

Structural Joins: A Primitive for Efficient XML Query Pattern Matching

Shurug Al-Khalifa

Univ of Michigan
shurug@eecs.umich.edu

H. V. Jagadish

Univ of Michigan
jag@eecs.umich.edu

Nick Koudas

AT&T Labs–Research
koudas@research.att.com

Jignesh M. Patel

Univ of Michigan
jignesh@eecs.umich.edu

Divesh Srivastava

AT&T Labs–Research
divesh@research.att.com

Yuqing Wu

Univ of Michigan
yuwu@eecs.umich.edu

Abstract

XML queries typically specify patterns of selection predicates on multiple elements that have some specified tree structured relationships. The primitive tree structured relationships are parent-child and ancestor-descendant, and finding all occurrences of these structural relationships in an XML database is a core operation for XML query processing.

In this paper, we develop two families of structural join algorithms for this task: *tree-merge* and *stack-tree*. The tree-merge algorithms are a natural extension of traditional merge joins and the recently proposed multi-predicate merge joins, while the stack-tree algorithms have no counterpart in traditional relational join processing. We present experimental results on a range of data and queries using (i) the TIMBER native XML query engine built on top of SHORE, and (ii) a commercial relational database system. In all cases, our structural join algorithms turn out to be vastly superior to traditional traversal-style algorithms. Structural join algorithms are also much more efficient than using traditional join algorithms, implemented in relational databases, for the same task. Finally, we show that while, in some cases, tree-merge algorithms can have performance comparable to stack-tree algorithms, in many cases they are considerably worse. This behavior is explained by analytical results that demonstrate that, on sorted inputs, the stack-tree algorithms have worst-case I/O and CPU complexities linear in the sum of the sizes of inputs and output, while the tree-merge algorithms do not have the same guarantee.

1 Introduction

XML employs a tree-structured model for representing data. Quite naturally, queries in XML query languages (see, e.g., [10, 7, 6]) typically specify patterns of selection predicates on multiple elements that have some specified tree structured relationships. For example, the XQuery path expression:

```
book[title = 'XML']//author[. = 'jane']
```

matches `author` elements that (i) have as content the string value “jane”, and (ii) are descendants of `book` elements that have a child `title` element whose content is the string value “XML”. This XQuery path expression can be represented as a node-labeled tree pattern with elements and string values as node labels.

Such a complex query tree pattern can be naturally decomposed into a set of basic parent-child and ancestor-descendant relationships between pairs of nodes. For example, the basic structural relationships corresponding to the above query are the ancestor-descendant relationship (`book`, `author`) and the parent-child relationships (`book`, `title`), (`title`, `XML`) and (`author`, `jane`). The query pattern can then be matched by (i) matching each of the binary structural relationships against the XML database, and (ii) “stitching” together these basic matches.

Finding all occurrences of these basic structural relationships in an XML database is clearly a core operation in XML query processing, both in relational implementations of XML databases, and in native XML databases. There has been a great deal of work done on how to find occurrences of such structural relationships (as well as the query tree patterns in which they are embedded) using relational database systems (see, for example, [14, 28, 27]), as well as using native XML query engines (see, for example, [21, 23, 22]). These works typically use some combination of indexes on elements and string values, tree traversal algorithms, and join algorithms on the edge relationships between nodes in the XML data tree.

More recently, Zhang et al. [30] proposed a variation of the traditional merge join algorithm, called the multi-predicate merge join (MPMGJN) algorithm, for finding all occurrences of the basic structural relationships (they refer to them as containment queries). They compared the implementation of containment queries using native support in two commercial database systems, and a special purpose inverted list engine based on the MPMGJN algorithm. Their results showed that the MPMGJN algorithm could outperform standard RDBMS join algorithms by more than an order of magnitude on containment queries. The key to the efficiency of the MPMGJN algorithm is the (`DocId`, `StartPos` : `EndPos`, `LevelNum`) representation of positions of XML elements, and the (`DocId`, `StartPos`, `LevelNum`) representation of positions of string values, that succinctly capture the *structural relationships* between elements (and string values) in the XML database (see Section 2.3 for details about this representation). Checking that structural relationships in the XML tree like ancestor-descendant and parent-child (corresponding to containment and direct containment relationships, respectively, in the XML document representation) are present between elements amounts to checking that certain *inequality conditions* hold between the components of the positions of these elements.

While the MPMGJN algorithm outperforms standard RDBMS join algorithms, we show in this paper that it can perform a lot of *unnecessary* computation and I/O for matching basic structural

relationships, especially in the case of parent-child relationships (or, direct containment queries). *In this paper, we take advantage of the (DocId, StartPos : EndPos, LevelNum) representation of positions of XML elements and string values to devise novel I/O and CPU optimal join algorithms for matching structural relationships (or, containment queries) against an XML database.*

Since a great deal of XML data is expected to be stored in relational database systems (all the major DBMS vendors including Oracle, IBM and Microsoft are providing system support for XML data), our study provides evidence that RDBMS systems need to augment their suite of physical join algorithms to include structural joins to be competitive on XML query processing. The alternative approach (used by commercial products such as Oracle InterMedia Text and IBM DB2 text extenders) of loosely coupling an IR engine with the database engine can result in sub-optimal query evaluation plans, in addition to other problems with locking, concurrency control and recovery [30]. Our study is equally relevant for native XML query engines, since structural joins provide for an efficient set-at-a-time strategy for matching XML query patterns, in contrast to the node-at-a-time approach of using tree traversals.

1.1 Outline and Contributions

We begin by presenting background material (data model, query patterns, basic structural relationships, positional representations of XML elements and string values) in Section 2. Our main contributions are as follows:

- We develop two families of join algorithms to perform matching of the parent-child and ancestor-descendant structural relationships efficiently: *tree-merge* and *stack-tree* (Section 3). Given two input lists of tree nodes, each sorted by (DocId, StartPos), the algorithms compute an output list of sorted results joined according to the desired structural relationship. The tree-merge algorithms are a natural extension of merge joins and the recently proposed multi-predicate merge join algorithm [30], while the stack-tree algorithms have no counterpart in traditional relational join processing.
- We present an analysis of the tree-merge and the stack-tree algorithms (Section 3). The stack-tree algorithms are I/O and CPU optimal (in an asymptotic sense), and have worst-case I/O and CPU complexities linear in the sum of sizes of the two input lists and the output list for both ancestor-descendant (or, containment) and parent-child (or, direct containment) structural relationships. The tree-merge algorithms have worst-case quadratic I/O and CPU complexities, but on some natural classes of structural relationships and XML data, they have linear complexity as well.
- We show experimental results on a range of data and queries using (i) the TIMBER native XML query engine built on top of SHORE, and (ii) a commercial relational database system

