# Practical Response Time Estimation in Parallel Relational Database Systems

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**Abstract**— An analytical approach to response time estimation in parallel relational database systems has been developed. It is based on a representation of database activity, in which queries are mapped to low-level patterns of resource consumption, capturing the execution logic of relational operators and mechanisms such as pipelined and partitioned execution. Resource usage profiles are mapped to open multi-class queueing networks. Queue waiting times are estimated using a heuristic rule, which labels resources as M/M/1 or M/G/1 queues. From these and the resource usage profile the average response time of a query is obtained. Synchronisation mechanisms such as pipelines between operators and partitioned parallelism are taken into account. The results of the analytical approach are compared against measurements of Informix XPS performance on a parallel system.

Index Terms— performance estimation, analytical model, queueing networks, pipeline parallelism, validation.

#### 1 Introduction

Parallel database systems are one of the more successful commercial applications of parallel computer technology. In recent years several prototype systems have been developed within research projects, e.g. Gamma [10], Bubba [5], DBS3 [2], EDS [36], Volcano [17], XPRS [18]. Many of the ideas from this work have been taken up by industry, and today, a number of major hardware and software vendors provide parallel database solutions (e.g. Informix, Oracle, IBM DB2, Teradata, NonStop SQL, Sybase, Microsoft SQL Server).

The ability to predict the performance of computer systems has always been important. This is especially true in the case of parallel database systems. It can be used before acquiring a parallel system to determine a suitable configuration, or once one has a parallel database system to tune its performance or decide how best to upgrade, e.g., there are many alternative ways of breaking up and distributing the data across the nodes and discs of a system, or scheduling an operator tree for execution. Performance prediction tools that automate the process of comparing and evaluating design decisions would benefit those involved with application design, system sizing and database administration. Due to the complexity of parallel databases, performance estimation of such systems is difficult. Correspondingly, the design of performance prediction tools and methodologies for this is a challenging task.

In estimating the performance of a parallel database system for a given set of queries, typical parameters of interest are the system bottlenecks, utilisation of different resources, maximum transaction throughput, and response time. Of these, the last is the most difficult to predict. This paper reports on a technique developed specifically for estimating response time in parallel database systems and its effectiveness compared with actual system measurements. The response time model is implemented as part of the core of a performance estimation tool called STEADY [38,9,37] that has been developed to predict the performance of three different parallel database systems.

The rest of this paper is organised as follows. Related work is discussed in Section 2. Section 3 is devoted to the technique used for response time estimation. The results from a validation exercise of the technique are reported in Section 4. Some conclusions and possible future work are given in Section 5.

# 2 Related Work

In the past research on measurement or estimation of response time of queries has ranged from studies of specific aspects of parallel database systems to commercial products for capacity planning or performance prediction. In the former case a number of researchers studying different aspects of parallel database operations such as skew, scheduling, hash-join algorithms, cache coherency, etc., have proposed and used models of query execution which produce an estimate of response time. In the latter case more general systems have been developed to predict performance for complete parallel database systems.

For example, in [33] data skew and its effect on response time of queries using joins is studied. A taxonomy of types of skew and a modelling methodology are proposed. A method for calculating response time is proposed and is applied to two existing hash join algorithms under skew. The response time analysis decomposes each algorithm into phases, each of which has a number of steps and can be partitioned across a number of processors. The time of each step is calculated based on the number of tuples involved, and is added to the total time of the resource responsible for the step (disc, cpu or communications). A bottleneck resource is established and its response time is used as the response time of the algorithm phase. The sum of the response times of all phases gives the response time of the algorithm.

In [14] scheduling in shared-nothing systems is studied and an analytical model of parallel execution and resource sharing is developed to estimate the response time of a given schedule (and, hence, the corresponding query). The model takes into account the overlapped use of multiple resources and captures both partitioned and pipeline parallelism. The response time of a parallel schedule is determined by either the slowest executing operator, or the load at the most heavily congested resource in the system, whichever is greater.

In [19] a method for efficient execution of multiple pipelined hash joins is proposed. The original execution tree is transformed into an allocation tree, each node of which is a multi-stage pipeline of hash joins. Several different hash-join strategies are studied. A combination of simulation and analytical techniques is used to derive the execution times of the different schemes. A (relatively simple) analytical formula is used to compute the execution time within each node. The formula sums the execution time for each phase of the joining of the relations i.e. reading, building hash tables, and probing. The simulation is used to traverse the allocation tree and carry out the join operations in parallel. Work on cache coherency policies uses models of response time to quantify the performance of alternative policies. For example, in [8] and [7] a comprehensive model of execution time is used, which attempts to account for the buffer hit probability, the concurrency control protocol, and the processing time and queueing delay at hardware resources. These three sub-models are analysed independently and their interactions are captured through a set of non-linear equations. An iterative process solves the system of equations to produce a solution to the overall model. Issues of pipelined execution are not considered.

There is some work specifically concerned with models of query execution which take account of the various factors affecting response time (e.g. query execution in a pipeline, contention for resources, effect of cache, etc.), and are intended for use in predicting overall response times of actual parallel database systems. An early attempt in the context of singleprocessor database systems is the work by Sevcik [26], which proposes an overall framework for predicting resource consumption, throughput and response time, and how these are affected by various physical and logical database design decisions.

A more recent example is the work by Salza et al [23,24], in which a modelling methodology for applications running on shared-nothing parallel database systems is developed. A workload model is used to characterise the database relations and set of transactions to be processed. From this, resource utilisation can be estimated. A buffer model is developed to capture the effect of caching. Through bottleneck analysis, maximum system throughput can be predicted. Response time is estimated as follows. First, by using queueing techniques, an estimate is made of how much the execution of each operator is slowed down by the concurrent running of other operators on the same node. A simplifying assumption of exponential service time distributions is made and standard queueing network techniques are applied. Second, the concurrent execution of different operators from the

same execution schedule is modelled, taking into account pipelined and partitioned execution. The approach is applied to a particular system (DB2 Parallel Edition on an IBM SP2 architecture), but no comparison results are reported.

