Advanced Databases

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Introduction Overview

Syllabus

- Introduction
- Relational databases overview
 - ► Data model, evaluation model
- Storage
 - ► Indexes. multidimensional data
- Query evaluation
 - ▶ Join evaluation algorithms, execution models
- Query optimisation
 - ► Cost models, search space exploration, randomised optimisation
- Concurrency control and recovery
 - ► Locking and transaction processing
- Parallel databases

Introduction Overview

Course logistics

• Lecturer: Stratis Viglas

• email: sviglas@inf.ed.ac.uk

• Days/Times: Mon & Thu, 11:10-12:00

• Location: 7 Bristo Square, Lecture Theatre 1

• Office hours: Mon, Thu 10:00-10:50 (or, by appointment)

▶ Room: IF, 5.11

• Course webpage: www.inf.ed.ac.uk/teaching/courses/adbs

• Mailing list: adbs-students@inf.ed.ac.uk

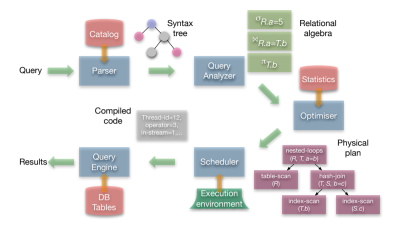
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Introduction Overview

Assignments and software

- Two programming assignments (slightly different assessment criteria for MSc students to account for different level offerings)
- The attica database system
 - ► Home-grown *RDBMS*, written in Java
 - ▶ Visit inf.ed.ac.uk/teaching/courses/adbs/attica to download the system and the API documentation
 - ▶ Both programming assignments will be using the attica front-end and code-base
- Plagiarism policy: You cheat, you're caught, you fail
 - ► No discussion

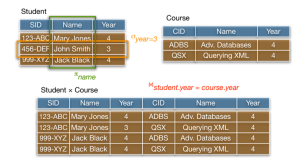
Query cycle



Relational databases overview

Data manipulation

- Operations to *isolate* a *subset* of a *single relation*: Selection (σ) , Projection (π)
- All set operations: Intersection, union, Cartesian product, set difference
- More *complex* operations: *Joins* (⋈), semi-joins, . . .



Three basic building blocks

- Attribute
 - ► A (name, value) pair
- Tuple
 - ► A *set* of attributes
- Relation
 - ► A set of tuples with the same schema

123-ABC

SID	Name	 Year
123-ABC	Mary Jones	 4

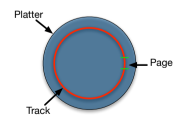
SID	Name	 Year	
123-ABC	Mary Jones	 4	
456-DEF	John Smith	 3	
999-XYZ	Jack Black	 4	

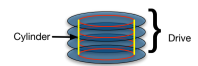
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Introduction

Relational databases overview

Data storage





- Disk drives are organised in records of 512 bytes
- The DB (and the OS) I/O unit is a disk page (typically, 4,096 bytes long)
- Pages (and records) are stored on tracks
- Tracks make up a platter (or a disk)
- Platters make up a drive
- The same tracks across all platters make up a cylinder
- The disk head (arm) reads the same block of all tracks on all platters

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Introduction

Relational databases overview

A bit of perspective

- The <u>dimensions</u> of the <u>head</u> are <u>impressive</u>¹. With a <u>width</u> of less than a <u>hundred nanometers</u> and a <u>thickness</u> of about <u>ten</u>, it flies above the platter at a <u>speed</u> of up to <u>15,000 RPM</u>, at a <u>height</u> that is the equivalent of <u>40 atoms</u>. If you start multiplying these infinitesimally small numbers, you begin to get an idea of their significance.
- Consider this little *comparison*: if the *read/write head* were a *Boeing* 747, and the *hard-disk platter* were the *surface of the Earth*
 - ► The *head* would *fly* at *Mach 800*
 - ▶ At less than *one centimeter* from the *ground*
 - ► And count every blade of grass
 - Making fewer than 10 unrecoverable counting errors in an area equivalent to all of Ireland

¹Source: Matthieu Lamelot, Tom's Hardware.

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Advanced Databas

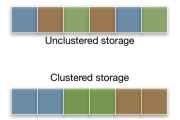
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Relational databases overview

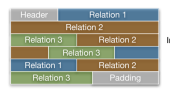
Advantages of clustering

- Scan a relation of X tuples, Y tuples per block
 - ► If *unclustered*, worst case scenario: *read X blocks*
 - ► Clustered: read X/Y blocks
- How about clustering disk blocks?
 - Reduces unnecessary arm movement

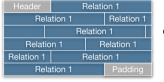


Storing tuples

- Every disk block contains
 - ► A header
 - ► Data (i.e., tuples)
 - ► Padding (maybe)
- Two ways of storing tuples
 - ► Either *interleave tuples* of multiple relations, or
 - ► Keep the tuples of the *same* relation clustered



Interleaved tuples



Clustered tuples

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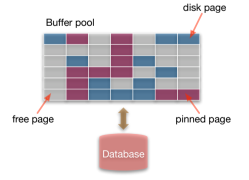
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Introduction

Relational databases overview

The buffer manager

- Though the data is on disk, real processing is in main memory
- Disk blocks are read and put into the *buffer pool*
 - ► A collection of *memory* pages
- The *buffer manager* manages the buffer pool
 - Keeping track of page references, replacing pages if full, . . .



Relational databases overview

Introduction

Relational databases overview

What does the buffer manager do?

- When a page is requested it:
 - ► Checks to see *if the page is in* the buffer pool; if so *it returns it*
 - ▶ If not, it *checks whether there is room* in the buffer pool: if so *it reads* it in and places it in the available room
 - ▶ If not, it picks a page for replacement; if the page has been "touched" it writes the page to disk and replaces it
 - ▶ In all three cases, it *updates the reference count* for the requested page
 - ▶ If necessary, it *pins the new page*
 - ▶ It returns a handle to the new page

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Relational databases overview

Why not use the OS

- The OS implements virtual memory, so why not use it?
 - ▶ Page reference patterns and pre-fetching: the RDBMS in most cases knows which page will be accessed later (think of a clustered sequential scan)
 - ► Different page replacement policies according to the reference pattern (check p. 322 of your book)
 - ▶ Page pinning: certain pages should not be replaced
 - ► Control over when a page is written to disk: at times, pages need to be forced to disk (we'll revisit that when discussing crash recovery)

Page replacement

- Least recently used (LRU): check the number of references for each page; replace a page from the group with the lowest count (usually implemented with a priority queue)
 - ► Variant: clock replacement
- First In First Out (FIFO)
- Most recently used (MRU): the inverse of LRU
- Random!

Storage and indexing Overview

Indexing and sorting

- Can be summarised as:
 - ► Forget whatever you've learned about indexing, searching and sorting in main memory (well, almost ...)
- Remember, we are operating over disk files
 - ► The main idea is to *minimise disk I/O* and *not number of comparisons* (i.e., complexity)
 - ▶ Just an idea: comparing two values in memory costs 4.91 · 10⁻⁸ seconds; Comparing two values on disk costs $18.2 \cdot 10^{-5}$ seconds (3) orders of magnitude more expensive.)

Indexing functionality

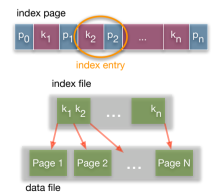
- Indexes can be used for:
 - ► Lookup queries (e.g., [...] where value = ''foo'')
 - ▶ Range queries (e.g., [...] where value between 20 and 45)
 - ▶ Join processing (after all, predicates are value-based, aren't they?)
- The above uses, and much more, are what we call access methods

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Storage and indexing One-dimensional indexing

Sorted indexes

- The basic idea:
 - ► An *index* is on an (collection of) attribute(s) of a relation (called the *index key*)
 - ▶ It is *much smaller* than the relation
 - ► Index pages contain (key, pointer) pairs
 - ★ key of the index
 - ★ pointer to the data page
 - ▶ Plus one additional pointer (low key)



Two main classes

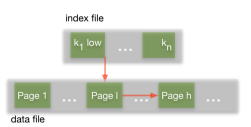
- Tree-structured indexes
 - ▶ Much like you would use a binary tree to search, but with a higher key-per-node cardinality
 - ▶ Retains *order*
 - ► Great for *range queries*
 - ▶ Both *one*-dimensional and *multi*-dimensional
- Hash-based indexes
 - ► Fully *randomized* (i.e., no order)
 - ► Great for single *lookup queries*

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How does it answer range queries?

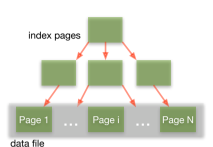
- Query is $low \le value \le high$
- Do a binary search on the index file to identify the page containing the low
- Keep scanning the data file until the *high key* is found
- All done!



Storage and indexing One-dimensional indexing

Potential problem (and the solution)

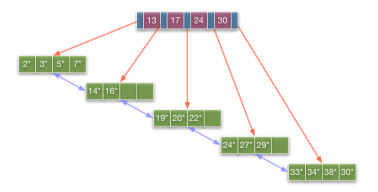
- The index is much smaller than the *relation*, but it's *still big*
- Binary search on it is still expensive
 - ▶ Remember. data is on disk
 - ► Have to access half the index file pages, plus the pages satisfying the predicate, all doing random I/O
- Why not build an index on the index?
 - ► Tree!



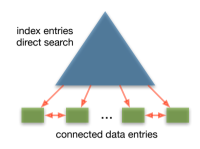
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B+tree example



B+trees: the most widely used indexes



- Insertion/deletion at log_f N cost (f = fanout, N = # leaf pages)
- Tree is height-balanced
- Minimum 50% occupancy (except for root)
- Characterised by its *order d*; each node contains d < m < 2dentries
- Equality and range searches are efficient

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B+trees in practice

- Typical *order*: 100, typical *fill-factor*: 67%
 - ► Average *fan-out*: 133
- Typical capacities
 - ► Height 3: 2,532,637
 - ► Height 4: 312,900,700 (!)
- The top levels can often be kept in memory
 - ▶ 1st level: 4,096, or 8,192 bytes (1 page)
 - ▶ 2nd level: 0.5, or 1MB (133 pages)
 - ▶ 3rd level: 62, or 133MB

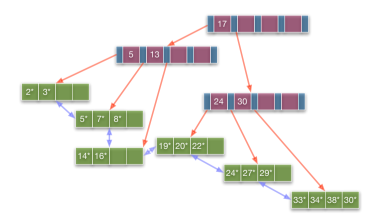
B+tree insertion

- Find correct leaf L
- Put data entry into L
 - ▶ If there is enough space in L, done!
 - ▶ If there is no space, L needs to be split into L and L'
 - ightharpoonup Redistribute entries evenly in L and L'
 - ► Insert index entry pointing to L' into the parent of L
- Ascend the tree recursively, splitting and redistributing as needed
- Tree tries to grow horizontally, worst case scenario: a root split increases the height of the tree

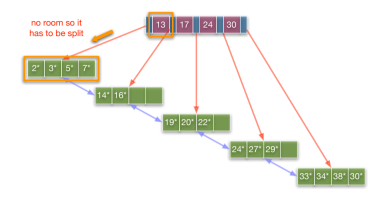
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B+tree insertion: 8*



B+tree insertion: 8*



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Insertion observations

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- Minimum occupancy is guaranteed at both leaf and non-leaf pages
- A leaf split leads to copying the key; a non-leaf split leads into pushing up the key (why?)
- The tree tries to first grow horizontally and if this is not possible, then vertically
 - ▶ In the example we could have avoided the extra level by redistributing
 - ▶ But in practice this is hardly ever done (why?)

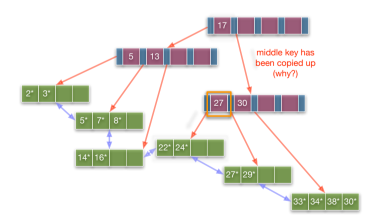
B+tree deletion

- Find leaf L where entry belongs
 - ► *Remove* the entry
 - ▶ If L is half-full, done!
 - ▶ If L only has d-1 entries
 - ★ Try to redistribute entries, borrowing from an adjacent sibling of L
 - * If redistribution fails, merge L and its sibling
 - * If merge has occurred, delete the entry for the merged page from the parent of L
- Ascend the tree recursively, performing the same algorithm
- Merge could propagate to the root, decreasing the trees height

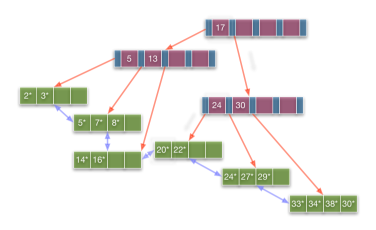
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B+tree deletion: 20*



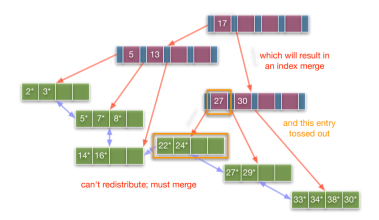
B+tree deletion: 19*



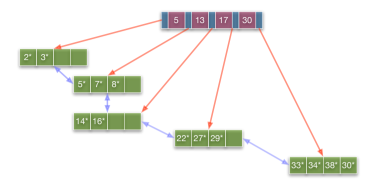
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B+tree deletion: 24*



B+tree after deletion of 24*



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Storage and indexing One-dimensional indexing

Hash indexes

- Hash-based indexes are good for equality selections, not for range selections
 - ▶ In fact, they *cannot support range* selections (why?)
- Static and dynamic techniques exist here as well
 - ► Trade-offs similar to those between ISAM and B+trees

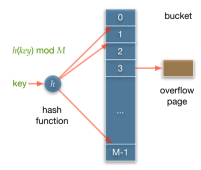
Summary of B+tree indexes

- Ideal for range searches, good for equality searches
- Highly *dynamic* structure
 - ► Insertions and deletions leave tree height-balanced, log N cost
 - ► For most typical implementations, height is rarely greater than 3 or 4, occupancy at 67%
 - ▶ Which means that the *index is almost always in memory*! (remember the buffer pool?)
 - ▶ Almost always better than maintaining a sorted file
 - ► The most optimised RDBMS structure

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Storage and indexing One-dimensional indexing

Static hashing



- Number of primary pages fixed
 - ► Allocated sequentially, never de-allocated
 - ► Overflow pages if needed
- $h(k) \mod M = \text{bucket}$ to which data entry with key k belongs (M =number of buckets)

Static hashing observations

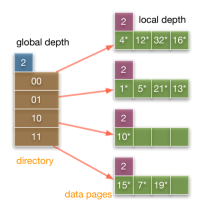
- The buckets contain the actual data!
 - ▶ But *only* the *key* is *hashed*
 - ► No secondary index like in the tree case
- The hash function must uniformly distribute the keys across all buckets
 - ▶ Lots of ways to *tune* the hash function
- Again, *long overflow chains* of pages will develop, and pretty soon we're doing random I/O
 - ▶ Need a dynamic technique (big surprise here...)
 - ► Extendible hashing to the rescue

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Storage and indexing One-dimensional indexing

Extendible hashing example

- Directory: array of size 4
- Key k, apply hash function h(k) and translate the result to binary
 - e.g., h(k) = 5 = 101
- Last global depth number of bits identify the bucket



Extendible hashing

- Problem: bucket (i.e., primary page) becomes full
- Solution: re-organize the file by doubling the number of buckets
 - ► Are you crazy? Reading and writing out everything is expensive!
 - ▶ Why not keep a directory of buckets and double only the directory? Only read the bucket that overflowed
 - ► Directory much smaller; operation much cheaper

Storage and indexing One-dimensional indexing

Global, local depth and doubling

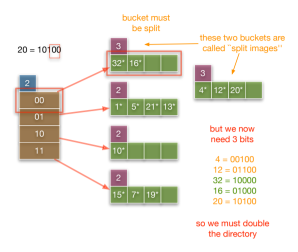
- Global depth (pertains to directory): maximum number of bits needed to tell which bucket an entry belongs to
- Local depth (pertains to bucket): maximum number of bits needed to tell whether an entry belongs to this bucket
- Before insertion (local = global) holds; if insertion causes (local > global) then directory needs to be doubled

Storage and indexing One-dimensional indexing

Storage and indexing

One-dimensional indexing

Insertion example: h(k) = 20



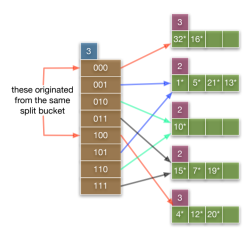
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Storage and indexing One-dimensional indexing

Extendible hashing observations

- Directory fits in memory: equality search answered with only one disk I/O (two in the worst case!)
 - ▶ 100MB file, 100 bytes/tuple, 4kB pages, 1,000,000 data entries, 25,000 directory entries: fits in memory!
 - ▶ If the *value distribution* is *skewed*, *directory grows large*
 - ► Same hash-value entries are a problem (why?)
- Deletion: if removal empties bucket, then it can be merged with split image; if each directory entry points to the same bucket as its split image, the directory is halved

Doubling the directory



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Storage and indexing One-dimensional indexing

Linear hashing

- Extendible hashing directory: even if it is small, it is still a materialised level of indirection
- Though the *number of buckets grows linearly*, the size of the directory grows exponentially
- Objective: no directory, linear growth
- Linear hashing gets the job done

Why one, when you can have many?

