Keywords: Local bisimulation, Relevant set, A(k)-Index, Partial order relation.

Abstract: Local bisimilarity has been proposed as an approximate structural summary for XML and other semi-structured databases. Approximate structural summary, such as A(k)-Index and D(k)-Index, reduce the index's size (and therefore reduce query evaluation time) by compromising on the long path queries. We introduce the A(k)-Simplified and the A(k)-Relevant, approximate structural summaries for graph documents in general, and for XML in particular. Like A(k)-Index and D(k)-Index, our indexes are based on local bisimilarity, however, unlike the previous indexes, they support the removal of non-relevant nodes. We also describe a way to eliminate false drops that might occur due to nodes removal. Our experiments shows that A(k)-Simplified and A(k)-Relevant are much smaller then A(k)-Index, and give accurate results with better performance, for short relevant path queries.

1 INTRODUCTION

XML (eXtensible Markup Language) has rapidly become the data representation language of choice for information exchange on the web (Bray, T., et al.), (Busse, R., et al, 2001). XML document is a semi-structured database (Buneman, P., et al, 1997), (Li, Q. and Moon, B., 2001) , i.e. can be described as a directed graph. In Figure 1 we see an example of a graph representing a semi structured database. As with relational databases, evaluating queries using only the original database can be very costly in terms of time and resources needed. Clearly, an index is needed for an efficient evaluation. Several types of indexes were suggested, such as connection indexes, which are designed to support ancestor-descendent connection, like HOPI (Schenkel, R. et al., 2004), but most of the indexes proposed are based on structure summary of the document. Existing structure summaries are based on NFA to DFA transition like Strong DataGuide (Goldman, R. and Widom, J., 1997), graph index with hash tree to support common prefixes, like APEX (Chung, C. W. et al., 2002), and bisimilarity based summaries like 1-Index (Milo, T. and Suciu, D., 1999), (Kaushik, R. et al., 2003) , T-Index (Milo, T. and Suciu, D., 1999), F&B-Index (Kaushik, R. et al., 2002), A(k)-Index (Kaushik, R. et al.) and D(k)-Index (Qun, C., Lim A. and Ong, K. W., 2003).

In this paper we introduce A(k)-Simplified and A(k)-Relevant – a structure summary of a semi structured database, based on local bisimilarity, which supports removing nodes with non relevant tags (and therefore significantly decreasing the index size), while saving the document structure, so short queries can still be answered accurately. The main assumption our indexes is based upon is that we can classify the tags into 2 groups: relevant and non relevant (as mentioned in (Kaushik, R. et al., 2002),(Qun, C., Lim A. and Ong, K. W., 2003)). By declaring a tag as a relevant tag we either mean that the expression paths are built from relevant tags only – and we have A(k)-Simplified to support query evaluation, or, we mean that the results will be composed of relevant tags only – and we have A(k)-Relevant to support these queries. The proposed approximate indexes are much smaller then A(k)-Index, and therefore give much better processing time results. Additional contributions in this paper are:

1. Presenting a method to eliminate false drops in A(k)-Simplified and A(k)-Relevant (with respect to A(k)-Index) using partial order. This method increases the size of the indexes only by a tiny amount, and uses data saved on a small percent of the edges.

2. Showing experimentally that A(k)-Simplified and A(k)-Relevant reduce both the size of the index and the processing cost (compared to A(k)-Index). We also examine the indexes behavior when evaluating queries with length greater then k.

The idea of replacing irrelevant tags with the label “other”, as a method to reduce index size, appeared in this paper.
already both in (Milo, T. and Suciu, D., 1998) and (Kaushik, R. et al., 2002). However, both papers do not elaborate on the algorithmic implications of this idea, and in the context of A(k)-Index. This algorithmic elaboration is the main contribution of this paper (see also section 2).

The rest of the paper is organized as follows: We review existing work in section 2, then we elaborate on the data model and other related issues in section 3 — preliminaries. Afterwards we describe A(k)-Simplified and A(k)-Relevant in section 4, and then we give a way to remove false drops in section 5. We then conclude with experimental results (section 6) and further research (section 7).

