Integrating Association Rule Mining with Relational Database Systems: Alternatives and Implications

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Abstract

Data mining on large data warehouses is becoming increasingly important. In support of this trend, we consider a spectrum of architectural alternatives for coupling mining with database systems. These alternatives include: loose-coupling through a SQL cursor interface; encapsulation of a mining algorithm in a stored procedure; caching the data to a file system on-the-fly and mining; tight-coupling using primarily user-defined functions; and SQL implementations for processing in the DBMS. We comprehensively study the option of expressing the mining algorithm in the form of SQL queries using Association rule mining as a case in point. We consider four options in SQL-92 and six options in SQL enhanced with object-relational extensions (SQL-OR). Our evaluation of the different architectural alternatives shows that from a performance perspective, the Cache option is superior, although the performance of the SQL-OR option is within a factor of two. Both the Cache and the SQL-OR approaches incur a higher storage penalty than the loose-coupling approach which performance-wise is a factor of 3 to 4 worse than Cache. The SQL-92 implementations were too slow to qualify as a competitive option. We also compare these alternatives on the basis of qualitative factors like automatic parallelization, development ease, portability and interoperability. As a byproduct of this study, we identify some primitives for native support in database systems for decision-support applications.

1 Introduction

An ever increasing number of organizations are installing large data warehouses using relational database technology. There is a huge demand for mining nuggets of knowledge from these data warehouses.

The initial research on data mining was primarily concentrated on defining new mining operations and developing algorithms for them. Most early mining systems were developed largely on file systems and specialized data structures and buffer management strategies were devised for each algorithm. Coupling with database systems was at best loose, and access to data in a DBMS was provided through an ODBC or SQL cursor interface (e.g. [Int96, AAB+96, HFK+96, IM96]).

This paper is an attempt to understand the implications of various architectural alternatives for coupling data mining with relational database systems. In particular, we are interested in studying how competitive

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can a mining computation expressed in SQL be compared to a specialized implementation of the same mining operation.

There are several potential advantages of a SQL implementation. One can make use of the database indexing and query processing capabilities thereby leveraging more than a decade of effort in making these systems robust, portable, scalable, and concurrent. One can also exploit the underlying SQL parallelization, particularly in an SMP environment. The DBMS support for checkpointing and space management can be valuable for long-running mining algorithms.

![SQL architecture for mining in a DBMS](image)

Figure 1: SQL architecture for mining in a DBMS

The architecture we have in mind is schematically shown in Figure 1. We visualize that the desired mining operation will be expressed in some extension of SQL or a graphical language. A preprocessor will generate appropriate SQL translation for this operation. We consider translations that can be executed on a SQL-92 [MS92] relational engine. We also consider translations that take advantage of some of the newer object-relational capabilities being designed for SQL [MM96]; specifically, we assume availability of blobs, user-defined functions, and table functions [PR98]. We do not assume mining specific extensions in the underlying relational engine; however, identification of such extensions is a secondary goal of this study.

We compare the performance of the above SQL architecture with the following alternatives:

**Read directly from DBMS:** Data is read tuple by tuple from the DBMS to the mining kernel using a cursor interface. Data is not copied to a file system. We consider two variations of this approach. One is the loose-coupling approach where the DBMS runs in a different address space from the mining process. This is the approach followed by most existing mining systems. A potential problem with this approach is the high context switching cost between the DBMS and the mining process [AS96b]. In spite of the block-read optimization present in many systems (e.g. Oracle [Ora92], DB2 [Cha96]) where a block of tuples is read at a time, the performance could suffer. The second is the stored-procedure approach where the mining algorithm is encapsulated as a stored procedure [Cha96] that runs in the same address space as the DBMS. The main advantage of both these approaches is greater programming flexibility and no extra storage requirement. Also, any previous file system code can be easily transferred to work on data stored in the DBMS. The mined results are stored back into the DBMS.

**Cache-mine:** This option is a variation of the Stored-procedure approach where after reading the entire data once from the DBMS, the mining algorithm temporarily caches the relevant data in a lookaside buffer on a local disk. The cached data could be transformed to a format that enables efficient future accesses. The cached data is discarded when the execution completes. This method has all the advantages of the stored procedure approach plus it promises to have better performance. The disadvantage is that it requires additional disk space for caching. Note that the permanent data continues to be managed by the DBMS.

**User-defined function (UDF):** The mining algorithm is expressed as a collection of user-defined functions (UDFs) [Cha96] that are appropriately placed in SQL data scan queries. Most of the processing happens in the UDF and the core DBMS engine is used primarily to provide tuples to the UDFs. Little use is made of
the query processing capability of the DBMS. The UDFs are run in the unfenced mode (same address space as the database). Such an implementation was presented in [AS96b]. The main attraction of this method over Stored-procedure is the execution time since passing tuples to a stored procedure is slower than passing it to a UDF. Otherwise, the processing happens in almost the same manner as in the stored procedure case. The main disadvantage is the development cost since the entire mining algorithm has to be written as UDFs involving significant code rewrites [AS96b].

We do both quantitative and qualitative comparisons of the architectures stated above with respect to the problem of discovering Boolean Association rules [AIS93] defined as follows:

Given a set of transactions, where each transaction is a set of items, an association rule is an expression \( X \implies Y \), where \( X \) and \( Y \) are sets of items. The intuitive meaning of such a rule is that the transactions that contain the items in \( X \) tend to also contain the items in \( Y \). An example of such a rule might be that “30% of transactions that contain beer also contain diapers; 2% of all transactions contain both these items”. Here 30% is called the confidence of the rule, and 2% the support of the rule. The problem of mining association rules is to find all rules that satisfy a user-specified minimum support and minimum confidence.

This basic framework has been extended to handle hierarchies on items [SA95] and time-based orderings on purchases for discovering sequential patterns [SA96]. We concentrate on the basic boolean associations problem for the most part. In Section 5.4 we discuss these extensions.

We compare the performance of these different approaches using four real-life datasets. We also use synthetic datasets to better understand the behavior of different algorithms. Our conclusions about the comparison of different architectures are based on the current associations algorithms and DBMS technology (DB2/UDB version 5). If the algorithms are made significantly faster or the DBMS performance characteristics change drastically these conclusions should be re-evaluated.

1.1 Related work

Lately researchers have started to focus on issues related to integrating mining with databases. There have been language proposals to extend SQL to support mining operators. For instance, the query language D-MQL [HFK+96] extends SQL with a collection of operators for mining characteristic rules, discriminant rules, classification rules, association rules, etc. The M-SQL language [IVA96] extends SQL with a special unified operator Mine to generate and query propositional rules. Another example is the Mine Rule [MPC96] operator for a generalized version of the association rule discovery problem. Query flocks [TAC+98] for association rule mining also discusses ways of generalizing association rule in the database context. Our focus in this paper is on the performance of various architectures. We do not consider the issues of the language constructs required to extend SQL with mining features.

The issue of tightly coupling a mining algorithm with a relational database system from a systems point of view was addressed in [AS96b]. This proposal makes use of user-defined functions (UDFs) in SQL statements to selectively push parts of the computation into the database system. The SETM algorithm [HS95] for finding association rules was expressed in the form of SQL queries. However, as shown in [AMS+96], SETM is not efficient and there are no results reported on running it against a relational DBMS. Another approach to integrating association rule mining with a DBMS is presented in [RIC97]. We discuss this work later in the paper. Issues in coupling classification algorithms with a database system have been discussed in [M.W98, S.C98, GUS98].
1.2 Paper Layout

The rest of the paper is organized as follows. In Section 2, we present the overview of the SQL implementations. In Sections 3 and 4, we discuss different ways of doing the support counting phase of Associations in SQL — Section 3 presents SQL-92 implementations whereas Section 4 gives implementations in SQL-OR. In Section 5 we present performance comparison of the different architectural alternatives using real-life and synthetic datasets. We also include qualitative comparisons of the different architectures along the dimensions of development and maintenance ease, storage and memory requirements, portability, inter-operability, and parallelizability. In Section 6, we propose primitives in a relational DBMS that we believe would be useful for a large class of mining algorithms. We conclude with a summary of results and directions for future work in Section 7.

2 Associations in SQL

The association rule mining problem can be decomposed into two subproblems [AIS93]:

- Find all combinations of items, called frequent itemsets, whose support is greater than minimum support.
- Use the frequent itemsets to generate the desired rules. The idea is that if, say, $ABCD$ and $AB$ are frequent, then the rule $AB \implies CD$ holds if the ratio of support($ABCD$) to support($AB$) is at least as large as the minimum confidence. Note that the rule will have minimum support because $ABCD$ is frequent.

We first present a brief survey of existing file-based algorithms for finding frequent itemsets. We then address various issues that arise when implementing these algorithms in SQL including input-output representations (Section 2.2), frequent itemset computation (Section 2.3) and rule generation (Section 2.4).

2.1 Existing algorithms

2.1.1 Apriori Algorithm

We use the Apriori algorithm [AMSS+96] as the basis for our presentation. The Apriori algorithm for finding frequent itemsets makes multiple passes over the data. In the $k$th pass it finds all itemsets having $k$ items called the $k$-itemsets. Each pass consists of two phases. Let $F_k$ represent the set of frequent $k$-itemsets, and $C_k$ the set of candidate $k$-itemsets (potentially frequent itemsets). First, is the candidate generation phase where the set of all frequent $(k-1)$-itemsets, $F_{k-1}$, found in the $(k-1)$th pass, is used to generate the candidate itemsets $C_k$. The candidate generation procedure ensures that $C_k$ is a superset of the set of all frequent $k$-itemsets. A specialized in-memory hash-tree data structure is used to store $C_k$. Then, data is scanned in the support counting phase. For each transaction, the candidates in $C_k$ contained in the transaction are determined using the hash-tree data structure and their support count is incremented. At the end of the pass, $C_k$ is examined to determine which of the candidates are frequent, yielding $F_k$. The algorithm terminates when $F_k$ or $C_{k+1}$ becomes empty.