- Key idea: instead of having a single hash function and using a set of bits, have multiple hash functions
 - ▶ *Multiple* hash functions implement the *progressive doubling* of the directory
- Allocate buckets not when they become full, but whenever we reach some pretetermined load factor
- Single bucket allocation
- Each bucket allocation results in another hash function to be used
- Keep track of the number of buckets and the number of times the number of buckets has doubled
- Discard unused hash functions.

Storage and indexing One-dimensional indexing

Bookkeeping

- Two variables: Next, and Level
 - ▶ N points to the *bucket* to be *split next*
 - ▶ L keeps track of the number of *times* the *range* of the *hash function* has doubled
- Splitting proceeds in 'rounds
 - ▶ Round ends when all M_R initial (for round R) buckets are split
 - ▶ Buckets 0 to N-1 have been split
 - ▶ Buckets N to M_R have yet to be split
- Current round is L

In more detail

- Use a family of hash functions $h_0, h_1, h_2, ...$
 - $h_i(key) = g(key) \mod (2^i M)$
 - ightharpoonup M = initial number of buckets
 - g is some hash function (range is not [0, ..., N-1])
 - ▶ If $M = 2^{d_0}$, for some d_0 , h_i consists of applying g and looking at the last d_i bits, where $d_i = d_0 + i$.
 - $ightharpoonup h_{i+1}$ doubles the range of h_i (similar to directory doubling)

Storage and indexing One-dimensional indexing

Search and insert

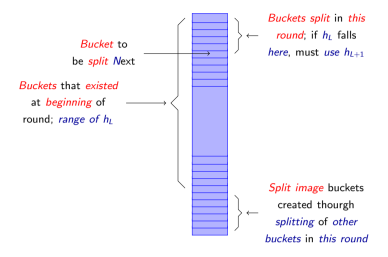
Search

- To *find* bucket for key K, find $h_L(K)$)
 - ▶ If $h_I(K) \in [N, ..., M_R]$, r belongs here
 - Else, r could belong to bucket $h_L(K)$ or bucket $h_L(r) + M_R$; we must apply $h_{L+1}(K)$ to find out.

Insert

- Find bucket as above, by applying h_L or h_{L+1}
- If bucket to insert is full
 - Add overflow page and insert entry
 - (Maybe) Split bucket N and increment N

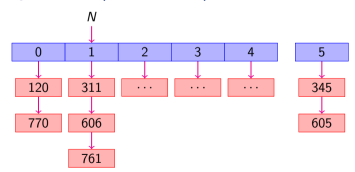
Linear hashing file



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Storage and indexing One-dimensional indexing

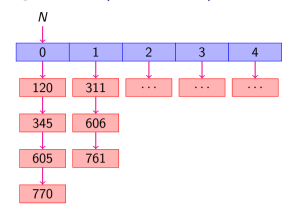
Splitting a bucket (0 in this case)



Hash functions

• $h_1(K) = K \mod 10$

Splitting a bucket (0 in this case)



Hash functions

• $h_0(K) = K \mod 5$

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Storage and indexing One-dimensional indexing

Algorithms in more detail

```
Lookup for key K
bucket := h_L(K);
if bucket < N then bucket = h_{L+1}(K)
```

Expansion

```
N := N + 1:
if N = M2^L then
L := L + 1; \quad N := 0;
```

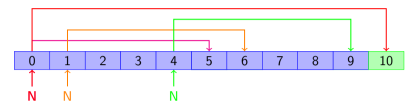
Contraction

```
N := N - 1;
if N < 0 then
 L := L - 1; \quad N := M2^{L} - 1;
```

Storage and indexing One-dimensional indexing

The expansion process (round 0)

Expansion N := N + 1: if $N = M2^L$ then $L := L + 1; \quad N := 0;$

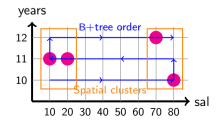


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Storage and indexing Multidimensional indexing

Why more than one dimensions?

- Single-dimensional indexes are not enough
 - ► Consider a *composite search* key *e.g.*, an index on ⟨*sal*, *years*⟩
 - ► The 2-dimensional space is linearised
 - ▶ We sort entries first by sal and then by years
- A multidimensional index clusters entries
 - **Exploits nearness** in multidimensional space.
 - ▶ Balanced index structures in multiple dimensions are challenging



 $\langle 10, 11 \rangle$, $\langle 20, 11 \rangle$

 $\langle 70, 12 \rangle$, $\langle 80, 10 \rangle$

Linear hashing observations

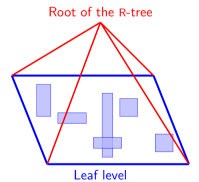
- Can choose any criterion to trigger split
 - ► Typically, we want to maintain some load factor
- Since buckets are split round-robin, long overflow chains do not develop!
- Doubling of directory in extendible hashing is similar
 - ► Switching of hash functions is implicit in how the number of bits examined is increased

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Storage and indexing Multidimensional indexing

The R-tree

- The R-tree is a tree-structured index that remains balanced on insertions and deletions
- Each key stored in a leaf entry is intuitively a box, or collection of intervals. with one interval per dimension



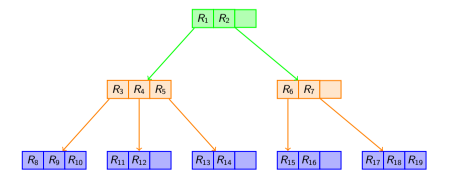
R-tree properties

- Leaf entry format: \(\langle \text{n-dimensional bounding box}\), pointer to record \(\rangle\)
 - ▶ Bounding box is the tightest bounding box for a data object
- Non-leaf entry format: \(\ n dim \ box, \ pointer to \ child \ node \)
 - ► The box covers all boxes in child node (in fact, subtree)
- All leaves at same distance from root
- Nodes can be kept 50% full (except root)
 - \blacktriangleright Can choose some parameter m that is < 50%, and ensure that every node is at least m% full

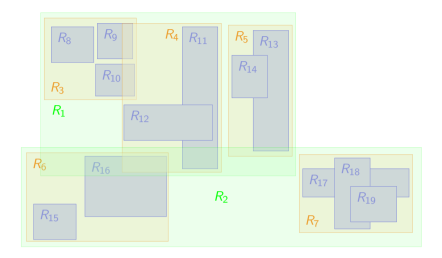
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Storage and indexing Multidimensional indexing

R-tree example (cont.)



R-tree example



Storage and indexing Multidimensional indexing

Search for objects overlapping box Q

Start at root

If current node is non-leaf

For each entry $\langle E, ptr \rangle$, if box E overlaps Q, search subtree identified by ptr If current node is leaf

For each entry $\langle E, rid \rangle$, if E overlaps Q, rid identifies an object that might overlap

Note

May have to *search several subtrees* at each node! (In *contrast*, a *B+tree* equality search goes to just one leaf.)

Storage and indexing Multidimensional indexing

Insert entry $\langle B, ptr \rangle$

Start at root and go down to "best-fit" leaf L

Go to child whose box needs least enlargement to cover B; resolve ties by going to smallest area child

If best-fit leaf L has space, insert entry and stop. Otherwise, split L into L_1 and L_2

Adjust entry for L in its parent so that the box now covers (only) L_1 Add an entry (in the parent node of L) for L_2 . (This could cause the parent node to recursively split.)

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Storage and indexing Multidimensional indexing

Comments on R-trees

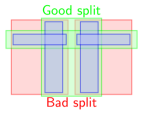
- Deletion consists of searching for the entry to be deleted, removing it, and if the node becomes under-full, deleting the node and then re-inserting the remaining entries
- Overall, works quite well for 2- and 3-D datasets
- Several variants (notably, R+ and R* trees) have been proposed; widely used
- Can improve search performance by using a convex polygon to approximate query shape (instead of a bounding box) and testing for polygon-box intersection.

Storage and indexing

Multidimensional indexing

Splitting a node

- The entries in node L plus the newly inserted entry must be distributed between L_1 and L_2
- Goal is to reduce likelihood of both L_1 and L_2 being searched on subsequent queries
- Redistribute so as to minimize area of L_1 plus area of L_2



Redistribution

Exhaustive algorithm is too slow; quadratic and linear heuristics are used in practice

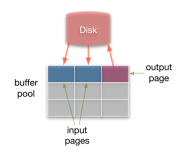
Storage and indexing Sorting

Overview

- Sorting is probably the most classic problem in CS
 - ► Simple idea: impose a total order on a set of values
- It is a *classic problem* in *databases* too
 - ► Remember *ISAM*? First step is to sort the file
 - ▶ In fact, if you're bulk loading a B+tree, you're better off sorting the file first
- Useful as well for duplicate elimination
- Useful for *join evaluation* (*sort-merge* algorithm)
- But what if I have a 1GB relation and 1MB of physical memory?
 - ► Remember, its all about *minimising I/O*
 - ► (Or, why your algorithms class didn't tell you the whole truth)

Two-way external merge sort

- Requires a maximum of three buffer pages and multiple passes over the data
- First pass: read one page, sort it. write it out
- Subsequent passes: read two pages, merge them, write out the result



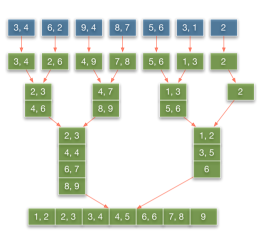
Storage and indexing Sorting

But why only three pages?

- We have an entire buffer pool of more than three pages, can we utilise it?
 - ► Yes: N-way merge sort
- To sort a file of N pages using B buffer pool pages:
 - ▶ First pass: sorted runs of B pages each $(\lceil \frac{N}{B} \rceil)$
 - ▶ Subsequent passes: merge B-1 runs (why?)

How it works

- Each pass will read and write each page in the file
- N pages, so the number of passes is $\lceil \log_2 N \rceil + 1$
- So, the total I/O cost is $2N(\lceil \log_2 N \rceil + 1)$



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Storage and indexing Sorting

What is the I/O cost?

- Number of passes: $1 + \lceil \log_{B-1} \lceil \frac{N}{B} \rceil \rceil$
- I/O cost: 2N· (Number of passes)
- For example: 108 pages in the file, 5 buffer pool pages
 - ▶ Pass 0: $\lceil \frac{108}{5} \rceil = 22$ sorted runs of 5 pages each
 - Pass 1: $\lceil \frac{22}{4} \rceil = 6$ sorted runs of 20 pages each
 - ► Pass 2: 2 sorted runs, 80 pages and 28 pages
 - ► Pass 3: final merge, done!

Storage and indexing Sorting

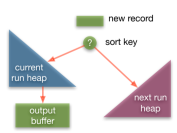
Storage and indexing Sorting

A bit of perspective

		257 * 4,096 = 1,052,672				
N	B=3	B=5	B=9	B=17	B=129	B=257
100	7	4	3	2	1	1
1,000	10	5	4	3	2	2
10,000	13	7	5	4	2	2
100,000	17	9	6	5	3	3
1,000,000	20	9	7	5	3	3
10,000,000	23	12	8	6	4	3
100,000,000	26	14	9	7	4	4
1,000,000,000	30	15	10	8	5	4

Storage and indexing Sorting

How does heapsort work?



- Keep *two heaps* in memory, one for each run (the *current* and the *next* one)
- Sum of memory needed for the two heaps equals the buffer size
- Keep adding to the *current* run *until* we are out of buffer space
- When buffer is full, swap heaps and *iterate*

Are we done?

- No! We can actually do much better than this
- Key observation: we are using main memory algorithm (e.g., quicksort) to sort pages in memory
 - ▶ But that *doesn't minimise I/O*, does it?
 - ▶ Wouldn't it be nice if we could *generate sorted runs longer than*
 - ► Solution: heapsort (a.k.a. tournament or replacement sort)

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Storage and indexing Sorting

The algorithm

```
Initialisation: read B pages into the current heap
while (not finished) do {
 while (r = lowest key from current heap) {
  write r to the current run
  \max = r
  get k from input
  if (k > max) insert k into current heap
  else insert k into next heap
 swap current and next heaps, max = 0
```

Storage and indexing Sorting

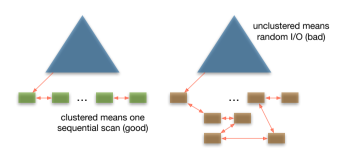
Storage and indexing Sorting

Heapsort observations

- What is the average length of a run?
 - ► Proven to be 2B (!)
- Quicksort is computationally cheaper
- But *heapsort* produces *longer runs*
 - ► Minimises 1/0
 - ▶ Remember, you should "forget" main memory methods when it comes to databases!