M. Spiliopoulou et al [30,28,29] study the problem of estimating execution time for queries composed of multiple pipelined operators scheduled to run in a parallel database system. The intention is to incorporate the execution time prediction mechanism into a generic optimiser for parallel query processing. The developed cost model uses a comprehensive set of analytical formulae to compute the cost of a large variety of query operators, both in isolation and running in a pipeline. The model does not take into account contention for physical resources.

A number of commercial products also exist, which have a response time prediction capability. These are usually capacity planning or performance prediction tools, developed for existing parallel database systems. Tools vary in complexity from a simple set of cost formulae to a detailed simulation of the DBMS. Both analytical and simulation models are used. Examples include the Athene Performance Management System from Metron Technology [15,16,12] (an analytical capacity planning tool, developed for Oracle), a simulation-based performance prediction tool from Platinum Technology [22], an analytical tool from BEZ Systems [27,3], and a simulation tool for Oracle from SES Inc [25].

Another example is the DB2 Estimator [11] project at IBM, which has produced an analytical performance estimation tool, designed specifically for DB2 for OS/390 V5 and V6. It runs on a PC and calculates estimated costs using formulae obtained from an analysis of real DB2 code and performance measurements. For simple OLTP workloads, Estimator aims to predict the cost of 90% of the transactions within a 10% error. For decision support workloads these figures are 80% and 20%, respectively.

SMART (Simulation and Model of Application based Relational Technology) [6,1] is a

tool developed for predicting the performance of relational database applications using simulation. This tool is currently being superceded by its re-engineered successor, SWAP. SMART/SWAP is a sophisticated and versatile tool which is able to model complex real applications running on a variety of different platforms. Currently it models the performance of Oracle V7 and is in the process of being extended to model Oracle V8.

Most of this work suffers from two major drawbacks. Firstly, in many cases the service processes are assumed to be exponential whereas in practice they are not. This produces significant differences in terms of the results of the queueing network model. Secondly, most of the work is not supported by actual system measurements. The model described in this paper is an analytical one, intended for use in rapid estimation of response time, and it takes account of both these issues.

# **3** Estimating Response Time

The method proposed for estimation of query response time consists of three stages. The first stage (*preparation*) transforms a query into a suitable form. Database queries given as execution plans are reduced to a collection of low-level resource usage specifications, detailing patterns of resource consumption during query execution. Section 3.1 provides a brief overview of the reduction process and the *resource usage profile* representation.

The second stage (*mean resource response time estimation*) is concerned with the prediction of response time of individual hardware resources (CPUs, discs, etc), given their pattern of use during query execution (specified by the resource usage profile). For this stage, the profile is taken as the specification of a type of open, multi-class queueing network. Synchronisation between query execution phases, such as pipelined execution or partitioned parallelism, is not taken into account during this stage. Queueing network techniques are used to solve the network and obtain estimates for resource response times. Section 3.2 summarises the method used, which is based on a heuristic rule.

The third stage (*mean query response time estimation*), calculates the mean response time of a query, given its resource usage profile and estimated response times of individual resources. This is achieved through a process of traversing the resource usage profile and accumulating usage time according to the structure of the profile. Intra-operator parallelism such as pipelined or partitioned execution is taken into account during traversal and determines the way usage time is accumulated. Full details of this are given in Section 3.3.

# 3.1 Preparation

The starting point for the analysis is an *operator tree* [14,13], created from a query execution tree by decomposing each of its nodes into one or more constituent operator nodes, e.g. SCAN, AGGREGATE, BUILD, PROBE. Operator trees are generated by the compiler of the DBMS and given as input to the model. An operator tree together with a mapping of system processors to operators of the tree constitutes a *parallel execution schedule* [14].

An example parallel execution schedule is presented in Fig. 1. This shows an execution schedule for a query joining relations A, B and C and computing an aggregate function (e.g. max) over the resulting set of tuples. The execution schedule is typical of Informix XPS, and describes the following sequence of actions. Relation A is scanned and its tuples are used to build a hash table in preparation for joining with B. B is scanned and tuples from it are used to probe the hash table; the resulting join tuples then build a second hash table. C's tuples are scanned and used to probe the second hash table. Join tuples resulting from the last probe phase are aggregated. Redistribution of tuples takes place between a scanning and corresponding building phase in accordance with the requirements of the hash-join algorithm.

Arrows between nodes of the schedule represent flow of data as well as timing dependencies. Solid arrows denote *blocking*. For example, the probing of a hash table with tuples from C can begin only after the hash table has been built – hence the solid arrow between nodes "BUILD A $\otimes$ B" and "PROBE C". Dotted arrows denote *pipelined execution*.

For example, the scanning of tuples from A and their insertion into a hash table can be done in a pipeline (nodes "SCAN A" and "BUILD A"). Similarly a 3-stage pipeline can be established for the scanning of tuples from B, probing a hash table with them, and using resulting tuples to build a new hash table ("SCAN B", "PROBE B" and "BUILD A $\otimes$ B"). *Partitioned parallelism* is achieved by spreading an operator across multiple machine processors, as specified in the figure. For example, the scanning of A is carried out on PEs 0-2, as dictated by data placement constraints. Similarly, after redistributing the tuples selected from A, the building of the hash table takes place across the same three PEs.



Fig. 1 Parallel execution schedule

A process akin to "macro expansion" is then applied to each of the operators at the nodes of the parallel execution schedule. The process transforms each node into a detailed sequence of resource consumption items, termed here a *resource usage block*. The set of resource usage blocks and their interdependencies (taken from the execution schedule) form the *resource usage profile* of the query.

As an example, consider Fig. 2, which shows the resource usage block corresponding to the "SCAN A" node of the execution schedule. This represents the process of consecutive fetching of database pages from disc, selecting relevant tuples from the page, and sending them on to another block. This activity is given as a detailed sequence of resource usage.