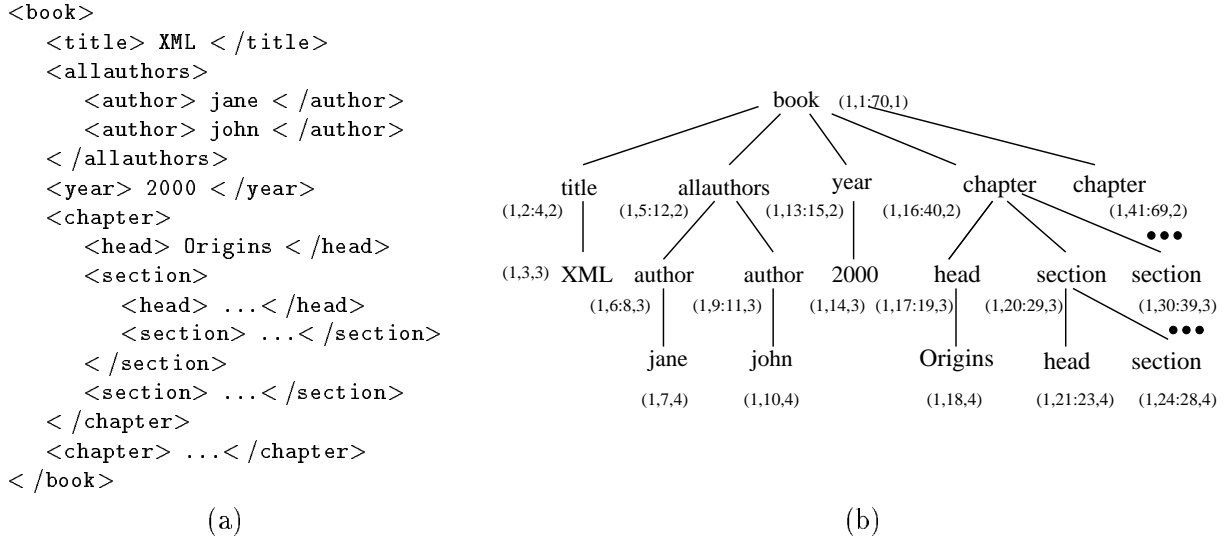


Figure 1: (a) A sample XML document fragment, (b) Tree representation

(Section 4).

- In all cases, our structural join algorithms turn out to be vastly superior to traditional traversal-style algorithms.
- Structural join algorithms are also much more efficient than using traditional join algorithms, implemented within relational databases, for the same task.
- We show that while, in some cases, the performance of tree-merge algorithms can be comparable to that of stack-tree algorithms, in many cases they are considerably worse. This is consistent with the analysis presented in Section 3.

We describe related work in Section 5, and discuss ongoing and future work in Section 6.

2 Background and Overview

2.1 Data Model and Query Patterns

An XML database is a forest of rooted, ordered, labeled trees, each node corresponding to an element and the edges representing (direct) element-subelement relationships. Node labels consist of a set of (attribute, value) pairs, which suffices to model tags, PCDATA content, etc. Sibling nodes (children of the same parent node) are ordered. The ordering of sibling nodes implicitly defines a total order on the nodes in a tree, obtained by a depth-first, left-right traversal of the tree nodes. For the sample XML document of Figure 1(a), its tree representation is shown in Figure 1(b). (The utility of the numbers associated with the tree nodes will be explained in Section 2.3.)

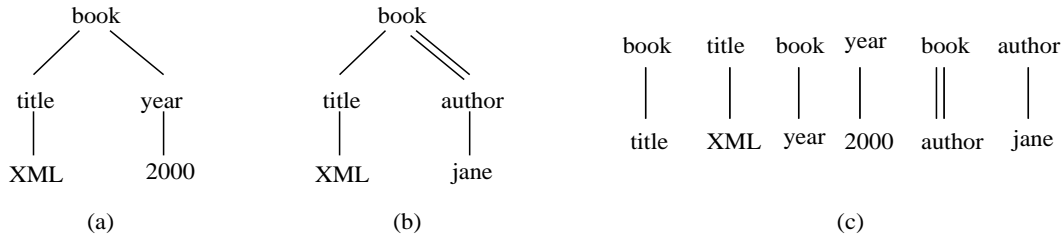


Figure 2: (a) and (b) Rooted Tree Patterns, (c) Basic Structural Relationships

Queries in XML query languages like XQuery [6], Quilt [7], and XML-QL [10] make fundamental use of (node labeled) tree patterns for matching relevant portions of data in the XML database. The query pattern node labels include element tags, attribute-value comparisons, and string values, and the query pattern edges are either parent-child edges (depicted using single line) or ancestor-descendant edges (depicted using a double line). For example, the XQuery path expression:

```
book[title = 'XML' AND year = '2000']
```

which matches `book` elements that (i) have a child `title` element with content `XML`, and (ii) have a child `year` element with content `2000`, can be represented as the rooted tree pattern in Figure 2(a). Only parent-child edges are used in this case. This query pattern would match the document in Figure 1. Similarly, the XQuery path expression:

```
book[title = 'XML']//author[. = 'jane']
```

which matches `author` elements that (i) have content `jane`, and (ii) are descendants of `book` elements that have a child `title` element with content `XML`, can be represented as the rooted tree pattern in Figure 2(b). Note that an ancestor-descendant edge is needed between the `book` element and the `author` element. Again, this query pattern would match the document in Figure 1.

In general, at each node in the query tree pattern, there is a *node predicate* that specifies some predicate on the attributes (e.g., tag, content) of the node in question. For the purposes of this paper, exactly what is permitted in this predicate is not material. It suffices for our purposes that there be the possibility of constructing efficient access mechanisms (such as index structures) to identify the nodes in the XML database that satisfy any given node predicate.

2.2 Matching Basic Structural Relationships Using Joins

A complex query tree pattern can be decomposed into a set of basic binary structural relationships such as parent-child and ancestor-descendant between pairs of nodes. The query pattern can then be matched by (i) matching each of the binary structural relationships against the XML database, and (ii) “stitching” together these basic matches. For example, the basic structural relationships corresponding to the query tree patterns of Figures 2(a) and 2(b) are shown in Figure 2(c).

Finding all occurrences of these basic structural relationships in an XML database (like that in Figure 1) is clearly a core operation in XML query processing, both in relational implementations of XML databases, and in native XML databases. There has been a great deal of work done on how to find occurrences of such structural relationships (as well as the query tree patterns in which they are embedded) using relational database systems (see, for example, [14, 28, 27]), as well as using native XML query engines (see, for example, [21, 23, 22]). These works typically use some combination of indexes on elements and string values, tree traversal algorithms, and join algorithms on the edge relationships between nodes in the XML data tree.

A straightforward approach to matching structural relationships against an XML database is to use traversal-style algorithms by using either child-pointers or parent-pointers. Such “tuple-at-a-time” processing strategies are known to be inefficient compared to the set-at-a-time strategies used in database systems. Pointer-based joins [29] have been suggested as a solution to this problem in the context of object-oriented databases, and shown to be quite efficient.

In the context of XML databases, nodes may have a large number of children, and the query pattern often requires matching ancestor-descendant structural relationships (for example, the `(book, author)` edge in the query pattern of Figure 2(b)), in addition to parent-child structural relationships. In this case, there are two options: (i) explicitly maintaining only (parent, child) node pairs and identifying (ancestor, descendant) node pairs through repeated joins; or (ii) explicitly maintaining (ancestor, descendant) node pairs. The former approach would take too much query processing time, while the latter approach would use too much (quadratic) space. In either case, using pointer-based joins is likely to be infeasible.

The key to an efficient, uniform mechanism for set-at-a-time (join-based) matching of structural relationships is a positional representation of occurrences of XML elements and string values in the XML database, which we discuss next.

2.3 Representing Positions of Elements and String Values in an XML Database

The classic inverted index data structure in information retrieval [26] maps a word (or string value) to a list of `(DocId, StartPos)` pairs, where the word occurs at position `StartPos` in the document identified by `DocId`. In order to process documents with a nested structure such as XML, the inverted index does not suffice since `StartPos` in a document alone cannot capture the structural relationships between elements and string values. However, there are elegant extensions that do suffice.

In one such extension [8, 9, 30], the position of an element occurrence in the XML database can be represented as the 3-tuple `(DocId, StartPos : EndPos, LevelNum)`, and the position of a string occurrence in the XML database can be represented as the 3-tuple `(DocId, StartPos, LevelNum)`, where (i) `DocId` is the identifier of the document; (ii) `StartPos` and `EndPos` can be

generated by counting word numbers from the beginning of the document with identifier `DocId` until the start of the element and end of the element, respectively;¹ and (iii) `LevelNum` is the nesting depth of the element (or string value) in the document. Figure 1(b) depicts a 3-tuple with each tree node, based on this representation of position. (The `DocId` for each of these nodes is chosen to be 1.)