2.1.2 Other algorithms

A number of variants [SON95, ZPOL97, Toi96, BMUT97] of this basic algorithm have been proposed. Most of them have the same basic dataflow structure as the Apriori algorithm. Our goal in this work is to understand how best to integrate this basic structure within a database system. Therefore, instead of discussing each of these
variants in detail we only focus on some improvements that might have an impact on our overall architectural comparison. One class of algorithms [Toi96, BMU'T97] reduce the number of data passes that in Apriori is equal to the maximum length of a frequent itemset (or one more). We discuss a promising sampling based algorithm presented in [Toi96].

2.1.3 Sampling-based approach

This approach first selects a small sample of the dataset and uses that to find the frequent itemsets using Apriori. The frequent sets are then augmented with their negative border that consists of infrequent itemsets all of whose subsets are frequent. The entire database is then used to find the exact support values for itemsets in the augmented set. If no itemset from the negative border becomes frequent, the algorithm is done having found the frequent items in just one pass of the entire database. Otherwise, one or more subsequent passes may be necessary to find support count of all newly discovered candidate itemsets.

We analyze the work savings through this sampling algorithm vis-a-vis Apriori. First, we compare the number of itemsets counted against the entire dataset. The best case for the sampling algorithm is when the sample finds the true frequent set. In this case, the itemset augmented with the negative border consists of the union of candidate itemsets in all passes of Apriori. Thus, the itemsets counted by Sampling is a super set of the itemsets counted by Apriori. There is no saving on the number of itemsets to be counted. The only saving is the reduction in the number of passes. The sample helps to identify the potential candidate itemsets for all levels in advance, hence a single pass (in the best case) can be used to count all frequent itemsets. This also implies that the one data pass of the Sampling algorithm will require at least the maximum of the time required by any single Apriori pass.

2.2 Input-output formats

Input format  The input is a transaction table $T$ with two column attributes: transaction identifier (tid) and item identifier (item). For a given tid, typically there are multiple rows in the transaction table corresponding to different items in the transaction. The number of items per transaction is variable and unknown during table creation time. Thus, alternative schemas may not be convenient. In particular, assuming that all items in a transaction appear as different columns of a single tuple (e.g. [RIC97]) is not practical because often the number of items per transaction can be more than the maximum number of columns that the database supports. For instance, for one of our real-life datasets the maximum number of items per transaction is 872 and for another it is 700. In contrast, the corresponding average number of items per transaction is only 9.6 and 4.4 respectively.

Output format  The output is a collection of frequent itemsets and rules of varying length. The maximum length of the frequent itemsets and rules is much smaller than the number of items and is rarely more than a dozen. Therefore, a frequent itemset is represented as a tuple in a fixed-width table where the extra column values are set to NULL to accommodate smaller itemsets. The schema of an itemset is (item$_1, \ldots, $item$_k$, support) and for a rule is (tail$_1, \ldots, tail_{k-1}, head_2, \ldots, head_{k-1}, support, confidence$) where $k$ is the size of the largest frequent itemset. For instance, if $k = 5$, the frequent itemset ABCD which has 30% support and the rule $AB \implies CD$ which has 90% confidence are represented by the tuples $(A, B, C, D, NULL, 0.3)$ and $(A, B, NULL, NULL, C, D, NULL, NULL, 0.9, 0.3)$ respectively. Note that for rules a more compact representation would be (item$_1, \ldots, $item$_k$, rule, confidence, support) where rule denotes the separation between the tail and head items. We preferred the more redundant former schema because it made subsequent
queries on the rules table considerably simpler.

2.3 Computing frequent itemsets in SQL

In each pass \( k \) we first generate a candidate itemset set \( C_k \) from frequent itemsets \( F_{k-1} \) of the previous pass and then count the support of itemsets in \( C_k \) using transaction table \( T \). We discuss SQL implementations of these steps in Sections 2.3.1 and 2.3.2 respectively.

2.3.1 Candidate generation

Given \( F_{k-1} \), the set of all frequent \((k-1)\)-itemsets, the Apriori candidate generation procedure [AMS+96] returns a superset of the set of all frequent \( k \)-itemsets. We assume that the items in an itemset are lexicographically ordered. Since, all subsets of a frequent itemset are also frequent, we can generate \( C_k \) from \( F_{k-1} \) as follows:

In the join step, a superset of the candidate itemsets \( C_k \) is generated by joining \( F_{k-1} \) with itself:

insert into \( C_k \) select 1.\textit{item}_1, \ldots, 1.\textit{item}_{k-1}, 2.\textit{item}_{k-1}
from \( F_{k-1} \) 1.\textit{item}, \( F_{k-1} \) 2.\textit{item}
where 1.\textit{item}_1 = 2.\textit{item}_1 and

\[ \vdots \]

1.\textit{item}_{k-2} = 2.\textit{item}_{k-2} and 1.\textit{item}_{k-1} < 2.\textit{item}_{k-1}

For example, let \( F_3 \) be \{\{1 2 3\}, \{1 2 4\}, \{1 3 4\}, \{1 3 5\}, \{2 3 4\}\}. After the join step, \( C_4 \) will be \{\{1 2 3 4\}, \{1 3 4 5\}\}. Next, in the prune step, all itemsets \( c \in C_k \), where some \((k-1)\)-subset of \( c \) is not in \( F_{k-1} \), are deleted. Continuing with the example above, the prune step will delete the itemset \{1 3 4 5\} because the subset \{1 4 5\} is not in \( F_3 \). We will then be left with only \{1 2 3 4\} in \( C_4 \).

We can perform the prune step in the same SQL statement as the join step by writing it as a \( k \)-way join as shown in Figure 2. A \( k \)-way join is used since for any \( k \)-itemset there are \( k \) subsets of length \((k-1)\) for which \( F_{k-1} \) needs to be checked for membership. The join predicates on \( I_1 \) and \( I_2 \) remain the same. After the join between \( I_1 \) and \( I_2 \) we get a \( k \) itemset consisting of \((I_1.\textit{item}_1, \ldots, I_1.\textit{item}_{k-1}, I_2.\textit{item}_{k-1})\). For this itemset, two of its \((k-1)\)-length subsets are already known to be frequent since it was generated from two itemsets in \( F_{k-1} \). We check the remaining \( k-2 \) subsets using additional joins. The predicates for these joins are enumerated by skipping one item at a time from the \( k \)-itemset as follows: We first skip \( \textit{item}_1 \) and check if subset \((I_1.\textit{item}_2, \ldots, I_1.\textit{item}_{k-1}, I_2.\textit{item}_{k-1})\) belongs to \( F_{k-1} \) as shown by the join with \( I_3 \) in the figure. In general, for a join with \( I_r \), we skip \( \textit{item}_{r-2} \). Figure 3 gives an example for \( k = 4 \). We construct a primary index on \((\textit{item}_1, \ldots, \textit{item}_{k-1})\) of \( F_{k-1} \) to efficiently process these \( k \)-way joins using index probes.

\( C_k \) need not always be materialized before the counting phase. Instead, the candidate generation can be pipelined with the subsequent SQL queries used for support counting.

2.3.2 Counting support

This is the most time-consuming part of the association rules algorithm. We consider two different categories of SQL implementations:

(A) The first one is based purely on SQL-92. We discuss four approaches in this category in Section 3.
The second utilizes object-relational extensions like UDFs, BLOBs (Binary large objects) and table functions. Table functions [PR98] are virtual tables associated with a user defined function which generate tuples on the fly. They have pre-defined schemas like any other table. The function associated with a table function can be implemented as a UDF. Thus, table functions can be viewed as UDFs that return a collection of tuples instead of scalar values.

We discuss six approaches in this category in Section 4. UDFs in this approach are lightweight and do not require extensive memory allocations and coding like the UDF architectural option.

2.4 Rule generation

In order to generate rules having minimum confidence, minconf, we first find all non-empty proper subsets of every frequent itemset \( I \). Then, for each subset \( m \), we find the confidence of the rule \( m \Rightarrow (I - m) \) and output the rule if it is at least \( \text{minconf} \).

In the support counting phase, the frequent itemsets of size \( k \) are stored in table \( F_k \). Before the rule generation phase, we merge all the frequent itemsets into a single table \( F \) having the schema discussed earlier in Section 2.2.

We use a table function \( \text{GenRules} \) to generate all possible rules from a frequent itemset. The input argument to the function is a frequent itemset \( \text{item}_1, \ldots, \text{item}_k \) and its support. The output is a set of rules of the form \( (tail_1, \ldots, tail_{k-1}, head_1, \ldots, head_{k-1}, support) \) where the \( tail_1, \ldots, tail_{k-1} \) part consists of all non-empty proper subsets of that itemset with NULLs used to pad smaller subsets as described in Section 2.2. The output is joined with \( F \) to find the support of the antecedent (tail part) and the confidence of the rule is calculated by taking the ratio of the supports of the full frequent itemset and the antecedent as shown in the query below. Figure 4 shows the query diagramatically.

We can also do rule generation without using table functions and base it purely on SQL-92. The rules are generated in a level-wise manner where in each level \( k \) we generate rules with consequents of size \( k \). Further, we make use of the property that for any frequent itemset, if a rule with consequent \( c \) holds then so do rules with consequents that are subsets of \( c \) as suggested in [AMS+96]. We can use this property to generate rules in level \( k \) using rules with \( (k-1) \) long consequents found in the previous level, much like the way we did candidate generation in Section 2.3.1.

The fraction of the total running time spent in rule generation is very small. Therefore, we do not discuss
3 Support counting using SQL-92

We studied four approaches in this category — KwayJoin, 3wayJoin, 2GroupBy and Subquery.

3.1 K-way joins

In each pass $k$, we join the candidate itemsets $C_k$ with $k$ transaction tables $T$ and follow it up with a group by on the itemsets as shown in Figure 5. The figure 5 also shows a tree diagram of the query. These tree diagrams should not be confused with the plan trees that could look quite different.

\[
\begin{align*}
\text{insert into } F_k & \text{ select } \text{item}_1, \ldots, \text{item}_k, \text{count}(*) \\
\text{from} & \quad C_k, \ T \ t_1, \ldots, \ T \ t_k \\
\text{where} & \quad \text{t}_1.\text{item} = C_k.\text{item}_1 \text{ and} \\
& \quad \vdots \\
& \quad \text{t}_k.\text{item} = C_k.\text{item}_k \text{ and} \\
& \quad \text{t}_1.\text{tid} = \text{t}_2.\text{tid} \text{ and} \\
& \quad \vdots \\
& \quad \text{t}_{k-1}.\text{tid} = \text{t}_{k}.\text{tid} \\
\text{group by} & \quad \text{item}_1, \text{item}_2, \ldots, \text{item}_k \\
\text{having} & \quad \text{count}(*) > :\text{minsup} \\
\end{align*}
\]

This SQL computation, when merged with the candidate generation step, is similar to the one proposed in [TAC+98] as a possible mechanism to implement query flocks.

