Relational-Style XML Query

Taro L. Saito
leo@cb.k.u-tokyo.ac.jp

Shinichi Morishita
moris@cb.k.u-tokyo.ac.jp

Department of Computational Biology
University of Tokyo, Japan
Japan Science and Technology Agency (JST)

ABSTRACT

We study the problem of querying relational data embedded in XML. Relational data can be represented by various tree structures in XML. However, current XML query methods, such as XPath and XQuery, demand explicit path expressions, and thus it is quite difficult for users to produce correct XML queries in the presence of structural variations.

To solve this problem, we introduce a novel query method that automatically discovers various XML structures derived from relational data. A challenge in implementing our method is to reduce the cost of enumerating all possible tree structures that match the query. We show that the notion of functional dependencies has an important role in generating efficient query schedules that avoid irrelevant tree structures.

Our proposed method, the relational-style XML query, has several advantages over traditional XML data management. These include removing the burden of designing strict tree-pattern schemas, enhancing the descriptions of relational data with XML’s rich semantics, and taking advantage of schema evolution capability of XML. In addition, the independence of query statements from the underlying XML structure is advantageous for integrating XML data from several sources. We present extensive experimental results that confirm the scalability and tolerance of our query method for various sizes of XML data containing structural variations.

Categories and Subject Descriptors:
H.2.4 [Database Management]: Systems—Query processing

General Terms: Design, Management

1. INTRODUCTION

XML (eXtensible Markup Language) [6] is a text format for tree-structured data. While it is suitable for describing any type of data, there is no such common data format for relational databases. Hence, XML is a promising portable format for relational data. However, there is no obvious simple manner for making queries of relational data embedded in tree-structured XML.

With regard to the expressibility of data, there is no significant difference between XML and relational data [16]. For example, node and edge tables are sufficient to describe tree-structured data in relational databases. Even so, the tree structure of XML is necessary in several cases. XHTML [9], which is an XML version of HTML, uses a tree structure for data layout, which would not work in relational form. Another case is the use of user-defined tags in XML for organizing data groups or appending additional information to extend the schema of XML dynamically.

Other than these two cases involving data layout and the schema-evolution facilities of XML, various types of data can be expressed in relational format. Hierarchical data, often mentioned as an ideal XML application, are not difficult to describe in relational form using simulated data hierarchies with keys to expand multiple columns. An example of this is shown in Figure 1, illustrating relational and XML data with corresponding hierarchies. A triplet of company, section and employee (IDs) comprises a primary key in the following relational data:

```
<table>
<thead>
<tr>
<th>company</th>
<th>section</th>
<th>employee</th>
</tr>
</thead>
<tbody>
<tr>
<td>cl</td>
<td>s1</td>
<td>e1</td>
</tr>
<tr>
<td>cl</td>
<td>s2</td>
<td>e2</td>
</tr>
<tr>
<td>cl</td>
<td>s3</td>
<td>e3</td>
</tr>
</tbody>
</table>
```

Figure 1: Hierarchical data in relational and XML format

This translation from relational data to XML is quite natural and straightforward for the hierarchy of companies through to sections and employees. However, by changing the viewpoints of this relational data, other XML representations are also possible. In Figure 2, the XML data on the left-hand side organize the above relational data for each section, and those on the right-hand side are for each employee:

```
<company id="cl">
  <section id="s1">
    <employee id="e1"/>
  </section>
  <section id="s2">
    <employee id="e2"/>
  </section>
  <section id="s3">
    <employee id="e3"/>
  </section>
</company>

<employeeList>
  <employee id="e1"/>
  <employee id="e2"/>
  <employee id="e3"/>
</employeeList>
```

Figure 2: Various XML representations of relational data
Although the meaning of these XML data is the same, XML queries using path expressions are dependent on the specific XML structures. For example, an XPath [8] query to retrieve all employees in a company c1 and section s1 is completely different for each XML dataset, as shown below:

\[ P1: //company[@id='c1']/section[@id='s1']/employee \] (Fig. 1)
\[ P2: //section[@id='s1'][company[@id='c1']]/employee \] (lhs of Fig. 2)
\[ P3: //employee[company[@id='c1'][section[@id='s1']] \] (rhs of Fig. 2),

where the descendant-axis (\(/\)) traverses an arbitrary-depth of XML data, while the child-axis (\(/\)) for child nodes, the attribute-axis (@) for attribute nodes (data contained in start tags), and the brackets([]) enclose twig nodes to test. This example indicates that without knowledge of the precise XML structure, users cannot produce correct XML queries.

This problem of the structural variations of XML data is common when translating relational data into XML. One possible solution to this problem is to disallow structural variations using an XML schema [25], DTD [6], or RelaxNG [18]. However this greatly limits the flexibility of XML data modeling, and prevents dynamic schema evolution or the population of XML nodes with user-defined tags. The requirement for XML schemas comes mainly from the existing standard XML processing methods (e.g., SAX [20], DOM [6], XPath [8], XQuery [5], etc.). These query methods are based on tree navigation, so without detailed knowledge of the underlying XML structure, it is quite difficult to traverse tree-structured XML data correctly.

A brute-force solution would be to cover all structural variations with a single XPath expression by exhaustively concatenating all possible tree patterns. For the above example, the query would be \( P1 \lor P2 \lor P3 \). However, a slight change in the XML structure, for example when some employees join a project team, and thus XML data are modified as in Figure 3, would still force the user to modify query statements or XML reader programs to accommodate this new structure:

```
<company id="c1">
  <section id="s1">
    <team project="p1">
      <employee id="e1"/>
      <employee id="e2"/>
    </team>
  </section>
  <section id="s2">
    <employee id="e3"/>
  </section>
</company>
```

Figure 3: Decorating employee data with a custom tag, team.

Unlike the examples of the above XPath queries, SQL query statements are stable after this sort of schema evolution. For example, the following SQL select statement,

```
SELECT company, section, employee FROM ...
```
can be used without any modification, because a relation consisting of company, section and employee nodes still holds after insertion of the team node.

This observation motivated us to develop a means of querying XML data in relational style. For example, we use a simple expression (company, section, employee) to specify node names in a relation without reference to the tree structure, and retrieve variously structured relational data embedded in XML. A key insight in this development is that even if XML representations vary according to the specific viewpoint of relational data, these XML structures are all derived from the same relational data. To describe these variously structured relational data with a simple expression, we define a class of tree structures that construct relations in XML. Given a query expression, e.g., (company, section, employee), our query method covers all possible tree structures that can be generated from input company, section and employee nodes.

A challenge in implementing our query method is to discover the appropriate tree structures from the XML data. In general, the number of possible structural variations of \( n \) XML nodes is \( n^{n-1} \), which is identical to the number of labeled trees with \( n \) nodes. To improve query performance, we must avoid issuing \( n^{n-1} \) queries. Another challenge is that even for a single tree pattern, its instances in XML data could be numerous. For example, XML data in Figure 4 has a hierarchical pattern with one company node, three section nodes and five employee nodes. While there are \( 1 \times 3 \times 5 = 15 \) instances of (company, section, employee) pairs, only 5 of those are appropriate in that they connect each employee node with its corresponding parent section node. This shows that naive enumeration of tree instances is inefficient for larger volumes of XML data. Therefore, eliminating incorrect tree structures is another key to achieving good query performance.

