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Revisiting the Role of Pipelined Parallelism in Multi-Join Query Processing

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Abstract

Multi-join queries are the core of any integration service that integrates data from multiple distributed data sources. Due to the large number of data sources and possibly high volumes of data, the evaluation of multi-join queries faces increasing scalability concerns. Parallel processing has been applied to tackle this problem. State-of-the-art parallel multi-join query processing commonly assume that the application of maximal pipelined parallelism leads to superior performance. In this paper, we instead illustrate that this assumption does not generally hold. We investigate how best to combine pipelined parallelism with alternate forms of parallelism to achieve an overall effective parallel processing strategy. An m-way bushy parallel processing strategy is proposed. Experimental studies are conducted on an actual software system over a cluster of high-performance PCs. The experimental results confirm that the proposed parallel processing strategy leads to an on average of 50% improvement in terms of total processing time in comparison to existing state-of-the-art solutions.

1 Introduction

Motivation. Many applications such as data integration services, decision support systems, and ETL middleware have their results specified in terms of complex multi-join queries across distributed data sources. Efficient processing of such multi-join queries is thus critical to the success of these applications. The evaluation of multi-join queries can take a prohibitively long time due to the following reasons: (1) the distributed nature of data sources, (2) the possibly large number of data sources, and (3) the large volume of data in each data source. Thus, there is an increasing demand for scalable multi-join query processing solutions.

Parallelizing query processing over a shared-nothing architecture, i.e., a computing cluster, has been shown to have a high degree of scale up and speed up [6]. Here, we use the term machine to refer to each computation device in a shared-nothing architecture. Three types of parallelism have been identified in the parallel query processing [12]. First, query operators none of which use data produced by the others may run simultaneously on distinct machines. This is termed independent parallelism. Second, query operators may be composed by a producer and consumer relationship such that tuples output by a producer can be fed to a consumer as they get produced. This is termed pipelined parallelism. The third, termed partitioned parallelism, refers to running several instances of one single operator on different machines concurrently, with each instance only processing a partitioned portion of the complete data.

Two processing strategies at opposite ends of the spectrum, namely, sequential processing and pipelined processing, have been proposed in the literature [22]. For example, we process a four-way join query $R_1 \bowtie R_2 \bowtie R_3 \bowtie R_4$ on 2 machines. Here, we assume $R_1 \sim R_4$ are not in these 2 machines originally. Figure 1(a) illustrates an example of sequential processing. That is, we first evaluate $R_1 \bowtie R_2$ over 2 machines and get the intermediate result $I_1$. We then process $I_1 \bowtie R_3$ on the same 2 machines (indicated by the dashed rectangle) and get the intermediate result $I_2$. This process repeats until we get the final query results. Figure 1(b) shows an example of pipelined processing of this four-way join query. For example, we first distribute $R_2$, $R_3$, and $R_4$ over the 2 machines. Then, tuples read from $R_1$ probe these relations in a pipelined fashion and generate query results. This pipelined processing of multi-join queries has been shown to be superior to the sequential processing [22]. As we will discuss shortly, state-of-the-art parallel multi-join query processing solutions tend to maximally apply this pipelined processing as its core execution strategy [22, 29, 4].

However, does this commonly accepted solution of maximally applying pipelined parallelism always perform effectively when evaluating multi-join queries? Or put it differently, are there methods that enable us to generate even more efficient parallel execution strategies than this fully pipelined processing?
In this work, we show via a cost analysis as well as real system evaluations that such maximally pipelined processing is not always effective. We propose an m-way bushy parallel processing strategy for multi-join queries that outperforms state-of-the-art solutions.

Focus of the Work. We focus on complex multi-join queries, i.e., they involve 10 or more source relations. We target application scenarios in which all data will be first taken to and then processed in the cluster. This requirement of processing joins outside the data sources is a rather common in many applications. For example, in a data warehouse loading environment (e.g., ETL [20]), operating data sources may be too busy to process such complex join queries or even simply may not be willing to give control to outsiders. Or data sources may not have the advanced query processing capabilities necessary to evaluate complex join queries, i.e., web servers.

We focus on hashing join algorithms [17] since they are among the most popular ones in the literature due to their proven superior performance [21, 17]. Hashing joins provide the possibility of a high degree of pipelined parallelism. Other join algorithms such as sort-merge join do not have this natural property of pipelined parallelism [21]. Furthermore, hashing joins also naturally fit partitioned parallelism.

The key research question that we propose to address in this work is whether maximally pipelined multi-join query processing is indeed a superior solution as commonly assumed in the literature. This pipelined process implies main memory based processing. Hence, we assume that the aggregated memory of all available machines is sufficient to hold the hash tables of the join relations \(^1\). The rationale behind this is that both the main memory of each machine and the number of machines in the cluster are getting increasingly large at affordable cost.

Due to possibly large volumes of data in each source relation, the main memory of one machine may not be enough to hold the full hash table of one source relation. Thus, partitioned parallelism is applied to each join operation whenever it is necessary. That is, a partition (exchange) operator [11] will be inserted into the query plan to partition the input data tuples to multiple machines to conduct a partitioned hash join processing.

Contributions. To highlight, the main contributions of this work include:

- We question the commonly accepted model of maximally pipelined parallelism in parallel multi-join query processing by both an analytical argument as well as experimental observations.
- We propose an m-way bushy parallel processing strategy that aims to balance all three forms of parallelism for complex multi-join queries. This has not been carefully explored in the literature.
- We provide optimization algorithms to generate the above m-way bushy processing strategies.
- We build a distributed query engine to back up our claims. We incorporate our proposed strategies and algorithms into the system. Extensive experimental studies show that the m-way bushy parallel processing has on average a 50% improvement in terms of total processing time in comparison to state-of-the-art solutions.

The remainder of the paper is organized as follows. Section 2 describes the state-of-the-art. Section 3 discusses a multi-phase parallel optimization approach. Section 4 analyzes the cost factors and tradeoffs that affect the parallel processing performance. Section 5 presents the proposed m-way bushy tree processing and optimization algorithms. Experimental results are provided in Section 6. While Sections 7 and 8 discuss related work and conclusions respectively.