Structural relationships between tree nodes (elements or string values) whose positions are recorded in this fashion can be determined easily, as discussed below. For simplicity of exposition, we shall refer uniformly to the `(DocId, StartPos : EndPos, LevelNum)` representation for both (non-leaf) elements and (leaf-level) strings; for strings `EndPos` is the same as `StartPos`.

Ancestor-Descendant: A tree node n_2 whose position in the XML database is encoded as $(D_2, S_2 : E_2, L_2)$ is a descendant of a tree node n_1 whose position is encoded as $(D_1, S_1 : E_1, L_1)$ iff (i) $D_1 = D_2$; and (ii) $S_1 < S_2$ and $E_2 < E_1$. For example, the author node with position $(1, 9 : 11, 3)$ is a descendant of the book node with position $(1, 1 : 70, 1)$ in Figure 1(b).

Parent-Child: A tree node n_2 whose position in the XML database is encoded as $(D_2, S_2 : E_2, L_2)$ is a child of a tree node n_1 whose position is encoded as $(D_1, S_1 : E_1, L_1)$ iff (i) $D_1 = D_2$; (ii) $S_1 < S_2$ and $E_2 < E_1$; and (iii) $L_1 + 1 = L_2$. For example, the string “john” with position $(1, 10, 4)$ is a child of the author element with position $(1, 9 : 11, 3)$ in Figure 1(b).

A key point worth noting about this representation of node positions in the XML data tree is that checking an ancestor-descendant structural relationship is as easy as checking a parent-child structural relationship. The reason is that one can check for an ancestor-descendant structural relationship without knowledge of the intermediate nodes on the path. Also worth noting is that this representation of positions of elements and string values allow for checking order and proximity relationships between elements and/or string values; this issue is not explored further in our paper.

2.4 An Overview

In the rest of this paper, we take advantage of the `(DocId, StartPos : EndPos, LevelNum)` representation of positions of XML elements and string values to (i) devise novel, I/O and CPU optimal (in an asymptotic sense) join algorithms for matching basic structural relationships (or,

¹We would like to remark that `(DocId, StartPos : EndPos, LevelNum)` is not the only representation of positions of elements and string values that allows for direct checking of structural relationships. An alternative representation is the 4-tuple `(DocId, PrePos, PostPos, LevelNum)`, where `PrePos` is the number given to a tree node in a pre-order traversal of the tree, and `PostPos` is the number given to a tree node in a post-order traversal of the tree.

```

Algorithm Tree-Merge-Anc (AList, DList)
/* Assume that all nodes in AList and Dlist have the same DocId */
/* AList is the list of potential ancestors, in sorted order of StartPos */
/* DList is the list of potential descendants in sorted order of StartPos */

begin-desc = DList->firstNode; OutputList = NULL;
for (a = AList->firstNode; a != NULL; a = a->nextNode) {
  for (d = begin-desc; (d != NULL && d.StartPos < a.StartPos); d = d->nextNode) {
    /* skipping over unmatchable d's */
    begin-desc = d;
    for (d = begin-desc; (d != NULL && d.EndPos < a.EndPos); d = d->nextNode) {
      if ((a.StartPos < d.StartPos) && (d.EndPos < a.EndPos)
          [&& (d.LevelNum = a.LevelNum + 1)]) {
        /* the optional condition is for parent-child relationships */
        append (a,d) to OutputList; }
    }
  }
}

```

Figure 3: Algorithm Tree-Merge-Anc with Output in Sorted Ancestor/Parent Order

containment queries) against an XML database; (ii) present an analysis of these algorithms; and (iii) show their behavior in practice using a variety of experiments.

The task of matching a complex XML query pattern then reduces to that of evaluating a join expression with one join operator for each binary structural relationship in the query pattern. Different join orderings may result in different evaluation costs, as usual. Finding the optimal join ordering is outside the scope of this paper, and is the subject of future work in this area.

3 Structural Join Algorithms

In this section, we develop two families of join algorithms for matching parent-child and ancestor-descendant structural relationships efficiently: *tree-merge* and *stack-tree*, and present an analysis of these algorithms.

Consider an ancestor-descendant (or, parent-child) structural relationship (e_1, e_2) , for example, (book, author) (or, (author, jane)) in our running example. Let $AList = [a_1, a_2, \dots]$ and $DList = [d_1, d_2, \dots]$ be the lists of tree nodes that match the node predicates e_1 and e_2 , respectively, each list sorted by the (DocId, StartPos) values of its elements. There are a number of ways in which the Alist and the Dlist could be generated from the database that stores the XML data. For example, a native XML database system could store each element node in the XML data tree as an object with the attributes: ElementTag, DocId, StartPos, EndPos, and LevelNum. An index could be built across all the element tags, which could then be used to find the set of nodes that match a given element tag. The set of nodes could then be sorted by (DocId, StartPos) to produce the lists that serve as input to our join algorithms.

```

Algorithm Tree-Merge-Desc (AList, DList)
/* Assume that all nodes in AList and DList have the same DocId */
/* AList is the list of potential ancestors, in sorted order of StartPos */
/* DList is the list of potential descendants in sorted order of StartPos */

begin-anc = AList->firstNode; OutputList = NULL;
for (d = DList->firstNode; d != NULL; d = d->nextNode) {
  for (a = begin-anc; (a != NULL && a.EndPos < d.StartPos); a = a->nextNode) {
    /* skipping over unmatchable a's */ }
  begin-anc = a;
  for (a = begin-anc; (a != NULL && a.StartPos < a.StartPos); a = a->nextNode) {
    if ((a.StartPos < d.StartPos) && (d.EndPos < a.EndPos)
        [&& (d.LevelNum = a.LevelNum + 1)]) {
      /* the optional condition is for parent-child relationships */
      append (a,d) to OutputList; }
    }
  }
}

```

Figure 4: Algorithm Tree-Merge-Desc with Output in Sorted Descendant/Child Order

Given these two input lists, **AList** of potential ancestors (or parents) and **DList** of potential descendants (resp., children), the algorithms in each family can output a list **OutputList** = $[(a_i, d_j)]$ of join results, sorted either by $(\text{DocId}, a_i.\text{StartPos}, d_j.\text{StartPos})$ or by $(\text{DocId}, d_j.\text{StartPos}, a_i.\text{StartPos})$. Both variants are useful, and the variant chosen may depend on the order in which an optimizer chooses to compose the structural joins to match the complex XML query pattern.

3.1 Tree-Merge Join Algorithms

The algorithms in the *tree-merge* family are a natural extension of traditional relational merge joins (which use an equality join condition) to deal with the multiple inequality conditions that characterize the ancestor-descendant or the parent-child structural relationships, based on the $(\text{DocId}, \text{StartPos} : \text{EndPos}, \text{LevelNum})$ representation. The recently proposed multi-predicate merge join (MPMGJN) algorithm [30] is also a member of this family.

The basic idea here is to perform a modified merge-join, possibly performing multiple scans through the “inner” join operand to the extent necessary. Either **AList** or **DList** can be used as the inner (resp., outer) operand for the join: the results are produced sorted (primarily) by the outer operand. In Figure 3, we present the tree-merge algorithm for the case when the outer join operand is the ancestor; this is similar to the MPMGJN algorithm. Similarly, Figure 4 deals with the case when the outer join operand is the descendant. For ease of understanding, both algorithms assume that all nodes in the two lists have the same value of **DocId**, their primary sort attribute. Dealing with nodes from multiple documents is straightforward, requiring the comparison of **DocId** values and the advancement of node pointers as in the traditional merge join.