Storage and indexing Sorting

Clustered vs. unclustered storage



Good-old B+trees

- What if the table to be sorted has a B+tree index on sort field?
- Traverse the leaf pages and we're done!
 - ► Follow the *left-most pointers*, find the *low key*, *scan forward*
- Is this always a good idea?
 - ▶ If the *B*+tree is clustered, it's a great idea
 - ▶ Otherwise, it could lead to random I/O

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Storage and indexing Sorting

Summary of sorting

- Databases spend a lot of their time sorting
- In fact, they might dedicate part of their buffer pool for sorting data
 - ► Remember *pinning buffer pool pages*?
- External sorting minimises I/O cost
 - First you produce sorted runs, then you merge them
- The choice of internal sort matters as well
 - ► Yes, quicksort is computationally cheap
 - ► Though heapsort is computationally more expensive, it produces longer runs, which means less I/O
- Finally, *clustered B+trees* (when they exist) are a good way of *sorting* in one sequential scan

Overview

- A physical plan is what the query engine uses in order to evaluate queries
- In most cases, it is a tree of physical operators
 - ▶ Physical in the sense that they access and manipulate the raw, physical
- Plenty of ways to formulate this tree
 - ▶ Identifying the "best" tree is the job of the query optimiser

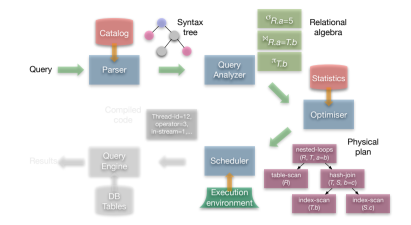
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Query evaluation Physical plans

Algebraic operators vs. physical operators

- A relational algebraic operator is a procedural abstraction of what should be retrieved
- The physical operator specifies how the retrieval will take place
- The same algebraic operator may map to multiple physical operators
- Physical operators for the same algebraic operator may be implemented using different algorithms
 - ▶ For instance: $join \rightarrow physical\ join \rightarrow sort-merge\ join$

Query cycle



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Query evaluation Physical plans

Example

SQL query

student.id, student.name select

from student, course

student.cid = course.cid and where

course.name = 'ADBS'

Algebraic expression

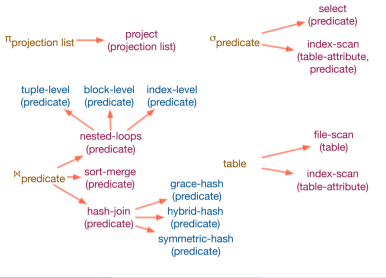
 $\pi_{student.id.student.name}$

(student ⋈_{student.cid=course.cid} $\sigma_{course.name='ADBS'}$ (course))

Algebraic operations

- $\pi_{student.id.\ student.name}$
- \bullet $\bowtie_{student.cid} = course.cid$
- $\sigma_{course.name} = `ADBS'$

Mappings to/of various physical operators



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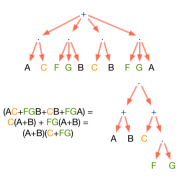
Query evaluation Physical plans

Physical plans

- Physical plans are trees of physical operators over the physical data
 - ▶ Just as arithmetic expressions are trees of arithmetic operators over numbers
- There are different ways of organising trees of physical operators
 - ▶ Just as there are different ways to organise a mathematical expression
- Physical plans are what produce query results

Math analogy

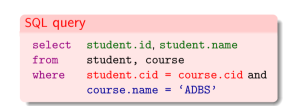
- Remember *factoring*?
- Same arithmetic expression can be evaluated in different ways
- If you map arithmetic expressions to infix notation, you have different "plans"

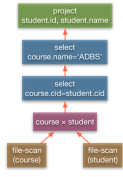


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Query evaluation Physical plans

Here's a plan





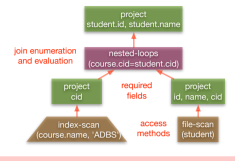
Query evaluation Physical plans

Observations

• Certain selection predicates can be incorporated into the access method

Query evaluation Physical plans

- If a *field* is *not needed*, it is *thrown out* (why?)
- More than two sources need to be combined (even through a Cartesian product)
- The query plan includes operators not present in the original query
- Yes, the query specifies what should be retrieved
 - ▶ But *how it is retrieved* is an *entirely different* business



SQL query

```
student.id, student.name
select
from
        student, course
where
        student.cid = course.cid and
        course.name = 'ADBS'
```

Here's a better plan

Query evaluation Physical plans

Issues

- Choice of order in which the physical operators are executed
 - ► Heuristics, access methods, optimisation
- Choice of algorithms whenever there are more than one
 - ► Again, optimisation (join enumeration, mainly)
- How are physical operators connected?
 - ► Different execution models
- What does a *connection* actually *imply*?
 - ► *Pipelining* (sometimes)
- What about *multiple readers* or even *concurrent updates* of the data?
 - ► Concurrency control (be patient ...)
- Finally, how is it all executed?
 - ► Query engine

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Query evaluation Physical plans

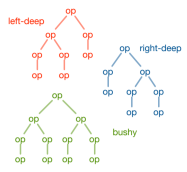
A note on duplicates

- The *relational model* calls for *sets of tuples*
- The query language (SQL) does not
 - ► Remember "distinct"?
- Sets can be guaranteed on base relations by specifying key (integrity) constraints
- But what happens with *intermediate results*?
 - ► Set semantics are lost, intermediate results have bag semantics
 - ▶ But set semantics can always be imposed; they are just more expensive to ensure

Query evaluation Physical plans

Query evaluation Physical plans

Types of plan



- There are two types of plan, according to their shape
 - ► Deep (left or right)
 - ► Bushy
- Different shapes for different objectives

Query evaluation Physical plans

Summary

- A plan is what the query engine accepts as input
 - ▶ ... and what produces the query results
- The same algebraic expression can produce multiple plans
 - ▶ Because the *same algebraic operator* maps to *multiple physical* operators
- A physical operator implements an evaluation algorithm
- A physical plan does not necessarily contain all the algebraic operators of the query
 - ▶ More or fewer, depending on the available physical operators
- The optimiser chooses the best physical plan
- Types of plans are classified according to their shape and evaluation objectives

Plan objectives

- A deep plan is better for pipelining
 - ▶ Because, let's face it, it's a line!
- A bushy plan is better for parallel computation
 - ► Different branches can be executed concurrently
- But all of these depend on the algorithms chosen
 - ▶ And on the execution model

Overview

- Physical plans are trees of connected physical operators
- The execution model defines the interface of the connections
 - ► And *how data* is *propagated* from one operator to the next
- It also defines *how operators* are *scheduled* by the query engine
 - ▶ Different execution models map to different process execution paradigms

Operator connections

- Operator functionality: relation in, relation out
- The connections are the interface through which the input is read and propagated
- In fact, there is a producer/consumer analogy



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Query evaluation

Execution models

What happens in practice

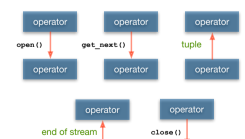
- Pipelining is simulated through the operator interface
- But different operations have different evaluation times
 - ▶ So there will be some need for buffering
- If we have joins, chances are the plan will block
 - ▶ We will see why that happens when talking about join algorithms

Pipelining

- Pipelining is the following process: read, process, propagate
- The opposite is to materialise intermediate results
- Pipelining works in theory, but in practice certain intermediate relations need to be materialised
 - ► This is called **blocking** (e.g., sorting)
- The benefits of pipelining include
 - ► No buffering
 - * No intermediate relation is materialised
 - ► Faster evaluation
 - ★ Since nothing is materialised, no disk I/O
 - ▶ Better resource utilisation
 - ★ No disk I/O means more in-memory operation

The iterator model

- Also known as a cursor
- Three basic calls
 - ▶ open()
 - ▶ get_next()
 - ► close()
- Have you ever accessed a database through external code?
 - ► For example: exec sql declare cursor in embedded SQL. ResultSets in Java/JDBC, etc.



operator

Query evaluation Execution models

Pure implementation

- The *iterator interface*, as described, is a *completely synchronous* interface
- A pure implementation means that all operators reside in the same

Query evaluation Execution models

- ► So *calls* can be *propagated downstream*
- But *certain operators* are "faster" than others
 - ▶ It could be the case that an asynchronous implementation could be more beneficial

get next() get_next() open() get next() open() select get next() open() scan

• All *calls* are propagated downstream

• The query engine makes the calls to the *topmost* operator only

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Call propagation

Query evaluation

Execution models

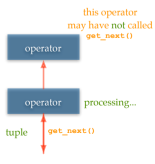
Different implementations

- The *iterator interface* is what *operators* use to *communicate*
- But how it is implemented, can be entirely different
 - ▶ The *reason* is that there might be *need for buffering*
 - ► Three possibilities
 - * Push model (buffering in the operator making the calls)
 - ★ Pull model (buffering in the operator accepting the calls)
 - * Streams (buffering in the connections)

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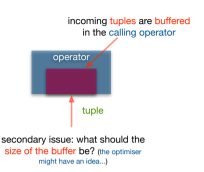
The push model

- Tuple propagation begins at the lower levels of the evaluation tree
- A lower operator propagates a tuple as soon as it is done with it
 - ► Does not "care" if the receiving operator has called get_next()



Buffering

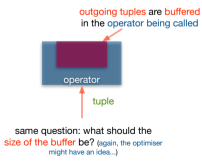
• The main issue: what happens if the *lower operator* has propagated the tuple before the operator above it has called get_next()?



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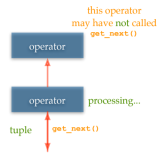
Query evaluation Execution models

Buffering — again



• The question this time: what happens if the lower operator is done processing the tuple before the operator above it calls get_next()?

The pull model



- The *inverse* of the *push model*
- If the *lower operator* is *done* processing a tuple it does not propagate it
 - ▶ It waits until the operator above it makes a get_next() call

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Query evaluation Execution models

The stream model

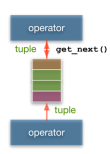
- The *connections* become first-class citizens
- Streams are queues of tuples connecting the operators
- Propagations and get_next() calls are synchronised on each stream



Query evaluation Execution models

Buffering — third time

- This time, there is no question!
- When the *lower operator* is done, it propagates the tuple
- When the top operator is ready. it calls get_next() on the incoming stream



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Query evaluation

Execution models

Benefits of each model

- Push model
 - ▶ *Minimises idle time* of the operators (why?)
 - ► Great for pipelining
- Pull model
 - ► Closest to a pure implementation
 - ▶ But still on-demand
- Streams model
 - ▶ Fully asynchronous to the operators, the synchronisation is on the streams
 - ► Highly parallelisable

Query evaluation Execution models

Why all this?

- Pure iterator implementation
 - ▶ If an operator receives get_next() and is not ready, it blocks
 - ► In fact, the entire plan blocks (why?)
 - ► Assume there is a *sort operation somewhere* in the plan
 - ★ Congratulations, your plan is officially blocked
- *Non-pure* implementations
 - ► Operators act (almost) independently of one another
 - ▶ Depending on the implementation of the interface (push-, pull-, stream-based) there are different benefits
 - * There could still be blocking, but the time during which a plan is blocked is minimised
 - ▶ It could lead to a each operator running in its own process thread
 - ★ Though this is not always a good idea

Summary

- A physical plan is a tree of connected operators
- Operators need to communicate data to one another
- The *iterator interface* is the *means* of this *communication*
 - open(), close(), get_next()
- As with any *interface* there are different ways of implementing it, known as execution models
 - Push model
 - ★ Data propagated as soon as they are available
 - ► Pull model
 - * Data retrieved on demand
 - Stream model
 - * Asynchronous communication on the connections between operators

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Query evaluation Join algorithms

Overview

- The *join operation* is *everywhere*
 - ► Any single query with two or more sources will need to have a join (even in the form of a Cartesian product)
 - ► So *common* that certain *DBMSs implement join indexes*
- As a consequence, a DBMS spends a lot of time evaluating joins
- Probably the most optimised physical operator
- A physical operator can be mapped to different algorithms
- As is always the case, a good join algorithm minimises I/O
- Choosing a join algorithm is not as straightforward; the choice might depend on
 - ▶ The *cardinality* of the input, its *properties* (clustered, sorted, *etc.*) and any available indexes
 - ► Available *memory*

Query evaluation

Join algorithms

Three classes of algorithms

- Iteration-based
 - ► Namely, *nested loops join* (in three flavours)
- Order-based
 - Sort-merge join (essentially, merging two sorted relations)
- Partition-based
 - ► Hash join (again, in three flavours)

Overview (cont.)

- Choosing how to evaluate a single join is different than choosing the order in which joins should be evaluated
- The query optimiser spends most of its time enumerating (ordering) the joins in a query
 - ▶ In fact, the order in which joins are evaluated affects the choice of algorithm
 - ► The two are largely interconnected (more on that when discussing query optimisation)

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Query evaluation Join algorithms

Terminology

- We want to evaluate $R \bowtie S$, shorthand for R.a = S.b
 - ► Also known as an equi-join
- In algebra: $R \bowtie S = S \bowtie R$
 - ▶ Not true for the physical join: $cost(R \bowtie S) \neq cost(S \bowtie R)$
- Three factors to take into account
 - ▶ Input cardinality in tuples T_R and pages P_R
 - ► *Selectivity factor* of the predicate
 - * Think of it as the percentage of the Cartesian product propagated
 - Available memory

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Query evaluation Join algorithms

Query evaluation Join algorithms

Nested loops join

- The simplest way to evaluate a join
- But it can still be optimised so that it minimises I/O
- Very useful for non-equi joins (the other two approaches will not work)
- Three variations
 - ► *Tuple-level* nested loops
 - ► Block-level nested loops
 - ► *Index* nested loops

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Query evaluation Join algorithms

What is the cost?

- One scan over the outer relation
- For every tuple in the outer relation, one scan over the inner relation
- If relations are *not clustered*, then
 - $ightharpoonup cost(R \bowtie S) = T_R + T_R \cdot T_S$
 - * Assume $T_R = 100,000, T_S = 50,000, \text{ then } cost = 5,000,100,000 \text{ I/Os}$
 - ★ At 10ms an I/O, that is 50,001,000 seconds, or, 14,000 hours

It doesn't get simpler than this...

Tuple-level nested loops

```
for each tuple r \in R do
 for each tuple s \in S do
  if r.a == s.b then add \langle r, s \rangle to the result
```

- R is the outer relation
- S is the inner relation

Query evaluation Join algorithms

What about clustered storage?

- Much, much better, I/O is at a page level
- So, the total cost will be
 - $ightharpoonup cost(R \bowtie S) = P_R + P_R \cdot P_S$
 - ▶ In the previous example, for 100 tuples per page, then $P_R = 1,000$, $P_S = 500$, cost = 501,000 I/Os
 - ▶ At 10ms an I/O, that is 5010 seconds, or about an hour and a half
- But we can *improve* that *even more*!
 - ► Block-level I/O and the buffer pool will work wonders

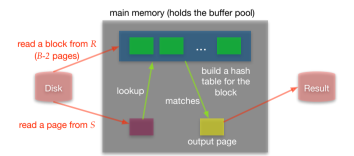
Here's an idea

- Assume we have *B pages available* in the buffer pool
- Read as many outer relation pages as possible; this constitutes a block
 - ▶ Put the pages of the block in the buffer pool, pin them
- Read the inner relation in pages
- Block size is B-2 pages (why?)
- Even more I/O savings

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Query evaluation Join algorithms

How it works



The Algorithm

Block-level nested loops

add $\langle r, s \rangle$ to result

```
Assumption: B dedicated pages in the buffer pool, block size is B-2
pages
for each block of B-2 pages of R do
  for each page of S do {
   for all matching in-memory tuples r \in R-block and s \in S-page
```

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Query evaluation Join algorithms

How much does it cost?