2 RELATED WORK

In semi-structured databases, indexes are structural summaries, made to reduce time for query evaluation. Some of the indexes are approximate, which means that for some queries, verification is needed in order to remove false drops (by the term false drops we mean false positive — some of the index's results are incorrect). Examples for such indexes are: Approximate DataGuide (Goldman, R. and Widom, J., 1999), which is based on Strong DataGuide in which “similar” nodes are merged, APEX (Chung, C. W. et al., 2002) — An index that adjusts its structure using data mining strategies in order to efficiently evaluate queries with common suffixes, D(k)-Index (Qun, C., Lim A. and Ong, K. W., 2003), which uses different local bisimilarity values in order to emphasize important tags, and A(k)-Index. In our work we are interested in the A(k)-Index and local bisimilarity on which it is based. We say that two nodes are bisimilar if the two groups of incoming label path are similar. Every node in a bisimilarity based index is an equivalence class for the bisimilar relation. The first bisimilarity based index was 1-Index, which is safe and accurate but tends to be very large, especially with "complex" documents. A(k)-Index offers a way to reduce the index size by relaxing the bisimilarity demand for having all the incoming label paths, and declaring a new relation – local bisimilarity. Two nodes are k local bisimilar, if their two groups of incoming label paths, not longer than k, are similar. Local bisimilarity is based on the assumption that long queries are rare, and therefore we can build a structure that gives accurate results for short queries, while in long queries the results are approximate, and sometimes need verification (i.e. A(k)-Index is safe). Experiments show that A(k)-Index is much smaller then 1-Index.

In A(k)-Index both relevant and non-relevant tags gets the same treatment, because we define one k for all the index, in D(k)-Index we can emphasize relevant tags by giving the nodes with the desired tags different 'k' (local similarity) value, however, in spite the fact that some tags are known to be non relevant at all, D(k)-Index still includes all the nodes whether their tags are relevant or not.

As seen in (Kaushik, R. et al., 2002),(Milo, T. and Suciu, D., 1998) using partial data can improve evaluation time. The schema in (Milo, T. and Suciu, D., 1998) is based on simulation and is used for creating efficient regular path expressions, which improves DB scanning time. "other" edges are used as a replacement for unknown tag's edges which exist in the DB. However, the goal of the data structure in (Milo, T. and Suciu, D., 1998) differs from the goal of (Kaushik, R. et al., 2002) and ours. We use a bisimulation data structure to efficiently evaluate the query, rather then creating a more efficient one, both (Kaushik, R. et al., 2002) and us offer a method to reduce the index size, by replacing irrelevant tags with the tag "other", and removing isolated "other" nodes. This method allows an efficient evaluation of queries based on relevant tags only. (Kaushik, R. et al., 2002) uses this method to reduce the F&B index size, but does not give detailed algorithms for constructing the index and evaluating the query using "other". We present these algorithms in depth, especially in the context of A(k)-Index structure, and exploit the local bisimulation in treating "other" nodes. A(k)-Simplified uses this idea and adds a method to remove "other" node's path, and A(k)-Relevant exploits local bisimilarity and relaxes the demand for queries with relevant tags only, by just requiring a relevant tag at the end of the query and therefore supports all queries which return relevant data. Both indexes support the collapse of "other" paths without introducing false results.

3 PRELIMINARIES

XML or other semi-structured databases are modeled as a directed, labeled graph G, with each edge indicating an object-subobject or object-value relationship. Each node in G has a label, and an OID (as described in (Papakonstantinou, Y., 1995)), with simple objects having a distinguished label, VALUE. In Figure 1 we can see an example of a semi structured data graph. The id-idref edges are sketched with dotted line, but in our model, we consider all the edges in the same manner. Queries in semi structured databases are based on regular path expression. Let G be a data graph, and Σ₀ be
Definition 1. (Safe and Accurate) Let $R$ be a regular path expression, we say that an index $I_G$ is safe if for each node $v \in V_G$, if $R$ matches $v$, then $v$ is in the result of evaluating $R$ on $I_G$. We say that $I_G$ is accurate if for each node $v \in V_G$, matches $v$ if and only if $v$ is in the result of evaluating $R$ on $I_G$.