For pass 2 we use a special optimization where instead of materializing $C_2$, we replace it with the 2-way joins between the $F_1$s as shown in the candidate generation phase in section 2.3.1. This saves the cost of materializing $C_2$ and also provides early filtering of the $T$s based on $F_1$ instead of the larger $C_2$ which is almost a cartesian product of the $F_1$s. In contrast, for other passes corresponding to $k > 2$, $C_k$ could be smaller than $F_{k-1}$ because of the prune step.
3.2 Three-way joins

The above approach requires \((k + 1)\)-way joins in the \(k\)th pass. We can reduce the cardinality of joins to 3 using the following approach which bears some resemblance to the AprioriTid algorithm in [AMS+96]. Each candidate itemset \(C_k\), in addition to attributes \((item_1, \ldots, item_k)\) has three new attributes \((oid, id_1, id_2)\). \(oid\) is a unique identifier associated with each itemset and \(id_1\) and \(id_2\) are \(oids\) of the two itemsets in \(F_{k-1}\) from which the itemset in \(C_k\) was generated (as discussed in Section 2.3.1). In addition, in the \(k\)th (for \(k > 1\)) pass we generate a new copy of the data table \(T_k\) with attributes \((tid, oid)\) that keeps for each tid the oid of each itemset in \(C_k\) that it supported. For support counting, we first generate \(T_k\) from \(T_{k-1}\) and \(C_k\) and then do a group-by on \(T_k\) to find \(F_k\) as follows:

\[
\text{insert into } T_k \text{ select } t_1.\text{tid, oid} \\
\text{from } C_k, T_{k-1} t_1, T_{k-1} t_2 \\
\text{where } t_1.\text{oid} = C_k.id_1 \text{ and } t_2.\text{oid} = C_k.id_2 \text{ and } t_1.\text{tid} = t_2.\text{tid} \\
\]

\[
\text{insert into } F_k \text{ select oid, item}_1, \ldots \text{ item}_k, \text{count(*)} \\
\text{from } C_k, \\
\quad (\text{select oid as cid, count(*) as cut from } T_k) \\
\quad \text{group by oid having count(*) > :minsup} \quad \text{as temp} \\
\quad \text{where } C_k.\text{oid} = \text{cid} \\
\]

3.3 Subquery-based

This approach makes use of common prefixes between the itemsets in \(C_k\) to reduce the amount of work done during support counting. The support counting phase is split into a cascade of \(k\) subqueries. The \(l\)-th subquery \(Q_l\) (see Figure 6) finds all tids that match the distinct itemsets formed by the first \(l\) columns of \(C_k\) (call it \(d_l\)). The output of \(Q_l\) is joined with \(T\) and \(d_{l+1}\) (the distinct itemsets formed by the first \(l + 1\) columns of \(C_k\)) to get \(Q_{l+1}\). The final output is obtained by a group-by on the \(k\) items to count support as above. Note that the final “select distinct” operation on the \(C_k\) when \(l = k\) is not necessary.

For pass-2 the special optimization of the KwayJoin approach is used.

3.4 Two group-bys

This approach avoids the multi-way joins used in the previous approaches, by joining \(T\) and \(C_k\) based on whether the “item” of a \((tid, item)\) pair of \(T\) is equal to any of the \(k\) items of \(C_k\). Then, do a group by on \((item_1, \ldots, item_k, tid)\) filtering tuples with count equal to \(k\). This gives all (itemset, tid) pairs such that the tid supports the itemset. Finally, as in the previous approaches, do a group-by on the itemset \((item_1, \ldots, item_k)\) filtering tuples that meet the support condition. For pass-2, we apply the same optimization as in the KwayJoin approach where we avoid materializing \(C_2\).

\[
\text{insert into } F_k \text{ select item}_1, \ldots \text{item}_k, \text{count(*)} \\
\text{from } (\text{select item}_1, \ldots \text{item}_k, \text{count(*)} \\
\quad \text{from } T, C_k \\
\quad \text{where } \text{item} = C_k.\text{item}_1 \text{ or} \\
\quad \quad \vdots \\
\]

\[9\]
insert into $F_k$ select $item_1, \ldots, item_k$, count(*)
from (Subquery $Q_k$) $t$
group by $item_1, item_2 \ldots item_k$
having count(*) > $\minsup$

Subquery $Q_l$ (for any $l$ between 1 and $k$):
select $item_1, \ldots, item_k$, tid
from $T$ $t_l$, (Subquery $Q_{l-1}$) as $r_{l-1}$,
(select distinct $item_1, \ldots, item_l$ from $C_k$) as $d_l$
where $r_{l-1}.item_1 = d_l.item_1$ and $\ldots$ and
$r_{l-1}.item_{l-1} = d_l.item_{l-1}$ and
$r_{l-1}.tid = t_l.tid$ and
$t_l.item = d_l.item_l$

Figure 6: Support counting using subqueries

<table>
<thead>
<tr>
<th>Datasets</th>
<th># Records in millions</th>
<th># Transactions in millions</th>
<th># Items in thousands</th>
<th>Avg.#Items per transaction (R/T)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dataset-A</td>
<td>2.5</td>
<td>0.57</td>
<td>85</td>
<td>4.4</td>
</tr>
<tr>
<td>Dataset-B</td>
<td>7.5</td>
<td>2.5</td>
<td>15.8</td>
<td>2.62</td>
</tr>
<tr>
<td>Dataset-C</td>
<td>6.6</td>
<td>0.21</td>
<td>15.8</td>
<td>31</td>
</tr>
<tr>
<td>Dataset-D</td>
<td>14</td>
<td>1.44</td>
<td>480</td>
<td>9.62</td>
</tr>
</tbody>
</table>

Table 1: Description of different real-life datasets.

item = $C_k.item_k$
group by $item_1, \ldots, item_k$, tid
having count(*) = $k$ as temp
group by $item_1, \ldots, item_k$
having count(*) > $\minsup$

3.5 Performance comparison of SQL-92 approaches

In this section we compare the performance of the four SQL-92 approaches. Our experiments were performed on Version 5 of IBM DB2 Universal Server installed on a RS/6000 Model 140 with a 200 MHz CPU, 256 MB main memory and a 9 GB disk with a measured transfer rate of 8 MB/sec.

We use four real-life datasets obtained from mail-order companies and retail stores for the experiments. These datasets differ in the values of parameters like the number of (tid, item) pairs, number of transactions (tids), number of items and the average number of items per transaction. Table 1 summarizes characteristics of these datasets.

For these experiments we built a composite index $(item_1 \ldots item_k)$ on $C_k$, $k$ different indices on each of the $k$ items of $C_k$ and a (tid, item) and a (item, tid) index on the data table. The goal was to let the optimizer
Figure 7: Comparison of four SQL-92 approaches. The time taken is broken down by each pass and the candidate generation time.

choose the best plan possible. We do not include the index building cost in the total time.

In Figure 7 we show the total time taken by the four approaches: KwayJoin, 3wayJoin, Subquery and 2GroupBy. For comparison, we also show the time taken by the Loose-coupling approach because this is the approach used by existing systems. The graph shows the total time split into candidate generation time (Cgen) and the time for each pass. The open bars in the figure correspond to approaches that did not complete even within the maximum time value on the y-axis. From these set of experiments we can make the following observations:

- Overall, the best approach in the SQL-92 category is the Subquery approach. An important reason for its superior performance over the KwayJoin approach is the exploitation of common prefixes between candidate itemsets. Although the Subquery approach is comparable or better than the Loose-coupling approach in some cases, for other cases involving low support values it did not complete even after taking ten times more time than the Loose-coupling approach. For instance, for Dataset-C for the high support value of 2% the Subquery approach is better than the Loose-coupling approach. But as we decrease the support to 1% the Subquery approach becomes almost ten times worse than the Loose-coupling approach and on further decreasing the support to 0.25% none of the SQL-92 approaches could be taken to completion.
within 20 times the Loose-coupling time and finally got killed due to storage overflow.

- The 2GroupBy approach is significantly worse than the other two approaches because it involves an index-Oring operation on $k$ indices for each pass $k$ of the algorithm. In addition, the inner group-by requires sorting a large intermediate relation. The outer group-by is comparatively faster because the sorted result is of size at most $C_k$ which is much smaller than the result size of the inner group-by. The DBMS does aggregation during sorting therefore the size of the result is an important factor in the total cost.

- The 3wayJoin approach is worse than the KwayJoin approach because it cannot use the pass-2 optimization of KwayJoin and it requires writing large intermediate relations. As shown in [AMS+96] there might be other datasets especially ones where there is significant reduction in the size of $T_k$ as $k$ increases where 3wayJoin might perform better than KwayJoin. One disadvantage of the 3wayJoin approach is that it requires space to store and log the temporary relations $T_k$ generated in each pass.

- Based on these experiments we can also extrapolate on how the Sampling approach discussed in Section 2.1.3 will affect the performance. As observed in Section 2.1.3 by using sampling, for each approach the time taken will be greater than the maximum of the time taken by a single pass. For the Loose-coupling approach the Sampling approach would take close to the second pass time whereas for the Sampling version of the Subquery approach (best performing SQL-92 approach) the time taken would be dominated by the second or third pass time as seen in Figure 7. The other SQL-92 approaches compare even less favourably with Loose-coupling as we use sampling.

The important conclusion to draw from this study, is that implementations based on pure SQL-92 are too slow to be considered a general-purpose alternative to the existing Loose-coupling approach.

4 Support counting using SQL with object-relational extensions

In this section, we study approaches that use object-relational extensions in SQL to improve performance. We first consider an approach we call GatherJoin and its three variants in Section 4.1. Next we present a very different approach called Vertical in Section 4.2. Finally, in Section 4.3 we present an approach based on SQL-bodied functions. For each approach, we also outline a cost-based analysis of the execution time to choose between these different approaches. In Section 4.4 we present performance comparisons.