2 State-of-the-Art

Various solutions have been investigated for parallel multi-join query processing in the literature [22, 29, 4]. To illustrate, we use the 10-join query depicted in Figure 2 to explain the core ideas. The multi-join query is depicted by its join graph. Each node in the graph \((R_0 \sim R_9)\) represents one join relation (data source), while an edge denotes a join between two respective data sources.

2.1 Sequential vs. Pipelined Processing

Two strategies at opposite ends of the spectrum, namely, sequential processing and pipelined processing, have been proposed [22]. Note that partitioned parallelism is applied by default for each join operator. Sequential processing is

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\(^1\)In situations when main memory is not enough to hold all hash tables at the same time, we follow the typical approach to divide the query into several pieces with each piece being processed sequentially. We defer this discussion to Section 5.4.
based on a left-deep query tree. Figure 3(a) illustrates one example of sequential processing for the query defined in Figure 2. Here $B_i$ represents the building phase of the $i$-th join operation, while $P_i$ denotes the corresponding probing phase. This processing can be described by the following steps: (1) scan $R_0$ and build $B_1$, (2) scan $R_1$, probe $P_1$, and build $B_2$, (3) scan $R_2$, probe $P_2$, and build $B_3$, and so on. This is repeated until all the join operations have been evaluated. As can be seen, it processes joins sequentially and only partial operations, namely, the probing and the successive building operations, are pipelined.

Pipelined processing is based on a right-deep query tree [22]. Figure 3(b) illustrates an example of pipelined processing for the same query in Figure 2. In this case, all the building operations such as scan $R_1$ and build $B_1$, scan $R_2$ and build $B_2$, ..., scan $R_9$ and build $B_9$ can be run concurrently. After that, the operation of scan $R_0$ and all the probing operations, probe $P_1$, probe $P_2$, ..., probe $P_9$ can be done in a pipelined fashion. As demonstrated above, it achieves fully pipelined parallelism.

Note that a pipeline process implies main memory based processing. That is, it requires there to be enough main memory to hold all the hash tables of the building relations ($R_1 \sim R_9$ in this case) throughout the duration of processing the query.

As identified in [22], pipelined processing is preferred whenever main memory is adequate. This is because (1) intermediate results in pipelined processing exist only as a stream of tuples flowing through the query tree, and (2) even though sequential processing in general may require less memory, this is not always true due to intermediate results have to be stored. A large intermediate result may consume even larger memory than the sum of all building relations.

The simulation results in [22] confirm that the pipelined processing (right-deep) is more efficient than the sequential one (left-deep) in most of the cases they considered. Without loss of generality, we thus associate the pipelined processing with a right-deep query tree, and the sequential processing with a left-deep query tree in the following discussions.

2.2 Maximally Pipelined Processing

State-of-the-art parallel multi-join query processing solutions maximally pursue the above pipelined parallelism to improve the overall performance [22, 29, 4]. If the main memory is not enough to hold all the hash tables of the building relations, they commonly take the approach of dividing the whole query into “pieces”, with the expectation that the building relations of each piece fit into the main memory. That is, pieces are processed one by one with each piece utilizing the entire memory applying fully pipelined parallelism.

For example, zigzag processing [29] takes a right-deep query tree and slices it into pieces based on the memory availability. As an example, the right-deep tree in Figure 3(b) is cut into two pieces, one is $R_0 \sim R_3$, and the other is $I_1, R_4 \sim R_9$ (Figure 4(a)). Here, $I_1$ corresponds to the result of the first piece $R_0 \sim R_3$. These two pieces are processed sequentially with fully pipelined parallelism in each piece.

Segmented right-deep processing [4] proposes heuristics, namely, balanced-consideration and minimized-work, to generate pieces directly from the query graph based on the memory constraint. The query tree is similar to the zigzag tree. However, each piece can be attached not only at the first join operation of the next piece, but instead also in
the middle of it. For example, Figure 4(b) illustrates one example of segmented right-deep processing. As can be seen, the output (from \( P_3 \)) is attached as the building relation of \( B_8 \).

To summarize, all the above approaches take the common model of pursuing a maximally pipelined processing of multi-joins via a right-deep query tree, with the number of join relations in the right-deep tree primarily being determined by the main memory available in the cluster.

We now question the performance of such a maximally pipelined processing model. As mentioned earlier, this pipeline process implies a main memory based processing. Clearly, more efficient main memory based processing strategies would lead to an improved overall performance. Without loss of generality, we use the term pipelined segment to refer a right-deep query tree that can be fully processed in the main memory.

3 A Multi-Phase Optimization

Multi-join query optimization is an expensive process because the number of alternative query plans for a query grows at least exponentially in the number of relations participating in the query [26]. Parallel multi-join query optimization is even harder [14, 23, 9]. Complications arise because the cost to be optimized, either total amount of work to be processed or total processing time, are no longer closely correlated since a query plan with minimal work may have a high sequential dependency that results in high overall processing time. Second, even one sequential query plan can in turn have a huge number of parallel solutions.

We take a multi-phase optimization approach in this work to cope with the complexity of parallel multi-join query optimization. That is, we break the whole optimization task into several phases and then optimize each phase individually. While a single-phase optimization approach such as [23] could also be applied, our multi-phase approach enables us to focus our attention on the research task we are tackling.

Breaking the Optimization Task. We divide the whole optimization task into the following three phases, (1) generating an optimized query tree, (2) allocating query operators in the query tree to machines, and (3) choosing pipelined execution methods. We note that even if we divide the optimization task into multiple phases, the complexity of each phase, i.e., phases (1) and (2), still remains exponential in the number of join relations.

The main focus of this work is on investigating the impact of query trees (phase (1)) and different forms of parallelism on the overall performance. To proceed, we first describe the design choices we will assume in the reminder of our work for phases (2) and (3) below. We simplify the operator-machine allocation (for phase(2)) and choose the concurrent execution approach [22] as the pipeline execution method (for phase(3)).