Two types of notation are used in the body of the block (lines 4-28). The first is

concerned with the specific uses of individual resources for particular amounts of time. The usage of a particular hardware resource (e.g. processing unit - pu, system support unit - ssu, interconnect - net, disc) for a specified amount of time is represented in italics. Simple examples are lines 5 and 6, where resource *ssu* is used for time  $t_1$  and resource *pu* for time  $t_2$ . Line 12 gives an example of a resource (disc0) used for different amounts of time depending on the processing element (PEs 0-2) using the resource. The service time requirements for simple resource usage ( $t_i$ ) are extracted from the particular parallel database architecture modelled. For example,  $t_5$ ,  $t_6$ , and  $t_7$  in line 12 represent the average time spent by *disc0* of PE0, PE1, and PE2 in the process of fetching a data page, while  $t_3$  (line 11) is the amount of service time consumed by the *pu* resource during the page fetch.

1	BLOCK: SCAN A	16	group {
2	HOME: PE0, PE1, PE2	17	option {
3	RESOURCE TIME	18	pu t <sub>9</sub> : 0.6
4	group {	19	$pu t_{10}: 0.4;$
5	ssu t <sub>1</sub> ;	20	};
6	$pu t_2$	21	group {
7	} PE0:0.0;PE1:1.0;PE2:1.0;	22	$pu t_{11};$
8	loop {PE0:1710;PE1:1501;PE2:1802} {	23	$ssu t_{12};$
9	mean_shared_lock_waiting_time;	24	net t <sub>13</sub>
10	group {	25	}PE0:0.67;PE1:0.67;PE2:0.67
11	<i>pu</i> t <sub>3</sub> ;	26	}0.75
12	$disc0$ {PE0(t <sub>5</sub> ), PE1(t <sub>6</sub> ), PE2(t <sub>7</sub> )}:	27	}
13	} PE0:0.00;PE1:0.76;PE2:0.98;	28	}
14	loop {PE0:20;PE1:20;PE2:20} {	29	END TIME
15	nu t <sub>e</sub> :		

Fig. 2 Resource usage block

The second type of notation represents templates used to structure simple resource usage items. Templates are represented in bold and are of three types: *group*, *option*, and *loop*. The group template is used to designate a sequence of resource usage items, which has an associated probability of occurring. For example, the group template in lines 21-25 encloses three simple resource items, which represent the resource consumption required for the sending of a selected tuple for processing by the "BUILD A" operator. The probability associated with this group template is intended to rule out the physical sending across the interconnect of tuples destined for the processor where they are produced.

The option template is used to denote choice. It can have a number of branches, each

of which has an associated probability and contains a sequence of resource usage items. The grouped items occur together with the given probability. An example of option template use is given in lines 17-20, where the *pu* resource is used for either time  $t_9$  (with probability 0.6) or time  $t_{10}$  (with probability 0.4).

The loop template is used to designate repeated use of resources. The consecutive fetching of data pages is conveniently expressed with the loop construct since similar actions are carried out for each fetched page. The loop on line 8 represents the processing of 1710, 1501 and 1802 pages from PEO, PE1, and PE2, respectively. For each page read, a second loop (line 14) is used to represent the processing of individual tuples within the newly read page.

# 3.2 Mean Resource Response Time Estimation

The aim of the second stage of the method is to estimate the response time of individual hardware resources, given their pattern of use as specified by the resource usage profile. The approach used here is to take the resource usage profile of a query as specifying a queueing network [21,4], and "solve" the network for resource response time. The network queueing stations are the hardware resources of the machine and are treated as FCFS servers. The transitions among stations are determined by the templates within the resource usage blocks.



Fig. 3 Queueing network for simple example

An example of a queueing network for two simple resource usage blocks representing two hypothetical queries is given in Fig. 3. The pattern of resource consumption is represented as an open queueing network with two customer classes Q1 (solid line) and Q2 (dashed line), as illustrated. Shown are the three resources, the probabilities of transitions among them, and the service times for each customer class and resource.

Two difficulties present themselves when working with queueing networks in this context. One concerns the service time requirements at the queueing stations. Typically, exponentially distributed service times with a given mean are assumed. However, in practice assuming non-exponential service times is more realistic. The queueing network is thus composed of *non-exponential servers*, which renders it a non-product-form network.

The second difficulty is to do with the dependencies among resource usage blocks (e.g. pipeline execution and blocking). This imposes a synchronisation mechanism on the queueing network, which governs the timing of transitions between stations.

There have been numerous studies that address the non-exponential service time problem and lead to analytical approximation techniques [4]. Similarly, many studies are concerned with the synchronisation issues [4]: fork-join queues, blocking queues, etc. We are not aware of any studies, which address both issues simultaneously.

The approach used here is to tackle the two problems separately. In this section the first problem is addressed without regard for resource block inter-dependencies (i.e. the transitions between queueing centres are assumed to be of the "classical" type); the synchronisation issue is dealt with in the next section where overall query response time is estimated from the resource usage profile.

The non-product-form nature of the networks due to non-exponential service time requirements means that no exact analytical solutions for the response time or queue length of queueing centres can be found. However, numerous approximation techniques for non-product-form networks have been proposed in the literature [4]. In [31] we conduct a comprehensive study of approximation techniques for open multi-class networks. A range of

simple models was used and for each model the different approximation techniques were used to predict the response times for varying inter-arrival rates. The results were compared with those obtained from discrete event simulation. From the numerous examples considered in the study, best results were produced when the queueing network was considered to consist of a combination of M/M/1 and M/G/1 queueing centres.

This observation was incorporated into a simple heuristic rule, which accounted for all the cases tried. The rule is used to assign the resources of the network with either an M/M/1 or M/G/1 label. In accordance with the assigned label, the queue length or waiting time of each resource can be computed easily from the Khinchin-Pollaczek [21] formula. The heuristic rule used is based on the idea of resource dominance, and is summarised below.