4.1 GatherJoin

The GatherJoin approach (see Figure 8) generates all possible $k$-item combinations of items contained in a transaction, joins them with the candidate table $C_k$, and counts the support of the itemsets by grouping the join result. It uses two table functions Gather and Comb-K. The data table $T$ is scanned in the $(tid, item)$ order and passed to the table function Gather, which collects all the items of a transaction in memory and outputs a record for each transaction. Each record consists of two attributes: the tid and item-list which is a collection of all items in a field of type VARCHAR or BLOB. The output of Gather is passed to another table function Comb-K which returns all $k$-item combinations formed out of the items of a transaction. A record output by Comb-K has $k$ attributes $T_{i1tm_1}, \ldots, T_{i1tm_k}$, which can be used to probe into the $C_k$ table. An index is constructed on all the items of $C_k$ to make the probe efficient.

This approach is analogous to the KwayJoin approach except that we have replaced the $k$-way self join of $T$ with the table functions Gather and Comb-K. These table functions are easy to code and do not require a large
amount of memory. It is also possible to merge them into a single table function GatherComb-K, which is what we did in our implementation. Note that the Gather function is not required when the data is already in a horizontal format where each tid is followed by a collection of all its items.

```
insert into F_k select item_1, ..., item_k, count(*)
from C_k,
(select t_2.T_item_1, ..., t_2.T_item_k from T,
table (Gather(T.tid, T.item)) as t_1,
table (Comb-K(t_1.tid, t_1.item-list)) as t_2)
where t_2.T_item_1 = C_k.item_1 and
... t_2.T_item_k = C_k.item_k
group by C_k.item_1, ..., C_k.item_k
having count(*) > xminsup
```

Figure 8: Support Counting by GatherJoin

Special pass 2 optimization: For k = 2, the 2-candidate set C_2 is simply a join of F_1 with itself. Therefore, we can optimize the pass 2 by replacing the join with C_2 by a join with F_1 before the table function (see Figure 9). The table function now gets only frequent items and generates significantly fewer 2-item combinations. We apply this optimization to other passes too. However, unlike pass 2 we still have to do the final join with C_k and therefore the benefit is not as significant.

```
insert into F_2 select t_2.T_item_1, t_2.T_item_2, count(*)
from (select * from T, F_1 where
T.item = F_1.item_1) as t_1,
table (GatherComb-2(t_1.tid,item)) as t_2)
group by t_2.T_item_1, t_2.T_item_2
having count(*) > xminsup
```

Figure 9: Support Counting by GatherJoin in the second pass

4.1.1 Variations of GatherJoin approach

GatherCount: One variation of the GatherJoin approach for pass two is the GatherCount approach where we perform the group-by inside the table function GatherComb-2. We will refer to this extended table function as Gather-Cnt. The candidate 2-itemset C_2 is represented as a two dimensional array (as suggested in [AMS+96]) inside function Gather-Cnt. Instead of outputting the 2-item combinations, the function uses the combinations to directly update support counts in memory and outputs only the frequent 2-itemsets, F_2 and their support after the last transaction.

The attraction of this option is the absence of the outer grouping. The UDF code is small since it only needs to maintain a 2D array. We could apply the same trick for subsequent passes but the coding becomes considerably more complicated because of the need to maintain hash-tables to index the C_k. The disadvantage of this approach is that it can require a large amount of memory to store C_2. If enough memory is not available,
$C_2$ needs to be partitioned and the process has to be repeated for each partition. Another problem with this approach is that it cannot be automatically parallelized.

**GatherPrune:** A problem with the **GatherJoin** approach is the high cost of joining the large number of item combinations with $C_k$. We can push the join with $C_k$ inside the table function and thus reduce the number of such combinations. $C_k$ is converted to a BLOB and passed as an argument to the table function.

The cost of passing the BLOB for every tuple of $R$ can be high. In general, we can reduce the parameter passing cost by using a smaller Blob that only approximates the real $C_k$. The trade-off is increased cost for other parts notably grouping because not as many combinations are filtered. A problem with this approach is the increased coding complexity of the table function.

**Horizontal:** This is another variation of **GatherJoin** that first uses the **Gather** function to transform the data to the horizontal format but is otherwise similar to the **GatherJoin** approach. Rajamani et al. [RIC97] propose finding associations using a similar approach augmented with some pruning based on a variation of the **GatherPrune** approach. Their results assume that the data is already in a horizontal format which is often not true in practice. They report that their SQL implementation is two to six times slower than a UDF implementation.

<table>
<thead>
<tr>
<th>$R$</th>
<th>number of records in the input transaction table</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T$</td>
<td>number of transactions</td>
</tr>
<tr>
<td>$N$</td>
<td>average number of items per transaction = $\frac{R}{T}$</td>
</tr>
<tr>
<td>$F_1$</td>
<td>number of frequent items</td>
</tr>
<tr>
<td>$S(C)$</td>
<td>sum of support of each itemset in set $C$</td>
</tr>
<tr>
<td>$R_f$</td>
<td>number of records out of $R$ involving frequent items = $S(F_1)$</td>
</tr>
<tr>
<td>$N_f$</td>
<td>average number of frequent items per transaction = $\frac{R_f}{T}$</td>
</tr>
<tr>
<td>$C_k$</td>
<td>number of candidate k-itemsets</td>
</tr>
<tr>
<td>$C(N, k)$</td>
<td>number of combinations of size $k$ possible out of a set of size $n = \frac{n!}{k!(n-k)!}$</td>
</tr>
<tr>
<td>$s_k$</td>
<td>cost of generating a $k$ item combination using table function Comb-k</td>
</tr>
<tr>
<td>group($n$, $m$)</td>
<td>cost of grouping $n$ records out of which $m$ are distinct</td>
</tr>
<tr>
<td>join($n$, $m$, $r$)</td>
<td>cost of joining two relations of size $n$ and $m$ to get a result of size $r$</td>
</tr>
<tr>
<td>blob($n$)</td>
<td>cost of passing a BLOB of size $n$ integers as an argument</td>
</tr>
</tbody>
</table>

**Table 2: Notations used for cost analysis of different approaches**

### 4.1.2 Cost analysis of GatherJoin and its variants

The relative performance of these variants depends on a number of data characteristics like the number of items, total number of transactions, average length of a transaction etc. We express the costs in each pass in terms of parameters that are known or can be estimated after the candidate generation step of each pass. The purpose of this analysis is to help us choose between the different options. Therefore, instead of including all I/O and CPU costs, we include only those terms that help us distinguish between different options. We use the notations of Table 2 in the cost analysis.

The cost of **GatherJoin** includes the cost of generating $k$-item combinations, joining with $C_k$ and grouping to count the support. The number of $k$-item combinations generated, $T_k$ is $C(N, k) * T$. Join with $C_k$ filters out the non-candidate item combinations. The size of the join result is the sum of the support of all the candidates denoted by $S(C_k)$. The actual value of the support of a candidate itemset will be known only after the support
counting phase. However, we approximate it to the minimum of the support of all its \((k - 1)\)-subsets in \(F_{k-1}\). The total cost of the GatherJoin approach is:

\[
T_k * s_k + \text{join}(T_k, C_k, S(C_k)) + \text{group}(S(C_k), C_k), \text{ where } T_k = C(N, k) * T
\]

The above cost formula needs to be modified to reflect the special optimization of joining with \(F_1\) to consider only frequent items. We need a new term \(\text{join}(R, F_1, R_f)\) and need to change the formula for \(T_k\) to include only frequent items \(N_f\) instead of \(N\).

For the second pass, we do not need the outer join with \(C_k\). The total cost of GatherJoin in the second pass is:

\[
\text{join}(R, F_1, R_f) + T_2 * s_2 + \text{group}(T_2, C_2), \text{ where } T_2 = C(N_f, 2) * T \approx \frac{N_f^2 * T}{2}
\]

Cost of GatherCount in the second pass is similar to that for basic GatherJoin except for the final grouping cost:

\[
\text{join}(R, F_1, R_f) + \text{group, internal}(T_2, C_2) + F_2 * s_2
\]

In this formula, “group, internal” denotes the cost of doing the support counting inside the table function.

For GatherPrune the cost equation is:

\[
R * \text{blob}(k * C_k) + S(C_k) * s_k + \text{group}(S(C_k), C_k).
\]

We use \(\text{blob}(k * C_k)\) for the BLOB passing cost since each itemset in \(C_k\) contains \(k\) items.

The cost estimate of Horizontal is similar to that of GatherJoin except that here the data is materialized in the horizontal format before generating the item combinations.

### 4.2 Vertical

We first transform the data table into a vertical format by creating for each item a BLOB containing all tids that contain that item (Tid-list creation phase) and then count the support of itemsets by merging together these tid-lists (support counting phase). This approach is similar to the approaches in [SON95, ZPOL97]. For creating the Tid-lists we use a table function Gather. This is the same as the Gather function in GatherJoin except that we create the tid-list for each frequent item. The data table \(T\) is scanned in the (item, tid) order and passed to the function Gather. The function collects the \(tids\) of all tuples of \(T\) with the same item in memory and outputs a (item, tid-list) tuple for items that meet the minimum support criterion. The tid-lists are represented as BLOBs and stored in a new TidTable with attributes (item, tid-list).

In the support counting phase, for each itemset in \(C_k\) we want to collect the tid-lists of all \(k\) items and use a UDF to count the number of \(tids\) in the intersection of these \(k\) lists. The tids are in the same sorted order in all the tid-lists and therefore the intersection can be done efficiently by a single pass of the \(k\) lists. This step can be improved by decomposing the intersect operation to share these operations across itemsets having common prefixes as follows.

We first select distinct \((item_1, item_2)\) pairs from \(C_k\). For each distinct pair we first perform the intersect operation to get a new result-tid-list, then find distinct triples \((item_1, item_2, item_3)\) from \(C_k\) with the same first two items, intersect result-tid-list with tid-list for \(item_3\) for each triple and continue with \(item_4\) and so on until all \(k\) tid-lists per itemset are intersected. This approach is analogous to the Subquery approach presented for SQL-92.