To remove irrelevant tree structures from query results, it is necessary to know the implied semantics in the XML data, e.g., each employee node belongs to a section node. We describe these semantics with functional dependencies (FDs) [17] tailored to XML. For example, an FD could be employee \( \rightarrow \) section, meaning that each employee node belongs to a unique section node. Our definition of FD is flexible to allow structural variations, as a section node may be a child of an employee node (Figure 2), or there may be another node inserted between them, as shown in Figure 3.

Our proposed method, the relational-style XML query, provides new insight into XML query processing. While the de facto standards for XML query processing languages, such as XPath [8] and XQuery [5], require explicit path expressions to perform queries, we use FDs to define XML data structures, and thus have no need to specify tree structures in query statements. All we need to query XML data is to describe target nodes of interest with tag names, predicates, keywords, etc. Relational-style XML queries enable the user to perform queries without detailed knowledge of the XML structure. This means query expressions are much simpler than those for path-based query methods.

The outline and contributions of this paper are as follows:

- We present a compelling example of the relational-style XML query, which does not use explicit path structures for either queries or schemas (Section 2).
- We define a relation in XML that can capture structural variations in XML data, and present an XML algebra to describe XML queries (Section 3).
• We define FDs for XML, and create a relationship between XML structures and FDs (Section 4).
• We present optimization techniques based on our XML algebra to expedite retrieval of XML structures satisfying FDs (Section 5).
• We present experimental evaluation of our proposed methods to confirm the scalability and tolerance of our proposed method. (Section 6).

We present a survey of related work in Section 7, and conclude this work in Section 8.

2. RELATIONAL-STYLE XML QUERY

Relational-style XML query allows structural variations in XML databases. This capability provides a great impact on XML query processing. For example, by detecting relational-part from existing XML data, we call this a relation in XML, query expressions of XML becomes much simpler than path-based query methods. In addition, in creating a XML database from scratch, its schema design becomes straightforward translation from an ER-diagram [17], which is far simpler than defining a comprehensive tree schema. To illustrate these benefits, let us consider an XML database of a company data that has several employees and working projects. Figure 5 illustrates an ER-diagram of this database:

![ER-diagram of a company data](image)

Figure 5: An ER-diagram of a company data, and its decomposition into relations.

To create an XML database from this model, we first decompose this ER-diagram into several relations:

R1: (company, section, employee)
R2: (project, task, employee)
R3: (employee, name)

We choose these relations so that each of these node pairs organizes a reasonable unit in this data model, so a relation can be a much smaller fragment, e.g., (company, section), (section, employee), etc. This decomposition process is similar to the design of table schemas in relational databases.

One-to-Many Relationship. In this ER-model, a company has several sections, and each employee belongs to one of these sections. This is an example of one-to-many relationships between a company and sections, and a section and employees. To describe these relationships in the ER-diagram, we extract the following functional dependencies (FDs):

employee → section (Each employee belongs to a section)
section → company (Each section belongs to a company)

An one-to-many relationship between P and Q corresponds to an FD Q → P, meaning that from each node Q we can uniquely determine another node P. Stated in another way, a node P may have several associated nodes Q.

Many-to-Many Relationship. This data model has a project node, each of them has several tasks. Each task is assigned to an employee, and employees may be assigned several tasks in several projects. This is a many-to-many relationship between projects and employees. In general, we can divide such many-to-many relationships into one-to-many relationships [17]. The following FDs represent two one-to-many relationships (project-task and employee-task):

\[
\text{task} \rightarrow \text{project}
\]
\[
\text{task} \rightarrow \text{employee}
\]

Relation to XML Structures. Relations and FDs are sufficient to describe a schema of XML. Figure 6 shows an example of XML data generated from the ER-diagram. This example involves various tree structures that denote data in the same relation. The node pairs of (project, task, employee) are hierarchically organized when ignoring the employee list node. The tree structures of (project, task, employee) pairs are different under the project list and task list nodes. In the traditional XML schema design, we have to decide which structure to use, even though this structural difference has no significant meaning. The relational-style XML query completely does away with the inconvenience, because query expressions for retrieving these distinct tree structures are the same as follows:

\[
(\text{project}, \text{task}, \text{employee})
\]

From a given set of definitions of relations and FDs, our query processor automatically finds XML structures that form a relation.

Querying Relational Data Enhanced with XML. XML has rich-data semantics that can enhance the meanings of relational data. For example, a relation (project, task, employee) in Figure 6 is decorated with an intermediate node, active (17), which does not appear in the ER-diagram. The other nodes employee list (2), project list (14) and task list (26) also enhance relational data by grouping the XML structures representing relations.

In XML, it is required to handle database queries that contain both relational and XML semantics. Consider a query for employee names who are working for active tasks. In Figure 6, two task nodes 18, 22 are marked as active, but the ER-diagram has no information of the active node. A query Q1 in Figure 7, which is written in XQuery [5], has to traverse several paths, then performs a value-based join operation on employee@id. To produce this XQuery statement, the user must know that the active node appears only under the project list node, and employee names are under the employee list. However, learning such knowledge requires a great deal of efforts and demands the ability to make a complex query.

In the relational-style XML query, this query expression becomes much simpler as shown in Q2 in Figure 7, which first retrieves two relations (employee, name) and (active, task, employee), then joins them by using employee@id values. Since we have the knowledge of the FD task → employee, we can avoid invalid node pairs such as (employee (20), active (17), task (22)), which connects irrelevant employee and task nodes. In processing XML queries, we have to correctly extract relations embedded in XML, such as (employee, task), and at the same time to locate XML nodes (e.g., active) associated to these relations.

3. RELATION IN XML

In this section, we define a relation in XML that specifies XML structures of interest using a pattern tree, which allows various structure organizations by using the notion of amoeba [19]. On this basis, we define an XML algebra, which is the foundation for describing XML queries with a nested form of expressions.

Throughout this paper, we use a tree model of XML data, made up of tree nodes with text values and edges. To distinguish element nodes (general tree nodes) and attribute nodes [6], attribute node
names are prefixed with "@." Each element and attribute node has a global ID, which is unique in the XML data.

**Amoeba Structure.** To describe various tree structures that can be generated from XML nodes, the notion of amoeba has been proposed [19] as a relaxed definition of trees from the graph theory:

**Definition 3.1 [Amoeba].** Given a set \( r = \{r_1, \ldots, r_k\} \) of XML nodes, where \( r_i \) is an XML node, we say \( r \) is an amoeba if one of \( r_1, \ldots, r_k \) is a common ancestor of the others, denoted by \( \langle \langle r_1, \ldots, r_k \rangle \rangle \).

For example, every structural variation in Figure 8 is an amoeba. To describe a set of amoebas consisting of three types of nodes, project, task, and employee, we use a notation \( \langle\langle \text{project, task, employee} \rangle\rangle \). It is important that regardless of the structure of a node set in the XML data, the node set can be considered to be an amoeba as long as it contains a common root node. The root node of an amoeba is usually an element or attribute node, but a singleton node set, e.g., \( r = \{r_1\} \), can also form an amoeba. This definition of amoeba allows node insertions. Figure 3 shows an example of this where a team node is inserted into the tree structure of company, section and employee nodes.