**Definition** The dominance of a resource is a weighted average of its utilisation and its relative visit ratio, i.e.

$$dom(R_i) = \frac{1}{2} \times util(R_i) + \frac{1}{2} \times \frac{vr(R_i)}{\sum vr(R_j)}$$

where  $vr(R_i)$  is the average number of times resource  $R_i$  is visited for each arrival from outside. This corresponds to the visit ratio defined in [21] where the reference queue is  $q_0$  (the outside world).

Since both utilisation and relative visit ratio are numbers in the range 0 to 1, so also is dominance. Although relative visit ratio does not depend on query arrival rate, utilisation does. Thus, the utilisation of each resource must be taken for some particular query arrival rate (e.g. when the bottleneck resource has utilisation of 80%). The heuristic rule is now as follows:

- (a) Find the dominance of each resource.
- (b) If there is a single dominant resource, i.e. dom(R<sub>i</sub>) > dom(R<sub>j</sub>) for all j ≠ i, then according to the squared coefficient of variation of service time for each resource, label the resources as follows:

- label dominant resource  $R_i$  as max(M/M/1, M/G/1)
- label other resources  $R_i$  as min(M/M/1, M/G/1)
- (c) If there is more than one dominant resource, i.e.  $dom(R_{i_1}) = ... = dom(R_{i_n}) > dom(R_j)$ for all  $j \neq i_1...i_n$  then label all resources as min(M/M/1, M/G/1).
- (d) According to the label, use the Khinchin-Pollaczek formula to compute resource waiting time.

For parts (b) and (c) of the rule, the M/G/1 formula collapses into the M/M/1 formula when the squared coefficient of variation of the service time is 1.0; however, if the latter is less than 1.0, max(M/M/1, M/G/1) = M/M/1 whereas if it is greater than 1.0, max(M/M/1, M/G/1) = M/G/1.

# 3.3 Mean Transaction Response Time Estimation

Given the estimated queue waiting time of individual resources according to the above rule, the response time of the query can be estimated. This process is essentially a traversal of the resource block profile. During the traversal, response time is accumulated according to the structure of the blocks and the dependencies among blocks. In particular, account is taken of the templates within each block (*group*, *option*, *loop*, etc.), the partitioned parallelism, and the blocking or pipeline dependency among blocks. This is detailed below. First the response time estimation within a resource usage block not involved in a pipeline with others is discussed. This is extended to the case of such a block executing in parallel across several nodes (partitioned parallelism). Next, pipelined execution of blocks is considered. Finally, an overall response time for the complete resource usage profile of the query is obtained.

# 3.3.1 Templates

Within a resource usage block not involved in a pipeline with other blocks and running on a single processor, the response time of a sequence of one or more resource usage items is the sum of their response times. Each of these items may be a simple usage or a template (*group*,

*option* or *loop*). The response time of a simple usage involving resource r is the sum of the particular service time and the mean waiting time of the resource, as estimated by the rule. Denote this response time as RT(r).

For a *group* template with probability p (**group** { $usg_1$ ;...;  $usg_n$ } p), the response time is computed as  $RT(group) = p \times (RT(usg_1) + ... + RT(usg_n))$ . In the case of an *option* template (**option** { $usg_1:p_1;...; usg_n:p_n$ }), the response time is given by  $RT(option) = p_1 \times RT(usg_1) + ... + p_n \times RT(usg_n)$ .

The *loop* template denotes pipeline execution internal to the block. The response time of a loop usage is computed from the pipeline *initialisation* time and *production* time. A simple example of a pipeline is given in Fig. 4 to illustrate the idea. Three resources (*resA*, *resB* and *resC*) are organised in a pipeline that performs 10 iterations. The resources take 3, 4 and 1 unit of time per iteration, respectively. These represent the actual service time added to the mean waiting time for the resource as estimated by the rule; the service times are not shown. This applies to all subsequent examples.

The execution pattern is presented on the right hand side. The first iteration completes after 3 + 4 + 1 time units. Thereafter, an iteration completes every 4 time units as determined by *resB*, the pipeline bottleneck. This time (4 time units) is termed the *production time* of the pipeline and is the highest response time among the resources in the loop. The time from start to end of an iteration (in this case 3 + 4 + 1 = 8 time units) is termed the pipeline *initialisation time*. It is equal to the sum of the response times of all resources involved. The response time of a pipeline loop of *n* iterations is approximated as:

$$RT(loop) = \sum_{r \in res(L)} rt(r, L) + (n-1) \times rt(bn(res(L), L), L)$$

where:

- *L* is the set of resource usage items within the body of the loop;
- res(U) is the set of resources involved in the set of usage items U;

- rt(r, U) is the response time of resource r given its involvement in usage items from U;
- *bn*(*R*, *U*) is the bottleneck resource (the one with highest response time) among those in set *R* given their involvement in usage items from *U*.



Fig. 4 Pipelined execution

Informally, the response time is the sum of the initialisation time and the product of (n-1) and the production time. For the given example the formula evaluates to  $(3+4+1)+9\times4=44$  time units. This example is for a very simple loop. Typically, more complicated patterns of resource usage exist within the body of the loop, involving group or option templates. In this case, resource consumption is accumulated for each resource involved in the loop (by rt(r, U) and bn(R, U)) according to the templates and probabilities involved. The production and initialisation times are calculated based on these accumulated quantities.

In some cases, such as the resource block shown previously in Fig. 2, one loop template occurs within another. The inner loop is the last template of the outer loop. A simpler example is given in Fig. 5(a). The overall response time depends on the response times of resources from the inner and outer loops and the number of iterations of the inner loop. Suppose the inner loop, considered in isolation, has smaller overall response time than that of the bottleneck resource from the outer loop. In this case, the bottleneck of the outer loop determines the overall response time of the system. Otherwise, the inner loop, of which there are 15 iterations altogether, determines the performance. This type of analysis is carried out

in the discussion of pipelined blocks in Section 3.3.3. The same reasoning can be applied here. In order to do so, this case of loop usage is treated as being equivalent to two resource blocks working in a pipeline, as shown in Fig. 5(b). With this representation, the analysis of Section 3.3.3 can be reused.