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The above sequence of operations can be written as a single SQL query for any $k$ as shown in Figure 10. The final intersect operation can be merged with the count operation to return a count instead of the tid-list — we do not show this optimization in the query of Figure 10 for simplicity.

```sql
insert into $F_k$ select $item_1, \ldots, item_i$, count(tid-list) as cnt
from (Subquery $Q_k$) t where cnt > $\minsup$
```

Subquery $Q_l$ (for any $l$ between 2 and $k$):

```sql
select $item_1, \ldots, item_i$,
Intersect($r_{l-1}.tid-list, t_i.tid-list$) as tid-list
from TidTable $t_i$, (Subquery $Q_{l-1}$) as $r_{l-1}$,
(select distinct $item_1, \ldots, item_i$ from $C_k$) as $d_i$
where $r_{l-1}.item_1 = d_i.item_1$ and \ldots and
$r_{l-1}.item_{i-1} = d_i.item_{i-1}$ and
$t_i.item = d_i.item_i$
```

Subquery $Q_1$: (select * from TidTable)

**Special pass 2 optimization:** For pass 2 we need not generate $C_2$ and join the TidTables with $C_2$. Instead, we perform a self-join on the TidTable using predicate $t_1.item < t_2.item$.

```sql
insert into $F_k$ select $t_1.item, t_2.item$, cnt
from (select $t_1.item, t_2.item$, CountIntersect($t_1.tid-list, t_2.tid-list$) as cnt
from TidTable $t_1, TidTable t_2$
where $t_1.item < t_2.item$) as t
where cnt > $\minsup$
```

**4.2.1 Cost analysis**

The cost of the Vertical approach during support counting is dominated by the cost of invoking the UDFs and intersecting the tid-lists. The UDF is first called for each distinct item pair in $C_k$, then for each distinct item triple and so on. Let $d_j^k$ be the number of distinct $j$ item tuples in $C_k$ Then the number of UDF invocations is $\sum_{j=2}^{k} d_j^k$. In each invocation two BLOBs of tid-list are passed as arguments. The UDF intersects the tid-lists by a merge pass and hence the cost is proportional to $2 \times \text{average length of a tid-list}$. The average length of a tid-list can be approximated to $\frac{R_L}{F_1}$. Note that with each intersect the tid-list keeps shrinking. However, we ignore such effects for simplicity.

The total cost of the Vertical approach is:

$$
\sum_{j=2}^{k} d_j^k \times \left( 2 \times \text{Blob} \left( \frac{R_L}{F_1} \right) + \text{Intersect} \left( \frac{2R_L}{F_1} \right) \right)
$$

In the above formula Intersect$(n)$ denotes the cost of intersecting two tid-lists with a combined size of $n$. We are not including the join costs in this analysis because it accounted for only a small fraction of the total cost.
The total cost of the second pass is:

\[
C_2 \times \{ 2 \times \text{Blob} \left( \frac{R_f}{F_1} \right) + \text{Intersect} \left( \frac{2R_f}{F_1} \right) \}
\]

### 4.3 SQL-bodied functions: SBF

This approach is based on SQL-bodied procedures commonly known as SQL/PSM [MM96]. SQL/PSMs extend SQL with additional control structures. We will make use of such constructs for `do .. end`.

We use the `for` construct to scan the transaction table \( T \) in the \((\text{tid}, \text{item})\) order. Then, for each tuple \((\text{tid}, \text{item})\) of \( T \), we update those tuples of \( C_k \) that contain one matching item. \( C_k \) is extended with 3 extra attributes \((\text{prevTid}, \text{match}, \text{supp})\). The \text{prevTid} attribute keeps the \text{tid} of the previous tuple of \( T \) that matched that itemset. The \text{match} attribute contains the number of items of \text{prevTid} matched so far and \text{supp} holds the current support of that itemset.

```sql
for this as select * from \( T \) do
  update \( C_k \) set prevTid = tid,
  match = case when tid = prevTid then match+1 else 1 end,
  supp = case when match = k-1 and tid = prevTid then supp+1 else supp end
  where item = \text{item}_1\ or
  \vdots
  item = \text{item}_k
end for;
insert into \( F_k \) select \text{item}_1, \ldots, \text{item}_k, \text{supp}
  from \( C_k \) where supp > :\text{minsupp}
```

We omit a cost analysis of this approach because it was not competitive compared to other approaches.

### 4.4 Performance comparison of SQL-OR approaches

We studied the performance of six SQL-OR approaches using the datasets summarized in Table 1. Figure 11 shows the results for only four approaches: \text{GatherJoin}, \text{GatherCount}, \text{GatherPrune} and \text{Vertical}. For the other two approaches \((\text{Horizontal and SBF})\) the running times were so large that we had to abort the runs in many cases. The reason why the \text{Horizontal} approach was significantly worse than the \text{GatherJoin} approach was the time to transform the data to the horizontal format. For instance, for \text{Dataset-C} it was 3.5 hours which is almost 20 times more than the time taken by \text{Vertical} for 2% support. For \text{Dataset-B} the process was aborted after running for 5 hours. After the transformation, compared to \text{GatherJoin} the time taken by \text{Horizontal} was also significantly worse when run without the frequent itemset filtering optimization. However with the optimization the performance was comparable. The \text{SBF} approach had significantly worse performance because of the expensive indexing ORing of the \( k \) join predicates. Another problem with this approach is the large number of updates to the \( C_k \) table. In DB2, all of these updates are logged resulting in severe performance degradation.

We first concentrate on the overall comparison between the different approaches. Then we will compare the approaches based on how they perform in each pass of the algorithm.

The \text{Vertical} approach has the best overall performance and it is sometimes more than an order of magnitude better than the other three approaches.
Figure 11: Comparison of four SQL-OR approaches: Vertical, GatherPrune, GatherJoin and GatherCount on four datasets for different support values. The time taken is broken down by each pass and an initial “prep” stage where any one-time data transformation cost is included.
The majority of the time of the Vertical approach is spent in transforming the data to the Vertical format in most cases (shown as “prep” in figure 11). The vertical representation is like an index on the item attribute. If we think of this time as a one-time activity like index building then performance looks even better. The time to transform the data to the Vertical format was much smaller than the time for the horizontal format although both formats write almost the same amount of data. The reason is the difference in the number of records written. The number of frequent items is often two to three orders of magnitude smaller than the number of transactions.

Between GatherJoin and GatherPrune, neither strictly dominates the other. The special pass-2 optimization in GatherJoin had a big impact on performance. With this optimization, for Dataset-B with support 0.1%, the running time for pass 2 was reduced from 5.2 hours to 10 minutes.

When we compare these approaches based on time spent in each pass no single approach emerges as “the best” for all passes with the different datasets.

For pass three onwards, Vertical is often two or more orders of magnitude better than the other approaches. Even in cases like Dataset-B, support 0.01% where it spends three hours in the second pass, the total time for next two passes is only 14 seconds whereas it is more than an hour for the other two approaches. For higher passes, the performance degrades dramatically for GatherJoin, because the table function Gather-Comb-K generates a large number of combinations. For instance, for pass 3 of Dataset-C even for support value of 2% pass 3 did not complete after 5.2 hours whereas for Vertical pass 3 finished in 0.2 seconds. GatherPrune is better than GatherJoin for third and later passes. For pass 2 GatherPrune is worse because the overhead of passing a large object as an argument dominates the cost.

The Vertical approach sometimes spends too much time in the second pass. In some of these cases the GatherJoin approach was better in the second pass (for instance for low support values of Dataset-B) whereas in other cases (for instance, Dataset-C with minimum support 0.25%) GatherCount was the only good option. In the latter case, both GatherPrune and GatherJoin did not complete after more than six hours for pass 2. Further, they caused a storage overflow error because of the large size of the intermediate results to be sorted. We had to divide the dataset into four equal parts and ran the second pass independently on each partition to avoid this problem.

![Figure 12: Effect of increasing transaction length (average number of items per transaction)](image)

Two factors that affect the choice amongst the Vertical, GatherJoin and GatherCount approaches in different passes and pass 2 in particular are: number of frequent items ($F_1$) and the average number of frequent items per transaction ($N_f$). From Figure 11 we notice that as the value of the support is decreased for each dataset causing the size of $F_1$ to increase, the performance of pass 2 of the Vertical approach degrades rapidly. This
trend is also clear from our cost formulae. The cost of the **Vertical** approach increases quadratically with $F_1$. **GatherJoin** depends more critically on the number of frequent items per transaction. For **Dataset-B** even when the size of $F_1$ increases by a factor of 10, the value of $N_f$ remains close to 2, therefore the time taken by **GatherJoin** does not increase as much. However, for **Dataset-C** the size of $N_f$ increases from 3.2 to 10 as the support is decreased from 2.0% to 0.25% causing **GatherJoin** to deteriorate rapidly. From the cost formula for **GatherJoin** we notice that the total time for pass 2 increases almost quadratically with $N_f$.

We validated this observation further by running experiments on synthetic datasets for varying values of the number of frequent items per transaction. We used the synthetic dataset generator described in [AMS+96] for this purpose. We varied the transaction length, the number of transactions and the support values while keeping the total number of records and the number of frequent items fixed. In Figure 12 we show the total time spent in pass 2 of the **Vertical** and **GatherJoin** approaches. As the number of items per transaction (transaction length) increases, the cost of **Vertical** remains almost unchanged whereas the cost of **GatherJoin** increases.

### 4.5 Final hybrid approach

The previous performance section helps us draw the following conclusions: Overall, the **Vertical** approach is the best option especially for higher passes. When the size of the candidate itemsets is too large, the performance of the **Vertical** approach could suffer. In such cases, **GatherJoin** is a good option as long as the number of frequent items per transaction ($N_f$) is not too large. When $N_f$ is large **GatherCount** may be the only good option even though it may not easily parallelize.

The hybrid scheme chooses the best of the three approaches **GatherJoin**, **GatherCount** and **Vertical** for each pass based on the cost estimates outlined in Sections 4.1.2 and 4.2.1. The parameter values used for the estimation are available at the end of the previous pass. In Section 5 we plot the final running time for the different datasets based on this hybrid approach.

### 5 Architecture comparisons

In this section our goal is to compare the five alternatives: **Loose-coupling, Stored-procedure, Cache, UDF**, and the best **SQL** implementation.