### 3.1 Relation in XML

To describe a set of XML data fragments all of which match a specific tree pattern, we need a pattern expression, such as XPath [8]. Such an XPath expression is typically modeled as a pattern tree [11]. However, in the presence of structural variations, it is too restrictive to demand that data structures obey a single tree pattern. Furthermore, concatenating all possible path structures into a single XPath expression can be tedious. To represent both strict and flexible tree structures easily, we introduce the notion of a **relation in XML**, which can express various path structures, including twigs and amoebas:

**Definition 3.2 [Relation in XML].** A relation \( R \) in XML is a k-ary tuple of nodes (for element and attribute nodes) with a Boolean conjunction of conditions of the following types:

- A condition to specify a subset of nodes in \( R \), say \( [a, b, c] \), constructs an amoeba, denoted \( \langle a, b, c \rangle \).
- For two XML nodes \( u \) and \( v \) in \( R \), \( u \) is a child (or descendant) of \( v \).
- Comparison of a text value of a node in \( R \) with a constant using one of the operators \( =, >, \leq, \geq \).

Although it is possible to use other types of conditions (e.g. document orders of nodes, sibling axes in XQuery [8], etc.), we limit the condition types in a relation for the purpose of illustration.

**Definition 3.3 [An instance of a relation].** An instance of a relation \( R \) in an XML data, denoted \( [R] \), is a set of node tuples \( \{ [r_1], \ldots, [r_k] \} \) such that each XML node \( r_i \) matches a corresponding node name in \( R \), and satisfies all conditions in \( R \). We denote a node tuple \( r \) contained in an instance of \( R \) as \( r = \{r_1, \ldots, r_k\} \in [R] \).

A relation in XML can be used to describe a fixed tree structure, which is common in XPath expressions. For example, by using an XPath expression, we simply denote a relation in XML as \( R_1 = //\text{employee/name} \) to specify a tree pattern consisting of employee and name nodes, where name nodes must be a child of a employee node in the XML document. We denote an instance of \( R_1 \) as \( [R_1] \) or \( //\text{company/name} \). Figure 9 shows another example of a relation in XML that has element nodes project, task and employee, and a text value \( \text{employee@id} \). Its predicates are \( \langle \text{project, task, employee} \rangle \) and a path constraint that an employee@id node is a child of an employee node. Its instance is shown in the table in
3.2 XML Algebra

We present three essential algebraic operations for XML queries: selection, projection and amoeba join.

Selection. First, we introduce the selection operation for XML:

Definition 3.4 [Selection]. Let R be a relation in XML, and C be a Boolean conjunction of conditions listed in Definition 3.2. A selection operator, denoted by \( \sigma_C(R) \), applies a condition C to a relation \( R \), i.e.,

\[
\sigma_C(R) = \{ r \mid r \in R \land r \text{ satisfies } C \}.
\]

Node Labels. It is essential to have the capability of specifying some nodes in a relation in XML. In relational databases, a table has columns and each column has a name. Hence, users of the relational database can perform algebraic operations by specifying data columns by name. Node names can be used as equivalents in XML. For example, in an XML relation \( R_1 = \langle \text{project}, \text{task}, \text{employee} \rangle \), node names project, task and employee can be used to specify nodes in \( R_1 \). To avoid ambiguity of node names between several relations in XML, we use a dot notation. For example, when \( R_3 = \langle \text{task}\_id, \text{id} \rangle \) and \( R_4 = \langle \text{employee}\_id \rangle \), we can distinguish these two \( \text{id} \) nodes as \( R_3\_\text{id} \) and \( R_4\_\text{id} \), or we simply denote these node labels as \( \text{task}\_\text{id} \) and \( \text{employee}\_\text{id} \). We use a label for a text value of a node as \( \text{[node name]} \). For example, \( \text{[task}\_\text{id}] \) and \( \text{[name]} \) specify text values for \( \text{task}\_\text{id} \) and \( \text{name} \), respectively.

In this paper, we consider that the inputs and outputs of an XML query are relations in XML, and that a query is evaluated using instances of each input relation. Then, the query produces an instance of another relation. In particular, XML queries often involve intermediate relations, which are themselves relations. Assigning new temporary node names to all intermediate relations can be a daunting task. Therefore, for readability, we assume node names are inherited by the intermediate relations. For example, if we perform a selection operation on a relation \( R = \langle \text{book}\_\text{isbn} \rangle \) and generate another relation \( R' = \langle \text{book}\_\text{isbn} = \text{xx7} \rangle \), then we can use the node names book and book\_isbn that exist in both the relations \( R \) and \( R' \).

Projection. To retrieve a specific set of XML nodes from a relation, we define the projection of a relation \( R \), denoted by \( \pi_X(R) \), where \( X \) is a list of node labels. For example, when \( R = \langle \text{employee}\_name, \text{[name]} \rangle \) and \( \mathbb{R} = \{(4,6),(7,9),(11,13)\} \), then the result of a projection \( \pi_{\text{name}}(R) \) is \( \{(4,6),(9,13)\} \).

Ameoba Join. Given a list of relations in XML, \( R_1 = \langle \text{project}, \text{task} \rangle \), \( R_2 = \langle \text{employee} \rangle \), for example, we need an operation to construct their amoebas. This operation is called an amoeba join [19]. A similar operation is a structural join [1], which concatenates two nodes \( p \) and \( q \) if \( p \) is an ancestor of \( q \). The structural join is generally used to process descendant-axis (/) queries. However, to handle structural variation, we also must consider both the case where \( p \) is an ancestor of \( q \), or \( p \) is a descendant of \( q \). In addition, there are indirect structural relationships involving more than two nodes, for example, nodes \( p \) and \( q \) connected through another node \( r \). To collect instances of variously structured XML data, we define the amoeba join operation as an operator in the XML algebra:

Definition 3.5 [Amoeba Join]. Given a list of node labels \( L_1, ... , L_m \) and a list of input relations \( R_1, ... , R_k \), an amoeba join operation \( AJ_{L_1,...,L_m}(R_1, ... , R_k) \) is a selection with an amoeba condition for \( L_1, ... , L_m \), i.e.,

\[
AJ_{L_1,...,L_m}(R_1, ... , R_k) = \sigma_{\langle L_1, ... , L_m \rangle}(R_1 \times \cdots \times R_k).
\]

For example, when \( R_1 = \langle \text{project} \rangle \), \( R_2 = \langle \text{task} \rangle \) and \( R_3 = \langle \text{employee} \rangle \), then an amoeba join \( AJ_{\langle \text{project}, \text{task}, \text{employee} \rangle}(R_1, R_2, R_3) \) is a selection with a condition \( \langle \text{project}, \text{task}, \text{employee} \rangle \), and generates all instances of amoebas in the XML document, matching one of the structures in Figure 8.

4. Functional Dependencies

A relation in XML has the capability of handling variously structured XML data. However, without knowledge of the semantics hidden in XML data, it is not possible to retrieve correct XML structures. For example, Figure 9 shows invalid tuples (colored in blue) that connect irrelevant task and employee nodes (18, 24) and (22, 20). To resolve this problem, we need information of data semantics, such as each task belongs to a project and is assigned to an employee. These data semantics are described with FDs, \( \text{task} \rightarrow \text{project} \) and \( \text{task} \rightarrow \text{employee} \). In this section, to incorporate data semantics into XML, we define FDs in XML and a class of relations that can be used to describe XML structures satisfying FDs.