Allocating Query Operators. Query operators (joins) need to be allocated to machines in the cluster. However, resource allocation itself is a research problem of high complexity that has been extensively investigated in the literature [16, 10, 15]. Like most work in parallel multi-join processing literature [22, 29, 4], we focus on main memory in the allocation phase. This is because main memory is the key resource in the above hash-based join processing. Other factors such as CPU capabilities of computation nodes are assumed to have less impact on the allocation, i.e., they are often assumed to be sufficient.

The allocation is performed based on pipelined segments to promote the usage of pipelined parallelism [16]. For example, if a right-deep tree is cut into pieces with each piece being processed sequentially due to insufficient memory, then all machines are allocated to each piece. Thus, the whole allocation is performed in a linear fashion. As it can be seen, all previous processing strategies described in Section 2 fall into this type of linear allocation.

Pipelined Execution Method. The building relations of each pipelined segment can entirely fit into the memory of the machines that have been allocated to it. We apply a concurrent execution approach [22] to process a pipelined segment. In this execution method, all scan operations are scheduled concurrently. For example, in Figure 5, we process a 4 way pipelined segment on 3 machines. Each building relation \( (R_2 \sim R_4) \) is evenly partitioned across all 3 machines. Thus, each machine houses the appropriate partitions from all building relations, denoted as \( P_j \). Here, subscript \( i (2 \leq i \leq 4) \) denotes join relations, while superscript \( j (1 \leq j \leq 3) \) represents machine ID. The probing relation \( (R_1) \) is also partitioned into all 3 machines to probe the appropriate hash tables to generate results.

Other pipelined execution strategies such as staged partitioning [4] have also been proposed. The detailed discussion of these strategies and their impact on parallel processing strategies are beyond the scope of this paper. They can be found in our technical report.
4 Cost Analysis of Pipelined Segment

4.1 Identifying Tradeoffs

The following two factors need to be considered when analyzing the performance of parallel multi-join query processing via a partitioned hashing: (1) redirection costs between join operations, and (2) optimal degree of parallelism.

Redirection Costs. The basic idea behind the partitioned hash join algorithm is that the join operation can be evaluated by a simple union of joins on individual partitions. For example, an equi-join \( A \bowtie B \) can be computed via \( (A_1 \bowtie B_1) \cup (A_2 \bowtie B_2) \cup \ldots \cup (A_n \bowtie B_n) \) if \( A \) and \( B \) are first divided into \( n \) partitions \( (A_1 \sim A_n, B_1 \sim B_n) \) using the same hash function. Assume the two partitions in a pair \( (A_i, B_i) \) are put in the same machine, while different pairs are spread over the distinct machines. This way, all pairs can be evaluated in parallel.

However, for a right-deep tree segment, it is not possible to always have all the matching partitions reside in the same machine. For example, assume a query tree is defined by “A.\( A_1 = B. B_1 \) and B.\( B_2 = C. C_1 \)””. \( A \) and \( B \) are partitioned based on their common attribute \( A.A_1 \) (or \( B.B_1 \)), while \( C \) has to be partitioned based on the common attribute between \( B \) and \( C \), namely, \( B.B_2 \) (or \( C.C_1 \)). If we assume \( A \) is the probing relation, then the partition function of \( B.B_2 \) has to be re-applied to the intermediate result of \( A_i \bowtie B_i \) to find the corresponding partitions \( C_i \). However, this corresponding partition \( C_i \) might exist in a machine different from where the current \( B_i \) resides. Thus redirection of intermediate results is necessary in this situation. For the special case of a right-deep tree when only one attribute per source relation is involved in the join condition, i.e., “A.\( A_1 = B. B_1 = C. C_1 \)””, the same partition function can be applied to all relations. In that case, all the corresponding partitions can be put into the same machine to avoid such redirections.

Optimal Degree of Parallelism. Startup and coordination overhead among machines will counteract the benefits that could be gained from parallel processing. [27, 19] discuss the basics on how to choose the optimal degree of parallelism for a single partitioned operator, meaning the idea number of machines that need to be assigned to one operator. As one example, if a relation only has 1,000 tuples, it is not a good idea to have it evenly distributed across a large number of machines (i.e., 100) since the startup and coordination costs among these machines might be higher than the actual processing cost. Given the processing of more than one join operators (pipelined segment), we expect this factor has a major impact on the overall performance.

4.2 Pipelined Processing Cost Model

For pipelined processing of a right-deep segment, the cost in terms of total work versus the overall processing time may not be that closely correlated. We thus derive two separate cost models. To facilitate the description of cost models, we assume \( R_0 \) is the probing relation, while \( R_1, \ldots, R_n \) are the building relations of the pipelined segment. We also assume \( k \) machines are available to process the pipelined segment. These machines are denoted by \( M_1, M_2, \ldots, M_k \). Without loss of generality, we use \( I_1 \) to represent the intermediate result after joining with \( R_0 \). For example, \( I_1 \) denotes the result of \( R_0 \bowtie R_1 \), while \( I_2 \) represents \( I_1 \bowtie R_2 \). Thus \( I_n \) represents the final output of these joins.

Estimating Total Work. The total work of pipelined processing can be described as the sum of the work in the building phase \( (W_b) \) and the work in the probing phase \( (W_p) \), as listed below.

\[
W_b = (t_{\text{read}} + t_{\text{partition}} + t_{\text{network}} + t_{\text{build}}) \sum_{i=1}^{n} |R_i|
\]

\[
W_p = (t_{\text{read}} + t_{\text{partition}} + t_{\text{network}} + t_{\text{probe}}) |R_0| + \frac{k - 1}{k} \sum_{i=1}^{n-1} |I_i| \cdot t_{\text{network}} + \left( \sum_{i=1}^{n-1} |I_i| \right) \cdot t_{\text{probe}}
\]

\[
t_{\text{read}}, t_{\text{partition}}, t_{\text{network}}, t_{\text{build}}, \text{ and } t_{\text{probe}} \text{ in the above formulae represent the unit cost of reading a tuple from a source relation, partitioning, transferring the tuple across the network, inserting the tuple into the hash table, and probing the hash tables respectively. They represent the main steps involved in a partitioned hash join processing. In the probing phase, } \frac{k - 1}{k} \sum_{i=1}^{n-1} |I_i| \cdot t_{\text{network}} \text{ denotes the redirection cost assuming the redirection occurs after each join operation and the output of each join operation is uniformly distributed across all the machines. The cost of outputting the final results is omitted since it is the same for all processing strategies.}