3.1.1 An Analysis of the Tree-Merge Algorithms

Traditional merge joins that use a single equality condition between two attributes as the join predicate can be shown to have time and space complexities $O(|input| + |output|)$, on sorted inputs, while producing a sorted output. In general, one cannot establish the same time complexity when the join predicate involves multiple equality and/or inequality conditions. In this section, we identify the criteria under which tree-merge algorithms have asymptotically optimal time complexity.

Algorithm Tree-Merge-Anc for ancestor-descendant Structural Relationship:

Theorem 3.1 *The space and time complexities of Algorithm Tree-Merge-Anc are $O(|AList| + |DList| + |OutputList|)$, for the ancestor-descendant structural relationship. ■*

The intuition is as follows. Consider first the case where no two nodes in `AList` are themselves related by an ancestor-descendant relationship. In this case, the size of `OutputList` is $O(|AList| + |DList|)$. Algorithm `Tree-Merge-Anc` makes a single pass over the input `AList` and at most two passes over the input `DList`.² Thus, the above theorem is satisfied in this case.

Consider next the case where multiple nodes in `AList` are themselves related by an ancestor-descendant relationship. This can happen, for example, in the `(section, head)` structural relationship for the XML data in Figure 1. In this case, multiple passes may be made over the same set of descendant nodes in `DList`, and the size of `OutputList` may be $O(|AList| * |DList|)$, which is quadratic in the size of the input lists. However, we can show that the algorithm still has optimal time complexity, i.e., $O(|AList| + |DList| + |OutputList|)$.

One cannot establish the I/O optimality of Algorithm `Tree-Merge-Anc`. In fact, repeated paging can cause its I/O behavior to degrade in practice, as we shall see in Section 4.

Algorithm Tree-Merge-Anc for parent-child Structural Relationship: When evaluating a parent-child structural relationship, the time complexity of Algorithm `Tree-Merge-Anc` is the same as if one were performing an ancestor-descendant structural relationship match between the same two input lists. However, the size of `OutputList` for the parent-child structural relationship can be much smaller than the size of the `OutputList` for the ancestor-descendant structural relationship. In particular, consider the case when all the nodes in `AList` form a (long) chain of length n , and each node in `AList` has two children in `DList`, one on either side of its child in `AList`; this is shown in Figure 5(a). In this case, it is easy to verify that the size of `OutputList` is $O(|AList| + |DList|)$, but the time complexity of Algorithm `Tree-Merge-Anc` is $O((|AList| + |DList|)^2)$; the evaluation is pictorially depicted in Figure 5(b), where each node in `AList` is associated with the sublist of `DList` that needs to be scanned. The I/O complexity is also quadratic in the input size in this case.

²A clever implementation that uses a one node lookahead in `AList` can reduce the number of passes over `DList` to just one.

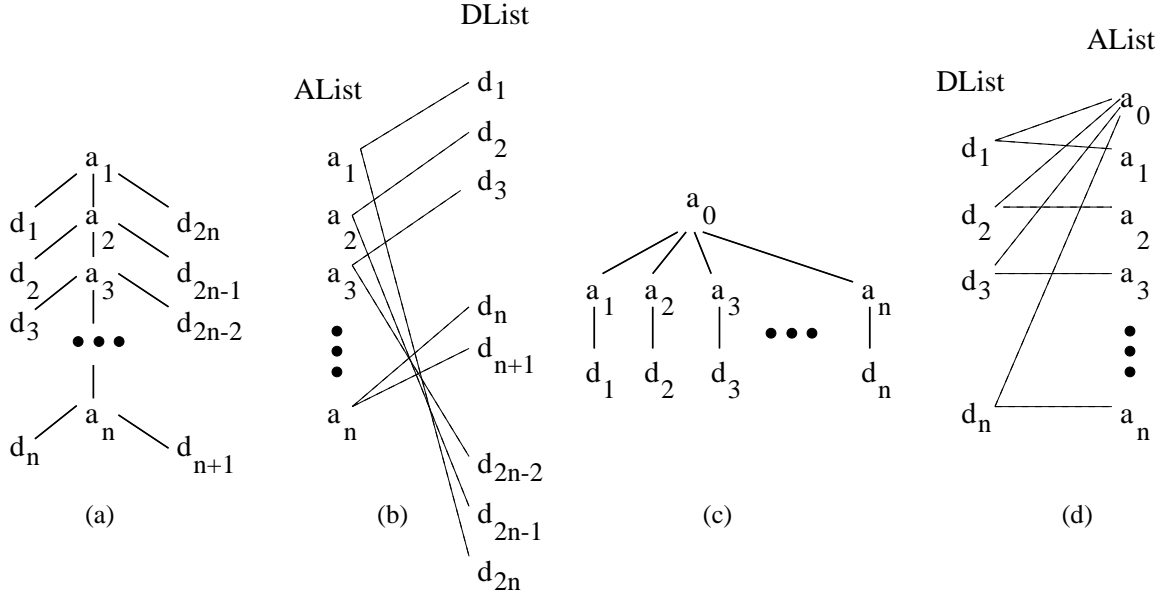


Figure 5: (a), (b) Worst case for `Tree-Merge-Anc` and (c), (d) Worst case for `Tree-Merge-Desc`

Algorithm `Tree-Merge-Desc`: There is no analog to Theorem 3.1 for Algorithm `Tree-Merge-Desc`, since the time complexity of the algorithm can be $O((|\text{AList}| + |\text{DList}| + |\text{OutputList}|)^2)$ in the worst case. This happens, for example, in the case shown in Figure 5(c), when the first node in `AList` is an ancestor of each node in `DList`. In this case, each node in `DList` has only two ancestors in `AList`, so the size of `OutputList` is $O(|\text{AList}| + |\text{DList}|)$, but `AList` is repeatedly scanned, resulting in a time complexity of $O(|\text{AList}| * |\text{DList}|)$; the evaluation is depicted in Figure 5(d), where each node in `DList` is associated with the sublist of `AList` that needs to be scanned.

While the worst case behavior of many members of the tree-merge family is quite bad, on some data sets and queries they perform quite well in practice. We shall investigate the behavior of Algorithms `Tree-Merge-Anc` and `Tree-Merge-Desc` experimentally in Section 4.

3.2 Stack-Tree Join Algorithms

We observe that a depth-first traversal of a tree can be performed in linear time using a stack of size as large as the height of the tree. In the course of this traversal, every ancestor-descendant relationship in the tree is manifested by the descendant node appearing somewhere higher on the stack than the ancestor node. We use this observation to motivate our search for a family of stack-based structural join algorithms, with better worst-case I/O and CPU complexity than the tree-merge family, for both parent-child and ancestor-descendant structural relationships.

Unfortunately, the depth-first traversal idea, even though appealing at first glance, cannot be used directly since it requires traversal of the whole database. We would like to traverse only the candidate nodes provided to us as part of the input lists. We now describe our *stack-tree* family of

```

Algorithm Stack-Tree-Desc (AList, DList)
/* Assume that all nodes in AList and DList have the same DocId */
/* AList is the list of potential ancestors, in sorted order of StartPos */
/* DList is the list of potential descendants in sorted order of StartPos */

a = AList->firstNode; d = DList->firstNode; OutputList = NULL;
while (the input lists are not empty or the stack is not empty) {
  if ((a.StartPos > stack->top.EndPos) && (d.StartPos > stack->top.EndPos)) {
    /* time to pop the top element in the stack */
    tuple = stack->pop(); }
  else if (a.StartPos < d.StartPos) {
    stack->push(a)
    a = a->nextNode }
  else {
    for (a1 = stack->bottom; a1 != NULL; a1 = a1->up) {
      append (a1,d) to OutputList
    }
    d = d->nextNode
  }
}

```

Figure 6: Algorithm Stack-Tree-Desc with Output in Sorted Descendant Order

structural join algorithms; these algorithms have no counterpart in traditional join processing.