We describe a functional dependency for XML with node labels in relations. Let \( X \) and \( Y \) be lists of node labels. Then, a functional dependency for XML is expressed as \( X \rightarrow Y \). Now, we give the definition of FDs in XML:

Definition 4.1 [FDs in XML]. We say a relation \( R \) satisfies an FD relation \( X \rightarrow Y \) if for each pair of instances \( p, q \in \mathbb{R} \), \( p.X = q.X \) implies \( p.Y = q.Y \), where \( p.X \) denotes a list of nodes (or text values) in \( p \) corresponding to each node label in \( X \). The equality of two nodes (or text values) \( n_1 \) and \( n_2 \) is defined as follows:

\[
\begin{align*}
\text{when } n_1 \text{ and } n_2 \text{ are XML nodes}, & \\
\text{when } n_1 \text{ and } n_2 \text{ are text values}, & \\
\end{align*}
\]

where \( n.d \) is a unique node ID in the XML data.

Intuitively, an FD relation \( X \rightarrow Y \) specifies that a node set belonging to \( X \) uniquely determines a node belonging to \( Y \). For example, some instances in Figure 9 violate the FD relation \( \text{task} \rightarrow \text{employee} \): two distinct employee nodes 20 and 24 are associated to each of the task nodes in different manners.
Given a list tree relation class of XML structures, called a tree relation. To avoid these irrelevant node connections, while allowing various and thus invalid nodes may be connected, as illustrated in an amoeba structure itself is a connected component of tree nodes, and thus invalid nodes may be connected, as illustrated in Figure 9. To avoid these irrelevant node connections, while allowing various tree structures in describing XML data, we introduce a restricted class of XML structures, called a tree relation.

Before defining a tree relation, we introduce some notations. Let $F$ be a set of FDs, $NL(F)$ is the set of node labels appearing in $F$. Given a list $L$ of relations $R_1, ..., R_k$, then if each $R_i$ contains at least one node label in $NL(F)$, and all node labels in $NL(F)$ are contained in $L$, we say that $I$ covers $NL(F)$. For example, for $F = \{\text{employee} \rightarrow \text{name}\}$, then $NL(F) = \{\text{name}\}$, and thus the pair of relations $R_1 = /employee$ and $R_2 = /name$ covers $NL(F)$.

Now, we define a tree relation in XML:

**Definition 4.2 [Tree Relation].** Let $F$ be a set of FDs and $R_1, ..., R_k$ be a list of relations that covers $NL(F) = \{L_1, ..., L_m\}$. A tree relation $R$ for $F$ is a result of selection $\sigma_{\text{project}}(R_1 \times \cdots \times R_k)$ such that $R$ satisfies all FDs in $F$, and $C$ is a conjunction of the following amoeba conditions:

1. $\{L_1, ..., L_m\}$
2. $\forall X, Y \in NL(F)$, $\forall \text{FD } X \rightarrow Y_i \in F$ for each $\text{FD } X \rightarrow Y_i$.

where $X$ is a list of node labels, and each $Y_i$ is a single node label.

For example, when $F = \{A \rightarrow B, B \rightarrow CD\}$, then $NL(F) = \{A, B, C, D\}$, and its tree relation for $F$ has the following condition:

$$\langle A, B, C, D \rangle \land \langle A, B \rangle \land \langle B, C \rangle \land \langle B, D \rangle.$$ 

As another example, an FD with the form $AB \rightarrow C$, which has several node labels in the left hand side, imposes the constraint $\langle A, B, C \rangle$.

The first constraint (P1) $\langle L_1, ..., L_m \rangle$ confirms that nodes in $NL(F)$ construct an amoeba, i.e., a node set of $L_1, ..., L_m$ must at least form a tree structure in the XML data. The second constraint (P2) indicates that nodes appearing in an FD must also have an amoeba structure. Intuitively, to establish the correspondence between FDs and XML structures, we consider XML nodes that construct an amoeba structure are semantically related. If there are partial dependencies (FDs) within a relation, XML structures must represent all of these relationships. Figure 10 illustrates variations of tree relations for several sets of FDs; A tree relation of nodes $A, B$ and $C$ must form a tree structure but allows several tree shapes. When FDs are defined in this relation, tree shapes are restricted so that these FDs can be represented in these tree structures.

These structural constraints imposed by FDs have an important role in eliminating incorrect XML structures that do not match the data semantics. For example, when $F$ has two FDs $\text{task} \rightarrow \text{project}$ and $\text{task} \rightarrow \text{employee}$, then a tree relation for $F$ must satisfy the following condition:

$$\langle \text{project}, \text{task}, \text{employee} \rangle \land \langle \text{task}, \text{project} \rangle \land \langle \text{task}, \text{employee} \rangle.$$ 

In Figure 6, an instance of a relation $R_1$ satisfies all of these conditions. Thus, we say $R_2$ is a tree relation for $F$. The first constraint $\langle \text{project}, \text{task}, \text{employee} \rangle$ allows all possible tree structures consisting of these three nodes. However, a node pair $(15, 18, 24)$ in Figure 9 satisfies $\langle \text{task}, \text{project}, \text{employee} \rangle$ but connects irrelevant task (18) and employee (24) nodes. Hence, the other constraints $\langle \text{task}, \text{project} \rangle$ and $\langle \text{task}, \text{employee} \rangle$, which are imposed by FDs, are needed to remove such inappropriate tree structures.

Next, we present some examples of FDs in XML:

- $\text{employee} \rightarrow \text{employee@id}$: Each employee node must have an @id attribute node.
- $\text{employee@id} \rightarrow \text{employee}$: The is the opposite of the FD above. In XML, every attribute must belong to a single element, so this type of FD always holds for all attribute nodes.
- $\text{author} \rightarrow \text{paper}$: Each author belongs to a paper. In other words, a paper may have several authors. The rationale to use an amoeba structure $\langle \text{author}, \text{paper} \rangle$ to represent this one-to-many relationship is that, for each paper node, its author nodes should be ancestor or descendant nodes, not sibling or other nodes. The amoeba condition $\langle \text{author}, \text{paper} \rangle$ covers such tree structures. If several paper nodes are found for an author node, such XML data violate this FD, and needs to be modified.
- $\langle \text{book@isbn} \rangle \rightarrow \text{book}$: Given an book@isbn value, we can uniquely determine a book node. In this case, the book@isbn value is a key (global ID) of book node, no duplicate value of book@isbn is allowed in the XML document.
- $\text{country, ssn} \rightarrow \text{person}$: Any person node is identified by a pair of country and ssn (social security number) nodes. This is an example of a primary key with two nodes. Either of the country or ssn nodes is not sufficient to locate a person node, as an ssn may not be unique outside of a country.
- $\text{country, [person@ssn]} \rightarrow \text{person}$: With the information of an country node and person@ssn value, a unique person node can be determined. This example can be considered as a relative key [7], which localizes the key definition under the specified path, as uniqueness of [person@ssn] values is also localized in the context of the country node, but various data structures are allowed compared to the relative keys proposed in [7]. For example, a country node can be a parent or child of a person node in our definition of FDs.