Estimating Processing Time. Similarly, estimation of the processing time can be divided into two parts: one, the hash table building time \( (T_b) \) and two, the probing time \( (T_p) \). The building time of the pipelined processing \( T_b \) can be estimated as follows:

\[
T_b = \max_{1 \leq i \leq n} (t_{\text{read}} + t_{\text{partition}} + t_{\text{network}} + t_{\text{build}}) \cdot \frac{f(k)}{k} |R_i|
\]

The processing time of the building phase can be estimated as the maximal building time of each individual relation over \( k \) machines. Here, \( f(k) \) represents the contention factor of the network since the more machines are involved,
the more contention of the network caused by transferring tuples of join relations arises. This is used to reflect the optimal degree of parallelism as discussed in Section 4.1.

The processing time of the probing phase is more difficult to analyze because of the pipelined processing. We use the following formula to estimate the pipeline processing time.

\[ T_p = I_{\text{setup}} + \frac{W_p}{k} + I_{\text{delete}} \]

Here \( I_{\text{setup}} \) represents the pipeline setup time, while \( I_{\text{delete}} \) denotes the pipeline depletion time. The steady processing time of the pipeline can be estimated by the average processing time of one tuple \( \frac{W_p}{|R_0|} \) multiplied by the number of tuples \( |R_0| \) that need to be processed over the total of \( k \) machines. Clearly, this is a simplified model representing the ideal steady processing time without including for example variations in the network costs.

As we will discuss in Section 5.3, the above cost model is used in finding the most efficient pipelined processing strategies of each subgraph.

5 Breaking Pipelined Parallelism

5.1 Bushy Trees and Independent Parallelism

Query trees of a multi-join query can be classified into two types: sequential trees (i.e., a right-deep tree or a left-deep tree as discussed above), and bushy trees. A right-deep tree has a better performance over a left-deep tree since it has a high potential of pipelined parallelism for a hash-based join algorithm. Thus we now use a right-deep tree as the representative of sequential trees (e.g., Figure 6(a)).

A bushy tree has a height of at least \( \log_2 n \) (given a binary bushy tree that is balanced) with \( n \) being the number of join relations involved in the multi-join query. A bushy tree brings new flexibility to the style of processing, such as having multiple probing relations and composing different pipelined segments. Moreover, a bushy tree has the potential of processing independent subtrees (segments) concurrently. However, such flexibility may also bring dependencies to the execution. This dependency may both affect the allocation of query operators and the corresponding parallel processing performance.

For example, Figure 6(b) illustrates one bushy tree and its possible pipeline segments (each pipeline segment is denoted by one dashed oval). Four segments (\( P_1 \sim P_4 \)) can be identified. As can be seen, \( P_1 \) and \( P_3 \) can be processed in parallel by processing them on different machines. While the execution of \( P_2 \) depends on \( P_1 \), the execution of \( P_4 \) depends both on \( P_2 \) and \( P_3 \).

As can be seen, a right-deep tree has the highest degree of pipelined parallelism without any dependencies because each subtree is a join relation. However, there is no opportunity for independent parallelism except during the initial building phase of the join relations. While a wide bushy tree has many subtrees, it also has up to \( \log_2 n \) layers of dependencies with \( n \) being the number of source relations. These dependencies are likely to impact the overall performance.

5.2 M-way Bushy Tree

Seen from the cost model, if the results of pipelined segments in a bushy tree are smaller than those of the original join relations, then the bushy tree processing may have less total work \( (W_b + W_p) \) when compared with the fully right-deep processing. Here we assume all the intermediate results are kept in main memory.

Comparing the overall parallel processing time of fully right-deep and bushy trees is more complicated. As we can see, each pipelined segment in a bushy tree only gets one portion of the total available machines. Thus the network contention \( f(k) \) in the building phase may be less severe than that of the full right-deep case. As a consequence, given the independent processing of these smaller pipelined segments, the processing time of a bushy tree may be better than that of fully pipelined processing. However, as we identified earlier, a bushy tree style processing may be affected by the dependencies among subtrees. Moreover, there may be subtrees (up to \( \lceil n/4 \rceil \)) that have short pipelined processing. For example, \( P_1 \) and \( P_3 \) only have a pipeline of one probing followed by the building for the next join. These two factors may eventually counteract the benefits gained by introducing the independent parallelism and smaller network contention in each segment.

Thus, the key question now is how to balance independent parallelism and pipelined parallelism in parallel multi-join query processing. By reducing each pipelined segment (i.e., identified by dashed oval in Figure 6(b)) into one ‘mega-node’, we can build a dependency tree out of the original query tree. We note that the dependencies are associated with the height of this dependency tree. Thus reducing the height of the dependency tree should effectively
reduce the dependencies. We thus propose to utilize an \textit{m-way bushy} query tree. An m-way bushy tree can be controlled to have a dependency tree with height of 2 as long as we increase the number of subtrees of the root node.

Figure 7 illustrates the example of an m-way bushy tree of the join query in Figure 6. In this example, the whole query is cut into three groups, \(R_1 \sim R_4\), \(R_4 \sim R_7\), and \(R_8\). Three pipelined segments \(P_1\), \(P_2\), and \(P_3\) can be identified correspondingly. \(P_1\) and \(P_2\) can be processed independently, each with pipelined parallelism. The output from these two segments can be directly fed into \(P_3\). Without loss of generality, the pipelined segment that contains outputs of all other segments is referred to as the \textit{final pipelined segment}. In this case, \(P_3\) is the final pipelined segment. Thus, all pipelined segments except the final one can be executed concurrently without any dependencies. We can see that an m-way bushy tree processing applies independent parallelism with minimal dependencies among subtrees (groups) since it only has one layer of dependencies among pipelines.