For **Loose-coupling**, we use the implementation of the Apriori algorithm [AMS+96] for finding association rules provided with the IBM data mining product, Intelligent Miner [Int96]. For **Stored-procedure**, we extracted the Apriori implementation in Intelligent Miner and created a stored procedure out of it. The stored procedure is run in the unfenced mode in the database address space. For **Cache**, we used an option provided in Intelligent Miner that causes the input data to be cached as a binary file after the first scan of the data from the DBMS. The data is copied in the horizontal format where each item is followed by an encoding of all its frequent items. For the **UDF-architecture**, we use the UDF implementation of the Apriori algorithm described in [AS96b]. In this implementation, first a UDF is used to initialize state and allocate memory for candidate itemsets. Next, for each pass a collection of UDFs are used for generating candidates, counting support, and checking for termination. These UDFs access the initially allocated memory, address of which is passed around in BLOBs. Candidate generation creates the in-memory hash-trees of candidates. This happens entirely in the UDF without any involvement of the DBMS. During support counting, the data table is scanned sequentially and for each tuple a UDF is used for updating the counts on the memory resident hash tree.
Figure 13: Comparison of four architectures: Cache, Stored-procedure, UDF and SQL on four real-life datasets. Loose-coupling is similar to Stored-procedure. For each dataset three different support values are used. The total time is broken down by the time spent in each pass.
5.1 Timing comparison

In Figure 13, we show the performance of Cache, Stored-procedure, UDF and the hybrid SQL-OR implementation for the datasets in Table 1. We do not show the times for the Loose-coupling option because its performance was very close to the Stored-procedure option.

We can make the following observations:

- **Cache** has the best or close to the best performance in all cases. 80-90% of its total time is spent in the first pass where data is accessed from the DBMS and cached in the file system. Compared to the SQL approach this approach is a factor of 0.8 to 2 times faster.

- The **Stored-procedure** approach is the worst. The difference between Cache and Stored-procedure is directly related to the number of passes. For instance, for Dataset-A the number of passes increases from two to three when decreasing support from 0.5% to 0.35% causing the time taken to increase from two to three times. The time spent in each pass for Stored-procedure is the same except when the algorithm makes multiple passes over the data since all candidates could not fit in memory together. This happens for the lowest support values of Dataset-B, Dataset-C and Dataset-D.

- UDF is similar to Stored-procedure. The only difference is that the time per pass decreases by 30-50% for UDF because of closer coupling with the database.

- The SQL approach comes second in performance after the Cache approach for low support values and is even somewhat better for high support values. The cost of converting the data to the vertical format for SQL is typically lower than the cost of transforming data to binary format outside the DBMS for Cache. However, after the initial transformation subsequent passes take negligible time for Cache. For the second pass SQL takes significantly more time than Cache particularly when we decrease support. For subsequent passes even the SQL approach does not spend too much time. Therefore, the difference between Cache and SQL is not very sensitive to the number of passes because both approaches spend negligible time in higher passes.

The SQL approach is 1.8 to 3 times better than Stored-procedure or Loose-coupling approach. As we decreased the support value so that the number of passes over the dataset increases, the gap widens.

Note that we could have implemented Stored-procedure using the same hybrid algorithm that we used for SQL instead of using the 1M algorithm. Then, we expect the performance of Stored-procedure to improve because the number of passes of the data will decrease. However, we will pay the storage penalty of making additional copy of the data as we did in the Cache approach. The performance of Stored-procedure cannot be better than Cache because as we have observed that most of the time of Cache is spent in the first pass which cannot be changed for Stored-procedure.

We now analyze the effect of Sampling (refer Section 2.1.3) on each of the four alternatives. The total time of the Cache approach remains almost unchanged. The time taken by the Stored-procedure and UDF approaches is in most cases close to the maximum of the time spent in a single pass which happens to be the second pass in our case. The time of the SQL approach remains almost unchanged because most of the time is spent either in the second pass which cannot be reduced by sampling or in the first pass when we create the vertical bit maps. The bitmap creation time cannot be reduced because without the bitmaps the time spent in itemset counting for subsequent passes becomes much more than the bitmap creation time. In summary, with sampling the Stored-procedure and UDF approaches can be made to be close to the Cache approach except in cases like the lower support values of Dataset-B, Dataset-C and Dataset-D where pass-2 required multiple data scans.
5.1.1 Scale-up experiment

Our experiments with the four real-life datasets above has shown the scaling property of the different approaches with decreasing support value and increasing number of frequent itemsets. We experiment with synthetic datasets to study other forms of scaling: increasing number of transactions and increasing average length of transactions. Figure 14 shows how Stored-procedure, Cache and SQL scale with increasing number of transactions, UDF and Loose-coupling have similar scale-up behavior as Stored-procedure, therefore we do not show these approaches in the figure. We used a dataset with 10 average number of items per transaction, 100 thousand total items and a default pattern length (defined in [AMS+96]) of 4. Thus, the size of the dataset is 10 times the number of transactions. As the number of transactions is increased from 10K to 3000K the time taken increases proportionately. The largest frequent itemset was 5 long. This explains the five fold difference in performance between the Stored-procedure and the Cache approach. Figure 15 shows the scaling when the transaction length changes from 3 to 50 while keeping the number of transactions fixed at 100K. All three approaches scale linearly with increasing transaction length.

5.1.2 Impact of longer names

In these experiments we assumed that the tids and item ids are all integers. Often in practice these are character strings longer than four bytes. Longer names need more storage and cost more during comparisons. This could hurt all four of the alternatives. For the Stored-procedure, UDF and Cache approach the time taken to read data from the DBMS will increase. Intelligent Miner [Int96] maps all character fields to integers using an in-memory hash-table. Since bulk of the time is spent in reading data and mapping them to integers we expect the time of these three alternatives to increase. For the SQL approach we cannot assume an in-memory hash-table for doing the mapping. We propose how to efficiently do the mapping in SQL without creating expensive extra copies of the data.

We discuss the impact of longer names only on the winning hybrid approach. For mapping the item names we use table $F_1$ — after the first pass we assign unique integer identifiers to the frequent items. Since for subsequent passes of all horizontal approaches like GatherJoin (see Figure 9) we anyway join the transaction table $T$ with $F_1$, we can get integer item ids from the join. Since $F_1$ is much smaller than $T$, the storage overhead of this mapping is negligible. For mapping the transaction ids ($tids$), we pipeline the original data table through a table function in the $tid$ order. The table function remembers the previous $tid$ and maintains
a counter. Every time the \textit{tid} changes, the counter is incremented. This counter value is the mapping assigned to each \textit{tid}. In the \textit{Vertical} approach we need to do the \textit{tid} mapping only once before creating the TidTable and therefore we can pipeline these two steps. The mapping never needs to be stored explicitly. After these two transformations, the tid and item fields are integers for all the remaining steps of the algorithm including candidate generation and rule generation.

By mapping the fields this way, we expect longer names to have similar performance impact on all of our architectural options.

5.2 Space overhead of different approaches

In Figure 16 we summarize the space required for three options: \textit{Stored-procedure}, \textit{Cache} and SQL. We assume that the tids and items are integers. The space requirements for UDF and \textit{Loose-coupling} is the same as that for \textit{Stored-procedure} which in turn is less than the space needed by the \textit{Cache} and SQL approaches. The \textit{Cache} and SQL approaches have comparable storage overheads. For \textit{Stored-procedure} and UDF we do not need any extra storage for caching. However, all three options \textit{Cache}, \textit{Stored-procedure} and UDF require data in each pass to be grouped on the \textit{tid}. In a relational DBMS we cannot assume any order on the physical layout of a table, unlike in a file system. Therefore, we need either an index on the data table or need to sort the table every time to ensure a particular order. Let \( R \) denote the total number of (tid, item) pairs in the data table. Either option has a space overhead of \( 2 \times R \) integers. The \textit{Cache} approach caches the data in an alternative binary format where each tid is followed by all the items it contains. Thus, the size of the cached data in \textit{Cache} is at most: \( R + T \) integers where \( T \) is the number of transactions. For SQL we use the hybrid \textit{Vertical} option. This requires creation of an initial TidTable of size at most \( I + R \) where \( I \) is the number of items. Note that this is slightly less than the cache required by the \textit{Cache} approach. The SQL approach needs to sort data in pass 1 in all cases and pass 2 in some cases where we used the \textit{GatherJoin} approach instead of the \textit{Vertical} approach. This explains the large space requirement for \textit{Dataset-B}.

In summary, the UDF and \textit{Stored-procedure} approaches require the least amount of space followed by the \textit{Cache} and the SQL approaches which require roughly as much extra storage as the data. When the item-ids or tids are character strings instead of integers, the extra space needed by \textit{Cache} and SQL is a much smaller fraction of the total data size because before caching we always convert item-ids to their compact integer representation and store in binary format.

5.3 Summary of comparison between different architectures

We present a summary of the pros and cons of the different architectures on each of the following yardsticks: (a) performance (execution time); (b) storage overhead; (c) potential for automatic parallelization; (d) development and maintenance ease; (e) portability (f) inter-operability.

In terms of execution time, the \textit{Cache} approach is the best option. The SQL approach is a close second — it is always within a factor of two of \textit{Cache} for all of our experiments and is sometimes even slightly better. The UDF approach is better than the \textit{Stored-procedure} approach by 30\textit{Stored-procedure} and \textit{Cache}, the performance difference is a function of the number of times the data is scanned — if we make four scans of the data the \textit{Stored-procedure} approach is four times slower than \textit{Cache}. With sampling we can reduce the number of scans required by the UDF and \textit{Stored-procedure} options to the maximum required in any one pass of Apriori. Thus, we can hope to make the \textit{Stored-procedure} and UDF approaches close to the \textit{Cache} option except in cases where a single algorithm pass required multiple scans of the data because of memory shortage as we observed for some
Figure 16: Comparison of different architectures on space requirements. The first part refers to the space used in caching data and the second part refers to any temporary space used by the DBMS for sorting or alternately for constructing indices to be used during sorting. The size of the data is the same as the space utilization of the Stored-procedure approach.

of the lower support values on our datasets.

In terms of space requirements, the Cache and the SQL approach lose to the UDF or the Stored-procedure approach. The Cache and SQL approaches have similar storage requirements.

