# Non-Standard Database Systems

Parallel Databases

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### Introduction

- Parallel machines are becoming quite common and affordable
  - prices of microprocessors, memory, and disks have dropped sharply
  - recent desktop computers feature multiple processors and this trend is projected to accelerate
- Databases are growing
  - large volumes of transaction data are collected and stored for later analysis
  - large objects like multimedia data are increasingly stored in databases
- Large-scale parallel database systems increasingly used for:
  - storing large volumes of data
  - processing time-consuming decision-support queries
  - providing high throughput for transaction processing

### Parallelism in Databases

### Databases naturally lend themselves to parallelism:

- Parallel I/O: data can be partitioned across multiple disks.
- Parallel execution: execute individual relational operations in parallel
  - e.g., sort, join, aggregation can be executed in parallel
  - each processor can work independently on its own data partition
- Queries are expressed at the logical level and in a high level language:
  - SQL is declarative and is translated to relational algebra
  - separation of logical and physical level makes parallelization easier
- Different queries can run in parallel:
  - concurrency control takes care of conflicts

## Outline

- 1/O Parallelism
- 2 Interquery Parallelism
- Intraquery Parallelism
  - Intraoperation Parallelism
  - Interoperation Parallelism
- Query Optimization and System Design

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# I/O Parallelism

 Reduce the time required to retrieve relations from disk by partitioning the relations on multiple disks.

 Horizontal partitioning — tuples of a relation are divided among many disks such that each tuple resides on one disk.

## Horizontal Partitioning

Let *n* be the number of disks.

- Round-robin:
  - send the *i*-th tuple inserted in the relation to disk *i* mod n.
- Hash partitioning:
  - choose one or more attributes A as the partitioning attributes
  - choose hash function h with range 0...n-1
  - send tuple t with hash value i = h(t[A]) to disk i
- Range partitioning:
  - choose one or more attributes A as the partitioning attributes
  - choose a partitioning vector  $[v_0, v_1, \dots, v_{n-2}]$ 
    - tuples t with  $t[A] < v_0$  got to disk 0
    - tuples with  $v_i \le t[A] < v_{i+1}$  to to disk i+1
    - tuples with  $v_{n-2} \le t[A]$  go to disk n-1
  - Example: with partitioning vector [5,11] on attribute A, a tuple t with partitioning attribute value of t[A] = 2 will go to disk 0, a tuple with t[A] = 8 will go to disk 1, while a tuple with t[A] = 20 will go to disk 2.

- We distinguish three different types of data access:
  - 1. sequential scan: scan the entire relation
  - 2. point query: locate a specific tuple
    - predicate is equality, zero or one result tuple
    - e.g., tuple of relation r with r.A = 25 (A is a key)
    - multi point query: zero or more result tuples (A is not a key)
  - 3. range query: locate all tuples within a specified value range
    - e.g., all tuples of relation r with  $10 \le r.A < 25$ .

#### Round robin:

- Good for sequential scan:
  - all disks have almost an equal number of tuples
  - retrieval work is thus well balanced between disks
- Point queries and range queries are difficult to process
  - no clustering relevant tuples are scattered across all disks

### Hash partitioning:

- Good for sequential access
  - assuming hash function is good, and partitioning attributes form a key, tuples will be equally distributed between disks
  - retrieval work is then well balanced between disks
- Good for point queries on partitioning attribute
  - lookup single disk, leaving others available for answering other queries
- No clustering, so difficult to answer range queries

### Range partitioning:

- Provides data clustering by partitioning attribute value.
- Good for sequential access.
- Good for point queries:
  - lookup single disk, leaving others available for answering other queries
- Good for range queries on partitioning attribute:
  - lookup single or few disks
  - good if result tuples are from one to a few blocks of a disk
- Execution skew: affects range queries and multi point queries
  - if many blocks are to be fetched, they may still be fetched from one to a few disks: potential parallelism in disk access is wasted
  - e.g., partition by order date, then tuples with recent order dates will be accessed more frequently