- The outer relation is still scanned once (P_R pages)
- The inner relation is scanned $\lceil \frac{P_R}{R} \rceil$ times
 - ► Each scan costs P_S I/Os

 - ► So, $cost(R \bowtie S) = P_R + P_S \cdot \lceil \frac{P_R}{B-2} \rceil$ ► Same example, $P_R = 1,000$, $P_S = 500$, assume a block size of 100 pages, then number of I/Os is 6,500
 - ► At 10ms per I/O, it will take 65 seconds

Key observation

- The inner relation is scanned a number of times that is dependent on the size of the outer relation
- So, the *outer relation* should be the *smaller one*
- Let's forget the ceilings and assume two relations: big and small
- Then we are *comparing*
 - $big + small \cdot \frac{big}{B-2}$
 - \rightarrow small + $big \cdot \frac{small}{B-2}$
- And big > small
- Remember, $cost(R \bowtie S) \neq cost(S \bowtie R)$ when it comes to physical operators

Query evaluation Join algorithms

The algorithm

Index nested loops

Assumption: there is an index on S.b

for each tuple $r \in R$ do for each tuple $s \in S$ where r.a == s.badd $\langle r, s \rangle$ to the result

• Predicate evaluation is an *index lookup* in the *index* over *S.b*

What if there is an index?

- Suppose the inner relation has an index on the join attribute
- We can use the index to evaluate the join
 - ▶ Remember, the *join predicate*, if we fix one of the join attribute values, is just a selection
- Scan the outer relation
 - ► Look at the join attribute's value and use it to perform an index lookup on the inner relation

Query evaluation Join algorithms

What is the cost?

- Depending on whether the outer relation is clustered or not, P_R or T_R I/Os to scan it
- Selectivity factor f: percentage of the Cartesian product propagated; this means that every outer tuple joins with $f \cdot T_S$ tuples
 - ► Depending on the index, each lookup will be, say, avg_lookup I/Os
- If R is clustered
 - $ightharpoonup cost(R \bowtie S) = P_R + T_R \cdot f \cdot T_S \cdot avg_lookup$
- If R is not clustered
 - $ightharpoonup cost(R \bowtie S) = T_R + T_R \cdot f \cdot T_S \cdot avg_lookup$

Index nested loops

- If the selectivity factor and the average lookup cost are small, then the cost is essentially a (few) scan(s) of the outer relation
- If the outer relation is the smaller one, it leads to significant I/O savings
- Again, it is the job of the query optimiser to figure out if this is the case

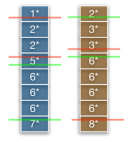
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Query evaluation

Join algorithms

How it works

- Key idea: there exist groups in the sorted relations with the same value for the join attribute
- We need to take that into account when merging
 - ► The *reason* is that we will have to do some backtracking when generating the complete result



Sort-merge join

- Really simple idea
- The join is evaluated in two phases
 - First, the two input relations are sorted on the join attribute
 - ▶ Then, they are merged and join results are propagated
- External sorting can be used to sort the input relations
- The merging phase is a straightforward generalisation of the merging phase used in merge-sort

Query evaluation Join algorithms

The algorithm

```
Merge-join
 r \in R, s \in S, gs \in S
 while (more tuples in inputs) do {
  while (r.a < gs.b) do advance r
                                        // a group might begin here
   while (r.a > gs.b) do advance gs
   while (r.a == gs.b) do \{
    s = gs // mark group beginning
    while (r.a == s.b) do // while in group
        add \langle r, s \rangle to the result; advance s // produce result
    advance r // move forward
              // candidate to begin next group
```

Query evaluation Join algorithms

Query evaluation Join algorithms

What is the cost?

- We know the cost of externally sorting either relation: $2 \cdot P_R \cdot \log P_R$, or $2 \cdot P_S \cdot \log P_S$
- The merge phase is essentially one scan of each sorted input: P_R or Ps (these scans are always clustered)
- $cost(R \bowtie S) = P_R \cdot (2 \cdot \log P_R + 1) + P_S \cdot (2 \cdot \log P_S + 1)$
 - ▶ Running example: $P_R = 1,000, P_S = 500, 100$ buffer pool pages to sort, the *number of I/Os is 7.500*
 - ▶ At 10ms an I/O, this is *one minute and fifteen seconds* (about the same as block nested loops)

Query evaluation

Join algorithms

Hash join

- Partition-based join algorithms
- Key idea: partition R and S into m partitions, R_i and S_i , so that every R_i fits in memory
 - ▶ Observation: *joining tuples* will fall into the *same partition*
- Then, for every R_i load it in memory, scan S_i and produce the join results
- Three flavours: Simple hash join, grace hash join, hybrid hash join

A few issues

- If there are <u>large groups</u> in the <u>two relations</u>, then we <u>may</u> have to <u>do</u> a lot of backtracking
 - ► Performance will suffer due to possible extra I/O
 - ► Hopefully, pages will be in the buffer pool
- Most relations can be sorted in 2-3 passes
 - ▶ Which means that we can compute the join in 4 passes max (almost regardless of input size!)
 - ▶ In fact, we can *combine* the *final merge of external sorting* with the merging phase of the join and save even more I/Os

Query evaluation Join algorithms

The simple algorithm

Simple hash join

Assumption: m partitions, each partition P_i fits in main memory

for all partitions $P_i, i \in [1, m]$

for each $r \in R$ read r and apply hash function $h_1(r.a)$

if r falls into P_i apply hash function $h_2(r,a)$ and put it in an in-memory hash table for Pi

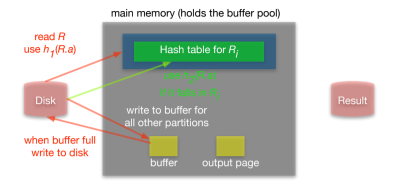
otherwise, write it back out to disk

for each $s \in S$ read s and apply hash function $h_1(s,b)$

if s falls into P_i apply hash function $h_2(s,b)$ and for all matching tuples $r \in P_i$, add $\langle r, s \rangle$ to the *result*

otherwise, write it back out to disk

How it works — partitioning R, iteration i



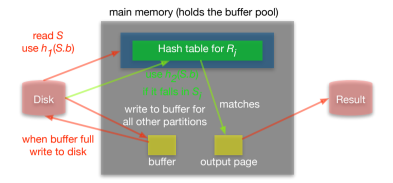
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Query evaluation Join algorithms

What is the cost?

- Assume equal partition sizes, input T, P_T pages
- For *m partitions*, we will make *m passes* over each input
 - ► For the first pass:
 - * Read P_T pages, write $P_T \frac{P_T}{m}$ pages: $2P_T \frac{P_T}{m}$ I/Os
 - ► For the second pass:
 - * Read $P_T \frac{P_T}{m}$, write $P_T \frac{P_T}{m} + P_T 2\frac{P_T}{m}$ pages: $2P_T 3\frac{P_T}{m}$ I/Os
 - ► Pass *i*: $2P_T (2i 1) \frac{P_T}{m}$ I/Os
- In the end, $m(m+1)P_T$ I/Os
- For two relations R and S, total cost is $m(m+1)(P_R + P_S)$
- Makes sense if *m* is *small*, or we have a lot of memory
- Effectively, this is nested loops join
 - ▶ But the number of iterations is decided by the number of partitions, not the input sizes!

How it works — paritioning and joining S, iteration i



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Query evaluation Join algorithms

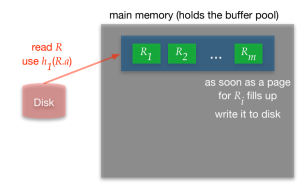
The "grace" algorithm

```
Grace hash join
 for each r \in R read r and add it to the buffer page for h_1(r,a)
 for each s \in S read s and add it to the buffer page for h_1(s,b)
 for i = 1, ..., m do {
  for each r \in R_i read r and insert it into a hash table using h_2(r.a)
  for each s \in S_i do {
    read s, probe the hash table using h_2(s.b)
    for all matching tuples r \in R_i add \langle r, s \rangle to the result
 clear hash table
```

Query evaluation Join algorithms

Query evaluation Join algorithms

How it works — partitioning R

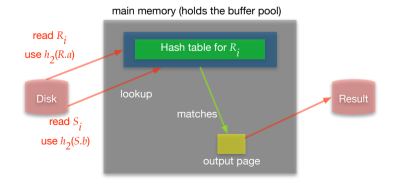


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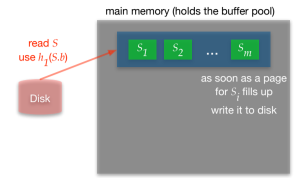
Query evaluation

Join algorithms

How it works — joining



How it works — partitioning S



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Query evaluation Join algorithms

What is the cost?

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- Scan R and write it to disk, so $2 \cdot P_R$
- Do the same for S, so $2 \cdot P_S$
- Read R in partition-by-partition, so P_R
- Scan S partition-by-partition and probe for matches, so P_S
- $cost(R \bowtie S) = 3 \cdot (P_R + P_S)$
 - ▶ Same example, $P_R = 1,000$, $P_S = 500$, cost is 4,500 I/Os
 - ▶ At 10ms an I/O the join will take 45 seconds to evaluate

Memory requirements

- Objective: the hash table for a partition must fit in memory
 - ► Minimise partition size by maximising number of partitions
- What are the *optimum sizes*?
 - For B buffer pool pages, maximum number of partitions m = B 1(why?)
- Size of each partition is $\left\lceil \frac{P_R}{R-1} \right\rceil$
- Size of the hash table is $\lceil \frac{f \cdot P_R}{R-1} \rceil$ (f = fudge factor to capture the increase in partition size due to the hash table)
- During the *probing phase*, in addition to the hash table, we need *one* page to read S, plus one page for output
 - ▶ So, $B > \left\lceil \frac{f \cdot P_R}{P-1} \right\rceil + 2 \Rightarrow B > \sqrt{f \cdot P_R}$

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Query evaluation

Join algorithms

How it works

- Suppose that $B (k+1) > \frac{f \cdot P_R}{k}$
 - ▶ We have *enough memory* during partitioning to *hold* an *in-memory* hash table of size B - (k + 1) pages
- Idea: keep R₁ in memory at all times
- While partitioning S, if a tuple falls into S_1 , don't write it to disk; instead probe the hash table for R_1 for matches
- For all partitions R_i , S_i , i > 2, continue as in hash join

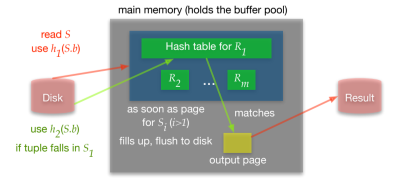
Hybrid hash join

- An improvement over hash join if there is extra memory
 - Minimum amount of memory for hash join $B > \sqrt{f \cdot P_R}$
 - ► Suppose that $B > \frac{f \cdot P_R}{k}$, for some integer k
 - ▶ Divide R into k partitions of size $\frac{P_R}{k}$ (k + 1 buffer pool pages needed)
 - ▶ This leaves B (k+1) extra buffer pool pages

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Query evaluation Join algorithms

How it works — partitioning and joining



Query evaluation Join algorithms

On predicates

- Essentially, *reduces* the *number* of *full passes*
- Running example, $P_R = 1,000$, $P_S = 500$, assume 300 pages in the buffer pool
- Choose the smaller relation, S

Savings over grace hash join

- Two partitions for it, each 250 pages
 - ▶ But one will stay in memory; so, cost is 500+250=750 I/Os
- Scan R, use two partitions, each 500 pages
 - ▶ But the *first one* is *not written* to disk: so cost is 1.000+500=1500 I/Os
- Join the two on-disk partitions, cost 250+500=750 I/Os
- Total cost 750+1500+750=300 I/Os
- At 10ms an I/O, this is half a minute

Query evaluation

Join algorithms

On pipelining

- Pipelining is great, but it cannot always be achieved
- All three algorithms will essentially block at some point
 - ▶ In the best case, between matches
 - ▶ In the worst case, until after a few scans of the input relations
- This is not necessarily bad; in fact, even if the algorithms block, the time needed to compute the complete join result might be less
- In reality, more than two stages of pipelining can rarely be obtained in a single plan

Query evaluation Join algorithms

- ► If there are no equi-join predicates (inequality joins) the only algorithm that will work is *nested loops* (why?)
- ▶ If there are indexes on the inequality join predicate's attributes, we can use index nested loops and revert the join to multiple scans
 - * Hoping that we will have buffer pool hits
 - * Remember access patterns and page replacement policy?
- ► Luckily, in a typical query workload there will mostly be equi-join predicates

Query evaluation Join algorithms

Summary

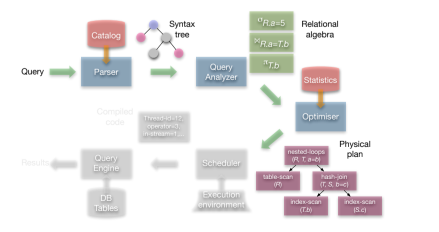
- The physical join is the most optimised physical evaluation operator
 - ▶ Because a DBMS spends most of its time evaluating joins
- Three main classes of algorithms
 - ► Iteration-based, order-based, partition-based
- Three main choice criteria
 - ► Physical layout, indexes, available memory

Summary (cont.)

- *Iteration-based* methods
 - ► Essentially, *nested loops*
 - ▶ Very simple to implement, but if implemented poorly very inefficient
 - ▶ But also *very useful* because they *evaluate non-equi-join predicates*
- Order-based methods
 - ► *Sort* the inputs, *merge* them afterwards
 - ▶ Well-behaved cost 3-4 passes over the data will do the trick

Query optimisation Overview

Query cycle



Summary (cont.)