The SQL implementation has the potential for automatic parallelization particularly on a SMP machine. Parallelization could come for free for SQL-92 queries. Unfortunately, the SQL-92 option is too slow to be a candidate for parallelization. The stumbling block for automatic parallelization using SQL-OR could be queries involving UDFs that use scratch pads. The only such function in our queries is the Gather table function. This function essentially implements a user defined aggregate, and would have been easy to parallelize if the DBMS provided support for user defined aggregates or allowed explicit control from the application on how to partition the data amongst different parallel instances of the function. On a MPP machine, although one could rely on the DBMS to come up with a data partitioning strategy, it might be possible to better tune performance if the application could provide hints about the best partitioning [AS96a]. Further experiments are required to assess how the performance of these automatic parallelizations would compare with algorithm-specific parallelizations (e.g. [AS96a]).

The development time and code size using SQL could be shorter if one can get efficient implementations out of expressing the mining algorithms declaratively using a few SQL statements. Thus, one can avoid writing and debugging code for memory management, indexing and space management all of which are already provided in a database system. We illustrate this point further by showing SQL implementations of extensions of the basic Boolean association rules problems with hierarchies and sequential patterns [SA96] in Section 5.4. However, there are some detractors to easy development using the SQL alternative. First, any attached UDF code will be harder to debug than stand-alone C++ code due to lack of debugging tools. Second, stand-alone code can be debugged and tested faster when run against flat file data. Running against flat files is typically a factor of five to ten faster compared to running against data stored in DBMS tables. Finally, some mining algorithms (e.g. neural-net based) might be too awkward to express in SQL.
The ease of porting of the SQL alternative depends on the kind of SQL used. Within the same DBMS, porting from one OS platform to another requires porting only the small UDF code and hence is easy. In contrast the Stored-procedure and Cache alternatives require porting larger lines of code. Porting from one DBMS to another could get hard for SQL approach, if non-standard DBMS-specific features are used. For instance, our preferred SQL implementation relies on the availability of DB2’s table functions, for which the interface is still not standardized across other major DBMS vendors. Also, if different features have different performance characteristics on different database systems, considerable tuning would be required. In contrast, the Stored-procedure and Cache approach are not tied to any DBMS specific features. The UDF implementation has the worst of both worlds — it is large and is tied to a DBMS.

One attraction of SQL implementation is inter-operability and usage flexibility. The adhoc querying support provided by the DBMS enables flexible usage and exposes potential for pipelining the input and output operators of the mining process with other operators in the DBMS. However, to exploit this feature one needs to implement the mining operators inside the DBMS. This would require major rework in existing database systems. The SQL approach presented here is based on embedded SQL and as such does not provide operator pipelining and inter-operability. Queries on the mined result is possible with all four alternatives as long as the mined results are stored back in the DBMS.

5.4 Extensibility of SQL approaches
In this section we demonstrate the extensibility of the SQL framework by showing how the SQL approaches can be modified to mine generalized association rules and sequential patterns.

5.4.1 Generalized association rules

Often real-life applications have taxonomies (is-a hierarchies) over items, Figure 17 shows an example taxonomy that indicates that Pepsi is-a soft drink is-a beverage and so on. We represent the taxonomy as a table Tax with the schema (parent, child). Each record in Tax corresponds to an edge in the taxonomy DAG(Directed Acyclic Graph). The antecedent and consequent of an association rule can contain items from any level of the taxonomy (for example, soft drinks $\implies$ snacks).

![Diagram of a sample taxonomy and the corresponding taxonomy table](image-url)

The best known algorithm [SA95] for finding frequent itemsets with hierarchies has the same basic structure as Apriori (see Section 2.1.1) for boolean associations, except for the following two differences. (1) For counting support, we augment each transaction with all ancestors of its items. (2) From the candidate itemsets we prune all itemsets that contain an item and its ancestor. We discuss how these extensions can be reflected in our SQL implementations.
We first use the taxonomy table above to precompute ancestors of items.

**Pre-computing ancestors** We call \( \hat{x} \) an ancestor of \( x \) if there is a directed path from \( \hat{x} \) to \( x \) in \( \text{Tax} \). We use the transitive closure operation in SQL3 as shown in Figure 18 for the ancestor computation. The result of the query is stored in table \( \text{Ancestor} \) having the schema \((\text{ancestor}, \text{descendant})\).

```sql
INSERT INTO Ancestor WITH R-Tax (ancestor, descendant) AS
    (SELECT parent, child FROM Tax UNION ALL
     SELECT p.ancestor, c.child FROM R-Tax p, Tax c
     WHERE p.descendant = c.parent)
    SELECT ancestor, descendant FROM R-Tax
```

![Figure 18: Pre-computing Ancestors](image)

**Candidate pruning** The candidate generation can be formulated as explained in Section 2.2.3 except that we need to prune from \( C_k \) itemsets containing an item and its ancestor. [SA95] proves that this pruning needs to be done only in the second pass (for \( C_2 \)). In the SQL formulation as shown in Figure 19, we prune all \((\text{ancestor}, \text{descendant})\) pairs from \( C_2 \) which is generated by joining \( F_1 \) with itself.

```sql
INSERT INTO C_2 (SELECT I_1.item_1, I_2.item_1 FROM F_1 I_1, F_1 I_2
    WHERE I_1.item_1 < I_2.item_1) EXCEPT
    (SELECT ancestor, descendant FROM Ancestor UNION
     SELECT descendant, ancestor FROM Ancestor)
```

![Figure 19: Generation of \( C_2 \)](image)

**Support counting** All the support counting approaches for boolean association rules presented in Sections 3 and 4 can be extended to handle taxonomies by replacing the original transaction table \( T \) with \( T^* \) that is \( T \) augmented to include \((\text{tid}, \text{item})\) entries for all ancestors of items appearing in \( T \). This can be formulated as a SQL query as shown in Figure 20. The \text{select distinct} clause is used to eliminate duplicate records due to extension of items with a common ancestor.

**Query to generate \( T^* \)**

```sql
SELECT item, tid FROM T UNION
SELECT distinct A.ancestor as item, T.tid
FROM T, Ancestor A
WHERE A.descendant = T.item
```

![Figure 20: Transaction extension subquery](image)

When counting support we do not materialize \( T^* \). Instead we use the SQL support for common subexpressions (with construct) to pipeline the generation of \( T^* \) with the join operations. For instance, for the \text{KwayJoin} join query (section 3) the query would be modified as follows:

```sql
INSERT INTO F_k WITH T^*(tid, item) AS (Query for \( T^* \))
    (KwayJoin query with \( T \) replaced by \( T^* \))
```

Thus, we notice that the introduction of hierarchies on itemsets required very few changes to the SQL implementations of the algorithms.

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5.4.2 Sequential patterns

Given a set of data-sequences each of which is a list of transactions ordered by the transaction time, the problem of mining sequential patterns is to discover all sequences with a user-specified minimum support. Each transaction contains a set of items. A sequential pattern is an ordered list (sequence) of itemsets. The itemsets that are contained in the sequence are called elements of the sequence. For example, \(((\text{computer}, \text{modem})(\text{printer}))\) is a sequence with two elements \((- (\text{computer}, \text{modem}) \text{ and } (\text{printer}))\). The support of a sequential pattern is the number of data-sequences that contain the sequence. A sequential pattern can be further qualified by specifying maximum and/or minimum time gaps between adjacent elements and a sliding time window within which items are considered part of the same sequence element. These time constraints are specified by three parameters, \(\text{max-gap, min-gap and window-size}\). The basic structure of the algorithm [SA96] for sequential pattern mining is again similar to that of the Apriori algorithm. We discuss the differences and SQL formulations to handle them next.

**Input-output formats** The input table \(D\) of data-sequences has three column attributes: sequence identifier (\(sid\)), transaction time (\(time\)) and item identifier (\(item\)). The output is a collection of frequent sequences. A good representation of the sequences is crucial for easy generation of SQL queries at various stages. After considering a few alternatives, we chose the following representation: A sequence is a tuple in a fixed-width table with schema \((\text{item}_1, \text{eno}_1, \ldots, \text{item}_k, \text{eno}_k)\). The \(\text{eno}\) attributes store the element number of the corresponding items. For sequences of smaller length, the extra column values are set to NULL. For example, if \(k = 4\), the sequence \(((\text{computer}, \text{modem})(\text{printer}))\) is represented by the tuple \((\text{computer}, 1, \text{modem}, 1, \text{printer}, 2, \text{NULL, NULL})\).

**Candidate generation** Candidates are generated in two steps. The join step generates a superset of \(C_k\) by joining \(F_{k-1}\) with itself. A sequence \(s_1\) joins with \(s_2\) if the subsequence obtained by dropping the first item of \(s_1\) is the same as the one obtained by dropping the last item of \(s_2\). This is expressed in SQL as follows:

\[
\text{insert into } C_k \text{ select } I_1.\text{item}_1, I_1.\text{eno}_1, \ldots, I_1.\text{item}_{k-1}, I_1.\text{eno}_{k-1}, I_2.\text{item}_{k-1}, I_2.\text{eno}_{k-1} + I_2.\text{eno}_{k-2} - I_2.\text{eno}_{k-3} \text{ from } F_{k-1} I_1, F_{k-1} I_2
\]

where \(I_1.\text{item}_2 = I_2.\text{item}_1\) and \(I_1.\text{item}_{k-1} = I_2.\text{item}_{k-2}\) and

\[
\begin{align*}
I_1.\text{eno}_{k-2} - I_1.\text{eno}_{k-3} &= I_2.\text{eno}_{k-2} - I_2.\text{eno}_{k-3} \\
I_1.\text{eno}_{k-1} - I_1.\text{eno}_{k-2} &= I_2.\text{eno}_{k-1} - I_2.\text{eno}_{k-2} - I_2.\text{eno}_{k-3}
\end{align*}
\]

In the above query, subsequence matching is expressed as join predicates on the attributes of \(F_{k-1}\). Note the special join predicates on the \(\text{eno}\) fields that ensure that not only do the joined sequences contain the same set of items but that these items are grouped in the same manner into elements. The result of the join is the sequence obtained by extending \(s_1\) with the last item of \(s_2\). The added item becomes a separate element if it was a separate element in \(s_2\), and part of the last element of \(s_1\) otherwise (note the last item in the \text{select} list above).