## Partitioning a Relation across Disks

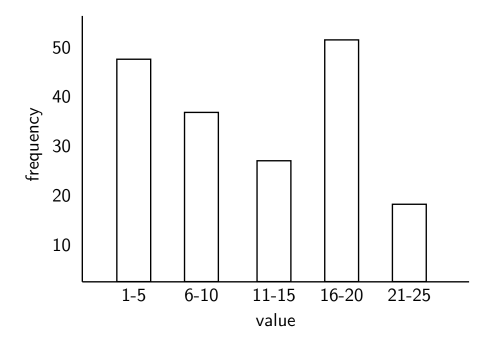
- If a relation contains only a few tuples which will fit into a single disk block, then assign the relation to a single disk.
- Large relations are preferably partitioned across all the available disks.
- If a relation consists of m disk blocks and there are n disks available, then the relation should be allocated to min(m, n) disks.

## Handling of Data Skew

- Distribution of tuples to disks may be skewed: some disks have many tuples, while others have fewer tuples.
- Skew limits speedup. Example:
  - relation with 1000 tuples is partitioned to 100 disks (10 tuples/disk)
  - expected speedup for scan: ×100
  - skew: one disk has 40 tuples  $\Rightarrow$  max. speedup is  $\times 25$
- Types of data skew:
  - Attribute-value skew:
    - Some values appear in the partitioning attributes of many tuples; all the tuples with the same value for the partitioning attribute end up in the same partition.
    - Can occur with range-partitioning and hash-partitioning.
  - Partition skew:
    - With range-partitioning, badly chosen partition vector may assign too many tuples to some partitions and too few to others.
    - Less likely with hash-partitioning if a good hash-function is chosen.

## Handling Skew using Histograms

- Balanced partitioning vector can be constructed from histogram in a relatively straightforward fashion
  - assume uniform distribution within each range of the histogram
- Histogram can be constructed by scanning relation, or sampling (blocks containing) tuples of the relation



## Handling Skew Using Virtual Processor Partitioning

- Skew in range partitioning can be handled elegantly using virtual processor partitioning:
  - create a large number of partitions (say  $10\times$  the number of processors)
  - assign virtual processors to partitions either in round-robin fashion or based on estimated cost of processing each virtual partition
- Basic idea:
  - If any normal partition would have been skewed, it is very likely the skew is spread over a number of virtual partitions.
  - Skewed virtual partitions get spread across a number of processors, so work gets distributed evenly.

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- 1 I/O Parallelism
- 2 Interquery Parallelism
- Intraquery Parallelism
  - Intraoperation Parallelism
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## Interquery Parallelism

- Queries/transactions execute in parallel with one another.
- Increases transaction throughput; used primarily to scale up a transaction processing system to support a larger number of transactions per second.
- Easiest form of parallelism to support, particularly in a shared-memory parallel database, because even sequential database systems support concurrent processing.
- More complicated on shared-disk or shared-nothing architectures:
  - locking and logging: coordinate by passing messages between processors.
  - data in a local buffer may have been updated at another processor.
  - cache-coherency has to be maintained: reads and writes of data in buffer must find latest version of data.

## Cache Coherency Protocol

- Example of a cache coherency protocol for shared-disk systems:
  - before reading/writing to a page, the page must be locked in shared/exclusive mode
  - on locking a page, the page must be read from disk
  - before unlocking a page, the page must be written to disk if it was modified.
- More complex protocols with fewer disk reads/writes exist.
- Cache coherency protocols for shared-nothing systems are similar. Each database page is assigned a home processor. Requests to fetch the page or write it to disk are sent to the home processor.

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## Intraquery Parallelism

- Execution of a single query in parallel on multiple processors/disks;
  important for speeding up long-running queries.
- Two complementary forms of intraquery parallelism:
  - Intraoperation Parallelism parallelize the execution of each individual operation in the query.
  - Interoperation Parallelism execute the different operations in a query expression in parallel.
- Intraoperation parallelism scales better with increasing parallelism because the number of tuples processed by each operation is typically more than the number of operations in a query.