Definition 2. (Reversed Bisimulation) Let $G$ be a data graph in which the symmetric, binary relation $\approx$, the reversed bisimulation, is defined as : two nodes $u$ and $v$ are bisimilar ($u \approx v$) recursively:
1. $u = v$ iff $u$ and $v$ have the same label.
2. $u \approx v$ iff $u =_{k+1} v$ and for every parent $u'$ of $u$, there is a parent $v'$ of $v$, s.t. $u' =_{k} v'$, and vice versa.

If two nodes are bisimilar then all the incoming label paths are equal, while if two nodes are $k$-bisimilar, then all the incoming label paths not longer than $k$ are equal. $1$-Index and $A(k)$-Index are based on bisimulation and $k$-bisimilarity respectively. We note that $1$-Index is accurate while $A(k)$-Index is safe. In Figure 1 we see an example of $A(0)$-Index (label split index), $A(1)$-Index and $1$-Index for the data graph shown in Figure 1. Each index node has its extent attached.

A command is the basic instruction that a script file contains. Some commands require parameters that further define what the command should do. An expression is a combination of operators and arguments that create a result. Expressions can be used as values in any command. Examples of expressions include arithmetic, relational comparisons, and string concatenations.

3.1 The Set $L$

Normally we can split our DB into relevant and irrelevant data (with respect to the queries on the database). For example, in our movie database we might want to query only movies or actors so director and director’s name are irrelevant. We split the database by defining $L$ ($L \subseteq \Sigma_G$) to be the set of relevant tags in the database. We will see later in this paper that by exploiting the existence of $L$, we can reduce the index size and query evaluation time significantly. By defining $L$, we should consider two types of queries containing relevant tags:

1. Relevant queries. Queries built only on relevant tags. i.e. the query is a regular path expression $R$, the set of labels in $G$, we define a regular path expression $R$ as follows: $R := L \mid \_ \mid R \cdot R \mid (R \| R) \mid R^*$, where $L \subseteq \Sigma_G$, ‘$\_$’ denotes any label (one label only), ‘$\cdot$’ denotes sequence, ‘$\|$’ alternation and ‘$^*$’ denotes repetition. From now on, we will use a common abbreviation – the sign “$^*$” will replace “$\cdot \ldots \cdot$”. Before explaining the usage of regular path expressions, we give some formal definitions. A node path in $G$ is a sequence of nodes, $n_1, n_2, \ldots, n_m$, such that $n_0 \in G$, and an edge exists between $n_i$ and $n_{i+1}$ for $1 \leq i \leq m-1$. A label path is a sequence of labels $l_1, l_2, \ldots, l_n$, such that $n_0 \in G$, and an edge exists between $n_i$ and $n_{i+1}$ for $1 \leq i \leq m$. We denote $L(R)$ to be the regular language specified by $R$, and we say that $R$ matches a graph node $n$ if there is a label path for some word in $L(R)$ that matches a node path ending in $n$. Now, the result of evaluating a query $R$ on a data graph $G$, is the set of nodes in $G$ that matches $R$. For example, the path expression $\_\cdot$ movie.title evaluated on Figure 1 data graph, is all the movie’s titles.

Each node in the index graph is associated with an extent, which is the set of nodes in the matching equivalence class. If an edge exists from a node in one equivalence class to a node in another equivalence class, we add the appropriate edge to the index. We denote the index for data graph $G$ as $I_G$. Evaluating a query, which is a regular path expression $R$ on $I_G$, is a union of the extents of index nodes in $I_G$ that matches $R$.

<table>
<thead>
<tr>
<th>a</th>
<th>b</th>
<th>c</th>
<th>d</th>
</tr>
</thead>
<tbody>
<tr>
<td>Figure 1: Semi structured database represented as a graph, $1$-Index and $A(k)$-Index</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
R=r₁,r₂, ..., rₙ, and for i=1,...,n-1, rᵢ ∈ L ∪ {"*", (rᵢ₁|rᵢ₂)}, rₙ ∈ L.