In the prune step which is similar to the association rule case (section 2.3.1), all candidate sequences that have a non-frequent contiguous \((k-1)\)-subsequence are deleted. While joining \(F_1\) with itself to get \(C_2\), we need to generate sequences where both the items appear as a single element as well as two separate elements.

**Support counting** The KwayJoin approach for sequential patterns is very similar in structure to that for association rules except for these two key differences:

1. We use `select distinct` before the `group by` to ensure that only distinct data-sequences are counted.
2. Second, we have additional predicates \( PRED(k) \) between sequence numbers. The predicates \( PRED(k) \) is a conjunct (and) of terms \( p_{ij}(k) \) corresponding to each pair of items from \( C_k \). \( p_{ij}(k) \) is expressed as:

\[
(C_k.\text{enol}_j = C_k.\text{enol}_i \text{ and } \text{abs}(d_j.\text{time} - d_i.\text{time}) \leq \text{window-size}) \text{ or } \\
(C_k.\text{enol}_j = C_k.\text{enol}_i + 1 \text{ and } d_j.\text{time} - d_i.\text{time} \leq \text{max-gap} \text{ and } d_j.\text{time} - d_i.\text{time} > \text{min-gap}) \\
\text{ or } (C_k.\text{enol}_j > C_k.\text{enol}_i + 1)
\]

Intuitively, these predicates check (a) if two items of a candidate sequence belong to the same element, then the difference of their corresponding transaction times is at most \text{window-size} and (b) if two items belong to adjacent elements, then their transaction times are at most \text{max-gap} and at least \text{min-gap} apart.

The \text{Vertical} approach for association rules is extended as follows. For each item, we create a BLOB (s-list) containing all \((\text{sid}, \text{time})\) pairs corresponding to that item. The sequence table \( D \) is scanned in the \((\text{item}, \text{sid}, \text{time})\) order and passed to the table function \text{Gather}, which collects the \((\text{sid}, \text{time})\) attribute values of all tuples of \( D \) with the same item in memory and outputs a \((\text{item}, \text{s-list})\) pair for all the items that meet the minimum support criterion. The \text{s-lists} are maintained sorted using \( \text{sid} \) as the major key and \( \text{time} \) as the minor key, and is stored in a new \text{SlistTable} with the schema \((\text{item}, \text{s-list})\). The support counting query has the same structure as in \text{boolean association rules} (section 4.2) except that the user defined function to intersect the \text{s-lists} is quite different. Here the candidate containment in data sequences has to account for the timing constraints as well. We use an algorithm similar to the one described in [SA96] for this purpose.

In the \text{GatherJoin} approach, we generate all possible \( k \)-sequences using a \text{Gather} table function, join them with \( C_k \) and group the join result to count the support of the candidate sequences. The time constraints are checked on the table function output using join predicates \( PRED(k) \) as in the \text{KwayJoin} approach.

We augmented the boolean association rule framework to implement \text{generalized association rule} and \text{sequential pattern mining}. The major addition for \text{generalized association rule} was to “extend” the input transaction table (transform \( T \) to \( T^* \)). For \text{sequential patterns}, the join predicates for candidate generation and support counting were significantly different.

6 Proposed extensions

One of the goals of our work has been to “unbundle” complex mining operations like associations and classifications into smaller primitives that can be supported efficiently by a \text{general purpose DBMS} and be useful for a large class of mining algorithms. Our exercise with association rules has helped us identify a few such primitives that we believe could be generally useful in a number of other decision support applications.

\textbf{Richer set operations:} We expect a richer collection of set operations to be useful for mining. For association rules, we used three different versions of the basic \text{subset} operation. For candidate generation we needed to find all \( k - 1 \) subsets of a set of \( k \) elements for doing the pruning. We enumerated the subsets explicitly in the query predicate because the size of the set and the subsets was \text{fixed} and known in advance. For support counting, we used the Comb-K table function which generated all \( k \) subsets of a \text{variable} length set. For rule generation, we used the GenSubset table function to generate \text{all} subsets of a \text{variable} length set. The subset operation would also be useful in building decision trees on categorical attributes [MAR96].

Another important set operation was the \text{intersect} operation used in the \text{Vertical} approach. Most systems already have \text{internal implementations} of this operation for doing \text{AND} and \text{OR} operations on \text{RID} (record identifier) lists obtained during multiple index scans. In current OLAP and data warehousing systems this
operation is rampant in the popular bit-mapped indices. Our Vertical approach is similar to this bit-mapped approach with one important difference. Instead of performing ANDs on RIDs, we perform the ANDs on another attribute, the transaction identifier.

**Gather-scatter operations** Another common operation is the Gather operation that can transform two attributes in a data table to a form where for distinct values of one of the attributes (called the grouping attribute) we collect together in a set all values of the other attribute. We can think of Gather as a glorified aggregate function. Its reverse operation scatter is a special case of the subset operation where the subset size is 1. Recently, similar operators called UnPivot have also been proposed in the context of classification in [GUS98]

We used this operation in four of our SQL approaches: GatherJoin, GatherPrune, Vertical and Horizontal. To implement the Gather table function in our queries we required data to appear in a particular order as inputs to the table function. This would be trivial to express in SQL if order by clauses were allowed in inner subqueries. In the absence of such a feature, we relied on our knowledge of DB2’s internals to force such an order in our experiments. Another way is to treat the Gather function as an aggregate function that simply concatenates its arguments. Systems that have support for user-defined aggregate functions [SK91] can easily support such a functionality.

There are several other decision support applications where gather could be extremely useful. For instance, suppose we are trying to classify customers of a credit card company and the data, in addition to static customer attributes like age, salary consists of detailed transaction data about each purchase activity of a customer. In this case, we would like to gather all activities of a customer as one time series and extract features from these time series for classification. In OLAP applications too, data often needs to be converted back and forth from a format where different measures are gathered together as different attributes of the same row to a format where different measures are scattered as different rows.

**Blank operators** The one functionality that we most missed from the database engine is easy application level support for addition of new operators in a SQL query tree. The existing support for user extensibility in terms of User Defined Functions(UDFs), User Defined Aggregates and Table Functions is rather limited. UDFs can only be invoked on one tuple at a time, they cannot return multiple values and they cannot control the order in which they are invoked. User defined aggregate functions allow limited control on the order in which they are invoked but they are restricted to return exactly one row per group. Table functions provide flexibility in terms of the number of rows that can be returned per input argument but have no control on the order in which input arguments are supplied to it.

An operator that could be treated by the programmer as a general-purpose node of a query’s data flow would be extremely useful for the integration of mining functionality with SQL operators. The table function definition (say as used in DB2/UDB) can be easily extended to meet our requirements. We propose additional clauses that control the order of passing input arguments. First, is the “group-by or order-by” clause on part of the input arguments. Unlike in SQL queries, the group-by clause should not imply that only one record per group is output. Second is the “Parallelizable” clause that indicates if that operator can be decomposed and evaluated simultaneously on different parts of the argument. This clause is already provided in the definition but it needs reinterpretation in view of the first clause. We illustrate how such a blank operator can be used to implement some of the above mentioned primitives. All of the subset operators can be easily implemented using existing table functions. The Gather operator required additional clauses.

**Gather:**
Create function Gather(tid integer, item integer) returns table (items varchar())
This implies that the Gather operator can be parallelized as long as the input arguments are grouped on tid such that all rows with the same tid go to the same node. Another example is the MapTid() operator used in Section 5.1.2 to map long transaction identifiers expressed as strings to unique integers in a pipelined manner. In contrast to Gather this operator cannot be parallelized and it generates multiple output rows per group. Hence the User defined aggregate model would not work.

MapTid:
Create function MapTid(transId varchar()) returns table (tid integer)
<other details about function definition>
Disallow parallelization
Group by tid

In conclusion, we feel that what the core database engine should provide is support for easy addition of powerful blank operators that can be interposed anywhere in the data stream of SQL queries. Using this blank operator a mining extender can be used to implement some of the basic primitives like “subset”, “gather” and scatter mentioned above.

7 Conclusion and future work

We explored various architectural alternatives for integrating mining with a relational database system. As an initial step in that direction we studied the association rules algorithms with the twin goals of finding the trade-offs between architectural options and the extensions needed in a DBMS to efficiently support mining. We experimented with different ways of implementing the association rules mining algorithm in SQL to find if it is at all possible to get competitive performance out of SQL implementations.

We considered two categories of SQL implementations. First, we experimented with four different implementations based purely on SQL-92. Experiments with real-life datasets showed that it is not possible to get good performance out of pure SQL based approaches alone. We next experimented with a collection of approaches that made use of the new object-relational extensions like UDFs, BLOBs, Table functions etc. With this extended SQL we got orders of magnitude improvement over the SQL-92 based-implementations.

We compared the SQL implementation with different architectural alternatives. We concluded that based just on performance the Cache approach is fastest. A close second is the SQL-OR approach that was sometimes slightly better than Cache and was never worse than a factor of two on our datasets. Both these approaches require additional storage for caching, however. The Stored-procedure approach does not require any extra space (except possibly for initially sorting the data in the DBMS) and can be made to be within a factor of two to three of Cache with sampling. The UDF approach is 30% faster than Stored-procedure but is significantly harder to code. The SQL approach offers some secondary advantages like easier development and maintenance and potential for automatic parallelization. However, it might not be as portable as the Cache approach across different database management systems.

This exercise has demonstrated that with a good understanding of the performance profile of a DBMS engine it is possible to express the associations computation efficiently in SQL. Often, the SQL formulation may not be a one to one translation of the file system code, therefore it might be necessary to design alternative methods. This
exercise also helped us identify some easy-to-implement primitives that are useful for other mining algorithms as well. The work presented in this paper points to several directions for future research. What kind of a support is needed for answering short, interactive, adhoc queries involving a mix of mining and relational operations. How much can we leverage from existing relational engines? What data model and language extensions are needed? Some of these questions are orthogonal to whether the bulky mining operations are implemented using SQL or not. Nevertheless, these are important in providing analysts with a well-integrated platform where mining and relational operations can be inter-mixed in flexible ways.

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