5. QUERY PROCESSING

5.1 Pushing Structural Constraints

Using the operations defined so far, we are able to implement the
“pushing selection” technique developed for relational databases, which makes it efficient to process tree relations for a set of FDs.

Queries for a tree relation for a set of FDs contain several amoeba predicates. As we explained in Section 4, amoeba constraints imposed by FDs eliminate irrelevant structures to the tree relation. The order in which these conditions are applied is an important factor in reducing the size of the intermediate query results.

In this section, we present optimization techniques that translate a query operation into a nested form of several amoeba joins so that temporary results can be minimized by gradually applying structural constraints imposed by FDs. This method enables selective retrieval of XML structures that satisfy each amoeba constraints, and avoids extraction of unwanted XML structures. To give an equivalent translation of a query expression, we incorporate the commutative law and cascading selection of relational algebra [17] into XML:

**Theorem 5.1** [Pushing Selection]. Let \( R \) and \( S \) be input relations, and \( C \) be a condition. When a relation \( S \) contains no node label that appears in \( C \), the following translations hold:

\[
\sigma_C(R \times S) = \sigma_C(R) \times S \quad \text{(commutative law)}
\]

\[
\sigma_{c_1,c_2}(R) = \sigma_{c_1}(\sigma_{c_2}(R)) \quad \text{(cascading selection)},
\]

where \( c_1 \) and \( c_2 \) are conditions.

**Proof (Sketch).** The proof is an induction on the number of conditions based on the fact that relations of the left-hand side and right-hand side in the above expressions have the same set of conditions.

Using the rules in Theorem 5.1, we can decompose a selection operation to retrieve a tree relation into a nested form of selections. If \( A \) is a set of amoeba conditions, and \( a \) is an amoeba condition in \( A \), then the following translation holds:

\[
\sigma_A(R \times S) = \sigma_A(\sigma_{A \setminus \{a\}}(R) \times S) = \sigma_A(\sigma_{A \setminus \{a\}}(R) \times S),
\]

where a relation \( S \) does not contain node labels in \( A - \{a\} \).

When \( a = \{X, Y\} \), a selection operation \( \sigma_a \) is an amoeba join \( AJ_{X,Y} \) that connects nodes \( X \) and \( Y \). Hence, this decomposition technique can be used repeatedly to derive a series of amoeba joins equivalent to the original query. Figure 11 illustrates query schedules generated by this decomposition. In this example, we have three amoeba conditions, \( \{A, B, C\} \), \( \{A, B\} \) and \( \{A, C\} \). When we choose one of the conditions, say \( \{A, B, C\} \) (\( a_1 \)), as a decomposition target, the query schedule becomes like the left-hand schedule in Figure 11. This schedule effectively reduces the search space of possible relations by evaluating amoeba conditions \( a_3 \) and \( a_2 \) in earlier steps, thus decreasing the input size of the final \( AJ_{A,B,C} \) operation. On the other hand, the right-hand schedule evaluates the condition \( a_1 \) first, which enumerates all possible structural variations, and subsequently makes selections with \( \{A, B\} \) and \( \{A, C\} \).

This translation is equivalent to the so-called pushing-down selection in relational algebra. This technique is also useful in XML query processing to eliminate instances of irrelevant XML structures from intermediate query results.

**Parent-Child Join Decomposition.** Functional dependencies are frequently observed between parent and child nodes, e.g., \( \text{task}@\text{id} \rightarrow \text{task} \), which imposes \( \{\text{task}@\text{id}, \text{task}\} \), and the task node must be the parent of the \( \text{task}@\text{id} \) node. In this case, we can explicitly decompose the query using a parent-child join:

**Corollary 5.2.** Let \( R \) and \( S \) be input relations, and \( A \) be a set of amoeba conditions. For an amoeba condition \( a = \{P, C\} \in A \), where \( P \) is the parent node of \( C \), and when a relation \( S \) does not contain node labels in \( A - \{a\} \), the following translation holds:

\[
\sigma_A(R \times S) = PC_{PC}(\sigma_{A \setminus \{a\}}(R) \times S),
\]

where \( PC_{PC} \) denotes the parent-child join, which is a specialized version of an amoeba join that connects parent nodes \( P \) and child nodes \( C \).

### 5.2 Minimal Relation

Unlike relational databases that use flat tables, relations in XML have tree structures. This structural discrepancy often demands an XML query to involve extra nodes that do not necessarily appear in the final results. For example, consider a query for a pair of project and employee nodes from a relation \( \text{relation}(\text{project}, \text{task}, \text{employee}) \).

In SQL, simply specifying project and employee labels is sufficient to produce this query statement. In XML, however, we also have to include task node label in the query operation, because when a task node is a root node of the amoeba structure, the project and employee nodes cannot be connected without the task node. Therefore, project, task, customer is a minimal relation required to answer this query.

The algorithm to compute minimal relation for a given list \( L \) of input node labels is simple. Let \( F_q \) be a subset of pre-defined FDs such that each FD in \( F_q \) contains some node label appeared in \( L \). Then, the minimal relation of the node list \( L \) is \( \text{NL}(F_q) \), which is a list of all node labels that appear in \( F_q \) and \( L \). For example, when \( L = \{\text{project}, \text{employee}\} \) and a pre-defined set \( F \) of FDs is \( \{\text{task} \rightarrow \text{project}, \text{task} \rightarrow \text{employee}\} \), then \( F_q \) is identical to \( F \), and \( \text{NL}(F_q) = \{\text{project}, \text{task}, \text{employee}\} \), which is the minimal relation of \( \{\text{project}, \text{employee}\} \).

Its query operation is described as follows:

\[
\pi_{\text{project}, \text{employee}}(\sigma_C(\text{project} \times \text{task} \times \text{employee})) \quad (S_1)
\]

where a condition \( C = \{\text{project}, \text{task}, \text{employee}\} \cup \{\text{project}, \text{task}\} \cup \{\text{task}, \text{employee}\} \), which is derived from \( F_q \). This query correctly locates the minimal relation for the project and employee nodes, and then the projection eliminates the task node, which is not contained in the original input. The notion of minimal relations can be utilized to complement some missing nodes in the query, so the users can produce queries without considering structural differences between relational and XML structures. For example, a query statement for \( S_1 \) is simple as follows:

\[
(\text{project}, \text{employee}) \quad (S_1)
\]

There is a case that some node labels in a query do not appear in any FDs. This is usual for relational data enhanced with XML syntax, as explained in Section 2. For example, a minimal relation of \( \{\text{task}, \text{list}, \text{task}, \text{employee}\} \) has no additional node, since an only FD related to this relation is \( \text{task} \rightarrow \text{employee} \), but its node labels are already contained in this relation. This query is evaluated with
nested form of amoeba joins using the query translation technique described in the previous section:

$$AJ_{task, list, task, employee} \quad AJ_{task, employee(task, employee), task, list} \quad (S_2)$$

which first retrieves a relation (task, employee), then finds task list nodes associated to this relation.