2. Relevant end queries. Queries that end with a relevant tag. This means we have relevant tags as our results, but irrelevant tags can be a part of the queries. We define those queries as a regular path expression \( R = r_1.r_2. \ldots r_n \), and for \( i=1,\ldots,n-1 \), \( r_i \in L \cup \{ \ast, (r_i_1|r_i_2) \} \), \( r_n \in L \).

The symbol "_" was not mentioned, but since we are using a relevant set, it can be used in queries only in order to replace a relevant tag.

In this paper we will propose 2 indexes: A(k)-Simplified and A(k)-Relevant. These indexes exploit the existent of L in order to reduce index size and query evaluation time. A(k)-Simplified is very efficient when evaluating queries from the first type (relevant tags only), while A(k)-Relevant supports both type of queries, and is preferred when dealing with queries which contains irrelevant tags along the path. A(k)-Simplified and A(k)-Relevant are very efficient when L is defined, and the queries comply with one of the two types. If this is not the case, using these indexes may result in false drops. The term 'relevant' will be used both when discussing tags and nodes with relevant tags. The meaning will be clear from the context.

4 A(K)-SIMPLIFIED AND A(K)-RELEVANT

4.1 Introduction to A(k)-Simplified and A(k)-Relevant

A(k)-Simplified index is based on k-bisimilarity by the fact that we preserve all the incoming paths with length less or equal to k, furthermore, giving the set L, each tag in A(k)-Simplified is either from L or "other". A(k)-Simplified is built by replacing all the irrelevant tags by the tag "other", and then replacing an "other" tree with one "other" node. A(k)-Relevant is also based on k-bisimilarity, but in this index, besides the relevant tags and the tag "other", we preserve the tags of nodes that have a relevant descendant in distance not greater than k. A(k)-Relevant is also built by replacing all the irrelevant tags by the tag "other", and then replacing an "other" tree with one "other" node. A(k)-Simplified and A(k)-Relevant have the following properties (w.r.t A(k)-Index (Kaushik, R. et al.), (Qun, C., Lim A. and Ong, K. W., 2003)):

1. For relevant tags, which are our main (and usually all) interest, A(k)-Relevant keeps all paths with length ≤ k (and therefore accurate for those queries). Using false drops removal, A(k)-Simplified is also accurate for relevant queries with length ≤ k.
2. A(k)-Simplified and A(k)-Relevant are safe.

For k values large enough, A(k)-Simplified and A(k)-Relevant do not change (they have a steady structure) and they can be considered as 1-Index with relevant support.

4.2 Construction Algorithms

We first describe the A(k)-Simplified construction algorithm and then the modifications needed for building A(k)-Relevant. The construction algorithm has 4 steps:

1. **Collapse** – replacing irrelevant tags with the tag "other", and removing irrelevant nodes which have no relevant descendant.
2. **A(k)-Index** – creating an index structure where k-bisimilar nodes are combined to the same extent.
3. **MaxOtherPath** – marking the "other" trees.
4. **Replacing other trees** – replacing the trees marked in step 3 by one "other" node.

**Algorithm 1: Collapse**

**Input:** The inverted data graph G=(V,E).

**Output:** The data graph G with irrelevant tags replaced with the tag "other", and irrelevant nodes with no relevant descendant removed.

1. Empty (Q)
2. for each vertex v ∈ V(G) do
3. If v ∈ L (Relevant) then Distance(v) = 0;
   Status(v) = "Visited"; Enqueue(Q,v)
4. Else Distance(v) = ∞; Status(v) = "Not Visited"; Label(v) = "other"
5. While Q is not Empty
6. u = head(Q)
7. for each s son of u do
8. if Status(s) = "Not Visited" then
   Status(s) = "Visited"; Distance(s) = Distance(u) + 1; Enqueue(Q,s)
9. Dequeue(Q)
10. Status(u) = "Finished"
11. Remove all vertices with Distance = ∞ and Reverse G' back to G

We can see the collapse running example in Figure 2. We remark that the input is G’ (G reversed) because the distance is calculated from the descendents to their parents. We will also remark that in step 8 we save the distance from a irrelevant node to a relevant node. This is not important here (in A(k)-Simplified we are only interested to know whether it is ∞ distance (no path) or not). We will use this parameter later, when we describe A(k)-Relevant. The algorithm is based on BFS with some modifications – we traverse G’ – the reversed graph and initialize the next node queue with all the relevant nodes (the grey nodes in Figure 2 will become clearer when we describe A(k)-Relevant)

As stated earlier, the next step is applying A(k)-Index to Collapse’s result. The algorithm is described in (Kaushik, R. et al.). The third step is MaxOtherPath - this algorithm is based on depth first search, where for each “other” node we encounter – we either relate it to the current path (if exist) or start a new path. The motivation is to mark “other” nodes trees before replacing each one with a single node.