- Partition-based methods
 - ► Simple hash join, Grace hash join, and hybrid hash join
 - ▶ If there is extra memory, hybrid hash join's behaviour is excellent
- Figuring out the best join algorithm for a particular pair of inputs is the job of the query optimiser
- Which, along with good implementations, will choose the one that evaluates a join in 30 seconds and not in 14,000 hours

Query optimisation Overview

Query optimiser

- The query optimiser is the heart of the evaluation engine
 - ▶ Yes, the *physical operators* get the *job done*
 - ▶ Yes, the execution model makes sure the operators actually run
 - ▶ But, unless the *query optimiser decides on those things*, the query will
 - ► And the *decision* needs to be a *good one*

Query optimisation Overview

Query optimisation Overview

Decisions

- Two crucial decisions the optimiser makes
 - ► The *order* in which the *physical operators* are applied on the inputs (i.e., the plan employed)
 - ► The algorithms that implement the physical operators
- These two decisions are not independent
 - ▶ In fact, one affects the other in more ways than one

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Query optimisation Overview

Plan enumeration

- Plan enumeration consists of two parts (again, not necessarily independent from one another)
 - ► Access method selection (i.e., what is the best way to access a relation that appears in the query?)
 - ▶ Join enumeration (i.e., what is the best algorithm to join two relations, and when should we apply it?)
- Access methods, join algorithms and their various combinations define a search space
 - ► The *search space* can be *huge*
 - ▶ Plan enumeration is the exploration of this search space

Cost-based query optimisation

- The paradigm employed is cost-based query optimisation
 - ► Simply put: enumerate alternative plans, estimate the cost of each plan, pick the plan with the minimum cost
- For cost-based optimisation, we need a cost model
 - ► Since what "hurts" performance is I/O, the cost model should use I/O as its basis
 - ► Hence, the cardinality-based cost model
 - ★ Cardinality is the number of tuples in a relation

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Query optimisation Overview

Search space exploration

- As was stated, the *search space* is *huge*
 - **Exhaustive exploration** is out of the question
 - ▶ Because it *could be the case* that *exploring* the search space might *take* longer than actually evaluating the query
 - ► The way in which we explore the search space describes a query optimisation method
 - ★ Dynamic programming, rule-based optimisation, randomised exploration, ...

Query optimisation Overview

Query optimisation Overview

Just an idea ...

- A query over five relations, only one access method, only one join algorithm, only left-deep plans
 - ▶ Remember, $cost(R \bowtie S) \neq cost(S \bowtie R)$
 - ▶ So, the number of *possible plans* is 5! = 120
 - ▶ If we add one extra access method, the number of possible plans becomes $2^5 \cdot 5! = 3840$
 - ▶ If we add one extra join algorithm, the number of possible plans becomes $2^4 \cdot 2^5 \cdot 5! = 61440$

Query optimisation Overview

Cardinality estimation

- An entire area of query optimisation
- Largely a matter of statistics
- It has *triggered* the "percentage wars"
 - "This estimation technique is within x% of the true value with a y%probability"
- Fact: the better the statistics, the better the decisions
- Another fact: errors in statistics propagate exponentially; after 4 or 5 joins, you might as well flip a coin
- Third fact: cost functions are discontinuous, so in certain scenarios only perfect statistics will help

Cardinality-based cost model

- A cardinality-based cost model means we need good ways of doing the following
 - ▶ Using cardinalities to estimate costs (e.g., accurate cost functions)
 - ► Estimating output cardinalities after we apply certain operations (e.g., after a selection the cardinality will change: it will not change after a projection)
 - * Because these output cardinalities will be used as inputs to the cost functions of other operations

Query optimisation Overview

Are we done?

- The previous issues were only a subset of the problems an optimiser solves
 - ► We also need to worry about certain properties of the data
 - ★ For instance, if we use a B+tree as an access method, then we won't have to sort (e.g., interesting orders in System R)
 - * If we use a hash join later on the order is spoiled
 - * So we will have to sort again
 - ▶ Depending on the algorithm and the environment, we need to allocate
- And as if all these were not enough, optimisation time assumptions do not necessarily hold at run time

Query optimisation Overview

Query optimisation Overview

The final nail ...

- These are all for one query
- Now, imagine a system doing that for 1000 queries
 - ► Simultaneously
- And it all has to be done fast
 - ▶ Once a decision is made, it cannot be undone

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Query optimisation Overview

The agenda

- Mapping SQL queries to relational algebra
 - ▶ Query blocks, uncorrelated *vs.* correlated queries
- Optimisation of a single query block
- Equivalence rules
- Statistics and cardinality estimation
- Search space exploration
 - ► Dynamic programming (System-R)

Conclusion

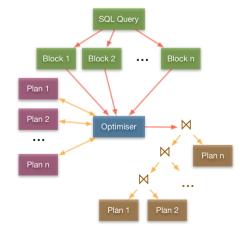
- Query optimisation is a very, very hard problem
- But without it a DBMS is doomed to seriously sub-optimal performance
- The problem is not nearly solved
 - ► All we *have* is *decent* optimisation strategies
 - ► And decent sub-problem solutions
- Fact: rarely will an optimiser pick the "best" plan
 - ▶ But it will almost always pick a plan with good performance and stay away from bad choices
 - ▶ At the end of the day, thats what counts

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Query optimisation Block decomposition and equivalence rules

SQL decomposition

- SQL queries are optimised by *decomposing* them into a *collection* of *query* blocks
- A block is optimised in isolation, resulting in a plan for a block
- Plans for blocks are combined to form the complete plan for the query



What is a block?

- An SQL query with no nesting
- Exactly one select-clause
- Exactly one from-clause
- At most one
 - ▶ Where-clause in conjunctive normal form
 - ► *Group by-/sort by-*clause
 - Having-clause

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Query optimisation

Block decomposition and equivalence rules

Two blocks in the query

```
select s.sid, min(r.day)
                    sailors s, reserves r, boats b
                   s.sid = r.sid and r.bid = b.bid and
                    b.color = 'red' and
                    s.rating = ()
           group by s.sid
           having count(*) > 1
                                           reference
        outer
                                           select max(s2.rating)
        block
                                           from sailors s2
select s.sid, min(r.day)
        sailors s, reserves r, boats b
                                                  nested
       s.sid = r.sid and r.bid = b.bid and
where
                                                  block
        b.color = 'red' and
        s.rating = (select max(s2.rating))
                    from sailors s2)
group by s.sid
having count(*) > 1
```

Example

Sample schema

- Sailors (sid, sname, rating, age)
- Boats (bid, bname, color
- Reserves (sid, bid, day, rname)

Example

For each sailor with the highest rating over all sailors, and at least two reservations for red boats, find the sailor id and the earliest date on which the sailor has a reservation for a red boat.

SQL query

```
select
           s.sid, min(r.day)
from
           sailors s, reserves r, boats b
           s.sid = r.sid and r.bid = b.bid and
           b.color = 'red' and
           s.rating = (select max(s2.rating) from sailors s2)
group by
           s.sid
           count(*) > 1
having
```

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Query optimisation Block decomposition and equivalence rules

Single block optimisation — step 1

```
SQL query
 select
            s.sid, min(r.day)
 from
             sailors s, reserves r, boats b
             s.sid = r.sid and r.bid = b.bid and
            b.color = 'red' and
             s.rating = (select max(s2.rating) from sailors s2)
 group by
            s.sid
            count(*) > 1
 having
```