Without loss of generality, we always assume the right-most pipeline of an m-way bushy tree to serve as the probing relation of the final pipelined segment. For example, \(P_1\) is the probing relation of the final segment \(P_3\) in Figure 7.

![Figure 7. A M-way Bushy Tree](image)

### 5.3 Composing m-way Bushy Tree

Now, we address the question how to generate the above m-way bushy tree for a multi-join query. Algorithm 1 sketches our proposed algorithm. It consumes a connected join graph \(G\) and the maximal number of nodes \(m\) per group (we will discuss how to get this \(m\) shortly). We would choose the largest join relation as the probing relation of each group since this reduces the time and the memory consumption of the building phase. Once we select the probing relation, we then begin to enumerate all possible groups having a maximum of \(m\) join nodes starting from this probing relation. Enumeration is possible since \(m\) is usually much smaller than the number of nodes in the join graph. Some of the groups may not contain exactly \(m\) nodes due to the nodes in the group being no longer connected by a join edge. Our goal is to avoid Cartesian products given that each data source may be large, thus resulting in huge intermediate results. After that, we choose the best graph, a partition of the original join graph, from these candidates generated from the enumeration based on the cost model we developed in Section 4.2. The selection can also be based on heuristics, i.e., choosing the group in which the join attributes are the same to reduce the possible redirection costs, or selecting the one with the smallest output results.

#### Algorithm 1: ComposeBushyTree(G,m)

**Input:** A connected join graph \(G\) with \(n\) nodes. Number \(m\) that specifies the maximum number of nodes in each graph.

**Output:** An m-way bushy tree that has at least \((\lceil n/m \rceil - 1)\) subtrees.

1: completed = false
2: while (!completed) do
3:   Choose a node with largest cardinality that has not yet been grouped as probing relation
4:   Enumerate all subgraphs starting from node selected in Step 3 with at most \(m\) nodes
5:   Choose best subgraph, mark the nodes in this group have been selected in original join graph
6:   if (\(\exists K, K\) is a connected subgraph of \(G\) with unselected nodes) \&\& (\(K\).size() \(\geq 2\)) then
7:     completed = true
8:   end if
9: end while
10: Compose an m-way bushy tree

![Figure 8. An Example of the Algorithm](image)

Figure 8 illustrates how the example join graph depicted in Figure 2 is divided by applying Algorithm 1 when \(m = 4\). For example, we start from the relation with largest cardinality, say relation \(R_7\). The enumeration in Step 4 generates all the possible connected groups with 4 nodes starting from \(R_7\), as illustrated in Figure 8(a). In this case, we choose \(R_7\), \(R_9\), \(R_4\), and \(R_8\) as the nodes in the first group (pipelined segment). For simplicity, we call this group \(G_1\). After this, if \(R_1\) is the one with the largest cardinality among the nodes that have not yet been grouped, we then choose \(R_1\) as the
probing relation for the second group $G_2$. We repeat the process as illustrated by Figures 8(b)-(c). After these steps, only $R_0$ and $R_5$ are left. They are not connected. We thus end up with 4 groups. An example m-way bushy tree with these 4 groups can be built as shown in Figure 9(a).

Allocating machines to an m-way bushy is based on the number of building relations in each pipelined segment. For example, for the m-way bushy tree shown in Figure 9(a), three pipelined segments can be identified (see dashed cycles in Figure 9(b)). The number of machines that are assigned to each pipelined segment, denoted by $k_1$, $k_2$, and $k_3$, can be computed as follows.

$N_b = \sum_{0 \leq i \leq 9, i \neq 1,7} |R_i| + |I_1|$

$k_1 = \left(\left\lfloor \frac{|R_0| + |R_8| + |R_9|}{N_b} \right\rfloor\right)$

$k_2 = \left(\left\lfloor \frac{|R_2| + |R_3| + |R_4|}{N_b} \right\rfloor\right)$

$k_3 = k_1 - k_2$

Here, $I_1$ and $I_2$ denote the outputs of groups $G_1$ and $G_2$ respectively. $N_b$ represents the total number of tuples that need to be built assuming $R_7$, $R_1$, and $I_2$ are the probing relations of $G_1$, $G_2$, and the final pipelined segment respectively. Note that the selection of the probing relation for the final pipeline segment is not straightforward. We will discuss this in more detail in Section 6.5.

![Figure 9. M-way Tree and Node Allocation](image)

However, the question remains how to decide the right number of groups given a join graph. Let us now use $g$ to represent this number. Note that the input of Algorithm 1, the maximum number of nodes in each group $m$ can be estimated by $m = \lceil n/g \rceil$ with $n$ being the number of join relations in the query. There are two ways to address this issue. The first is a heuristics-based selection approach. For example, we can choose $g$ as the number of nodes that have cardinality larger than $3/2$ of the average cardinality. Here, we assume that $g$ has to be bound within $2 \sim n/2$. The rationale behind this selection criterion is that in the best case, we can choose all these large join relations as the probing relations for the generated groups. The second is a cost-based selection approach. Again we note that the range of the number of groups $g$ is between 2 to $n/2^4$. We thus can repeatedly call the function $ComposeBushyTree$ (Algorithm 1) with the number $m$ ranging from $n/2$ to 2 ($g$ changes from 2 to $n/2$ correspondingly). We then estimate the cost of processing strategy from $ComposeBushyTree$. The final output will be the one with the best estimated cost. While this may increase the optimization cost, it has the potential to result in a better processing strategy.

5.4 Handling Insufficient Memory

The problem of handling insufficient memory can be addressed using the “cutting” principle as in [22, 4]. That is, we divide the whole query (joins) into pieces such that each piece can be run in the main memory. Note that in the extreme case, the multi-join query processing would have to be sequentialized due to not enough memory being available to hold more than one join. As we mentioned in Section 1, we assume that the aggregated memory can hold at least 2 or more building relations.