Fig. 5 Example of loop within a loop

#### 3.3.2 Partitioned parallelism

The home field of a resource usage block specifies the processing elements used to execute the block and thus is an indication of intra-operation parallelism. For example, in Fig. 2, execution is spread over PE0, PE1 and PE2. The total response time of a block is obtained in two steps. First, the response time of each processing element from the block's home is calculated by accumulating time for the sequence of resource usage items in the body of the block. This is done following the procedure outlined above. Second, the processing element with the largest accumulated response time is found. This value is the response time of the block. Since all processing elements in its home are running in parallel, the block will complete when the processing element with the longest response time completes.

# 3.3.3 Pipelined parallelism

A pipeline may span two or three blocks within a query tree, which is typical of the queries used for validation in Section 4. Consider the case of two single-home blocks in a pipeline with one sending tuples and the other receiving them. A simple example of two such blocks is shown in Fig. 6. PE1 of block1 sends n tuples in a pipeline (in a similar way to the example in Fig. 4). The sending of a tuple requires work from all three resources *resA1*, *resB1* and

*resC1*. Let  $t_2$  be  $max(t_1, t_2, t_3)$ . The bottleneck resource of this pipeline is thus *resB1*; the pipeline production time is  $t_2$  and the initialisation time is  $t_1 + t_2 + t_3$ . Block2 on PE2 receives the *n* tuples and performs further operations using resources *resA2* and *resB2*. The production time of this pipeline is  $t_5$ . The full processing time needed to receive a tuple (the initialisation time of this loop) is  $t_4 + t_5$ .



Fig. 6 Two blocks in a pipeline

Time *T1* in Fig. 6 is the time at which the first iteration of the loop in block1 is complete and the first tuple has been sent to and is about to be received by block2. Similarly, *T2* is the time when the last tuple has just been sent and the work needed for its processing in block2 begins. Between times *T1* and *T2* the sending and receiving process execute together in parallel. The response time of the system depends on the relative speeds of sender and receiver. Let the bottlenecks of the two loops be  $r_1^*$  and  $r_2^*$ , i.e.  $bn(res(L_1), L_1) = r_1^*$  and  $bn(res(L_2), L_2) = r_2^*$ . Since the resources required by the two blocks are disjoint, if the sending process can send its tuples in less time than the receiving one can process them, i.e.  $rt(r_1^*, L_1) < rt(r_2^*, L_2)$ , then the total time of the system is governed by the speed of the receiver and may be approximated as  $\sum_{r \in res(L_1)} rt(r_2^*, L_2) + \sum_{r \in res(L_2)} rt(r, L_2)$ .

Here,  $L_1$  and  $L_2$  stand for the set of resource usage items in the loops of block1 and

block2, respectively. The first component in the formula accounts for the time prior to *T1*. The second covers the period between *T1* and *T2*, and the third from *T2* to the end of execution. For the example this would be  $(t_1 + t_2 + t_3) + (n - 1) \times t_5 + (t_4 + t_5)$ .

On the other hand, if the receiving process takes less time than the sending process, i.e.  $rt(r_1^*, L_1) > rt(r_2^*, L_2)$ , then the total time of the system is governed by the sender:

$$\sum_{r \in res(L_1)} rt(r, L_1) + (n-1) \times rt(r_1^*, L_1) + \sum_{r \in res(L_2)} rt(r, L_2)$$

If the resources required by the two blocks are not disjoint, then the calculation of the production time of each loop must take into account any usage of shared resources taking place in the other loop. Suppose the bottleneck resources of the two loops are  $r_1^*$  and  $r_2^*$ , i.e.

$$bn(res(L_1), L_1 \cup L_2) = r_1^*$$
 and  $bn(res(L_2), L_2 \cup L_1) = r_2^*$ .

Note that the bottlenecks are determined taking into account resource consumption from both loops. In the case when  $r_1^* \neq r_2^*$  a similar analysis to that for the case of disjoint resources applies. In other words, if  $rt(r_1^*, L_1 \cup L_2) < rt(r_2^*, L_2 \cup L_1)$ , then the response time of the system is determined by that of the receiver and may be approximated as:

$$\sum_{r \in res(L_1)} rt(r, L_1) + (n-1) \times rt(r_2^*, L_2 \bigcup L_1) + \sum_{r \in res(L_2)} rt(r, L_2)$$

In the case of a slower sender, the overall response time is determined by the sender's production time  $\sum_{r \in res(L_1)} rt(r, L_1) + (n-1) \times rt(r_1^*, L_1 \cup L_2) + \sum_{r \in res(L_2)} rt(r, L_2)$ .

If  $r_1^* = r_2^*$  (i.e. the bottleneck is a shared resource for the two loops), the response time of the sender/receiver system is approximated as the sum of the response times of the two loops:  $\sum_{r \in res(L_1)} rt(r, L_1) + (n-1) \times rt(r^*, L_1) + \sum_{r \in res(L_2)} rt(r, L_2) + (n-1) \times rt(r^*, L_2)$ , where  $r^* = r_1^* = r_2^*$ .

The idea that the overall time of a sender/receiver system is determined by the slower of the two can also be extended to apply to the case of multiple senders and receivers. This represents the case of a block with a multiple processor home sending tuples to another block executing on multiple processors, with senders and receivers working in parallel. Typically, redistribution of tuples takes place between the two blocks and different numbers of tuples may be sent/received by different sending/receiving loops, as illustrated in Fig. 7. The total time of this system is determined by the slowest of all sending and receiving loops.



Fig. 7 Multi-home blocks in a pipeline.

First, all sending loops are analysed to determine the slowest one, as follows. Consider sending loop  $L_S$ . Suppose there is no receiving loop  $L_R$  sharing resources with  $L_S$  and let  $bn(res(L_S), L_S) = r_{L_S}^*$ . The response time of  $L_S$  can be approximated as before:

$$RT(L_S) = \sum_{r \in res(L_S)} rt(r, L_S) + (iter(L_S) - 1) \times rt(r_{L_S}^*, L_S)$$
(1)

where iter(L) is the number of iterations in loop L.