**Algorithm 2: MaxOtherPath**

**Input:** The result of the second step (A(k)-Index) – the index graph G.

**Output:** The index graph G with each “other” node related to an “other” tree. Each “other” tree has a set of all incoming and outgoing edges.

1. Global-Current-Tree = 0
2. For each vertex u ∈ V(G) Visited(u) = Never; Path(u) = Global-Current- Tree
3. MaxOtherPath-DFS-Visit(root, False, 0)
   MaxOtherPath-DFS-Visit(u, Path-Indication, Current-Tree)
   1. Visited(u) = Visited
   2. if u ∈ L (i.e. Relevant) then Tree-Indication = False
   3. Else
   4. If Tree-Indication = False then Path-Indication = True; Global-Current-Tree = Global-Current-Tree + 1; Current-Tree = Path(u) = Global-Current-Tree
   5. Else Path(u) = Current-Tree
   6. For each v ∈ Son(u)
   7. if Visited(v) = Never then
   8. MaxOtherPath-DFS-Visit(v, Path-Indication, Current-Tree)
   9. For each path found – walk through the nodes
     and for every edge not part of the path add it to incoming set or outgoing set according to the edge origin.

We can see an example of MaxOtherPath in Figure 3. The last step of the construction algorithm, is replacing each “other” nodes tree by one “other” node. This is done simply by removing all the “other” nodes and their edges and replacing them with one node with tag “other”. For every incoming and outgoing edge we saved in step 9 in MaxOtherPath, we add the appropriate edge to our new node (see Figure 3d). Note that if instead of replacing every “other” tree, we replace all the “other” nodes with one node, we may add many false drops, and this may cause a lot of verifications when we process the queries, and therefore substantially increase time for evaluating the queries. We will also note that self edges will be created when we have 2 (or more) “other” paths (paths built on “other” nodes only) from one “other”
node to another. For now, self edges can be omitted; we will discuss this in the next section – removing false drops.

Until now we discussed A(k)-Simplified building algorithm. As we have already seen, the difference between A(k)-Simplified and A(k)-Relevant is that the former replaces the tag of all the irrelevant nodes with the tag “other”, while the latter only replaces the tag of irrelevant nodes with distant relevant descendent (distance > k). In order to achieve this, we have to modify the first step (collapse). The modifications are: k will be added to the input for the algorithm, and instead of replacing all the irrelevant nodes, we will replace only those nodes which have distance > k. In Figure 2f, we marked with gray dots the irrelevant nodes which will not be tagged as “other”. If the data graph is G=(V,E), the total constructing time is O(k|E|) (A(k)-Index time complexity). In more details, we have the time complexity O(|E|) for collapse (BFS complexity), O(k|E|) (Kaushik, R. et al.) for A(k)-Index, O(|E|) for MaxOtherPath (DFS complexity) and O(|E|) for replacing other trees (removing irrelevant nodes, and restoring all the affected edges).

Theorem 1. A(k)-Simplified is safe for l ≤ k relevant queries. A(k)-Relevant is safe and accurate for l ≤ k relevant end queries.

Proof. Obvious from the indexes definitions.

5 REMOVING FALSE DROPS

As mentioned earlier, replacing the “other” trees may result in increasing number of false drops (w.r.t. A(k)-Index). In Figure 4 we can see that replacing the irrelevant tree increases the number of false drops in A(k)-Simplified for paths with length < k (in Figure 4b the dotted line shows the path which does not exist in the original graph). A(k)-Relevant is accurate for short relevant end queries, however, in Figure 4 we see an example where for path longer than k, A(k)-Relevant adds false drops which are not returned by a lookup in the A(k)-Index. In Figure 4c the dotted line marks the result of the query R.A.*.B. In Figure 4d we see the result of this query on A(4)-Relevant, which includes one false drop.