### 5.3 Database Integration

XML is a tree-structured data, however, a single tree is not sufficient to describe data models of the real world that often should be described as a graph structure. As we illustrated in Section 2, any graph-structured data model can be decomposed into several trees (relations). To integrate several trees into a single XML document, H. V. Jagadish et al. introduced the notion of colorful XML [12], which appends a color property to each XML node so that a projection of each colored tree represents one of the trees decomposed from a graph-structured data. However, the colorful XML requires a significant extension of the XML specification [6], and also to edit multiply colored XML data is quite difficult for standard text editors or simple script programs.

Our solution to this problem is to store several aspects of the data model separately in the form of XML data fragments, and to retrieve them using relational-style XML queries. These query results are joined using keys defined for XML. This approach does not require any extension of the XML specification. Figure 12 illustrates this approach. This XML data has some employee data (left), and associated office and section XML data (right) that wrap employees:

![Figure 12: Employee data (left) and additional information (office and section) described in two separated trees (right).](image)

These three XML fragments might be placed in the same XML document, or in different XML files. The colorful XML [12] merges these three XML fragments into a single tree while tolerating employee nodes with different colors. This method enables an XML query processor to traverse name, office, and section nodes from an employee node. Our solution to this problem is much simpler and leaves the XML data as they are, because a query for employee names, office and section can be expressed as a join operation of relations using employee@id values, described as follows:

$$(employee, name) \bowtie employee@id (office, employee) \bowtie employee@id (section, employee)$$

Let $R, S$ be relations, and $p$ be a node label for a join target, then a join operation $R \bowtie_p S$ is a selection $\sigma_{R.p=S.p}(R \times S)$. Therefore, without actually materializing a merged form of XML fragments, we can integrate the above XML data from the knowledge that employee@id values connect three relations; namely, employee@id is a key (or foreign key) for relations (employee, name), (office, employee), and (section, employee).

A key is a special case of an FD, and it can be used to uniquely locate XML nodes. In this example, we have the following key definitions for these three relations:

$$[\text{employee@id} \rightarrow \text{employee name}] \quad [\text{employee@id} \rightarrow \text{office employee}] \quad [\text{employee@id} \rightarrow \text{section employee}]$$

These keys (FDs) mean that an employee@id value is sufficient to uniquely locate all nodes in each relation (employee, name), (office, employee) and (section, employee). Buneman et al. have proposed keys for XML [7], however, their definition cannot handle structural variations. Our definition of FD allows both the cases that an office node is a child (descendant) of an employee node, or vice versa.

Integration of variously structured XML data is also useful for handling schema evolution. Figure 12 illustrates a process of enhancing employee data by appending supplementary information. Suppose that, first, we have only the employee name data, and subsequently these employees are assigned to some office and section, which is described as the right-hand side XML data in Figure 12. When creating a new database, it is usual that some data are missing or not available yet. With the capability to query variously structured XML data, schema evolution of XML databases can be managed with a simple join operation of several XML data. In addition, it is flexible to allow various XML structures in designing new XML data for enhancing existing databases.

Related to database integration, we mention several open problems that still need further study:

#### Handling Variations of Tag Names

There may be variations of tag names in describing the same data model in XML. For example, an XML tag employee may be named worker in another location. To handle these variations of tag names, one can use, for example, a simple mapping function that translates worker into employee or some dictionary that groups synonym words. In general, however, we have to consider a more difficult problem, called semantic integration [4], which needs to resolve semantic heterogeneity of XML tag names under specific paths.

#### Semantics of Nested Elements

When XML data has a recursive structure, its data semantics may be ambiguous. Figure 13 illustrates this problem; two name nodes are located under the manager node. To query a manager name, the amoeba condition $\langle manager, name \rangle$ cannot be used, since the manager is associated with its correct name Lucy as well as its employee’s name David unexpectedly. A solution to this problem is to clarify the data semantics by using XML namespace e.g., manager:name, employee:name, etc. XML attributes, such as manager@name, also can be used to avoid the problem of the semantic ambiguity. Although it is quite easy to capture the amoeba structure $\langle manager, manager:name \rangle$, the problem of automatic assignment of these namespace labels remains open.

![Figure 13: Clarifying semantics of the name tags by using XML namespace.](image)

### 5.4 Querying Incomplete Relations

Although the relational-style XML query manages structural variations of XML data, the user who only has a limited knowledge of
the underlying XML structure may fail to retrieve necessary information from the XML data. For example, a query for employee names who are working for active tasks can be described as follows:

(employee, name, (active, task)),

which has a nested query (active, task) to retrieve task nodes marked as active. This query has to find a relation (employee, name, active, task), but there is no matching tree structure for this relation in the XML data. In reality, many partially matching structures are available and would provide useful information.

To detect these partial matches, we present a query operation that collects incomplete relations allowing null values. For example, the query process of (employee, name, (active, task)) involves node pairs (4, 6, null, null), (20, null, 17, 18) and (29, null, null).

Figure 14 shows these nodes tuples. Then, to fill null values in these node pairs, we merge employee nodes 4, 20 and 29 by using equality of the employee@id value “e1”, and generate a node tuple ((4, 20, 29), 6, 17, 18) as one of the query results. In this query process, employee@id values work as object IDs of employee nodes.

We extend the definition of the amoeba join to tolerate null values in the query result:

Definition 5.3 \[ (A^J) \] Let NL be a list of node labels, and R be an input relation, an amoeba join allowing null values, denoted AI_{NL}(R), generates the same relation with an amoeba join A_{NL}(R), except that each result instance in AI_{NL}(R) is allowed to have null nodes other than the node corresponding a first node label in NL.

The A^J operator has a flavor of the outer join in relational databases, but is different in that A^J considers structural variations of input nodes.

Figure 14 illustrates a query schedule of (employee, name, (active, task)) that uses A^J operations instead of A J. First, to merge employee nodes using employee@id values, this schedule performs PC-join of these nodes (P1). Then, to retrieve task nodes that are marked active, we simply compute their amoeba join (P2). Among the inputs of the query, a pair of employee and task has a structural constraint imposed by the FD task -> employee, so we have to connect them by using a A^J operation (P3) allowing null values for the task nodes. In the similar manner, we perform A^J operation between employee and name to compose a relation (employee, name) (P4). The upper-right table in Figure 14 shows the intermediate query results up to (P4) phase. In (P5), employee nodes that have the same employee@id values are merged to fill the blank column in the table, and incomplete rows that still have null values are eliminated. Finally, using projection π, the query reports only requested nodes by the user, excluding employee@id column (P5), and the result is the lower-right table in Figure 14.

5.5 Amoeba Join Processing

The amoeba join processing depends on the capability to detect an ancestor-descendant relationships of two nodes, because to test an amoeba condition \((a, b, c)\), we need to check one of the nodes among \(a, b\) and \(c\) is a common ancestor of the others. If node \(a\) is a common ancestor in the amoeba structure, then the node \(a\) is an ancestor of nodes \(b\) and \(c\).

To make faster the detection of ancestor-descendant relationships, we use indexes that label each XML node with an interval \((\text{start, end})\) [14]. The tree structure of XML is encoded so that every interval of an ancestor node subsumes all its descendant nodes, and all intervals are disjoint. Using this node label, the detection of the ancestor-descendant relationship becomes a containment test of two intervals, i.e. a node \(p\) is an ancestor of a node \(q\) if \(p_{\text{start}} < q_{\text{start}} \land q_{\text{end}} < p_{\text{end}}\).