## Parallel Processing of Relational Operations

- Our discussion of parallel algorithms assumes:
  - read-only queries
  - shared-nothing architecture
  - n processors,  $P_0, \ldots, P_{n-1}$ , and n disks  $D_0, \ldots, D_{n-1}$ , where disk  $D_i$  is associated with processor  $P_i$ .
- If processor has multiple disks: simulate a single disk  $D_i$ .
- Shared-nothing architectures can be efficiently simulated on shared-memory and shared-disk systems.
  - Algorithms for shared-nothing systems can thus be run on shared-memory and shared-disk systems.
  - However, some optimizations may be possible.

## Parallel Sort/1

### Range-Partitioning Sort

- Choose processors  $P_0, \ldots, P_{m-1}$ , where  $m \leq n$  to do sorting.
- Create range-partition vector with *m* ranges, on the sorting attributes
- Redistribute the relation using range partitioning
  - all tuples that lie in the  $i^{th}$  range are sent to processor  $P_i$
  - $P_i$  stores the tuples it received temporarily on disk  $D_i$
  - this step requires I/O and communication overhead
- Each processor  $P_i$  sorts its partition of the relation locally.
- Each processors executes same operation (sort) in parallel with other processors, without any interaction with the others (data parallelism).
- Final merge operation is trivial: range-partitioning ensures that, for  $0 \le i < j < m$ , the key values in processor  $P_i$  are all less than the key values in  $P_j$ .

# Parallel Sort/2

### Parallel External Sort-Merge

- Assume the relation has already been partitioned among disks  $D_0, \ldots, D_{n-1}$  (in whatever manner).
- Each processor  $P_i$  locally sorts the data on disk  $D_i$ .
- Sorted runs of processors are merged to get the final sorted output.
- Parallelize the merging of sorted runs as follows:
  - The sorted partitions at each processor  $P_i$  are range-partitioned across the processors  $P_0, \ldots, P_{m-1}$ .
  - Each processor  $P_i$  performs a merge on the streams as they are received, to get a single sorted run.
  - The sorted runs on processors  $P_0, \ldots, P_{m-1}$  are concatenated to get the final result.

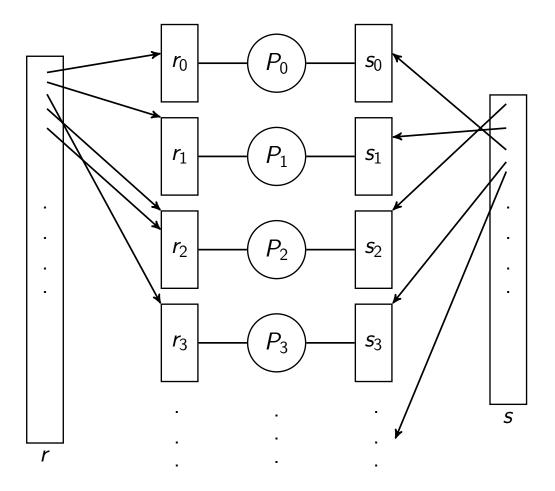
### Parallel Join

- The join operation requires pairs of tuples to be tested to see if they satisfy the join condition, and if they do, the pair is added to the join output.
- Parallel join algorithms attempt to split the pairs to be tested over several processors. Each processor then computes part of the join locally.
- In a final step, the results from each processor can be collected together to produce the final result.

## Partitioned Join/1

- For equi-joins and natural joins, it is possible to partition the two input relations across the processors, and compute the join locally at each processor.
- Let r and s be the input relations, and we want to compute  $r \bowtie_{r,A=s,B} s$ .
- r and s each are partitioned into n partitions, denoted  $r_0, r_1, \ldots, r_{n-1}$  and  $s_0, s_1, \ldots, s_{n-1}$ .
- Can use either range partitioning or hash partitioning.
- r and s must be partitioned on their join attributes (r.A and s.B), using the same range-partitioning vector or hash function.
- Partitions  $r_i$  and  $s_i$  are sent to processor  $P_i$ ,
- Each processor  $P_i$  locally computes  $r_i \bowtie_{r_i.A=s_i.B} s_i$ . Any of the standard join methods can be used.