3.2.1 Stack-Tree-Desc

Consider an ancestor-descendant structural relationship (e_1, e_2) . Let $\mathbf{AList} = [a_1, a_2, \dots]$ and $\mathbf{DList} = [d_1, d_2, \dots]$ be the lists of tree nodes that match the node predicates e_1 and e_2 , respectively, each list sorted by the $(\text{DocId}, \text{StartPos})$ values of its elements.

We first discuss the stack-tree algorithm for the case when the output list $[(a_i, d_j)]$ is sorted by $(\text{DocId}, d_j.\text{StartPos}, a_i.\text{StartPos})$. This is both simpler to understand and extremely efficient in practice. The algorithm is presented in Figure 6 for the ancestor-descendant case.

The basic idea is to take the two input operand lists, \mathbf{AList} and \mathbf{DList} , both sorted on their $(\text{DocId}, \text{StartPos})$ values and conceptually merge (interleave) them. As the merge proceeds, we determine the ancestor-descendant relationship, if any, between the current top of stack and the next node in the merge, i.e., the node with the smallest value of StartPos . Based on this comparison, we appropriately manipulate the stack, and produce output.

The stack at all times has a sequence of ancestor nodes, each node in the stack being a descendant of the node below it. When a new node from the \mathbf{AList} is found to be a descendant of the current top of stack, it is simply pushed on to the stack. When a new node from the \mathbf{DList} is found to be a descendant of the current top of stack, we know that it is a descendant of all the nodes in the stack. Also, it is guaranteed that it won't be a descendant of any other node in \mathbf{AList} . Hence, the join results involving this \mathbf{DList} node with each of the \mathbf{AList} nodes in the stack are output. If the

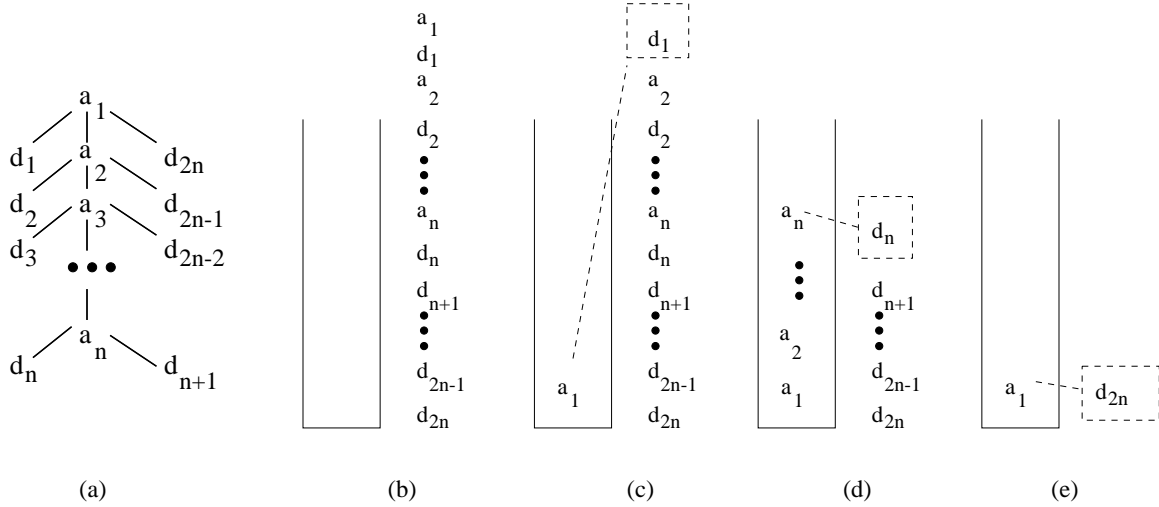


Figure 7: (a) Dataset (b)–(e) Steps during evaluation of **Stack-Tree-Desc**

new node in the merge list is not a descendant of the current top of stack, then we are guaranteed that no future node in the merge list is a descendant of the current top of stack, so we may pop stack, and repeat our test with the new top of stack. No output is generated when any element in the stack is popped.

The parent-child case of Algorithm **Stack-Tree-Desc** is even simpler since a **DList** node can join only (if at all) with the top node on the stack. In this case, the “for loop” inside the “else” case of Figure 6 needs to be replaced with:

```
if (d.LevelNum = stack->top.LevelNum + 1) append (stack->top,d) to OutputList
```

Example 3.1 [**Algorithm Stack-Tree-Desc**]

Some steps during an example evaluation of Algorithm **Stack-Tree-Desc**, for a parent-child structural relationship, on the dataset of Figure 7(a), are shown in Figures 7(b)–(e). The a_i ’s are the nodes in **AList** and the d_j ’s are the nodes in **DList**. Initially, the stack is empty, and the conceptual merge of **AList** and **DList** is shown in Figure 7(b). In Figure 7(c), a_1 has been put on the stack, and the first new element of the merged list, d_1 , is compared with the stack top (at this point (a_1, d_1) is output). Figure 7(d) illustrates the state of the execution several steps later, when a_1, a_2, \dots, a_n are all on the stack, and d_n is being compared with the stack top (after this point, the **OutputList** includes $(a_1, d_1), (a_2, d_2), \dots, (a_n, d_n)$). Finally, Figure 7(e) shows the state of the execution when the entire input has almost been processed. Only a_1 remains on the stack (all the other a_i ’s have been popped from the stack), and d_{2n} is compared with a_1 . Note that all the desired matches have been produced while making only a single pass through the *entire* input. Recall that this is the same dataset of Figure 5(a), which illustrated the sub-optimality of Algorithm **Tree-Merge-Anc**, for the case of parent-child structural relationships. ■

```

Algorithm Stack-Tree-Anc (AList, DList)
/* Assume that all nodes in AList and DList have the same DocId */
/* AList is the list of potential ancestors, in sorted order of StartPos */
/* DList is the list of potential descendants in sorted order of StartPos */

a = AList->firstNode; d = DList->firstNode; OutputList = NULL;
while (the input lists are not empty or the stack is not empty) {
  if ((a.StartPos > stack->top.EndPos) && (d.StartPos > stack->top.EndPos)) {
    /* time to pop the top element in the stack */
    tuple = stack->pop();
    if (stack->size == 0) { /* we just popped the bottom element */
      append tuple.inherit-list to OutputList }
    else {
      append tuple.inherit-list to tuple.self-list
      append the resulting tuple.self-list to stack->top.inherit-list
    }
  }
  else if (a.StartPos < d.StartPos) {
    stack->push(a)
    a = a->nextNode }
  else {
    for (a1 = stack->bottom; a1 != NULL; a1 = a1->up) {
      if (a1 == stack->bottom) append (a1,d) to OutputList
      else append (a1,d) to the self-list of a1
    }
    d = d->nextNode
  }
}

```

Figure 8: Algorithm Stack-Tree-Anc with Output in Sorted Ancestor Order

3.2.2 Stack-Tree-Anc

We next discuss the stack-tree algorithm for the case when the output list $[(a_i, d_j)]$ needs to be sorted by $(\text{DocId}, a_i.\text{StartPos}, d_j.\text{StartPos})$.

It is not straightforward to modify Algorithm `Stack-Tree-Desc` to produce results sorted by ancestor because of the following: if node a from `AList` on the stack is found to be an ancestor of some node d in the `DList`, then every node a' from `AList` that is an ancestor of a (and hence below a on the stack) is also an ancestor of d . Since the `StartPos` of a' precedes the start position of a , we must delay output of the join pair (a, d) until after (a', d) has been output. But there remains the possibility of a new element d' after d in the `DList` joining with a' as long a' is on stack, so we cannot output the pair (a, d) until the ancestor node a' is popped from stack. Meanwhile, we can build up large join results that cannot yet be output. Our solution to this problem is described in Figure 8 for the ancestor-descendant case.