The details of the amoeba join algorithm are described in [19], thus we present its outline. The amoeba join can be processed efficiently by sorting input nodes in advance in the order of start values, since the root node of an amoeba always has the smallest start value. By sweeping the sorted input nodes, the amoeba join chooses a node \(p\) that has the smallest start value as a candidate of the root node of an amoeba. Then, for each input node list of the amoeba join except that contains \(p\), it searches the descendant nodes of \(p\) from range between \(p_{\text{start}}\) and \(p_{\text{end}}\) for the other components of the amoeba. After the search, this algorithm enumerates all amoeba structures rooted by \(p\), therefore, it sweeps the node \(p\) off from the input, then proceed to the next smallest node.

6. EXPERIMENTAL RESULTS

We evaluated the performance of the relational-style XML query to show the scalability of our method for various sizes of XML data, and the tolerance to structural variations.

Implementation. We implemented a prototype of our database management system in C++, which consists of several components, such as XML reader, index generator, query processor, etc. Our implementation of database indexes uses B+-trees provided by the Berkeley DB library [22]. On top of the B+-tree, we stored XML nodes labeled with (start, end, level, path ID, text), where the start and end are the interval labels [14] to efficiently detect ancestor-descendant relationships, and the level is the depth of a node in the XML tree, which is required to detect parent-child relationships of XML nodes. The path ID represents an ID assigned to each distinct path. The text is a text content encapsulated by tags or attributes.

XML nodes are stored in a B+-tree in ascending order of their start values. To make node retrieval faster, we also generated a secondary B+-tree index using a compound key (path ID, start), which aligns XML nodes first in the order of path IDs, then that of start values. This secondary index is used to efficiently locate nodes belonging to specific paths, e.g. //A, //A/B, etc.

Machine Environment. As a test vehicle, we used a Windows XP machine; dual Xeon 3.0GHz processors, 2GB memory and 250GB 7,200 rpm HDD.

Experimental Methodology. We run each query six times and take the average of the last five runs, because OS caches of the database files are quite different between the first run and the others. The standard deviation of the query performance is at most 0.02 (3σ = 0.06 seconds) or a far smaller value. It is sufficiently small to measure differences of the query performance.

Query Performance on XMark. To evaluate the query performance on standard XML data, we used XMark [21] benchmark.
program. We have changed its scalability parameter \( f \) from 0.1 to 1 to produce various sizes of XML data, which are almost 10M, 25M, 50M and 100M bytes. Figure 15 shows query schedules used in this experiment (\( Q_1 \) to \( Q_6 \)). This query set is designed so that the characteristics and scalability of the amoeba join algorithm become clear, so simple path queries and join (=) operation that can be processed with the standard techniques are not presented.

The XMark database contains 83 types of tag names. A relation in XML is a subset of these tag names. To detect FDs in the XMark data, we created a simple program that investigates one-to-many or one-to-one relationships that hold in the XMark data. For example, under the root node “site” in the XMark data, there are many person nodes, and each person node has many descendant interest nodes. These relationships correspond to FDs person \( \rightarrow \) site and interest \( \rightarrow \) person.

Query \( Q_1 \) and \( Q_2 \) are amoeba joins of two nodes that have one-to-many relationships. Figure 16 shows the performance of these queries and their result sizes. The performance of \( Q_1 \) and \( Q_2 \) scales in proportion to the XML data sizes.

Here, we present two examples that emphasize the significant benefit of query optimization. When more than two nodes involved in the amoeba join operation (\( Q_3 \)), its performance significantly deteriorates. Our implementation of the query processor does not use secondary storages to store intermediate results of a query. The permutation size of site, person and interest nodes is quite huge, and consequently the query Q3, which simply computes all possible tree structures consisting of these nodes, exhausted the main memory storage, and stopped after an out of memory error was observed. Query \( Q_{3F} \) is an optimized query schedule of \( Q_3 \) using the pushing-structural constraint technique described in Section 5, and the amoeba constraints derived from the FDs person \( \rightarrow \) site and person \( \rightarrow \) interest are pushed into the sub queries. Although both \( Q_3 \) and \( Q_{3F} \) has the same amoeba join operation \( \text{AJ}_{\text{site}, \text{person}, \text{site}} \), the performance of \( Q_{3F} \) scales well with increase in XML data sizes. This is because nested amoeba join queries in \( Q_{3F} \) construct appropriate tree structures in a bottom-up fashion, and efficiently avoids irrelevant tree structures. This result indicates that the right-hand schedule in Figure 11, which first processes an amoeba condition with more than two nodes, must be avoided. Query \( Q_4 \) and \( Q_{4F} \) are more complex examples of nested query schedules. To retrieve the relation (regions, item, mail, date), \( Q_4 \) considers an FD mail \( \rightarrow \) date in the path mail\text{date}, so PC-join can be used in this query. However, the relation (regions, item, mail, date) in the XMark data has several other FDs as shown in \( Q_{4S} \). Similar to the results of \( Q_1 \) and \( Q_{1F} \), computation of \( Q_4 \) could not be completed in the main memory, and \( Q_{4F} \), which considers all of these FDs, is scalable to the database size.

Query \( Q_5 \) and \( Q_{5F} \) show that amoeba join is not always slow; In XMark data, the mail object is a parent of two child nodes, 'from' and 'date', so the amoeba join of these nodes never reports incorrect results. In this case, the decomposed schedule \( Q_{5F} \) is less efficient due to the overhead of pipelining. Query \( Q_6 \), \( Q_{6F} \) and \( Q_{6S} \) retrieve nested relations in which each open\text{auction} node has current price information and several bidders associated with the bid time and amount of increase data. Query \( Q_6 \) misses the one-to-many relationship between open\text{auction} and bidder, so \( Q_{6F} \), which totally decomposes the schedule, becomes efficient. Considering that two relations (open\text{auction}, current) and (bidder, increase, time) comprise distinct objects, and are connected through an FD bidder \( \rightarrow \) open\text{auction}, we can produce a more efficient query schedule \( Q_{6S} \), which reduces the number of sub queries. This type of query optimization needs to be exploited but is left as a future work.

**Tolerance to Structural Variations.** To further study the tolerance of our method for variously structured XML data, we developed an XML data generator that produces three types of structural variations: simple, hierarchical, and random. Figure 17 illustrates these tree-structures generated from the same input table. The simple structure converts each row in the table into an XML fragment organized from the first column data to the last one. A column value in the input table is described as an XML attribute. The hierarchical structure aggregates column values that have the same value. For example, all values in the column \( a \) are aggregated into a single tag. This aggregation process is repeated recursively from column \( a \) to \( c \). This type of aggregation is frequently observed in the real-world XML data. The random structure is generated in almost the same manner with the hierarchical structure, but it randomly chooses target columns of aggregation, so the random XML data contains many structural variations. The generated XML data is a collection of a relation (a, b, c) that satisfies two FDs \( c \rightarrow b \) and \( b \rightarrow a \), representing two one-to-many relationships. The fanout parameter controls the number of associated nodes in these relationships. For example, when fanout = 5, each a node has 5 b nodes, and each b node has 5 c nodes. We programmed this data generator so that all three types of XML data consist of the same number of instances of the relation (a, b, c).