# Partitioned Join/2



# Partitioned Parallel Hash-Join/1

### Parallelizing partitioned hash join:

- Assume s is smaller than r and therefore s is chosen as the build relation.
- A hash function  $h_1$  takes the join attribute value of each tuple in s and maps this tuple to one of the n processors.
- Each processor  $P_i$  reads the tuples of s that are on its disk  $D_i$ , and sends each tuple to the appropriate processor based on hash function  $h_1$ . Let  $s_i$  denote the tuples of relation s that are sent to processor  $P_i$ .
- As tuples of relation s are received at the destination processors, they are partitioned further using another hash function,  $h_2$ , which is used to compute the hash-join locally.

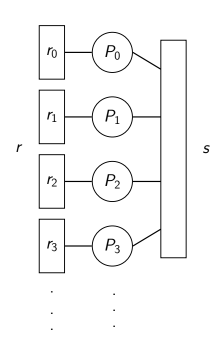
## Partitioned Parallel Hash-Join/2

- Once the tuples of s have been distributed, the larger relation r is redistributed across the m processors using the hash function  $h_1$ 
  - Let  $r_i$  denote the tuples of relation r that are sent to processor  $P_i$ .
- As the r tuples are received at the destination processors, they are repartitioned using the function  $h_2$ 
  - (just as the probe relation is partitioned in the sequential hash-join algorithm).
- Each processor  $P_i$  executes the build and probe phases of the hash-join algorithm on the local partitions  $r_i$  and  $s_i$  to produce a partition of the final result of the hash-join.
- Note: Hash-join optimizations can be applied to the parallel case
  - e.g., the hybrid hash-join algorithm can be used to cache some of the incoming tuples in memory and avoid the cost of writing them and reading them back in.

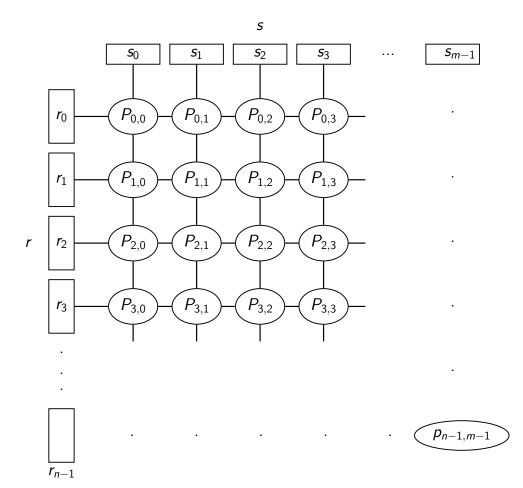
- Partitioning not possible for some join conditions
  - E.g., non-equijoin conditions, such as r.A > s.B.
- For joins were partitioning is not applicable, parallelization can be accomplished by fragment and replicate technique
- Special case asymmetric fragment-and-replicate:
  - One of the relations, say r, is partitioned; any partitioning technique can be used.
  - The other relation, s, is replicated across all the processors.
  - Processor  $P_i$  then locally computes the join of  $r_i$  with all of s using any join technique.

## Parallel Nested-Loop Join

- Assume that
  - relation s is much smaller than relation r
  - r is stored by partitioning (partitioning technique irrelevant)
  - there is an index on a join attribute of relation r at each of the partitions of relation r.
- Use asymmetric fragment-and-replicate, with relation s being replicated, and using the existing partitioning of relation r.
- Each processor  $P_j$  where a partition of relation s is stored reads the tuples of relation s stored in  $D_j$ , and replicates the tuples to every other processor  $P_i$ .
  - At the end of this phase, relation s is replicated at all sites that store tuples of relation r.
- Each processor  $P_i$  performs an indexed nested-loop join of relation s with the  $i^{th}$  partition of relation r.