If the set of processors on which the sending and receiving blocks operate are not disjoint (as in Fig. 7), a receiving loop,  $L_R$ , may share resources with  $L_S$ . In this case, let  $r_{L_S}^*$  be the bottleneck resource of  $L_S$ , taking into account any shared use. Formally,

$$r_{L_s}^* = BN(res(L_s), (L_s, iter(L_s)), (L_R, iter(L_R))),$$
(2)

where  $BN(res(L_1), (L_1, n_1), (L_2, n_2))$  returns resource  $r_i \in res(L_1)$  such that  $n_1 \times rt(r_i, L_1) + n_2 \times rt(r_i, L_2) > n_1 \times rt(r_j, L_1) + n_2 \times rt(r_j, L_2)$  for all  $j \neq i$ . Then the response time can be approximated according to the following formula, which takes into account the use of  $r_{L_s}^*$ 

within both loops:

$$RT(L_{S}) = \sum_{r \in res(L_{S})} rt(r, L_{S}) + (iter(L_{S}) - 1) \times rt(r_{L_{S}}^{*}, L_{S}) +$$
(3)  
$$iter(L_{R}) \times rt(r_{L_{S}}^{*}, L_{R})$$

The same analysis can be carried out on each receiving loop,  $L_R$ , in order to obtain an estimation of its response time,  $RT(L_R)$ .

The overall response time of the system of senders and receivers is approximated as follows. Let  $L_S$  and  $L_R$  be the sending and receiving loops with the highest response times as determined by the above procedure. Let their bottlenecks be  $r_{L_S}^*$  and  $r_{L_R}^*$ , respectively. Suppose that  $r_{L_S}^* \neq r_{L_R}^*$ . If  $RT(L_S) < RT(L_R)$  then:

$$RT = \underset{L_{S_i}}{\operatorname{avg}}(\sum_{r \in res(L_{S_i})} rt(r, L_{S_i})) + RT(L_R)$$
(4)

On the other hand, if  $RT(L_S) > RT(L_R)$ , then:

$$RT = RT(L_S) + \underset{L_{R_i}}{\operatorname{ver}} (\sum_{r \in res(L_{R_i})} rt(r, L_{R_i}))$$
(5)

In the above two formulae, the average initialisation time of all sending/receiving loops is used to account for the time needed for the first/last tuple to be sent/received.

If, however,  $r_{L_s}^* = r_{L_R}^*$ , then

$$RT = \sum_{r \in res(L_S)} rt(r, L_S) + (iter(L_S) - 1) \times rt(r_{L_S}^*, L_S) + \sum_{r \in res(L_R)} rt(r, L_R) + (iter(L_R) - 1) \times rt(r_{L_R}^*, L_R)$$

Consider the example given in Fig. 7, which can be used to illustrate the process of estimating the response time according to the above analysis. Consider first each sending loop. The loop on PE1 shares a resource (*resB1*) with the receiving loop on the same node and therefore (2) and (3) are used to calculate the response time. The bottleneck resource of this loop ( $r_{L_s}^*$ ), calculated according to (2), is the shared resource *resB1*, since its combined usage within both loops is greater than those of the three other non-shared resources (*resA1*,

*resC1* and *resD1*). The response time of the sending loop can be calculated from (3) as  $(1+3+4)+(16-1)\times 3+4\times 2=61$ .

The sending loop on PE2 does not share any resource usage, and its response time can be calculated from (1). The bottleneck resource of this loop is resB2 (with response time of 6) and therefore the response time of this sending loop is  $(2+6+4) + (8-1)\times 6 = 54$ .

Consider next the receiving loops. Their response times are estimated in a similar way. The receiving loop on PE3 does not share resources with a sending loop and evaluates (via (1)) to  $(4+5+2) + (20-1)\times 5 = 106$ . The receiving loop on PE1, however, shares resource *resB1* with the sending loop on the same node. Resource *resB1* is the bottleneck of this loop and the loop's response time was computed above as 61. Now consider the sending and receiving loops with the highest response times. These are the sending loop on PE1 with response time of 61 and bottleneck *resB1* and the receiving loop on PE3 with response time of 106 and bottleneck *resB3*. Since the two bottlenecks are different, and the receiving loop has a higher response time, the overall response time of the system may be approximated according to (4) as avg(1+3+4, 2+6+4) + 106 = 10 + 106 = 116.

This approach can be extended to a pipeline of three or more stages, as follows. First, the slowest stage of the pipeline is identified. This stage could be the initial sending stage of the pipeline, an intermediate sending/receiving stage, or the final receiving stage. The response time of each stage of the pipeline is determined by the response time of the slowest sending/receiving loop within the stage, taking account of shared resource usage as needed. Then, the total time of the pipeline is approximated as the total time of the slowest stage, plus the average initialisation times within each of the remaining phases, in a similar way to that explained above.

# 3.3.4 Overall response time

Thus far, the focus has been on estimating response time for various stages of the execution

schedule of the query. The response time for the query as a whole can be determined as follows.

The pipeline dependencies among blocks are considered in order to determine which blocks can be analysed together as a single pipeline. The full dependencies are considered to establish when a block (or group of blocks engaged in a pipeline) must wait for other blocks to complete before it can begin execution.

Consider the example execution schedule from Fig. 1. The overall response time of the query is the sum of the response times of three distinct phases within the execution schedule. The first phase is the two-stage pipeline involving blocks 'SCAN A' and 'BUILD A'. The second stage consists of the three-stage pipeline involving blocks 'SCAN B', 'PROBE B' and 'BUILD A $\otimes$ B'. The final phase is the three-stage pipeline involving blocks 'SCAN C', 'PROBE C', and 'AGGREGATE A $\otimes$ B $\otimes$ C'.

# **4** Validating the Technique

The ICL Goldrush MegaServer [34,35] is a parallel platform developed as a back-end database server to host several different database systems (Ingres, Oracle, Informix). In this section results from the validation of the model are discussed.