Figure 18 shows the query performance of \( Q_7 \) grouped by various query result sizes, and next by fanout values. Even in the presence of structural variations, the query performance between the simple and random format is stable. This characteristic is suited for integrating variously structured XML data. When the fanout parameter is between 2 to 100, the hierarchical data is more efficient for query processing, because it efficiently aggregates one-to-many relationships, and thus its database sizes are smaller than those of the others. However, when the fanout values are 500 and 1000, their query performance becomes slower. This is because our query processor expands the aggregated XML data into node tuples to report intermediate results, so many duplicate nodes are instantiated. For example, in Figure 17, a single a node in the hierarchical data is copied three times to generate intermediate node tuples. This inefficiency can be improved by holding intermediate results as a tree structure.

Our experiments demonstrate the scalability of our query optimization techniques to process queries of relatively large amount of results. If value conditions are involved, input data sizes of the amoeba join will be squeezed, so naive application of the amoeba join probably works well even for multiple input nodes. It still needs further study to estimate costs of amoeba join operations for various input data. Other than this cost estimation methodology, we can leverage the existing techniques of System R style query optimization on our XML algebra. In addition, the relational-style XML query provides independence of query statements from the underlying XML data structure. This property can be utilized to reorganize XML data structure for efficient query processing or minimizing database sizes. Although it might be possible to use relational databases as a storage scheme for relations in XML, it must have capabilities to query and store other XML nodes associated to relations.

7. RELATED WORK

The use of relational model to query complex structured data, including XML, has been studied in [16]. Our approach is unique in that it allows structural variations of XML data, and utilizes functional dependencies to capture data semantics of XML.

**Finding Relations in XML.** There have been several studies of the problem in finding relations in XML; Y. Li et al. [15] attempted
<table>
<thead>
<tr>
<th>Relation (Query Expression)</th>
<th>FD</th>
<th>Query Schedule</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Q_1$</td>
<td>(site, person)</td>
<td>person $\rightarrow$ site</td>
</tr>
<tr>
<td>$Q_2$</td>
<td>(person, interest)</td>
<td>interest $\rightarrow$ person</td>
</tr>
<tr>
<td>$Q_{1r}$</td>
<td>(site, person, interest)</td>
<td>interest $\rightarrow$ person, person $\rightarrow$ site</td>
</tr>
<tr>
<td>$Q_{2r}$</td>
<td>(regions, item, mail, date)</td>
<td>mail $\rightarrow$ date</td>
</tr>
<tr>
<td>$Q_3$</td>
<td>(mail, from, date)</td>
<td>mail $\rightarrow$ from date</td>
</tr>
<tr>
<td>$Q_6$</td>
<td>(open_auction, current, (bidder, increase, time))</td>
<td>open_auction $\rightarrow$ current, bidder $\rightarrow$ open_auction, bidder $\rightarrow$ increase time</td>
</tr>
<tr>
<td>$Q_{6r}$</td>
<td>(open_auction, current, (bidder, increase, time))</td>
<td>open_auction $\rightarrow$ current, bidder $\rightarrow$ open_auction, bidder $\rightarrow$ increase time</td>
</tr>
<tr>
<td>$Q_7$</td>
<td>(a, b, c)</td>
<td>c $\rightarrow$ b, b $\rightarrow$ a</td>
</tr>
</tbody>
</table>

Figure 15: Query schedules for retrieving relations with several FDs in XMark ($Q_1$ to $Q_{6r}$) and synthetic data set ($Q_7$).

Figure 16: Query performance (Left) and result sizes (Right) of $Q_1$ to $Q_{6r}$. Performance and result sizes of $Q_1$, $Q_4$ and $Q_6$ could not be measured due to out of memory errors.

Figure 17: Synthetic XML data of simple (left), hierarchical (center) and random (right) structures, generated from the same input table data.

Figure 18: Query performance of $Q_7$ for variously structured XML data, which have the same number of relations (a, b, c).
to extract particular patterns, containing the smallest least common ancestor (slea) of a given set of XML nodes. The slea, which was coined in [27], is a least common ancestor (lca) that contains no other lca nodes among its descendants. This definition of slea is an attempt to exclude the XML root node from query results. This is because XML is a single-rooted tree, and thus irrelevant nodes that never belong to the same relation may be connected through the root node. However, the slea approach is highly dependent on the query input. For example, when two unrelated nodes are the inputs of an slea query, the root node will be wrongly reported as a query result. The amoeba join [19] successfully avoids such unintentional results, since it does not rely on any additional lca nodes. However, the cost of enumerating all tree structures is prohibitive without the knowledge of functional dependencies. Query methods that retrieve XML structures without using knowledge of the schema or FDs do return incorrect results. Several such cases were presented in [23].

Another approach to querying variously structured XML data is to search the data to the level of ancestor or descendant nodes [2, 10] or nearest neighbor nodes [26]. However, these methods cannot address all possible tree structures derived from relational data. In addition, they are optimized for keyword-search queries, and are thus not suited to rigid database queries.

Functional Dependencies for XML. FDs and keys have been well studied to find ways of reducing data redundancy and avoiding update anomalies [17]. In recent years, these concepts have been applied to XML in the form of XML keys [7] and XML FDs [3, 13, 24, 28]. These approaches are based on paths; given sets X and Y of paths, an FD for XML is defined as $X \rightarrow Y$. However, these path-based definitions of FDs cannot handle XML documents containing structural variations, which require multiple path expressions.

In summary, previous work on FDs for XML [3, 7, 13, 24, 28] inferred FDs from a path structure of an XML document. In contrast, our approach that assumes FDs are defined outside the XML data, and are specified using node names (e.g., tag or attribute names) on a relation, rather than on paths. Unlike path-based definitions, our definition of FD allows various XML data expressions, and therefore makes the design of XML databases much easier.

8. CONCLUSIONS

The presence of structural variations is a serious problem for the traditional XML query processors, because path-expression queries are dependent to the underlying XML tree structures. We overcome this problem by introducing the relational-style XML query, which uses the notion of a relation in XML that allows amoeba structures. In addition, to capture the data semantics implied in the XML structure, we incorporated the well-known notion of functional dependencies into XML, and devised efficient query processing techniques for retrieving relations satisfying FDs. With these capabilities, we can utilize heterogeneous XML structures to design and integrate several XML databases. The contributions described in this paper include:

- The notion of the relation in XML. With this capability, FDs and keys are smoothly incorporated into XML.
- A class of XML structures, called a tree relation, which can be used as an XML counterpart of relational tables.
- A departure from path-expression queries. XML structures of interest are automatically determined from a set of FDs.
- Capability of integrating variously structured XML data.
- Experimental results that confirm the scalability and tolerance of our query method in the presence of structural variations.

Repeatability Assessment Result

All the results (except $Q_3$ to $Q_{10}$) in this paper were verified by the SIGMOD repeatability committee. Results of query $Q_3$ to $Q_{10}$ were added after the submission of the code in order to reflect a reviewer’s comment. Code and data used in the paper are available at http://www.sigmod.org/codearchive/sigmod2008/.

9. REFERENCES