The reason for increasing number of false drops w.r.t. A(k)-Index is that we replace the “other” trees with one node and add paths which do not exist in the original graph. From these observations we see that removing invalid paths involving “other” nodes can reduce the number of false drops for A(k)-Simplified (for queries in any length) and for A(k)-Relevant (query’s length > k). In this section we will present a method to remove these invalid paths. For each “other” tree replaced by one node, we mark the location of every incoming and outgoing edge, and use these marks in query calculation. We now describe a way to mark the edges using a partial order relation.

5.1 Partial Order Relation

The intuition for using partial order relation is the observation that it stands for the relation “descendent of” between 2 nodes in an “other” tree discovered by collapse, and therefore, by keeping the partial order relation between the nodes we can reconstruct the original paths, and remove the false paths.

We save the location of every node in each “other” tree in the following way: p₁;p₂;...;pₖ;d (n>0). pᵢ represents the path from the root to the node, d represents the node’s depth. By the term “other” tree we mean the sequence of “other” nodes discovered during the MaxOtherPath algorithm. The meaning of pᵢ is: We mark the first “other” node discovered in the “other” path (the path’s root) by ;1 which stands for depth 1 (d=1, i=0). Assuming we are in an “other” node u which is marked with p₁;p₂;...;pₖ;d then:
1. If u does not have an “other” child node there is nothing to do.
2. If u has exactly one “other” child node, it will be marked with p₁;p₂;...;pₖ;d+1
3. If u has k>1 “other” children nodes, their mark will be: p₁;p₂;...;pₖ;d;1;d+1, ...
We can now define our partial order relation. Assume that $A=a_1; a_2; \ldots; a_n; d_a$, $B=b_1; b_2; \ldots; b_n; d_b$ ($A, B$ represent two locations in an "other" path), we define the relation $\leq$: $A \leq B$ if $a_1; a_2; \ldots; a_n \subset b_1; b_2; \ldots; b_n$ ($a_1; a_2; \ldots; a_n$ is not equal but is a prefix of $b_1; b_2; \ldots; b_n$) or if $a_1; a_2; \ldots; a_n = b_1; b_2; \ldots; b_n$ and $d_a \leq d_b$. This is a partial order relation – it is reflexive, antisymmetric and transitive. We will see later on, that given 2 nodes $u, v$ in an "other" tree there are three possibilities, either there is a path from $u$ to $v$ (which translates to $\text{Mark}(u) \leq \text{Mark}(v)$), a path from $v$ to $u$ ($\text{Mark}(v) \leq \text{Mark}(u)$) or there is no path at all (no relation between the $\text{Mark}(u)$ and $\text{Mark}(v)$). After defining the relation $\leq$, we should mark the "other" nodes and take this mark into consideration when querying the index. In MaxOtherPath, when we traverse an "other" tree we keep two parameters: trail and depth, and we mark each node with trail;depth, the trail corresponds to $p_1, p_2, \ldots, p_k$ and the depth corresponds to $d$. When replacing the "other" nodes tree with one "other" node we give each ingoing/outgoing edge two values: ingoing mark and outgoing mark. If the origin of the edge is from a relevant node (not an "other" node) its origin mark will be null, else, its origin mark will be the "other" node’s mark. We treat the node’s destination mark with a same manner. In Figure 5 we can see an example of how we mark each node in the "other" tree, and how we use those marks to mark the origin and destination of the appropriate edges.

5.2 Handling Self Edges

Both in $A(k)$-Simplified and $A(k)$-Relevant construction algorithms, self edges could have been formed when there where 2 (or more) "other" paths (paths built on "other" nodes only) from one "other" node to another. Those self edges were removed because they did not contribute to query analysis. However, when we add the false drops treatment, self edges play an important role. More information regarding self edges and self edges removal is given in (Biber, P. and Gudes, E., 2005).