Relational algebra

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```
\pi_{s.sid,\min(r.day)}
having_{count(*)>2}
group by_{s,sid}
\sigma_{s.sid=r.sid \land r.bid=b.bid \land b.color=red \land s.rating=nested-value}
sailors \times reserves \times boats))))
```

- Express the query in relational algebra
- More specifically, extended relational algebra

Single block optimisation — step 2

Relational algebra — before

 $\pi_{s.sid,min(r.day)}$ ($having_{count(*)>2}($ group bys.sid($\sigma_{s.sid=r.sid \land r.bid=b.bid \land b.color=red \land s.rating=nested-value}$ $sailors \times reserves \times boats))))$

Relational algebra — after

 $\pi_{s.sid}($ $\sigma_{s.sid=r.sid \land r.bid=b.bid \land b.color=red \land s.rating=nested-value}$ $sailors \times reserves \times boats)$

- *Ignore* the aggregate operations
 - ► They only have meaning for the complete result
 - ► Convert the *query* into a subset of relational algebra called $\sigma\pi\times$

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Query optimisation Block decomposition and equivalence rules

Equivalence rules

- Essentially, every query block consists of three things
 - ► Cartesian product of all relations in the from-clause
 - ► Selection predicates of the where-clause
 - ▶ *Projections* of the *select-clause*
- The equivalence rules define the space of alternative plans considered by an optimiser
 - ▶ In other words, the *search space of a query*

Single block optimisation — step 3

- Use equivalence rules to identify alternative ways of formulating the query
- "Plug in" algorithms
- Enumerate plans
- Estimate the cost of each plan
- Pick the one with the minimum cost

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Query optimisation Block decomposition and equivalence rules

Selection and projections

- Cascading of selections
- Commutativity

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- Cascading of projections

 - ▶ iff $a_i \subseteq a_{i+1}$, i = 1, 2, ..., n-1

Cartesian products and joins

- Commutativity
 - $ightharpoonup R \times S = S \times R$
 - $ightharpoonup R \bowtie S \equiv S \bowtie R$
- Assosiativity
 - $ightharpoonup R imes (S imes T) \equiv (R imes S) imes T$
 - $ightharpoonup R \bowtie (S \bowtie T) \equiv (R \bowtie S) \bowtie T$
- Their combination
 - $ightharpoonup R \bowtie (S \bowtie T) \equiv R \bowtie (T \bowtie S) \equiv (R \bowtie T) \bowtie S$ $\equiv (T \bowtie R) \bowtie S$

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Query optimisation Block decomposition and equivalence rules

Among operations (cont.)

- Projection-Cartesian product commutativity
 - $\blacktriangleright \pi_a (R \times S) \equiv \pi_{a_1}(R) \times \pi_{a_2}(R)$
 - iff a_1 is the subset of attributes in a appearing in R and a_2 is the subset of attributes in a appearing in S
- Projection-join commutativity
 - $\blacktriangleright \pi_a (R \bowtie_c S) \equiv \pi_{a_1}(R) \bowtie_c \pi_{a_2}(R)$
 - iff same as before and every attribute in c appears in a
- Attribute elimination

 - iff at subset of attributes in R appearing in either a or c and as is the subset of attributes in S appearing in either a or c

Among operations

- Selection-projection commutativity
 - \bullet π_a (σ_c (R)) $\equiv \sigma_c$ (π_a (R))
 - iff every attribute in c is included in the set of attributes a
- Combination (join definition)
 - $\triangleright \sigma_c (R \times S) \equiv R \bowtie_c S$
- Selection-Cartesian/join commutativity
 - \bullet $\sigma_c(R \times S) \equiv \sigma_c(R) \bowtie S$
 - ▶ iff the attributes in c appear only in R and not in S
- Selection distribution/replacement
 - $\sigma_{c_1}(R) \bowtie \sigma_{c_2}(S)$
 - \blacktriangleright iff c_1 is relevant only to R and c_2 is relevant only to S

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Query optimisation Block decomposition and equivalence rules

What do we have and what do we need?

- We have
 - ► A way to decompose SQL queries into multiple query blocks
 - ▶ A way to map a block to relational algebra
 - ► Equivalence rules between different algebraic expressions, i.e., a search space
- We need
 - ▶ A way to estimate the cost of each alternative expression
 - ★ Depending on the algorithms used
 - ► A way to *explore* the *search space*

Cost estimation

- A *plan* is a tree of operators
- Two parts to estimating the cost of a plan
 - ► For each node, estimate the cost of performing the corresponding operation
 - ► For each node, estimate the size of the result and any properties it might have (e.g., sorted)
- Combine the estimates and produce an estimate for the entire plan



Selectivity factor

- The maximum number of tuples in the result of any query is the product of the cardinalities of the participating relations
- Every *predicate* in the *where-clause* eliminates some of these *potential* results
- Selectivity factor of a single predicate is the ratio of the expected result size to the maximum result size
- Total result size is estimated as the maximum size times the product of the *selectivity factors*
- Key assumption: the predicates are statistically independent

Cost and cardinality

- We have seen various storage methods and algorithms
 - ► And *know the cost* of *using each* one, *depending* on the *input*
- The problem is estimating the output cardinality of the operations
 - ► Namely, *selections* and *joins*

How it works

SQL query

select $a_1, a_2, \dots a_k$ $R_1, R_2, \ldots R_n$ where P_1 and P_2 and ... and P_m

Maximum output cardinality

 $|R_1| \cdot |R_2| \cdot \ldots \cdot |R_n|$

Selectivity factor product

 $f_{P_1} \cdot f_{P_2} \cdot \ldots \cdot f_{P_m}$

Estimated output cardinality

$$(f_{P_1} \cdot f_{P_2} \cdot \ldots \cdot f_{P_m}) \cdot |R_1| \cdot |R_2| \cdot \ldots \cdot |R_n|$$

Various selectivity factors

- $column = value \rightarrow \frac{1}{\#keys(column)}$
 - ► Assumes *uniform distribution* in the values
 - ▶ Is itself an approximation
- $column_1 = column_2 \rightarrow \frac{1}{\max(\#keys(column_1), \#keys(column_2))}$
 - ► Each value in column₁ has a matching value in column₂; given a value in column₁, the predicate is just a selection
 - ► Again, an approximation

Query optimisation

Cardinality estimation

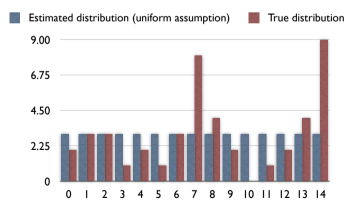
Key assumptions made

- The values across columns are uncorrelated
- The values in a single column follow a uniform distribution
- Both of these assumptions rarely hold
- The first assumption is hard to lift
 - ▶ Only recently have researchers started tackling the problem
- The uniform distribution assumption can be lifted with better statistical methods
 - ► In our case, *histograms*

Various selectivity factors (cont.)

- $column > value \rightarrow \frac{(high(column) value)}{(high(column) low(column))}$
- $value_1 < column < value_2 \rightarrow \frac{(value_2 value_1)}{(high(column) low(column))}$
- column in list \rightarrow number of items in list times s.f. of column = value
- column in sub-query → ratio of subquery's estimated size to the number of keys in column
- not (predicate) \rightarrow 1 (s.f. of predicate)
- $\bullet P_1 \lor P_2 \to f_{P_1} + f_{P_2} f_{P_1} \cdot f_{P_2}$

What we would like



Lifting the uniform distribution assumption

- At the basic level, all we need is a collection of (value, frequency) pairs
- Which is just a relation!
 - ► So, *scan* the *input* and *build* it
- But this is unacceptable
 - ▶ Because the *size* might be comparable to the size of the relation
 - ► And we *need* to *answer* queries about the value distribution fast

parts		
name	color	stock
bolt	red	10
bolt	green	5
nut	blue	4
nut	black	10
nut	red	5
nut	green	10
cam	blue	5
cam	green	10
cam	black	10

parts.color

value	freq		
red	2		
green	3		
blue	2		
black	2		
parts.stock			

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Desirable histogram properties

- Small
 - ▶ Typically, a DBMS will allocate a *single page* for a histogram!
- Accurate
 - ► Typically, less than 5% error
- Fast access
 - ► Single lookup access and simple algorithms

Histograms

- Elegant data structures to capture value distributions
 - ▶ Not affected by the uniform distribution assumption (though this is not entirely true)
- They offer *trade-offs* between *size* and *accuracy*
 - ▶ The *more memory* that is dedicated to a histogram, the *more accurate* it is
 - ▶ But also, the *more expensive* to manipulate
- Two basic classes: equi-width and equi-depth

Query optimisation

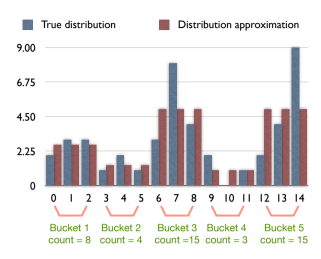
Cardinality estimation

Mathematical properties

- A histogram approximates the value distribution for attribute X of table T
- The value distribution is partitioned into a number of b subsets, called buckets
- There is a partitioning constraint that identifies how the partitioning takes place
 - ▶ Different constraints, lead to different classes of histograms
- The values and frequencies in each bucket are approximated in some common fashion

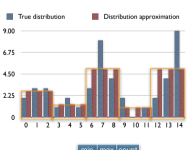
Query optimisation Cardinality estimation

Equi-width histogram



Equi-width histogram estimation

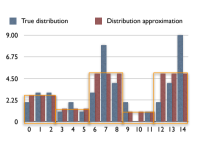
- To estimate the output cardinality of a range query
 - ► The *starting bucket* is identified
 - ► The *histogram* is then scanned forward until the ending bucket is identified
 - ► The *numbers of tuples* in the buckets of the range are summed
 - ▶ Within each bucket the uniform distribution assumption is made
- $6 \le v \le 10$: $\frac{3}{3} \cdot 15 + \frac{2}{3} \cdot 3 = 17$



min	max	count
0	2	8
3	5	4
6	8	15
9	11	3
12	14	15

Equi-width histogram construction

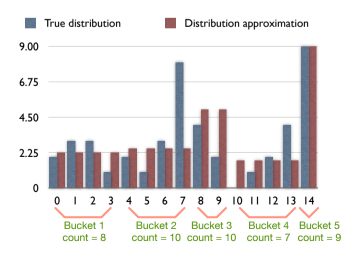
- The total range is divided into sub-ranges of equal width
- Each *sub-range* becomes a bucket
- The total number of tuples in each bucket is stored



min	max	count
0	2	8
3	5	4
6	8	15
9	11	3
12	14	15

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Equi-depth histogram



• Equi-depth histograms are generally better than equi-width

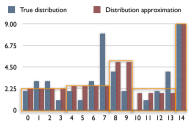
► Buckets with frequently occurring values contain fewer values

► Infrequently occurring values are approximated less accurately (but the

► So the uniform distribution assumption within each bucket leads to

Equi-depth histogram construction and estimation

- The total range is divided into sub-ranges so that the number of tuples in each range is (approximately) equal
- Each *sub-range* becomes a bucket
- The same schema as in equi-width histograms is used
- In fact, the same algorithm is used for *estimation* (!)
- 6 < v < 10: $\frac{2}{4} \cdot 10 + \frac{2}{3} \cdot 10 + \frac{1}{4} \cdot 7 \approx 17$



min	max	count
0	3	8
4	7	10
8	9	10
10	13	7
14	14	9

What do we have and what do we need?

- We have
 - ► A way to *decompose* a *query*
 - ▶ A way to *identify* equivalent, *alternative representations* of it (i.e., a search space)
 - ► A statistical framework to estimate cardinalities
 - ▶ A cost model to estimate the cost of an alternative
- We need
 - ► A way to *explore* the *search space*
 - Dynamic programming

min	max	cour
0	3	8
4	7	10
8		10
10	13	7
14	14	9

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Comparison

Query optimisation Plan enumeration

Dynamic programming

error is less significant)

better approximation

- In the beginning, there was System R, which had an optimiser
- System R's optimiser was using dynamic programming
 - ▶ An *efficient way* of *exploring* the search space
- Heuristics: use the equivalence rules to push down selections and projections, delay Cartesian products
 - ▶ Minimise input cardinality to, and memory requirements of the joins
- Constraints: left-deep plans, nested loops and sort-merge join only
 - ► Left-deep plans took better advantage of pipelining
 - ► Hash-join had not been developed back then

Interesting orders

- If there is an order by or group by clause on an attribute, we say that this attribute has an interesting order associated with it
 - ▶ Interesting, because depending on the access method we can get away with fewer physical operations (e.g., sorting)
- The same holds for attributes participating in a join
 - ▶ Again, interesting because we can use the access method in evaluating the join

Query optimisation Plan enumeration

An example

emp	^	^	
name	dno	job	salary
Smith	50	12	8500
Jones	50	5	15000
Doe	51	5	9500

	12	8500		50	MFG	Edinburgh
)	5	15000		51	Billing	London
	5	9500		52	Shipping	Glasgow
		_	nam	ne. salarv.	iob title. d	epartment nar

job	
job	title
5	clerk
6	typist
8	sales
12	mechanic

of employees who are clerks and work in departments in Edinburgh

select name, title, salary, dname from emp, dept, job where job.title='Clerk' and dept.location = 'Edinburgh' and emp.dno = dept.dno and emp.job = job.job

interesting orders

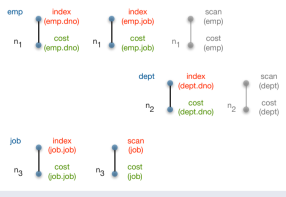
Dynamic programming steps

- Identify the cheapest way to access every single relation in the query. applying local predicates
 - ► For every relation, keep the cheapest access method overall and the cheapest access method for an interesting order
- For every access method, and for every join predicate, find the cheapest way to join in a second relation
 - ► For every join result keep the cheapest plan overall and the cheapest plan in an interesting order
- Join in the rest of the relations using the same principle

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Query optimisation Plan enumeration

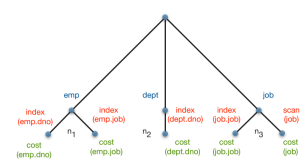
Access methods and local predicates



- Scanning emp is the most expensive method for emp; emp.dno and emp.job are interesting orders
- Scanning dept is the most expensive method for dept; dept, dno is an interesting order
- Scanning job is the cheapest method for job; but, job.job is an interesting order

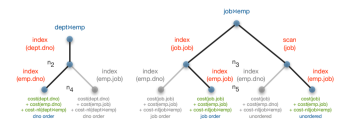
join predicates

Search tree for access methods



Query optimisation Plan enumeration

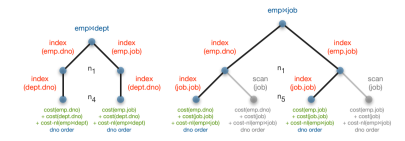
Join enumeration for relations dept, job (nested loops)



- $cost(emp \bowtie dept) \neq cost(dept \bowtie emp)$ so we will enumerate dept's joins even though we have an alternative for generating the same result (same for $job \bowtie emp$)
- Both dept ⋈ emp results in the same order, only one propagated
- Since there is no dept ⋈ job predicate in the query, that join is not enumerated (same for job ⋈ dept)
- The unordered result for job ⋈ emp is propagated because it is the cheapest overall

Query optimisation Plan enumeration

Join enumeration for relation emp (nested loops join)



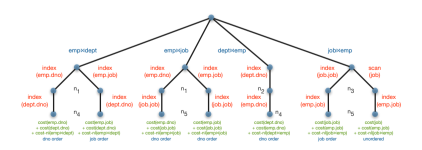
- Both emp ⋈ dept results are in different interesting orders so they are propagated
- Only the cheapest result in any interesting order is propagated for each pair of inputs

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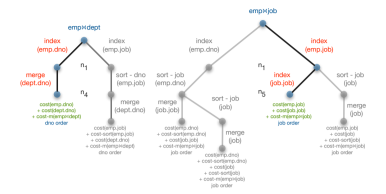
Query optimisation Plan enumeration

Search tree — 2 relations, nested loops join



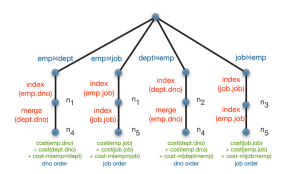
Join enumeration for relations dept, job (sort-merge)

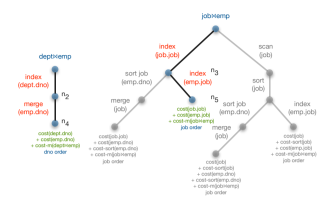




Query optimisation Plan enumeration

Search tree — 2 relations, sort-merge join

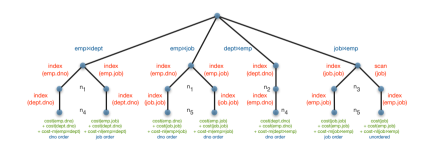




Query optimisation Plan enumeration

Query optimisation Plan enumeration

Search tree — 2 relations, both join methods



- For each pair or relations, for each different join order and for each interesting order for that pair one plan is propagated
- An unordered result is only propagated if it is the cheapest overall for a pair in a given join order

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Three relations

- Repeat the process
 - ► For *every pair* of two relations
 - ► For *every join* method
 - ► For every access method of the remaining relation
 - Find the cheapest way to join the third relation with the pair
 - **★** Estimate cardinalities
 - * Estimate the cost of computing the join
 - ▶ Keep the cheapest choice for every interesting order and the cheapest for the unordered case if it is the cheapest overall

Query optimisation

Plan enumeration

Randomised exploration

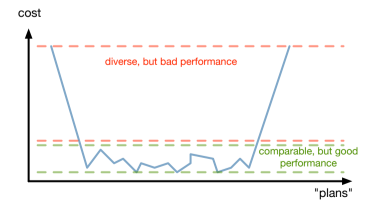
- Mostly useful in big queries (more than 15 joins or so)
- The *problem* is one of *exploring a bigger portion* of the search space
 - ▶ So, every once in a while the optimiser "jumps" to some other part of the search space with some probability
- As a *consequence*, it gets to *explore parts* of the search space it would not have explored otherwise

Rule-based optimisation

- Basically an issue of if-then rules
 - ► If (condition list) then apply some transformation to the plan constructed so far
 - * Estimate the cost of the new plan, keep it only if it is cheaper than the original
 - ► The *order* in which the *rules are applied* is *significant*
 - ► As a *consequence*, rules are applied *by precedence*
 - ★ For instance, *pushing down selections* is given *high precedence*
 - * Combining two relations with a Cartesian product is given low precedence

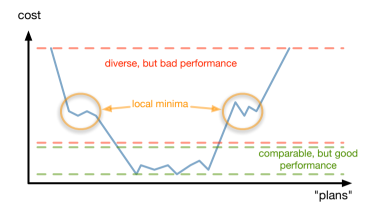
Query optimisation Plan enumeration

The "well"



Query optimisation Plan enumeration

The "well" and local minima

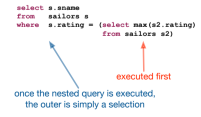


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Query optimisation Plan enumeration

Uncorrelated queries

- Usually, they can be executed in isolation
- The *nested query* feeds the outer query with results



Final step — the entire plan

- The optimiser has produced plans for each query block
- The *question* is now one of *combining* the *sub-plans* to *formulate* the entire query plan
- The strategy used depends on whether the outer and nested queries are correlated or not
 - ▶ If they are, then in all probability the two sub-plans will be combined through a join

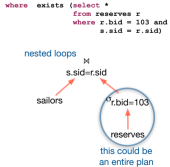
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Query optimisation Plan enumeration

select s.sname from sailors s

Correlated queries

- Sometimes, it is not possible to execute the nested query just once
- In those cases the optimiser reverts to a nested loops approach
 - ► The *nested query* is executed once for every tuple of the outer query



Query optimisation Plan enumeration

In practice

- Before breaking up the query into blocks, most systems try to rewrite the *query* in some *other way* (*de-correlation*)
 - ▶ The idea is that there will probably be a join, so it will be better if the query is optimised in its entirety