Algorithm 2 sketches an incremental approach to address this problem. This incremental approach is based on the static right deep tree [22] or segmented right-deep tree [4] which divides the join query into right-deep segments based on the main memory of the cluster. After that, we further compose each right-deep segment into an m-way bushy tree if it is necessary, i.e., the number of building relations in each piece is larger than a certain threshold. Since each right-deep segment is likely to be more efficiently processed, the performance of the whole query is also expected to be better than the static right-deep or segment-right-deep tree processing.

**Algorithm 2 SimpleInclMwayTree(G,M)**

**Input:** A connected join graph $G$ with $n$ nodes, total main memory of cluster $M$. **Output:** A sequence of m-way bushy trees, each processable in main memory of cluster.

1: Compose Static or Segmented Right-Deep Tree
2: for each right-deep segment $r$ do
3:    $m \leftarrow$ Maximal number of relations per group
4:    $t \leftarrow$ ComposeBushyTree($r$, $m$)
5:    Put $t$ into result sequence
6: end for
7: Return result sequence

A “top-down cut” approach, dividing the join graph directly such that each group can be processed in the main

\(^4\)In extreme cases, the actual number of groups may be larger than $n/2$. However, we assume that we have less interests in these cases when a large number of groups with only one join relation in it.
memory, can also be devised. We then select the groups and process them iteratively. However, as mentioned earlier, the essence of our work is to re-examine the performance of a main memory based maximal pipelined processing. We argue that having a more efficient main memory based processing strategies will also lead to improved overall performance even if we apply a simple incremental optimization algorithm such as Algorithm 2. This claim is confirmed by our experimental studies discussed below.

6 Experiments

6.1 Prototype System

We have implemented a distributed query engine to test out our hypothesis. The system is implemented using Java. It is capable of optimizing and executing multi-join queries across a set of shared nothing machines connected by network. The basic architecture of the system is depicted in Figure 10. The architecture consists of two main modules, one is the controller module and the other is the execution module. The controller module is in charge of managing the computation process. It can be installed on a standalone machine or on the machine that has other modules. The controller module contains packages that compose multi-join queries, generate parallel execution query plans, and distribute query plans to the participating machines. The parallel query plans (processing strategies) are specified by query operators such as scan, partition, hash join, union and load in an xml file format. The query is executed in the execution module. This execution engine is installed on each participant machine in the cluster that is involved in the computation process. The execution engine in each node waits for incoming query plans sent by the controller module. Once the execution engine receives the query plan, it parses the query plan, initializes it and starts up the query operators. After that, query operators in different computation machines automatically connect to each other and begin the query processing.

The system is deployed on a cluster composed of 10 machines, as described Figure 11. Each machine in the cluster has dual 2.4GHz Xeon CPUs with 2GB RAM. They are connected by a private gigabit ethernet switch. In our experimental setting, all source (join) relations are stored in an oracle database that reside in a different machine outside the cluster having 2 PIII 1G Hz CPUs and 1G main memory. The query results are sent to an application server with one PIII 800M Hz CPU and 256M Memory. This setup follows a typical data warehouse loading environment (e.g., ETL [20]) where the process has to be performed outside the data sources. This is because the operating data sources may be too busy to process complex join queries or even simply may not be willing to give control to the outsiders.

6.2 Experimental Setup

As done in [4], we use generated data sets and queries in our experiments. This is because benchmark queries such as TPC-H [25] only have a limited number of queries (around 20), and most of them have less than 5 joins. The multi-join queries used in the experiments are randomly generated with the number of join relations ranging from 8, 12, to 16. The cardinality of each join relation ranges from 1K ∼ 100K tuples, and the average size of each source tuple is about 40 bytes. Each result tuple has about 320 ∼ 640 bytes on average, by simply concatenating all tuples from join relations. Data size in our experiment is chosen to make sure all the hash tables can fit in the main memory since our main focus of this work is the main memory based processing.

\footnote{We actually generate random connect acyclic graphs given a specified number of nodes. Each node represents join relations, while each edge denotes the join condition.}
6.3 Impact of the Number of Data Servers

Initial experiments have been conducted to evaluate the impact of the number of Oracle data servers in the experimental setup on the overall performance. We compare the performance of multi-join queries using a pure right-deep tree (pipelined) processing given different numbers of data servers. The test queries are generated randomly with \(8 \sim 16\) join relations. For each query, we vary the number of data servers from 1 to 4. Thus, if we have \(i\) data servers with \(1 \leq i \leq 4\) and \(k\) (either 8, 12, or 16) join relations, then we have each data server hold on average \(\lceil k/i \rceil\) join relations. These data servers are deployed on different machines with similar configurations having Oracle 8i installed. The result is shown in Figure 12. Each data point in Figure 12 reflects an average of 50 randomly generated queries for each query type (queries have the same number of join relations). In Figure 12, x-axis denotes the number of join relations in the query, while y-axis represents the total processing time. From Figure 12, we can see that the number of data servers in the system only has a minor impact on the overall performance. This is because the total time spend on reading the tuples from data servers only represents a small fraction of the total query processing time in our current experimental settings. Thus, the improvement due to shared read by multiple data servers does not play a major role in the overall performance. This indicates that the data server is not the bottleneck in our experimental environment. Without loss of generality, we report our following experimental results with a setup that stores all join relations in one data server.

![Figure 12. Vary the Number of Data Servers](image)

6.4 Pipelined vs. M-way Bushy Processing

Experiments have been conducted to compare the performance (total processing time) of a pure right-deep tree processing having fully pipelined processing to our proposed m-way bushy tree processing that mixes both pipelined and independent parallelism. Figure 13 shows the results of 20 randomly generated queries with 8 join relations. Here, the m-way bushy tree has a maximum of 3 join relations per group. In Figure 13, we see that an m-way bushy tree processing almost consistently outperforms fully pipelined processing.