Theorem 2. False drops removal eliminates the invalid paths formed when "other" tree is replaced by one "other" node.

Proof. In order to show that no invalid path is added, we will see that there is a path from a node $u$ to a node $v$ (which translates to $\text{Mark}(u) \leq \text{Mark}(v)$), a path from $v$ to $u$ ($\text{Mark}(v) \leq \text{Mark}(u)$) or there is no path at all (no relation between the $\text{Mark}(u)$ and $\text{Mark}(v)$). After defining the relation $\leq$, we should mark the "other" nodes and take this mark into consideration when querying the index. In MaxOtherPath, when we traverse an "other" tree we keep two parameters: trail and depth, and we mark each node with trail;depth, the trail corresponds to $p_1, p_2, \ldots, p_k$ and the depth corresponds to $d$. When replacing the "other" nodes tree with one "other" node we give each ingoing/outgoing edge two values: ingoing mark and outgoing mark. If the origin of the edge is from a relevant node (not an "other" node) its origin mark will be null, else, its origin mark will be the "other" node’s mark. We treat the node’s destination mark with a same manner. In Figure 5 we can see an example of how we mark each node in the "other" tree, and how we use those marks to mark the origin and destination of the appropriate edges.

5.3 Querying the Indexes

Without false drops treatment, the way to compute a query is equal to the one described in (Kaushik, R. et al.). In Figure 6a we can see an example of an index (it can be Relevant or Simplified), a DFA that matches the query *.A.B (Figure 6b), and the results of the index scan (Figure 6c) without taking the marks into consideration. The returned set is the extent of the index nodes that get an accepting state (state 3, marked with an arrow). In order to take advantage of the false drops removal’s markings added to the index, the scanning must consider two
factors: the edges marking and the self edges. In Figure 6d we can see the results of the same query evaluated with marks. In Figure 4c when the nodes C6, C7, C1 were translated to “other” nodes, they got the marking: ;1, ;1;1;2 and ;1;2;2 respectively, so in Figure 4d the edge between A and O has a destination mark 1;1;2, the edge from O to C2 has an origin mark 1;1;2 and the edge from O to C8 an origin mark 1;2;2. Now, since 1;1;2 has no ≤ relation with 1;2;2 when we process the query R.A.*.B and enter the “other” node from A, we will not be able to go to C8 and the false drop will be removed.

6 EXPERIMENTAL STUDY

In this section we present the results of the conducted experiments. This section layout is as follows: first we give some background regarding queries, the cost model, the selection of the relevant set L and the databases used in the experiments, next we examine the indexes size and the increase of size due to false drops removal, afterwards we compare the time needed to scan the indexes when processing queries and in the last section we examine the number of false drops.

6.1 Experimental Framework

Query classification

In our experiments we use a slightly different classification than the one presented in (Kaushik, R. et al.). We split the queries into 3 groups:

- **Short queries** which are queries built on labels only – for example L1.L2.L3. It is clear that for short and - relevant or relevant end query - A(k)-Simplified with false drops removal and A(k)-Relevant (respectively) are accurate.

- **Simple queries** which are queries that have only one “*” located at the beginning of the queries – for example *.L1.L2, although when processing these queries we use paths longer then k, it can be easily shown that the indexes are still accurate.

- **Complex queries** which have more then one “*” or only one “*” but not as the query’s starting label. When processing these queries all of the indexes (including A(k)-Index) may return false drops, so verification might be needed.

The experiments results include false drops verification (whenever it is need). As expected, verification is needed for complex queries, or when k is very small.

The cost model

We will use the cost model offered in (Kaushik, R. et al.). The cost of evaluating a query is the number of nodes visited in the index during automaton execution plus the number of nodes visited in the DB during false drops verification. Note that for precise queries no verification is needed. The graph size is calculated as the sum of the graph nodes and edges.

Selecting the relevant set L

In our experiments, we randomly selected various sets of relevant labels. There are two ways to examine the relevant set: the percent of nodes out of all the data graph nodes which have the relevant tags or, the percent of the relevant tags out of all the tags in the database. Our experiments show that the former is a better measure to predict the indexes size and accuracy. We also randomly selected the relevant set, so usually the nodes will be spread all over the graph, however, in some cases the nodes will not be spread homogeneously so the results can have some variation, but the experiments always show a tendency.