As with Algorithm `Stack-Tree-Desc`, the stack at all times has a sequence of ancestor nodes, each node in the stack being a descendant of the node below it. Now, we associate two lists with each node on the stack: the first, called *self-list* is a list of result elements from the join of this node

with appropriate `DList` elements; the second, called *inherit-list* is a list of join results involving `AList` elements that were descendants of the current node on the stack. As before, when a new node from the `AList` is found to be a descendant of the current top of stack, it is simply pushed on to the stack. When a new node from the `DList` is found to be a descendant of the current top of stack, it is simply added to the self-lists of the nodes in the stack. Again, as before, if no new node (from either list) is a descendant of the current top of stack, then we are guaranteed that no future node in the merge list is a descendant of the current top of stack, so we may pop stack, and repeat our test with the new top of stack. When the bottom element in stack is popped, we output its self-list first and then its inherit-list. When any other element in stack is popped, no output is generated. Instead, we append its inherit-list to its self-list, and append the result to the inherit-list of the new top of stack.

A small, but key, optimization to the algorithm (incorporated in Figure 8) is as follows: no self-list is maintained for the bottom node in the stack. Instead, join results with the bottom of the stack are output immediately. This results in a small space savings, but more importantly it renders the stack-tree algorithm partially non-blocking.

3.2.3 An Analysis of Algorithm `Stack-Tree-Desc`

Algorithm `Stack-Tree-Desc` is easy to analyze. Each `AList` element in the input may be examined multiple times, but these can be amortized to the element on `DList`, or the element at the top of stack, against which it is examined. Each element on the stack is popped at most once, and when popped, causes examination of the new top of stack with the current new element. Finally, when a `DList` element is compared against the top element in stack, then it either joins with all elements on stack or none of them; all join results are immediately output. In other words, the time required for this part is directly proportional to the output size. Thus, the time required for this algorithm is $O(|input| + |output|)$ in the worst case. Putting all this together, we get the following result:

Theorem 3.2 *The space and time complexities of Algorithm `Stack-Tree-Desc` are $O(|AList| + |DList| + |OutputList|)$, for both ancestor-descendant and parent-child structural relationships.*

Further, Algorithm `Stack-Tree-Desc` is a non-blocking algorithm. ■

Clearly, no competing join algorithm that has the same input lists, and is required to compute the same output list, could have better asymptotic complexity.

The I/O complexity analysis is straightforward as well. Each page of the input lists is read once, and the result is output as soon as it is computed. Since the maximum size of stack is proportional to the height of the XML database tree, it is quite reasonable to assume that all of stack fits in memory at all time. Hence, we have the following result:

Theorem 3.3 *The I/O complexity of Algorithm Stack-Tree-Desc is $O(\frac{|AList|}{B} + \frac{|DList|}{B} + \frac{|OutputList|}{B})$, for ancestor-descendant and parent-child structural relationships, where B is the blocking factor. ■*

3.2.4 An Analysis of Algorithm Stack-Tree-Anc

The key difference between the analyses of Algorithms Stack-Tree-Anc and Stack-Tree-Desc is that join results are associated with nodes in the stack in Algorithm Stack-Tree-Anc. Obviously, the list of join results at any node in the stack is linear in the output size. What remains to be analyzed is the appending of lists each time the stack is popped. If the lists are implemented as linked lists (with start and end pointers), these append operations can be carried out in unit time, and require no copying. Thus one comparison per AList input and one per output are all that are performed to manipulate stack. Combined with the analysis of Algorithm Stack-Tree-Desc, we can see that the time required for this algorithm is still $O(|input| + |output|)$ in the worst case.

The I/O complexity analysis is a little more involved. Certainly, one cannot assume that all the lists of results not yet output fit in memory. Careful buffer management is required. It turns out that the only operation we ever perform on a list is to append to it (except for the final read out). As such, we only need to have access to the tail of each list in memory as computation proceeds. The rest of the list can be paged out. When list x is appended to list y , it is not necessary that the head of list x be in memory, the append operation only establishes a link to this head in the tail of y . So all we need is to know the pointer for the head of each list, even if it is paged out. Each list page is thus paged out at most once, and paged back in again only when the list is ready for output. Since the total number of entries in the lists is exactly equal to the number of entries in the output, we thus have that the I/O required on account of maintaining lists of results is proportional to the size of output (provided that there is enough memory to hold in buffer the tail of each list: requiring two pages of memory per stack entry — still a requirement within reason). All other I/O activity is for the input and for the output. This leads to the desired linearity result.

Theorem 3.4 *The space and time complexities of Algorithm Stack-Tree-Anc are $O(|AList| + |DList| + |OutputList|)$, for both ancestor-descendant and parent-child structural relationships.*

The I/O complexity of Algorithm Stack-Tree-Anc is $O(\frac{|AList|}{B} + \frac{|DList|}{B} + \frac{|OutputList|}{B})$, for both ancestor-descendant and parent-child structural relationships, where B is the blocking factor. ■

4 Experimental Evaluation

In this section, we present the results of an actual implementation of the various join algorithms for XML data sets. We consider the following algorithms in our performance evaluation:

- Traversal-style algorithms that locate one node in the pattern by means of an index, and then navigate to find the rest. TOP-DOWN TRAVERSAL (TDT) starts from the root node of

the pattern, and `BOTTOM-UP TRAVERSAL (BUT)` starts from a candidate leaf node of the pattern. Output in these algorithms is naturally sorted by the starting node of the pattern, the root (ancestor) node for `TDT` and the leaf (descendant) node for `BUT`.

- Traditional relational join algorithms. We have a choice of output sort order, which could affect the specific join algorithm chosen. Since relational join algorithms have been studied extensively, rather than re-implement these ourselves, we use these algorithms from a leading commercial database.
- The structural join algorithms we introduce in this paper, namely, `TREE-MERGE JOIN(TMJ)` and `STACK-TREE JOIN (STJ)`. Once more, the output can be sorted in two ways, based on the “ancestor” node or the “descendant” node in the join. Correspondingly, we consider two flavors of these algorithms, and use the suffix “-A” and “-D” to differentiate between these. The four algorithms are thus labeled: `TMJ-A`, `TMJ-D`, `STJ-A` and `STJ-D`.

4.1 Experimental Testbed

We implemented the XML join algorithms, as well as `BUT` and `TDT`, in the `TIMBER` XML query engine. `TIMBER` is a native XML query engine that is built on top of the `SHORE` [5] Storage Manager. Since the goal of `TIMBER` is to efficiently handle complex XML queries on large data sets, we implemented our algorithms so that they could participate in complex query evaluation plans with pipelining. All experiments using `TIMBER` were run on a 500MHz Intel Pentium III processor running WindowsNT Workstation v4.0. `SHORE` was compiled for a 8KB page size. `SHORE` buffer pool size was set to 32MB, and the container size in our implementation was 8000 bytes.

All numbers presented here are produced by repeatedly running the experiments multiple times and averaging all the execution times except for the first run (i.e., these are warm cache numbers).

We also present a comparison of the algorithms when implemented on top of a relational database management system. The DBMS system is running on an SUN 20 Enterprise server with SunOS 5.7.

