the property `LUBM:headOf` is a subproperty of `LUBM:worksFor`. This removal triggers a reasoning process that derives a number of triples proportional to the input size. To simulate the same reasoning for the addition, we add two new triples that indicate that `LUBM:headOf` is a subproperty of a new property `LUBM:responsibleOf`, which is also a subproperty of a new `LUBM:Manager`.
- **Update 5 and 6.** Add/Remove respectively one and two entire universities.

We chose Update 1 and Update 2 to evaluate the insertion cost and the overhead of checking that no reasoning can be applied. Update 3 represents a small update that produces only limited reasoning. Update 4 consists of schema information, which has a consistent impact on the knowledge base. Update 5 and Update 6 represent large updates, which trigger the execution of multiple rules. Notice that significantly larger updates are not possible since we store the update in main memory. For them, an incremental procedure might not be the best choice, and a complete re-computation would probably be more efficient.

Table 2 shows the execution time of these updates using the three algorithms explained above. We noticed a certain fluctuation in the runtime, so we repeated every measurement five times and report their average. First of all, we notice that the runtime for the addition ranges from 117 ms (insertion of one triple) to 31.8 s (addition of two universities, i.e., $\sim 250k$ triples). Even though these runtimes cannot always guarantee a real-time processing, they are significantly lower than recomputing a complete materialization. A significant result is represented by the results obtained in Update 1 and in Update 2. In both cases, no reasoning is triggered; however, Update 2 is significantly slower, since it requires *DynamiTE* to add 16k triples into the BTrees. Once again, this experiment demonstrates how accessing the B-Trees on disk represents the main cost for *DynamiTE*.

Considering the removal, we immediately observe that the DRed algorithm is very slow, with a runtime that is always larger than 30 minutes. This is due to the fact that the *Rederive* phase needs to access the entire input to re-compute possible conclusions that were incorrectly removed in the *Delete* phase. In contrast, our counting algorithm is much faster, with a runtime that ranges from 117 ms (removal of a triple) to 135 s (removal of two universities). We must remark that the procedure of enabling the counting slows down the initial preprocessing by about 49%⁵. However, the advantage obtained during a removal is so significant that this additional cost is quickly amortized after only a few updates.

⁴ For reproducibility, also these updates are available in the repository of the project.

⁵ This overhead is already included in the results presented in Fig. 5.

Update	Addition (sec.)	Removal (sec.)	
		DRed	Counting
Update 1	0.117	2902.7	0.117
Update 2	8.2	2049.6	25.7
Update 3	3.7	2121.9	25.4
Update 4	31.8	2132.2	51.0
Update 5	16.8	2196.0	74.6
Update 6	30.5	3830.2	135.8

Table 2: Runtime of four type of updates on 138 million triples (LUBM(1000)).

Performance Bottleneck. To further investigate the behavior of *DynamiTE*, and to identify the main performance bottlenecks, we launched the complete materialization and the incremental Update 4 changing two critical settings of the system: the kind of disk adopted and the number of threads used for processing.

First, we changed the disk storage from an SSD to a normal HDD. The full closure was only 6% slower, but the incremental addition and removal became 30 times and 15 times slower, respectively. This clearly indicates that the disk speed is a performance bottleneck for the B-Tree indices.

Second, we decreased the degree of parallelism by decreasing the number of concurrent processing threads from eight to four (the SSD disk was used for the storage). The runtime of the materialization became 12% slower, while the runtime of the incremental addition and removal became respectively at most 6% and 30% slower.

From these experiments, we can conclude that the degree of parallelism is an important component in shaping the performance, especially for the full materialization. However, disk throughput remains the largest performance bottleneck for the incremental updates, since the disk-based data structure heavily relies on it to retrieve the data.

7 Related Work

The problem of updating derived information upon changes in the knowledge base has been widely studied by the database community, in the context of view maintenance and deductive databases. More in particular, the idea of *incrementally* updating derived information has been studied since the beginning of the 1980s [5], leading to two main algorithms: DRed (Delete and Rederive) [6], and PF (Propagate Filter) [7]. Both algorithms share the same idea: when some base facts are removed, they first compute an overestimation of the derived knowledge that needs to be deleted, and then rederive the information that is still valid.

A declarative version of the DRed algorithm was first introduced in [14] and then extended in [17, 18] to consider also updates in the ruleset. These algorithms, however, create an update program that can be significantly larger than the original program (in terms of number of rules). Moreover, the update program can include negations, even if they are not present in the original program.

Our implementation of the DRed algorithm follows the work presented in [10], which overcomes the limitations listed above and manages incremental updates without changing the set of derivation rules. As we show in the evaluation, DRed has the disadvantage of always requiring to read full knowledge base.

Our counting algorithm is significantly more efficient than DRed, since it avoids (when possible) a complete scan over the input. This algorithm is based on the operation of counting and decrementing the number of possible derivations. This operation was also introduced in the original DRed paper [6], but it was not implemented nor designed for a declarative language.

Recently, new solutions for incremental materialization in the domain of stream reasoning were being proposed [3]. In particular, in [2], the authors proposed a novel algorithm and implemented it into the C-SPARQL execution engine. In this algorithm all data structures are stored in the main memory. The evaluation of this approach, based on the transitive property, proved that it is faster than DRed for updates that involve less than 13% of changes into the knowledge base. However, this algorithm is tailored for a specific application scenario (stream reasoning in C-SPARQL), and relies on some strong assumptions, e.g., that the expiration time of triples is known a-priori. In practice, this only happens if triples are observed through a *fixed* time window; unpredictable changes or other kinds of observation windows are not supported.

8 Conclusions

In this paper, we presented *DynamiTE*, a system designed to efficiently compute and maintain the materialization of a knowledge base in the presence of addition or removal of triples.

For data addition, *DynamiTE* implements a parallel version of the well-known semi-naive evaluation. For data removal, it implements two algorithms, one that is among the state of the art in the literature, and a more efficient one. *DynamiTE* is designed to exploit multi-core hardware for improved performance, adopting data structures that enable fast retrieval of information, e.g., to efficiently execute queries. Our evaluation shows the efficiency of *DynamiTE*, both in computing a complete materialization, and in managing incremental updates. Furthermore, it shows how the removal algorithm that we propose significantly outperforms existing state of the art approaches.

As future work, we plan to extend the algorithms implemented in *DynamiTE* to support different types of rules, and dynamic changes in the ruleset. Moreover, we will investigate how distributed processing can further increase the scalability.

To conclude, we have shown how our system encodes efficient parallel methods to perform full and incremental materialization. The throughput is higher than state of the art methods (per machine), while the response time to updates ranges from hundreds of milliseconds to a few minutes. This allows the system to perform a large-scale materialization on much more dynamic inputs than currently possible.

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