Asymmetric fragment and replicate



Fragment and replicate

- General case: reduces the sizes of the relations at each processor.
  - r is partitioned into n partitions  $r_0, r_1, \ldots, r_{n-1}$ ; s is partitioned into m partitions,  $s_0, s_1, \ldots, s_{m-1}$ .
  - Any partitioning technique may be used.
  - There must be at least m \* n processors.
  - Label the processors as
  - $P_{0,0}, P_{0,1}, \ldots, P_{0,m-1}, P_{1,0}, \ldots, P_{n-1,m-1}$
  - $P_{i,j}$  computes the join of  $r_i$  with  $s_j$ . In order to do so,  $r_i$  is replicated to  $P_{i,0}, P_{i,1}, \ldots, P_{i,m-1}$ , while  $s_i$  is replicated to  $P_{0,i}, P_{1,i}, \ldots, P_{n-1,i}$
  - Any join technique can be used at each processor  $P_{i,j}$ .

- Both versions of fragment-and-replicate work with any join condition, since every tuple in r can be tested with every tuple in s.
- Usually has a higher cost than partitioning, since one of the relations (for asymmetric fragment-and-replicate) or both relations (for general fragment-and-replicate) have to be replicated.
- Sometimes asymmetric fragment-and-replicate is preferable even though partitioning could be used.

# Other Relational Operations/1

### Selection $\sigma_{\theta}(r)$

- If  $\theta$  is of the form  $a_i = v$ , where  $a_i$  is an attribute and v a value.
  - If r is partitioned on  $a_i$  the selection is performed at a single processor.
- If  $\theta$  is of the form  $l \leq a_i \leq u$  (i.e.,  $\theta$  is a range selection) and the relation has been range-partitioned on  $a_i$ 
  - Selection is performed at each processor whose partition overlaps with the specified range of values.
- In all other cases: the selection is performed in parallel at all the processors.

# Other Relational Operations/2

### Duplicate elimination

- Perform by using either of the parallel sort techniques
  - eliminate duplicates as soon as they are found during sorting.
- Can also partition the tuples (using either range- or hash-partitioning) and perform duplicate elimination locally at each processor.

### Projection

- Projection without duplicate elimination can be performed as tuples are read in from disk in parallel.
- If duplicate elimination is required, any of the above duplicate elimination techniques can be used.

## Grouping/Aggregation

- Partition the relation on the grouping attributes and then compute the aggregate values locally at each processor.
- Can reduce cost of transferring tuples during partitioning by partly computing aggregate values before partitioning.
- Consider the sum aggregation operation:
  - Perform aggregation operation at each processor  $P_i$  on those tuples stored on disk  $D_i$ 
    - results in tuples with partial sums at each processor.
  - Result of the local aggregation is partitioned on the grouping attributes, and the aggregation performed again at each processor  $P_i$  to get the final result.
- Fewer tuples need to be sent to other processors during partitioning.

## Cost of Parallel Evaluation of Operations

- If there is no skew in the partitioning, and there is no overhead due to the parallel evaluation, expected speedup will be *n*
- If skew and overheads are also to be taken into account, the time taken by a parallel operation can be estimated as

$$T_{part} + T_{asm} + max(T_0, T_1, ..., T_{n-1})$$

- $\bullet$   $T_{part}$  is the time for partitioning the relations
- $\bullet$   $T_{asm}$  is the time for assembling the results
- $T_i$  is the time taken for the operation at processor  $P_i$ 
  - this needs to be estimated taking into account the skew, and the time wasted in contentions.

## Interoperator Parallelism

- Two types of interoperation parallelism:
  - pipelined parallelism
  - independent parallelism

## Pipelined Parallelism

Example: Consider a join of four relations

$$r_1 \bowtie r_2 \bowtie r_3 \bowtie r_4$$

- Set up a pipeline that computes the three joins in parallel
  - Let  $P_1$  be assigned the computation of  $temp_1 = r_1 \bowtie r_2$
  - And  $P_2$  be assigned the computation of  $temp_2 = temp_1 \bowtie r_3$
  - And  $P_3$  be assigned the computation of  $temp_2 \bowtie r_4$
- Each operation can execute in parallel sending result tuples to the next operation even while it is computing further results
- Requires pipelineable (non-blocking) join evaluation algorithm (e.g., indexed nested loops join)