4.2 Workload

For our workload, we used the IBM XML data generator to generate a number of data sets, of varying sizes and other data characteristics, such as the fanout (`MaxRepeats`) and the maximum depth, using the Organization DTD presented in Figure 9. We also used the `XMach-1` [1] and `XMark` [2] benchmarks, and some real XML data. The results obtained were very similar in all cases, and in the interest of space we present results only for the largest organization data set that we generated. This data set consists of 6.3 million element nodes, corresponding to approximately

```

<!ELEMENT manager (name,(manager|department|employee)+)>
<!ATTLIST manager id CDATA #FIXED "1">
<!ELEMENT department (name, email?, employee+, department*)>
<!ATTLIST department id CDATA #FIXED "2">
<!ELEMENT employee (name+,email?)>
<!ATTLIST employee id CDATA #FIXED "3">
<!ELEMENT name (#PCDATA)>
<!ATTLIST name id CDATA #FIXED "4">
<!ELEMENT email (#PCDATA)>
<!ATTLIST email id CDATA #FIXED "5">

```

Figure 9: DTD used in our experiments

Node	Count
manager	25,880
department	342,450
employee	574,530
email	250,530

Query	XQuery Path Expression	Result Cardinality
QS1	employee/email	140,700
QS2	employee//email	142,958
QS3	manager/department	16,855
QS4	manager//department	587,137
QS5	manager/employee	17,259
QS6	manager//employee	990,774
QC1	manager/employee/email	7,990
QC2	manager//employee/email	232,406

Table 1: Description of Queries and Characteristics of the Data Set

800MB of XML documents in text format. The characteristics of this data set in terms of the number of occurrences of element tags are summarized in Table 1.

We evaluated the various join algorithms using the set of queries shown in Table 1. The queries are broken up into two classes. QS1 to QS6 are *simple structural relationship queries*, and have an equal mix of parent-child queries and ancestor-descendant queries. QC1 and QC2 are *complex chain queries*, and are used to demonstrate the performance of the algorithms when evaluating complex queries with multiple joins in a pipeline.

4.3 Experiment 1: Behavior of Traversal-Style Algorithms

In this section we comment on the performance of the two navigation-based algorithms using a simple structural join query. In order to better understand the behavior of these algorithms, we used the query QS1 from Table 1, and a 10% (600K elements) extract from our 6.3 million elements dataset. Even with this small data set and a large buffer size, we observed that BUT and TDT performed rather poorly. BUT took 283.5s and TDT took 717.8s; the result size was 14,070. Both BUT and TDT have high execution times, since both have to navigate through suitable nodes. Their relative behavior is explained by the fact that TDT needs to traverse *all* children of each *employee* element (by following child- and sibling-pointers), whereas BUT needs to traverse only parent-pointers.

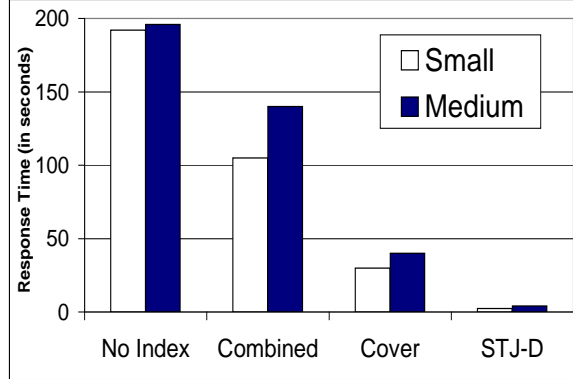


Figure 10: Comparison With Relational Databases Using QS1

In comparison, as we shall see in Section 4.6, both the STJ and TMJ algorithms execute under 15s, even though they are joining much larger lists. The `employee` list has about 0.5 million node entries and the `email` list is about half that size, for the 6.3 million node dataset.

Since the performance of the traversal-style algorithms degrades considerably with the size of the dataset, we concluded that the traversal-style algorithms yield very poor performance compared with the structural join algorithms, and, consequently do not consider the traversal-style algorithms for the remainder of this section.

4.4 Experiment 2: Comparison With Relational Databases

We also experimented with the performance of our algorithms, when implemented on top of a commercial database management system. We implemented STJ-D as an application program interfacing through the use of cursors to a relational database management system and we measured the total time to execute the join operation when the data resides on a DBMS and essentially the algorithm uses the DBMS as a data store, extracts the data as needed and performs the tree matching operation entirely in the application space.

We also formed SQL queries capable of performing tree matching operations by using native (to the DBMS) join algorithms. We execute the simplest query we experimented with, query QS1. We conducted various experiments, studying the impact of alternate access paths to the underlying relations. More specifically, we measure the time to execute the query in the following cases: (a) no index is available in the underlying relations; (b) a combined index on the start and end positions of the numbering is available [30], and (c) a combined index on all attributes is available (cover index in [30]). As argued in [30], the relational approach could offer performance benefits when the selectivity of the nodes involved is very high. To study the impact of selectivity of the underlying nodes, in the performance of the algorithms we experimented with queries complying to the simple structural join pattern, involving nodes of varying selectivity. The first experiment of Figure 10

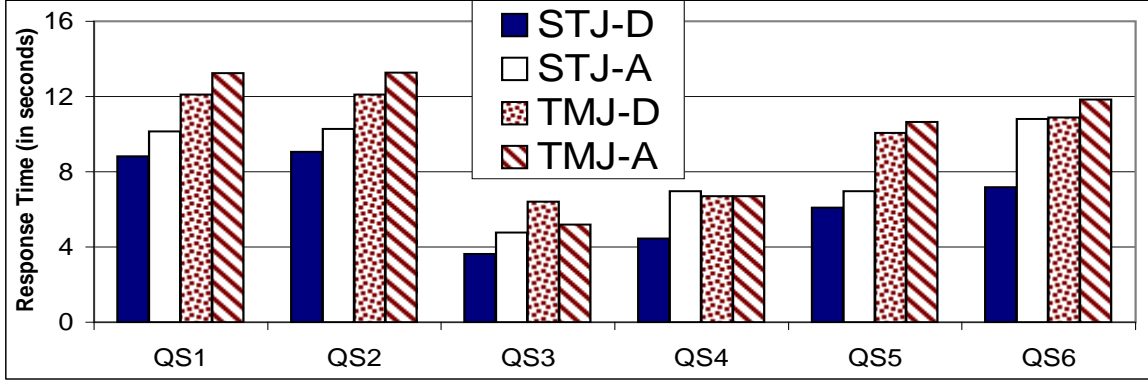


Figure 11: STJ and TMJ, Simple Queries: QS1-QS6

involves a structural join query with a small value of selectivity of each node (at most 10%), and the second involves nodes with a medium value of selectivity (at most 25%). It is evident that the STJ-D approach offers very large performance benefits.

4.5 Detailed Implementation of Structural Join Algorithms

Based on the results presented above, it is evident that traversal algorithms are not very effective. It is also evident that traditional relational join algorithms are not well-suited to exploit the structure in XML patterns. As such, the focus in the rest of the experiments is to characterize the performance of the four new algorithms, and understand their differences. Before doing so in the following subsections, we present here some additional detail regarding the manner in which these were implemented for the experiments reported. Note that our choice of implementation, on top of Shore and TIMBER, was driven purely by the need for sufficient control — the algorithms themselves could just as well have been implemented on many other platforms, including (as new access methods) on relational databases.

All join algorithms were implemented using the iterator model [15]. In this model, each operator provides an *open*, *next* and *close* interface to other operators, and allows the database engine to construct an operator tree with an arbitrary mix of query operations (different join algorithms or algorithms for other operations such as aggregation) and naturally allows for a pipelined operator evaluation. To support this iterator model, we pay careful attention to the manner in which results are passed from one operator to another. Algorithms such as the TMJ algorithms may need to repeatedly scan over one of the inputs. Such repeated scans are feasible if the input to a TMJ operator is a stream from a disk file, but is not feasible if the input stream originates from another join operator (in the pipeline below it). We implemented the TMJ algorithms so that the nodes in a current sweep are stored in a temporary SHORE file. On the next sweep, this temporary SHORE file is scanned. This solution allows us to limit the memory used by TMJ implementation,

as the only memory used is managed by the SHORE buffer manager, which takes care of evicting pages of the temporary file from the buffer pool if required. Similarly for the STJ-A algorithm, the inherit- and self-lists are stored in a temporary SHORE file, again limiting the memory used by the algorithm. In both cases, our implementation turns logging and locking off for the temporary SHORE files. Note that STJ-D can join the two inputs in a single pass over both inputs, and, never has to spool any nodes to a temporary file.