# 4.1 Goldrush and Informix XPS

The basic Goldrush hardware architecture consists of a number of Processing Elements (PEs) and Communication Elements (CEs) linked by a high speed Deltanet network. Each PE is connected to its own disc storage subsystem and runs a version of the UNIX operating system and DBMS code. A CE provides external links for Goldrush to clients via LANs. The particular Goldrush configuration used here has 1 CE, 8 PEs, 6 discs per PE and a cache of 16MBytes on each PE.

This type of architecture is an ideal platform for the shared-nothing Informix Extended

Parallel Server (XPS) [20]. It contains a set of internal components called co-servers, which are installed on each of the PEs of Goldrush. A co-server provides a user entry point to Goldrush on a given PE. Locks on data items and all data processing are managed locally within each co-server. Where deemed suitable, single queries are broken into subtasks and processed concurrently by threads within a single co-server and across co-servers. Data can be partitioned across discs and PEs so that parallel I/O operations can take place. The system can detect that certain partitions are irrelevant for a particular query and would not consider them when executing the query.

# 4.2 Tables

The experiments detailed in this section are carried out on a subset of the AS<sup>3</sup>AP benchmark [32] tables. In particular, the Uniques relation is used, the structure of which is given in Table 1. A number of variations of this relation were produced as shown in Table 2.

Attribute name	Attribute type	Name	Rows	Placement
key	integer(4)	un80	80	1 disc
int	integer(4)	un30k	30,000	1 disc
signed	integer(4)	un90k	90,000	1 disc
float	real(4)	un120k	120,000	1 disc
double	double(8)	un270k	270,000	1 disc of each PE
decim	numeric(18,2)	un540k	540,000	1 disc of each PE
date	datetime(8)	Table 2	Table 2 Uniques relations used	
code	char(10)			
name	char(20)			
address	varchar(20)			

 Table 1 Attributes of Uniques relations

*Un270k* and *un540k* are fragmented into 8 fragments by a simple hash function on the *key* primary key attribute with one hash fragment placed on a single disc of each of the PEs. Each tuple has a unique value for attributes *key* and *int*. Attribute *signed*, however, is modified from the benchmark specification so that for each relation tuples have *signed* values in the range 1 to 10 with an equal number of tuples for each value. Scans involving tables placed on one PE are not parallelised. Those involving tables *un270k* and *un540k*, however,

are performed in parallel by all co-servers containing the table data.

# 4.3 Queries

A number of different queries, classified in the following groups, were considered:

**Simple select-project-aggregate queries.** This group of queries exercises the ability of the method to work for the simple relational operations select, project and aggregate. Different tables (X), predicates (Y) and aggregation operators (max, count, etc) are used, as shown in Fig. 8(a).

**Simple hash-join queries.** Each of these queries (Fig. 8(b)) finds the maximum value of the *int* attribute from the result of a join of *un80* with the other *Uniques* tables where the *int* values are the same. Each of the other tables contains the tuples of *un80* so that the size of the resulting join is 80 tuples. The query employs a hash-join algorithm, which builds a hash table across all co-servers using *un80* and subsequently probes this with the tuples from X.

(1)	<b>select</b> max( <i>int</i> ) <b>from X</b> <b>where</b> <i>key</i> > 0;	select $max(un80.int)$ from $un80$ , X where $un80.int = X.int$			
(2)	<pre>select max(int), min(int) from X where Y;</pre>	$\frac{\mathbf{X} = un30k, un90k, un270k, un540k}{(b)}$			
(3)	<pre>select max(int), min(int), avg(int), count(*) from X where Y;</pre>	<pre>select max(int) from un30k where int &gt; (select max(key)</pre>			
	$\mathbf{X} = un30k, un90k, un270k, un540k;$ $\mathbf{Y} = "signed not in (1)",$ "signed not in (1, 3, 5)"	from $un30k$ where signed not in (1,3,5)); $max_1 = select max(key)$ from $un30k$			
	(a)	where <i>signed</i> not in (1,3,5); select max( <i>int</i> ) from <i>un30k</i> where int > max <sub>1</sub>			

(c)

Fig. 8 Queries used for validation

**Simple nested query and equivalent non-nested version.** This is a simple example of the use of a sub-query. The sub-query is used to return the maximum value of the key attribute from a subset of the tuples of *un30k*. This value is then used in the predicate of the

outer query. Note that this is not a correlated sub-query, and the execution plan indicates that Informix executes the sub-query first followed by the outer query. An equivalent "flat" formulation of the query was also considered, as shown in Fig. 8(c). The sub-query executes first as an independent query. Following it, a second query uses its result to compute the maximum int value. Although equivalent in their execution, the two queries have different response time characteristics, as discussed briefly in Section 4.5.

A union query. This query, shown in Fig. 9(a) illustrates the execution of the Informix union operator when there is opportunity for parallelism. The query groups together the maximum int values from tables *un30k* and *un270k*. It was expected that some form of parallelism would be used when executing the query. However, the execution plans indicate a sequential execution, with *un30k* scanned first, followed by the scanning of *un270k*, and followed by the union operator. To investigate this, the same query was executed with two *un90k* tables, specifically created on different processing elements and discs of Goldrush. Even in this case, where clearly the query can execute in half the time if the two relations are scanned in parallel, Informix chooses to perform the two scans in sequence.

A three-way hash-join. The execution schedule of this query was discussed in Section 3.1 and shown in Fig. 1. The query is shown in Fig. 9(b). Tables A, B and C, used in the discussion, correspond to *un80*, *un90k*, and *un30k*, respectively.

select max( <i>int</i> )	select max(un30k.int),			
from un30k	count(un30k.int)			
where $int > 0$	<b>from</b> un90k, un30k, un80			
union all	where un90k.key = un30k.key and un30k.key = un80.key			
select max(int)				
from un270k	(b)			
where $int > 0$				
(a)				

#### Fig. 9 Queries used for validation

### 4.4 Taking Measurements

Calibration of the model required running a carefully constructed set of queries in order to

estimate the cost of basic operations. The parallelised query execution plans produced by the optimiser were used to determine the way in which queries were broken down by the system and what basic operations were involved. Their costs were then obtained using the Informix XPS performance measuring tool onstat.