![Figure 13. Performance of 20 Example Queries](image)

Figure 14 shows the results of queries with an increasing number of join relations in the query. The number of relations in a query ranges from 8, 12 to 16. The experimental results reflect an average processing time over 50 different randomly generated queries per query type. For example, for queries with 8 join relations, we generate 50 queries randomly. We then produce both the fully pipelined processing and the m-way bushy processing strategies for each generated query. In this experimental setup, queries with 8 relations are divided into groups having a maximum of 3 relations, while queries with 12 and 16 relations are divided into groups having a maximum of 4 relations.

In Figure 14, we can see that m-way bushy tree processing is consistently better than maximal pipelined parallelism. The performance improvement is around 50% in terms of the total processing time.

![Figure 14. Right-Deep vs. M-way Bushy](image)

6.5 Probing Relation Selection for Final Pipelined Segment

Selection of the probing relation of a pipelined segment is usually based on the cardinality of the join relations.
This is because choosing a large relation as probing relation can effectively reduce the work and processing time of the building phase. However, for a pipelined segment that involves outputs from other segments (assuming main memory is enough to hold these building relations), the cardinality of the relation alone may no longer be the best choice in general. Changing the probing relation of a pipelined segment that only involves source join relations does not change the number of probes in the probing phase. It only changes the number of probing and building tuples. Here we define the number of probe steps as the maximum number of hash tables that a tuple from the probing relation needs to probe to produce the final output. However, for a pipeline segment having outputs from other segments, changing the probing relation will also change the total number of probes.

For example, if we change the probing relation for the pipeline segment \( P_1 \) as shown in Figure 15(a) from \( R_7 \) to \( R_6 \), no changes in the number of probe steps occur. Both of them are 3 (Figures 15(a)-(b)). However, if we change the probing relation of pipeline \( P_3 \) (exchanging \( P_1 \) and \( P_2 \)), then the total number of probe steps changes from 4 to 5 in this case. This is because \( P_1 \) itself has 3 probe steps while \( P_2 \) only has 2.

**Figure 15. Probing Relation Selection**

Figure 16 shows the experimental results of the impact of the probing relation selection for the final pipelined segment. Here, the number on the x-axis denotes the number of relations in the probing relation of the final pipelined segment. The generated queries have 16 join relations. In Figure 16, we see that in our current environment, the larger the number of relations in the probing relation of the final pipelined segment, the worse the total processing performance will be. This is because the longer probe steps in the final pipelined segments impair the processing performance. This again confirms our observation that a full pipeline may not be the best performer. Note that the performance degradation for a pipeline that is longer than 8 can be explained by the experiments shown in Figure 14. Hence, in Figure 16, we conveyed the scope of smaller pipeline sizes.

**6.6 Number of Join Relations per Group**

Figure 17 illustrates the impact of the maximal number of join relations per group in our environment. Here, all the tested queries have 16 join relations. We vary the number of join relations per group from 3 to 6. As we can see, if the number of join relations per group increases, the total processing time also increases. This is mainly because given our ComposeBushyTree algorithm, the final pipelined segment tends to choose the largest subgraph (the one with the largest number of join relations) as the probing relation since it usually has the largest intermediate results. As shown in Section 6.5, a long pipeline of the final pipelined segment degrades the overall performance. We thus revise our algorithm to choose the subgraph with the smallest number of probing steps as the probing relation of the final pipelined segment. As can be seen, the revised algorithm is less sensitive to the number of join relations in a group.

**Figure 17. Exchanging the Probing Relation**

**6.7 Handling Insufficient Memory**

Figure 18 shows the experimental results when the aggregated main memory is not sufficient to hold all the hash tables of the building relations. We deploy join queries with 32 join relations. Assume the query will be cut into three
pieces with each piece being executed sequentially. Here, the intermediate results of each piece will be first written to the data server, while the next piece will read the intermediate results back into the main memory. We compare the performance of the segmented right-deep tree with our m-way bushy tree generated by Algorithm 2. Note that the segmented right-deep tree has each piece fully pipelined, while the m-way bushy will have the same right-deep segment (piece) further composed into an m-way bushy tree with a maximum of 3 join relations per group. Figure 18 reports the comparison between these two approaches for 10 randomly generated queries. As can be seen, the m-way bushy tree processing consistently outperforms the segmented right-deep processing. This is expected because each piece is processed more efficiently given our m-way bushy tree approach. Thus, the overall performance of the query is correspondingly improved.

![Figure 18. M-way Bushy vs. Segmented Right-Deep](image)

6.8 Concluding Remarks

As can be seen, these experimental results clearly highlight the main message of our work, namely, the long-standing assumption that “maximal pipelining is preferred” is shown to be wrong. Our proposed m-way bushy processing almost consistently beats full pipelined processing. Given the massive application of pipelined processing, especially in growing areas such as continuous query processing, this observation can also shed some new light on how best to optimize distributed pipelined query plans when the optimization function is related to total processing time.

7 Related Work

Parallel query processing has been extensively studied in the literature [6, 27, 13, 19, 22, 14, 5, 18, 11]. Many different research efforts have been conducted in this area. For example, GAMMA [7], Bubba [2], PRISMA/DB [27] are examples of parallel database systems. Many papers were written studying their performance. [13] proposes solutions for scheduling pipelined query operators to minimize the total work. Task scheduling and allocation in general also have been extensively studied [15]. Other focuses such as load balancing [8, 3] and resource allocation [16, 10] are also topics closely related to parallel query processing. As can be seen, these works provide the necessary background for the work presented in this paper. In this work, we instead focus on a specific area of parallel query processing, namely, the parallel multi-join query processing via hashing.

Evaluating a multi-join query via hashing in parallel (applying partitioned and pipelined parallelism) over a shared-nothing environment also has been investigated in the literature before [22, 24, 18]. Different parallel processing strategies such as left-deep and right-deep [22], segmented right-deep [4], and zigzag tree [29] have been proposed, as we have provided an in-depth discussion in Section 2. However, these proposed solutions all share the common approach which is to maximally use pipelined parallelism (i.e., maximally divide a right-deep tree into segments) based on certain objective functions (i.e., memory constraints), and each segment is processed one by one. In this work, we instead consider more tradeoffs in optimizing such parallel multi-join query processing, i.e., other types of query tree shapes, independent parallelism and its dependencies, properties of the join definitions to reduce redirection costs, etc. Moreover, most of the previous works report their results based on simulations, while we report our results based on a working distributed system.