To amortize the storage and access overhead associated with each SHORE object, in our implementation we group nodes into a large *container* object, and create a SHORE object for each container. The join algorithms write nodes to containers and when a container is full it is written to the temporary SHORE file as a SHORE record. The performance benefits of this approach are substantial; we do not go into details for lack of space.

4.6 Experiment 3: STJ and TMJ, Simple Structural Join Queries

Next we compare the performance of the STJ and the TMJ algorithms using all the six simple queries, QS1–QS6, shown in Table 1. Figure 11 plots the performance of the four algorithms. As shown in the Figure, STJ-D outperforms the remaining algorithms in all cases. The reason for the superior performance of STJ-D is because of its ability to join the two data sets in a single pass over the input nodes, and it never has to write any nodes to intermediate files on disk. From Figure 11, we can also see that STJ-A usually has better performance than both TMJ-A and TMJ-D. For queries QS4 and QS6, the STJ-A algorithms and the two TMJ algorithms have comparable performance. These queries have large result sizes (approximately 600K and 1M tuples respectively as shown in Table 1). Since STJ-A keeps the results in the lists associated with the stack, and can output the results only when the bottom-most element of the stack is popped, it has to perform many writes and transfers of the lists associated with the stack elements (recall in our implementation, these lists are maintained in temporary SHORE files). With larger result sizes this list management slows down the performance of STJ-A in practice. Figure 11 also shows that the two TMJ algorithms have comparable performance.

We also ran these experiments with reduced buffer sizes, but found that for this data set the execution time of all the algorithms remained fairly constant. Even though the XML data sets that we use are large, after applying the predicates, the candidate lists that we join are not very large. Furthermore, the effect of buffer pool size is likely to be critical when one of the inputs has nodes that are deeply nested amongst themselves, *and* the node that is higher up in the XML tree has many nodes that it joins with. For example, consider the TMJ-A algorithms, and the query “`manager/employee`”. If many `manager` nodes are nested below a `manager` node that is higher up in the XML tree, then after the join of the `manager` node at the top is done, repeated scans of the descendant nodes will be required for the `manager` nodes that are descendants of the `manager` node

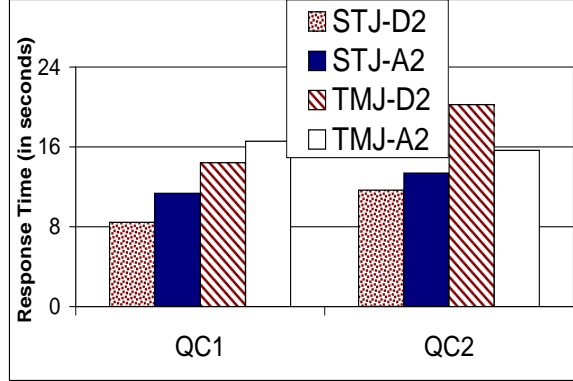


Figure 12: STJ and TMJ, Complex Queries: QC1 and QC2

at the top. Such scenarios are rare in our data set, and, consequently, the buffer pool size has only a marginal impact on the performance of the algorithms.

4.7 Experiment 4: Complex Queries

In our final experiment, we evaluate the performance of the algorithms using the two complex chain queries, QC1 and QC2, from Table 1. Each query has two joins and for this experiment, both join operations are evaluated in a pipeline. For each complex query one can evaluate the query by using only ancestor-based join algorithms or using only descendant-based join algorithms. These two approaches are labeled with suffixes “-A2” and “-D2” for the ancestor-based and descendant-based approaches respectively. The performance comparison of the STJ and TMJ algorithms for both query evaluation approaches (A2 and D2) is shown in Figure 12.

From the figure we see that STJ-D2 has the highest performance once again, since it is never has to spool nodes to intermediate files.

5 Related Work

Matchings between pairs of trees in memory has been a topic of study in the algorithms community for a long time (e.g., see [3] and references therein). The algorithms developed deal with many variations of the problem but unfortunately are of high complexity and always assume that trees are entirely memory resident. The problem also has been considered in the programming language community, as it arises in various type checking scenarios but once again solutions developed are geared towards small data collections processed entirely in main memory.

Many algorithms are known to be very efficient over tree structures. Most relevant to us in this literature are algorithms for checking the presence of sets of edges and paths. Jacobson et al. [16] present linear time merging-style algorithms for computing the elements of a list that are

descendants/ancestors of some elements in a second list, in the context of focusing keyword-based searches on the Web and in UNIX-style file systems. Jagadish et al. [17] present linear time stack-based algorithms for computing elements of a list that satisfy a hierarchical aggregate selection condition wrt elements in a second list, for the directory data model. However, none of these algorithms compute *join results*, which is the focus of our work.

Join processing is central to database implementation and there is a vast amount of work in this area [15]. For inequality join conditions, band join [11] algorithms are applicable when there exists a fixed arithmetic difference between the values of join attributes. Such algorithms are not applicable in our domain as there is no notion of fixed arithmetic difference. In the context of spatial and multimedia databases, the problem of computing joins between pairs of spatial entities has been considered, where commonly the predicate of interest is overlap between spatial entities [18, 24, 19] in multiple dimensions. The techniques developed in this paper are related to such join operations. However, the predicates considered as well as the techniques we develop are special to the nature of our structural join problem.

In the context of semistructured and XML databases, the issue of query evaluation and optimization has attracted a lot of research attention. In particular, work done in the context of the Lore database management system [25, 20, 21], and the Niagara system [23], has considered various aspects of query processing on such data. XML data and various issues in their storage as well as query processing using relational database systems have recently been considered in [14, 28, 27, 4, 12, 13]. In [14, 28, 13], the mapping of XML data to a number of relations was considered along with translation of a select subset of XML queries to relational queries. In subsequent work [27, 4, 12], the authors considered the problem of publishing XML documents from relational databases. Our work is complementary to all of these since our focus is on the join algorithms for the primitive (ancestor-descendant and parent-child) structural relationships. Our join algorithms can be used by these previous works to advantage.

The representation of positions of XML elements used by us, (`DocId`, `StartPos` : `EndPos`, `LevelNum`), is essentially that of Consens and Milo, who considered a fragment of the PAT text searching operators for indexing text databases [8, 9], and discussed optimization techniques for the algebra. This representation was used to compute containment relationships between “text regions” in the text databases. The focus of that work was solely on theoretical issues, without elaborating on efficient algorithms for computing these relationships.

Finally, the recent work of Zhang et al. [30] is closely related to ours. They proposed the multi predicate merge join (MPMGJN) algorithm for evaluating containment queries, using the (`DocId`, `StartPos` : `EndPos`, `LevelNum`) representation. The MPMGJN algorithm is a member of our Tree-Merge family. Our analytical and experimental results demonstrate that the Stack-Tree family is considerably superior to the Tree-Merge family for evaluating containment queries.

6 Conclusions

In this paper, our focus has been the development of new join algorithms for dealing with a core operation central to much of XML query processing, both for native XML query processor implementations as well for relational XML query processors. In particular, the Stack-Tree family of structural join algorithms was shown to be both I/O and CPU optimal, and practically efficient.

There is a great deal more to efficient XML query processing than is within the scope of this paper. For example, XML permits links across documents, and such “pointer-based joins” are frequently useful in querying. We do not consider such joins in this paper, since we believe that they can be adequately addressed using traditional value-based join methods. There are many issues yet to be explored, and we currently have initiated efforts to address some of these, including the piecing together of structural joins and value-based joins to build effective query plans.

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