Once this was complete, a transaction generator was created to emulate a parallel database system workload with many users independently querying the data. It was used to fire queries from the communication element (CE) to a given co-server at a specified rate. Two generators were created. The first fired transactions with constant inter-arrival times and was used to determine the arrival rate for which the maximum throughput of the machine can be achieved. The throughput was computed by dividing the number of completed transactions by the time between the start time of the first transaction and the end time of the last one. Initially, the measured throughput is equal to the arrival rate at which the queries are fired. As the arrival rate is increased, a value is reached beyond which the measured throughput does not increase further. This is taken to be the maximum system throughput.

The second version of the transaction generator fired transactions with exponentially distributed inter-arrival times and was used to obtain transaction response times. The highest arrival rate used for the exponential generator was equal to 80% of the rate at which the maximum throughput (determined using the deterministic generator) was achieved. For each arrival rate the generator was set to fire transactions until 100 transactions were fired. The response time of each query was recorded, and the average was formed to compare against the value estimated by the analytical method. At the higher arrival rates, when queues would build quickly and the system would take longer to settle into a steady state, more than 100 queries were run when needed. Note that due to the good repeatability of measurements obtained from Informix, it was generally found necessary to repeat experiments no more than three times.

### 4.5 Results

**Simple select-project-aggregate queries.** Fig. 10 shows the maximum throughput prediction against actual measured throughput for one of the queries. The x-axis represents arrival rate and the y-axis – throughput. The prediction produced by the tool is a single figure: the maximum possible throughput, which is independent of the arrival rate. The predicted maximum throughput is higher than the one achieved by the system. This can be attributed to additional operating system overhead for which no account has been taken in the model, or to inaccuracies in the measured basic costs. Despite this, the predicted maximum throughput is an acceptable upper limit of the achieved maximum system throughput.



The response time predictions for this query are shown in Fig. 11. The x-axis measures arrival rate, while the y-axis represents the response time. The prediction is accurate to within 10% for the entire range of arrival rates.

All the experiments reported here were performed with the data present in cache. By comparison, consider a query of type (1) using table *un120k*, for which all data pages are read from disc. The results from two experiments performed are presented in Table 3. The arrival rates of the experiments correspond to 40% and 80% of the maximum throughput.

experiment	arrival rate	response	time (sec)	relative error (%)
	(trx/sec)	measured	predicted	
1	0.03	24.9	23.9	-4.0
2	0.06	74.8	83.4	11.5

Table 3 Results for query reading pages from disc

Simple hash-join queries. Fig. 12 shows the maximum throughput prediction against

actual measured throughput for this query on table *un540k*. Again the tool's prediction is higher than the achieved maximum throughput. Fig. 13 shows the response time prediction compared with measured response time.



**Simple nested query and equivalent non-nested version.** The graph in Fig. 14 shows the response time prediction and the corresponding measured values for the non-nested version of this query. The tool compares better with the non-nested query. The nested version of the query seems to involve extra resource usage, which can be attributed to the query optimiser 'un-nesting' the original query. The performance model does not currently account for this type of activity.

**Three-way hash-join query.** The response time results obtained for this query are presented in Fig. 15. The predicted values are within 15% of the measured values.



version of query



The results of the response time prediction method for all queries are summarised in Table 4. The table gives the number of experiments and indicates the magnitude of the relative percentage error computed from the predicted and measured response time for each query category.

	experiments							
Query	0% - 60% max throughput			70% - 80% max throughput				
	no. of	no. with	no. with	no. with	no. of	no. with	no. with	no. with
	exp.	err. below	err. 10-	err. 20-	exp.	err. below	err. 10-	err. 20-
		10%	20%	30%		10%	20%	30%
type 1	11	9	2	-	5	4	1	-
type 1 (from disc)	1	1	-	-	1	-	1	-
type 2	21	17	4	-	9	5	4	-
type 3	15	3	12	-	8	-	3	5
simple hash-join	8	-	8	-	6	-	3	3
non-nested	5	4	1	-	11	-	6	5
union	3	3	-	-	5	-	-	5
triple join	3	3	-	-	2	-	2	-

## Table 4 Summary of results

The experiments are divided into two groups depending on the value of the arrival rate for the experiment. The first category is for arrival rates that are up to and including 60% of predicted maximum throughput, while the second is for rates between 70% and 80%.

A two-query application. The results of a two-query application are also presented. The two chosen queries are a type (3) query on relation un30k with predicate 'signed not in (1, 3, 5)' and a type (1) query on relation un30k (see Fig. 8). The queries are assigned different frequencies: 70% for the first and 30% for the second. The measured and approximated response times of the two queries are presented in Fig. 16 and Fig. 17. There is a reasonable match between the approximation and the measured values.



#### 5 Conclusions

In this paper an analytical technique is presented for response time estimation of queries executing within a shared-nothing parallel database management system. This technique deals with the problem of non-exponential servers as well as with dependencies such as pipelining. It is based on a heuristic approach, used to obtain a better approximation of the behaviour of a network of non-exponential servers as used in parallel database systems than other approximation techniques. The techniques were validated by comparing their predictions against actual performance measurements on a parallel database system. The experiments indicate that the estimated and measured quantities are in reasonably good agreement. In particular, the predicted maximum throughput is typically higher than the measured maximum throughput by up to 15%. A possible explanation for this is additional resource consumption due to activity by non-DBMS processes or the operating system itself. Such activity is not accounted for within the models at present. Despite this, the approximation to the maximum system throughput is acceptable.

The experiments also show that the response time approximation technique produces reasonably good results. In particular, a total of 114 separate experiments were performed involving different transactions and arrival rates. In 96 of the experiments (over 84%) the relative error is less than 20%. In half of these, the error is less than 10%. For the remaining 18 experiments, the highest error is under 30%. Generally, the error is smaller for the lower transaction arrival rates and increases as the arrival rate is increased.

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