- If de-correlation is not possible, then it is nested loops all the way
 - ▶ Usually, *compute* the *nested guery*, *store* it in a temporary relation and do nested loops with the outer

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Query optimisation Summary

Summary

- The *query optimiser* is the *heart* of the *query engine*
 - ▶ If it does *not* do a *good job*, the engine is doomed to *sub-optimal* performance
- Two key, closely related decisions
 - Order in which operations are performed
 - ► Algorithms that perform the operations
- The paradigm used is cost-based optimisation
 - ▶ Three steps: generate alternative plans, estimate the cost of each plan, pick the cheapest
- The cost model used is the cardinality-based cost model
 - ▶ Because *cardinality* is a *good I/O metric*
 - ► As a *consequence*, we need *good ways of doing* two things
 - * Estimating the cost of an algorithm
 - ★ Estimating the output cardinality of operations

What do we have and what do we need?

We have

- ► A way to decompose a guery
- ► A way to *identify* equivalent, *alternative representations* of it (i.e., a search space)
- ► A statistical framework to estimate cardinalities
- ▶ A cost model to estimate the cost of an alternative
- ▶ Ways of *exploring* the *search space*
- We need
 - ► Nothing!

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Query optimisation Summary

Summary (cont.)

- Cardinality estimation is 50% of the problem
 - ► Two approaches: uniform distribution assumption, or histograms
 - ► The uniform distribution assumption essentially does not "care" about the values themselves, they all have an equal probability of appearing
 - ► Histograms are a better and more elegant distribution approximation technique
 - ★ Equi-width and equi-depth histograms are the two dominant classes

Query optimisation Summary

Summary (cont.)

- The remaining 50% is search space exploration
 - ► Largely based on the equivalence rules of relational algebra
 - ► Dynamic programming is the dominant approach
 - ★ Find the *cheapest way* to access single relations
 - * Find the *cheapest way* to join two relations
 - * For each pair, find the cheapest way to join in a third relation
 - ★ Keep going ...

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Transactions, concurrency, and recovery Overview

Overview

- So far, we have focussed on query processing
 - ▶ In other words, reading and manipulating data
- A database system, however, not only reads, but also stores data
 - ► At the same time as others are querying it
- We need a way to ensure concurrent access to the data
 - ▶ Without compromising system performance

Query optimisation Summary

Summary (cont.)

- Other approaches include rule-based optimisation, randomised exploration, ...
- All approaches aim at one thing
 - ► Picking a good evaluation plan
 - ▶ It might not be the cheapest overall, but it usually is of comparable cost
- Query optimisation is still an open issue
 - ▶ We have good ways of solving sub-problems, but the entire problem remains largely unsolved

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Transactions, concurrency, and recovery Overview

Overview (cont.)

- The basic concept is transaction processing
- Every transaction needs to satisfy four basic properties
 - ► Atomicity, consistency, isolation, durability
- How does the system guarantee these properties?
 - ▶ Remember, without compromising performance
 - ► Solution: by interleaving transactions

Transactions, concurrency, and recovery Overview

Transactions, concurrency, and recovery Overview

Overview (cont.)

- How can we decide if, after we have interleaved transactions, the result is correct?
 - ► Interleaving transactions actually causes certain anomalies
 - ► Solution: the system uses locks to ensure correctness
- How are locks used?
 - ► Lock granularity, degrees of consistency and two-phase locking
- What *impact* do *locks* have on *performance*?

Transactions

- A DBMS spends a lot of time waiting on I/O
 - ▶ It is important to keep the CPU busy while waiting
 - ▶ In other words, execute other operations concurrently
- Fact: the DBMS does not "care" what the user does with the data that is being read or written
 - ▶ All it cares about is that data is being read or written
- A transaction is the DBMS's abstract view of user programs: a sequence of reads and writes

Overview (cont.)

- Locking poses significant overhead
 - Luckily, however, this overhead can be "tuned" by the user
 - ► Transaction isolation levels
- But what if the worse comes to worst?
 - ► System crashes
 - ► Transactional semantics and recovery
 - ▶ Write-ahead logging and the ARIES algorithms

Concurrent execution

- The transaction user abstraction: when a user submits a transaction it is as if the transaction is executing by itself
 - ► The *DBMS* achieves concurrency by interleaving transactions
 - ▶ If the transaction begins with the DB in a consistent state, it must leave the DB in a consistent state after it finishes
- The *semantics* of the *transactions* are *unknown* to the *system*
 - ▶ Whether the transaction updates a bank account or it fires a rocket missile, the DBMS will never know!

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ACID properties

- Atomicity: all the actions in a transaction are executed as a single atomic operation; either they are all carried out or none are
- Consistency: if a transaction begins with the DB in a consistent state. it must finish with the DB in a consistent state
- Isolation: a transaction should execute as if it is the only one executing; it is protected (isolated) from the effects of concurrently running transactions
- Durability: if a transaction has been successfully completed, its effects should be *permanent*

Example

- Consider two transactions
 - ► First transaction transfers funds, second transaction pays 6% interest
- If they are submitted at the same time, there is no guarantee as to which is executed first
 - ▶ But the *end effect* should be equivalent to the transactions running serially

T1



T2



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Example (cont.)

Acceptable schedule

T1	A = A+100		B = B-100	
T2		A = 1.06*A		B = 1.06*B

Problematic schedule

T1	A = A+100			B = B-100
T2		A = 1.06*A	B = 1.06*B	

DBMS's view

T1	R(A), W(A)			R(B), W(B)
T2		R(A), W(A)	R(B), W(B)	

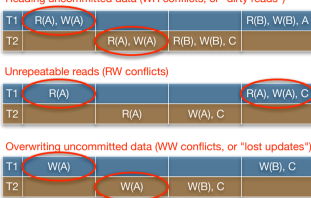
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Scheduling

- A schedule is a sequence of reads and writes for some transaction workload incorporating all actions of the workload's transactions
 - ▶ Serial schedule: the actions of different transactions are not interleaved
 - ► Equivalent schedules: for any database state, the effect of executing the first schedule is identical to the effect of executing the second schedule
 - ► Serialisable schedule: a schedule that is equivalent to a serial schedule

Conflicts

Reading uncommitted data (WR conflicts, or "dirty reads")



What can go wrong?

- If a transaction T_i is aborted, then all its actions have to be undone; not only that, but if T_i reads an object written by T_i , T_i needs to be aborted as well (cascading aborts)
- Most systems avoid cascading aborts with the following rule:
 - ▶ If T_i writes an object T_i can read this object only after T_i commits
- In order to know what needs to be undone, the system keeps a log, recording all writes
- The *log* is also *helpful* when *recovering* from *system crashes*

The solution: locks

- Before a transaction "touches" a DB object it has to obtain a lock for
 - ► S (Shared) lock for reading
 - ► X (eXclusive) lock for writing
- Strict two-phase locking (Strict 2PL)
 - ► Each transaction must obtain an S lock for everything it reads before it starts reading it and an X lock for everything it writes before it starts writing
 - ► All locks held by a transaction are released only when the transaction
 - ▶ Once a transaction obtains an X lock for a DB object no other transaction can obtain an X or an S lock for that object
- Strict 2PL produces only serialisable schedules

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Transactions, concurrency, and recovery Transaction processing

The log

- The following actions are recorded in the log
 - ▶ Whenever a transaction writes an object
 - * The log record must be on disk before the data record reaches the disk
 - ▶ Whenever a *transaction commits/aborts*
- Log records are chained by transaction ID (why?)
- All log-related activities (in fact, all concurrency control related activities) are handled by the DBMS
 - ► The user does not know anything

Transactions, concurrency, and recovery Transaction processing

Transactions, concurrency, and recovery Concurrency control

Crash recovery

- Three phases to recovery (ARIES)
 - ► Analysis: scan log forward, identifying committed and aborted/unfinished transactions
 - ▶ Redo: all committed transactions are made durable
 - ▶ Undo: the actions of all aborted and/or unfinished transactions are undone

Transactions, concurrency, and recovery Concurrency control

Dependency graphs

- Given a schedule S
 - ► One *node* per *transaction*
 - \blacktriangleright An edge from T_i to T_i , if T_i reads or writes an object written by T_i
- Theorem: a schedule S is conflict serialisable if and only if its dependency graph is acyclic

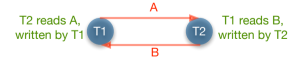
Concurrency control

- Serial schedule: the actions of different transactions are not interleaved
- Equivalent schedules: for any database state, the effect of executing the first schedule is identical to the effect of executing the second schedule
- Serialisable schedule: a schedule that is equivalent to a serial schedule
- Two schedules are conflict equivalent if:
 - ► They *involve* the *same actions* of the *same transactions*
 - Every pair of conflicting actions is ordered the same way
- Schedule S is conflict serialisable if S is conflict equivalent to some serial schedule

Transactions, concurrency, and recovery Concurrency control

Example: not conflict serialisable schedule

T1	R(A), W(A)			R(B), W(B)
T2		R(A), W(A)	R(B), W(B)	



Transactions, concurrency, and recovery Concurrency control

Review: Strict 2PL

- Strict two-phase locking (Strict 2PL)
 - ► Each *transaction* must obtain an *S* (*Shared*) lock for everything it reads before it starts reading it and an X (eXclusive) lock for everything it writes before it starts writing
 - ► All locks held by a transaction are released only when the transaction commits
 - ▶ Once a transaction obtains an X lock for a DB object no other transaction can obtain an X or an S lock for that object
- Strict 2PL produces only serialisable schedules
 - ▶ In other words: schedules with acyclic dependency graphs

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Transactions, concurrency, and recovery Concurrency control

Lock management

- Lock and unlock requests are handled by the lock manager that maintains a lock table
- Lock table entry:
 - ► Number of transactions currently holding a lock
 - ► Type of lock held (shared or exclusive)
 - ► Pointer to queue of lock requests
- Locking and unlocking have to be atomic operations
- Lock upgrade: transaction that holds a shared lock can be upgraded to hold an exclusive lock

Simple 2PL

- Two-phase locking (2PL)
 - ► Each transaction must obtain an S (Shared) lock for everything it reads before it starts reading it and an X (eXclusive) lock for everything it writes before it starts writing
 - ► A transaction cannot request additional locks once it releases any locks
 - ▶ Once a transaction obtains an X lock for a DB object no other transaction can obtain an X or an S lock for that object

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Transactions, concurrency, and recovery Concurrency control

Deadlocks

- As always, where there are locks, there are deadlocks
- Deadlocks: cycle of transactions waiting for locks to be released by each other
- Two ways of dealing with deadlocks
 - ► Deadlock prevention
 - ► Deadlock *detection*

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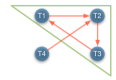
Deadlock prevention

- The solution involves timestamps; a timestamp is the transaction's priority
- If T_i wants a lock that T_i holds, there are two possible policies
 - \blacktriangleright Wait-Die: if T_i has higher priority, T_i waits for T_i ; otherwise T_i aborts
 - \blacktriangleright Wound-Wait: if T_i has higher priority, T_i aborts; otherwise T_i waits
- If a transaction re-starts, it has its original timestamp

Deadlock detection

- Create a waits-for graph
 - ► Nodes are transactions
 - ► There is an *edge* from T_i to T_i if T_i is waiting for T_i to release a lock
- Periodically check for cycles in the waits-for graph

T1	S(A)	R(A)			S(B)					
T2			X(B)	W(B)				X(C)		
ТЗ						S(C)	R(C)			X(A)
T4									X(B)	

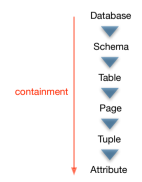


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Multiple granularity locks

- What should we lock? Tuples, pages, tables, ...
- But there is an *implicit* containment
- Idea: lock DB objects hierarchically



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Hierarchical locks and new locking modes

- Allow transactions to lock at each level of the hierarchy
- Introduce "intention" locks: IS and IX
 - ▶ Before locking an item, a transaction must introduce intention locks on all the item's ancestors in the hierarchy
 - ► Release locks in reverse order

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• One extra lock: SIX — "share, with intention to write"

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Compatibility matrix

held lock SIX

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A few examples

- T1 scans R, and updates a few tuples
 - ► T1 gets an SIX lock on R, then repeatedly gets an S lock on tuples of R, and occasionally upgrades to X on the tuples
- T2 uses an index to read only part of R
 - ► T2 gets an IS lock on R, and repeatedly gets an S lock on tuples of R
- T3 reads all of R
 - ► T3 gets an S lock on the entire relation
 - ► Or, it gets an IS lock on R, escalating to S lock on every tuple

In more detail

- Each transaction starts from the root of the hierarchy
- To obtain S or IS lock on a node, must hold IS or IX on parent node
 - ▶ What if a transaction holds SIX on parent? S on parent?
- To obtain X or IX or SIX on a node, must hold IX or SIX on parent node
- Must release locks in bottom-up order

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Here's the catch (the phantom problem)

- If we relax the assumption that the DB is a fixed collection of objects, even Strict 2PL will not assure serialisability!
 - ► T1 locks all pages containing sailor records with rating = 1, and finds oldest sailor (say, age = 71)
 - \blacktriangleright Next, T2 inserts a new sailor. rating = 1, age = 96
 - ► T2 also deletes oldest sailor with rating = 2 (and, say, age=80), and
 - ightharpoonup T1 now locks all pages containing sailor records with rating = 2, and finds oldest (say, age=63)
- No lock conflicts, but also no consistent DB state where T1 is "correct"!

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The problem

- T1 implicitly assumes that it has locked the set of all sailor records with rating = 1
 - ▶ The assumption only holds if no sailor records are added while T1 is executing!
 - ▶ We *need* some *mechanism* to *enforce* this *assumption*
 - **★** Index locking
 - * Predicate locking
- The example shows that conflict serialisability guarantees serialisability only if the set of objects is fixed!

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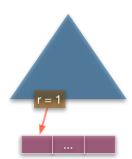
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Predicate locking

- Grant lock on all records that satisfy some logical predicate, e.g., $salary > 2 \cdot salary$
 - ► Index locking is a special case of predicate locking for which an index supports efficient implementation of the predicate lock
 - ▶ What is the *predicate* in the *sailor example*?
- In general, predicate locking imposes a lot of locking overhead

Index locking

- If there is an index on the rating field, T1 should lock the index page containing the data entries with rating = 1
 - ► If there are no records with rating = 1, T1 must lock the index page where such a data entry would be, if it existed!
- If there is no suitable index, T1 must lock all pages, and lock the file/table to prevent new pages from being added, to ensure that no new records with rating = 1 are added



Transactions, concurrency, and recovery Concurrency control

B+tree locking

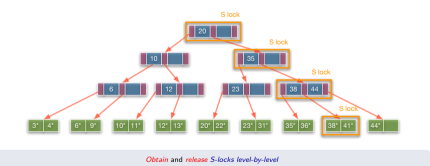
- How can we efficiently lock a particular node?
 - ► This is *entirely different* than *multiple granularity locking* (why?)
- One solution: ignore the tree structure, just lock pages while traversing the tree, following 2PL
 - ► Terrible performance
 - ▶ Root node (and many higher level nodes) become bottlenecks because every tree access begins at the root

Key observations

- Higher levels of the tree only direct searches to leaf pages
- For *insertions*, a *node* on a *path* from the *root* to a modified *leaf* must be locked (in X mode, of course), only if a split can propagate up to it from the modified leaf (similar point holds for deletions)
- We can *exploit* these *observations* to design *efficient locking protocols* that guarantee serialisability even though they violate 2PL

Transactions, concurrency, and recovery Concurrency control

Example: search 38*

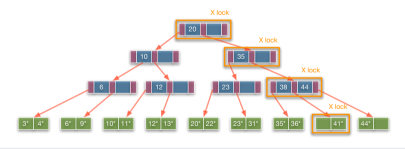


The basic algorithm

- Search: start at root and descend; repeatedly, S lock child then unlock parent
- Insert/Delete: start at root and descend, obtaining X locks as needed; once *child* is *locked*, *check* if it is *safe*:
 - ► Safe node: a node such that changes will not propagate up beyond this
 - * Insertion: node is not full
 - ★ Deletion: node is not half-empty
 - ▶ If *child* is *safe*, *release* all *locks* on *ancestors*

Transactions, concurrency, and recovery Concurrency control

Example: delete 38*

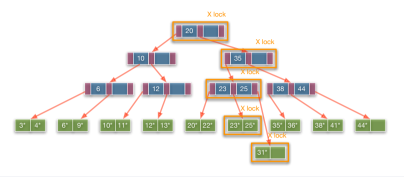


Obtain X-locks while descending; release them top-down once the node is designated safe

Transactions, concurrency, and recovery

Concurrency control

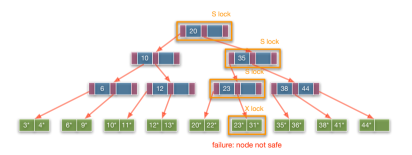
Example: insert 25*



Obtain X-locks while descending; leaf-node is not safe so create a new one and lock it in X-mode; first release locks on leaves