[28] experimentally compares five types of query shapes and various execution strategies based on the PRISMA/DB system [27]. However, it does not explore how to generate optimized parallel processing query plans. In this work, we propose algorithms to generate efficient parallel processing solutions.

8 Conclusion

In this work, we have revisited the common assumption that has been taken by practically all prior work in the literature, namely, to pursue maximal pipelined parallelism when processing multi-join query processing in parallel. We have shown both experimentally and via a cost analysis that the introduction of independent parallelism at the cost of reducing the pipeline can greatly impact the parallel performance. A heuristic-driven optimization algorithm for generating a new class of processing strategies incorporating independent parallelism and yet controlling its dependencies has been proposed in this paper. A working distributed query engine has been implemented. Experimental studies confirm our claim that maximal pipelined parallelism is not
always the best.

The observation we made in this work also sheds some new light on how best to optimize pipelined query plans in general given the optimization function is related to the total processing time. This optimization is bound to get increasingly attention due to new and growing research areas such as continuous query processing [1].

References


A M-way Processing Cost Model

We provide the m-way processing cost model. The assumptions we made are the same as discussed in Section 4.2.

Estimating Total Work. Assume join relations are divided into m groups (pipelines) connected by an m-way bushy tree. Without loss of generality, we assume all these groups are denoted by its join relation indices, \((0 \sim m_1), (m_1 + 1 \sim m_2), \ldots, (m_{m-1} + 1 \sim n)\). The intermediate result of each group is represented by \(I_{m_1}, \ldots, I_{m_m}\). Correspondingly, we assume each group will be assigned \(k_{m_i}\) machines based on its building relation sizes. The final pipelined segment gets \(k_f\) machines.

The query result is also represented by \(I\). Without loss of generality, we also assume that \(I_{m_1}\) will be the probing relation of the final pipelined segment. Given these, the total work of building phase of an m-way bushy processing \((W'_b)\) and the total work of the probing phase \((W'_p)\) can be described by the following formulae.

\[
W'_b = (t_{\text{read}} + t_{\text{partition}} + t_{\text{network}} + t_{\text{build}}) * \left( \sum_{m=1}^{m} \sum_{j=m+1}^{m_1} |R_j| + \sum_{i=2}^{m} |I_{m_i}| \right)
\]

\[
W'_p = (t_{\text{read}} + t_{\text{partition}} + t_{\text{network}} + t_{\text{probe}}) * (|I_{m_1}| + |R_0| + \sum_{i=1}^{m-1} |R_{m_i+1}|)
+ t_{\text{network}} * \left( \sum_{i=1}^{m} \frac{k_{m_i} - 1}{k_{m_i}} \sum_{j=m+1}^{m+1} |I_j| + \sum_{i=2}^{m} k_f - 1 |I_{m_i}| \right)
+ t_{\text{probe}} * \left( \sum_{i=1}^{m} \sum_{j=m+1}^{m+1} |I_j| + \sum_{i=2}^{m} |I_{m_i}| \right)
\]

Estimating Processing Time. The overall processing time of the bushy tree can be treated as the sum of two phases. The first phase, \(T_{f1}\), estimates the time of processing all the pipelined segments (groups) with the results of these pipelines being directly fed into the building phase of the final pipelined segment. The second phase, denoted as \(T_{f2}\), estimates the time of probing the final pipelined segment and outputting the query results.

The processing time of each pipelined segment \((m_i)\) is composed by the following three components. (1) The building phase time of the building relations in \(m_i\), denoted by \(B_{m_i}\). (2) The probing phase time of the group \(m_i\), represented by \(P_{m_i}\). (3) The building time to the final pipelined segment from the output of group \(m_i\) \((I_{m_i})\), denoted as \(B'_{m_i}\). The processing time estimations of these components are given below.

\[
B_{m_i} = \max_{m_{i+1} \leq j \leq m_i} \left\{ \frac{f(k_{m_i})}{k_{m_i}} * |R_j| * (t_{\text{read}} + t_{\text{partition}} + t_{\text{network}} + t_{\text{build}}) \right\}
\]

\[
P_{m_i} = I_{\text{setup}} + \frac{W_{p_{m_i}}}{k_{m_i}} + I_{\text{delete}}
\]

\[
W_{p_{m_i}} = (t_{\text{read}} + t_{\text{partition}} + t_{\text{network}} + t_{\text{probe}}) * |I_{m_{i-1}}|
+ t_{\text{network}} * \left( \frac{k_{m_i} - 1}{k_{m_i}} \sum_{j=m+1}^{m_i-1} |I_j| \right)
+ t_{\text{probe}} * \left( \sum_{j=m+1}^{m_i-1} |I_j| \right)
+ (t_{\text{partition}} + t_{\text{network}} + t_{\text{build}}) * |I_{m_i}|
\]

\[
B'_{m_i} = \frac{f(k_f)}{k_f} * |I_{m_i}| * (t_{\text{read}} + t_{\text{partition}} + t_{\text{network}} + t_{\text{build}})
\]

The cost of the first phase is estimated by \(T_{f1} = \max_{1 \leq i \leq m} \{B_{m_i} + P_{m_i} + B'_{m_i}\}\). Note that the \(P_{m_i}\) and \(B'_{m_i}\) are actually processed in a pipelined fashion, we simply it by adding the cost directly.
The processing time of the second phase ($T_{f2}$) is composed basically the probing of first group ($I_{m_1}$), and the rest of the intermediate results. We estimate the time as $\frac{W'_i}{k_f}$. $W'_i$ can be described below.

$$W'_i = t_{network} \ast \frac{k_f - 1}{k_f} \sum_{i=2}^{m-1} |I_{m_i}| + t_{probe} \ast \sum_{m=2}^{m-1} |I_{m_i}|$$