The Databases

In the experiments described, two XML databases were used. The first source is the Internet Movie Database (IMDB) (The Internet Movie Database Ltd, “Internet Movie Database”, Available: http://www.imdb.com) and the second is a synthetic database generated for performing our experiments. IMDB was selected because it represents a real life database with many cycles (Goldman, R. and Widom, J., 1997). The portion of the database selected is organized around movie. The synthetic database is a highly complex database with a lot of irregularity that allows examining the indexes in highly irregular conditions, where the index size is usually very large.
6.2 Index Size

The size of A(k)-Simplified and A(k)-Relevant increases as k increases but it remains stable compared to the A(k)-Index’s size. High variation of the database increases the size of A(k)-Simplified and A(k)-Relevant (compared to A(k)-Index). The size of the indexes w.r.t. A(k)-Index is displayed in Figure 7. The size is calculated for different values of k, the A(k)-Simplified and A(k)-Relevant are calculated for different L sets. The size shown in the graph is the indexes average size over different L sets, and as we can see A(k)-Simplified is 40%-52% of A(k)-Index, and A(k)-Relevant is 55%-69% of A(k)-Index. The increase of size due to false drops removal (because of self edges addition) is displayed in the lower figure. As expected, the increase of A(k)-Relevant’s size is almost zero for small k values and zero for large k values – since we preserve the tags of incoming paths with length smaller or equal to k, the size and complexity of the “other” paths is reduced and therefore the number of self edges is also reduced. In A(k)-Simplified the size increases for larger k, because in the second step of the building algorithm (A(k)-Index) fewer “other” nodes merge into one index node, and so in the fourth step (replacing the “other” trees) more nodes are replaced and so more self edges are formed. This trend stops at a certain k value.

6.3 Query Cost

Figure 8 shows the cost of evaluating various queries. As expected, the queries evaluation cost for A(k)-Simplified and A(k)-Relevant with or without false drops removal is much smaller than evaluating the query on A(k)-Index. Also as k reaches a certain value, the evaluation cost stops growing, since the indexes reach their stable state. As mentioned earlier, the result here are the index scanning cost plus verification cost (when needed). For complex queries, there is always a need to verify the query results (for all the queries). Since complex queries, like simple queries, contain “*” (one or more) the proportion between the A(k)-Index scanning time to the A(k)-Simplified and A(k)-Relevant scanning time remains quite similar to those of the simple queries evaluation. The important factor here is the number of false drops which causes costly verification. Figure 9 displays the percent of false drops for complex queries. It is clear that A(k)-Simplified without false drops removal has more false drops (and therefore has a costly evaluation time). With false drops removal the number of false drops in A(k)-Simplified is reduced by an order of tens of percents. As expected, for A(k)-Relevant the increase of false drops number (w.r.t. A(k)-Index) is very small – less than 0.3% in average (actually it is 0% most of the time) – here the false drops removal is less effective though we can still see a small improvement in the synthetic database. We can also see that for relevant end queries, A(k)-Relevant has much less false drops than A(k)-Simplified, which explains A(k)-Relevant’s better results.
7 CONCLUSION AND FUTURE RESEARCH

A(k)-Simplified and A(k)-Relevant are small, efficient indexes evolved from A(k)-Index. The indexes offer a way to reduce index size significantly when prior knowledge of the relevant tags exists. Though based on A(k)-Index, A(k)-Simplified and A(k)-Relevant are flexible, and can be modified to be based on other structured indexes, such as 1-Index and D(k)-Index. A(k)-Simplified is very efficient when queries are known to be based on relevant tags only, while A(k)-Relevant gives good results when only the ending tags are known. Performance examination shows that the indexes give good results when only the ending tags are known. More work is needed to modify existing indexes for supporting irrelevant node’s removal. It seems appropriate to adjust bisimulation based indexes to support exploiting relevant set declaration, and removing irrelevant nodes. Further work is needed to make A(k)-Simplified and A(k)-Relevant support DB functionality such as update, insert and delete.

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REFERENCES