Transactions, concurrency, and recovery Concurrency control

Example: insert 25*



Obtain S-locks while descending, and X-lock at leaf; the leaf is not safe, so abort, release all locks and restart using the previous

Optimistic B+tree locking

- Search: as before
- Insert/delete: set locks as if for search, get to the leaf, and set X lock on the leaf
 - ▶ If the leaf is not safe, release all locks, and restart transaction, using previous insert/delete protocol
- "Gambles" that only leaf node will be modified; if not, S locks set on the first pass to leaf are wasteful
 - ▶ *In practice*, *better* than previous algorithm

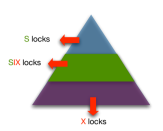
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Even better algorithm

- Search: as before
- Insert/delete: use original insert/delete protocol, but set IX locks instead of X locks at all nodes
 - ▶ Once leaf is locked, convert all IX locks to X locks top-down: i.e., starting from the unsafe node nearest to root
 - ► Top-down reduces chances of deadlock
 - * Remember, this is *not the same* as *multiple granularity locking!*

Transactions, concurrency, and recovery Concurrency control

Hybrid approach



- The *likelihood* that we will need an X lock decreases as we move up the tree
- Set S locks at high levels, SIX locks at middle levels, X locks at low levels

Transactions, concurrency, and recovery

Review: ACID properties

- Atomicity: all the actions in a transaction are executed as a single atomic operation; either they are all carried out or none are
- Consistency: if a transaction begins with the DB in a consistent state, it must finish with the DB in a consistent state
- Isolation: a transaction should execute as if it is the only one executing; it is protected (isolated) from the effects of concurrently running transactions
- Durability: if a transaction has been successfully completed, its effects should be *permanent*

Atomicity and durability are ensured by the recovery algorithms

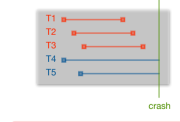
Transaction isolation

Isolation level	Dirty read	Unrepeatable read	Phantoms	
Read uncommitted	Maybe	Maybe	Maybe	
Read committed	No	Maybe	Maybe	
Repeatable reads	No	No	Maybe	
Serialisable	No	No	No	

Transactions, concurrency, and recovery Recovery

What can go wrong?

- Atomicity
 - ► Transactions may abort: their effects need to be undone
- Durability
 - ▶ What if the *system* stops running?



Transactional semantics

- T1, T2, T3 should be durable
- T4, T5 should be aborted

Problem statement

- *Updates* are happening in place
 - ► There is a *buffer pool*
 - * Data pages are read from disk
 - ★ Data pages are modified in memory
 - ★ Overwritten on, or deleted from disk
- We need a simple scheme to guarantee atomicity and durability

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The problems

- Steal's problems are all about atomicity
 - ▶ What if a *transaction modifying a page aborts*?
 - ▶ If we steal a page, we need to remember its old value so it can be restored (UNDO)
- No force's problems are all about durability
 - ▶ What if a system crashes before a modified page is written to disk?
 - ▶ We need to record enough information to make the changes permanent (REDO)

More on the buffer pool

- Two issues: force and steal
- Force: when a data page is modified it is written straight to disk
 - ► Poor response time
 - ► But durable
- Steal: effects of uncommitted transactions reach the disk
 - ► Higher throughput
 - ► But not atomic



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Transactions, concurrency, and recovery Recovery

The solution: logging

- Record REDO and UNDO information in a record of a separate structure: the log
 - ► Sequential writes for every update
 - ▶ *Minimal information* written (more efficient!)
 - Keep it on a separate disk!
- Log: a list of REDO and UNDO actions
 - ► Each *log record* contains *at least*:
 - * Transaction id, modified page, old data, new data

Write-ahead logging

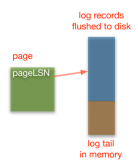
- The log adheres to the write-ahead protocol (WAL)
 - Must force the log record for an update before the corresponding data page gets to disk
 - 2 Must force all log records for a transaction before it commits
- #1 guarantees atomicity
- #2 guarantees durability

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Transactions, concurrency, and recovery Recovery

WAL and the log

- Each log record has a unique log sequence number (LSN)
 - ► LSNs are always increasing
- Each data page contains a pageLSN
 - ► The *LSN* of the *most recent* log record for an update to that page
- The system keeps track of flushedI SN
 - ► The max LSN flushed so far
- WAL: before a page is written, pageLSN ≤ flushedLSN



Normal execution

- Series of reads and writes
- Followed by a commit (success) or abort (failure)
- Steal, No-force management
- Adherence to the WAL protocol
- Checkpoints: periodically, the system creates a checkpoint to minimise the time taken to recover
 - ► Assume the DB is consistent after a checkpoint

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Transactions, concurrency, and recovery Recovery

Log records

- Possible log records types
 - ► Update
 - ► Commit
 - ► Abort
 - ► End (signifies commit or abort!)
 - Compensation Log Records
 - ★ Logging UNDO actions!
 - ★ But we will not talk about them in more detail



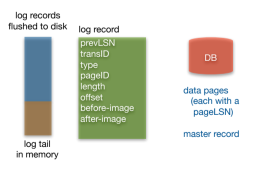
Other log-related state

- Transaction table: one entry per active transaction
 - ► Contains *transaction id*, *status* (running/committed/aborted) and lastLSN — log sequence number of the last log record for that transaction
- Dirty page table: one entry per dirty page in buffer pool
 - ► Contains recLSN the LSN of the log record which first caused the page to be dirty

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Transactions, concurrency, and recovery Recovery

What's stored where



Checkpoint records

- begin_checkpoint record: indicates when checkpoint began
- end_checkpoint record: contains current transaction table and dirty page table
- This is a "fuzzy checkpoint"
 - ▶ Other transactions continue to run; so these tables accurate only as of the *time* of the begin_checkpoint *record*
 - ▶ No attempt to force dirty pages to disk; effectiveness of checkpoint limited by oldest unwritten change to a dirty page
 - ► So it's a good idea to periodically flush dirty pages to disk
- Store LSN of checkpoint record in a safe place (master record)

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Transactions, concurrency, and recovery Recovery

Simple transaction abort

- For now, consider an *explicit abort* of a transaction
 - ► No crash involved

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- We want to "play back" the log in reverse order, UNDO ing updates
 - ► Get *lastLSN* of *transaction* from *transaction table*
 - ► Follow chain of log records backward via the prevLSN field
 - ► Before starting UNDO, write an Abort log record
 - ★ For recovering from crash during UNDO!

transaction table

dirty page table

lastLSN

status

recLSN flushedLSN Transactions, concurrency, and recovery Recovery

Transactions, concurrency, and recovery Recovery

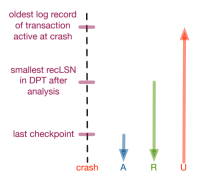
Abort (cont.)

- To perform UNDO, must have a lock on data
 - ► No problem
- Before restoring old value of a page, write a CLR
 - ► Continue logging while you UNDO!
 - ► CLR has one extra field: undonextLSN
 - ★ Points to the next LSN to undo (i.e., the prevLSN of the record we're currently undoing)
 - ► CLRs are never undone (but they might be redone when repeating history: guarantees atomicity)
- At the end of UNDO, write an "end" log record

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Transactions, concurrency, and recovery Recovery

Recovery: big picture



- Start from a checkpoint (found via master record)
- Three phases
 - ► Analysis: figure out which transactions committed since the checkpoint, and which failed
 - ► *REDO* all actions
 - ★ Repeat history
 - ► UNDO effects of failed transactions

Transaction commit

- Write commit record to log
- All log records up to the transaction's lastLSN are flushed
 - ► Guarantees that flushedLSN > lastLSN
 - ▶ Note that *log flushes* are *sequential*, *synchronous writes* to disk
 - ► Many log records per log page
- Commit() returns
- Write end record to log

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Transactions, concurrency, and recovery Recovery

Additional issues

- What happens if the system crashes during the analysis phase? During REDO phase?
- How can the amount of work during REDO be limited?
 - ► Flush asynchronously in the background
- How can the *amount of work* during *UNDO* be *limited*?
 - ► Avoid long-running transactions

Transactions, concurrency, and recovery Summary

Transactions, concurrency, and recovery Summary

Summary

- Concurrency control and recovery are key concepts of a DBMS
- Both are ensured by the system itself, the user does not (and should not!) know of their existence
- The key abstraction is the transaction
 - ► The *processing unit* of the *system*
 - ► Four key properties
 - ★ Atomicity, consistency, isolation, durability

Transactions, concurrency, and recovery Summary

Summary (cont.)

- Multiple granularity of locks
 - ▶ Leads to an escalation of locks, as we are descending the hierarchy
- Special protocols for indexes and predicates
- Transactions help after recovering from a crash
 - ▶ As the processing unit, we know what needs to be repeated or deleted

Summary (cont.)

- A transaction is viewed by the system as a series of reads and writes
- To improve throughput, the system interleaves the actions of the transactions (i.e., a schedule)
 - ► At all times, *ensuring serialisability* of the *produced schedules*
- Locks are the mechanism that ensures serialisability
 - ▶ Before reading, obtain a Shared lock
 - ▶ Before writing, obtain an eXclusive lock

Transactions, concurrency, and recovery Summary

Summary (cont.)

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- Steal, no-force buffer pool management
 - ► Higher response time (steal)
 - ► Higher throughput (no-force)
- Need to use it, without satisfying correctness
- Use a log to record all actions
 - ► Employ the Write-Ahead Logging protocol

Summary (cont.)

- Use *checkpoints* to *periodically record consistent states* and *limit* the amount of the *log* that needs to be *scanned during recovery*
- Recovery in three phases
 - ► Analysis: from checkpoint, figure out REDO and UNDO extents
 - ► REDO: repeat entire history
 - ► UNDO: delete effects of failed transactions
- Repeating history simplifies the logic

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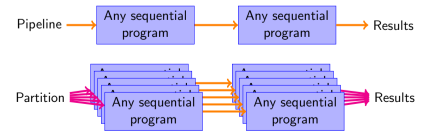
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Parallel data management

Parallelism and DBMSs

- Parallelism is natural to DBMS processing
 - Pipeline parallelism: many machines each doing one step in a multi-step process
 - ► Partition parallelism: many machines doing the same thing to different pieces of data.
 - ▶ Both are natural in a DBMS



Partitioning: split inputs, merge outputs

Why parallelism?

- The very *definition* of *parallelism*: *divide* a big *problem* into many *smaller* ones to be *solved in parallel*
- Consider we have a *terabyte* of data to *scan*
 - ▶ With *one pipe* of 10MB/s, we need 1.2 days
 - ▶ By partitioning the data in disjoint subsets and having 1,000 parallel pipes of the same bandwidth, we need 90s

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Parallel data management

The parallelism success story

- DBMSs are the most (only?) successful application of parallelism
 - ► Teradata, Tandem vs. Thinking Machines, KSR, ...
 - ► Every *major DBMS vendor* has some *parallel server*
 - Workstation manufacturers now depend on parallel DB server sales
- Reasons for success
 - ► Bulk-processing (partition parallelism)
 - ► Natural pipelining
 - ► Inexpensive hardware can do the trick
 - ► Users/app-programmers do *not* need to *think in parallel*

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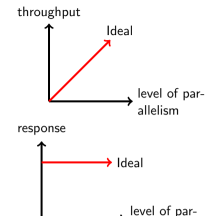
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dvanced Database

Terminology

Speed-up

More resources means proportionally less time for given amount of data (throughput)



Scale-up

If resources increased in proportion to increase in data size, time is constant

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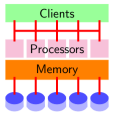
allelism

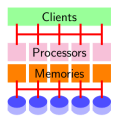
Parallel data management

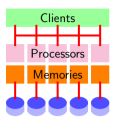
Different types of parallelism

- Intra-operator parallelism
 - ► All machines working to compute a single operation (scan, sort, join)
- Inter-operator parallelism
 - ► Each operator may run concurrently on a different site (exploits pipelining)
- *Inter-query* parallelism
 - ► Different gueries run on different sites
- We shall focus on intra-operator parallelism

Architecture: what to share?







Shared memory

- Easy to program
- Expensive to build
- Difficult to scale up

Shared disk

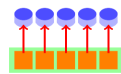
- *Middle* of the road
- Distributed file system
- *Cluster* computing

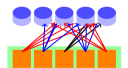
Shared nothing

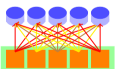
- Hard to program
- Cheap to build
- Easy and ideal to speed/scale up

Parallel data management

Automatic data partitioning







Range

- Good for equi-joins
- Range-queries
- Good for aggregation

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Hash

- Good for equi-joins
- No range-queries
- Problematic with skew

Round-robin

- Indifferent for equi-joins
- Range-queries complicated
- Load-balanced

Parallel scans

- Scan in parallel, and merge
- Selections may not require all sites for range or hash partitioning
- Indexes can be built at each partition
- Question: how do indexes differ in the different schemes?
 - ► Think about **both** lookups and inserts!
 - ▶ What about *key* indexes?

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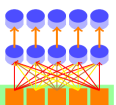
Parallel data management

Parallel aggregation

- For each aggregate function, need a decomposition
 - $count(S) = \sum_{i} count(s(i)), ditto for sum()$
 - $\rightarrow avg(S) = (\sum_{i} sum(s(i))) / \sum_{i} count(s(i))$
 - ▶ and so on ...
- For groups
 - ► Sub-aggregate groups close to the source
 - ▶ Pass each sub-aggregate to its group's site
 - ★ Chosen via a hash function

Parallel sorting

- Key idea: sorting *phases* are intrinsically parallelisable
 - ► Scan in parallel, range-partition as you go
 - ► As *tuples come in*, begin "local" sorting using standard algorithm
 - ► Resulting data is sorted, and range-partitioned
- Problem: skew
 - ► Solution: *sample* the data to determine partition points



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Parallel data management

Parallel joins

Nested loops

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- ► Each outer tuple must be compared with each inner tuple that might
- ► Easy for range partitioning on join columns, hard otherwise
- Sort-merge (or plain merge-) join
 - ► Sorting gives range-partitioning
 - Merging partitioned tables is local

Parallel hash join

- During the first phase, partitions are distributed to different sites
 - ► A good *hash function automatically* distributes work *evenly*
- Second phase is local at each site
 - ► Almost *always* the *winner* for *equi-join*
- Good use of split/merge makes it easier to build parallel versions of sequential join code

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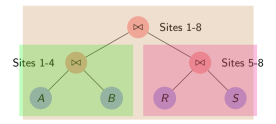
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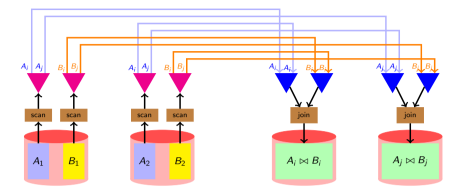
Parallel data management

Complex parallel query plans

- Complex queries: inter-operator parallelism
 - ► *Pipelining* between operators
 - * Note that *sorting* and *phase one* of *hash-join block* the pipeline (yet again!)
 - ► *Bushy* execution trees



Dataflow network for parallel join



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Parallel data management

Observations

- It is relatively easy to build a fast parallel query executor
- It is hard to write a robust and world-class parallel query optimizer
 - ► There are many *tricks*
 - ► One quickly hits the *complexity barrier*
 - ► Still open research

Parallel data management

Parallel query optimization

- Common approach: two phases
 - ► Pick best sequential plan (System R algorithm)
 - ▶ Pick *degree of parallelism* based on current system parameters
- Allocate operators to processors
 - ► Take *query tree*, *decorate* as in previous example

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Parallel data management

Parallel databases summary

- Parallelism natural to query processing
 - ▶ Both *pipeline* and *partition parallelism*
- Shared-nothing vs. Shared-memory
 - ► Shared-disk too, but less standard
 - ► Shared-mem easy, costly; does not scaleup
 - ► Shared-nothing cheap, scales well, harder to implement
- Intra-operator, inter-operator, and inter-query parallelism all possible.

What can go wrong?

- Best sequential plan \neq best parallel plan
- Trivial counter-example
 - ► Table partitioned with local secondary index at two nodes
 - ► Range query: all of node 1 and 1% of node 2
 - ★ e.g., select * from telephone_book where name < "NoGood"
 - ▶ Node 1 should do a scan of its partition
 - ► Node 2 should use secondary index

Table scan Index scan

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Parallel data management

Parallel database summary (cont.)

- Data layout choices important
- Most database operations can be done using partition-parallelism
 - Sort
 - ► Sort-merge join, hash-join
- Complex plans
 - ► Allow for *pipeline-parallelism*, but sorts, hashes *block* the *pipeline*
 - ► Partition-parallelism achieved through bushy trees

Parallel database summary (cont.)

- Hardest part: optimization
 - ► Two-phase optimization simplest, but can be ineffective
 - ► More *complex schemes* still at the *research* stage
- We have not discussed transactions, logging
 - ► Easy in shared-memory/shared-disk architecture
 - ► Takes *some care* in *shared-nothing*
 - ► Some ideas from *distributed transactions* are *handy*

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