## Factors Limiting Utility of Pipeline Parallelism

- Pipeline parallelism is useful since it avoids writing intermediate results to disk
- Useful with small number of processors, but does not scale up well with more processors. One reason is that pipeline chains do not attain sufficient length.
- Cannot pipeline operators which do not produce output until all inputs have been accessed (e.g., aggregate and sort)
- Little speedup is obtained for the frequent cases of execution skew in which one operator's execution cost is much higher than the others.
- Advantage: avoids writing intermediate results to disk

## Independent Parallelism

• Example: Consider a join of four relations

$$r_1 \bowtie r_2 \bowtie r_3 \bowtie r_4$$

- Independent parallelism:
  - Let  $P_1$  be assigned the computation of  $temp_1 = r_1 \bowtie r_2$
  - And  $P_2$  be assigned the computation of  $temp_2 = r_3 \bowtie r_4$
  - And  $P_3$  be assigned the computation of  $temp_1 \bowtie temp_2$
  - $\bullet$   $P_1$  and  $P_2$  can work independently in parallel
  - $P_3$  has to wait for input from  $P_1$  and  $P_2$ 
    - Can pipeline output of  $P_1$  and  $P_2$  to  $P_3$ , combining independent parallelism and pipelined parallelism
- Does not provide a high degree of parallelism
  - useful with a lower degree of parallelism.
  - less useful in a highly parallel system.

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# Query Optimization/1

- Query optimization in parallel databases is significantly more complex than query optimization in sequential databases.
- Cost models are more complicated, since we must take into account partitioning costs and issues such as skew and resource contention.
- When scheduling execution tree in parallel system, must decide:
  - How to parallelize each operation and how many processors to use for it.
  - What operations to pipeline, what operations to execute independently in parallel, and what operations to execute sequentially, one after the other.
- Determining the amount of resources to allocate for each operation is a problem.
  - E.g., allocating more processors than optimal can result in high communication overhead.
- Long pipelines should be avoided as the final operation may wait a lot for inputs, while holding precious resources

# Query Optimization /2

- Use heuristics: Number of parallel evaluation plans much larger than number of sequential evaluation plans.
- Heuristic 1: No pipelining, only intra-operation parallelism:
  - Parallelize every operation on all processors
  - Use standard optimization technique, but with new cost model
- Heuristic 2: First choose most efficient sequential plan and then choose how best to parallelize the operations in that plan.
  - Volcano parallel database popularized the exchange-operator model
  - exchange operator is introduced into query plans to partition and distribute tuples
  - each operation works independently on local data on each processor, in parallel with other copies of the operation
- Choosing a good physical storage organization (partitioning technique) is important to speed up queries.

# Design of Parallel Systems/1

### Some issues in the design of parallel systems:

- Parallel loading of data from external sources is needed in order to handle large volumes of incoming data.
- Resilience to failure of some processors or disks.
  - Probability of some disk or processor failing is higher in a parallel system.
  - Operation (perhaps with degraded performance) should be possible in spite of failure.
  - Redundancy achieved by storing extra copy of every data item at another processor.

# Design of Parallel Systems/2

- On-line reorganization of data and schema changes must be supported.
  - For example, index construction on terabyte databases can take hours or days even on a parallel system.
    - Need to allow other processing (insertions/deletions/updates) to be performed on relation even as index is being constructed.
  - Basic idea: index construction tracks changes and "catches up" on changes at the end.
- Also need support for on-line repartitioning and schema changes (executed concurrently with other processing).

## Examples of Parallel Database Systems

- Teradata (1979), appliance, still large market share
- IBM Netezza (1999), appliance
- Microsoft DATAllegro / Parallel Data Warehouse (2003), appliance
- Greenplum (2005), Pivotal, open source
- Vertica Analytic Database (2005) commodity hardware
- Oracle Exadata (2008), appliance
- SAP Hana (2010), main memory, appliance